Soil pH determines the alpha diversity but not beta diversity of soil fungal community along altitude in a typical Tibetan forest ecosystem

Jun-Tao Wang · Yuan-Ming Zheng · Hang-Wei Hu · Li-Mei Zhang · Jing Li · Ji-Zheng He

Received: 18 December 2014 / Accepted: 15 January 2015 / Published online: 5 February 2015
© Springer-Verlag Berlin Heidelberg 2015

Abstract
Purpose Despite their symbiotic relationship with trees and the vital role as decomposer in forest, soil fungi received limited attention regarding their changes with altitude in forest ecosystems. This study aimed to determine the diversity patterns of soil fungi along an altitudinal gradient on Mt. Shegyla, a typical forest ecosystem on the Tibetan Plateau.

Materials and methods High-throughput barcoded pyrosequencing and quantitative PCR approaches were employed to measure the community composition, diversity, and abundance patterns of soil fungal 18S ribosomal RNA (rRNA) gene in 20 samples collected along the altitudinal gradient of Mt. Shegyla.

Results and discussion Abundant taxa in the fungal community were Agaricomycetes and Leotiomyceta on Mt. Shegyla. Fungal abundance decreased significantly with increasing altitude. Beta diversity of the fungal community, as measured using weighted UniFrac distance, was significantly related to altitude. Significant correlation was observed between altitude and alpha diversity including richness and phylodiversity, but not with evenness. Network analysis revealed that Ceramothyrium and Clavulina were two important hubs in the community, and an uncultured fungal taxon that previously detected in glacier forefront dominated this network. Distance-based linear model identified soil pH as the dominant driver which significantly related with fungal alpha diversity including richness, phylodiversity, and evenness. However, fungal abundance and the first component of PCoA on weighted UniFrac matrix (beta diversity) did not change significantly with pH.

Conclusions These results provided strong evidence that soil pH was the dominant driver for structuring altitudinal alpha diversity pattern but not beta diversity pattern or community abundance of soil fungi in this typical forest on the Tibetan Plateau.

Keywords Altitude · Diversity · Forest · Fungi · Soil pH · Tibetan Plateau

1 Introduction
The Tibetan Plateau harbours vast areas of alpine meadow and forest (Luo et al. 2002). The uplift of the Tibetan Plateau generated an altitudinal gradient of almost 5,000 m and thereby inevitably modified the global climate (Spicer et al. 2003). Intensified monsoon from the Indian Ocean brought plentiful precipitation in the southern areas across the latitude, facilitating the formation of typical altitudinal forest belts along the steep environmental gradient (Wang et al. 2014). Forest biomes on the Tibetan Plateau suffered from harsh environmental conditions including strong UV, low temperature and low oxygen content, and the support from the underground communities was especially important in sustaining the biological diversity and ecosystem functions in a scenario of global change (Li et al. 2013; Bardgett and van der Putten 2014;...
Shen et al. 2014b). Fungi are recognized as a vital component of the belowground community intimately related with plant communities in a forest (He et al. 2005; Mueller et al. 2014; Peay et al. 2013). Soil fungi play a key role as decomposer to accelerate degradation of soil organic carbon and nitrogen input (McGuire et al. 2010; Schneider et al. 2012), and they also have a symbiotic relationship with aboveground vegetation, which benefits plant with more resistance against extreme circumstances like oligotrophic or arid habitats (Compan et al. 2010).

Altitudinal distribution patterns of biodiversity could be interpreted into alpha diversity and beta diversity. The former was usually characterized by richness, evenness, and phylodiversity. Richness pattern of fungi evaluated through the operational taxonomic unit (OTU) counts across individual community demonstrated a species-area relationship along the altitudinal gradient of the Alps (Pellissier et al. 2014). Phylodiversity and evenness pattern demonstrated the fungal variance across altitudinal forest types in the Andes (Geml et al. 2014). The latter reflects the shifts of community composition triggered the turnover of fungi (beta diversity) along altitude (Geml et al. 2014). A previous study revealed that variation of prokaryotic community composition (beta diversity) among different altitudinal belts was much larger than in the same belt (Wang et al. 2014). Soils from different forest types with discrete edaphic properties in the Amazon basin harbored distinct fungal communities (Peay et al. 2013). A general knowledge on fungal community, abundance, and diversity patterns along an altitudinal gradient is essential to precisely interpret the fungal functions and responses to environmental factors.

Factors driving the fungal diversity pattern could be quite variable under different soil conditions. A previous experiment demonstrated pH as the most powerful driver in structuring the richness and abundance of fungi in an arable soil (Rousk et al. 2010). The superiority of soil nutrients in structuring fungal community was recognized across land-use types (Lauber et al. 2008). Situation might be more complicated for the interaction of various factors along altitude in spontaneous habitat, since different environmental factors co-vary with altitude (Sundqvist et al. 2013). The Tibetan Plateau was considered as the third pole with glacier and permafrost (Yang et al. 2013). Its harsh environment developed distinct plant/animal taxa in the Tibetan Plateau from the plain areas (Guo et al. 2011). However, we have limited information on the soil fungal community variance and diversity patterns along altitude.

This study aimed (i) to determine the community variance (beta diversity), abundance, and alpha diversity patterns (including richness, evenness, and phylodiversity) of fungi along the altitudinal gradient and (ii) to evaluate fungal effect of different environmental factors in structuring fungal altitudinal distribution patterns in the forest soils on the Tibetan Plateau. To achieve these goals, we collected soil samples along a 1,300-m altitudinal gradient on Mount (Mt.) Shegyla, a typical forest ecosystem on the Tibetan Plateau, and performed quantitative PCR (qPCR) and high-throughput barcoded pyrosequencing analyses on the soil fungal community.

2 Materials and methods

2.1 Soil sample collection, DNA extraction and physicochemical analyses

The Nyingchi District in the southeast Tibet harbors the largest area of forest on the Tibetan Plateau. Different from other regions, Nyingchi has a wetter climate with mean annual precipitation higher than 650 mm, which facilitates the proliferation of forest. Soil samples were collected in July 2011 on Mt. Shegyla (94° 25’–94° 45’ E, 29° 35’–29° 57’ N) characterized by the typical altitudinal vegetation in Nyingchi. The base of Mt. Shegyla is 2,100 m (above sea level) and the peak is 5,300 m. Seven altitudinal sites were identified ranging from 3,351 to 4,477 m where no obvious anthropogenic disturbance was observed. At each site, three surface soil samples (0–10 cm) were collected from individually separated 10 m×10 m plots by pooling eight soil cores randomly taken from each plot. Stones and plant residues were removed. Fresh samples were transported to the laboratory in a freezer and sieved to 2 mm before storage at 4 °C. A small portion of each soil sample was freeze-dried and preserved at −80 °C before DNA extraction. The aboveground vegetation and the altitude of each plot were also recorded.

DNA was extracted from 0.25 g of fresh soil samples using a MOBIO Ultracean Soil DNA Isolation Kit (MOBIO laboratories, Carlsbad, CA, USA) following the manufacturer’s instructions as previously described (Wang et al. 2014). DNA extracts were qualified using a NanoDrop® ND-2000c UV-Vis spectrophotometer (Thermo Fisher Scientific, Wilmington, USA), and tenfold dilutions were used in the downstream molecular analyses. Soil moisture content was determined by oven-drying fresh soil samples at 105 °C for 12 h. Soil pH was measured using a soil to water ratio of 1:2.5. Soil organic carbon (SOC) was determined using the K2Cr2O7 oxidation method, and total nitrogen (TN) was determined using a Vario EL III analyzer (Elementar Analysensysteme GmbH, Hanau, Germany). Soil clay content (<0.002 mm) was determined using a Mastersizer 2000 Laser Diffraction Particle Analyzer (Malvern Instruments Ltd., Malvern, UK). Soil cation exchange capacity (CEC) was measured with the Kjeldahl method (Kitsopoulos 1999). Detailed edaphic properties of the soil samples are listed in Table 1.
2.2 Quantitative PCR analyses and barcoded pyrosequencing of fungal communities

Quantitative PCR assays were performed targeting the 18S ribosomal RNA (rRNA) genes on a BIO-RAD IQ5 thermal cycler (Bio-Rad Laboratories, Inc., CA, USA) to determine the fungal abundance. Primer pair NS1 (GTAGTCATATGC TTGTCTC) and FUN (ATTCCCCGTTACCGTTG) was used in the quantification of 18S rRNA genes (May et al. 2001). The 25-μl PCR reaction mixture contained 12.5 μl Premix Ex Taq™ (Takara Biotechnology, Dalian, China), 1 μl DNA template, 2.5 μl of each primer (10 μM), and 6.5 μl sterilized H2O.

The primer pair EF4 (GGAGGGGRTGTATTATTAG) and FUN5 (GTAAAAGTCCTGTTCCCCTTCCC) targeting 18S rRNA gene was used in the barcoded pyrosequencing (Smit et al. 1999). Primer A-Key (CGTATCCGTCTTACCGCCTCAG) and Primer B-Key (CTATGCGCTTGCCAGCC CGCTCAG) were respectively linked to the 5′ end of the forward and reverse primer as the adaptor. Between the forward primer and the Primer A-Key, a 10-bp MID sequence was added for barcode identification. The 25-μl PCR reaction system contained 12.5 μl Premix Ex Taq™ Version 2.0 (Takara Biotechnology), 2 μl DNA template (1–10 ng), 1 μl of each primer (10 μM), and 8.5 μl sterilized H2O. The resultant PCR amplicons were purified with a Wizard SV Gel and PCR Clean-Up System (Promega Corporation, Madison, USA) following the manufacturer’s instructions. The purified amplicons were mixed equimolarly in a 2-ml tube and sequenced on a 454 GS FLX Titanium pyrosequencer (Roche 454 Life Sciences, Branford, USA).

2.3 Pyrosequencing data processing

Processing of the raw sequences obtained through high-throughput barcoded pyrosequencing was performed using the Quantitative Insights into Microbial Ecology (QIIME) pipeline (Caporaso et al. 2010b). Reads with quality scores lower than 20, ambiguous bases and improper primers were discarded before clustering. Chimeras were eliminated by performing the identify_chimeric.seqs.py and filter_fast.py scripts within QIIME. The resultant high-quality sequences were then clustered into operational taxonomic units (OTUs) at 97% similarity using UPARSE (Edgar 2013). Taxonomic classification of representative sequences from individual OTU was performed by BLASTing against the QIIME-compatible versions of the Silva database (Pruesse et al. 2007). PyNAST was chosen as the alignment algorithm against the Silva database (Caporaso et al. 2010a). Fungal community composition was described by the mean relative abundance of sequences assigned to different taxa.

Before diversity calculation, a resampling procedure was employed at a depth of 285 sequences per sample to eliminate the bias on diversity comparison as caused by unevenly sequencing. Fungal richness was demonstrated using the OTU counts at the 97% similarity, and Faith’s phylodiversity index was calculated to compare the phylodiversity. Evenness of the fungal community was evaluated using Simpson’s evenness index. The weighted UniFrac matrix was calculated as the beta diversity metrics on community variance along altitude (Lozupone et al. 2007).

2.4 Statistical analysis

Spearman’s correlation analysis was performed to examine the relationships among altitude, edaphic factors, and mean annual temperature (MAT) (Hijmans et al. 2005). Nonmetric multidimensional scaling (NMDS) analysis based on the weighted UniFrac distances matrix was performed to uncover the variance of fungal communities along altitude using the nmds.py script in QIIME. Linear and quadratic fittings were performed in SPSS 18.0 to explore the altitudinal patterns of fungal abundance and diversity, and the significance was examined through analysis of variance (ANOVA). A distance-based multivariate linear model (DistLM) was performed to evaluate the effects of different environmental factors on fungal distribution pattern (Anderson 2001) using the Vegan package (Oksanen et al. 2007) in R (R Core Team. 2013).

A network analysis based on the 97% OTU identity was performed in CoNet (Faust et al. 2012) to explore the linkage among different fungal taxa. A filtering procedure was performed in advance to exclude OTUs with less than 10
sequences or three occurrences to avoid spurious correlations. Correlation scores were calculated using Spearman correlation, Pearson correlation, Bray-Curtis dissimilarity, mutual information, and Kullback-Leibler dissimilarity, and only those that reached significant level ($P<0.05$) by at least three methods were considered to be effective. The importance of certain taxon in the network was evaluated using eigenvector centrality (EC) based on the nod’s connections with other nods in Gephi (Bastian et al. 2009). High degree nod within module was considered as a module hub that got higher EC score than other nods in the module, while connector of module hub was considered as the most important networks hub with the highest EC score (Deng et al. 2012). Calculation on topological properties and polish of the network were also performed in the Gephi.

3 Results

3.1 Variations of environmental factors along the altitudinal gradient

The correlations among altitude, MAT, and edaphic properties were examined using Spearman’s correlation matrix. The MAT showed a significant decreasing trend with the increasing altitude ($r^2=0.82$, $P<0.001$). No significant relationship could be observed between altitude and any edaphic factors. It was notable that soil pH, varying from 4.01 to 6.03, exhibited a concave pattern while other factors exhibited unimodal distributions with increasing altitude. Significant correlations were found among nearly all the edaphic factors, especially that soil pH exhibited significantly negative correlations with clay content ($r^2=0.48$, $P<0.05$), TN ($r^2=0.48$, $P<0.05$), SOC ($r^2=0.65$, $P<0.01$), and CEC ($r^2=0.67$, $P<0.01$), while the latter four factors exhibited significantly positive correlations with each other.

3.2 Fungal community composition, variations of abundance, and diversity along the altitudinal gradient

For fungal community analysis along the altitudinal gradient, we obtained 152,991 high-quality 18S rRNA gene sequences through the high-throughput barcoded pyrosequencing to the 20 soil samples and 7,649 sequences on average were assigned to each sample. Community composition was visualized using the mean relative abundance of each taxon in all the samples (Fig. 1). Agaricomycetes (52 %) was the most abundant phylum in the fungal community, followed by Leotiomyceta (29 %), and uncultured soil fungi (9 %). A small proportion of Glomeromycota (2 %) and Chytridiomycota (1 %) were also detected. Quantitative PCR assays revealed that fungal 18S rRNA gene abundance on Mt. Shegyla ranged from $4.0 \times 10^7$ to $2.0 \times 10^9$ copies g$^{-1}$ soil.

Using the weighted UniFrac distance, a nonmetric multidimensional scaling (NMDS) analysis was performed to illustrate the fungal community variance (beta diversity) along the altitude (Fig. 2). Communities from the middle altitudinal sites tended to cluster together, while those from high and low altitudinal sites were more similar to each other. This finding was further corroborated by Mantel test through which a significant relationship was detected between altitude and the weighted UniFrac matrix of the fungal communities (Spearman $r=0.34$, $P<0.01$). Fungal abundance and alpha diversity (including richness, evenness, and phylodiversity) patterns along the altitude were also characterized (Table 2). Abundance ($P<0.05$) and phylodiversity ($P<0.05$) decreased significantly with the increasing altitude, while richness was lowest at middle elevations ($P<0.05$). No significant pattern was observed on fungal evenness.

![Fig. 1](image1.png) Fungal community composition in forest soils on Mt. Shegyla. The relative abundance of each taxon was represented using its mean value in all the samples. Taxa were arranged in the pie chart in a decreasing order of sequence counts.

![Fig. 2](image2.png) Nonmetric multidimensional scaling (NMDS) analysis (Stress=0.05) using the weighted UniFrac distances showing the fungal community variance along the altitudinal gradient on Mt. Shegyla. Colors of the diamonds indicated the altitudinal sites of the samples.
3.3 Network analysis of soil fungal communities

The relationships among different fungal taxa were explored through construction of an OTU network. This network revealed 37 significant correlations (edges) of 28 OTUs (nodes) (Fig. 3). Short average path length (1.83) and network diameter (4) showed that the network of different taxa in the fungal community held a characteristic of small world. Notably, by calculating the eigenvector centrality (EC), an index that measures the importance of nod in a certain network, several important taxa were identified in the soils. Cyanodermella (OTU_51, EC 0.24), Ceramothyrium (OTU_156, EC 0.58), Gibberella (OTU_16, EC 0.45) in Leotiomyceta, and Clavulina (OTU_90, EC 0.48) in Agaricomycetes were important nods closely related to the other OTUs. More importantly, an uncultured soil fungi OTU_160 (EC 1.0) was identified as a vital hub closely related to OTU_51 and OTU_90.

3.4 Effect of environmental factors in structuring fungal abundance, alpha and beta diversity patterns along the altitude

A distance-based multivariate linear model (DistLM) was used to distinguish the effects of edaphic properties (including soil pH, SOC, TN, moisture, CEC, and clay content), mean annual temperature (MAT), and vegetation in shaping the fungal abundance, alpha and beta diversity pattern along the altitudinal gradient (Table 3). Although soil pH (11 %) and moisture (10 %) explained the largest part of the community composition variance as measured by the weighted UniFrac matrix, neither of them had a statistically significant effect on fungal beta diversity along altitude. The abundance pattern was neither significantly influenced by any factors examined. However, alpha diversity pattern was closely related with environmental factors, for example, fungal richness was significantly affected merely by soil pH ($P<0.01$), and evenness and phylodiversity were dominated by both soil pH ($P<0.001$, $P<0.05$) and MAT ($P<0.01$, $P<0.05$), respectively.

3.5 Fungal abundance, alpha and beta diversity patterns along the soil pH gradient

To further investigate the effect of soil pH on fungal distribution, beta diversity, abundance, and alpha diversity were regressed against soil pH by the quadratic function (Fig. 4). The first axis of a principal coordinate analysis (PCoA) using the weighted UniFrac matrix, which explained 44.0 % of the community variance, was not significantly related with soil pH ($r^2=0.09$, $P>0.05$). No significant relationship was detected between fungal abundance and soil pH ($r^2=0.12$, $P>0.05$). Fungal alpha diversity exhibited strong correlation...
with soil pH. A significant increasing trend of richness was observed with increasing pH ($r^2=0.63$, $P<0.001$), while an opposite trend was detected as for the evenness ($r^2=0.65$, $P<0.001$). Fungal phylodiversity showed a significant increasing trend with the increasing soil pH ($r^2=0.22$, $P<0.05$).

### Table 3

Distance-based multivariate linear model (DistLM) revealed the effect of different environment factors on soil fungal community, abundance and diversity on Mt. Shegyla

| Community | Abundance | Diversity |
|-----------|-----------|-----------|
| pH        | 0.11      | 0.25**    | 0.25*** |
| MAT       | 0.06      | 0.11      | 0.15*   |
| Moisture  | 0.10      | 0.04      | 0.04    |
| SOC       | 0.09      | 0.02      | 0.03    |
| TN        | 0.05      | 0.00      | 0.01    |
| Clay      | 0.05      | 0.06      | 0.07    |
| CEC       | 0.02      | 0.03      | 0.05    |
| Vegetation| 0.07      | 0.09      | 0.07    |

MAT mean annual temperature, SOC soil organic carbon, TN total nitrogen, CEC cation exchange capacity

* $P<0.05$; ** $P<0.01$; *** $P<0.001$

4 Discussion

#### 4.1 Characteristics of fungal community in typical forest soils on the Tibetan Plateau

In this study, fungal community composition in the Tibetan forest soils was determined by performing a high-throughput barcoded pyrosequencing on fungal 18S rRNA gene. Our results revealed that the fungal community was mostly dominated by Agaricomycetes (Basidiomycota) and Leotiomyceta (Ascomycota), while the relative abundance of Chytridiomycota was quite low. This finding is in good agreement with a previous investigation on fungal communities in the Tibetan Plateau meadow soils (Zhang et al. 2013) where Basidiomycota and Ascomycota were also found to be abundant. However, another study using the clone library approach reported that the fungal communities in the high-elevation soils at the Rocky, Andes, and Himalayan mountains were dominated by Chytridiomycota (Freeman et al. 2009). It should be noted that the latter investigation was carried out on soils of the unvegetated habitat. Because the fungal community was closely related with vegetation (Mueller et al. 2014), the aboveground plant community could be an important...

![Fig. 4](#)

Fig. 4 Effect of pH in structuring the abundance (a), beta diversity (b), richness (c), evenness (d), and phylodiversity (e) patterns of fungi in the soils on Mt. Shegyla. Beta diversity of fungal community was characterized by the first component of PCoA on weighted UniFrac distance. Quadratic fittings were performed between soil pH and the indices mentioned.
factor that determines fungal community composition. Taxonomic compositions of fungal community in the Tibetan forest soil were also different from that in the Amazon rainforest soils dominated by Sordariomycetes and Saccharomycetes (Peay et al. 2013).

Detailed information on the characteristics of the fungal community in the Tibetan forest soils was obtained through network analysis, a systems biology approach that explored the relationships of different taxa using the high-throughput sequencing data (Raes and Bork 2008). Topologically, the high degree of Cyanodermella (OTU_51) and Clavulina (OTU_90) indicated that they were frequently related to other taxa. A central hub OTU_160 linked Cyanodermella (OTU_51) and Clavulina (OTU_90) was identified through the network analysis. The representative sequence of OTU_160 detected in the Tibetan forest soil was BLASTed to be highly similar to another sequence from a primary successional glacier forefront in the North Cascade Mountains (Jumpponen 2007). This finding suggested that the habitat for fungi in the Tibetan forest soil was, to some extent, similar to that in primary deglaciated environment and highlighted the importance of belowground fungal communities in sustaining aboveground ecosystem under the harsh conditions.

4.2 Drivers of the fungal diversity patterns along the altitudinal gradient

Global patterns of biodiversity were considered to be determined by both stochastic and determinative processes (Martiny et al. 2006). In the case of altitudinal biodiversity patterns, these two processes were further interpreted as a mid-domain effect and an altitudinal environmental gradient, respectively (Colwell and Lees 2000; Sundqvist et al. 2013). Since no unimodal pattern was observed in our research, the altitudinal fungal distribution on Mt. Shegyla could be mostly due to the determinative process (Zhang et al. 2009). Decreased temperature with increasing altitude was the most significant environmental gradient. The temperature gradient largely limited the plant/animal distribution at high elevations through controlling the physiological activity (Hodkinson 2005). Effect of temperature on the altitudinal distribution of microbes was also observed (Bahram et al. 2012; Wang et al. 2012). The MAT did exhibit a significant effect on evenness and phylodiversity patterns of fungi in our research. However, the concave richness pattern emphasized that the spatial heterogeneity of the soils was strongly deterministic in structuring altitudinal distribution patterns of the soil microbes (Lauber et al. 2008).

Among all the edaphic factors examined, soil pH was found to be the most dominant driver that structured fungal richness, phylodiversity, and evenness. The finding that a lower soil pH supported fewer fungal taxa was quite similar to that from a pH controlled experiment (Rousk et al. 2010). A pH-driven diversity pattern was also detected on soil ammonia oxidizers and eukaryotes (Hu et al. 2013; Shen et al. 2014a). Evenness was another important diversity index that microbiologist should take into account when studying altitudinal biodiversity. In our research, we observed a significant effect of soil pH on fungal evenness along the altitudinal gradient on Mt. Shegyla. Higher evenness of communities in lower pH habitat indicated that different taxa were relatively equal in quantity. However, lower pH was stressful for most taxa, which supported the maintenance but not the prosperity of the community, for instance, acidic mine drainages accommodated much fewer microbial taxa than normal habitat (Kuang et al. 2013). With the increasing pH, some endurable taxa became abundant and the balance of taxa quantity in previous habitat was broken, resulting in the decrease of the community evenness.

Although soil pH explained more community variance (beta diversity) than other factors like vegetation or nutrients (Table 2), it was still much weaker to identify pH as a determinant on fungal beta diversity. Contrastingly, 40-50 % of prokaryotic community variance could be explained by soil pH in our previous study (Wang et al. 2014), indicating that fungi was less affected by soil pH compared with prokaryotes. This finding could be supported by previous pH-manipulated field experiment, in which soil pH was found to exert much weaker influence on fungal community than on bacterial community (Rousk et al. 2010).

5 Conclusions

In conclusion, by performing qPCR and barcode pyrosequencing analyses on the fungal communities of soils from Mt. Shegyla, soil pH was found to be a more important driver than other edaphic factors that determined the alpha diversity (including richness, evenness, and phylodiversity) pattern, rather than the beta diversity or abundance pattern of soil fungi along the altitudinal gradient in this typical Tibetan forest ecosystem. This finding strengthened our knowledge on belowground biodiversity in sustaining aboveground community and would better help interpret the biodiversity change of the Tibetan Plateau under future environmental scenarios.

Acknowledgments This work was financially supported by grants from National Science Foundation of China (41230857, 51221892), MOST (2013CB956300) and STSN-21-02. We appreciated Drs. Mu Wang and Xi Zha from Agricultural and Animal Husbandry College of Tibet for their assistance in soil sampling.

References

Anderson MJ (2001) A new method for non-parametric multivariate analysis of variance. Austral Ecol 26:32-46
Wang J-T, Cao P, Hu H-W, Li J, Han L-L et al (2014) Altitudinal Distribution patterns of soil bacterial and archaeal communities along Mt. Shegyla on the Tibetan Plateau. Microb Ecol 69:135–145
Yang Y, Gao Y, Wang S, Xu D, Yu H et al (2013) The microbial gene diversity along an elevation gradient of the Tibetan grassland. ISME J 8:430–440
Zhang LM, Wang M, Prosser JI, Zheng YM, He JZ (2009) Altitude ammonia-oxidizing bacteria and archaea in soils of Mount Everest. FEMS Microbiol Ecol 70:208–217
Zhang XF, Zhao L, Xu SJ, Liu YZ, Liu HY, Cheng GD (2013) Soil moisture effect on bacterial and fungal community in Beilu River (Tibetan Plateau) permafrost soils with different vegetation types. J Appl Microbiol 114:1054–1065