The transcriptional regulator NtrC controls glucose-6-phosphate dehydrogenase expression and polyhydroxybutyrate synthesis through NADPH availability in *Herbaspirillum seropedicae*

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The NTR system is the major regulator of nitrogen metabolism in Bacteria. Despite its broad and well-known role in the assimilation, biosynthesis and recycling of nitrogenous molecules, little is known about its role in carbon metabolism. In this work, we present a new facet of the NTR system in the control of NADPH concentration and the biosynthesis of molecules dependent on reduced coenzyme in *Herbaspirillum seropedicae* SmR1. We demonstrated that a ntrC mutant strain accumulated high levels of polyhydroxybutyrate (PHB), reaching levels up to 2-fold higher than the parental strain. In the absence of NtrC, the activity of glucose-6-phosphate dehydrogenase (encoded by zwf) increased by 2.8-fold, consequently leading to a 2.1-fold increase in the NADPH/NADP⁺ ratio. A GFP fusion showed that expression of zwf is likewise controlled by NtrC. The increase in NADPH availability stimulated the production of polyhydroxybutyrate regardless the C/N ratio in the medium. The mutant ntrC was more resistant to H₂O₂ exposure and controlled the propagation of ROS when facing the oxidative condition, a phenotype associated with the increase in PHB content.

Poly-3-hydroxybutyrate (PHB) is an aliphatic polyester member of the polyhydroxyalkanoates (PHA) family synthesised by some bacteria as carbon and reducing equivalents storage⁴. Usually, bacteria produce PHB under conditions of carbon excess and low levels nitrogen, phosphate and oxygen⁵. At least three enzymes are involved in its synthesis: 3-ketothiolase, acetoacetyl-CoA reductase and PHA synthase encoded by phaA, phaB and phaC respectively⁴. These enzymes catalyse the condensation of acetyl-CoA forming acetoacetyl-CoA, then reduction of acetoacetyl-CoA to 3-hydroxybutyryl-CoA (3HB-CoA) and finally polymerisation of 3HB-CoA to yield PHB⁵.

PHB is a thermoplastic biodegradable polymer with physicochemical properties comparable to recalcitrant oil-based plastics such as polypropylene and polystyrene⁶. Although PHB is a sustainable alternative to such plastics, its production cost is still considerably higher, creating a necessity for engineering of PHB overproducer strains and process optimisation. So far, several studies of metabolic engineering for improving microbial PHB production have been reported⁷⁻¹¹. The majority of these studies focused on carbon metabolism pathways engineering and the improvement of NAD(P)H availability. Although the carbon to nitrogen (C/N) ratio is a major factor controlling PHB accumulation in several bacteria¹², little attention has been given to re-engineering the nitrogen metabolism. In bacterial cultivation for the production of PHB, the addition of high ammonium concentration to the medium improves the cell growth but reduces the production of PHB¹³. The negative effect is due to that with a high concentration of ammonium the bacterium diverts much of the carbon skeleton to produce...
C/N ratio is one of the key factors controlling PHB accumulation in bacteria. Accordingly, we anticipated that low C/N ratio leads to the uridylylation of GlnB (GlnB-UMP) by GlnD. GlnB-UMP stimulates the deuridylylation of GlnB, turning GlnB active. The 2-oxoglutarate (2-OG) is an effector of PhaP, and its concentration is high under nitrogen-limiting conditions.

Figure 1. The NTR system and its regulation. In nitrogen excess, high glutamine concentration activates GlnD, stimulating the dephosphorylation of NtrC by NtrB and hence the transcriptionally active form of NtrC. On the other hand, when the bacterium faces a nitrogen-limiting condition, the low glutamine concentration leads to the uridylylation of GlnB (GlnB-UMP) by GlnD. GlnB-UMP stimulates the deuridylylation of GlnB-AMP, turning GlnB active. Therefore, a better understanding of the carbon and nitrogen metabolism interrelationships can contribute to engineering better PHB producers. 

**Results**

The *ntrC* mutant of *H. seropedicae* SmR1 produces more PHB than the parental strain. The C/N ratio is one of the key factors controlling PHB accumulation in bacteria. Accordingly, we anticipated that mutant strains of the NTR system would be a useful tool to investigate its involvement in PHB synthesis. We measured the content of PHB in the *ntrC* mutant of *H. seropedicae* SmR1 and found that it produces more PHB than the parental strain. The authors found that the *ntrC* mutant could produce PHB in both low- and high-C/N ratio media, while the wild-type had no significant PHB production in low-C/N ratio. Also, mutants in both PII proteins genes (*glnB* and *glnZ*) or *glnB* had higher contents of PHB than the wild type under low-C/N ratio. These findings indicate that PHB synthesis is coupled with the nitrogen levels via the components of the NTR system. To better understand how the NTR system affects PHB production in bacteria, we investigated the PHB synthesis in NTR mutants of *H. seropedicae* SmR1. The results obtained in this work reveal a new facet of how the NTR system can influence the carbon metabolism, especially in the synthesis and accumulation of PHB.
late stationary phase (OD$_{600}$ 1.6). The PHB contents of the parental, $glnK$ and $glnD$ strains were very similar since their maximum PHB content was around 25%/cdw (cell dry weight) in high C/N (Fig. 2). The PHB was reduced to 16%/cdw when the strains grew in low C/N. Interestingly, the $glnB$ and $amtB$ mutants presented a significant reduction of PHB accumulation compared to the parental strain (Fig. 2). This effect was remarkable in low C/N where $glnB$ and $amtB$ mutants produced only half of the PHB level of the parental strain. In contrast, PHB production in the $ntrC$ mutant was unregulated and presented the highest content of PHB amongst all strains analysed, reaching 32% of PHB regardless the nitrogen level in the medium suggests that $ntrC$ knock-out decouples the synthesis of PHB of the C/N ratio.

The $ntrC$ mutant also produces more PHB with monosaccharides as carbon source. As malate is not a conventional substrate for biotechnological applications due to its high price compared to other sources, we measured PHB accumulation in the $ntrC$ mutant on D-glucose and D-xylose, which are monosaccharides highly abundant in cheap feedstocks and agro-industrial residues. The C/N ratios were the same. When grown in the presence of D-glucose, the maximum PHB production of the $ntrC$ mutant was 1.8- and 1.7-fold higher than the parental production, in low and high C/N ratio, respectively (Fig. 3A and D). On D-fructose, the maximum PHB production of the $ntrC$ mutant was 2.0- and 2.15-fold higher than the parental production, in low and high C/N ratio respectively (Fig. 3B and E). Interestingly, when D-xylose was used as the carbon source, in low C/N ratio there was no difference of PHB between both strains, but in high C/N ratio, the $ntrC$ mutant produced 1.5-fold more PHB (Fig. 3C and F). Table 1 shows the maximum content of PHB and the productivities obtained for both strains in the different conditions assayed. The profile of sugar consumption and the data of PHB concentration (g/L) and yield of g PHB/g of substrate for the SmR1 and $ntrC$ strains are shown in the supplemental information (Figure S1 and Tables S1 and S2).
Complementation of the ntrC mutant restores PHB production to the parental level. The ntrC gene is clustered in an operon downstream from glnA and ntrB. The ntrC mutant was complemented through pKRT1 conjugation. The pKRT1 is a pLAFR3-derivative containing a glnAntrBC operon copy from H. seropedicae SmR1. PHB accumulated in strains harbouring pKRT1 was reduced when grown in malate or glucose, regard-less the C/N ratio applied (Fig. 4). Therefore, the complementation of the ntrC mutant demonstrates that the NtrC is directly involved in the higher PHB production observed.

The derepression of the zwf (glucose-6-phosphate dehydrogenase) gene results in high NADPH in the ntrC mutant. Previous works have shown that NtrC is involved in the regulation of the expression of glucose-6-phosphate dehydrogenase (G6PDH) and glutamate dehydrogenase (GDH). Specifically, the G6PDH activity is widely implicated in the generation and maintenance of the NADPH/NADP⁺ ratio. The activity of the NADP⁺-dependent malic enzyme (ME) is also correlated with NADPH/NADP⁺ balance. This observation led us to determine the activity of these enzymes in the parental strain SmR1 and ntrC mutant. The G6PDH and NADP⁺-dependent ME activities were 2.3- and 1.6-fold higher in the ntrC mutant when grown in NFb-glucose with 20 mM NH₄Cl (Fig. 5A and C). GDH activity did not differ statistically between both strains (Fig. 5B). The specific activities of G6PDH, ME and GDH enzymes for the SmR1 and ntrC strains are shown in the Supplemental Figure S2. The repression of zwf transcription by NtrC has been previously reported for

Figure 3. PHB accumulation profiles of H. seropedicae SmR1 and ntrC mutants in D-glucose, D-fructose and D-xylose as sole carbon sources. Strains grew in NFbHP medium amended with 25 mM D-glucose (A and D), 25 mM D-fructose (B and E) or 30 mM D-xylose (C and F). Low C/N ratio media had 20 mM NH₄Cl (blue circles), while high C/N ratio media had 5 mM NH₄Cl (red circles). PHB contents were determined in three independent samples. Strains were cultivated at 30 °C at 120 rpm (orbital shaking).
Table 1. Maximal PHB productivities of the parental strain SmR1 and the ntrC mutant of *H. seropedicae* SmR1.

| Strain   | Carbon Source | Parameter | Nitrogen concentration (NH₄Cl) |
|----------|---------------|-----------|---------------------------------|
|          |               |           | 5 mM                              | 20 mM                          |
| SmR1     | Glucose       | CDW (mg/Lh) | 30.6                              | 38.3                            |
|          |               | PHB (mg/Lh) | 12.4                              | 11.5                            |
|          |               | % PHB      | 40.4 ± 1.8                        | 30.1 ± 0.7                      |
|          | Fructose      | CDW (mg/Lh) | 28.9                              | 36.1                            |
|          |               | PHB (mg/Lh) | 8.8                               | 9.0                             |
|          |               | % PHB      | 30.5 ± 1.8                        | 25.2 ± 0.7                      |
|          | Xylose        | CDW (mg/Lh) | 37.0                              | 36.9                            |
|          |               | PHB (mg/Lh) | 14.7                              | 10.8                            |
|          |               | % PHB      | 40.4 ± 1.8                        | 30.1 ± 0.7                      |
| ntrC     | Glucose       | CDW (mg/Lh) | 53.3                              | 47.3                            |
|          |               | PHB (mg/Lh) | 27.3                              | 21.1                            |
|          |               | % PHB      | 51.2 ± 0.9                        | 44.5 ± 0.2                      |
|          | Fructose      | CDW (mg/Lh) | 47.4                              | 55.7                            |
|          |               | PHB (mg/Lh) | 31.2                              | 29.2                            |
|          |               | % PHB      | 65.8 ± 0.7                        | 52.3 ± 1.0                      |
|          | Xylose        | CDW (mg/Lh) | 29.5                              | 38.0                            |
|          |               | PHB mg/Lh  | 17.6                              | 11.5                            |
|          |               | % PHB      | 59.8 ± 0.2                        | 30.1 ± 0.5                      |

Figure 4. PHB content of ntrC mutant strain carrying an additional copy of the operon *glnAntrBntrC* of *H. seropedicae* SmR1. Strains harbouring the plasmids pLAFR3 (empty vector) or pKRT1 (pLAFR3 with the *glnAntrBntrC* as an insert) grew in NFbHP medium amended with 25 mM D-glucose. Graphs A and C represent the data for SmR1, while B and D for the ntrC mutant at low and high C/N ratio, respectively. Red symbols are non-complemented strains, while blue symbols correspond to the complemented ones. Data represent the average ± standard deviation of three independent samples.
the bacterium Pseudomonas putida\(^3,30\). The zwf of H. seropedicae is located downstream from pgi (Hsero1099, encoding a phosphoglucoisomerase) and upstream from a gene encoding a transcriptional regulator of the HexR family (Hsero1097) and talB (Hsero1096, encoding a transaldolase). Since the 76 bp region between pgi and zwf seems not to contain a promoter region, the upstream region of pgi was cloned with GFP in the pEKGFP01 to determine the expression profile of the operon carrying zwf in H. seropedicae SmR1 and ntrC mutant. The activity of the Ppgi-gfp fusion in the ntrC mutant was higher than that of the parental strain during all growth phases, achieving a maximum difference of 2.8-fold (Fig. 5D). The transcription of zwf and other genes involved in the Entner-Doudoroff pathway and in the PHB metabolism were compared between the SmR1 and ntrC strains.
through RNA-seq analysis. The data corroborate the higher expression of zwf in the ntrC mutant (Table S3). Also, the transcription of genes involved in the PHB metabolism was lower in the ntrC mutant (Table S4), indicating that the higher PHB production measured is a consequence of a metabolic factor. This prompted us to measure the NAD(P)H/NAD(P)+ ratio in both strains. The NADH/NAD+ ratio had no statistically significant difference between the parental strain SmR1 complemented or not with an additional copy of the operon glaNtrBntrC cloned into pLAFR3 (pKRT1) (Fig. 5E). The same was observed for the NADH/NAD+ ratio in the ntrC mutant (Fig. 5F). However, the NADPH/NADP+ ratio was statistically higher for both strains when they were not complemented with pKRT1. For the ntrC mutant, the NADPH/NADP+ ratio was 2.1-fold greater than in the parental strain (Fig. 5E and F, comparing the red bars). Taken together these results indicate that the knock-out in ntrC increases the expression of G6PDH and the regeneration of NADP+ in NADPH.

The ntrC mutant of H. seropedicae SmR1 is more resistant to oxidative stress caused by hydrogen peroxide. A high NADPH/NADP+ ratio stimulates PHB production35,36. High [NADPH] increases the flux towards PhaB (NADPH-dependent acetoacetyl-CoA reductase), yielding PHB not only as a carbon stock but also a redox sink35,36. Furthermore, NADPH is crucial to anti-oxidative defences in most organisms, ensuring a reductive cellular environment to mitigate the deleterious effects of oxidative species as hydrogen peroxide, hydroxyl radical and superoxide37. Therefore we determine the resistance to oxidative stress of both strains. Cellular growth was assessed in media amended with increasing concentrations of hydrogen peroxide (H2O2). The addition of the oxidant at 6 h of cultivation impaired the growth of the parental strain in concentrations above 0.2 mM, while the ntrC mutant resisted the oxidative shock, recovering growth after one hour even at 5 mM of H2O2, as observed in Fig. 6B. The serial dilution onto agar plates containing H2O2 showed that the ntrC mutant was able to grow in dilution of 104-fold while the parental SmR1 growth only until 102-fold dilution at 0.1 mM H2O2 (Fig. 6C, lower panel). The ROS measurement applying the fluorescent probes H2-DCDFDA showed that the ntrC mutant was able to control ROS propagation, while the parental strain could not maintain the ROS at low levels (Fig. 6D). The fluorescence value of the oxidised H2-DCDFDA increased 2.5-fold from 0 to 2 mM of H2O2 in the parental SmR1, while in the ntrC mutant the oxidation of the probe increased 1.3-fold applying the same treatment with H2O2. Taken these results together, we conclude that the ntrC mutant has a more efficient defence against the oxidative stress, probably due to the NADPH accumulation which can be applied to mitigate the deleterious effects of the oxidative insult. The high NADPH generation also stimulated the PHB synthesis observed in the mutant.

Discussion

The nitrogen level in the growth medium is a key factor interfering with PHB production in bacteria38–40. Therefore, the interruption of nitrogen regulatory systems can be a useful strategy to reduce the C/N ratio effect on PHB production and ultimately to improve PHB production. Since the NTR system is the master regulator of the nitrogen metabolism in several bacterial species41, wherein we investigate the PHB accumulation using as a model the bacterium H. seropedicae SmR1 and a set of mutants defective in the expression of regulatory proteins of the NTR system. Among all strains evaluated only the ntrC mutant produced higher PHB contents than the parental strain. The ntrC mutant produced around the double of PHB than the parental strain (in % PHB/mg of cdw), in all tested conditions. Similar results were also reported to ntrB and ntrC mutants of A. brasilense Sp724,42 suggesting that possibly the effect of NtrC on PHB synthesis is widespread among prokaryotes. Based on these findings, we anticipated some possible explanations for our results.

The synthesis of PHB is largely dependent on high levels of acetyl-CoA and NADPH. Normally, both metabolites are in high intracellular concentrations when the bacterium faces a condition of carbon excess and limitation in another nutrient, such as nitrogen. This condition is permissive for PHB synthesis due to elevating acetyl-CoA level generated by carbon overflow and high NADPH/NADP+ ratio as consequence of nitrogen limitation42,43.

Considering that H. seropedicae ntrC mutant produced more PHB than the parental strain even under unfavourable conditions (low C/N ratio) suggests that NtrC somehow modulating acetyl-CoA and/or NADPH concentrations. A role of NtrC on directly activating transcription of genes involved in PHB synthesis is unlikely since PHB production is higher in the ntrC mutant even under an excess of ammonium, a condition where NtrC is mainly dephosphorylated and therefore inactive22 (Fig. 2A and B). A transcriptomic analysis of the ΔntrC mutant of the PHA-producing bacterium Pseudomonas putida KT2442 brought important elements to understand the link between NtrC and PHA metabolism9. In the ΔntrC strain, the transcription of zwf-1 (encoding glucose-6-phosphate dehydrogenase – locus tag PP1022) and gap-1 (glyceraldehyde-3-phosphate dehydrogenase - PP1009) was upregulated 5.7- and 2.6-fold, indicating that NtrC represses their expression. Herrvás et al. suggested that repression of zwf shows that NtrC controls hexose catabolism in bacteria, likely to prevent a carbon overflow under nitrogen-limiting conditions for growth. In another work, the ntrC deletion in P. putida KT2440 also rendered cells more resistant to oxidative stress44. Furthermore, zwf-1 (PP1022) was also up-regulated in the ΔntrC mutant of P. putida KT2440, as determined by transcriptomic analysis. H. seropedicae SmR1 and P. putida have the same incomplete glycolytic pathway since it lacks 6-phosphofructokinase (PFK-1). As a consequence, the Entner-Doudoroff (ED) pathway converges glucose and fructose into pyruvate and glyceraldehyde-3-phosphate. Particularly, in mutants defective in ntrC expression, it is expected a higher metabolic flux through glucose-6-phosphate dehydrogenase, generating more acetyl-CoA and NADPH, which in turn increases PHB accumulation.

These findings suggest that the up-expression of zwf could be a major factor leading ntrC defective mutants to accumulate more PHB under permissive conditions. The observation that the H. seropedicae ntrC mutant produced more PHB even in unfavourable conditions (low C/N ratio) suggests a higher pool of reducing power as compared to the parental strain. Therefore is likely that the bacterium switches the synthesis of PHB as an electron sink to avoid the deleterious effect of a high reductive environment.
In fact, we have already reported that mutant strains defective in PHB synthesis or accumulation presented a severe growth penalty on glucose\(^1\). Probably, as a consequence of the redox imbalance caused by the inability of those mutants to divert electrons towards PHB synthesis.

NtrC is widely recognised as a transcriptional regulator restricted to regulate genes involved in nitrogen metabolism. Indeed, a significant portion of genes corresponding to 2% of the *E. coli* genome was determined to be under control of the NtrC transcriptional activity, most of them involved in amino acids and ammonium transport, nitrogen assimilation and nitrogen recycling\(^4\). Our results demonstrate that NtrC interferes with PHB synthesis, bringing to the discussion if NtrC could be one of the connection points between nitrogen and carbon metabolism. Alternatively, the NtrC control of the NADPH recycling may help cells dealing with a stressful condition such as nitrogen limitation. In fact, the glutamate synthase (GOGAT) and the glutamate dehydrogenase (GDH), which are important reactions for ammonium assimilation require NADPH as an electron donor to catalyse the amination of 2-oxoglutarate\(^4\). In *E. coli*, NADPH concentration was homeostatic after ammonium upshift\(^4\). Since NADPH consumption increases when ammonium assimilation is higher, it is likely that other pathways are generating the NADPH demand\(^4\). Whether or not the phosphorylation state of NtrC interferes in NADPH production through *zwf* expression is uncertain. The repression of *gdhA* encoding the glutamate dehydrogenase in *P. putida* KT2442 by NtrC was shown to be independent of the phosphorylation state, since both

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Figure 6. The effect of H\(_2\)O\(_2\) on growth and ROS propagation in the parental strain and the *ntrC* mutant of *H. seropedicae* SmR1. Growth kinetics of parental strain SmR1 (A) and *ntrC* mutant (B) in NFbHP-malate medium in the presence of increasing hydrogen peroxide concentrations. The growth kinetics were determined in three independent samples for each strain. At the time of 6 h hydrogen peroxide was added to the cultures. (C) SmR1 and *ntrC* mutant grew to stationary phase and serial dilutions were plated onto NFbHP-malate medium containing H\(_2\)O\(_2\) at the indicated concentrations. After two days of incubation at 30 °C, the SmR1 strain presented higher hypersensitivity to H\(_2\)O\(_2\) than the *ntrC* mutant. (D) Median fluorescence of H\(_2\)-DCFDA treated cultures during H\(_2\)O\(_2\) stress in SmR1 and *ntrC* strains. The median fluorescence was determined in three independent samples for each strain. Where appropriate, statistical significance is shown (* p-value ≤ 0.05, independent two-sample t-test).
of NH\textsubscript{4}Cl as a nitrogen source. The microplate was incubated in an orbital shaker (Incubator Shaker Series I26, 160 rpm). E. coli strains grew in LB medium at 37 °C and shaken at 180 rpm.

Promoter, 2\textsuperscript{pgi} pBBR1MCS-3 vector were inoculated in a 96-well microplate containing 200 μL of saturated cultures containing either the transcriptional fusion pEKGFP01 or an empty pBBR1MCS-3 vector were inoculated in a 96-well microplate containing 200 μL of NFbHP-glucose with 20 mM of NH\textsubscript{4}Cl as a nitrogen source. The microplate was incubated in an orbital shaker (Incubator Shaker Series I26, 160 rpm).

Table 2. Bacterial strains and plasmids used in this work.

| Strain or plasmid | Relevant characteristics | Reference/source |
|-------------------|--------------------------|------------------|
| E. coli Top10     | Cloning host             | Thermo Fisher Scientific |
| S17-1             | Conjugational transfer of plasmids | 45 |
| H. seropedicae SmR1 | Parental strain, Nif\textsuperscript{+}, Sm\textsuperscript{R} | 46 |
| ntrC(DCP286A) SmR1 | containing ntrC::Tn5-b20 | 23 |
| glnB SmR1         | containing glnB::Tc\textsuperscript{R} | 47 |
| glnK SmR1         | with a chromosomal deletion of the glnK | 47 |
| glnD SmR1         | containing glnD::Tc\textsuperscript{R} | Unpublished result |
| amtB SmR1         | containing amtB::lacZ::Km\textsuperscript{R} | 47 |

Plasmids

pBBR1MCS-3

pLAFR3 Broad-host-range cloning vector, IncP1, Tc\textsuperscript{R} 29

pKRT1 23 kb fragment from H. seropedicae SmR1 harbouring the glnA, ntrB and ntrC genes 46

pEK07 pBBR1MCS-3 containing pgi upstream region (P\textsubscript{pgi}) from H. seropedicae SmR1 cloned between Xhol and SpeI restriction sites This work

BBa_113504 Plasmid pSB1C3-derivative containing the gfpmut3b gene (BBa_E0040) with the strong rbs (BBa_B0034) and the double terminator (BBa_B0015). Registry of Standard Biological Parts (partsregistry.org)

pEKGFP01 pBBR1MCS-3 containing pgi upstream region (P\textsubscript{pgi}) from H. seropedicae SmR1 and gfpmut3b downstream. This work

Table 2. Bacterial strains and plasmids used in this work.

wild-type and the mutant NtrC\textsuperscript{DD55E,S161F} (mimicking the phosphorylated protein) can bind to the gdhA promoter and repress transcription\textsuperscript{31}. The effect of phosphorylation could be further studied employing a ntrB mutant of H. seropedicae and complementation of the ntrC mutant with NtrC variants unable to be phosphorylated. Such experiments will determine if the reduction in PHB production observed in the glnB and amtB mutants derived from the phosphorylation state of NtrC.

It would be interesting to investigate further the role of NtrC on PHB synthesis in other PHB-producing models, such as Ralstonia eutropha, Azotobacter vinelandii and P. putida. It would point if the involvement of NtrC on PHB synthesis is conserved among other classes of bacteria, serving as a strategy for metabolic engineering aiming to improve PHB production.

Methods

Bacterial Strains, Plasmids, and Growth Conditions. Strains and plasmids used are listed in Table 2. Escherichia coli strain Top10 (Thermo Fisher Scientific Inc., Waltham, MA, USA) and E. coli S17-1\textsuperscript{45} were used for cloning and conjugational procedures, respectively. E. coli strains grew in LB medium at 37 °C and shaken at 160 rpm. H. seropedicae parental strain SmR1\textsuperscript{46} and mutant strains were cultured in NFbHP media with 37 mM DL-malate and the indicated concentration of NH\textsubscript{4}Cl at 30 °C and shaken at 120 rpm\textsuperscript{49}. D-glucose at 25 mM, D-fructose at 25 mM and D-xylose at 30 mM were applied as alternative carbon sources for PHB production measurement as a replacement for malate. Two regimes of carbon-to-nitrogen (C/N) ratio were used at the start of cultivation: high C/N ratio with 5 mM of NH\textsubscript{4}Cl and low C/N with 20 mM of NH\textsubscript{4}Cl.

Quantification of PHB. PHB was quantified by methanolysis, followed by GC-FID (gas chromatography coupled to a flame ionisation detector) analyses as described previously\textsuperscript{50} with 5 to 10 mg of lyophilised bacteria. Amounts of PHB in each sample were normalised to cell dry weight (cdw; the weight of the lyophilised bacterial pellet) and expressed as % of PHB cell dry weight\textsuperscript{−1}.

Construction of P\textsubscript{pgi-gfp} transcriptional fusion. The intergenic region of the pgi (locus-tag Hsero1099) was amplified using the primers Fw\textsubscript{pgi-Hs} 5’TATTCCTGGGTGTGGTGCGGTTCTGGTATACGG 3’, containing a XhoI site (underlined) and Rev\textsubscript{pgi-Hs} 5’TATACTAGTCATATGGGGTCTTGCGTGTCGGTCGGG 3’, containing a SpeI site (underlined) as previously described\textsuperscript{31}. The amplified product was cloned into the sites XhoI and SpeI of the pBRR1MCS-3\textsuperscript{52}, generating the pEK07. The reporter gene gfp containing upstream the rbs site B0034 and downstream the double terminator B0015 was extracted from the plasmid BBa_113504 ( Registry of Standard Biological Parts, partsregistry.org) digested with the EcoRI and SpeI enzymes and cloned into pBlueScript II KS\textsuperscript{+} digested with EcoRI and Xbal. Then, the gfp cassette was removed by digestion with the XbaI and SacI enzymes and cloned into the pEK07 digested with the same enzymes, generating the pEKGFP01. The pEKGFP01 was transformed in E. coli S17-1 and conjugated to H. seropedicae by bi-parental mating.

Measurement of P\textsubscript{pgi-gfp} transcriptional activity. To measure the transcriptional activity of the pgi promoter, 2 μL of saturated cultures containing either the transcriptional fusion pEKGFP01 or an empty pBBR1MCS-3 vector were inoculated in a 96-well microplate containing 200 μL of NFbHP-glucose with 20 mM of NH\textsubscript{4}Cl as a nitrogen source. The microplate was incubated in an orbital shaker (Incubator Shaker Series I26,
New Brunswick™) at 30 °C and 120 rpm. The fluorescence was measured using a Berthold™ TriStar LB 941 using 355 nm filter for excitation and a 535 nm for the emission wavelength. Arbitrary fluorescence units were normalized by OD readings at 600 nm using a Bio-Rad iMark™ Microplate Reader.

**Complementation of ntrC mutant.** The pKRT1 cosmids was conjugated by bi-parental mating between E. coli S17-1 and H. seropedicae strains. The transconjugant colonies were selected in NFbHP-malate with 20 mM NH₄Cl agar with 10 μg/mL of tetracycline. The complemented strains were cultivated in NFbHP-malate and NFbHP-glucose at high and low-C/N ratios.

**Determination of enzymatic activities.** Glucose-6-phosphate dehydrogenase (G6PDH), glutamate dehydrogenase (GDH) and malic enzyme (ME) activity assays were performed by measuring the formazan production at 585 nm, as previously described. Formazan is the insoluble product formed by MTT (3-(4,5-dime-thyl-thiazolyl-2)-2,5-diphenyl-tetrazolum bromide) reduction. Cells were lysed by sonication on an ice bath. The lysates were centrifuged at 4,000 × g for 10 min at 4 °C. The supernatants were maintained on ice until assay setup. The reaction contained 300 μM NADP⁺, 300 μM of the substrate (D-glucose-6-phosphate, DL-malate or L-glutamate), 300 μM MTT and 30 μM PES (phenazine ethosulphate). The components were diluted up to 900 μL in 50 mM Tris-HCl buffer at pH 8 with 0.13% (m/v) gelatin. The gelatin was employed to prevent the formazan precipitation. The reactions were carried out in 1 mL-cuvettes, starting by addition of 100 μL of the supernatant. The reactions were monitored for 5 min at 585 nm in a Shimadzu™ spectrophotometer. The activity was expressed as μmol of formazan/min of reaction per OD₆₀₀ of the culture.

**Determination of NAD(P)H/NAD(P)⁺ ratio.** Intracellular levels of NAD⁺, NADP⁺, NADH and NADPH were determined by the improved cyclic assay using either ADH (Sigma #A3263) or G6PDH (Sigma #G6378), respectively. The dinucleotides were extracted using cell pellets from 1 mL of culture, cultivated either until the mid-log (OD₆₀₀ of 0.4–0.5) or late-log (OD₆₀₀ of 1.0–1.2) phases. Reduced and oxidised nicotinamide adenine dinucleotides were differentially extracted by treatment with alkali or acid, respectively, followed by extract neutralisation. The assays were performed in 200 μL in a water bath for 30 minutes at 37 °C containing the following components: 0.1 M Tricine–NaOH buffer (pH 8.0); 4.2 mM MTT; 40 mM EDTA (disodium salt); 16.6 mM PES; 5 mM ethanol as substrate for alcohol dehydrogenase to determine NADH/NAD⁺ or 25 mM glucose 6-phosphate (dipotassium salt) as substrate for G6PDH to determine NADPH/NADP⁺. To determine NADPH/NADP⁺ and NADH/NAD⁺, 10 μL of a baker’s yeast G6PDH solution (14 units/mL) or 10 μL of a baker’s yeast alcohol dehydrogenase solution (100 units/mL) were added per reaction, respectively. Reactions were stopped by adding 100 μL of 5 M NaCl followed by 5 minutes of ice incubation. The precipitated formazan was centrifuged for 5 min at 14,000 x g and solubilised in 500 μL of 96% ethanol. The formazan was quantified as a function of the absorbance at 550 nm of 200 μL of sample in 96-well plates in a Biotek ELX-800 microplate reader. The standard calibration curve was run in triplicate using up to 30 pmol/assay of either NAD(P)H or NAD(P)⁺ standards.

**Analysis of intracellular ROS levels using flow cytometry.** Cells from 1 mL of culture were collected by centrifugation at 14,000 × g for 1 min and then resuspended in 500 μL of PBS buffer supplemented with 1 mM EDTA, 0.01% Tween 20 and 0.1% Triton X-100. Cells were subsequently incubated with 50 μM 2′,7′-dichlorofluorescein diacetate (H₂DCFDA) for 30 min at 30 °C in the dark. Control experiments without H₂DCFDA addition were also set up under the same conditions. Treatment with H₂O₂ was performed by pre-incubation of cells with increasing concentration of H₂O₂, for 30 min at 120 rpm and 30°C, before addition of H₂DCFDA. The samples were analysed by flow cytometry using a BD Accuri™ C5 flow cytometer equipped with a 488 nm argon laser and a 533/30 nm bandpass filter (FL1-H). The median fluorescence intensity was used to determine the intracellular ROS levels.

**Data availability.** The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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**Author Contributions**

E.N.M.S., E.Y.S.K. and H.L.R.C. designed, performed the experiments and analysed the data. M.M.S., L.S.C., E.M.S., and F.O.P. conceived, supervised the study and analysed the data. M.M.S. wrote the manuscript. All authors reviewed the manuscript.

**Additional Information**

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