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 This is a pre print version of the following article:

 Original Citation:

 Availability:

 This version is available http://hdl.handle.net/2318/1622737

 since 2017-01-20T11:42:51Z

 Published version:

 DOI:10.1016/j.funeco.2016.08.006

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This is the author's final version of the contribution published as:

[Nascimbene J., Ackermann S., Dainese M. Garbarino M., Carrer M. 2016. Finescale population dynamics help to elucidate community assembly patterns of epiphytic lichens in alpine forests. Fungal Ecology 24: 21-26. doi: 10.1016/j.funeco.2016.08.006.]

The publisher's version is available at:

[http://www.sciencedirect.com/science/article/pii/S1754504816300873]

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Link to this full text: [http://hdl.handle.net/2318/1622737]

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- Fine-scale population dynamics help to elucidate community assembly patterns of epiphytic
 lichens in alpine forests
- 3
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14

15 Abstarct

We examined the main and interactive effects of factors related to habitat filtering, dispersal 16 dynamics, and biotic interactions, on tree-level population dynamics of a subset of species 17 composing the epiphytic lichen pool in an alpine forest. We tested these processes evaluating the 18 population size of 14 lichen species on six hundred and sixty-five trees within a 2 ha plot located in 19 a high elevation alpine forest of the eastern Italian Alps. Our results indicate that community 20 assembly patterns at the tree-level are underpinned by the simultaneous effects of habitat filtering, 21 dispersal, and biotic interactions on the fine-scale population dynamics. These processes determine 22 how the single species are sorted into community assemblages, contributing to tree-level 23 community diversity and composition patterns. This corroborates the view that the response of 24 lichen communities to environmental gradients, in terms of compositional and diversity shifts, may 25 reflect differential species responses to different drivers. 26

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Key-words: biotic interactions; dispersal dynamics; facilitation; habitat filtering; host tree; spatial
distribution; species distribution modelling

31 Introduction

32 Ecological communities are dynamic assemblages of species whose patterns in space and time are regulated by different interacting processes. The relative importance of these processes may depend 33 on environment type, organism traits, and spatial scale of analysis (e.g. Guisan & Thuiller 2005). 34 Habitat filtering, dispersal dynamics, and biotic interactions are the main processes that interact to 35 determine community patterns (Lortie et al. 2004). Habitat filtering, invoked in niche-based models, 36 37 emphasizes the role of environmental factors and habitat quality in determining species distribution patterns, especially at the fine-scale. Among the stochastic processes (i.e., neutral theory of 38 biodiversity; Hubbell 2001), dispersal dynamics play a crucial role in promoting and maintaining 39 40 diversity, acting mainly at broad spatial scale (Wiszt et al. 2013). However, according to the metapopulation theory (Hanski 1999), dispersal processes could be also influential at the fine-scale. 41 Habitat connectivity and dispersal traits may influence species patterns due to the dynamics of their 42 43 patches (Snäll et al. 2004; Snäll, Ehrlén, & Rydin 2005). For instance, poorly dispersed species may be negatively affected by scarce connectivity among habitat patches. Biotic interactions are 44 45 recognized to contribute to community structure, both at fine and broad spatial scale (Wiszt et al. 2013), based on the concept that species are not stand-alone entities and interact positively (e.g., 46 facilitation) and/or negatively (e.g., competitive exclusion) with other species. Interactions 47 48 influence the patterns of each species and, in turn, influence community patterns (Wiszt et al. 2013). The relative effects of these three processes could also depend on the species, or on the 49 successional stage of the habitat. For instance, stochastic processes such as dispersal can initially 50 determine which species arrive at a particular site, while non-random processes, such as habitat 51 filtering or biotic interactions, can determine the persisting of the species (Lortie et al. 2004). 52 Fine resolution studies, which simultaneously investigated the role of habitat filtering, dispersal 53 54 dynamics, and biotic interactions are almost lacking for epiphytic lichens (Ellis 2012), one of the most diverse and functionally important forest organisms. The evaluation of the processes 55 determining their distribution patterns may provide information to prevent loss of forest diversity 56

and ecosystem functions. There is evidence that in forest ecosystems lichen patterns are influenced 57 58 by host tree features, such as tree species, size, age, crown dimension (e.g. Nascimbene et al. 2009; Nascimbene, Marini, Nimis, 2009) and microclimatic conditions (Nascimbene, Marini, Ódor 2012). 59 However, dispersal dynamics may also play a key role resulting in patterns that could differ 60 between spore- and vegetatively-dispersed species (Löbel, Snäll & Rydin 2006a). The former are 61 62 considered good dispersers due to the small size of the spores, while vegetatively-dispersed species 63 have lower dispersal capacity due to the larger size of vegetative propagules (e.g. Werth *et al.* 2006). For these species, patch connectivity could be important even at a fine spatial scale, since the 64 establishment and development of a population are density-dependent processes affected by 65 66 distance and size of propagule sources. The role of biotic interactions in structuring lichen communities is scarcely explored (Ellis 2012), although autogenic processes such as competition 67 and facilitation are likely to contribute in determining lichen patterns. For example, along a 68 69 chronosequence small and slow growing crustose lichens could be outcompeted by large and fast 70 growing macrolichens, favoring an ecological succession. Facilitation was never demonstrated for 71 epiphytic lichen communities (e.g., Belinchón et al. 2012), though biotic interactions are considered to be important drivers of lichen structure (Maestre et al. 2008). For instance, in soil lichen 72 communities facilitation would be dominant under stressful conditions (Maestre et al. 2008, 2009), 73 74 or moss carpets are known to improve the performance of high humidity demanding species (Öckinger, Niklasson & Nilsson 2005), or photobiont sharing (Rikkinen, Oksanen, & Lohtander 75 2002) is a plausible mechanism contributing to the success of the species in forest ecosystems. 76 This research aims at reacting to a scarcity of studies simultaneously incorporating the analysis of 77 different processes (namely habitat filtering, dispersal dynamics, and biotic interactions) in shaping 78 lichen distribution, explicitly dealing with spatial patterns (see e.g. Schei et al. 2012). Our study 79 80 focuses on fine-scale patterns of selected species in a high elevation Alpine forest, using a fine resolution analysis that is expected to be highly predictive for fixed epiphytic organisms (Guisan & 81 Thuiller 2005). After a preliminary analysis describing the main spatial patterns of the species 82

(clumped vs random) we explicitly tested the influence of factors indicative of different processes 83 84 and evaluated their relative importance in determining species patterns. Specifically, we hypothesized that: (i) Habitat filtering is the main process determining fine-scale lichen distribution 85 due to the influence of multiple drivers related to tree features and microclimate. We expected that 86 habitat filtering influences all the species, even if associated with different drivers. (ii) The 87 relevance of dispersal dynamics and biotic interactions should be species-specific. Specifically, we 88 89 expected that the dispersal dynamics depended on the dispersal traits of the species. For example, vegetatively-dispersed species, having lower dispersal capacity, should be positively affected by 90 patch connectivity. Considering biotic interactions, positive interactions (i.e. facilitation) may 91 92 explain the pattern of the most abundant species that usually co-occur on the same tree. Conversely, negative interactions (i.e. competition) may explain the pattern of ecologically more demanding 93 species that are likely to be outcompeted by more plastic lichens, especially in benign environments 94 95 (Bertness & Callaway, 1994). (iii) The spatial distribution of lichens (clumped vs random) may correspond to a different response of the species to habitat filtering, dispersal dynamics, and biotic 96 97 interactions.

98

99 Materials and methods

100 *Study site*

101 The study site is a 2-ha plot located in the Italian Eastern Alps at an elevation of 1900 m a.s.l

102 (Latitude: 46.23 N; Longitude: 11.32 E; Figure 1). The climate is temperate-cold to continental,

103 characterized by strong daily and annual temperature fluctuations. Mean annual temperature is

4.6°C, while mean annual precipitation is c. 950 mm, with a peak during summer and a dip between

105 December and February. On average, a solid precipitation of 260 cm per winter period has been

recorded at the nearest nivological station of Obereggen (1872 m a.s.l.), forming a permanent snow

107 cover during 110-131 days per year.

108 Vegetation belongs to Vaccinio-Piceetea (Larici-Cembretum), with Norway spruce (*Picea abies*

109 (L.) Karst.), stone pine (Pinus cembra L.), and European larch (Larix decidua Mill.) as main tree

species. The shrub layer is mainly composed of *Daphne striata*, *Juniperus communis* subsp. *alpina*,

111 Rhododendron hirsutum and R. ferrugineum, Ribes alpinum, Vaccinium myrtillus and V. vitis-idaea

and the herbal layer of Adenostyles alliariae, Calamagrostis villosa, Luzula sylvatica,

113 Maianthemum bifolium, Melampyrum sylvaticum, Petasites albus, Saxifraga sp.

The area is subject to the typical dynamics of many high-elevation forests in the Alps, where the significant reduction of livestock activities and the decreased intensity of silvicultural practices during the last centuries triggered a change in forest composition where larch, the initial dominant species, is decreasing its presence respect to stone pine and spruce (Carrer & Urbinati 2001). These dynamics couple with increasing tree density and canopy closure. Management activities ceased in the 90s and currently the area is completely left to natural evolution and used for long-term ecological studies.

121

122 Sampling design and data collection

All the trees taller than 130 cm were mapped with a total station and georeferenced using an 123 electro-optical distance meter and their species (Figure 1), DBH and crown dimension recorded. 124 125 Tree age has been also determined through increment coring. Further details on the sampling protocol for forest structure can be found in Carrer & Urbinati (2001) and in Carrer, Soraruf. & 126 Lingua (2013). After an exhaustive floristic survey that yielded 84 species (Nascimbene 2013), we 127 selected a subset of 14 species (Table 1). Precondition to be included in our sampling design was 128 that the lichen species could be readily identified in the field with naked eye or the help of a 129 magnifier. The species were also selected as to represent different dispersal strategies, including 130 both sexually (i.e. by spores) and vegetatively (i.e. by lichenized propagules) dispersed species. 131 On each tree with a DBH >15cm, the abundance of each species was estimated as value of total 132 coverage (in cm²) on the stem surface up to a height of 1.80 m. Six hundred-sixty-five trees were 133

surveyed, including 311 spruce, 239 stone pine, and 115 larch. The lichen survey was carried out insummer 2012.

136

137 *Explanatory variables*

We quantified 7 explanatory variables indicative of three different processes: environmental 138 filtering, dispersal dynamics, and biotic interactions (see Appendix S1 in Supporting Information). 139 140 To account for the environmental filtering process we selected four tree-level variables that are known to be among the most meaningful descriptors of forest lichen patterns: tree species, tree size 141 (DBH), tree age, and crown volume. As a proxy for micro-topographic conditions, we calculated a 142 143 curvature index in a GIS environment. A fine resolution (1-m) DEM was computed by using geographic position (x, y) and elevation (z) of each mapped tree. In this way, we were able to assess 144 if a given tree was located on a linear, concave or convex surface. 145

For each lichen species on each tree we quantified the Incidence Function Model (IFM; Hanski 147 1994) to account for dispersal dynamics. The IFM is a typical connectivity measure used in 148 metapopulation ecology (Moilanen & Nieminen 2002). Connectivity (S_i) was calculated for each 149 tree by

150
$$S_i = \Sigma_{j \neq i} \exp(-\alpha \, d_{ij}) A_j$$

where d_{ij} is the Euclidean distance between the tree *i* and each neighbor *j* and *A* the surface area occupied by a lichen species on a tree trunk. The parameter α was estimated separately for each species based on tree occupancy data, by testing different α values and selecting the value that gave the best model fit in a logistic regression model (Oksanen 2004; Jönssonn, Edman & Jonsson 2008). The value of S_i was computed using the software R version 2.15.2 (R Core Team 2012) with the add-on package 'metapop' (Oksanen 2004).

157 For each species at tree level we quantified the cover of the co-occuring species assuming lichen158 cover to be a reasonable proxy for biotic interactions (Roux et al. 2014).

159

160 *Statistical analyses*

To disentangle the different distribution behavior of the lichens we performed a preliminary analysis to test the spatial autocorrelation of the distribution patterns. We used the Moran's I index, a global index which computes the degree of correlation between the values of a variable (in our case, the abundance) as a function of spatial lags (Fortin, Dale & ver Hoef 2002). The analyses were computed with a lag distance of 10 m, up to 100 m that corresponds to the shortest size of the plot. We considered values of |z(I)| > 1.96 (p < 0.05).

Depending on the occurrence of the lichen species, two different approaches were used to test the 167 effect of environmental filtering, dispersal dynamics, and biotic interactions on lichen cover. The 168 169 following covariates were included in the models: tree species, age, DBH, crown volume, curvature, connectivity, and lichen cover. We also tested the interaction between age and DBH. Given the 170 structure of our data (skew distribution), we opted to use generalized linear models (GLM). For 171 172 common species (n = 5; frequency > 44%), lichen cover was analyzed using GLM with a negative binomial distribution to account for the overdispersion of the data (Zuur et al. 2009) (see Appendix 173 174 S2). For relatively rare species (n = 9; frequency < 43%) with an excess of zero cases in the dataset, a hurdle regression model was performed (also called zero-altered or two-part models; Zuur et al. 175 2009). Ignoring zero inflation can create problems in model inference by biasing the estimated 176 177 parameters and standard errors, as well as overdispersion (Martin et al. 2005; Zuur et al. 2009). In our case, the zero inflation was the result of a large number of 'true zero' observations caused by 178 the real ecological effect of interest (i.e., unsuitable habitat; Martin et al. 2005) Specifically, we 179 applied a zero-truncated negative binomial (ZANB) to account for the overdispersion of the data 180 (for more details of the models specification see Appendix S2). In both the models (GLM and 181 ZANB), all predictors were standardized by mean-centering and dividing by two standard 182 deviations to improve interpretability of parameter estimates, particularly when interactions were 183 fitted, and continuous and categorical factors are combined in the same model (Gelman 2008). 184 Negative binomial GLM was analyzed using the 'MASS' package (Venables & Ripley 2002) in R, 185

while the hurdle model using the 'pscl' package (Zeileis, Kleiber & Jackman 2008; Jackman 2012)in R.

We used an information-theoretic model selection procedure to evaluate alternative competing 188 models (Burnham & Anderson 2002). We compared the fit of all possible candidate models 189 obtained by the combination of the predictors using second-order Akaike's information criterion 190 (AICc). Models were chose that differed from the AICc of the best fitting model by < 4. We used 191 192 the Akaike weights (w_i) to measure the relative importance of each predictor, summing the w_i across the models $(\sum w_i)$ in which the predictor occurred. For each parameter, we used model averaging in 193 order to incorporate model selection uncertainty into our parameter estimates (Burnham & 194 195 Anderson 2002; Grueber et al. 2011). Individual predictor variables that had an Akaike weight > 0.75 or model averaged confidence intervals that did not include 0 were considered as most 196 important predictors. Model comparison was implemented using the 'MuMIn' package (Barton 197 198 2013) in R. Finally, the variation in lichen cover was decomposed for each species using a series of (partial) 199

200 regression analyses implemented in the 'vegan' package for R (Oksanen et al. 2013). The total explained variation (TVE) in lichen cover was partitioned into seven components (adjusted R^2 ; 201 Peres-Neto et al. 2006): the pure effect of environmental filtering (E), dispersal dynamics (D), and 202 203 biotic interactions (B); three first-order joint components ($E \cap D$, $E \cap B$, $D \cap B$); and the joint component among the three groups $(E \cap D \cap B)$. In the environmental filtering component (E) we 204 included tree species, tree size (DBH), tree age, canopy volume, and curvature. In the dispersal 205 dynamics components (D) we included the connectivity index, while in the biotic interactions 206 component (B) the cover of the other lichen species. 207

All the statistical analyses were performed separately for each species.

209

210 **Results**

211 Lichen species and spatial patterns

The 14 species widely differed in frequency (Table 1), ranging from a minimum of 3% of colonized
trees for *Letharia vulpina* up to a maximum value of 97,6 % for *Parmeliopsis ambigua*. Three
species were extremely common, since they were recorded on more than 90% of the trees, while
five species were relatively rare, being recorded on less than 20% of the trees.
After the spatial autocorrelation analysis the species were equally distributed in two groups (Table
1; Appendix S2): i) lichens with a clumped spatial pattern and ii) lichens with a random spatial
pattern. Both groups included vegetatively- and spore-dispersed species.

219

220 Drivers of fine-scale lichen patterns

221 Among the variables related with habitat filtering, tree species was by far the most important for both clumped and randomly distributed groups (Figures 2, 3; Appendix S4, S5), only the extremely 222 generalist species *Hypogymnia physodes* being not significantly influenced by this factor. Age and 223 224 DBH mainly influenced clumped distributed species also by a significant interaction (Figure 4; Appendix S6). While DBH had in general a positive effect (except for one species), Age had 225 226 contrasting effects with species preferring either young or old trees. Crown dimension had a significant influence on only two species with clumped distribution, with contrasting effects. 227 For the remaining exploratory variables: i) microclimatic conditions, as inferred by the effect of 228 microtopography, influenced the distribution of five species, two of them preferring trees located on 229 exposed sites (i.e. relatively sun exposed and dry conditions) and three of them preferring trees in 230 small depressions, i.e. sheltered and humid conditions (Figures 2, 3; Appendix S4, S5); ii) dispersal 231 dynamics, as inferred by the role of connectivity, influenced the distribution of five species, mainly 232 reproducing by vegetative propagules (4 species), including two randomly distributed lichens 233 (Figures 2, 3; Appendix S4, S5). 234

Biotic interactions, as inferred by the role of lichen cover, positively influenced four common
species with clumped distribution, while had a negative effect on one relatively rare species with
random distribution (Figures 2, 3; Appendix S4, S5).

239 The relative role of environmental filtering, dispersal dynamics and biotic interactions The variation partitioning analysis indicated that the total variation in species abundance patterns 240 explained by the models was higher for clumped species (explained variation range between 6 and 241 242 37%) than for randomly distributed species (explained variation range between 3 and 12%) (Table 2). Habitat filtering was the most important process for almost all the species, except for 243 244 Hypogymnia physodes and Pseudevernia furfuracea for which biotic interaction was the main process determining their patterns (explained variation 15% and 17%, respectively) with an 245 additional impact of the shared component between biotic interaction and environmental filtering, 246 247 summing up to 12% of the total variance. Dispersal dynamics seemed to have a negligible influence in shaping lichen distribution in our study system (explained variation range between 1 and 3%). 248 249

250 Discussion

Our results reveal that habitat filtering is the main process accounting for the fine-scale patterns of our selected species, indicating that habitat features are the main drivers of lichen distribution for both clumped and randomly distributed species. Dispersal dynamics and biotic interactions play a significant role mainly for clumped species.

255

256 Habitat filtering

Tree species is the most important environmental factor whose effect is mainly related with speciesspecific differences in the chemical and physical traits of the bark, chiefly pH and texture (e.g. Fritz & Heilmann-Clausen 2010; Király *et al.*, 2013). These differences may be relevant even among relatively similar host trees, such in the case of our three coniferous species. Besides tree species, tree size and age are also important drivers of local lichen patterns (Nascimbene *et al.* 2009), with both direct and interactive effects. According to an 'area effect', tree size positively influences abundance patterns fostering the population size. Tee age seems to have species-specific effects

with some lichens alternatively preferring young or old trees, according with either a pioneer or a 264 265 late-successional behavior. The interaction between tree size and tree age indicates a decrease of the positive effect of tree size on lichen cover with increasing tree age, even to become neutral on older 266 trees (> 180 years). On these old trees, lichen dynamic are more influenced by a 'time per se' effect 267 (i.e. time available for colonization and increase of population size) than by an 'area effect'. In 268 addition, tree size gains importance on large trees, while age gains importance on small ones, 269 corroborating the hypothesis that the 'area effect' and "time per se" effect are two complementary 270 mechanisms influencing lichen patterns in forest ecosystems. Crown dimension influenced the 271 distribution of only two species, with contrasting effects. Chaenotheca chrysocephala, preferring 272 273 environmental conditions protected from rain, was positively affected by crown dimension, while Hypogymnia physodes, which prefers well-lit conditions, was negatively affected by this driver. In 274 general, the effect of this tree level factor is poorly explored in the lichen literature (e.g. 275 276 Nascimbene et al. 2008), although it is likely to interact with dispersal dynamics and to influence microclimatic conditions (e.g. Nascimbene et al. 2008, 2009). Further evidence for the importance 277 278 of microclimatic conditions (Nascimbene, Marini & Ódor 2012) is provided by the significant 279 contribution of microtopography to the abundance pattern of five species, discriminating between those preferring very humid-shaded (e.g. Schismatomma pericleum) or relatively dry and well-lit 280 conditions (i.e. Pseudevernia furfuracea). 281

282

283 Dispersal dynamics and biotic interactions

Dispersal dynamics scarcely affect the abundance patterns of our selected species. This confirms the
hypothesis that dispersal plays a minor role in determining fine-scale patterns (Schei *et al.* 2012).
However, the positive effect of habitat connectivity on several vegetatively-dispersed species
suggests a trait-mediated response predicting that at fine-scale dispersal dynamics are influential for
poor dispersers (Löbel, Snäll & Rydin 2006b). For these species, an excessive distance from

propagule sources may hinder the density-dependent processes of establishment and populationgrowth.

A positive effect of biotic interactions was found for the most common species, indicating 291 that their success may depend on some kind of facilitation. To the best of our knowledge, this is the 292 first time that this processes is detected for epiphytic lichens, although, our data did not allow a 293 direct evaluation of the mechanisms behind this effect (e.g., Belinchón et al. 2012). Anyway, the 294 295 high relative importance of biotic interactions in explaining the abundance patterns of *Hypogymnia* physodes and Pseudevernia furfuracea suggests that photobiont sharing could be a plausible 296 mechanism. Indeed, these two species host phylogenetically close-related photobionts (Hauck, 297 298 Helms & Friedl 2007) that could be alternatively used to promote their occurrence along wide ecological gradients (Blaha, Baloch & Grube 2006). Contrary to our expectations, we found low 299 support to competition hypothesis. We found indeed that only one species, such as 300 301 Tuckermannopsis chlorophylla, showed a negative effect of biotic interactions. Also, the effect of biotic interactions seems to play a minor role in determining fine-scale patterns. This could be due 302 to a presence of moderate stress levels in the study area that determine a neutral effect of biotic 303 interactions, as suggested by Maestre et al. (2009a, b). 304

305

306 *Clumped vs randomly distributed species*

The two different patterns of abundance distribution (clumped vs random) correspond to a different 307 response of the species to the drivers indicative of the three processes. The group of clumped 308 species includes lichens that are very common in different types of alpine forests (Nascimbene, 309 310 Nimis & Dainese 2014) where they constitute the keystones of epiphytic lichen communities. Our results indicate that their patterns are determined by a multiple and complex (i.e. interactive effects) 311 response to several drivers indicative of habitat filtering, dispersal and biotic interactions. This 312 complex behavior may ensure a high degree of adaptation enhancing the resistance and resilience of 313 their populations to forest dynamics induced by natural and anthropogenic disturbances. On the 314

contrary, the group of randomly-distributed species mainly includes relatively rare lichens that have 315 316 more specific ecological requirements (Nascimbene, Nimis & Dainese 2014). Their abundance patterns are ruled by more simple dynamics, mainly related to habitat filtering. In particular, their 317 strict dependence on the host tree species suggests that they are strongly influenced by tree 318 319 dynamics. In our dynamic forest, species related with open, larch-dominated stands (e.g. Letharia *vulpina*; Nascimbene, Nimis & Dainese 2014) could be relicts restricted to remnant patches whose 320 321 connectivity is fundamental for their maintenance, such in the case of Tuckneraria laureri and Tuckermannopsis chlorophylla. On the other hand, spruce-related species (e.g. Schismatomma 322 pericleum; Nascimbene, Nimis & Dainese 2014) may be in an expansion phase enhanced by the 323 324 increasingly available substrate.

325

326 Conclusions

The insights provided by this study on the processes determining fine-scale spatial patterns of 327 epiphytic lichens may contribute to a more conservation-oriented forest management. The high 328 dependence of lichen patterns on habitat filtering highlights the importance of forest management in 329 shaping the dynamics of these organisms at the local level. Indeed, most of the main factors 330 331 affecting habitat conditions relevant for lichens are controlled by management practices (Nascimbene, Thor & Nimis 2013). On this basis, conservation-oriented management should 332 improve local habitat heterogeneity favouring the coexistence of various tree species with different 333 334 size and age (i.e., mixed multi-layered and uneven-aged stands). Microtopography could further contribute to habitat heterogeneity, providing fine-scale variability of microclimatic conditions that 335 determine the local occurrence of species with different ecological requirements. Forest 336 337 management is also responsible for connectivity between trees, that favours the dispersal dynamics of several, mainly vegetatively dispersed, lichens. Yet, the relationships of many species with tree 338 dynamics suggest that habitat heterogeneity should be maintained also at the landscape level 339 enhancing the presence of forest patches at different successional stages. 340

Besides external processes, our study also highlights the importance of autogenic processes related with biotic interactions for few species. Research in this field is still in its infancy but promising results are expected from specific investigations aimed to reveal the biological mechanisms driving biotic interactions. The case of the photobiont sharing (Rikkinen, Oksanen & Lohtander 2002) that could explain the ecological plasticity of keystone species is just a first example.

346

347 Acknowledgements

The study was conducted in the framework of the project "Biodiversità, biomonitoraggio e conservazione dei licheni epifiti negli ambienti forestali della provincia di Bolzano", funded by the Autonomous Province of Bolzano (Ripartizione Diritto allo studio, Università e Ricerca scientifica). The Forest planning office of the Autonomous Province of Bolzano (project partner) is thanked for providing logistic and technical support. In particular, we are grateful to Günther Unterthiner, Fabio Maistrelli and their collaborators.

354

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- 478 SUPPORTING INFORMATION
- 479 Additional supporting information may be found in the online version of this article:480
- 481 **Appendix S1.** Descriptive statistics of the covariates.
- 482 Appendix S2. Description of the species distribution modeling.
- 483 Appendix S3. Moran's I correlograms.
- 484 Appendix S4. Model averaged coefficients for variables predicting species with clumped
 485 distribution.
- 486 Appendix S5. Model averaged coefficients for variables predicting species with random
- 487 distribution.
- 488 Appendix S6. The effect of tree age on the response of *Parmeliopsis ambigua* and *Vulpicida*
- 489 *pinastri* cover to tree size (DBH).

Table 1. Study species. Nomenclature and information on growth form and dispersal strategy were 491 retrieved from Nimis & Martellos (2008). In the last two columns the frequency (F) of the species 492 (expressed as percentage of trees on which they occurred) and the type of spatial pattern (SP; C = 493 clumped, R = random) are reported.

Species name	Dispersal strategy	F (%)	SP
Calicium viride Pers.	Sexual/spores	44.2	С
Chaenotheca chrysocephala (Ach.) Th.Fr.	Sexual/spores	68.1	С
Chaenotheca trichialis (Ach.) Th.Fr.	Sexual/spores	42.1	С
Evernia divaricata (L.) Ach.	Asexual/fragmentation	28	R
Hypogymnia physodes (L.) Nyl.	Asexual/soredia	97	С
Letharia vulpina (L.) Hue	Asexual/soredia	3.4	R
Parmeliopsis ambigua (Wulfen) Nyl.	Asexual/soredia	97.6	С
Platismatia glauca (L.) W. L. Culb. & C. F. Culb.	Asexual/isidia	12.6	R
Pseudevernia furfuracea (L.) Zopf	Asexual/isidia	90.1	С
Ramalina obtusata (Arnold) Bitter	Asexual/soredia	17.4	R
Schismatomma pericleum (Ach.) Branth & Rostr.	Sexual/spores	28.7	R
Tuckermannopsis chlorophylla (Willd.) Hale	Asexual/soredia	17.8	R
Tuckneraria laureri (Kremp.) Randlane & Thell	Asexual/soredia	18.6	R
Vulpicida pinastri (Scop.) J.E.Mattsson & M.J.Lai	Asexual/soredia	35.3	С

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507	Table 2. Variation partitioning of (a) species with clumped distribution and (b) species with random
508	distribution. The total variation explained was partitioned among environmental filtering (E),
509	dispersal dynamics (D), and biotic interactions (B). Values are adjusted R ² in %. Adjusted fractions
510	of total variation explained (TVE, in %) were estimated following the procedure of Peres-Neto et al.
511	(2006).

	Pure components		Shared components					
	Е	D	В	E∩D	E∩B	D∩B	E∩D∩B	TVE
(a) Species with clumped distribution								
Calicium viride	3	-	-	-	-	3	-	6
Chaenotheca chrysocephala	4	2	-	0	0	-	-	6
Chaenotheca trichialis	16	1	1	0	0	-	0	18
Hypogymnia physodes	4	-	15	0	12	-	0	31
Parmeliopsis ambigua	17	1	6	-	7	1	-	32
Pseudevernia furfuracea	8	-	17	-	12	-	-	37
Vulpicida pinastri	9	3	1	1	1	0	-	15
(b) Species with random distribution								
Evernia divaricata	3	-	1	-	2	-	-	6
Letharia vulpina	1	-	1	-	2	-	-	4
Platismatia glauca	4	-	0	-		-	0	4
Ramalina obtusata	12	-	-	0	0	0	-	12
Schismatomma pericleum	6	-	-	-		0	-	6
Tuckermannopsis chlorophylla	3	-	-	-	0	-	0	3
Tuckneraria laureri	3	-	-	-	0	-	0	3

515 **Figure captions:**

Figure 1. (a) Study area, (b) study site: a 2-ha plot located in the Italian Eastern Alps at an elevation
of 1900 m a.s.l (Latitude: 46.23 N; Longitude: 11.32 E).



Figure 2. Sum of model weights $(\sum w_i)$ for each variable estimated by the multi-model inference 518 519 procedure for species with clumped distribution. Predictors that consistently occurred in the most likely models ($\sum w_i > 0.75$) or model averaged confidence intervals that did not include 0 were 520 considered well supported by our data and considered as most important predictors (in grey). The 521 distribution of lichen species was modeled using hurdle regression (a-c) or GLM (d-g). The 522 direction of the relationship is indicated by (+) or (-) for continuous variables. For tree species, the 523 main host species, resulting from Tukey contrasts, is indicated: (L) larch, (P) stone pine, (S) spruce, 524 and (n.s.) not significant. 525



526

Figure 3. Sum of model weights $(\sum w_i)$ for each variable estimated by the multi-model inference 527 procedure for species with random distribution. Predictors that consistently occurred in the most 528 likely models ($\sum w_i > 0.75$) or model averaged confidence intervals that did not include 0 were 529 considered well supported by our data and considered as most important predictors (in grey). The 530 distribution of lichen species was modeled using hurdle regression (a-c and e-g) or GLM (d). The 531 direction of the relationship is indicated by (+) or (-) for continuous variables. For tree species is 532 indicated the main host species resulting from Tukey contrasts: (L) larch, (P) stone pine, (S) spruce, 533 and (n.s.) not significant. 534







SUPPORTING INFORMATION

Appendix S1. Descriptive statistics of the continuous factors used in the models.

	Mean \pm SD	Min	Max
(a) Environmental variables			
DBH (cm)	35.4 ± 11.9	6	70
Tree age (years)	149.1 ± 28.4	51	260
Crown volume (m ³)	63.6 ± 53.5	0.7	449.1
Curvature index	-1.3 ± 93.0	-541.8	487.8
(b) Dispersal dynamics (IFM)			
Calicium viride	0.4 ± 2.0	0.0	29.0
Chaenotheca chrysocephala	122.9 ± 389.1	0.0	6778.0
Chaenotheca trichialis	0.9 ± 3.9	0.0	51.2
Evernia divaricata	0.2 ± 1.0	0.0	13.0
Hypogymnia physodes	441.4 ± 1465.3	0.0	33399.2
Letharia vulpina	0.1 ± 0.5	0.0	7.1
Parmeliopsis ambigua	719.5 ± 1098.6	0.0	9679.3
Platismatia glauca	0.2 ± 0.8	0.0	9.3
Pseudevernia furfuracea	4.5 ± 8.1	0.0	62.3
Ramalina obtusata	0.1 ± 0.3	0.0	4.8
Schismatomma pericleum	0.2 ± 1.2	0.0	16.2
Tuckermannopsis chlorophylla	0.3 ± 1.1	0.0	10.8
Tuckneraria laureri	1.8 ± 8.2	0.0	95.9
Vulpicida pinastri	0.9 ± 4.1	0.0	90.7
(c) Biotic interaction (cover co-occurin	g species in cm ²)		
Calicium viride	5772.3 ± 5362.3	15	45520
Chaenotheca chrysocephala	5694.0 ± 5370.9	20	45515
Chaenotheca trichialis	5285.5 ± 5149.6	0	45520
Evernia divaricata	5994.9 ± 5392.0	20	45520
Hypogymnia physodes	3415.7 ± 3493.5	0	21250
Letharia vulpina	5996.7 ± 5388.4	20	45515
Parmeliopsis ambigua	3773.7 ± 4045.0	0	41505
Platismatia glauca	5984.2 ± 5387.8	20	45520
Pseudevernia furfuracea	5052.3 ± 4444.4	20	44020
Ramalina obtusata	6039.1 ± 5417.2	20	45520
Schismatomma pericleum	5647.8 ± 5259.6	20	45520
Tuckermannopsis chlorophylla	5992.2 ± 5388.4	20	45515
Tuckneraria laureri	5992.4 ± 5387.2	20	45520
Vulpicida pinastri	5994.6 ± 5388.5	20	45515

Appendix S2. Description of the species distribution modeling.

Three steps were considered in defining the GLM models (see Zuur et al. 2009): (i) the choice of the distribution for the response variable (Y_i) and the definition of its mean and variance; (ii) the definition of a predictor function specifying the covariates; and (iii) the link between the predictor function and the mean of the distribution (Zuur et al. 2009). In our case, the following GLM was applied:

542

543

1. Y_i , the lichen cover at tree i, was negative binomial distributed (NB) with mean μ_i and a dispersion parameter *k*.

544 2. The predictor function (η_1) included the following covariates: tree species, age, DBH, crown 545 volume, curvature, connectivity, and lichen cover. We also tested the interaction between 546 age and DBH.

547 3. There was a logarithm link between the mean of Y_i and the predictor function

548 The mathematical formulation was:

$$549 \qquad Lichen\ cover\ _i \sim NB(\mu_i,k)$$

 $E(Lichen \ cover_i) = \mu_i \quad \text{and} \quad \text{var}(Lichen \ cover_i) = \mu_i + \frac{\mu_i^2}{k} = \mu_i + \alpha \times \mu_i^2$

$$\frac{1}{551} \log(\mu_i) = \eta_i$$

 $\eta_i = \alpha + fTree \ species_i + Age_i \times DBH_i + Crown \ volume_i + Curvature_i + Connectivity_i$ $+ Lichen \ cover_i$

Hurdle model includes two components: (1) a count model for the positive values and (2) a binomial probability model for the distribution of zero values. The count component was modeled using a truncated negative binomial (ZANB) model with a logarithmic link function $log(\mu_i)$ to account for the overdispersion of the data. The binomial component was instead modeled using a binomial error distribution and a logit link function $logit(\mu_i)$. Applying the hurdle model we assumed that a species absence or zero abundance was due to changes in host trees and dispersal dynamics. Thus, the predictor function of binomial component (η_{bi}) included tree species and 560 connectivity as covariates, while the predictor function of count component (η_{ci}) included all

561 covariates (as for GLM).

562 The mathematical formulation was:

563 Lichen cover $_i \sim ZANB(\mu_i, \pi_i, k)$

$$E(Lichen \ cover_i) = \frac{1 - \pi_i}{1 - P_0} \times \mu_i \quad where \quad P_0 = \left(\frac{k}{\mu_i + k}\right)^k$$

$$\operatorname{var}(Lichen\ cover_i) = \frac{1 - \pi_i}{1 - P_0} \times \left(\mu_i^2 + \mu_i + \frac{\mu_i^2}{k}\right) - \left(\frac{1 - \pi_i}{1 - P_0} \times \mu_i\right)^2$$

565

568

564

$$\log\left(\mu_{i}\right) = \eta_{c_{1}}$$

567
$$logit(\mu_i) = \eta_{b_i}$$

$$\begin{split} \eta_{e_i} &= a_i + fTree\ species_i + Age_i \times DBH_i + Crown\ volume_i + Curvature_i + Connectivity_i \\ &+ Lichen\ cover_i \end{split}$$

569
$$\eta_{b_1} = \alpha_2 + fTree\ species_i + Connectivity_i$$

570 where π_i is the probability that $Y_i = 0$ (*Lichen cover*_i = 0).

Appendix S3. Moran's I correlograms for the 14 studied species, using a lag distance of 10 m. For *Chaenotheca trichialis* the graph based on a lag distance 5m is also given. Global significance, after applying Bonferonni correction, is reported. Species are grouped according to spatial distribution in (a) clumped and (b)

random.



(b) Species with random distribution



Appendix S4. Model averaged coefficients, standard errors (SE), confidence intervals (CI) and relative importance (cumulative Akaike weight) for variables predicting species with clumped distribution: (a) *Calicium viride*, (b) *Chaenotheca chrysocephala*, (c) *Chaenotheca trichialis*, (d) *Hypogymnia physodes*, (e) *Parmeliopsis ambigua*, (f) *Pseudevernia furfuracea*, and (g) *Vulpicida pinastri*.

(a) Calicium viride

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance
(a) Count componer	ıt				
(Intercept)	6.214	0.190	5.841	6.586	
Pine ³	-0.409	0.240	-0.878	0.061	0.89
Spruce	0.143	0.221	-0.291	0.576	"
Age	0.267	0.210	-0.145	0.679	0.54
DBH	0.136	0.166	-0.190	0.461	0.41
$Age \times DBH$	-0.571	0.352	-1.261	0.119	0.14
Canopy	0.123	0.186	-0.242	0.487	0.31
Curvature	-0.175	0.141	-0.451	0.102	0.44
Connectivity	-0.275	0.244	-0.753	0.202	0.38
Lichen cover	0.147	0.179	-0.203	0.498	0.32
(a) Binomial compo	nent				
(Intercept)	-0.079	0.200	-0.471	0.314	
Pine ³	-0.762	0.248	-1.247	-0.277	1.00
Spruce	0.303	0.233	-0.153	0.758	"
Connectivity	-0.194	0.178	-0.542	0.154	0.41

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance
(a) Count compone	ent				
(Intercept)	5.985	0.161	5.670	6.300	
Pine ³	0.432	0.198	0.044	0.819	1.00
Spruce	-0.582	0.195	-0.964	-0.199	
Age	-0.298	0.145	-0.581	-0.015	0.83
DBH	-0.505	0.192	-0.881	-0.129	0.93
$Age \times DBH$	-0.312	0.202	-0.707	0.083	0.40
Canopy	0.609	0.200	0.216	1.001	0.95
Curvature	-0.264	0.108	-0.476	-0.052	0.85
Connectivity	1.005	0.221	0.572	1.439	1.00
Lichen cover	-0.146	0.170	-0.478	0.187	0.35
(a) Binomial comp	onent				
(Intercept)	1.381	0.250	0.890	1.871	
Pine ³	-0.562	0.289	-1.129	0.005	0.95
Spruce	-0.832	0.278	-1.377	-0.286	
Connectivity	0.242	0.240	-0.229	0.712	0.41

(b) Chaenotheca chrysocephala

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)
 ³ Larch was the reference category.

(c) Chaenotheca trichialis

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative
(a) Count component					Importance
(Intercept)	5.590	0.334	4.935	6.244	
Pine ³	0.335	0.387	-0.424	1.094	1.00
Spruce	1.726	0.349	1.041	2.410	"
Age	0.008	0.224	-0.431	0.446	0.35
DBH	1.060	0.184	0.700	1.420	1.00
$Age \times DBH$	0.277	0.319	-0.349	0.903	0.12
Canopy	0.114	0.245	-0.365	0.594	0.28
Curvature	-0.496	0.157	-0.803	-0.188	0.98
Connectivity	0.151	0.132	-0.107	0.409	0.44
Lichen cover	-0.136	0.193	-0.515	0.243	0.31
(a) Binomial component	-1.744	0.281	-2.295	-1.193	
(Intercept)	0.368	0.268	-0.157	0.894	
Pine ³					1.00
Spruce	0.196	0.333	-0.456	0.849	"
Connectivity	2.768	0.311	2.159	3.378	0.54

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008). ² Unconditional standard errors (SE)

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance
(Intercept)	6.931	0.141	6.655	7.207	
Pine ³	-0.284	0.139	-0.556	-0.011	0.58
Spruce	-0.247	0.139	-0.520	0.026	"
Age	-0.246	0.098	-0.439	-0.054	0.91
DBH	0.467	0.147	0.179	0.755	0.98
Age \times DBH	-0.167	0.143	-0.448	0.114	0.36
Canopy	-0.246	0.123	-0.488	-0.004	0.72
Curvature	0.349	0.088	0.177	0.522	1.00
Connectivity	0.073	0.087	-0.097	0.243	0.31
Lichen cover	1.360	0.100	1.163	1.556	1.00

(d) Hypogymnia physodes

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)

³ Larch was the reference category.

(e) Parmeliopsis ambigua

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance
(Intercept)	8.058	0.109	7.844	8.273	
Pine ³	-0.304	0.131	-0.562	-0.046	1.00
Spruce	-0.787	0.126	-1.034	-0.540	"
Age	-0.115	0.094	-0.301	0.071	0.83
DBH	0.651	0.119	0.417	0.886	1.00
Age \times DBH	-0.441	0.140	-0.716	-0.167	0.76
Canopy	-0.113	0.123	-0.355	0.128	0.35
Curvature	0.132	0.084	-0.034	0.297	0.51
Connectivity	0.217	0.084	0.053	0.381	0.94
Lichen cover	0.473	0.091	0.294	0.652	1.00

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)

³ Larch was the reference category.

(f) Pseudevernia furfuracea

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance
(Intercept)	6.837	0.153	6.538	7.137	Importance
Pine ³	-0.089	0.184	-0.451	0.273	1.00
Spruce	-0.642	0.177	-0.989	-0.295	"
Age	0.217	0.133	-0.046	0.479	0.96
DBH	0.417	0.162	0.098	0.735	0.98
$Age \times DBH$	-0.467	0.195	-0.850	-0.084	0.78
Canopy	-0.145	0.171	-0.480	0.191	0.34
Curvature	0.336	0.118	0.105	0.567	0.95
Connectivity	0.032	0.116	-0.196	0.261	0.27
Lichen cover	1.321	0.129	1.069	1.573	1.00

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance
(Intercept)	-0.459	0.254	-0.958	0.040	
Pine ³	-0.077	0.310	-0.686	0.533	1.00
Spruce	1.577	0.288	1.012	2.142	"
Age	-0.762	0.218	-1.190	-0.335	1.00
DBH	1.049	0.260	0.538	1.560	1.00
Age \times DBH	-0.858	0.338	-1.521	-0.195	0.81
Canopy	0.226	0.257	-0.279	0.730	0.36
Curvature	0.014	0.184	-0.348	0.376	0.26
Connectivity	0.754	0.168	0.426	1.083	0.95
Lichen cover	0.476	0.197	0.089	0.863	0.85

(g) Vulpicida pinastri

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008). ² Unconditional standard errors (SE) ³ Larch was the reference category.

Appendix S5. Model averaged coefficients, standard errors (SE), confidence intervals (CI) and relative importance (cumulative Akaike weight) for variables predicting species with random distribution: (a) *Evernia divaricata*, (b) *Letharia vulpina*, (c) *Platismatia glauca*, (d) *Ramalina obtusata*, (e) *Schismatomma pericleum*, (f) *Tuckermannopsis chlorophylla*, and (g) *Tuckneraria laureri*.

(a) Evernia divaricata	ı
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Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance			
(a) Count component								
(Intercept)	1.871	0.405	1.076	2.665				
Pine ³	-0.383	0.386	-1.140	0.374	0.73			
Spruce	-0.820	0.347	-1.501	-0.140	"			
Age	0.781	0.299	0.195	1.367	0.91			
DBH	-0.325	0.287	-0.888	0.238	0.45			
Age \times DBH	0.276	0.421	-0.549	1.102	0.13			
Canopy	-0.237	0.295	-0.815	0.341	0.32			
Curvature	-0.231	0.335	-0.889	0.427	0.29			
Connectivity	-0.028	0.242	-0.502	0.446	0.27			
Lichen cover	0.245	0.301	-0.345	0.835	0.37			
(a) Binomial component								
(Intercept)	-0.877	0.146	-1.163	-0.591				
Pine ³	-0.389	0.261	-0.900	0.121	0.28			
Spruce	-0.239	0.248	-0.724	0.247	"			
Connectivity	0.096	0.168	-0.233	0.426	0.30			

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance			
(a) Count component								
(Intercept)	0.232	18.270	-35.576	36.040				
Pine ³	-2.155	1.667	-5.421	1.112	0.58			
Spruce	2.295	1.945	-1.518	6.108				
Age	0.261	3.596	-6.787	7.309	0.44			
DBH	1.422	1.650	-1.812	4.657	0.52			
Age \times DBH	4.752	4.193	-3.466	12.970	0.14			
Canopy	-0.302	1.852	-3.931	3.328	0.41			
Curvature	0.059	0.879	-1.664	1.781	0.28			
Connectivity	-7.508	7.336	-21.886	6.870	0.63			
Lichen cover	0.152	0.458	-0.745	1.050	0.17			
(a) Binomial component								
(Intercept)	-1.900	0.297	-2.483	-1.317				
Pine ³	-2.121	0.588	-3.273	-0.969	1.00			
Spruce	-3.082	0.770	-4.591	-1.573				
Connectivity	0.323	0.315	-0.295	0.941	0.34			

(b) Letharia vulpina

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)
 ³ Larch was the reference category.

(c) Platismatia glauca

(C) Futismatia glauca							
Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance		
(a) Count component							
(Intercept)	4.099	0.421	3.274	4.925			
Pine ³	-1.745	0.727	-3.170	-0.320	0.43		
Spruce	-0.092	0.625	-1.317	1.134	"		
Age	0.894	0.827	-0.727	2.516	0.42		
DBH	-0.448	0.610	-1.643	0.747	0.38		
$Age \times DBH$	-1.165	0.813	-2.759	0.430	0.07		
Canopy	0.055	0.546	-1.014	1.124	0.29		
Curvature	0.726	0.621	-0.490	1.943	0.42		
Connectivity	0.029	0.364	-0.684	0.742	0.27		
Lichen cover	-0.276	0.373	-1.007	0.455	0.31		
(a) Binomial component							
(Intercept)	-0.182	0.202	-0.578	0.214			
Pine ³	-2.996	0.396	-3.772	-2.221	1.00		
Spruce	-2.385	0.305	-2.981	-1.788	"		
Connectivity	0.291	0.212	-0.125	0.707	0.44		

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)
 ³ Larch was the reference category.

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance
(Intercept)	-3.044	0.431	-3.891	-2.197	
Pine ³	1.838	1.510	-1.128	4.804	1.00
Spruce	6.986	1.397	4.245	9.728	"
Age	1.235	0.408	0.435	2.036	0.99
DBH	2.173	0.337	1.512	2.834	1.00
Age \times DBH	-1.165	0.638	-2.417	0.087	0.58
Canopy	0.125	0.352	-0.567	0.817	0.29
Curvature	-0.128	0.270	-0.659	0.402	0.29
Connectivity	-0.235	0.523	-1.261	0.791	0.28
Lichen cover	-0.145	0.303	-0.740	0.451	0.29

(d) Ramalina obtusata

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)
 ³ Larch was the reference category.

(e) Schismaiomma pericieum							
Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance		
(a) Count component							
(Intercept)	4.873	0.914	3.082	6.664			
Pine ³	1.231	0.876	-0.487	2.949	0.84		
Spruce	2.120	0.810	0.532	3.708	"		
Age	0.343	0.454	-0.547	1.233	0.43		
DBH	-0.490	0.433	-1.339	0.358	0.52		
$Age \times DBH$	1.142	0.628	-0.090	2.374	0.19		
Canopy	0.584	0.376	-0.153	1.322	0.65		
Curvature	-0.633	0.313	-1.245	-0.020	0.72		
Connectivity	-0.027	0.388	-0.787	0.734	0.27		
Lichen cover	0.296	0.366	-0.420	1.013	0.33		
(a) Binomial component							
(Intercept)	-3.168	0.510	-4.168	-2.167			
Pine ³	0.909	0.560	-0.188	2.006	1.00		
Spruce	3.334	0.524	2.307	4.360	"		
Connectivity	0.025	0.187	-0.342	0.392	0.26		

(e) Schismatomma pericleum

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE)
 ³ Larch was the reference category.

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance			
(a) Count component								
(Intercept)	3.540	0.549	2.465	4.615				
Pine ³	-4.436	0.654	-5.717	-3.154	1.00			
Spruce	-3.011	0.485	-3.961	-2.061	"			
Age	0.363	0.439	-0.498	1.224	0.38			
DBH	-0.113	0.448	-0.992	0.766	0.34			
Canopy	0.460	0.315	-0.158	1.077	0.55			
Curvature	0.082	0.328	-0.561	0.725	0.28			
Connectivity	0.053	0.219	-0.376	0.483	0.28			
Lichen cover	-0.765	0.301	-1.355	-0.175	0.81			
(a) Binomial component								
(Intercept)	-0.704	0.218	-1.130	-0.277				
Pine ³	-2.039	0.358	-2.741	-1.337	1.00			
Spruce	-0.701	0.264	-1.219	-0.183	"			
Connectivity	0.923	0.214	0.503	1.343	1.00			

(f) Tuckermannopsis chlorophylla

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008).

² Unconditional standard errors (SE) ³ Larch was the reference category.

(g) Tuckneraria laureri

Predictor	Estimate ¹	SE ²	Lower CI	Upper CI	Relative importance			
(a) Count component								
(Intercept)	0.564	0.452	0.279	0.865				
Pine ³	-1.579	0.558	-2.672	-0.486	0.88			
Spruce	0.140	0.577	-0.991	1.271	"			
Age	-0.287	0.652	-1.564	0.991	0.37			
DBH	0.942	0.552	-0.139	2.023	0.69			
$Age \times DBH$	1.066	0.961	-0.816	2.949	0.11			
Canopy	-0.346	0.442	-1.212	0.520	0.33			
Curvature	-0.938	0.696	-2.301	0.426	0.44			
Connectivity	-0.353	0.263	-0.867	0.162	0.38			
Lichen cover	-0.012	0.397	-0.790	0.766	0.27			
(a) Binomial component								
(Intercept)	-0.274	0.203	-0.671	0.123				
Pine ³	-1.708	0.289	-2.274	-1.141	1.00			
Spruce	-1.514	0.263	-2.030	-0.998	"			
Connectivity	0.598	0.185	0.234	0.961	0.99			

¹ Effect sizes have been standardized on two standard deviation (SD) following Gelman (2008). ² Unconditional standard errors (SE) ³ Larch was the reference category.



Appendix S6. The effect of tree age on the response of (a) Parmeliopsis ambigua and (b) Vulpicida pinastri cover to tree size (DBH). DBH have