SPX1 is an important component in the phosphorus signalling network of common bean regulating root growth and phosphorus homeostasis

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Abstract

Proteins containing the SPX domain are believed to play vital roles in the phosphorus (P) signalling network in plants. However, the functions of SPX proteins in legumes remain largely unknown. In this study, three SPX members, *PvSPX1–PvSPX3* were cloned from common bean (*Phaseolus vulgaris* L.). It was found that the transcripts of all three *PvSPX* members were significantly enhanced in both bean leaves and roots by phosphate (Pi) starvation. Among them, the expression of nuclear localized *PvSPX1* showed more sensitive and rapid responses to Pi starvation. Consistently, only overexpression of *PvSPX1* resulted in increased root P concentration and modified morphology of transgenic bean hairy roots, such as inhibited root growth and an enlarged root hair zone. It was further demonstrated that *PvSPX1* transcripts were up-regulated by overexpressing *PvPHR1*, and overexpressing *PvSPX1* led to increased transcripts of 10 Pi starvation-responsive genes in transgenic bean hairy roots. Taken together, it is suggested that *PvSPX1* is a positive regulator in the P signalling network of common bean, and is downstream of *PvPHR1*.

Key words: Bean, hairy roots, phosphate starvation, phosphorus homeostasis, root growth, SPX domain.

Introduction

Phosphorus (P) is an essential element for plant growth, and is easily fixed by soil particles due to its chemical properties. Therefore, low P availability adversely affects crop growth and production, especially on acid soils (Raghothama, 1999; Vance et al., 2003). To cope with low P stress, plants have developed a wide range of adaptive strategies, such as changes in root architecture and morphology (Liao et al., 2004; Zhou et al., 2008; Péret et al., 2011; Tian et al., 2012), increased exudation of protons and organic acids (Fox and Comerford, 1990; Ström et al., 2005; Taghipour and Jalali, 2012), and enhanced secreted or root-associated acid phosphatase activities (Del Pozo et al., 1999; Bozzo et al., 2002; Ligaba et al., 2004; C. Wang et al., 2009; Liang et al., 2010; Robinson et al., 2012; L.S. Wang et al., 2012). These adaptive strategies are tightly mediated by the P signalling network, which is composed of a wide array of regulators (Raghothama, 1999; Vance et al., 2003; Chiou and Lin, 2011).

Proteins containing the SPX domains have been demonstrated to play vital roles in the P signalling networks of yeast (*Saccharomyces cerevisiae*), Arabidopsis (*Arabidopsis thaliana*), rice (*Oryza sativa*), and rape (*Brassica napus*) (Ligaba et al., 2004; Duan et al., 2008; C. Wang et al., 2009, 2012; Z. Wang et al., 2009; Liu et al., 2010; Secco et al., 2012a, b; Yang et al., 2012). The SPX domain is named after SYG1/Pho81/XPR1 proteins, which contain a conserved domain in the N-terminal peptides of yeast SYG1 and PHO81, and human XPR1 proteins (Spain et al., 1995; Lenburg and O'Shea, 1996; Battini et al., 1999; Wang et al., 2004).

In yeast, several SPX domain-containing proteins involved in P acquisition and the signalling pathway have been identified (Secco et al., 2012b). PHO81 is a cyclin-dependent kinase
(CDK) inhibitor (Lenburg and O’Shea, 1996). Under phosphate (Pi) starvation conditions, PHO81 inhibits the kinase activity of the PHO80–PHO85 complex against the Pho4 transcription factor, which subsequently regulates transcripts of several Pi starvation-responsive genes (Lenburg and O’Shea, 1996). Many yeast Pi transporters, such as Pho84, Pho87, Pho89, Pho90, and Pho91, also possess the SPX domain (Secco et al., 2012b). It is interesting that most SPX domain-harbouring proteins, including Vtc2, Vtc3, Vtc4, and Gde1, appear to play key regulatory roles in Pi homeostasis in yeast (Secco et al., 2012b).

In plants, four groups of proteins were also found to contain the SPX domain. Among them, three groups of proteins have the SPX domain in the N-terminus and other domains in the C-terminus, including an EXS (ERD1, XPR1, and SYG1), a major facility superfamily (MFS), or a RING-type zinc finger domain (Hamburger et al., 2002; Stefanovic et al., 2007; Lin et al., 2010; Secco et al., 2010; Kant et al., 2011; C. Wang et al., 2012). Similar to the functions of proteins containing the SPX domain in yeast, most of these plant members are involved in regulating Pi homeostasis in plants. Examples include OsSPX-MFS1 in rice (C. Wang et al., 2012), along with AtPHO1:1 and AtNLA in Arabidopsis (Stefanovic et al., 2007; Secco et al., 2010; Kant et al., 2011).

Recently, a specific group of proteins only containing the SPX domain have been characterized in plants, such as Arabidopsis and rice (Duan et al., 2008; C. Wang et al., 2009; Z. Wang et al., 2009; Liu et al., 2010; Yang et al., 2011). In Arabidopsis, four members only contain the SPX domain, namely AtSPX1, AtSPX2, AtSPX3, and AtSPX4 (Duan et al., 2008). Furthermore, expression patterns of several Pi starvation-responsive genes were positively and negatively regulated by AtSPX1 and AtSPX3, respectively (Duan et al., 2008). Similarly, the negative regulatory role of OsSPX1 was also suggested in rice, because transcription of several Pi starvation-responsive genes (e.g. OsPT2, OsPT6, and OsPAP10) was suppressed through OsSPX1 overexpression (C. Wang et al., 2009). Furthermore, it has recently been determined that OsSPX1 is downstream of OsPHR2 and OsPHO2 in the rice Pi signalling pathway (Liu et al., 2010).

Despite accumulated knowledge of the Pi signalling network in model plants (i.e. Arabidopsis and rice) (Chiou and Lin, 2011), information on Pi starvation-responsive pathways in other crops remains fragmentary. A group of Pi starvation-responsive genes (e.g. PvmiR399 and PvPS2:1) have been cloned and characterized in common bean (Phaseolus vulgaris L.), an important legume crop (Tian et al., 2007; Valdes-Lopez et al., 2008; Hernández et al., 2009; Liang et al., 2012a,b). This has facilitated elucidation of the Pi signalling network in bean, although this knowledge remains incomplete. Recently, essential roles for PvPHR1 and PvmiR399 have been suggested in Pi deficiency signalling (Valdes-Lopez et al., 2008). Nevertheless, other regulators are probably required as well. In a previous study, three expressed sequence tags (ESTs) with high homology to AtSPX1 were identified through screening a suppression subtractive hybridization library constructed from P-deficient bean (Tian et al., 2007). Among them, the full-length cDNA of PvIDS4-1 (i.e. PvSPX1) was cloned, and its expression levels were found to be up-regulated by Pi starvation in bean (Tian et al., 2007). However, the functions of PvSPX1 and other PvSPX genes in bean adaptation to P deficiency remain unknown. In this study, the full-length cDNA of the other two PvSPX genes (i.e. PvSPX2 and PvSPX3) was cloned. Subsequently, the expression patterns and functions of all three PvSPX gene family members as related to P availability were characterized in bean.

Materials and methods

Plant material and growth conditions

Seeds of common bean genotype G19833 were surface sterilized for 1 min using 10% (v/v) H2O2 and then germinated in the dark on germination paper moistened with 1/4 strength modified nutrient solution as described previously (Yan et al., 2001). Five days after germination, seedlings were transferred to nutrient solution supplied with 5, 50, 100, or 500 μM KH2PO4 for a P dosage experiment. After 10 d, young leaves and roots were harvested. For time course experiments, seedlings were pre-treated in 1/4 strength nutrient solution for 7 d and then transplanted to nutrient solution containing 5 μM KH2PO4. Shoots and roots were each harvested at 0, 4, and 8 d after treatment for determination of fresh weight, total root length, and P content. Young leaves and roots were separately harvested for RNA extraction. Nutrient solution was well aerated and its pH was maintained between 5.8 and 6.0. Four biological replicates were included in all of the experiments.

Analysis of total root length and P content

Roots were scanned, and then the digital images were analysed using Win-Rhizo software (Régent Instruments, Canada) to measure total root length. Shoots and roots were kept separately at 75 °C until completely dry, and then were ground into powder for total P content analysis. P content was determined using the phosphorus–molybdate blue colour reaction as previously described (Murphy and Riley, 1962)

Cloning full length cDNAs of PvSPX2 and PvSPX3

Gene-specific primers were designed according to the EST sequences of PvSPX2 (EG594307) and PvSPX3 (EG594308) (Tian et al., 2007) (Supplementary Table S1 available at JXB online). Using the full-length cDNA library constructed from the roots of G19833 as a template, the 5’ and 3’ termini of each gene were amplified by the specific primers paired with T3 and T7 primers, respectively. The amplified DNA fragments were then cloned into the pMD18-T vector (TaKaRa, Japan) and sequenced. Sequences of PvSPX2 and PvSPX3 were analysed at the National Center for Biotechnology Information (NCBI) website (http://www.ncbi.nlm.nih.gov/) and deposited in GenBank with accession numbers GU189405 for PvSPX2 and GU189406 for PvSPX3. Multiple sequence alignments were conducted using ClustalW 1.8. The phylogenetic tree was established using the Neighbor-Joining method of the MEGA 4.1 program.

RNA extraction and quantitative real-time PCR

Total RNA was isolated from young leaves and roots using RNAiso Plus reagent (TaKaRa) and treated with DNase I (TaKaRa). The first-strand cDNA was synthesized from total RNA using MMLV reverse transcriptase following the manual (Promega Inc., USA). The first-strand cDNA was then used for SYBR Green-monitored quantitative real-time PCR (qPCR) analysis, which was performed using a Rotor-Gene 3000 (Corbett Research, Australia). Expression levels of the tested genes were quantified relative to expression levels
of the reference gene EF-1α (PvTC3216) using arbitrary units. The primer pairs used for qPCR analysis are shown in [supplementary Table S1](#) at JXB online. All of the gene expression analyses had four biological replicates.

**Subcellular localization analysis**

The coding regions of *PvSPX1* (EF191350), *PvSPX2*, and *PvSPX3* without stop codons were separately cloned into the transient expression vector (pBEGFP-N1) and fused with green fluorescent protein (GFP; [Liang et al., 2010](#)). For subcellular localization of *PvSPXs* in onion (*Allium cepa*) epidermal cells, the *PvSPX–GFP* fusion constructs and GFP empty vector control were separately transformed into onion epidermal cells using a helium-driven accelerator (PDS/1000, Bio-Rad). After the transformed cells were cultured on Murashige and Skoog (MS) medium for 16 h, the GFP fluorescence was observed using a confocal scanning microscope system (TCS SP2, Leica, Germany) with 488 nm excitation and 500–525 nm emission filter wavelengths. For subcellular localization of *PvSPXs* in leaf epidermal cells of tobacco (*Nicotiana tabacum*), the *PvSPX–GFP* fusion constructs and GFP empty vector control were separately transformed into *Agrobacterium tumefaciens* strain GV3101, which were further used for transformation as previously described ([Sparkes et al., 2006](#)). After the transformation, plants were grown under normal conditions for 48 h and the GFP fluorescence was observed using a fluorescence microscope (Leica DM5000B). The GFP fluorescence was imaged using a Leica DFC 480 camera.

**Transformed genes in common bean hairy roots**

The coding regions of *PvSPX1*, *PvSPX2*, *PvSPX3*, and *PvPHR1* (EU500763) were inserted separately into the unique BamHI and MluI sites of the binary vector pYLRNai as previously described ([Liang et al., 2010](#)). For *PvSPX1* RNA interference (RNAi) construction, the same binary vector pYLRNai was used by inserting the *PvPSX1*-specific fragment into the BamHI and HindIII, and the *PvI* and MluI sites, respectively. The overexpression, RNAi constructs, and the empty vector control (CK) were then separately transformed into *Agrobacterium rhizogenes* containing the SPX domain could be divided into four groups, [Liang et al., 2012b](#). Briefly, sterilized bean seeds were germinated on half-strength MS medium. After 35 h, the abaxial sides of cotyledons were wounded with a scalpel previously dipped into the overnight cultures of the transgenic *A. rhizogenes* strain K599. The wounded cotyledons were cultured in solid MS medium to develop hairy roots. The expression levels of the corresponding genes in hairy roots grown under high P conditions. Subsequently, qPCR analysis for *PvPHR1*, *PvPT1*, *Pv4*, *PvPAP1*, *PvPAP2*, *PvPAP3*, *PvPAP4*, *PvPAP5*, and *PvP2–1* were increased duration of P deficiency, plant fresh weight and total P content gradually decreased. After 8 d of Pi starvation, the total P contents of bean shoots and roots were reduced by 32% and 30%, respectively, as compared with high P conditions (Supplementary Table S2). Similarly, total root length was also significantly decreased by P deficiency. Total root length at 4 d and 8 d of P deficiency was reduced by 39% and 55%, respectively, as compared with under high P conditions (Supplementary Table S2).

**Identification and bioinformatics analysis of *PvSPX2* and *PvSPX3***

Based on the reported EST sequences of *PvSPX2* and *PvSPX3*, the full-length cDNAs of both *PvSPX2* and *PvSPX3* were cloned from a full-length cDNA library of G19833 subjected to P deficiency. The coding regions of *PvSPX2* and *PvSPX3* were 861 bp and 756 bp in length, respectively. Alignment analysis showed that *PvSPX2* and *PvSPX3* exhibited 75% and 50% similarity to *PvSPX1*, respectively.

**Phylogenetic analysis**

Phylogenetic analysis showed that plant proteins containing the SPX domain could be divided into four groups, namely SPX, SPX-EXS, SPX-MFS, and SPX-RING (Fig. 1). Furthermore, SPX proteins could be further subdivided into three groups. Among them, *PvSPX1*, *PvSPX2*, and *PvSPX3* belong to group I, which includes AtSPX1 and AtSPX2 in *Arabidopsis*, as well as OsSPX1 and OsSPX2 in rice (Fig. 1).

**Temporal expression patterns of *PvSPX* genes in response to Pi starvation**

The temporal expression patterns of the three *PvSPX* genes in bean leaves and roots were analysed by qPCR. As shown in Fig. 2, their expression levels were significantly increased over time and reached their highest levels after 8 d of low P treatment (Fig. 2). However, their expression patterns varied in leaves and roots at 4 d of P deficiency (Fig. 2). After 4 d of P deficiency, significantly increased transcription was observed for *PvSPX1* and *PvSPX2* in leaves, while for *PvSPX3* transcription was not increased either in leaves or in roots (Fig. 2).
This suggests that \( \text{PvSPX1} \) and \( \text{PvSPX2} \) respond to \( \text{Pi} \) starvation earlier than \( \text{PvSPX3} \) in bean.

**Dosage responses of \( \text{PvSPX} \) genes to \( \text{P} \) availability**

Expression patterns of the three \( \text{PvSPX} \) members studied here were tightly dependent on \( \text{P} \) availability in the medium (Fig. 3). Their highest transcript levels were observed in both leaves and roots supplied with 5 \( \mu \text{M} \) \( \text{P} \), and were decreased with increased \( \text{P} \) availability (Fig. 3). When the applied \( \text{P} \) concentration was increased to 500 \( \mu \text{M} \), transcription of each \( \text{PvSPX} \) gene was negligible (Fig. 3). However, slight differences existed among their expression patterns as related to \( \text{P} \) availability. Transcript levels of \( \text{PvSPX1} \) and \( \text{PvSPX2} \) in both leaves and roots declined significantly when the applied \( \text{P} \) concentration was increased from 100 \( \mu \text{M} \) to 500 \( \mu \text{M} \), but that of \( \text{PvSPX3} \) did not (Fig. 3), suggesting that expression of \( \text{PvSPX1} \) and \( \text{PvSPX2} \) might be more sensitive to \( \text{P} \) availability than that of \( \text{PvSPX3} \).

**Subcellular localization of \( \text{PvSPX} \) proteins**

To determine the subcellular localization, the coding regions of the three \( \text{PvSPX} \) genes were fused with the \( \text{GFP} \) reporter gene and transiently expressed in onion and tobacco epidermal cells. Subcellular localization was visualized by detecting \( \text{GFP} \) signal in the transformed onion and tobacco epidermal cells. The empty vector containing \( \text{35S:GFP} \) was used as a control. The results showed that the three \( \text{PvSPX} \) members were found in various subcellular localizations (Fig. 4). Signals of \( \text{GFP} \) fusion with \( \text{PvSPX1} \) and \( \text{PvSPX2} \) were only detected in the nuclei of onion and tobacco epidermal cells (Fig. 4). However, \( \text{GFP} \) fusion with \( \text{PvSPX3} \) was observed in many areas in onion and tobacco epidermal cells, suggesting...
that PvSPX3 might be localized in the cytoplasm and nuclei (Fig. 4).

Functional analysis of PvSPX genes in transgenic hairy roots

The functions of PvSPX genes were further analysed in bean transgenic hairy roots by overexpressing PvSPX1, PvSPX2, and PvSPX3. Significantly increased transcripts of the three PvSPX genes in the transgenic bean hairy roots were verified through qPCR analysis (Supplementary Fig. S1 at JXB online). Subsequently, the transgenic hairy roots were grown in MS medium with or without P application for 14 d. The results showed that only overexpressing PvSPX1 could inhibit hairy root growth, as reflected by reduced fresh weight of hairy roots under both P conditions (Fig. 5A, B). Compared with the control lines, the fresh weight of the PvSPX1 overexpression line was reduced by ~60% in high P and 40% in low P (Fig. 5B). Furthermore, the P concentration in the PvSPX1 overexpression line was higher than that in the control line by ~45% in high P and 30% in low P (Fig. 5C). In contrast, the fresh weight and P concentration of both PvSPX2 and PvSPX3 overexpression lines were similar to those in the control line at the two P levels (Fig. 5B, C). Similarly, suppressed PvSPX1 did not affect hairy root fresh weight and P concentration, compared with those in the empty vector (CK) controls (Supplementary Figs S2, S3).

Root morphology was further investigated in all hairy root lines at the two P levels through determination of the percentage of the root hair zones (i.e. the part of root zone with >10 root hairs per 1 mm root) in bean hairy roots. The percentage of the root hair zones in all hairy root lines was ~80% without P application (Fig. 6). With P application, the percentage of the root hair zones of CK, and PvSPX2 and PvSPX3 overexpression lines was decreased by >50% (Fig. 6). However, for the PvSPX1 overexpression line, applied P did not affect the percentage of the root hair zone (Fig. 6). Interestingly, similar results were also observed in PvSPX1 RNAi lines, in which the percentage of the root hair zone was not affected by P application (Supplementary Fig. S4 at JXB online). The results suggest that expression of PvSPX1 might regulate enlargement of the root hair zones at a high P level.

PvSPX participates in the P signalling network in bean

The expression patterns of 11 genes were investigated in the transgenic hairy roots overexpressing PvSPX1 in order to illustrate the regulatory role of PvSPX1 in the P signalling network.
Fig. 3. Dosage response of *Pv*SPX genes to P deficiency. Seedlings were grown in nutrient solution supplied with 5, 50, 100, or 500 μM KH₂PO₄. After 10 d, total RNA was isolated from leaves and roots for qPCR analysis. Expression levels of the tested genes were quantified relative to expression levels of the reference gene *EF-1α* (*Pv*TC3216) using arbitrary units. Each bar is the mean of four replicates with the standard error. Different letters represent significant differences at the 0.05 level.

Fig. 4. Subcellular localization of *Pv*SPXs. (A) Transient expression of the pBEGFP construct and *Pv*SPX–GFP fusion in onion epidermal cells. Scale bars=50 μm. (B) Transient expression of the pBEGFP construct and *Pv*SPX–GFP fusion in tobacco epidermal cells. Scale bars=20 μm. The first row shows the empty vector control, followed by *Pv*SPX1–GFP, *Pv*SPX2–GFP, and *Pv*SPX3–GFP constructs. Cells were observed by the green fluorescence of GFP and the *Pv*SPX–GFP proteins.
network in bean. Among them, nine genes were previously characterized as Pi starvation-responsive genes, namely two Pi transporters (\textit{PvPT1} and \textit{PvPHT2}), five purple acid phosphatases (\textit{PvPAP1}–\textit{PvPAP5}), \textit{Pv4}, and \textit{PvPS2:1}. The other two genes (\textit{PvLPR1-like} and \textit{PvPDR2-like}) exhibit high homology with \textit{AtLPR1} and \textit{AtPDR2}, respectively, which both regulate root growth in \textit{Arabidopsis}. The qPCR analysis showed that overexpressing \textit{PvSPX1} led to significantly increased transcription of \textit{PvSPX1} led to significantly increased transcription of 10 genes compared with the control line, namely \textit{PvPT1}, \textit{PvPHT2}, \textit{Pv4}, \textit{PvPAP1}–\textit{PvPAP5}, \textit{PvPS2:1}, and \textit{PvLPR1-like} (Fig. 7). Consistently, suppressed transcripts of \textit{PvSPX1} resulted in lower expression patterns of several genes—\textit{PvPHT2}, \textit{PvPAP3}, \textit{PvPS2:1}, and \textit{PvLPR1-like} (Supplementary Fig. S5 at JXB online). The results suggest that expression of these genes is positively regulated by \textit{PvSPX1}. However, expression levels of \textit{PvPDR2-like} were inhibited in the \textit{PvSPX1} overexpression lines and increased in the \textit{PvSPX1} RNAi lines, compared with those in the control line (Fig. 7; Supplementary Fig. S7), suggesting that \textit{PvPDR2-like} is negatively regulated by \textit{PvSPX1} in bean. Similarly, \textit{PvSPX2} overexpression resulted in increased transcripts of several genes downstream of \textit{PvSPX1}, except \textit{PvPDR2-like} (Supplementary Fig. S6), suggesting that \textit{PvSPX2} might have a similar regulatory role to \textit{PvSPX1}. However, overexpression of \textit{PvSPX3} did not affect expression patterns of genes downstream of \textit{PvSPX1} (Supplementary Fig. S6). Furthermore, significantly increased transcription of \textit{PvSPX1} was obviously observed in the transgenic bean hairy roots with overexpression of \textit{PvPHR1} (Fig. 8), suggesting that \textit{PvSPX1} lies downstream of \textit{PvPHR1}.

![Fig. 5. Growth and P concentration of bean hairy roots at two P levels.](http://jxb.oxfordjournals.org/)

(A) Photograph of bean hairy roots grown at two P levels. Scale bars=1 cm. (B) Fresh weight of bean hairy roots at two P levels. (C) P concentration in bean hairy roots. Each bar is the mean of four replicates with the standard error. Asterisks represent significant differences between overexpressing \textit{PvSPX} and CK for the same trait in t-tests. *0.01<P≤0.05; **P≤0.01. OX1 and OX2 indicate two transgenic bean hairy root lines overexpressing \textit{PvSPX1}, \textit{PvSPX2}, or \textit{PvSPX3}. CK1 and CK2 indicate the two transgenic lines transformed with the empty vector.)
Proteins containing the SPX domain have been well documented to be involved in the P signalling pathway of yeast and model plants, including *Arabidopsis* and rice (Lenburg and O’Shea, 1996; Duan et al., 2008; C. Wang et al., 2009; Z. Wang et al., 2009; Lin et al., 2010; Secco et al., 2012a). However, involvement of SPX proteins in P signalling remains largely unknown in legumes. In this study, three *PvSPX* genes were cloned and comparatively characterized as related to Pi starvation in bean. The results demonstrated that *PvSPX1* is an important regulator in the P signalling network of common bean, which shows several novel functions in regulating root growth, P homeostasis, and downstream gene transcription.

Since the transcription of several Pi starvation-responsive genes was noticeably increased and decreased in the *PvSPX1* overexpression and RNAi transgenic bean hairy roots, respectively, *PvSPX1* appears to be a positive regulator in the bean P signalling network (Fig. 7; Supplementary Fig. S7 at JXB online). Furthermore, *PvSPX1* appears to be a downstream gene of *PvPHR1*, because overexpressing *PvPHR1* led to increased transcription of *PvSPX1* in bean hairy roots (Fig. 8). Similarly, it has been demonstrated that *AtSPX1* and *OsSPX1* were downstream genes of *AtPHR1* and *OsPHR2* in the P signalling pathways of *Arabidopsis* and rice, respectively (Duan et al., 2008; C. Wang et al., 2009; Liu et al., 2010). However, regulatory roles of *PvSPX1*, *AtSPX1*, and *OsSPX1* in the P signalling pathways seemed to vary among species despite them showing several similar properties, such as nuclear localization and Pi starvation-induced expression patterns (Figs 2–4). In rice, *OsSPX1* has been considered as a negative regulator in the P signalling network because overexpressing *OsSPX1* significantly suppressed the expression levels of 10 Pi starvation-induced genes (C. Wang et al., 2009). Also, *OsSPX1* suppression resulted in increased transcripts of *OsPT2* and *OsPT8* in rice (C. Wang et al., 2009; Liu et al., 2010). However, *AtSPX1* was considered as a positive regulator in the *Arabidopsis* P signalling pathway because overexpressing *AtSPX1* led to increased transcription of several genes increased by Pi starvation, such as *AtACP5* and *AtRNS1* (Duan et al., 2008). Therefore, it seems that the regulatory roles of *SPX1* in dicots might differ from those in monocot plants, which needs to be further studied.

Consistent with the enhanced expression levels of two Pi transporter genes (*PvPHT2* and *PvPT1*), a significantly increased P concentration was observed in bean hairy roots overexpressing *PvSPX1*, especially under high P conditions (Fig. 5C). This suggests that *PvSPX1* is involved in regulating P homeostasis in bean roots. Similarly, it has been documented that suppressed expression of *OsSPX1* led to more P accumulation in both leaves and roots in rice under high P conditions (C. Wang et al., 2009; Liu et al., 2010). Taken together,
Fig. 7. Transcription levels of downstream genes of \( PvSPX1 \) in CK and \( PvSPX1 \)-overexpressing bean hairy roots. Expression patterns of downstream genes were determined by qPCR in CK and two \( PvSPX1 \) overexpression hairy root lines grown in MS medium containing 1.25 mM P. Expression levels of the tested genes were quantified relative to expression levels of the reference gene \( EF-1 \alpha (PvTC3216) \) using arbitrary units. OX1 and OX2 indicate two transgenic bean hairy root lines overexpressing \( PvSPX1 \). CK indicates the transgenic line transformed with the empty vector. Asterisks represent significant differences of downstream gene expression levels between \( PvSPX1 \)-overexpressing and CK in \( t \)-tests: * \( 0.01 < P \leq 0.05 \); ** \( 0.001 < P \leq 0.01 \); *** \( P \leq 0.001 \).
these results suggest that SPX might control P homeostasis in plants through regulating expression of Pi transporter (PT) genes. However, the molecular mechanisms underlying SPX regulation of PT transcription remain largely unknown. Since PvSPX1 has regulatory roles which appear to contrast with those of OsSPX1 and AtSPX3 in P signalling pathways, it is plausible that SPX might not directly control downstream gene expression. It will be important to clarify the functions of SPX through identification of other P signalling regulators interacting with SPX in plants.

Another novel feature of PvSPX1 is its involvement in regulating root growth and root morphology in bean roots. Changes in root morphology, such as inhibition of root elongation and stimulation of root hair growth, are well accepted as typical responses of plant roots to Pi starvation (Péret et al., 2011). It was found here that overexpressing PvSPX1 significantly inhibited hairy root growth at two P levels (Fig. 5B), but led to enlarged root hair zones (Fig. 6). This suggests that overexpression of PvSPX1 could enhance root morphological modifications in adaptation to P deficiency.

In bean, it has been documented that the P-efficient genotype G19833 has greater root hair density and longer root hair length than the P-inefficient genotype DOR364 in low P conditions (Yan et al., 2004). In this study, the Pi starvation-induced PvSPX1 was originally cloned from G19833. Furthermore, higher PvSPX1 expression levels were found in G19833 than in DOR364 (data not shown) at low P, suggesting positive contributions of PvSPX1 to superior P efficiency in G19833 through regulation of root morphology. Subsequently, two genes regulating root growth in transgenic bean hairy roots, PVPAP3-like and PVPDR2-like, were cloned and their transcription was investigated. Overexpression of PvSPX1 led to increased expression of PVPAP3-like and reduced expression of PVPDR2-like (Fig. 7). Since it has been documented that AtLPR1 and AtPDR2 are two critical components regulating root growth in opposite ways (Ticconi et al., 2006; Svistoonoff et al., 2007; Wang et al., 2010; Miura et al., 2011), it is conceivable that changes in root morphology in bean result from up-regulation of PvSPX1, with consequent effects on transcripts of PVPAP3-like and PVPDR2-like.

Although the three PspX proteins studied here exhibit high homology, and belong to the same subgroup in phylogenetic tree analysis (Fig. 1), diverse properties and functions of PspXs were observed in response to Pi starvation, as reflected by different expression patterns, variations in subcellular localization, and dissimilar growth of transgenic bean hairy roots. In response to Pi starvation, it seems that PvSPX1 and PvSPX2 might be earlier responsive genes which are more sensitive to Pi starvation than PvSPX3 in bean leaves. At 4 d of Pi starvation, transcripts of both
Functional characterization of PvSPX1

Table S2. Effects of phosphorus availability on bean growth.

<table>
<thead>
<tr>
<th>Phosphorus Availability</th>
<th>Root Length (cm)</th>
<th>Root Fresh Weight (g)</th>
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<tbody>
<tr>
<td>Control</td>
<td>5.2</td>
<td>0.8</td>
</tr>
<tr>
<td>Low Pi</td>
<td>3.8</td>
<td>0.6</td>
</tr>
<tr>
<td>High Pi</td>
<td>6.4</td>
<td>1.2</td>
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References


Supplementary data

Supplementary data are available at JXB online.

Figure S1. Expression of PvSPX genes in transgenic bean hairy roots.

Figure S2. Expression of PvSPX1 in PvSPX1 RNAi transgenic bean hairy roots.

Figure S3. Growth and P concentration of bean hairy roots in CK and PvSPX1 RNAi transgenic lines at two P levels.

Figure S4. Percentage of root hair zone in bean hairy roots with suppressed PvSPX1 at two P levels.

Figure S5. Transcription levels of downstream genes of PvSPX1 in CK and PvSPX1 RNAi transgenic lines.

Figure S6. Transcription levels of downstream genes of PvSPX1 in CK and overexpression transgenic lines of PvSPX2 or PvSPX3.

Figure S7. Expression patterns of PvSPX1 and PvSPX2 through their promoter::GUS analysis.

Table S1. List of primers used in the study.


