Genomic Identification and Expression Analysis of the Phosphate Transporter Gene Family in Poplar

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Inorganic phosphate is one of key macronutrients essential for plant growth. The acquisition and distribution of phosphate are mediated by phosphate transporters functioning in various physiological and biochemical processes. In the present study, we comprehensively evaluated the phosphate transporter (PHT) gene family in the latest release of the Populus trichocarpa genome (version 3.0; Phytozome 11.0) and a total of 42 PHT genes were identified which formed five clusters: PHT1, PHT2, PHT3, PHT4, and PHO. Among the 42 PHT genes, 41 were localized to 15 Populus chromosomes. Analysis of these genes led to identification of 5–14 transmembrane segments, most of which were conserved within the same cluster. We identified 234 putative cis elements in the 2-kb upstream regions of the 42 PHT genes, many of which are related to development, stress, or hormone. Tissue-specific expression analysis of the 42 PtPHT genes revealed that 25 were highly expressed in the roots of P. tremula, suggesting that most of them might be involved in Pi uptake. Some PtPHT genes were highly expressed in more than six of the twelve investigated tissues of P. tremula, while the expression of a few of them was very low in all investigated tissues. In addition, the expression of the PtPHT genes was verified by quantitative real-time PCR in four tissues of P. simonii. Transcripts of 7 PtPHT genes were detected in all four tested tissues of P. simonii. Most PtPHT genes were expressed in the roots of P. simonii at high levels. Further, PtPHT1.2 and PtPHO9 expression was increased under drought conditions, irrespective of the phosphate levels. In particular, PtPHT1.2 expression was significantly induced by approximately 90-fold. However, the transcriptional changes of some PtPHT genes under drought stress were highly dependent on the phosphate levels. These results will aid in elucidation of the functions of PtPHT in the growth, development, and stress response of the poplar plant.

Keywords: phosphate transporter, poplar, genome-wide analysis, expression profile, phosphate levels, drought

INTRODUCTION

Phosphorus is a major macronutrient required for plant growth and development. It constitutes up to 0.2% of the dry weight of plant cells and consequently is required in significant quantities (Schachtman et al., 1998). However, it is often a limiting factor for plant growth because the soil inorganic phosphate (Pi) concentration is usually lower than 10μM (Shen et al., 2011).
Plants have adopted several strategies to increase the acquisition of poorly available P source (Bucher, 2007). For example, plants could produce root exudates such as organic acids, sugars, and phenolic acids in the rhizosphere to improve the acquisition of poorly available P from soil (Cesco et al., 2010; Valentunuzzi et al., 2015). Additionally, plants have evolved specific transport systems for the uptake and partitioning of Pi (Poirier and Bucher, 2002). Both low- and high-affinity Pi-transporter systems have been identified in plants (Clark et al., 2000; Misson et al., 2004). The genes encoding for low-affinity Pi transport are constitutively expressed and function at relatively high Pi in the millimolar range, whereas high-affinity Pi transporters are transcriptionally induced at low Pi levels in the micromolar range (Raghothama, 2000).

Generally, the phosphate transporters are classified into PHT1, PHT2, PHT3, and PHT4 families (Rausch and Bucher, 2002; Liu et al., 2011). PHT1s belonging to the major facilitator superfamily are homologs of the yeast PHO84 Pi transporter (Pao et al., 1998). Since the initial identification of PHT1 genes in Arabidopsis (Muchhal et al., 1996), homologs of PHT1 transporters have been characterized in many species, such as tomato (Chen et al., 2014), rice (Liu et al., 2011), and poplar (Loth-Pereda et al., 2011). The PHT1 family, as H⁺/Pi symporters, has been reported to include proteins involved in the Pi uptake from the soil and distribution within plants (Nussaume et al., 2011). Regarding the PHT2 family of Pi transporters, one member (PHT2:1) has been functionally studied in Arabidopsis (Daram et al., 1999). PHT2 family proteins function as H⁺/Pi cotransporters in the plastids of plants (Versaw and Harrison, 2002). In addition, the PHT3 family was reported after the cloning of the first putative mitochondrial Pi transporter gene in plants (Takahatake et al., 1999). The PHT3 family, which are supposed to act as a Pi/H⁺ symporter Pi/OH⁻ antiport, catalyze Pi/Pi exchange between the matrix and cytosol (Stappen and Krämer, 1994). This family included at least three members in Arabidopsis to date, and all of them show high conservation within the mitochondrial transporter family (Rausch and Bucher, 2002). The PHT4 family gene, which includes six members in Arabidopsis, has been reported to be involved in the Pi transport in plastids and the Golgi apparatus (Guo et al., 2008). Recently, the PHO1 gene in Arabidopsis was identified by a map-based positional cloning strategy (Hamburger et al., 2002). The members of PHO1 family were mainly expressed in root stellar cells and in the lower part of the hypocotyl, which is consistent with the role of PHO1 in the loading of Pi to the xylem (Poirier and Bucher, 2002). The expression patterns of 11 PHO1 homologs in the Arabidopsis genome suggest that PHO1 proteins likely play a role in transferring Pi to the vascular cylinders in various tissues, as well as in acquiring Pi into cells, such as pollen, root epidermal cells, and cortical cells (Wang et al., 2004). Most studies of PHT have focused on a single cluster of the PHT family, such as the PHT1 family (Loth-Pereda et al., 2011; Nussaume et al., 2011); however, to our knowledge, there is no report of the whole PHT family, including PHT1, PHT2, PHT3, PHT4, and PHO, in any plant species. In addition, there are few reports of coordinated expression of the entire set of PHTs on a global scale in woody tree species.

Poplar is a model woody tree species used in functional genomic studies due to its small genome, easy transformation, fast growth, and vegetative propagation (Brunner et al., 2004a). Populus simonii is an important poplar species in the northwestern area of China with a wide distribution, and it is capable of growing under low- or high-Pi conditions (Weisgerber and Han, 2001). In this study, we conducted comprehensive genome-wide analysis to identify phosphate transporter gene family in poplar, including PHT1, PHT2, PHT3, PHT4, and PHO, in the P. trichocarpa genome (version 3.0; Phytozome 11.0); in addition, we analyzed the phylogeny, gene structures, genomic locations, and cis elements of these genes. We also investigated the expression patterns of all of the PHT genes in various tissues, as well as their expression profiles at different Pi levels (Pi starvation, low and high Pi levels) in P. simonii. Moreover, we examined the transcriptional profiles of PHT genes in P. simonii roots under drought conditions. The results of this study may provide a basic foundation for future functional genomic studies of the PHT family in poplar.

**MATERIALS AND METHODS**

**Plant Seedlings and Growth Conditions**

The cuttings of P. simonii (ca. 15 cm in length, 2 cm in diameter) were rooted in pots (10 L) filled with fine sand. The plants were cultivated in a glasshouse (natural light, 75% relative humidity) and irrigated with 50 ml nutrient solution (Hoagland and Arnon, 1950) (10 µM EDTA-FeNa, 5 µM MnSO₄·H₂O, 1 µM ZnSO₄·7H₂O, 1 µM CuSO₄·5H₂O, 30 µM H₂BO₃, 0.5 µM H₂MoO₄, 1000 µM MgSO₄·7H₂O, 1000 µM CaCl₂, 1000 µM Na₂SO₄, and 1000 µM NH₄NO₃, pH 6.5) containing Pi at one of three different levels (0 µM, 1000 µM, and 3000 µM KH₂PO₄) separately every other day. Each Pi treatment was applied to 12 plants for a period of 8 weeks. In addition, a phosphate-free solution was prepared by replacing KH₂PO₄ with K₂SO₄. Six plants for each Pi treatment were subjected to drought stress for 2 weeks before harvesting, and the other six plants for each treatment were not subjected to drought stress and served as controls. After the treatments, whole plantlets were harvested and separated into roots, stems, new leaves, and old leaves. Then, samples from the same tissues were pooled for each treatment, frozen in liquid nitrogen for further analysis.

**Identification and Phylogenetic Analysis of PHT Gene Family Members in Populus**

To identify phosphate transporters in poplar, the reported protein sequences from both Arabidopsis and Oryza sativa were used as query sequence with E-value cutoff set as 1e⁻¹⁰ to perform a local BLASTP against P. trichocarpa JGI gene catalog (phytozome v11.0, https://phytozome.jgi.doe.gov/pz/portal.html), respectively. The choice of candidate phosphate transporters was based on the E-value, the sequence homology value (>60%) and the value of score (>450). All corresponding DNA and protein sequences of poplar phosphate transporters were obtained from Phytozome 11.0 (http://phytozome.jgi.doe.gov/pz/portal.html) and also verified with the Pfam database (http://pfam.xfam.org/). Various splicing variants of one gene
and incomplete genes with short length were discarded. In addition, redundant sequences were removed, and the remaining sequences were used in further analyses.

For each putative protein, the grand average of hydropathicity (GRAVY) was calculated using ProtParam (http://web.expasy.org/protparam/). Further, TMPred (http://www.ch.embnet.org/software/TMPRED_form.html) was used to predict the transmembrane domains (TMDs) of each PHT protein. In addition, the subcellular location of each PHT was predicted using WolF PSORT (http://www.genscript.com/wolf-psort.html). Multiple amino acid alignments of the PHTs were generated using ClustalW (Larkin et al., 2007). Phylogenetic tree was constructed using the neighbor-joining method with the 1000 bootstrap in MEGA 6 (http://www.megasoftware.net/).

**Chromosomal Locations and Gene Structures**
The chromosomal locations and gene structures of the PHT genes were obtained from Phytozome 11.0. The chromosomal locations of the PHT genes were mapped with MapDraw program (Liu and Meng, 2003). The exon/intron organization of the individual PHT gene was drawn using Gene Structure Display Server (GSDS) program (Hu et al., 2015).

**Cis-Regulatory Element Identification**
Promoter sequences located 2 kb upstream of the transcription start sites (ATG codons) of the *Populus* PHT genes were retrieved from the *Populus trichocarpa* genome v3.0 in Phytozome 11.0 (http://phytozome.jgi.doe.gov/pz/portal.html). The cis-regulatory elements were scanned in PLACE (Higo et al., 1999).

**RNA Isolation and Quantitative Real Time PCR Analysis**
Total RNA was isolated from roots, stems, new leaves, and old leaves using a plant RNA extraction kit (R6827, Omega Bio-Tek, GA, USA) according to the manufacturer’s instructions. The first-strand cDNA was synthesized using a PrimeScript RT Reagent Kit (DRR037S, Takara, Dalian, China) following the manufacturer's instructions. The specific primers (Table S1) for quantitative real time PCR analysis were designed by Primer Premier 6.0 (Premier Biosoft, Palo Alto, CA, USA). We performed PCR in a 20 µl reaction mixture containing 10 µl of 2X SYBR Green Premix Ex Taq II (Biotek, China), 2 µl of cDNA, and 1 µl of 20 mM of each primer (Table S1) using a Roche LightCycle 96 machine (Roche, Germany). PCR amplification was performed under the following conditions: one cycle of 2 min at 95°C, followed by 45 cycles at 95°C for 10 s, 55°C for 20 s, and 72°C for 20 s. Actin2/7 was used as a reference gene (Brunner et al., 2004b). Three biological replicates with three technical replicates were assayed for each sample. Reactions for the reference gene were included in each plate. The relative expression levels of all the *PtPHT* genes were calculated using the $2^{-\Delta\Delta CT}$ method (Livak and Schmittgen, 2001).

**RESULTS**

**Identification and Phylogenetic Analysis of the *PtPHT* Gene Family**
We identified a total of 42 putative *PtPHT* genes, including 12 *PHO*, 14 *PtPHT1*, 2 *PtPHT2*, 6 *PtPHT3*, and 8 *PtPHT4* genes from the genome of *P. trichocarpa* according to the full *AtPHT* family in *Arabidopsis* (Table 1). The lengths of the encoded proteins ranged from 314 amino acids to 802 amino acids, and their sequences contained 5 to 14 TMDs. The molecular weight (MW) of 42 putative proteins ranged from 33.5 to 93.5 kDa. The GRAVY value of putative phosphate transporter protein (PHT family) was positive and ranged from 0.128 to 0.795, while that of PHO family was negative ranging from $-0.255$ to $-0.064$.

The amino acid sequences of all of the PHT genes of *P. trichocarpa* were aligned with those of *A. thaliana*. The resulting phylogenetic tree contained five clusters with PHO, PHT1, PHT2, PHT3, and PHT4, respectively (Figure 1). Based on the phylogenetic analysis results, the *P. trichocarpa* PHT genes were named after the *A. thaliana* PHT genes (Figure 1, Table 1).

**Chromosomal Location and Structural Analyses of *PtPHT* Genes**
The *PtPHT* genes were marked on a physical map of *Populus* linkage groups based on the location information for *PHTs* in Phytozome 11.0. The *Populus* *PtPHT* genes showed a heterogeneous distribution pattern among the chromosomes (Figure 2). A total of 41 of the 42 *PtPHT* genes were localized to 15 of the 19 linkage groups of *Populus* (Figure 2). Only *PtPHO12* was localized to unattributed scaffold fragments.

The lengths of the 42 *PtPHT* genes ranged from 945 to 2409 bp, and they contained 1 to 16 exons (Table 1). These genes were distributed across 15 of the 19 poplar chromosomes excluding chromosomes 3, 7, 11, and 13 (Table 1, Figure 2). In addition, the lengths of the 12 *PtPHO* genes ranged from 1890 to 2409 bp, and they contained 11 to 14 exons (Table 1). Most of these genes were localized to chromosomes 8 and 10 (Table 1). Further, the lengths of the 14 *PtPHT1* genes ranged from 1311 to 1620 bp, and they contained 1 to 3 exons. Most of these genes were located on chromosomes 5 and 10 (Table 1). Two members of the PHT2 family, *PtPHT2.1* and *PtPHT2.2*, had lengths of 963 and 1773 bp and contained 2 and 3 exons, respectively. The PHT3 family included six *PtPHT* genes in poplar, with lengths ranging from 945 to 1125 bp and containing 6 exons. Further, the PHT4 family included nine *PtPHT* genes, with lengths ranging from 864 to 1662 bp and containing 1 to 11 exons.

We further analyzed the exon/intron structures of the 42 *PtPHT* genes. The genes belonging to the same PHT family, with the exception of the PHT2 family, were well conserved in terms of their exon/intron structures, with similar exon lengths (Figure 3). Unexpectedly, compared with other members of the PHT4 family, which contained 7–16 exons, *PtPHT4.6* possessed only one long exon, and its coding sequence was shorter. These substantial differences in gene structure are due to differences in the numbers of exons and introns among the various genes.
| Gene name | ID     | Gene Genome location | ORF (bp) | Protein size | MW (kDa) | PI | GRAVY | Exon Number | Sub-cellular location |
|-----------|--------|----------------------|----------|--------------|----------|----|-------|--------------|-----------------------|
| PtPHT1.1  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.2  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.3  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.4  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.5  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.6  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.7  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.8  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.9  | Potri.001G322300 PHT1 Chr01:25927053-2593243  | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.10 | Potri.001G322300 PHT1 Chr01:25927053-2593243 | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.11 | Potri.001G322300 PHT1 Chr01:25927053-2593243 | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.12 | Potri.001G322300 PHT1 Chr01:25927053-2593243 | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.13 | Potri.001G322300 PHT1 Chr01:25927053-2593243 | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.14 | Potri.001G322300 PHT1 Chr01:25927053-2593243 | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.15 | Potri.001G322300 PHT1 Chr01:25927053-2593243 | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
| PtPHT1.16 | Potri.001G322300 PHT1 Chr01:25927053-2593243 | 6 1587 528 58 10.07 0.396 8 10 11 chlo: 9, mito: 4 |  |  |  |  |  |  |  |
### Table 1 (Continued)

| Gene Name ID | Gene name | Gene family | Genome location | ORF (bp) | Protein size | MW (kDa) | PI | GRAVY | Exon Number | TMD (Outside to Inside) | TMD (Inside to Outside) | Subcellular location prediction |
|--------------|-----------|-------------|-----------------|----------|--------------|---------|----|-------|-------------|------------------------|--------------------------|----------------------------------|
| PtPHO7       | Potri.010G164900 | PHO | Chr10:16885666-16889577 | 229      | 763          | 88.7    | 9.47 | −0.246 | 11         | 10, E.R.: 2, nucl: 1   | 0                       | plas: 10, E.R.: 2, nucl: 1, E.R.: 1, perox: 1 |
| PtPHO8       | Potri.010G165300 | PHO | Chr10:16905446-16909093 | 240      | 802          | 93.5    | 9.33 | −0.192 | 12         | 10, cyto: 1, nucl: 1   | 0                       | plas: 10, E.R.: 1, perox: 1, cyto: 1, nucl: 1 |
| PtPHO9       | Potri.010G165800 | PHO | Chr10:16935711-16945240 | 238      | 795          | 92.8    | 9.29 | −0.197 | 10         | 10, E.R.: 1, perox: 1, ER: 1, nucl: 1 | 0                       | plas: 10, E.R.: 1, cyto: 1, nucl: 1, ER: 1 |
| PtPHO10      | Potri.016G032900 | PHO | Chr16:1872547-1876343  | 2325     | 774          | 90.5    | 9.44 | −0.177 | 12         | 10, cyto: 1, nucl: 1   | 0                       | plas: 11, cyto: 1, ER: 1, nucl: 1 |
| PtPHO11      | Potri.017G086800 | PHO | Chr17:10433862-10439838 | 23       | 79          | 92.6    | 9.26 | −0.064 | 11         | 11, E.R.: 1, cyto: 1, nucl: 1 | 0                       | plas: 12, vac: 1, E.R.: 1, nucl: 1 |
| PtPHO12      | Potri.T146700  | PHO | scaffold_467:326-5065  | 1890     | 629          | 72.8    | 9.32 | −0.209 | 12         | 7                       | 0                       | E.R.: 4, nucle: 3, cyto: 2, nucl: 2, plas: 2 |

### Figure 1

**Phylogenetic Tree of P. trichocarpa and A. thaliana PHT Proteins.** The phylogenetic tree was constructed using the neighbor-joining method with the 1000 bootstrap in MEGA 6 (http://www.megasoftware.net/).

### cis-Regulatory Elements of PtPHT Genes in Poplar

The 2 kb upstream regions of the 42 PtPHT genes were analyzed using PLACE signal scan program. A total of...
234 putative cis elements were detected, many of which are involved in development, stress, or hormone activity (Table S2). Twenty-six of the detected cis elements were found in all of the PtPHT genes involved in energy, light, hormone, defense, carbon metabolism, and tissue-specific expression. For example, GATABOX, GT1CONSENSUS, and IBOXCORE are light-responsive cis elements, and CACTFTPPCAL is carbon metabolism related cis element (Gowik et al., 2004).

Some Pi-response and other stress-related cis regulatory elements including ABRE elements, helix–loop–helix elements, PHO-like, PHR1, TATA-box-like, and WRKY1 were detected in the upstream regions of PtPHT genes. AACACOREOSGLUB1, a WRKY1 element, and ACGTABREMOTIFA2OSEM, an ABRE element, were present in 22 and 7 PtPHT genes, respectively. In addition, a PHO-like element (AACACOREOSGLUB1) and a TATA-box-like element (TATABOX2) was detected in 22 and 34 PtPHT genes, respectively; A PHR1 element (P1SB) was observed in 20 PtPHT genes; and a helix–loop–helix element, (CATATGGMSAUR) was present in 11 PtPHT genes. Furthermore, 17 PtPHT genes had typical cis elements that were present especially in the respective genes, and the total number of these cis elements was 32.

Transcription Patterns of PtPHT Genes in Various Tissues

The specificities of tissues or organs may allow for the detailed elucidation of their functions. We measured the transcript levels of all of the PtPHT genes in 12 various tissues of Populus tremula using a public database, PopGenIE (http://popgenie.org/) (Figure 4). These tissues included roots, dormant cambium-phloem, mature petiole, freshly expanded leaves, expanding young leaves, mature leaves, suckers, dormant buds, dormant flowers, expanded flowers, expanding leaves, and mature seeds. Twenty-five PtPHT genes were highly expressed in the roots of P. tremula, suggesting that most PtPHT genes might be involved in Pi uptake. Six PtPHT genes, including PtPHO3, PtPHO4, PtPHO5, PtPHO6, PtPHT1.4, and PtPHT1.14, were highly expressed in more than six of the twelve investigated tissues. For example, PtPHO6 was highly expressed in nine tissues, including cambium, petiole, expanded freshly leaves, expanding young leaves, mature leaves, suckers, dormant buds, dormant flowers, expanding flowers, and mature seeds. In contrast, the expression levels of several PtPHT genes, including PtPHO3.3 and PtPHT3.3b, were very low in all of the investigated tissues. In particular, PtPHT1.9 and PtPHT1.10 were not expressed in
any of the 12 tissues examined; this finding is not shown in Figure 4. Further, some PtPHT genes were only highly expressed in a few specific tissues. For example, PtPHT1.6 was exclusively expressed in the roots of *P. tremula*, while PtPHT1.5 and PtPHO9 were only highly expressed in the roots and suckers of *P. tremula*.

Real-time quantitative RT-PCR was also performed to analyze the transcript levels of all 42 PtPHT genes in the roots, stems, young leaves, and mature leaves of *P. simonii*. Transcripts of 35 PtPHT genes were detected in the examined tissues (Figure 5), but some of them were present at relatively low levels (such as PtPHT1.5/1.6, PtPHO1, and PtPHO2) (Figures 5A-D). Most of the PtPHT genes were confirmed to be expressed in the roots of poplar. Transcripts of 7 PtPHT genes (such as PtPHT1.3, PtPHT2.2, and PtPHT4.1a) were detected in all four tested tissues (Figures 5A-D). Most PtPHT genes were expressed in the roots of *P. simonii* at high levels. PtPHT1.8 and PtPHO7 exhibited increased transcript accumulation in the leaves compared to the roots (Figures 5A,D). In addition, PtPHT3.1a/b expression was similar in all four investigated tissues (Figure 5B).

**PtPHT Transcript Patterns in Response to Different Phosphate Levels**

To better understand the functions of PtPHT genes in relation to phosphate uptake, the transcription patterns of
PtPHT genes in the roots of *P. simonii* were examined at varying phosphate levels. There were 33 *PtPHT* genes in total expressed in roots (Figure 6), but the expression of a few *PtPHT*5s, such as *PtPHT3.2a/b* and *PtPHO1/2/3*, was very low (Figures 6B,D). In the roots, the expression of *PtPHT1.12* was up-regulated and that of *PtPHT1.5* was only slightly altered under phosphate starvation conditions; in addition, the expression of *PtPHT1.2, PtPHT1.3, PtPHT2.2, PtPHT3.1b*, and *PtPHT4.1a* was markedly up-regulated under low-phosphate conditions, and that of *PtPHT3.3a, PtPHT4.1b, PtPHT4.5b, PtPHO8*, and *PtPHO9* was up-regulated under high-phosphate conditions (Figures 6A–D).

Most of the *PtPHT* genes were expressed at high levels in the roots of *P. simonii* in the presence of low phosphate levels, indicating that they are high-affinity genes. The accumulation of transcripts of many *PtPHT* genes was reduced when the level of H$_2$PO$_4^{-}$ was increased in the roots of *P. simonii*, with the exception of *PtPHO8* and *PtPHO9*.

### Changes in PtPHT Transcription in Response to Drought Stress

To understand the transcriptional changes of *PtPHT* genes induced by drought, *P. simonii* plants supplied with different phosphate levels were subjected to drought stress for 2 weeks. *PtPHT1.2* and *PtPHO9* expression was found to be increased under drought conditions, irrespective of the phosphate level. In particular, *PtPHT1.2* expression was significantly induced by approximately 90-fold. However, changes in the transcription of some *PtPHT* genes under drought stress were largely dependent on the phosphate level. For example, after 14 days of drought stress, the expression of *PtPHT3.3b, PtPHT4.3*, and *PtPHT4.5a* was significantly down-regulated, while that of *PtPHT1.9, PtPHO4*, and *PtPHO9* was only slightly increased when no phosphate was supplied (Figure 7). However, the expression of these genes was up-regulated under drought stress in the presence of a low level of phosphate. Further, in the presence of a high level of phosphate, the expression of *PtPHT1.9, PtPHT1.11, PtPHT1.3, PtPHT1.4, PtPHT2.2, PtPHT4.4, PtPHT4.6, PtPHO1*, and *PtPHO8* was increased under drought conditions. These data suggested that the changes in the transcription of some *PtPHT* genes under drought conditions were strongly influenced by the phosphate level.

### DISCUSSION

Phosphorus is a mineral nutrient that is essential for plant growth and development. Pi was assimilated in roots via PHT transporters. In this study, we identified the whole PHT gene family in poplar and examined the expression patterns of these genes in different plant tissues and in response to differing Pi levels and drought. The PHT family, the cis elements of the PHT genes, and transcriptional changes in the PHT genes in response to different Pi levels and drought are discussed below.

### PHT Gene Family in Populus

In the current study, we conducted comprehensive analysis of PHT genes in poplar. A total of 42 *PtPHT* genes were retrieved from *Populus* genome (Phytozome 11.0, *P. trichocarpa* v3.0) with improved annotation, and they were classified into five clusters: PHT1, PHT2, PHT3, PHT4, and PHO. In a subsequent study, Loth-Pereda et al. (2011) reported the identification of 12 PHT1 family members in an earlier version of the *Populus* genome (v3), and we expanded upon their findings with the identification of 14 PHT1 family members in the latest version of *Populus* genome (v3). Notably, the PHT1 family is larger in poplar (14 genes) than in *Arabidopsis* (9 genes), barley (8 genes), rice (13 genes), and tomato (8 genes) (Muchhal et al., 1996; Rae et al., 2003; Liu et al., 2011; Chen et al., 2014).

The PHT2, PHT3, PHT4, and PHO families contained more members of *PtPHT* genes in poplar than that in *Arabidopsis*. One PHT2 family gene has been identified in *Arabidopsis* (Daram et al., 1999), while two (*PtPHT2.1* and *PtPHT2.2*) were detected in poplar. These two genes are located on different chromosomes (Chr8 and Chr10), and the lengths of their encoded proteins greatly differ which is consistent with
the previous report that AtPHT2.1 cDNA encodes a 61-kD protein that is structurally similar but different from the PHT1 family proteins in Arabidopsis (Daram et al., 1999). However, phylogenetic analysis showed that PtPHT2.1 and PtPHT2.2 share 100% similarity (Table S3). At least three AtPHT3 genes have been identified in Arabidopsis to date (Rausch and Bucher, 2002), while we found six PHT3 family genes in poplar. Further, six PHT4 family genes have been characterized in Arabidopsis (Guo et al., 2008), while 8 PtPHT4 genes were detected in the present study. Ten genes with homology to AtPHO1 are present in the Arabidopsis genome (Hamburger et al., 2002), and in this study, 12 PHO family genes were detected with similarities to the PHO1 protein in Arabidopsis.

The presence of a large number of genes within a family reflects the successive expansion and rearrangement of the genome by the extensive duplication and diversification that frequently occur over the course of evolution (Wang et al., 2007). The P. trichocarpa genome is duplicated and contains more protein-coding genes than the Arabidopsis genome; on average, there are 1.4 to 1.6 putative Populus genes for every Arabidopsis gene (Tuskan et al., 2006). Whole-genome duplication increases gene gains and losses, which greatly affects the amplification of members of gene families in the genome (Guo, 2013). The expanded genes in the PHT family, including PHT1, PHT2, PHT3, PHT4, and PHO, in poplar compared with Arabidopsis might be the result of gene duplication. Gene duplication has been reported in P. trichocarpa in many studies, including studies of the glutamine synthetase family, nitrate transporter gene family, and CCCH zinc finger family (Castro-Rodriguez et al., 2011; Chai et al., 2012; Bai et al., 2013). In the present study gene expansion and loss might also occur in the PHT family of P. trichocarpa similar to that of Arabidopsis. For example, two closely related orthologs of Arabidopsis, AtPHT4.1, AtPHT4.5, and AtPHO1, were found in Populus suggesting that duplications might be a partial reason for the expansion of PHT gene family in Populus. Forty-three PtPHT genes were localized to 15 poplar chromosomes, and some PHT genes were found to be clustered together as closely related genes (Figure 2). The presence of these closely related clusters suggests that these genes might evolve from regional duplications (e.g., PtPHO6, PtPHO7, PtPHO8, and PtPHO9).

### The cis Elements Identified in Poplar

Cis elements may control the efficiency of promoters and thus regulate the expression of the genes that they control by interacting with the corresponding trans-regulatory factors (Liu et al., 2011). Studies of cis elements might play a crucial role in dissecting the functions of genes. A previous study has described the cis elements of PHT1 family members in poplar (Loth-Pereda et al., 2011), but the cis elements of all PHT gene family members in poplar have not yet been identified. In this study, the upstream regions of all the PtPHTs contain cis elements affecting energy, light, hormone, defense, carbon metabolism, and tissue-specific expression. These findings suggest that the expression of these PtPHT genes might be regulated by many factors, such as light-, hormone-, and defense-related factors. For instance, GATABOX and GT1CONSENSUS are cis elements required for light regulation (Lam and Chua, 1989; Terzaghi and Cashmore,
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**FIGURE 6 |** The expression of PtPHT genes (A: PHT1; B: PHT2 and PHT3; C: PHT4; D: PHO) under different phosphate levels in roots of *P. simonii* seedlings. The relative expression of PHT1.2 was set as a standard (100). Total RNA was isolated using a plant RNA extraction kit (R68 27, Omega Bio-Tek, GA, USA) according to the manufacturer’s instructions. The RNA was used for qRT-PCR with PHT-specific primers. The signal intensities were calibrated based on a constitutively expressed poplar actin gene (Brunner et al., 2004b). The data are presented as the mean ± SD of three separate measurements.

1995), and they have also been identified in the upstream regions of all OsPHT genes in rice (Liu et al., 2011). Further, putative PHO-like (CGCGTGGG) and TATA-box-like (TATAAATA) cis regulatory elements have been identified in the promoter regions of *Arabidopsis* genes (Hammond et al., 2003), and they were observed in 22 and 34 of the 42 PtPHT promoters, respectively. Alternatively, because the core sequence of the *Arabidopsis* PHO-like element (CACGTG) is also similar to that of the ABRE element (ABREOSRAB21), which is present in 6 of the 42 PtPHT promoters, proteins in the bZIP class of transcription factors might bind to them (Schindler et al., 1992). The presence of these cis elements in poplar PtPHT genes and their likely roles in regulating gene expression suggest that PtPHT genes may be involved in various stress responses in poplar.

Some cis elements function under phosphate deficiency, for example, the P1BS element can be bound to PHR1 (a MYB transcription factor) to potentially regulate the phosphate starvation response in *Arabidopsis* (Rubio et al., 2001), PHR1 has been suggested as a central integrator to role in the transcriptional regulation of phosphate starvation responses (Bustos et al., 2010). In the current study, P1BS-like elements were found in 20 of the 42 PtPHT promoters examined suggesting that expression of these genes might be influenced by Pi starvation. Similarly, P1BS-like elements have also been detected in OsPHT genes in rice (Liu et al., 2011). These elements are found in the promoters of phosphate regulated genes, not only in *Arabidopsis* but also in other plant species such as barley and rice (Schünmann et al., 2004; Liu et al., 2011), which suggests that they may participate in a conserved signaling pathway for the phosphate starvation response in plants. The existence of P1BS-like elements in poplar, rice, and *Arabidopsis* suggests that they may serve a similar role in both woody species and herbs.

**Transcript Profiles of PHT Genes in Various Tissues**

The number of Pi transporters present in roots may reflect the complexity and significance of the process of Pi transport. In the current study, most PHT family members were expressed in the roots of *P. tremula* and *P. simonii*. In particular, most PHT1 family members were highly expressed in the roots of poplar, consistent with a previous study conducted by Loth-Pereda et al. (2011) suggesting that they may function in Pi uptake. Notably, most PHT1 genes in other plant species are also expressed in
FIGURE 7 | The expression of PHT genes affected by drought at different phosphate levels (A: No P; B: low P; C: high P) in roots of *P. simonii* seedlings. The control is defined as 1 in the figures. Total RNA was isolated using a plant RNA extraction kit (R6827, Omega Bio-Tek, GA, USA) according to the manufacturer’s instructions. The RNA was used for qRT-PCR with PHT-specific primers. The signal intensities were calibrated based on a constitutively expressed poplar actin gene (Brunner et al., 2004b). The data are presented as the mean ± SD of three separate measurements.

the roots. For example, eight of the nine PHT1 family genes in *Arabidopsis* are expressed in the roots (Bucher et al., 2001; Mudge et al., 2002), in addition, 8 PHT1 genes in tomato (Chen et al., 2014) and 10 of the 13 PHT1 genes in rice are expressed in the roots (Paszkowski et al., 2002). These results indicate that PHT1 family genes may play similar roles in Pi uptake in different species.

Homologous PHT1, PHT2, PHT3, PHT4, and PHO genes showed substantial differences in their regulation in different tissues. Nine PHT1 family genes were highly expressed in the roots of *P. tremula*, but the expression of the other five genes was very low, and two of these genes (*PtPHT1.9* and *PtPHT1.10*) were not expressed in any tissues examined. *PtPHT2.1* was highly expressed in the roots, cambium-phloem, mature petiole, and dormant buds, while *PtPHT2.2* was mainly expressed in the roots, cambium-phloem, and mature leaves. Six homologous *PtPHT3* genes exhibited substantial differences in their expression in the various tissues examined. Three of them (*PtPHT3.1b, PtPHT3.2b, and PtPHT3.3a*) were highly expressed in the roots of *P. tremula*, while the others (*PtPHT3.1a, PtPHT3.2a, and PtPHT3.3b*) were expressed at very low levels. Although most PHT4 family members, excluding *PtPHT4.3* and *PtPHT4.6*, were highly expressed in the roots and leaves, *PtPHT4.1* was also highly expressed in the cambium-phloem, dormant buds, and dormant flowers. Further, *PtPHO5* and *PtPHO6* were highly expressed in nine tissues examined, while *PtPHO7* and *PtPHO8* were exclusively highly expressed in only a few of the tissues. Expression of the *PtPHT* genes varied among the different tissues examined, suggesting that these genes may have different functions in Pi uptake, transport, and storage. These results suggest that the PHT genes may have different functions since they expressed in a wide range of tissues, which may be partly crucial for the growth of perennial species.

The differential expression of *PtPHT* genes in different tissues might be related to their subcellular locations (Table 1). For example, two PHT2 family members were expressed in the leaves of *P. simonii*, in agreement with a previous study demonstrating that the *AtPHT2.1* gene, which is located in the chloroplast, is predominantly expressed in green tissues in *Arabidopsis* (Liu et al., 2011). In addition, the expression patterns of *PtPHT4* genes in both roots and leaves, and the phosphate transport activities as well as their subcellular locations, indicate that PHT4 proteins play roles in transferring Pi between the cytosol and chloroplasts, heterotrophic plastids, and the Golgi apparatus (Guo et al., 2008). For example, *AtPHT4.6* transports Pi out of the Golgi lumenal space to recycle Pi released from glycosylation processes in *Arabidopsis* (Cubero et al., 2009). In addition, the PHO1 protein has been reported to be localized to stellar cells of the roots as well as the lower part of the hypocotyl in *Arabidopsis* (Ribot et al., 2008). In the present study, PHO family members were found to be highly expressed in many of the tissues examined, suggesting that they have diverse functions in loading Pi to different parts of plants.

**Transcript Profiles of PHT Genes in Response to Different Phosphate Levels and Drought in *Populus***

The transcription of *PtPHT* genes differed at different Pi levels. Upon Pi starvation, *PtPHT1.9, PtPHT1.11, PtPHT1.12,
PtPHT3.3b, PtPHT4.3, and PtPHT4.5a expression was up-regulated in P. simionii (Figure 5). The up-regulation of some PtPHTs, such as PtPHT9 and PtPHT11, under Pi starvation is consistent with the findings of a previous study of P. trichocarpa conducted by Loth-Pereda et al. (2011). Most PHT1 family genes expressed in roots show up-regulation in phosphate deprived Arabidopsis plants; in particular, PtPHT9 and PtPHT11 in poplar are highly homologous to AtPHT8 and AtPHT9, which have been shown to be strongly up-regulated during Pi starvation (Mudge et al., 2002). In rice, high-affinity Km values of OsPHT9 and OsPHT10 for Pi transport have been reported, as well as specific induction of their expression by Pi starvation (Wang et al., 2014). Therefore, some of these PHT transporters likely play roles in Pi scavenging under conditions of low soil Pi availability. PtPHT1.2, PtPHT1.3, PtPHT2.2, PtPHT3.1b, and PtPHT4.1a expression was markedly up-regulated under low-Pi conditions, suggesting that their expression might be induced under these conditions. The transcript levels of a few PHT genes in Arabidopsis, such as AtPHT1, AtPHT2, and AtPHT4, were increased by Pi deficiency (Ribot et al., 2008); however, the expression of most PHO family members was not induced by Pi deficiency in poplar, suggesting that PHT genes might have different functions in woody species and herbs. PtPHT3.3a, PtPHT4.1b, PtPHT4.5b, PtPHO8, and PiPHO9 expression was up-regulated under high-phosphate conditions, indicating that these proteins might be low-affinity transporters. In addition, some PHT4 family members might be low-affinity transporters, consistent with the relatively high Pi levels that have been reported in the cytosol and other subcellular compartments (Mimura, 1999).

The expression of PHT genes varied at different Pi levels, suggesting that PHT family members are likely to encode both high- and low-affinity transporters. For example, OsPHT2 shows low-affinity characteristics on the mM scale in oocytes (Ai et al., 2009). By contrast, a Km value of 19 μM has been demonstrated to be a high-affinity transporter (PtPHT3.3) in barley (Buchert, M., Rausch, C., and Daram, P. (2001). Molecular and biochemical mechanisms of phosphorus uptake into plants. J. Plant Nutr. Soil Sci. 164, 209–217. doi: 10.1002/1522-2624(200104)164:2<209::AID-JPLN209>3.0.CO;2-F

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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