Some Metabolic, Diversity and Toxicity Aspects of Bacterial Communities Life in Aquatic Sediments

Abstract

Sediment is a special realm in aquatic environments and has high microbial diversity. Addition of organic electron donors to sediments stimulates heterotrophic activity. Bacteria utilize the more available fractions of the organic detritus in their own metabolism, leaving indigestible residues known as marine humus. Input of nitrogen and carbon from vegetated regions into freshwater sediments also stimulate aerobic, anaerobic and chemolithotrophic sediment microbes. Because bacteria respond dramatically to changes in their chemical environment, can be predicted rapid responses by the aquatic sediment microbial community to any change in plant species that affects sediment porewater chemistry. Temperature and rainfall can affect the some microorganisms growth and permanence such as Escherichia coli and Enterococcus spp. The increase of pollutants load (e.g. Pb and Zn) in water can be expected with the reduction of flow, whereby the decrease of pathogens concentration can be due to high concentrations of metals (e.g. Hg and Pb) discharges in rivers. The enrichment of metals and nutrients concentration in sediments, especially in those with the fine texture, highlights that the adsorption process is the prevailing self-purification mechanism. Investigations of short-term variations of microbiological and chemical parameters in the water column have implied that, despite seasonal and nitrogen variations, microbial standing stock biomass and activity in one and the same body of water do not change drastically from one day to the other. Marine sediments contain as much as 18 per cent organic matter and bacteria in bottom deposits may grow and multiply. Enteric bacteria may be concentrated in bottom sediments of rivers as a result of sedimentation of bacteria-bound particles. Although it has been demonstrated that microbial communities can affect chemical pollutants, the presence of toxicants can also affect microbial community structure.

Keywords: Sediment; Bacteria; Water; Environment; Communities

Abbreviations: T-RFLP: Terminal Restriction Fragment Length Polymorphism; POM: Particular Organic Matter; DOM: Dissolved Organic Matter; ALAN: Artificial Light at Night; PCR: Polymerase Chain Reaction; PLFA: Phospholipid Fatty Acid; DTPA: Diethylenetriamine Pentaacetic Acid+TEA Buffered to pH 7.3

Introduction

Bacterial communities have a great importance in aquatic sediments and water. They are main degraders of organic and some inorganic compounds. Bacteria participate in cycling of nitrogen through nitrification and denitrification, as in cycling of some other elements. Microbial biomass may be significantly in relation to rate of C source consumption in all lake sediments, suggesting that microbial activity is limited by C availability. Findlay (2003) showed that the addition of different carbon sources, i.e., glucose, bovine serum albumin and natural leaf leachate to hyporheic biofilms enhanced microbial activities [1]. Carbon sources may influence nitrate reducing activity of the biofilm. Methanol is the most widely used carbon source for denitrification in reactors. Higher denitrification rates have been obtained with acetate as a carbon source. Also, abundance of denitrifying bacteria in the biofilm may be influenced by different carbon sources. This can be monitored by investigating the density of nosZ gene which codes for the nitrous oxide reductase, the last step of denitrification. The structural parameters of the biofilm formed under different carbon sources were examined and it was observed that the acetate and ethanol biofilms were mosaic-structured with the characteristic voids, whereas confluent biofilms were formed with glucose and methanol. The community structure of the biofilm established under different carbon sources were analysed by the DGGE fingerprinting method. According to the analysis, a 27% change in community occurred when the carbon source was changed from acetate to glucose, but 71.5% community change was observed from glucose to methanol transition [2].

Since gradients in microbial function may correspond to chemical gradients, microbial biodiversity may also respond to changes in sediment chemistry that are linked to above-sediment plants. Freshwater sediments may protect allochthonous bacteria (Escherichia coli, Enterococcus faecalis, Vibrio cholerae, V. parahaemolyticus, V. salmonicida, Pseudomonas fluorescens, Staphylococcus epidermidis) from variety of stressors and enteric bacteria are often found at higher concentrations in sediment...
than in overlying water column. The diversity and composition of bacterial communities have been shown to be affected by metal contamination of their habitat.

Therefore, toxic metals can be classified as a stressor that according to ecological theory should cause a shift in community composition toward lower diversity and metal tolerant bacterial strains. Bacterial tolerance to heavy metals is commonly coded by genes located on extra chromosomal elements (plasmids).

Discussion

Metabolic patterns

Addition of organic electron donors to sediments stimulated heterotrophic activity. Wang et al. [3] showed that addition of electron donors (glucose, sucrose and potato starch and sodium acetate) stimulated denitrification in Lake Taihu (China) sediments. In the study of benthic microbial response to the deposition of natural seston in Lake Erken (Sweden), Törnblom and Rydin (1998) showed that seston addition caused an immediate increase in bacterial production, activity and total sediment metabolism. Generally, sediments responded rapidly to addition of most electron donors by increasing CO₂ production rates. Sediments from site KR in Lake Okeechobee with the lowest microbial biomass exhibited the longest lag phase before responding to electron donor addition. Microbial biomass was significantly related to rate of C source consumption in all lake sediments, suggesting that microbial activity is limited by C availability. Although the magnitude of response to electron donor additions was related to microbial biomass, varied responses in sediments were probably related to catabolic diversity of microorganisms utilizing these substrates. Results of Principal Component Analysis 1 showed that Lake Apopka (Central Florida) sediments had the highest respiration per unit of microbial biomass with most of the electron donor additions (propionate, lake-SS, butyrate, acetate, and formate), indicating that these sediments respired most of the added C. These sediments support high rates of microbial activity, as indicated by rapid utilization of a wide range of electron donors. There were found that, after seston addition to sediment, bacterial biomass doubled indicating assimilation of C into microbial biomass. King and Klug (1982) reported that the addition of glucose into microbial biomass was low (20%) in a eutrophic lake sediment (Wintergreen Lake). Basal CH₄ production rates were highest in sediments of hypereutrophic Lake Apopka. Low rates of basal CH₄ production found in eutrophic Lake Okeechobee sediments may be explained by electron donor limitation, or by the presence of iron oxides which potentially can inhibit methanogenesis. Methanogens have the ability to use only a limited number of substrates, including H₂, CO₂ formate, acetate, methanol, and methylated amines. The most important substrates for methanogens are H₂/CO₂ and acetate, and they often depend on other anaerobic bacteria for these substrates. Most methanogenic species use H₂/CO₂ and a fewer number of species can use acetate. Circumneutral to alkaline sediment pH creating good conditions for hydrogenotrophic methanogenesis. Other studies have shown that CH₄ production at low temperature (4°C) in sediments was mainly from acetate, however, an increase in temperature (20-25°C) lead to an increase in contribution of CH₄ production from H₂/CO₂. Slightly higher temperature at the sediment-water surface (26.3-30.7°C) creates good conditions for hydrogenotrophic methanogenesis. Study has shown that sediments with different biogeochemical properties have microbial communities that exhibit distinct catabolic responses to a range of C-sources. The hypereutrophic lake with higher C availability in its sediments had the highest catabolic diversity, where the microbial communities were able to efficiently use a broader range of substrates.

The oligo-mesotrophic lake sediment microorganisms had lowest efficiency in use of energy [1]. The lack of water circulation in stratified water bodies (with a thermocline and halocline) in conjunction with continuous O₂-consumption by remineralization leads eventually to complete O₂ depletion. First nitrate and then sulphate is used for the degradation of organic materials in anoxic water. As a result, H₂S is then produced by dissimilatory reduction of sulphate by sulphate-reducing bacteria [4]. Bacteria change indigestible forms such as chitin or celluloses into more digestible bacterial protoplasm. Bacteria may serve as food for metazoa, and it is known that they are an important food for protozoa. The bacteria, however, also utilize the more available fractions of the organic detritus in their own metabolism, leaving indigestible residues known as marine humus [5].

Input of nitrogen and carbon from vegetated regions into freshwater sediments stimulate aerobic, anaerobic and chemolithotrophic sediment microbes. Since gradients in microbial function may correspond to chemical gradients, microbial biodiversity may also respond to changes in sediment chemistry that are linked to above-sediment plants. Bacteria are abundant both in water (more than 10⁶/ml) and in aquatic sediments (more than 10⁹/gdw), although not all bacteria are metabolically active at any one time. Specifically, aerobic metabolism, denitrification, metal reduction, sulfate reduction and methanogenesis all appear to vary over these spatial scales. These results suggest that a diverse assemblage of bacterial functional groups coexists in freshwater sediments. Work by Dahm has shown that the availability of specific forms of electron donors and electron acceptors in ground water affects microbial composition and dynamics in freshwater sediments. Under the appropriate conditions, one or more microbial functional groups may become active and dominant. Thus, functional diversity, and perhaps species diversity of freshwater sediment microbes must be very dynamic over time and space, and may respond rapidly to changes in water chemistry induced by changes in the overlying plant species composition [5].

Temperature and rainfall can affect the microorganism's

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growth and permanence such as *E. coli* and *Enterococcus spp*. The enrichment of metals and nutrients concentration in sediments, especially in those with the fine texture, highlights that the adsorption process is the prevailing self-purification mechanism. As well as the sediments represent a memory of aquatic ecosystem, their role of sink of pollutants and nutrients is correlated to fine size (silt and clay) and iron and manganese oxides. The *Clostridium* spores in sediments indicate a fecal contamination and their growth depends on the S and C content because they are sulphur-reductive bacteria involving in the sulphur-compound demobilization in anoxic environment. The high concentration of metals in sediments can compromise the life of the aquatic ecosystem, but the pollution risk from metals depends on their chemical speciation rather their total elemental contents. The extraction with EDTA and DTPA solutions reveals a different percentage of availability of the metals in sediments and the high amount extracted by EDTA is due to the acid pH, while the DTPA with neutral pH does not extract the metals immobilized in sediments. High percentage of Pb and Cu are extracted by both solutions highlighting a greater pollution risk than the other metals which are poorly extractable. Adsorption of pollutants in sediments seems to be the main self-purification mechanism while the low pathogenic contamination is related to severe drought during the spring-summer period that, lowering the river flow, will increase the concentration of pollutants in water. Sediment is the sink of nutrients and pollutants and their hazard can be evaluated by metals speciation and by their availability [6].

The profound sediment of deep lakes is characterized by constant temperature throughout the year. Addition of algae to the profundal sediment of a freshwater lake resulted in a systematic sequential change in both activity and composition of the active microbial community, which eventually led to methanogenic decomposition of the added algal organic matter. Increased metabolic activity in summer thus can be caused only by change in chemistry, for example, input of fresh organic matter by sedimentation of algal biomass. The change in chemistry is reflected, for example, in simulated sedimentation of *Peridinium gatunense* cells onto intact sediment cores, resulting in an enhanced CH4 production rate and in the formation of products of organic matter fermentation (propionate, acetate), especially within the upper sediment layers. Bacterial populations are involved in decomposition of the added algal biomass. Addition of algal biomass to the sediment core resulted in increased concentrations of acetate and in doubling of the CH4 production rate from within the upper sediment layers. Dissolved organic matters is probably converted by the concerted activity of fermenting bacteria, syntrophic H2-producing bacteria and methanogenic archaea, resulting in increase of acetate, propionate, H2, CO2 and CH4 concentrations immediately after addition of algal biomass. The quantity as well as quality of available organic material caused changes in the structure of the bacterial community. Analysis of the microbial community by fluorescence in situ hybridization and denaturing gradient gel electrophoreses showed that addition of algal biomass resulted in a fast and selective increase of distinct bacterial groups. In marine sediments, respiratory processes, sulfate reduction in particular, dominate the decomposition of organic matter. Here has been analyzed reverse-transcribed 16S rRNA on addition of algal biomass to sediment sampled. Although the data cannot be directly compared because of different sampling times and targeting rRNA versus rDNA, they are consistent with respect to the main features of the archaeal and bacterial communities. In particular, always observed a relatively high abundance of members of the *Deltaproteobacteria* and *Bacteroidetes* phylum, but also of *Chloroflexi*, *Planctomycetes*, *mmproteobacteria* and *Epsilonproteobacteria*. After 1 d of incubation, T-RFs allocated to *Desulfobacterales*, *trophobacterales*, and *Mycococcales* dominated the bacterial T-RFLP (terminal restriction fragment length polymorphism) profile of the algal amended slurries. Since sulfate was not detectable after 1 d of incubation, the high relative abundance of *Desulfobacterales* must be due to metabolism other than sulfate reduction. The detection of *Syntrophobacterales* is reasonable since members of this order, such as *Syntrophobacter wolinii* (X70906), may have contributed to the syntrophic degradation of propionate that transiently accumulated on addition of algae. After 6 d of incubation, the structure of the active bacterial population changed again with members of the *Bacteroidetes* phylum (formally known as *Cytophaga-Flavobacteria-Bacteroides* group) becoming dominant. The *Bacteroidetes* phylum has already been described as one of the most abundant bacterial groups in aquatic habitats including freshwater lakes sediments. Therefore can be assumed that *Bacteroidetes*-related bacteria were involved in the degradation of *P. gatunense* thecae which consist mainly of polysaccharides. *Bacteroidetes* related bacteria (designated as “*Cytophaga*”) became abundant in algal-amended marine sediments. Although the next relatives of the clone sequences retrieved gave no clue to a particular phenotype, it can be assumed that populations of *Bacteroidetes*-related bacteria were primarily involved in the degradation of POM (particular organic matter) and became abundant as soon as the more labile DOM (dissolved organic matter) had been consumed by *Clostridiales* and *Deltaproteobacteria*. This study has shown that the addition of algae to the profundal sediment of a freshwater lake resulted in a systematic sequential change in both activity and composition of the active microbial community, which eventually leaded to methanogenic decomposition of the added algal organic matter [7].

Between dissolved and particulate organic matter, mutual exchange processes can be expected as indicated by the inverse variation pattern of glucose and organic matter content of the sediments. Variations in the total number of microbial cells correspond to organic nutrients and microbial activity. However, the higher the number of cells, the lower the concentration and the turnover time of glucose. From the calculations of microbial uptake activity, it is understandable that variations of both actual uptake rate and turnover time of glucose correspond to variations of glucose and total monosaccharides. The physicochemical parameters obviously influence microbial activity indirectly by various relationships existing between salinity, temperature, or both and parameters of dissolved organic carbon, which in turn relate to microbial activity. Saprophytes do not correlate well with total number and biomass. The only pronounced
relationship exists between saprophytes and respiratory rate of glucose. Pigmented saprophytes (percentage of the total number of saprophytes) show interrelationships with the concentration of glucose and total monosaccharides and actual uptake rates of glucose. Investigations of short-term variations of microbiological and chemical parameters in the water column have implied that, despite seasonal and daily variations, microbial standing stock biomass and activity in one and the same body of water do not change drastically from one day to the other [8].

Artificial illumination near surface freshwaters has the potential to be a viable night-time energy source for microbial phototrophs, some of which can carry out photosynthesis at light levels that are near to those generated by artificial light at night (ALAN) [9]. Artificial light can most certainly be used for photosynthesis, as a photon from the sun may be no different from one emitted by a light bulb, they may only be different in intensity or colour. However, the rate at which photosynthesis occurs is dependent on the spectral emissions of the light source [10]. Phototrophic organisms are able to carry out photosynthesis at low artificial light levels (e.g. after five months of illumination). This could result in changes to the microbial community as conditions may favor taxa that benefit from nocturnal (low-level) light. These changes could, in turn, influence ecosystem-level processes such as carbon turnover. In general, community respiration and net ecosystem production incorporates biochemical pathways that make organic carbon molecules and energy available to microorganisms and exhibit a strongly positive relationship with temperature. Such changes may also vary temporally because day length and light levels (along with temperature) act as seasonal cues for many organisms in natural systems. There have been carried out field and laboratory experiments to examine changes to microbial community structure and carbon mineralization in sediments of an agricultural drainage system. Such a long-term approach, including multiple generations of most microbes, potentially provides an important insight into temporal change in community structure and associated process rates arising from light. In the current study has been observed increased abundance in some phototrophic taxa in sediments after five months of illumination. Phototrophs again showed an increase in response to ALAN, but this was much more pronounced in the pre-exposed site. After 1 year of artificial illumination in the lit site, this trend continued with the pretreated sediment no longer showing a clear seasonal response to temperature. Results from the pre-treated sediment community suggest that ALAN may be the sole trigger and that it may override other seasonal triggers such as temperature. Consequently, this results in a loss of temporal structure and may lead to a loss of diversity [9].

Diversity of bacterial communities in various types of sediments

Illumina tags may be used for the study of bacterial diversity. Wang et al. [11] determined millions of Illumina reads for a comparison of bacterial communities in freshwater, intertidal wetland and marine sediments along Pearl River, China, using a technically consistent approach. Statistical analysis (LEfSe) demonstrated that the freshwater sediment was enriched with

_Acidobacteria, Nitrospirae, Verrucomicrobia, Alphaproteobacteria, and Betaproteobacteria_. The biomass and taxon richness of microbes in sediment are much higher than those of the corresponding water bodies. They determined sediments along the Pearl River in south China and performed a detailed comparison of the bacterial diversity and indicator bacterial groups in each type of sediment. Even though there has been used the Illumina instrument with stringent quality controls to determine the tag sequences, there are still many biases introduced by PCR error, sequencing error, sequencing depth and the bioinformatics pipeline. As the most diverse and even community, the freshwater sediment had the highest number of indicator taxa distributed in a variety of lineages, of which _Acidobacteria, Nitrospira_, and _Verrucomicrobia_ were the three major indicators. Although the _Acidobacteria_ phylum accumulated in the freshwater sediment, not all subdivisions of the phylum were the same. Result and recent reports suggest that subdivision 6 (Gp6), Gp1, Gp3, Gp4, and Gp18 (from high to low confidence) are enriched in the freshwater environment while subdivisions Gp10, Gp22, Gp21, and Gp26 are more abundant in marine sediments. In addition to _Acidobacteria, Nitrospira_ is another strong indicator. _Proteobacteria_ was the most abundant and largest phylum in all three sediments, but its classes showed different tendencies. Within the phylum, _Alphaproteobacteria_ and _Betaproteobacteria_ were enriched in the freshwater sediment, _Gammaproteobacteria_ in marine sediments, _Epsilonproteobacteria_ in intertidal wetland, and _Deltaproteobacteria_ in all sediments. In comparison, the groups enriched in marine sediment were significantly enriched in intertidal environments. In addition, many chemolithoautotrophic microbes, e.g., _Chloroflexi, Bacillariophyta_ and _Ectothiorhodspiraceae_ (from the order of Chromatiales) were significantly enriched in intertidal environments. In addition, many chemolithoautotrophic bacteria from _Epsilonproteobacteria_ were also enriched in intertidal sediment, such as _Sulfurovum_ and _Arcobacter_ (_Campylobacter_). All of these results suggest that the intertidal sediment is highly productive and has various types of primary producers which contribute to the high nutrient concentration in the intertidal environment. Microbial communities from three different habitats showed obvious differences, which could not be fully explained by the limited factors determined in the present study. The freshwater sediment had the highest diversity, with _Acidobacteria, Nitrospira, Verrucomicrobia, Alphaproteobacteria_, and _Betaproteobacteria_ as indicators. The intertidal sediment had medium diversity, with many primary producers (such as _Chloroflexi, Bacillariophyta, Gammaproteobacteria, and Epsilonproteobacteria_) and eutrophic microbes (such as _Bacteroidetes, Firmicutes, and Actinomycetales_). The marine sediments had the lowest diversity and was enriched with _Gammaproteobacteria_ and some _Deltaproteobacteria_ orders, which were mainly involved with sulfate reduction under anaerobic conditions [11].

Molongoski et al. [12] examined anaerobic heterotrophic bacteria from freshwater Wintergreen Lake sediment. The most striking feature of the bacterial strains isolated from Wintergreen Lake sediments is the predominance of clostridia among the
organisms examined. The genus *Clostridium* accounts for 71.8% of the total isolates. Greater than 50% of the clostridial strains, in turn, were identified as *C. bifermentens* and *C. sporogenes* (47.7 and 17.8%, respectively). Although toxicity tests were not performed, *C. sporogenes* was nevertheless identifiable due to its characteristic “medusa-head” colonial morphology. A saccharolytic isolate tentatively identified as *C. butyricum* represented a minor portion of the clostridial isolates (6.3%). The largely proteolytic nature of the clostridial isolates obtained is likely due to the predominance of nitrogenous organic substances in Wintergreen Lake sediments. Matches and Liston (1974) examined the sediments of Puget Sound for the presence of clostridia. They reported anaerobic plate counts ranging from 0.73x10^4 to 23.5x10^4 cells/ml of sediment-water slurry. Approximately 30% of these organisms were determined to be clostridia, the three species isolated in greatest numbers being *C. perfringens*, *C. bifermentens* and *C. novyi*. Finne and Matches (1974) have recently reported the isolation of low-temperature growing clostridia from marine sediments that can grow at 5°C or less, well within the in situ temperatures of most marine and freshwater sediments. The production of characteristic clostridial fermentation end products from [U-14C] glucose and a U-13C-labeled amino acid mixture by natural sediment microbial communities provides additional presumptive evidence that clostridia, and possibly *C. bifermentens* and *C. sporogenes* in particular, are active in the degradation of organic substances in Wintergreen Lake sediments. *C. bifermentens* and *C. sporogenes* elaborate an array of characteristic volatile metabolites, including the distinctive valeric and caproic series of fatty acids. With the exception of a few species of anaerobic cocci (*Peptostreptococcus sp.* and a small number of *Bacteroides sp.* and *Eubacterium sp.*), the production of valeric and iso-valeric acids as fermentative end products is generally limited to the genus *Clostridium*. These results support the hypothesis that a resident clostridial community is actively metabolizing in Wintergreen Lake sediments, particularly during periods in the summer and early fall when sediment temperatures are most favorable for growth of the clostridia. The diverse modes of anaerobic energy-generating metabolism displayed by members of this genus, their demonstrated ability to grow at environmental temperatures and their ability to form dormant spores whenever environmental conditions become unsuitable for growth should afford the clostridia a definite advantage in organically enriched anoxic habitats such as lake sediments [12].

In addition to living organisms, clear sea water contains 0.2 to 2.0 ppm of both inorganic and organic particles in suspension. Since approximately half of this particulate material is inorganic and claylike it may be expected that bacteria will be adsorbed and that the particles will eventually settle and be deposited on the ocean floor. Attached organisms thus become removed from surface waters and accumulate in sediments. Marine sediments contain as much as 18 per cent organic matter and bacteria in bottom deposits may grow and multiply. Russell (1892) demonstrated that there were greater numbers of bacteria in marine sediments than in overlying waters, and Rubentschik reported that coliform bacteria were absorbed by muds from salt lakes. More recently, it was confirmed that at TraA, the major filament protein of the F-pilus, is essential for the development of highly organized, mature *E. coli* K-12 biofilms. The presence of TraA pili could support biofilm maturation even in the absence of flagella, type 1 fimbriae or Ag43, thus providing further confirmation that the latter are surface appendages that promote the initial adsorption of the cells to the solid surface, but are dispensable for later stages of biofilm development. Conjugative pili can accelerate initial adsorption depending on the nature of the abiotic surface and growth conditions but they are also specifically involved in biofilm maturation [13]. The adsorption capacity of muds varied and there were differences in the susceptibility of different bacteria to adsorption. Similar results were obtained by Walsman and Vartiovaara (1938) who demonstrated that the clay fraction of marine sediments adsorbed more bacteria than did silt or sand. Weiss (1951) reported that sea water flocculated and decreased the capacity of silt to adsorb bacteria [14]. Clay accumulation is so important to the total sediment specific surface area and accumulation of clays in intertidal sediments evidently requires biological incorporation [15]. Diérent and Guillerd (1940) and Nusbaum and Garver (1955) attributed reduced bacterial counts to adsorption and sedimentation of coliform bacteria in sea water. Large numbers of coliform bacteria were detected in marine sediments underlying the paths of two oil spills that discharged unchlorinated sewage (Rittenberg et al. 1958), and it appeared that sedimentation was responsible for the distribution observed [14]. Enteric bacteria may be concentrated in bottom sediments of rivers as a result of sedimentation of bacteria-bound particles. Auer and Niehaus (1993) showed that greater than 50% of fecal coliform in lake water was associated with small-sized particles in the range of 0.45-5.5μm. Such particle associations influence the sedimentation rate of bacteria generally by increasing the rate of loss compared to unattached cells. Freshwater sediments may protect allochthonous bacteria form variety of stressors and enteric bacteria are often found at higher concentrations in sediment than in overlying water column. Sediment-bound enteric bacteria may become re-suspended by variety of perturbations including dredging, storm events and ship traffic. For example, Grimes (1980) reported an increase in indicator bacteria densities of 4-50 times in waters below dredging operations as compared to upstream waters [16].

**Responses and adaptations of bacterial communities on toxic agents**

Although it has been demonstrated that microbial communities can affect chemical pollutants, the presence of toxicants can also affect microbial community structure. Chemicals can alter the community structure through selection of pollutant degraders or through acute toxicity to microorganisms. Readily biodegradable pollutants can increase population densities by promoting growth through providing carbon and energy to microorganisms in otherwise oligotrophic environments. The greater the oil contamination the more numerous were the hydrocarbon-degrading bacteria. Chronic exposure to petroleum hydrocarbons have been proposed as a partial explanation for variability in bacterial response to these chemicals. Perhaps of even more importance are the factors, such as type of bacteria present at the
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Bacterial communities react in various ways in response to toxic factors. Frostegård et al. [21] studied the effect of heavy metal pollution on the microbial community in the soil by performing a phospholipid fatty acid (PLFA) analysis. They found that addition of Cd, Pb, Zn and Ni induced similar changes in the PLFA pattern, while the effect of Cu addition on the PLFA pattern differed from the effects of the other metals. Positive correlations between metal contents and the percentages of bacteria tolerant to heavy metals have also been observed with sediments and in aquatic environments. The opposite result was observed by Dean-Ross and Mills (1989) in a study of the responses of planktonic, sediment and epilithic bacterial communities to heavy metal pollution. However, these authors suggested that the lack of correlation between metal concentration and resistance in the sediment bacterial community was due in part to the high pH of the river water, which reduced the toxic effect of the heavy metals present. In polluted soils, particularly soils contaminated with Cu, Zn, and Cd, the appearance of colonies was delayed compared with unpolluted soil. The data support the conclusions of Doelman and Haanstra (1986, 1989), however, who found that the slopes of the dose-response curves for Cd, Cr, Cu, Ni and Zn for urease and phosphatase activities obtained in soils polluted with these metals differed from the slopes obtained in unpolluted soils. To overcome this problem, Doelman and Haanstra suggested the use of an ecological dose range (defined as the dose range in which activity decreases from 90 to 10% of the undisturbed activity) instead of the IC

values used in ecotoxicological studies. IC

values were closely related (r = 0.909, P < 0.001). The levels of tolerance of the bacterial community to the different metals in soils polluted with low doses were in some cases even lower than the levels of tolerance in the unpolluted control. These results and those reported by Frostegård et al. [21] seem to support this assumption since (i) an increase in the tolerance of the bacterial community and changes in the PLFA pattern were found at similar metal concentrations and (ii) similar effects of heavy metal additions were deduced after the bacterial communities were grouped on the basis of the results of a PLFA pattern analysis and tolerance pattern measurements [21].

Nayar et al. [22] studied reactions of heterotrophic bacteria on heavy metals in mangrove sediment environment in Ponggol Estuary. This study revealed very high concentrations of heavy metals in suspended particulates and sediments compared to the levels in water. It is a mangrove ecosystem and the fact that mangrove mud binds metals explains the elevated heavy metal levels in particulates and sediments. Also, fine sediments have a larger surface area, which allows heavy metals and other contaminants to be adsorbed easily. In the aquatic and sediment environment, bacteria have been reported to have a wide range of metal detoxification or resistant mechanisms. Bacteria immobilize metals and nullify their toxicity by sequestering the metals to their extracellular polymer matrix, excluding or by precipitating the metals into an insoluble mineral matrix. The toxic effects of metals have been stated to be attenuated by the precipitation and complexing capacities of the bacteria. Other detoxification mechanisms in bacteria include the formation of an extracellular polysaccharide capsule that acts as a metal trap,

Dutka and Kwan (1982) compared the results of different methods of detecting toxicity to bacteria and found that each method had its own sensitivity pattern to toxicants. The relative toxicities of the compounds compared very well whether the acetate incorporation or the glucosidase activity method was used. With the acetate incorporation method, toxic compounds were inhibitory to microbial communities taken from above a sewage treatment plant outfall than to those taken from below. There are two possible explanations for this.

Much organic material is added to the stream with the treatment plant effluent, and this could cause chelation of the toxicants. Also, the organisms below the plant could be more resistant to perturbation due to selection since they are more continuously exposed to more toxic compounds than are those from above. This could be especially true of the sediment community since it is less mobile than the water column community [20].
the presence of metal-resistant membrane proteins, and plasmid encoded metal efflux. In the presence of sulphate or phosphate, bacterial assemblages have been reported to detoxify cadmium by forming insoluble cadmium–phosphate or cadmium-sulphide complexes, which are precipitated out of solution. Observations along similar lines were made of an increased incidence of zinc-tolerant bacteria, linking the zinc tolerance to multiple-metal resistance. These observations provide evidence that heterotrophic bacteria can adapt well when exposed to high heavy metal concentrations and make use of the dissolved organic carbon pool available from the disintegration of the less-tolerant organisms. This explains the enhanced rates of heterotrophic bacterial production and increased total bacterial biomass in treatments with high concentrations of metals. It was apparent that bacterial heterotrophs were not affected by the background concentrations of heavy metals that were biologically available from the sediments re-suspended by dredging [22].

Studies of the effect of sludge application on the microbial flora have focused on the discharge of pathogenic organisms into the environment and the changes in microbial mediated nutrient cycling. Even those studies that examined the response of the endogenous microbial community to accumulated heavy metals have assessed soil enzyme activities, rather than microbial community structure. The diversity and composition of bacterial communities have been shown to be affected by metal contamination of their habitat. Numerous reports are available which correlate the extent of metal pollution with bacterial resistance in soil, water and sediment. Therefore, toxic metals can be classified as a stressor that according to ecological theory should cause a shift in community composition toward lower diversity and metal tolerant bacterial strains. Bacterial tolerance to heavy metals is commonly coded by genes located on extra chromosomal elements (plasmids). Among natural bacterial communities the development of resistance to stressors could be greatly enhanced by the horizontal spread of resistance plasmids. Elucidation of the relationship that exists between bacterial plasmid patterns and the response of the community to pollutants may contribute toward a better understanding of the role of plasmids in adaptation to environmental stressors [23].

Conclusion

Bacterial communities in aquatic sediments have some specific properties since they live with lack of light, in low temperature and with variable amount of oxygen. But, their metabolic role remains the same as in above water because significant part in element cycling belongs to them. Bacteria may degrade almost all organic compounds during the time, including some inorganic, creating important conditions for life in their habitat. Various factors affecting their diversity and shifts during the circadian and season period.

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