Diversity and extracellular enzyme activities of heterotrophic bacteria from sediments of the Central Indian Ocean Basin

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Sedimentary bacteria play a role in polymetallic nodule formation and growth. There are, however, limited reports on bacterial diversity in nodule-rich areas of the Central Indian Ocean Basin (CIoB). In this study, bacterial abundance in thirteen sediment cores collected from the CIoB was enumerated, followed by phylogenetic characterisation and, screening of select heterotrophic bacteria for extracellular enzyme activities. Total bacterial counts (TBC) were in the order of $10^7$ cells g$^{-1}$; there was a significant difference ($p > 0.05$) among the cores but not within the sub-sections of the cores. The retrievable heterotrophic counts ranged from non-detectable to $5.33 \times 10^5$ g$^{-1}$; the heterotrophic bacteria clustered within the phyla Firmicutes, Proteobacteria and Actinobacteria. Bacillus was the most abundant genus. The extracellular enzyme activities were in the order: amylase > lipase > protease > phosphatase > Dnase > urease. Major findings are compared with previous studies from the CIoB and other areas.

Polymetallic nodules (PMNs) are found on the seabed and believed to replace the land-based deposits as sources of metals in future¹. PMNs are majorly composed of concentric layers of iron and manganese oxides around a core, and they are abundantly found at 4000–6000 m water-depth in major oceans². PMNs are rich in metals such as manganese (Mn), nickel (Ni), copper (Cu), cobalt (Co) etc. Regions in the Pacific and Indian Oceans are under consideration for industrial-level mining of PMNs³. India is presently conducting environmental impact assessment (EIA) studies in the CIoB (0°–25°S and 70°–90°E)³ to evaluate the possible impact of mining of PMNs on the deep-sea environment (Fig. 1).¹ PMNs are formed abiotically, involving hydrogenetic and diageneric processes, on the seabed. It is hypothesised that microbial activities, especially of sedimentary bacteria, may contribute towards precipitation of metal hydroxides and play a role in formation and growth of the nodules⁴. Sedimentary bacteria are already known to play crucial roles in nutrient cycling in the deep-sea environment⁷, with the help of extracellular enzymes⁸.

Chandramohan et al.⁹, in one of the earliest studies from the Indian Ocean (02°N, 58°E), characterised bacteria present in nodules and associated sediments. They reported the presence of both Mn(II)-oxidising and MnO₂-reducing psychrophilic bacteria in the nodules. Bacterial taxa Arthrobacter, Bacillus, "coryneforms", Enterobacteriaceae, Micrococcus, Staphylococcus and Vibrio were identified from the nodules and associated sediments. Later, Loka Bharathi & Nair¹⁰ reported Acinetobacter, "coryneforms", Enterobacter, Micrococcus, Moraxella, Pseudomonas and Staphylococcus from the sediments of the CIoB. Naik et al.¹¹ isolated and identified 17 bacterial genera from the sediment samples of the southern and northern regions of the CIoB (Supplementary Table S1).

There are few reports on gene sequence-based identification and taxonomy of culturable bacteria from the CIoB, though there are reports on metagenomic studies [e.g. Das et al., Sujith et al.¹³ and Wang et al.¹⁴]. Das et al.¹²

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performed 454 pyrosequencing of bacterial communities of red clays of the CIOB (16°S, 75.5°E) and reported 14 phyla, 25 classes, 38 orders, 52 families and 62 genera. Sujith et al. studied bacteria from sea-mount samples from the CIOB (03°S, 83°E) and reported the dominance of Proteobacteria and bacterial genera Halomonas and Thiobiospira. Recently, Wang et al. characterised the microbial communities in a gravity core sediment sample from rare earth element (REY)-rich mud of the Indian Ocean (22°S, 81°E) using Illumina high-throughput sequencing of 16S rRNA genes and revealed the dominance of Proteobacteria, followed by Firmicutes and Actinobacteria (Supplementary Table S1).

During a cruise (SSD-013) to collect baseline geochemical, biochemical and microbiological data from the First Generation Mining site and the Preservation Reference Zone (PRZ) site, we aimed to enumerate down-core variation of bacterial abundance in the deep sub-seafloor sediments of the CIOB. This was followed by 16S rRNA gene-based phylogenetic characterisation and screening of heterotrophic bacterial cultures for extracellular enzyme activities, including amylase, protease, lipase, DNAse, urease, alkaline phosphatase and nitrate reduction.

Results

Abundance of sedimentary bacteria. The water depth at the sampling locations ranged from 4856–5297 m. Down-core variation of total bacterial counts (TBC) in the 13 sediment box-cores is presented in Fig. 2. The TBC were in the order of 10^7 cells g^-1 of dry sediment and higher at 0–4 cm bsf. Highest TBC of 8.12 × 10^7 cells g^-1 of dry sediment were observed in the BC-10, subsection 6–8 cm bsf. Conversely, lowest TBC of 1.10 × 10^7 cells g^-1 of dry sediment were observed in the BC-13, subsection 25–30 cm bsf. Two way ANOVA analysis showed that there is a significant difference (p < 0.05) in the TBC between the sediment cores. However, no significant difference in the TBC was observed within the core sub-sections.

The retrievable counts of heterotrophic bacteria (colony forming units, CFUs) performed in duplicates and averaged later are presented in Fig. 3. The CFUs ranged from non-detectable to 5.33 × 10^5 g^-1 of dry sediment. Down-core variation in the CFUs was higher at 4–6 cm bsf. The highest CFUs were recovered from the BC-02, subsection 4–6 cm bsf. Higher CFUs were recovered from the upper 10 cm, compared to the deeper sub-sections of the cores (Fig. 3). The heterotrophic bacteria were not detectable in the cores BC-09 and BC-10, though the TBC from the cores were in the order of 10^7 g^-1 of dry sediment.

Phylogenetic characterisation of heterotrophic bacteria. In total, 62 heterotrophic bacteria were isolated. Only 43 bacterial cultures were included in the phylogenetic analysis due to poor viability of the 19 cultures. Phylogenetic relationships of the bacteria, inferred based on 16S rRNA gene sequence-data analysis, are presented in Fig. 4. The system output details retrieved from MEGA 7 about the tree construction are in Supplementary Document. In the NJ tree (Fig. 4), 43 newly generated gene-sequences clustered within three bacterial phyla, Actinobacteria, Firmicutes and Proteobacteria (subphyla α-Proteobacteria and γ-Proteobacteria). The phylogenetic analysis recovered 15 bacterial clades, including Bacillus clades I-V, Oceanobacillus, Staphylococcus, Pseudomonas (Proteobacteria), Brachybacterium, Brevibacterium, Micrococcus and Streptomyces (Actinobacteria) (Fig. 4). Location-wise and subsection-wise groupings of the identified clades are presented in Supplementary Tables S2 and S3, respectively.

Screening for extracellular enzyme activities. The enzyme activities of the 43 bacterial cultures followed the order: amylase > lipase > protease > phosphatase > Dnase > urease. Nine cultures were positive for nitrate reduction. Nine Bacillus cultures isolated from 4–6 cm subsection bsf were positive for most of the enzyme...
activities (Table 1). Streptomyces sp. VJ21 was positive for all the extracellular enzyme activities and nitrate reduction. Staphylococcus sp. VJ01 was positive for only nitrate reduction. Alteromonas sp. VJ25 was negative for all the enzyme activities and nitrate reduction.

**Discussion**

This is the first study reporting 16S rRNA gene-based phylogenetic characterisation of heterotrophic bacteria from sub-seafloor sediments of PMN-rich areas in the CIOB. The forty-three heterotrophic bacteria clustered within three phyla and 11 known genera (Table 1). Bacillus species were majorly isolated, which formed 5 distinct clades. Bacillus is widely-reported from the CIOB and other parts of the Indian Ocean9–13,16,17, especially in PMNs and associated sediments. Further, four bacteria grouped within Oceanobacillus. Oceanobacillus is a new record from the sediments of the CIOB. It appears that culturable bacilli are abundant in the PMN-rich sediments of the CIOB. Similar patterns of dominance by Firmicutes, including bacilli, were reported by Sujith et al.13, Wang et al.14, and Khandeparker et al.16 from the Indian Ocean. Bacilli are known to survive and grow under physico-chemical extremities and with high metabolic versatility. This possibly can explain the abundance of culturable bacilli in the CIOB sediments.

To the best of our knowledge, Brachybacterium, a member of the phylum Actinobacteria, has not been reported from the CIOB (Supplementary Table S1), even though they are known from deep-sea sediments of Southern Ocean18 and Pacific Ocean deep-sea Mn nodule sediment19. This genus is reported for the first time from the CIOB sediment samples. Interestingly, Brachybacterium sp. isolated from the nodules in the Pacific Ocean19 reportedly showed high Mn(II) resistance and Mn(II)-oxidizing/removing abilities. Brevibacterium, a member of “coryneforms” bacteria reported in this study, might have been also isolated from the CIOB sediments and classified under “coryneforms” by Loka Bharathi & Nair10. Chandramohan et al.3 and Sujith et al.17 reported “coryneforms” and Brevibacterium members from the nodules and basal fragments collected from the non-CIOB regions of the Indian Ocean, respectively (Supplementary Table S1).

Glycocaulis, to the best of our knowledge, has not been previously reported from the CIOB (Supplementary Table S1). The genus was originally described from hot water plume of a deep-sea hydrothermal vent from
Canada and it reportedly lacks phospholipids. Role of bacterial phospholipids in deep-sea ecology is subject to future studies. *Micrococcus* is previously reported from the sediments of the CIOB by Loka Bharathi & Nair and Naik et al. Chandramohan et al. reported this genus from nodules and associated sediments from a non-CIOB region in the Indian Ocean. *Micrococcus* species are known to have the ability to adapt to elevated pressure, while some species of *Micrococcus* may produce antibiotics.

*Pseudomonas* species have been reported from the CIOB sediments (Supplementary Table S1) and also from a few non-CIOB regions. *Pseudomonas* species are reportedly involved in Mn-oxidation, and they possibly adsorb metals such as Co, Cu, and Ni. Interestingly, they may form biofilms which help in the growth of nodules. *Paracoccus* has not been previously reported from the CIOB sediments, though the genus is known from deep sediments of the Arctic Ocean.

*Staphylococcus* has been reported from the CIOB and the non-CIOB regions (Supplementary Table S1). Recently, Wang et al. reported the whole genome sequence of a Mn(II) tolerant *Staphylococcus* sp. isolated from deep-sea sediment in the Clarion-Clipperton Zone in the Pacific ocean, which exhibited efficient Mn(II) oxidation. Further studies are required from the CIOB about molecular mechanisms and pathways involved in Mn(II) oxidation by bacteria such as *Staphylococcus* spp. Bacterial genus *Streptomyces* is reported for the first time from the CIOB sediments. *Streptomyces* is known for the production of secondary metabolites. Recently, Rodriguez et al. reported *Streptomyces cyaneofuscatus* isolated from a Gorgonian coral collected at 1500 m depth in the Avilés submarine Canyon which produced antibiotic Anthracimycin B. Role of bacterial secondary metabolites, including antibiotics, in deep-sea ecology of the CIOB, needs further investigations.

Figure 3. Down-core variation in the heterotrophic bacterial counts (CFUs) in the CIOB sediments.
In our study, we observed that the TBC were two orders higher than the CFUs. The TBC were in the order $10^7$ g$^{-1}$ of dry sediment. Similar counts were reported by Naik et al.$^{11}$ from the CIOB, while Raghukumar et al.$^{27}$ reported TBC two orders higher than our findings. There was no significant difference in the TBC within the core sub-sections, suggesting a homogeneous distribution pattern of total bacterial abundance. Loka Bharathi and Nair$^{10}$ and Raghukumar et al.$^{27}$ reported similar homogeneous distribution patterns. Geological studies from the CIOB indicate that sediment characteristics (e.g. porous, mottled and loose)$^{28}$ and disturbance of sediment layers by biological activities such as bioturbation$^{11}$ could be responsible for this kind of homogeneous distribution pattern. ISBA guidelines (2018) recommend the use of molecular tools to quantify bacterial abundance in the

Figure 4. Phylogenetic characterization of select heterotrophic bacteria based on 16S rRNA gene sequence analysis in MEGA.
proposed PMN-mining areas. Traditional methods, therefore, may be replaced with modern molecular tools in microbial abundance studies, in near future E.g. Cho et al.29.

The cultivability of heterotrophic bacteria in our study was about 1% of the TBC. Similar results were reported by Nair et al.30 and Raghukumar et al.27 from the CIOB and by Chandramohan et al.28 from a non-CIOB region. Naik et al.11, however, reported CFU counts one order lesser than the present study. The retrievable counts of heterotrophic bacteria have been used to quantify variations in bacterial abundance under the influence of disturbance in environmental conditions. For example, Loka Bharathi and Nair10 observed a significant increase in retrievable nitrate reduction. Table 1. List of bacterial cultures included in the phylogenetic analysis, their extracellular enzyme activities and nitrate reduction.

| Sl. no. | Clade name | Isolate no. | Station no. | Core depth (cm) | Amylase | Protease | Lipase | Dnase | Urease | Alkaline | Phosphatase | Nitrate | Genbank no. |
|---------|------------|-------------|-------------|----------------|---------|----------|--------|-------|--------|----------|-------------|---------|-------------|
| 1       | Alteromonas | VJ25 | BC-06 | 35–40 | -- | - | - | - | - | - | - | -- | MH605399 |
| 2       | Bacillus clade I | VJ10 | BC-02 | 4–6 | + | -- | + | + | - | -- | -- | -- | MH605389 |
| 3       | Bacillus clade I | VJ28 | BC-07R | 2–4 | + | -- | -- | -- | -- | -- | -- | -- | MH605402 |
| 4       | Bacillus clade I | VJ30 | BC-07R | 4–6 | + | -- | -- | -- | -- | -- | -- | -- | MH605404 |
| 5       | Bacillus clade I | VJ31 | BC-07R | 6–8 | + | -- | -- | -- | -- | -- | -- | -- | MH605405 |
| 6       | Bacillus clade I | VJ32 | BC-07R | 6–8 | + | -- | -- | -- | -- | -- | -- | -- | MH605406 |
| 7       | Bacillus clade I | VJ35 | BC-07R | 6–8 | -- | -- | -- | -- | -- | -- | -- | -- | MH605408 |
| 8       | Bacillus clade I | VJ37 | BC-07R | 6–8 | + | -- | + | -- | -- | -- | -- | -- | MH605410 |
| 9       | Bacillus clade I | VJ45 | BC-08R | 15–20 | -- | -- | -- | -- | -- | -- | -- | -- | MH605415 |
| 10      | Bacillus clade I | VJ48 | BC-10 | 30–35 | + | -- | + | -- | -- | -- | -- | -- | MH605417 |
| 11      | Bacillus clade I | VJ47 | BC-10 | 35–40 | + | -- | + | -- | -- | -- | -- | -- | MH605416 |
| 12      | Bacillus clade I | VJ53 | BC-12 | 4–6 | + | -- | + | -- | -- | -- | -- | -- | MH605420 |
| 13      | Bacillus clade I | VJ54 | BC-12 | 4–6 | + | -- | + | -- | -- | -- | -- | -- | MH605421 |
| 14      | Bacillus clade II | VJ41 | BC-08R | 2–4 | -- | -- | -- | -- | -- | -- | -- | -- | MH605413 |
| 15      | Bacillus clade II | VJ39 | BC-08R | 4–6 | -- | -- | + | -- | + | -- | -- | -- | MH605412 |
| 16      | Bacillus clade III | VJ07 | BC-02 | 30–35 | + | -- | + | -- | -- | -- | -- | + | MH605387 |
| 17      | Bacillus clade IV | VJ38 | BC-08R | 4–6 | -- | + | -- | -- | -- | -- | -- | -- | MH605411 |
| 18      | Bacillus clade IV | VJ44 | BC-08R | 8–10 | -- | -- | -- | -- | -- | -- | -- | -- | MH605414 |
| 19      | Bacillus clade V | VJ11 | BC-03 | 40–45 | + | -- | + | -- | -- | -- | + | -- | MH605390 |
| 20      | Bacillus clade V | VJ13 | BC-04 | 8–10 | + | -- | + | -- | -- | -- | + | -- | MH605392 |
| 21      | Bacillus clade V | VJ27 | BC-06 | 20–25 | + | -- | + | -- | -- | -- | -- | + | MH605401 |
| 22      | Bacillus clade V | VJ29 | BC-07R | 2–4 | + | + | + | -- | -- | -- | -- | + | MH605403 |
| 23      | Brachybacterium | VJ26 | BC-06 | 20–25 | + | -- | + | -- | -- | -- | -- | + | MH605400 |
| 24      | Brachybacterium | VJ37 | BC-13 | 4–6 | + | + | + | -- | -- | -- | + | -- | MH605423 |
| 25      | Brevibacterium | VJ30 | BC-04 | 0–2 | -- | -- | -- | -- | -- | -- | -- | -- | MH605384 |
| 26      | Brevibacterium | VJ05 | BC-01 | 0–2 | -- | -- | -- | -- | -- | -- | -- | -- | MH605385 |
| 27      | Glycocaulis | VJ19 | BC-05 | 0–2 | -- | -- | -- | -- | -- | -- | -- | -- | MH605396 |
| 28      | Micrococcus | VJ14 | BC-04 | 23–30 | -- | -- | -- | -- | -- | -- | -- | -- | MH605393 |
| 29      | Micrococcus | VJ32 | BC-11R | 0–2 | + | -- | + | -- | -- | -- | + | -- | MH605419 |
| 30      | Micrococcus | VJ61 | BC-13 | 4–6 | + | -- | + | -- | -- | -- | -- | -- | MH605425 |
| 31      | Micrococcus | VJ60 | BC-13 | 0–2 | + | -- | + | -- | -- | -- | -- | -- | MH605424 |
| 32      | Oceanobacillus | VJ09 | BC-02 | 4–6 | + | + | + | -- | -- | -- | -- | -- | MH605388 |
| 33      | Oceanobacillus | VJ17 | BC-04 | 8–10 | -- | -- | + | -- | -- | -- | -- | -- | MH605394 |
| 34      | Oceanobacillus | VJ18 | BC-04 | 35–40 | -- | -- | + | -- | -- | -- | -- | -- | MH605395 |
| 35      | Oceanobacillus | VJ23 | BC-06 | 2–4 | -- | -- | + | -- | -- | -- | -- | -- | MH605398 |
| 36      | Paracoccus | VJ36 | BC-13 | 6–8 | + | -- | -- | -- | -- | -- | -- | -- | MH605422 |
| 37      | Pseudomonas | VJ34 | BC-07R | 6–8 | + | -- | -- | -- | -- | -- | -- | + | MH605407 |
| 38      | Pseudomonas | VJ36 | BC-07R | 6–8 | + | -- | + | -- | -- | -- | -- | + | MH605409 |
| 39      | Staphylococcus | VJ01 | BC-01 | 0–2 | -- | -- | + | -- | -- | -- | + | -- | MH605383 |
| 40      | Staphylococcus | VJ06 | BC-01 | 35–40 | -- | -- | -- | -- | -- | -- | -- | -- | MH605386 |
| 41      | Staphylococcus | VJ49 | BC-11R | 2–4 | -- | -- | -- | -- | -- | -- | -- | -- | MH605418 |
| 42      | Streptomyces | VJ12 | BC-04 | 8–10 | -- | -- | -- | -- | -- | -- | -- | + | MH605391 |
| 43      | Streptomyces | VJ21 | BC-05 | 0–2 | + | -- | + | + | ++ | -- | -- | + | MH605397 |

Table 1. List of bacterial cultures included in the phylogenetic analysis, their extracellular enzyme activities and nitrate reduction.
The 43 heterotrophic bacterial cultures screened for seven extracellular enzymatic activities showed promising results. Hydrolytic degradation of polymers such as starch, protein and lipids by bacterial cultures is a possible indicator of mineralisation of polymers and regeneration of nutrients in the deep-sea. In this study, 25 bacterial cultures exhibited amylase activity. Chandramohan et al., Rajhu kumar et al. and Loka bharathi and Nair have previously reported amylase activity from the bacterial cultures from the Indian Ocean. Loka Bharathi and Nair suggested that amylase activity by bacterial cultures from the sediments indicates the availability of free sugar in the CIOB sediments. Lipase activity was observed in 20 bacterial cultures (Table 1), suggesting in situ lipolytic activity of the bacteria. Seventeen cultures were positive for protease activity, and possibly these bacteria are involved in break-down of complex proteins present in the sediments into simpler peptides and amino acids. Fourteen cultures were positive for phosphatase activity. Recently, Biche et al. reported phosphatase activities of the sediment samples as an indication of available phosphate in the CIOB sediments. The bacterial enzymatic activities of phosphatases and nitrate reductases in the present study indicate that the sedimentary bacterial communities possibly participate in nutrient recycling. These are preliminary observations revealed in this study, which need to be subjected to further research for confirmation.

**Conclusion**

This is the first study reporting 16S rRNA gene-based phylogenetic characterization of heterotrophic bacteria from the sediments of potential PMN-mining areas in the CIOB. As evident from the present study and other studies, PMNs and associated sediments support phylogenetically-diverse bacterial diversity, which is reportedly distinct from the deep water column (Supplementary Table S1). A comprehensive understanding of microbial diversity present in nodule-rich areas of the CIOB is required to assess the impact of large scale removal of nodules and sediments from the deep sea-bed for industrial mining of the PMNs, as suggested Schulse et al. in the context of Pacific Ocean. Microbial abundance, diversity and functions can vary across the seabed, therefore studies employing culture-dependent and -independent methods are required to improve our understanding of microbial diversity and their role in deep-sea ecology.

**Methods**

**Sampling.** Sampling was carried out during July–August 2015 with Research Vessel Sindhu Sadhana (SSD-013), as part of ongoing Polymetallic Nodules Program (PMN). The sub-seafloor sediment samples were collected from 13 locations in the CIOB (Fig. 1; Supplementary Table S4), using USNEL-type box-corer of 50 cm × 50 cm × 50 cm dimension.

The box-corer was partitioned into four parts by fixing metallic plates. After siphoning the overlying water, sub-samples were collected using clean 6.3 mm inner diameter plexiglass tubes. On an average, 45 cm bsf long cores were obtained. The cores were sliced at 2 cm intervals up to 10 cm depth and at 5 cm intervals thereafter and extruded aseptically to sterile polythene bags on board. The bags with sediment samples were preserved at 4 °C until further processing.

**Abundance of bacteria in deep sub-seafloor sediment samples.** For TBC, approximately 1 g of sediment sample was suspended in 9 ml of sterile seawater and processed as detailed in Das et al. The DAPI method was used to count the cells. Total bacterial counts were expressed as cells g⁻¹ of dry sediment.

For retrievable counts of heterotrophic bacteria, each sub-sample was serially diluted to 10⁻² with filtered sterile seawater as diluent on-board. Zobell marine agar (ZMA, 12.5%) medium (HiMedia, India) prepared with 50% seawater was spread-plated with 100 μl of original and 10⁻² inocula, in duplicates, and incubated at 4 °C for a month and later transferred to 25 °C for another 15 days. After incubation, the plates were examined for retrievable CFUs and calculated per gram of dry sediment.

Bacterial colonies with apparent unique morphology were selected and purified by quadrant streaking method. All the purified bacterial cultures were maintained on ZMA by sub-culturing after every four months as recommended by Bowman for further studies. The bacterial cultures were also preserved in 80% glycerol stocks and stored -80 °C for long-term.

**DNA extraction and PCR amplification of 16S rRNA gene.** Select bacterial cultures were cultivated in Zobell marine broth (ZMB, 12.5%) for 3–7 days at 25 °C, followed by DNA extraction using ZM Fungal/Bacterial DNA MiniPrep Kit (Zymo Research, D6005). 16S rRNA gene was amplified by PCR using the primers 27F (5′-AGAGTTTGATCCTGGCTCAG-3′) and 1492R (5′-TACGGYTACCTTGTTACGACTT-3′). PCR amplification was performed in a reaction volume of 50 μl, including 5 μl of 10X concentrated buffer (with 15 mM MgCl₂) (Genei, Bangalore, India), 1 μl of 10 mM dNTP mix (Genei), 1 μl each of 20 pmol/μl 27F and 1492R primers, 1.5 μl of Taq polymerase (1 unit/μl) (Chromous Biotech, India), 3 μl 20–50 ng template DNA and 37.5 μl nuclease free water. PCR conditions used were: initial denaturation of 5 min at 95 °C, followed by 35 cycles of 1 min at 95 °C, 1 min at 50 °C and 2 min at 72 °C, and a final extension of 72 °C for 10 min. Purification of PCR product was carried out using QIAquick PCR purification columns (QIAGEN, Catalogue number 28106), following the manufacturer’s instructions. The PCR amplicons (DNA) were stored at -20 °C.

**Gene sequencing and phylogenetic analysis.** The purified PCR products were sequenced using Genetic Analyzer 3130xl (ABI) based on the Big Dye termination v 3.1 (Chain termination) chemistry using the same primer set. For each isolate, a consensus sequence was generated in DNA Dragon (SequentiX, Germany) using both forward and reverse sequences, where possible.

A multiple alignment of 16S rRNA gene sequences, including newly generated sequences from this study and homologous sequences retrieved from NCBI GenBank (Fig. 2), was prepared in MEGA. The 16S rRNA gene sequences generated in this study have been deposited in the NCBI GenBank under accession numbers MH605383–MH605425 (Table 1). Phylogenetic relationships of the bacteria from deep-sea sediments were evaluated in MEGA.
Screening for extracellular enzymatic activities. The 43 bacterial cultures were screened for extracellular hydrolytic enzyme activities, including amylases, proteases, lipases, DNases and ureases on agar medium having 1% substrates, i.e. starch, casein, tween 80, toluidine blue and urea, respectively. Alkaline phosphatase and Nitrate reductase were also checked for these cultures. After spot inoculation, the plates were incubated for 4 days at 25 ± 1°C.

Statistical analysis. Two-way analysis of variance (ANOVA) was carried out using Statistica ver. 6 (Statsoft) to find out significant variations in bacterial abundance within the depths and between the cores.

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Acknowledgements
The authors thank the following: Director, CSIR-NIO; Head, Biological Oceanography Division; Ministry of Earth Sciences (MoES), Govt. of India for funding the project GAP 2128; CSIR India for funding under project PSC0206 for DNA sequencing; Drs Rahul Sharma, LokaBharathi, H.N. Khadge, T.N.R. Srivivas and Moturi Shri Ramakrishna for their support and encouragement; and the Master and crew who provided support in scientific operations, during the cruise SSD013. This manuscript has NIO Communication Number 6395.

Author Contributions
M.S.S., S.R.D. and B.D.S. conceived the idea, All authors (V.S.G., M.S.S., S.R.D., D.P., R.M.M. and B.D.S.) conducted experiments; V.S.G., M.S.S. and B.D.S. analysed the results. All authors reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-45792-x.

Competing Interests: The authors declare no competing interests.

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