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The relative importance of plant-soil feedbacks for plant-species performance increases with decreasing intensity of herbivory

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Authors contribution:

J.H. conceived the idea, designed the study and performed the experiment; J.H. collected data, with N.K.S., S.S. and M.M.G. provided additional data; J.H., D.P. and J.J. analyzed the data; J.H. led the writing of the manuscript. All authors contributed critically to the draft.

1 **Abstract**

2 Under natural conditions, aboveground herbivory and plant-soil feedbacks (PSFs) are
3 omnipresent interactions strongly affecting individual plant performance. While recent
4 research revealed that aboveground insect herbivory generally impacts the outcome of PSFs,
5 no study tested to what extent the intensity of herbivory affects the outcome. This, however,
6 is essential to estimate the contribution of PSFs to plant performance under natural conditions
7 in the field.

8 Here, we tested PSF effects both with and without exposure to aboveground herbivory for
9 four common grass species in nine grasslands that formed a gradient of aboveground
10 invertebrate herbivory. Without aboveground herbivores, PSFs for each of the four grass
11 species were similar in each of the nine grasslands – both in direction and magnitude. In the
12 presence of herbivores, however, the PSFs differed from those measured under herbivory
13 exclusion, and depended on the intensity of herbivory. At low levels of herbivory, PSFs were
14 similar in the presence and absence of herbivores but differed at high herbivory levels. While
15 PSFs without herbivores remained similar along the gradient of herbivory intensity,
16 increasing herbivory intensity mostly resulted in neutral PSFs in the presence of herbivores.
17 This suggests that the relative importance of PSFs for plant-species performance in grassland
18 communities decreases with increasing intensity of herbivory. Hence, PSFs might be more
19 important for plant performance in ecosystems with low herbivore pressure than in
20 ecosystems with large impacts of insect herbivores.

21

22

23 Key-words: plant-soil feedback, herbivorous insects, field conditions, selective herbivory,
24 nutritional quality

25

26

27 **Introduction**

28 Under natural field conditions the performance (i.e. biomass production) of a plant is
29 influenced by many abiotic and biotic environmental factors that act simultaneously above-
30 and belowground (e.g., Bazzaz 1996, Wardle et al. 2004). Biotic environmental factors such
31 as belowground microbiota and mesofauna as well as aboveground insect herbivory have
32 profound effects on plant performance (Heinze and Joshi 2018).

33 Via litter production, exudation and uptake processes plants induce changes in abiotic and
34 biotic soil properties that, in turn, influence subsequent seedling establishment and plant
35 growth. These plant-soil feedbacks (PSFs; Bever et al. 1997) are typically examined by
36 evaluating the growth of a plant species in response to its own, 'home' (i.e. conspecific) soil
37 compared to growth with other, 'away' (i.e. heterospecific) soil (e.g., Kulmatiski et al. 2008,
38 van der Putten et al. 2013). Besides abiotic soil effects, soil biota are important drivers of
39 PSFs (e.g., DeLong et al. 2019). Since microbial soil biota can function as pathogens or
40 parasites (e.g., pathogenic fungi, bacteria or nematodes) or as mutualists [e.g., arbuscular
41 mycorrhizal fungi (AMF), plant-growth promoting rhizobacteria (PGPR)] (see e.g., van der
42 Heijden et al. 2008, van der Putten et al. 2013, Bever et al. 2015), PSFs can be negative,
43 neutral or positive. Positive PSFs, for example, increase plant-biomass production and thus
44 enhance competitiveness of plant species, whereas negative PSFs weaken their competitive
45 ability. Therefore, PSFs are suggested to influence plant competition and community
46 composition (e.g., Klironomos 2002, Kulmatiski et al. 2008, van der Putten et al. 2013) and
47 have been the subject of intense research (see e.g., Brinkman et al. 2010, Smith-Ramesh and
48 Reynolds 2017). Besides influencing plant biomass, studies on PSFs revealed that soil biota
49 also can influence the nutritional quality of plants (Kos et al. 2015) as well as the composition
50 of secondary metabolites that are involved in herbivory defense (Kostenko et al. 2012,
51 Bezemer et al. 2013). Hence, it is likely that plants growing in home versus away soils,

52 mediated by soil biota, differ in nutritional quality and palatability, which in turn influences
53 aboveground herbivory.

54 Aboveground insect herbivory can affect plant performance directly (e.g., Hulme 1996), but
55 can also influence the composition of plant communities by altering competitive asymmetry
56 between plant species via selective herbivory (Borgström et al. 2016). Therefore, insect
57 herbivory is considered a prominent factor influencing plant species performance and
58 community diversity (Crawley 1989, Branson and Sword 2009). Due to their metabolic
59 requirements, herbivorous insects are known to prefer plants with low carbon (C) to nitrogen
60 (N) ratios [i.e. high N content] and high phosphorus content (Schädler et al. 2003, Berner et
61 al. 2005, Huberty and Denno 2006, Behmer, 2009). Therefore, changes in plant nutritional
62 quality due to soil conditioning in home and away soils (Kos et al. 2015) may alter
63 aboveground herbivore preferences (e.g., Mattson 1980, Massey et al. 2007) and finally the
64 amount of biomass reduction.

65 As calculations of PSFs are mostly based on biomass ratios ('home' vs. 'away'; see Brinkman
66 et al. 2010) it is likely that any disproportional reduction of plant biomass in home relative to
67 away soils by herbivores, due to soil-mediated differences in plant nutritional quality, will
68 influence the results (i.e. outcome) of PSFs. A previous study found that herbivory influences
69 the outcome of PSFs (Heinze and Joshi 2018), but it is currently unknown how the outcome
70 and thus importance of PSFs for plant performance is affected by the strength of this biomass
71 reduction, i.e. by the intensity of herbivory. Thus, 1) if home or away soils increase plant
72 nutritional quality, the resulting increase in aboveground herbivory could mask PSF effects on
73 plant growth and 2) the strength of this masking effect will depend on the intensity of
74 herbivory.

75 Both PSFs and herbivory affect the performance of plants and can act as mechanisms
76 enabling coexistence in plant communities (e.g., i.e. Janzen-Connell-effects; see Petermann et
77 al. 2008; selective herbivory; see Borgström et al. 2016). Understanding the relative impact of

78 PSF-herbivory interactions on plant performance is key to understanding the contexts in
79 which these interactions contribute to coexistence. However, to the best of our knowledge,
80 whether and how the intensity of herbivory influences the outcome of PSFs in the field has
81 never been tested.

82 Most previous studies on PSF-herbivory interactions were performed under controlled
83 greenhouse conditions (e.g., Morriën et al. 2011; Kostenko et al. 2012; Bezemer et al., 2013;
84 but see Heinze and Joshi, 2018). There is, however, high agreement that PSFs should be
85 tested together with herbivory under field conditions in order to gain a comprehensive
86 understanding on the importance of PSFs for plant performance (see van der Putten et al.
87 2016), especially because PSFs differ between greenhouse and field conditions (Heinze et al.
88 2016). Therefore, for the first time we tested PSF (i.e. home vs. away) effects with a
89 standardized comparative PSF pot-experiment in nine grasslands that differed in intensity of
90 aboveground herbivory. We focussed on effects of soil biota (i.e. biotic PSFs) to avoid
91 confounding effects with abiotic soil properties that can also influence nutrient content of
92 plants (e.g., Mattson 1980). In each of the nine grasslands, we manipulated the
93 presence/absence of aboveground herbivorous insects with an herbivore-exclusion treatment.
94 The intensity of herbivory (i.e. the density/abundance of insects) corresponded to the natural
95 condition (i.e., was not experimentally manipulated) to avoid restricting the herbivory effects
96 to one or only a few types of herbivores. To assess the impact of home and away soils on the
97 nutritional quality of plants, we analysed C and N concentrations in roots and shoots of the
98 experimental plants. We hypothesized that: 1) Home and away soils differentially influence
99 plant nutritional quality; 2) As herbivorous insects chose plants selectively consume plants
100 according to their nutritional quality, these home and away soil effects will consequently
101 affect aboveground herbivory by insects; and 3) The outcome of PSFs is influenced by the
102 intensity of herbivory, due to herbivore-induced changes in home vs. away biomass ratios.
103

104 **Material and Methods**

105 *Study region*

106 The comparative PSF experiment was performed in the Biodiversity Exploratories Project
107 (Fischer et al. 2010) in nine grasslands within the Hainich-Dün region (Thuringia, Central
108 Germany). The studied grasslands are located on calcareous mineral soils with high clay
109 content (Fischer et al. 2010).

110

111 *Plant-soil feedback experiment*

112 We selected four common grass species that are widespread within Central Europe (Klötzli et
113 al. 2010): *Arrhenatherum elatius* (L.) J. Presl. et C. Presl., *Anthoxanthum odoratum* L.,
114 *Dactylis glomerata* L. and *Holcus lanatus* L.. All four species are perennial tussock grasses
115 that are frequently found in grasslands within the Biodiversity Exploratories (Heinze et al.
116 2015a,b). Seeds of all four grass species were collected in 2016 in a meadow at a field site of
117 the University of Potsdam (N52° 24' 29.76", E13° 1' 13.74", Brandenburg, Germany). In May
118 2017 seeds of all four species were surface-sterilized for 3 min in 7% sodium hypochlorite
119 solution and subsequently rinsed with sterile water to prevent microbial contaminations.
120 Afterwards, seedlings were germinated on autoclaved sand (5 times within 24 h; 20 min,
121 121°C) in sterile plastic chambers (32 cm × 50 cm × 14 cm; Meyer; Germany) in a
122 greenhouse at the University of Potsdam.

123 We used the “self vs. other” approach (Kulmatiski 2016) to investigate PSF effects for the
124 four grass species. Although this approach does not provide insight into soil mediated
125 interactions between species pairs it focuses on conspecific PSF effects and minimizes the
126 sample size (Kulmatiski 2016). We used species-specific field conditioned rhizosphere soils
127 of all species for our PSF experiment in accordance with the “natural-experiment” approach
128 (Kulmatiski and Kardol 2008). All four species are perennials that form persistent tussocks
129 and therefore generated PSFs over longer time periods. Immediately before the start of the

130 experiment species-specific rhizosphere soils were sampled in the same meadow (size
131 approximately 1 ha) that served as origin for the seeds. For each species we selected 20
132 patches (30 cm x 30 cm), spaced at least 2 m apart from each other, in which the vegetation
133 was solely covered (i.e., 100 %) by the respective species (see Heinze et al. 2016 for
134 description on vegetation structure). Within each patch, we collected 1 L of species-specific
135 soil (top 20 cm) from the rhizosphere and directly adjacent to the rhizosphere following
136 Brandt et al. (2014). As we were interested in general PSF effects rather than within-site
137 variation in PSFs we mixed the 20 replicate soil samples per species to one bulk soil for each
138 species and split in two halves with one half serving as ‘home’ soil (i.e. conspecific soil),
139 whereas the other half was used to create ‘away’ soils (i.e. soils of the remaining
140 heterospecific species) for the other species. Although this mixing procedure decreases
141 variance in plant responses among individual soil samples (Reinhart and Rinella 2016) this
142 procedure was appropriate for our specific research question as we were interested in general
143 (rather than within-site variation of) PSF effects and how they are influenced by the intensity
144 of herbivory. Furthermore, this mixing procedure is reported to produce similar PSFs
145 compared to independent soil samples (see e.g., Kulmatiski 2016, Cahill et al. 2017, Gundale
146 et al. 2019). In total there were eight soils: four home soils (one for every species) and four
147 away soils that each consisted of equal proportions of soils from the three heterospecific
148 species. To reduce potential differences in soil nutrient availability among the eight soils, the
149 soils were inoculated (10%) into an autoclaved soil:sand mixture. The soil:sand mixture
150 consisted of a 1:1 mixture of sieved (mesh size: 5 mm) field soil collected from the same
151 meadow at the field site of the University of Potsdam and purchased sand (grain size: 2 mm;
152 Brun & Böhm; Potsdam, Germany).

153 Pots (Deepots D25L: volume 0.41 L; height 25 cm; diameter 5 cm; Stuewe & Sons; USA)
154 were prepared with an autoclaved fleece strip (3 cm x 25 cm) covering 10 cm of the pots’
155 inside and hanging out 15 cm to enable watering from below. The pots were subsequently

156 filled with the inoculated soils. To prevent cross-contamination between the pots, each pot
157 was placed in a separate plastic cup (volume 0.3 L; height 15.2 cm; diameter 5.9 cm) and
158 received an additional layer (1 cm) of sterilized sand on top.

159 In early June 2017, two-week old similar-sized seedlings of all four species were planted in
160 the prepared pots, one seedling per pot. Each species was grown in pots inoculated with
161 'home' soil or with 'away' soil. Immediately after planting, the pots were moved from the
162 greenhouse to a protected outdoor site near the field study site of the University of Potsdam.
163 There, seedlings were allowed to acclimatize for one week. Seedlings that died during this
164 week were replaced.

165

166 *Herbivore-exclusion treatment*

167 To compare the outcome of PSFs for the four grass species in the presence vs. absence of
168 aboveground insect herbivores we performed a herbivory-exclusion treatment in accordance
169 with Heinze and Joshi (2018). This herbivore-exclusion treatment was established in nine
170 grasslands in the Hainich-Dün region (see below). In each grassland we established two plots
171 (120 cm x 160 cm) that were spaced 80 cm apart. The plots were equipped with cages (length
172 160 cm × width: 120 cm × height 100 cm) that were either completely covered with fly mesh
173 (mesh size: 1.3 mm; Meyer; Germany) or only shaded (i.e. no fly mesh at the lower 50 cm).
174 The fully covered cages excluded herbivorous insects (see MacDonald and Kotanen 2010),
175 whereas the shaded cages allowed aboveground herbivorous insects to reach the experimental
176 plants while providing the same levels of shade and precipitation as the cage treatment (see
177 Heinze and Joshi, 2018). In both plots we removed the sward to slightly (ca.5 cm) sink the
178 prepared pots (in boxes; see below) into the soil and for the fully covered plots to exclude
179 non-developed aboveground herbivorous insects whose eggs might be attached to plants or
180 buried in the soil. In the fully covered plots the fly screen was buried into the soil. One side
181 was prepared as a door to water the plants. The fully covered plots that excluded aboveground

182 herbivorous insects (> 1.3mm) are referred to as ‘– herbivory’, whereas the shaded plots are
183 referred to as ‘+ herbivory’ treatment throughout the manuscript. The plots within each
184 grassland were fenced off (3 m x 3 m) to prevent herbivory by vertebrates as well as
185 disturbances by land-use activities (e.g., mowing).

186

187 *Intensity of aboveground insect herbivory*

188 To test our hypothesis that the intensity of aboveground insect herbivory gradually affects the
189 outcome of PSF effects under natural conditions, we selected nine grasslands along a gradient
190 of land-use intensity. For this gradient it has been shown that land-use intensification
191 influences the abundance and diversity of multiple taxa (Manning et al., 2015), especially
192 herbivorous insects (Simons et al., 2014a,b; Chisté et al., 2016). These land-use effects were
193 found to ultimately affect the severity of aboveground insect herbivory (Börschig et al. 2014;
194 Egerov et al. 2017), which decreases with increasing land-use intensity (Gossner et al. 2014).
195 We used information about past land-use practices (2006–2015), abundance of herbivorous
196 insects (2011–2013) and herbivory on plants measured in the grasslands in 2013 to select nine
197 grasslands along the land-use gradient that are supposed to form a gradient of aboveground
198 insect herbivory (see Online Resource 1: Table S1). The nine grasslands differed in average
199 amount of fertilizer application as well as mowing and grazing intensity, factors that were
200 previously shown to affect abundance and diversity of insect herbivores as well as
201 invertebrate herbivory (Gossner et al. 2014; Simons et al., 2014a,b, Chisté et al., 2016; see
202 Online Resource 1: Table S1).

203 Between and within years land-use practices and their frequency that influences abundance of
204 herbivorous insects, and thus intensity of herbivory, can be highly dynamic and dependent on
205 climate conditions (Blüthgen et al., 2012). Therefore, we also used information regarding
206 planned land use management in 2017 (personal communications from farmers) for the

207 selection of grasslands. We also recorded land-use practices (e.g., mowing events) before and
208 during the experiment (Online Resource 1: Table S2).

209 To test whether land-use intensity affects the intensity of aboveground herbivory in our
210 experiment, we calculated an index of land-use intensity (LUI) according to Blüthgen et al.
211 (2012). This index integrates three components of land use: mean amount of fertilizer (kg N
212 $\text{ha}^{-1} \text{ year}^{-1}$), mean frequency of mowing (number cuttings year^{-1}) and mean intensity of
213 grazing (live-stock units days of grazing $\text{ha}^{-1} \text{ year}^{-1}$) per grassland, that are standardized by
214 the mean of each component per region. The index is square-root transformed, to achieve
215 more evenly distributed values. High values indicate intense land use and vice versa (see also
216 Online Resource 1: Table S1).

217

218 *PSF experiment and herbivore-exclusion treatment along the gradient of herbivory*

219 In mid-June 2017, the planted pots (PSF experiment) were transported to the Hainich-Dün
220 region and positioned in the prepared – and + herbivory plots (herbivore-exclusion treatment)
221 at the nine grasslands (Fig. 1). In each of the nine grasslands, each treatment [herbivory-
222 exclusion and soil treatment (home vs. away)] was replicated nine times for every species,
223 resulting in 1296 pots (9 grasslands x 4 species x 2 soils x 2 herbivory treatments x 9
224 replicates). In the experiment each of the nine grasslands was equipped with exactly the same
225 experimental setup. The planted pots were placed in individual plastic cups (see above) to
226 enable watering from below and were arranged in a randomized block design [i.e. one block
227 contained a single replicate per species and soil treatment (home vs. away)].

228 As we were interested in the effects of aboveground invertebrates (excluding slugs) and as we
229 wanted to exclude direct competition between experimental and neighbouring plants in our
230 experiment, pots and plastic cups were placed in boxes (78 cm × 50 cm × 30 cm). To protect
231 the pots from slug herbivory, these boxes were filled with water (height: 5 cm). In addition to
232 this water barrier, the edges of these boxes are effective barriers against slugs (personal

233 observation J. Heinze). In every grassland, each + herbivory and – herbivory plot contained
234 three boxes, which again contained three blocks of pots each (see Fig. 1). At the beginning of
235 the experiment all plants were watered and all plastic cups underneath every pot were filled
236 with 200 ml water. Every third week the water level in the plastic cups was checked and water
237 was added if necessary.

238

239 *Measurements*

240 We were interested in damage caused by herbivorous insects on the four grass species during
241 the experimental time. We therefore measured herbivory on experimental plants, as these
242 plants were not exposed to destructive land-use practices (like mowing) or slug herbivory. In
243 early September 2017, after 11 weeks of variable invertebrate herbivory intensity exposure,
244 we recorded herbivory on experimental plants. To check whether aboveground herbivory
245 differed between the nine grasslands and the different home vs. away soils, we assessed the
246 damage by aboveground chewing insect herbivores without any further discrimination of
247 feeding guilds. We visually estimated biomass removal (in percent; severity) at ten randomly
248 chosen leaves per individual plant (see e.g., Johnson et al. 2016). Furthermore, in accordance
249 with Russel et al. (2010) for each single experimental plant we also determined the proportion
250 of damaged leaves by counting the number of damaged as well as total leaves (incidence). We
251 used severity and incidence to assess the shoot biomass removal by aboveground insect
252 herbivores for whole experimental plants according to Smith et al. (2005).

253 After herbivory measurements were complete, the pots were brought back to the University of
254 Potsdam where the shoots were harvested and the roots were washed. Shoot and root biomass
255 was dried (shoot 48h, 80°C; root 72h, 70°C) and weighed.

256 To check whether inoculated soils differed in nutrient concentration, we analysed abiotic soil
257 conditions of the eight different inoculated soils (four home soils and four away soils) prior to
258 the experiment according with Heinze et al. (2017). To test whether the different home and

259 away soils affected the nutritional quality in plant shoots and roots, we analysed C and N (see
260 Berner and Law 2016 for C and Cornelissen et al. 2003 for N). As the same soils were used in
261 all of the nine grasslands, we analysed C and N in plant shoots and roots for subsamples of
262 three grasslands. One replicate per species, soil and herbivory treatment was sampled within
263 these three chosen grasslands (see Online Resource 1: Table S1), resulting in 48 samples (4
264 species x 2 soils x 2 herbivory-exclusion treatments x 3 grasslands). Complete shoots and
265 roots were dried at 80°C (48 h), separately ground (Retsch MM200; Germany) and
266 subsequently analysed for C and N concentrations using an elemental analyser (HEKAtech
267 GmbH; Wegberg; Germany; Euro EA 3000).

268

269 *Statistical analysis*

270 All analyses were performed in R version 3.1.2 (R Development Core Team 2014). To
271 account for the split-plot design and the nesting of factors, we analysed the data on shoot-,
272 root- and total biomass, herbivory, PSFs, and C:N ratios of plants with linear mixed effects
273 models using the “nlme” package (Pinheiro et al. 2017). Data on soil nutrients were analysed
274 with linear models, as we tested initial conditions of soils prior to the experiment. Residuals
275 were checked for homogeneity of variance and tested for normality.

276 We used ANOVAs and Tukey HSD tests to check whether the eight inoculated soils [i.e. the
277 sterilized soil:sand mixture (90%) that was inoculated (10%) with the different home and
278 away soils of all four species] differed in abiotic characteristics.

279 To test the first hypothesis that home and away soils differentially affect plant nutritional
280 quality, we performed ANOVAs for N and C concentration as well as C:N ratios in shoots
281 and roots. The ANOVAs included species (*A. elatius*, *A. odoratum*, *D. glomerata*, *H.*
282 *lanatus*), soil treatment (home and away), and herbivory-exclusion treatment (+ herbivory and
283 – herbivory) as well as their interactions as predictor variables. We used “grassland” (three;

284 see “Measurements”) as random factor. Afterwards, differences in N, C and C:N between
285 home and away soils were tested with two sample t-tests for every species.
286
287 To test the second hypothesis, that home and away soils affect aboveground herbivory, and to
288 verify whether intensity of aboveground herbivory differed between the nine grasslands along
289 the land-use intensity gradient we analysed the herbivory (i.e. estimated shoot biomass
290 removal) of experimental plants that were exposed to herbivory (experimental plants in the –
291 herbivory plots showed no damage by herbivores).
292 The ANOVA tested effects and interactions between the predictor variables ‘species (S)’,
293 ‘soil treatment (T)’, ‘herbivory-exclusion treatment (H)’ and ‘land-use intensity (LUI)’ as
294 fixed factors on herbivory, as response variable. We used ‘boxes’ (three) nested in ‘grassland’
295 (nine) as random factors. Additionally, we integrated shoot biomass as co-variable into the
296 model, to test whether herbivory was related to shoot biomass. We used linear regressions to
297 check whether herbivory was related to land-use intensity, for 1) all experimental plants and
298 2) separately for all species.
299 We used average percentage of estimated shoot biomass removal per grassland as a
300 continuous variable in the following analyses to test for the effects of herbivory intensity on
301 PSFs and biomass production (see below). Average percentage of estimated shoot biomass
302 removal is therefore referred to as ‘intensity of herbivory’ throughout the manuscript.
303
304 PSFs were calculated using log biomass ratio of ‘home vs. away’ contrasts, that has the
305 advantage of directly comparing positive and negative feedback effects (see Brinkman et al.
306 2010): $PSF_A = \log(\text{home}_A / \text{away}_A)$; where ‘home_A’ is the biomass of species A with its
307 own soil biota and ‘away_A’ is the biomass of species A with soil biota of the three remaining
308 heterospecific species. PSFs were calculated pairwise per block (i.e. replicate) for shoot, root
309 and total biomass for the + and – herbivory treatments in every grassland.

310

311 To test the third hypothesis, that the intensity of aboveground herbivory influences the
312 outcome of PSFs, we performed ANOVAs using linear mixed effects models. The model
313 included the predictors ‘species (S)’, ‘herbivory-exclusion treatment (H)’ and ‘intensity of
314 herbivory (I)’ (average percentage of estimated shoot biomass removal per grassland) as fixed
315 factors, as well as their interactions and tested their effects on PSFs. We used ‘boxes’ (three
316 per herbivory plot), ‘herbivory plot’ (two per grassland) and ‘grassland’ (nine) as random
317 factors that were nested as follows: boxes nested in herbivory plots and herbivory plot nested
318 in grassland. Whether PSFs for the four species differed within the herbivory treatments along
319 the gradient of herbivory intensity (S x I interaction) was checked by separate ANOVAs for +
320 herbivory and – herbivory. The relationship between intensity of herbivory and PSFs in the
321 two herbivore-exclusion levels were analysed for each species using linear regressions, and
322 differences in slopes were tested with ANOVAs (H x I interaction).

323 The main focus of this study was to investigate effects of herbivory intensity on the outcome
324 of PSFs. However, as PSFs are based on biomass ratios it is likely that data on biomass
325 (shoot) in home vs. away soils in response to herbivory intensity contain valuable
326 information. These results are presented in the supporting information, along with the
327 respective ANOVAs (see Online Resource 1: Table S3; Fig. S1).

328

329 **Results**

330 *Home and away soil effects on plant nutritional quality and herbivory*

331 At the beginning of the experiment the eight inoculated soils neither differed in plant-
332 available nor total nutrient concentrations (Table S4). However, plant shoot N concentration
333 but not C concentration was affected by the different home and away soils for all four species,
334 resulting in different C:N ratios (S x T: shoot N: $F_{3,30} = 10.06$, $P < 0.001$; shoot C:N: $F_{3,30} =$
335 15.15 , $P < 0.001$; Online Resource 1: Table S5a). *A. elatius*, *D. glomerata* and *H. lanatus*

336 showed higher shoot N concentration in away soils, whereas for *A. odoratum* N concentration
337 was highest in shoots when grown on home soils (Fig. 2 a-h). N, C and C:N ratios in roots
338 were not affected by the different soils (Online Resource 1: Table S5b).

339 All four grass species showed differences in aboveground herbivore damage when grown in
340 home vs. away soils (S x T: $F_{3,603} = 13.96$, $P < 0.001$; Online Resource 1: Table S6). *A.*
341 *elatius*, *D. glomerata* and *H. lanatus* showed highest shoot biomass removal in away soils,
342 where their shoots had the highest N concentration (Fig. 2 i, k, l), whereas for *A. odoratum*
343 damage by aboveground herbivores was highest in home soils where its shoots had the
344 highest N concentration (Fig. 2 j).

345

346 *Aboveground herbivory on experimental plants along the gradient of land-use intensity*

347 The estimated shoot biomass removal was highest in less intensively managed grasslands and
348 decreased with increasing land-use intensity ($F_{1,7} = 12.71$; $P = 0.009$; Tables S6; Fig. 3). This
349 pattern of herbivore damage in response to land-use intensity was similar for all four species
350 (S x LUI: $F_{3,603} = 1.74$; $P > 0.05$; Online Resource 1: Table S6; Fig. S2). When grown without
351 herbivores, shoot biomass was similar in all grasslands along the land-use gradient, but
352 decreased with decreasing land-use intensity in the presence of herbivores (see Online
353 Resource 1: Fig. S3).

354

355 *Impact of intensity of aboveground herbivory on PSFs*

356 For all four grass species, the presence of aboveground herbivory influenced the outcome of
357 PSFs for total plants (shoots and roots), but these effects differed among the four species and
358 along the gradient in intensity of herbivory (S x H x I: PSF total: $F_{3,566} = 4.53$, $P = 0.004$; see
359 Online Resource 1: Table S7). Without aboveground herbivores, the four species exhibited
360 different individual PSFs (Fig. 4a-d). *A. elatius* and *H. lanatus* exhibits negative PSFs in
361 home soils (i.e., showed higher biomass production in away soils), *A. odoratum* in contrast

362 responded positively to home soils (i.e. showed positive PSFs), and *D. glomerata* showed
363 neutral PSFs (Fig. 4a-d). Importantly, for all species these PSFs remained similar in
364 magnitude and direction along the gradient of aboveground herbivory intensity (S x I: $F_{3,278} =$
365 0.9, $P > 0.5$; Online Resource 1: Table S7a; Fig. 4a-d). In contrast, when plants were exposed
366 to aboveground herbivory, the direction and magnitude of PSFs for all four species were
367 significantly altered by herbivory intensity (S x I: $F_{3,288} = 8.57$, $P < 0.001$; Online Resource 1:
368 Table S7; Fig. 4a-d). The mostly negative and neutral PSFs of *A. elatius*, *H. lanatus* and *D.*
369 *glomerata* became more positive with increasing intensity of herbivory, whereas for *A.*
370 *odoratum* positive PSFs decreased. Increasing intensity of herbivory increased the difference
371 between PSFs measured with and without herbivores, whereas in the presence of herbivores
372 increasing intensity resulted in mostly neutral PSF effects (Fig. 4a-d).

373

374 **Discussion**

375 The results of our study confirm all three hypotheses and reveal four important findings.
376 First, shoot N concentration of the four grass species was influenced by whether the plants
377 were growing in home or away soils. Second, herbivory by aboveground invertebrate
378 herbivores differed between home and away soils, with all species exhibiting most damage in
379 soils in which their shoots contained highest N concentrations. Third, home and away soils
380 also affected biomass production (i.e. PSFs) of all four species, with highest biomass
381 production in soil in which the species also exhibited highest shoot-N-concentration. Forth
382 and most important, in the presence of herbivores these PSFs changed in magnitude and in
383 direction with increasing intensity of aboveground herbivory, while without herbivores these
384 PSFs remained similar along the gradient of herbivory. These results suggest that that the
385 relative importance of PSFs for individual plant biomass production and thus for the
386 performance in plant communities increases with decreasing intensity of herbivory.

387

388 *Effect of home and away soils on plant quality*

389 In our PSF experiment, all eight home and away soils did not differ in total or plant-available
390 nutrients at initial conditions, an advantage of the inoculation method (Brinkman et al. 2010).
391 Hence, the observed differences in plant nutritional quality (i.e. N and C concentrations) and
392 biomass production of the grass species in the different soils (i.e. home and away) appear to
393 be caused by soil biota.

394 In this study we examined whether the N and C concentrations in plants, chemical plant traits
395 that were broadly overlooked in the past and rarely tested in the context of PSF (see
396 Baxendale et al. 2014, Cortois et al. 2016) were affected when grown in the different soils.
397 We observed that the grass species exhibited highest shoot N concentration in soils where
398 also their biomass production benefitted from soil biota (positive away soil effects for *A.*
399 *elatius* and *H. lanatus* and positive home soil effects for *A. odoratum*). This result is in
400 accordance with findings of Stajković-Srbinović et al. (2016), who showed that inoculation
401 with plant PGPRs enhances both plant biomass and N content in shoots of grass species (see
402 also Baltensperger et al. 1978 and White et al. 2015). In our experiment N concentration was
403 enhanced in shoots in soils where the species benefitted from soil biota but not in roots, a
404 pattern also found in previous inoculation studies with grasses (e.g., Baltensperger et al. 1978,
405 Djonova et al. 2016). Overall, shoots show high turnover rates during growth and thus are
406 sinks for N (Mattson 1980; Xu et al. 2012). This might explain why increased N
407 concentration was confined to shoots.

408

409 *Plant quality and aboveground insect herbivory*

410 In general, due to their high protein content and poor N use efficiency, herbivorous insects
411 need to ingest relatively large amounts of N (Mattson 1980; Bernay and Chapman 1994).
412 Insect herbivores therefore generally prefer to feed on plants with high N content (Berner et
413 al. 2005, Behmer 2009). In our experiment, all four grass species showed highest shoot

414 damage (i.e. estimated shoot biomass removal) caused by aboveground herbivorous insects in
415 soils in which they had highest shoot-N-concentrations. This result is consistent with studies
416 showing that the quantity of herbivore damage is positively related with plant N content
417 (Cebrian and Lartigue 2004, Berner et al. 2005). A reverse pattern was observed for C:N
418 ratios. In line with Schädler et al. (2003) we found all species to have lowest levels of shoot
419 damage in soils where plants had highest C:N ratios. This suggests that beside shoot-N-
420 concentration the palatability is influenced by other physical and/or chemical plant properties
421 (Massey et al. 2007). Soil conditioning can influence other primary and secondary
422 compounds such as amino acids, glycosides, and pyrrolizidine alkaloids (e.g., Kostenko et al.
423 2012, Kos et al. 2015, Zhu et al. 2018) and therefore might affect the palatability of a plant.
424 Furthermore, there are also indications that biotic or abiotic soil characteristics can affect the
425 leaf toughness of plants (Orwin et al. 2010). However, to what extent physical anti-herbivore
426 plant properties are influenced by soil conditioning remains unknown. Although we did not
427 determine specific N-containing secondary metabolites, amino acids or silica content in our
428 study, we nevertheless provide empirical evidence that soil-mediated differences in total N
429 concentration in shoots can strongly affect herbivory by aboveground arthropods. Such
430 specific home and away soil effects on aboveground plant damage and their intensity
431 subsequently affected the outcome of PSFs in our experiment (see below).

432

433 *Intensity of herbivory and its effects on the outcome and importance of PSFs*

434 Increasing intensity of herbivory increased the difference between PSFs measured with and
435 without aboveground insect herbivores. These results confirm previous studies on PSF and
436 herbivory that aboveground herbivores can have negative direct effects on plant growth in the
437 feedback phase (Bezemer et al. 2013). Hence, herbivory has the potential to affect the
438 outcome of PSFs (Heinze and Joshi 2018), most likely due to soil-mediated differences in
439 plant quality. However, more importantly with our present study we were able to show, for

440 the first time, that the intensity of herbivory gradually affected the outcome of PSFs. The
441 change in direction and magnitude of PSFs in response to increasing herbivory intensity
442 mostly resulted in neutral PSFs for the grass species, suggesting that aboveground herbivores
443 reduce the soil-mediated benefits for biomass production depending on herbivore intensity.
444 This is supported by analyses of shoot biomass along the gradient of herbivory intensity:
445 herbivores solely reduced shoot biomass on one specific soil type, namely soil in which the
446 species showed highest shoot N concentration (in away soil for *A. elatius*, *D. glomerata* and
447 *H. lanatus* and in home soil for *A. odoratum* see Online Resource 1: Fig. S1).

448 Grasses are known to have a large and often finely branched root systems with a large surface
449 area and therefore may be more susceptible to root pathogens (Newsham et al. 1995). That *A.*
450 *odoratum* in comparison to the other species exhibited positive PSFs might be due to its high
451 concentrations of coumarin they exudate via roots in comparison to other species (Tava
452 2001). Coumarin was recently found to have a negative effect on soil pathogens but a positive
453 impact on beneficial rhizobacteria (Stringlin et al. 2018) that are important for nutrient uptake
454 and thus plant N concentrations (e.g., Adesemoye et al. 2010). This might also explain the
455 neutral and negative PSFs of the other species, as the away soils they grew in most likely
456 contained coumarin exudates from *A. odoratum*. However, we did not determine soil
457 microbial communities in our experiment. Therefore future studies should use sequence
458 techniques to better understand the role of soil biota in PSF-herbivore interactions.

459 Nevertheless, the findings of our study provide new insights and allow assessments of the
460 importance of PSFs for plant performance in relation to the intensity of herbivory, which has
461 only been considered within a theoretical framework so far (see Smith-Ramesh and Reynolds
462 2017). Based on results of this study we propose that 1) PSFs might be more important for
463 plant performance in ecosystems where the influence of aboveground herbivores is low and 2)
464 as the magnitude and direction of PSFs are altered by herbivory, mostly resulting in neutral

465 PSFs, the importance of PSFs will be changed or overridden by aboveground herbivores in
466 ecosystems where herbivorous insects have a large impact on plant communities (see Fig. 5).
467 In our experiment, species were best supplied with N in soils from which they received the
468 highest biomass gain, indicating that biotic PSFs influence plant performance and quality
469 (Fig. 5a). As larger plants with more biomass are considered to be better competitors in plant-
470 plant interactions (e.g., Aarsen 2015; Heinze et al. 2015a), aboveground herbivores, via
471 specific selection of well-supplied plants (i.e., high N concentration), might prevent the
472 development of dominance structures within plant communities (Fig. 5b). Potential soil-
473 mediated competitive advantages might therefore be attenuated by selective herbivory, thus
474 promoting coexistence in plant communities (see Fig. 5).

475 We suggest that negative density-dependent soil effects (i.e. Janzen-Connell effects) such as
476 negative PSFs for more competitive plant species (*A. elatius*, *D. glomerata* and *H. lanatus*;
477 see Pierce et al. 2017) can act as a stabilizing mechanism (see Chesson 2000) enabling species
478 coexistence in ecosystems with low abundances of herbivorous insects. However, in
479 ecosystems with high abundance of herbivorous insects plant species coexistence might be
480 elevated due to additional equalizing mechanisms, such as selective herbivory that neutralizes
481 soil-mediated competitive advantages, thus influencing the competitive asymmetry between
482 competing plants (Borgström et al. 2016).

483 In our study, we focused on effects of intensity of aboveground insect herbivory on the
484 outcome of PSFs. Soils in our experiment were conditioned with one specific herbivore
485 community (i.e., intensity of herbivory). As the intensity of herbivory is suggested to
486 influence PSFs (Smith-Ramesh and Reynolds 2017) further studies should perform soil
487 conditioning under different intensities of herbivory and investigate these conditioning effects
488 in a feedback phase. Furthermore, we solely excluded insect herbivores > 1.3 mm in our
489 experiment. However, slugs or smaller insect herbivores such as aphids can also have large
490 effects on plant performance (Crawley 1989, Rodriguez and Brown 1998). Therefore, further

491 studies should examine PSF-herbivory interactions by using stepwise exclusion of herbivores
492 and test these interactions across different habitat types as well as with other functional
493 groups to elucidate the relative contribution of herbivores on biomass production and thus
494 their impact on the outcome PSFs.

495

496 *Conclusions*

497 This study is the first to provide empirical evidence that the outcome of PSFs depends on the
498 intensity of aboveground insect herbivory even in our short-term experiment. Soil-mediated
499 differences in plant quality affected herbivory. The intensity of herbivory in turn influenced
500 the shoot biomass in home and away soils for all species and therefore the overall outcome of
501 PSFs. We propose that PSF effects might be more important for plant performance in
502 ecosystems with low insect herbivore pressure compared to ecosystems with high insect
503 herbivory pressure, where soil-mediated advantages for plants might be attenuated via
504 selective herbivory. In addition to the stabilizing effect of negative PSFs, soil-mediated
505 selective herbivory might act as an equalizing mechanism between competing species and
506 might thus promote coexistence in plant communities (Fig. 5). Since under natural conditions
507 both PSFs and herbivory interact and affect plant biomass production over longer time
508 periods PSF-herbivory interactions might be stronger and may change over time. Future
509 studies should therefore test potential changes in these interactions in long-term experiments
510 and assess their impact for competitive outcomes. However, from the present results we
511 suggest that in general the relative importance of PSFs for plant species performance in
512 grassland communities increases with decreasing intensity of herbivory.

513

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528

529 **Conflict of Interest**

530 The authors declare that they have no conflict of interest.

531

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751 FIGURE LEGENDS

752 **Fig. 1** Conceptual figure of the experimental design. To test plant-soil feedback (PSF) effects,
753 four grass species were grown in pots in their ‘home’ and ‘away’ soils. To investigate the
754 effect of herbivory on PSFs, nine replicates of each ‘home’ vs. ‘away’ contrast were exposed
755 to a herbivory treatment in which aboveground insects could either reach the plants (+
756 herbivory plot) or not (– herbivory plot). Within each of the + and - herbivory plots, the nine
757 replicates were arranged in a randomized complete block design and distributed over three
758 boxes (i.e. one box contained 3 replicates/blocks). The boxes were necessary to prevent
759 herbivory by slugs and competition with surrounding plants, and to enable the watering from
760 below. To test whether the intensity of herbivory affect the outcome of PSF effects this set-up
761 (i.e. PSF experiment x herbivory treatment) was installed at nine grasslands that formed a
762 gradient in aboveground herbivory intensity. In total, the experiment contained 1296 plants
763 (4 species x 2 soils x 9 replicates x 2 herbivory treatments x 9 grasslands). For further details
764 see "Material and Methods". Color version of this figure is available online

765

766 **Fig. 2** a-d) Shoot nitrogen (N) concentration, e-h) shoot carbon (C) to nitrogen ratio (C:N) as
767 well as i-l) estimated shoot biomass removal by aboveground insect herbivores of *A. elatius*
768 (left), *A. odoratum* (middle left), *D. glomerata* (middle right) and *H. lanatus* (right) grown in
769 “home” (left bars) and “away” (right bars) soils. Data represent mean \pm SE; with n = 6 for a –
770 h and n = 81 for i – l. Asterisks between bars represent significance: (*) $P < 0.1$; * $P < 0.05$;
771 ** $P < 0.01$; *** $P < 0.001$

772

773 **Fig. 3** Relationship between land-use intensity and estimated shoot biomass removal of all
774 experimental plants exposed to herbivory. Data represent mean \pm SE (n = 72)

775

776 **Fig. 4** Relationship between intensity of herbivory (i.e. average shoot biomass removal by
777 aboveground herbivores per grasslands) and plant-soil feedback [PSF; log total biomass ratio
778 (“home”/”away”)] in the presence (full circles) and absence (open circles) of aboveground
779 herbivorous insects; for a) *Arrhenatherum elatius*, b) *Anthoxanthum odoratum*, c) *Dactylis*
780 *glomerata* and d) *Holcus lanatus*. Statistics shown are interactions of herbivory-exclusion (H)
781 and intensity of herbivory (I) derived from ANOVAs, and for lines derived from linear
782 regressions. Asterisks represent significance: (*) $P < 0.1$; * $P < 0.05$; ** $P < 0.01$; *** $P <$
783 0.001 . Data represent mean \pm SE (n = 9)

784

785 **Fig. 5** Diagram showing how PSF may differently affect plant performance and plant-plant
786 competition in ecosystems with a) low vs. b) high herbivore pressure. In general, soils can
787 have negative or positive effects on nutrient uptake [e.g., nitrogen (N)] resulting in smaller
788 plants with lower nutrient quality in shoots (left plant) or larger and better-supplied plants
789 (right plant). These soil-mediated differences in plant quality and performance might affect
790 competition between competing plants. In ecosystems with low herbivore pressure (a) this
791 soil-mediated advantage in plant growth might be maintained due to marginal damage by
792 insect herbivores resulting in enhanced competition effects for the larger plant. However, in
793 ecosystems with large herbivore pressure (b) effects of insect herbivores might be larger for
794 better-supplied plants. This selective herbivory might dampen the soil-mediated gain of plant
795 growth (grey shadowed) and therefore attenuate competition between plants. Overall, effects
796 from soils influence plant performance and competition, but depending on the intensity and
797 selectivity of herbivory these effects might be influenced by herbivory. The width of arrows
798 and the size of letters indicated the strength or impact of the processes (nutrient uptake,
799 competition, herbivory). Color version of this figure is available online

Fig.1

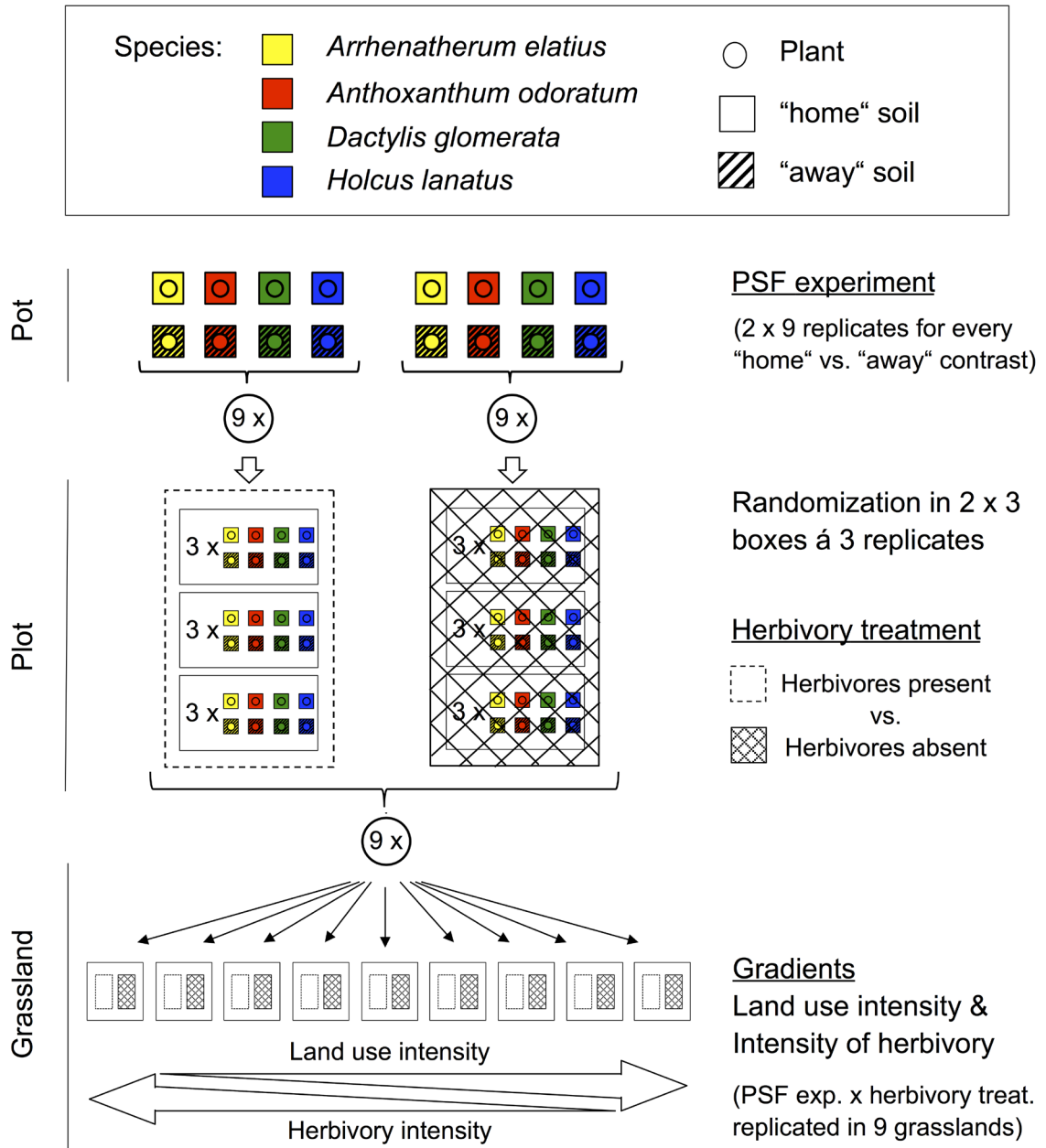


Fig. 2

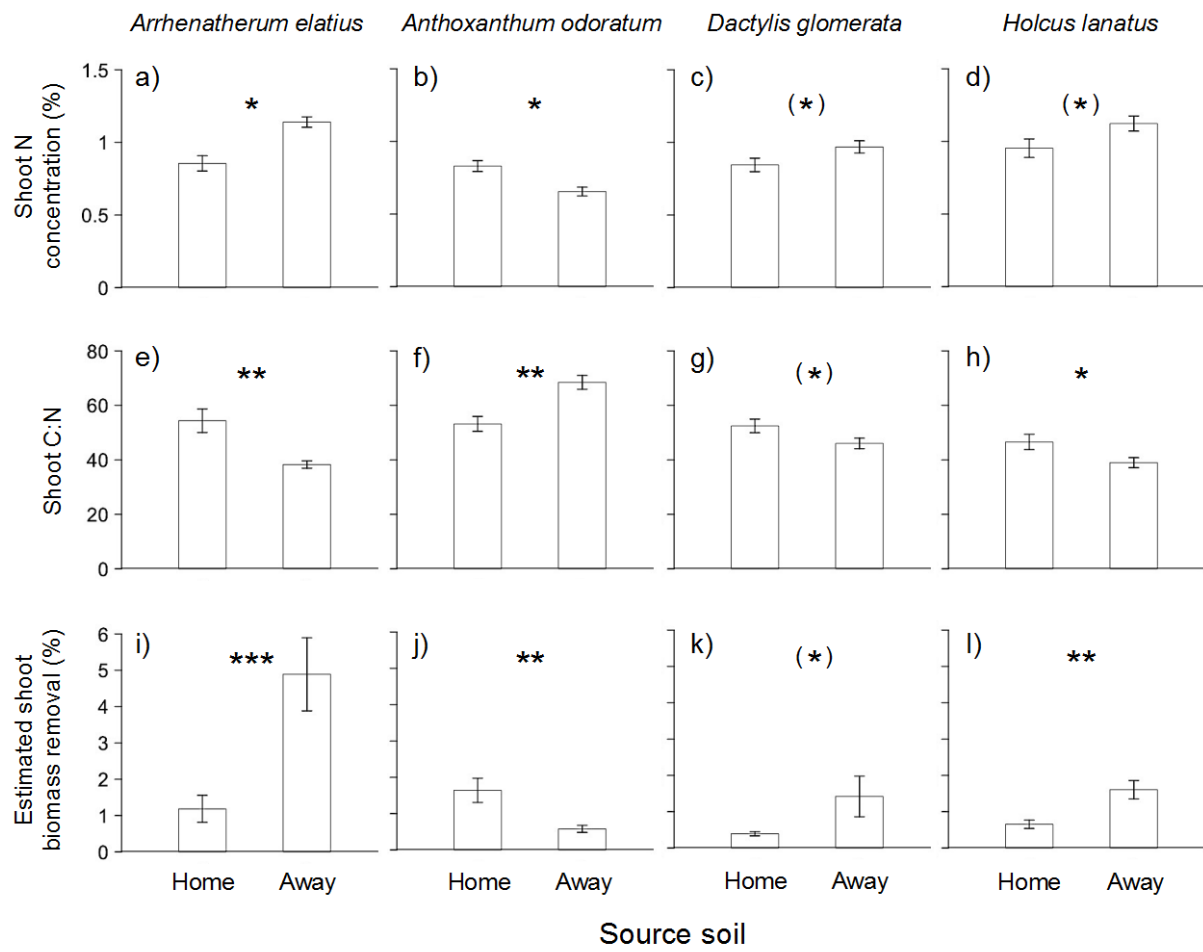


Fig. 3

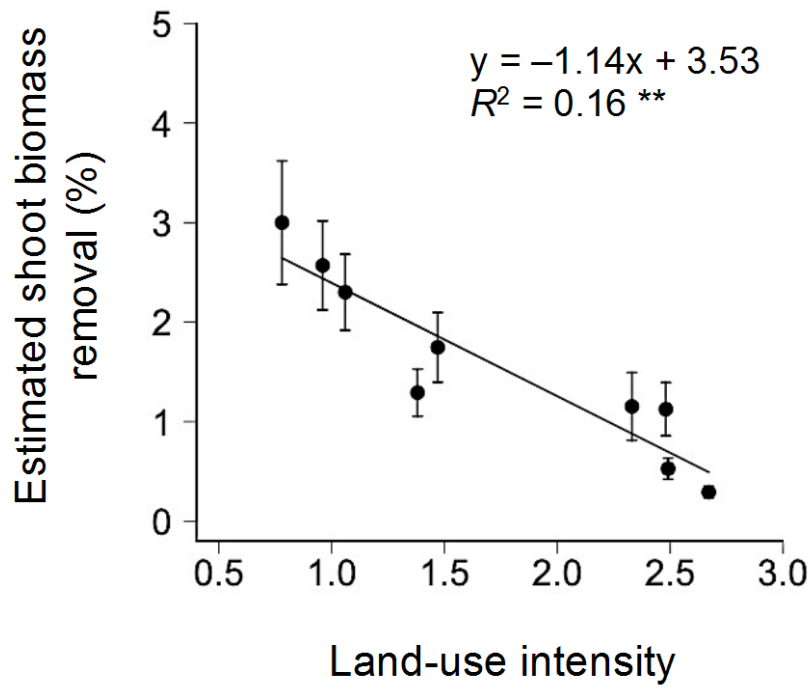


Fig. 4

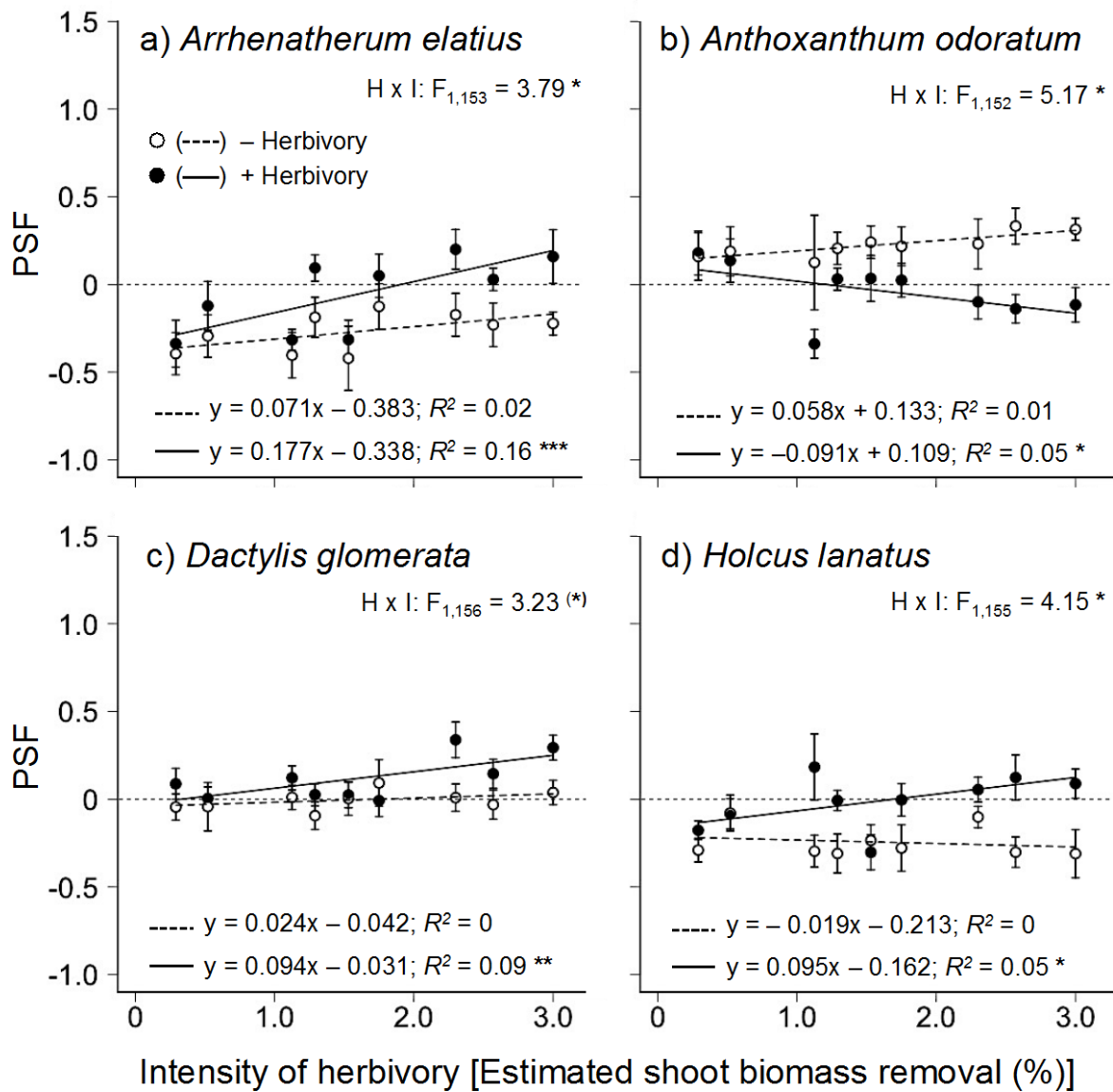
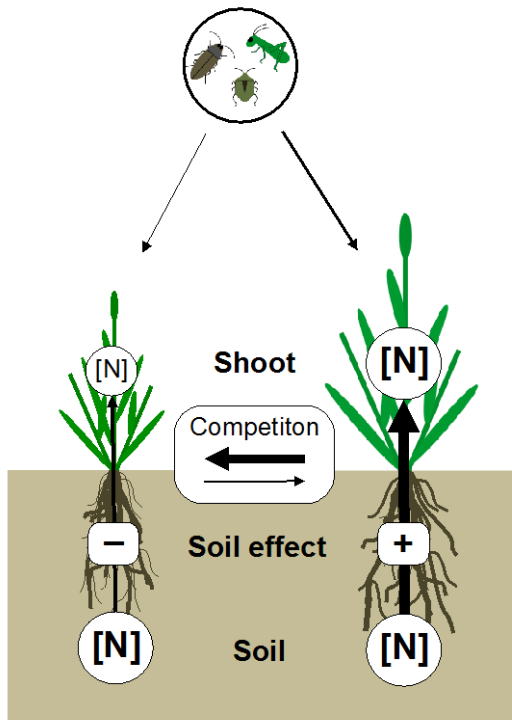


Fig. 5

a) Low herbivore pressure



b) High herbivore pressure

