Diversity of Community Soil DNA and Bacteria in Degraded and Undegraded Tropical Forest Soils of North-Eastern India as Measured by ERIC–PCR Fingerprints and 16S rDNA-DGGE Profiles

Sorokhaibam S. Singh¹, Michael Schloter², Subhash Chandra Tiwari^{3*} and Mamtaz S. Dkhar⁴

¹Department of Forestry, North Eastern Regional Institute of Science & Technology, Nirjuli, Arunachal Pradesh, INDIA
²Institute of Soil Ecology, GSF-National Research Centre for Environment & Health, Neuherberg, GERMANY
³Department of Forestry, Wildlife & Environmental Sciences, Guru Ghasidas Vishwavidyalaya, Bilaspur, Chhattisgarh, INDIA
⁴Department of Botany, North Eastern Hill University, Shillong 793 022, INDIA

ABSTRACT

The analysis of the denaturated gradient gel electrophoresis (DGGE) of the polymerase chain reaction (PCR) amplified 16S rDNA fingerprints demonstrated a high diversity of bacterial communities in the soils from the three study sites at the surface and subsurface soil layers. The undegraded site possessed maximum number of 16S rDNA fingerprints than the degraded and undegraded sites at both soil depths. The surface soil of the degraded site displayed minimum 16S rDNA fingerprints thereby suggesting the reduced bacterial diversity in this site as a consequence of the long term utilization of the soil through shifting cultivation in the past. The cluster analysis of the DGGE bands of 16S rDNA genes revealed a clear separation of the degraded and moderately degraded sites from that of the undegraded site in terms of bacterial genomic communities.

Keywords: Community Soil DNA, DNA fingerprinting, microbial diversity, soil degradation, Tropics

INTRODUCTION

Microorganisms possess a large size of diversity in terms of number and genetic makeup while constituting an essential part of the biological diversity on the earth. As in higher organisms, microorganisms have some values economic, agricultural, medicinal, food, fibre, industrial, ecological, etc. (Dilly and Munch 1998; Handelsman and Wackett 2002; Hunter-Cevera 1998; Robe et al 2003; Robert and Szakacs 1998; Rondon et al 1999: Roose-Amslag et al 2001, Lauber et al 2009). So, their understanding and maintenance is of equal importance to that of higher organisms. Despite vast size in number of expected species our knowledge on the number of described species of these organisms is less than five percent of the total expected species (DIVERSITAS; Groombridge1992). On the other hand these organisms are apparently depleting from the biosphere due to changes made by natural and manmade activities (Ahn et al 2009, Caracciolo et al 2011). Lodge et al (1996) described various threats to microbial diversity in tropical forests which include forest fragmentation, loss of hosts caused by logging and other human activities, air pollutants, fungicides, disturbances that alter microclimates and exposure to sunlight, global climatic changes, etc. Deforestation and intensification of agricultural practices have been the most important causes among various manmade activities for fragmentation of natural habitats leading to rapid decline in natural forests, subsequent soil erosion, accelerated loss of fertile top soil, nutrient depletion, lowered crop productivity, loss of rare flora and fauna including the microorganisms (Girvan et al 2003; Saxena and Ramakrishnan 1986; Singh 2002; Tiwari et al 2002). Understanding the changes in microbial diversity and their activities in functioning of the tropical forest ecosystems following perturbations is of crucial importance. This is because of the reason that microorganisms in soil play a major role in ecosystem functioning though the microbial estimates do not necessarily reflect the characteristics of microbial populations or processes under field conditions (Dilly and Munch 1998). Moreover, microbial diversity of soil is important to sustainable agriculture because microbes mediate many processes that support agricultural productions and even may indicate disturbances or beneficial effects of amendments or management strategies (Lupiwayi et al 1998; Sharma et al 1998).

Studying the microbial components of the soil systems to understand their role in ecosystem functioning require using both the traditional cultivation, biochemical and recently developed molecular techniques. As an outcome of efforts being made during the last few decades, to explore the complex microbial resources, a number of techniques have been developed based on nucleic acid (DNA and RNA) probes and improvements have been made over the existing methods for screening, isolation and characterization of

^{*} Corresponding author: sct_in@yahoo.com

microorganisms (Hill et al 2000; Ogram 2000; Robe et al 2003; Schloter et al 2000; Trosvik 1980; Volossiouk et al 1995; Yeates et al 1998). The use of DNA based techniques have been proved to be better by many folds than the traditional plate culture methods but a polyphasic approach comprising of both the techniques while studying microbial diversity and their role in soil ecosystems is a prerequisite (Crecchio et al 2004; Hugenholtz and Pace 1996; Roose-Amsleg et al 2001; Trosvik 1980).

The small-subunit ribosomal DNA gene (16S rDNA) which codes for 16S rRNA in prokaryotic ribosomes has been used as a useful bio-molecular marker and presently employed as routine technique for cultureindependent technique in microbial ecology. The sequence analysis of the gene, isolated and purified from many environmental samples, has shown this molecular probe as one of the most powerful tools in microbial diversity research, molecular evolution and phylogenetic classification of the living organisms (Zhou et al 1997, Sorensen et al 2005, Liebner et al 2008). The analysis of the 16S rDNA nucleotide sequence after isolation and purification from the community soil DNA with the help of polymerase chain reaction (PCR) can easily determine the identity and diversity of the bacterial communities in soil as compared to the traditional plate culture methods which is considered to underestimate culture irresponsive species (Dunbar et al 2000). Some of the reasons for use of 16S rDNA as a molecular marker in microbial diversity studies are due to the universal distribution of this gene in all the communities of the domain bacteria, structural and functional conservation and the size which allows for sufficient sequence divergence (Ludwig and Schliefer 1994; Goebel 1995). The 16S rDNA gene has regions which are highly conserved while other regions display considerable sequence variation even within closely related taxa (Gobel 1995; Santros and Ochman 2004). These conserved sequences have been used as a phylogenetic marker for classification of bacteria into different taxa. The inference of phylogenies based on the comparative sequence analysis of the 16S rDNA provides, from the deepest separation of the different branches of life to the genus or even species or strain level, and facilitates identification and classification of microorganisms with little effort (Ludwig and Schliefer 1994; Olsen et al 1986, 1994). In fact, categorizing of the 16S rDNA gene from the community DNA of environmental samples has become a popular alternative to characterise microbial communities because it avoids the limitations of cultivability and directly provide information on phylogenetic diversity (Zhou et al 1997). However, the cloning and sequencing strategies are rather cost, time and labour consuming and thus not suitable for monitoring a large number of samples, e.g., in studies on the succession of microbial communities during the growing season, or following shifts of microbial communities after perturbations (Heuer and Smalla 1997). Recently use of the denaturing gradient gel electrophoresis (DGGE) of the PCR amplified DNA fragments has become a new approach to study the structural diversity of microbial communities which overcomes the disadvantages in cloning and sequencing of the DNA fragments. DGGE was initially developed for use in the medical research for detection of point mutations (Fisher and Lerman 1983; Myers et al 1985, 1987) but it was introduced in the microbial ecology by Muyzer et al (1993).

Enterobacterial repetitive intergenic consensus (ERIC) sequences are short interspersed repetitive elements found in the genome of eubacteria (Gillings and Holley 1997). ERIC elements are 126 bp in size and are distributed throughout extragenic regions of the genomes of many gram negative enteric bacteria and closely related phyla including vibrios (Giovanni et al 1999a; Hulton et al 1991; Versalovic et al 1991; Wilson and Sharp 2006). The unique locations of ERIC elements in bacterial genomes allows discrimination at genus, species, and even strain level based on the electrophoretic pattern of amplification products (de Bruijn 1992). PCR amplified ERIC elements generates amplicons of varying sizes ranging from approximately 50 to 3000 bp which collectively constitute a DNA fingerprint (Giovanni et al 1999b). Selective amplification of ERIC elements using oligonucleotide primers and comparative study of electrophoretic patterns of the fingerprints are used for identification, discrimination and classification of bacterial strains or communities (Bhattacharya et al 2003; de Bruijn 1992; Giovanni et al 199b; Niemann et al 1999).

Application of these techniques had been successfully employed in diverse microbial habitats on the biosphere ranging from agricultural and Siberian tundra soils in the terrestrial ecosystems towards marine sediments and hot springs in aquatic ecosystems (Cifuentes et al 2000; Dunbar et al 2000; Ennahar et al 2003; Hobel et al 2005; Zhou et al 1997). There has been numerous reports on successful use of PCR based DGGE profiles of 16S rDNA as a technique in studying described and undescribed microbial diversity in agricultural soil (Ovreas et al 1998; Girvan et al 2003), grassland soil (McCaig et al 2001), rhizosphere soil (Duineveld et al 2001; Heuer and Samalla 1997), marine sediments (Ogram et al 1987), polluted soil (Kirk et al 2004; Maila et al 2005), water biofilms (Lyauley et al 2005), pine forest soil (Laverman et al 2005), etc. Similarly,

ERIC-PCR has been used generally to identify and discriminate pathogenic bacterial strains or mixed bacterial communities and including the rhizosphere bacterial communities of genetically modified organisms (Finger et al 2006; Hulton et al 1991; Giovanni et al 1999a, 199b). However, irrespective of the type of methods employed, remarkably very little has been published on the numerous and often crucial ways in which microorganisms (fungi and bacteria) influence tropical forest ecosystems (Lodge et al 1996). Moreover, reports are scarce on application of such molecular based techniques (16S rDNA DGGE and ERIC- PCR) in studying the microbial communities in sub-tropical humid forest soils where the practices of shifting cultivation is a prominent system which leads to large scale environmental degradation problems of north-eastern region. Therefore, the present study was made to investigate the impact of soil degradation due to shifting cultivation and selective logging on distribution of community soil DNA and bacterial diversity by using ERIC-PCR fingerprints and 16S rDNA-DGGE profiles.

MATERIALS AND METHODS

Study site description

Banderdewa forest reserve in the Papum-pare District of Arunachal Pradesh (north-eastern India) was the area of the reported study. The area is located between 27°6' N latitude and 93°49' E longitude at an elevation of 350m amsl. The average annual (maximum) ambient temperature and average total annual rainfall were 26°C and 2609 mm. The vegetation consists of evergreen to semi-evergreen mixed natural forests of varying tree species ranging from the primitive Magnolia to newly introduced teak plantations.

Two study sites, namely degraded forest (DF) and moderately degraded forest (MDF) sites were selected based on the level of disturbance due to agricultural practice (shifting cultivation or "jhum") and selective logging. A protected tract in the reserve forest was selected as an undegraded natural forest (UDF) as control site in this study. The various criteria for classification of the two sites includes land use history, living tree stump density, dominant tree species composition, exposure of soil surface, thickness of soil profile (Ah) and important physico-chemical properties of soil (tables 1 and 2).

Parameters	DF	MDF	UDF
Soil texture (%) Sand	72.8	73.0	74.9
Silt	20.1	18.0	18.1
Clay	7.1	9.0	7.0
Textural Class	Loamy sand	Loamy sand	Loamy Sand
Soil profile thickness (A _h , cm) Tree density (trees/100 m ²) Porosity at surface layer (%)	3-5 12 49	7-10 84 52	13-17 121 55
Land use history (years)*	20	0	0
Erosion class [#]	3	1	0
Extent of degradation ^{\$}	Dominant (.5)	Common (.3)	0

*Agriculture practices

[#]Soil Survey Manual: Handbook No. 18 (USDA, Revised Edition, 1995)

Sehgal and Abrol (1994)

Parameters	DF		MDF		UDF	
	S	SS	S	SS	S	SS
Bulk density (g/cm ³)	1.3±0.06	1.34±0.07	1.1±0.09	1.2±0.05	1.0±0.07	1.12±0.05
Moisture content (%)	19.7±1.2	16.6±0.38	20.9±0.98	17.8±1.3	22.7±0.34	20.5±1.6
pH	5.5±0.38	5.2±0.21	5.7±0.23	5.3±0.21	6±0.45	5.7±0.44
Organic C (%)	1.7 ± 0.51	1.2±0.42	2.4±0.41	1.5±0.46	3.1±0.45	2±0.44
Total N (%)	0.22±0.14	0.17±0.13	0.33±0.18	0.21±0.14	0.43±0.20	0.29±0.15

Table 2. Physico-chemical characteristics of soil in DF, MDF and UDF sites at surface (S) and sub-surface (SS) soil layers

(Values are means of triplicate analysis with standard deviation)

DF was a nine years old regenerating jhum fallow when the research was conducted in 2002. This site had been used for cultivation of rice, maize, finger millet and tuber crops etc. for about fifteen years without proper input of nutrients based on either organic or inorganic fertilizers. Thus, the nutrients available to the crops were obtained only from the debris of plant residues and ash after burning of dried slash prior to cropping in each jhum cycle. This site has not been used for further cultivation of crops since last jhum practiced in 1993 due to lower crop productivity and was lying as an abandoned regenerating fallow land. Luxuriant growth of fern, weeds and grasses covers the soil surface during summer rainy season while dried plant residues cover the soil surface in winter season. MDF was a forest tract where removal of mature and large sized timber trees and introduction of new plantations i.e. selective logging were practiced by forest personnel as routine silvicultural operations. Annual cutting and removal of trees, cleaning, burning of dried grasses and weeds on forest floor are the main causes of soil disturbance in this site. However, no sign of agricultural practice, either in the form of jhum or other practice, was observed form this site. The soil in the study site falls under Karsingsa series, which is a member of mixed loamy sand of hyperthermic family, typic Haplustalfs i.e. class of Alfisol (Singh 1999). The geology of the soil consists of sedimentary (sandstone) parent rock which is drained by small tributaries of Dikrong River towards Brahmaputra River in Assam.

Soil sampling and laboratory analysis

Soil sampling was done from triplicate plots of 50 m2 each from all the three sites in the middle of April 2002. Soil samples were collected separately using metal soil corer having a diameter and height of 6cm and 20cm respectively from two soil depths i.e. surface (0-20 cm) and subsurface (20-40 cm.) layers after discarding approximately 0.5 cm of soil in between the two depths. A total of 15 soil core samples were collected from each triplicate plots of DF, MDF and UDF sites and were mixed to obtain a composite sample for each site. The field moist soil collected from each study site was processed and passed through a 2mm mesh screen for further analyses. The physico-chemical properties of soil were determined on field fresh soil or after air drying of the soil samples (Okalebo et al 1993).

Isolation of community DNA from Soil

Isolation of community soil DNA was done using the Fast DNA[®] SPIN Kit for Soil (BIO 101). Approx. 0.5 g of fresh soil (stored at -20°C) was taken in a 2 ml E-tube containing lysing mixture. 978 μ l SPB (sodium phosphate buffer) and 122 μ l MT buffer were added in the tube and homogenized at maximum speed for 1 minute. The suspension was centrifuged for 1 minute at 14000x g and the supernatant was transferred into a clear 2 ml tube and 250 μ l PPS (protein precipitation solution) was added followed by mixing the tube for 2 minutes. Then the tube was centrifuged again for 5 minutes at 14000x g and the supernatant was transferred into a 15 ml tube. 1 ml of binding matrix suspension was added and the tube was turned upside down for at least 5 times to allow binding of DNA to the matrix. About 500 μ l of the supernatant at the surface layer of the tube was discarded and resuspended the remaining supernatant in the binding matrix. The slurry was transferred into a Spin Tube in two aliquots of 500 μ l of SEWS-M (salt/ethanol wash solution, DNase-free) at 14000 x g for 1 minute and the filtrate was discarded. The Spin was removed from the tube and dried for 5 minutes at room temperature. The Spin was replaced into a fresh catch tube and 50 μ l of DNA eluting solution (DES, Dnase /Pyrogen free water)

J. BIOL. ENVIRON. SCI., 2011, 5(15), 183-194

was added while gently stirring the filter membrane with the pipette tip. Then, the tube was centrifuged at 14000 x g for 1 minute to elute the DNA into the catch tube. The DNA content of the extract was checked at 1% agarose gel.

Enterobacterial repetitive intergenic consensus (ERIC)-PCR

The DNA sample was amplified using ERIC-I universal primer (5'ATGTAAGCTCCTGGGGATTCAC-3'; Versalovic et al 1991). The PCR reaction mixture of 50µl contained 5µl 1x PCR buffer, 2.5µl of 2.5µM MgCl₂, 5µl of 3% bovaine serum albumin (BSA), 5µl 0.2 mM dNTP, 2µl of ERIC-I universal primer, 29 µl RNase free water, 1µl DNA and 0.5 µl Taq DNA polymerase (Invitrogen). The PCR was done in two reactions, first with 10 min hot start at 95°C, pause at 80°C and add Taq DNA polymerase. Second reaction starts with denaturation at 94°C for 1 min, annealing at 53°C for 1 min, elongation at 65°C for 2 min for a total of 30 cycles followed by final stabilisation of the products at 65°C for 10 minutes. The PCR product was electrophoresed in 1% agarose gel to check the presence of DNA amplicons followed by further electrophoresis of the ERIC-PCR products on a polyacrylamide gel electrophoresis (PAGE) for analysis of microbial community soil DNA

16S rDNA amplification for DGGE

Denaturing Gradient Gel Electrophoresis (DGGE)

DGGE was performed by using 6% acrylamide gel (ratio of acrylamide to bisacrylamide, 37:1, Bio-Rad) with a 45 to 65% denaturing gradients {the 100 % denaturing solution comprised of 15 ml of 40 % PAA dissolved in 40 ml of formamide with 42 g of urea (Sigma) and 2 ml of 50x TAE buffer and the final volume maintained at 100 ml with milli- Q water}. 15 μ l of the purified 16Sr DNA was loaded in the gel using one-third volume of the DGGE buffer dye in each lane for the six different samples. The gel was electrophoresed at 60°C for 17 h at a constant voltage (70V) by using the universal mutation detector system (Bio-Rad). The gel was fixed in 25% glacial acetic acid solution for 30 min. followed by three times washing for 2 min. each with milli-Q water. Then, the gel was stained in silver nitrate solution (AgNO3, SIGMA) for 25 min, washed twice with milli-Q water for 30 sec. and developed the images in sodium thiosulphate+sodium carbonate solution. The gels were then put in Na-EDTA solution for 10 min to stop darkening of the gel. The gel was dried at 45 to 50°C for 48 hrs in dark and the images were captured using HP Scanjet II scanner.

Gel compare analysis

The DGGE images were processed for removal of background colors and for sharper bands before analysis using Gel Compare-II software (Applied Maths, Belgium).

RESULTS

ERIC profiles of community soil DNA

The community soil DNA profiles of the soil samples from surface and subsurface soil layers of the three study sites as revealed by the ERIC-PCR are depicted in figure 1. The fingerprint of the ERIC elements shows the presence of a variety of community soil DNA derived from various organisms predominantly microorganisms in these soils. The surface soil layer (DF) of the degraded site showed minimum number of ERIC bands as compared to other samples. The surface (MDF and UDF) and subsurface (mdf and udf) soil layers of moderately degraded and undegraded forest sites displayed more dense and higher number of ERIC bands than other soil samples. Gel compare analysis (Dice similarity coefficient-UPGMA based dendrogram) of the community soil DNA from ERIC-PCR fingerprints resulted in two groups of genomic diversity (figure 3A). Group I consists of the repetitive intergenic consensus sequences from surface (MDF and UDF) and subsurface (mdf and udf) soil layers of the degraded site. Group II consists of the community soil DNA from surface (MDF and UDF) and subsurface (mdf and udf) soil layers of moderately degraded and undegraded sites respectively. The surface layers formed the subgroup IIa while the the ERIC bands from subsurface soil layers formed subgroup IIb. There was a subdivision of the two soil layers into subgroups, IIa and IIb respectively. Therefore, there was a clear discrimination between the soil layers layers of moderately degraded and undegraded sites and between degraded site and other two sites in terms of community soil DNA composition represented by ERIC-PCR fingerprints.



Figure 1. Community soil DNA profiles of six soil samples on PAGE after ERIC-PCR. Surface soil layer (DF) and subsurface soil layer (df) of degraded site; surface soil layer (MDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (udf) of undegraded site. M=1.3 kb DNA ladder

DGGE profiles of 16S rDNA

The PCR products of 16S rDNA from the surface and subsurface soil layers of the three study sites produced a clear band of 473 bp against a 1kb ladder as shown in figure 2(A). The purified product of this PCR amplicon was used for separation of the different 16S rDNA nucleotide sequences by DGGE.

J. BIOL. ENVIRON. SCI., 2011, 5(15), 183-194

The DGGE profiles of the 16S rDNA sequences of the six soil samples are shown in figure 2(B). Gel compare analysis of the 16S rDNA sequences in all the soil samples showed clear variation among the soil samples of the three study sites (figure 3B). The dendrogram of the 16S rDNA-DGGE bands resulted in three different clusters. The group I consist of 16S rDNA bands from surface layer (DF) of degraded site. The cluster II is formed by surface (UDF) and subsurface (udf) layers of undegraded forest site. Similarly, the cluster III is formed by three soil samples and subdivided into two subgroups, IIIa and IIIb respectively. The subgroup IIIa consists of only 16S rDNA bands derived from the subsurface (df) soil layer of degraded site. There was a 91% similarity index of the banding patterns (16S rDNA bands) between the cluster II and III and 86% among the three cluster groups in this analysis.



Figure 2. (A) 16S rDNA band (473bp) on 1% agarose gel. (B) DGGE profiles of 16S rRNA bands. Surface soil layer (DF) and subsurface soil layer (df) of degraded site; surface soil layer (MDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (udf) of undegraded site.



Figure 3. (A) Dendrogram of ERIC-PCR amplified DNA bands (Dice similarity coefficient- UPGMA). (B) Dendrogram of 16S rDNA bands. Surface soil layer (DF) and subsurface soil layer (df) of degraded site; surface soil layer (MDF) and subsurface soil layer (mdf) of moderately degraded site; surface soil layer (UDF) and subsurface soil layer (udf) of undegraded site. M = DNA ladder (1.3 Kbp)

DISCUSSION

Diversity of community soil DNA

Generally, ERIC bands are expected to be generated from microbial genomes, particularly gram negative enteric bacteria and closely related phyla (Giovanni et al 1999a; Hulton et al 1991; Versalovic et al 1991). However, presence of the ERIC elements only in the bacterial genome had been disagreed since ERIC primers sometimes acts as RAPD primers to amplify any DNA under low stringency PCR conditions (Gillings and Holley 1997; personal communication with G.D. Di Giovanni). Generally, soil is a heterogenous matrix of various genomes comprising of DNA derived from plant, animal and predominantly microorganisms, the ERIC elements obtained in the present study were considered to be total community soil DNA with more emphasis on bacterial sources.

Use of Dice similarity coefficient index takes into account of the band position together with more weight on matching bands when the data was analysed for discrimination of variables to form groups with highly similar fingerprints in such a way that the fingerprints in different groups are as much as dissimilar as possible (Rademaker and de Bruijn, 1997). The results from the gel compare analysis of ERIC-PCR products clearly indicated that the soil in the degraded forest site is quite less in composition in terms of total community soil DNA bands as compared to moderately degraded and undegraded forest sites. The degraded site with comparatively less number of ERIC bands are separated from moderately degraded and undegraded forest sites with more number of ERIC bands. The surface and subsurface soil layers of degraded site in cluster I revealed presence of approximately less number of microbial genomes and less diverse groups in terms of community soil DNA contents represented by ERIC elements, However, the separation of the moderately degraded and undegraded forest sites in a separate cluster (group IIa & IIb) suggests that more number of ERIC bands and similar diversity of community soil DNA existing in these two sites despite clear difference between the surface and subsurface soil layers. It can be mentioned here that the degraded site in the present study was a regenerating jhum fallow land (shifting cultivated fallow). This site had been used for about 15 years for jhum during which the soil had been disturbed with repeated burning, tillage and cropping. Moreover, this site being a hill slope, the top soil has been lost considerably during the course of cultivation in addition to runoff losses through heavy rainfall during summer seasons in the region. The surface layer of the soil shows very thin Ah horizon and characterised by increased bulk density. Vazquez et al (1993) have reported suppresses growth of aerobic bacterial population due to compaction of soil following forest burning of an Atlantic soil as compared to unburnt soil. Changes in physiological groups and even complete sterilization of microorganisms have been reported after forest wild fire in an Atlantic soil and soil heating in a pine forest stand. The degraded site supports no more favourable soil environment conditions for growth and multiplication microorganisms in absence of required moisture and nutrients in the form of humus or organic matter as revealed by reduced organic carbon contents (table 2). These enditions might have caused decline in community soil DNA population and diversity in this site as compared to undegraded forest site where no soil disturbance has occurred.

The moderately degraded forest site with no cultivation history in the past was disturbed by selective logging of forest timber and routine silviculture practices. There was removal of ground vegetal cover through cleaning and subsequent burning of the dried biomass on the forest floor for every year or alternate year by forest personnel. These practices must have a profound impact on the distribution of community DNA in the soil resulting in an altered diversity status as compared to the subsurface layer where impact of fire and other disturbance are less. The undegraded forest site on the other hand had a similar composition of total genomic composition to the moderately degraded site at both the surface and subsurface soil layers. This suggests that selective logging has lower impact on the distribution of soil community DNA as compared to shifting cultivation or jhum. Decline in population and diversity of bacteria and fungi as measured by cultivation and culture independent methods have been reported from degraded forest soils (Singh 2002).

Bacterial diversity as revealed by 16S rDNA-DGGE profiles

The 16S rDNA sequence pattern of the different soil samples showed different profiles of bacterial genomic diversity. Since, the DNA sequence of the 16S rDNA is derived from the bacterial communities using specific primer, the DNA bands are expected to represent the available bacterial communities in the soil samples. The gel compare analysis uses the number of DNA bands available in a particular lane which represents a soil sample in the gel image. The cluster analysis is based on the number and similarity of bands in each lane with respect to other lanes. Therefore, the resulting classifications of the clusters are the indices of similarity or dissimilarity of the bacterial genomic compositions among the samples. The similarity index of the cluster analysis reveals that 91% of the total genomic diversity is same or 9% dissimilarity among degraded, moderately degraded and undegraded forest sites at both the surface and subsurface layers. However, the surface and the subsurface soil layers of the undegraded site had the maximum similarity index of 96% or the least dissimilarity value of only 4%. This reveals that the bacterial communities between the two soil samples were more similar than the rest of the samples in terms of genetic makeup. This suggests that the microbial communities in the undegraded site possess intact natural population and higher diversity while degraded and moderately degraded sites supports altered microbial communities in terms of 16S rDNA profiles following shifting cultivation and selective logging practices. The surface soil layer of the degraded site displayed minimum 16S rDNA bands in comparison to the other soils thereby revealing reduced microbial composition as a result of consistent jhum cultivation in the past. The lower percentage of similarity index (86%) or higher value of dissimilarity index (14%) in the number of bands and banding patterns of soil samples in this site than other soil are direct influence of lowered microbial population as well as less diversity of bacterial community inhabiting in this disturbed site. These results are in correspondence with the lower bacterial population of the degraded site determined by cultivation method in this site (Singh 2002). Clear variation in bacterial count, physiological function and molecular diversity (16S rDNA-DGGE) have also been reported from intensively managed horticultural site and another recently established organically farmed soils in southern Italy (Crecchio et al 2004).

The surface and subsurface soil layers of the moderately degraded site and subsurface layer of degraded site were clustered in one group revealing that these soils contained similar bacterial communities different from the other soil samples. It may be mentioned that the surface soil layer of moderately degraded site has also been disturbed for every year or for alternate year by clear-cutting of ground vegetation and burning of the forest floor biomass for proper growth of the new plantations. This practice must have some detrimental impact on the bacterial communities at the surface soil layers though the impact was comparatively lower at the subsurface soil layers, hence less disturbance in the bacterial community structure. Lundgren (1992) also reported a strong and long term detrimental effect of clear cutting of forest on the bacterial population of Ao horizon as compared to an undisturbed natural forest stand.

Therefore, a conclusion may be drawn that shifting cultivation or jhum practice, as a dominant agricultural practice in the north-eastern hill forest soils, has a long term detrimental impact on the distribution of community soil DNA and bacterial genomic diversity as revealed by ERIC-PCR and 16S rDNA-DGGE profiles. The cluster analysis of the DGGE bands of 16S rDNA fragments displayed a clear separation of the degraded and moderately degraded forest sites from that of the undegraded site in terms of community soil DNA content and bacterial diversity. It is clear from the present study that shifting cultivation causes a severe impact than selective logging practices on the distribution of total soil DNA content as well as bacterial genomic diversity in the long term. The use of ERIC-PCR and 16S rDNA-DGGE as culture independent techniques can be successfully applied to the assessment of impact of environmental degradation on microbial communities in soil.

ACKNOWLEDGEMENTS

Dr. S. Sureshkumar Singh is thankful to the German Academic Exchange Service (DAAD), Bonn for awarding Short Research Fellowship and the Council of Scientific & Industrial Research (CSIR), New Delhi for awarding Senior Research Fellowship during the work. The authors also gratefully acknowledge Dr. Shilpi Sharma and Dr. Vibiane Radl, Institute of Soil Ecology for their technical support during the work.

REFERENCES

Acea M J and Caraballas T 1996 Changes in physiological groups of microorganisms in soil following wild fire; FEMS Microbiol. Ecol. 20 33-39.

Acea M J and Caraballas T 1999 Microbiological fluctuations after soil heating and organic amendment. Bires. Tech. 67 65-71.

- Ahn C W, Gillevet P M Sikaroodi M and Wolf K L 2009 An assessment of soil bacterial community structure and physiochemistry in two microtopographic locations of a palustrine forested wetland. Wetland Ecology and Management 17(4) 397-407.
- Bhattacharya D, Sarma P M, Krishna S, Mishra S and Lal B 2003 evaluation of genetic diversity among Pseudomonas citronellolis isolated from oily sludge-contaminated sites; Appl. Environ. Microbiol. 69 1435-1441
- Caracciolo A M, Grenni P, Falconi F, Caputo M C, Ancona V and Uricchio V F 2011 Pharmaceutical waste disposal; assessment of its effect on bacterial communities in soil and ground water. Chem Ecol 27(S1) 43-51.
- Cifuentes A, Anton J, Benlloch S, Donnelly A, Herbert R A and Rodrigues-Valera F 2000 Prokaryotic diversity in Zostera nolti-colonized sediments; Appl. Environ. Microbiol. 66 1715-1719
- Crecchio C, Gelsomino A, Ambrosoli R, Minatti J L and Ruggiero P 2004 Functional and molecular responses of soil microbial communities under differing soil management practices; Soil Biol. Biochem. 36 1873-1883
- de Bruijn F J 1992 Use of repetitive (repetitive extragenic plindromic and enterobacterial repetitive intergenic consensus) sequences and polymerase chain reaction fingerprint of the genomes of Rhizobium meliloti isolates and other soil bacteria; Appl. Environ. Microbiol. 58 2180-2187
- Dilly O and Munch J C 1998 Ratio between microbial estimates of microbial biomass contents and microbial activity in soils; Biol. Fert. Soils 37 374-379.

DIVERSITAS Programme on Biodiversity IUBS-SCOPE-UNESCO

- Dunbar J, Ticknor L O and Kuske C R 2000 Assessment of microbial diversity in our southwestern United States soils by 16S rRNA gene terminal restriction fragment analysis. App. Environ. Microbiol. 66 2943-2950.
- Duineveld B M, Kowalchuk G A, Keizer A, van Elas J D and van Veen JA 2001 Analysis of bacterial communities in the rhizosphere of Chrysanthemum via denaturing gradient gel electrophoresis of PCR-amplified 16S rRNA as well as DNA fragments coding for 16S rRNA; Appl. Environ. Microbiol. 67 172-178.

J. BIOL. ENVIRON. SCI., 2011, 5(15), 183-194

Engelen B, Meinken K, von Wintzingerode F, Heuer H, Malkomes H P and Backhaus H 1998 Monitoring impact of a pesticide treatment on bacterial soil communities by metabolic and genetic fingerprinting in addition to conventional testing procedures; Appl. Environ. Microbiol. 64 2814-2821

Ennahar S, Cai Y and Fujita Y 2003 Phylogenetic diversity of lactic acid bacteria associated with paddy rice silage as determined by16S ribosomal DNA analysis; Appl. Environ. Microbiol. 69 444-451

Fisher S G and Lerman L S 1983 DNA fragments differing by single base pair substitutions are separated by denaturing gradient gel electrophoresis; Proc. Nat. Acad. Sci. USA 80 1579-1583

Goebel, U.B. 1995. Phylogenetic amplification of the detection of uncultured bacteria and the analysis of complex microbiota. J. Microbiol. Methods. 23 117-128

Gillings M and Holley M 1997 Repetitive element PCR fingerprinting (rep-PCR) using enterobacterial repetitive intergenic consensus (ERIC) primers is not necessarily directed at ERIC elements; Lett. Appl. Microbiol. 25 17-21

Giovanni G D Di, Watrud L S, Seidler R J and Widmer F 1999a Fingerprinting of mixed bacterial strains and BIOLOG gram-negative (GN) substrate communities by enterobacterial repetitive intergenic consensus sequence-PCR (ERIC-PCR); Curr. Microbiol. 38 217

Giovanni G D Di, Watrud L S, Seidler R J and Widmer F 1999b Comparision of parental and transgenic Alfalfa rhizosphere bacterial communities using Biolog GN metabolic fingerprinting and enterobacterial repetitive intergenic consensus sequence-PCR (ERIC-PCR); Microbial Ecol. 37 129-139

Girvan M S, Bullimore J, Pretty J N, Osborn A M and Ball A S 2003 Soil type is the primary determinant of the composition of the total and active bacterial communities in arable soils; Appl. Environ. Microbiol. 69 1800-1809

Groombridge B 1992 Global Biodiversity: Status of the Earth's Living Recources; World Conservation Monitoring Centre Staff, Chapman & Hall, London, UK.

Handelsman J and Wackett L P 2002 Ecology and industrial microbiology: Microbial diversity-sustaining the Earth and industry; Curr. Opinion Microbiol. 5 237-239

Heuer H and Smalla K 1997 Application of denaturing gradient gel electrophoresis and temperature gradient gel electrophoresis for studying soil microbial communities; in Modern Soil Microbiology (eds) J D Elas, J T Trevors and E M H Wellington (Marcel Deker, Inc.) pp 353-373

Hill G T, Mitowski N A, Aldrich-Wolfe L, Emele L R, Jurkone D D, Ficke A, Maldonando-Ramierez S, Lynch S T and Nelson E B 2000 Methods for assessing the composition and diversity of soil microbial communities; Appl. Soil Ecol. 15 25-36

Hobel C F V, Marteinsson V T, Hreggvidsson G O and Kristjansson J K 2005 Investigation of the microbial ecology of intertidal hot springs by using diversity analysis of 16S rRNA and Chitinage genes; Appl. Environ. Microbiol. 63 4516-4522

Hugenholtz P and Pace N R 1996 Identifying microbial diversity in the natural environment: a molecular approach; T. Biotech. 14 190-198

Hulton C S, Higgins C F and Sharp P M 1991 ERIC sequences: a novel family of repetitive elements in the gonomes of Escherichia coli, Salmonella typhimurium and other entrobacteria; Mol. Microbiol. 5 825-834

Hunter-Cevera J C 1998 The value of microbial diversity; Curr. Opinion Microbiol. 1 278-285

Kirk J L, Klironomos J N, Lee H and Trevors J T 2004 The effect of perennial rhygrass and alfa alfa on microbial abundance and diversity in petroleum contaminated soil; J. Environ. Poll. 133 455-465

Lauber C L, Hamady M, Knight R and Fierer N 2009. Pyrosequencing based assessment of pH as a predictory soil bacterial community structure at the continental scale. Appl Environ Microbiol 75 (15) 5111-5120.

Laverman A M, Braster M, Roling W F M and van Verseveld H W 2005 Bacterial community structure and metabolic profiles in a forest soil exhibiting spatially variable net nitrate production; Soil. Biochem. 37 1581-1588

Liebner S, Harder J and Wagner D 2008 Bacterial diversity and community structure in polygonal Tundra soils from Samoylov island, Lena Delta Siberia. Int Microbiol 11 195-202.

Lodge D G, Hawksworth D L and Ritchie B J 1999 Microbial Diversity and tropical forest functioning; Ecol. Studies 122 69-100

Ludwig W and Schliefer K H 1994 Bacterial phylogeny based on 16S and 23S rRNA sequence analysis; FEMS Microiol. Rev. 15 2-3

Lundgren B 1982 Bacteria in a pine forest soil as affected by clear cutting; Soil Biol. Biochem. 14: 537-542

Lupiwayi N Z, Rice W A and Clayton G W 1998 Soil microbial diversity and community structure under wheat as influenced by tillage and crop rotation; Soil Biol. Biochem. 30 1733-1741.

Lyautey E, Lacoste B, Ten-Hage L, Rols J and Garabetian F 2005 Analysis of bacterial diversity in river biofilms using 16S rDNA PCR-DGGE: methodological settings and fingerprints interpretation; J. Water Res. 39 380-388

Maila M P, Randima P, Surrindge K, Dronen K and Cloete T E 2005 Evaluation of microbial diversity of different soil layers at a contaminated dielsel site; Int. Biodet. Biodegrad. 55 39-44

McCaig A E, Glover L A and Prosser J I 2001 Numerical analysis of grassland bacterial community structure under different land management regimes by using 16S ribosomal DNA sequence data and denaturing gradient gel electrophoresis; Appl. Environ. Microbiol. 67 4554-4559

Meyers R M, Fischer S G Lerman, L S and Maniatis T 1985 Nearly all single base substituions in DNA fragments joined to GC-clamp can be detected by denaturing gradient gel electrophoresis; Nucleic Acids Res. 13 3131-3145.

Meyers R M, Maniatis T and Lerman L S 1987 Detection and localization of single base changes by denaturing gradient gel electrophoresis; Methods Enzymol. 155 501-527

Muyzer G, Waal E C D and Uitterlinden A G 1993 Profiling of complex microbial populations by denaturing gradient gel electrophoresis and analysis of polymerase chain reaction-amplified genes coding for 16S rRNA; Appl. Environ. Microbiol. 59 695-700

Niemann S, Dammann-Kalinowski T, Nagel A, Puhler A and Selbitschka W 1999 Genetic basis of enterobacterial repetitive intergenic consensus (ERIC)-PCR) fingerprint pattern in Sinorhizobium meliloti and identification of S. meliloti employing PCR primers derived from an ERIC-PCR fragment; Arch. Microbiol. 1 22-30

Ogram A 2000 Soil molecular ecology at age 20: methodological challenges for the future; Soil Biol. Biochem. 32 1499-1504

Ogram A, Sayler G S, and Barkay T 1987 The extraction and purification of microbial DNA from sediments. J. Microbiol. Methods. 7 57-66

J. BIOL. ENVIRON. SCI., 2011, 5(15), 183-194

Okalebo J R, Gathua K W and Woomer P L 1993 Laboratory Methods of Soil and Plant Analysis: A working Mannual. Soil Science Society of East Africa. Technical publication No.1,TSBF. Marvel EPZ (Kenya) Ltd, Nairobi, Kenya

Ovreas L, Jinsen S, Daae F L and Torsvik V 1998 Microbial community changes in perturbed agricultural soil investigated by molecular and physiological approaches; Appl. Environ. Microbiol. 64 2739-2742

Olsen G J, Lane D J, Giovannoii S J, Pace N R and Stahl D A 1986 Microbial ecology and evolution: a ribosomal RNA approach; Ann. Rev. Microbiol. 40 337-365.

Olsen G J, Woese C R and Overbeek R 1994 The winds of (evolutionary) change: breathing new life into microbiology; J. Bacteriol 176 1-2

Rademaker J L W and de Bruijn F J Characterization and classification of microbes by rep- PCR genomic fingerprinting and computerassisted pattern analysis; "In: Cateno-Annoles G and Gresshoff P M (Eds), DNA Markers: Protocols, Applications and Overview, John Wiley & Sons, New York, pp 151-171."

Robe P, Nalin R, Capellano C, Vogel T. M and Simonet P 2003 Extraction of DNA from soil; Europ. J. Soil Biol. 39 133-190

Rondon M R, Goodman R M and Handesman J 1999 The earth's bounty: assessing and accessing soil microbial diversity; T. Biotech. 17 403-409

Roose-Amsaleg C L, Garnier-Sillam E and Harry M 2001 Extraction and purification of microbial DNA from soil and sediment samples; Appl. Soil Ecol. 18 47-60

Santros S R and Ochman H 2004 Identification and phylgenetic sorting of bacterial lineages with universally conserved genes and proteins; Environ. Microbiol. 6 754-759

Saxena K G and Ramakrishnan P S 1986 Nitrification during slash and burn agriculture (jhum) in North-eastern India. Acta Oecologica/ Oecolagia Plantaraum; 7 319-331

Schloter M, Lebuhn M, Thiarry H and Hartmann A 2000 Ecology and evolution of bacterial microdiversity; FEMS Microbiol. Rev. 24 647-660

Sehgal J and Abrol I P 1994 Soil degradation in India: Status and impact; Oxford & IBH Publishing Co. Pvt. Ltd., New Delhi, pp 80.

Singh S 1999 A Resource Atlas of Arunachal Pradesh. Deptartment of Planning, Govt. of Arunachal Pradesh, Itanagar, India

Singh S S 2002 Microbial communities and their activities in degraded and undegraded forest soils of Arunachal Pradesh, Ph.D. Thesis, North eastern Hill University, Shillong, India

Sharma S, Rangger A, Von Lutzow M and Insam H 1998 Functional diversity of soil bacterial communities increases after maize litter amendment; Europ. J. Soil Biol. 34 53-60.

Sorensen S R, Rasmussen J Jacobsen C S, Jacobsen O S, Juhler R K and Aamand J 2005

Elucidatiing the key member of a Linuron-Mineralizing bacterial community by PCR and Reverse Transcription-PCR Denaturing Gradient Gel Electrophoresis 16S rDNA gene Fingerprinting and cultivation. Appl Environ Mirobio 71 (7) 4144-4148.

Tengerdy R P and Szakacs G 1998 Perspectives in agrobiotechnology; J. Biotech. 66 91-99

Tiwari S C, Sorokhaibam S S, Dkhar M S and Mishra R R 2002 Soil degradation affects microbial biomass carbon and dehydrogenase activity in humid tropical hilly forest soils; Asian J. Microbiol. Biotech. Env. Sc. 4 143-148

Trosvik V L 1980 Isolation of bacterial DNA from soil; Soil Biol. Biochem. 12 15-21

Trosvik V, Daae F L, Sandaa R and Ovreas L 1998 Novel techniques for analysing microbial diversity in natural and perturbed environments; J. Biotech. 64 53-62

Vazquez F J, Acea M J and Caraballas T 1993 Soil microbial population after wildfire; FEMS Microbiol. Ecol. 13 93-104.

Versalovic J, Koeuth T and Lupski J R 1991 Distribution of repetitive DNA sequences in eubacteria and application to fingerprinting of bacterial genomes; Nucleic Acids Res.19 6823-6831

Volossiouk T, Robb E J and Nazer R N 1995 Direct DNA extraction for PCR-mediated assays of soil organisms; Appl. Environ. Microbiol. 61 3972-3976

Wilson L A and Sharp P M 2006 Enterobacterial repetitive intergenic consensus (ERIC) sequences in Escherichia coli: Evolution and implications for ERIC-PCR; Mol. Biol. Evol. 23 1156-1168.

Yeates G, Davison A D Altavilla N and Veal D A 1998 Methods for microbial DNA extraction from soil for PCR amplification; Biol. Proc. Online 1 40-47

Zhou J, Davey M E, Figueras J B, Rvkina E, Gilchinsky D and Tiedje J M 1997

Phylogenetic diversity of a bacterial community determined from Siberian tundra soil DNA; Microbiol. 143 3913-3919