

nosX is essential for whole-cell N₂O reduction in *Paracoccus denitrificans* but not for assembly of copper centres of nitrous oxide reductase

Sophie P. Bennett¹, Maria J. Torres², Manuel J. Soriano-Laguna², David J. Richardson², Andrew J. Gates^{2,*} and Nick E. Le Brun^{1,*}

Abstract

Nitrous oxide (N₂O) is a potent greenhouse gas that is produced naturally as an intermediate during the process of denitrification carried out by some soil bacteria. It is consumed by nitrous oxide reductase (N₂OR), the terminal enzyme of the denitrification pathway, which catalyses a reduction reaction to generate dinitrogen. N₂OR contains two important copper cofactors (Cu_A and Cu_Z centres) that are essential for activity, and in copper-limited environments, N₂OR fails to function, contributing to rising levels of atmospheric N₂O and a major environmental challenge. Here we report studies of *nosX*, one of eight genes in the *nos* cluster of the soil dwelling α -proteobacterium *Paracoccus denitrificans*. A *P. denitrificans* Δ *nosX* deletion mutant failed to reduce N₂O under both copper-sufficient and copper-limited conditions, demonstrating that NosX plays an essential role in N₂OR activity. N₂OR isolated from *nosX*-deficient cells was found to be unaffected in terms of the assembly of its copper cofactors, and to be active in *in vitro* assays, indicating that NosX is not required for the maturation of the enzyme; in particular, it plays no part in the assembly of either of the Cu_A and Cu_Z centres. Furthermore, quantitative Reverse Transcription PCR (qRT-PCR) studies showed that NosX does not significantly affect the expression of the N₂OR-encoding *nosZ* gene. NosX is a homologue of the FAD-binding protein ApbE from *Pseudomonas stutzeri*, which functions in the flavinylation of another N₂OR accessory protein, NosR. Thus, it is likely that NosX is a system-specific maturation factor of NosR, and so is indirectly involved in maintaining the reaction cycle of N₂OR and cellular N₂O reduction.

INTRODUCTION

Nitrous oxide is a potent greenhouse gas which has rapidly increased in the atmosphere over the past century [1]. The rise in N₂O coincides with the introduction and application of anthropogenic nitrogen species in agriculture, to improve crop yield and ultimately feed the growing global population [2, 3]. Of the total N₂O released, 40% is produced by soil bacteria [4]. Soil dwelling denitrifying micro-organisms such as *Paracoccus denitrificans* consume nitrate as an alternative electron acceptor during anaerobic growth conditions. N₂O is an intermediate substrate in the denitrification pathway; it is reduced to N₂ by the copper enzyme nitrous oxide

reductase (N₂OR). N₂OR-containing bacteria can be separated into two clades, and a feature that distinguishes the clades is the ability of the micro-organism to produce and consume, or only consume, N₂O [5, 6]. Clade-I members are complete denitrifiers with the nitrite reductase genes *nirS* or *nirK* present in their genome. In contrast, about half of the clade-II members are non-denitrifying N₂O reducers, and are therefore N₂O sinks [5]. Ammonia-oxidizing bacteria (AOB) are another microbial source of N₂O in coastal ecosystems, through a process named 'nitrifier denitrification'. However, they do not harbour genes encoding N₂O reduction activity [7]. Environmental factors such as soil pH, Cu content, and moisture impact on N₂O emissions from soil [8–10]. In order

Received 13 May 2020; Accepted 07 July 2020; Published 04 September 2020

Author affiliations: ¹Centre for Molecular and Structural Biochemistry, School of Chemistry, University of East Anglia, Norwich Research Park, Norwich, NR4 7TJ, UK; ²Centre for Molecular and Structural Biochemistry, School of Biological Sciences, University of East Anglia, Norwich Research Park, Norwich, NR4 7TJ, UK.

***Correspondence:** Nick E. Le Brun, n.le-brun@uea.ac.uk; Andrew J. Gates, a.gates@uea.ac.uk

Keywords: copper; denitrification; nitrous oxide; nitrous oxide reductase.

Abbreviations: BCS, bathocuproinedisulfonic acid; cDNA, complementary DNA; LC-MS, liquid chromatography-mass spectrometry; MV, methyl viologen; NGC, *nos* gene cluster; N₂OR, nitrous oxide reductase; OD, optical density; qRT-PCR, quantitative reverse transcription-polymerase chain reaction.

One supplementary table and two supplementary figures are available with the online version of this article.

000955 © 2020 The Authors



This is an open-access article distributed under the terms of the Creative Commons Attribution License. This article was made open access via a Publish and Read agreement between the Microbiology Society and the corresponding author's institution.

to identify N₂O mitigation strategies, we are trying to understand the optimal genetic components needed to biologically remove N₂O.

Nitrous oxide reductase (N₂OR) is a homo-dimeric, ~120 kDa, multi-Cu protein. Each monomer contains two Cu cofactors, the Cu_A and Cu_Z centres, responsible for electron transfer and the catalytic reduction of N₂O, respectively. The Cu_A centre is a bis-thiolate-bridged di-nuclear Cu centre, accommodated within a cupredoxin fold domain, similar to that of subunit II of cytochrome *c* oxidase. The Cu_Z centre is a unique [Cu-S] cluster ligated by seven conserved histidine residues within a β-barrel domain. It comprises four Cu atoms and one or two sulphur atoms, depending on the purification method [11–13]. Notably, the subunits of the active homodimer are orientated in a head to tail configuration, with one Cu_A centre in close proximity to the Cu_Z centre of the other monomer. N₂OR is encoded by the *nosZ* gene, which, in denitrifying organisms such as *Paracoccus denitrificans* and *Pseudomonas stutzeri*, is translated and exported through the twin-arginine transport [14] pathway to the periplasm, as a folded apo-protein, before acquiring its Cu cofactors. Consistent with this, a TAT signal leader sequence mutant accumulated unprocessed, dimeric, apo-protein in the cytoplasm of the cell [15]. In contrast, the N₂OR of clade-II members are transported through the Sec pathway [16]. The functional significance of this is currently unknown.

The *nosZ* gene is found among the *nos* gene cluster (NGC), which comprises eight genes in *P. denitrificans*: *nosCRZD-FYLX*. The *nosC* and *nosR* genes are copper responsive in *P. denitrificans* and function in the regulation of *nosZ* transcription. During Cu limitation, *nosCR* transcription is increased, whilst *nosZ* transcription is reduced [17]. In *Pseudomonas stutzeri*, NosR is a cytoplasmic membrane protein with two soluble domains located at either side of the membrane: the N-terminal periplasmic domain covalently binds a flavin mononucleotide, while the C-terminal cytoplasmic domain binds two [4Fe-4S] clusters [18]. The *P. denitrificans* homologue (44.3% identical) is predicted to have similar features. The function of NosR is not well understood; in addition to the regulatory role mentioned above, it is important for whole-cell N₂O reduction [17, 18], with evidence indicating that it is not involved in the assembly of the Cu centres of N₂OR, but may be the physiological electron donor to NosZ [18].

nosDFY encode a cytoplasmic membrane spanning ABC-type transporter that functions in the maturation of the Cu_Z centre of N₂OR, as illustrated by an insertional mutation in *P. stutzeri* *nosD*, which produced an N₂OR without the key spectroscopic signal of the Cu_Z centre [19]. Similarity to mitochondrial ABC transporters that export a sulphur species to the cytoplasm for iron-sulphur cluster biogenesis suggests a role for NosDFY in providing the essential sulphur atoms of the catalytic Cu_Z centre [20]. The *nosL* gene is well conserved across NGCs and is essential for whole-cell N₂O reduction in *P. denitrificans*. NosL is a Cu-binding lipoprotein, putatively anchored to the outer membrane of the cell. The properties of N₂OR purified

from a PdΔ*nosL* strain revealed that Cu-binding NosL is a component of the Cu_Z maturation apparatus under Cu replete conditions and, more importantly, is an essential maturation factor for both Cu centres during Cu limitation [21].

The *nosX* gene is predominantly found in α- and β-proteobacterial NGCs in clade I but does not feature among γ-proteobacteria nor clade-II NGCs (Fig. 1). NosX is a soluble protein of ~30 kDa, which is exported to the periplasm by the Tat pathway. Previously, it was reported that insertional mutagenesis of *P. denitrificans* *nosX* resulted in wild-type-like growth [22]. Interruption of both *nosX* and the homologue *nirX* did, however, present a Nos-negative (Nos⁻) phenotype, leading to the conclusion that NosX and NirX are functional homologues [22]. Furthermore, this study demonstrated that the *nosX nirX* double mutant strain contained N₂OR that was deficient in its Cu_A centre, implicating these proteins in copper cofactor assembly [22].

The γ-proteobacterium *P. stutzeri*, which does not feature *nosX* in its NGC, instead contains a NosX homologue encoded elsewhere on the genome. The protein, called ApbE, was shown to be a FAD-binding flavinyl transferase that serves as a flavin donor to NosR, which in turn activates N₂OR [23]. PsApbE and PdNosX share 32% amino acid homology, in particular the conservation of key amino acid residues associated with flavin binding suggest that their roles are similar while their genetic context implies they may differ in system specificity. Here, we present a re-examination of the role of NosX in *P. denitrificans*, through the analysis of full *nosX* deletion in *P. denitrificans*, in terms of cell growth and the properties of N₂OR purified from an unmarked mutant background. The data show that NosX is essential for N₂OR activity and cannot be substituted by NirX. Furthermore, NosX plays no role in assembly of the NosZ Cu cofactors, nor does it have a major function in the regulation of *nosZ* expression. Instead, the role of NosX is consistent with a system-specific maturation factor for NosR to support the activity of NosZ *in vivo*.

METHODS

Construction and complementation of a *nosX*-deficient strain of *P. denitrificans*

A double allelic exchange method was employed to generate a whole *nosX* gene deletion strain (Table S1, available in the online version of this article), as described previously [17, 21]. Briefly, the suicide plasmid pK18*mobsacB* containing DNA regions that flank the *nosX* gene (pSPBN4) was conjugated into PD1222 using the *E. coli* helper plasmid pRK2013. Single cross-over recombination events resulted in Spec^R/Km^R transconjugants, from which a double cross over mutant (Spec^R), named PD2502, was generated. The mutated region was PCR amplified and confirmed by sequencing.

PdΔ*nosX* (PD2502) was complemented *in trans* using pSPBN5, which contains the coding sequence of Pden_4214. The gene was synthesized by Genscript with flanking 5' *NdeI* and 3' *EcoRI* restriction sites and subcloned into a taurine inducible modified pLMB509 derivative with gentamycin

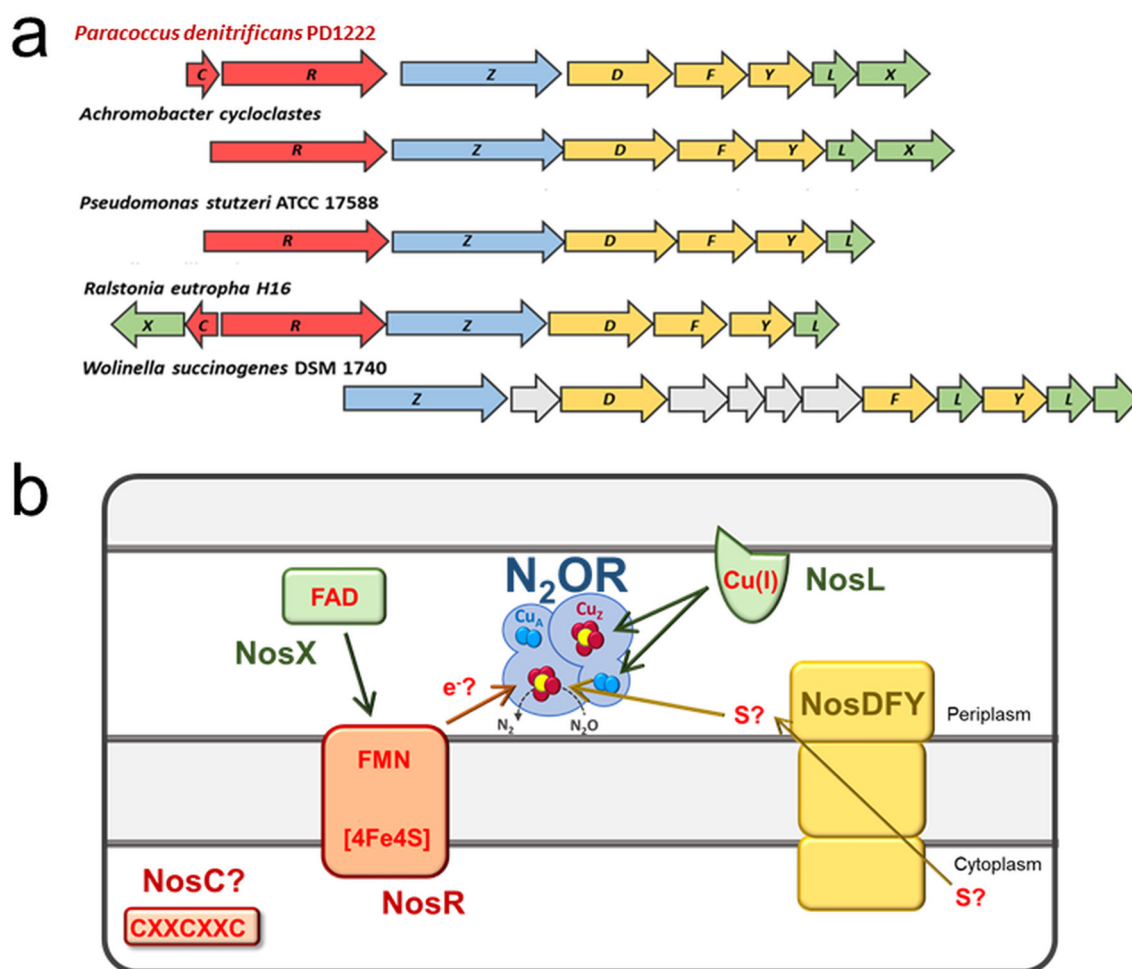


Fig. 1. (a) Comparison of NGCs from clade-I nitrous-oxide-reducing bacteria (*P. denitrificans*, *Achromobacter cycloclastes*, *Pseudomonas stutzeri*, *Ralstonia eutropha*) and the clade-II member *Wolinella succinogenes*. (b) The core *nosZDFYL* genes encode the nitrous oxide reductase polypeptide (NosZ), an ABC transporter complex (NosDFY) that is essential for Cu_2 centre maturation, and a Cu centre maturation factor (NosL). The *nosR* and *nosX* genes are less conserved across the two clades. NosR is a transmembrane iron-sulfur cluster containing protein with an FMN moiety, which is obtained from an ApbE-type flavinyltransferase (proposed as NosX here). Together the function of these proteins may involve supplying electrons to N_2OR for catalytic turnover and, where absent in the NGC, a homologue is likely to be found elsewhere in the genome.

resistance ($20 \mu\text{g ml}^{-1}$) to generate pSPBN5. The complementation plasmid was conjugated into the mutant strain using the helper *E. coli* pRK2013 strain, with successful conjugants identified as $\text{Spec}^{\text{R}}/\text{Gm}^{\text{R}}$. Expression of *nosX* from the plasmid was induced by adding 1 mM taurine to the medium at the start of growth.

Growth and phenotypic analysis of cultures

Anaerobic minimal media batch cultures (400 ml) were grown in sealed Duran flasks fitted with a septum seal to allow for gas-tight sample extraction. Minimal media consisted of: 30 mM succinate, 20 mM nitrate, 11 mM dihydrogen orthophosphate, 29 mM di-sodium orthophosphate, 0.4 mM magnesium sulphate, 1 mM ammonium chloride, pH 7.5. The minimal media was supplemented with a 2 ml l^{-1} Vishniac and Santer trace element solution [24] where copper sulphate

was present (Cu-sufficient, $12.8 \mu\text{M}$) or excluded (Cu-limited, $<0.5 \mu\text{M}$) from the original recipe. Media were inoculated using a 1% inoculum from a starter culture to give a starting $\text{OD}_{600 \text{ nm}}$ of ~ 0.02 and incubated at 30°C . Samples of the liquid culture were taken in 1 ml aliquots and $\text{OD}_{600 \text{ nm}}$ measured. The 3 ml gas samples were removed from the headspace of the cultures and stored in pre-evacuated 3 ml Exetainer vials. A 50 μl gas sample was injected into a Clarus 500 gas chromatograph (PerkinElmer) equipped with an Elite-PLOT Q (30 m \times 0.53 mm internal diameter) and an electron capture detector. Carrier gas was N_2 , make-up gas was 95% (v/v) argon, 5% (v/v) methane. Standards containing N_2O at 0.4, 5, 100, 1000, 5000 and 10000 p.p.m. (Scientific and Technical Gases) were measured and total N_2O was determined as previously described [17].

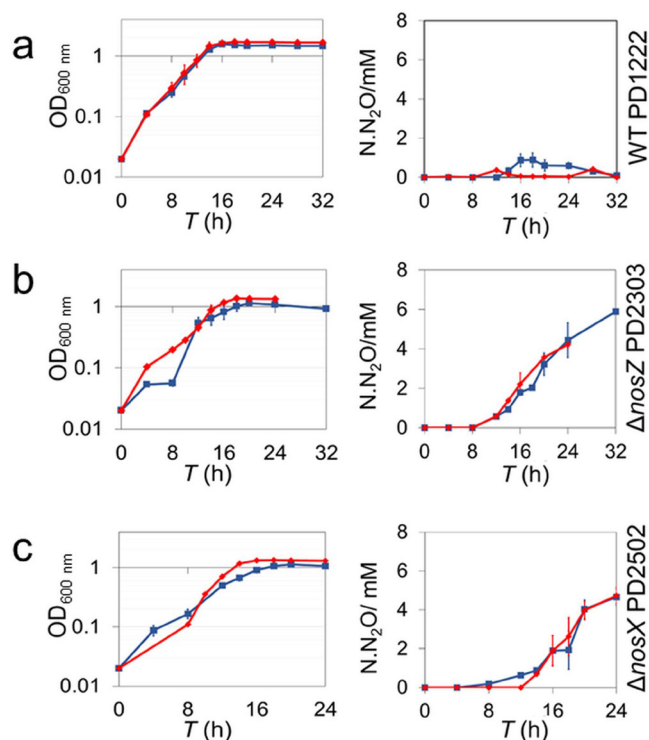


Fig. 2. Growth and N_2O production characteristics of *P. denitrificans* strains. (a) $OD_{600\text{ nm}}$ as a function of time (left) and N_2O emissions as $N.N_2O$ (millimolar N in the form of N_2O , right) for wild-type PD1222 grown in anaerobic batch culture in Cu-sufficient media and Cu-limited media. (b) and (c) As in (a) but for $\Delta nosZ$ deletion mutant PD2303, and $\Delta nosX$ deletion mutant PD2502, respectively. Cultures were grown in triplicate and bars represent SE.

Purification and characterization of affinity-tagged N_2OR from *P. denitrificans* strains

Plasmid pMSL002, which encodes NosZ (N_2OR) with a C-terminal Strep-tag II, was conjugated into wild-type (PD1222), $Pd\Delta nosZ$ (PD2303) and $Pd\Delta nosX$ (PD2502) strains using the *E. coli* pRK2013 helper strain. Strep-tagged N_2OR was overproduced and purified as previously described [21]. Briefly, this involved applying the soluble portion of cell lysates to a Hi-Trap HP Strep II affinity column (5 ml, GE Healthcare) and eluting with 20 mM HEPES, 150 mM NaCl and 2.5 mM desthiobiotin, pH 7.2, before exchanging into 20 mM HEPES, 150 mM NaCl, pH 7.2. Sample purity was confirmed using SDS-PAGE analysis and LC-MS. Protein concentrations were determined using the Bradford assay (BioRad) [25] and bovine serum albumin as a protein standard.

UV-visible absorbance spectra of N_2OR -Strep-tag II from different backgrounds were recorded on a Jasco V-550 spectrophotometer. Samples were made anaerobic by sparging with nitrogen gas for 5 min and oxidized or reduced with 5 mg ml⁻¹ stocks of potassium ferricyanide and sodium dithionite, respectively, in 20 mM HEPES, 150 mM NaCl, pH 7.5, by titrating concentration equivalents. Total copper content of the protein was determined using a colorimetric

bathocuproinedisulfonic acid (BCS) assay as previously described [21].

Activities of N_2OR -Strep-tag II isolated from different backgrounds were determined using an adapted methyl viologen assay [26, 27] in which samples were pre-incubated with a 500-fold excess of reduced methyl viologen for 150 min. Reaction was initiated by adding N_2O saturated buffer and the oxidation of blue (reduced) methyl viologen to its oxidized colourless form was followed at 600 nm as a function of time and data converted to specific activity using $\epsilon_{600\text{ nm}} = 13600\text{ M}^{-1}\text{ cm}^{-1}$ for the reduced methyl viologen cation radical [27].

RNA isolation, cDNA synthesis and qRT-PCR experiments

Expression of the *nosZ* gene was determined by qRT-PCR, using an AriaMx Real-Time PCR System G9930A (Agilent Technologies). The *nosX* mutant and PD1222 wild-type strains were cultivated under anoxic conditions as mentioned above for 12 h, reaching final $OD_{600\text{ nm}}$ of 0.6. Total RNA extraction, RNA quality and integrity assays, and RNA quantification were performed using the methodology previously described [17]. Briefly, 2 μg of total RNA were used for cDNA synthesis using RevertAid First Strand cDNA synthesis kit (Thermo Scientific) and random hexamers following the supplier's instructions. qRT-PCR reactions were run in triplicate in a total volume of 20 μl containing 10 μl of SensiFAST SYBR No-ROX Mix (Bioline), 0.7, 7 or 70 ng of cDNA and 2 μM of each primer. Melting curves were generated to verify the specificity of each amplification reaction. Expression of *nosZ* gene was determined using the oligonucleotide pair nosZ2F/nosZ2R [17] and normalized against the housekeeping gene *gapA* (glyceraldehyde-3-phosphate dehydrogenase; GAPDH1F/GAPDH1R [17]). The changes in gene expression were analysed accordingly to Pfaffl methodology [28]. The data presented correspond to the average of three independent biological replicates.

RESULTS

NosX is essential for whole-cell N_2O reduction in *P. denitrificans*

Wild-type *P. denitrificans* (PD1222), $\Delta nosZ$ (PD2303, missing the gene encoding N_2OR) and $\Delta nosX$ (PD2502, missing the gene *Pden_4214*) were grown in batch culture, in minimal medium, under Cu-sufficient and limited conditions. The wild-type culture produced a small amount of N_2O (~1 mM) in Cu-deficient conditions, but this was no longer detected as the culture moved into the stationary phase of growth Fig. 2. A N_2OR -negative phenotype (Nos⁻), in terms of growth and N_2O production, was observed in the $\Delta nosZ$ strain under both Cu regimes. For the $\Delta nosX$ strain, growth was affected both under Cu-sufficient and limited conditions, and N_2O levels were similar to those of the $\Delta nosZ$ strain, demonstrating the absence of a functioning enzyme.

The Nos⁻ phenotype of the $\Delta nosX$ strain was almost fully complemented under both Cu regimes by a plasmid-borne

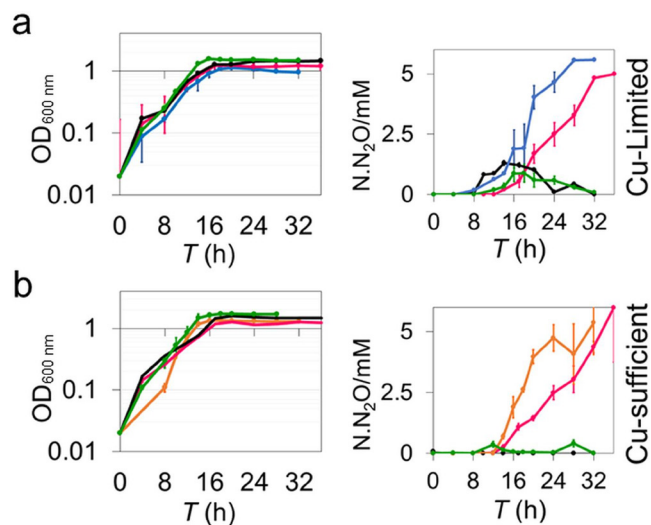


Fig. 3. Complementation of the *nosX* mutant. (a) Growth characteristics (optical density, OD_{600 nm}), left, and N₂O production (N₂O, mM N in the form of N₂O), right for the mutant Δ*nosX* PD2502 complemented under (a) Cu-limited, and (b) Cu-sufficient conditions in anaerobic batch culture. The pSPBN5 plasmid was conjugated into the Δ*nosX* PD2502 strain and cultured in the absence of taurine and in the presence of 1 mM taurine. For reference, the Δ*nosZ* PD2303 strain and wild-type PD1222 are shown. Experiments were repeated in triplicate and bars represent SE.

nosX gene copy (pSPBN5) expressed *in trans* from a taurine inducible promoter (Fig. 3), demonstrating that the Nos⁻ phenotype is associated with the absence of *nosX* and not a downstream effect of the deletion. The data demonstrate that the *nosX* deletion mutant strain of *P. denitrificans* is unable to catalyse N₂O reduction. This is in contrast to a previous study by Saunders and co-workers [22] involving a marked *nosX* deletion, where it was concluded that that NosX and NirX are functionally redundant, such that only one is required for N₂O reduction.

NosX is not involved in maturation of either Cu cofactor in N₂OR

Three possible explanations for the Nos⁻ phenotype in the Δ*nosX* mutant are apparent: the incomplete maturation/assembly of copper centres of N₂OR; the failure to activate N₂OR catalytic activity, for example through disruption of supply of electrons; or, the severe down-regulation of *nosZ* transcription. To investigate this further, a C-terminal strep II-tagged N₂OR was purified from the Δ*nosX* mutant strain and the properties of the N₂OR analysed with respect to the status of the Cu_A and Cu_Z centres.

Aerobically purified N₂OR, also known as the pink form of N₂OR, has been spectroscopically well characterized and all oxidized spectra were normalised to ε_{580 nm} 5000 M⁻¹ cm⁻¹ per monomer, as described by Rasmussen *et al.* [13]. Absorbance spectra of N₂OR enzymes isolated from cultures grown under Cu-sufficient conditions are shown in Fig. 4a. Spectra of N₂OR from wild-type cells and Δ*nosX* and Δ*nosZ* mutants

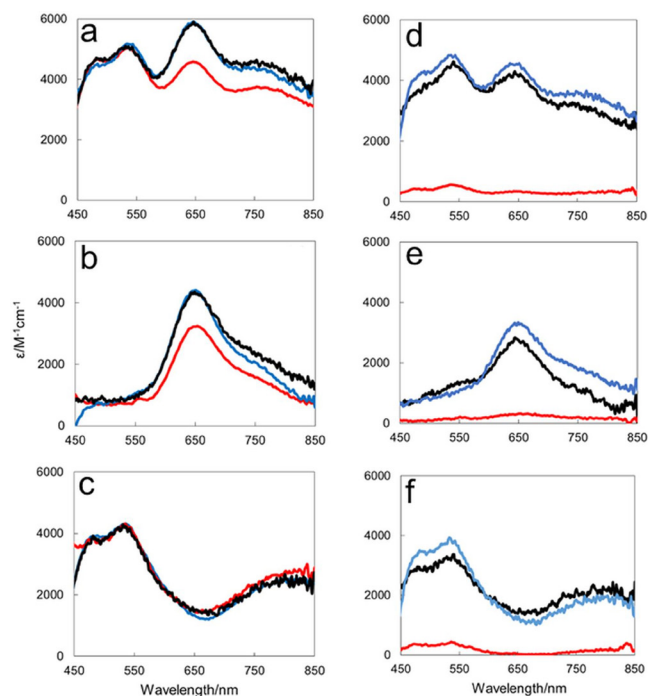


Fig. 4. UV-visible absorbance characterisation of strep-tagged N₂OR purified from different *P. denitrificans* backgrounds. Data are shown for N₂OR from wild-type PD1222, Δ*nosX* PD2502 and Δ*nosZ* PD2303 in 20 mM HEPES, 150 mM NaCl, pH 7.2. Spectra of ferricyanide-oxidized (a), sodium dithionite-reduced (b) and the oxidized minus reduced difference (c) are shown for enzymes isolated from cultures grown under Cu-sufficient conditions. Equivalent spectra (d–f), respectively, were measured for enzymes isolated from cultures grown under Cu-limited conditions.

have features at 480, 540 and 640 nm, in agreement with the previous literature on N₂OR from *P. denitrificans* [21], *P. pantotrophus* (PpN₂OR) [13], *Pseudomonas stutzeri* (PsN₂OR) [29], *Pseudomonas nautica* (PnN₂OR) [30], *Achromobacter cycloclastes* (AcN₂OR) [31] and *Marinobacter hydrocarbonoclasticus* (MhN₂OR) [32]. Features in the absorption spectrum at these wavelengths arise from S²⁻ to Cu(II) charge-transfer bands and additional optical bands due to interactions between the Cu(I) and Cu(II) ions of the centres [13]. Spectra of N₂OR isolated from wild-type cells have lower extinction coefficients than those from the mutant strains, suggesting that it contains lower levels of Cu cofactors.

The Cu content of all isolated N₂OR enzymes was determined (Table 1), confirming that enzymes isolated from Δ*nosZ* and Δ*nosX* mutants are replete with Cu, while that from wild-type cells contains slightly lower amounts, consistent with the absorption spectra. N₂OR activity was measured using a methyl viologen assay in which the reduced MV extinction coefficient, ε_{600 nm} = 13600 M⁻¹ cm⁻¹ [27], was used to quantify activity, and N₂OR was pre-incubated with a 500-fold excess reduced methyl viologen (MV) before initiating the reaction with N₂O (Table 1). Each N₂OR sample was active, with values for the enzyme from the wild-type and Δ*nosZ*

Table 1. Summary of some characteristics of strep-tagged N₂OR purified from *P. denitrificans* strains PD1222, PD2502 and PD2303

	Cu ions/monomer*		Specific activity† ($\mu\text{mol N}_2\text{O min}^{-1}$ mg^{-1})
	Cu-sufficient	Cu-limited	
Wild-type PD1222/ pMSL002 (StrepII tagged-NosZ)	5.6 \pm 0.1	0.4 \pm 0.27	171 \pm 13
Δ nosX/pMSL002	6.4 \pm 0.2	4.2 \pm 0.2	172 \pm 12
Δ nosZ/pMSL002	5.9 \pm 0.6	4.8 \pm 0.4	196 \pm 9

*Total copper per monomer was determined using the BCS Cu assay (see Methods).

†N₂O reductase activity was determined for enzymes isolated from cultures grown under Cu-sufficient conditions using a reduced methyl viologen assay ($\mu\text{mol N}_2\text{O min}^{-1}$ enzyme). Proteins were pre-incubated with a 500-fold excess reduced methyl viologen for 150 min prior to activity assay. All reactions were carried out in triplicate and SD is shown. ND. The data show that even though the Δ nosX strain has a Nos⁻ phenotype, N₂OR isolated from it is fully or close to fully active in an *in vitro* assay.

strains consistent with those previously reported [21, 26, 30]. Activity for N₂OR from the Δ nosX mutant was similar to that from wild-type, even though it contained significantly more Cu, suggesting that the enzyme from the Δ nosX mutant has a slightly lower activity.

Reduction of N₂OR samples with dithionite leads to reduction of the Cu_A centre to a [Cu^I:Cu^I] diamagnetic species, which is colourless and thus does not contribute in the visible region of the absorbance spectrum. Thus, in Fig. 4b, bands at 480, 540 and 900 nm are lost to leave a Cu_Z⁺ signature, consisting of a peak at 640 nm, in agreement with the literature for pink N₂OR [13]. The oxidized minus reduced difference spectrum, Fig. 4c, revealed the spectrum due to the Cu_A centre. The close similarity of spectral form and absorption extinction coefficients for N₂OR from Δ nosZ and Δ nosX mutants demonstrate that the assembly of the Cu cofactors of N₂OR is not affected by the *nosX* deletion when grown under Cu sufficiency [13].

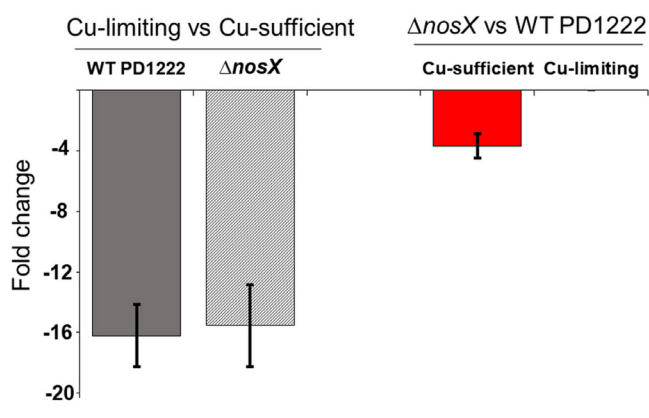


Fig. 5. *nosZ* relative expression determined by qRT-PCR. Left side shows *nosZ* expression change under Cu-limited versus Cu-sufficient conditions in the WT PD1222 and Δ nosX mutant strains. Right side shows *nosZ* expression change in Δ nosX mutant versus WT PD1222 strains under Cu-limiting and Cu-sufficient growth conditions.

An equivalent spectroscopic analysis of N₂OR enzymes isolated from cultures grown under Cu limitation (Fig. 4d–f) revealed spectra similar to those of Fig. 4a–c for enzymes from Δ nosZ and Δ nosX mutants, but with lower extinction coefficients, suggesting lower incorporation of Cu. Spectra for enzyme isolated from wild-type cultures, however, indicate very low levels of Cu incorporation. Determination of Cu content (Table 1) revealed that N₂OR from Δ nosZ and Δ nosX mutants contain ~4 Cu per N₂OR monomer, while that recovered from wild-type cells contains <1 Cu per monomer, consistent with absorbance data (Fig. 4d–f). As above, the close similarity between N₂OR enzymes isolated from Δ nosZ and Δ nosX mutants demonstrate that NosX does not play a role in assembly of the Cu cofactors of N₂OR under Cu-limited conditions.

NosX has a minor effect on transcription of *nosZ* under Cu-sufficient conditions

The data presented in Fig. 4, Table 1 revealed some variability in the extent to which Cu_Z centres are assembled in enzymes isolated from different strains and grown under different conditions; specifically, plasmid-encoded strep-tagged N₂OR isolated from wild-type cells contained fewer Cu_Z centres than that from the two mutants. Thus, the Δ nosX mutant behaves similarly to the Δ nosZ mutant, in which chromosomal *nosZ* is missing. This suggests that there may be fewer chromosomally encoded versions of N₂OR in the *nosX* mutant than in wild-type cells, as would be expected if *nosZ* expression is perturbed in the *nosX* mutant. Less chromosomally encoded N₂OR would provide less competition for Cu, leading to greater incorporation of Cu into the plasmid-encoded N₂OR.

To investigate this, qRT-PCR experiments were performed to determine the differential expression of *nosZ* in the *nosX* mutant compared to wild-type cells. Under Cu-sufficient conditions, a twofold decrease (1.9 \pm 0.2) in expression of *nosZ* was measured in Δ nosX compared to wild-type cells (Fig. 5). This likely contributes to the observed increased incorporation of Cu into strep-tagged N₂OR isolated from the Δ nosX mutant compared to that from wild-type cells. However, no significant difference in expression of *nosZ* was detected between Δ nosX and wild-type grown under Cu-limiting conditions. In both cases, the *nosZ* expression in Cu-limiting conditions was ~15-fold lower than that under Cu-sufficient conditions (Fig. 5), consistent with previous report on the effect of Cu on *nosZ* expression in wild-type cells [17]. Thus, effects on *nosZ* expression do not account for the very low incorporation of Cu into strep-tagged N₂OR in wild-type cells compared to in the Δ nosX mutant.

DISCUSSION

The *nosX* gene is conserved across the NGC of α - and β -proteobacteria, but not among γ - or clade-II members of N₂O-reducing bacteria. Here, we have demonstrated a Nos⁻ phenotype for a *nosX* deletion mutant in *P. denitrificans* (PD2502), which was complemented *in trans* using a functional *nosX* plasmid-borne gene copy under taurine

inducible control. NosX is a member of the AbpE protein family, which bind flavin adenine dinucleotide [33, 34]. Some AbpE proteins are flavinyl transferases, functioning in the post-translational maturation of another flavin-requiring protein. For example, *Vibrio cholera* AbpE transfers a flavin mononucleotide (FMN) to a threonine residue in NqrC [35]. In *P. denitrificans* there are three *abpE* homologues: *nosX*, encoded by *pden_4214*, *nirX* (*pden_2485*) and *pden_3291*. NosX and NirX are exported to the periplasm via the Tat pathway while Pden_3291 is predicted to be cytoplasmic.

An earlier study of an antibiotic cassette insertion mutation in the *P. denitrificans nosX* gene reported no effect on N₂OR activity [22]. This led to the proposal that the *nirX* gene in *P. denitrificans* is a functional homologue of *nosX*, such that mutation of both genes are required in order to observe a Nos⁻ phenotype. This previous conclusion is clearly at odds with the data presented here. One possibly important observation is that the previous mutagenesis study did not involve full *nosX* deletion. Conserved residues within the putative FAD binding pocket in NosX are now known, including Ser68, Tyr70, Thr174 and Gly256, based on sequence similarities with the SeAbpE (Fig. S1) [34]. If these residues are important for NosX function, then the previous mutation strategy for *P. denitrificans nosX*, in which a kanamycin insertion was made 469 bp into the gene, would not have disrupted the conserved Ser68 and Tyr70 residues. The resulting truncated NosX may have retained some function, which would account for why a clear phenotype was not observed in the single *nosX* insertional mutant. We note that the requirement for *nosX* in N₂O reduction has also been demonstrated in *Sinorhizobium meliloti*. In that case, a Tn5-mediated mutation 31 nucleotides into the total 966 nucleotide sequence downstream of *nosDFYL*, a region now recognized as *nosX*, abolished N₂OR activity [36].

In the earlier report of a double *nosXnirX* mutant of *P. denitrificans*, it was reported that the N₂OR present in unfractionated periplasm from this mutant was deficient in the Cu_A centre, leading to the conclusion that NosX and NirX play a role in assembly of this cofactor [22]. However, subsequent studies of anaerobically purified N₂OR from the double *nirXnosX* mutant and a single *nirX* mutant indicated that the absence of NosX resulted in N₂OR with both Cu cofactors assembled, but with Cu_Z exhibiting a spectroscopically distinct form, termed pink Cu_Z^{*}, that is normally only observed upon reaction with O₂ [37]. This Cu_Z form is not catalytically active, but is proposed to represent a catalytically relevant intermediate oxidation state of the Cu_Z centre ([4CuS]³⁺), which binds N₂O and proceeds through a state denoted as Cu_Z⁰ [38].

Here, to determine the effect of the absence of *nosX*/NosX alone on N₂OR, we utilized a previously reported plasmid-encoded Strep-tagged N₂OR that can be readily isolated from different background strains and characterized in terms of its Cu cofactor content and spectroscopic properties. These experiments demonstrated unequivocally that the assembly of the Cu_A and Cu_Z centres was unaffected in the absence of *nosX*. Thus, the phenotype exhibited by the mutant does not

arise because of a deficiency in the insertion of Cu into N₂OR. We note that the spectroscopic properties of N₂OR from the Δ *nosX* mutant strain are the same as those of the Cu_Z^{*} centre from purified from the *nosXnirX* mutant. This may suggest the Cu_Z centre was purified in a catalytically inactive redox state. However, the pink form reported in this work was generated by aerobic purification, with *nirX* remaining in the genome and under conditions where we expect to observe the Cu_Z centre is this pink Cu_Z^{*} form, as demonstrated by the control experiments with N₂OR isolated from the wild-type strain.

AbpE from the N₂O-reducing bacterium *P. stutzeri* is a monomeric FAD-binding protein [23]. In the absence of *nosX* in the NGC of *P. stutzeri*, AbpE functions as a flavin donor, catalysing the covalent flavinylation of a threonine residue of NosR [23]. Importantly, the post-translationally modified, FMN-bound NosR is proposed to be the electron donor to N₂OR, such that in the absence of NosR N₂OR is not functional. Our data indicate that N₂OR Cu cofactor maturation is unaffected by the loss of NosX, and we conclude that in *P. denitrificans* it most likely functions as the main system-specific maturation factor for NosR, and thus as an indirect activator of N₂OR. If this is the case, then a Nos⁻ phenotype would be expected for a Δ *nosR* strain. This was recently demonstrated: a *P. denitrificans* Δ *nosR* strain exhibited a vastly decreased capacity to reduce N₂O, irrespective of the levels of Cu in the cell [17]. However, we note that the Δ *nosR* strain did retain some ability to reduce N₂O, whereas the *nosX* mutant investigated here did not, and so the *nosX* phenotype is actually more dramatic than the *nosR* phenotype. Why this is the case is not clear. One possibility is that NosX does not only mature NosR, such that in the absence of NosX, there is a further effect on NosZ activity. Alternatively, having a non-flavinylated NosR present might somehow inhibit NosZ more severely than having no NosR present at all. We also note that the previously reported transcription data revealed the loss of Cu-responsive transcription of *nosZ* in the *nosR* deletion strain [17], suggesting that NosR itself may be multifunctional, or that its absence leads to pleiotropic effects, some of which may be indirect. Clearly, further studies are needed to investigate directly the role of NosX in NosR maturation, and more generally other possible roles of NosX and the function(s) of NosR.

An intriguing observation reported here is the lower levels of Cu cofactor incorporation observed under Cu-limited conditions for the Strep-tagged N₂OR from wild-type cells compared to that from the *nosZ* and *nosX* mutants. One possibility that we examined was that *nosX*/NosX is involved in the regulation of *nosZ*, such that in the absence of *nosX*/NosX, lower amounts of chromosomally encoded N₂OR were present, perhaps leading to less competition for copper and higher incorporation of Cu into the plasmid-encoded Strep-tagged form. While the absence of *nosX* did result in a twofold reduction of *nosZ* expression under Cu-sufficient conditions, no significant difference between the wild-type and *nosX* mutant strains was detected under Cu-limited conditions where the incorporation of Cu was most pronounced. The very low expression of the chromosomal *nosZ* gene under Cu-limited conditions suggests that a simple competition

between chromosomal- and plasmid-encoded N₂O_R enzymes for Cu is unlikely. A further possibility is that the presence of the Strep-tag required for rapid recovery and biochemical analysis of NosZ results in modest perturbation of Cu cofactor assembly factor interactions such that the wild-type enzyme is a preferred substrate, an effect that only becomes apparent under very low Cu conditions. Clearly, further studies are needed to explore this possibility.

In summary, the data presented here show that *nosX* is essential for whole-cell N₂O reduction in the α -proteobacterium *P. denitrificans*, and that the *nosX* and *nirX* gene products are not functionally redundant under our experimental conditions. The function of NosX is not associated with the assembly of the Cu cofactors of N₂O_R. Instead, based on homology between NosX and ApbE proteins, and the recent demonstration of an essential role for an ApbE family flavin transferase in the maturation of NosR in *P. stutzeri*, it is likely that NosX is involved in indirectly maintaining the reaction cycle of N₂O_R through the flavinylation of another accessory protein, NosR.

Funding information

This work was supported by Biotechnology and Biological Sciences Research Council through the award of a DTP PhD studentship to S.P.B., grants BB/L022796/1, BB/M00256X/1 and BB/S008942/1. M.J.S.-L. was funded by the Marie Skłodowska Curie Initial Training Network, Nitrous Oxide Research Alliance (NORA), grant 316472, under EU's seventh framework programme.

Conflicts of interest

The authors declare that there are no conflicts of interest.

References

- Ravishankara AR, Daniel JS, Portmann RW. Nitrous oxide N₂O: the dominant ozone-depleting substance emitted in the 21st century. *Science* 2009;326:123–125.
- Fowler D, Coyle M, Skiba U, Sutton MA, Cape JN et al. The global nitrogen cycle in the twenty-first century. *Philos Trans R Soc Lond B Biol Sci* 2013;368:20130164.
- Thomson AJ, Giannopoulos G, Pretty J, Baggs EM, Richardson DJ. Biological sources and sinks of nitrous oxide and strategies to mitigate emissions. *Philos Trans R Soc Lond B Biol Sci* 2012;367:1157–1168.
- Hu H-W, Chen D, He J-Z, HW H, JZ H. Microbial regulation of terrestrial nitrous oxide formation: understanding the biological pathways for prediction of emission rates. *FEMS Microbiol Rev* 2015;39:729–749.
- Hallin S, Philippot L, Löffler FE, Sanford RA, Jones CM. Genomics and ecology of Novel N₂O-reducing microorganisms. *Trends Microbiol* 2018;26:43–55.
- Jones CM, Graf DRH, Bru D, Philippot L, Hallin S. The unaccounted yet abundant nitrous oxide-reducing microbial community: a potential nitrous oxide sink. *ISME J* 2013;7:417–426.
- Stein LY. Insights into the physiology of ammonia-oxidizing microorganisms. *Curr Opin Chem Biol* 2019;49:9–15.
- Domeignoz-Horta LA, Philippot L, Peyrard C, Bru D, Breuil M-C et al. Peaks of in situ N₂O emissions are influenced by N₂O-producing and reducing microbial communities across arable soils. *Glob Chang Biol* 2018;24:360–370.
- Griffis TJ, Chen Z, Baker JM, Wood JD, Millet DB et al. Nitrous oxide emissions are enhanced in a warmer and wetter world. *Proc Natl Acad Sci U S A* 2017;114:12081–12085.
- Liu B, Frostegård Åsa, Bakken LR. Impaired reduction of N₂O to N₂ in acid soils is due to a posttranscriptional interference with the expression of *nosZ*. *mBio* 2014;5:e01383–01314.
- Brown K, Djinic-Carugo K, Haltia T, Cabrito I, Saraste M et al. Revisiting the catalytic Cu₂ cluster of nitrous oxide (N₂O) reductase. Evidence of a bridging inorganic sulfur. *J Biol Chem* 2000;275:41133–41136.
- Pomowski A, Zumft WG, Kroneck PMH, Einsle O. N₂O binding at a [4Cu:2S] copper-sulphur cluster in nitrous oxide reductase. *Nature* 2011;477:234–237.
- Rasmussen T, Berks BC, Butt JN, Thomson AJ. Multiple forms of the catalytic centre, Cu₂, in the enzyme nitrous oxide reductase from *Paracoccus pantotrophus*. *Biochem J* 2002;364:807–815.
- Mattatall NR, Jazairi J, Hill BC. Characterization of YpmQ, an accessory protein required for the expression of cytochrome c oxidase in *Bacillus subtilis*. *J Biol Chem* 2000;275:28802–28809.
- Heikkilä MP, Honisch U, Wunsch P, Zumft WG. Role of the Tat transport system in nitrous oxide reductase translocation and cytochrome *cd*₁ biosynthesis in *Pseudomonas stutzeri*. *J Bacteriol* 2001;183:1663–1671.
- Simon J, Einsle O, Kroneck PMH, Zumft WG. The unprecedented *nos* gene cluster of *Wolinella succinogenes* encodes a novel respiratory electron transfer pathway to cytochrome *c* nitrous oxide reductase. *FEBS Lett* 2004;569:7–12.
- Sullivan MJ, Gates AJ, Appia-Ayme C, Rowley G, Richardson DJ. Copper control of bacterial nitrous oxide emission and its impact on vitamin B₁₂-dependent metabolism. *Proc Natl Acad Sci U S A* 2013;110:19926–19931.
- Wunsch P, Zumft WG. Functional domains of NosR, a novel transmembrane iron-sulfur flavoprotein necessary for nitrous oxide respiration. *J Bacteriol* 2005;187:1992–2001.
- Riester J, Zumft WG, Kroneck PM. Nitrous oxide reductase from *Pseudomonas stutzeri*. Redox properties and spectroscopic characterization of different forms of the multicopper enzyme. *Eur J Biochem* 1989;178:751–762.
- Zumft WG. Biogenesis of the bacterial respiratory Cu_A Cu-S enzyme nitrous oxide reductase. *J Mol Microbiol Biotech* 2005;10:154–166.
- Bennett SP, Soriano-Laguna MJ, Bradley JM, Svistunenko DA, Richardson DJ et al. NosL is a dedicated copper chaperone for assembly of the Cu₂ center of nitrous oxide reductase. *Chem Sci* 2019;10:4985–4993.
- Saunders NF, Hornberg JJ, Reijnders WN, Westerhoff HV, de Vries S et al. The NosX and NirX proteins of *Paracoccus denitrificans* are functional homologues: their role in maturation of nitrous oxide reductase. *J Bacteriol* 2000;182:5211–5217.
- Zhang L, Trncik C, Andrade SLA, Einsle O. The flavinyl transferase ApbE of *Pseudomonas stutzeri* matures the NosR protein required for nitrous oxide reduction. *Biochim Biophys Acta Bioenerg* 2017;1858:95–102.
- Vishniac W, Santer M. The thiobacilli. *Bacteriol Rev* 1957;21:195–213.
- Bradford MM. A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. *Anal Biochem* 1976;72:248–254.
- Ghosh S, Gorelsky SI, Chen P, Cabrito I, Moura JGG et al. Activation of N₂O reduction by the fully reduced μ_4 -sulfide bridged tetranuclear Cu₂ cluster in nitrous oxide reductase. *J Am Chem Soc* 2003;125:15708–15709.
- Kristjansson JK, Hollocher TC. First practical assay for soluble nitrous oxide reductase of denitrifying bacteria and a partial kinetic characterization. *J Biol Chem* 1980;255:704–707.
- Pfaffl MW. A new mathematical model for relative quantification in real-time RT-PCR. *Nucleic Acids Res* 2001;29:e45–.
- Coyle CL, Zumft WG, Kroneck PM, Körner H, Jakob W. Nitrous oxide reductase from denitrifying *Pseudomonas perfectomarina*. Purification and properties of a novel multicopper enzyme. *Eur J Biochem* 1985;153:459–467.
- Prudêncio M, Pereira AS, Tavares P, Besson S, Cabrito I et al. Purification, characterization, and preliminary crystallographic study

- of copper-containing nitrous oxide reductase from *Pseudomonas nautica* 617. *Biochemistry* 2000;39:3899–3907.
31. Paraskevopoulos K, Antonyuk SV, Sawers RG, Eady RR, Hasnain SS. Insight into catalysis of nitrous oxide reductase from high-resolution structures of resting and inhibitor-bound enzyme from *Achromobacter cycloclastes*. *J Mol Biol* 2006;362:55–65.
 32. Dell'Acqua S, Pauleta SR, Moura JJG, Moura I. Biochemical characterization of the purple form of *Marinobacter hydrocarbonoclasticus* nitrous oxide reductase. *Philos Trans R Soc Lond B Biol Sci* 2012;367:1204–1212.
 33. Bertsova YV, Fadeeva MS, Kostyrko VA, Serebryakova MV, Baykov AA *et al.* Alternative pyrimidine biosynthesis protein ApbE is a flavin transferase catalyzing covalent attachment of FMN to a threonine residue in bacterial flavoproteins. *J Biol Chem* 2013;288:14276–14286.
 34. Boyd JM, Endrizzi JA, Hamilton TL, Christopherson MR, Mulder DW *et al.* FAD binding by ApbE protein from *Salmonella enterica*: a new class of FAD-binding proteins. *J Bacteriol* 2011;193:887–895.
 35. Hayashi M, Nakayama Y, Yasui M, Maeda M, Furuishi K *et al.* FMN is covalently attached to a threonine residue in the NqrB and NqrC subunits of Na(+)-translocating NADH-quinone reductase from *Vibrio alginolyticus*. *FEBS Lett* 2001;488:5–8.
 36. Chan YK, McCormick WA, Watson RJ. A new *nos* gene downstream from *nosDFY* is essential for dissimilatory reduction of nitrous oxide by *Rhizobium (Sinorhizobium) meliloti*. *Microbiology* 1997;143:2817–2824.
 37. Wunsch P, Körner H, Neese F, van Spanning RJM, Kroneck PMH *et al.* NosX function connects to nitrous oxide (N₂O) reduction by affecting the Cu₂ center of NosZ and its activity *in vivo*. *FEBS Lett* 2005;579:4605–4609.
 38. Carreira C, Pauleta SR, Moura I. The catalytic cycle of nitrous oxide reductase - The enzyme that catalyzes the last step of denitrification. *J Inorg Biochem* 2017;177:423–434.

Edited by: I. Martin-Verstraete and J. A Galnick

Five reasons to publish your next article with a Microbiology Society journal

1. The Microbiology Society is a not-for-profit organization.
2. We offer fast and rigorous peer review – average time to first decision is 4–6 weeks.
3. Our journals have a global readership with subscriptions held in research institutions around the world.
4. 80% of our authors rate our submission process as 'excellent' or 'very good'.
5. Your article will be published on an interactive journal platform with advanced metrics.

Find out more and submit your article at microbiologyresearch.org.