

## DIFFERENCES IN THE MICROBIAL POPULATION ASSOCIATED WITH THREE WETLAND TYPES IN THE SANJIANG PLAIN, NORTHEAST CHINA

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**Abstract.** The Sanjiang Plain is the largest freshwater wetland area in Northeast China and plays an important role in regulating climate stability for this region. However, agricultural activity has decreased wetland coverage with about 84 % since the 1950s. This has resulted in lowered water tables and degraded wetlands, with dryer marsh meadows and dry meadow coverage replacing wet marshlands. Here, we report investigations towards the soil microbial community composition and diversity in the different degeneration wetlands types. Bacterial and fungal communities in the soil types were compared using high resolution bar-coded pyrosequencing technology. The results revealed that the bacterial and fungal diversity was lower in wet marshland than in drier marsh meadow and dry meadows. The distribution of sequence reads into different bacterial and fungal phyla further differed between the soil samples. The wet marsh soil displayed a higher abundance of Proteobacteria but lower abundance of Acidobacteria, while the higher abundance of unclassified fungi but smaller fractions of Ascomycota and Basidiomycota than the other soil types. The results reported here demonstrate that soil bacterial and fungal communities change as a result of differences in the soil environment in the Sanjiang Plain.

**Keywords:** *community structure, microorganism; Miseq; wetland degradation; bacterial diversity; fungal diversity*

### Introduction

Wetlands are the most important terrestrial ecosystems in the world, having crucial environmental functions such as regulation of the carbon cycle (Keller, 2011), maintaining fresh water capacities (McJannet et al., 2012) and protecting biodiversity (Burton and Uzarski, 2009). Wetlands are areas with land and shallow water bodies, where the water tables permanently or periodically higher than surface level and with specific wetland ecological communities (Mausbach and Parker, 2001; Mitsch and Gosselink, 2007). Although natural wetlands occupy only 5–8 % of Earth's land surfaces (Mitsch and Gosselink, 2007), they are regarded as the “kidney” of the Earth, playing several key roles in biogeochemical processes such as pollutant degradation,

nitrification, denitrification, methanogenesis, methanotrophy, and iron and sulfate reduction (Davidsson et al., 1997; Gutknecht et al., 2006).

The Sanjiang Plain, covering an area of 10.89 million hectare, contains the largest freshwater wetland in Northeast China. It is also named *Deyeuxia angustifolia* (Kom.) wetland because this plant is the dominant species in the area. The local wetland environment is inevitable for climate stability, biodiversity protection, and greenhouse gas emission reduction in Northeast Asia. However, growing human populations have resulted in a decline of the wetlands. Whereas half of the Sanjiang Plain was covered by freshwater in 1950 (Zhao et al., 1999; Liu and Ma, 2000), approximately 84% of the wetlands have since been converted to farmland, especially to paddy fields (Liu and Ma, 2000). As a consequence of agricultural water use and decreased precipitation, the amount of water and water-covered surface area has decreased steadily, resulting in degraded wetlands. The original wet marshlands have changed to drier marsh meadows (recognized as a transitional state in wetland degeneration) and eventually into dry meadows, a fully degenerative state of wetland. Ecosystems have changed as well, with a decreased vegetation diversity, a fall in methane emission and a rise in carbon dioxide and nitrous oxide emission being prominent.

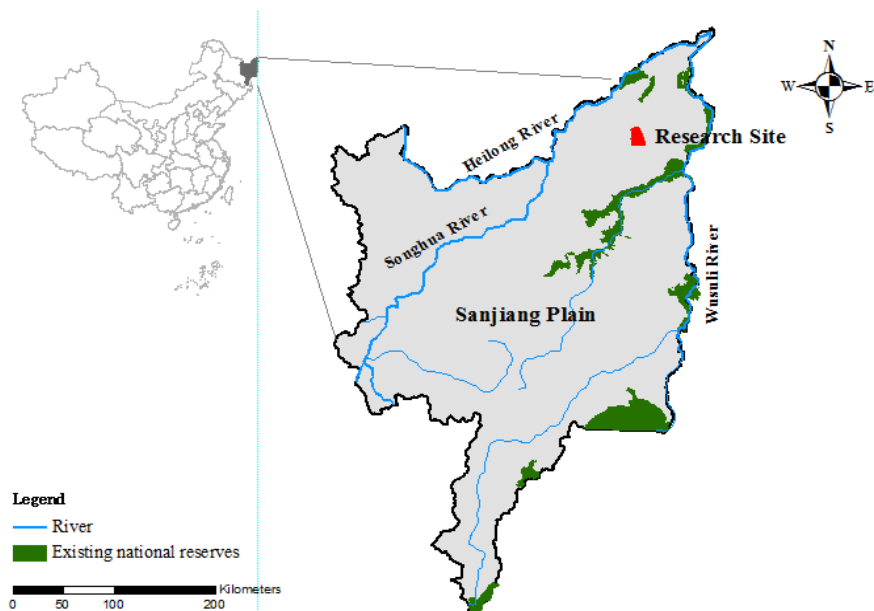
Several studies have evaluated the effects of wetland degeneration in the Sanjiang Plain on greenhouse gas emission (Song et al., 2013), nitrogen cycling between the atmosphere, vegetation and the soil (Sun, 2007), or soil microbial biomass and soil respiration (Huang et al., 2012), but little attention has been paid to the composition and diversity of soil microbial communities and how the microbial composition responds to wetland degeneration. With the complete array available from pristine wetlands to severely degenerative states, including three well-recognized wetland types along a water decline gradient within a small area, the site of the Sanjiang Wetland Experimental Station provided ideal site to study the effects on soil microbial ecosystems.

We hypothesized that changes in the soil microbial communities would be visible between wet marshlands, drier marsh meadows and dry meadows in Sanjiang Plain, and further that the variation in composition of soil fungi might exceed that of bacteria. To test these hypotheses, we collected soil samples from the three wetland types and estimated the composition, diversity and phylogeny of soil bacterial and fungal communities in these samples using high-throughput sequence analysis.

## Material and methods

### *Site description and soil sampling*

The study was conducted at the Sanjiang Wetland Experimental Station (47°35'N, 133°31'E), property of the Institute of Nature and Ecology of Heilongjiang Academy of Sciences, China (*Figure 1*). The local average monthly temperature ranges from -21.6 °C in January to 21.5 °C in July, with an annual average of 1.9 °C. The average annual precipitation is about 560 mm, with approximately 80% occurring between May and October. Three sites inside the station were selected for this study. Wet marshland was designated K0, a drier marsh meadow as K1 and dry meadow was called K2. The K0 site was approximately 20 m away from K2, and these were approximately 1 km away from the K1 site. All three sites are characterized by Quaternary sediments, and their soils were classified as Albic Boric Luvisols with a silty clay texture (Xi, 1993).



**Figure 1.** Map of the Research Site in the Sanjiang Plain, China

Soil samples were taken at a depth of 0–20 cm on 15 June 2014. Approximately 1kg soil was collected from five locations within each site and stored in polyethylene bags, placed in a container with ice and immediately transported to the laboratory. Upon arrival, approximately 2g of each soil sample was placed in a sterile micro centrifuge tube (2 mL) and stored at  $-80^{\circ}\text{C}$  for DNA extraction. The remaining of the samples was air-dried with the weight difference used as a measure of soil water content. Dried soil was used for the determination of other physiochemical properties: Soil pH was measured using a pH meter after mixing the soil with water (1:5 w/v) for 30 min. The soil total carbon (TC) and total nitrogen (TN) composition was determined using an Elemental analyzer (Vario EL III, Elementar Analyses system, Hanau, Germany). Nitrogen fractions  $\text{NH}_4^+\text{-N}$  and  $\text{NO}_3^-\text{-N}$  were measured by use of FLAstr 5000 analyzer (Foss Tecator AB Sweden Supply Company, Hoganas, Sweden).

### ***Soil DNA extraction and high-throughput sequencing***

DNA was extracted from 0.5g of each frozen soil sample with a MOBIO PowerSoil DNA Isolation Kit (USA) according to the manufacturer's instructions. The extracted DNA was diluted in 100  $\mu\text{L}$  TE (10 mM Tris-HCl, 1 mM EDTA, pH 8.0) and stored at  $-20^{\circ}\text{C}$  until used.

For PCR amplification and pyrosequencing we selected the V1-V3 region of bacterial 16S rRNA and the ITS1 region of fungal rDNA (Chakravorty et al., 2007; Bellemain et al., 2010). High-throughput sequencing was performed by the Shanghai Majorbio Biotechnology Company, Shanghai, China.

### ***Data analysis***

Obtained high-throughput sequences were analysed by using Mothur Software. The UniFrac statistical analysis tool was used to compare bacterial and fungal community compositions (Lozupone and Knight, 2005).

### ***Statistical analysis of effective and optimized gene sequences***

Using the simultaneous sequencing method for multi-samples, a barcode labeled gene sequences and forward primer sequences were introduced in the sequences of all samples. Sequences containing both barcode and forward primer sequences were then selected as effective sequences, after which the sequencing connectors and barcode sequences were removed. Subsequent data analyses were performed using the treated, effective sequences.

To obtain high-quality, accurate results from bio-informatic analyses, sequences were optimized by discarding sequences with lengths less than 150bp, those containing imprecise base calling, or those for which primer bases contained more than two mismatching sequences. The optimized sequences were used for subsequent statistical analyses.

### ***OTU-based analysis***

All sequences were identified to operational taxonomic units (OTU) for bio-informatic statistical analysis. Optimized gene sequences with gene lengths greater than 350bp were selected, compared with the SILVA database and then clustered. Clustering analysis was performed using the software packages mothur and chopseq ([http://www.mothur.org/wiki/Main\\_Page](http://www.mothur.org/wiki/Main_Page)).

### ***Bacterial community diversity and rarefaction curve***

Species richness and diversity of the bacterial community were characterized by Chao1 and the Shannon index, and the sequencing depth index was expressed as Coverage. Alpha-diversity of the bacterial community was measured at significance levels of 97% (0.03). The estimates were calculated by employing the tools Aligner, Complete Linkage Clustering, and Rarefaction of the RDP pyrosequencing pipeline.

The optimized gene sequences were randomly sampled. The sub-sampled sequences and the number of OTUs present in each were used to calculate a rarefaction curve. If the rarefaction curve tends to be flat then the sampling process is considered to be rational and further sampling is likely to produce few new OTUs, otherwise increased sampling will produce more new OTUs.

### ***Analysis on whole-sample similarity***

The Jost algorithm was used to compare the differences in OTUs from the three soil samples and to calculate the number of sequences from each OTU, thus obtaining a similarity relation between the samples. The selected OTUs had a similarity level of 0.03. Canonical Correlation Analysis for bacteria was conducted by R software.

## **Results**

### ***Soil Physicochemical Properties***

Soil characteristics such as pH, organic carbon content, total nitrogen, ammonium nitrogen, nitrate nitrogen, and soil water content of the three wetland types K0 (wet marshland), K1 (drier marsh meadow) and K2 (dry meadow) are listed in *Table 1*. All but one determined variables followed an increasing trend, from K0 to K2, while, as expected, soil water content was lower in K2 than in K0.

**Table 1.** chemical properties of soil samples

Sites	pH	Organic C(g.kg-1)	Total N (g.kg-1)	Ammonium nitrogen(mg.kg-1)	Nitrate nitrogen (mg.kg-1)	Soil water content (%)
K0	5.56±0.01 <sup>A</sup>	42.32±0.12 <sup>A</sup>	2.27±0.01 <sup>A</sup>	17.47±0.56 <sup>A</sup>	4.25±0.07 <sup>A</sup>	185±0.11 <sup>C</sup>
K1	5.66±0.02 <sup>B</sup>	44.23±0.19 <sup>B</sup>	2.70±0.02 <sup>B</sup>	18.51±0.56 <sup>B</sup>	4.41±0.08 <sup>B</sup>	86±0.08 <sup>A</sup>
K2	5.82±0.01 <sup>C</sup>	47.91±0.16 <sup>C</sup>	2.88±0.02 <sup>C</sup>	20.17±0.56 <sup>C</sup>	5.15±0.05 <sup>C</sup>	75±0.10 <sup>A</sup>

Different capital letters in the same column identify significant differences at 0.05 level among parameters. K0, K1, K2 represent wet marshland, drier marsh meadow and dry meadow, respectively.

### ***Diversity of bacterial and fungal communities***

The diversity of bacterial and fungal communities was calculated from 16S rRNA sequences and ITS rDNA sequences at the 3 % level, respectively, among the three samples. In total 660, 683, and 636 OTUs for bacteria and 199, 291, and 260 OTUs for fungi were identified in samples K0, K1 and K2, respectively (*Table 2*). The diversity indices showed that the drier marsh meadow K1 had the highest Shannon's diversity index and the lowest Simpson's index compared with the other samples. The  $S_{\text{chao}}$  estimator of the three samples was in the order K1>K2>K0. Thus, all diversity indices showed that the bacterial and fungal community compositions varied between the three types in that they were most diverse in drier marsh meadow soil (*Table 2*).

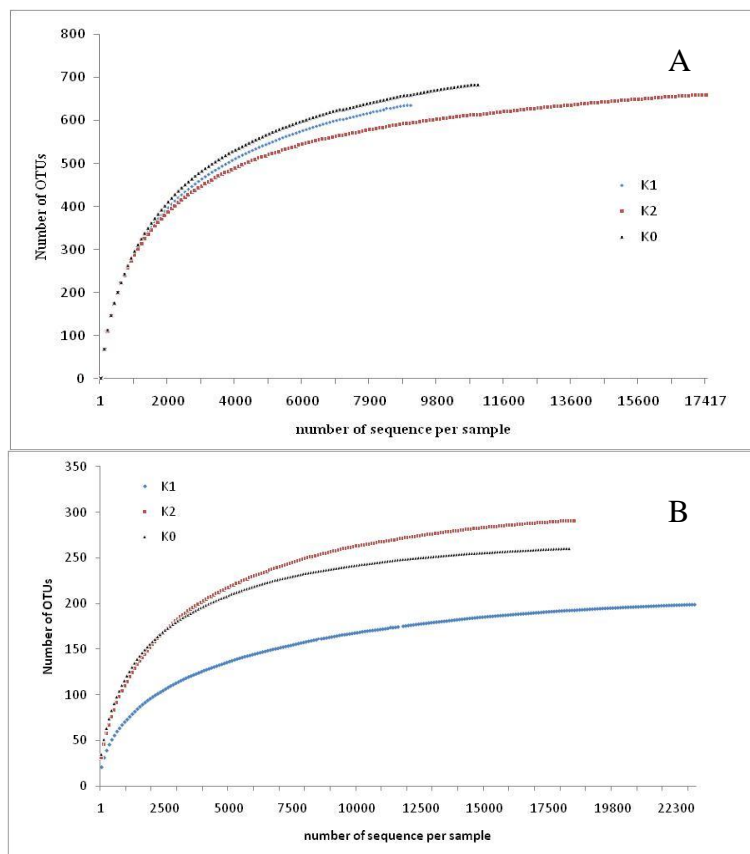
### ***Rarefaction curves of bacterial and fungal sequences***

Rarefaction curves were calculated by plotting the number of OTUs at the 3% level (*Figure 2*). At that level, the curves were increased at the rate of OTUs detection, indicating that the reads analysis evaluated almost the full extent of taxonomic diversity at the species level, and the coverage of the reads of bacterial and fungal sequences was estimated as above 98% (*Table 2*).

**Table 2.** Diversity indices for obtained bacterial and fungal sequences from three wetland soils in Sanjiang plain, NE China.

	Bacteria						Fungi					
	Reads	OTUs	Coverage (%)	$H'$	$D$	$S_{chao}$	Reads	OTUs	Coverage (%)	$H'$	$D$	$S_{chao}$
Wet marsh wetland(K0)	17417	660	99%	5.19	0.0156	709	18375	199	99.8%	2.33	0.0920	206
Drier marsh meadow wetland (K1)	10816	683	98%	5.26	0.0129	750	18299	291	99.8%	3.25	0.0782	301
Dry meadow wetland (K2)	8955	636	98%	5.22	0.0131	745	22953	260	99.8%	3.49	0.0762	268

OTUs:Operational taxonomic units,  $H'$ : Shannon's diversity index,  $D$ :Simpson's index



**Figure 2.** Rarefaction curves for the bacterial 16S (panel A) and fungal ITS (panel B) rRNA sequences obtained. Operational taxonomic units (OTUs) were calculated based on the 3% level.

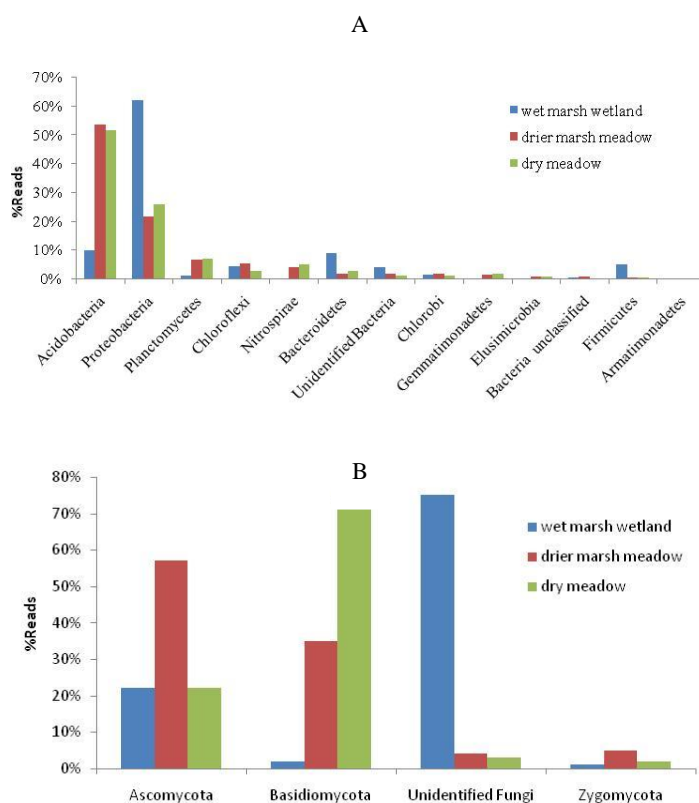
### **Compositions of bacterial and fungal communities**

Bacterial phyla were determined using the classifier tool at the RDP website. The obtained sequences were classified into (in decreasing order): 15432 Proteobacteria, 12118 Acidobacteria, 1984 Bacteroidetes, 1533 Chloroflexi, 1530 Planctomycetes, 924 Firmicutes, 899 Nitrospirae, 471 Chlorobi, 332 Gemmatimonadetes and 150 Elusimicrobia; 933 clones remained unclassified. About 90% of all 37,188 bacterial clones belonged to six taxonomic phyla: Proteobacteria (42%), Acidobacteria (33%), Bacteroidetes (5%), Chloroflexi (4%), Planctomycetes (4%), Firmicutes (3%) and Nitrospirae (2%).

The distribution of clones into the different bacterial divisions among the three high-throughput clone libraries was uneven, as shown in *Figure 3A*. Notably, Proteobacteria were highly over-represented in wet marshland, while the abundance of Acidobacteria, Chloroflexi and Chlorobi was higher in the drier marsh meadow soil (K1) than in the other two types. In contrast, Planctomycetes, Nitrospirae and Gemmatimonadetes were more abundant in dry meadow soil K2 in comparison to the other two types. Highest proportions of Bacteroidetes, Firmicutes and unidentified bacteria were observed in wet marshland (K0).

Fungal sequences were divided into 19470 Ascomycota, 23384 Basidiomycota, and these were again unevenly distributed among the three sample types (*Figure 3B*). The abundance of Ascomycota (56.56%) and Zygomycota (72.65%) was higher in the drier

marsh meadow soils(K1) than in the other two types, while Basidiomycota were remarkably abundant in dry meadow samples(K2), and the highest proportion of unidentified fungal sequences(75.12%) was obtained from wetmarshland (K0). Thus, the dominant fungal phylum was different in each type of wetland.

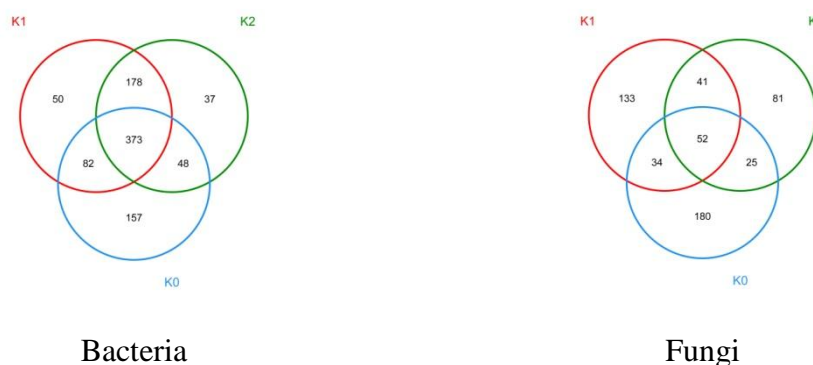


**Figure 3.** Distribution of the bacterial (panel A) and fungal (panel B) rRNA gene sequences in phyla obtained from wet marsh wetland (K0), drier marsh meadow wetland (K1) and dry meadow wetland (K2)

### **Comparison of bacterial and fungal community composition between the three wetlands**

Shared OTUs of bacteria and fungi among the three wetlands are shown in a Venn diagram in *Figure 4*. The analysis identified that 52 fungal and 373 bacterial OTUs were shared among all three wetland samples, while 41 OTUs of fungi and 178 OTUs of bacteria were shared between drier marsh meadow and dry meadow samples, 34 fungal and 82 bacterial OTUs were found in both wet marshland and drier marsh meadows, and 25 fungal and 48 bacterial OTUs were shared between wet marshland and dry meadow soils. Numbers of unique OTUs were highest in wet marshland K0, both for bacteria (180) and fungi (157), and lowest in dry meadow K2 (81 and 37, respectively).



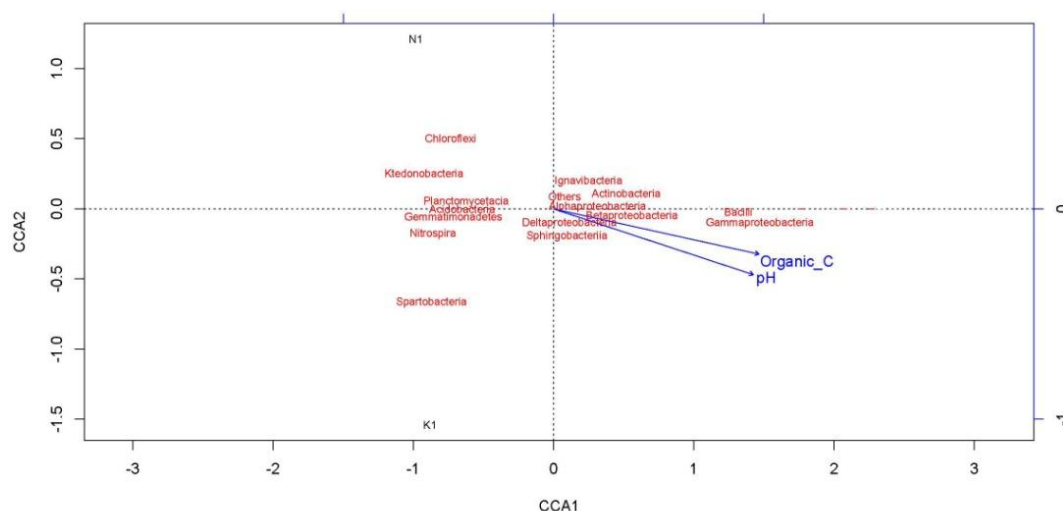


**Figure 4.** Venn diagram showing shared and unique OTUs identified for bacteria (left) and fungi (right) obtained from wet marsh wetland(K0),drier marsh meadow (K1) and dry meadow soils (K2).

### Long-term wetland degeneration alters soil microbial community composition

The statistical significance of differences in bacterial and fungal community compositions for all sequences was analysed by UniFrac (Table 3).The difference between total clones obtained from K0 and K1 was highly significant for bacteria. Highly significant differences at the phyla level are indicated in bold in the table. Only one difference was highly significant for Fungi (Agaricomycetes between K0 and K2).

Finally, a Canonical Correlation Analysis was performed, the results of which are presented in Figure 5. A negative correlation was observed between abundance of Acidobacteria and pH and soil organic carbon content. Thus, soil pH and carbon content are the major factors governing the abundance of Acidobacteria in the wetlands of Sanjiang Plain.



**Figure 5.** Canonical Correlation Analysis for bacteria in Sanjiang Plain

**Table 3.** UniFrac *P*-test values for bacterial and fungal sequences at total clones and individual phylum level for the three wetland soils in the Sanjiang Plain, NE China

Bacteria												
Sample	Total clones		Acidobacteria		Alpha-proteobacteria		Planctomycetacia		Delta-proteobacteria		Beta-proteobacteria	
	K1	K0	K1	K0	K1	K0	K1	K0	K1	K0	K1	K0
K1	-	<b>0.001</b>	-	0.007	-	<b>0.001</b>	-	0.010	-	<b>0.003</b>	-	0.022
K2	0.313	0.012	0.461	<b>0.002</b>	0.532	<b>0.002</b>	0.767	0.031	0.695	<b>0.004</b>	0.632	0.017
Fungi												
Sample	Total clones		Dothideomycetes		Sordariomycetes		Agaricomycetes		Leotiomycetes		Zygomycota	
	K1	K0	K1	K0	K1	K0	K1	K0	K1	K0	K1	K0
K1	-	0.823	-	0.415	-	0.151	-	0.069	-	0.812	-	0.620
K2	0.779	0.996	0.047	0.095	0.322	0.537	0.162	<b>0.005</b>	0.274	0.237	0.447	0.339

Statistically significant findings at or below the 0.005 level are given in bold.

## Discussion

The diversity of bacterial and fungal communities as determined in this study was lower in samples from wet marshland than in the degraded wetland types, as estimated by the number of OTUs, and as indicated by the Shannon's diversity, Simpson's and  $S_{chao}$  indices at phylum level (Table 2). This suggests that during the process of marsh degenerating into dry meadow the diversity of soil bacteria and fungi increases. Marsh wetland is characterized by perennial water levels that are characteristically low in oxygen. This restricts the growth of soil microbes and limits microbial diversity. Drier marsh meadow and dry meadows, however, provide better conditions for a diversity of microbes because their soils have lower water content and much more oxygen, which promoted soil microbial diversity. This is consistent with the results of Li (2011), who found that increased water content in wetlands decreased the oxygen amount and restrained the growth of fungi. Xu (2004) also found that a high soil water content restrains fungal diversity of valleys wetlands in Changbai Mountains, China (Xu et al., 2004). The results presented here also suggest that soil microbial diversity increases as a result of degeneration of wetlands in the Sanjiang Plain.

The difference in bacterial and fungal community compositions between the three wetland types was shown in Fig. 1, and their statistical significance presented in Table 3, suggest that the degeneration from marsh wetlands to drier environments significantly alters the soil bacterial and fungal community compositions. Liu (2014) also observed changes in the bacterial composition between such environments. Several studies have been conducted on soil microbial composition of various kinds of wetlands around the world (Dedysh, 2011). The majority of available research demonstrated that the proportion of Acidobacteria and Proteobacteria was higher than any other bacterial phyla in wetlands (Hartman et al., 2008; Ausec et al., 2009; Pankratov et al., 2011). For example, Ausec et al. (2009) observed Acidobacteria as the dominant phylum present in bog soils (41.6%) and in fen soils (23.7%) sampled in Slovenia. Hartman et al. (2008) determined 38.1% Acidobacteria, 17.4% Alpha proteobacteria and 9.7% Actinobacteria as the major bacterial phyla in wetland soils of a North Carolina coastal plain, but they did not detect any Chloroflexi (Hartman et al., 2008). Kanokratana et al. (2011) found that Acidobacteria (35.0%) and Proteobacteria (37.9%) dominated in soil from a tropical peat swamp forest in Thailand, by analyzing 280 clones of full length 16s rRNA sequences. The result of our research is consistent with those studies, as we found about 90% of all bacterial clones belonged to six taxonomic phyla, of which Proteobacteria dominated (Figure 3). The proportion of Proteobacteria was higher in the wet marshland than in the degraded types, but the fraction of Acidobacteria was higher in the two dryer types. This indicates that the composition of bacteria has changed during wetland degeneration. Li (2015) pointed out that Proteobacteria was the dominant phyla in the wetland of the Wuliangshuai eutrophic lake, with an even higher fraction of Proteobacteria detected than in our findings (Li et al., 2015). Proteobacteria seem to dominate in water-covered surfaces but their numbers decrease while Acidobacteria increase during transition to drier environments. The results presented here strongly suggest that the composition of bacterial communities in the three wetland types of the Sanjiang Plain have undergone changes due to a fall of the water table.

The composition of the fungal community seems to be less complex than that of bacteria. The proportions of the three fungal phyla Ascomycota, Basidiomycota, and

Zygomycota as well as unclassified fungi all varied in the three analysed wetland types. The dominant fungi in the types differed, from unclassified fungi in the wet marshland to Ascomycota in the drier marsh meadow and Basidiomycota in the dry meadow samples.

We thus infer that: 1) the soil nutrients in the three wetland types are different. For example, the wet marshland soil had the highest water content, which combined with low oxygen levels does not support fungal growth very well. This would explain the low fungal diversity and relatively large fraction of unclassified fungi observed. In contrast, a lower soil water content and higher oxygen levels in the drier marsh meadow and dry meadow sites better supported fungal growth and stimulated fungal diversity. 2) The composition of dominant vegetation correlated with the soil fungal composition. Wang (2016, unpublished) observed that with the decrease of the wetland area of the Sanjiang Plain the composition of plants significantly changed due to agricultural development and other human activities. Although *Deyeuxia angustifolia* was the dominant species in all sampled environments, its proportion was different for each type. As a result, the vegetative was the composition and decomposition ratio differed as well. For example, the litter composition was relatively simple and decomposed rather slowly in the marsh wetland. This correlated with a simple fungal composition and low diversity. The drier marsh meadow and dry meadow lands had a better soil environment to promote higher plant diversity, with a more complex litter composition that decomposed faster, correlating with a fungal composition dominated by Ascomycota and Basidiomycota. Tang (2012) found that soil microbial community compositions change according to water logging time, plant diversity and altitude. Zhao (2011) inferred that soil microbial community compositions were different in different plant diversity wetlands. Nevertheless, other studies reported that plant communities, waste composition and soil nutrients did not affect the composition of soil microbial communities (Tscherko, 2005; Andersen, 2010). Hence, the underlying causes for variation of soil microbial community composition can be complex and would require more research to be explained.

As reported by others, the abundance of Acidobacteria correlated negatively with soil organic carbon levels (Smit et al., 2001; Fiereret al., 2007) and soil pH (Jones et al., 2009; Rousk et al., 2010); the Canonical Correlation Analysis presented here also suggest this negative correlation. Thus soil pH and soil carbon are major factors governing the abundance of Acidobacteria in the wetlands of the Sanjiang Plain.

In conclusion, the conversion from wet marshlands to dry meadow increased the diversity of bacterial and fungal community and altered the community composition. The permanent conversion of submerged marsh wetland into dry meadow increased the abundance of Acidobacteria, Planctomycetes, Ascomycota, and Basidiomycota but decreased the abundance of Proteobacteria and unclassified fungi. These changes correlated with soil pH and organic carbon content, which were considered main impact factor on soil microbial composition.

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