



# Grazing exclusion-induced shifts, the relative importance of environmental filtering, biotic interactions and dispersal limitation in shaping desert steppe communities, northern China

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**Abstract:** Grazing exclusion is one of the most efficient approaches to restore degraded grassland but may negatively affects the recovery of species diversity. Changes in plant species diversity should be a consequence of the ecological assembly process. Local community assembly is influenced by environmental filtering, biotic interactions, and dispersal. However, how these factors potentially contribute to changes to species diversity is poorly understood, especially in harsh environments. In this study, two management sites within a Stipa breviflora desert steppe community (typical natural steppe) were selected in northern China. In one of the two management sites, grazing has been excluded since 2010 and in the other with open grazing by sheep. In August 2016, three plots were established and 100 sampling units were created within each plot in a 5 m×5 m area at the two management sites. To assess the effects of grazing exclusion on S. breviflora steppe, we analyzed the vegetation biomass, species diversity, soil organic carbon, and soil particle size distribution using paired T-tests. In addition, variation partitioning was applied to determine the relative importance of environmental filtering and dispersal limitation. Null mode analysis was used to quantify the influence of biotic interactions in conjunction with EcoSim niche overlap and co-occurrence values. Our results demonstrated that (1) species diversity significantly decreased and the main improvements in soil quality occurred in the topsoil 0-10 cm after the grazing exclusion; (2) environmental filtering was important for community assembly between grazed and fenced grassland and this appears particularly true for soil particle size distribution, which may be well correlated with soil hydrological processes; and (3) however, competitive exclusion may play a significant role within the exclusion. The multiple pathways of assembly may collectively determine negative effects on the restoration of species diversity. Therefore, designers should be aware of the risk of reducing grazing exclusion-induced species diversity and account for manipulating processes. This in turn will reduce dominant species and promote environmental heterogeneity to maximize species diversity in semi-arid regions.

Keywords: community assembly; environmental filtering; Stipa breviflora; desert steppe; semi-arid grassland

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#### 1 Introduction

Grasslands have experienced dramatic degradation under human disturbances and global climate change, both of which threaten biodiversity and ecosystem functioning (Bai et al., 2012; Shao et al., 2016). Recently, many ecological engineering approaches have been used to restore degraded grassland such as shrub planting, grazing exclusion, species introduction, conversion of land use, or altered grassland management. Biodiversity is critical for ecosystem resilience (Coiffait-Gombault et al., 2012). Thus, the ultimate goal of ecological restoration should be to maintain biological diversity and the associated ecological services (Cadotte et al., 2011; Peco et al., 2012; Wu et al., 2014). Species diversity is a component of biodiversity that should be highly correlated to ecosystem properties such as resource use, primary production and stability (Balvanera et al., 2006). Therefore, restoration of plant species diversity is crucial for degraded grassland ecosystem and has attracted significant attentions in recent years. Grazing exclusion is a simple way to induce self-recovery and has been regarded worldwide as one of the most efficient approaches to achieve this (Wu et al., 2009; Wu et al., 2010; Jing et al., 2014). However, grazing exclusion may have negative effects on plant diversity (Jing et al., 2014). Although the negative effects of grazing exclusion on species diversity are well documented, the processes that underlie variations in species diversity and how this relates to natural vegetation restoration are not well understood.

Species diversity, measured as species richness, abundance, and Shannon diversity index, is regarded as an outcome of species interactions or community adaptation to the biotic or abiotic environment in conjunction with evolutionary factors (Rice and Westoby, 1983). The alteration of species diversity is a consequence of the ecological assembly process (Mayfield et al., 2010). The rule of ecological assembly describes community structure and composition by passing them through the following ecological filters: the external abiotic environment (e.g., topography, soil, and precipitation), internal dynamic including biotic interactions, and dispersal (Götzenberger et al., 2012; Nathan et al., 2016). Indeed, ecological assembly theories, advances, and challenges have been extensively reviewed (Silvertown, 2004; Vellend, 2010; Weiher et al., 2011; Götzenberger et al., 2012; Hillerislambers et al., 2012), and the mechanisms of community assembly that are influenced by processes operating in diverse, spatiotemporal ranges are well-known. Species found over continental scales are constrained by evolutionary history and biophysical conditions (Virtanen et al., 2006); whereas, those found within local scale community assemblies reflect external variables and internal dynamics (Irvine et al., 2009). The intra and interspecific interactions within small spatial scales that have only a few centimeters or millimeters between neighbors are likely to be influenced by biotic filters (Götzenberger et al., 2012). The community assembly processes are also disturbance dependent. Grazing is the most common disturbance regime that has a pronounced effect on species assembly (De Bello et al., 2013). Grazing directly influences competition and introduces heterogeneity into grasslands via selective grazing, treading, seed distribution, and nutrient redistribution (Jerrentrup et al., 2015). For example, selective foraging causes a local decrease in the competitive exclusion of species, thereby allowing subordinate species with low competitive abilities to coexist and increase species richness (Grime and Mackey, 2002; Mayfield et al., 2010). Moreover, trampling by grazing animals and subsequent fine-scale soil disturbances may help to create niches for gap-colonizing species (Hofmann and Isselstein, 2004). In addition, patches of bare soil lacking vegetation cover may offer space for the propagation and dispersal of seeds (Jerrentrup et al., 2015). In contrast, grazing exclusion can be followed by a spontaneous recovery of soil processes, disturbance regime, and plant-plant interactions (Coiffait-Gombault et al., 2012; Wu et al., 2014; Wang et al., 2015). Shifts in species coexistence patterns and community assembly processes may occur after grazing pressures are released through the use of mesh fencing. For example, long-term grazing exclusion enhanced the litter layer accumulation, competition for water and light resources, and competitive exclusion between different species (Guo, 2005; Wu et al., 2009). In contrast, a lack of grazing in grasslands may improve niche differentiation (Mason et al.,

2011), which promotes species coexistence and community diversity (Siepielski and Mcpeek, 2010; Thompson et al. 2010; Freschet et al., 2011). In addition, internal self-organization is a key driver for community assembly; whereas, external variables play a less significant role in community assembly after grazing pressures are released over a long term (Lawley et al., 2013).

Natural grasslands cover 40% of area in China (Hua and Squires, 2015), making it the largest ecological system in the country with important ecological functions for ecological security, including water and soil conservation, biodiversity, wind resistance, sand fixation, and air purification (Shao et al., 2016). Approximately 80% of the grasslands in the regions of western China have undergone the worst degradation in the world as a consequence of overgrazing and other human activities (Wu et al., 2014). Restoration, followed by community self-organization in arid and semi-arid grassland ecosystems, has complex outcomes driven by internal interactions and does not follow a linear, predictable trajectory (Lawley et al., 2013). Importantly, the trajectories of community development may not necessarily lead to the desired state (Andrade et al., 2015). Previous studies that focused on grassland restoration have examined changes in vegetation (e.g., composition and structure, diversity, and biomass) and soil characteristics (e.g., water content and carbon storage) in different grassland type, primarily focus on restoration evaluation (Zhang et al., 2011; Jing et al., 2014). These studies reported that grazing exclusion in semi-arid grasslands significantly improved vegetation with higher species diversity in the early stages of restoration, but decreased species diversity and biomass with increasing time of restoration. This may be due to the long-term grazing inducing litter layer accumulation, competition for water and light resources, and competitive exclusion between different species (Wu et al., 2009). Recent studies have focused on patterns of beta diversity and alteration in ecosystem functioning in China's grasslands (Bai et al., 2012; Chi et al., 2014). However, little is known about the relative contributions of environmental filtering, biotic interactions and dispersal on community assembly at a local scale. For the maintenance and restoration of diversity, understanding the relative importance of different factors in the process of community assembly at a local scale is necessary and required for establishing guidelines for restoration of degraded grassland eco-environments (Lawley et al., 2013). In this study, two management sites of Stipa breviflora steppe (typical natural steppe) were selected in a desert steppe ecosystem in northern China. One was a fenced grassland where grazing had been excluded since 2010, and the other was a sheep-grazed grassland at a level of grazing intensity of 1.5 sheep/hm<sup>2</sup>). Because S. breviflora is a typical species in continental steppes, it is regarded as a mature ecosystem that can be considered a reference system for restoration (Bai et al., 2012; Jing et al., 2013, 2014). The objectives of this study were (1) to investigate the effect of grazing exclusion on vegetation characteristics and soil quality; (2) to assess the relative contributions of environmental filtering, biotic interactions, and dispersal limitation on community assembly between the two management sites; and (3) to compare the shifts in the different processes in order to gain insight into appropriate strategies for degraded grassland restoration.

# 2 Materials and methods

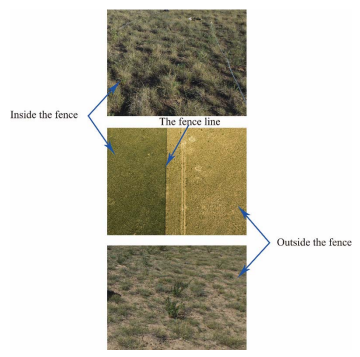
### 2.1 Study area

The experiment was conducted in Maerzhuang village (37°27′30″–37°37′30″N, 106°37′30″–106°56′15″E; 1450 m a.s.l.) in Yanchi County, Ningxia Hui Autonomous Region, China. The area is located in the southwestern fringe of Mu Us Sandy Land and in the mid-western regions of the Loess Plateau. Harsh climatic conditions in this area make up the majority of the underlying forces causing land desertification. However, overgrazing is also known to exacerbate desertification. The annual mean temperature is 8.1°C, ranging from –8.7°C in January to 22.4°C in July. The mean annual precipitation is 295.1 mm, mainly occurring in June to September (Tang et al., 2015). The mean annual potential evaporation is 2136 mm. The main soil types in this area are sierozem, loess, and orthi-sandic entisols, all of which have low fertility and loose structures (Liu et al., 2014). The representative vegetation include *S. breviflora*, *Leymus chinensis*, and

Agropyron cristatum communities. However, the representative vegetative communities have been greatly altered, primarily due to the long-term overgrazing.

# 2.2 Experimental design and data collection

The management status of the studied grasslands in the region ranges from fenced grasslands (Fig. 1), where grazing has been excluded since 2010, to grasslands that are grazed by domestic animals (mainly sheep). In late July 2016, we selected two study sites for sampling: (1) inside the fenced grassland, where livestock had been excluded for six years (from 2010 to 2016); and (2) outside the fenced grassland, where grazing was allowed all year round. The predominant plant species at the two sites were *S. breviflora* (perennial grass), and *Cleistogenes squarrosa*. Three 50 m×50 m plots (500 m apart from each other) were selected at both the grazed and fenced sites. A total of six plots were established and each plot was divided into three parts.



**Fig. 1** The landscape of study sites. The middle picture was taken using unmanned aerial vehicle (DJIS1000+Canon5D Mark III). The fence line is clearly visible with a sharp contrast in the vegetation density and cover on its both sides.

# **2.2.1** Sampling for species-area relations

There were 13 subplots (0.01, 0.02, 0.04, 0.08, 0.16, 0.32, 0.64, 1.28, 2.56, 5.12, 10.24, 20.48, and 40.96 m<sup>2</sup>) created within a nested quadrat for measurements of species-area curves. Species presence and absence were recorded in all 13 subplots. This provided information in determining the minimum sampling scale in Section 2.2.3 (Fig. 2).

# **2.2.2** Sampling for vegetation and soil inside and outside the fence

Five randomly distributed subplots (1 m×1 m each) were established within each 50 m×50 m plot to measure vegetation and soil properties. Above-ground biomass of green plants within each subplot were harvested, and the number of species, plant height, cover, abundance, and frequency for each plant species were recorded. The specific operations for a detailed vegetation survey and biomass measurement were given by a study in the same region (Tang et al., 2015). At the same time, three random soil samples in each quadrat were collected with a 5-cm diameter soil auger from four depths: 0–5, 5–10, 10–20 and 20–40 cm. Soil samples at the same depth in each subplot were then mixed to a single sample.

# **2.2.3** Sampling for variation partitioning and the pattern of species coexisting inside and outside the fence

One 5 m×5 m subplot was established in each 50 m×50 m plot for assessing variation in community assembly between the grasslands inside and outside the fence. Vegetation surveys were carried out within the subplots, which were further divided into 100 sampling units (50 cm×50 cm). Trampling from domestic animals can cause soil compaction, which alters soil structure, soil bulk density, and the transport of oxygen and water. These changes may be detrimental to plants in grasslands (Van Klink et al., 2015). Soil compaction and water content can be detected directly in the field using a stratameter and time domain reflectometry, respectively at a depth of 0-10 cm. At the same time, soil samples were taken at 0-10 cm depth. We collected a total of 100 soil samples and established 100 vegetation sampling units within each 5 m×5 m subplot. The plot scales, 0.5 and 5.0 m<sup>2</sup>, were used for assessing variation in community composition because: (1) the species-area curve saturates at 5.0 m<sup>2</sup> and the minimum size of sampling unit should be 0.5 m<sup>2</sup> (Fig. 2); and (2) co-existing species divergence may occur at the finest spatial scale. At those scales, individuals from different species are more competitive with neighbors for resources and space (Luzuriaga et al., 2012). For example, co-existing species at 50 cm×50 cm showed a convergence pattern caused by environmental filtering in dry semi-natural grasslands (De Bello et al., 2013).

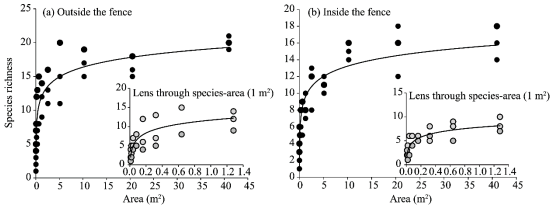


Fig. 2 Species-area curves inside and outside the fence. Two saturation points at plot size 0.5 and 5.0 m<sup>2</sup> were determined, respectively.

### **2.2.4** Laboratory analysis

In the laboratory, soil samples were air-dried and passed through a 2-mm sieve. The roots and debris were carefully picked out and discarded. The samples were then divided into three parts. One part was stored at room temperature. The second part was passed through a 0.15-mm sieve for the analysis of soil organic carbon by dichromate. The third part was used for the particle size distribution, which was described in relative percentage of clay (<0.002 mm), silt (0.05–0.002 mm), fine sand (0.1–0.05 mm), or coarse sand (2–0.1 mm). This analysis was determined using a Mastersizer 3000 particle size analyzer (Malvern Instruments, Malvern, England).

# 2.3 Data analysis

# **2.3.1** Species diversity, biomass and soil quality changes

We used the average mean of the plots to determine the differences between the treatments (inside and outside the fence). Vegetation and soil properties were analyzed using paired-sample t-tests in SPSS software. Biomass measured by Grass, Leguminosae, Compositae and weeds, respectively. Plant diversity was assessed using the Richness index (R), Shannon-Wiener index (H'), Simpson index (D), and Pielou index (J). Plant diversity indexes were calculated using the "vegan" library of the R software (Oksanen et al., 2016).

# **2.3.2** Relative roles of spatial factors and environmental filtering

Based on the mean values of species diversity matrix for each site, we employed two methods to

disentangle the relative importance of the environmental and spatial process independently or in combination. The mantel and partial Mantel test, based on the environmental differences (as a proxy for environmental filtering) and spatial distance (as a proxy for dispersal limitation), were used to detect whether the species diversity depended on the environmental and spatial distance, which was based on Euclidean distance (Wang et al., 2013).

Variation partitioning was then used to examine the relative contribution between environmental filtering and dispersal limitation based on Hellinger-transformed species data and standardized environmental data (Wang et al., 2013). First, the spatial eigenvectors (as a proxy for spatial factors) were obtained through the Principal Coordinates of Neighbor Matrices (PCNM) analysis across the coordinates of all samples. For a complete description of the method, see Dray et al. (2006). In a region where soil is nutrient deficient, some plants induce the build-up of organic matter and buffer against abiotic factors. Thus, their location may be more important in determining future spatial patterns than the soil substrate (Butterfield et al., 2010). At our study sites, the soil was of low fertility and the predominant species, S. breviflora had higher individual cover, density, and productivity in comparison with the other species. In this type of environment, S. breviflora is more likely to improve environmental conditions or enhance the survival and growth of the other species than in open spaces (Luzuriaga et al., 2012). Environmental variables tested included soil characteristics (organic carbon, water content and particle size) and aboveground biotic factors (cover and density of S. breviflora). Since there was strong collinearity between some environment variables, we first removed environment variables that were highly correlated and then used forward selection ("forward.sel" function in the R package version 0.0–8.0) to retain environment variables with a significant effect on plant diversity (Wang et al., 2013). Similarly, forward selection was also performed for the PCNM variables. The significant environmental and the PCNM variables ( $\alpha$ =0.05, Monte Carlo permutation, n=999) retained in the final model were then applied to partition variation in plant diversity into four fractions: [a]=variation explained by the environmental factors independent of space, [b]=variation explained by the spatially structured environmental factors, [c]=variation explained by spatial factors independent of any environment factors, and [d]=residual variation (Oksanen et al., 2016). All analyses were conducted in R software.

# 2.3.3 Pattern of species co-existing

To determine if the communities inside and outside the fenced grasslands were randomly or non-randomly structured, we used null model systems to analyze species co-occurrence patterns based on the mean values of the presence-absence matrices for each site (McGee and Eaton, 2015; Miguel et al., 2016). Random patterns were compared with those observed in the community to obtain information about the underlying mechanisms (abiotic filters and biotic interactions) in community composition. To quantify patterns of species co-occurrence and identify the relative strength of these relationships, we applied a standardized effect size of the co-occurrence index in EcoSim 7.0 simulation software (Gotelli and Entsminger, 2001; Zhang et al., 2009). For a complete description, see Gotelli (2000). Assuming a normal distribution of standardized effect size (SES-C), 95% of the observation was located between -2.0 and 2.0. For example, for an SES-C score, an observed value of 1.5 or larger indicates nonrandom segregation and suggests a significantly less co-occurrence than expected by chance among species. This model demonstrates whether species co-occurrence in a community may be more competitive for space and resources; however, the opposite can be said when a SES-C score is less than -1.5 (Mcgee and Eaton, 2015). The P-values in the report of EcoSim 7.0 provided information about the statistically significant difference of the observed index in relation to those obtained from 50,000 simulations (Gotelli and Entsminger, 2001). We performed the fixed-fixed null model, which has a low risk of Type I error, to simulate the randomly constructed assemblages (Gotelli and Entsminger, 2001; De Miguel et al., 2016). In this null model, both row sums (species) and all columns (plots) remain constant and equal to the original data matrix.

The niche overlap analysis was performed in EcoSim 7.0 to determine if the communities inside and outside the fenced grasslands were structured around an environmental condition or a

limited resource. The niche overlap model tests variation among species in a niche utilization. We selected the randomization algorithm conditions to exam the possibility that changes in certain conditions or resource states may be involved in structuring the community (McGee and Eaton, 2015). Furthermore, in niche overlap simulated communities, niche utilization values were randomly shuffled for each species through 50,000 interactions. The observed mean niche overlap and the mean overlap in the simulated communities, the probability of the observed mean overlap and the standard effect size (SES) were calculated. The hypotheses stated that niche overlap does not differ significantly from the expected level of overlap, even in the absence of species interactions. In general, the species partitioned the niche (an assemblage that was structured by resource partitioning) when the overlap value was significantly lower than the expected level of overlap, even in the absence of species interactions. In general, the species shared the niche when niche overlap was greater than the expected level of overlap, even in the absence of species interactions (Miguel et al., 2016).

#### 3 Results

# 3.1 Species diversity, biomass and soil quality changes

In our study sites, plant coverage increased within the exclusion (Fig. 1). Species diversity significantly decreased (P<0.05), herbage mass significantly increased (P<0.05) in grass while significantly decreased (P<0.05) in Leguminosae, Compositae and weeds within the exclusion (Table 1). The main improvements in soil quality occurred in the topsoil (0–10 cm) within the fenced grassland (S1 Table). In general, grazing exclusion may be an efficient way to restore grassland but this practice may sacrifice biodiversity.

Vegetation property	Index	Outside the fence	Inside the fence	t-test
Diversity index	R	13.73±1.87	10.73±1.58	P=0.00
	H'	$10.14\pm1.21$	7.41±1.28	P=0.00
	D	$8.04 \pm 0.94$	5.50±1.22	P=0.00
	J	$0.88 \pm 0.03$	$0.84 \pm 0.04$	P=0.00
Biomass of different family $(g/m^2)$	Grass	43.57±10.23	74.14±18.99	P=0.00
	Leguminosae	37.84±16.75	$11.89\pm6.24$	P=0.00
	Compositae	5.24±5.62	1.11±1.21	P=0.03
	Ruderal	37.79±11.86	18.99±11.98	P=0.00
	Total biomass	124.44±28.49	106.13±18.69	P=0.05

**Table 1** Changes in species diversity and biomass inside and outside the fence

Note: Plant diversity of grassland was assessed using the richness index (R), Shannon-Wiener index (H'), Simpson index (D), and Pielou index (J). Biomass measured by Grass, Leguminosae, Compositae and Ruderal, respectively. Differences in mean diversity between two sites were assessed with paired-sample t-tests. Mean $\pm$ SD.

### 3.2 Relative contributions of environmental filtering and spatial factors

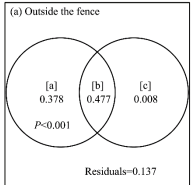
The total variation ([a+b+c]) explained by environmental and spatial factors outside the fence was greater than inside the fence. The separate components [a], [b], and [c] were different inside and outside the fence (Fig. 3). The pure effects of environmental variables [a] and spatial structure [c] turnover increased by 31.5% and by 25%, respectively, while the variation controlled by spatially structured environment variables [b] decreased by 53.5% inside the fence. Only the pure effect of environment variables [a] was statistically significant and explained a large percentage of the variation in diversity both inside and outside the fence (Fig. 3). The environmental variables were significantly related to plant diversity between the two sites (S2 Table). For example, the silt content explained 81.0% of total variation in diversity outside the fenced grassland; whereas, the coarse sand and silt content contributed 63.6% and 15.3% of total variation in diversity inside the fenced grassland, respectively. Outside the fenced grassland, the plant density was significantly correlated with space distance (P<0.05), even after controlling for the effect of environment

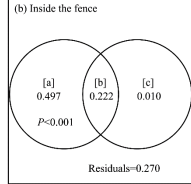
distance in a partial Mantel test, and the correlation coefficients were higher than those inside the grassland (Table 2). In contrast, the correlation inside the fence was not significant (P>0.05), and the plant diversity displayed higher correlation coefficients with environment distance, even after controlling for the space distance (Table 2).

**Table 2** Mantel test and partial Mantel test correlations for community diversity, space distance, and environmental dissimilarity inside and outside the fence

Matriana	Outside	the fence	Inside the fence		
Matrices -	R	P	R	P	
Environments distance	0.0094	0.423	0.078	0.065	
Environments space distance <sup>a</sup>	0.0095	0.394	0.061	0.161	
Space distance	0.1372	0.002	0.056	0.050	
Space environments distance <sup>b</sup>	0.1370	0.001	0.060	0.072	

Note: <sup>a</sup>, plant community dissimilarity with environments distance, controlling for spatial distance; <sup>b</sup>, plant community dissimilarity with spatial distance, controlling for environments distance.





**Fig. 3** Variation partition results for diversity inside and outside the fence. [a]=variation explained by the environmental factors independent of space, [b]=variation explained by the spatially structured environment factors, and [c]=variation explained by spatial factors independent of any environmental factors.

### 3.3 Pattern of species co-existing

The EcoSim 7.0 analysis (Table 3) showed that the community was non-randomly structured inside the fence, and there was an inter-specific competition between members of this community, suggesting a competitively structured community (SES=2.29). It also showed that the observed niche overlap was significantly less than the expected overlap inside and outside the fence (SES=-28.17 and SES=-56.2, respectively), suggesting that the differential use of resources or limitation of some environmental resource helped to structure the community.

Table 3 Co-occurrence (C-score) and niche overlap analysis of all species inside and outside the fence

C-score analysis	Comparison of Monte Carlo	<ul> <li>C<sub>obs</sub> and C<sub>exp</sub> results</li> </ul>	SES	Structure	
	P values of obs and exp	Cobs and Cexp results	SES	Structure	
Inside	$P_{\text{obs}} > P_{\text{exp}} = 0.01566$	$C_{obs} > C_{exp} = 174.7044/50000$	2.29	CCS	
Outside	$P_{\text{obs}} > P_{\text{exp}} = 0.27968$	$C_{obs} > C_{exp} = 177.4621/50000$	0.55	NCCS	
Nich overlap analysis	Comparison of Monte Carlo  P values of obs and exp	Obs and exp results	SES	Structure	
Inside	$P_{\rm obs} < P_{\rm exp} = 0.0000$	Obs <exp=0.75063 50000<="" td=""><td>-28.17</td><td>PLR</td></exp=0.75063>	-28.17	PLR	
Outside	$P_{\rm obs} < P_{\rm exp} = 0.0000$	Obs <exp=0.75082 50000<="" td=""><td>-56.20</td><td>PLR</td></exp=0.75082>	-56.20	PLR	

Note: The C-score and niche overlap values of the observed community data (obs) were compared to those of the Monte Carlo generated pseudo-community (exp), and the number of times out of 5000 iterations and the probability (*P* value) that one was greater or lower than other were calculated. From this a standardized effect size (SES) was determined, and whether the plant communities were non-competitively or competitively structured (NCCS or CCS) and were not partitioning or partitioning some limited resource (PLR or NPLR).

### 4 Discussion

The deterministic processes, such as environmental filtering and competitive exclusion, are the primary processes in shaping plant communities in the harsh environment of semi-arid grasslands (Laliberté et al., 2013; Wang et al., 2013). For instance, individual interactions and environmental filtering jointly determine community structure at the local community scale (Laliberté et al., 2013). As expected, the most significant contribution by an environment variable was the important process in community assembly inside and outside the fence (Fig. 3). Among the examined environment factors, soil particle size distribution, such as silt content outside the fence and coarse content inside the fence, explained a significant fraction of the total variance that was detected in the plant community assembly based on redundancy analysis (S2 Table). Grazing exclusion increased the fine sand and silt+caly content in the topsoil (Su et al., 2005; Wu et al., 2014). Moreover, the fenced management improved the soil structure and soil-water capability (Wu et al., 2014). Soil particles should be well correlated with soil hydrology physical processes. For example, soils that have large particles and pores can hold large amounts of water but do not retain the water since the water leaches through the pores. In contrast, clay and silt soil have low porosity and high water retention (Wu et al., 2014). Therefore, soil particle size distribution may play an important role in determining the soil water availability, which largely determines the structure of community assembly in arid and semi-arid habitats (Chesson et al., 2004; Miranda et al., 2011). Our finding showed that there were more silt and clay in topsoil inside the fence in comparison to outside the fence (S1 Table), resulting in a higher water retention in the topsoil and an increase in the vertical segregation of soil water availability. In other words, variation in the vertical soil profile may create a spatially heterogeneous water resource.

Niche overlap value of the observed community data was significantly less than the expected value inside and outside the fence (Table 3), suggesting a differential use of resources or limitation of some environment resource that could be in help to structure the community. However, both types of processes (environmental filtering and competitive exclusion) did not work together to determine species diversity and coexistence outside the fence (Fig. 3; Table 3). Disturbance has been acknowledged as one of the important factors affecting species assembly by creating ecological niches, changing soil processes, and leading to competition (Grime, 2006; Bello et al., 2013). Therefore, free grazing by sheep outside the fence may cause a dilutive effect of competition among different species, especially between annual species and perennials. For instance, selective foraging exerts a local decrease in the chance of a competitive exclusion of species, thereby allowing the subordinate species to coexist and increasing species richness (Grime and Mackey, 2002; Bello et al., 2013; Nathan et al., 2016). Grazing also introduces more heterogeneity into grasslands, which may be beneficial for structural complexity and diversity (Jerrentrup et al., 2015). Heterogeneous structures may allow different species to occupy different microhabitats and provide ecological niches promoting biodiversity (Mayfield et al., 2010). This was confirmed as the spatially structured environmental variation explained 47.7% of species diversity; whereas its explanatory power (22.2%) decreased inside the fence (S2 Table). In contrast, grazing exclusion led to an increase in relative biomass of grass, while the biomass of Leguminosae, Compositae and ruderal decreased. The transition of above-ground productivity may reflect competition among different families (Table 1), suggesting that the grass family has a greater competitive ability for space and resources. Therefore, competition between co-existing species plays a significant role inside the fence (Table 3).

The different processes, such as niche and dispersal processes, may be not mutually exclusive and jointly determine community assembly across different scales, regions and vegetation types (Wang et al., 2013; Li et al., 2015). Contrary to this viewpoint, however, the community composition in this study was controlled by a negligible effect of space that was independent of environment (Fig. 3). Seed limitation strongly influences plant community assembly in grassland (Marteinsdóttir, 2014). The dispersal filter was not acting in a semi-arid grassland system dominated by a tussock-forming *Stipa tenacissima*, because there was a dense and evenly distributed soil seed bank (Luzuriaga, 2012). In our study region, mean annual wind velocity was

2.8 m/s and the prevailing winds mainly occurred in April and May. If we assume that the seed bank was evenly distributed in this region, the climate condition may have been beneficial for seed dispersal. In addition, patches of bare soil created by tramping may have offered space for propagation and dispersal of seeds (Luzuriaga, 2012).

#### 5 Conclusions

Our findings indicate that in semi-arid grassland ecosystems dominated by tussock-forming *S. breviflora* (perennial grass), rapid changes in topsoil quality that occur following grazing exclusion were associated with a fast recovery of biomass production in grass. The different processes, environmental filtering and competitive exclusion processes, were not mutually exclusive and collectively determined community assembly after grazing the exclusion. Shifts in the relative importance of different processes may be due to certain ecological changes. First, grazing exclusion increased the fine sand and silt+clay content in the topsoil, which may correlate with soil hydrology and influence the soil water availability. Second, a rapid increase in the relative biomass of the grass family may enhance its competitive ability for space and resources. It can be inferred that grazing exclusion may be an efficient way to restore grassland but this practice may sacrifice biodiversity. For the maintenance and restoration of diversity, designers should be aware of this risk and account for the steering community assembly to maximize species diversity, especially when manipulating processes that introduce new species and promote environmental heterogeneity.

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fence

Appendix
S1 Table Soil organic carbon content and particle size distribution to a depth of 40 cm inside and outside the

Parameter	Sites	Soil depth				
raiametei	Sites	0–5 cm	5-10 cm	10-20 cm	20–40 cm	
	Outside the fence	0.27±0.07	0.27±0.09	0.4±0.12	0.32±0.12	
Soil organic carbon (%)	Inside the fence	$0.29\pm0.07$	$0.34 \pm 0.22$	$0.37 \pm 0.11$	$0.27 \pm 0.08$	
	t-test (P value)	0.41	0.27	0.58	0.12	
	Outside the fence	$2.06\pm0.48$	$2.92\pm0.73$	4.96±1.35	$5.36 \pm 0.92$	
Clay (<0.002 mm) (%)	Inside the fence	2.44±0.69	4.36±1.85	7.23±1.78	$2.34\pm0.96$	
	t-test (P value)	0.10	0.01	0.00	0.00	
	Outside the fence	18.52±3.63	$25.37 \pm 6.75$	$41.41\pm9.82$	44.22±8.09	
Silt (0.002-0.05 mm) (%)	Inside the fence	19.79±4.88	$30.95 \pm 7.99$	$44.31\pm4.09$	19.83±7.4	
	t-test (P value)	0.46	0.06	0.31	0.00	
	Outside the fence	35.35±1.21	$31.75\pm2.33$	24.94±4.11	23.7±3.47	
Fine sand (0.05–0.1 mm) (%)	Inside the fence	$32.43\pm3.2$	28.16±3.74	22.94±2.18	$34.5 \pm 3.63$	
	t-test (P value)	0.01	0.00	0.14	0.00	
	Outside the fence	44.06±3.5	39.97±5.33	$28.69 \pm 6.96$	$26.72\pm5.72$	
Coarse sand (0.1–2 mm) (%)	Inside the fence	45.34±4.45	$36.53 \pm 6.03$	25.51±3.33	43.33±5.77	
	t-test (P value)	0.38	0.15	0.12	0.00	

Note: The difference between soil organic carbon (SOC) contents wasn't statistically significant inside versus outside the fence. SOC inside the fence increased at depths of 0-5 and 5-10 cm, and declined at depths of 10-20 and 20-40 cm when compared with outside the fence. The soil particle contents inside the fence, such as clay (<0.002 mm) content, was significantly (P<0.05) increased at depth of 0-20 cm but decreased at a depth of 20-40 cm. The silt content at depths of 0-5, 5-10 and 20-40 cm exhibited differences similar to those of the clay contents inside the fence. The fine sand content decreased significantly (P<0.05) at depths of 0-5 and 5-10 cm but increased significantly (P<0.05) at depth of 20-40 cm compared to outside the fence lnside the fence had higher coarse sand content (0.1-2 mm) at all depths than the outside, but only at depth of 20-40 cm was statistically significant (P<0.05).

**S2 Table** Explanatory environmental and spatial variables selected by the forward selective procedure in the redundancy analyses inside and outside the fence

Sites	Variables	Order	Variables	$R^2$	$R^2$ Cum	Adj R <sup>2</sup> Cum	F	P value
	Environment	1	Silt	0.810	0.810	0.808	417.013	0.001
		2	Coarse sand	0.006	0.852	0.849	28.039	0.001
		3	Cover	0.043	0.859	0.854	4.417	0.035
		1	PCNM4	0.137	0.137	0.129	15.614	0.001
		2	PCNM20	0.069	0.206	0.190	8.389	0.003
		3	PCNM9	0.061	0.267	0.244	7.950	0.005
Outside		4	PCNM8	0.059	0.325	0.297	8.239	0.007
the fence	Space	5	PCNM2	0.037	0.362	0.329	5.481	0.022
		6	PCNM10	0.036	0.399	0.360	5.639	0.019
		7	PCNM21	0.033	0.432	0.388	5.278	0.015
		8	PCNM17	0.029	0.460	0.413	4.868	0.018
		9	PCNM28	0.029	0.489	0.438	5.036	0.024
		10	PCNM40	0.028	0.517	0.463	5.241	0.016
		11	PCNM14	0.025	0.542	0.485	4.722	0.030
Inside the fence		1	Coarse sand	0.636	0.636	0.633	171.536	0.001
	Environment	2	Silt	0.153	0.789	0.785	70.099	0.001
		3	Cover	0.010	0.799	0.793	4.782	0.022
		4	Fine sand	0.012	0.811	0.803	6.138	0.015

To be continued

								Continued
Sites	Variables	Order	Variables	$R^2$	R <sup>2</sup> Cum	Adj R <sup>2</sup> Cum	F	P value
	Space	1	PCNM25	0.070	0.070	0.061	7.426	0.006
Inside the fence		2	PCNM35	0.062	0.133	0.115	6.952	0.003
		3	PCNM39	0.051	0.183	0.158	5.973	0.015
		4	PCNM13	0.046	0.230	0.197	5.732	0.015
		5	PCNM6	0.041	0.271	0.233	5.342	0.020

Note: We used forward selection for all environmental and PCNM variables to obtain the significant environmental and PCNM variables, respectively. Outside the fence, the silt, coarse sand content, and the cover of *S. breviflora* were the main environmental variables; and eleven PCNM variables (#2, 4, 8, 9, 10, 14, 17, 20, 21, 28, and 40) were selected as the main spatial variables. Together, these explanatory environmental and PCNM variables explained 85.4% and 48.5% of the total plant diversity variances, respectively. Inside the fence, the coarse sand, silt, fine sand content, and the cover of *S. breviflora* were the major environmental factors extracted; and five PCNM variables (#6, 13, 25, 28, 35, and 39), representing the major spatial predictor were generated. These explanatory environmental and PCNM variables accounted for a total of 80.3% and 23.3% of the plant diversity variance, respectively.