The effect of simulated sheep grazing on male and female reproductive performance in *Caragana microphylla* Lam. (Leguminosae)

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Abstract Understanding the reproductive response of host plants to herbivores is important in grazing ecology and grassland management. Simulated grazing experiments were conducted to determine the influence of different grazing intensities on reproductive performance of a shrub, *Caragana microphylla* Lam. The total leaf mass, total flower mass, total flower mass allocation, and single flower mass allocation decreased with increased grazing intensity. The total spine mass, single flower mass and total spine mass allocation increased with increased grazing intensity. The stem mass, stem mass allocation and total leaf mass allocation had no significant change with the increasing grazing intensity. Under heavy grazing treatments, the host plants significantly decreased their investment in reproduction and increased investment in physical defense organs. Although there were no significant differences in the number of ovaries among different grazing intensities, herbivory negatively affected reproductive performance, including the number of flowers, the number of pollen grains per flower, the number of ripe seeds and the rate of pod-set in host plants. These results indicate that there are trade-offs among vegetative and reproductive and defensive organs. Compared with male reproduction, female reproductive performance was less sensitive to herbivory and grazing intensity. Moreover, pollen grains from heavily browsed plants seemed to be less likely to sire pods and ripe seeds than those from unbrowsed plants, indicating that herbivory not only decreased pollen production, but also adversely affected pollen performance.

Keywords *Caragana microphylla*; herbivory; Inner Mongolia steppe; reproduction; saliva; trade-off

Introduction

The relationship between animal and plant is one of the important aspects of research in ecology (Nagaoke, 2012; Massad, 2013). For individual plants, herbivore leads not only to a loss of photosynthetic area, but also to a loss of stored nutrients that would otherwise be available for growth and reproduction (Quesada et al., 1995; Mundim et al., 2012). Therefore, grazing animals can affect plant metabolism, growth and reproduction (Rebek and O’Neil, 2005). Several studies have provided experimental evidence of the morphological and secondary metabolic response of host plants to herbivory (Maleck and Dietrich, 1999; Sell, 2000; Brookshire et al., 2002; Musser et al., 2012; Francois, 2013). However, these browsing studies were based only on one stocking rate, and few considered differences in sexual reproduction (Rebek and O’Neil, 2005; Zhang, 2017).

Reproduction is an important stage of plant life-history (Buchanan and Underwood, 2013), and compensatory growth may happen if the plant is browsed. Therefore, compensatory growth is produced by browsed plants at the expense of sexual reproduction (du Toit and Olff, 2014). Many reports have described the effects of herbivory on the reproduction of host plants, such as flowering phenology (Brody, 1997; Juenger and Bergelson, 1997; van Kleunen et al., 2004), the number of flowers and flower size (Krupnick and Weis, 1999), sexual expression (Leather, 2000), fruit-set and seed-set (Mueller et al., 2005), indicating that herbivory negatively affects plant reproduction (Stephenson, 1982; Schemske, 1988; Krupnick and Weis, 1999; Maschinski, 2001). Yet, there is little information on the effects of herbivory on pollen quality.

We explored the mechanisms through which a simulated sheep grazing experiment affected male and female reproductive success in a perennial shrub, *Caragana microphylla* Lam. *C. microphylla* is a widely distributed shrub species in the northern steppe and agro-pastoral ecotone of China, and propagate through sexual reproduction (He et al., 2010). In these regions, *C. microphylla* has a great ecological and economic value, and plays a key role in vegetation succession from active dune to sandy grassland (Zhang, 1994). *C. microphylla* is a perennial xerophytic shrub with a plant height of 40–70 cm, stipules persistent, hardened needle-like, pinnate compound leaves, solitary yellow flowers, bell-shaped calyx and cylindrical pods (Cong, 2017). Generally, it germinates in mid-early April, spreads its leaves in mid-early May, blooms in mid-late May, bears fruit in mid-early June and matures in mid-late July.
The objective of our study was to estimate three aspects of herbivore effects on the reproduction of *C. microphylla*. First, what effects do different grazing intensities exert on biomass allocation of the plant? Second, do male and female reproductive performances respond differently to grazing intensity? Third, does herbivory affect the success of reproduction, especially, pollen quality?

**Materials and methods**

**Study area**

The Inner Mongolia Grassland Ecosystem Research Station established an experimental grazing field in 1989. It is located at Bayan Siler County, Xilinhot City (N43°37', E116°43'), mostly at an elevation of 1000m above sea level or more (Li et al., 1999). The regional climate is continental, with a 30-year average annual rainfall of approximately 350 mm (200–500 mm), 60 to 70% of which falls between July and August. Annual mean temperature is –0.4°C, and average monthly temperature is –23°C in January and 17.9°C in July. There are 150 to 180 favorable days for plant growth per year. The predominant plant species are *Artemisia frigida* Willd. (Asteraceae) and *Cleistogenes squarrosa* (Trin.) Keng (Poaceae) (Wang et al., 2001). Grazing plots were set up using a randomized complete block design with different grazing intensities, namely no grazing, light grazing, moderate grazing, and over grazing, and each treatment had three 1 hm² plots (Wang et al., 1998).

**Plant materials**

*C. microphylla* is a typical C₃ legume shrub with compound leaves and spines in its stem, varying in height from 0.3 to 1.3 m depending on grazing intensity, and shows great resistance to ungrazed herbivory. Adult plants of *C. microphylla* blossom out many simple flowers in its stem very early in June, which will stand for about 10 days. In the host plant flower, ten stamens pose diadelphous stamen and there are one carpel and one locule, but the number of ovules in its ovary is variable. Male and female organs mature at nearly the same time. Flowers of *C. microphylla* are self-incompatible and require pollinator visitation or wind agitation to set pod and seed (Xu et al., 1988). Pods mature in early September.

**Experimental design**

The grazing experiment was ceased in 2012, after which we simulated grazing by applying sheep saliva in the experiment so as to control the defoliation intensity of *C. microphylla*. Twenty shrubs in each grazing intensity plot were chosen randomly and tagged. Different defoliation intensities were simulated by clipping an appropriate length of stems on May 20 in 2012. In the light grazing plot, we removed 20% of stem length to simulate light grazing (LB), 35% for moderate grazing (MB) and 55% for heavy grazing (HB).

In terms of the effect of saliva on plant growth (Zhang et al., 2007), we clipped the chosen branches and applied saliva immediately. Sheep saliva was collected by inserting a sponge into the mouth of a two-year old wether at the experimental site. The sponge was sterilized with 70% alcohol and then dried before use. When enough saliva had been collected in a tube, the chosen plants were clipped and saliva was applied immediately to the damaged parts. About 1.5 ml saliva was applied to each branch.

To estimate the biomass allocation of the shrub, 10 branches of different individual shrubs of similar diameter were chosen and tagged in four grazing densities (no grazing (NB), LB, MB and HB) on May 31, 2012. On 10 June, when the host plant was in bloom, we clipped 10 cm length branches (excluding the annual branches) and divided them into four parts, namely the stem, leaf, flower, and spine. Then all samples were oven-dried at 65°C 48 h to a constant weight. At the same time, we collected and counted the pollen from 10 flowers produced on 10 experimental branches in each plot in order to estimate male reproductive performance. We carefully removed anthers from flower buds and developing anthers (prior to pollen dehiscence) were pulled from the corolla using fine-point tweezers. The anthers removed from each flower were placed into a labeled 1.5 ml open Eppendorf tube in a closed dry cabinet. On June 25, tubes were sealed in 1% NaCl solution and samples were later processed and counted using a hemocytometer with a microscope (Solomon, 1986). Then we quantified the total number of pollen grains produced per flower.

On June 10, we counted the number of flowers on 10 cm length branches on another 10 branches chosen in each plot. In order to ensure that pollen is derived from the same treatment conditions and to avoid pollen of other treatments, when estimate the pod-set and seed-set for each treatment, flowers were emasculated and pollinated by rubbing dehiscing anthers from other plants onto their stigmas, and were then bagged. The pollen donor was several meters away from the pollen recipient in a plot of the same grazing density. After the flowers withered, they were sprayed with insecticide to prevent insects grazing the pods. Mature pods were collected from each chosen branch prior to pod dehiscence (September, 5–15). The number of pods, seeds in undeveloped ovules, aborted seeds and ripe seeds in each pod were counted (the total ovule number per pod being the sum of the latter three variables). We randomly selected a sample of 20 seeds from each plot with 10 replications to estimate mean seed mass. The rate of pod-set per chosen branch was calculated as the ratio of mature pods to the total number of flowers. The rate of seed set was calculated as the ratio of the number of ripe seeds to the total number of ovules per pod.

To determine the effect of different grazing intensities on pollen performance, outcrossing rates were compared for 3 plants between NB and HB treatments. We chose and tagged 45 flowers from three plants in each plot, then the following treatments were applied: (1) open pollination...
we conducted a diagnostic test of normal distribution and homoscedasticity, and the result is very satisfactory. In addition, to estimate the difference of experimental traits among different grazing intensities, post-hoc multiple comparisons were performed using Tukey B test (Tukey’s-b) at 0.05 level. Correlation coefficients for the biomass of host plants in different modules were calculated using partial correlation.

When estimating the effects of pollination experiments, data were statistically analyzed using a two-way ANOVA procedure with Type III sums of squares to test the effects of both grazing intensity (NB and HB) and different pollen donor (pollen from NB or HB anthers) on female performance, such as the rate of pod-set and seed-set. In addition, correlations between the parameters tested were analyzed using Pearson’s correlation matrix.

Results

Biomass allocation

On the branches chosen, both total leaf and flower biomass decreased with grazing intensity and the only significant difference of the two index was observed between NB and HB treatments ($F_{3,12} = 3.4, P = 0.034$; $F_{3,12} = 6.698, P = 0.023$, respectively), whereas total spine biomass significantly increased with increasing grazing intensity ($F_{3,12} = 45.04, P < 0.001$) (Table 1). Therefore, host plants that were heavily browsed significantly decreased investment in vegetative (leaves) and reproductive (flowers) organs at the cost of an increase in defensive investment (spines). Significantly negative correlations were found between the biomass of vegetative and defensive organs ($R = -0.521, P = 0.015$) (Table 2). Although herbivory evidently influenced the total number of flowers (Table 1), the single flower mass of HB plants was significantly higher than that of the three other grazing intensities ($F_{3,12} = 5.488, P = 0.013$). These results indicate trade-offs among the vegetative, reproductive, and defensive modules for *C. microphylla* under grazing.

Male and female reproductive performances

Grazing intensity affected the reproductive performances of male and female organs (Table 3). There were significant

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Table 2. Partial correlation matrix among all counted biomass modules for *Caragana microphylla* under different browsing intensities.

| Variable                  | Stem mass | Total leaf mass | Total spine mass | Total flower mass |
|---------------------------|-----------|-----------------|------------------|------------------|
| Total leaf mass allocation| 0.036     | 0.400           | -0.521*          | -0.472*          |
| Total spine mass allocation| -0.207    | -0.521*         | 0.000            | 0.000            |
| Total flower mass allocation| -0.153    | -0.400*         | 0.599*           | -0.197           |

*Correlation is significant at the 0.05 level; **Correlation is significant at the 0.01 level.

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Table 1. Results of one-way ANOVA for the effect of different browsing intensities on biomass allocation of chosen branches.

| Variable                  | $F(df_1,df_2)$ | No browsing | Light browsing | Moderate browsing | Heavy browsing |
|---------------------------|----------------|-------------|----------------|-------------------|---------------|
| Stem mass                 | 0.912 (3,12)   | 0.466±0.02a | 0.457±0.085a   | 0.468±0.008a      | 0.466±0.033a  |
| Total leaf mass           | 3.4 (3,12)     | 0.318±0.033a| 0.245±0.011ab  | 0.245±0.020ab     | 0.210±0.004b  |
| Total spine mass          | 45.04 (3,12)** | 0.037±0.002d| 0.057±0.0002c  | 0.145±0.009b      | 0.193±0.005a  |
| Total flower mass         | 6.698 (3,12)*  | 0.424±0.039a| 0.310±0.035a   | 0.340±0.110a      | 0.123±0.024b  |
| Single flower mass        | 5.488 (3,12)   | 0.024±0.002b| 0.017±0.001b   | 0.022±0.005b      | 0.041±0.011a  |
| Stem mass allocation      | 0.318 (3,12)   | 0.373±0.021a | 0.421±0.012a   | 0.394±0.044a      | 0.423±0.079a  |
| Total leaf mass allocation| 1.962 (3,12)   | 0.249±0.016a | 0.226±0.014a   | 0.201±0.003a      | 0.208±0.018a  |
| Total spine mass allocation| 31.27 (3,12)** | 0.030±0.002d | 0.053±0.003c   | 0.123±0.019b      | 0.193±0.024a  |
| Total flower mass allocation| 7.148 (3,12)** | 0.330±0.019a | 0.285±0.028a   | 0.266±0.065a      | 0.127±0.037b  |
| Single flower mass allocation| 6.295 (3,12)*  | 0.018±0.001a | 0.158±0.001a   | 0.017±0.002a      | 0.048±0.012b  |

*The mean difference is significant at the 0.05 level; **The mean difference is significant at the 0.001 level. The different letter within a row indicates the significant difference at the 0.05 level. DF1 is the freedom of between groups; DF2 is the freedom of within groups.

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Data analysis

To estimate biomass allocation, all data on the rates of pod and seed-set, and pollen production, were statistically analyzed using the one-way ANOVA procedure of SPSS version 10.0 (Chicago, IL, USA). Before the analysis, all data were log transformed to improve the homogeneity of variances. We first performed a diagnostic test of normal distribution and homoscedasticity, and the result is very satisfactory. Then, we conducted a diagnostic test of normal distribution and homoscedasticity, and the result is very satisfactory. In addition, to estimate the difference of experimental traits among different grazing intensities, post-hoc multiple comparisons were performed using Tukey B test (Tukey’s-b) at 0.05 level. Correlation coefficients for the biomass of host plants in different modules were calculated using partial correlation.

When estimating the effects of pollination experiments, data were statistically analyzed using a two-way ANOVA procedure with Type III sums of squares to test the effects of both grazing intensity (NB and HB) and different pollen donor (pollen from NB or HB anthers) on female performance, such as the rate of pod-set and seed-set. In addition, correlations between the parameters tested were analyzed using Pearson’s correlation matrix.
Pollination treatment (No browsing). From the partial Eta squared results, ** = 0.253 > Eta^2 = 0.05. The mean difference is significant at the 0.05 level; The different letter within a row indicates the mean difference is significant at the 0.01 level. In addition, ** = 0.749 > Eta^2 = 0.342) for NB plants showed the number of ripe seeds produced of C. microphylla, increased. Mainly as a result of the reallocation of available resources, this adaptive grazing strategy is a trade-off strategy for plants to survive which reduce the efficiency of plant sexual reproduction caused by total flowers and total flower biomass (Feller, 1996). Wang et al. (2015) showed that plants avoid excessive grazing damage to their growth.

Table 3. Results of one-way ANOVA for the effect of browsing intensity on sexual reproduction.

| Variable               | F (df, df_e) | No browsing       | Light browsing    | Moderate browsing | Heavy browsing |
|------------------------|--------------|--------------------|-------------------|-------------------|---------------|
| Number of flowers      | 89.99(3,15)** | 18.80±0.41a        | 18.00±0.82ab      | 15.50±1.43b       | 3±0.63c       |
| Rates of pod-set       | 11.903(3,15)** | 0.79±0.02a         | 0.78±0.01a        | 0.70±0.04a        | 0.29±0.07b    |
| Number of pollen grains per flower | 3.242(3,36) | 63125±22214.64a    | 53500±3578.49ab   | 51500±4349.33b    | 46500±1848.94b |
| Number of ovules per flower | 3.41(3,31) | 17.12±0.74a        | 16.85±0.78a       | 17.48±0.42a       | 18.24±0.66a   |
| Number of ripe seeds per pod | 12.269(3,31)** | 5.30±0.19a        | 3.14±0.35b        | 3.20±0.65bc       | 2.56±0.40c    |
| Seed mass (mg)         | 24.67(3,36)** | 2.5±0.57a         | 2.1±0.1b          | 2.0±0.12bc        | 1.9±0.1c      |

*The mean difference is significant at the 0.05 level; **The mean difference is significant at the 0.01 level; The different letter within a row indicates the significant difference at the 0.05 level. DF1 is the freedom of between groups; DF2 is the freedom of within groups.

Pollen performance

For NB plants, T0 pollination significantly increased the rate of pod-set and the number of ripe seeds compared with T1 pollination treatment (P = 0.007) (Fig. 1). However, for the HB treatment, the rate of pod-set and the number of ripe seeds in T2 treatments were 1.54 and 1.63 times greater than under T1 pollination, respectively (Fig. 1). In addition, there were no significant differences between T0 and T1 pollination treatments (P = 0.057, 0.188, respectively) for HB plants (Fig. 1) and pollination method did not significantly influence the number of ripe seeds (P = 0.342) for NB plants (Fig. 1). Heavy grazing significantly decreased the rate of pod-set and the number of ripe seeds for both open and T1 treatments, whereas there were no significant differences between NB and HB treatments for T2 pollination (P = 0.31, 0.698, respectively; Fig. 1).

Interactive effects of grazing intensity and pollination method on the rate of pod-set and number of ripe seeds were observed (Table 4). From the partial Eta squared results, these effect were influenced to a greater extent by grazing intensity than by pollination method (for rate of pod-set, Eta^2_grazing intensity = 0.749 > Eta^2_pollination = 0.599; for ripe seed, Eta^2_grazing intensity = 0.253 > Eta^2_pollination = 0.187).

Discussion

In a browsing environment, grazing usually stimulates host plants to invest more resources in defensive organs (Simms and Rausher, 1987) and compensatory growth (Noy-Meir, 1993; Wang et al., 2015). In our study, similar results were found for the biomass of leaves and flowers of the host plant, C. microphylla, decreased, and the biomass of spine and single flower increased with the increase in grazing intensity (Table 1). The total flower biomass and the number of flowers of the host plant are significantly decreased under heavy grazing, but the single flower biomass increased. Mainly as a result of the reallocation of available resources, this adaptive grazing strategy is a trade-off...
Table 4. The effect of browsing intensity and hand pollination on pod-set per experimental branch and number of ripe seed per pod. Data is obtained through correlation analysis.

| Factors                           | d.f. | Pod-set per experimental branch | Number of ripe seed per pod |
|-----------------------------------|------|---------------------------------|-----------------------------|
|                                   | F    | Partial Eta²                    | F                           | Partial Eta² |
| Browsing intensity                | 1    | 35.89*                         | 0.749                       | 15.21**      | 0.253       |
| Hand pollination                  | 2    | 8.98**                         | 0.599                       | 5.03*        | 0.187       |
| Browsing intensity × hand pollina- | 2    | 13.81*                         | 0.697                       | 4.49*        | 0.166       |
| tion error (d.f.)                 | 0.02(12) | 1.588(12)                  |                             |              |              |

The residual mean squares with their degrees of freedom in parentheses are also given in the bottom row. *Correlation is significant at the 0.05 level; **Correlation is significant at the 0.01 level.

by increasing their investment in thorns (compensatory growth). Rusch et al. (2009) also suggest that grazing significantly reduces the traits of the plants, including the biomass of the entire plant, the biomass of the leaves, the biomass of the stems, and the biomass of the seeds.

Sexual reproduction is considered to be an important carbon sink for plants. (Gowda and Palo, 2003). Due to grazing, the plant photosynthetic organs were damaged (Krupnick and Weis, 1999; Milchunas and Sternberg, 2011), and the available resources of sexual reproduction of plants were reduced (Pollice et al., 2013; González-Megías, 2016). Herbivory negatively influenced male and female reproductive performance (Juenger, 2000). In this research, the number of flowers and pollen grains decreased with the increase of grazing intensity, while the ovules did not change significantly (Table 3). The results were consistent with the performance of Isomeris aborea, whose performance of male reproduction was found to be far stronger than its female reproduction when browsed (Krupnick and Weis, 1999). This is because ovules are packaged in the ovary and are not easily influenced by the environment (Knight et al., 2005), while pollen directly emerges from the anthers and can be browsed by herbivores. Therefore, a decrease in the number of pollen grains, decreased investment in male reproduction and in the probability that ovule fertilization would increase the fitness of the host plant (Miao, 2016).

Threshold grazing intensity may exist because low grazing intensity has no significant effect on reproductive performance (Lee and Bazzaz, 1980; Mueller et al., 2005). Moreover, pollen quality is affected by herbivory damage (Strauss, 1997). Our results also showed that pollen from flowers on heavily browsed plants set fewer pods and ripe seeds compared with pollen from unbrowsed plants (Table 3), and different grazing intensities lead to different pollen performance (Fig. 1). Grazing and the type of pollination and their interaction significantly influenced reproductive success (N’Guessan and Hartnett, 2011). On the other hand, our study found that the proportion of fertilized ovules may not differ depending on the source of pollen from plants browsed at different intensities, and that ovules fertilized by pollen from undamaged plants were more likely to mature. Rosas et al. (2011) suggested that pollen quality is the most likely factor that affects the probability of siring fruits and seeds, such as differences in the speed of germination and/or growth rates of pollen tubes. Moreover, pollen quality could affect the proportion of ovules fertilized under different treatments. At the same time, self-incompatibility in plants may lead to different results for different pollination treatments. That is, autogamy would not usually sire pods and seeds (Xu et al., 1988; Thrall et al., 2014).

In conclusion, biomass allocation to reproductive organs was reduced with increasing grazing intensity. Grazing significantly reduced the number of pollen grains of single flowers, the total number of flowers, riped seeds and the rate of pod-set, and seed mass. Pollen grains from heavily grazed plants that were donated to conspecifics seem to be less likely to set mature seeds if they must compete against pollen grains from non-grazed plants for ovules access.

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References
Brody, A. (1997). Effects of pollinators, herbivores, and seed predators on flowering phenology. Ecology 78, pp. 1624–1631.
Brookshire, E.N.J., Kauffman, J.B., Lytjen, D., Otting, N. (2002). Cumulative effects of wild ungulate and livestock herbivory on riparian willows. Oecologia 132, pp. 559–566.
Buchanan, A.L., Underwood, N. (2013). Attracting pollinators and avoiding herbivores: insects influence plant traits within and across years. Oecologia 173, 473–482.
Cong, X.L., Yang, J.Y., Li, X.M. (2017). Genetic diversity analysis of Caragana Micnphylla LAM. and Caranaga Stenophylla Poojar in inner mongola plateau. Chinese Journal of Grassland 7, pp. 85–92.
Du Tott, J.T., Olff, H. (2014). Generalities in grazing and grazing ecology: using across-guild comparisons to control contingencies. Oecologia 174, 1075–1083.
Feller, I.C. (1996). Effects of nutrient enrichment on leaf anatomy of dwarf Rhizophora mangle L. (Red mangrove). Biotropica 28, 13–22.
Francois, P. (2013). Early physiological responses of Abies alba and Rubus fruticosus to ungulate herbivory. Plant Ecology 214, 127–138.
González, M.A. (2016). Within-and trans-generational effects of herbivores and detritivores on plant performance and reproduction. *Journal of Animal Ecology 85*, 283–290.

Gowda, J.H., Palo, R.T. (2003). Age-related changes in defensive traits of *Acacia tortilis* Hayne. *Africa Journal of Ecology 41*, 218–223.

He, M., Hu, J.H. (2010). Influence of grazing on seed production of *Caragana microphylla*. *Chinese Journal of Plant Science 45*, 59–65.

Juenger, T., Bergelson, J. (1997). Pollen and resource limitation of compensation to herbivory in scarlet gilia, *Ipomopsis aggregata*. *Ecology 78*, 1684–1695.

Juenger, T., Bergelson, J. (2000). Does early season grazing influence the effect of self-pollination in scarlet gilia? *Ecology 81*, 41–48.

Kaitaniemi, P., Neuvonen, S., Nyystyssonen, T. (1999). Effects of cumulative defoliations on growth, reproduction, and insect resistance in mountain birch. *Ecology 80*, 524–532.

Karban, R., Strauss, S. (1993). Effects of herbivores on growth and reproduction of their perennial host, *Erigeron glaucus*. *Ecology 74*, 39–46.

Knight, T.F., Steets, J.A., Vamosi, J.C., Mazer, S.J., Burd, M., Campbell, D.R., Dudash, M.R., Johnston, M.O., Mitchell, R.J., Ashman, T.L. (2005). Pollen limitation of plant reproduction: pattern and process. *Annual Review of Ecology, Evolution, and Systematics 36*, 467–497.

Krupnick, G.A., Weis, A.E. (1999). The effect of floral herbivory on male and female reproductives success in *Isomeris arborea*. *Ecology 80*, 135–149.

Leather, S.R. (2000). Herbivory, phenology, morphology and the expression of sex in trees: who is in the driver’s seat? *Oikos 90*, 194–196.

Lee, T.D., Bazzaz, F.A. (1980). Effects of defoliation and competition on growth reproduction in the annual plant *Abutilon theophrasti*. *Ecology 68*, 813–821.

Li, Y.H., Chen, Z.Z., Wang, S.P., Huang, D.H. (1999). Grazing experiment for sustainable management of grassland ecosystem of Inner Mongolia steppe: experimental design and the effects of stocking rates on grassland production and animal live weight. *Acta Agricultural Sinica 7*, 173–182 (in Chinese with English abstract).

Liu, L.D., Zhang, L., Gao, Y.B. (2004). *Caragana microphylla* Lam. *Bulletin Biology 39*, 21–22 (in Chinese).

Maleck, K., Dietrich, R.A. (1999). Defense on multiple fronts: how do plants cope with divers enemies? *Trends in Plant Science 4*, 215–219.

Maschiinski, J. (2001). Impacts of ungulate herbivores on a rare willow at the southern edge of its range. *Biological Conservation 101*, 119–130.

Massad, T.J. (2013). Ontogenetic differences of herbivory on woody and herbaceous plants: a meta-analysis demonstrating unique effects of herbivory on the young and the old, the slow and the fast. *Oecologia 172*, 1–10.

Miao, F.Z., Xue, W., Guo, Z.G. (2016). Response of niche characteristics of alpine meadow plant populations on the northeastern margin of the Tibetan Plateau to grazing grazing. *Acta Pratacultura Sinica 25*, 88–97.

Milchunas, D.G., Sternberg, M. (2011). Plant community dynamics in shortgrass steppe with grazing relaxation and imposition by large and small herbivores. *Israel Journal of Ecology & Evolution 57*, 23–41.

Mothershead, K., Marquis, R.J. (2000). Fitness impacts of herbivory through indirect effects on plant-pollinator interactions in *Oenothera macrocarpa*. *Ecology 81*, 30–40.

Mueller, R.C., Wade, B.D., Gehring, A.A., Whitham, T.G. (2005). Chronic herbivory negatively impacts cone and seed production, seed quality and seedling growth of susceptible pinyon pines. *Oecologia 143*, 558–565.

Mundim, F.M., Bruna, E.M., Vieira-Neto, E.H.M., Vasconcelos, H.L. (2012). Attack frequency and the tolerance to herbivory of Neotropical savanna trees. *Oecologia 168*, 405–414.

Musser, R.O., Hum-Musser, S.M., Lee, H.K., DesRochers, B.L., Williams, S.A., Vogel, H. (2012). Caterpillar labial saliva alters tomato plant gene expression. *Journal of Chemical Ecology 38*, 1387–1401.

Nagaike, T. (2012). Effects of grazing by sika deer (*Cervus nippon*) on subalpine vegetation at Mt. Kita, central Japan. *Ecological Research 27*, 467–473.

N’Guessan, M., Hartnett, D.C. (2011). Differential responses to defoliation frequency in little bluestem (*Schizachyrium scoparium*) in tallgrass prairie: implications for herbivory tolerance and avoidance. *Plant Ecology 212*, 1275–1285.

Noy-Meir, I. (1993). Compensating growth of grazed plants and its relevance to the use of rangelands. *Ecological Applications 3*, 32–34.

Pollice, J., Marcora, P., Renison, D. (2013). Seed production in *Polyplepis aurialis* (Rosaceae) as influenced by tree size, live stock and interannual climate variations in the mountains of central Argentina. *New Forests 44*, 233–247.

Rebek, K.A., O’Neil, R.J. (2005). Impacts of simulated herbivory on *Alliaria petiolata* survival, growth, and reproduction. *Biolcontrol 34*, 283–289.

Rosas, F., Quesada, M., Lobo, J.A., et al. (2011). Effects of habitat fragmentation on pollen flow and genetic diversity of the endangered tropical tree *Swietenia humilis* (Melianaceae). *Biological Conservation 140*, 3082–3088.

Ruhomaki, K., Haukoja, E., Repka, S., Lehtila, K. (1997). Leaf value: effects of damage to individual leaves on growth reproduction of mountain birch shoots. *Ecology 78*, 2105–2117.

Rusch, G.M., Skarpe, C., Halley, D.J. (2009). Plant traits link hypothesis about resource-use and response to herbivory. *Basic & Applied Ecology 10*, 466–474.

Schemske, D.W. (1988). Plant-animal interactions and fruit production in a neotropical herb: a path analysis. *Ecology 69*, 1128–1137.

Sell, A.F. (2000). Morphological defenses induced in situ by the invertebrate predator *Chaoborus*: comparison of responses between *Daphnia pules* and *D. rosea*. *Oecologia 125*, 150–160.

Simms, E.L., Rausher, M.D. (1987). Costs and benefits of plant resistance to herbivory. *American Naturalist 130*, 570–581.

Solomon, B.P. (1986). Sexual allocation and andromonoecy: resource investment in male and hermaphrodite flowers of *Solanum carolinense* (Solanaceae). *American Journal of Botany 73*, 1215–1221.

Stephenson, A.G. (1982). The role of the extrafloral nectaries of *Catapla speciosa* in limiting herbivory and increasing fruit production. *Ecology 63*, 663–668.

Strauss, S. (1997). Floral characters link herbivores, pollinators, and plant fitness. *Ecology 78*, 1640–1645.

Thrall, P.H., Encinasviso, F., Hoeebee, S.E., et al. (2014). Life history mediates mate limitation and population viability in self-incompatible plant species. *Ecology & Evolution 4*, 673–687.

van Kleunen, M., Ramponi, G., Schmid, B. (2004), Effects of herbivory simulated by clipping and jasmonic acid on *Solidago Canadensis*. *Basic and Applied Ecology 5*, 173–181.

Vazquez, D.P., Simberloff, D. (2004). Indirect effects of an introduced ungulate on pollination and plant reproduction. *Ecological Research 19*, 288–299.

Wang, L.H., Liu, W., Wang, J.N., et al. (2015). Compensatory growth of grassland communities, stratification and species under different cutting strengths. *Acta Pratacultura Sinica 24*, 35–42.

Wang, S.P., Li, Y.H., Wang, Y.F., Chen, Z.Z. (2001). Influence of different cutting strengths on plant community in Inner Mongolia steppe. *Acta Agricultural Sinica 30*, 89–96 (in Chinese with English abstract).

Wang, S.P., Li, Y.H., Wang, Y.F., Han, Y.H. (1998). The succession of *Artemisia frigida* rangeland and multivariation analysis


under different stocking rates in Inner Mongolia. *Acta Agriculturae Sinica* 6, 299–305 (in Chinese with English abstract).

Xu, L.R., Hao, X.Y., Yang, X.L., Li, Y.J. (1988). On the hybridization and classification of *Caragana korshinskii* Kom. and *C. intermedia* Kung et H. C. Fu. Abstracts of the papers presented at the 55th Anniversary of the Botanical Society of China, pp. 188.

Zhang, Z., Wang, S.P., Patton, B., Nyren, P., Jiang, G.M. (2007). Responses of *Artemisia frigida* and *Leymus chinensis* to sheep saliva. *Journal of Arid Environment* 70, 111–119.