

# Global control of bacterial nitrogen and carbon metabolism by a PTS<sup>Ntr</sup>-regulated switch

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**The nitrogen-related phosphotransferase system (PTS<sup>Ntr</sup>) of *Rhizobium leguminosarum* bv. *viciae* 3841 transfers phosphate from PEP via PtsP and NPr to two output regulators, ManX and PtsN. ManX controls central carbon metabolism via the tricarboxylic acid (TCA) cycle, while PtsN controls nitrogen uptake, exopolysaccharide production, and potassium homeostasis, each of which is critical for cellular adaptation and survival. Cellular nitrogen status modulates phosphorylation when glutamine, an abundant amino acid when nitrogen is available, binds to the GAF sensory domain of PtsP, preventing PtsP phosphorylation and subsequent modification of ManX and PtsN. Under nitrogen-rich, carbon-limiting conditions, unphosphorylated ManX stimulates the TCA cycle and carbon oxidation, while unphosphorylated PtsN stimulates potassium uptake. The effects are reversed with the phosphorylation of ManX and PtsN, occurring under nitrogen-limiting, carbon-rich conditions; phosphorylated PtsN triggers uptake and nitrogen metabolism, the TCA cycle and carbon oxidation are decreased, while carbon-storage polymers such as surface polysaccharide are increased. Deleting the GAF domain from PtsP makes cells “blind” to the cellular nitrogen status. PTS<sup>Ntr</sup> constitutes a switch through which carbon and nitrogen metabolism are rapidly, and reversibly, regulated by protein:protein interactions. PTS<sup>Ntr</sup> is widely conserved in proteobacteria, highlighting its global importance.**

bacterial metabolism | regulatory network | nitrogen | plant–host interactions

Maintaining proper intracellular carbon and nitrogen levels is crucial in cell physiology to maximize nutrient utilization and cell growth. Among bacterial regulatory circuits, the well-conserved and ubiquitous P<sub>II</sub> family proteins (i.e., *glnB*, *glnK*, and *nifl* products) play a major role in coordinating nitrogen metabolism by signal transduction via posttranslational modifications. Among other targets, GlnK controls the activity of the ammonium transporter AmtB; GlnB regulates the expression and activity of glutamine synthetase as well as the NtrBC transcriptional cascade regulating the Ntr operon, and Nifl interacts with nitrogenase, posttranslationally inactivating this enzyme in response to ammonium (1). Upon nitrogen starvation NtrC also up-regulates the gene encoding RelA, which synthesizes guanosine tetraphosphate (ppGpp), the effector molecule of the bacterial stringent response (2). Besides transcriptional and posttranslational regulation, small RNAs (sRNAs) have also been identified as a third level of regulation for fine tuning the nitrogen network (3).

In proteobacteria, P<sub>II</sub> proteins are uridylylated under nitrogen limitation and rapidly deuridylylated under N-excess (4). This switch is mediated by GlnD, an uridylyltransferase/uridylyl-removing enzyme that senses glutamine/ $\alpha$ -ketoglutarate (the major metabolic signals reflecting the nitrogen and carbon status of the cell) and ATP/ADP ratios (reflecting the cellular energy status) (4, 5).  $\alpha$ -Ketoglutarate is a key intermediate of the tricarboxylic acid (TCA) cycle and the major carbon skeleton for nitrogen assimilation (4, 6). As the  $\alpha$ -ketoglutarate pool responds within minutes to a change in extracellular nitrogen availability

(7), bacteria have therefore evolved a variety of regulatory mechanisms to sense this effector molecule. The accumulation of  $\alpha$ -ketoglutarate under nitrogen (N)-limiting conditions also inhibits carbohydrate uptake in *Escherichia coli* by binding to the first component of the carbohydrate-phosphoenolpyruvate phosphotransferase system (PTS) (8).

In gram-negative bacteria there are two common PTS variants: the carbohydrate-PTS, which coordinates carbohydrate transport (9, 10), and the nitrogen-related PTS (PTS<sup>Ntr</sup>), a signal transduction cascade with various regulatory roles (11). Most proteobacteria implement an integrated PTS comprising PTS<sup>Ntr</sup>, encoded by the genes *ptsP* (EI<sup>Ntr</sup>), *ptsO* (NPr), and *ptsN* (EIIA<sup>Ntr</sup>), and an EIIA component remaining from the carbohydrate-PTS, *manX* (EIIA<sup>Man</sup>). The importance of PTS<sup>Ntr</sup> is suggested by its wide conservation in  $\alpha$ -,  $\beta$ -, and  $\gamma$ -proteobacteria (12, 13) and, because *ptsN* is often found contiguous with *rpoN*, which codes for the nitrogen-responsive RNA polymerase  $\sigma^{54}$  sigma factor (11, 12). PtsP contains a GAF domain homologous to the sensory domain of the NifA protein, which binds small molecules at its N terminus (14). In *E. coli*, glutamine and  $\alpha$ -ketoglutarate control phosphorylation of PTS<sup>Ntr</sup> through allosteric binding to the GAF domain of PtsP. PtsP acquires high-energy phosphate from phosphoenolpyruvate (PEP), which then phosphorylates the small carrier protein NPr (or HPr in the carbohydrate-PTS) on a conserved histidine residue (15). In *E. coli* and *Rhizobium leguminosarum* the GAF domain was reported to be dispensable for PEP-dependent PtsP autophosphorylation (16, 17).

## Significance

**Bacteria have evolved intricate regulatory networks to coordinate their metabolism with internal and external signals of their status. The regulatory phosphotransferase systems (PTSs) constitute a key part of these intricate circuits, with their signal transduction cascades participating in multiple regulatory functions. Although two major systems have been described as being involved in regulating carbon and nitrogen pools, there is very little information on their physiological role in vivo under real-time conditions. In this work we demonstrate the role of PTS as an integrated system, widely conserved in proteobacteria, acting as a complex biological sensor-actuator device enabling bacterial cells to posttranslationally alter bacterial physiology and balance carbon and nitrogen availability.**

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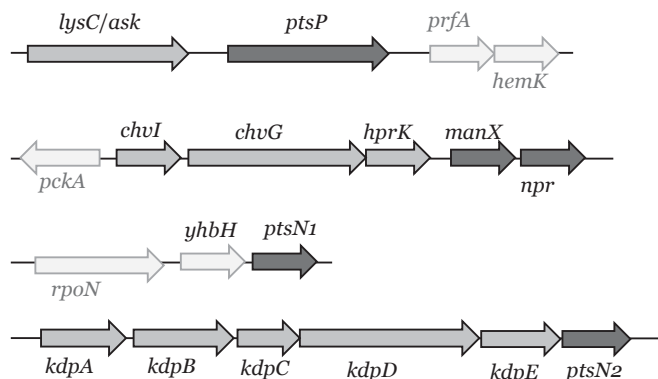
The authors declare no competing interest.

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**Fig. 1.** Schematic genetic organization of the PTS components in *R. leguminosarum* bv. *viciae* 3841. In dark, PTS components; in gray, relevant genes interacting with PTS; and in white, other neighboring genes.

Extensive previous analyses have shown that carbohydrate-PTS transport systems are the exception rather than the rule in bacteria. Most well-studied  $\alpha$ - and  $\beta$ -proteobacteria, Chlamydiae and Planctomycetes, do not contain the PTS permease proteins (EIIB and EIIC) responsible for sugar translocation (9, 13, 18, 19), although there are exceptions, such as *E. coli*. The ultimate acceptors of P<sub>i</sub> from NPr~P are therefore ManX and PtsN (EIIA components), with both proteins having an exclusively regulatory role. It is also striking that  $\alpha$ -proteobacteria mainly contain PTS<sup>Ntr</sup> components (18) organized in three operons (Fig. 1). In cluster one, aspartokinase (*lysC* or *ask*) is located upstream of *ptsP*, while in the second cluster, *npr* is located downstream of *manX*, encoding the EIIA<sup>Man</sup> homolog. Further upstream are *hprK*, the two-component regulatory system *chuI/chuG*, and phosphoenolpyruvate carboxykinase (*pckA*). HPrK is a kinase/phosphatase common in gram-positive bacteria, but absent in many gram-negative bacteria, including *E. coli* (12). It is a major regulator of carbon catabolite repression in firmicutes, where PTS<sup>Ntr</sup> genes are absent (20). Interestingly, proteobacteria of the  $\alpha$ -subdivision possess a truncated HPrK missing the N-terminal domain (about 130 amino acids) (19, 21). *ptsN* is present in the third gene cluster downstream of *yhbH*, which codes for a putative  $\sigma^{54}$ -modulating protein that can associate with ribosomes. Intriguingly, in 10% of the genomes (most of them rhizobial) there is a second copy of *ptsN*, always located downstream of the high-affinity potassium (K<sup>+</sup>) transport system *kdpABCDE*.

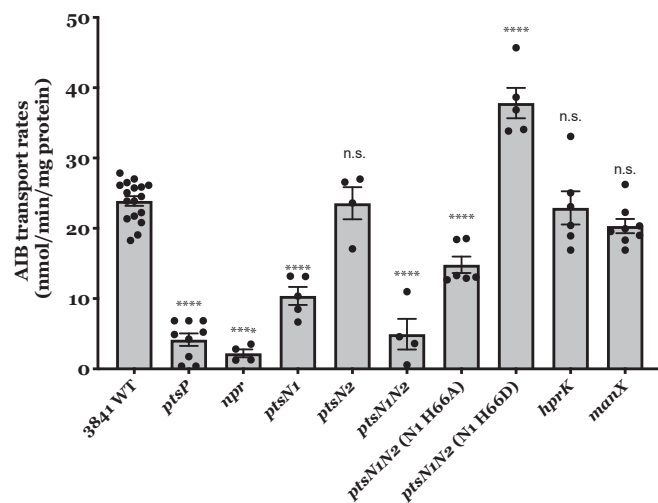
PTS systems, via their EIIA components, exert their regulatory role in numerous bacteria at the protein:protein level (9), coordinating essential processes for cell survival such as K<sup>+</sup> homeostasis and phosphate starvation, ATP-binding cassette (ABC) transporters, or central metabolic enzymes, including pyruvate dehydrogenase (PDH) and  $\alpha$ -ketoglutarate dehydrogenase ( $\alpha$ -KGDH) (22–26). Although EIIA proteins can alter transcription, their regulation is usually exerted by binding to histidine kinases such as KdpD or PhoR (24, 25). The sheer range of regulatory roles assigned to PTS that have been studied independently in different bacteria has made it difficult to understand how phosphorylation coordinated by NPr acts at the global level to control responses in bacterial cells. Therefore, we characterized the entire regulatory network of the integrated PTS in *Rhizobium leguminosarum* bv. *viciae* 3841 (Rlv3841) which, as a model  $\alpha$ -proteobacterium, lacks the carbohydrate-PTS and sugar translocation components (EIIB and EIIC). Rlv3841 also has a second *ptsN* copy named *ptsN2* and is located on plasmid pRL11 (pRL110376) (23) (Fig. 1). This highly conserved operon arrangement highlights the relevance of the PTS<sup>Ntr</sup> branch in  $\alpha$ -proteobacteria and emphasizes the importance of

using a model bacterium to unravel the mechanism of action of PTS. We show in this study that this integrated PTS system is another major posttranscriptional regulator, with its two output regulators, PtsN and ManX, acting reciprocally to integrate carbon and nitrogen metabolism signals at the cellular level.

## Results and Discussion

**PTS<sup>Ntr</sup> Phosphorylation Is Essential for ABC Transport Activation.** The PTS<sup>Ntr</sup> phosphorelay starts with PEP as the high-energy phosphate donor to PtsP on His367. This phosphate is then transferred to NPr on His17 and PtsN on His66 (15). PtsN (EIIA<sup>Ntr</sup>) mediates the effects of PTS in Rlv3841 on the activation of ABC transporters and K<sup>+</sup> uptake (17, 23). Mature bacteroids become symbiotic auxotrophs and their host plant must provide branched-chain amino acids to support their development and persistence (27). Active amino acid transport is therefore essential for nitrogen fixation in pea nodules; mutations of the main ABC-type broad-specificity amino acid uptake systems (Aap and Bra) led to severely N-starved plants (27, 28). In both Rlv3841 and *E. coli*, nonphosphorylated PtsN interacts with KdpD, the sensor kinase that activates the transcription of the high-affinity K<sup>+</sup> transporter KdpABC via the response regulator KdpE (23, 24). However, EI (PtsP) phosphorylation on His367 and NPr on His17 are required for ABC transport activation, suggesting that PtsN~P is the active species (17).

By examining the complete PTS network, we now demonstrate that a *manX* mutant is not significantly affected in transport, while confirming that PtsN, as well as PtsP and NPr (required for phosphoryl-group transfer), are essential for transport activation (Fig. 2). Amino acid transport was measured with  $\alpha$ -aminoisobutyric acid (AIB) because, like glutamine, it is transported exclusively by Aap and Bra, but cannot be metabolized (29). As previously seen, the *ptsN1* mutant (LMB271) had a large reduction in transport, whereas the *ptsN2* mutant (RU4193) maintained wild-type levels (23). The essentiality of the PTS<sup>Ntr</sup> branch in transport activation was also shown for glucose (SI Appendix, Fig. S1). These results agree with our initial work on PTS in Rlv3841, where we already measured AIB



**Fig. 2.** Transport rates by PTS mutants. Standard rates were obtained from cultures grown on UMS with 10 mM glucose and 10 mM NH<sub>4</sub>Cl. Rlv3841 wild type, *ptsP* (PtsP107), *npr* (AA031), *ptsN1* (LMB271), *ptsN2* (RU4391), *ptsN1N2* (AA047), *ptsN1N2* (N1 H66A, OPS1102), *ptsN1N2* (N1 H66D, OPS1104), *hprK* (AA081), and *manX* (LMB692). All rates are expressed in nmol min<sup>-1</sup> mg protein<sup>-1</sup>. Data are averages ( $\pm$ SEM) from at least three independent cultures analyzed by one-way ANOVA with Dunnett's posttest for multiple comparisons (\*\*\*\*)  $P < 0.0001$  and n.s., not significant.

transport (taken up with very similar kinetics to glutamate),  $\gamma$ -amino butyric acid (GABA),  $\delta$ -aminolevulinic acid (ALA), glucose, and myo-inositol (23). These substrates, taken up by ABC transporters, were all greatly reduced in the *ptsP* mutant compared with wild-type Rlv3841, whereas this was not the case for succinate transport, driven by the proton-coupled transporter DctA.

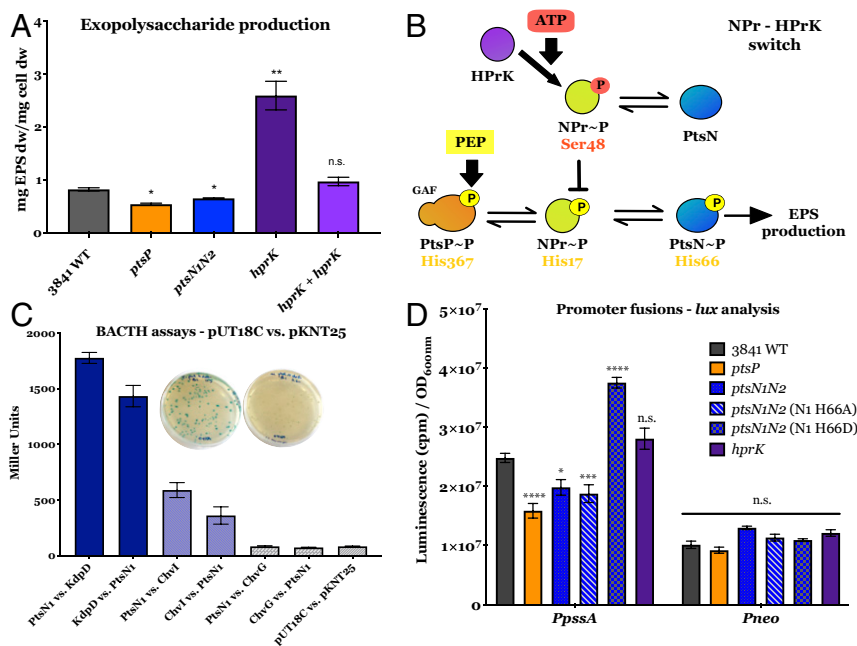
To confirm the role of phosphorylated PtsN1 in activating ABC transporters, we replaced the chromosomal *ptsN1::* $\Omega$ Spec insertion with a permanent phosphorylation (H66D, strain OPS1104) or nonphosphorylation mimic (H66A, strain OPS1102) in the double *ptsNIN2* mutant (AA047). The PtsN1 H66D phosphomimic (OPS1104) significantly increased AIB transport rates above the level observed with the native protein. While H66A shows a partial activation, it does not restore transport rates to wild-type level (Fig. 2). This is consistent with PtsN1~P being required for full activation of a wide range of ATP-dependent ABC transport systems in *R. leguminosarum*.

**Exopolysaccharide Secretion Is Regulated by PtsN via Interaction with the ChvI/ChvG System.** Extracellular polysaccharides are carbon-based surface polymers indispensable for the invasion of a large number of host plants which form indeterminate nodules (30). We had previously shown that a functional PTS is required for normal production of exopolysaccharide (EPS) and therefore a mucoid surface phenotype in *R. leguminosarum* that produces a high-molecular-weight acidic EPS (17, 23). When grown on agar plates, we now show that mutants that disrupt the phosphorylation cascade of PTS<sup>Ntr</sup> (*ptsP*, *npr*, or *ptsNIN2*) have a dry surface compared to wild type (SI Appendix, Fig. S2). Total EPS was quantified in Rlv3841 and several PTS mutants (Fig. 3A), showing a reduction of 35% in PtsP107 (*ptsP::*Tn5) and 21% in

AA047 (*ptsN1::* $\Omega$ Spec;  $\Delta$ *ptsN2*). Since *ptsNIN2* and mutants in the phosphotransfer cascade have a dry surface, PtsN~P is likely to be required for EPS production (Fig. 3B). Plasmid complementation of AA047 with either wild-type *ptsN* or *ptsN* H66A (permanent nonphosphorylated version) showed that only complementation with *ptsN* gives a mucoid surface phenotype (SI Appendix, Fig. S2), indicating that PtsN~P is the likely species that activates both ABC transport and EPS production in Rlv3841.

The gene encoding NPr is located close to those that encode a two-component global regulatory system, ChvI/ChvG, which modifies the cell surface (Fig. 1); the homologous system in *Sinorhizobium meliloti*, ChvI/ExoS, controls succinoglycan and galactoglycan production (31, 32). *R. leguminosarum* synthesizes only an acidic EPS and a *chvG* mutation was shown to decrease its production, among other pleiotropic effects (33). We investigated the potential link between PTS and ChvI/ChvG using bacterial two-hybrid (BACTH) analysis. PtsN1 and the response regulator ChvI interacted strongly (Fig. 3C), while PtsN1 and the membrane sensor kinase ChvG did not. This interaction suggests that the effect of PTS on EPS is mediated by ChvI, which acts at the transcriptional level (34). A plasmid-borne *lux* fusion to the promoter of *pssA*, the gene responsible for the polymerization of EPS (30), was conjugated into the *ptsP* mutant (dry phenotype) as well as in the double mutant *ptsNIN2* and its derivatives with the different versions of *ptsN1* (H66A, dry; H66D, mucoid). Increased expression of *pssA* in the *ptsN1* H66D strain and decreased expression in *ptsNIN2* and *ptsN1* H66A (Fig. 3D) is consistent with PtsN~P stimulating EPS production via the interaction with the ChvI regulator.

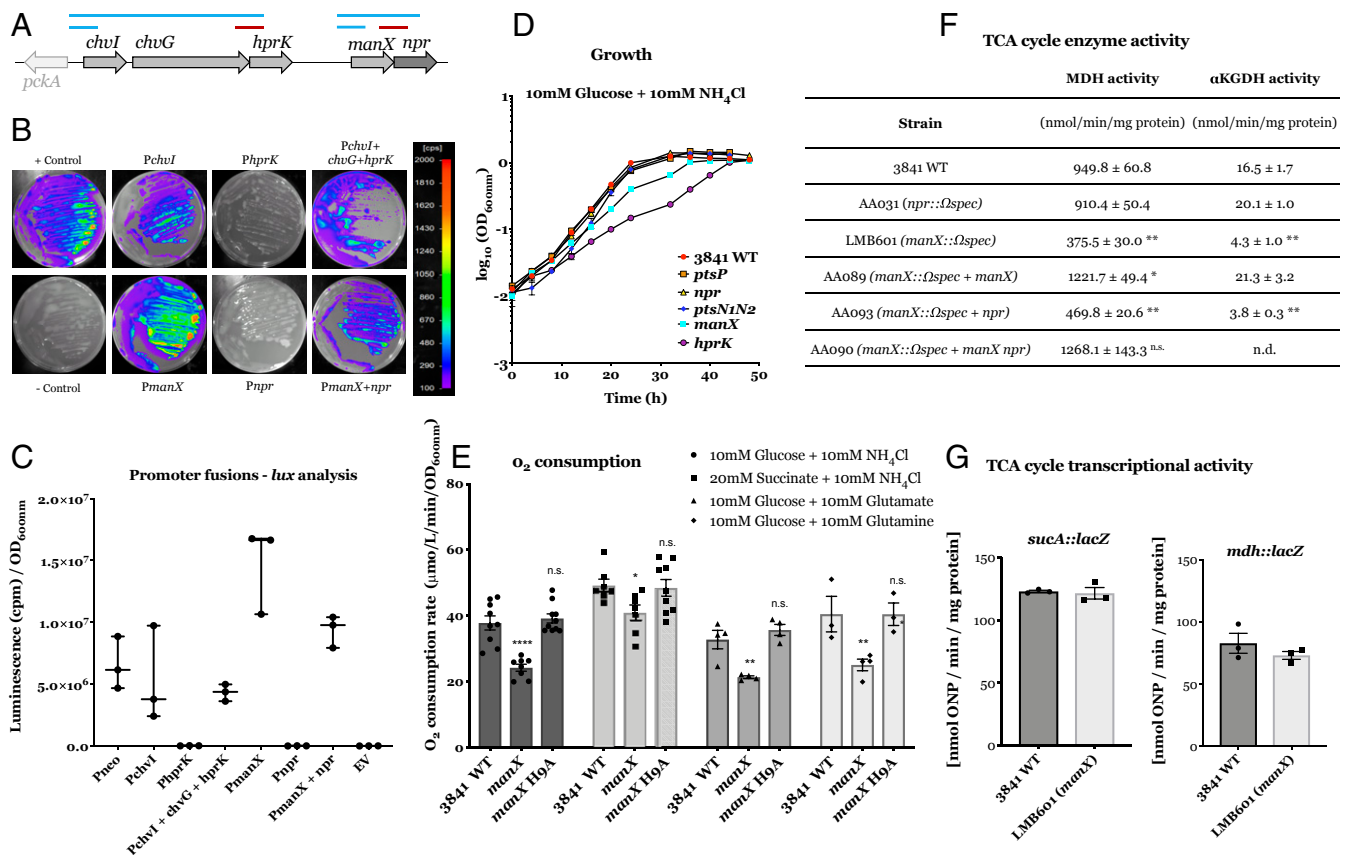
In summary, while the first defined role assigned to dephosphorylated PtsN was to control K<sup>+</sup> homeostasis by direct binding



**Fig. 3.** EPS secretion and regulation. (A) EPS production measurements for Rlv3841 wild type, *ptsP* (PtsP107), *ptsNIN2* (AA047), *hprK* (AA081), and *hprK + hprK* (AA088 complemented strain). Data are averages ( $\pm$ SEM) from at least four independent cultures analyzed by one-way ANOVA with Dunnett's posttest for multiple comparisons (\*)  $P < 0.01$ , (\*\*)  $P < 0.01$  and n.s., not significant. All rates are expressed in mg EPS dry weight  $\text{mg}^{-1}$  cell dry weight. (B) Npr-HPrK switch model. PTS<sup>Ntr</sup> components are phosphorylated on histidine residues, initially (PtsP) from PEP as indicated, with PtsN~P inducing EPS production. Conversely, HPrK phosphorylates NPr on Ser48 from ATP, blocking subsequent phosphorylation of PtsN. (C) BACTH assays of the interaction of PtsN with the ChvI/ChvG system. Interactions expressed in Miller units of LacZ activity. Plates show blue colonies indicating a positive interaction (PtsN1-ChvI) and white colonies indicating a negative interaction (PtsN1-ChvG). PtsN1-KdpD and KdpD-PtsN1 are positive controls. Data are averages ( $\pm$ SEM) from at least three independent cultures. (D) Quantification of promoter activity for *PpsaA* compared to the constitutive promoter *nptII* (*Pneo*) in Rlv3841 wild type, *ptsP* (PtsP107), *ptsNIN2* (AA047), *ptsNIN2* (N1 H66A, OPS1102), *ptsNIN2* (N1 H66D, OPS1104), and *hprK* (AA081). All rates are expressed in counts per minute (cpm). Data are averages ( $\pm$ SEM) from at least five independent cultures. Statistical analyses are indicated following the level of significance, with (\*\*\*\*) as  $P$  value  $< 0.0001$ , (\*\*\*) as  $P$  value  $< 0.001$  and n.s., not significant.

to KdpD in *R. leguminosarum* (23), as it does in *E. coli* (23, 24), we have now confirmed that PtsN~P regulates Aap and Bra and most likely a wide range of ABC transporters. We have also demonstrated the direct interaction of PtsN with ChvI (Fig. 3C), suggesting that the surface phenotype of PTS mutants is mediated through this global regulator of EPS production. The *chvIG* operon encodes a two-component regulatory system involved in virulence or symbiosis [BvrR/BvrS in *Brucella* spp (35), ChvI/ChvG in *Agrobacterium tumefaciens* (36), and ChvI/ExoS in *S. meliloti* (32)]. Cross-talk between this system and PTS<sup>Ntr</sup> had been proposed in *Brucella melitensis*, suggesting that PTS communicates the metabolic state of the cell to the virulence gene *virB*, by phosphorylating or interacting with the BvrR/S system (26). In rhizobia, this two-component regulatory system has already been described as a master transcriptional regulator of

EPS (31, 32, 37). Indeed, based on the genetic proximity of *chvGI* to *hprK* and the opposite phenotypes between these mutants in terms of growth (*chvG* and *chvI* mutants are unable to grow on complex media, while the *hprK* mutant is able to) and EPS production (*chvG* and *chvI* show reduced EPS, *hprK* increased EPS), PTS was suggested to play a role in the dephosphorylation of ExoS/ChvI in *S. meliloti* (32). This suggestion fits our model, where HPrK phosphorylation of Ser48 on NPR reduces phosphorylation on its His17, resulting in reduced EPS (Fig. 3B). The BACTH data point toward PtsN~P binding to ChvI and stimulating EPS production at the transcriptional level, a hypothesis reinforced after showing that *ptsA* is transcriptionally up-regulated when PtsN1 is present as a permanent phosphomimic (strain OPS1104, PtsN1 H66D, Fig. 3D). This differential role of PtsN driven by its phosphorylation status



**Fig. 4.** Promoter activity analysis, growth phenotype, and TCA enzymatic activity and regulation. (A) Genetic organization of *chvI/chvG/hprK* and *manX/npr* operons. In blue, lines indicate regions with promoter activity and in red, those without. (B) Qualitative luminescence imaging showing the activity of the six different potential promoter regions fused to the *luxCDABE* cassette: *PchvI* (pOPS0298) encompasses the region immediately upstream of the *chvI* operon, and *PchvI+chvG+hprK* (pOPS0313) contains the same region with the *chvI*, *chvG*, and *hprK* genes included. *PmanX* (pOPS0606) and *PmanX+npr* (pOPS0296) contain the putative promoter-encoding region in front of *manX* and this region with the entire *manX* operon included, respectively. Regions with potential promoters for *npr* (*Pnpr*) (pOPS0605) and *hprK* (*PhprK*) (pOPS0603) were also included for analysis. (Scale bar on the Right, in counts per minute [cpm].) (C) Quantification of promoter activity from the regions selected for testing. The constitutive *Pneo* promoter was used as a positive control on the plasmid pJ11282 and the empty vector pJ11268 (EV) as a negative control. All rates are expressed in cpm. Data are averages (±SEM) from three independent cultures. (D) Growth curves on UMS supplemented with 10 mM glucose + 10 mM NH<sub>4</sub>Cl for strains wild-type Rlv3841, *ptsP* (ptsP107), *npr* (AA031), *ptsN1N2* (AA047), *manX* (LMB692), and *hprK* (AA081). (E) O<sub>2</sub> consumption assay on UMS supplemented with 10 mM glucose + 10 mM NH<sub>4</sub>Cl, 20 mM succinate + 10 mM NH<sub>4</sub>Cl, 10 mM glutamate + 10 mM glucose, or 10 mM glutamine + 10 mM glucose. Strains tested are *ptsP* (ptsP107), *npr* (AA031), *ptsN1N2* (AA047), *manX* (LMB692), *hprK* (AA081), and *manX* H9A (OPS1012). O<sub>2</sub> consumption rates are expressed in μmol L<sup>-1</sup> min<sup>-1</sup> OD<sub>600nm</sub><sup>-1</sup>. Data are averages (±SEM) from at least three independent cultures; two-way ANOVA with Dunnett's posttest for multiple comparisons (\*) *P* < 0.05 (\*\*) *P* < 0.01 (\*\*\*) *P* < 0.0001 and n.s., not significant. (F) MDH and α-KGDH, alpha-ketoglutarate dehydrogenase activities for Rlv3841 wild-type, *npr* (AA031), *manX* (LMB601), and complemented strains AA089, AA093, and AA090. Values expressed as nmol<sup>-1</sup> min<sup>-1</sup> mg protein. Data are averages (±SEM) from three independent cultures. Statistical analyses are indicated following the level of significance, (\*) *P* value < 0.01, (\*\*) *P* value < 0.001, and n.s., not significant; n.d., not determined. (G) Transcriptional activity assays in Rlv3841 (black) and LMB601 (*manX::Ωspec*, gray) by using transcriptional LacZ fusions in the *mdh-sucCDAB* operon located on two different cosmids, pRU3068 *sucA::lacZ* and pRU3070 *mdh::lacZ*. Data are averages (±SEM) from three independent cultures. Values expressed as nmol ONP min<sup>-1</sup> mg protein<sup>-1</sup>. No significant differences were observed.

suggests that a phosphorylation switch on PtsN might operate to activate one system while inhibiting another. Given that Rlv3841, along with most  $\alpha$ -proteobacteria, has a second EIIA component (ManX) as part of the integrated PtsP-NPr phosphorylation cascade, there may be a more comprehensive switch operating to control a wide range of ABC transporters and metabolism in bacteria.

**Expression of *manX* Is Independent of the *chvI* Promoter.** To investigate the role of ManX in *R. leguminosarum*, we first analyzed its operon structure. *manX* and *npr* genes are colocalized with *chvI*, *chvG*, and *hprK*, potentially as part of an operon (Fig. 1). Lux-fusion analysis showed that *manX* and *npr* are transcribed together, as are *chvI*, *chvG*, and *hprK* (Fig. 4 A–C). The carbohydrate-PTS system is known to be regulated by changes in carbon source in other bacterial species, where it controls their uptake (38). While the *manX* promoter showed small but significant differences in expression in minimal medium after 22 h but not after 18 h of growth, the main difference was the increased expression in rich medium (tryptone-yeast [TY]) for both the *chvI* and *manX* promoters (SI Appendix, Fig. S3). We have therefore confirmed that *manX* and *npr* are transcribed from a single promoter in front of *manX*, independently of the *chvI*-*chvG*-*hprK* operon, with this genomic region conserved across different host-invading bacteria and characteristic of the Rhizobiales (39).

**ManX Is Required for the Activation of the TCA Cycle.** *S. meliloti* *manX* mutants have reduced growth on several carbon sources and altered succinate-mediated catabolite repression (40). Previous work in other systems had already reported direct binding of PTS proteins to enzymatic components (26, 41). This suggested that ManX regulation might be due to direct binding to TCA cycle enzymes. To investigate ManX in Rlv3841, a markerless in-frame *manX* mutant (LMB692) was generated to prevent polar effects on *npr*. LMB692 showed reduced growth rates in all of the conditions tested (Fig. 4D and SI Appendix, Table S1), whereas this was not the case for mutants in components needed for phosphorylation (*ptsP*, *npr*), nor *ptsN1N2*. This defect in growth was further investigated by measuring oxygen (O<sub>2</sub>) consumption, which indicates reductant generated by the TCA cycle feeding directly into the respiratory chain in free-living cultures of the obligate aerobe *R. leguminosarum*. The *manX* mutant had a reduced O<sub>2</sub> consumption rate, regardless of the carbon or nitrogen source tested (Fig. 4E).

The contiguous arrangement of *manX* and *npr* in Rlv3841 suggested the possibility of polar effects between the genes. Although LMB692 is a markerless in-frame mutant, we tested for polarity effects by introducing *manX* and *npr* genes individually in LMB692 (*manX* mutant) and AA031 (*npr* mutant) under their native promoters on the low-copy plasmid pRK415. AA031 complemented by *npr* recovered the mucoid colony morphology of the wild-type strain. Likewise, complementation of LMB692 with *manX*, but not *npr*, restored growth and O<sub>2</sub> consumption to wild-type levels (SI Appendix, Fig. S4). Thus, as expected, the markerless mutation in *manX* is not polar on *npr*, indicating that the effect on O<sub>2</sub> consumption is exclusively due to the absence of ManX.

It had already been demonstrated by a yeast two-hybrid approach in the  $\alpha$ -proteobacterium *B. melitensis* that ManX interacts with SucA, the E1 component of  $\alpha$ -KGDH (26). We therefore analyzed the two halves of the TCA cycle by measuring  $\alpha$ -KGDH and malate dehydrogenase (MDH) enzyme activities in Rlv3841, AA031 (*npr*:: $\Omega$ Spec), and LMB601 (*manX*:: $\Omega$ Spec). Both enzyme activities were reduced by 70 and 60%, respectively, in the *manX* mutant (Fig. 4F), whereas in the *npr* mutant,  $\alpha$ -KGDH and MDH activities were similar to wild-type levels. LMB601 (*manX*:: $\Omega$ Spec) was complemented for enzyme

activity by plasmid-borne *manX*, but not *npr*. Thus, ManX alone is required for the full activation of these two TCA cycle enzymes. The reduction of TCA cycle activity, as measured by O<sub>2</sub> consumption (Fig. 4E) and enzyme activity (Fig. 4F), explains the high generation times of the *manX* mutant (SI Appendix, Table S1).

The structural genes for MDH (*mdh*) and  $\alpha$ -KGDH (*sucC-DAB*) are organized in the same cluster of genes in *R. leguminosarum* (42). Transcriptional down-regulation could therefore explain a concerted reduction in activity of  $\alpha$ -KGDH and MDH. This cluster of genes has a promoter in front of *mdh*, suggesting an operon, but there is also a second promoter in front of *sucA*. We compared their transcriptional activity in LMB601 (*manX*:: $\Omega$ Spec) and wild-type strains using transcriptional Tn5-*lacZ* fusions located in *mdh* or *sucA* in two different cosmids containing the entire *mdh-sucABCD* cluster, pRU3070 *Pmdh*::*lacZ* and pRU3068 *PsucA*::*lacZ* (42). Both LMB601 and Rlv3841 showed identical levels of *lacZ* expression for *mdh* and *sucA* fusions (Fig. 4G). Therefore, down-regulation of *mdh-sucC-DAB* transcription does not account for the reduced enzyme activities, suggesting that this effect is due to posttranscriptional regulation and agrees with the demonstrated protein:protein interaction between ManX and SucA in *B. melitensis* (26). We therefore suggest that ManX interacts with these TCA cycle enzymes directly or indirectly to regulate their activity in Rlv3841. There are other established protein:protein interactions of EIIA proteins with enzymes. In *Salmonella typhimurium* phosphorylated PtsN tightly binds to GlnS (D-glucosamine-6-phosphate synthase) inhibiting the enzyme (43), while in *Pseudomonas putida* unphosphorylated PtsN inhibits pyruvate dehydrogenase (41). In *R. leguminosarum* the unphosphorylatable *manX* mutant (H10A) shows growth at wild-type levels (Fig. 4D) and activates respiration associated with the TCA cycle regardless of the carbon or nitrogen source (Fig. 4E), indicating that nonphosphorylated ManX acts on the TCA cycle.

**The Phosphorylation Switch through NPr Controls ManX and PtsN Activity.** It was reported from in vitro experiments that *B. melitensis* NPr also phosphorylates ManX on His9, similarly to PtsN (26). An *hprK* homolog upstream of *manX* and *npr* increases the complexity of PTS-mediated regulation by interfering in this phosphoryl transfer pathway (Figs. 1 and 3B). HPr proteins like NPr have been shown to be phosphorylated on two different residues: His17 and Ser48 (44). In *S. meliloti* and *B. melitensis*, phosphorylation of NPr on the serine residue by HPrK slows down or prevents phosphorylation on the histidine by PtsP (26, 45). Thus, histidine phosphorylation of ManX and PtsN should increase in an *hprK* mutant. Accordingly, we hypothesized that NPr phosphorylation on Ser48 by HPrK would raise the amount of dephospho-EIIAs (ManX and PtsN), while an *hprK* mutation would show hyperphosphorylation of EIIAs (PtsN and ManX). Insertion of an  $\Omega$ Spec marker cassette into *hprK* (strain AA081) led to colonies that were hypermucoid and grew extremely slowly on TY and universal minimal salts (UMS) media. As observed in *S. meliloti* (45), AA081 (*hprK*:: $\Omega$ Spec) showed a large increase in EPS, which was restored to wild-type levels in the complemented strain AA088 (*hprK*:: $\Omega$ Spec + *hprK*) (Fig. 3A and SI Appendix, Fig. S2). The hypermucoid phenotype is consistent with PtsN~P promoting EPS synthesis (Fig. 3B).

In *Ralstonia eutropha* the elevated histidine-phosphorylated NPr formed in the *hprK* mutant negatively affects growth of this organism (46). Consistent with this, an *hprK* mutant (strain AA081), unable to prevent the phosphorylation of ManX or PtsN, has the most defective growth of all of the mutants tested, with the longest mean generation time (SI Appendix, Table S1 and Fig. 4D) and excess EPS production. These phenotypes are as expected for hyperphosphorylated PtsN, i.e., a wide range of ABC transporters fully active and EPS production up-regulated,

while the absence of dephosphorylated ManX suppresses TCA cycle activity. As in Rlv3841, deletion of *hprK* in *S. meliloti* also causes a strong growth defect on minimal medium and up-regulation of EPS production (45). Absent in gram-negative enteric bacteria, the HPrK/P regulator is encoded in the genomes of many proteobacteria (19). However, it has a shorter sequence than its counterpart in gram-positive bacteria, lacking about 130 amino acids including an N-terminal region important for the phosphatase activity of HPrK/P (21, 47). This indicates that HPrK in proteobacteria might not be able to efficiently dephosphorylate P~Ser~NPr (26, 45), highlighting the importance of the phosphorylation switch coordinated by NPr in these organisms. Indeed, HPrK was shown to be essential in *R. eutropha* due to the PTS phosphorelay imbalance caused by an elevated amount of P~His~NPr (46). We note that protein SixA, a well-conserved protein found in proteobacteria, actinobacteria, and cyanobacteria (48), has recently been reported as a phosphohistidine phosphatase in *E. coli* acting on NPr (49). It might play an additional role in dephosphorylating P~His~NPr.

As seen above, the lack of effect of *ptsP* and *npr* mutations on the TCA cycle enzyme activity compared with the strong reduction observed in the *manX* mutant suggests that ManX in its nonphosphorylated form is needed for full TCA activity. Accordingly, the hyperproduction of EPS and activation of a wide range of ABC transporters (due to PtsN~P) combined with a lower TCA cycle enzyme activity (elevated ManX~P depletes nonphosphorylated ManX) would explain why the *hprK* mutant (AA081) shows the highest generation times of all strains tested (SI Appendix, Table S1). Thus, ManX should activate the TCA cycle in its nonphosphorylated version and, indeed, a genomic *manX* H9A mutant (OPS1012) restored the growth rate under all conditions tested (SI Appendix, Table S1). O<sub>2</sub> consumption in minimal medium with different carbon and nitrogen sources was restored to wild-type levels in the *manX* H9A strain (Fig. 4E), confirming that nonphosphorylated ManX activates the TCA cycle in Rlv3841 as measured by carbon oxidation. Therefore, while PtsN~P activates uptake and carbon storage into the EPS layer, carbon catabolism via the TCA cycle will be down-regulated by reduced unphosphorylated ManX (depleted by the formation of ManX~P). Consequently, PTS<sup>Ntr</sup> acts as a central switch coordinating metabolism of both carbon and nitrogen.

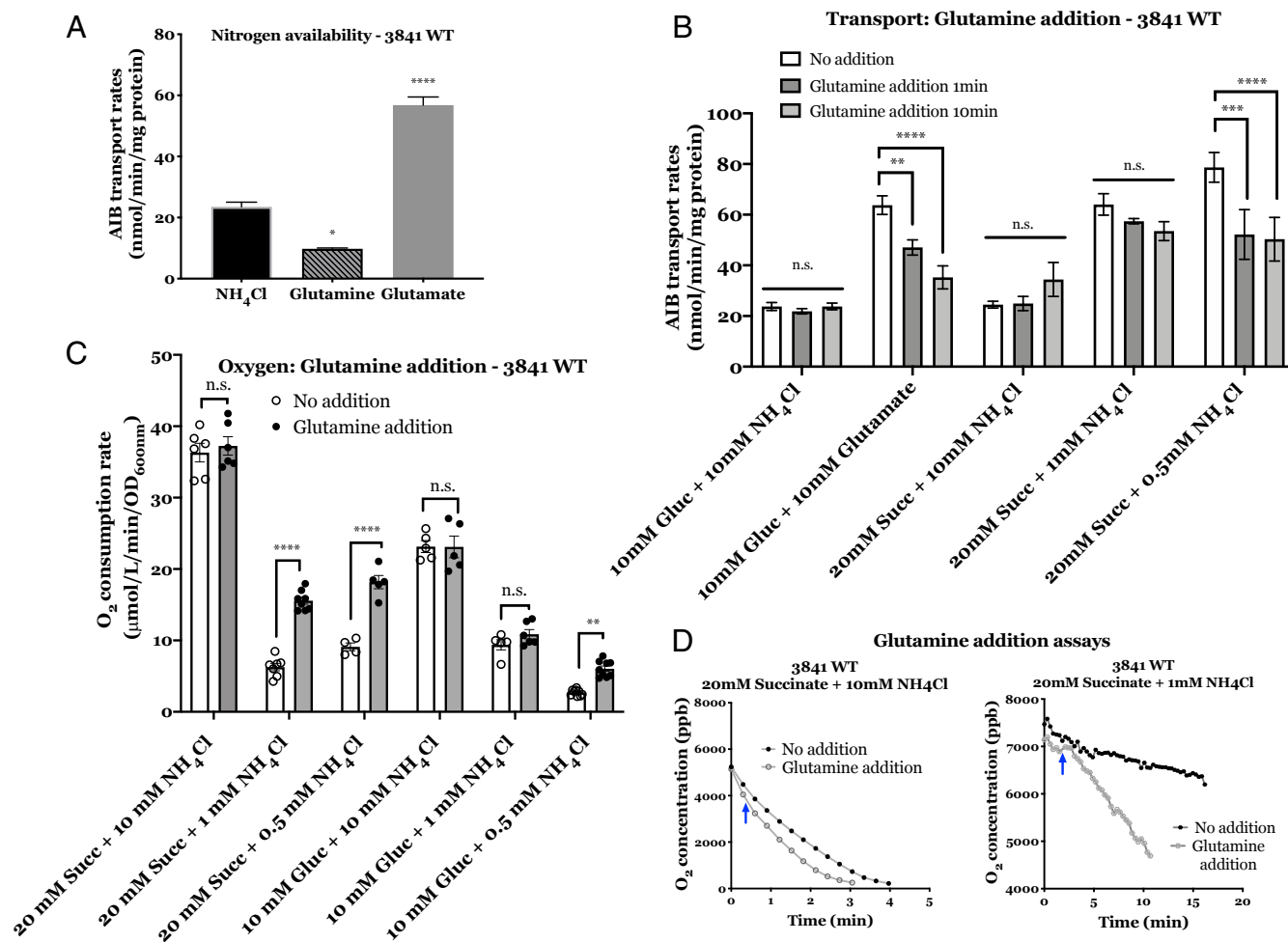
#### **In Vivo Experiments Show PTS<sup>Ntr</sup> Is Regulated by Nitrogen Availability.**

The complete carbohydrate-PTS senses the presence or absence of carbohydrates in the medium related to the intracellular PEP/pyruvate ratio (50), increasing the ratio of nonphosphorylated/phosphorylated PTS components with the intake of carbohydrates (38). In the case of the PTS<sup>Ntr</sup> system, the N-terminal GAF sensory domain modulates its autophosphorylation (16). In *E. coli* PtsP is regulated by its GAF domain binding glutamine and  $\alpha$ -ketoglutarate (16), whereas in *S. meliloti*, only glutamine binds the GAF domain of PtsP in vitro and inhibits its phosphorylation (51). In *S. typhimurium*, PtsN was also reported to be dephosphorylated in response to nitrogen excess and rapidly degraded by Lon protease upon depletion of cellular amino sugars (43). In Rlv3841, PtsP lacking its GAF domain activates ABC transport (17) and complements the dry surface phenotype of PtsP107 (*ptsP* mutant), suggesting that the ligand binding to GAF negatively regulates PTS phosphorylation. Overall, while in vitro protein work and mutant complementation studies suggest a role of PtsP in nitrogen signaling (8, 16, 51, 52), in vivo physiological evidence has been lacking. To develop definitive in vivo assays, wild-type cells were grown in minimal medium (UMS) supplemented with either succinate (20 mM) or glucose (10 mM) as carbon sources, and either 10 mM NH<sub>4</sub>Cl or 10 mM glutamine for N-rich conditions, or 10 mM glutamate and 0.5 to 1 mM NH<sub>4</sub>Cl as N-limiting conditions. While ammonia either diffuses freely across the membrane or is incorporated via the AmtB permease, glutamate

and glutamine are transported actively into cells by the ABC systems Aap and Bra (53).

As expected, Rlv3841 cultures grown overnight on glutamine (N-rich) had decreased amino acid transport rates via Aap and Bra relative to Rlv3841 cultures grown on glutamate (N-limiting) (Fig. 5A). This decrease would currently be attributed to transcriptional regulation by NtrC in most bacteria, including Rlv3841 (54). However, we reasoned that it might be due to PtsP regulation, with glutamine signaling via PTS and unrelated to NtrC-dependent transcription. To test this, cells grown on glucose or succinate with glutamate (10 mM) or different concentrations of NH<sub>4</sub>Cl (10, 1, and 0.5 mM) were subsequently harvested and resuspended without carbon or nitrogen sources. Cells were then incubated in glutamine (100  $\mu$ M), for 1 or 10 min, harvested, and washed to remove extracellular glutamine, which sterically inhibits AIB transport. Just 1 min after glutamine exposure, AIB transport was significantly decreased in N-limiting conditions (10 mM glucose + 10 mM glutamate, 20 mM succinate + 0.5 mM NH<sub>4</sub>Cl; Fig. 5B). The speed of the response in this in vivo test of amino acid uptake indicates a rapid protein:protein interaction inhibiting uptake via Aap and Bra. This is consistent with glutamine binding the GAF domain of PtsP to reduce its phosphorylation and subsequently that of PtsN and ManX (16, 51, 52). Reduced PtsN~P explains the decreased amino acid uptake (Fig. 2) and critically predicts that increased dephosphorylated ManX should activate the TCA cycle and O<sub>2</sub> consumption. O<sub>2</sub> consumption was therefore measured in liquid cultures growing in exponential phase on succinate or glucose with different NH<sub>4</sub>Cl concentrations (10, 1, and 0.5 mM). The initial rates of O<sub>2</sub> consumption were much higher under N-rich conditions (10 mM NH<sub>4</sub>Cl) than N-limiting conditions (1 and 0.5 mM NH<sub>4</sub>Cl), showing that the activity of the TCA cycle in Rlv3841 is reduced when cells are N-starved (Fig. 5C and D). The O<sub>2</sub> consumption rate did not increase significantly when glutamine was added to Rlv3841 grown under N-rich conditions (Fig. 5C). However, the rate increased immediately when glutamine was added to cells grown under N-limiting conditions. We observed an increased O<sub>2</sub> consumption when glutamine was added to cells growing on 20 mM succinate with 1 mM and 0.5 mM NH<sub>4</sub>Cl, whereas when glucose was the carbon source, the increase was only observed when glutamine was added at the lowest NH<sub>4</sub>Cl concentration, 0.5 mM NH<sub>4</sub>Cl. This indicates that cells were more severely N-deprived on succinate. Thus, as predicted, the TCA cycle responds dramatically to an up-shift in nitrogen status (i.e., glutamine), indicating that it is immediately and tightly regulated by nitrogen availability. As a control, we carried out both transport and O<sub>2</sub> consumption assays adding minimal medium or water and glutamate in place of glutamine to N-limiting conditions (SI Appendix, Figs. S5 and S6). Whereas no significant effect was observed with the addition of plain media or water, there was a small increase when glutamate was added that can be accounted for by the rapid conversion of glutamate to glutamine (7), consistent with the role of glutamine in PTS regulation.

**GAF-Deletion Mutants Are Blind to Nitrogen Status.** The results of the glutamine addition experiments are consistent with the proposed model of reciprocal regulation of PtsN and ManX. However, we still needed to show that the effects observed were dependent on the presence of the GAF domain binding glutamine to prevent PTS phosphorylation as previously described (16, 51, 52). Therefore, chromosomal PtsP::Tn5 was replaced with a version of PtsP lacking the GAF domain, generating *ptsP* $\Delta$ GAF (OPS1010), a strain that recovered the wild-type mucoid surface, growth on glutamate, and the ability to transport amino acids (SI Appendix, Table S1 and Fig. 6A), indicating that PtsP $\Delta$ GAF is a functional protein. Remarkably, OPS1010 no longer responded to the nitrogen status of the growth



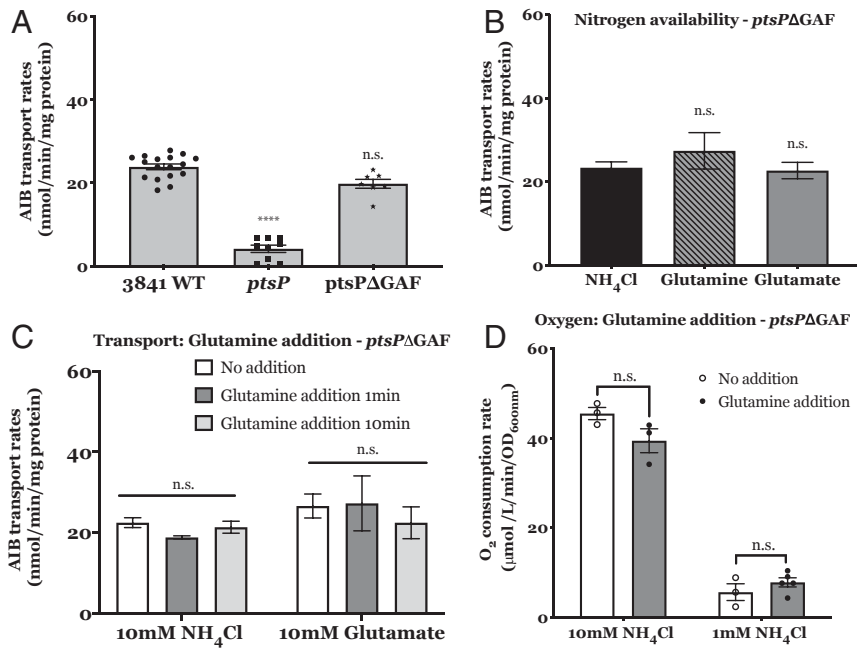
**Fig. 5.** Effect of glutamine on the PTS regulatory cascade of Rlv3841. (A) Effect of nitrogen availability on AIB transport. Membrane transport of wild-type cells grown on UMS with 10 mM glucose as carbon source and 10 mM NH<sub>4</sub>Cl, glutamine (N-rich conditions) or glutamate (N-limiting conditions). (B) Effect of glutamine addition on AIB transport. Wild-type cells grown on UMS with 10 mM glucose as carbon source and 10 mM NH<sub>4</sub>Cl (N-rich conditions) or 10 mM glutamate (N-limiting conditions). Transport rates measured at 1 min and 10 min after glutamine addition. All transport rates are expressed in nmol min<sup>-1</sup> mg protein<sup>-1</sup>. (C) Effect of glutamine addition on O<sub>2</sub> consumption of cells grown under different nitrogen conditions. Wild-type cells grown on UMS with 20 mM succinate as carbon source and 10 mM NH<sub>4</sub>Cl (N-rich conditions) or 1 mM NH<sub>4</sub>Cl (N-limiting conditions). Values show the rates of O<sub>2</sub> consumption expressed in μmol L<sup>-1</sup> min<sup>-1</sup> OD<sub>600nm</sub><sup>-1</sup>. (D) Effect of glutamine addition on O<sub>2</sub> concentration over time. Graphical representation of O<sub>2</sub> concentration (ppb) over time (min) for cells with no glutamine added (Top line, filled dots) compared to cells after the addition of glutamine (100 μM) indicated by a blue arrow (Bottom line, empty dots). Conditions are as above. Data are averages (±SEM, at least three independent cultures). Two-way ANOVA with Sidak's multiple comparisons test (\*)  $P < 0.05$ , (\*\*)  $P < 0.01$ , (\*\*\*)  $P < 0.001$  (\*\*\*\*),  $P < 0.0001$  and n.s. not significant.

medium (Fig. 6B). The lack of any significant effect of glutamine addition on amino acid uptake (Fig. 6C) or TCA cycle activity as measured by O<sub>2</sub> consumption (Fig. 6D) supports our hypothesis of OPS1010 being nitrogen blind and points toward GAF-mediated nitrogen sensing in Rlv3841. Together, these data support the role of glutamine as the signal inhibiting both PtsP phosphorylation and subsequently, that of PtsN and ManX, down-regulating amino acid uptake and up-regulating TCA cycle activity, respectively. This PtsP-sensing mechanism would allow the cell to balance nitrogen and carbon metabolism.

**PTS Systems Are Essential for Symbiosis.** Several rhizobial PTS mutants have symbiotic phenotypes (40, 45, 55). All of the individual PTS mutants derived from Rlv3841 were able to nodulate and fix nitrogen (Fig. 7A and B), although *ptsP*, *npr*, and *ptsNIN2* mutants did this at a lower rate and the occurrence of suppressor mutations was observed at a high rate. This suggested that a single blockage through mutation of *ptsN* or *manX* does

not block nitrogen fixation. However, the double mutant *ptsNmanX* had white nodules (Fig. 7C) that fixed significantly less nitrogen (Fig. 7B). These data show that either branch of the pathway (PTS<sup>Nit</sup> or carbohydrate-PTS) is adequate for nitrogen fixation, but disrupting both at the same time blocks it.

In this work, we have determined the physiological consequences of the cross-talk between the two historical PTS branches in rhizobia. Our study provides evidence that this complex biological sensor-actuator device enables bacterial cells to posttranslationally alter physiology to balance carbon and nitrogen availability. By analyzing the PTS components of *R. leguminosarum* as a single integrated system and identifying the internal signal that activates it, we have been able to assign a role to each of the EIIA regulatory proteins to demonstrate that PTS coordinates the main metabolic cellular functions. The biological relevance of this phosphorylation switch between ManX and PtsN is the ability to balance internal fluxes of carbon and nitrogen. In our model system, the absolute absence of these two

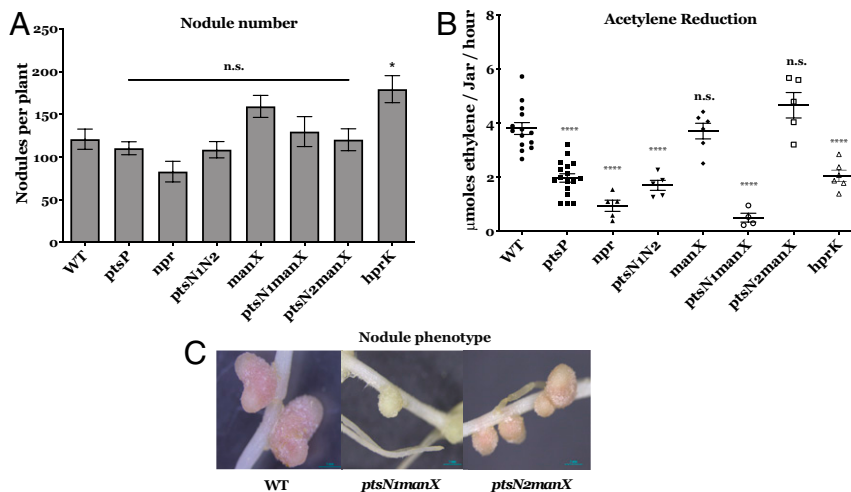


**Fig. 6.** Effect of deletion of the GAF domain on the PTS regulatory cascade. (A) Membrane transport of AIB in wild-type cells (3841 WT), *ptsP* mutant (PtsP107), and *ptsPΔGAF* mutant (OPS1010) grown on UMS with 10 mM glucose as the carbon source and 10 mM  $\text{NH}_4\text{Cl}$ . (B) Effect of nitrogen availability on AIB transport. Membrane transport of *ptsPΔGAF* mutant (OPS1010) cells grown on UMS with 10 mM glucose as carbon source and 10 mM  $\text{NH}_4\text{Cl}$ , glutamine (N-rich conditions) or glutamate (N-limiting conditions). (C) Effect of glutamine addition on AIB transport. *ptsPΔGAF* mutant (OPS1010) cells grown on UMS with 10 mM glucose as carbon source and 10 mM  $\text{NH}_4\text{Cl}$  (N-rich conditions) or 10 mM glutamate (N-limiting conditions). Transport rates measured at 1 min and 10 min after the glutamine addition. All transport rates are expressed in  $\text{nmol min}^{-1} \text{mg protein}^{-1}$ . (D) Effect of glutamine addition on  $\text{O}_2$  consumption. *ptsPΔGAF* mutant (OPS1010) cells grown on UMS with 20 mM succinate as the carbon source and 10 mM  $\text{NH}_4\text{Cl}$  (N-rich conditions) or 1 mM  $\text{NH}_4\text{Cl}$  (N-limiting conditions). Values show the rates of  $\text{O}_2$  consumption expressed in  $\mu\text{mol L}^{-1} \text{min}^{-1} \text{OD}_{600\text{nm}}^{-1}$ . Data are averages ( $\pm$ SEM,  $n = 3$  independent cultures). Statistical analyses are indicated following the level of significance, with (\*\*\*\*)  $P < 0.0001$  and n.s., not significant.

output regulators leads to an inefficient symbiosis. Whereas individual mutants are still able to fix nitrogen, potentially due to a partial affinity between EIAs, the disruption of the phosphotransfer cascade affected negatively the rates of fixation, indicating that the regulatory mechanisms of  $\text{PTS}^{\text{Ntr}}$  are essential.

## Conclusion

The PTS system has been described as an autonomous biochemical device that modulates a variety of cell functions in response to the intracellular PEP/pyruvate ratio, the N/C ratio, and the intracellular/extracellular fraction of  $\text{K}^+$  ions (56). However, despite its apparent universal importance in proteobacteria,

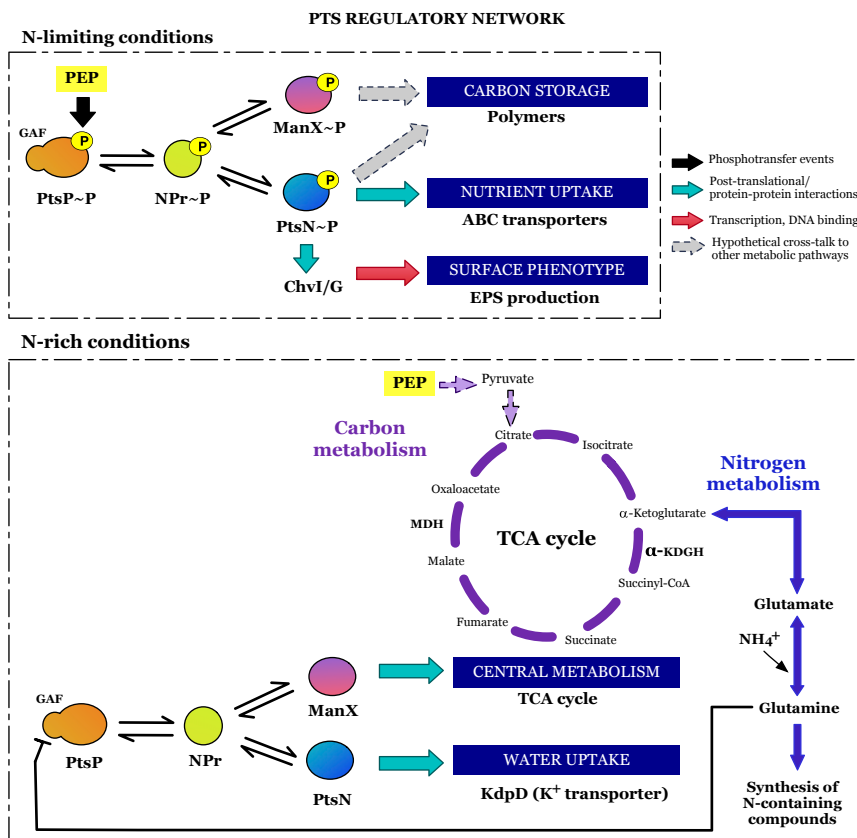


**Fig. 7.** Symbiotic phenotype of Rlv3841 PTS mutants. (A) Nodule number and (B) acetylene reduction assay carried out 21 d postinoculation (dpi) for Rlv3841 wild type, *ptsP* (PtsP107), *npr* (AA031), *ptsN1N2* (AA047), *manX* (LMB692), *ptsN1manX* (OPS0374), *ptsN2manX* (OPS0849), and *hprK* (AA081) mutants. Data are averages ( $\pm$ SEM) from at least three plants inoculated with independent cultures. Statistical analyses are indicated following the level of significance, with (\*)  $P < 0.01$ , (\*\*\*\*)  $P < 0.0001$  and n.s., not significant. (C) Pea nodules formed by wild-type Rlv3841 and double mutants *ptsN1manX* (OPS0374) and *ptsN2manX* (OPS0849). (Scale bar, 1 mm.)



it was unclear how the nitrogen and carbon balance was globally integrated. We reasoned that to understand PTS at the global level, all of the components needed to be tested in a single system. Furthermore, it was crucial to have single-copy integrated versions of PtsN and ManX that were either wild type or non-phosphorylatable and, in the case of ptsN, to have a functional phosphomimic version. It was also essential to have single chromosomal copies of full-length PtsP and Pts $\Delta$ GAF. With these strains as tools it was then possible to design real-time *in vivo* experiments to test how the components work. To illustrate our findings, we propose a model for an integrated PTS regulatory network in Fig. 8. This two-component EIIA switch modulated by NPr integrates PTS<sup>Ntr</sup> and the carbohydrate-PTS systems as a single PTS system in  $\alpha$ -proteobacteria, where glutamine is the signal binding the GAF domain of PtsP. Our results on transport and O<sub>2</sub> consumption rates for both wild type and the GAF deletion mutant (Figs. 5 and 6) point toward a model where, as seen in other  $\alpha$ -proteobacteria (51, 52), glutamine prevents PtsP phosphorylation by binding to GAF under N-rich conditions. This binding would drive a reduction in amino acid transport (due to a decrease in PtsN~P) while increasing the activity of the TCA cycle (ManX also dephosphorylated). We hypothesize that carbon catabolism increases to match the nitrogen status of the cell. By contrast, under N-limiting and carbon-rich conditions PtsP~P would increase as would PtsN~P and ManX~P. We predict that PtsN~P will increase amino acid uptake while the drop in dephosphorylated ManX will reduce the activity of the TCA cycle, increasing nitrogen acquisition

while slowing down carbon catabolism. Concomitantly, PtsN~P stimulates extracellular EPS production (Fig. 3A) but is also likely that either PtsN~P or ManX~P increase the production of intracellular carbon storage polymers such as glycogen or polyhydroxybutyrate (PHB). Effectively, when carbon cannot be catabolized because of limiting nitrogen, carbon surplus is stored as polymers. Accordingly, PTS mutants in *R. eutropha*, *P. putida*, and *Sinorhizobium fredii* that cannot phosphorylate EIIA have significantly reduced PHB levels (46, 55, 57). Therefore, this two-component EIIA switch allows bacteria to balance the availability of nitrogen and carbon skeletons and, therefore, nitrogen and carbon metabolism, by means of PTS phosphorylation (Fig. 8) and independently of the Ntr system. The speed of response observed after addition of glutamine also confirmed that the regulation exerted by PTS follows the model of allosteric binding of signaling metabolites and rapid protein post-translational modifications, allowing regulatory mechanisms to act at different timings relative to nitrogen levels. Thus, PTS constitutes another major regulatory system that is able to interact with both metabolic and transcriptional networks (50), where cross-talk between PTS systems could increase the power and connectivity of signal-response networks (58). Indeed, it was recently revealed that in *Caulobacter crescentus* and *R. eutropha* PTS interacts with the global stress stringent response (52, 59), emphasizing its profound importance to bacterial signaling and control. In *C. crescentus*, this PTS<sup>Ntr</sup>-dependent regulation of (p)ppGpp accumulation upon N-starvation is mediated by direct binding of SpoT to the



**Fig. 8.** Schematic model for PTS interactions according to nitrogen availability. Under N-limiting conditions, PtsP autophosphorylation via PEP modulates the phosphorylation of NPr and subsequently, PtsN (EIIA<sup>Ntr</sup>) and ManX (carbohydrate-EIIA). Once phosphorylated, PtsN activates ABC transporters and interacts with ChvI to control EPS production, with both phosphorylated EIIs possibly acting on carbon storage. Under N-rich conditions, intracellular glutamine inhibits PtsP autophosphorylation and, therefore, the upcoming phosphorylation of PTS components. Dephosphorylated PtsN interacts with KdpD, controlling K<sup>+</sup> homeostasis, and ManX acts on the TCA cycle. Dual control of PtsN and ManX mediated by the NPr switch is affected by nitrogen availability, balancing carbon metabolism.

phosphorylated version of PtsN interfering with its (p)ppGpp hydrolase activity, a regulatory mechanism also conserved in *S. meliloti* (60).

Given the broad conservation of this system, the reciprocal regulation of carbon and nitrogen metabolism by an EIIA switch as proposed here is likely to be another global regulatory mechanism, enabling bacterial cells to alter their physiology in real time to balance carbon and nitrogen availability. PTS and P<sub>II</sub> proteins coordinate posttranslationally many facets of bacterial metabolism by interacting with and regulating the activities of enzymes, transcription factors, and transport proteins. These regulatory mechanisms work in parallel using the glutamine/ $\alpha$ -ketoglutarate level to time their responses differentially, reinforcing the importance of coordinating nitrogen and carbon sources as essential elements for all life. Such behavior allows bacteria to efficiently adapt to nutritional adversity and reveals how versatile they are at modifying their metabolism in response to nutrient availability. An attractive line of future investigation would be to model regulatory networks, exploring how bacteria integrate PTS regulation with P<sub>II</sub> proteins, the stringent response, and Ntr regulation to unravel how nitrogen levels drive these bacterial metabolic connections.

## Methods

**Bacterial Strains and Growth Conditions.** The bacterial strains and plasmids used in this study are listed in *SI Appendix, Table S2*. *E. coli* strains were grown in liquid or solid Luria–Bertani (LB) medium (61) at 37 °C supplemented with appropriate antibiotics ( $\mu\text{g ml}^{-1}$ ): ampicillin, 100; tetracycline, 10; and kanamycin, 20. *R. leguminosarum* strains, listed in *SI Appendix, Table S2*, were grown at 28 °C in TY extract (62) or UMS (63) with appropriate carbon and nitrogen sources at 10 mM unless otherwise stated. Antibiotics were used at the following concentrations ( $\mu\text{g ml}^{-1}$ ) unless otherwise stated: gentamicin, 20; kanamycin, 20; neomycin, 40; spectinomycin, 50; streptomycin, 500; and tetracycline (2 in UMS, 5 in TY). Mean generation times (MGTs) for Rlv3841-derived strains were obtained from cells grown in 50 mL UMS with the corresponding carbon and nitrogen sources at an initial optical density (OD<sub>600nm</sub>) of 0.01. OD<sub>600nm</sub> measurements were taken at 4-h time intervals until growth reached stationary phase from at least three biological replicates. The MGT of Rlv strains was calculated as the number of hours it takes the population to double while in exponential growth phase.

**Mutant and Plasmid Construction.** All routine DNA analyses were done using standard protocols (61). The cloning details are described in *SI Appendix*. Primers used in this work are listed in *SI Appendix, Table S3*. Conjugations and transductions with bacteriophage RL38 were performed as previously described (64, 65).

**Transport Assays.** Transport assays were performed with 25  $\mu\text{M}$  (4.625 kBq) of <sup>14</sup>C-labeled solute (53, 66), with cultures grown in UMS with 10 mM glucose and 10 mM NH<sub>4</sub>Cl unless otherwise specified. Glutamine and glutamate addition experiments were done by adding 5  $\mu\text{L}$  of a 100  $\mu\text{M}$  stock to 5 mL of starved cells. The same procedure was followed by adding 5  $\mu\text{L}$  of rhizobium minimal salts (RMS) medium as control. At two different time points after their addition, 1 min and 10 min, 2 mL of cells were filtered using a Millipore filtration unit with vacuum suction, washed twice, and resuspended in RMS with no carbon or nitrogen sources for subsequent transport assays.

**Exopolysaccharide Measurements.** The 250-mL cultures were grown up to an OD<sub>600nm</sub> of 0.5. Cells were spun down, dried, and weighted. The supernatant was treated with two volumes (500 mL) of cold isopropanol. After precipitation, the EPS was spun down and the pellet dried at 37 °C. Values are expressed in mg/mg dry weight of cells.

**Promoter Analysis.** Promoter analyses with *lux* fusions were done by growing rhizobial strains on UMS agar slopes or liquid cultures (with appropriate

antibiotics) following the protocol by Pini et al. (67). Sensitivity of each promoter fusion was measured either with the NightOWL in Vivo Imaging System (Berthold Technologies) with agar plates or the Promega GloMax Microplate Reader with liquid cultures.  $\beta$ -Galactosidase fusions were assayed according to Miller (68), with modifications as described by Poole et al. (69).

**O<sub>2</sub> Consumption Assays.** O<sub>2</sub> consumption rates were obtained from UMS cultures grown with 10 mM glucose and 10 mM NH<sub>4</sub>Cl unless otherwise specified to early exponential phase. A 25-mL sealed glass universal, containing an OxyDot was quickly filled completely with the liquid culture, and O<sub>2</sub> measurements were taken every 15 s for 1,000 s, or until the O<sub>2</sub> level decreased below 1%. These noninvasive measurements were performed with the O<sub>2</sub> electrode OxySense 3251 System and the data analyzed with the OxySense Gen III software. O<sub>2</sub> consumption rates were calculated as the time it takes the population to decrease the O<sub>2</sub> concentration by 3%. Glutamine and glutamate addition experiments were done by measuring O<sub>2</sub> consumption after injecting 250  $\mu\text{L}$  of a 100-mM stock into the sealed universal. The same procedure was followed by adding 250  $\mu\text{L}$  of water as control.

**Enzyme Assays.** Cultures of *R. leguminosarum* strains were harvested at exponential growth, washed, and resuspended in 10 mL 40 mM Hepes pH 7, containing 1 mM dithiothreitol. Cells were disrupted by two passages on a FastPrep-24 5G ribolyser (MP Biomedicals), following centrifugation. Oxoglutarate dehydrogenase (EC 1.2.4.2) was assayed according to Reeves et al. (70) and malate dehydrogenase (EC 1.1.1.37) by the technique of Saroso et al. (71). The protein concentration of whole cells was determined by the method of Lowry et al. (72), using bovine serum albumin as standard.

**BACTH Assays.** Interacting partners on the high-copy pUT18C vectors were transformed into MAE01 ( $\Delta\text{cyaA}::\text{Apra}^R$ ) cells. Single transformants were grown at 37 °C in 5 mL LB to an OD<sub>600nm</sub> of 0.3 to 0.5 and subsequently transformed with the interaction partners on the low-copy pKNT25 vector, plated onto LB plates with 0.5 mM isopropyl- $\beta$ -D-1-thiogalactopyranosid (IPTG) and X-gal, and incubated for 2 d at 28 °C. To quantify BACTH interactions, these colonies were grown overnight at 28 °C in 10 mL LB with ampicillin and kanamycin at standard concentrations (73). Overnight cultures were then used to inoculate 5-mL cultures of LB with 1% wt/vol glucose, ampicillin, and kanamycin to an initial OD<sub>600nm</sub> of 0.1 and grown until they reached an OD<sub>600nm</sub> of 0.4. Cultures were then induced with 2 mM IPTG and grown for 6 h at 28 °C. A standard  $\beta$ -galactosidase assay (68) was used to quantify the interaction between proteins. KdpD/PtsN1 pairs were used as positive controls (23), while empty vectors, as negative controls.

**Plant Growth and Acetylene Reduction.** *Pisum sativum* cv. Avola seeds were surface sterilized using 95% ethanol and 2% sodium hypochlorite at the time of sowing. Plants were inoculated with  $1 \times 10^7$  cells of the appropriate rhizobial strain and grown in 1-L beakers filled with sterile medium-grade vermiculite and N-free nutrient solution as previously described (69) in a growth room (16 h light/8 h dark). They were harvested at pea flowering (3 wk) and acetylene reduction was determined as previously described (74).

**Statistical Analysis.** All analyses were performed using GraphPad Prism 8 (GraphPad Software). Significant differences between pairs of parameters were determined by Student's *t* tests. Comparisons of more than two groups were done by ANOVA followed by multiple comparisons post hoc corrections as indicated in each figure legend. A *P* value of less than 0.05 was considered as statistically significant.

**Data Availability.** The data supporting the findings of the study are available in this article and its *SI Appendix*.

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