The genome of *Nitrospina gracilis* illuminates the metabolism and evolution of the major marine nitrite oxidizer

Sebastian Lücker1*, Boris Nowka2, Thomas Rattei2, Eva Spieck2 and Holger Daims1*

1 Department of Microbial Ecology, Ecology Centre, University of Vienna, Vienna, Austria  
2 Department of Microbiology and Biotechnology, BioCenter Klein Flottbek, University of Hamburg, Hamburg, Germany  
3 Department of Computational Systems Biology, Ecology Centre, University of Vienna, Vienna, Austria

**INTRODUCTION**

Nitrite-oxidizing bacteria (NOB) are chemolithoautotrophic organisms catalyzing the oxidation of nitrite to nitrate, which is the second step of nitrification and a major biogeochemical process. Nitrite is substrate not only for nitrite oxidation but also for denitrification and anaerobic ammonium oxidation (anammox), both leading to nitrogen losses from ecosystems to the atmosphere. Therefore, the activity of NOB is a key factor controlling the retention of fixed nitrogen in the form of nitrate in the biosphere.

NOB are a phylogenetically diverse functional group. Known representatives belong to the genera *Nitrobacter*, *Nitrotaga*, and *Nitroccocus* within the Alpha-, Beta-, and Gammaproteobacteria, respectively (Winogradsky, 1890; Watson and Waterbury, 1971; Alawi et al., 2007), to the genus *Nitrospina* within the *Nitrospiraceae* (Daims et al., 2001), and to the genus *Nitrolancinetus* within the *Chloroflexi* (Sorokin et al., 2012). The genus *Nitrospina* was provisionally assigned to the *Deltaproteobacteria* (Teske et al., 1994), but this affiliation has been questioned due to difficulties to classify this genus solely by 16S rRNA-based phylogenetic inference (Schloss and Handelsman, 2004). Most NOB lineages are highly versatile and can be found in terrestrial, limnic, and marine ecosystems as well as in man-made habitats such as wastewater treatment facilities. However, the dominant nitrite oxidizers in the oceans appear to be *Nitrosopina*. The type species of this genus, *Nitrospina gracilis*, was isolated from ocean surface waters (Watson and Waterbury, 1971). It forms long, slender, Gram-negative rods, grows purely chemolithoautotrophically with nitrite and CO₂ as the sole source of energy and carbon, respectively, and was described to be strictly aerobic using oxygen as terminal electron acceptor. No heterotrophic potential was observed and many organic substrates even inhibited the growth of this organism (Watson and Waterbury, 1971). *Nitrospina*-like bacteria were frequently detected by molecular methods in open ocean water (Fuchs et al., 2005; DeLong et al., 2006) as well as in marine sediments (Hunter et al., 2006; Jorgensen et al., 2012), but surprisingly also in marine oxygen minimum zones (OMZs; Labrenz et al., 2007; Fuchsen et al., 2011) and anoxic marine sediments (Davis et al., 2009;
Jorgensen et al., 2012). Moreover, nitrite oxidation has been measured in OMZs (Füssel et al., 2012). Interestingly, the distribution profiles of Nitrospina and ammonia-oxidizing archaea (AOA) were found to correlate in some coastal and open ocean habitats (Mincer et al., 2007; Santoro et al., 2010). Prolonged coexistence of Nitrospina and AOA closely related to "Candidatus Nitrosopumilus maritimus" was even observed in an enrichment culture (Park et al., 2010), further strengthening the functional link between these nitriﬁers. The only non-marine habitats Nitrospina-like 16S rRNA sequences were derived from are uranium mill tailings (Kadeva and Selenska-Pobell, 2005) and an alpine subsurface radioactive thermal spring (Weidler et al., 2007). Thus, Nitrospina apparently are conﬁned mainly to marine environments.

In marine ecosystems, nitrate accounts for as much as 88% of the ﬁxed nitrogen, corresponding to 5.8 × 10^9 Tg (Gruber, 2008). While rapidly assimilated by phytoplankton in ocean surface waters, nitrate accumulates in the deep sea where it constitutes the largest pool of ﬁxed inorganic nitrogen in the biosphere. The only biochemical reaction known to form nitrate is bacterial nitrite oxidation, which takes place during nitrification and, to a lesser extent, during anammox (Strous et al., 2006). Contrasting the obviously immense importance of Nitrospina as a major source of nitrate in marine environments, very little biochemical and genomic data is available for these enigmatic organisms, and even the phylogenetic afﬁliation of the genus is still unclear. This embarrassing lack of knowledge is mainly caused by the recallintrace of Nitrospina to cultivation in the laboratory. Their slow growth rates with a minimum generation time of 24 h (Watson and Waterbury, 1971) further impede biochemical characterization. In this study, we employed a genomic approach to gain deeper insights into the metabolic potential of Nitrospina gracilis strain 3/211. The sequenced genome of this organism shed light on the core metabolism enabling Nitrospina of chemolithoautotrophic growth with nitrite. Comparative genomic analyses illuminated the evolutionary descent of Nitrospina and its nitrite-oxidizing capacity. By utilizing concatenated datasets of established marker proteins, the phylogenetic afﬁliation of Nitrospina could ﬁnally be assessed.

MATERIALS AND METHODS

CULTIVATION

Cells of N. gracilis were grown in 51 batch cultures in mineral salts medium prepared with 70% seawater containing 0.3 mM nitrite as the sole energy source. The marine medium (Watson and Waterbury, 1971) was composed of 700 ml ﬁltered seawater and 300 ml distilled water, supplemented with 0.005 g l−1 CaCl2 × 2 H2O, 0.1 g l−1 MgSO4 × 7 H2O, 0.001 g l−1 FeSO4 × 7 H2O, 0.0017 g l−1 KH2PO4, 6 µg l−1 CuSO4 × 5 H2O, 25 µg l−1 Na2MoO4 × 2 H2O, 50 µg l−1 MnCl2 × 4 H2O, 0.5 µg l−1 CoCl2 × 6 H2O, and 25 µg l−1 ZnSO4 × 7 H2O. The pH was adjusted to 6.5–7.0 and changed to 7.4–7.6 within 2 days after autoclaving. Cultures were started with 1% inoculum and incubated at 28°C in the dark. When nitrite was consumed for the first time, the culture bottles were started to be stirred moderately. Nitrite consumption was measured and substrate was replenished regularly. To check for purity of the culture, aliquots were incubated in heterotrophic marine medium (0.15 g l−1 bactopeptone, 0.15 g l−1 yeast extract, 0.055 g l−1 sodium pyruvate in marine mineral salts medium) without nitrite.

DNA EXTRACTION, GENOME SEQUENCING, AND ANNOTATION

High-molecular weight genomic DNA was isolated following the hexadecltrimethylammonium bromide (CTAB) protocol as provided by the DOE Joint Genome Institute (JGI, http://my.jgi.doe.gov/general/protocols.html) with minor modiﬁcations. Shortly, cells were resuspended in TE buffer and digested by the sequential addition of lysozyme and 10% (w/v) sodium dodecylsulfate (SDS) and Proteinase K. Following the subsequent incubation in CTAB/NaCl solution, the DNA-containing aqueous phase was extracted in Roti-Phenol/Chloroform/Isomylalcohol (Carl Roth, Karlsruhe, Germany) and washed in Roti-C/I once to remove residual phenol. After DNA precipitation with isopropanol and washing with 70% ethanol, the DNA was suspended in TE buffer and stored at −20°C. Genome sequencing was performed at LGC Genomics (Berlin, Germany) using GS FLX Titanium sequencing technology. One 1/4 picotiter plate was run with an 8 kb paired-end large-insert library. Sequence read ﬁltering and correction, contig assembly, and scaffold formation were done by LGC Genomics.

The draft genome of N. gracilis was integrated into the MicroScope annotation platform (Vallenet et al., 2009). After automated prediction and annotation of coding sequences (CDS), the annotation of all CDS in key pathways, including those for nitrite oxidation, respiration, and carbon ﬁxation, was manually reﬁned by using the respective tools of MaGe (Vallenet et al., 2006) as described in detail elsewhere (Lücker et al., 2010).

The draft genome sequence of Nitrospina gracilis strain 3/211 has been deposited at EMBL-EBI under project number PRJEB1269.

AMPLIFICATION, CLONING, AND SEQUENCING OF nxrB GENES

As both copies of the operon containing the genes of nitrite oxidoreductase (NXR) were incomplete and lacked the nxrB gene coding for the beta subunit of NXR, these regions were speciﬁcally PCR-ampliﬁed from genomic DNA, cloned in E. coli by TOPO-TA Cloning (Life Technologies, Paisley, UK), and Sanger-sequenced (3130xl Genetic Analyzer, Applied Biosystems, Vienna, Austria). Operon-speciﬁc PCR was carried out using the newly designed primers Nspn_nxrA1_3064f (CATTCTTGCT-GAGGTCA; forward) in combination with Nspn_nxrB1_111r (CATATCCAACACCACTGCTG; reverse) and Nspn_nxrA2_3064f (CATTCCAGATCCCAGAGG; forward) with Nspn_nxrB2_111r (CAAATGCTGACACACTCC; reverse). These primer pairs target the C-terminal ends of the alpha subunit genes nxrA and the N-terminal parts of the gamma subunits nxrC. Cycling conditions were as follows: an initial denaturation step (94°C, 5 min) was followed by 35 cycles of template denaturation (94°C, 30 s), primer annealing (52°C, 30 s), and product elongation (72°C, 1.5 min). Cycling was completed by a ﬁnal elongation step (72°C, 20 min). The cloned amplicons were sequenced using M13 primers.

nxrB sequences obtained in this study have been deposited at GenBank under accession numbers KC262217 and KC262218.
**PHYLOGENETIC ANALYSES**

An amino acid sequence database of heme-copper oxidase-family enzymes was established using the software ARB (Ludwig, 2004). Multiple sequence alignments were generated automatically using ClustalW2 (Larkin et al., 2007) and manually refined using the sequence editor included in ARB. Protein sequences of the NxrA and NxrB subunits of NXR were imported into existing datasets of type II DMSO reductase-family molybdopterin cofactor-binding proteins (Lücker et al., 2010). The sequences were aligned using the automatic aligner integrated in ARB, followed by manual refinement.

Amino acid sequences of 49 established phylogenetic marker proteins (Table A1 in Appendix) were downloaded from the Microbial Genome Database for Comparative Analysis (Uchiyama et al., 2010). Ortholog clusters were manually filtered to include bacterial sequences only. To avoid erroneous phylogenetic signals caused by horizontal gene transfer (HGT) or incorrect annotations, organisms with more than one entry for a query protein were excluded. This resulted in datasets with 443–502 entries for each protein. The single-protein datasets were automatically aligned with ClustalO (Siers et al., 2011), running two iterations, followed by filtering for badly aligned regions or sites of rare insertions by Gblocks (Talavera and Castresana, 2007) with settings “Minimum Length of an Initial Block (b0) = 2,” “Maximum Number of Contiguous Non-conserved Positions (b3) = 50,” “Minimum Length of A Block (b4) = 3,” and “Allowed Gap Positions (b5) = all” to obtain less stringent filtering. The filtered alignment files were subsequently concatenated using an in-house generated C++ program and imported into ARB.

Phylogenetic analyses were performed by applying maximum-likelihood, maximum-parsimony, and Bayesian interference methods: RAxML [version 7.0.3 (Stamatakis et al., 2005); 100 rapid or 100 advanced bootstrap iterations], PhyML (Felsenstein, 2005), ProML and ProtPars (PHYLIP package, version 3.6 (Guindon and Gascuel, 2003); 100 bootstrap iterations), TreePuzzle [version 5.0 (Schmidt et al., 2002)], and MrBayes [version 3.1 (Ronquist and Huelsenbeck, 2003); 2–5 M generations] using the JTT, WAG, or Blossum62 substitution models. Where applicable, N-terminal signal peptide sequences were excluded from the analyses and manually created indel filters were used.

**PHYLOME CONSTRUCTION**

For the calculation of phylogenetic trees for each protein of the in silico deduced proteome of *N. gracilis*, the software PhyloGenie (Frickey and Lupas, 2004) was used as previously described (Lücker et al., 2010). The reference database for PhyloGenie was generated from the National Center for Biotechnology Information non-redundant protein database NCBI nr (Sayers et al., 2011), in which taxon names were edited to remove characters that would interfere with the export of phylogenetic trees in the Newick file format. The NCBI taxonomy database name file was adapted in a similar manner. PhyloGenie analysis was performed for each query protein using the default parameters with the following modification: −blammerparams = −taxid f. For the BLAST (Altschul et al., 1990) searches in PhyloGenie, NCBI BLAST (version 2.2.19) was used. Protein phylogenies were calculated on the basis of full or partial automatic alignments produced by the BLAMMER program included in PhyloGenie. Trees were calculated using RAxML (Stamatakis et al., 2005) with the JTT substitution model and 100 bootstrap iterations. All trees were subject of specific taxonomic queries using Phat (Frickey and Lupas, 2004). Furthermore the taxonomy of closest homologs was determined from these trees since this strategy is less prone to alignment artifacts than best hits from automatic multiple sequence alignments. An in-house Perl script determined the tree node closest to the respective query proteins from *N. gracilis* and consecutively sorted all operational taxonomic units behind this node by their distances to the query. The organism with the smallest distance was considered to contain the closest homologous protein.

**RESULTS**

**GENOME RECONSTRUCTION**

The obtained sequence reads were assembled into 109 contigs, which were arranged in four scaffolds. The largest scaffold encompassed 106 contigs, whereas the remaining three scaffolds consisted of single contigs whose location relative to the other contigs could not be determined unambiguously. The presence of frameshift mutations in some essential and highly conserved genes indicates that some pyrosequencing-associated errors could not be detected and corrected during read filtering and assembly. The obtained genome sequence of *N. gracilis* has a size of 3,067,213 bp and an average G + C content of 56.2%. It contains 3,147 predicted coding sequences (CDS; Table 1), but the large number of gene fragments (due to sequence gaps within genes and frameshifting) likely causes an overestimation of the CDS count. The genome sequence contains 1 complete rrn operon and 45 tRNA genes (1–5 for each of the 20 amino acids). The sequenced genome is near-complete based on the full set of tRNAs, the presence of all 127 clusters of orthologous groups (COGs) of proteins conserved in all 50 bacterial genomes currently in the COG database (Tatusov et al., 2003), and the estimated small gap sizes within the large scaffold.

Notably, the smallest of the four scaffolds contained only an *nxrB* gene coding for the beta subunit of nitrite oxidoreductase (NXR). The NXR operon, which contains the genes of the alpha (*nxrA*), beta (*nxrB*), and gamma (*nxrC*) subunits, is duplicated in the *N. gracilis* genome. Since the sequences of the two *nxrB* genes are highly similar, the complete NXR operons could not be unambiguously assembled from the sequence reads. Therefore, operon-specific primers where designed and used to amplify the genomic regions between *nxrA1* and *nxrC1*, and *nxrA2* and *nxrC2*, respectively, in order to obtain by cloning and Sanger-sequencing the complete operon sequences including *nxrB*. When translated into their protein sequence, the two NxrB copies had a length of 425 amino acids and an identity of 100% (with five synonymous single-base substitutions on the nucleotide level).

**PHYLGENY**

A set of 49 established phylogenetic marker proteins (Table A1 in Appendix) was extracted from the genome of *N. gracilis* and a range of other complete bacterial genomes covering all bacterial phyla for which genome sequences are available. The concatenated dataset of these proteins enabled a detailed phylogenetic analysis of the affiliation of *Nitrospina* within the *Bacteria*. Preliminary
trees containing sequence information from 502 bacteria indicated deep phylogenetic branching of *N. gracilis* between the Proteobacteria and a group consisting of the Acidobacteria and Nitrospirae phyla (data not shown). Consecutively, a reduced dataset containing representatives of these and few other closely affiliated phyla was generated and used for more detailed analyses employing different tree calculation methods. These analyses clearly indicated a position of *N. gracilis* close to, but outside the Proteobacteria (Figure 1) and not within any other bacterial phylum either. Notably, the Epsilonproteobacteria did not cluster with the other Proteobacteria, a branching pattern that has been observed previously (Ludwig, 2010), and formed a separate branch together with the Aquificae. This grouping of Aquificae and Epsilonproteobacteria is consistent with other marker protein-based analyses that also found the Aquificae not to group as one of the most basal bacterial phyla (Korbel et al., 2002; Cavalier-Smith, 2006). The Acidobacteria and Nitrospirae formed a superphylum in the Bayesian interference analysis that grouped as the next most closely related phyla to Nitrospina (Figure 1A), followed by the aforementioned Aquificae/Epsilonproteobacteria group and the Deferribacteres. Maximum-likelihood analysis grouped Nitrospina as sister phylum of the Acidobacteria, with the Nitrospirae forming a separate branch (Figure 1B), and the maximum-parsimony approach indicated a superphylum of all three groups (Figure 1C). The separation of Nitrospina from the Proteobacteria thus is supported by all analyses, and the grouping with the Acidobacteria and Nitrospirae in some analyses strongly contradicts the possibility that Nitrospina forms a basal new class within the Proteobacteria.

To further assess the separate phylogenetic position of *N. gracilis* as member of a novel phylum, two additional methods were employed: comparison of shared conserved insertion and deletion signature sequences (indel analysis; Gupta and Griffiths, 2002) and gene transposition analysis (Kunisawa, 2006). Both analyses were consistent with the calculated phylogenetic trees by suggesting that *N. gracilis* belongs to a phylum related to, but separate from the Proteobacteria. *N. gracilis* carries the characteristic insertions in

---

**Table 1 | Overview of key features of the *Nitrospina gracilis* strain 3/211 genome.**

| Feature | Value |
|---------|-------|
| Draft genome size | 3,067,213 bp |
| Estimated genome size | 3,101,305 bp |
| Estimated genome completeness | 98.90% |
| Number of scaffolds | 4 |
| Number of contigs | 109 |
| Average G+C content | 56.21% |
| Number of genomic objects [CDS, fragment CDS (r,t)RNA] | 3202 |
| Number of coding sequences (CDS) | 3147 |
| rRNA genes | 3 |
| tRNA genes | 45 |

**CLUSTERS OF ORTHOLOGOUS GROUPS (COG) AUTOMATED CLASSIFICATION**

| COG | Function | CDS | CDS (%) |
|-----|----------|-----|---------|
| D   | Cell cycle control, cell division, chromosome partitioning | 43  | 1.37 |
| M   | Cell wall/membrane/envelope biogenesis | 222 | 7.05 |
| N   | Cell motility | 97  | 3.08 |
| O   | Posttranslational modification, protein turnover, chaperones | 176 | 5.59 |
| T   | Signal transduction mechanisms | 166 | 5.27 |
| U   | Intracellular trafficking, secretion, and vesicular transport | 91  | 2.89 |
| V   | Defense mechanisms | 59  | 1.87 |
| A   | RNA processing and modification | 1   | 0.03 |
| B   | Chromatin structure and dynamics | 1   | 0.03 |
| J   | Translation, ribosomal structure, and biogenesis | 181 | 5.75 |
| K   | Transcription | 114 | 3.62 |
| L   | Replication, recombination and repair | 140 | 4.45 |
| C   | Energy production and conversion | 219 | 6.96 |
| E   | Amino acid transport and metabolism | 295 | 9.37 |
| F   | Nucleotide transport and metabolism | 79  | 2.51 |
| G   | Carbohydrate transport and metabolism | 138 | 4.39 |
| H   | Coenzyme transport and metabolism | 114 | 3.62 |
| I   | Lipid transport and metabolism | 78  | 2.48 |
| P   | Inorganic ion transport and metabolism | 196 | 6.23 |
| Q   | Secondary metabolites biosynthesis, transport, and catabolism | 94  | 2.99 |
| R   | General function prediction only | 456 | 14.49 |
| S   | Function unknown | 238 | 7.56 |
the transcription termination factor Rho and the alanine-tRNA ligase AtaS, which are common to all Proteobacteria and a range of phyla including the Chlamydiae and Aquificae (Gupta, 2000; Gupta and Griffiths, 2002), but lacks signature sequences in the inorganic phosphatase Ppi, molecular chaperone DnaK, and CTP synthetase PyrG that distinguish the Proteobacteria from the other phyla (Gupta, 2000, 2001). Analyses of gene transposition events (Kunisawa, 2010, 2011) also indicated a close relationship to the Proteobacteria, Nitrospirae, and to a lesser extent the Deferribacteres and Aquificae, but clearly distinguished N. gracilis from these established phyla based on unique differences in bordering genes.

Another way to look at ancestry and identify HGT events at the same time is by phylogeic reconstruction, which is the determination of the relationship of every protein encoded in a genome by phylogenetic inference (Frickey and Lupas, 2004). If the exact phylogenetic position of an organism is known this is an elegant method to determine the fraction of horizontally transferred genes and to identify potential donor organisms. For organisms of uncertain affiliation, the phylomic information may reveal ancestral relationships among homologs in Nitrospirae. However, in most cases the phylogeny of N. gracilis was reconstructed and the closest relative (based on the distance to the respective homolog in N. gracilis) was assigned according to the distance to the closest relative based on the distance to the respective homolog in N. gracilis) was extracted from each single-protein tree. Of the 2,354 proteins of N. gracilis that have homologs in public databases, a large part clustered on the class level with Deltaproteobacteria (n = 514), followed by Gammaproteobacteria (n = 225) and Nitrospirae (n = 211). On the genus level Nitrospira had most hits (n = 161), followed by the deltaproteobacterial Geobacter (n = 81), the recently described “Candidatus Methylomirabilis” (Ettwig et al., 2016; n = 68), and “Candidatus Kuenenia” (Strous et al., 2006; n = 51). The numerous closest homologs in Proteobacteria and Nitrospirae are consistent with the phylogenetic trees as well as the indel and gene transposition analyses. The large number of proteins clustering with Nitrospira may also reflect a high degree of functional similarity in Nitrospira and Nitrospira. In contrast, the large number of proteins affiliated to homologs in the Methylomirabilis and Kuenenia was unexpected and might indicate a large extent of HGT between these organisms and Nitrospira. One possible caveat of the applied tree query approach is that the next neighbors in a tree may not always have the shortest distance to each other. However, manual reassessment by using Phat within the PhyloGenie program (Frickey and Lupas, 2004) revealed the same trends for the relationship of Nitrospira proteins to the aforementioned groups (Figure 2). These results strongly corroborate the phylogenetic placement of Nitrospira in a distinct phylum with the Proteobacteria and Nitrospirae as most closely related known phyla.

NITRITE OXIDATION AND NITROGEN METABOLISM
N. gracilis is a chemolithoautotrophic nitrite-oxidizing bacterium. It gains all energy required for growth from the oxidation of nitrite and fixes CO₂ as sole carbon source, using electrons derived from nitrite oxidation. The key enzyme for nitrite oxidation is nitrite oxidoreductase, which shuttles two electrons per oxidized nitrite and fixes CO₂. The key enzyme for nitrite oxidation is nitrite oxidoreductase, which shuttles two electrons per oxidized nitrite and fixes CO₂. The key enzyme for nitrite oxidation is nitrite oxidoreductase, which shuttles two electrons per oxidized nitrite and fixes CO₂.
NO$_3^-$ into the electron transport chain. The NXR belongs to the type II group of molybdopterin-binding enzymes within the DMSO reductase enzyme family (Jormakka et al., 2004). These enzymes are membrane-associated complexes of three subunits. The catalytic alpha subunit contains the molybdenum bis molybdopterin guanine dinucleotide (Mo-bis-MGD) cofactor and one [4Fe–4S] iron-sulfur (Fe-S) cluster. The beta subunit contains one [3Fe–4S] and three [4Fe–4S] cluster and transfers the electrons to and from the membrane-integral respiratory chain (Lowe et al., 2010). In Nitrospina, however, the NXR complex is membrane-associated (Bartosch et al., 1999; Spieck and Bock, 2005b). Thus, an additional membrane anchor likely is required for the formation of active NXR in this organism. In addition to the two linked copies of NxrC, N. gracilis has genes of two other putative $\gamma$-subunits (Table S1 in Supplementary Material). NxrC3 is located in the direct vicinity of a TorD-like chaperone that most likely functions in Mo cofactor insertion into NxrA prior to Tat export, as has been proposed for other Mo-bis-MGD-containing enzymes such as selenate reductase (Guymer et al., 2009). This NxrC has one predicted membrane helix as well as the DMSO reductase-like heme $b$-binding domain. NxrC4 is encoded in a genetic context unrelated to nitrite oxidation and shows only low similarities to the other NxrC copies, but also contains one predicted transmembrane helix and the heme $b$-binding domain.

Aside from these candidate NXR $\gamma$-subunits, the genome of N. gracilis codes for at least one putative alternative NxrC (alt_NxrC1, Table S1 in Supplementary Material). This protein is remarkable since it contains not only two transmembrane helices, but also two cytochrome $c$-type $c$-binding sites and an iron-sulfur (Fe-S) cluster. However, the NXR complex contains three additional copies of this putative alternative $\gamma$ subunit (Table S1 in Supplementary Material), which all contain 1–3 cytochrome $c$ binding sites and an iron-sulfur (Fe-S) cluster. How-
be a functional nitrite reductase, while NirK2 might have reduced or altered activity (Wherland et al., 2005). Interestingly, all analyzed genomes of NOB except *Nitrolobactes hollandiacus* (Sorokin et al., 2012) contain *nirK* genes, but the function of nitrite reduction in NOB has not clearly been determined yet. No known nitric oxide (NO) reductase is encoded in the *N. gracilis* genome and a role of NirK in denitrification therefore seems unlikely. NO-stimulated NADH formation in *Nitrobacter* and was speculated to be an intermediate of nitrite oxidation in this organism (Freitag and Bock, 1996; Poughon et al., 2001). A later study proposed an alternative role of NO in the regulation of the cellular redox state in *Nitrobacter* (Starkenburg et al., 2008). In this model the terminal cytochrome *c* oxidase is reversibly inhibited by NO under low-oxygen conditions, leading to an increased flow of electrons through the reverse electron transport chain. This causes higher CO₂ fixation rates and may increase the biosynthesis of the storage compound poly-β-hydroxybutyrate (Starkenburg et al., 2008). For NOB other than *Nitrobacter* no function of NirK has been proposed, and denitrification of nitrite to NO under anoxic conditions has not been observed yet. *N. gracilis* grows on nitrate as sole source of both energy and nitrogen. Consistently, the genome contains the *nirK* gene for assimilatory ferredoxin-nitrite reductase, which is located directly next to *nirC* encoding a nitrite transporter of the formate/nitrite transporter family (Figure 3; Table S1 in Supplementary Material). A second nitrite transporter of the NarK nitrate/nitrite antiporter type, as well as two Amt-type ammonium transporters, likely is also involved in nitrogen uptake for assimilation (Table S1 in Supplementary Material).

### ENERGY METABOLISM AND REVERSE ELECTRON TRANSPORT

*N. gracilis* encodes a terminal oxidase of the cytochrome *cbb₃*-type (Figure 3; Table S1 in Supplementary Material). As the *cbb₃*-type terminal oxidases have high affinities for O₂ (Cosseau and Batut, 2004), its presence will allow *Nitrospina* to sustain respiration in low-oxygen environments. The electrons derived from nitrite oxidation most likely are shuttled via cytochrome *c* to this terminal oxidase for reduction of O₂ as terminal electron acceptor, yielding the energy for the formation of a proton gradient across the cytoplasmic membrane. The proton motive force drives ATP formation by a canonical F₁F₀ ATPase (Figure 3; Table S1 in Supplementary Material). Interestingly, the three subunits of the cytochrome *cbb₃* oxidase of *Nitrospina* are fused into one gene encoding a protein of 954 aa. The N-terminal part of this protein corresponds to subunit I, has 13 predicted transmembrane helices, and contains most of the characteristic binding sites for two heme *b* and the low-potential copper (C₄₉) center. The C-terminal part corresponds to subunits II and III with one and two heme *c* binding sites, respectively. Interestingly, one amino acid residue in the C₄₉ center is unprecedented in heme-copper oxidases (Figure 4). In all members of this family the copper is complexed by three histidine residues, which are numbered His<sup>207</sup>, His<sup>257</sup>, and His<sup>258</sup> in the cytochrome *cbb₃* oxidase of *Pseudomonas stutzeri* whose structure has been determined (Buschmann et al., 2010). At position 203 (*P. stutzeri* numbering) most of these enzymes contain a highly conserved tryptophan that is involved in passing electrons from the high-sphin heme *b₃* to the C₄₉ center where oxygen is reduced (Zaslavsky and Gennis, 2000; Pereira et al., 2001). This tryptophan is replaced by glycine in the *N. gracilis* enzyme, thus clearly impeding any charge transfer. Directly upstream, however, is an additional histidine residue found only in *N. gracilis* (His<sup>202</sup>, *P. stutzeri* numbering) whose positively charged side chain may adopt the role of the hydrophobic Trp<sup>203</sup> in electron transfer. In addition, two uncommon tyrosine residues flanking the His<sup>207</sup> at positions 205 and 208 (*P. stutzeri* numbering) might also function as electron carriers. Besides *Nitrospina*, the His<sup>203</sup>, Gly<sup>206</sup>, and Tyr<sup>208</sup> residues are found only in subunit I of putative heme-copper oxidases from various *Sinorhizobium* genomes (e.g., *Sinorhizobium meliloti* SM11, Figure 4), where they occur in one of several homologous cytochrome *cbb₃* oxidases. Their presence in other organisms strongly suggests that these unusual amino acid substitutions are not sequencing errors in the *N. gracilis* genome.

*Nitrospina* depends on the electrons derived from nitrite oxidation for the regeneration of reducing equivalents and for autotrophic CO₂ fixation by the reductive tricarboxylic acid (rTCA) cycle. These electrons enter the respiratory chain at the level of cytochrome *c* due to the high redox potential of the NO₂⁻/NO₃⁻ couple. Therefore, reverse electron transport to NAD<sup>+</sup> (E<sub>red</sub> = −320 mV) and to ferredoxin (E<sub>red</sub> ≤ −500 mV) is required and most likely powered by the proton motive force across the cytoplasmic membrane. Mechanistically, the cytochrome *bc* complex and NADH-quinone oxidoreductase (complexes III and I of the respiratory chain, respectively; Elbehti et al., 2000) are involved in electron transport from cytochrome *c* to NAD<sup>+</sup>, whereas a novel type of complex I may catalyze the reduction of ferredoxin. In addition, electrons can be shuttled from the quinone pool into the rTCA cycle by the succinate dehydrogenase/fumarate reductase (complex II; Lemos et al., 2002). Moreover, the use of organic storage compounds such as glycogen as energy source requires a respiratory chain with the fission of the cytochrome *bc* complex rather than to the *b/b₃* second copy of the cytochrome *bc* (Table S1 in Supplementary Material). Notably, *Nitrospina* carries elsewhere in the genome a second copy of the cytochrome *c* subunit that is not split, but also appears to be of the *b/b₃*-type (Table S1 in Supplementary Material). The only putative cytochrome *c* subunit encoded next to the cytochrome *bc*-like operon is an atypical octaheme cytochrome *c*. Interestingly, the exactly same operon structure including a similar
Nitrospina gracilis 3/211

| organism          | Gene Product                        | Location       |
|-------------------|-------------------------------------|----------------|
| Sinorhizobium meliloti SM11 (AEH81668) | cbb, a, b, c, d, e, f, g, h, i, j, k, l, m, n, o, p, q, r, s, t, u, v, w, x, y, z | nuoA-M          |
| Rhizobium sp. TGN441 (X66992)           | cbb, a, b, c, d, e, f, g, h, i, j, k, l, m, n, o, p, q, r, s, t, u, v, w, x, y, z | nuoA-M          |
| Pseudomonas aeruginosa PA01 (AAG04943)  | cbb, a, b, c, d, e, f, g, h, i, j, k, l, m, n, o, p, q, r, s, t, u, v, w, x, y, z | nuoA-M          |
| Pseudomonas aeruginosa PA01 (AAG04943)  | cbb, a, b, c, d, e, f, g, h, i, j, k, l, m, n, o, p, q, r, s, t, u, v, w, x, y, z | nuoA-M          |
| Nitrococcus mobilis NB-231 (2P_01127177) | a, a, b, c, d, e, f, g, h, i, j, k, l, m, n, o, p, q, r, s, t, u, v, w, x, y, z | sulfoXY         |
| Sulfolobus acidocaldarius (P39481)      | SoxM, SoxF, SoxC, SoxB, SoxA | sulfoXY         |
| Aeropyrum pernix K1 (BAA80623)          | ba, c, d, e, f, g, h, i, j, k, l, m, n, o, p, q, r, s, t, u, v, w, x, y, z | sulfoXY         |
| Geobacillus stearothermophilus (BAA35056) | bo, c, d, e, f, g, h, i, j, k, l, m, n, o, p, q, r, s, t, u, v, w, x, y, z | sulfoXY         |
| Sulfolobus acidocaldarius (CA44510)     | SoxB, SoxF, SoxC, SoxB, SoxA | sulfoXY         |
| Geo bacter metallireducens GS-15 (YP_386423) | qNOR | sulfoXY         |
| Haloracula marismortui (YP_138606)      | qNOR | sulfoXY         |
| Nitrosomonas europaea (NP_842019)      | eNOR | sulfoXY         |
| Pseudomonas stutzeri (YP_004715865)    | eNOR | sulfoXY         |

FIGURE 4 | Partial multiple sequence alignment of selected heme-copper oxidase-family members. Relationship of the enzymes is indicated by a cladogram. The oxidase type is indicated for each enzyme. Amino acid residues involved in the formation of the copper (Cu-binding site) are highlighted in color and by a symbol indicating function: dark blue, histidine residues involved in Cu₃-binding (c); purple, residues involved in electron transport from the high-spin heme to Cu₃(C); red, uncommon amino acids; orange, crosslinking tyrosine residue stabilizing one of the Cu₃-binding histidines (h), green and turquoise, alternative conserved residues in this position.

octaheme protein is conserved in the genome of the anammox organism “Candidatus Kuenenia stuttgartiensis” (de Almeida et al., 2011) and in “Candidatus Methylocamibalis oxyfera,” a noteworthy bacterium capable of nitrite-driven anaerobic methane oxidation (Ettwig et al., 2010).

The genome of N. gracilis contains the genes for all 14 subunits of NADH:quinone oxidoreductase (Nuo; complex I) needed for the forward transport of electrons from NADH to quinone (Figure 3; Table S1 in Supplementary Material). All Fe-S clusters required for electron channeling between NADH and quinone are preserved, as are the NADH and flavin mononucleotide (FMN)-binding patterns within subunit NuoE. The mnoA-M genes are clustered in one operon on the lagging strand, whereas mnoN is located on the opposite strand five genes downstream of mnoA. Besides the canonical NADH:quinone oxidoreductase (NUO-1), a second copy is present, but the genes coding for this NUO-2 are distributed across the genome (Table S1 in Supplementary Material). While mnoB2 and mnoF2 are not organized in an operon structure, the other genes of NUO-2 form two clusters consisting of mnoACHJIKLMN and mnoEGD, respectively. Intriguingly, this complex shows substantial differences to NUO-1 in the binding patterns for cofactors in the catalytic subunits. NADH in canonical...
NUOs (including NUO-1 of N. gracilis) binds to NuoF, where two electrons are transferred onto the primary electron acceptor FMN that serves as a two-to-one-electron converter (Brandt, 2006). Consecutively, FMN feeds single electrons into a chain of seven conserved Fe-S clusters distributed throughout subunits NuoE, G, I, and B in the hydrophilic arm of the complex, at the end of which quinone reduction occurs (Sazanov and Hinchliffe, 2006). The N. gracilis NuoF2 lacks the FMN binding site, but contains one [2Fe-2S] and two [4Fe-4S] ferredoxin-like clusters in addition to the Fe-S cluster N3 and the NADH binding domain. Similarly, NuoG2 has the characteristic cysteine binding patterns for Fe-S clusters N1b, N4, N5, and N7, along with one additional [4Fe-4S] cluster. Without the FMN cofactor a function as NUO is unlikely, but a possible alternative role for NUO-2 might be in the reverse electron transport from quinol to low-potential ferredoxin. Two enzymes of the rTCA cycle (see below) belong to the 2-oxoacid:ferredoxin oxidoreductase family and require reduced low-potential ferredoxin as electron donor for the reductive carboxylation of their substrates. Common mechanisms to reduce low-potential ferredoxin with electrons from NADH (Rnf complex; Biegel and Müller, 2010) or H2 (Ech complex; Welte et al., 2010) employ membrane-integral complexes that use the proton motive force for reverse electron transport to ferredoxin (E°’ < −500 to −420 mV). These complexes evolved from a NUO-like ancestor (Buckel and Thauer, 2013). Nitrospina lacks these known pathways, but the NUO-2 complex is an interesting candidate to perform ferredoxin reduction with quinol as electron donor. Due to the lack of the FMN binding site, the NuoF2 subunit cannot interact with a two-electron carrier such as NADH/NADH. However, as NADH and ferredoxin binding sites display homology in some proteins (Hanke et al., 2011), the predicted NADH binding site of NUO-2 might interact with the single-electron carrier ferredoxin. Powered by proton motive force and with the help of the four additional Fe-S clusters in NuoG2 and NuoF2, two of which are of the bacterial [4Fe-4S] ferredoxin type, it might be possible to elevate electrons from the quinol pool not only to the redox potential of NADH, but further to the potential of ferredoxin. Indeed, the ferredoxin-like [4Fe-4S] clusters 6a and 6b of NuoI display a negative shift of their redox midpoint potential of <600 mV when expressed heterologously without incorporation into the NUO complex (Yano et al., 1999), and the NuoI homolog in E. coli is responsible for ferredoxin reduction (Forzi et al., 2005). The extra [4Fe-4S] clusters in NuoF2 could thus be sufficiently electronegative to interact with ferredoxin. Interestingly, in Ca. K. stuttgartiensis a highly similar complex is present that also contains all additional Fe-S clusters in the NuoGF subunits and apparently lacks a FMN binding site. The organization of all corresponding genes in one operon in Kuenenia strongly suggests that their products are subunits of the same enzyme complex. The reduction of ferredoxin by reverse electron transport is important also in Kuenenia, which employs the acetyl-coenzyme A pathway for CO₂ fixation (Strous et al., 2006). The Nuo-2 complex of Kuenenia has been tentatively annotated as formate:quinone oxidoreductase (Strous et al., 2006), but biochemical data endorsing this function is lacking. Future experimental work may reveal whether the NUO-2 complex indeed is a novel mechanism for reverse electron transport spanning the large difference in redox potential between quinol and ferredoxin.

Complex II, the succinate:quinone oxidoreductase (SQR), links the TCA cycle to the quinone pool. This complex can work in both directions. It catalyzes either the oxidation of succinate in the course of the oxidative TCA cycle and oxidative phosphorylation, or the reduction of fumarate to channel electrons into the rTCA cycle (Cecchini et al., 2002). SQRs are classified in five types (A–E; Lemos et al., 2002) and consist of three (type B and E) or four (type A, C, and D) subunits. Many organisms encode a separate succinate dehydrogenase (SDH) and fumarate reductase (FRD), which are expressed differentially under oxic and anoxic conditions, but can also replace each other functionally (Maklashina et al., 1998). SDH and FRD in the same organism usually are similar in sequence and subunit composition, or only one enzyme is present. In contrast, N. gracilis contains two highly different versions of SQRs. One enzyme is of type B that is found also in bacilli and Epsilonproteobacteria, whereas the other one belongs to type E found mainly in Archaea (Lemos et al., 2002). Both types consist of three subunits (Table S1 in Supplementary Material). The type B SQR-1 of N. gracilis consists of a FAD-binding flavoprotein, a Fe-S protein containing all conserved cysteine residues for binding one [2Fe-2S], one [4Fe-4S], and one [3Fe-4S] cluster, and a diheme cytochrome b membrane subunit with five predicted transmembrane helices (Lancaster and Simon, 2002). This SQR type is a good candidate to catalyze the reverse reaction, because it interacts with menaquinone (MQ) rather than ubiquinone (P. Shirawski and Unden, 1998) and MQ has a lower redox potential (E°’ = −74mV) than the succinate/fumarate couple (E°’ = +30mV). In Bacillus subtilis fumarate reduction with menaquinol catalyzed by a type B SQR is electrogenic, so that the reaction contributes to the proton gradient over the cytoplasmic membrane (Schnorpfeil et al., 2001; Lancaster et al., 2008). However, the quinone types present in Nitrospina are not known yet. The type E SQR-2 has some fundamental differences to the type B SQR-1 in its subunit composition and predicted binding sites for prosthetic groups. The large subunit also is a FAD-binding flavoprotein, but it has only low similarity to the corresponding subunit of SQR-1. The Fe-S subunit contains a second [4Fe-4S] cluster instead of the [3Fe-4S] cluster (Gomes et al., 1999), resulting in a slightly lower redox potential of the binuclear center (Lemos et al., 2001). Further, type E SQRs contain an unusual membrane subunit (SdhE) that is unrelated to the membrane anchors of the SQR types A–D. The respective gene is split in the N. gracilis genome sequence, but this may be caused by an uncorrected sequencing error. SdhE does not bind heme b, has only one small putative membrane-spanning segment at the C-terminus, and contains two tandem repeats of the conserved cysteine motif CX₃HX₄CX₅C (Jansen et al., 1997). These motifs bind an additional high-potential [2Fe-2S] cluster that functionally replaces the missing heme b (Iwasaki et al., 2002). Despite these differences, SdhE is essential for membrane attachment of the SQR complex, possibly mediated by amphipathic helices, and (mena-)quinone interaction (Lemos et al., 2002). Although type E SQRs can work reversibly, they appear to preferentially catalyze succinate oxidation in most organisms carrying these enzymes (Gomes et al., 1999). The slightly lower redox potential of the active center allows this SQR to efficiently donate
According to cultivation-based experiments, *N. gracilis* catalyze the reversible oxidation of H$_2$ (Vignais and Billoud, 2007). Members of this group and NAD(P). Characteristic amino acid patterns in the large subunit contains predicted binding sites for one [2Fe-2S] cluster, FAD, electrons to quinone or directly to Rieske-type Fe-S proteins as described by Lücker et al. Genome analysis of *N. gracilis* reveals that most type 3b hydrogenases can reduce electrons to quinone or directly to Rieske-type Fe-S proteins as found in complex III (Janssen et al., 1997). Hence, the SQR-2 of *N. gracilis* is the more likely candidate than SQR-1 for a SDH that is active during the degradation of carbon storage compounds.

Interestingly, *N. gracilis* contains genes for a cytoplasmic NiFe-hydrogenase (Figure 3; Table S1 in Supplementary Material). These genes are organized in a single operon and encode the four subunits of the holoenzyme and six accessory proteins involved in hydrogenase assembly and maturation. Sequence analysis revealed binding sites for nickel in the α and β subunit and for one and three [4Fe-4S] clusters in the δ subunit, respectively. The γ subunit contains predicted binding sites for one [2Fe-2S] cluster, FAD, and NAD(P). Characteristic amino acid patterns in the large subunit clearly classify the enzyme as a type 3b bidirectional (NADP) hydrogenase (Vignais and Billoud, 2007). Members of this group catalyze the reversible oxidation of H$_2$ with NAD(P)$^+$ (Silva et al., 2000). Furthermore, most type 3b hydrogenases can reduce elemental sulfur (S$_0$) or polysulfide to H$_2$. Since they display higher affinities for polysulfides than for H$_2$, they are also referred to as polysulfhydrogenases (Ma et al., 2000). The physiological role of this enzyme in *N. gracilis* is unclear, but three functions appear to be possible: (i) When present, H$_2$ may be oxidized to provide reduced NAD(P)H for biosynthetic processes and energy generation. (ii) Polysulfide (or H$_2$S) can be used as terminal electron acceptor during fermentation of intracellular glycogen deposits in absence of more favorable electron acceptors. (iii) Polysulfide (or S$_0$) can be reduced to H$_2$S for sulfur assimilation during protein biosynthesis if neither sulfate nor sulfite is available. None of these reactions has been reported for *N. gracilis* so far and all three possibilities seem unlikely in the aerobic marine habitat. However, the hydrogenase may be advantageous under hypoxic or even anoxic conditions (see also below).

*N. gracilis* encodes a periplasmic sulfite:cytochrome c oxidoreductase that may couple sulfite oxidation to sulfate with the reduction of two cytochromes (Figure 3; Table S1 in Supplementary Material). In eukaryotes this reaction mainly serves as a detoxification mechanism for sulfite formed during the degradation of sulfur-containing amino acids, but it can be used for energy conservation by sulfur-oxidizing prokaryotes (Kappler et al., 2000). Since *N. gracilis* also has all genes needed for sulfate assimilation via 3'-phosphoadenylyl sulfate and the subsequent reduction to sulfide, the sulfite:cytochrome c oxidoreductase is probably not involved in sulfur assimilation. Hence, *N. gracilis* might be able to use sulfite as alternative energy and/or electron source, but physiological data showing this metabolic capacity are lacking.

### CO$_2$ Fixation and Carbon Metabolism

According to cultivation-based experiments, *N. gracilis* is an obligate autotroph that meets all carbon demands by CO$_2$ fixation (Watson and Waterbury, 1971). Surprisingly, *Nitrospina* employs the rTCA cycle for this task. This CO$_2$ fixation pathway is unexpected in an aerobic organism, because two of its key enzymes belong to the 2-oxocid oxireductase (OR) family whose members normally are highly oxygen-sensitive (Berg, 2011). However, *Hydrogenobacter thermophilus* possesses unusual five-subunit types of 2-oxoglutarate:ferredoxin (OGOR) and pyruvate:ferredoxin oxidoreductase (POR) that are functional under oxic conditions (Yoon et al., 1996, 1997). Moreover, the aerobic nitrite oxidizer Ca. *N. defluvii* also utilizes this pathway using such five-subunit OGOR and POR (Lücker et al., 2010). *N. gracilis* encodes a five-subunit POR with high identity of the subunits (60–76%) to the *Nitrosira* holoenzyme. The OGOR of *N. gracilis* lacks the δ subunit, but the remaining subunits also share high identities with their homologs in *Nitrosira* (53–68%; Table S1 in Supplementary Material). The δ subunit was required for heterologous expression of the *H. thermophilus* holoenzyme in *E. coli*, but it does not contain any known binding motifs for prothetetic groups (Ikeda et al., 2006). A function of this subunit in the expression or assembly of the active OGOR complex might be fulfilled by the δ subunit of POR or by a yet unidentified chaperone. The reduced ferredoxins required by the two ORs could either be provided by the alternative NUO-2 complex (see above) or by a complex involving electron transfer flavoproteins (ETFs). *Clostridium kluveri* can couple the reduction of ferredoxin with NADH to crotonyl-CoA reduction, similar to the reaction in the ETF:quinone reductase in mammals (Herrmann et al., 2008). In this electron bifurcation reaction, the energy derived from the electron transfer from NADH to crotonyl-CoA is used to simultaneously catalyze the unfavorable reduction of a [4Fe-4S] ferredoxin (Buckel and Thauer, 2013). The two FAD-containing ETGs as well as a putative FAD-containing acyl-CoA dehydrogenase necessary are encoded in the *N. gracilis* genome (Table S1 in Supplementary Material), but it appears counterintuitive that *Nitrospina* should rely on the fatty acid degradation pathway for the reduction of ferredoxin during autotrophic growth. Alternatively, the ETF proteins might interact with an unidentified flavin-containing ETF:quinone reductase, that couples the ferredoxin reduction by NADH to the reduction of quinone.

Besides OGOR and POR, the presence of ATP citrate lyase (ACL) is generally considered to be a prerequisite of the rTCA cycle (Hügler et al., 2007). The *N. gracilis* genome contains two CDS with very low similarities to the large and small subunits of known ACLs (Table S1 in Supplementary Material). They contain the characteristic succinyl-CoA synthase-like NAD(P)- and CoA-binding domains and the ATP-grasp fold (Kanao et al., 2001) in the large and small subunit, respectively. The alternative pathway of ATP cleavage in *Aquificaceae* and *Leptospirillum*, which involves citryl-CoA synthase and citryl-CoA lyase (Hügler et al., 2007; Levin et al., 2008), could not be identified in *N. gracilis*. Therefore, we assume that the predicted ACL is functional despite its low similarity to validated ACLs from other organisms.

Each of the other enzymes involved in the reductive and/or oxidative TCA cycles (Berg, 2011) has one homolog in the *N. gracilis* genome (Table S1 in Supplementary Material). There are, however, notable differences to the typical enzyme set for the anaerobic reactions replenishing the oxaloacetate pool: usually, oxaloacetate is formed from pyruvate either in one single step by pyruvate carboxylase or via phosphoenolpyruvate (PEP) by the enzymes pyruvate, water dikinase, and PEP carboxylase. Intriguingly, *N. gracilis* lacks these enzymes, but alternatives for these routes are present. Firstly, instead of pyruvate carboxylase *Nitrospina* appears to employ a membrane-bound oxaloacetate decarboxylase. This enzyme complex usually performs the oxidative decarboxylation of oxaloacetate and directly uses the released
energy for the extrusion of sodium ions across the membrane. In the presence of a sodium gradient, however, the reaction can be reversed to carboxylate pyruvate (Dimroth and Hilpert, 1984). Such dependency on a sodium gradient to drive this reaction might also explain why Nitrospina is restricted to marine habitats. Secondly, the two-step route via PEP can be mediated by the enzymes pyruvate, phosphate dikinase, and the ATP-utilizing PEP carboxykinase. Pyruvate, phosphate dikinase replaces pyruvate, water dikinase in many phototrophic organisms, but usually functions in ATP synthesis from PEP in chemoautotrophic bacteria (Lim et al., 2007). The second enzyme, PEP carboxykinase, is mainly involved in gluconeogenesis by replenishing PEP from oxaloacetate, but can also catalyze the carboxylation of PEP (Zamboni et al., 2004).

Observation by electron microscopy of N. gracilis cells revealed glycogen deposits as intracellular storage compounds (Watson and Waterbury, 1971). Accordingly, automatic genome annotation predicted all genes required for gluconeogenesis and glycogen formation (Figure 3; Table S1 in Supplementary Material). The stored glycogen can be catabolized and used as energy source via the Embden–Meyerhof–Parnas pathway and the oxidative (oTCA). Organotrophic growth of N. gracilis was not observed previously (Watson and Waterbury, 1971). Consistently, no transport systems for the import of organic substrates were identified in the genome except one, merely putative, ABC-type sugar transporter of unknown specificity.

RESISTANCE AND DEFENSE
Although N. gracilis is an aerobic organism, it lacks classical defense mechanisms against reactive oxygen species (ROS) usually found in aerobic bacteria. No homolog of superoxide dismutase (SOD) or reductase (SOR) could be identified in the genome, and catalase appears to be absent, too. Hydrogen peroxide might be degraded by a cytochrome c peroxidase and peroxiredoxins, possibly assisted by the thioredoxin and glutaredoxin systems, which all are encoded in the genome. The protection mechanism against superoxide radicals is more puzzling. Ca. N. defluvii also lacks SOD and catalase, and the presence of a manganese import system has led to speculations that some resistance of this organism against ROS may be conferred by manganese (Lücker et al., 2010). Although the N. gracilis genome contains no known specific manganese import system, protection from ROS in the periplasm might be mediated by a periplasmic multicopper protein that is highly conserved in the Ca. N. defluvii genome, too. This putative multicopper oxidase shows similarity to an enzyme of Bacillus sp. strain SG-1 that is capable of manganese oxidation to MnO2 (van Waasbergen et al., 1996). Since multicopper oxidases are one-electron oxidants, the oxidation of Mn(II) to Mn(IV) proceeds via a Mn(III) intermediate (Spio et al., 2009). Mn(III) can be stabilized in the cell by binding to siderophores (Park et al., 2004) or pyrophosphatase (Tebo et al., 2004), and Mn(III) pyrophosphates contribute up to 100% of the total dissolved manganese pool in marine suboxic zones (Trouwborst et al., 2006). Mn(III) can oxidize O2 to harmless O2-, becoming reduced back to Mn(II) (Barnese et al., 2008; Batinic-Haberle et al., 2011). Interestingly, N. gracilis also has genes for siderophores that may function in Mn(III) stabilization. One could speculate that siderophores loaded with Mn(III) may also be transported into the cell, extending ROS protection from the periplasmic space into the cytoplasm. Here, pyrophosphate for further Mn(III) stabilization could be provided by inorganic phosphatase. Thus, Mn(III) produced by the multicopper oxidase could be stabilized and may contribute to ROS protection. Aside from manganese, polyamines can confer protection against oxidative stress and scavenging free radicals (Chattopadhyay et al., 2003; Tkachenko et al., 2012). As the N. gracilis genome contains the genes needed for producing the polyamines putrescine and spermidine, the organism might also employ this ROS defense mechanism.

N. gracilis appears to be well protected against a plethora of toxic compounds (Table S1 in Supplementary Material). Arsenate reductase (Ji and Silver, 1992) and mercuric reductase (Laddaga et al., 1987) were identified in the genome. Thiosulfate sulfurtransferase and cyanate hydratase convert thioycanate to cyanate and then to ammonia and CO2 for cyanate detoxification (Kamennaya et al., 2008). This mechanism may also allow N. gracilis to use (thio)cyanate as source of ammonium for assimilation. A range of transporters for monovalent copper, diverse mono- and divalent cations, acrilavin, as well as multidrug export systems complete the diverse defense mechanisms of N. gracilis. Multiple Na+ export systems, including an putative eight subunit Mnh-type secondary Na+/H+ antipporter (Swartz et al., 2007), likely contribute to salt tolerance necessary for growth in the marine environment.

DISCUSSION
PHYLGENY AND DIVERSITY OF NITROSPINA
As each gene is shaped by a unique set of functional constraints through evolution, phylogenetic trees based on different single genes, which are calculated for the same set of organisms, may show conflicting topologies (Rokas et al., 2003). Therefore, comparative phylogenetic analyses using different individual marker genes often suffer from incongruent trees that hamper firm conclusions on the affiliation of an organism or even a whole lineage. This problem can be ameliorated by using concatenated datasets of multiple marker genes or proteins, as analyses based on such datasets are not biased toward the evolutionary history of a single marker (Rokas et al., 2003; Strous et al., 2006). This approach and the now available genome sequence of N. gracilis allowed us to assess the phylogenetic affiliation of Nitrospina with an unprecedented high degree of confidence. Contrasting previous phylogenies, which were based on 16S rRNA gene sequences only (Teske et al., 1994), our analyses strongly suggest that N. gracilis does not belong to the Deltaaproteobacteria but represents a distinct, deep-branching line of descent within the Bacteria. This conclusion gains further support from analyses of conserved insertions and deletions in highly conserved genes (Gupta and Griffiths, 2002) and of gene transposition events (Kunisawa, 2006), which are not affected by the quality of sequence alignments and other possible biases of tree calculation methods. For the novel phylogeny we therefore propose the tentative name Nitrospinae phyl. nov. and the reclassification of the family Nitrospiraceae (Garrity et al., 2005) in the Nitrospini class. nov., and Nitrospinales ord. nov., with N. gracilis as the type species. The species description remains as published by Watson and Waterbury (1971). However, the exact position of the Nitrospinae in the bacterial tree could
not be resolved unambiguously. Some trees suggested a position between the *Proteobacteria* and a group consisting of *Nitrospirae* and *Acidobacteria*, whereas others indicated a superphyllum comprising the *Nitrospirae*, *Acidobacteria*, and *Nitrospinae* (Figure 1). Ambiguous branching patterns of underrepresented lineages in phylogenetic trees can sometimes be resolved by adding more reference sequences related to these lineages (Peplies et al., 2008). However, this is currently not possible for *Nitrospinae* since no closely related organisms are known.

**EVOLUTIONARY HISTORY OF NITRITE OXIDATION**

Two main groups of NXR are known which can be distinguished based on their subcellular localization and phylogenetic affiliation within the type II DMSO reductase-family molybdopterin cofactor-binding enzymes. In *Nitrobacter*, *Nitrococcus*, and *Nitrolancetus* NXR faces the cytoplasmic side of the cytoplasmic membrane. This type of NXR is closely related to the respiratory nitrite reductase (NAR) of *E. coli* and many denitrifying bacteria (Lücker et al., 2010; Sorokin et al., 2012). Furthermore, the proteobacterial NOB *Nitrobacter* and *Nitrococcus* contain stacks of intracytoplasmic membranes (ICM), which are densely packed with NXR complexes also facing the cytoplasm (Spieck and Bock, 2005a). Since these ICM resemble those of related phototrophic proteobacteria (purple bacteria), Teske et al. (1994) hypothesized that *Nitrobacter* and *Nitrococcus* evolved from phototrophic ancestors where nitrite was used as electron donor for anoxygenic photosynthesis. Indeed, phototrophic nitrite oxidizers from the genera *Rhodopseudomonas* and *Thiothrix*, which are closely related to *Nitrobacter* and *Nitrococcus*, were discovered recently (Schott et al., 2010). In contrast, *Nitrospira* lack ICM. Their NXR is also attached to the cytoplasmic membrane, but it is oriented toward the periplasm (Spieck et al., 1998; Lücker et al., 2010). Previously, we showed that this periplasmically oriented NXR is phylogenetically distinct from the cytoplasmically oriented enzyme and clusters together with the NXR of the anammox organism *Ca. K. stuttgartiensis* (Lücker et al., 2010). Anammox bacteria oxidize nitrite not for energy generation but to replenish the respiratory chain with electrons needed for CO2 fixation (Jettten et al., 2009). Interestingly, the different subcellular orientation and phylogenetic separation of the two NXR types mirror differences in the nitrite requirements of the respective NOB, which were determined experimentally (Schramm et al., 1999; Bartosch et al., 2002; Freitag et al., 2005; Sorokin et al., 2012). All known NOB with the periplasmic NXR achieve growth with lower nitrite concentrations than NOB with the cytoplasmic enzyme (Sorokin et al., 2012). A likely explanation is the higher efficiency of periplasmic NXR, which conserves more energy per oxidized nitrite. Since the additional O atom in nitrate stems from water, two protons are released in the periplasmic space if nitrite is oxidized by a periplasmic NXR. An additional two protons are consumed in the cytoplasm during O2 reduction to water by the terminal oxidase. Thus, the activity of a periplasmic NXR directly contributes to the proton motive force. This is not the case with a cytoplasmically oriented NXR, where the protons are released and consumed at the same side of the membrane (Lücker et al., 2010). *N. gracilis* lacks ICM and is adapted to low nitrite concentrations (Watson and Waterbury, 1971). Consistently, its two copies of the NxrA subunit are highly similar to the periplasmic NxrA of *Ca. N. defluvii* (54–55% identity). Surprisingly, however, their sequence identity to the NxrA of *Ca. K. stuttgartiensis* is even higher (62%), indicating a closer evolutionary relationship between the NXRs of *Nitrospinae* and anammox bacteria. Indeed, phylogenetic analysis showed that the NxrA of *Nitrospinae* clusters together with the marine *“Ca. Scalindula profunda”* within the anammox group, whereas *Nitrospinae* branches off at the ancestral node of the NXR branch (Figure 5A). This branching pattern implies that the genes coding for the periplasmic NXR were horizontally transferred from an ancestor of *Nitrospinae* to an ancestral anammox organism (or vice versa) and from a (putatively marine) anammox bacterium to *Nitrospinae*. Vertical inheritance can be excluded due to the large phylogenetic distance between these organisms at the inter-phyllum level. The first HGT between ancestral *Nitrospinae* and anammox requires the coexistence of these organisms in an anoxic or hypoxic environment, which was already proposed based on the first analysis of a *Nitrospinae* genome (Lücker et al., 2010). Since *Nitrospinae*-like bacteria often co-occur with anammox organisms in low-oxygen marine environments (Füsel et al., 2012), the second HGT from anammox to *Nitrospinae* appears possible as well. The use of the rTCA cycle for carbon fixation also is fully consistent with a microaerophilic or even anaerobic evolutionary origin of *Nitrospinae*. The periplasmic NXR might originally have evolved in ancestral *Nitrospinae* or anammox bacteria, or it was received by one of these lineages from an unknown donor. As the closest known branch contains bacterial and archaeal nitrate reductases (Figure 5A), one may speculate that such a donor was a nitrate-reducing or nitrite-oxidizing bacterium or archaeon. In this context it appears striking that the reaction of recent periplasmic NXR is reversible and the enzyme still catalyzes both nitrite oxidation and nitrate reduction (Ehrich et al., 1995).

In phylogenetic trees based on the smaller and highly conserved NxrB subunit, *N. gracilis* and *Nitrospinae* cluster together and the anammox lineage branches off first (Figure 5B). This topology is unclear with respect to the aforementioned order of HGT events, because the common ancestral node may represent a member of the anammox lineage, *Nitrospinae*, *Nitrospirae*, or an unknown donor. Additional NxrB sequences from each lineage, once available, might disambiguate the branching order. However, the clustering of the NxrB of *Nitrospinae* and *Nitrospinae* may also reflect similar functions in the respiratory chain. NxrB is more downstream than NxrA and channels electrons to the gamma subunit, NxrC. In both aerobic NOB, the electrons from nitrite are shuttled either toward a terminal oxidase or into a reverse electron transport chain. This bifurcated electron flow from nitrite may require different forms of NxrC, which interact with the respective downstream acceptors. Consistently, the genomes of both *Ca. N. defluvii* and *N. gracilis* contain several different NxrC-like CDS (Lücker et al., 2010 and this study). NxrB must then interact with different types of NxrC. This flexibility may require adaptations in the sequence of NxrB not found in anammox bacteria, where nitrite oxidation is always followed by reverse electron flow. Alternative explanations for the differences between the NxrA and NxrB trees include the possibility that these NXR subunits...
do not share the same evolutionary history but were subject to different HGT events. Interestingly, a member of the _Clostridium_ family also encodes an NXR-like protein that branches between _Nitrospina/Nitrosospira_ and the anammox group (Figure 5B). A closer inspection of its publicly available genome sequence revealed that _C. aerophila_ also carries genes for the TorD-like chaperone and NxrC, but the catalytic NxrA is lacking. Thus, the functions of its NXR-like proteins remain to be investigated.

Phylogenomics (Frickey and Lupas, 2004), further illuminated the extent of HGT between _Nitrospina, Nitrosospira_, and the anammox lineage (represented by _Kuenenia_, Figure 2). A set of genes is highly conserved among all three organisms (Figure 6). In _Ca. K._ stuttgartiensiises these genes form one cluster that codes for the NXR subunits and other proteins that may form a membrane-bound complex involved in nitrite oxidation and electron transfer into the respiratory chain (de Almeida et al., 2011). Exactly these genes are conserved between _Kuenenia_ and _Nitrospira_ (Lücker et al., 2010), and their conservation in _Nitrospina_ strongly supports their importance for nitrite oxidation in these bacteria. Besides the NXR α, β, and γ subunits, these genes encode an alternative NxrC subunit, the TorD-like chaperone, one mono- and one diheme cytochrome c, one putative zinc finger- and two cupredoxin-like copper-binding proteins, and one conspicuous cytochrome bd-like oxidase. The two cytochromes c likely are involved in electron transport from NXR to the terminal cytochrome c oxidase for energy generation and to the cytochrome bc complex for reverse electron transport. Alternatively, the interaction with the terminal oxidase or the bc complex might involve copper instead of heme-based electron carriers. This would explain the presence of genes coding for cupredoxin-like proteins in the cluster (see also de Almeida et al., 2011). The function of the _bd-like_
oxidase is more enigmatic. The predicted protein contains 14 transmembrane helices and shows partial similarity to subunit I of canonical cytochrome bd terminal oxidases. In comparative analyses, the N-terminal parts of the canonical bd and bd-like oxidases align well and the ligands for heme b-binding (Borisov et al., 2011) are conserved. In place of the periplasmically oriented Q-loop found in canonical bd oxidases, however, the bd-like oxidases have three predicted membrane helices, and all residues responsible for quinol binding in the canonical enzyme (Borisov et al., 2011) are lacking. Thus, unless the bd-like oxidases have evolved an alternative quinol binding site, a role comparable to canonical cytochrome bd terminal oxidases is unlikely. However, if the enzymes can interact with quinone they might function like an alternative complex III and enable reverse electron transport from nitrite to the quinone pool, either through direct interaction with NXR, or mediated by the copper- or heme-based electron carriers also encoded in the gene cluster. Interestingly, the gene cluster also contains the gene for the aforementioned multicopper oxidase that might be involved in ROS defense in *Nitrospina* and *Nitrospira* (Figure 6). The lack of this gene in *Ca. K. stuttgartiensis* would be in line with a function in ROS defense, which would be largely obsolete in strictly anaerobic anammox bacteria.

**CONCLUSION**

*N. gracilis* is the most abundant and widespread nitrite oxidizer in marine systems and is of central importance to the oceanic nitrogen cycle. The extended phylogenetic analyses performed in this study have shown that *N. gracilis* represents a novel bacterial phylum. Intriguingly, *N. gracilis* extends an evolutionary model of nitrite oxidation that is based on lateral gene transfer across phylum boundaries, and it strengthens the previously proposed evolutionary link between anammox bacteria and NOB with a periplasmic NXR (Lücker et al., 2010; Sorokin et al., 2012). Like with *Nitrospira*, the use of an oxygen-sensitive CO₂ fixation pathway and the lack of canonical ROS defense mechanisms suggest an evolutionary origin of *Nitrospina* from microaerophilic or even anaerobic ancestors. Consistently, *Nitrospina*-like organisms are frequently encountered in hypoxic marine habitats. However, nitrite accumulation in the truly anoxic core regions of OMZs (Thamdrup et al., 2012) indicates that the observed nitrite oxidation across OMZs,
and also in marine sediments, still requires oxygen and probably is caused by the activity of microaerophilic NOB including *Nitrospina*.

**ACKNOWLEDGMENTS**

We thank Christiane Dorninger and Jasmin Schwarz for help with DNA extraction and for cloning of the NXR operons and Ilia Lakougaard for providing the Perl script for phylome sorting. Martina Grill and Florian Goldenberg are acknowledged for technical support, as is the LABGeM team at Genoscope. This work was supported by the Austrian Science Fund (FWF; grant P24101-B22) and the Vienna Science and Technology Fund (WWTF; grant LS09-40).

**REFERENCES**

Alawi, M., Lipski, A., Sanders, T., Pfeiffer, E. M., and Speck, E. (2007). Cultivation of a novel cold-adapted nitrite oxidizing betaproteobacterium from the Siberian Arctic. ISME J. 1, 256–261.

Altschul, S. F., Gish, W., Miller, W., Myers, E. W., and Lipman, D. J. (1990). Basic local alignment search tool. J. Mol. Biol. 215, 403–410.

Backlund, A. S., Bohlin, J., Gustavsson, N., and Nilsson, T. (2009). Periplasmic cysteine chlorores and chloride reduction in *Idella decleratans*. Appl. Environ. Microbiol. 75, 2439–2445.

Barnese, K., Gralla, E. B., Cabelli, D. E., and Selverstone Valentine, J. (2008). Manganous phosphate acts as a superoxide dismutase. J. Am. Chem. Soc. 130, 4604–4606.

Bartosch, S., Hartweg, C., Speck, E., and Bock, E. (2002). Immunological detection of *Nitrospira*-like bacteria in various soils. Microb. Ecol. 43, 26–33.

Bartosch, S., Wolgast, I., Speck, E., and Bock, E. (1999). Identification of Nitrite-oxidizing bacteria with monoclonal antibodies recognizing the nitrite oxidoreductase. Appl. Environ. Microbiol. 65, 4126–4133.

Batnic-Haberle, I., Rajic, Z., Tovmasyan, A., Reboucas, J. S., Ye, X., Leong, K. W., et al. (2011). Different functions of cationic Mn(III) N-substituted pyridyl-porphyrins, recognized as SOD mimics. Free Radic. Biol. Med. 51, 1035–1053.

Berg, I. A. (2011). Ecological aspects of the distribution of different autotrophic CO₂ fixation pathways. Appl. Environ. Microbiol. 77, 1925–1936.

Biegler, E., and Müller, V. (2010). Bacterial Na⁺-translocating ferredoxin:NAD⁺ oxidoreductase. Proc. Natl. Acad. Sci. U.S.A. 107, 18138–18142.

Borrisov, V. B., Gennis, R. B., Hemp, J., and Verkhovsky, M. I. (2011). The cytochrome bd respiratory oxygen reductases. Biochim. Biophys. Acta 1807, 1398–1413.

Brandt, U. (2006). Energy converting NADH:quinone oxidoreductase (complex I). Ann. Rev. Biochem. 75, 69–92.

Buckel, W., and Thauer, R. K. (2013). Energy conservation via electron bifurcating ferredoxin reduction and proton/Na⁺ translocating ferredoxin oxidation. Biochim. Biophys. Acta 1827, 94–113.

Buschmann, S., Warkentin, E., Xie, H., Langen, J. D., Ermel, U., and Michel, H. (2010). The structure of cbh3 cysteine oxidase provides insights into proton pumping. Science 329, 327–330.

Cavalier-Smith, T. (2006). Rooting the tree of life by transition analyses. Biol. Direct 1, 19.

Cechini, G., Schröder, I., Gunsalus, R. P., and Malkhasiana, D. (2003). Succinate dehydrogenase and fumarate reductase from *Escherichia coli*. Biochim. Biophys. Acta 1553, 140–157.

Chattopadhyay, M. K., Tabo, S., and Tabor, H. (2003). Polymamines protect *Escherichia coli* cells from the toxic effect of oxygen. Proc. Natl. Acad. Sci. U.S.A. 100, 2261–2265.

Cosseau, C., and Batut, J. (2004). Genomes of the ccoNOQP-encoded cbh3 oxidase complex in bacteria. Arch. Microbiol. 181, 89–96.

Daims, H., Nielsen, J. L., Nielsen, P. H., Schleifer, K. H., and Wagner, M. (2001). In situ characterization of *Nitrospira*-like nitrite-oxidizing bacteria active in wastewater treatment plants. Appl. Environ. Microbiol. 67, 5273–5284.

Davis, R. E., Stakes, D. S., Wheat, C. G., and Moyer, C. L. (2009). Bacterial variability within an iron-silica-manganese-rich hydrothermal mound located off-axis at the cleft segment, Juan de Fuca Ridge. Geomicrobiol. J. 26, 570–580.

de Almeida, N. M., Maalcke, W. J., Keljins, J. T., Jen, T. M. S. M., and Kartal, B. (2011). Proteins and protein complexes involved in the biochemical reactions of anaerobic ammonium-oxidizing bacteria. Biochem. Soc. Trans. 39, 303.

DeLong, E. F., Preston, C. M., Minner, T., Rich, V., Hallam, S. J., Frigaard, N.-U., et al. (2006). Community genomics among stratified microbial assemblages in the ocean's interior. Science 311, 496–503.

Dimroth, P., and Hilpert, W. (1984). The structure of the sodium pumps oxaloacetate decarboxylase and proton/Na⁺ translocating NADH:quinone oxidoreductase. Arch. Microbiol. 139, 134–140.

Diverse functions of cationic iron-silica-manganese-rich hydrothermal mound located off Namibian oxygen minimum zone. ISME J. 6, 1200–1209.

Ettwig, K. F., Butler, M. K., Le Paslier, D., Ehrich, S., Behrens, D., Lebedeva, C. G., and Moyer, C. L. (2009). Structural conservation in the [4Fe-4S] clusters of Ech hydrogenase. Proc. Natl. Acad. Sci. U.S.A. 106, 19428–19433.

Elbehti, A., Brasseur, G., and Lemesle-Vicot, A. (2006). Community genetic relationship of *Nitrospira* sp. nov. and its phylogenetic relationship. Arch. Microbiol. 164, 16–23.

Fuchs, B. M., Woebken, D., Zubkov, M. V., Burkill, P., and Amann, R. (2005). Molecular identification of picoplankton populations in contrasting waters of the Arabian Sea. Aquat. Microb. Ecol. 39, 145–157.

Fuchsman, C. A., Kirkpatrick, J. B., Brazelton, W. J., Murray, J. W., and Staley, J. T. (2011). Meta-omic identification of functional communities inferred from biologic and chemical profiles in the Black Sea suboxic zone. FEMS Microbiol. Ecol. 78, 586–603.

Füssel, J., Lamm, P., Lavik, G., Jensen, M. M., Holtappels, M., Gunter, M., et al. (2012). Nitrite oxidation in the Namibian oxygen minimum zone. ISME J. 6, 1200–1209.

Felsenstein, J. (2005). PHYLIP (Phylogeny Inference Package) version 3.6. Seattle: Department of Genome Sciences, University of Washington.

Finn, R. D., Mistry, J., Tate, J., Coggill, P., Eger, A., and Böck, J., et al. (2013). The Pfam protein families database. Nucleic Acids Res. 38, D211–D222.

Forzi, L., Koch, J., Guss, A. M., Radovčič, S. M., and Hedderich, R. (2005). Assignment of the [Fe–Fe] clusters of Ech hydrogenase from Methanosarcina barkeri to individual subunits via the characterization of site-directed mutants. FEMS J. 272, 4741–4753.

Freitag, A., and Bock, E. (1990). Energy conservation in *Nitrobacter*. FEMS Microbiol. Lett. 66, 157–162.

Freitag, T. E., Chang, L., Clegg, C. D., and Prosser, J. I. (2005). Influx of inorganic nitrogen management regime on the diversity of nitrite-oxidizing bacteria in agricultural grassland soils. Appl. Environ. Microbiol. 71, 8323–8334.

Frickey, T., and Lupas, A. N. (2004). PhyloGenie: automated phylogenee generation and analysis. Nucleic Acids Res. 32, 5231–5238.

Table S1 | *Nitrospina gracilis* strain 3/211 proteins with predicted functions in key metabolic pathways.
diversity associated with carbon and nitrogen cycling in permeable shelf sediments. Appl. Environ. Microbiol. 72, 5689–5701.

Ikeda, T., Ochiai, T., Morita, S., Nishiyama, A., Yamada, E., Arii, H., et al. (2006). Anabolic five subunit-type pyruvate:ferredoxin oxidoreductase from Hydrogenobacter thermophilus TK-6. Biochim. Biophys. Acta. 1764, 70–82.

Ilbert, M., and Bonnefoy, V. (2012). Insight into the evolution of the iron oxidation pathways. Biochim. Biophys. Acta. doi:10.1016/j.bbabio.2012.10.001. [Epub ahead of print].

Iwasaki, T., Kounou, A., Aoshima, M., Ohmori, D., Inai, T., Urushiyama, A., et al. (2002). Novel [Fe2S2] type redox center C in SdhC of archaeological respiratory II from Sulfolobus tokodaii strain 7. J. Biol. Chem. 277, 39642–39648.

Jansen, S., Schafar, G., Anнемüller, S., and Moll, R. (1997). A succinate dehydrogenase with novel structure and properties from the hyperthermophilic archaeon Sulfolobus acidocaldarius: genetic and biophysical characterization. J. Bacteriol. 179, 5560–5609.

Jetten, M. S., Nietlik, L., Strous, M., Kartal, B., Keldjens, J. T., and Op Den Camp, H. J. (2009). Biochemistry and molecular biology of anammox bacteria. Crit. Rev. Biochem. Mol. Biol. 44, 65–84.

Ji, G., and Silver, S. (1992). Reduction of arsenate to arsenite by the ArcC protein of the arsenic resistance operon of Staphylococcus aureus plasmid p258. Proc. Natl. Acad. Sci. U.S.A. 89, 9474–9478.

Jorgensen, S. L., Hannisdal, B., Lanzen, A., Baumberger, T., Flesland, K., Fonseca, R., et al. (2012). Correlating microbial community profiles with geochemical data in highly stratified sediments from the Arctic Mid-Ocean Ridge. Proc. Natl. Acad. Sci. U.S.A. 109, E2846–E2855.

Jormakka, M., Richardson, D., Byrne, E. L. L. (2007). Advantages of combining bioinformatic and phylogenetic approaches, metal centres and membrane attachment. Biochim. Biophys. Acta. 1553, 158–170.

Jenkinson, S., Wagner, M., Maass, A., and Parada, P. (2008). Comparative genomic analysis of carbon and nitrogen assimilation mechanisms in three indigenous bioleaching bacteria: predictions and validations. BMC Genomics 9, 581. doi:10.1186/1471-2164-9-581

Lim, K., Read, R. I., Chen, C. C. H., Tempczyk, A., Wei, M., Ye, D., et al. (2007). Swiveling domain mechanism in pyruvate phosphate dikinase. Biochemistry 46, 14845–14853.

Lowe, E. C., Beldner, S., Hartshorne, R. S., Tape, H. L. U., Bridge, E. J., Debieux, C., et al. (2010). Quinol-cytochrome c oxidoreductase and cytochrome c4 medium electron transfer during sulfate respiration in Thiothrix selena. J. Biol. Chem. 285, 18433–18442.

Löcker, S., Wagner, M., Maass, A., Pelletier, E., Koch, H., Vacherie, B., et al. (2010). A Nitrospira metagenome illuminates the physiology and evolution of globally important nitrite-oxidizing bacteria. Proc. Natl. Acad. Sci. U.S.A. 107, 13479–13484.

Ludwig, W. (2004). ARB: a software environment for sequence data. Nucleic Acids Res. 32, 1363–1371.

Ludwig, W. (2010). “Molecular phylogeny of microorganisms is rRNA still a useful marker?”, in Molecular Phylogeny of Microorganisms, eds A. Oren and R. T. Papke (Norfolk: Caister Academic Press), 65–84.

Ma, K., Weiss, R., and Adams, M. W. (2000). Characterization of hydrogenase II from the hyperthermophilic archaeon Pyrococcus furiosus and assessment of its role in sulfur reduction. FEMS Microbiol. Lett. 182, 177–190.

Lancaster, C. R. D., Herzog, E., Juhneke, H. D., Madej, M. G., Müller, F. G., Paul, R., et al. (2008). Electrophoretic and electronic catalysis by diheme-containing succinate-quinone oxidoreductases. Biochim. biophys. Acta. 1786, 996–1000.

Lancaster, C. R. D., and Simon, J. (2002). Succinate-quinone oxidoreductases from Epsilonproteobacteria. Biochim. Biophys. Acta. 1553, 84–101.

Larkin, M. A., Blackshields, G., Brown, N. P., Chenna, R., McGettigan, P. A., McWilliam, H., et al. (2007). Clustal W and Clustal X version 2.0. Bioinformatics 23, 2947–2948.

Lemos, R. S., Fernandes, A. S., Pereira, M. M., Gomes, C. M., and Teixeira, M. (2002). Quinol oxidoreductases and succinate-quinone oxidoreductases: phylogenetic relationships, metal centres and membrane attachment. Biochim. Biophys. Acta. 1553, 158–170.

A., Onsen, P. R. T. Papke (Norfolk: Caister Academic Press), 65–84.

Löcker, S., Wagner, M., Maass, A., Pelletier, E., Koch, H., Vacherie, B., et al. (2010). A Nitrospira metagenome illuminates the physiology and evolution of globally important nitrite-oxidizing bacteria. Proc. Natl. Acad. Sci. U.S.A. 107, 13479–13484.

Ludwig, W. (2004). ARB: a software environment for sequence data. Nucleic Acids Res. 32, 1363–1371.

Ludwig, W. (2010). “Molecular phylogeny of microorganisms is rRNA still a useful marker?”, in Molecular Phylogeny of Microorganisms, eds A. Oren and R. T. Papke (Norfolk: Caister Academic Press), 65–84.

Ma, K., Weiss, R., and Adams, M. W. (2000). Characterization of hydrogenase II from the hyperthermophilic archaeon Pyrococcus furiosus and assessment of its role in sulfur reduction. FEMS Microbiol. Lett. 182, 177–190.

Makishima, E., Berthold, D. A., and Cecchinii, G. (1998). Anaerobic expression of Escherichia coli succinate dehydrogenase: functional
replacement of fumarate reductase in the respiratory chain during anaerobic growth. J. Bacteriol. 180, 5989–5996.

Martinez-Espinosa, R. M., Dridge, E. J., Bonete, M. I., Butt, J. N., Butler, C. S., Sargent, F., et al. (2007). Look on the positive side! The orientation, identification and bioenergetics of ‘Archaeal’ membrane-bound nitrate reductases. FEMS Microbiol. Lett. 276, 129–139.

McDevitt, C. A., Hugenholtz, P., Hanson, G. R., and McSwain, A. G., (2002). Molecular analysis of dimethyl sulfide dehydrogenase from Rhodovulum sulfidophilum: its place in the dimethyl sulfide reductase family of microbial molybdenopterin-containing enzymes. Mol. Microbiol. 44, 1575–1587.

Mincer, T. J., Church, M. J., Taylor, L. T., Preston, C., Karl, D. M., and Delong, E. F. (2007). Quantitative distribution of presumptive archaeal and bacterial nitritifiers in Monterey Bay and the North Pacific Subtropical Gyre. Environ. Microbiol. 9, 1162–1175.

Park, B.-J., Park, S.-J., Yoon, D.-N., et al. (2004). Activity, abundance and diversity of nitrifying archaea and bacteria in the central Californian Current. Environ. Microbiol. 12, 1989–2006.

Sayers, E. W., Barrett, T., Benson, D. A., Bolton, E., Bryant, S. H., Canese, K., et al. (2011). Database resources of the National Center for Biotechnology Information. Nucleic Acids Res. 39, D38–D51.

Sazanov, L. A., and Hinchliffe, P. (2006). Structure of the hydrophobic domain of respiratory complex I from Thermus thermophilus. Science 311, 1430–1436.

Schartl, W. and Unden, G. (1998). Menaquinone-dependent succinate dehydrogenase of bacteria catalyzes reversed electron transport driven by the proton potential. Eur. J. Biochem. 257, 210–215.

Schloss, P. D., and Handelsman, J. (2004). Status of the microbial census. Microbiol. Mol. Biol. Rev. 68, 868–691.

Schmidli, H. A., Strimmer, K., Vingron, M., and Von Haeseler, A. (2002). TREE-PUZZLE: maximum likelihood phylogenetic analysis using quartets and parallel computing. Bioinformatics 18, 502–504.

Schonperf, M., Ianausch, G., Biel, S., Kriger, A., and Unden, G. (2001). Generation of a proton potential by succinate dehydrogenase of Bacillus subtilis functioning as a fumarate reductase. Eur. J. Biochem. 268, 3069–3074.

Schott, J., Griffin, B. M., and Schink, B. (2010). Anaerobic phototrophic nitrite oxidation by Thiothrix sp. strain KS1 and RoseobacterDenudatus sp. strain LQ1. Microbiology 156, 2428–2437.

Schramm, A., De Beer, D., Van Den Heuvel, J. C., Ottengraf, S., and Amann, R. (1999). Microscale distribution of populations and activities of Nitrospina and Nitrospira spp. along a macroscale gradient in a nitrifying biofilm: quantification by in situ hybridization and the use of microsensors. Appl. Environ. Microbiol. 65, 3690–3696.

Schütz, M., Brugna, M., Lebrun, E., Baymann, F., Huber, R., Stetter, K.-O., et al. (2000). Early evolution of cytochrome bc complexes. J. Mol. Biol. 300, 663–675.

Siers, F., Wilm, A., Dineen, D., Gibson, S. J., Karplus, K. L., Li, W., et al. (2011). Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. Mol. Syst. Biol. 7:539. doi: 10.1038/msb.2011.75

Silva, P. J., Van Den Ban, E., Wassen, H., Haaker, H., De Castro, B., Robb, F. E., et al. (2000). Enzymes of hydrogen metabolism in Pyrococcus furiosus. Eur. J. Biochem. 267, 6541–6551.

Sone, N., Sawa, G., Sone, T., and Noguchi, S. (1995). Thermophilic Bacilli have split cytochrome b genes for cytochrome b6 and subunit IV. J. Biol. Chem. 270, 10162–10167.

Sorokin, D. Y., Lücker, S., Vejmelkova, R., Klotz, S., and Amann, R. (1999). Microscale approaches to resolving incongruence in molecular phylogenies. Mol. Phylogenet. Evol. 3, 8–17.

Tolonen, T., Teisala, J., and Voutilainen, E. (2005). Widespread occurrence of nitrite oxidoreductase of Nitrospina: evidence for a nitrite reductase system. Arch. Microbiol. 180, 153–158.

Tavare, G., and Casteel, J. (2007). Improvement of phylogeny after diverging and ambiguously aligned blocks from protein sequence alignments. Syst. Biol. 56, 564–577.

Tatusov, R., Fedorova, N., Jackson, I., Jacobs, A., Kiryutin, B., Koonin, E., et al. (2003). The COG database: an updated version includes eukaryotes. BMC Bioinformatics 4:41. doi:10.1186/1471-2105-4-41

Tebo, B. M., Barg, J. R., Clement, B. G., Dick, G. L., Murray, K. J., Parker, D. E., et al. (2004). Bio- genic manganese oxides: properties and mechanisms of formation. Annu. Rev. Earth Planet. Sci. 32, 247–328.

Teske, A., Ahm, E., Regan, J. M., Toze, S., Rittmann, B. E., and Stahl, D. A. (1994). Evolutionary relationships among ammonia- and nitrite- oxidizing bacteria. J. Bacteriol. 176, 6623–6630.

Thamdrup, B., Dalsgaard, T., and Revsbech, N. P. (2012). Widespread functional anoxia in the oxygen minimum zone of the Eastern South Pacific. Deep Sea Res. Part I Oceanogr. Res. Pap. 65, 36–46.

Tkachenko, A. G., Akhova, A. V., Shumkov, M. S., and Nesterova, L. Y. (2012). Polyamines reduce oxidative stress in Escherichia coli cells exposed to bactericidal antibiotics. Res. Microbiol. 163, 83–91.
Trouwborst, R. E., Clement, B. G., Tebo, B. M., Glazer, B. T., and Luther, G. W. (2006). Soluble Mn(III) in suboxic zones. Science 313, 1955–1957.

Uchiyama, I., Higuchi, T., and Kawai, M. (2010). MBGD update 2010: toward a comprehensive resource for exploring microbial genome diversity. Nucleic Acids Res. 38, D361–D365.

Vallenet, D., Labarre, L., Rouy, Z., Barbe, Vallenet, D., Engelen, S., Mornico, D., Vignais, P. M., and Billoud, B. (2007). Evolutionary and Genomic Microbiology. Frontiers in Microbiology.

van Waasbergen, L. G., Hildebrand, M., Trouwborst, R. E., Clement, B. G., Tebo, B. M., Glazer, B. T., and Luther, G. W. (2005). Communities of Archaea and Bacteria in a subsurface radioactivative thermal spring in the Austrian Central Alps, and evidence of ammonia-oxidizing Crenarchaeota. Appl. Environ. Microbiol. 73, 259–270.

Welte, C., Kallnik, V., Grapp, M., Bender, G., Rapidale, S., and Deppenmeier, U. (2010). Function of Ech hydrogenase in ferredoxin-dependent, membrane-bound electron transport in Methanosarcina mazei. J. Bacteriol. 192, 674–678.

Whelan, S., Forrer, O., and Pecht, I. (2005). Intramolecular electron transfer in nitrite reductases. Chemphyschem 6, 805–812.

Winogradsky, S. (1890). Recherches sur les organismes de la nitrification. Ann. Inst. Pasteur 6, 213–231.

Yano, T., Magnitsky, S., Sed, V. D., Ohnishi, T., and Yagi, T. (1999). Characterization of the putative 2x(4Fe–4S)-binding NQO9 subunit of the proton-translocating NADH-quinone oxidoreductase (NDH-1) of Paracoccus denitrificans. Expression, reconstitution, and EPR characterization. J. Biol. Chem. 274, 28598–28605.

Yoon, K. S., Ishii, M., Igarashi, Y., and Kodama, T. (1996). Purification and characterization of 2-oxoglutarate:ferredoxin oxidoreductase from a thermophilic, obligately chemolithoautotrophic bacterium, Hydrogenovibacter thermophilus TK-6. J. Bacteriol. 178, 3365–3368.

Yoon, K. S., Ishii, M., Kodama, T., and Igarashi, Y. (1997). Purification and characterization of pyruvate:ferredoxin oxidoreductase from Hydrogenovibacter thermophilus TK-6. Arch. Microbiol. 167, 275–279.

Zamboni, N., Maahmeimo, H., Szyperski, T., Hohmann, H.-P., and Sauer, U. (2004). The phosphoenolpyruvate carboxylase in C3 carboxylation at the interface of glycolysis and the TCA cycle of Bacillus subtilis. Metab. Eng. 6, 277–284.

Zaslavsky, D., and Gennis, R. B. (2000). Proton pumping by cytochrome oxidase: progress, problems and postulates. Biochim. Biophys. Acta 1458, 164–179.

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 05 December 2012; paper pending published: 26 December 2012; accepted: 02 February 2013; published online: 21 February 2013.

Citation: Lücker S, Nowka B, Rattei T, Spieck E and Daims H (2013) The genome of Nitrospina gracilis illuminates the metabolism and evolution of the major marine nitrite oxidizer. Front. Microbio. 4:27. doi: 10.3389/fmicb.2013.00027

This article was submitted to Frontiers in Evolutionary and Genomic Microbiology, a specialty of Frontiers in Microbiology. Copyright © 2013 Lücker, Nowka, Rattei, Spieck and Daims. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in other forums, provided the original authors and source are credited and subject to any copyright notices concerning any third-party graphics etc.
## Table A1 | Proteins used for phylogeny.

| Gene   | Protein                | Identifier         | Alignment positions |
|--------|------------------------|--------------------|---------------------|
| rplM   | Ribosomal protein L1   | NIGRAClv1_180012   | 244                 |
| rplB   | Ribosomal protein L2   | NIGRAClv1_280027   | 305                 |
| rplC   | Ribosomal protein L3   | NIGRAClv1_280024   | 288                 |
| rplD   | Ribosomal protein L4   | NIGRAClv1_280025   | 233                 |
| rplE   | Ribosomal protein L5   | NIGRAClv1_280036   | 181                 |
| rplF   | Ribosomal protein L6   | NIGRAClv1_280039   | 165                 |
| rplL   | Ribosomal protein L7/12| NIGRAClv1_280016   | 137                 |
| rplI   | Ribosomal protein L9   | NIGRAClv1_620016   | 164                 |
| rplJ   | Ribosomal protein L10  | NIGRAClv1_280015   | 139                 |
| rplK   | Ribosomal protein L11  | NIGRAClv1_280013   | 156                 |
| rplM   | Ribosomal protein L13  | NIGRAClv1_180012   | 164                 |
| rplN   | Ribosomal protein L14  | NIGRAClv1_280034   | 163                 |
| rplO   | Ribosomal protein L15  | NIGRAClv1_280042   | 230                 |
| rplP   | Ribosomal protein L16  | NIGRAClv1_280031   | 146                 |
| rplQ   | Ribosomal protein L17  | NIGRAClv1_280051   | 150                 |
| rplR   | Ribosomal protein L18  | NIGRAClv1_280040   | 126                 |
| rplS   | Ribosomal protein L19  | NIGRAClv1_380018   | 119                 |
| rplT   | Ribosomal protein L20  | NIGRAClv1_220025   | 114                 |
| rplU   | Ribosomal protein L21  | NIGRAClv1_1040055  | 117                 |
| rplV   | Ribosomal protein L22  | NIGRAClv1_280029   | 111                 |
| rplW   | Ribosomal protein L23  | NIGRAClv1_280026   | 128                 |
| rplX   | Ribosomal protein L24  | NIGRAClv1_280035   | 165                 |
| rpmA   | Ribosomal protein L27  | NIGRAClv1_1040054  | 85                  |
| rpmB   | Ribosomal protein L28  | NIGRAClv1_500015   | 47                  |
| rpmC   | Ribosomal protein L29  | NIGRAClv1_280032   | 44                  |
| rpmF   | Ribosomal protein L32  | NIGRAClv1_740027   | 62                  |
| rpmI   | Ribosomal protein L35  | NIGRAClv1_220024   | 46                  |
| rpsB   | Ribosomal protein S2   | NIGRAClv1_250010   | 224                 |
| rpsC   | Ribosomal protein S3   | NIGRAClv1_280030   | 257                 |
| rpsD   | Ribosomal protein S4   | NIGRAClv1_280049   | 248                 |
| rpsE   | Ribosomal protein S5   | NIGRAClv1_280041   | 149                 |
| rpsF   | Ribosomal protein S6   | NIGRAClv1_620012   | 116                 |
| rpsG   | Ribosomal protein S7   | NIGRAClv1_280020   | 212                 |
| rpsH   | Ribosomal protein S8   | NIGRAClv1_280038   | 170                 |
| rpsI   | Ribosomal protein S9   | NIGRAClv1_180011   | 154                 |
| rpsJ   | Ribosomal protein S10  | NIGRAClv1_280023   | 98                  |
| rpsK   | Ribosomal protein S11  | NIGRAClv1_280048   | 114                 |
| rpsL   | Ribosomal protein S12  | NIGRAClv1_280019   | 151                 |
| rpsM   | Ribosomal protein S13  | NIGRAClv1_280047   | 123                 |
| rpsO   | Ribosomal protein S15  | NIGRAClv1_950034   | 84                  |
| rpsP   | Ribosomal protein S16  | NIGRAClv1_380023   | 66                  |
| rpsQ   | Ribosomal protein S17  | NIGRAClv1_280033   | 71                  |
| rpsR   | Ribosomal protein S18  | NIGRAClv1_620014   | 76                  |
| rpsS   | Ribosomal protein S19  | NIGRAClv1_280028   | 88                  |
| rpsT   | Ribosomal protein S20  | NIGRAClv1_270042   | 110                 |
| rpoA   | RNA polymerase, alpha  subunit | NIGRAClv1_280050 | 336            |
| rpoB   | RNA polymerase, beta subunit | NIGRAClv1_280017 | 1211            |

(Continued)