Bacterial Community Profiling of the Arabian Sea Oxygen Minimum Zone Sediments using Cultivation Independent Approach

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Abstract
The eastern Arabian Sea has a unique and permanent oxygen minimum zone (OMZ) that extends along the western continental margin of India. In order to understand the bacterial community structure and diversity of OMZ sediment of the Arabian Sea (AS), PCR-DGGE analysis were carried out for samples collected off Ratnagiri, Goa and Karwar at 50m, 200m, 500m and 1000m depths. PCR-DGGE analysis showed that Proteobacteria dominated all the sites ie 50m (non-OMZ) and 200m, 500m and 1000m (OMZ) along the depth gradient. Majority of the bands were attributed to the Proteobacteria. The other groups obtained belonged to Nitrospira, Bacteriodetes, Actinobacteria, Chloroflexi and Acidobacteria. The species diversity index was higher in the OMZ sites compared to the non-OMZ sites.

Keywords: Arabian sea; OMZ; Bacterial diversity; DGGE

Introduction
Oxygen minimum zones are common and permanent features in the most productive regions of all oceans where dissolved oxygen concentrations are as low as 0.5mL L−1, present at different water depths ranging from shelf to upper bathyal zones (10-1300m). Oxygen depletion has become widespread in the world oceans occurring as permanent, seasonal and episodic features [1]. In the present scenario of the dramatic global climate change, the likelihood of the expansion of OMZs, thereby acidifying the oceans may not be unanticipated, thus posing destructive effects to oceanic ecosystems [2,3]. Of the total OMZ area 59 % occurs in Indian Ocean i.e. Arabian Sea and Bay of Bengal [4]. They are associated with the continental margins where burial of organic carbon takes place in the sediments and contribute to the global carbon cycle [5]. The oxygen depleted bottom water affects the characteristics of the sediment pore water as they are in close association [6]. Thus the region where oxygen minimum zone impinges on the sea floor creates a strong gradient in dissolved oxygen concentrations and organic carbon thus serving as a peculiar habitat for the organisms. These gradients are known to strongly influence the composition and structure of prokaryotic assemblages [7]. Culture-dependent methods are known to be inadequate for the analysis of microbial communities because only a small fraction of bacteria in the environmental samples are culturable [8]. The advent of molecular methods based on 16S rRNA genes has now been widely used for providing new insights into microbial diversity and into structure and dynamics of microbial communities. PCR-DGGE of 16S rRNA gene fragments has been found suitable to analyze multiple samples simultaneously and to assess temporal and spatial dynamics of microbial community [9]. In the present study an attempt has been made to understand the spatial variation in bacterial diversity associated with sediments along a depth gradient in the OMZ in three latitudes such as off Ratnagiri, off Goa and off Karwar. This may provide important insights into the composition and variability of bacterial communities in these regions and processes occurring therein.

Methods
Collection of sediments
Sediment samples were collected from the Arabian Sea (AS) off Ratnagiri, Goa, and Karwar at 50m, 200m, 500m and 1000m depths, onboard FORV Sagar Sambad #254. The stations at the latter three sites (200m, 500m and 1000m) fall in the OMZ while the one at 50m depth was a control site which was a coastal station. The samples were handled aseptically and preserved in liquid nitrogen until DNA extraction.
DNA extraction from sediments

DNA was extracted using modification of earlier method [10]. The purified DNA was estimated using Nanodrop spectrophotometer (NanoDrop Technologies, Inc, USA).

PCR-DGGE analysis

Table 1: Coordinates of the sampling stations.

| Transects   | Stations | Depth (m) | Latitude   | Longitude   |
|-------------|----------|-----------|------------|-------------|
| Karwar      | K1       | 50        | 14°32'26.7"N | 73°5'44.7"E |
|             | K2       | 200       | 14°32'14.5"N | 73°11'28.3"E |
|             | K3       | 500       | 14°32'148"N  | 73°07" E     |
| Goa         | K4       | 1000      | 14°32'223"N  | 73°9'30.42"E |
|             | G1       | 50        | 15°26'088"E  | 73°29'131"E  |
|             | G2       | 200       | 15°25'988"E  | 72°52'702"E  |
|             | G3       | 500       | 15°25'465"E  | 72°47'215"E  |
|             | G4       | 1000      | 15°25'529"N  | 72°40'877"E  |
| Ratnagiri   | R1       | 50        | 16°17'491"N  | 73°07'436"E  |
|             | R2       | 200       | 16°18'133"N  | 72°21'941"E  |
|             | R3       | 500       | 16°18'201"N  | 72°19'140"E  |
|             | R4       | 1000      | 16°16'25"N   | 72°16'64"E   |

16S rRNA gene amplification of the purified DNA from all the sites was performed using the primers, 341F-CCT AGC GGA GCC AGC AG (59bp GC clamp attached) and 907R-ATT ACC GCG GCC AGC AG (GC clamp attached) and 907R-ATT ACC GCG GCC AGC AG respectively [9]. The reaction was carried out using the touchdown protocol-5min at 94 °C, 15 cycles with 1 cycle consisting of 40sec at 94 °C, decreasing of the annealing temperature from 65-55 °C for 40 sec and extension of 72 °C for 40sec. In addition another 20 cycles (1 cycle consisting of 40sec at 94 °C, 40sec at 50 °C and 1min 30sec at 72 °C) and a final extension at 72 °C for 5min. The PCR amplification was performed in a Thermal cycler (Eppendorf, Germany). The amplicons were inspected on 1.5% agarose gel and visualized using the gel documentation system (Eastman Kodak Company, USA). Sequencing was carried out by a commercial company (Chromous Biotech Pvt. Ltd., India). 16S rRNA gene amplicons were analyzed by DGGE using D code Universal Mutation Detection System (Biorad, USA). The sequences obtained were deposited in the Genbank database under accession numbers GU968618-GU968638.

Nucleotide sequence accession numbers

The partial 16S rRNA gene environmental sequences obtained from this study were deposited in the Genbank database under accession numbers GU968618-GU968638.

Results

DGGE analysis of bacterial community structure

The DGGE profiles obtained for the 12 sampling sites in the three transects i.e. Goa, Karwar and Ratnagiri is shown in Figure 1. The total number of band positions detected was 82 and the number of bands per sample ranged from 47-62 in all the samples. However, the maximum number of bands was obtained in the OMZ sites 200m, 500m and 1000m at all latitudes. In control sites, the number of bands was more or less similar with 47, 51 and 49 in Goa, Karwar and Ratnagiri respectively. Statistical analysis of the DGGE data showed that at the level of 69% similarity two clusters were formed, which did not show any clustering of similar sites.

The diversity index was also calculated and all the indices showed slight variation among transect and stations it was high in the OMZ sites compared to the non-OMZ sites in all latitudes (Table 2). The Shannon-Weiner diversity index (H’) was calculated. Phylogenetic tree was constructed using 22 sequences obtained from DGGE analysis based on neighbour-joining method using MEGA software [11].
Figure 1: Hierarchical cluster analysis of the 16S rRNA DGGE band profiles of three transects (G-Goa, R-Ratnagiri and K-Karwar) and depths.

Figure 2: Variation in Shannon-Weiner diversity index at various depths (m).

Table 2: Diversity indices of three transect Goa (G), Karwar (K) and Ratnagiri (R).

| Depth       | Abundance | Richness | Evenness | Shannon | Simpson |
|-------------|-----------|----------|----------|---------|---------|
| G 50m       | 47        | 9.9      | 0.92     | 3.55    | 0.97    |
| G 200m      | 51        | 10.8     | 0.89     | 3.51    | 0.96    |
| G 500m      | 61        | 13.02    | 0.92     | 3.82    | 0.98    |
| G 1000m     | 50        | 10.4     | 0.89     | 3.5     | 0.98    |
| K 50m       | 51        | 10.8     | 0.92     | 3.6     | 0.97    |
| K 200m      | 52        | 11.1     | 0.9      | 3.57    | 0.97    |
| K 500m      | 54        | 11.5     | 0.89     | 3.58    | 0.97    |
| K 1000m     | 54        | 11.5     | 0.93     | 3.7     | 0.98    |
| R 50m       | 49        | 10.4     | 0.92     | 3.6     | 0.97    |
| R 200m      | 60        | 12.8     | 0.92     | 3.7     | 0.98    |
| R 500m      | 56        | 11.9     | 0.89     | 3.6     | 0.97    |
| R 1000m     | 62        | 13.2     | 0.92     | 3.8     | 0.98    |
Phylogenetic analysis of excised bands

Figure 3: Neighbor-joining tree showing the phylogenetic affiliations of bacteria derived from the sequences of excised DGGE bands. The sequences obtained in this study are shown in bold. The bar represents 5% sequence divergence. Planctomyces maris was used as the out group.
Twenty two bands were excised from gels corresponding to the different stations in the three transect based on the relative band intensity. Same results were obtained for the bands that were excised from the same position, but in different lanes. Unique bands detected at different sites were also excised. Only dominant bands were sequenced that produced good results, suggesting the presence of other bacteria that may be present in lower numbers. The phylogenetic analysis revealed that the sequences belonged to Proteobacteria, Acidobacteria, Actinobacteria, Chloroflexi, Nitrospira and Bacteriodetes (Figure 3). The number of phylogenetic groups (> 4) was higher in the OMZ region. Proteobacteria was the dominant phylum at all depths with deltaproteobacteria and gammaproteo bacteria forming the major classes. All the groups were invariably observed at all the depths without any differentiation of the OMZ and non-OMZ regions. In the OMZ region the density of the phyla such as Proteobacteria, Nitrospira and Chloroflexi were higher than the non-OMZ region (Figure 4). Proteobacteria accounted for 44% of the population in the OMZ as compared to 39% in the non-OMZ. Nitrospira and Chloroflexi comprised of 9% and 4% respectively in the OMZ and 6% and 3% respectively in the non-OMZ region. However, in the non-OMZ region Bacteriodetes and Acidobacteria, each contributing to 5% of the population. The abundance of Actinobacteria was similar (15%) in both the regions.

![Figure 4: Percentage of phylogenetic groups in OMZ and non-OMZ sites.](image)

**Discussion**

The sampling stations in the present study form a spatial gradient along the continental margin. The site at 50m is located in the continental shelf while the sites at 200m, 500m and 1000m are situated in the continental slope. In this region there is a vertical decrease in the dissolved oxygen concentrations from 0.5mL-1 to as low as 0mL-1. On the other hand, the site in the continental shelf is well ventilated with dissolved oxygen concentrations 3-5mL-1. Oxygen is a strong evolutionary force and the dominating factor determining functional interactions and spatial structure of many microbial communities [12]. Though studies have been carried out on the diversity of bacteria in the OMZ water systematic studies on the bacterial diversity in the sediments is very low [12-16]. Few studies have been conducted in the OMZ sediments of Pacific and Atlantic Ocean [17,18]. Bacterial diversity of the Arabian Sea OMZ water column using cultivation-independent methods have been conducted [13,14,16], but those in Arabian Sea OMZ sediments is meagre [19]. This is the first report of the bacterial diversity in the surficial sediments using PCR-DGGE analysis.

The important groups obtained from this study were Proteobacteria, Acidobacteria, Actinobacteria, Chloroflexi, Nitrospira and Bacteriodetes. Proteobacteria was the dominant group observed in all sites and transects. Proteobacteria is ubiquitous in distribution and its dominance has been reported in AS OMZ water [16]. The Deltaproteobacteria, which includes the anaerobic sulphate reducing bacteria, were the most abundant class, suggesting the prevalence of sulphate reduction in this region. Sulphate reducing bacteria falling in genera Desulfosarcina and Desulfofrigus have been retrieved from the OMZ waters of Arabian Sea [13,20]. 73% of the sequences showed close affiliation to Syntrophobacterales and Desulfovibrionales. Sulphate concentrations varied from 39.8 to 12.7mm in the pore waters of the eastern AS indicating sulphate reduction [21]. Similar occurrence of high sulphate reducers has been reported in other marine sediments [22]. The dominance of this group was not restricted only to Mid Atlantic Ridge [23] but also in Tokyo and Sagami bays [24,25]. However, a few did not show any resemblance to cultured representatives but were closely related to clones detected at cold seeps and mud volcano [26,27].

Gammaproteobacteria formed a major group after Deltaproteobacteria in the phylum Proteobacteria in the both the sites with higher occurrence in the OMZ. Occurrence of this has been reported among culturable and non-culturable bacteria of pelagic waters and sediments [28,29]. There have also been reports of active diastrophic planktonic Gammaproteobacteria in the Arabian Sea [30]. Thus the possibilities of the presence of sediment diastrophic gammaproteobacterial communities cannot be ruled out and future studies should be conducted using functional probes. Also the largest bacteria which have been reported such as Thioploca sp. falling in the Gammaproteobacteria in the OMZ sediments off Oman have not been observed in OMZ sediments.
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Chloroflexi was restricted to the deeper two OMZ sites. The presence of Chloroflexi related SAR 202 cluster, which forms a monophyletic subgroup in the phylum, has been reported in the oxic and the oxygen minimum zone waters although these were not found in the present study [14,36]. The members of this phylum exhibit unusual metabolic diversity like anoxygenic photosynthesis, organotrophy and reduction of chlorinated hydrocarbons [37]. Schafer et al. [18] in their work on the microbial diversity of the sediment of Benguela upwelling system reported the presence of this phylum in sulphate-depleted depths [18]. Their presence seems to be a common feature of many deep sub-seafloor sediments [38,39]. Thus the present studies demonstrate that eastern AS OMZ region possesses a phylogenetically distinct bacterial community that may be playing an important role in the prevailing biogeochemical processes. The data obtained from DGGE profiling is more accurately considered as the structure of dominant populations rather than a general measure of bacterial diversity [40]. The deeper depths i.e., OMZ region lodged more bacterial populations than non-OMZ region suggesting that the environmental conditions prevalent in these regions provide a favorable condition for the survival of bacteria.

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