Characterization of the Depth-Related Changes in the Microbial Communities in Lake Hovsgol Sediment by 16S rRNA Gene-Based Approaches

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The undisturbed sediment of Lake Hovsgol (Mongolia) is scientifically important because it represents a record of the environmental changes that took place between the Holocene (the present age) and Pleistocene (the last ice age; 12,000 ¹⁴C years before present day). Here, we investigated how the current microbial communities change as the depth increases by PCR-denaturing gradient gel electrophoresis (DGGE) analysis of the 16S rRNA genes of the microbial communities. The microbial diversity, as estimated by the Shannon index, decreased as the depth increased. In particular, significant changes in archaeal diversity were observed in the middle depth (at 39~42 cm depth of total 60 cm depth) that marks the border between the Holocene and Pleistocene. Phylotype belonging to Beta- and Gamma-Proteobacteria were the predominant bacteria and most of these persisted throughout the depth examined. However, as the depth increased, some bacteria (some genera belonging to Beta-Proteobacteria, Nitrospira, and OP8-9) were not detectable while others (some genera belonging to Alpha-, Beta-, Gamma-Proteobacteria) newly deteced by DGGE. Crenarchaea were the predominant archaea and only one phylotype belonging to Euryarchaea was found. Both the archaeal and bacterial profiles revealed by the DGGE band patterns could be grouped into four and three subsets, respectively, subsets that were largely divided by the border between the Holocene and Pleistocene. Thus, the diversity of the current microbial communities in Lake Hovsgol sediments decreases with increasing depth. These changes probably relate to the environmental conditions in the sediments, which were shaped by the paleoclimatic events taking place between the Holocene and Pleistocene.

Keywords: Lake Hovsgol, sediment, DGGE, microbial diversity, 16S rRNA gene

Benthic sediment represents one of the most complex microbial habitats on Earth (Priscu *et al.*, 1998). The microorganisms in benthic sediments play important roles in nutrient cycles such as methane production (Chan *et al.*, 2005), sulfate reduction (Li *et al.*, 1999) and ammonia oxidation (Hastings *et al.*, 1998). Consequently, investigating the microbial communities and their function in freshwater sediments will greatly enhance our understanding of the aquatic ecosystem (Tamaki *et al.*, 2005). Since most freshwater sediments are quite heterogeneous ecosystems that give rise to many different environmental niches even on a millimeter scale (Spring *et al.*, 2000), only few studies have analyzed how microbial communities change as sediment depth increases (Urakawa *et al.*, 2000; Koizumi *et al.*, 2003).

Lake Hovsgol (Khubsugul, Mongolia) is one of the largest lake in Asia (Lake Baikal is the largest) (Karabanov *et al.*, 2004). It is famous for being one of the very few ultra-oligotrophic pure lakes in the world (Belykh *et al.*, 2005), and has characteristics of low mineral content (total ion concen-

tration is 210~230 mg/L) and alkalic pH water (pH 8.1~8.5) (Kozlov et al., 1989). Lake Hovsgol is also very interesting from a scientific point of view because it is known to be 2.5~4.0 Ma (million years) old. Since the last glacial maximum (LGM; last ice age) ended 12,000 to 13,000 years ago (Goulden et al., 2003), it is certain that Lake Hovsgol experienced the LGM. Analysis of the lake's sediments (Nara et al., 2005) has revealed that during the LGM represented by the >40~50 cm-depth sediments, (Fedotov et al., 2004), there was no accumulation of planktonic/benthic diatoms, chrysophyte cysts, sponge spicules and zooplankton in the lake (Fig. 2). After the LGM, however, nutrient loading from the watershed resumed and the organisms listed above began to accumulate again, as revealed by the $0 \sim 40$ cm-deepth sediments (Fig. 2). Thus, the stratified sediments of Lake Hovsgol represent a record of past climatic changes that have impacted significantly on the ecology of the lake.

We speculated that these different depth-related environmental conditions may constitute specific niches for microorganisms and that the microbial communities currently living in the Lake's sediments may consequently vary with sediment depth. Since the current microbial diversity in the stratified

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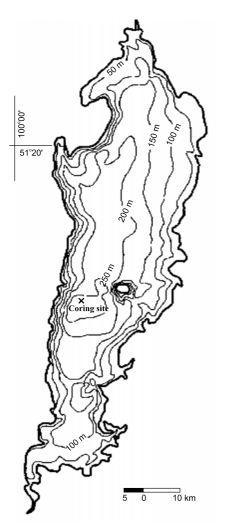


Fig. 1. Map of Lake Hovsgol with sampling site.

sediments of Lake Hovsgol has not yet been investigated, we subjected the microbial communities in various sediment depths to PCR-denaturing gradient gel electrophoresis (DGGE) analysis.

The 'great plate count anomaly' paradigm refers to the observation that often the most prevalent microorganisms cannot be cultivated (Staley and Konopka, 1985). This anomaly has caused microbial ecologists to move from agar mediato PCR-based approaches to estimate microbial diversity and analyze the structure of microbial communities (Rondon et al., 1999; Hugenholtz, 2002). Such molecular biological techniques employ certain molecular markers, such as 16S rRNA gene or its encoding gene, to detect and identify particular microorganisms. Analysis of the structure of microbial communities has been aided in particular by the introduction of DGGE (Muyzer et al., 1993). In this study, we used DGGE analysis of 16S rRNA genes to study the depth-related changes in the microbial communities in the sediment layers of Lake Hovsgol. How the paleoclimatic history of the lake shaped the physico-chemical properties of the sediments, and how these properties have in turn molded the microbial populations we detected is discussed.

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Materials and Methods

Sample collection

Sampling of the sediment was carried out in March 2004 at one central location (50°57'18.4"N, 100°21'32.7"E, water depth; about 260 m) of Lake Hovsgol by using a Benthos gravity corer (Fig. 1). The core immediately was transported to the local laboratory, and frozen until next procedure. The core in the plastic pipe was sectioned with 3 cm interval down to a depth of approximately 60 cm. The each sliced core was re-sampled at each center position of core and packed into sterile plastic tubes and stored at -70°C until analysis.

DNA extraction from sediment samples and purification Total genomic DNA of the frozen sediment samples was isolated by using the bead-beating method as described previously (Yeates *et al.*, 1998). All DNA samples were treated with RNase A (Sigma, USA) and purified by using the ethanol precipitation method. The extracted DNAs were further purified by using the UltraClean[®] Microbial DNA Isolation Kit (Mo Bio Laboratories, USA) with the following modifications: the bead-beating step was excluded, and DNA solution was added to Solution MD1 instead of the Micro-Bead Solution.

PCR-Amplification of bacterial and archaeal 16S rRNA genes PCR amplification of bacterial 16S rRNA genes was performed by using the GC338f-518r primer combination as described Muyzer *et al.* (1993) and performed using amplifying condition of Henckel *et al.* (1999). Nested PCR amplification of archaeal 16S rRNA genes was carried out by using the arch20f-958r primer set (DeLong, 1992; DeLong *et al.*, 1999) for the first round of amplification and the archGC340f-519r set (Ovreas *et al.*, 1997) for the second round of amplification as described previously (Ferris *et al.*, 1996).

DGGE analysis of bacterial and archaeal diversities DGGE was performed by using the D-Code System (Biorad, USA). Amplified DNAs were loaded onto 8% (w/v) polyacrylamide gels with 40 to 60% (Archaea) and 30 to 60% (Bacteria) denaturing gradient (where 100% of the denaturant consists of 7 M urea and 40% formamide) in $1 \times$ TAE (20 mM Tris, 10 mM acetate, 0.5 mM EDTA, pH 8.3) and subjected to 14 h of electrophoresis at 60°C and 80 V. Thereafter, the gels were soaked in SYBR Green I nucleic acid stain (Bioneer, Korea) and photographed with UV transillumination.

Sequencing and phylogenetic analysis of excised DGGE bands Dominant bands were excised with razor blades from the DGGE gels and soaked overnight in 20 μ l of 0.1× TE buffer solutions. One microliter of each solution was PCR-reamplified and directly sequenced. The sequences were taxonomically assigned by using BLAST on the GenBank nucleotide database [National Center for Biotechnology Information (NCBI)] (Altschul *et al.*, 1990). Sequences of 16S rRNA genes were aligned and a phylogenetic tree was constructed by using CLUSTAL X software (Thompson *et al.*, 1997).

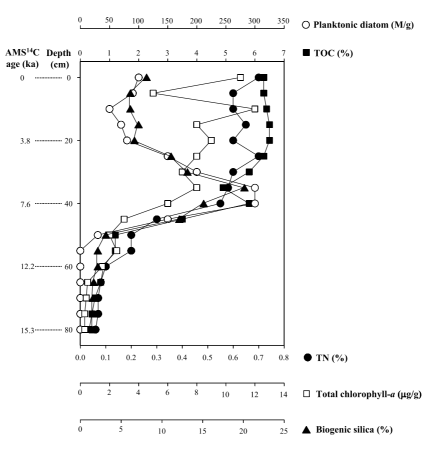


Fig. 2. Depth-related changes in total organic carbon (TOC, black squares), total nitrogen (TN, black circles), planktonic diatoms (open circles), total chlorophyll-a (open squares) and biogenic silica (black triangles) in Lake Hovsgol sediment (redrawn from references 29 and 42).

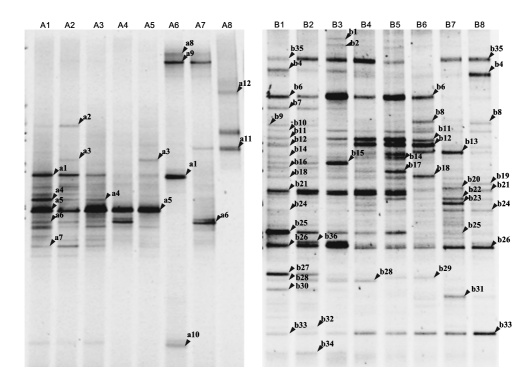


Fig. 3. DGGE analysis of PCR-amplified 16S rDNA fragments from the archaeal (A) and bacterial (B) communities in the sediments of Lake Hovsgol. Lanes 1, $0 \sim 3$ cm; 2, $6 \sim 9$ cm; 3, $12 \sim 15$ cm; 4, $21 \sim 24$ cm; 5, $30 \sim 33$ cm; 6, $39 \sim 42$ cm; 7, $48 \sim 51$ cm; 8, $57 \sim 60$ cm.

Nucleotide sequence accession numbers

The nucleotide sequences obtained detected in this study are available from the GenBank nucleotide database at the NCBI website (http://www.ncbi.nlm.nih.gov) under accession numbers DQ452578-DQ452612.

Clustering and diversity analysis of DGGE profiles

Similarities between the DGGE profiles of samples and the band intensities of each lane were analyzed by BioNumerics software (BioSystematica, UK). For DGGE profile analysis, each lane was chosen separately, common bands were selected as positions for normalization to compensate for differences in migration distance due to gel heterogeneity. The normalized banding patterns were then used to generate dendrograms by calculating the Pearson product moment correlation coefficient and by employing the unweighted pair group method with arithmetic averages (UPGMA) clustering. For diversity analysis, densitometric curves from each lane were acquired by using the Bionumerics software. A matrix was then constructed with this information and used to calculate a set of numerical values that described the diversity of the prokaryotic communities. As a measure of the structural diversity of prokaryotic communities, the Shannon index (Koizumi et al., 2003) of general diversity (H) was calculated by using the formula $H=-\Sigma Pi \cdot \ln (Pi)$, in which Pi is the importance probability of the bands in a track. H was calculated on the basis of the bands on the gel tracks, namely, on the intensities of the bands (as judged by

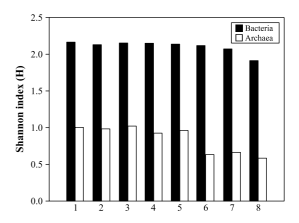


Fig. 4. Change of Shannon index values throughout the depth of lake Hovsgol sediment. Shannon index values (H) were calculated on the basis of the number and intensity of the bands on the gel tracks. Lanes 1, $0 \sim 3$ cm; 2, $6 \sim 9$ cm; 3, $12 \sim 15$ cm; 4, $21 \sim 24$ cm; 5, $30 \sim 33$ cm; 6, $39 \sim 42$ cm; 7, $48 \sim 51$ cm; 8, $57 \sim 60$ cm.

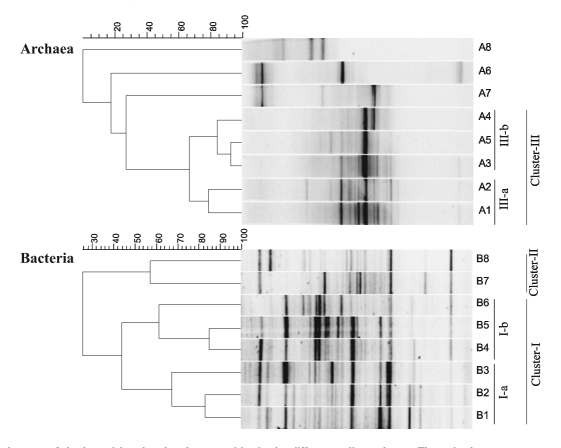


Fig. 5. Dendrogams of the bacterial and archaeal communities in the different sediment layers. These dendrograms were constructed by using Pearson's curve-based correlations and the UPGMA clustering method (BioNumerics software; BioSystematica, UK). Lanes 1, 0-3 cm; 2, 6-9 cm; 3, 12-15 cm; 4, 21-24 cm; 5, 30-33 cm; 6, 39-42 cm; 7, 48-51 cm; 8, 57-60 cm.

Pearson correlation (Opt:0.50%) [15.8%~95.4%]

peak heights in the densitometric curves). The importance probability (Pi) was calculated as Pi=ni/N, where ni is the height of the ith peak and N is the sum of all peak heights in the densitometric curve.

Results

DGGE analysis of the bacterial and archaeal communities in the sediments of Lake Hovsgol

To investigate how the microbial population in Lake Hovsgol changes with sediment depth, we subjected eight 3 cm-thick layers of the sediments (up to about 60 cm deep) to DGGE analysis using 16S rRNA gene. Although bacterial 16S rRNA genes were successfully amplified by a direct PCR step, archaeal 16S rRNA gene amplification only occurred upon nested PCR using the arch20f-958r primer set for the first round amplification step and the archGC340f-519r set for the second round amplification step. This indicates that the archaeal community in the sediments is quantitatively much smaller than the bacterial community.

The DGGE patterns revealed by the bacterial PCR products were also more diverse than those of the archaea (Fig. 3), as on average 20.8 and 9.1 DGGE bands were obtained from each sediment layer, respectively. Since the number and thickness of the bands in a DGGE profile can represent species diversity (Muyzer, 1999), the DGGE band patterns obtained from each 3 cm thick-sediment were used to calculate the Shannon index of diversity (H). This index allowed us to determine and compare the microbial diversity in each environmental niche. Figure 4 shows that H dropped with increasing sediment depth, suggesting that diversities of both bacterial and archaeal communities decreased as the sediment depth increased. In particular, the archaeal diversity index fell markedly at the 39~42 cm depth; this low diversity was also observed in the deeper sediments. It is also clear that the archaeal DGGE profiles of the deeper sediments (samples 6~8; 39~60 cm) differ from the profiles of the upper layers (Fig. 3A), as many of bands of the upper layers disappeared and small number of new bands detected. In contrast, most of the bacterial DGGE bands were observed in all layers, although a few bands did disappear with others newly appearing as depth increased (Fig. 3B). Thus, the composition of the archaeal community is more clearly affected by the depth-related environmental changes of the sediments than the bacterial community.

To measure more clearly how the microbial communities change with sediment depth, we subjected the DGGE profiles to UPGMA cluster analysis (Fig. 5). This revealed that the members of the bacterial communities in the two deepest layers (B7; 48~51 cm and B8; 57~60 cm) fall into a different cluster (Cluster II) relative to the communities in the upper layers, which make up Cluster I (B1-6; 0~42 cm). The entire densitometric curves for each lane were then compared numerically by calculating Pearson product moment correlation coefficients (r). When r equals 100, this indicates that the dominant species in the samples being compared are identically distributed. For the Lake Hovsgol sediments, the bacterial r values ranged from 43.77% to 84.64% (Fig. 5). This analysis revealed that the Cluster I bacterial profiles could be further subdivided into two sub-clusters, namely, Cluster I-a (B1-3; $0\sim15$ cm) and Cluster I-b (B4-6; $21\sim42$ cm). When the lanes within Cluster I-a or Cluster I-b were compared (intra-cluster comparisons), the r values exceeded 61.15%, which indicates very similar DGGE profiles. Moreover, these two clusters were related more closely to each other (r=43.77%) than to Cluster II (r=25.49%).

In the case of the archaea, UPGMA cluster analysis revealed that the DGGE profiles of the top five layers were very similar to each other and differed from those of the three bottom layers; the bottom three layers also did not resemble each other. As a result, the top five layers could be grouped into Cluster III while the lowest three layers formed individual groups. Calculation of the Pearson coefficients to compare the archaeal DGGE patterns revealed r values that ranged from 17.60% to 92.29% (Fig. 5). These r values allowed Cluster III to be divided into two sub-clusters, namely, Cluster III-a (A1-2; 0~9 cm) and Cluster III-b (A3-6; 12~42 cm). These observations support our observations above that suggest the depth-related environmental changes in Lake Hovsgol sediments affect the archaeal communities.

Phylogenetic profiling of DGGE band clones

All DGGE bands that were obvious to the naked eye were excised from the gel, PCR-reamplified and subjected to nucleotide sequencing. Repeatedly excised bands located at the same position in the gel that showed identical nucleotide sequences were named, counted, and indicated as one representative. Thirty six sequences were obtained from bacterial DGGE bands, and twelve sequences from archaea (Fig. 3). Since some full-length bacterial sequences cannot be determined due to chimerism or co-migration of DGGE fragments (Qiu *et al.*, 2001), incomplete sequences were not used for analysis (Table 1). In conclusion, total 23 and 12 sequences were obtained from the bacterial and archaeal DGGE bands, respectively.

Comparison of the 16S rRNA gene sequences derived from DGGE bands with those in the GenBank database revealed the sequenced bacterial fragments were clustered into eight phylogenetic groups (Table 1 and Fig. 6A). These consisted of Alpha-Proteobacteria (YDB31), Beta-Proteobacteria (YDB11, 13, 14, 17, 20, 23, 24, 25, and 26), Gamma-Proteobacteria (YDB1, 2, 6, 8, 21, and 35), Firmicutes (YDB4 and 29), Nitrospira (YDB27 and 30), Actinobacteria (YDB33), and OP8 & 9 (YDB28 and YDB15). The Proteobacteriarelated sequences were most similar to those of uncultured proteobacterial clones that had been isolated from marine sediment (YDB31) (Asami et al., 2005), an uranium-contaminated aquifer (YDB14 and YDB17), deep subsurface sediment (YDB24), subseafloor (YDB26), cold-tolerant alpine soil (YDB1 and YDB6) (Meyer et al., 2004), the Northern Baltic sea (YDB2) (Kisand et al., 2002) and the Sub-Antarctica (YDB21). The sequences of YDB4 and YDB29 in the Firmicutes group matched those of the facultative psychrophilic low G+C Gram-positive bacterium HTA563 (100%) from deep sea mud (Takami et al., 1997) and the uncultured bacterium Neu2P3-93 (99%) from the sediment of a reservoir (this latter bacterium was closely related to Clostridium scatologenes). The DGGE bands of YDB27 and YDB30, which could be grouped into the phy-

Table 1. Phylogenetic analysis of the 16S rRNA sequences of the DGGE bands obtained from the sediment layers of Lake Hovsgol (The position of each band in the DGGE gel is shown in Fig. 3.)

Clone	Clone Accession Phylogenetic number group		Nearest neighbor	Identity (%)	Source of the closest relative	Nearest isolate	Identity (%)
YDB1	DQ452578	Gamma	Pseudomonas sp. BG2dil (AY263468)	97% alpine soil (Meyer <i>et al.</i> , 2004)		Pseudomonas freder- iksbergensis isolate OUCZ24 (AY785733)	97%
YDB2	DQ452579	Gamma	Pseudomonas sp. GOBB3-207 (AF321048)	97%	estuary (Kisand <i>et al.</i> , 2002)	Pseudomonas sp. B65 (AF332541)	97%
YDB4	DQ452580	Firmicutes	Low G+C Gram-positive bacte- rium isolate HTA563 (AB002641)	100%	sea mud (Takami <i>et al.</i> , 1997)	Staphylococcus epi- dermidis (AY308046)	100%
YDB6	DQ452581	Gamma	Pseudomonas sp. BE3dil (AY263472)	100%	alpine soil (Meyer <i>et al.</i> , 2004)	Pseudomonas sp. An23 (PSP55116)	100%
YDB8	DQ452582	Gamma	Pseudomonas sp. NMX (AF482685)	98%	soil sediment (Coates et al., 2002)	Pseudomonas sp. HS300 (AY940129)	98%
YDB11	DQ452583	Beta	Uncultured beta proteobacte- rium clone Gitt-KF-125 (AJ532686)	100%	uranium mill tailings (Selenska-Pobell, 2002)	Janthinobacterium agar- icidamnosum (Y08845)	99%
YDB13	DQ452584	Beta	Uncultured beta proteobacte- rium clone AKYG404 (AY922005)	98%	farm soil (Tringe et al., 2005)	Zoogloea ramigera (D14257)	96%
YDB14	DQ452585	Beta	Uncultured beta proteobacte- rium clone B-BH93 (AY622261)	98%	subsurface soil (Reardon <i>et al.</i> , 2004)	Janthinobacterium sp. Rud09 (AY788980)	98%
YDB15	DQ452586	OP9	Uncultured bacterium clone Amsterdam-2B-61 (AY592418)	98%	deep-sea mud volcanoes	NA	
YDB17	DQ452587	Beta	Uncultured beta proteobacte- rium clone C-CY80 (AY622237)	95%	subsurface soil (Reardon <i>et al.</i> , 2004)	Janthinobacterium agar- icidamnosum (Y08845)	95%
YDB20	DQ452588	Beta	Uncultured bacterium clone 4-21 (AF351229)	98%	aquifer waters (Bakermans and Madsen, 2002)	Dechloromonas sp. FL9 (AF288773)	95%
YDB21	DQ452589	Gamma	Vibrio sp. U32 (AY864627)	96%	Sub-Antarctica	<i>Vibrio</i> sp. U32 (AY864627)	96%
YDB23	DQ452590	Beta	Uncultured bacterium clone 2-7 (AY548943)	99%	ANAMMOX sludge	Ralstonia solanacearum (AY216796)	97%
YDB24	DQ452591	Beta	Uncultured proteobacterium clone NANK-B57 (AY436534)	100%	marine sediment core (Newberry et al., 2004)	<i>Ralstonia</i> sp. Q3-8/14 (AY216798)	100%
YDB25	DQ452592	Beta	Uncultured <i>Thiobacillus</i> sp. clone 44a-B2-21 (AY082471)	98%	mine drainage	Thiobacillus thioparus (M79426)	96%
YDB26	DQ452593	Beta	Uncultured bacterium clone 33-FL54B99 (AF469368)	100%	subseafloor (Huber <i>et al.</i> , 2003)	Ralstonia sp. 50 (AY177368)	100%
YDB27	DQ452594	Nitrospira	Uncultured bacterium clone ITKB-228 (AB198852)	98%	subsurface sediment	Leptospirillum ferroox- idans (X86776)	84%
YDB28	DQ452595	OP8	uncultured bacterium clone SH050-1C2 (AY781382)	100%	lake sediment	NA	
YDB29	DQ452596	Firmicutes	Unidentified bacterium clone Neu2P3-93 (AJ518357)	99%	sediment (Wobus et al., 2003)	Clostridium scatologenes (AJ427628)	99%
YDB30	DQ452597	Nitrospira	Uncultured bacterium clone ITKB-228 (AB198852)	99%	subsurface sediment	Leptospirillum ferroox- idans (X86776)	84%
YDB31	DQ452598	Alpha	Uncultured alpha proteobacte- rium clone KY40 (AB116424)	98%	marine sediment (Asami et al., 2005)	Rhodobium bactotapeta- rum (AJ510235)	94%
YDB33	DQ452599	Actinobacteria	Uncultured bacterium isolate SSDG gel band 9 (AY362838)	100%	waste water (Lee et al., 2004)	Propionibacterium acnes (AY642054)	99%
YDB35	DQ452600	Gamma	Pseudomonas sp. An15 (AJ551153)	100%	deep sea sediment	Pseudomonas sp. An15 (AJ551153)	100%

Table	1.	Continued
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Clone	Accession number	Phylogenetic group	Nearest neighbor	Identity (%)	Source of the closest relative	Nearest isolate	Identity (%)
YDA1	DQ452601	Marine Benthic group C	Uncultured crenarchaeote VAL81 (AJ131315)	99%	forest lake (Jurgens et al., 2000)	NA	
YDA2	DQ452602	Marine Benthic group C	Uncultured archaeon clone St_D_05 (AY531752)	99%	lake sediment	NA	
YDA3	DQ452603	Marine Benthic group C	Uncultured crenarchaeote VAL81 (AJ131315)	97%	forest lake (Jurgens et al., 2000)	NA	
YDA4	DQ452604	Marine Benthic group C	Uncultured archaeon clone MKCSM-bx10 (DQ363782)	97%	mangrove soil	NA	
YDA5	DQ452605	Marine Benthic group C	Uncultured archaeon clone ARC-U3SP-10 (AY456738)	99%	soil (Castro <i>et al.</i> , 2004)	NA	
YDA6	DQ452606	Marine Benthic group C	Uncultured crenarchaeote clone X4Ar42 (AY607267)	99%	rice field soil (Lueders et al., 2004)	NA	
YDA7	DQ452607	Marine Benthic group C	Uncultured archaeon clone ARC-U3SP-10 (AY456738)	95%	soil (Castro <i>et al.</i> , 2004)	NA	
YDA8	DQ452608	Group 1.1a	Uncultured archaeon clone ODP1230A22 (AB177110)	99%	marine sediment (Inagaki et al., 2006)	NA	
YDA9	DQ452609	Group 1.1a	Uncultured archaeon clone 1.8mBB.27 (AY661852)	99%	marine sediment (Sorensen et al., 2004)	NA	
YDA10	DQ452610	Euryarchaea	Uncultured archaeon cloneASN24 (AB161351)	100%	soil (Kasai et al., 2005)	NA	
YDA11	DQ452611	Group I.1b	Uncultured Cenarchaeote clone HPA-85 (AY430142)	99%	stromatolite (Burns <i>et al.</i> , 2004)	NA	
YDA12	DQ452612	Group I.1b	Uncultured crenarchaeote clone MWS38 (AY522902)	100%	mesophilic soil (Sliwinski and Goodman, 2004)	NA	

* Organisms for which a related isolate in their own phylum could not be found on the basis of the 16S rRNA sequence are marked as NA

Table 2. The number of microorganisms belonging to the major bacterial and archaeal groups found in the sediment layers of La	ke					
Hovsgol (Lanes 1, 0~3 cm; 2, 6~9 cm; 3, 12~15 cm; 4, 21~24 cm; 5, 30~33 cm; 6, 39~42 cm; 7, 48~51 cm; 8, 57~60 cm)						

	Lane 1	Lane 2	Lane 3	Lane 4	Lane 5	Lane 6	Lane 7	Lane 8
α-Proteobacteria	0	0	0	0	0	1	1	1
β-Proteobacteria	7	7	7	8	8	6	6	5
γ-Proteobacteria	3	3	5	5	5	4	3	3
Actinobacteria	1	1	1	1	1	1	1	1
Firmicutes	1	1	1	1	2	2	1	1
Nitrospira	2	2	2	0	0	0	0	0
OP8,9	2	2	2	2	1	0	0	0
Marine Benthic Group C	6	7	5	3	5	1	1	0
Group 1.1a	0	0	0	0	0	2	2	0
Group 1.1b	0	0	0	0	0	0	1	2
Euryarchaea	1	1	1	0	0	1	0	0

lum *Nitrospira* (\geq 98% similarity), were most similar in sequence to isolate of *Leptospirillum ferrooxidans* (84%); *L. ferrooxidans* is known as an iron-oxidizing bacterium that is often found in the deep underground of mines (Schrenk *et al.*, 1998). YDB15 and YDB28 resembled OP9 and OP8, respectively, which were detected in the anoxic zone of the Cariaco Basin (Madrid *et al.*, 2001) and in anaerobic sludge (Chouari *et al.*, 2005). Table 2 reveals that the diverse bac-

terial phylotypes are found in most of the sediment layers. However, the bacterial community did change somewhat as the depth of the sediments increased (Table 2). While most Proteobacteria were found throughout the sediments, the *Janthinobacterium*-like β -proteobacterial clones were only found in the shallower layers, the deepest being the 39-42 cm layer. In contrast, the *Zoogloea*-like *Beta*-Proteobacteria and the *Rhodobium*-like *Alpha*-Proteobacteria were not

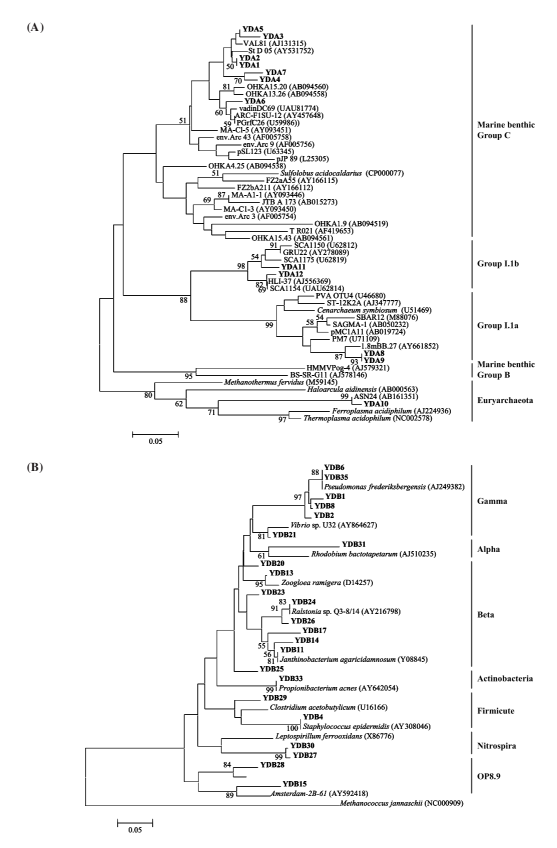


Fig. 6. Neighbor joining phylogenetic trees of the Lake Hovsgol archaea (A) and bacteria (B) that were identified on the basis of partial 16S rDNA sequences. The phylogenetic tree was constructed by using the CLUSTAL X package program as described in 'Materials and Methods'. The scale bar represents 0.05 substitutions per nucleotide position. Bootstrap values (expressed as percentages of 100 replications) greater than 50% are shown at the branch points.

Archaeal DGGE bands were also sequenced and analyzed. The Crenarchaea formed the predominant archaeal group observed in Lake Hovsgol sediments and only one phylotype belonging to Euryarchaea (YDA10) was found. This euryarchaeal clone was grouped with the uncultured freshwater sediment archaeon ASN24 (100%), which was isolated from petroleum-contaminated soil (Kasai et al., 2005). With regard to the Crenarchaea, the YDA1, 2, 3, 4, 5, 6, and 7 clones were grouped into marine benthic Crenarchaea group C, while YDA8 and 9 were grouped into marine Crenarchaea group 1.1a and YDA11 and 12 were grouped in the soil Crenarchaea group 1.1b (Table 1 and Fig. 6A). Unlike the bacterial community, however, each band was found in only a limited number of sediment samples rather than in all or many of them (Table 2). In addition, each phylogenetic group was found at a particular depth (Table 2). For example, while the marine benthic group C Crenarchaea predominated in the sediments, this group gradually became less frequently detected as the depth decreased. In addition, while the Group 1.1a and 1.1b Crenarchaea found at the depth of 39~42 cm, the Euryarchaea disappeared.

Discussion

Lake Hovsgol is 2.5~4.0 Ma (million years) old (Goulden et al., 2003) and has never been subjected to anthropogenic stress (Belykh et al., 2005). Analyses of its sediment layers have revealed that consecutive paleoclimatic transitions have generated different environmental niches. Lake Hovsgol was glaciated during the LGM of the Pleistocene, at which point the level of the lake was 200 m lower than it is at present (Fedotov et al., 2002). During this period, the water of the lake did not run out and water circulation did not occur. The low water level and volume during the LGM drastically changed the lake water chemistry as it elevated the carbonate concentrations (Karabanov et al., 2004). Such high salinity, along with the sudden change of climate, the cold temperatures, and limited water circulation, markedly reduced the productivity of the organisms living in the lake. The water level and circulation in the lake recovered around 12¹⁴C ka (about 12,000 years ago, as determined by radiocarbon C¹⁴ dating), at which point the ecosystem of the lake is believed to have regenerated (Prokopenko et al., 2003).

Nutrient availability is a primary factor that can change the microbial communities in lake sediments, as Parkes *et al.* (2005) have revealed that high levels of organic matter in the sediment greatly stimulates prokaryotic activity and results in highly diverse prokaryotic populations. In Lake Hovsgol during the LGM, the environmental conditions resulted in a marked decline in nutrient availability. This is revealed by measurements of the chlorophyll-*a* concentrations in the sediments; Nara *et al.* (2005) found that these concentrations were 10-fold higher in the upper sediments of Lake Hovsgol than in its >50 cm sediments. This change was paralleled by decreases in total organic carbon, total nitrogen, biogenic silica concentrations, planktonic diatom numbers, and benthic diatom numbers (Karabanov *et al.*, 2004), which indicates that macro-organisms like diatoms largely disappeared during the LGM and only returned at a depth of about 50 cm. Significantly, the 40~50 cm deep Lake Hovsgol sediments have been estimated to be 12,000 years old (Fedotov *et al.*, 2004); this is exactly the border between the Holocene (present age) and Pleistocene (the last glacial age).

In our study, we found the microbial communities in the Lake Hovsgol sediments are much less diverse in terms of both bacteria and archaea in the 39~60 cm sediment lavers (as estimated by the Shannon index) compared to the communities in the upper layers (Fig. 4). In particular, the archaeal communities in the $0 \sim 33$ cm layers were almost twice diverse than the communities found in the 39~60 cm layers. UPGMA analysis of the DGGE profiles of the sediment communities also showed that both prokaryotic communities could be separated at the border between Holocene and Pleistocene that is represented by the 39~42 cm layer. This suggests that the phylogenetic analysis of DGGE clones from Lake Hovsgol sediments reveals the existence of a fairly diverse and stratified bacterial and archaeal community, and the different nutrient concentrations in the sediment strata, which were shaped by paleoclimatic events, may modulate the diversity of the microbial communities currently living in the sediment strata of Lake Hovsgol.

The 16S rRNA gene sequences retrieved from the DGGE bands allowed us to partially identify the microorganisms in the various depth sediments and to determine their phylogenetic affiliations. With regard to the bacterial community in the Lake Hovsgol sediments, it generally resembled the communities found in other lakes with similar habitats. However, although Delta-Proteobacteria and CFB bacteria are frequently detected in the sediments of other lakes (Glockner et al., 2000; Humayoun et al., 2003; Demergasso et al., 2004), they were not detected in the sediments of Lake Hovsgol. Phylogenetic analysis of the archaeal 16S rRNA sequences obtained from Lake Hovsgol sediments revealed two groups, namely, euryarchaea and uncultured crenarchaea (Fig. 6). These crenarchaeal and euryarchaeal groups have also been reported to be present in the sediments of other lakes. Although we tried to detect methanogen using PCR with specific primers (Marchesi et al., 2001), we could not detect the presence of methanogen (data not shown), which suggests that anaerobic digestion coupled with sulfate reduction and methane production may not occur in Lake Hovsgol sediments. According to the study of Fedotov et al. (2004), the sediments of Lake Hovsgol are anoxic from just below the surface and contain many metals; in particular, P, Mn, Fe, and Ba levels increase from the bottom to the surface. It is possible that these elements are reduced in the organic matter-rich sediments into their ionic forms and become precipitated in the oxygenated zones near the surface. That metal elements such as Fe increase in concentration as the sediments get closer to the surface may also be related to the presence of bacteria belonging to the genus Leptospirillum. These iron-oxidizing bacteria occupy the upper sediments (0~15 cm) of Lake Hovsgol

(Table 1 and Fig. 3B). In addition, *Rhodobium*, which is capable of photosynthetic hydrogen production, was absent from the sediments overlying the 39~42 cm layer while *Janthinobacterium*, a nitrate reducer, is only present in this layer and the sediments above it. The appearance of *Leptospirillum* and *Janthinobacterium* in the upper sediments is probably related to changed nutrient levels in the sediments. Thu, the climatic changes that took place between the Holocene and Pleistocene promoted the bloom of primary producers such as diatoms and consequently altered the microbial communities in the sediments and increased their growth.

DGGE method can clearly indicate the existence of common but potentially uncultivated groups of microorganisms in natural environment, it does not permit the more precise identification of these microbes to the species or sub-species level (Table 1 and Fig. 6). Moreover, we cannot determine from the partial sequence information yielded by DGGE how the microorganisms in the lake sediments function or interact. A more comprehensive understanding of the physiology of these organisms and their complex biogeochemical processes will require their cultivation, isolation and characterization (Torsvik *et al.*, 2002; Zengler *et al.*, 2002; Bae *et al.*, 2005).

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