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Published in: Molecular Microbiology

DOI: 10.1111/j.1365-2958.1996.tb02644.x

IMPORTANT NOTE: You are advised to consult the publisher's version (publisher's PDF) if you wish to cite from it. Please check the document version below.

Document Version Publisher's PDF, also known as Version of record

Publication date: 1996

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):

Gier, J-W. L. D., Schepper, M., Reijnders, W. N. M., Dyck, S. J. V., Slotboom, D. J., Warne, A., ... Oost, J. V. D. (1996). Structural and functional analysis of aa3-type and cbb3-type cytochrome c oxidases of Paracoccus denitrificans reveals significant differences in proton-pump design. Molecular Microbiology, 20(6), 1247-1260. DOI: 10.1111/j.1365-2958.1996.tb02644.x

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# Structural and functional analysis of *aa*<sub>3</sub>-type and *cbb*<sub>3</sub>-type cytochrome *c* oxidases of *Paracoccus denitrificans* reveals significant differences in proton-pump design

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#### Summary

In Paracoccus denitrificans the aa<sub>3</sub>-type cytochrome c oxidase and the bb<sub>3</sub>-type quinol oxidase have previously been characterized in detail, both biochemically and genetically. Here we report on the isolation of a genomic locus that harbours the gene cluster ccoNOQP, and demonstrate that it encodes an alternative *cbb*<sub>3</sub>-type cytochrome *c* oxidase. This oxidase has previously been shown to be specifically induced at low oxygen tensions, suggesting that its expression is controlled by an oxygen-sensing mechanism. This view is corroborated by the observation that the ccoNOQP gene cluster is preceded by a gene that encodes an FNR homologue and that its promoter region contains an FNR-binding motif. Biochemical and physiological analyses of a set of oxidase mutants revealed that, at least under the conditions tested, cytochromes aa<sub>3</sub>, bb<sub>3</sub> and cbb<sub>3</sub> make up the complete set of terminal oxidases in P. denitrificans. Protontranslocation measurements of these oxidase mutants indicate that all three oxidase types have the capacity to pump protons. Previously, however, we have reported decreased H<sup>+</sup>/e<sup>-</sup> coupling efficiencies of the *cbb*<sub>3</sub>-type

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oxidase under certain conditions. Sequence alignment suggests that many residues that have been proposed to constitute the chemical and pumped proton channels in cytochrome  $aa_3$  (and probably also in cytochrome  $bb_3$ ) are not conserved in cytochrome  $cbb_3$ . It is concluded that the design of the proton pump in cytochrome  $cbb_3$  differs significantly from that in the other oxidase types.

#### Introduction

In mitochondrial respiration the reduction of oxygen to water is catalysed by the  $aa_3$ -type cytochrome *c* oxidase. This membrane-bound protein complex is the last component of a linear respiratory pathway in which electrons from NADH and succinate are transferred, in a sequence of redox reactions, to oxygen. The aerobic respiration in bacteria, on the other hand, often proceeds via a more complex electron-transfer network. Alternative respiratory pathways enable a bacterium to adjust the composition and/or the efficiency of oxidative phosphorylation in response to changes in environmental conditions (Anraku and Gennis, 1987; van der Oost *et al.*, 1994a; Garcia-Horsman *et al.*, 1994a; van Spanning *et al.*, 1995a).

In addition to a number of alternative guinone-reducing dehydrogenases, the terminal oxidases appear to represent an important site of flexibility in respiratory networks. To date, the best-characterized bacterial respiratory system is the one in Escherichia coli. In this bacterium two distinct quinol oxidases (cytochromes bo3 and bd) are expressed during aerobic growth, but a cytochrome c branch is absent (Anraku and Gennis, 1987). Paracoccus denitrificans, in turn, possesses three distinct terminal oxidases. An aa<sub>3</sub>-type cytochrome c oxidase (Raitio et al., 1987; 1990; Steinrücke et al., 1987; van Spanning et al., 1990; van der Oost et al., 1991), as well as a bb3/ba3-type quinol oxidase, hereafter called cytochrome bb<sub>3</sub>, have been characterized previously (de Gier et al., 1994; Richter et al., 1994). In addition, the expression of an alternative cytochrome c oxidase has been suggested (Bosma, 1989; de Gier et al., 1992; 1994; Raitio and Wikström, 1994).

In the present study, a *P. denitrificans*  $\Delta aa_3/bb_3$  mutant

was used for the purification of an alternative cytochrome c oxidase: cytochrome  $cbb_3$ . Using the N-terminal amino acid sequence of one of its subunits, a genomic locus, ccoNOQP, was isolated that contained a gene cluster with four open reading frames. Sequence analysis of the flanking regions revealed the presence of a ccoGH cluster downstream of ccoNOQP. In the upstream region two open reading frames were found, the derived sequences of which are homologous to HemN and FNR.

Analysis of the oxygen consumption by single and multiple oxidase mutants of *P. denitrificans* suggests that, besides cytochrome  $aa_3$  and  $cbb_3$ , no additional cytochrome *c* oxidase is present. The currently generated set of oxidase mutants of *P. denitrificans* was used to demonstrate that all three terminal oxidases have the capacity to pump protons. It is discussed, however, that in the  $cbb_3$ type cytochrome *c* oxidase the coupling between oxygen reduction and proton translocation may be less tight than in cytochromes  $aa_3$  and  $bb_3/bo_3$ .

#### Results

#### Purification of the cbb3-type cytochrome c oxidase

In a previous study, the simultaneous deletion of cytochromes  $aa_3$  and  $bb_3$  in P. denitrificans ( $\Delta aa_3/bb_3$ ) has been reported. Analysis of this mutant revealed (i) the presence of an alternative cytochrome c oxidase, (ii) the presence of only protohaem IX (haem B) in membrane extracts, and (iii) an increased expression of cytochromes c and b. No quinol oxidase activity was detectable in this mutant (de Gier et al., 1994). Here, we report the isolation of a cb-type cytochrome c oxidase from membranes of the Paracoccus  $\Delta aa_3/bb_3$  mutant. Both the activity and the subunit composition of the oxidase complex turned out to be rather unstable during the purification, as described in the Experimental procedures. As will be discussed below, a cytochrome c subunit with an apparent molecular mass of 45 kDa is lost during the purification of Paracoccus cytochrome cbb<sub>3</sub> (as described in the Experimental procedures).

Analysis of the purified cytochrome *c* oxidase by a Coomassie brilliant blue-stained SDS-PAGE gel indicates that it consists of two components with apparent molecular masses of 45 kDa and 30 kDa (Fig. 1A). Only the fast-migrating component has been identified as a cytochrome *c* because of the presence of covalently bound haem, as demonstrated by haem-stained SDS-PAGE (not shown). Sequencing of this 30 kDa subunit revealed the N-terminal sequence AILEKHKVLEKNATLLLVFSFLVVRI. The optical spectrum of this oxidase preparation indicates that it involves a *cb*-type cytochrome *c* in this spectrum relatively small amount of cytochrome *c* in this spectrum relative to a previous preparation (de Gier *et al.*, 1994) can best be explained as the loss of a second haem



Fig. 1. A. SDS-PAGE of purified cytochrome  $cbb_3$  (approx. 7.5 µg of protein). The molecular mass standards are: 97.4, 66.2, 45.0, 31.0, 21.5, and 14.4 kDa.

B. Absorption spectrum of purified cytochrome  $cbb_3$  after reduction with Na-dithionite (approx. 0.01 mg protein ml<sup>-1</sup>).

C-containing subunit which would co-migrate with the 45 kDa polypeptide, as demonstrated below.

#### Isolation and characterization of the ccoNOQP cluster

Based on the N-terminal sequence of the 30 kDa subunit, degenerate primers were designed (see the *Experimental procedures*, primers 171 and 172) and used to perform the polymerase chain reaction (PCR) on genomic DNA from the  $\Delta aa_3/bb_3$  mutant. A PCR product of the expected size (0.1 kb) was obtained. Sequence analysis of the PCR product indicated that the DNA sequence corresponded to the N-terminal peptide of the 30 kDa subunit (not shown). With the PCR product as a probe, the complete *ccoNOQP* cluster could be cloned (Fig. 2A). Fragments were subcloned into M13mp18/19 for sequence analysis. The complete DNA sequence has been submitted to the GenBank database Data Library (Accession Number Banklt12589 U34353).

Four open reading frames (ORFs) in the isolated genomic locus share a high degree of homology with the *ccoNOQP* gene cluster from *Rhodobacter capsulatus* (Thöny-Meyer *et al.*, 1994) and to the *fixNOQP* operons of *Rhizobium meliloti* (Kahn *et al.*, 1993; D. Kahn, 1993, EMBL Data Library Accession No. 221854), *Bradyrhizobium japonicum* (Preisig *et al.*, 1993; Fig. 3) and *Azorhizobium caulinodans* (Mandon *et al.*, 1994). As *P. denitrificans* is not known to be able to fix molecular nitrogen, the *ccoNOQP* rather than the *fixNOQP* nomenclature



(XhoI)

Km<sup>R</sup>

Fig. 2. A. Genetic organization and physical map of the *P. denitrificans ccoNOQP* region. Arrows represent individual genes. The positions of FNR-boxes are indicated by arrows.

B. Insertion mutants have been generated by substituting part of the *ccoNOQP* operon with a kanamycin-resistance ( $Km^R$ ) cassette, deleting the 3' half of *ccoN* and the two 5' codons of *ccoO*.

has been adopted for these genes. The four ORFs have a GC-rich codon preference which is typical of P. denitrificans. The ccoNOQP genes are tightly clustered, with the exception of the ccoQ and ccoP genes which are 41 bases apart. As no transcription-termination signal was found here, the four genes are probably transcribed as a single operon, as has been demonstrated for R. meliloti (Batut et al., 1989). The proposed initiation codons of all four genes are six to eight bases downstream of putative Shine-Dalgarno sequences. A putative anaerobox (TTGAC-N<sub>4</sub>-ATCAA) is located upstream of ccoN. Its sequence is very similar to the binding sites of FNR in E. coli and of FixK in R. meliloti and B. japonicum (TTGAT-N<sub>4</sub>-ATCAA; Spiro, 1994), and identical to the anaerobox found upstream of the A. caulinodans fixNOQP operon (Mandon et al., 1994).

(XhoI

Attempts to sequence the N-terminus of subunit I (CcoN, apparent molecular mass 45 kDa (Fig. 1A)) were unsuccessful, probably because of blocking of the first amino group. The N-terminal methionine of CcoN (Fig. 3A) has been chosen because the corresponding ATG codon is downstream of a Shine-Dalgarno sequence, and because no alternative in-frame initiation codons are present. Moreover, the size of the gene correlates with that of ccoN/fixN in the aforementioned bacteria. The putative anaerobox is located at positions -110 and -124 of the chosen start codon. Based on sequence alignment with the subunits I of the well-characterized cytochromes aa3 from Rhodobacter sphaeroides and P. denitrificans, as well as with the subunit I of cytochrome bo3 from E. coli (Hosler et al., 1993), four histidines in the CcoN sequence (H271, H321, H322 and H409) are the most probable ligands of the binuclear catalytic centre formed by a penta

co-ordinated haem B and a copper atom,  $Cu_B$  (Fig. 3A). The ligands of the hexa co-ordinated haem B are H411 and (probably) H122 in the *Paracoccus* CcoN (Fig. 3A).

CcoO has an N-terminal amino acid sequence that corresponds to the N-terminal sequence determined for the 30 kDa subunit (Fig. 3B). Apart from this hydrophobic, putative membrane-spanning fragment, CcoO contains a hydrophilic, probably periplasmic, domain with a Cx<sub>2</sub>CHx<sub>n</sub>M motif which is the consensus for haem C binding (Fig. 3B). Apart from this motif, CcoO/FixO does not share much similarity with the available cytochrome c sequences (Moore and Pettigrew, 1990). CcoQ is a small protein with one hydrophobic region (Fig. 3C); apart from CcoQ and FixQ, no significant homologues have been found in sequence databases. CcoP appears to be a membrane-anchored di-haem cytochrome c. Its C-terminal hydrophilic domain, probably facing the periplasmic side of the plasma membrane, contains two motifs (Gx<sub>3</sub>Fx<sub>3</sub>Cx<sub>2</sub>CHx<sub>n</sub>M) that are well conserved among soluble mono-haem cytochromes of photosynthetic bacteria and eucaryotic algae (Mandon et al., 1994). In contrast to the FixP/CcoP sequences from other sources, an N-terminal hydrophilic extension is present in the Paracoccus sequence (Fig. 3D), probably located at the cytoplasmic side of the membrane.

#### Flanking regions

Upstream of the *ccoNOQP* cluster of *P. denitrificans* is an ORF that potentially codes for a polypeptide of 278 amino acids (orf278 in Fig. 2A). It shares a rather high degree of homology (28% identical amino acids, not shown) with orf277, which is located adjacent to the *fixNOQP* operon

Α					
Pd CcoN 1	MLDTIKLIALGTIAVLAAIAANYARPDDLAYLVNALIIMLAAGIMFLRVLROMGNEOPA				
Bj FixN 1	MSQPSISKSMTIGESGLAVVFAATAFLCVIAAAKALDAPFAFHAALSAAASVAAVFCIVNRYFERPA				
Pd.CtaD 1	MADAAVH				
Ec CyoB 1	MFGKLSLDAVPFHEPIVMVTIAGIILGGLALVG				
	my r				
Pd CooN 60					
Bi FixN 68	ALPPAEINGRPNYNMGPIKESSEMAMEWGTAGELVGT I LASOLAWDALNEDL				
Pd CtaD 8	GHGDHHDTRGFFTRWFMSTNHKDIGTLYLFTAGIVGLTSVCFTVYMRMFLOHPGVOYMCLFGARLTA				
Ec CyoB 34	LITYFGKWTYLWKEWLTSVDHKRLGIMYIIVAIVMLLRGFADAIMMRSOOALA				
	TM II TM III				
Pd CcoN 104	ITMGYTNFGKLRPLHTSAVIFAFGGNGLIATSFYVVQRTSAARLWGGNAAWFVFWG				
BJ FIXN II6	LAGDLAPWFVVVG				
FG CUOB 87	DASALCIPNGHLWNVMIIIHGVLMMFFVVIPALFGGFGNIFMPLHIGAPDMAFPRLNNLSYWMYVCG SACFACFLDDHHYDOTFTAHCVIMIFEVAMDFVIC-IMNIUUDIOTCADDVAFDFYNNI STNDTUDUC				
De eyeb er					
	94 113 124 131				
Dd CooN 164	TM IV				
Bi FixN 173	YNFFILVAGTGYLLGVTOSKFVAPPPWYNDIWLTVVWVVYLAVFLGTILK				
Pd CtaD 142	VALGVASLLAPGGNDOMGSGVGWVLYPPLSTTEAGYSMDIATFAVHUSCASSTICATNITTETIN				
Ec CyoB 153	VILVNVSLGVGEFAQTGWLAYPPLSGIEYSPGVGVDYWIWSLOLSGIGTTLTGINFFVTILK				
1	N T				
	199 203 TM 17				
Pd CcoN 214	RKEP-HTYVANWEVT SETUPTIMI HTUNNI ATPUST ECSKSVOT E				
Bi FixN 223	RKEP-HIFVANWFYLAFTVTTAVIHIGNNPALPVSAFGSKSYVAWCGTODAMFOWW				
Pd CtaD 207	MRAPGMTLFKVPLFAWSVFITAWLILLSLPVLAGAITMLLMDRNFGTOFFDPAGGGDPVLYOHILWF				
Ec CyoB 216	MRAPGMTMFKMPVFTWASLCANVLIIASFPILTVTVALLTLDRYLGTHFFTNDMGGNMMMYINLIWA				
	TM VII TM VIII				
Pd CcoN 269	YGHNAVGFFLTAGFLGMMYYFIPKQAERPVYSYKLSIIHFWALIFLYIWAGPHHLHYTALPDWASTL				
BJ FIXN 278 Pd Ctap 274	IGHNAVGFFLIAGFLAIMYYFIPKRAERPIYSYRLSIIHFWALIFLYIWAGPHHLHYTALPDWTQTL				
Ec CyoB 282	CORE AVIIIIAE OF GIISRVISIF - ARREIF CYTELUNATAALGILGE VVWAHHMIIAGMSLTQQAY				
1	HEY S HH				
	276 278 280 291 325 326				
	TM IX				
Pd CcoN 336	GMVFSIILWMPSWGGMINGLMTLSGAWDKLRTDPIIRMMVVAVGFYGMATFEGPMMSIKAVNFVSHY				
Bj FixN 345	GMTFSIMLWMPSWGGMINGLMTLSGAWDKLRTDPVLRMLVVSVAFYGMSTFEGPMMSIKVVNSLSHY				
Pd CtaD 340	FMLATMTIAVPTGIKVFSWIATMWGGSIEFKTPMLWAFGFLFLFTVGGVTGVVLSQAPLDRVYHD				
Ec CyoB 348	FGITTMIIALPTGVKIFNWLFTMYQGRIVFHSAMLWTIGFIVTFSVGGMTGVLLAVPGADFVLHN				
	T T K L D				
	TM X TM XI				
Pd CcoN 403	TDWTIGHVHSGALGWNGMITFGALYYLVPRLWGR-ERLYSTGLVSWHFWLATIGLVLYAASMWVSGI				
Bj FixN 412	TDWTIGHVHSGALGWVGFVSFGALYCLVPWAWNR-KGLYSLKLVNWHFWVATLGIVLYISAMWVSGI				
Fa Cuap 413	TIIVVAHFHIVMSLGAVFGIFAGVYYWIGKMSGRQYPEWAGQLHFWMMFIGSNLIFFPQHFLGR				
ве субв 413	BELETARE REVISES VEGETARMIT WWPKAFGFKLNETWGKRAFWFWIIGFFVAFMPLYALGF				
	411.413				
-	TM XII				
Pd CcoN 469	MEGLMWREVDAQGFLVNAFADTVAAKFPMNVVRALGGVLYLFFALIMCYNLWATVAKQPKTQSTAAA				
BJ FIXN 478	LQGLMWKAYTSLGFLEYSFIETVEAMHPFYIIRAAGGGLFLIGALIMAYNLWMTVRVGEAEVQMPVA				
EC CYOB 477	Q-ONE ANTIDIEVERAIMMNISSIGATISEASELEFIGIVEYTILFAGKRVNVPNYWNEHADT				
DC CYOD 4//	M GHIRRDSQUIDEQE RINDHIAASGAVLIALGILULVIQMIVSIRDRDQNRDLTGDPWGGRT				
Pd CcoN 536	VPAE				
Bj FixN 545	LQPAE				
Pd CtaD 530	LEWTLPSPPPEHTFETLPKREDWDRAHAH				
Ес СуоВ 539	LEWATSSPPPFYNFAVVPHVHERDAFWEMKEKGEAYKKPDHYEEIHMPKNSGAGIVIAAFSTIFGFA				
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В			
Pd CcoO Bj FixO	1 1	MAILEKHKVLEKNATLLLVFSFLVVTIGGIVEIAPLFYLQNTIE MSFWTRHQVFEKNSIILIVGILLVIAIGGLVEITPLFYLKSTIE	44 44
Pd CcoO Bj FixO	45 45	KVQGMRPYTPLELKGRDIYVREGCYVCHSQMIRPM-RDEVERYG KVDGVRPYTPLELAGRNVYVREGCYLCHSQMIRPLRDEV-ERYG	87 87
Pd CcoO Bj FixO	88 88	H Y S L A A E S M Y D H P F Q W G S K R T G P D L A R V G G R Y S D E W H L D H L V D P H F S L A A E S M F D H P F Q W G S K R T G P D L A R V G A K Y S D D W H V T H L T N P	131 131
Pd CcoO Bj FixO	132 132	QAVVPESIMPKYGF – LLNRQVDASNMQQRLKTDALG – GVPYDDA RAIVPQSVMPGYPFLSATEVDPDTIADHMRTLRTVGVPYTDDQI	173 175
Pd CcoO Bj FixO	174 176	M I A A A G E D F R V Q A A P D A D A S G L E E R Y P G – A Q Q R N F D R R P G – V S E A N A S A D L K A Q A D P D N A G A D A F N K R Y A K A V V R N F D – G K T G T P – T E	215 217
Pd CcoO Bj FixO	216 218	MDALIAYLQVLGTMVDFSTFEPDPNR MDALIAYLQMLGTLVDFKIYNEKANLR	241 244
С			
Pd CcoQ Bj FixQ	<b>1</b> 1	M D R Y S F L R E L A D S W V L L L L V V F F L G T I V F A F R P G F A A A A S M K A I L T L D N L A S G L V T T I W T P V F V A I F L A I I A Y A F W P R N K A A F D	40 44
Pd CcoQ	41	RRGRKHLP	48
Bj FixQ	45	EAAHLPLIREE	54
Bj FixQ	45	EAAHLPLIREE	54
Bj FixQ D	45	MADTDDEHASPONDDNRIELEROAADEAHKAKILAHPPEAGGDP	54
Bj FixQ D Pd CcoP Bj FixP	45 1 1	E AAH L PLREE MADT D D E HASP QNPD NRIELER QAAD EAH KAKILAH PPEAGG DP MTDHSEFD S	54 44 9
D Pd CcoP Bj FixP Pd CcoP Bj FixP	1 1 45 10	E AAH LPLREE MADTDDEHASPQNPDNRIELERQAADEAHKAKILAHPPEAGGDP MTDHSEFDS	54 44 9 88 30
D Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP	45 1 1 45 10 89 31	E AAH L PLREE MADT D D E HASP QNPD NRIELER QAAD EAHKAKILAHPPEAGGDP MTDHSEFDSVSTGHSWDGIEEYDNPL VSGKTTTGHEWDGIKELNTPL PRWWLWTFYATIVWGVLYLIAYPAIPLVNGATQGLLGQNYRSDV PRWWVICFYLTIVWAIGYWIVYPAWPLISSNTTGLFGYSSRADV	44 9 88 30 132 74
D Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP	45 1 1 45 10 89 31 133 75	E AAH L PLREE MADTDDEHASPONPDNRIELERQAADEAHKAKILAHPPEAGGDP MTDHSEFDSVSGRTGHSWDGIEEYDNPL VSGKTTTGHEWDGIKELNTPL PRWWLWTFYATIVWGVLYLIAYPAIPLVNGATOGILGONYRSDV PRWWVICFYLTIVWAIGYWIVYPAWPLISSNTTGLFGYSSRADV AAEIORFNEANAPIOAKLVETPLEEIAADPELANYTANAGAAIF AVELANLEKIRGDKMAALGAASLADVEKDPALLALARAKGKTVF	54 44 9 88 30 132 74 176 118
D Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP	45 1 1 45 10 89 31 133 75 177 119	E AAH L PLREE MADTD DE HASP ON PDNRIELER QAADEAHKAKILAHPPEAGGDP MTDHSEFDS	44 9 88 30 132 74 176 118 220 162
D Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP Pd CcoP Bj FixP	45 1 1 45 10 89 31 133 75 177 119 221 163	E AAH L PLREE MAD T D D E HA S P Q N P D N R I E L E R Q A AD E A H K A K I L A H P P E A G G D P M T D H S E F D S VS G K T T G H S W D G I E E Y D N P L VS G K T T G H E W D G I K E L N T P L P R W W L W T F Y A T I V W G V L Y L I A Y P A I P L V N G A T Q G L L G Q N Y R S D V P R W W V I C F Y L T I V W A I G Y W I V Y P A W P L I S S N T T G L F G Y S S R A D V A A E I Q R F N E A N A P I Q A K L V E T P L E E I A AD P E L A N Y T A N A G A A I F A V E L A N L E K I R G D K M A A L G A A S L A D V E K D P A L L A L A R A K G K T V F R T W C A Q C H G S G A G G A T G Y P S L L D N D W L W G G T L E E I H T T V M H G I R G D N C A P C H G S G G A G A K G F P N L N D D D W L W G G T L D Q I M Q T I Q F G A R D P K D A D T R Y S E M P R F G I D G L L E N A Q I S Q V V N H V L E L G G L P H D A A S G H A K T H E G Q M L A F G K D G V L K G D E I V T V A N Y V R S L S G L P T R K G Y	44 9 88 30 132 74 176 118 220 162 264 206
D Pd CcoP Bj FixP Pd CcoP Bj FixP	45 1 45 10 89 31 133 75 177 119 221 163 265 207	EAAHLPEREE MADTDDEHASPQNPDNRIELERQAADEAHKAKILAHPPEAGGDP MTDHSEFDS	44 9 88 30 132 74 176 118 220 162 264 206 308 250

Fig. 3. Alignments of the P. denitrificans ccoNOQP-encoded polypeptides.

A. CcoN of *P. denitrificans* (cytochrome *cbb*<sub>3</sub>), FixN of *B. japonicum* (cytochrome *cbb*<sub>3</sub>; Preisig *et al.*, 1993), CtaDII of *P. denitrificans* (cytochrome *aa*<sub>3</sub>; Raitio *et al.*, 1990), and CyoB of *E. coli* (cytochrome *bo*<sub>3</sub>; Chepuri *et al.*, 1990). Transmembrane helices are shaded and conserved residues discussed in the text are indicated.

B. CcoO of P. denitrificans and FixO of B. japonicum (Preisig et al., 1993). Identical residues are boxed.

C. CcoQ of P. denitrificans and FixQ of B. japonicum (Preisig et al., 1993). Identical residues are boxed.

D. CcoP of P. denitrificans and FixP of B. japonicum (Preisig et al., 1993). Identical residues are boxed.

of *B. japonicum*. No homologue has yet been reported in other members of the Rhizobiaceae family. As no mutants are available, a link between this gene product and cytochrome  $cbb_3$  remains to be demonstrated. Further upstream, two genes are located that encode homologues of *E. coli* HemN and FNR (Fig. 2A). In the latter bacterium both polypeptides play an important role during oxygen limitation: HemN catalyses an oxygenindependent bypass in haem biosynthesis (Plunkett *et al.*, 1993), and FNR is a transcription activator of genes involved in anaerobic metabolism (Spiro and Guest, 1990).

Without exception, the gene cluster fixGHIS is located downstream of the fixNOQP genes of the Rhizobiaceae family (Kahn et al., 1993; Preisig et al., 1993; Mandon et al., 1994) and ccoNOQP genes of R. capsulatus (Thöny-Meyer et al., 1994). In R. meliloti, a number of transposon insertions in fixGHI have been isolated which had a 'fixminus' phenotype (Kahn et al., 1989). Also, in Paracoccus at least part of this cluster is located adjacent to the ccoNOQP genes. Two genes homologous to fixG and fixH (Fig. 2A) have been found downstream of ccoNOQP but, for the reasons given above, these genes are designated ccoG and ccoH. In all instances an anaerobox is located in the fixG and in the ccoG promoter regions (Fig. 2A), suggesting expression of this oxidase type at low oxygen tensions. The ccoG product is a ferredoxin-like protein, with two typical 4Fe4S-binding motifs. FixGHIS has been suggested to be a polypeptide complex consisting of, at least, a redox protein (FixG) and a cation pump (FixI) (Kahn et al., 1989).

#### Mutagenesis of ccoNO

Insertion mutants were generated by substituting part of the *ccoNOQP* operon by a kanamycin-resistance (Km<sup>R</sup>) cassette, deleting the 3' half of ccoN and the two 5' codons of ccoO (Fig. 2B). The ccoNOQP operon was deleted in wild-type Paracoccus (Pd1222) and in the ctaDI/ctaDII double mutant Pd92.20 ( $\Delta aa_3$ ), yielding Pd27.21 ( $\Delta cbb_3$ ) and Pd93.12 ( $\Delta aa_3/cbb_3$ ), respectively. Southern blot analysis of chromosomal DNA of selected Km<sup>R</sup> clones confirmed the appropriate recombination (not shown). Analysis of the mutant strains generated, grown on minimal medium with succinate, did not show significant spectroscopic changes relative to their respective parent strain. SDS-PAGE and subsequent haem staining of cytoplasmic membranes showed that the 30 kDa cytochrome c is absent, and the 45 kDa cytochrome c is only just detectable in the  $\Delta cbb_3$  mutant strains (Fig. 4).

#### Oxygen consumption

The oxygen-consumption rates in cell suspensions of wild

type and oxidase mutants were measured polarographically, with endogenous as well as exogenous substrates (succinate or ascorbate with *N*,*N*,*N'*,*N'*-tetramethyl-*p*phenylenediamine (TMPD)). The oxidation rates are shown in Table 1. In separate assays the electron flow to oxygen via cytochrome *c* reductase (the *bc*<sub>1</sub> complex) was inhibited by addition of either antimycin A or myxothiazol. The endogenous respiration of the wild type was clearly affected by these inhibitors. In contrast, the endogenous respiration of the  $\Delta aa_3$  mutant and the  $\Delta cbb_3$  mutant was only slightly inhibited, and that of the  $\Delta aa_3/cbb_3$ mutant strain appeared to be unaffected.

The effect of the cytochrome  $bc_1$  inhibitors on succinate oxidation in the  $\Delta cbb_3$  mutant and in the  $\Delta aa_3$  mutant was much less than in the wild type. Again, in the  $\Delta aa_3/cbb_3$  mutant the succinate oxidation appeared to be unaffected by these inhibitors. More or less equal values of ascorbate/TMPD oxidation indicated the presence of cytochrome *c* oxidase(s) in the wild type, as well as in the  $\Delta cbb_3$  and the  $\Delta aa_3$  mutant. In the  $\Delta aa_3/cbb_3$  mutant, the endogenous respiration was not enhanced at all upon the addition of ascorbate/TMPD, indicating that no cytochrome *c* oxidase was expressed.

#### Proton translocation

The proton-pumping capacity of the set of oxidase mutants was measured by using the oxygen-pulse method as described elsewhere (Raitio and Wikström, 1994; de Gier *et al.*, 1994). In the proton-translocation assay, succinate was used as the electron-donating substrate and either oxygen or potassium ferricyanide was used as the terminal electron acceptor (Table 2). The main goal was to study the proton-pumping capacity of the *cbb*<sub>3</sub>-type cytochrome *c* oxidase. The H<sup>+</sup>/e<sup>-</sup> ratios of the  $\Delta aa_3$  mutant and the  $\Delta bb_3/aa_3$  mutant clearly show that the *cbb*<sub>3</sub>-type cytochrome *c* oxidase does have the capacity to translocate protons. In Fig. 5, representative proton-translocation traces of the  $\Delta bb_3/aa_3$  mutant and the  $\Delta aa_3/cbb_3$ 





**Table 1.** Oxygen-consumption measurements of whole-cell suspensions of different *P. deni-trificans* strains: Pd1222 (wild type), Pd92.20 ( $\Delta aa_3$  mutant), Pd27.21 ( $\Delta cbb_3$  mutant), and Pd93.12 ( $\Delta aa_3/cbb_3$  mutant).

	Substrate inhibitor	Endogenous		Succinate		e		
Strain		_	AA	myx	-	AA	myx	Ascorbate/ IMPD -
Wild type		1.5	0.9	0.9	2.4	0.6	0.5	3.7
Δaa <sub>3</sub> mutant		1.3	1.2	1.1	2.9	2.6	2.7	2.5
$\Delta cbb_3$ mutant		1.4	1.2	1.3	2.7	1.9	1.7	2.6
$\Delta aa_3/cbb_3$ mutant		2.2	2.2	2.2	2.1	2.1	2.1	0.0

Rates after addition of exogenous substrate have been corrected by subtraction of the corresponding endogenous rates. Values are the means of three independent assays (nmol  $O_2$  per mg of protein per s). Inhibitors: antimycin A (AA), myxothiazol (myx).

mutant confirm that the  $aa_3$ -type cytochrome c oxidase and the  $bb_3$ -type quinol oxidase both translocate protons. Upon the addition of inhibitors of the  $bc_1$  complex, the H<sup>+</sup>/e<sup>-</sup> ratio drops from 3 to 2 (not shown). This is consistent with the decreased ratio of  $2H^+/e^-$  that is detected only in the  $\Delta aa_3/cbb_3$  mutant. This suggests that ubiquinol apparently was preferentially oxidized by ubiquinol:cytochrome c oxidoreductase ( $bc_1$  complex) rather than by ubiquinol:oxygen oxidoreductase (cytochrome  $bb_3$ ).

With the artificial electron-donating couple ascorbate/ TMPD, proton translocation was barely detectable in the absence of cytochrome  $aa_3$ . However, at elevated TMPD concentrations (final concentration: 2–3 mM) some proton translocation was observed, albeit never more than 50% of the theoretical maximum (not shown). The oxygenconsumption experiments indicate that, in the absence of cytochrome  $aa_3$  and cytochrome  $cbb_3$ , no electron flow from ascorbate to oxygen is detectable in succinategrown cells of *P. denitrificans*. Accordingly, no proton extrusion was observed with ascorbate/TMPD in the  $\Delta aa_3/cbb_3$  mutant.

#### Discussion

*P. denitrificans* contains a respiratory network in which electrons are transferred from specific dehydrogenases to different types of terminal oxidases. One electron

 Table 2. Proton-translocation measurements of whole-cell suspensions of different *P. denitrificans* strains.

Strain	Succinate O <sub>2</sub> (H*/e <sup></sup> )	Succinate Fe(CN) <sub>6</sub> <sup>3-</sup> (H*/e <sup>-</sup> )
Wild type	2.93 ± 0.13	1.96 ± 0.09
$\Delta aa_3$ mutant	$2.96 \pm 0.15$	1.94±0.18
∆bb <sub>3</sub> mutant	$2.97 \pm 0.12$	1.96±0.12
$\Delta aa_3/bb_3$ mutant	$2.95 \pm 0.12$	1.98±0.16
$\Delta cbb_3$ mutant	$3.01 \pm 0.14$	1.94±0.13
$\Delta aa_3/cbb_3$ mutant	$1.98 \pm 0.11$	$1.97 \pm 0.09$

 $H^+/e^-$  ratios are averages ± standard deviation (*n*=8). Cells were cultivated as aerobic batch cultures in minimal medium with succinate. The electron donor was succinate; anaerobic suspensions were pulsed with either oxygen (O<sub>2</sub>) or ferricyanide (Fe(CN)<sup>3-</sup><sub>6</sub>).

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transfer route closely resembles the mitochondrial respiratory chain in which ubiquinol, reduced by either NADH or succinate, is oxidized by a supercomplex consisting of cytochrome *c* reductase ( $bc_1$  complex), cytochrome  $c_{552}$ and cytochrome *c* oxidase (cytochrome  $aa_3$ ) (Berry and Trumpower, 1985; Trumpower, 1991). In addition, it has been demonstrated that *Paracoccus* expresses a distinct



Fig. 5. Proton translocation in whole cells from strain Pd93.11 ( $\Delta aa_3/bb_3$ ) with succinate as substrate. An anaerobic cell suspension was pulsed with equal volumes (10 µl) of air-saturated water (258 µM O<sub>2</sub>), ferricyanide (1.0 mM FIC) and acid (1.0 mM HCl) to calibrate the system. The H<sup>+</sup>/e<sup>-</sup> ratio with oxygen is 3.0 and with ferricyanide the H<sup>+</sup>/e<sup>-</sup> ratio is 2.0.

respiratory branch in which electrons are directed to a second terminal oxidase, the quinol-oxidizing cytochrome  $bb_3$  (de Gier *et al.*, 1994).

Analysis of a recently generated  $\Delta aa_3/bb_3$  mutant revealed that the aerobic respiration proceeds via an alternative cytochrome *c* oxidase (de Gier *et al.*, 1994). In the present study, the allocated gene cluster of this *cbb*<sub>3</sub>type oxidase has been cloned and sequenced. It has previously been demonstrated that no quinol oxidase activity remains in a  $\Delta bb_3$  mutant (de Gier *et al.*, 1994), and here we show that a  $\Delta aa_3/cbb_3$  double mutant does not express cytochrome c oxidase activity (Table 1) (van der Oost *et al.*, 1995). Hence, it is concluded that, at least under a variety of tested conditions, these three types form the complete set of terminal oxidases expressed in *P. denitrificans*.

#### Characterization of cytochrome cbb3

The products of the isolated P. denitrificans ccoNOQP gene cluster share a high degree of homology with the cytochrome c oxidase of R. capsulatus (Thöny-Meyer et al., 1994) and the 'high-affinity' oxidase from several members of the Rhizobiaceae (Kahn et al., 1993; Preisig et al., 1993; Mandon et al., 1994) (Fig. 3). Although the cbb3type oxidase apparently represents a distant member of the haem-copper oxidase family (van der Oost et al., 1994; Saraste and Castresana, 1994), its cytochrome b subunit (CcoN) still shares some typical features with the catalytic subunit (subunit I) of these oxidases. Six histidines that are invariant in the available CcoN/FixN sequences have been aligned with the conserved histidine ligands of the redox centres of subunit I of cytochromes aa<sub>3</sub> and bo<sub>3</sub>. The alignment shown in Fig. 3A suggests that H271, H321, and H322 are the most probable ligands of Cu<sub>B</sub>, H409 of the penta co-ordinated haem B, and H411 and H122 of the hexa co-ordinated haem B in P. denitrificans CcoN.

Garcia-Horsman et al. (1994b) have recently reported a thorough biochemical analysis of the cbb<sub>3</sub>-type oxidase from R. sphaeroides and have demonstrated the presence of a binuclear centre typical of haem-copper oxidases and the absence of Cu<sub>A</sub>. Moreover, metal analysis revealed the ratio of haemC:haemB:copper to be 3:2:1. This agrees very well with the sequence data of cytochrome cbb<sub>3</sub>: two haems B and a copper ion are located in subunit I (CcoN), and three haems C reside in the two cytochrome c subunits (one in CcoO and two in CcoP). Bosma (1989) has reported a 30 kDa cytochrome c associated with the alternative oxidase from P. denitrificans with an E<sub>m7</sub> value of 322 mV. Gray et al. (1994) purified cytochrome cbb<sub>3</sub> from R. capsulatus and reported a 28 kDa cytochrome c (CcoO) with an  $E_{m7}$  value of 320 mV, a 32 kDa cytochrome c (CcoP) with an  $E_{m7}$ value of 265 mV, and a low-spin cytochrome b with an  $E_{m7}$  value of 385 mV. These data may indicate that CcoO is the direct electron donor of the redox centres (haems B and Cu<sub>B</sub>) in CcoN. Indeed, the purified 2 subunit CcoNO complex (Fig. 1) retains TMPD oxidase activity. Apparently CcoP is less tightly bound to the cytochrome cbb<sub>3</sub> core and may play a role as electron carrier from the bc1 complex to the CcoO-CcoN complex, which would be in agreement with the reported midpoint potential values.

#### Expression

Comparison of membrane proteins from the wild type and from the  $\Delta cbb_3$  mutant of *P. denitrificans* by haem-stained SDS-PAGE (Fig. 4) identified the 30 kDa and 45 kDa cytochromes c as the CcoO and CcoP subunits of cytochrome cbb<sub>3</sub>, respectively. Previously, Bosma (1989) performed a detailed analysis of the expression of *c*-type cytochromes in chemostat-grown cells of P. denitrificans, using the same haem analysis (reviewed by van Spanning et al., 1995a). Bosma demonstrated that both the 30 kDa (CcoO) and the 45 kDa (CcoP) cytochromes c were abundantly present in membrane fractions when cells were grown in either nitrate-limited (anaerobically) or oxygenlimited (microaerobically) conditions. In succinate-limited (aerobic) cultures, on the other hand, the 45 kDa cytochrome c was absent and the 30 kDa cytochrome c was barely detectable (Bosma, 1989). It was concluded that the cbb<sub>3</sub>-type oxidase plays an important role at decreased oxygen tensions.

In the promoter regions of all presently known cco/ fixNOQP and cco/fixGH(IS) clusters is a sequence motif that closely resembles the binding site of the well-known anaerobic/microaerobic transcription activator FNR (Spiro and Guest, 1990). In R. meliloti, the 'oxygen state' is sensed by the haem protein FixL. In the absence of oxygen, this protein is autophosphorylated (FixL~P), Subsequently, the phosphate is transferred from the sensor to the regulator, FixJ, inducing its activation (FixJ~P). The latter modification results in transcription activation of fixK, the gene that encodes an FNR homologue. FixK, in turn, binds at the anaerobox upstream of the fixNOQP and fixGHIS clusters, thereby activating their transcription (David et al., 1988; Batut et al., 1989; Kahn et al., 1989; Fischer, 1994). To date, no FixLJ system has been demonstrated in any non-nitrogen-fixing organism. However, a FixK-like protein, NNR, has been identified in P. denitrificans as a transcription activator of nitrite reductase and nitric oxide reductase, two polypeptide complexes that are involved in anaerobic respiration (van Spanning et al., 1995b). Mutagenesis of the nnr gene, however, did not affect expression of cytochrome cbb<sub>3</sub>. Here we show that in the flanking region of the cco locus of Paracoccus, a gene is located that potentially codes for a second transcription regulator: fnrP (Fig. 2A). Unlike NNR and FixK, this FNR homologue has an N-terminal cysteine cluster which, in the case of E. coli FNR, has been demonstrated to be involved in the binding of iron. The redox state of this iron ion determines whether the regulator is active or not (Spiro and Guest, 1990).

#### Energy conservation

Cytochromes aa3 and bb3 from P. denitrificans are proton-

pumping terminal oxidases (van Verseveld *et al.*, 1981; Solioz *et al.*, 1981; Puustinen *et al.*, 1989). Analysis of proton translocation in cell suspensions of the *Paracoccus* oxidase mutants offers the unique opportunity to measure proton translocation of individual oxidases in whole-cell suspensions. During succinate oxidation, a minimum  $H^+/e^-$  stoichiometry of 2 is theoretically expected when ubiquinol is oxidized directly by cytochrome  $bb_3$  (Fig. 6). On the other hand, a ratio of  $3H^+/e^-$  is the theoretical maximum when ubiquinol is oxidized via cytochrome  $bc_1$ and cytochrome  $aa_3$  (Fig. 6).

Measurements of the set of P. denitrificans mutants are in perfect agreement with the theoretical values (Table 2). Succinate oxidation by cytochrome  $bb_3$  in the  $\Delta aa_3/cbb_3$ mutant results in 2 H<sup>+</sup>/e<sup>-</sup>. In the conditions used for analysis of proton translocation (0.4–1.7  $\mu$ M O<sub>2</sub>), ubiquinol appears to be oxidized preferentially via the cytochrome c branch ( $bc_1$  complex, cytochrome c, cytochrome c oxidase) rather than by the guinol oxidase cytochrome bb<sub>3</sub>. This observation may be due to the low affinity for oxygen that has been measured for cytochrome  $bb_3$  ( $K_m$ ) 10-20 µM O<sub>2</sub>; de Gier, 1995). This may explain why, in the  $\Delta cbb_3$  mutant, electrons mainly flow to cytochrome aa3, resulting in a stoichiometry of 3H<sup>+</sup>/e<sup>-</sup>. The proton pumping capacity of cytochrome cbb<sub>3</sub>, as first reported by Raitio and Wikström (1994) in a  $\Delta aa_3$  mutant, has been confirmed here with the  $\Delta aa_3/bb_3$  mutant during electron transfer from succinate to oxygen (Table 2).

The conclusion that cytochrome  $cbb_3$  does translocate protons, however, is in conflict with the previous measurements by de Gier *et al.* (1994). In the latter study an H<sup>+</sup>/e<sup>-</sup> stoichiometry of 2 has been measured during succinate oxidation of the  $\Delta aa_3/bb_3$  mutant, suggesting that cytochrome  $cbb_3$  does not pump protons. The main technical difference is that, in the experiment presented here, HEPES (0.5 mM) rather than glycylglycine (1.5 mM) has been used as buffer. The 'decoupling' of cytochrome  $cbb_3$  in glycylglycine (de Gier *et al.*, 1994) has been confirmed by analysis of cells from a single batch in both buffers (not shown). The apparent buffer sensitivity of cytochrome  $cbb_3$ , a phenomenon that is not observed in the case of cytochromes  $aa_3$  and  $bb_3$ , is not understood at present (see below).

The finding that all three terminal oxidases of Paracoccus ( $aa_3$ ,  $cbb_3$  and  $bb_3$ ) couple the reduction of oxygen to the translocation of protons with maximal efficiency (1 H<sup>+</sup>/e<sup>-</sup>) also does not agree with analyses of growth efficiency in chemostat cultures of (wild-type) P. denitrificans, grown under a variety of conditions. The maximal growth efficiency is observed only under conditions in which cytochrome aa<sub>3</sub> is expressed (reviewed by Stouthamer, 1991). To address the discrepancy between the latter physiological studies and the above-described proton-pumping capacity of cytochrome cbb<sub>3</sub>, we are currently cultivating a number of Paracoccus oxidase mutants under well-defined conditions. Preliminary data from a comparison of the growth yields of these mutants suggest that the energy-transducing efficiency of cytochrome cbb<sub>3</sub> is indeed lower than that of cytochrome aa<sub>3</sub> (de Gier, 1995).

The latter conclusion would be consistent with the aforementioned observation that the H<sup>+</sup>/e<sup>-</sup> stoichiometry of cytochrome *cbb*<sub>3</sub> is not a fixed number, and may be sensitive to, for example, buffer composition (de Gier *et al.*, 1994; Table 2). Moreover, proton pumping analysis of the  $\Delta aa_3/bb_3$  mutant with ascorbate/TMPD as substrate, either in HEPES or in glycylglycine, results in H<sup>+</sup>/e<sup>-</sup> values that never exceed 50% of the ratio that is measured in the presence of cytochrome  $aa_3$  (de Gier *et al.*, 1994; Raitio and Wikström, 1994). This apparent variability in the H<sup>+</sup>/e<sup>-</sup> ratio of cytochrome *cbb*<sub>3</sub> might be the result of certain structural deviations of subunit I within the regions that have been demonstrated to be important for the coupling between oxygen reduction and proton pumping (see below).

#### Residues involved in proton translocation

A major challenge of cytochrome oxidase research is elucidation of the proton-translocation mechanism at a

**Fig. 6.** Aerobic respiratory pathways in *P. denitrificans.* The proton-translocation ratios  $(H^+/e^-; protons per electron detectable in the medium) and the charge-separation ratios <math>(q^+/e^-)$  for each of the contributing respiratory complexes are shown. NADH-dehydrogenase is represented by NADH.dh, succinate dehydrogenase by suc.dh, cytochrome reductase by cyt. *bc*<sub>1</sub>, cytochrome *c*<sub>552</sub> by cyt. *c*<sub>552</sub>, *aa*<sub>3</sub>-type cytochrome *c* oxidase by cyt. *cbb*<sub>3</sub>, type quinol oxidase by cyt. *bb*<sub>3</sub>.



molecular level. This requires the identification of residues that are involved in the transfer of consumed (scalar) and/ or pumped (vectorial) protons. Among the highly conserved residues in subunits 1 of cytochrome aa3 and cytochrome bo<sub>3</sub>, a large number of polar residues have been substituted by means of site-directed mutagenesis (reviewed by Hosler et al., 1993). An aspartic acid in the domain between transmembrane helices II and III (loop II-III) has recently been proposed to participate in the influx of protons into the oxidase core (Garcia-Horsman et al., 1995; Fetter et al., 1995). Substitution of this conserved aspartate with an asparagine (D124N; numbering for P. denitrificans cytochrome aa3, CtaDII, is used throughout) in both cytochromes aa<sub>3</sub> and bo<sub>3</sub> results in decoupling of proton pumping from electron transfer. In addition, substitution of two conserved asparagine residues (N113, N131) results in a decrease in the H<sup>+</sup>/e<sup>-</sup> ratio. Interestingly, proton pumping is recovered in a double mutant of E. coli CyoB, D124N/N131D (Garcia-Horsman et al., 1995). It has been suggested that this cytoplasmic loop plays an important role as part of the entry to the pumped proton pathway, and moreover, that there should be a second channel, the chemical proton pathway. Residues that may be part of the latter pathway are a tyrosine (Y280) in helix VI, as well as two threonines (T344, T351) and a lysine (K354) in helix VIII. Non-polar substitutions for each of these residues result in a severe loss of oxidase activity (Hosler et al., 1993). Both T351 and K354 have been proposed to participate in a protonconducting channel towards the binuclear centre (Hosler et al., 1993; Fetter et al., 1995). Spectroscopic analyses suggest that T344 is in close proximity to the binuclear centre. Y280 may be a ligand to Cu<sub>B</sub>, at least during part of the catalytic cycle of oxygen reduction (Hosler et al., 1993). In addition, one of the invariant histidines that has been identified as a Cu<sub>B</sub> ligand has recently been proposed to play a key role in the proton-translocation machinery. In the 'histidine cycle' model, a Cu<sub>B</sub> co-ordinating histidine has been proposed to move back and forth in the binuclear 'pocket', thereby passing two protons from the inside to the outside (Morgan et al., 1994).

Analysis of the recently resolved structure of *P. denitrificans* cytochrome  $aa_3$  (lwata *et al.*, 1995) supported many of the conclusions that were based on previous studies of the *E. coli* and *R. sphaeroides* oxidases (Hosler *et al.*, 1993; Fetter *et al.*, 1995; Garcia-Horsman *et al.*, 1995). Indeed, two suggested proton channels have been envisaged in the structural model (lwata *et al.*, 1995). A pathway for consumed protons, from the cytoplasmic surface to the oxygen-binding site, appears to be made up of S291, K354, T351, the hydroxyl group of haem  $a_3$ , and, ultimately, Y280, which may donate a proton to oxygen (Fig. 3A, Table 3). With the exception of K354, all residues in this channel appear to be

connected by hydrogen bonds, either directly or via putative solvent molecules. A second series of hydrophilic residues may constitute the pathway for pumped protons. A gate is formed by D124, N199, and T203; subsequently, a number of hydrophilic residues, again with additional putative solvent molecules, may form a pathway to E278 (Table 3). Beyond E278 the proton pathway is less clear. However, as proposed before (Morgan et al., 1994), the structural model suggests that one of the histidine ligands, H325, may not be in a fixed position, but rather may switch between different conformational states. One option is that two subsequent protons, approaching the catalytic site via E278, convert the H325 imidazolate, via imidazole, into imidazolium. In the latter state, the positively charged H325 moves away from the binuclear centre, and no longer ligates Cu<sub>B</sub>. The two protons of H325 imidazolium would leave the system via the exit pathway, in which groups of the haem a<sub>3</sub> molecule and/ or residues in the periplasmic loop XI-X may be involved (Table 3), and H325 imidazole would return to its original position (for details, see Iwata et al., 1995).

As cytochrome  $cbb_3$  has the capacity to pump protons (Raitio and Wikström, 1994; Table 2), comparison of its sequence with that of the much better characterized cytochrome  $aa_3$  is of particular interest. From the alignment presented in Fig. 3A, it is obvious that the sequences from CcoN/FixN show some significant deviation from both the  $aa_3$ -type and the  $bo_3$ -type oxidases, especially

**Table 3.** Comparison of residues proposed to be involved in the chemical and the pumped proton channels in cytochrome  $aa_3$  (Hosler *et al.*, 1993; Fetter *et al.*, 1995; Iwata *et al.*, 1995), with counterparts in cytochrome  $cbb_3$ , based on the alignment presented in Fig. 3.

aa3	Location	cbb <sub>3</sub>	Consensus
Chemical channel			
a <sub>3</sub> -hvdroxvl	Haem	No hvdroxvl	_
Y280	TM-VI	G	
T351	TM-VIII	S	+
K354	TM-VIII	G	_
S291	TM-VI	М	
Pump channel			
D399	Loop-IX/X	N	+
L393-carbonyl	Loop-IX/X	М	+
a <sub>3</sub> -propionate	Haem	b <sub>3</sub> -propionate	+
a <sub>3</sub> -formyl	Haem	No formyl	_
H325	TM-VII	Η	+
E278	TM-VI	А	_
N113	TM-II	Х	
N131	Loop-11/111	Х	
N199	TM-IV	V	
T203	TM-IV	Т	+
D124	Loop-II/III	х	_

In transmembrane helix II (TM-II) and the interconnecting loop between helices II and III (loop II-III) a reliable alignment is not possible; in this case the *cbb*<sub>3</sub> residues are marked 'X'.

with respect to the residues that have been predicted to be part of either one of the two proton channels (Table 3). Although alignment of the oxidase sequences at their Ntermini is difficult, it is clear that the invariant aspartate residue in loop II-III of cytochrome aa<sub>3</sub> is not conserved in cytochrome cbb<sub>3</sub> (Table 3). In helix VI of cytochrome cbb<sub>3</sub>, the tyrosine (Y280) and the glutamate (E278) that are conserved in all aa3- and bo3-type oxidases (and appear to be essential in the aforementioned proton pump model) are substituted by a glycine and an alanine, respectively. In helix VIII, the threonines are conserved as serines, but a glycine is found at the position of the conserved lysine (Fig. 3A, Table 3). In conclusion, comparison of the primary structures suggests that many residues that have been proposed to constitute the chemical and pumped proton channels in cytochrome  $aa_3$  (and probably also in cytochromes  $bo_3$  and  $bb_3$ ) are not conserved in cytochrome cbb<sub>3</sub>. These proton-conducting pathways are undoubtedly an essential component of the proton-translocation machinery. Hence, it is concluded that the design of the proton pump in cytochrome cbb<sub>3</sub> differs significantly from that present in the other oxidase types. Although site-directed mutagenesis would provide more definitive evidence, it is tempting to assume that these deviations in cytochrome cbb3 might correlate with its apparently variable H<sup>+</sup>/e<sup>-</sup> coupling.

#### Experimental procedures

#### Bacterial strains, plasmids and growth conditions

The strains of *P. denitrificans* and *E. coli* as well as the plasmids used in this study are listed in Table 4. Cells of wildtype *P. denitrificans* (Pd1222) and mutants were cultivated in aerobic batch cultures (0.51 bottles with 100 ml of culture, on a rotary shaker at 30°C), with either brain-heart infusion (BHI) broth or minimal medium supplemented with 25 mM succinate, as described previously (van Spanning *et al.*, 1990). For microaerobic cultivation, cultures were not shaken. *E. coli* strains were cultivated in YT medium at 37°C. When appropriate, antibiotics were added: rifampicin (Rif, 40 mg1<sup>-1</sup>), streptomycin (Sm, 50 mg1<sup>-1</sup>), tetracycline (Tc, 12.5 mg1<sup>-1</sup>), kanamycin (Km, 50 mg1<sup>-1</sup>) and ampicillin (Amp, 100 mg1<sup>-1</sup>).

## Purification of the cbb<sub>3</sub>-type cytochrome c oxidase from P. denitrificans

Cells from succinate-grown Pd93.11 ( $\Delta aa_3/bb_3$ ) were harvested in the late log phase, and resuspended in 20 mM Bis-Tris propane (pH7.3), 2 mM EDTA and 0.5 mM PMSF. Membranes were prepared by passage of the cell suspension twice through a French Pressure Cell and subsequent centrifigation (1.5 h, 150 000 × g). Membranes were solubilized in 20 mM Bis-Tris propane (pH7.3), with 1.0% (w/v) dodecyl maltoside and 0.15 mM PMSF (30 min, 4°C). The suspension contained approx. 10 mg protein per mI and was cleared by ultracentrifugation (30 min,  $150\,000 \times g$ ). The solubilized membrane proteins were applied to a Q-Sepharose HP (Pharmacia) column in 20 mM Bis-Tris propane (pH 7.3), with 0.03% (w/v) dodecyl maltoside, and bound protein was eluted with a linear NaCl gradient (0.0-1.0 M). Oxidase activity was measured spectrophotometrically with TMPD as described previously (Lübben et al., 1994). Fractions with cytochrome c oxidase activity eluted at 0.3 M NaCl. Pooled fractions were applied onto a sucrose gradient, as described previously (Gray et al., 1994). Fractions with TMPD oxidase activity were applied to a Chelating-Sepharose FF (Pharmacia) column, saturated with Cu2+. A linear ammonium chloride gradient (0.0-1.0 M) in 20 mM Tris (pH8.0), 0.5 M NaCl, 15 mM PMSF and 0.03% (w/v) dodecyl maltoside was used to elute bound proteins. Cytochrome c oxidase activity eluted at 0.3 M ammonium chloride. As a last purification step the active fractions were loaded onto a Q-Sepharose HP column (Pharmacia) in 20 mM Bis-Tris propane (pH 7.3), with 0.03% (w/v) dodecyl maltoside, and bound protein was

Table 4. Strains and plasmids.

Strain/Plasmid	Relevant characteristics	Reference/Source
Strain		
E. coli		
TG1	supE hsdD5 thi Δ(lac⁻proAB) F′ (traD36 proAB lacl° lacZΔM15)	Sambrook <i>et al.</i> (1989)
HB101	F <sup>-</sup> hsdS20 (r <sub>B</sub> m <sub>B</sub> ) lacY1 proA2 recA13	Boyer <i>et al.</i> (1969)
S17.1	Sm <sup>R</sup> pro (r <sub>B</sub> <sup>-</sup> m <sup>+</sup> <sub>B</sub> ) RP4-2, integrated (Tc::Mu) (Km::Tn7)	Simon <i>et al.</i> (1983)
P. denitrificans		
Pd1222	Rif <sup>R</sup> , enhanced conjungation frequencies, (ramt)	de Vries <i>et al.</i> (1989)
Pd92.20	Pd9218 derivative, ΔctaDl ΔctaDl	de Gier <i>et al.</i> (1994)
Pd26.21	Pd1222 derivative, <i>goxB</i> ::Km <sup>R</sup>	de Gier <i>et al.</i> (1994)
Pd93.11	Pd9220 derivative, Δ <i>ctaDl</i> Δ <i>ctaDll cvoB</i> ::Km <sup>R</sup>	de Gier <i>et al.</i> (1994)
Pd27.21	Pd1222 derivative, ccoNO::Km <sup>R</sup>	This work
Pd93.12	Pd9220 derivative, Δ <i>ctaDI</i> Δ <i>ctaDII ccoNO</i> ::Km <sup>R</sup>	This work
Plasmid		
pUC19	Amp <sup>R</sup> <i>lacZ′</i>	Yanisch-Perron et al. (1985)
pUC4KIXX	Km <sup>R</sup> (Tn5)	Pharmacia
M13mp18/19	pUC18/19 mcs, lacZ'	Sanger <i>et al.</i> (1980)
pRK2020	Tc <sup>R</sup> pRK2013 Km::Tn <i>10</i>	Ditta et al. (1985)
pGRPd1	oriV (colE1) Amp <sup>R</sup> oriT Sm <sup>R</sup> (Tn1831)	van Spanning et al. (1990)
pCco1	pUC19 derivative, ccoNOQP	This work
pCco2	pUC19 derivative, ccoNAorfN	This work
pCco3	pUC19 derivative, ccoNOQPGH'	This work
pCcoNO::Km <sup>R</sup>	pCco1 derivative, ccoNO::Km <sup>R</sup>	This work
pRTd27.21	pGRPd1 derivative, ccoNO::Km <sup>R</sup>	This work

eluted with a linear NaCl gradient (0.0-1.0 M). Active fractions eluted at 0.3 M NaCl. The purified oxidase was analysed by SDS–PAGE and spectrophotometrically (Laemmli, 1970; de Gier *et al.*, 1994). Haem staining was performed as described by Thomas *et al.* (1976).

#### DNA manipulation

General cloning techniques were carried out as described by Ausubel *et al.* (1993). Conjugations were performed as described previously (de Vries *et al.*, 1989). The matings of *Paracoccus* host strains were performed either directly with an *E. coli* S17.1 derivative that carried the plasmid of interest or via a triparental mating using any *E. coli* strain transformed with the plasmid of interest, in combination with *E. coli* HB101(pRK2020) containing the 'helper plasmid'. Mutagenesis was performed using pGRPd1 as described by van Spanning *et al.* (1990).

The PCR was performed using Super-Taq polymerase (HT Biotechnology Ltd.), essentially as described by de Gier *et al.* (1994). Primers were synthesized according to the amino acid sequence of the N-terminal peptide of the 30 kDa subunit of the *Paracoccus cbb*<sub>3</sub> complex (CcoO) (nos 171 and 172). These were degenerate primers with a bias for G:C at the third position of a codon. The sequences of these primers are as follows: 171 (sense), 5'-AAAAAAGTCTGCNAT[T/C/A]CTNGA[A/G]CA[T/C]-3'; and 172 (antisense), 5'-AAAA-GAATTC[A/G/T]ATNCGNGNAACNACNAG[A/G]AA-3'.

DNA sequencing was performed using the dideoxy method (Sanger *et al.*, 1977) on fragments subcloned into M13mp18/ 19 (Sanger *et al.*, 1980) with the *Taq* dye dideoxy terminator cycle sequencing kit, and the *Taq* dye primer cycle sequencing kit (Applied Biosystems). Sequence analysis was carried out on a 370A DNA sequencer (Applied Biosystems). Editing of the sequences was performed with the SEQUENCHER 2.1 Program (Gene Codes Corp.).

#### Oxygen-consumption analysis

Oxygen consumption by bacterial cell suspensions was performed polarographically with a Clark-type oxygen electrode, as described by de Gier *et al.* (1994).

#### Proton translocation

Proton translocation was studied by using the oxygen-pulse method in intact cell suspensions (de Gier *et al.*, 1994; Raitio and Wikström, 1994). Cells were harvested in the late log phase from succinate batch cultures, washed twice and resuspended in 150 mM KCl (final OD<sub>660</sub> is 100), and diluted four times in reaction medium (pH7.4) in the stirred anaerobic reaction vessel (3.0 ml). The buffer contained 100 mM KCl, 100 mM KSCN, 0.5 mM HEPES (pH7.4) and 30  $\mu$ M rotenon, as described by Raitio and Wikström (1994). Succinate was used as the electron donor (final concentration 2.5 mM). The reactions were started with small pulses (5–20  $\mu$ l) of air-saturated water (containing 258  $\mu$ M O<sub>2</sub> at 25°C) or 1.0 mM potassium ferricyanide, made anaerobic with argon. Pulses were calibrated with 1.0 mM HCl or 0.5 mM oxalic acid, made anaerobic with argon. Controls with the protonophore FCCP were

routinely included to ascertain if the observed proton ejection was due to proton translocation.

#### Acknowledgements

We thank M. Wikström and H. V. Westerhoff for critical reading of the manuscript. This research was supported by the Netherlands Foundation for Chemical Research, with financial aid from the Netherlands Organization for the Advancement of Science. J.v.d.O. had a Fellowship of the Royal Netherlands Academy of Science. EMBO supported visits of J.W.d.G. to the laboratories of M. Wikström and M. Saraste.

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