Short title: Medicago PHO1 members transport Pi to bacteroids

PHO1 family members transport phosphate from infected nodule cells to bacteroids in *Medicago truncatula*

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One-sentence summary
Two members of the PHO1 family in *Medicago truncatula* are involved in the transport of phosphate from the infected nodule cells to the *Sinorhizobium meliloti* bacteroids.

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NNTN, AJV, and YP conceived the project. JC and DJV contributed to plant transformation and analysis of gene expression; PVV and DdB performed confocal and electron microscopy, respectively; MJ and ML helped with measurement of nitrogen fixation; and MU contributed essential mutant plant lines. NNTN performed all other experiments. NNTN and YP wrote the manuscript, and ML, AJV, JC and MU read and provided essential manuscript feedback. All authors read and approved the final manuscript. YP agrees to serve as the author responsible for contact and ensures communication.
Abstract

Legumes play an important role in soil nitrogen availability via symbiotic nitrogen fixation (SNF). Phosphate deficiency severely impacts SNF because of the high phosphate requirement of symbiosis. Whereas PHT1 transporters are involved in phosphate uptake into nodules, it is unknown how phosphate is transferred from the plant infected cells to nitrogen-fixing bacteroids. We hypothesized that Medicago truncatula genes homologous to Arabidopsis PHO1, encoding a vascular apoplastic phosphate exporter, are involved in phosphate transfer to bacteroids. Among the seven MtPHO1 genes present in M. truncatula, we found that two genes, namely MtPHO1.1 and MtPHO1.2, were broadly expressed across the various nodule zones in addition to the root vascular system. Expression of MtPHO1.1 and MtPHO1.2 in Nicotiana benthamiana mediated specific phosphate export. Plants with nodule-specific down-regulation of both MtPHO1.1 and MtPHO1.2 were generated by RNA interference to examine their roles in nodule phosphate homeostasis. Nodules of RNAi plants had lower phosphate content and a 3-fold reduction in SNF, resulting in reduced shoot growth. Whereas the rate of $^{33}$P phosphate uptake into nodules of RNAi plants was similar to control, transfer of $^{33}$P phosphate from nodule cells into bacteroids was reduced and bacteroids activated their phosphate-deficiency response. Our results implicate plant MtPHO1 genes in bacteroid phosphate homeostasis and SNF via the transfer of phosphate from nodule infected cells to bacteroids.
Introduction

Biological fixation of nitrogen by rhizobia is of prime importance in the global nitrogen cycle and a major provider of fixed nitrogen in agricultural soils (Herridge et al., 2008). The rhizobia–legume interaction occurs within the nodule, a specialized root structure, whereby rhizobia fixes atmospheric nitrogen in exchange for organic acids and other nutrients provided by the plant (Udvardi and Poole, 2013). Rhizobia enters the root via an infection thread initiated within the root hairs that eventually reaches cortical cells that divide and multiply in response to Nod factors released by the bacteria (Oldroyd et al., 2011; Mergaert et al., 2020; Roy et al., 2020). Bacteria released from the infection thread enter the cortical cells via a process generating an organelle-like structure, the symbiosome, whereby the bacteria are surrounded by a membrane derived from the plant plasma membrane (PM). This membrane, named the peribacteroid or symbiosome membrane (SM), separate the bacteria, which ultimately differentiate into nitrogen-fixing bacteroids, from the plant cytosol (Whitehead and Day, 1997). Proteomic analysis of SM identified a range of transporters, pumps, and channels that are thought to contribute to the movement of nutrients and ions between the plant cells and the bacteroids (Saalbach et al., 2002; Wienkoop and Saalbach, 2003; Clarke et al., 2015). Despite this, our current knowledge of the proteins involved in the movement of specific nutrients and ions from the cytosol of infected nodule cells to the bacteroids is very limited. For example, whereas malate is thought to be the main source of carbon provided by the plant to the bacteroid in exchange for ammonia, the malate transporter involved in this exchange is still unknown (Clarke et al., 2014; Poole et al., 2018). However, proteins involved in the transport of nitrate and citrate to the bacteroids have been described (Vincill et al., 2005; Kryvoruchko et al., 2018).

Beyond carbon and nitrogen, the plant cell must also provide bacteroids with numerous elements essential for proper metabolism, including calcium, sulfate (SO₄), potassium, and several metals such as iron, zinc, and copper. *Glycine max* ZIP1 and DMT1 were shown to be transporters for zinc and iron, respectively, when synthesized in yeast, and found to be expressed in nodules and localized to the SM (Moreau et al., 2002; Kaiser et al., 2003). Similarly, a protein from *Lotus japonicus* that can mediate SO₄ transport when expressed in yeast was shown to be essential for SNF (Krusell et al., 2005).

Phosphorus (P) is one of the most limiting nutrients for plant growth, especially in legumes that can access atmospheric nitrogen via SNF. P level in soybean nodules are three-fold
higher than in roots and leaves (Gaude et al., 2004a) and legumes require higher levels of P for growth supported by SNF compared to growth on nitrate (Israel, 1987; Gunawardena et al., 1992; Israel, 1993). Nodules are highly active metabolically, with protein concentrations more than an order of magnitude higher than in roots (Gaude et al., 2004a). Comparison of phosphorylated metabolites in free living *Bradyrhizobium japonicum* and bacteroids isolated from soybean nodules showed a large increase in P-containing metabolites in bacteroids, ranging from free inorganic phosphate (Pi) to ATP and numerous phosphorylated intermediates (Vauclare et al., 2013). Pi deficiency can affect SNF either indirectly, e.g. by reducing photosynthesis and carbon supply from the plant to the bacteroids, or directly by constraining nodule development and metabolic activity through its impact on Nod factor production and nitrogenase activity (McKay and Djordjevic, 1993; Hernandez et al., 2009).

Adaptation of legumes to Pi deficiency shares many similarities to that of non-leguminous plants, including changes in root development, secretion of organic acids and phosphatases into the rhizosphere, and the activation of pathways that bypass the need for ATP and some P-containing compounds (O'Rourke et al., 2013; Sulieman and Tran, 2015). The main Pi-deficiency signaling components identified in Arabidopsis, including the transcription factor PHR1, the small RNA mi399, and SPX signaling components, are also found in legumes (Xu et al., 2013; Zhang et al., 2016; Xue et al., 2017). Nevertheless, it is likely that adaptation of nodules to Pi deficiency allowing the maintenance of nitrogen fixation involves nodule-specific mechanisms (Valentine et al., 2017).

Plants acquire P from their environment via Pi transporters coupled to a proton gradient generated by H⁺-ATPases associated with the PM. The principal transporters of Pi into plant cells belong to the PHT1 family (Nussaume et al., 2011; Poirier and Jung, 2015). Two pathways for Pi entry into nodules have been described, namely a direct uptake by the nodule and an indirect pathway involving Pi transfer from roots of the host plant to the nodule (Al-Niemi et al., 1998). Expression of two *PHT1* genes in soybean nodules was shown to be essential for nodulation and nitrogen fixation, highlighting their implication in the uptake of Pi into nodules (Qin et al., 2012; Chen et al., 2019). Whereas entry of Pi into bacteroids is mediated by low-affinity Pit- and high-affinity Pst bacterial Pi transporters (Bardin et al., 1998; Botero et al., 2000; Yuan et al., 2006), plant transporters required to transfer Pi from the host cell cytoplasm to the bacteroids are unknown.
The *Arabidopsis thaliana* and rice (*Oryza sativa*) *AtPHO1* and *OsPHO1.2* genes are involved in loading of Pi into the xylem vessel for transfer to the shoot (Hamburger et al., 2002; Secco et al., 2010). *AtPHO1* is also expressed in the chalazal seed coat where it is involved in the transfer of Pi from the maternal seed coat to the filial developing embryo (Vogiatzaki et al., 2017). *AtPHO1* expressed in Arabidopsis or *Nicotiana benthamiana* mediates Pi export across the plasma membrane to the apoplastic space (Arpat et al., 2012; Wege et al., 2016). Given the role of *AtPHO1* in Pi transfer between tissues, we hypothesized that homologues of *PHO1* in legumes could be involved in Pi homeostasis in the nodule, and more specifically in Pi transfer to the bacteroid. In this work, we characterize the *PHO1* gene family in *Medicago truncatula* and report the role of two *MtPHO1* transporters in the transfer of Pi from infected plant cells to the bacteroids.
Results

Identification of MtPHO1 genes expressed in nodules

Seven genes encoding proteins with high similarity to AtPHO1, including conserved SPX and EXS domains, were identified using the TBLASTN function of the *M. truncatula* Genome Database (http://blast.jcvi.org/Medicago-Blast/index.cgi; Fig. S1). Phylogenetic analysis grouped the MtPHO1 homologues into two distinct clades, in agreement with previous analysis of the PHO1 gene family in monocots and dicots (Secco et al., 2010). Clade 1 contains three *M. truncatula* PHO1 homologues, named MtPHO1.1–1.3, whereas 4 genes named MtPHO1.4–1.7 belong to clade 2 (Fig. 1a).

The expression profiles of MtPHO1 genes was assessed using the *M. truncatula* Gene Expression Atlas (mtgea.noble.org/v3/) compiling data obtained using a dedicated Affymetrix Medicago Gene Chip (affymetrix.com) that contained probes for 6 MtPHO1 genes, namely MtPHO1.1–1.6. The three MtPHO1 genes belonging to clade 1, namely MtPHO1.1, MtPHO1.2, and MtPHO1.3, were broadly expressed in the shoot, stem, flower, seed, root, and nodule, with MtPHO1.1 and MtPHO1.2 being the genes expressed most robustly in nodules (Fig. 1b). By contrast, MtPHO1.4–1.6 genes were not expressed, or only very weakly, in various tissues, including nodules. RT-PCR analysis of nodules from *M. truncatula* inoculated with *S. meliloti* and grown under nitrogen fixing conditions confirmed the expression of MtPHO1.1, MtPHO1.2, and MtPHO1.3, whereas expression of all other genes, including MtPHO1.7, was either very weak or undetectable (Fig. S1b). The genomic structure of the MtPHO1.1 and MtPHO1.2 loci as well as MtPHO1.1 and MtPHO1.2 amino-acid alignment with the *A. thaliana* PHO1 are shown in Supplementary Figure S1.

The spatial expression patterns of the three main MtPHO1 genes expressed in nodules was investigated further by gene-GUS fusion analysis (Fig 1c). All constructs included a 2 kb promoter region and the cDNA sequence of MtPHO1.1, MtPHO1.2 or MtPHO1.3 fused at the C-terminus to the *uidA* gene, and histochemical analysis for GUS activity was performed on roots and nodules from composite transgenic plants grown under nitrogen fixing conditions. All three genes were expressed in the vascular cylinder of roots (Fig. 1c, upper panels). Whereas GUS staining for the pMtPHO1.3:cMtPHO1.3-GUS construct was detected only in the vascular tissue of nodules, GUS expression from the pMtPHO1.1:cMtPHO1.1-GUS and pMtPHO1.2:cMtPHO1.2-GUS constructs was not limited to the vascular tissue but observed...
across the whole nodule (Fig. 1c, lower panels). Since transporters involved in Pi transfer from the infected cortical cells to the bacteroids are expected to be broadly expressed in cells of the nodule and not be confined to the vascular cylinder, further analysis was restricted to the MtPHO1.1 and MtPHO1.2 genes. Analysis of the expression patterns of these two genes using data from the Symbimics database (https://iant.toulouse.inra.fr/symbimics), derived from laser-capture microdissection of nodules (Roux et al., 2014), confirmed that both genes were broadly expressed in the nodules, from the meristematic zone to the nitrogen-fixing zone (Fig. S2).

**MtPHO1.1 and MtPHO1.2 are Pi exporters**

Pi transport activity of the MtPHO1.1 and MtPHO1.2 proteins fused to GFP was tested by transient expression in *N. benthamiana* leaves. Arabidopsis PHO1-GFP and free GFP were expressed as positive and negative controls, respectively. Transient expression of MtPHO1.1-GFP and MtPHO1.2-GFP fusions led to greater export of Pi into the extracellular medium than either free GFP or non-infiltrated controls, and to a similar extent to the expression of the Arabidopsis PHO1-GFP (Fig. 2a). By contrast, nitrate export was unaffected by expression of any of these transporters, indicating specificity for Pi export of MtPHO1.1 and MtPHO1.2.

Expression of MtPHO1.1-GFP and MtPHO1.2-GFP fusion proteins in *M. truncatula* hairy...
root transformants using the gene’s endogenous promoters did not show reliable GFP expression above the autofluorescence background in either nodules or root vascular cylinder. Subcellular localization of MtPHO1.1-GFP and MtPHO1.2-GFP was thus determined by transient expression in *N. benthamiana*. MtPHO1.1-GFP or MtPHO1.2-GFP were co-expressed with either the Golgi marker ManI-RFP (Langhans et al., 2011) or the plasma membrane marker CBL1-OFP (Batistič et al., 2010). AtPHO1-GFP has previously been shown to colocalize with the Golgi marker (Arpat et al., 2012), and a similar co-localization occurred with the MtPHO1.2-GFP construct. By contrast, MtPHO1.1-GFP co-localized primarily with the plasma membrane marker and not with the Golgi marker (Fig. 2b).

**MtPHO1.1 and MtPHO1.2 contribute to nitrogen fixation**

In order to identify roles of *MtPHO1.1* and *MtPHO1.2* in nodulation and nitrogen fixation, *M. truncatula* mutants for each gene were obtained from the *Transposable Element from N. tabacum (Tnt1)* collection of the Noble Research Institute (Tadege et al., 2008). Two homozygous mutant lines for each gene were identified and evaluated. Mutants *mtpHO1.1a*
(NF14242) and mtpho1.1b (NF14056) harbor Tnt1 insertions in the second and seventh introns of the MtPHO1.1 gene, respectively, leading to an approximately 50–60% reduction in gene expression levels (Fig. 3 a, c). The two independent mutants mtpho1.2a (NF1102) and mtpho1.2b (NF10573) contain Tnt1 insertions in the first and second exons, respectively, and resulted in undetectable levels of gene expression (Fig. 3 a, c). Growth of all four Tnt1 mutants under nitrogen fixing conditions were undistinguishable from either the parental R108 or WT segregants obtained from the heterozygous mutants, with no significant change in either shoot fresh weight or Pi content of shoot or nodule, nodule number or nodule size (Fig. 3 b, d-h).

Because of the potential functional redundancy between MtPHO1.1 and MtPHO1.2, we aimed to create transgenic lines where both genes are down-regulated. Since both MtPHO1.1 and MtPHO1.2 are not only expressed in nodules but also various other tissues, including in the vascular cylinder of roots, we chose to down-regulate the expression of both genes specifically in nodules with an RNAi approach using the nodule-specific promoter of the NCR001 gene (Mergaert et al., 2003). Three distinct hairpin RNAi constructs were designed using sequences outside the PHO1 SPX and EXS domains, corresponding to regions of highest divergence between members of the MtPHO1 family (Fig. S3a). Each RNAi construct was a hybrid containing approximately 500 bp of coding sequence of both MtPHO1.1 and MtPHO1.2 genes, and thus designed to down-regulate both genes simultaneously (Fig S4). Composite plants with transgenic roots expressing the RNAi construct or transformed with an
empty vector control (EV) were inoculated with *S. meliloti* and grown in a clay-based substrate in the presence of either full nutrient containing a source of nitrogen (NO$_3^-$ and NH$_4$NO$_3$) or full nutrient without mineral nitrogen. Several independent composite transgenic plants transformed with each of the three RNAi constructs gave plants that grew poorly under mineral nitrogen-free conditions compared to EV controls (Fig. S3b). These plants exhibited shoot Pi content comparable to EV controls but had lower nodule Pi contents and nodules had significantly higher fresh weights than those of controls (Fig. S3c-e). Because the phenotypes observed with the three RNAi constructs were similar, further detailed characterization of the RNAi phenotype was limited to composite plants transformed with the RNAi3 construct (Figure 4).

Reverse transcription quantitative PCR (RT-qPCR) analysis of transgenic roots transformed with RNAi3 showed an approximate 55% reduction in *MtPHO1.1* and *MtPHO1.2* expression; however, this was confined to nodules and gene expression in root sections without nodules was unchanged relative to the EV control (Fig. 4a). The reduction in shoot fresh weight of the RNAi-transformed roots was only observed in plants grown under SNF conditions and not when plants were grown in the presence of a fixed nitrogen source in the nutrient solution (Fig. 4b-c). Under both growth conditions, shoot Pi content remained similar between RNAi and EV (Fig. 4d). Although nodules from RNAi roots showed an increase in the fresh weight.
as compared to EV and contained less Pi per mass, the number of nodules per plant remained unchanged (Fig 4d-g).

Nitrogen fixation in isolated nodules was measured using $^{15}\text{N}_2$. Plants with roots transformed with the RNAi3 construct showed a 3.5-fold reduction in the specific nitrogen fixation activity ($\text{N}_2$ fixation per nodule biomass) relative to the EV control (Fig. 4h). Although this reduction was partially compensated by an increase of nodule biomass in the RNAi plants, the reduced expression of MtPHO1.1 and MtPHO1.2 specifically in nodules decreased the nitrogen fixation potential of the plant, resulting in reduced growth of plants in the absence of mineral nitrogen.

Nodules from roots transformed with the EV control or RNAi3 construct roots appeared similar in gross nodule structure (Fig. 5a). However, bacteroids in the nitrogen fixation zone (zone 3) showed a significant reduction in cell length in nodules derived from RNAi roots compared to EV controls, as measured by electron microscopy (Fig. 5b, c).

**Reduction of MtPHO1.1 and MtPHO1.2 expression results in Pi-deficient bacteria**

The dynamics of Pi and SO$_4$ entry into nodules and its transfer to bacteroids were measured using radio-isotopes. Excised nodules from roots transformed with the RNAi3 construct imported similar amounts of either $^{33}\text{Pi}$ or $^{35}\text{SO}_4$ from the bathing solution compared to nodules from EV controls (Fig. 6a, b). However, there was a 2-fold reduction in the amount of $^{33}\text{Pi}$ acquired by bacteroids in the RNAi lines compared to EV controls, whereas the amount of $^{35}\text{SO}_4$ acquired by the bacteroids was not different between RNAi and EV-control root nodules (Fig. 6c). These data indicated a deficiency in Pi transfer from plants cells to the bacteroids.

To test whether the bacteria present in nodules on roots transformed with the RNAi3 construct experienced Pi deficiency, we inoculated plants with the *S. meliloti* RCR2011 strain containing *pstS::gusA* or *orfA-pit::gusA* reporter genes. *PstS* encodes a high-affinity Pi transporter and its promoter is activated under Pi deficiency, whereas *Pit* encodes a low-affinity Pi transporter and is repressed under Pi deficiency (Yuan et al., 2006). Compared to the EV control, nodules from roots transformed with the RNAi3 construct containing *S. meliloti* with *orfA-pit::gusA* showed a decrease in GUS expression throughout the nodule whereas an increase in GUS expression was associated with *S. meliloti* expressing *pstS::gusA*.
in RNAi3-expressing nodules compared to the control (Fig. 6d). Altogether, these experiments showed that down-regulation of \textit{MtPHO1.1} and \textit{MtPHO1.2} expression was accompanied by a reduction of Pi transfer from infected nodule cells to bacteroids, resulting in Pi-deficient bacteroids and a reduction in nitrogen fixation.

\textbf{Fig. 5} Bacteroid size and distribution in nodules. A, Panoramic reconstruction of 3X3 and 5X5 pictures showing a 200 \textmu m thick longitudinal section of nodules from empty vector (EV) control and RNAi plants grown for 28 days under symbiotic nitrogen fixation (SNF) condition. Cells heavily colonized by bacteroids are shown as electron-dense areas. Bar = 250 \textmu m. B, Electron micrographs of cells colonized with differentiated elongated bacteroids in the nitrogen fixation zone of nodules from EV control and RNAi plants grown for 28 days under SNF condition. Bar = 10 \textmu m. C, The distribution of bacteroid size measured in electron micrographs of infected cells in the nitrogen fixing zone. For both nodules from EV control and RNAi plants, the size of 1000 bacteroids were measured from 30-40 distinct cells.
Fig. 6 Flux of Pi and SO$_4$ into nodules and bacteroids. Isolated nodules from empty vector (EV) control plants and RNAi plants grown in clay-based substrate under symbiotic nitrogen fixation (SNF) conditions were incubated in a nutrient solution containing either $^{32}$Pi (A) or $^{35}$SO$_4$(B) and the amount of $^{32}$Pi and $^{35}$SO$_4$ absorbed by the nodule was measured. Error bars represent SD (n=4). C. The amount of $^{32}$Pi and $^{35}$SO$_4$ acquired by bacteroids in the RNAi line was normalized to the number of bacteroids and expressed relative to the EV control. Error bars represent SD (n=4). Asterisks represent statistically significant differences for RNAi compared to the EV control (t-test, p < 0.05). D. Cross-section of nodules stained for β-glucuronidase activity taken from RNAi and EV plants infected with S. meliloti strain Sm2011 harboring the promoter fusion orfA::gusA or ppsS::gusA and grown under SNF conditions. Bar = 500 μm.
Discussion

Phylogenetics analysis of the PHO1 gene family in angiosperms showed the presence of two major clades (Secco et al., 2010; He et al., 2013). Genes in Clade 1 are found in both monocots and dicots and include genes with highest homology to AtPHO1 and AtPHO1;H1, the only two genes in A. thaliana shown to mediate Pi export to the root xylem vessels for long-distance Pi transport (Stefanovic et al., 2007). PHO1-like genes belonging to Clade 1 in rice and tomato (Solanum lycopersicum) are also involved in the long-distance transfer of Pi from roots to shoots (Secco et al., 2010; Zhao et al., 2019). By contrast, genes in Clade 2 are present only in dicots, and none have been shown to be directly involved in long-distance Pi transport (Stefanovic et al., 2007). The only functionally characterized members of this clade are AtPHO1;H4 (SHB1), implicated in hypocotyl elongation in blue light (Kang and Ni, 2006); and AtPHO1;H3, thought to negatively modulate AtPHO1 activity under zinc deficiency (Khan et al., 2014). MtPHO1.1, MtPHO1.2, and MtPHO1.3 all belong to Clade 1 and are the only M. truncatula PHO1 genes with robust expression in a range of tissues, including nodules. Similar to the Pi exporters AtPHO1 and AtPHO1;H1 in Arabidopsis and OsPHO1.2 in rice (Hamburger et al., 2002; Stefanovic et al., 2007; Jabnoune et al., 2013), these three M. truncatula genes are expressed in the root vascular cylinder. These data, combined with the Pi export activity of MtPHO1.1 and MtPHO1.2 expressed in N. benthamiana, suggest that these genes are also likely involved in long-distance Pi transfer via Pi export into the xylem. Since the null mtpho1.2 mutants showed no decrease in shoot Pi content, the typical phenotype of null pho1 mutants in A. thaliana, rice, and tomato (Poirier et al., 1991; Secco et al., 2010; Zhao et al., 2019), it is likely that other members of the MtPHO1 gene family expressed in the root vascular cylinder contribute to this activity, including MtPHO1.1 and MtPHO1.3. Analysis of higher-order knock-out mutants for these genes should help clarify their respective contribution to long-distance, root-to-shoot Pi transport.

The concomitant down-regulation of MtPHO1.1 and MtPHO1.2 in nodules by RNAi led to a ~3.5-fold reduction in nitrogen fixation in nodules. This significantly impacted growth of plants relying on nitrogen fixation as the primary source of usable N. Interestingly, neither the knocked-down or null Tntl mutants in MtPHO1.1 or MtPHO1.2, respectively, showed growth phenotypes or perturbations in Pi content in nodules in plants grown under SNF conditions. These results would suggest that these genes are likely to have redundant function in nodules. The similar expression profiles of these two genes in nodules and the ability of the respective
proteins to mediate Pi export when expressed in *N. benthamiana* supports this conclusion. However, the observation that the majority of MtPHO1.1 and MtPHO1.2 proteins are not localized in the same sub-cellular membrane when expressed in *N. benthamiana* raises the possibility that, whereas both proteins contribute to nodule Pi homeostasis and SNF, their mode of action is not completely redundant (see further discussion on MtPHO1.1 and MtPHO1.2 localization below).

Whereas shoot Pi content was not affected in RNAi plants compared to EV controls, irrespective of the growth conditions, nodule Pi content was decreased by ~2-fold in RNAi plants. Experiments with $^{33}$Pi showed that the transfer of Pi acquired by nodules into the bacteroids was significantly decreased, despite that direct Pi uptake from the bathing medium into nodules was not affected in the RNAi plants. These data support the hypothesis that *MtPHO1.1* and *MtPHO1.2* expression in nodules directly contributes to Pi transfer from the plant cells to the bacteroids. No perturbation in the transfer of SO$_4$ from the nodule cells to the bacteroids was observed, in agreement with the specificity of PHO1 for Pi export (Arpat et al., 2012; Wege et al., 2016). Interestingly, RNAi plants showed a 1.5-fold increase in nodule weight although the number of nodules per plant remained the same. The increase in nodule weight in RNAi plants could potentially be a consequence of the reduction of SNF.

Previous studies have shown that under nitrogen deficit, symbiotic plants can stimulate nodule expansion via a systemic signal as a compensatory mechanism aimed at increasing nitrogen acquisition capacities (Jeudy et al., 2010; Laguerre et al., 2012).

Pi transport in *S. meliloti* is mediated by both low-affinity Pit- and high-affinity Pst transporters. As in *E. coli*, the Pit-encoding gene OrfA is strongly expressed under Pi-sufficient conditions but repressed under Pi-deficient conditions. Conversely, the PstSCAB operon involved in high-affinity Pi transport is repressed under Pi-sufficient conditions but induced under Pi starvation (Bardin et al., 1998; Yuan et al., 2006; diCenzo et al., 2017). Relative expression of the Pit- and Pst-encoding operons in bacteroids has previously been used to deduce the level of Pi availability in indeterminate nodules, e.g. in *M. truncatula*, and determinate nodules, e.g. soybean (Bardin et al., 1998; Yuan et al., 2006; diCenzo et al., 2017; Hu et al., 2018). In the present study, bacteroids containing the OrfA::gusA reporter showed a reduction in expression in nodules of the *PHO1* RNAi plants compared to the control, whereas the reverse was observed for bacteroids containing the Pst::gusA reporter.
These data show that reduction of MtPHO1.1 and MtPHO1.2 expression in nodules leads to an activation of the Pi-starvation response in bacteroids, consistent with the lower level of Pi transfer from the nodule plant cells to bacteroids. Nodules are metabolically highly active and bacteroids fixing nitrogen contain high amount of Pi, ATP, and other phosphorylated intermediates (Gaude et al., 2004b; Vauclare et al., 2013). Rhizobium mutants affected in the expression of Pi importers show defect in nodulation and SNF, highlighting the importance of an adequate supply of Pi to bacteroids for optimal SNF (Bardin et al., 1996; Jiao et al., 2016; Hu et al., 2018). Taken together, these data indicate that the reduction of Pi supply from the plant nodule to the bacteroids is likely to be the primary effect of the down-regulation of MtPHO1.1 and MtPHO1.2 expression, which in turn negatively impacts nitrogen fixation and nodulation. It is unknown whether Pi-deficiency in bacteroids causes a reduction in bacteroid size or whether other more indirect effect are involved, such as changes in the expression of cysteine-rich peptides (NCRs) known to affect bacteroid differentiation and size (Horvath et al., 2015).

A. thaliana PHO1 is primarily localized to the Golgi and trans-Golgi network (TGN) either when expressed under control of its native promoter in the root vascular cylinder or under control of the CaMV35S promoter in N. benthamiana leaves, and this expression pattern is concomitant with its Pi export activity to the apoplast (Arpat et al., 2012; Liu et al., 2012). The absence of detectable amounts of AtPHO1 at the PM could be part of a post-translational retrieval mechanism to control its activity. This would be analogous to IRT1, an iron transporter that acts at the PM but partitions primarily to the endomembrane system, which is part of a mechanism to control iron import through constitutive IRT1 endocytosis regulated via monoubiquitin- and clathrin-dependent mechanisms (Barberon et al., 2011). We have been unable to detect expression of MtPHO1.1-GFP or MtPHO1.2-GFP in nodules, likely due to the combination of weak expression and high level of tissue autofluorescence. However, functional expression in N. benthamiana showed that Pi export activity was associated with PM and Golgi localization of MtPHO1.1-GFP and MtPHO1.2-GFP, respectively. Although it is unknown whether MtPHO1.1 and MtPHO1.2 have distinct subcellular localizations within nodules, these results in N. benthamiana show that Pi export mediated by MtPHO1.1 and MtPHO1.2 is not constrained by their predominant steady-state sub-cellular localization either at the PM or Golgi. These results imply that the distinct subcellular localization of MtPHO1.1 and MtPHO1.2 at steady-state levels may not necessarily reflect distinct modes of action as Pi exporters, but perhaps distinct modes of regulation. For example, whereas both Pi exporters
would transit via the Golgi-TGN-exocytic vesicle pathway to reach their final destination, MtPHO1.2 would be more partitioned, at steady-state level, in the Golgi-TGN (similar to IRT1), while MtPHO1.1 would be more partitioned to the PM (or SM in nodules).

Transfer of Pi from the cytoplasm of infected plant nodule cells to the bacteroids must ultimately occur across the SM, which acts as the gate keeper to the bacteroids (Roy et al., 2020). A proteomic study aimed at identifying SM proteins in soybean reported the presence of the Pi transporters GmPT5 (Glyma10g04230) and GmPT7 (Glyma10g33030), both belonging to the PHT1 family (Clarke et al., 2015). However, further studies of these proteins found them expressed either in the nodule vascular bundle for GmPT5 (Qin et al., 2012), or the plasma membrane of cells of the nodule outer cortex and fixation zones for GmPT7 (Chen et al., 2019), making them unlikely candidates for SM Pi transporters. Although the present work demonstrates the implication of MtPHO1.1 and MtPHO1.2 in the transfer of Pi transfer from the nodule cells to the bacteroids, further work is necessary to elucidate more precisely their mode of action. In particular, it will be interesting to determine if MtPHO1.1 and/or MtPHO1.2 can be localized at the SM, either at steady-state level or transiently. It will also be interesting to determine whether the sub-cellular localization of MtPHO1.1 and MtPHO1.2 is distinct between infected cells of nodules and cells of the vascular cylinder.
Material and Methods

Identification of PHO1 Family Members in *Medicago truncatula*

*Medicago truncatula* PHO1 genes were identified by sequence similarity to the complete *Arabidopsis thaliana* PHO1 using the TBLASTN algorithm at the *Medicago truncatula* Genome Database (http://blast.jcvi.org/Medicago-Blast/index.cgi) and the Noble Foundation *Medicago truncatula* IMGAG V3.0 (https://gb.noble.org/cgi-bin/gb2/gbrowse/medicago). The gene nomenclature and associated annotation numbers are given in Figure S1. The phylogenetic tree was constructed through the MABL bioinformatics platform (https://www.phylogeny.fr) (Dereeper et al., 2008).

Plant material, bacterial strains, growth conditions, and hairy root transformation

*M. truncatula* plants were derived from the ecotypes A17 and R108 whereas the *Sinorhizobium meliloti* strain was Rm2011. The *M. medicago* Tnt1 mutant population (Tadege et al., 2008) was screened for insertions in the MtPHO1.1 and MtPHO1.2 genes by PCR as previously described (Cheng et al., 2011). Oligonucleotides sequences used as primers for PCR and genotyping are provided in Table S1.

Seeds were scarified for 10 min in H$_2$SO$_4$, rinsed 5 times with sterile water, followed by surface sterilization for 10 min with 33% (v/v) commercial bleach. Seeds were then rinsed 5 times with sterile water and left in the dark in 50-mL tubes for 1 h to imbibe. Surface-sterilized seeds were placed on inverted plates containing 0.7% (w/v) agarose in the dark for 3 d at 4°C and 1 d at 20°C. Germinated seedlings were transferred to agar-solidified Fahräeus medium (0.132 g/L CaCl$_2$, 0.12 g/L MgSO$_4$.7H$_2$O, 0.1 g/L KH$_2$PO$_4$, 0.075 g/L Na$_2$HPO$_4$.2H$_2$O, 5 mg/L Fe-citrate, and 0.07 mg/L each of MnCl$_2$.4H$_2$O, CuSO$_4$.5H$_2$O, ZnCl$_2$, H$_3$BO$_3$, and Na$_2$MoO$_4$.2H$_2$O, adjusted to pH 6.5 before autoclaving) and grown for 10 days for hairy root transformation using *Agrobacterium rhizogenes* ARqua1 essentially as previously described (Boisson-Dernier et al., 2001). Following hairy root transformation, seedlings were grown vertically on Fahräeus medium supplemented with 25 μg/mL kanamycin in a growth chamber at 21°C (16 h light/8 h dark cycles) for 14 d, before being transferred to pots in a clay-based substrate (Seramis; https://www.seramis.com/) with no additional fertilizer. Seven days after transfer to pots, plants were inoculated with *S. meliloti* grown in TY medium (Beringer, 1974) and resuspended in nitrogen-free, half-strength B&D
medium (Broughton and Dilworth, 1971) to an OD$_{600}$ of 0.02. Each plant was supplied with 50 mL of the inoculum at the base of the stem. Inoculation with *S. meliloti* was repeated after 10 days.

Non-inoculated control plants were grown in the clay-based substrate fertilized every 10 days with nitrogen-free half-strength B&D medium (Broughton and Dilworth, 1971) containing 3 mM of KNO$_3$ and 2.5 mM NH$_4$NH$_3$, whereas plants grown under nitrogen fixation conditions were fertilized with a similar nutrient solution but without any source of nitrogen. Plants were grown in a greenhouse with a 16 h, 25°C light/8 h, 20°C dark cycle.

**RNA preparation and RT-qPCR**

Plant tissues were frozen in liquid nitrogen, then ground using a mortar and pestle and the powder was used to extract the RNA using the RNeasy Plant Mini Kit (Qiagen). Samples were treated with DNase on columns during the RNA extraction using the RNase-free DNase kit (Qiagen). RNA samples were used for reverse transcription using SuperScript II reverse transcriptase (Invitrogen). Quantitative PCR (qPCR) was performed using SYBER Select Master Mix (Invitrogen) and the Stratagene Mx3005 qPCR system using gene-specific primer (Supplemental Table S2)

**Generation of PHO1-GUS fusion genes and GUS staining**

Plasmid pMDC163 (Curtis and Grossniklaus, 2003) was digested by restriction enzyme *XhoI* to excise the hygromycin-resistance cassette to replace it with the neomycin phosphotransferase cassette amplified from plasmid pMDC100 (Curtis and Grossniklaus, 2003) and inserted by In-Fusion (Takara Bio). The modified pMDC163 plasmid was named pMDC163K. A 2-kb DNA fragment upstream of the starting ATG of *MtPHO1.1* (*Medtr1g041695.1*), *MtPHO1.2* (*Medtr1g075640.1*), or *MtPHO1.3* (*Medtr8g069955.1*) was amplified by PCR using gene-specific primers (Supplemental Table S3) from genomic DNA of *M. truncatula* A17, whereas the coding sequence of genes was amplified by PCR using gene-specific primers (Supplemental Table S3) from cDNA derived from whole plants. The promoter and coding cDNA fragments were cloned into the pENTR2B (Invitrogen) by In-Fusion, checked by sequencing and then fused to the *GUS* reporter gene in the destination vector pMDC163K or pKGWFS7-RR (Karimi et al., 2002) by In-Fusion (Takara Bio). The various DNA constructs were introduced in plants via hairy root transformation using *A.*
rhizogenes ARqua1, as described above. Nodules were stained for GUS activity by infiltrating, under mild vacuum, a solution containing 2 mM 5-bromo-4-chloro-3-indolyl-β-glucoronic acid (X-Gluc), 10 mM EDTA, 2 mM potassium ferricyanide, 2 mM potassium ferrocyanide, 0.1% (v/v) Triton X-100, and 100 mM sodium phosphate, pH 7. The enzymatic reaction was performed at 37°C in the dark. Stained tissues were washed with a graded ethanol series (10–80% (v/v)) and cleared in chloral hydrate solution (2.7 g/mL in 30% (v/v) glycerol).

Expression of MtPHO1-GFP fusion in Nicotiana benthamiana, ion export and confocal analysis
The MtPHO1.1 and MtPHO1.2 genes were amplified from genomic DNA using gene specific primers (Table S3), cloned into the pENTR2B (Invitrogen) by In-Fusion polymerase (Takara Bio), and inserted into pMDC83 plasmid by Gateway cloning (Invitrogen). For transient expression, Nicotiana benthamiana plants were infiltrated with Agrobacterium tumefaciens as previously described (Arpat et al., 2012). Pi and nitrate export assays were performed as previously described (Arpat et al., 2012). Pi concentration in solution was quantified by the molybdate assay (Ames, 1966) whereas nitrate was quantified by first converting nitrate to nitrite using commercial nitrate reductase from Aspergillus niger (Sigma) followed by nitrite quantification using sulfanilamide (Barthes et al., 1995). Confocal imaging was performed using a Zeiss LSM 700 instrument with Apochromat 63 × NA1.2 water immersion or multi-immersion objective.

Generation of RNAi construct
RNAi construct were designed by fusing together PCR fragments from MtPHO1.1 and MtPHO1.2 by In-Fusion (Takara Bio) and cloning the product in pENTR2B. The chimeric fragments were inserted into pK7GWIWG5D(II) harboring the promoter of the MtNCR001 gene (Mergaert et al., 2003) using Gateway Technology (Invitrogen). Sequences of oligonucleotides used for this cloning are listed in Table S4.

Nitrogen fixation assay
The 15N2 fixation was measured essentially as previously described (Ruffel et al., 2008). Freshly excised nodulated roots were placed in air-tight, 10-mL tubes containing 2 mL of basal nutrient solution. Ten minutes labeling was achieved by replacing in each tube 5 mL of air by 5 mL of 80%15N/20%O2 mix (99 atom% 15N). Samples (100 μL) of 15N2-enriched air
were harvested at the beginning and at the end of the labeling, for precise analysis of the atom % $^{15}$N of the $^{15}$N$_2$ source and leak check. After labeling, nodules were separated from roots, both organs were dried and analyzed for $^{15}$N and total N contents using a continuous-flow isotope ratio mass spectrometer (Isoprime mass spectrometer, GV instruments) coupled to a nitrogen elemental analyzer (Euro vector S.P.A) at the stable isotope platform of B&PMP (https://www1.montpellier.inra.fr/wp-inra/bpmp/en/platform/stable-isotope-analytical-platform/). N$_2$ fixation capacity was expressed as the number of μmole of N atoms acquired per g of nodule dry weight and hours.

$^{33}$PO$_4$ and $^{35}$SO$_4$ flux measurement in nodules
Nodules from transgenic roots were harvested, weighed, and placed in a solution containing 10 μM KH$_2$PO$_4$, 10 μM Mg$_2$SO$_4$, 100 μM CaCl$_2$, and 1 mM KCl, pH 5.6 and supplemented with either 10 μCi/ml of $^{33}$PO$_4$ or $^{35}$SO$_4$. After 8 h, nodules were washed extensively with the same solution without radioisotopes, and total radioactivity in nodules measured by crushing nodules and adding 5 mL of scintillation cocktail (Ultima Gold, Perkin Elmer). To measure the uptake of $^{33}$PO$_4$ or $^{35}$SO$_4$ into bacteroids, the nodules were crushed in ice-cold bacteroid extraction buffer (BEB=125 mM KCl, 50 mM Na-succinate, 50 mM TES, pH 7.0) containing 1% (w/v) BSA. The slurry was first centrifuged at 100 g for 10 min to remove plant debris and the suspension was then centrifuged at 3,600 g for 20 min at 4°C. The pellet was resuspended in ice-cold BEB at OD600=1, loaded on top of a 70% (v/v) percoll solution in BEB buffer and centrifuged for 30 min at 48,000 g at 4°C. The band containing bacteroids was carefully removed by pipetting, diluted in 1.4 mL of ice-cold BEB and centrifuged at 3,600 g for 10 min at 4°C. The pellet representing a purified bacteroid preparation was resuspended in a small volume of ice-cold BEB. A portion was mixed with scintillation cocktail for radioactivity measurement whereas another portion was examined by microscopy to count the number of bacteroids per volume. The amount of $^{33}$PO$_4$ or $^{35}$SO$_4$ measured was then normalized to the amount of bacteroids in the solution. Radioactivity was measured in the Perkin Elmer tri-carb 2800TR Scintillation counter.

Electron and light microscopy
Nodules were fixed in a 2.5% (w/v) glutaraldehyde solution (EMS) in 0.1 M phosphate buffer (PB) (pH 7.4) for 12 h at RT under vacuum. They were then washed several times in PB, and post-fixed in a fresh mixture of osmium tetroxide 1% (w/v) (EMS) with 1.5% (w/v) of potassium ferrocyanide in PB buffer for 3 h at RT under vacuum. The samples were then
washed twice in distilled water and dehydrated in a graded acetone solution. This was followed by infiltration in LR White resin (EMS) and finally polymerized for 48 h at 60°C in an oven under nitrogen atmosphere. Semi-thin sections 200 µm thick (for light microscopy) and ultrathin sections 50 nm thick (for TEM microscopy) were cut longitudinally in the middle of the nodule using a Leica Ultracut (Leica Mikrosysteme GmbH) and picked up on cover glass (for light microscopy) or a copper slot grid 2 x 1 mm (for TEM microscopy) coated with a polystyrene film (Sigma). Ultrathin sections were post-stained with 4% (w/v) uranyl acetate (Sigma) in water for 10 min, rinsed several times with water followed by Reynolds lead citrate in water (Sigma) for 10 min and rinsed several times with water.

Micrographs were taken with a transmission electron microscope FEI CM100 (FEI, Eindhoven, The Netherlands) at an acceleration voltage of 80 kV with a TVIPS TemCamF416 digital camera (TVIPS GmbH, Gauting, Germany). For light imaging, micrographs were taken using a Zeiss Axio Imager Z2 (Zeiss, Oberkochen, Germany) with a Hamamatsu digital camera C11440 (Hamamatsu Photonics K.K., Hamamatsu, Japan).

**In situ GUS expression in S. meliloti**

*S. meliloti* strain Rm2011 containing the plasmids orf-pit::gusA and pstS::gusA (Yuan et al., 2006) were obtained from T. M. Finan (University of McMaster, Canada) and used to infect plants transformed with EV or the RNAi3 construct. Nodules from transgenic roots of plants grown under SNF conditions were fixed in 3% (w/v) paraformaldehyde under vacuum, stained for GUS activity as described above, embedded in 6% (w/v) agarose, and cut at 100-µM thickness using a vibratome.

**Quantification and statistical analysis**

Values are presented as mean ± SD. Student’s *t*-test was used for statistical analyses, with *p* < 0.05 considered significant. The number of replicates for all experiments is indicated in the figure legends.

**Accession Numbers**

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers listed in Supplemental Figure S1.

**Supplemental Data**
**Supplemental Figure S1.** Nomenclature and structure of the *M. truncatula PHO1* gene family.

**Supplemental Figure S2.** Expression pattern of the *MtPHO1* genes.

**Supplemental Figure S3.** Analysis of plants with composite transgenic roots transformed with RNAi constructs targeting the *MtPHO1.1* and *MtPHO1.2* genes.

**Supplemental Figure S4.** Sequence of the three RNAi constructs used to down-regulate *MtPHO1.1* and *MtPHO1.2* genes.

**Supplemental Table S1.** Gene-specific primer pairs used in *Tnt1* mutant genotyping.

**Supplemental Table S2.** Gene-specific primer pairs used in RT-PCR analysis.

**Supplemental Table S3.** Gene-specific primer pairs used in making gMtPHO1
MtPHO1::GUS fusions.

**Supplemental Table S4.** Gene-specific primer pairs used in RNAi constructs.

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**Competing interest**

None
Figure Legends

Fig. 1 Phylogeny and expression of the PHO1 family in *M. truncatula*. A, Unrooted phylogenetic tree of PHO1 from Arabidopsis (green) and *M. truncatula* (black) showing the two main clades. Numbers at the nodes indicates the bootstrap values on neighbor joining analysis. B, Expression of the *M. truncatula* PHO1 genes obtained from the *M. truncatula* Gene Expression Atlas using an Affymetrix Medicago Gene Chip and RNA isolated from shoots, stems, flowers, and roots from 28-day-old plants (28d), seeds 20 days after pollination (20dap), and nodules 20 days after inoculation (20dpi) with *S. meliloti* (mtgea.noble.org/v3/). The tissues analyzed are indicated by a color code (top right corner) and the order is kept the same for all genes. C, GUS staining in *M. truncatula* roots (upper panels) and nodules (lower panels) transformed with promoter:cDNA-GUS fusions for the *M. truncatula* genes MtPHO1.1, MtPHO1.2, and MtPHO1.3. Bars represent 200 µM.

Fig. 2 MtPHO1.1 and MtPHO1.2 genes mediate Pi export. A, Measurement of Pi and nitrate export mediated by the transient expression in *N. benthamiana* leaf discs of the Arabidopsis PHO1-GFP, or *M. truncatula* PHO1.1-GFP and PHO1.2-GFP fusion proteins. As controls, Pi and nitrate export was measured in leaf discs expressing either free GFP or not infiltrated. Error bars represent SD (n=4). Values on the y axis represent the percentage of total ion present in the tissue at time 0 that is exported in the medium after 2 hours. Asterisks represent statistically significant differences compared to the GFP or non-infiltrated control (*t*-test, *p* < 0.05). B, Subcellular localization of MtPHO1.1-GFP and MtPHO1.2-GFP fusion proteins transiently expressed in *N. benthamiana*. MtPHO1.1-GFP and MtPHO1.2-GFP were co-expressed with the plasma membrane (PM) marker CBL1-OFP (upper two panels) and the Golgi marker Man1-RFP (lower two panels). Scale bars = 10 µm.

Fig. 3 Analysis of MtPHO1.1 and MtPHO1.2 insertion mutants. A, Localization of Tnt1 insertions within the MtPHO1.1 and MtPHO1.2 genes. Black boxes and lines represent exons and introns, respectively. B, Visual appearance of Tnt1 mutant plants and controls when grown for 28 days under SNF conditions. Control plants were either the R108 genotype or WT-segregants derived from heterozygous Tnt1 mutant lines. C, Expression level of the MtPHO1.1 and MtPHO1.2 genes in the various Tnt1 mutants relative to the WT control R108 (set at 1) as detected by RT-qPCR. REL, relative expression level. Error bars represent SD.
(n=3). Measurement of shoot fresh weight (D), shoot Pi content (E), nodule fresh weight (F), number of nodules per plants (G), and nodule Pi content (H) in plants grown in clay-based soil under SNF conditions for 28 days. For D to H, error bars represent SD (n ≥ 6).

**Fig. 4** Analysis of RNAi lines with reduced expression of *MtPHO1.1* and *MtPHO1.2* genes. A, Expression level of the *MtPHO1.1* and *MtPHO1.2* genes in the nodules and sections of roots devoid of nodules relative to the empty vector (EV) control as detected by RT-qPCR. REL, relative expression level. Error bars represent SD (n=3). B, Visual appearance of RNAi plants and EV controls for plants grown for 28 days in clay soil supplemented with full nutrient (FN) including nitrate and ammonium or grown under symbiotic nitrogen fixation (SNF) conditions. Measurement of shoot fresh weight (C), shoot Pi content (D), nodule fresh weight (E), number of nodules per plants (F), nodule Pi content (G), and nitrogen fixation activity of nodules (H) in plants grown in SNF conditions for 28 days. For C to H, error bars represent SD (n ≥ 6). Asterisks represent statistically significant differences compared to the EV control (*t*-test, *p* < 0.05).

**Fig. 5** Bacteroid size and distribution in nodules. A, Panoramic reconstruction of 3X3 and 5X5 pictures showing a 200 µm thick longitudinal section of nodules from empty vector (EV) control and RNAi plants grown for 28 days under symbiotic nitrogen fixation (SNF) conditions. Cells heavily colonized by bacteroids are shown as electron-dense areas. Bars = 250 µm. B, Electron micrographs of cells colonized with differentiated elongated bacteroids in the nitrogen fixation zone of nodules from EV control and RNAi plants grown for 28 days under SNF condition. Bars = 10 µm. C, The distribution of bacteroid size measured in electron micrographs of infected cells in the nitrogen fixing zone. For both nodules from EV control and RNAi plants, the size of 1,000 bacteroids was measured from 30–40 distinct cells.

**Fig. 6** Flux of Pi and SO₄ into nodules and bacteroids. Isolated nodules from empty vector (EV) control plants and RNAi plants grown in clay-based substrate under symbiotic nitrogen fixation (SNF) conditions were incubated in a nutrient solution containing either ³²³Pi (A) or ³¹⁵SO₄ (B) and the amount of ³²³Pi and ³¹⁵SO₄ absorbed by the nodule was measured. Error bars represent SD (n=4). C, The amount of ³²³Pi and ³¹⁵SO₄ acquired by bacteroids in the RNAi line was normalized to the number of bacteroids and expressed relative to the EV control. Error bars represent SD (n= 4). Asterisks represent statistically significant differences for RNAi compared to the EV control (*t*-test, *p* < 0.05). D, Cross-section of nodules stained for ß-
glucuronidase activity taken from RNAi and EV plants infected with *S. meliloti* strain Sm2011 harboring the promoter fusion orfA::gusA or pstS::gusA and grown under SNF conditions. Bars = 500 µm.


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