Rhizosphere soil bacterial communities and nitrogen cycling affected by deciduous and evergreen tree species

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Abstract
Deciduous and evergreen trees differ in their responses to drought and nitrogen (N) demand. Whether or not these functional types affect the role of the bacterial community in the N cycle during drought remains uncertain. Two deciduous tree species (Alnus cremastogyne, an N2-fixing species, and Liquidambar formosana) and two evergreen trees (Cunninghamia lanceolata and Pinus massoniana) were used to assess factors in controlling rhizosphere soil bacterial community and N cycling functions. Photosynthetic rates and biomass production of plants, 16S rRNA sequencing and N-cycling-related genes of rhizosphere soil were measured. The relative abundance of the phyla Actinobacteria and Firmicutes was higher, and that of Proteobacteria, Acidobacteria, and Gemmatimondaetes was lower in rhizosphere soil of deciduous trees than that of evergreen. Beta-diversity of bacterial community also significantly differed between the two types of trees. Deciduous trees showed significantly higher net photosynthetic rates and biomass production than evergreen species both at well water condition and short-term drought. Root biomass was the most important factor in driving soil bacterial community and N-cycling functions than total biomass and aboveground biomass. Furthermore, 44 bacteria genera with a decreasing response and 46 taxa showed an increased response along the root biomass gradient. Regarding N-cycle-related functional genes, copy numbers of ammonia-oxidizing bacteria (AOB) and autotrophic ammonia-oxidizing archaea (AOA), N2 fixation gene (nifH), and denitrification genes (nirK, nirS) were significantly higher in the soil of deciduous trees than in that of the evergreen. Structural equation models explained 50.2%, 47.6%, 48.6%, 49.4%, and 37.3% of the variability in copy numbers of nifH, AOB, AOA, nirK, and nirS, respectively, and revealed that root biomass had significant positive effects on copy numbers of all N-cycle functional genes. In conclusion, root biomass played key roles in affecting bacterial community structure and soil N cycling. Our findings have important implications for our understanding of plants control over bacterial community and N-cycling function in artificial forest ecosystems.

Keywords
actinorhizal plants, co-occurrence network, mixed plantations, photosynthetic carbon, plant-soil interactions
1 | INTRODUCTION

Global climate change is likely to increase frequencies and severity of droughts, imposing soil water deficit stress on trees, which will markedly reduce productivity of plantations and natural forests (Gillespie et al., 2020; Pardos et al., 2021). Compared to monocultures, diverse plant communities are more likely to increase productivity and be more resistant and resilient to drought (Chen et al., 2021; Pardos et al., 2021). Increasing evidence suggests that the different drought-resistant abilities have been deeply impacted by soil microbial communities (Castro et al., 2019; Osburn et al., 2021). Changes in the bacterial community in the face of drought decline soil nitrogen (N) availability through impacting soil N cycling process (Chen et al., 2021; Wang et al., 2021). However, greater microbial diversity in more diverse forests is thought to be beneficial to facilitate efficient nutrient cycling process thus potentially better sustaining forest functions and stability compared to monospecific forests (Gillespie et al., 2020; Naylor & Coleman-Derr, 2018).

The N fixation, nitrification, and denitrification are key N cycling processes in soils, which are largely determined by specific microbial guilds and tightly connected with soil properties including water availability and pH (Bowen et al., 2020; McCulloch et al., 2020). Symbiotic N fixation by N$_2$-fixing plants constitutes the largest natural input of N into forest ecosystems (Ngom et al., 2016). Ammonia-oxidizing bacteria (AOB) and autotrophic ammonia-oxidizing archaea (AOA) have been primarily used to study nitrifying microbial communities (Guo et al., 2021; Trivedi et al., 2019). Genes (nirK and nirS) reflecting denitrifying microbial communities encode nitrite reductase to transform nitrite to nitric oxide (Bowen et al., 2020; Moreau et al., 2015). A lower soil water availability reduced N cycling by limiting extracellular enzymatic activities and impacting the N fixation, denitrifying or nitrifying microbial communities (Bowen et al., 2020; Castro et al., 2019; Guo et al., 2021).

In addition to effects of soil properties, plants can also strongly regulate soil microbial communities and microbiially driven N-cycling processes through litter or diverse root-derived inputs of labile organic compounds (Henneron et al., 2020; Mushinski et al., 2020). The rhizosphere soil hosts diverse microbial communities that are crucial to promote plant nutrient acquisition and resistance/ tolerance to abiotic stressors (Moreau et al., 2015; Naylor & Coleman-Derr, 2018). The N$_2$-fixing plants enhance N fixation ability by promoting photosynthesis and root-derived carbon releasing to rhizosphere soil. Symbiotic N fixation is costly, and allocation of labile carbon (C) to root nodules of N$_2$-fixing plants can be greatly reduced due to decreased photosynthetic C fixation during drought (Minucci et al., 2019). Plants are proposed to control soil N cycling in the rhizosphere to sustain their nutrition and growth (Henneron et al., 2020). However, the impacts on microbial community and N-cycling functions among plants with different N demand in response to drought need more research.

McCulloch et al. (2020) showed that drought reduced symbiotic N fixation by reducing nodule biomass and nitrogenase activity in eight N$_2$-fixing species. Root-nodule symbiosis of actinorhizal plants and bacteria of the genus Frankia is an important pathway of new N input into forest ecosystems, in addition to symbioses of legumes and Rhizobium bacteria (Ngom et al., 2016; Tobita et al., 2015). The actinorhizal plants play crucial roles in replenishing soil nutrients of agroforestry and improving bacterial community in degraded land (Farias et al., 2009; Ngom et al., 2016). Among actinorhizal plants, species belonging to the genus Alnus are distributed worldwide and have been cultivated in monoculture for revegetating or rehabilitating disturbed habitats and commercial production (Tobita et al., 2015; Uri et al., 2014). Henneron et al. (2020) concluded that acquisitive species with higher photosynthesis and N uptake induced a stronger acceleration of N cycling than conservative species in rhizosphere soil. Deciduous and evergreen trees represent opposite extremes along "leaf economics spectrum," which ranks from acquisitive and fast-growing to conservative and slow-growing traits (Wright et al., 2004). Deciduous trees with greater specific root length and specific leaf area show higher N uptake and photosynthetic capacity, growing faster than evergreen trees (Baldocchi et al., 2010; Cantarel et al., 2015; Wright et al., 2004). Growth of the deciduous species is more closely coupled to soil nutrient availability than evergreen species (Gray & Schlesinger, 1983). Higher N demand of deciduous trees may accelerate N cycling by imposing different selective pressures on soil bacterial community structures compared with evergreen species. Therefore, in the present study, we examined responses of soil bacterial community and functions associated with deciduous and evergreen trees to short-term drought. We predicted higher number of N-cycling-related bacteria species in soil of deciduous trees due to higher N demands than evergreen trees. In addition, the crucial factors driving soil bacterial community and N-cycling function of deciduous and evergreen trees under soil water deficit were elucidated.

2 | MATERIAL AND METHODS

2.1 Experimental design

Cunninghamia lanceolata, Pinus massoniana, Alnus cremastogyne, and Liquidambar formosana are important tree species in monocultures and mixed plantations (Wang et al., 2009; Wu et al., 2021).
The first two are coniferous evergreen trees, and the latter two are broadleaf deciduous species. Soils collected from a degraded land (0–20 cm depth) were homogenized after removing gravels, roots and plant litter. One-year-old C. lanceolata, P. massoniana, A. cremastogyne and L. formosana trees collected from a local nursery garden near the experiment site (30°19 N, 120°23 E) and 30 individuals per species were planted in plastic pots (30 cm external diameter and 21 cm height, one individual per pot) in mid-December 2018. After 5 months, we selected 16 individuals of similar size and growth performance per species (May 28, 2019). We subjected all species to two treatments (water vs. drought); soil water content was maintained at approximately 80% field capacity (approximately 27.8% soil water) in the well water treatment and 30% field capacity (approximately 10.6% soil water) in the drought treatment. The pots were weighed every day to monitor soil water content and the lost water was replenished in time. Eight replicates per treatment were used. The experiment was conducted in a greenhouse at Hangzhou Normal University, China (30°19 N, 120°23 E).

2.2 | Gas exchange measurements and harvesting

The net photosynthetic rate per treatment (eight replicates) was measured using an LI-6400 photosynthesis system (Li-Cor) under the following conditions: photosynthetic photon flux density 1500 μmol m⁻² s⁻¹, 28°C leaf temperature, 70% relative humidity, and 400 μmol mol⁻¹ CO₂. Measurements were made from July 29 to August 1, 2019. We harvested all plants on August 30, 2019, and all samples (root, stem, and leaf) were dried at 75°C for 72 h.

2.3 | Rhizosphere soil physicochemical properties

Soil adhering to root was defined as rhizosphere soil. Rhizosphere soil samples were collected by carefully removing soil from the root surface after harvesting the plants. Four soil replicates of P. massoniana in the water treatment were discarded because of technical errors during transport. Subsamples of rhizosphere soil were air-dried to measure pH, total phosphorus, available phosphorus, total N, and ammonium (NH₄⁺-N) and were determined according to Guo et al. (2019). In brief, soil-water suspension (1:2.5 w/v) was used for pH determination. Soil extractions with 50 ml 2 M KCl were used to determine NH₄⁺. Soil digested by H₂SO₄ and HClO₄ was used to measure total phosphorus through molybdenum blue colorimetry. Soil extractions with sodium bicarbonate was used to determine available phosphorus. The soil organic matter was measured according to Guo et al. (2021). Summary of soil properties is shown in Table S1. Further subsamples of fresh rhizosphere soil were stored at 4°C, and activities of β-1,4-N-acetylglucosaminidase (NAG) and β-D-glucosidase were assessed using respective ELISA kits. NAG and β-D-glucosidase are related to chitin and cellulose degradation, respectively (Jing et al., 2020).

2.4 | 16S rRNA sequencing and real-time quantitative PCR

Soil subsamples were stored at −80°C until DNA extraction. Total DNA was extracted from fresh soil (0.5 g) using a kit (Omega BioTek), and DNA quality was assessed using 1% agarose gel electrophoresis. The 16S primers 338F (5′-ACTCTACGGGAGGCAGCA-3′) and 806R (5′-GGACTACHVGGGTWTCTAA-3′) targeting the V3–V4 region were used for PCR amplification, followed by paired-end sequencing on a MiSeq platform (Illumina; Guo et al., 2021). The PCR amplification steps followed: initial denaturation (3 min) at 95°C, 27 cycles (30 s) at 95°C, annealing (30 s) at 55°C, elongation (45 s) at 72°C, and a final extension (10 min) at 72°C. PCR reactions were performed in triplicate 20 µl mixture. Raw FASTQ reads were demultiplexed, quality filtered. More details concerning the process on raw reads are shown in the supplementary materials. All bacterial sequences were assigned to operational taxonomic units (OTUs) using the SILVA database and the UPARSE pipeline at 97% similarity (Quast et al., 2013).

DNA extracts were also subjected to quantitative PCRs (qPCRs) to quantitate copy numbers of N-cycling genes: ammonia monoxygenase (amoA) genes of AOA, AOB, nifH, nirS and nirK. The following primer combinations were used: AOA forward: 5′-ATGGTCGGCTWAGAGC-3′; reverse: 5′-GCCATCCATCTGTATGTTCA-3′; AOB forward: 5′-GGAGRAAAGCAGGATCTG-3′; reverse: 5′-CTAGGTTTGTAGTTTCAAACGC-3′; nifH forward: 5′-TGCGAYCCSAARGCBGACTC-3′; reverse: 5′-ATSGCCATCAYTCRCCGGA-3′; nirS forward: 5′-AACGYSAGGARACS-3′; reverse: 5′-GASTTCGRTGSGTCTTTSAYGAA-3′; and nirK forward: 5′-TGCACATCGCCAAAGGNNATGWYGG-3′. More details concerning the process on qPCR are shown in the supplementary materials. Standard curves for each functional gene were produced according to Frey and Rieder (2013).

2.5 | Statistical analyses

We used a two-way analysis of variance (ANOVA) to identify the effects of species, drought, and their interaction terms on plant biomass, enzyme activity, and bacterial alpha-diversity. We named a factor ‘leaf habit’ to indicate deciduous and evergreen tree species in the present study in order to compare their effects combined with drought on net photosynthetic rate, enzyme activity, qPCR results, and bacterial alpha-diversity using two-way ANOVA following Tukey’s b post hoc test in case of significant interactions. Relative abundances of bacterial were analyzed at phylum and genus level between deciduous and evergreen trees with independent-samples T test. Bacterial beta-diversity was ordinated using nonmetric multidimensional scaling (NMDS) based on Bray–Curtis distances. PERMANOVA using the “adonis” function of the “vegan” software was used to find significant differences in soil bacterial community dissimilarity (Oksanen et al., 2020).
“Ward. D2” function was used to perform hierarchical clustering analysis based on Bray–Curtis distances of bacterial OTUs from all samples, and the silhouette width method in R decided the number of clusters to be used (Xiong et al., 2021). The Spearman correlations between characters of bacterial community (the relative abundance of the major bacterial phyla), soil properties and plant growth versus functions in enzyme activities and N-cycling-related genes. The multiple regression model with variance decomposition analysis was applied to estimate the importance of soil, plant, and bacterial characteristics in driving the studied enzyme activities and N-cycling-related functional genes following descriptions in Jiao et al. (2020). After checking potential collinearity, distance-based redundancy analysis (db-RDA) was also performed using ‘vegan’ package. Network analysis was conducted and visualized using Cytoscape v. 3.5 and Gephi, respectively (Bastian et al., 2009; Shannon et al., 2003). Robust Spearman correlation scores (Spearman’s |r| > .5) and statistically significant correlations were tested (p < .01). Network complexity and hub nodes in co-occurrence networks were tested as described by Xiong et al. (2021). Soil bacterial community change along the root biomass gradient was analyzed using threshold indicator taxa analysis (Baker et al., 2015). We used bootstrapping (n = 500) to estimate uncertainty around the environmental change points for each indicator taxa. Structural equation models (SEM) were used to test the effects of root biomass on each N-cycling function using the ‘lavaan’ package (Rosseel, 2012). The model was fitted according to chi-squared tests, a high comparative fit index (≥0.90), a low root means square error of approximation index (≤0.1), and a low standardized root means square residual index (≤0.1).

3 | RESULTS

3.1 | Plant performance in carbon fixation and biomass production

The largest root biomass was produced by A. cremastogyne in the water treatment (Table 1). Root biomass of L. formosana and A. cremastogyne was significantly smaller in the drought than in the water treatment, but root biomass of C. lanceolata and P. massoniana was not significantly smaller in the drought than in the water treatment (Table 1). Deciduous trees showed significantly higher net photosynthetic rates than evergreen both at well water (12.86 vs. 7.46 μmol m⁻² s⁻¹) and drought treatment (5.45 vs. 1.63 μmol m⁻² s⁻¹; Figure S1).

3.2 | Composition, diversity, and structure of bacterial community

Proteobacteria, Actinobacteria, Chloroflexi, Acidobacteria, Firmicutes, and Gemmatimonadetes were the six most abundant soil bacterial phyla. The drought treatment did not significantly

| TABLE 1 Growth characteristics of four tree species in response to two soil water levels |
|---------------------------------|----------------|----------------|
|                                 | Aboveground biomass (g) | Root biomass (g) | Total biomass (g) |
| Liquidambar formosana           |                          |                 |
| Control                         | 59.09 ± 1.77a           | 6.86 ± 0.38c    | 65.95 ± 1.64a     |
| Drought                         | 30.95 ± 1.92b           | 5.22 ± 0.29d    | 36.17 ± 1.97b     |
| Alnus cremastogyne              |                          |                 |
| Control                         | 23.70 ± 1.54c           | 12.66 ± 0.46a   | 36.35 ± 1.80b     |
| Drought                         | 14.33 ± 1.20de          | 8.39 ± 0.35b    | 22.72 ± 1.42c     |
| Cunninghamia lanceolata         |                          |                 |
| Control                         | 19.60 ± 1.64 cd         | 4.50 ± 0.29de   | 24.10 ± 1.87c     |
| Drought                         | 12.13 ± 1.47e           | 3.02 ± 0.29e    | 15.15 ± 1.70d     |
| Pinus massoniana                |                          |                 |
| Control                         | 4.32 ± 0.55f            | 0.56 ± 0.05f    | 4.87 ± 0.54e      |
| Drought                         | 4.04 ± 0.69f            | 0.49 ± 0.07f    | 4.53 ± 0.68e      |
| Significant effects             |                          |                 |
| Species                         | (243.616)***            | (303.968)***    | (255.620)***      |
| Drought                         | (105.581)***            | (62.706)***     | (122.493)***      |
| Species × drought               | (29.325)***             | (13.596)***     | (26.346)***       |

Note: A two-way analysis of variance was used to test the effect of plant species, drought, and their interactions on growth characteristics. Different lowercase letters in the column indicate significant differences. Significant effects of factors and interactions (x) are indicated. Numbers in brackets indicate F values. Abbreviation: NS, not significant. ***p ≤ .001.
affect the relative abundances of each phylum (Figure 1a). However, Actinobacteria and Firmicutes in the soil of deciduous trees showed higher relative abundances, whereas those of Proteobacteria, Acidobacteria, and Gemmatimonadetes were lower in deciduous than in coniferous tree soils (Figure 1a). The dominant bacterial taxa at genus level were also significantly different between deciduous and coniferous tree soils (Figure 1b). For example, the relative abundance of Arthrobacter and Bacillus in deciduous tree soils were significantly higher than in coniferous.

Shannon diversity in the L. formosana soil community after the drought was significantly higher than under well-watered conditions (Figure 2a). Shannon diversity of C. lanceolata and P. massoniana soil communities tended to be higher than those of communities associated with L. formosana and A. cremastogynge. Soil bacterial alpha-diversity in the soil of deciduous species was significantly increased by drought, whereas that in the soil of coniferous species did not differ significantly between water and drought treatments (Figure 2b). Plants significantly affected richness of bacterial community (Figure S2). Regarding bacterial beta-diversity, clustering and NMDS analyses showed a clear separation between broadleaf deciduous and coniferous tree soil communities (Figure 2c,d).

Drought and species exerted specific effects on network traits, such as clustering coefficients, positive/negative relationships, modularity, and hub node number (Figure 3; Figure S2; Table 2). Regarding plant leaf habit, drought declined positive relationships, average degree, clustering coefficients, and hub nodes but increased negative relationships and path distance. Modularity was reduced in broadleaf deciduous trees but increased in coniferous evergreen trees in the drought treatment (Table 2; Figure S3).

### 3.3 Nitrogen-cycling functions

Soil β-D-glucosidase in P. massoniana and NAG in C. lanceolata showed the highest activity in the water treatment (Figure S4a,b), and NAG activity was significantly higher in the soil of evergreen than in that of deciduous trees (Figure S4). Soil AOA, AOB, nifH, and nirS copy numbers were significantly higher in the soil of deciduous trees than in evergreen (Figure 4). The ratio of AOA to AOB was also higher in the soil of deciduous trees (Figure 4f). These functional genes were less affected by drought than by plant leaf habit, and significant interaction effects of the two factors were found on the denitrification-related genes nirS and nirK. Copy numbers of nirS in the soil of deciduous species were significantly higher in the drought than in the water treatment (Figure 4d), whereas in evergreen, nirK copy numbers were significantly lower in the drought than in the water treatment (Figure 4e).

### 3.4 Effects of root on bacterial community and N-cycling functions

Root biomass was a key factor in predicting diverse functions of soil microbial community (Figure 5). NAG activity, copy numbers of nifH, AOB, AOA, nirK and nirS was positively related with root biomass,
and abundance of Actinobacteria and Firmicutes, while negatively related with the abundance of Gemmatimonadetes (Figure 5). Root biomass and soil properties explained 89.77% of the variation in bacterial communities, and RDA axis 1 explained 79.92%, with root biomass as the main factor affecting the variation of this axis (Figure S5).

Threshold indicator taxa analysis (TITAN) identified 44 bacteria genera with a decreasing response along the root biomass gradient and 46 taxa showed an increasing response (Figure 6). The negative responding bacteria taxa declined sharply at small root biomass (2.265 g), while the positive responding bacteria taxa increased at high root biomass (8.885 g; Figure S6). The most sensitive negative responding bacteria taxon (lowest change point value) was *Pseudoduganella* while the most sensitive positive responding bacteria taxon (highest change point value) was *Chitinimonas* (Figure 6).

The SEM explained 50.2%, 47.6%, 48.6%, 49.4%, and 37.3% of the variability in copy numbers of *nifH*, AOB, AOA, *nirK*, and *nirS*, respectively (Figure 7). Root biomass negatively affected bacterial Shannon diversity, but the effect was not statistically significant. However, root biomass exerted significant positive effects on copy numbers of all N-cycle functional genes (Figure 7); the correlation coefficients of *nifH*, AOB, AOA, *nirK*, and *nirS* were 0.48, 0.50, 0.55, 0.59, and 0.60, respectively.

**FIGURE 2** Bacterial alpha- and beta-diversity in the soil of two deciduous (*Alnus cremastogyne* and *Liquidambar formosana*) and two evergreen (*Cunninghamia lanceolata* and *Pinus massoniana*) tree species in response to two water regimes. (a) Bacterial Shannon diversity of each species at operational taxonomic unit level, (b) Bacterial Shannon diversity of both deciduous and evergreen species combined. A two-way analysis of variance was used. Post hoc tests were used to test differences among treatments with Tukey’s b tests. Different letters indicate significant differences. (c) Hierarchical clustering analysis and (d) nonmetric multidimensional scaling (NMDS) based on Bray–Curtis distances

**FIGURE 3** Co-occurrence network of the four tree species and two plant leaf habits in the control (water) and drought treatments. *Cunninghamia lanceolata* and *Pinus massoniana* belong to the evergreen species, *Alnus cremastogyne* and *Liquidambar formosana* belong to the deciduous species.
TABLe 2 Traits of bacterial co-occurrence networks of four tree species and two plant leaf habits in response to control (water) and drought treatments

| Treatment  | Positive edge | Negative edge | Average degree | Modularity | Average clustering coefficient | Average path distance | Hub node |
|------------|---------------|---------------|----------------|------------|-------------------------------|----------------------|---------|
| Leaf habit                                                                                     |
| Broadleaf deciduous                           | 7547          | 20            | 51.128        | 0.440      | 0.592                         | 2.146                | 123     |
| Drought                                            | 5726          | 72            | 39.044        | 0.348      | 0.525                         | 2.408                | 77      |
| Coniferous evergreen                           | 8304          | 119           | 56.341        | 0.343      | 0.607                         | 2.204                | 129     |
| Drought                                            | 4502          | 150           | 31.327        | 0.416      | 0.482                         | 2.631                | 47      |
| Species                                                                                       |
| Liquidambar formosana                           | 2283          | 14            | 15.896        | 0.512      | 0.472                         | 3.444                | 0       |
| Drought                                            | 2438          | 107           | 17.196        | 0.434      | 0.496                         | 3.668                | 1       |
| Alnus cremastogyne                              | 4002          | 1             | 27.311        | 0.492      | 0.586                         | 3.123                | 50      |
| Drought                                            | 1706          | 65            | 12.621        | 0.589      | 0.479                         | 3.888                | 0       |
| Cunninghamia lanceolata                         | 2017          | 105           | 14.937        | 0.558      | 0.478                         | 3.487                | 0       |
| Drought                                            | 1388          | 68            | 10.189        | 0.634      | 0.431                         | 4.065                | 0       |
| Pinus massoniana                                | 4273          | 83            | 30.455        | 0.370      | 0.561                         | 3.124                | 56      |
| Drought                                            | 1146          | 178           | 9.355         | 0.614      | 0.416                         | 4.391                | 0       |

4 | DISCUSSION

Soil microbiomes could play essential roles in forest ecosystem responses to increasing anthropogenic global climate changes. We tested the individual and interactive effects of plant leaf habit and drought on soil bacterial community structure and N cycling function. Our results showed that bacterial community and N cycling function were interactively affected by plant leaf habit and drought.

4.1 | Effects of plant and drought on bacterial community

The drought-induced changes in diversity and taxonomic composition of soil bacterial community depend on water availability, soil physicochemical properties and plant physiological responses (Bastida et al., 2017; Han et al., 2021; Naylor & Coleman-Derr, 2018). Drought significantly reduced SOM, TN, and NH₄⁺ of rhizosphere soils (Table S1) and induced changes in the abundance, diversity, and network structure of the bacterial community in this study (Figures 1-3; Figure S2). Nutrient competition among microbes and osmotic stress affects bacteria species (Chodak et al., 2015; Xie et al., 2021). The bacterial genera, for example, Arthrobacter and Bacillus, were reduced by drought, suggesting drought-sensitive species belonging to the two genera were strongly inhibited by the declining water availability (Chodak et al., 2015; Xie et al., 2021). Plants also impose strong selective pressure on the bacterial community by recruiting species-specific microbes in response to water deficit (Ehlers et al., 2020; Guo et al., 2019, 2021). In this study, co-occurrence network results revealed a better connected and more stable bacterial network of rhizosphere soil surrounding L. formosana compared to A. cremastogyne with a greater decline in positive relationships, average degrees, and clustering coefficients but an increase in average path lengths after drought (Figure 3). Mutualistic relationships in the plant-soil continuum have been reported between bacterial communities and plants considering resistance to drought (Naylor & Coleman-Derr, 2018). The more stable and connected network of bacterial community can be less affected by drought (Jiao et al., 2020).

4.2 | Crucial role of root biomass in shaping soil bacterial community and function

Our results provided convincing evidences that the plant leaf habit imposed stronger impacts on bacterial community than short-term drought. Greater variations happened in the relative abundance of dominant bacteria, richness and diversity (alpha- and beta), and N-cycling function between deciduous and evergreen plants than drought (Figures 1, 2, and 4). Plant leaf habit had no significant effects on soil physiochemical traits (Table S1), which demonstrated an important role of plant characteristics in shaping the divergent bacterial community. The two deciduous species recruited higher relative abundance of plant growth-promoting bacteria genera Arthrobacter and Bacillus than the two evergreen species (Figure 1). The random forest analysis and db-RDA results confirmed the important role of plant characteristics, particularly root biomass, in driving bacterial structure and functions (Figure 5; Figure S5). Further analysis identified by TITAN showed that different indicators of bacterial taxa at the genus level along the smaller to larger root biomass gradient from evergreen to...
deciduous species (Figure 6; Figure S6). The hub nodes from the co-occurrence network were considered as potential keystone taxa in microbial community (Jiao et al., 2020). The changing bacterial taxa indicators affected by the root mass were probably the main reasons in driving changes in keystone taxa, connection, and modularity of the network (Figure 3; Figure S3).

We found that root biomass was a crucial factor in determining the composition and structure of the soil bacterial community. Effects of deciduous and evergreen trees on soil bacterial communities can be attributed to differences in plant photosynthetic carbon supply as well as to the varying amounts of carbon released to the soil through root exudates (Guo et al., 2019, 2021; Keller et al., 2021; Zhalnina et al., 2018). Analogous to the leaf economic spectrum aboveground (Donovan et al., 2011; Wright et al., 2004), the root of deciduous and coniferous species ranges from more competitive and fast-growing traits to more conservative slow-growing traits (Sun et al., 2021). Deciduous trees with higher growth rates and biomass production show higher root exudation rates and release more root-derived carbon to the rhizosphere compared to evergreen trees (Emmett et al., 2020; Sun et al., 2021; Wang et al., 2021). Our results were consistent with previous studies showing that deciduous trees typically have a higher net photosynthetic rate than evergreen trees (Balodocchi et al., 2010; di Francescantonio et al., 2020; Wang et al., 2021), which implies a higher quantity of root-derived carbon released into rhizosphere soil.

4.3 Deciduous trees accelerate soil N cycle

We hypothesized that deciduous trees would elicit N cycling more strongly by promoting N-cycling-related bacteria, compared to evergreen; significantly higher copy numbers of N-cycle-related genes in the soil of deciduous species compared to that of
evergreen supported this hypothesis. Plant–soil microbe interactions in the rhizosphere are a predominant link in N cycling and supply (Moreau et al., 2019). Plants seem to be able to control N transformations mediated by diverse soil microbes in and near the rhizosphere (Bardgett et al., 2014; Galland et al., 2019; Moreau et al., 2019; Mushinski et al., 2020). Random forest analysis identified root biomass as having a key role in affecting N-cycling functions (Figure 5). All studied N-cycling genes were positively correlated with the relative abundance of Actinobacteria and Firmicutes but negatively correlated with Gemmatimonadetes (Figure 5). The SEM also demonstrated strong positive effects of root biomass on N-cycling genes (Figure 7). Higher N demand of deciduous trees requires more N input to meet plant biomass accumulation by increasing the abundance of nifH. Soil nitrate (NO$_3^-$) is the major plant-available form of N, and higher AOB and AOA levels accelerate nitrification. However, the denitrification-related bacteria showed higher gene copies. Cantarel et al. (2015) observed that denitrification depends on the relative growth rate of plants. Higher nirK and nirS in the rhizosphere of deciduous species can stimulate transformation from NO$_3^-$ to nitrous oxide and increase N loss, which may imply that bacteria outcompete plants regarding the use of NO$_3^-$ resources (Moreau et al., 2015). In addition, denitrification inhibitors or stimulators through the release of organic compounds have been reported (Bardon et al., 2016; Moreau et al., 2019). However, the exact mechanisms underlying the simulation of denitrification in the plant rhizosphere remain to be elucidated.

The four tree species studied in this study are widely used to construct artificial plantations (Wang et al., 2009; Wu et al., 2021). Our results have important implications for our understanding of plants control over bacterial community and N-cycling function in artificial forest ecosystems. A recent study across nine European countries assessed responses of mixed and monoculture forests to drought events between 1975 and 2015 (Pardos et al., 2021), which showed that drought resistance of mixed conifer-deciduous broadleaf forests exceeded that of mixed broadleaf forests; this suggests an important role of plant leaf habit diversity in mixed forests (Figure S7). Our simple framework showed that
different tree species with different roots (biomass, architecture, or exudates) recruited more diverse microbes to influence forest productivity (Figure S7). The diversity of mycorrhizal fungi associating with roots of different plant leaf habits contributes to niche separation with respect to acquisition and transport of soil N and water, which suggests reduced competition for these resources due to symbiotic mycorrhiza. Roots of N$_2$-fixing plants colonized by mycorrhiza show increased N fixation and drought resistance (Liu et al., 2020), and N$_2$-fixing plants increase soil N availability and promote the growth of adjacent plants (Chen et al., 2021; Minucci et al., 2019; Tang et al., 2019). Mixed forests greatly increase soil N availability compared to monocultures, by promoting N mineralization, increasing soil N retention, or reducing denitrification (Chen et al., 2021; Wang et al., 2021).

**CONCLUSIONS**

Overall, declining soil water availability and plant leaf habit impacted the bacterial community. However, stronger effects of deciduous and evergreen tree species on bacterial community and N-cycling function than short-term drought had been proved in this study. Deciduous and evergreen tree species showed different plant traits in carbon fixation and growth. As a linkage between above and belowground, root biomass played key roles in affecting bacterial community structure and soil N cycling. However, our research is constrained by low sample size of deciduous and evergreen tree species. Further research on N-cycling microbes affected by plant functions and plant–soil interactions is required to predict the respective consequences of climate change.

**FIGURE 6** Threshold Indicator Taxa Analysis of soil bacterial community at genus level in responding to a root biomass gradient. Only taxa with pure (purity ≥ 0.95) and reliable (reliability ≥ 0.95) responses are plotted. Bootstrapping-based ($n = 500$) probability density functions of negative responding bacteria taxa (blue) (a) and positively responding bacteria taxa (red) (b) along the root biomass gradient.

**FIGURE 7** Structural equation models to assess relationships and effects of root biomass on soil alpha-diversity and N-cycling-related genes (a–c). AOA, ammonia-oxidizing archaea; AOB, ammonia-oxidizing bacteria.
AUTHOR CONTRIBUTIONS
Jiantong Liu: Formal analysis (equal); visualization (equal); writing – original draft (equal). Xinyu Wang: Data curation (equal); software (equal); writing – original draft (equal). Lin Liu: Data curation (equal); software (equal); visualization (equal). Xuefeng Wu: Visualization (equal). Zhichao Xia: Funding acquisition (supporting); writing – review and editing (equal). Qingxue Guo: Data curation (equal); funding acquisition (lead); methodology (lead); project administration (lead); writing – original draft (equal).

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CONFLICT OF INTEREST
The authors declare that they have no conflict of interest.

DATA AVAILABILITY STATEMENT
All bacterial data are available at the National Center for Biotechnology Information under BioProject ID PRJNA821928. Data of soil properties and plant biomass were input in Dryad. https://doi.org/10.5061/dryad.2547d7wqs.

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REFERENCES

Baker, M.E., King, R.S., & Kahle, D. (2015). TITAN2: Threshold indicator taxa analysis. R package version 2.1. https://CRAN.R-project.org/package=TITAN2

Baldocchi, D.D., Ma, S., Rambal, S., Misson, L., Ourcival, J.M., Limousin, J.M., Pereira, J., & Papale, D. (2010). On the differential advantages of evergreenness and deciduousness in Mediterranean oak woodlands: A flux perspective. Ecological Applications, 20, 1583-1597. https://doi.org/10.1890/08-20471

Bardgett, R.D., Mommer, L., & De Vries, F. (2014). Going underground: Root traits as drivers of ecosystem processes. Trends in Ecology & Evolution, 29, 692–699. https://doi.org/10.1016/j.tree.2014.10.006

Bardon, C., Poiy, F., Piola, F., Pancton, M., Comte, G., Meiffren, G., & Haichar, F. Z. (2016). Mechanism of biological denitrification inhibition: Procyandins induce an allosteric transition of the membrane-bound nitrate reductase through membrane alteration. FEBS Microbiology Ecology, 92, fiw034. https://doi.org/10.1038/femsec/fiw034

Bastian, M., Heymann, S., & Jacomy, M. (2009). Gephi: An open source software for exploring and manipulating networks. ICWSM, 8, 361–362. https://doi.org/10.13140/2.1.1341.1520

Bastida, F., Torres, I., Andrés-Abellán, M., Baldrian, P., López-Mondéjar, R., Větrovský, T., Richnow, H.H., Starke, R., Ondofo, S., Garcia, C., López-Serrano, F. R., & Jehmlich, N. (2017). Differential sensitivity of total and active soil microbial communities to drought and forest management. Global Change Biology, 24, 155. https://doi.org/10.1111/gcb.13790

Bowen, H., Maul, J.E., Cavigelli, M.A., & Yarwood, S. (2020). Denitrifier abundance and community composition linked to denitrification activity in an agricultural and wetland soil. Applied Soil Ecology, 151, 103521.

Cantarel, A. A., Pommer, T., Desclos-Theveniau, M., Diquelou, S., Dumont, M., Grassein, F., Kasti, E. M., Grigulis, K., Lainé, P., Lavorel, S., Lemauviel-Lavenant, S., Personeni, E., Schloter, M., & Poly, F. (2015). Using plant traits to explain plant-microbe relationships involved in nitrogen acquisition. Ecology, 96, 788–799. https://doi.org/10.1890/13-2107.1

Castro, S. P., Cleland, E. E., Wagner, R., Sawad, R. A., & Lipson, D. A. (2019). Soil microbial responses to drought and exotic plants shift carbon metabolism. The ISME Journal, 13, 1776–1787. https://doi.org/10.1038/s41396-019-0389-9

Chen, X., Chen, H. Y. H., Searle, E. B., Chen, C., & Reich, P. B. (2021). Negative to positive shifts in diversity effects on soil nitrogen over time. Nature Sustainability, 4, 225–232. https://doi.org/10.1038/s41371-020-0418-y

Chodak, M., Gołębiewski, M., Morawska-Płoskonka, J., Kuduk, K., & Niklińska, M. (2015). Soil chemical properties affect the reaction of forest soil bacteria to drought and rewetting stress. Annales de Microbiologie, 65, 1627–1637. https://doi.org/10.1007/s11231-014-1002-0

Chorrea, D., Villagra, M., Goldstein, G., & Campanello, P. I. (2020). Drought and frost resistance vary between evergreen and deciduous Atlantic forest canopy trees. Functional Plant Biology, 47, 779–791. https://doi.org/10.1071/FP19282

Donovan, L. A., Maherali, H., Caruso, C. M., Huber, H., & de Kroon, H. (2011). The evolution of the worldwide leaf economics spectrum. Trends in Ecology & Evolution, 26, 88–95. https://doi.org/10.1016/j.tree.2010.11.011

Ehlers, B. K., Berg, M. P., Staudt, M., Holmstrup, M., Glasius, M., Ellers, J., Tomio, S., Madsen, R. B., Slotsbo, S., & Penuelas, J. (2020). Plant secondary compounds in soil and their role in belowground species interactions. Trends in Ecology & Evolution, 35, 716–730. https://doi.org/10.1016/j.tree.2020.04.001

Emmett, B. A., Buckley, D. H. R., & Drinkwater, L. E. (2020). Plant growth rate and nitrogen uptake shape rhizosphere bacterial community composition and activity in an agricultural field. The New Phytologist, 225, 960–973. https://doi.org/10.1111/nph.16171

Farías, F., Orlando, J., Bravo, L., Guevara, R., & Carú, M. (2009). Comparison of soil bacterial communities associated with actinorhizal, non-actinorhizal plants and the interspaces in the sclerophyllous matorral from Central Chile in two different seasons. Journal of Arid Environments, 73, 1117–1124. https://doi.org/10.1016/j.jaridenv.2009.06.010

Frey, B., & Rieder, S. R. (2013). Response of forest soil bacterial communities to mercury chloride application. Soil Biology and Biochemistry, 65, 329–337. https://doi.org/10.1016/j.soilbio.2013.06.001

Galland, W., Piola, F., Burlet, A., Mathieu, C., Nardy, M., Poussineau, S., Blazère, L., Gervaix, J., Pujialon, S., Simon, L., & Haichar, F. Z. (2019). Biological denitrification inhibition (BDI) in the field: A strategy to improve plant nutrition and growth. Soil Biology and Biochemistry, 136, 107513. https://doi.org/10.1016/j.soilbio.2019.06.009

Gillespie, L. M., Fromin, N., Milcu, A., Buatois, B., Pontoizeau, C., & Hättenschwiler, S. (2020). Higher tree diversity increases soil microbial resistance to drought. Communications Biology, 3, 377. https://doi.org/10.1038/s42003-020-1112-0

Gray, J. T., & Schlesinger, W. H. (1983). Nutrient use by evergreen and deciduous shrubs in Southern California. II. Experimental investigations of the relationship between growth, nitrogen uptake, and nitrogen availability. Journal of Ecology, 71, 43–56.

Guo, Q. X., Liu, J. T., Yu, L., Korpelainen, H., & Li, C. Y. (2021). Different sexual impacts of dioecious Populus euphratica on microbial communities and nitrogen cycle processes in natural forests. Forest Ecology and Management, 496, 119403. https://doi.org/10.1016/j.foreco.2021.119403
Xie, J., Dawwam, G. E., Sehim, A. E., Li, X., Wu, J., Chen, S., & Zhang, D. (2021). Drought stress triggers shifts in the root microbial community and alters functional categories in the microbial gene pool. *Frontiers in Microbiology*, 12, 744897. [https://doi.org/10.3389/fmicb.2021.744897](https://doi.org/10.3389/fmicb.2021.744897)

Xiong, C., Zhu, Y. G., Wang, J. T., Singh, B., Han, L. L., Shen, J. P., Li, P. P., Wang, G. B., Wu, C. F., Ge, A. H., Zhang, L. M., & He, J. Z. (2021). Host selection shapes crop microbiome assembly and network complexity. *The New Phytologist*, 229, 1091-1104. [https://doi.org/10.1111/nph.16890](https://doi.org/10.1111/nph.16890)

Zhalnina, K., Louie, K. B., Hao, Z., Mansoori, N., da Rocha, U. N., Shi, S. J., Cho, H. J., Karaoz, U., Loque, D., Bowen, B. P., Firestone, M. K., Northen, T. R., & Brodie, E. L. (2018). Dynamic root exudate chemistry and microbial substrate preferences drive patterns in rhizosphere microbial community assembly. *Nature Microbiology*, 3, 470-480. [https://doi.org/10.1038/s41564-018-0129-3](https://doi.org/10.1038/s41564-018-0129-3)

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