Effects of reindeer grazing and recovery after cessation of grazing on the ground-dwelling spider assemblage in Finnish Lapland

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The effect of reindeer *Rangifer tarandus* L. grazing on the ground-dwelling spider assemblage in Northern Finland was studied. Changes in species richness, abundance and evenness of spider assemblages were analyzed in relation to changes in vegetation and environmental factors in long term grazed and ungrazed sites as well as sites that had recently switched from grazed to ungrazed and vice versa. Grazing was found to have a significant impact on height and biomass of lichens and other ground vegetation. However, it seemed not to have an impact on the total abundance of spiders. This is likely caused by opposing family and species level responses of spiders to the grazing regime. Lycosid numbers were highest in grazed and linyphiid numbers in ungrazed areas. Lycosidae species richness was highest in ungrazed areas whereas Linyphiidae richness showed no response to grazing. Four Linyphiidae, one Thomisidae and one Lycosidae species showed strong preference for specific treatments. Sites that had recovered from grazing for nine years and the sites that were grazed for the last nine years but were previously ungrazed resembled the long term grazed sites. The results emphasize the importance of reindeer as a modifier of boreal forest ecosystems but the impact of reindeer grazing on spiders seems to be family and species specific. The sites with reversed grazing treatment demonstrate that recovery from strong grazing pressure at these high latitudes is a slow process whereas reindeer can rapidly change the conditions in previously ungrazed sites similar to long term heavily grazed conditions.
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Abstract

The effect of reindeer *Rangifer tarandus* L. grazing on the ground-dwelling spider assemblage in Northern Finland was studied. Changes in species richness, abundance and evenness of spider assemblages were analyzed in relation to changes in vegetation and environmental factors in long term grazed and ungrazed sites as well as sites that had recently switched from grazed to ungrazed and vice versa. Grazing was found to have a significant impact on height and biomass of lichens and other ground vegetation. However, it seemed not to have an impact on the total abundance of spiders. This is likely caused by opposing family and species level responses of spiders to the grazing regime. Lycosid numbers were highest in grazed and linyphiid numbers in ungrazed areas. Lycosidae species richness was highest in ungrazed areas whereas Linyphiidae richness showed no response to grazing. Four Linyphiidae, one Thomisidae and one Lycosidae species showed strong preference for specific treatments. Sites that had recovered from grazing for nine years and the sites that were grazed for the last nine years but were previously ungrazed resembled the long term grazed sites. The results emphasize the importance of reindeer as a modifier of boreal forest ecosystems but the impact of reindeer grazing on spiders seems to be family and species specific. The sites with reversed grazing treatment demonstrate that recovery from strong grazing pressure at these high latitudes is a slow process whereas reindeer can rapidly change the conditions in previously ungrazed sites similar to long term heavily grazed conditions.
Introduction

Wild reindeer have been a natural part of the northern boreal ecosystems in Eurasia since the last ice age, but in northern Fennoscandia and many parts of northern Russia they have been replaced by reindeer husbandry with a sustained high density of semi-domesticated reindeer. Contrary to the reindeer herding practices in Norway and Sweden, in Finland, semi-domesticated reindeer herds can no longer have the traditional long migration between their separate summer and winter ranges. The Finnish reindeer herding area is divided into 57 herding districts defined by legislation (Suominen & Olofsson 2000). These districts are relatively small (about 1 000 to 5 000 km²) compared to the herding areas in Norway and Sweden and prevent long term migration. At present, modern herding practices force reindeer herds in Finland to graze in restricted areas. It is especially detrimental to the lichen dominated winter pastures, if reindeer are forced to use them at summertime since that leads to strong trampling impact. Kumpula et al. 2000 showed that the condition of lichen rich winter pastures is not explained by reindeer density in relation to the total land area of a herding district but by the density of reindeer in relation to the area of lichen ranges. Nowadays with the inclusion of supplementary feeding which prevents natural mortality owing to shortage of food, the grazing pressure in relation to the carrying capacity of pastures can rise excessively high (Kojola et al. 1991, Helle & Kojola 1993; Evans 1996; Kumpula et al. 1997; Suominen & Olofsson 2000). Local herders' cooperatives are trying to save their best winter grazing areas from overgrazing by a system of pasture rotation system (Kumpula et al. 1999), but the success of this is limited.

In the boreal ecosystem, the semi-domesticated reindeer (*Rangifer tarandus tarandus* L.) can be seen as an ecosystem engineer or modifier as well as a disturbance. The ecosystem
engineer concept (Jones et al. 1994; 1997) implies that, in addition to impacts of reindeer through trophic interactions, they control the physical properties of the ecosystem as well. The concept of disturbance, according to the intermediate disturbance hypothesis (IDH) (Connel 1978), implies that the species diversity is at its peak when the level of disturbance is intermediate. IDH has often been applied to the impact of herbivores on vegetation diversity. Both roles stem from grazing behavior and are not mutually exclusive.

Sustained high density of deer has been shown to have a negative impact on vegetation and this impact is further cascades to the whole ecosystem including animal assemblages (Côté et al. 2004). Several studies have demonstrated that the changes caused by large herbivores can have an impact on species composition and abundance of several other animal taxa through various direct and indirect mechanisms (reviewed in Suominen & Danell 2006 and Foster et al. 2014). At the current constantly high densities reindeer herds have substantial influence on the environment; irrespective of whether reindeer considered to be a native part of the ecosystem or a disturbance due to high densities and lack of natural pasture rotation migration. For instance, grazing can change, sustain and create habitats (Putman 1994), accelerate (Kielland et al. 1997; Augustine & McNaughton 1998; Stark et al. 2000; 2002; Olofsson et al. 2004) and even decelerate nutrient flux by affecting the detritus food web via changes in microclimate (Väre et al. 1996; Pastor & Cohen 1997; Augustine & McNaughton 1998; Olofsson et al. 2001). In addition, grazing has an impact on the community structure and species diversity of vegetation (e.g. Väre et al. 1996; Pastor & Naiman 1992; Pastor & Cohen 1997; Milchunas et al. 1998; Suominen 1999) and can influence the post-disturbance succession rate of vegetation (Oksanen et al. 1995; Kielland et al. 1997; Helle et al. 1998). The impacts of reindeer on vegetation and soil properties as well as on the interaction between soils, vegetation and climate change have
been studied a lot in recent years (e.g. Eskelinen et al. 2017; Kaarlejärvi et al. 2017; Ylänne et al. 2017; Egelkraut et al. 2018; Maliniemi et al. 2018). Especially in winter ranges of reindeer in boreal forests, which are typically dry pine forests with *Cladonia*-lichen dominated bottom layer vegetation, it has been shown that the main impact of reindeer is a strong sift from thick lichen carpets to much less bottom and field layer vegetation (e.g. Väre et al. 1996, Suominen & Olofsson 2000, Köster et al. 2013). In addition to biomass and composition of forest floor vegetation this then has cascading impacts on other ecosystem properties such as tree recruitment, fungal communities, litter decomposition (Köster et al. 2013, 2015, Santalahti et al. 2018). Variation observed in vegetation can subsequently mediate the impact of grazing on invertebrate partners in a community (Owen-Smith 1987; DeCalesta 1994; Baines et al. 1994; Bromham et al. 1999; Suominen 1999; Wardle et al. 2001; Suominen et al. 2003; Suominen & Danell 2006; Foster et al. 2014).

The forest floor spider community is greatly influenced by the habitat’s abiotic conditions, such as moisture, light and temperature (Uetz 1991) as well as by the structure of the vegetation and other three dimensional habitat microhabitat features (Colebourn 1974, Hatley & Macmahon 1980, Koponen 1995), and the presence of reindeer can change these conditions and these changes can have an impact on the availability of prey for spiders. Since spiders act both as abundant predators in the invertebrate community and as important prey for numerous forest animals, the changes in spider assemblage can further cascade through the food web via numerous direct and indirect ways.

In this study, we show the impact of reindeer grazing and trampling on the ground-dwelling spider fauna in a special area with four different types of reindeer grazing history. It is also unique in the way that moose (*Alces alces*) is present in both sites with and without reindeer
grazing. Our hypothesis is that reindeer through the changes caused in vegetation height,
structure and biomass as well as in the abiotic conditions has an impact on the assembly structure
of spiders. More specifically, we hypothesize that sites with recently reversed reindeer grazing
status would have intermediate environmental conditions and assembly structure compared long
term grazed and ungrazed sites. Following the IDH assumptions we expected a peak in diversity
of spider assemblages at intermediate grazing treatments compared to long term grazed or
ungrazed pastures.

**Study area and methods**

The study area (Fig. 1) was in eastern Finnish Lapland, in Raja-Jooseppi Inari (68°28’N, 28°
28’E) where semi-domesticated reindeer have been traditionally herded by tradition by Sámi
people for centuries on the Finnish territory. The study site was in dry lichen dominated pine
heath of typical reindeer winter pasture on Lappi reindeer herding district’s eastern boarder. The
district has got an area of 4396 km², and its maximum number of overwintering reindeer is 8000
individuals. On the Russian side of the national border there are no wild or semi-domesticated
reindeer in the area close to the border. The study sites were in the restricted border zone on the
Finnish side of the border. We had a permission for the work from the Finnish Boarder Guard
(permit number 2717/2015).

A reindeer fence along the Finnish-Russian border was erected in the 1940s, and in 1997
a new fence crossing the old one was built in our study area. With time, the old fence was torn
down so that herds could enter untouched pastures. The junction of the new fence and the
position of the old fence forms a cross, where four different grazing regions are distinctly present: i) UGraz - continuously ungrazed from 1940. ii) Graz - continuously grazed from at least 1940. iii) (new ungrazed) NUGraz - ungrazed from 1997, but previously grazed. iii) (new grazed) NGraz - grazed from 1997 but previously ungrazed.

Reindeer fences prevent reindeer from entering the Russian side, where reindeer are locally extinct due to historical causes. Moose, however, occur on both sides and are capable of crossing the fences. This gives a unique opportunity to solely study the grazing effects of reindeer.

**Pitfall trapping**

Spiders were collected using pitfall traps. The traps were plastic cups 170 ml in volume each with a mouth diameter of 70 mm. The killing / preserving fluid was a mix of polypropylene glycol (60%) and tap water, with a hint of detergent added. With pitfall traps it is possible to passively collect a large proportion of actively moving fauna, and being abundant and actively moving creatures, spiders are frequently encountered in pitfalls. A comparison of spider sampling techniques has proved pitfall trapping to maintain its catching efficiency most constantly in time (Churchill & Arthur 1999).

**Trapping design**
A total of 36 trap lines, each consisting of five traps positioned three meters apart, were placed in six plots in the vicinity of the reindeer fence junction in an area of 150 x 500 meters. Each plot included three treatments: the upper region of the fence crossing had Graz, NGraz and UGraz; the lower region had Graz, NU Graz and UGraz. Each treatment within a plot had two trap lines had two trap lines, one five meters from the fence and another 20 meters from the fence. All the trap lines were parallel to the fence. Due to the possible fence effect, i.e. packing of the deer against the fence and thus higher grazing and trampling impact (e.g. Oksanen 1978, Olsofsson et al. 2001), we tested prior to the further analyses whether the lines with different distance to the fence differed from each other (paired t-test or Wilcoxon signed-ranks test).

The trapping period was two months from early June to early August 2005 and due to the short summer of northern boreal regions it practically covered the entire growing season. The emptying interval was one month.

Identification of mature specimens was done to species level. As juveniles often lack distinctive species characters (Norris 1999) they were identified to family level.

Vegetation and environmental factors

Species composition and height of field and bottom layer vegetation was evaluated with a 50 x 50 cm square situated in the middle of each trap line. Dry biomass of the vegetation and soil moisture of soil samples were collected in mid-July, and measured in the laboratory. In each trap line, temperature was measured at the depth of 5 cm in the soil (June 7th 2005, to nearest 0.1 °C).

Since the impacts of reindeer on Cladina lichen dominated forest floor vegetation have been documented in several other studies (e.g. reviews by Suominen & Olofsson 2000, Bernes et al. 2015), and the main focus of the present study is on the spider community, we present the
vegetation results here only as major functional groups pooled over species: lichens, mosses and vascular plants. In our study sites, all other vascular plants than ericaceous dwarf shrubs, were too rare to be statistically tested, and all these shrubs had similar response to treatments. Thus, the result on vascular plants is in fact equal to the response of these shrubs.

Data analysis

Data was analyzed using SAS Enterprise Guide 3.0, EcoSim (Gotelli & Entsminger 2001), and Canoco for Windows 4.5 (ter Braak & Smilauer 1998). Equality of variances was tested with Levene’s test. Normality of residuals was tested with the Shapiro-Wilk test. Variables that were not normally distributed were analyzed with the nonparametric Kruskal-Wallis $\chi^2$-test or Spearman rank correlation (see the table captions). Since many key variables such as spider abundances and plant biomasses could not be analysed with parametric tests the effect of treatments on even those response variables that fulfilled the requirements of parametric tests were also tested with simple one-way ANOVA. In the statistical analyses we used trapping lines as sampling units (observations) i.e. data over the individual traps on each line was combined. The vegetation and physical variables were also measured at trapping line level.

Interactions between environmental variables and species were analyzed using linear regression. The Detrended Correspondence Analysis (DCA) ordination method was used to analyze community structure. The two options "downweighting of rare species" and "trend correction by segments" of Canoco for Windows were turned on.

As the number of species and the number of individuals per sample are often strongly correlated, the number of individuals caught in each trap line was adjusted by rarefaction (see Gotelli & Colwell 2001) to make trap lines of dissimilar catch size comparable.
Results

The distance of the trap line from the fence, i.e. "fence effect", was found to have no effect on spiders and vegetation (paired t-test or Wilcoxon signed-ranks test, p>0.1) and in the subsequent analysis this classifying factor was removed.

General abundance and species numbers

A total of 4225 spider specimens were collected during the whole sampling period. Of these, 3426 specimens were collected from early June till early July and 804 from early July to early August. Mature spiders that were identified to species level numbered 3241 individuals, and 984 specimens were juveniles and therefore identified to family level only (see Supplement for a complete list of specimens). In total 73 spider species were caught, representing 11 families (Table 1).

In terms of specimen numbers (number of spider individuals, abundance), the most common families were wolf spiders (Lycosidae) and sheet weavers (Linyphiidae). Nearly one third of the mature spiders were wolf spider *Pardosa eiseni* or *Alopecosa aculeata* individuals. The twelve most common species comprised 76% of the total number of spiders (Table 2). Eight species were encountered only once.

Vegetation and environmental variables by treatment
Total lichen dry biomass (Fig. 2) and lichen height differed significantly between treatments (df=5, $\chi^2=26.65, p<0.0001$; df=5, $\chi^2=43.18, p<0.0001$) Soil moisture did not differ between treatments, but soil temperature did (df=5, $F=4.09, p=0.0212$) (Fig. 2). The impact of treatment on response variables is shown in Table 3. Dry biomasses of mosses or vascular plants were not significantly affected by grazing (Table 3). Vegetation height differed significantly between treatments for all functional groups i.e. lichens, mosses, and dwarf shrubs (practically all vascular plants in the field layer were dwarf shrubs) all being highest in ungrazed plots (Table 3).

Most common vascular plants were dwarf shrubs ling (Calluna vulgaris), lingonberry (Vaccinium vitis-idaea), crowberry (Empetrum nigrum) and bilberry (V. myrtillus). In addition to those, some graminoids (Poaceae), willow (Salix spp.), bog bilberry (Vaccinium uliginosum) and marsh Labrador tea (Rhododendron tomentosum, ex. Ledum palustre) grew on some sites. Among these plants there were three, which typically characterize more moist or even marshy sites, that were found only or almost exclusively in the long term ungrazed sites. Bog bilberry and Sphagnum-mosses were found only in the ungrazed sites where their occurrence frequencies were 36% and 20% of the studied plots. For marsh Labrador tea the occurrence frequency was 64% in ungrazed and 18% in grazed sites.

Soil moisture and soil temperature (df=16, $F=8.18, p=0.011$), and lichen biomass and soil temperature (df=16, $F=8.93, p=0.009$) correlated negatively, whereas no correlation was found between soil moisture and lichen biomass (Figs 3 a-c).

**Impacts of grazing on spiders**
Abundance

The total number of all spider individuals was similar between treatments, but at the family level there were differences (Table 3), correlations between spider families are shown in Table 4).

Wolf spiders (Lycosidae) were most numerous in long grazed and recently grazed treatments; sheet weavers (Linyphiidae) were most abundant in ungrazed areas, while crab spiders (Thomisidae) showed a more complex yet family specific pattern (Table 3, Fig. 4).

Sheet weavers *Palliduphantes antroniensis* and *Hilaira herniosa* clearly occurred most often in ungrazed areas, whereas the sheet weaver *Agyneta gulosa* and the crab spider *Ozyptila arctica* were as clearly restricted to grazed areas (Fig. 5).

Species richness

Rarefied trap line species richness (n=123 for each trap line) did not differ significantly between treatments (Table 3). The rarefied wolf spider species richness (n=21) was affected by treatments. The highest number of species (6, SD 0.4) was detected in ungrazed sites, the lowest in sites with a long history of grazing (3.5, SD 0.2).

Species richness of Thomisidae was also affected (F=4.27, df=5, p=0.018) by treatments so that the highest richness was found in grazed areas. In sheet weavers the species richness showed no response to grazing.
268 Impact of vegetation and environmental factors on spiders
269
270 With increasing Cladina dry mass and height, the abundance of linyphiids and species richness
271 of lycosids increased but the abundance of thomisids decreased (Table 5). Biomass of vascular
272 plants did not correlate with the abundance or richness of any of the tested spider taxa (Spearman
273 correlation n=36, r<0.27, P>0.11). Moss biomass correlated negatively with Lycosidae richness
274 (Spearman correlation n=36, r=-0.460, P=0.005), but not with other spider variables.
275 Of the most common species Ozyptila arctica (Thomisidae) and Agyneta gulosa
276 (Linyphiidae) decreased in abundance with increasing lichen dry mass and height, whereas
277 Palliduphantes antroniensis (Linyphiidae) was clearly more abundant in areas with thick lichen
278 cover. Also, Hilaira herniosa (Linyphiidae) showed a similar pattern to P. antroniensis. Other
279 common species showed no response to changes in lichen dry mass or height (Table 5). With
280 increasing soil moisture, the abundance of Gnaphosidae and the total species richness of spiders
281 increased significantly. Micrargus herbigradus and Pocadicnemis pumila (Linyphiidae) were
282 more abundant in moister areas, whereas A. gulosa numbers decreased with increasing moisture.
283 The other common species did not respond significantly to soil moisture.
284 Higher soil temperature was connected to a decrease in linyphiid abundance, rarefied
285 lycosid species richness and the total rarefied species richness of the spider assemblage in each
286 treatment (Table 5). M. herbigradus and P. antroniensis decreased in abundance, while A. gulosa
287 abundance was significantly higher in sites with warmer soil.
288
289 Correlations between spider families
290
Correlations between abundances and species richnesses of spider the two most numerous spider families, Lycosidae and Linyphiidae, are shown in Table 4. For the overall spider abundance and richness we used the abundance and richness of all other spider families than the one that was tested for, i.e. for Lycosidae all spiders except lycosids and for Linyphiidae all other spiders than linyphids. The abundance of wolf spider individuals correlates negatively with their own rarefied species richness and positively with the abundance of other spiders and the richness of linyphid spiders. The species richness of Lycosidae spiders correlated positively with the abundance of Linyphiidae and negatively with the richness other spiders and especially that of linyphids. High species richness of linyphids was associated with high abundance of spiders of other families. Earlier, Marusik & Koponen (2002) have found that rich lycosid fauna correlates with high overall species richness in the spider community as well analogous patterns. In our case the opposite was found when the lycosids themselves were excluded from the overall spied richness.

**Multidimensional scaling**

A DCA-ordination visualizing the effect of environmental factors and treatments on spider communities is presented in Fig 6. Treatments are distinctly located on different sides of the graph depending on grazing history. Recently grazed and recently ungrazed treatments are grouped with long grazed areas on the first ordination axis, leaving the ungrazed area at the other end of the gradient. On the second ordination axis, no clear differences are detectable, except a more narrow distribution of ungrazed areas. Soil temperature and soil moisture varies between trap lines fairly independently of lichen dry biomass, but covary with each other to some extent.
The position of spider species on ordination axis 1 and 2 is presented in Fig. 7. Axis 1 mainly represents lichen biomass as in Fig. 6. *Agyneta conigera* and *A. gulosa* (Linyphiidae) are clearly detached from other species, indicating a preference for grazed areas over ungrazed. The lichen rich end of the first axis is dominated by *H. herniosa* (Linyphiidae), *Robertus lividus* (Theridiidae) and especially *P. antroniensis* (Linyphiidae).

On the second axis *Pardosa hyperborea* (Lycosidae) represents a distinct end point. *H. herniosa* and *Minyriolus pusillus* (Linyphiidae) amongst others are found at the opposite end. *Xysticus obscurus* (Thomisidae) is a species that did not react to the first axis, but reacted strongly to the second, while other species were situated somewhere between.

**Discussion**

**Impact of grazing on spiders**

Both family and species level responses to grazing were extensive in lycosids and linyphids, even though the combined abundance or species richness of all spiders did not differ between treatments.

As expected, wolf spiders were more abundant in grazed areas, and the response of sheet weavers was entirely the opposite. Several explanations for this pattern can be found. Wolf spiders chase their prey at ground level, thus making open habitat the optimal foraging ground. Wolf spiders are also relatively large in body size, making it hard to exploit thick, porous lichen carpet as a hunting ground. Also, because this group hunts actively by running, warmer and drier ground may be more suitable for them. Lower numbers of lycosids in ungrazed areas can also be
explained by a lower number of suitable prey (Suominen 1999), in addition to physically more demanding substrate. Linyphiidae spiders catch their prey by using webs, thus their habitat should comprise suitable anchoring sites for web structures. The abundance of web-building spiders has been found to be limited by both the availability of suitable web sites and prey availability (Minoshima et al. 2013, and references therein). Grazing has been shown to decrease both abundance and species richness of web-hunting spiders in Japan and in Kenya, by removing suitable web-building sites (e.g. Miyashita et al. 2004; Warui et al. 2005; Takada et al. 2008; Minoshima et al. 2013). Some studies have suggested that the impact of large herbivores could also be mediated through the abundance of prey (Suominen et al. 2008; Takada et al. 2008) or direct physical disturbance to the spider webs (Foster et al. 2015). In several of these studies the method obtaining spider counts in the field has been based on direct observations of spider webs. This partially circularly reasoning methodology might overemphasize the importance of web building structures for the abundance of web-building spiders. Sweep net sampling in an area close to the present study site during the same summer did not show significant differences in the amount of insect prey (see supplementary material). Since the study area is typical winter pasture for reindeer it is unlikely that there would be substantial direct disturbance of the webs by reindeer during summer. A lack of suitable construction sites for webs and possibly also the different microclimate conditions (i.e. humidity and temperature) seem feasible explanations in this study.

Thomisidae abundance in both intermediate treatments strongly reflected the grazing history of the intermediate treatments. The observed similarity between recently ungrazed and continuously grazed sites could be explained by the slow recovery of vegetation. However, as
Species richness of spiders

The response of rarefied wolf spider species richness to treatments was as expected. In the presumably favourable sites species richness was low and abundance high, whereas in low quality sites species richness was high and abundance low. This is well in line with the competitive exclusion theory, according to which strong competitors in good quality habitats can outcompete subdominants with a similar ecological niche, hence leading to lower species richness (Gause 1934; Hardin 1960). In suboptimal, ungrazed habitats, competitive exclusion is not effective as other factors than inter-species competition are more limiting. A high number of species could be seen as a sign of a more heterogeneous habitat, where more species can live together. However, a monotonous thick and dense carpet of lichen is not likely to be more heterogeneous than grazed areas or provide more suitable environments for ground dwelling wolf spiders.

Species richness within Linyphiidae family did not differ between treatments. As linyphiids are highly efficient in their dispersal capacities (ballooning method), they can respond efficiently to environmental changes. Yet, due to wind-driven and rather erratic dispersal, linyphiids are also often dispersed to suboptimal habitats, hence the species present in one spot might reflect chance only. Linyphiid abundance, however, can tell more about the actual quality of the habitat.
Spider species responses

* A. gulosa and A. conigera, linyphid species found in grazed areas in high numbers, typically prefer areas in an early successional phase just after disturbance and are even considered pioneer species (Koponen 2004; Koponen & Koneva 2006). Occurrence of pioneer species in an area grazed continuously from at least the 1940s shows that grazing pressure is high enough to keep an area constantly in its early stage of succession. Linyphiidae spiders. *P. antroniensis* and *H. herniosa* are typical northern boreal bottom layer species (Koponen 1999), and here strictly occurred in great abundance in ungrazed, reindeer-free areas.

Of the species that are not web-spinning, *Xysticus obscurus* (Thomisidae) was restricted to humid areas, but was rather infrequent in abundance and was previously found in multiple habitats (Koponen 1977). *P. hyperborea* also indicated predominantly lichen rich, humid areas, but was also highly abundant in one site with low lichen biomass. The crab spider *O. arctica* was exceptional in its abundance by being most common in long term grazed sites as well as in newly ungrazed site.

Linear regression revealed the affinity of *Micrargus herbigradus* (Linyphiidae) to cool and moist sites. However, its occurrence was not affected by lichen biomass.

DCA-ordination did not group species into families but instead showed greatly differing environmental affinities, even within a genus. Hence, use of higher taxa than species to describe changes in environmental attributes can be misleading if the responses of the species contradict each other and the researcher is not utterly aware of the ecology of the focal species.
Importance of microclimate for spiders

In addition to causing differences in lichen biomass and height as well as other direct and indirect impacts on vegetation, grazing induces differences in microclimate, which may also explain the observed patterns in spider fauna (Koponen et al. 1975; Samu et al. 1999; Frick et al. 2007). Large, actively moving wolf spiders require a warmer habitat than small, relatively motionless web-hunting spiders. Due to their small body size, linyphiids are also more vulnerable to drying out. Inside and on top of the spongy, absorbent lichen carpet, the risk of desiccation is probably less important. Also, Riechert and Tracy (1975) showed that by building webs in physically optimal sites instead of sites maximizing prey catch, web-building spiders could significantly increase their fitness.

Impact of grazing on vegetation and environmental variables

As expected and shown by numerous previous studies (e.g. Väre et al. 1996, Suominen & Olofsson 2000, Köster et al. 2013), the main impact of reindeer on vegetation was a reduction in reindeer lichens. Reindeer significantly reduced both the biomass and the height of their main winter food, the Cladina lichens, that otherwise dominate ground layer vegetation in these forests. Grazing also raised the soil temperature in summer by removing the isolating and reflecting layer of white thick lichen. This differs from the results of the study by Köster et al. (2015). They did not find significant difference in soil temperature in fairly similar conditions not so far away from our study site. We did not detect significant differences in soil moisture
between treatments, and this is similar to the results of Köster et al. (2015). This may have been a result of small-scale topographical differences at the study site. However, occurrence of typical peatland species on the sandy soil of our study area like the spider *Pardosa sphagnicola* and plants like peat moss (*Sphagnum* spp.) and bog bilberry in the long term ungrazed area implies that there are actual differences in moisture levels between treatments. Also *Pardosa hyperborea*, being relatively abundant in our samples in ungrazed areas, is known to prefer moist conditions (Koponen 1977), even to the extent of being one of the dominant species in peatlands of the boreal zone (Koponen 2002).

The DCA ordination of the spider community split the treatments along two gradients: a first ordination axis that mainly represented lichen biomass and a second axis representing soil moisture and temperature. The latter, however, could not divide the species as clearly as the first. Most of the long grazed sites, along with intermediate treatments, were located at a relatively dry section of the gradient. The first axis clearly separated the ungrazed sites from the others, while variation within axis two was to some extent greater. Since the treatments with a recently altered grazing regime were grouped with long grazed treatments, eight years of protection from grazing appears not to be a long enough time to alter the composition of spider assemblage at the study site. Furthermore, areas with only a decade of grazing already resembled the long grazed areas in multiple ways.

Next to non-existent differences in vegetation between continuously grazed areas and areas with a decade of recovery from grazing were in contrast to our hypothesis emphasizing the importance of the level of grazing. Generally, harsh climatic conditions in northern boreal forests, leading to a slow succession rate and therefore minute detectable changes in vegetation in the time span of only one decade, may explain some of the findings. The study and modelling
by Kumpula et al. (2000) showed that if lichen has been grazed and trampled to extremely low
level its recovery is much lower than that of more moderately depleted lichen stand. This might
well be the case in our study area as well. Also, changing interspecific competition in the plant
community due to changes in microclimate may slow down recovery from grazing (Olofsson
2006). Patterns in Thomisidae abundance, however, imply that there might be real differences
also between long grazed and intermediate treatments.

Conclusion

Based on our results, we could not find evidence of the hypothesized increase in spider diversity
under intermediate grazing impact by reindeer in lichen dominated dry pine forests in the
northern boreal zone, but the hypothesis cannot be rejected based on our data. Our data, of
course, comes from only one year and a limited area, and as such is has some limitations.
However, it seems rather that instead of being intermediate in relation reindeer impact the
treatments where grazing impact had been reversed eight years ago were fairly similar to long
term grazed habitats.

Focal species, sexes or even life stages can experience the same disturbance in dissimilar
ways. However, in sites where one species was highly abundant, the species richness within that
family was always lower than the species richness in sites of more even species distribution. This
suggests active competitive exclusion.
Even though Linyphiidae and Lycosidae spiders responded to reindeer grazing in the same way as to clear-cut forest management in Pajunen et al. (1995), oversimplification of the detrimental effects of heavy grazing on the forest floor spider community should be avoided. The findings only support the existence of substantial habitat quality differences between grazed and ungrazed areas for spiders. Furthermore, it is vital to remember that foraging habits and thus the impact on vegetation also differ between large herbivore grazers (Huntly 1991; Adler et al. 2001). The distinct differences in feeding habits of reindeer between seasons can lead to completely different impacts of the same herbivore species, through winter grazing on ground lichens and summer grazing on grasses and deciduous leaves. Moreover, even though the responses to treatments showed clear family-level habitat preferences among linyphiids and lycosids, certain species within these families showed opposite responses. Thus, oversimplification of family-level habitat preferences should also be avoided.

What our results clearly demonstrate is an extremely slow recovery rate of the community after heavy grazing, which extends from the lichens even to invertebrates such as spiders. Both of our intermediate treatments with eight years of reindeer exclusion or eight years of grazing were strikingly similar to each other and to the long term grazed treatment. Thus we could not see signs of noticeable recovery from heavy grazing after eight years of protection from grazing, but on the other hand formerly ungrazed habitat was very similar to habitat that had been grazed for decades after eight years of grazing. The recovery of lichen pasture after heavy grazing is a long process (Olofsson 2006; Hansen et al. 2007) and not only for slow-growing lichens, but also for other vegetation, the physical environment, and for faunistic components like the spiders studied here.
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**Figure and Table captions**
Figure 1. Map of Finland showing the study area.

Figure 2. The differences between treatments in soil temperature (°C at the depth of 5 cm), soil moisture (top layer of mineral soil), and total lichen biomass (dry mass g/0.25 m²) (mean ± SD). The treatments are Graz – long term grazed area, UGraz – long term ungrazed, NGraz – grazed last 9 years, but previously ungrazed, NUGraz – ungrazed last 9 years, but previously grazed (the number (1) in treatments Graz and UGraz refers to sites that act as a reference to the NGraz treatment and number (2) to those that act as reference to the NUGraz treatment).

Figure 3. The relationship between lichen biomass (dry mass g/0.25 m²), soil moisture (% top layer of mineral soil), and soil temperature (°C at the depth of 5 cm). Statistically significant linear regressions are included into the figures.

Figure 4. Pooled family level abundances of spider individuals of the three most abundant spider families in different treatments (mean number of individuals per trapping line ± SD). For the abbreviations of treatments see the legend of figure 2.

Figure 5. Abundances of spider individuals of four common species between different treatments (mean number of individuals per trapping line ± SD). For the abbreviations of treatments see the legend of figure 2.

Figure 6. DCA-ordination visualizing the effect of environmental factors and treatments on spider communities. The symbols for treatments: long term grazed = □, sites where grazing
status changed in 1997 = ◊, long term ungrazed = ○. The axes 1 and 2 explained 33.4% and 16.7%
% of the variation (relative eigenvalue portions 0.40 and 0.16 respectively).

Figure 7. The location of spider species on DCA-ordination axes 1 and 2. Only species with
more than 20% weight on the ordination are shown. The axes 1 and 2 explained 14.5% and 9.3%
% of the variation in the species data (relative eigenvalue portions 0.198 and 0.127 respectively).

Table 1. List of spider species found in the study.

Table 2. Twelve most common species and their specimen numbers.

Table 3. Impact of treatment on response variables (mean and SD, or estimate with 95%
confidence limits). Variables that fulfill the assumptions for a parametric test were tested with
ANOVA and the F-test (DF= 5). Spider abundances and lichen biomass were tested with the
non-parametric Kruskal-Wallis χ²-test (DF=5). Species richness for pooled families, Lycosidae
and Linyphiidae, are rarefied values. Due to the low number of species and specimens, species
richnesses of Thomisiidae and Gnaphosidae are absolute values without rarefaction. Site-level
richness (α-diversity) is the rarefied number of species per trapping line. γ-diversity is the
rarefied number of species per treatment (with 95% confidence limit). Bold font indicates
variables with statistically significant (p<0.05) treatment impact.
Table 4. Spearman rank correlations of the abundances (n) and species richnesses (S) of spider families. (* = p<0.05, **p<0.01)(n=18). Others n and Others S mean the abundance and richness respectively of all other spider families than the one tested against.

Table 5. The responses of spider family species richnesses (s) and abundances (n), as well as the abundances of the most numerous species to environmental variables. Linear regression results are also included (* = p<0.05, **p<0.01)(df=16). S = number of species, n = number of specimen.
Figure 1

Map of Finland showing the study area.
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Figure 2

The differences between treatments in soil temperature (°C at the depth of 5 cm), soil moisture (top layer of mineral soil), and total lichen biomass (dry mass g/0.25 m²) (mean ± SD).

The treatments are Graz – long term grazed area, Ugraz – long term ungrazed, NGraz – grazed last 9 years, but previously ungrazed, NUgraz – ungrazed last 9 years, but previously grazed (the number (1) in treatments Graz and Ugraz refers to sites that act as a reference to the NGraz treatment and number (2) to those that act as reference to the NUgraz treatment).
Figure 3

The relationship between lichen biomass (dry mass g/0.25 m²), soil moisture (% top layer of mineral soil), and soil temperature (°C at the depth of 5 cm).

Statistically significant linear regressions are included into the figures.
Figure 4

Pooled family level abundances of spider individuals of the three most abundant spider families in different treatments (mean number of individuals per trapping line ± SD).

For the abbreviations of treatments see the legend of figure 3.
Figure 5

Abundances of spider individuals of four common species between different treatments (mean number of individuals per trapping line ± SD).

For the abbreviations of treatments see the legend of figure 3.
Figure 6

DCA-ordination visualizing the effect of environmental factors and treatments on spider communities.

The symbols for treatments: long term grazed = □, sites where grazing status changed in 1997 = ◊, long term ungrazed = ○. The axes 1 and 2 explained 33.4 % and 16.7 % of the variation (relative eigenvalue portions 0.40 and 0.16 respectively).
Figure 7

The location of spider species on DCA-ordination axes 1 and 2.

Only species with more than 20% weight on the ordination are shown. The axes 1 and 2 explained 14.5 % and 9.3 % of the variation in the species data (relative eigenvalue portions 0.198 and 0.127 respectively).
Table 1 (on next page)

List of species found in the study.
| 1  | Araneidae          |
|----|-------------------|
| 2  | Hypsosinga albovittata (Westring, 1851) |
| 3  | Cercidia prominens (Westring, 1851) |
| 4  | Dictynidae         |
| 5  | Hackmania prominula (Tullgren, 1948) |
| 6  | Gnaphosidae        |
| 7  | Gnaphosa lapponum (L. Koch, 1866) |
| 8  | Gnaphosa montana (L. Koch, 1866) |
| 9  | Gnaphosa muscorum (L. Koch, 1866) |
| 10 | Haplodrassus signifer (C. L. Koch, 1839) |
| 11 | Haplodrassus soerensen (Strand, 1900) |
| 12 | Micaria alpina (L. Koch, 1872) |
| 13 | Zelotes subterraneus (C. L. Koch, 1833) |
| 14 | Hahnidae           |
| 15 | Hahnia ononidum (Simon, 1875) |
| 16 | Linyphiidae        |
| 17 | Agyneta cauta (O. P.-Cambridge, 1902) |
| 18 | Agyneta conigera (O. P.-Cambridge, 1863) |
| 19 | Agyneta gulos (L. Koch, 1869) |
| 20 | Agyneta subtilis (O. P.-Cambridge, 1863) |
| 21 | Agyneta trifurcata (Hippa & Oksala, 1985) |
| 22 | Bolephyphantes index (Thorell, 1856) |
| 23 | Ceratinella brevipes (Westring, 1851) |
| 24 | Centromerus arcanus (O. P.-Cambridge, 1873) |
| 25 | Cnephalocotes obscurus (Blackwall, 1834) |
| 26 | Decipiphantes decipiens (L. Koch, 1879) |
| 27 | Diplocentria bidentata (Emerton, 1882) |
| 28 | Diplocentria rectangulata (Emerton, 1915) |
| 29 | Hilaira herniosa (Thorell, 1875) |
| 30 | Macrargus rufus (Wider, 1834) |
| 31 | Maso sundevalli (Westring, 1851) |
| 32 | Micrargus herbigradus (Blackwall, 1854) |
| 33 | Minyriolus pusillus (Wider, 1834) |
| 34 | Moebelia penicillata (Westring, 1851) |
| 35 | Mughiphantes cornutus (Schenkel, 1927) |
| 36 | Neriene clathrata (Sundevall, 1830) |
| 37 | Oreoneta sinuosa (Tullgren, 1955) |
| 38 | Palliduphantes antroniensis (Schenkel, 1933) |
| 39 | Pocadinemis pumila (Blackwall, 1841) |
| 40 | Porromma pallidum (Jackson, 1913) |
| 41 | Scandichrestus tenuis (Holm, 1943) |
| 42 | Semljicola latus (Holm, 1939) |
| 43 | Sisicus apertus (Holm, 1939) |
| 44 | Tapinocyba pallens (O. P.-Cambridge, 1872) |
| 45 | Tenuiphantes alacris (Blackwall, 1853) |
| 46 | Tenuiphantes mengei (Kulczynski, 1887) |
| 47 | Tenuiphantes tenebricola (Wider, 1834) |
| 48 | Tobioplodius arcuatus (Tullgren, 1955) |
| 49 | Tobiplus diversus (L. Koch, 1879) |
| 50 | Walckenaeria antica (Wider, 1834) |
| 51 | Walckenaeria capito (Westring, 1861) |
| 52 | Walckenaeria cuspidata (Blackwall, 1833) |
| 53 | Walckenaeria dysderoides (Wider, 1834) |
| 54 | Walckenaeria karpinskii (O. P.-Cambridge, 1873) |
| 55 | Walckenaeria obtusa (Blackwall, 1836) |
| 56 | Walckenaeria unicornis (O. P.-Cambridge, 1861) |
| 57 | Zornella cultrigera (L. Koch, 1879) |
| 58 | Liocranidae         |
| 59 | Agroeca proxima (O. P.-Cambridge, 1871) |
| 60 | Lycosidae           |
| 61 | Acantholycosa lignaria (Clerck, 1757) |
| 62 | Alopecosa aculeata (Clerck, 1757) |
| 63 | Alopecosa pinetorum (Thorell, 1856) |
| 64 | Pardosa eiseni (Thorell, 1875) |
| 65 | Pardosa hyperborea (Thorell, 1872) |
| 66 | Pardosa lasciva (L. Koch, 1879) |
| 67 | Pardosa lugubris (Walckenaer, 1802) |
| 68 | Pardosa palustris (Linnaeus, 1758) |
| 69 | Pardosa sphagnicola (Dahl, 1908) |
| 70 | Philodromidae       |
| 71 | Thanatus formicinus (Clerck, 1757) |
| 72 | Salticidae          |
| 73 | Evarcha falcata (Clerck, 1757) |
| 74 | Neon reticulatus (Blackwall, 1853) |
| 75 | Therididae          |
| 76 | Robertus lividus (Blackwall, 1836) |
92  *Theonoe minutissima* (O. P.-Cambridge, 1879)

95  **Thomisidae**

96  *Ozyptila arctica* (Kulczynski, 1908)

97  *Xysticus audax* (Schrank, 1803)

98  *Xysticus luctuosus* (Blackwall, 1836)

99  *Xysticus obscurus* (Collett, 1877)
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Table 2 (on next page)

Twelve most common species and their specimen numbers.
| Species       | Pardosa    | Alopecosa    | Pardosa    | Ozyptila    | Hilaira    | Pocadienemis |
|---------------|------------|--------------|------------|-------------|------------|--------------|
| n             | 622        | 344          | 276        | 235         | 200        | 159          |

| Species       | Agyneta    | Pardosa      | Walckenaeria | Palliduphantes | Hahnia    | Micrargus   |
|---------------|------------|--------------|--------------|----------------|-----------|-------------|
| n             | 148        | 118          | 117          | 103            | 80        | 67          |
Table 3 (on next page)

Impact of treatment on response variables (mean and SD, or estimate with 95% confidence limits).

Variables that fulfill the assumptions for a parametric test were tested with ANOVA and the F-test (DF= 5). Spider abundances and lichen biomass were tested with the non-parametric Kruskal-Wallis χ²-test (DF=5). Species richness for pooled families, Lycosidae and Linyphiidae, are rarefied values. Due to the low number of species and specimens, species richnesses of Thomisiidae and Gnaphosidae are absolute values without rarefaction. Due to the same reason the Pielou’s evenness index J’ could not be calculated for Gnaphosidae. Site-level richness (α-diversity) is the rarefied number of species per trapping line. γ-diversity is the rarefied number of species per treatment (with 95% confidence limit). JK-species no. is the first order jackknife estimate of the total number of species in each treatment (with 95% confidence limits). Bold font indicates variables with statistically significant (p<0.05) treatment impact.
| SPIDER RICHNES | Mean (SD) | UGraz (1) | Graz (1) | Graz (2) | NU Graz | NGraz | F(χ²) | P   |
|----------------|-----------|-----------|----------|----------|---------|-------|-------|-----|
| α              | 26.8 (4.1)| 28.5 (2.3)| 23.13 (2.1)| 27.12 (5.9)| 28.8 (6.0)| 21.4 (2.8)| 1.53 | 0.251 |
| γ              | 41.5 (3.1)| 49.3 (1.5)| 38.7 (4.1)| 47.0 (0.2)| 46.0 (0.5)| 36.9 (2.5)| -   | -    |
| Lycosidae      | 4.1 (0.2)| 6.1 (0.4)| 3.6 (0.3)| 4.5 (0.7)| 4.7 (1.5)| 3.6 (0.8)| 4.37 | 0.017 |
| Linyphiidae    | 11.5 (1.4)| 9.8 (1.4)| 11.2 (1.7)| 10.2 (3.1)| 11.2 (1.7)| 12.6 (2.0)| 0.72 | 0.623 |
| Thomisidae     | 1.7 (0.6)| 2.3 (0.6)| 2.3 (0.6)| 3.3 (0.6)| 2.3 (0.6)| 1.3 (0.6)| 4.27 | 0.018 |
| Gnaphosidae    | 2.6 (0.6)| 1.3 (0.6)| 2.6 (0.6)| 3.0 (1.7)| 3.3 (1.5)| 3.0 (1.0)| 1.2  | 0.366 |

| SPIDER ABUNDANCE | Mean (SD) | UGraz (1) | Graz (1) | Graz (2) | NU Graz | NGraz | F(χ²) | P |
|-------------------|-----------|-----------|----------|----------|---------|-------|-------|---|
| Total             | 203.7 (66.0)| 166.7 (37.3)| 225.3 (41.4)| 135.6 (14.2)| 144.6 (13.7)| 206.0 (121)| 5.63 | 0.344 |
| Lycosidae         | 93.3 (55.9)| 22.3 (1.2)| 123.3 (35.5)| 47.7 (9.2)| 40.0 (16.5)| 147.7 (117.4)| 12.09 | 0.034 |
| Linyphiidae       | 81.7 (3.8)| 120.2 (35.3)| 58.0 (3.5)| 56.3 (9.3)| 67 (7.2)| 35.7 (10.0)| 15.27 | 0.009 |
| Thomisidae        | 8.7 (7.6)| 10.0 (6.1)| 26.0 (4.0)| 22.3 (5.5)| 22.7 (3.1)| 7.3 (2.1)| 12.74 | 0.026 |
| Gnaphosidae       | 9.0 (2.6)| 2.0 (1.0)| 6.3 (3.5)| 4.7 (3.1)| 6.3 (2.5)| 7.7 (6.7)| 7.45  | 0.189 |
| P. antroniensis   | 10.7 (5.3)| 22 (7.5)| 0 (0)| 0.7 (0.6)| 0.7 (0.6)| 0.3 (0.6)| 24.41 | <0.0001 |
| H. herniosa      | 10 (2.4)| 47.3 (27.1)| 1.7 (2.1)| 0.7 (1.2)| 5.3 (2.1)| 1.7 (0.6)| 7.88  | 0.002 |
| O. arctica       | 8 (5.3)| 7.7 (5.5)| 22.7 (2.9)| 15.3 (8.1)| 17.7 (2.9)| 7 (1.7)| 5.24  | 0.009 |
| A. gulosa        | 5.3 (6.8)| 0.7 (1.2)| 11.7 (6.7)| 17 (13.1)| 7.3 (6.8)| 7.3 (8.5)| 1.46  | 0.273 |

| ENV.VAR. | Mean (SD) | UGraz (1) | Graz (1) | Graz (2) | NU Graz | NGraz | F(χ²) | P |
|-----------|-----------|-----------|----------|----------|---------|-------|-------|---|
| Soil °C   | 3.48 (0.45)| 2.77 (2.07)| 4.9 (0.56)| 5.7 (0.80)| 5.4 (1.84)| 6.5 (0.24)| 4.09 | 0.0212 |
| Moist %   | 0.18 (0.04)| 0.16 (0.05)| 0.15 (0.08)| 0.09 (0.04)| 0.13 (0.04)| 0.10 (0.06)| 1.27 | 0.338 |

| VEGET.HEIGHT | Mean (SD) | UGraz (1) | Graz (1) | Graz (2) | NU Graz | NGraz | F(χ²) | P |
|--------------|-----------|-----------|----------|----------|---------|-------|-------|---|
| Lichen       | 13.33 (1.32)| 13.61 (0.78)| 5.00 (1.12)| 4.94 (1.24)| 7.22 (0.79)| 5.10 (0.95)| 43.18 | <0.0001 |
| Moss         | 3.89 (2.15)| 4.89 (1.76)| 1.63 (0.74)| 1.88 (0.78)| 3.00 (1.87)| 1.13 (0.23)| 8.49  | <0.0001 |
| Vascular plants | 23.11 (8.08)| 18.33 (3.84)| 14.00 (2.35)| 10.67 (3.31)| 14.44 (4.25)| 10.67 (2.35)| 10.38 | <0.0001 |

| VEGET.BIOM. | Mean (SD) | UGraz (1) | Graz (1) | Graz (2) | NU Graz | NGraz | F(χ²) | P |
|-------------|-----------|-----------|----------|----------|---------|-------|-------|---|
| Lichen      | 165.29 (68.14)| 181.62 (49.12)| 29.09 (9.84)| 31.96 (11.96)| 41.17 (10.50)| 50.50 (17.27)| 26.65 | <0.0001 |
| Moss        | 8.36 (14.28)| 11.23 (9.52)| 4.29 (8.46)| 8.62 (13.75)| 21.40 (39.24)| 2.38 (3.15)| 0.77  | 0.580 |
| Vascular plants | 43.40 (16.39)| 35.45 (18.82)| 28.09 (12.11)| 36.64 (20.37)| 34.45 (19.20)| 23.43 (6.94)| 1.09  | 0.384 |
Table 4 (on next page)

Spearman rank correlations of the abundances (n) and species richnesses (S) of spider families. (* = p<0.05, **p<0.01)(n=18).
|               | Lycosidae n | Lycosidae S | Linyphiidae n | Linyphiidae S |
|---------------|-------------|-------------|---------------|---------------|
| Lycosidae n   | 1           |             |               |               |
| Lycosidae S   | -0.842**    | 1           |               |               |
| Linyphiidae n | -0.418      | 0.495*      | 1             |               |
| Linyphiidae S | 0.712**     | -0.628**    | -0.203        | 1             |
| Others n      | -0.294      | 0.376       | -0.362        | 0.710**       |
| Others S      | 0.539*      | -0.602**    | 0.221         | -0.005        |
Table 5

The responses of spider family species richnesses (s) and abundances (n), as well as the abundances of the most numerous species to environmental variables.

Linear regression results are also included (* = p<0.05, **p<0.01)(df=16). S = number of species, n = number of specimen.
|                      | Soil moist. | Soil temp. |                      | Soil moist. | Soil temp. |
|----------------------|-------------|------------|----------------------|-------------|------------|
|                      | Lichen bm   | %          | °C                   | Lichen bm   | %          | °C                   |
| Tot. species (S)     | 0.47        | 0.28       | 5.68 *               | A.aculeata  | 1.06       | 0.65     1.38 |
| Lycosidae S          | 3.64        | 0.33       | 10.71 **             | O.arctica   | 4.92*      | 0.1       | 2.02 |
| Linyphiidae S        | 0.41        | 2.31       | 0.02                 | H.ononidum  | 0.26       | 0.69     0.05 |
|                       |             |            |                      | P.antroniensis |           |           | |
| Thomisidae S         | 1.09        | 0.11       | 0.01                 | A.gulosa    | 37.17**    | 1.96     11.98** |
| Gnaphosidae S        | 2.8         | 0.44       | 1.32                 |             | 6.28*      | 4.94*    6.52** |
| Tot. specimens (n)   | 0.43        | 3.32       | 0                    | P.pumila    | 2.11       | 4.42*    1.87 |
| Lycosidae n          | 0.33        | 0.85       | 1.16                 | H.herniosa  | 17.7**     | 0.25     | 3.28 |
| Linyphiidae n        | 20.69 **    | 1.09       | 8.09 *               | W.karpinskii | 0.86       | 0.02     0.35 |
| Thomisidae n         | 6.54 **     | 0.25       | 2.42                 | M.herbigrad | 0.27       | 12**     6.57** |
| Gnaphosidae n        | 0.11        | 3.954      | 0                    |             |            |           |