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The crystal structure of heme d_1 biosynthesis-associated small ctype cytochrome NirC reveals mixed oligomeric states *in crystallo*

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Synopsis The crystal structure of the c-type cytochrome NirC from *Pseudomonas aeruginosa* has been determined and reveals the simultaneous presence of monomers and 3D domain-swapped dimers in the same asymmetric unit.

Abstract C-type cytochromes are important electron transporters in all domains of life. They possess a common fold hallmarked by three α -helices that surround a covalently attached heme. An intriguing feature of many c-type cytochromes is their capacity to form oligomers by exchanging at least one of their α -helices, which is often referred to as 3D domain swapping. Here, we have determined the crystal structure of NirC, a c-type cytochrome co-encoded with other proteins involved in nitrite reduction by the opportunistic pathogen *Pseudomonas aeruginosa*. Crystals diffracted anisotropically to a maximum resolution of 2.12 Å (spherical resolution 2.83 Å) and initial phases were obtained by Fe-SAD phasing, revealing the presence of eleven NirC chains in the asymmetric unit. Surprisingly, these protomers arrange into one monomer and two different types of 3D domain-swapped dimers, one showing pronounced asymmetry. While the simultaneous observation of monomers and dimers probably reflects the interplay between high protein concentration required for crystallization and the structural plasticity of c-type cytochromes, the identification of conserved structural motifs in the monomer together with a comparison to similar proteins may offer new leads to unravel the unknown function of NirC.

1. Introduction

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Acta Crystallographica Section D

research papers

C-type cytochromes are well-known electron transporters in the respiratory chain of organisms in all domains of life. In bacteria, they are also involved in H_2O_2 -scavenging and work as electron entry point for the nitrate-, nitrite-, and nitric oxide reductases, which function as terminal oxidases in the respiratory chain under anaerobic conditions (denitrification) (Bertini et al., 2006). The monoheme cytochrome c domain features three α -helices that surround one c-type heme covalently attached to the cysteines of the characteristic sequence motif CXXCH via thioether bonds. The central iron cation of heme is always coordinated by a histidine residue at its proximal site, whereas a methionine ligates the other side in most cases. Many c-type cytochromes can form dimers and oligomers by exchanging the N- or C-terminal α -helix, which offers attractive protein engineering opportunities (Nagao *et al.*, 2015; Hayashi et al., 2012; Hirota et al., 2010). We have recently embarked on a structural biology program to investigate enzymes involved in the biosynthesis of heme d_i , an essential cofactor of the cd_1 nitrite reductase NirS, which catalyses the reduction of nitrite to nitric oxide in e.g. the opportunistic pathogen Pseudomonas aeruginosa (Schobert & Jahn, 2010). The nir-operon of P. *aeruginosa*, which encodes most of the enzymes involved in heme d_1 biosynthesis in addition to *nirS* itself, also contains open reading frames for two c-type cytochromes termed *nirM* and *nirC* (Figure 1) (Arai et al., 1990; Kawasaki et al., 1995). Whereas NirM is absent in many NirS-producing bacteria and has been assigned to function as an electron transporter for the nitrite reductase (Hasegawa et al., 2001), orthologs of NirC are found in nearly all NirS producers, but its function is subject to ongoing discussions. It has been demonstrated that NirC can act as an electron donor for the nitrite reductase, but its redox potential is not optimal (Zajicek et al., 2009; Hasegawa et al., 2001) and other electron transporters such as cytochrome c550 or pseudoazurin are better suited for this function in vivo (Pearson et al., 2003). Interestingly, mutants of of Paraccocus denitrificans and Pseudomonas *fluorescens* impaired in their ability to express *nirC* are unable to produce active nitrite reductase due to the lack of heme d_1 , implicating a role in heme d_1 biosynthesis (Boer et al., 1994; Ye et al., 1992), which could, however, not be confirmed in *P. aeruginosa* (Hasegawa et al., 2001). To further investigate NirC, we have determined the crystal structure of heterologously produced P. aeruginosa NirC. We find a unique crystal packing that contains monomeric NirC together with two different dimeric forms. To our knowledge, this is the first time that monomers and dimers of a c-type cytochrome have simultaneously been observed in the same asymmetric unit.

Acta Crystallographica Section D

research papers

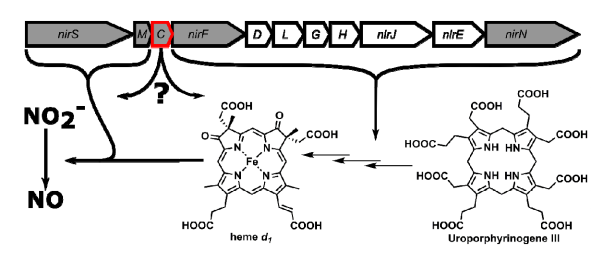


Figure 1 Depiction of the *nir*-operon indicating the function of each gene product in nitrite reduction and heme d_1 biosynthesis. A grey background highlights the periplasmic location of the gene product.

2. Materials and methods

2.1. Macromolecule production

The gene encoding NirC was amplified by polymerase chain reaction from genomic DNA extracted from *P. aeruginosa* PA14, omitting the signal peptide for periplasmic export as determined by SignalP 5.0 (Almagro Armenteros *et al.*, 2019). The fragment was cloned between the restriction sites NotI/KpnI of a modified pCOLA-Duet plasmid (pVP008-ompA), yielding an expression vector that produces NirC with an N-terminal ompA signal peptide for periplasmic export in *E. coli* followed by a Tobacco Etch Virus (TEV) protease-cleavable StrepII-tag. Residue E71 was mutated to alanine by QuikChange Mutagenesis for reasons outlined below. The success of cloning and mutagenesis was verified by the Eurofins Genomic sequencing service.

E. coli C43(DE3) (Miroux & Walker, 1996) was cotransformed with the NirC expression plasmid and with pEC86, a plasmid containing the cytochrome *c* maturation system of *E. coli*, which is necessary for the covalent attachment of heme to a suited polypetide chain (Arslan *et al.*, 1998). LB medium (8 x 1 l) supplemented with 34 mg/l chloramphenicol, 30 mg/l kanamycin, 200 μ M δ -aminolevulinic acid and 12.5 μ M Fe(II)SO₄ was inoculated with 10 ml overnight culture, cultivated at 37 °C with mild shaking till an optical density between 0.6 and 0.8 was reached. Cultures were then induced with 1 mM isopropyl β -D-1-thiogalactopyranoside and further incubated at 20 °C for 20 h. Cells were harvested by centrifugation and pellets were stored at -20 °C until needed.

Cell pellets were thawed, resuspended in lysis buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl) supplemented with protease inhibitors (cOmplete mini EDTA-free) and lysed by sonication. The lysate was centrifuged 30 min at 36,000 g followed by an additional centrifugation of the supernatant for 60 min at 100,000 g to remove cell debris.

Acta Crystallographica Section D

research papers

The cleared cell lysate was loaded on a StrepTactin-HC column (IBA, Göttingen, Germany) equilibrated in protein buffer (10 mM Tris-HCl pH 8.0, 150 mM NaCl) and connected to an Äkta Purifier FPLC system (GE Healthcare, Boston, U.S.A.). The column was washed with five column volumes and subsequently eluted with three column volumes of protein buffer containing 5 mM d-desthiobiotin. Red colouration of elution fractions and UV-Vis absorption at 410 nm confirmed the presence of heme. Fractions containing no impurities as judged by SDS-PAGE analysis were pooled and digested with TEV-protease at 4 °C overnight with simultaneous dialysis against protein buffer. To remove undigested protein, the resulting solution was again loaded onto the StrepTactin column and the flowthrough was collected. As final purification step, NirC was subjected to size exclusion chromatography using a HiLoad 16/60 Superdex 75 column (GE Healthcare, Boston, U.S.A.). Pure protein was concentrated to 45 mg/ml with Vivaspin 20 (MWCO 3000) ultrafiltration units (Sartorius AG, Göttingen, Germany) and concentration was determined photometrically, using a NanoDrop2000 (ThermoFisher Scientific, Waltham, U.S.A.).

Source organism	Pseudomonas aeruginosa
Cloning Primer	AAA <u>GCGGCCGC</u> GACGAGCATCCCGATGCCC
6	AAA <u>GGTACC</u> TCATGGGGCGATCTCTCCTTCG
Mutagenesis Primer	GCCTGCTCAGCGCAGACGACGCCGG
	CCGGCGTCGTCTGCGCTGAGCAGGC
Cloning vector	pVP008-ompA modified from p-COLA-Duet
Expression vector	pVP008-ompA-NirC-E71A
Expression host	Escherichia coli C43
Complete amino	<i>MKKTAIAIAVALAGFATVAQ</i> AMAS WSHPQFEK VDENLYF <u>QG</u> GGRDEH
acid sequence of the	PDARRQAQLRHLLLQDCGSCHGLRLTGGLGPALTPEALRGKPRESLV
recombinant protein	ATVLMGRPQTPMPPWAGLLSADDAGWLVDRLIEGEIAP

TEV-protease cleavage site or NotI/KpnI restriction sides underlined; StrepII-tag in bold, ompA leader peptide in italics. E75A mutation as red bold letter.

2.2. Crystallization

Crystallization experiments were set up using a HoneyBee 961 pipetting robot (Digilab Genomic Solutions, Hopkinton, U.S.A.) and monitored with a Rock Imager 1000 automated microscope (Formulatrix, Bedford, U.S.A.). Optimization screens were prepared utilizing a Formulator pipetting robot (Formulatrix, Bedford, U.S.A.). Experiments to crystallize wild type NirC never yielded well-diffracting crystals. Therefore, a mutation (E71A) as suggested by the surface entropy reduction server (Goldschmidt *et al.*, 2007) was introduced, leading to crystal growth in a few conditions of the

Acta Crystallographica Section D

Cryos sparse matrix screen (Qiagen, Hilden, Germany). The respective precipitants were further optimized by varying the concentration of precipitant and salts as well as the pH. Crystals usually began to appear after one week and took three more weeks to grow to full size (50-100 μ m) (Supplementary Figure 1). Crystals were fished and directly flash-cooled in liquid nitrogen without the addition of cryoprotectants.

Table 2Crystallization

	Crystal used for SAD phasing	Crystal used for final structure refinement
Method	Sitting drop vapour diffusion	Sitting drop vapour diffusion
Plate type	Intelli 96-3	Intelli 96-3
Temperature (K)	293	293
Protein concentration (mg/ml)	27	40
Buffer composition of	10 mM Tris-HCl pH 8.0	10 mM Tris-HCl pH 8.0, 150 mM
protein solution	150 mM NaCl	NaCl
	15% glycerol	15.33% glycerol
Composition of reservoir	8.5 mM NiCl ₂	0.1 M 2-(N-
solution	85 mM Tris pH 8.5	morpholino)ethanesulfonic acid
501401011	17% Poly(ethylene) glycol	рН 6.3
	monomethyl ethers 2000	10.8% Poly(ethylene) glycol 20000
Volume and ratio of drop	200 nl : 200 nl	200 nl : 200 nl
Volume of reservoir	60 µl	60 µl

2.3. Data collection and processing

Data were collected at beamline P11 of the PETRAIII synchrotron (DESY, Hamburg, Germany) (Burkhardt *et al.*, 2016). The anomalous data set was processed using DIALS (Winter *et al.*, 2018), POINTLESS (Evans, 2011) and AIMLESS (Evans & Murshudov, 2013) of the CCP4 suite (Winn *et al.*, 2011). Despite low completeness of the highest resolution shell the data set was sufficient to obtain initial phases via SAD phasing. The data set used for final refinement was processed using autoPROC (Vonrhein *et al.*, 2011) executing XDS (Kabsch, 2010), POINTLESS (Evans, 2011), AIMLESS (Evans & Murshudov, 2013) and STARANISO (Tickle *et al.*, 2018). Anisotropic data

Acta Crystallographica Section D

research papers

truncation was applied assuming a local $I/\sigma(I)$ of 1.2 include useable data beyond the spherical resolution of 2.83 Å.

Table 3 Data collection and processing

Values for the outer shell are given in parentheses.

	Anomalous Dataset	Native dataset
Diffraction source	Petra III beamline P11	Petra III beamline P11
Wavelength (Å)	1.738	1.033
Temperature (K)	100K	100K
Detector	Pilatus 6M	Pilatus 6M
Crystal-detector distance (mm)	544	457.5
Rotation range per image (°)	0.1	0.1
Total rotation range (°)	1080	360
Exposure time per image (s)	0.1	0.04
Space group	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁
<i>a</i> , <i>b</i> , <i>c</i> (Å)	77.7, 80.6, 200.0	76.8, 80.2, 195.7
α, β, γ (°)	90, 90, 90	90, 90, 90
Resolution range (Å)	199.97-3.42	spherical:99.16-2.83
Diffraction limit along principal axes of fitted ellipsoid	-	2.98 (a) 2.91 (b) 2.12 (c)
Total No. of reflections	368556 (6018)	460561 (18799)
No. of unique reflections	13722 (873)	35413 (1772)
Completeness spherical (%)	77.3 (24.9)	53.6 (8.9)
Completeness ellipsoid (%)	-	92.7 (74.8)
Multiplicity	26.9 (6.9)	13.0 (10.6)
<i>〈 Ι/</i> σ(<i>I</i>) <i>〉</i>	10.5 (2.2)	10.9 (1.8)
R _{p.i.m.}	0.046 (0.404)	0.058 (0.416)

Acta Crystallographica Section D)		research papers
Overall <i>B</i> factor from Wilson plot (\AA^2)	63	23	

2.4. Structure solution and refinement

The structure was solved by SAD phasing with anomalous differences caused by the heme iron atoms, using the phasing pipeline Crank2 (Skubák & Pannu, 2013) executing ShelxC/D (Sheldrick, 2008) for substructure determination, Parrot (Cowtan, 2010) for density modification and Buccaneer (Cowtan, 2006) for model building. The iron substructure determination showed an occupancy drop for the 12^{th} atom and beyond, hinting at the presence of eleven chains in the asymmetric unit. Automated model building produced a model consisting of 940 amino acids and having an R_{work} of 35%. Inspection of the electron density showed that not all of the eleven chains were built correctly, requiring manual adjustments in coot (Emsley & Cowtan, 2004) and further refinement in REFMAC5 (Murshudov *et al.*, 2011) with enabled NCS restraints. Final refinement was performed against the higher resolution native data set, using Phenix.refine (Afonine *et al.*, 2012) while lifting the applied NCS restraints, adding water molecules and applying TLS refinement. Depictions of the structure were made with PyMol (Schrödinger, 2015).

Table 4 Structure solution and refinement

No. of reflections, working set	35402 (172)
No. of reflections, test set	1746 (12)
Final R _{cryst}	0.211 (0.391)
Final <i>R</i> _{free}	0.247 (0.539)
No. of non-H atoms	
Protein	6879
Ligand	473
Water	154
Total	7506
R.m.s. deviations	
Bonds (Å)	0.004
Angles (°)	0.71

Values for the outer shell are given in parentheses.

Acta Crystallographica Section D		research paper
Average <i>B</i> factors (Å ²)	32	
Protein	33	
Ligand	27	
Water	29	
Ramachandran plot		
Most favoured (%)	97.44	
Allowed (%)	2.56	

2.5. Size exclusion chromatography coupled to an multi angle laser light detector

To assess the molecular weight of the isolated protein, we performed SEC-MALS experiments with an Agilent Technologies 1260 Infinity II HPLC system (Santa Clara, U.S.A.) equipped with a Wyatt Optilab rEX diffraction index detector and a Treos II multi angle laser light scattering detector (Wyatt, Santa Barbara, U.S.A.). Separation was achieved with a Superdex 75 Increase 10/300 GL column (GE Healthcare, Boston, U.S.A.) with protein buffer as eluent. For calculation of the molecular mass, a dn/dc of 0.195 ml/g was assumed.

2.6. Structural Bioinformatics

Functional orthologs of NirC were identified by extracting orthologs from the OMA database (Altenhoff *et al.*, 2018) and followed by manually filtering for the presence of the genes encoding nitrite reductase NirS and heme d_1 biosynthesis enzyme NirF in their genetic neighbourhood. After aligning the functional orthologs with Clustal Ω , sequence conservation was mapped onto the NirC monomer structure utilizing the ConSurf server (Figure 3 and Supplement (Figure 2) (Ashkenazy *et al.*, 2016) (Sievers *et al.*, 2011). Structural homologs were identified with the DALI server (Holm & Laakso, 2016).

3. Results and discussion

3.1. Simultaneous presence of NirC monomers and dimers in the asymmetric unit

Approximately 2 mg of pure heme-bound NirC per litre of *E. coli* culture could reliably be obtained with the procedure described above. Size exclusion chromatography as well as multi-angle light scattering suggested that the purified protein is monomeric with a molecular mass of 11.5 kDa (\pm 0.9%, 10.5 kDa expected; Fig. 2). Because the wildtype protein failed to crystallize, we attempted to reduce the surface entropy by mutating E71 to alanine, finally yielding cube-shaped crystals that diffracted anisotropically and possessed a very large orthorhombic unit cell (Supplementary Figure 1).

Acta Crystallographica Section D

research papers

To obtain initial phases, we made use of the presence of anomalous differences in diffraction data collected at the iron K-edge, leading to the positioning of eleven iron atoms. By assuming that NirC contains one iron atom bound to the covalently attached heme, this suggests a solvent content of 58%. A combination of automated model building and manual adjustments led to a final model with R-factors $R_{work} = 21.1\%$ and $R_{free} = 24.7\%$, indeed consisting of eleven chains in the asymmetric unit. Interestingly, despite using monomeric NirC for crystallization, ten of these chains form dimers by exchanging their N-terminal α -helices recognizable by different traces in the 2Fo-Fc density for the "hinge loop" (Figure 5). The dimers fall into two different types as outlined below. Surprisingly, the eleventh chain is monomeric, i.e. the crystal form obtained here contains both dimers and monomers of NirC at the same time.

The formation of dimers from monomeric c-type cytochromes is a well-known phenomenon (Hirota, 2019). It has been shown that the folding of horse cytochrome *c* starts with hydrophobic interactions between the N- and C-terminal helix, yielding 3D-domain swapped oligomers if the helices belong to different chains (Parui *et al.*, 2013). Therefore, denaturation and subsequent refolding at high protein concentration is often utilized to produce such dimers (Nagao *et al.*, 2015; Hayashi *et al.*, 2012). This indicates that the folding landscape of c-type cytochromes contains at least two stable states. The high NirC concentration and the presence of precipitants in the crystallization experiments performed here is apparently sufficient to allow an equilibrium between both states.

Acta Crystallographica Section D

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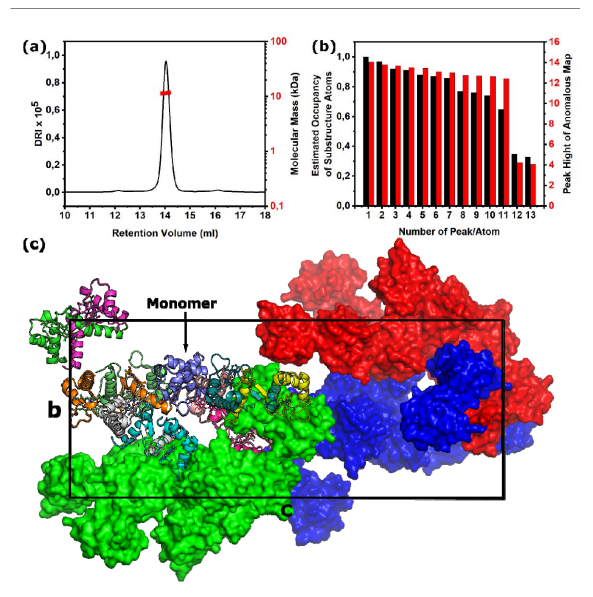


Figure 2 Panel (a) shows the SEC-MALS chromatogram. Bar diagrams in (b) show the occupancy of atoms of the initially determined substructure as calculated with ShelxD (Sheldrick, 2008) and the height of the anomalous map peak calculated with phases of the final structure by ANODE (Thorn & Sheldrick, 2011). The drop after the 11th site indicates that only eleven iron atoms are present. Panel (c) depicts the unit cell. One asymmetric unit is shown as in cartoon representation of the peptide backbone with the covalently attached heme as ball-and-stick model and each chain coloured individually. The other asymmetric units are depicted as surface model coloured in red, blue and green.

3.2. NirC shows high similarity to other c-type cytochrome domains of the nir-operon

Acta Crystallographica Section D

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NirC displays a typical c-type cytochrome fold containing of three σ -helices ($\alpha 1, \alpha 3, \alpha 4$) that surround a thioether-bound heme c moiety (Fig. 3). The iron cation is coordinated by H24 and M62, which is consistent with mutagenesis studies performed with the orthologue from Paracoccus panthotrophus (Zajicek et al., 2009). The heme-binding crevice consists of a predominantly hydrophobic pocket established by L32, P34, L36, T52, V53, T60, P61 and W65, which are also highly conserved in NirC proteins from other species. In P. aeruginosa NirC, the propionate groups of heme form H-bonds with K44 and R57, two residues that are replaced with hydrophobic amino acids in some species, probably reflecting the fact that the heme propionate groups are solvent-exposed and do not require stabilizing interactions with the protein to be accommodated. Sequence alignment reveals that in NirC the heme attachment motif (CXXCH) is always led by a conserved aspartate (D19), whose sidechain is solvent exposed, and followed by a conserved GGLG motif, residing on a loop connecting helix $\alpha 2$ and $\alpha 3$ (Supplementary Figure 3). Together with residues in the neighbourhood of the iron-ligating M62, these motifs create a conserved surface patch, which may be required to form an interaction site for other proteins such as e.g. the heme d_1 biosynthesis protein NirF, whose function is not clear but may lie in an oxidoreduction within the last steps of the pathway (NirF-paper). Alternative hypotheses suggest an involvement as electron transporter for nitrite reductase NirS, although, as outlined above, its redox potential is not optimal for this purpose (Zajicek et al., 2009; Hasegawa et al., 2001). A role in heme d_1 biosynthesis itself, on the other hand, seems to be supported by the finding that similarity searchers with DALI (Holm & Laakso, 2016) identify the cytochrome c domains of the nir-operon proteins nitrite reductase NirS and heme d_1 biosynthesis enzyme NirN as most similar to NirC, while NirM and cyt c552, the known electron transporters that interact with NirS in nitrite reduction, differ to a larger extent (Figure 4 and supporting Table S1). In NirN and NirS, the cytochrome *c*-domains are covalently attached to a β -propeller domain and function as electron shuttles in the reactions catalysed by these enzymes. This suggests that NirC may perform a similar function with NirF, which is a standalone β -propeller protein with high similarity to the β -propeller domains of NirN and NirS. Interestingly, superposition of NirC and NirF onto the respective domains of NirS leads to a complex in which the conserved surface patch of NirC is oriented towards the propeller of NirF and L32 of the conserved GGLG motif is buried in a hydrophobic pocket (Supplementary Figure 4). This may further corroborate NirC's involvement in heme d_1 biosynthesis.

Acta Crystallographica Section D

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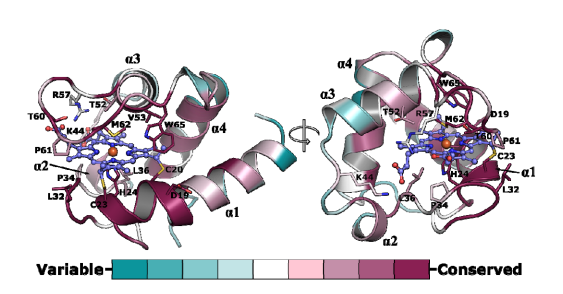


Figure 3 The NirC monomer shown as a cartoon with residues at a maximum distance of 4 Å to the covalently attached heme presented as sticks. Colours are based on sequence conservation as defined by Consurf (Ashkenazy *et al.*, 2016). The sequence alignment used for determining conservation is shown in Supplementary Figure 3.

Acta Crystallographica Section D

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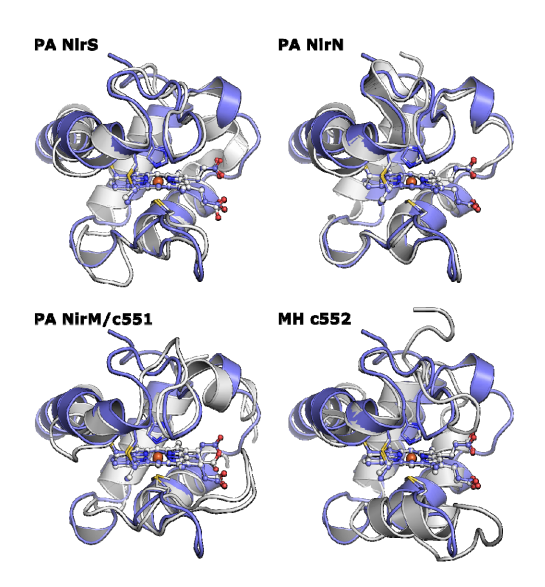


Figure 4 Comparison of the structure of monomeric NirC with the cytochrome *c* domains of NirS and NirN as well as with NirM from *Pseudomonas aeruginosa* or *cyt c552* of *Marinobacter hydrocarbonoclasticus* after superposition of the covalently attached heme. All structures are depicted as cartoons and the covalently attached heme as a ball-and-stick model. NirC is shown in blue and the compared structure in white.

3.3. NirC forms two different types of dimers

The dimers of NirC found here form by exchanging the N-terminal helix (P4-H24), which results in heme being coordinated by H24 and M62 coming from two different chains, similar to what has been observed for dimers of cyt c551 (NirM) from *P. aeruginosa* and cyt c552 from *Hydrogenobacter thermophiles* (HT) (Hayashi *et al.*, 2012)(Nagao et al., 2015). Notably, dimerization does not have a

Acta Crystallographica Section D

research papers

major impact on the structure of the heme binding site, indicating that monomers and dimers could display similar biochemical properties as was already found for other N-terminally domain-swapped cytochromes (Nagao *et al.*, 2015; Hayashi *et al.*, 2012).

Closer inspection reveals that the five dimers present in the asymmetric unit fall into two groups distinguishable by heme iron distance, global symmetry as quantified by the GloA-score (Swapna et al., 2012), and relative position of the hinge loop (25-GLRLTGGLG-33), which produced welldefined electron density both dimeric and the monomeric state of NirC (Figure 5). In the three "fardimers", the average iron-to iron distance is 28.5 ± 0.2 Å with a GloA-score of 0.9 ± 0.2 Å, and L26, R27 and L28 of the hinge loop interacting in an anti-parallel β-sheet-like fashion. The two "closedimers" have an iron-to-iron distance of 24.9 ± 0.1 Å, a GloA-score of 3.05 ± 0.01 Å and no interactions between the hinge-loops. The extent of this asymmetry is readily discernible after superposition of one promoter of a dimer onto the other (Supplementary Figure 2). Naturally occurring homodimers are usually highly symmetric with GloA-scores under 0.4 Å, and scores above 3 Å are indicative of substantial asymmetry, which is often associated with oligomers that bind other asymmetric macromolecules (Swapna et al., 2012). The basis for the formation of different dimers in NirC seems to be rooted in the hinge loop, which with nine residues is much longer and allows more flexibility than the three residues found in PA cyt c551 or HT cyt c552 (Hayashi et al., 2012; Nagao et al., 2015). This is further supported by the observation of different domain-swapped dimers after elongation of the hinge loop in HT cyt c552 (Ren et al., 2015), although this led to a mixture of Nand C-terminally exchanged dimers.

The relevance of dimer formation in NirC and in particular the observation of asymmetry is not clear at present. Previous work with eukaryotic c-type cytochromes suggests that 3D-domain swapped oligomers are involved in apoptosis (Junedi *et al.*, 2014), but their importance in prokaryotes is not known (Hirota, 2019). We therefore assume that our findings reflect the artificially high concentrations used in crystallization experiments together with the structural plasticity of these proteins rather than physiologically significant states.

In summary, our work provides new insight into NirC, a c-type cytochrome likely involved in the biosynthesis of heme d_1 , the essential cofactor in nitrite reduction by denitrifying bacteria such as the opportunistic pathogen *Pseudomonas aeruginosa*. While the simultaneous observation of NirC monomers and dimers in the same asymmetric unit of the crystal form obtained here may reiterate typical properties of this protein family, the mapping of conserved sequence motives onto the structure of the monomer may offer new avenues to decipher the function of the NirC.

Acta Crystallographica Section D

research papers

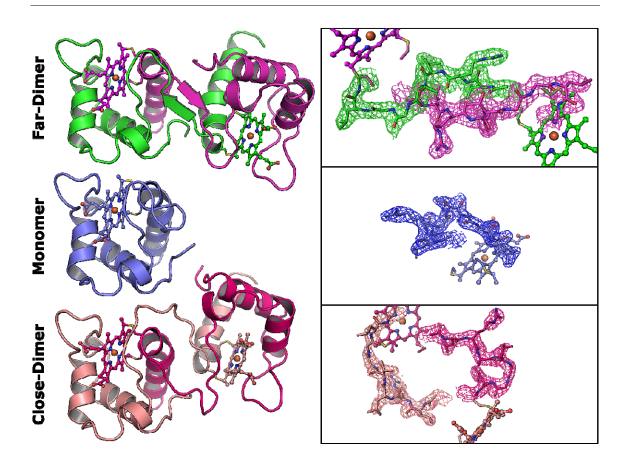


Figure 5 Depiction of the two different oligomerization states found in the crystal structure of NirC. On the left the NirC monomer and both dimer conformations are shown as described in Figure 1 after superposition of one protomer. The backbone of the NirC is shown as a cartoon with the covalently attached heme as ball-and-stick model. On the right, residues of the hinge loop are depicted as stick models with a $2F_oF_c$ map at a σ -level of one, coloured according to the associated chain and individually orientated to allow unobstructed view.

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Acta Crystallographica Section D

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Supporting information

Organism and	PDB-ID	Z-score	rmsd	%id	Citation
Name					
PA NirN	6rtd	12.2	1.6	37	(Klünemann <i>et</i> <i>al.</i> , 2019)
PA NirS	1nno	12	2.3	26	(Nurizzo <i>et al.</i> , 1998)
PA NirM/ cyt c551	351c	7.8	2.5	25	(Matsuura <i>et</i> <i>al.</i> , 1982)
MH cyt c552	1cno	5.2	2.7	23	(Brown <i>et al.</i> , 1999)

Table S1Selected high scoring results of the Dali server discussed in this publication.

Acta Crystallographica Section D

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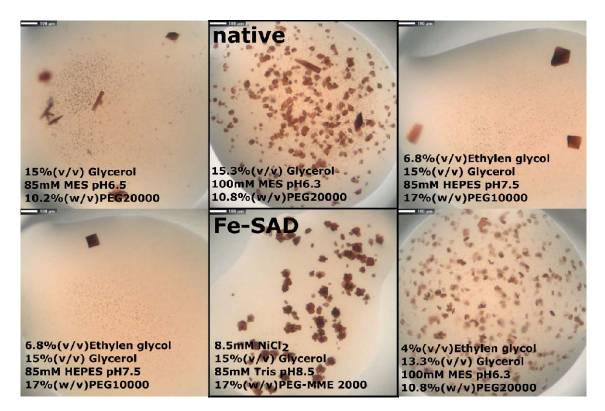


Figure S1 Crystals of NirC^{E71A} grown in different conditions used for diffraction experiments.

Conditions used in this study are indicated by a black frame.

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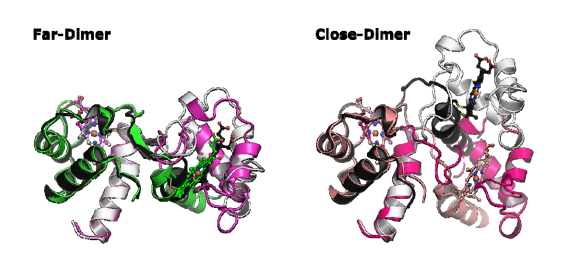


Figure S2 Depiction of NirC dimers. To highlight the asymmetry, one protomer was superposed with the copy of the opposite one. Except for the colouration of the copied dimer (black and white), the colour and representation of the model is the same as lied out in Figure 5 in the main text.

Acta Crystallographica Section D

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PERABOOSSO	LACEN	DAIRCAOLRHIL	LODCGSCHGLT	LTGGLGPALTPEALRG LTGGLGPALTPEALRG
PSEA700617	LACEH?	DARRCAQLRHIL	FODCeQCHC_S	LTGGLGPALT?EALRG
PSEBN03357				TIGGLGPPLTHEALAG
BORFD03982 PSEU503413				LTGGLGPPLTREALAG XTGGLGPPLTRAALAD
PSEUT03467				ATGGLGPPLTRAALAD
CUPNJ04970	TSAAQ	GPARGAELARI L	RCDCGACHGMT	LCGGLG3PLTAAALTG
CUPTR05130	RAADA	APARÇAQLAHNL	REDCGACHGMT	LRGGLGPPLT PAGLAG
Cupnh05899 Cupnn05794				LGGGLGPPLTREALSG LRGGLGPPLTREALSG
THIDA00076				LRGGLGPALTSETLKD
ALIDK02236	TLAAAF	AP SRAAQLVRKV	RODCGOCHGMR	LTGGLGPALTRQALAA
ACIETO1671				LTGGLGPALT RALAG
ACISJ01764 ACCPU02313				LTGGLGPALTREALAG LCGGLGPSLKPDALRD
LEPCP01428				LTGGLGPALTNEALAE
RUBGI00659				LGGGLGPALT?QALAD
HAHCH04183				IKGGLGP SIR PERNOA
POLGS00564 PARDP02406				RXGGLGSPLTAEALAH
ROSIO01453				NTGGLGRPITAAALAG NTGGLGSPITJEALEG
ROSLO03035				TGGLGSAITFCALLG
DINSH03144				RKGGLGPCLRARALAH
RUEPO04047	TATA.A	ETLDPDALR <mark>RIV</mark>	QODCGSCHGLT:	LXGGLGPCIRPEALGH
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Figure S3 Sequence alignment of NirC orthologs from different denitrifying species (for identification refer to the OMA server (Altenhoff *et al.*, 2018)). The sequence at the top is from *Pseudomonas aeruginosa* PA01, which was used in this study. Green boxes mark heme-interacting residues. A cyan box indicates E71 mutated to alanine to enable crystal formation and the black box highlights residues belonging to the hinge loop, which adopts different conformation in the 3D domain swapped dimers. Bold blue letters indicate identity in all residues and red letters are equivalent to 70%. Sequence numbering is based on whole protein including the signal peptide. The figure was prepared with ESPrit3.0 (Robert & Gouet, 2014).

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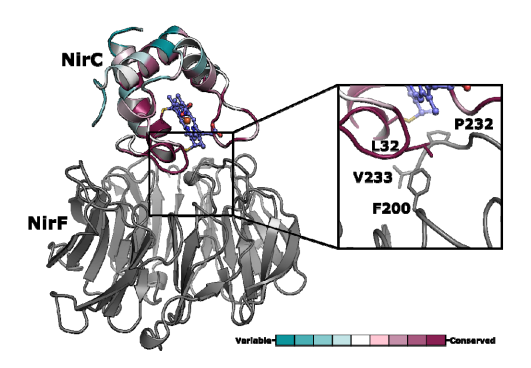


Figure S4 Depiction of NirC (coloured according to sequence conservation) and NirF (grey) (PDB: ####) after superposition onto the cytochrome *c* and *d*₁-domains of nitrite reductase NirS (PDB: 1nir), respectively.