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Permanent residents or temporary lodgers: characterizing intracellular bacterial communities in the siphonous green alga *Bryopsis*

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The ecological success of giant celled, siphonous green algae in coastal habitats has repeatedly been linked to endophytic bacteria living within the cytoplasm of the hosts. Yet, very little is known about the relative importance of evolutionary and ecological factors controlling the intracellular bacterial flora of these seaweeds. Using the marine alga *Bryopsis* (Bryopsidales, Chlorophyta) as a model, we explore the diversity of the intracellular bacterial communities and investigate whether their composition is controlled by ecological and biogeographic factors rather than the evolutionary history of the host. Using a combination of 16S rDNA clone libraries and denaturing gradient gel electrophoresis analyses, we show that *Bryopsis* harbours a mixture of relatively few but phylogenetically diverse bacterial species. Variation partitioning analyses show a strong impact of local environmental factors on the presence of *Rickettsia* and *Mycoplasma* in their association with *Bryopsis*. The presence of Flavobacteriaceae and Bacteroidetes, on the other hand, reflects a predominant imprint of host evolutionary history, suggesting that these bacteria are more specialized in their association. The results highlight the importance of interpreting the presence of individual bacterial phylotypes in the light of ecological and evolutionary principles such as phylogenetic niche conservatism to understand complex endobiotic communities and the parameters shaping them.

1. Introduction

Variation in traits across species or populations is influenced by their ecology and evolutionary history [1]. Organisms are shaped by the environment in which they live, with species residing in similar environments having common adaptations [2]. They are also the product of their evolutionary history, and closely related species have the tendency to be more similar than distantly related species [3]. This tendency for related species to resemble each other more in a trait than expected by chance is referred to as phylogenetic signal or phylogenetic conservatism [4]. Applying these principles to host–bacterial relationships, one might presume that obligate bacteria are phylogenetically structured, while facultative endobiotic bacteria are expected to be more randomly dispersed among host species ([5]; figure 1). In this study, we assess for the first time, to our knowledge, the combined effect of host dependency, ecology and biogeography on the structure of a complex endobiotic community in an algal model.

Marine macroalgae (seaweeds) are commonly associated with bacteria that either live on the surface or in the cytoplasm and/or vacuolar systems of the cells [6–8]. These bacteria are able to influence the morphogenesis and life cycle of their algal host [9–11] and are linked with various metabolic functions such as the production of growth factors, fixed nitrogen and antimicrobial

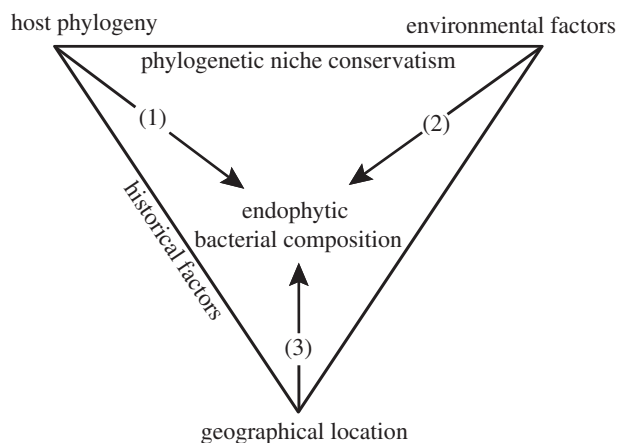


Figure 1. Relationships between host phylogeny, environment and geography on endophytic bacterial composition in *Bryopsis* seaweeds and relations between these three factors. (1) Phylogenetic structured variation, (2) ecological structured variation and (3) geographical structured variation. The shared influence of phylogeny and environment (1 + 2) is known as ‘phylogenetically structured environmental variation’.

compounds [12–14]. Siphonous green seaweeds, belonging to the Bryopsidales and Dasycladales of the chlorophytan class Ulvophyceae, consist of a single giant tubular cell and form a benevolent biotic environment for endobiotic bacterial communities [15–17]. The siphonous cells, which range from centimetres to metres in length, typically exhibit vigorous cytoplasmic streaming to transport organelles, photosynthates and nutrients [18]. Chisholm *et al.* [19] demonstrated that siphonous algae take up nutrients from the sediment by a root-like system containing intracellular bacteria and translocate them throughout the thallus. These cellular innovations alongside unique mechanisms of wounding response [20,21] and the close interactions with bacteria may provide a physiological explanation for the successful spread of invasive siphonous green algae such as *Caulerpa* and *Codium* in marine coastal habitats [19,22,23].

Very little is known about the factors controlling the presence of bacteria inside siphonous seaweeds. Two host-related mechanisms may affect the intracellular bacterial composition. First, siphonous seaweeds readily regenerate from protoplasts, facilitating environmental uptake of bacteria into the cell [24,25]. Second, endogenous bacteria can persist by vertical inheritance through gametes [26]. Beside the question of whether the endobionts are acquired vertically or horizontally from the environment, ecological parameters and geographical aspects may also need to be considered to explain the bacterial composition, as some bacteria (or hosts) are likely to be geographically restricted or occur only in particular niches. Although a previous study suggested that seaweed-associated bacterial communities are biogeographically structured [23], it is not known whether ecological or historical factors cause this structure.

The goal of this study is to investigate the relative roles of host, environment and geography in determining the intracellular bacterial flora of siphonous seaweeds, focusing on the genus *Bryopsis* (Bryopsidales, Chlorophyta) as a case study. This genus is known to harbour several types of endogenous bacteria and protocols are in place to study them [17,27]. *Bryopsis* is known to possess mechanisms for environmental uptake as well as vertical inheritance of bacteria [25,26]. This combination of features, combined with

the large collection of available cultures, makes the genus an ideal case study to address our goal. The experimental approach consisted of molecular characterization of host samples and their intracellular bacterial flora. The molecular identification of bacterial phylotypes, along with the host phylogeny and environmental data, was explored and analysed with statistical techniques designed to disentangle the effects of host phylogeny, geography and the external environment on the intracellular bacterial composition.

2. Material and methods

(a) Algal material

The 20 *Bryopsis* samples analysed in this study are listed in the electronic supplementary material, table S1, and their sampling sites are depicted in the electronic supplementary material, figure S1. All samples were transferred to and maintained as unialgal cultures under the conditions described by Hollants *et al.* [17].

(b) Molecular approach

Bryopsis samples were subjected to a surface sterilization step to eliminate epiphytic bacterial contamination [27] prior to total DNA extraction [28]. The host *rbcL* and bacterial 16S rRNA genes were PCR amplified as described by Hollants *et al.* [17]. The endophytic bacterial diversity was assessed by creating 16S rRNA gene clone libraries and performing nested PCR denaturing gradient gel electrophoresis (DGGE) analyses as described previously [17,25]. Sequences were submitted to EMBL under accession numbers HE648924–HE648948.

(c) Sequence data analyses

Bryopsis rbcL and bacterial 16S rRNA gene sequences were assembled, checked for chimaeras, compared with nucleotide databases and aligned as previously described [17]. Phylogenetic trees were inferred with maximum-likelihood implemented in PHYML v. 3.0 [29] and Bayesian inference using MRBAYES [30], via the University of Oslo Biportal website [31]. Both the analyses were performed under a HKY + G model as determined by the Akaike Information Criterion in JMODELTEST v. 0.1.1 [32]. Bacterial phylotypes or operational taxonomic units (OTUs) were identified based on 97 per cent sequence similarity.

(d) Statistical analysis

The influence of environmental, geographical and host phylogenetic factors on the endophytic bacterial diversity in *Bryopsis* was analysed using multivariate statistical and comparative phylogenetic approaches. The response table was represented by a presence/absence matrix of the seven bacterial phylotypes in the 20 host samples (figure 2). The three explanatory matrices (environment, geography and phylogeny) were prepared as follows. The environmental component was represented by seven macro-ecological variables (figure 2) extracted from Bio-ORACLE [33]. The geographical component was represented by a set of orthogonal spatial variables extracted from geographical coordinates by Moran’s Eigenvector Maps (MEM) analysis [34] using ‘codep’ in R [35]. The geographical matrix was represented by the first two eigenvectors, which were the only ones having positive eigenvalues (6.54 and 1.52). The phylogenetic component was expressed as principal coordinates via a principal coordinate analysis (PCoA) [36] computed from a distance matrix [37]. A corrected distance matrix of the *Bryopsis rbcL* alignment was calculated in MEGA [38]; the PCoA analysis was performed in PCO [39]. The phylogenetic matrix was represented by the

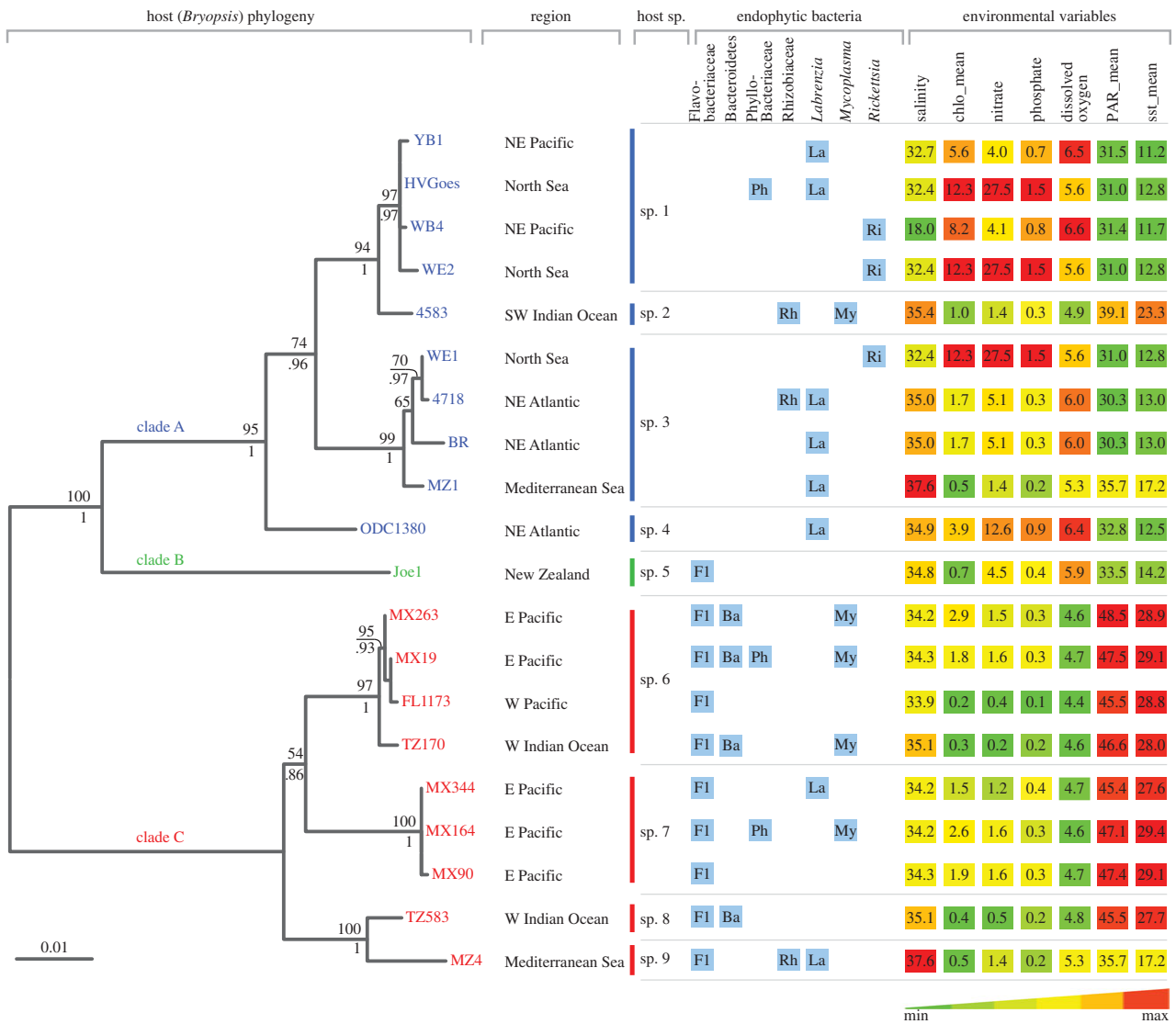


Figure 2. Endophytic diversity results, geographical data and environmental variables plotted against the *Bryopsis* host phylogram. The endophytic bacterial diversity, displayed by blue boxes, summarizes the diversity results from the 16S rRNA gene clone libraries and DGGE analyses. Environmental variables were extracted from the host sampling sites using Bio-ORACLE, salinity (PSS); chlo_mean, annual mean chlorophyll (mg m^{-3}), nitrate ($\mu\text{mol l}^{-1}$), phosphate ($\mu\text{mol l}^{-1}$), dissolved oxygen (ml l^{-1}); PAR_mean, annual mean photosynthetically available radiation ($\mu\text{mol m}^{-2} \text{s}^{-1}$); sst_mean, annual mean sea surface temperature ($^{\circ}\text{C}$). The phylogram on the left classifies the 20 algal samples for which endophytic bacterial data are available in nine different *Bryopsis* species and three distinct clades (i.e. A, B and C). These clades seem more consistent with the ecology of the host samples (environmental variables depicted on the right) than with their geographical origin (sample region). Maximum-likelihood bootstrap values and Bayesian inference posterior probabilities, respectively, are indicated above and below the branch nodes. The scale bar indicates 0.01 nucleotide changes per nucleotide position. NE, northeast; SW, southwest; E, east; W, west.

first four principal coordinates, representing 98 per cent of the total variation.

To study the influence of environment, geography and host phylogeny on the endophytic bacterial diversity, we first performed data ordinations and calculated phylogenetic signals of the bacterial community composition. Ordination of *Bryopsis* samples based on endophytic bacterial community composition was performed using a principal component analysis (PCA) in CANOCO for WINDOWS v. 4.5 [40]. Environmental variables were plotted on the PCA graph as electronic supplementary material. Phylogenetic signal was assessed for: (i) the environmental variables, (ii) the geography, (iii) the total endophytic bacterial community (i.e. represented by principal components 1 and 2 calculated as described above), and (iv) the presence/absence data of each of the endophytic bacterial OTUs separately. p -values were calculated using randomizations of the K-statistic [41] in the R package Picante [42] (for (i)–(iii)) and the D statistic [43] in the R package 'caper' (<http://cran.r-project.org/web/packages/caper/>) (for (iv)). We quantified the common and unique influences of host phylogeny, geography and environment

on the endophytic flora variation using variation partitioning analyses [2,44], using the varpart function in the R package 'vegan'. The total bacterial diversity as well as presence/absence data of the seven individual phylotypes was considered separately as response tables. We performed variation partitioning analyses using three (phylogeny, environment and geography) and two (phylogeny and environment) explanatory tables, respectively.

3. Results

(a) *Bryopsis* host phylogeny

Based on the phylogenetic analysis of host *rbcl* sequences (figure 2), we assigned the seaweed samples to nine *Bryopsis* species, numbered sp. 1–9. The host phylogeny shows three main clades. Clades A and B include *Bryopsis* samples isolated from cold to temperate regions, whereas clade C is warm-temperate to tropical. The phylogenetic signal in

annual mean sea surface temperature, as well as annual mean photosynthetically available radiation and dissolved oxygen levels, which are inversely proportional to each other, is statistically significant ($p < 0.01$, electronic supplementary material, table S2), suggesting that the structure of the *Bryopsis* phylogeny reflects temperature-related environmental variables. Conversely, geographical location (represented by MEM) did not show a significant phylogenetic structure (see the electronic supplementary material, table S2).

(b) Endophytic bacterial diversity

The results from the clone libraries and DGGE analyses showed the presence of seven unique endophytic bacterial phylotypes or OTUs within *Bryopsis* (see the electronic supplementary material, table S3). Five could be identified as Flavobacteriaceae (OTU-1), *Mycoplasma* (OTU-2), Bacteroidetes (OTU-3), Phyllobacteriaceae (OTU-4) and *Labrenzia* (OTU-7) species, which were previously shown to occur in *Bryopsis* [17] (see the electronic supplementary material, table S3 and figure S2). In addition, two new endophytic phylotypes were identified, OTU-5 and OTU-6 (see the electronic supplementary material, table S3 and figure S2). OTU-5 showed high sequence similarities with Rhizobiaceae strains isolated from root nodules of leguminous plants, and represents two distinct clusters that include *Rhizobium leguminosarum* and *Ensifer meliloti*-type strains, respectively. OTU-6 is allied to uncultured Rickettsiales bacteria associated with the coral *Montastraea faveolata* and the marine ciliate *Diophrys appendiculata*. All OTU-6 sequences formed a distinct and well-supported clade closely related to the genus *Rickettsia*, and most probably represent a new species based on their low sequence similarities (less than or equal to 93%) with *Rickettsia*-type strains.

(c) Endophytic bacterial composition

Figure 2 schematizes the endophytic bacterial diversity (blue boxes) in *Bryopsis*. Composition of the endophytic community varied between host species, and samples from the same host species harboured diverse combinations of one to four different endophytic phylotypes. Different host species with the same geographical origin commonly displayed differences in their intracellular bacterial community composition (e.g. samples MZ1 and MZ4). This apparent lack of correlation between total bacterial diversity and *Bryopsis* host species and geography is confirmed by the PCA plot, which illustrates that the ordination of the different *Bryopsis* species is not fully explained by their similarity in endophytic bacterial community composition (figure 3). This PCA plot, however, clearly indicates a correlation between the presence of individual endophytic phylotypes and certain environmental variables. Flavobacteriaceae, Bacteroidetes and *Mycoplasma* endophytes were only present in *Bryopsis* species isolated from tropical or warm-temperate seas, *Labrenzia* species were more often found in algal samples isolated from temperate regions, and *Rickettsia* endophytes were only present in *Bryopsis* species inhabiting seas with a low mean sea surface temperature (11.7°C–12.8°C) and high chlorophyll, nitrate and phosphate levels (figures 2 and 3). These correlations suggest that the distribution of individual bacterial OTUs may be more predictable than the total bacterial community composition. Individual bacterial endophyte groups also appear to be more strongly correlated

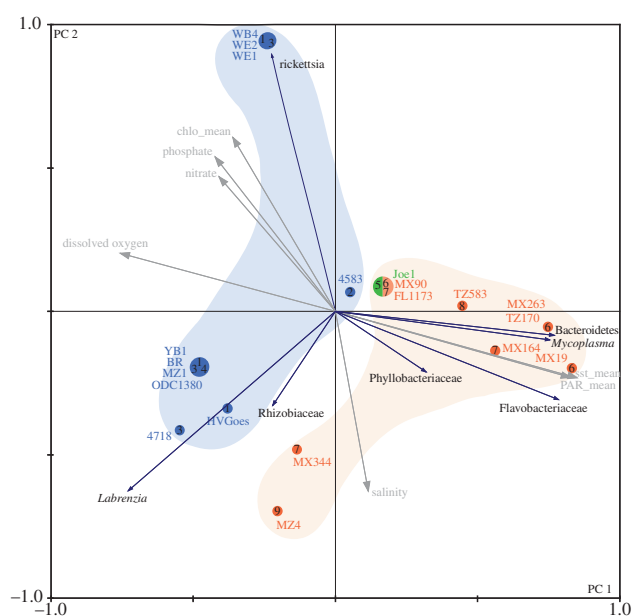


Figure 3. PCA of the 20 *Bryopsis* samples for which endophytic bacterial information is available. The PCA plot spreads the host samples in direction of maximum variance in endophytic bacterial community composition with principal component 1 explaining 41.7% and principal component 2 explaining 19.9% of the variance. *Bryopsis* species are indicated as numbers 1–9 and phylogenetic clades A, B and C are shown in blue, green and red, respectively. Environmental variables (in grey) were plotted on the PCA graph in the electronic supplementary material.

with the host phylogeny than the overall bacterial composition. Flavobacteriaceae and Bacteroidetes species displayed a significant phylogenetic signal ($p \leq 0.01$, see electronic supplementary material, table S2), while Rhizobiaceae, Phyllobacteriaceae, *Mycoplasma*, *Rickettsia* and *Labrenzia* species did not. Because the host phylogeny is correlated with ecological features as a consequence of niche conservatism (figure 1), it is not obvious whether the latter pattern is due to ecological preferences of the endophytic bacteria or their host.

(d) Host versus environmental influences

In order to disentangle the influences of different factors shaping the endophytic bacterial diversity, we performed variation partitioning analyses. In the first set of analyses, we partitioned the variation of the bacterial diversity data with respect to the ecological, geographical and host phylogenetic factors into different portions: a part strictly influenced by environmental variables, a part strictly influenced by the *Bryopsis* host phylogeny, a part strictly explained by geography, four parts explained by the shared influence of these three factors and an unexplained part of the variation. When considering the total endophytic bacterial diversity, approximately equal parts of the variation (ca 30%) were explained by environmental and phylogenetic factors, while the strict influence of geography was low; most of the variance, however, remained unexplained (figure 4). Analyses of the seven bacterial phylotypes separately showed that the influence of environment, phylogeny and geography was very different between the seven phylotypes. The influence of geography was, in most cases, low and highly correlated with environment and/or host phylogeny (see figure 4 and electronic supplementary material, table S4). For this reason, we excluded geography in a

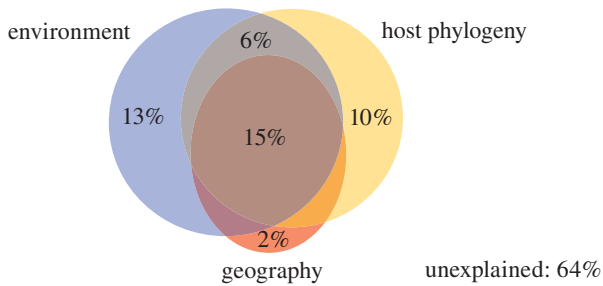


Figure 4. Partitioning the variation in total bacterial species composition explained by phylogenetic, environmental and geographical variables. Adjusted R^2 values are given. A detailed overview of the variation partitioning results is provided in the electronic supplementary material, table S4.

second set of analyses (figure 5). The independent effects of host phylogeny and environment had little influence on the presence of Phyllobacteriaceae, Rhizobiaceae and *Labrenzia* phylotypes. The shared influence of host phylogeny and environment was larger than their individual effects for these bacterial types. The occurrence of *Mycoplasma* and *Rickettsia* species, on the other hand, was in part strictly determined by environmental factors, whereas the distribution of Bacteroidetes could to a large extent be explained by host phylogenetic factors only. Most of the variance in presence of these six endophytic phylotypes, however, remained unexplained, suggesting that factors other than host phylogeny and environment (at least the seven variables sampled) determine their occurrence within particular *Bryopsis* samples (figure 4). This is in contrast with the situation for Flavobacteriaceae endophytes, whose presence could be almost entirely explained by host phylogenetic factors, which partly overlapped with environmental factors.

4. Discussion

Community structure and variation in traits across species are the outcome of environmental, geographical and historical factors which are clearly interwoven with each other. Bacterial communities associated with eukaryotic hosts are influenced by similar factors which need to be identified separately. Besides serving as baseline knowledge of the bacterial diversity occurring inside the siphonous cells of *Bryopsis*, our results provide insights into the various elements that contribute to the composition of the endogenous bacterial flora of siphonous green seaweeds.

(a) Diversity of endogenous bacteria

Characterization of the endobiotic microbial communities of 20 *Bryopsis* samples from widely divergent geographical localities indicates that *Bryopsis* harbours endophytic bacterial communities that are not very complex, but nevertheless taxonomically diverse. The constraint of living, unialgal samples as starting material necessitated extended cultivation of the algal samples prior to molecular processing [17]. These artificial culture conditions might have unknown effects on the endophytic bacterial community, potentially altering its complexity and this needs to be taken into account when interpreting these findings. Although they may not fully represent the total microbial variety present within the alga in its natural environment, the bacteria identified in this study are part of the natural *Bryopsis* endobiotic flora. Supplementary

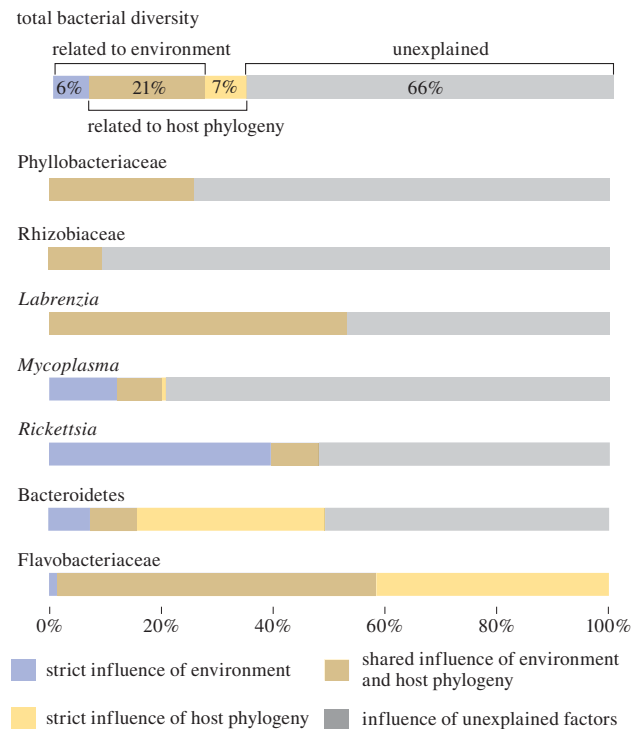


Figure 5. Partitioning the variation in bacterial species composition explained by phylogenetic and environmental variables. Diagrams show the unique and shared influences of both factors on the variation in total endophytic bacterial diversity and the individual endophytic phylotypes. Adjusted R^2 values are given or illustrated. Negative fractions (which indicate that two explanatory variables have strong and opposite effects on the dependant variable) are treated as zeros in the graphs. A detailed overview of the variation partitioning results is given in the electronic supplementary material, table S4.

time stability experiments [25] demonstrated that *Bryopsis* endophytic bacterial communities are rather stable and clearly distinct from the epiphytic and surrounding cultivation water bacterial communities.

Besides the five bacterial phylotypes that were previously characterized in *Bryopsis* (*Labrenzia*, *Mycoplasma*, Phyllobacteriaceae, Bacteroidetes and Flavobacteriaceae) [17], we identified two additional phylotypes related to Rhizobiaceae and *Rickettsia* species. These bacteria have been especially well studied from terrestrial habitats [45,46], but have also been reported from marine habitats. Rhizobiales are common epiphytes of *Ulva* seaweeds [47–50] and have also been isolated from the surface of kelps where they display antimicrobial activity [13]. Additionally, a *Rhodopseudomonas* species with the potential to fix nitrogen was isolated from the rhizoidal cytoplasm of the siphonous green seaweed *Caulerpa taxifolia* [19]. We presume that *Bryopsis* also hosts Rhizobiaceae species with nitrogen fixing capacities, as we were able to amplify *Ensifer*-like nitrogenase reductase genes (EMBL accession numbers HE649370–HE649371) from *Bryopsis* samples 4718 and MZ4 [51]. Obligate intracellular *Rickettsia* species, on the other hand, have not previously been described from macroalgae but have been characterized through 16S rRNA gene analysis within freshwater green microalgae [52], marine ciliates [53] and coral tissue [54].

(b) Factors affecting bacterial composition

Even though each bacterial phylotype was encountered in at least three *Bryopsis* samples, the total endophytic bacterial

diversity per host sample showed no clear pattern. All algal samples harboured diverse combinations of one to four endophytic phylotypes regardless of their phylogenetic affiliation, geographical origin or macro-ecological niche. On the other hand, when the presence of individual endophytic phylotypes rather than the total bacterial composition was analysed, host phylogenetic, geographical and environmental influences could be determined more clearly. These three factors, however, are inevitably interrelated as a result of phylogenetic niche conservatism, i.e. the tendency of closely related species to be ecologically similar [55], and historical factors such as dispersal limitation, resulting in geographical proximity of closely related species (figure 1). The *Bryopsis* host phylogeny was found to be mainly correlated with temperature-dependent variables and to a lesser extent with geography (see the electronic supplementary material, table S2). To disentangle the effects of host phylogeny, geography and environment on the endophytic bacterial community, we performed a variation partitioning analysis [2,56]. This technique has proved useful in quantifying independent influences of host phylogeny and other traits such as habitat and morphology in host–parasite [57–59] and arbuscular mycorrhizal symbiosis studies [60]. Our analyses shed light on the symbiotic nature and on potential modes of transmission of the individual endophytic phylotypes.

(c) Geography and environment on the endophytic bacterial community

The presence of endophytic Phyllobacteriaceae, Rhizobiaceae and *Labrenzia* phylotypes was not separately determined by host phylogenetic, geographical and ecological factors, suggesting that these endophytes are true generalists adapted to both free-living and host associated lifestyles along with a wide variety of environmental conditions. This is consistent with our previous observations that *Labrenzia* and Phyllobacteriaceae endophytes can survive outside their *Bryopsis* host and are reacquired from the local environment after repeated wounding events in culture [25]. Also the close phylogenetic relatedness of all three endophytic phylotypes with sequences from free-living bacteria (see the electronic supplementary material, figure S2) indicates a facultative association with the *Bryopsis* host. These generalist phylotypes may be selectively acquired by *Bryopsis* hosts to fulfil specific metabolic requirements, such as nitrogen-fixation (Rhizobiaceae; [45]), anoxygenic photosynthesis (Phyllobacteriaceae; [17]) or CO-oxidation (*Labrenzia*; [61]).

The occurrence of *Mycoplasma* and *Rickettsia* endophytes was to some extent strictly influenced by environmental factors. *Mycoplasma* endophytes were only present in *Bryopsis* samples from tropical regions, whereas *Rickettsia* bacteria were only found in algal samples isolated from temperate seas. This environmental influence suggests the acquisition of habitat-specific endophytes by *Bryopsis* hosts. In addition, the phylogenies of these more specialized endophytic phylotypes show a close relatedness with symbiotic *Rickettsia* and *Mycoplasma* species isolated from the cytoplasm of the marine ciliate *D. appendiculata* [53] and the intestinal bacterial flora of the *Bryopsis*-feeding abalone *Haliotis diversicolor* [62], respectively, suggesting the uptake of these endophytes could be vector dependant. This hypothesis is likely as both endophytes belong to the orders that are well known as obligate intracellular parasites of plants and animals [63,64].

Also within sponge hosts, horizontal symbiont transmission has been proposed to occur through vectors, including sponge-feeding animals [65].

The presence of Bacteroidetes endophytes within *Bryopsis* was to a large degree influenced by host phylogenetic factors, which may result from the fact that related algae have evolved similar traits that select for the uptake of these bacteria. Alternatively, the presence of Bacteroidetes endophytes in related hosts may be explained through vertical inheritance of bacteria either during sexual reproduction or asexual proliferation by fragmentation or extruded protoplasts that regenerate into new *Bryopsis* plants [66]. The absence of Bacteroidetes endophytes in the culture medium suggests that they may be obligate and hence vertically inherited endosymbionts [25], but further research is needed to exclude alternative hypotheses.

The presence of Flavobacteriaceae was found to be influenced by host phylogenetic factors in part combined with environmental influences, suggesting that these bacteria are specialized and obligate endosymbionts [67]. This is in line with results from culture experiments, which showed that these bacterial species are strictly dependant on the *Bryopsis* host for their growth and survival [25].

Overall, the variation partitioning analyses showed a high fraction of unexplained variation. This was true when considering the total endophytic bacterial diversity and the individual bacterial phylotypes, with the notable exception of the Flavobacteriaceae endophytes. This unexplained variation indicates that other factors may be important in explaining endophytic bacterial composition in *Bryopsis*. The variables included in this study are situated at the macro-ecological level and are considered suitable for explaining broad scale distribution patterns [33]. Ecological variables associated with microhabitat preferences, biotic interactions (e.g. bacterial transmission as a result of grazer-induced wounding) and physiological state of the host might also be important for interpreting fine scale bacterial community structure. In addition, endophytic community composition may be a result of stochastic processes. The few samples collected from one region and the same host species in this study indicate that a considerable variation in bacterial community composition may exist. Further studies, including many samples from a single host species at a single locality will be required to shed light on variation between co-occurring *Bryopsis* plants. However, to do so, technical difficulties inherent to this study (e.g. time consuming culturing and surface sterilization of host plants, and a large (greater than 95%) fraction of host chloroplast 16S rDNA in clone libraries) will need to be overcome, for example by applying species specific primers or high-throughput sequencing techniques. The unexplained variation in endophytic bacterial species composition is also relevant in the light of recent studies providing evidence that functional genes and transcripts, rather than species identified through rRNA taxonomy may be important in understanding bacterial community structure [68,69]. For example, bacterial communities associated with the green seaweed *Ulva* were found to be largely determined by function, rather than taxonomic identity [68]. It might thus be possible that functional genes rather than symbiont species or phylotypes are influenced by evolutionary (host phylogeny) and ecological factors [69].

In conclusion, characterization of *Bryopsis* algae sampled worldwide revealed the presence of variable endobiotic

communities of relatively few but taxonomically diverse bacterial species. Variation partitioning analyses show a strong impact of local environmental factors on the presence of some bacteria, while the presence of others reflects a predominant imprint of host evolutionary history. These observations, however, were only evident when subdividing the total endophytic diversity into its individual bacterial phylotypes, suggesting that both the whole community and

individual community members need to be considered in host–symbiont studies.

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