Original Research

Links between Arbuscular Mycorrhizal Fungal and Nitrogen-Fixing Bacterial Communities from Plant Rhizosphere Soils in the Karst Region of Southwest China

Yueming Liang², Yirong Su¹,³, Xiangbi Chen¹,³, Fujing Pan⁴, Xunyang He¹,³*

¹Key Laboratory of Agro-ecological Processes in Subtropical Region, Institute of Subtropical Agriculture, Chinese Academy of Sciences, China
²Institute of Karst Geology, CAGS, Karst Dynamics Laboratory, MLR, China
³Huanjiang Observation and Research Station for Karst Eco-systems, Chinese Academy of Sciences, Huanjiang, China
⁴College of Environmental Science and Engineering, Guilin University of Technology, China

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Abstract

Arbuscular mycorrhizal (AM) fungi and nitrogen-fixing bacteria (NFB) are critical to plant growth and recovery in degraded ecosystems. However, information about the relationships between these microorganisms in the rhizosphere soils of natural ecosystems is limited. Rhizosphere soils from six common plant species: (Loropetalum chinense, Syzygium championii, Alchornea trewioides, Bauhinia purpurea, Pterolobium punctatum and Albizia odoratissima), were sampled in the karst region of Southwest China. The abundance and community composition of AM fungi and NFB were measured using quantitative polymerase chain reaction (qPCR) and terminal restriction fragment length polymorphism (T-RFLP). The community composition and abundance of AM fungi and NFB varied among plant rhizosphere soils, but the richness of these microbes was not significantly different. The abundances of AM fungi and NFB significantly related to each other. Specifically, dominant 278-bp and 105-bp terminal restriction fragments (T-RFs) of AM fungi positively linked with dominant 184-bp and 180-bp T-RFs of NFB (P<0.05), respectively. Redundant analyses indicated that soil organic carbon, available phosphorus, and total nitrogen significantly correlated with the composition of fungal and bacterial communities. These results suggest that AM fungi and NFB have host-plant specificity and that these microbes are closely linked in plant rhizosphere soils of the karst region.

Keywords: arbuscular mycorrhizal fungi, nitrogen-fixing bacteria, soil microbial diversity, karst, rhizosphere

*e-mail: hbbpjh@isa.ac.cn
Introduction

The soil microbial community is critical for plant growth and development. In particular, two major groups of rhizosphere microbes, arbuscular mycorrhizal (AM) fungi and nitrogen-fixing bacteria (rhizobia), increase the supply of nitrogen, phosphorus, and other essential nutrients to plants [1-3], thus, promoting their establishment in a given soil habitat. Moreover, AM fungi strengthen the physical composition of soil by stabilizing soil aggregate structure [4]. Thus, these microbes play major complementary roles in the successful recolonization of plants in severely disturbed soil systems by establishing a sustainable nutrients cycle and improving soil stabilization [5].

Given that their beneficial effects both derive from altering soil properties to improve nutrient uptake, the two microbial taxa may act synergistically in promoting plant growth. For example, increased phosphorus uptake stemming from the action of AM fungi yields more energy for rhizobia to fix nitrogen [6]. Indeed, co-inoculation with AM fungi and NFB appeared to stimulate plant growth and biomass increase [2, 3, 7], more so than either inoculation with AM fungi or NFB alone [2]. However, other experiments have yielded inconsistent results. Larimer et al. [8] reported that rhizobia inoculation suppresses AM fungal colonization of roots, even though AM fungal infection increases root nodule number and plant mass. Additionally, these two microbial taxa were not synergy due to plants did not perform better than expected given symbiont individual effects [9, 10]. As can be seen from these results, no clear consensus exists on the relationship between AM fungi and NFB at the species level. Furthermore, most of the studies examining the interaction of these symbiotic microorganisms are usually controlled pot experiments, with few field studies that validate laboratory findings. The lack of field data is particularly notable in soil-nutrient deficient ecosystems, which are especially dependent on AM fungi and rhizobia for plant growth.

Karst landscapes are ideal for such field research because the soil nutrient content is considerably lower than other regions at the same latitude [11] and is easily lost due to the prominent underground drainage system characteristic of this topography [12, 13]. The karst region of Southwest China, one of the largest in the world, covers 540,000 km² [14]. These regions are characterized as low vegetation cover and a high ratio of bedrock outcrop to shallow soil. These features cause the region’s ecological systems to be fragile and extremely vulnerable to human activities. Recently, an increasing human population and the resultant over-exploitation of natural resources have accelerated desertification [15]. In response, local Chinese governments have begun to promote ecological restoration of karst regions, and successful recovery is currently evaluated based on the degree of plant establishment [16]. Dominant plant and legume are widely used for vegetation restoration. However, the relationships between AM fungi and NFB from plants rhizosphere soils in the karst region are poor, although these two microbial taxa have confirmed the benefits of AM fungi and NFB on plant establishment in poor environmental conditions based on pot experiments.

In this study, we hypothesized that a high abundance of AM fungi will be associated with a high abundance of NFB in rhizosphere soils of the Chinese karst region. Our objectives were 1) to characterize AM fungal and NFB communities in rhizosphere soils from six typical plant species of a karst shrub ecosystem and 2) to explore the relationships between AM fungal and NFB communities.

Materials and Methods

Study Site and Sample Collection

The study site is located in Huanjiang County, in the Guangxi autonomous region of southwest China (107°51' to 108°43'E, 24°44' to 25°33'N). This region is dominated by a subtropical mountainous monsoon climate, with a mean annual rainfall of 1,389 mm and a mean annual air temperature of 18.5°C. The wet season, during which 70% of the annual precipitation occurs, lasts from April to August [17].

Six representative shrub species: *Loropetalum chinense* (Hamamelidaceae), *Syzygium championii* (Myrtaceae), *Alchornea trewioides* (Euphorbiaceae), *Bauhinia purpurea* (Leguminosae), *Pterolobium punctatum* (Leguminosae), and *Albizia odoratissima* (Leguminosae), were selected in each plot. *Loropetalum chinense, S. championii,* and *A. trewioides* were defined as the dominant species, whereas *B. purpurea, P. punctatum,* and *A. odoratissima* were considered non-dominant and leguminous plants [18]. Considering the higher spatial heterogeneity of soil nutrients in karst ecosystems, plots were established and located in the same topographic position (southeast aspect and middle slope) and rock type (dolomites).

The rhizosphere soils were sampled as previously described by Bell et al. [19] in December 2011. Five individual rhizosphere soils of each host plant in each plot (10 m × 10 m) were carefully excavated, collected, and thoroughly mixed to form a composite soil sample (18 samples in total: 6 plant species × 3 sampling plots). Because plant age influences AM fungal community composition, we attempted to select plants of comparable age, based on similar basal diameter (approximately 2 cm). Additionally, to minimize the effects between hosts, plants were separated by approximately 50 cm. We confirmed that the plants could form nodules when rhizosphere soils were sampled. Each composite soil sample was divided into two subsamples. One subsample was frozen in liquid nitrogen and transported to the laboratory for molecular analysis; the other was air-dried for analysis of soil.
physicochemical properties. The field study sites did not involve any privately-owned land or protected area of land (such as national park), and the sampling did not involve any endangered or protected species. Thus, no specific permits were required for the study field sites.

Soil Physicochemical Properties

Total nitrogen (TN), available phosphorus (P), soil organic carbon (SOC), and pH were measured. Soil pH was determined using a solution of soil and distilled water (1:2.5 w/v). We measured SOC using K$_2$Cr$_2$O$_7$·H$_2$SO$_4$ oxidation-reduction titration and TN using the Kjeldahl method [20]. Available P was extracted using 0.5 M sodium bicarbonate and measured using the Mo-Sb colorimetric method [21]. Soil physicochemical properties are shown in Table 1.

DNA Extraction from Soil

Microbial DNA was extracted in triplicate from 500 mg of freeze-dried soil, following the methods of a previous study [17]. The extracted DNA was dissolved in 50 μL water, quantified by spectrophotometry, and stored at -20°C until further use. Prior to real-time quantitative polymerase chain reaction (qPCR), the DNA template was prepared by diluting the solution to a final concentration of approximately 5 ng DNA μL$^{-1}$ using sterilized water.

PCR Amplification and Terminal Restriction Fragment Length Polymorphism (T-RFLP) Analyses

The composition of the AM fungal and NFB communities in the 18 soil samples was estimated with terminal restriction fragment length polymorphism (T-RFLP) analysis. The extracted DNA was subjected to nested PCR with primers Geo11F/GeoA2R and NS31/AM1 (Table 2) for amplification of an 18S rRNA gene fragment of AM fungi. The latter forward primer was labeled at the 5’ ends with 6-carboxy-fluorescein (FAM; Invitrogen, China). The first PCR (50-μL volume) of Geo11F/GeoA2R contained 25 μL 2× PCR Premix (0.1 U Prime STAR HS DNA polymerase, 0.5 mM dNTPs; Tiangen, China), 10 pM of each primer, 20 ng genomic DNA, and 19 μL H$_2$O. The cycling conditions were as follows: 95°C for 2 min, 35 cycles of 60 s at 94°C, 60 s at 60°C, and 60 s at 72°C, followed by 72°C for 10 min. The first amplification product was diluted with double-distilled water (1:10) and a 1-μL subsample was used as a template for the second PCR amplification under the same conditions, with the following exceptions: 30 PCR cycles were performed instead of 35 and the annealing temperature was 64°C instead of 60°C.

The PoI/PoIR primer set was used to amplify the NFB nifH gene (Table 2). The reaction was performed in a 50-μL volume with 25 μL 2× PCR Premix (0.1 U Prime STAR HS DNA polymerase, 0.5 mM dNTPs), 10 pM of each primer, 20 ng genomic DNA, and 19 μL H$_2$O. Cycling conditions for the nifH gene were an initial denaturation step at 95°C for 2 min, 35 cycles at 94°C for 30 s, 55°C for 60 s, and 72°C for 60 s, followed by a final 10-min extension at 72°C.

Triplicate reaction mixtures per sample were pooled, purified using the QIAquick PCR purification kit (Tiangen Biotech Ltd., China), and quantified using NanoDrop ND-1000 (Thermo Scientific, USA). Approximately 200 ng of each amplicon was digested with 5 U restriction enzyme. Restriction enzymes HinFI and HaeIII were used to digest the fungal 18S rRNA gene and the bacterial nifH gene, respectively. The digestion products were analyzed using an automated sequencer (model 373A; Applied Biosystems, Weiterstadt, Germany) by the Sunny Company (Shanghai, China).

| Plant species | AP (g.kg$^{-1}$) | SOC (g.kg$^{-1}$) | TN (g.kg$^{-1}$) | pH | Richness | Evenness |
|--------------|----------------|-----------------|----------------|----|----------|----------|
|              |                |                 |                |    | AM fungi | NFB      | AM fungi | NFB      |
| L. chinense  | 8.23±0.66b     | 57.78±2.12b     | 4.70±0.35b     | 7.47±0.16a | 15±1a    | 9±1a     | 0.92±0.02a | 0.77±0.03b |
| S. championii| 8.27±0.89b     | 59.41±6.61b     | 5.46±1.23b     | 7.53±0.14a | 13±3a    | 9a       | 0.88±0.03a | 0.86±0.02a |
| A. trewioides| 8.74±0.76b     | 67.53±8.05b     | 5.87±1.26b     | 7.29±0.16a | 14a      | 8±1a     | 0.92±0.02a | 0.84±0.01a |
| B. purpurea  | 10.25±0.36b    | 80.44±12.44b    | 7.11±0.20a     | 12±1a    | 9a       | 0.89±0.04a | 0.88±0.01a |
| P. punctatum | 12.6±0.37b     | 101.97±23.03b   | 10.11±2.73b    | 7.09±0.13a | 14±1a    | 9a       | 0.91±0.01a | 0.88±0.02a |
| A. odoratissima| 20.81±6.23a   | 224.99±82.14a   | 20.39±8.02a    | 7.19±0.37a | 13±1a    | 9a       | 0.90±0.01a | 0.89±0.79a |

Values are means±SE of three replicates.
Within rows, means with the same letter do not differ significantly at $p<0.05$ (LSD).
SOC, soil organic carbon; TN, total nitrogen; NFB, nitrogen-fixing bacteria.
The size and relative abundance of terminal restriction fragments (T-RFs) were evaluated using Gene Scan 2.1 (Applied Biosystems). Peak areas of T-RFs that differed by ±2 bp were summed and considered as one fragment. The relative abundance (RA) of each T-RF was calculated as previously described [26]: where \( n_i \) represents the peak area of one distinct T-RF, and \( N \) is the sum of all peak areas in one sample. T-RFs that were > 50 bp in length and with RA>1% in all three replicates were analyzed. Peaks with RA>5% were regarded as dominant T-RFs.

As Aldrich-Wolfe et al. [27] described, using database T-RFLP identified AM fungal and NFB species: (i) T-RFLP profiles of AM fungi were determined for our 454 pyrosequencing sequences that were submitted to the MG-RAST public database (http://metagenomics.anl.gov/) under ID 4540338.3. And (ii) T-RFLP profiles of NFB were determined for 40 \( nifH \) sequences from karst region (accession numbers KF859859 to KF859898).

Quantification of 18S rRNA and \( nifH \) Genes

Abundances of the AM fungal 18S rRNA gene and the NFB \( nifH \) gene were determined using qPCR (ABI 7900, Foster City, CA) with primers AMV4.5NF/AMDGR and PolF/PolR, respectively. Although AMV4.5NF/AMDGR can amplify non-AM fungal sequences, more than 70% of sequences obtained from the 454 pyrosequencing with these primers belonged to AM fungi in karst region [28].

The 10-μL reaction mixture contained 5 μL 1× SYBR Premix ExTaq, 0.2 μL Rox (Takara Bio, Shiga, Japan), 0.2 μM of each primer (Invitrogen, China), 1 μL 5-ng DNA template, and 3.4 μL sterilized water. The thermocycling protocol for the fungal 18S rRNA gene was 20 s at 95°C, followed by 30 cycles of 10 s at 95°C, 15 s at 62°C, and 15 s at 72°C. The thermocycling protocol for the bacterial \( nifH \) gene was 20 s at 95°C, followed by 5 cycles of 15 s at 95°C, 20 s at 64°C, and 15 s at 72°C, then 35 cycles of 15 s at 95°C, 25 s at 60°C, and 15 s at 72°C.

A standard curve was established from a 10-fold dilution series (10^{-2}-10^{8} copies) of the plasmids containing the target gene fragment. The efficiency of real-time qPCRs was 98% for the AM fungal 18S rRNA and 103% for the NFB \( nifH \) gene, while the \( R^2 \) value for the curves of both two genes were 0.99. The qPCR assays were performed using four technical replicates per sample for all samples in one plate. A single, sharp peak was observed for the melt curves of each assay. Data analysis was performed automatically using the SDS 2.3 software included with the real-time qPCR system.

### Statistical Analyses

Statistical analyses were performed using SPSS 19.0 for Windows (SPSS Inc., Chicago, IL). Differences were considered significant at \( p<0.05 \). Differences in soil physicochemical properties between plants were evaluated using least significant difference (LSD) analysis. Data that were not normally distributed were \( \log_{10}(x+1) \)-transformed. Diversity indices of AM fungi and NFB were calculated using PC-ORD 5.0 (MJM Software Design, Gleneden Beach, OR). Relationships between soil physicochemical properties, and abundances of fungal and bacterial were evaluated using the Pearson product-moment correlation coefficient. Redundancy analysis (RDA) was used to analyze the effects of soil physicochemical characteristics on microbial community composition, in CANOCO 4.5 (Microcomputer Power, Inc., Ithaca, NY). Before the RDA analysis, detrended correspondence analysis (DCA; gradient length <3) was performed to confirm that the linear ordination method was appropriate for analyzing the T-RFLP data. Monte Carlo permutation tests were also used to compute statistical significance. The relationships between community composition of AM fungi and NFB were assessed based on Pearson's
correlations using Mantel tests with the Ecodist package in R software.

Results

Abundance and Diversity of AM Fungi and NFB in Rhizosphere Soils

The range of AM fungal abundance was 5.57 from to 6.77 log copies⁻¹ soil, which varied among the plant rhizosphere soils (Fig. 1); *A. odoratissima* and *A. trewioides* exhibited the highest and lowest abundance, respectively. The evenness and richness indices of AM fungi were not significantly different among plants (Table 1).

The range of NFB abundance was 6.46 from to 7.05 log copies⁻¹ soil, which varied among the plant rhizosphere soils (Fig. 1); again, *A. odoratissima* and *A. trewioides* exhibited the highest and lowest abundance, respectively. The evenness index of NFB was lower in *L. chinense* than in other plants, but the richness index was similar across all examined plants (Table 1).

Composition and Structure of AM Fungal and NFB Community in Rhizosphere Soils

We used 28 T-RFs in the ribotypes to analyze AM fungal community composition and structure in rhizosphere soils (Fig. 2a). There were 16, 14, 20, 14, 16, and 14 T-RFs for AM fungi from *L. chinense*, *S. championii*, *A. trewioides*, *P. punctatum*, *B. acuminata*, and *A. odoratissima*, respectively. Six T-RFs (105, 130, 138, 189, 278, and 300 bp in length) were predominant in the rhizosphere soils of all plant species and accounted for 55% of all T-RFs, and the dominant 300 bp T-RF for AM fungi was the significantly different T-RFs among plants. Significant variation existed among the plant species in 27 AM fungal T-RFs (Fig. 2a). Although there are limitations to extrapolating species identities from T-RFs, the dominant 138-, 189-, and 278-bp T-RFs were closely related to *Glomus* sp. M20, *Glomus* sp. MUCL, and *Glomus macrocarpum*, respectively.

We obtained 13 T-RFs for NFB (Fig. 2b) and 11 T-RFs significantly differed among these soils; of these, 6, 7, 9, 7, 9, and 8 were found for *L. chinense*, *S. championii*, *A. trewioides*, *P. punctatum*, *B. acuminata*, and *A. odoratissima*, respectively. Five dominant T-RFs (66, 75, 157, 180, and 184 bp) accounted for 80% of the NFB T-RFs in rhizosphere soils, and the dominant 180 bp T-RF for NFB was the significantly different T-RFs among plant soils. Our analyses indicated that the dominant 75, 157, 180, and 184-bp T-RFs were closely related to *Bradyrhizobium* sp. CCBAU 101065, *Bradyrhizobium* sp. ISA1601, *Bradyrhizobium japonicum*, and *Bradyrhizobium* sp. ISA0508, respectively.

Effects of Host Plant and Soil Physicochemical Properties on Microbial Communities

Results of the RDA revealed that the fungal community composition differed among plants, especially between *A. odoratissima* and the other plants (Fig. 3a). The NFB community composition had a similar distribution as AM fungi (Fig. 3b). These results indicated that host plant species had a strong effect on microbial community composition. Available phosphorus was significantly correlated with AM fungal (F = 2.80, p = 0.016) and NFB community composition (F = 4.80, p = 0.002). Additionally, SOC and TN were significantly correlated with AM fungi (Fig. 3a; F = 2.80, p = 0.016 and F = 2.80, p = 0.016, respectively) and with NFB community composition (Fig. 3b; F = 4.71, p = 0.002 and F = 3.44, p = 0.004, respectively).

![Fig. 1. Copy numbers for AM fungi and NFB in rhizosphere soils collected from different plant species. Different letters indicate significant differences between plants (LSD test; p<0.05). Bars indicate SE (n = 3).](image-url)
The contents of available phosphorus, soil organic carbon, and total nitrogen were positively correlated to the AM fungal abundance \((p<0.05, \text{Table 3})\). Similarly, the contents of soil organic carbon and available phosphorus were negatively correlated with the NFB abundance \((p<0.05, \text{Table 3})\).

**Links between AM Fungi and NFB Communities in Rhizosphere Soils**

The Mantel test showed that AM fungal community composition was not significantly correlated with NFB community composition in rhizosphere soils \((r=0.093, \ p=0.115)\). However, RDA analysis found that the dominant 278-bp T-RF \((\text{Glomus macrocarpum})\) from AM fungi was highly correlated with the dominant 184-bp T-RF \((\text{Bradyrhizobium sp. ISA0508})\) from NFB. In addition, the 105-bp T-RF \((\text{uncultured Glomus})\) from AM fungi positively correlated with the 180-bp T-RF \((\text{Bradyrhizobium japonicum})\) from NFB (Fig. 4).

No significant correlation was found between the fungal and bacterial richness \((r=0.110, \ p=0.661)\) or evenness \((r=-0.388, \ p=0.112)\) indices, but the abundances of AM fungi and NFB were highly

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**Fig. 2.** Average relative abundance of AM fungi a) and NFB b) terminal restriction fragments (T-RFs) in rhizosphere soils of different plant species, as determined by endonuclease digestion with \(\text{Hin} \text{fI} \) (fungi) and \(\text{Hae} \text{III} \) (bacteria). The relative abundance of T-RFs is given as a percentage of the total peak area. Fragment sizes in the graph indicate the size of the experimental T-RFs. Bars indicate SE \((n=3)\). Among the plant species, 27 T-RFs \((66, 76, 86, 94, 105, 110, 130, 134, 138, 144, 158, 189, 196, 252, 260, 271, 278, 284, 289, 300, 304, 315, 325, 348, 365, \) and 375 bp) from AM fungi and 11 T-RFs \((62, 85, 97, 125, 151, 157, 169, 180, 184, 201, \) and 315 bp) from NFB differed significantly.
Links between Arbuscular Mycorrhizal Fungal...

Discussion

Effects of Host Plant Species and Soil Nutrients on Rhizosphere AM Fungal Community

AM fungi can improve soil nutrient content, thus, promoting plant growth, and studies have demonstrated that higher AM fungal colonization tends to increase soil nutrient contents [2, 29]. In the present case,
AM fungal abundance was positively correlated with available phosphorus content as well as the contents of total nitrogen and soil organic carbon. Our result was not in agreement with the previous study by Gosling et al. [30]. Several explanations are possible for this inconsistent result. First, the soil nutrient content in the karst region may not have been lower than the threshold that leads to reducing AM fungal abundance. Second, other characteristics, such as soil structure, may alter how soil nutrients affect AM fungal diversity [31].

Fungal abundance varied among the plants and was higher in non-dominant plants than in dominant ones (except for *S. championii*). This outcome suggests that non-dominant species were more dependent on AM fungi in karst regions. Additionally, the AM fungal community composition varied across plants (Fig. 3a), suggesting the presence of host-plant specificity [32]. The previous study has found that specificity in AM fungi-plant interactions may occur more often on the level of plant functional groups than individual species [32]. While we did not address this possibility in the current study, future research will attempt to verify whether the relevance of plant functional groups also applies for fungi-plant relations in karsts.

Despite plant-specific variation in abundance, the dominant fungal taxon in all rhizosphere soils was Glomerales. This order is also dominant in many host plants from other regions [33, 34], likely because these fungi can colonize plants from fragments of mycelium or mycorrhizal roots. Compared with other taxa, Glomerales appear to be more flexible and can better adapt to a range of environments [34, 35], including karst regions [28].

The richness of AM fungi did not vary significantly among rhizosphere soils. This outcome is likely due to the ability of AM fungal species to form symbioses with several host plant species [36], even when they have host-plant preferences. As a result, the richness of AM fungi is generally lower and less variable compared with host plant richness: approximately 100 species of AM fungi have been described based on morphology, versus more than 100 000 terrestrial plant species [37, 38]. We obtained lower AM fungal richness than other studies that used the same methods [39]. This disparity may due to variation in the degree of fungal interspecific competition across the different study sites. In our study, we found six predominant T-RFs for AM fungi associated with all of the plant species. These dominant species suppress the growth of non-dominant species, thus reducing and homogenizing total microbial richness across rhizospheres.

Effects of Host Plant Species and Soil Nutrient Content on NFB in the Rhizosphere

The abundance of NFB was higher in the rhizospheres of non-dominant plants than dominant plants (except for *S. championii*). We believe this result is likely due to differences in soil properties between the rhizospheres of non-dominant versus dominant plants. Among soil properties, soil carbon availability especially affects NFB abundance [40]. Thus, higher soil carbon availability in the rhizosphere of non-dominant plants can improve growth and survival of NFB, increasing their abundance.

Among the non-dominant legume plants, bacterial abundance was highest in the *S. championii* rhizosphere, which also exhibited high AM fungal abundance. Previous research has found that AM fungi enhance the nutrient-uptake capacity and thus the abundance of NFB [41], explaining our results. Therefore, plant species-dependent variation in soil nutrients and the presence of AM fungi may be the major factors influencing the abundance of NFB.

Host plants also have a great effect on the NFB community composition [18]. Plants affect microbial community composition mainly through the composition and concentration of root exudates [42, 43]. Root exudates influenced soil available phosphorus and further had great effect on bacterial community composition, corroborating previous results [44, 45]. Despite bacterial community composition variation in all rhizosphere soils, the dominant bacterial taxon was the genus *Bradyrhizobium*. This genus is widely distributed in China [46, 47], and typically prevalent in high-pH soils [39].

Relationships between AM Fungi and NFB in Rhizosphere Soils

Despite numerous studies on the topic, researchers disagree about the nature of interactions between AM fungi and NFB [1, 48]. Here, we found significant correlations between the dominant T-RFs of AM fungi and NFB (Fig. 4), implying that these microorganisms exhibit similar preferences for host-plant species. One possible explanation for this preference is that species-specific exudates from AM fungi may activate particular NFB species, resulting in selective and stable cooperation between distinct species of both microbial
taxa on the same host plant. This interaction can help explain why different effects on plant growth were observed in co-inoculation experiments with AM fungi and NFB [2, 9, 10]. We suggest that the compatibility of fungi and bacterial communities should be considered when targeting plants for co-inoculation as a means of restoring vegetation in karst regions.

We found a concurrent increase in the abundance of AM fungi and NFB (Fig. 5), supporting the close relationship of the two taxa. Their combined effects on plant growth may explain the concurrent increase. AM fungi enhances phosphorus uptake [49], and phosphorus is essential for bacterial N-fixation and further leads to their increased abundance [6]. This increase in available nitrogen then serves to improve plant metabolism and productivity. As a result, plants are able to provide large quantities of carbohydrates to AM fungi, increasing fungal abundance in a positive feedback loop [39]. If this interaction occurs in the low-nutrient karst soils, then the observed positive correlation between AM fungal and NFB abundance is probably important to vegetation restoration in the karst region. While our findings have potential applications for all nutrient-poor soils, further study is needed to determine whether the relationship between fungal and NFB abundances occurs in other regions.

Conclusions

In conclusion, the composition of AM fungal and NFB communities varied among plant species and was closely related to soil properties (total nitrogen, soil organic carbon, and available phosphorus). Fungal and bacterial taxonomic abundance increased concurrently. Moreover, these microbes are similar in their host-plant specificity. Therefore, inoculating plants with compatible species of AM fungi and NFB should promote target plant growth and establishment, improving the success of vegetation restoration. However, limitations inherent to the T-RFLP analysis meant we could not definitively identify which microbial taxon was dominant in soil. Thus, future studies aiming to characterize AM fungal and NFB communities should combine morphological and molecular methods.

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Conflicts of Interest
The authors declare no conflict of interest.

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