Geographic distance and pH drive bacterial distribution in alkaline lake sediments across Tibetan Plateau

Jinbo Xiong,1 Yongqin Liu,2 Xiangui Lin,1 Huayong Zhang,1 Jun Zeng,1 Juzhi Hou,2 Yongping Yang,2 Tandong Yao,2* Rob Knight3,4 and Haiyan Chu1*
1State Key Laboratory of Soil and Sustainable Agriculture, Institute of Soil Science, Chinese Academy of Sciences, Nanjing 210008, China.
2Key Laboratory of Tibetan Environment Changes and Land Surface Processes, Institute of Tibetan Plateau Research, Chinese Academy of Sciences, Beijing 100085, China.
3Department of Chemistry and Biochemistry, and 4Howard Hughes Medical Institute, University of Colorado, Boulder, CO 80309, USA.

Summary

Continent-scale biogeography has been extensively studied in soils and marine systems, but little is known about biogeographical patterns in non-marine sediments. We used barcode pyrosequencing to quantify the effects of local geochemical properties and geographic distance for bacterial community structure and membership, using sediment samples from 15 lakes on the Tibetan Plateau (4–1670 km apart). Bacterial communities were surprisingly diverse, and distinct from soil communities. Four of 26 phyla detected were dominant: Proteobacteria, Bacteroidetes, Firmicutes and Actinobacteria, albeit 20.2% of sequences were unclassified at the phylum level. As previously observed in acidic soil, pH was the dominant factor influencing alkaline sediment community structure, phylotype richness and phylogenetic diversity. In contrast, archaeal communities were less affected by pH. More geographically distant sites had more dissimilar communities (\( r = 0.443, P = 0.030 \)). Variance partitioning analysis showed that geographic distance (historical contingencies) contributed more to bacterial community variation (12.2%) than any other factor, although the environmental factors explained more variance when combined (28.9%). Together, our results show that pH is the best predictor of bacterial community structure in alkaline sediments, and confirm that both geographic distance and chemical factors govern bacterial biogeography in lake sediments.

Introduction

Understanding which environmental factors influence the variation in microbial communities across different scales is a key goal of ecology. Many studies show that contemporary environmental factors, such as trophic status (Hansel et al., 2008), soil pH (Rousk et al., 2010), metalloid contamination, and the rhizosphere (Xiong et al., 2010), influence microbial community structure. In contrast, and because of these large effects of local factors, links between microbial community structure and geographic distance remain elusive and controversial (Martiny et al., 2006), and are mainly linked to studies of single species, such as Bacillus (Connor et al., 2010), and Cyanobacteria (Bahl et al., 2011). Cho and Tiedje (2000) first showed that the genetic divergence of Pseudomonas isolates correlated with spatial distance at regional (5 m to 80 km) but not greater scales, whereas a larger-scale (1–1000 km) survey of microbial eukaryotes revealed a taxa–area relationship (Hillebrand et al., 2001). Similarly, microbial endemism (Whitaker et al., 2003), and distance effects on microbial spatial dispersal (Martiny et al., 2006), have been reported. However, there is still active debate about the relative contribution of geographic distance to microbial community dissimilarity in soils; it is unclear whether geographic distance is a trivial (Fierer and Jackson, 2006; Ramette and Tiedje, 2007), important (Zhou et al., 2008), or dominant factor (Ge et al., 2008) relative to local environmental factors.

Recent studies suggest a consensus that the microbial community structure and diversity correlate significantly with soil pH across local (Rousk et al., 2010), regional (Hollister et al., 2010) and continent (Fierer and Jackson, 2006; Lauber et al., 2009; Chu et al., 2010) scales. However, these studies focused on acidic soils, and did not directly test the relative contribution of pH...
and geographic distance to community structure. A meta-analysis showed that sediments are more phylogenetically diverse than any other environment type (Lozupone and Knight, 2007). However, limited evidence exists for spatial patterns in sediments, although directional would be expected to be lower than in soils (Martiny et al., 2006). Bacteria also use different strategies to survive acidic and alkaline conditions (Oilworth and Glenn, 1999; Yohannes et al., 2004). In the present work, we, therefore, examine alkaline lake sediments, testing whether predictable relationships between bacterial communities and factors such as pH and geographic distance mirror those observed in soils.

The Tibetan Plateau is the Earth’s largest (2 × 106 km2) and highest (average ~4500 m a.s.l.) plateau. It contains several thousand saline and alkaline lakes, and is remote from centers of human activity. The region has a dry climate, and salinity increases from south to north due to decreased annual precipitation (Yang et al., 2003). The Tibetan Plateau thus provides an excellent natural laboratory for studying microbial distribution patterns on a regional scale. Several studies have attempted to characterize microbial communities in individual hypersaline lake sediments based on techniques with limited resolution, phospholipid fatty acid and/or 16S rDNA clone libraries (Dong et al., 2006; Jiang et al., 2007). The higher throughput permitted by pyrosequencing allows us to better characterize the diversity in this understudied system and to test the direct effects of geographic distance and environmental factors. We collected pristine sediment samples from 15 alkaline lakes across the Tibetan Plateau, and used barcoded pyrosequencing to evaluate the bacterial communities with respect to three broad and related aims: (i) to explore the taxonomic diversity of the bacteria on Tibetan Plateau alkaline lake sediments, (ii) to determine key factors in shaping the bacterial communities distribution, environmental heterogeneity, geographic distances, or both? and (iii) to quantify their relative importance to bacterial community variation.

Results

Sediment geochemical characteristics

The major geographical and physicochemical characteristics of the lake sediments are summarized in Table S1. Sediment pH varied from 6.88 to 10.37. Pairwise distances between sampling sites ranged from 4 to 1670 km (Table S2). There was no autocorrelation (P = 0.862) observed between sediments pH and geographic distance. While C/N ratios were significantly correlated with altitude (r = 0.561, P = 0.019), and pH was positively (r = 0.506, P = 0.038) correlated with Mg2+.

Distribution of taxa and phylotypes

Across all sediment samples, we obtained 139,714 quality sequences in total and 4173–19,050 sequences per sample (mean = 8218), and were able to classify 79.8% of those sequences. The dominant phyla across all sediments were Bacteroidetes, Firmicutes, Gammaproteobacteria, Deltaproteobacteria, Alphaproteobacteria, Actinobacteria and Chloroflexi (relative abundance > 5%), accounting for more than 61% of the bacterial sequences (Fig. 1). In addition, Chloroflexi, Cyanobacteria, Euryarchaeota, Deinococcus-Thermus and Acidobacteria were present in most sediments at low abundance, and 15 other rarer phyla were identified (Table S3). Interestingly, 2.5% of the detected sequences were affiliated to Halobacteria in the Archaea (5.3% of classified OTUs). Alpha diversity, measured as the number of OTUs (~97% similarity) detected per 4000 sequences was negatively correlated with sediment pH (r = −0.706, P = 0.002). Thus observed OTU richness doubled from high diversity (c. 2239 OTUs) at pH 6.88 and low diversity (c. 1174 OTUs) at pH 10.37 (Fig. 2A).

This observation was confirmed using phylogenetic diversity, also negatively (r = −0.731, P = 0.001) correlated with pH (Fig. 2B), but not correlated with other sediment or site factors (P > 0.10 in all cases, Table S4), including salinity (Fig. S1). In contrast, the distribution of Archaea was not significantly correlated with pH (Fig. S2), indicating that bacterial community members are more affected by sediment pH than archaeal community members. This observation confirms patterns observed in a recent study, in which archaeal and bacterial communities responded differently to environmental gradients in hypersaline lake sediments (Swan et al., 2010).

Bacterial community structure

Sediment pH strongly affected bacterial community structures across the gradient. The PCA (principal component analyses) biplot clearly shows that bacterial communities of different pH align along RDA1 axis (Fig. 3). This interpretation was supported by correlating bacterial community UniFrac distance with difference in sediment pH (P = 0.001, Mantel test). Other measured physical and geochemical factors, including salinity, altitude, C/N ratio of the samples, did not significantly improve the model over pH alone. The effect of sediment pH was evident even at a very coarse level of taxonomic resolution because the relative abundances of the dominant bacterial phyla (e.g. Alphaproteobacteria and Deltaproteobacteria), rarer phyla (e.g. Cyanobacteria), and classes (e.g. Chromatiales) were significantly correlated across the pH gradient (Fig. 4), although sometimes in opposite directions. For
example, the relative abundance of Deltaproteobacteria decreased as pH increased, but Alphaproteobacteria increased. Some ions that correlate with pH, such as Na$^+$ and Mg$^{2+}$, showed significant correlations with the relative abundance of Bacteroidetes, Firmicutes or Cyanobacteria. Sediment TC (total carbon) and TP (total phosphorus) were significantly ($P < 0.05$ in all cases) correlated with the relative abundances of Acidobacteria, Firmicutes or Nitrospira, (Fig. 5). Taken together, the results strongly demonstrated that local sediment pH is, directly or indirectly,
shaping the bacterial community structure among sites across the Tibetan Plateau lake sediments.

The samples contained 1166–2303 OTUs (phylotypes) per 4000 sequences (Fig. 2A); 49.0%–84.3% of phylotypes were unique to each sample, and few phylotypes overlapped in any pair of samples (0.24–12.27%, average 2.95%) (Table S5), indicating strict site endemism. In contrast, duplicate samples collected from the same lake shared more phylotypes: for example, KZC1 and KZC2 shared 11.5% of the OTUs, and those between LBC1 and LBC2 were 5.4% (Table S5). The unweighted UniFrac distances of bacterial communities (beta diversity) were significantly \((P = 0.030)\) correlated with the geographic distance (Fig. 6). Partial Mantel tests further supported the distance effect when the effect of sediment environmental similarity matrix was held constant \((P = 0.042)\). Together, the results revealed that spatial isolation affects community structure, presumably via dispersal, at this level of spatial and phylogenetic resolution.

**Linking bacterial communities to sediment properties and geographic distance**

Variance partitioning analysis (VPA) was performed to quantify the relative contributions of geographic distance and sediment environmental parameters to the taxonomic structure of the bacterial communities. A subset of environmental parameters (pH, Na\(^+\), Mg\(^{2+}\) and C/N ratio) was selected by the BioEnv procedure, which provides the highest Pearson correlation with bacterial communities. The combination of selected sediment properties and geographic distance showed a significant \((P = 0.014)\) correlation with the bacterial community structures. These variables explained 41.1% of the observed variation, leaving 58.9% of the variation unexplained. The sediment properties explained 28.9% \((P = 0.003)\), and geographic distance alone explained 12.2% \((P = 0.029)\) variations, and no interaction effect was detected (Fig. 7). Although the sediment properties together explained more of the variation, geographic distance by itself explained 12.2% of the variation observed, more than any of the other 4 of the individual sediment variables (Fig. 7). Thus both sediment properties and geographic distance are important factors in shaping bacterial community structures.

**Discussion**

Collecting sediment from pristine sites usually involved long drives over unfinished roads, making sampling hazardous and expensive. Consequently, most studies of microbial community composition in lake sediments have been limited to single lake (Jiang et al., 2007; Swan et al., 2010), preventing large-scale tests of bacterial spatial patterns in this habitat type.

In this study, we found that Bacteroidetes and Firmicutes were the main phyla in lake sediments, and high
relative abundances of Gammaproteobacteria and Deltaproteobacteria were observed (Table S3); in contrast, these phyla are rare in soils (Chu et al., 2010; Griffiths et al., 2011), indicating that bacterial communities in sediments are highly distinctive relative to soils (Lozupone and Knight, 2007).

The fundamental role of soil pH in shaping bacterial community structure has recently been demonstrated across North America (Lauber et al., 2009), in the Arctic (Chu et al., 2010), and across British soils (Griffiths et al., 2011). However, the samples examined in these studies were acidic to neutral, leaving open the question of whether pH also dominates the bacterial distribution in alkaline sediments, especially because some individual species can grow in environments spanning 4 pH units (Rosso et al., 1995). Here, we showed that the bacterial community structure, phylotype richness and phyloge-
netic diversity were primarily correlated with a single parameter, sediment pH (Figs 2 and 3, Table S4), although there was substantial variation in bacterial community structure across the range of surveyed sediments. The pH was the main factor controlling the bacterial community structure even within a limited range of 2.4 pH units if regardless the site pH 6.88 (Fig. 3), and nutrient concentrations were not significantly associated (Table S4). In addition, the ways in which specific phyla change across this pH gradient are similar to pH responses observed in soils, suggesting that they may be general across habitats. For instance, the relative abundance of Alphaproteobacteria has been shown to increase with higher pH (Chu et al., 2010; Rousk et al., 2010). The results imply that pH is a strong, universal predictor of bacterial community structure both for acidic soils and for alkaline lake sediments.

The distribution pattern of some phyla differed from previous observations. For example, previous work found Acidobacteria to be the dominant phylum across acidic soils (Lauber et al., 2009; Chu et al., 2010), but this phylum accounts for just 1.04% (average, Table S3) of the sequences in the current study. This result was expected, as many groups of Acidobacteria are more abundant at low pH (Griffiths et al., 2011) and the sediments in the present study are alkaline. Similar strong significant correlations between the relative abundance of Acidobacteria and Bacteriodetes and pH have been reported in soils from different biomes (Lauber et al., 2009; Chu et al., 2010), but were not detected in the Tibetan Plateau lake sediments. It should be noted that importance of pH in structuring bacterial communities does not completely erase the influence of other sediment or local features in driving the distribution of particular bacterial phyla. Indeed, we observed that TC and TP, Na⁺ and Mg²⁺, were significantly correlated with some dominant or subdominant phyla, matching the results obtained in earlier studies (Hollister et al., 2010; Griffiths et al., 2011). In particular, the relative abundances of Bacteriodetes and Acidobacteria were negatively correlated with Na⁺ and TC respectively. Na⁺ ions are critical to alkaliphilic bacteria (e.g. Bacteriodetes) as proton replacements to cope with high external pH (Skulachev et al., 1999). These phyla also play important roles in soil, where their relative abundances correlate with carbon mineralization rate (Fierer et al., 2007). Therefore, the changes in these dominant taxa over the pH gradient likely have consequences for ecosystem functions.

Salinity displays a strong gradient along our sampling sites. Given the key role of salinity in shaping bacterial community both in lake water (Wu et al., 2009) and globally (Lozupone and Knight, 2007), we expected to see a strong influence of salinity on the microbial communities in these sediments. However, there was no significant correlation between lake water salinity and bacterial diversity (Fig. S1, Table S4). Although the salinity of surface sediment was not directly measured, it is typically correlated with the lake water salinity because of its near water saturation. Similarly, Jiang and colleagues (2007) showed that the succession of proteobacterial groups along the salinity gradient was typically observed in free-living bacterial communities but not in sediments, and a recent study performed in hypersaline soils and sediments showed that site pH and organic carbon rather than salinity were the primary factors in controlling the bacterial structure on this local scale (Hollister et al., 2010). Thus,
the main factors driving bacterial community structure in sediments differ in the details from those in lake water and soil. Our results emphasize the requirement of comparing the bacterial distribution pattern among local soils, lake water and sediments simultaneously in future work.

An issue relevant to our study, and to many other biogeographic studies, is whether spatial distance correlates strongly with genetic variation (Ramette and Tiedje, 2007; Ge et al., 2008). Here, we employed a more powerful phylogenetic approach that takes into account different levels of similarity between different pairs of taxa (Lozupone and Knight, 2005) to compare more subtle differences among the samples. Consequently, a significant correlation was observed between bacterial community variation and geographic distance (Fig. 6). Meanwhile, there was little taxonomic overlap across the sites (Table S5). These results suggest that geographic distance is an indicator of bacterial dispersal among sediments. The geographic distance explained more variation in community structure than any other individual factor (Fig. 7). The unexplained variation may be due to additional factors not measured in this study, such as sediment temperature and redox state (Pett-Ridge and Firestone, 2005), which can influence bacterial communities.

In contrast, a recent continent-scale survey of soils showed that geographic distance was not correlated with changes in community structure (Chu et al., 2010), whereas Ge and colleagues observed 60.3% of the variation among communities was explained by spatial isolation (Ge et al., 2008). The former study was performed in an open environment, where wind, animal and other vectors may spread the bacteria, leading to random spatial distributions (Green and Bohannan, 2006); another similar study also showed that when the geographic distance increased, the correlation between community dissimilarity and geographic distance disappeared (Griffiths et al., 2011). Similarly, no ammonia-oxidizing bacterial diversification was detected in salt marsh sediments across the continental scale (Martiny et al., 2011). However, the latter study only collected samples from three locations (Ge et al., 2008), limiting its statistical power, especially because the relative influence of historical and environmental factors depends on the sampling scale (Rickels, 2004; Martiny et al., 2006). A small-scale (0.03 m to 300 m) survey in salt marsh sediments indicated that, at least on this scale and in this system, the bacterial taxon–area relationship was primarily driven by environmental heterogeneity (Horner-Devine et al., 2004); similar results were obtained in a gradient of hypersaline lake sediments along a 140 m transect (Hollister et al., 2010). Therefore, studies need to be performed across a range of spatial scales and in a range of systems, as is being performed in projects such as the Earth Microbiome Project (Gilbert et al., 2010).

In conclusion, this study represents a large attempt to investigate bacterial spatial patterns in pristine sediments across the Tibetan Plateau. We show that bacterial community structure, phylotype richness and phylogenetic diversity are predicted by sediment pH, even within a narrow pH range. This is the first study to demonstrate that pH predicts microbial community structure in alkaline sediments, mirroring results in acidic soils. Our results clearly show that local geochemical features are the dominant factors in driving the bacterial community variation, whereas geographic distances as a single factor also a larger contributor in community variation across a regional spatial scale (about 1670 km). From a practical point, future research in this area would benefit from using appropriate sampling scale and relative less disturbed ecosystem to study microbial biogeography (Green et al., 2004; Ramette and Tiedje, 2008). These observations thus provide baseline information for predicting regional-scale responses to future environmental changes.

**Experimental procedure**

**Sample collection**

The Tibetan Plateau lies at a critical and sensitive junction of three climatic systems: the East Asian Monsoon, the cold polar airflow from the Siberian high pressure, and the Indian monsoon (Jiang et al., 2007). The ages of lakes located at the Tibetan Plateau were estimated to be 2 to 8 million years (Zheng and Yao, 2004), sediments in lake record past climatic and environmental changes.

In the summer of 2010, we collected 17 surface sediments (0–5 cm) from 15 lakes across the Tibetan Plateau, and duplicate samples were collected from two lakes to compare local variances of bacterial community. The lakes were chosen to cover a salinity gradient from 0.32 g l⁻¹ (0.03%, freshwater) to 308.0 g l⁻¹ (30.8%, hypersaline). The sediment samples were mixed thoroughly and immediately shipped to lab on ice packs. Subsamples were archived at 4°C for sediment geochemical characterization and ~80°C for genomic DNA extraction. GPS coordinates recorded at each sampling point (varied from 29°13′–34°36′N, 79°47′–96°47′E), imported into the NOAA website (http://www.nhc.noaa.gov/gccalc.shtml) to calculate the pairwise geographic distance. The pH values were measured using pH strips with 1:1 (wt/vol) of sediment to water. Sediments were immediately freeze-dried to detect geochemical characteristics. Concentrations of sodium (Na⁺) and magnesium (Mg⁺) ions and TP were measured using inductively coupled plasma mass spectrometry (ICP-MS) (ELAN 9000/DRC-e, PerkinElmer, USA). TC and total nitrogen (TN) were measured by CN Analyzer (Vario Max CN, Elementar, Germany).

**DNA extraction and purification**

Community DNA was extracted from sediments using a FastDNA® Spin kit (Bio 101, Carlsbad, CA, USA) according to the manufacturer’s protocol. The raw DNA was purified using...
a 0.8% (w/v) low melting point agarose gel. The DNA bands were excised, then extracted using an agarose gel DNA purification kit (TaKaRa), and quantified with a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA). Purified DNA was stored at −20°C until use.

**Bacterial 16S rRNA amplification and 454 sequencing**

An aliquot (50 ng) of purified DNA from each sample was used as template for amplification; the V4–V5 hypervariable regions of bacterial 16S rRNAs (Escherichia coli positions 515–907; Biddle et al., 2008) were amplified using the primer set: F515: GTGCCAGCMGCCGCGG with the Roche 454 ‘A’ pyrosequencing adapter and a unique 7 bp barcode sequence, and primer R907: GGTTCAACCTTGTTATGATGT with the Roche 454 ‘B’ sequencing adapter at the 5′-end of each primer respectively. Each sample was amplified in triplicate with 50 μl reaction under the following conditions: 30 cycles of denaturation at 94°C for 30 s, annealing at 55°C for 30 s, and extension at 72°C for 30 s; with a final extension at 72°C for 10 min. PCR products were pooled together and purified by Agarose Gel DNA purification kit (TaKaRa) as described above.

The PCR products from each sample were combined in equimolar ratio in a single tube and to run on a Roche FLX 454 pyrosequencing machine (Roche Diagnostics Corporation, Branford, CT, USA), producing reads from the forward direction F515.

**Processing of pyrosequencing data**

Data were processed as described previously (Fierer et al., 2008; He et al., 2010), using the Quantitative Insights Into Microbial Ecology (QIIME) pipeline (http://qiime.sourceforge.net/) (Caporaso et al., 2010). Specifically, bacterial sequences with the same barcode were assigned into the same sample, and then the barcode and primer sequences were removed. Only the first 350 bp after primer-F515 of each sequence were included to improve reads quality for further analysis. Bacterial phylotypes were identified using uclust (Edgar, 2010) and assigned to operational taxonomic units (OTUs, 97% similarity). Representative sequences from each phylotype were aligned using PyNAST (DeSantis et al., 2006a), and the most abundant sequence in the OTU was selected as the representative sequence. Taxonomic identity of each phylotype was determined using the Greengenes database (DeSantis et al., 2006b). To correct for sampling effort, we used a randomly selected subset of 4000 sequences per sample to calculate distances between samples. The differences in overall community composition between each pair of samples were determined using the UniFrac metric (Lozupone and Knight, 2005), which provides a more robust index of community distance than taxon-based methods.

**Statistical analysis**

Phylogenetic diversities were estimated using Faith’s index, which incorporates the phylogenetic breadth across taxonomic levels (Faith, 1992). The relationships between the taxonomic diversity for the group with geochemical features were tested with linear regression analyses using spss 17.0 for Windows. Raw environmental data were standardized to make the different environmental factors comparable. Euclidean distances were used to construct a similarity matrix for sediment properties. Partial Mantel tests were used to calculate the correlation between the UniFrac distances of bacterial communities and the sediment characteristics or geographic distance in PASSaGE (Rosenberg and Anderson, 2011). BioEnv and canonical correspondence analysis (CCA) were also used to identify the abiotic factors most important to bacterial community composition, and they were used to construct the sediment property matrix for variation partitioning analysis in R v.2.8.1 with the vegan package (R Development Core Team, 2008).

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Supporting information

Additional Supporting Information may be found in the online version of this article:

Fig. S1. Lake water salinity in relative to bacterial phylotype richness (a) and phylogenetic diversity (b) based on 97% sequence similarity in Tibetan Plateau lake sediments. The communities were randomly sampled at the 4000 sequences level.

Fig. S2. Sediment pH in relative to Archaea phylotype richness based on 97% sequence similarity in Tibetan Plateau lake sediments. The communities were randomly sampled at the 4000 sequences level.

Table S1. Sediment physical and chemical properties and lake water salinity.

Table S2. Pairwise geographic distance (km) matrix.

Table S3. Relative average abundances of phyla across all sediments and sediments grouped into various pH categories (values represent % of total non-redundant sequences). Asterisks indicate sequences classified to the domain Bacteria, but not to a specific phylum.

Table S4. Correlations (r) between bacterial phylotype richness and phylogenetic diversity and sediment and site characteristics and lake water salinity. Values in bold indicate significant correlations (*P* < 0.05).

Table S5. The detected OUTs, percentages of overlap, unique and diversity indices for each sample per 4000 sequences.

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