Soil pH: a key edaphic factor regulating distribution and functions of bacterial community along vertical soil profiles in red soil of pomelo orchard

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Abstract

Background: Soil microbes exist throughout the soil profile and those inhabiting topsoil (0–20 cm) are believed to play a key role in nutrients cycling. However, the majority of the soil microbiology studies have exclusively focused on the distribution of soil microbial communities in the topsoil, and it remains poorly understood through the subsurface soil profile (i.e., 20–40 and 40–60 cm). Here, we examined how the bacterial community composition and functional diversity changes under intensive fertilization across vertical soil profiles [(0–20 cm (RS1), 20–40 cm (RS2), and 40–60 cm (RS3)] in the red soil of pomelo orchard, Pinghe County, Fujian, China.

Results: Bacterial community composition was determined by 16S rRNA gene sequencing and interlinked with edaphic factors, including soil pH, available phosphorous (AP), available nitrogen (AN), and available potassium (AK) to investigate the key edaphic factors that shape the soil bacterial community along with different soil profiles. The most dominant bacterial taxa were Proteobacteria, Acidobacteria, Actinobacteria, Chloroflexi, Crenarchaeota, and Bacteriodetes. Bacterial richness and diversity was highest in RS1 and declined with increasing soil depth. The distinct distribution patterns of the bacterial community were found across the different soil profiles. Besides, soil pH exhibited a strong influence (pH > AP > AN) on the bacterial communities under all soil depths. The relative abundance of Proteobacteria, Actinobacteria, Crenarchaeota, and Firmicutes was negatively correlated with soil pH, while Acidobacteria, Chloroflexi, Bacteriodetes, Planctomycetes, and Gemmatimonadetes were positively correlated with soil pH. Co-occurrence network analysis revealed that network topological features were weakened with increasing soil depth, indicating a more stable bacterial community in the RS1. Bacterial functions were estimated using FAPROTAX and the relative abundance of functional bacterial community related to metabolic processes, including C-cycle, N-cycle, and energy production was significantly higher in RS1 compared to RS2 and RS3, and soil pH had a significant effect on these functional microbes.

Conclusions: This study provided the valuable findings regarding the structure and functions of bacterial communities in red soil of pomelo orchards, and highlighted the importance of soil depth and pH in shaping the soil bacterial community.

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Background

Soil is one of the most diverse ecosystems consisting of microorganisms, organic matter, minerals, and water [1]. Soil microbes, including bacteria, fungi, macrofauna, mesofauna, and microfauna (e.g., protozoans and nematodes), are considered as the major soil biota [2]. Among these, bacteria have the most abundant community structure in the soil [3]. The soil microbial diversity is a distinguishing attribute of the agricultural systems and a variety of biogeochemical cycles, e.g., organic matter decomposition and soil nutrients cycling [4–6]. Therefore, it is necessary to understand the community composition and its response to environmental changes that could significantly affect the ecological functions.

Despite the widespread occurrence of soil microbiota in different soil depths, the current knowledge about the distribution of soil microbes is limited to surface soil (0—20 cm) [7] owing to higher contents of soil organic matter, mineral nutrients, and rich in microbial diversity compared with subsurface soil [8]. Various studies have investigated the microbial diversity and community composition in the surface and subsurface soils, e.g., in paddy soils or Alaskan soil cores [9–11]. The subsurface soil microbial community has great significance owing to its role in the soil formation process relative to the surface soil profile [12]. The soil microbial communities in subsurface soil also play a crucial role in soil carbon sequestration due to the storage of organic carbon in the subsurface soil profiles [13]. Thus, characterization of the soil microbial communities along various soil gradients would enable us to better understand the key characteristic of soil microbial communities and their potential functions in the red soil.

Red soil is mainly distributed in the southern regions of China (2.18 × 10^5 ha), covering 22% of the total land area [14]. These soils are characterized by low pH, low nutrient contents, and high aluminum ion activity that results in poor soil physical properties that are key limiting factors for plant growth [15], especially for pomelo orchards in the red soil of Pinghe County (Fujian Province), which is the most famous area for pomelo production [16, 17]. The cultivated area and yield of pomelo have been increased in recent decades, and total production of pomelo reached over 5 × 10^4 ha with an annual production of over 130 × 10^4 t (Pinghe County Statistical Yearbook, 2017). Recently, excessive use of fertilizers for higher yield has resulted in serious environmental problems such as soil acidity [18], soil contamination [19], greenhouse gases emission [20], water contamination [21], and detrimentally influenced the soil microbial diversity [22]. Evidence suggests that the soil microbial community has a significant role in the maintenance of soil structure and nutrients cycling, hence soil microbial diversity is an important index to assess soil health [23]. The diversity and community composition of soil microbes are significantly influenced by soil type, fertilizer type, application methods, and other various factors [23, 24]. For example, the application of inorganic fertilizers, especially N-fertilizer plays a significant role in improving crop productivity, however, its long-term application deteriorates crop productivity and soil quality [25, 26]. Nevertheless, the contradictions and uncertainty persist regarding the substantial impacts of inorganic fertilizers on the functional diversity and composition of soil microbes. Besides, inorganic fertilizers have a significant effect in improving biomass carbon and nitrogen [27], but decreases the functional diversity of soil microbes under long-term N application [28]. Some studies have found no effect of N application on community composition and functional diversity of soil microbes [24, 29]. In contrast, the inorganic fertilizers have shown the direct effects on functional diversity as well as community composition of soil microbiota [30]. However, in pomelo orchards, the effects of intense inorganic fertilization on soil microbial richness and community structure have not been fully elucidated.

Although several previous findings have validated the potential role of soil microbes in soil functioning and ecosystem services under various ecological conditions [31–33], however, very limited information is available about the important edaphic factors of soil bacterial community structure and functional diversity in the red soils. Hence, we investigated the bacterial diversity and community structure by 16S rRNA sequencing and conducted detailed analyses for microbial community structure through co-occurrence networks and prediction of functional studies. The key research objectives were as follows: (1) to explore the distribution of bacterial diversity and community composition along with the vertical soil profile under intensive use of fertilizers in red soil; (2) to investigate the important edaphic factors that shape the soil bacterial community; (3) to predict the functional

Keywords: Red soil, Bacterial diversity, Functional analysis, Soil pH, Spatial distribution
potential of bacterial community and their relationship with the edaphic factors.

Results
Changes in soil physicochemical properties under different soil depths
Soil physicochemical properties were significantly affected along soil profile gradient. Soil pH was significantly higher in topsoil RS1 (4.47) compared with RS2 (3.99) and RS3 (3.96) (Fig. 1A). Similarly, the available nutrients contents including AN (Fig. 1B), AP (Fig. 1C), and AK (Fig. 1D) were also significantly higher in RS1 compared with RS2 and RS3. Overall, we found that with decreasing the soil pH, the availability of the nutrients also decreased.

Effect of soil depth on specific bacterial community
In total, 6778 OTUs were extracted from 57 soil samples and categorized based on phylum (34), class (109), order (214), family (345), and genus (598). The dominant phyla included the Proteobacteria, Acidobacteria, Actinobacteria, Chloroflexi, Crenarchaeota, and Bacteroidetes, which accounted for 92% of all microbial communities (Fig. 2A). The relative abundance (RA) of Proteobacteria, Acidobacteria, and Bacteroidetes decreased with increasing soil depth, while RA of Actinobacteria and Crenarchaeota increased along with the soil depth, maximum at 40–60 cm (RS3) depth soil layer (Supplementary Fig. S1). Similarly, the differences in bacterial communities were observed at the genus level (Fig. 2B). Overall, these findings show that changes at the phylum level for specific

![Fig. 1 Changes in soil physicochemical properties. The variation in soil physicochemical properties were observed for A Soil pH; B Available nitrogen (AN); C Available phosphorous (AP); D Available potassium (AK) along vertical soil depths. The different lowercase letters indicate the significant differences (p < 0.05) among different soil depths, i.e., RS1 (0–20 cm), RS2 (20–40 cm), RS3 (40–60 cm)]
bacterial are contributed by the soil depth. According to the Venn diagram, 4232 OTUs were found to be prevalent across the three soil depths (Fig. 2C) and accounted for 93.8% of the total number of reads. It implies that the similarity of bacterial community composition was high in all depths of the red soil.

Richness and diversity of bacterial communities decreased along the soil profile gradient

We assessed the four α-diversity indices, including the observed number of OTUs, Shannon index, Chao1 index, and ACE index. The species richness indices, i.e., the observed number of OTUs, Chao1 and ACE were significantly higher in the RS1 (0–20 cm) than RS2 (20–40 cm), and RS3 (40–60 cm) soil depth. Similarly, Shannon’s was consistent and found the higher diversity in RS1, and it was significantly different from the RS3 (Fig. 3). It implies that bacterial richness and diversity were high in the topsoil and declined with increasing soil depth.

Changes in bacterial communities across the soil depth

To assess whether the soil depth influenced the microbial community composition across the different soil depths, the principal coordinates analysis (PCoA) was performed. The results showed that bacterial communities were separated by the different soil depths, where PCo1 accounted for 14.4% and PCo2 9.4% of the variation (Fig. 4). PERMANOVA analysis of the complete dataset suggested the significant differences in bacterial communities of different soil depths (Table 1). Furthermore, we
used PERMANOVA analysis to analyze the differences between the different soil depths for each paired group and discovered that bacterial populations were segregated by soil depth (Table 2). Soil bacterial communities in the RS1 were significantly different than RS2 and RS3, while no significant difference was found between the RS2 and RS3 bacterial communities (Table 2). These findings showed that soil depth has a significant effect on bacterial community composition.

**Soil properties correlated with bacterial community**

The distance-based redundancy analysis (RDA) exhibited that all the soil properties except AK, were significantly positively correlated in shaping the bacterial community’s structure, and the soil pH, AP, and AN had significant effects in changing the bacterial community’s structure (Fig. 5A). Among these edaphic factors, we found that bacterial populations were most affected by soil pH in the red soils (Table 3). Furthermore, we investigated these effects in each soil depth and found that soil pH was the most influential factor affecting the soil bacterial communities in each soil depth, including RS1 (Fig. 5B), RS2 (Fig. 5C), and RS3 (Fig. 5D), Table 3. Correlation analysis was further performed to explore the effect of soil pH on each bacterial community. Overall, we found that the relative abundance of Proteobacteria, Actinobacteria, Crenarchaeota, and Firmicutes was negatively affected by soil pH. In contrast, soil pH had a positive effect on the relative abundance of Acidobacteria, Chloroflexi, Bacteriodetes, Planctomycetes, and Gemmatimonadetes. Hence, soil pH plays a significant role in shaping the soil bacterial communities (Fig. 6).

**Co-occurrence network analysis**

The taxonomic properties of the bacterial populations in the red soil were studied using a co-occurrence network analysis. In all soil depths, the network analysis revealed a significant correlation between the bacterial communities. The high-abundance nodes in the co-occurrence network were separated into six phyla. Among them, Proteobacteria, Acidobacteria, Actinobacteria, Chloroflexi, Gemmatimonadetes, and Bacteriodetes represented the most dominant bacterial community at phylum level in RS1. Meanwhile, topological properties were calculated to explore the complex interrelationship patterns among the nodes [34]. RS1 had the highest number of nodes and edges (671 and 1813, respectively, Fig. 7A) compared with RS2 (449

Fig. 3 The α-diversity indices in different soil depths. The α-diversity indices were calculated; A Observed OTUs, B Shannon index, C Chao1, D ACE index. The differences between any two soil depths were tested by Wilcon test. The star represents the significance (*, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001, while ns, non-significant. RS1, RS3, and RS3 represents the soil depths of 0–20, 20–40, and 40–60 cm, respectively
and 600, respectively, Fig. 7B) and RS3 (478 and 805, respectively, Fig. 7C). It revealed that the soil bacterial network was more complex and bacterial associations were tight in RS1, while in RS2 and RS3 bacterial networks were less complex and bacterial associations were less tight.

**Table 1** Permutational analysis of variance (PERMANOVA) for all groups

|               | Df | Sum of sqs | Mean Sq | F. Model | R²   | Pr(> F) |
|---------------|----|------------|---------|----------|------|---------|
| Group         | 2  | 1.249      | 0.624   | 2.717    | 0.092| 0.001   |
| Residuals     | 54 | 12.418     | 0.229   | 0.908    |      |         |
| Total         | 56 | 13.668     | 1       |          |      |         |

Significant P-value (≤ 0.05) is bolded

**Table 2** Significance test of differences among bacterial communities in different soil depths using permutational analysis of variance based on Bray–Curtis distance

| Groups        | Measure | Permutations | R²      | P. value | Significance |
|---------------|---------|--------------|---------|----------|--------------|
| RS1 vs RS2    | bray    | 999          | 0.064   | 0.002    | ***          |
| RS1 vs RS3    | bray    | 999          | 0.0956  | 0.001    | **           |
| RS2 vs RS3    | bray    | 999          | 0.0428  | 0.066    |              |

The star represents the significance (**, P ≤ 0.01; ***, P ≤ 0.001)
Functional analysis

The expected functions of soil bacterial communities in pomelo orchards were determined using FAPROTAX analysis and predicted major functions were majorly attributed by C-cycle, N-cycle, and energy and their relative abundance were higher under RS1 compared with RS2 and RS3 (Supplementary Fig. S2). When the minor functions were predicted with the soil factors, we found that the metabolic functions of the C-cycle (e.g., photoautotrophy, photoheterotrophy, etc.), N-cycle (denitrification, nitrate and nitrite denitrification, etc.) were significantly affected by the soil pH (Fig. 8).

Discussion

The bacterial community is most dominated in the soil habitat owing to their diversity and functioning in different soil biochemical processes [35]. Their community composition, spatial patterns, and functional profiling have yet to be well investigated in the red soil of pomelo orchards. Recent researches of soil microbial biodiversity and long-term fertilizer studies have revealed that soil bacterial diversity is significantly affected by the soil depth and edaphic factors that shape the soil bacterial community [7]. For example, soil pH is a primary determinant of microbial community composition and
functional diversity [31, 36]. Thus, in the current study, we examined the bacterial community composition and functional diversity along with different soil profiles and identified the key edaphic factors regulating the soil bacterial diversity.

We found that different soil profiles representing variation in edaphic factors, e.g., soil pH, AN, AP, and AK were significantly higher in RS1 compared with RS2 and RS3 (Fig. 1), and had significant effects on bacterial community structure (Fig. 2). These findings are in concordance with the previous studies that soil depth plays an important role to shape the soil bacterial communities owing to differences in soil properties [7, 37]. Although soil properties (e.g., pH, AP, AK, AN) along different soil depths varied from each other but resulted in a high percentage of the shared OTUs accounted for 93.8% of total reads and it implies that similarity of bacterial community composition was high along the soil gradient. On the contrary, some taxa were still significantly different from each other in different soil profiles. For example, Proteobacteria and Bacteriodetes showed higher relative abundance in RS1 (Supplementary Fig. S1), and these are often categorized as copiotrophic group [38] and shows the higher growth rate with abundant resources [39], while Acidobacteria and Chloroflexi belonging to the oligotrophic group are highly abundant under low nutrients environment [40]. Hence, this is the reason that we found the higher relative abundance of Proteobacteria and Bacteriodetes in RS1 and Acidobacteria and Chloroflexi in RS3. So, these changes in bacterial communities along the soil depth could be attributed to differences in the soil properties [41]. The previous findings also showed the change in the relative abundance of bacterial communities along with the soil depth [42], although the changes in abundance patterns of bacterial communities were highly inconsistent. For instance, the relative abundance of Acidobacteria and Actinobacteria varied throughout the soil depth in one previous study [43]. However, we found that the RA of Acidobacteria was decreased with increasing soil depth, whereas Actinobacteria RA was at the peak in RS3 (40–60 cm), and consistent with the previous results where similar trends were observed in agricultural soils of Iowa, USA [7].

The differences in bacterial diversity and community composition were also revealed by α-diversity indices. The α-diversity of RS1 was significantly different from RS2 and RS3, but we found that RS2 and RS3 showed no difference in bacterial community structures although the differences in edaphic factors were high including pH, AP, and AK. However, the average soil pH was about the same in RS2 and RS3, i.e., 3.99 and 3.96, and as a result, we couldn’t find differences in RS2 and RS3. Overall, we found that the species richness and alpha diversity was decreased along the soil depth gradient and consistent with the previous findings of Hao et al. (2020) who also found the decrease in bacterial diversity along with soil profile [7]. Furthermore, redundancy analysis revealed that soil pH had a substantial impact on the structure of bacterial communities (Fig. 5, Table 3), and the importance of soil pH in shaping the bacterial populations along with vertical soil profile has long been recognized [44].

### Table 3
Pearson correlation between the Bray–Curtis dissimilarity score and soil characteristics using the mantel test

| Variable name | Corr-method | Corr_res | p_res | significance |
|---------------|-------------|----------|-------|--------------|
| Overall       | pH          | Pearson  | 0.366 | 0.001        | ***          |
|               | AP (Available Phosphorous) | Pearson  | 0.217 | 0.002        | **           |
|               | AK (Available Potassium)    | Pearson  | -0.007| 0.466        |              |
|               | AN (Available Nitrogen)     | Pearson  | 0.127 | 0.037        | *            |
| RS1           | pH          | Pearson  | 0.403 | 0.008        | **           |
|               | AP (Available Phosphorous)  | Pearson  | 0.113 | 0.132        |              |
|               | AK (Available Potassium)    | Pearson  | -0.182| 0.993        |              |
|               | AN (Available Nitrogen)     | Pearson  | -0.047| 0.639        |              |
| RS2           | pH          | Pearson  | 0.288 | 0.008        | **           |
|               | AP (Available Phosphorous)  | Pearson  | 0.322 | 0.005        | **           |
|               | AK (Available Potassium)    | Pearson  | -0.226| 0.963        |              |
|               | AN (Available Nitrogen)     | Pearson  | 0.224 | 0.087        |              |
| RS3           | pH          | Pearson  | 0.192 | 0.008        | **           |
|               | AP (Available Phosphorous)  | Pearson  | 0.247 | 0.005        | **           |
|               | AK (Available Potassium)    | Pearson  | -0.163| 0.963        |              |
|               | AN (Available Nitrogen)     | Pearson  | -0.119| 0.087        |              |

The star represents the represented the significance (*, $P \leq 0.05$; **, $P \leq 0.01$; ***, $P \leq 0.001$)
relationship has also been cited in previous studies [45, 46]. Schreiter et al. (2014) explained that besides some available nutrients, soil pH also drives the soil bacterial communities [45]. Sun et al. (2015) has demonstrated that soil pH is the key factor in shaping the soil bacterial communities [46]. The correlation analysis also showed that soil pH was a key factor for determining the soil microbial community composition, and our results agree with numerous previous studies [47, 48]. The close relationship of soil pH with the bacterial community is owing to the dependency of the majority of the bacterial community under narrow pH [49], and the slight change in pH significantly affects the bacterial community [50]. Many studies have reported the response of specific bacterial taxa to soil pH [51]. For instance, a negative correlation is found between the relative abundance of Proteobacteria and soil pH [52], a positive correlation between Acidobacteria and soil pH [53], we also found similar results in the present study. Furthermore, Gemmatimonadetes also showed

Fig. 6 Correlation analysis between the soil pH and bacterial communities. The effect of soil pH was tested on abundant taxa of bacterial community at phylum level

Fig. 7 Co-occurrence network of bacterial communities in red soil based on correlation analysis. The Co-occurrence network of bacterial communities was investigated in A 0–20 cm (RS1); B 20–40 cm (RS2); C 40–60 cm (RS3). The nodes in the network are colored based on phylum. Size of each node is proportional to the relative abundance of specific taxa. The connections showing the strong (spearman’s ρ ≥ 0.6) and significant (P ≤ 0.05) correlations
Fig. 7 (See legend on previous page.)

(A) RS1
- Proteobacteria (38.6%)
- Acidobacteria (20.1%)
- Actinobacteria (9.8%)
- Chloroflexi (8.35%)
- Gemmatimonadetes (5.0%)
- Bacteroidetes (4.9%)

Network Topological Features
- Nodes = 671
- Edges = 1813
- Avg. Degree = 5.40

(B) RS2
- Proteobacteria (31.1%)
- Acidobacteria (17.3%)
- Chloroflexi (16.7%)
- Actinobacteria (12.2%)
- Crenarchaeota (6.6%)
- AD3 (5.1%)

Network Topological Features
- Nodes = 449
- Edges = 600
- Avg. Degree = 2.67

(C) RS3
- Proteobacteria (28.4%)
- Chloroflexi (21.3%)
- Actinobacteria (15.6%)
- Acidobacteria (14.0%)
- AD3 (6.2%)
- Crenarchaeota (5.0%)

Network Topological Features
- Nodes = 478
- Edges = 805
- Avg. Degree = 3.36
a positive correlation with soil pH in the present study and supported by the results of Yang et al. (2020) [52].

Soil microorganisms, in general, do not function individually and instead establish a complex relationship network. Therefore, the relationship between diverse soil microbes has played a significant role in the functioning and stability of the microbial ecosystems [54]. The use of network analysis is a powerful technique for better understanding how related soil microbiomes adapt to environmental changes. In the current research, the network analysis revealed that bacterial taxa belonging to the phylum Proteobacteria, Acidobacteria, Actinobacteria were most dominant in the red soil, and consistent with the previous studies [55]. The highly connected nodes in RS1 compared with RS2 and RS3, revealed more complex and bacterial association in RS1, and such taxa are predictable as keystone bacterial taxa [54]. These keystones taxa are very crucial for maintaining the co-occurrence network structure [56]. Hence, the bacterial community in RS1 was more tolerant to environmental disturbances owing to complex connectivity because it has been widely accepted that bacterial communities with higher connectivity are more resilient to environmental changes compared with weaker connectivity of simple networks [57].

Numerous studies have confirmed that the functional composition of soil microbiota rather than taxonomic composition in natural environments appears to be closely related to environmental factors [58–60]. Although, the ecological functions of soil microbes inhabiting similar environments are closely related but their community composition may differ greatly. Hence, it implies that besides identifying which soil microorganisms are present in the environment, it is also a matter of great importance to reveal the functional profile of soil microbial communities. The FAPROTAX predicted functional analysis of bacterial communities exhibited the greatest number of OTUs for metabolic processes related to C-cycle and N-cycle and energy production in all soil depth (Fig. 8, Supplementary Fig. 2). Hence, these fundamental resource pathways were considered as potential drivers of bacterial community structure and the key functions of the bacterial community in the red soil [61]. The C- and N-cycles are a matter of great concern to understand their regulation in soil microbial ecology because soil bacterial communities play a key role in the regulation of biogeochemical cycles. Our findings suggest that the functional profile of the bacterial community was significantly shaped by the soil pH [62].
general, the FAPROTAX analysis revealed the microbial community’s expected functions, provided a brief outline of functional potential within the microbial community. Therefore, we recommend that further in-depth studies of metagenomic sequencing are needed for this system to fully evaluate the potential functional gene categories for a comprehensive understanding of the microbial ecology under different soil profiles.

**Conclusions**

The ultimate objective of exploring the bacterial diversity is to obtain a deeper understanding of who is there and doing what? To elucidate these questions, reproducible, reliable, quantitative, and statistically effective experimental information is required. Here, 16S rRNA gene high-throughput sequencing technology was deployed to analyze the bacterial diversity in the red soil of the pomelo orchard of Fujian Province. Our results showed that soil depth strongly influenced the soil bacterial communities. The relative abundance of *Proteobacteria*, *Acidobacteria*, and *Bacteroidetes* decreased with increasing soil depth. Principal coordinates analysis also showed that bacterial community composition was significantly affected across the different soil depths. Among the measured edaphic factors, soil pH was the most significant dominant factor in elucidating the dissimilarity in bacterial community composition in red soil. The FAPROTAX revealed that predicted functions of bacterial communities related to the C and N cycle were dominated in the red soil and also significantly affected by the soil pH. Our results indicated that soil pH rather than soil nutrients was the main factor for soil bacterial diversity, community composition, and functional profile in the red soil of pomelo orchard. We recommend that further studies of metagenomics sequencing are needed to evaluate the potential functional gene categories. Overall, these results provide valuable findings regarding the structure and functions of bacterial communities and suggest alleviation of soil acidification by adopting integrated management practices to preserve the soil microbial communities for better ecological functioning.

**Methods**

**Sampling site and soil sampling collection**

Soil samples were collected from the red soil of pomelo orchards of Pinghe County Fujian province, Southeast China (24°02′–24°35′ N, 116°54′–117°31′ E). This region is characterized by subtropical monsoon climatic conditions with an annual precipitation 1600–2000 mm, and an average annual temperature ranges from 17.5–21.34°C. The soil type is classified as haplic ferrasol with sand, silt and clay about 39.2%, 35.8%, and 25.1%, respectively [63, 64]. In June 2019, we collected 57 soil samples of the red soil from three different vertical soil depths, i.e., 0–20 cm (RS1), 20–40 cm (RS2), and 40–60 cm (RS3) and each depth contained 22, 17, and 18 soil samples, respectively, with the removal of top 5 cm soil layer to evade the exogenous disturbance. Each soil sample was a composite of 10 subsamples collected from the dripline, and two subsamples from each tree and trees were apart from 3 m from each other. The soil samples were immediately moved to the laboratory on ice. Subsequently, sieved through a mesh of 2 mm diameter, and remaining straw residues and fine roots were removed manually. Each soil sample was separated into two portions, one of which was used to determine soil physicochemical characteristics (stored at 4°C), and the other part for molecular analyses (stored at -80°C until use). The detail of intensive input of N.P.K has been shown in Supplementary Table S1.

**Soil physicochemical properties**

For determination of soil pH, soil/water (1:2.5, w/v) suspension was prepared, and pH meter (ORION A215 STAR, Thermo Ltd., USA) was used for its determination. Available phosphorous (AP) was measured by using sodium bicarbonate (NaHCO₃, 0.5 mol·L⁻¹) at pH 8.5 (2.5 g of soil was mixed with 50 mL solution, and shaken for 30 min) [65]. For Available potassium (AK), ammonium acetate (NH₄CH₃CO₃) extraction and subsequent flame photometer analysis were performed [66]. Available nitrogen (AN) was extracted by sodium hydroxide (NaOH) hydrolysis [67].

**Soil DNA extraction**

Total soil DNA was extracted from the 0.5 g of a soil sample by Soil DNA-Extraction-Kit (MO-BIO Laboratories, Carlsbad, CA-USA) using the manufacturer’s protocol. The agarose gel-electrophoresis and Nanodrop (Thermo Scientific NanoDrop 2000) were used to detect DNA purity and concentration. An appropriate amount of sample was taken in a centrifuge tube, diluted the sample with sterile water to 10 ng/μl, and stored in a refrigerator (−40°C) for subsequent analyses.

**PCR assays and high-throughput sequencing**

The extracted total soil DNA was further amplified by using the specific primers. The bacterial 16S V4-V5 region primers 515-F (5'-GTG GCCAGCMGGC GC GTAA-3') with 909-R (5'-CCCGYCAA TCM TTTT RAGT-3') were used [68]. The mixture of PCR (25 µl) contained the 1 × -PCR buffer, MgCl₂ (1.5 mM), deoxy-nucleoside triphosphate (0.4 µM), TaKaRa Ex-Taq of 0.5 U, 1.0 µM concentration for each primer, and 10 ng of soil genomic DNA. The PCR reaction included the following steps; initial denaturation at 94 °C for 3 min, followed by
30-cycles at 94°C for 40 s, 56°C for 60 s, and 72°C for 10 mint for the final extension. The two PCR reactions were run for each sample and then combine after PCR-amplification. The PCR products were mixed and subjected to gel-electrophoresis (1% agarose); targeted DNA bands were excised and a gel extraction kit was used for purification. For quantification of PCR products, a NanoDrop was used and then pooled together with an equal molar amount from each sample. For sequencing, the samples were prepared according to manufacturer instruction of the TruSeq DNA kit. The constructed library was quantified by Qubit and qPCR and then sent for sequencing on Illumina Miseq system (Bobett Biotechnology Co., Ltd, Sichuan, China).

**Processing of sequencing data**
Quantitative Insights Into Microbial Ecology (QIIME v1.9.0) program was used to process the raw sequences with the default settings [69] and UPARSE pipeline [70]. The primers, barcode sequences, and low-quality reads were removed from the analysis. The Operational Taxonomic Units (OTUs) were clustered at 97% sequence similarity. The OTU representative sequence (the sequence with the highest frequency appears in the OTU) was selected and Greenenes was used as reference database for annotation. The sequencing data is available at NCBI BioProject SRA database under the accession number PRJNA714448.

**Data analyses**
For downstream statistical analyses, we used the R V_4.0.3 [71]. In total, 4 83 879 high-quality sequences were achieved from all 57 samples with 7914 to 8709 sequences per sample. To strengthen the influence of the sequencing complexity on soil microbial diversity and community composition, the OTUs table was rarified so that each sample having 7900 reads. The Observed, Shannon, Chao1, and ACE indices were used for the determination of species richness and alpha diversity. The Kruskal–Wallis (KW) rank-sum test was used to see if there were any significant differences in alpha diversity between different groups of soil depths. For microbial beta diversity, the principal coordinate analysis (PCoA) based on Bray–Curtis dissimilarity was performed to check the differences in microbial diversity among different soil depths. Furthermore, permuational analysis of variance (PERMANOVA) was performed to assess the significant differences of bacterial communities among different soil depths. The relationship between soil bacterial communities and soil physicochemical parameters, including soil pH, available nitrogen (AN), available phosphorous (AP), and available potassium (AK), was investigated using distance-based redundancy analysis (dbRDA). The mantel test, based on Pearson correlation between soil attributes and Bray–Curtis dissimilarity score, was used to determine which soil factors had a substantial impact on bacterial communities. Correlation analysis between soil properties and the bacterial population is very important in analyzing and inferring the most important factors shaping the community structure. All these statistical analyses were performed using the R package "microco v0.2.0" [72]. Analysis of variance was applied using Statistix 8 (Version 8.1) to study soil physiochemical properties under different soil depth and significant differences between different soil depths were compared based on least significant difference (LSD, $P \leq 0.05$).

The co-occurrence network of bacterial communities was determined according to Spearman's correlation. OTUs with significant correlations ($P < 0.01$, $\rho > 0.7$) were chosen. These correlations were revealed by pairwise analysis of taxa abundance and resulted in a highly complex network whereby each node described the phylum, whereas internode (stand among the nodes) represented the significant correlation between the nodes. For visualization and modularity of co-occurrence, Gephi V0.9.2 was used. Nodes with a high degree and relative abundance were categorized as keystone species in the co-occurrence network [73]. Finally, for predicting potential functions of soil bacterial community, functional annotation of Prokaryotic taxa (FAPROTAX) was used with default settings [74].

**Abbreviations**
AH: Available nitrogen; AP: Available phosphorous; AK: Available potassium; RS: Red soil; OTU: Operational taxonomic unit; RA: Relative abundance; PCoA: Principal coordinates analysis; PERMANOVA: Permuational multivariate analysis of variance; dbRDA: Distance-based redundancy analysis; FAPROTAX: Functional annotation of prokaryotic taxa.

**Supplementary Information**
The online version contains supplementary material available at https://doi.org/10.1186/s12866-022-02452-x.

**Additional file 1: Supplementary Figure S1.** The relative abundance of the bacterial communities at the phylum level in different soil profiles. The least significant test (LSD test, $P < 0.05$) was applied to check the significance of bacterial relative abundance between the different groups.

**Supplementary Figure S2.** Functional analysis of bacterial communities on the basis of % OTUs. The symbol M represents the module, and contains a set of OTUs. **Supplementary Table S1.** The intensive application of N, P, K in Pinghe County.

**Acknowledgements**
The authors would like to thank Prof. Chunjian Li, Prof. Shwei Guo and Prof. Xuexian Li for providing useful suggestions and guidance to improve the study.
Authors’ contributions
MAM, WH, XH and CZ designed the experiment. MAM, WH, JL helped in sampling and performed experiments. MAM, MRK, YC and WY analyzed the data, and prepared the tables and figures. MAM and WH wrote the manuscript. UX, BJ, and CZ review and edited the manuscript. All authors have read and approved the final manuscript.

Funding
This research was funded by the National Natural Science Foundation of China (41601244) and the Open Research Foundation of International Magnesium Institute (ML2018-09).

Availability of data and materials
All data generated or analyzed during this study are included in this article and its supplementary information files. The raw reads of sequencing data is available at NCBI BioProject SRA database under the accession number PRJNA714448.

Declarations

Ethics approval and consent to participate
Not applicable

Consent for publication
Not applicable

Competing interests
There is no conflict of interest.

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Received: 26 August 2021 Accepted: 24 January 2022
Published online: 02 February 2022

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