Long-Term Exclusion of Grazing Increases Soil Microbial Biomass but Not Diversity in a Temperate Grassland

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ABSTRACT

Restoration of grassland such as exclusion of grazing has been considered to increase aboveground plant diversity and soil fertility. However, knowledge on the effect of long-term exclusion of grazing on soil bacterial community structure and diversity is not well understood. The two sites were selected in the Inner Mongolian grassland, i.e., one fenced off since 1979 (UG79) and the other continually grazed by sheep (FG) all along. Soil microbial biomass was measured using fumigation method and bacterial community structure and diversity were assessed using methods of denaturing gradient gel electrophoresis (DGGE) and clone library. Results showed that the UG79 soil had significantly higher microbial biomass carbon and nitrogen compared with the FG soil. There was a clear separation in soil bacterial community structure, but not in bacterial diversity between the two sites. Moreover, 55 clones from the UG79 soil and 56 clones from the FG soil were selected and sequenced. Phylogenetic analysis of all clone sequences indicated that bacterial communities were dominated by the groups of \textit{Actinomycetes}, \textit{Proteobacteria} and \textit{Bacteroidetes}, but there were no significant differences in bacterial diversity between the two sites, consistent with the results obtained from DGGE. The results highlighted that although long-term exclusion of grazing increased soil microbial biomass, but it did not harbor higher bacterial diversity compared with freely grazed site.

Keywords: Grazing; Microbial Biomass; Community Structure; Diversity; DGGE; Clone Library; Inner Mongolia

1. Introduction

Ungulate grazing plays a vital role in carbon (C) and nitrogen (N) cycling in grassland ecosystems [1]. Overgrazing may lead to degradation of grassland [2]. Restoration of grassland by exclusion of grazing has usually been used to increase aboveground plant biomass and soil fertility, which greatly affect belowground microbial community structure and diversity [3]. Many studies have documented the effects of grazing intensity on aboveground plant communities [4], belowground microbial communities [5,6], and C and N mineralization [7,8]. However, few studies have conducted the effect of long-term exclusion of grazing on soil bacterial community structure and diversity.

Grazing has been found to decrease soil microbial biomass irrespective of changes in both the quantity and quality of litter [9]. Recently, Klumpp \textit{et al.} [6] have reported that compared with low level grazing, intensive grazing leads to a change in soil microbial communities and a proliferation of Gram(+) bacteria in a controlled experiment. However, change in bacterial communities in response to grazing are more complicated than we expected [10]. For example, McCaig \textit{et al.} [11] did not observe obvious changes in soil bacterial community structure between the grazed and non-grazed sites using method of denaturing gel gradient electrophoresis (DGGE). Jangid \textit{et al.} [12] found that soil bacterial community structure rather than diversity significantly differed along a chronosequence of restoration of grassland using methods of phospholipid fatty acid and clone libraries. Zhou \textit{et al.} [13] found that soil bacterial diversity changed monotonically with increasing grazing intensities. Therefore, it is necessary to investigate the effect of long-term exclusion of grazing on soil bacterial community structure and diversity in order to improve the sustainable management of grassland ecosystems.

The Inner Mongolian temperate grassland approximately accounts for 12% of the national area in China and acts as an important and representative part of the Eurasian grasslands [3,14]. There are two sites in Inner Mongolia selected for this study; one has been fenced since 1979 (UG79) representing an undisturbed and climax steppe community, and the other (FG) has been subjected to continually grazing outside the UG79 site.
The objectives of this study were to compare soil microbial biomass and bacterial community structure and diversity using methods of PCR-DGGE and clone libraries between the two sites. We hypothesized that long-term exclusion of grazing would increase soil microbial biomass and bacterial diversity.

2. Materials and Methods

2.1. Site Description and Soil Sampling

The experimental sites are located in Xilin River Watershed of Inner Mongolia Autonomous Region (43°32’N, 116°40’E, 1200 m a.s.l.). The region is characterized by a continental climate with dry, cold winters and warm, wet summers and the mean annual air temperature is −0.7°C. The mean annual precipitation is ca. 350 mm according to local record and most of them occurs from April to September. The soil type is a dark chestnut (Mollisol) according to FAO classification with a texture of ca. 21% clay, 60% sand and 19% silt [2].

The area being grazed by herds consisted of 70% - 90% sheep and 10% - 30% goats. Prior to 1979, the whole experimental area was generally grazed under a low intensity. In 1979, the UG79 site (24 ha) was fenced and excluded from grazing. The grassland outside the UG79 site was continually grazed (FG). The FG site is assumed to represent a heavily grazing intensity of 2 sheep units (1 SU = 1 ewe and 1 lamb) ha⁻¹·yr⁻¹ [3].

The UG79 site is dominated by perennial rhizome grasses such as Leymus chinensis and Stipa grandis which represents a widely distributed grassland community in the Eurasian grasslands [3,14]. However, the FG site is dominated by herbaceous species such as Artemisia frigida and Pedicularis acaulis. The UG79 soil had a pH of 7.1 and 21.7 mg·kg⁻¹ of soil organic C, while the FG soil had a pH of 7.2 and 18.7 mg·kg⁻¹ of soil organic C.

Three plots (20 × 30 m) within the two sites were selected at 50 m intervals. Soil samples were taken from a depth of 0 to 5 cm with a 3 cm diameter auger within each plot in August 2004. All soil cores within each plot were immediately mixed thoroughly and kept in a cooler (ca. 4°C). After passing through a 2-mm sieve, the soil samples were stored at 4°C for a week prior to analysis.

2.2. Measurement of Soil Biochemical Properties and Microbial Biomass

Soil moisture content was determined after being oven-dried at 105°C overnight. Soil pH was measured at a 1:2.5 dry soil/water ratio. Soil organic C was analyzed using a H₂SO₄-K₂Cr₂O₇ oxidation method. Soil organic N was measured using the Kjeldahl digestion method (Kjektec System 1026 Distilling Unit, Sweden).

Microbial biomass C (MBC) and N (MBN) were measured by the chloroform fumigation-extraction method [16]. Briefly, two portions of 10 g field moist soil samples were weighed, and one portion of them was fumigated with chloroform for 24 h and extracted with 0.5 M K₂SO₄ for 1 h, and then filtered through a Whatman No. 42 paper. The other proportion of soil was directly extracted as above. The amounts of total soluable organic C and total soluable N in the fumigated and unfumigated soil extracts were determined using SHIMA-DZU TOC-VCPH/CPN analyzer (fitted with a TN unit). Microbial quotient was the ratio of MBC to soil organic C.

2.3. Extraction and PCR Amplification of Soil DNA

Nucleic acids were extracted from soil samples (0.5 g dry weight equivalent) based on the procedure described in Zhou et al. [13]. Briefly, after soil samples were homogenized for 30 s in a FastPrep bead beater cell disrupter (Bio101), nucleic acids were precipitated and washed twice in 75% (v/v) ethanol. The final crude extracts were purified with Qiagen gel extraction Kit (Qiagen, Inc.).

The PCR amplification of bacterial communities was performed in a 25 μl (total volume) reaction mixture using primers of 16S rRNA gene P338F with a 40-mer GC clamp and P518R (Muyzer et al., 1993). The final concentration of different components in the mixture included ca. 50 ng of purified DNA, 0.4 μM of each primer, 200 μM of each deoxynucleoside triphosphate, 1.5 mM MgCl₂, 1 × thermophilic DNA polymerase 10 × reaction buffer (MgCl₂-free), 1.25 U per 50 μl of Taq DNA polymerase (Promega, Madison, WI, USA), and DNAse and RNAse free filter sterilized water (Liuyan Apparatus Co. Beijing, China). PCR was performed using the following procedure: 95°C for 5 min followed by 25 cycles of 94°C for 1 min, 55°C for 1 min, and 72°C for 1 min and a final extension step of 72°C for 10 min. The PCR samples were amplified on a PTC-200 thermal cycler (MJ Research Com., USA) and the final PCR products were checked on a 1.2% agarose gel stained with ethidium bromide.

2.4. DGGE Analysis

DGGE analysis was performed using a Dcode system (Bio-Rad, Hercules, CA, USA) [13,17]. The bacterial community profiles were generated on 8% polyacrylamide gels with a gradient denaturant of 40% - 60%. Electrophoresis was run at 60°C and 110 V overnight for 12 h. The gels were soaked for 30 min in SYBR Green I nucleic acid gel stain (1:10,000 dilution, FMC Bioproducts, Rockland, ME, USA), then photographed on a UV trans-illumination table with a Hewlett Packard Scanjet 5370C. The DGGE patterns were determined using the Labworks software (Labworks TM software version 4.0).
UG79 and FG sites at P
microbial quotient as well as bacterial diversity between
were performed
2.6. Statistical Analysis and Phylogenetic Analysis

The PCR products of bacterial communities from the
UVP, UK. Soil bacterial diversity index, \textit{i.e.}, Shannon
index \(H\), was calculated based on band intensity on the
gel tracks.

\[
H = -\sum P_i \log P_i
\]

(1)
\[
P_i = n_i / N
\]

(2)
where \(n_i\) is the height of the peak and \(N\) the sum of all
peak heights in the densitometric curve [13].

2.5. 16S rRNA Gene Cloning and Sequencing

The PCR products of bacterial communities from the
UG79 and FG soils were also cloned into pGEM-T easy
vector (Promega) following the manufacturers’ proce-
dure. Ligations were transformed into \textit{Escherichia coli}
DH5\(\alpha\) supercompetent cells. White colonies were screen-
ed directly for inserts and plasmid DNAs were isolated
from randomly selected clones and screened for inserts
of the expected sizes. Finally, 55 and 56 clones were at
last randomly selected and sequenced from UG79 and
FG soils, respectively. The plasmid DNAs were se-
quenced using primers T7 and SP6 on an ABI PRISM
3700 DNA analyzer (ABI Biosystems, Inc.).

2.6. Statistical Analysis and Phylogenetic Analysis

Least Significant Difference (LSD) tests were performed
to compare the differences in microbial biomass C and N,
microbial quotient as well as bacterial diversity between
the UG79 and FG sites at \(P < 0.05\). Principal components
analysis (PCA) was used to separate the DGGE patterns
of soil bacterial communities between the two sites. All
LSD tests and PCA analyses were performed using SPSS
12.0 software (SPSS Inc., USA).

The sequences from the two sites were compared to a
current database of genetic sequences (GenBank) with
the Blast (basic local alignment search tool) program of
the Ribosomal Database Project to determine their ap-
proximate phylogenetic affiliation. Sequence assembly and
manual refinement of alignments were carried out using
the multiple-alignment algorithm as implemented in
CLUSTALX 1.81. Phylogenetic analysis was carried out
using Neighbor joining (NJ) method with Jukes-Cantor
model in the PHYLIP 3.5 package. Nodal robustness on
the NJ tree was estimated by the nonparametric bootstrap
(1000 replicates). The clones were clustered into opera-
tional taxonomic units (OTUs) at a level of sequence
similarity of \(>97\%\). Finally, collector’s curves or species
abundance curves were constructed to compare bacterial
diversitys between the two sites. The nucleotide sequences of clone libraries had been
deposited in the Genbank Data Library under accession
numbers of DQ414821 to DQ414826, DQ414832 to
DQ414844 and DQ973190 to DQ973281.

3. Results

3.1. Soil Microbial Biomass

The UG79 soil had significantly higher microbial bio-
mass C (Figure 1(a)) and N (Figure 1(b)) than the FG
site. However, there were no significant differences in
microbial quotient, \textit{i.e.,} the ratio of microbial biomass C
to soil organic C between the two sites (Figure 1(c)).

3.2. PCR-DGGE Analysis of Soil Bacterial Communities

The DGGE patterns of soil bacterial communities from
each plot were highly reproducible. Although a few
dominant bacteria were observed in the both soils, some
smears occurred in the DGGE patterns, which could be
due to large amounts of different bacterial species pre-
sent in the soils. Given that each band of the DGGE pat-
ttern represented one unique ribotype, there were similar
and numbers in the UG79 (36 \(\pm\) 2) and FG (34 \(\pm\) 2.1)
soils. The results of PCA analysis indicated that the
UG79 site was clearly separated from the FG site (Fig-
ure 2). However, there were no significant differences in
diversity of the dominating members of the bacterial
communities in the UG79 (1.88 \(\pm\) 0.08) and FG (2.02 \(\pm\)
0.14) soils.

3.3. Soil Bacterial Clone Library Analyses

Prior to phylogenetic analysis, two clone libraries were
identified chimeras which showed not only typical bell-
shaped histograms from the CHECK-CHIMERA pro-
gram but also low percentages of matching with se-
quencies in the database. The UG79 and FG clone libra-
ries contained 55 and 56 sequences, respectively. Based
on comparison of the two clone libraries, we found that
YC53 from the UG79 soil and LW32 from the FG soil
were identical based on BLAST analysis (Figure 3).

Table 1 lists the phylogenetic relationships of all clone
sequences from the two sites. Many sequences belonged
to characterized groups such as groups of \textit{Proteobacteria}
and \textit{Actinomycetes}, while some sequences fell into re-
cently recognized groups such as groups of \textit{Acidobacte-
ria, Firmicutes} and \textit{Gemmataceae}. Soil bacterial
diversity did not differ between the two sites based on
the collector’s curves (data not shown), consistent with
the DGGE results. However, the groups of \textit{Proteobacte-
rria, Acidobacteria} and \textit{Firmicutes} showed large differ-
ences between the two sites (Table 1). For example, \(\beta-
Proteobacteria\) and \(\gamma\)-\textit{Proteobacteria} only occurred in the
FG soil and \(\gamma\)-\textit{Proteobacteria} only occurred in the
UG79 soil.

<table>
<thead>
<tr>
<th>UG79</th>
<th>FG</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{Proteobacteria}</td>
<td>50</td>
</tr>
<tr>
<td>\textit{Actinomycetes}</td>
<td>30</td>
</tr>
<tr>
<td>\textit{Acidobacteria}</td>
<td>15</td>
</tr>
<tr>
<td>\textit{Firmicutes}</td>
<td>5</td>
</tr>
<tr>
<td>\textit{Gemmataceae}</td>
<td>3</td>
</tr>
</tbody>
</table>

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Figure 1. Soil microbial biomass C (a) and N (b) as well as microbial quotient (c), i.e., the ratio of microbial biomass C to soil organic C at the UG79 site fenced since 1979 and the FG site subjected to continually grazing all along. * indicates significant differences at \( P < 0.05 \).

Figure 2. Principal components analysis (PCA) of soil bacterial DGGE patterns from the UG79 site fenced since 1979 and the FG site subjected to continually grazing all along.

Table 1. Relative abundance of soil bacterial clones from the UG79 site fenced since 1979 and the FG site subjected to continually grazing all along.

<table>
<thead>
<tr>
<th>Phylogenetic group</th>
<th>Relative clone abundance (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>UG79</td>
</tr>
<tr>
<td>Proteobacteria</td>
<td></td>
</tr>
<tr>
<td>Alfa-proteobacteria</td>
<td>5.5</td>
</tr>
<tr>
<td>Beta-proteobacteria</td>
<td>0</td>
</tr>
<tr>
<td>Gamma-proteobacteria</td>
<td>1.8</td>
</tr>
<tr>
<td>Delta-proteobacteria</td>
<td>5.5</td>
</tr>
<tr>
<td>Acidobacteria</td>
<td>10.9</td>
</tr>
<tr>
<td>Actinomycetes (Gram+)</td>
<td>50.9</td>
</tr>
<tr>
<td>Gemmatimonadetes</td>
<td>3.6</td>
</tr>
<tr>
<td>Firmicutes (Gram+)</td>
<td>0</td>
</tr>
<tr>
<td>Bacteroidetes</td>
<td>14.5</td>
</tr>
<tr>
<td>Unclassfied bacteria</td>
<td>7.3</td>
</tr>
</tbody>
</table>

The results of phylogenetic analysis of 111 sequences are shown in Figure 3. Seven individual clones were not related to cultured or uncultured representatives of the sequence databases and were considered novel groups. Cluster analysis of clone sequences revealed six major clusters with numerous diverse groups. Clones that exhibited >97% sequences similarity were clustered on the trees. In all 11 clusters containing two or three sequences with >97% sequence homology were observed, the UG79 and FG soils contained 8 and 7 clusters, respectively.

Among the all clone sequences, the largest major group was the Actinomycetes (50.9% and 50%, respectively). Four clones (YC49, YC52, LW45 and LW48) of
Figure 3. Neighbor-joining tree showing relationship of soil bacterial communities from the UG79 (VC) site fenced since 1979 and the FG (LW) site subjected to continually grazing all along, based on analysis of 111 clone sequences of aligned 16S rRNA sequences. Clones exhibiting >97% sequence similarity are included in numbered clusters. The scale bar indicates an estimated change of 5%. The bootstrap values (>50%) are shown below the branches.
this group exhibited 100% similarity to the culturable bacteria *Arthrobacter globiformis* (X80736), *Streptosporangium roseum* (X89949), *Amycolatopsis* sp. (AF453718) and *Bradyrhizobium* sp. (AF230720), respectively. The *Proteobacteria* was the second most dominant group (12.8% and 19.7%). In this group, LW24 had a similarity of 100% to uncultured bacterium (AY218681) in the GenBank. The third major group belonged to *Bacteroidetes* (14.5% and 14.3%). Two clones (YC4 and LW9) in this group exhibited 99% similarity to the culturable bacteria *Pseudomonas* sp. (DQ339153) and *Sphingomonas aquatilis* (AF131295), respectively.

The group of *Acidobacteria* is a recently recognized bacterial division with only several cultivated representatives; the majority of sequences in this division come from environmental clones. Phylogenetic analysis indicated that bacterial YC16, YC34 and YC43 fell into the representative subdivision (Figure 3).

4. Discussion

4.1. Soil Microbial Biomass

In this study, the heavily grazing site caused a decrease in soil organic C and N, which was consistent with previous studies in this region [2,13]. Compared with soil organic C, labile organic C such as microbial biomass was more sensitive to changes in management practices [18]. The heavily grazing significantly decreased soil microbial biomass C and N compared with the ungrazed site (Figure 1). This might be associated with larger amounts of the aboveground plant biomass at the non-grazed site, as plants provided available C for microbial communities via root exudates and litter decomposition [19]. Recently, Klumpp *et al.* [6] reported that grazing triggered soil C losses through changing plant communities and biomass and subsequently increasing decomposition of soil organic C. In this region, Cui *et al.* [20] reported that heavily grazing removed plant biomass by 67% relative to the UG79 site, which could contribute to lower soil microbial C at the FG site. However, microbial quotient did not significantly differ between the two sites, indicating a similar efficiency in conversion of organic C into microbial C [21].

4.2. DGGE Patterns and Clone Library Analysis

Soil bacterial communities have widely been assessed using methods of DGGE and clone library [5,11,13,17]. Studies have shown that the results obtained from DGGE are consistent with those from clone libraries and this finding was also confirmed in this study.

Our results showed that there was a clear separation between the non-grazed and grazed sites, which was in contrast to those reported by McCaig *et al.* [11]. Clegg [5] found that grazing affected community structure of specific groups of microorganisms compared with the non-grazed site. Klumpp *et al.* [6] reported that the grazed treatment led to a proliferation of Gram(+) bacteria rather than Gram(–) bacteria. Although the results from clone libraries did not found an obvious difference in the group of *Actinomycetes* with high G + C content, the grazed site had an obvious increase in the group of *Firmicutes* with low G + C content. Among the clone libraries, β-proteobacteria were only detected at the FG site, while γ-proteobacteria were only detected at the UG79 site (Table 1).

Actually, soil bacterial communities were greatly affected by many factors such as pH [22], soil organic C and N [5] and aboveground plant communities [23]. Jangid *et al.* [12] found that during restoration of grassland, two similar grasslands with different soil pH, organic C and N contents had significantly different bacterial communities. On the other hand, previously we found that soil bacterial diversity reached peak at the moderate grazing intensity sites, but decreased with increasing grazing intensities [13]. In light of this statement, there was similar bacterial diversity between the long-term exclusion and non-grazed sites, which supplemented our results.

The results of phylogenetic analysis of all clone sequences indicated that soil bacterial communities were dominated by the groups of *Actinomycetes*, *Proteobacteria* and *Bacteroidetes*. The group of *Actinomycetes* was also found to be dominant in a grassland soil of England [24]. However, in the Inner Mongolians steppe, the group of *Actinomycetes* accounted for a fraction of bacterial communities in this study or using plate count [25]. Barns *et al.* [26] reported that the group of *Acidobacteria* was as environmentally widespread and ecologically important as *Proteobacteria*, and the ratio of ribotype numbers of *Proteobacteria* and *Acidobacteria* to total clone numbers might reflect the nutrient status of soil [27,28]. In this study, this ratio at the UG79 (10.9) site was higher than that at the FG (1.8) site, which was consistent with the higher soil organic C at the FG site (Table 1).

5. Conclusion

Long-term exclusion of grazing increased soil microbial biomass and changed bacterial community structure compared with the grazed site using the methods of DGGE and clone library. Phylogenetic analysis of the clone libraries from the two sites indicated that the grassland soils were dominated by the groups of *Actinomycetes*, *Bacteroidetes* and *Proteobacteria*.

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