Six-year removal of co-dominant grasses alleviated competitive pressure on subdominant grasses but dominant shrub removal had neutral effects in a subalpine ecosystem

Wenjin Li, Johannes M.H. Knops, G. Kenny Png, Xi Yan, Huan Dong, Jinhua Li, Huakun Zhou, Rubén Díaz Sierra

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1	Six-year removal of co-dominant grasses alleviated competitive pressure on		
2	subdominant grasses but dominant shrub removal had neutral effects in a		
3	subalpine ecosystem		
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5	Wenjin Li ¹ *, Johannes M. H. Knops ² , G. Kenny Png ^{3,4} ,		
6	Xi Yan ¹ , Huan Dong ¹ , Jinhua Li ¹ , Huakun Zhou ⁵ , Rubén Díaz Sierra ⁶		
7 8	1.State Key Laboratory of Grassland Agro-ecosystems, School of Life Sciences,		
9	Lanzhou University, Lanzhou 730000, China		
10	2. Health and Environmental Sciences, Xi'an Jiaotong-Liverpool University,		
11	Suzhou 215123, China		
12	3. Soil and Ecosystem Ecology Laboratory, Department of Earth and		
13	Environmental Sciences, The University of Manchester, Manchester M13 9PT,		
14	United Kingdom		
15	4. Asian School of the Environment, Nanyang Technological University, 50		
16	Nanyang Ave, Singapore 639798, Singapore		
17	5. Qinghai Province Key Laboratory of Restoration Ecology of Cold Area, the		
18	Northwest Institute of Plateau Biology, Chinese Academy of Sciences, Xining,		
19	810008, Qinghai, China		
20	6.Mathematical and Fluid Physics Department, Faculty of Sciences,		
21	Universidad Nacional de Educacion a Distancia (UNED), Madrid 28040,		
22	Spain		
23			
24	*Corresponding author: Wenjin Li		
25	E-mail addresses: liwj@lzu.edu.cn or wenjinli@163.com		
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29			

30	Abstract
31	The 'stress-gradient hypothesis' predicts increasing facilitative interactions with
32	increasing environmental stress, but it remains unclear if the prevailing interaction (i.e.
33	facilitative or competitive) of dominant plant species on subordinate plant species
34	occurring in harsh environments is dependent on the plant functional type. In addition,
35	most plant-species removal experiments in grasslands are short-term (1-2 years),
36	which may imprecisely reflect transient effects arising from methodological
37	limitations. We conducted a dominant species removal experiment in a subalpine
38	ecosystem, containing a mosaic of grass-dominated and shrub-dominated community
39	patches, both of which are common in the subalpine zone of the Qinghai-Tibetan
40	Plateau. We examined the direction and magnitude of the effects of three co-dominant
41	grass and a dominant shrub species on subordinate species richness and biomass over
42	a 6-year period. Removal of the dominant grass species alleviated their competitive
43	pressure on subdominant grasses, which resulted in similar total and grass biomass
44	detected in the final year of the study. By contrast, shrub removal showed no effects
45	on its subordinate species biomass. Furthermore, neither the removal of the dominant
46	shrubs nor the grasses altered their respective subordinate species richness. Thus, in
47	subalpine ecosystems that experience harsh environmental conditions, our results
48	showed that the direction of interactive effects of dominant plant species on
49	subordinate species may be dependent on the plant functional type and are not
50	necessarily facilitative. Furthermore, we showed that longer-term plant-removal
51	experiment observations may be required to better determine the effects of species
52	removal for subalpine and montane ecosystem(s).
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54	
55	Keywords: stress-gradient hypothesis; removal experiments; plant functional type;
56	competition and facilitation; grassland; Qinghai-Tibetan Plateau;
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58 1. Introduction

59	Environmental changes such as drought, warming, nitrogen deposition or land
60	management practices (e.g. fire-suppression and grazing intensity changes) are
61	increasingly common in grassland ecosystems globally (e.g. Shaw et al. 2002; Menge
62	and Field 2007; Wardle et al. 2011). These changes can strongly influence plant-plant
63	interactions that can, in turn, lead to detrimental shifts in the vegetation community
64	composition (Eldridge et al. 2011; Loranty and Goetz 2012; Naito and Cairns 2011;
65	Ratajczak et al. 2012). Grassland vegetation communities are typically dominated by
66	a few plant species, typically grasses (Grime 1998; Gibson 2009), which are considered
67	to be superior competitors compared with other co-occurring forb species (Collins and
68	Glenn 1990; Miles and Knops 2009). As such, dominant grass species are expected to
69	drive plant community dynamics via competitive interactions (in benign
70	environmental conditions) (e.g., Bertness & Callaway 1994; Grime 1998; Smith and
71	Knapp 2003; Smith et al. 2020). However, it is unclear if competitive or facilitative
72	interactions prevail in driving vegetation dynamics in ecosystems that contain
73	dominant plants from different plant functional groups and typically experience
74	stressful environmental conditions. This is because contrasting plant
75	functional/physiological trait(s) may confer different types of plant-plant interaction
76	with respect to the environmental stress factor(s) the community experiences, and
77	field experiments examining interactions in stressful environments have produced
78	inconsistent findings (Olofsson et al. 1999; Pugnaire et al. 2015; Michalet et al. 2015).
79	The relatively short duration of many of these studies may further confound the
80	findings due to the possible detection of 'transient' effects (e.g. unintentional
81	short-term effects arising from possible methodological limitations associated with
82	plant removals or inter-annual environmental variability) (He and Bertness 2014;
83	Pugnaire et al. 2015). Thus, it is valuable to accurately determine the predominant
84	type of plant-plant interaction occurring in grasslands experiencing stressful abiotic
85	environments over a longer period of time, which can in turn, inform their
86	management and restoration.
87	The expected net effect of a dominant species on neighboring plants is likely to be
88	influenced by their respective functional/physiological trait(s) with respect to the
89	environmental stress factor(s) the community experiences (Schöb et al. 2013;
90	Soliveres 2014; Soliveres et al. 2015). For example, in dry alpine regions, net

91	facilitation on neighboring plants is expected to be detected for drought-tolerant
92	shrubs whose canopy-form (e.g. shelter from wind, high ultraviolet irradiation, and/or
93	improve organic matter accumulation; Padilla and Pugnaire 2006) and resource
94	acquisition strategies (e.g. deeper roots that help transport water from lower to upper
95	soil layers via the hydraulic lift process; Horton and Hart 1998; Prieto et al. 2011)
96	benefit neighboring understory plant species more than the detriments arising from the
97	shrubs' competitive abilities. Conversely, in arid grasslands, a hypothetical dominant
98	grass species that is relatively fast-growing, shallow-rooted and less water-use
99	efficient is expected to be highly competitive (e.g. spatial, water, and light
100	availability). However, facilitative effects have been hypothesized to increase with
101	increasing environmental stress (e.g. Callaway et al. 2002; Maestre et al. 2009), with
102	many studies supporting the 'stress-gradient hypothesis' (Aksenova et al. 1998; Lortie
103	and Callaway. 2006). On the other hand, there are also studies showing the
104	importance of competitive interactions in harsh conditions (e.g. del Moral 1983;
105	Moral 1985; Olofsson et al. 1999). As such, in ecosystems containing several
106	dominant plant species from contrasting plant functional groups, more experimental
107	tests are required to examine if facilitative effects generally prevail under harsh
108	environmental conditions as hypothesized by the stress-gradient hypothesis (Bertness
109	& Callaway 1994).
110	Plant-plant interactions are frequently examined by comparing the performance of a
111	target plant with and without a neighboring plant (Aarssen and Epp 1990; Diaz-Sierra
112	et al. 2017; Diaz et al. 2003). At present, there is an increasing trend towards
113	examining the interactions between a target plant species and its community using a
114	similar plant removal approach (Michalet et al. 2015) to determine how the target
115	species influence co-occurring species, plant diversity (Brooker 2006; Xu et al. 2010),
116	community dynamics, and ecosystem functioning (Grime 1998; Mariotte 2014;
117	Huang et al. 2020). However, the use of species removal experiments to determine
118	how plant species influence/interact with its community in cold and stressful
119	environments can be difficult. This is because ecosystem processes (e.g. soil microbial
120	activity, decomposition, nutrient cycling, plant growth rates) in cold and stressful
121	environments tend to be relatively slow (e.g. Atkin et al. 1996; Körner, 2003;
122	Bradford et al. 2016; Wang et al. 2016), and many experimental removal studies were
123	conducted over a relatively short period of time (1-2 years) (e.g. Smith and Knapp
124	2003; Xu et al. 2010; Michalet et al. 2015). Thus, it is unclear if previous results

125	reflected transient effects (e.g., short-term local microclimatic disturbances, soil
126	legacy effects from slowly decomposing roots of clipped plants, and/or climatic
127	fluctuations between years) or indicated the definite interactive effect(s) of the
128	removed species (Kardol et al. 2018; Lepš 2014). Furthermore, plant removal studies
129	that examined plant interactions over sufficiently long time periods are rare, and may
130	be essential to be able to detect interactive effects accurately (Adler et al. 2010; Lepš
131	2014).
132	A subalpine ecosystem consisting of a mosaic of vegetation patches dominated by
133	either the shrub Potentilla fruticosa or co-dominated by three grasses Roegneria
134	nutans, Elymus nutans and Leymus secalinus is common within the subalpine zone of
135	the Qinghai-Tibetan Plateau (Michalet et al. 2015). A previous study in this study area
136	showed that the removal of a dominant shrub species led to a decrease in the biomass
137	of subdominant grass species, but no net effect on forbs (Xu et al. 2010), while a
138	separate study at the same site showed that the removal of dominant grass species had
139	no net effect on the species richness of forbs (Michalet et al. 2015). However,
140	considering observations by both these studies were done over a relatively short
141	period of time (Xu et al. 2010; Michalet et al. 2015), it remains unclear if the plant
142	community responses (to consistent shoot removal of target species) detected in these
143	studies are indeed due to the removed species. This is because clonal regeneration of
144	clipped species from unremoved belowground plant organs (Qian et al. 2017; Ott et al
145	2019) or transient effects may persist (Kardol et al. 2018) as a result of relatively slow
146	ecosystem processes in cold environments (Atkin et al. 1996; Körner, 2003; Germino
147	2014; Bradford et al. 2016; Wang et al. 2016).
148	To better determine if the net effect by dominant species on the subordinate
149	herbaceous species is dependent on the contrasting plant functional traits of the
150	dominant species in a cold and harsh environment, we conducted a longer-term
151	removal experiment over 6 growing seasons within a subalpine ecosystem. Our study
152	site is a subalpine meadow grass-shrubland ecosystem, located at an altitude of 2900m
153	consisting of a mosaic of meadow grass- and shrub-dominated patches. We
154	hypothesized that: (1) the plant functional type of the dominant species is important to
155	consider under stressful environmental conditions, because the dominant grass species
156	are expected to be superior competitors (Collins and Glenn 1990; Miles and Knops
157	2009), and the removal of the dominant grasses, due to competitive release, will lead
158	to an increase in the richness and biomass of herbaceous subordinate species at the

159	end of the study; while (2) the dominant shrub <i>Potentilla fruticosa</i> is expected to
160	function as a facilitative 'nurse' species, because its shrub growth form may alter the
161	microclimate of the herbaceous understory layer to facilitate co-occurring herbaceous
162	understory species in this climatically harsh subalpine environment, and shrub
163	removal may lead to decreases in both the herbaceous understory richness and
164	biomass. Furthermore, (3) the net direction and magnitude of interaction measured
165	between the first and final year of the study will not be consistent due to transient
166	effects arising from the possible methodological limitations associated with plant
167	(aboveground) removals.
168	
169	2. Materials and methods
170	2.1 Study site
171	This experiment was conducted at the Alpine Meadow and Wetland Ecosystem
172	Research Station of Lanzhou University (34°44′N, 102°53′E), located on the eastern
173	Qinghai-Tibetan Plateau in Hezuo, China, at an altitude of 2900 m. The study site has
174	been fenced since 2007 and is used as winter grazing pastures. The site has an annual
175	grazing intensity of 8-10 Tibetan sheep per ha for 20 days in the winter, and such
176	grazing regime is typical for the regional area (Michalet et al. 2015). The growing
177	season is short with 54 frost-free days annually, and temperature varies from -8.3 $^{\circ}$ C in
178	December and 11.9°C in July (Institute of Hezuo Meteorology, China). Mean annual
179	precipitation is 558 mm with 85% of the precipitation occurring during the growing
180	season and with an annual potential evapotranspiration of 1222 mm (Institute of
181	Hezuo Meteorology, China).
182	The soil type is classified as subalpine meadow soil according to the Chinese soil
183	classification system (Gong 1999), with a mean soil organic carbon content of 34.9 \pm
184	$0.6~g~kg^{-1}$ dry soil and soil total nitrogen of $3.3\pm0.8~g~kg^{-1}$ dry soil. The mean soil pH
185	is 7.2 ± 0.1 .
186	The study site has a subalpine ecosystem consisting of a mosaic of two subalpine
187	vegetation patch types: subalpine meadow grass and shrub patches. The subalpine
188	meadow grass patch typically contains three co-dominant grass species, Roegneria
189	nutans, Elymus nutans, and Leymus secalinus, which combine to contribute 45 \pm 5 %
190	to the total plant cover (Li et al. 2018). The shrub patch is dominated by the only
191	shrub species occurring at our study site, <i>Potentilla fruticosa</i> , which accounts for 59 \pm
192	11 % of the total plant cover (Xu et al. 2010).

2.2 Experimental design and plot selection

Twenty-four 50 cm \times 50 cm plots were established in total within the study site containing a mosaic of subalpine grass- and shrub- dominant meadow patches in 2012. Twelve plots were selected in patches with the dominant shrub present (hereafter also referred to as "shrub plots") to test for the effects of the dominant shrub species on co-occurring subordinate plant species, while another 12 plots were selected within patches containing all 3 co-dominant grass species (hereafter also referred to as "grass plots") to test their effects on the subordinate species. All plots were also selected to contain other plant species from the four plant functional groups of legumes, grasses, sedges and forbs. Plots were permanently marked, and all plots were at least 1.5 m apart and situated within a 1-ha area. Each removal treatment was randomly assigned to 6 of the 12 plots for each vegetation patch-type, with the remaining 6 plots as control. For both the grass and shrub removal plots, we clipped the 3 co-dominant grasses or the shrub to ground level in early June using a similar approach by Michalet et al. (2015) and Elumeeva et al. (2017). Briefly, all removal treatments were repeated every two weeks between early June and late July each year.

2.3 Vegetation sampling

Species richness of all vascular plant for each $50~\rm cm \times 50~\rm cm$ plot was recorded at the peak of every growing season in early-August (prior to winter grazing). Each plot area was subdivided into four equal sized subplots. Aboveground biomass was estimated by clipping a $25~\rm cm \times 25~\rm cm$ area in one of the four subplots to ground level each year, in early-August, from 2012 to 2017, except for 2016. The clipped vegetation from each plot was further sorted into legumes, grasses, sedges or forbs, and then weighed to obtain biomass data for each plant functional group. All clipped samples were dried for 48 hours at 75 °C prior to weighing.

We choose to keep plots relatively small to capture the proximate area that the dominant shrub and grasses are likely to influence their understory. Although we clipped a relatively large proportion within each plot to estimate the plant biomass of each plot, we do not think that this impacted our results. This is because the site is grazed by sheep during winter, and the clipped and unclipped areas of our plots

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2.4 Statistical analyses

For each patch type, we assessed the effects of dominant species removal 229 treatments (i.e. removal, non-removal), and the possible interaction with temporal 230 effects (i.e. inter-annual environmental variability), on the following response 231 232 variables: (i) total aboveground biomass and (ii) total species richness of the herbaceous layer, and the respective herbaceous plant functional groups' aboveground 233 234 biomass [(iii) grass, (iv) sedge, (v) legume, and (vi) forb biomass] and species richness [(vii) grass (dominant + subordinate species), (viii) subordinate grass, (ix) 235 sedge, (x) legume, and (xi) forb species richness]. We used a repeated measures 236 analysis of variance (ANOVA) test ($\alpha = 0.05$) to analyze each response variable, and 237 performed post-hoc Tukey HSD tests ($\alpha = 0.05$) if significant differences were 238 detected. For each statistical analysis, we visually assessed data for normality and 239 heterogeneity of variance, and applied appropriate transformations when necessary. 240 All statistical analyses were performed in SPSS 20.0 (IBM Corp., New York, USA). 241 Shrub data were not included in the calculations or analyses for two reasons: (1) 242 243 the community responses of the herbaceous layer were of interest in our study; and, (2) comparisons involving the dominant shrub and herbaceous plants are partial because 244 245 many herbaceous plants are generally smaller (relative to P. fruticosa) and are limited spatially within the herbaceous layer stratum, whereas the dominant shrub is less 246 247 palatable to large herbivore grazers (Elkington & Woodell 1963) and is intrinsically taller/larger. That is, the shrubs' upper canopy position, woody growth and less 248 grazing allows the accumulation of a large proportion of its woody biomass in the 249 vertical shrub layer stratum over a longer period of time. 250

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3. Results

3.1 Co-dominant grass species removal

Removal of co-dominant grasses significantly reduced grass (dominant + subordinate species) biomass (~21% of total) and total aboveground biomass in the initial four sampling years, but not in the final year (Fig. 1A; Table 1). By contrast, the removal of the co-dominant grasses had little or no effect on the biomass of legumes and sedges (Fig. 1A; Table 1). In addition, within each given year, there was no significant difference in the biomass of forbs between the removal treatments (Fig. 1A; Table 1).

Forbs comprised between 67-83% of the total species richness (6-26 total species
per plot), and this functional group represented the largest proportion of plant species
in the grass patches of this subalpine ecosystem (Fig. 2A; Table 1). Within each given
year, the removal of co-dominant grasses had little or no effect on the total, legume or
forb species richness between the removal treatments (Fig. 2A; Table 1). Further, the
subordinate grass species richness was generally similar between removal treatments
in most years, except in 2015 (Fig. 2B; Table 1).

3.2 Dominant shrub species removal

Only forb biomass varied significantly among years, but the total plot biomass and the biomass of each plant functional group were, respectively, similar between the shrub removal treatments within each given year (Fig. 1B; Table 1).

The total species richness of each shrub plot ranged between 6 to 26, and the forb functional group also represented the largest proportion of plant species in the shrub patches (ranging between 50-87%) of this subalpine ecosystem (Fig. 2C; Table 1). The total biomass and the biomass of grass and forbs varied significantly among years, but not the biomass of sedges or legumes (Fig. 2C; Table 1). However, within each given year, total species richness and the species richness of each plant functional group were respectively similar between both shrub removal treatments (Fig. 2C; Table 1).

4. Discussion

In this study, we found some indication of competitive effects by co-dominant grasses as hypothesized. However, the competitive effects of the co-dominant grasses were limited only to the subordinate grass species (i.e. marginal aboveground biomass compensation by subordinate grass species). This suggests that plant functional type is important to consider, and that facilitative effects are not necessarily predominant under harsh abiotic environments (e.g. del Moral 1983; Moral 1985; Olofsson et al. 1999). In addition, these competitive effects were only detected in the 6th and final year of the study, which could support our hypothesis that observations of transient effects may diminish over a longer term and suggest that observations over a longer period may be required to verify effects. Contrary to our hypothesis, the dominant shrub species did not show any impact on the herbaceous understory layer (i.e. neutral effects). Findings from our longer-term study were in contrast with two shorter-term

294	studies in the same study area that detected neutral (i.e. Michalet et al. 2015) or
295	facilitative effects (Xu et al. 2010), and studies elsewhere detecting facilitative effects
296	in ecosystems experiencing harsh abiotic environments (e.g. Callaway et al. 2002;
297	Soliveres et al. 2015). Furthermore, the absence of a strong response from subordinate
298	species to dominant grass or shrub species removal, in terms of species richness, may
299	also suggest that factors other than plant-plant interactions could be important drivers
300	of plant community structure in this subalpine ecosystem (Mitchell et al. 2009).
301	Together, our results showed that the plant functional types of the dominant plant
302	species as well as transient effects are important to consider to some extent. However,
303	other factors such as grazing intensity, growing season length, rainfall and
304	temperature may also be important at influencing plant community dynamics in this
305	subalpine and other montane ecosystem(s) (Mitchell et al. 2009). Thus, we
306	recommend future studies to explore the importance of these factors (and their
307	interactions) in combination with species removal in montane environments.
308	
309	4.1 Biomass responses of the herbaceous layer
310	We found that the competitive effects of the co-dominant grass species were limited to
311	subordinate species that were functionally similar (i.e. grasses) and likely to occupy
312	the same niche (McKane et al. 2002). This is because the removal of co-dominant
313	grass species showed marginal biomass compensation by subordinate grass species in
314	the 6 th year, but had little or no effect on both the biomass and plant species richness
315	of all other plant functional groups examined. In addition, the similar overall
316	aboveground biomass between plot removal treatments in the final year may also
317	suggest that the subordinate grass species may play a functionally equivalent role as
318	the co-dominant grass species to maintain the ecosystem function of biomass
319	production (Smith and Knapp 2003). Thus, highlighting the importance of functional
320	redundancy of less dominant species in ecosystem functionality (Fetzer et al. 2015)
321	and biodiversity conservation (Wohl et al. 2004). Nonetheless, because biomass
322	compensation was only observed in the 6 th and final year, we recommend that future
323	studies should examine removal effects over a longer period of time to validate that
324	observations were not due to transient effects associated with inter-annual
325	environmental variability.
326	On the other hand, shrub removals had little or no net effect on the total and

respective plant functional group biomass of the herbaceous layer in our study, which

328	is in contrast to a number of other studies that show either a nurse/facilitative or
329	competitive effects on understory species in ecosystems exposed to harsh climatic
330	effects (e.g. Bråthen and Lortie 2016; Pajunen et al. 2011; Xu et al. 2010). However,
331	our results are in line with a six-year shrub-removal experiment in the Alaskan
332	tussock tundra (Bret - Harte et al. 2008). We speculate that many herbaceous
333	understory species in the shrub plots, as well as in our study system, are well-adapted
334	to the harsh conditions of the subalpine environment (Sun et al. 2014). Moreover,
335	resource and vertical spatial partitioning may also occur among the shrub and other
336	understory species (McKane et al. 2002; Hu et al. 2013). Thus, based on our results,
337	the shrub <i>P. fruticosa</i> appeared to have little or no influence on herbaceous forage
338	vegetation production in our study system. Nevertheless, because P. fruticosa has a
339	widespread distribution in the northern hemisphere, future studies should examine
340	how the effects of climate (e.g. climate warming) may promote shrub expansion into
341	higher altitude ecosystems (Myers-Smith et al. 2011; Eldridge et al. 2011), which in
342	turn has negative impacts on forage production (Pajunen et al. 2011).
343	
344	4.2. Transient belowground effects of remnant roots
345	The initially $(2012 - 2015)$ lower total aboveground and grass biomass in the
346	co-dominant grass removal plots followed by a shift to similar levels of biomass of
347	the control plots in the final year could indicate transient effects. The transient effects
348	could be due to the short-term recurrence of plant clonal regeneration from remnant
349	belowground organs (Ott et al. 2019) and/or generation of soil legacies through
350	decomposition of the remnant roots of the clipped grasses (Kardol et al. 2018; Lepš
351	2014). For example, the unremoved remnant roots could continue to allow dominant
352	grass species to exert their effects on the community via spatial exclusion,
353	maintenance of interactions with subordinate species via clones (Ott et al. 2019),
354	and/or suppressing establishment of new species (Kardol et al. 2018; Lepš 2014).
355	Although the lack of biomass compensation by subordinate grasses during the initial
356	years could also be, in part, explained by the slower, conservative growth traits of
357	subalpine plants that may constrain their growth responses to the increase in available
358	resources (Bret-Harte et al. 2004), we consider this to be less likely. This is because
359	the dead standing aboveground biomass vegetation is entirely consumed by sheep
360	during the winter (i.e. unclipped areas of non-removal plots visually similar to clipped

subplots), but the vegetation appeared to recover to similar functional proportions in the following season.

However, we did not measure root biomass nor root competition in this study, and are unable to ascertain if the subordinate species may have been suppressed by the negative effects from the remnant roots of dominant plant species, or if non-detection of effects were due to conservative growth traits of subalpine plants (Atkin et al. 1996; Körner, 2003; Germino 2014). Moreover, plant removal studies in this subalpine study system are limited and previous shorter-term experiments have yielded inconsistent results (Xu et al. 2010; Michalet et al. 2015). Thus, little is known about the intensity and relative importance of belowground interactions in this subalpine meadow ecosystem. Therefore, future studies should involve more greenhouse or mesocosm studies that allow the examination of both aboveground and belowground plant interactions.

4.3 Seed propagation strategies and grazing intensity may explain the lack of species richness responses

The lack of responses in terms of subordinate species richness, particularly forbs (Li et al 2019; Siebert et al.2019), could be due to the seed propagation strategy of many plant species in this subalpine meadow ecosystem. Many non-graminoid species within this subalpine vegetation mosaic are long-lived, slow-growing perennials that likely rely on seed reproduction (*cf.* clonal reproduction) (Weppler et al. 2006). This is because seeds provide the advantage of being able to remain dormant and persist during extended periods of unfavorable environmental conditions or climatic variability common to montane ecosystems (Bewley 1997; Baskin and Baskin 2014). As such, changes in species richness after plant removals may only be detected when new individuals are episodically recruited in response to (a series of) specific environmental cue(s) that occasionally occur (e.g. shorter cold growing seasons; Körner, 2003; Mondoni et al. 2012). As such, species richness responses of the plant community to dominant plant species removal in this study system may require longer-term observations (i.e. >6 years).

Conversely, the current grazing intensity of the study area – while considered typical (Michalet et al. 2015) – may also work to obscure the plant species richness response to dominant species removals. This is because hand-clipped and unclipped areas of our plots tended to appear similar post-winter grazing, which suggests that

395	the current level of sheep herbivory may inhibit seedling establishment of new plant		
396	species. However, because grazing only took place in our sites during winter, do note		
397	that the effects of herbivory on seedling establishment in our study is likely to affect		
398	seeds that germinate in fall, and not those in spring. Nevertheless, grazing is clearly an		
399	important factor, and should be considered in future plant-plant interaction studies		
400	(Graff et al. 2007; Le et al 2012).		
401			
402	5. Conclusions		
403	After six years of dominant grass species removal in a subalpine ecosystem, our		
404	results showed significant biomass compensation of subordinate grass species in		
405	response to dominant grass species removals only in the final (i.e. 6 th) year. In contrast		
406	shrub removals had little or no effect on the herbaceous layer, which indicated that the		
407	direction of interactive effects of dominant plant species on subordinate species may		
408	be dependent on the plant functional type. Furthermore, the competitive effects of		
409	dominant grass species detected in this subalpine ecosystem suggested that plant-plant		
410	interactions are not necessarily facilitative in cold, stressful environments. Taken		
411	together, our results highlight the importance of considering plant functional groups as		
412	well as conducting longer-term observations of plant-removal experiments to better		
413	understand plant-plant interactions in this subalpine and other montane ecosystem(s).		
414			
415	Author contributions WL conceptualized this study, led field surveys and wrote the		
416	first draft. JK, GKP, RS, LJ and HZ interpreted the results and revised the manuscript		
417	for readability. XY and HD collected and analyzed data. All authors contributed to		
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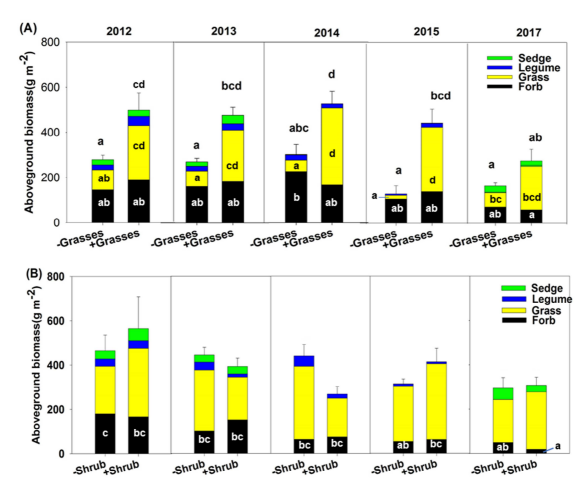
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Table 1. Test statistics of repeated measures analysis of variance tests on plant aboveground (dry) biomass and species richness of all species or respective plant functional groups of the herbaceous layer in response to co-dominant grass or dominant shrub species removals between 2012 to 2017. Biomass data were analyzed using ln(x+1) transformed values. F- and p-values in bold indicate significant differences ($p \le 0.05$; n=6). ns = not significant, p > 0.05, * p < 0.05, ** p < 0.01, ***

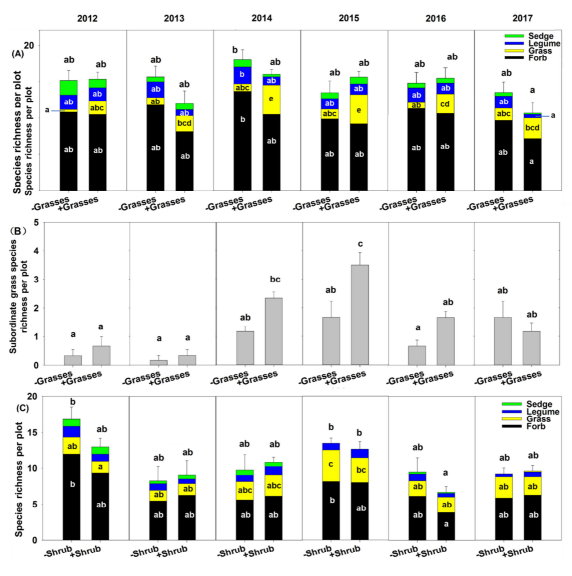
7 8 p < 0.001, / no data.

	Co-dominant grass species removal			Dominant shrub species removal		
Variables	Removal Treatment (T)	Year (Y)	Y*T	Removal Treatment (T)	Year (Y)	Y*T
D:	F-, p-values			F-, p-values		
<u>Biomass</u>						
Total biomass (g m ⁻²)	19.64***	4.68ns	16.62**	0.01ns	6.84*	0.01ns
Grass (dominant + subordinate) biomass (g m ⁻²)	21.80**	0.52ns	27.10**	0.01ns	0.01ns	0.12ns
Sedge biomass (g m ⁻²)	0.91ns	0.02ns	9.53*	0.002ns	0.85ns	0.79ns
Legume biomass (g m ⁻²)	0.03ns	8.43*	1.72ns	0.74ns	17.53**	0.11ns
Forb biomass (g m ⁻²)	0.18ns	16.84**	1.81ns	0.09ns	38.35***	0.48ns
Species richness						
Total species richness (plot ⁻¹)	0.06ns	0.25 ns	0.77ns	1.44ns	2.46ns	0.04ns
Grass (dominant + subordinate) species richness (plot ⁻¹)	11.49**	6.01*	0.79ns	0.10ns	21.28***	1.14ns
Subordinate grass species richness only (plot ⁻¹)	9.11*	18.76**	0.07ns	/	/	/
Sedge species richness (plot ⁻¹)	0.09ns	0.75ns	14.03**	0.08ns	36.80***	0.01ns
Legume species richness (plot ⁻¹)	3.67ns	7.90*	0.16ns	0.07ns	1.78ns	0.52ns
Forb species richness (plot ⁻¹)	2.05ns	2.43ns	0.04ns	0.95ns	4.35ns	0.04ns



Dominant species removal treatment

Figure 1. Effects of (A) co-dominant grass or (B) dominant shrub species removals on the aboveground (dry) biomass of the four plant functional groups of the herbaceous layer from 2012 to 2017, except for 2016. The grasses removed were *Roegneria nutans, Elymus nutans, Leymus secalinus*, and the shrub removed was *Potentilla fruticosa*. Bars represent means and error bars represent \pm 1 standard error. Different letters above each bar indicate significant differences among total aboveground biomass, while different letters within the respective plant functional group-sections indicate differences within the same functional group (*post-hoc* Tukey HSD test following significant repeated measures analysis of variance test on ln(x+1) transformed biomass values; $p \le 0.05$; Table 1); letters not shown when no significant differences was detected.



Dominant species removal treatment

Figure 2. Effects of (A) co-dominant grass species removals on the species richness of four different plant functional groups and (B) subordinate grass species richness, or the effects of the (C) dominant shrub species removals on the species richness of the herbaceous layer from 2012 to 2017. The grasses removed were *Roegneria nutans*, *Elymus nutans*, *Leymus secalinus*, and the shrub removed was *Potentilla fruticosa*. Bars represent means and error bars represent ± 1 standard error. Different letters above each bar indicate significant differences among plot total species richness, while different letters within the respective plant functional group-sections indicate differences within the same functional group (*post-hoc* Tukey HSD test following significant repeated measures analysis of variance test; $p \le 0.05$; Table 1); letters not shown when no significant differences was detected.

WL conceptualized this study, led field surveys and wrote the first draft. JK, PC, RS, LJ and HZ interpreted the results and revised the manuscript for readability. XY and HD collected and analyzed data. All authors contributed to this work and approved the final submission. The authors declared no competing interests.