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The microbially mediated soil organic carbon loss under degenerative succession in an alpine meadow

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Abstract

Land-cover change has long been recognized as having marked effect on the amount of soil organic carbon (SOC). However, the microbially mediated processes and mechanisms on SOC are still unclear. In this study, the soil samples in a degenerative succession from alpine meadow to alpine steppe meadow in the Qinghai–Tibetan Plateau were analysed using high-throughput technologies, including Illumina sequencing and GEOCHIP functional gene arrays. The soil microbial community structure and diversity were significantly ($p < .05$) different between alpine meadow and alpine steppe meadow; the microbial α -diversity in alpine steppe meadow was

significantly ($p < .01$) higher than in alpine meadow. Molecular ecological network analysis indicated that the microbial community structure in alpine steppe meadow was more complex and tighter than in the alpine meadow. The relative abundance of soil microbial labile carbon degradation genes (e.g., pectin and hemicellulose) was significantly higher in alpine steppe meadow than in alpine meadow, but the relative abundance of soil recalcitrant carbon degradation genes (e.g., chitin and lignin) showed the opposite tendency. The Biolog Ecoplate experiment showed that microbially mediated soil carbon utilization was more active in alpine steppe meadow than in alpine meadow. Consequently, more soil labile carbon might be decomposed in alpine steppe meadow than in alpine meadow. Therefore, the degenerative succession of alpine meadow because of climate change or anthropogenic activities would most likely decrease SOC and nutrients mediated by changing soil microbial community structure and their functional potentials for carbon decomposition.

1 INTRODUCTION

Land-cover change is a common phenomenon in land ecosystems, and it has been recognized that this aboveground change would markedly affect the belowground soil organic carbon (SOC) pool. The aboveground change includes processes such as the plant succession and degradation, reflecting the ecological processes caused by the combined effects of natural climate change, overgrazing, deforestation and other human activities (Yan et al., [2005](#); Yu & Xie, [2013](#)). About 1.2 Pg carbon (C) in every year, or about 12%–15% of total anthropogenic fluxes, was released as CO₂ to the atmosphere by land-cover change (Powers, Corre, Twine, & Veldkamp, [2011](#)). Both soil fertility loss and CO₂ release (Powers et al., [2011](#)) were the consequences of the changes in plant residues and the immobilization of organic C mediated by microorganisms (Tate, [1987](#); Van der Werf et al., [2009](#)). Therefore, the study of the effect of land-cover change on SOC and its effect on processes and mechanisms is critically important to understand the global C balance and contributes to sustainable land-cover management.

Microorganisms are one of the most abundant and diverse organisms and are essential to soil ecological function, particularly in SOC and nutrient cycling (Feeney et al., [2006](#); Van der Heijden, Bardgett, & van Straalen, [2008](#)). Many studies have revealed the changes in SOC, CO₂ release, microbial biomass and microbial species diversity by land-cover changes (Fierer, Nemergut, Knight, & Craine, [2010](#); López-Lozano et al., [2013](#); Lundquist, Jackson, Scow, & Hsu, [1999](#); Michelsen, Andersson, Jensen, Kjoller, & Gashew, [2004](#); Wang, Dalal, Moody, & Smith, [2003](#)). Because of the high microbial diversity in soil ecosystem and technical limitations, soil microbial activities and processes involved in soil C cycling have been assessed by indirect indicators in most previous studies, such as soil respiration (Bastida, Moreno, Hernandez, & Garcia, [2006](#); Li, Xu, Zou, & Xia, [2010](#)), metabolic quotient (Bini et al., [2013](#)), exoenzyme activities (Nayak, Babu, & Adhya, [2007](#)) and microbial phospholipid fatty acid (Smith, Marin-

Spiotta, de Graaff, & Balser, [2014](#)). In recent years, some researchers have focused on the changes in soil microbial community based on 16S rDNA, ITS or functional gene sequencing (Xue et al., [2016](#); Zifcakova, Vetrovsky, Howe, & Baldrian, [2015](#)). However, the soil microbial activities and processes mediating the conversion of SOC to CO₂ and biomass are still a “black box” (Ding et al., [2013](#); Lange et al., [2015](#); Sulman et al., [2014](#); Wardle et al., [2004](#)). Therefore, although land-cover changes significantly affect SOC, little is known about the influence of land-cover changes on the metabolic activities and processes of the belowground microbial community (Reeve et al., [2010](#)). Further study on the change in soil microbial potential and metabolic traits following land-cover change is needed (López-Lozano et al., [2013](#)).

The Qinghai–Tibetan Plateau is the highest and the largest low-latitude plateau in the world (Wang et al., [2012a](#); Wang, Wu, Tian, Niu, & Tan, [2012b](#)), and it is an extremely sensitive region to the impact of global warming and environmental changes (Zhang et al., [2013](#)). The alpine meadow (AM), widely distributed on the Tibetan Plateau, occupies over 40% of the Qinghai–Tibetan Plateau area and plays a critical role in regional sustainable development, biodiversity and water resource conservation (Kang, Han, Zhang, & Sun, [2007](#); Zhou, Zhao, Tang, Gu, & Zhou, [2005](#)). The AM is also a large SOC pool. Wang, Cheng, and Shen ([2002](#)) found that soil (0–75 cm) organic C content reached 23.2 Pg in the meadow and steppe grasslands in the Tibetan Plateau, accounting for 23.44% of China's total organic soil-stored C or 2.5% of the global soil C pool. As one of the most important and vulnerable soil C pools, about 3.02 Pg of C has been emitted from the grasslands of the Qinghai–Tibetan plateau because of the changes in land-cover and grassland degeneration in the last 30 years (Wang et al., [2002](#)). In recent decades, succession and degradation have been gradually occurring between different AM types, as AM has appeared in the alpine steppe meadow (ASM) region. This might be the consequences of the climate warming and anthropogenic activities (Guo et al., [2011](#); Wang et al., [2012a](#), [2012b](#); Zhou et al., [2005](#)).

In this study, we adopted Illumina sequencing and functional gene microarray (GEOCHIP) to analyse the processes and mechanisms of changes in microbially mediated SOC in the degenerative succession from AM to ASM in Qinghai–Tibetan Plateau. The aims of this study were to determine (i) the effect of degenerative succession from AM to ASM on SOC and soil microbial community structure; (ii) the divergence of soil C utilization by microbes and microbial functional gene diversity related to C cycling; and (iii) the major environmental factors affecting soil microbial community structure and microbially mediated SOC loss.

2 MATERIALS AND METHODS

2.1 Site and sampling

The study sites were situated in Sanjiangyuan Natural Reserve (97°40'22"–100°05'27"E, 34°08'16"–35°56'06"N), Qinghai Province, China, which was located in the centre of Qinghai–Tibetan Plateau (Zhang et al., [2013](#)). The annual mean air temperature is –5.6~3.8, and the average precipitation is 262.2~772.8 mm (Lu et al., [2015](#)).

Soil sampling sites were set up in AM (35°41'26"N, 99°33'01"E, elevation: 3,880 m) and ASM (35°40'10"N, 99°55'13"E, elevation: 3,490 m). At each site, 10 plots (1 m × 1 m) were established and the diagonal method was used to collect soil samples at the depth of 0–10 cm in each plot. Ten to 15 soil cores were taken from each plot and combined to obtain about 400 g of soil. Roots and stones were removed from samples, and then, the samples from each plot were thoroughly mixed. Ten replicate soil samples were collected from the same site. To avoid contamination during sampling, the sterile gloves, sterilized paper and water were used for sampling from each plot. At the same time, plant properties were investigated and recorded in each plot, including the plant species, plant number, canopy of each grass and plant height (Fang, Shen, Tang, & Wang, [2004](#)). To survey the plant biomass, all the grass was harvested in each plot, dried in the oven at 65°C for about 24 hr and weighed.

2.2 Soil property measurements

All soil samples were air-dried and then sieved to 2 mm. Soil moisture was measured by the drying method (Bao, [1999](#)). Soil pH was measured by pH meter according to the ratio of 1:2.5 soil:H₂O. Total organic C, total nitrogen (TN), total phosphorus (TP), total sulphur (TS), rapidly available phosphorus (RAP), available N (AN), nitrate N (NO₃⁻-N) and ammonium N (NH₄⁺-N) were measured (Bao, [1999](#)). The vegetation properties and soil physicochemical properties are presented in Table [S1](#).

2.3 Soil microbial carbon utilization

A Biolog Ecoplate experiment was performed to examine the microbial functional diversity of carbon metabolism (Cookson, Murphy, & Roper, [2008](#)). Each well of the plate was scanned at the wavelength of 595 nm with the Biolog plate reader (Microlog ReL 3.5) at 12-hr intervals through to 168 hr (Liu, Zhang, Hu, & Li, [2013](#)). C utilization was monitored by average well colour development (AWCD) = $\sum(C_i - R)/n$, where C_i denotes the absorption value of the i th well, R denotes the control absorption well and n denotes the number of plates ($n = 31$). AWCD values of 168 hr were used to calculate the microbial functional diversity of C metabolism (Garland & Mills, [1991](#)). Several indexes were used to analyse diversity and richness of the

communities: the Shannon–Wiener diversity (H): $H = -\sum P_i \times (\ln P_i)$, where P_i was the ratio of the relative absorption of the i th divided by the sum of all relative colour development of the plate at 168 hr; the McIntosh index (U): $U = \sqrt{\sum ni^2}$, where n_i denotes the relative colour development of the i th and the richness index (S) denotes the number of wells with $C_i - R > 0.25$.

2.4 Soil microbial DNA extraction, purification and quantification

Soil microbial DNA extraction was conducted by the Fast DNA Spin kit for soil following the manufacturer's instructions (MP Biomedical, Carlsbad, CA, USA). Soil microbial DNA was further purified twice by using 0.5% low melting point agarose gels and was determined by analysing the ratios of absorbance at 260/280 nm and 260/230 nm. Finally, microbial DNA was quantified using a FIUOstar Optima (BMG Labtech, Jena, Germany).

2.5 Illumina sequencing and data processing

Purified DNA extracts from soil samples were used as a template, and the primers were designed for amplification according to the V4 hypervariable region of the bacterial 16S rDNA gene. The sequence of forward primer was 5'-GTGCCAGCMGCCGCGGTAA-3' (515F), and that of the reverse primer was 5'-GGACTACHVGGGTWTCTAAT-3' (806R) (Caporaso et al., [2011](#), [2012](#)). The reverse primer was combined with a barcode sequence. PCR amplification was used in a 25- μ l reaction mixture, containing 1 μ l of each primer, 2.5 μ l of AccuPrime PCR buffer II (Invitrogen, Grand Island, NY, USA), 5 μ l of DNA and 0.1 μ l of AccuPrime Taq Polymerase. PCR cycle consisted of denaturation at 94°C for 1 min, followed by 30 cycles of 94°C for 20 s, 53 °C for 25 s and 68°C for 45 s and extension at 68°C for 10 min (Ding et al., [2015](#)). The PCR products were purified and run using a MISEQ (Illumina, San Diego, CA, USA; Cong et al., [2015](#); Ding et al., [2015](#)).

Raw data were separated into samples according to the barcode sequence. Adapters, low-quality and ambiguous reads (“N”) were trimmed; for example, reads that did not perfectly match the PCR primer, had nonassigned tags or had reads <250 bp were removed (Kong, [2011](#)). The forward and reverse reads were integrated into a whole sequence by FLASH (Magoč & Salzberg, [2011](#)). Operational taxonomic units (OTUs) were defined at 97% similarity level by using UCLUST (Edgar, [2010](#)). The singletons were removed. The ribosomal database project (RDP) classifier was used to determine the taxonomic identity of each phylotype (Wang, Garrity, Tiedje, & Cole, [2007](#)). The number of detected OTUs and sequences at different levels of classification was counted. Random resampling was processed with 15,000 sequences per soil sample. All these data were tested with the Galaxy Illumina sequencing pipeline.

2.6 GEOCHIP hybridization and data processing

GEOCHIP 4.0 was used for detecting soil microbial DNA functional gene diversity. GEOCHIP 4.0 contained 82,000 oligonucleotide probes covering 141,995 functional genes involved in 410 gene categories related to C, N cycling and other biogeochemical processes. The detailed GEOCHIP information is presented on the website (<http://ieg.ou.edu>). Purified DNA was labelled with Cy3 fluorescent dye using a random priming method (Tu et al., 2014). All hybridizations were carried out at 42°C for 16 hr using a hybridization station (MAUI; BioMicro Systems, Salt Lake City, UT, USA), and arrays were scanned at full laser power and 100% photomultiplier tubes with a NimbleGen MS200 Microarray scan (Roche, Madison, WI, USA). Scanned images were gridded by NIMBLESCAN software (Tu et al., 2014).

Raw GEOCHIP data were uploaded to the GEOCHIP data analysis manager (<http://ieg.ou.edu/microarray/>). Data were preprocessed data using the following steps: (i) the poor-quality spots with a signal-to-noise ratio of less than 2.0 or the signal intensity value less than 1,000 were discarded; (ii) genes that were detected in no more than six of 10 replicate samples from the same sampling site were removed; (iii) normalizing the signal intensity of each spot by dividing the mean value of each sample of total signal intensity; and (iv) transformation of the data to the natural logarithmic form (Cong et al., 2015; Ding et al., 2015; He et al., 2010).

2.7 Statistical analysis

Plant diversity was calculated by Simpson index, and the number of plant species was calculated in all samples based on the survey data in the fields. Shannon index, Simpson's index, Pielou's evenness, Simpson evenness and OTUs richness index were used to test soil microbial diversity based on Illumina sequencing data of 16S rDNA and *gyrB* genes in GEOCHIP4.0. Data analysis was performed by *t* test analysis, and *p* values of *t* tests were adjusted by a false discovery rate (FDR) of <5% (Kong et al., 2013). Principal coordinate analysis (PCoA) was used to assess the distribution of microbial communities based on the Bray–Curtis dissimilarity matrix. The Mantel test was used to analyse the correlation between microbial community structure and environmental factors; variance partitioning analysis (VPA) was performed to analyse the contributions of environmental variables to the microbial community structure. Canonical correspondence analysis (CCA) was used to determine the major environmental attributes contributing to the microbial community structure. Before performing CCA, the environmental variables were first filtrated according to the variance inflation factors (VIF) (Yang et al., 2014). All data were tested in R version 3.1.2 using the VEGAN package (version 3.1.2).

2.8 Soil microbial network construction

Based on random matrix theory (Deng et al., [2012](#)), ecological networks were constructed using sequencing data of 16S rDNA. In the network construction, only three of 10 replicates of OTUs data were used. Various network properties, such as average clustering coefficient, average degree, modularity index and average path distance, were determined. Among the topological properties in the ecological network, modularity could be used to measure the extent of species interactions and it could characterize the ecosystem quality and stability (Alon, [2003](#); Olesen, Bascompte, Dupont, & Jordano, [2007](#)). Average degree was used to describe the properties of nodes (Guimera, Sales-Pardo, & Amaral, [2007](#)), and average clustering coefficient was used to measure the extent of module structure present in a network (Deng et al., [2012](#)), while harmonic geodesic distance (HD) could represent the path length of different nodes in disjointed graph (Deng et al., [2012](#)).

The network modules were generated using rapid greedy modularity optimization. Hub and connector genes were determined by among-module connectivity (P_i) and within-module connectivity (Z_i) (Olesen et al., [2007](#)). The Z_i described the degree of connectivity between a node and other nodes in its own module, and P_i reflected the extent that a node was connected to the other modules. The network parameters and properties were obtained from the website (<http://ieg2.ou.edu/mena/>). According to the parameters and properties, the visualized network graphs were constructed by CYTOSCAPE 2.8.0 software (Cline et al., [2007](#)).

3 RESULTS

3.1 SOC and soil geochemical properties

The soil and plant characteristics were remarkably different between AM and ASM (Table [S1](#)). *Kobresia pygmaea*, *Potentilla bifurca* and *Leontopodium pusillum* were the dominant species in AM, while *Poa annua*, *Oxytropis deflexa* and *Carex tristachya* were the dominant in ASM. The plant biomass and plant α -diversity were significantly lower ($p < .01$) in AM sites than in ASM sites.

Among the measured soil parameters (Table [S1](#)), SOC content was significantly ($p < .01$) higher in AM samples than in ASM samples, and most of the other soil nutrient contents followed the same trend, such as soil TP, TN and RAP (Table [S1](#)). All these results indicated that the soil degenerative process from AM to ASM resulted in decrease in SOC and nutrient contents, even though the aboveground grass biomass might be temporarily increased.

3.2 Soil microbial community composition and structure between AM and ASM

To compare soil microbial community composition and structure in the two meadow sites, 16S rDNA high-throughput sequencing was performed. A total of 13,307 and 15,754 operational taxonomic units (OTUs) were separately obtained at 97% similarity level, ranging from 2,663 to 4,407 OTUs per sample in AM and from 3,615 to 4,759 OTUs per sample in ASM. For taxonomic identification, all detected OTUs could be classified into 34 bacterial phyla and two archaeal phyla. The dominant phyla were *Acidobacteria*, *Proteobacteria*, *Actinobacteria* and *Planctomycetes* in both AM and ASM; the soil microbial richness (the number of OTU) was significantly ($p < .05$) higher in ASM than in AM (Table S2). Total 21 subgroups of phylum *Acidobacteria* were detected and seven of them were dominant (Table S3). At the family classification level, a total of 176 families (average number of OTU over one in 10 replicate samples) were detected in the two sites, with 154 families in AM and 171 families in ASM. The most dominant families (average number of OTU over 100 in 10 replicate samples) in these sites were *Planctomycetaceae*, *Actinomycetales*, *Solirubrobacterales*, *Chitinophagaceae*, *Sphingomonadaceae* and *Acidimicrobiales* (Data S1). At the genus classification level, a total of 369 genera (average number of OTU over 1 in 10 replicate samples) were found in the two sites, and 311 genera in AM and 342 genera in ASM. The most dominant genera (average number of OTU over 80 in 10 replicate samples) in these sites were *Acidimicrobineae*, *Conexibacteraceae*, *Zavarzinella* and *Gemmatimonas* (Data S2). However, the microbial diversity based on *gyrB* gene was lower than on the 16S rDNA and was not significantly different between the two alpine meadow soils (Table 1).

Table 1. Overall microbial community diversity detected by Illumina sequencing and Biolog Ecoplate data in the two meadow sites

	Indices	AM	ASM	FDR
Taxonomic diversity (16S rDNA)	Shannon index (H)	7.33 ± 0.06	7.59 ± 0.03	0.003
	Simpson index (D)	636.15 ± 36.47	844.31 ± 22.45	0.000
	Pielou's evenness (J)	0.89 ± 0.003	0.91 ± 0.001	0.003

	Indices	AM	ASM	FDR
	Simpson evenness (Si)	0.17 ± 0.01	0.19 ± 0.01	0.012
	Richness index	3680.80 ± 559.69	4374.40 ± 301.34	0.006
Phylogenetic diversity (gyrB)	Shannon index (H)	6.25 ± 0.16	6.05 ± 0.16	0.333
	Simpson index (D)	585.32 ± 78.45	463.02 ± 61.50	0.283
Biolog data	Shannon index	3.22 ± 0.01	3.29 ± 0.01	0.006
	McIntosh index	6.85 ± 0.28	8.69 ± 0.31	0.000
	Richness index	25.13 ± 0.44	27.40 ± 0.40	0.003

- Data present the mean value and standard error.
- AM, alpine meadow; ASM, alpine steppe meadow.

The relative abundance of *δ-Proteobacteria*, *Planctomycetes Chloroflexi* and *Firmicutes* was significantly ($p < .05$) higher in ASM than in AM. The relative abundance of α -*Proteobacteria* and β -*Proteobacteria* was significantly ($p < .05$) higher in AM than in ASM, while the others had no significant difference between the two meadow sites (Table [S2](#)).

Therefore, the composition and relative abundance of soil microbial community were significantly different between AM and ASM.

The α -diversity indexes of microbial community structures were calculated (Table [1](#)). The Shannon index and Simpson index were significantly ($p < .01$) higher in ASM (7.59 and 844.31, respectively) than in AM (7.33 and 636.15, respectively). PCoA of the overall microbial

community structure showed that the microbial communities of the two meadow sites were well separated (Figure 1). Furthermore, three nonparametric multivariate statistical tests (MRPP, ANOSIM and Adonis) indicated that there were significant ($p < .01$) differences between these two sites (Table S4). Therefore, the diversity and structure of the soil microbial communities were significantly different between AM and ASM soil.

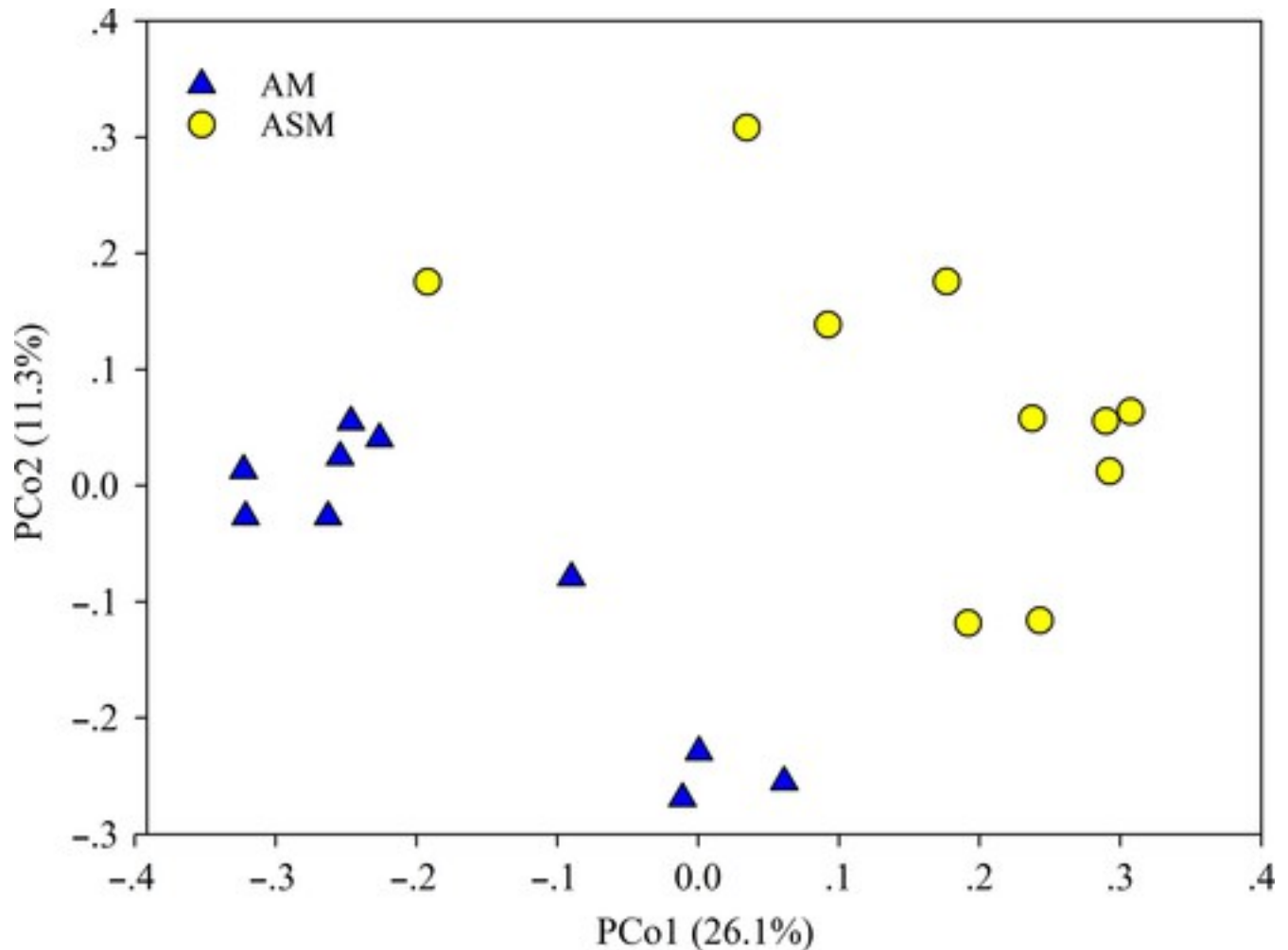


Figure 1

[Open in figure viewerPowerPoint](#)

Principal coordinate analysis (PCoA) based on Bray–Curtis dissimilarity matrix of Illumina sequencing between the two meadow sample sites. AM, alpine meadow; ASM, alpine steppe meadow [Colour figure can be viewed at wileyonlinelibrary.com]

3.3 Ecological network analysis of soil microbial communities between AM and ASM

The ecological networks that were constructed had 613 and 828 nodes for AM and ASM, respectively, under the identical thresholds (0.89; Table S5). In this study, modularity, average

degree and average clustering coefficient were higher in the network of ASM than in AM (Table S5), which indicated that soil microbial community structure in ASM site might be more complex and tighter than in AM site.

In the $Z P$ -plot, peripherals representing a node in this category have lower connectivity and lower value of P_i and Z_i . According to the network topological structure graph, the majority of nodes belonged to the peripherals and did not contact with the external module ($P_i = 0$). The connector category describes the nodes with lower Z_i , but higher P_i . In AM sites, no nodes were detected that belonged to connector category, while seven connectors were observed in ASM site (Figure 2). Among these connectors, four of seven connectors were derived from *Proteobacteria*, and the other three connectors were derived from *Acidobacteria*, *Gemmatimonadetes* and *Actinobacteria*, respectively. Module hubs represented the nodes with higher Z_i , but lower P_i . For AM, five nodes were detected that belonged to the module hub category, which were composed of three *Proteobacteria*, one *Acidobacteria* and one *Actinobacteria*. In ASM, seven module hubs were observed, which were composed of three *Actinobacteria*, three *Acidobacteria* and one *Chloroflexi*. According to the results, the network interaction of soil microbial taxa had been substantially changed in the process of degenerative succession from AM to ASM.

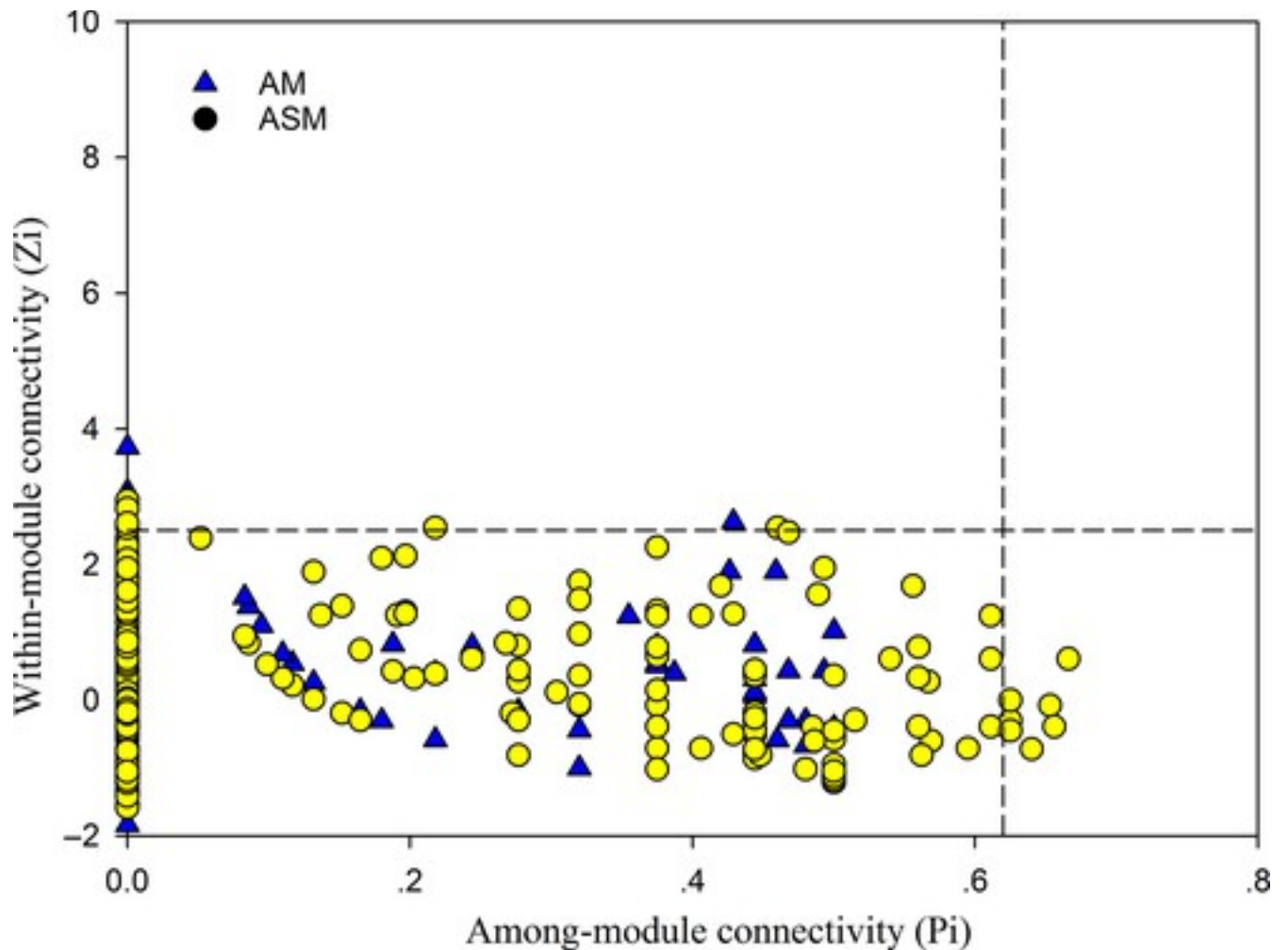


Figure 2

[Open in figure viewer](#)[PowerPoint](#)

Z P-plot showing the distribution of OTUs based on their topological roles. Each colour represents a kind of meadow type, ASM (yellow) and AM (blue). The topological role of each OTU or gene was determined according to the scatter plot of within-module connectivity (Z_i) and among-module connectivity (P_i). AM, alpine meadow; ASM, alpine steppe meadow [Colour figure can be viewed at wileyonlinelibrary.com]

3.4 Carbon utilization of soil microbial communities

The average well colour development (AWCD) showed that the C sources were rapidly used from 24- to 168-hr incubation and reached the maximum values at 168 hr (Figure 3). Compared with AM, samples in ASM had higher AWCD values across all the incubation time points (Figure 3). The Shannon index, McIntosh index and richness index were significantly ($p < .01$) higher in ASM than in AM (Table 1). These results indicated that soil microbial diversity and activity to SOC utilization were higher in ASM than in AM.

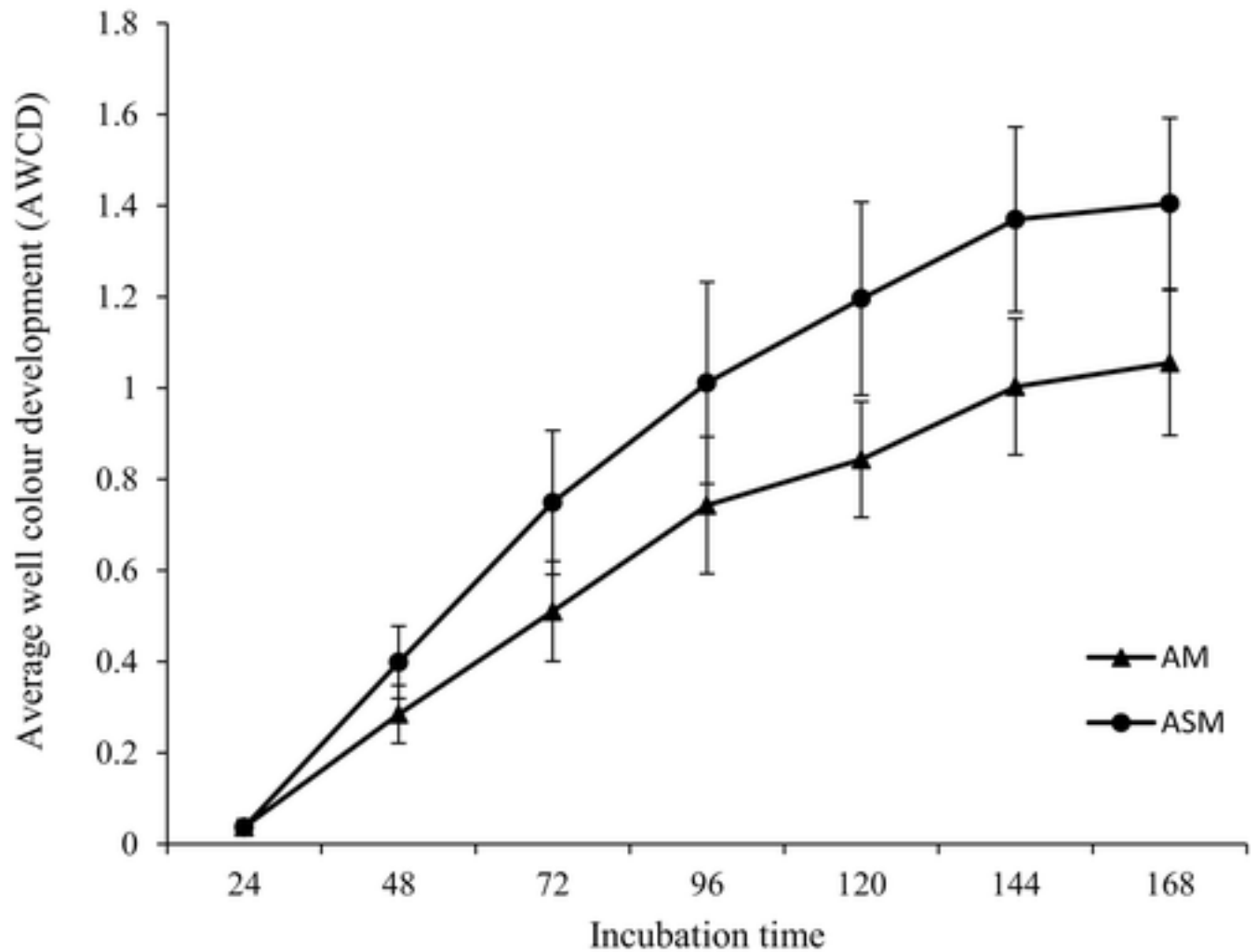


Figure 3

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Average well colour development (AWCD) between alpine meadow and alpine steppe meadow. AM, alpine meadow; ASM, alpine steppe meadow

Carbon sources analysis showed that the use of most C sources was significantly ($p < .01$) higher in ASM than in AM, such as polymers, carbohydrates, phenolic acids, carboxylic acids and amino acids (Fig. [S1](#)). These results implied that the soil microbial communities in ASM might consume a broader range of C substrates to satisfy their ecological function. Furthermore, the top three C sources utilized by microbes in ASM were polymers, amino acids and phenolic acids, while the top three C substrates in AM were polymers, amino acids and carbohydrates (Fig. [S1](#)). The range of C sources metabolized indicates different ecological functions of the soil microbial communities in these two meadow sites.

3.5 Differences in soil microbial functional gene related to C and N cycling

A total of 6,425 microbial genes involved in different C degradation pathways, such as starch, pectin, hemicellulose, cellulose, chitin and lignin degradations, were detected by GEOCHIP 4.0 in the meadow samples. The detected relative abundance of many genes involved in labile C degradation was significantly ($p < .05$) higher in ASM than in AM (Figure 4), such as the *pectinase*, *rgl* and *rgl* genes involved in pectin degradation and the *ara* gene involved in hemicellulose degradation. However, the detected relative abundance of genes involved in recalcitrant C degradation was significantly ($p < .05$) higher in AM than in ASM, such as endochitinase gene involved in chitin degradation, *mnp* and phenol oxidase genes involved in lignin degradation (Figure 4). These results apparently indicated that soil microorganisms in AM might have a higher potential ability to use some recalcitrant C (e.g., chitin and lignin), while after conversion of AM to ASM, the microorganisms may tend to decompose more labile C (e.g., pectin and hemicellulose).

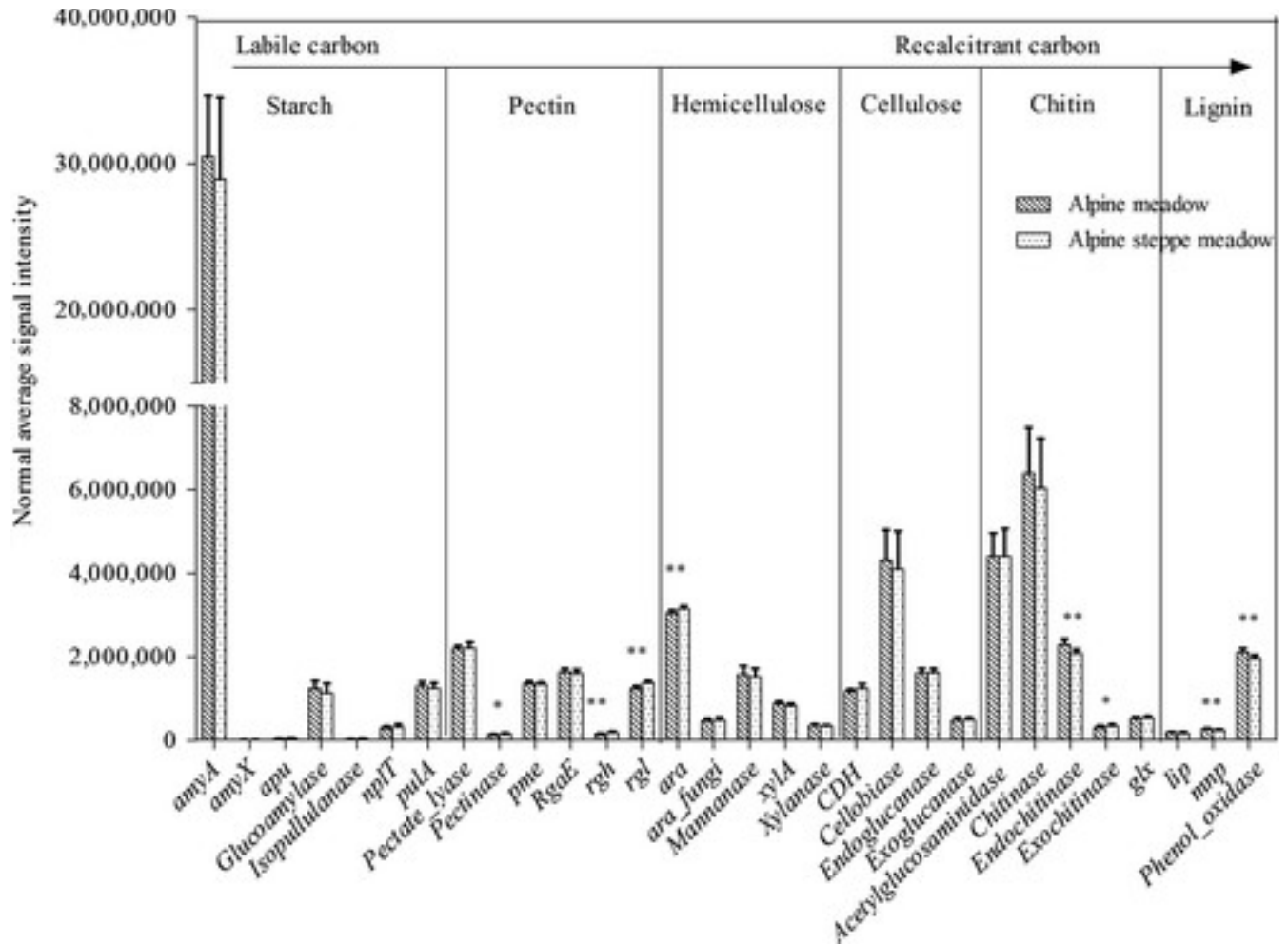


Figure 4
[Open in figure viewer](#) [PowerPoint](#)

The normalized average signal intensity of detected gene probes for carbon degradation in AM and ASM sites based on GEOCHIP 4.0 data. Error bars represent standard error. The difference between AM and ASM was tested by two-tailed unpaired *t* test. **p* < .05; ***p* < .01. AM, alpine meadow; ASM, alpine steppe meadow

Microbial genes related to N cycling were analysed. The detected relative abundance of *amoA* gene related to nitrification was significantly higher (*p* < .05) in ASM than in AM. In contrast, the detected relative abundance of *napA* and *nrfA* genes related to N reduction and *hzo* gene involved in Anammox was significantly (*p* < .05) higher in AM than in ASM (Figure 5). These variations in N cycling genes might lead to the difference in transformation from NO₃⁻ synthesis to NH₄⁺ synthesis. These results indicated that many N cycling genes might be changed and influence the N bioprocess under the degenerative succession from AM to ASM.

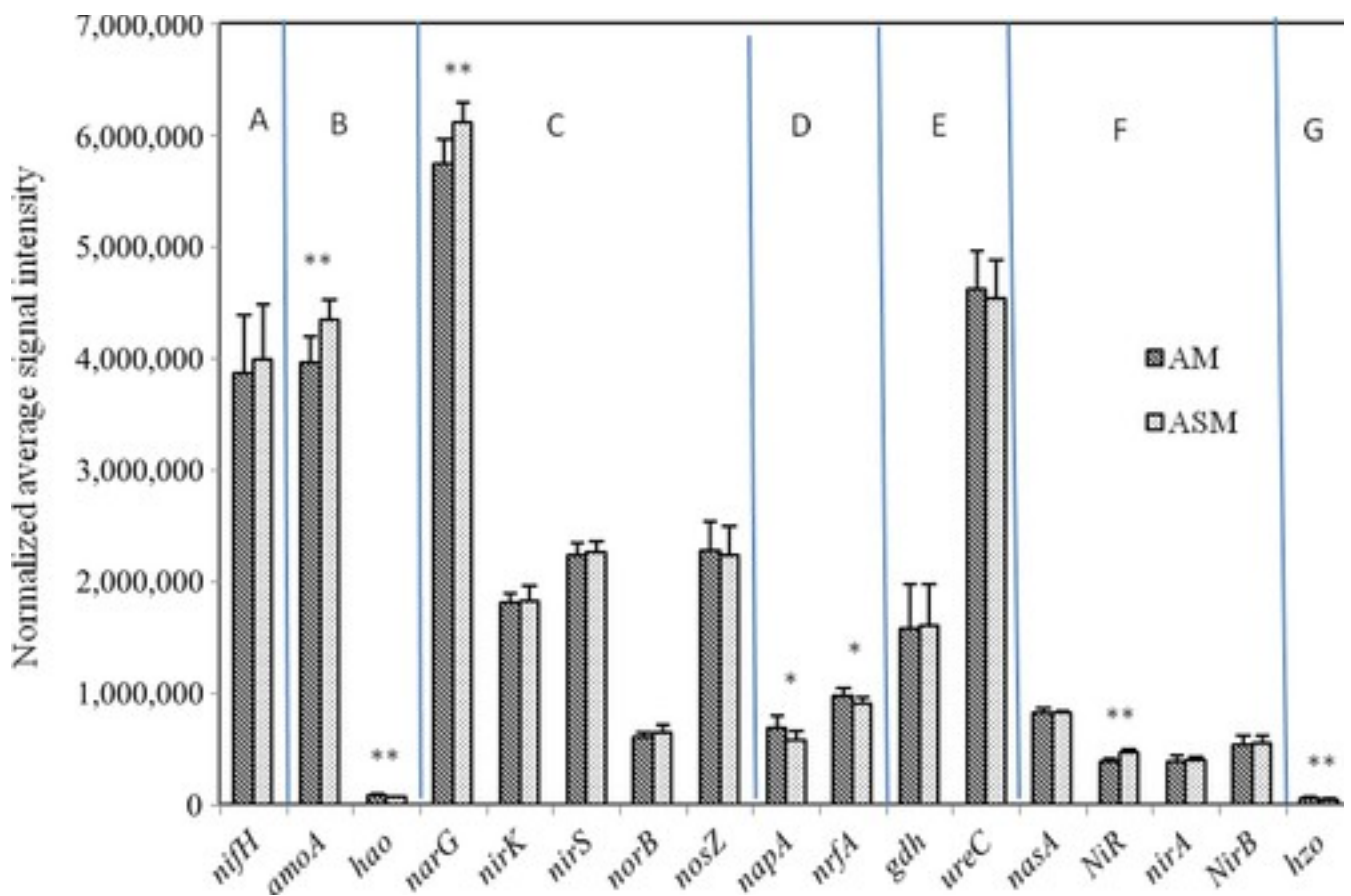


Figure 5

[Open in figure viewerPowerPoint](#)

The normalized average signal intensity of N cycling genes in AM and ASM sites based on GEOCHIP 4.0. Among the detected genes, (a) N₂ fixation, (b) nitrification, (c) denitrification, (d) dissimilatory N reduction, (e) ammonification, (f) assimilatory N reduction and (g) Anammox. The differences in functional genes in the two meadow system were tested by two-tailed paired *t* test. **p* < .05, ***p* < .01. AM, alpine meadow; ASM, alpine steppe meadow [Colour figure can be viewed at wileyonlinelibrary.com]

3.6 Relationship between soil microbial community and environmental factors

To identify the relationship between environmental factors and soil microbial community, Mantel test and CCA were performed (Table 2). The results indicated that soil properties such as pH, TN, TP, TS, SOC and RAP were significantly ($p < .05$) affected by both soil microbial taxonomic and functional gene structures. The CCA results indicated that SOC, plant diversity and TP might be the most important factors in forming microbial taxonomic structure ($p = .05$; Fig. S2A) and microbial functional genes ($p = .01$; Fig. S2B).

Table 2. Mantel test between 16S rDNA OTUs and functional genes of carbon and nitrogen cycling genes with environmental factors

Environmental parameters	16S OTUs		Functional genes	
	<i>r</i>	<i>p</i>	<i>r</i>	<i>p</i>
Vegetation properties				
Plant biomass	.102	.124	.444	.001
Plant diversity	.181	.027	.391	.001
Plant species	-.055	.719	.007	.410
Soil properties				
Moisture	-.046	.654	.001	.428
pH	.213	.015	.338	.002

Environmental parameters	16S OTUs		Functional genes	
	<i>r</i>	<i>p</i>	<i>r</i>	<i>p</i>
Total nitrogen	.317	.001	.654	.001
Total phosphorus	.376	.001	.707	.001
Total sulphur	.294	.004	.435	.001
NH ₄ ⁺ -N	.012	.434	-.017	.549
NO ₃ ⁻ -N	.104	.156	.028	.311
Soil organic carbon	.352	.003	.674	.001
Available nitrogen	.080	.198	.289	.004
Rapidly available phosphorus	.393	.002	.739	.001

VPA was performed to analyse the contributions of environmental variables to microbial community structure. A substantial proportion (52.92%) of the variations in soil microbial community structure could be explained by the selected environmental factors; specifically, 9.88%, 33.39% and 4.76% of the variations could be explained by vegetation factors (including plant biomass and plant diversity), soil nutrients (including TN, TP, NH₄⁺-N, NO₃⁻-N, SOC and RAP) and soil pH, respectively (Fig. [S3A](#)). Even higher proportion (86.2%) variations could be explained for the microbial functional gene structure (Fig. [S3B](#)). These results showed that soil

nutrients were highly associated with soil microbial taxonomic structure and potential metabolic function in both sampling sites.

4 DISCUSSION

In recent decades, our knowledge on soil microbial communities has expanded rapidly with the development of new sequencing methods by passing the need for isolations of microorganisms (Drenovsky, Vo, Graham, & Scow, [2004](#); Torsvik & Ovreas, [2002](#)). In this study, the microbial taxonomic composition obtained by Illumina sequencing showed that species diversity significantly increased under the degeneration from AM to ASM, but the responses of different phyla could be varied. The phylum *Acidobacteria* is one of the most abundant soil bacteria and the relative abundance of dominant subgroups 3, 4, 6, 7 and 10 was increased in ASM, with low soil organic matter content, when compared with AM. Recent studies showed that the phylum *Acidobacteria* are in general oligotrophic ecosystems and revealed their adaptation to low substrate environments; for example, the proportion of *Acidobacteria* was reported to be significantly lower in nutrient-rich rhizosphere than in bulk soil (Kielak, Pijl, van Veen, & Kowalchuk, [2009](#)), and they have low abundance in nutrient-rich agricultural soil (Kielak et al., [2009](#); López-Lozano et al., [2013](#)). However, some dominant subgroups of *Acidobacteria* were also known to have a decreasing response to the soil environments with decreased available nutrients (Navarrete et al., [2013](#), [2015](#); Zhang et al., [2014a](#)). In our study, the dominant subgroup 17 was significantly decreased in ASM. These results suggested that a differential response of the *Acidobacteria* subgroups to ecosystem or environmental changes could be used as early warning indicator of soil management and plant-type successions (Navarrete et al., [2013](#), [2015](#); Zhang et al., [2014a](#)).

For the complicated and diversified interactions among different species (Olesen et al., [2007](#)), ecological network analysis is a sensitive, reliable and robust tool to reveal the interactions of microorganisms in complex biogeochemical processes (Deng et al., [2012](#); Zhou, Deng, Luo, He, & Yang, [2011](#)). The modularity, average degree and harmonic geodesic distance (HD) are crucial indicators reflecting the stability, robustness and resistance of complicated ecosystem networks (Deng et al., [2012](#)). According to the analysis results, the ecological network relationship was significantly different between AM and ASM. The network topological properties were more complicated in ASM than in AM, implying that the microorganisms in ASM might have more complicated interactions, in which microbial species could more stably coexist (Ding et al., [2015](#)). With the degenerative succession of AM to ASM, soil microbial community structures were changing towards a more complicated ecosystem, and this succession might be conducive to strengthening the resistance to external disturbance.

Understanding the mechanism of land-cover changes on the soil microbially mediated C cycling is essential for the estimation of the soil C pool. Analysing the differences in microbial C utilization ability was helpful to understand functional changes in the soil microbial community (Liu et al., [2013](#)). In the Biolog C utilization study, AWCD represented the utilization of C sources by microbes, and reflected the activity and physiological function of microbial communities (Liu et al., [2013](#)), while the Shannon index, McIntosh index and richness index could reflect the functional diversity of microbial metabolism (Wang et al., [2011](#)). Our studies indicated that soil microbial communities in ASM had higher activity than in AM for C utilization of different C components, such as polymers, carbohydrates, phenolic acids and carboxylic acids. The analysis of utilization of sole C substrates by microbial community structure indicated that soil microbes might have greater ability to decompose SOC in ASM than in AM, which could be the reason why SOC decreased in ASM.

Directly revealing the microbial metabolic activities and processes that mediated the SOC cycles caused by land-cover change is still difficult. In our study, the microbially mediated soil C cycling processes were further analysed using GEOCHIP technology. In previous studies, GEOCHIP was used to show that the detected functional gene signal intensities had significant correlations with environmental nutrient contents and that GEOCHIP could be used to link microbial communities with ecosystem processes and functions to a certain extent; for example, Yergeau, Kang, He, Zhou, and Kowalchuk ([2007](#)) showed a significant correlation between cellulase enzyme activity and the number of cellulase gene variants; Reeve et al. ([2010](#)) found a significant correlation between cellulose gene signal intensity and cellulose activity in the soil; Zhang et al. ([2014b](#)) showed that oxidizable organic carbon was significantly linked ($p < .05$) to the total abundance of genes involved in active organic carbon degradation (cellulose, hemicellulose and starch); Ding et al. ([2015](#)) explored that the total abundance of nitrification genes (*amoA* and *hao*) was negatively correlated ($r = -.46, p = .023$) with soil NH_4^+ -N, and total abundance of denitrification genes (*nirS* and *nirK*) was also negatively correlated ($r = -.54, p = .008$) with soil NO_3^- -N. In this study, the relative abundance of microbial C degradation genes related to labile C degradation was significantly higher in ASM site than in AM site, but the relative abundance of C degradation genes related to recalcitrant C was significantly lower in ASM site than in AM site. Therefore, a significant difference in soil C metabolic processes might occur with degenerative succession and soil microbes could be the facilitators of this process in the AM. Compared with 16S rDNA sequencing, however, GEOCHIP provided limited information for complex ecosystems due to the limitation of microbial functional gene probe number and type, sensitivity and quantitative capability; it might

be preferable to use a combination of technologies to better reveal the interaction in complex soil ecosystems in future.

Plant diversity and soil nutrients are important environmental factors that influence the soil microbial community and their ecological functions (Liu, Liu, Fu, & Zheng, 2008; Wardle et al., 2004). Species diversity and productivity of vegetation might greatly affect organic compounds and the litter diversity, which are the major soil resources and substrate satisfying microbial requirements (Bardgett & Shine, 1999). Changes in the plant community often lead to a corresponding change in both quantity and quality of soil organic matter (Carney & Matson, 2005; Chabrierie, Laval, Puget, Desaire, & Alard, 2003; Yang et al., 2014). SOC, as an important intermediate substance between plant and microorganisms, could be considered as the primary driving force in shaping microbial diversity and activity (Benizri & Amiaud, 2005; Eilers, Lauber, Knight, & Fierer, 2010). Some studies have confirmed that soil C availability, which could effectively regulate the changes in microbial community structure and microbial growth, had a substantial impact on microbial community composition and activity (Liu et al., 2014; Zhang et al., 2014b). Consistently, SOC was not only significantly correlated with microbial community composition, but influenced the functional genes involved in C and N cycling. Soil phosphorus as a limiting factor might indirectly affect the availability of other nutrients, such as N element (Janssens et al., 1998). Recent studies have reported that the availability of phosphorus not only limited microbial growth, but was also an important factor in driving microbial community structure and biogeochemical function (DeForest, Smemo, Burke, Elliott, & Becker, 2012; Demetz & Insam, 1999; Kuramae et al., 2010; Liu et al., 2014). In this study, we found that soil nutrients, including TN, TP, $\text{NH}_4^+\text{-N}$, $\text{NO}_3^-\text{-N}$, SOC and RAP, were highly associated with soil microbial taxonomic structure and potential metabolic function in both AM and ASM sites, suggesting that the soil degenerative succession might dramatically affect both soil nutrients and microbial communities synchronously.

In summary, to understand the effect of land-cover change on the soil microbial community and microbially mediated SOC loss, the soil microbial community structure and metabolic function related to C cycling in AM and ASM on Qinghai–Tibetan Plateau were analysed by Illumina sequencing, Biolog Ecoplate and GEOCHIP technologies. The results showed that the soil microbial community structure and diversity were significantly increased under degenerative succession from AM to ASM. Both microbial functional genes involved in C cycling and Biolog experiments indicated that ASM might decompose more SOC and released it in the form of CO_2 , which could further intensify the greenhouse effect. Therefore, the changes in land cover not only affected soil microbial community structure, but also affected their functional potential for

C decomposition. This might alter organic C dynamics, leading to increasing soil C losses and greenhouse gas emissions with degenerative succession of vegetation based on climate change or anthropogenic activities.

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AUTHORS' CONTRIBUTIONS

Y.Z., J.Z. and D.L. designed the experiments. Y.Z., X.L. and Y.D. wrote the main manuscript text. Y.Z., J.C., H.L., H.Y., X.W. and Y.S. performed the experiments. Y.Z., X.L. and D.Y. analysed the data. All authors reviewed the manuscript.

DATA ACCESSIBILITY

Sequencing data are accessible in NCBI SRA database with Accession No. [SRP096658](#). GEOCHIP data are accessible in NCBI database with Accession No. [GSE93158](#). The OTU table as well as the input and output files of the network analysis is accessible in Dryad database with <https://doi.org/10.5061/dryad.h781v>.