

## ORIGINAL ARTICLE

# The microbial gene diversity along an elevation gradient of the Tibetan grassland

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Tibet is one of the most threatened regions by climate warming, thus understanding how its microbial communities function may be of high importance for predicting microbial responses to climate changes. Here, we report a study to profile soil microbial structural genes, which infers functional roles of microbial communities, along four sites/elevations of a Tibetan mountainous grassland, aiming to explore the potential microbial responses to climate changes via a strategy of space-for-time substitution. Using a microarray-based metagenomics tool named GeoChip 4.0, we showed that microbial communities were distinct for most but not all of the sites. Substantial variations were apparent in stress, N and C-cycling genes, but they were in line with the functional roles of these genes. Cold shock genes were more abundant at higher elevations. Also, *gdh* converting ammonium into urea was more abundant at higher elevations, whereas *ureC* converting urea into ammonium was less abundant, which was consistent with soil ammonium contents. Significant correlations were observed between N-cycling genes (*ureC*, *gdh* and *amoA*) and nitrous oxide flux, suggesting that they contributed to community metabolism. Lastly, we found by Canonical correspondence analysis, Mantel tests and the similarity tests that soil pH, temperature,  $\text{NH}_4^+-\text{N}$  and vegetation diversity accounted for the majority (81.4%) of microbial community variations, suggesting that these four attributes were major factors affecting soil microbial communities. On the basis of these observations, we predict that climate changes in the Tibetan grasslands are very likely to change soil microbial community functional structure, with particular impacts on microbial N-cycling genes and consequently microbe-mediated soil N dynamics.

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## Introduction

As the roof of the world, the Tibet plateau is considered the third ‘pole’ (Qiu, 2008). Rich in glaciers and permafrost, the region has Earth’s third largest ice store after Antarctic and Arctic (Qiu, 2010). Recent decades have witnessed strong

effects of climatic warming as this region has been experiencing three times the average global warming rate since 1960 (Li and Tang, 1988). It is predicted that the Tibetan alpine tundra could decrease to 20% of its current level with further climate warming (Walker *et al.*, 2001). These observations and *in silico* modeling render Tibet as one of the most threatened regions by climate warming.

Tibetan ecosystems are particularly vulnerable to climate warming, which can partially be attributed to the pronounced role of climate in structuring alpine system (for example short growth season of plants) (Klein *et al.*, 2004) and the snow-albedo or ice-albedo feedback that amplifies climate warming

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(Déry and Brown, 2007). The alpine grassland, accounting for roughly 35% of Tibet plateau, is a representative landscape of the Tibetan region (Cao *et al.*, 2004). Studies have been carried out to analyze vegetation (Klein *et al.*, 2004, 2007) and soil biogeochemical processes (Cao *et al.*, 2001; Hastings *et al.*, 2005; Wang *et al.*, 2009b). However, soil microbial communities remain little understood. This knowledge gap calls for attention as it has been well documented that soil microbes drive a variety of biogeochemical processes such as carbon (C), nitrogen (N), phosphorus (P) and sulfur (S) cycling, and consequently provide critical ecosystem services such as soil fertility and greenhouse gas mitigation (Whitman *et al.*, 1998; Green *et al.*, 2008).

The recent, rapid development of high-throughput metagenomics technologies has enabled scientists to conduct detailed investigation of microbial communities. Although advancements have been achieved (Huber *et al.*, 2007; Pointing *et al.*, 2009; Dick and Tebo, 2010), most studies of this kind focus on microbial taxonomy. However, it is crucial to examine microbial functional signatures such as structural genes relevant to metabolic pathways, energetics and regulatory circuits, which directly address microbial functional potentials related to ecosystem processes. To fulfill this need, a microarray-based metagenomics tool named GeoChip is an excellent choice and has been widely adopted to examine microbial gene diversity in various environments (Van Nostrand *et al.*, 2009; Waldron *et al.*, 2009; Wang *et al.*, 2009a; He *et al.*, 2010; Lu *et al.*, 2012). The most advanced version of GeoChip, GeoChip 4.0 can target a wide array of 152 414 microbial genes belonging to 410 gene families.

In this study, we used GeoChip 4.0 to survey microbial gene diversity along the elevation/temperature gradient of a mountainous grassland of the Tibet plateau. The observational investigations along an elevation/temperature gradient reveal potential microbial responses to climate changes, using a strategy of space-for-time substitution (Dunne *et al.*, 2004). Although there have been numerous studies on profiling microbial distribution at the phylogenetic level (Bryant *et al.*, 2008; Fierer *et al.*, 2011; Wang *et al.*, 2012), there has been no such study at the functional gene level. Here we define microbial genes present in all of the elevations as ubiquitous genes and those present at only one elevation as endemic genes. We are interested in testing the following hypotheses: (i) the functional structure of microbial community differs among elevations, given high soil heterogeneity and variations of vegetation and climatic conditions, and soil and vegetation attributes contribute significantly in shaping the functional structure; (ii) variations of microbial stress genes are consistent with the increasing alpine stress at higher elevations; (iii) variations of C- or N-cycling genes are significantly linked to soil C and N dynamics in the alpine grassland; and (iv) the pattern of endemic gene

abundances not only differs from that of ubiquitous gene abundances but also differs among elevations. The study is based on field investigations of four fenced sites/elevations in a mountainous grassland located at the Haibei Alpine Meadow Ecosystem Research Station of Qinghai province, which has typical climate and vegetation of Tibetan alpine grasslands (Wang *et al.*, 2009b, 2010a, b).

## Materials and methods

### *Site and sampling*

Sampling was carried out at the Haibei Alpine Meadow Ecosystem Research Station in Qinghai province, located in the northeast of Qinghai-Tibet Plateau (37°37'N, 101°12'E) in a valley surrounded by the Qilian Mountains. The local climate is highland continental, characterized to be cold and long in winter but warm and short in summer. The annual mean air temperature and precipitation are  $-1.7^{\circ}\text{C}$  and 560 mm, respectively (Zhao *et al.*, 2006). Soil type at the research sites is dominated by Mat Cryic Cambisols typical of alpine grassland soil, with pH value of 7.3 and 7.4 at depths of 10 cm and 20 cm, respectively. The growing season of vegetation is from May to September, during which more than 80% of the precipitation occurs. Vegetation reaches its maximal aboveground biomass level in late July and early August and starts withering in early October. Approximately 87% of vegetation species at the sites use the  $\text{C}_3$  photosynthetic pathway (Zhao *et al.*, 2006).

The study sites, as part of a comprehensive project to examine the effects of artificial warming and/or livestock grazing on Tibetan grasslands, were initiated at four sites/elevations of 3200, 3400, 3600 and 3800 m in May 2006. The spatial distances between adjacent elevations are 6.2 km (3200–3400 m), 4.2 km (3400–3600 m) and 1.3 km (3600–3800 m), respectively. These sites were selected for their representative vegetation and soil attributes for the elevations. The plant community at 3200 m is dominated by *Kobresia humilis*, *Elymus nutans*, *Stipa aliena*, *Potentilla anserine* and *Thalictrum alpinum*. The plant community at 3400 m consists primarily of a species of hardy deciduous flowering shrub (*Potentilla fruticosa*) with 50–60% total coverage and beneath grass species of *Kobresia humilis*, *Elymus nulan* and *Festuca ovina* with 80% total coverage. The plant community at 3600 m site is dominated by *Kobresia humilis*, *Potentilla nivea*, *Thalictrum alpinum*, *Carex atrofusca*, *Poa crymophila* and *Potentilla fruticosa*. The plant community at 3800 site is dominated by *Kobresia humilis*, *Poa crymophila*, *Androsace mariae*, *Polygonum macrophyllum* and *Kobresia pygmaea*. Owing to the alpine environment, the plant communities are characterized by short growth period, low primary production and diversity.

At each elevation, three plots comprising typical vegetation and soil attributes were fenced to minimize disturbance. The dimensions of the plots were 1.0 m, 1.0 m and 0.3 m in length, width and height, respectively. The distance between adjacent plots was roughly 0.6 meter. In August 2009, soil samples were collected from all of the plots. Five soil cores at a depth of 0–20 cm and diameter of 1.5 cm were randomly taken from each plot and mixed thoroughly to generate a soil sample representing the plot. Soil samples were kept on ice when transporting to laboratory and sieved with 2 mm mesh to remove roots and stones. Soil samples were preserved at  $-80^{\circ}\text{C}$  before DNA extraction.

#### *DNA extraction, purification and quantitation*

Soil genomic DNA was extracted using a FastDNA spin kit for soil (MP Biomedical, Carlsbad, CA, USA) following the manufacturer's instructions. To purify it, the DNA extract was mixed with 2.5 volume of 100% ice-cold ethanol and 0.1 volume of 3 M NaOAc (pH 5.2) before overnight incubation at  $-20^{\circ}\text{C}$ . DNA was precipitated by centrifugation for 30 min at 13 000 *g*. Then supernatant was decanted and washed with 1 ml of 70% ethanol. DNA was air-dried and dissolved in 50  $\mu\text{l}$  of nuclease-free water. DNA quality and quantity were measured using NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies Inc., Wilmington, DE, USA) and with PicoGreen (Ahn *et al.*, 1996) using a FLUOstar Optima (BMG Labtech, Jena, Germany), respectively.

#### *GeoChip 4.0 experiments*

As previously described (Yang *et al.*, 2013), DNA samples were labeled with the fluorescent dye Cy-5 using a random priming method and purified using the QIA quick purification kit (Qiagen, Valencia, CA, USA). Then DNA was dried in a SpeedVac (ThermoSavant, Milford, MA, USA) at  $45^{\circ}\text{C}$  for 45 min. The hybridization was carried out at  $42^{\circ}\text{C}$  for 16 h on a MAUI hybridization station (BioMicro, Salt Lake City, UT, USA). After purification, GeoChip microarrays were scanned by a NimbleGen MS200 scanner (Roche, Madison, WI, USA) at 633 nm using a laser power and photomultiplier tube gain of 100% and 75%, respectively.

#### *Data analyses*

Signal intensities were quantified and processed using the data analysis pipeline as previously described (Yang *et al.*, 2009; He *et al.*, 2010; Yang *et al.*, 2013). Then processed GeoChip data were analyzed using the following steps: (i) remove the poor quality spots, which were flagged as 1 or 3 by ImaGene (Arrayit, Sunnyvale, CA, USA) or with a signal to noise ratio of less than 2.0; (ii) normalize the signal intensity of each spot by dividing the

signal intensity by the total intensity of the microarray followed by multiplying by a constant; (iii) transform the data to the natural logarithmic form; and (iv) remove genes detected in only one out of three samples from the same elevation.

Principal component analysis was used to determine overall functional changes in the microbial communities. Bray-Curtis distance was used to obtain dissimilarity matrices in the adonis algorithm of the dissimilarity test for comparing GeoChip data of four elevations. All the analyses were performed using functions in the Vegan package (v.1.15-1) in R v. 2.8.1 (RDC Team, 2006).

The similarity test, Mantel test, Canonical correspondence analysis (CCA) and Variation partitioning analysis were used to evaluate the linkages between microbial gene compositions and environmental attributes. In the similarity test, Euclidean distance was used to calculate the distance between samples, followed by calculation of Pearson correlation coefficient. To select attributes in CCA modeling, we used variation inflation factors to examine whether the variance of canonical coefficients was inflated by the presence of correlations with other attributes. If an attribute had a variation inflation factor value larger than 20, we deemed it to be affected by other attributes and consequently removed it from the CCA modeling. The Variation partitioning analysis analysis resulted in five soil attributes (soil pH, moisture,  $\text{NH}_4^+-\text{N}$ ,  $\text{NO}_3^--\text{N}$  and soil temperature at the depth of 5 cm) and two vegetation attributes (Shannon diversity and total coverage of vegetation). As temperature was measured per site, one way analysis of variance was not conducted. All the analyses were performed using functions in the Vegan package (v.1.15-1) in R v. 2.8.1 (RDC Team, 2006).

#### *Soil and vegetation attribute measurements*

At the time of soil sampling, soil temperature was measured by type-K thermocouples (Campbell Scientific, Logan, UT, USA) coupled to a CR1000 datalogger at depths of 5, 10 and 20 cm. Soil moisture at depths of 5 and 10 cm were recorded by time domain reflectometry (Model Diviner-2000, Sentek Pty Ltd., Stepney, SA, Australia).

In the lab, total soil organic C and N were quantified by a TOC-5000 A analyzer (Shimadzu Corp., Kyoto, Japan) and a Vario EL III Elemental Analyzer (Elementar, Hanau, Germany), respectively. To measure  $\text{NH}_4^+-\text{N}$  and  $\text{NO}_3^--\text{N}$ , 10 g of fresh soil was suspended in 50 ml of  $2\text{ mol l}^{-1}$  KCl solution, then shaken at room temperature for 1 hour and placed for 30 min under unstirred conditions. Subsequently, clear supernatant was passed through a filter paper of 30–50  $\mu\text{m}$  pore size and analyzed using continuous flowing SAN<sup>++</sup> Analyzer (SKALAR, Breda, the Netherlands).

The greenhouse gases ( $\text{CO}_2$ ,  $\text{N}_2\text{O}$  and  $\text{CH}_4$ ) were measured by chamber per plot setting at the 5 cm

depth of soil as previously described (Wang *et al.*, 2009b). During sampling, chambers were closed for half an hour. Hundred milliliters of air samples were taken by plastic syringes every 10 min and transported to the lab. Greenhouse gas concentrations were measured by gas chromatograph (Agilent 4890D, Agilent Co., Santa Clara, CA, USA) equipped with a flame ionization detector and an electron capture detector as previously described (Yuesi and Yinghong, 2003). As there were many rat holes in the soil at the elevation of 3600 m, the measurement of the greenhouse gas fluxes data ( $\text{CO}_2$ ,  $\text{CH}_4$  and  $\text{N}_2\text{O}$ ) was unreliable. Therefore, we excluded those points in the correlation analysis with the greenhouse gas fluxes.

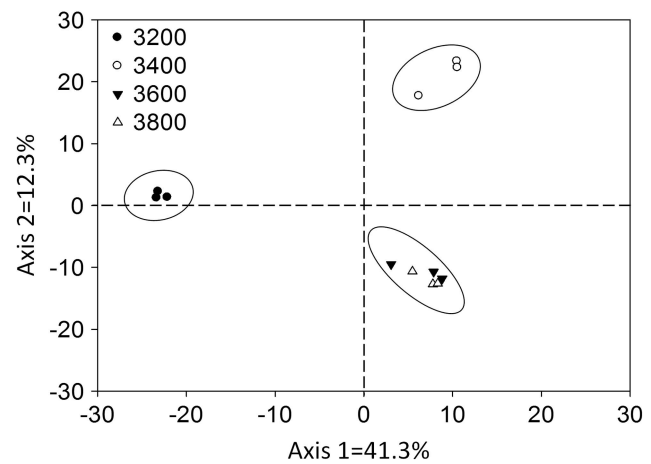
One quadrant of  $1 \times 1$  m was selected at each plot to measure vegetation attributes. The vegetation attribute indices included vegetation species number, diversity, total coverage and aboveground biomass, which were all recorded according to established protocols (Klein *et al.*, 2007). Vegetation was weighted instantly for aboveground biomass data after mowing. Vegetation diversity was calculated by Shannon index.

## Results

### *Distribution patterns of soil microbial communities along the elevation gradient*

A total of 49 520 genes were detected by GeoChip 4.0. Detected genes were the fewest at the 3200 m site (23 369 genes) and roughly doubled at the 3400 m site (45 685 genes). Slightly fewer genes were detected at higher elevations than those at the 3400 m site (Supplementary Table S1). Alpha-diversities, as shown by Shannon and Simpson indices, varied concurrently with gene numbers. Alpha-diversities differed significantly ( $P < 0.05$ ) among 3200, 3400 and 3600 m sites, which was supported by the dissimilarity test using adonis algorithm (Supplementary Table S2). However, microbial communities at 3600 and 3800 m sites appeared to be similar by the statistical test ( $P > 0.05$ ). Consistently, principal component analysis with the whole GeoChip data set showed that samples of 3200 and 3400 m sites were well separated from the others, but samples of 3600 and 3800 m sites were similar (Figure 1).

Gene overlap between elevations was also calculated. Higher percentages of shared genes were observed among 3400, 3600 and 3800 m sites (81.1–83.8%) (Supplementary Table S3), whereas only 50.3–53.4% of the genes were shared between 3200 m and other sites. The highest percentages of shared genes (82.7–92.3%) were observed for the plots within the same elevation (data not shown), suggesting that the within-elevation variation was small.



**Figure 1** Principle component analysis (PCA) of GeoChip data. The values of Axis 1 and 2 are percentages of total variations that can be attributed to the corresponding axis.

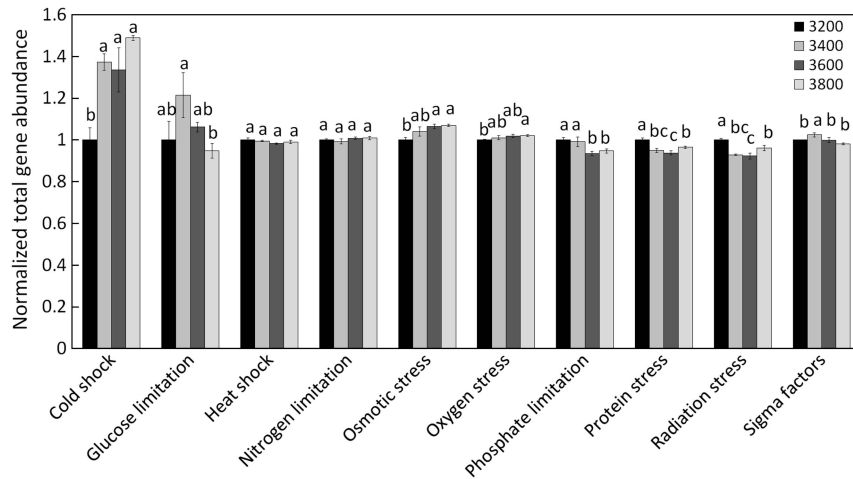
### *Selected gene categories*

Genes involved in stress response, N and C cycling were examined below. Only selected genes were presented or described.

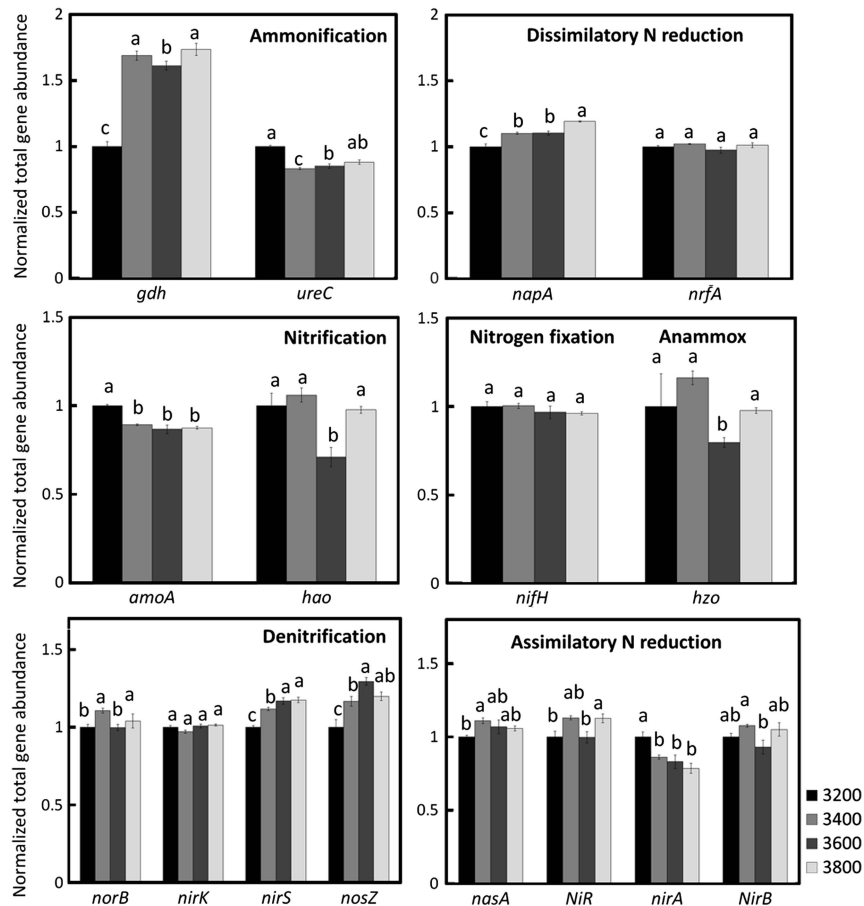
**Stress response.** Considerable variability of stress genes was observed (Figure 2). Cold shock genes, *CspA*, *cspB*, *desK* and *desR* genes, which are cold shock genes required for adaption to cold environment (Ermolenko and Makhatadze, 2002), were more abundant at higher elevations. In contrast, heat shock genes (*dnaK*, *groEL*, *groES*, *grpE* and *hrcA*) remained unchanged.

**N cycling.** N-cycling genes differed substantially among the four elevations (Figure 3). For example, total abundances of *ureC* and *gdh* genes involved in ammonification, *napA* genes involved in dissimilatory N reduction and *nosZ* genes involved in denitrification exhibited distinct differences for each site, whereas no significant difference was observed for the abundances of *nifH* and *nrfA* genes. Notably, changes of *ureC* and *gdh* abundances were largely reverse, which was consistent with their opposite functional role in N mineralization. *nirS* and *nosZ* genes involved in denitrification were more abundant at higher elevations but *amoA* gene involved in nitrification was less abundant. Consistently, soil  $\text{NO}_3^-$ -N and  $\text{NH}_4^+$ -N contents were increased and decreased at higher elevations, respectively (Supplementary Table S4).

Grassland nitrous oxide ( $\text{N}_2\text{O}$ ) emission is a microbe-mediated process. Thus, we explored whether N-cycling genes were correlated with  $\text{N}_2\text{O}$  flux. The results showed that total abundances of *ureC* and *amoA* were positively correlated ( $P < 0.01$ ) with  $\text{N}_2\text{O}$  flux, whereas those of *gdh* ( $P < 0.01$ ), *nosZ* ( $P < 0.01$ ) and *nirS* ( $P < 0.1$ ) were negatively correlated (Figure 4).



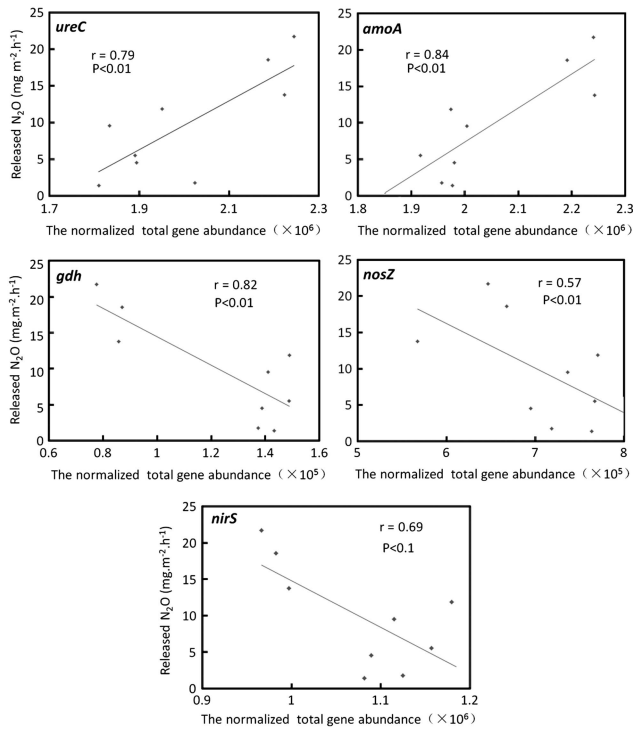
**Figure 2** Total gene abundances, normalized to the abundance at 3200 m, of stress gene categories at each elevation. All data are presented as mean  $\pm$  s.e. calculated from biological triplicates. Significant ( $P < 0.05$ ) differences among elevations are indicated by alphabetic letters above the bars.



**Figure 3** Total gene abundances, normalized to the abundance at 3200 m, of N genes at each elevation. All data are presented as mean  $\pm$  s.e. calculated from biological triplicates. Significant ( $P < 0.05$ ) differences among elevations are indicated by alphabetic letters above the bars.

*C* cycling. Variability was apparent for C cycling genes (Supplementary Figure S1). For C-fixation genes, the abundance of *rubisco* (ribulose-1, 5-bisphosphate carboxylase/oxygenase) was the

lowest at 3200 m, but no other C-fixation genes exhibited similar patterns. For C degradation (starch, hemicellulose, cellulose, aromatics, chitin and lignin degradation), isopullulanase and



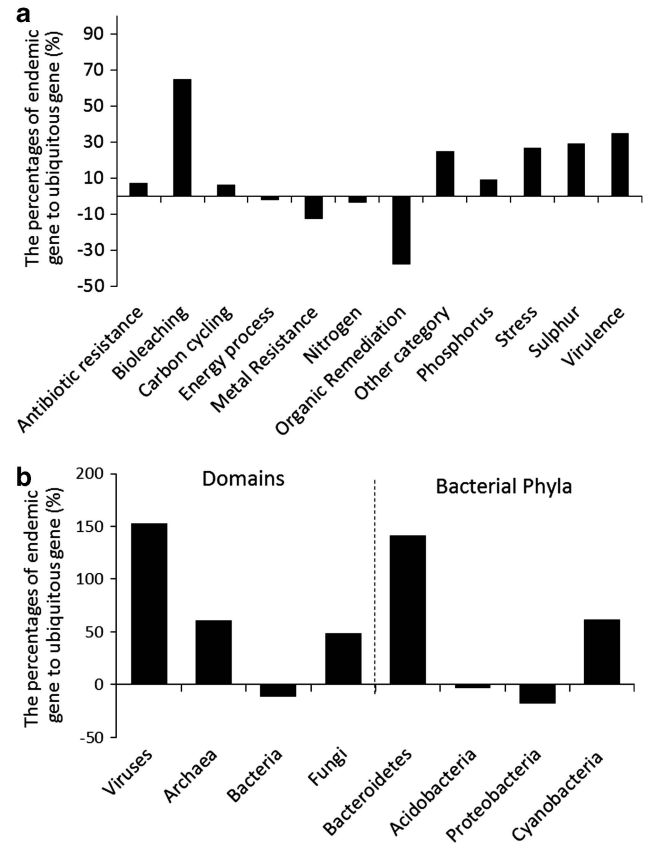
**Figure 4** Correlations between N<sub>2</sub>O flux and the abundance of N-cycling genes.

exoglucanase were more abundant at higher elevations, whereas exochitinase and *glx* were less abundant. For methane (CH<sub>4</sub>) cycling, our results indicated that methane-producing gene *mcrA* and methane-oxidizing gene *mmoX* were more abundant at higher elevations, whereas methane-oxidizing gene *pmoA* remained unchanged. Although these results might suggest that functional potentials of CH<sub>4</sub> cycling were enhanced, no correlation was observed between CH<sub>4</sub> flux and abundances of *mmoX* ( $r = -0.05$ ,  $P > 0.11$ ), *mcrA* ( $r = -0.10$ ,  $P > 0.25$ ) and *pmoA* ( $r = -0.43$ ,  $P > 0.84$ ) genes. Similarly, there was no correlation between CO<sub>2</sub> flux and abundances of C-degradation genes ( $r = 0.29$ ,  $P > 0.36$ ).

#### Ubiquitous and endemic genes

A total of 21842 microbial genes were present in all of the four sites, whereas 5962 genes were present at only one site. That is, ubiquitous and endemic genes accounted for 44.1% and 12.0% of total detected genes, respectively (Supplementary Figure S2). Ubiquitous genes were predominantly derived from the phyla of *Proteobacteria*, *Actinobacteria* and *Firmicute*, which was consistent with observations in a wide range of soil types that these bacteria were prevalent in soil (Roesch *et al.*, 2007; Monroy *et al.*, 2012; Yergeau *et al.*, 2012).

The average abundance of endemic genes was roughly 33% of that of ubiquitous genes, suggesting



**Figure 5** Comparison of percentage changes of (a) each gene category of endemic genes to that of ubiquitous genes and (b) microbial domains or phyla of endemic genes to those of ubiquitous genes.

that endemic genes tended to be rare genes that were typically low in abundance. Several gene categories enriched in the endemic genes (Figure 5a). Among them, there were a total number of 1718 endemic stress genes, including 151  $\sigma^{24}$  genes responsive to extreme temperature stress, 122 *glnA* genes encoding a glutamine synthetase required for N-limiting conditions (Merrick and Edwards, 1995) and 162 *fnr* genes encoding a transcriptional activator required for oxygen-limiting conditions (Spiro and Guest, 1987).

It was also noted that endemic genes enriched genes derived from viruses, archaea and fungi (Figure 5b). Major endemic viruses included *Enterobacteria* phages and *Bacillus* phages. Major endemic archaea included *Halobacteriaceae*, *Methanosarcinaceae* and *Sulfolobaceae*, and major endemic fungi included *Aspergillus niger* and *Gibberella zeae*.

The numbers of endemic genes differed substantially among the four elevations, with the highest at the 3400 m site and the lowest at the 3200 m site (Supplementary Table S5). At higher elevations, stress, N cycling and S cycling genes occupied larger portions of endemic genes as compared with the 3200 m site, suggesting that they were needed to

cope with the environments at higher elevations (Supplementary Figure S3). In contrast, C cycling, metal resistance and bacterial phage occupied smaller portions. Endemic stress genes included chaperone *dnaK*, *groEL*, *hsp10/groES* and *grpE*, response regulators *desR* and *hrcA* of temperature stress, heme catalase *katE*, response regulator *oxyR* of oxygen stress, proline transporter *proU* of osmotic stress and GTPase *obgE* of radiation stress involved in final steps of the 50S ribosomal subunit assembly and *glnA* and *glnR* genes responsive to N stress.

#### The tight linkage between the microbial gene composition and environmental attributes

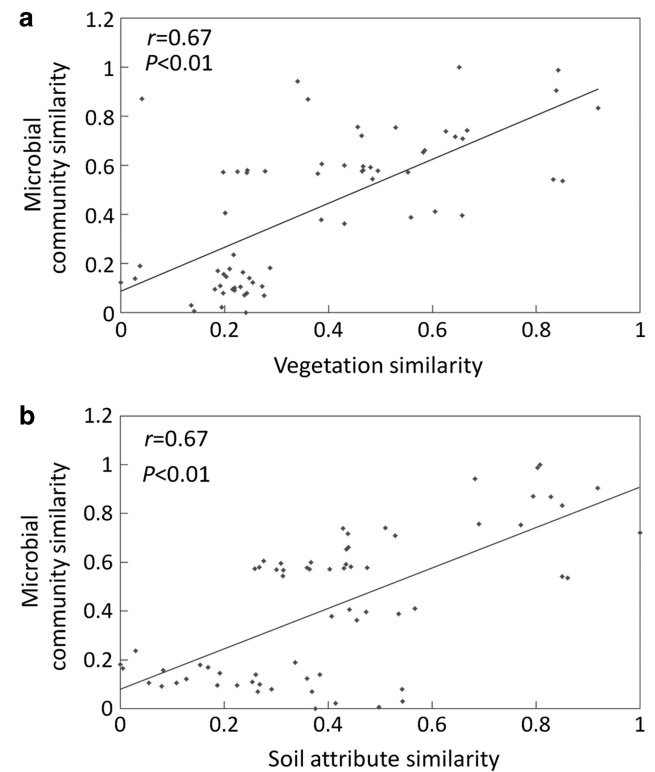
All the environmental attributes were generated by soil geochemical measurements, and four vegetation attributes including total coverage of vegetation, vegetation species number, vegetation diversity were calculated with the Shannon index and vegetation biomass (Supplementary Table S4). To explore the linkages between the microbial gene composition and environmental attributes, the similarities between microbial gene composition and aboveground vegetation and soil attributes were examined, which in turn showed very strong and significant correlations ( $P < 0.01$ ) (Figure 6). To verify it, Mantel tests (Table 1) were conducted with the whole data set of vegetation and soil attributes, showing that the microbial community was significantly ( $P < 0.01$ ) correlated with vegetation and soil attributes with  $r$  values of 0.75 and 0.67, respectively.

CCA was used to identify the major environmental attributes controlling soil microbial gene composition. The results of CCA, a significant model at the confidence level of  $P < 0.01$ , indicated that soil pH, temperature,  $\text{NH}_4^+ - \text{N}$  and vegetation diversity were important environmental attributes controlling the microbial community structure as they were significantly correlated with Axis 1 ( $P < 0.01$ ), which represented the major variations among microbial communities (Figure 7a). The importance of these environmental attributes was verified by Mantel tests on the basis of their significant correlation ( $P < 0.01$ ) with GeoChip data (Table 1).

To partition the contributions of environmental attributes, Variation partitioning analysis was carried out. The results showed that a small set of seven environmental attributes accounted for 81.4% of microbial community variations of GeoChip data (Figure 7b), indicating that they were major factors shaping microbial gene composition. Soil climatic, geochemical and aboveground vegetation attributes appeared to impose strong effects on microbial community. In addition, interaction between vegetation and soil geochemical attributes was substantial (13.2%).

## Discussion

Our first hypothesis, namely that the functional structure of a microbial community differs across elevations and that soil and vegetation attributes contribute significantly to shaping the functional structure, was verified in part, as significant differences were detected except for 3600 and 3800 m samples. The number of detected microbial genes at



**Figure 6** The correlations between (a) microbial community and vegetation and (b) microbial community and soil attributes. Euclid distances among samples were calculated and plotted to generate Pearson correlation values.

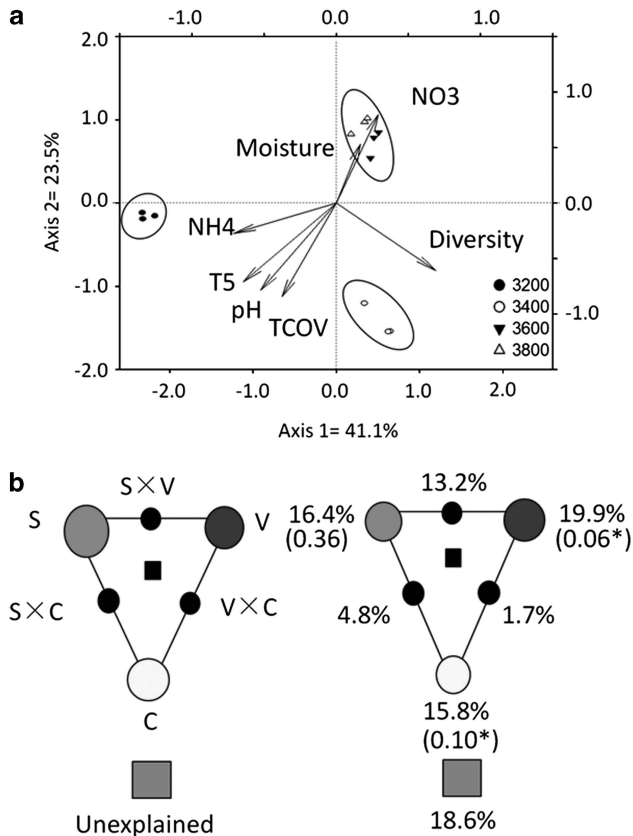
**Table 1** Mantel test of GeoChip data with environmental attributes

Environmental attribute (unit)	$r$ value	Environmental attribute (unit)	$r$ value
Elevation (m)	<b>0.58**<sup>b</sup></b>	TOC20 (%)	<b>0.52**</b>
Soil pH	<b>0.42**</b>	TN20 ( $\text{g kg}^{-1}$ )	<b>0.30*</b>
Moisture (%)	0.10	SIN ( $\text{g kg}^{-1}$ )	0.10
T5* ( $^{\circ}\text{C}$ )	<b>0.69**</b>	Total coverage of vegetation (%)	0.15
T10 ( $^{\circ}\text{C}$ )	<b>0.74**</b>	Vegetation species	0.16
AT ( $^{\circ}\text{C}$ )	<b>0.25*</b>	Vegetation diversity <sup>c</sup>	<b>0.54**</b>
$\text{NO}_3^- - \text{N}10$ ( $\text{mg kg}^{-1}$ )	0.07	Vegetation biomass (g)	<b>0.74**</b>
$\text{NH}_4^+ - \text{N}10$ ( $\text{mg kg}^{-1}$ )	<b>0.47*</b>	Released $\text{CH}_4$ ( $\mu\text{g m}^{-2} \text{h}^{-1}$ )	0.04
TOC10 (%)	<b>0.65**</b>	Released $\text{CO}_2$ ( $\mu\text{g m}^{-2} \text{h}^{-1}$ )	-0.13
TN10 ( $\text{g kg}^{-1}$ )	0.09	Released $\text{N}_2\text{O}$ ( $\mu\text{g m}^{-2} \text{h}^{-1}$ )	-0.12

<sup>a</sup>Abbreviations: T, temperature; AT, air temperature; TOC, total organic C; TN, total N; SIN, soil inorganic N; 5, 10 and 20, soil characteristics measured at depths of 5, 10 or 20 cm.

<sup>b</sup>Significant differences ( $P < 0.05$ ) are indicated in bold. \*\* $P < 0.01$ , \* $P < 0.05$ .

<sup>c</sup>Vegetation diversity was calculated by Shannon index.



**Figure 7** (a) CCA of GeoChip data and geochemistry attributes: soil pH,  $\text{NO}_3^-$ -N,  $\text{NH}_4^+$ -N, the temperature of 5 cm depth in soil (T5), the total coverage of vegetation (TCOV), the alpha diversity of vegetation (diversity), soil moisture (moisture). The percentage of variation explained by each axis is shown and the relationship is significant ( $P < 0.05$ ). (b) Partitioning of microbial diversity variance among important geochemical attributes, soil chemistry including soil pH, moisture, ammonia and nitrate (S), soil climate attributes including soil temperature and soil moisture of 5 cm depth (C), vegetation including total coverage and diversity (V), and their interactions ( $S \times V$ ,  $S \times C$  and  $V \times C$ ). The values in parentheses are  $P$ -values.

the 3200 m site was roughly half of those at the other sites, suggesting that its microbial gene diversity was substantially different. This observation can be explained by soil heterogeneity linking to geographic distance, as the distance between 3600 and 3800 m sites is the shortest (1.3 km), and the 3200 m site is distant from the other sites (6.2–11.5 km). Notably, soil pH might have an adverse effect on microbial gene diversity of the 3200 m site, as it was the highest (pH of 7.9). Alternatively, it might be caused by aboveground vegetation as the vegetation diversity was the lowest at this site. Vegetation affects soil microbial community through a number of mechanisms (Wardle *et al.*, 2004). As producer, it supplies organic C and other resources to soil microbial community, which comprises most of the decomposer system. The quantity and quality of resources depend on vegetation species, thus individual vegetation species can influence the composition and function of soil microbial

communities. However, effects of the vegetation diversity and productivity on soil biodiversity depend on context and vary substantially between locations (Wardle, 2002). In our case, productivity appeared not to be the determinant of microbial gene diversity. Rather, the results of CCA (Figure 7a) and Mantel tests (Table 1) demonstrated that the vegetation diversity had an important role in controlling microbial community structure of the Tibetan grassland. Also, we found that the vegetation diversity appeared to be strongly correlated ( $r = 0.58$ ,  $P < 0.04$ ) with the microbial gene diversity (Supplementary Figure S4), which provided further support.

Soil attributes of pH, temperature and  $\text{NH}_4^+$ -N were crucial for shaping microbial community structure, as shown by CCA (Figure 7a) and Mantel tests (Table 1). Consistently, soil pH and temperature were key drivers in shaping bacterial communities along elevation gradients (Wang *et al.*, 2012; Shen *et al.*, 2013). It has also been observed that the productivity of alpine grassland was mainly controlled by the available N pool (Baumann *et al.*, 2009). Generally, microbial abundance and composition depended on environmental attributes of soil nutrient availability (Lozupone and Knight, 2007; Jesus *et al.*, 2009). These previous studies demonstrated that the soil pH, temperature and N were important for shaping phylogenetic compositions of microbial communities, whereas our results supported it at the level of microbial functional genes.

Notably, the two axes of CCA together (Figure 7a) explained 64.6% of community variance, which was more substantial than most other GeoChip studies (Liang *et al.*, 2011; Lu *et al.*, 2012). It has recently been documented that harsh environments could reduce microbial community variation and consequently correlate microbial community with environmental attributes well (Pointing *et al.*, 2009; Yergeau *et al.*, 2012). Consistently, our recent experiments to examine effects of anthropogenic perturbation on this site indicated that microbial community structure was fragile (Yang *et al.*, 2013). However, it should be noted that a limitation in our study exists in the use of a small area as a proxy to assess the Tibetan alpine grassland. Although, we take comfort in the fact that the study site, located at Haibei Alpine Meadow Ecosystem Research Station, has a representative landscape of a large portion of Tibetan grasslands (Wang *et al.*, 2009b, 2010b), there will be exceptions in other alpine grasslands, given the typically high soil heterogeneity.

Our second hypothesis, namely that variations of microbial stress genes are consistent with the increasing alpine stress at higher elevations, was verified by our observation that microbial stress genes responsible for coping with a cold environment were more abundant at higher elevations (Figure 2), which was apparently a response to low temperature. However, our third hypothesis, namely



that variations of C- or N-cycling genes are significantly linked to soil C and N dynamics in the alpine grassland, was only partially supported by the correlation between several N-cycling genes and N<sub>2</sub>O flux (Figure 4), but no correlation was detected for CO<sub>2</sub> and CH<sub>4</sub> fluxes. An explanation was that the C/N ratio of plants was higher than that of microbes, rendering microbial communities sensitive to N as a limiting factor. For N-cycling genes, *ureC* converted urea into the substrate of N<sub>2</sub>O flux (NH<sub>4</sub><sup>+</sup>-N), whereas *gdh* reversed the reaction. Accordingly, *ureC* and *gdh* were positively and negatively correlated to N<sub>2</sub>O flux, respectively. Interestingly, a positive correlation was observed for *amoA* gene involved in nitrification, whereas negative correlations were observed for *nirS* and *nosZ* involved in denitrification. This finding might implicate that nitrification, but not denitrification, was the major process of N<sub>2</sub>O emission of the alpine grassland. As a consequence, the change of N<sub>2</sub>O emission negatively affected soil NO<sub>3</sub><sup>-</sup>-N content, which in turn affected the abundance of denitrification genes *nirS* and *nosZ*. In support of this view, previous studies indicated that nitrification was the dominant mechanism influencing grassland N<sub>2</sub>O emission (Ryden, 1981; Klemetsson *et al.*, 1988). Meanwhile, it is not surprising to detect no correlation between C-cycling genes and CO<sub>2</sub> flux because the latter includes soil autotrophic respiration, making it unlikely to correlate CO<sub>2</sub> flux with soil microbial genes. However, it remains unclear why no correlation was detected for CH<sub>4</sub> flux.

Our findings supported our fourth hypothesis that the pattern of endemic gene abundances not only differs from that of ubiquitous gene abundances but also differs among elevations. Distribution patterns vary by microbial functional or taxonomic groups (Horner-Devine *et al.*, 2004; Zhou *et al.*, 2008). Thus, we examined endemic genes to identify those with relatively high habitat specificity, which reflected sensitivity to environmental variations. Interestingly, endemic genes were usually low in abundance, raising the possibility that they are part of a 'rare biosphere'. Meanwhile, bearing in mind a caveat that the identification of endemic genes is subjected to the detection level of GeoChip, we noted the enrichment of several microbial groups. Enrichment of stress genes (Figure 5a) in the endemic group implicates their important roles in coping with different alpine environments characterized by low temperature, oxygen and nutrient levels and high UV irradiation. For phylogenetic groups, the enrichment of bacteriophage was not surprising, as the appearance of new bacteria induced new bacteriophage species. For fungi, the narrow dispersal has been reported (Herlemann *et al.*, 2011; Yergeau *et al.*, 2012), which is attributed to its relatively large size compared with bacteria. However, size could not

explain the enrichment of archaea, which are of comparable size to bacteria. Notably, there were reports that archaea were sensitive to certain environments such as N-limited or dry environment (Rothrock and Garcia-Pichel, 2005; Pointing *et al.*, 2009). Our results showed that many endemic genes differed among elevations, which reflected habitat specificity. It was noted that vegetation biomass was the highest for 3200 m (Supplementary Table S4), which might require additional C-cycling genes to utilize C deposited to soil. Meanwhile, stress of temperature, oxygen and nutrient differs by elevations, which induced different stress genes.

Global climate warming effects are most pronounced in the polar and alpine regions, where ecosystems are fragile (Li and Tang, 1988; Hansen, 2000). It is thus necessary to document the ecosystem, particularly the poorly understood microbial communities, because they may encounter catastrophic shifts, should warming continue to occur. Here we report the profiling of microbial distribution at the gene level, which provides valuable insights for potential effects of climate changes on this alpine ecosystem. On the basis of the strategy of space-for-time substitution, we predict that the warming will upshift vegetation and microbial communities along the elevation, thus affecting microbial community structures as well as the particularly sensitive N cycling in the Tibetan grasslands.

## Conflict of Interest

The authors declare no conflict of interest.

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