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Microbial communities of *Schisandra sphenanthera* Rehd. et Wils. and the correlations between microbial community and the active secondary metabolites

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Background: *Schisandra sphenanthera* Rehd. et Wils. is a kind of Traditional Chinese Medicine (TCM) which can protect the kidney and liver. However, great differences exist in the content of active secondary metabolites in different parts of *S. sphenanthera*. Do microorganisms critically influence the accumulation of active components in different parts of *S. sphenanthera*?

Methods: In this study, 16S/ITS amplicon sequencing analysis was applied to unravel microbial communities in rhizospheric soil and different parts of wild *S. sphenanthera*. The active secondary metabolites were measured in different parts, and the correlation with microorganisms was estimated.

Results: The contents of essential oil components and lignans in fruit were much higher than that in stem and leaf, and the dominant essential oil components in different parts were different. The microbial amplicon sequences were taxonomically grouped into 8 (bacteria) and 7 (fungi) different phyla. Community diversity and composition analyses showed that different parts of *S. sphenanthera* had similar and unique microbial communities, and functional prediction analysis showed that host-related functions are associated with metabolic processes. Results showed that the accumulation of secondary metabolites in *S. sphenanthera* was closely related to the microbial community composition, especially bacteria. Our results provided a new opportunity to further understand the effects of microorganisms on the active secondary metabolites and provided a basis for further research on the sustainable utilization of *S. sphenanthera*. 
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ABSTRACT

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generated a basis for further research on the sustainable utilization of *S. sphenanthera*.

**Keywords** Plant microbiome, Rhizospheric soil microorganism, Endophytes, Active secondary

**Introduction**

The relationship between microorganisms and plants or animals is one of the most studied
research areas in biology or microbiology in recent years, including studies of human gut
microfauna (Liu et al. 2021) and plant microbiome (Bai et al. 2015). In most cases,
interactions maintain a close mutualistic relationship with animals and plants, through which
they provide nutrients, and the related microbial community can play an important role in the
immune system of animals and plants (Haasquard et al. 2015). Plant-microbe interactions are
important to better understand their role in plant growth and development (Verma et al. 2022).
Within plant microbiome research, most attention has been dedicated to endophyte and
rhizospheric soil microorganisms (Bai et al. 2015; Zhu et al. 2020). Differentiation of
rhizospheric soil microorganisms and endophytes in plants has been reported using high-
throughput sequencing methods (Sun et al. 2022).

Microorganisms that colonize plant organs and do not cause obvious plant diseases are
depicted as endophytes (Shao et al. 2023). The diversity and community structures of endophytes are closely
related to the species, growth stage, different parts, living environment, and genotype of the host
plants (Xia et al. 2023). In addition, endophytes can produce antibiotics, enzymes, plant growth
regulators, alkaloids, and a series of metabolites (Shao et al. 2023). The rhizosphere environment
is an important place for plant growth, metabolism, and absorption of soil nutrients, as well as
the most direct interaction between roots and soil (Li et al. 2023). Rhizospheric soil contains a
large number of rhizosphere microorganisms, whose quantity and diversity are significantly
higher than that of bulk soil (Yuan et al. 2022). Rhizospheric soil microorganisms can increase
plant tolerance to biotic and abiotic stresses, improve soil nutrient absorption, and affect plant
yield and quality (Vries et al. 2020). At the same time, rhizospheric soil microorganisms and
plant endophytes interact with each other (He et al. 2021), and rhizospheric soil microorganisms
directly affect soil biochemical activities (Huong et al. 2022).

The active secondary metabolites in most medicinal plants are affected by environmental,
genetic, and other factors. Endophytes are an important part of the internal environment of
medicinal plants, and rhizospheric soil microorganisms are a significant part of the external
environment of plants (Korenblum et al. 2022; Hou et al. 2022). They can increase the active
components of medicinal plants by producing products that are the same or similar to the active
secondary metabolites of medicinal plants, or by transforming the original active secondary
metabolites of medicinal plants into new compounds to increase the types of active components (Korenblum et al. 2022; Hou et al. 2022). Therefore, the study of endophytes and rhizospheric soil microorganisms of medicinal plants, as well as the influence of microorganisms on active secondary metabolites, has become an important content in the production of Chinese medicine and the development of new drugs.

_Schisandra sphenanthera_ Rehd. et Wils., a perennial deciduous woody vine of the genus _Schisandra_, which is a high-value Traditional Chinese Medicine (TCM) (Editorial Committee of Flora of China, 1986; Smith, 1947; The State Pharmacopoeia Commission of P. R. China, 2020). The dried and ripe fruit is often used as medicine, known as "Nanwuweiizi", to treat chronic cough, asthma, night sweats, and palpitations insomnia (Li et al. 2020; The State Pharmacopoeia Commission of P. R. China, 2020). The active ingredients in fruits include lignans, essential oils, polysaccharides, and so on (Gu et al. 2008; Lu et al. 2012; Wang et al. 2018). In addition, the fruits of _S. sphenanthera_ can also be used in cosmetics and health products, and listed as a functional food by the Ministry of Health of the P. R. China (Huang et al. 2021). Since its main source is wild resources, the genetic improvement and extensive commercial use of _S. sphenanthera_ are limited by the depletion of natural herbal resources. We know from the literature that the specific endophytes isolated from _S. sphenanthera_ by You et al. can promote the growth of the host (You et al. 2021). However, up to now, minimal studies are available that involve the _S. sphenanthera_ endophytes, both domestically and abroad.

In the present study, we focused on two main questions: (i) How variables are rhizospheric soil microorganisms and endophytes in different parts of _S. sphenanthera_? (ii) Whether the microbial communities influence the accumulation of active secondary metabolites? Accordingly, this study evaluates the diversity and composition of microorganisms in the rhizospheric soil, root, stem, leaf, and fruit of wild _S. sphenanthera_ by high-throughput 16s rRNA and ITS amplicon sequencing. Furthermore, the correlation between the active secondary metabolites and the microorganisms of _S. sphenanthera_ is also investigated. The purpose of this study is to provide a reference for the exploitation of microorganisms and the improvement of the yield and quality of medicinal plants by analyzing the interaction between the active secondary metabolites and the microorganisms of _S. sphenanthera_.

**Materials & Methods**

**Sampling and sample processing**

Healthy plant materials (_S. sphenanthera_) were collected from Zhashui County, Shaanxi province of China, south of the Qinling Mountains, in August 2019. Rhizospheric soil, root, stem, leaf, and fruit of five individual plants lying approximately 10 m apart were sampled using sterile
tools. The roots of similar thickness (about 2 mm) were collected at a depth of 10-20 cm below the ground, without damaging the taproot. The rhizospheric soil was soil particles adhered to the root system. Particles separated during the shaking of the roots were collected and then sifted through a 2 mm sieve. For stem and leaf, one complete branch was randomly collected from each plant, and all leaves were collected from the sampled offshoot. And fruits were collected from each individual. The plant materials were transported to the laboratory with the sterile bag in ice boxes, and stored at -80 °C. All voucher specimens of plant materials were kept in the College of Life Sciences, Shaanxi Normal University (Voucher number: SN-ZS-YP-ZJW 001-003).

Part of the sample was air-dried in the shade at room temperature, while the other part was cleared and sterilized from epiphytic bacteria (surface sterilization) according to the following methods (Liu et al. 2023). 5 g of root, stem, leaf, and fruit were divided into small pieces with a sterile scalpel, and soaked in 75 % ethanol for 3 min. Then, the surface-sterilized samples were washed 3-5 times with sterile water to remove excess ethanol. To evaluate the efficiency of the surface sterilization, 100 μl of the last washed distilled water was plated on trypsin soybean agar (TSA) and potato dextrose agar (PDA) plates, and plates were incubated at 28 °C and 35 °C, respectively. Finally, samples corresponding to plates without bacterial and fungal growth were used for sequencing.

**DNA extraction, PCR amplification, high-throughput sequencing, and sequencing data processing**

The total genomic DNA of samples (root, stem, leaf, and fruit) was extracted using the cetyltrimethylammonium bromide (CTAB) method. And the DNA of rhizospheric soil microorganisms was extracted using Soil Kit (Thermo Scientific) following the manufacturer’s instructions. Primers designed according to conserved regions (bacterial 16S rRNA gene primer (V4) and fungal ITS1-5F rRNA gene primer) were used for community analysis of bacteria and fungi (Table S1). PCR reactions were performed with 10 ng of template DNA, 2 μM of primers, and 15 μL of Phusion High Fidelity PCR Master Mix (New England Biolabs). PCR was initiated at 98°C for 1 min, followed by 30 cycles (98°C for 10 s, 50°C for 30 s, 72°C for 30 s), and finally extended at 72°C for 5 min. After the detection of PCR products with 2% agarose gel, PCR products were purified, quantified, and homogenized to form the microbial sequencing library.

The purified amplicon libraries were pooled in equal concentration and sequenced with a Miseq (Illumina, USA) system following the manipulation instructions at SAGENE Guangzhou (China). Sequences with ≥97% similarity were assigned to the same OTUs and were analyzed by Uparse software (Uparse v7.0.100). The SILVA database was used based on the Mothur method.
to annotate OTUs sequences. To investigate the phylogenetic relationships of different OTUs, multiple sequence alignment was performed using MUSCLE software (Version 3.8.31). Subsequent analyses of alpha diversity and beta diversity were performed after the normalization of OTUs abundance information.

**Extraction of essential oils and component identification**

As too many root samples would cause damage to plants, the essential oil components were not extracted and analyzed from root samples. The dried stem, leaf, and fruit were comminuted to dried powders separately in a pharmaceutical disintegrator and then used to extract essential oil after 100 meshes sieve. The dried stem, leaf, and fruit were extracted with petroleum ether at a designed time, temperature, power, and solid-liquid ratio through the ultrasonic-assisted extraction method. The supernatant was collected by centrifugal extraction solution (1200 rpm for 3 min). Essential oils were analyzed on a Thermo GC-MS Trace U3000 system using a chromatographic column, TG-5MS (30 m × 0.25 mm × 0.25 μm). The oven column heating procedure had been slightly modified to maintain the initial temperature of 50 °C for two minutes and then increased to 180 °C at a rate of 6 °C/min (Wang et al. 2018). The mass spectra were obtained by automatic scanning at m/z 35–550 amu after injection of 1.0 μL sample. Components were identified based on matching their recorded mass spectra to the main library database, and their relative concentrations were calculated by comparing their GC peak areas to the total area.

The extraction of fruit essential oil was carried out according to the optimal extraction method (the solid-liquid ratio was 1:10, 30 °C, 240 W ultrasonic for 30 min) obtained from the orthogonal optimization experiment conducted in the laboratory earlier (Wang et al. 2018). The orthogonal L_9(3)^4 design was used to extract essential oils from the stem and leaf, and the optimum extraction process was studied. Four factors with three variation levels were listed in Table S2. The experimental conditions and extraction rates for each test and the results of the analysis of variance were presented in Table S3 and Table S4. The yield of essential oil (%) was calculated by dividing the essential oil content by the weight of the dried pretreated sample weight. The contents of essential oils extracted from the stem, leaf, and fruit by the optimum extracting technology reached 4.87 %, 4.42 %, and 6.33 %, which were the average value of the triplicate experiments.

**Statistical analysis**

Alpha diversity (Chao1 index, ACE index, Shannon, and Simpson index) in our samples was calculated with QIIME (v1.9.1), which was used to analyze the complexity of species diversity of rhizospheric soil and four parts (root, stem, leaf, and fruit). Chao1 and ACE were used to
evaluate the microbial species richness. Shannon and Simpson were used to evaluate the community diversity of microflora. Beta diversity analysis was also calculated with QIIME. Principal component analysis (PCoA) and hierarchical cluster analysis were performed to evaluate differences of rhizospheric soil and four parts (root, stem, leaf, and fruit) in species complexity. The obtained OTUs classification information was used to plot the structure and composition histogram of each sample and the visual heatmap. Finally, the metabolic and ecologically relevant functions of endophyte and rhizospheric soil microorganisms were annotated by Tax4Fun for 16S rDNA OTUs and FunGuild for ITS OTUs. Spearman correlation analysis was mainly used to reflect the relationship between the active secondary metabolites of the sample and the changes in the relative abundance of microorganisms (Top20) in different plant organs. All statistical analyses were carried out using the R language.

Results

OTU analysis and the diversity of bacteria and fungi

To understand the microbial taxa distribution among plant organs (root, stem, leaf, and fruit), and rhizospheric soil, we analyzed the microbiomes by 16S/ITS amplicon sequencing analysis. The total OTUs of bacteria were significantly different from that of fungi, especially in rhizospheric soil (total OTUs of bacteria are 17 times more than fungi) (Fig. S1a). OTUs cluster analysis showed that the microorganisms in the above-ground parts (stem, leaf, and fruit) of *S. sphenanthera* were grouped into one group, and the microorganisms in the below-ground parts (rhizospheric soil and root) into another category (Fig. S1b, c). The Venn diagram results showed that there were differences in composition and genetic relationships among rhizospheric soil and different parts of *S. sphenanthera* (Fig. S1d, e). There were 10 common OTUs of bacteria in rhizospheric soil and four parts (root, stem, leaf, and fruit) of *S. sphenanthera* (Fig. S1d), which belonged to three phyla (Proteobacteria, Actinobacteria, and Cyanobacteria) respectively. Among the fungi of *S. sphenanthera*, there were 6 common OTUs in rhizospheric soil and four parts (Fig. S1e) belonging to two phyla (Ascomycota and unclassified_Fungi).

Alpha diversity indices were applied for the estimating of endophytic community complexity, on the basis of the OTUs sequence, the diversity of the community was further reflected (Table 1). Among the endophytic and rhizosphere bacteria identified, the diversity of the below-ground parts was higher than the above-ground parts, and the soil Shannon index was nearly two times as high as that of the endophytes index. It indicated that the abundance of microbial species in the rhizospheric soil was high and distributed evenly. In the identified fungal microbial communities, the Simpson index was above 0.9, and the Shannon indexes in the stem and leaf were higher, and the microbial diversity of the fungi was lower than that of the bacteria (Table 1).
Chao1 also revealed that the diversity of root samples was higher than that in the leaf samples. On the contrary, fungi (Chao 1) showed an increasing trend of species richness from root to stem and leaf samples (Table 1). The diversity of endophytic microbial in different parts of the host and the specificity of the rhizospheric soil were fully reflected.

The Beta diversity of the microbial community structure of rhizospheric soil and four parts of *S. sphenanthera* was evaluated according to the principal coordinate analysis diagram, which could intuitively show the microbial community difference of rhizospheric soil and different parts (Fig. S2). The distribution of the three points representing stem, leaf, and fruit in the figure was not close to each other, and the separation distance was not very large, which illustrated that the endophyte community structures in the above-ground parts were different, but the differences were small. The part of above-ground parts (stem, leaf, and fruit) was far apart from the below-ground parts (root and rhizospheric soil), indicating that there were great differences in the composition of microbial between the above-ground parts and the below-ground parts (Fig. S2).

**The microbial composition and community structure of bacteria and fungi**

The histogram abundance map could analyze the composition and proportion of species between rhizospheric soil and different parts more intuitively, taking the level of the phylum as an example (Fig. 1). The endophytic bacteria were mainly composed of Cyanobacteria and Proteobacteria, and the bacteria of rhizospheric soil were mainly composed of Acidobacteria and Proteobacteria (Fig. 1a). The fungi in rhizospheric soil and different parts of *S. sphenanthera* were mainly composed of Ascomycota and Basidiomycota (Fig. 1b). Ascomycota had higher compositions in fruit, stem, leaf, and root, while Basidiomycota had higher concentrations in rhizospheric soil (Fig. 1b). However, there were differences in the dominant flora of endophytes at lower levels in the root, stem, leaf, and fruit (Fig. 2, S3, and S4, Table S5 and S6).

There were 12 common bacterial genera in the rhizospheric soil and four parts of *S. sphenanthera* (Fig. 2a). *Achromobacter* was the dominant genus of leaf and stem, and *Methylobacterium* was the dominant genus of leaf. In addition, *Methylobacterium* was also found in fruit and stem (Fig. 2a). *Rhodoplanes* was the dominant genus in root and rhizospheric soil, and only existed in below-ground parts (Table S5). There were 11 common fungal genera belonging to rhizospheric soil and four parts of *S. sphenanthera* (Fig. 2b). Among the common genera, *Alternaria* was found with high concentrations in root, stem, leaf, and fruit, especially in stem. And *Cercospora* was found with high concentrations in stem and leaf (Fig. 2b). In addition, *Hebeloma* (rhizospheric soil), *unclassified_Helotiales* (root), *Stomiopeltis* (stem), and *Botrytis* (fruit) ran to greater than 10 % in their respective domains (Table S6).

The heat map of species level clustering showed that rhizospheric soil and root were still clustered as one group, indicating that microbes of rhizospheric soil and root were relatively...
close (Fig. S4). The clustering results of rhizospheric soil and four parts were consistent with OTUs clustering results (Fig. S1c). It indicated that rhizospheric soil and root microbes were relatively close, and the microbes of stem, leaf, and fruit were close (Fig. S4). According to the above results, there were significant differences in the composition and diversity of microflora in rhizospheric soil and different parts.

**Functional Annotations**

To explore the function of microorganisms in different tissue parts of *S. sphenanthera*, Tax4Fun and FunGuild software were used to predict the functions of bacteria and fungi based on 16S rDNA and ITS sequences (Fig. 3). The results showed that there was no significant difference in the function of bacteria between rhizospheric soil and different parts of *S. sphenanthera*, and most of the bacteria were concentrated in metabolism, environmental information processing, cellular processes and so on (Fig. 3a). Among them, more than 70 percent of the bacteria were involved in metabolic function, and among metabolic functions, global and overview maps accounted for about half (Fig. 3a, and S5). The relative abundance of amino acid metabolism and biosynthesis of amino acid pathway of endophytic bacteria was higher in leaf and stem than in other parts. In addition, the relative abundance of bacteria involved in amino acid synthesis in fruit was higher, but the relative abundance of amino acid metabolism was lower (Fig. S5).

Compared with other parts, the relative abundance of bacteria about two functions (xenobiotics biodegradation and metabolism (Fig. S5a), and microbial metabolism in diverse environments (Fig. S5b)) in rhizospheric soil and root was higher. In addition, the relative abundance of bacteria in root about three functions (lipid metabolism, metabolism of terpenoids and polyketides, and fatty acid metabolism) was higher (Fig. S5).

However, there were differences in fungal functions between rhizospheric soil and different parts of *S. sphenanthera* (Fig. 3b). Besides the unidentified groups, the fungi in rhizospheric soil and different parts of *S. sphenanthera* included 11 ecological functional groups (Fig. 3b). The relative abundance of fungi about four functions (arbuscular mycorrhizal, bryophyte parasite, ectomycorrhizal, and lichenized) in below-ground parts were much higher than that in the other three parts. Among the four functions, only the relative abundance of fungi with arbuscular mycorrhizal function was greater in root than in rhizospheric soil. Endophytic fungi (root, stem, leaf, and fruit) were mainly pathogenic functional groups and saprophytes, among which the ecological functional groups with higher relative abundance were mainly plant pathogen, animal pathogen, wood saprotroph, and undefined saprotroph. Only endophytic fungi in stem had algal parasite function (0.25 %) (Fig. 3b). The results also showed that the fungal groups in the samples are mainly predicted to have seven nutrient types, such as pathotroph, symbiotroph, saprotroph and pathotroph-saprotroph (Fig. 3c). In the rhizospheric soil, saprotroph-symbiotroph
and symbiotroph were the main nutrient forms, accounting for more than 79%. Pathotroph was the main nutrient form in leaf (31.67%) and fruit (63.68%), while the relative abundance of the fungal pathotroph-saprotoph trophic form in leaf was also high (40.27%). Pathotroph-saprotoph-symbiotroph and saprotroph were the main trophic forms of stem endophytic fungi, accounting for more than 88%. And the highest trophic pattern of endophytic fungi in root was Pathotroph-Saprotroph-Symbiotroph (39.03%) (Fig. 3c).

Content of active secondary metabolites in *S. sphenanthera*

The optimum extraction method for the highest extraction rate of the stem (1:20 ratio, 30 min, 30°C and 300 W) and leaf (1:15 ratio, 20 min, 35°C, 240 W) were different (Table S2 and S3). Except that ultrasonic temperature had a small impact on leaf, other factors had a significant influence on the yield of essential oils from the stem and leaf (Table S4). The factors affecting the yield of essential oils from stem and leaf were listed in order as follows: ultrasonic time > ratio of material to solvent > ultrasonic temperature > ultrasonic power, and the ratio of material to solvent > ultrasonic time > ultrasonic power > ultrasonic temperature (Table S3).

It could be found that the component and content of essential oil varied greatly from stem, leaf, and fruit (Fig. S6, Table S7). The components of essential oils (sesquiterpenes and oxygenated sesquiterpenes) from all samples with low content (< 0.1%) were not listed in Table S7. A total of 41 sesquiterpenoids were identified from all the samples, and 23 essential oils components were identified from the three parts (stem, leaf, and fruit) of *S. sphenanthera*, but the contents were different (Fig. 4 and Table S7). Each part had its high content of dominant components, but there were obvious differences in the dominant components of the three parts. For example, γ-muurolene, δ-cadinol, and trans-farnesol were characteristic components of the stem. α-Cadinol and neoisolongifolene-8-ol were characteristic components of leaf (Fig. 4). Isospathulenol, α-santalol, cedrenol, and Longiverbenone were characteristic components of fruit (Table S7).

Correlation Analysis

The correlation analysis between microbial communities (Top 20 at genus level) and the common components in different parts (Fig. S7 and S8) showed that 12 genera of bacteria and fungi had significant correlation with the common components (Fig. 5). Of the 23 essential oil components, only 16 components were associated with bacteria and 10 components were related to fungi (Fig. 5). *Staphylococcus* and *Hyphomicrobium* had positive correlation with 5 common components (γ-muurolene, β-bisabolene, germacrene D-4-ol, trans-farnesol, and ledene oxide). In addition, *Staphylococcus* showed a positive correlation with δ-cadinene, and *Hyphomicrobium* showed a significant positive correlation with β-bisabolene and ledene oxide. *Methylobacterium* and *Propionibacterium* were correlated with the contents of 3 common components, respectively.
Achromobacter, Burkholderia, and Planctomyces had positive effects on farnesol and negative effects on β-chamigrene, and there was a significant positive correlation among the three genera. Except for the bacteria genera mentioned above, the remaining genera were only related to one common component (Fig. 5a). Among the genera of fungi, only Stomiopeltis had the most influence on the essential oil components (farnesyl alcohol, δ-cadinene, and δ-cadinol). Botrytis, Cortinarius, and Zygophiala had negative effects on α-amorphene and positive effects on β-chamigrene, and there was a significant positive correlation relationship among the three genera. Unclassified_Agaricales, unclassified_Thelephoraceae, and unclassified_Chaetothyriales had a negative correlation with α-bisabolene, and there was a positive correlation relationship between the three genera. In addition, except germacrene D, β-himachalene, α-cadinol, and α-bisabolol were affected by only one genus, and the remaining components were associated with two or three genera (Fig. 5b). These results suggested that bacterial communities played a greater role than fungal communities in the accumulation of active secondary metabolites.

Discussion

Microbial diversity analysis of rhizospheric soil and different parts in S. sphenanthera

Endophytes existed in specific tissues of plants and interacted with plants through the exchange of nutrients, enzymes (catalase, oxidase, etc.), functional factors (biosurfactants, etc.), and signal transmission (Hacquard et al. 2015). They colonized plant tissues for a long time and did not produce negative effects similar to pathogens, such as the destruction of photosynthesis, nutrient transfer, etc. On the contrary, the presence of these endophytes in host plants had beneficial effects on their health and growth. In recent years, high-throughput sequencing technology had been widely used to study the diversity of environmental, food, animal, plant, and microbial communities (Liu et al. 2023). Endophyte determination by high-throughput sequencing technology could obtain detailed information about the microbial structure in plants, and analysis of plant internal micro level of ecological and environmental relation. At the same time, the study of endophytes of medicinal plants provided the possibility of screening high-quality strains and fermentation for the production of drug-active ingredients and established a new mode of genuine identification of medicinal materials, which had gradually become a research focus of microbial resources of medicinal plants (Adeleke and Babalola, 2021).

In this study, Proteobacteria, Cyanobacteria, and Acidobacteria were the main bacteria, and Ascomycota and Basidiomycota were the main fungi at the phylum level (Fig. 1). According research found that most microbial communities had little difference at the phylum level, which was consistent with the results of this study, which fully demonstrated the similarity of microbes.
in larger taxonomic units (Sun et al. 2022; Shao et al. 2023). At present, endophytes could be isolated from various parts and organs of studied plants, and the structure composition and abundance of endophytes would change with different plant varieties, parts, and development periods (Xia et al. 2023). This study revealed that there were significant differences between the microbial communities screened from rhizospheric soil and endophytic communities screened from different parts of S. sphenanthera (Table S5 and S6). There were also significant differences in the community structure and composition of endophytes in different parts of S. sphenanthera, which might be related to the different physiological structures of different tissues and the diversity of endophyte sources. The microbial communities of rhizospheric soil and root were more similar than those of endophytes from stem, leaf, and fruit, suggesting that endophytes of root invaded from rhizospheric soil and reached the root through transport tissue. However, the endophytic communities of the stem and fruit were similar to the leaf, suggesting that the microorganisms of the stem and fruit might enter through the phyllosphere (Fig. S1b, c).

**Active secondary metabolites of different parts in S. sphenanthera**

In this study, the component and content of essential oil extracted from different parts of S. sphenanthera were different. Due to the different components and content of essential oil, their characteristics were formed, such as the colors of essential oil in leaf (yellowish green), stem (slightly yellow), and fruit (light yellow) were different. As a Traditional Chinese medicine, the dried fruit of S. sphenanthera was the main part to be used as medicine. Compared with stem and leaf, there were many studies on the essential oil of fruit. The main components of essential oil were sesquiterpenoids and oxidized sesquiterpenoids, which were the same as those reported in previous literature (Yu et al. 2017). However, the main components of sesquiterpenoids were different, which might be related to the extraction method and the origin of materials (Yu et al. 2017). However, the non-medicinal parts (stem and leaf) were abundant and usually had the same or similar biological active ingredients as the medicinal parts, so they had broad development potential and application prospects.

The biological activities of the essential oils were related to the major components. The contents of γ-muurolene, δ-cadinol, and trans-farnesol were the highest in the essential oil of stem. γ-Muurolene and δ-cadinol had anti-microbial activity, and δ-cadinol had cytotoxic activity against MCF7 cells (Ringel et al. 2022; Silva et al. 2009). Farnesol could improve serum allergy antibody titers in mice (Ku and Lin, 2016). As the characteristic components of leaves, α-dienol and neoisonolene-8-ol had antibacterial activities (Hassan et al. 2020; Hsu et al. 2020). In addition, neoisonolene-8-ol could also be used in the synthesis of spices (Tang et al. 2021). Among the dominant components of fruit, isspathulenol had an immunosuppressive effect on activated lymphocytes (Paksoy et al. 2016), α-santalol had antibacterial activity (Bommareddy et
al. 2018), cedarol could be used as a flavorful ingredient (Bhatia et al. 2008), and longiverbenone was an active toxic compound (Khani and Heydarian, 2014).

Effects of microorganisms on active secondary metabolites of S. sphenanthera

Plant secondary metabolites were important agents of plant-microbial interaction (Sasse et al. 2018). Endophytes could promote the production of plant secondary metabolites or produce metabolites with medicinal value (Jia et al. 2016). In addition to the endophytes, microorganisms from plant rhizospheric soil also played an important role in energy conversion and material cycling in plants (Perez-Jaramillo et al. 2019). This indicated that it was important to explore the relationship and influence of microorganisms on the quality of S. sphenanthera. In this study, we found that the content of secondary metabolites in S. sphenanthera was correlated with the presence of microorganisms. Bacteria had a stronger correlation with the accumulation of active components than fungi. These results highlighted the involvement of specific bacterial communities in plant secondary metabolic pathways, suggesting that a variety of bacterial flora may promote the production of plant secondary metabolites (Fig. 3a; Wu et al. 2021).

In this study, Achromobacter, Methylobacterium, and Propionibacterium with high abundance had been detected in above-ground parts of S. sphenanthera (Fig. 2a). Achromobacter could inoculate vetiver (Chrysopogon zizanioides) with aromatic compounds as the only carbon source (Ho et al. 2013). Methylobacterium and Propionibacterium had the greatest influence on the composition of essential oils (Fig. 5a), and Methylobacterium can protect soybean (Glycine max) against pathogens by inducing endophytic community changes (Christian et al. 2021). Propionibacterium was a kind of bacterial microorganism in the stem that could synthesize propionic acid using special carboxylic enzymes, and the study showed that Propionibacterium was widely used in the production of vitamin B12, four pyrrole compounds, propionic acid, acid, as well as probiotics and cheese industry (Guyomarc'h et al. 2020). Staphylococcus and Hyphomicrobium also had the most influence on essential oil composition. Hyphomicrobium could be used to synthesize vitamin B12 (Dudko et al. 2022). In addition, Rhodoplanes, Candidatus Solibacter, and Gemmata were also the dominant flora in the below-ground parts (Table S5). Rhodoplanes were facultative photoorganics and potential nitrate fixation bacteria (Hiraishi and Ueda, 1994). Candidatus Solibacter, Gemmata, and Staphylococcus were pathogenic (Backman and Sikora, 2008; Muriuki et al. 2021; Othman et al. 2021). The remaining bacteria were involved in plant protection, growth promotion, and functional secondary metabolites. The results also confirmed that endophytic bacteria and rhizospheric soil microorganisms had a low risk to plants.

The relative abundances of unclassified_Thelephoraceae, unclassified_Agaricales, and Cortinarius were higher in the below-ground part of the 12 fungal genera that influenced the
components of essential oil, while the relative abundances of other genera were higher in the above-ground part (Fig. 5b). Some of the fungi in this study were pathogenic microorganisms, and although they temporarily lost pathogenicity, they might regain it under environmental selection, such as Botrytis (Kan et al. 2014) and Zygodiala (Fig. 4b; Batzer et al. 2008). At the same time, pathogens could be used as inducers to enhance the host's resistance to disease, activate the host’s defense system, and improve the host's defense ability against pathogens (Backman and Sikora, 2008). In addition to the above two pathogens, Stomiopeltis and Cortinarius also had higher effects on essential oils (Fig. 5b; Aijitomi et al. 2017). Most Cortinarius were edible mycorrhizas and could also form ectomycorrhizas (Bödeker et al. 2014). In addition, some endophytic fungi and their metabolites in above-ground parts could promote the growth and development of host plants, such as Trametes and Eremothecium (Table S6). Although Trametes was a wood saprotroph and Eremothecium was a plant pathogen (Fig. 4b), the secondary metabolites of Eremothecium and Trametes had medicinal value (Semenova et al. 2022; Zmitrovich et al. 2012). Moreover, Hebeloma and Clavulinopsis were dominant genera in the rhizospheric soil of S. sphenanthera (Table S6). Hebeloma affected the flow of phosphorus in the litter, the transformation of soil organic matter itself, and the retardation of humification by participating in fungal interactions (Mrnka et al. 2020). Clavulinopsis, a member of the Clavariaceae family, assimilated and transferred 15N-depleted N form to the leaves of host plants, and built fungal cell walls by absorbing 15N-enriched N (Birkebak et al. 2013; Mayor et al. 2009).

Because of the roles of endophytes in plant growth, our research results were helpful to expand the application of endophytes in the production of S. sphenanthera and its important metabolites. The information on the differences in endophytes between above-ground and below-ground parts could serve as the basis for the selection of functional microorganisms.

Conclusions

In conclusion, this is the first study to analyze the microbial diversity in different parts and rhizospheric soil of S. sphenanthera and the effects of microorganisms on the secondary metabolites. Our findings demonstrated that the components of essential oils in different parts of S. sphenanthera were different, especially the dominant components. At the same time, plant organs have important effects on the composition and structure of the rhizosphere and endophytic communities of S. sphenanthera. In addition, positive and negative correlations were found among active secondary metabolites and microbial operating taxonomic units based on the correlation coefficient matrix, indicating that the accumulation of secondary metabolites in S. sphenanthera was closely related to microbial community composition. As a Traditional Chinese medicine, S. sphenanthera had important practical significance in medicine and the economy.
Our findings provide new insights into the distribution and resources of endophytic communities in different parts of the medicine plant. There are still some limitations in this study, the migration mechanism between microorganisms in different parts and how microorganisms affect the active secondary metabolites are not very clear, and further experiments are needed to solve these problems.

ADDITIONAL INFORMATION AND DECLARATIONS

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Author Contributions
Xiaolu Qin conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.
Han Pu performed the experiments, prepared figures and/or tables, and approved the final draft.
Xilin Fang performed the experiments, prepared figures and/or tables, and approved the final draft.
Qianqian Shang analyzed the data, prepared figures and/or tables, and approved the final draft.
Jianhua Li analyzed the data, prepared figures and/or tables, and approved the final draft.
Qiaozhu Zhao performed the experiments, prepared figures and/or tables, and approved the final draft.
Xiaorui Wang conceived and designed the experiments, performed the experiments, analyzed the data, authored or reviewed drafts of the article, and approved the final draft.
Wei Gu conceived and designed the experiments, reviewed drafts of the article, and approved the final draft.
All authors have read and agreed to the published version of the manuscript.

Competing Interests
The authors declare there are no competing interests.

Data Availability
The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Supplementary Information
Supplemental information for this article can be found online at

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Table 1 (on next page)

Alpha diversity index of bacteria and fungi (m ± sd).

Different lowercase letters (a-c) indicate significant differences ($p < 0.05$) of the same alpha diversity index between rhizosphere soil and different parts of $S.$ *sphenanthera*, one-way ANOVA, Tukey test.
### Table 1. Alpha diversity index of bacteria and fungi (m ± sd).

<table>
<thead>
<tr>
<th>Alpha index</th>
<th>Rhizosphere soil</th>
<th>Root</th>
<th>Stem</th>
<th>Leaf</th>
<th>Fruit</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bacteria</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACE</td>
<td>7480.27±931.55 a</td>
<td>7336.57±679.33 b</td>
<td>8624.06±787.37 b</td>
<td>5099.82±824.41 c</td>
<td>5380.41±817.88 c</td>
</tr>
<tr>
<td>Chao1</td>
<td>7052.79±875.83 a</td>
<td>8624.06±745.22 b</td>
<td>4257.31±562.71 c</td>
<td>4964.72±765.88 c</td>
<td>4413.24±357.68 c</td>
</tr>
<tr>
<td>Shannon</td>
<td>11.59±0.23 a</td>
<td>5.97±1.32 b</td>
<td>4.97±2.13 b</td>
<td>4.68±0.78 b</td>
<td>4.48±0.64 b</td>
</tr>
<tr>
<td>Simpson</td>
<td>1.00±0.00 a</td>
<td>0.81±0.15 a</td>
<td>0.84±0.17 a</td>
<td>0.80±0.16 a</td>
<td>0.78±0.22 a</td>
</tr>
<tr>
<td><strong>Fungi</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACE</td>
<td>1502.15±312.88 bc</td>
<td>1184.42±307.64 c</td>
<td>1989.66±268.35 b</td>
<td>2541.61±323.54 a</td>
<td>1210.33±237.67 c</td>
</tr>
<tr>
<td>Chao1</td>
<td>1555.06±244.68 c</td>
<td>1192.01±277.37 c</td>
<td>2142.18±245.33 b</td>
<td>2659.47±356.57 a</td>
<td>1283.38±231.64 c</td>
</tr>
<tr>
<td>Shannon</td>
<td>5.15±0.40 bc</td>
<td>4.85±0.61 c</td>
<td>6.21±0.51 b</td>
<td>7.39±0.78 a</td>
<td>5.69±0.56 bc</td>
</tr>
<tr>
<td>Simpson</td>
<td>0.93±0.05 ab</td>
<td>0.90±0.04 b</td>
<td>0.96±0.03 a</td>
<td>0.98±0.01 a</td>
<td>0.93±0.02 ab</td>
</tr>
</tbody>
</table>

Different lowercase letters (a-c) indicate significant differences (p < 0.05) of the same alpha diversity index between rhizosphere soil and different parts of *S. sphenanthera*, one-way ANOVA, Tukey test.
Figure 1

Stack diagram at the level of the phylum of the sample.

a: Bacteria. b: Fungal
Figure 2

The differences in the abundance of the common genus of microbial in rhizosphere soil and different parts of *S. sphenanthera* (m ± sd).

a: Bacteria. b: Fungal. Different lowercase letters (a-c) indicate significant differences (*p* < 0.05) of the same genera between rhizosphere soil and different parts of *S. sphenanthera*, one-way ANOVA, Tukey test.
Figure 3

Horizontal clustering tree-stacking diagram of microbial function annotation (a: bacteria, b: fungal) and trophic mode of fungal (c) in rhizosphere soil and different parts of *S. sphenanthera*.

a-e and A-E: Rhizosphere soil, Root, Stem, Leaf, and Fruit
Common chemical compositions (sesquiterpenes and oxygenated sesquiterpenes) identified in the volatile oils of different parts of *S. sphenanthera*
Figure 5

Pearson correlations between genera of microbial and common chemical compositions of *S. sphenanthera*.

The *, **, and *** mean significant correlation estimates at the level of 0.05, 0.01, and 0.001, respectively.