RESEARCH ARTICLE

Endosphere microbiome comparison between symptomatic and asymptomatic roots of *Brassica napus* infected with *Plasmodiophora brassicae*

Ying Zhao¹,², Zhixiao Gao¹,², Binnian Tian¹,², Kai Bi¹,², Tao Chen², Huiquan Liu³, Jiatao Xie¹,², Jiasen Cheng¹,², Yanping Fu², Daohong Jiang¹,²*

¹ State Key Laboratory of Agricultural Microbiology, Huazhong Agricultural University, Wuhan, Hubei Province, P-R China, ² The Provincial Key Lab of Plant Pathology of Hubei Province, College of Plant Science and Technology, Huazhong Agricultural University, Wuhan, Hubei Province, P-R China, ³ State Key Laboratory of Crop Stress Biology for Arid Areas, College of Plant Protection, Northwest Agriculture and Forestry University, Yangling, Shanxi, P-R China

* daohongjiang@mail.hzau.edu.cn

Abstract

Clubroot caused by *Plasmodiophora brassicae*, is a severe disease of cruciferous crops that causes large hypertrophic galls in the roots. The plant microbiome is important for growth promotion and disease suppression. In this study, using 16S rRNA and internal transcribed spacer (ITS) sequencing techniques, we compared the endosphere microbiome of symptomatic and asymptomatic *B. napus* roots infected with *P. brassicae* collected from the same natural clubroot field. The results showed that the microbial population and its relative abundance in the asymptomatic roots was far higher than that in the symptomatic roots, and that many microorganisms in asymptomatic roots have biological control and plant growth promotion functions that may be related to clubroot symptoms. These results suggest the importance of the endosphere microbiome in clubroot disease and provide potential biocontrol resources for its prevention.

Introduction

Although studies have been conducted for many years, the plant microbiome has gained public attention as a new concept only recently [1, 2]. Turner et al. considered the plant microbiome to mainly include phyllosphere microorganisms, rhizosphere microorganisms, and endogenous microorganisms [3]. Edwards et al. clarified that spatial resolution distinguished three root-associated compartments: the endosphere (root interior), the rhizoplane (root surface), and the rhizosphere (soil close to the root surface). Compared to the rhizosphere and the rhizoplane, the endosphere is highly specific in terms of the microbial communities that it contains [4].

The plant microbiome has been identified as an important determinant of plant health [5]. In its capacity to do harm, the microbiome contains many plant pathogens such as...
Magnaporthe oryzae, Sclerotinia sclerotiorum, Botrytis cinerea, and Fusarium sp. In terms of benefits, among others, it contributes to the promotion of growth and resistance to disease [2, 5, 6]. Textbook examples of the microbiome contributing to plant growth promotion include the assistance by mycorrhiza and rhizobia to plants in phosphorus and nitrogen uptake [7, 8]. Some nitrogen-fixing bacteria belong to Azorhizobium, Bradyrhizobium, Ensifer, Mesorhizobium, Rhizobium, and Sinorhizobium [9], and most mycorrhizae belong to the phylum Glomeromycota, which only comprises endophytes known as arbuscular mycorrhizal fungi [10, 11]. Many microorganisms in the microbiome have been used to prevent and control plant diseases. For example, Trichoderma controls a broad range of diseases such as Fusarium wilt, grey mold, and white rot disease [12, 13]; Pseudomonas can be used to control cotton blight and other soil-borne diseases [14, 15]; and Streptomyces has been useful in controlling Phytophthora root rot [16]. Overall, with increasing research on this topic, the plant microbiome is gaining more attention.

Clubroot is caused by a protist, Plasmodiophora brassicae, and is a significant disease in cruciferous crops [17, 18]. It endangers plants by forming finger, bar, or spherical galls in their roots. In a field survey conducted, in the same field contaminated with P. brassicae, we found that some rapeseed seedlings were seriously diseased with typical symptoms whereas some of the neighboring seedlings showed no obvious symptoms. One of the reasonable explanations for this phenomenon is the uneven distribution of the pathogen throughout the field; however, P. brassicae could be detected in the root cortical cells of asymptomatic plants using polymerase chain reaction (PCR) amplification. Therefore, symptoms manifestation may be associated with other factors. In previous studies, a higher frequency of Curtobacterium flaccumfaciens was observed in asymptomatic plants than that in symptomatic plants, which may contribute toward resistance to citrus variegated chlorosis (CVC) [19]. Asymptomatic or symptomatic CVC in citrus plants may have a relationship with the microorganism population balance (mainly Methylobacterium spp., C. flaccumfaciens, and X. fastidiosa) [20]. The frequency of bacterial isolates possessing various plant-beneficial properties was higher in asymptomatic citrus plants affected by Huanglongbing than that in symptomatic citrus plants [21]. Bruz indicated that diverse fungi and bacteria colonized the woody tissues of asymptomatic and symptomatic esca-diseased grapevines [22]. Therefore, we assume that the microbiome might be associated with the asymptomatic and symptomatic presentation of clubroot under certain conditions.

While 16S rRNA is the most suitable appraisal indicator for the systemic development and classification of bacteria, as the use of the internal transcribed spacer (ITS) region in fungal identification has also gradually increased [23, 24]. Sequencing of 16S rRNA and ITS has become an important means for studying the plant microbiome [25–29]. In this study, the root endosphere microbiome in symptomatic and asymptomatic seedlings was analyzed by sequencing the 16S rRNA and ITS sequencing to clarify the relationship between symptoms and microbiome.

Materials and methods
Preparation of samples for sequencing

P. brassicae-infected B. napus samples were collected (December 18, 2014) from a natural field large approximately 3 Mu (1 Mu = 0.0667 hectares) in Dangyang County, Hubei Province, PR China (30.50 N, 111.47 E), where clubroot disease had been occurring for five years. The plants were at the five- to ten-leaf stage. Two sample groups were collected: RS1 (i.e., asymptomatic root samples, no obvious swelling) and RS2 (i.e., symptomatic root samples, swollen). In each group, 30 samples were randomly collected and divided equally into three sub-groups. Each
sub-group (including 10 plant roots) was regarded as a single sample for sequencing (in total six samples, denoted as RS1.1, RS1.2, RS1.3, RS2.1, RS2.2, and RS2.3).

**PCR and quantitative (q)-PCR detection**

All roots were carefully washed with tap water before being peeled with a sterilized razor. The genomic DNA of all samples was extracted using the CTAB (hexadecyl trimethyl ammonium bromide) method [30]. DNA concentration and purity were monitored on 1% agarose gel. DNA was diluted to 1 ng/μL using sterile distilled water. The PCR protocol described by Wal lenhammar and Arwidsson (2001) was used for detection of *P. brassicae* in RS1 and RS2 samples. PCR amplification was performed using the primers PbITS6: 5′-CAACGAGTCAGCTTG AATGC-3′ and PbITS7: 5′-TGTTTCGGCTAGGATGTTTC-3′ [31]. The conditions for PCR amplification included a denaturation step at 95˚C for 5 min and 32 cycles of 94˚C for 30 s, 59˚C for 30 s, and 72˚C for 1 min. The products of PCR amplification were sequence analyzed by BGI (Beijing Genomics Institute, Beijing).

For quantitative detection of *P. brassicae* in RS1 and RS2 samples, q-PCR analysis was performed using the primers Pb actin-F: 5′-CACCGACTACCTGATGAA-3′ and Pb actin-R: 5′-CAGCTTCCTCTTGATGTC-3′, according to the method described by Chen et al. [21]. The DNA samples were diluted to 500 ng/μL with sterile distilled water and 10-μL reactions were analyzed in triplicate using the CFX96 Real-Time PCR Detection System (Bio-Rad, USA). Each reaction mixture contained 5 μL of 2× SYBR Green Super mix (Bio-Rad, USA), 1 μL of sample DNA, 0.15 μL of forward primer, and 0.15 μL of reverse primer (10 μmol/L). Sterile distilled water was added to make up the final volume. The program was as follows: denaturation at 95˚C for 3 min followed by 49 amplification cycles of 95˚C for 15 s, 58.5˚C for 20 s, and 72˚C for 15 s. A melt curve was generated to verify the specificity of amplification from 65˚C to 95˚C with an increment of 0.5˚C per cycle, with each cycle held for 5 s. The primer sequences are provided in S1 Table. The *B. napus* actin gene (primers: Bn actin-F: 5′-TGAAG ATCAAGGTGGTCGCA-3′ and Bn actin-R: 5′-GAAGGCAGAAACACTTAG AAG-3′) was used as the internal control for normalization.

**16S rRNA and ITS sequencing**

Using the genomic DNA as a template, the 16S rRNA gene and ITS fragments were amplified using specific primers (515F: 5′-GTGCCCAGCMGCCGCGGTAA-3′ and 806R: 5′-GGAACTACHVGGGTATCTTAAAC-3′ for 16S rRNA; ITS1-1F-F: 5′-CTTGTCATTGATATGGCTGC-3′ and ITS1-1F-R: 5′-GCTGCGTTCTTCATCGATGC-3′ for ITS) tagged with abarcode. PCR was conducted in a total reaction volume of 30 μL containing 15 μL of Phusion High-Fidelity PCR Master Mix (New England Biolabs), 0.2 μM forward and reverse primers, and approximately 10 ng of template DNA. The thermal cycling consisted of initial denaturation at 98˚C for 1 min, followed by 30 denaturation cycles at 98˚C for 10 s, annealing at 50˚C for 30 s, elongation at 72˚C for 60 s, and finally 72˚C for 5 min. An equal volume of 1X loading buffer (containing SYBR green) was mixed with the PCR products and electrophoresed on 2% agarose gel for detection. Samples with bright main bands between 400 and 450 bp were chosen for further experiments. The PCR products were mixed in equidensity ratios and purified using the Gene JET Gel Extraction Kit (Thermo Scientific). Sequencing libraries were generated using the NEB Next Ultra DNA Library Prep Kit for Illumina (NEB, USA) according to the manufacturer’s recommendations, and index codes were added. The quality of the library was assessed using the Qubit 2.0 Fluorometer (Thermo Scientific) and the Agilent 2100 Bioanalyzer system. The library was sequenced on an Illumina HiSeq 2500 platform by Beijing Novogene Bioinformatics Technology Co. Ltd. The raw sequencing data for 16S rRNA and ITS
sequencing results have been deposited at the Sequence Read Archive (SRA, http://www.ncbi.nlm.nih.gov/sra/) under accession numbers PRJNA388067 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA388067) and PRJNA388081 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA388081), respectively.

Processing of the sequencing data
Since the original raw data from the Illumina sequencing platform included some low-quality points, pre-processing was necessary for further analysis. The specific processing steps were as follows:

1. Data resolution: according to barcode sequences, each sample sequence was resolved from the raw data and the barcode sequences and primers were truncated;

2. Raw read splicing: the split data of each sample read were spliced using FLASH (V1.2.7, http://ccb.jhu.edu/software/FLASH/) for Mosaic [32], as it was the original tag data (raw tags);

3. Tag filtering: to obtain clean tags, the raw tags required stringent filter processing [33] according to the methods described by Caporaso et al. [34]. The raw tags were truncated at the first low quality base site when the continuous length of low-quality value (quality value ≤3) bases was up to three. The truncated tags in which continuous high quality base length was less than 75% of the tag lengths were excluded.

4. Chimera removal: the clean tags were contrasted with the Gold database (http://drive5.com/uchime/uchime_download.html) using the UCHIME Algorithm (http://www.drive5.com/usearch/manual/uchime_algo.html) to test the chimeric sequences. Final valid data (effective tags) were obtained after removing the chimeric sequences [35, 36]. Finally, the effective tags were used for subsequent analysis.

Operational taxonomic units statistics and classification
To study species diversity, the effective tags were clustered using the UPARSE software [37]. Sequences with ≥97% similarity were assigned to the same operational taxonomic units (OTUs). Since the number of taxon tags was different for each of the six sequenced samples, the OTUs clustering results were standardized for subsequent analysis according to the minimum value of the sequence. The OTUs statistical information of the RS1 and RS2 samples was rendered using the Wayne figure.

Sequences in the same OTUs are considered as a corresponding taxon. In the process of assigning OTUs, the highest frequency sequence in the same OTUs was selected as the representative sequence. The representative sequences were used for classification. The 16S rRNA sequences were assigned with the Greengenes database using the RDP Classifier (threshold value is 0.8–1), and the ITS sequences were assigned with the UNITE/INSDC database using QIIME (threshold value is 0.8–1) [38–40].

Microorganism community statistics and cluster analysis
After assigning the corresponding database, the OTUs were classified into different classification levels (i.e., Kingdom, Phylum, Class, Order, Family, Genus, and Species). The level into which more than 80% of the OTUs could be classified was chosen for microorganism community analysis. The relative microorganism abundance in each sample was determined according to the number of OTU reads.
To determine the difference in the microorganisms between the two groups, microorganism community cluster analysis was performed based on the Z value using the software MeV 4.9. The Z value of one sample in a classification was equal to the value calculated as the difference between the relative abundance in the sample and the average relative abundance in all samples, divided by the standard deviation of all samples.

**Results**

**Symptoms of clubroot in the field**

The symptoms of the clubroot samples are shown in Fig 1. The roots of RS1 samples were not obviously swollen and the lateral roots remained healthy, whereas the roots of RS2 samples were obviously swollen, and few lateral roots could be observed (Fig 1A). PCR amplification results showed that *P. brassicae* could be successfully detected in both RS1 and RS2 samples (Fig 1B). To evaluate the relative *P. brassicae* content in RS1 and RS2 samples, expression of the *P. brassicae* actin gene was measured using q-PCR. In the symptomatic (RS2) samples, the *P. brassicae* numbers were thousand-fold higher than those in the asymptomatic (RS1) samples (Fig 1C).

**Sequencing quality assessment**

Raw data were generated by Illumina sequencing. Impurities such as joint and primer sequences, and low-quality reads were first removed to obtain effective tags for subsequent

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**Fig 1. Symptoms of Brassica napus samples and PCR detection.** (a), Symptoms of *B. napus* samples collected from the same clubroot field; photos of asymptomatic (above) and symptomatic (underneath) samples. (b), Detection of *P. brassicae* in surface-peeled roots of RS1 and RS2 samples; Lane 7: sterile water; Lane 8: *P. brassicae* DNA; Lane M: DNA marker. (c), q-PCR detection of *P. brassicae* in surface-peeled roots of RS1 and RS2 samples; The *B. napus* actin gene served as the internal control for normalization.

https://doi.org/10.1371/journal.pone.0185907.g001
analyses. Sequencing quality was determined using the number of effective tags, their proportion in the raw data, and the base percentage of the sequencing error rate, which is less than 0.1% (Q30). In the case of 16S rRNA sequencing (S1 Table), 45471, 27897, 82715, 57064, 25939, and 20842 tags were generated as raw data for the six samples (RS1.1, RS1.2, RS1.3, RS2.1, RS2.2, and RS2.3, respectively). For each sample, the percentage of effective tags in the raw data was up to 92%, and the Q30 values were above 96%. In the case of ITS sequencing (S2 Table), 66344, 66782, 55796, 44365, 44576, and 66344 tags were generated as raw data for the six samples (RS1.1, RS1.2, RS1.3, RS2.1, RS2.2, and RS2.3, respectively). The percentage of effective tags in the raw data was up to 96% and the Q30 values were above 99% in each sample. This demonstrated that the data were suitable for subsequent analyses.

OTU statistics and analysis

All effective tags from the six samples were clustered into OTUs according to sequence similarity (≥97%) using the UPARSE software (Edgar & Robert, 2013). According to the OTU statistical results, most effective tags (Taxon Tags) could be clustered to OTUs, with the exception of only a few data (Unclassified Tags and Unique Tags) (S1 Fig). The number of OTU clusters formed for the RS1 samples (RS1.1, RS1.2, and RS1.3) as a result of 16S rRNA and ITS sequencing were 114, 108, and 123, and 135, 133, and 148, respectively; and for the RS2 samples (RS2.1, RS2.2 and RS2.3) the number of OTU clusters formed were 93, 63, and 88, and 87, 81, and 69, respectively (S1 Fig).

Since the number of taxon tags in each sequencing sample was different, the OTU clusters formed above could not be directly used to compare the two samples. The OTU clustering results were standardized for subsequent analysis on the basis of the minimal sequence read number. The standardized OTU information for RS1 and RS2 samples was rendered using the Wayne figure (Fig 2). In both RS1 and RS2 groups, three samples were used for sequencing. Shared and unique OTUs were found among the three samples. To ensure the accuracy and completeness of the data, OTUs that existed in at least two of the three samples were chosen for further study. In the RS1 group, 96 OTUs (49 shared and 47 in two samples) from 16S rRNA and 110 OTUs (61 shared and 49 in two samples) from ITS were used for subsequent studies (Fig 2A and 2D). In the RS2 group, 58 OTUs (31 shared and 27 in two samples) from 16S rRNA and 54 OTUs (31 shared and 23 in two samples) from ITS were used for further studies (Fig 2C and 2F). Comparison of the OTUs in at least two samples from each group showed that, in 16S rRNA 48 OTUs were common to both groups, 48 OTUs were only in the group RS1, and 10 OTUs were only in the group RS2 (Fig 2B). In ITS, 48 OTUs were common to both groups, 62 OTUs existed only in the group RS1, and 6 OTUs were only in the group RS2 (Fig 2D). Therefore, 106 OTUs from 16S rRNA and 116 OTUs from ITS were used for further analysis. The number of OTUs in the group RS1 was obviously greater than that in the group RS2, both from 16S rRNA and ITS sequencing results.

OTU classification and microorganism community analysis

After the OTU cluster analysis, the same OTU sequences were assumed to derive from a certain taxon, and a representative sequence for each OTU was classified into different classification levels (Kingdom, Phylum, Class, Order, Family, Genus, and Species).[38, 39] The OTU classification information and statistics results are shown in S3 and S4 Tables and S2 Fig. More than 80% of the OTUs could be classified at the family level from the 16S rRNA sequencing results (S2A Fig) and at the genus level from the ITS sequencing results (S2B Fig); thus, the family and genus taxa were used to analyze the bacterial and fungal communities. The 106 OTUs from 16S rRNA and 116 OTUs from ITS as obtained above were classified into 45
bacterial families and 71 fungal genera (S3 and S4 Tables). To determine the differences in endosphere microorganism communities between the asymptomatic (RS1) and symptomatic (RS2) *B. napus* samples, the relative abundance of bacteria and fungi in each sample was calculated based on the number of OTU reads and used for analysis (S3 and S4 Tables).

Based on the classification results, 39 and 28 bacterial families were found in the asymptomatic RS1 and symptomatic RS2 groups, respectively (S3 Table). The two most abundant families (i.e., Streptophyta and Mitochondria) were excluded from the analysis. Except these two families, the relative abundances of the top 15 bacterial families in each sample is shown in Fig 3A. The most abundant family was Oxalobacteraceae, followed by Pseudomonadaceae, Comamonadaceae, Xanthomonadaceae, Methylophilaceae, Rhizobiaceae, Flavobacteriaceae. The relative abundance of these bacteria in the asymptomatic samples (RS1 ≥2) and the RS2 (asymptomatic) samples (RS2 ≥2) detected from 16S rRNA (b) and ITS (e) sequencing results.

https://doi.org/10.1371/journal.pone.0185907.g002

**Fig 2. Number of operational taxonomic units (OTUs) in different samples.** Venn diagrams were used to show the number of standardized OTUs in the three RS1 (asymptomatic) samples (RS1.1, RS1.2 and RS1.3) detected from 16S rRNA (a) and ITS (d) sequencing results, as well as the OTU numbers in the three RS2 (asymptomatic) samples (RS2.1, RS2.2 and RS2.3) detected from 16S rRNA (c) and ITS (f) sequencing results; The Venn diagrams were also used to express the number of OTUs present in at least two samples from each group between the RS1 (asymptomatic) samples (RS1 ≥2) and the RS2 (asymptomatic) samples (RS2 ≥2) detected from 16S rRNA (b) and ITS (e) sequencing results.
The ITS sequencing results indicated the endosphere fungal communities and *P. brassicae*. The most abundant species in all six samples was undoubtedly *P. brassicae*, with its content in the symptomatic samples (0.986, 0.991, and 0.989) was slightly higher than that in the asymptomatic samples (0.952, 0.975, and 0.908). Based on the classification, 65 and 39 fungal genera were found in RS1 and RS2 groups, respectively (*S4 Table*). The most abundant fungal genera were Un—s-Tremellomycetes sp., *Tetracladium*, Un—s-Nectriaceae sp., *Verticillium*, *Fusarium*, *Gibberella*, *Alternaria*, Un—s-Sordariomycetes sp., and *Mortierella*. The relative abundance of these fungi in the asymptomatic samples was also much greater than that in the symptomatic samples (*Fig 3B*). In the asymptomatic samples, other fungi (0.013, 0.007, and 0.026) were present in greater amounts than those in the symptomatic samples (0.003, 0.003, and 0.003).

**Microorganism community clustering and specific microorganism analysis**

For further analysis of the differences in the endosphere communities between the asymptomatic and symptomatic samples, the relative abundance of all bacterial families and fungal genera was clustered. For the bacterial endosphere community, 16 families existed only in the asymptomatic samples, five families only in the symptomatic samples, and 21 families common to both samples (*Fig 4*). Among the 16 families that were present only in the asymptomatic samples, Streptomycetaceae, Pseudonocardiaceae, Rhodospirillaceae, Sphingomonadaceae, and Verrucomicrobiaceae were present in all three groups (*Fig 4AI*), and the remaining families (i.e., Intrasporangiaceae, Nocardioidaceae, Mycobacteriaceae, Cryomorphaceae, Burkholderiaceae, Turicibacteraceae, Alteromonadaceae, Bacillaceae, Chitinophagaceae, Nannocystaceae, and Promicromonosporaceae) were present in two of the three samples. The families present only in the symptomatic samples included Peptostreptococcaceae, Moraxellaceae, Acetobacteraceae, Deinococcaceae, and an unclassified bacterium (auto67_4W) (*Fig 4AII*).

Of the 21 shared families, the relative abundance of Nocardiaceae, Phyllobacteriaceae, and Enterobacteriaceae in the symptomatic samples was greater than that in the asymptomatic samples (*Fig 4BII and 4BIII*). The abundance of Methylophilaceae, Oxalobacteraceae, and Microbacteriaceae was similar in all the samples (*Fig 4BIII*). The remaining 15 families were more abundant in the asymptomatic samples than in the symptomatic samples, especially Xanthomonadaceae, Comamonadaceae, Flavobacteriaceae, Sphingobacteriaceae, Rhizobiales, Caulobacteraceae, and Bradyrhizobiaceae (*Fig 4BIII, 4BIV and 4BV*).

In the fungal endosphere community, 30 genera were unique to the asymptomatic samples, including *Tetraplosphaeria*, *Phaeoseptoria*, *Cistella*, *Phoma*, *Ilyonectria*, *Glomus*, *Ambispora*, and Un—s-Hypocreales sp. (*Fig 5AI*). Only four genera were detected in the symptomatic samples, namely Un—s-Pleosporales sp. REFI07, *Aspergillus*, Un—s-fungal sp. DG16, and Un—s-Helotiales sp. r427 (*Fig 5AII*). Among the 37 shared genera, seven, including *P. brassicae*, were much more abundant in the symptomatic samples than in the asymptomatic samples (*Fig 5BIII and 5BIV*). The abundance of the remaining 30 genera including *Tetracladium*, *Fusarium*, *Alternaria*, Un—s-fungal sp. 3 EO_2010, and *Sporobolomyces* was much greater in the asymptomatic samples than in the symptomatic samples (*Fig 5BIII and 5BIV*).

Overall, the bacterial and fungal endosphere communities in the asymptomatic samples were more abundant than those in the symptomatic samples. Among the unique and highly
abundant bacteria and fungi in the asymptomatic samples, six bacterial families and six fungal genera have been attributed with biological control functions, three bacterial families with...
plant growth activity, and four bacterial families and seven fungal genera causing plant disease. Interestingly, Rhizobiaceae and Bradyrhizobiaceae were detected in both the asymptomatic and symptomatic samples, and their relative abundance in the asymptomatic samples was much greater than that in the symptomatic samples. Furthermore, *Glomus* was detected only in the asymptomatic samples, whereas *Aspergillus* was detected only in the symptomatic samples (Tables 1 and 2).

**Discussion**

Recently, research on microbiome has attracted increasing attention. Previously, 16S rRNA and ITS sequencing have been widely used in plant microbiome studies [35–42]. In this study, we analyzed the root endosphere microbiome of *B. napus* infested with *P. brassicae* using 16S rRNA and ITS sequencing, and tailored the sequencing quality to the requirement of the analysis. The 16S rRNA sequencing results revealed 106 OTUs (OTUs that existed in at least two of the three samples), and 116 OTUs were detected from the ITS sequencing results. The majority of OTUs (except OTU3) from the 16S rRNA sequencing results could be classified to the family level and most OTUs from the ITS sequencing results could be classified to the genus level (S2 Fig).

Although the results showed that 16S rRNA and ITS sequencing work well in microbiome research, some difficulties persist. For instance, in the 16S rRNA sequencing results, 11, 15, and 10 OTUs in the asymptomatic samples (RS1.1, RS1.2, and RS1.3) and 9, 6, and 7 OTUs in the symptomatic samples (RS2.1, RS2.2, and RS2.3) could not be classified into any known family. In the symptomatic samples, OTU 3 was classified as an uncultured bacterial clone and its relative content was 70–90% of all the OTUs. The ITS sequencing results also showed 18, 23, and 25 OTUs in the asymptomatic samples (RS1.1, RS1.2 and RS1.3) and 14, 8, and 8 OTUs in the symptomatic samples (RS2.1, RS2.2 and RS2.3) that could not be classified into any known genera (S2 Fig). This might suggest that new microorganisms were being hosted by *P. brassicae*-infected *B. napus* roots including unknown bacteria (i.e., OTU 3). There may be two possible reasons for this. First, our knowledge of microorganisms is still very limited especially for those that cannot be purified and cultivated. Secondly, only 500 bp sequences were created and analyzed in the 16S rRNA sequencing platform, which is a little short for bacterial identification. Moreover, it is difficult to overcome the chloroplast and mitochondrial sequence interference in 16S rRNA sequencing [43]. Moreover, since fungal ITS sequences in the public database are relatively poor compared to the bacterial 16S rRNA gene sequences, it is necessary to include other genes such as EF-1α, β-tubulin, and RNA polymerases to complete the identification [44]. Therefore, achieving a comprehensive understanding of the microbiome and its functions will take a long time.

The main purpose of this study was to clarify the difference between the endosphere microbiomes of symptomatic and asymptomatic *B. napus* roots from the same natural clubroot field, using 16S rRNA and ITS sequencing. Based on the results, we first found that *P. brassicae* was common in both the symptomatic and asymptomatic samples. Second, many bacterial families and fungal genera were unique to the asymptomatic samples. Moreover, most of the shared bacteria and fungi present across both groups were more abundant in the asymptomatic...
samples. These results were similar to those of previous studies. *Pseudomonas syringae* pv. *actinidiae* was detected in both the symptomatic and asymptomatic tissues of kiwifruit [45]. The microbial population balance (mainly *Methylobacterium* spp., *C. flaccumfaciens*, and *X. fastidiosa*) was also related to the CVC asymptomatic or symptomatic presentation in citrus plants [20]. Besides, in previous studies, some beneficial microorganisms were found to be much more salient in asymptomatic plants. For example, numerous plant-beneficial bacterial isolates were found in the asymptomatic plants of Huanglongbing diseased citrus compared to the symptomatic plants [21]; *C. flaccumfaciens* abundance in asymptomatic plants was higher than

### Table 1. Bacterial classification statistics based on function.

| Function                  | Family Name             | Relative content  |
|---------------------------|-------------------------|-------------------|
|                           |                         | RS1 (Avg) | RS2 (Avg) |
| **Biological Control**    | Streptomycetaceae^[50]  | 2.08E−04 | 0         |
|                           | Sphingomonadaceae^[52]  | 1.09E−03 | 0         |
|                           | Intrasporangiaceae^[52] | 1.91E−04 | 0         |
|                           | Bacillaceae^[47]        | 3.47E−05 | 0         |
|                           | Flavobacteriaceae^[51]  | 1.01E−03 | 1.91E−04 |
|                           | Pseudomonadaceae^[52]   | 3.33E−03 | 2.11E−03 |
| **Plant Growth Promotion**| Burkholderiaceae^[44]  | 5.20E−05 | 0         |
|                           | Chitinophagaceae^[43]   | 3.47E−05 | 0         |
|                           | Nannocystaceae^[47]     | 3.47E−05 | 0         |
| **Disease Inducing**      | Enterobacteriaceae^[19] | 6.93E−05 | 2.77E−04 |
|                           | Xanthomonadaceae        | 1.68E−03 | 9.01E−04 |
|                           | Mycobacteriaceae        | 8.67E−05 | 0         |
|                           | Micrococccaceae         | 3.12E−04 | 6.93E−05 |
| **Others**                | Rhizobiaceae            | 1.02E−03 | 3.99E−04 |
|                           | Bradyrhizobiaceae       | 1.04E−04 | 6.93E−05 |

https://doi.org/10.1371/journal.pone.0185907.t001

### Table 2. Fungal classification statistics based on function.

| Function                  | Genus Name             | Relative content  |
|---------------------------|------------------------|-------------------|
|                           |                        | RS1 (Avg) | RS2 (Avg) |
| **Biological Control**    | *Rhodotorula*^[55]     | 6.38E−04 | 9.33E−05 |
|                           | *Sporobolomyces*^[35]  | 1.17E−04 | 2.33E−05 |
|                           | *Sporidiobolus*^[36]   | 8.55E−05 | 0         |
|                           | *Leucoosporidium*^[57] | 7.00E−05 | 0         |
|                           | Un—*s-Sporidiobolales*^[36] | 1.79E−04 | 4.67E−05 |
|                           | Un—*s-Hypocreales* sp. | 3.89E−04 | 0         |
| **Disease Inducing**      | *Fusarium*             | 2.65E−03 | 3.19E−04 |
|                           | *Gibberella*           | 2.49E−03 | 4.43E−04 |
|                           | *Alternaria*^[1]       | 2.37E−03 | 3.19E−04 |
|                           | *Sclerotinia*          | 5.29E−04 | 2.02E−04 |
|                           | *Phoma*                | 1.63E−04 | 0         |
|                           | *Leptosphaeria*        | 9.33E−05 | 0         |
|                           | *Cylindrocarpon*       | 3.89E−05 | 0         |
| **Others**                | *Glomus*               | 7.00E−05 | 0         |
|                           | *Aspergillus*          | 0         | 4.67E−05 |

https://doi.org/10.1371/journal.pone.0185907.t002
that in symptomatic plants, and may contribute to CVC resistance [19]. In our study, many of the unique and highly abundant bacteria and fungi in the asymptomatic samples have been attributed with biological control or plant growth promotion functions. *Bacillus* sp. from the family Bacillaceae, was reported as a biocontrol agent for *P. brassicae* [44, 46, 47]; *Streptomyces* spp. [48, 49] and *Streptomyces griseoruber* A316 [50] from the family Streptomycetaceae have been reported to reduce the severity of clubroot disease; *Flavobacterium hercynium* EPB-C313, an endophytic bacteria of the family Flavobacteriaceae, has been used to control Chinese cabbage clubroot [51]. In this study, we identified 1, 1, and 7 OTUs in the asymptomatic samples belonging to Bacillaceae, Streptomycetaceae, and Flavobacterium families, respectively. Strains of *Sphingomonadaceae* can inhibit *Pythium spinosum*, whereas strains of *Pseudomonadaceae* efficiently control fusarium wilt disease [14, 15, 52]. Species from the Chitinophagaceae and Burkholderiaceae families can promote plant growth [53, 54]. Yeasts from *Sporobolomyces*, *Leucosporidium*, and *Sporidiobolus* can be used to control gray mold [55–57]. In asymptomatic samples, 9 OTUs were detected as belonging to the six above-mentioned families. They may have similar effects on oilseed rape and clubroot disease. Above all, we assume that the differences in the endosphere microbiome of the symptomatic and asymptomatic samples in the endosphere microbiome may influence the development of clubroot symptoms.

Another purpose of this study was to exploit the endosphere microbiome analysis technique to mine microorganism resources for resistance to clubroot disease. Little is known about Rhodocyclaceae, Verrucomicrobiaceae, Pseudonocardiaceae, and “Un—s—” fungi, all of which were only present in the asymptomatic samples, along with the unknown bacteria (OUT 3) in the symptomatic samples. These microorganisms may be potential bio-control resources for preventing clubroot disease. Moreover, in the asymptomatic *B. napus* root samples, a variety of pathogenic plant microorganisms from other crops were detected including Xanthomonadaceae, Fusarium, Gibberella, Alternaria, Sclerotinia, Phoma, Leptosphaeria, and Cylindrocarpon. The impact of these pathogens on non-host plants remains a topic for further study.

A few anaerobic or facultative anaerobic bacteria like Acetobacteraceae [58] were unique to symptomatic *B. napus* root samples, suggesting that *P. brassicae* might consume oxygen or need environmental hypoxia when growing in *B. napus* roots [59]. Interestingly, some fungi from the genus *Aspergillus* are typical saprophytes or opportunistic pathogens [60], and *Aspergillus* could only be detected in the symptomatic samples. Therefore, we postulate that *P. brassicae* needs to decay and disintegrate the host roots with the help of saprophytic microorganisms, in order to release its resting spores into the soil at the later stages of clubroot development. Furthermore, we also found an interesting phenomenon that bacteria from the family Rhizobiaceae and Bradyrhizobiaceae and fungi from the genus Glomus were detected in *B. napus* infested with *P. brassicae*, although *B. napus* is not a plant that hosts rhizobia and arbuscular mycorrhizal fungi. Further analysis on this phenomenon is underway.

In summary, this study was mainly conducted to compare the differences in the endosphere microbiomes of symptomatic and asymptomatic *B. napus* roots collected from the same natural clubroot field using 16S rRNA and ITS sequencing. The microbe population and their relative abundance in asymptomatic roots are far greater than that in the symptomatic *B. napus* roots. Many microorganisms detected in the asymptomatic roots have biological control and plant growth promotion functions. Furthermore, we also found many pathogenic plant microorganisms to which *B. napus* is not a host, and the related mechanism underlying this observation needs further research. These results provide a new basis for studying the endosphere microbiomes of roots and some potential bio-control resources for clubroot prevention.
Supporting information

S1 Fig. The number of Tags and OTUs in different samples. The statistical figures show the number of Tags and OTUs for all samples from the 16S rRNA sequencing results (a) and ITS sequencing results (b). Total Tags, the filtered splicing sequence numbers; Taxon Tags, the number of tags used to build the OTUs and gain classification information; Unclassified Tags, the number of tags used to build OTUs but which did not provide any classification information; Unique Tags, the number of tags for which the frequency is one and cannot be clustered into OTUs; OTUs, the final OTU numbers.

S2 Fig. The number of OTUs at each sample classification level. (a), The OTUs were classified by the taxonomic information into different bacterial classification levels detected in the 16S rRNA sequencing results. The above row of numbers refers to the number of OTUs classified to the bacterial Kingdom level and the lower row of numbers refers to the number of OTUs classified to the bacterial Family level. (b), The OTUs were classified by the taxonomic information into different fungal classification levels detected in the ITS sequencing results. The above row of numbers refers to the number of OTUs classified into the fungal Kingdom level and the lower row of numbers refers to the number of OTUs classified into the fungal Genus level.

S1 Table. Data preprocessing statistics and quality analysis of rapeseed involving 16S rRNA sequencing of the root. Raw data, the number of PE reads; Raw Tags, Tag number of patchwork sequence; Clean Tags: Tags taken off the low quality tag number; Effective Tags, Tag number for aftershock; Base, The number of bases of the Effective Data; AvgLen, The average length of the Effective Tags; Q30, Base percentage of the sequencing error rate is less than 0.1% in Effective Tags; Effective (%), Effective Tags/PE Reads.

S2 Table. Data preprocessing statistics and quality analysis of rapeseed involving ITS sequencing of the root. Raw data: the number of PE reads; Raw Tags: Tag number of patchwork sequence; Clean Tags: Tags taken off the low quality tag number; Effective Tags: Tag number for aftershock; Base: The number of bases of the Effective Data; AvgLen: The average length of the Effective Tags; Q30: Base percentage of the sequencing error rate is less than 0.1% in Effective Tags; Effective (%): Effective Tags/PE Reads.

S3 Table. Classification information and relative abundance of each OTU in the 16S rRNA sequencing results.

S4 Table. Classification information and relative abundance of each OTU in the ITS sequencing results.

Acknowledgments

The study was financially supported by the earmarked fund for China Agriculture Research System (CARS-13) and the Programme of Introducing Talents of Discipline to Universities in China (the 111 Project no. B14032). We also wish to thank the reviewers for their valuable comments.
Author Contributions

Formal analysis: Ying Zhao, Huiquan Liu.
Investigation: Ying Zhao, Zhixiao Gao, Binnian Tian, Kai Bi, Tao Chen.
Methodology: Yanping Fu.
Project administration: Daohong Jiang.
Resources: Jiatao Xie, Jiasen Cheng, Yanping Fu.
Supervision: Daohong Jiang.
Writing – original draft: Ying Zhao, Yanping Fu.
Writing – review & editing: Yanping Fu, Daohong Jiang.

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