1	Grazing <mark>increases</mark> litter decomposition <mark>rate</mark> but <mark>decreases</mark> nitrogen release <mark>rate</mark> in an
2	alpine meadow
3	
4	
5	Yi Sun ^{1,3} , Xiong Z. He ² , Fujiang Hou ¹ , Zhaofeng Wang ¹ , and Shenghua Chang ¹
6	
7	
8	
9	
10	¹ State Key Laboratory of Grassland Agro-ecosystems, Key Laboratory of Grassland Livestock Industry Innovation, Ministry
11	of Agriculture, College of Pastoral Agriculture Science and Technology, Lanzhou University, Lanzhou, 730000, Gansu, China
12	² School of Agriculture and Environment, College of Science, Massey University, Private Bag 11-222, Palmerston North, New
13	Zealand
14	³ State Key Laboratory of Cryospheric Sciences, Cold and Arid Regions Environmental and Engineering Research Institute,
15	Chinese Academy of Sciences, 320 Donggang West Road, Lanzhou 730000, China
16	Correspondence to: Fujiang Hou (cyhoufj@lzu.edu.cn)
17	

19	Abstract. Litter decomposition and N release are the key processes that strongly determine the nutrient cycling at the soil-
20	plant interface; however, how these processes are affected by grazing or grazing exclusion in the alpine grassland ecosystems
21	on the Qinghai-Tibetan Plateau (QTP) is poorly understood. So far few studies have simultaneously investigated the influence
22	of both litter quality and incubation site on litter decomposition and N release. Moreover, previous studies on the QTP
23	investigating how grazing exclusion influences plant abundance and biodiversity usually lasted for many years, and the short-
24	term effects have rarely been reported. This work studied the short-term (6 month) effects of grazing and grazing exclusion on
25	plant community composition (i.e., plant species presented) and litter quality, and long-term (27 – 33 month) effects on soil
26	chemical characteristics and mixed litter decomposition and N release on the QTP. Our results demonstrate that: (1) shorter
27	term grazing exclusion had no effect on plant community composition but increased plant palatability and total litter biomass;
28	(2) grazing resulted in higher N and C content in litter; and (3) grazing accelerated litter decomposition, while grazing exclusion
29	promoted N release from litter and increased soil organic carbon. In addition, incubation site had significantly more impact
30	than did litter quality on litter decomposition and N release, while litter quality affected decomposition in the early stages. This
31	study provides insights into the mechanisms behind the nutrient cycling in alpine ecosystems. We suggest that periodic grazing
32	and grazing exclusion is beneficial in grassland management on the QTP.

1 Introduction

37	The Qinghai-Tibetan Plateau (QTP) represents an important eco-region in China (Wen et al., 2010), in which alpine grasslands
38	cover more than 85 % of total area and are regarded as the major land unit of natural pastures in China (Dong et al., 2010).
39	However, the grassland systems in this region have suffered from severe degradation driven by a range of factors including
40	climate change, overgrazing, over-cultivation and poor management (Han et al., 2008; Li et al., 2009; Wu et al., 2009; Feng et
41	al., 2010; Wu et al., 2010), and with the degraded land area increasing at 1.2-7.44 % per year (Ma et al., 2007). Since the
42	1990's, the restoration of degraded grasslands has attracted considerable attention (Kang et al., 2007; Han et al., 2008), and
43	some efforts has recently been directed towards grassland restoration and maintenance by increasing aboveground plant
44	abundance (Niu et al., 2009) and biodiversity (Wu et al., 2009; Niu et al., 2010), and improving soil organic matter content
45	and nutrient availability (Cao et al., 2004; Wu et al., 2010; Sun et al., 2011). It is well known that grazing may change the
46	vegetation community structure, soil structure and nutrient cycling processes, and that such changes have important
47	consequential impacts on the structure and functioning of the ecosystem as a whole. However, litter decomposition and N
48	release, the key factors regulating the nutrient cycle and availability at the soil-plant interface through grazing (Carrera and
49	Bertiller, 2013), are as yet little studied in these alpine ecosystems (Luo et al., 2010; Zhu et al., 2016).
50	Herbivore grazing may induce short-term ecophysiological changes in plant tissues which in turn may translate into
51	litter quality changes, and long-term shifts in plant community composition. At the short-term ecophysiological level, grazing
52	may promote the plant species producing high-quality litter (Holland and Detling, 1990; Sirotnak and Huntly, 2000; Olofsson

53	and Oksanen, 2002; Semmartin et al., 2004, 2008), because the consuming of plant tissues by herbivores may favour the grazed
54	species with a higher re-growth rate and greater nutrient contents in plant tissues due to the higher nutrient uptake (see Holland
55	and Detling, 1990; Olofsson and Oksanen, 2002; Semmartin et al., 2008). At the long-term community level, selective foliar
56	grazing may alter the competitive interactions and recruitment patterns of plant species, which may change their abundance
57	and life form structure (Bardgett and Wardle, 2003; Semmartin et al., 2008; Wu et al., 2009; Niu et al., 2010). For instance,
58	herbivores preferentially feed on the most palatable plants (e.g., species with high nutrient and low fibre contents), which may
59	promote dominance of unpalatable species (Garibaldi et al., 2007), resulting in the high inputs of low-quality litter to soil and
60	thus a reduction of decomposition rate, nutrient availability and nutrient cycling (Ritchie and Knops, 1998; Moretto et al.,
61	2001; Olofsson and Oksanen, 2002). Therefore, litter in grassland subject to long-term grazing may decompose more slowly.
62	However, some studies demonstrate that grazing per se may accelerate litter decomposition by modifying site conditions for
63	litter turnover through physical changes in the soil by herbivore activities, such as trampling and urine/dung deposition (Takar
64	et al., 1990; Fahnestock and Knapp, 1994; Semmartin et al., 2008; Luo et al., 2010; Liang et al., 2018). Empirical evidence of
65	variance in litter quality input and decomposition caused by grazing is still subject to debate (Garibaldi et al., 2007).
66	It is often assumed that higher nutrient content in plant tissue usually results in faster litter decomposition, and in higher
67	nutrient mineralization and availability in soil (Olofsson and Oksanen, 2002). At the ecosystem scale, the chemical
68	characteristics of plant litter, for example the carbon:nitrogen ratio (C:N) and/or nitrogen and lignin content, are often regarded
69	as the indicators of litter quality (Aerts, 1997; Strickland et al., 2009). Many studies have demonstrated a positive correlation
70	between litter decomposition rate and litter N content, or a negative relationship between litter decomposition rate and initial

- 71 litter lignin content and C:N or lignin:N ratio (e.g., Wardle et al., 2002; Aerts et al., 2003; Semmartin et al., 2004; Garibaldi et
- 72 al., 2007; Luo et al., 2010; Vaieretti et al., 2013).

73	In addition to litter quality (its chemical composition), two further factors controlling litter decomposition are the
74	climate (mainly temperature and humidity) and decomposing organisms (their abundance and activity) (Coûteaux et al., 1995;
75	Aerts, 1997; Semmartin et al., 2004; Keeler et al., 2009; Berg and McClaugherty, 2014; Zhu et al., 2016). Climate usually
76	regulates decomposition processes at global and regional scales (Coûteaux et al., 1995; Silver and Miya, 2001), but microbial
77	activity regulates decomposition processes through soil temperature and moisture effects (modified by grazing) at a local scale
78	(Coûteaux et al., 1995; De Santo et al., 1993; Luo et al., 2010; Orsborne and Macauley, 1988). Generally, climatic influence
79	dominates litter quality and decomposer activity in areas where weather conditions are unfavourable (Coûteaux et al., 1995),
80	due to the dependence of decomposer activity on microclimate (De Santo et al., 1993). Under favourable conditions, litter
81	quality may largely prevail as the regulator and remain important until the late decomposition stages (Coûteaux et al., 1995).
82	However, specific temperature and moisture conditions and litter quality may interact strongly and thus the rate of litter
83	decomposition is difficult to predict.
84	Most studies evaluating the effect of grazing on litter decomposition usually focus on forest, grassland or crop
85	ecosystems in temperate areas (e.g., Aber and Melillo, 1980; Berg and Staaf, 1981; Luo et al., 2010; McCurdy et al., 2013),
86	largely ignoring those in the alpine zones. On the QTP, previous studies prove that long-term grazing exclusion (> 2 years)
87	may promote plant abundance and biodiversity (Niu et al., 2009, 2010; Wu et al., 2009, 2010); however, exclusion may limit
88	the efficient use of grassland. By contrast, the short-term effect of grazing exclusion is seldom studied. Other previous studies

89	have focused on the dominant species only (e.g., Wu et al., 2009; Luo et al., 2010) and this approach provides less insight into
90	the nature of nutrient cycling in the grasslands than work on mixed litter (Zhu et al., 2016; this study). Moreover, few studies
91	have simultaneously investigated how both litter quality and incubation site affect litter decomposition (e.g., Luo et al. 2010;
92	Zhu et al. 2016; Liang et al. 2018). In this study, we examined the short-term effect (6 month) of grazing and grazing exclusion
93	on plant community composition and litter quality and their longer-term effect on mixed litter decomposition and N release
94	(27 month) and soil chemical characteristics (33 month). Based on the above, this research aimed to test three hypotheses: (1)
95	short-term grazing exclusion does not change plant community composition and litter quality (i.e., nutrient content as N and
96	biomass of palatable plant species), (2) grazing may accelerate litter decomposition and N release and thus increase soil organic
97	carbon and N, and (3) litter quality has less effect on litter decomposition and N release compared to incubation site. Results
98	of the present study may improve our understanding of nutrient cycling in alpine regions in general, and this study may also
99	further provide knowledge relevant to the development of strategies for restoring the degraded grasslands on the QTP in
100	particular.
101	

2 Materials and methods

2.1 Experimental site

This study site was an alpine meadow on the eastern QTP, SW China (N33°59′, E102°00′, altitude 3,500 m above sea level).
The mean annual temperature is 1.2 °C, ranging from –10 °C in January to 11.7 °C in July, with approximately 270 days with

106	frost per year. The mean annual precipitation over the last 35 years, occurring mainly during the short and cool summer is 620
107	mm (Niu et al., 2010). The years during this study (i.e., 2009 – 2012) were climatically typical (Sun et al., 2015; Supplemental
108	Fig. 1).
109	The grassland selected for experiments was > 9 ha in area (including 6 ha of experimental plots and 3 ha buffer areas)
110	and regularly used for Tibetan sheep and yak grazing during the grazing season (May – October). The slopes at the site are
111	less than 5 %, typical of the gently undulating topography of the region. The soil properties in the experimental plots were
112	similar after a long-term grazing history with the same grazing pattern. The soil type at the experiment site is an alpine meadow
113	soil, similar to the Mat-Cryic Cambisols described by Wu et al. (2010).
114	
115	2.2 Litter composition and quality
116	To measure the annual litter composition and determine whether plants could recover without grazing, three grazing (GP, 100
117	$m \times 200 \text{ m}$) and three grazing exclusion paddocks (GEP, $30 \text{ m} \times 20 \text{ m}$) were established when all aboveground plants were
118	dormant in October 2009. Grazing in GP started with an optimal moderate stocking rate of 4 Tibetan sheep/ha from April 2010.
119	The mean body weight of sheep was about 38 kg when used for the experiment. Before grazing started, 20 quadrats (0.5 m \times
120	0.5 m) identified by GPS coordinates were randomly established within the GP or GEP, and litter was cleared soon after the
121	establishment of quadrats. In October 2010, we collected all plant litter from each quadrat of the GP and GEP for two purposes:
122	
	(1) measurement of litter composition and quality in this experiment, and (2) measurement of litter decomposition and N

124	$m \times 0.25$ m) along diagonal; see Supplemental Fig. 2] were designed to minimize the sample variance caused by the uneven
125	litter distribution and to ensure the similar composition and quality of litter used for this and the next experiment.
126	To measure the litter composition, litter of different species collected from each quarter was identified. After litter
127	species identification, litter was separated into two groups of contrasting palatability to the Tibetan sheep (Niu et al., 2009,
128	2010; Wu et al., 2009; see Supplementary Table 1): (1) palatable species - preferred and desirable species, and (2) unpalatable
129	species - undesirable and toxic species. To measure the dry biomass, the palatable and unpalatable litter was separately oven-
130	dried at 60 °C for 48 h and then weighed.
131	To measure the quality of litter collected from GP (GP-litter treatment) or from GEP (GEP-litter treatment), the
132	palatable and unpalatable litter from a quarter was mixed again, then ground and stored in a zip-lock bag with 10 g per bag for
133	a quality test. There were six replicates for each treatment. The contents of lignin, cellulose and hemicellulose were measured
134	as descripted by van Soest et al. (1991). Organic carbon concentration (C) was measured by the $K_2Cr_2O_7-H_2SO_4$ oxidation
135	method of Walkley-Black (Nelson and Sommers, 1996). The total Kjeldahl nitrogen (N) and total phosphorus (P) were
136	analyzed using a FIAstar 5000 flow injection analyzer (Foss Tecator, Högnäs, Sweden) (Chen et al., 2016). We also calculated
137	the ratios of C:N, lignin:N, cellulose:N and hemicellulose:N.
138	We also examined the effects of grazing and grazing exclusion on soil characteristics. We randomly collected five soil
139	samples in each experimental paddock (n=30 in total) from the $0 - 10$ cm depth using a bucket auger (10 cm in diameter) in
140	October of 2010, 2011 and 2012. The same methods used to test litter quality (i.e., Nelson and Sommers, 1996; Chen et al.,
141	2016) were applied to estimate the soil organic carbon (SOC), total nitrogen (TN) and total phosphorus (TP).

143 **2.3 Litter decomposition and N release**

144 In this experiment, we included four treatments: (1) GP-GP, litter of all species was collected from and incubated in the GP;

- 145 (2) GEP-GEP, litter of all species was collected from and incubated in the GEP; (3) GP-GEP, litter of all species was collected
- 146 from the GP but incubated in the GEP; (4) GEP-GP, litter of all species was collected from the GEP but incubated in the GP.

147 Treatments 1 and 2 were designated "in situ" incubation treatments, while treatments 3 and 4 were designated "across" grazing

148 category incubation treatments, and these were included to improve understanding of the "home-field advantage" effect on

149 litter deposition (John et al., 2011).

150 For each sample soil particles attached to the litter were cleaned off with a soft brush, and samples were air-dried for three 151 days. Dry litter collected from each quadrat was cut to ≈ 5 cm length and 10 g litter was packed into a nylon litter-bag (15 cm 152 \times 20 cm with mesh size of 0.35 mm) (Cornelissen, 1996), which should have prevented any loss of material and had no effect 153 on litter decomposition (Cornelissen et al., 1999). On 20th Oct 2010, the packed litter was incubated above the soil surface by 154 fastening to the ground surface with four steel stakes to prevent removal by sheep and small animals (Vaieretti et al., 2013), 155 such as the plateau pika *Ochotona curzoniae* (Hodgson). For each treatment, 24 litter-bags were incubated with 20 cm apart 156 from each other to reduce the mutual interference. Three litter bags from each treatment were retrieved after incubation periods of 56, 141, 247, 391, 444, 582, 695 or 799 days (i.e., on 15th December 2010, 10th March, 24th June and 4th September 2011, 157 and 7st January, 24th May, 14th September and 27th December 2012, respectively). There were a total of 96 litter-bags used in 158

this experiment. Retrieved litter was brought back to the laboratory, cleaned by removing any extraneous material attached,

160	and then weighed after being oven-dried at 60 °C for 48 h. Samples were ground and stored in a zip-lock bag for further
161	chemical analyses as mentioned above. We estimated the litter decomposition and N release as the percentage of dry weight
162	lost during each incubation period (Cornelissen et al., 1999; Vaieretti et al., 2013).
163	
164	2.4 Statistical analyses
165	A goodness-of-fit test (Shapiro-Wilk test, Univariate Procedure) was used to test the normality of data before mean comparison
166	using analysis of variance (ANOVA, GLM Procedure). All data were normally distributed. Data on the initial chemical
167	characteristics of litter (Table 1) were analyzed using ANOVA followed by Tukey's Studentized multiple range test. Data on
168	the biomass of palatable or unpalatable species and that on the total biomass between GP and GEP were also analyzed using
169	ANOVA, while for GP or GEP the difference in litter biomass between the palatable and unpalatable species was compared
170	by paired-t test (TTEST Procedure) (Fig. 1). Data on the final proportion of litter biomass or N remaining (Fig. 4), litter quality
171	(content of organic carbon, nitrogen, phosphorous and other chemical characteristics of litter) (Table 1), and soil SOC, TN and
172	TP (Fig. 2) were analyzed using ANOVA followed by least significant difference test (LSD test) for multiple comparisons.
173	The decomposition rate $(k, g \cdot 10 g^{-1} \cdot day^{-1})$ of litter biomass during the incubation period (Table 2) was assessed using a
174	negative exponential model according to Swift et al. (1979): $y = a_*e^{-(t*k)}$, where y is the dry biomass of litter remaining in the
175	litter bags at time t (days), a is the initial litter biomass (i.e., 10 g in this study). The difference in decomposition rate between
176	treatments was compared according to Julious (2004), i.e. there is deemed to be no significant difference in decomposition
177	rate if their 83.4 % CL overlap. The decomposition rate and 83.4 % CL were estimated by fitting the negative exponential

178 model to a nonlinear least square regression model (NLIN Procedure).

179	A multivariate regression model (GLM Procedure) employed by Vaieretti et al. (2013) was applied to quantify the
180	effect of incubation site and litter quality (the two independent factors) on the final litter decomposition or N release (the
181	dependent factor) (Table 3): litter decomposition or N release = Site + Quality + Site×Quality + ϵ , where 'Site' is the paddock
182	category where the litter was incubated (i.e., incubation site: GP and GEP), 'Quality' is the litter quality reflecting the sources
183	where the litter was collected from (i.e., GP and GEP), and ϵ is the model error. The proportional contribution of incubation
184	site, litter quality and their interaction to variability of litter decomposition or N release was calculated as the sum of squares
185	for each of the terms, divided by the total sum of squares. The Type I sum of squares was used because of the balanced design
186	of this experiment. All analyses were done using SAS 9.3 (SAS Institute Inc., Cary, NC, USA). The rejection level for H_0 was
187	set at $\alpha < 0.05$. Values of the means (± SE) are presented in Figs. 1-4.
188	
189	3 Results
190	3.1 Litter composition and quality and soil property
191	Fifty-five plant species (mostly forbs and grasses along with several legumes and sedges) were identified, and all were found
192	in both GP and GEP, except Gentiana macrophylla Pallas, which was only found in the GP (Supplemental Table 1). However,
193	even though the annual litter biomass of unpalatable species in both GP and GEP was similar (ANOVA: $F_{1,38} = 3.43$, P =
194	0.0717), litter biomass of palatable species was significantly greater in the GEP than in the GP (ANOVA: $F_{1,38} = 75.32$, P <

^{195 0.0001),} and this difference, which contributed significantly more to the total litter biomass in the GEP than in the GP (ANOVA:

196	$F_{1,38} = 114.66$, P < 0.0001) (Fig. 1). The litter biomass was not significantly different between palatable and unpalatable species
197	in the GP (Paired-t test: $t_{19} = 0.96$, P = 0.3510); however, in the GEP, litter biomass of palatable species was significantly
198	greater than that of unpalatable species (Paired-t test: $t_{19} = 7.17$, P < 0.0001) (Fig. 1).
199	GP-litter had significantly higher C and N but significantly lower hemicellulose and hemicellulose:N than GEP-litter
200	(Table 1). Although other quality characteristics were lower in GP-litter than in GEP-litter, the differences were not significant
201	(Table 1).
202	The concentrations of soil TN and TP were not significantly different between the GP and GE for any year (LSD =
203	0.0002 - 0.0015 and $0.0003 - 0.0004$ for TN and TP, respectively; P > 0.05) (Fig. 2a-b). Similarly, there was no significant
204	difference in SOC concentration between the GP and GE in 2010 and 2011 (LSD = 0.0169 and 0.0111 for 2010 and 2011,
205	respectively; $P > 0.05$), while in 2012 SOC was significantly higher in GEP than in GP (LSD = 0.0138, P = 0.0279) (Fig. 2c).
206	
207	3.2 Litter decomposition
208	The proportion of litter biomass remaining continuously decreased with incubation duration and litter decomposed faster in
209	the first year (i.e., 44.6, 38.5, 41.46 and 31.6 % decomposition in GP-GP, GEP-GEP, GEP-GP and GP-GEP, respectively)
210	than in the second year (i.e., 18.8, 24.1, 27.6 and 23.4 % decomposition in GP-GP, GEP-GEP, GEP-GP and GP-GEP,
211	respectively) (Fig. 3a). As shown in Table 2, the decomposition rate (k) of litter incubated in GP was significantly higher than
212	that in GEP (non-overlap of 83.4 % CL), i.e., for the "in situ" treatments k in GP-GP > k in GEP-GEP and for the "across"
213	treatments k in GEP-GP > k GP-GEP. The final proportion of litter biomass remaining was significantly lower in GP-GP and 12

214 GEP-GP than in GP-GEP (LSD = 2.51, P < 0.0001) (Fig. 4a).

- 215
- 216 **3.3 N release**
- 217 Generally, the percentage of total N release did not change during the first winter when temperature was < 0 °C, except that it
- increased during the incubation period of December 2010 and March 2011 (first winter) in the treatment GEP-GP (Fig. 3b).
- 219 From January in the second winter (2012), the percentage of total N remaining decreased steadily until the end of the
- 220 experiment (Fig. 3b). The final proportion of total N remaining was significantly higher in GEP-GP and significantly lower in

221 GEP-GEP and GP-GEP (LSD test: LSD = 5.36, P = 0.0002) (Fig. 4b).

222

223 **3.4** Contribution of incubation site and litter quality to litter decomposition and N release

- The multivariate regression model indicates that both incubation site and litter quality significantly affected litter decomposition and N release (Table 3). Incubation site contributed respectively near 25 % and 50 % more to litter decomposition and N release than did litter quality (Table 3). Furthermore, the model predicts that GP resulted in significantly greater litter decomposition (8.13 %) but significantly lower N release (9.73 %) than did GEP ($F_{1,8} = 62.48$ and 57.49 for litter decomposition and N release, respectively; P < 0.0001). Results show that GEP-litter decomposed significantly faster (2.5 %) but released N significantly more slowly (9.27 %) than GP-litter ($F_{1,8} = 34.99$ and 15.80 for litter decomposition and N release, respectively; P < 0.01).
- A significant interaction between incubation site and litter quality for litter decomposition was found ($F_{2,8} = 7.10$, P =

232	0.0286), i.e., litter collected from GEP but incubated in GP decomposed significantly faster (also see Fig. 4a). However, the
233	interaction between incubation site and litter quality on N release was not significant ($F_{2,8} = 2.76$, P = 0.1350).
234	
235	4 Discussion
236	4.1 Litter Composition and Quality
237	Grazing or grazing exclusion of herbivores may indirectly alter the species composition and functioning of grasslands by
238	inducing shifts in plant competitive interactions and recruitment patterns, and thus changes in species abundance and life form
239	structure (Bardgett and Wardle, 2003; Garibaldi et al., 2007; Semmartin et al., 2008; Wu et al., 2009; Niu et al., 2010; Chaneton,
240	2011). However, our results indicate that herbivore grazing or grazing exclusion did not alter plant community composition in
241	terms of species inventory, as species found in the GEP mostly also occurred in the GP. On the QTP, species composition is
242	grazing-intensity-dependent (Niu et al. 2010; Sun et al., 2011) and/or grazing exclusion-period-dependant (Wu et al., 2009).
243	Thus our results imply that a stocking rate of 4 Tibetan sheep/ha in the GP or a short period of grazing exclusion (i.e., from
244	April to October) in the GEP did not change the species composition.
245	However, our results show that herbivore grazing significantly altered species composition in terms of species
246	abundance or palatability with significantly less palatable as well as total litter produced in the GP compared to the GEP (Fig.
247	1). The low biomass of palatable species in GP may be attributed to the more palatable species (mostly the grasses and sedges,
248	see Supplemental Table 1) on the QTP being taller (Sun et al., 2011), and therefore more accessible to herbivore grazing, in
249	addition to being more likely to be grazed by preferential grazing. Through these two mechanisms, the biomass of palatable 14

250	species in the GP would subsequently be reduced. Results of this study indicate that grazing exclusion for a short period may
251	allow the recovery of palatable species in the alpine meadows. However, there was no significant difference in the litter
252	biomass of unpalatable species between the GP and GEP (Fig. 1), which provides evidence against the assumption that
253	removing the canopy of palatable species may allow the intra- and inter-specific competition for light which ultimately favours
254	the establishment of short, less-palatable species (Sternberg et al., 2000; Pavlů et al., 2008; Wu et al., 2009; Sun et al., 2011).
255	It is generally accepted that litter quality is usually determined by the levels of various chemical compounds such as
256	soluble C, N and P, as well as lignin or lignin:N ratio, and litter of high quality usually has higher N content but lower lignin
257	and lignin:N ratio (e.g., Aerts, 1997; Olofsson and Oksanen, 2002; Wardle et al., 2002; Garibaldi et al., 2007; Strickland et al.,
258	2009). In this study, grazing might have improved litter quality at least to some extent by significantly increasing the N content
259	and potentially reducing the hemicellulose content and C:N, lignin:N, cellulose:N and hemicellulose:N ratios (Table 1).
260	Therefore, our results <mark>align with</mark> previous studies that <mark>have indicated</mark> grazing may promote litter quality <mark>due to the high nutrient</mark>
261	uptake (e.g., Sirotnak and Huntly, 2000; Olofsson and Oksanen, 2002; Semmartin et al., 2004).
262	
263	4.2 Litter decomposition
264	For a given climatic region, the ecological processes of litter decomposition are regulated by incubation
265	microenvironment (i.e., grazing/grazing exclusion and soil property in this study) and litter quality. Our results suggest that
266	herbivore grazing played a major role in litter decomposition on the QTP. Firstly many studies have demonstrated that litter

267 quality is one of the most important factors affecting the litter decomposition, and litter with higher N content but lower lignin

268	and lignin:N ratio will decompose faster (Aerts, 1997; Olofsson and Oksanen, 2002; Wardle et al., 2002; Garibaldi et al., 2007;
269	Strickland et al., 2009). Therefore, it may be expected that regardless of incubation site, GP-litter that had significantly higher
270	N content should decompose faster than GEP-litter. However, evidence for this expectation was only detected for the "in situ"
271	incubation treatments (i.e., greater decomposition rate in GP-GP than in GEP-GEP), while for the "across" incubation
272	treatments opposite results were found (i.e., greater decomposition rate in GEP-GP than in GP-GEP) (Table 2 and Fig. 4a).
273	Our experimental site is located at high altitude with a low mean annual temperature of 1.2 °C; hence the activity of
274	decomposers may be inhibited during the cold seasons (Coûteaux et al., 1995). Therefore, litter quality may not be a good
275	predictor of litter decomposability in cold temperate regions (Aerts, 1997), where climate is more important than litter quality
276	in the regulation of litter decomposition (Coûteaux et al., 1995). On the QTP, decomposition rate ($k = 1.04 \times 10^{-3} - 1.34 \times 10^{-3}$
277	³ g·10 g ⁻¹ ·day ⁻¹ in Table 2, i.e. $k = 0.38 - 0.48$ g·g ⁻¹ ·year ⁻¹) was much lower than that of the global mean ($k \approx 0.75$ g·g ⁻¹ ·year ⁻¹)
278	with similar latitudes (N30° - 40°) (Zhang et al., 2008).
279	Additionally, incubation site had significantly greater effect on litter decomposition ($\approx 25\%$) than did litter quality
280	(Table 3), and regardless of litter quality, litter decomposed faster in GP (19.9 % for GP-litter and 11.8 % for GEP-litter) than
281	in GEP (Fig. 4a; Table 2). This may be attributed to the effect of herbivore grazing activity, which modifies the incubation site
282	conditions for litter turnover (Takar et al., 1990; Fahnestock and Knapp, 1994; Semmartin et al., 2008; Luo et al., 2010; Liang
283	et al., 2018). Our results further demonstrate that regardless of incubation site and litter quality, litter decomposed faster in the
284	first year (31.6 – 44.6 % decomposition) than in the second year (18.8 – 27.6 % decomposition) (Fig. 3a). Berg (2014) and
285	Berg and McClaugherty (2014) have stated that litter decomposition rate varies at different stages. In the early stages of 16

286	decomposition litter mass is lost rapidly via the leaching of soluble compounds, while in the later stages decomposition can
287	even cease after only recalcitrant litter compounds remain (Berg 2014; Berg and McClaugherty 2014). Therefore, litter quality
288	regulates decomposition processes mainly in the early stages of decomposition (Berg and McClaugherty, 2014), which is
289	supported by our results, i.e., when incubated in GP in the first year, the percentage decomposition of GP-litter (44.6 %) was
290	higher than that of GEP-litter (41.5%). However, we could not reject the above conclusion that incubation site is the dominant
291	factor affecting litter decomposition over litter quality, as in the first year litter always decomposed faster in GP (i.e., 44.6 %
292	in GP-GP and 41.5 % in GEP-GP) than in GEP (i.e., 38.5 % in GEP-GEP and 31.6 % in GP-GEP) (Fig. 3a). These data also
293	imply that a home-field advantage is detected only for GP-litter during the first year of incubation (i.e., percentage
294	decomposition 44.6 % in GP-GP > 31.6 % in GP-GEP) with no evidence for GEP-litter (i.e., percentage decomposition 38.5 %
295	in GEP-GEP < 41.5 % in GEP-GP).
296	In a long-term (9 year) study on the QTP, Wu et al. (2009) reported that grazing exclusion favors the increase of soil
297	total nitrogen, soil organic matter, soil organic carbon, soil microbial biomass carbon and soil carbon storage. It is intersting
298	that in the present study, only soil organic carbon (SOC) significantly increased after three-years grazing exclusion (Fig. 2).
299	The increass in SOC in GEP may be because grazing exclusion prevents the reduction of removal of palatable litter by the
300	herbivores (Fig. 1), and the organic C locked within plant tissues may be returned to the soil during litter decomposition instead
301	(Bardgett and Wardle, 2003; Wu et al., 2009). Holland and Detling (1990) and Ågren et al. (1999) stated that increasing carbon
302	availability in soil may promote decomposer growth and activity even at low nitrogen concentrations. However, the expected
303	results, i.e. <mark>significantly</mark> higher litter decomposition rate caused by the possible increasing decomposer mass and/or activity in 17

304	the grazing exclusion grasslands (Wu et al., 2009), were not observed in the GEP in this study (Table 2, Fig. 4a). Thus during
305	a relatively short period the soil properties are unlikely to be significantly changed through grazing or grazing exclusion, and
306	thus limited duration grazing events have a smaller effect on litter decomposition under cool environments on the QTP, than
307	in experiments conducted in warmer climates.
308	
309	4.3 N release
310	N release is a more complex process, compared to litter decomposition. N release may involve any one or both processes of N
311	immobilization and N mineralization, where the former results in the accumulation of N in the litter and the latter causes the
312	release of N from the litter (Manzoni et al., 2008). Swift et al. (1979) and Berg and McClaugherty (2014) reported that the
313	biological decomposition of litter is mainly carried out by microbial decomposers, which per se have a higher N:C ratio
314	compared with most litter types. This property of decomposers creates a high N demand for decomposer growth (Manzoni et
315	al., 2008). Results show that regardless of litter quality or source, N remaining was significantly higher when litter was
316	incubated in GP than in GEP (Fig. 4b). Bosatta and Balesdent, (1996) and Manzoni et al. (2008), a promising candidate
317	mechanism may be that the faster decomposition rate of litter in GP increases the utilization of C by the decomposers which
318	in turn increases the N:C ratio in litter; when N:C ratio is high, large amounts of mineral N are immobilized, increasing the N
319	concentration in litter. This mechanism may also account for the dynamics of N release over the incubation period (Fig. 3b).
320	For instance, decomposer activity resulted in continuous C consumption and litter decomposition (March – June 2011), while
321	high N:C ratio due to decomposition progress induced the release of accumulated N (June – August 2011). The inverse pattern

322	of N release and litter decomposition found in this study (Fig. 4) is frequently reported (e.g., Aber and Melillo, 1980; Fahey et
323	al., 1991; Gallardo and Merino, 1992).
324	It is not surprising that because both litter decomposition and N release are regulated by decomposers synchronously,
325	incubation site also had a significantly greater effect ($\approx 50\%$) on N release as on litter decomposition (Table 3) than did litter
326	quality, indicating the latter is not a good predictor on litter N release in the cold temperate region (Aerts, 1997).
327	
328	5 Conclusion
329	Results of our study are not completely consistent with previously proposed hypotheses. On the cold QTP, short-term grazing
330	exclusion did not promote species abundance but increases plant palatability and total litter biomass. Grazing improves litter
331	quality through higher N content but lower hemicellulose and hemicellulose:N ratio. Grazing significantly accelerated litter
332	decomposition, while grazing exclusion promoted N release and increased SOC. Although litter quality may affect
333	decomposition at the early stages, incubation site had significantly more impact on both litter decomposition and N release.
334	The different effects of grazing and grazing exclusion functioning on the grassland ecosystems may have implications in the
335	management of alpine meadows on the QTP. For example, periodic grazing and grazing exclusion may be a good option which
336	allows plant recovery and promotes nutrient cycling, and thus contributes to the restoration of degraded grasslands.
337	
338	Author contributions. YS and FH designed, and FH supervised the experiments, YS, FH, ZFW and SHC performed research
339	and collected data. XZH, FH and YS analysed data and prepared the manuscript, and all authors contributed to the writing.

341 *Competing interests.* The authors declare that they have no conflict of interest.

2	1	0
э	4	2

343	Acknowledgments.	We are very	grateful	to Dr.	JA.	Subke	(Editor)	and	two a	anonymous	reviewers	for	their	constructive
	0									2				

- 344 comments and suggestions, which have significantly improved the paper. We are also grateful to Professor C. Matthew for his
- 345 valuable comments and time spent in English editing of the manuscript. This study was financially supported by the National
- 346 Key Project of Scientific and Technical Supporting Programs (2014CB138706), National Natural Science Foundation of China
- 347 (No. 31672472), Program for Changjiang Scholars and Innovative Research Team in University (IRT13019) and the
- 348 independent grants from the State Key Laboratory of Grassland Agro-ecosystems (SKLGAE201708).
- 349

350 References

- 351 Aber, J. D. and Melillo, J. M.: Litter decomposition: measuring relative contributions of organic matter and nitrogen to forest
- 352 soils, Can. J. Biochem., 58, 416-421, doi:10.1139/b80-046, 1980.
- 353 Aerts, R., De Caluwe, H. and Beltman, B.: Plant community mediated vs. nutritional controls on litter decomposition rates in
- 354 grasslands, Ecology, 84, 3198-3208, doi:10.1890/02-0712, 2003.
- 355 Aerts, R.: Climate, leaf litter chemistry and leaf litter decomposition in terrestrial ecosystems: a triangular relationship, Oikos,
- 356 79, 439-449, doi:10.2307/3546886, 1997.

- 357 Ågren, G. I., Shaver, G. R. and Rastetter, E. B.: Nutrients: dynamics and limitations, in: Luo, Y., Mooney, H. A. (Eds.): Carbon
- dioxide and environmental stress, Academic Press, San Diego, 333-345 pp., 1999.
- 359 Bardgett, R. D. and Wardle, D. A.: Herbivore-mediated linkages between aboveground and belowground communities,
- 360 Ecology, 84, 2258-2268, doi:10.1890/02-0274, 2003.
- Berg, B.: Decomposition patterns for foliar litter A theory for influencing factors. Soil Biol. Biochem., 78, 222-232,
 doi:org/10.1016/j.soilbio.2014.08.005, 2014.
- 363 Berg, B. and McClaugherty, C.: Plant litter: decomposition, humus formation, carbon sequestration, 3rd ed., Springer, 315 p.,
- 364 2014.
- 365 Berg, B. and Staff, H.: Leaching, accumulation and release of nitrogen in decomposing forest litter, in: Terrestrial Nitrogen
- 366 Cycles, Clark, F. E. and Rosswall, T. (Eds.): Processes, Ecosystem Strategies and Management Impacts, Ecological

367 Bulletins, Stockholm, Sweden, 33, 163-178 pp., 1981.

- 368 Bosatta, E. and Balesdent, J.: Isotope discrimination during decomposition of organic matter: a theoretical analysis, Soil Sci.
- 369 Soc. Am. J., 60, 1121-1126, doi:10.2136/sssaj1996.03615995006000040023x, 1996.
- 370 Cao, G. M., Tang, Y. H., Mo, W. H., Wang, Y. S., Li, Y. N. and Zhao, X. Q.: Grazing intensity alters soil respiration in an
- alpine meadow on the Tibetan plateau, Soil Biol. Biochem., 36, 237-243, doi:10.1016/j. soilbio.2003.09.010, 2004.
- 372 Carrera, A. L. and Bertiller, M. B.: Combined effects of leaf litter and soil microsite on decomposition process in arid
- 373 rangelands, J. Environ. Manage., 114, 505-511, doi:10.1016/j.jenvman.2012.10.059, 2013.

- 374 Chaneton, E. J.: Linking grazing-induced changes in plant biodiversity to rangeland ecosystem function, IX International
- 375 Rangeland Congress, Lugar, Rosario, Año, 286-291 pp., 2011.
- 376 Chen, X. J., Hou, F. J., Matthew, C. and He, X. Z.: Soil C, N, and P stocks evaluation under major land uses on China's Loess
- 377 Plateau, Rangeland Ecol. Manage., 70, 341-347, doi:10.1016/j.rama.2016.10.005, 2016.
- 378 Cornelissen, J. H. C., Harguindeguy, N. P., Díaz, S., Grime, J. P., Marzano, B., Cabido, M., Vendramini, F. and Cerabolini,
- B.: Leaf structure and defence control litter decomposition rate across species and life forms in regional floras on two
- 380 continents, New Phytol., 143, 191-200, doi:10.1046/j.1469-8137.1999.00430.x, 1999.
- 381 Cornelissen, J. H. C.: An experimental comparison of leaf decomposition rates in a wide range of temperate plant species and
- 382 types, J. Ecol., 84, 573-582, doi:10.2307/2261479, 1996.
- 383 Coûteaux, M. M., Bottner, P. and Berg. B.: Litter decomposition, climate and litter quality, Trends Ecol. Evol., 10, 63-66,
- 384 10.1016/S0169-5347(00)88978-8, 1995.
- 385 De Santo, A. V., Berg, B., Rutigliano, F. A., Alfani, A. and Floretto, A.: Factors regulating early-stage decomposition of needle
- 386 litters in five different coniferous forests, Soil Biol. Biochem., 25, 1423-1433, doi:10.1016/0038-0717(93)90057-I, 1993.
- 387 Dong, S. K., Wen, L. U., Zhu, L. and Li, X. Y.: Implication of coupled natural and human systems in sustainable rangeland
- 388 ecosystem management in HKH region, Front. Earth Sci. China, 4, 42-50, doi:10.1007/s11707-010-0010-z, 2010.
- 389 Fahey, T. J., Stevens, P. A., Hornung, M. and Rowland, P.: Decomposition and nutrient release from logging residue following
- 390 conventional harvest of Sitka spruce in North Wales, Forestry, 64, 289-301, doi:10.1093/forestry/ 64.3.289, 1991.

- Fahnestock, J. T. and Knapp, A. K.: Plant responses to selective grazing by bison: interactions between light, herbivory and
 water stress, Vegetatio, 115,123-131, doi:10.1007/BF00044867, 1994.
- 393 Feng, R. Z., Long, R. J., Shang, Z. H., Ma, Y. S., Dong, S. K., Wang, Y. L.: Establishment of *Elymus natans* improves soil
- quality of a heavily degraded alpine meadow in Qinghai-Tibetan Plateau, China, Plant Soil, 327, 403-411,
 doi:10.1007/s11104-009-0065-3, 2010.
- 396 Gallardo, A. and Merino, J.: Nitrogen immobilization in leaf litter at two Mediterranean ecosystems of SW Spain,
- 397 Biogeochemistry, 15, 213-228, doi:10.1007/BF00002937, 1992.
- 398 Garibaldi, L. A., Semmartin, M. and Chaneton, E. J.: Grazing-induced changes in plant composition affect litter quality and
- nutrient cycling in flooding Pampa grasslands, Oecologia, 151, 650-662, doi:10.1007/s00442-006-0615-9, 2007.
- 400 Han, J. G., Zhang, Y. J., Wang, C. J., Bai, W. M., Wang, Y. R., Han, G. D., Li, L. H.: Rangeland degradation and restoration
- 401 management in China, Rangeland J., 30, 233-239, doi: 10.1071/RJ08009, 2008.
- 402 Holland, E. A. and Detling, J. K.: Plant response to herbivory and belowground nitrogen cycling, Ecology, 71, 1040-1049,
- 403 doi:10.2307/1937372, 1990.
- 404 John, M. G. S., Orwin, K. H. and Dickie, I. A.: No 'home' versus 'away' effects of decomposition found in a grassland-forest
- 405 reciprocal litter transplant study, Soil Biol. Biochem., 43, 1482-1489, 10.1016/j.soilbio. 2011.03.022, 2011.
- 406 Julious, S. A.: Using confidence intervals around individual means to assess statistical significance between two means, Pharm.
- 407 Stat., 3, 217-222, doi:10.1002/pst.126, 2004.

- 408 Kang, L., Han, X. G., Zhang, Z. B. and Sun O. J. X.: Grassland ecosystems in China: review of current knowledge and research
- 409 advancement, Philos. Trans. R. Soc. Lond. B. Biol. Sci., 362, 997-1008, doi:10.1098/rstb. 2007.2029, 2007.
- 410 Keeler, B. L., Hobbie, S. E. and Kellogg, L. E.: Effects of long-term nitrogen addition on microbial enzyme activity in eight
- 411 forested and grassland sites: implications for litter and soil organic matter decomposition, Ecosystems, 12, 1-15, doi:
- 412 10.1007/s10021-008-9199-z, 2009.
- 413 Li, X. G., Zhang, M. L., Li, Z. T., Shi X. M., Ma, Q. and Long, R. J.: Dynamics of soil properties and organic carbon pool in
- 414 topsoil of zokor-made mounds at an alpine site of the Qinghai-Tibetan Plateau, Biol. Fert. Soils, 45, 865-872,
- 415 doi:10.1007/s00374-009-0398-3, 2009.
- Liang, D. F., Niu, K. C., Zhang, S. T.: Interacting effects of yak dung deposition and litter quality on litter mass loss and
 nitrogen dynamics in Tibetan alpine grassland, Grass Forage Sci., 73, 123-131, 10.1111/gfs.12280, 2018.
- 418 Luo, C. Y., Xu, G. P, Chao, Z. G., Wang, S. P., Lin, X. W., Hu, Y. G., Zhang, Z. H., Duan, J. C., Chang, X. F., Su, A. L., Li,
- 419 Y. N., Zhao, X. Q., Du, M. Y., Tang, Y. H. and Bruce, K.: Effect of warming and grazing on litter mass loss and
- 420 temperature sensitivity of litter and dung mass loss on the Tibetan plateau, Global Change Biology, 16,1606-1617,
- 421 doi:10.1111/j.1365-2486.2009.02026.x, 2010.
- 422 Ma, Y. S., Zhang, Z. H., Dong, Q. M., Shi, J. J., Wang, Y. L. and Sheng L.: Application of restoration ecology in "black soil
- 423 type" degraded grassland rebuilding, Journal of Gansu Agricultural University, 42, 91-97, doi:
- 424 10.13432/j.cnki.jgsau.2007.02.021, 2007.

- 425 Manzoni, S., Jackson, R. B., Trofymow, J. A. and Porporato, A.: The global stoichiometry of litter nitrogen mineralization,
- 426 Science, 321, 684, doi:10.1126/science.1159792, 2008.
- 427 Moretto, A. S., Distel, R. A. and Didoné, N. G.: Decomposition and nutrient dynamic of leaf litter and roots from palatable
- 428 and unpalatable grasses in a semi-arid grassland, Appl. Soil Ecol., 18, 31-37, doi:10.1016/S0929-1393(01)00151-2, 2001.
- 429 Nelson, D. and Sommers, L.: Total carbon, organic carbon and organic matter, in: Sparks, D. L., Page, A. L., Helmke, P. A.,
- 430 Loeppert, R. H., Soltanpour, P. N., Tabatabai, M. A., Johnston, C. T. and Sumner, M. E. (Eds.): Methods of soil analysis.
- 431 Part 3: chemical methods, Soil Science Society of America Book Series, Number 5, Soil Science Society of America,
- 432 Wisconsin, 961-1010 pp., 1996.
- 433 Niu, K. C., Choler, P., Zhao, B. B. and Du, G. Z.: The allometry of reproductive biomass in response to land use in Tibetan
- 434 alpine grasslands, Funct. Ecol., 23, 274-283, doi:10.1111/j.1365-2435.2008.01502.x, 2009.
- 435 Niu, K. C., Zhang, S. T., Zhao, B. B. and Du, G. Z.: Linking grazing response of species abundance to functional traits in the
- 436 Tibetan alpine meadow, Plant Soil, 330, 215-223, doi:10.1007/s11104-009-0194-8, 2010.
- 437 Olofsson, J. and Oksanen, L.: Role of litter decomposition for the increased primary production in areas heavily grazed by
- 438 reindeer: a litterbag experiment, Oikos, 96, 507-515, doi:10.1034/j.1600-0706.2002.960312.x, 2002.
- 439 Orsborne, J. L. and Macauley, B. J.: Decomposition of *Eucalyptus* leaf litter: influence of seasonal variation in temperature
- 440 and moisture conditions, Soil Biol. Biochem., 20, 369-375, doi:10.1016/0038-0717(88)90018-1, 1988.
- 441 Pavlů, V., Hejcman, M., Pavlanduring, L. and Gaisler, J.: Restoration of grazing management and its effect on vegetation in
- 442 an upland grassland, Appl. Veg. Sci., 10, 375-382, doi: 10.1111/j.1654-109X.2007.tb00436.x, 2008.

- 443 Ritchie, M. E. and Knops, J. M. H.: Herbivore effects on plant and nitrogen dynamics in oak savannah, Ecology, 79, 165-177,
- 444 doi:10.1890/0012-9658(1998)079[0165:HEOPAN]2.0.CO;2, 1998.
- 445 Semmartin, M., Aguiar, M. R., Distel, R. A., Moretto, A. S. and Ghersa, C. M.: Litter quality and nutrient cycling affected by
- 446 grazing-induced species replacements along a precipitation gradient, Oikos, 107,148-160, doi:10.1111/j.0030-
- 447 1299.2004.13153.x, 2004.
- 448 Semmartin, M., Garibaldi, L. A. and Chaneton, E. J.: Grazing history effects on above-and below-ground litter decomposition
- 449 and nutrient cycling in two co-occurring grasses, Plant Soil, 303, 177-189, doi:10.1007/s 11104-007-9497-9, 2008.
- 450 Sirotnak, J. M. and Huntly, N. J.: Direct and indirect effects of herbivores on nitrogen dynamics: voles in riparian areas,

451 Ecology, 81,78-87, doi:10.1890/0012-9658(2000)081%5B0078:DAIEOH%5D2.0.CO;2, 2000.

- 452 Sternberg, M., Gutman, M., Perevolotsky, A., Ungar, E. D. and Kigel, J.: Vegetation response to grazing management in a
- Mediterranean herbaceous community: a functional group approach, J. Appl. Ecol., 37, 224-237, doi:10.1046/j.13652664.2000.00491.x, 2000.
- 455 Strickland, M. S., Osburn, E., Lauber, C., Fierer, N. and Bradford, M. A.: Litter quality is in the eye of the beholder: initial
- 456 decomposition rates as a function of inoculum characteristics, Funct. Ecol., 23, 627-636, doi:10.1111/j. 1365-
- 457 2435.2008.01515.x, 2009.
- 458 Sun, D. S., Wesche, K., Chen, D. D., Zhang, S. H., Wu, G. L., Du, G. Z. and Comerford, N. B.: Grazing depresses soil carbon
- 459 storage through changing plant biomass and composition in a Tibetan alpine meadow, Plant Soil Environ., 57, 271-278,
- 460 doi:2-s2.0-79959230740, 2011.

461	Sun, Y., Angerer, J. P. and Hou, F. J.: Effects of grazing systems on herbage mass and liveweight gain of Tibetan sheep in
462	Eastern Qinghai-Tibetan Plateau, China, Rangeland J., 37, 181-190, doi:10.1071/RJ14062, 2015.
463	Swift, M. J., Heal, O. W. and Anderson, J. M.: Decomposition in terrestrial ecosystems, Blackwell, Cambridge, Mass, 1979.
464	Takar, A. A., Dobrowolski, J. P. and Thurow, T. L.: Influence of grazing, vegetation life-form, and soil type on infiltration
465	rates and interrill erosion on a Somalion [Somalian] rangeland, J. Range. Manage., 43, 486-490, doi:10.2307/4002350,
466	1990.
467	Vaieretti, M. V., Cingolani, A. M., Harguindeguy, N. P. and Cabido, M.: Effects of differential grazing on decomposition rate
468	and nitrogen availability in a productive mountain grassland, Plant Soil, 371, 675-691, doi:10.1007/s11104-013-1831-9,
469	2013.
470	van Soest, P. J., Robertson, J. B., Lewis, B.A.: Methods for dietary fiber, neutral detergent fiber, and nonstarch polysaccharides
471	in relation to animal nutrition, J. Dairy Sci., 74, 3583-3597, doi:10.3168/jds. S0022-0302(91) 78551-2, 1991.
472	Wardle, D., Bonner, K. and Barker, G.: Linkages between plant litter decomposition, litter quality, and vegetation responses
473	to herbivores, Funct. Ecol., 16, 585-595, doi:10.1046/j.1365-2435.2002.00659.x, 2002.
474	Wen, L., Dong, S. K., Zhu, L., Li, X. Y., Shi, J. J., Wang, Y. L. and Ma, Y. S.: The construction of grassland degradation index
475	for alpine meadow in Qinghai-Tibetan Plateau, Procedia Environ. Sci., 2, 1966-1969, doi:10.1016/j.proenv.2010.10.210,
476	2010.
477	Wu, G. L., Du, G. Z., Liu, Z. H. and Thirgood, S.: Effect of fencing and grazing on a Kobresia-dominated meadow in the
478	Qinghai-Tibetan Plateau, Plant Soil, 319, 115-126, doi:10.1007/s11104-008-9854-3, 2009. 27

479	Wu, G. L., Liu, Z. H., Zhang, L. and Chen, J. M.: Long-term fencing improved soil properties and soil organic carbon storage									
480	in an alpine swamp meadow of western China, Plant Soil, 332, 331-337, doi:10.1007/s11104-010-0299-0, 2010.									
481	Zhang, D. Q., Hui, D. F., Luo, Y. Q. and Zhou G. Y.: Rates of litter decomposition in terrestrial ecosystems: global patterns									
482	and controlling factors, J Plant Ecol., 1, 85-93, doi:10.1093/jpe/rtn002, 2008.									
483	Zhu, W. Y., Wang, J. Z., Zhang, Z. H., Ren F., Chen L. T. and He, J. S.: Changes in litter quality induced by nutrient addition									
484	alter litter decomposition in an alpine meadow on the Qinghai-Tibet Plateau, Sci. Rep.UK, 6, 34290,									
485	doi:10.1038/srep34290, 2016.									
486										
487										
488										
489										
490										
491										
492										
493										
494										
495										
496	28									

497 Tables

Table 1. Initial chemical characteristics (mean ± SE) of litter collected grazing paddocks (GP-litter) and grazing exclusion

499 paddocks (GEP-litter). Unit of chemical characteristics is mg/g litter for C, N, P, lignin, cellulose and hemicellulose. Different

Compound	GP-litter		GEP-litter		LSD	Р
С	576.44 ± 4.20	a	553.03 ± 3.35	b	14.92	0.0121
Ν	7.41 ± 0.32	a	5.35 ± 0.67	b	2.05	0.0494
Р	1.39 ± 0.27	a	1.05 ± 0.27	a	1.05	0.4197
Lignin	22.94 ± 4.57	a	18.83 ± 1.67	a	13.51	0.4456
Cellulose	328.61 ± 11.55	a	385.18 ± 19.27	a	62.38	0.0655
Hemicellulose	296.76 ± 6.82	b	324.56 ± 5.52	a	24.37	0.0340
C:N	78.08 ± 3.62	a	106.82 ± 13.96	a	40.05	0.1171
Lignin:N	3.15 ± 0.72	a	3.70 ± 0.74	a	2.87	0.6210
Cellulose:N	44.40 ± 1.35	a	75.23 ± 13.15	a	36.70	0.0800
Hemicellulose:N	40.12 ± 1.21	b	62.58 ± 7.69	a	21.61	0.0447

500 letters in each row indicate significant difference (ANOVA: P < 0.05).

- 504 **Table 2.** Litter decay rate (k, g·10 g⁻¹·day⁻¹) in different incubation environments. GP-GP, mixed litter was collected from
- 505 grazing paddocks (GP) and incubated in GP; GEP-GEP, mixed litter was collected from grazing exclusion paddocks (GEP)
- 506 and incubated in GEP; GEP-GP, mixed litter was collected from GEP and incubated in GP; GP-GEP, mixed litter was collected
- 507 from GP and incubated in GEP. *k* values followed by different letters are significantly different (non-overlap of 83.4 % CL).
- 508 The R², F and P are estimated from the negative exponential model of Swift et al. (1979).

Treatment	$\frac{k}{k} \pm SE (\times 10^{-3})$	83.4 % CL (× 10 ⁻	$^{-3}$) R ²	F _{1,26}	Р
GP-GP	$1.34\pm0.04\ a$	1.28 - 1.40	0.9666	6716.09	< 0.0001
GEP-GEP	$1.20\pm0.04\ b$	1.14 -1.27	0.9545	5646.09	< 0.0001
GEP-GP	$1.30\pm0.07\ ab$	1.12 - 1.44	0.9149	2382.19	< 0.0001
GP-GEP	$1.04\pm0.02~\text{c}$	0.10 - 1.07	0.9809	8524.80	< 0.0001

Table 3. Contribution (%) of incubation site (Site: GP, grazing paddocks; GEP, grazing exclusion paddocks) and litter quality

511 (Quality: GP-litter, mixed litter collected from grazing paddocks; GEP-litter, mixed litter collected form grazing exclusion

512 paddocks) to litter decomposition and N release.

Parameter	df	Type I SS	Contribution	F	Р
Litter decomposition					
Site	1	110.84	55.51	62.48	< 0.0001
Quality	1	62.06	31.08	34.99	0.0004
Site×Quality	1	12.59	6.30	7.10	0.0286
Error	8	14.19	7.11		
N release					
Site	1	466.25	68.39	57.49	< 0.0001
Quality	1	128.18	18.80	15.81	0.0041
Site×Quality	1	22.41	3.29	2.76	0.1350
Error	8	64.88	9.52		

517 Supplemental table

Species	Palatability	Functional	Species	Palatability	Functional
		group			group
Agrostis hugoniana	Р	GG	Euphorbia esula	UP	FG
Agrostis micrantha	Р	GG	Euphrasia regelii	UP	FG
Allium sikkimense	Р	FG	Gentiana macrophylla*	UP	FG
Astragalus polycladus	Р	LG	Gentiana sino-ornata	UP	FG
Blysmus sinocompressus	Р	SG	Gentianopsis barbata	UP	FG
Carex brunnescens	Р	SG	Geranium wilfordii	UP	FG
Elymus nutans	Р	GG	Halenia elliptica	UP	FG
Festuca sinensis	Р	GG	Lamiophlomis rotata	UP	FG
Gueldenstaedtia	Р	LG	Lancea tibetica	UP	FG
diversifolia					
Kobresia graminifolia	Р	SG	Leontopodium souliei	UP	FG
Kobresia humilis	Р	SG	Ligularia virgaurea	UP	FG
Kobresia kansuensis	Р	SG	Oxytropis kansuensis	UP	LG
Kobresia tibetica	Р	SG	Parnassia palustris	UP	FG
Plantago depressa	Р	FG	Pleurospermum pulszkyi	UP	FG
Poa chalarantha	Р	GG	Polygonum viviparum	UP	FG
Poa poophagorum	Р	GG	Potentilla anserine	UP	FG
Potentilla fragarioides	Р	FG	Ranunculus tanguticus	UP	FG
Roegneria nutans	Р	SG	Rumex patientia	UP	FG
Scirpus pumilus	Р	SG	Saussurea hieracoides	UP	FG
Stipa aliena	Р	GG	Saussurea pachyneura	UP	FG
Anaphalis lactea	UP	FG	Saussurea stella	UP	FG
Anemone obtusiloba	UP	FG	Swertia bimaculata	UP	FG
Anemone rioularis	UP	FG	Thalicturn alpinum	UP	FG
Anemone trullifolia	UP	FG	Thermopsis lanceolata	UP	LG
Arenaria serpyllifolia	UP	FG	Thlaspi arvense	UP	FG
Aster alpinus	UP	FG	Trollius farreri	UP	FG
Cerastium arvense	UP	FG	Veronica eriogyne	UP	FG
Cremanthodium lineare	UP	FG			

Table S1. The list of all species in grazing and grazing exclusion paddocks (Niu et al., 2009, 2010; Wu et al., 2009).

519	Bold font indicates the dominant species. P, palatable species; UP, unpalatable species; GG, group of grass species; SG, group
520	of sedge species; LG, group of leguminous species; FG, group of forbs species; *G. macrophylla presented in GP only.
521	
522	
523	
524	
525	
526	
527	
528	
529	
530	
531	
532	
533	
534	
535	
536	22





- 554 GEP for SOC in 2012 (LSD test: P < 0.05).



558 from grazing paddocks (GP) and incubated in GP; GEP-GEP, mixed litter was collected from grazing exclusion paddocks

559	(GEP) and incubated in GEP; GEP-GP, mixed litter was collected from GEP and incubated in GP; GP-GEP, mixed litter was
560	collected from GP and incubated in GEP. Grey lines under months indicate the mean air temperatures < 0 °C.
561	
562	
563	
564	
565	
566	
567	
568	
569	
570	
571	
572	
573	



583 Supplemental figures



585 Figure S1. Mean temperature (°C) and total precipitation (mm) during experiments.





587 Figure S2. Field sampling methods: (a) half alongside, (b) half along diagonal, (c) two sub-quarters along diagonal, and (d)

