Morphological and Chemical Traits of Cladonia Respond to Multiple Environmental Factors in Acidic Dry Grasslands

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Abstract: Terricolous lichen communities in lowlands occur especially in open dry habitats. Such communities are often dominated by species of the genus Cladonia, which are very variable in morphology, reproduction strategies, and secondary metabolites. In this work, we investigated traits-environment relationships considering vegetation dynamics, substrate pH, disturbance, and climate. A total of 122 plots were surveyed in 41 acidic dry grasslands in the western Po Plain (Northern Italy). Relationships between Cladonia traits and environmental variables were investigated by means of a model-based Fourth Corner Analysis. Thallus morphology and metabolites responded to vegetation dynamics, substrate pH, disturbance, and climate, whereas reproduction strategies responded only to vegetation dynamics. Traits’ correlations with vegetation dynamics elucidate their colonization patterns in open dry habitats or suggest biotic interactions with bryophytes and vascular plants. In addition, correlations between metabolites and environmental factors support interpretations of their ecological roles. Our results also stress the importance of studying traits’ relationships with climatic factors as an alert towards lichen reactions to climate change.

Keywords: lichens; open dry habitats; reproduction strategy; secondary metabolites; species traits; thallus growth forms; vegetation dynamics

1. Introduction

The analysis of functional traits to explore species’ responses to environmental factors is increasingly applied also to lichens [1,2]. However, most studies have addressed epiphytes (e.g., [3–5]), whereas terricolous lichens are relatively less investigated (e.g., [6–8]) despite their ecological importance (e.g., [9]) and conservation concern [6,10]. It is therefore of utmost importance to understand the relationships driving terricolous species’ responses to environmental stresses, particularly in the current context of global change [2,5,6].

Terricolous lichen communities are often dominated by species of the genus Cladonia. In this genus, the thallus is composed of two parts: a basal primary thallus and a secondary thallus with a very variable morphology across species [11–13]. This high variability leads to a range of possible morphological combinations that have been almost overlooked in previous literature (e.g., [6,7,14,15]). Additionally, reproduction,
which is a relevant trait in species life history [5,6], is achieved in variable ways, not only sexually (by means of ascospores), but also with different types of vegetative propagules [11,12], as well as by dispersion of thallus fragments [16]. The position of pycnidia may also vary, developing either on the primary or secondary thallus [12,13]. 

Cladonia lichens produce several secondary metabolites, some of them being widely studied due to their potential cytotoxic activity [17–19]. However, their relationships with environmental factors have been little explored [20–22].

In lowland landscapes of central-southern Europe, Cladonia-dominated communities occur especially in open dry habitats on acidic substrates [10,14,23–26]. Open dry habitats are among the most threatened by human activities – land-use change, land consumption, pollution – and climate change [27], but also by abandonment in less productive regions [28], especially in densely inhabited lowland areas, such as the Po Plain [29,30]. Among them, there is a relevant role for terricolous lichen diversity in acidic dry grasslands [6,7,10,24–26,31], which develop on acidic, mineral, shallow, and oligotrophic soils [32]. In this habitat, terricolous lichen communities are influenced by small-scale disturbance, e.g., by trampling or invasive species, and by climate features [8,10].

This study aims at exploring the relationships between Cladonia functional traits and environmental factors that drive community composition in lowland acidic dry grasslands. We hypothesized that (i) the abundance of different functional traits, such as growth forms, reproductive strategies, and secondary metabolites, can be shaped by the main environmental factors, i.e., vegetation dynamics, disturbance, substrate pH, and climatic features (temperature and precipitation); and that (ii) discriminating traits like growth forms and reproduction strategy in detail could give better insights on the responses of these lichens to environmental factors.

2. Materials and Methods

2.1. Study Area

The study was carried out in the central-western Po Plain (northern Italy), in an area located on the two sides of the boundary between the regions of Lombardy and Piedmont (Figure 1). The mean annual temperature ranges between 13.3 and 14.2 °C. Annual rainfall ranges between 788 and 1104 mm. The altitude varies between 61 and 189 m a.s.l.

In this area, 41 sites with lichen-rich acidic dry grasslands were searched for and located (Figure 1). They were clustered along the course of two main rivers, i.e., the Sesia (7 sites) and the Ticino (26 sites), in a stretch of the Po river where the substrate is acidic (3 sites) and in a small area in Lomellina in which residual inland sand dunes still occur (5 sites). These grasslands are attributed to the Natura 2000 Habitat 2330 (“Inland dunes with open Corynephorus and Agrostis grasslands”) or to a pioneer and acidic facies of Habitat 6210 (“Seminatural dry grasslands and scrubland facies on calcareous substrates”). Acidic dry grasslands are severely threatened in Europe [27,32], and therefore these grasslands have a relevant conservation value. Habitat 2330 has also a biogeographical value in this area since it is at the southernmost edge of its distribution range [30].

In the study area, these grasslands are often fragmented and located in marginal and unproductive areas that are not actively managed and are sometimes degraded due to human disturbance – typically uncontrolled grazing and motorbike riders – with the colonization of invasive species [29,33]. Nevertheless, they frequently host terricolous lichen communities, and though species-poorer than in similar habitats in central Europe, they include some species with a more Mediterranean distribution pattern [10,25,26,34,35].
2.2. Sampling

At each site, a linear transect connecting the two furthest vertices of the grassland was laid out. Along each transect, from 1 to 7 circular plots with a 3 m radius were placed at regular intervals proportionally to the site area. This resulted in a total of 122 plots.

Vegetation was recorded in each plot between April and June 2016. We recorded the cover (%) and the mean height (cm) of the five vegetation layers – cryptogamic, herbaceous, lower-shrubby (shrubs up to 1.5 m), higher-shrubby (shrubs between 1.5 and 3 m), arboreal (woody species over 3 m high) – and the cover (%) of each vascular plant, lichen and bryophyte species. Easily recognizable species were identified in the field, whereas difficult specimens were collected and identified in the laboratory. All the specimens are retained in the first author’s personal herbarium.

In each plot, we also recorded the occurrence of disturbance that could impact lichens. Human trampling was estimated according to a categorical scale: 0 (no trampling), 1 (<5 m$^2$ showing evidence of trampling on vegetation), 2 (5–10 m$^2$), 3 (10–15 m$^2$), 4 (>15 m$^2$). The impact by lagomorphs was estimated using the abundance of fecal pellets as a proxy [33], according to a categorical scale: 0 (no pellet), 1 (<2 pellet/m$^2$), 2 (3–5 pellet/m$^2$), 3 (>5 pellet/m$^2$). Additionally, soil pH was recorded on the field by means of a portable kit.

2.3. The Genus Cladonia

*Cladonia* (Hill.) P. Browne (Cladoniaceae, Lecanorales, Ascomycetes) is a cosmopolitan and megadiverse genus with a wide altitudinal and ecological range, most species being terricolous and ranging from mineral to humus-rich soils [11,12]. *Cladonia* species are characterized by a thallus composed of two parts: a squamulose or crustose primary thallus and a fruticose secondary thallus with a very variable morphology that can include stick-shaped, club-shaped, cup-shaped, sparingly branched or richly branched structures called podetia [11–13]. The primary thallus can be ephemeral or persistent,
and, in this case, even dominant; on the other hand, the secondary thallus, which is often dominant, in some species can be inconspicuous or even absent [11–13].

Reproduction can be sexual or asexual. Cladonia has biterine apothecia which produce simple ascospores; apothecia develop at the tips of capitiform, bacillary and branched podetia, on the edges of scyphipherous podetia or, in few species, directly on primary squamules [11–13]. Vegetative propagules can include soredia, schizidia, blastidia, and microsquamules [11,12], but also thallus fragments can act efficiently as propagules [16,36]. Vegetative reproduction involving only the mycobiont is carried out by conidia; conidia are produced in pycnidia, which in Cladonia can be located either on the primary squamules or on podetia [11–13].

Cladonia lichens produce several compounds, chiefly aliphatic acids (e.g., rangifor-mic and bourgeanic acids), dibenzofurans (e.g., usnic acid), depsides (e.g., homosekikaic and perlatic acid, atranorin), and depsidones (e.g., fumarprotocetraric, norstictic, and psoromic acids) [37–43]. Some of them show cytotoxic activity and are much studied for their pharmacological potential [17–19], but their ecological roles have been addressed more rarely. More investigated are allelopathic [44–46] and anti-herbivorous [47–49] effects, rarer are ecological studies that investigated their roles in photoprotection [50–52] and in regulating species’ preferences for substrate pH [20–22,53].

2.4. Functional Traits

Three groups of functional traits were considered: growth form, reproductive strategy, and secondary metabolites.

Previous literature that considered the growth forms of Cladonia in the analysis of functional traits used a weak differentiation just between foliose/squamulose (for species without secondary thallus) and fruticose thalli, or between foliose/squamulose, fruticose with simple podetia, and fruticose with branched podetia [6,7,14,15]. The huge diversity occurring within Cladonia, not only in morphology but also in size, deserves a sharper and more precise distinction since different shapes and sizes can potentially give different benefits or disadvantages under different environmental conditions, at a microhabitat scale. On the basis of morphological data reported in the main literature sources [11–13,54] and many personal observations on the specimens collected for this work, we considered six different growth forms (Table 1).

| Abbreviation       | Description                                                                 |
|--------------------|-----------------------------------------------------------------------------|
| Small squamules    | Squamulose thallus with small squamules: species usually without podetia and with squamules usually shorter than 5 mm; e.g., Cladonia strepsilis. |
| Big squamules      | Squamulose thallus with big squamules: species usually without podetia and with squamules usually longer than 5 mm; e.g., Cladonia foliacea. |
| Small simple podetia| Thallus with small simple podetia: species with persistent primary thallus and usually with bacillar or capitiform podetia shorter than 10 mm; e.g., Cladonia peziziformis. |
| Big simple podetia | Thallus with big simple podetia: species with persistent or ephemeral primary thallus and usually with bacillar of scyphipherous podetia taller than 10 mm; e.g., Cladonia pyxidata, Cladonia rei. |
| Branched podetia   | Thallus with branched podetia: species with ephemeral primary thallus and with sparingly branched podetia; e.g., Cladonia furcata. |
| Richly branched podetia | Thallus with richly branched podetia: species with ephemeral |
primary thallus and with richly branched, bush-shaped podetia; e.g., *Cladonia rangiformis* and species belonging to subgenus *Cladina*.

The reproduction strategy was described by means of the main reproduction type and the position of pycnidia on thallus. The main reproduction type of each species was retrieved from the database ITALIC [55]. Two types were considered: sexual reproduction by ascospores or asexual reproduction by soredia. The position of pycnidia was assessed consulting the main literature sources [11–13,54] and also through personal observations on the specimens collected for this work. Two cases were considered: pycnidia on the primary squamules or pycnidia on the podetia. Species with pycnidia on the primary thallus could be expected to have a faster development, being earlier colonizers of pioneer situations, whereas species with pycnidia on podetia could be expected to have slower development and to occur later in the succession since podetia develop after primary thallus.

The occurrence of secondary metabolites was assessed by means of thin-layer chromatography (TLC) performed with the solvents A, B’, and C [56]. The eight most frequent metabolites were atranorin, fumarprotocetraric acid, homosekikaic acid, norstictic acid, perlatic acid, rangiformic acid, usnic acid, and zeorin. In addition, baeomycesic acid, squamatic acid, and strepsilin occurred only in one rarely recorded species, *Cladonia strepsilis*, and therefore they were not considered in the analysis.

The attribution of the considered functional traits to the 14 *Cladonia* species recorded in the 122 plots is shown in Table 2.

For each trait, the abundance in each plot was calculated as the sum of the abundances of the species with that trait in the plot. The abundances, originally recorded in the field in percent values, were converted in a scale ranging from 1 to 10, as follows: 1–10% = 1; 11–20% = 2; 21–30% = 3; 31–40% = 4; 41–50% = 5; 51–60% = 6; 61–70% = 7; 71–80% = 8; 81–90% = 9; 91–100% = 10.

**Table 2.** Functional traits of the 14 *Cladonia* species recorded in the 122 plots. Nomenclature follows Nimis and Martellos (2020).

| Species                      | Growth Form         | Position of Pycnidia | Reproduction | Metabolites                                  |
|------------------------------|---------------------|----------------------|--------------|----------------------------------------------|
| *Cladonia cariosa*           | Small simple podetia| Squamules            | Spores       | Atranorin, rangiformic acid                  |
| (Ach.) Spreng.               |                     |                      |              |                                              |
| *Cladonia chlorophaea*       | Big simple podetia  | Podetia              | Soredia      | Fumarprotocetraric acid                     |
| (Sommerf.) Spreng.           |                     |                      |              |                                              |
| *Cladonia coccifera*         | Big simple podetia  | Podetia              | Spores       | Usnic acid, zeorin                          |
| (L.) Willd.                  |                     |                      |              |                                              |
| *Cladonia fimbriata*         | Big simple podetia  | Podetia              | Soredia      | Fumarprotocetraric acid                     |
| (L.) Fr.                     |                     |                      |              |                                              |
| *Cladonia foliacea*          | Big squamules       | Squamules            | Spores       | Fumarprotocetraric acid, usnic acid         |
| (Huds.) Willd.               |                     |                      |              | Atranorin, fumarprotocetraric acid          |
| *Cladonia furcata*           | Branched podetia    | Podetia              | Spores       | Atranorin, fumarprotocetraric acid          |
| (Huds.) Schrad.              |                     |                      |              |                                              |
| *Cladonia peziziformis*      | Small simple podetia| Squamules            | Spores       | Fumarprotocetraric acid                     |
| (With.) J.R.Laundon Nyl.     |                     |                      |              |                                              |
| *Cladonia polyarceoides*     | Big squamules       | Squamules            | Spores       | Norstictic acid                              |
| (Dufour) Coem.               |                     |                      |              |                                              |
| *Cladonia portentosa*        | Richly branched podetia | Podetia              | Spores       | Perlatolic acid, usnic acid                 |
| (Dufour) Coem.               |                     |                      |              |                                              |
| *Cladonia pyxidata*          | Big simple podetia  | Podetia              | Spores       | Fumarprotocetraric acid                     |
| (L.) Hoffm.                  |                     |                      |              |                                              |
| *Cladonia rangiformis*       | Richly branched podetia | Podetia              | Spores       | Atranorin, rangiformic acid                 |
| Hoffm.                       |                     |                      |              |                                              |
| *Cladonia rei*               | Big simple podetia  | Podetia              | Soredia      | Fumarprotocetraric acid, homosekikaic acid  |
| Schaer.                      |                     |                      |              |                                              |
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2.5. Environmental Variables

The cover values (%) of the vascular plant biological forms [57] were calculated for each plot based on the floristic composition recorded. Therophytes (annual/biennial herbs), hemi-cryptophytes, and geophytes (perennial herbs) are part of the herbaceous layer, chamephytes are generally found in the lower-shrubby layer and phanerophytes, typical of higher-shrubby and arboreal layers, can be found in the lower-shrubby layer when young. Biological forms were considered in the analysis as a proxy of vegetation dynamics since it is known that therophytes dominate pioneer stages, also indicating ongoing vegetation dynamics due to disturbance in some cases; hemi-cryptophytes and geophytes dominate intermediate stages, indicating less active but still ongoing vegetation dynamics; chamephytes and phanerophytes dominate more mature stages, i.e., scrub and forest, indicating the passage from grassland to more developed vegetation types [58].

Climatic variables, i.e., mean annual temperature and annual precipitation (considered as a proxy of humidity), were retrieved for each sampling plot from CHELSA [59].

2.6. Data Analysis

To explore the relationships between functional traits and environmental variables, a model-based fourth corner analysis was used. This method is aimed at solving the “fourth corner problem”, by analyzing the relationships between the three matrices (i) species x sites, (ii) species x traits, and (iii) sites x environmental variables, to estimate a matrix with environment-trait associations [60]. In particular, we followed the framework proposed by [61] and implemented it in the R package “mvabund” [62]. This approach proceeds by fitting a GLM with species abundances as a function of species’ traits, environmental variables, and their interactions. The model was fitted using a Poisson distribution with LASSO penalty to enhance prediction accuracy; this latter sets to zero all the coefficient terms that do not explain any variation [61]. In the end, the model was evaluated through diagnostic plots.

3. Results

Significant relationships were found between morphological traits and vegetation dynamics, substrate features, disturbance, and climate (Figure 2). Small squamules, small simple podetia, branched, and richly branched podetia correlated with variables describing vegetation dynamics. Big squamules, small and big simple podetia, branched podetia correlated with climatic variables. Small simple podetia, branched and richly branched podetia correlated with substrate pH, and richly branched podetia correlated with trampling.

Reproduction traits were less responsive to environmental factors (Figure 2). Only sexual reproduction and pycnidia located on primary squamules correlated with few variables describing vegetation dynamics.

Significant relationships were found also between secondary metabolites and various predictors associated with vegetation dynamics, substrate, disturbance, and climate (Figure 2). Atranorin, homosekikaic, norstictic, perlatoilic, rangiformic, and usnic acids correlated with variables describing vegetation dynamics. Atranorin, fumarprotocetraric acid, and zeorin correlated with climatic variables. Fumarprotocetraric acid and zeorin correlated with substrate pH, while fumarprotocetraric acid correlated with fecal pellets and rangiformic acid with trampling.
4. Discussion

In accordance with our hypothesis, morphological, reproduction, and chemical traits of terricolous Cladonia species in acid dry grasslands were involved in the responses of these organisms mainly to vegetation dynamics and climate, but also, to a lesser extent, to disturbance and substrate pH.

Some relationships with vegetation dynamics were particularly evident, e.g., species with small simple podetia and species with pycnidia on primary squamules were more frequent in stages dominated by therophytes (pioneer grasslands), decreasing in stages dominated by other biological forms (intermediate-mature grasslands). An opposite pattern was found for species with branched and richly branched podetia and species with perlatolic acid, related to intermediate-mature stages. An allelopathic activity against vascular plants was demonstrated for perlatolic acid [45], but if this was the case, an evident correlation should have been observed for other biological forms considered, not only with therophytes; this negative correlation could better be seen as a link with mature stages of dry grasslands.

Research has suggested photoprotection activity for usnic acid [52] and facilitation in exploiting low light intensities for atranorin [63]. This can explain their correlations
with vascular plants which produce a thicker canopy than herbs and forbs, i.e., chame-
phytes and phanerophytes, which occur in mature stages of dry grasslands. These two
metabolites showed opposite patterns in relation to the canopy, i.e., usnic acid correlated
negatively and atranorin correlated positively. Considering the 83 Cladonia taxa reported
in Italy so far [55], none of them contain both these metabolites at the same time [11,12].
This could suggest that the production of one compound instead of another can help
these lichens to cope with site-specific light conditions—and, consequently, with a dif-
ferent stage of vegetation succession.

Species with homosekikaic and rangiformic acid correlated positively with the cover
of therophytes, and the species which produce at least one of these two metabolites – C.
cariosa, C. rangiformis, C. rei, which have different growth forms and reproductive strate-
gies – are widely recognized as early colonizers of prohibitive substrates which often
dominate the communities where they develop [13,64–66]. However, they also occur, and
often dominate in intermediate-mature stages [25,26], as backed by the positive cor-
relations with other biological forms of vascular plants. The role of these two compounds
in fostering colonization of primitive substrates and long-lasting dominance could be
hypothesized.

At a smaller scale, the dynamics of cryptogam communities in dry grasslands are
linked also to bryophytes, which are more abundant in intermediate-mature stages
[25,26]. However, correlations with bryophyte cover could also suggest facilitation/competition dynamics. The positive relationship with small squamules could be related to water provisioning [67]: for example, inconspicuous Cladonia with small squamules could be easier to fit within higher covers of mosses and profit from the water they retain. In contrast, negative correlations with richly branched podetia and reproduction by apothecia suggest competition between mosses and lichens. Richly branched Cladonia are typical of intermediate-mature stages of vegetation succession, in which bryophytes also reach high cover values. A high bryophyte cover can make it more difficult for spores to encounter a photobiont for the regeneration of a new lichen thallus.

Competition with bryophytes and vascular plants is not the only stressful factor for
lichens in dry grasslands, however, we found only a few correlations between traits and disturbance factors. Trampling is regarded as the main threat to Cladonia lichens [68], but the positive correlation with richly branched podetia suggests that a moderate trampling could be a positive factor in open habitats, e.g., as a major driver of dispersal [16,36,69]. Trampling has the positive effect of producing and dispersing thallus fragments and, therefore, it could be particularly beneficial for lichens with large and fragile thalli. Additionally, the abundance of fecal pellets by lagomorphs can have different effects on li-
chens [33]. In our case, their positive correlation with fumarprotocetraric acid could suggest that this metabolite allows lichens to deal with a nitrified substrate since it is al-
ready known that this compound helps in tolerating substrates containing heavy metals
[70] and has an antimicrobial activity [71].

Substrate pH is a limiting factor also. Small simple podetia, branched podetial, and
fumarprotocetraric acid correlated positively with pH, suggesting that these traits are
fostered by subneutral soils; contrariwise, richly branched podetia and zeorin occurred in
very acidic substrates. These growth forms include species ranging from acidic to cal-
careous substrates, e.g., C. cariosa, C. furcata, C. rangiformis, which is the case also for some
species with fumarprotocetraric acid; therefore, these correlations could be spurious and
due to the incomplete pH range included in our data, limited to acidic substrates. The
 correlation with zeorin is backed by [21], who suggested that this metabolite could play a
 role in the interaction with strongly acidic substrates.

Our results indicate that climatic factors may also contribute to the selection of spe-
cies traits in local communities, determining species dynamics and community compos-
tion in a climate change scenario [8,72,73]. An outstanding example is that of species with
atranorin, which may be fostered by increasing temperatures and decreasing precipita-
tion. The size of simple podetia increased with increasing precipitation, with small sim-
ple podetia correlating negatively and large simple podetia correlating positively, which could be related to a better water exploitation capacity by larger podetia [74]; however, small simple podetia can also be expected to decrease at increasing temperatures. Partial responses were found also for species with big squamules and fumarprotocetraric acid, expected to increase respectively at increasing temperatures and decreasing precipitations, and by species with big simple podetia and zeorin, expected to decrease with decreasing precipitation.

5. Conclusions

*Cladonia* shows a wide variety of thallus growth forms, reproductive strategies, and chemotypes [11,12], being, therefore, a suitable model genus to assess the relationships between environmental factors and species traits of terricolous communities. Therefore, our results can have broader applicability for a better understanding of these communities from a functional standpoint, also considering that *Cladonia*-dominated communities often have a similar composition across different biogeographical contexts (cf. [11,12,55]).

The relationships between species traits and vegetation dynamics can help elucidate their colonization patterns in open dry habitats, also reflecting the effect of biotic interactions between lichens and bryophytes or vascular plants. Furthermore, the correlations of some metabolites with certain environmental factors could help in addressing future research aimed at understanding their ecological roles, which are still largely unexplored.

From a conservation standpoint, the contrasting relationships between some traits and vegetation dynamics support the view that management of acidic dry grasslands should aim at maintaining patches at different dynamics stages (pioneer, intermediate, mature) to maximize taxonomic and functional diversity of lichen communities [10].

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**References**

1. Nimis, P.L.; Martellos, S.; Chiarucci, A.; Ongaro, S.; Peplis, M.; Pittao, E.; Nascimbene, J. Exploring the Relationships between Ecology and Species Traits in Cyanolithicens: A Case Study on Italy. *Fungal Ecol.* 2020, 47, 100950, doi:10.1016/j.funeco.2020.100950.
2. Saiz, H.; Dainese, M.; Chiarucci, A.; Nascimbene, J. Networks of Epiphytic Lichens and Host Trees along Elevation Gradients: Climate Change Implications in Mountain Ranges. *J. Ecol.* 2020, doi:10.1111/1365-2745.13538.
3. Giordani, P.; Brunialti, G.; Bacaro, G.; Nascimbene, J. Functional Traits of Epiphytic Lichens as Potential Indicators of Environmental Conditions in Forest Ecosystems. *Ecol. Indic.* 2012, 18, 413–420, doi:10.1016/j.ecolind.2011.12.006.
4. Jüriado, I.; Paal, J. Epiphytic Lichen Synusiae and Functional Trait Groups in Boreo-Nemoral Deciduous Forests Are Influenced by Host Tree and Environmental Factors. *Nord. J. Bot.* 2019, 37, e01939, doi:10.1111/njb.01939.
5. Hurtado, P.; Prieto, M.; Martínez-Vilalta, J.; Giordani, P.; Aragón, G.; López-Angulo, J.; Košuthová, A.; Merinero, S.; Díaz-Peña, E.M.; Rosas, T.; et al. Disentangling Functional Trait Variation and Covariation in Epiphytic Lichens along a Continent-Wide Latitudinal Gradient. *Proc. R. Soc. B* 2020, 287, 20192862, doi:10.1098/rspb.2019.2862.
6. Leppik, E.; Jüriado, I.; Suja, A.; Liira, J. Functional Ecology of Rare and Common Epigeic Lichens in Alvar Grasslands. *Fungal Ecol.* 2015, 13, 66–76, doi:10.1016/j.funecc.2014.08.003.

7. Jüriado, I.; Kämärä, M.-L.; Oja, E. Environmental Factors and Ground Disturbance Affecting the Composition of Species and Functional Traits of Ground Layer Lichens on Grey Dunes and Dune Heaths of Estonia. *Nord. J. Bot.* 2016, 34, 244–255, doi:10.1111/njbo.00936.

8. Balveese, C.; Nascimbene, J.; Giordani, P.; Benesperi, R.; Casazza, G. Modelling Range Dynamics of Terricolous Lichens of the Genus Peltigera in the Alps under a Climate Change Scenario. *Fungal Ecol.* 2021, 49, 101014, doi:10.1016/j.funecc.2020.101014.

9. Maestre, F.T.; Bowker, M.A.; Cantón, Y.; Castillo-Monroy, A.P.; Cortina, J.; Escolar, C.; Escudero, A.; Lázaro, R.; Martínez, I. Ecology and Functional Roles of Biological Soil Crusts in Semi-Arid Ecosystems of Spain. *J. Arid Environ.* 2011, 75, 1282–1291, doi:10.1016/j.jaridenv.2010.12.008.

10. Gheza, G.; Assini, S.; Lelli, C.; Marini, L.; Mayrhofer, H.; Nascimbene, J. Biodiversity and Conservation of Terricolous Lichens and Bryophytes in Continental Lowlands of Northern Italy: The Role of Different Dry Habitat Types. *Biodivers. Conserv.* 2020, 29, 3533–3550, doi:10.1007/s10531-020-02034-1.

11. Burgaz, R.A.; Ahti, T. *Cladoniaceae*; Flora liquenológica ibérica; Sociedad Española de Lichenología: Madrid, Spain, 2009; ISBN 1696-0513.

12. Ahti, T.; Stenroos, S.; Moberg, R. Nordic Lichen Flora; Museum of Evolution, Uppsala University, Uppsala, 2013; Volume 5.

13. Gheza, G. Chiave Alle Specie Italiane Del Genere Cladonia. *Not. Della Soc. Lichenol. Ital.* 2018, 31, 70–96.

14. Bültmann, H. Zeigerwerte von Erdflachen in Trockenrasen: Vorschläge zur Ergänzung und Korrektur. In *Trockenrasen auf Unterschiedlichen Betrachtungsseben*; Bültmann, H., Fartmann, T., Hasse, T., Eds.; Institut fur Landschaftsökologie Münster: Münster, Germany, 2006; Volume 15, pp. 127–143.

15. Sulavik, J.; Auestad, I.; Halvorsen, R.; Rydgren, K. Assessing Recovery of Alpine Spoil Heaps by Vascular Plant, Bryophyte, and Lichen Functional Traits. *Restor. Ecol.* 2020, doi:10.1111/rec.13257.

16. Heinken, T. Dispersal Patterns of Terricolous Lichens by Thallus Fragments. *Lichenologist* 1999, 31, 603–612, doi:10.1006/lich.1999.0219.

17. Molnár, K.; Farkas, E. Current Results on Biological Activities of Lichen Secondary Metabolites: A Review. *Z. Für Nat. C* 2010, 65, 157–173, doi:10.1515/znc-2010-3-401.

18. Stanojkovic, T. *Lichen Secondary Metabolites: Bioactive Properties and Pharmaceutical Potential*; Ranković, B., Ed.; Springer International Publishing: New York, NY, USA, 2015; ISBN 978-3-319-35516-0.

19. Goga, M.; Elekt, J.; Marcinčinová, M.; Ručová, D.; Bačkorová, M.; Bačkor, M. Lichen Metabolites: An Overview of Some Secondary Metabolites and Their Biological Potential. In *Co-Evolution of Secondary Metabolites*; Méryillon, J.-M., Ramawat, K.G., Eds.; Reference Series in Phytochemistry; Springer International Publishing: Cham, Switzerland, 2020; pp. 175–209, ISBN 978-3-319-96397-6.

20. Park, Y.S. Habitat Selection in a Pair of Sibling Chemospecies of the Lichen Genus Cladonia. *Am. Midl. Nat.* 1985, 114, 180–183, doi:10.2307/2425253.

21. Zraik, M.; Booth, T.; Piercey-Normore, M.D. Relationship between Lichen Species Composition, Secondary Metabolites and Soil PH, Organic Matter, and Grain Characteristics in Manitoba. *Botany* 2018, 96, 267–279, doi:10.1139/cjb-2017-0176.

22. Hauck, M.; Jürgens, S.-R.; Leuschner, C. Norstictic Acid: Correlations between Its Physico-Chemical Characteristics and Ecology. *Phytocoenologia* 2021, 66–76, doi:10.1016/j.funeco.2020.101014.

23. Ketner-Oostra, R.; Sykora, K.V. Vegetation Change in a Lichen-Rich Inland Drift Sand Area in the Netherlands. *Phytocoenologia* 2009, 267–286, doi:10.1127/0340-269X/2009/0038-0267.

24. Gheza, G.; Assini, S.; Valcuvia Passadore, M. Terricolous Lichen Communities of Corynephorus Canescens Grasslands of Northern Italy. *Tuscania* 2016, 36, 121–142, doi:10.14471/2016.36.005.

25. Gheza, G.; Barcella, M.; Assini, S. Terricolous Lichen Communities in Thero-Airion Dry Grasslands of the Po Plain (Northern Italy): Syntaxonomy, Ecology and Conservation Value. *Tuscania* 2019, 39, 377–400, doi:10.14471/2019.39.011.

26. Janssen, J.A.M.; Rodwell, J.S.; Criado, M.G.; Arts, G.H.P.; Bijlsma, R.J.; Schaminee, J.H.J.; European Red List of Habitats: Part 2. *Terrestrial and Freshwater Habitats*; European Union: Luxembourg, 2016.

27. Jentsch, A.; Beyschlag, W. Vegetation Ecology of Dry Acidic Grasslands in the Lowland Area of Central Europe. *Flora* 2003, 198, 3–25, doi:10.1078/0367-2530-00071.
33. Gheza, G.; Assini, S.; Marini, L.; Nascimbene, J. Impact of an Invasive Herbivore and Human Trampling on Lichen-Rich Dry Grasslands: Soil-Dependent Response of Multiple Taxa. *Sci. Total Environ.* 2018, 639, 633–639, doi:10.1016/j.scitotenv.2018.05.191.

34. Gheza, G. Terricolous Lichens of the Western Pianian Plain: New Records of Phytogeographical Interest. *Acta Bot. Gall.* 2015, 162, 339–348, doi:10.1080/12538078.2015.1108667.

35. Gheza, G. Licheni Terricoli Degli Ambienti Aperti Aridi Della Pianura Piemontese. *Riv. Piemont. Stor. Nat.* 2020, 41, 23–31.

36. Lange, O.L.; Green, T.G.A.; Türk, R. An Unusual Growth Form of *Cladonia Furcata*: The Trampling-Resistant Primary Thallus Colonizing a Paved Pathway. *Lichenologist* 1998, 30, 583–588, doi:10.1006/lich.1998.0169.

37. Huovinen, K.; Ahti, T. Biosequential Patterns for the Formation of Depsides, Depsidones and Dibenzofurans in the Genus *Cladonia* (Lichen-Forming Ascomycetes). *Ann. Bot. Fenn.* 1982, 19, 225–234.

38. Huovinen, K.; Ahti, T. The Composition and Contents of Aromatic Lichen Substances in the Genus *Cladina*. *Ann. Bot. Fenn.* 1986, 23, 93–106.

39. Huovinen, K.; Ahti, T. The Composition and Contents of Aromatic Lichen Substances in *Cladonia*, Section Unciales. *Ann. Bot. Fenn.* 1986, 23, 173–188.

40. Huovinen, K.; Ahti, T. The Composition and Contents of Aromatic Lichen Substances in *Cladonia* Section *Perviae*. *Ann. Bot. Fenn.* 1988, 25, 371–383.

41. Huovinen, K.; Ahti, T.; Stenroos, S. The Composition and Contents of Aromatic Lichen Substances in *Cladonia*, Section *Cocciferae*. *Ann. Bot. Fenn.* 1989, 26, 133–148.

42. Huovinen, K.; Ahti, T.; Stenroos, S. The Composition and Contents of Aromatic Lichen Substances in *Cladonia* Section *Helodium* and Subsection *Folisae*. *Ann. Bot. Fenn.* 1989, 26, 297–306.

43. Huovinen, K.; Ahti, T.; Stenroos, S. The Composition and Contents of Aromatic Lichen Substances in *Cladonia* Section *Cladagia* and Group *Furcatae*. *Bibl. Lichenol.* 1990, 209–241. Available online: https://www.cabdirect.org/otrag?%2fcabdirect%2fabstrac%2f20198610189 (accessed on 26 January 2021).

44. Lokajová, V.; Bačkórová, M.; Bačko, M. Allelopathic Effects of Lichen Secondary Metabolites and Their Naturally Occurring Mixtures on Cultures of Aposymbiotically Grown Lichen Photobiont Trebouxia Erici (Chlorophyta). *S. Afr. J. Bot.* 2014, 93, 86–91, doi:10.1016/j.sajb.2014.03.015.

45. Peres, M.T.; Cândido, A.C.S.; Facenda, O.; Gianini, A.S.; Honda, N.K.; Peres, M.T.; Cândido, A.C.S.; Facenda, O.; Gianini, A.S.; Honda, N.K. Phytotoxicity of Perlatolic Acid and Derivatives. *Braz. Arch. Biol. Technol.* 2016, 59, doi:10.1590/1678-4324-2016160118.

46. Goga, M.; Antreich, S.J.; Bačkor, M.; Weckwerth, W.; Lang, I. Lichen Secondary Metabolites Affect Growth of Physcomitrella Patens by Allelopathy. *Protoplasma* 2017, 254, 1307–1315, doi:10.1007/s00709-016-1022-7.

47. Slansky, F., Jr. Effect of the Lichen Chemicals Atranorin and Vulpinic Acid upon Feeding and Growth of Larvae of the Yellow-Striped Armyworm, Spodoptera Orinchogallii. *Environ. Entomol.* 1979, 8, 863–868, doi:10.1093/ee/8.5.865.

48. Cook, W.E.; Raisbeck, M.F.; Cornish, T.E.; Williams, E.S.; Brown, B.; Hiatt, G.; Kreeger, T.J. Paresis and Death in Elk (Cervus Elaphus) Due to Lichen Intoxication in Wyoming. *J. Wildl. Dis.* 2007, 43, 498–503, doi:10.7589/0090-3558-43.3.498.

49. Cetin, H.; Tufan-Cetin, O.; Turk, A.O.; Tay, T.; Canadan, M.; Yanikoglu, A.; Sumbul, H. Insecticidal Activity of Major Lichen Compounds, (+)- and (-)-Usnic Acid, against the Larvae of House Mosquito, Culex Pipiens L. *Environ. Pollut.* 2017, 225–234, doi:10.1016/j.envpol.2017.04.022.

50. James, P.W.; Cladonia, P.B. Lichens of Great Britain and Ireland; Smith, C.W., Aptom, R., Copps, B.J., Fletcher, A., Gilbert, O.L., James, P.W., Wolseley, P.A., Eds; The British Lichen Society & The Natural History Museum: London, UK, 2009; pp. 309–338.

51. Nimis, P.L.; Martellos, S. *ITALIC 6.0*—The Information System on Italian Lichens Available online: http://dryades.units.it/italic (accessed on 20 January 2021).

52. Elix, J.A. *A Catalogue of Standardized Chromatographic Data and Biosynthetic Relationships for Lichen Substances*; The Author: Canberra, Australia, 2014.

53. Pipatti, S. *Flora d’Italia*; Edagricole: Bologna, Italy, 1982.
60. Dray, S.; Legendre, P. Testing the Species Traits–Environment Relationships: The Fourth-Corner Problem Revisited. *Ecology* 2008, 89, 3400–3412, doi:10.1890/08-0349.1.

61. Brown, A.M.; Warton, D.I.; Andrew, N.R.; Binns, M.; Cassis, G.; Gibb, H. The Fourth-Corner Solution—Using Predictive Models to Understand How Species Traits Interact with the Environment. *Methods Ecol. Evol.* 2014, 5, 344–352, doi:10.1111/2041-210X.12163.

62. Wang, Y.; Naumann, U.; Eddelbuettel, D.; Wilshire, J.; Warton, D.; Byrnes, J.; dos Santos Silva, R.; Niku, J.; Renner, I.; Wright, S. *Mvabund: Statistical Methods for Analyzing Multivariate Abundance Data*; R Package Version 4.1.3.; R Foundation for Statistical Computing: Vienna, Austria, 2020.

63. Rao, D.N.; LeBlanc, F. A Possible Role of Atranorin in the Lichen Thallus. *Bryologist* 1965, 68, 284–289, doi:10.2307/3240598.

64. Paas, S. Die Erdfechtenvegetation Nordwestdeutschlands Und Einiger-Randgebiete. *Bibl. Lichenol.* 1997, 66, 1–222.

65. Osyczka, P.; Rola, K. Cladonia Lichens as the Most Effective and Essential Pioneers in Strongly Contaminated Slag Dumps. *Cent. Eur. J. Biol.* 2013, 8, 876–887, doi:10.2478/s11535-013-0210-0.

66. Rola, K.; Osyczka, P.; Nobis, M. Cryptogamic Communities Dominated by the Lichen Cladonia Rei—A Case Study of Polish Post-Smelting Dumps in a Worldwide Context. *Herzogia* 2014, 27, 121–135, doi:10.13158/heria.27.1.2014.121.

67. Benesperi, R.; Nascimbene, J.; Lazzaro, L.; Bianchi, E.; Tepsich, A.; Longinotti, S.; Giordani, P. Successful Conservation of the Endangered Forest Lichen Lobaria Pulmonaria Requires Knowledge of Fine-Scale Population Structure. *Fungal Ecol.* 2018, 33, 65–71, doi:10.1016/j.funeco.2018.01.006.

68. Bayfield, N.G.; Urquhart, U.H.; Cooper, S.M. Susceptibility of Four Species of Cladonia to Disturbance by Trampling in the Cairngorm Mountains, Scotland. *J. Appl. Ecol.* 1981, 18, 303–310, doi:10.2307/2402497.

69. Heggenes, J.; Odland, A.; Chevalier, T.; Ahlberg, J.; Berg, A.; Larsson, H.; Bjerketvedt, D.K. Herbivore Grazing—or Trampling? Trampling Effects by a Large Ungulate in Cold High-Latitude Ecosystems. *Ecol. Evol.* 2017, 7, 6423–6431, doi:10.1002/ece3.3130.

70. Hauck, M.; Huneck, S. The Putative Role of Fumarprotocetraric Acid in the Manganese Tolerance of the Lichen Lecanora Conizaeoides. *Lichenologist* 2007, 39, 301–304, doi:10.1017/S0024282907006664.

71. Ranković, B.; Mišić, M. The Antimicrobial Activity of the Lichen Substances of the Lichens Cladonia Furcata, Ochrolechia Androgyyna, Parmelia Caperata and Parmelia Conspresa. *Biotechnol. Biotechnol. Equip.* 2008, 22, 1013–1016, doi:10.1080/13102818.2008.10817601.

72. Aptroot, A.; van Herk, C.M. Further Evidence of the Effects of Global Warming on Lichens, Particularly Those with Trentepohlia Phycobionts. *Environ. Pollut.* 2007, 146, 293–298, doi:10.1016/j.envpol.2006.03.018.

73. Concostrina-Zubiri, L.; Valencia, E.; Ochoa, V.; Gorazlo, B.; Mendoza, B.J.; Maestre, F.T. Species-Specific Effects of Biocrust-Forming Lichens on Soil Properties under Simulated Climate Change Are Driven by Functional Traits. *New Phytol.* 2020, doi:10.1111/nph.17143.

74. Gauslaa, Y. Rain, Dew, and Humid Air as Drivers of Morphology, Function and Spatial Distribution in Epiphytic Lichens. *Lichenologist* 2014, 46, 1–16, doi:10.1017/S0024282913000753.