Effect of Different Ammonia Concentrations on Community Succession of Ammonia-oxidizing Microorganisms in a Simulated Paddy Soil Column

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Abstract

Ammonia oxidation is performed by both ammonia-oxidizing bacteria (AOB) and ammonia-oxidizing archaea (AOA). To explore the effect of ammonia concentration on the population dynamic changes of ammonia-oxidizing microorganisms, we examined changes in the abundance and community composition of AOA and AOB in different layers. Most of the archaeal amoA sequences were Nitrosotalea-related and the proportion that Nitrosotalea cluster occupied decreased in the surface layer and increased in the deep layer during the cultivation process. Nitrosopumilus-related sequences were only detected in the deep layer in the first stage and disappeared later. Both phylogenetic and quantitative analysis showed that there were increased Nitrosomonas-related sequences appeared in the surface layer where the ammonia concentration was the highest. Both AOA and AOB OTU numbers in different layers decreased under selective pressure and then recovered. The potential nitrification rates were 25.06 μg-N·L⁻¹·g⁻¹·dry soil·h⁻¹ in the mid layer which was higher than the other two layers. In general, obvious population dynamic changes were found for both AOA and AOB under the selective pressure of exogenous ammonia and the changes were different in three layers of the soil column.

Introduction

Nitrification is critical in the global nitrogen cycle and is a unique pathway linking reduced inorganic nitrogen with oxidised inorganic nitrogen in nature [1]. Ammonia oxidation is the first and rate-limiting step of nitrification. It was assumed that all ammonia oxidisers were bacteria until ammonia-oxidizing archaea were found. The findings of Venter et al. [2] and Treusch et al. [3] indicated the possibility of archaea involving in ammonia oxidation and the exist of ammonia-oxidation archaea was finally certified by the enrichments and pure cultures of several strains of AOA [4–10]. Recent research has demonstrated that ammonia-oxidizing archaea may form a separate and deep-branching phylum, the Thaumarchaeota [11–13].

Molecular biological research based on the functional gene amoA and the 16S rRNA gene has revealed a widespread distribution of AOA [14–17]. Quantitative PCR analyses found that AOA normally outnumbered their substrate competitors, such as AOB, in both soil [17] and water [18], but the relative contributions of AOA and AOB in the soil nitrogen cycle remained undetermined. Some researchers believed that AOB played a more important role than AOA in soil nitrification. Jia and Conrad [19] found that changes in potential nitrification rates were only correlated with the number of AOB amoA genes when they added substrate (NH₄⁺) or inhibitor (C₂H₂) of ammonia oxidation to the soil. Subsequently, Di et al. [20] discovered the same phenomenon in New Zealand grassland. In contrast, other researchers regarded AOA as the main drivers of ammonia oxidation in soil. Gubry-Rangin et al. [21] reported that the potential nitrification rates were positively correlated with the number and transcriptional activity of AOA amoA genes. Zhang et al. [22] confirmed the dominant role of AOA in soil nitrification by means of stable isotope probe and the quantitative analysis of archaial amoA genes. Olfre et al. [23] also demonstrated that the growth of only archaeal but not bacterial ammonia oxidizers occurred in microcosms with active nitrification. Differences in cell size, specific cell activity and other physiological characteristics may explain the different contributions of AOA and AOB to soil nitrification. In addition, AOA and AOB may compete mutually or exhibit functional redundancy under some conditions, whereas under other conditions the fundamental physiological differences between these organisms may lead to niche separation of AOA and AOB [24]. Therefore, environmental factors are important in determining the different nitrification activities and relative contributions of AOA and AOB.

Among all environmental factors, ammonia is the substrate of ammonia oxidation for which concentrations will directly affect nitrification activity. It was found that AOA were able to grow well and the growth of AOA was coupled with soil nitrification when the concentration of ammonia was relatively low or the supply of ammonia was through the mineralization of organic matter; however, AOB were more competitive in soil nitrification and the number of AOB amoA gene copies was greater than that of AOA.
when the concentration of ammonia was higher [19,20,23–26]. Adaptation to long-term energy stress is believed to be a crucial factor that distinguishes AOA from AOB [27]. Different specific affinities for substrate between AOA and AOB may explain their different growth patterns under low or high ammonia concentrations. *N. maritimus* exhibited a high affinity for ammonia and was able to grow and convert ammonia at an extremely low ammonia concentration (0.14 μg/L), while the ammonia-oxidizing activity was completely inhibited when the ammonia concentration reached 28 mg/L. The affinity of *N. maritimus* for ammonia can be more than 1,000-fold greater than that of *N. europaea* [29]. The affinity for ammonia of *N. viennensis* fell in between *N. maritimus* and *N. europaea*, and their growth was totally inhibited at ammonia concentration of 290 mg/L [7]. Other than ammonia concentration, other environmental traits such as pH [29], oxygen concentration [18] and organic carbon [30] could affect the abundance and diversity of ammonia-oxidizing microorganisms and consequently lead to niche separation of AOA and AOB.

Paddy soils are an important component of agricultural ecosystems in Asia, and these soils represent one of the major cultivated soil types in China. China has a long history of rice production, and the total area of the paddy soils in the country is 46 M ha [31]. Thus, research on nitrification in paddy soils is of great significance for improving our understanding of nitrogen fluxes in agricultural and soil ecosystems. The abundance and community structure of ammonia-oxidizing microorganisms in paddy soils have been described [32], but the mechanism underlying the response of these microorganisms to ammonia stress is unclear. Therefore, the main objective of this study was to verify the significant role that ammonia concentrations play in the population shifts of AOA and AOB in paddy soils.

**Materials and Methods**

**Ethics Statement**

No specific permits were required for the described field studies.

**Simulated Soil Column and Sampling**

The flooded paddy soil used in this study was collected from paddy land (30°26’N, 120°19’E) in Zhejiang Province, China. The upper 50 cm of paddy soil was collected and filled into the soil column. The soil column was divided into three layers due to the definition and classification of soil layer in pedology and the setting of sampling ports, the surface layer (2–5 cm), the mid layer (17–20 cm) and the deep layer (47–50 cm), to investigate the differences in ammonia-oxidizing microorganisms at different depths.

The simulated soil column (Fig. 1) was made of plexiglass and had a height of 70.0 cm and a diameter of 12.5 cm. It had a working volume of 2.5 L and a total volume of 3.0 L. Spherical fillings placed at the bottom of the column acted as a filter to avoid loss of biomass. Paddy soil in agricultural ecosystem were usually irrigated with water from river or rainfall, so natural river water was used in this study to simulate the in situ environment of paddy soil. Therefore, the experimental medium was prepared with natural river water (NH4+–N, 0.53±0.1 mg/L; NO2–N, 0.1±0.03 mg/L; NO3–N, 3.72±0.5 mg/L), filtrated by filter) adding ammonia, with final ammonia concentration of 42 mg/L, medium was introduced at the top of the column by spraying (5 L per day), and then permeated to the deeper layer. Due to the consumption of microorganisms and adsorption by the soil, an ammonia concentration gradient formed in the soil column from the surface layer to the deep layer (surface layer, 143 μg NH4+–N g⁻¹ dry soil; mid layer, 61.9 μg NH4+–N g⁻¹ dry soil; deep layer, 39.9 μg NH4+–N g⁻¹ dry soil). The soil column was placed in thermostatic chamber and the temperature was 30±1°C. From the top layer to the deep layer, pH of the soil were 7.24±0.12, 7.41±0.1, 7.09±0.15 and the humidity were 38.7±1%, 36.5±0.8% and 34.4±1.5%, respectively. The oxygen concentration in the top soil of the column was 2.48 mg/L.

**Potential Nitrification Rates**

The potential nitrification rates in the soil layers were determined via batch tests using 150 mL conical flasks containing 80 mL of liquid. The medium for determining the rates contained 3.5 mg/L NH4+-N. Medium with sterilized soil was incubated as a control. Tests were performed in triplicate for 8 hours at 30°C in the dark, and the concentrations of NH4+-N, NO2–N and NO3–N were periodically monitored per 1.5 hours during incubation. Potential nitrification rates were characterized on the basis of the consumption of NH4+-N. In consideration of some ammonia-oxidizing microorganisms which could accumulate substrates intracellularly [33], the results of potential nitrification rates we obtained may be to some extent higher than the real situation.

**DNA Extraction and PCR Amplification**

DNA was extracted using the Power Soil DNA Kit (Mo Bio Laboratories, California, USA) following the manufacturer’s instructions. The extracted DNA was examined in a 1.0% agarose gel by electrophoresis.

![Figure 1. The simulated paddy soil column used in this study.](image-url)

(1) inlet pail, (2) peristaltic pump, (3) timer, (4) main body of soil column, (5) sprayer, (6) beaumontage, (7) outlet pipe, (8) sample connection.
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Primers sets Arch-amoAF, Arch-amoAR [34] and amoA-1F, amoA-2R [35] were used for the amplification of archaeal and bacterial amoA genes, respectively. The PCR thermal cycling programs employed were described previously [36]. The amplified products were examined in a 1.0% agarose gel by electrophoresis.

Cloning and Sequencing

The obtained PCR products were cloned using the pMD19-T vector (TaKaRa, Bio Inc., Shiga, Japan) according to the manufacturer’s instructions. Plasmid DNA was isolated using the Gene JET™ Plasmid Miniprep kit (Fermentas Life Sciences, Burlington, Canada) and was digested with 5 U of EcoRI enzyme in EcoRI buffer for 1.5 h at 37°C. The digestion products were examined for an insert of the expected size using agarose (1.0%) gel electrophoresis. At least 20 positive clones from each library were randomly selected for sequencing using an ABI3100 automated sequencer (Applied Biosystems, California, USA).

Phylogenetic Analyses

All of the amoA gene sequences obtained were imported into MEGA4.1 to construct alignment files in combination with the most similar environmental sequences and sequences of known strains of AOA and AOB. Phylogenetic trees were constructed using the neighbour-joining method. Bootstrap analysis with 1000 replications was applied to estimate the confidence values for the tree nodes.

Statistical Analyses

Operational taxonomic unit (OTUs) was used to determine the archaeal and bacterial amoA gene diversity in different soil samples. For both AOA and AOB, researchers found that 1%–3% 16S rRNA distance equates to a 15% amoA gene distance [37,38]. 85% sequence identity at the amoA gene level could be regarded as an approximate threshold below which ammonia-oxidizing microorganisms can be assigned to different species. Hence, OTUs were defined by 15% differences for both archaeal amoA gene and bacterial amoA gene in nucleotide sequences. OTU numbers were determined by the farthest neighbour algorithm in the DOTUR program [39]. The coverage of the clone libraries was calculated as $C = [1 - (n_{i}/N)] \times 100$, where $n_{i}$ is the number of unique OTUs and $N$ is the total number of clones in a library.

Real-time Quantitative PCR (qPCR)

The primer used to quantify the copy numbers of amoA genes of AOA and AOB were the same with those used in the clone library construction. Primer pair P128r, P365r designed by Dang [40] was used to quantify the copy numbers of Nitrosomonas amoA gene. qPCR was performed using an iCycler iQ5 thermocycler and a real-time detection system (Bio-Rad, California, USA) according to the method described by Francis et al. [34], McTavish et al. [41] and Dang [40]. Quantification standards consisted of $10^{-3}$, $10^{-4}$, $10^{-5}$, $10^{-6}$ copy number dilution series of plasmid containing inserted amoA genes of AOA, AOB and Nitrosomonas, respectively. $R^2$ and the amplification efficiencies of the AOA, AOB and Nitrosomonas standard curves were 0.996, 0.998 and 0.996 (as shown in Figure S3, S4 and S5) and 96%, 89% and 95%, respectively.

Analytical Methods

Measurements of NO$_3$-N, NO$_2$-N and NH$_4$+-N in the experimental medium were performed according to standard methods [42]. The total solid (TS) content was measured by the weighing method after drying at 105°C. The inorganic nitrogen contents in the soils were determined colorimetrically via flow injection analysis of soil-KCl extracts as previously described [43].

Accession Numbers

Sequences were deposited in GenBank under accession numbers JQ320498 to JQ320554.

Results

Diversity and Phylogenetic Analysis of AOA and AOB

105 derived archaeal amoA genes belonged to Nitrosotalea cluster, Nitrososphaera sister cluster, Nitrospumilus cluster and Nitrospumilus cluster according to a new nomenclature system for archaeal amoA genes reported by Pester [37] (Figure 2), but no Nitrosocatulue-related sequences was detected. The sequences affiliated with Nitrosotalea cluster were the most and accounted for 69.2% of all the archaeal amoA genes obtained. Only two sequences were detected belonging to Nitrospumilus cluster and the rest sequences belonged to Nitrososphaera cluster and Nitrososphaera sister cluster. A total of 9 OTUs were obtained using the DUTOR program based on a 15% amoA gene sequence distance cutoff [37] for all 105 AOA amoA sequences detected in this study. Nitrospumilus cluster contained only one OTU and the other three clusters contained at least two OTUs. Sequences belonging to OTU1 were found in all three soil layers during the entire cultivation process, totalling 121 sequences, which accounted for 65.4% of the detected AOA amoA gene sequences.

As to AOB, all 201 sequences obtained for the AOB amoA gene in the clone libraries belonged to β-proteobacteria, as shown in Figure 3. Among these sequences, 91.0% were grouped into the Nitrosopumilus genus, and the remainders were grouped into Nitrosomonas. A total of 8 OTUs were obtained for the bacterial amoA genes using the DUTOR program based on a 15% amoA gene sequence distance cutoff [38]. Nitrososphaera cluster contained 2 OTUs, and the remainders were grouped into Nitrosopumilus cluster.

Community Succession of AOA and AOB

The community structure of the AOA and AOB in the soil column changed during the cultivation process and was closely related to soil depth. An obvious population shift was found for AOA in different layers when the proportions that each cluster occupied in the first stage and the last stage were compared. In the surface layer, the proportion of Nitrosotalea cluster decreased from 66.7% to 52.9% (Figure 4). Accordingly, an increase from 20% to 47.1% was found for Nitrosopumilus cluster. The sequences affiliated with Nitrososphaera sister cluster disappeared during the 18-month cultivation process. Interestingly, in the mid layer, we found that cluster Nitrospumilus-related sequences represented more than 94.4% of the sequences detected, and this proportion did not change during the cultivation process. Accordingly, the proportion of Nitrosomonas-related sequences was less than 5.6% throughout the cultivation process. In the deep layer, the observed changes were dramatic. In the first stage, the diversity of the population was high and all the four AOA amoA clusters were found, the Nitrospumilus-related sequences were only detected in the first stage of the deep layer and represented 9.1% of the sequences detected. However, 18 months later, the sequences belonging to Nitrosopumilus disappeared and the proportion that Nitrospumilus cluster occupied increased from 27.3% to 72.2%. In the 18-month cultivation process, the number of OTUs in all three layers decreased first and then increased. The number of OTUs in the deep layer was greater than in the other two layers in the first stage (Figure S1). In the second stage, the numbers of OTUs in all three layers were reduced dramatically. In the surface layer, the number of OTUs
dropped from 4 to 1, while it decreased from 2 to 1 and from 7 to 3 in the mid and deep layer, respectively. The only difference was that the number of OTUs in the surface and mid layers recovered more rapidly than was observed for the deep layer. The number of OTUs in the surface and mid layers began to increase in the third stage; however, the number of OTUs in the deep layer did not increase until the last stage.

With respect to AOB, increasing sequences belonging to *Nitrosomonas* were detected in all three layers in the soil column, and the proportions that *Nitrosomonas*-related sequences occupied were 25%, 15.8% and 18.8% from the surface layer to the deep layer (Figure 5). The numbers of AOB OTUs varied in a similar way as was observed for the AOA OTUs. The AOB OTU numbers in the mid and deep layers decreased in the first three stages and increased in the last stage. However, in the surface layer, the number of OTUs remained stable at three after the third stage (Figure S2).

Real-time Quantitative PCR (qPCR)
The AOA and AOB *amoA* genes in the three soil layers were quantified every 6 months. AOA were more abundant than AOB in almost all soil samples, with the exception of the sample from the surface layer in the last stage, in which the AOB/AOA ratio was 16.95. The ratios of AOA/AOB ranged from 1.03–12.96 in the other 11 soil samples (Figure 6). The copy number of *Nitrosomonas amoA* genes increased substantially in the surface layer during the cultivation process, from $2.2 \times 10^6$ to $1.2 \times 10^8$. As for the other two layers, the change of was not obvious (Figure 7).

Nitrification Activity
Nitrification activity was tested after the 18-month cultivation period. Samples were collected from the three different layers of the soil column. Different nitrification activities were observed for the three layers of the soil column. The maximum specific nitrification activity in the mid layer was
layers were 16.13, 9.97, 25.06, 10.80, 11.14, 16.13, 9.97 μg·N·L⁻¹·g⁻¹·h⁻¹, respectively. Therefore, ammonia-oxidizing microorganisms in the mid layer exhibited the highest nitrification activity among the three layers of the soil column.

**Discussion**

**Community Succession of AOA and AOB**

It was found that 69.2% sequences derived in our soil column were related to *Nitrosotalea devanaterra*-the only ammonia-oxidizing archaea from acid soil [6]. This was in accordance with the high-throughput sequencing results reported by Gubry-Rangina et al. [44], which demonstrated that *Nitrosotalea devanaterra*-related sequences may be predominant in some agricultural soils. pH in our soil column were neutral or slightly alkaline and was not in accordance with the acid environment where *Nitrosotalea devanaterra* was found. The contradictory could be explained by that there may be existed AOA belonging to *Nitrosotalea devanaterra* cluster which could survive and oxidize ammonia to nitrite in neutral or slightly alkaline environment. The community composition of the AOA in the different layers of the soil column underwent great changes during the cultivation process. In the surface layer, the sequences belonging to *Nitrososphaera* sister cluster disappeared after 18-month cultivation. Since there was no isolated or enrichment strain of this AOA cluster [37], the physiochemical characteristics of *Nitrososphaera* sister cluster was unknown. Combined with the increased *Nitrososphaera*-related sequences in the surface and the decreased *Nitrososphaera*-sister-related sequences in the deep layer, we deduced that the AOA affiliated with *Nitrososphaera* sister cluster had relative high affinity for ammonia and prefer to live in the environment with relative low ammonia concentration. *Nitrososphaera viennensis* [7] could grow well in the media containing ammonia concentration as high as 15 mM, indicating a low affinity of this strain of AOA. So, it was reasonable that the proportion *Nitrososphaera*-related sequences occupied increased in the last stage compared with the first stage in the surface layer, since it preferred conditions with relatively higher ammonia concentration. Correspondingly, in the deep layer, the proportion *Nitrososphaera*-related sequences occupied decreased from 59.1% to 16.7%. *Nitrosopumilus maritimus* showed extremely high affinity for ammonia and the ammonia inhibition concentration was as low as 2 mM to 3 mM [5], so it was not strange that the *Nitrosopumilus*-related sequences disappeared in the deep layer after exogenous ammonia was added during the cultivation process. No *Nitrososphaera*-related sequence was detected in the surface and mid layer during the whole 18 months. *Nitrosotalea*-related sequences were the dominant species in our soil column throughout the cultivation process. In the surface layer, the proportion that the *Nitrosotalea*-related sequences occupied decreased from 66.7% to 52.9%; on the contrary, the proportion in the deep layer increased from 27.3% to 72.2%. It was reported that the acid AOA could grow well with the total NH₃&NH₄⁺ concentration ranged from 10 μM to 10 mM (equivalent to 0.18 nM to 0.18 μM of NH₄⁺) at pH 4.5 and no growth was detected at 50 mM and 100 mM (equivalent to 9 μM to 18 μM NH₃, pKₐ for NH₃: NH₃⁺, 9.25) [6]. The available NH₃ concentration in the medium was 44μM (pH, 7.24; 30°C), which was much higher than the ammonia concentration.
inhibition concentration (9 µM) of Nitrosotalea decantarrens. So, the sequences affiliated with Nitrosotalea cluster decreased in the surface layer where the ammonia concentration was relatively higher during 18 months cultivation process.

In our simulated soil column, we found that all three layers contained a certain amount of Nitrosomonas-like sequences after 18 months. In the surface layer the proportion occupied by Nitrosomonas-like sequences was 25.0%, which was higher than in the other two layers (mid layer, 15.8%; deep layer, 18.8%). Quantitative analysis also reached the similar conclusion that the copy number of Nitrosomonas amoA genes increased from 2.2 × 10^6 to 1.2 × 10^7 in the surface layer, and the copy numbers were much higher than that in the other two layers. This could be explained by differences in physiological characteristics between Nitrosomonas and Nitrosospira. Schramm et al. [45,46] have reported that the growth of Nitrosomonas species followed an r-strategy. The Nitrosomonas genus had a low ammonia affinity and was adaptable to conditions where the concentration of ammonia was high. In contrast, the growth of Nitrosospira species followed a k-strategy. The Nitrosospira genus had a high ammonia affinity and was adaptable to conditions where the concentration of ammonia was low. The appearance of and increases in Nitrosomonas-like sequences were reflection of the competitive advantage of Nitrosomonas species under higher ammonia concentrations when exogenous ammonia was added. The obtained Nitrosomonas-like sequences were primarily found in the surface layer because of the relatively higher ammonia concentration in this layer compared with the other two layers. Notably, no Nitrosomonas-related sequences was found in the first stage through construction of AOB clone libraries, but we indeed detected Nitrosomonas amoA gene using qPCR, this may be due to the different specificities of the primers used in phylogenetic and quantitative analysis.

The richness of AOA and AOB decreased under ammonia stress and then recovered after a period of adaptation, as reflected in the variation in OTU numbers. It seemed that the AOA in the deep layer were less sensitive to exogenous ammonia than those in the other two layers and had a relative longer adaptation period. Another explanation of the OTU numbers variation was the cultivation effect that led to the survival of only a few adapted species which take over the community. For instance, in the deep layer of the first stage, AOA population was constituted with OTU1-3 and OTU6-9, and 6 months later just OTU1, OTU7 and OTU9 were left. In the last stage, the population was composed with 4 OTUs and a new OTU was found even though it had never been derived in the first three stages, so finally the population was taken over by the survival species who could adapt to the selective pressure of the environment.

**Conclusion**

In general, obvious population dynamic changes were found for both AOA and AOB under the selective pressure of exogenous ammonia and the changes were different in three layers of the soil column.

**Supporting Information**

Figure S1 Changes in AOA OTU numbers in different layers during the 18-month cultivation process. (TIF)

Figure S2 Changes in AOB OTU numbers in different layers during the 18-month cultivation process. (TIF)

Figure S3 The standard curve of qPCR for AOA amoA genes. (TIF)

Figure S4 The standard curve of qPCR for AOB amoA genes. (TIF)

Figure S5 The standard curve of qPCR for Nitrosomonas amoA genes. (TIF)
Author Contributions
Conceived and designed the experiments: LLP XXX ZP. Performed the experiments: HBL LS SLD. Analyzed the data: HBL LS LLP. Contributed reagents/materials/analysis tools: LLP XXX ZP HBL. Wrote the paper: HBL LS.

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