Abstract
The Arabian Sea oxygen minimum zone (OMZ), the largest suboxic region in the world’s oceans, is responsible for up to half of the global mesopelagic fixed nitrogen (N) loss from the ocean via denitrification and anammox. Dinitrogen (N₂) fixation is usually attributed to cyanobacteria in the surface ocean. Model prediction and physiological inhibition of N₂ fixation by oxygen, however, suggest that N₂ fixation should be enhanced near the oxygen-deficient zone (ODZ) of the Arabian Sea. N₂ fixation and cyanobacterial nifH genes (the gene encoding dinitrogenase reductase) have been reported in surface waters overlying the Arabian Sea ODZ. Here, water samples from depths above and within the Arabian Sea ODZ were examined to explore the distribution, diversity, and expression of nifH genes. In surface waters, nifH DNA and cDNA sequences related to Trichodesmium, a diazotroph known to occur and fix N₂ in the Arabian Sea, were detected. Proteobacterial nifH phylotypes (DNA but not cDNA) were also detected in surface waters. Proteobacterial nifH DNA and cDNA sequences, as well as nifH DNA and cDNA sequences related to strictly anaerobic N-fixers, were obtained from oxygen-deficient depths. This first report of nifH gene expression in subsurface low-oxygen waters suggests that there is potential for active N₂ fixation by several phylogenetically and potentially metabolically diverse microorganisms in pelagic OMZs.

Introduction
Productivity in much of the world’s ocean is limited by the availability of fixed nitrogen (N). Observations and extrapolations from models suggest either that fixed N losses via denitrification and anammox greatly exceed N inputs to the ocean via dinitrogen (N₂) fixation (Codispoti et al., 2001; Codispoti, 2007), or that these opposing processes are balanced, at least on geological timescales (Gruber & Sarmiento, 1997; Altabet et al., 2002; Deutsch et al., 2007; Altabet, 2007). Although both processes can be variable in time and space, it is hypothesized that the apparent imbalance is owing to an underestimation of N₂ fixation (Codispoti, 2007). To test this hypothesis, the relative distributions of microorganisms that fix N₂ and their activity are being investigated using observations and models.

Denitrification occurs in anoxic and suboxic areas of the world’s oceans including marine sediments, anoxic basins and fjords, and pelagic oceanic oxygen minimum zones (OMZs) (Devol, 2008). Here, the term OMZ denotes the oceanographic region, for example the Arabian Sea, where low-oxygen waters are found, while oxygen-deficient zone (ODZ) refers to the depths where oxygen concentrations are low enough to induce anaerobic metabolisms, estimated at 1–2 µM. The Arabian Sea OMZ contains the largest suboxic region in the world’s oceans and is responsible for up to half of the global mesopelagic fixed N loss (Naqvi, 2008). While low oxygen is thought to be beneficial for N₂ fixation because oxygen inhibits nitrogenase (the enzyme complex that mediates N₂ fixation), high nitrate concentrations, such as those present in ODZs, are thought to be inhibitory to N₂ fixation in these systems (Postgate, 1998). Nonetheless, even Trichodesmium, the best-known marine N-fixing cyanobacterium, fixes N₂ in the presence of dissolved inorganic nitrogen (DIN) (Mulholland et al., 1999, 2001; Holl &
Further, $N_2$ fixation has recently been documented in areas where substantial DIN concentrations were measured, including coastal waters (Grosse et al., 2010; Mulholland et al., 2012), the Pacific Ocean (Moißander et al., 2010), and the eastern tropical Atlantic Ocean (Voss et al., 2004). In addition, models suggest that $N_2$ fixation may occur in closer proximity to zones of denitrification than previously thought (Deutsch et al., 2007; Mulholland & Capone, 2009). Denitrification results in the depletion of fixed N relative to phosphorus (P), thereby lowering N : P ratios of dissolved nutrient pools, which may create conditions conducive to $N_2$ fixation (Deutsch et al., 2007; Capone & Knapp, 2007; Naqvi, 2008). Consistent with the hypothesis that $N_2$ fixation should occur in proximity to denitrification, low rates of $N_2$ fixation, as well as gene sequences of $nifH$ (the gene encoding a critical component of the nitrogenase enzyme complex) representing several proteobacterial $nifH$ phylotypes, were detected in the low-oxygen, nitrate-rich waters of coastal California (Hammersley et al., 2011) and in the OMZ region of the Eastern Tropical South Pacific (Fernandez et al., 2011).

$N_2$ fixation is likely an important process in surface waters of the Arabian Sea. Microscopic observations and measurements of $nifH$ abundance and diversity suggest that there is a diverse assemblage of diazotrophs in surface waters of the Arabian Sea that includes *Trichodesmium*, unicellular cyanobacteria, *Gammaproteobacteria*, and diatom–diazotroph assemblages (Capone et al., 1998; Mazard et al., 2004; Bird et al., 2005; Parab et al., 2006; Mulholland & Capone, 2009; Padmakumar et al., 2010). Direct measurements of $N_2$ fixation rates, however, have been reported only once for the Arabian Sea (Capone et al., 1998). To investigate the capacity for $N_2$ fixation within the Arabian Sea, we examined $nifH$ diversity and gene expression within the ODZ as well as in the overlying surface waters.

**Materials and methods**

Water samples were collected from three stations in the Arabian Sea (Fig. 1) during September and October of 2007 aboard the R/V Roger Revelle. Water column nitrate, nitrite, and oxygen concentrations at these stations were reported previously (Ward et al., 2009) and are shown here for one station in Fig. 2. Oxygen concentrations were $< 1 \mu M$ between 100 and 750 m depth at Station 1 and between 140 and 750 m depth at Station 2. The depleted oxygen layer at Station 3 was much thinner, spanning only 120–260 m in depth. Water samples (up to 12 L) were collected from 2 to 4 depths per station (Table 1): near the surface or in the chlorophyll maximum layer (upper mixed layer), within the oxycline where the oxygen gradient was steepest, and within the ODZ at the depth of the secondary nitrite maximum. Whole water from CTD-mounted Niskin bottles was filtered through Sterivex-GP capsule filters (Millipore), using a peristaltic pump, quick frozen in liquid nitrogen,
Table 1. Stations, depth, and oceanographic features from which DNA and cDNA sequences were obtained

<table>
<thead>
<tr>
<th>Station</th>
<th>Depth (m)</th>
<th>Feature</th>
<th>DNA clones analyzed</th>
<th>DNA clone ID (number of clones)</th>
<th>cDNA clones analyzed</th>
<th>cDNA clone ID (number of clones)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (19°N 67°E)</td>
<td>10</td>
<td>Euphotic zone</td>
<td>17</td>
<td>S1D010M (3)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>Oxycline</td>
<td>24</td>
<td>S1D060M (20)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>150</td>
<td>Secondary nitrite maximum (ODZ)</td>
<td>24</td>
<td>S1D150M (23)</td>
<td>47</td>
<td>mRNA S1D150M (25)</td>
</tr>
<tr>
<td></td>
<td>175</td>
<td>Secondary nitrite maximum (ODZ)</td>
<td>23</td>
<td>S1D175M (11)</td>
<td>24</td>
<td>mRNA S1D175M (22)</td>
</tr>
<tr>
<td>2 (15°N 64°E)</td>
<td>2.5</td>
<td>Euphotic zone</td>
<td>32</td>
<td>S2D2.5M (0)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>150</td>
<td>Secondary nitrite maximum (ODZ)</td>
<td>32</td>
<td>S2D150M (5)</td>
<td>28</td>
<td>mRNA S2D150M (25)</td>
</tr>
<tr>
<td>3 (12°N 64°E)</td>
<td>10</td>
<td>Euphotic zone</td>
<td>32</td>
<td>S3D010M (25)</td>
<td>20</td>
<td>mRNA S3D10M (4)</td>
</tr>
<tr>
<td></td>
<td>65</td>
<td>Chlorophyll Maximum</td>
<td>32</td>
<td>S3D065M (0)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td></td>
<td>110</td>
<td>Oxycline</td>
<td>32</td>
<td>S3D110M (5)</td>
<td>28</td>
<td>mRNA S3D110M (23)</td>
</tr>
</tbody>
</table>

Clones analyzed = total number of clones sequenced. (number of clones) = number of clones that were determined by phylogenetic analysis to represent nifH genes. NA = cDNA clone libraries were not made from these samples.

and shipped in liquid nitrogen to Princeton University (PU). The samples were stored at −80 °C before being transported to Old Dominion University (ODU) in liquid nitrogen for DNA extraction.

The DNA was extracted at ODU from Sterivex filters using the PUREGENETM Genomic DNA Isolation Kit (Qiagen, Germantown, MD). The RNA was extracted at PU from replicate Sterivex filters using the ALLPrep DNA/RNA Mini Kit (Qiagen). The cDNA was synthesized immediately following purification of RNA using a SuperScript III First Strand Synthesis System (Invitrogen, Carlsbad, CA) following the procedure described by the manufacturer. DNA and cDNA were quantified using PicoGreen dsDNA quantification kit (Invitrogen) according to the manufacturer’s specifications. The nifH sequences were amplified from environmental DNA and cDNA using Phusion™ High-Fidelity DNA Polymerase (Finnzymes), on an MJ100 Thermal Cycler (MJ Research). A nested reaction was used, as previously described (Zehr et al., 1998), with slight modification: 25 μL PCRs were amplified for 30 cycles (1 min at 98 °C, 1 min at 57 °C, 1 min at 72 °C), first with the outer PCR primers (Zani et al., 2000), followed by amplification with the inner PCR primers (Zehr and McReynolds, 1989). To minimize the possibility of amplifying contaminants (Zehr et al., 2003), negative controls (autoclaved and UV-irradiated water) were run with every PCR experiment, reagents were diluted in freshly autoclaved water, the PCR preparation station was UV irradiated for 1 h before each daily use, and the number of cycles was limited to 30 for each reaction. The PCR reagents were also irradiated where possible. Each reagent was then tested separately for amplification in negative controls.

Amplified fragments were electrophoresed on 1.2% agarose gels, and nifH bands were excised and then cleaned using a QIAGen Quick Nucleotide Removal Kit (Qiagen). Cleaned fragments were inserted into a pCR®2.1-TOPO® vector using One Shot® TOP10 Chemically Competent E. coli, TOPO TA Cloning® Kit (Invitrogen) according to manufacturer’s specification. Clones were picked randomly and amplified using M13 forward (−20) and M13 reverse primers. The products were sequenced at the DNA Analysis Facility at Yale University using Big DyeTM terminator chemistry (Applied Biosystems, Carlsbad, CA). Sequences were edited using CHIMERA ver. 1.4.0 (Geospiza Inc.) and checked for identity using BLAST. Consensus nifH sequences (359 bp) were translated to amino acid (aa) sequences (108 aa after trimming the primer region) and aligned using CLUSTALX (Thompson et al., 1997) along with published nifH sequences from the NCBI database. Neighbor-joining trees were produced from the alignment using distance matrix methods (PAUP 4.0; Sinauer Associates). Bootstrap analysis was used to estimate the reliability of phylogenetic reconstruction (1000 iterations). The nifH sequence from Methanosarcina lacustris (AAL02156) was used as an outgroup. The new Arabian Sea nifH sequences have been deposited in GenBank, DNA sequences accession numbers JF429940–JF429973 and cDNA sequences accession numbers JQ358610–JQ358707.

Results and discussion

Nitrate, nitrite, and oxygen profiles were characteristic of an open ocean OMZ, with a strong nitrate deficit and a clear secondary nitrite maximum within the depth interval where oxygen concentrations were < 1 μM (Fig. 2; Ward et al., 2009). All major nutrients were depleted in surface waters. The stable stratification of warm surface waters likely resulted in little if any communication between the euphotic zone and the depth of the secondary nitrite maximum, with the exception of the downward flux of sinking particulate material. Thus, the dramatically different chemical environments at the
sample depths imply quite different conditions for microbial growth and suggest that the diazotrophic assemblages would likely differ among depths.

A total of 92 DNA clones containing sequences identified as homologous with known nifH sequences were retrieved from seven of the nine depths from which clone libraries were made (Table 1). The cDNA clone libraries were made from five depths, and a total of 99 cDNA sequences were obtained (Table 1). Phylogenetic analysis of translated nifH DNA and cDNA fragments grouped the sequences into four of the major clusters previously used to characterize nifH phylogeny (Chien & Zinder, 1996). Cluster I, including proteobacteria and cyanobacteria, is shown in Fig. 3, and sequences from Clusters II, III, and IV are shown in Fig. 4.

Three sequences from surface waters (10 m) at Station 1 had high identity with nifH sequences from Trichodesmium spp. (Fig. 3). These sequences clustered with uncultured marine cyanobacterial sequences reported by Mazard et al. (2004) from surface waters of the Arabian Sea, as well as with sequences from surface waters in many other locations, including the North Pacific, North Atlantic, and South China Sea.

Although several clones (32 from each depth) of the PCR products obtained with the nested nifH primers from the euphotic zone at Station 2 and the chlorophyll maximum at Station 3 (Table 1) were sequenced, no nifH DNA sequences were detected among those clones. This suggests that N2-fixing cyanobacteria were not abundant in these waters. Nevertheless, we did detect nifH gene expression: four cDNA sequences from 10 m at Station 3 were almost identical to the Trichodesmium-like DNA sequence from 10 m at Station 1.

No sequences related to the recently identified ubiquitous, single-celled N2-fixing ‘Group A’ and ‘Group B’ cyanobacteria (Zehr et al., 2001, 2008) were recovered from any of the DNA or cDNA clone libraries (Fig. 3). Using PCR primers for the 16S rRNA gene designed to be specific for members of the Groups A and B N2-fixing uncultured cyanobacteria (UCYN2-fix lineage), Mazard et al. (2004) detected these relatives of Crocosphaera and Cynothecce only in the most southern and most northern regions of their N-S transect across the Arabian Sea. However, no sequences related to these groups were detected at their transect stations, which essentially coincide with Stations 1 and 2 in the current study. Mazard et al. (2004) suggested that the UCYN2-fix lineage was restricted to very warm (~ 30 °C) oligotrophic waters and that temperature was the factor that limited their distribution. Trichodesmium-like DNA sequences, for both nifH and 16S rRNA genes, were detected by Mazard et al. (2004) at the central Arabian Sea stations on their transect. Thus, although based on a small number of clones, the two studies are consistent in suggesting that Trichodesmium-like cyanobacteria are the most important N2 fixers in the open ocean surface waters of the Arabian Sea. No cyanobacteria-like nifH sequences were detected in any of the clone libraries from depths below the surface mixed layer.

Twenty-seven nifH DNA sequences from 150 and 175 m at Station 1 and three DNA sequences from 150 m at Station 2 clustered in the alpha-proteobacterial section of Cluster I (Fig. 3). These were most closely related to sequences derived from other marine environments, such as the South China Sea and the English Channel. Of particular interest is the near perfect identity of several clones from 150 and 175 m at Station 1 (S1D150M21 and S1D175M21) with a single sequence (ABW80585) previously reported from water that had been upwelled from oxygen-deficient depths in the Eastern Tropical South Pacific, an environment analogous to the one sampled here (Fernandez et al., 2011). Eighteen clones (including 16 identical sequences; S1D60M24) from Station 1 surface waters (60 m) were also identified as alpha-proteobacterial sequences, but with closest identities to Sphingomonas azotifigens (isolated from rice roots) and several uncultured sequences from terrestrial environments.

One group of five nifH DNA sequences from 150 m (Station 2) and 175 m (Station 1) was nearly identical to 25 sequences from 10 m at Station 3. All of these were closely related to sequences (ABM67093, associated with aerobic alpha-proteobacteria) retrieved from mangrove sediments (Zhang et al., 2008). No alpha-proteobacterial cDNA sequences were obtained from any of the samples.

Six DNA clones, including representatives from all three stations from oxycline and secondary nitrite maximum depths, were identified as likely beta-proteobacterial nifH sequences, with closest identities to cultivated symbiotic or soil bacteria (e.g. Burkholderia phytophylum, Sinorhizobium meliloti, Cupriavidus spp., and Methylocycnis rosae). This cluster included one sequence retrieved as mRNA from the Mediterranean Sea (Man-Aharonovich et al., 2007) that was most closely related to Burkholderia xenovorans (Chain et al., 2006). This sequence was thought to represent an active clade of diverse N2 fixers in the open ocean region of the Mediterranean Sea, where N limitation in ultra-oligotrophic surface waters is thought to control productivity and microbial community composition. By contrast, no beta-proteobacterial cDNA sequences were obtained from any of the samples from the Arabian Sea.

Four DNA sequences, two from 110 m at Station 3 and two from 175 m at Station 1, were identified as likely gamma-proteobacterial nifH fragments. These had closest identities to sequences retrieved from aquatic marine
environments, including tropical and subtropical surface waters and the English Channel. Fifteen identical cDNA sequences (mRNA S1D175M-A1) were also identical to the DNA sequences from the ODZ at Station 1, suggesting that this gamma-proteobacterial clade is indeed active in N$_2$ fixation in the ODZ.
The gamma-proteobacterial subcluster also contains sequences obtained from zooplankton guts (Braun et al., 1999) and from soils and marine sediments. Several sequences in this cluster were identified as belonging to heterotrophic bacterial symbionts of dinoflagellates in surface waters of the southern Indian Ocean (Farnelid et al., 2010). The largest group of expressed gamma-proteobacterial \( nifH \) genes that was detected, however, was not closely related to any of the published DNA sequences (Fig. 3). These cDNA sequences (73 in total) comprised a separate subcluster and were retrieved from all three stations at both oxycline and ODZ depths.

A cluster of anomalous cDNA sequences (top of tree, Fig. 3) were clearly associated with Cluster I, but did not fall within the cyanobacteria or any specific proteobacterial group. The translated sequences were unambiguously recognized as \( nifH \) sequences, but their affiliation is unknown. They were not associated with DNA sequences from this study but were identical to DNA sequences from unpublished studies, retrieved from Chesapeake Bay water column (accession no. AAZ06675) and marine microbial mats (accession no. ADD26204) from sandy intertidal beach on a Dutch barrier island.

Only six of the \( nifH \) DNA sequences reported here were identified phylogenetically as falling outside of Cluster I (Fig. 4). Three DNA sequences from 175 m at Station 1 grouped in Cluster III with heterotrophic bacterial genera typical of anaerobic environments, such as...
sulfate reducers and free living Spirochetes from freshwater sediments (Lilburn et al., 2001). A group of four cDNA sequences from the ODZ at Station 2 grouped into Cluster III, but were not identical to any of the DNA sequences. Cluster III nifH sequences typically derive from anaerobic gut symbionts and anoxic marine sediments (Chien & Zinder, 1996). Although the ODZ is not a sulfidic environment, the total absence of oxygen may be conducive to both free living and symbiotic N₂ fixers, at least some of which were expressing the nifH gene when sampled. Two DNA sequences from 60 m at Station 1 and one sequence from 150 m at Station 2 were only distantly related to other nifH sequences, but were identified with strong bootstrap support within Cluster IV. These sequences had greatest identity at the protein level (although still < 70% identity) with a sequence previously reported as a contaminant in PCR reagents from an unpublished study investigating nifH genes in surface ocean waters (accession no. ADI24166).

Proteobacterial nifH sequences have been reported as contaminants of PCR reagents (Zehr et al., 2003), and such contaminants probably occur much more frequently than reported. It is therefore important to consider whether the Cluster IV and proteobacterial nifH sequences reported here do indeed represent native N₂-fixing microorganisms in the Arabian Sea, or whether they might be due to contamination of reagents. The possibility of contamination was minimized by rigorous inclusion of negative controls in the PCR analysis and a relatively low number of amplification cycles (as recommended by Zehr et al., 2003), but even failure to amplify negative controls does not guarantee that sequences obtained in samples were derived from the sample alone. Amplification of trace contaminants may be enhanced in the presence of sample DNA, such that greater sensitivity is obtained in the sample than in the negative control reactions. Use of 30 cycles was intended to minimize this problem without compromising sensitivity. The proteobacterial sequences reported here are not related to others previously reported as contaminants, but this is no guarantee because almost any microbe could be present as a trace contaminant. We suggest that the sequences affiliated with sequences derived from known anaerobic species and environments may be less likely to derive from contaminants, as the reagents were not anoxic and full oxygenation of the reaction conditions likely prevailed prior to analysis.

It would be most compelling in support of active N₂ fixation if the same sequences were detected in both DNA and RNA. Some examples of identical DNA and cDNA sequences were found, but more often, the cDNA sequences were not very closely related to DNA sequences from the same samples. Detection of cDNA nifH sequences implies nifH gene expression in the cyanobacteria (Trichodesmium like), the Gammaproteobacteria, and among the sequences associated with strict anaerobes in Cluster III. The dominant cDNA clade in the Gammaproteobacteria contained numerous sequences from all three stations at oxycline and ODZ depths. While these sequences were not closely related to DNA sequences obtained from the same samples, the fact that they were retrieved from multiple depths, excluding the surface waters, suggests that they are characteristic of the subsurface low-oxygen waters of the Arabian Sea. That these sequences were not detected in the DNA implies that cells containing them were not abundant relative to other potential diazotrophs. Their expression, however, was apparently favored by the conditions in the ODZ at the time of sampling. Such disparity between diversity and abundance of sequences in DNA vs. cDNA clone libraries is not unusual (Short and Zehr, 2007), and even PCR and qPCR analyses of the same sample can yield quite different abundances of nifH sequences in clone libraries (Turk et al., 2011). Thus, PCR bias could explain the differential representation in our DNA and cDNA libraries. Other explanations, for example dependent upon transient responses to environmental conditions, are possible and could be tested with further experimental work.

We conclude from this discussion of the contamination issue that the nifH DNA sequences reported here in the gamma-proteobacterial and alpha-proteobacterial regions of Cluster I likely represent microorganisms present at the depths of the oxycline and secondary nitrite maximum of the ODZ, based on their affinity with many other sequences from aquatic marine environments. Although we did not detect gene expression in the alpha-proteobacterial clade, the DNA sequences were abundant and phylogenetically consistent with nifH detection from marine environments. The small number of sequences representing beta-proteobacterial nifH sequences is somewhat more suspect, given that they are most closely related to sequences previously reported from soil and wastewater treatment sites. Nonetheless, several sequences related to Burkholderia sp. were reported as dinoflagellate symbionts (Farnelid et al., 2010) and identified as actively expressed in surface waters of the Mediterranean (Man-Aharonovich et al., 2007), suggesting that members of this cluster are indeed natural inhabitants of the marine environment. Further exploration of the activity and regulation of these nifH genes are required to determine whether they contribute to N₂ fixation in the ODZ.

Direct measurements of N₂ fixation by Trichodesmium (Capone et al., 1998; Mulholland & Capone, 2009) and indirect evidence from geochemical tracers and models (Deutsch et al., 2007) all indicate that N₂ fixation occurs in surface waters of the Arabian Sea. Isotope mass
balances derived from stable isotope measurements of nitrate in the surface waters of the Arabian Sea indicate that a significant fraction of primary production depends on N2 fixation (Brandes et al., 1998). The occurrence of N2 fixation in surface waters of the Arabian Sea is supported by the detection of both 16S rRNA and nifH genes representing N2-fixing cyanobacteria (Mazard et al., 2004; Bird et al., 2005), as well as the detection of Trichodesmium-like nifH DNA and cDNA sequences in the present study. A recent geochemical model suggested that denitrification and N2 fixation are closely coupled in space and time in large regions of the ocean (Deutsch et al., 2007), including in the Arabian Sea (Mulholland & Capone, 2009). Thus, evidence of N2 fixation in both surface and subsurface waters presented here is consistent with recent models of N budgets in this region and with recent findings in the Eastern Tropical South Pacific (Fernandez et al., 2011). The nifH sequences reported here are the first found in the ODZ of the Arabian Sea and suggest that the genetic potential for N2 fixation occurs at the same sites where fixed N loss occurs by denitrification and anammox.

The nifH sequences retrieved from three stations in the open ocean Arabian Sea suggest that a diverse diazotrophic community including Alpha-, Beta-, and Gamma-proteobacteria is present in the ODZ. Detection of numerous cDNA sequences from these stations implies that at least some of the nifH containing organisms are active in N2 fixation. Differences in identity between DNA and cDNA may be due to clone library biases or transient environmental conditions that induced expression by low abundance members of the assemblage. While the presence of nifH genes in the ODZ suggests the capacity for N2 fixation within and adjacent to oxygen-deficient waters, active N2 fixation has not yet been measured in subsurface waters or within the ODZ. Molecular studies examining the distribution and abundance of nifH genes have recently shown the presence of nifH genes in a wide range of environments not previously thought to harbor N2 fixers: archaeal N2 fixers near hydrothermal vent systems (Mehta et al., 2005; Mehta & Baross, 2006), heterotrophic bacteria associated with flagellates in surface waters (Farnelid et al., 2010), and cyanobacteria in nitrate replete surface waters (Voss et al., 2004; Grosse et al., 2010; Moisander et al., 2010). The diversity of diazotrophs and their geographical range is clearly broader than previously appreciated.

Through their opposing impacts on regional nitrogen cycling, two aspects of the Arabian Sea underscore this region’s biogeochemical complexity. The first is the extensive mid-depth ODZ within which denitrification and anammox occur, comprising a major loss term for fixed N in the global fixed N budget. Second, the demonstrated presence of Trichodesmium spp. and other diazotrophs in surface and subsurface waters suggests that N2 fixation may counter the ODZ-related losses of fixed N, at least to some degree (Mulholland & Capone, 2009). Geochemical evidence and rate estimates suggest that rates of N2 fixation in the Arabian Sea could be high (Somasundar et al., 1990; Gruber & Sarmiento, 1997; Brandes et al., 1998; Capone et al., 1998; Bange et al., 2000; Deutsch et al., 2007). Models estimate the annual input of new N via N2 fixation to be 3.3 Tg N year−1 in the Arabian Sea (Bange et al., 2000, 2005). New evidence for the presence of diazotrophic organisms in oxygen-deficient waters supports the need for direct measurement of N2 fixation rates, as well as gene expression, to assess the net contribution of this process to the Arabian Sea’s nitrogen and carbon budgets.

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References


