

# Regulation of Polyhydroxybutyrate Synthesis in the Soil Bacterium *Bradyrhizobium diazoefficiens*

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## ABSTRACT

Polyhydroxybutyrate (PHB) is a carbon and energy reserve polymer in various prokaryotic species. We determined that, when grown with mannitol as the sole carbon source, *Bradyrhizobium diazoefficiens* produces a homopolymer composed only of 3-hydroxybutyrate units (PHB). Conditions of oxygen limitation (such as microoxia, oxic stationary phase, and bacteroids inside legume nodules) were permissive for the synthesis of PHB, which was observed as cytoplasmic granules. To study the regulation of PHB synthesis, we generated mutations in the regulator gene *phaR* and the phasin genes *phaP1* and *phaP4*. Under permissive conditions, mutation of *phaR* impaired PHB accumulation, and a *phaP1 phaP4* double mutant produced more PHB than the wild type, which was accumulated in a single, large cytoplasmic granule. Moreover, PhaR negatively regulated the expression of *phaP1* and *phaP4* as well as the expression of *phaA1* and *phaA2* (encoding a 3-ketoacyl coenzyme A [CoA] thiolases), *phaC1* and *phaC2* (encoding PHB synthases), and *fixK<sub>2</sub>* (encoding a cyclic AMP receptor protein [CRP]/fumarate and nitrate reductase regulator [FNR]-type transcription factor of genes for microoxic lifestyle). In addition to the depressed PHB cycling, *phaR* mutants accumulated more extracellular polysaccharides and promoted higher plant shoot dry weight and competitiveness for nodulation than the wild type, in contrast to the *phaC1* mutant strain, which is defective in PHB synthesis. These results suggest that *phaR* not only regulates PHB granule formation by controlling the expression of phasins and biosynthetic enzymes but also acts as a global regulator of excess carbon allocation and symbiosis by controlling *fixK<sub>2</sub>*.

## IMPORTANCE

In this work, we investigated the regulation of polyhydroxybutyrate synthesis in the soybean-nodulating bacterium *Bradyrhizobium diazoefficiens* and its influence in bacterial free-living and symbiotic lifestyles. We uncovered a new interplay between the synthesis of this carbon reserve polymer and the network responsible for microoxic metabolism through the interaction between the gene regulators *phaR* and *fixK<sub>2</sub>*. These results contribute to the understanding of the physiological conditions required for polyhydroxybutyrate biosynthesis. The interaction between these two main metabolic pathways is also reflected in the symbiotic phenotypes of soybeans inoculated with *phaR* mutants, which were more competitive for nodulation and enhanced dry matter production by the plants. Therefore, this knowledge may be applied to the development of superior strains to be used as improved inoculants for soybean crops.

*Bradyrhizobium diazoefficiens* is an important soil bacterium that fixes atmospheric N<sub>2</sub> in symbiosis with soybean plants, a key crop for food production worldwide (1). In addition to its agricultural relevance, *B. diazoefficiens* may be of industrial interest because it produces significant quantities of polyhydroxybutyrate (PHB), a polymer that accumulates as granules in the cytoplasm and has potential use as biodegradable plastic (2–6). PHB granules are synthesized as sinks of excess carbon and reducing power and are used as carbon and energy reserves when bacteria face starvation conditions (7). However, this cycle of synthesis and degradation must be regulated because if the two pathways occur simultaneously (8), the net result would be consumption of energy and reducing power. The proteins involved in the different steps of the PHB cycle are well-characterized (9–12), and all of them are present in *B. diazoefficiens* (3, 13, 14); however, regulation of the cycle in this bacterium was not yet studied. In other rhizobium species, such as *Rhizobium etli* and *Ensifer meliloti*, the gene product of *phaR* (PHA [polyhydroxyalkanoate] regulator, previously known as *aniA* for anaerobically induced gene A) was reported to control, at least in part, PHB synthesis (15, 16).

The regulatory circuit in which PhaR takes part was best stud-

ied in *Ralstonia eutropha* (17–19). In this bacterial species, PhaR binds to the promoter of its own gene and to the promoter of *phaP*, which encodes a protein that binds to the surface of PHB granules called phasins (PhaP). Phasins contain amphipathic  $\alpha$ -helices whereby these proteins may coat the hydrophobic granules during their growth, exposing a polar surface to the bulk cytoplasm (20, 21). It is believed that, in this way, the phasins may control the final size of the PHB granules (22). In addition, PhaR also binds to the surface of PHB granules, although with less af-

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finitly than phasins (18). Thus, the model proposed for the regulation of PHB granule growth through PhaR/PhaP activities suggests that, while the granules are growing, their surface area is enough to accommodate PhaR and PhaP, thus maintaining a low cytoplasmic concentration of free PhaR, which keeps the *phaR* and *phaP* promoters free. As the granules reach a critical volume, their surface areas become limiting for PhaR and PhaP, which are continuously synthesized. At this point, PhaP displaces PhaR from the granule surface, raising the PhaR cytoplasmic concentration. The free PhaR binds *phaR* and *phaP* promoters, and the expression of the two genes is inhibited, arresting PHB granule growth (18, 23). However, a recent study reported that the association constant of PhaR to its target DNA sequence in the *phaR* promoter is similar to that of an unspecific DNA sequence (24).

In *E. meliloti*, the symbiont of alfalfa, PhaP and PhaR (AniA) activities are necessary for PHB synthesis (16, 25). In addition, *phaR* (*aniA*) may play a more ample regulatory role in the carbon flow in *R. etli*. A *phaR::Tn5* mutant synthesized around 40% of the PHB level of the wild type and had significantly increased extracellular polysaccharide (EPS) levels, with extensive changes in its proteome (15). However, the regulatory circuit through which *phaR* may control EPS synthesis is unknown.

The genome of *B. diazoefficiens* USDA 110 harbors one copy of *phaR* (*blr0227*), which is located adjacent to PHB-related genes *phaA2* (*bll0226*) and *phaB2* (*bll0225*) but is transcribed in the opposite direction (see Fig. S1A in the supplemental material). Meanwhile, at least four paralogs of *phaP* occur as isolated genes at locations elsewhere in the genome. The expression of these genes depends on the culture conditions. In particular, conditions permissive for PHB synthesis, such as microoxia (26) and growth in yeast extract-mannitol (YM) (14), increase the expression of *phaP1* and *phaP4*, the last being the paralog with the highest expression in YM. PhaR and the four PhaP paralogs bind to PHB *in vitro* (14), with PhaP4 being the phasin with the highest affinity. Further, an increasing concentration of PhaP4 triggered a competitive displacement of PhaR bound to PHB fine powder in suspension (14). These results suggested that PhaP4 may be the main phasin responsible for PhaP biological function in *B. diazoefficiens*. However, mutations of these genes were not obtained, and thus their implication in the regulation of PHB synthesis could not be directly established *in vivo*.

To better understand the regulation of PHB in *B. diazoefficiens* USDA 110, we constructed mutant strains in *phaR* and in two *phaP* genes and evaluated their roles in PHB and EPS syntheses as well as in the symbiotic interaction with soybean plants.

## MATERIALS AND METHODS

**Bacterial strains and culture conditions.** Strains and plasmids are summarized in Table S1 in the supplemental material. *B. diazoefficiens* was grown oxically or microoxically in Götz minimal medium with mannitol as the sole carbon source (27). For oxic growth, 50-ml cultures were grown in 250-ml Erlenmeyer flasks at 30°C with rotary shaking at 180 rpm in contact with the air. For microoxic growth, cells were inoculated into 500-ml serum bottles containing 35 ml of Götz medium to an optical density at 500 nm ( $OD_{500}$ ) of 0.03. The bottles were closed with rubber septa and maintained at 30°C and 60 rpm under microoxic conditions by exchange of gas phase (0.5% O<sub>2</sub> and 99.5% N<sub>2</sub>) every 8 to 16 h until the desired growth state (26). Total biomass was estimated by  $OD_{500}$  and by the number of viable bacteria by CFU counts on YM agar plates (28). Conjugation was performed in a modified peptone salts-yeast extract medium (26). *Escherichia coli* was grown in Luria-Bertani medium (29). An-

tibiotics were added to the media at the following concentrations: 400 (*B. diazoefficiens*) or 100 (*E. coli*)  $\mu\text{g ml}^{-1}$  streptomycin (Sm); 200 (*B. diazoefficiens*) or 100 (*E. coli*)  $\mu\text{g ml}^{-1}$  spectinomycin (Sp); 150 (*B. diazoefficiens*) or 25 (*E. coli*)  $\mu\text{g ml}^{-1}$  kanamycin (Km); 200  $\mu\text{g ml}^{-1}$  (*E. coli*) ampicillin (Ap); 100 (*B. diazoefficiens*) or 10 (*E. coli*)  $\mu\text{g ml}^{-1}$  gentamicin (Gm); 20 and 100 (*B. diazoefficiens* liquid and solid cultures, respectively) or 10 (*E. coli*)  $\mu\text{g ml}^{-1}$  tetracycline (Tc); and 20  $\mu\text{g ml}^{-1}$  (*B. diazoefficiens*) chloramphenicol (Cm).

**Genetic techniques and DNA manipulation.** Cloning procedures, including DNA isolation, restriction digestion, ligation, and transformation, were performed as described previously (29). Tri- or biparental matings were performed with *E. coli* DH5 $\alpha$  or S17-1 strains as described previously (3). Electroporation was performed with a Gene Pulser system (Bio-Rad, Hercules, CA) at 1.5 V, 25  $\mu\text{F}$ , and 200  $\Omega$  in a 0.1-cm-gap-width electroporation cuvette.

Oligonucleotide primers were purchased from Life Technologies (Buenos Aires, Argentina) or Sigma-Aldrich (Madrid, Spain). DNA amplification was performed by PCR using *Taq* DNA polymerase (Life Technologies, Buenos Aires, Argentina) for routine PCR or Kapa HiFi HotStart (HS) DNA polymerase (Kapa Biosystems, Woburn, MA) or Pfx (Life Technologies, Buenos Aires, Argentina) for the amplification of blunt-ended amplicons and/or targets longer than 1,000 bp. DNA sequencing was performed at Macrogen Corp. (Seoul, South Korea).

To construct the *B. diazoefficiens phaR* mutant, primers specific for the *blr0227* locus tag were designed (see Table S2 in the supplemental material). An internal 244-bp PCR fragment was amplified with primers Fw1 and Rv1 and cloned into pGEM-T Easy, creating pIQ35. This plasmid was digested with EcoRI and ligated into pG18*mob2*, generating pIQ36, which was conjugated into *B. diazoefficiens* LP 3004 by biparental mating using *E. coli* S17-1 as the donor. Transconjugants were selected in Gm, Sm, and Cm. We obtained two independent clones by single homologous recombination of pIQ36 in the middle of the *phaR* coding sequence, leading a predicted truncated polypeptide of 108 amino acids (58% of the full-length protein sequence). The results obtained regarding growth, PHB synthesis, and extracellular polysaccharide production were similar in the two clones, and, therefore, we selected one of them that was designated LP 0227 (here referred to as the *phaR* mutant) (see Fig. S1A in the supplemental material).

For complementation, the complete sequence of *phaR* (amplified with primers Fw1c/Rv1c) was integrated into a replicative vector. Briefly, a 1,245-bp target sequence and its own putative promoter (inferred by the BPROM program) were amplified from *B. diazoefficiens* LP 3004 and cloned into the *Sma*I site of pBBR1-MCS-5 to create pIQ37. Then, pIQ37 was digested with XbaI and PstI, and the *phaR*-containing fragment was inserted in the replicative broad-host-range promoter-probe vector pCB303 (30), generating plasmid pIQ38. The construction was confirmed by sequencing. The plasmid was transferred into the LP 0227 strain by biparental mating, selected by growth on Tc, Gm, and Cm. The LP 0227 strain carrying pIQ38 turned blue in YM agar plates supplemented with X-phosphate, thus confirming that the 1,245-bp fragment, cloned in the same orientation as the promoterless *phaA* gene in pIQ38, carried both the *phaR* promoter and the *phaR* coding sequence.

To construct the *B. diazoefficiens* phasin mutants, primers specific to the *bls155* (*phaP1*) or *bl7395* (*phaP4*) locus tags were designed (see Table S2 in the supplemental material). Fragments of the *B. diazoefficiens* USDA 110 genomic DNA were generated from the two sides of the target coding sequences. For the two phasins, the strategy was similar. An upstream fragment of 249/253 bp and a downstream fragment of 235/222 bp were amplified using primer pair Fw2/Rv2 or Fw3/Rv3 (fragments *phaP1-1* and *phaP4-1*, respectively) and primer pair Fw4/Rv4 or Fw5/Rv5 (fragments *phaP1-2* and *phaP4-2*, respectively). The PCR fragments *phaP1-2* and *phaP4-2* were cloned into the *Sma*I site of pBBR1-MCS-4, creating pIQ39 and pIQ40. These plasmids were then digested with EcoRV, and the fragments *phaP1-1* and *phaP4-1* were introduced, creating pIQ41 and pIQ42, respectively. These plasmids were linearized with

EcoRI. In parallel, pHP45 $\Omega$ SmSp and pHP45 $\Omega$ Km were EcoRI-digested, and the resulting fragments  $\Omega$ Sm-Sp and  $\Omega$ Km were cloned into pIQ41 and pIQ42, creating pIQ43 and pIQ44, respectively. Finally, these two plasmids were digested with XbaI and KpnI, and the fragments of pIQ43 and pIQ44 were ligated into XbaI/KpnI-digested pK18mob and pG18mob2, creating pIQ45 and pIQ46, respectively.

For *phaP1*, gene replacement was performed by introducing pIQ45 into *B. diazoefficiens* USDA 110 by biparental mating, selecting by growth on Sm-Sp and Cm, and screening for Km sensitivity. The resulting strain was designated LP 5155 ( $\Delta$ *phaP1*); this strain carries the  $\Omega$ -Sm-Sp interposon in place of the genomic DNA from base 5711611 to base 5711972, thus removing 301 bp from the 339-bp open reading frame (ORF) bll5155 (see Fig. S1B in the supplemental material). For *phaP4*, gene replacement was performed by introducing pIQ46 into *B. diazoefficiens* USDA 110 by triparental mating using the pRK2013 helper plasmid, selecting by growth on Km and Cm, and screening for Gm sensitivity. The resulting strain was designated LP 7395 ( $\Delta$ *phaP4*); this strain carries the  $\Omega$ -Km interposon in place of the genomic DNA from base 8130454 to base 8130880, thus removing 426 bp from the 435-bp ORF bll7395 (Fig. S1B). The double mutant LP 5173 ( $\Delta$ *phaP1*  $\Delta$ *phaP4*) was constructed as described above, introducing pIQ45 in the background of LP 7395.

All of the mutations were confirmed by PCR using external primers (see Table S2 and Fig. S1 in the supplemental material) as described previously (3) and by DNA sequencing.

**Bioinformatic characterization of PhaR and PhaP.** PhaR (blr0227), PhaP1 (bll5155), and PhaP4 (bll7395) were aligned with their putative homologs in *E. meliloti* 1021, *R. etli* CFN42, *Azorhizobium caulinodans* ORS571, and *Rhizobium leguminosarum* bv. *viciae* 3841 using T-Coffee (31). The secondary structures of PhaP1 and PhaP4 were predicted using Phyre2 alpha helix prediction (<http://www.sbg.bio.ic.ac.uk/phyre2>) and the PSIPRED v 3.3 server (<http://bioinf.cs.ucl.ac.uk/psipred/>).

**RNA isolation, cDNA synthesis, and qRT-PCR analysis.** *B. diazoefficiens* LP 3004 and *phaR* mutant cultures were grown microaerobically as described above. The cultures were grown to mid-exponential phase at OD<sub>500</sub> values of 0.25 to 0.33 (*phaR* mutant) or 0.50 to 0.62 (LP 3004), stopped by adding cool stop solution (equilibrated phenol and ethanol at a ratio of 1:9), collected by centrifugation at 4°C, and frozen at -80°C as previously described (26, 32). This material was used for RNA isolation and cDNA synthesis as described elsewhere (33). Primers for quantitative reverse transcription-PCRs (qRT-PCRs) were designed (see Table S2 in the supplemental material) to have a melting temperature of 57°C to 62°C and to generate PCR products of 87 to 134 bp. Each PCR mixture contained 9.5  $\mu$ l of iQ SYBR green supermix (Bio-Rad, Hercules, CA), 2  $\mu$ M individual primers, and appropriate dilutions of different cDNAs in a total volume of 19  $\mu$ l. Reactions were run in triplicate in a Bio-Rad iCycler. Melting curves were generated with Bio-Rad iQ5 software to verify the specificity of the amplification. Relative changes in gene expression (fold changes) were calculated as described elsewhere (34). The fold changes observed were considered significant when their absolute values were higher than 2.0. Expression of the primary sigma factor gene *sigA* was used as a reference for normalization.

**Quantification and analysis of PHB and extracellular polysaccharides.** For quantification of PHB, *B. diazoefficiens* cultures were centrifuged twice at 12,000  $\times$  g for 40 min. Then, the pellets were homogenized overnight with sodium hypochlorite at room temperature, washed with double-distilled water, precipitated with 1:1 alcohol-acetone, and resuspended in chloroform. Then, PHB was determined as crotonic acid in H<sub>2</sub>SO<sub>4</sub> (35). For analysis of composition, PHB was extracted from lyophilized *B. diazoefficiens* cells as described previously (36), and monomer composition was analyzed by gas chromatography (GC) after conversion of PHB to 3-hydroxybutyryl-methyl esters via acid methanolysis (36). Benzoate-methyl ester was used as an internal standard.

To determine the PHB content of bacteroid cells, 10 fresh mature nodules (70 to 90 mg) were washed, crushed and homogenized, suspended in 0.5 ml of double-distilled water, and centrifuged for 5 min at

500  $\times$  g (3). The supernatants were then processed for PHB determination as above.

Extracellular polysaccharides (EPSs) include exopolysaccharide and capsular polysaccharide, which have the same sugar composition in *B. diazoefficiens* USDA 110 (37). Exopolysaccharides were precipitated from the culture supernatants as described previously (38), and capsular polysaccharides were released from bacterial cells and precipitated as described previously (37). Finally, the polysaccharides were hydrolyzed, and the reducing sugars were quantified with anthrone as described previously (38).

**TEM.** Transmission electron microscopy (TEM) was performed essentially as described previously (3, 38). Briefly, bacteria from liquid cultures or nodules (each one transversally cut in half) were collected and fixed in 2% (vol/vol) glutaraldehyde. Next, the samples were dehydrated, infiltrated with epoxy resin, sectioned, and stained. Glutaraldehyde-fixed samples were placed in grids and viewed at 80 kV in a JEM 1200 EX II transmission electron microscope (JEOL Ltd., Tokyo, Japan).

Some TEM micrographs of free-living cells were used to determine the diameters of PHB granules, using ImageJ software (v 1.49). The largest diameter of every individual granule was registered from each individual cell ( $n = 132$  to 411).

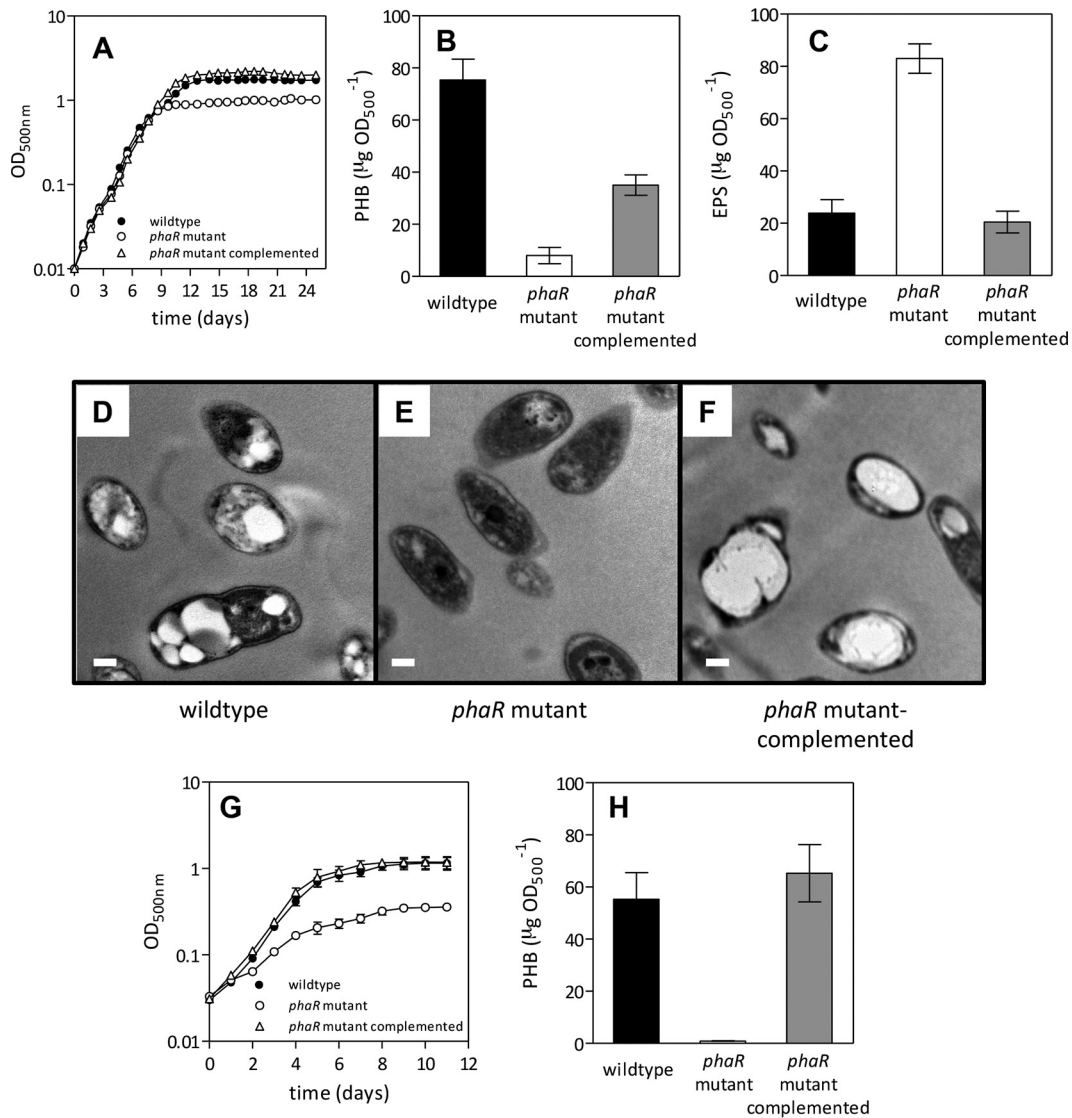
**Plant experiments.** DM 3810 soybean seeds, kindly provided by DonMario SA, Argentina, were surface sterilized, germinated, and cultivated in vermiculite pots as described previously (3). The stability of plasmid pIQ38 was verified by bacteroid isolation and CFU determination on plates containing Cm or Cm and Tc. Competition for nodulation was studied by inoculating LP 3004 and *phaR* mutant strains in a 1:1 proportion (as CFU ml<sup>-1</sup>) to soybean plants as described previously (39). Nodule occupation was evaluated on the basis of the antibiotic resistance of the strains recovered from the nodules 21 days after inoculation. Statistical analysis was carried out with the  $\chi^2$  test, considering 45% nodule occupation by each strain alone and 10% double occupation as the null hypothesis (38).

Plant growth parameters were assessed in 67-day-old plants. Shoots from each plant were cut and dried in an oven at 60°C to a constant weight, and the materials from each plant were weighed individually. The statistical analysis was performed by one-way analysis of variance (ANOVA), with an  $\alpha$  of <0.05. All results shown are representative of duplicate experiments.

## RESULTS

***B. diazoefficiens* produces PHB homopolymer.** The polyhydroxyalkanoates extracted from the lyophilized cells of *B. diazoefficiens* USDA 110 (wild type), LP 3004 (wild-type, spontaneous Sm-resistant derivative of USDA 110), and *phaR* mutant strains from exponential-phase (7 days) and late-stationary-phase (25 days) cultures under oxic conditions (Fig. 1A) were characterized for composition of their monomer units by GC analysis. GC chromatograms from the two culture ages were similar, and those corresponding to the stationary phase are shown in Fig. S2 in the supplemental material. As observed, polyhydroxyalkanoates of the three strains tested possess only 3-hydroxybutyrate units. This finding allowed us to quantify the polyhydroxyalkanoate contents by the detection of crotonic acid at 235 nm because this specific analysis method applies only for poly(3-hydroxybutyrate) [poly(3HB)] (35).

**PHB and EPS syntheses are controlled by *phaR* under oxygen-limiting conditions.** We measured total biomass (as OD<sub>500</sub>) daily in oxic cultures for 25 days in the wild-type, *phaR* mutant, and mutant-complemented strains (Fig. 1A). There were no differences in exponential growth rates, but the stationary phase was reached earlier in the *phaR* mutant, resulting in reduced biomass with respect to the wild-type and mutant-complemented strains.



**FIG 1** Physiological characterization of the *phaR* mutant under oxic or microoxic conditions. (A to F) Oxic conditions, with rotary shaking at 180 rpm in contact with the air. (A) Growth kinetics. (B) PHB extracted from 25-day-old cells. (C) EPS obtained from 25-day-old cultures. (D to F) Transmission electron micrographs of cells from 25-day-old cultures showing bright PHB granules (scale bar, 0.2  $\mu\text{m}$ ). (G and H) Microoxic conditions, with rotary shaking at 60 rpm in contact with an atmosphere of 0.5%  $\text{O}_2$  and 99.5%  $\text{N}_2$ . (G) Growth kinetics. (H) PHB extracted from 11-day-old cells. In all cases, quantitative results are the average  $\pm$  standard error of the mean (SEM) from four independent cultures. When error bars are not visible, they are smaller than the symbol.

Next, we measured PHB levels at 7 (logarithmic), 13 (early stationary phase), 17 (middle stationary phase), and 25 (late stationary phase) days of growth. The production of PHB at 7 days of growth was higher in the *phaR* mutant than in the wild type, but later, there was a strong increase in PHB production in the wild type at the early, middle, and late stationary phases that was not observed in the *phaR* mutant (see Fig. S3A in the supplemental material). At the late stationary phase, the mutant-complemented strain accumulated more PHB than the mutant but only half of the wild-type PHB levels (Fig. 1B). In addition, we measured EPS contents at the same growth states. While there were no differences between strains at the logarithmic and early and middle stationary phases (data not shown), at the late stationary phase the *phaR* mutant produced about 4-fold more EPS than the wild-type and *phaR*-complemented strains (Fig. 1C), which was also re-

flected macroscopically in the high mucoidy of the *phaR* mutant cell pellets after centrifugation (see Fig. S3B in the supplemental material). TEM of 25-day-old cells (Fig. 1D to F) showed that, while the *phaR* mutant cells were devoid of PHB granules, the wild-type and the *phaR* mutant-complemented strains had PHB granules (in the complemented strain, most cells had a single large granule).

In contrast, in microoxic cultures, the growth rate was lower and total biomass of the *phaR* mutant was smaller than those of the wild-type strain or the mutant-complemented strain from the beginning of the culture (Fig. 1G). Likewise, PHB production was affected in the mutant at the early stationary phase (11 days of growth) and was restored in the mutant-complemented strain (Fig. 1H). These results, in agreement with previous investigations (3, 27, 40), indicated that the microoxia and the stationary phase

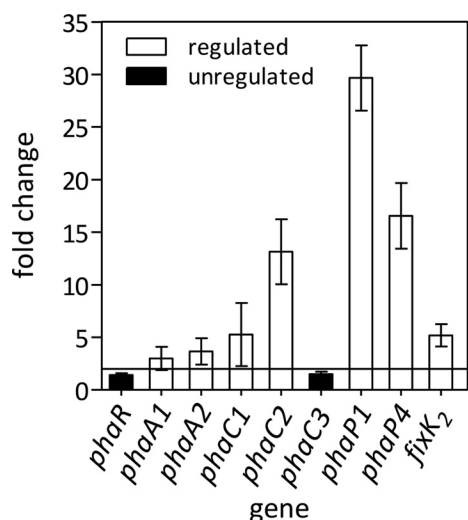


FIG 2 Differential expression of selected genes among the wild type and the *phaR* mutant, evaluated by qRT-PCR in cells grown microaerobically. Fold change refers to the relative expression in the mutant compared to that in the wild type. An absolute value fold change of  $\geq 2$  (line of reference) was considered statistically significant. Results are the average from three runs  $\pm$  SEM.

of oxic cultures were permissive conditions for PHB synthesis while the exponential phase of the oxic cultures was nonpermissive. In addition, the oxygen-limiting bacteroid state inside root nodules is also a permissive condition (3). Our results indicate that the influence of PhaR on PHB production is mainly exerted under these permissive conditions.

**PhaR exerts a negative control on genes related to PHB and *fixK2* expression.** The permissive conditions indicated above both involve oxygen limitation. Because oxic cultures of the *phaR* mutant at the late stationary phase produce too much EPS, which interferes with RNA extraction, we evaluated the role of PhaR as a transcriptional regulator in mid-exponential-phase microoxic cultures. We compared the levels of certain transcripts in the *phaR* mutant and the wild type by qRT-PCR as summarized in Fig. 2. The levels of most of the transcripts were lower in the wild type, indicating that *phaR* is a repressor, which is in agreement with previous reports (41). However, the mutation of *phaR* had no effect on its own transcription, which is in line with the low association constant of PhaR with respect to its target DNA sequence at the *phaR* promoter region described in *R. eutropha* H16 (24).

Phasin genes are known as direct targets of *phaR*. In *B. diazoefficiens*, there are at least four paralogs encoding phasins, of which *phaP1* and *phaP4* have the highest expression levels (14, 26) and PhaP4 has the highest affinity for PHB *in vitro* (14). Therefore, we assessed the expression of these two phasin genes and found that transcription of *phaP1* and *phaP4* was strongly derepressed in the *phaR* mutant, with *phaP1* being the most affected.

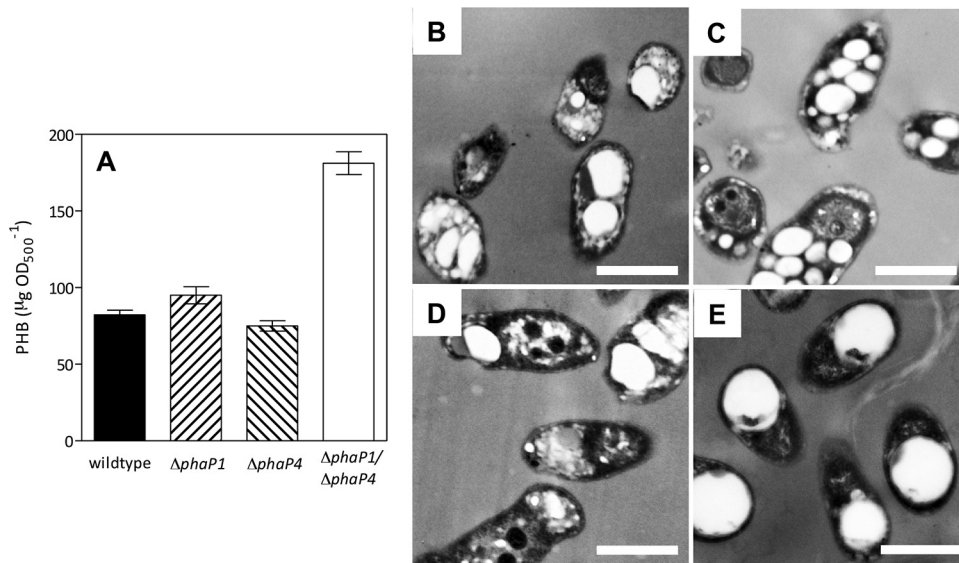
In addition, we evaluated the transcript levels of genes encoding enzymes of the PHB biosynthetic pathway. Particularly, we assessed the 3-ketoacyl-coenzyme A (CoA) thiolase paralogs *phaA1* and *phaA2* and the PHB synthase paralogs *phaC1*, *phaC2*, and *phaC3*. Of these three PHB synthase-coding genes, only *phaC1* is required (3). In turn, a deletion of *phaC2* causes significant increases in PHB synthesis, while PhaC3 seems active only in a *phaC2* mutant background (3). We observed that *phaC2* was significantly (13-fold) upregulated, and transcription of *phaC1*

was increased 5-fold while transcription of *phaC3* was not affected in the *phaR* mutant. The most upregulated gene in the mutant was *phaC2*, and interestingly, this gene was previously reported as positively controlled by the global regulator *fixK2* (26). In the same line as the  $\Delta$ *phaC2* strain, a  $\Delta$ *fixK2* strain also accumulated more PHB than the wild type in the stationary phase (see Fig. S4 in the supplemental material). Therefore, we also measured the amounts of *fixK2* transcripts and found that, as with the other genes, mutation of *phaR* resulted in a  $\sim$ 6-fold upregulation of *fixK2* transcription.

**Phasin mutants showed altered patterns of PHB granule size and accumulation.** The secondary structure prediction analyses of PhaP1 and PhaP4 indicated high prevalences of alpha helix structures (91% and 86%, respectively, data not shown), which are compatible with known phasins. Moreover, growth kinetics of the  $\Delta$ *phaP1*,  $\Delta$ *phaP4*, and  $\Delta$ *phaP1*  $\Delta$ *phaP4* deletion mutants were similar to the wild type, indicating that none of the phasin genes are essential for growth in *B. diazoefficiens*. In addition, PHB accumulations in the single mutants were similar to that of the wild type at the middle stationary phase, but the  $\Delta$ *phaP1*  $\Delta$ *phaP4* strain accumulated  $\sim$ 3 times more PHB (Fig. 3A). However, although the  $\Delta$ *phaP4* mutant produced PHB granules that did not differ significantly from those seen with the wild type ( $0.420 \pm 0.134 \mu\text{m}$  and  $0.521 \pm 0.145 \mu\text{m}$  in diameter, respectively), the  $\Delta$ *phaP1* mutant produced smaller granules ( $0.304 \pm 0.115 \mu\text{m}$ ) but in greater quantity than the wild type (see Fig. S5 in the supplemental material). In addition,  $\Delta$ *phaP1*  $\Delta$ *phaP4* cells had large PHB granules ( $0.774 \pm 0.140 \mu\text{m}$ ) (Fig. 3B to D; see also Fig. S5 in the supplemental material) that occupied a considerable portion of the cytoplasm (Fig. 3E), similar to a *phaP1* mutant in *R. eutropha* (23, 42). Thus, an influence of PhaP1 on granule surface/volume ratio as well as an interaction between PhaP1 and PhaP4 for controlling the growth of the granules may be appreciated.

**The *phaR* mutant has a better symbiotic performance than the wild type.** Previous reports on the nodulation of *phaR* mutants are contradictory. On the one hand, the *E. meliloti* *phaR* mutant formed many pseudonodules in alfalfa that were negatively altered in  $\text{N}_2$  fixation (16); on the other hand, the *R. etli* *phaR* mutant produced macroscopically normal nodules in the common bean without significant differences in nitrogenase activity with respect to that of the wild type (15). However, nodulation differs among these host plants because alfalfa produces indeterminate nodules without PHB while the common bean produces determinate nodules, which have typical PHB granules in the bacteroidal cytoplasm. Since soybean plants produce determinate nodules, we wanted to know if our *phaR* mutant behaves similarly to *R. etli* for its symbiotic interaction with soybean roots.

As in the common bean inoculated with *R. etli*, soybean plants inoculated with the *B. diazoefficiens* *phaR* mutant produced morphologically normal nodules. However, when nodules from 67-day-old plants were observed by TEM, it became evident that they were completely devoid of PHB granules (Fig. 4B). When this mutant was complemented with the wild-type *phaR* gene in *trans*, the production of PHB granules in soybean nodules was restored, indicating that the mutation in *phaR* was responsible for the lack of PHB granule production in the mutant bacteroids (Fig. 4C) as observed above under free-living conditions. Furthermore, the analytic determination of PHB levels from crushed nodules corroborated these observations (Fig. 4D). It is possible that the low PHB levels detected in nodules infected by the *phaR* strain ( $0.03$



**FIG 3** PHB accumulation in the wild type and three  $\Delta phaP$  mutant derivatives, evaluated at 17 days under oxic conditions. (A) PHB extracted from cells, expressed as the average from four independent determinations  $\pm$  SEM. Transmission electron micrographs of the wild-type (B),  $\Delta phaP1$  (C),  $\Delta phaP4$  (D), and  $\Delta phaP1 \Delta phaP4$  (E) strains. Note the PHB granules as bright spots inside the cytoplasm. Scale bars, 0.5  $\mu$ m.

$\mu$ g · mg nodule fresh weight<sup>-1</sup>) correspond to PHB granules from bacteria (not bacteroids) that remained in the infection threads.

Competition for nodulation was evaluated by inoculating soybean plants with 1:1 mixtures of wild-type and *phaR* mutant strains, each one at a concentration on the order of 10<sup>5</sup> CFU ml<sup>-1</sup>. Because late-stationary-phase cultures are poor competitors (40) and we wanted to compare permissive and nonpermissive conditions, we prepared the inocula from oxic cultures either in the exponential phase (nonpermissive) or in the middle stationary phase (permissive). As a result, we observed in two independent experiments that under nonpermissive conditions the levels of competitiveness of the two strains were similar, while under permissive conditions the *phaR* mutant was more competitive than the wild type, occupying 74% of the nodules (Fig. 4E).

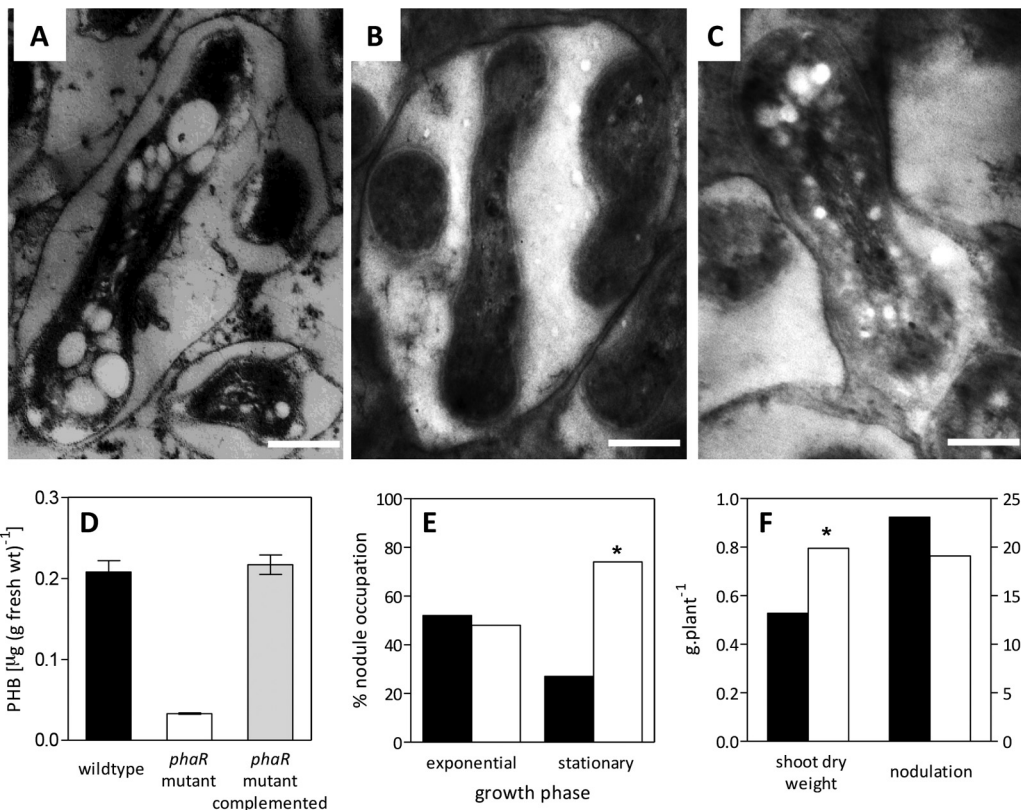
Despite differences in competitiveness, there were no differences in the numbers of nodules produced by the wild type and the *phaR* mutant (Fig. 4F). However, the shoot dry weights of 67-day-old plants cultured in nutrient solution without combined nitrogen were significantly higher in the plants inoculated with the *phaR* mutant (Fig. 4F). This result suggests that each nodule occupied by the *phaR* mutant was more active than those occupied by the wild type.

## DISCUSSION

As observed in other bacterial species, *phaR* was necessary for the production of PHB in both microoxic and stationary-phase oxic cultures of *B. diazoefficiens*. In addition, mutation of *phaR* impaired biomass accumulation in stationary-phase oxic cultures and compromised the growth of microoxic cultures at all growth phases. These results indicate that PhaR exerts its activity under conditions of oxygen limitation, in which accumulation of the substrates for PHB synthesis, similarly to reducing power and acetyl-CoA levels, may take place. In addition, the lack of PHB granules in our *phaR* mutant was accompanied by a higher production of EPS, a correlation previously observed in *E. meliloti* (16), *R. etli* (15), and N-limited *B. diazoefficiens* wild-type cultures

(27, 40). However, a plausible explanation of how *phaR* also controls EPS synthesis is lacking. Interestingly, PhaR also appeared as a repressor of *fixK<sub>2</sub>*, a global regulator which activates the expression of more than 200 genes needed for life under low-oxygen tension, including essential operons such as *fixNOQP* and *fixGHIS* (26, 43, 44). Because the expression of *phaR* is not affected in a  $\Delta fixK_2$  strain grown under oxygen-limiting conditions (26), it seems that *phaR* is at the highest hierarchy of this regulatory circuit. Therefore, the regulation scope of PhaR may be wider than previously thought. In Fig. 5, we propose an extended model for the regulation of PHB synthesis, including the findings obtained in this work.

At least three models may explain the initiation of PHB synthesis in *R. eutropha* (6), but it is unknown which of them is functional in *B. diazoefficiens*. Our results indicate that after initiation under permissive conditions, PHB synthesis would be catalyzed by basal levels of 3-ketoacyl-CoA thiolase, acetoacetyl-CoA reductase, and PHB synthase using microoxically accumulated reducing power and acetyl-CoA as the substrates. At this point, *phaR* may be expressed, but since the surface/volume ratio of the PHB granule is large, most of the PhaR produced should bind the PHB granule surface, leaving the *phaC1*, *phaC2*, *phaP1*, *phaP4*, and *fixK<sub>2</sub>* promoters free. FixK<sub>2</sub> stimulates the expression of *phaC2*, which encodes an inactive PHB synthase subunit that, by mass action, may impair the formation of PhaC1/PhaC1 fully active homodimers while increasing the concentration of PhaC1/PhaC2 less-active heterodimers and PhaC2/PhaC2 inactive homodimers (3, 14, 17, 45). Thus, the effect of the juxtaposition of *phaR* and *fixK<sub>2</sub>* on *phaC2* regulation may allow for a fine-tuning of the whole PHB synthase activity at the stage of PHB granule growth. Interestingly, *fixK<sub>2</sub>* and *phaC2* are also induced under conditions of oxidative burst, where an increase in EPS production was also observed (46). Therefore, we can speculate that PHB synthesis might be diminished under oxidative damage, perhaps by down-regulation of *phaR*. In addition to PhaC2, the phasins PhaP1 and



**FIG 4** Effects of *phaR* on the symbiotic interaction with soybean plants. (A to C) Transmission electron micrographs of ultrathin cuts of 67-day-old soybean nodules showing bacteroids. (A) Wild type; (B) *phaR* mutant; (C) *phaR* mutant complemented. Note the PHB granules as bright spots in panels A and C. Scale bars, 0.5  $\mu\text{m}$ . (D) Average PHB content ( $\pm$  SEM) from soybean nodules obtained from ten 67-day-old plants inoculated with the indicated strains. (E) Percentages of nodules occupied by the wild type (black bars) or the *phaR* mutant (white bars) inoculated in 1:1 mixtures on 10 soybean plants. Inoculants were prepared with cultures harvested at the physiological states indicated. (F) Shoot dry weight and nodulation of 10 soybean plants inoculated with the wild type (black bars) or the *phaR* mutant (white bars) after 67 days of growth. Values shown are the results determined by subtracting the value corresponding to the uninoculated control (shoot dry weight,  $0.658 \pm 0.09 \text{ g plant}^{-1}$ ; nodules, 0). Asterisks in panels E and F indicate statistically significant differences ( $\alpha < 0.05$ ).

PhaP4 may play important roles in controlling the size of PHB granules. Although PhaP4 seems to be the main protein covering PHB granules (14), only the double mutant  $\Delta\text{phaP1 } \Delta\text{phaP4}$  synthesized more PHB than the wild type. Moreover, when the two phasins were missing, the PHB granule surface/volume ratio decreased, and the bacteria possessed only one large granule per cell, resembling other bacterial phasin mutants (23, 47). Hence, in *B. diazoefficiens*, the combined function of PhaP1 and PhaP4 may prevent PHB granule coalescence, as reported in *R. eutropha* (42). This combined function suggests that in *B. diazoefficiens*, phasins may form trimers and/or tetramers as in other PHB-producing bacteria (48–50) as homo- and/or hetero-oligomers. In this way, the relative concentrations of PhaC1 and PhaC2 may modulate PHB synthase activity, while the production of PhaP1 and PhaP4 may control the number and size of PHB granules. As the granules reach their final sizes, PhaR would be released from their surface, exerting its inhibitory effect toward all of these PHB synthesis proteins. According to this model, the higher accumulation of PHB in the  $\Delta\text{phaP1 } \Delta\text{phaP4}$  strain might be explained by a high PhaR sequestration at the PhaP1- and PhaP4-devoid PHB granule in formation, which might promote higher expression of PHB biosynthetic genes before the PhaC2/PhaC1 ratio reaches its critical value to inhibit PHB synthase activity and free PhaR begins to increase. The presence of one large granule per cell indicates that each granule grows rapidly within a single cell cycle.

In addition to its action upon *phaC2* expression, FixK<sub>2</sub> was reported to stimulate the expression of blI3998 (a succinate-semi-

aldehyde dehydrogenase) and blr4655 (a phosphoenolpyruvate synthase) (26). The succinate-semialdehyde dehydrogenase, which catalyzes the oxidation of succinate-semialdehyde to succinate using NADP<sup>+</sup> as an electron acceptor, was characterized in a *sucA* mutant of *B. diazoefficiens* as part of a bypass of the tricarboxylic acid (TCA) cycle (51). The phosphoenolpyruvate synthase is a key enzyme of gluconeogenesis and may participate in the secretion of excess carbon as EPS. Thus, induction of fixK<sub>2</sub> under permissive conditions may increase the provision of NADPH to PHB synthesis while this pathway is active or may contribute to the switch of the carbon flow to gluconeogenesis when PHB synthesis is arrested by PhaP coverage of PHB granules and PhaC2 relative increased participation in the PHB synthase dimers (Fig. 5). This hypothesis might explain the inhibition of PHB synthesis and the increase in EPS content observed with the *phaR* mutants in *E. meliloti* (16), *R. etli* (15), and in this work. If *phaR* is knocked out, the expression of *phaC2* and *phaP* becomes uncontrolled, and therefore, the accumulation of their products would lead to rapid inhibition of PHB synthesis and granule growth. In parallel, fixK<sub>2</sub> expression is induced, which would stimulate the expression of the succinate-semialdehyde dehydrogenase and the phosphoenolpyruvate synthase. The precursors provided by the first enzyme could not be used for PHB synthesis because it is arrested and, as gluconeogenesis may be stimulated by the second enzyme, those precursors could be channeled to the synthesis and secretion of EPS.

In symbiosis, the phenotype observed here also indicated that

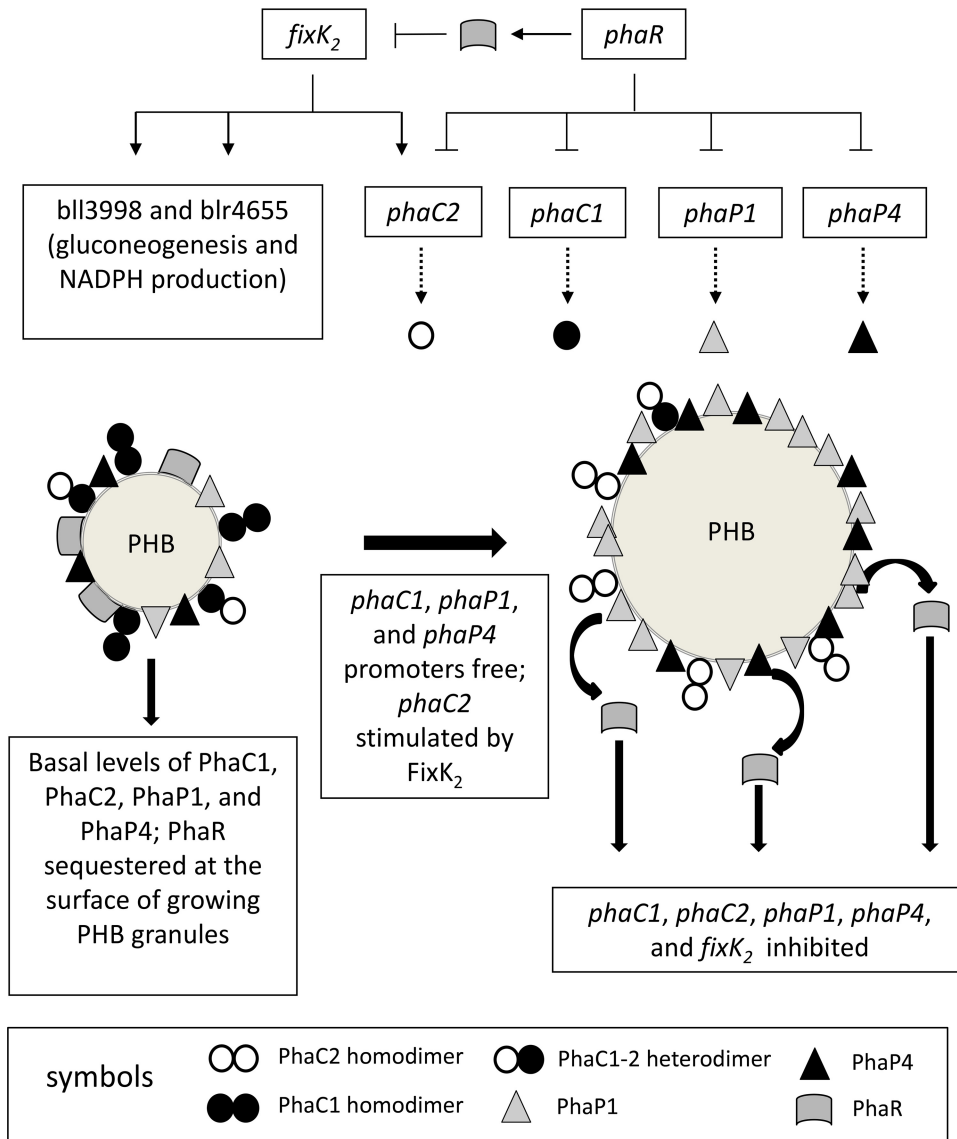


FIG 5 Proposed model for the regulation of PHB synthesis in *B. diazoefficiens*, including the possible role of FixK<sub>2</sub> (→, stimulation; ⊥, inhibition; solid lines indicate transcription; dashed lines depict translation). Permissive conditions, e.g., low O<sub>2</sub>, lead to an increase of NADPH and acetyl-CoA concentrations; thus, PHB biosynthesis starts using these metabolites as the substrates. This biosynthetic process is catalyzed initially by high levels of 3-ketoacyl-CoA thiolase, acetoacetyl-CoA reductase, and PHB synthase due to PhaR sequestration at the PHB granule surface. Meanwhile, *phaC2* expression is stimulated by FixK<sub>2</sub>, increasing the relative PhaC2/PhaC1 concentration, which progressively decreases total PhaC activity during PHB granule growth. As granules grow, less PHB surface becomes available for PhaR and PhaP, which may provoke a competitive displacement of PhaR by PhaP from the surface that in turn may repress PhaR-controlled promoters. In the *phaR* mutant, these promoters are not repressed; therefore, PhaP1, PhaP4, and FixK<sub>2</sub> levels are increased. Hence, FixK<sub>2</sub> stimulates the expression of *phaC2* and the biosynthesis of EPS. Altogether, these changes result in low levels of PHB in the *phaR* mutant. Moreover, in the  $\Delta$ *phaP1*  $\Delta$ *phaP4* strain, the very large PHB granule devoid of PhaP1 and PhaP4 may be able to allocate more PhaR than the wild type at the beginning of PHB biosynthesis. This PhaR sequestration may promote more PHB biosynthetic activity, leading to higher PHB accumulation before the PhaC2/PhaC1 ratio reaches its critical value and PhaR begins to accumulate in the cytoplasm. Protein symbols are indicated at the bottom of the figure. For more details, see the text.

*phaR* has a pleiotropic function. Previously, we observed that the lack of PHB in *B. diazoefficiens*  $\Delta$ *phaC1* strains led to decreased competitiveness for nodulation but had no effect on dry matter accumulation in plants cultured without combined nitrogen, indicating that N<sub>2</sub> fixation was not affected (3). In contrast, here we observed an increase in competitiveness for nodulation of the *phaR* mutant grown under permissive conditions for PHB synthesis as well as an increase in dry matter accumulation in plants inoculated with the *phaR* mutant and cultured under N<sub>2</sub>-fixing

conditions. Because the *phaR* mutant bacteroids also had no PHB granules in their cytoplasm, the comparison of the results obtained with *phaC1* and *phaR* mutants indicated that the symbiotic phenotypes of the *phaR* mutants are not related only to PHB depletion. It is possible that in  $\Delta$ *phaC1* strains, the cytoplasmic levels of PhaR are high because of the absence of PHB granules, and in this way, the expression of the *fixK2* regulon, which includes the regulators *rpoN1*, *fixK1*, and *nnrR* (26), might be repressed by PhaR. In contrast, derepression of *fixK2* in the *phaR* mutants



would favor the expression of these genes, and in particular, it might lead to higher N<sub>2</sub> fixation via the expression of the alternative sigma factor RpoN<sub>1</sub> (26). Nevertheless, we should not rule out an active role of EPS in the symbiotic performance of the *phaR* mutant due to its higher production in this strain.

Thus, the present results confirm the role of *phaR* as a global regulator in both free-living and symbiotic states, suggesting that this role may be exerted at two levels: directly on PHB synthesis and indirectly on other pathways through regulation of the transcription factor *fixK*<sub>2</sub>.

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