# Transcriptional and translational adaptation to aerobic nitrate anabolism in the denitrifier *Paracoccus denitrificans*

Victor M. Luque-Almagro<sup>1</sup>, Isabel Manso<sup>1</sup>, Matthew J. Sullivan<sup>2</sup>, Gary Rowley<sup>2</sup>, Stuart J. Ferguson<sup>3</sup>, Conrado Moreno-Vivián<sup>1</sup>, David J. Richardson<sup>2</sup> Andrew J. Gates<sup>2</sup>, M. Dolores Roldán<sup>1\*</sup>

<sup>1</sup>Departamento de Bioquímica y Biología Molecular, Universidad de Córdoba, Edificio Severo Ochoa, 1<sup>a</sup> planta, Campus de Rabanales, Córdoba, 14071, Spain. <sup>2</sup>School of Biological Sciences, University of East Anglia, Norwich Research Park, Norwich, NR4 7TJ, UK. <sup>3</sup>Department of Biochemistry, University of Oxford, South Parks Road, Oxford, OX1 3QU, UK.

\* Corresponding author address:
Departamento de Bioquímica y Biología Molecular
Universidad de Córdoba
Córdoba 14071
Spain
Tel: +34 957218318
Fax: +34 957218592
E-mail: bb2rorum@uco.es

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#### SUMMARY STATEMENT

Transcriptomic, proteomic and mutational analyses of the nitrate assimilatory system in *P. denitrificans* PD1222 reveals that nitrate exerts a very tight regulation at the transcriptional and translational levels through the NtrBC and the NasTS systems.

#### **AUTHOR CONTRIBUTION**

Matthew J. Sullivan (MJS) and Gary Rowley (GR) contributed to the microarray construction under the supervision of David J. Richardson (DJR), Stuart J. Ferguson (SJF) and Andrew J. Gates (AJG). Proteomic analysis and generation of mutant strains were carried out by Victor M. Luque-Almagro (VLA) under the supervision of Conrado Moreno-Vivián (CMV) and M. Dolores Roldán (MDR). RT-PCR and qPCR was performed by Isabel Manso (IM) under the supervision of CMV, MDR and AJG. The final version of this manuscript has been written by MDR and discussed and revised in agreement with DJR, CMV and SJF.

# 1 ABSTRACT

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3 Transcriptional adaptation to nitrate-dependent anabolism by *Paracoccus* 4 denitrificans PD1222 was studied. A total of 74 genes were induced in cells grown 5 with nitrate as N-source compared to ammonium, including *nasTSABGHC* and 6 *ntrBC* genes. The *nasT* and *nasS* genes were cotranscribed, although *nasT* was more strongly induced by nitrate than nasS. The nasABGHC genes constituted a 7 8 transcriptional unit, which is preceded by a non-coding region containing hairpin 9 structures involved in transcription termination. The *nasTS* and *nasABGHC* 10 transcripts were detected at similar levels with nitrate or glutamate as N-source, but *nasABGHC* transcript was undetectable in ammonium-grown cells. The nitrite 11 reductase NasG subunit was detected by 2D-PAGE in cytoplasmic fractions from 12 13 nitrate-grown cells, but it was not present when either ammonium or glutamate 14 was used as N-source. The *nasT* mutant lacked both *nasABGHC* transcript and 15 NADH-dependent nitrate reductase activity. On the contrary, the nasS mutant showed similar levels of the *nasABGHC* transcript to the wild-type strain and 16 17 displayed NasG protein and NADH-nitrate reductase activity with all N-sources 18 tested, except with ammonium. Ammonium repression of *nasABGHC* was 19 dependent on the Ntr system. The *ntrBC* and *ntrYX* genes were expressed at low 20 levels regardless of the nitrogen source supporting growth. Mutational analysis of 21 the *ntrBCYX* genes indicated that while *ntrBC* genes are required for nitrate 22 assimilation, *ntrYX* genes can only partially restore growth on nitrate in absence of 23 *ntrBC* genes. The existence of a regulation mechanism for nitrate assimilation in *P*. 24 *denitrificans*, by which nitrate induction operates at both transcriptional and 25 translational levels, is proposed.

26

#### **1 INTRODUCTION**

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3 The soil denitrifier Paracoccus denitrificans PD1222 makes multiple metabolic uses of nitrate: (i) as a respiratory electron acceptor to support anaerobic growth, catalysed by a 4 5 membrane-bound nitrate reductase (Nar); (ii) as an electron sink to dispose of excess 6 reductant during aerobic metabolism of highly reduced carbon substrates, catalysed by the periplasmic nitrate reductase (Nap); and (iii) as nitrogen source for anabolism 7 during both oxic and anoxic growth. The first two processes have been widely studied 8 9 [1], but the biochemistry and regulation of the third process, nitrate assimilation, has 10 received much less attention. The nitrate assimilation system (Nas) is encoded by the nasABGHC gene cluster, which contains a nitrate transporter (NasA), an NADH-11 12 dependent nitrite reductase (NasB), a small Rieske-type protein required for both nitrate and nitrite reduction (NasG), a nitrite transporter (NasH), and a nitrate reductase 13 14 (NasC), which receives electrons from NADH via NasB and NasG [2]. The NasB and 15 NasG proteins are essential for growth with nitrate or nitrite as sole nitrogen source 16 under both aerobic and anaerobic conditions. However, the NasA and NasC proteins are 17 required for nitrate import and reduction only under aerobic conditions because of a 18 functional overlap with biochemical components of the respiratory nitrate reductase 19 system that are synthesised under anaerobic conditions [2,3].

20 Nitrate assimilation is regulated in response to nitrate by a two-component 21 system, encoded by the *nasTS* genes, which are located directly upstream from the *nasA* 22 gene. NasS acts as nitrate/nitrite sensor whereas NasT has an ANTAR (AmiR and NasR 23 transcription antitermination regulators) domain for acting as a transcriptional 24 antiterminator [4,5]. In the absence of nitrate or nitrite, NasS and NasT form an inactive 25 tetrameric complex (two units of each), leading to the premature termination of the 26 nasABGHC gene transcription. In the presence of nitrate or nitrite, the NasTS complex 27 becomes dissociated, and thereby NasT allows expression of nas genes [5]. Recently, 28 the P. dentrificans PD1222 whole genome has been analyzed by using the Quadparser 29 programme, revealing the existence of a guanine-rich region located upstream of the 30 *nasT* gene that forms a canonical G-quadruplex structure. Stabilization of this secondary 31 structure in DNA has been suggested to act as a negative regulator for nitrate-dependent 32 growth [6].

33 Nitrate assimilation is a widespread metabolic capacity in proteobacteria that 34 usually is controlled at the transcriptional level by nitrate and nitrite induction and by 35 ammonium repression [7,8]. In cyanobacteria, the catabolite activator protein (CAP) 36 family transcription factor NtcA represses the nitrate reductase genes in the presence of 37 ammonium, whereas it activates transcription of these genes at a high C/N ratio 38 (nitrogen depletion), reflected by high 2-oxoglutarate levels [9-11]. In some 39 cyanobacteria, the LysR family transcription regulator NtcB is required for the 40 nitrate/nitrite-dependent induction of the nitrate reductase gene [11]. In Gram positive 41 bacteria like Bacillus subtilis and Streptomyces coelicolor the TnrA and GlnR regulators 42 respond to nitrogen starvation [12,13]. In Klebsiella oxytoca, expression of the nitrate 43 assimilation genes is activated under low nitrogen conditions through the global 44 nitrogen regulatory Ntr system, including NtrA, NtrB and NtrC proteins, and by nitrate/nitrite induction through NasR, a transcription antitermination protein that also 45 46 binds to nitrate [14,15]. The crystal structure of the *Klebsiella oxytoca* NasR protein has 47 been solved; it is a dimer with a large N-terminal nitrate and nitrite-sensor (NIT) domain and a C-terminal ANTAR domain necessary for specific binding to leader 48

1 mRNA. The NIT domain binds nitrate and nitrite between two conserved arginine 2 residues located on adjacent helices [16]. In *Azotobacter vinelandii* and *Pseudomonas* 3 *putida*, as well as in *P. denitrificans*, two different proteins NasT and NasS act as 4 transcriptional antiterminator and nitrate sensor, respectively [17-20].

5 The NtrBC two-component system has been extensively characterized in enteric 6 bacteria [21]. NtrB is a sensor kinase that autophosphorylates on a histidine residue 7 under low nitrogen concentrations and transfers a phosphoryl group to the NtrC 8 response regulator protein on a specific aspartate residue [22]. Phosphorylated NtrC acts 9 as a transcriptional activator that oligomerizes on the DNA template with ATPase activity [23]. The NtrC members are usually dependent on the  $\sigma^{54}$  factor (NtrA) and 10 they are involved in transcription of genes related to nitrogen metabolism, such as the 11 12 glutamine synthetase glnA gene. However, in Rhodobacter capsulatus a regulatory twocomponent NtrBC system has been described in which the NtrC component is not  $\sigma^{54}$ -13 14 dependent to activate transcription of nifA1 and nifA2 genes, which code for 15 transcriptional activators that induce nitrogen fixation gene expression, and the glnB 16 gene that is a negative regulator of the R. capsulatus NtrBC system under nitrogen 17 excess [24].

18 The NtrYX system is also a two-component regulatory system with similarity to 19 the sensor/kinase NtrB and the regulatory protein NtrC, respectively. This system has 20 been investigated in diazotrophs with a proposed role in nitrogen assimilation. In 21 Azorhizobium caulinodans, a mutant in the ntrX gene was found to be defective in using 22 nitrate as nitrogen source, and showed also a reduced *nifA* expression under nitrogen 23 fixation conditions with a disturbed symbiotic phenotype. In the ntrC mutant strain, 24 expression of the *ntrYX* operon was derepressed in the presence of nitrate, suggesting an 25 interaction between both NtrBC and NtrYX systems [25]. The NtrBC system in 26 Azospirillum brasilense is involved in regulation of nitrate assimilation, ammonium 27 transport, and nitrogenase switch-off by ammonium. The NtrYX system may be 28 involved in nitrate utilization through a possible substitution of the NtrBC system by the 29 NtrYX sensor-regulator pair [26,27]. Herbaspirillum seropedicae is a diazotrophic  $\beta$ -30 proteobacterium with both NtrBC and NtrYX systems displaying a role in regulation of 31 nitrate assimilation [28]. The photosynthetic bacterium Rhodobacter capsulatus also has 32 both NtrBC and NtrYX systems, with a function of the NtrBC system in urea 33 assimilation and nitrogen fixation, but with an unclear physiological function of the 34 NtrYX system [29]. The NtrYX system of *Brucella* spp. is involved in redox sensing 35 and regulation of denitrification genes. Expression of narGHIJK, nirKV, norBCDEFQ 36 and nosDFLRXYZ genes are down-regulated in an ntrY mutant strain under aerobic and 37 microaerobic conditions, with a marked down-regulation of the *nir*, *nor* and *nos* genes 38 under microaerobic conditions. The Brucella spp. NtrY protein contains one haem 39 group for sensing the oxygen status in the cell and shows its maximal activity as an 40 autohistidine kinase in the ferrous state under low oxygen tension [30,31].

Analysis of the whole genome sequence of *P. denitrificans* reveals the presence of genes encoding NtrBC and NtrYX proteins. In this work, we explore the global transcriptomic changes that underpin the transition from ammonium-dependent to nitrate-dependent aerobic growth, including the relationships within the different twocomponent regulatory systems NasTS, NtrBC and NtrYX. An additional mechanism to the transcriptional regulation of nitrate assimilation, based on a nitrate-dependent translation control of the *P. denitrificans* PD1222 Nas proteins, is also proposed.

# **1 MATERIALS AND METHODS**

2 3

## Bacterial strains, media and growth conditions

4 All strains used in this study are listed in Table S1. P. denitrificans PD1222 was grown 5 at 30 °C in Luria-Bertani (LB) medium [32] or defined mineral salts medium as 6 previously described [33], containing 50 mM succinate as carbon source. Ammonium 7 chloride, potassium nitrate, potassium nitrite or L-glutamate were used as nitrogen 8 source (10 mM each), as stated in the text. Bacteria were cultured in 250 mL flasks 9 containing 50 mL of medium that were rotated at 200 rpm. E. coli strains were grown at 37 °C in LB medium. Cell growth was followed measuring the absorbance of the 10 cultures at 600 nm ( $A_{600}$ ) or by protein determination [34]. Antibiotic supplements were 11 used at the following concentrations (µg·mL<sup>-1</sup>): ampicillin (Amp), 100; kanamycin 12 13 (Km), 25; rifampicin (Rif), 100; spectinomycin (Sp), 25; streptomycin (Sm), 60; 14 tetracycline (Tet), 10; chloramphenicol (Cm), 50.

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#### 16 Analytical methods

17 Upon reaching exponential growth phase, a 50 mL culture volume was subject to cell fractionation. Cells were harvested, washed twice with 50 mM Tris-HCl (pH 8.0) and 18 re-suspended in 10 mM Tris-HCl (pH 8.0), 500 mM sucrose and 3 mM EDTA. 19 Lysozyme (chicken egg white, EC 3.2.1.17) at 0.2 mg $\cdot$ mL<sup>-1</sup> final concentration and a 20 few grains of DNase I (bovine pancreas, EC 3.1.21.1) were added. This mixture was 21 22 incubated with shaking for 30 minutes at 30 °C. The sphaeroplasts formed were 23 harvested by centrifugation (13000 x g) for 15 minutes at 4 °C and the periplasm was 24 recovered. Sphaeroplasts were re-suspended in a volume of 10 mL 100 mM Tris-HCl 25 (pH 8.0) and disrupted by sonication, and cytoplasmic and membrane fractions were 26 separated by ultracentrifugation (40000 x g). Assimilatory NADH-dependent nitrate 27 reductase activity was assayed in cytoplasmic fractions in the presence of NADH as 28 electron donor by measuring the nitrite formed from nitrate [35].  $\beta$ -galactosidase 29 activity was determined spectrophotometrically as previously described [36]. Protein 30 concentration was measured by using a Bradford protein assay kit (Bio-Rad, UK) with a 31 BSA standard (Fraction V, Sigma, UK).

32

# 33 Microarray analysis of P. denitrificans

34 P. denitrificans genomic DNA was isolated using a Genomic DNA extracting kit and 35 100/G columns (Qiagen) from 10 mL exponential phase cells according with the 36 specification of the manufacturer. For RNA extractions, 30 mL of early-exponential 37 phase cells (A<sub>600</sub> ~ 0.4) was added to 12 mL of ice-cold 95% ethanol/5% phenol (v/v) 38 solution, and incubated on ice for 30 minutes to prevent RNA degradation. Cells were 39 then pelleted and stored at -80 °C until RNA was extracted by using SV Total RNA 40 isolation kit (Promega). Trace DNA contamination was removed by treatment with 41 Turbo DNA-free<sup>TM</sup> DNase (Ambion), and this was confirmed by PCR amplification of 42 RNA samples using MyFi<sup>™</sup> DNA polymerase (Bioline). Nucleic acids were quantified 43 spectrophotometrically in a Nanodrop 2000 (Thermo Scientific), and integrity of RNA 44 samples was analyzed using an Experion<sup>™</sup> Automated Electrophoresis platform 45 (BioRad) using RNA StdSens chips (BioRad). All standard protocols were carried out 46 according with the instructions of the manufacturers.

47 For labelling and hybridisation of microarray slides, total RNA (10  $\mu$ g) from 48 three independent bacterial cultures were reverse-transcribed to cDNA using

1 AffinityScript multiple temperature reverse transcriptase (Agilent), and fluorescently 2 labelled using random primers (Invitrogen) to incorporate Cy5-dCTP (Amersham). P. 3 denitrificans genomic DNA (10 µg) was labelled with Cy3-dCTP (Amersham) using a Bioprime<sup>®</sup> DNA labelling system (Invitrogen), prior to mixing (1:5) with labelled 4 cDNA and hybridised to custom-designed 4 x 44K oligonucleotide array slides 5 6 (Agilent). Hybridisation buffer (50 mM morpholine-4-ethanesulfonic acid pH 6.5, 1 M NaCl, 20% w/v formamide, 20 mM EDTA, 1% w/v, Triton-X-100) mixed with Cy5-7 8 and Cy3-dCTP labelled nucleic acids were loaded onto the GASKET slide prior to 9 placing the microarray slide in contact with the hybridization mix, which were sealed in 10 a tight chamber and incubated at 55 °C in a rotary hybridisation oven at 8 rpm, for 11 approximately 60 hours in the dark. Following hybridizations, slides were removed and washed for 5 minutes in a microscope-slide chamber using a solution of 6 x SSPE (0.2 12 13 M phosphate buffer, 2.98 M NaCl, 20 mM EDTA, pH 7.4) supplemented with 0.005% 14 N-lauryl-sarcosine, followed by 5 minutes in a solution of 0.6 x SSPE supplemented 15 with 0.18% polyethylene glycol 200. Slides were then dried by centrifugation for 30 16 seconds.

17 For analyses and interpretation, microarray slides were scanned using a scanner 18 (GenePix 4000A, Axon Instruments) with excitation wavelengths of 532 nm and 635 19 nm. Fluorescent spots and background intensities were quantified using GenePix Pro 20 software (Axon Instruments) and filtered to omit those with a reference signal lower 21 than two standard-deviations from the background intensity in further analyses. Signal 22 intensities were corrected by subtracting the background and the red/green (Cy5/Cy3) 23 ratios. All datasets were normalised using Batch Anti-Banana Algorithm in R (BABAR) 24 which uses cyclic loess to normalise across the complete dataset [37] and analyzed 25 using Gene Spring 7.3 (Agilent) to filter genes that were differentially expressed  $\geq 2$ -26 fold with significance of  $\geq 95\%$  across three independent cultures. These data are 27 represented in Figure 1, where  $\log_2$  of normalised expression values are shown as a 28 heat-map. Microarrays were validated by qPCR with oligonucleotide primers that 29 annealed to internal regions of the nas genes (Table S2), as described below. Primers were designed using Primer<sup>3</sup> Plus software [38], to amplify products between 100 and 30 150 bp, with a  $T_m$  of about 60 °C and used at a final concentration of 0.4  $\mu$ M. The 31 32 relative fold-change values were calculated by using amplification efficiencies, as 33 described previously [39].

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#### 35 **Proteomic analysis**

Two-dimensional gels electrophoresis (2D-PAGE) was performed with sample 36 37 preparations that were obtained from P. denitrificans cells grown to mid-log phase in 38 minimal medium with different nitrogen sources as previously described [2]. 39 Subcellular fractionation was carried as described above, and isolated cytoplasmic 40 fractions containing 250 µg protein were used to rehydrate 11 cm strips (Immobiline 41 DryStrips with the appropriate pH range; Amersham Biosciences) for 12 h. Isoelectric focusing was carried out in an IPGphor (Pharmacia) until 20000 volt-hours were 42 43 reached. After isoelectric focusing, strips were equilibrated as previously described [40] 44 and applied to 12% (v/v) polyacrylamide gels. Second dimension SDS-PAGE was 45 performed by using the Hoefer SE600 system (Amersham Biosciences) and gels were 46 stained using the Brilliant Blue G-colloidal concentrate (Sigma) and scanned with a 47 Molecular Image FX (Bio-Rad). Triplicate 2D-PAGE separations were generated for 48 each sample. Protein identification was carried out in the UCO-SCAI Proteomic Centre,

1 University of Córdoba (Spain), a member of ProteoRed network. Protein spots of 2 interest were excised automatically in a ProPic station (Genomic Solutions, UK) and 3 samples were automatically digested with trypsin according with standard protocols in a 4 ProGest station (Genomic Solutions), and analyzed in a 4700 Proteomics Analyzer MALDI-TOF/TOF mass spectrometer (Applied Biosystems), in the m/z range 800 to 5 4000, with an accelerating voltage of 20 kV, in reflectron mode and with delayed 6 extraction set to "on" and an elapsed time of 120 ns. Proteins were identified by peptide 7 8 mass fingerprinting (PMF) and confirmed by MS/MS analysis of the three most abundant peptide ions. MASCOT searching engine (Matrixscience, UK) was used for 9 10 protein identification over the non-redundant NCBI database of proteins. The 11 confidence in the peptide mass fingerprinting matches (p < 0.05) was based on the MOWSE score (higher than 65) and CI > 99.8%, and confirmed by the accurate 12 13 overlapping of the matched peptides with the major peaks of the mass spectrum.

14

#### 15 Routine DNA manipulations and site-directed mutagenesis of *nasA* leader 16 sequence

Genomic and plasmid DNA were routinely isolated and purified using the Wizard<sup>®</sup> Genomic DNA purification kit (Promega, USA) and the Qiagen plasmid kit (Qiagen, Germany), respectively. Custom oligonucleotide primers, listed in Table S2, were supplied by Invitrogen (Paisley, UK) and the polymerase chain reaction (PCR) was performed using the Expand High Fidelity PCR system (Roche, Switzerland) with 5% DMSO added as standard.

23 In the mutational analysis carried out in trans of the nasA leader region, a nasA promoter transcriptional fusion was constructed by using an intergenic nasS-nasA 392-24 25 bp fragment amplified by PCR with oligonucleotides FA1 A/RA1, which were located 26 at the 3'-end of nasS and at the 5'-end of nasA, respectively. This fragment was cloned 27 within the pSparkII vector and subcloned as SphI-PstI in the promoter probe vector 28 pMP220, generating pMP220-PnasA (Table S1). Different mutated versions of PnasA-29 *lacZ* were also constructed by using oligonucleotides containing substitutions of three 30 bases for two hairpins, HI or HII, identified in the *nasA* gene leader region (HI 137 bp 31 and HII 91 bp upstream 5'-end nasA gene, respectively). Oligonucleotides FA1 B/RHI 32 and FHII/RA1 were used for mutagenesis of hairpins I or II, respectively, and the PCR 33 products amplified with these oligonucleotides were cloned into the pSparkII vector. 34 The pMP220-PnasA vector was digested with PstI/AscI and used to clone the mutated 35 fragment of the hairpin I previously liberated from pSparkII with PstI/AscI restriction enzymes, obtaining the pMP220-PnasA1 construct. The pMP220 plasmid is a non-36 37 integrative vector. For mutagenesis of hairpin II, the corresponding mutated fragment 38 was liberated from pSparkII with AscI/SphI and cloned into pMP220-PnasA previously 39 digested with AscI/SphI, obtaining pMP220-PnasA2 plasmid. All constructs were 40 sequenced (UCO-SCAI, University of Córdoba) to confirm the native or the mutated 41 sequences and the final constructs were introduced into P. denitrificans PD1222 by 42 triparental mating [2].

In the mutational analysis carried out *in cis* of the ANTAR region (hairpin I), the DNA regions upstream and downstream this secondary structure were amplified by PCR with oligonucleotide pairs AIF1/AIR1a and AIF2/AIR2, respectively. These two fragments were cloned separately into the pSparkII cloning vector. The upstream fragment (524 bp) was subcloned as *EcoRI/Sal*I into pK18mobsacB, a kanamycinresistant suicide vector in *Paracoccus denitrificans*. The downstream fragment (578 bp) was subcloned as *SalI/Hind*III into the previous construction. The final construction

1 contained a 35 bp sequence corresponding to the multicloning site of the pSparkII 2 vector replacing the native hairpin I sequence (37 bp). This plasmid was used for 3 triparental mating with the receptor strain Paracoccus denitrificans and the helper strain 4 E. coli containing the pRK2013 plasmid. First transconjugant selection was carried out 5 in LB media with spectinomycin and kanamycin. After that, a sucrose selection for double-crossover events was performed. Finally, mutation was confirmed by PCR and 6 enzyme restriction analyses. The wild-type *nasA* leader region with hairpin I was 7 8 amplified with oligonucleotides FA1 A/RA1. This wild-type fragment was not digested by SalI, whereas the fragment amplified from the mutant was digested by this restriction 9 10 enzyme because it contains the SalI recognition sequence of the pSparkI multicloning 11 site.

12

## 13 Site-directed mutagenesis of the NtrC binding sequence

14 The native *nasT* promoter, as well as a mutated version affected in the putative NtrC 15 binding site located upstream of nasT gene, were used to generate transcriptional 16 fusions by PCR-driven overlap extension. Initial PCRs were carried out with 17 oligonucleotides PTF/PTNtrR and PTR/PTNtrF for the native construct, and PTF/PTNtrR Mut and PTR/PTNtrF Mut for the mutated version. Internal primers 18 19 (PTNtr) were used to generate overlapping and complementary 3'-ends on the 20 intermediate segments. In the case of the mutated *nasT* promoter, the PTNtr primers 21 were also used to introduce the mutated bases. Overlapping strands hybridize at this 22 region in a subsequent PCR and were extended to generate the full-length product (619-23 bp) amplified by flanking primers that include restriction enzyme sites for cloning as 24 *PstI/SphI* into the promoter probe vector pMP220. The final constructs for the analysis 25 of the native *nasT* promoter and the NtrC-binding site mutated version of *nasT* 26 promoter were called pMP220-PnasT and pMP220-PnasT1, respectively. All plasmids 27 were checked by sequencing (UCO-SCAI, University of Córdoba) and the final constructs were introduced into P. denitrificans PD1222 by triparental mating [2]. 28

29

## 30 RT-PCR and qPCR reactions

31 P. denitrificans wild-type and mutant strains were cultured in minimal medium with 32 succinate as carbon source and different nitrogen sources. Cells were harvested (A<sub>600</sub>  $\sim$ 0.4) and washed in TEG buffer with 25 mM Tris-HCl (pH 8.0), 1% glucose and 10 mM 33 34 EDTA. RNA isolations were performed following the Qiagen RNA extraction kit 35 (RNeasy midi kit). DNase incubation was carried out in the column with RNase-free 36 DNase set (Qiagen) and an additional post-column treatment was required with DNase I 37 (Ambion). The concentration and purity of the RNA samples were measured in a 38 ND1000 spectrophotometer (Nanodrop Technologies). Synthesis of total cDNA was 39 achieved in 20 µL final volume, containing: 500 ng RNA, 0.7 mM dNTPs, 200 U 40 SuperScript II Reverse Transcriptase (Invitrogen) and 3.75 µM random hexamers 41 (Applied Biosystems). Samples were initially heated at 65 °C for 5 min and then 42 incubated at 42 °C for 50 min, followed by incubation at 70 °C for 15 min. To carry out 43 PCR reactions, 2 µL of each cDNA were initially heated at 98 °C for 2 min, followed by 44 30 cycles of amplification: 98 °C, 30 s; 60 °C, 30 s and 69 °C, 1 min. Polymerase extension reactions were completed by an additional incubation at 69 °C for 10 min. For 45 46 real-time assays, the cDNA was purified using Favorprep Gel/PCR purification kit 47 (Favorgen) and the concentration was measured using a ND1000 spectrophotometer. 48 The iQ5 Multicolor Real-Time PCR Detection System (Bio-Rad) was used in a 25 µL 49 reaction (final volume), containing 2 µL diluted cDNA (12.5, 2.5 and 0.5 ng), 0.2 µM of

3 Samples were initially denatured by heating at 95 °C for 3 min, followed by 40 cycles of

4 amplification (95 °C, 30 s; test annealing temperature, 60 °C, 30 s; elongation and signal

5 acquisition, 72 °C, 30 s). For relative quantification of the fluorescence values, a

6 calibration curve was made using dilution series from 100-0.001 ng of *P. denitrificans* 

7 genomic DNA sample. Represented data were normalized by using the *rpoB* and the 8 *dnaN* genes as housekeeping (Table S2). Primer design and relative fold-change values

- analygenes as housekeeping (Table 32). Finner design and relative for
   calculation were carried out as previously described [38,39].
- 10

# 11 Generation of *ntrB* and *ntrY* mutant strains of *P. denitrificans*

12 Mutant strains of *P. denitrificans* were generated by significant allelic replacement with 13 specific antibiotic resistance markers. The *ntrB* mutant (*ntrB* $\Delta$ ::Sm) was generated by 14 PCR amplification of the 3'-end (682 bp) and 5'-end (533 bp) of the *ntrB* gene with the respective set of primers ntrB1/ntrB2 and ntrB3/ntrB4 (Table S2). The PCR products 15 were cloned within the pGEM-T Easy vector and then assembled within pUC18 vector 16 17 (Table S1). A unique BamHI restriction site was generated at the interface of the 3'-end 18 and the 5'-end of ntrB gene to generate a 685 bp deletion and to insert a streptomycin 19 resistance cassette from the pSRA2 vector (Table S1). A single EcoRI fragment 20 (*ntrB* $\Delta$ ::Sm) was transferred to pSUP202 to produce the final mobilizable vector.

21 For the *ntrY* mutant strain (*ntrY* $\Delta$ ::Km), fragments containing the 3'-end (510) 22 bp) and 5'-end (717 bp) of ntrY were produced by PCR using the respective primer 23 pairs ntrY1/ntrY2 and ntrY3/ntrY4 (Table S2) and cloned within the pGEM-T vector 24 (Table S1). The upstream region was screened by restriction digestion and selected by 25 using an *Eco*RI recognition site and the downstream region of *ntrY* was introduced as an AatII-BamHI fragment, yielding a unique BamHI restriction site between both regions, 26 27 generating a 1035 bp deletion. A kanamycin resistance cassette obtained from the 28 pSRA2 vector (Table S1) was inserted as a BamHI fragment between these regions of 29 ntrY. Finally, the AatII-EcoRI fragment (ntrYA::Km) was cloned into the mobilizable 30 vector pSUP202. This construct was transferred to either the wild-type strain or the ntrB 31 mutant to generate the single *ntrY* or the double NtrB/NtrY mutant, respectively. 32 Plasmids were analyzed by DNA-sequencing (UCO-SCAI, University of Córdoba) and 33 mutants were checked by PCR by using primer pairs that amplify for a product 34 containing the deletion-insertion mutation. Mutants showed bands higher than those 35 presented in the wild type because the inserted resistance cassettes were larger than the 36 deleted fragments, confirming allelic exchange by double-crossover mechanism.

37

# 1 **RESULTS**

2

## 3 Transcriptomic differences between bacteria cultured aerobically under nitrate-4 versus ammonium-dependent anabolism

5 To investigate the underlying biochemical adaptation to nitrate assimilation compared to ammonium-dependent anabolism, transcriptomic analysis was undertaken through 6 microarray analysis of RNA isolated from P. denitrificans cells grown in batch culture 7 8 with 20 mM succinate and either 10 mM nitrate or 10 mM ammonium as the sole N-9 source. Of the 5077 protein-coding genes that are present in the genome of P. 10 denitrificans PD1222, only 83 genes (1.64 % of the whole genome) displayed at least 2fold expression differences at a 95% significance level between the two growth 11 conditions tested. Among these, 74 genes were induced during nitrate-dependent 12 13 growth, while 9 genes were down-regulated (Fig. 1A).

14 The genes induced when nitrate was the only nitrogen source belonged to 30 15 gene clusters, including among others, the gltB and gltD genes encoding the small and 16 large subunits of the glutamate synthase (GOGAT), the amtB gene that codes for a high-17 affinity ammonium transporter, the ntrB and ntrC genes that code for the general 18 nitrogen regulatory Ntr system, the glnB gene that codes for the nitrogen regulatory 19 protein PII, the glnA gene encoding the glutamine synthetase (GS), the bztA-D gene 20 cluster involved in glutamate/aspartate transport, the ureA-GJ gene cluster for urea 21 metabolism, the urtA-E gene cluster for urea transport, the potDF-I gene cluster 22 involved in polyamine transport, the braC-G gene cluster involved in branched-chain 23 amino acids transport, the *dctPQM* genes encoding a TRAP dicarboxylate transporter 24 system, the respiratory nitrate reductase narGJ genes, and the nasTSABGHC gene 25 cluster for nitrate/nitrite assimilation (Fig. 1A). The genes down-regulated during 26 growth with nitrate included the gdhA gene that encodes the glutamate dehydrogenase 27 (Fig. 1A). Expression of a selection of these genes was also quantified by qPCR and 28 corroborated well with the microarray data (Table 1). The genes for which expression 29 was highly increased on nitrate included nasA, nasH, narG, narJ, nasT, nasS, urtC, 30 amtB, ureD, glnA, glnB, gltD, and ntrC. The nasT gene was more strongly induced by 31 nitrate than the nasS gene (Table 1).

32 Qualitative proteomic analysis by 2D-PAGE of soluble fractions of P. 33 denitrificans cells grown with nitrate or ammonium as the sole N-source confirmed that 34 a significative number of the key changes in the transcriptome could be also indentifed 35 in the proteome (Fig. 1B). The up-regulated proteins for which higher levels of 36 synthesis in the presence of nitrate could be confirmed included the nitrogen regulatory 37 protein GlnB (PII), the periplasmic component of the urea ABC-type transporter (UrtA), 38 the polyamine transporter component (PotD), the glutamate/aspartate transporter 39 component (BztA), and two proteins encoded by the nas gene cluster of P. denitrificans 40 (NasG and NasB). A decreased synthesis of the glutamate dehydrogenase (GdhA) was 41 observed in the nitrate grown cells compared to the ammonium-grown cells (Fig. 1B).

42

#### 43 Analysis of the expression of the *nasTS and nasABGHC* transcripts

The transcriptome differences between nitrate- and ammonium-anabolising cells suggest a general adaptation to metabolism of secondary (non-ammonium) nitrogen sources, nitrogen scavenging and the need to search for organic carbon to provide the reductant for nitrate assimilation. Alongside this is a more specific adaptation to nitrate anabolism through expression of the *nasABGHC* genes. Therefore, the control of the

1 nas gene expression was investigated. The regulatory pattern for P. denitrificans nas 2 gene expression was analyzed at the transcriptional level by RT-PCR analysis with 3 RNA from cells grown with nitrate, glutamate, ammonium, or ammonium plus nitrate 4 as N-source. Primer pairs were designed to explore transcription across the 3'- and the 5'-ends of different nas gene boundaries (Fig. 2A). The PCR products were detected 5 across the nasTS, nasAB, nasBG, nasGH, nasHC, and nasGHC boundaries (Fig. 2B). 6 However, PCR product was undetectable when the primers located at the 3'-end of the 7 8 nasS gene and the 5'-end of the nasA gene were used. These results suggest that the 9 nasTSABGHC gene cluster comprises two different transcriptional units, one 10 corresponding to the regulatory *nasTS* genes and the other constituted by the structural 11 nasABGHC genes.

12 The RT-PCR analysis also indicated that the *nasA-C* transcript is found in cells 13 grown with either nitrate or glutamate as sole N-source, but not when ammonium was 14 present (Fig. 2). However, a high glutamine concentration (5 mM) or a 2-15 oxoglutarate/glutamine 5 mM/5 mM ratio did not repress expression of the nas genes. The nasA gene expression, determined by RT-qPCR, was similar in cells grown with 16 17 glutamate or glutamate plus nitrate (Fig. 2C). The *nasTS* transcript was detected in cells 18 grown with all nitrogen sources tested, although only at very low levels in the presence 19 of ammonium (Fig. 2C).

20 A proteomic approach was applied to confirm the presence of polypeptides 21 encoded by the nas genes in cytoplasmic fractions of P. denitrificans cells grown with 22 different nitrogen sources. The Rieske-type protein NasG required for both nitrate and 23 nitrite reductase activities (Fig. 3A) and the catalytic subunit of the nitrite reductase 24 NasB (not shown) were only detected when nitrate was present in the media and 25 ammonium was absent. NADH-dependent nitrate reductase activity was determined in 26 the cytoplasmic fraction from cells grown with different nitrogen sources. In accordance 27 with the proteomic analysis, the nitrate reductase activity was detected only in presence 28 of nitrate and absence of ammonium (Fig. 3A). It is notable that although the 29 nasABGHC transcript was detected in cells grown with glutamate in absence of nitrate 30 (Fig. 2B), the NasG and NasB proteins and the nitrate reductase activity were not 31 detectable under these conditions (Fig. 3A).

32

#### 33 Analysis of *nas* gene expression in the *nasT* and *nasS* mutants

34 It has been previously described that a *nasT* mutant of *P. denitrificans* is unable to grow 35 with nitrate as the sole nitrogen source, whereas a P. denitrificans nasS mutant grows 36 with nitrate as the sole N-source with similar rate and yield to the wild-type strain [5]. 37 To analyze the regulatory effect of nitrate in the *nasS* and *nasT* mutants, a comparison 38 of nas gene expression between cells grown with glutamate versus cells grown with 39 glutamate plus nitrate was performed. RT-qPCR analysis showed that the nasT mutant 40 lacks the nasABGHC transcript (Fig. 3B). Accordingly, 2D-PAGE revealed that this 41 mutant strain is also devoid of the NasG polypeptide (Fig. 3C). It is noticeable that the 42 respiratory nitrate reductase *narG* and *narJ* genes were up-regulated in the *nasT* mutant 43 strain, whereas genes coding for glutamine synthetase (glnA) and PII regulatory protein 44 (glnB) were down-regulated (Table 2), suggesting that NasT might also serve as part of 45 a more complex global regulatory system.

46

In the *P. denitrificans nasS* mutant, the *nasABGHC* transcript was detected in all
nitrogen sources, except when ammonium was used as N-source, similarly to the wildtype strain (Fig. 3B). The NasG polypeptide and the nitrate reductase activity were

detected in the *nasS* mutant strain grown with glutamate both in presence and absence of nitrate (Fig. 3C), whereas in the wild-type strain the NasG polypeptide and the nitrate reductase activity were detected only in the presence of nitrate (Fig. 3A). In addition, RT-qPCR analysis revealed that the *nasS* mutant strain showed significantly increased

5 levels of the *narG* and *gdhA* genes in response to nitrate (Table 2).

6

## 7 Identification of putative *cis*-regulatory RNA secondary structures in the *nasA* 8 leader region

9 Between the regulatory *nasTS* genes and the structural *nasABGHC* genes lies a ~200 bp 10 non-coding region with putative hairpin structures that could lead to transcription termination (Fig. 4A). Secondary structures in the RNA leader sequence of nasA are 11 12 thought to terminate transcription [14,15], but binding of NasT could allow 13 transcription antitermination. Putative hairpin structures that could be a target for the 14 transcriptional antiterminator NasT in the *nasA* leader region were identified by using 15 the RNAfold web server http://rna.tbi.univie.ac.at/cgi-bin/RNAfold.cgi. To explore the 16 importance of this region, several lacZ transcriptional fusions were constructed and 17 expressed in P. denitrificans cells grown with either nitrate or ammonium as sole N-18 source. In cells grown under nitrate anabolizing conditions, the mutation of the HI 19 hairpin resulted in a severe deficiency in the transcriptional activity. However, 20 transcriptional activity was relatively unaffected in the HII mutation (Fig. 4B). In cells 21 grown with ammonium the transcriptional activity was 100-fold lower than in nitrate-22 assimilating cells, but there was a two-fold increase in cells carrying the HII mutation. 23 These results suggest that the HI hairpin is important for NasT binding, and that HII 24 structure may contribute to transcription attenuation. To corroborate the importance of 25 HI, a specific chromosomal mutation in cis of the putative NasT binding region was 26 introduced. This chromosomal mutant lost the parental capacity to grow with nitrate as 27 the sole nitrogen source (Fig. 4C). Controls established that the chromosomal HI mutant 28 strain showed a similar growth to the parental strain with ammonium as nitrogen source. 29

#### 30 The contribution of *ntrBCYX* to the regulation of nitrate assimilation

31 In addition to *nasTS* genes, the *ntrBC* genes were also up-regulated during nitrate-32 dependent anabolism compared to cells grown with ammonium (Fig. 1A, Table 1). This 33 suggests an integrated regulation of nitrate metabolism between these two systems. The 34 ntrBC genes are clustered together with the ntrYX genes. The ntrB gene encodes a 35 signal transduction protein (386 residues) with histidine kinase activity involved in nitrogen metabolism, and the *ntrC* gene codes for a Fis-like  $\sigma^{54}$ -dependent regulator 36 37 receiver protein (486 residues). Similarly, the *ntrY* gene codes for a multi-sensor signal 38 transduction regulator (709 residues) with histidine kinase activity and the *ntrX* gene 39 encodes a  $\sigma^{54}$ -dependent regulatory protein (463 residues). A non-coding region of 217 bp lies between *ntrC* and *ntrY* genes, whereas *ntrY* and *ntrX* genes overlap by 3 bp. 40 41 NtrB and NtrC homologues have been found in many different microorganisms (Fig. 42 S1). Although the NtrYX system is not widely distributed among bacteria, the P. 43 denitrificans NtrY protein showed the highest identity (41%) with Azorhizobium 44 caulinodans NtrY, whereas the P. denitrificans NtrX protein showed the highest 45 identity (56%) with its homologues in Azorhizobium caulinodans and Azospirillum 46 brasilense (Fig. S2).

47 Isolation and characterization of single *ntrB* and *ntrY* mutants and double 48 *ntrB/ntrY* mutant of *P. denitrificans* were carried out. All mutant strains displayed 49 similar growth rates and yields to that shown by the wild-type strain when ammonium 1 was used as nitrogen source. However, the *ntrY* mutant was only slightly affected on its 2 growth on nitrate, whereas the *ntrB* mutant showed a longer lag-phase than the wild-3 type, and the double *ntrB/ntrY* mutant was unable to grow with nitrate (Fig. 5A).

4 Expression of the nasT, nasS, ntrB, ntrY regulatory genes and the structural nasA 5 gene was analyzed by RT-qPCR in the *ntrB* and *ntrY* mutants and the wild-type strain 6 of P. denitrificans. In the ntrB mutant nasTS and nasA were down-regulated (Fig. 5B), 7 whereas in the *ntrY* mutant all these genes were expressed similarly to the wild-type 8 strain (not shown). The assimilatory NADH-dependent nitrate reductase activity was 9 determined in cytoplasmic fractions from wild-type, *ntrB* and *ntrY* mutants grown with 10 nitrate, when all strains reached similar biomass (protein concentration about 150 mg mL<sup>-1</sup>). The *ntrY* mutant showed similar levels of activity to that presented by the wild-11 type, about 4 nmol NO<sub>2</sub><sup>-</sup> formed min<sup>-1</sup> mg<sup>-1</sup>, whereas the *ntrB* mutant only presented an activity of about 1 nmol NO<sub>2</sub><sup>-</sup> formed min<sup>-1</sup> mg<sup>-1</sup>. 12 13

Transcriptional fusions to the β-galactosidase-encoding gene *lacZ* were carried out in the *nasT* leader sequence and expressed in the wild-type, *ntrB* and *ntrY* mutant strains of *P. denitrificans*. The β-galactosidase activities in the *ntrY* mutant were very similar to those found in the wild-type strain (Table 3). However, β-galactosidase activities in the *P. denitrificans ntrB* mutant were much lower than those described for the wild-type strain (Table 3).

20

21 These data strongly suggest a role for NtrBC in the regulation of nitrate 22 assimilation at the level of expression of *nasT* and so the possibility of a *cis*-acting 23 binding site for NtrC was explored. The RegPrecise program for collection of manually 24 curated inferences of regions in prokaryotic genomes identified a very well conserved 25 putative NtrC binding site (5'- (T/C)GC(C/A)NNNTTNNT(G/A)GCA-3') in the 26 leader regions of a number of the different genes in P. denitrificans, including the 27 glnBA, amtB, and nasTS genes. These genes were upregulated during nitrate 28 assimilation, as revealed by the transcriptomic analyses (Fig. 1A, Table 1). Several base 29 pairs were mutated in the putative NtrC binding site located in the nasT leader 30 sequence, resulting in the loss of the up-regulation observed under nitrate- or glutamate-31 dependent growth conditions (Table 3). These results highlight the importance of this 32 sequence motif in the regulation of nitrate assimilation.

33 34

#### 1 **DISCUSSION**

2

3 The soil denitrifier P. denitrificans can use nitrate as the sole N-source by an 4 assimilatory nitrate reduction system. The process of nitrate assimilation in this bacterium has been comprehensively investigated by comparative transcriptomic and 5 proteomic analyses in cells grown under nitrate- or ammonium-assimilating conditions 6 (Figs. 1, 3 and S3). Comparing nitrate- to ammonium-grown cells, a key metabolic 7 8 change was the decrease in the expression of glutamate dehydrogenase, the enzyme related with the low affinity ammonium assimilation pathway. This was accompanied 9 10 by induction of components for organic nitrogen scavenging, urea uptake and 11 utilisation, high affinity ammonium uptake and assimilation, regulatory and structural 12 components of inorganic nitrate and nitrite assimilation, and scavenging of 13 dicarboxylate organic acids to provide the additional reducing equivalents required for 14 nitrate assimilation (eight electrons per nitrate ion converted into ammonia).

15 In *P. denitrificans*, the regulatory *nasTS* genes are located at the 5'-end of the *P*. 16 denitrificans nas gene cluster for nitrate assimilation (Fig. 2A). RT-PCR and qPCR 17 analyses of the P. denitrificans nasTS and nasABGHC genes demonstrate that these 18 genes are transcribed as separated transcriptional units. The existence of two 19 independent transcriptional units in the P. denitrificans nas region is also supported by the presence of a non-coding region between the regulatory *nasTS* genes and the 20 21 structural *nasABGHC* genes, with two putative hairpins (HI and HII) that could lead to 22 secondary structures in this mRNA region (Fig. 4). These two transcriptional units show 23 different regulation, since expression of the regulatory nasT and nasS genes was 24 observed in all N-sources tested, including ammonium, although in general at very low 25 levels, whereas *nasABGHC* transcript was detected either with nitrate or glutamate, but 26 it was undetectable with ammonium independently of the presence or absence of nitrate 27 (Fig. 2). However, although the *nasABGHC* transcript was present in wild-type cells 28 grown with glutamate as the sole N-source, the Rieske-type NasG protein and the large 29 subunit of the assimilatory nitrite reductase NasB, required for both nitrate and nitrite 30 reductase activities, were not detected by 2D-PAGE and these proteins were only 31 observed in the presence of nitrate. Accordingly, the assimilatory NADH-dependent 32 nitrate reductase activity was not detected when glutamate was used as the sole nitrogen 33 source (Fig. 3A). These results suggest that nitrate is not an obligate inducer of the 34 nasABGHC gene expression, and that it may also play a positive role at the translational 35 level.

36 Mutational analysis of the nasT gene of P. denitrificans reveals that NasT could 37 act as transcription antiterminator because the *nasT* mutant strain is unable to grow with 38 nitrate as the sole N-source and lacks both nasABGHC transcript and NADH-nitrate 39 reductase activity (Fig. 3BC). Also, the strain harbouring the chromosomal mutation in 40 a putative *cis*-acting NasT binding site (hairpin I) was unable to grow with nitrate (Fig. 41 4C). Therefore, in this work we have identified the RNA region where NasT is likely to 42 bind as a transcriptional antiterminator. On the contrary, the *nasS* mutant strain of *P*. 43 denitrificans is capable to use nitrate as the sole N-source, similarly to the wild-type 44 strain, but shows a deregulated nitrate reductase activity that can be detected 45 independently of the presence or absence of nitrate (Fig. 3BC). In addition, assimilatory 46 nitrate reductase activity in the P. denitrificans nasS mutant is abolished in the presence 47 of ammonium, as previously described in the wild-type strain [5]. Expression of the 48 nasT gene is higher than nasS gene expression (Tables 1 and 2), probably because strong secondary structures present at the 5'-end of the *nasS* gene might provoke RNA polymerase to be occasionally released without completing the whole transcript.

3 P. denitrificans NasS and NasT proteins expressed in E. coli have been purified as a tetrameric complex in the absence of nitrate or nitrite, but in addition to the NasT-4 5 NasS complex, free NasT protein can be also detected [5]. Therefore, in the absence of 6 nitrate, NasS and NasT form a protein-protein complex, probably with two units of each 7 protein, which limits binding of NasT to the nas mRNA to act as transcription antiterminator. However, in the presence of nitrate, NasS clamps the oxyanion, 8 9 changing conformation and dissociates from NasT, thus increasing the size of the free 10 NasT pool available to serve for *nasABGHC* transcription antitermination [5]. However, 11 the elevated *nasT* gene expression over the *nasS* gene could lead to an excess of NasT 12 protein over the NasS protein. This unbalance implies that there is always enough free 13 NasT to allow transcription of nasABGHC genes in the absence of both nitrate and 14 ammonium (i.e., glutamate as N-source), suggesting additional global regulatory roles. 15 In the *P. denitrificans nasT* mutant the structural *nasABGHC* transcript was absent in all 16 N-sources tested (Fig. 3B). This is consistent with a role for NasT as transcription antiterminator and with the phenotype of the P. denitrificans nasT mutant, which is 17 18 unable to grow with nitrate or nitrite as the sole N-source. In addition, up-regulation of 19 narG and narJ genes and down-regulation of glnAB genes in the nasT mutant also 20 indicated that NasT may be involved in other regulatory processes. Recently, it has been 21 suggested in the soybean endosymbiont Bradyrhizobium japonicum that nasST genes 22 regulate respiratory nitrous oxide and periplasmic nitrate reductases [41]. Therefore, 23 additional targets for NasT may exist in P. denitrificans, which could function as 24 negative regulators of nar gene expression.

25 In the *nasS* mutant the *nasABGHC* gene expression is no longer under nitrate 26 control because in the absence of the NasS protein, all the NasT pool is free to serve as 27 transcription antiterminator under all growth conditions (Fig. 3BC). Thus, in contrast to 28 the wild-type strain, the nasS mutant of P. denitrificans displayed in 2D-PAGE gels the 29 assimilatory nitrite reductase NasB and the Rieske-type NasG polypeptides when 30 glutamate was used as nitrogen source, independently of the presence or absence of 31 nitrate (Fig. 3C). These results corroborate that nitrate exerts a regulatory control at 32 post-transcriptional (translational) level in addition to the role of NasTS in the 33 transcription antitermination.

34 The phenotypes of the *ntrB*, *ntrY* and *ntrB/ntrY* mutants of *P*. *denitrificans* 35 suggest that the NtrBC system is mainly required for nitrate assimilation, while the 36 NtrYX system only has a slight contribution (Fig. 5A). The regulatory nasTS genes and the structural nasA gene are down-regulated in the ntrB mutant of P. denitrificans (Fig. 37 38 5B), suggesting that these genes are under the control of the NtrBC system. This idea 39 has been corroborated by using the lacZ gene transcriptionally fused to native and 40 mutated *nasT* leader sequence (Table 3). These results indicate that, in the absence of 41 ammonium, phosphorylated NtrC could bind to the nasT promoter region, where a 42 consensus NtrC-binding site has been identified. The absence of the nasABGHC 43 transcript in cells grown with ammonium also suggests that NtrC or NrtX may bind to an, as yet unidentified, site in the nasA leader region. According with the RegPrecise 44 45 program, very well conserved putative NtrC binding sites are found in the promoter 46 regions of the *nasT* and *ntrBC* genes, suggesting that these two gene clusters are under 47 the regulation of the general nitrogen control.

1 The increased synthesis of P. denitrificans nasABGHC genes reflects a system-2 specific response to ammonium-limited growth in which nitrate is initially the sole 3 available N-source [2]. This bacterium also uses nitrate as terminal electron acceptor by 4 a membrane-bound nitrate reductase [1]. This respiratory nitrate reductase (Nar) can substitute for NasC (assimilatory nitrate reductase) during anaerobic growth. The 5 present study shows that transcription of *nasTS* occurs at a low level in the presence of 6 ammonium, whereas transcription of *nasABGHC* genes is fully repressed in cells grown 7 8 with ammonium (Fig. 2). In K. oxytoca, nasR expression is sensitive to ammonium 9 since it is induced by the NtrBC system when ammonium is absent [14]. In Azotobacter 10 vinelandii expression of nasST genes is also sensitive to ammonium, although is not 11 under the control of the Ntr system [17]. Nevertheless, the capacity for nitrate assimilation is lost in several ntr mutants [19]. Recently, it has been demonstrated that 12 13 NtrBC and NasST co-regulate the *nasAB* genes required for nitrate/nitrite uptake and reduction in A. vinelandii [20]. Bioinformatic analysis has revealed that the two-14 15 component NasTS system is a regulatory mechanism for nitrate assimilation that is widely spread among bacteria. Particularly, NasT is mainly present in  $\alpha$ -proteobacteria 16 like Rhizobiales and Rhodobacterales, and  $\beta$ -proteobacteria like Burkholderiales [5]. 17 NasTS is involved in the regulation of the assimilatory nitrate/nitrite reductase genes in 18 19 response to nitrate or nitrite in Azotobacter vinelandii, Pseudomonas putida and 20 Rhodobacter capsulatus [17,18,42]. However, in Azotobacter vinelandii these genes are 21 arranged in the *nasST* orientation, instead of the *nasTS* organization found in *P*. 22 denitrificans, and they do not cluster together with the nas genes encoding the 23 nitrate/nitrite uptake and reduction system [43]. The NasS protein is homologous to 24 proteins which belong to the ATP-dependent nitrate transport systems found in 25 cyanobacteria, although NasS lacks the signal peptide for periplasmic translocation. This is consistent with the role of NasS as a nitrate/nitrite sensor rather than acting as 26 27 the periplasmic component of the transport system for these oxyanions. The NasT 28 protein is a regulator with the ANTAR binding domain characteristic of transcription 29 antiterminators like the Klebsiella oxytoca NasR protein [4,14,16]. However, the nasTS 30 genes are absent in the *nas* cluster of *Klebsiella*, in which they are functionally replaced 31 by *nasR*, a gene mainly found in  $\gamma$ -proteobacteria. It has been proposed that NasR is 32 both a nitrate/nitrite sensor and a transcription antiterminator. A hairpin structure in 33 mRNA upstream from the K. oxytoca nas structural genes causes early termination of 34 transcription [14]. It has been suggested that in the presence of nitrate or nitrite NasR binds to the mRNA transcript preventing hairpin formation. Additional novelty on the 35 regulatory mechanisms for nitrate assimilation was found in P. denitrificans because 36 37 repression of structural nas genes by ammonium could be directly exerted by 38 ammonium itself rather than by glutamine or the glutamine/2-oxoglutarate ratio. Under 39 low nitrogen conditions (absence of ammonium), the NtrB and NtrY proteins become 40 active to phosphorylate their respective partners, NtrC or NtrX, which in turn, become active for binding to promoter regions of the regulated genes. In R. capsulatus the 41 42 histidine kinases NtrB and NtrY can substitute for each other as phosphodonors towards 43 the response regulator NtrC [29]. Also, a cross-talk between the NtrB/C and NrtY/X 44 sensor/regulator pairs has been suggested in A. brasilense during nitrate-dependent 45 growth [26]. However, NtrY protein is much larger than NtrB suggesting that they may 46 play different functions. The NtrBC system is widespread among microorganism and 47 displays known functions, like the control of genes for glutamine synthetase, amino 48 acids transport, nitrate and nitrite assimilation, nitrogen fixation and, also for the 49 expression of other nitrogen regulation genes. Although the role of the NtrBC system seems to be more related to nitrogen metabolism, in A. brasilense there is also a link 50

1 between this two-component regulatory system and polyhydroxybutyrate production 2 [44]. To summarize, Fig. 6 represents the integration of mutational, transcriptomic and 3 proteomic data to establish a novel regulation model of the nitrate/nitrite assimilation 4 process in P. denitrificans, in which nitrate exerts a transcriptional and translational 5 control that has no precedents in the literature as far as we know. In this model three conditions have been considered: i) presence of ammonium, in which the nasABGHC 6 genes are repressed and there is a very low expression of the *nasTS* genes; ii) absence of 7 8 both ammonium and nitrate (i.e. glutamate as N-source), in which nasABGHC and 9 *nasTS* genes are expressed, but Nas polypeptides and nitrate reductase reductase activity 10 are not detected; and iii) absence of ammonium and presence of nitrate, in which 11 nasABGHC and nasTS genes are expressed and Nas proteins are synthesized, leading to 12 an active nitrate assimilation system. Our experimental data supports the existence of a 13 regulatory cascade with three levels of control. At level 1, the phosphorylated NtrC protein (and probably NtrX in a lesser extent) interacts with the nas promoter to activate 14 15 nas gene expression in absence of ammonium, as described in other bacteria (general 16 nitrogen control). Accordingly, a conserved NtrC-binding sequence has been found in 17 the promoter region of the *nasT* gene. At level 2, the NasTS system controls expression 18 of the *nasABGHC* genes by transcription antitermination. In the absence of nitrate, an 19 inactive NasTS complex is formed but there is enough free NasT protein to allow 20 transcription antitermination for the synthesis of a complete *nasABGHC* transcript, 21 which is detectable in cells grown with glutamate. This NasT/NasS unbalance is 22 probably a consequence of the high level of expression of the *nasT* gene compared to 23 the nasS gene. At level 3, translation of the Nas proteins only occurs in the presence of 24 nitrate. This regulation at the translational level may occur when NasS binds to nitrate 25 and the NasTS complex dissociates, leading to an increase of free NasT protein that, in 26 turn, may lead to up-regulation of systems involved in post-translational regulation.

This is a novel regulatory mechanism, but bioinformatics analyses may suggest that NasTS regulation is emerging in a wide number of bacteria, as previously reviewed [8]. As the *P. denitrificans* Nas system is very well characterized [2,5], this work may now lead to comparative studies in other bacteria, in which the NasTS system regulated nitrate assimilation and even other processes like nitrous oxide reduction, as recently described in *Bradyrhizobium japonicum* [41].

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# **ABBREVIATIONS**

ANTAR, Ami and NasR transcription antitermination regulator; Cy5-dCTP and Cy3dCTP, deoxycytidine triphosphate containing either Cy5 or Cy3 dyes; dNTPs, deoxyribose-nucleoside triphosphates; EDTA, ethylenediaminetetraacetic acid; MALDI-TOF/TOF, matrix-assisted laser desorption/ionization (time of flight); NADH, nicotinamide adenine dinucleotide; NIT, N-terminal nitrate/nitrite sensor domain; Ntr, nitrogen regulator; RT-PCR, reverse transcription polymerase chain reaction; RTqPCR, quantitative reverse transcription polymerase chain reaction; 2D-PAGE, twodimensional polyacrylamide gel electrophoresis.

Gene	NO <sub>3</sub> <sup>-</sup> /NH <sub>4</sub> <sup>+</sup>
Pden_0490 gltD	$2.8\pm0.5$
Pden_0794 bztB	$1.9\pm0.4$
Pden_1212 ureD	$5.1 \pm 1.8$
Pden_2032 amtB	$5.3\pm1.7$
Pden_2151 potD	$2.6\pm0.2$
Pden_4019 urtC	$6.2\pm0.4$
Pden_4129 ntrC	$4.6\pm0.5$
Pden_4130 ntrB	$1.6\pm0.2$
Pden_4234 narJ	$24.3\pm11.1$
Pden_4236 narG	$100.5\pm23.3$
Pden_4450 nasH	$185.7\pm9.8$
Pden_4453 nasA	$385.0\pm11.9$
Pden_4454 nasS	$7.7\pm1.5$
Pden_4455 nasT	$23.3\pm 6.3$
Pden_4461 glnB	$2.6\pm0.6$
Pden_4462 glnA	$3.4\pm 0.7$

Table 1. Analysis by RT-qPCR of nitrate-induced genes detected in the microarrays of *P. denitrificans*.

Data correspond to the mean  $\pm$  standard deviation of the nitrate/ammonium gene expression ratio.

Gene		Wild-type Glu+NO <sub>3</sub> -/Glu	<i>nasT</i> mutant Glu+NO <sub>3</sub> <sup>-</sup> /Glu	<i>nasS</i> mutant Glu+NO <sub>3</sub> -/Glu
Pden_0490	gltD	$0.6 \pm 0.1$	$0.7\pm0.1$	$2.6\pm0.2$
Pden_0794	bztB	$2.2\pm1.6$	$0.7\pm0.4$	-
Pden_1212	ureD	$1.1\pm0.4$	$1.0\pm0.1$	$1.8\pm0.1$
Pden_2032	amtB	$0.4\pm0.0$	$0.6\pm0.1$	$0.3\pm0.1$
Pden_2151	potD	$0.5\pm0.1$	$0.6\pm0.1$	-
Pden_4019	urtC	$0.5 \pm 0.1$	$0.5\pm0.1$	-
Pden_4129	ntrC	$0.4 \pm 0.1$	$0.5\pm0.1$	$0.8\pm0.2$
Pden_4130	ntrB	$0.7\pm0.1$	$0.6\pm0.2$	$0.6 \pm 0.1$
Pden_4234	narJ	$7.7 \pm 1.2$	$57.0 \pm 14.4$	-
Pden_4236	narG	$2.5\pm0.2$	$67.5\pm12.2$	$173.8\pm\!\!12.3$
Pden_4450	nasH	$0.6 \pm 0.2$	$0.9\pm0.1$	$0.2\pm0.1$
Pden_4453	nasA	$0.9\pm0.3$	$0.3\pm0.1$	$0.2\pm0.1$
Pden_4454	nasS	$1.0\pm0.3$	$1.4\pm0.1$	$0.0\pm0.0$
Pden_4455	nasT	$2.0\pm0.4$	$0.0\pm\!0.0$	$1.8\pm0.1$
Pden_4461	glnB	$2.0\pm0.2$	$0.4 \pm 0.1$	$1.0 \pm 0.1$
Pden_4462	glnA	$1.7\pm0.1$	$0.4 \pm 0.1$	$1.2 \pm 0.1$
Pden_3872	gdhA	$2.0\pm0.3$	1.1 ±0.2	$11.4\pm1.0$

Table 2. Analysis by RT-qPCR of nitrate-induced genes in the wild-type, *nasT* and *nasS* mutant strains of *P. denitrificans*.

Data correspond to the mean  $\pm$  standard deviation of the glutamate plus nitrate/glutamate gene expression ratio.

	N-source				
Strain/construct	$\mathrm{NH_4}^+$	NO <sub>3</sub> -	Glu	$Glu + NO_3^-$	
WT/PnasT (native)	$71 \pm 5$	$14500\pm143$	$11000\pm120$	$15000 \pm 152$	
WT/PnasT (mutated)*	$50\pm3$	$220\pm14$	$53 \pm 5$	$600\pm25$	
<i>ntrY</i> / PnasT (native)	n.d.	n.d.	$11500\pm115$	$15200\pm135$	
<i>ntrB</i> /PnasT (native)	n.d.	n.d.	$10 \pm 1$	$120\pm4$	

Table 3. Analysis of the *nasT* promoter by *lacZ* ( $\beta$ -galactosidase activity) gene fusion in *P. denitrificans* wild-type and *ntrB* and *ntrY* mutant strains.

\* The base substitutions are specified in Table S2; n.d.: not determined.

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#### **FIGURE LEGENDS**

Figure 1. Transcriptomic and proteomic analysis of aerobically cells grown with nitrate or ammonium as nitrogen source. (A) Heat map from DNA microarrays analyses of *P. denitrificans* PD1222 grown aerobically with either ammonium or nitrate as the sole nitrogen source. Colours indicate  $\log_2$  of normalised expression values in either condition. Genes were selected based on at least 2-fold change in expression with *p*-values  $\leq 0.05$ . A total of 74 genes were up-regulated more than 2-fold under nitrate assimilation conditions, whereas nine genes were down-regulated more than 2-fold under these conditions. (B) 2D-PAGE analysis of *P. denitrificans* PD1222 soluble fractions from cells grown aerobically with ammonium or nitrate (10 mM each) as the sole nitrogen source. Spots present or absent in gels are highlighted by black or white triangles, respectively. The presented gels are representative of three independent replicants.

Figure 2. Analysis by RT-PCR and qPCR of the *P. denitrificans* nitrate assimilation nas genes. (A) The nas gene cluster of *P. denitrificans* and the location of the oligonucleotides used in this study are shown. (B) RT-PCR reactions with RNA from the wild-type strain of *P. denitrificans* grown in the presence of different N-sources. To isolate RNA, cells were cultured in minimal medium with different N-sources as described in Materials and Methods section and harvested at an  $A_{600}$  of about 0.4. The *rpoB* gene was used as housekeeping. -/+, without/with reverse transcriptase (C) RT-qPCR analysis of the *nasT*, *nasS* and *nasA* genes from cells grown with different nitrogen sources. Error bars represent standard deviation calculated from the results of three independent experiments.

Figure 3. Characterization of the wild-type strain and the *nasT* and *nasS* mutants of P. denitrificans with different nitrogen sources. (A) NADH-dependent nitrate reductase activity and NasG polypeptide detection by 2D-PAGE analysis in the wildtype strain of *P. denitrificans*. The 2D-PAGE analyses were carried out in cytoplasmic fraction from cells grown with different nitrogen sources obtained by subcellular fractionations as indicated in Materials and Methods. The different nitrogen sources were nitrate, ammonium, ammonium plus nitrate, glutamate or glutamate plus nitrate (10 mM each). Isoelectric focusing was performed from IPG strips range 4-7 and second dimension was carried out onto 12% polyacrylamide gels. The nitrate reductase activity (NR) was assayed in cytoplasmic fractions and was undetectable with glutamate or ammonium as N-source, and expressed as nmol NO<sub>2</sub><sup>-</sup> formed min<sup>-1</sup> mg<sup>-1</sup> (n.d, not detected). (B) RT-PCR analysis of structural nas genes in the wild-type and nasT and nasS mutant strains of P. denitrificans. To isolate RNA, wild-type and nasT and nasS strains were cultured in minimal medium with different N-sources as described in Materials and Methods section and harvested at an  $A_{600}$  of about 0.4. The *rpoB* gene was used as housekeeping. -/+, without/with reverse transcriptase. (C) NasG polypeptide detection by 2D-PAGE analysis and NADH-nitrate reductase activity in cytoplasmic fractions from *nasT* and *nasS* mutant strains of *P. denitrificans*. The 2D-PAGE analysis and the NR activity assays were performed as indicated in (A) for the wild-type strain.

Figure 4. Analysis of the *nasA* leader region of *P. denitrificans*. (A) Intergenic *nasSnasA* region from *P. denitrificans*. (B) Expression of the transcriptional  $P_{nasA}$ -lacZ fusions and effect of mutations within the leader sequence on *nasA*-lacZ expression in cells grown with ammonium or nitrate (10 mM each). The  $\beta$ -galactosidase activity was measured when the  $A_{600}$  was about 1.0 and it was represented as Miller units (M.U.). Control represents the wild type (native) construction from -321 to +71 nucleotides of the *nasS-nasA* intergenic region. HI and HII represent the  $P_{nasA}$ -*lacZ* constructions with 3 bp mutations in each hairpin. (C) Growth curves with nitrate as sole nitrogen source of the *P. denitrificans* wild-type strain and the mutants in the *nasT* gene and the *cis*-acting NasT binding region (hairpin I, HI).

Figure 5. Characterization of the *ntrB*, *ntrY* and double *ntrB/ntrY* mutant strains of *P. denitrificans* in media with nitrate. (A) Growth-curves of *P. denitrificans* wild-type strain ( $\bullet$ ), and *ntrB* ( $\blacksquare$ ), *ntrY* ( $\blacktriangle$ ) and double *ntrB/ntrY* ( $\blacklozenge$ ) mutants with 10 mM nitrate as nitrogen source. (B) RT-qPCR analysis of the regulatory genes *nasT*, *nasS*, *ntrB* and *ntrY* and the structural *nasA* gene in the wild-type strains and the *ntrB* mutant strains of *P. denitrificans* grown with 10 mM nitrate as N-source. The oligonucleotides used in the qPCR analysis for the *ntrB* and *ntrY* genes hybridized upstream the mutation site. Represented data were normalized by using the *rpoB* and *dnaN* genes as housekeeping (Table S2). Error bars represent standard deviation calculated from the results of three independent experiments.

**Figure 6. Proposed model of the nitrate assimilation regulation in** *P. denitrificans.* The model describes the regulatory cascade controlling expression of nitrate assimilation *nas* genes by the *ntrBCYX* gene cluster in response to ammonium availability and by the *nasTS* genes for nitrate control. Levels 1 and 2 include regulatory mechanisms that control transcription of the *nas* genes, whereas level 3 involves a control on the translation of the Nas proteins. A circle with a plus symbol means activation of the transcription and a circle with the capital letter A indicates transcription antitermination.

Α

Gene ID	Name	NH₄⁺	NO <sub>3</sub>	Function / Annotation	Gene ID	Name	NH4	NO <sub>3</sub>	Function / Annotation
281	ehuB			ABC transporter ectoine binding protein	3705	argT			Amino acid ABC transporter substrate-binding protein
286	eutC			Ectoine utilisation protein	3928	braE			Branched-chain amino acid ABC transporter permease
288	eutE			Succinylglutamate desuccinylase/aspartoacylase	3929	braD			Branched-chain amino acid ABC transporter permease
488	gltB			Glutamate synthase Large Subunit (GOGAT)	3930	braG			ABC transporter ATP-binding protein
489				conserved hypothetical protein	3931	braF			ABC transporter ATP-binding protein
490	gltD			Glutamate synthase Small Subunit (GOGAT)	3932	braC			Branched-chain amino acid binding protein
552				Putative inner membrane protein	4017	urtA			Urea-binding protein
553				hypothetical protein	4018	untB			Urea ABC transporter permease
556				hypothetical protein	4019	urtC			Urea ABC transporter permease
567	sdhC			succinate dehydrogenase subunit C	4020	urtD			Urea ABC transporter ATP-binding protein
607	dpp8			Peptide/nickel/opine ABC-transporter permease	4021	untE			Urea ABC transporter ATP-binding protein
608	dppA			Peptide/nickel/opine ABC-transporter binding protein	4110	citE			Citrate lyase
794	btzB			Glu/Asp ABC-transporter Permease	4129	ntrC	_		Two-component o54 specific transcriptional regulator
795	btzC			Glu/Asp ABC-transporter Permease	4130	ntrB			Nitrogen specific signal transduction histidine kinase
796	btzD			Glu/Asp ABC-transporter ATP-binding protein	4131				hypothetical protein
823				protein of unknown function DUF404	4234	narJ			Respiratory nitrate reductase chaperone (NarJ)
824				protein of unknown function DUF403	4236	narG			Respiratory nitrate reductase alpha subunit (NarG)
825				transolutaminase. N-terminal domain protein	4449	nasC			Assimilatory nitrate reductase (NasC)
945	proX			Glycine/betaine ABC-transporter binding protein	4450	nasH			Nitrite transporter (NasH)
1205	ureG			Urease cobalamin synthesis protein	4451	nasG			Assimilatory nitrite reductase small subunit (NasG)
1206	ureE			Urease accessory protein	4452	nasB			Assimilatory nitrite reductase large subunit (NasB)
1207	ureF			Urease accessory protein	4453	nasA			Nitrate/nitrite transporter (NasA)
1208	ureC			Urease o-subunit	4454	nasS		_	Putative nitrate binding protein
1209	ureB			Urease B-subunit	4455	nasT			Response regulator reciever and ANTAR domain protein
1210	und			Urease accessory protein	4461	alnB	-		nitrogen regulatory protein P-II
1211	ureA			Urease v-subunit	4462	olnA			Glutamine synthetase (GS)
1212	umD			Linease accessory protein	4619	datP		_	TRAP dicarbowlate transporter, binding protein
1248	dodA	_	_	Pertide/nickel/opine ABC-transporter binding protein	4620	detO		_	TRAP dicarboxylate transporter small membrane protein
1508				Lytic transolycosylase, catalytic	4621	ddM			TRAP dicarboxylate transporter large membrane protein
2032	amtR	_		Ammonium transporter AmtR	4864	hisl			Amino acid ABC transporter biording protein
2149	poti		_	Polyamine ABC transporter permease	4888	omX			Givcine/betaine ABC-transporter binding protein
2150	notH			Polyamine ABC transporter permease	5095	liv <b>K</b>	_		Branched chain amino acid ABC transporter hinding protein
2151	potD			Polyamine periolasmic binding protein	0000		_	_	contract count of the count of conspected entering present
2152	potE			Polyamine periplasmic binding protein	317				hypothetical protein
2153	potG			ABC transporter ATP-binding protein	623			_	protein of unknown function DUF1127
2472	unnR		_	Carbohydrate ABC transporter binding protein	650	matF	_	-	MATE efflux family protein
2543				hypothetical protein	1552			_	conserved hypothetical protein
2610	hhuð			Hemin TonB-dependent transporter	3196	MAR			conjunal transfer protein TrhB
2961	ind	-	_	isocitrate dehydrogenase (NADP)	3872	orthA			Gistamate debudronenase (GDH)
3349	hne			nucleoid protein H-NS	4121	det0			TRAP dicarbowlate transporter small membrane rootein
3461	notR			Polyamine ARC transporter permease	4718				hypothetical notain
3462	notC			Polyamine ABC-transporter permease	4725	_			conserved hyrothetical notein
0102	puto			2 2 4 0 4	3 3		6	-	onider real information protein
				-3 -2 -1 0 1	2 3	4 3	Ů	<i>.</i>	

В



Figure 1









В



Figure 2





Figure 3











Figure 4



Α

Figure 5



Strain or plasmid	Genotype and description	Source or reference
Paracoccus denitrificans PD1222		
Wild-type	Sp <sup>R</sup> , Rif <sup>R</sup> , enhanced conjugation frequencies	[1]
nasT	<i>nasT</i> ∆::Sm <sup>R</sup>	[2]
nasS	nasSΔ::Km <sup>R</sup>	[2]
ntrB	$ntrB\Delta::Sm^{R}$	This work
ntrY	<i>ntrY</i> Δ::Km <sup>R</sup>	This work
ntrB/ntrY	$ntrB\Delta::Sm^{R}/ntrY\Delta::Km^{R}$	This work
HI (hairpin I)	Unmarked mutant	This work
Escherichia coli		
DH5a	deoR endA1 gyrA96 hsdR17( $r_k^- m_k^+$ ) recA1 relA1 supE44 thi-1 $\Delta$ (lacZYA-argFV169) Ø80 $\delta$ lacZ $\Delta$ M15 F <sup>-</sup> $\lambda^-$	[3]
pRK2013	Km <sup>R</sup> , Tra <sup>+</sup> (pRK2013 encoded) helper strain	[4]
Plasmids		
pUC18	Amp <sup>R</sup> , cloning vector	[5]
pGEM-Teasy	Amp <sup>R</sup> , cloning vector	Promega (USA)
pGEM-T	Amp <sup>R</sup> , cloning vector	Promega (USA)
pSparkII	Amp <sup>R</sup> , cloning vector	Canvax (Spain)
pSUP202	Amp <sup>R</sup> , Tet <sup>R</sup> , Cm <sup>R</sup> , Mob <sup>-</sup> , mobilizable suicide vector	[6]
pSUP202*	pSUP202::Km-∆Tet, mobilizable suicide vector	[7]
pSUP2021	pSUP202::Tn5, source of Km <sup>R</sup> cassette	[6]
pK18mobsacB	Gene knockout vector, Km <sup>R</sup>	[8]
pSRA2	Amp <sup>R</sup> , Sp <sup>R</sup> , Sm <sup>R</sup> , source of Sm <sup>R</sup> cassette	[9]
pMP220	Promoter probe vector, $lacZ$ gene, IncP1 Tc <sup>R</sup>	[10]
pMP220-PnasA	pMP220-based transcriptional fusion spanning the intergenic region between $nasS$ and $nasA$ (Tet <sup>R</sup> )	This work
pMP220-PnasA1	pMP220-based transcriptional fusion spanning the intergenic region between <i>nasS</i> and <i>nasA</i> containing the hairpin I mutated (Tet <sup>R</sup> )	This work
pMP220-PnasA2	pMP220-based transcriptional fusion spanning the intergenic region between $nasS$ and $nasA$ containing the hairpin II mutated (Tet <sup>R</sup> )	This work
pMP220-PnasT	pMP220-based transcriptional fusion spanning the $nasT$ leader region (Tet <sup>R</sup> )	This work
pMP220-PnasT1	pMP220-based transcriptional fusion spanning the <i>nasT</i> leader region and containing the	This work

# Table S1. Bacterial strains and plasmids used in this study.

	NtrC-binding site mutated (Tet <sup>R</sup> )	
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Table S2.	Oligonuc	leotide	primers	used	in	this	study.
			p · · · · · · · ·				search .

Primer	Sequence $(5' \rightarrow 3')^a$	Used for
ntrB1-HB ( <i>Hin</i> dIII- <i>Bam</i> HI)	CCC <u>AAGCTT</u> AC <u>GAATTC</u> TCGCCAATGGCGACGTGGTGGA TGCGGCCTCGC	ntrB mutagenesis
ntrB2-B (BamHI)	CG <u>GGATCC</u> CAGGATCGCGCCGAGCCGGGGCACGATG	ntrB mutagenesis
ntrB3-B (BamHI)	CCCTGC <u>GGATCC</u> AGATCGAGGACGACGGCCC	ntrB mutagenesis
ntrB4-K (KpnI)	ACGG <u>GGTACC</u> CGAAGGGCTTGGGCAGGTAGTCGAAGGC	ntrB mutagenesis
ntrY1-A (AatII)	GAT <u>GACGTC</u> GCGAGAGTCGCTGAACGGCCACAGAA	ntrY mutagenesis
ntrY2-B (BamHI)	CGC <u>GGATCC</u> CGAGGTCGAAACCACCTGCTGCACCC	ntrY mutagenesis
ntrY3-B (BamHI)	GCC <u>GGATCC</u> CTGGAGGGCTATGTGGTGGCGCTGGA	ntrY mutagenesis
ntrY4-E (EcoRI)	GCC <u>GAATTC</u> CTCGGGGGGCATCGGTCAGCATCAGGC	ntrY mutagenesis
PTF	CGCCACGCCGCATAATGGTCGAAGGGATAGT	<i>nasT::lacZ</i> fusion
PTR	GCGAGGGATTGGCCAGGTCGATCAGCA	<i>nasT::lacZ</i> fusion
PTNtrR	GCGGCGCTGCCCATGGTTCGGTAGGCG	<i>nasT::lacZ</i> fusion
PTNtrR_Mut	GCGGCGCT <b>AT</b> (GC)CC <b>C</b> (A)TGGTTCGGTAGGCG	Mutated <i>nasT</i> promoter:: <i>lacZ</i> fusion
PTNtrF	CGCCTACCGAACCATGGGCAGCGCCGC	<i>nasT::lacZ</i> fusion
PTNtrF_Mut	CGCCTACCGAACCA <i>G</i> (T)GG <i>AT</i> (CG)AGCGCCGC	Mutated and native <i>nasT</i> promoter:: <i>lacZ</i> fusion
FA1_A	CGACGGCACCATCTTCGA	nasA::lacZ fusion
RA1	GTAAAGGCGAAGGTCGAGAG	Mutated <i>nasA</i> promoter:: <i>lacZ</i> fusion
RA1-S (SphI)	TA <u>GCATGC</u> GTAAAGGCGAAGGTCGAGAG	Mutated <i>nasA</i> promoter :: <i>lacZ</i> fusion
FA1_B-P ( <i>Pst</i> I)	AT <u>CTGCAG</u> ACGGCACCATCTTCGAACCACC	Mutated <i>nasA</i> promoter (hairpin I):: <i>lacZ</i> fusion
RHI-A (AscI)	GCCGGT <u>GGCGCGCC</u> GTT <i>CC</i> (GC)G <i>C</i> (C)CGGGCCAT	Mutated <i>nasA</i> promoter (haipin I):: <i>lacZ</i> fusion

FHII-A (AscI)	AAC <u>GGCGCGCC</u> ACCGGCCCGCTCGCAATGATGCGGCAC	Mutated nasA
	GGAT <b>A</b> (T)C <b>GG</b> (CC)GCGA	promoter (haipin
		II):: <i>lacZ</i> fusion
AIF1 (EcoRI)	AA <u>GAATTC</u> GCGAATATCTGGACCTGCCGGCGGAA	Unmarked hairpin
		I mutant
AIR1a	ATGGGATCAGACTGGGTCAGGACGCA	Unmarked hairpin
		I mutant
AIF2 (SalI)	AA <u>GTCGAC</u> CGCTCGCAATGATGCGGCACGGATTC	Unmarked hairpin
		I mutant
AIR2 (HindIII)	TT <u>AAGCTT</u> CATTGCCGGTGCCGAAGATGCCCAAC	Unmarked hairpin
		I mutant
nasTR	GATGCGGCCATCGCACGCTTTCACA	RT-PCR
nasSF	GCGGCGTCGATCAGCGGCACATAGC	RT-PCR
CD		
nasSR	GUGATGGUGUATTICUGUAUUGATUTTAUU	RI-PCR
nasAF	ATGCGTTGCTGATGGCGAAGGCTGTCGGAAA	RT-PCR
		iti i oit
nasAR	CTGGGCAAGGCGGCCGTCTACAAGCACAT	RT-PCR
nasBF	CCCGACAGGACCGGCGACAGCATCAG	RT-PCR
		DT DCD
nasBK	TGGTCTGCGTCGACAGCGGCTACCAGATCAG	RI-PCK
nasGF	GCGGTGCCCGAGTTCATGTCGAACACCCAGTT	RT-PCR
nasGR	CTGCCCGCTGCACAACTGGGTGTTCGACATGA	RT-PCR
nasHF	GCGAAATTGCCGCAAAAGACCAGGCCCCAGTT	RT-PCR
magIID		
nasnk		KI-PCK
nasCF	CGTCGCCAGCACGCCGCAGCCCACGCCGCAATA	RT-PCR
RT-RpoB5'	TTTGTCGCGAAAGACATCATCAACGAGCAGAC	RT-PCR
		(housekeeping)
RT-RpoB3'	TGGGTATCCGGCGCATCCAGGTCAAGG	RT-PCR
DT 45'		(housekeeping)
KI-AJ	GATGTTCGTGGACCGCTCGGATGAG	qPCK (nas1 gene)
RT-A3'	GGAACATGTGAAAGCGTGCGATGGC	aPCR ( <i>nasT</i> gene)
		1(8)
RT-K5′	GGACGAGCCCGAGAACCGC	qPCR (nasS gene)
RT-K3'	CGGCGTGGAAGGCGATGAA	qPCR (nasS gene)
PT C5'		aDCP (nas 1 apro)
KI-CJ	GUGUIGAIGIAIGUGACUIICAICG	qi UK (nush gene)
RT-C3′	GCTGGGGATGTGCTTGTAGACGGC	gPCR (nasA gene)
-		1 (

RT-H5′	GCGACGTTCATCCGACCCATTACG	qPCR (rpoB-
		housekeeping)
RT-H3'	TGCCCTCGACCACCTTGCGGTAGG	qPCR (rpoB-
		housekeeping)
dnaN-1F'	CATGTCGTGGGTCAGCATAC	qPCR (dnaN-
		housekeeping)
dnaN-1R'	CTCGCGACCATGCATATAGA	qPCR (dnaN-
		housekeeping)
ntrB10	GCCATCATGGGCGAGGTTTTCGAGC	qPCR
ntrB11	CGGGGCGGGATTCCAGTGCGAT	qPCR
ntrY4	CGGTGAATATCGGGCTGGAGGGCTGG	qPCR
ntrY5	TGGCGTCGTTGGTCAAGTCTCGGCG	qPCR
nasT2F	TCGATCGTCGTCATCGAAT	qPCR to verify
		microarrays
nasT2R	GGTTTCTTCCGAGATGATGC	qPCR to verify
	01004001001041011001	microarrays
nasS2F	CIGGACCIGGIGAIGIICCI	qPCR to verify
nasS2R	ΑΤΓΩΓΩΑΔΑΩΤΓΩΑΔΑΤΓΑΤ	aPCR to verify
1105521		microarrays
nasA1F	CCTGACGCAGGTCTATGGTT	qPCR to verify
		microarrays
nasA1R	GACGATGAAGGTCGCATACA	qPCR to verify
		microarrays
nasB2F	TCTGGACCGGCCCTATAAAT	qPCR to verify
		microarrays
nasB2R	CGATAGACCGACTGGCTGAT	qPCR to verify
marC1E	CTCTTTCCCCTCCACCAC	microarrays
nasGIF	GICTIIGUGUIGGAUGAU	qPCR to verify
nasG1R	ΔΤΩΤΟΩΔΔΟΔΟΟΟΔΩΤΤΩΤΩ	aPCR to verify
moutiv	ATTECONOLOGICATION	microarravs
nasH1F	TCATGTGCAACTGGATGGTT	aPCR to verify
		microarrays
nasH1R	CATGTTCACGATGGAATGCT	qPCR to verify
		microarrays
nasC1F	CTATGTCGCCAACAAGCTGA	qPCR to verify
		microarrays

nasC1R	CCAGCTCCAGATCCTCGTAG	qPCR to verify microarrays
ntrB10	GCCATCATGGGCGACGGTTTTCGAGC	qPCR to verify microarrays
ntrB11	CGGGGCGGGATTCCAGTGCGAT	qPCR to verify microarrays
ntrC1F	ACCGCTATTTCGACCTGCAT	qPCR to verify microarrays
ntrC1R	GGTTTATGCCCAGAAGATCG	qPCR to verify microarrays
GIDH1F	ACGGCAACACCTATTTCACC	qPCR to verify microarrays
GIDH1R	ATTCCTGCACCTTCTCGATG	qPCR to verify microarrays

<sup>a</sup>Enzyme restriction sites are underlined. Bases changed are marked in bold and italics

and native bases are in brackets.





B



Figure S1

A		_		
NtrY	Pd Ab	1	-MAGTISKGTWERVARLERLYRNTATLGLALLGPALAGLTFRVMGPFADSTSGG-TVERLULADLLYLIVLTGLVVARMAQ	82
NtrY	AC	1	- MTOA BEDOAS DNAGHTPSGSSEG IFAD - ADVILALISALATELISALATELINGI. TPUVDEH - OVUSULUMAAAVI IT SAMYGEE WE	81
NtrY	Ba	1	-MNRANLTTDMNGTROSSODEGOGEGRRLLALPGIITVVSALVTASFSFAILIGITPITPDR-TVTLALVIINVALILFLILLICREVYR	88
NtrY	Cc	1	-MGGGYALAVILTVT	59
NtrY	Sm	1	MVDGMALPLGTEDGVTAAQDRRASFALPGLMLATGALICATLSLLVLLGLTPIRPER-NIVIACAGINGLFVVGLIYLIAREIFR	84
NtrY	Ml	1	MLALPGVVAVVGALVMAAISFTILVGATPIAPDA-RTTWALIALNAAFVLFLMALVGREVHR	61
NtrV	Ъđ	83		172
NtrY	Ab	87	WIGERBEIAGSOM WILWAVISTIAUVERTIMETESTVERVUCVOSVISERVERTAVNES LAVASAVILEEHOON TRADALAMANDIUNDEA	176
NtrY	Ac	82	IAKARARGRAAALIFIRIWGLGAVYSVVGAILVEVVASLTLDRCLDRWGSMRTOEIVASSVSVAOTWVREHALNIRGDILAMSADUTRLK	171
NtrY	Ba	89	IITARRLGKAASRLEVRIVALESLIAAIEAIVVEIVASVTLNLCLDRWEDQNTRDIISSSQNFYTAMLQETALNLQSTSYSMLQDLDAQR	178
NtrY	Cc	60	LIDAR-ASDAGARIH LRFVGLESLAAVAEAVIVELFFGVLVNRCVDGVESERVQTVVGNSAKVANSYVEQQKNYISEHIGPMAASINQAA	148
NtrY	Sm	85	LLRARSKGRAAARIH VRIVAIDSIVAIDSIVAID IFASIDUCLDRWHSLRTQAIVRSSLNVAQAVVLENASYLQGQDVSMANDHERNR	174
NtrY	Ml	62	IVMARRHGKRASRIH VRIMAMBALVAAIRAIMVRIIASITLDICLDRWBEIRTKTIVNSELSIADAVVQENARNLQGTTLSMAVDIDASR	151
NtrY	Pd	173	RONPMI.DDGET.RI.I.GOGOAL.I.ORGI.REZVUTD-GRGTTRARGERSYOFWYRRDSPUOFDRAATRGI.UL.I.RDWONNEFRA.UVDLVDI.ADR	261
NtrY	Ab	177	ARLASDPERFEQVVATQ AMLRALSER IVFNGTTGAIVARSGYTFALEFDPIPDDKLATARRGEVAMIVSENDDRVRALVRLDRFADT	263
NtrY	Ac	172	SVYegdrsrfnQiltaQaalrnlpgmlir-rdlsvveranvnigrefivpanlaigdatpdQpviylpndadyvaavvplkdyddlsveranvnigrefivpanlaigdatpdQp	257
NtrY	Ва	179	AIYSLDRNGFIQLMTLQVRGRALLGPTVR-DNGDVVVQSDTGMESHLPPPTKDSLEKAPDGKPVIIPPGNTNFVGALIKMREIPDM	264
NtrY	Cc	149	PALAQSPVAFGHFLAGLTQDNGFSAFYVLD-RDGRILARTEAEGAPPFLAPPPASFDWADGGEASAQRFVSTDLFRTLYKLKGFDDG	234
NtrY	Sm Ml	152	QLYSLDRTGFVELMTRQ ARGRGMLGFFLVR-ADGSALLQANISTDRPLPAIPQDALKSTIAGQPTLIPPGVTNLVGAVIPLENIPGT SLYGLDRTGFLDILNKF AVGRSLAHPALIK - DGSFVMSAOTDADFAMDRPPEGSVSSATDGKPVLTEDRTRNIMGATVKLRETEGL	260
		100		237
NtrY	Pd	262	YLYVTRDVDGSLLGLLDDTRQTVGDYQRLEQERGRVLFEFSLLYLGFATLLVAA?MMLGLWFADRLSRFIGRTAEBSEQNGECNDDFQIP	351
NtrY	Ab	264	Y MY YGR WVEPRVLSHMASAEGAVREFGALESORGSLQITETLIFLCVATLLLAR VNAGLIFATRLVR ISATIGRADRURACOT TVRVT	353
NtrY	Ac	258	YLYVARLIDPRVIGYLKTTOETLADYRSLEERRFGVQVAFALMYAVITTIVLLSZVMLGLNFSKWLVAFIRRIMSMADHVAEGNTDVRVP	347
NtrY	Ba	265	YLYTYN SYPPOIIDAINLMNANARRYDEMDANRIPTOIAALLYYGETNIVLESEINTGIAVADRLVRFIRIDIGESDIVAACA DVSVP	354
NtrY	Sm	235	HEYVYR FERSELENDELEIQDAUS IRDARKSRGRIQA HUGIS ILDIAEUVLIKE INVGIAAANS LAGEVACIV BAARNS GODDARVS VYTYR VND PEVNR SMEIMEENTARVKTI RAGETSIO I AGUVI VIGRAI IVII LAZIMTA IAVAD IVDE IROLI GAADSI SKUD VVVD	324
NtrY	Ml	238	YLYTIRLVDPEVIKAROIVESNTDEYRNLEDNRRTSOVABALPYLSITTIIILSEINTGIAVADRLVRIIRGIIGAADEVATENTDVAVP	327
			* * * #	
NtrY	Pd	352	APDTGDEIQTIGESENRMTROLKOROELVESYRAADDORRLFDSVISSYTAGVIGLDAAGEIDFLNRSATRLLGL-DPATAHD-RLFS	439
NtrV	AD Ac	304	EKPADULALDSRAHKNII ISIESSRAALDSRAHKIIDSRAKIISIOOVSEGNUGUDAESKIIDSRAKADU VADASSIIO-MKIAS	442
NtrY	Ba	355	TIRAGULASARI HANIRELKSCRAFTI IARUTI SATUR UN SATURAL TIRAGULASARI TIRAGULASA	442
NtrY	Cc	325	VERGPEEIRALSNAFNMMTSELQLQQAALKAASLDAESRRQFIETVLSGVSAGVVSLDDRGKISAVNRRAVALLGLPDDALGVDLIA	411
NtrY	Sm	351	$v \texttt{Ravdgdvgnlsrtenkw} v \texttt{Ravdgdvgnlsrtenkw} \\ Ravdgdvgnlsrtenk$	438
NtrY	Ml	328	VRSSDGDVAS GDTEN KULLELKSCRNEILSAKDLIDERRESEAVIACVTAGVIGVDPYGIVTIVNRSAESMLAISASAALGQNISA	415
			*	
NtrY	Pd	440	AVEEFAPLFERLAQSVNESVQDEIRLTRECRV-ESLLVRMAIRR-GAAGGLEGYNVALDDVTDLVS-AORMAANGDVARRVRHEIKNE	524
NtrY	Ab	443	LVPEMGGLLHEAPGRPGLVVQDQIKIR <mark>RDC</mark> TTPLTLLVRISTE-GRGSGMMRGYVVFDEHHRTWSPAQRKAAMARRRPIAASPTRVKNP	531
NtrY	Ac	436	VVETAGLEEAEHARQRSVQGNITLTRDCRE-RVFAVRVTTE-QSPEAEHGWVTIDDTELIS-AQRTSAMADVARRIAHEIKNP	519
NErY	Bа	443	IAPEEVGQAFEVARATGRIVHREQVSMARGEGS-KSFNOVIVE-DAESEDHSIVVIVDDIIDLVQ-AQRSSAWADVARKIMHEIKNP	100
NtrY	Sm	439	LARGE EI WIIDSESSEZED ID EI DIRACETE - RINAVE - AGDESHGSVUITVDITDIVII-BORSTÄMADVARTIR - EL KNE	522
NtrY	Ml	416	ILPHVGRVFEIGRQSGKPVYREQVFFFRAGTE-RTFNVQITIEAGDDGSEEKSYNVTVDDITDLVQ-AQRSSAWADVARRIAHEIKNP	501
	- 1			
NErY	rd Ah	525 520	ITELOISANGKUKKKEGEIAG-BEKEALEQIIEVIIKQIND EKRIVDENSKEARMEELDKKETDIAKULKDALLMQKDALQGALVSDIP	67U
NtrV	Ac	520	TTPIOLSARELIKERGENT ODERTERGENTIGENTIGENTIGEN VOLTAGEN VERFORMEREN VERFORMER	607
NtrY	Ba	527	LTPIOLSAERIRRRYGKVIT-EDREVFDQCDDTIIRCVGDIGENVDEFSAFARMEKEEMRHMDLREALREASFLIEVSRS-DIHFDNDFG	614
NtrY	Cc	491	LTPIQLSAERLRRKYRKEIA-SDLETFDRCTDT <mark>IIRQ</mark> VGDIG <mark>RWVDEFS</mark> A <mark>FARMPA</mark> ERFEPSNLTEMLRQAVFAQRFQDAEIEVRLEEPG	579
NtrY	Sm	523	LTPIQLSAERLKRRYGKQIDQEDRAVFDQCHETIVRCVEDIG <mark>RWVDEFS</mark> AFARMEK <mark>E</mark> TKEKSDLRAILKDAVFLREMGNS-HINFVRDFG	611
NtrY	Ml	502	<b>ITPIQISAE</b> RIKRRYGKVIT-EDREVFDQCDDT <b>UIRQVEDI</b> G <b>RMVDDES</b> AF <b>ARVE</b> KEEMKAIDLRESLREASFLVEVSRA-DITFERIFG	589
			+ + + + +++ ++	
NtrY	Pd	612	DEPWIVDABAGMMRGVFTKLLKNAGBERIDELRENPPEDWTPOVHVKLRVAPDSVAIRTIDKERGHA-DDRTRLFARENVAMMKPG	698
NtrY	Ab	621	QGPLTVPORSRQISCALTNLLQNAADAIEGRPPPAEGTELPPGHVAIRVEADAERIAMIIEDNCKSLPTEERDRLTEPYVTTRAKCTGLC	710
NtrY	Ac	608	P-AMPARFORRLVSCALTNILKNAABAIEAVPPDVRGQGRIRVSANRVGEDLVIDIIDNGTGLPQESRNRLLBPYVUTREKGTGLG	692
NtrY	Ba	615	SEKLIGSFOSKLIG AFGNVIKNASETIDAVAKEERGEGHIRIHAVRADGQUVVVUDVIDNEKCIPBDDRQKILEPMMTREKGFGIG	700
NEYN	CC Sm	58U 612	DEDUNTISERNINGVALINILAWEGENVGARKANEPELQGRITATEVCEGDELCVIVEDNGVEDEAKDREIDEI DEV VITREKENGLE DEDIRGOED GRMICCAEGNIVENEVENTRAVDAGTSRG-APTVVIPSPRDDTTGPRVDDVIDNGVEDEAKDREIDEI DEMAMMODVEDELCVIV	555 700
NtrY	Ml	590	NEPLKGTFUSRLAGAFGUVIKWASEPIDGLEOKDGSHGIIRIOAGRONGAIRIDVIDMCKEIPRENRORIJEDMINTREKENRIG	675
	-			
			+ + + + + +	
NtrY	Pd	699	PUYKKUYEEHGGSIMUTDAPEGRCAMAETRLPRERQPVRLSARRHKHEQETLKRQPVRLSARRHKHEQETLK	752
NtrY	Ab Ac	711	AALVIKKHWEDUGGVETTEDREGGCREVGVIPUHIPASGTAAGDAPGGVGTPAETGEEKRHAAHGA	777
NtrV	ас Ва	693 701	INTERNET BOTT DE TRANSFERENT ET STELLEN BEREKEN DE STELLEN BEREKEN BEREKEN DE STELLEN BEREKEN DE STELLEN	760
NtrY	Cc	667	ALLYKRIMED IGGSLALVDARE	716
NtrY	Sm	701	ALVKKIIPDHGGDLELHDAPPDFDGGAGAMIRVVLPPAGETGGEDNLKDKGNTNGG	757
NtrY	Ml	676	AAIVKKUVEDHGGRLENHDAPADFHGGRG2MISIILPLAAVTPLRGEGKTEHERETEKVGNGV	738

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		*	
NtrX Pd NtrX Ab NtrX Ac NtrX Ba NtrX Cc NtrX Sm NtrX Ml	1 1 1 1 1 1	-MSEITIVDDERDIRELISDITEDEGYSTRLMANSDOAVEELNAAEFATMILDIWLKDSKMOGIDILKOVKRNNPDVPVVIISGHGNIEI MAHDIIIVDDEADIRMLIAGITNDEGMKTREAADADOAFAOVSARRESIVVLDIWLOGSRLDGLOIDEOLMRDHRNLPVIMISGHGNIET MAHDIIIVDDEPDISGLVAGITEDEGYSARTARDADOALAELAARRENIFLDIWLOGSRLDGLEIDDIKKEHPEVPVVMISGHGNIET MAADIIVVDDEVDIRDLVAGITSDEGHETRTAFDADSALAAINDRAPRIVFLDIWLOGSRLDGLEIDDIKKEHPEVPVVMISGHGNIET MAADIIVVDDEVDIRDLVAGITSDEGHETRTAFDADSALAAINDRAPRIVFLDIWLOGSRLDGLEIDDIKKALDADLPVVMISGHGNIET MAADIIVVDDEADIRCLVAGITSDEGHETRTAFDSSALAAINDRAPRIVFLDIWLOGSRLDGLALIDEIKKNHPDLPVVMISGHGNIET MAADIIVVDDEADIRCLVAGITSDEGHETRTAFDSSALAAINDRAPRIVFLDIWLOGSRLDGLALIDEIKKNHPDLPVVMISGHGNIET MAADIIVVDDEADIRCLVAGITSDEGHETRTAFDSSALAAINDRVPRIIFLDIWLOGSRLDGLALIDEIKNRHPDLPVVMISGHGNIET MAADIIVVDDEDIRCLVAGITSDEGHETRTAFDSSALAAINDRVPRIIFLDIWLOGSRLDGLALIDEIKNRHPDLPVVMISGHGNIET	89 90 90 90 90 90 90
		* ** +++++++	
NtrX Pd NtrX Ab NtrX Ac NtrX Ba NtrX Cc NtrX Sm NtrX Ml	90 91 91 91 91 91 91	AVAAI KOGAYDFIEKPFNID OLLUVIRRAMETARI RRENATIKRGEARAAEMIGNSAPFRIREQLDKWAKSNGR VMLTGEFGAGKEIAA AVSAI KIGAYDFIEKPFKAD RULLMVDRAIBAARI KRENEELKLRAGGEVELIGRSTAVNHVROSIEKVSPTGSEVLITG PAGSGKBUVA AVAAI KRGAYDFIEKPFKAD RUVITERALETLRI RREVRELKQLT-OPHTWYGRSSVLOURAVVDROGPTNSEILIV GPSGSGKBUTA AVSAI RRGAYDFIEKPFKAD RULLVABRALETSKI KREVSDURKRTGDOLELVGTSLAMNQIR QTIERVAPTNSEIMITGPSGAGKBUVA AVSAI RRGAYDFIEKPFKAD RULLVABRALETSKI KRENSELRKSGDPVELIGTSVAVSQURQMIEKVAPTNSEIMIGEPSGAGKBUVA AVSAI KRGAYDFIEKPFKAD RULLVERALBAAGI REENRRIRAGTLSPDGLIG KSAPAQAI RQLILKVAPANSE VLVG PFGAGKBUVA AVSAI KRGAYDFIEKPFKAD RULLVERALBAAGI REENRRIRAGTLSPDGLIG KSAPAQAI RQLILKVAPANSE VLVG PFGAGKBUVA AVSAI KRGAYDFIEKPFKAD RULLIVERALBAAGI REENRRIRAGTLSPDGLIG KSAPAQAI RQLIKVAPANSE VLVG PFGAGKBUVA AVSAI KRGAYDFIEKPFKAD RULLIVERALBAAGI REENRIKRAGTLSPDGLIG KSAPAQAI RQLIKVAPANSE VLVG PFGAGKBUVA AVSAI RRGAYDFIEKPFKAD RULLIABRALBASKI KRENSELRRKSGDPVELIGTSVAVSQURQMIEKVAPTNSE IMIG PSGSGKBLAA	179 180 179 180 180 180 180
NtrX Pd NtrX Ab NtrX Ac NtrX Ba NtrX Cc NtrX Sm NtrX Ml	180 181 180 181 181 181 181	RYVHAQSPRAREPEVTVDCATIFEEHMEEVLEGRETAEREIEPGLIEOAHGCVIYFDEVADMHLGTOPKILRVETEGOBCRAGGTDKV RLIHARSRRAGGPEVGLNCATMREDRLEMELFGIEAGVDCGGRKIGTFEOAHGCTLLLDEVADMHLETOGKIVRALOEOVSERVGGGORV RWIHAASARAQGPEVVINAAATTEERLEYELFGUE-GEGRERKGALEEAHGCTLFIDEIADMERETONRVLRVEVCETESRIGSSEXV RAIHAQSSRANGPEVVINAATTEERLEYELFGIE-OMGGERKVGALEEAHGCILFIDEVADMERETONRVLRVEVCETESRIGSSEXV RLIHGASTRARQEFVAVSAFGMAFERLDVELFGEG-EGGRERKIGVFERHGCILFIDEVADMERETONKILRVEVCEGERVGGTKRV RWIHRKSARANGEFVALNAAATTEERLEYELFGIE-OMGGERKVGALEEAHGCILFIDEVADMERETONKILRVEVCEGERVGGTKRV RLIHGASTRARQEFVAVSAFGMAFERLDVELFGEG-EGGRERKIGVFERHGCILFIDEVADMERETONKILRVEVCEGERVGGTKRV RMIHRKSARANGEFVALNAAATTEERMEIELFGIE-TTCOPRECALEEAHGCILFIDEVADMERETONKILRVEVCEGERVGGKRV RAIHTLSARKGAPFVTLSAANTTEERMEIELFGIE-SNEVERKVGALEEAHRGILVIDEVADMERETONKILRVEVCEGERVGGTKRV	267 270 268 268 269 269 269
NtrX Pd NtrX Ab NtrX Ac NtrX Ba NtrX Cc NtrX Sm NtrX Ml	268 271 269 269 270 270 269	RVDIR VISSTNRDIAAEIAASREROE VDRINVVE VAVESIAERE DDIEVLATHEIELFHASGGLPCRALPEETSBALGAMNWEGNIROL EVDVR VVATSNRDIOAEIDOSREROD FYRIAVVE IRVESIAERE EDIELLAREFMORSAEAGLPARDFGEDAMAALGAYDWEGNVROL RVDVR IISSTGRHEEEIAASRERED YHRISVVE IRVEPLAERE EDIELLAREFMORSAEAGLPARDFGEDAMAALGAYDWEGNVROL KVDVR IISSTGRHEEEIAASRERED FHRISVVE VVEPLAERE EDIELVDFFINGIAEGAGIKERKIGPDAMAVLGAHSWEGNIROL KVDVR IISSTGNIEGMIAESTERED FHRISVVE VVEPLAERE EDIESLVEFFMKOIAEGAGIKERKIGPDAMAVLGAHSWEGNIROL KVDVR IISSTANTEGMIAESTERED FHRISVVE VVERLAERE EDIESLVEFFMKOIAEGAGIKERKIGEDALAVLGAHSWEGNIROL KVDVR IISSTANTEENTERED FHRISVVE VVERLAERE EDIESLVEFFMKOIAEGAGIKERKIGEDALAVLGAHSWEGNIROL KVDVR IISSTANTEENTEELERED FHRISVVE VVERLAERE EDIESLVDFFMROVSEGAGIKERKIGEDALAVLGAHSWEGNIROL KVDVR IISSTSONTEAMIAGRERED YHRIAVVE VMVEGLAERE EDIESLVDFFMROVSEGAGIKERRIGEDALAVLGAHSWEGNIROL	357 360 358 358 359 359 359 358
NtrX Pd NtrX Ab NtrX Ac NtrX Ba NtrX Cc NtrX Sm NtrX Ml	358 361 359 359 360 360 359	RNVIERVIE LGDGTGFIQPSEIESQNGS-AGTSEALALGPOITSLALREARE LFEREYLLAQINRFGGNISRTAQE VGMERSALHRK RNVVDWLLIMAQGD-PKEPIRADQIPPEIGAITPTVLKWDKGGEIMGUELREAREVFEREYLLAQINRFGGNISRTASE VGMERSALHRK RNVVERLILLGGD-PDAEVTASHIPPDVGALVPTLPNGNGGEHLMGUELREAREVFEREYLAQINRFGGNISRTASE VGMERSALHRK RNVVERLILLGGD-PDAEVTASHIPPDVGALVPTLPNGNGGEHLMGUELREARERFEKEYLAQINRFGGNISRTASE VGMERSALHRK RNVVERLILLGGD-PDDELVTADIIPAEIGDTLPRAPT-ESDQHIMALELREARERFEKEYLAQINRFGGNISRTASE VGMERSALHRK RNNVERMILLSGE-PGDVITAEMISGAEQFSGNAGA-IGAERIALELREARERFEKEYLAQINRFGGNISRTASE VGMERSALHRK RNNVERMILLSGE-PGDVITAEMISGAEQFSGNAGA-IGAERIIALELREARERFEKEYLAQINRFGGNISRTASE VGMERSALHRK RNNIERLMILLASGE-PGDVITAEMISGAEQFSGNAGA-IGAERIIALELREAREMPERDYLIAQINRFGGNISRTASE VGMERSALHRK RNNIERLMILASGDPDTPITADMIPNEVGDTLPKISA-QGDQHIMALELREAREMPERDYLIAQINRFGGNISRTASE VGMERSALHRK RNNVERLMILAGDDVDAPITADIIPSEIGDVMPRTPN-QSDQHIMALELREAREMPERDYLIAQINRFGGNISRTASE IGMERSALHRK	443 449 447 447 447 448 448 447
NtrX Pd NtrX Ab NtrX Ac NtrX Ba NtrX Cc	444 450 448 448 448	LKSLGVVGGMRVEEEMMGK 462 LKSLGVHGSEKGKLFVE 466 LKALGVG 454 LKSLGV 453 LKSLGVSGARGDEEE 462	

NtrX Sm 449 LKSLGV----- 454 NtrX M1 448 LKSLGV----- 453

Figure S2



Figure S3

### FIGURE LEGENDS

**Figure S1. Phylogenetic trees with NtrB (A) and NtrC (B) homologues.** The evolutionary history was inferred using the Neighbor-Joining method [11]. The bootstrap consensus tree was inferred from 3000 replicates [12]. Branches corresponding to partitions reproduced in less than 50% bootstrap replicates are collapsed. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (3000 replicates) are shown next to the branches [12]. The evolutionary distances were computed using the p-distance method and are in the units of the number of amino acid differences per site. The analysis involved 18 amino acid sequences. All positions containing gaps and missing data were eliminated. There were a total of 330 positions in the final dataset. Evolutionary analyses were conducted in MEGA5 [13]. The *P. denitrificans* sequences are highlighted by arrows.

Figure S2. Sequence alignment of the two-component regulatory system NtrY (A) and NtrX (B). Alignment was performed using ClustalW. Identical and similar residues are shaded in black and grey respectively. Regions of selected sequences are highlighted by boxes (located below their respective sequences) and show significant matches with the protein family database (Pfam-A, http://pfam.sanger.ac.uk/) for the following domains: A. HAMP linker domain (blue box), PAS domain involved in many signaling proteins where they are used as a signal sensor domain (red box), phospho-acceptor domain (green box), ATPase domain of histidine kinases (black box). B. Response regulator receiver domain (blue box), sigma54 interaction domain (red box); and helixturn-helix (HTH) structural motif capable of binding DNA (green box). Asterisks define the position of conserved residues at the active site, triangles indicate the phosphorylation site, crosses define the ATP binding site, and points indicate the heme binding pocket. Pd, Paracoccus denitrificans PD1222 (NtrY, gi|119377428 and NtrX, gi|119377427); Ab, Azospirillum brasilense Sp245 (NtrY, gi|20451353 and NtrX, gi|119377427); Ac, Azorhizobium caulinodans ORS 571 (NtrY, gi|38737 and NtrX, gi|38738); Ba, Brucella melitensis biovar Abortus 2308 (NtrY, gi|123754544 and NtrX, gi|82699956); Cc, Caulobacter crescentus CB15 (NtrY, gi|16125986 and NtrX, gi|16125987); Sm, Sinorhizobium meliloti SM11 (NtrY, gi|336033438 and NtrX, gi|336033437); Ml, *Mesorhizobium loti* MAFF303099 (NtrY, gi|14021375 and NtrX, gi|14021376).

Figure S3. Schematic overview of the nitrate-regulated systems in *Paracoccus denitrificans*. Transcriptomic data reveal systems that are up-regulated ( $\uparrow$ ) or down-regulated ( $\downarrow$ ) by nitrate. The fold changes are indicated. Proteomic analysis also corroborated these results.

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