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# The effects of exotic weed *Flaveria bidentis* with different invasion stages on soil bacterial community structures

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A new exotic weed, *Flaveria bidentis*, is spreading in central China where it forms dense monospecific patches modifying the structure of different native ecosystems and threatening native aboveground biodiversity. However, little is known about the consequences of such an invasion for soil bacterial community, especially its effect pattern at different invasion stages. In this study, soil samples were taken in native ecosystems that were uninvaded, partially invaded (transition), and severely invaded by *F. bidentis*. The bacterial richness and diversity in *F. bidentis* in rhizospheres soil was evaluated using denaturing gradient gel electrophoresis (DGGE) analysis. Different stages of *F. bidentis* invasion can trigger changes in soil physicochemical properties in particularly in available N and P *F. bidentis* invasion significantly decreased the richness of soil bacterial community, and the decline contents were positively correlated with invasion progress. In the invaded soils, bacterial species in *Proteobacteria*, *Chloroflexi* and *Actinomycetes* decreased with invasion, with the greatest reduction in relative abundance occurring for *Proteobacteria*, which was the dominant species in the native soils. Invasion of *F. bidentis* corresponded with an alteration in the structure of soil bacterial community, and soil microbial biomass as well, thus soil environment modification was expected to promote spreading of this exotic weeds in turn.

**Key words:** Biological invasion, *Flaveria bidentis*, soil nutrients, soil bacteria, polymerase chain reaction denaturing gradient gel electrophoresis (PCR-DGGE).

# INTRODUCTION

Biological invasion has become a global ecological and economic problem. Understanding the impact of invasive species on the local ecological systems has gained increased attention in invasive ecology (Ehrenfeld et al., 2001). Invasive species employs very complex and multiple mechanisms and strategies (Mack et al., 2000). Once establishing in a new environment, exotic weeds can replace the native species, causing above vegetable

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Abbreviations: MBC, Microbial biomass carbon; MBN, microbial biomass nitrogen.

Author(s) agree that this article remains permanently open access under the terms of the <u>Creative Commons Attribution License 4.0</u> International License structural changes in the native ecological system. Soil microorganisms play an important role in the successful spread of exotic weeds (Hierro et al., 2005). Altered soil microbial communities and resulting effects on ecosystem processes may be an invisible legacy of exotic weed invasions by rhizosphere microbe and the host plants an inter-dependent and mutual constraint have relationship, which is the reason that why certain rhizosphere microbial communities only co-exist with specific plant species (Schloter, 2003). It has also been suggested that exotic weeds could disrupt mutualistic associations within native microbial communities (Richardson et al., 2000; Callaway and Ridenour, 2004; Stinson et al., 2006). Alteration in the structure and function of soil microbial communities could eventually lead to changes in the vegetation structures (Callaway et al., 2013). Among soil microbial communities, bacteria associated with plant roots are fundamentally important in plant nutrition. arowth promotion. and disease interactions (Marschner et al., 2001). For this reason, there has been considerable interest in characterizing the structure and function of rhizosphere communities. The bacterial community composition in the rhizosphere is important for the performance of the plant, as bacterial species can have beneficial, neutral or harmful relationships with the roots (Atkinson and Watson, 2000; Sylvia and Chellemi, 2001). It is well accepted that different plant species can be selected for specific rhizosphere communities (Burke et al., 2002; Costa et al., 2006). The differences in root-derived substrates are claimed to explain the plant specific rhizosphere bacterial communities that have been observed for different plant species grown under similar conditions (Marschner et al., 2001; 2002; Smalla et al., 2001).

A new exotic weed, Flaveria bidentis (L.) Kuntze, commonly called "vellowtop", is increasing spreading in central China. This species originated from South Africa and was first found in 2001 in suburbs of Tianjin and a few cities of Hebei province (Liu, 2005). It invades roadsides, abandoned field or even arable fields, outcompetes natural vegetation, and forms a dense population (Huangfu et al., 2011). This weed tolerates environmental stresses of salinity and cold temperature, and could become troublesome for the development of sustainable agriculture (Gao, 2004). The function and population of rhizosphere microbes also undergo various alterations to allow the establishment of the invasive species. However, there is little information on the impact on soil bacterial diversity and mechanism upon invasion by the exotic weeds (Lorenzo et al., 2010).

This study aimed (1) to examine the effects of different stages of *F. bidentis* invasion on soil physicochemical properties and (2) detect the effects of different degrees of *F. bidentis* invasion on the community structure of bacteria in soils. We hypothesize that (1) increasing stages of *F. bidentis* invasion enhance soil nutrient element concentrations (especially soil N) because

invasive plants have high nutrient cycling rates, especially for N (van Kleunen et al., 2010; Laungani and Knops, 2009; Jones and Chapman, 2011), and that (2) *F. bidentis* invasion significantly increases the richness and diversity of the soil bacterial community along the invasion gradients. Also, the changes in soil bacterial communities were associated with soil physicochemical properties. Towards these aims, we used the PCR-DGGE approach together with cloning and sequence analysis of 16S rRNA fragments of soil bacteria upon the invasion process by *F. bidentis*. In our findings will provide fundamental knowledge for soil bacteria diversity upon invasion by alien plant species.

#### MATERIALS AND METHODS

#### Site description and sampling

The sampling sites were collected in wasteland ecosystems, a typical system F. bidentis infestation (Zhang et al., 2010), located in the Xian County in north China (38° 15' 30"N, 115° 57' 50"E) with temperate continental monsoon climate, mean annual precipitation of 560 mm, mean annual temperature of 12.3°C, and its average frostless period lasts 189 days. Geographically, the experimental site had flat land, a uniform landscape, and a similar terrain and soil origin with very minor disturbance by human and animals, and very minimal habitat variation. The soil is alluvial type where F. bidentis plants grown as monocultures had formed alternate successions. The following three sites (soil types) with three different levels of invasion by F. bidentis were sampled: (a) native soils (the control) mainly dominated by native herbaceous plants, including Setaria viridis (L.) Beauv, Digitaria ciliaris, Phragmites australis and Echinochloa crusgalli with coverage of over 60%; (b) transition soils where F. bidentis plants covered 10 to 30% of the plot, and (c) invaded soils where F. bidentis covered over 60% of the plot. Soil samples were collected on August 10 in 2009. In each of the sampling sites, six plots (repetitions) were randomly chosen each covering 3 × 3 m area with about 10 to 20 m apart from each other. The five points Quincunx sampling scheme was used to collect soil samples in each plots at the 0 to 20 cm depths, and soils within the same plot were pooled and mixed together equally as one replication, thus 18 soil samples collected in total were placed in plastic bags for transport to laboratory. Prior to sampling, all plants and organic matter debris on the ground were removed. Samples were stored at -20°C until analysis. From 1000 g of each soil sample, 20 g were homogenized and subsamples of 5 g were taken for further analysis. To verify the impact pattern found with this invasive plant, sampling was done in following year. Twice sampling data was pooled for soil physicochemical parameters analyses and only once PCR-DGGE fingerprinting was presented given the fact that there were no inter-year differences between treatments detected.

#### Determination of soil nutrients

Soil NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> were extracted by shaking 20 g of fresh soil in 100 ml of 2 M KCl solution for 1 h. Soil extracts were analyzed with the FIAstar 5000 Auto Analyzer system. Total N and P in soil samples were analyzed with oven-dried samples, 48 h at 70°C. The Kjeldahl method was used for analyzing the total nitrogen (N) content of the soil. Soil mineral N was extracted using 2 mol L<sup>-1</sup> KCl, then the concentrations of NO<sub>3</sub><sup>-</sup>-N in the KCl extracts were determined by hydrazine sulfate colorimetry and the concentrations

of  $NH_4^+-N$  by indophenol blue colorimetry (Mulvaney, 1996). Total phosphorus (P) was extracted using the  $HCIO_4-H_2SO_4$  method, and available P was determined using the sodium bicarbonate method.

#### **DNA extraction from soil samples**

Total DNA was isolated from soil samples using the PowerSoil<sup>®</sup> DNA Isolation Kit (MO BIO Laboratories, Inc., CA), following the instruction of the maximum yield method. After a final purification, the soil DNA was visualized on 1% (w/v) agarose gels to assess its purity and molecular size. The final DNA extracts obtained from the soils were color-free, indicating that they did not contain high amounts of humic compounds.

#### Specific PCR of 16s rRNA gene fragments

For amplification of 16s rRNA fragments, a pair of universal primers consisting of the 357f- GC and 518r (Muyzer et al., 1993) were used to amplify the V3 region of bacterial 16S rRNA. Primer sequences were 357f-GC (5'-GCclamp-CCTACGGGAGGCAGCAG-3'), and 518r (5'-ATTACCGCG GCTGCTGG-3'). The PCR reaction were carried out in a final volume of 50 µl containing 2 µM of MgCl<sub>2</sub>, 200 µM of each dNTP, 0.5 µM of each primer, 50 ng of isolated DNA, 5 µl 10 × PCR buffer, and 2.5U of Ex Taq<sup>TM</sup> polymerase (TaKaRa Inc., Dalian, China). Touch-down PCR procedure was performed for increasing both the specificity and sensitivity of PCR assays in a thermal cycler (Bio-Rad) (Labbe et al., 2007). After preincubation at 95°C for 5 min, samples were amplified with denaturation for 1 min at 94°C, annealing for 1 min (temperature decreasing 0.5°C per cycle from 65 to 55°C, and then 15 cycles at 55°C), primer extension for 3 min at 72°C, followed by one final extension at 72°C for 15 min. Aliquots (5 µL) of PCR mixture were examined by electrophoresis in an agarose gel (1%, w/v) stained with ethidium bromide to check fragment size and integrity.

#### **DGGE Patterns**

DGGE was performed with 8%(w/v) acrylamide gels containing a linear chemical gradient ranging from 40 to 60%. The gels were allowed to polymerize overnight. DNA samples containing 20 µl of the PCR products were electrophoresed in 1× TAE buffer at 60°C at a constant voltage of 120 V for 8 h, and all DGGE analysis was done in the Dcode Universal Mutation Detection System (Bio-Rad, Hercules, CA, USA). After electrophoresis, the gels were stained for 30 min with SYBR gold nucleic acid gel stain (Invitrogen Molecular Probes, Eugene, USA) (10,000-fold diluted in 1× TAE) and photographed under UV light with a video imaging system. Band detection and quantification of band intensity was performed using Quantity One 4.62 software (Bio-Rad, USA). DNA band intensity was normalized by dividing the band intensity of each band by the mean band intensity of the gel. Therefore, both band position and intensity are expressed as relative values. Each peak represents individual groups of species having 16S rRNA sequences with similar melting behavior. The band intensity indicates the relative abundance of the group under these PCR conditions.

#### Sequence analysis of DGGE bands and phylogenetic analysis

Selected DGGE bands that occurred in majority of samples were excised from the gels and eluted (Kowalchuk et al., 1997). The criteria for selection of bands were that (i) they appeared as a single band in the pool of lanes, (ii) they represented bands in high abundance in the community or (iii) they were of relatively low abundance in the DGGE pattern. It is noteworthy that, in some

cases where multiple clones were generated from a given excised band, more than one phylotype was detected from that band. In total, we sequenced 20 different clones, corresponding to 17 excised DNA fragments. These sequences of 16S rRNA genes obtained were submitted to the GenBank to determine the closest known relatives of the partial 16S rRNA sequences and the phylogenetic affiliations are shown in Table 2. Eluted DNA was then amplified using the 518r and 357f primer pair without GC clamp, and PCR products were ligated onto pMD19-T vector (Takara) and transferred into Escherichia coli JM<sub>109</sub> competent cells. After positive cloning selection, the white colonies were further screened with vector primer pMD19-T to confirm the positive clones. The positive colonies were cultured in LB broth overnight at 37°C with constant shaking. Aliquots of 500 µL bacterial stocks were mixed with sterile glycerine (50%) and stored at -70°C. The clones of each of excised bands were chosen for sequencing. Sequencing was carried out at Shanghai Biotech Company. Searches in GenBank with the BLAST program (Altschul et al., 1997) were performed to determine the closest known relatives of the partial 16S rRNA sequences obtained. Multiple alignments of the sequences were performed using Clustal X (Thompson et al., 1997). A phylogenetic tree was constructed by the neighbor-joining method in MEGA 4.1 (Tamura et al., 2007). The confidence values for the branches of the phylogenetic tree were determined using bootstrap analysis (Felsenstein, 1985) based on 1000 resamplings. The similarity between sequences was calculated using the GENETYX computer program (Yumoto et al., 1999).

#### Measurement of soil microbial biomass carbon and nitrogen

Microbial biomass carbon (MBC) and nitrogen (MBN) were measured by a chloroform-fumigation extraction method modified from Vance et al. (1987). Six aliquots of wet soil from each sampling replications equal to 20 g dry weight were placed into 100 ml beakers. Three samples were fumigated whereas the other three were not. Soil was placed on top of the internal shelf in a vacuum desiccators drier that had internal diameter of 29 cm. Below the shelf, 60 ml HPLC grade chloroform and glass beads (to prevent explosion) were put into a 100 ml beaker. After addition of 50 ml 1 mol/L NaOH, the soil was covered with a few layers of wet filter paper to maintain moisture content of the steamed soil. After sealing the drier with Vaseline, vacuum was turned on and chloroform started to boil. Degas was topped after 5 min, and the samples were stored in darkness at 25°C for 24 h. After removal of the beaker containing chloroform, the soil was degassed again to remove chloroform residuals. The non-steaming treated soil was placed into a separate drier, and the chloroform was replaced with distilled water. After putting the fumigated soil into a 150 ml flask and addition of 60 ml 0.5 mol/l  $K_2SO_4$  (soil: water = 1:4), the mixture was shaken at 25°C and 200 rpm for 30 min. The extracts were filtered through mid-speed filter paper, and the filtrates were measured immediately or stored at -15°C for later analysis. Soil microbial biomass carbon and nitrogen contents were measured using a multi NC3100 TOC/TN instrument (Analytik Jena AG. Germany). MBC and MBN of soil microbes were calculated using the differences in organic carbon and nitrogen between fumigated and non-fumigated soil, divided by the conversion factor of 0.45 (Joergensen, 1996). Data from the same soil type were pooled for analysis.

#### Statistical analysis

One-way ANOVA and Duncan's Test as post hoc test were used to check the differences in soil nutrients and microbial biomass C and N, bacterial richness and diversity between different soil types. Richness, defined as number of species, was calculated as the total

Type of soils	NS <sup>a</sup>	TS	IS
Organic matter (g⋅kg <sup>-1</sup> )	76.02±0.83 <sup>b</sup>	77.56±0.06 <sup>b</sup>	80.36±0.51 <sup>a</sup>
Total N (g⋅kg⁻¹)	0.81±0.02 <sup>b</sup>	0.86±0.01 <sup>b</sup>	1.00±0.01 <sup>a</sup>
Total P (g⋅kg⁻¹)	0.75±0.05 <sup>a</sup>	0.73±0.41 <sup>ª</sup>	0.64±0.13 <sup>a</sup>
NH₄-N (mg⋅kg⁻¹)	7.40±0.49 <sup>c</sup>	12.92±1.53 <sup>b</sup>	19.33±0.81 <sup>ª</sup>
NO₃-N (mg⋅kg⁻¹)	2.30±0.03 <sup>c</sup>	4.72±0.06 <sup>b</sup>	6.28±0.06 <sup>a</sup>
Available P (mg⋅kg⁻¹)	9.20±0.25 <sup>a</sup>	6.50±0.10 <sup>b</sup>	4.63±0.03 <sup>c</sup>

 Table 1. Changes in soil nutrient contents of different type of soil in *F. bidentis* invaded area.

<sup>a</sup>IS, invaded soils; TS, transition soils; NS, native soils; <sup>b</sup>different lowercase letter within same row means difference at P=0.05 level.

Table 2. Phylogenetic affiliation of sequences retrieved from DGGE bands.

Band	Closest relative	Similarity %	Accession number
1	Kaistobacter sp.	99	FJ889336.1
2	Bacteroidetes bacterium clone	100	GU552199.1
3-7	Actinobacterium	98	EU328001.1
3-9	<i>Legionella</i> sp.	98	GU598197.1
4-1	Uncultured bacterium clone	99	EU133920.1
4-2	Nitrosospira sp. clone	100	GU472862.1
5	Acinetobacter sp.	97	GU827519.1
6	Mycobacterium houstonense	94	EU977810.1
7	Chloroflexus sp.	100	AB257633.1
8	gamma proteobacterium	94	FJ568411.1
9	<i>Devosia</i> sp.	100	FN594685.1
10	Streptomyces rochei strain	100	HM153701.1
11	Rhodocyclaceae bacterium	99	EF606817.1
12	<i>Bacillus</i> sp.	98	EU043767.1
13	Chloroflexaceae bacterium enrichment	94	EU918581.1
14	Acidobacteriaceae	99	AM936873.1
15-17	Chelatococcus sp	98	AM412118.1
15-19	Uncultured bacterium clone	97	EU546566.1
16	Nocardioides sp.	100	GU202324.1
17	Uncultured alpha proteobacterium clone	98	GU552196.1

number of bands per sample, and diversity, defined as number of different species and their relative frequency (Lorenzo et al., 2013). Based on DGGE results, cluster analysis of soil bacteria was performed using the Ward's method by the Quantity-One software (Bio-Rad). The Shannon–Wiener index of general diversity, *H* was calculated using the following function:

$$H=-\Sigma P_{i}\log_{2}P_{i} \tag{1}$$

Where, Pi is the importance probability of the bands in a track. H was calculated on the basis of the bands on the gel tracks using the intensity of the bands as judged by peak heights in the densitometric curves (Ampe and Miambi, 2000). The importance probability Pi was calculated as:

 $P_i = n_i / \Sigma n_i$  (2)

ANOVA was performed with the SPSS 17.0 for Windows software package, and mean comparison was done using the least significant difference (LSD) test at P < 0.05.

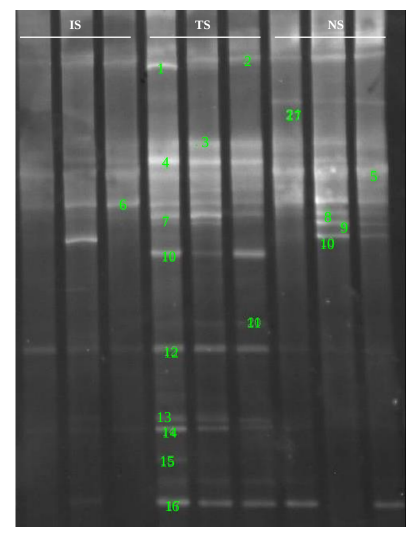
#### RESULTS

#### Changes in soil nutrient contents status

Compared with native soils, the *F. bidentis* invaded soils had obviously higher contents of soil organic carbon, total nitrogen,  $NO_3^-$ , and  $NH_4^+$ , but lower contents of soil available phosphorus (*P*<0.05). For example, it was increased by 5.7 and 23.4% in soil organic carbon and total nitrogen, respectively, while available phosphorous was reduced by 49.6% in invaded soil (*P* < 0.05, Table 1).

### Impact of invasion by F. bidentis on bacteria diversity

16S rRNA fragments amplified from DNA extracted directly from soil samples were compared and only three



**Figure 1.** DGGE analytical result of bacterial 16S rRNA V3 fragments in different type of soil in *Flaveria bidentis* invaded area. IS, invaded soils; TS, transition soils; NS, native soils.

replications of each site were presented due to a relatively high similarity of the DGGE patterns obtained for each of the replicates, which also suggested a low degree of variability caused by sampling (Figure 1). DGGE profiles of amplified 16S rRNA fragments from DNA extracted from the rhizosphere bacterial fractions significant differences of the bacterial revealed fingerprints from different F. bidentis invasion stages. Both the strains and number of bacterial reduced in wellinvaded soils (Figure 1). Bands that were shared among all the soil samples included bands 2, 4, 6, 12 and 16, indicating that bacteria carrying these genes were common to all types of soil and were not affected by the invasion. The H' were ranked in descending order as native soils (2.96) > transition soils (2.58) > invaded soils (2.33) (P < 0.05). Consequently, compared with the control (native soils), the invasion of F. bidentis reduced bacterial diversity with invasion progress.

Analysis of the DGGE profiles found were different in soil bacterial community between different soils. The transition soils and the native soils were clustered firstly at similarity index of 0.89 as one group, while the invaded soils distinctly separated from them with similarity of 0.68 (P < 0.05). The impact of this exotic weed on soil bacteria was a continuous process; as the invasion intensified some bacterial strains diminished in the soil. By comparing sequencing results, it was found that 20 sequences belonged to six different bacterial phyla with the majority in the division of Proteobacteria (Table 2). The similarity of the closest relatives of the partial 16S rRNA fragments of all sequenced bands ranged between 94 and 100%. Bands 2, 4-2, 7, 9, 10 and 16 showed the highest sequence similarity of 100%, respectively. On the other hand, other bands have various similarities with the

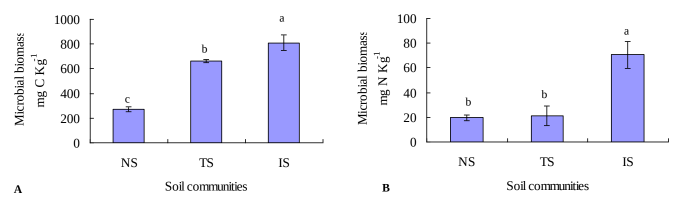


Figure 2. Mean microbial biomass (±SE) carbon (a), and nitrogen (b) for the different soil communities. NS, native soils; TS, transition soils; IS, invaded soils.

similar sequences in the NCBI database. The bands 1, 11 and 14 showed the higher sequence similarity (99%) with those of genus *Kaistobacter*, *Rhodocyclaceae* and *Acidobacteriaceae*, respectively, while bands 6, 8 and 13 only got 94% similarities with assigned sequences, respectively. According to the intensity of the band, a bacterial species that exhibited the higher sequence similarity to genus *Kaistobacter* (99% similarity) (band 1) was one of the most predominant during invasion of *F. bidentis*.

#### Impact of F. bidentis invasion on soil MBC and MBN

This result indicates that *F. bidentis* invasion led to alteration of microbial carbon metabolism in the soil (Figure 2). Upon the invasion of *F. bidentis*, soil MBC increased accordingly. The MBC content was ranked in descending order as following: invaded soils>transition soils> native soils. MBC was almostly 200% higher in the invaded soils than that of native soils (Figure 2a, P < 0.05). The same pattern was found for MBN which increased significantly after invasion by the exotic weed (Figure 2b). However, non-significant effect was detected between the transition soils and the native ones in MBN. Based on MBC and MBN, invasion of *F. bidentis* increased soil nutrient level as suggested in Table 1.

# DISCUSSION

The results obtained partially support our original hypothesis. Firstly, invasion of *F. bidentis* lead to significant increases in soil N, organic matter, but decrease in available P. Because the genus of *Flaveria* is extensively cloned by arbuscular mycorrhizal fungi, the fungi known to help plant uptaking phosphorus from soils that are P-deficient for plant growth (Aziz et al., 1995; Bagayoko et al., 2000), further research should focus on possible competitive relationship in uptake of phosphorus

between F. bidentis and native plant species. As a result, the decrease of phosphorus was attributed to the high uptake by this exotic weeds and the competition of exotic weeds with the soil community. Secondly, we found that invasion of F. bidentis was associated with significant increases in total soil N, C, organic matter and exchangeable P, but reduced soil bacterial diversity index, contrary to findings of Sanon et al. (2009). According to the positive feedback hypothesis, exotic weeds may cause soil-based ecosystem processes change following invasion, and such changes could establish positive feedbacks that enhance the spread of the exotic (Ehrenfeld et al., 2001). Our study suggests that number and diversity of soil bacteria changed, that the invasion of F. bidentis propagated certain groups of bacteria while suppressing others. Even some bacterial were common to all type soils, but plant species will eventually alter structure of soil bacterial community (Briones et al., 2002).

Many phytopathogenic organisms, bacteria as well as fungi, have coevolved with plants and show a high degree of host specificity (Raaijmakers et al., 2009). As the invasion progressed, the soil bacterial community structures also underwent significant changes as suggested by similarity analysis of the DGGE fingerprints where invaded soils clustered separately from native soils in the cluster analysis.

A decrease of bacterial diversity and increase of microbial biomass could be also caused by an increase of fungal biomass (Schimel et al., 1999). Alteration of native microbial community composition may further decrease competition from native plants and therefore support *F. bidentis* dominance as suggested by Rudgers and Orr (2009). *F. bidentis* can release allelopathic compounds (kaempferol, quercetin) (Xie et al., 2010; Iwashina, 2003), which may inhibit the growth of many microorganisms. It was found that extracts of *F. bidentis* from both leaves and roots reduced seed germination and seedling growth of native plant species (Huangfu et al., 2011). Therefore, allelopathic compounds produced

by *F. bidentis* may be responsible for alterations in microbial biomass pools but further study is needed.

Our previous study has shown that *F. bidentis* invasion significantly decreases soil pH values (Zhang et al., 2010). This result may be mainly attributed to the fact that this invasive plant has high ammonium uptake rates as suggested by our study (unpublished data). The metabolic activities and community structure of soil microorganisms were highly correlated with soil pH values (Hackl et al., 2005; Högberg et al., 2007). Thus, we believe that changes in soil pH values mediated by *F. bidentis* invasion can enhance the succession of soil microbial communities in the rhizosphere and facilitate further invasion. With the widespread introduction and invasion of exotic weeds there are many studies that investigate alteration of basic ecosystem structure and function.

However, studies concerning invasive processes, information about changes in the impact over time is rarely available (Souza-Alonso et al., 2015). Some studies found that changes in soil properties as C or N contents and microbial properties soil ecosystem parameters are more pronounced after a long period of invasion (Marchante et al., 2008). Nevertheless, recent findings suggest that both ecological and adaptation processes may increase or attenuate the impact of invaders on the resident community, and that the impact of an invasive species on soil characteristics and on the structure and function of microorganisms does not necessarily remain constant or accumulate over the course of invasion (Strayer, 2012; Dostál et al., 2013). Our study sought to determine the effects of different stages of plant invasion on soil bacterial communities to better understand the mechanism of plant invasion. Different stages of F. bidentis invasion can trigger changes in soil physicochemical properties, in particularly in available N and P. F. bidentis invasion significantly decreased the richness of soil bacterial community, and the decline contents were positively correlated with invasion progress. Changes in the soil physicochemical properties and community structure of soil bacterial communities mediated by F. bidentis invasion may play an important role in facilitating further invasion.

# **Conflict of interests**

The authors did not declare any conflict of interest.

# ACKNOWLGEDMENTS

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