Effects of drought stress on soil bacterial community in the West foot of Daxing'an Mountains

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Abstract. Global warming poses a serious threat to agriculture and natural systems, in part because of the change of soil moisture content, which changes soil microbial communities and ecological processes. Soil water content is the main factor limiting the growth of plants in soil. Microbial communities rely on soil water to complete their activities, and reveal the changes of underground microbial communities under different soil moisture content, which will help us to further understand the potential impact of climate change on soil ecosystem. To investigate the soil bacterial community structure, we established experiment indoor in the West foot of Daxing'an Mountains with manipulative water content treatments consisting of 20%, 15%, 10%, 5%, 0%. Results showed that bacterial community composition varied significantly with altered drought stress, but community richness did not. The relative abundance of Actinobacteria increased with the increase of drought stress. Proteobacteria, Acidobacteria and Gemmatimonadota decreased with the increase of drought stress, actinobacteria was more likely to accumulate or maintain stable under drought stress, bacterial communities can responding directly to changes in soil moisture.

1 Introduction

Global climate change may have considerable impacts on hydrological cycles worldwide, which may result in increased precipitation, higher evaporation rates, and the uneven distribution of rainfall[1]. Some regions may witness significant changes in the timing of dry and wet seasons, which might trigger increases in both droughts and floods[2]. Global warming poses serious threats to agricultural and natural systems, partly through its capacity to alter soil microbial communities and ecological processes due to variations in the content of soil water content[3]. Immense diversity of microorganisms that live belowground contributes significantly to shaping aboveground biodiversity and the functioning of terrestrial ecosystems[4]. Zhang et al analyzed the distribution and variation of temperature and precipitation in the Western foot of Daxing'an Mountains of China in 1960—2015, over the past 56 years, it has shown a trend of warming and drying[5]. Consequently, uncovering the effect of drought stress on the microbial community composition in the Western foot of Daxing'an Mountains will considerably enhance our understanding of the potential impact of climate change on arid ecosystems and their functions.

Soil microorganisms are active factors in the soil ecosystem. They play an important role in soil formation and development, soil organic matter decomposition, material and energy input, nutrient conversion, and fertility evolution. They are important links in the material cycle ecological chain[6,8]. The Soil water content is one of critical environmental factors that influence microbial community structure and function [7], and the microbial community rely on soil moisture to complete their activities and life spans[6]. A high SWC decreases microbial degradation rates because of low oxygen supply; a low soil water content decreases both microbial mobility and microbial degradative activity by reducing the diffusion of soluble substrates[9]. Researchers have observed increased rainfall might alter the elemental compositions of karst soils, and bacterial communities are likely to be more sensitive to variations in soil moisture in contrast to their fungal counterparts[10]. Microcosm-based experiments with tropical or subtropical soil have shown that soil water content exerts more significant effects on bacterial diversity and structure compared to Soil temperature [11-12]. Soil bacterial communities can provide biologically relevant insights on the impacts of land use on soil ecosystems[13]. A diverse adaptation strategy of
different bacterial groups. Actinobacterial could play an important role in nitrogen fixation, Bacillus and Pseudomonas can form endospores under drought stress, thereby resisting the variation caused by altered precipitation regimes,[14], the phylogenetic shifts in the composition of soil microbial communities are correlated with community function [15,16]. How the bacterial community structure and potential function without plants respond to water changes is poorly understood.

For this study, we conducted a controlled soil water content experiment in 5 gradient about the black soil of Daxing'an Mountains in China. Control variable method, high-throughput sequencing and bioinformatics were combined. The main aim is to determine the effects of the drought stress increased on soil bacterial communities and potential functions, to explore the direct effect of water content on soil microbial community.

2 Materials and methods

2.1 Experimental design

In the scientific observation and Experiment Station of soil management and ecological restoration of tnixe, chenbalhu banner, Hulunbuir City, the West foot of Daxinganling Mountains, 160kg soil was taken, mixed evenly, and dried in the room.

Five treatments were set up in the experiment. The relative water content of soil was 20% (water needed for normal plant growth), 15%, 10%, 5% and 0% respectively. Each treatment was repeated three times. The soil after shade drying was divided into 15 parts, each 10kg. The soil was placed in a plastic flowerpot of 48.4cm×32.2cm×25.7cm. The soil without water was marked W0. The other groups were marked W20, w15, W10 and W5. After 72 hours of cultivation in gradient incubator (lt-36vl) (the preliminary experiment found that the soil microbial community would maintain a new stable state after 72 hours of cultivation), the soil was taken after cultivation, the three repeated samples were grouped at the level of 97% similarity, 3941 bacterial OTUs were formed. At the lower level of classification, there were 37Class, 104 orders, 265 families, 426 genera, and 1512 species found in the West foot of Daxing'an Mountains soils. Across all samples, the dominant soil bacterial phyla were Actinobacterial(31.94%), Proteobacterial(26.70%), Acidobacterial(10.22%), Bacteroidota(8.93%), Gemmatimonadota(6.92%), Chloroflexi(6.64%), Verrucomicrobiota(2.27%), Myxococcota(1.82%), and Planctomycetes(1.10%), which collectively accounted for 96.08%~97.18% of all taxon sequences(Fig.1B).

2.2 16S metabarcoding and bioinformatics

Total soil DNA was extracted from replicate samples (n=5) by using DNeasy® PowerSoil® Pro Kit. After monitoring the DNA integrity and purity on 1% agarose gels, PCR was carried out using the bacterial-specific primer pair 515F (GTGCCAGCMGCCGCGGTAA) and 806R (GGACTACHVGG GTWTCTAAT) with a 6bp error correcting barcode unique to each sample. DNA was amplified and then agarose gel was used to detect the quality of DNA. Then the concentration of PCR product was mixed with equal concentration, and then the GeneJET gel Recovery Kit (Thermo Scientific) and 1 Gene JET were used. The PCR product was purified by agarose gel electrophoresis with TAE concentration of 2%, and the target band (size of 400–450bp sequence) was recovered by tapping. The library was constructed by using trueq DNA PCR free library preparation kit of illustra company. After qubit quantification and library test, the constructed library was sequenced by novaseq 6000 (Beijing Nuohu Zhiyuan bioinformatics Co., Ltd.).

2.3 Statistical analysis

The alpha diversity of samples was calculated with alpha_diversity.py in QIIME with the lowest number of sequences of all samples. The difference was analyzed by Kruskal Wallis h test, and Simpson and ACE index were calculated to characterize the bacterial community α Diversity. Based on Bray Curtis distance matrix PCA was used to analyze the bacterial community structure.Lefse multi-level species difference discriminant analysis was used to indicate species, and nonparametric Kruskal Wallis (kw) sum rank test was used to detect species richness differences among different groups, and significant species were obtained and the significance difference of species in multiple groups of samples is analyzed.SigmaPlot was used to fit the quadratic function of different species with soil water content.

3 Rresult

3.1 Community composition of bacterial in soils of in the West foot of Daxing'an Mountains

A total of bacterial sequences from the 15 soil samples were obtained from a total of 1167155 sequences. The amount of sequencing data is enough(Fig.1A). When grouped at the level of 97% similarity, 3941 bacterial OTUs were formed. At the lower level of classification, there were 37Class, 104 orders, 265 families, 426 genera, and 1512 species found in the West foot of Daxing'an Mountains soils. Across all samples, the dominant soil bacterial phyla were Actinobacterial(31.94%), Proteobacterial(26.70%), Acidobacterial(10.22%), Bacteroidota(8.93%), Gemmatimonadota(6.92%), Chloroflexi(6.64%), Verrucomicrobiota(2.27%), Myxococcota(1.82%), and Planctomycetes(1.10%), which collectively accounted for 96.08%~97.18% of all taxon sequences(Fig.1B).

3.2 Dissimilarity of bacterial communities between different drought stress

We used the ACE index to estimate the richness of bacterial communities, and the Simpson index to estimate the diversity of bacterial communities in the 5 different water content. The results indicated that
richness of the bacterial communities was no significant change (Fig. 1C), but the diversity followed the different trend. The highest was for the W0, followed by W5, W10, W15, and W20. The diversity of samples belonging to the W0 exhibited a greater variation range significantly than for the other soil water content, reflecting an association between community structure and soil water content, with the increase of drought stress, the species diversity increased (Fig. 1D).

Dissimilarities in the bacterial communities from the 5 soil water content were identified with a PCA biplot (Fig. 2). Notably, the composition of the bacterial communities was significantly affected by the soil water content (ANOSIM test, \( R = 0.8652, p=0.001 \)). Comparing the differences of bacterial communities between each soil water content, indicated that W5 and W10 were the most similar, whereas W0 compared with other soil water content exhibited the largest dissimilarity.

Fig. 1. Rank-Abundance curves (A) and Soil bacterial community abundance at the phylum level (B) and the Significance analysis of ACE(C) Simpson index (D) index under different soil water content. at the phylum level. 

Using LefSe analysis, we obtained bacterial biomarkers from phyum to species for each soil water content. There were 19 biomarkers identified in W20, 11 in W15, two in W10, two in W5 and 28 in W0 (Fig. 3).
3.3 Changes of bacterial community under different drought stresses

The significance test results demonstrated that altered soil water content significantly affected the abundance of Actinobacteria, Proteobacteria, Acidobacteria and Gemmatimonadota from the top 10 phyla of abundance (Table 1).

The significance analysis of the top 27 genera (Relative abundance>1%) with relative abundance showed that soil water content had a significant effect on 16 genera, and 16 genera belonged to 6 different phylum, which 8 belonged to Actinobacteria, 3 belonged to Proteobacteria, 2 belonged to Acidobacteria, gemmatimonadota and bacteroidota only have 1 respectively (Table 2). Among the 16 genera, the relative abundance of 8 genera of Actinobacteria and ellin6055 increased with the increase of drought stress, which they may belong to drought tolerant bacteria. The relative abundance of Lysobacter and g_norank_f__norank_o__vicinamibacteria decreased with the aggravation of drought stress that was sensitive to water content. With the increase of drought stress, the relative abundance of Sphingomonas, g_norank_f__Microscillaceae and g_norank_f__norank_o__norank_e__Gitt-GS-136 decreased first and then increased, and the adaptability was strong. They could adjust its adaptability with the change of soil water content; g_norank_f__Microscillaceae increased first and then decreased with the increase of drought stress (Fig.5).

| Species name     | P Value | Kruskal-Wallis H test |
|------------------|---------|----------------------|
| Actinobacteriota | 0.020   | *                    |
| Proteobacteria   | 0.036   | *                    |
| Acidobacteriota  | 0.016   | *                    |
| Bacteroidota     | 0.453   | ns                   |
| Gemmatimonadota  | 0.012   | *                    |
| Chloroflexi      | 0.106   | ns                   |
| Verrucomicrobiota| 0.264   | ns                   |
| Myxococciata     | 0.277   | ns                   |
| Patescibacteria  | 0.122   | ns                   |
| Firmicutes       | 0.271   | ns                   |

NOTE: * has significant difference at P<0.05 level, and "ns" has no significant difference, the same as below.
In these four phyla, the relative abundance of Actinobacteria increased with the increase of drought stress. Compared with W20, the abundance of W15, W10, W5 and W0 increased respectively 12.06%, 37.65%, 43.40% and 57.66%, the highest abundance was in extreme drought; However, the abundance of Proteobacteria, acidobacteria an gemmatimonadota decreased with the increase of drought stress, indicating that actinobacteria was more likely to accumulate or maintain stable under drought stress (Fig. 4).

### 4 DISCUSSION

#### 4.1 Dissimilarity of bacterial communities between different drought stress

Our analyses indicate that soil water content significantly influence the composition of bacterial communities in the the West foot of Daxing'an Mountains. With the increase of drought stress increased species diversity and reached the maximum at W0. some studies have found that microbial community structure[17]and richness[18]change in response to increased precipitation, whereas other works did not observe substantial effects on microbial community...
composition[19-20]. Maybe the factor of water alter soil microbial communities by changing soil physical and chemical properties, nutrient cycling[21-23].

4.2 A diverse adaptation strategy of different bacterial groups

Four major response patterns were detected across all bacterial samples[21], the first pattern was that resource-limited populations, some bacterial increased in relative abundance with the increase in precipitation. By contrast, an opposite result was found which suggested that these genera belong to drought-tolerant populations. The third pattern showed that bacterial genera were most abundant.

According to the trend analysis, they also had drought resistance characteristics, and the relative abundance increased with the increase of drought stress. Lysobacter and g_norank_f_norank_o_vicina-mibacteria decreased with the aggravation of drought stress that were sensitive to water content. Sphingomonas, g_norank_f_Microscillaceae and g_norank_f_norank_o_norank_c_Gitt-GS-136

Consistent with the results of previous studies, but because there are no different plants and soil types, so the types of bacteria are different. The relative abundance of Proteobacteria, Acidobacteria and Gemmatimonadota decreased with the increase of drought stress. Actinobacteria were more likely to accumulate or remain stable under drought stress. Blastococcus, Arthrobacter, Nocardioides, Aeromicrobium bacteria belonged to Actinobacteria,
decreased first and then increased, and the adaptability was strong.

4.3 Actinobacteria have a rich abundance for under drought stress

Actinobacteria are known for their metabolic potential of producing diverse secondary metabolites such as antibiotics. Actinobacteria play an important role in nitrogen cycling in both desert and cultivated farm ecosystems. Actinobacteria is associated with environmental factors, such as soil type [24]. They are important for the cycling of carbon, nitrogen, phosphorus, potassium, and several other elements in the soil [25]. Some members of the phylum Actinobacteria may be able to live in an oligotrophic environment. Arocha-Garza [26] isolated 350 different strains from an oligotrophic desert oasis. The Actinobacteria can produce a variety of extracellular hydrolytic enzymes to degrade animal and plant residues, litter and other organic compounds in soil, enabling them to thrive in oligotrophic environments [27]. Our research indicated that drought stress had a significant effect on soil actinobacteria. The highest abundance of actinobacteria was found in all the varieties, and its abundance increased with the increase of drought stress. After significance analysis and trend fitting, it was found that 50% of the 16 genera responding to drought stress belonged to actinobacteria. It can be enriched to cope with drought in the natural precipitation. Lastly, relative abundance did not vary among different precipitation regimes for several soil bacterial populations. These bacterial genera were defined as the inactive population, thereby indicating a diverse adaptation strategy of different bacterial groups [21].

5 Conclusion

This detailed description of bacterial communities in 5 soil water content experiment of the West foot of Daxing'an Mountains soil indicates that community structures vary according to the drought stress. Actinobacteria could play an important role in arid soil. In the absence of crops, soil water content has a direct impact on bacterial communities. Next, we will further explore the function of actinomycetes in black soil and find out the beneficial bacteria in drought tolerant bacteria.

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