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Four years of experimental warming do not modify the interaction between subalpine shrub species

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Author Contributions: AAR, JMN, SP, OG and EC conceived and designed the experiments. AAR, JN and EC performed the experiments in the field. AAR, SP, MCS and EN performed laboratory analyses. AAR and EN analyzed the data. AAR wrote the manuscript with the substantial advice, corrections and comments of SP, JMN, OG, EC, SN and EN. All the authors contributed to the discussion of the results.

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1    **Abstract**

2    Climate warming can lead to changes in alpine plant species interactions through modifications in  
3    environmental conditions, which may ultimately cause drastic changes in plant communities. We  
4    explored the effects of four years of experimental warming with open-top chambers (OTC) on  
5    *Vaccinium myrtillus* performance and its interaction with neighbouring shrubs at the Pyrenean treeline  
6    ecotone. We examined the effects of warming on height, above-ground (AG) and below-ground (BG)  
7    biomass and the C and N concentration and isotope composition of *V. myrtillus* growing in pure stands  
8    or in stands mixed with *Vaccinium uliginosum* or *Rhododendron ferrugineum*. We also analysed  
9    variations in soil N concentrations, rhizosphere C/N ratios and the functional diversity of the microbial  
10   community, and evaluated whether warming altered the biomass, C and N concentration and isotope  
11   composition of *V. uliginosum* in mixed plots. Our results showed that warming induced positive  
12   changes in the AG growth of *V. myrtillus* but not BG, while *V. uliginosum* did not respond to  
13   warming. *Vaccinium myrtillus* performance did not differ between stand types under increased  
14   temperatures, suggesting that warming did not induce shifts in the interaction between *V. myrtillus* and  
15   its neighbouring species. These findings contrast with previous studies in which species interactions  
16   changed when temperature was modified. Our results show that species interactions can be less  
17   responsive to warming in natural plant communities than in removal experiments, highlighting the  
18   need for studies involving the natural assembly of plant species and communities when exploring the  
19   effect of environmental changes on plant-plant interactions.

20

21    **Keywords:** dwarf shrub, plant interactions, Pyrenees, *Vaccinium myrtillus*, passive warming

22    **Abbreviations:** AG (above-ground), BG (below-ground),  $\delta^{13}\text{C}$  (carbon isotope composition),  $\delta^{15}\text{N}$   
23    (nitrogen isotope composition)

24      **Introduction**

25            In recent decades, climate warming and land-use change (i.e. abandonment of extensive  
26        livestock grazing and tree logging) have led to shrub encroachment processes in the alpine treeline  
27        ecotone worldwide (Dullinger et al. 2003; Rundqvist et al. 2011; Ropars and Boudreau 2012). The  
28        forecasted global air temperature increase of 1.0–3.7 °C by the end of the century could accelerate  
29        these processes; especially at high elevations and latitudes, where change is predicted to be larger  
30        (Nogués-Bravo et al 2007; Collins et al. 2013; Rangwala et al. 2013). This could have a dramatic  
31        impact on alpine and Arctic tundra ecosystems due to shifts in community composition and potential  
32        feedbacks to warming, such as decreasing albedo due to the higher radiation absorption by shrub  
33        canopies, reducing radiative cooling at night through the reflection of the thermal energy emitted by  
34        the soil, or through the inputs of more recalcitrant litter in the ecosystem (Hobbie 1996; Cornelissen et  
35        al. 2007; Myers-Smith et al. 2011; D’Odorico et al. 2013).

36            Many studies in Arctic and alpine ecosystems have shown the need to conduct species-specific  
37        studies to understand vegetation changes with warming, since coexisting species may differ in their  
38        responses to increasing temperatures (Kudo and Suzuki 2003; Klanderud 2008; Anadon-Rosell et al.  
39        2014; Little et al. 2015; Yang et al. 2015). However, it is also important to consider plant–plant  
40        interactions, since they are crucial for plant community dynamics (Callaway and Walker 1997). The  
41        stress-gradient hypothesis (Bertness and Callaway 1994) postulates that competition is the major  
42        selective force in habitats with more benign environmental conditions, whereas facilitation dominates  
43        in more severe environments. Many studies in cold regions across the globe have shown that plant  
44        interactions shift from facilitation to competition as temperature increases, or in the opposite direction  
45        when temperature decreases (Shevtsova et al. 1997; Choler et al. 2001; Klanderud 2005; Pugnaire et  
46        al. 2015; Wheeler et al. 2015; Olsen et al. 2016). Nevertheless, most of these studies involved plant  
47        removal experiments, and studies focusing on the effects of temperature changes on plant interactions  
48        within natural communities are scarce (but see Dormann et al. 2004).

49            Shrubs are major components of alpine and Arctic tundra ecosystems. Amongst them, clonal  
50        dwarf shrub species are of great importance in terms of vegetation cover, structure and functionality.  
51        They present a complex network of subterranean rhizomes bearing fine roots, and producing

52 individual above-ground (AG) ramets. Thus, the below-ground (BG) system of clonal shrubs is  
53 essential for their persistence and vegetative expansion, as well as an important source of soil carbon  
54 (C) (Cornelissen et al. 2014). Changes in the BG structure of dominant clonal shrubs could translate  
55 into major changes in the community and ecosystem functioning. Consequently, the study of BG  
56 responses to warming is an essential part of the complex responses to temperature increase in Arctic  
57 and alpine areas. However, the destructive nature of BG sampling and the difficulty to identify and  
58 separate roots from different species, together with the compromise of having studies running for the  
59 longest term possible, explain why warming experiments including both AG and BG plant  
60 measurements are infrequent (but see Hollister and Flaherty 2010 and Yang et al. 2015, amongst  
61 others).

62 Global warming may also induce shifts in the composition and function of the soil microbial  
63 community (Streit et al. 2014; Classen et al. 2015; DeAngelis et al. 2015), which can have strong  
64 impacts on ecosystem functioning (Schimel and Schaeffer 2012). For instance, rising temperatures can  
65 alter nitrogen (N) mineralization, with effects on N availability and, ultimately, plant growth (Bardgett  
66 and Wardle 2010). Several studies in cold ecosystems have found an increase in the soil N pool size  
67 with warming (Chapin et al. 1995; Hartley et al. 1999; Dijkstra et al. 2010; Dawes et al. 2011; Bai et  
68 al. 2013), which has been related to a stimulation of mineralization and decomposition processes.  
69 Since coexisting species show different N preferences and N-acquisition strategies (Körner et al. 2003;  
70 Pöron et al. 2007), shifts in N pools may affect interspecific interactions by altering relative niche  
71 and fitness differences between species (Chesson 2000; Tilman and Lehman 2001).

72 *Vaccinium myrtillus* L. forms shrub patches that colonize subalpine and alpine grasslands in  
73 the Pyrenees, where it grows close to the upper altitudinal limit of its distribution (Bolòs et al. 2005),  
74 subjected to low temperatures and short growing seasons. Warmer temperatures could favour its  
75 growth at the treeline ecotone, as has been reported in warming experiments in the Alps (Dawes et al.  
76 2011; Anadon-Rosell et al. 2014) and in the Arctic tundra (Rinnan et al. 2009; Taulavuori et al. 2013).  
77 However, co-occurring species such as *Vaccinium uliginosum* or *Empetrum hermaphroditum* have not  
78 been found to respond to temperature increase (Richardson et al. 2002; Kudo and Suzuki 2003;  
79 Anadon-Rosell et al. 2014) and, consequently, interactions between these species might shift with  
80 warming. On the other hand, in line with the stress-gradient hypothesis, a modification of the

81 environment through air temperature increase could induce changes in the interaction between this  
82 species and its neighbours towards increased competition. Despite the numerous studies focusing on  
83 *V. myrtillus* in tundra ecosystems, to our knowledge the potential effects of warming on the interaction  
84 with its neighbours have not been reported. Moreover, the previously mentioned experiments on *V.*  
85 *myrtillus* have mainly focused on its AG responses to warming, whereas BG effects have not been  
86 assessed.

87 At the treeline ecotone in the Central Pyrenees, *V. myrtillus* grows in pure patches (stands  
88 hereafter) or in mixed stands together with *Vaccinium uliginosum* L. subsp. *microphyllum* (Lange)  
89 Tolm. (hereafter *V. uliginosum*), or *Rhododendron ferrugineum* L. A previous study did not find  
90 evidence that the co-occurrence with these neighbouring shrubs had major effects on *V. myrtillus*  
91 structure and functioning (Anadon-Rosell et al. 2016). The objective of the present study was to  
92 investigate the AG and BG effects of four years of passive warming on *V. myrtillus*, and whether  
93 warming induced changes in interactions between *V. myrtillus* and its neighbouring species: a shrub of  
94 a very similar size (*V. uliginosum*) and a taller shrub (*R. ferrugineum*). For this purpose we assessed *V.*  
95 *myrtillus* phenology, AG and BG biomass, C and N concentration and isotopic signature ( $\delta^{13}\text{C}$  and  $\delta$   
96  $^{15}\text{N}$ ), soil inorganic N concentrations ( $\text{NO}_3^-$  and  $\text{NH}_4^+$ ) and C and N availability in the rhizosphere in  
97 different types of *V. myrtillus* stands subjected to warming treatments. We also assessed the effects of  
98 warming on *V. uliginosum* AG biomass and C and N concentration and isotopic signature in mixed  
99 stands. Moreover, we evaluated the effects of warming on the functional diversity of the microbial  
100 community in *V. myrtillus* pure stands. We hypothesized that warming will (i) benefit *V. myrtillus* AG  
101 and BG performance (i.e. biomass and physiological traits), but that it will also (ii) modify interactions  
102 with co-occurring species, which will be manifested as different responses to warming of *V. myrtillus*  
103 when growing in pure stands than when growing in mixtures. We also hypothesized that (iii) the soil  
104 inorganic N pool will increase under warming with different magnitude across stand types.

105

## 106 **Materials and methods**

### 107 *Study area*

108 The study site was located at Eth Corrau des Machos (Val d'Aran), in the buffer zone of the  
109 Aigüestortes and Estany de Sant Maurici National Park (Central Pyrenees, Catalonia, UTM

110 coordinates 31N 329, 472), on a N-facing 10-15° steep slope at 2250 m a.s.l. The vegetation consisted  
111 of *Festuca eskia* Ramond ex DC. and *Nardus stricta* L. grasslands mixed with patches of dwarf shrub  
112 heath dominated by *V. myrtillus*, *V. uliginosum* and *R. ferrugineum*. For the period 2001-2013, the  
113 mean annual precipitation and mean annual temperature ( $\pm$  SD) were 1146.4 ( $\pm$  58.3) mm and 3.0 ( $\pm$   
114 0.2) °C, respectively. For the study period (2010-2013) the mean annual precipitation and mean annual  
115 temperature were 1223.1 ( $\pm$  244.8) mm and 2.7 ( $\pm$  0.9) °C, respectively. The mean monthly  
116 precipitation and the mean temperature for the main months of the growing season (June–August)  
117 were 99.1 ( $\pm$  28.5) mm and 10.2 ( $\pm$  1.0) °C (obtained from a meteorological station at a nearby  
118 location: La Bonaigua, 6.3 km away from the study site and at a similar altitude, run by the  
119 Meteorological Service of Catalonia, [www.meteo.cat](http://www.meteo.cat), accessed in May 2014).

120 *Experimental design*

121 In July 2010 we established 30 plots of 1.1 m<sup>2</sup> combining a stand type and a warming treatment. We  
122 selected 10 pure stands of *V. myrtillus* (M stands), 10 mixed stands of *V. myrtillus* and *V. uliginosum*  
123 subsp. *microphyllum* (U stands), and 10 mixed stands of *V. myrtillus* and *Rhododendron ferrugineum*  
124 (R stands). In each stand, shrubs were the dominant species (Table S1), but other grasses and forbs  
125 were also present. Amongst them, the most abundant species were *Festuca eskia*, *Festuca nigrescens*,  
126 *Nardus stricta*, *Trifolium alpinum*, *Phleum alpinum* and *Meum athamanticum*. The distance between  
127 two plots ranged from one to a few metres (< 20 m), always ensuring that the studied patches were  
128 independent from each other. Soil organic matter and organic C concentrations, pH and colour  
129 (according to Munsell System) were similar across plots (Table S2). In half of the plots (i.e. 15 plots,  
130 five for each stand type) we placed an open-top chamber (OTC) made of transparent polycarbonate,  
131 similar to the model used in the International Tundra Experiment (ITEX; Marion et al. 1997). The  
132 other 15 plots served as ambient air temperature controls. The air temperature increase inside the  
133 OTCs in summer was 1.1 °C, which we measured with temperature loggers (*iButton* 1-wire  
134 Thermochron, Embedded Data Systems, USA) placed at ground level in two plots of each stand type x  
135 warming combination during the growing season 2013 (recording every hour). The snow  
136 accumulation was high and homogenous along the study site and our phenological survey did not

137 reveal substantial irregularities in the snowmelt pattern, even despite the presence of OTCs. Thus, we  
138 left the OTCs in place throughout the experiment.

139 *Phenology and community composition*

140 In 2011 we labelled six *V. myrtillus* ramets per plot, which we monitored during the growing seasons  
141 of 2011 and 2012 for a phenological survey. We recorded the following phenophases: winter state,  
142 bud swelling, bud bursting, leaf expansion, shoot elongation, vegetative state, leaf colour change, leaf  
143 shedding, leafless state and shoot winter colouring (brown-red coloration). We visited the plots *ca.*  
144 once a month starting after snowmelt until late Autumn, when ramets were leafless, and we recorded  
145 the presence of different phenophases in the six marked ramets. We assigned an ordinal numeric code  
146 to all phenophases and calculated the average numeric code per plot as the average score of the six  
147 ramets at each visit.

148 Plant community composition within the study plots was first recorded in 2011, by estimating  
149 the percentage cover of the main plant groups in each plot, i.e. shrubs and grasses. This was re-  
150 assessed in September 2013 before the end of the experiment (Table S1). Lichen and bryophyte cover  
151 was very low and was not recorded.

152 *AG and BG biomass*

153 On the 3rd September 2013 we harvested five *V. myrtillus* ramets per plot (not corresponding with  
154 those phenologically surveyed) plus five *V. uliginosum* ramets in U plots, avoiding sampling close to  
155 the edges of the OTCs. We also dug out their rhizomes (down to *ca.* 20 cm long) and the roots  
156 attached, and collected six soil cores of 12 cm length x 4 cm diameter in each plot (corresponding to 0-  
157 15 cm depth), which were kept in sealed plastic bags in a cool box until they arrived to the lab. Two of  
158 these cores were kept frozen and were used for BG biomass measurements at the plot scale. The rest  
159 were kept refrigerated at 4 °C and were used for measurements of soil NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> concentration  
160 and analyses of the functional diversity of the microbial community (two cores); and for rhizosphere C  
161 and N concentration and isotope composition analyses (two cores). Soil cores obtained for the same  
162 purpose from each plot were pooled together. Therefore, we had one composite soil sample per plot  
163 for each type of measurement.

164 Once in the lab, we measured the ramet height of both *Vaccinium* species and counted the  
165 scars left by the buds in each ramet to estimate their age. Then, we separated leaves, new shoots (i.e.  
166 shoots grown in 2013), rhizomes and roots of each species, and dried them at 60 °C for 48 hours.  
167 Leaves and new shoots were weighed for AG biomass measurements and subsequently used for N and  
168 C concentration and isotope composition analyses. Rhizomes and roots were only used for N and C  
169 concentration and isotope composition analyses because BG biomass was measured at the plot scale  
170 on material obtained from the soil cores. We conducted BG biomass measurements referring to a  
171 specific soil volume to make comparisons between warming treatments and stand types possible. Soil  
172 cores for BG biomass measurements were sieved to separate rhizomes, coarse roots ( $\geq 1$  mm diameter)  
173 and fine roots ( $< 1$  mm diameter). We dried them in the oven at 60 °C for 48 h and weighed them for  
174 BG biomass analyses.

175 *Carbon and nitrogen concentration and isotope composition*

176 For the analyses of C and N concentration and isotope composition of leaves, new shoots, rhizomes  
177 and roots, we pooled together the material from all the harvested ramets of each plot for each plant  
178 part of each *Vaccinium* species. Then we ground the material and weighed *ca.* 1 mg subsamples in  
179 small tin capsules. The C and N concentrations of samples were determined using an Elemental  
180 Analyzer Flash 1112 (Carbo Erba, Milan). The C and N isotope composition of samples were  
181 determined using the Elemental Analyzer coupled to an IRMS Delta C isotope ratio mass spectrometer  
182 through a Conflo III Interface (Thermo-Finnigan, Germany). The results of C isotope analyses are  
183 reported in per thousand (‰) on the relative  $\delta$ -scale as  $\delta^{13}\text{C}$ , and refer to the international standard V-  
184 PDB (Vienna Pee Dee Belemnite) according to the following equation:

$$185 \quad \delta^{13}\text{C} = \left( \frac{R_{\text{sample}}}{R_{\text{standard}}} \right) - 1 \quad (\text{Eq. 1})$$

186 where  $R$  is the  $^{13}\text{C}/^{12}\text{C}$  ratio.

187 N isotopic composition results are also expressed in  $\delta$  notation ( $\delta^{15}\text{N}$ ) using international  
188 secondary standards of known  $^{15}\text{N}/^{14}\text{N}$  ratios (IAEA N<sub>1</sub> and IAEA N<sub>2</sub> ammonium sulphate and IAEA  
189 NO<sub>3</sub> potassium nitrate) relative to N<sub>2</sub> in air:

$$190 \quad \delta^{15}\text{N} = \left( \frac{R_{\text{sample}}}{R_{\text{standard}}} \right) - 1 \quad (\text{Eq. 2})$$

191 where  $R$  is the  $^{15}\text{N}/^{14}\text{N}$  ratio.

192 We used C and N concentrations to calculate C/N ratios of each plant part of the two  
193 *Vaccinium* species. We calculated total N pools for all *V. myrtillus* plant parts (for BG parts we only  
194 used data from pure stands, referred to soil volume) and for *V. uliginosum* AG plant parts. All EA-  
195 IRMS analyses were performed at the Scientific and Technological Centers (CCiT) of the University  
196 of Barcelona. The  $\delta^{13}\text{C}$  of CO<sub>2</sub> of the air and the  $\delta^{15}\text{N}$  of the bulk soil were analysed in 2013 and were  
197 *ca.* -10.91‰ and *ca.* 7.33 ‰, respectively (see Anadon-Rosell et al. 2016 for more information on air  
198 and soil sampling).

199 *Soil inorganic N concentrations*

200 NO<sub>3</sub><sup>-</sup> concentrations were measured following the UV method described by Kaneko et al. (2010) by  
201 measuring the absorbance of KCl extracts from soils at 220 nm and 260 nm wavelengths. Soil NH<sub>4</sub><sup>+</sup>  
202 concentrations were measured by the conversion of ammonium into the intense blue indophenol  
203 complex (IPC) using salicylate, following the methods used by Kempers and Kox (1989).

204 Rhizospheric soil analyses and functional diversity of the microbial community

205 We carefully selected rhizomes and roots from the two soil cores collected for rhizosphere analyses  
206 and separated the soil that was attached using a small paint brush. We ground the soil and weighed *ca.*  
207 3.5 mg subsamples in small tin capsules and analysed its C and N concentration and isotope  
208 composition following the same procedure as for plant tissues.

We assessed the impact of the warming treatment on the use of different C sources by soil microbial communities using Biolog EcoPlates (Insam 1997). Every plate had 96 wells containing 31 different C sources plus a blank well, in three replications. The rate of utilization of the C sources by microorganisms results in the increase of the optical density ( $OD_{590}$ ) (Pohland and Owen 2009). We analysed the use of C by soil microbial communities in *V. myrtillus* pure stands only, using three replicates per warming treatment. Soils were sieved at 2 mm before extracting the bacterial community. Bacterial cells were extracted by mixing 10 g fresh soil with 95 ml of sterilized Milli-Q water inside 100 ml Erlenmeyers (see details in Muñiz et al. 2014). The mixture was magnetically

shaken for 30 minutes, followed by one-hour rest. Afterwards, 10 ml of the soil suspension was put into 50 ml Falcon tubes and, after one-minute sonication, the tubes were centrifuged (1000 g, 10 minutes). 9.5 ml of the supernatant were separated and the remaining was resuspended after adding 9.5 ml of water. 47.5 ml of soil extract was obtained from each sample after five cycles of sonication-centrifugation. The extracts were kept at 4 °C for a few hours. Just before inoculating the Biolog plates, soil particles were removed by a low-speed centrifugation (500 g, 2 minutes). 150 µl of the soil extract were put into each well of the plates by a multipipete. All the laboratory material was sterile or it was previously autoclaved (121 °C, 20 minutes) and the operations were made inside a biological laminar flow chamber. The plates were incubated in the dark at 25 °C for 120 hours. The OD<sub>590</sub> of each well was measured just after the inoculation (at 0, 5, 72, 96, 101 and 120 h) using the Anthos 2010 microplate reader and ADAP 2.0 Software (Biochrom, Ltd. Cambridge Science Park, Cambridge, CB4 0FJ. England).

#### 229 Statistical analyses

We tested the effects of warming and stand type on *V. myrtillus* phenology, ramet height and AG biomass using linear mixed effects models fitted with the restricted maximum likelihood estimation method (REML). We included warming and stand type as fixed factors and plot as a random factor to account for the multiple sampling within a plot. We used the same models for *V. uliginosum* variables, but in this case we only used warming as a fixed factor. To test the effects of warming and stand type on the C and N concentration and isotope composition of the different AG and BG tissues, BG biomass, soil NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> concentration, rhizosphere C and N concentration and isotope composition we used simple linear model functions. We included ramet age as a covariate when it significantly contributed to improve model fits (which we tested with likelihood ratio tests) to account for possible age effects on our growth-related response variables. This was the case in the models for *V. myrtillus* height and AG biomass, plus the models for height, number of shoots and dry weight per shoot unit for *V. uliginosum*. We tested for significance with analysis of variance tests and graphically evaluated the assumptions of normality and homoscedasticity of residuals (Zuur et al. 2009). We double-checked with Shapiro and Bartlett tests when the visual evaluation of graphs was difficult. We log-transformed data when necessary to satisfy these assumptions. Moreover, when homoscedasticity

245 of residuals was not met, we used the varIdent structure (Zuur et al. 2009; Pinheiro et al. 2016) to  
246 account for the heterogeneity of variances among factor levels. In *V. uliginosum* analyses, when both  
247 normality and homoscedasticity were not met, we used the non-parametric Wilcoxon test. We  
248 considered effects significant at  $P < 0.05$  and trending towards significance at  $0.05 > P < 0.10$  to  
249 account for the relatively low replication. When we found significant differences between stand types,  
250 we run Tukey HSD post hoc tests to determine those factor levels that differed significantly. We  
251 performed all the analyses with R 3.1.2 (R Core Team, 2015). For linear mixed effects models we  
252 used the *nlme* package (Pinheiro et al. 2016); for graphical evaluation of model assumptions we used  
253 the *lattice* package (Sarkar 2008); and for multiple comparisons we used the *multcomp* package on  
254 linear mixed effects models (Hothorn et al. 2008) and the *agricolae* package on simple linear models  
255 (de Mendiburu 2010).

256 The microbial activity of each microplate was expressed as the average well-colour  
257 development (AWCD) and was determined as previously described (Garland and Mills 1991) as  
258 follows:

259 
$$\text{AWCD} = \sum OD_i / 31 \quad (\text{Eq. 3})$$

260 where  $OD_i$  is the optical density value from each well at any given time. The AWCD curves were  
261 adjusted to a log-logistic 4-parameters model using R Software. This was done first for each replicate  
262 and also for the warming treatment levels, each including all replicates (in this case with the raw data  
263 of  $OD_{590}$ ). Then curve parameters (curve slope -slop-, maximal degradative capacity -max- and time to  
264 reach half of the slope -TM50-) were compared using the *CompParm* function in *drc* package (Ritz  
265 and Streibig 2005): the null hypothesis was that the ratio between parameters equalled 1. The ratio was  
266 obtained by dividing the same parameters from each curve by the other parameters; if the ratio  
267 significantly differed from 1, the null hypothesis was rejected, meaning that values were significantly  
268 different ( $P < 0.05$ ).  $P$ -values were adjusted using Bonferroni correction for multiple T-tests.

269

270 **Results**

271 *Phenology*

272 Warming advanced early-season vegetative phenology through an earlier onset of bud burst and leaf  
273 expansion (Fig. S1). On the 21st May 2011 (day of year, DOY, 141), *V. myrtillus* ramets in unwarmed  
274 plots were at the bud swelling phase, whereas in warmed plots had already started bursting ( $F_{1,24} =$   
275 3.92,  $P = 0.059$ ). In 2012, *V. myrtillus* ramets in warmed plots were expanding their leaves on the 14th  
276 June (DOY 166), whereas ramets in unwarmed plots were still in the bud burst phenophase ( $F_{1,24} =$   
277 6.59,  $P = 0.017$ ). Monitoring later in the season for both years did not show any other significant  
278 differences between warming treatments (see Fig. S1 for visit dates). We only found significant  
279 differences between stand types (regardless of the warming treatment) in September 2011, when  
280 ramets in M stands were already shedding their leaves whereas ramets in the other two stand types had  
281 just started changing their colour prior to leaf shedding ( $F_{2,24} = 9.31$ ,  $P = 0.001$ ). This advancement in  
282 senescence in M stands with respect to the other stand types was especially obvious in warmed plots  
283 (the interaction stand type x warming trended towards significance,  $F_{2,24} = 3.13$ ,  $P = 0.062$ ).

284 *Age and AG biomass of Vaccinium species*

285 Our age analyses confirmed that there were no differences in *V. myrtillus* ramet age between warming  
286 treatments ( $F_{1,24} = 0.16$ ,  $P = 0.696$ ) nor between stand types ( $F_{2,24} = 1.89$ ,  $P = 0.173$ ). Likewise *V.*  
287 *uliginosum* did not show differences in ramet age between warming treatments ( $F_{1,6} = 0.01$ ,  $P =$   
288 0.930).

289 After four years of warming, *V. myrtillus* ramets were 15% taller in warmed plots than in  
290 unwarmed plots. There were no differences in ramet height between stand types or an interaction  
291 between warming and stand type (Table 1). Similarly, there was no warming effect on *V. uliginosum*  
292 height ( $F_{1,6} = 0.08$ ,  $P = 0.784$ ).

293 *Vaccinium myrtillus* leaf biomass per ramet did not differ between warming treatments (Table  
294 1, Fig. 1a) but new shoot biomass was higher under warming than in control plots (Fig. 1b). The total  
295 above-ground biomass per ramet was also higher in warmed plots than in unwarmed plots (Table 1,  
296 Fig. 1c). There were no differences between stand types or a stand type x warming interaction for *V.*  
297 *myrtillus* AG biomass (Table 1). There were no differences between warming treatments in terms of *V.*  
298 *uliginosum* leaf biomass ( $F_{1,6} = 2.77$ ,  $P = 0.147$ ), new shoot biomass ( $F_{1,6} = 0.04$ ,  $P = 0.849$ ) or total  
299 AG biomass ( $F_{1,6} = 0.39$ ,  $P = 0.554$ , Fig. S2), but we found contrasting effects of warming on the dry

300 weight per shoot and the number of new shoots. Dry weight per shoot in *V. uliginosum* was higher  
301 inside the OTCs than in control plots ( $F_{1,6} = 6.42, P = 0.044$ ), whereas the number of new shoots was  
302 higher in ramets from unwarmed plots ( $F_{1,6} = 14.81, P = 0.009$ ).

303 *Vaccinium myrtillus BG biomass*

304 There were no effects of warming on *V. myrtillus* BG biomass (Fig. 1d, e, f). We only found  
305 differences in rhizome and coarse root biomass between stand types. R stands showed lower rhizome  
306 biomass per soil volume than in the other two stand types ( $F_{2,24} = 6.93, P = 0.004$ , Fig. 1d). U stands  
307 showed a trend towards significantly greater coarse root biomass than M stands ( $F_{2,19} = 3.04, P =$   
308 0.071, Fig. 1e). Fine root biomass did not differ between stand types ( $F_{2,24} = 0.41, P = 0.667$ , Fig. 1f).  
309 We did not find any warming x stand type interaction for any of the BG plant parts analysed (Table  
310 S3).

311 *C and N concentration and isotope composition of AG and BG plant fractions*

312 C concentration in *V. myrtillus* organs was similar across warming treatments and stand types for  
313 leaves, shoots and roots. Rhizomes, however, had greater C concentration under warming than in  
314 control plots (Table 2, Fig. 2), which was not related to any rhizome biomass increase under warming  
315 (see above). C concentration values of *V. uliginosum* new shoots, rhizomes and roots did not show any  
316 response to warming, but there was a trend towards a significantly positive effect of warming on leaf  
317 C concentration (Table 3, Fig. 3).

318 The  $\delta^{13}\text{C}$  of *V. myrtillus* and *V. uliginosum* tissues did not differ between warming treatments  
319 (Fig. 2, 3) but we found significant differences in the  $\delta^{13}\text{C}$  of *V. myrtillus* tissues between stand types.

320 *Vaccinium myrtillus*  $\delta^{13}\text{C}$  was lower in R stands than in the other two stand types for leaves (only  
321 trending towards significance), shoots and rhizomes. There were no significant differences between  
322 stand types for the  $\delta^{13}\text{C}$  composition of roots (Table 2, Fig. 2), or any warming x stand type  
323 interaction.

324 There was no warming effect on the N concentration and  $\delta^{15}\text{N}$  of any of the *V. myrtillus*  
325 organs, and only a very marginal trend towards significance of the interaction between warming and  
326 stand type in the N concentration of *V. myrtillus* rhizomes, which was higher in control plots than in  
327 warmed plots in U stands (Table 2, Fig. 2). However, we found significant differences between stand

types. Leaf N concentration was higher in R stands than in U stands, but this was not the case for any of the other plant organs. Leaf and shoot  $\delta^{15}\text{N}$  values were higher in M stands than in the other two stand types. Finally, rhizome  $\delta^{15}\text{N}$  values were also higher in M stands than in the other two stand types, but only significantly higher than in R stands (Table 2, Fig. 2). The N pool in *Vaccinium myrtillus* rhizomes ( $0.037 \pm 0.006 \text{ mg/cm}^3$ ) and roots ( $0.010 \pm 0.002 \text{ mg/cm}^3$ ) did not differ between warming treatments ( $F_{1,7} = 0.51, P = 0.497; F_{1,8} = 0.60, P = 0.462$ , respectively). Leaf N pools did not differ between warming treatments ( $F_{1,24} = 2.21, P = 0.150$ ) or stand types ( $F_{2,24} = 1.34, P = 0.282$ ) (average across warming and stand type treatments of  $9.16 \pm 0.65 \text{ mg}$ ). The N pool in *V. myrtillus* new shoots was higher in warmed ( $5.46 \pm 0.62 \text{ mg}$ ) than in unwarmed plots ( $3.72 \pm 0.38 \text{ mg}$ ) ( $F_{1,24} = 5.82, P = 0.024$ ), but did not differ between stand types ( $F_{2,24} = 0.93, P = 0.408$ ). We did not find any warming effect on C/N ratios, only a trend towards significance for rhizomes in U stands, which showed higher values under warming than in unwarmed plots (Table 2, Fig. S3). We only found significant differences in C/N ratios between stand types in leaves, which showed higher values in U stands than in the other two (Table 2, Fig S3). New shoots and roots did not show significant differences in their C/N ratios for any of the treatments.

*Vaccinium uliginosum* shoots showed significantly lower N concentrations under warming than in unwarmed plots. This seemed associated with an increase in leaf N concentrations under warming (although the latter was not significant; Table 3).  $\delta^{15}\text{N}$  values did not differ significantly between warming treatments (Table 3, Fig. 3). Leaf and new shoot N pools in *V. uliginosum* did not differ between warming treatments either ( $F_{1,6} = 1.74, P = 0.235; F_{1,6} = 0.04, P = 0.843$ , respectively). The total amount of N in leaves was  $9.25 \pm 0.78 \text{ mg}$  and in new shoots  $1.92 \pm 0.23 \text{ mg}$  (averaged across warming treatments). C/N ratios did not differ significantly between warming treatments in leaves ( $24.58 \pm 0.74, F_{1,6} = 1.59, P = 0.254$ ), rhizomes ( $93.12 \pm 6.30, F_{1,6} = 0.03, P = 0.874$ ) or roots ( $53.87 \pm 1.96, F_{1,6} = 0.90, P = 0.379$ ) (averages across warming treatments shown). However, in new shoots C/N values were higher under warming ( $52.37 \pm 0.95$ ) than in unwarmed plots ( $46.28 \pm 1.84; F_{1,6} = 8.65, P = 0.026$ ).

*Soil inorganic N concentrations, rhizosphere C and N and functionality of the soil microbial community*

356 Soil  $\text{NO}_3^-$  concentration decreased by 36% in warmed plots compared with unwarmed plots ( $F_{1,24} = 5.87, P = 0.023$ , Fig. 4a), but the  $\text{NH}_4^+$  concentration remained similar between warming treatments ( $F = 0.45, P = 0.508$ , Fig. 4b). As a consequence, the  $\text{NO}_3^-/\text{NH}_4^+$  ratio decreased by 27% under warming  
357 with respect to control conditions. There was no difference between stand types or any interaction  
358 between warming and stand type for any of the two N forms analysed.

360       The rhizosphere C/N ratio did not differ between warming treatments. However, it differed  
361 between stand types, as it was higher in U stands than in the other two ( $F_{2,24} = 7.99, P = 0.002$ , Fig.  
362 S4). Both rhizosphere soil C and N concentration were significantly higher in U stands than in R and  
363 M stands ( $F_{2,24} = 5.81, P = 0.009$  and  $F_{2,24} = 3.64, P = 0.042$ , respectively), but the difference in the C  
364 concentration was greater than the difference in N (data not shown). There was no significant warming  
365 x stand type interaction on the rhizosphere C/N ratio ( $F_{2,24} = 0.89, P = 0.422$ ), but the high dispersion  
366 in the data could have masked possible differences between warming treatments in U stands. Neither  
367 warming nor stand type or their interaction had any effects on rhizospheric soil  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$  values  
368 ( $P > 0.28$ ).

370       The warming treatment did not change the ability of the microbial communities of *V. myrtillus*  
371 M stands to degrade the different sources of C present in the Ecoplates (Fig. S5 and S6), indicating  
372 that the exposure to 1.1 °C warmer temperatures during four years was not enough to result in  
373 significant changes in the soil C cycling by microbia.

374

## 375 **Discussion**

376 *Vaccinium myrtillus* slightly benefitted from increased temperatures regardless of neighbourhood  
377 composition. Thus, our results evidence that four years of passive warming did not lead to changes in  
378 the interaction between *V. myrtillus* and its neighbour species at our study site. Although coexistence  
379 theory predicts that intraspecific competition should be stronger than interspecific competition  
380 (Chesson 2000), we did not find evidence of stronger competition in pure than in mixed stands nor that  
381 this changed with warming, even when the percentage cover of *V. myrtillus* ramets in pure stands was  
382 higher than in mixed stands (and also ramet density, which was measured in 2010 in similar plots at  
383 the same study site; Table S1). A previous study at the same area also found that the performance of *V.*

384 *Vaccinium myrtillus* was similar across stand types, indicating that ramets did not perform worse in pure stands  
385 than in mixtures (Anadon-Rosell et al. 2016). Furthermore, other studies have reported that  
386 intraspecific facilitation can be more important than competition at moderate to high levels of stress  
387 (Chu et al. 2008; Fajardo and McIntire 2011).

388 The lack of warming effects on species interactions in our study contrasts with previous  
389 studies in which the interaction between plant species shifted when temperatures changed (both  
390 naturally and experimentally). Dormann et al. (2004) found that the interaction between the rush  
391 *Luzula confusa* and the deciduous shrub *Salix polaris* changed with warming in favour of *S. polaris* in  
392 Svalbard. In a removal experiment in Finse, Norway, Klanderud and Totland (2005) found that the  
393 removal of the neighbour species negatively affected *Thalictrum alpinum* in unwarmed plots, but not  
394 inside OTCs, indicating that warming affected the interaction between these species. Callaway et al.  
395 (2002) also reported evidence of a shift from facilitation at higher elevation sites to competition at  
396 lower elevations when removing neighbours of target individuals at different mountain sites across the  
397 world. In addition, a study in seminatural grasslands in southern Norway found increased competitive  
398 interactions with increasing temperature across precipitation and temperature gradients (Olsen et al.  
399 2016). Most of these studies consisted of removal experiments, which provide very important  
400 ecological and functional information about the community and species studied, and allow to test the  
401 effect of species composition on the community response to the environment (see review by Díaz et al.  
402 2003). However, despite their numerous advantages and outcomes, removal experiments cannot avoid  
403 the disturbance caused by the extraction of selected species. In contrast, our approach was based on  
404 naturally established populations and species and allowed us to assess their natural responses to  
405 warming. According to our results, species interactions seem less responsive to warming when studied  
406 under natural conditions and within the natural distributions of the study species. While the removal  
407 experiments mentioned above captured changes in species interactions after shorter periods of  
408 warming than our experiment, four years of passive warming were not sufficient to cause changes in  
409 species interactions at our study site.

410 The AG biomass increase of *V. myrtillus* inside the OTCs could be the result of a longer  
411 growing period caused by the advancement of its early-vegetative phenology. A previous study on the  
412 same species in the Swiss Alps showed that its increased growth after six years of soil warming with

heating cables was not related to a longer growing period (Anadon-Rosell et al. 2014). The above-ground phenology of ramets could be more affected by warming through OTCs than by soil warming, since air temperature at canopy level may be higher inside the OTC. However, other factors related to warming but not directly linked to a longer growing season could have also influenced *V. myrtillus* growth in our study, such as direct warming effects on photosynthetic rates (Heskel et al. 2013; Fu et al. 2015) or through the stimulation of N uptake rates, which would be supported by the lower soil  $\text{NO}_3^-$  values in warmed plots at our study (see below). In addition, although OTCs are a robust tool for the study of warming effects on plant species (Hollister and Webber 2000; see review by Bokhorst et al. 2013), they might potentially cause confounding effects on microclimate variables that could have influenced the response of our study species to the warming treatment, such as modifications in wind patterns around the study plants (DeBoek et al. 2012) and changes in soil moisture and vapour pressure deficit (Marion et al. 1997; Piiki et al. 2008).

In our study, the BG biomass of *V. myrtillus* did not change with warming, and this was consistent across stand types. Thus, the AG growth stimulation under warming did not result in increased BG growth, indicating an uncoupling between AG and BG responses to warming. This could potentially be related to different phenological responses to increased temperatures. In fact, an uncoupling between AG and BG phenology has been reported along an Arctic elevation gradient (Blume-Berry et al. 2016). Another explanation could be that OTCs mainly increase ground-level and air temperature. However, they have been found to slightly increase soil temperature at 5 cm depth (Hollister et al. 2006) and even at 10 cm in steppe ecosystems in Northern Mongolia (Sharkhuu et al. 2013). Hollister and Flaherty (2010) found a BG biomass increase in *Salix rotundifolia* at the Alaskan tundra after 3–4 years of warming with OTCs, but Shaver et al. (1998) found no BG biomass increase after 6–9 years of passive warming in another Alaskan wet sedge tundra site, indicating contrasting BG responses to warming depending on the study site and community composition. *Vaccinium myrtillus* can expand its rhizomes laterally several metres below-ground (Flower-Ellis, 1971); therefore our warming treatment might have not captured the potential response of a whole functional unit to warming, or a possible transfer of assimilates from AG parts might have been diluted by the complex BG network of this species. The lack of differences between warming treatments in the

441 rhizosphere C/N ratio and in the soil microbial C source use suggests that the degree of warming  
442 applied in our study was not enough to induce significant BG changes. Moreover, the similar substrate  
443 utilization by the microbial community between warming treatments indicates that the soil C pools  
444 were not altered by the increased temperature (Rinnan et al. 2009). Numerous studies on the effect of  
445 temperature on the composition and functioning of soil microbial communities have led to contrasting  
446 results (see review by Classen et al. 2015). Temperature shifts of as much as 10 °C did not  
447 significantly alter the physiological functioning of the microbiota of humic soils (Pettersson and Bååth  
448 2003). However, a temperature increase of 5 °C in temperate forests resulted in a significant alteration  
449 of soil microbial communities (DeAngelis et al. 2015), and an increase of 4 °C in a warming  
450 experiment at the Swiss treeline led to changes in the microbial substrate use (Streit et al. 2014). All  
451 these contrasting results indicate that warming effects on microbial metabolism and BG processes may  
452 be mediated through other biotic and abiotic factors (see also Christiansen et al. 2017), and certainly  
453 deserve further study.

454         *Vaccinium uliginosum* has been shown to be less plastic in response to warming than *V.*  
455 *myrtillus* (Richardson et al. 2002; Kudo and Suzuki 2003; Anadon-Rosell et al. 2014). This can be  
456 attributed to the better adaptation of *V. myrtillus* to higher temperatures, which is evidenced by the fact  
457 that it grows at lower altitudes than *V. uliginosum* (Bolòs et al. 2005). Although the dry weight of new  
458 individual shoots of *V. uliginosum* increased with warming, the number of shoots decreased, probably  
459 as a trade-off, which led to an overall lack of AG biomass response to warming in this species. In fact,  
460 only the leaf C concentration of *V. uliginosum* increased slightly with warming, and was not  
461 accompanied by any other changes in the shrub performance. Our study provides evidence that  
462 although *V. myrtillus* is more responsive to warming than *V. uliginosum*, it does not benefit more from  
463 warming when it grows in mixtures than when it grows in pure stands.

464         The slightly lower N concentration in *V. myrtillus* rhizomes in warmed plots than in  
465 unwarmed plots when coexisting with *V. uliginosum* suggests that there could be an increase in  
466 competition for N with warming. In fact, competition for N was found in mixed stands of the two  
467 *Vaccinium* species under natural conditions in a previous study at the same site (Anadon-Rosell et al.  
468 2016), and is supported by the higher rhizosphere and leaf C/N ratios in mixed stands of these two  
469 species than in pure stands found in this study. Moreover, the higher  $\delta^{15}\text{N}$  values in pure stands than in

470 mixed stands, which might be explained by lower N uptake through mycorrhiza or larger ecosystem  
471 losses of  $^{15}\text{N}$ -depleted N (leaving an enriched remaining pool), are indicative of larger N availability  
472 (Craine et al. 2009). A study in the Swiss Alps found a positive response to warming in *V. uliginosum*  
473 leaf N concentration but only a short-term positive response in *V. myrtillus* (Dawes et al. 2011). On the  
474 other hand, in the Swedish Lapland, *V. myrtillus* increased leaf N concentration in response to  
475 warming whereas the opposite was found for *V. uliginosum* (Richardson et al. 2002). However, these  
476 studies did not test warming effects on interspecific interactions. Our study suggests that although  
477 warming may increase the competition for N between *V. myrtillus* and *V. uliginosum*, this does not  
478 outweigh the positive growth response of *V. myrtillus* to warming.

479 A meta-analysis of experimental warming effects on N pools in terrestrial ecosystems showed  
480 that warming increased N mineralization rates and N pools across different ecosystem types (Bai et al.  
481 2013). However, in our experiment soil  $\text{NO}_3^-$  decreased with warming (regardless of the stand type).  
482 This could be due to greater  $\text{NO}_3^-$  uptake rates promoted by warming, since temperature is a modulator  
483 of plant N assimilation (Laine et al. 1994; Volder et al. 2000). The lack of an increase in the N  
484 concentration of *V. myrtillus* tissues could be due to a dilution effect caused by the greater biomass,  
485 which is supported by the higher total N pool found in new shoots under warming. Additionally, other  
486 species (especially grasses, due to their abundance) could have increased their  $\text{NO}_3^-$  assimilation under  
487 warming, which was not assessed in this study. Another explanation for the reduced soil  $\text{NO}_3^-$   
488 concentrations in the OTCs may be earlier consumption of  $\text{NO}_3^-$  through an advanced root phenology  
489 promoted by warming (Sullivan and Welker 2005; Nord and Lynch 2009). In the Finnish tundra,  
490 Rinnan et al. (2009) detected no increase in soil N concentration with warming either, but there was a  
491 decrease in soil  $\text{NH}_4^+$  concentration inside the OTCs. The authors argued that this reduction could  
492 reflect the increased efficiency of N uptake with warming. The differing responses in the N form  
493 between that study and ours might reflect the differential use of specific N forms at different sites with  
494 different community composition, or a greater availability of  $\text{NO}_3^-$  than  $\text{NH}_4^+$  at our study site.

495 In conclusion, four years of experimental warming had no effect on the interaction between *V.*  
496 *myrtillus* and *V. uliginosum* or *R. ferrugineum*. *Vaccinium myrtillus* showed a positive AG biomass  
497 response to warming regardless of the neighbouring species, but no BG responses were found.  
498 Although warming seemed to increase the competition for N between the two *Vaccinium* species, their

499 overall performance was not affected. This study shows that species interactions are not altered by  
500 warming at this treeline site and, thus, the performance of these populations will probably not change  
501 due to mild warming in the near future.

502

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511

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513

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## Tables

Table 1. Results of ANOVA for effects of warming and stand type on *Vaccinium myrtillus* height and above-ground (AG) biomass parameters at the ramet scale

AG Variable	Factor/covariate	df	F	P
Height	Age	1, 118	26.47	<0.001
	Warming	1, 24	5.33	0.030
	Stand type	2, 24	1.89	0.173
	Stand type x warming	2, 24	0.16	0.857
Leaf biomass	Age	1, 118	12.00	0.001
	Warming	1, 24	2.38	0.136
	Stand type	2, 24	1.49	0.246
	Stand type x warming	2, 24	1.20	0.320
New shoots biomass	Age	1, 118	8.68	0.004
	Warming	1, 24	5.02	0.035
	Stand type	2, 24	1.38	0.271
	Stand type x warming	2, 24	0.37	0.693
Total AG biomass	Age	1, 118	32.41	<0.001
	Warming	1, 24	4.74	0.040
	Stand type	2, 24	1.77	0.193
	Stand type x warming	2, 24	0.16	0.857
No. of shoots	Age	1, 114	23.45	<0.001
	Warming	1, 23	0.26	0.613
	Stand type	2, 23	0.84	0.443
	Stand type x warming	2, 23	0.83	0.450
Dry weight/shoot	Age	1, 114	2.87	0.093
	Warming	1, 23	3.88	0.061
	Stand type	2, 23	2.24	0.129
	Stand type x warming	2, 23	0.55	0.584

Table 2. Results of ANOVA testing the effects of warming (W), stand type (ST) and their interaction (W x ST) on the C and N concentration and isotope composition ( $\delta^{13}\text{C}$ ,  $\delta^{15}\text{N}$ ) and C/N ratio of *Vaccinium myrtillus* leaves, new shoots, rhizomes and roots. *F*-values and *P*-values (in parentheses) are given. Between-groups degrees of freedom were 1 for W, 2 for ST and 2 for ST x W. Within-groups degrees of freedom were 24, except for root N concentration and C/N (22), rhizome C and N concentration (22), root  $\delta^{15}\text{N}$  (23) and rhizome C/N (21)

Organ	Variable	W	ST	W x ST
Leaves	C concentration	0.93 (0.344)	1.51 (0.242)	1.57 (0.228)
	N concentration	0.02 (0.884)	4.93 (0.016)	0.07 (0.931)
	$\delta^{13}\text{C}$	0.90 (0.352)	2.72 (0.086)	0.85 (0.441)
	$\delta^{15}\text{N}$	0.08 (0.780)	10.28 (0.001)	0.04 (0.960)
	C/N	0.06 (0.796)	3.74 (0.038)	0.23 (0.798)
New shoots	C concentration	1.68 (0.207)	0.94 (0.404)	0.94 (0.404)
	N concentration	0.07 (0.793)	0.63 (0.540)	0.77 (0.472)
	$\delta^{13}\text{C}$	0.07 (0.794)	8.16 (0.002)	1.85 (0.179)
	$\delta^{15}\text{N}$	0.33 (0.571)	9.39 (0.001)	0.00 (1.000)
	C/N	0.70 (0.410)	0.84 (0.444)	0.23 (0.794)
Rhizomes	C concentration	5.71 (0.026)	0.33 (0.723)	0.7 (0.509)
	N concentration	0.05 (0.829)	0.46 (0.637)	2.57 (0.099)
	$\delta^{13}\text{C}$	0.42 (0.522)	8.78 (0.001)	0.03 (0.972)
	$\delta^{15}\text{N}$	0.02 (0.884)	6.53 (0.005)	0.08 (0.921)
	C/N	2.71 (0.114)	0.26 (0.775)	2.70 (0.091)
Roots	C concentration	0.21 (0.653)	0.43 (0.656)	0.56 (0.578)
	N concentration	1.69 (0.207)	0.62 (0.545)	1.19 (0.323)
	$\delta^{13}\text{C}$	1.59 (0.218)	0.15 (0.860)	2.53 (0.101)
	$\delta^{15}\text{N}$	0.21 (0.650)	2.04 (0.153)	0.19 (0.826)
	C/N	1.58 (0.222)	0.31 (0.739)	0.38 (0.688)

Table 3. Results of ANOVA or Wilcoxon tests for the effects of warming on the C and N concentration and isotope composition ( $\delta^{13}\text{C}$ ,  $\delta^{15}\text{N}$ ) of *Vaccinium uliginosum* leaves, new shoots, rhizomes and roots

Organ	Parameter	<i>df</i>	<i>F/W</i>	<i>P</i>
Leaves	C concentration	1, 6	3.89	0.096
	N concentration	1, 6	2.57	0.160
	$\delta^{13}\text{C}$	1, 6	0.01	0.934
	$\delta^{15}\text{N}$	-	<i>W</i> = 6	0.686
New shoots	C concentration	-	<i>W</i> = 6	0.686
	N concentration	1, 6	13.91	0.010
	$\delta^{13}\text{C}$	1, 6	0.03	0.871
	$\delta^{15}\text{N}$	1, 6	0.35	0.575
Rhizomes	C concentration	1, 6	0.99	0.357
	N concentration	-	<i>W</i> = 8	1.000
	$\delta^{13}\text{C}$	-	<i>W</i> = 8	1.000
	$\delta^{15}\text{N}$	-	<i>W</i> = 8	1.000
Roots	C concentration	1, 6	0.00	0.997
	N concentration	1, 6	0.93	0.373
	$\delta^{13}\text{C}$	1, 6	1.08	0.339
	$\delta^{15}\text{N}$	1, 6	3.86	0.097

1    **Figure legends**

2    **Figure 1.** Effects of stand types (ST) and warming treatment (W) on a-c) aboveground and d-f)  
3    belowground biomass (means + 1 SE; n = 5) of *Vaccinium myrtillus*. Stand types included: *V.*  
4    *myrtillus* pure stands (M); *V. myrtillus* mixed with *R. ferrugineum* stands (R) and *V. myrtillus* mixed  
5    with *V. uliginosum* stands (U). Note that BG biomass data are reported as unit volume of soil at the  
6    plot scale. Symbols indicate significance levels: ( $\dagger$  0.1 > P > 0.05; \* 0.05 > P > 0.01; \*\* P < 0.01)

7

8    **Figure 2.** Effects of stand types (ST) and warming treatment on C and N concentrations and isotope  
9    compositions ( $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$ ) of leaves, new shoots, rhizomes and roots (mean  $\pm$ 1 SE; n = 5) of *V.*  
10   *myrtillus* organs in different stand types (ST) and warming (W) treatments. Stand types included: *V.*  
11   *myrtillus* pure stands (M); *V. myrtillus* mixed with *R. ferrugineum* stands (R) and *V. myrtillus* mixed  
12   with *V. uliginosum* stands (U). Symbols indicate significance levels: ( $\dagger$  0.1 > P > 0.05; \* 0.05 > P >  
13   0.01; \*\* P < 0.01). Y-axis values for nitrogen concentration in roots differ from the rest of the tissues,  
14   and are indicated at the right side of the panel

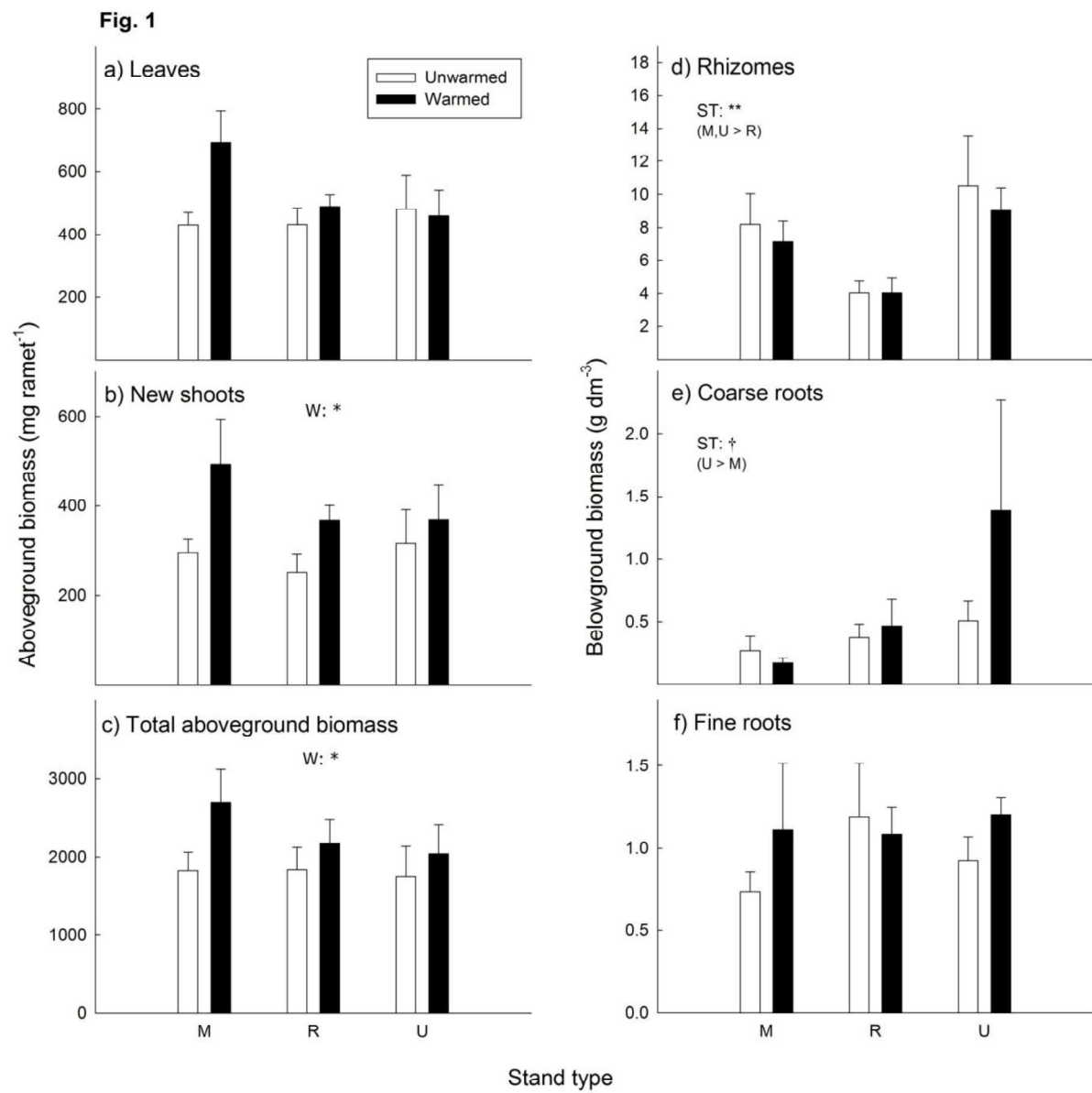
15

16   **Figure 3.** Effects of warming treatment on a) C and b) N concentrations and c)  $\delta^{13}\text{C}$  and d)  $\delta^{15}\text{N}$  of *V.*  
17   *uliginosum* organs (leaves, shoots, rhizomes and roots; mean  $\pm$ 1 SE; n = 4). Symbols indicate  
18   significance levels: ( $\dagger$  0.1 > P > 0.05; \* 0.05 > P > 0.01; \*\* P < 0.01)

19

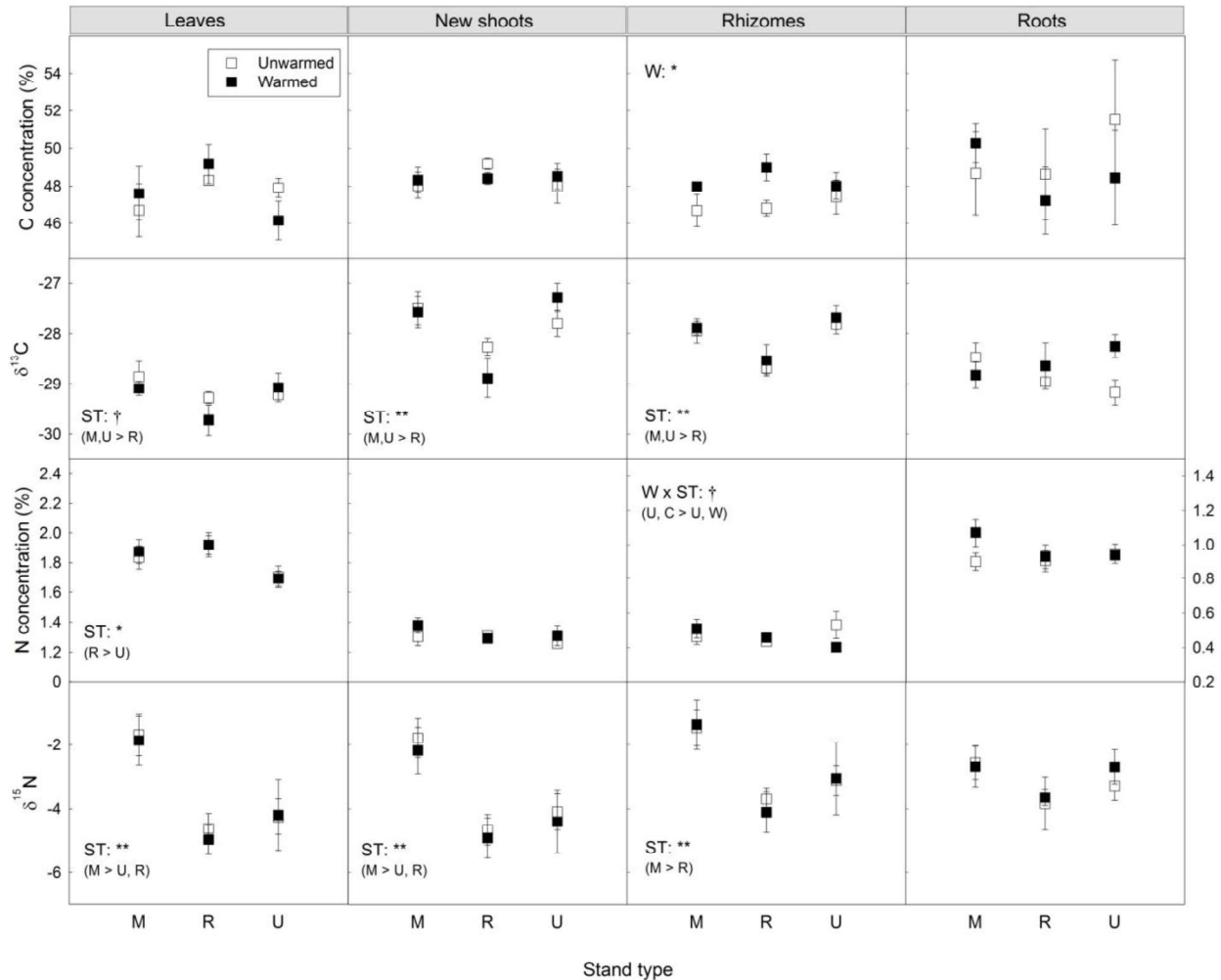
20   **Figure 4.** Soil a) nitrate ( $\text{NO}_3^-$ ) and b) ammonia ( $\text{NH}_4^+$ ) concentration in our study plots in different  
21   stand types (ST) and warming treatments in September 2013 (W; n = 5, means + 1 SE are shown).  
22   Stand types included: *V. myrtillus* pure stands (M); *V. myrtillus* mixed with *R. ferrugineum* stands (R)  
23   and *V. myrtillus* mixed with *V. uliginosum* stands (U). Asterisks (\*\*) show significant differences at P  
24   < 0.01

**Figure 1**

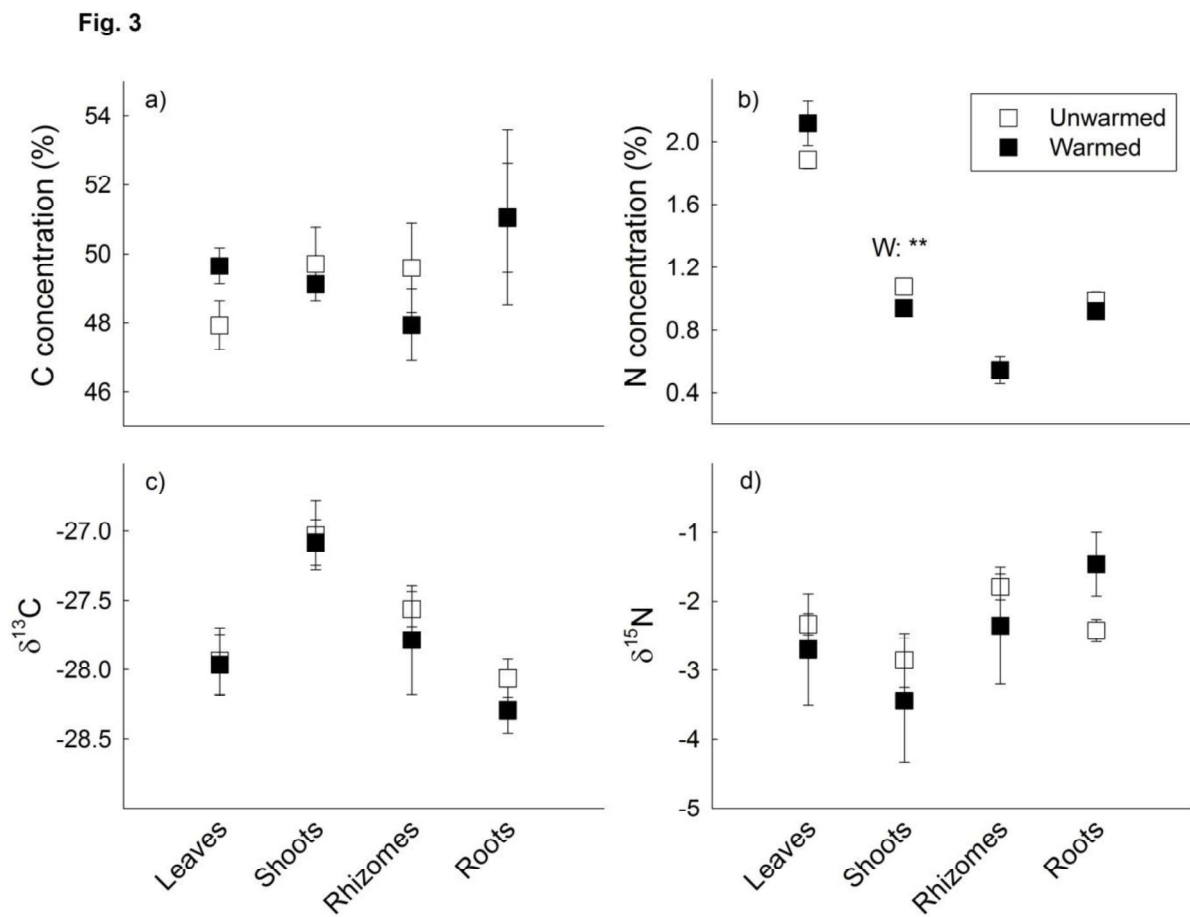


**Figure 2**

**Fig. 2**



**Figure 3**



**Figure 4**

**Fig. 4**

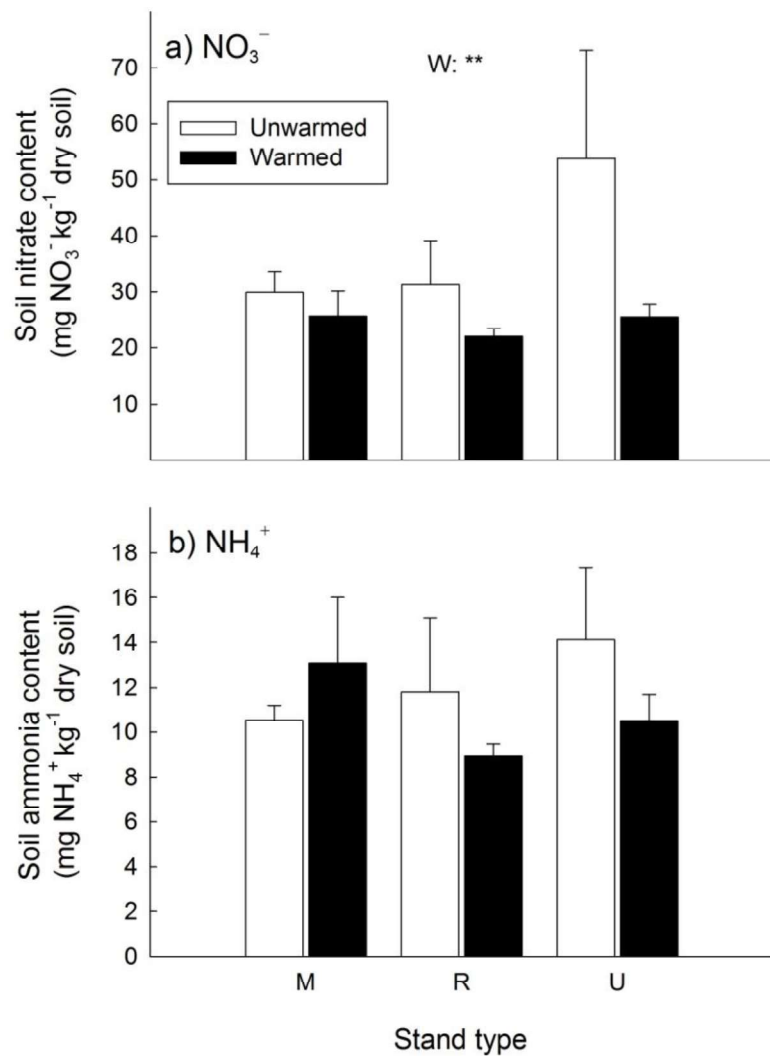


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Total AG biomass	Age	1, 118	32.41	<0.001
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	Stand type x warming	2, 24	0.16	0.857
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	Stand type	2, 23	2.24	0.129
	Stand type x warming	2, 23	0.55	0.584

Table 2. Results of ANOVA testing the effects of warming (W), stand type (ST) and their interaction (W x ST) on C and N concentration and isotope composition ( $\delta^{13}\text{C}$ ,  $\delta^{15}\text{N}$ ) and C/N ratio of *Vaccinium myrtillus* leaves, new shoots, rhizomes and roots. *F*-values and *P*-values (in parentheses) are given. Between-groups degrees of freedom were 1 for W, 2 for ST and 2 for ST x W. Within-groups degrees of freedom were 24, except for root N concentration and C/N (22), rhizome C and N concentration (22), root  $\delta^{15}\text{N}$  (23) and rhizome C/N (21)

Organ	Variable	W	ST	W x ST
Leaves	C concentration	0.93 (0.344)	1.51 (0.242)	1.57 (0.228)
	N concentration	0.02 (0.884)	4.93 (0.016)	0.07 (0.931)
	$\delta^{13}\text{C}$	0.90 (0.352)	2.72 (0.086)	0.85 (0.441)
	$\delta^{15}\text{N}$	0.08 (0.780)	10.28 (0.001)	0.04 (0.960)
	C/N	0.06 (0.796)	3.74 (0.038)	0.23 (0.798)
New shoots	C concentration	1.68 (0.207)	0.94 (0.404)	0.94 (0.404)
	N concentration	0.07 (0.793)	0.63 (0.540)	0.77 (0.472)
	$\delta^{13}\text{C}$	0.07 (0.794)	8.16 (0.002)	1.85 (0.179)
	$\delta^{15}\text{N}$	0.33 (0.571)	9.39 (0.001)	0.00 (1.000)
	C/N	0.70 (0.410)	0.84 (0.444)	0.23 (0.794)
Rhizomes	C concentration	5.71 (0.026)	0.33 (0.723)	0.7 (0.509)
	N concentration	0.05 (0.829)	0.46 (0.637)	2.57 (0.099)
	$\delta^{13}\text{C}$	0.42 (0.522)	8.78 (0.001)	0.03 (0.972)
	$\delta^{15}\text{N}$	0.02 (0.884)	6.53 (0.005)	0.08 (0.921)
	C/N	2.71 (0.114)	0.26 (0.775)	2.70 (0.091)
Roots	C concentration	0.21 (0.653)	0.43 (0.656)	0.56 (0.578)
	N concentration	1.69 (0.207)	0.62 (0.545)	1.19 (0.323)
	$\delta^{13}\text{C}$	1.59 (0.218)	0.15 (0.860)	2.53 (0.101)
	$\delta^{15}\text{N}$	0.21 (0.650)	2.04 (0.153)	0.19 (0.826)
	C/N	1.58 (0.222)	0.31 (0.739)	0.38 (0.688)

Table 3. Results of ANOVA or Wilcoxon tests for the effects of warming on the C and N concentration and isotope composition ( $\delta^{13}\text{C}$ ,  $\delta^{15}\text{N}$ ) of *Vaccinium uliginosum* leaves, new shoots, rhizomes and roots

Organ	Parameter	df	F/W	P
Leaves	C concentration	1, 6	3.89	0.096
	N concentration	1, 6	2.57	0.160
	$\delta^{13}\text{C}$	1, 6	0.01	0.934
	$\delta^{15}\text{N}$	-	$W = 6$	0.686
New shoots	C concentration	-	$W = 6$	0.686
	N concentration	1, 6	13.91	0.010
	$\delta^{13}\text{C}$	1, 6	0.03	0.871
	$\delta^{15}\text{N}$	1, 6	0.35	0.575
Rhizomes	C concentration	1, 6	0.99	0.357
	N concentration	-	$W = 8$	1.000
	$\delta^{13}\text{C}$	-	$W = 8$	1.000
	$\delta^{15}\text{N}$	-	$W = 8$	1.000
Roots	C concentration	1, 6	0.00	0.997
	N concentration	1, 6	0.93	0.373
	$\delta^{13}\text{C}$	1, 6	1.08	0.339
	$\delta^{15}\text{N}$	1, 6	3.86	0.097

Figure 1

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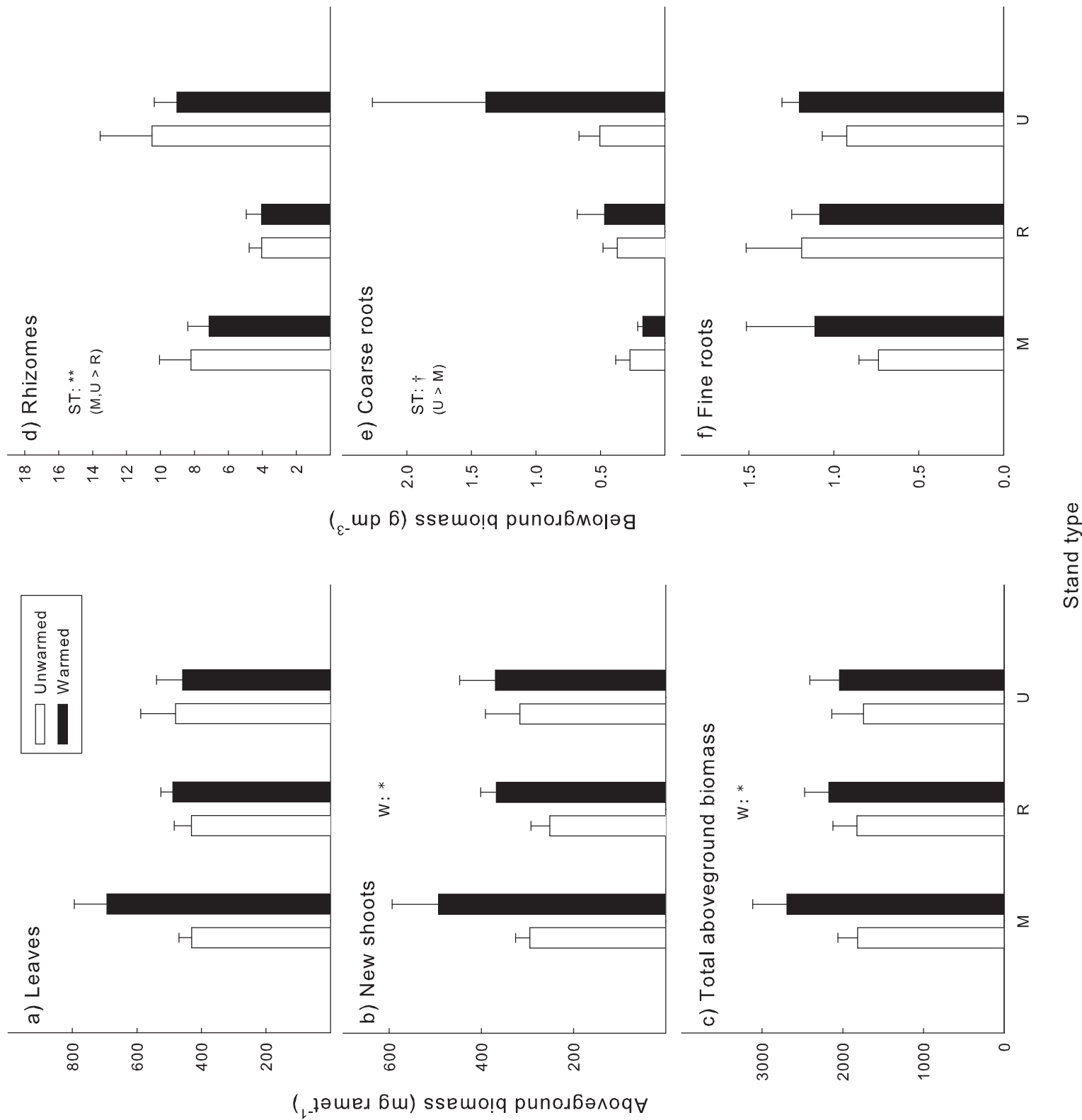
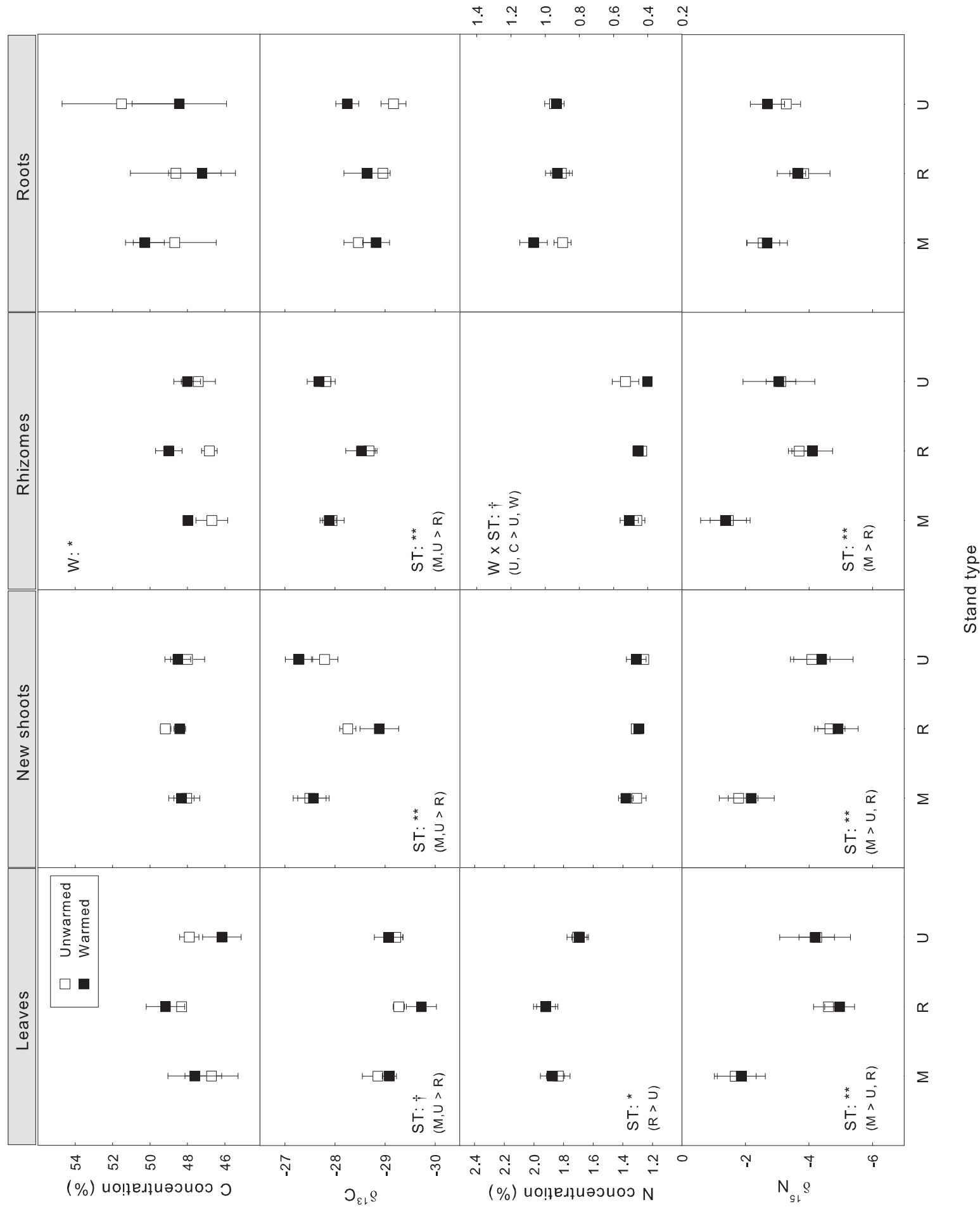
**Fig. 1**

Figure 2

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Fig. 2



**Fig. 3**

Figure 3

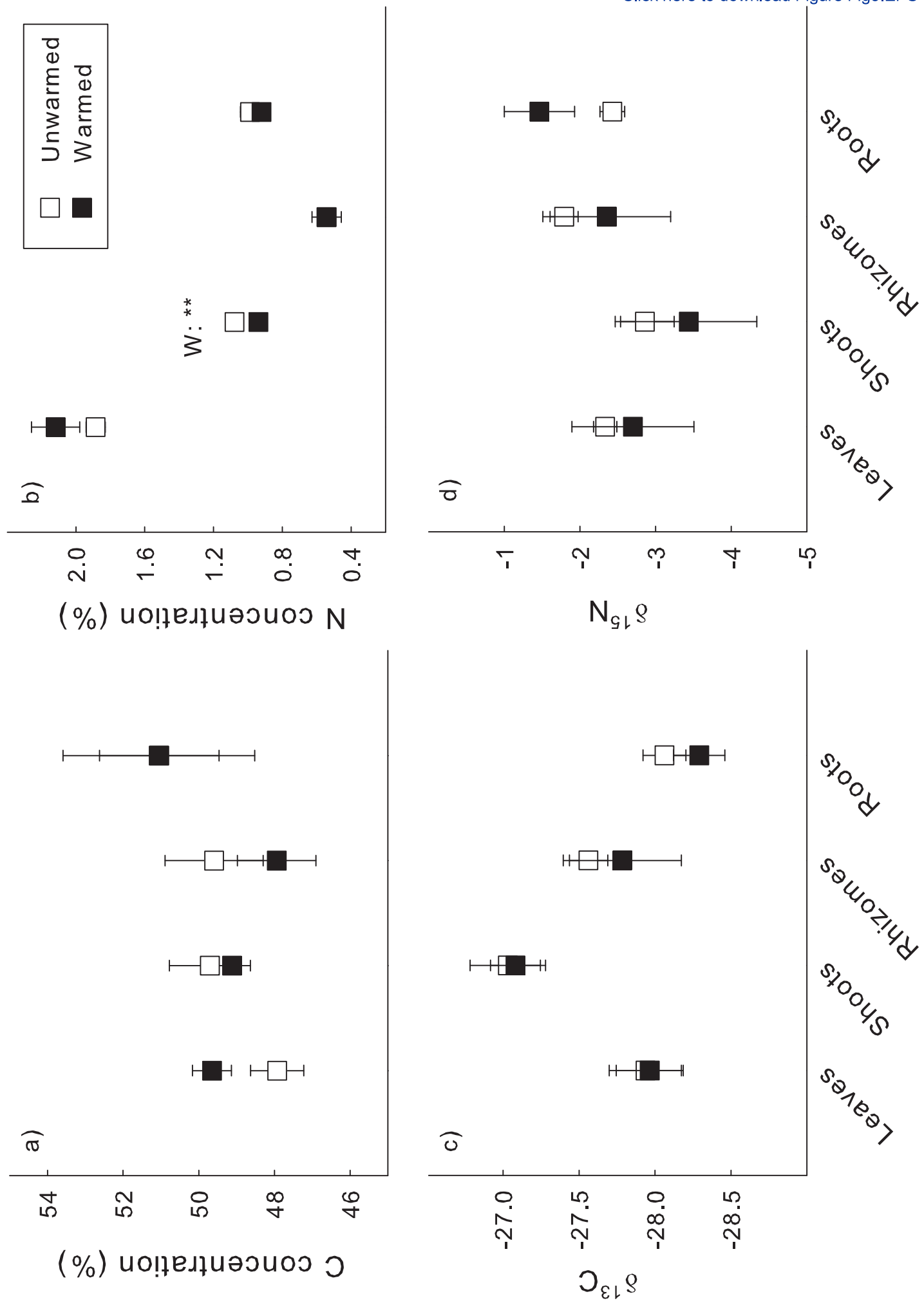
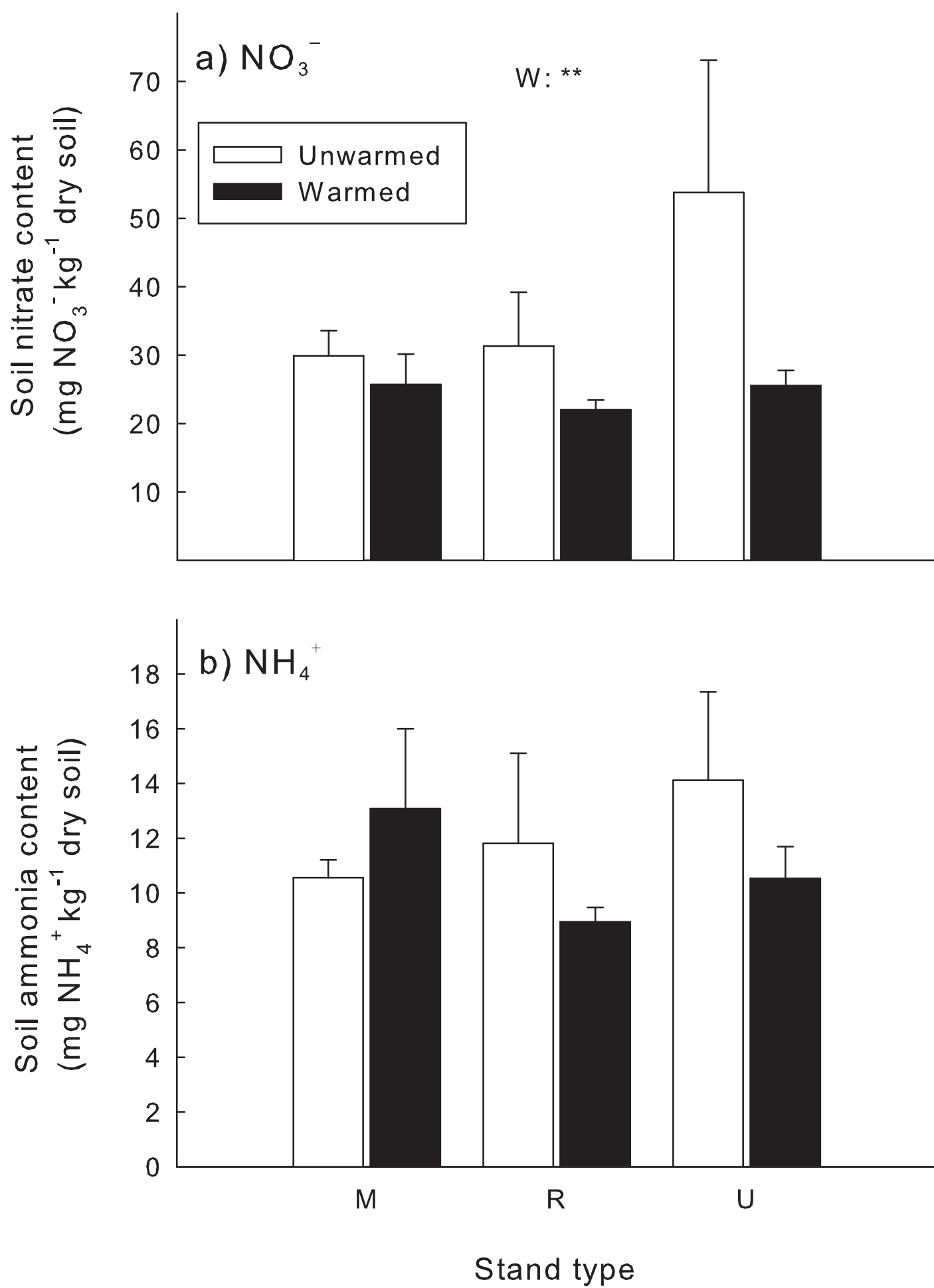
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Figure 4  
**Fig. 4**

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