Modulation of Quorum Sensing as an Adaptation to Nodule Cell Infection during Experimental Evolution of Legume Symbionts

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ABSTRACT
Over millions of years, changes have occurred in regulatory circuitries in response to genome reorganization and/or persistent changes in environmental conditions. How bacteria optimize regulatory circuitries is crucial to understand bacterial adaptation. Here, we analyzed the experimental evolution of the plant pathogen Ralstonia solanacearum into legume symbionts after the transfer of a natural plasmid encoding the essential mutualistic genes. We showed that the Phc quorum sensing system required for the virulence of the ancestral bacterium was reconfigured to improve intracellular infection of root nodules induced by evolved Ralstonia. A single mutation in either the PhcB autoinducer synthase or the PhcQ regulator of the sensory cascade tuned the kinetics of activation of the central regulator PhcA in response to cell density so that the minimal stimulatory concentration of autoinducers needed for a given response was increased. Yet, a change in the expression of a PhcA target gene was observed in infection threads progressing in root hairs, suggesting early programming for the late accommodation of bacteria in nodule cells. Moreover, this delayed switch to the quorum sensing mode decreased the pathogenicity of the ancestral strain, illustrating the functional plasticity of regulatory systems and showing how a small modulation in signal response can produce drastic changes in bacterial lifestyle.

IMPORTANCE
Rhizobia are soil bacteria from unrelated genera able to form a mutualistic relationship with legumes. Bacteria induce the formation of root nodules, invade nodule cells, and fix nitrogen to the benefit of the plant. Rhizobial lineages emerged from the horizontal transfer of essential symbiotic genes followed by genome remodeling to activate and/or optimize the acquired symbiotic potential. This evolutionary scenario was replayed in a laboratory evolution experiment in which the plant pathogen Ralstonia solanacearum successively evolved the capacities to nodulate Mimosa pudica and poorly invade, then massively invade, nodule cells. In some lines, the improvement of intracellular infection was achieved by mutations modulating a quorum sensing regulatory system of the ancestral strain. This modulation that affects the activity of a central regulator during the earliest stages of symbiosis has a huge impact on late stages of symbiosis. This work showed that regulatory rewiring is the main driver of this pathogeny-symbiosis transition.

KEYWORDS
symbiosis, rhizobium, experimental evolution, infection, quorum sensing, regulation

Most bacterial cellular functions are regulated by complex multilayered genetic regulatory networks that include environmental and internal sensors and a wide variety of signaling pathways. Fine-tuned regulation allows rapid shifts in metabolism,
physiology, and behavior in response to environmental fluctuations (1–3). Over millions of years, the regulation circuits have evolved to adapt to genome reorganization and/or persistent changes in environmental conditions (4). Deciphering how bacteria optimize regulatory circuitries is crucial to understand bacterial adaptation and the biosphere.

Quorum sensing (QS) systems are key components of regulatory networks in bacteria which allow the regulation of gene expression in a population-dependent manner. QS involves the production of extracellular signals, called autoinducers, whose concentration is a function of cell population density. Above a system-specific threshold level, these molecules rapidly activate or repress the transcription of hundreds of genes, thus modifying and synchronizing bacterial behavior on a population-wide scale. These signal response systems are widespread in both Gram-positive and Gram-negative bacteria, where extensive variations regarding the type of autoinducers and cognate receptors, signal transduction mechanisms, and associated cellular responses have been described (5). Biological functions regulated by QS are numerous, including bioluminescence, virulence, symbiosis, motility, biofilm formation, or exopolysaccharide (EPS) production (6–10). The plant pathogen *Ralstonia solanacearum* possesses a virulence-related QS system, the Phc system, which controls the activity of the LysR-type transcription regulator PhcA (11). One component of this system is PhcB, a protein harboring methyltransferase activity essential for the synthesis of autoinducers, either (R)-3-hydroxypalmitic acid methyl ester [(R)-3-OH PAME] or (R)-3-hydroxymyristic acid methyl ester [(R)-3-OH MAME], depending on the *R. solanacearum* strains. In strain GMI1000, only (R)-3-OH MAME was detected (12). How this signal is transduced to PhcA has not been fully elucidated. At low cell density, the unphosphorylated two-component system PhcSR is thought to inhibit the activity of the central virulence regulator PhcA via an unknown mechanism. When the QS molecules reach the minimal stimulatory concentration (around 10 nM, matching the cell density of 10^7 CFU/ml in liquid culture), they likely trigger the autophosphorylation of the sensor histidine kinase PhcS, which, in turn, phosphorylates the response regulator PhcR (13), and this releases the inhibition of PhcA (14). The *phcQ* gene, which is likely cotranscribed with *phcBSR*, encodes a protein composed of a receiver domain similar to the receiver domain of PhcR (see Fig. S1A in the supplemental material). The role of this protein in the QS-dependent signaling cascade is not known (14). Activation of PhcA in *R. solanacearum* allows the switch from metabolic versatile cells to nonmotile EPS-producing virulent cells (15). Three independent transcriptomic studies conducted under three different experimental conditions (minimal medium, rich medium, and *in planta*) indeed identified hundreds of genes under the positive or negative control of PhcA (16–18).

Rhizobia, the nitrogen-fixing symbionts of legumes, are an excellent biological system to investigate how regulatory circuits rewire in response to genome reorganization and colonization of new niches. Rhizobia are soil bacteria able to induce the formation of root nodules where internalized bacteria fix nitrogen for the benefit of the plant (19). Extant rhizobia belong to hundreds of species distributed in 18 different genera of alpha- and betaproteobacteria (20–22). They have evolved through the horizontal acquisition of essential symbiotic genes, the nodulation (*nod*) and nitrogen fixing (*nif-fix*) genes, likely followed by subsequent adaptation of the recipient genome to the new legume endospheric environment (20). To understand posthorizontal gene transfer adaptation, we previously experimentally evolved *Ralstonia solanacearum* into legume symbionts by mimicking the natural evolutionary scenario. The symbiotic plasmid pRalta of the rhizobium *Cupriavidus taiwanensis* LMG19424 harboring the *nod* and *nif-fix* genes (23) was transferred to *R. solanacearum* strain GMI1000, and the resulting chimera was then evolved in parallel lineages using serial plant-bacteria cocultures. While the ancestral strain was strictly extracellular and unable to form root nodules, most lines sequentially acquired the capacity to form root nodules, poorly infect nodules, and massively infect nodule cells (24, 25). Acquisition of nodulation relied on the inactivation of the *R. solanacearum* pathogenic type III secretion system (T3SS) (24). The first level of infection was gained via inactivation of virulence regula-

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tors, either HrpG or VsrA (24, 26). In some lines, infection was optimized via modifications in the EfpR regulatory pathway (27). EfpR was shown to be a global virulence regulator and a metabolic repressor (27, 28). In other very infectious lines, no mutation was detected in known components of the EfpR pathway, suggesting that evolution targeted either unknown components of this path or another pathway.

Here, we show that in some of these lines, massive intracellular accommodation was reached via modifications in the R. solanacearum quorum sensing system. Mutations in either phcB or phcQ allowed improved infection via a fine modulation of the sensory pathway that delays to different degrees the cell density-dependent activation of the central regulator PhcA. While the highly similar gene expression profiles of efpR and phcA mutants suggested a connection between the two pathways, present results indicated that EfpR interferes with the PhcA regulon independently from the PhcBSRQ sensory cascade. Both efpR and phc late infection-adaptive mutations transiently affect the expression of PhcA target genes during early symbiotic stages corresponding to the entry and progression of bacteria in root hairs. EfpR and PhcA may thus control the expression of a common set of genes whose deregulation helps change the lifestyle of the bacterium, improving its symbiotic infection capacity while decreasing its pathogenicity.

**RESULTS**

Infection-adaptive mutations affect the phc system in E, K, and M lineages. Our previous work showed that the evolved clones E16, K16, and M16 bore adaptive mutations in genes encoding components of the Ralstonia solanacearum Phc quorum sensing system. Nonsynonymous mutations in the phcB (phcBR22C), phcQ (phcQR154C), and phcS (phcSL161R) genes (Table 1) improved the in planta fitness (evaluated by the number of bacteria in nodule populations) of the evolved clones compared to that of the derivatives harboring the wild-type alleles (29). Genome resequencing or PCR analysis of intermediate clones along the E, K, and M lineages allowed the detection of the phcBR22C mutation in half clones from cycle 10 in the E lineage and the detection of phcQ R154C and phcS L161R mutations in all clones from cycles 13 and 2 in the K and M lineages, respectively (Fig. 1). This suggested that these mutations have fixed rapidly in the K and M lineages and less rapidly in the E lineage.

To further determine whether these mutations improve nodule cell infection, we inoculated Mimosa pudica individually with E16, K16, M5, their derivatives harboring the wild-type (WT) phc alleles, their respective nodulating ancestors CBM212, CBM349, and CBM356, and mutants of these ancestors carrying the phc mutations (M5, which is statistically as infectious as M16 [26], was used rather than M16, since genetic transformation failed in the latter clone in spite of many trials). We measured the size of the intracellularly infected and necrotic zones, identified as light-brown and dark-brown zones, respectively (25, 30), in sections of nodules collected at 21 days postinoculation (dpi). These two parameters were used as indicators of the quantity and quality of infection. The intracellular infection capacity of E16, CBM212 phcBR22C, K16, and CBM349 phcQR154C increased significantly compared to that of E16 phcBW, CBM212, K16 phcQWT, and CBM349, respectively (Fig. 2A). At the same time, the capacity to induce necrosis decreased in the strains bearing the mutated alleles (Fig. 2B). Reconstructing the phcB and phcQ mutations in the chimeric Ralstonia strain GMI1000(pRalta) harboring an hrpG mutation (CBM1627), conferring nodulation and partial infection ability, provided this strain a capacity to infect nodule cells and induce necrosis similarly.

**TABLE 1 Validated infection-adaptive mutations in E, K, and M lineages**

| Strain(s) | Gene | Product | Mutation | Position on the chromosome | Protein modification |
|-----------|------|---------|----------|---------------------------|---------------------|
| E16       | RSc2735, phcB | Regulatory protein, SAM-dependent methyltransferase domain | C/T | 2943477 | R22C |
| K16       | RSc2738, phcQ | Response regulator receiver, CheY family | C/T | 2947813 | R154C |
| M5, M16   | RSc2736, phcS | Sensor protein histidine kinase, repressor of PhcA | T/G | 2945312 | L161R |

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to that of the evolved E16 or K16 clones, respectively, indicating that these mutations are the major infection-adaptive ones in E and K lines (Fig. 2).

The effect of the \textit{phcS} L161R mutation depended on the genetic background (Fig. 2). It improved intracellular infection and reduced necrosis in GMI1000(pRalta) hrpG but failed to do so in \textit{hrcV}-mutated backgrounds [M5, CBM356, and GMI1000(pRalta) \textit{hrcV}]. This result is not necessarily contradictory to the previously observed increase in relative \textit{in planta} fitness of CBM356 \textit{phcSL161R} and M5 compared to that of CBM356 and M5 \textit{phcSWT} (29), which could be due to an increase in nodulation competitiveness or extracellular infection rather than an increase in intracellular infection \textit{per se}.

The reconstruction of the \textit{phcBQS} mutations in GMI1000(pRalta) did not allow the original chimeric strain to nodulate \textit{M. pudica}, indicating that the effect of these mutations is conditional on the presence of the \textit{hrpG} stop mutation.

Altogether, this showed that the \textit{phcBR22C} and \textit{phcQR154C}, but not \textit{phcSL161R}, were the main infection-enhancing mutations in their respective lineages. We thus further analyzed the effect of the \textit{phcB} and \textit{phcQ} mutations on the Phc quorum sensing system.

\textbf{Infection-adaptive \textit{phcBQ} mutations do not constitutively inactivate the PhcA transcription regulator.} The cell density-responsive regulatory system PhcBSRQ controls the PhcA central regulator (11). To evaluate whether the infection-adaptive \textit{phc} mutations led to the inactivation of the Phc quorum sensing system, we evaluated the activity of PhcA in wild-type and mutated strains. For that, we constructed a plasmidic fusion between the promoter region of \textit{xpsR}, a direct transcriptional target of PhcA (31), and the \textit{Escherichia coli} reporter gene \textit{lacz} encoding \(\beta\)-galactosidase. We introduced this plasmid in GMI1000(pRalta) \textit{hrpG}, in its \textit{phcBR22C}, \textit{phcQR154C} single nucleotide polymorphism (SNP) derivative mutants, and in its \textit{phcB}-(in frame), \textit{phcQ}-, and \textit{phcA}-deleted mutants. Since \textit{xpsR} expression is under positive control of PhcA, the dosage of \(\beta\)-galactosidase activity positively correlates with PhcA activity.

The \(\beta\)-galactosidase activity was determined from mid-exponentially grown liquid cultures of each strain in rich medium. In agreement with the literature (31), deletion of \textit{phcA} completely abolished \textit{xpsR-lacz} expression (Fig. 3). In contrast, the expression of \textit{xpsR-lacz} in the \textit{phcB} SNP mutants was high and similar to that of strain GMI1000(pRalta) \textit{hrpG}, and the \textit{phcQ} SNP mutant only exhibited a lower level of reporter gene expression (Fig. 3). The \textit{phcB} SNP mutant was clearly different from a \textit{phcB} in-frame deletion mutant, indicating that the mutation did not inactivate the
FIG 2 Quantification of the relative infected and necrotic areas of nodules induced by evolved clones and mutants. Distributions of the percentages of infected areas (A) and necrotic areas (B) per nodule section recovered at 21 dpi. The central rectangles span the first quartiles (Continued on next page)
protein. In contrast, the \textit{phcQ} deletion mutant exhibited a similar level of \textit{xpsR-lacZ} reporter gene expression as the \textit{phcQ}R154C mutant. Surprisingly, the three Δ\textit{phcA}, Δ\textit{phcB} in frame, and Δ\textit{phcQ} mutants induced on \textit{M. pudica} the formation of nodules with infection and necrosis zones similar to that of nodules induced by the \textit{phcBQ} SNP mutants (around 25% to 30% of the nodule section) (Fig. 2). Moreover, the \textit{phcB} and \textit{phcQ} SNP mutants display either no or a minor difference in relative \textit{in planta} fitness from that of a Δ\textit{phcA} mutant, indicating that a \textit{phcA} deletion is almost as adaptive as the \textit{phcBR22C} and \textit{phcQR154C} mutations for symbiosis (Fig. 4). This suggested that the point mutations in \textit{phcB} and \textit{phcQ} affect respective protein functions, resulting in an altered PhcA activity in symbiosis.

\textit{Infection-adaptive phc mutations decrease responsiveness to cell density}. The \textit{phcBR22C} and \textit{phcQR154C} mutations are not located in the known functional domains of the respective proteins (see Fig. S1A in the supplemental material). The R22C mutation does not map to the S-adenosyl-L-methionine-dependent methyltransferase domain of PhcB nor does the R154C mutation map to the receiver domain of PhcQ. Yet, mutations alter conserved amino acids of \textit{Ralstonia}, \textit{Cupriavidus}, and \textit{Paraburkholderia} homologous proteins and would influence protein function as suggested by the Phyre2 protein structure prediction server (Fig. S1B and C) (32).

We hypothesized that these mutations affect protein conformation, stability, or efficiency and thus may alter the dynamics of PhcA responsiveness to cell density. To test this hypothesis, we monitored the expression kinetics of the \textit{xpsR-lacZ} reporter gene fusion in various genetic backgrounds along with growth in rich liquid medium (Fig. 5A). In the nodulating chimeric strain GMI1000(pRalta) \textit{hrpG}, the expression of \textit{xpsR-lacZ} sharply rose at an optical density at 600 nm (OD$_{600}$) of 0.04 (corresponding to ca. $5 \times 10^7$ CFU/ml) (see Fig. S2), reached a peak at an OD$_{600}$ of 0.12 (ca. $10^8$ CFU/ml),

\begin{figure}
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\includegraphics[width=\textwidth]{figure3}
\caption{Expression of the \textit{xpsR-lacZ} plasmidic fusion in the chimeric \textit{R. solanacearum} nodulating strain and derivative mutants at mid-exponential phase. β-Galactosidase activities were measured on bacteria grown to mid-exponential phase (OD$_{600}$ was around 0.2) in rich BG medium. Data are from 3 independent experiments. Values correspond to means \pm standard deviations. *$P < 0.05$; **$P < 0.001$ versus the wild-type strain (Student’s \textit{t} test).}
\end{figure}

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\caption{Legend (Continued)}
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and then decreased and stabilized at an OD$_{600}$ of 0.2 (ca. 2 x 10$^8$ CFU/ml). No peak of expression was observed in either the $phcB$ or $phcQ$ SNP mutant. In the $phcB$R22C mutant, the expression was slightly delayed and reached a maximum at an OD$_{600}$ of 0.2 (ca. 2 x 10$^8$ CFU/ml). We confirmed that the $phcB$R22C mutant does not behave like a $phcB$ in-frame deletion mutant, which exhibited no PhcA activity regardless of the cell density. In contrast, the $phcQ$R154C mutant closely resembled a $phcQ$ deletion mutant. In both mutants, the induction of $xpsR$-$lacZ$ was slow and linear until an OD$_{600}$ of 1 (ca. 10$^9$ CFU/ml). These dynamics of $xpsR$ expression suggested that PhcQ may be involved in a positive feedback activation loop of PhcA, implying that the $phcQ$ mutation may affect the synthesis of autoinducing molecules.

To substantiate whether the expression of other PhcA target genes was affected by the $phcB$ and $phcQ$ SNP mutations, we checked by reverse transcription-quantitative PCR (qRT-PCR) the expression of seven genes known to be positively or negatively controlled by PhcA ($egl$, $flhF$, $motA$, Rsc1817, Rsp0983, Rsp0178, and $xpsR$) (16–18) (see Fig. S3). For this, we harvested cells grown in rich liquid medium to an OD$_{600}$ of around 0.1, a cell density at which the expression of $xpsR$ was lower in the $phcB$ and $phcQ$ SNP mutants than in the wild-type strain. All these genes were differentially expressed in the $phcB$R22C, $phcQR154C$, and $ΔphcA$ mutants compared to that in the GMI1000(pRalta) $hrpG$ wild-type strain. Consistently with the $xpsR$ expression data, expressions of these genes were less affected in the $phcB$R22C mutant than in the $phcQ$R154C mutant.

These results showed that the $phcB$ and $phcQ$ mutations delay, to different degrees, the timing of Phc quorum sensing regulation, likely by decreasing the activity of the corresponding proteins.

The $phcB$R22C and $phcQR154C$ mutations affect the synthesis of QS molecules. To evaluate the efficiency of $phcB$ and $phcQ$ SNP mutants to produce autoinducing molecules, we set up a bioassay in which a strain deficient for the synthesis of the QS molecules [i.e., a $phcB$ in-frame deletion mutant in the GMI1000(pRalta) $hrpG$ background] carrying the reporter plasmidic $xpsR$-$lacZ$ fusion was incubated with supernatants of cultures harvested at an OD$_{600}$ of 0.1 from either the GMI1000(pRalta) $hrpG$ wild-type strain or the $phcB$R22C, $phcQR154C$, $ΔphcB_{in-frame}$, $ΔphcQ$, or $ΔphcA$ derivative mutants. As expected, after 4 h of incubation with the supernatant of the wild-type strain, the $xpsR$-$lacZ$ fusion was well expressed, while this fusion was not expressed after incubation with the supernatant of a $ΔphcB_{in-frame}$ mutant. In agreement with previous works showing that some PhcA targets positively control the production or the stability of 3-OH MAME (33), the $xpsR$-$lacZ$ fusion was less expressed after incubation with the supernatant of a $ΔphcA$ mutant. Interestingly, the expression of $xpsR$-$lacZ$ was strongly
FIG 5  Kinetics of the ppsR-lacZ expression in the chimeric R. solanacearum nodulating strain (GMI1000(pRalta) hrg) and derivative mutants according to the OD600. β-Galactosidase activities were measured on bacteria grown in rich BG medium. Tendency curves with their 95% confidence intervals were calculated using local polynomial regressions (locally estimated scatterplot smoothing [LOESS] method). Data were from 4 to 12 independent experiments. The same set of data for efpRE66K was used in panels A and B.
Reduced or near the background level after incubation with the supernatants from the \textit{phcB} \textit{R22C}, \textit{phcQ} \textit{R154C}, and \textit{ΔphcQ} mutants (Fig. 6), indicating that the amount of active QS molecules present in the supernatants of these mutants was reduced compared to that in the supernatant of the wild-type strain taken at the same OD$_{600}$. Moreover, we tested the involvement of PhcQ in signal transduction by incubating the double \textit{phcQ} \textit{R154C} Δ\textit{phcB} in frame or Δ\textit{phcQ} Δ\textit{phcB} in frame mutants with the supernatant of a wild-type strain. We found that the \textit{xpsR-lacZ} expression was not affected in these strains, indicating that PhcQ is involved in the production of QS molecules but not in the signal transduction to PhcA (Fig. 6).

\textbf{Infection-adaptive phc mutations alter PhcA activation during the uptake of bacteria in root hairs.} In \textit{M. pudica}, rhizobial partners enter roots via transcellular infection threads that progress toward the cortex and ultimately release bacteria within cells of the developing nodules, where they are subsequently accommodated in membrane-enclosed compartments called symbiosomes. In the evolved \textit{Ralstonia-M. pudica} interaction, intercellular bacteria were also observed associated with necrotic zones that usually occur in nodules induced by GMI1000(pRalta) \textit{hrpG} and occasionally in nodules induced by the \textit{phcB} and \textit{phcQ} SNP mutants. To know whether the delay in quorum sensing responsiveness also operates under symbiotic conditions, we assessed the expression of the \textit{xpsR-lacZ} fusion at three distinct symbiotic stages, following inoculation with either GMI1000(pRalta) \textit{hrpG} or its \textit{phcB} \textit{R22C}, \textit{phcQ} \textit{R154C}, Δ\textit{phcB} \textit{in frame} Δ\textit{phcQ}, or Δ\textit{phcA} derivative mutants (i) in root hairs during bacterial uptake, (ii) in young nodules when bacteria are released from infection threads into the cytoplasm of plant cells, and (iii) in mature nodules accommodating intracellular bacteria, called bacteroids. As expected, in the \textit{phcA}-deleted mutant, no expression of \textit{xpsR-lacZ} was observed under any symbiotic conditions (Fig. 7F). Whatever the inoculated strain, no \textit{xpsR} expression was observed in bacteroids. However, as the number of infected cells was very low in nodules induced by the GMI1000(pRalta) \textit{hrpG} strain, it was difficult to evaluate the level of \textit{xpsR} expression in bacteroids formed by this strain. In nodules induced by the...
FIG 7 Expression of pxpsR-lacZ in the chimeric R. solanacearum nodulating strain [GMI1000(pRalta) hrpG] and derivative mutants during symbiotic interaction with Mimosa pudica. Roots and nodules were (Continued on next page)
poorly infecting strain GMI1000(pRalta) hrpG, xpsR-lacZ expression was visualized in root hair and nodule infection threads (ITs) as well as in intercellular bacteria (Fig. 7A). When roots were inoculated by any of the two phcB and phcQ SNP mutants, loss of xpsR-lacZ expression was only observed in ITs progressing in root hairs, xpsR-lacZ being normally expressed in nodule ITs (Fig. 7B and C). These results suggested that activation of PhcA in the late infection-adapted phc mutants was impaired during a very early symbiotic stage, corresponding to the bacterial uptake in root hairs and initiation of infection threads. This early stage thus appeared crucial for late intracellular accommodation of bacteria in nodules.

**QS modulation as an alternative adaptive path to EfpR inactivation.** The phc mutations have the same impact on intracellular infection as mutations in the EfpR path previously detected in other evolved lines. Indeed, the size of infected and necrotic zones in nodules induced by phc SNP mutants was similar to that of nodules induced by efpR mutants (Fig. 2) (27). To compare the adaptiveness of the efpR and phc SNP mutations in a global and more integrative way, we measured their relative in planta fitness following plant coinoculation. We showed that the in planta fitness of the efpRE66K mutant was not significantly different from that of the phcBR22C or phcQ154C mutant in a GMI1000(pRalta) hrpG context (Fig. 4). Moreover, the double phcBR22C-efpRE66K or phcQ154C-efpRE66K mutant was as fit as the single efpRE66K mutant, indicating that efpR and phc adaptive mutations do not have cumulative effects (Fig. 4). Similar to that in phc SNP mutants, in an efpRE66K mutant, the expression of xpsR was strongly affected but not abolished under in vitro conditions (Fig. 5A), while under in planta conditions, expression of the xpsR-lacZ fusion was visualized in nodule infection threads but not in root hair infection threads (Fig. 7G1).

In a previous study, we determined the EfpR regulon in a GMI1000(pRalta) hrpG context (27). Strikingly, this EfpR regulon largely overlaps with the PhcA regulon determined recently in three independent studies (16–18). Among the 200 putative efpR targets, 160 (80%) were also found differentially expressed in at least one of the PhcA regulons (see Table S2). These genes include 17 for EPS synthesis, 52 for motility, 14 for hemin/siderophore transport and metabolism genes, the prhI sigma factor, and 10 genes encoding Hrp and T3 effector proteins. In addition, both PhcA and EfpR were shown to be metabolic repressors (15, 17, 27, 28). The metabolic profiles of the two mutants also display high similarities, since 88% of substrates found to be more efficiently metabolized in GMI1000(pRalta) hrpG efpRE66K (27) were also better metabolized in a phcA mutant (17) (see Table S3). Hence, the EfpR path may either affect PhcA activity directly, interfere with its activating pathway, PhcBSRQ, or act on some genes commonly targeted by PhcA.

To assess whether the EfpR pathway could interfere with the QS sensory cascade PhcBSRQ, we first determined the production of QS molecules in an efpRE66K mutant using the bioassay described above. We showed that the efpRE66K mutant produced similar amounts of active QS molecules as the wild-type GMI1000(pRalta) hrpG strain (Fig. 6). Then, we combined a phcB polar deletion mutation, which downregulated the expression of the full phcBSRQ operon (see Fig. S4), with an efpRE66K mutation in the GMI1000(pRalta) hrpG strain and monitored the expression of the xpsR-lacZ reporter gene fusion by β-galactosidase assay along growth as described above. In the single

**FIG 7 Legend (Continued)**

harvested at 6 and 7 or 14 dpi, respectively, and fixed with 2% glutaraldehyde. Roots and 55±um sections of nodules were then stained in the presence of X-Gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) to reveal beta-galactosidase activity. The xpsR-lacZ expression was detected in infection threads and extracellularly invaded spaces of young and mature nodules induced by all strains (A2 to C2, E2, G2 to I2, A3 to C3, G3 to I3) except by the phcA and phchΔinframe deletion mutants (D and F). Only the poorly infectious strains, GMI1000(pRalta) hrpG and its derivatives ΔphcBΔpolar and ΔphcBpolar-efpRE66K mutants, expressed the fusion in root hairs (A1, H1, and I1 versus B1 to G1). The phcBR22C and phcQ154C mutants constitutively expressed the mCherry fluorophore. Bacteria fluorescing in red can be visualized in ITs (B and C). Black arrows indicate infection threads. White arrows indicate extracellular bacteria. Light brown cells and black stars indicate intracellularly infected nodule cells. Images are representative of data obtained from at least 3 independent experiments.
ΔphcB polar deletion mutant, probably because of the absence of the PhcSR repressive action (14), PhcA was constitutively active at an intermediary level throughout growth, although with some variations (Fig. 5B). Interestingly, the xpsR-lacZ expression in the double ΔphcB polar-efpR E66K mutant was always lower than in the single ΔphcB polar mutant, indicating that the efpRE66K mutation can repress xpsR in the absence of the PhcBSRQ proteins and thus may alter the PhcA regulon in a PhcBSRQ-independent manner. Notwithstanding, when the OD600 was below 0.1, xpsR-lacZ expression was higher in the double ΔphcB polar-efpR E66K mutant than in the single efpR E66K mutant, suggesting an additive effect of the two mutations at low cell density. Consistently, both ΔphcB polar and ΔphcB polar-efpRE66K mutants were poorly infectious (Fig. 2) and harbored a detectable xpsR-lacZ expression in epidermal ITs (Fig. 7H and I), thus reinforcing the correlation between the absence of early xpsR expression in root hair ITs and a nice late intracellular infection phenotype.

**xpsR deletion did not improve intracellular infection.** To assess whether the downregulation of xpsR itself could solely account for the infection phenotype of the efpR and phc mutants, we constructed a xpsR deletion mutant in the GMI1000(pRalta) hrpG background and quantified its nodule cell infection ability. This mutation did not improve the intracellular infection (Fig. 2), suggesting that either EPS biosynthesis is not involved in infection or repression of EPS biosynthesis alone is not sufficient to improve infection.

**Quorum sensing modulation decreased the pathogenicity of R. solanacearum.** To evaluate whether the phc mutations adaptive for *M. pudica* nodule cell infection affect the pathogenicity of *R. solanacearum*, we introduced the phcBR22C, phcQR154C, ΔphcB_in_frame, ΔphcQ, and ΔphcA derivative mutants. *, the survival curves are significantly different from the wild-type curve; #, the survival curves are significantly different from the corresponding phc mutant curves ($P < 0.001$, Gehan-Breslow-Wilcoxon test). Values were obtained from 3 independent series of 16 plants.

**DISCUSSION**

Quorum sensing is a widespread bacterial regulatory system allowing rapid shifts in behavior in response to changes in cell density or confinement (34). The mode and tempo of signal transduction and target outputs of each quorum sensing system reflect
the unique biology carried out by a particular bacterial species. Here, we show that a system optimized to allow survival in a specialized niche can be rapidly redesigned to promote thriving in a very different environment, i.e., change in QS responsiveness helps the extracellular plant pathogen *R. solanacearum* having acquired a set of essential symbiotic genes to turn into an intracellular legume symbiont. During experimental evolution under legume (*M. pudica*) selection pressure, point mutations affecting two components of the Phc QS regulatory cascade, the autoinducer synthase PhcB and the regulatory protein PhcQ, modified the kinetics of activation of the downstream central regulator PhcA. The delay in PhcA activation in the *Ralstonia phc*-mutated evolved clones observed in vitro results in an inactivation of PhcA in epidermal ITs, which correlates with their nice nodule intracellular infection phenotypes on *M. pudica*. Conversely, these mutations, when introduced in the *Ralstonia* ancestor, reduced the virulence capacity of the strain, reemphasizing the antagonism between pathogeny and symbiosis (24, 26, 27). Unlike other infection-adaptive mutations we previously identified (*hrpG* and *efpR*) (24, 26, 27), the virulence regulator PhcA was not constitutively inactivated. Potentially, this conditional inactivation could be more favorable for subsequent adaptation steps. Interestingly, in *Cupriavidus taiwanensis*, a *Ralstonia*-neighboring taxon that evolved into *Mimosa* symbionts, the *phcB* and *phcS* orthologous genes were found under positive selection and associated with the transition to symbiosis during the natural evolution of this rhizobium species (29), likely reflecting the need to adjust QS to the legume endosymbiotic niche. Adaptive mutations altering QS responsiveness were also evidenced during experimental and natural evolution of other pathogenic and symbiotic bacteria such as *Vibrio fischeri* and *Staphylococcus aureus* (35–38), exemplifying the role of these sensory systems in bacterial adaptation and phenotypic diversification. The role of QS in rhizobial symbiosis is, however, variable, either neutral (39), positive (40–42), or negative (43, 44). Rhizobial QS systems antagonistically control EPS biosynthesis and motility-chemotaxis but also affect a wide spectrum of other physiological traits that vary among rhizobial species, including biofilm formation, swarming motility, the type III secretion system, plasmid transfer, cell division, metabolism, and transport (6, 39, 45).

Studies identifying the molecular mechanisms that underlie convergent evolution have shown that mutations involved in a given phenotype often target the same pathways, making experimental evolution a powerful tool to decipher regulatory networks (27, 46). In this line, our experiment allowed us to uncover a new component of the *R. solanacearum* Phc QS system and provided additional information on the functioning of the Phc network. Although the PhcBSR(Q) pathway was identified approximately 20 years ago (14), the roles of several components were not clearly elucidated. This was the case for the regulatory protein PhcQ, which was suspected to be involved in the QS cascade because it belongs to the PhcBSRQ operon. Here, we demonstrated that PhcQ is involved in the dynamics of activation of PhcA in response to the quorum. In addition, the synthesis of QS molecules was found to be affected in a *phcQ* mutant, suggesting that the activity of the autoinducer synthase PhcB is directly or indirectly regulated by PhcQ. The PhcQ protein is composed of a receiver domain and an uncharacterized C-terminal part. Several different mechanisms of action have been reported for proteins containing a single-receiver domain, including a direct regulatory effect on the activity of a downstream protein, the transfer of phosphate in phosphorelays, or a role as phosphate-sinks that redirect phosphate flux away from histidine kinases (47). It is also possible that PhcQ, through its receiver domain, interacts with PhcB and modulates its activity. Such a role in the synthesis and transduction of a diffusible signal factor (DSF) has been demonstrated for the receiver domain of the *Xanthomonas campestris* sensor kinase RpfC in a QS cascade (48). In a previous study, we showed that in some evolved *Ralstonia* lines, massive nodule cell infection capacity was gained through the inactivation of another central regulator, EfpR, controlling the membrane associated RSc3146 to -3148 genes (27). The *phc* and *efpR* mutations displayed similar in planta fitness, and their combination had no cumulative effect on this fitness. This, together with strong similarities in the transcriptomic and metabolo-
mic profiles of the two regulators, suggested a connection between the EfpR and PhcA regulatory pathways. Our findings indicated that the EfpR regulatory cascade does not interfere with the PhcBSRQ-mediated QS signal transduction and thus that the two paths act independently on the PhcA regulon. However, the mechanism by which the EfpR-Rsc3146-Rsc3148 pathway represses the PhcA regulon remains unknown. It may either repress PhcA activity, act independently on PhcA target genes, or impair one of the positive feedback loops of activation of PhcA (16, 33) (Fig. 9). Dashed lines indicate potential indirect effects.

In rhizobium-legume symbiosis, the accommodation of bacteria within nodule cells is a crucial step, since it protects internalized bacteria from plant defense reactions, while massive bacterial multiplication allows for nitrogen fixation levels that sustain plant growth (49). How this property is acquired during evolution is only partly understood. An intrinsic infection capacity is gained through the acquisition of nod genes that determine the synthesis of Nod factors. These lipochitooligosaccharide signal molecules specifically recognized by plant LysM receptors trigger the developmental program required for nodule organogenesis, bacterial infection (50, 51), and partial suppression of plant defenses (52). Other bacterial factors, such as surface polysaccharides, are likely recruited in recipient genomes to achieve infection. In Lotus japonicus, for instance, compatible bacterial EPS are specifically recognized by the plant LysM receptor kinase EPR3, which might activate a signaling cascade that controls and maintains an intracellular infection mode (53–55). In this study, we showed that inactivation of the central regulator PhcA was an alternative way to the inactivation of another global regulator, EfpR, to improve intracellular infection during experimental evolution of R. solanacearum into legume symbionts. Since PhcA and EfpR control a common set of more than 150 genes, including those for exopolysaccharides and motility and virulence factors, it is likely that the concerted change of expression of several genes rather than a single gene is involved in infection improvement. Indeed, the sole inactivation of XpsR, the activator of EPS biosynthesis genes, did not lead to an improved infection phenotype. Other virulence genes controlled by these systems may trigger plant defense reactions and thus hamper symbiosis establishment (16–18).
Another common interesting feature of both regulators is that their repression/inacti-
vation enhances the metabolism of the bacterium, increasing its growth rate on many
substrates. Future investigation of other intracellular lines, where EfpR and Phc path-
ways do not seem targeted by evolution, may narrow the number of possible functions
responsible for phenotypic changes. Yet, we were able to identify at which symbiotic
stage modifications were required, by monitoring the in planta expression of a direct
PhcA target, xpsR. A clear change induced by the phc mutations was a specific loss of
xpsR expression in infection threads of root hairs, suggesting that this step may
condition the success of the subsequent accommodation of bacteria within nodule
cells. The link between early and late events of the infection process is supported by the
fact that several plant genes (NFP/NFR5, LYK3/NFR1, SYMRK, or IPD3/CYCLOPS) in-
volved in early symbiotic signaling in the root epidermis are also required for the
release of bacteria in nodule cells (56–58). Furthermore, a tight coordination between
the progression of ITs and the periclinal divisions of specific root cell layers was
proposed to determine the intracellular infection of nodules (59). It is thus possible that,
for either signaling or metabolic reasons, the primary ITs formed by efpR and phc
mutations progress more rapidly than those made by a single hrpG mutant, allowing a
better invasion of nodule cells. Future fine cellular analyses of the progression rate of
ITs as well as a comparison of molecular plant responses differently induced by poor
and nice infectious evolved clones should help us to understand intracellular infection,
the distinctive feature of nitrogen-fixing root nodule symbioses and possibly the first
trait acquired by the common ancestor of the root nodule forming plant clade
FaFaCuRo (Fabales, Fabaceae, Cucurbitales and Rosales) during evolution (60).

**MATERIALS AND METHODS**

**Bacterial strains, plasmids, and growth conditions.** Bacterial strains and plasmids used in this work
are listed in Table S1 in the supplemental material. *R. solanacearum* strains were grown at 28°C either on
rich BG medium (61) or on MP minimal medium (62) supplemented either with 20 mM glutamate for
gene expression analyses or with glycerol 2% for bacterial transformation. Antibiotics were used at the
following concentrations: trimethoprim, 100 μg/ml; spectinomycin, 40 μg/ml; kanamycin, 50 μg/ml;
tetracycline, 10 μg/ml.

**Plants assays and cytological analyses.** *Mimosa pudica* seedlings from Australian origin (B&T World
Seed, Pauqignan, France) were cultivated as previously described (29).

For cytology, nodules were harvested at 21 days postinoculation (dpi) and cut into 55-μm sections
with a vibrating blade microtome (VT1000 S; Leica, Wetzlar, Germany). For each nodule, longitudinal
sections were observed using an inverted microscope (DM IRB/E; Leica, Wetzlar, Germany), and images
were acquired using a charge-coupled-device (CCD) camera (Color Coolview; Photonic Science, Milham,
UK). Quantification of infection and necrosis areas on nodule sections was performed as described
previously (25, 30) using the Image-Pro Plus software and based on the HIS method (Media Cybernetics,
Rockville, MD, USA). Necrotic areas are characterized by a dark-brown color, while intracellularly infected
cells are characterized by a light-brown color. Results of infection and necrosis quantifications were
obtained from 2 to 3 independent experiments for each strain. At least 16 plants were analyzed per
experiment.

For relative in planta fitness assays, we coinoculated *M. pudica* plants with a pair of different strains
in equivalent proportions (5 × 10⁶ bacteria of each strain per plant). Coinoculated strains were carrying
the same antibiotic marker and expressing or not the mCherry fluorophore. Nodules from 20 plants were harvested
21 days after inoculation, pooled, surface sterilized, and crushed together. Dilutions of nodule
crushes were spread on solid medium. In cases where one of the coinoculated strains was expressing the
mCherry fluorophore, colonies formed by each strain were counted under a stereo zoom microscope
with a vibrating blade microtome (VT1000 S; Leica, Wetzlar, Germany), and images
were acquired using a charge-coupled-device (CCD) camera (Color Coolview; Photonic Science, Milham,
UK). Quantification of infection and necrosis areas on nodule sections was performed as described
previously (25, 30) using the Image-Pro Plus software and based on the HIS method (Media Cybernetics,
Rockville, MD, USA). Necrotic areas are characterized by a dark-brown color, while intracellularly infected
cells are characterized by a light-brown color. Results of infection and necrosis quantifications were
obtained from 2 to 3 independent experiments for each strain. At least 16 plants were analyzed per
experiment.

**Beta-galactosidase assays.** Expression of the plasmidic *xpsR-lacZ* fusion was assessed by β-
galactosidase assay of 100-μl aliquots of bacterial cultures grown in rich BG medium. Samples
were collected either at an OD₆₀₀ of 0.2 (Fig. 3) or every 2 to 3 h along a growth curve from an OD₆₀₀ of
0.005 to an OD₆₀₀ of 1. β-Galactosidase activities were measured in Miller units as described
previously (63). The correspondence between the OD₆₀₀ units and the number of bacteria (CFU) was
determined for each strain throughout growth in rich BG medium by plating serial dilutions of
cultures (Fig. 52).

To test the production of QS molecules by the chimeric strains, the GMI1000(pRalta) *hrpG
ΔphcBΔm* strain unable to produce QS molecules and expressing the reporter gene fusion *xpsR-lacZ*
was grown in rich BG medium until the OD₆₀₀ reached ca. 0.2. In parallel, strains producing QS molecules
[GMI1000(pRalta) hrpG and its phcBR22C, phcQR154C, ΔphcBΔ_phs frame, ΔphcQ, ΔphcA, and efpR66K derivative mutants] were grown in rich BG medium until the OD₆₀₀ reached 0.1. Supernatants from cultures of QS molecule donor strains were prepared by centrifugation for 15 min at 5,000 rpm followed by filtration at 0.22 μm. One volume of filtered supernatant was added to 1 volume of culture of the reporter strain and incubated for 4 h at 28°C under agitation before β-galactosidase activity analysis. β-Galactosidase activities were measured from 100-μl aliquots of bacterial cultures. Background expression level of the xpsR-lacZ reporter fusion was measured by incubating the reporter strain with the supernatant of a culture of the GMI1000(pRalta) ΔphcBΔ_phs frame strain.

To test the QS signal transduction in the phcQR154C and ΔphcQ mutants, a similar experiment was performed using either the GMI1000(pRalta) hrpG phcQR154C ΔphcBΔ_phs frame xpsR-lacZ or GMI1000(pRalta) hrpG ΔphcQ ΔphcBΔ_phs frame xpsR-lacZ strains as reporter strains and the GMI1000(pRalta) hrpG as donor strain.

**Statistical analyses.** Statistical analyses of infection and necrosis sizes of nodule sections were performed using R software (version 3.5.1) and the nonparametric multiple-comparison test of Kruskal-Wallis. For the comparisons of in planta relative fitness, β-galactosidase activities in mid-exponential-phase cultures, and gene expression, we did two-sided t tests. The local polynomial regressions and confidence intervals of β-galactosidase activities throughout growth were performed using the R package ggplot2 and the geom_smooth function (64).

Other materials and methods concerning genome resequencing and detection of mutations in evolved clones, constructions of mutants and plasmids, detection of xpsR-lacZ expression in planta, gene expression analyses by quantitative reverse transcription-PCRs, pathogenicity assays, protein structure prediction, and protein alignments are provided in Text S1 in the supplemental material.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available only online.

TEXT S1, DOCX file, 0.1 MB.

FIG S1, PDF file, 0.6 MB.

FIG S2, PDF file, 0.6 MB.

FIG S3, PDF file, 0.3 MB.

FIG S4, PDF file, 0.6 MB.

TABLE S1, DOCX file, 0.1 MB.

TABLE S2, XLSX file, 0.1 MB.

TABLE S3, XLSX file, 0.1 MB.

TABLE S4, XLSX file, 0.1 MB.

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We declare no conflict of interest.

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