Autotrophic carbon fixation strategies used by nitrifying prokaryotes in freshwater lakes

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One sentence summary: CO₂ fixation strategies used by nitrifiers were investigated in lakes, suggesting that Thaumarchaeota using the 3-hydroxypropionate/4-hydroxybutyrate pathway were dominant in deep and oligotrophic lakes, whereas bacteria employing the Calvin cycle were important in samples with an elevated nutrient status.

ABSTRACT

Niche specialization of nitrifying prokaryotes is usually studied with tools targeting molecules involved in the oxidation of ammonia and nitrite. The ecological significance of diverse CO₂ fixation strategies used by nitrifiers is, however, mostly unexplored. By analyzing autotrophy-related genes in combination with amoA marker genes based on droplet digital PCR and CARD-FISH counts targeting rRNA, we quantified the distribution of nitrifiers in eight stratified lakes. Ammonia oxidizing (AO) Thaumarchaeota using the 3-hydroxypropionate/4-hydroxybutyrate pathway were dominant in deep and oligotrophic lakes, whereas Nitrosomonas-related taxa employing the Calvin cycle were important AO bacteria in smaller lakes. The occurrence of nitrite oxidizing Nitrospira, assimilating CO₂ with the reductive TCA cycle, was strongly correlated with the distribution of Thaumarchaeota. Recently discovered complete ammonia-oxidizing bacteria (comammox) belonging to Nitrospira accounted only for a very small fraction of ammonia oxidizers (AOs) present at the study sites. Altogether, this study gives a first insight on how physicochemical characteristics in lakes are associated to the distribution of nitrifying prokaryotes with different CO₂ fixation strategies. Our investigations also evaluate the suitability of functional genes associated with individual CO₂ assimilation pathways to study niche preferences of different guilds of nitrifying microorganisms based on an autotrophic perspective.

Keywords: chemoautotrophs; nitrifiers; lakes; CO₂ fixation pathways
INTRODUCTION

Nitrifiers are chemolithoautotrophs, which are defined by their ability to use reduced inorganic nitrogen compounds as an energy source and inorganic carbon to fulfill the carbon need. Until recently, the two-step nitrification process has been considered to be catalyzed by two separate groups: ammonia oxidizing (AO) organisms, which include ammonia-oxidizing archaea (AOA, Könneke et al. 2005; Hatzenpichler 2012), ammonia-oxidizing bacteria (AOB, Kowelchuk and Stephen 2001) and nitrite-oxidizing bacteria (NOB). However, the discovery of bacteria that catalyze complete nitrification (complete ammonia oxidizers; ‘comammox’), members of the genus Nitrospira, has fundamentally expanded our view of the nitrification process (Daims et al. 2015; van Kessel et al. 2015). The first and rate-limiting step in nitrification, the oxidation of ammonia to hydroxylamine, is catalyzed by the enzyme ammonia monoxygenase (AMO). As all known bacterial and archaeal ammonia oxidizers harbour AMO, the gene encoding the alpha subunit of this enzyme (amoA) has become a well-established functional marker to analyze the distribution, diversity and ultimately the niche preferences of AOA and AOB in the environment (e.g. Bouskill et al. 2012; Meinhardt et al. 2015).

On the other hand, the ecological importance of different carbon fixation strategies used by nitrifying prokaryotes has not been paid much attention. This is quite surprising, considering the diversity of biochemistries of CO2 assimilation pathways that also might influence the distribution of nitrifiers in the environment. To date, six carbon assimilation pathways are known, whereof three are used by nitrifying prokaryotes (Berg 2011). The Calvin-Benson-Bassham (CBB) cycle is present in different genera of NOB including Nitrobacter, Nitrococcus, Nitrotroga and Nitrolancea (Daims, Lückner and Wagner 2016) and generally found in AOB belonging to Proteobacteria (Badger and Bek 2008). The enzyme responsible for the actual fixation of CO2 in the CBB cycle, ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco), occurs in different forms characterized by specific catalytic properties (Berg 2011). Most AOB possess form IA Rubisco. This enzyme is known for its poor catalytic affinity for CO2, which is in some AOB (e.g. Nitrosomonas europaens C91) compensated by carbon concentrating mechanisms that support better growth at low CO2 concentrations (Stein et al. 2007; Badger and Bek 2008). Form IC Rubisco is present in several AOB affiliated with different Nitrospira species and the Gammaproteobacterium Nitrosococcus oceani. Several AOB, including Nitrosomonas sp. Is79 and Nitrosomonas sp. AL212, even encode two copies of the Rubisco operon in their genomes (Bollmann et al. 2013). However, there is a clear lack of environmental studies addressing the ecological adaptation and specific niches occupied by AOB using different forms of Rubisco. A variant of the 3-hydroxypropionate/4-hydroxybutyrate (HP/HB) cycle operates in AOA (Könneke et al. 2014). This pathway was described as the most energy efficient aerobic carbon fixation cycle, which fits in well with the adaptation of AOA to nutrient-limited conditions. For environmental investigations, molecular tools targeting genes coding for the key enzymes 4-hydroxybutyryl-CoA dehydratase and acetyl-CoA/propionyl-CoA carboxylase are highly specific instruments to explore the diversity of autotrophic Thaumarchaeota using the HP/HB cycle for CO2 fixation (e.g. Yakimov, La Cono and Denaro, 2009, 2011; Bergauer et al. 2013; Hu et al. 2013; La Cono et al. 2013; Tolar, King and Hollibaugh 2013; Alfreider et al. 2017). Nitrifying bacteria using the reductive citric acid cycle (rTCA) are found in members of the genus Nitrospira, which include NOB and comammox (Daims, Lückner and Wagner 2016). In marine systems, the rTCA cycle is employed by the nitrite oxidizing Nitrospira species (Pachiadaki et al. 2017). The rTCA cycle is a reversal of the oxidative citric acid cycle (Krebs cycle) and forms acetyl-CoA from two CO2 (Berg 2011). This carbon fixation strategy was originally known to occur in Epsilonproteobacteria and Aquificae in anaerobic and microaerobic environments, due to the oxygen sensitivity of key enzymes in the cycle (Hügler and Sievert 2011). However, for Nitrospira it has been shown that enzymatic adaptations strengthen the O2 robustness of the rTCA cycle that allows the pathway to function also in aerobic habitats (Berg 2011). In environmental samples, the rTCA cycle is usually detected by targeting genes coding for the alpha or beta subunit of the ATP citrate lyase and the alpha subunit of 2-oxoglutarate:ferredoxin oxidoreductase enzymes (Hügler and Sievert 2011; Kovaleva et al. 2011; Noguerola et al. 2015; Alfreider et al. 2017).

The goal of this research has been developed based on the results of a former study, where the diversity of sequences coding for selected key enzymes in the HP/HB, CBB and rTCA cycle in six lakes were analyzed (Alfreider et al. 2017). In that study, the authors demonstrated that a significant part of the sequences was related to nitrifiers, suggesting that nitrification is a major source of energy for chemoaotrophs in these lakes. Specifically, sequences affiliated with the genus Nitrospira and Thaumarchaeota, using the rTCA and HP/HB cycle respectively, were mostly found in deep lakes. Rubisco form IA genes, related to members of the N. oligotropha lineage (cluster 6A) have been detected in different lake types and depths. However, as the study of Alfreider et al. (2017) was mostly based on sequence analysis from selected samples, the abundance, distribution and consequently the ecological niche preferences of different guilds of nitrifiers remained unexplored. For the present work, different strategies were developed in order to quantify the distribution of three CO2 fixation pathways of AOA, AOB and NOB in a variety of eight stratified lakes of different sizes and environmental characteristics. These ecosystems are characterized by distinct and stable concentration gradients of oxygen and different redox states of nitrogen, thus allowing the investigation of nitrifying prokaryotes in the ecological framework of measurable habitat heterogeneity. In this respect, we expect a vertical niche separation of nitrifying Thaumarchaeota and bacteria with different energy and substrate requirements, which is also linked to their different strategies for CO2 fixation. Digital droplet PCR (ddPCR) with specifically designed primers were applied to target the most abundant clades of sequences affiliated with nitrifiers in lakes. These values were compared with ddPCR derived amoA gene abundances of AOB and AOA, which also includes the analysis of comammox-Nitrospira based on a recently developed qPCR assay (Pjevac et al. 2017). Furthermore, catalyzed reporter deposition fluorescence in situ hybridization (CARD-FISH) enabled the microscopic analysis based on the taxonomic affiliation of dominant nitrifiers.

MATERIALS AND METHODS

Field work and chemical analysis

Eight lakes (Attersee-ATT, Hallstättersee-HAL, Millstätter See-MIL, Traunsee-TRA, Faaksee-FAA, Irsee-RR, Mondsee-MON, Weißensee-WEI), all of them located in Austria, were chosen based on their different thermal and chemical stratification patterns. The geographical location and the key morphometric and
At least 400 DAPI stained cells were counted with a Zeiss Axioplan epifluorescence microscope.

Probe Nitro878 was designed in ARB (Ludwig et al. 2004) using the SILVA database LSU Ref 123 (Pruesse et al. 2007). A bootstrapped maximum likelihood tree (GTR-GAMMA model) of all 235 rRNA sequences affiliated with Nitrosomonas (Fig. 53, Supporting Information) served as backbone for probe design with the ARB tools probe_design and probe_check. The resulting probe targets all publically available sequences affiliated with the target group (Fig. 53, Supporting Information), but has also nine outgroup hits within Xanthomonadaceae (Greekproteobacteria). However, these microbes were obtained from non-aquatic habitats and can be thus neglected when using the probe for lake samples only. The probe and its competitor oligonucleotide were evaluated in silico with the web-tool mathFISH (Yilmaz, Parniske et al. 2007). A bootstrap-based alignment of the target group (Fig. S3, Supporting Information) served as backbone for probe design with the ARB tools probe_design and probe_check. The resulting probe targets all publically available sequences affiliated with the target group (Fig. S3, Supporting Information). CARD-FISH and probe design

CARDFISH of water samples was done following the protocol of Wendeberg (2010). In brief, water samples were fixed with 0.2 μm filtered formaldehyde (2% final concentration) and between 20 and 50 mL were filtered onto 0.2 μm polycarbonate filters (Poretics, 47 mm filter diameter) and stored at −20 °C until use. Before hybridization, filters were embedded in 0.1% ultrapure agarose (wt./vol., SeaKem LE Agarose, Lonza, Basel, Switzerland). Filters used for hybridization with bacterial oligonucleotide probes were incubated in lysozyme solution for 55 min at 37 °C. Filters hybridized with archaeal oligonucleotide probes were pre-treated in lysozyme solution (see above), but with the addition of proteinase K (75 μL Proteinase K stock 1:100) for 35 min at 37 °C. Taxonomic specificity, sequence information and hybridization conditions (formamide concentration) of HRP-labeled oligonucleotide probes (Biomers.net, Germany) are listed in Table S2 (Supporting Information). Alexa488 tyramides (Thermofisher) were used for signal amplification. The filters were counterstained with 4,6-Diamidin-2-phenylindol (DAPI) and embedded in a 5:1:1 mix of Citifluor (Citifluor Ltd., London), Vectashield (Vector Laboratories, Inc., Burlingame, CA) and PBS.

DNA-extraction and droplet digital PCR

Lake water samples (800 - 1170 mL) were filtered through polyethersulfone membrane filters (0.22 μm pore size and 47 mm diameter; Merck Millipore Ltd., Ireland) and stored at −20 °C until use. DNA was extracted with the PowerWater©DNA Isolation Kit (MO BIO Laboratories Inc., USA) according to the manufacturer’s protocol. A Quantus Fluorometer (Promega Corporation, USA) and QuantiFluor©dsDNA chemistry (Promega Corporation, USA) was used to measure the DNA content in the samples. Quantitative PCR analysis was performed using a QX200 Droplet Digital PCR system (Biorad) in combination with an automated droplet generator (AutoDG Instrument, Biorad). DdPCR reactions were set up with QX200 ddPCR EvaGreen Supermix (Biorad) to a final volume of 20 μL in 96-well plates following the manufacturer’s instructions. Primer pairs used for qPCR targeting genes coding for key enzyme 4-hydroxybutyryl-CoA dehydratase in the HP/HB cycle were published in Alfreider et al. (2017). Primers targeting aclA genes of Nitrospira in the rTCA cycle and the putative cbbL form IA genes of the N. oligotropha
Cluster in the Calvin cycle were designed for this study (Table S1, Supporting Information). The optimal annealing temperature for both primer pairs was determined empirically with DNA extracted from samples of different lakes. Specification of all other primers used are also listed in Table S1 (Supporting Information). Optimal primer concentration and annealing temperature for most primers was (re)evaluated for ddPCR based on different primer concentrations and temperature gradient experiments. After automated droplet generation using the standard protocol provided by the manufacturer, PCR plates were heat sealed (Pierceable Foil Heat Seal, Biorad) and placed in a T100 thermal cycler (Biorad) for PCR amplification using the following cycling conditions: initial enzyme activation step of 5 min at 95 °C, followed by 40 cycles including 30 s denaturation at 95 °C, 30 s of primer annealing at primer specific temperatures (see Table S1, Supporting Information) and 1 min of primer extension at 72 °C. Signal stabilization of the reaction was accomplished by final steps at 4 °C for 5 min and 90 °C for 5 min. A 2.5 °C/sec ramp rate was used to guarantee each droplet reaches the correct temperature for each step. For signal measurement, the plates were placed into the reader and droplets were examined according to manufacturer’s recommendations. Raw data were further analyzed using QuantaSoft Software 1.7.4. (Biorad). As the recommended dynamic range of the ddPCR system is from 1 to 120 000 copies of the target molecule/20 μL reaction, samples containing over 100 000 copies were diluted accordingly and analyzed again. Quality check included non-template controls, the evaluation of the fluorescence amplitude of positive and negative droplets and the examination of the reliability of the automated threshold settings by the QuantaSoft software.

**Sequence analysis for the evaluation of newly designed qPCR primers**

In order to test the coverage and specificity of the newly designed qPCR primers pairs q_cbbL_IA_Nit and q_aclA_Nit.
Figure 2. Vertical distribution of autotrophy-related (hcd, cbbL, and aclA) and amoA gene copy numbers. Please note the logarithmic scale in the left side plots. The grey shaded area represents the approximate dimension of the metalimnion.
Figure 3. Box plots summarizing gene copy abundances (panels A and B) and CARD-FISH counts (panels C and D) in lakes dominated by AOA (panels A and C) and AOB (panels B and D). Note the logarithmic scale on the x-axis in panels A and B. Probe Ntspa0476 was originally designed to specifically detect \textit{Ca. N. nitrosa} and \textit{Ca. N. nitrificans} (van Kessel et al. 2015), but the probe also covers potential NOB. The vertical distribution of functional marker genes numbers and CARD-FISH counts are shown in detail in Fig. 2 and Fig. S1 (Supporting Information).

Table 2. Spearman rank correlation comparing different primers and probes used for the detection of specific guilds of nitrifiers. The taxonomic coverage is given in Tables S1 and S2 (Supporting Information).

| Variable | hcd | aclA | cbbL | amoA | amoA | comaA | comaB | Ntspa476 | Ntspa662 | Nso1225 |
|----------|-----|------|------|------|------|-------|-------|----------|----------|----------|
| hcd      |     |      |      |      |      |       |       |          |          |          |
| aclA     | 0.71** |      |      |      |      |       |       |          |          |          |
| cbbL     | -0.29 | -0.06|      |      |      |       |       |          |          |          |
| amoA     | 0.98** | 0.74**|      |      |      |       |       |          |          |          |
| amoA     | 0.12 | 0.27 | 0.54**| 0.12|      |       |       |          |          |          |
| comaA    | 0.21 | 0.18 | 0.15 | 0.18 | 0.16|       |       |          |          |          |
| comaB    | -0.01 | -0.03 | 0.24 | -0.01 | 0.15 | 0.45**|       |          |          |          |
| Ntspa476 | 0.01 | -0.03 | -0.07 | -0.02 | -0.08 | -0.10 | -0.11 |          |          |          |
| Ntspa662 | 0.53** | 0.15 | -0.33 | 0.48**| -0.11 | 0.00 | -0.10 | 0.27     |          |          |
| Nso1225  | -0.14 | 0.00 | 0.47**| -0.17 | 0.29 | -0.01 | -0.03 | 0.15 | 0.25     |          |
| MGI-535  | 0.67** | 0.54**| -0.22 | 0.66**| 0.05 | 0.11 | -0.09 | -0.06 | 0.38** | -0.11    |

Significance level: *p<0.01, p<0.001**

AOA AOB NOB \textit{Nitrosira} Comammox \textit{Nitrospira}

(Table S1, Supporting Information), sequence analysis of selected PCR-products from different lakes was performed (results are shown in Figs 4 and 5). All PCR products selected for sequencing analysis were separated on 1.5% agarose gels. Bands with proper size were selected for subsequent cloning, cut out of the gel and purified using a MinEluteVR Gel Extraction Kit (Qiagen Inc., Valencia, CA). PCR products were ligated into pGEM-T-Easy Vector plasmid (Promega, Madison, WI) and transformed into JM109 competent cells following the manufacturer’s instructions. Clones were screened for the presence of
Figure 4. Evaluation of the specificity and coverage of qPCR primers targeting aclA genes in Nitrospira based on amino acid sequence analysis (shown in black). DNA was extracted from lake water samples obtained from this (ATT, TRA) and a previous study (Achensee-ACH, Starnberger See-STA; Alfreider et al. 2017). Numbers in brackets indicate the number of clones analyzed. Closest NOB-Nitrospira aclA genes are shown in red color, comammox-Nitrospira are designated in blue.

proper inserts by PCR using vector-specific primers M13-F/R and GoTaqVR G2 Hot Start Master Mix (Promega, Madison, WI) following the protocol provided by the manufacturer. Selected reactions were Sanger sequenced by a sequencing service enterprise (Eurofins MWG Operon, Ebersberg, Germany).

Closest relatives to nucleotide sequences and deduced amino acid sequences were obtained using NCBI’s sequence similarity search tools BLASTN and BLASTP (https://blast.ncbi.nlm.nih.gov/Blast.cgi). Deduced amino acids were aligned using MUSCLE algorithm as implemented in MEGA 6.0 software (Tamura et al. 2013), followed by visual inspection of the alignment. Neighbor-Joining trees applying gamma distribution as the distance method were also computed with the MEGA 6 software package. Bootstrap analysis (1000 replicates) was used to obtain confidence estimates for tree topology. The phylogenetic tree was condensed by compressing subtrees with highly similar sequences.

Sequence data deposition
Sequences data have been submitted to GenBank databases under accession numbers MG600595 - MG600653 (acl) and MG600654 - MG600709 (cbbl-Form IА).

RESULTS AND DISCUSSION
Lake characteristics
Vertical profiles of temperature revealed that at the time of sampling all lakes were stratified, with a thermocline established in different depths of each lake (Fig. 1). Deep lake ATT was well oxygenated over almost the entire water column, whereas all other lakes showed distinct declines in DO concentrations in the hypolimnion. Samples taken close to the bottom of the lakes FAA, HAL, IRR, MIL and WEI were characterized by DO concentration <1 mg L$^{-1}$ (Fig. 1). Accordingly, most lakes revealed an increase in ammonium concentrations at the deepest sampling depths. Nitrate concentrations showed the highest values in the metalimnion, nitrate depletion caused by photoautotrophs occurred in the epilimnion of the lakes. In the hypolimnion, nitrate generally decreased with depth. A sharp decline in nitrate concentration was observed close to the lake bottom. Among the lakes, significant differences in hydrogen-carbonate concentrations occur as this parameter is strongly influenced by the geology of the catchment (data not shown). However, hydrogencarbonate values range between 89 and 220 mg L$^{-1}$, suggesting sufficient inorganic carbon concentrations for chemoautotrophs in all lakes. TRA exhibited high chloride concentrations in the hypolimnion (11.4 mg L$^{-1}$ at 30 m increasing to and 49.3 mg L$^{-1}$ at 191 m), caused by waste disposal of soda and salt industries.

Thaumarcheota are the dominant nitrifiers in deep oligotrophic lakes
The quantification of hcd genes coding for the 4-hydroxybutyryl-CoA dehydratase in the HP/HB cycle indicates that Thaumarchaeota were the dominant nitrifiers in deep oligotrophic lakes (ATT, HAL, MIL and TRA). With exception of lakes HAL and TRA, the vertical distribution of hcd abundances was characterized by very low numbers in the upper water layers (Fig. 2). Thaumarchael hcd gene numbers in the hypolimnion were at least one magnitude higher than functional genes of bacterial nitrifiers (Figs 2 and 3). Thaumarcheal hcd gene numbers in the hypolimnion were at least one magnitude higher than functional genes of bacterial nitrifiers (Figs 2 and 3). The maximum hcd abundance was observed in ATT at 170 m water depth (6.91 × 10$^3$ genes mL$^{-1}$). Smaller lakes (FAA, IRR, MON and WEI) were generally characterized by very low archaeal hcd gene numbers (<10 mL$^{-1}$). The only exception was the hypolimnion of lake WEI, where hcd abundances reached values up to 0.21 × 10$^3$ gene copies mL$^{-1}$ (Fig. 2). Distribution patterns of archaeal amoA gene numbers were tightly correlated with hcd genes ($r = 0.98, P < 0.001$, Table 2) and on average almost equally abundant as hcd genes (Fig. 3). In
accordance with the autotrophic gene marker, genes coding for archaeal AMO also showed the highest values in 170 m depth of ATT ($5.93 \times 10^4$ genes mL$^{-1}$). Relative thaumarchaeal CARD-FISH counts (% of DAPI stained cells) were also significantly correlated with hcd ($r = 0.67, P < 0.001$) and amoA gene numbers ($r = 0.66, P < 0.001$, Table 2). With exception of the hypolimnion of lake MON, CARD-FISH numbers of Thaumarchaeota were close to the detection limit or absent in all smaller lakes (Fig. 3; Fig. S1, Supporting Information).

In freshwater systems, most research focusing on AOA are based on amoA gene surveys, suggesting that Thaumarchaeota are major players in the nitrogen cycle in deep lakes (e.g. August et al. 2012; Vissers et al. 2013a,b; Hugoni et al. 2013). These results are also supported by studies based on RNA analysis (including CARD-FISH) targeting Thaumarchaeota (Callieri et al. 2016). Callieri et al. (2014) amongst others already pointed out that the hypolimnion of deep oligotrophic lakes is also a place of important microbial metabolisms in regard to the carbon cycle and that the magnitude of dark CO2 fixation rates are comparable with photosynthetic fixation of inorganic carbon in the photic zone. Nevertheless, caution is required when linking thaumarchaeal abundances with potential archaeal chemosynthetic activity, because several studies suggest that not all thaumarchaeal representatives are oxidizing ammonium and exhibiting hetero trophic or mixotrophic lifestyles (Herndl et al. 2005; Ingalls et al. 2006; Agogue et al. 2008; Alonso-Sáez et al. 2012). However, the highly similar distribution patterns of amoA and hcd gene numbers in this investigation provide strong evidence for a chemosynthetic lifestyle of the thaumarchaeal communities at our study sites.

**Nitrospira observed in the lakes were mostly NOB, correlated with the distribution of AOA**

Phylogenetic analysis of aclA genes, coding for the ATP-citrate lyase in the rTCA cycle, showed that aclA is not a suitable phylogenetic marker to differentiate comammox-Nitrospira from nitrite oxidizing Nitrospira (Fig. 4, Alfreider et al. 2017). The same is true for RNA gene sequences, because comammox-Nitrospira do not form a monophyletic clade within Nitrospira (Pjevac et al. 2017). Consequently, our analysis based on CARD-FISH and aclA targeting information revealed that all form IA Nitrospira, whereas amoA-targeted primers recently developed by Pjevac et al. (2017) allowed the specific quantification of comammox clades A and B. However, as both clades of comammox-Nitrospira were rare in all lakes (average amoA gene abundances $< 10$ mL$^{-1}$, Fig. 3), the majority of Nitrospira detected at our study sites are most probably related to NOB.

In accordance with thaumarchaeal hcd gene numbers, the highest abundances of aclA genes were also detected in the hypolimnion of the deep lakes, with the maximum number registered in MIL at 80 m depth ($1.5 \times 10^4$ genes mL$^{-1}$). In low AOA lakes, however, Nitrospira were also almost one magnitude less abundant (Fig. 3). Correlation analysis indicate that the distribution pattern of nitrite oxidizing Nitrospira based on aclA gene counts closely followed AOA ($r = 0.74, P < 0.001$; Table 2). This significant correlation in the distribution of AOA and NOB suggests biological interactions between both groups of nitrifiers. The co-occurrence of AOA and different guilds of NOB was also shown in other studies including freshwater ecosystems (Mukherjee et al. 2016) and a broad range of terrestrial habitats (Ke et al. 2013; Daebeler et al. 2014; Stempfhuber et al. 2015; Stempfhuber et al. 2017). However, the exact nature of this potential relationship is not completely elucidated, as microbe-microbe interactions were so far mostly studied between AOB and NOB (Daims, Lücke and Wagner 2016). The hypolimnion of deep lakes was also the environment where Nitrospira-like bacteria detected with CARD-FISH (probe Ntspa 662) were most abundant (Fig. 3), though the distribution patterns did not concur with the vertical gradients observed for aclA gene abundances (Table 2, Fig. 1 and Fig. S1, Supporting Information). With exception of the hypolimnion of WEI, aclA gene numbers were rare in smaller lakes ($< 100$ aclA genes mL$^{-1}$; Fig. 3).

**Nitrospomas-related taxa are the major nitrifying group in smaller lakes**

Primers specifically designed to target the CBB cycle for CO2 fixation in the Nitrospomas lineage revealed that these microbes were the most abundant nitrifiers in smaller lakes (Figs 2 and 3), with highest abundances of $1.1 \times 10^3$ genes mL$^{-1}$ observed in MON in 20 m depth (Fig. 2). In large and deep lakes, numbers of cbbL-form IA genes were about two magnitudes lower than thaumarchaeal hcd abundances, however higher values were usually encountered in the deepest water layers (Figs 2 and 3). The maximal abundance was observed in ATT ($6.9 \times 10^3$ genes mL$^{-1}$ in 170 m depth, Fig. 2). Abundances of AOB, quantified with a bacterial amoA primer set, were of similar magnitude with cbbL IA in both lake types (Fig. 3). The highest numbers were measured in MON in 20 m depth ($2.05 \times 10^3$ genes mL$^{-1}$, Fig. 2). In contrast to archaeal marker genes for autotrophy and ammonia oxidation, cbbL IA and bacterial amoA genes showed a less pronounced correlation between and within the lakes ($r = 0.54; P < 0.001$, Table 2). Highest cell counts of AO-Betaproteobacteria (probe Nso1225) were usually observed in the deep water layers (Fig. S1, Supporting Information). CARD-FISH counts based on probe Nitro878, which was specifically designed to detect the N. oligotropha group, were mostly below the detection limit (data not shown). It is important to note, however, that other AOB might be present in the lakes that are not covered by both markers.

One explanation for the discrepancy between different marker genes targeting AOB might be the taxonomic coverage and specificity of the primer sets used (Table S1, Supporting Information). Sequence analysis of PCR amplicons with the primer set q.cbbL.IA.Nit.J/q.cbbL.IA.Nit.r, which was specifically designed for the present study, showed that all form IA Rubisco sequences were affiliated with representatives of the targeted N. oligotropha lineage (Nitrosomomas cluster 6A) and closely related sister clades (Fig. 5). This group of AOB was found to be the dominating group of nitrifiers based on sequences derived by broad range primers for Form IA Rubisco genes in a variety of lakes (Alfreider et al. 2017). On the other hand, primers used for the quantification of amoA genes cover a wide range of proteobacterial AOB (Meinhardt et al. 2015), but primers do not perfectly match with the cbbL genes of several representatives of the N. oligotropha lineage (data not shown). Another reason might be the incongruent cbbL phylogeny of ammonia oxidizing Proteobacteria compared with tRNA based taxonomy, the latter gene was targeted by a CARD-FISH probe also designed to detect the N. oligotropha lineage (Table S2, Supporting Information). Although different forms of Rubisco are conserved proteins with distinct sequence differences, both horizontal gene
transfer and gene duplication in proteobacterial lineages complicate the interpretation of systematic and physiological relationships based on RuBisCO phylogeny (Delwiche and Palmer 1996; Tabita et al. 2008).

Influence of environmental factors on distribution patterns

There is a fast growing number of studies investigating the environmental factors determining the distribution of different nitrifying guilds in nature. Particularly niche preferences of AOB and AOA have been extensively surveyed in the last years and several biotic and abiotic factors have been identified that determine their distribution in nature. So far, most investigations have been performed in soil and marine ecosystems, although patterns of niche differentiation of AOA and AOB were also studied in freshwater environments (Jiang et al. 2009; French et al. 2012; Small et al. 2013; Vissers et al. 2013a; Hayden and Beman 2014; Mukherjee et al. 2016; Pajares et al. 2017). In general, AOB dominantly contribute to nitrification under high substrate concentrations while AOA are the most abundant group in oligotrophic systems.

Potential niche preferences of AOA and AOB are usually not discussed from the perspective of different CO2 fixation strategies used by AOs, although different environmental conditions control their distribution in nature. The availability of ammonium and DO concentration were the determining factors for the occurrence of AOB at the study sites (Fig. 6). In deep lakes, at sampling depths where ammonium levels were very low, AOB were outnumbered by AOA by two or even three orders of magnitude (Fig. 2). The distribution of AOB was also positively correlated with the concentration of total phosphorus, suggesting that AOB and the high energy demand of the CBB cycle are better adapted to water depths characterized by an elevated nutrient status. Several studies have already reported that environments favor AOB development with higher substrate availability, including cultivation based investigations specifically targeting the N. oligotropha group (French et al. 2012). On the other hand, AOA using the most energy efficient aerobic carbon cycle (Könneke et al. 2014) were the dominant group in the hypolimnion of deep oligotrophic lakes and their distribution was positively correlated with depth (Fig. 6). Although not directly measured, depth dependent parameters such as competition for substrates with phototrophs, heavy grazing pressure on slow-growing nitrifiers and inhibition by light might be responsible for the low numbers of AOA in the surface waters of lakes (Hugoni et al. 2013; Small et al. 2013; Vissers et al. 2013a; Alfreider et al. 2017).

Several studies have revealed that AOA are better adapted to low oxygen concentrations than AOB (Lam et al. 2007; Martens-Habbena et al. 2009; French et al. 2012; Hugoni et al. 2013). These findings are not in accordance to our results, where DO was negatively correlated with cbbL gene numbers in AOB (Fig. 6). A low Km for O2 in AOA supports the hypothesis that AOA are well adapted to low O2 concentrations. On the other hand, the magnitude of DO is also of fundamental importance for the efficiency of the CBB cycle in organisms using this pathway (Berg 2011). Increased O2 levels in the catalytic environment of RuBisCO enzymes in AOB have a negative effect on CO2 fixation abilities. In this context, one of the most important biochemical characteristics between different forms of Rubisco is the ability to discriminate between CO2 and O2 at a given CO2:O2 concentration ratio (Tabita 1999).

Although sequence analysis of different forms of Rubisco did not reveal a major cluster affiliated with NOB using the CBB cycle for CO2 fixation in lakes (Alfreider et al. 2017), we can for the moment only speculate that the genus Nitrospira is the most dominant NOB group at our study sites. In contrast to AOA and AOB, very little is known about the environmental factors determining the niche partitioning of NOB including Nitrospira in natural freshwater systems (Pester et al. 2014). Temperature, DO and nitrite levels have been shown to be key factors in niche differentiation of different NOB lineages (Daims, Lücke and Wagner 2016). At our study sites, multivariate statistical analysis showed that temperature is negatively correlated with the distribution of Nitrospira aclA gene abundances in the lakes, although different Nitrospira strains are known to grow in a broad temperature range (Alawi et al. 2009). It has been proposed that Nitrospira strains preferentially thrive under hypoxic conditions (Park and Noguer 2008) due to the lack of reactive oxygen species as a
classic defense mechanism against oxidative stress (Lücke et al. 2010). Our results demonstrated the presence of Nitrospira-like bacteria (which also includes potential comammox-Nitrospira) in both high and low DO habitats and statistical analysis suggests that oxygen concentrations had no crucial effect on the distribution of Nitrospira-like bacteria in the lakes (Fig. 6).

Redundancy analysis of the relationship between environmental parameters and CARD-FISH counts showed a similar trend observed with the ddPCR results of AOA and NOB based on functional genes (Fig. S2, Supporting Information). However, in contrast to the RDA analysis based on amoA and cbbL abundances, the availability of ammonium and DO concentration were not the controlling variables for the distribution of AOB in lakes based on CARD-FISH numbers. One explanation for this disagreement might be taxonomic coverage and specificity of the different markers used, which was already discussed above. Consequently, it can also not be ruled out that some representatives detected by CARD-FISH have a heterotrophic lifestyle. Another reason for discrepancy are the different detection limits of the two approaches (Alfreider et al. 2017). Gene copy numbers analyzed by ddPCR were measured as low as one target gene mL$^{-1}$ in the lake water samples, while microscopic-based techniques do not allow accurate counts at this low magnitude. As AOB mostly occur in very low abundances at the study sites, the efficiency of both methods might cause corresponding differences in the results.

One major question, however, remains unclear: Why is the third group of AOs, comammox-Nitrospira, so rare in the studied lakes? This stands in contrast to a recent investigation suggesting that the comammox bacterium N. inopinata is highly adapted to oligotrophic habitats, at least based on results derived from substrate competition kinetics (Kits et al. 2017). Studies that investigate the influence of environmental variables on the structure or distribution of comammox are rare. Beside the proposed adaptation of comammox-Nitrospira to microaerophilic and low substrate fluxes (Lawson and Lücke 2018), Fowler et al. (2018) identified temperature to have a positive impact on Nitrospira in rapid sand filters. However, the authors could not distinguish if comammox and nitrite-oxidizing taxa were affected. In principle, the high adaptation of comammox-Nitrospira to oligotrophic conditions is also reflected by the rTCA cycle, a pathway that is considered to be far more energy efficient than other CO$2$ fixation cycles employing non-reducing carboxylases (Berg 2011; Mangiapia and Scott 2016). However, some taxa including the thermophilic Hydrogenobacter thermophilus and the genus Nitrospira, have developed enzymatic adaptations for oxygen tolerance, in contrast to most other bacterial phyla operating this cycle in anaerobic or microaerobic environments (Yamamoto et al. 2006; Lücke et al. 2010; Berg 2011; Daims et al. 2015). The specific biochemical adaptations are poorly understood and some still unknown mechanisms may strengthen the O$_2$ robustness of the rTCA pathway in Nitrospira (Berg 2011). However, if these protection mechanisms are accompanied by a lower specific activity, it may significantly increase the energy requirements of the pathway when used in an oxic environment (Berg 2011). Certainly, more research is necessary to determine the niche preferences of comammox-Nitrospira, focusing on habitats characterized by different trophic states and oxygen levels.

**SUPPLEMENTARY DATA**

Supplementary data are available at FEMSEC online.

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**REFERENCES**

Agogue H, Brink M., Dinasquet J et al. Major gradients in putatively nitrifying and non-nitrifying Archaea in the deep North Atlantic. Nature 2008;456:788–91.

Alawi M, Off S, Kaya M et al. Temperature influences the population structure of nitrite-oxidizing bacteria in activated sludge. Environ Microbiol Rep 2009;1:184–90.

Alfreider A, Baumer A, Bogensperger T et al. CO₂ assimilation patterns in stratified lakes: diversity and distribution patterns of chemolithoautotrophs. Environ Microbiol 2017;19:2754–68.

Alonso-Sáez L, Waller AS, Mende DR et al. Role for urea in nitrification by polar marine Archaea. Proc Natl Acad Sci USA 2012;109:17989–94.

Auguet JC, Triado-Margarit X, Nomokonova N et al. Vertical segregation and phylogenetic characterization of ammonia-oxidizing archaea in a deep oligotrophic lake. ISME J 2012;6:1786–97.

Badger MR, Bek EJ. Multiple Rubisco forms in proteobacteria: their functional significance in relation to CO₂ acquisition by the CBB cycle. J Exp Bot 2008;59:1525–41.

Berg IA. Ecological aspects of the distribution of different autotrophic CO₂ fixation pathways. Appl Environ Microbiol 2011;77:1925–36

Bergauer K, Sintes E, Bleijswijk J et al. Abundance and distribution of archaeal acetyl-CoA/propyionyl-CoA carboxylase genes indicative for putatively chemolithoautotrophic Archaea in the tropical Atlantic’s interior. FEMS Microb Ecol 2013;84:461–73.

Bollmann A, Sedlacek CJ, Norton J et al. Complete genome sequence of Nitrosomonas sp. Is79, an ammonia oxidizing bacterium adapted to low ammonium concentrations. Stand Genomic Sci 2013;7:469–82.

Bouskill NJ, Eveillard D, Chien D et al. Environmental factors determining ammonia-oxidizing organism distribution and diversity in marine environments. Environ Microbiol 2012;14:714–29.

Callieri C, Coci M, Eckert EM et al. Archaea and bacteria in deep lake hypolimnion: in situ dark inorganic carbon uptake. J Limnol 2014;73:31–8.

Callieri C, Hernández-Avilés S, Salcher MM et al. Distribution patterns and environmental correlates of Thaumarchaeota abundance in six deep subalpine lakes. Aquat Sci 2016;78:215–25.

Daebeler A, Bodelier PLE, Yan Z et al. Interactions between Thaumarchaeota, Nitrospira and methanotrophs modulate autotrophic nitrification in volcanic grassland soil. ISME J 2014;8:2397–410.

Daims H, Lücker S, Wagner M. A new perspective on microbes formerly known as nitrite-oxidizing bacteria. Trends Microbiol 2016;24:699–712.

Daims H, Lebedeva E, Pfevac P et al. Complete nitrification by Nitrospira bacteria. Nature 2015;528:504–9.

Delwiche CF, Palmer JD. Rampant horizontal transfer and duplication of rubisco genes in eubacteria and plastids. Mol Biol Evol 1996;13:873–82.

Fowler SJ, Palomo A, Dechesne A et al. Comammox Nitrospira are abundant ammonia oxidizers in diverse groundwater-fed rapid sand filter communities. Environ Microbiol 2018;20:1002–15.

French E, Kozlowski JA, Mukherjee M et al. Ecophysiological characterization of ammonia-oxidizing archaea and bacteria from freshwater. Appl Environ Microbiol 2012;78:5773–80.

Hatzenpichler R. Diversity, physiology, and niche differentiation of ammonia-oxidizing archaea. Appl Environ Microbiol 2012;78:7501–10.

Hayden CJ, Beman JM. High abundances of potentially active ammonia-oxidizing bacteria and archaea in oligotrophic, highaltitude lakes of the Sierra Nevada, California, USA. PLoS ONE 2014;9:1–9.

Hernndi GJ, Reinthaler T, Teira E et al. Contribution of Archaea to total prokaryotic production in the deep Atlantic Ocean, Appl Environ Microbiol 2005;71:2303–9

Hu A, Yang Z, Yu CP et al. Dynamics of autotrophic marine planktonic Thaumarchaeota in the East China Sea. PLoS ONE 2013;8:e61087.

Hügler M, Sievert SM. Beyond the Calvin Cycle: autotrophic carbon fixation in the ocean. Ann Rev Mar Sci 2011;3:261–89.

Hugoni M, Etien S, Bourges A et al. Dynamics of ammonia-oxidizing archaea and bacteria in contrasted freshwater ecosystems. Res Microbiol 2013;164:360–70.

Ingalls AE, Shah SR, Hansman RL et al. Quantifying archaeal community autotrophy in the mesopelagic ocean using natural radiocarbon. Proc Natl Acad Sci USA 2006;103:6442–7.

Jiang HC, Dong H, Yu BS et al. Diversity and abundance of ammonia-oxidizing archaea and bacteria in Qinghai Lake, northwestern China. Geomicrobiol J 2009;26:199–211.

Ke X, Angel R, Lu Y et al. Niche differentiation of ammonia oxidizers and nitrite oxidizers in rice paddy soil. Environ Microbiol 2013;15:2275–92.

Kits KD, Sedlacek CJ, Lebedeva EV et al. Kinetic analysis of a complete nitrifier reveals an oligotrophic lifestyle. Nature 2017;549:269–72.

Könneke M, Bernhard AE, de la Torre JR et al. Isolation of an autotrophic ammonia-oxidizing marine archaeon. Nature 2005;437:543–6.

Könneke M, Schubert DM, Brown PC et al. Ammonia-oxidizing archaea use the most energy efficient aerobic pathway for CO₂ fixation. Proc Natl Acad Sci USA 2014;111:8239–44.

Kovalchuk GA, Stephen JR. Ammonia-oxidizing bacteria: a model for molecular microbial ecology. Annu Rev Microbiol 2001;55:485–529.

Kovalova OL, Tourova TP, Muyzer G et al. Diversity of RuBiSCo and ATP citrate lyase genes in soda lake sediments. FEMS Microbiol Ecol 2011;75:37–47.

La Cono V, La Spada G, Arcadi E et al. Partaking of Archaea to biogeochemical cycling in oxygen-deficient zones of meromictic saline Lake Faro (Messina, Italy). Environ Microbiol 2013;15:1717–33.
Lawson CE, Lücke S. Complete ammonia oxidation: an important control on nitrification in engineered ecosystems. Curr. Opin. Biotechnol. 2018;50:158–65 S0958-1669(17)30161-1 29414055

Pruesse E, Quast C, Knittel K et al. SILVA: a comprehensive online resource for quality checked and aligned ribosomal RNA sequence data compatible with ARB. Nucleic Acids Res 2007;35:7188–96.

Small GE, Bullerjahn GS, Sterner RW et al. Rates and controls of nitrification in a large oligotrophic lake. Limnol Oceanogr 2013;58:276–86.

Stein LY, Arp DJ, Berube PM et al. Whole-genome analysis of the ammonia-oxidizing bacterium, Nitrosonomas eutrophica C91: implications for niche adaptation. Environ Microbiol 2007;9:2993–3007.

Stempfhuber B, Richter-Heitmann T, Regan KM et al. Spatial interaction of archaeal ammonia-oxidizers and nitrite-oxidizing bacteria in an unfertilized grassland soil. Front Microbiol 2015;6:1567.

Stempfhuber B, Richter-Heitmann T, Bieneck L et al. Soil pH and plant diversity drive co-occurrence patterns of ammonia and nitrite oxidizer in soils from forest ecosystems. Biol Fertil Soils 2017;53:691–700.

Tabita FR. Microbial ribulose 1,5-bisphosphate carboxylase/oxygenase: a different perspective. Photosynth Res 1999;60:1–28.

Tabita FR, Satogapan S, Hansson TE et al. Distinct form I, II, III, and IV Rubisco proteins from the three kingdoms of life provide clues about Rubisco evolution and structure/function relationships. J Exp Bot 2008;59:1515–24.

Tamura K, Stecher G, Peterson D et al. MEGA6: molecular evolutionary genetics analysis version 6.0. Mol Biol Evol 2013;30:2725–9.

Tolar B, King GM, Hollibaugh JT. An analysis of Thaumarchaeota populations from the Northern Gulf of Mexico. Front Microbiol 2013;4:72.

van Kessel M, Speth D, Albertsen M et al. Complete nitrification by a single microorganism. Nature 2015;528:555–9.

Vissers EW, Anselmetti FS, Bodelier PLE et al. Temporal and spatial coexistence of archaeal and bacterial amoA genes and gene transcripts in Lake Lucerne. Archaea 2013a;2013:289478.

Vissers EW, Blaga CI, Bodelier PLE et al. Seasonal and vertical distribution of putative ammonia-oxidizing thaumarcheotal communities in an oligotrophic lake. FEMS Microbiol Ecol 2013b;83:515–26.

Wendeborg A. Fluorescence in situ hybridization for the identification of environmental microbes. Cold Spring Harb Protoc 2010; DOI:10.1101/pdb.prot5366.

Yakimov MM, La Cono V, Denaro R. A first insight into the occurrence and expression of functional amoA and accA genes of autotrophic and ammonia oxidizing bathypelagic Crenarchaeota of Tyrrhenian Sea. Deep-Sea Res 2009;56:748–54.

Yakimov MM, La Cono V, Smedile F et al. Contribution of crenarchaeal autotrophic ammonia oxidizers to the dark primary production in Tyrrhenian deep waters (Central Mediterranean Sea). ISME J 2011;5:945–51.

Yamamoto M, Ariai H, Ishii M et al. Role of two 2-oxoglutarate: ferredoxin oxidoreductases in Hydrogenobacter thermophilus under aerobic and anaerobic conditions. FEMS Microbiol Lett 2006;263:189–93.

Yilmaz LS, Parmenkar S, Noguera DR. mathFISH, a web tool that uses thermodynamics-based mathematical models for in silico evaluation of oligonucleotide probes for fluorescence in situ hybridization. Appl Environ Microbiol 2011;77:1118–22.