Divergent Changes in Plant Community Composition under 3-Decade Grazing Exclusion in Continental Steppe

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Abstract

An understanding of the factors controlling plant community composition will allow improved prediction of the responses of plant communities to natural and anthropogenic environmental change. Using monitoring data from 1980 to 2009, we quantified the changes in community composition in *Leymus chinensis* and *S. grandis* dominated grasslands in Inner Mongolia under long-term grazing-exclusion and free-grazing conditions, respectively. We demonstrated that the practice of long-term grazing exclusion has significant effects on the heterogeneity, the dominant species, and the community composition in the two grasslands. The community composition of *L. chinensis* and *S. grandis* grasslands exhibited directional changes with time under long-term grazing exclusion. Under free grazing, the *L. chinensis* community changed directionally with time, but the pattern of change was stochastic in the *S. grandis* community. We attributed the divergent responses to long-term grazing exclusion in the *S. grandis* and *L. chinensis* grasslands to litter accumulation and changes in the microenvironment after grazing exclusion, which collectively altered the growth and regeneration of the dominant species. The changes in the grazed grasslands were primarily determined by the selective feeding of sheep during long-term heavy grazing. Overall, the responses of the community composition of the Inner Mongolian grasslands to long-term grazing exclusion and heavy grazing were divergent, and depended primarily on the grassland type. Our findings provide new insights into the role of grazing in the maintenance of community structure and function and therefore have important implications for grassland management.

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Introduction

The composition of plant communities is controlled by many environmental factors and disturbances [1–4]. The practice of grazing is the most important land use in Inner Mongolia grasslands and has important impacts on plant community composition [5–7]. Therefore, understanding the changes in community composition under long-term grazing exclusion and continued grazing is very important for ecologists. This information can not only enhance our ability to predict the response of plant communities to natural and anthropogenic environmental change but also provide a scientific foundation for future grassland management.

Grasslands are typically characterized by complex disturbance regimes, including grazing, fire, and drought, each of which differs in scale, frequency, and intensity [4,8–10]. The effects of disturbances on grassland vegetation can vary, and empirical evidence suggests that these disturbances often interact with each other to affect the community structure [11–14]. Generally, Inner Mongolian grasslands are primarily affected by complex grazing regimes [11,15], and fire plays a relatively weak role [16,17]. The Inner Mongolian grasslands have adapted to light grazing during their millennia-long evolution. Therefore, a marked increase in grazing intensity or a decades-long period of grazing exclusion should be somewhat equivalent to a disturbance and may influence the community structure of grasslands in the region.

Currently, improved grassland management, particularly grazing exclusion is commonly used to protect or restore the Inner Mongolian grasslands of northern China. Improved structure and function of the grasslands, such as productivity [18], community composition, C and N storage [12,19] are expected to result from the implementation of measures aimed at grassland protection. However, there is little information about the long-term changes in plant community composition caused by the removal of large animals (grazing exclusion) from Inner Mongolia grasslands. In practice, understanding the grassland dynamics under such conditions is necessary for the sustainable management of the grasslands.

*L. chinensis* and *S. grandis* grasslands are widely distributed both in northern China and in the Eurasian steppe, and have been millenary subjected to graze by sheep and cattle. Using long-term data on plant community composition from 1980 to 2009, we analyzed changes in the community composition of *Leymus chinensis* and *S. grandis* dominated semiarid grasslands under grazing-exclusion and grazing-free conditions. The objectives of the study were to explore the temporal patterns in plant community dynamics over the past three decades and to assess the response of different grassland communities to long-term grazing exclusion and heavy grazing.
Materials and Methods

Study area

The study was conducted in two semiarid grasslands in Inner Mongolia, northern China, a *S. grandis* grassland (43°32′25″N, 116°33′18″E) and a *L. chinensis* grassland (43°33′06″N, 116°40′20″E), respectively. The mean annual temperature (1980–2009) for the area is 1.1°C with mean monthly temperature ranging from −21.4°C in January to 19.0°C in July. The average annual precipitation is approximately 333.5 mm (Fig. 1). The soil is chestnut (i.e., Calcic kastanozems), equivalent to Calcic-orthic Aridisol in the US soil taxonomy classification system. The vegetation of the region consists primarily of grassland plants such as *L. chinensis, S. grandis, Cleistogenes squarrosa,* and *Koeleria cristata* [16,17]. *S. grandis* grassland and *L. chinensis* grassland are widely distributed both in northern China and in the Eurasian steppe [16].

Experimental plots

In 1979, the Inner Mongolia Grassland Research Station (IMGERS) established long-term experimental plots (25 hectares each) in *L. chinensis* grassland and *S. grandis* grassland. Both of which have been fenced since 1979 to prevent large-animal grazing. Simultaneously, scientists selected an adjacent area outside of each enclosures to form (fenced-grazing) paired experimental plots. These sites therefore included two-pair experimental plots (four experimental plots in all), designated as fenced *S. grandis* grassland (SG–F), grazed *S. grandis* grassland (SG–G), fenced *L. chinensis* grassland (LC–F) and grazed *L. chinensis* grassland (LC–G). The *S. grandis* and *L. chinensis* grasslands were considered to be in excellent condition and representative of lightly disturbed, climax steppe communities when the grazing exclosures were established in 1979 [17]. From 1979–2009, grazing by large animals was prevented by fences in SG–F and LC–F; the grazing grasslands (SG–G and LC–G) were grazed freely by sheep as according to the traditional grazing practices in the region. *L. chinensis* and *S. grandis* grasslands have adapted to light grazing during their millennia-long evolution. However, accompanying the economic reforms that have taken place in China since the late 1970s, the stocking rate increased from 0.51 sheep per ha in 1980 to 1.34 sheep per ha in 2000. The stocking rate was relatively stable thereafter. Therefore, SG–G and LC–G have been subjected to light degradation owing to long-term heavy grazing during the past three decades [15]. No natural or prescribed fires have occurred in these four plots since 1979. The detail characteristics of the four experimental plots are described in Table 1.

Field sampling and measurement

Within each plot, an east-west transect of 200 m × 100 m was established and divided into five equal-sized replicate subplots (40 m × 100 m each) [18]. Field monitoring was conducted in mid-August. In each subplot, we randomly setup one 1 m × 1 m sampling quadrat during field sampling. The aboveground biomass in these 1 m × 1 m quadrats was clipped at the ground level. All living vascular plants in the sample were sorted into species and dried and weighed. Because the standing biomass of these steppe communities reached its annual peak in mid-August, our estimated community biomass approximated the aboveground net primary productivity of these ecosystems [17].

In 2009, we collected soil samples from a depth of 0–10 cm using a soil sampler (diameter 4 cm), with 5 replicates in each of the four experimental plots. The samples were air-dried in a ventilated room and cleared of roots and organic debris. Air-dried soils passed through 2-mm sieve were ground for further analysis. The soil organic C content (%) was measured using a modified Mehbius method [20]. Briefly, 0.5-g soil samples were digested with 5 ml of 1 N K₂Cr₂O₇ and 10 ml of concentrated H₂SO₄ at 180°C for 5 min, followed by titration of the digests with standardized FeSO₄. Soil organic N (%) was analyzed using the Kjeldahl acid-digestion method with an Alpkem autoanalyzer (2300 Kjeltc Analyzer Unit, FOSS, Sweden). Total P was determined by molybdenum antimony blue colorimetry. The pH of the 0–10 cm soil samples (soil:water ratio 1:5) was tested with a PHS-35 pH meter (Sartorius, Germany). The soil moisture was measured gravimetrically using 3 soil samples from the 0–10 cm layer at 10-day intervals in the four experimental plots in 2007. Similarly, the soil temperature was measured using sensors in 2007 in plots LC–F and LC–G.

Figure 1. Changes in precipitation (P) and temperature (T) from 1980 to 2009 in the region of the study. The top panel shows the average monthly precipitation (mm) and temperature ( ºC); the solid line shows the average annual precipitation and the dashed line shows the average Jan–Aug precipitation.

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Table 1. Characteristics of 4 experimental plots.

| Type               | Treatment | Location          | Soil type  | Soil organic C (g kg⁻¹) | Total N (g kg⁻¹) | Total P (g kg⁻¹) | pH  | Grassland condition | Land-use history |
|--------------------|-----------|-------------------|------------|-------------------------|-----------------|-----------------|-----|---------------------|-----------------|
| Stipa grandis      | Fenced    | 43°32′25″N, 116°33′18″E | Chestnut⁹ | 15.91±0.46²⁳          | 1.57±0.07⁶       | 0.23±0.01⁹      | 7.99±0.06⁶ | Excellent           | Fenced and no fire since 1979 |
|                    | Grazed    | 43°32′25″N, 116°33′18″E | Chestnut   | 13.94±0.58⁸          | 1.51±0.06⁴       | 0.26±0.01⁷      | 8.07±0.07⁷ | Light degraded      | Long-term free grazing and no fire since 1979 |
| Leymus chinensis   | Fenced    | 43°32′25″N, 116°33′23″E | Dark chestnut | 17.19±0.41*           | 1.66±0.05⁶ | 0.29±0.01⁸ | 7.34±0.05⁸ | Excellent           | Fenced and no fire since 1979 |
|                    | Grazed    | 43°32′25″N, 116°33′23″E | Dark chestnut | 15.47±0.75⁲⁰        | 1.59±0.10⁶ | 0.29±0.01⁸ | 7.82±0.12⁷ | Light degraded      | Long-term free grazing and no fire since 1979 |

1The soil was chestnut (i.e., Calcic kastanozems), which is equivalent to Calcic–orthic Aridisol in the US soil taxonomy classification system.
20–10 cm soil samples were sampled and measured in August 2009, and values were represented as mean ± SE (n=5).
3Values with same superscript letter in the same column denoted non-significant differences at P=0.05 level (t-test).

Data analysis

For statistical analysis, 150 non-independent samples were available from plot SG–F from 1980 through 2009. A total of 145 non-independent samples were available for plot LC–F from 1980 to 2009 because plant species data were missing for LC–F in 1986. Field sampling for the two grazed plots, SG–G and LC–G, was conducted in 1980, 1991, 1992, 1993, 1997, 1998, 1999, 2006, 2007, 2008, and 2009. Moreover, we derived the data on the community composition of the S grandis grassland and L. chinensis grassland in 1979 from the literature [17], which reported measurements made prior to the installation of the fences. In this study, the relative biomass (RB, %) of each plant species was selected to characterize the long-term changes in community composition.

To determine the small-scale compositional variability of the communities, we calculated the plant community heterogeneity at the plot scale for each year as the mean dissimilarity in the species composition of RB among the five subplots [21]. Euclidean distance (ED) was then calculated as our measure of heterogeneity. Therefore, community heterogeneity was designated as HED and calculated with equation 1.

\[
ED = \left( \sum_{i,j} \left( x_{ik} - x_{jk} \right)^2 \right)^{1/2}
\]

\(x_{ik}\) is the RB of the i-th species in the j-th sample, \(x_{ik}\) is the RB of the i-th species in the k-th sample, and S is the total number of species. Because the differences are squared, ED emphasizes the change in the abundant species from one sample to another [21].

Furthermore, a time-lag analysis of euclidean distance in community composition (designated as \(C_{ED}\)) among different years was performed to determine the rate of change in community composition under different treatments over time [22,23]. First, for comparison with the community composition in 1979, we calculated a triangular dissimilarity matrix from the species-by-time rectangular data matrix from 1979 to 2009 (the relative biomass (RB, %) of species being the variables and samples in time being the observations). Euclidean distance is also calculated as the equation 1, where \(x_{ik}\) is the RB of the i-th species in the j-th sample, and \(x_{ik}\) is the RB of the i-th species in the k-th sample, and S is the total number of species. As used here, samples correspond to measurements of community composition over time. Because differences are squared, ED emphasizes change in species from one sample to the next from 1979 to 2009. Next, the Euclidean distance values are plotted against time lag for all lags below the diagonal in the triangular resemblance matrix. Linear regressions can then be calculated for Euclidean distance as a function of the square root of the time lag. The square root transformation reduces the probability that the smaller number of points at larger time lags will bias the analysis. If the regression line is significant, positive, and linear, then it implies that the assemblage in question is undergoing directional change. If the regression line is not significant or the slope is not significantly different from zero, then it implies fluctuation or stochastic variation over time. If the slope of the line is negative, then it implies species composition is converging on a community-type characteristic of one of the early sample periods.

For statistical analysis, 150 non-independent samples were available from plot SG–F from 1980 through 2009. A total of 145 non-independent samples were available for plot LC–F from 1980 to 2009 because plant species data were missing for LC–F in 1986.
precipitation and Jan–Aug precipitation on \( H_{ED} \) and \( C_{ED} \) by performing a correlation analysis. We conducted a partial correlation analysis to evaluate the changes in the dominant and sub-dominant species and PFGs with time by controlling either the total or the Jan–Aug precipitation. All analyses were conducted using SPSS version 13.0.

**Results**

The long-term grazing exclusion enhanced the soil organic C and N concentration in \( S. grandis \) and \( L. chinensis \) grasslands, but the soil pH decreased to some extent (Table 2). Moreover, the soil moisture was significantly higher in the fenced grasslands than in the grazed grasslands, but the soil temperature was significantly lower in the fenced grasslands (Table 2).

Community heterogeneity \( (H_{ED}) \) increased significantly with time in plots SG–F \( (R^2 = 0.394, P = 0.001) \) and LC–F \( (R^2 = 0.145, P = 0.042) \) \( (\text{Fig. 2}) \). However, \( H_{ED} \) did not appear to change in plots SG–G and LC–G. For all four sites, \( H_{ED} \) was not significantly related to the total precipitation or the Jan–Aug precipitation (data not shown). A comparison of the paired-plot correlation analysis to evaluate the changes in the dominant and sub-dominant species and PFGs with time by controlling either the total or the Jan–Aug precipitation. All analyses were conducted using SPSS version 13.0.

A time-lag analysis of \( C_{ED} \) revealed a strong linear, positive regression for \( C_{ED} \) in plots SG–F and LC–F \( (\text{Fig. 3}) \). This result indicated that strong directional changes in the community composition occurred over time. The community composition of the grazed \( S. grandis \) grassland fluctuated and varied stochastically over time. However, in the grazed \( L. chinensis \) grassland, the changes in community composition were directional over time, as shown by the time-lag analysis of \( C_{ED} \) \( (\text{Fig. 3}) \). Moreover, the value of the slope was higher in plot LC–F than in plot SG–F, indicating that the changes in community composition in the \( L. chinensis \) grassland were more responsive to the long-term grazing exclusion. Interestingly, changes in LC–F, as measured by the value of the slope, were also greater than in LC–G \( (\text{Fig. 3}) \). Moreover, the results showed that an unexpected apparent decrease of the dominant species \( (L. chinensis) \) occurred in 2001 in plot LC–F \( (\text{Fig. 4}) \).

Different time trends were found for the dominant or sub-dominant plant species in the two fenced grasslands (Table 1). In SG–F, the relative biomass \( (RB) \) of \( S. grandis \) \( (\text{former dominant species}) \) decreased significantly with time, while that of \( L. chinensis \) increased significantly with time \( (\text{Fig. 4}; \text{Table 3}) \). In contrast, the

![Figure 2. Changes in the heterogeneity \( (H_{ED}) \) of the plant community with relative biomass \( (RB, \%) \). SG–F, fenced \( S. grandis \) grassland; SG–G, grazed \( S. grandis \) grassland; LC–F, fenced \( L. chinensis \) grassland; LC–G, grazed \( L. chinensis \) grassland. doi:10.1371/journal.pone.0026506.g002](https://www.plosone.org/doi/fig/10.1371/journal.pone.0026506.g002)

Table 2. Changes in soil temperature \( (\degree C) \) and moisture (%) during the growth period.

| Month | Site | Soil moisture (%) | Soil temperature \( (\degree C) \) |
|-------|------|------------------|--------------------------|
|       |      | \( \text{SG–F} \) | \( \text{SG–G} \) | \( \text{LC–F} \) | \( \text{LC–G} \) | \( \text{LC–F} \) | \( \text{LC–G} \) |
|       |      | 13.90±1.28\(^{a}\) | 10.58±1.27\(^{b}\) | 17.40±0.95\(^{a}\) | 13.40±1.33\(^{b}\) | 10.29±0.27\(^{a}\) | 11.93±0.39\(^{b}\) |
|       |      | 9.82±0.78\(^{a}\)  | 7.77±0.57\(^{b}\)  | 11.26±0.59\(^{a}\) | 8.24±0.40\(^{b}\)  | 16.68±0.47\(^{a}\) | 19.40±0.55\(^{b}\)  |
|       |      | 7.51±0.82\(^{a}\)  | 6.36±0.85\(^{b}\)  | 9.94±1.79\(^{a}\)  | 8.17±1.80\(^{b}\)  | 20.75±0.28\(^{a}\) | 23.91±0.37\(^{b}\)  |
|       |      | 5.79±0.54\(^{a}\)  | 5.05±0.51\(^{b}\)  | 9.18±1.57\(^{a}\)  | 9.10±1.79\(^{b}\)  | 18.50±0.36\(^{a}\) | 20.35±0.48\(^{b}\)  |
|       |      | 7.59±0.93\(^{a}\)  | 6.96±0.71\(^{b}\)  | 9.18±1.40\(^{a}\)  | 8.81±1.11\(^{b}\)  | 13.41±0.36\(^{a}\) | 14.90±0.42\(^{b}\)  |

\(^{a}\)SG–F, fenced \( S. grandis \); \(^{b}\)SG–G, grazed \( S. grandis \); LC–F, fenced \( L. chinensis \); LC–G, grazed \( L. chinensis \); \(^{1}\)Data were measured in 2007, and values with the same superscript letter between fenced and grazed grasslands in same site do not differ significantly at the \( P = 0.05 \) level (t-test).

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The analysis of the PFG showed that PB \( (\text{the former dominant PFG}) \) in SG–F decreased significantly with time, accompanied by a significant increase in PR \( (\text{Fig. 5 and Table 3}) \). In contrast, PR \( (\text{the former dominant PFG}) \) in LC–F decreased significantly as PB increased \( (\text{Fig. 5 and Table 4}) \). Under grazing, there were no apparent changes in the dominant species in plot SG–F. In plot LC–F, however, \( L. chinensis \) decreased significantly and that of \( S. grandis \) increased significantly \( (\text{Table 3}) \).
The heterogeneity of the plant community in the Inner Mongolia grassland increased significantly under long-term grazing exclusion. The value of heterogeneity in the fenced grassland was higher than that in the grazed grassland. Previous studies have reported that rational grazing can enhance plant diversity and evenness in the region [24,25]. The heterogeneity of the plant community was negatively correlated with the species richness in fenced grasslands, but no apparent trend was observed for grazed grasslands. Some studies have demonstrated that an intermediate frequency of disturbance (e.g., grazing and fire) can decrease the heterogeneity of the plant community in the North American prairie [1,21]. Therefore, we concluded that the exclusion of large-animal grazing from these long-term fenced grasslands resulted in the observed increase in heterogeneity, which partly supported our assumption that a decades-long period of grazing exclusion should be somewhat equivalent to a strong disturbance and may influence the community structure of grasslands in the region.

The responses of community composition to long-term grazing exclusion differed in *S. grandis* grassland and in *L. chinensis* grassland. After the 30-yr grazing exclusion, accumulation of litter in the surface soil was 172.0 ± 8.1 g m$^{-2}$ in plot SG–F and 233.8 ± 14.4 g m$^{-2}$ in plot LC–F. Accompanying the increasing accumulation of litter, soil temperature was significantly lowered in the fenced grasslands than grazing grasslands (Table 2); simultaneously, the soil moisture was significantly enhanced because the fenced grasslands could retain more snow during the winter. The accumulated litter and the decreased soil temperature impeded the formation of buds or the rhizome...
Table 3. Changes in 12 dominant and sub-dominant species (relative biomass, %) with time, controlling for Jan–Aug precipitation (partial correlation, two-tailed test).

| Species | Fenced (SG–F) | Grazing (SG–G) | Fenced (LC–F) | Grazing (LC–G) |
|---------|---------------|----------------|---------------|----------------|
| LC      | 0.845**       | 0.026          | 0.942NS       | 0.689 <0.001** |
| SG      | −0.642 <0.001** | 0.332          | 0.348NS       | 0.524 0.004**  |
| AC      | 0.606 <0.001** | −0.083         | 0.0819NS      | 0.576 0.001**  |
| CS      | 0.300         | 0.114NS        | 0.014         | 0.969NS       |
| KC      | −0.046        | 0.814NS        | −0.347        | 0.326NS       |
| AS      | 0.191         | 0.320NS        | —             | —             |
| CK      | 0.591         | 0.001**        | 0.364         | 0.301NS       |
| AF      | −0.037        | 0.849NS        | −0.644        | 0.044*        |
| AS2     | −0.057        | 0.768NS        | −0.187        | 0.604NS       |
| AP      | −0.720 <0.001** | −0.800         | 0.005**       | −0.473 0.011** |
| HA      | −0.690 <0.001** | −0.800         | 0.005**       | −0.008        |
| PP      | 0.201         | 0.296NS        | —             | —             |

Abbreviations for plant species: LC, Leymus chinensis (Trin.) Tzvel; SG, Stipa grandis P. Smirn.; AC, Agropyron cristatum (L.) Gaertn.; CS, Cleistogenes squarrosa (Trin.) Keng; KC, Koeleria cristata (L.) Pers; AS, Achnatherum sibiricum (L.) Keng; CK, Carex korshinskyl Kom.; AF, Artemisia frigida Willd.; AS2, Artemisia scoparia Waldst et Kit; AP, Artemisia pubescens Ledeb.; HA, Heteropappus altaicus (Willd.) Novopokr.; PP, Poa pratensis L.

NS, P>0.05; * P<0.05; ** P<0.01.

Partial correlation analysis was not conducted because the species was not present in the plot.

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Figure 5. Changes in the relative biomass (%) of plant functional groups in four grasslands. The plant functional groups were classified as PB (perennial bunchgrasses), PR (perennial rhizome grass), and others. SG–F, fenced S. grandis grassland; SG–G, grazed S. grandis grassland; LC–F, fenced L. chinensis grassland; LC–G, grazed L. chinensis grassland.

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Table 4. Changes in relative biomass (RB, %) of plant functional groups (PFGs) with time, controlling for Jan–Aug precipitation (partial correlation, two-tailed test).

|                    | S. grandis grasslands | L. chinensis grasslands |
|--------------------|-----------------------|-------------------------|
|                    | Fenced (SG–F)         | Grazing (SG–G)          | Fenced (LC–F) | Grazing (LC–G) |
| PFGs1              |                       |                         |               |                |
| PB                 | −0.592                | 0.001***                | 0.376         | 0.284**        |
|                    |                       |                         | 0.696         | <0.001**       |
| PR                 | 0.895                 | <0.001**                | 0.076         | 0.834**        |
|                    |                       |                         | −0.515        | 0.005**        |
|                    |                       |                         | −0.934        | <0.001**       |

1PB, Perennial bunchgrasses; PR, Perennial rhizome grass; Others, plant species except PB and PR;

NS, P>0.05; *, P<0.05; **, P<0.01.
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renewal of L. chinensis (a perennial species with rhizomes), but their effects were far less for S. grandis (a perennial bunchgrass) [26]. At both sites, the removal of litter and increased soil temperatures resulting from prescribed fires will help to increase the L. chinensis population; simultaneously, the S. grandis population can be suppressed by prescribed fires because fire can directly damage the buds of S. grandis [27,28]. Therefore, we assumed that changes in biotic factors, abiotic factors (especially in litter accumulation and its associated effects), and their interaction foster the divergent responses to long-term grazing exclusion.

Increased stocking rates had an apparent influence on the community composition of L. chinensis grasslands, but their effect on the S. grandis grasslands was minor. L. chinensis (the former dominant species) decreased significantly over time in the grazed plots. This decrease determined the directional changes in community composition (Fig. 3 and Table 3). Under natural grazing, domestic animals (primarily sheep) preferred L. chinensis to S. grandis owing to the former’s higher palatability and nutrient content [29]. Therefore, under long-term heavy grazing, the changes in L. chinensis were mainly attributed to selective feeding by sheep [24,25]. However, the increased stocking rates did not have the same effects on the community composition of S. grandis grassland because the importance of L. chinensis in this community is minor [17]. Generally, long-term heavy grazing should affect apparently the community composition of Inner Mongolian grassland, but the type of grassland will determine the outcome.

Community composition should show abrupt changes owing to changes in the dominant species under long-term grazing exclusion in arid grassland regions. An unexpected outcome was that L. chinensis decreased sharply in the fenced L. chinensis grassland after a 21-yr grazing exclusion. A plausible explanation for the sharp decrease was that the accumulated litter decreased soil temperature decreased the availability of water from small rainfall events and affected nutrient cycles. These changes impaired the formation of buds or the rhizome renewal of L. chinensis and depleted the resources for growth previously accumulated in the rhizome. Some previous studies have demonstrated that many types of disturbances, including grazing, mowing, fire, plowing and harrowing, can apparently boost the formation of rhizome and rhizome buds. The enhanced rhizome and buds would cause the density and the biomass of the L. chinensis population to increase significantly in the following year [13,26,27]. Litter accumulation has multiple effects on the community structure. These effects depended on the amount of litter [11,30,31]. Litter has been identified as the primary mechanism for structuring grassland diversity. Richness and evenness were shown to decline with increasing litter cover [32].

The foundation provided by this information is essential for the development of general conclusion about the relationship between grazing exclusion and community composition in terrestrial ecosystems.

Further studies are required to distinguish among the divergent effects of grazing exclusion on the community structure and function in semiarid grasslands. Long-term grazing exclusion should interfere with the community composition of L. chinensis grassland. Therefore, long-term grazing exclusion from those high-quality grasslands should be employed cautiously. Of course, we do not deny that grazing exclusion is a beneficial approach to the restoration of degraded Inner Mongolian grassland [11,13]. Moreover, we need to conduct further tests our plausible explanations suggested by the study by manipulating grazing, prescribed fires and other factors in the future.

Figure 6. Changes in plant species richness under different treatments. SG–F, fenced S. grandis grassland; SG–G, grazed S. grandis grassland; LC–F, fenced L. chinensis grassland; LC–G, grazed L. chinensis grassland. Results of significant tests are reported as NS, P>0.05; *, P<0.05; **, P<0.01.
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The overall results of this study indicated that long-term grazing exclusion apparently influenced the heterogeneity, dominant species, and composition of the plant communities. The response of the community composition to long-term grazing exclusion differed in different grassland types. The community composition reflected sharp changes in the dominant species of the *L. chinensis* grassland under the long-term grazing exclusion. The divergent responses of the community composition can be attributed to the properties of the dominant plant species, litter accumulation, soil temperature, soil moisture, and plant-animal interaction; however, the underlying mechanism is not yet fully understood. These divergent responses and their underlying mechanisms should be important not only for empirical and theoretical research on community dynamics and for predicting the response of community composition to climate change but also for grassland management in the future.

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**Author Contributions**

Conceived and designed the experiments: NH XH QC. Analyzed the data: NH. Wrote the paper: NH GY QC.

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