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Bark traits affect epiphytic bryophyte community assembly in a temperate forest

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Abstract

Bark traits of trees often serve as a key factor determining the community structure of epiphytes. However, the extent to which barks modulate the relative importance of abiotic and biotic assembly processes of epiphytes is poorly understood. Here, using a community phylogenetic approach, we aimed to infer the assembly processes of epiphytic mosses and liverworts on tree species with varying bark traits in a temperate forest of central Japan. We observed a total of 56 moss and 35 liverwort species on 150 trees. Moss communities showed decreasing species richness and a tendency toward phylogenetic overdispersion, that is, higher phylogenetic diversity than expected by chance, in relation to increasing bark roughness and acidity. Along the same bark gradients, liverwort communities became phylogenetically clustered. Species richness of both mosses and liverworts increased with the nitrogen content of barks. The results indicate non-random assembly processes such as abiotic filtering associated with environmental harshness and microhabitat variety determined by barks. Our findings imply that bark traits modulate community assembly processes through which epiphyte diversity is maintained.

Keywords Biodiversity · Competition · Environmental filtering · Evolutionary history · Mosses · Phylogenetic diversity

Introduction

Epiphytes are essential components of forest biodiversity (Burns and Zotz 2010; Mendieta-Leiva and Zotz 2015; Tatsumi et al. 2017). Understanding the processes through which epiphyte species assemble on host trees provides a crucial step toward developing effective conservation strategies and preserving the functional roles epiphytes play in forest ecosystems (Ellis 2012). Notably, the characteristics of barks have been recognized as a key determinant of epiphyte community structure (Callaway et al. 2002; Wyse and

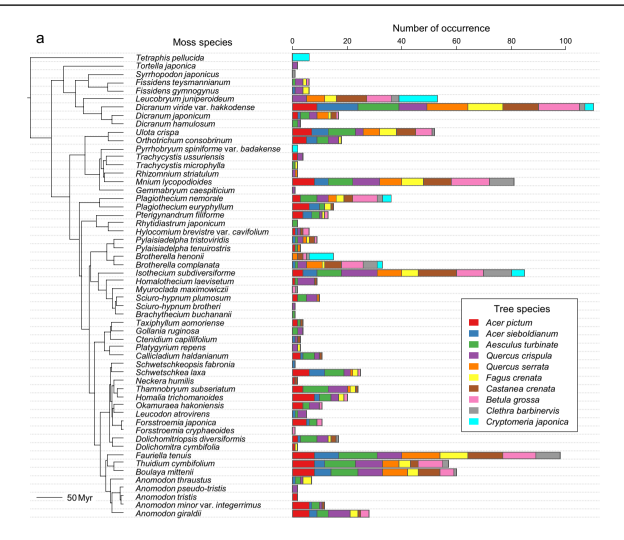
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Burns 2011; Mendieta-Leiva and Zotz 2015). However, despite extensive research describing the composition and distribution patterns of epiphytes on various barks, comparatively little is known about how bark traits modulate the relative importance of assembly processes (e.g., abiotic filtering or biotic interactions) driving such patterns (Spicer and Woods 2022).

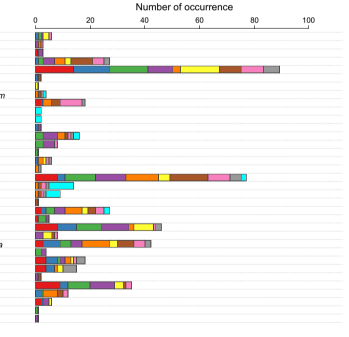
Phylogenetic diversity has been widely employed to account for evolutionary and ecological relatedness among species within a community. In particular, the sign and magnitude of phylogenetic diversity deviating from null expectations have commonly served as proxies representing the relative strengths of different assembly processes (Webb et al. 2002; Cavender-Bares et al. 2004; Gerhold et al. 2015). Under evolutionary niche conservatism, phylogenetic diversity lower or higher than expected by chance, referred to as phylogenetic clustering and overdispersion, respectively, has been interpreted as indicative of abiotic and biotic assembly (Webb 2000; Webb et al. 2002). In combination with demonstrable environmental gradients, phylogenetic diversity can provide insights into ecological processes through which species assemble into communities (Cadotte and Tucker 2017; Cadotte et al. 2019; Tatsumi et al. 2019).



b



Liverwort species
Metzgeria temperata
Trichocoleopsis sacculata
Mylia verrucosa
Syzygiella autumnalis
Liochlaena subulata
Cephaloziella spinicaulis
Odontoschisma denudatum
Schiffneria hyalina
Cephalozia otaruensis
Cahyoogeia tosana
Blepharostoma minus
Plagiochila solophila
Plagiochila solophila
Plagiochila solophila
Plagiochila solophila
Neteroscyphus pinous
Chiloscyphus profundus
Chiloscyphus minor
Bazzania tridens
Bazzania denudata
Porella vernicosa
Macvicaria ulophylla
Radula japonica
Radula japonica
Frullania tamarisci subsp. obscura
Frullania davurica
Nipponelejeunea pilifera
Lejeunea ulicina
Lejeunea sinosa
Acrolejeunea sinosa
Acrolejeunea sinosa



◄Fig. 1 Reconstructed phylogenies and the number of occurrences of a mosses and b liverworts. Three chloroplast genes (*rbcL*, *rps*4, and *trnL*-F) were used for reconstruction

For example, the acidity of barks can serve as a gradient determining the environmental harshness for epiphytes (Mitchell et al. 2021). Abiotic filtering can thus become more pronounced on barks with lower pH, resulting in phylogenetic clustering of communities under niche conservatism and overdispersion under convergent evolution (Webb 2000; Webb et al. 2002). On the other hand, the significance of biotic interactions often increase with resource availability (Briones et al. 1998). Therefore, on barks that can retain resources such as water (Zamfir and Goldberg 2000), competitive exclusion among closely related species may become a dominant assembly process, leading to phylogenetic overdispersion under niche conservatism (Webb 2000; Cavender-Bares et al. 2004; Cadotte et al. 2019).

Here, we explore community assembly of epiphytic bryophytes on barks. Specifically, we analyze phylogenetic diversity of mosses and liverworts, which constitute two major clades of bryophytes, on multiple tree species that represent gradients of bark traits in a temperate forest. Using null models, we test whether communities show tendency toward phylogenetic clustering or overdispersion along the gradients. Based on the phylogenetic community structure observed, we infer underlying assembly processes and their links to bark traits.

Methods

Study site and tree species

This study was conducted in the Ashiu Forest Research Station of the Kyoto University, western Japan (4186 ha; 35.3° N, 135.8° E; 355-959 m elevation) (Fig. S1). The study site is covered by primary forests and part of it is designated as a National Bryophyte Heritage Site of Japan for the rich bryophyte flora. The study site is located in a warm- and cool-temperate ecotone dominated by an evergreen conifer *Cryptomeria japonica* and deciduous broadleaves including *Aesculus turbinata, Fagus crenata*, and *Quercus crispula*. The mean monthly temperature ranges from $-0.4 \, ^{\circ}$ C in January to 24.0 $^{\circ}$ C in August. The mean annual precipitation is 2568 mm.

We selected 10 tree species for our study: Acer pictum subsp. mono, Acer sieboldianum, Aesculus turbinata, Betula grossa, Castanea crenata, Clethra barbinervis, Cryptomeria japonica, Fagus crenata, Quercus crispula, and Quercus serrata. These species were selected to cover a large variety of bark traits as possible. For each tree species, we surveyed bryophyte communities on 15 trees, totalling 150 trees, in six plots distributed across the study area (Fig. S1). The surveyed trees were selected in such a way that most tree species had similar levels of variation in tree sizes (except for *A. sieboldianum* and *C. barbinervis* which are shrub species; Fig. S2) and among-individual geographical distances (Fig. S1). We selected trees in closed-canopy stands that were at least ~ 20 m away from the nearest forest edge to minimize the potential variation in light environment. We measured the diameters at breast height (DBH) of the trees using diameter tapes.

Bryophyte survey and diversity

In October 2016, we surveyed epiphytic bryophytes in four 10-cm wide, 200-cm high quadrats positioned at the cardinal directions of each tree, totalling 8000 cm² per tree. We recorded the presence or absence of bryophyte species on each tree. Species were identified in the field or in the laboratory under a microscope. To prevent epigeic species from being included, the quadrats were placed approximately 5–30 cm above the ground surface, depending on the inclination of stems and slopes. We used quadrats with a fixed size so that bryophyte diversity would be comparable among trees of different sizes, without being affected by variation in the survey area per se. All trees were surveyed at their cardinal directions to keep the possible influences of aspect consistent.

A bryophyte phylogeny was reconstructed based on three chloroplast genes (*rbcL*, *rps*4, and *trnL*-F), which are commonly used in bryophyte phylogenetics (Stech and Quandt 2010). See Supplementary text 1 for details on phylogeny reconstruction.

We quantified phylogenetic diversity of bryophyte communities using mean pairwise distance (MPD) (Webb 2000). We calculated the standardized effect size of MPD, referred to as net relatedness index (NRI), based on null modelling (Webb et al. 2002). The NRI was defined as $-1 \cdot (x - \mu_{null})/\sigma_{null}$, where x is the observed MPD, μ_{null} is the mean MPD of a null distribution, and σ_{null} is the standard deviation of a null distribution (Webb et al. 2002). The null distributions were generated based on 999 iterations of presence–absence randomizations across 150 communities using the independent swap algorithm (Gotelli 2000). Randomizations were conducted separately for mosses and liverworts. To examine for possible effects of tree sizes on bryophyte community structure, we compared models with and without DBH as an explanatory variable.

Bark traits

For each of the 10 tree species, we measured bark roughness, water holding capacity, pH, and inorganic nitrogen content. These traits were selected based on previous research that

has shown their associations with epiphyte community structure (Gustafsson and Eriksson 1995; reviewed by Ellis 2012). We measured each trait on three trees per species and used the mean value for statistical analyses. See Supplementary text 2 for details of the measurement methods and Table S1 for the observed bark trait values. To account for correlations between some pairs of traits (Table S2), we performed a principal component analysis to derive composite measures of bark traits.

Regression analyses

We tested the changes in bryophyte species richness along bark trait gradients using generalized linear mixed models with a Poisson error distribution and a log-link function. Changes in MPD were tested using log-normal linear mixed models. Changes in NRI were tested using linear mixed models. We included 'plots' as a random variable in all models. We used R 4.3.0 (R Core Team 2023) for all statistical analyses.

Results and discussion

We observed a total of 56 moss and 35 liverwort species on 150 trees, with 1016 occurrences of mosses and 515 occurrences of liverworts (Fig. 1). Regarding bark traits, more than half of the variation was captured by the first principal component (PC 1) (Fig. 2). Bark roughness and pH showed a negative correlation (Table S2), likely due to the tendency for rougher barks to capture more atmospheric materials, resulting in increased acidity (Oka et al. 2021). The PC 1 represented a composite gradient of bark roughness, pH, and water holding capacity (Fig. 2, Table S3), along which we found significant changes in species richness of mosses (Figs. 3a, S3). This result may reflect the impact of bark acidity (ranging from pH 4.16 to 6.18; Table S1), which often reduce germination and growth rates of mosses (Löbel and Rydin 2010), thereby leading to a decrease in species richness (Kaufmann et al. 2019; Mitchell et al. 2021).

The MPD and NRI of mosses increased and decreased along the PC 1 axis (Fig. 3e, 3i), respectively, suggesting changes in assembly processes. Specifically, moss communities became phylogenetically overdispersed (NRI < 0) on rough and acid barks (Figs. 3i, S3i, S3j); that is, communities became composed of species belonging to a larger variety of lineages than would be expected by chance. A possible reason for this pattern is that rough barks, which often have a greater heterogeneity of microhabitats than smooth barks (Wyse and Burns 2011; Lamit et al. 2015), allowed moss species from different lineages favouring different microhabitats to coexist. Alternatively, the observed pattern of overdispersion (Fig. 3i) may reflect independent adaptations

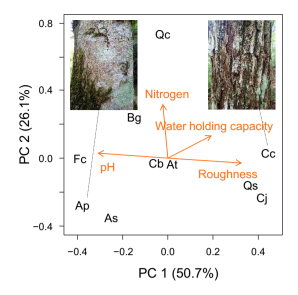


Fig. 2 Ordination plot of tree bark traits. Values in parentheses indicate the proportion of variation explained by the first and second principal components (PC). Ap=*Acer pictum* subsp. *mono*, As=*Acer sieboldianum*, At=*Aesculus turbinata*, Bg=*Betula grossa*, Cc=*Castanea crenata*, Cb=*Clethra barbinervis*, Cj=*Cryptomeria japonica*, Fc=*Fagus crenata*, Qc=*Quercus crispula*, and Qs=*Quercus serrata*. The pictures show barks of Ap and Cc as examples with contrasting barks

among moss lineages to harsh environments. Convergent evolution of plants to harsh environments is a commonly observed phenomenon, including adaptations of alpine plants to high elevations (Bryant et al. 2008) and mangrove trees to salinity (Shi et al. 2005). In our study, we observed moss species from distant lineages (e.g., *Tetraphis pellucida* [order Tetraphidales], *Dicranum viride* var. *hakkodense* [Dicranales], *Brotherella complanata* [Hypnales]) on *C. japonica* trees (Fig. 1a) that have rough, acidic, and wet barks (Fig. 2). Among these bark traits, acidity (pH=4.16 for *C. japonica*) may have acted as an environmental filter representing harshness, given the fact that many extant moss species favour neutral pH (Robinson et al. 1989).

Contrary to our expectation, we found no significant effect of water holding capacity on moss community assembly (Fig. S3k). Water often serves as key resource for which mosses compete (Zamfir and Goldberg 2000). We therefore expected biotic interactions to intensify with increasing water availability on barks, leading to phylogenetic overdispersion (Webb 2000; Cavender-Bares et al. 2004; Cadotte et al. 2019). However, such competition-mediated assembly was not evident in our study site, possibly because it receives ample precipitation, making water a non-limiting resource regardless of bark traits. It is also worth noting that the water holding capacity only serves as a rough proxy for hydrological environment on barks. Future studies are thus needed to examine whether other hydrological variables, such as

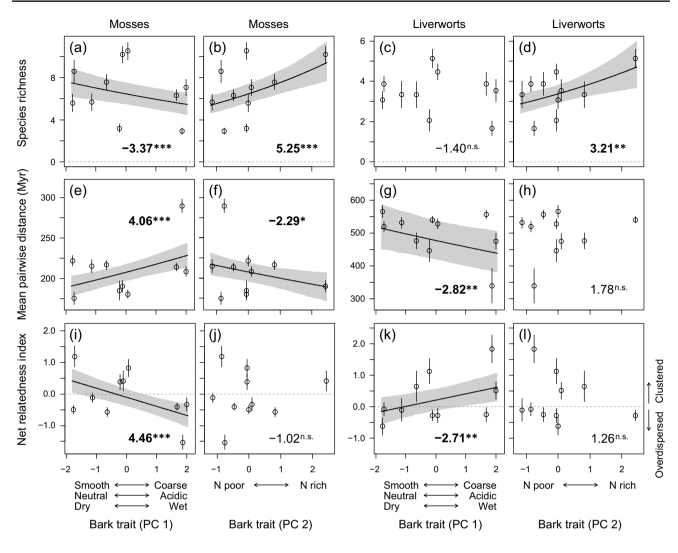


Fig. 3 Changes in bryophyte diversity along bark trait gradients. **a**, **b**, **c**, **d** species richness, **e**, **f**, **g**, **h** mean phylogenetic diversity, and **i**, **j**, **k**, **l** net relatedness indices of mosses and liverworts along the first and second principal components (PC 1 and PC 2) of bark traits. Circles and vertical bars represent the mean and standard errors for each tree species (n = 15 trees surveyed for each of 10 tree species,

totalling n = 150). Lines show fitted models with significant slopes (P < 0.05). Grey areas represent 95% confidence intervals of the fitted models. The values in each panel indicate the z-statistics of the slope of the fitted model. Significance: *, P < 0.05; **, P < 0.01; ***, P < 0.001; not significant (n.s.), $P \ge 0.05$

cortical runoff which quantifies stem flow in relation to treecanopy morphology (González-Mancebo et al. 2003), can better explain the impacts of water availability on epiphytic bryophyte communities.

In contrast to mosses, liverwort communities showed decreasing MPD and a tendency toward phylogenetic clustering (NRI>0) in relation to the increased roughness and acidity of barks (Figs. 3g, 3k, S4). According to Fiz-Palacios et al. (2011), liverworts experienced a relatively slow diversification process from the mid-Cretaceous to the early Cenozoic era, during which mosses and ferns rapidly diversified in habitats created by angiosperms (as proposed by the "shadow of angiosperms" hypothesis; Schneider et al. 2004). It is possible that liverwort species, due to this constrained

niche evolution, have maintained their specific habitats over time, leading to phylogenetic niche conservatism. The combination of closely related species having similar habitat preferences and environmental filtering associated with rough and acidic barks could have contributed to the observed pattern of phylogenetic clustering (Figs. 3k, S4i, S4j). A logical next step of our study would be to test the assumption of niche conservatism based on relationships between functional traits, phylogeny, and community structure of bryophytes.

Ample evidence has shown that excess nitrogen owing to human activities (e.g., fertilization and atmospheric deposition) can reduce bryophyte richness (Oishi and Hiura 2017), both directly by posing toxic impacts and indirectly by enhancing the competitiveness of vascular plants (Turetsky 2003). In contrast, we found increasing species richness of both mosses and liverworts in response to inorganic nitrogen content (Figs. 3b, 3d, S3d, S4d). The observed pattern could be attributed to the fact that our study was conducted in a primary temperate forest where anthropogenic nitrogen inputs are kept minimal and epiphytic vascular plants are rare. In a nitrogen-limited environment with few competitors like our study site, barks with a high nitrogen content may serve as a hotspot for bryophytes. Nevertheless, it should be noted that we observed low levels of MPD on nitrogen-rich barks (Fig. 3f), indicating that only a restricted number of moss lineages could utilize such habitats.

While we identified significant relationships between bark traits and bryophyte community patterns (Fig. 3), there are still some uncertainties associated with our findings. Specifically, the absolute values of NRI that we observed were smaller than 1.96 (Fig. 3i, 3j, 3k, 3l), suggesting relatively weak signals of community assembly processes (Webb et al. 2002). Moreover, we were only able to measure bark traits at the species level, without accounting for the possible variations that derive from individual-tree characteristics (e.g., tree size) and environments (Burns and Zotz 2010; Lamit et al. 2015; Rosell 2019). In fact, some bark traits showed relatively large levels of variations among trees (Table S1). Nevertheless, we confirmed that the signs and statistical significance of the estimated relationships between bark traits and bryophyte communities remained consistent whether or not DBH was included as an explanatory variable (Table S4). Moving forward, future studies should verify the robustness of the relationships using a larger dataset than ours, while accounting for variations in barks and environmental conditions at individual-tree levels.

In this study, we found that epiphytic bryophyte communities assemble non-randomly along gradients of bark traits (Fig. 3). Our study provides an important step toward understanding how host trees, as living patches, determine epiphyte assembly processes. Trees with different bark traits respond differently to environments (Rosell and Olson 2014), implying that potential future changes in tree bark diversity under environmental change can have cascading effects on epiphytic bryophyte diversity. While our study was based on snapshot data, future research should incorporate long-term monitoring and investigate the dynamics of host trees and epiphytes over time. Doing so would provide a more comprehensive understanding of epiphyte community assembly, which is essential for informing effective strategies for their conservation in the face of changing environments.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s11258-023-01363-9.

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Author contributions ST conceived and led the study. All authors conducted the field survey. TO identified the bryophyte species. ST and WAA measured the bark traits. ST analyzed the data. ST wrote the manuscript with inputs from other authors.

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Data availability The bryophyte community matrix (presence-absence of 91 bryophyte species on 150 trees), the reconstructed bryophyte phylogenetic tree (Newick format), and the bark trait data are available at FigShare (https://doi.org/10.6084/m9.figshare.23673258).

Declarations

Conflict of interest The authors declare no conflict of interest.

References

- Briones O, Montaña C, Ezcurra E (1998) Competition intensity as a function of resource availability in a semiarid ecosystem. Oecologia 116:365–372. https://doi.org/10.1007/s004420050599
- Bryant JA, Lamanna C, Morlon H, Kerkhoff AJ, Enquist BJ, Green JL (2008) Microbes on mountainsides: contrasting elevational patterns of bacterial and plant diversity. Proc Natl Acad Sci USA 105:11505–11511. https://doi.org/10.1073/pnas.0801920105
- Burns KC, Zotz G (2010) A hierarchical framework for investigating epiphyte assemblages: networks, meta-communities, and scale. Ecology 91:377–385. https://doi.org/10.1890/08-2004.1
- Cadotte MW, Carboni M, Si X, Tatsumi S (2019) Do traits and phylogeny support congruent community diversity patterns and assembly inferences? J Ecol. https://doi.org/10.1111/1365-2745.13247
- Cadotte MW, Tucker CM (2017) Should environmental filtering be abandoned? Trends Ecol Evol 32:429–437. https://doi.org/10. 1016/j.tree.2017.03.004
- Callaway RM, Reinhart KO, Moore GW, Moore DJ, Pennings SC (2002) Epiphyte host preferences and host traits: mechanisms for species-specific interactions. Oecologia 132:221–230. https://doi.org/10.1007/s00442-002-0943-3
- Cavender-Bares J, Ackerly DD, Baum DA, Bazzaz FA (2004) Phylogenetic overdispersion in Floridian oak communities. Am Nat 163:823–843. https://doi.org/10.1086/386375
- Ellis CJ (2012) Lichen epiphyte diversity: a species, community and trait-based review. Perspect Plant Ecol Evol Syst 14:131–152. https://doi.org/10.1016/j.ppees.2011.10.001
- Fiz-Palacios O, Schneider H, Heinrichs J, Savolainen V (2011) Diversification of land plants: insights from a family-level phylogenetic analysis. BMC Evol Biol 11:341. https://doi.org/10.1186/ 1471-2148-11-341

- Gerhold P, Cahill JF, Winter M, Bartish IV, Prinzing A (2015) Phylogenetic patterns are not proxies of community assembly mechanisms (they are far better). Funct Ecol 29:600–614. https://doi.org/10. 1111/1365-2435.12425
- González-Mancebo JM, Losada-Lima A, McAlister S (2003) Host specificity of epiphytic bryophyte communities of a laurel forest on Tenerife (Canary islands, Spain). Bryologist 106:383–394
- Gotelli NJ (2000) Null model analysis of species co-occurrence patterns. Ecology 81:2606–2621. https://doi.org/10.2307/177478
- Gustafsson L, Eriksson I (1995) Factors of importance for the epiphytic vegetation of aspen *Populus tremula* with special emphasis on bark chemistry and soil chemistry. J Appl Ecol 32:412. https:// doi.org/10.2307/2405107
- Kaufmann S, Weinrich T, Hauck M, Leuschner C (2019) Vertical variation in epiphytic cryptogam species richness and composition in a primeval *Fagus sylvatica* forest. J Veg Sci 30:881–892. https:// doi.org/10.1111/jvs.12775
- Lamit LJ, Lau MK, Næsborg RR, Wojtowicz T, Whitham TG, Gehring CA (2015) Genotype variation in bark texture drives lichen community assembly across multiple environments. Ecology 96:960– 971. https://doi.org/10.1890/14-1007.1
- Löbel S, Rydin H (2010) Trade-offs and habitat constraints in the establishment of epiphytic bryophytes. Funct Ecol 24:887–897. https:// doi.org/10.1111/j.1365-2435.2010.01705.x
- Mendieta-Leiva G, Zotz G (2015) A conceptual framework for the analysis of vascular epiphyte assemblages. Perspect Plant Ecol Evol Syst 17:510–521. https://doi.org/10.1016/j.ppees.2015.09. 003
- Mitchell RJ, Hewison RL, Beaton J, Douglass JR (2021) Identifying substitute host tree species for epiphytes: the relative importance of tree size and species, bark and site characteristics. Appl Veg Sci 24:1–13. https://doi.org/10.1111/avsc.12569
- Oishi Y, Hiura T (2017) Bryophytes as bioindicators of the atmospheric environment in urban-forest landscapes. Landsc Urban Plan 167:348–355. https://doi.org/10.1016/j.landurbplan.2017.07.010
- Oka A, Takahashi J, Endoh Y, Seino T (2021) Bark effects on stemflow chemistry in a Japanese temperate forest I. The role of bark surface morphology. Front for Glob Change 4:1–10. https://doi. org/10.3389/ffgc.2021.654375
- Robinson AL, Vitt DH, Timoney KP (1989) Patterns of community structure and morphology of bryophytes and lichens relative to edaphic gradients in the subarctic forest-tundra of northwestern Canada. Bryologist 92:495–512. https://doi.org/10.2307/3243674
- Rossell JA (2019) Bark in woody plants: understanding the diversity of a multifunctional structure. Integr Comp Biol 59:535–547
- Rosell JA, Olson ME (2014) The evolution of bark mechanics and storage across habitats in a clade of tropical trees. Am J Bot 101:764–777. https://doi.org/10.3732/ajb.1400109

- Schneider H, Schuettpelz E, Pryer KM, Cranfill R, Magallón S, Lupia R (2004) Ferns diversified in the shadow of angiosperms. Nature 428:553–557. https://doi.org/10.1038/nature02361
- Shi S, Huang Y, Zeng K, Tan F, He H, Huang J, Fu Y (2005) Molecular phylogenetic analysis of mangroves: Independent evolutionary origins of vivipary and salt secretion. Mol Phylogenet Evol 34:159–166. https://doi.org/10.1016/j.ympev.2004.09.002
- Spicer ME, Woods CL (2022) A case for studying biotic interactions in epiphyte ecology and evolution. Perspect Plant Ecol Evol Syst 54:125658. https://doi.org/10.1016/j.ppees.2021.125658
- Stech M, Quandt D (2010) 20,000 species and five key markers: the status of molecular bryophyte phylogenetics. Phytotaxa 9:196–228. https://doi.org/10.11646/phytotaxa.9.1.11
- Tatsumi S, Cadotte MW, Mori AS (2019) Individual-based models of community assembly: neighbourhood competition drives phylogenetic community structure. J Ecol 107:735–746. https://doi.org/ 10.1111/1365-2745.13074
- Tatsumi S, Ohgue T, Azuma W, Tuovinen V, Imada Y, Mori AS, Thor G, Ranlund Å (2017) Tree hollows can affect epiphyte species composition. Ecol Res. https://doi.org/10.1007/ s11284-017-1468-x
- Turetsky MR (2003) New frontiers in bryology and lichenology: the role of bryophytes in carbon and nitrogen cycling. Bryologist 106:395–409
- Webb CO (2000) Exploring the phylogenetic structure of ecological communities: an example for rain forest trees. Am Nat 156:145– 155. https://doi.org/10.1086/303378
- Webb CO, Ackerly DD, McPeek MA, Donoghue MJ (2002) Phylogenies and community ecology. Annu Rev Ecol Syst 33:475–505. https://doi.org/10.1146/annurev.ecolsys.33.010802.150448
- Wyse SV, Burns BR (2011) Do host bark traits influence trunk epiphyte communities? N Z J Ecol 35:296–301
- Zamfir M, Goldberg DE (2000) The effect of initial density on interactions between bryophytes at individual and community levels. J Ecol 88:243–255. https://doi.org/10.1046/j.1365-2745.2000. 00442.x

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