

The *nirSTBM* region coding for cytochrome cd_1 -dependent nitrite respiration of *Pseudomonas stutzeri* consists of a cluster of mono-, di-, and tetraheme proteins

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Genes for respiratory nitrite reduction (denitrification) of *Pseudomonas stutzeri* are clustered within 7 kbp. A 4.6-kbp *Hind* III-*Kpn* I fragment carrying *nirS*, the structural gene for cytochrome cd_1 , was sequenced. An open reading frame immediately downstream of *nirS* codes for a 22.8-kDa protein with four heme c -binding motifs. Mutagenesis of this gene causes an apparent defect in electron donation to cytochrome cd_1 . Following this ORF are the structural genes for cytochrome c_{551} , cytochrome c_{553} , and ORFs that codes for a 11.9-kDa monoheme protein. All cytochromes have a signal sequence for protein export.

Nitrite respiration; Cytochrome cd_1 ; Cytochrome c_{551} ; Cytochrome c_{553} ; Tetraheme protein; Signal peptide; *Pseudomonas stutzeri*

1. INTRODUCTION

Pseudomonas stutzeri [1] and *Pseudomonas aeruginosa* [2] synthesize cytochrome cd_1 as respiratory nitrite reductase (ferrocytochrome c_{551} :oxygen oxidoreductase; EC 1.9.3.2) for their denitrification system [3,4]. The structural gene, *nirS*, for cytochrome cd_1 of *P. stutzeri* is vicinal to genes for nitrous oxide respiration, which are part of the overall denitrification process [5]. The primary structure of *nirS* from *P. aeruginosa* was recently elucidated [6]. Contiguous to *nirS* is the gene coding for cytochrome c_{551} [7,8], the *in vitro* electron donor for cytochrome cd_1 (in certain systems interchangeable with azurin). Here we report the substantially extended nucleotide sequence of the *nirS* region from *P. stutzeri* with the primary structure of five heme proteins and their respective precursor forms, and establish a gene order different from that of *P. aeruginosa*.

2. MATERIALS AND METHODS

The *nirS* region of *P. stutzeri* strain ZoBell (formerly *P. perfringens* ATCC 14405) was cloned from a lambda gtl1 expression library [5]. Generation of subclones from cosmids, DNA sequencing,

and other details of recombinant DNA work are described elsewhere [5,9,10]. Cytochrome cd_1 , cytochrome c_{551} and cytochrome c_{553} were purified from anaerobically, nitrate-grown cells [4]. The conditions of Iso-Dalt electrophoresis were as in [11]. Iron was determined by atomic absorption spectroscopy. Molecular masses were estimated by gel filtration, by pore-gradient electrophoresis and by SDS-PAGE. Manual and automated Edman degradations were used to determine the N-terminal sequences of Cm-cytochrome cd_1 and peptides [12].

3. RESULTS

3.1. Gene map, sequencing strategy, open reading frames

Fig. 1 shows the restriction map of the 4.6-kbp *Hind* III-*Kpn* I fragment and the sequencing strategy. Both directions were sequenced by the dideoxy-chain termination method; GC-rich regions were verified with dITP. The sequenced DNA region revealed five ORFs in the same transcriptional direction (Fig. 1), which showed a codon preference typical of *Pseudomonas* genes. Three ORFs could be assigned to known cytochromes.

3.2. Sequence of the *nirS* region

The nucleotide sequence and the derived gene products of the *nirSTBM* region are shown in Fig. 2. The first ORF codes for the structural gene of cytochrome cd_1 . The *nirS* gene product carries a presequence of 26 amino acids, 14 of which are positionally identical to those of the *P. aeruginosa* nitrite reductase. Overall, the two proteins have 56.4% identity and 13.6% conser-

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Abbreviations: ORF(s), open reading frame(s); PAGE, polyacrylamide gel electrophoresis; SDS, sodium dodecyl sulfate

vative replacements when aligned by the algorithm of [13]. Fig. 3A shows an alignment of the N-terminal regions which have the greatest divergence due to insertions adjacent to the heme *c*-binding site without equivalent in the other protein; the binding site itself, however, is highly conserved. Of 14 histidine residues in cytochrome *cd₁* II are invariant. Histidine (besides methionine) coordinates heme *c* and is a likely candidate also for the axial ligand of the *d₁* heme. His-167 resides within the most conserved neighbourhood; other conserved regions are around the histidine residues 261, 311, 353, and 362.

The *nirS* gene is followed by ORF2, assigned to the *nirT* gene whose mutagenesis leads to a Nir⁻ phenotype [4]. Its derived gene product carries four heme *c*-binding motifs. These motifs show an apparent pairwise arrangement (Fig. 2), yet there is no conspicuous sequence similarity between the N-terminal and C-terminal domains. A screening of the MIPSX data bank (Martinsried Institute for Protein Sequence Data, release 17) by the FASTA program [14] did not reveal significant matches with other multiheme cytochromes including that of the photosynthetic reaction center. In all four motifs the cysteines are spaced by two residues and not by four as found in the *c₁*-type cytochromes of the sulfate-reducing bacteria [15]. The motif starting with Cys-88 is unusual in having a proline residue [16]. No heme-binding motif with proline was found in the cytochrome entries of the SWISS-PROT Protein Sequence Databank (release 15). A *c*-type cytochrome with four heme groups within a single polypeptide chain has not been identified so far in any pseudomonad. The *nirT* product has a presumed leader sequence with a predicted cleavage site at Gly-28 [17] and a hydrophobic, potential membrane anchor of 17 amino acids directly following the N-terminus.

Immediately downstream of *nirT* the structural gene for the diheme cytochrome *c₅₅₂*, termed *nirB*, was found (Fig. 2). A comparison of the N-terminus of the purified protein [18] and the *nirB* gene product revealed a presequence of 23 amino acids (Table I). One of the two heme-binding sites of cytochrome *c₅₅₂* is unusual with tryptophan as a third residue spacing the thioether-forming cysteines. The derived amino acid sequence confirms this unique heme-binding motif of the cytochrome family and extends the C-terminus 21 residues beyond the previously reported, incomplete sequence of this protein [18]. The two uncertain amino acids Ser-245 and Asn-247 of that sequence were identified as arginine and histidine; for Asp-36 and Asn-206 the derived NirB sequence indicates alanine and aspartate, respectively.

The fourth ORF in row, assigned to *nirM*, codes for cytochrome *c₅₅₁*. The amino acid sequence of this cytochrome from *P. stutzeri* strain 221 [19] shows 11 replacements (5 of which are of conservative nature) versus the strain ZoBell used in this work (Fig. 2).

Cytochrome *c₅₅₁* carries a signal peptide of 22 amino acids (Table I). The *nirM* gene is immediately followed by ORF3. This ORF is likely to code for a further monoheme *c*-type cytochrome. Its derived gene product is a hydrophilic protein with a canonical heme-binding motif and a predicted export signal sequence.

3.3. Properties of derived gene products

Properties of the derived gene products are summarized in Table I. The molecular mass of cytochrome *cd₁* determined from iron analysis, SDS-PAGE, non-denaturing pore-gradient PAGE, and gel filtration, gave values of 35.3, 61.2, 135, and 119 kDa, respectively. Like the cytochrome *cd₁* from other denitrifying bacteria the *P. stutzeri* protein is in solution a dimer, carrying pairs of heme *c* and *d₁* groups. Its experimentally determined isoelectric point is 6.5 ± 0.25 ($n = 3$). Cytochrome *cd₁* (see also [20]), cytochrome *c₅₅₁* and cytochrome *c₅₅₂* were localized in the periplasm by two-dimensional electrophoretic analysis of this compartment. Purified cytochrome *c₅₅₁* and cytochrome *c₅₅₂* separated in isoelectric focusing into several charge isomers. Their positions were identified on the gels by comparison with the purified components, and for cytochrome *cd₁* also by immunoblotting (C. Siegel, B. Goos and W.G. Zumft, unpublished). In agreement with the periplasmic location of the three cytochromes is the presence of signal peptides (Table I). Mature cytochrome *c₅₅₂* has a calculated M_r of 29 413 Da including the two heme groups. A value of 25 800 Da was determined from sedimentation equilibrium [21]. The same method indicated 7660 Da for cytochrome *c₅₅₁* [22] versus 9180 Da from the composition.

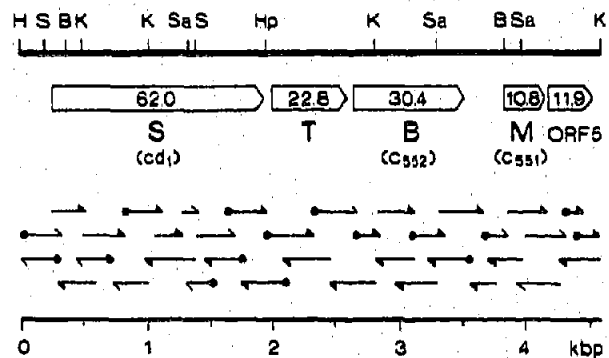


Fig. 1. Gene map, open reading frames, and sequencing strategy of the *nirS* region from *P. stutzeri*. Open arrows show the location and direction of transcription of the genes for *nirS* (cytochrome *cd₁*), *nirT* (tetraheme protein), *nirB* (cytochrome *c₅₅₂*), *nirM* (cytochrome *c₅₅₁*), and ORF3 (monoheme protein). Numbers indicate molecular masses of the precursor forms. Small arrows show the sequenced part of subclones. Arrows beginning with a dot identify clones sequenced with synthetic, others with M13 universal primers; filled-in arrowheads denote clones sequenced also with dITP. Physically mapped restriction sites are B, *Bgl* II; H, *Hind* III; Hp, *Hpa* I; K, *Kpn* I; S, *Sma* I; and Sa, *Sal* I.

Fig. 2. Nucleotide and derived amino acid sequence of the *nirSTBM* genes and ORFs of *P. stutzeri* ZoBell. Potential ribosome-binding sites (SD) are overlined. Stop codons are termed *end*. Inverted repeats are shown by opposing arrows. Underlined amino acids of the *nirS* product were determined by peptide sequencing. Asterisks denote identical, dots similar residues in comparison with the NirS protein from *P. aeruginosa* [6]; gaps for alignment are not shown. The triangle at nucleotide position 2474 of *nirT* indicates the Tn5 insertion site of mutant MK201, which was determined by sequencing. The amino acids which differ in cytochrome *c₅₅₁* from *P. stutzeri* strain 221 [19] are shown as subscripts in the *nirM* sequence. Boxed residues indicate heme *c*-binding motifs. The N-terminal amino acid of a processed protein is circled. The nucleotide sequence data will appear in the EMBL, GenBank and DDBJ Nucleotide Sequence Databases under accession no. X56813.

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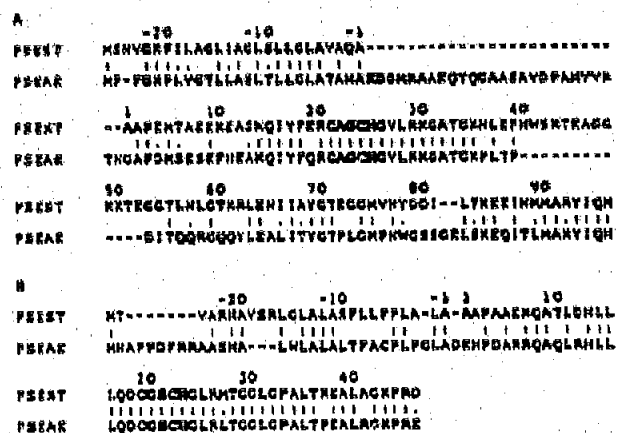


Fig. 3. Alignment of the N-terminal domains of (A) cytochrome *cd*₁, and (B) ORF5 from *P. stutzeri* (PSEST) and *P. aeruginosa* (PSEAE). Identical amino acids are shown by colons, conservative replacements by dots; heme c-binding sites and N-termini are printed in bold-face. Negative numbers label the signal sequence. Data for *P. aeruginosa* cytochrome *cd*₁ and ORF5 are from ref. 6 and 8, respectively.

4. DISCUSSION

The genes for nitrite respiration, *nir*, and the *nos* genes required for nitrous oxide respiration of *P. stutzeri* are closely linked. Of the former genes, *nirD* (presumably involved in heme *d*₁ synthesis or processing) and *nirS* have been previously identified. The two genes are separated by about 4 kbp [5]. Sequencing the *nirS* region towards *nirD* has revealed five ORFs all coding for heme proteins. The gene order of *P. stutzeri* with respect to cytochrome *cd*₁ and cytochrome *c₅₅₁* (cf. Fig. 1) is unexpectedly different from that found in *P. aeruginosa*, given the systematic relation between these bacterial species as members of the tight rRNA homology group I. The *nirS* gene of *P. aeruginosa* is followed by *nirM* [7,8] and ORF5. Arai et al. [8] reported 232 nontranslated nucleotides beyond the 3'-end of *nirM*. We found that this sequence harbors an

ORF that overlaps four nucleotides with the preceding *nirM* gene. A potential ribosome-binding site is located -12 nucleotides upstream of the putative start codon. Translation of this region revealed that it corresponds to the ORF5 gene product of *P. stutzeri* showing 62.8% identity (Fig. 3B).

In *P. stutzeri* the genes *nirS* and *nirM*-ORF5 are separated by intercalation of the genes *nirT* and *nirB*. The *nirT* gene product of *P. stutzeri* has features of a novel tetraheme protein. Mutagenesis of this gene with the transposon Tn5 leads to a Nir⁻ phenotype (mutant strain MK201), yet catalytically active cytochrome *cd*₁ is still synthesized [4]. This is indicative of a defect in electron donation. As a result of the Tn5 insertion, mutant MK201 also overproduces cytochrome *c₅₅₂*. The hydrophobic part at the mature N-terminus of the NirT protein might anchor it in the membrane, and allow electron flow between the respiratory chain and a periplasmic acceptor. Electrons could pass directly from NirT to the reductase or, as deduced from in vitro evidence, via cytochrome *c₅₅₁*. Given the proximity of *nirT* to *nirS*, it will depend on the transcriptional control exerted by the inverted repeat downstream of *nirS* with a free energy of formation of -126 kJ/mol [23], whether readthrough is possible and *nirT* expression will be regulated like *nirS*. Anaerobic regulation is indicated from the organizational feature that the *nirT* gene forms probably a transcriptional unit with the immediately following *nirB* gene. The *nirB* product (cytochrome *c₅₅₂*) is not found in aerobically grown cells [22].

Cytochrome *c₅₅₂* shows peroxidase activity on proteolytic modification [18]. The function of this component in vivo and of its acquired peroxidase activity is unknown. Like cytochrome *cd*₁ the synthesis of cytochrome *c₅₅₂* depends on anaerobic culture conditions and the concentration of this component increases with rising cellular nitrite reductase activity [22,24]. A role in nitrite reduction has been suggested. The location of the *nirB* gene next to a gene involved in electron

Table I
Properties of the derived products of the *nirS*-*nirB* gene cluster and ORF5 of *P. stutzeri* strain ZoBell

Gene product	Residue number ^a	M _r ^{a,b}	Presequence ^c	pI ^{a,b}	Heme-c binding motifs
NirB	268	28 180	MKKTLMASAVGAV	4.93	CAGCH, CWGSCH
			IAFOTHGAMA-A		
NirM	82	8 563	MKKLIPMLALG	5.46	CAACH
			GALAMQPALA-Q		
NirS	534	39 532	MSNVGKPLAGLIA	6.43	CAGCH
			GLSLLGLAVAQQA-A		
NirT	173	19 738	MTDKDGKQQKGIL	6.32	CISCH, CPDCH
			ALLRRPSTRYSLG-G ^d		CRNCH, CIACH
ORF5	87	9 202	MTVARHVSRLGLA	5.79	CGSCH
			LASFLLFPLALA-A ^e		

^aValue for mature protein ^bM_r value without heme groups ^cCleavage site indicated by a dash ^dCalculated value ^ePresequence not confirmed by N-terminal sequencing

donation to cytochrome c_{bb} , and perhaps within the same operon as *nirS*, adds further circumstantial evidence to this implication. Transposon mutants MK202 and MK206/207 which had lost cytochrome c_{bb} or had only a low content also had a reduced cytochrome c_{bb} level [4].

Unlike cytochrome c_{bb} , cytochrome c_{aa} is not regulated anaerobically [22,23], questioning an exclusive role in nitrite reduction. The intergenic region of 320 nucleotides between *nirB* and *nirM* has multiple inverted repeats (Fig. 2) with the potential to form various transcriptional termination, attenuation or other regulatory structures. We assume from this that the genes for cytochrome c_{bb} and cytochrome c_{aa} of *P. stutzeri* will belong to separate operons.

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