Article title: Variation of rhizosphere bacterial community diversity in the desert ephemeral plant Ferula sinkiangensis across environmental gradients

Authors: Zhang tao[1], Wang Zhongke[2], Lv Xinhua[3], Dang Hanli[4], Zhuang Li[5]

Affiliations: Shihezi University[1]

Orcid ids: 0000-0001-5003-2519[1]

Contact e-mail: 1838856694@qq.com

License information: This work has been published open access under Creative Commons Attribution License http://creativecommons.org/licenses/by/4.0/, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. Conditions, terms of use and publishing policy can be found at https://www.scienceopen.com/.

Preprint statement: This article is a preprint and has not been peer-reviewed, under consideration and submitted to ScienceOpen Preprints for open peer review.

Funder: Chinese National Basic Research Program

DOI: 10.14293/S2199-1006.1.SOR-.PPEUPDD.v1

Preprint first posted online: 16 June 2020

Keywords: Ferula sinkiangensis, rhizosphere bacterial community, high-throughput sequencing, soil depth, slope positions, soil physicochemical factors
Variation of rhizosphere bacterial community diversity in the desert ephemeral plant *Ferula sinkiangensis* across environmental gradients

Tao Zhang, Zhongke Wang, Xinhua Lv, Hanli Dang* and Li Zhuang*

1 College of Life Sciences, Key Laboratory of Xinjiang Phytomedicine Resource Utilization, Ministry of Education, Shihezi University, Xinjiang Shihezi, 832003, China.

* Correspondence: Hanli Dang email: 3497404295@qq.com; Li Zhuang email: 3033573705@qq.com

Abstract

*Ferula sinkiangensis* is a desert short-lived medicinal plant, and its number is rapidly decreasing. Rhizosphere microbial community plays an important role in plant growth and adaptability. However, *Ferula sinkiangensis* rhizosphere bacterial communities and the processes that drive its assembly remain unclear. On this study, based on Illumina HiSeq high-throughput sequencing, we explored the diversity, structure and composition of *Ferula sinkiangensis* rhizosphere bacterial communities at different slope positions and soil depths and their correlation with soil physicochemical properties. Our results revealed the heterogeneity and variation trends of *Ferula sinkiangensis* rhizosphere bacterial community diversity and abundance on a fine spatial scale (Slope position, soil depth, rhizosphere and non-rhizosphere) and Found Actinobacteria (22.7%), Proteobacteria (18.6%), Acidobacteria (14.0%), Gemmatimonadetes (10.1%) and Cyanobacteria (7.9%) were the dominant bacterial phyla in *Ferula sinkiangensis* rhizosphere soil. Among the physiochemical variables, there was a strong positive correlation between phosphorus (AP) and the diversity of rhizosphere bacterial community in *Ferula sinkiangensis* (p < 0.01). In addition, Soil physicochemical factors jointly explained 29.81% of variation in *Ferula sinkiangensis* rhizosphere bacterial community structure. Among them, pH largely explained the variation of *Ferula sinkiangensis* rhizosphere bacterial community structure (5.58%), followed by altitude (5.53%) and total salt (TS, 5.21%).

Keywords: *Ferula sinkiangensis*; rhizosphere bacterial community; high-throughput sequencing; soil depth; slope positions; soil physicochemical factors

1 Introduction

*Ferula sinkiangensis* is a short-lived medicinal desert herb, which was included in the
pharmacopeia of the People's Republic of China in 1977 [1]. Currently, it is only distributed in the yining region of xinjiang, China. Ferula sinkiangensis has important medicinal values, such as anti-cancer [2, 3], antioxidant [4], antibacterial [5, 6], anti-inflammatory [7], anti-influenza [5, 8] and anti-diabetes activities [9]. However, high medicinal value of Ferula sinkiangensis leads to the excessive exploitation by people [10]. As a result, the ecological environment of Ferula sinkiangensis habitat is severely damaged, and the population size decreases sharply. In addition, pests [11] and the low natural reproduction capacity of Ferula sinkiangensis (low seed yield and male sterility) [12] result in weak population recovery, which leads to a shrinkage of the distribution area of Ferula sinkiangensis, from 1400/ha in 1987 to 133/ha in 2012, and now the species number is still decreasing continuously [13]. Hence, the need to protect Ferula sinkiangensis is imminent.

Soil microbes play a major role in the function of terrestrial ecosystems. This includes decomposition of organic matter, nutrient cycling (carbon, nitrogen and phosphorus) and pollutant conversion [14-17]. In 1904, Hiltner first proposed the term “rhizosphere region” to describe soil regions affected by root exudates and litter [18, 19]. Root exudates and litter provide the microorganisms with the carbon and energy needed for survival [20, 21]. In turn, microorganisms regulate the morphology and physiology of plants [22, 23], promote plant growth and damage repair [24-26], increase plant tolerance to biological and abiotic stresses [27-29], and improve plant ecological adaptability. Thus, the rhizosphere microbiome is considered as the second genome to plant condition [30] and serves on a highly evolved external functional environment for plants [31], which are conducive to plant survival and population recovery. Although, we have learned that rhizosphere microbes have a positive effect on plant growth and development. However, it must be said that the information about the distribution and composition of the rhizosphere microbial community of Ferula sinkiangensis is not clear at present. Therefore, our primary purpose in this study was to explore the structure and diversity of the bacterial community in the Ferula sinkiangensis rhizosphere.

We also need to be aware of the interaction between microbes, plants and environmental factors. Some scholars founded that changing water and nutrient availability can affect root exudates and regulate the rhizosphere microbial diversity [26, 32, 33]. The soil pH also has a significant effect on the growth of microbial communities [34, 35]. Nitrogen deposition and precipitation can impact microorganisms [36]. Of course, there are many scholars who make full use of the relationship between plants, microbes and the environment. They modify the soil microbial structure by changing environmental factors to improve agricultural soil fertility. For example, Qin et al. regulated soil microorganisms and improved soil environment by adjusting biochar [37]. Zhou Jing et al. influenced fungal communities by increasing nitrogen. Similarly, we also hope to make use of the interaction between Ferula sinkiangensis and the environment and microorganisms to carry out the protection work of Ferula sinkiangensis. Unfortunately, it is not clear what kind of relationship exists between Ferula sinkiangensis and the environment and soil microorganisms. On this study, we investigated the effect of slope position...
and soil depth on rhizospheric soil microorganisms in *Ferula sinkiangensis* and the relationship between these microorganisms and soil physical and chemical factors.

Giving the spatial heterogeneity of microorganisms [39-41] and the influence of numerous soil physicochemical factors on microorganisms, we hypothesized that (1) the bacterial community structure and diversity of *Ferula sinkiangensis* rhizosphere are significantly different at various slope positions and soil depths and (2) there are only a few Soil physicochemical properties having dominant influence on the diversity and structure of the bacterial community in the *Ferula sinkiangensis* rhizosphere soil.

2 Results

2.1 Phylum and genus of dominant bacteria in *Ferula sinkiangensis* rhizosphere and non-rhizosphere soil

All samples contained 4,634,264 raw tags in total, filtering out the low quality tags and removing the chimeric sequences to obtain the final 4,405,120 effective sequences. On average, these reads were grouped into 4,017 bacterial OTUs and contained a total of 75,570 taxon tags per sample (Supplementary Table S1). In addition, the results of the rarefaction curves drawn based on OTUs (Supplementary Figure S1) suggested that the curve representing each soil sample eventually tends to be flat. With the increasing Sequences number, it is rarely accompanied by the increase of the number of new bacteria OTUs, which reflects the reasonable amount of data we sequenced.

Species sequence analysis identified 60 phyla and 901 genera of bacteria. The most abundant phyla were (Figure 1): Actinobacteria (22.7%), Proteobacteria (18.6%), Acidobacteria (14.0%), Gemmatimonadetes (10.1%), Cyanobacteria (7.9%), Bacteroidetes (6.9%), Planctomycetes (3.9%), Verrucomicrobia (3.5%), Firmicutes (3.4%), and Chloroflexi (3.2%). The most abundant genera were (Supplementary Figure S2): *RB41* (3.03%), *Sphingomonas* (1.64%), *Rubrobacter* (1.43%), *Gaiella* (1.10%), *Pseudarthrobacter* (1.01%), *Solirubrobacter* (0.85%), *Gemmatimonas* (0.71%), *Bacteroides* (0.63%), *Acinetobacter* (0.57%), *Adhaeribacter* (0.57%), *Haliangium* (0.47%), *Staphylococcus* (0.43%), *Rhodococcus* (0.42%), *Pseudomonas* (0.40%), *Bacillus* (0.38%); the remaining 76.59% include genera outside the top 30 of the annotated list and the parts not annotated.
Figure 1 Rarefaction curves of OTUs at 97% similarity for each sample. Legend: Longitudinal direction is the number of OTUs constructed based on the number of sequencing strips, the horizontal direction is the number of sequencing strips randomly extracted from a sample. Different samples are represented by curves with different colors.

2.2 Relationships between rhizosphere and non-rhizosphere, slope position, soil depth and dominant bacteria (phyla and genera)

Multivariate analysis of variance and LDA effect size analysis showed that Cyanobacteria, Actinobacteria, Acidobacteria, Chloroflexi and Firmicutes were significantly changed in the rhizosphere and non-rhizosphere. The specific abundance of Actinobacteria, Chloroflexi and Acidobacteria in the rhizosphere was significantly higher than that observed in the non-rhizosphere. However, the relative abundance of Cyanobacteria and Firmicutes in the non-rhizosphere was significantly higher than that observed in the rhizosphere. Moreover, Firmicutes, Planctomycetes and Verrucomicrobia were significantly affected by the slope position, as shown by a significantly higher relative abundance of Firmicutes at the bottom of the slope compared to the top of the slope. The opposite was observed for Verrucomicrobia. In addition, the relative abundance of Planctomycetes in the middle and top of the slope was significantly higher than at the bottom of the slope. Soil depth had a significant effect on the relative abundance of Cyanobacteria, Bacteroidetes, Acidobacteria and Planctomycetes. The relative abundance of Planctomycetes and Acidobacteria at a 0-10cm soil depth was significantly higher than at 25-40cm, and Bacteroidetes were more abundant at 0-10cm than at a 10-25cm soil depth. However, the relative abundance of Cyanobacteria at 10-40cm soil depth was significantly higher than that observed at 0-10cm soil depth (Figure 2 and Supplementary Table S2). Figure 2 also
shows specific differences in the distribution of different bacterial genera.

Figure 2 Results of intergroup LDA effect size (LEfSe) analysis revealed the specific bacteria phylum and genus distributed in the rhizosphere and non-rhizosphere regions under different slope positions and soil depths. Description: Each small circle at a different classification level represents a classification at this level and the diameter of the small circle is proportional to the relative abundance. Species with no significant differences are uniformly colored in yellow, the different species Biomarker follows the group for coloring.

2.3 Effects of slope position, soil depth, rhizosphere and non-rhizosphere on bacterial diversity and richness

The results of variance analysis (Table 1) based on the diversity indexes of each sample indicate that Shannon and Chao1 indexes, which represent diversity and richness, have significant differences at the top and bottom of the slopes. This means that the diversity and richness of the bacterial community were affected by the slope position. Specifically, the bacteria diversity and richness at the top of the slope were higher than that at the bottom of the slope. In addition, Shannon and Chao1 indexes showed significant differences in different soil depths, rhizosphere and non-rhizosphere. This indicates that the diversity and richness of bacterial community are influenced by soil depth, rhizosphere and non-rhizosphere. Specifically, the diversity and richness of bacterial community in the soil layer of 0-10cm was significantly higher than that in the soil layer of 10-25cm and 25-40cm. Furthermore, the diversity and richness of bacteria were significantly higher in the rhizosphere than in the non-rhizosphere.

Table 1. Variance analysis based on the diversity index of each sample.
### Table 1: Rhizosphere and Non-rhizosphere Soil Physicochemical Properties

|                | Shannon Index | Chao1 Index |
|----------------|---------------|-------------|
| Rhizosphere    | 9.690±0.848a  | 4801.479±525.306a |
| Non-rhizosphere| 8.365±2.163b  | 3633.340±1595.432b |
| E              | 9.423±1.145a  | 4657.891±540.111a |
| R              | 9.209±1.256ab | 4145.907±1338.738ab |
| S              | 8.452±2.492b  | 3848.430±1720.369b |
| 1              | 9.790±0.601a  | 4847.895±287.097a |
| 2              | 8.707±1.89b   | 3981.225±1463.036b |
| 3              | 8.587±2.189b  | 3823.108±1590.955b |

Legend: E, R and S represent the top, middle and bottom of slope respectively. 1, 2 and 3 represent soil depths of 0-10cm, 10-25cm and 25-40cm respectively.

#### 2.4 Relationship between slope position, soil depth and *Ferula sinkiangensis* rhizosphere soil physicochemical properties

Spearman correlation analysis showed that most variables including total phosphorus content (TP), ammonium nitrogen content (AN), nitrate nitrogen content (NN) and available phosphorus content (AP) are associated with altitude (Table 2). Specifically, altitude was negatively correlated with AN and NN. On the other hand, a significant positive correlation was observed between altitude and TP or AP. Total organic carbon content (TOC), total nitrogen content (TN), and total potassium content (TK) were positively correlated with AP. TOC was positively correlated with TN, total salt content (TS) and AP. TS was negatively correlated with pH, and TP was positively correlated with altitude. However, TP was negatively correlated with AN (Table 2). In addition, multivariate analysis of variance showed that slope position was significantly correlated with most soil physicochemical properties including TOC, TP, TK, NN, AN, AP, and TS. However, only a few soil physicochemical properties, including AP, TS and pH, had a significant correlation with depth. Specifically, TOC, TP, TK, AP, and TS were significantly higher at the top than the middle of slope. The contents of TP and AP at the top of slope were significantly higher than those at the bottom and middle of slope. Interestingly, the content of AN was significantly higher at the bottom than at the top and middle of slope. In terms of depth, the content of AP and pH in 0-10cm soil were significantly higher than those in 25-40cm soil (Supplementary Table S3).
Table 2. Spearman correlation coefficients between soil physicochemical properties across all samples.

|       | TOC  | TN   | TP   | TK   | NN   | AN   | AP   | TS   | pH   | Altitude |
|-------|------|------|------|------|------|------|------|------|------|----------|
| TOC   | 1.000|      |      |      |      |      |      |      |      |          |
| TN    | 0.792**| 1.000|      |      |      |      |      |      |      |          |
| TP    | 0.163 | 0.193| 1.000|      |      |      |      |      |      |          |
| TK    | 0.171 | 0.333| 0.067| 1.000|      |      |      |      |      |          |
| NN    | 0.356 | 0.277| -0.149| -0.079| 1.000|      |      |      |      |          |
| AN    | -0.200| -0.269| -0.448*| 0.012| 0.031| 1.000|      |      |      |          |
| AP    | 0.568**| 0.639**| 0.271| 0.472*| 0.069| -0.230| 1.000|      |      |          |
| TS    | 0.411*| 0.232| 0.004| 0.125| 0.287| -0.031| 0.073| 1.000|      |          |
| pH    | 0.038| 0.149| 0.002| 0.129| 0.077| 0.047| 0.213| -0.390*| 1.000|          |
| Altitude | -0.006| 0.157| 0.443*| 0.221| -0.629**| -0.425*| 0.419*| -0.093| -0.245| 1.000 |

Legend: TS, total salt content; AP, available phosphorus content; AN, ammonium nitrogen content; NN, nitrate nitrogen content; TP, total phosphorus content; TN, total nitrogen content; TK, total potassium content; TOC, total organic carbon content. \( r \)-value represents Spearman correlation coefficient, between -1 and 1, \( r < 0 \) is negative correlation, \( r > 0 \) is positive correlation *\( p < 0.05 \), **\( p < 0.01 \).
2.5 pH largely explained the variation of *Ferula sinkiangensis* rhizosphere bacterial community structure

Across all samples, soil AP showed a significant positive correlation with α-diversity of the bacterial community (r=0.538, p<0.01). Soil TP and TK and the α-diversity displayed a significant positive correlation (Shannon index, r = 0.495 and 0.405, respectively, p < 0.05, Table 3). In addition, distance-based redundancy analysis (db-RDA) also showed a correlation between soil physicochemical properties and the distribution of bacterial communities in the rhizosphere of *Ferula sinkiangensis* (Figure 3). All the soil physicochemical factors explained 29.81% of the variation in the rhizosphere bacterial community structure of *Ferula sinkiangensis*. The pH explained 5.58% of variation, altitude explained 5.53%, TS 5.21%, TP 4.90%, NN 3.89% and AP explained 3.60%. Among them, pH, altitude, TS and TP explained the largest proportion of the variation of the bacterial community structure in the rhizosphere of *Ferula sinkiangensis* (Supplementary Table S4).

Table 3. The spearman correlation analyses between the α-diversity index and soil physicochemical properties.

|                | Shannon index | Chao1 index |
|----------------|---------------|-------------|
|                | r             | p           | r             | p           |
| Altitude       | 0.136         | 0.107       | 0.175         | 0.383       |
| TOC            | 0.234         | 0.240       | -0.037        | 0.854       |
| TN             | 0.185         | 0.356       | -0.152        | 0.449       |
| TP             | 0.459         | 0.016*      | 0.114         | 0.571       |
| TK             | 0.405         | 0.036*      | 0.225         | 0.259       |
| NN             | -0.312        | 0.113       | -0.102        | 0.613       |
| AN             | -0.312        | 0.113       | 0.078         | 0.698       |
| AP             | 0.538         | 0.004**     | 0.240         | 0.228       |
| TS             | 0.027         | 0.892       | -0.035        | 0.862       |
| pH             | 0.317         | 0.498       | -0.190        | 0.343       |

Legend: r-value represents spearman correlation coefficient, between -1 and 1, r<0 is negative correlation, r>0 is positive correlation *p < 0.05, **p < 0.01.
Figure 3 Ordination diagram (samples-environment biplot) of db-RDA depicting environmental drivers of rhizosphere bacterial community composition of *Ferula sinkiangensis*.

2.6 Spearman correlation analysis between relative abundance of dominant bacteria (phyla and genera) and soil physicochemical properties

Although the relative abundance of bacterial phylum has a significant correlation with slope position, soil depth and root zone, the relative abundance of bacterial also has a significant correlation with soil Physicochemical Properties. Table 4 shows the relationship between the top ten dominant bacteria phyla and soil Physicochemical Properties. Specifically, TN, AP and TS have a significant positive correlation with the relative abundance of Actinobacteria; AP and TS showed significant positive relationships with Chloroflexi; pH showed significant positive relationships with Bacteroidetes; Altitude have significant positive relationships with Gemmatimonadetes; AN have significant positive relationships with Verrucomicrobia. Conversely, NN and TS significantly negative correlated with the relative abundances of Gemmatimonadetes; TN and AP showed significant negative relationships with Cyanobacteria. Interestingly, Firmicutes, Proteobacteria, Planctomycetes and Acidobacteria were not significantly related to each variable (Table 4). Moreover, the relationship between the relative abundance of the first thirty-five bacterial phyla and the Physicochemical Properties of the soil is shown in Supplementary Figure 3. In addition, correlation analysis between relative abundance of bacteria genera and soil
physicochemical properties showed that pH and altitude were significantly correlated with most bacteria genera. For example, pH has significant positive correlation with the relative abundance of *Adhaeribacter*, *Altererythrobacter*, *Gemmatimonas*, *Hymenobacter*, *Massilia*, *Opitutus*, *Rubellimicrobium* and *Sphingomonas* (Figure 4A). Altitude has significant negative correlation with *Blastococcus*, *Opitutus*, *RB41*, and *Rubellimicrobium* (Figure 4B).

**Figure 4** Spearman correlation between pH (A), altitude (B) and dominant bacteria. Description: “+” indicates positive correlation; “-” indicates negative correlation *p < 0.05, **p < 0.01.*
Table 4. The spearman correlation analysis between relative abundance of dominant bacteria and soil physicochemical properties.

|            | Actinobacteria | Firmicutes | Proteobacteria | Bacteroidetes | Planctomyetes | Acidobacteria | Gemmatimonadales | Chloroflexi | Verrucomicrobia | Cyanobacteria |
|------------|----------------|------------|----------------|---------------|---------------|---------------|------------------|--------------|-----------------|---------------|
| pH         | -0.355         | -0.003     | 0.054          | 0.434*        | 0.106         | 0.13          | 0.027            | -0.263       | 0.183           | 0.21          |
| TOC        | 0.346          | -0.026     | -0.101         | 0.049         | 0.026         | 0.026         | -0.377           | 0.311        | -0.049          | -0.291        |
| TN         | 0.420*         | -0.122     | 0.087          | -0.035        | -0.224        | -0.096        | -0.151           | 0.349        | -0.035          | -0.384*       |
| TP         | -0.026         | -0.279     | 0.314          | 0.278         | 0.012         | 0.12          | 0.178            | -0.065       | -0.125          | -0.12         |
| TK         | 0.246          | -0.114     | 0.026          | 0.071         | -0.006        | 0.3           | -0.007           | 0.337        | -0.089          | -0.302        |
| NN         | 0.018          | -0.083     | -0.004         | 0.351         | -0.04         | 0.028         | -0.564**         | 0.007        | 0.094           | 0.123         |
| AN         | 0.226          | -0.024     | -0.162         | -0.137        | 0.174         | 0.049         | 0.067            | 0.196        | 0.395*          | 0.073         |
| AP         | 0.445*         | -0.206     | 0.187          | 0.201         | -0.156        | -0.002        | -0.012           | 0.411*       | -0.123          | -0.492**      |
| TS         | 0.443*         | -0.169     | -0.238         | -0.206        | -0.09         | 0.027         | -0.394*          | 0.394*       | 0.031           | -0.119        |
| Altitude   | 0.169          | -0.093     | 0.175          | -0.186        | -0.274        | -0.076        | 0.478*           | 0.169        | -0.335          | -0.297        |

Legend: r-value represents spearman correlation coefficient, between -1 and 1, r<0 is negative correlation, r>0 is positive correlation *p < 0.05, **p < 0.01.
3 Discussion

One noteworthy result of our study was that the diversity and richness of the bacterial community in the rhizosphere of *Ferula sinkiangensis* were significantly different at different slope positions and soil depths (Table 1). This result supported our first hypothesis on the bacterial diversity and abundance of *Ferula sinkiangensis* being sensitive to slope and soil depth variation; importantly, the spatial heterogeneity of bacterial communities was detectable on a small scale; these findings were consistent with the results of Qi Sun et al. [58] in exploring the microbial diversity of steep slope soils in the semi-arid Loess Plateau and also Martina et al. [59] who explored the small-scale spatial diversity of temperate forests.

The variation in diversity and richness of bacterial communities along the slope might be attributed mainly to two aspects. 1. Redistribution of nutrients down the slope during rainfall erosion [60, 61], causing differences in soil properties (Supplementary Table S3), indirectly affecting the diversity and structure of bacterial communities. 2. The illumination time and intensity differ at various slope positions [62]. For example, there was a longer illumination time at the top of the slope than at the bottom, which was conducive to increased photosynthetic formation of organic compounds by plants, thus increasing the input of organic matter into soil and affecting the bacterial community.

Stratification differences in the diversity and richness of bacterial communities with soil depth were similar to those in the studies of Lopez-Mondejar et al. and Eilers et al. [63, 64] mainly due to spatial heterogeneity of nutrients. The topsoil has greater contents of nutrients than the deeper soil layers, with a consequent increase in microbial α-diversity [65]. Secondly, the topsoil microorganisms have higher activity than the deep soil microorganisms. Microbial decomposition and metabolic processes result in a large number of metabolites (including extracellular enzymes), contributing to topsoil nutrient accumulation and thus growth of bacteria.

Soil pH is considered as the best predictor of bacterial community structure and diversity at different spatial scales and soil types [66-68], and this was consistent with the results we found in this study regarding the composition and variation of the rhizosphere bacterial community in the rhizosphere of *Ferula sinkiangensis* (Figure 3 and Supplementary Table S4). However, the mechanism behind this model is little known, and we propose three relevant hypotheses. 1. Bacteria have a certain range of tolerance to pH, beyond which they cannot survive. The intracellular pH of many microorganisms is close to neutral [69]. Therefore, any degree of pH change exerts pressure on some bacterial groups with narrow pH tolerance and inadaptability, interfering with their growth, affecting their competitiveness [70], and eventually leading to changes in the composition of bacterial communities. 2. The pH may not directly affect the structure of bacterial communities, but it may affect the physicochemical properties of soils directly and indirectly. For example, there was a significant negative correlation between pH and salt content (Table 2); in addition,
pH affects the availability of nutrients and heavy metal ions in soil [71, 72]. Therefore, pH can be used as an integrated variable to combine with other factors in influencing the community structure of bacteria. Dakora et al. showed that some plants actively affect soil bacterial communities by changing soil pH through root exudates [22]. In addition, studies have shown that plant growth was passively influenced by pH, regulating the amount and composition of root secretions [72, 73], which further influences nutrient availability, enzyme activity and microbial abundance [74, 75]. Of course, pH, whether actively or passively, plays an important role in influencing the structure of bacterial communities.

Interestingly, we found in this study there was a significant correlation between AP (but not pH) and *Ferula sinkiangensis* rhizosphere bacterial diversity. This suggested that pH was not always a universal predictor of bacterial community diversity. Depending on habitat environment and soil type, there might be factors that explain bacterial diversity better than pH. The reason why AP has a significant relationship with bacterial community diversity was a relationship between C mineralization and P availability. Previous studies showed that in the absence of P, the retention time of litter was longer and the mineralization of unstable C was inhibited. This even had a negative effect on the carbon sequestration in cultivated lands [76, 77]. However, C is the main source of energy for microorganisms and interferes with their growth [78]. The response of C mineralization to P availability directly impacts the biodiversity.

Microorganisms can be closely involved in soil nutrient cycle, improved soil nutrient status and optimized soil structure [79-81]. Provide suitable soil environment for plant growth. Considering the important role of microorganisms in plant growth and development, Enkatachalam et al. put forward the "Belowground Solutions to an Aboveground Problem" viewpoint [31]. That is, by trying to solve some of the problems faced by plants on the ground by relying on microbes in the soil below. Meanwhile, the research of some scholars also confirmed this viewpoint. For example, Hoflich et al. stimulated plant growth by inoculation with symbiotic and associative rhizosphere microorganisms [82], and the microorganisms discovered by Horace et al. regulated plant growth by producing various secondary metabolites [83]. In addition, some scholars applied phosphate, potash and nitrogen fertilizers and organic matter to alter the microbial community diversity and influence plant growth [84, 85]. Therefore, we hope that the discovery of pH, AP, TP and TK significantly influencing the bacterial community diversity and structure in the rhizosphere of *Ferula sinkiangensis* (Table 3 and Supplementary Table S4) can be applied in conservation and commercial cultivation of *Ferula sinkiangensis*.

4 Conclusion

Although *Ferula sinkiangensis* population continues to decline, the research conducted on *Ferula sinkiangensis* is still focused mainly on medicinal properties; in contrast, the research on habitat ecology, plant physiology and soil is rare, restricting
the capacity to conserve *Ferula sinkiangensis*. In this study, we explored the
diversity and structural composition of the rhizosphere bacterial community in the
rhizosphere of *Ferula sinkiangensis*. However, bacterial communities are made up of
a wide variety of bacteria, some of which are beneficial to plant growth and
development, some of which are neutral, and some of which are not conducive to
plant growth and development. This requires us to conduct further characterization
in subsequent studies, to identify the bacterial populations conducive to the growth
and development of *Ferula sinkiangensis*. The spatial heterogeneity of
microorganisms has been shown, but there are few studies on sloping land and soil
depth. Our study revealed trends in bacterial community diversity at different slopes
and soil depths, which will contribute to our understanding of the
microbially-mediated carbon cycle in soil. Our study demonstrated the universality
of pH prediction of bacterial community structure, but also showed that pH effects
on bacterial community diversity might be limited under the influence of specific
habitats and soil types. In addition, the dominant effects of pH, AP, TK and TP on the
bacterial community were identified, and we considered using them in the
conservation and commercial culture of *Ferula sinkiangensis*. By adjusting pH value
and applying phosphate and potassium fertilizers, the diversity and structure of
bacterial community might be affected, and soil structure might be improved to
provide suitable soil environment for the growth of *Ferula sinkiangensis*. Exploring
the optimal amount and proportion of fertilizers needs to be done in the future
studies.

5 Materials and methods

5.1 Site description and sampling

Since *Ferula sinkiangensis* is a short-lived plant, we chose to study it during its
growing season (April-May). We set the place of Byshidun, Yili, Xinjiang, China as
the research plot (desert gray calcareous soil, slope: 43°, longitude: 82.083359,
latitude: 43.723121). On the slopes where *Ferula sinkiangensis* grew, we designed
rectangular plots of 10m×5m at the top, middle and bottom of the slope respectively.
Riley and Barber shaking methods [42, 43] were used to collect rhizosphere and
non-rhizosphere soil samples in 10m×5m plots circled at the top, middle and bottom
of the slope. Taking the slope top as an example, 3 *Ferula sinkiangensis* with the same
growth potential were randomly selected in the sample plot of 10m×5m. Then the
rhizosphere and non-rhizosphere soil samples were collected by shaking[42, 43] at
the roots of 0-10cm, 10cm-25cm and 25cm-40cm of each *Ferula sinkiangensis*. Collected soil was immediately placed in a 5 ml test tube with sterile tweezers, and
then marked on a tube and placed in a liquid nitrogen tank for storage. A total of 18
soil samples were collected at the top of the slope. Specifically, two types
(rhizosphere and non-rhizosphere) ×3 depths (0-10 cm, 10 cm-25 cm and 25 cm-40
cm) ×3 replicates. We explored three different plots of top, middle and bottom, with
a total of 54 soil samples (3×18). In addition, we used a mixed labeling system to label the samples. The first group of letters was used to represent the position of the slope, rhizosphere and non-rhizosphere (E, R and S represent the rhizosphere region of the top, middle and bottom of the slope respectively), (NE, NR and NS represent the non-rhizosphere region of the top, middle and bottom of the slope respectively). The second number indicates the depth (1, 2, and 3 represent 0-10cm, 10-25cm, and 25-40cm depth, respectively) and the third number indicates the number of repetitions. For example, S2.1 represents the first replicate soil sample with a root depth of 10-25cm in the roots of *Ferula sinkiangensis* at the bottom of the slope (Supplementary Table S5).

### 5.2 Soil analysis

Obtained soil samples were naturally air-dried, sieved to 2mm, and plant impurities were removed for determining physical and chemical properties. Organics matter contents were determined by external heating with potassium bichromate [44]. Nitrogen content was determined by the perchlorate-sulfuric acid digestion method, and the use of a fox 1035 automatic nitrogen determination apparatus [44]. Total phosphorus contents were assayed by the molybdenum method with an agilent Cary60 UV spectrophotometer r [44]. Total potassium was measured using the acid dissolution - atomic absorption method and a Thermo Scientific Series Atomic Absorption Spectrometer. pH measurements were performed with a mettler tolido FiveEasy Plus pH-meter. Available phosphorus was determined by molybdenum inverse colorimetry after extraction with sodium bicarbonate. Nitrate and ammonium nitrogen was determined by the 0.01M calcium chloride extraction method using a BRAN + LUEBBE flow analyzer. Total salt was determined by using dry-residue method (Supplementary Table S6).

### 5.3 DNA extraction, amplification and library generation

DNA was isolated from each sample using a Centrifugal Soil Genomic DNA Extraction Kit, and the purity and concentration of the DNA were measured by 1% agarose gel electrophoresis. An appropriate amount of DNA sample was placed in a sterile centrifuge tube and diluted to 1 ng/μL with sterile water. Partial 16S rDNA-based high-throughput sequencing was used to determine the bacterial diversity and community composition according to Caporaso et al [45, 46]. Using diluted DNA as a template, PCR amplification of the V4 region of 16S rRNA gene was performed using barcode-specific primers 806R (5’-GGACTACHVGGGTWTCTAAT-3’) and 515F (5’-GTGCCAGCMGCGGTAA-3’). The V4 region was used because 806R and 515F primers produced the greatest diversity at the bacterial level [47]. Phusion® High-fidelity PCR Master Mix with GC buffer from New England Biolabs was used to ensure the efficiency and accuracy of amplification. PCR products were detected by electrophoresis with 2% agarose gel, and target bands between 200-300bp were cut for further experiments. PCR products were mixed at equal density ratios. The
PCR products were purified using Qiagen Gel Extraction Kits (Qiagen, Germany). The recovered purified products were placed in a 1.5 mL sterile centrifugal tube in a dry ice box and sent to Beijing Compass Biotechnology Co., Ltd for high-throughput sequencing. The library was constructed using a TruSeq® DNA PCR-Free Sample Preparation Kit. The constructed library was quantified by Qubit and qT-PCR. After the library was qualified, it was sequenced using HiSeq2500 PE250.

### 5.4 Sequence analysis

After barcode sequences and primer sequences were cut off, the reads of each sample were spliced with a fast and accurate analysis tool FLASH [48] to obtain the raw tags. Raw tags were carefully filtered [49] to get clean tags of high quality using Qiime (V1.9.1) [50]. Tags quality control process comprised the following operations: a) Tags intercept: raw tags with low quality value (default quality threshold was ≤19) of the set length (default length value was 3) were truncated; b) Tags length filtering: the tagged data set obtained after interception of tags was further filtered out of the continuous high quality tags whose base length was less than 75% of the tags length. The tag sequences obtained after the above treatment are compared with the Gold database by UCHIME Algorithm [51] to detect chimeric sequences that were removed [52], to obtain the final effective tags.

Uparse software (Uparse v7.0.1001) [53] was used to cluster the effective tags of all samples, with 97% sequence identity. Grouping sequences into Operational Taxonomic Units (OTUs). The sequences with the highest occurrence frequency in OTUs were selected as representative of OTUs by using the Mothur method. SILVA [54] SSUrRNA database [55] was used for OTUs annotation to represent species sequence analysis (setting threshold of 0.8-1) and acquire information on taxonomy. The MUSCLE (Version 3.8.31) software [56] was used for fast multi-sequence alignment to obtain the system relationship of all OTUs representative sequences. Finally, the sample with the smallest amount of data was homogenized. Subsequent alpha diversity (α-diversity) and beta diversity (β-diversity) analyses were both based on the homogenized data.

### 5.5 Data analysis

QIIME software (Version 1.9.1) was used to calculate the Observed-species, Chao1, Shannon, and Goods-coverage indices. Chao1, Shannon and Goods-coverage indices can be used to evaluate the richness, diversity and sequencing depth of samples respectively [57]. In addition, R software (Version 2.15.3) was used to draw the dilution curve to reflect the rationality of sequencing data volume. β-diversity analysis was performed on weighted and unweighted unifac with the QIIME software (version 1.7.0). The latter software was used as a hierarchical clustering method to interpret the distance matrix using average link, and arithmetic mean (UPGMA) clustering was used for the unweighted pair method. The linear discriminant analysis (LDA) effect size (LEfSe) was used to identify statistically significant differences between groups.
One-way ANOVA (Tukey test) was used to assess differences in bacterial relative abundance or diversity between samples. Spearman correlation analysis was performed to assess the interrelationship between soil properties and the dominant bacterial species. In addition, distance-based redundancy analysis (db-RDA) and variance partitions were performed to study the relationship between the composition of the rhizosphere, soil properties and non-soil parameters. This was carried out with the Canoco statistical software (version 5.0) and default parameters (vegan package).

Availability of materials and data

We declare that all experimental data involved in the manuscript have been added to the manuscript and supplementary materials. If readers need further data related to the manuscript, please feel free to contact our corresponding author at any time.

References

1. Qian ZZ, Dan Y, Liu YZ, Peng Y: Pharmacopoeia of the People's Republic of China (2010 Edition): A Milestone in Development of China's Healthcare. Chinese Herbal Medicines 2010, 02(2):157-160.
2. Suzuki K, Okasaka M, Kashiwada Y, Takaishi Y, Honda G, Ito M, Takeda Y, Kodzhimatov OK, Ashurmetov O, Sekiya M: Sesquiterpene lactones from the roots of Ferula varia and their cytotoxic activity. Journal of Natural Products 2007, 70(12):1915-1918.
3. Mansour Z, Ben Jannet H, Sylvie C, Jalloul B: Chemical composition, biological and cytotoxic activities of plant extracts and compounds isolated from Ferula lutea. Molecules 2014, 19(3):2733-2747.
4. Kartal N, Sokmen M, Tepe B, Daferera D, Polissiou M, Sokmen A: Investigation of the antioxidant properties of Ferula orientalis L. using a suitable extraction procedure. Food Chemistry 2007, 99(2):584-589.
5. El-Razek MHA, Ohta S, Hirata T: Terpenoid Coumarins of the Genus Ferula. Cheminform 2003, 34(22).
6. Matejić JS, Džamić AM, Mihajilov-Krstev T, Randelović VN, Krivošej ZD, Marin PD: Total phenolic content, flavonoid concentration, antioxidant and antimicrobial activity of methanol extracts from three Seseli L. taxa. Central European Journal of Biology 2012, 7(6):1116-1122.
7. Motai T, Kitanaka S: Sesquiterpene chromones from Ferula fukanensis and their nitric oxide production inhibitory effects. Journal of Natural Products 2004, 67(3):432-436.
8. Lee CL, Chiang LC, Cheng LH, Liaw CC, Abd El-Razek MH, Chang FR, Wu YC: Influenza A (H1N1) Antiviral and Cytotoxic Agents from Ferula assa-foetida. Journal of Natural Products 2009, 72(9):1568-1572.
9. Bagheri SM, Mohammadsadeghi H, Dashtir MH, Mousavian SMM, Aghaei ZA: Effect of Ferula assa-foetida oleo-gum-resin on renal function in normal Wistar rats. Indian J Nephrol 2016, 26(6):419-422.

10. Xu E, Zhang H: Spatially-explicit sensitivity analysis for land suitability evaluation. Applied Geography 2013, 45(45):1-9.

11. Zhu J, Li XJ, Jia XG: A preliminary study on the ecological habit of the pest of Ferula sinkiangensis -- longicorn longicornis with white hair. In: Proceedings of the 9th national symposium on natural drug resources: 2010.

12. Tan GY, He S, Yao F: Male sterility and morphological observation of Ferula sinkiangensis. In: 70th anniversary annual meeting of Chinese botanical society: 2003.

13. Xie C, Shi M, Guo B, Shi L, Zeng F, Fu D, Li X, Jia X: Resource Investigation for Endangered Wild Ferula sinkiangensis Based on Low Altitude Remote Sensing. 2014.

14. Rodriguez H, Fraga R: Phosphate solubilizing bacteria and their role in plant growth promotion. Biotechnology Advances 1999, 17(4-5):319-339.

15. Bardgett RD, Freeman C, Ostle NJ: Microbial contributions to climate change through carbon cycle feedbacks. The ISME Journal 2008.

16. Schimel DS: Terrestrial ecosystems and the carbon cycle. Global Change Biology 2010, 1(1):77-91.

17. Thielebruhn S: Linking soil biodiversity and agricultural management. Current Opinion in Environmental Sustainability 2012, 4(5):523-528.

18. Bais HP, Weir TL, Perry LG, Gilroy S, Vivanco JM: THE ROLE OF ROOT EXUDATES IN RHIZOSPHERE INTERACTIONS WITH PLANTS AND OTHER ORGANISMS. Annual Review of Plant Biology 2006, 57(1):233-266.

19. Hiltner, L: Über neuere Erfahrungen und probleme auf dem gebiete der Bodenbakteriologie unter besonderer Berü cksichtigung der grü ndü ngung und Brache. Arbeit. Deut. Landw. Ges. Berlin 1904,98:59–78.

20. De Deyn GB, Cornelissen JH, Bardgett RD: Plant functional traits and soil carbon sequestration in contrasting biomes. Ecology Letters 2008, 11(5):516-531.

21. Bulgarelli D, Garrido-Oter R, Münch PC, Weiman A, Dröge J, Pan Y, Mchardy AC, Schulze-Lefert P: Structure and function of the bacterial root microbiota in wild and domesticated barley. Cell Host & Microbe 2015, 17(3):392-403.

22. Dakora FD, Phillips DA: Root exudates as mediators of mineral acquisition in low-nutrient environments. Plant & Soil 2002, 245(1):201-213.

23. Matiru VN, Dakora FD: Xylem transport and shoot accumulation of lumichrome, a newly recognized rhizobial signal, alters root respiration, stomatal conductance, leaf transpiration and photosynthetic rates in legumes and cereals. New Phytologist 2010, 165(3):847-855.

24. Gull M, Hafeez FY, Saleem M, Malik KA: Phosphorus uptake and growth promotion of chickpea by co-inoculation of mineral phosphate solubilising bacteria and a mixed rhizobial culture. Australian Journal of Experimental Agriculture 2004, 44(6):142-153.

25. Rodriguez RJ, Henson J, Van VE, Hoy M, Wright L, Beckwith F, Kim YO, Redman RS: Stress tolerance in plants via habitat-adapted symbiosis. Isme Journal 2008, 2(4):404.

26. Liu L, Liu S, Chen F, Yang X, Yang C, Bingqi WU, Zhang M, Zhao J: Effect of endophytic bacteria inoculation on cadmium uptake in Solanum nigrum L. Acta Scientiae Circumstantiae 2013, 33(12):3368-3375.
27. Mayak S, Tirosh T, Glick BR: Plant growth-promoting bacteria confer resistance in tomato plants to salt stress. *Plant Physiology & Biochemistry* 2004, 42(6):565-572.

28. Shimon, TIROSH, Tsipora, GLICK, Bernard R: Plant growth-promoting bacteria that confer resistance to water stress in tomatoes and peppers. *Plant Science* 2004, 166(2):525-530.

29. Rolli E, Marasco R, Vigani G, Ettoumi B, Mapelli F, Deangelis ML, Gandolfi C, Casati E, Previtali F, Gerbino R: Improved plant resistance to drought is promoted by the root-associated microbiome as a water stress-dependent trait. *Environmental Microbiology* 2014, 17(2):316-331.

30. Berendsen RL, Pieterse CMJ, Bakker PAHM: The rhizosphere microbiome and plant health. *Trends in Plant Science* 2012, 17(8):478-486.

31. Lakshmanan V, Selvaraj G, Bais HP: Functional Soil Microbiome: Belowground Solutions to an Aboveground Problem. *Plant Physiology* 2014, 166(2):689-700.

32. Howard AG, Comber SDW, Kifle D, Antai EE, Purdie DA: Arsenic Speciation and Seasonal Changes in Nutrient Availability and Micro-plankton Abundance in Southampton Water, U.K. *Estuarine Coastal & Shelf Science* 1995, 40(4):435-450.

33. Nessner KV, Taketani RG, Lançoni MD, Andreote FD, Mendes R, Soares dMI: Water regime influences bulk soil and Rhizosphere of Cereus jamacaru bacterial communities in the Brazilian Caatinga biome. *Plos One* 2013, 8(9):e73606.

34. Schnittler M, Stephenson SL: Myxomycete biodiversity in four different forest types in Costa Rica. *Myologia* 2000, 92(4):626-637.

35. Nascimento AL, Souza AID, Andrade PAMD, Diniandreote F, Gomes ARC, Oliveira FC, Regitano JB: Sewage Sludge Microbial Structures and Relations to Their Sources, Treatments, and Chemical Attributes. *Frontiers in Microbiology* 2018, 9(1462):1462-.

36. Treseder KK: Nitrogen additions and microbial biomass: a meta-analysis of ecosystem studies. *Ecology Letters* 2010, 11(10):1111-1120.

37. Yao Q, Liu J, Yu Z, Li Y, Jin J, Liu X, Wang G: Three years of biochar amendment alters soil physiochemical properties and fungal community composition in a black soil of northeast China. *Soil Biology & Biochemistry* 2017, 110:56-67.

38. Jing Z, Xin J, Zhou B, Zhao B, Ma M, Guan D, Li J, Chen S, Cao F, Shen D: Thirty four years of nitrogen fertilization decreases fungal diversity and alters fungal community composition in black soil in northeast China. *Soil Biology & Biochemistry* 2016, 95:135-143.

39. Franklin RB, Mills AL: Multi-scale variation in spatial heterogeneity for microbial community structure in an eastern Virginia agricultural field. *Fems Microbiology Ecology* 2010, 44(3):335-346.

40. Porter SS, Rice KJ: Trade-offs, spatial heterogeneity, and the maintenance of microbial diversity. *Evolution;international journal of organic evolution* 2013, 67(2):599-608.

41. Choi JH, Kim GB, Cha CJ: Spatial heterogeneity and stability of bacterial community in the gastrointestinal tracts of broiler chickens. *Poultry Science* 2014, 93(8):1942-1950.

42. Riley D, Barber SA: Bicarbonate Accumulation and pH Changes at the Soybean (Glycine max (L.) Merr.) Root-Soil Interface1. *Soil Science Society of America Journal* 1969, 33(6):905-908.
43. Riley D, Barber SA: Salt accumulation at the soybean (Glycine max (L.) Merr) root-soil interface. *Soil Science Society of America Journal* 1970, 34(1):154-155.

44. Bao, S. D. Soil Agro-chemical Analysis. 22–196 (China Agriculture Press, 2008).

45. Caporaso JG, Lauber CL, Walters WA, Berglyons D, Lozupone CA, Turnbaugh PJ, Fierer N, Knight R: Global patterns of 16S rRNA diversity at a depth of millions of sequences per sample. *Proc Natl Acad Sci U S A* 2011, 108(Supplement_1):4516-4522.

46. Caporaso JG, Lauber CL, Walters WA, Berglyons D, Huntley J, Fierer N, Owens SM, Betley J, Fraser L, Bauer M: Ultra-high-throughput microbial community analysis on the Illumina HiSeq and MiSeq platforms. *Isme Journal Multidisciplinary Journal of Microbial Ecology* 2012, 6(8):1621-1624.

47. Peiffer JA, Spor A, Koren O, Jin Z, Tringe SG, Dangl JL, Buckler ES, Ley RE: Diversity and heritability of the maize rhizosphere microbiome under field conditions. *Proceedings of the National Academy of Sciences of the United States of America* 2013, 110(16):6548-6553.

48. Magoč T, Salzberg SL: FLASH: fast length adjustment of short reads to improve genome assemblies. *Bioinformatics* 2011, 27(21):2957-2963.

49. Bokulich NA, Subramanian S, Faith JJ, Gevers D, Gordon JL, Knight R, Mills DA, Caporaso JG: Quality-filtering vastly improves diversity estimates from Illumina amplicon sequencing. *Nature Methods* 2013, 10(1):57-59.

50. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, Fierer N, Peña± A AG, Goodrich JK, Gordon JI: QIIME allows analysis of high-throughput community sequencing data. *Nature Methods* 2010, 7(5):335-336.

51. Edgar RC, Haas BJ, Clemente JC, Quince C, Knight R: UCHIME improves sensitivity and speed of chimera detection. *Bioinformatics* 2011, 27(16):2194.

52. Haas BJ, Gevers D, Earl AM, Feldgarden M, Ward DV, Giannoukos G, Ciulla D, Tabbaa D, Highlander SK, Sodergren E: Chimeric 16S rRNA sequence formation and detection in Sanger and 454-pyrosequenced PCR amplicons. *Genome research* 2011, 21(3):494-504.

53. Edgar RC: UPARSE: highly accurate OTU sequences from microbial amplicon reads. *Nature Methods* 2013, 10(10):996.

54. Christian Q, Elmar P, Pelin Y, Jan G, Timmy S, Pablo Y, J?Rg GC: The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. *Nucleic Acids Research* 2013, 41(Database issue):590-596.

55. Wang Q, Garrity GM, Tiedje JM, Cole JR: Naive Bayesian classifier for rapid assignment of rRNA sequences into the new bacterial taxonomy. *Applied & Environmental Microbiology* 2007, 73(16):5261.

56. Edgar RC: Local homology recognition and distance measures in linear time using compressed amino acid alphabets. *Nucleic Acids Research* 2004, 32(1):380-385.

57. Li B, Zhang X, Guo F, Wu W, Zhang T: Characterization of tetracycline resistant bacterial community in saline activated sludge using batch stress incubation with high-throughput sequencing analysis. *Water Research* 2013, 47(13):4207-4216.

58. Guo S: Spatial variations of soil respiration and temperature sensitivity along a steep slope of the semiarid Loess Plateau. *Plos One* 2018, 13(4):e0195400.
59. M Š, Bárt a J, H Š, Baldrian P: Small-scale spatial heterogeneity of ecosystem properties, microbial community composition and microbial activities in a temperate mountain forest soil. *Fems Microbiology Ecology* 2016, 92(12):fw185.

60. Lal R: Soil erosion and carbon dynamics. *Soil & Tillage Research* 2005, 81(2):137-142.

61. Cheng Y, Shao-Shan AN, Yun-Fei MA: Soil Microbial Biomass and Enzymatic Activities in the Loess Hilly Area of Ningxia under Different Slope Positions. *Research of Soil & Water Conservation* 2010.

62. Guo C, Jinchuan LI, Yue J, Yang S, Ning LU: Diurnal changes in the photosynthetic characteristics of two high yield and high quality grasses during different stages of growth and their response to changes in light intensity. *Acta Ecologica Sinica* 2013, 33(6):1751-1761.

63. Fierer N, Schimel JP, Holden PA: Variations in microbial community composition through two soil depth profiles. *Soil Biology & Biochemistry* 2003, 35(1):167-176.

64. Eilers KG, Debenport S, Anderson S, Fierer N: Digging deeper to find unique microbial communities: The strong effect of depth on the structure of bacterial and archaeal communities in soil. *Soil Biology & Biochemistry* 2012, 50(7):58-65.

65. Voříšková J, Brabcová V, C ajthaml T, Baldrian P: Seasonal dynamics of fungal communities in a temperate oak forest soil. *New Phytologist* 2013, 201(1):269-278.

66. Rousk J, Båå th E, Brookes PC, Lauber CL, Lozupone C, Caporaso JG, Knight R, Fierer N: Soil bacterial and fungal communities across a pH gradient in an arable soil. *Isme Journal* 2010, 4(10):1340-1351.

67. Bartram AK, Xingpeng J, Lynch MDJ, Masella AP, Nicol GW, Jonathan D, Neufeld JD: Exploring links between pH and bacterial community composition in soils from the Craibstone Experimental Farm. *Fems Microbiology Ecology* 2013, 87(2):403-415.

68. Fierer N: Embracing the unknown: disentangling the complexities of the soil microbiome. *Nature Reviews Microbiology* 2017, 15(10):579-590.

69. Abed RMM, Kohls K, Palinska KA, Golubic S: Diversity and Role of Cyanobacteria and Aerobic Heterotrophic Microorganisms in Carbon Cycling in Arid Cyanobacterial Mats; 2010.

70. Fierer N, Jackson RB: The diversity and biogeography of soil bacterial communities. *Proceedings of the National Academy of Sciences of the United States of America* 2006, 103(3):626-631.

71. Harter, nbsp, Robert D: Effect of Soil pH on Adsorption of Lead, Copper, Zinc, and Nickel. *Soil Scisocamj* 1983, 47(1):47-51.

72. Weil R, Brady N: The Nature and Properties of Soils, Global Edition; 1974.

73. Ibekwe AM, Poss JA, Grattan SR, Grieve CM, Suarez D: Bacterial diversity in cucumber ( Cucumis sativus ) rhizosphere in response to salinity, soil pH, and boron. *Soil Biology & Biochemistry* 2010, 42(4):567-575.

74. Hong CQ: Effects of root exudates on plant nutrition. *Ecological environment* 2003, 12:508-511.

75. Zhao XL, Liu XH, Jiang-Zhou HE, Wan CX, Gong MF, Zhang LL: Effects of Cotton Root Exudates on Available Soil Nutrition,Enzyme Activity and Microorganism Quantity. *Acta Botanica Boreali-Occidentalia Sinica* 2009:1426-1431.
76. Cleveland CC, Townsend AR, Schmidt SK: Phosphorus limitation of microbial processes in moist tropical forests: Evidence from short-term laboratory incubations and field studies. Ecosystems 2002, 5(7):680-691.

77. Cruz AF, Hamel C, Hanson K, Selles F, Zentner RP: Thirty-seven years of soil nitrogen and phosphorus fertility management shapes the structure and function of the soil microbial community in a Brown Chernozem. Plant & Soil 2009, 315(2):173-184.

78. Eilers KG, Lauber CL, Knight R, Fierer N: Shifts in bacterial community structure associated with inputs of low molecular weight carbon compounds to soil. Soil Biology & Biochemistry 2010, 42(6):896-903.

79. Henis Y: Soil microorganisms, soil organic matter and soil fertility; 1986.

80. Sparling GP, Pankhurst C, Doube BM, Gupta VVSR: Soil microbial biomass, activity and nutrient cycling as indicators of soil health; 1997.

81. Webster EA, Hopkins DW, Chudek JA, Haslam SF, Simek M, Pícek T: The relationship between microbial carbon and the resource quality of soil carbon. Journal of Environmental Quality 2001, 30(1):147.

82. Castrosoiwinski S, Herschkovitz Y, Okon Y, Jurkevitch E: Effects of inoculation with plant growth-promoting rhizobacteria on resident rhizosphere microorganisms. Fems Microbiology Letters 2010, 276(1):1-11.

83. Cutler HG, Wells JM: Unusual plant growth regulators from microorganisms. Critical Reviews in Plant Sciences, 6(4):323-343.

84. Bourke RM: Influence of nitrogen and potassium fertilizer on growth of sweet potato (Ipomoea batatas) in Papua New Guinea. Field Crops Research 1985, 12(85):363-375.

85. Sarathchandra SU, Lee A, Perrott KW, Rajan S, Oliver E, Gravett IM: Effects of phosphate fertilizer applications on microorganisms in pastoral soil. Australian Journal of Soil Research 1993, 31(3):641-648.

Acknowledgements

In this study, we would like to thank professor LZ for his guidance, all the authors for their joint efforts, and Ferula sinkiangensis research enthusiasts who provided guidance services in this experiment.

Author Contributions

T.Z.: Methodology, Data curation, Investigation, Writing-Original draft preparation.
Z.W.: Validation, Investigation. X.Lv.: Validation, Investigation. H.D.: Supervision, Methodology, Validation, Investigation. L.Z.: Supervision, Project administration, Conceptualization, Writing-Reviewing and Editing. All authors have read and agreed to the published version of the manuscript.
Conflicts of Interest

We declare that the research was conducted in the absence of any commercial or financial relationships.