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Ecophysiological Characterization of Ammonia-Oxidizing Archaea and Bacteria from Freshwater

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Aerobic biological ammonia oxidation is carried out by two groups of microorganisms, ammonia-oxidizing bacteria (AOB) and the recently discovered ammonia-oxidizing archaea (AOA). Here we present a study using cultivation-based methods to investigate the differences in growth of three AOA cultures and one AOB culture enriched from freshwater environments. The strain in the enriched AOA culture belong to thaumarchaeal group I.1.a, with the strain in one enrichment culture having the highest identity with “Candidatus Nitrosoarchaeum koreensis” and the strains in the other two representing a new genus of AOA. The AOB strain in the enrichment culture was also obtained from freshwater and had the highest identity to AOB from the Nitrosomonas oligotropha group (Nitrosomonas cluster 6a). We investigated the influence of ammonium, oxygen, pH, and light on the growth of AOA and AOB. The growth rates of the AOB increased with increasing ammonium concentrations, while the growth rates of the AOA decreased slightly. Increasing oxygen concentrations led to an increase in the growth rate of the AOB, while the growth rates of AOA were almost oxygen insensitive. Light exposure (white and blue wavelengths) inhibited the growth of AOA completely, and the AOA did not recover when transferred to the dark. AOB were also inhibited by blue light; however, growth recovered immediately after transfer to the dark. Our results show that the tested AOB have a competitive advantage over the tested AOA under most conditions investigated. Further experiments will elucidate the niches of AOA and AOB in more detail.

Nitration, the microbial oxidation of NH₃ (ammonia) to NO₃⁻ (nitrate), is one of the key processes of the global nitrogen cycle. The first and rate-limiting step of nitrification is the oxidation of NH₃ to NO₂⁻ (nitrite). Until recently, aerobic ammonia oxidation was attributed to only a small subset of the Proteobacteria; most freshwater and terrestrial ammonia-oxidizing bacteria (AOB) belong to a distinct group in the Betaproteobacteria, while a few marine AOB species belong to the Gammaproteobacteria (29, 32, 33). The AOB have a chemolithoautotrophic metabolism, oxidizing NH₃ to NO₂⁻ via the intermediate NH₂OH (hydroxylamine) and fixing carbon from CO₂ (carbon dioxide) via the Calvin cycle (1).

Recently, genes encoding ammonia monooxygenase (amoA), the first enzyme in the process of ammonia oxidation, were discovered together with archaeal 16S rRNA genes in a metagenomic study (60) and a soil fosmid library (57). At the same time, Nitrosopumilus maritimus, the first archaeal ammonia oxidizer, was isolated in pure culture from a saltwater aquarium (30). Ammonia-oxidizing archaea (AOA) in pure and enrichment cultures have essentially the same metabolism as AOB; they oxidize NH₃ stoichiometrically to NO₂⁻ and fix carbon from bicarbonate (HCO₃⁻) (15, 20, 30, 36, 43, 56). However, the genomes of N. maritimus and “Candidatus Nitrosoarchaeum limnia” revealed differences between AOA and AOB, such as the use of the 3-hydroxypropionate/4-hydroxybutyrate pathway for HCO₃⁻ fixation, the absence of hydroxylamine oxidoreductase, and the presence of many copper-containing enzymes (5, 63).

AOA and AOB often co-occur in the same environment, but the contributions of AOA and AOB to the total ammonia oxidation still need to be elucidated. Many previous studies focused on the influence of environmental factors on niche differentiation between AOA and AOB using cultivation-independent molecular methods. From those studies, it can be concluded that AOA are frequently found in environments with lower substrate (NH₄⁺ and O₂) availability and AOB are frequently found in environments with higher substrate availability (see references 4, 13, 17, 24, 42, and 55, among others). However, most of these studies were conducted using methods that target the abundance and/or expression of the archaeal and bacterial amoA genes. Unfortunately, it is not possible to draw direct conclusions about the activity of the AOA and AOB on the basis of the abundance and expression of the amoA gene, because amoA mRNAs has been detected in AOB for weeks and 16S rRNA (ribosomes) has been detected for up to a year after the onset of starvation (8, 25, 26). The response of AOA toward starvation and resuscitation has not yet been investigated. In addition, it has been shown that not all amoA-recording Thaumarchaeota are autotrophic ammonia oxidizers (39, 64). While studies focusing on the analysis of abundance and activity of microbes using molecular methods give very valuable insights, it is also necessary to investigate the response of microbes to environmental factors using cultivation-based approaches, because these experiments will demonstrate changes in physiological activity more conclusively.

Here we present a study that used a cultivation-dependent approach to investigate the responses of AOA and AOB to environmental factors. Cultures of three phylogenetically distinct AOA from freshwater sediments in Ohio were enriched, and the growth
of the AOA was characterized under different conditions and compared with that of AOB from freshwater in an enrichment culture. Factors of interest include the NH$_4^+$ concentration, pH, O$_2$ concentration, and light wavelength and intensity. These factors have strong effects on the physiology and niche differentiation between AOA and AOB.

**MATERIALS AND METHODS**

**Sampling.** Near-shore sediment samples were taken from Lakes Acton (AC; 39°55’N, 84°74’W) and Delaware (DW; 40°59’N, 83°05’W) in fall 2008. Additional sediment core samples were collected from Lake Acton in summer 2009.

**Medium.** The mineral salts (MS) medium used to enrich and cultivate AOA and AOB contained 10 mM NaCl, 1 mM KCl, 1 mM CaCl$_2$, 2 mM H$_2$O$_2$, 0.2 mM MgSO$_4$ · 7H$_2$O, and 1 ml liter$^{-1}$ trace elements solution (9, 61). HEPES buffer was added in a 4-fold molar ratio to the NH$_4^+$ concentration, and the pH was adjusted to 7.5 before autoclaving. After autoclaving, sterile KH$_2$PO$_4$ solution was added to obtain a final concentration of 0.4 mM (9, 61).

**Enrichment of AOA (AOA-AC2, AOA-AC5, and AOA-DW enrichment cultures).** Sediment samples (1 g) were inoculated into 50 ml MS medium with 0.25 mM NH$_4^+$ immediately upon arrival in the laboratory. The enrichments were incubated at 27°C in the dark. NH$_4^+$ levels were monitored weekly using a colorimetric assay (9, 28). When the cultures reached late logarithmic growth phase (depletion of about 80% of the initial NH$_4^+$ concentration), they were transferred to fresh medium using a 10% (vol/vol) inoculum. The cultures were passed through 0.45-μm-pore-size filters for the first five to six transfers to exclude AOB (9; Annika Mosier, personal communication). In addition to filtration, the enrichment cultures from DW were also treated with 100 μg ml$^{-1}$ streptomycin to eliminate AOB. After several transfers, when the cultures depleted NH$_4^+$ at regular intervals, 20 ml was collected on 0.1-μm-pore-size nitrocellulose filters for molecular characterization. The filters were stored at −20°C.

**AOB culture.** We used the previously described AOB strain from a freshwater enrichment culture G5-7 (AOB-G5-7) to compare the growth of AOA to that of AOB (6, 7). The strain belongs to the Nitrosonomas oligotropha cluster and is adapted to low NH$_4^+$ concentrations (6, 7). Members of this AOB cluster have been found in many freshwater environments around the world (11, 12, 14, 22, 53; E. French and A. Bollmann, unpublished data).

**Growth experiments.** All growth experiments were conducted in MS medium with 0.5 mM NH$_4^+$ at pH 7.5 in 125-ml Erlenmeyer flasks with cotton stoppers unless otherwise noted. We tested the influence of different factors (NH$_4^+$ concentration, O$_2$ concentration, pH, and light) on the rate of NO$_3^-$ (NO$_2^-$) production of the three AOA enrichment cultures (AOA-AC2, AOA-AC5, and AOA-DW) and the AOB enrichment culture (AOB-G5-7). All cultures were inoculated with 10% (vol/vol) conditioned late-log-phase cells and incubated in the dark at 27°C. Samples (1 ml) were taken at regular intervals and centrifuged at 16,000 rpm for 20 min. The supernatant was stored at −20°C for further chemical analysis. To investigate the influence of different NH$_4^+$ concentrations, medium with 15 μM to 5 mM NH$_4^+$ was prepared with the corresponding HEPES concentrations. The influence of pH was investigated by adjusting the initial pH in the medium to values between 6 and 9. The influence of the O$_2$ concentration was investigated by equilibrating the medium in serum bottles under anaerobic conditions overnight. After equilibration, the bottles were sealed with rubber stoppers. Different calculated O$_2$ concentrations in the headspace were achieved by exchanging the corresponding volume of the headspace with sterile filtered air. The influence of light was investigated by incubating the cultures 18 cm above light-emitting diode panels emitting 30 μmol photons m$^{-2}$ s$^{-1}$ at the wavelengths 5,000 to 7,000K (white light), 623 ± 3 nm (red light), and 470 ± 5 nm (blue light) and 3 μmol photons m$^{-2}$ s$^{-1}$ at the wavelength 470 ± 5 nm (blue light). The light intensity inside the glass bottles was 25 μmol photons m$^{-2}$ s$^{-1}$ (high light conditions) and 2.5 μmol photons m$^{-2}$ s$^{-1}$ (low light conditions), as measured with a LI-250A light meter (LI-COR Biosciences, Lincoln, NE), indicating that the glass filtered approximately 15% of the light. To investigate the influence of light-to-dark and dark-to-light transitions on the growth of AOA and AOB, cultures were incubated in the dark until 50% of the NH$_4^+$ was consumed and then transferred to the light. At the same time, cultures that were incubated in the light were transferred from the light to the dark. Controls were incubated for the complete cycle in the dark.

**Evaluation of growth experiments.** NO$_3^-$ and NO$_2^-$ concentrations were determined in the supernatants using colorimetric assays (9, 49). NO$_3^-$ /NO$_2^-$ concentrations were log transformed and plotted against time (see Fig. S1 in the supplemental material). Growth rates were calculated from the linear increase (slope) of the log-transformed NO$_3^-$ /NO$_2^-$ concentrations over time, assuming that NO$_2^-$ /NO$_3^-$ production in the cultures is correlated with the growth of AOA and AOB (9, 3, 30). The increase in NO$_3^-$ /NO$_2^-$ production was linear for several days to 1 week, and the correlation coefficients were always ≥0.97 but in most cases were even ≥0.99.

**Molecular analysis.** (i) DNA isolation from AOA enrichment cultures. DNA was isolated from the nitrocellulose filters using a Qiagen DNeasy blood and tissue kit (Valencia, CA) with the following modifications. Acid-washed zirconium beads (1 g) and 500 μl high-salt buffer (1 M NaCl, 5 mM MgCl$_2$ · 2H$_2$O, 10 mM Tris, pH 8) (2) were added to the nitrocellulose filters. The filters were homogenized using a bead beater (Biospec Products, Bartlesville, OK) at 4,800 rpm for 30 s. This was repeated three times, and the samples were stored in between cycles on ice for 10 min. After bead beating, 500 μl Qiagen buffer AL and 50 μl proteinase K were added and the mixture was incubated at 56°C for 30 min. The reaction mixture was spun down at 8,000 rpm for 1 min and transferred to spin columns supplied by the manufacturer. The spin columns were treated according to the manufacturer’s recommendations, and the DNA was eluted with 100 μl elution buffer AE (Qiagen).

(ii) PCR. GoTaq green master mix (Promega, Madison WI) was used for all standard PCRs, according to the manufacturer’s recommendations, using the primers and protocols summarized in Table S1 in the supplemental material.

(iii) Cloning and sequencing. PCR products were cleaned using Wizard SV gel and PCR product cleanup system (Promega, Madison, WI) and cloned into the pGEM-T Easy vector system (Promega, Madison, WI) with the following modifications: the hybridization temperature was 46°C, and the first wash conditions were 0.2% (v/v) SDS and 0.9 M NaCl, 7 mM MgCl$_2$ · 2H$_2$O, 10 mM Tris, pH 8. (2) were added to the nitrocellulose filters. The filters were homogenized using a bead beater (Biospec Products, Bartlesville, OK) at 4,800 rpm for 30 s. This was repeated three times, and the samples were stored in between cycles on ice for 10 min. After bead beating, 500 μl Qiagen buffer AL and 50 μl proteinase K were added and the mixture was incubated at 56°C for 30 min. The mixture was spun down at 8,000 rpm for 1 min and transferred to spin columns supplied by the manufacturer. The spin columns were treated according to the manufacturer’s recommendations, and the DNA was eluted with 100 μl elution buffer AE (Qiagen).

(iv) DNA sequence analysis. All sequences were edited with the 4Peaks program (A. Griekspoor and T. Groothuis, The Netherlands Cancer Institute). The sequences were aligned using ARB software (35). Phylogenetic trees were constructed using the neighbor-joining algorithm in ARB, and parsimony and maximum likelihood methods were performed using the PHYLIP program (16). Trees constructed with all three methods showed the same overall grouping; therefore, only the tree constructed with the neighbor-joining method has been presented.

(v) FISH. The catalyzed reporter deposition (CARD)–fluorescence in situ hybridization (FISH) protocol (45, 48) was used with the following modifications: the hybridization temperature was 46°C, and the first wash was performed at 48°C, followed by an amplification step at 46°C. All probes (see Table S2 in the supplemental material) were labeled at their 3' ends with horseradish peroxidase and used at a final concentration of 50 ng μl$^{-1}$. All filters were counterstained with DAPI (4',6-diamidino-2-phenylindole) for total cell counts. Direct microscopic counts were performed by fluorescence microscopy (Zeiss Axiohot HBO100; Carl Zeiss Inc., North America) at ×1,000 magnification.
Nucleotide sequence accession numbers. All sequences were deposited in GenBank under the numbers JQ669389 to JQ669394.

RESULTS

Enrichment of AOA. AOA were enriched from the sediment of Lakes Acton (AOA-AC2 and AOA-AC5 cultures) and Delaware (AOA-DW culture) under autotrophic conditions with NH4⁺ as the sole electron donor in the medium. On the basis of the AOA amoA sequences, all enrichment cultures belong to water column/sediment group I.1a of the Thaumarchaeota (Fig. 1). The strain in the AOA-AC2 culture was 81 to 81.7% (amoA) and 92.8 to 93.1% (16S rRNA gene) identical to the strains in the other two enrichment cultures, while the strains in AOA-AC5 and AOA-DW were 87.1% (amoA) and 97.9% (16S rRNA gene) identical to each other. The amoA sequences of the strain in AOA-DW were 98.2 to 98.5% identical to those of clones from the sediment of Lakes Acton, Delaware, and Pleasant Hill (C. Li and A. Bollmann, unpublished), 98.5% identical to those of clones from the freshwater sediment in the San Francisco Bay (38), and 98.1% identical to those of clones from a drinking water distribution system in the Netherlands (59). The amoA sequences of the strain in AOA-AC5 were 99.9% identical to the amoA sequence of a clone from a paddy soil in Japan (18). The strain in the third enrichment culture, AOA-AC2, is closely related to "Ca. Nitrosoarchaeum koreensis" (99.4% identity for amoA and 98.3% identity for the 16S rRNA gene) and "Ca. Nitrosoarchaeum limnia" (94.3% identity for amoA and 98.5% identity for the 16S rRNA gene) (5, 27). In contrast to the strain in AOA-AC2, the strains in AOA-AC5 and AOA-DW were not closely related to described AOA isolates or strains in enrichment cultures, such as N. maritimus and Nitrososphaera viennensis, among others (70 to 82% identity for amoA and 81 to 93% identity for the 16S rRNA gene) (Table 1).

CARD-FISH was used to determine the proportion of AOA in the enrichment cultures at the end of the logarithmic growth phase. AOA-DW contained 85% AOA, AOA-AC2 contained 91% AOA, and AOA-AC5 contained 81% AOA (Table 2). AOB and nitrite-oxidizing bacteria (NOB) were not detected, as tested by PCR amplification with AOB-specific 16S rRNA and amoA primers (see Table S1 in the supplemental material) (results not shown) and FISH with AOB- and NOB-specific 16S rRNA probes (Table 2; see Table S2 in the supplemental material).

Influence of NH4⁺ concentration on growth rates of AOA and AOB. During stratification in the summer, the NH4⁺ concentration in Lake Acton increases to up to 400 μM (41), which falls within the tested range of NH4⁺ concentrations of 15 μM and 5 mM NH4⁺. Increasing NH4⁺ concentrations up to 1 mM NH4⁺ doubled the growth rate of the AOB in AOB-G5-7, while the growth rates of the AOA in the enrichment cultures decreased or remained constant (Fig. 2). The growth rate of the AOA in
AOA-DW at the lowest NH₄⁺ concentration (15 μM) was significantly higher than the growth rate at higher NH₄⁺ concentrations (see Table S3 in the supplemental material). The same tendency was observed for the other two cultures, although the statistical support was less strong (Fig. 2; see Table S3 in the supplemental material). The AOA strains in the enrichment cultures exhibited different tolerances to high NH₄⁺ concentrations; the strain in AOA-DW grew at NH₄⁺ concentrations up to 1 mM, the strain in AOA-AC2 grew at NH₄⁺ concentrations up to 2 mM, and the strain in AOA-AC5 grew at NH₄⁺ concentrations up to 5 mM (Fig. 2). The lag phase of AOA and AOB differed; the strain in AOA-DW culture was significantly lower in the red light and after transfer from the light to the dark (Fig. 5; see Tables S7 and S8 in the supplemental material). The same tendency was observed for the other two cultures, although the statistical support was less strong (Fig. 2; see Table S3 in the supplemental material). The strains in all cultures showed bell-shaped curves in relation to the pH, with maximum growth rates at pH 7 to 7.5 (Fig. 4). The strain in the AOA-AC2 culture did not grow at pH 6, while the other AOA strains and the AOB strain did. The growth rates of the strains in AOA-AC5 and AOA-DW cultures at pH 9 were similar to the growth rates at pH 7.5, while the growth rates of the strains in AOA-AC2 and AOB-G5-7 cultures differed significantly from their respective rates at pH 7.5 (Fig. 4; see Table S6 in the supplemental material).

### Influence of pH on growth of AOA and AOB.

Lake Acton stratifies during the summer and has an anaerobic zone as well as a zone with low oxygen availability (1 mg liter⁻¹ O₂) (41). We therefore investigated the response of the strains in our enrichment cultures to 0.5 to 2% O₂ (calculated) in the headspace, which corresponded to 0.2 to 0.8 mg liter⁻¹ O₂ in the medium. The growth rate of the AOB in AOB-G5-7 decreased with decreasing O₂ concentration, and the growth rates at all different O₂ concentrations were significantly different from each other (Fig. 3; see Table S5 in the supplemental material). The strains in the AOA enrichment cultures grew at all O₂ concentrations in the headspace, with the exception of that in the AOA-AC2 enrichment culture at 0.5% O₂. The decrease of the growth rates with decreasing O₂ concentration in the AOA cultures was less steep than the decrease of the growth rates in the AOB enrichment culture. However, at low O₂ concentrations the growth rates in the AOA-AC2 and AOA-AC5 enrichment cultures were significantly lower than the growth rates at 21% O₂ (Fig. 3; see Table S5 in the supplemental material).

### Influence of O₃ concentration on growth of AOA and AOB.

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### Influence of light on growth of AOA and AOB.

The investigated intensities represent a range of light, but below light saturation, at which phytoplankton in freshwater systems are able to grow (51). White light (30 μmol photons m⁻² s⁻¹) strongly inhibited the growth of the AOA in the AOA-DW culture but had no effect on the AOB in the AOB-G5-7 culture (Fig. 5). The AOA did not grow in white light and did not begin to grow after being transferred from the light to the dark. However, growth continued when the AOA cultures were transferred from the dark to the light. To get a better insight into which wavelength of light had the strongest influence on the growth of AOA and AOB, we conducted similar experiments with red (623 ± 3 nm) and blue (470 ± 5 nm) light. Strains in both cultures grew in the red light, but while the growth of the AOB in the AOB-G5-7 culture was not influenced by the red light, the growth rate of the AOA in the AOA-DW culture was significantly lower in the red light and after transfer from the light to the dark (Fig. 5; see Tables S7 and S8 in the supplemental material). Blue light at 30 μmol photons m⁻² s⁻¹ had the strongest effect on the growth of strains in both cultures (Fig. 5). In the blue light, strains did not grow in any of the cultures, and growth of the AOA in the AOA-DW culture did not recover after transfer from the light to the dark. In contrast, the AOB in the AOB-G5-7 culture recovered immediately after transfer from the light to the dark, but the growth rate was significantly lower than the growth rate in the continuous dark (see Table S7 in the supplemental material). Transfer of the cultures from the dark into blue light stopped growth immediately. Strains in both cultures grew in the less intense blue light (3 μmol photons m⁻² s⁻¹), but the growth rate of strain in the AOA-DW culture was significantly lower in the low blue light than in the dark (Fig. 5; see Table S8 in the supplemental material).

### DISCUSSION

Enrichment of AOA cultures AOA-DW, AOA-AC2, and AOA-AC5. We enriched and characterized the growth of three different freshwater AOA belonging to thaumarchaeal group I.1a within the newly described phylum *Thaumarchaeota* (10, 52). The AOA in one of the cultures, AOA-AC2, is closely related to *Ca. Nitrosoarchaeum koreensis,* while the strains in the other two cultures, AOA-AC5 and AOA-DW, are only 70 to 82% (amoA) and 81 to 93% (16S rRNA gene) identical to other cultivated isolates and
enrichment culture strains, such as *N. maritimus* and *Nitrososphaera viennensis* (Table 1). This finding indicates that the AOA in these two enriched AOA cultures belong to a new genus of the ammonia-oxidizing *Thaumarchaeota*, assuming that the identity between the two genera is, on average, 96.4% on the basis of the 16S rRNA gene sequence (66). This new genus/group includes many ribotypes from non-salt water systems, such as freshwater systems (38; Li and Bollmann, unpublished) and drinking water systems (59), as well as soil and hot spring environments (67), as indicated by highly identical clones (Fig. 1).

**Pure cultures.** In this study, no pure cultures of the AOA were obtained. It is safe to assume that the heterotrophic satellite community is providing some compound that enabled the AOA to grow in the enrichment culture. Similar observations have been made with other AOA as well as with AOB. Potential compounds that positively influence the growth of AOA could be small organic compounds such as pyruvate, which improved growth and enabled isolation of *N. viennensis* (56). However, the addition of pyruvate during serial dilution has not led to isolation of any of these strains to date, indicating that different compounds might be important for different AOA. Further research will be necessary to elucidate the interactions between AOA (and AOB) and the heterotrophic satellite bacteria in ammonia-oxidizing enrichment cultures.

**Growth of AOA and AOB.** Overall, the growth experiments showed that the growth rates of the AOA were almost always lower than the growth rates of the AOB. All our experiments have been conducted under strict chemolithoautotrophic conditions. The results indicate that the strain in AOB-G5-7 had an advantage over the strains in the three tested AOA cultures under the conditions investigated. In nature, however, conditions are often less defined with respect to energy-generating processes. It has been suggested that not all *Thaumarchaeota* are chemolithoautotrophic ammonia oxidizers; some carry the *amoA* gene but are not actively oxidizing NH$_4^+$, and others utilize mixotrophic or heterotrophic lifestyles in pure and enrichment cultures (39, 56, 65). On the basis of these observations and our data, one could speculate that AOA in natural samples utilize a mixotrophic and/or heterotrophic lifestyle rather than a completely autotrophic lifestyle, which could explain their success in nature compared to that in the laboratory.

Increasing NH$_4^+$ concentrations had different influences on the growth rates and lag phases of AOA and AOB, with AOB growing faster and having shorter lag phases than AOA (Fig. 2; see Fig. S2 and Tables S3 and S4 in the supplemental material). After...
comparing these results with data provided by other studies that determined the $K_m$ of AOA for $\text{NH}_4^+$ to be approximately 1,000 times lower than the $K_m$ of AOB (27, 36, 43), we suggest that AOB have an advantage over AOA at higher $\text{NH}_4^+$ concentrations ($>10 \mu\text{M}$). This assumption is supported by the detection of high abundances of AOB in environments with higher $\text{NH}_4^+$ input due to fertilization and other processes, while AOA are more abundant in low-$\text{NH}_4^+$ and unfertilized environments (17, 21, 24, 62, 64).

The AOA in enrichment cultures AOA-DW and AOA-AC5 showed lower tolerance to high $\text{NH}_4^+$ concentrations than the strain in the AOA-AC2 culture, with the highest concentrations supporting growth at 1 mM $\text{NH}_4^+$ (AOA-DW) and 2 mM $\text{NH}_4^+$ (AOA-AC5). These concentrations are lower than the highest tolerances toward $\text{NH}_4^+$ observed for $N. \text{viennensis}$ (15 mM), “Ca. Nitrosoarchaeum koreensis” (10 mM), and the strain in the AOA-AC2 enrichment culture (5 mM), which is closely related to “Ca. Nitrosoarchaeum koreensis” (27, 56). These results indicate that the strains in the AOA-DW and AOA-AC5 cultures are less tolerant to high $\text{NH}_4^+$ concentrations than other AOA isolates and isolates in other enrichment cultures. Similar observations have been made for AOB; members of the Nitrosomonas oligotropha cluster, which are also commonly found in freshwater environments, are less tolerant to high $\text{NH}_4^+$ concentrations and better adapted to low $\text{NH}_4^+$ concentrations, while members of the Nitrosomonas europaea/N. eutropha cluster are primarily found in environments with high $\text{NH}_4^+$ concentrations (6, 7, 31, 32, 33).

AOA and AOB responded differently when cultured over a range of $O_2$ concentrations. The strains in the AOA-AC5 and AOA-DW cultures grew at all tested $O_2$ concentrations at the same rate, while the strain in the AOA-AC2 culture did not grow at 0.5% $O_2$, and the growth rate of the strain in the AOB-G5-7 culture decreased with decreasing $O_2$ concentrations (Fig. 3; see Table S5 in the supplemental material). Environmental surveys often detected AOA at the oxic-anoxic interface (4, 13, 17, 47), indicating an adaptation to low oxygen conditions. The low $K_m$ for $O_2$ found for $N. \text{maritimus}$ as well as other AOA (27, 36, 43) and the environmental data support the hypothesis that AOA are very likely better adapted to low $O_2$ than AOB and may therefore have a competitive advantage at the oxic-anoxic interface, while AOB are active under more aerobic conditions.

AOA and AOB grew at most of the tested pH values, with AOA growing at almost the same rate over a wide pH range and AOB showing a more bell-shaped curve, with the highest growth rate occurring at pH 7 to 7.5 (Fig. 4; see Table S6 in the supplemental material). AOA are found over a wide pH range in different environments, such as soils and hot springs (15, 19, 21, 40, 46), but most cultivated AOA, such as $N. \text{maritimus}$, $N. \text{viennensis}$, “Ca. Nitrosoarchaeum koreensis,” and “Ca. Nitrosotalea devanaterra,”
have rather narrow pH ranges for growth and activity compared with the AOA strains tested in enrichment cultures (27, 30, 56, 58).

The strain in AOB-G5-7 was more tolerant to light than the strain in AOA-DW and also recovered faster after exposure, while the strain in AOA-DW did not fully recover from light exposure (Fig. 5; see Tables S7 and S8 in the supplemental material). In the environment, maximum numbers of thaumarchael amoA were detected at levels below where photosynthetically active radiation (PAR) in the water column dropped to 0, indicating that no light was penetrating to this depth (4, 47). In the same study, AOB and AOA were detected in low abundance in more shallow waters of the Pacific, indicating that AOB as well as some AOA strains could be more tolerant to light than those that are the most abundant in the lower parts of the water column (47). The light response of AOB and AOA could be due to differences in the reaction of the copper-containing enzymes to light. AOB are very sensitive to blue near-UV light (23, 50). The authors discussed that this inhibition could be attributed to the absorption of light by the oxygcnated state of the copper-containing ammonioxygenase, which leads to inactivation of the enzyme (50). Genome studies of AOA showed a large number of copper-containing enzymes such as multicopper oxidases and blue copper proteins (5, 63), suggesting that some of the copper-containing enzymes in AOA could be sensitive to light as well, leading to inhibition of overall metabolism in AOA by light. During the preparation of the manuscript, Merbitz et al. (2012) published a study investigating the response of two AOB (Nitrosomonas europaea and Nitrosospira multiformis) and two AOA (N. maritimus and “Ca. Nitrosotalea devanaterra”) to white light (37). The study confirmed our findings.

Conclusion. The results of this study show that AOB are able to outcompete AOA under almost all conditions tested. These findings are in accordance with those of other cultivation-based studies, as well as observations made in the environment using molecular approaches. Further investigation must be done using other cultivation-based experiments, such as continuous cultures, which enable us to cultivate AOA and AOB under more stringently controlled conditions, and in situ incubations, which enable us to investigate the response of AOA and AOB to environmental changes under conditions which allow AOA and AOB to utilize metabolic functions as they would naturally in the environment.

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REFERENCES


