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OPEN Identification of the rhizospheric microbe and metabolites that led by the continuous cropping of ramie (Boehmeria nivea L. Gaud)

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Continuous cropping lowers the production and quality of ramie (Boehmeria nivea L. Gaud). This study aimed to reveal the metagenomic and metabolomic changes between the healthy- and obstacle-plant after a long period of continuous cropping. After 10 years of continuous cropping, ramie planted in some portions of the land exhibited weak growth and low yield (Obstacle-group), whereas, ramie planted in the other portion of the land grew healthy (Health-group). We collected rhizosphere soil and root samples from which measurements of soil chemical and plant physiochemical properties were taken. All samples were subjected to non-targeted gas chromatograph-mass spectrometer (GS/ MS) metabolome analysis. Further, metagenomics was performed to analyze the functional genes in rhizospheric soil organisms. Based on the findings, ramie in Obstacle-group were characterized by shorter plant height, smaller stem diameter, and lower fiber production than that in Healthgroup. Besides, the Obstacle-group showed a lower relative abundance of Rhizobiaceae, Lysobacter antibioticus, and Bradyrhizobium japonicum, but a higher relative abundance of Azospirillum lipoferum and A. brasilense compared to the Health-group. Metabolomic analysis results implicated cysteinylglycine (Cys-Gly), uracil, malonate, and glycerol as the key differential metabolites between the Health- and Obstacle-group. Notably, this work revealed that bacteria such as Rhizobia potentially synthesize IAA and are likely to reduce the biotic stress of ramie. L. antibioticus also exerts a positive effect on plants in the fight against biotic stress and is mediated by metabolites including orthophosphate, uracil, and Cys-Gly, which may serve as markers for disease risk. These bacterial effects can play a key role in plant resistance to biotic stress via metabolic and methionine metabolism pathways.

Abbreviations

GS/MS	Gas chromatograph-mass spectrometer
PCoA	Principal co-ordinates analysis
NMDS	Nonmetric multidimensional scaling
Cys-Gly	Cysteinylglycine
TICs	Total ion currents

Continuous cropping is defined as the agricultural practice of producing or growing a single crop on a piece of land. Long-term continuous cropping causes plant growth obstacles, mainly manifesting as decreased leaf photosynthesis and metabolism, soil-borne diseases, deteriorated soil environment, and disturbance of microbial and nematode community¹⁻⁵, which consequently contaminate the soil, inhibits plant growth and reduces crop quality^{5,6}.

Increasing evidence has revealed the changes in soil microbial diversity and structure under continuouscropping conditions^{1,7,8}. Continuous cropping usually lowers the abundance of beneficial microbes, such as Azotobacter, Rhizobium, and Nitrobacteria, while similarly enhancing the growth of harmful microbe in soils⁹⁻¹². For instance, Tan et al. in their study reported that the continuous cropping of Panax notoginseng reduced the fungal diversity in the rhizospheric soil and roots³. Also, Han et al. that continuous cotton cropping for 5 years not only reduced the diversity of nutrition related bacteria but also lowered the relative abundance of ammonifying

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Parameters	Health-group	Obstacle-group	p
Total N (%)	1.35 ± 0.03	1.30 ± 0.05	0.152
Total P (%)	0.08 ± 0.00	0.07 ± 0.01	0.104
Total K (%)	3.42 ± 0.03	3.50 ± 0.02	0.018
Available P (mg/kg)	29.03 ± 0.15	20.27 ± 0.15	< 0.0001
Available K (mg/kg)	101.33 ± 2.08	86.67±1.53	0.0006

Table 1. Differences in soil chemical parameters between Health-group and Obstacle-group. Differences areanalyzed using a t-test.



Figure 1. Changes in ramie growth and fiber traits through continuous cropping based on data collected from 30 individual plants in each group. Differences are analyzed using a t-test.

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bacteria, nitrifying bacteria, and aerobic nitrogen-fixing bacteria. Changes in the diversity of soil bacteria are correlated with soil physicochemical properties, for example, pH value, moisture, salinity, porosity, available nitrogen, and organic matter^{13–15}. Accordingly, continuous cropping practices lower crop yields and quality by affecting the biosynthesis and metabolism of active plant components¹⁶.

Ramie (*Boehmeria nivea* L. Gaud), also known as "China grass," is a perennial, diploid (2n = 28) and herbaceous plant of the *Urticaceae* family. Ramie is widely cultivated owing to its bast fiber, which offers excellent properties such as excellent thermal conductivity and high tensile strength¹⁷. Ramie fiber is extracted from stem bast and poses numerous excellent characteristics, for example, long strands, smooth texture, and high tensile strength¹⁸. Previous studies reported that continuous cropping of ramie reduces plant height, stem diameter, and fiber yield^{5,12,19}. In our previous studies, we showed that the stem growth of ramie is impeded by continuous cropping¹². Moreover, we found that the abundance of *Firmicutes* was correlated positively with the plant height, stem diameter, and ramie fiber yield. However, as the duration of continuous cropping of ramie. However, none of the reports indicated whether soil microbial communities and metabolites were associated with continuous cropping obstacles.

In production practice, we found an interesting phenomenon: After 10 years of continuous cropping (single variety), ramie planted in some portions of the land were characterized by weak growth and low yield (Obstaclegroup), on the other hand, ramie planted in the other portions exhibited healthy growth (Health-group). Thus, we hypothesized that this difference may be attributed to the effects of soil microorganisms and could be revealed by soil metagenomics and metabolome analysis. This present study attempted to provide a better understanding of the relationship between soil bacteria and metabolites and continuous cropping obstacles and to elucidate on the growth of ramie under continuous cropping practice.

Results

Continuous cropping of ramie. After 10 years of continuous cropping (single variety), ramie planted in some portions of the grew healthy (defined as Health-group) whereas, ramie planted in the other portions grew weak in and with low yields (defined as Obstacle-group). Soil samples in Obstacle-group had lower contents of available P and K, and higher content of total K than Health-group (p < 0.05, Table 1). The values of plant height, stem diameter, bark thickness, weight, and fresh and dry fiber weight in the Obstacle-group were lower compared to those in Health-group (p < 0.0001, Fig. 1). These findings confirmed that growing the same ramie variety in different continuous cropping lands greatly varies.

Illumina gene sequencing. Illumina sequencing generated a total of 326.58 M clean reads from 6 rhizosphere soil samples with an average Q30 value of 94.59% and GC content of 64.37%. A total of 10,450,4244 scaffolds and 80,931 open reading frames (ORFs, longer than 200 bp) were retrieved from the data (Table S1). Additionally, 435,887 non-redundant gene catalogs or clusters were identified, including 49,444 annotated genes



Figure 2. Differences in microbial gene number and kingdom abundance between the two groups. Boxplots indicate the minimum, maximum, median and upper and lower quartiles of the number of genes in each group. In a boxplot, the black line indicates the median.

to bacterial taxonomy. Expression analysis showed that 5517 clusters were differentially expressed between the two groups. Notably, the numbers of annotated genes in the two groups were different (Fig. 2). About 98.26% of the clean reads originated from bacteria, followed by 1.28% and 0.3% from archaea and virus, respectively.

Identification of different microbe between the two groups. Principal coordinates analysis (PCoA) and Nonmetric Multidimensional Scaling (NMDS) analyses demonstrated different beta diversity among all samples (Fig. 3A,B). Samples in Health-group were clustered together in both PCoA (PC1=79.15%, PC2=8.62%, and PC3=6.19%) and NMDS. Similarly, samples in Obstacle-group were clustered together. Among the top 10 taxonomies (from species to phylum), no significant difference was found in the abundance of microbes. Then, we further analyzed the microbes with a lower relative abundance. We found that the relative abundances of 8 phyla (Fig. 4, two Archaea phyla), 41 families (Fig. S1), 132 genera and 493 species (Table S2) were different between Health- and Obstacle-group. For instance, the relative abundance of Acidithiobacillaceae $(0.023 \pm 0.002\%)$, Proteobacteria), Ectothiorhodospiraceae $(0.520 \pm 0.016\%)$, Proteobacteria) and Xanthomonadaceae $(0.973 \pm 0.020\%, Proteobacteria)$ family in Obstacle-group was higher than that in Healthgroup (0.019±0.000%, 0.480±0.055%, and 0.921±0.023% correspondingly; p<0.05, Fig. S1). The abundance of *Rhizobia* (*Rhizobiaceae*) was lower in Obstacle-group than in Health-group $(1.073 \pm 0.014\% \text{ vs. } 1.120 \pm 0.008\%,$ p < 0.05). Besides, the abundance of two virus families *Podoviridae* and *Myoviridae* was higher and lower in Obstacle-group compared with Health-group, respectively (*Podoviridae* $0.030 \pm 0.003\%$ vs. $0.038 \pm 0.003\%$; *Myo*viridae $0.018 \pm 0.001\%$ vs. $0.015 \pm 0.000\%$, p<0.001). At the species level, the relative abundance of Janthino*bacterium lividum* $(0.019 \pm 0.001\%$ vs. $0.014 \pm 0.001\%$, p < 0.05), *Bradyrhizobium japonicum* $(0.119 \pm 0.002\%$ vs. 0.102±0.002%, p<0.05) and Lysobacter antibioticus (0.030±0.002% vs. 0.023±0.001%, p<0.05; Table S2) were higher in Obstacle-group compared to Health-group (p < 0.05). These findings demonstrated that the growth of plants under continuous-cropping conditions is correlated with the changing diversity of soil microbe, especially the bacteria.

GC–MS/MS metabolomics. TICs (total ion currents) revealed 554 peaks detected from all samples (including quality control samples). Metabolite annotation results showed 190 identified metabolites. PCA (Fig. S2A) and OPLS-DA (Fig. S2B) results showed that there were different metabolites in soil and root between the two groups. Moreover, we identified 20 and 31 differential metabolites in the soil and root samples between the two groups, respectively (Table S3), including 3 overlapping metabolites (threitol, glycerol, and 2-Deoxyerythritol).

Identification of the differential metabolites in soil samples. From the soil samples, there were higher levels of threitol, 2-Deoxyerythritol, uracil, malonic acid, and Cys-Gly (Cysteinylglycine and lower glycerol level in Obstacle-group than in Health-group (Table S3, Fig. 5A). KEGG (Kyoto Encyclopedia of Genes and



Figure 3. Beta diversity among samples sequenced. (**A**) The principal co-ordinates analysis (PCoA) results. (**B**) The Nonmetric Multidimensional Scaling (NMDS) analysis results. Both analyses were performed based on the Euclidean distance algorithm.



Figure 4. A heatmap of the 8 phyla among samples. The dendrogram to the left and top of the heatmap shows the clustering tree. The box was colored based on the standardized relative abundances data. The redder color shows a higher abundance of the phyla, and the blue color shows a lower abundance.

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Figure 5. Different metabolites from the soil and root samples between the two groups. Several differential metabolites in soil (**A**) and root (**B**) samples between the two groups, respectively. VIP, variable importance in projection. VIP scores estimate the importance of each variable in the projection used in a PLS model. The p-value is analyzed using a t-test. N/B: the Y axes have been divided into two or three segments for a clearer representation of smaller bars. P values are marked in the figure (n=6).

Genomes) pathway enrichment analysis showed that these 20 different metabolites in soil samples were enriched in 17 pathways. Notably, Cys-Gly, orthophosphate, uracil, malonate, and glycerol had a higher frequency in these pathways. Moreover, Uracil was involved in multiple pathways, such as metabolic pathway, pantothenate and CoA biosynthesis, pyrimidine metabolism, among others, whereas malonate was associated with pyrimidine metabolism and beta-Alanine metabolism.

Identification of the differential metabolites in root samples. From the root samples, the level of glycerol and threitol was higher in Obstacle-group compared to Health-group (Fig. 5B). On the contrary, the level of 2-Deoxyerythritol, maleic acid, and raffinose in the Obstacle-group was higher than in Health-group (p < 0.05). The KEGG enrichment analysis showed that 90 pathways were enriched, including Carbon metabolism (e.g. fumarate, 3-Hydroxypropanoate, citrate, D-Gluconic acid, D-Glucono-1,5-lactone, and pyruvate), Metabolic pathways (e.g. fumarate, 3-Hydroxypropanoate, citrate, glycerol, myo-Inositol, alpha, alpha-Trehalose, D-Gluconic acid, cysteamine, pyruvate, and Phytosphingosine), Biosynthesis of phenylpropanoids, and Biosynthesis of plant secondary metabolites (e.g. fumarate, citrate, gallate, L-Phenylalanine and pyruvate; Table S4). Of all the enriched differential metabolites in the pathways, orthophosphate, uracil and Cys-Gly were found to exhibit a high frequency.

Integrated analysis of the metagenome and metabolomics. To identify the correlation between the differential microbial genes and metabolites in the two groups, we identified the same pathways associated with the differential genes and metabolites in soil and root samples. Of note, the oxidative phosphorylation (ko00190), pyrimidine metabolism (ko00240), beta-alanine metabolism (ko00410), glutathione metabolism (ko00480), metabolic pathways (ko01100), ABC transporters (ko02010), and two-component system (ko02020) were found to be correlated with the content of metabolites (e.g., orthophosphate, uracil, malonate and Cys-Gly) through the genes expressed in 5 Archaea, one virus and 206 bacteria (Table S3). For instance, Archaea Nitrososphaera viennensis, Candidatus Nitrososphaera gargensis and Candidatus Nitrososphaera evergladensis regulated two-component system (ko02020) and metabolic pathways (ko01100) by changing the metabolomic profiles of orthophosphate, uracil, and Cys-Gly. Genes expressed in Archaea, for example, cmpC glutamate dehydrogenase (GDH2) and glrR are listed in Table S3. Of which, WspR, expressed in Azoarcus sp. PA01 and Planococcus sp. PAMC_21323, regulated orthophosphate metabolism and was involved in the two-component system pathway; trxB was expressed by Nitrospira sp. SCGC_AG-212-E16 and Bradyrhizobium elkanii, which regulated pyrimidine metabolism by regulating the metabolism of uracil and malonate. The f accD (Nitrospira moscoviensis) and accC (Janthinobacterium lividum) genes could regulate orthophosphate, uracil, and Cys-Gly and were enriched in metabolic pathway (ko01100) (Table S3). Besides, we identified several genes involved in methionine metabolism, which regulates orthophosphate, uracil and Cys-Gly metabolism, they include gshB (encodes glutathione synthase, GSH) in Azospirillum lipoferum; pepA (encodes leucyl aminopeptidase, LAP) in Elusimicrobium minutum; phoB1 (encodes alkaline phosphatase synthesis response regulator PhoP) in A. brasilense; and mmuM (encodes homocysteine S-methyltransferase, HMT) in Desulfitobacterium metallireducens (Fig. 6A). In addition, an interaction network was constructed to reveal the relationship between metabolites and candidate genes (Fig. 6B). Results showed that Threitol, Cys-Gly, oxamide and D-(glycerol 1-phosphate) were highly associated with the genes. These findings uncovered the crucial roles of soil bacterial diversity in the rhizospheric and root metabolites, in particular, Cys-Gly, which participated in multiple pathways and showed rich interactions with other compositions.



Figure 6. The integrated analysis of different bacteria, metabolites and bacterial genes. (**A**) The relationship between different metabolites and different genes enriched in the same pathway. The heatmap shows the relative abundance of bacteria in different groups, the redder the color, the higher the abundance. The colored scale shows the level of abundance based on the standardized relative abundances data. Gene name and bacterial name horizontally correspond to each other, the larger the size, the higher the interaction. (**B**) The interacting network among the top 50 metabolites and genes, the darker and bigger the node, the richer the interaction with the genes.

Discussion

Continuous cropping obstacles are primarily related to disorders or deterioration of rhizosphere microorganisms^{20,21}. Our present study confirmed that continuous cropping of ramie is correlated with the changing rhizospheric soil microbes and soil metabolomics as well as the root metabolism. The present study identified that the abundance of some bacteria, including Rhizobia, L. antibioticus, and Bradyrhizobium japonicum, were lower in Obstacle-group compared to Health-group, a finding that concurs with previous reports²²⁻²⁵. A higher abundance of Rhizobia can increase soil biomass, as was confirmed in previous experiments²⁶⁻²⁹. Among many soil bacteria, Rhizobia can produce higher levels of indolacetic acid (IAA)³⁰, which is one of the most physiologically active auxins. IAA is a common product of L-tryptophan metabolism produced by several microorganisms including Plant Growth Promoting Rhizobacteria (PGPR)³¹. The endogenous or exogenous IAA can stimulate cell elongation by modifying certain conditions, for example, increasing in osmotic contents of the cell, increasing the permeability of water into cells, reducing wall pressure, increasing cell wall synthesis and inducing specific RNA and protein synthesis³². In addition to *Rhizobia*, we identified two bacteria (including A. lipoferum and A. brasilense) that were highly abundant in the Obstacle-group compared to Health-group. Notably, A. lipoferum and A. brasilense are growth-promoting bacteria and are capable of impoving plant resistance to abiotic stresses partially via enhancing plant abscisic acid level^{33,34}. Both A. lipoferum and A. brasilense can also produce IAA³⁵. For instance, Mazhar et al. showed that the inoculation of A. lipoferum ameliorated salinity-induced growth stress in wheat by promoting production of plant osmolytes (proline and soluble sugars). However, a recent study performed by Rivera et al. demonstrated that L-amino acids (e.g., L-Leu, L-Phe, L-Ala, L-Ile, and L-Pro) inhibited the growth of bacteria and biosynthesis of IAA. Elsewhere, A. brasilense inoculation could increase the uptake of nitrate, a substance of GSH³⁶, whereas it elevated nitrogenase release, thereby revealing superior plant growth promotion³⁷. Moreover, the overexpression of GSH in plants has been shown to contribute to improved plant stress tolerance and productivity³⁸. It is worth noting that A. lipoferum expresses the GSH encoding gene gshB, while D. metallireducens expresses a gene mmuM encoding HMT involving in plant tolerance and defense response^{39,40}. Although no L-amino acids were detected among the differential metabolites in soil and ramie root, our study uncovered that these bacterial genes regulated the metabolism of orthophosphate, uracil, and Cys-Gly, thus participates in metabolic pathway and methionine metabolism. Therefore, we speculated that the increased abundance of A. lipoferum and A. brasilense in the soil may be associated with the resistance to the continuous cropping induced abiotic stress in ramie root. In the present study, L. antibioticus was revealed to play important roles in various metabolic pathways via the regulation of metabolites such as orthophosphate, uracil, and Cys-Gly. Moreover, L. antibioticus could produce phenazine antibiotics⁴¹. In the recent past, Laborda et al. showed that L. antibioticus OH13 produced antifungal p-aminobenzoic acid. Another study reported that L. antibioticus exhibited biocontrol potential as it produced 4-hydroxyphenylacetic acid and several lytic enzymes against phytophthora blight⁴². In summary, these studies showed that *L. antibioticus* exerts a positive effect on plants in fighting biological stress. Low levels of L. antibioticus in the Obstacle-group may imply a decline in plant defense mechanism. Moreover, metabolites such as orthophosphate, uracil, and Cys-Gly may serve as mediators for L. antibioticus to play an active role. In addition, Bradyrhizobium japonicum, a microsymbiotic nitrogen-fixing bacteria, is known to enhance legume-root nodulation, which when introduced when planting, improve crop yields, especially legumes⁴³. As far as we know, no report has explored whether Bradyrhizobium japonicum is related to the growth of ramie. However, based onour findings, we speculate that Bradyrhizobium japonicum may also play an active role in maintaining the health of ramie.

Our metagenome analysis showed a lower relative abundance of several soil probiotics in the Obstacle-group, which may have been induced by long-term continuous cropping. Additionally, the NMDS and PCoA analyses using different methods revealed the distinct separations among samples from different groups. This showed that differences exist in the microenvironment of rhizosphere soil between the Obstacle-group and Healthgroup. The PCoA based on the weighted UniFrac algorithm clearly demonstrated the variations among different groups. And in the dimension of PC1, the two groups can be distinguished to the greatest extent (79.15%). These differences in soil microorganisms might be the most important factor that ultimately led to a greater difference between the two groups. Han et al. showed that 5-year continuous cropping of cotton lowered not only the richness of nutrition bacteria type in soil but also the abundance of ammonifying bacteria, nitrobacteria, and aerobic nitrogen-fixing bacteria. Also, a different study reported that two plant growth-promoting bacteria Azospirillum brasilense Cd and B. pumilus ES4 could enhance the growth of microalga by inducing the production of total carbohydrates, chlorophyll a and total lipids in microalga³⁴. Besides, numerous previous studies found that Bradyrhizobium and Rhizobium (Proteobacteria) are symbiotic bacterial partners that promote the formation of nitrogen-fixing nodules on legumes, thereby enhancing the plant growth, resistance, and tolerance to abiotic stresses and pathogens⁴⁴⁻⁴⁶. Similarly, our present study reported higher relative abundances of R. leguminosarum, A. chroococcum and R. mesoamericanum (Proteobacteria) in the Health-group than that in the Obstacle-group, which was inversely associated with the inhibition of ramie growth. The above-mentioned bacteria potentially play a key role in keeping plants healthy. Previous findings revealed the Lysobacter species as a potential source of novel antimicrobial plant metabolites^{41,47,48}. Notably, L. antibioticus has been revealed to be capable of producing phenazine antibiotics⁴¹. For example, a recent study by Laborda et al. showed that *L*. antibioticus OH13 produced antifungal p-aminobenzoic acid. In our study, we suggested that the lower level of L. antibioticus in the Obstacle-group may signify a decline in plant defense.

Conclusions

Soil bacteria may primarily affect health and obstacles to the growth ramie after long-term continuous cropping. Bacteria such as *Rhizobia* can synthesize IAA and are likely to reduce the biotic stress of ramie. For instance, *L. antibioticus* exerts a positive effect on plants in fighting biotic stress. In this study, we showed that lower levels of *L. antibioticus* in the Obstacle-group may imply a decline in plant defense. Metabolites including orthophosphate, uracil, and Cys-Gly might serve as mediators for *L. antibioticus* to play an active role. Notably, subjecting ramie to biological stress may increase in abundance of *A. lipoferum and A. brasilense*. Consequently, the bacterial changes are associated with various metabolic pathways through the regulation of metabolites such as orthophosphate, uracil and Cys-Gly. Particularly, Cys-Gly participated in multiple pathways and showed rich interaction in the interaction analysis. A low abundance of beneficial bacteria in Obstacle-group might be associated with the resistance to the continuous cropping induced abiotic stress in ramie root. These bacterial changes may play a key role in plant resistance to biotic stress via the metabolic and methionine metabolism pathways. Thus, this present study provides novel insights into the association of soil microbe and metabolome. These findings might show some indications on the agricultural practice for continuous ramie cropping.

Methods

Plant materials and sample collection. The ramie cultivar (Zhongzhu NO.1) was planted at the experimental field in the Institute of Bast Fiber Crops, Chinese Academy of Agricultural Sciences, Changsha, China (N 28° 12' 35.91", E 112° 41' 51.07"). The seeds of Zhongzhu NO.1 were obtained from the Seed resource bank of the Institute of Bast Fiber Crops. The experiment commenced in March 2008 and proceeded up to June 2017, during which a single-variety Zhongzhu NO.1 was grown through a continuous cropping system (3 seasons per year, 10 consecutive years). The test field was 15 acres (areas were close to each other with a similar initial soil structure). All the acres were farmed under the same cropping system and conditions, including annual planting time, harvest time, fertilization and irrigation measures, etc. Routine agronomic management and fertilization were performed annually during the ramie growing period. In the later period of the experiment, we found that the ramie planted in some portions of the land grew well (after 10 years of continuous cropping, some ramie were healthy, and were defined as Health-group), while the ramie planted the other portions had obvious growth obstacles due to continuous cropping (mainly characterized by short plants and low fiber production, and were defined as Obstacle-group). Rhizosphere soil samples were collected from the Health-group and Obstacle-group whereby a total of 30 soil samples were obtained in each group on June 23, 2017. The fields in the two groups were about 10 m apart, and each field was separated by a 1.5-m-wide ditch. Every 5 samples were pooled, and a total of 6 rhizosphere soil samples were prepared. The pooled soil samples in each group were divided into 2 portions, for metagenomics sequencing analysis (randomly selecting three samples, n = 3) and metabolome analysis, respectively. We also collected the ramie root samples. A total of 30 root samples were collected, every 5 samples were pooled, and a total of 6 ramie root samples were prepared. All the 6 root samples were used for metabolome analysis. All collected samples were stored at – 80 °C immediately after preparation.

Detection of soil chemical properties. The soil chemical properties, including the contents of available nitrogen (N), phosphorus (P), potassium (K), and total N, P, K were measured according to the methods reported previously⁴⁹. In each group, 15 samples were randomly collected from 0 to 60 cm soil, mixed into three pools, air dried and ground. Soil total N, P and K was detected using the GB 7173-1987 (NY/T 53-1987, semi-micro Kjeldahl method), GB 9837-1988 (NaOH melt-colorimetry method with molybdenum and antimony) and GB 9836-1988 (NY/T 87-1988) national standard of China, respectively. Soil available P and K content was measured using the Bray and Kurtz 1 extraction solution (0.025 M HCl in 0.03 M NH4F) and colorimetry (660 nm), DB13/T 844–2007 (CH₃COONH₄ extraction-flame photometric method), respectively. Each determination was repeated at least three times.

Determination of the plant growth and ramie quality indexes. For the determination of ramie growth and quality, the plant height (cm), stem diameter (mm), bark thickness (mm), fresh bark weight (g), fresh and dry fiber weight (g) were measured and recorded according to the previous report⁵⁰.

DNA extraction, library preparation, and Illumina sequencing. Rhizospheric soil samples (n=6) were collected from the two groups in May 2017. Soil samples were subjected to DNA extraction using a MoBio PowerSoil DNA Isolation Kit (MoBio Laboratories, Carlsbad, CA, USA) following the manufacturer's instructions. DNA concentration, purity and integrity were determined using NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA) and 1% agarose gel electrophoresis. DNA samples were fragmented, ligated to adapters and enriched standardly. TruSeq DNA PCR-Free Sample Preparation Kit (Illumina, San Diego, CA, USA) was used for the construction of DNA libraries, which were then evaluated using Qubit 2.0 Fluorometer (Thermo Scientific, MA, USA) and Agilent Bioanalyzer 2100 system (Agilent Technologies, Santa Clara, CA, USA). Illumina HiSeq2500 platform (250 bp paired-end reads) was employed for the sequencing.

Sequencing data preprocessing. The paired-end raw reads in the format of FASTQ were quality controlled using Trimmomatic by trimming and filtering adaptors and low quality reads. SOAP denovo (v4.5.4; https://soap.genomics.org.cn/) was used for the metagenome assembly⁵¹. ORF prediction of assembled scaffolds was performed using prodigal (v2.6.3) and was translated into amino acid sequences. All effective tags were clustered by CDHIT (v4.5.4; https://www.bioinformatics.org/cd-hit/)⁵². With the thresholds of 95% identity and 90% coverage, a non-redundant gene set was assembled. The relative abundance of each gene in the clean reads was analyzed using fast gapped-read alignment with Bowtie2 (v2.1.0)⁵³ by blasting to the non-redundant gene catalog. GeneMark.hmm (v3.26) was used to predict the potential protein-coding region in a DNA sequence. Blastp (v2.2.28 + , https://blast.ncbi.nlm.nih.gov/Blast.cgi) was employed for the alignment in NCBI nonredundant (NR) database (e-value = 1e–5). Then the sum of the gene abundance was calculated and the abundance of microbe at different taxonomies was obtained accordingly. Heatmap of the abundance profile was constructed

using R (v3.2.0). The annotation of the gene products were performed in databases including NR, SWISSPROT, COG (Clusters of Orthologous Groups of proteins), Gene Ontology (GO) and KEGG with the threshold of $e \le 1e-5$. PCoA and NMDS analysis was performed based on the Euclidean distance algorithm to identify the beta diversity among samples and groups.

The differentially expressed genes in metagenome between the two groups were identified using ANOVA in QIIME (V1.8, https://qiime.org/scripts/alpha_rarefaction.html). The KEGG pathways associated with differentially expressed genes were identified in KEGG database (https://www.genome.jp/KEGG/pathway.html) with the threshold of p < 0.05.

Sample preparation for metabolome analysis. A total of 60 mg ramie root samples were collected from ramie plants monocultured continuously for 10 years in soil without (Health-group, n=6) and with (Obstacle-group, n = 6) continuous cropping, so did as the rhizosphere soil samples (n = 12, 1 g). 60 mg of root samples were placed into 1.5 mL EP tube, incubated with 360 µL pre-cooled methyl alcohol and 40 µL L-2-chloro-phenylalanine (Shanghai hengchuang biotechnology co., LTD., Shanghai, China) at 80 °C for 2 min, followed by grinding into homogenate using an automated grinder (JXFSTPRP-24/32, Shanghai Netcom Industries Development Co., Ltd. Shanghai, China; at 60 Hz for 2 min) and ultrasonic cleaner (SB-5200DT, Ningbo Scientz Biotechnology Co., Ltd, Ningbo, China; for 30 min). Samples were then incubated with 200 µL chloroform with sonication for 30 min, centrifuged at 12,000 rpm, 4°C for 10 min, and dried in a vacuum centrifuge concentrator (LNG-T98, Huamei Biochemistry Instrument, Taicang, China). For soil samples, 1 mL 50% pre-cooled methyl alcohol and $20 \,\mu\text{L}$ L-2-chloro-phenylalanine were used for sample extraction, followed by setting at $-20 \,\text{°C}$ for 2 min and grind (at 60 Hz for 2 min). Homogenates were centrifuged and supernatants were collected, dried, dissolved into 50% pre-cooled methyl alcohol, vortex oscillated for 60 s, ultrasonic-treated for 30 s, and finally centrifuged (at 12,000 rpm, 4°C for 10 min) and dried. All samples were subjected to treatments with methoxamine hydrochloride pyridine under rotary (2 min) and concussion (at 37 °C for 90 min); and further incubation with BSTFA (containing 1% TMCS) and 20 µL n-hexane by rotary (2 min) and concussion (at 70 °C for 60 min).

Preparation of quality control (QC) samples. QC samples were prepared for the non-targeted metabolomics analysis by pooling equal aliquots of individual sample (n = 6) within each group. Three QC samples were prepared.

Non-targeted chromatographic parameters. The analytical instrument used in this experiment is a 7890b-5977a gas chromatograph-mass spectrometer (GS/MS) from Agilent Technologies Inc. (USA). The chromatographic conditions are: DB-5MS column (30 m \times 0.25 mm \times 0.25 µm; Agilent J&W Scientific, Folsom, CA, USA); high purity helium (99.99%) with a flow rate of 1.0 mL/min; injection temperature of 260 °C. The temperature of column was increased from 60 to 125 °C at a speed of 8 °C/min, then increased to 210 °C at a speed of 4 °C/min, to 270 °C at a speed of 5 °C/min, to 305 °C at 10 °C/min and kept for 3 min. MS conditions are: electron impact ion (EI) source at 230 °C; quadrupole temperature at 150 °C; 70 eV high-energy. EI-MS spectra were recorded at a scan range of m/z 50–500. System stability and accuracy was validated using QC samples with an interval of 5 samples.

Non-targeted metabolomics data analysis. MS raw data (total ion current, TIC) was converted into file format using ChemStation (version E.02.02.1431, Agilent Technologies Inc). ChromaTOF (version 4.34, LECO, St Joseph, MI) was used to analyze the data, and NIST and Fiehn database were used for the annotation of the metabolites. After alignment with Statistic Compare component, the 'raw data array' (.cvs) was obtained from raw data including peak names, retention time-m/z and peak intensities. All internal standards and pseudo positive peaks were removed. Data was transformed by log2 and then imported into SIMCA software package (14.0, Umetrics, Umeå, Sweden). Unweighted principle component analysis (PCA) and (orthogonal) partial least-squares-discriminant analysis (OPLS-DA, with sevenfold cross validation and response permutation testing, 200 times randomly permutated) were performed to visualize the metabolism difference between groups. Metabolites with variable important in projection (VIP) > 1 and p value <0.05 by two-tailed Student's t-test were used for identification of differential metabolites. Metabolites between groups with |fold change (FC)|≥ 1 were considered as differential metabolites. The KEGG pathways associated with the differential metabolites were identified from KEGG database (https://www.genome.jp/KEGG/pathway.html) with the threshold of p <0.05.

Statistical analysis. GraphPad Prism 6 software was used for data statistical analysis. Data were expressed as the mean \pm SD and differences between groups were analyzed by t-test. A value of p < 0.05 was considered statistically significant.

Data availability

The DNA-seq data generated and analysed during the current study have been deposited at the NCBI Sequence Read Archive with the Bioproject ID: PRJNA560333.

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Author contributions

Y.W. planned and designed the research. T.L. and B.G. performed experiment. F.L. conducted the fieldwork, and prepared the reagents of experiments. X.B. performed the data analysis. S.Z. wrote the manuscript. All authors have read and approved the manuscript.

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Competing interests

The authors declare no competing interests.

Additional information

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