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# Abnormal Physiological and Molecular Mutant Phenotypes Link Chloroplast Polynucleotide Phosphorylase to the Phosphorus Deprivation Response in Arabidopsis<sup>1[C][W][OA]</sup>

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A prominent enzyme in organellar RNA metabolism is the exoribonuclease polynucleotide phosphorylase (PNPase), whose reversible activity is governed by the nucleotide diphosphate-inorganic phosphate ratio. In *Chlamydomonas reinhardtii*, PNPase regulates chloroplast transcript accumulation in response to phosphorus (P) starvation, and PNPase expression is repressed by the response regulator PSR1 (for PHOSPHORUS STARVATION RESPONSE1) under these conditions. Here, we investigated the role of PNPase in the Arabidopsis (*Arabidopsis thaliana*) P deprivation response by comparing wild-type and *pnp* mutant plants with respect to their morphology, metabolite profiles, and transcriptomes. We found that P-deprived *pnp* mutants develop aborted clusters of lateral roots, which are characterized by decreased auxin responsiveness and cell division, and exhibit cell death at the root tips. Electron microscopy revealed that the collapse of root organelles is enhanced in the *pnp* mutant under P deprivation and occurred with low frequency under P-replete conditions. Global analyses of metabolites and transcripts were carried out to understand the molecular bases of these altered P deprivation responses. We found that the *pnp* mutant expresses some elements of the deprivation response even when grown on a full nutrient medium, including altered transcript accumulation, although its total and inorganic P contents are not reduced. The *pnp* mutation also confers P status-independent responses, including but not limited to stress responses. Taken together, our data support the hypothesis that the activity of the chloroplast PNPase is involved in plant acclimation to P availability and that it may help maintain an appropriate balance of P metabolites even under normal growth conditions.

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Organisms require phosphorus (P) continually and in relatively high amounts, and in photosynthetic systems a major use is the regeneration of ribulose-1,6-bisphosphate, the acceptor for CO<sub>2</sub> fixation by Rubisco. Chloroplast inorganic phosphate (Pi) pools are also affected by starch biosynthesis, since the conversion of Glc-1-P to ADP-Glc, the penultimate step in the pathway, releases Pi through ATP hydrolysis. Starch is primarily synthesized during the day from excess triose phosphates and broken down at night, a step that consumes Pi through the action of starch phosphorylase and other enzymes (Zeeman et al., 2007). Thus, P is a major player in chloroplast metabolism and is intimately integrated into the carbon budget of plant cells.

Chloroplast P is also required for processes not directly related to photosynthesis, such as gene expression. In particular, the chloroplast ribonuclease polynucleotide phosphorylase (PNPase) both consumes and liberates P. PNPase in bacteria (Soreq and Littauer, 1977) and chloroplasts (Baginsky et al., 2001) is a homotrimer that degrades RNA through phosphorylytic attack, but it also readily synthesizes polynucleotides using nucleotide diphosphate or nucleotide triphosphate precursors, a reaction that generates Pi or inorganic pyrophosphate, respectively. The purified

chloroplast enzyme, like its bacterial counterpart, is readily reversible in vitro (Yehudai-Resheff et al., 2001). Several lines of work have clearly placed PNPase, in both prokaryotes and eukaryotic organelles (for review, see Slomovic et al., 2006), as a key player in a polyadenylation-stimulated RNA degradation pathway found in plant mitochondria and chloroplasts.

While the role of PNPase in RNA metabolism is established if incompletely understood, newer evidence has linked PNPase to somewhat unrelated functions. In human cells, for example, a cytosolic fraction of mitochondrial PNPase appears to influence cell differentiation and senescence (Sarkar and Fisher, 2006), and in the mitochondrion itself, PNPase is located in the intermembrane space while mitochondrial RNA is in the matrix, suggesting a metabolic rather than an RNA catalytic role for PNPase (Chen et al., 2006), through which it can nonetheless influence the accumulation of certain mitochondrial RNAs (Slomovic and Schuster, 2008). More closely related to this study, a genetic screen carried out in *Arabidopsis thaliana* for resistance to fosmidomycin, which inhibits the plastid methylerythritol phosphate (MEP) pathway, identified a chloroplast (cp)PNPase null mutant that was named *rif10* (Sauret-Gueto et al., 2006). This led to the suggestion that the MEP pathway might be regulated by plastid metabolic cues, which in turn could be influenced by PNPase.

This study was stimulated by our finding that in the green alga *Chlamydomonas reinhardtii*, reduced expression of cpPNPase rendered cells unable to acclimate to P deprivation, whereas the same strains had wild-type responses to other nutrient or environmental stresses (Yehudai-Resheff et al., 2007). We also found that both the PNPase transcript and protein were repressed under P deprivation, suggesting that reduced PNPase activity is part of the overall metabolic adjustment to P limitation. Furthermore, repression of PNPase expression required the general P deprivation response regulator *Psr1*, a likely transcription factor (Wykoff et al., 1999), demonstrating that PNPase regulation is integrated into the global P limitation response. The ortholog of *Psr1* in *Arabidopsis* is *PHR1* (Rubio et al., 2001), among whose functions is inducing microRNAs of the miR399 family, which through long-distance signaling (Pant et al., 2008) derepress a suite of genes including some encoding P transporters, through a ubiquitination pathway involving *PHO2* (Aung et al., 2006; Bari et al., 2006; Chiou et al., 2006). This pathway in turn is modulated by the noncoding RNA *IPS1* (Franco-Zorrilla et al., 2007) and possibly others.

Here, we examine the role of cpPNPase in *Arabidopsis* in the context of the P deprivation response using null mutant alleles. The most obvious growth defect under P limitation is in the elaboration of lateral roots, where the *pnp* mutant phenocopies *pdr2*, a mutant thought to define a signal needed for lateral root proliferation (Ticconi et al., 2004). When gene expression was examined using array technology, we found that the *pnp* mutants already induced, under normal

growth conditions, genes normally only induced upon P deprivation. These and other data suggest that *pnp* mutants exhibit elements of P starvation even when grown under P-replete conditions. Therefore, we propose that cpPNPase in plants, as in unicellular algae, has a key role in P metabolism.

## RESULTS

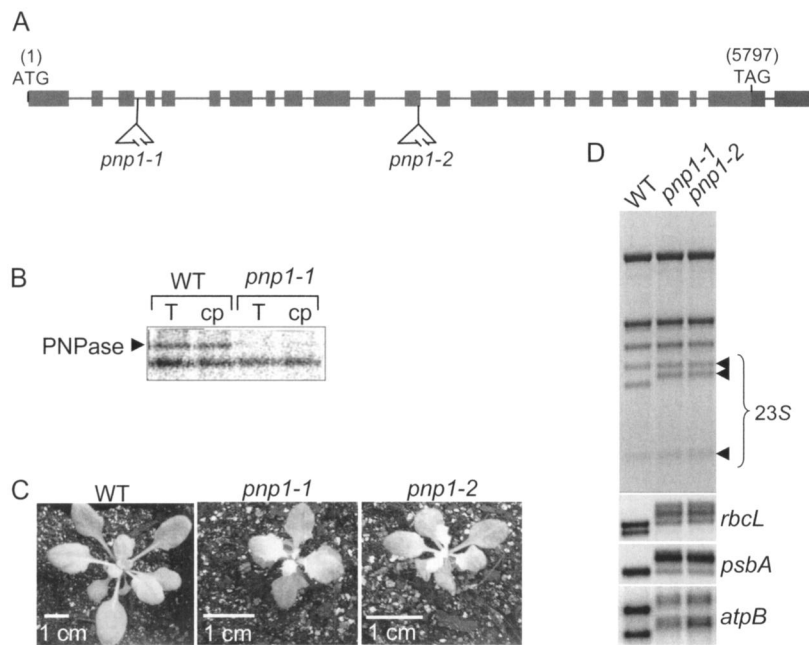
### PNP T-DNA Insertion Mutants

cpPNPase is encoded by the locus At3g03710, which specifies a 922-residue protein, consistent with the approximately 100-kD migration of PNPase previously described from spinach (*Spinacia oleracea*) and pea (*Pisum sativum*) chloroplasts (Hayes et al., 1996; Li et al., 1998). Limited data have been obtained on *Arabidopsis* lines reduced or totally deficient for cpPNPase (Walter et al., 2002; Sauret-Gueto et al., 2006). To pursue the analysis of *Arabidopsis* cpPNPase in the context of the nutrient stress response, we selected two T-DNA insertion alleles, naming them *pnp1-1* and *pnp1-2* (Fig. 1A). Homozygous mutants for each line were isolated following three outcrosses of the original T3 plants. UV cross-linking was performed to confirm that the *pnp1-1* mutant lacked cpPNPase, shown in Figure 1B as an approximately 100-kD RNA-binding activity. The mutants also lack complete *PNP* transcripts, as revealed by reverse transcription (RT)-PCR using primers flanking the respective T-DNA insertions (data not shown).

Figure 1C compares wild-type and *pnp* mutant plants after 21 d of growth on soil. The rosette leaves of *pnp* mutants emerged and remained pale green compared with those of wild-type plants, as was observed previously (Sauret-Gueto et al., 2006). In addition, the mutant plants were retarded in silique production, and both silique number and seed production were reduced (data not shown). Thus, cpPNPase is required for fully normal plant development, although in its absence a full life cycle is achieved. Another feature of cpPNPase deficiency is incomplete 3' end processing of certain chloroplast transcripts, including 23S rRNA and various mRNAs. These are exemplified in Figure 1D, and in the cases of 23S rRNA, *rbcL*, and *psbA*, the presence of 3' extensions had been confirmed previously (Walter et al., 2002).

### Lateral Root Development under P Limitation

It is well established that when faced with nutrient limitation, plants seek additional sources through altered root system architecture. In the case of P limitation, both root hair density and lateral root proliferation are observed (for review, see Lopez-Bucio et al., 2003). To see whether PNPase deficiency affected this process, we compared the root architecture of wild-type and mutant plants grown under +P and -P conditions. As exemplified in Figure 2, there were no substantive differences observed when several genotypes were com-

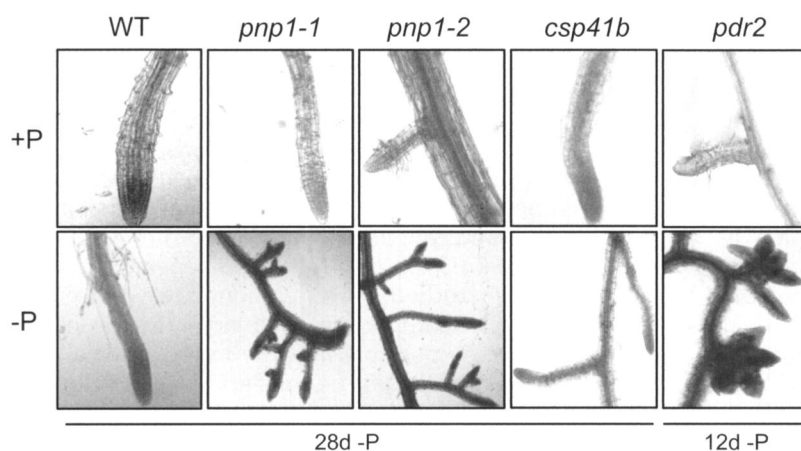


**Figure 1.** Characterization of *pnp1-1* and *pnp1-2* T-DNA insertion mutants. A, Diagram of the two T-DNA insertions in the *PNP* gene corresponding to the lines named *pnp1-1* and *pnp1-2* (SALK\_013306 and SALK\_070705, respectively). B, Total (T) or chloroplast (cp) proteins extracted from the wild type (WT) and the *pnp1-1* mutant were analyzed for RNA binding by the UV cross-linking assay as described in "Materials and Methods." The bottom band is an unknown RNA-binding protein whose signal is similar in the different samples. C, The wild type, *pnp1-1*, and *pnp1-2* homozygous mutants were grown on soil for 21 d. Note that the 1-cm bar size is different for the mutants. D, cpRNA patterns in the *pnp1-1* and *pnp1-2* mutants. Total RNA was isolated from the wild type, *pnp1-1*, and *pnp1-2* and analyzed by ethidium bromide staining to reveal 23S rRNA anomalies (top) or by gel blot using the chloroplast gene probes indicated at right. [See on-line article for color version of this figure.]

pared under +P conditions, although the *pnp* mutants had a slower overall root elongation rate (data not shown). We used two other mutants as controls. The first was *csp41b*, a nuclear mutant lacking two related chloroplast endoribonucleases (Bollenbach et al., 2009). We reasoned that if abnormal chloroplast RNA (cpRNA) metabolism had any pleiotropic effect on root architecture, that would be seen for both *pnp1-1* and *csp41b*. A second control was *pdr2*, a mutant that is known to exhibit abortive lateral root initiation under -P conditions (Ticconi et al., 2004). This control was useful because, as we discovered (Fig. 2, bottom row), the *pnp1-1* mutants phenocopied *pdr2*. In particular, when grown under -P conditions, the *pnp* mutants were unable to elaborate lateral roots, instead exhibiting a proteoid root-like phenotype after 4 weeks of P starvation. While the same effect was observed earlier for *pdr2*, it should be borne in mind that *pnp* mutants have an overall slower growth rate (Fig. 1).

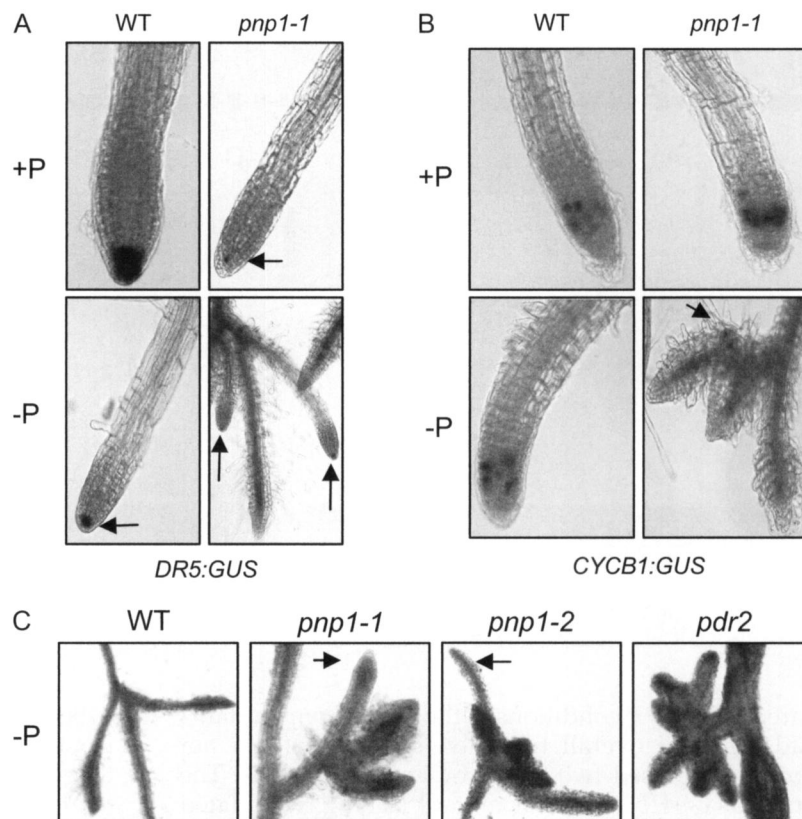
Because the *csp41b* mutant was indistinguishable from the wild type under -P conditions, we concluded that the aberrant root architecture phenotype of *pnp1-1/pnp1-2* was not related to altered cpRNA metabolism per se. Instead, we suspected that some function of PNPase was essential to lateral root elaboration. To investigate this phenomenon in more detail, we crossed the *pnp1-1* mutation into two backgrounds expressing GUS reporter genes, under the control of either the auxin-responsive promoter element DR5 (Ulmasov et al., 1997) or the cyclin B1 promoter (Colòn-Carmona et al., 1999), as an indicator of cell division. Both reporters have previously been used in studies of root physiology under P limitation conditions (Ticconi et al., 2004; Nacry et al., 2005; Sanchez-Calderon et al., 2005).

Figure 3, A and B, show results for plants expressing DR5:GUS and CYCB1:GUS, respectively. When grown on +P medium, *pnp1-1* exhibited considerably less staining in the primary root tip than did the wild type,



**Figure 2.** Lateral root phenotypes under P-replete conditions (top row) or under P deprivation (bottom row). Additional mutants used as controls were *csp41b*, which lacks two related chloroplast endoribonucleases (Bollenbach et al., 2009), and *pdr2*, which was previously demonstrated to exhibit the observed phenotype (Ticconi et al., 2004). For the experiment shown here, plants were grown on a full nutrient medium for 2 weeks, then transferred to +P or -P medium for the times indicated. WT, Wild type.

**Figure 3.** Cell division and mortality in lateral roots developing under P deprivation. A, Roots from plants expressing a *DR5:GUS* transgene were stained histochemically for GUS. Arrows point to smaller stained areas. Plants were germinated and grown for 2 weeks on +P medium, then transferred to +P or -P medium for 4 weeks. B, Roots from plants expressing a *CYCB1:GUS* transgene were stained histochemically, following growth as for A. C, Plants were grown as for A. Roots were stained with Evans blue. The arrows indicate *pnp* mutant root tips that are not stained. WT, Wild type. [See online article for color version of this figure.]

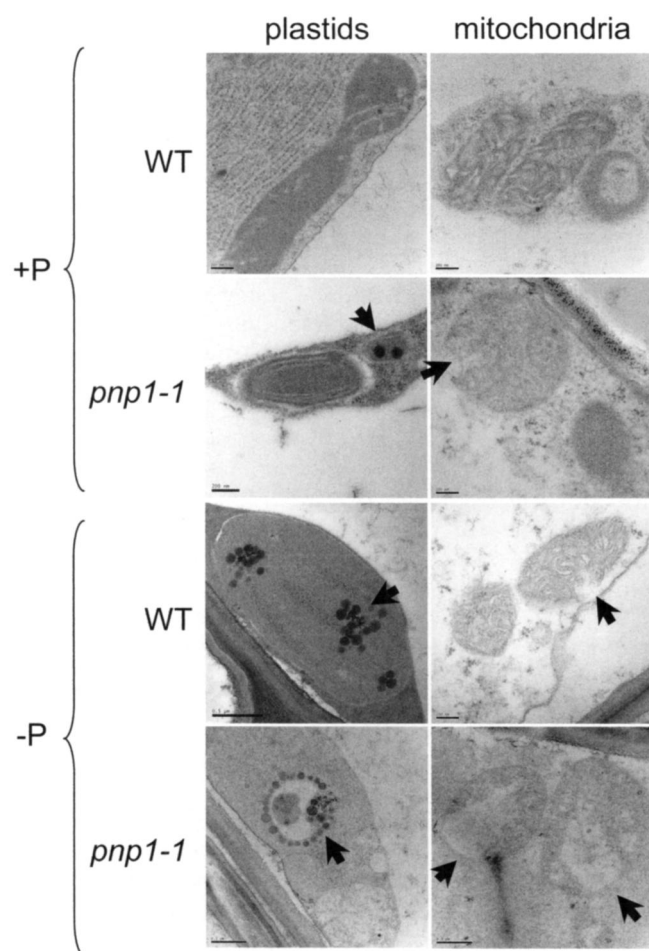


which is consistent with its slower overall root elongation rate. When grown on -P medium, the wild type exhibited reduced staining at the root tip, a result consistent with another study showing an age-dependent decrease *DR5:GUS* expression following P starvation as compared with seedlings grown on +P medium (Sanchez-Calderon et al., 2005). In the case of *pnp1-1*, we found that only one or two of the lateral root initiates in the proteoid-like clusters stained with GUS under -P conditions (Fig. 3A, arrows). This suggested that while some lateral roots in *pnp1-1* had a wild-type-like auxin environment, other roots in the clusters showed no evidence for this marker and were likely auxin deficient.

Concordant results were obtained with the cyclin marker (Fig. 3B). As observed in another study (Ticconi et al., 2004), a zone of mitotic cells was present in the root meristem under both +P and -P for the wild type, and this was also the case for *pnp1-1* when grown under +P conditions. When *pnp1-1* was starved for P, however, only a single early-stage initiate in the root clusters stained for GUS (Fig. 3B, arrow). This suggested that cell division had ceased in the majority of lateral root initiates, the same phenomenon that was documented for *pdr2* (Ticconi et al., 2004). Since the *pnp* mutant developed root clusters where the majority of roots appeared to lack both auxin and cell division, we used the dye Evans blue to see if cell death had occurred. As shown in Figure 3C, in the cases of both *pnp1-1* and *pdr2*, the majority of roots in P limitation-stimulated clusters stained with Evans blue, which

cannot pass intact membrane barriers. Thus, the majority of these cells are dead, and the affected root tips likely correspond to roots that also failed to stain with GUS driven by *DR5* or *CYCB1* promoters. In conclusion, *pnp1-1* exhibits abortive lateral root initiation under -P conditions, where the aborted roots cease division and undergo cell death.

Because PNPase is a plastid protein, we examined organellar ultrastructure in roots under +P and -P conditions, as exemplified in Figure 4. Under +P conditions, we observed an increased frequency of plastoglobules in *pnp1-1* as well as apparently ruptured mitochondria. Plastoglobules are associated with senescence and stress conditions and may be a pleiotropic consequence of the *pnp* mutation. The frequency of plastoglobules increased more dramatically in *pnp1-1* than in the wild type under -P conditions, also consistent with an enhanced stress response. In 60% of plastids in the mutant, they were arranged in a circle, as shown in Figure 4. We also noted increased mitochondrial disruption in the mutant, and occasionally in the wild type, under -P conditions. Mitochondrial rupture, as measured by cytochrome *c* release, has been associated with cell death in male-sterile sunflower (*Helianthus annuus*) anthers (Balk and Leaver, 2001) and subsequently studied in other contexts where programmed cell death occurs in plants (Reape et al., 2008). Thus, it is possible that the mitochondrial disruption observed in *pnp1-1* under -P conditions is symptomatic of the loss of cell integrity ultimately observed (Fig. 3C).



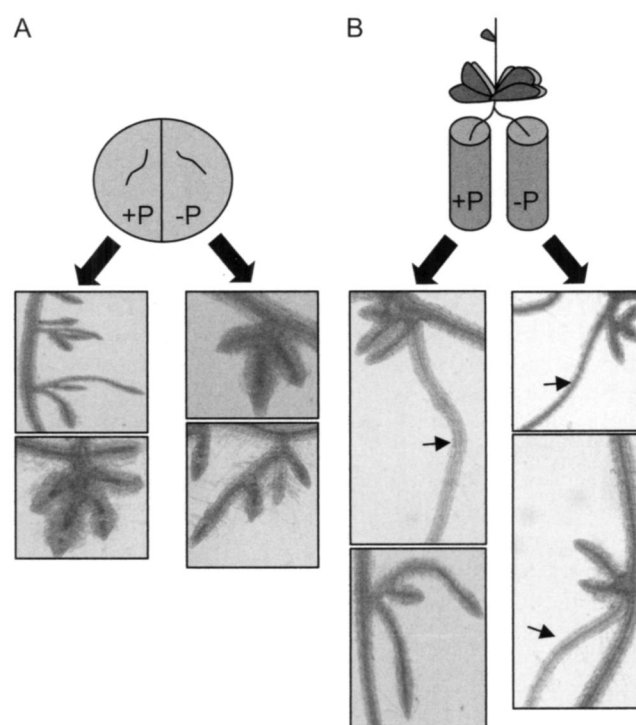
**Figure 4.** Ultrastructure of wild-type (WT) and *pnp1-1* mutant lateral root cortical cells. Plants were germinated and grown for 2 weeks on +P medium, then transferred to +P or -P medium. Plants were observed after 1, 2, 3, and 4 weeks on -P. Here, only images of root organelles after 1 week on -P are shown. Cortical cells of root tip sections in the longitudinal direction were observed using transmission electron microscopy as described in "Materials and Methods." Bars = 200 nm, except for *pnp1-1* on -P (plastids and mitochondria) and for the wild-type plastid on -P, where bars = 500 nm. Arrows indicate features referred to in the text.

The *PNP* gene is expressed in both roots and leaves (<http://mpss.udel.edu/at/>; S. Yehudai-Resheff and D.B. Stern, unpublished RT-PCR data), albeit at a much lower level in roots, raising the question of whether *pnp* mutants display an altered root architecture under -P conditions because of PNPase deficiency in roots, or in leaves, or both. To address this question, we carried out the experiments shown in Figure 5. We first excised roots from *pnp1-1* plants already exposed to -P conditions to observe whether providing them with P would correct the root architecture deficiency. Figure 5A shows that aberrant roots did not regain normal proliferation when placed on +P medium, although some growth occurred, suggesting that the P level encountered by the roots did not alone determine their phenotype. In contrast, when root masses from

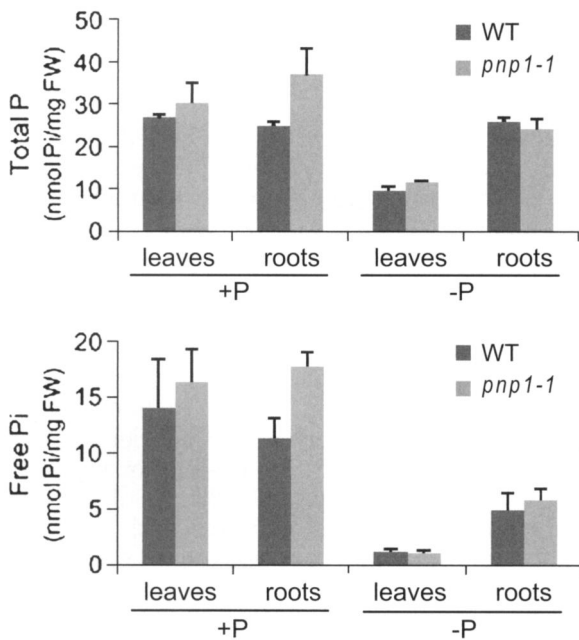
*pnp1-1* plants starved for P were divided and half placed on +P medium and the other half on -P medium, normal root elongation ensued in both samples (Fig. 5B). Thus, although half the *pnp* roots were in -P medium, P provided from the other roots and cycled through the aboveground tissues rescued the mutant phenotype. We conclude that altered root architecture in *pnp1-1* is not a cell-autonomous feature of root cells.

#### Metabolite Comparison of *pnp1-1* and the Wild Type

Based on the data described above, we hypothesized that PNPase deficiency either directly or indirectly affected production or transduction of a signal required for normal response to P deficiency. We first measured free or total P in wild-type or *pnp* mutant leaves and roots grown under +P and -P conditions. As shown in Figure 6, both free and total leaf P declined in both wild-type and mutant plants grown under -P conditions relative to +P, as was expected. For *pnp1-1*, we found that P levels were slightly higher in leaves, and significantly higher in roots, than those of the wild type when the plants were grown under +P conditions. On the other hand, P levels in *pnp1-1*



**Figure 5.** Split-root experiment using the *pnp1-1* mutant. A, Plants were germinated and grown for 2 weeks on +P medium and transferred to -P conditions for 3 weeks. The root segments were excised and placed on -P or +P agar medium. Roots were photographed after 2 additional weeks on these media. B, Plants were germinated and grown for 2 weeks on +P medium and transferred to -P conditions for 3 weeks. The roots were then divided into -P or +P liquid medium while remaining attached to the plants and grown for an additional 2 weeks. The arrows indicate lateral roots that have recovered a wild-type-like phenotype.



**Figure 6.** P content in leaves and roots. P content was measured in wild-type (WT) and *pnp1-1* mutant seedlings. Plants were germinated and grown for 2 weeks on +P medium, then transferred to -P or +P medium for 1 additional week. Free Pi and total P were assessed as described in "Materials and Methods." Error bars correspond to SE. FW, Fresh weight.

plants grown under -P conditions did not differ from those found in the wild type. We conclude that the *pnp1-1* mutation does not have a major effect on P content as related to fresh weight, although both leaf and root P were slightly elevated under +P conditions. We also measured P uptake in wild-type and mutant roots, in case differences in P uptake were masked in the P accumulation data. However, no significant differences were observed (Supplemental Fig. S3).

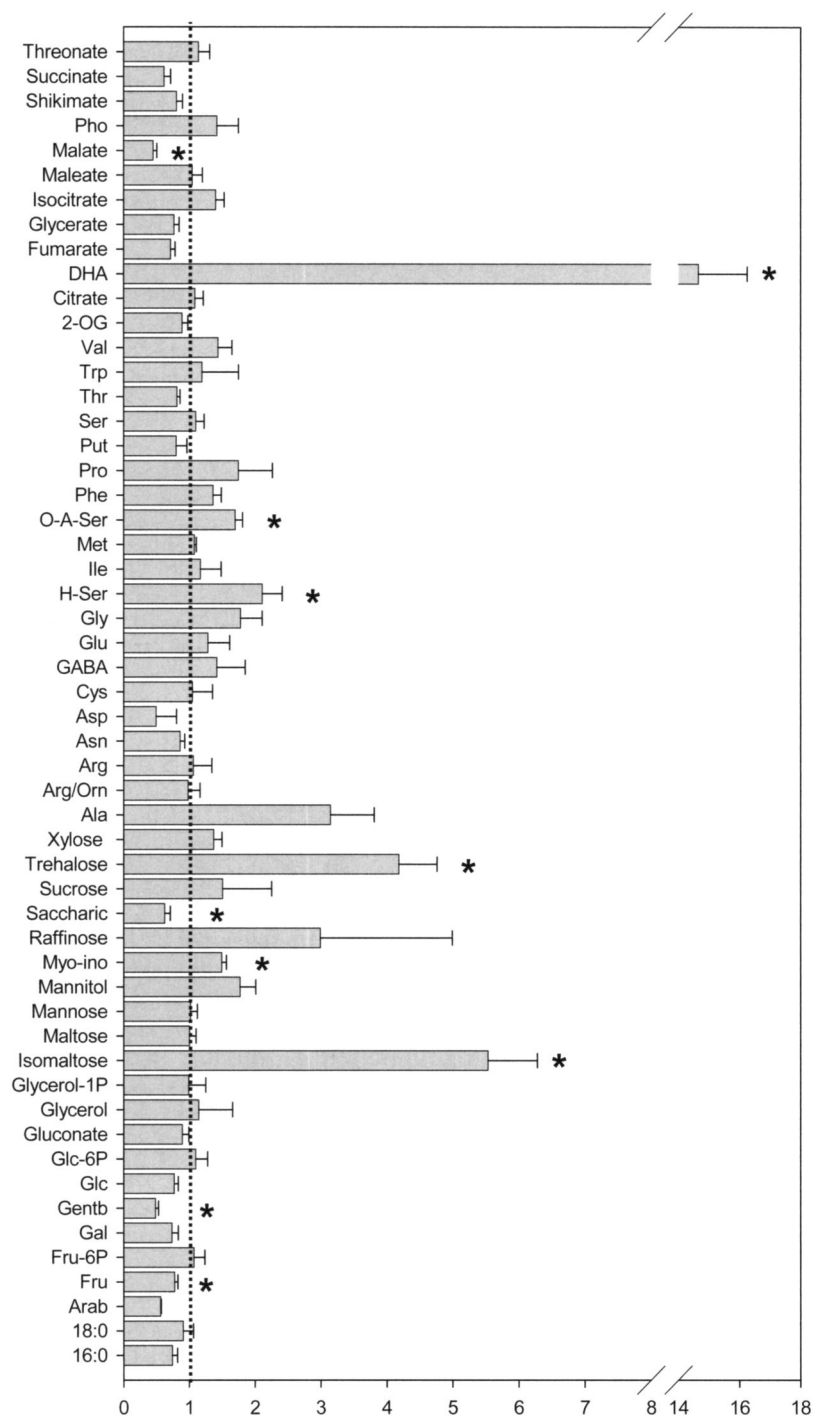
For a broader view of metabolic status, we quantified a panel of soluble, primary metabolites (Roessner et al., 2001) from wild-type and mutant plants grown under +P conditions or starved between 3 h and 3 weeks (Supplemental Tables S1 and S2; Supplemental Figs. S1 and S2). Under +P conditions, the metabolic differences between mutant and the wild type were limited (Fig. 7; Supplemental Table S1). Nonetheless, the sugars trehalose and isomaltose accumulated in the mutant to levels greater than 4-fold higher than in the wild type, whereas saccharate, gentiobiose, and Fru were present at significantly lower levels in the mutant. The mutant was additionally characterized by higher levels of O-A-Ser, H-Ser, and Ala. The levels of succinate, malate, and fumarate were considerably lower in the mutant than in the wild type, whereas dehydroascorbate (DHA) accumulated (although it is important to note that the DHA level measured here is not absolutely representative of the *in vivo* amount, since the extraction conditions used here do not exactly conserve the cellular redox poise of the ascorbate pools).

When looking from a global level, the impact of P starvation was fairly similar in both genotypes. The wild type displayed metabolic responses resembling those of many previous studies (Pieters et al., 2001; Uhde-Stone et al., 2003; Liu et al., 2005; Misson et al., 2005; Hernandez et al., 2007; Karthikeyan et al., 2007; Muller et al., 2007), while the *pnp* mutant generally also displayed many of these characteristic responses. Among observed differences was a far less dramatic change in the *pnp* mutant for the levels of Fru-6-P and Glc-6-P. Moreover, isomaltose, Man, saccharate, Suc, trehalose, and Xyl all increased in the wild type in response to P starvation, with raffinose being the only sugar that increased in both genotypes. Organic acids displayed mixed behavior in response to P deficiency: tricarboxylic acid (TCA) intermediates downstream of the reaction catalyzed by isocitrate lyase (one of the key enzymes of nitrate assimilation; Hodges et al., 2003) decreased in both genotypes, while those upstream increased. Taken together, these results suggest an altered TCA cycle activity on P stress and, as such, are largely in keeping with those of other recent studies (Morcuende et al., 2007).

With the exception of  $\gamma$ -aminobutyrate (GABA) and Pro, amino acids tend to accumulate with increasing periods of starvation in both genotypes. GABA, a stress-related metabolite (Fait et al., 2008), increased upon P starvation in the mutant, despite decreasing in the wild type, while the increase in the plastidial Asp family was visible following a mere 3 h of starvation in the mutant but only after 1 week in the wild type. Indeed, the changes in a wide range of amino acids were exacerbated in the mutant lines. To assess this statistically, two-way ANOVA tests were conducted (Supplemental Table S2); these tests revealed that the majority of the metabolites that discriminated between the genotypes were amino acids; however, also included in the top 25 discriminating metabolites were the phosphorylated intermediates Glc-6-P, glycerol 1-phosphate, and Fru-6-P, as expected (Morcuende et al., 2007, and refs. therein), and a range of sugars. These results were somewhat surprising, since the pattern of changes in many of these metabolites appears conserved between the genotypes; however, this clearly reflects that the altered metabolic response of *pnp1-1* is quite subtle. When studied from a functional rather than a chemical perspective, this list also revealed a high number of stress-related metabolites such as GABA, myoinositol, raffinose, gentiobiose, chlorogenate, DHA, and salicylate, which tended to respond more dramatically in the *pnp* mutant than in the wild type.

#### Transcriptome Characterization of *pnp1-1*

To gain further insight into how the lack of PNPase might affect plant responses to P starvation, we used microarray hybridizations to compare the transcriptomes of *pnp1-1* and wild-type plants grown under +P or -P conditions. To be able to relate transcriptome



**Figure 7.** Relative metabolite content of *pnp1-1* and the wild type grown under +P conditions expressed as mean fold change ( $\pm$ SE) to the wild type. Plants were grown for 2 weeks on +P medium, then transferred (at time 0) to +P or -P medium for 3 h, 6 h, 1 week, or 3 weeks. Metabolites were quantified as described in "Materials and Methods." The complete data set and statistical analysis are presented in Supplemental Table S2. Data are normalized with respect to the internal standard and the fresh weight. Values are presented as means  $\pm$  SE of determinations on six samples of bulked seedlings. Asterisks indicate values that were significantly different from the wild type when assessed by *t* tests ( $P < 0.05$ ).

data to the metabolite analysis, we chose two time points for which the two genotypes were most distinct, which were after 3 h and 1 week of P starvation, according to principal component analysis (Supplemental Fig. S2). Total rosette RNA was used with the Affymetrix ATH1 platform to facilitate comparison with previous studies. The threshold of 2-fold change was chosen with a false discovery rate (FDR)  $< 0.05$ . Supplemental Tables S3 and S4 include the full data sets of the 3-h and 1-week -P experiments, respectively. Because the 3-h experiment revealed no signif-

icant regulated genes in the wild type or *pnp1-1* in response to P starvation, we will only discuss the 1-week experiment here.

We first compared the two genotypes grown on +P medium (Table I). A total of 960 genes were found to be differentially expressed, which is perhaps not surprising given the slow growth and partial chlorosis of *pnp1-1*. What was noteworthy, however, was that the number of differentially expressed genes dramatically decreased when plants were grown on -P medium, to 224, indicating that the wild type and *pnp1-1* have more



**Table I.** The number of regulated genes in the wild type and/or *pnp1-1* under different growth conditions

Experiment <sup>a</sup>	Data <sup>b</sup>	Induced <sup>c</sup>	Repressed <sup>c</sup>	Total
<i>pnp1-1</i> +P/wild type +P	16,536	601	359	960
Wild type -P/wild type +P	16,660	409	99	508
<i>pnp1-1</i> -P/ <i>pnp1-1</i> +P	16,585	174	35	209
<i>pnp1-1</i> -P/wild type -P	16,751	154	70	224

<sup>a</sup>The genotype and the condition listed first correspond to the numerator of the gene expression mean ratio relative to the second, which is the denominator. For example, 601 genes are induced in the *pnp1-1* mutant when compared with the wild type under +P conditions.

<sup>b</sup>Data correspond to the genes that were tagged as "present" in at least one replicate as described in "Materials and Methods." <sup>c</sup>The induced or repressed genes exhibit at least 2-fold change in expression and FDR < 0.05.

similar gene expression patterns when P is not provided.

Subsequently, the effect of P deprivation on gene expression was assessed in the wild type. We found that 508 genes were regulated: 80.5% were up-regulated, whereas less than 20% were repressed. We compared our data set with other published experiments and found general consistency, although the experimental protocols were not identical (Misson et al., 2005; Morcuende et al., 2007). A third step of the analysis was to examine the effect of P deprivation in the mutant, and we found that the number of P-regulated genes in *pnp1-1* was dramatically reduced with respect to the wild-type (i.e. 209 versus 508; Table I). This is consistent with the data mentioned above, that 960 genes differentiate the two genotypes on +P but only 224 on -P. In terms of transcript regulation, we conclude that P starvation attenuates the differences between *pnp1-1* and the wild type.

#### Classification of Differentially Expressed Genes

We used MapMan to determine which functional categories were most affected in various pairwise comparisons (Supplemental Table S5). When the wild type and *pnp1-1* were compared on +P conditions, the categories of photosynthesis, RNA regulation, cell functions, and stress responses were all identified with a *P* value of  $10^{-10}$  or less, and various other metabolic functions were also identified with low *P* values ( $10^{-4}$  or less). Together, these categories likely include pleiotropic effects (e.g. stress responses) but may also include metabolic functions related to a particular role of PNPase. Under -P conditions and using a *P* value of  $10^{-4}$  as a cutoff, eight categories rather than 11 (as seen under +P conditions) were identified, consistent with the speculation above that P starvation attenuates differences between the wild type and mutant. Of these eight categories, only two were the same as for +P conditions (photosynthesis and mitochondrial metabolism), consistent with the known large-scale reprogramming of plant gene expression when facing abiotic stress. The eight categories also included three important metabolic networks: major carbohydrate metabo-

lism, oxidative pentose phosphate pathway, and TCA cycle/organic acid transformations. Thus, under P starvation, the *pnp* mutation affects normal gene expression as related to several major organellar and metabolic functions.

Another form of comparison was done, where we identified the 40 most regulated genes in *pnp1-1* versus the wild type when grown under +P conditions (Table II), 15 of which are related to defense or stress responses. Four genes related to photosynthesis are also strongly regulated. Most remarkable among them is *petD*, which encodes subunit IV of the cytochrome *b<sub>6</sub>/f* complex and is induced 100-fold or greater under both +P and -P conditions in the mutant. However, like other chloroplast-localized genes in *pnp1-1*, altered mRNA processing is likely to account for some of the observed increase, and furthermore, as our cDNA was primed with oligo(dT), we would amplify polyadenylated chloroplast transcripts, which have been reported to hyperaccumulate in PNPase-deficient Arabidopsis plants (Walter et al., 2002). In contrast, two nuclear genes encoding photosynthesis proteins are strongly repressed in *pnp1-1* versus the wild type; these are *PSBP2* (PSII oxygen-evolving enhancer) and *PORA* (chlorophyll biosynthesis).

#### P-Independent PNPase-Regulated Genes

We conducted further analysis to differentiate pleiotropic effects on gene expression related to the slower growth, chlorosis, and possible general stress responses of the *pnp1-1* mutant from those that could be directly attributed to the effect of the *pnp* mutation on chloroplast metabolism as possibly related to P deprivation responses. To do so, we compared sets of regulated genes as shown in Figure 8.

Figure 8A shows the overlap between *pnp1-1* versus the wild type on +P or -P, revealing 149 genes that are regulated in the same direction. These can be interpreted as nutrient-independent effects of the *pnp* mutation. Representatives of these 149 genes are listed in Table III, with major categories related to photosynthesis or chloroplast functions. Fourteen genes are chloroplast encoded, and 11 of them are very strongly induced in *pnp1-1*, which as discussed above likely relates to the RNA maturation function of PNPase. A second cluster of genes included in the overlap in Figure 8A is conspicuous, as it mainly encodes components of the PYK10 complex, which is named after an endoplasmic reticulum body-associated  $\beta$ -glucosidase thought to be involved in plant defense (Nagano et al., 2005). Like the gene encoding PYK10 itself, genes encoding eight jacalin-lectin proteins, three GDLS lipase-like proteins, TSA1-like (DNA topoisomerase), and a meprin and TRAF homology domain-containing protein are all up-regulated in the *pnp* mutant. All of these proteins except three jacalins (JAL4, -11, and -27) were found to be part of the PYK10 complex (Nagano et al., 2008), and their induction is consistent with a stress situation in the *pnp* mutant.

**Table II.** The 40 most up- and down-regulated genes in the *pnp1-1* mutant relative to the wild type when grown under +P conditions (FDR > 0.05) ns, The fold change is below the cutoff value of 2. Stress-related gene products are shown in boldface.

Probe Set Identifier	Arabidopsis Genome Initiative Code	Description	<i>pnp1-1</i> versus the Wild Type	
			Fold Change on +P	Fold Change on -P <sup>a</sup>
244977_at	AtCg00730	PetD, subunit IV of cytochrome <i>b<sub>6</sub>/f</i> complex	102.82	151.68
257673_at	At3g20370	Meprin and TRAF homology domain-containing protein	81.11	15.88
250515_at	At5g09570	Expressed chloroplast protein	39.15	1.61
263539_at	At2g24850	<b>Aminotransferase</b>	32.66	9.07 <sup>b</sup>
267472_at	At2g02850	Plastocyanin-like domain-containing protein	29.93	3.28 <sup>b</sup>
245002_at	AtCg00270	PsbD, D2 protein of PSII	28.93	34.44
258941_at	At3g09940	<b>Monodehydroascorbate reductase, putative</b>	27.35	ns
248434_at	At5g51440	23.5-kD mitochondrial small HSP	26.05	11.35 <sup>b</sup>
265051_at	At1g52100	<b>Jacalin-lectin family protein</b>	26.05	17.67
246888_at	At5g26270	Expressed protein	26.02	18.81
261930_at	At1g22440	Alcohol dehydrogenase	24.83	ns
265668_at	At2g32020	GCN5-related <i>N</i> -acetyltransferase family protein	22.45	20.99
260522_x_at	At2g41730	Expressed protein	22.34	9.65
266752_at	At2g47000	P-glycoprotein 6 (PGP6), multidrug-resistant transporter	21.49	11.89
266246_at	At2g27690	<b>Cytochrome P450</b>	20.82	5.00 <sup>b</sup>
252921_at	At4g39030	<b>Enhanced disease susceptibility 5 (EDS5)</b>	20.69	6.70 <sup>b</sup>
256159_at	At1g30135	<b>JAZ8/TIFY5A (jasmonate-ZIM-domain protein 8)</b>	20.60	ns
254385_s_at	At4g21830	<b>Met sulfoxide reductase domain-containing protein</b>	20.56	ns
255110_at	At4g08770	<b>Peroxidase</b>	18.24	ns
263153_s_at	At1g54010	<b>Myrosinase-associated protein</b>	18.11	6.18
247718_at	At5g59310	<b>Lipid transfer protein 4 (LTP4)</b>	0.01	0.03
258498_at	At3g02480	<b>ABA-responsive protein-related</b>	0.03	ns
262412_at	At1g34760	<b>14-3-3 protein GF14 omicron (GRF11)</b>	0.03	ns
266516_at	At2g47880	Glutaredoxin family protein	0.05	ns
266098_at	At2g37870	Protease inhibitor/lipid transfer protein (LTP) family protein	0.07	ns
259789_at	At1g29395	<b>Stress-responsive protein</b>	0.08	0.10 <sup>b</sup>
247095_at	At5g66400	<b>Dehydrin (RAB18)</b>	0.08	0.04 <sup>b</sup>
248197_at	At5g54190	NADPH-protochlorophyllide oxidoreductase A (PORA)	0.1	0.08 <sup>b</sup>
263981_at	At2g42870	Phytochrome rapidly regulated 1 (PAR1)	0.1	ns
247717_at	At5g59320	<b>Lipid transfer protein 3 (LTP3)</b>	0.1	0.13 <sup>b</sup>
260831_at	At1g06830	Glutaredoxin family protein	0.11	0.24
245353_at	At4g16000	Expressed protein	0.11	ns
264041_at	At2g03710	Agamous-like 3 (AGL3)	0.12	0.1
266873_at	At2g44740	CYCP4;1, cyclin family protein	0.13	ns
250933_at	At5g03170	Fasciclin-like arabinogalactan protein (FLA11)	0.13	ns
252612_at	At3g45160	Expressed protein	0.14	0.07
267569_at	At2g30790	PSII oxygen-evolving complex 23 (PSBP2)	0.14	0.17
261226_at	At1g20190	Expansin (EXP11)	0.14	ns
251039_at	At5g02020	Expressed chloroplast protein	0.14	0.15
251196_at	At3g62950	Glutaredoxin family protein	0.15	ns

<sup>a</sup>The corresponding data for -P are also indicated (i.e. *pnp1-1* on -P versus the wild type on -P). A full data set is given in Supplemental Table S4. <sup>b</sup>These values are not significant (FDR > 0.05).

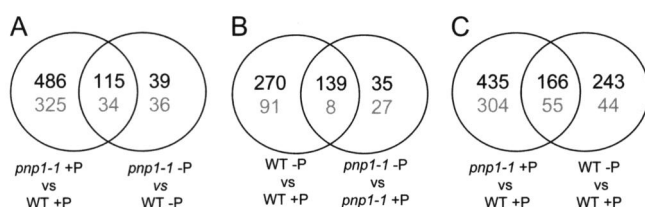
### Effect of the *pnp* Mutation on the P Starvation Response

In order to explore the apparent attenuation of P starvation-induced transcriptional responses in the *pnp1-1* mutant, we analyzed the overlap in regulated genes between *pnp1-1* -P versus +P and wild type -P versus +P (Fig. 8B). This revealed 147 genes whose expression changes in the same direction in the two genotypes, representing about 30% and 70% of wild-type and *pnp1-1* P-regulated genes, respectively. If one includes data below the 2-fold threshold we had chosen and allows an FDR > 0.5, 92% of the non-overlapping P-responsive genes in the wild type are also regulated by -P in the same way in *pnp1-1*.

Altogether, these data indicate that the wild type and *pnp1-1* possess largely parallel responses to P deprivation, but the average fold change of the 139 overlapping induced genes was globally lower for the *pnp1-1* plants (20.7-fold induction for *pnp1-1* and 32.4-fold for the wild type), whereas the eight overlapping repressed genes had a similar average regulation level (0.22-fold for *pnp1-1* and 0.20-fold for the wild type).

### Constitutive P Starvation Responses in *pnp1-1*

The attenuated transcriptional response of *pnp1-1* under P starvation raised the possibility that the plants



**Figure 8.** Venn diagrams showing overlap of significantly induced or repressed genes between the indicated data sets derived from DNA microarray analysis. Significant changes were defined as 2-fold or greater, with  $FDR < 0.05$ . A to C represent the overlap between pairs of comparisons. Numbers in black and gray correspond to induced and repressed genes, respectively. WT, Wild type.

were experiencing some degree of P stress, even when grown under nominally +P conditions, and thus had constitutive P starvation responses. Therefore, we compared the -P transcriptional responses in the wild type with the *pnp* versus wild-type effects on gene expression under +P conditions (Fig. 8C). The behaviors of some of the main known P-responsive genes are detailed in Table IV. The global result that emerges is that among the 508 P-responsive genes in the wild type, 43% are also regulated in the same direction in *pnp1-1* relative to the wild type under +P conditions. This 43%, or 221 genes, includes 166 that are induced and 55 that are repressed. Among them are several major genes normally induced during P stress, such as those encoding P transporters belonging to the PHT1 family (*PHT1;1* and *PHT1;2*, detected with the same probe; *PHT1;4* and *PHT1;7*, detected with the same probe), the ribonuclease *RNS1*, and the transcription factor *WRKY75*, a positive modulator of P starvation responses and root development (Devaiah et al., 2007). Messenger RNAs encoding at least 30 other plastid-targeted proteins are also regulated in *pnp1-1* as if it were already P starved (Supplemental Table S6). This strengthens the conclusion that plastid P metabolism is altered in *pnp1-1*.

#### Verification of Transcriptome Results

We performed quantitative RT-PCR to validate a portion of the expression data described above, as shown in Figure 9, selecting both P starvation-responsive genes and genes whose expression was independently affected by the *pnp1-1* mutation. Figure 9A shows various protein-coding genes. We examined *PHR1*, which encodes a key P response transcription factor. As shown previously (Rubio et al., 2001), *PHR1* expression is not regulated transcriptionally by P starvation in the wild type, and we found the same for the *pnp* mutant. *RNS1*, which encodes a P starvation-induced secreted ribonuclease (Bariola et al., 1999), showed the expected induction in the wild type; however, it was also constitutively (under +P conditions) expressed at 3-fold the wild-type level in *pnp1-1*. The transcripts encoding the transporters *Pht1;4* and *Pht1;1* are induced in the wild type, as expected (Mudge et al., 2002), and in the mutant. We also verified four genes whose

expression was regulated P independently by the *pnp1-1* genotype. We confirmed that genes encoding a jacalin-lectin and a potentially chloroplast-targeted RNase H domain-containing protein were induced, in agreement with microarray results (Supplemental Table S7), and that two photosynthetic genes, *PORA* and *PSBP2*, were repressed.

Figure 9B shows expression analysis of two P starvation-induced riboregulators, *At4* and *IPS1*, which are not represented on ATH1 arrays. Both were strongly induced in both genotypes. Although it is not evident because of the scale of the graph, *IPS1* was slightly but significantly induced in *pnp1-1* under +P conditions, approximately 2-fold relative to the wild type. Also, we examined the expression of *PDR2*, since mutations in that gene phenocopy *pnp1-1* in terms of lateral root abortion. *PDR2* expression, however, did not differ in *pnp1-1*. Finally, we examined the expression of the *PNP* gene itself, and any dependence on *PHR1*, because we had previously found that in *Chlamydomonas* (Yehudai-Resheff et al., 2007) the *PNP* gene is repressed by P starvation in a *Psr1*-dependent manner and *PHR1* is the ortholog of *Psr1*. As shown in Figure 9C, *PNP* transcripts decreased approximately 2-fold under P starvation conditions, which is comparable to the approximately 3-fold decrease observed in *Chlamydomonas*. This decrease appeared to be *PHR1* independent, however. The character of this mutant was verified by measuring *IPS1* transcripts, which, as expected (Nilsson et al., 2007), failed to be induced. In summary, our quantitative RT-PCR data supported the conclusions from the microarray approach and yielded additional information regarding the expression of riboregulators and the regulation of the *PNP* gene.

#### DISCUSSION

Our previous report illuminated a role for the cpPNPase in P starvation acclimation in *Chlamydomonas* (Yehudai-Resheff et al., 2007) and stimulated the studies reported here. While we conclude that cpPNPase also is important for a normal P starvation response in Arabidopsis, we used a different set of analytical tools to address issues specific to the multicellular context. In one perspective, the main commonality in the two experimental systems is that cell death occurs, which in the unicellular *Chlamydomonas*, of course, is lethal, whereas in Arabidopsis it is restricted to lateral root initiates. Both results, however, point to a key role for PNPase apart from RNA metabolism.

#### Altered cpRNA Maturation in *pnp* Mutants Is Associated with a Pale-Green Leaf Phenotype

Disruption of the cpPNPase gene in Arabidopsis was previously noted to be associated with 3' extensions of both mRNAs and 23S rRNA (Walter et al., 2002), which we confirmed and also observed for the

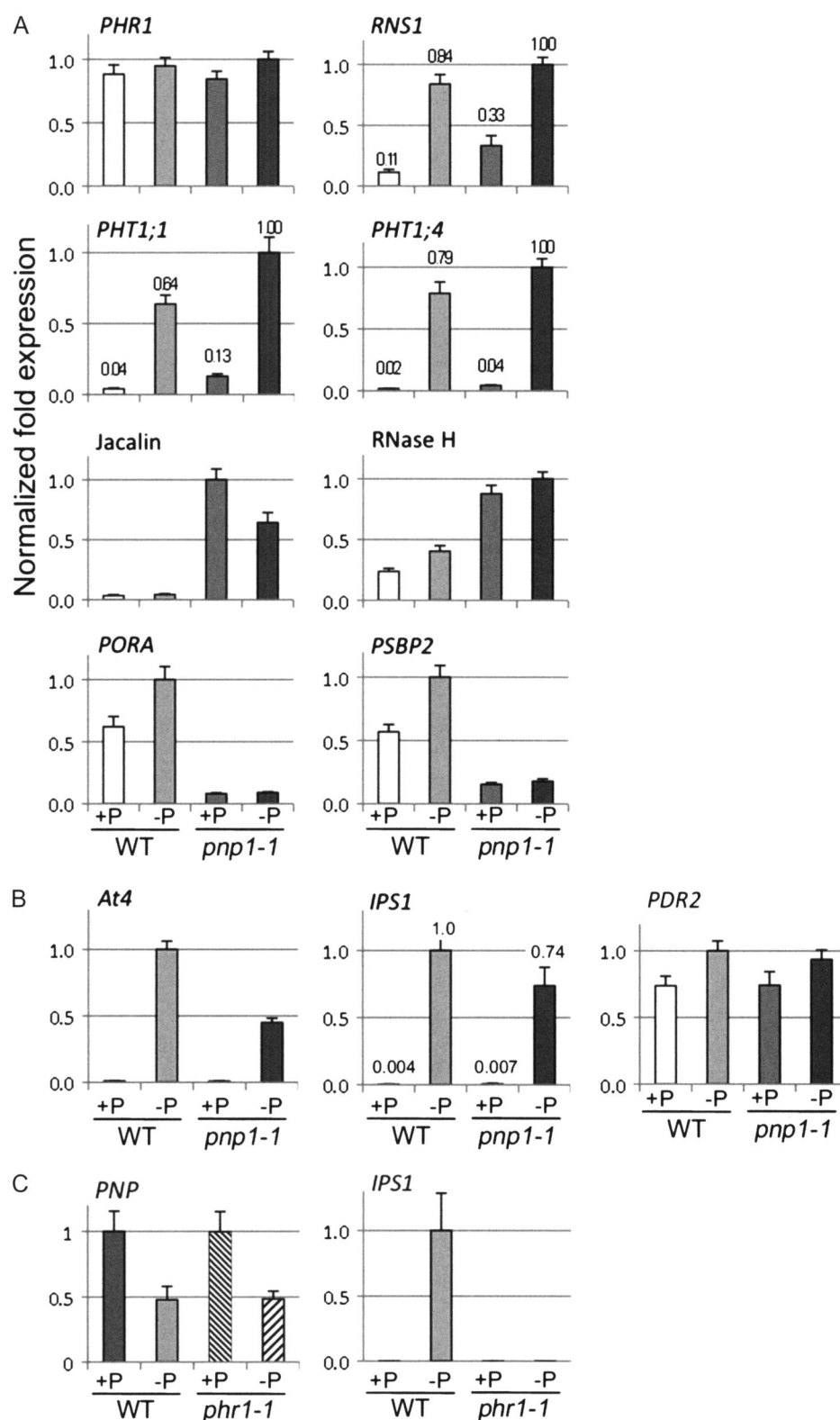
**Table III.** Selected genes regulated in *pnp1-1* relative to the wild type independent of P status

Probe Set Identifier	Arabidopsis Genome Initiative Code	Description	<i>pnp1-1</i> versus the Wild Type	
			Fold Change on +P	Fold Change on -P
Photosynthesis- and chloroplast-targeted proteins				
244977_at	AtCg00730	PetD	102.82	151.68
245002_at	AtCg00270	PsbD	28.93	34.44
245015_at	AtCg00490	Rbcl	16.12	18.88
244970_at	AtCg00660	Rpl20	15.03	26.41
245005_at	AtCg00330	Rps14	10.57	43.80
244991_s_at	AtCg00890	Ndh.B1	9.54	18.21
245003_at	AtCg00280	PsbC	6.35	9.79
244966_at	AtCg00600	PetG	5.57	7.03
245004_at	AtCg00300	Ycf9 (PsbZ)	3.24	4.99
245000_at	AtCg00210	Ycf6 (PetN)	2.79	3.08
244969_at	AtCg00650	Rps18	2.14	2.76
244988_s_at	AtCg00840	Rpl23.1	0.50	0.49
244964_at	AtCg00580	PsbE	0.40	0.47
245024_at	AtCg00120	AtpA	0.17	0.35
264868_at	At1g24090	RNase H domain-containing protein	7.19	5.56
256459_at	At1g36180	Acetyl-CoA carboxylase 2 (ACC2)	6.86	13.60
260648_at	At1g08050	Zinc finger (C3HC4-type RING finger) family protein	4.12	3.23
257667_at	At3g20440	BE1/EMB2729 (branching enzyme 1), $\alpha$ -amylase	3.58	4.98
245523_at	At4g15910	Drought-induced protein (Di21)	3.10	2.75
265479_at	At2g15760	Calmodulin-binding protein, similar to AR781	2.59	3.26
263325_at	At2g04240	Zinc finger (C3HC4-type RING finger) family protein	0.47	0.47
248040_at	at5g55970	Zinc finger (C3HC4-type RING finger) family protein	0.26	0.25
PYK10-associated proteins				
265051_at	At1g52100	JAL11, jacalin-lectin family protein	26.05	17.67
259384_at	At3g16450	JAL33, jacalin-lectin family protein, similar to ATMLP-300B (myrosinase-binding protein-like protein 300B)	13.28	4.97
266988_at	At2g39310	JAL22, jacalin-lectin family protein	8.73	3.46
259327_at	At3g16460	JAL34, jacalin-lectin family protein, similar to MBP1 (myrosinase-binding protein 1)	7.87	4.32
259381_s_at	At3g16390	JAL27, jacalin-lectin family protein, similar to ATMLP-470	5.25	2.71
262001_at	At1g33790	JAL4, jacalin-lectin family protein	3.28	4.63
259382_s_at	At3g16430	JAL30, PBP1 (PYK10-binding protein 1) JAL31	5.60	3.10
263153_s_at	At1g54000	GLL22, myrosinase-associated protein GLL23, myrosinase-associated protein	18.11	6.18
263156_at	At1g54010			
263156_at	At1g54030	GLL25, GDSL-motif lipase similar to ESM1 (epithiospecifier modifier 1)	3.57	2.19
259009_at	At3g09260	PYK10 (phosphate starvation response 3.1)	10.95	4.27
257798_at	At3g15950	TSA1-like (DNA topoisomerase), similar to TSK-associating protein 1 (TSA1)	12.46	5.78
249817_at	At5g23820	MD-2-related lipid recognition domain-containing protein	3.65	3.68
246855_at	At5g26280	Meprin and TRAF homology domain-containing protein	12.51	7.61
RNA- and DNA-related proteins				
264460_at	At1g10170	NF-X1-type zinc finger family protein	4.56	3.03
267140_at	At2g38250	DNA-binding protein-related, similar to transcription factor	3.17	2.55
260266_at	At1g68520	Zinc finger (B-box-type) family protein	0.40	0.46
262291_at	At1g70790	C2 domain-containing protein	0.38	0.35
248764_at	At5g47640	CCAAT box-binding transcription factor subunit B (NF-YB) (HAP3)	0.26	0.32
252504_at	At3g46590	TRFL1 (TRF-LIKE 1) telomere repeat-binding protein	0.26	0.40
263909_at	At2g36490	HhH-GPD base excision DNA repair family protein (ROS1)	0.16	0.24
264041_at	At2g03710	SEP4 (Sepallata 4); identical to agamous-like MADS-box protein AGL3 (AGL3)	0.12	0.10
Transport				
263402_at	At2g04050	MATE efflux family protein, similar to ATDTX1, antiporter	13.05	7.08
263401_at	At2g04070	MATE efflux family protein, similar to ATDTX1, antiporter	6.49	9.09
253732_at	At4g29140	MATE efflux protein-related	2.58	2.49
249188_at	At5g42830	Transferase family protein, similar to <i>N</i> -hydroxycinnamoyl/benzoyltransferase 6	10.47	5.68
255941_at	At1g20350	ATTIM17-1 (translocase mitochondrial inner membrane subunit 17-1)	10.44	5.51
253181_at	At4g35180	LHT7 (Lys/His transporter 7)	6.44	8.49
265479_at	At2g15760	Calmodulin-binding protein, similar to AR781	2.59	3.26
267483_at	At2g02810	ATUTR1 (UDP-Gal transporter 1)	2.45	2.34
265161_at	At1g30900	Vacuolar sorting receptor 6 precursor (AtVSR6)	2.33	3.60
254662_at	At4g18270	ATTRANS11 (translocase 11)	0.30	0.21
257294_at	At3g15570	Phototropic-responsive NPH3 family protein	0.15	0.03

**Table IV.** Behavior of major genes normally involved in the phosphate deprivation response

Values in bold are those for which the fold change is significant (fold change  $\geq 2$  and FDR  $< 0.05$ ); for those in normal type, the fold change is  $< 2$  and/or the FDR  $> 0.05$ .

Probe Set Identifier	Arabidopsis Genome Initiative Code	Description	<i>pnp1-1</i> +P versus Wild Type +P		Wild Type -P versus Wild Type +P		<i>pnp1-1</i> -P versus <i>pnp1-1</i> +P		<i>pnp1-1</i> -P versus Wild Type -P	
			Fold Change	FDR	Fold Change	FDR	Fold Change	FDR	Fold Change	FDR
247629_at	At5g60410	DNA-binding family protein (SIZ1)	1.82	0.324	1.97	NA	0.87	1.000	0.80	NA
253784_at	At4g28610	Phosphate starvation response regulator (PHR1)	0.94	0.820	0.86	0.589	0.91	1.000	1.00	0.999
245976_at	At5g13080	Transcription factor (WRKY75)	<b>7.41</b>	<b>0.022</b>	<b>4.77</b>	<b>0.114</b>	1.24	1.000	1.92	0.461
266743_at	At2g02990	Ribonuclease 1 (RNS1)	<b>4.04</b>	<b>0.024</b>	<b>6.66</b>	<b>0.008</b>	2.60	0.316	1.58	0.634
264893_at	At1g23140	C2 domain-containing protein	<b>3.55</b>	<b>0.014</b>	<b>48.17</b>	<b>0.000</b>	<b>11.83</b>	<b>0.000</b>	0.87	0.921
249152_s_at	At5g43350	Inorganic phosphate transporter (PT1) (Pht1;1/Pht1;2)	<b>7.83</b>	<b>0.001</b>	<b>40.68</b>	<b>0.000</b>	<b>7.65</b>	<b>0.001</b>	1.47	0.507
266184_s_at	At2g38940	Phosphate transporter (PT2) (Pht1;4/Pht1;7)	<b>9.54</b>	<b>0.003</b>	<b>367.71</b>	<b>0.000</b>	<b>51.03</b>	<b>0.000</b>	1.32	0.806
267646_at	At2g32830	Inorganic phosphate transporter (PHT5)	2.37	0.085	<b>12.00</b>	<b>0.001</b>	<b>6.39</b>	<b>0.011</b>	1.26	0.968
249151_at	At5g43360	Inorganic phosphate transporter (PHT3)	1.03	NA	1.02	NA	1.00	1.000	1.01	0.996
257311_at	At3g26570	Phosphate transporter family protein PHT2;1 (chloroplast)	0.74	0.782	0.74	0.876	0.75	1.000	0.75	0.902
258293_at	At3g23430	Phosphate transporter (PHO1)	0.74	0.656	<b>15.88</b>	<b>0.012</b>	6.43	0.186	0.30	0.223
267456_at	At2g33770	Ubiquitin-conjugating enzyme family protein (PHO2/UBC24)	1.40	0.315	1.76	0.164	1.10	1.000	0.87	0.926
264204_at	At1g22710	Suc transporter/Suc-proton symporter (PHO3/SUC2)	0.52	0.027	0.89	0.676	1.20	1.000	0.70	0.394
258158_at	At3g17790	Acid phosphatase type 5 (ACP5)	1.78	0.176	<b>42.25</b>	<b>0.000</b>	<b>20.71</b>	<b>0.002</b>	0.87	0.954
263083_at	At2g27190	Iron(III)-zinc(II) purple acid phosphatase (PAP12)	1.30	0.074	<b>5.35</b>	<b>0.000</b>	<b>4.11</b>	<b>0.000</b>	1.00	1.000
252006_at	At3g52820	Purple acid phosphatase (PAP22)	0.98	0.968	<b>15.16</b>	<b>0.000</b>	<b>12.50</b>	<b>0.000</b>	0.81	0.732
263594_at	At2g01880	Purple acid phosphatase (PAP7)	1.03	NA	10.00	0.307	10.88	0.597	1.12	0.950
255587_at	At4g01480	Inorganic pyrophosphatase (ATPPA5)	<b>2.01</b>	<b>0.013</b>	<b>5.08</b>	<b>0.000</b>	<b>2.36</b>	<b>0.016</b>	0.93	0.927
246071_at	At5g20150	SPX (SYG1/Pho81/XPR1) domain-containing protein (AtSPX1)	1.83	0.055	<b>99.05</b>	<b>0.000</b>	<b>46.20</b>	<b>0.000</b>	0.85	0.766
267611_at	At2g26660	SPX (SYG1/Pho81/XPR1) domain-containing protein (AtSPX2)	0.78	0.195	<b>5.17</b>	<b>0.000</b>	<b>4.85</b>	<b>0.000</b>	0.73	0.259
266132_at	At2g45130	SPX (SYG1/Pho81/XPR1) domain-containing protein (AtSPX3)	1.12	0.819	<b>139.62</b>	<b>0.000</b>	<b>123.14</b>	<b>0.000</b>	0.99	0.998
263599_at	At2g01830	His kinase (CRE1/AHK4/WOL)	<b>0.40</b>	<b>0.004</b>	0.55	0.033	0.98	1.000	0.71	0.273
258887_at	At3g05630	Phospholipase D (PLDP2)	0.89	NA	<b>69.58</b>	<b>0.000</b>	<b>62.57</b>	<b>0.000</b>	0.80	0.613
258452_at	At3g22370	Alternative oxidase 1a, mitochondrial (AOX1A)	<b>4.44</b>	<b>0.003</b>	1.87	0.107	0.99	1.000	2.35	0.074
257642_at	At3g25710	Basic helix-loop-helix family protein (bHLH32)	0.47	0.134	0.70	0.561	1.03	1.000	0.69	0.662
264783_at	At1g08650	Phospho <i>enol</i> pyruvate carboxylase kinase (PPCK1)	1.04	NA	<b>63.73</b>	<b>0.000</b>	<b>36.33</b>	<b>0.000</b>	0.59	0.404
258570_at	At3g04530	Phospho <i>enol</i> pyruvate carboxylase kinase (PPCK2)	1.02	NA	<b>31.95</b>	<b>0.005</b>	<b>9.09</b>	<b>0.222</b>	0.29	0.170
249846_at	At5g23630	ATPase E1-E2 type family protein (PDR2)	0.85	0.51	1.05	0.89	1.04	1	0.85	0.63
Plastid carbon transport										
248144_at	At5g54800	Glucose-6-phosphate/phosphate translocator (GPT1)	1.51	0.047	1.95	0.008	1.42	0.288	1.10	0.826
264400_at	At1g61800	Glucose-6-phosphate/phosphate translocator (GPT2)	<b>3.49</b>	<b>0.016</b>	<b>10.56</b>	<b>0.002</b>	<b>3.92</b>	<b>0.079</b>	1.29	0.806
252414_at	At3g47420	Glycerol-3-phosphate transporter	1.34	0.796	<b>33.88</b>	<b>0.016</b>	<b>24.66</b>	<b>0.057</b>	0.98	0.998
246445_at	At5g17630	Glucose-6-phosphate/phosphate translocator	0.89	0.666	0.77	0.419	0.75	0.880	0.87	0.785
248886_at	At5g46110	Phosphate/triose-phosphate translocator (TPT)	0.88	0.388	0.98	0.934	0.92	1.000	0.83	0.326
259185_at	At3g01550	Triose phosphate/phosphate translocator (PPT2)	0.80	0.831	0.35	0.856	0.36	1.000	0.82	0.954



**Figure 9.** Transcript levels for selected P starvation and *pnp1-1*-regulated genes as determined by quantitative RT-PCR. The fold change in expression was normalized to the actin gene *ACT2* and scaled to the sample with the highest expression level for each tested gene, which was defined as 1.0. Error bars represent se. Data correspond to three biological replicates and at least two technical replicates, as detailed in Supplemental Table S7. For selected genes, relative transcript abundance is shown above the bars. A, Validation of microarray data for selected genes. Gene symbols are used except where there is no gene name but the product is known. Plants were germinated and grown for 2 weeks on +P medium, then transferred to +P or -P medium for 1 additional week. B, Expression levels of *At4* and *IPS1*, two P starvation markers not represented on the ATH1 microarrays, and *PDR2*. Plants were grown as for A. C, Plants were germinated and grown for 1 week on +P medium, then transferred to +P or -P medium for 10 d, to replicate previously published growth conditions for the *phr1* mutant (Rubio et al., 2001). WT, Wild type.

*atpB* mRNA (Fig. 1D). The pale-green phenotype is evident in emerging leaves (Fig. 1C) but was not noted by Walter et al. (2002) because a cosuppressed line rather than T-DNA null mutants was used. However, the phenotype was noted where a *pnp* mutant allele

was obtained in a genetic screen for fosmidomycin resistance (Sauret-Gueto et al., 2006). The pale-green phenotype of young tissues progressively disappears, and cotyledons are indistinguishable from those of the wild type. This contrasts with deficiency in the prod-

uct of the paralogous At5g14580 locus, which encodes mitochondrial PNPase. In this case, null mutants are embryo lethal and knockdown lines hyperaccumulate antisense and intergenic transcripts, suggesting a perhaps fatal disruption in mitochondrial gene expression (Holec et al., 2006).

#### cpPNPase Deficiency Affects Root Adaptation to –P Stress

We observed that *pnp* plants grown under –P conditions were unable to elaborate lateral roots (Figs. 2 and 3), a phenomenon that phenocopies *pdr2*, albeit on a longer time scale. The *pdr2* mutation corresponds to a point mutation in At5g23630, which encodes a P-type ATPase of group V (Ticconi, 2005). Mutations in the same gene have been associated with reduced male fertility (Jakobsen et al., 2005). We also compared the root phenotype of *pnp1-1* with that of *csp41b-1*, the latter of which is deficient in two related chloroplast endoribonucleases and which has pale mature leaves (Bollenbach et al., 2009). Because *csp41b-1* resembled the wild type in terms of lateral roots, we conclude that neither the partial chlorosis nor the chloroplast gene expression defect of *pnp1-1* is likely to be responsible for its lateral root phenotype under –P conditions.

We used two reporter genes to gain additional insight into the *pnp1-1* root phenotype (Fig. 3). Using the *DR5* promoter to drive GUS expression, we observed reduced staining in *pnp1-1*, relative to the wild type, whether grown under +P or –P conditions. Moreover, in most cases only one or two lateral root initiates under –P conditions stained for GUS. Since the *DR5* construct essentially measures responsiveness to auxin in that tissue (Ulmasov et al., 1997), we conclude that auxin levels and/or activity are abnormally low. It has been shown that P starvation enhances auxin sensitivity in Arabidopsis roots and helps lead to higher lateral root density and inhibition of primary root elongation (Lopez-Bucio et al., 2002). Thus, our observations are consistent with the conclusion that the failure to elaborate lateral roots in *pnp1-1* under low-P conditions is at least in part related to a defective hormonal cue.

A second reporter gene, *CYCB1:GUS*, revealed no abnormalities under +P conditions but suggested that cell division was only occurring in a single lateral root initiate within the clusters that formed in *pnp1-1* under –P conditions. By staining with Evans blue, we concluded that, as in *pdr2*, mutant lateral root initiates die, followed by initiation of secondary, tertiary, and quaternary lateral roots. This cell death, or at least the loss of membrane integrity, was correlated with organellar abnormalities revealed by electron microscopy (Fig. 4). While we have not investigated whether lateral roots in *pnp1-1* undergo programmed cell death, it is worth noting that in plants, mitochondrial abnormalities in particular, but also chloroplast dysfunction or communication, have been associated with programmed cell death (Yao et al., 2004).

#### P Content and Uptake

To test the hypothesis that PNPase might have a role in P homeostasis, we measured total and free Pi in leaves, where PNPase is predominantly expressed, as well as in roots, where a defective growth phenotype was observed. A slight increase of total P and free Pi was evident under +P conditions for *pnp1-1*, whereas no significant differences were noted under –P conditions in the wild type (Fig. 6). While most P-containing metabolites that we measured did not exhibit differences between genotypes, phosphate was slightly elevated in *pnp1-1* (Fig. 7). Also, while P uptake did not differ between genotypes, we saw slight induction relative to the wild type of the gene encoding the P transporters *Ph1;1* and *Ph1;4*, under +P conditions (Fig. 9). This is consistent with a degree of P stress in *pnp1-1* under +P conditions, as discussed below.

#### Transcript and Metabolite Profiling

Microarray data revealed an obvious reorientation of the *pnp* mutant transcriptome toward expression of –P responses when grown on a full nutrient medium, comprising 221 out of the 508 P starvation-regulated genes in the wild type. Taken together with the P transporter data discussed above, we hypothesize that *pnp* mutant plants adjust their metabolism as if they were already, to some extent, under phosphate stress.

A second set of genes is responding to the absence of PNPase independent of P availability. Overlap analyses (Fig. 8A) between *pnp*-regulated genes on +P and on –P compared with the wild type revealed 149 genes that fall into two major metabolic classes: regulation of chloroplast activities and oxidative stress responses. Fourteen of these are encoded by the chloroplast genome and are generally strongly up-regulated. As mentioned earlier, the accumulation of these transcripts likely results from perturbation of normal cpRNA degradation pathways. The genotype also affected 12 nucleus-encoded chloroplast proteins. On the other hand, of several genes suggested to be responsible for the communication between chloroplast and nucleus, or retrograde signaling, none was significantly regulated. Altered photosynthesis-related gene expression in *pnp1-1* is in agreement with its partial chlorosis and slow-growth phenotype.

We also examined gene expression related to the chloroplast MEP pathway, since the *pnp* mutant *rif10* was identified using fosmidomycin, a strong inhibitor of deoxyxylulose 5-phosphate reductoisomerase (DXR), which catalyzes the second step (Sauret-Gueto et al., 2006). In the case of *rif10*, the authors reported a similar transcript accumulation for the deoxyxylulose 5-phosphate synthase (DXS), the first enzyme of the MEP pathway, and for DXR but increased accumulation for both at the protein level. Our experiments confirmed the lack of significant regulation for two DXS genes (*DXS1* and *DXS3*) and the DXR gene but revealed 5.8-fold activation of the third DXS

gene (*DXS2*; At3g21500), which is expressed at a low level under normal conditions, based on EST data (Rodriguez-Concepcion and Boronat, 2002), but in our study was also activated in the wild type during P starvation. That regulation of *DXS2* in *pnp1-1*, rather than a posttranslational effect, could explain at least in part the DXS enzyme accumulation described, although it should be noted that the activities of *DXS2* and *DXS3* remain to be established; indeed, it has been proposed that they do not encode functional proteins and should be renamed *DXL* (for DXS-like; Phillips et al., 2008).

Overall, in *pnp* mutants including *rif10* and *pnp1-1*, defects in the MEP pathway lead to less accumulation of chlorophyll and carotenoids. This is not only consistent with the observed chlorosis but may make the plants subject to oxidative stress due to a decrease in photoprotective carotenoids. Finally, because this pathway is also responsible for the biosynthesis of certain hormones, their levels may also be affected in *pnp* mutants.

Metabolic profiling revealed limited changes between *pnp1-1* and the wild type in the presence of P. However, *pnp1-1* displayed a different response to P starvation than wild-type Arabidopsis (Morcuende et al., 2007; this work) or common bean (*Phaseolus vulgaris*; Hernandez et al., 2007). For example, changes in the amino acid profile under  $-P$  conditions strongly hint of a perturbation in plastid metabolism, including large changes in the Asp family amino acids. Interestingly, these metabolites are those that cost the most ATP in their production, and their rate of biosynthesis has previously been demonstrated to be highly dependent on the plastidial energy charge (Regierer et al., 2002; Carrari et al., 2005). Importantly, these changes are considerably more dramatic in the *pnp* mutant than in the wild type, implying a functional role for the PNP protein in this process. Although less easy to rationalize there are also large changes in cytosolic and mitochondrially synthesized metabolites such as Gly, Ser, Ala, and GABA.

#### Oxidative Stress Responses and Organellar Integrity in *pnp1-1*

At the transcriptional and metabolic levels, several lines of evidence suggest activation of oxidative stress responses in *pnp1-1*. Among the metabolites, DHA is the most strongly regulated on  $+P$ , with its content increased about 15-fold. Several enzymes involved in the ascorbate cycle, and its biosynthesis, are strongly regulated in *pnp1-1*. Induced genes include *VTC5*, an ascorbate biosynthetic enzyme (Dowdle et al., 2007), and two ascorbate oxidases that convert L-ascorbate to mono-DHA (Ishikawa et al., 2006). In contrast, the gene encoding the chloroplast thylakoid enzyme responsible for the same conversion but via hydrogen peroxide rather than oxygen scavenging, tAPX, is repressed. Reduction of mono-DHA, which is a radical, is catalyzed by mono-DHA reductase, which is also

strongly induced in *pnp1-1*. Finally, the initiation and the activity of the ascorbate-glutathione cycle are strongly induced, probably leading to accumulation of DHA. However, the repression of the gene encoding tAPX might lessen the ability of the *pnp* mutant to detoxify the chloroplast (for review, see Asada, 2006). It should also be noted that a recent study associated oxidative stress resulting from a combination of drought and heat with enhanced malic enzyme activity and a decrease in malate, leading the authors to speculate that malate metabolism plays an important role in the response of Arabidopsis to this stress combination (Koussevitzky et al., 2008). Our findings of 50% lower malate in *pnp1-1* accompanying the accumulation of DHA hint of a similar response in this mutant.

The *pnp1-1*  $+P$  transcriptome revealed induction of additional genes involved in redox homeostasis, including peroxidases, thioredoxins, cytochromes P450, and glutathione S-transferases. We also note the induction of one cytosolic and one chloroplast copper/zinc superoxide dismutase: *CSD1* and *CSD2*, respectively (Kliebenstein et al., 1998). These are involved in superoxide radical detoxification and the positive and negative regulation of genes encoding glutaredoxins. Some of this regulation was also found in the wild type under P starvation.

A general consequence of reactive oxygen species accumulation can be the alteration of membrane lipids, which could lead in turn to organelle disorganization, as we observed in *pnp1-1* roots (Fig. 4). This loss of organelle integrity might be linked to the induction of a group of genes related to the PYK10 complex, as discussed in "Results." The genes encoding PYK10 complex components were found to be repressed in the *nai1* mutant (Nagano et al., 2008), which lacks a basic helix-loop-helix transcription factor (Matsushima et al., 2004). Another target of *NAI1* is the lipoyxygenase *LOX3*, which is induced more than 12-fold in *pnp1-1* on  $+P$ ; the *NAI1* gene itself is induced 1.8- and 1.5-fold in *pnp1-1* versus the wild type on  $+P$  and  $-P$ , respectively. One study of endoplasmic reticulum bodies suggested that the PYK10 complex might form only when subcellular structures are altered or disrupted, as the complex's partners would normally be localized in different compartments (Nagano et al., 2008). The induction of PYK10 complex genes in *pnp1-1* could be thus related to the loss of organelle integrity triggered by oxidative stress.

#### A Systemic Signal May Be Altered in *pnp1-1*

The split-root experiment (Fig. 5) raised the possibility that a systemic signal was affected in *pnp1-1* plants. Such a signal has been hypothesized to arise in the root cap during P starvation (Svistoonoff et al., 2007) and may be integrated in the aerial part of the plant to stimulate adaptation in root architecture. Our results suggest that in the case of *pnp1-1*, either a major metabolite or hormonal balance may be relevant to this issue.



One metabolite candidate would be sugars, as a tight relationship between sugar metabolism and P deprivation responses is well established and was evident in a transcriptomic comparison between P starvation responses and Suc-regulated metabolism (Muller et al., 2005). It was found that some P-responsive genes are more strongly induced in the presence of Suc and that these enhanced responses were correlated with increased lateral root density (Karthikeyan et al., 2007). It has additionally been proposed that phloem Suc transport may integrate root responses to phosphate deprivation (for review, see Hammond and White, 2008).

In the experiments reported here, *pnp1-1* differed from the wild type in particular with respect to early fairly dramatic decreases in Fru-6-P at 3 h and to a delayed increase in raffinose as compared with the wild type. The latter was found to accumulate in the mutant under control conditions. Additional differences in the sugar response to P starvation between the two genotypes are to be found in the opposite behavior of isomaltose and trehalose and the sugar derivative myoinositol, particularly following 1 week of starvation. The differences are reduced in the longest starvation period, suggesting an adjustment of sugar metabolism to P deprivation within the first hours.

Our microarray analysis also revealed gene expression changes in *pnp1-1* closely related to Suc metabolism. For example, under +P conditions, several carbohydrate transporters are induced: the plastid Glc-6-P translocator (GPT2), a mannitol transporter, a sugar transporter (STP4), two UDP-Gal transporters, and the Suc transporter SUC1. Also in *pnp1-1*, glycolysis appears to be globally activated. While it is challenging to link particular metabolite levels to any of these changes in gene expression, it is notable that both metabolite steady-state levels and relevant genes are fluctuating in tandem.

Two other sugars, isomaltose and trehalose, increased approximately 5.5- and 4-fold in *pnp1-1* versus the wild type, respectively, when grown on +P (Fig. 7). Their accumulation patterns also differ qualitatively upon P starvation (Supplemental Fig. S1). The role of isomaltose is unclear, as it is apparently not a major form of carbon exported from the chloroplast, at least in wild-type plants (Weise et al., 2004). Nor does isomaltose appear to be accumulating at the expense of maltose, which is the major exported form. However, we cannot exclude a deficiency of carbon export, as we have examined only total metabolites rather than their partitioning into subcellular compartments.

Trehalose is a disaccharide whose phosphorylated form, trehalose-6-phosphate (Tre-6-P), appears to be an important signaling molecule related to sugar metabolism (Paul et al., 2008). Trehalose is produced in a tandem reaction commencing with the synthesis of Tre-6-P from Glc-6-P and UDP-Glc catalyzed by Tre-6-P synthase (TPS); subsequently, Tre-6-P phosphatase (TPP) catalyzes the dephosphorylation of Tre-6-P to trehalose. The accumulation of trehalose in the mutant

under +P conditions might suggest an up-regulation of Tre-6-P dephosphorylation, a phenomenon that occurs in the wild type during the short periods of P starvation and might relate to P repartitioning within the cell. On the other hand, trehalose levels could increase in response to stress conditions (redox and/or P starvation). Indeed, although the role of trehalose as an osmoprotectant remains under debate, it is known to accumulate under a variety of environmental stresses, including cold, osmotic imbalance, and salt. Most of its biosynthetic genes respond to plant exposure to a wide range of abiotic and perhaps also biotic stresses (Iordachescu and Imai, 2008).

Consistent with our metabolite data and published gene expression results, transcriptome analyses revealed that five genes related to trehalose biosynthesis are differentially regulated on +P in *pnp1-1*. Four of them encode TPP (*TPPG* and *TPPD* are induced and *TPPA* and *TPPH* are repressed), whereas *TPS1* is repressed. *TPPA*, *TPPD*, and *TPPG* are predicted to be chloroplast targeted. Because both TPP and TPS are encoded by multigene families whose expression profiles vary widely (Paul et al., 2008), how this suite of expression changes results in increased trehalose content in *pnp1-1* is difficult to pinpoint. Furthermore, since Tre-6-P is an important metabolite whose concentration we have not measured directly, its relevance to the *pnp* root phenotype remains an open question.

#### Hormone Balance in *pnp1-1*

Many publications concerning P starvation responses highlight the roles of hormones in signal transduction, particularly auxins and cytokinins (Rubio et al., 2009; for review, see Yuan and Liu, 2008). It is not surprising, then, that many genes involved in hormone biosynthesis or degradation are regulated in the *pnp* mutant on +P (Supplemental Table S4). Furthermore, expression of DR5:GUS in *pnp1-1* (Fig. 3A) suggested a decreased auxin activity in *pnp1-1* root tips on +P as well as on -P. However, because we did not carry out detailed analyses of hormone contents, nor create genetic combinations with hormone pathway mutants, we feel it is premature to draw any direct connections between normal PNPase activity and hormone signaling.

In conclusion, the *Arabidopsis* and *Chlamydomonas* *pnp* mutants have few commonalities in their response to P starvation when examined in molecular detail. However, the importance of PNPase, and more generally the chloroplast, in conferring the ability to correctly respond to P starvation is conserved. Given the considerable differences in the survival strategies of a motile, unicellular organism and a sessile, multicellular one, differences in gene regulation and the consequences of PNPase inactivation are perhaps not surprising. Examining additional evolutionarily diverse photosynthetic species in this regard should similarly be interesting. How PNPase activity influ-

ences the ability of organisms to respond to P stress remains to be understood in detail, in particular what type of signal it generates and how that signal is integrated into the global response pathway.

## MATERIALS AND METHODS

### Plant Material

All of the *Arabidopsis* (*Arabidopsis thaliana*) plants used in this study are derived from the Columbia-0 ecotype, which was used as the wild type. The two mutant lines, *pnp1-1* (SALK\_013306) and *pnp1-2* (SALK\_070705), containing a T-DNA insertion in the *PNP* gene (At3g03710), were obtained from the SIGnAL collection (Alonso et al., 2003). The precise locations of the T-DNA left borders were confirmed by DNA sequencing. PCR with the left border primer (LBb1, 5'-GCGTGGACCGCTTGCTGCAACT-3') and gene-specific primers (*pnp1-1*, 5'-GCAAAGCTCGCTGTTAGATG-3' and 5'-CATAGCCATGTCA-ACITTTGCC-3'; *pnp1-2*, 5'-TACGTAGCGCAATTGTTGAGG-3' and 5'-CCAAACAACAGATGCCATACTG-3') was used to identify T-DNA insertion alleles of *pnp1-1* and *pnp1-2*, respectively. The *pnp1-1* line was crossed with the *Arabidopsis* *CYCB1:GUS* (Ulmasov et al., 1997) and *DR5:GUS* (Colón-Carmona et al., 1999) lines to generate the reporter lines used in Figure 3.

### Growth Conditions and Phosphate Starvation Treatment

Seeds were surface sterilized and stratified at 4°C for 3 to 4 d. Unless noted in the figure legends, plants were grown as follows. Seeds were germinated in a full nutrient Murashige and Skoog (MS) liquid medium in a controlled-environment chamber on a shaker at 25°C under fluorescent lights (100  $\mu\text{mol m}^{-2} \text{s}^{-1}$ ) with a long-day photoperiod (16 h of light). After 2 weeks, plantlets were transferred into Magenta boxes or onto petri plates with MS medium (1.25 mM  $\text{KH}_2\text{PO}_4$ ) containing 2% (w/v) Suc and 0.75% (w/v) phytagar (Murashige and Skoog, 1962) in the indicated periods. This medium is referred to as +P medium. For -P medium,  $\text{KH}_2\text{PO}_4$  was omitted from the nutrient solution and the plantlets were rinsed with the same liquid solution prior to transfer. The phytagar (commercial grade; Gibco-BRL) contributed about 25  $\mu\text{M}$  total P to the final medium.

For the microarray and the quantitative RT-PCR experiments, plants were germinated and grown for 2 weeks on a full nutrient MS medium (+P) containing 0.6% phytagar (w/v) and 0.5% (w/v) Suc (to limit the Suc effect on gene expression) at 22°C in a growth chamber under a 16-h photoperiod (with a fluorescent light intensity of 200  $\mu\text{mol m}^{-2} \text{s}^{-1}$ ). Then, they were transferred to fresh +P or -P MS medium. For the -P medium,  $\text{KH}_2\text{PO}_4$  was omitted but the potassium was compensated by  $\text{K}_2\text{SO}_4$ . Plantlets were rinsed with distilled water before transfer. On soil, plants were grown on Metro Mix 360, in a growth chamber, as described above.

### RNA Analysis

Total RNA was isolated using Tri Reagent according to the manufacturer's instructions (Molecular Research Center), separated by electrophoresis, and transferred onto a GeneScreen membrane (Perkin-Elmer) as described previously (Bollenbach et al., 2005). Gene-specific probes were labeled by random priming using 100 ng of DNA, 20  $\mu\text{Ci}$  of [ $\alpha$ - $^{32}\text{P}$ ]dCTP, and the Klenow fragment (Promega).

### UV Cross-Linking

UV cross-linking of proteins to radiolabeled RNA was performed as described previously (Lisitsky et al., 1997). The proteins (10 fmol) were mixed with [ $\alpha$ - $^{32}\text{P}$ ]UTP-RNA (10 fmol) in a buffer containing 10 mM HEPES-NaOH, pH 7.9, 30 mM KCl, 6 mM  $\text{MgCl}_2$ , 0.05 mM EDTA, 2 mM dithiothreitol, and 8% (v/v) glycerol and cross-linked immediately with 1.8 J of UV irradiation (Stratalinker 1800; Stratagene). The RNA was then digested with 10  $\mu\text{g}$  of RNase A and 30 units of RNase T1 at 37°C for 1 h. The proteins were fractionated by SDS-PAGE and analyzed by autoradiography.

### Root Analyses

Images of roots were recorded with a stereomicroscope (Olympus SZX12) high-performance CCD camera and imported into Photoshop Image software. Histochemical analysis of the GUS reporter enzyme activity was adapted from Jefferson (1987). Samples were stained using 2 mM 5-bromo-4-chloro-3-indolyl- $\beta$ -glucuronic acid (Sigma) dissolved in a 100 mM sodium phosphate buffer (pH 7.2) containing 0.2% Triton X-100, 2 mM  $\text{K}_4\text{Fe}(\text{CN})_6 \cdot \text{H}_2\text{O}$ , and 2 mM  $\text{K}_3\text{Fe}(\text{CN})_6 \cdot \text{H}_2\text{O}$ . For cell viability analysis, roots were stained with 10% (w/v) Evans blue (Fisher). After washing with distilled water and mounting in 50% glycerol, root tips were viewed using the Olympus SZX12 microscope.

### Transmission Electron Microscopy

Roots were embedded in an epoxy resin (Spurr, 1969). The root tips, from root cap to root hair zone (approximately 2 mm in length), were cut and fixed overnight in 5% (v/v) glutaraldehyde in 0.1 M sodium cacodylate buffer (pH 7.4) at 4°C, postfixed in 1% (w/v) aqueous osmium tetroxide for 3 h at room temperature, and rinsed three times with distilled water. After dehydration through a graded ethanol series, the samples were rinsed in propylene oxide, infiltrated, and embedded in Spurr's resin. Ultrathin sections were cut to a 50- to 70-nm thickness with a diamond knife on a Leica UCT ultramicrotome. The sections were stained with uranyl acetate (2.5%, w/v) and lead citrate. After staining, the sections in the middle longitudinal direction were viewed and photographed using a Philips FEI-Technai 12 microscope.

### Determination of Metabolite Levels

Free Pi and total P were assessed as described previously (Versaw and Harrison, 2002). For metabolic profiling, leaf samples were immediately frozen in liquid nitrogen and stored at -80°C. Extraction and quantification of metabolites were carried out using an established gas chromatography-mass spectrometry-based protocol (Roessner et al., 2001), with the exception that the metabolites studied also included subsequent additions to the mass spectral libraries (Schauer et al., 2005).

Principal component analysis was performed with the online tool MetaGeneAlyse ([www.metagenalyse.mpimp-golm.mpg.de](http://www.metagenalyse.mpimp-golm.mpg.de); Scholz et al., 2004) and TMEV software (Saeed et al., 2003). The data were normalized to the mean of the entire sample set for each metabolite and  $\log_{10}$  transformed before the analysis. This transformation reduces the influence of rare high-measurement values but does not change the discrimination in the data set. Statistical analysis of the data was performed by *t* test and two-way ANOVA.

### Microarray Hybridization and Data Analysis

Total RNA was isolated from seedlings, from which roots had been removed, using the RNeasy Plant Minikit (Qiagen), including DNase treatment according to the manufacturer's instructions. RNA quality check, linear amplification, labeling, hybridization, washing, and scanning were performed by the Cornell Microarray Core Facility (<http://cores.lifesciences.cornell.edu/brcinfo/?f=10>). Affymetrix ATH1 genome array GeneChips were used. Three biological replicates were used for each experimental condition.

Raw array data were normalized at the probe level using gcRMA (Wu et al., 2004). The detection calls (present, marginal, or absent) for each probe set were obtained using the mas5calls function in the Affy package (Gautier et al., 2004). Only genes with at least one present call across all of the compared samples were used to identify differentially expressed genes. Significance of gene expression was determined using the LIMMA test (Smyth, 2004), and raw *P* values of multiple tests were corrected using FDR (Benjamini and Hochberg, 1995). Genes with FDR < 0.05 and fold change  $\geq 2$  were identified as differentially expressed. The complete data set was deposited in the National Center for Biotechnology Information Gene Expression Omnibus under accession number GSE 18071.

### Quantitative RT-PCR

One microgram of DNase-treated RNA was reverse transcribed in a 20- $\mu\text{L}$  reaction using SuperScript III (Invitrogen) according to the instructions, including the RNase H treatment. One microliter of this cDNA was amplified using the Fast Sybr Green master mix (Applied Biosystems) and 0.66  $\mu\text{M}$  of each primer in a 15- $\mu\text{L}$  reaction. PCR amplification was performed using the

Bio-Rad CFX96 real-time PCR detection system with the following conditions: initial denaturation at 95°C for 3 min, followed by 40 cycles of 95°C for 10 s and 60°C for 30 s. The amplification specificity was checked using melting curves. The relative quantification of the samples was determined using the Bio-Rad CFX Manager software, integrating primer efficiencies calculated from a standard curve. For each gene, the sample showing the highest intensity level was used as reference with a value of 1. The final data result from averages of three biological replicates and at least two technical repetitions. Primers are listed in Supplemental Table S8.

## Supplemental Data

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

**Supplemental Figure S1.** Relative metabolite contents of *pnp1-1* and wild type grown under -P conditions for different time periods.

**Supplemental Figure S2.** Principal component analysis of metabolite profiles.

**Supplemental Figure S3.** Pi uptake in *pnp1-1* and wild type.

**Supplemental Table S1.** Raw data for metabolite contents of *pnp1-1* and wild type grown under +P or -P conditions.

**Supplemental Table S2.** Statistical analysis of metabolite data.

**Supplemental Table S3.** Microarray data for significantly regulated genes after 3 h of P starvation.

**Supplemental Table S4.** Microarray data for significantly regulated genes after 1 week of P starvation.

**Supplemental Table S5.** Statistical significance of the functional categorization of regulated genes using MapMan-defined BINs.

**Supplemental Table S6.** Genes encoding chloroplast-targeted proteins that are similarly regulated in *pnp1-1* +P versus wild type +P and in wild type -P versus wild type +P.

**Supplemental Table S7.** Normalized expression data for quantitative RT-PCR.

**Supplemental Table S8.** List of primers used for quantitative RT-PCR.

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