Assessment of the Diversity of *Pseudomonas* spp. and *Fusarium* spp. in *Radix pseudostellariae* Rhizosphere under Monoculture by Combining DGGE and Quantitative PCR

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*Radix pseudostellariae* is a perennial tonic medicinal plant, with high medicinal value. However, consecutive monoculture of this plant in the same field results in serious decrease in both yield and quality. In this study, a 3-year field experiment was performed to identify the inhibitory effect of growth caused by prolonged monoculture of *R. pseudostellariae*. DGGE analysis was used to explore the shifts in the structure and diversity of soil *Fusarium* and *Pseudomonas* communities along a 3-year gradient of monoculture. The results demonstrated that extended monoculture significantly boosted the diversity of *Fusarium* spp., but declined *Pseudomonas* spp. diversity. Quantitative PCR analysis showed a significant increase in *Fusarium* oxysporum, but a decline in *Pseudomonas* spp. Furthermore, abundance of antagonistic *Pseudomonas* spp. possessing antagonistic ability toward *F. oxysporum* significantly decreased in consecutively monocultured soils. Phenolic acid mixture at the same ratio as detected in soil could boost mycelial and sporular growth of pathogenic *F. oxysporum* while inhibit the growth of antagonistic *Pseudomonas* sp. CJ313. Moreover, plant bioassays showed that *Pseudomonas* sp. CJ313 had a good performance that protected *R. pseudostellariae* from infection by *F. oxysporum*. In conclusion, this study demonstrated that extended monoculture of *R. pseudostellariae* could alter the *Fusarium* and *Pseudomonas* communities in the plant rhizosphere, leading to relatively low level of antagonistic microorganisms, but with relatively high level of pathogenic microorganisms.

**Keywords:** *Radix pseudostellariae*, DGGE, *Fusarium*, *Pseudomonas*, quantitative PCR
INTRODUCTION

As much as 70% of medicinal plants suffer from consecutive monoculture problem, also known as replant disease or soil sickness. These problems are commonly observed in the production of many Chinese medicinal herbs, including *Radix pseudostellariae*, *Rehmannia glutinosa*, *Panax notoginseng*, etc (Zhang and Lin, 2009). *R. pseudostellariae*, a perennial tonic medicinal plant, belongs to the family Caryophyllaceae with extremely high medicinal value (Zhao W.O. et al., 2015). Consecutive monoculture of this plant in the same field leads to a serious decrease in both quality and yield of roots along with poor plant performance, which severely limited production and utilization of its medicinal plant virtues (Lin et al., 2015). Therefore, it is necessary to explore the mechanism of consecutive monoculture problems affecting the plant and develop effective control strategies for *R. pseudostellariae*.

*Fusarium* species is one of the most abundant, prevalent, and important soil fungi (Damicone and Manning, 1985). It is notorious due to the ability of attacking diversity of host plants and bring upon them diseases like vascular wilts, seedling damping off and rots of stem (Pietro et al., 2003; Punja and Parker, 2009; Chakravarty and Hwang, 2010). Similarly, the soil-borne disease caused by *F. oxysporum* in *R. pseudostellariae* fields were reported (Zhao Y.P. et al., 2015), however, the other *Fusarium* species are often overlooked. Therefore, in order to develop the full potential of the disease-suppressive microbial community in the biological control, we need more information to unravel the different roles of this potentially important species.

In recent years, more attentions were paid to develop the environment friendly and good agriculture practices for disease control. It has become important to explore the nature of microbial diversities in the soil, particularly *Pseudomonas* in different cropping periods or regime (Mendes et al., 2011). *Pseudomonas* species were reported to have a wide range of functional groups, such as plant pathogens (Samson et al., 1998), xenobiotic degraders (Clausen et al., 2002) and plant growth promoters (Patten and Glick, 2002). In addition, *Pseudomonas* species can be used as biological control agents for soil-borne pathogens, including black rot of tobacco, disease of wheat and *Fusarium* wilt (Raaijmakers and Weller, 1998; Patten and Glick, 2002; Mendes et al., 2011).

Recently, the increasing evidences suggest that plant–microbial interactions play many pivotal roles in soil quality and plant health (Lakshmanan et al., 2014; Macdonald and Singh, 2014). Li et al. (2014) reported that the peanut root exudates can selectively inhibit certain communal bacteria, such as *Geobacter metallireducens*, *Mitsuaria chitosanitabida*, and *Burkholderia*, but stimulate the bacterial taxon of *Desulfofotomaculum ruminis* and the fungal taxon *F. oxysporum* in soil. Wu et al. (2016c) found that the amount of two pathogenic fungi (*F. oxysporum* and *Aspergillus flavus*) in the rhizosphere significantly increased after *Rehmannia glutinosa* monoculture. Wu et al. (2015) indicated that long-term continuous cropping of black pepper (*Piper nigrum*) L. could lead to a significant decrease in soil bacterial content, especially the *Pseudomonas* spp., suggesting that the soil microbes might be responsible for soil health.

Denaturing gradient gel electrophoresis (DGGE) is considered as an effective technique to directly analyze the structural and diversity of microbial communities (Kozdrój and van Elsas, 2001). The traditional method of assessing the diversity of *Fusarium* is based on enumeration and isolation of strains which were grown on selective media (Vujanovic et al., 2002). However, morphological identification of *Fusarium* species is a time-consuming and formidable task. Yergeau et al. (2005) described a PCR-DGGE method to detect the presence of multiple *Fusarium* spp. from environmental samples. The method is based on the specific amplification and separation of the transcription elongation factor-1α (EF1α) gene. Similarly, Widmer et al. (1998) designed a primer set (PsR and PsF) which was based on the 16S rDNA gene of *Pseudomonas* spp. in 1998. When combining the PsR and PsF primers, Evans et al. (2004) developed a semi-nested PCR and DGGE to rapidly study the diversity within the genus *Pseudomonas*. Therefore, the role of soil microbial ecology in the prevention and control of plant diseases has been given more attention (Philippot et al., 2013; Cha et al., 2015). However, few studies have been carried out to understand the relationship between *Pseudomonas* and *Fusarium* of *R. pseudostellariae*, and the approaches to overcome diseases associated with this plant.

In this study, DGGE combined with qPCR technique was used to analyze the shifts of *Pseudomonas* and *Fusarium* communities in rhizosphere soil under *R. pseudostellariae* monoculture. Several microorganisms closely related to the problem of prolonged monoculture were isolated and performed for plant-microbe interactions study. Our study can help to illustrate the effects of ecological environment and root exudates on the selection of soil microbes in rhizosphere soil, and provide useful information on potential indigenous microflora for soil remediation and improvement.

MATERIALS AND METHODS

Field Experiment

In this study, the *R. pseudostellariae* cultivar ‘Zheshen 2’ was used as the test material. The experiment was carried out at the experimental station of Fuding City, Fujian Province (27°26′ N, 120°04′ E). The experimental field which previously planted *Oryza sativa* was performed for this study with four treatments: (1) control with no *R. pseudostellariae* cultivation (CK), (2) the newly planted *R. pseudostellariae* cultivation (FP), (3) 2-year consecutive monoculture (SP), (4) 3-year consecutive monoculture (TP). The physical and chemical properties of the soil were detected before the experiment was initiated: total nitrogen of 1.83 g kg⁻¹, available nitrogen of 26.23 mg kg⁻¹, total phosphorus of 0.47 g kg⁻¹, and available phosphorus of 96.34 mg kg⁻¹, total K of 8.46 g kg⁻¹, and available K of 365.21 mg kg⁻¹. The station has a subtropical oceanic monsoon climate, annual
mean temperature at 18.4°C. All treatments were treated with the same fertilization and field management during the experiment.

**Soil Sampling and DNA Extraction**

The above ground or below ground biomass of *R. pseudostellariæ* become significantly different after 5 months of planting (Figure 1A), according to our previously study (Wu et al., 2016b). Therefore, soil samples were randomly collected from five different points at each field on April 22nd, 2015. Additionally, we harvested the plants for yield determination on July 2nd, 2015 (Figure 1B).

Soil samples were collected after digging the plant samples. Firstly, the loosely adhering soil was shaken off, then scraping the soil that was still attached to the root as rhizosphere soil. DNA was immediately extracted from 0.5 g soil sample per treatment using Biofast Soil Genomic DNA Extraction Kit (BioFlux, Hangzhou, China) according to the manufacturer’s protocols. We further determined the DNA concentration using Nanodrop 2000C Spectrophotometer (Thermo Scientific, United States) and then diluted it to 20 ng µL⁻¹.

**PCR-DGGE and Analysis**

*Fusarium*-specific PCR was performed according to the nested amplification of the *Ef1α* gene. The first round of PCR reactions was performed by the *Ef*-1 and *Ef*-2 primers (O’Donnell et al., 1998). It was carried out in 50 µl volumes containing 25 µl of 2 × EasyTaq PCR SuperMix (Transgen Biotech, Beijing, China), 1 µl of each primer and 40 ng template soil DNA. The program of PCR was performed by the following protocol: 95°C for 5 min, 30 cycles of denaturation (95°C for 1 min), annealing (55°C for 1 min), extension (72°C for 1 min), and 1 cycle of final extension (72°C for 10 min). The amplicons were subsequently diluted (1:20) and used for the second PCR reaction via Alfie1-GC and Alfie2 (Yergeau et al., 2005) primers. Second round PCR protocol was similar to the method of the first reaction, except for the annealing (57°C for 50 s) and extension (72°C for 50 s).

*Pseudomonas*-specific PCR was based on the nested amplification of the V6/V7 region of *Pseudomonas* spp. The first round of PCR reactions was used the PsF and PsR primers (Tan and Ji, 2010). PCR reaction was carried out in 50 µl volumes containing 25 µl of 2 × EasyTaq PCR SuperMix (Transgen Biotech, Beijing, China), 1 µl of each primer and 20 ng template soil DNA. The program of PCR was performed by the following protocol: 95°C for 5 min, 30 cycles of denaturation (95°C for 1 min), annealing (64°C for 1 min), extension (72°C for 1 min), and 1 cycle of final extension (72°C for 10 min). The PCR products of first round was used to perform the second PCR reaction and the primers F968-GC1 and PsR were used (Garbeva et al., 2004). The following cycling protocol was performed for the second PCR: 1 cycle of initial denaturation at 94°C for 5 min, 10 cycle of denaturation (94°C for1 min), 1 min at 60°C (every subsequent one using a 0.5°C lower annealing temperature), and 2 min at 72°C, 1 cycle of 95°C for 5 min, 30 cycles of denaturation (95°C for 1 min), annealing (55°C for 1 min), and 1 cycle of final extension (72°C for 10 min). All PCR products were detected using 1.2% agarose gel and purified using a Gel Extraction Kit (OMEGA Bio-Tek, United States) according to the manufacturer’s instructions. The purified PCR products were used to perform DGGE experiments.

**DGGE Analysis**

We performed DGGE by using an 8% (w/v) polyacrylamide gel with 35–55% and 45–60% denaturant gradients for *Fusarium*-specific and *Pseudomonas*-specific communities, respectively, using the Junyi JY-TD331A system (JUNYI, Beijing, China). DGGE was carried out at 80 V and 60°C for 12 h and 15 h in 1x TAE buffer. After electrophoresis, gels were stained with silver stain. For analysis of the molecular community profiles, gels were digitized by using the Quantity One 4.0 software (BioRad). When bands were identified, they were excised from the DGGE gel by using a sterile scalpel. After incubation overnight at 4°C, DNA was eluted from the gel. The amplicons were amplified by using the Alfie1-GC/Alfie2 and F968-GC/PsR primer sets (as mentioned before). PCR amplicons were cloned into the pEASY-T1 Cloning vector (Transgen Biotech, Beijing, China) by using manufacturer’s instructions. Sequences were compared to the sequences on GenBank of NCBI using the BlastN search method.

**Quantitative PCR for Fusarium oxysporum and Pseudomonas spp.**

The fragments of *F. oxysporum* and *Pseudomonas* were cloned into the pEASY-T1 Cloning vector (TransGen Biotech Co., Beijing, China). Two plasmids were purified as described above. After determining DNA concentration, it was immediately diluted into 2, 1, 0.5, 0.1, 0.05, 0.01, 0.005, and 0.001 ng ml⁻¹. The reaction of standard curve was performed following the qPCR amplification protocol as described in Supplementary Table S1. In addition, the standard curve was generated by log10 value against the threshold cycle (Ct) value.

We further performed real-time PCR quantifications of *F. oxysporum* (primer sets ITS1F and AFP308R) and *Pseudomonas* (primer sets PsF and PsR) in four soil samples, and amplification protocol as described in Supplementary Table S1. Reaction of qPCR was performed in 15 µl mixture, containing 7.5 µl TransStart Green qPCR SuperMix (Transgen Biotech, Beijing, China), 0.6 µl of each primer (10 µ M) and 20 ng DNA.

**Isolation of Fusarium spp.**

For isolation of *Fusarium* spp., potato dextrose agar (PDA) was used to isolate and subculture the fungus. Soil suspensions were prepared by adding 10 g of fresh soil in a flask containing 90 ml of sterile water (10⁻¹ g l⁻¹), 100 µl soil suspensions were plated onto PDA. Plates were incubated at 30°C for 18 h, and then each single colony was isolated and purified. *Fusarium* genomic DNA extraction was done by using CTAB-based method as described by Rogers and Bendich (1985). The primer sets ITS1F and ITS4 (Supplementary Table S2) were used for ITS amplification. PCR amplicons were sent to Shanghai BoShang for sequencing. We
further used BlastN search method to compare sequences to the GenBank database.

Isolation and Counting of *Pseudomonas* spp. with Antagonistic Activity toward *Fusarium oxysporum*

For isolation of *Pseudomonas* spp., *Pseudomonas* selective isolation agar (PSIA) (Krueger and Sheikh, 1987) was used. As described above, each soil suspension was prepared (10−3 g l−1), after serial dilution, 60 µl soil suspensions (10−3 g l−1) were plated onto PSIA, incubated at 30°C for 30 h, and then each single colony was purified. Results were described as the numbers of CFU per g−1 (dry weight) soil.

For *in vitro* antagonism assays, we inoculated *F. oxysporum* to the center of the PDA plates and *Pseudomonas* isolates to the side of the plates at the same time. The results of antagonistic activity against *F. oxysporum* were recorded after 5 days of incubation at 30°C.

After incubation, we selected *Pseudomonas* isolates that had antagonistic activity against *F. oxysporum* for DNA extraction. *Pseudomonas* genomic DNA was extracted using the Bacteria Genomic DNA kit (CWbiotech, Beijing, China). The primer sets 27F and 1522R were used for 16S rRNA amplification. The thermal conditions are listed in the Supplementary Table S2. PCR amplicons were sent to Shanghai BoShang for sequencing. Finally, we used BlastN search method to compare sequences to the GenBank database for the identification purpose. Sequences were used Clustal X to align, and then phylogenetic trees were constructed with MEGA6.06 using a neighbor joining approach.

**Evaluation of the Pathogenicity of *Fusarium oxysporum* and Biocontrol Effects of *Pseudomonas* sp. CJ313**

*R. pseudostellariae* were planted in plastic pots and placed in a green house on December 15, 2015. The spore suspension of isolated *F. oxysporum* was added to the soil through pipette for observing the effects of *Fusarium* wilt in *R. pseudostellariae* after 5 months of planting. In order to assess biocontrol potential of *Pseudomonas* spp., the effect of isolated strain CJ313 was examined after 15 days of its exogenous addition. We added equal amount of LB as a control (CK) at the same time. Each treatment has three replicates. After 16 days, we collected rhizospheric soil from two treatments, then soil samples were immediately used to extract DNA and qPCR of *F. oxysporum* and *Pseudomonas* spp. as described above.

**The Effect of Phenolic Acids on the Growth of Isolated *Fusarium oxysporum*, *Pseudomonas* sp. CJ313 and *Pseudomonas* sp. CJ361**

Based on our previous HPLC results of phenolic acids in the *R. pseudostellariae* rhizosphere (Wu et al., 2016a), we prepared the solutions of eight phenolic acids (p-hydroxybenzoic acid, gallic acid, coumaric acid, syringic acid, vanillin acid, ferulic acid, vanillic acid and benzoic acid) and their mixture to assess its effect on the growth of isolated *F. oxysporum*. The ratio of their mixtures was the same as detected in the soil. We prepared the 10-fold dilution of soil extract agar medium (SEM), and added the phenolic acids into the SEM to reach final concentrations 30, 60, 120, 240, 480, 960 µ mol L−1. We inoculated isolated *F. oxysporum* onto the SEM plates to assess the mycelium growth mediated by phenolic acids. There were three replicates for each treatment. After incubation at 28°C for 8 days, we recorded the mycelium diameter. Likewise, isolated *F. oxysporum* was inoculated into 10-fold dilution of SEM by adding the phenolic acid mixtures, and solution was incubated at 200 rpm and 30°C for 7 days. *F. oxysporum* spores were counted by a hemocytometer.

We also detected the effects of eight phenolic acids and their mixtures on the growth of isolated *Pseudomonas* sp. CJ313 and CJ361. Specifically, the isolated *Pseudomonas* sp. CJ313 and CJ361 were determined by adding the phenolic acids to a LB medium with 8-fold dilution. After 8–10 h incubation at 200 rpm and 30°C, we determined the bacterial density at 600 nm using a microplate reader (Thermo Scientific Multiskan MK3, Shanghai, China).

**Statistical Analyses**

For all parameters, multiple comparison was carried out by one-way analysis of variance (ANOVA) followed by LSD’s test.

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**FIGURE 1 | (A) Photographs of above and below ground components of *R. pseudostellariae* under 1-year, 2-year, and 3-year consecutive monoculture. (B) Yield of *R. pseudostellariae* under 1-year (FP), 2-year (SP) and 3-year consecutive monoculture (TP).**
The Diversity of Fusarium-specific DGGE was also determined. The study revealed that Simpson, Shannon, evenness and Brillouin's index of Fusarium communities significantly increased with prolonged or increasing years of monoculture ($P \leq 0.05$) (Table 1).

Analysis of the DGGE Bands of Fusarium spp.
In order to further extract more detailed information from the DGGE bands in this study, excised bands from DGGE were sequenced. A total of 17 bands were identified in rhizospheric soil (Table 2). The Fusarium spp. belonged to 5 species, e.g., *F. oxysporum* (band a, b, c, g, h, k, l, m and n), *F. solani* (band p, q, r, s and t), *F. asiaticum* (band d), *F. falciforme* (band e), *F. foetens* (band f). Specifically, the bands of *F. oxysporum* significantly increased along with years of continuous cropping.

Pseudomonas-specific DGGE
Pseudomonas-specific PCR-DGGE analyses showed significantly changed Pseudomonas community structures in the rhizosphere with increasing years of monoculture (Figure 4). Likewise, we performed PCA to demonstrate the relative position of four soil samples. In PCA, the first principal component explained 83.30% of variance and second principal component 9.0% of total variance (Figure 3B). Furthermore, PCA showed the Pseudomonas community in CK, FP and SP were separated from the microbial community in TP by principal component 1, and the community in FP and CK was separated from the microbial communities in SP and TP by principal component 2 (Figure 3B).

The diversity of visible bands, Shannon and Brillouin’ index of Pseudomonas community significantly decreased with increasing years of monoculture ($P \leq 0.05$). However, the opposite was true for the Simpson’ index of the Pseudomonas community. There was no significant difference in evenness index among the four samples (Table 3).

Analysis of the DGGE Bands of Pseudomonas spp.
To further extract more detailed information from the DGGE bands, we excised and sequenced bands from DGGE. A total of 15 bands were identified in rhizospheric soil (Table 4). Pseudomonas spp. could be further divided into five species, e.g., *Pseudomonas lutea* (band c and k), *Pseudomonas fluorescens* (band d), *Pseudomonas aeruginosa* (band l), *Pseudomonas knackmussii* (band o and r), *Pseudomonas sp.* (band j) and uncultured bacterium (a, b, f, g, i, m, p and q).

Abundance of Pseudomonas and Fusarium oxysporum by Quantitative PCR
First, standard curves of $y = -0.2487x + 9.898$ ($R^2 = 0.997$) and $y = -0.271x + 9.8309$ ($R^2 = 0.990$) were developed for *Pseudomonas* and *F. oxysporum* qPCR analyses respectively. The first principal component, and FP was separated from SP by the second principal component (Figure 3A).
amount of *F. oxysporum* was significantly (*P* ≤ 0.05) higher in continuous monoculture soils (SP and TP) than in control (CK) and the newly planted soils (FP) (Figure 5A). The result of qPCR was consistent with the *Fusarium*-specific DGGE results (Table 1). However, the opposite was true for the qPCR result of *Pseudomonas* (Figure 5B).

### Isolation and Screening for *F. oxysporum* with High Pathogenicity

In our study, we separated and sequenced one strain of *F. oxysporum*. We found that the isolated *F. oxysporum* quickly led to wilt disease on the tissue culture of *R. pseudostellariae* (Figure 7A), and it also occurred in pots with *F. oxysporum* (Figure 7B). These results demonstrated that isolated *F. oxysporum* had the high pathogenicity on *R. pseudostellariae*.

### Screening for *Pseudomonas* Isolates with Antagonistic Activity toward *F. oxysporum*

For *in vitro* antagonism assays, we screened a total of 317 *Pseudomonas* isolates from four different soils. The results showed that the isolation frequencies of *Pseudomonas* were significantly higher in FP than SP and TP. The highest isolation frequencies were found in the newly planted (FP) soil (Figure 6A). *In vitro* antagonism assays, the number of *Pseudomonas* spp. with antagonistic activity toward *F. oxysporum* significantly declined with prolonged monoculture (Figure 6B). These isolation frequencies were similar to results of *Pseudomonas* obtained by qPCR. Approximately 17.4% (87 of 317) of all isolates showed the antagonistic activity. Strain
313 and 361 but not 117 had antagonistic activity against \textit{F. oxysporum} (Figure 6C). The sequences of \textit{Pseudomonas} sp. CJ313 and \textit{Pseudomonas} sp. CJ361 isolates were obtained to perform phylogenetic tree analysis. The neighbor-joining method generated a dendrogram with two main branches, where the first branch included \textit{Pseudomonas} sp. CJ361 and the second branch comprised \textit{Pseudomonas} sp. CJ313 (Supplementary Figure S1).

### Biocontrol Effects of \textit{Pseudomonas} sp. CJ313

We further evaluated the antagonism of \textit{Pseudomonas} CJ313 to \textit{F. oxysporum}. In the pot experiment, we found that the isolated \textit{Pseudomonas} CJ313 significantly inhibited the growth of \textit{F. oxysporum}, and the \textit{R. pseudostellariae} grew well without disease symptoms during the period of experiment (Figure 7B). Moreover, qPCR indicated that the abundance of \textit{Pseudomonas} was significantly higher in \textit{Pseudomonas}. CJ313 treatment than in control (CK), whereas \textit{F. oxysporum} showed the opposite trend (Figure 7C). The results clearly showed that strain \textit{Pseudomonas}. CJ313 has the potential of biological control. The results further suggested that exogenous antagonism of \textit{Pseudomonas} could be effective against \textit{F. oxysporum} infection. In addition, the results also demonstrated that the imbalance of these two strains (\textit{Pseudomonas} sp. CJ313 and \textit{F. oxysporum}) could be an important cause of the continuous cropping related diseases.

### The Effect of Phenolic Acids on the Growth of Isolated \textit{Fusarium oxysporum}, \textit{Pseudomonas} spp.

The results showed that mycelial and sporular growth of \textit{F. oxysporum} was significantly promoted by phenolic acid mixture (Figures 8A,B). Further analysis showed that p-hydroxybenzoic acid, vanillin, coumaric acid and ferulic acid could significantly promoted mycelial growth of \textit{F. oxysporum} among the eight phenolic compounds (Supplementary Figure S2). The results also indicated that...
the growth promotion by mixture was more than that of single phenolic acid on *F. oxysporum* (Figure 8A). However, the mixture significantly inhibited *Pseudomonas* sp. CJ313 (Figure 8C) and *Pseudomonas* sp. CJ361 (Figure 8D) growth. Among them, vanillic acid and syringic acid has the more inhibitory effects on *Pseudomonas* sp. CJ313 than others (Supplementary Figure S3). Likewise, coumaric acid, ferulic acid syringic acid had the greatest inhibitory effect on *Pseudomonas* sp. CJ361 (Supplementary Figure S4). The results indicated that certain allelechemicals of *R. pseudostellariae* root exudates possessed the selective effects on rhizosphere microbes.

**DISCUSSION**

Our studies presented a significant decline in the yield of *R. pseudostellariae* along with less aboveground biomass in consecutive monoculture field (Figure 1). Recently, researchers have focused on the biological relationships between plants and rhizosphere microorganisms, which are essential for plant growth and health (Haney and Ausubel, 2015; Lebeis et al., 2015). The study of *F. oxysporum* has become common due to its ability to cause diseases of important economic crops (Gordon et al., 1989; Gordon and Martyn, 1997). DGGE results revealed significant changes in *Pseudomonas* and *Fusarium* communities in the rhizosphere of *R. pseudostellariae* with prolonged monoculture (Figures 2, 4). Based on the DGGE analysis of *Fusarium*, we indicated that prolonged monoculture of *R. pseudostellariae* led to a significant increase in *Fusarium* species, especially *F. oxysporum* (Table 1). Quantitative PCR assay confirmed the increase in *F. oxysporum* with the increasing years of monoculture (Figure 3). These results are supported by the work of different researchers as stated that *F. oxysporum* is one of main pathogenic species to plants under monoculture regime (Wu et al., 2016b,c).

Due to an extensive distribution of *Pseudomonas* species in the environment, several studies reported an abundance of antagonistic *Pseudomonas* species, which controls specialized pathogens that are responsible for disease suppression in soils (Gorlach-Lira and Stefaniak, 2009; Mendes et al., 2011). Our study of *Pseudomonas-DGGE* revealed that the diversity of *Pseudomonas* spp. significantly declined with the prolonged monoculture. More importantly, it was found that the relative abundances of antagonistic *Pseudomonas* spp. declined in soils under consecutive monoculture, and a similar tendency was recorded for other *Pseudomonas* species studied in the selective medium assay. Similar effects of plants on the abundance of antagonistic *Pseudomonas* spp. under monoculture were found by Gorlach-Lira and Stefaniak (2009). Hence, the abundance of the *Pseudomonas* populations in soil of *R. pseudostellariae* were seriously affected by monoculture. In addition, it was also found that the abundance of *Pseudomonas* spp. having antagonistic activities against *F. oxysporum* significantly decreased with the increasing years of monoculture, and this was confirmed by the *in vitro* antagonism assays. This important antagonistic interaction effects between *Pseudomonas* and *F. oxysporum* need particular attention in disease management under a clear cropping system of *R. pseudostellariae*. Therefore, it is necessary to make robust inferences about balance between *Pseudomonas* communities and *Fusarium* of *R. pseudostellariae*.

Our previous study revealed that most phenolic acids of *R. pseudostellariae* from rhizosphere soil indicated no direct autotoxicity toward tissue culture seedlings of *R. pseudostellariae* (Wu et al., 2016a). Besides, many researchers did not support the assumption that the concentrations of allelochemicals in the soil were sufficient to directly influence the development of host plants or neighboring plants (Ehlers, 2011; Weidenhamer et al., 2013). A growing number of researchers reported that the microflora disorder mediated by plant root exudates was the crucial factor leading to plant consecutive monoculture problems (Wu et al., 2014). Root exudates have selective effects on certain microorganisms in the soil and can promote or inhibit the growth of a certain population (Haichar et al., 2008; Hartmann et al., 2009). In this study, the results indicated that the phenolic acid mixture had a significant improvement on the growth of mycelial and spore of pathogenic *F. oxysporum* (Figures 8A,B). However, phenolic acid mixture could greatly inhibit the growth of antagonistic *Pseudomonas* sp. CJ313 and CJ361.
FIGURE 6 | Pseudomonas populations (CFU g\(^{-1}\) of dry soil) in rhizosphere soils under four different samples (A). Number of Pseudomonas spp. with antagonistic activity against \(F.\) oxysporum in four different samples (B). Petri plates used for evaluation of antagonistic activity of Pseudomonas strains 313, 361 and 117 against \(F.\) oxysporum (C). CK, FP, SP and TP represent the control, newly planted, 2-year, and 3-year consecutively monoculture soils, respectively. Data are means ± standard errors (one-way analysis of variance, \(n = 3\)).

FIGURE 7 | Assessment of the pathogenic potential of isolated \(F.\) oxysporum (A), the biocontrol potential of Pseudomonas sp. CJ313 against \(F.\) oxysporum (FOX) (B). Quantification of \(F.\) oxysporum and Pseudomonas spp. from two samples. Data are means ± standard errors (one-way analysis of variance, \(n = 4\)).

(Figures 8C,D). Zhou and Wu (2012) observed that the abundance of \(F.\) oxysporum in soil was significantly increased by \(p\)-coumaric acid, which led to the severity of Fusarium wilt in field conditions. Wu et al. (2016a) reported that phenolic acid, such as syringic acid, significantly promoted the growth of Talaromyces helicus and Kosakonia sacchari, and inhibited growth of Bacillus pumilus. Bais et al. (2002) found that rosmarinic acid had a significant and deleterious effect on Pseudomonas aeruginosa. Furthermore, plant bioassays with representative isolates of Pseudomonas showed that Pseudomonas
CJ313 had a good performance that protected *R. pseudostellariae* from infection by *F. oxysporum* (Figure 7B). Combined with above-mentioned results, we can draw robust inferences that the imbalance of belowground microbial community resulted in the poor growth of monocultured *R. pseudostellariae* by root exudates.

**CONCLUSION**

Based on multifaceted approaches, such as cultural-independent and culture-dependent analyses, this study indicated that *R. pseudostellariae* biomass decreased under 3-year extended monoculture resulted from two important factors: (i) the decrease of antagonistic microorganisms (*Pseudomonas* sp. CJ313 and CJ361) against pathogens (*F. oxysporum*) might be due to selective inhibitory effect of root exudates, especially phenolic compounds and (ii) an increase of *F. oxysporum* which significantly induced the poor growth of *R. pseudostellariae* at a time when pathogenic microbes (*F. oxysporum*) have become dominant. (iii) Isolated *Pseudomonas* CJ313 of its exogenous addition could protect *R. pseudostellariae* from infection by *F. oxysporum*. These results are very important in the development of potential management approaches to solve the *R. pseudostellariae* problems under consecutive monoculture. However, additional works are still needed to explore the relationship between aboveground plant performance and belowground microbial diversity.

**AUTHOR CONTRIBUTIONS**

WL and JC conceived the study; JC wrote the paper; JC, and LW performed experiments; JC, SL, and ZX performed the statistical analyses; HW, XQ, YW, XW, and JW were involved in field management. MK assisted in English correction. All authors discussed the results and commented on the manuscript.

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**Conflict of Interest Statement**: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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