Difference of Bacterial Community Structure in the Meadow, Maize, and Continuous Cropped Alfalfa in Northeast China

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Maize and alfalfa (*Medicago sativa* L.) have been used extensively in the animal husbandry to compensate for the lack of livestock and fodder yields in the chilly northeast of China. Little is known, however, about the impact on soil characteristics of consecutive plantings in various crops and alfalfa. In this research, the soil characteristics, bacterial community diversity, and structure of the meadow, maize, and alfalfa continuous cropping fields (i.e., 6, 10, 14, 20, and 30 years) were measured. The results showed that maize cropping and continuous cropping of alfalfa increased the soil bacterial alpha diversity compared with meadow cropping, and alpha diversity of alfalfa increased with the continuous planting years. Soil pH, total phosphorus (TP), available P, total potassium (TK), and nitrate nitrogen (NO$_3^-$) content were soil variables significantly impacting the structure of soil bacterial communities in different plant types and different alfalfa continuous cropping systems. In addition, the relative abundance of some beneficial microbial species, such as *Arthrobacter* and *Gaiellales*, in the cropping maize and continuous cropping of alfalfa was much higher than that in the meadow field. Moreover, the networks differ among different plant types, and also differ among different continuous cropping years of alfalfa, and topologies of the networks suggested that continuous planting of alfalfa promotes cooperation between bacteria, which facilitates the long growth of alfalfa and is beneficial to the soil.

Keywords: meadow, maize, continuous cropping alfalfa, bacterial structure, network

INTRODUCTION

Alfalfa (*Medicago sativa* L.) is one of the most significant perennial herbaceous legume fodder in the world that is widely grown in many countries and contributes significantly to the development of agriculture and livestock (Han et al., 2005; Raiesi, 2007; Li and Huang, 2008). In China, alfalfa grows mainly in the arid and semi-arid soil of northern China and is grown on more than 4,000 ha per year (Zhang et al., 2016). The Northeastern portion of China is an agro-pastoral area with longer winters (Chen et al., 2013). As a result, animal feed in this region is almost entirely dependent on summer pasture and winter silage, and therefore, alfalfa can alleviate forage shortages for cattle in the winter in the Northeast with its more complete nutrition and high yield (Su, 2007; Chen et al., 2013). Thus, alfalfa has been continuously grown on a large scale in northeastern China to meet the winter demand for forage and thus increase livestock productivity (Dong et al., 2003). However, continuous cultivation of alfalfa has led to an increase in pathogenic bacteria and soil acidification, and these changes have been shown to be closely linked to soil microorganisms (Yan et al., 2012; Yao et al., 2019; Liu et al., 2020).
The study examined soil samples from three agricultural systems growing responses to long continuous cropping for 6, 10, 14, 20, and 30 years in their soil microorganisms and soil quality. It is meaningful to study the changes among different cropping systems, i.e., meadow, maize, and alfalfa, and to reveal the alfalfa growing responses to long continuous cropping for 6, 10, 14, 20, and 30 years in their soil microorganisms and soil quality. Therefore, more in-depth studies in different farming systems and environments are needed to explore the mechanisms of barriers to continue cropping.

In view of the changing soil microorganisms to promote sustainable animal feed industry in northeast China, and evaluated the soil bacterial structure and soil characteristics. This study aimed to explore the soil microbial community structure of diverse crop systems and alfalfa with continuous cropping time and to assess the complete connection between soil bacterial communities and physical and chemical characteristics.

**MATERIALS AND METHODS**

**Experimental Site and Design**

The experimental location is in Furalji District, Qiqihar City, Heilongjiang Province, China (4715°N, 12341'E). Fields that continuously planted alfalfa for 6, 10, 14, 20, and 30 years were selected and coded as C6, C10, C14, C20, and C30, respectively. Moreover, the soils of the meadows and maize field were selected as the controls, which encoded Me and Ma, respectively. Each treatment is over 900 m² in size. The sowing density of alfalfa is 4,000,000 seeds ha⁻¹. The chemical compound fertilizer (N 18%, P₂O₅ 18%, and K₂O 18%) of 280 kg/ha was applied to each treatment in June each year. The fields of alfalfa are maintained using standard planting and are not greased. The alfalfa was cut to the surface in June and August each year, except for the first year when it is sown.

**Soil Sampling and Soil Characteristic Measurement**

On June 30, 2019, during the flowering time of alfalfa, soil samples were collected at 0–15 cm ground depth. A combination of over five individual soil nuclei from a total area of 500 m² was obtained from each sample. A total of 42 soil samples of meadow, maize, and 5 alfalfa fields were collected. In field conditions, about 2 g of soil samples was placed in sterilized centrifuge tubes and stored at −80°C in a refrigerator for soil DNA extraction; fresh soil was used to measure soil Physicochemical properties.

Using a pH meter, the soil pH was determined in a soil water suspension (1:5 w/v). Fifteen grams of fresh soil was dried in an oven at 105°C for 24 h to a constant weight to determine the soil moisture content. An elemental analyzer was used to measure the soil total nitrogen and carbon contents (Jones and Willett, 2006). Using the continuous flow analysis system, 2.0 M KCl was used to extract ammonium (NH₄⁺–N) and nitrate (NO₃⁻–N). Moreover, 0.5 M NaHCO₃ and H₂SO₄–HClO₄ were used to extract the total and available phosphorus, respectively. Additionally, an inductively coupled plasma emission spectrometry (ICPS-7500), HNO₃–HClO₄–HF and 1.0 M CH₃COONH₄ were used to extract soil total and available potassium, respectively (Lu, 1999).

**DNA Extraction and High-Throughput Sequencing**

Using the Fast DNA Spin Kit (MP Biomedicals, Santa Ana, CA, United States), soil total DNA was extracted from soil samples. Primers of 515F/806R were used to amplify the bacterial
16S rRNA gene (White et al., 1990), and the forward primer being modified at the 5' end with a unique 6-nucleotide barcode was added. A 20-ml PCR mix with 0.5 ml of each 10 mM primer, 10 ng of DNA template, and 18 ml of Platinum PCR SuperMix were used to produce PCR. The PCR procedure was 95°C for 5 min; 94°C for 35 s, 55°C for 15 s, 72°C for 10 s for 32 cycles, and 75°C extension for 8 min (Liu et al., 2015). All the samples were standardized to equimolar levels and sequenced on the Majorbio Biotechnology Illumina MiSeq platform. All sequences are deposited in GenBank of NCBI with the reference PRJNA760979.

The raw FASTQ data were processed with QIIME Pipeline version 1.19.1 after sequencing. In brief, each barcode-based sample was allocated to all sequence reads. Preliminary analyses were performed to eliminate sequences of low quality (length < 200 bp and average basis quality score < 30). Use the UCHIME algorithm to find and eliminate chimeras of the trimmed sequences (Edgar et al., 2011). The RDP classification was used to allocate sequences phylogenetically based on their optimal match to the RDP databases. Operational taxonomic units (OTUs) with a CD-HIT sequence similarity of 97% were categorized (Cole et al., 2009; Li and Godzik, 2015).

For the alpha diversity, Shannon and Chao 1 indices were calculated in QIIME. Additionally, principal coordinate analysis, Adonis test, and canonical correspondence analysis have been carried out in R version 4.1.1 with the “vegan” package. GenStat 13 was used to perform the one-way analysis of variance (ANOVA) to assess differences in soil chemistry and the abundance of bacteria at different taxonomic levels. Bacterial symbiotic networks were analyzed for the Me, Ma, and AC treatments, and AC6-10, AC14-20, and AC30 groups. The raw data were statistically analyzed using the “psych” package in R and then visualized in Gephi (Jiang et al., 2017). The correlation between each of the two OTUs was chosen to be p < 0.05, with Spearman correlation coefficients greater than 0.7 (Mendes et al., 2018). Identification of keystone species was based on high nodality, high intermediate centrality, and high compact centrality (Berry and Widder, 2014; Agler et al., 2016).

RESULTS

Soil Physicochemical Characters
Soil pH, NO$_3^{-}$, TK, AK, and C/N were significantly higher in the alfalfa soils, compared with the soil of meadow and maize, while NH$_4^{+}$, TP, AP, and TN showed the opposite trend. Moreover, soil NO$_3^{-}$, AK, TC, and TN contents increased with the extension time in the soil of continuous crop alfalfa, whereas the contents of NH$_4^{+}$, TP, and AP decreased with the extension time in continuous cropping alfalfa soils (Table 1).

Soil Bacterial Diversity
According to the Chao index, soil microbial diversity was highest in the AC30 treatment and lowest in the Me treatment ($p > 0.05$; Figures 1A,B). Effect of crop type and continuous

| Treatment | pH | NO$_3^{-}$ | NH$_4^{+}$ | TP | TK | AK | AP | TC | TN | C/N |
|-----------|----|------------|------------|----|----|----|----|----|----|-----|
| Me        | 5.66 ± 0.04d | 7.8 ± 0.006e | 7.6 ± 0.035c | 22.4 ± 0.11a | 7.75 ± 0.006f | 8.23 ± 0.006b | 1.47 ± 0.010b | 23.96 ± 0.039d | 1.37a | 27.75 ± 0.006f |
| Ma        | 7.69 ± 0.04a | 7.81 ± 0.006a | 7.76 ± 0.033b | 22.18 ± 0.07a | 7.75 ± 0.006f | 8.23 ± 0.006b | 1.47 ± 0.010b | 23.96 ± 0.039d | 1.37a | 27.75 ± 0.006f |
| AC6       | 7.8 ± 0.006e | 7.8 ± 0.006e | 7.8 ± 0.006e | 22.18 ± 0.07a | 7.75 ± 0.006f | 8.23 ± 0.006b | 1.47 ± 0.010b | 23.96 ± 0.039d | 1.37a | 27.75 ± 0.006f |
| AC10      | 7.8 ± 0.006e | 7.8 ± 0.006e | 7.8 ± 0.006e | 22.18 ± 0.07a | 7.75 ± 0.006f | 8.23 ± 0.006b | 1.47 ± 0.010b | 23.96 ± 0.039d | 1.37a | 27.75 ± 0.006f |
| AC14      | 7.8 ± 0.006e | 7.8 ± 0.006e | 7.8 ± 0.006e | 22.18 ± 0.07a | 7.75 ± 0.006f | 8.23 ± 0.006b | 1.47 ± 0.010b | 23.96 ± 0.039d | 1.37a | 27.75 ± 0.006f |
| AC20      | 7.8 ± 0.006e | 7.8 ± 0.006e | 7.8 ± 0.006e | 22.18 ± 0.07a | 7.75 ± 0.006f | 8.23 ± 0.006b | 1.47 ± 0.010b | 23.96 ± 0.039d | 1.37a | 27.75 ± 0.006f |
| AC30      | 7.8 ± 0.006e | 7.8 ± 0.006e | 7.8 ± 0.006e | 22.18 ± 0.07a | 7.75 ± 0.006f | 8.23 ± 0.006b | 1.47 ± 0.010b | 23.96 ± 0.039d | 1.37a | 27.75 ± 0.006f |
cropping years on the bacterial phylum (Figure 2). Principal coordinate analysis (PCoA) revealed that cropping systems and alfalfa continuous cropping time significantly affected the soil bacterial communities (PERMANOVA, $p < 0.05$; Figures 3A–D and Table 2). According to the PCoA result, we divided all treatments into three groups—Me (Meadow), Ma (Maize), and AC (Alfalfa continuous cropping)—and further divided AC into three groups—AC6-10 (alfalfa continuous cropping for 6, 10, and 14 years), AC20 (alfalfa continuous cropping for 20 years), and AC30 (alfalfa continuous cropping for 30 years) (Figure 3 and Table 2). The results of CCA revealed that there was a close relationship between soil physicochemical and soil bacterial community composition (Figure 4). Specifically, total C ($r = 0.764; p < 0.01$) and N ($r = 0.654; p < 0.01$), C/N ($r = 0.876; p < 0.01$), TP ($r = 0.732; p < 0.05$), AK ($r = 0.732; p < 0.01$) and TK ($r = 0.804; p < 0.01$), NH$_4$ ($r = 0.677; p < 0.05$), pH ($r = 0.616; p < 0.01$), and NO$_3$ ($r = 0.677; p < 0.05$) seemed significantly associated with the microbial community composition.

### Specific Microbiomes

Actinobacteria, Acidobacteria, Proteobacteria, and Chloroflexi were the phyla with the highest relative abundance across all the treatments, accounting for 72.14–78.81% of the whole community (Figure 3). Overall, the relative abundance of Actinobacteria and Proteobacteria was higher in the
Ma and AC treatments compared with the Me treatment, while Acidobacteria showed the opposite trend. On the genera level, the relative abundance with Kruskal–Wallis $H$ test showed that some genera, such as *norank_Gaiellales*, *norank_Vicinambacterales*, *Rubrobacter*, and *Arthrobacter*, were significantly ($p < 0.05$) different among the Me, Ma, and AC fields. Moreover, some genera, such as *norank_JG30-KF-CM45*, *norank_Gaiellales*, *Arthrobacter*, *Sphingomonas*, *Microlunatus*, and *Lysobacter*, were significantly ($p < 0.05$) different among the cropping systems of alfalfa continuous cropping for AC6-10, AC20, and AC30 treatments (Figure 5). In more detail, the relative abundance of *Rubrobacter*, *norank_Vicinambacterales*, *norank_JG30-KF-CM45*, *norank_Vicinamibacteraceae*, *Arthrobacter*, and *norank_Gemmatimonadaceae* was significantly higher in the Ma and AC treatments compared with the Me treatment, while the relative abundance of *norank_Acidobacteria*, *Candidatus_Udaeobacter*, and *norank_TK10* showed the opposite trend (Figure 5A). Furthermore, the relative abundance of *norank_Gaiellales*, *norank_67-14*, *norank_Gemmatimonadaceae*, and *Lysobacter* were increased with the alfalfa continuous cropping time, while *norank_JG30-KF-CM45*, *Arthrobacter*, *norank_Geminicoccaceae*, *Sphingomonas*, and *Microlunatus* showed the opposite trend (Figure 5B).

**Co-occurrence Network**

The co-occurrence network based on OTU level shows the relationship between bacteria in different treatments (Figure 6). Comparing the Me, Ma, and AC treatments, the ranking of the number of negative correlations and modularity was Me > AC > Ma, while for the average degree (avgK) and clustering coefficient (avgCC), no significant differences were
found among the treatments. When comparing the AC6-10, AC20, and AC30 treatments, the number of negative correlations, modularity, and avgCC increased with the years of continuous cropping. For the keystone species, OTU1210 (Jatrophihabitans), OTU10961 (Blastococcus), and OTU8174 (norank_Gemmatimonadaceae) were identified in the Me, Ma, and AC networks, respectively, while OTU13196 (Microlunatus), OTU5705 (Paenibacillus), and OTU8419 (norank_Xanthobacteraceae) were identified in the AC6-10, AC20, and AC30 networks, respectively (Table 3).

**DISCUSSION**

In the present study, the Ma and AC treatments have higher microbial diversity than the Me treatment, and microbial diversities increased significantly in the long-term continuous (AC30) treatment. These results suggest that maize and alfalfa were enriched with more microbial species and were more conducive to soil conservation and sustainability, at least in terms of microbial diversity. Previous studies have found that, compared with corn-soybean rotation systems, there was less
rhizosphere bacterial diversity in continuously grown soybeans (Liu et al., 2020). A positive correlation between continuous cropping years and soil bacterial diversity has also been reported (Liu et al., 2020). Nevertheless, it has also been claimed that soil microbial diversity did not differ between soils grown in continuous soybean and soybean–maize rotations (Li et al., 2010). The different results of these studies might depend on the types of soil utilized and the different years of continuous cropping. Furthermore, differences in crop genotypes may also be responsible for this phenomenon, as microbial diversity has also shown different trends due to successive plantings of resistant and sensitive varieties (Yuan et al., 2021). Changes in soil pH can affect other soil physicochemical properties, and these changes directly or indirectly influence microbial diversity (Tan et al., 2017; Lian et al., 2019). In addition, microbial diversity in different farming systems can be affected by changes in plant root secretions, such as flavonoids and hormones (Tan et al., 2017; Lian et al., 2019; Liu et al., 2020; Shi et al., 2020).

From the results of PCoA and the PERMANOVA analysis, the crop types and years of continuous alfalfa were considered the two most important factors that changed the soil bacterial structure (p < 0.05). There is no doubt that different crops
have different microbial community structures (Lian et al., 2019). This was in line with some previous studies that have shown significant variation in soil bacterial communities in short- and long-term alfalfa continuous cropping field (Zhu et al., 2017; Yao et al., 2019). The CCA result demonstrated that the major factors in changing soil bacterial community structure in different treatments in this study were soil pH, $\text{NO}_3^-$, total K, total P, and available P. Similar results were found for the significant effect of soil characteristics, such as soil pH and AP, on the structure of the bacterial community. In our investigation, these soil parameters were impacted significantly by continuous cropping, showing that continuous crops modified their soil characteristics and subsequently changed their bacterial community.

In the Ma and AC soils compared with those of the Me system, the relative abundance of Actinobacteria and Proteobacteria was substantially enhanced, suggesting that the bacteria were increased with high nutrient availability (Li et al., 2014; Yuan et al., 2021). The relative abundances of *Arthrobacter* increased in the Ma and AC cropping field compared with Me, but then decreased in the AC20 and AC30 long continuous cropping field, compared with AC6-14. Hexavalent chromium can cause serious human irritation, while *Arthrobacter* can reduce hexavalent chromium, thus making the soil environment more beneficial. Some specific metabolites of *Arthrobacter* can promote amino acid secretion from plant roots (Romaniuk et al., 2018; Shi et al., 2020). Additionally, some microbial species, such as norank _Gaiellales_ and *Lysobacter*, which play a role in ecological function of ligninolysis and in soil suppression against the fungal root pathogen, were increased with the alfalfa continuous cropping time, suggesting that these bacteria might inhibit soil fungal diseases due to long-term continuous cropping (Gómez Expósito et al., 2015). Therefore, changes in these bacteria across treatments may be related to antagonistic activity of plant pathogens and improved soil nutrition. However, the contribution of these significantly responsive microbial species to the plant is speculative based on their abundance and reported function. Whether they have a definite role in continuous cropping for alfalfa requires further verification.

Association network analysis provides a more detailed understanding of bacterial community composition and associations (Xue et al., 2018; Xiong et al., 2021). The network negative correlations and modularity of the Me were higher than that in Ma and AC treatments, suggesting that continuous planting of alfalfa promotes cooperation between bacteria, which facilitates the long growth of alfalfa and is beneficial to the soil (Yao et al., 2019). This finding corresponds to an earlier research, showing that the soil microbial structure becomes increasingly healthy after a long period of continuous cropping (Yao et al., 2019; Liu et al., 2020).

In summary, maize cropping and continuous cropping of alfalfa increased the soil bacterial alpha diversity, and alpha diversity also increased in the long-term continuous planting system. Soil pH, $\text{NO}_3^-$, total K, and total P content were important factors influencing the structure of soil bacterial community in different plant types and different alfalfa continuous cropping system. Moreover, compared with planting meadow, maize and alfalfa continuous cropping significantly increases a number of beneficial bacterial species, such as *Arthrobacter* and _Gaiellales_, suggesting that the microbial community of maize and long-term alfalfa cropping shifts toward a healthy pattern. However, these microorganisms need to be isolated and formed into synthesized microbial communities to verify their specific benefits to the crop. Furthermore, the networks differ among different plant types and also differ among different continuous cropping years of alfalfa. The topology of the networks suggested that continuous planting of alfalfa promotes cooperation between bacteria, which facilitates the long growth of alfalfa and is beneficial to the soil.

### TABLE 4 | Keystone taxa identified in the co-occurrence network.

| OTU ID | Phylum          | Class           | Order     | Family       | Genus       | Species       |
|--------|-----------------|-----------------|-----------|--------------|-------------|---------------|
| Me     | OTU1210         | Actinobacteriota| Actinobacteria| Frankiales   | Frankiaceae | Jatropha_habitans norank |
|        | OTU1028         | Proteobacteria  | Gammaproteobacteria| Burkholderiales | Rhodocyclaceae | norank norank |
|        | OTU1098         | Actinobacteria  | Actinobacteria| Frankiaceae   | norank norank | norank norank |
|        | OTU10661        | Actinobacteria  | Actinobacteria| Frankiaceae   | Geodermatophilaceae | Blastoococcus norank |
|        | OTU11804        | Acidobacteriota | Blastocatella | Blastocatellales | Blastocatellales | norank norank |
|        | OTU6973         | Proteobacteria  | Alphaproteobacteria | Rhizobiales | Rhizobiales | Allorhizobium Pararhizobium |
| AC     | OTU8174         | Gemmatimonadota | Gemmatimonadetes | Gemmatimonadaceae | norank norank | norank norank |
|        | OTU10318        | Actinobacteriota| Thermophilina | Gaiellales     | norank norank | norank norank |
|        | OTU5813         | Chloroflexi     | Chloroflexia | Thermomicrobiales | norank norank | norank norank |
|        | OTU13196        | Actinobacteriota| Actinobacteria | Propinobacteriaceae | norank norank | norank norank |
|        | OTU8174         | Gemmatimonadota | Gemmatimonadetes | Gemmatimonadaceae | norank norank | norank norank |
|        | OTU12811        | Acidobacteriota | Vininamibacteria | Vininamibacteriales | norank norank | norank norank |
| AC20   | OTU5705         | Firmicutes      | Bacilli | Paenibacillales | Paenibacillus | norank norank |
|        | OTU12040        | Proteobacteria  | Gammaproteobacteria | Burkholderiales | norank norank | norank norank |
|        | OTU11037        | Actinobacteriota| Actinobacteria | Micrococcus | Microbacteriaceae | norank norank |
| AC30   | OTU8419         | Proteobacteria  | Alphaproteobacteria | Rhizobiales | Xanthobacteriaceae | norank norank |
|        | OTU11007        | Proteobacteria  | Thermophilina | norank norank | norank norank | norank norank |
|        | OTU9218         | Proteobacteria  | Gammaproteobacteria | Burkholderiales | norank norank | norank norank |
DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/supplementary material.

AUTHOR CONTRIBUTIONS

HL and ZY conceived and designed this study. ZY and YX performed the experiments and wrote the manuscript. SL, XW, and HC analyzed the data. All authors approved the final version of the manuscript.

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FUNDING

This work was supported by Outstanding Youth Fund of Heilongjiang Academy of Agricultural Sciences (2020CQX003), China Agriculture Research System of MOF and MARA (CR34), and the Grass-field Rotation Scientist Studio of Heilongjiang Province (202004).
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