Biogeography of anaerobic ammonia-oxidizing (anammox) bacteria

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INTRODUCTION

The anaerobic ammonia oxidation (anammox) process converts ammonia to N₂ gas by using nitrite as electron acceptor under anoxic conditions (van de Graaf et al., 1995). This process is important for removing fixed N from both engineered and natural systems and can be applied to wastewater treatment in order to replace conventional treatment systems. Anammox is cost effective and environmentally friendly because it does not require aeration or organic carbon inputs, and reduces the production of greenhouse gases (i.e., N₂O and CO₂) compared to conventional denitrification (Jetten et al., 1997; van Dongen et al., 2001); anammox was first implemented in a full-scale wastewater treatment plant (WWTP) in Rotterdam, Netherlands (van Dongen et al., 2001; Abma et al., 2007; van der Star et al., 2007). Although anammox bacteria were first discovered in WWTPs and their applications have been studied worldwide, they may account for more than 50% of N loss from marine environments (Arrigo, 2005; Francis et al., 2007). However, recent reports estimate that anammox bacteria contribute 23–30% to N loss from marine environments (Trimmer and Engström, 2011; Dalsgaard et al., 2012; Babbin et al., 2014). The contributions of anammox bacteria to biogeochemical N₂ production were measured as 18–36% in groundwater (Moore et al., 2011), 4–37% in paddy soils (Zhu et al., 2011), 9–13% in lakes (Schubert et al., 2006), and 1–8% in estuaries (Trimmer et al., 2003). These results indicate that anammox bacteria play a key role in the global N cycle.

Anammox bacteria branch deeply within the Plantomycetes phylum. There are five known anammox genera, with 16 species proposed to date. The first discovered anammox bacterium was Ca. Brocadiia anammoxidans, enriched from a denitrifying fluidized bed reactor (Mulder et al., 1995; Kuinen and Jetten, 2001). The three characterized species within the Ca. Brocadiia genera are Ca. Brocadiia fulgida (Kartal et al., 2008), Ca. Brocadiia sinica (Oshiki et al., 2011), and Ca. Brocadiia caroliniiensis (Rothrock et al., 2011); all of these were enriched in anammox bioreactors. The only species reported within the Candidatus Kuenenia genus is Ca. Kuenenia stuttgartiensis, which was isolated from a trickling filter biofilm (Schmid et al., 2000). The Ca. Scalindua genus consists of nine proposed species, six of which were discovered in marine environments (Kuyper et al., 2003; Woebken et al., 2008; Hong et al., 2011a; Fuchsman et al., 2012; Dang et al., 2013; van de Vossenberg et al., 2013). Ca. Scalindua sorokinii was the first anammox species found in a natural environment (the Black Sea; Kuyper et al., 2003). Ca. Scalindua richardii was also recovered from the Black Sea (Fuchsman et al., 2012). Although these two species originated from the Black Sea, they dominated in different zones. A cluster associated with Ca. Scalindua sorokinii was detected in the lower suboxic zone where ammonium concentration was high, but nitrite concentration was low.
whereas a cluster associated with *Ca.* Scalindua richardssii was found in the upper suboxic zone where ammonium concentration was low, but nitrite concentration was high (Fuchshman et al., 2012). *Ca.* Scalindua brodae and *Ca.* Scalindua wagneri were both identified in WWTPs (Schmid et al., 2003). *Ca.* Scalindua arabica originated in the Arabian Sea and the Peruvian oxygen minimum zone (OMZ; Woebken et al., 2008). *Ca.* Scalindua pacifica (Dang et al., 2013) and *Ca.* Scalindua profunda (van de Vossenberg et al., 2013) were retrieved from the Bohai Sea and a marine sediment of a Swedish fjord, respectively. Two additional species names were tentatively proposed from molecular surveys: *Ca.* Scalindua sinooilfield from a high temperature petroleum reservoir (Li et al., 2010) and *Ca.* Scalindua zhenghei from marine sediments (the South China Sea; Hong et al., 2011a). The only known species affiliated with the *Ca.* Anammoxoglobus genus was *Ca.* Anammoxoglobus propionicus, enriched from an anammox reactor (Kartal et al., 2007). *Ca.* Jettenia asiatica was retrieved from a granular sludge anammox reactor (Quan et al., 2008). Notably, known anammox bacteria species have mostly been discovered in engineered environments, but they have commonly been detected in various natural ecosystems and are more widespread than previously thought. However, it should be noted that *Ca.* Scalindua sinooilfield and *Ca.* Scalindua zhenghei are not in the category Candidatus on the list of prokaryotic names with standing in the nomenclature (LPSN) website. The classification and nomenclature of anammox *Ca.* species need to be better clarified and standardized in the future.

Observations of anammox bacterial diversity have demonstrated that *Ca.* Brocadia, *Ca.* Kuenenia, and *Ca.* Anammoxoglobus were commonly found in non-saline environments (i.e., Egli et al., 2001; Moore et al., 2011; Hu et al., 2013), whereas *Ca.* Scalindua dominated saline environments (i.e., Woebken et al., 2008; Hong et al., 2011a; Villanueva et al., 2014), including deep-sea methane seep sediments (Shao et al., 2014). Anammox bacteria have also been detected in extremely saline-related environments, including hydrothermal vents (Byrne et al., 2009; Russ et al., 2013), and cold hydrocarbon-rich seeps (Russ et al., 2013). However, because all previous molecular surveys of the anammox 16S rRNA genes were from individual studies of specific habitats, the overall understanding of global anammox bacterial diversities, distributions, and co-occurrences among lineages remains unclear.

Factors affecting anammox bacterial diversity and distribution have been investigated within individual habitat-specific studies. For example, organic carbon influenced anammox diversity in freshwater sediment (Hu et al., 2012b), soil (Shen et al., 2013), and an estuary (Hou et al., 2013). Ammonium and nitrite concentrations correlated with anammox diversity in a mangrove sediment (Li et al., 2011). Temperature impacted anammox communities in freshwater sediment (Osaka et al., 2012) and an estuary (Hou et al., 2013). Depth affected anammox diversity in marine sediment (Li et al., 2013). However, no comprehensive survey has previously explored factors that govern global anammox distributions.

The main objectives of this study were to investigate global anammox bacterial distributions and identify factors influencing anammox bacterial distributions and diversity. Over 6000 anammox 16S rRNA gene sequences from Genbank were collected and analyzed by both phylogenetic and multivariate statistical methods. An anammox 16S rRNA gene phylogenetic tree revealed broad anammox distributions across habitats, including marine sediment, marine water column, estuary, mangrove sediment, soil, freshwater, freshwater sediment, groundwater, reactor, WWTP, marine sponge, biofilter, fish gut, shrimp pond, and oil field. Co-occurrence analysis demonstrated strong relationships among dominant anammox phylotypes. Global distributions of anammox bacteria revealed factors that influence anammox bacterial distributions, with salinity being the most important environmental variable. This study provides a better understanding of the prevalence of anammox bacterial 16S rRNA genes across habitats and the key factors impacting their distribution patterns.

**MATERIALS AND METHODS**

**DATA COLLECTION AND PREPARATION**

All anammox 16S rRNA gene sequences available in Genbank were extracted on October 25th, 2013. In total, 14,790 potential anammox-related sequences were collected using the following keyword searches: “uncultured planctomycete 16S ribosomal RNA gene,” “anammox bacterium 16S ribosomal RNA gene,” “anaerobic ammonium-oxidizing bacterium 16S ribosomal RNA gene,” “Candidatus Brocadia 16S ribosomal RNA gene,” “Candidatus Scalindua 16S ribosomal RNA gene,” “Candidatus Kuenenia 16S ribosomal RNA gene,” “Candidatus Anammoxoglobus 16S ribosomal RNA gene,” and “Candidatus Jettenia 16S ribosomal RNA gene.” Most anammox bacterial 16S rRNA gene sequences were deposited in the Genbank with the definition “uncultured planctomycete 16S ribosomal RNA gene” (data not shown). However, this keyword-based search retrieved both anammox and non-anammox sequences. All collected sequences were searched by BLAST against known anammox species in Genbank core reference set and aligned by QIIME v1.7 (Caporaso et al., 2010) using Infernal (Nawrocki and Eddy, 2013) against the Greengenes database (May 2013 revision; DeSantis et al., 2006) to screen for anammox-related sequences. After removing non-anammox and low quality sequences, over 6000 sequences from >200 isolation sources were included in the analysis. All anammox sequences from across many specific “Isolation source” Genbank designations were assigned to 15 general habitats: marine sediment, marine water column, estuary, freshwater sediment, freshwater, groundwater, soil, mangrove sediment, WWTP, reactor, marine sponge, biofilter, fish gut, oil field, and shrimp pond.

Limitations of this analysis included metadata inconsistencies and missing environmental parameters across multiple studies. Consequently, metadata were qualitatively grouped into three broad categories: salinity (saline, mixed, and non-saline environments), ecosystem (natural and engineered), and habitat (listed above). Another limitation was that it was not possible to consistently determine relative abundances of anammox sequences within each study due to inconsistencies with reporting, sampling efforts, and methodologies. To address this shortcoming, all anammox 16S rRNA gene sequences were clustered into operational taxonomic units (OTUs) at 97% identity with cd-hit-est
was only counted as present or absent for each study.

**STATISTICAL AND MULTIVARIATE ANALYSES**

Individual studies that contributed anammox 16S rRNA gene sequences were usually associated with unique Genbank isolation sources. Because of this, the numbers of anammox 16S rRNA gene sequences contributed per study and/or unique isolation source were broad, ranging from 1 to 623 sequences. In order to ensure that dissimilarity matrices were generated from datasets derived from the same number of sequences from each study, multiple rarefied datasets were generated that varied in the number of sequences derived from each study/isolation source. In cases where multiple studies represented compatible isolation sources, we tested datasets rarefied to 10, 40, or 100 sequences from each isolation source. Subsequently, we tested datasets rarefied to 10, 40, or 100 sequences from each isolation source category.

After clustering the sequences at 97% identity, all sequences were aligned and trimmed in order to consider a single homologous spanning region of the 16S rRNA gene, which corresponded to the positions 384–834 of *Escherichia coli* (J01695;2; Brosius et al., 1978). Any sequences with less than 100 bases after trimming were discarded from the analysis. Consequently, the sequences from some isolation sources within five minor habitats (marine sponge, biofilter, fish gut, oil field, and shrimp pond) fell below the threshold for rarefied datasets 40 and 100 sequences. All five of these minor habitats were removed from further analysis. The minimum sequence threshold remained at 10, 40, and 100 after being trimmed. Consequently, 10 major habitats (marine sediment, marine water column, estuary, freshwater sediment, freshwater, groundwater, soil, mangrove sediment, WWTP, reactor) were considered in this analysis.

Principal coordinates analysis (PCoA) ordinations were generated from unweighted UniFrac distance matrices (Lozupone and Knight, 2005) through QIIME (Caporaso et al., 2010). Non-metric multidimensional scaling (NMDS) ordinations were calculated based on a Jaccard dissimilarity matrix, using the AXIOME pipeline (Lynch et al., 2013). To test treatment effects and within-group agreement, multi-response permutation procedures (MRPP) were tested on 999 permutations, using the R library vegan (Oksanen et al., 2008) from within AXIOME. Analyzed data for each rarefied dataset (10, 40, and 100 sequences), including the OTU table with taxonomic classifications and the analyzed sequences, a mapping file, and the source FASTA files, are in a single compressed Supplementary Material file (“Sonthiphand supp data files.zip”). All collected sequences, with corresponding Genbank accession numbers and metadata, are provided in a spreadsheet (sequences.xlsx) within the Supplementary Material.

**RAREFACTION CURVE AND DIVERSITY INDICES**

Rarefaction curves, observed species, phylogenetic diversity (PD), Chao1, and Shannon indices were generated by QIIME (Caporaso et al., 2010). The Wilcoxon Signed-rank test was performed by the R function wilcox.test (R Core Team, 2013). The null hypothesis was that the number of OTUs between habitats was the same. If $p$ was $\leq 0.05$, the null hypothesis was rejected.

**PHYLOGENETIC CONSTRUCTION**

Representative sequences for each OTU from each habitat were selected for phylogenetic analysis. A total of 505 OTU sequences from across all 15 habitats included all known anammox *Candidatus* species. Outgroups included cultured non-anammox species of *Planctomycetes*, including *Planctomyces maris* (X62910), *Isophaera* sp. (X81958), *Gemmatia obscuriglobus* (X85248), *Blastopirellula marina* (HE861893), *Rhodopirellula baltica* (FJ624346), and *Pirellula* sp. (X81942). Sequences were aligned using MUSCLE (Edgar, 2004) and trimmed to a final homologous length of ~310 bases. A maximum likelihood tree was constructed with the PhyML v.3.0.1, using the GTR model (Guindon and Gascuel, 2003). The tree topology was optimized at five random starts. The approximate likelihood ratio test (aLRT) was conducted to provide tree topology support. The phylogenetic tree was visualized by SEAVIEW (Galtier et al., 1996).

**CO-OCCURRENCE NETWORK ANALYSIS**

Anammox sequences were sorted by habitat and an OTU table was generated by AXIOME. Co-occurrence was assessed using a previously described method (Barberán et al., 2012). All singletons were discarded, and OTUs having a Spearman’s correlation $\geq 0.8$ were considered to have a strong co-occurrence relationship. Spearman’s correlation was used because it only checks if two OTUs are monotonically related, rather than having a linear relationship. As a result, it is less sensitive to differences in abundance, and this was desirable because abundance information may have been lost when the sequences were deposited in GenBank, as described above. The results were visualized with Gephi (Bastian et al., 2009).

**RESULTS**

**DISTRIBUTIONS OF ANAMMOX BACTERIA ACROSS HABITATS**

Anammox sequences were collected from multiple studies and isolation sources. The number of sequences was considerably different from one isolation source to another. Three rarefied sequence collections were generated to compare distribution patterns. Because the broad range of analyzed sequences (10–623 sequences) affected dissimilarity measurements, we chose to analyze set 40 in more detail to include as many isolation sources as possible in our analysis while maximizing sequence sample size (Figure 1). This was done because set 10 (i.e., 10 sequences per isolation source) showed poor groupings with low correlations (data not shown) and both set 40 and set 100 showed similar distribution patterns with high correlations (Figure 2).

All anammox sequences from 10 habitats were visualized within an ordination plot based on phylogenetic distances by using an unweighted UniFrac distance matrix (Figure 1). The percentage of PCoA principal coordinates (PC1 and PC2) explained 46% variability among all samples. The ordination demonstrated that anammox sequences clustered significantly by habitat (Figure 1B), which was supported by MRPP ($T = -7.6, A = 0.14, p < 0.001$; Figure 2). All anammox sequences clustered separately into two main groups (Figure 1B).
Marine sediment, marine water column, estuary, and mangrove sediment grouped together and were dominated by *Ca. Scalindua* cluster (Figures 1A,B). The WWTP, reactor, soil, freshwater, freshwater sediment, and groundwater grouped together and were dominated by *Ca. Brocadia*, *Ca. Jettenia*, and the unknown cluster. Four samples, one each from freshwater, freshwater sediment, soil, and WWTP, were present in both groups.

**KEY FACTORS AFFECTING GLOBAL ANAMMOX BACTERIAL DISTRIBUTION**

The strongest separation of anammox bacterial sequences was linked to sample salinity (Figure 1C), which we assigned qualitatively as saline, “mixed,” and non-saline environments. The mixed environments were generally river-marine transitional zones, mostly from mangrove and estuary habitats. Saline and mixed environments clustered together and differed significantly from non-saline environment (Figures 1C, 2B; $T = -12.1, A = 0.09, p < 0.001$). However, a few non-saline samples grouped with saline and mixed samples. The *Ca. Scalindua* cluster was clearly dominant in saline environments but almost never detected in non-saline environments (Figures 1A,C). The major complement of anammox bacteria found in non-saline environment was *Ca. Brocadia*, *Ca. Jettenia*, and the unknown clusters. The results indicated that salinity was the key factor governing global distributions of anammox bacteria.

**DISTINCT ANAMMOX BACTERIA IN NATURAL AND ENGINEERED ECOSYSTEMS**

Another factor that showed a significant correlation with the anammox bacterial distributions was ecosystem type. Although...
most anammox sequences were from natural ecosystems, those from engineered ecosystems grouped together (Figures 1D, 2C; $T = -9.1, A = 0.05, p < 0.001$). However, one sample from a WWTP grouped separately from other samples of engineered ecosystems (Figure 1D). This WWTP sample contained very few anammox sequences associated with Ca. Scalindua cluster. More robust group separation was visualized by the NMDS generated from an OTU-based Jaccard distance metric (Figures 2C,F). This observation demonstrated environmental selection of anammox bacteria in natural and engineered ecosystems.

**DIVERSITY RICHNESS OF ANAMMOX BACTERIA**

Rarefaction curves and diversity indices showed that freshwater possessed the highest anammox bacterial diversity, whereas the marine water column was associated with the lowest diversity (Figure 3 and Table 1). The diversity of anammox bacteria in freshwater and marine water column differed significantly ($p = 0.01$). The diversity of anammox bacteria in freshwater and freshwater sediment was not significantly different ($p = 0.22$). Rarefaction curves of freshwater showed no saturation, although only 170 sequences were analyzed. The majority of
freshwater anammox sequences were from unpublished data; only a few publications reported anammox bacterial 16S rRNA gene sequences from freshwater (Schubert et al., 2006; Hamersley et al., 2009; Pollet et al., 2011; Han and Gu, 2013; Sonthiphand and Neufeld, 2013). Consequently, more research on anammox bacteria in freshwater would be required to confirm this observation. Overall, the results imply that most novel anammox clusters remain undiscovered within freshwater habitats.

The diversity of anammox bacteria in marine sediments was higher than in the marine water columns \((p = 0.02; \text{Figure 3, Table 1})\). The reason for this observation might be higher physical and biogeochemical heterogeneity in marine sediments, associated with a greater overall microbial diversity (Table 1). The diversity of anammox bacteria among other isolation source samples, including freshwater sediment, estuary, mangrove sediment, soil, and marine sediment, showed no significant differences (Figure 3, Table 2). The diversity of anammox bacteria in engineered ecosystems, including WWTPs and reactors, were not significantly different \((p = 0.15)\), consistent with the observation that anammox bacteria from engineered ecosystems grouped together (Figures 1D, 2C,F).

Although groundwater, freshwater, and freshwater sediment were non-saline isolation sources, the diversity of groundwater was low and significantly different from freshwater \((p = 0.01)\) and freshwater sediment \((p = 0.01; \text{Table 1})\). However, the interpretation of this observation must be cautious because only a few publications have surveyed anammox bacterial 16S rRNA gene sequences in groundwater (Hirsch et al., 2011; Moore et al., 2011; Sonthiphand and Neufeld, 2013). Only 126 sequences were included in this analysis; however, 472 anammox sequences were collected from Genbank (Table 1). The majority of groundwater anammox sequences were from contaminated groundwater in Canada (Moore et al., 2011), and most sequences were excluded due to the region of analyzed 16S rRNA genes being outside of the region used to generate a phylogenetic tree, which was the basis of this analysis.

**PHYLOGENY AND CO-OCCURRENCE OF ANAMMOX BACTERIA**

The dominant anammox phylotypes recovered from across all isolation sources were *Ca. Scalindua* and *Ca. Brocadia*, in addition to lower abundance anammox phylotypes, including *Ca. Kuenenia*, *Ca. Anammoxoglobus*, and *Ca. Jettienia* (Figure 4A). The unknown cluster comprised of 76 OTUs; however, the average sequences per OTU were only 1.78 sequences. There was no majority of anammox sequences per OTU for the unknown cluster, reflecting that the unknown anammox clusters were likely low abundance but high diversity anammox bacteria, possibly representing part of the rare biosphere of these isolation sources.

Approximately 70% of total *Ca. Scalindua* OTU sequences were from saline-related environments, including marine sediment, marine water column, estuary, and mangrove sediment (Figure 4B). *Ca. Scalindua* was also detectable in soil and freshwater-related environments, representing 13 and 8% of all anammox OTUs from those isolation sources, respectively.

*Ca. Brocadia* was most commonly retrieved from non-saline environments, including freshwater sediment, freshwater, groundwater, and soil (Figure 4B). All freshwater-related environments and soil accounted for 38 and 24% of *Ca. Brocadia* OTU sequences, respectively. Engineered ecosystems, including WWTP and reactor, accounted for 15% of *Ca. Brocadia* OTU sequences. Although 16% of *Ca. Brocadia* OTU sequences were recovered from estuary isolation sources, only 1% of these OTUs were associated with marine sediment (Figure 4B). No *Ca. Brocadia* sequences were detected in marine water column data.

*Ca. Kuenenia* was the third most abundant cluster found across all isolation sources (Figure 4A). This cluster was detected across nine of the main habitats, but not the marine water column (Figure 4B). *Ca. Kuenenia* was also found in all five minor habitats, including marine sponge, biofilter, fish gut, shrimp pond, and oil field. Although *Ca. Kuenenia* was present in almost all...
habitats, a few OTUs (1–3 OTUs) per habitat were discovered. This observation indicated that Ca. Kuenenia cluster was not ubiquitous, but still widespread across habitats.

The Ca. Anammoxoglobus cluster was distributed similarly to the Ca. Brocadia cluster across isolation sources. For example, soil and freshwater-related environments accounted for 32 and 28% Ca. Anammoxoglobus OTU sequences (Figure 4B), respectively (compare to 24 and 38% for Ca. Brocadia, respectively). Estuary, WWTP, and reactor equally accounted for 14% of total Ca. Anammoxoglobus OTUs. Marine sediment and marine water column samples did not contribute OTUs from the Ca. Anammoxoglobus cluster.

The lowest abundance of known anammox bacterial genera was Ca. Jettenia, which comprised only eight OTUs (Figure 4A). Although Ca. Jettenia was not commonly detected within most isolation sources, the majority of this cluster was retrieved from engineered ecosystems, including WWTPs and reactors (Figure 4B). These engineered isolation sources accounted for 51% of all recovered Ca. Jettenia OTUs. Freshwater sediment, groundwater, and soil equally accounted for 13% of total Ca. Jettenia OTUs. None of Ca. Jettenia OTUs were associated with saline-related environments (Figure 4B).

The distributions of anammox bacterial OTUs of the unknown cluster were relatively similar to those of the Ca. Scalindua cluster (Figure 4B). The majority of sequences found in this cluster was from saline environments, including marine sediment, marine water column, estuary, and mangrove sediment; they accounted for 57% of the unknown OTU sequences. Freshwater, freshwater sediment, soil, and WWTPs accounted for 12, 9, 7, and 5% of unknown OTU sequences, respectively. As with the Ca. Scalindua cluster, the unknown cluster was present across nine of the main habitats, but not found in groundwater.

Co-occurrence patterns suggested that Ca. Scalindua OTUs correlated very well with other Ca. Scalindua OTUs (Figure 5). In some cases, Ca. Scalindua was found together with Ca. Brocadia, Ca. Kuenenia, and OTUs from the additional unknown cluster. Strong co-occurrences of Ca. Scalindua with Ca. Anammoxoglobus and Ca. Jettenia were not observed. Ca. Brocadia OTUs within the co-occurrence network were correlated with OTUs spanning all known genera and the unknown anammox cluster (Figure 5). Ca. Anammoxoglobus correlated consistently with Ca. Brocadia, indicating a close relationship between OTUs of these two genera. Although eight OTUs of Ca. Jettenia were reported (Figure 4A), singleton OTUs were removed from this network analysis. Only one main Ca. Jettenia OTU formed part of a co-occurrence network (Figure 5). A Ca. Jettenia OTU correlated with a Ca. Anammoxoglobus OTU, and these linked to a Ca. Brocadia OTU. Overall, the resulting network revealed the close relationships among OTUs of Ca. Jettenia, Ca. Anammoxoglobus, and Ca. Brocadia clusters. The closest co-occurring genus to Ca. Kuenenia was Ca. Brocadia (Figure 5). The co-occurrence of Ca. Kuenenia with Ca. Scalindua and one OTU of the unknown cluster was also observed.

**DISCUSSION**

Based on an ordination analysis and a non-parametric analysis of the distance matrix, we confirmed that salinity is the dominant factor governing the global distribution of anammox bacteria (Figures 1A,C). These results are not surprising given that within-study correlation analyses have previously demonstrated that salinity influenced the geographical distribution of anammox bacteria in estuary and marsh sediments (Dale et al., 2009; Hu et al., 2012a; Hou et al., 2013). Ca. Scalindua dominated saline environments, including marine sediment, marine water column, estuary, and mangrove sediment. The comprehensive phylogenetic analysis also supported that ~70% of Ca. Scalindua were from saline environments (Figures 4A,B). These results are consistent with previous observations that a lab-scale bioreactor community dominated by Ca. Kuenenia shifted toward Ca. Scalindua dominance after being enriched in high salt concentrations for 360 days (Kartal et al., 2006). In addition, salinity showed negative correlations with Ca. Scalindua diversity in the Bohai Sea sediment (Dang et al., 2013), which would be consistent with the low overall diversity we observed for saline environments surveyed here (Figure 3).

Although there is no pure anammox culture available so far, comparative metagenomic studies of Ca. Kuenenia (Strous et al., 2006; Speth et al., 2012), Ca. Brocadia (Gori et al., 2011), Ca. Jettenia (Hu et al., 2012b), and Ca. Scalindua (van de Vossenberg et al., 2013; Villanueva et al., 2014) revealed that Ca. Scalindua has unique characteristics that support marine environment adaptations. Ca. Scalindua has high-affinity ammonium absorption properties, which allows it to thrive in saline environments.
transport (\textit{amtB}) and formate/nitrite transport (\textit{focA}) proteins; both genes are highly expressed compared to those present in other anammox species (van de Vossenberg \textit{et al.}, 2013). These characteristics help \textit{Ca. Scalindua} adapt to marine environments where ammonium and nitrite may be limited (Lam and Kuypers, 2011). So far, only \textit{Ca. Scalindua} is known to contain genes involved in dipeptide and oligopeptide transport with moderate expression (van de Vossenberg \textit{et al.}, 2013). Consequently, \textit{Ca. Scalindua} has an alternative ammonium source from degraded and mineralized organic matter. \textit{Ca. Scalindua} also has a relatively versatile metabolism. \textit{Ca. Scalindua} can use \textit{NO}_2^-, \textit{NO}_3^-, and metal oxides as alternative electron acceptors (van de Vossenberg \textit{et al.}, 2008, 2013). In the presence of organic acids (i.e., propionate, acetate, formate), \textit{Ca. Scalindua} can perform dissimilatory nitrate reduction to ammonia (DNRA; Jensen \textit{et al.}, 2011). Lipid assays demonstrated that ladderane lipids with three cyclobutane

![phylogenetic tree and composition](image-url)
The links between salinity and anammox bacterial distributions are also more broadly observed for other microorganisms within a broad range of habitats. For example, the abundance and diversity of ammonia oxidizing bacteria (AOB) and archaea (AOA) were affected by salinity (Francis et al., 2003; Santoro et al., 2008; Biller et al., 2012). The diversity of denitrifying bacteria in WWTP systems was affected by salinity (Yoshie et al., 2004) and inhibitory effects of salinity on nitrification and denitrification rates were observed in estuary sediment (Rysgaard et al., 1999). Not only does salinity affect the distributions of specific groups of microorganisms, salinity impacted community fingerprints and species richness estimates for Bacteria, Archaea, and Eukaryotes within a solar saltern in Spain (Casamayor et al., 2002). The bacterial community composition along an estuary shifted due to a salinity gradient (Crump et al., 2004). Statistical and multivariate approaches have also confirmed salinity as the key factor driving global distribution patterns of Bacteria (Lozupone and Knight, 2007) and Archaea (Auguet et al., 2010).
Reasons for this finding include differences in the physiological properties of anammox bacteria in natural and engineered ecosystems. This observation suggests environmental selection because of a lower affinity for ammonia and nitrite, higher tolerance to O₂, and higher growth rate (Oshiki et al., 2011) are now characterized. These physiological properties apply O₂ to facilitate AOB activity so that the coexistence of anammox bacteria and AOB transforms fixed N to N₂ gas (Third et al., 2001; van Dongen et al., 2001).

After being enriched in fluctuating nitrite concentrations, a Ca. Brocadia dominated community shifted to a Ca. Kuenenia dominated community due to differences in affinity for NO₂⁻ (van der Star et al., 2008a). Ca. Scalindua from marine environment changed to Ca. Brocadia and Ca. Kuenenia after being enriched in a bioreactor (Nakajima et al., 2008). Either Ca. Brocadia or Ca. Kuenenia was commonly dominant in lab-scale bioreactors (Egli et al., 2001; Hu et al., 2010; Park et al., 2010). In this study, network co-occurrence analysis showed that Ca. Brocadia and Ca. Kuenenia OTUs are correlated with one other (Figure 5). However, more research on physiological properties, including kinetic and biochemical analyses, of other anammox species are needed to better understand niche differentiation of anammox bacteria in different ecosystems.

Although the diversity of anammox sequences from marine water column and marine sediment was significantly different (Figure 3, Table 2), marine environments harbored a low overall diversity of anammox bacteria, mostly restricted to Ca. Scalindua (i.e., Schmid et al., 2007; Woebken et al., 2008; Hong et al., 2011a,b). A microdiversity within Ca. Scalindua was previously discovered in marine OMZs, comprising several subclusters (Woebken et al., 2008). The microdiversity of Ca. Scalindua was also found in other marine environments, including the South China Sea (Hong et al., 2011a; Han and Gu, 2013), the Jiaozhou Bay (Dang et al., 2010), the Bohai Sea (Dang et al., 2013), the Columbian Pacific (Castro-González et al., 2014), and deep-sea methane seep sediments in the Okhotsk Sea (Shao et al., 2014). The novel subclusters, Ca. Scalindua zhenghei and Ca. Scalindua pacifica, were tentatively proposed after being identified in the South China Sea (Hong et al., 2011a) and the Bohai Sea (Dang et al., 2013), respectively. Ca. Scalindua showed strong connections within its cluster but relatively low connectivity to other known anammox clusters (Figure 5). This observation reflected the microdiversity within Ca. Scalindua cluster. However, co-occurrence of Ca. Scalindua and OTUs from the unknown cluster was high and consistent, reflecting the close relationship between the two. The unknown cluster might be a second dominant cluster found in marine environments that has yet to be assigned to a genus-level designation.

In contrast to marine environments, freshwater environments showed a high diversity of anammox bacteria. The coexistence of Ca. Brocadia with known and unknown anammox clusters was generally found in previously reported freshwater habitats (Zhang et al., 2007; Hamersley et al., 2009; Hirsch et al., 2011; Yoshinaga et al., 2011; Hu et al., 2012b; Sonthiphand and Neufeld, 2013). However, one dominant anammox phylotype, Ca. Brocadia, was detected in the sediments of the Dongjiang River, Hong Kong (Sun et al., 2014), Lake Taihu, China (Wu et al., 2012), and the Grand River, Canada (Sonthiphand et al., 2013). Network analysis also showed that Ca. Brocadia clusters connected to OTUs from all known genera and the unknown cluster (Figure 5). Ca. Scalindua was solely detected in Lake Tanganyika, which is meromictic with a sharp chemocline (Schubert et al., 2006). Overall, Ca. Brocadia OTUs were found in all previously reported freshwater habitats, except Lake Tanganyika.

As with other freshwater environments, Ca. Brocadia was the major anammox phylotype detected in contaminated groundwater. However, Ca. Kuenenia, Ca. Jettenia, Ca. Scalindua, and OTUs from the unknown cluster were also present (Moore et al., 2011). However, most of sequences from this study were removed from this current analysis, resulting in low diversity richness and underestimation of anammox phylotypes in groundwater. There is insufficient groundwater-specific information due to a paucity of anammox groundwater surveys to date. We recommend further surveys of ammonia-rich groundwater isolation sources for obtaining a better understanding of anammox bacterial diversity in this important low-oxygen and N-rich habitat.

The transitional zone between freshwater and marine environments, including estuary and mangrove sediment, is a dynamic habitat. River–sea interactions (i.e., river runoff, ocean tides, and inflow/outflow) possibly enhance the diversity of anammox bacteria. The mixture of known and unknown anammox clusters was evident in estuary habitats (Dale et al., 2009; Hirsch et al., 2011; Hu et al., 2012a; Hou et al., 2013) and mangrove sediment (Han et al., 2013; Li and Gu, 2013; Wang et al., 2013).

The combination of anammox OTUs associated with Ca. Brocadia, Ca. Kuenenia, Ca. Anammoxoglobus, and Ca. Jettenia was also found in various soil types, including peat soil (Hu et al., 2011), fertilized paddy soil (Zhu et al., 2011), a flooded paddy soil (Hu et al., 2013), and an agricultural soil (Shen et al., 2013). However, a single anammox phylotype was reported in some other soil types. Ca. Jettenia was recovered from manure pond soil (Sher et al., 2012) and permafrost soil (Humbert et al., 2010). Ca. Kuenenia was also detected in rhizosphere soil (Humbert et al., 2010). Interestingly, a rice paddy soil was dominated by Ca. Scalindua (Wang and Gu, 2013). The difference in soil properties (i.e., nutrients, O₂, and pH) and depth reflected a microniche of anammox bacteria within terrestrial habitats (Zhu et al., 2011; Sher et al., 2012).

Our findings revealed the global distributions and diversities of anammox bacteria. These results added to previous knowledge about the geographical distributions and abundances of anammox bacteria in various environments, including marine (Dang...
et al., 2013; Shao et al., 2014), estuary (Hu et al., 2012a; Hou et al., 2013), soil (Sher et al., 2012), and freshwater (Sonthiphand and Neufeld, 2013; Sun et al., 2014). The abundances of anammox bacteria in marine sediments were positively correlated with marine water depth (Jaeschke et al., 2010; Sokoll et al., 2012; Trimmer et al., 2013; Shao et al., 2014). Low temperature likely favored the abundance of anammox bacteria in marine sediments (Russ et al., 2013). Consequently, anammox bacteria likely play a key role in the deep sea, where temperature is usually low (Jaeschke et al., 2010; Shao et al., 2014). In contrast to marine environments, the abundance of anammox bacteria showed a negative correlation with soil depth (Sher et al., 2012). The suggested reason for higher anammox bacterial abundance in surface soils was higher nutrient availability in upper layers compared to bottom layers of the soil profile. Substrate availability (NO$_3^-$ and NH$_4^+$) influenced anammox bacterial abundance in marine (Dang et al., 2010), estuary (Hou et al., 2013), freshwater (Wu et al., 2012; Sun et al., 2014), and soil (Shen et al., 2013) environments. Because NO$_3^-$ can be generated from NO$_2^-$ reduction, NO$_3^-$ concentration also affected the anammox bacterial abundance in estuary sediments (Hu et al., 2012a) and marine sediments (Han and Gu, 2013). In addition to quantifying the abundance of anammox bacteria, their activity must still be assessed in many of the above habitats to better understand their contributions to ecosystem N loss as part of the global N cycle.

**CONCLUDING REMARKS**

The global distribution pattern of anammox bacteria is controlled primarily by salinity. Distinct partitioning of anammox bacterial communities among natural and engineered ecosystems was also observed in our sequence survey. Insufficient information on anammox genomes and physiological properties is available to draw conclusions on how extrinsic factors (i.e., salinity, NH$_4^+$, NO$_2^-$) affect possible anammox bacterial mechanisms. More additional metagenomic studies of other anammox species will help compare and contrast the specific genes and their functions that influence the distribution and co-occurrence of anammox bacteria. Further investigations on kinetic and biochemical properties of more anammox species are needed to better understand the ecological niche partitioning of anammox bacteria. Freshwater is a promising habitat in which to discover novel anammox species and groundwater, in particular, may be an ideal study habitat for discovering anammox bacterial contributions to N loss in freshwater-related environments. Multidisciplinary approaches, including both metagenomic studies and molecular anammox surveys, are needed to fill in missing knowledge gaps.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://www.frontiersin.org/journal/10.3389/fmicb.2014.00399/abstract

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