Community Composition of Nitrous Oxide Consuming Bacteria in the Oxygen Minimum Zone of the Eastern Tropical South Pacific

Xin Sun*, Amal Jayakumar and Bess B. Ward

Department of Geosciences, Princeton University, Princeton, NJ, United States

The ozone-depleting and greenhouse gas, nitrous oxide (N\textsubscript{2}O), is mainly consumed by the microbially mediated anaerobic process, denitrification. N\textsubscript{2}O consumption is the last step in canonical denitrification, and is also the least O\textsubscript{2} tolerant step. Community composition of total and active N\textsubscript{2}O consuming bacteria was analyzed based on total (DNA) and transcriptionally active (RNA) nitrous oxide reductase (nosZ) genes using a functional gene microarray. The total and active nosZ communities were dominated by a limited number of nosZ archetypes, affiliated with bacteria from marine, soil and marsh environments. In addition to nosZ genes related to those of known marine denitrifiers, atypical nosZ genes, related to those of soil bacteria that do not possess a complete denitrification pathway, were also detected, especially in surface waters. The community composition of the total nosZ assemblage was significantly different from the active assemblage. The community composition of the total nosZ assemblage was significantly different between coastal and off-shore stations. The low oxygen assemblages from both stations were similar to each other, while the higher oxygen assemblages were more variable. Community composition of the active nosZ assemblage was also significantly different between stations, and varied with N\textsubscript{2}O concentration but not O\textsubscript{2}. Notably, nosZ assemblages were not only present but also active in oxygenated seawater: the abundance of total and active nosZ bacteria from oxygenated surface water (indicated by nosZ gene copy number) was similar to or even larger than in anoxic waters, implying the potential for N\textsubscript{2}O consumption even in the oxygenated surface water.

Keywords: N\textsubscript{2}O consuming bacteria, nosZ gene, microarray, oxygen minimum zone, Eastern Tropical South Pacific

INTRODUCTION

N\textsubscript{2}O is a major ozone-depleting substance and a greenhouse gas whose radiative forcing per mole is 298 times that of carbon dioxide (IPCC, 2007; Ravishankara et al., 2009). Oxygen minimum zones (OMZs) are the most intense marine sources of N\textsubscript{2}O and are hot spots of rapid N\textsubscript{2}O cycling (Martinez-Rey et al., 2015). OMZs are marine regions with a strong O\textsubscript{2} gradient (oxycline) overlying an oxygen deficient zone (ODZ) where O\textsubscript{2} concentration is low enough to induce anaerobic processes. The global expansion and intensification of OMZs, which are
predicted to result from global warming, further stress the importance of understanding N\textsubscript{2}O cycling in these regions (Codispoti, 2010). N\textsubscript{2}O production and consumption are driven by marine bacteria (Naqvi et al., 2000). The dominant microbial process for N\textsubscript{2}O cycling is denitrification, the sequential reduction of NO\textsubscript{3} to NO\textsubscript{2}, NO, N\textsubscript{2}O and finally to N\textsubscript{2} (Zumft, 1997; Naqvi et al., 2000). Denitrification could stop at an intermediate step before N\textsubscript{2} if O\textsubscript{2} concentration exceeds the threshold for the latter step or if electron donors are depleted (Ward et al., 2008; Dalsgaard et al., 2012; Babbin et al., 2014). Thus O\textsubscript{2} concentration or electron donor availability could also control the N\textsubscript{2}O budget. N\textsubscript{2}O concentrations and net N\textsubscript{2}O production rates (N\textsubscript{2}O production minus N\textsubscript{2}O consumption) were found to peak at the oxic-suboxic interface in OMZs, due to excess production from nitrification and incomplete denitrification (Nicholls et al., 2007; Ji et al., 2015; Trimmer et al., 2016). However, while multiple processes can produce N\textsubscript{2}O, reduction by N\textsubscript{2}O consuming bacteria is the only known biological N\textsubscript{2}O sink.

N\textsubscript{2}O consumption is the final step of denitrification, and is the least O\textsubscript{2} tolerant step (Bonin et al., 1989; Körner and Zumft, 1989). N\textsubscript{2}O consumption rates have been measured in ODZs at depths where O\textsubscript{2} concentration ranged from very low to below the detection limit (Wyman et al., 2013; Babbin et al., 2015). N\textsubscript{2}O consumption by denitrification and genes involved in N\textsubscript{2}O reduction have also been detected in oxygenated seawater (Farias et al., 2009; Wyman et al., 2013). Characterizing the distribution and environmental regulation of this step is necessary for a complete quantification of the oceanic N\textsubscript{2}O budget and will improve our ability to predict oceanic N\textsubscript{2}O emissions under global climate change.

N\textsubscript{2}O consumption is catalyzed by the enzyme nitrous oxide reductase, encoded by the nosZ gene. A recent study of nosZ genes found a lower diversity of nosZ genes in ODZ waters than in the upper oxycline of the OMZ (Castro-González et al., 2015). The distribution of nosZ genes was related to O\textsubscript{2} concentration, which suggested that the quantity and composition of nosZ genes and the diversity of denitrifying bacteria might influence the microbial potential for N\textsubscript{2}O consumption.

We aimed to determine the distribution and community composition of total and transcriptionally active (abbreviated as ‘active’ hereafter) nosZ assemblages based on the presence (DNA) and expression (RNA) of nosZ genes in the OMZ of the Eastern Tropical South Pacific (ETSP), one of the three major OMZs in the world ocean. Three hypotheses were tested in the study: (1) the community compositions of total and active nosZ assemblages differ between coastal and off-shore stations because the quantity and quality of the nutrients at the two stations differ due to different contributions from land and sediment; (2) quantities and composition of total and active nosZ assemblages are related to O\textsubscript{2} concentration because N\textsubscript{2}O consumption is the least O\textsubscript{2} tolerant step in conventional denitrification; and (3) the distribution of the active nosZ assemblage is more related to N\textsubscript{2}O concentration than that of total nosZ assemblage because the former indicates live and active organisms.

**MATERIALS AND METHODS**

**Experimental Sites and Sampling**

Samples were collected on the R/V Nathaniel B. Palmer during June to July 2013 (cruise NBP 1305) in the OMZ of the ETSP at the off-shore station (BB1; 14.0°S, 81.2°W) and the coastal station (BB2; 20.50°S, 70.70°W) (Supplementary Figure 1). Particulate material was collected in Niskin bottles mounted on the standard conductivity-temperature-depth (CTD) rosette system (Seabird Electronics, Seattle, WA, United States) at four depths at each station (BB1: 60, 130, 300, and 1000 m; BB2: 60, 115, 300, and 1000 m) and concentrated by filtration (up to 4 L) through Sterivex filters (0.22 μm). Filters were flash frozen in liquid nitrogen onboard and stored at −80°C until DNA and RNA extraction was performed.

Temperature, salinity, sigma theta, bottom depth and pressure at each station were measured on the SBE 911+ CTD system. Fluorescence, representing chlorophyll a, was measured using a single channel fluorometer (Wet labs, Philomath, OR, United States) mounted on the CTD. Oxygen distributions were determined using the STOX sensor (detection limit = 10 nM) mounted on the CTD rosette (Revsebech et al., 2009). Ammonium, nitrite and nitrate concentrations were measured using standard colorimetric protocols (UNESCO, 1994). N\textsubscript{2}O concentration was determined using mass spectrometry (Ji et al., 2015). N\textsuperscript{*} is the deviation of measured dissolved inorganic nitrogen (DIN = nitrate + nitrite + ammonia) from predicted DIN by Redfield ratio and the world-ocean nitrogen to phosphate regression relationship (Deutsch et al., 2001). Environmental data were reported by Ji et al. (2015) and are provided in Supplementary Table 1.

**DNA and RNA Extractions**

Both DNA and RNA were extracted from eight Sterivex filters using the plant tissue protocol of the All Prep DNA/RNA Mini Kit (50) using a QIAcube (Qiagen). Reverse transcription from RNA to cDNA was performed using SuperScript® III First-Strand Synthesis System for RT-PCR (Invitrogen™ by Life Technologies™). Excess RNA was removed by RNase at the end of the synthesis.

**Quantitative PCR Assays**

The abundance of total and active nosZ assemblages were estimated by quantitative PCR (qPCR) using SYBR® Green based assays using protocols described previously (Jayakumar et al., 2013). Primers nosZIF and nosZIR (Henry et al., 2006) were used to amplify a 259-bp conserved fragment of the nosZ gene. Known quantities (~20–25 ng) of DNA and cDNA samples were assayed along with a minimum of five serial dilutions of plasmids containing nosZ gene, no template controls and no primer controls, all in triplicate on the same plate. To maintain continuity and consistency among qPCR assays, a subset of samples from the first qPCR assay was run with subsequent assays and fresh standard dilutions were prepared for each assay. DNA, cDNA and the concentrated standards were quantified prior to every assay using PicoGreen fluorescence (Molecular Probes,
Eugene, OR, United States) calibrated with several dilutions of phage lambda standards, to account for DNA loss due to freeze thaw cycles. qPCR assays were run on a Stratagene MX3000P (Agilent Technologies, La Jolla, CA, United States). Automatic analysis settings were used to determine the threshold cycle (Ct) values. The copy numbers (number of copies of the gene sequence detected in the sample) were calculated according to:

\[
\text{Copy number} = (\text{ng} \times \text{number/mole})/(\text{bp} \times \text{ng/g} \times \text{g/mole of bp})
\]

and then converted to copy number per ml seawater filtered, assuming 100% extraction efficiency.

**Microarray Experiments**

DNA and cDNA qPCR products were used as targets for microarray experiments to characterize the community composition of total and active nosZ assemblages, respectively. Triplicate qPCR products from each depth were pooled. nosZ gene targets were purified and extracted from agarose gels using the QIAquick gel extraction kit (Qiagen). Purified DNA qPCR products from eight depths and cDNA qPCR products from seven depths were used to prepare targets for microarray analysis.

Microarray targets were prepared from the qPCR products following the protocol of Ward and Bouskill (2011). Briefly, dUaa was incorporated into purified DNA and cDNA during linear amplification using the BioPrime kit (Invitrogen™). The dUaa-Klenow product was labeled with Cy3 (dissolved in dimethyl sulfoxide), purified using QIAquick columns (Qiagen) and quantified by Nanodrop 2000 (Thermo Scientific). Duplicate Cy3 products for each sample were hybridized at 65°C overnight (16 h) onto replicate microarrays under ozone free conditions. Hybridized microarrays were washed and scanned with an Axon 4300 laser scanner.

**nosZ Microarray**

The microarray (BC016) contains 114 nosZ archetype probes. Each probe is a 90-bp sequence comprised of a 70-bp nosZ gene fragment and a 20-bp control region. Each archetype probe represents, and hybridizes with, all nosZ sequences with >85% identity, based on published sequences available in 2013. There are 71 NosZ archetypes, which represent typical or Clade I nosZ genes, and 43 WNZ archetypes, which represent the atypical or Clade II nosZ genes. The development of the microarray is described in Jayakumar et al. (in preparation) and the sequences are shown in Supplementary Table 2.

**Data Analyses**

Fluorescence signal intensities for nosZ probes hybridized to the microarrays were obtained using GenePix Pro 7 software. The fluorescence ratio (FR) of each feature is defined as the ratio Cy3/Cy5 (70-mer probe/20-mer standard for each feature). The FR for each nosZ archetype was calculated as the average of probe signal intensities for duplicate features on the same microarray. Normalized fluorescence ratio (FRn) was calculated by dividing the FR of each nosZ probe by the maximum nosZ FR on the same microarray. FRn is the proxy of the relative abundance of each archetype and was used for further analyses.

Detrended correspondence analysis (DCA) was performed to analyze the overall microbial community composition. A dissimilarity test was performed using Permutational Multivariate Analysis of Variance (adonis). α-diversities (Shannon diversity indices) of total and active nosZ assemblages were calculated. β-diversities (Bray–Curtis dissimilarities) of total and active nosZ assemblages between different sites (i.e., depths) were calculated to perform a Mantel test. The Mantel test was used to determine significant environmental variables correlated with microbial community composition. These analyses were carried out using the vegan package in R (version 3.3.1). A maximum likelihood phylogenetic tree was built from aligned archetype sequences with MEGA 7 software. FRn values for each archetype at different depths from both stations were visualized on the phylogenetic tree by iTOL. The copy number of nosZ genes at each depth is given as mean (± standard error) of the qPCR triplicates.

**RESULTS**

**Abundance and Depth Distribution of Total and Active nosZ Assemblages**

At stations BB1 and BB2, the continuously undetectable O2 concentration, the local nitrite maximum and the nitrate deficit at intermediate depths (130–370 m at BB1; 75–400 m at BB2) all indicated the presence of ODZs (gray areas in Figure 1). Sampling depths were chosen to represent water column features defined by oxygen concentration, as measured with the in situ STOX sensor: oxygenated surface water, upper oxycline [characterized by sharp O2 concentration gradient ranging from saturation to below detection limit (<10 nM)], top of the ODZ (O2 concentration <10 nM), core of the ODZ (O2 concentration <10 nM) and lower oxycline (O2 concentration >10 nM). The abundance of the total nosZ genes ranged from 24.1 (±1.4) copies mL⁻¹ in a sample from the lower oxycline to 63.4 (±28.3) copies mL⁻¹ in a sample from the ODZ (Figure 1). As for the active nosZ assemblage, the lowest abundance of active nosZ genes was 5.1 (±0.5) copies mL⁻¹ in a sample from the lower oxycline and the highest abundance was 604.6 (±103.7) copies mL⁻¹ in a sample from the surface water.

The abundance of total and active nosZ assemblages showed different distribution patterns at the two stations (Figure 1). At station BB1, the abundance of both total and active nosZ genes was highest in a sample from the surface water, and decreased with depth. At station BB2, the abundance of both total and active nosZ genes peaked in samples from the ODZ and was lowest in samples from the lower oxycline. The active nosZ genes were most abundant in the sample from 300 m. However, the total nosZ genes were most abundant in the sample from 115 m, where the abundance of the active nosZ genes was only 3% of the total.

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1http://itol.embl.de/
Diversity and Dominant Archetypes of Total and Active nosZ Assemblages

The distribution of FRn of the total or active archetypes was similar across all depths within the same station (Figure 2). The average α-diversity was not significantly different (student’s t-test, $P = 0.102$) between the total assemblages (3.21) and the active assemblages (2.60) (Table 1). The least diverse total assemblage was from the lower oxycline (1000 m of station BB2), but the two least diverse active assemblages were from the ODZs (130 m of BB1 and 300 m of BB2).

The FRn distribution of nosZ archetypes showed that a very limited number of archetypes dominated the total or the active nosZ assemblages (Figure 2). Dominant archetypes were affiliated with bacteria from various environments, including salt marsh, soil, marine sediment, marine hot spring and activated sludge of a wastewater treatment plant (Supplementary Tables 3, 4). The FRn of the top five most abundant archetypes accounted for 48.9 to 83.3% of the total nosZ hybridization signal (Figure 3A and Supplementary Table 3). Notably, the highest percentage (83.3%) was from the sample from the lower oxycline (1000 m) at station BB2 and the most abundant archetype (NosZ42, an uncultured clone of nosZ gene derived from salt marsh sediments; Kearns et al., 2015) accounted for 31.6% of total FRn. However, this archetype was not among the top five archetypes of the active nosZ assemblage in the same sample (Figure 3B and Supplementary Table 4). The total nosZ assemblage was much more diverse than the active assemblage at this depth (Table 1). The representative sequences of WNZ21 and WNZ16 archetypes were derived from nosZ gene sequences of Anaeromyxobacter dehalogenans strain DCP18 (Chee-Sanford et al., unpublished) and an uncultured bacterium clone obtained from agricultural soils (Sanford et al., 2012), respectively. WNZ21 and WNZ16 archetypes were not only dominant in the active assemblage in one sample from the ODZ, but were among the top five abundant archetypes of both total and active assemblages in almost all samples (Figure 3).
Community Composition of Total and Active nosZ Assemblages

Functional gene microarrays were used to describe the community composition of nosZ assemblages. FRn values from duplicate microarrays replicated well ($r^2 = 0.802$–$0.997$) (Supplementary Figure 2) and each pair of duplicates clustered together in the DCA plots (Figure 4).

The two-dimensional DCA model including both DNA and RNA microarray results explained 43.7% of the community composition of nosZ assemblages with 31.6% explained by the first axis and 12.1% explained by the second axis (Figure 4A). The clearest pattern was the clear separation of total (filled symbols) and active (open symbols) nosZ assemblages, indicating that they were different from each other. The significance ($P < 0.001$) of the difference between total and active assemblages was confirmed by the dissimilarity test (Table 2). Therefore, the community composition of total and active nosZ assemblages was further analyzed by two DCA models separately to better examine other patterns.

The DCA model of DNA microarray results explained 63.9% of the composition of the total nosZ assemblage (Figure 4B).

**TABLE 1** | α-diversities of total (DNA) and active (RNA) nosZ genes at off-shore station BB1 and coastal station BB2.

<table>
<thead>
<tr>
<th>Depth(m)</th>
<th>DNA</th>
<th>RNA</th>
<th>Depth(m)</th>
<th>DNA</th>
<th>RNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off-shore station BB1</td>
<td>60</td>
<td>3.18</td>
<td>3.29</td>
<td>130</td>
<td>3.09</td>
</tr>
<tr>
<td></td>
<td>3.57</td>
<td>3.37</td>
<td></td>
<td>300</td>
<td>3.29</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>2.10</td>
<td></td>
<td>1000</td>
<td>2.10</td>
</tr>
<tr>
<td>Coastal station BB2</td>
<td>60</td>
<td>3.33</td>
<td>3.39</td>
<td>115</td>
<td>3.56</td>
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<td>1.63</td>
<td></td>
<td>300</td>
<td>3.57</td>
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<tr>
<td></td>
<td>1000</td>
<td>2.10</td>
<td></td>
<td>1000</td>
<td>2.10</td>
</tr>
</tbody>
</table>
The total nosZ community composition revealed site difference and O$_2$ dependence. Samples from the same station clustered together in the DCA model, indicating community composition was different between the off-shore station (BB1) and the coastal station (BB2). The site difference was statistically significant ($P < 0.001$) based on the dissimilarity test (Table 2). Besides the geographical pattern, composition of the total nosZ assemblage was also affected by O$_2$ concentration (Figure 4B). Samples from the ODZs clustered together, while samples with higher O$_2$ concentrations were distinct from the ODZ samples and different from each other. O$_2$ concentration of the seawater might not be the most important driver of microbial community composition, however, since the O$_2$ pattern was not captured by either axis of the DCA model.

The DCA model of RNA microarray results explained 40.2% of the community composition of the active nosZ assemblage (Figure 4C). Significant site difference ($P = 0.039$) of the active community composition was also revealed by the DCA model (Figure 4C) and the dissimilarity test (Table 2). However, the clustering based on O$_2$ concentration that was observed in the total nosZ assemblage was not observed for the active nosZ assemblage.

**DISCUSSION**

**Abundance and Diversity of nosZ Assemblages**

Oxygen minimum zones are sites of high N$_2$O flux to the atmosphere (Law and Owens, 1990; Arévalo-Martínez et al., 2015). N$_2$O consuming organisms are the only biological sink for N$_2$O. Hence their abundance and community composition in the OMZ may be important in understanding the N$_2$O flux. The abundance of total and active N$_2$O consuming bacteria in the OMZ of the ETSP was estimated by measuring nosZ gene copy number (Figure 1). The relationship between abundance of N$_2$O consuming bacteria and depth in this study differed from that of denitrifiers indicated by nirS gene copy number at the same stations (Ji et al., 2015): the abundances of the total and active N$_2$O consuming bacteria in the surface water were similar to or higher than those in the ODZs (Figure 1), but the abundance of denitrifiers in the surface water was two orders of magnitude smaller than that in the ODZs. This difference suggests that the two genes represent functionally different groups.

nirS and nosZ also differed in their absolute abundance. The highest abundance of N$_2$O consuming bacteria was only a few hundred copies mL$^{-1}$, which was three orders of magnitude smaller than the highest abundance of denitrifiers measured at the same stations (Ji et al., 2015). It is assumed that both genes are present in the genome as single copy genes, although there are exceptions for nosZ (Sanford et al., 2012). One possible explanation for the differences in both distribution and abundance of the nosZ assemblage and the nirS assemblage is that not all N$_2$O consuming bacteria contain the complete denitrification gene sequence (Sanford et al., 2012). The atypical nosZ genes are associated with bacteria that lack the other steps in the conventional denitrification pathway. Notably, bacteria with only nosZ genes but no other denitrification genes were overrepresented in the genomes of marine bacteria compared to other ecosystems (Graf et al., 2014). nirS, however, was
preferentially associated with bacteria that contained a complete denitrification pathway (Graf et al., 2014).

Another contributing factor may be the specificity or bias of the PCR primers. The nosZ primers used in this study were optimized to amplify all known nosZ sequences as of 2006, and should therefore represent the large database of both terrestrial and marine sequences available at the time. However, it is clear that they might underrepresent the atypical N₂O consuming bacteria, which were not known at the time. The nirS primers used in the previous analysis of these samples (Ji et al., 2015) are potentially biased toward marine sequences (Braker et al., 1998) and may underrepresent more diverse sequences now available from other environments. One way to improve the nosZ coverage is to use multiple primer sets targeting different groups of nosZ archetypes.

The N₂O consuming bacteria are a small component of the total microbial assemblage, but are still quite diverse (Jones et al., 2013), so they are difficult to characterize by pure culture or metagenomics. The microarray, which was designed specifically to target N₂O consuming bacteria using more than 100 nosZ gene probes (Figure 2), may be a better tool to capture these underrepresented organisms without cultivation or detection of rare sequences in complex metagenomic datasets. The high reproducibility of microarrays reported previously (Bulow et al., 2008) was confirmed in this study in that duplicates for each sample run on two different microarrays clustered together.
in DCA (Figure 4) and had high $r^2$ of linear regressions (Supplementary Figure 2).

Based on the FRn values of diverse nosZ archetypes determined by microarray hybridization (Figure 2), a very limited number of archetypes dominated the total or the active assemblages. Moreover, the top five active archetypes accounted for larger percentage of the assemblage than that of the total archetypes (Figure 3), consistent with the less diverse active assemblage compared to the total assemblage (Table 1). These findings imply that although the total nosZ assemblage is very diverse, a few nosZ archetypes might be the major contributors of N$_2$O consumption at the study sites. The relative abundance of active archetypes, however, might uncouple that of enzymes of different archetypes and/or the contribution of different archetypes to the N$_2$O consumption rate due to different stabilities of enzymes from different archetypes.

### Total and Active Community Compositions and Their Controlling Environmental Variables

The active nosZ assemblage was different from the total assemblage in both abundance profiles and community composition as detected by qPCR and nosZ microarray hybridization analysis, respectively. The highest abundance of the total N$_2$O consuming bacteria indicated by nosZ DNA copy number was 636.4 (±28.3) copies mL$^{-1}$ in the sample from the ODZ (115 m) at station BB2, but the abundance of the active bacteria in the same sample was only 21.1 (±5.2) copies mL$^{-1}$ (Figure 1). In the sample from oxygenated surface seawater (60 m) at station BB1, the abundance of the active N$_2$O consuming bacteria was 604.6 (±103.7) copies mL$^{-1}$, but the abundance of the total bacteria was only 357.2 (±12.5) copies mL$^{-1}$ (Figure 1).

### nosZ Assemblage in Oxygenated Seawater

The role of the nosZ assemblage in oxygenated seawater has been ignored because N$_2$O consumption is considered the least oxygen tolerant anaerobic step in the conventional denitrification pathway (Zumft, 1997). However, nosZ genes were abundant in oxygenated surface water in the Southern Indian Ocean (Raes et al., 2016) and nosZ mRNAs were detected in the

| Table 3 | Mantel tests between total (DNA) or active (RNA) nosZ genes and environmental factors. |
|---------|-----------------------------------|-----------------------------------|---------|-----------------------------------|-----------------------------------|
|         | **Total nosZ**                    | **Active nosZ**                   |         | **Total nosZ**                    | **Active nosZ**                   |
|         | $r$ | $P$ | $r$ | $P$ |
| Oxygen  | 0.169 | 0.190 | -0.046 | 0.636 |
| Relative Depth$^1$ | 0.848 | 0.001 | -0.170 | 0.886 |
| Nitrate | 0.355 | 0.009 | -0.149 | 0.913 |
| Nitrite | 0.090 | 0.178 | 0.300 | 0.019 |
| Nitrous Oxide | -0.089 | 0.684 | 0.415 | 0.014 |
| Temperature | 0.483 | 0.002 | -0.153 | 0.937 |
| Ammonium | 0.155 | 0.234 | -0.008 | 0.458 |
| Salinity | 0.258 | 0.069 | -0.165 | 0.941 |
| Sigma Theta$^2$ | 0.439 | 0.007 | -0.088 | 0.774 |
| Bottom Depth | 0.320 | 0.001 | 0.256 | 0.012 |
| Pressure | 0.492 | 0.003 | -0.188 | 0.949 |
| Fluorescence | 0.059 | 0.290 | 0.320 | 0.015 |

$^1$Relative depth was calculated by dividing measured depth by the bottom depth of each station. $^2$Sigma Theta was density calculated with in situ salinity and potential temperature at zero pressure. Bolded $P$ values indicate significant correlation ($P < 0.05$).
oxic regions in the Arabian Sea (Wyman et al., 2013). Our study confirmed that a nosZ assemblage was not only present but also active in oxygenated surface water in the OMZ of the ETSP. In particular, atypical nosZ archetypes, usually associated with N$_2$O consuming bacteria lacking a complete denitrification pathway, were present and active in surface waters. In addition, the most abundant archetypes of total and active nosZ communities were both atypical nosZ archetypes (WNZ21 and WNZ16), implying the significant contribution of atypical archetypes to the nosZ communities and the necessity to consider atypical archetypes while analyzing the potential of N$_2$O consumption.

N$_2$O reductase enzymes from denitrifiers had very low O$_2$ tolerance (Bonin et al., 1989; Körner and Zumft, 1989); on the contrary, nosZ assemblages were detected in the oxygenated surface waters and O$_2$ concentration was not significantly correlated with the active microbial community, as indicated by DCA and Mantel test. The survival of N$_2$O consuming bacteria in oxic layers and their O$_2$-independence might be attributed to anoxic micro-environments created by phytoplankton microaggregates or particles. Free-living and particle-associated microbes from the same seawater sample can have different community compositions (Delong et al., 1993). More specifically, a recent study in the OMZ of the ETSP showed that nosZ mRNAs were 28-fold more abundant on particles (>1.6 µm) compared to free-living microbes (0.2–1.6 µm) (Ganesh et al., 2015). Additionally, nosZ mRNA co-occurred with the cyanobacterium Trichodesmium in oxic water in the Arabian Sea (Wyman et al., 2013). Consistently, fluorescence, a proxy for chlorophyll a, was significantly correlated with the β-diversity of the active nosZ assemblages in this study (Table 3).

The active nosZ community in oxygenated surface water might capture N$_2$O produced in deeper seawater and thus reduce the flux into the atmosphere. Thus, evaluating the nosZ community is essential to the prediction of the oceanic N$_2$O emissions. Moreover, the oceanic N$_2$O emissions represent net fluxes, which are controlled by both N$_2$O production and N$_2$O consumption. Some N$_2$O flux models (Suntharalingam and Sarmiento, 2000; Martinez-Rey et al., 2015; Trimmer et al., 2016) do not parameterize N$_2$O consumption, and other models either consider N$_2$O consumption only in suboxic or anoxic waters (Cornejo and Farias, 2012; Babbin et al., 2015) or estimate N$_2$O consumption assuming it is constrained by O$_2$ concentration (Zamora et al., 2012). Failing to consider the O$_2$-independent, non-denitrification N$_2$O consumption potential in these O$_2$ forcing models might contribute to their uncertainty and the variation among different models. Additionally, the N$_2$O consuming organisms have not been fully investigated. Besides denitrifiers and atypical N$_2$O consuming bacteria analyzed in this study, other organisms (Trichodesmium and Crocosphaera) also exhibited N$_2$O consuming capacity under laboratory conditions (Farias et al., 2013), suggesting that their significance in the environment warrants further investigation.

**CONCLUSION**

The results described above support two (1 and 3) of the initial hypotheses. (1) Compositions of total and active nosZ assemblages were different between the coastal station and the off-shore station mainly due to their dramatic differences of distance to the sediment and to the shore, which are very likely to result in different environmental conditions (i.e., different phytoplankton assemblages, different nutrients and organic matter). (2) The abundances of total and active nosZ assemblages in oxygenated seawater were similar to or larger than those in the ODZs, implying the potential for N$_2$O consumption even in oxygenated surface water. Atypical nosZ archetypes, which may lack a complete denitrification pathway, dominated both total and active nosZ assemblages. (3) The total and active nosZ assemblages were significantly different from each other. The community composition of the total nosZ assemblage showed O$_2$ dependence and shifted along depth gradients and environmental gradients associated with depth, but fluorescence, N$_2$O and nitrite concentration were significantly correlated with the composition of the transcriptionally active community. We conclude that the difference between active and total nosZ assemblages may be related to differential response to environmental conditions by different components of the diverse natural assemblage and that the presence of nosZ assemblage in surface waters should be investigated to determine their actual N$_2$O reduction capabilities.

**AUTHOR CONTRIBUTIONS**

XS and BW designed the experiments. AJ and BW collected samples. XS and AJ performed experiments. XS analyzed the data. XS and BW wrote the paper.

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**SUPPLEMENTARY MATERIAL**

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