Environmental influence on cultivable microbial community in the sediment of Sundarban mangrove forest, India

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Accepted 20 August, 2013

INTRODUCTION

Mangroves are highly productive marine ecosystem where bacteria actively participate in biomineralization and biotransformation of minerals (Gonzalez-Acosta et al., 2006). The distribution of microbial activities in estuarine systems is clearly complex and variable. Much research remains to be done in order to define the distributions of microbial activities and the major factors involved in controlling these distributions in estuaries. Leaves and wood provided by mangrove plants to the sediment are degraded primarily by large variety of microbes which actively participated in the heterotrophic food chain (Thatoi et al., 2012; Alongi et al., 1989, 1993; Alongi,1994). Major products of general recycling of organic matter are detritus which is rich in enzymes and proteins and contains large microbial population (Holguin et al., 2001). Bacteria are the major participants in the carbon, sulphur, nitrogen and phosphorous cycles in mangrove forest (Toledo et al.,1995, Vazquez et al., 2000; Rojas et al., 2001). Bacterial activity is responsible for most of the carbon recycling in mangrove sediment under both oxic and anoxic condition (Das and Dangar, 2008). Many species of phosphate solubilizing rhizosphere
bacteria associated with black mangrove roots were found. The mechanism for phosphate solubilization probably involves the production of several organic acids (Vazquez et al., 2000).

Saprophytic fungi are fundamental to many aspects of decomposition and energy flow in mangrove forests (Nedwell, 1982; Radhakrishnan et al., 2011). Most investigation of anaerobic metabolism in natural ecosystem have dealt with sulfate rich marine sediments where sulfate reduction is the dominating process or eutrophic lake sediments where sulfate and nitrate is depleted in the hypolimnion and in the superficial sediment layers leaving terminal carbon mineralization principally to methane producing bacteria (Sahoo and Dhal, 2009; Senior et al., 1982; Lovley and Klug, 1982). Sulfate reduction, methane production, and denitrification are the important processes for the terminal electron removal during decomposition of organic matter in anoxic environment. The methanogens are characterized by their ability to produce methane from hydrogen and carbon dioxide, formate, acetate, methanol, etc (Chen et al., 2010; Mohanraju and Natarajan, 1992). Methanotrophs are a subset of a physiological group of bacteria known as methylotrophs. They are unique in their ability to utilize methane as a source of carbon and energy (Liebner et al., 2008; Chen-rul et al., 2003). Nitrogen fixing bacteria are the other group of bacteria that are involved in formation of ammonia or organic nitrogen from atmospheric nitrogen. They may be free living or symbiotic in nature. It has been studied that N2 fixation by heterotrophic bacteria are generally regulated by specific environmental factors like oxygen, combined Nitrogen and the availability of carbon source for energy requirement (Teri and Mary, 2005). Aerobic, autotrophic nitrifiers oxidize ammonia to nitrite and nitrate, with molecular oxygen as electron acceptor (Nicol et al., 2008). Nitrite and nitrate are reduced to dinitrogen gas by heterotrophic denitrifying bacteria that use NOx instead of oxygen as electron acceptor (Hayatsu et al., 2008; Riley et al., 1995).

The purpose of the present study was to examine seasonal and spatial variations in microbial population (bacteria and fungi) in mangrove soil and to find out the correlation between different microbes with nutrients.

Study area

Sundarban Mangrove forest is located geographically in between 21° 31’ N and 22° 30’ N and longitude 88° 10’ E and 89° 51’ E along the North East coast of Bay of Bengal, India. This mangrove forest is a part of the estuarine system of the River Ganges, NE coast of Bay of Bengal (Figure 1), which covers 9630 km2, out of which 4264 km2 of inter-tidal area, covered with thick mangroves, is subdivided as forest sub-ecosystem and 1781 km2 of water area as aquatic sub-ecosystem. The tide in this estuarine complex is semidiurnal in nature with spring tide range between 4.27 and 4.75 m and neap tide range between 1.83 and 2.83 m. It is a unique bioclimatic zone in land ocean boundaries of Bay of Bengal and the largest delta on the globe. Several numbers of discrete islands constitute Sundarbans. One of these Islands, Lothian Island covering an area of 38 km2 has been notified as a sanctuary and is situated at the confluence of Saptamukhi River and Bay of Bengal. In the southern part of the island, the ground level is high while in the northern areas, the land is low and gets inundated during highest high tide. Mangroves, Avicennia alba, Avicennia marina and Avicennia officinalis are the dominant species, Excoecaria agallocha and Heritiera fomes are thinly distributed and Ceriops decandra is found scattered all over the island. The deltaic soil of Sundarban Biosphere Reserve comprises mainly with saline alluvial soil consisting of clay, silt, fine sand and coarse sand particles. It is described as very deep, poorly drained, fine soils occurring on level to nearly level lower delta with loamy surface, severe flooding and very strong salinity (extensive extent) associated with very deep, very poorly drained, fine loamy soil. Sediment samples were collected from different sites namely rooted, un-rooted and deep forest regions at Lothian Island to critically examine the spatial variations of depth integrated microbial diversity in these mangrove ecosystem.

MATERIALS AND METHODS

Sediment cores were collected aseptically using a hand held stainless steel core sampler (3.2 cm diameter, 100 cm long) from the different tidal zone of Sundarban Mangrove Forest and from different depth (0-10, 10-20, 20-30, 30-40, 40-50, 50-60 cm) during pre-monsoon, monsoon and post monsoon. Samples were collected into sterilized container and were transferred to laboratory in iced condition for both chemical and microbiological assay. The three tidal zones were sediment from dense forest region, sediment from the region with pneumatophores near mid littoral zone (rooted), and the sediment from the lower littoral zone where no pneumatophore were found (un-rooted).

Quantification of bacteria and fungi

Sediment samples were stored at 4°C immediately after collection and transported to the laboratory, for analysis with adequate care. 10 g of sample from different depth of different regions were homogenized with sterilized phosphate buffer solution. Serial dilutions up to 10-4 were made and inoculation was done with 0.1 ml. Quantification of bacteria and fungi from mangrove sediments was carried out by spread plate method for different type of bacteria such as phosphorous solubilising bacteria (PSB), cellulose degrading bacteria (CDB), nitrifying bacteria, free living nitrogen fixing bacteria and fungi and they were incubated at different condition (Ramanathan et al., 2008). Sulfate reducing bacteria was carried out by spread plate method for different type of bacteria such as phosphorous solubilising bacteria (PSB), cellulose degrading bacteria (CDB), nitrifying bacteria, free living nitrogen fixing bacteria and fungi and they were incubated at different condition (Ramanathan et al., 2008). Sulfate reducing bacteria was cultured in Starkey’s medium in anaerobic condition (Fathul et al., 2008).

Sediment quality measurement

Sulphate, nitrite-nitrogen, nitrate-nitrogen, phosphate, silicate
concentration of the sediment sample was measured at 10 cm interval (from 0 to 60 cm depth). 30 g of soil subsample was collected from the different depth and was immediately extracted in 75 mL of 2 mol L\(^{-1}\) potassium chloride (KCl). The mixture was shaken until well mixed and allowed to stand overnight (Riley et al., 1995). After 24 h, 4 mL of the supernatant was collected for the estimation of different nutrients using standard spectrophotometric methods (Grasshoff, 1983). The pH value was measured in a 1:5 (w/w) soil water suspension using electric digital pH meter (Tiwari et al., 1989) and soil organic carbon was measured by standard methods (Walkley and Black, 1934).

RESULTS AND DISCUSSION

Mangrove sediment at Indian Sundarban showed seasonal variation with respect to both major nutrient concentrations and microbial population. Beside monsoonal addition of nutrients to the system mangrove, litters also played a significant role in regulating the nutrient status that in turn controls the microbial population (Das et al., 2012). Among several physical factors
tidal inundation, wave action, presence of mangrove roots and bioturbation are the important factors considered for determining microbial abundance in the mangrove sediment from surface to a depth up to 60 cm (Laing et al., 2009; Bharathkumar et al., 2008). During pre-monsoon, nutrient concentration in soil sample of deep forest region showed very weak stratification from surface to 30 cm of depth with almost uniform distribution. Intense bioturbation up to 30 cm depth by several benthic organisms could cause uniform mixing of soil nutrients. No significant variation of silicate concentration was found throughout the entire depth.

Again silicate concentration was found to be more in unrooted and rooted region than that of deep forest region, because rooted and un-rooted region is at the nearest part of the river and the ultimate source of silicate is riverine system. Gradual decrease in organic carbon and phosphate-phosphorous concentration was observed from 30 to 60 m. During transportation of organic matter from surface to bottom, it is decomposed by microbes. As a result, organic content of soil decreased with increasing depth (Kristensen et al., 2008). Organic carbon in the deep forest sediment was found to be more than that of rooted and un-rooted region. It could be attributed to mangrove litter fall with an annual rate of 1603 g m⁻² year⁻¹ (Ghosh et al., 1990). Both rooted and un-rooted regions regularly experience significant tidal flushing which carried away significant amount of mangrove litter. In contrary, the deep forest seldom gets inundated by tidal water resulting to organic carbon rich sediment. In all the sediments, organic carbon was found maximum during monsoon followed by post monsoon and pre-monsoon (Figure 2a).

The microbial population was also found maximum in the deep forest sediment relative to the other two sites (Figure 2a). Nitrate-Nitrogen concentration was increased from surface to 40 cm of depth but decreased from 60 to 40 cm. Vertical movement of materials, nutrient cycling and reuse driven by various burrowing organisms could have an effect on this nitrate-nitrogen distribution along the depth profile up to 40 cm. Less abundance of bioturbation below 40 cm could enhance the anoxic condition which in turn initiate denitrification causing sudden depletion of nitrate-nitrogen. The nitrate-nitrogen concentration showed no significant variation throughout depth but slight increased below 50 cm of depth which may be an indication of denitrification. Population of SRB was found to be increased with increasing depth. The Eh value of surface soil and soil from 60 cm of depth was found to be -78 mV and -163 mV respectively. Thus, more anoxic condition preferred the more population of SRB in the bottom soil than that of surface soil (Brune et al., 2000; Sass et al., 1997). Fungal population showed decreasing trend with increasing depth. Free living nitrogen fixing bacterial population that showed from surface to 30 cm depth increased again from 30 cm to 50 cm of depth. After the death of plant, the woods are carried carried away by tidal action or consumed by herbivorous animal but the root that remains attached to the bottom soil below the 50 cm depth seldom may act as the source of carbon to fungus and cellulose degrading bacteria (Figure 2a).

During monsoon, nitrate-nitrogen organic carbon content of soil showed decreasing pattern along with decrease in population of nitrifying bacteria with increase in depth. Silicate concentration showed little variation with increasing depth. Population of PSB was found to be decreased with increasing depth and at same time, phosphate-phosphorous concentration was also decreased with increase in depth. Population of CDB decreased with increase in depth as the organic carbon content of the soil was also decreased with increase in depth (Zemin et al., 2010). Population of SRB showed increase in trend as Eh value found in the surface and below 60 cm of depth were -83 mV and -169 mV (Figure 2b).

During postmonsoon, nitrate-nitrogen organic carbon content of soil showed decreasing pattern along with decrease in population of nitrifying bacteria with increase in depth. Silicate concentration showed little variation with increasing depth. Population of PSB was found to be decreasing with increasing depth and at same time phosphate-phosphorous concentration was also decreased with increasing depth. Free living nitrogen fixing bacteria showed decrease in population up to 30 cm of depth but below 30 cm to the next 60 cm of depth their population was increased. Population of CDB decreased with increase in depth as the organic carbon content of the soil was also decreased with increase in depth. Sulfate concentration did not show distinct stratification though population of SRB showed increase in trend as Eh value was found on surface and below 60 cm of depth were -87 mV and -198 mV (Figure 2c).

In the region of pneumatophores or rooted region during pre-monsoon, organic carbon content of soil showed decreasing pattern along with decrease in population of CDB from surface to 30 cm of depth and from 30 cm to the next 60 cm of depth, reverse image was found. Same profile was found for population of PSB and Phosphate-Phosphorous concentration. Silicate concentration showed little variation with increasing depth. Free living nitrogen fixing bacteria showed no graduation along the depth. Sulfate concentration did not show distinct stratification though population of SRB showed increase in trend as Eh value was found to surface and below 60 cm of depth were -89 mV and -198 mV (Figure 3a). Other nutrient and microbial population showed little graduation along the depth. This may be due to perforation of soil because of presence of pneumatophores (Figure 3a).

During monsoon season, sediment of rooted region showed more or less uniform distribution of nutrients along depth profile. During that season, effective tidal force becomes more active to mix up the nutrients vertically. Sulfate concentration increased suddenly at
**Figure 2.** Graphical representation of deep forest region that shows profile of nutrient concentration and CFU of microbes of different category along depth profile during pre monsoon (a), monsoon (b) and post monsoon (c) respectively.
Figure 3. Graphical representation of rooted region that shows profile of nutrient concentration and CFU of microbes of different category along depth profile during during pre monsoon (a), monsoon (b) and post monsoon (c) respectively.
about 40 cm and it may be due to sudden intrusion of inorganic sulfate from remote environment. Again C.F.U of SRB showed increase in their population from 30 to 50 cm of depth. Same type of trend was also found for population of fungus and CDB. It may be due to accumulation of such group of bacteria through the pores present in that zone (Figure 3b).

During postmonsoon, uniformity of nutrient concentration was found with increase in depth. Random change in silicate concentration was found along depth profile. Same time, fungal population, population of cellulose decomposing bacteria and population of nitrifying bacteria showed decreasing pattern with increasing depth. Free living nitrogen fixing bacteria showed more or less same population with increasing depth. Population of sulfate reducing bacteria was found to be increased with increase in depth (Figure 3c).

During pre-monsoon sulfate concentration was found to decrease with increasing depth followed by gradual increase of CFU of sulfate reducing bacteria. This may be due to more anoxic condition which is evident from sediment decreasing Eh value with increasing depth (Feng et al., 2003). Eh value was found maximum and minimum at the surface and 60 cm depth of un-rooted region with a value of -64.7 mV and -116.9 mV, respectively. Nitrite concentration did not vary with depth profile. Phosphate, nitrate, organic carbon content was found to show no proper gradation. Population of free living nitrogen fixing bacteria was found to show increase with depth. Other group of microbes showed no distinct gradation with increase in depth (Figure 4a).

During monsoon, organic carbon content and nitrate concentration showed a decreasing trend with increasing depth (Figure 4b). Monsoon played an important role in regulating the microbial population in Sundarban mangrove sediment. During post monsoon, no distinct gradation was found with respect to nutrient concentration and organic carbon content of soil along the depth profile. Microbial population showed same profile like nutrient concentration and organic carbon content of soil along the depth profile but only fungal population was found to decrease with increasing depth (Figure 4c).

Annual mean microbial population for rooted region (14.57±12.42 C.F.U ×10^6) was found relatively higher than un-rooted region (16.29±6.29 C.F.U ×10^5). The annual range of total microbial population fluctuated to maximum at the un-rooted exposed region (22.24±9.46 C.F.U ×10^6) and minimum in the deep forest region (15.21±6.14 C.F.U ×10^6). This indicates the occurrence of relatively stable condition over the deep forest region than that of the other two sites (Figure 5).

In the deep forest region, the most dominating group of microbe was found to be cellulose decomposing bacteria because that region contained more organic carbon content than that of the other two regions. Fungal population was less with respect to the other two regions. It may be attributed to less water content in deep forest region than that of other two regions. Population of SRB also showed more in proportion due to anoxic condition. In rooted region, fungal population was found to be the most dominant. CDB and SRB showed more or less same relative abundance. Population of PSB and free living nitrogen fixing bacteria was more in proportion than population of nitrifying bacteria. In un-rooted region, fungal population was found to be the most dominating group of microbe. The second highest was SRB. Un-rooted region was found to retain less organic carbon content for which population of CDB was less than population of fungus and SRB. Un-rooted region is inundated by water in most of the time and it increases water content of soil. More water content caused increase in fungal population. Anoxic condition also created ideal condition for much population of SRB (Figure 6).

Organic carbon content of the soil was found to be most significant on the growth rate of cellulose decomposing bacteria (Pearson correlation of Org.C (%) and C.D.B. (C.F.U ×10^6) = 0.500 P-Value = 0.000). The population of cellulose decomposing bacteria was found to be more in monsoon period than that of pre-monsoon and post monsoon. Again, the zone with more population of phosphate solubilising bacteria showed more concentration of available phosphate. Presence of phosphatase enzyme within such type of bacteria might be responsible for those findings (Cheng and Zhiping, 2007). It might be for availability of more organic carbon source. Rooted region showed a little stratification of nutrients along with microbial population with increasing depth. Rooted region makes the soil perforated for which during high tide nutrients and microbes present in sea water get mixed with soil vertically.

Again the little stratification in that region may be due to absorption of nutrients readily by the roots present in that region. Un-rooted region showed insignificant stratification of nutrient concentration and also for microbial population in those three seasons. That region experiences daily tidal action with great tidal wave and that high energy facilitates to mix the soil vertically (Cyr, 1998). Sulfate reducing bacteria was found to be correlated with sulfate concentration of soil sample (Pearson correlation of Sulfate-Sulfur (mg g^-1 dry wt of sediment) and S.R.B (C.F.U ×10^6) = 0.595 P-Value = 0.000). Phosphate solubilizing bacteria was also found to be correlated with phosphate concentration of the Sundarban mangrove soil (Pearson correlation of phosphate-phosphorous (µg g^-1 dry wt of sediment) and P.S.B (C.F.U ×10^6) = 0.766 P-Value = 0.000). No such correlation was found for nitrogen fixing bacteria with nitrate and nitrite concentration.

**Conclusion**

From the present study, the following conclusions have
Figure 4. Graphical representation of deep forest region that shows profile of nutrient concentration and CFU of microbes of different category along depth profile during pre monsoon (a), monsoon (b) and post monsoon respectively (c).
been drawn as a result of our research on depth integrated microbial diversity pattern of Sundarban Mangrove forest, along the shore of North East coast of Bay Of Bengal, India:

1) For deep forest region, a decrease in nutrient concentration was observed with increase in depth below 30 cm. Active bioturbation could result in vertical mixing of nutrients up to a depth of 30 cm below which effect of bioturbation become insignificant.

2) The overall concentration of organic C was found more in deep forest region than that of rooted and un-rooted
region. It might be attributed to undisturbed supply of mangrove litter which converts this zone to most suitable for microbial population.

3) Organic carbon content of the soil was found to be most significant on the population of cellulose decomposing bacteria (Pearson correlation of Organic C (%) and C.D.B C.F.U ($10^6$) = 0.500 P-Value = 0.000).

4) The zone with higher population of phosphate solubilising bacteria showed more concentration of available phosphate could be attributed to significant activity of phosphatase enzyme (Cheng and Zhiping, 2007). Phosphate solubilizing bacteria was also found to be correlated with phosphate concentration of the Sundarban mangrove soil (Pearson correlation of Phosphate-Phosphorous $\mu$g g$^{-1}$ dry wt of sediment and P.S.B C.F.U ($10^6$) = 0.766 P-Value = 0.000). No such correlation was found for nitrogen fixing bacteria with nitrate and nitrite concentration.

Organic carbon from the leaves, wood from forest and other organic dead or waste products from other living creatures are easily degraded by cellulose degrading bacteria in the mangrove sediment because they are the most dominating group of microbes prior to fungi. Other group of microbes also showed significant population which is a good sign for such mangrove forest with respect to mineralization of organic debris and as a result mangrove plants can easily get nutrient in simplest form. It can also be predicted that deep forest region is ecologically more stable than rooted region and un-rooted region. Sea level rise due to global warming may hamper the stable ecological zone of Sundarban Mangrove Forest which may ultimately reflect to net flux of several biologically active trace gases between soil and atmosphere.

Introduction of huge amount of nutrients during monsoon have a positive feedback on the bacterial population of mangrove sediment. Beside the changes in several physicochemical parameters, transport of huge amount of aquatic microbes could lead to the significant increase in the microbial population in the sediment of this mangrove ecosystem. This may contribute to the aquatic biogeochemistry of this tropical wetland.

REFERENCES


