Archaeal nitrification in the ocean


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Marine Crenarchaeota are the most abundant single group of prokaryotes in the ocean, but their physiology and role in marine biogeochemical cycles are unknown. Recently, a member of this clade was isolated from a sea aquarium and shown to be capable of nitrification, tentatively suggesting that Crenarchaeota may play a role in the oceanic nitrogen cycle. We enriched a crenarchaeote from North Sea water and showed that its abundance, and not that of bacteria, correlates with ammonium oxidation to nitrite. A time series study in the North Sea revealed that the abundance of the gene encoding for the archaeal ammonia monooxygenase alfa subunit (amoA) is correlated with a decline in ammonium concentrations and with the abundance of Crenarchaeota. Remarkably, the archaeal amoA abundance was 1–2 orders of magnitude higher than those of bacterial nitrifiers, which are commonly thought to mediate the oxidation of ammonium to nitrite in marine environments. Analysis of Atlantic waters of the upper 1,000 m, where most of the ammonium regeneration and oxidation takes place, showed that crenarchaeotal amoA copy numbers are also 1–3 orders of magnitude higher than those of bacterial amoA. Our data thus suggest a major role for Archaea in oceanic nitrification.

Crenarchaeota | enrichment culture | North Sea | North Atlantic Ocean | nitrifying bacteria

A rchaea constitute one of the three domains of life next to the Eukaryotes and Bacteria. Until a decade ago, Archaea were thought to mainly consist of organisms thriving in extreme environments such as sulfide hot springs, salt brines, and anoxic environments. With the advent of molecular biological techniques, it became clear that Archaea are actually widespread and occur in diverse environments such as oceans, lakes, and soils (1–4). However, these nonextremophilic Archaea are not closely related to cultured relatives, and thus not much is known about their physiology and role in biogeochemical cycling.

Planktonic Archaea in the ocean comprise both Crenarchaeota and Euryarchaeota, of which the former appears to be the most abundant (1, 5). Marine Crenarchaeota are typically relatively more abundant in deep neritic waters and in the meso- and bathypelagic zones of the ocean (1, 5–7) and are thought to account for ~20% of all prokaryotic cells in the global ocean (1). The metabolism of these planktonic Crenarchaeota is a subject of current debate. In situ labeling (8) and microautoradiography (7) experiments showed that marine Crenarchaeota can use dissolved inorganic carbon as carbon source but are also able to take up amino acids (7, 9), suggesting a heterotrophic lifestyle. Recently, a crenarchaeote, Candidatus “Nitrosopumilus maritimus,” was isolated from a sea aquarium and shown to be autotrophic and able to oxidize ammonium to nitrite (10). Positive correlations between the abundance of Crenarchaeota and nitrite were observed in the Arabian Sea (11) and the Santa Barbara Channel time series (12) and with particulate organic nitrogen in Arctic waters (13). Furthermore, a diverse set of putative archaeal ammonia monooxygenase encoding genes (amoA) were recently reported from shallow (<300 m) marine waters and sediments (14) and was found in fosmid sequences of Crenarchaeum symbiosum (15). These findings hint that Crenarchaeota may be involved in the marine nitrogen cycle, possibly as nitrifiers, but the relevance of crenarchaeotal nitrification for the marine nitrogen cycle is unknown. Until now it has been assumed that marine nitrification is mainly performed by two different groups of bacteria belonging to the β- and γ-proteobacteria (16).

Here, we provide experimental evidence that Crenarchaeota present in seawater are capable of aerobic ammonium oxidation (nitrification) by enrichment of a nitrifying crenarchaeote from coastal waters. We also show by quantitative analysis of both archaeal and bacterial amoA in coastal and open ocean waters that marine Crenarchaeota are likely important players of the present-day marine nitrogen cycle.

Results and Discussion

Enrichment Culture of a Nitrifying Crenarchaeote. We enriched a member of the Crenarchaeota from North Sea waters. The experimental setup was originally designed to examine the temperature adaptation of marine crenarcheotal membrane lipids (see ref. 17 for details). During this experiment, coastal North Sea water was incubated in the dark for 6 months at 25°C in an 850-liter mesocosm tank without addition of nutrients. A substantial increase in archaeal membrane lipids was observed, which coincided with an almost complete consumption of ammonium (17). This initial experiment tentatively suggested a link between North Sea Crenarchaeota and the oxidation of ammonium. To further investigate this possibility, water from this mesocosm tank was incubated in the dark at 22°C and 25°C, and inorganic nutrients, including ammonium, were added while pH was kept constant at 8.2, the regular pH of surface seawater. The abundance of Crenarchaeota was monitored with catalyzed reporter deposition–fluorescence in situ hybridization (CARD-FISH) (18) and revealed a substantial enrichment of crenarcheotal cells at both temperatures (Fig. 1 Lower; see also Fig. 4 and Table 2, which are published as supporting information on the PNAS web site). After a lag phase, crenarchaeotal abundance increased with a doubling time of 2 days to 4–5 × 10⁶ cells per ml in the incubation series and comprised at that time up to 40–70% of DAPI-stainable cells. Bacterial abundance increased during the first 3 days of incubation and then remained at <5% of DAPI-stainable cells during the whole incubation period (Fig. 1 Lower). Generally, 20–30% of the DAPI-stained cells are found to be composed of dead cell material (19), suggesting that our enrichment culture was dominated by Crenarchaeota at day 11 (Fig. 1 Lower). In fact, molecular analyses selective for Archaea revealed

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Abbreviations: AOB, ammonia-oxidizing bacteria; CARD-FISH, catalyzed reporter deposition–FISH; DGGE, denaturing gradient gel electrophoresis; QPCR, quantitative PCR.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database [accession nos. DQ659389–DQ659410 (archaeal 16S rDNA) and DQ784527–DQ784538 (amoA)].

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been nitrifiers and all of the nitrification activity had been mediated throughout the incubation. If all of the detected Proteobacteria had remained low (12318/H20841), the crenarchaeotal population within the ranges reported for bacterial aerobic ammonium oxidation (16) and compare very well with the -proteobacterial cell numbers (cells per milliliter) as determined by CARD-FISH (18). Aged seawater from a large mesocosm experiment (see text and Materials and Methods) with added inorganic nutrients was incubated at two different temperatures in the dark. The data shown are those obtained at 25°C. The crenarchaeotal population in our enrichment culture was shown to consist of a single species, phylogenetically closely related to North Sea Crenarchaeota (Figs. 3–5).

that the incubated waters were dominated by a single member of the crenarchaeotal phylogenetic cluster 1a (6) (see Fig. 5, which is published as supporting information on the PNAS web site) with 99% sequence identity over the nearly complete 16S rRNA gene (1,324 bp) compared with the nitrifying crenarchaeote, Candidatus “N. maritimus” (10). When ammonium levels dropped, nitrite concentrations increased concomitantly with the increase in crenarchaeotal abundance (Fig. 1 Upper), similar to what was previously observed in the large mesocosm tank (see above) (17). Importantly, the abundance of β- and γ-proteobacteria, which include the known nitrifying bacteria, remained low (<1% of DAPI-stained cells) throughout the incubation. If all of the detected Proteobacteria had been nitrifiers and all of the nitrification activity had been mediated by these nitrifiers, the activity per cell would have been >100 fmol of NH₃ cell⁻¹ day⁻¹, substantially more than ever reported. However, if Crenarchaeota were responsible for nitrification, the ammonium conversion rates were between 2 and 4 fmol of NH₃ cell⁻¹ day⁻¹ at 22° and 25°C, respectively. These values are well within the ranges reported for bacterial aerobic ammonium oxidation (16) and compare very well with the ~ 4 fmol of NH₃ cell⁻¹ day⁻¹ for Candidatus “N. maritimus” (estimated from figure 3 in ref. 10).

Further evidence for ammonia oxidation by the enriched marine crenarchaeote was provided by the identification of a single amoA. Its sequence is closely related (91% nucleotide and 98% amino acid sequence identities) to the amoA obtained from the archaean Candidatus “N. maritimus” (10) and to archaeal amoA recovered from the Sargasso Sea (4, 20) (90% nucleotide and 95% amino acid sequence identities; see Fig. 2) and only distantly related to known bacterial amoA sequences. Quantification of the abundance of the amoA and the marine crenarchaeotal 16S rRNA gene at day 7 in the enrichment culture yielded a relative ratio of 0.9:1, suggesting that the enriched crenarchaeote possesses a single copy of amoA. Thus, the distribution and coincidence of single phytypes of amoA and marine crenarchaeotal 16S rRNA gene in relation to the observed changes in nutrient concentrations strongly suggest that the enriched marine crenarchaeota was involved in nitrification. Our result is in agreement with recent findings regarding an ammonium oxidizing crenarchaeote isolated from an aquarium (10) and the presence and expression of amoA in soil Crenarchaeota (21). Importantly, our findings suggest that this metabolism may be widespread among Crenarchaeota thriving in marine waters.

Importance of Archaeal Nitrification in an Ocean Margin System. The ecological significance of the observed crenarchaeotal nitrification was investigated in the coastal waters of the North Sea from which the marine crenarchaeote was enriched. PCR amplification of crenarchaeal 16S rRNA gene by using a general archaeral primer followed by phylogenetic analysis of sequenced denaturing gradient gel electrophoresis (DGGE) fragments revealed that Crenarchaeota dominated the archaean community from late fall to early spring. The recovered 16S rRNA crenarchaeotal gene sequences from the North Sea during the crenarchaeotal winter bloom were closely related to each other and to the enriched crenarchaeote with sequence similarities of >96%, and all belonged to the Group I.1a (6) of the Crenarchaeota (Fig. 5). The abundance of Crenarchaeota varied coincidently with inorganic nitrogen species in North Sea waters: When ammonium levels dropped from 12.7 to 8.5 μM and nitrite levels were rising from 0.8 to 2.5 μM between November and December, crenarchaeotal abundance increased by 2 orders of magnitude (see Fig. 3 Middle). In December, abundances of Crenarchaeota decreased considerably, for reasons presently unclear, whereas ammonium levels remained relatively constant at ~9 μM. In early January, crenarchaeotal abundance again increased substantially by 1 order of magnitude, coinciding with a decrease in ammonium levels from 10 to 3 μM. In contrast to the crenarchaeotal cell abundance, the γ-proteobacterial cell abundance remained high but invariant throughout December to March with 4 ± 1 × 10⁶ cells per ml and did not show a distinct elevation in abundance when ammonium concentrations dropped.

Analyses of amoA genes of the North Sea time series yielded one dominant archaeal amoA gene. This gene was closely related to that of Candidatus “N. maritimus” (92% nucleotide and 97% amino acid identity) (Fig. 2). Importantly, quantification of the archaeal amoA copy numbers by quantitative PCR (QPCR) showed a strong positive linear correlation with both crenarchaeotal cell counts by CARD-FISH (r² = 0.81, n = 9) and marine crenarchaeotal 16S rRNA gene abundance (r² = 0.94, n = 39). The slopes of these correlation lines are 2.5 and 2.8, respectively, suggesting that each crenarchaeotal cell possessed two or three copies of amoA. This amount is significantly more than the single copy of amoA per cell in the enriched crenarchaeote. This finding may reflect either inaccuracies in our QPCR approach or variability in amoA copies within Crenarchaeota similar to ammonia-oxidizing bacteria (AOB) (22). Compared with archaeral amoA, β-proteobacterial amoA were present in equal or slightly higher numbers only during late spring and summer, while amoA of γ-proteobacteria could not be detected at all with our primer set. However, from late fall to early spring, during times of ammonium consumption, bacterial amoA was 1–2 orders of magnitude less abundant than Crenarchaeota-derived amoA (Fig. 3 Bottom). A recent study of ammonium-oxidizing β-proteobacteria in the North Sea reported the same seasonal abundance pattern (23). Thus, our results strongly suggest that the dominant cren-
 startActivityForResult and finish activity through intent.
copy number per cell, which is in between the ratios observed in the North Sea time series and the enriched Crenarchaeota. The recovered amoA sequences fall into two phylogenetic clusters, including the one containing the North Sea enrichment culture amoA sequence and that of Candidatus “N. maritimus” (Fig. 2). Quantification of bacterial amoA gene in the same set of water samples (Table 1) revealed that the amoA gene abundance of β-proteobacteria was lower by 1–3 orders of magnitude compared with archaeal amoA, whereas amoA gene abundance of γ-proteobacteria was below our detection limit. These combined results suggest that Crenarchaeota in the mesopelagic layer of the open ocean are also involved in nitrification and may play a more important role than bacterial nitrifiers.

Our data can be combined with literature data to give a rough estimate of the global importance of archaeal nitrification. Mineralization in the meso- and bathypelagic zones of the ocean [2.2 × 10^{13} mol-Cyr^{-1} (27)] releases ~3.3 × 10^{13} mol-Cyr^{-1}, assuming Redfield stoichiometry (28). If all of the generated ammonium would be oxidized by Crenarchaeota fixing one carbon atom for every ~10 nitrogen molecules oxidized (29), then one would expect an archaeal inorganic carbon fixation rate of ~3.3 × 10^{13} mol-Cyr^{-1}. This estimate is consistent with the estimated rate of global inorganic carbon fixation in the dark ocean by Archaea of 6.6 × 10^{13} mol-Cyr^{-1}, of which ~4.5 × 10^{13} mol-Cyr^{-1} may be taken up by Crenarchaeota (7), assuming that Euryarchaeota and Crenarchaeota are growing at equal rates. Archaeal nitrification thus may be an important process in the biogeochemical cycling of nitrogen in the ocean, although it remains uncertain whether all pelagic Crenarchaeota are nitrifiers. These data, together with the recently established importance of Planctomycetes in denitrification (30, 31) and unicellular cyanobacteria in dinitrogen fixation (32, 33), show the important role of hitherto unrecognized prokaryotes in the oceanic biogeochemical cycling of nitrogen.

**Materials and Methods**

**Incubation Experiment Setup.** Coastal North Sea water was kept in the dark at 25°C in an 850-liter mesocosm tank without addition of nutrients for 6 months (17). After these 6 months, batch cultures with aged mesocosm water were incubated in 20 liters of Clearboy tanks (Nalge Nunc, Rochester, NY) at 22° and 25°C in the dark. Nutrients were added before the incubation at concentrations of 150 μM NaNO₃, 150 μM NH₄Cl, 25 μM NaH₂PO₄, and 2,666 μM NaHCO₃ together with a sterile mix of 12.5 mg/liter yeast, 5 mg/liter peptone extract, vitamins, and trace elements. The pH was regularly adjusted to 8.2 by adding sterile 0.1 M NaOH or HCl, and salinity was maintained at 27 by addition of distilled water. The 20-liter tanks were continuously stirred and left open throughout the experiment, allowing constant gas exchange with the air. Samples for nutrient analysis, CARD-FISH, and DNA were taken every 3–4 days.

**Coastal North Sea Time Series.** The sampling site is situated at the western entrance of the North Sea into the Wadden Sea at the Island Texel (53°00′25″N, 4°78′27″E). Water samples for DNA and CARD-FISH were taken biweekly from August 2002 to July 2003.

**CARD-FISH Analyses.** Fifteen-milliliter water samples were fixed with formaldehyde (final concentration 4%) and stored at 4°C for at least 4 h. Thereafter, the samples were filtered onto 0.2-μm polycarbonate filters (25-mm filter diameter; Millipore, Billerica, MA) with 0.45-μm cellulose nitrate filters (Millipore) as supporting filters and stored frozen at −20°C until further analysis. Total picoplankton were enumerated after DAPI staining (34), and Bacteria and Archaea were enumerated by CARD-FISH (18) under the epifluorescence microscope. The oligonucleotide probes Eub338, BET42, and GAM42 were used for enumeration of Bacteria (35), β- and γ-proteobacteria (23), respectively, and specific probes were applied for the marine Crenarchaeota Group I,
Table 1. Crenarchaeotal cell abundance and archaeal and bacterial amoA copy numbers in water samples from the Atlantic Ocean

<table>
<thead>
<tr>
<th>Sample code</th>
<th>Lat., °N</th>
<th>Long., °E</th>
<th>Depth, m</th>
<th>Cren., cells·ml⁻¹</th>
<th>Arch. amoA, copies·ml⁻¹</th>
<th>β-Prot. amoA, copies·ml⁻¹</th>
<th>γ-Prot., amoA, copies·ml⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1S23</td>
<td>61.682</td>
<td>-16.750</td>
<td>100</td>
<td>1.8 × 10⁴</td>
<td>4.4 × 10⁴</td>
<td>1.4 × 10³</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S32</td>
<td>61.633</td>
<td>-20.187</td>
<td>1016</td>
<td>1.5 × 10⁴</td>
<td>1.2 × 10⁴</td>
<td>5.3 × 10¹</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S34</td>
<td>61.633</td>
<td>-20.187</td>
<td>100</td>
<td>2.8 × 10⁴</td>
<td>1.9 × 10⁴</td>
<td>8.4 × 10²</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S50</td>
<td>60.183</td>
<td>-25.700</td>
<td>150</td>
<td>1.7 × 10⁴</td>
<td>3.6 × 10⁴</td>
<td>8.5 × 10²</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S61</td>
<td>57.453</td>
<td>-27.919</td>
<td>100</td>
<td>3.9 × 10⁴</td>
<td>4.3 × 10⁴</td>
<td>1.5 × 10³</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S71</td>
<td>55.314</td>
<td>-30.432</td>
<td>400</td>
<td>2.7 × 10⁴</td>
<td>0.5 × 10⁴</td>
<td>6.7 × 10⁴</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S72</td>
<td>55.314</td>
<td>-30.432</td>
<td>150</td>
<td>3.1 × 10⁴</td>
<td>1.9 × 10⁴</td>
<td>3.6 × 10²</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S82</td>
<td>52.667</td>
<td>-34.167</td>
<td>100</td>
<td>2.2 × 10⁴</td>
<td>0.8 × 10⁴</td>
<td>1.7 × 10¹</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S83</td>
<td>55.200</td>
<td>-26.134</td>
<td>148</td>
<td>3.8 × 10⁴</td>
<td>3.1 × 10⁴</td>
<td>1.2 × 10³</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S120</td>
<td>49.734</td>
<td>-26.134</td>
<td>600</td>
<td>8.0 × 10³</td>
<td>3.0 × 10⁴</td>
<td>2.0 × 10²</td>
<td>n.d.</td>
</tr>
<tr>
<td>T1S121</td>
<td>49.734</td>
<td>-26.134</td>
<td>148</td>
<td>3.8 × 10⁴</td>
<td>3.1 × 10⁴</td>
<td>1.2 × 10³</td>
<td>n.d.</td>
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<tr>
<td>T1S154</td>
<td>41.600</td>
<td>-26.333</td>
<td>150</td>
<td>6.9 × 10³</td>
<td>3.6 × 10⁴</td>
<td>1.7 × 10³</td>
<td>n.d.</td>
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<tr>
<td>T2S14</td>
<td>40.104</td>
<td>-66.498</td>
<td>95</td>
<td>8.4 × 10⁴</td>
<td>2.4 × 10⁴</td>
<td>1.7 × 10³</td>
<td>n.d.</td>
</tr>
<tr>
<td>T2S25</td>
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<td>-62.433</td>
<td>95</td>
<td>2.7 × 10⁴</td>
<td>0.9 × 10⁴</td>
<td>2.5 × 10⁵</td>
<td>n.d.</td>
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<tr>
<td>T2S26</td>
<td>43.303</td>
<td>-58.083</td>
<td>100</td>
<td>2.1 × 10⁴</td>
<td>3.7 × 10⁴</td>
<td>9.8 × 10³</td>
<td>n.d.</td>
</tr>
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<td>T2S28</td>
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<td>-49.250</td>
<td>100</td>
<td>9.4 × 10³</td>
<td>2.0 × 10⁴</td>
<td>5.6 × 10²</td>
<td>n.d.</td>
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<tr>
<td>T2S27</td>
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<td>-45.067</td>
<td>243</td>
<td>2.9 × 10⁴</td>
<td>1.6 × 10⁴</td>
<td>1.4 × 10³</td>
<td>n.d.</td>
</tr>
<tr>
<td>T2S24</td>
<td>45.567</td>
<td>-45.067</td>
<td>100</td>
<td>5.2 × 10⁴</td>
<td>3.2 × 10⁴</td>
<td>3.9 × 10²</td>
<td>n.d.</td>
</tr>
<tr>
<td>T2S26</td>
<td>45.567</td>
<td>-45.067</td>
<td>100</td>
<td>5.2 × 10⁴</td>
<td>3.2 × 10⁴</td>
<td>3.9 × 10²</td>
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<tr>
<td>T2S84</td>
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<td>-42.217</td>
<td>266</td>
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<td>1.0 × 10⁴</td>
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<td>n.d.</td>
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<tr>
<td>T2S84</td>
<td>47.300</td>
<td>-42.217</td>
<td>103</td>
<td>1.0 × 10⁴</td>
<td>5.5 × 10⁴</td>
<td>1.7 × 10³</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

Crenarchaeotal cell abundance was determined by CARD-FISH, and archaeal and bacterial amoA copy numbers were determined by QPCR in water samples from the Atlantic Ocean. Samples were taken during the TRANSAT-1 and -2 cruises (7). Lat., latitude; Long., longitude; Cren., Crenarchaeota; Arch., Archaeal; prot., Proteobacterial.

* n.d., not detected, i.e., no specific products were formed upon QPCR or they were below the detection limit of ∼2.5 × 10² copies·ml⁻¹ as determined for γ-proteobacterial amoA.

Cren537 (5'–TGACCACTTGAGGTGCTG-3') (18). All probes were tested for their specificity before the study. Cell walls were permeabilized with lysozyme (Sigma, St. Louis, MO; 10 mg/ml in 0.05 M EDTA, 0.1 M Tris-HCl, pH 8) for Eub338 (18) or with proteinase-K for Cren537 ([Sigma; 1.844 units/mg, 10.9 mg/ml]; 0.2 μl/ml in 0.05 EDTA, 0.1 M Tris-HCl, pH 8) at 37°C for 1 h. Probe working solution (50 ng/μl) was added at a final concentration of 2.5 ng/μl. Hybridization was performed at 35°C for 8–12 h. Negative control counts (hybridization with HRP-Non338) averaged 1.5%. The average counting error in cell abundances for the enrichment culture was 29% for DAPI staining, 29% for Crenarchaeota, 40% for bacteria, and 92% for β- and γ-proteobacteria. For the North Sea time series the average counting error in cell abundance was 18% for DAPI staining, 42% for Crenarchaeota, and 28% for β- and γ-proteobacteria. The larger errors are usually associated with low cell numbers where slides contained substantially less than 200 cells (e.g., β- and γ-proteobacteria in the incubation experiments).

Total DNA Extraction. For the QPCR and phylogenetic analysis, 1 liter of coastal North Sea water or water from the incubation experiments was filtered through a 0.2-μm-pore-size polycarbonate filter and total DNA was extracted as described (17). The efficiency of cell lysis was ∼90% as determined by counting the percentage of DAPI-stained cells that remained in suspension or were still intact and attached to the filter or zirconium beads after the lyses steps during our extraction procedure. This whole procedure was performed three times per sample.

QPCR. The numbers of archaeal 16S rRNA, archaeal amoA, as well as bacterial amoA gene copy numbers in all samples were determined in duplicate by using an iCycler system (Bio-Rad, Hercules, CA). A total of 40 cycles were run with PCR conditions and reagents as described (36) but with annealing temperatures and primer combinations as listed in Table 3, which is published as supporting information on the PNAS web site. Fluorescently measured (Picogreen, Molecular Probes, Carlsbad, CA) known amounts (10 ng) of template DNA were added to the reaction mixtures. Accumulation of newly amplified double-stranded gene products was followed online as the increase in fluorescence due to the binding of the fluorescent dye SYBRGreen (Molecular Probes). Calibration of the samples was performed with known copies (between 10⁻⁶ and 10⁷) of Sulfolobus acidocaldarius DSMZ 639 (archaeal 16S rDNA), enriched marine Crenarchaeote from the North Sea (archaeal amoA), Nitrosomonas europaea (amoA of β-proteobacterial AOB), and Nitrosooccus oceanus (amoA of γ-proteobacterial AOB), which were generated during PCR with the same primers as used for the amplification of the environmental genes (Table 3). As a control of the specificity of the QPCR, the runs were repeated with only 32 cycles so that most amplicons reached the threshold cycle. In addition, 1 μl of the first reaction with 32 cycles was added to a fresh mixture of PCR ingredients and run for 12–15 cycles but this time with primers including the 40-bp-long GC clamp to allow subsequent DGGE analysis (36). Aliquots of these QPCR products were run on an agarose gel to identify unspcific PCR products such as primer dimers or fragments with unexpected fragment lengths (Table 3). Sequence analysis of the excised DGGE fragments (see below) revealed the diversity of the amplicons generated by QPCR and therefore was the ultimate proof that the QPCRs were in fact specific.

Phylogeny of Sequenced DGGE Fragments. Archaeal 16S rDNA amplicons were analyzed by DGGE as described (37). Archaeal amoA amplicons were analyzed by DGGE by using a similar protocol with the exception that the DGGE was run for 3 h, and the gradient of denaturants was 10–50%. To check the specificity of QPCR of β-proteobacterial AOB, amoA amplicons from β-proteobacterial AOB were analyzed by DGGE as described recently (38), and sequencing of DGGE fragments revealed the specificity of the QPCRs despite the low copy numbers. The community structure of β-proteobacterial AOB in the North Sea
clearly differed from that in the Atlantic Ocean (data not shown).

DGGE fragments were excised from the gels and subsequently sequenced for phylogenetic comparison with reference sequences from GenBank (39) by using the ARB software package (40). Archaeal 16S rDNA and amoA sequences obtained in this study have been deposited in GenBank under accession nos. DQ659389–DQ659410 and DQ784527–DQ784538, respectively.

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