The Role of Photobionts as Drivers of Diversification in an Island Radiation of Lichen-Forming Fungi

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Speciation in oceanic islands has attracted the interest of scientists since the 19th century. One of the most striking evolutionary phenomena that can be studied in islands is adaptive radiation, that is, when a lineage gives rise to different species by means of ecological speciation. Some of the best-known examples of adaptive radiation are charismatic organisms like the Darwin finches of the Galapagos and the cichlid fishes of the great African lakes. In these and many other examples, a segregation of the trophic niche has been shown to be an important diversification driver. Radiations are known in other groups of organisms, such as lichen-forming fungi. However, very few studies have investigated their adaptive nature, and none have focused on the trophic niche. In this study, we explore the role of the trophic niche in a putative radiation of endemic species from the Macaronesian Region, the Ramalina decipiens group. The photobiont diversity was studied by Illumina MiSeq sequencing of the ITS2 region of 197 specimens spanning the phylogenetic breadth and geographic range of the group. A total of 66 amplicon sequence variants belonging to the four main clades of the algal genus Trebouxia were found. Approximately half of the examined thalli showed algal coexistence, but in most of them, a single main photobiont amounted to more than 90% of the reads. However, there were no significant differences in photobiont identity and in the abundance of ITS2 reads across the species of the group. We conclude that a segregation of the trophic niche has not occurred in the R. decipiens radiation.

Keywords: Macaronesia, Ramalina, trophic niche, metabarcoding, adaptive radiation

INTRODUCTION

Oceanic islands, which are of volcanic origin and have never been connected to continental land masses, have attracted the interest of scientists since the 19th century, using them as natural laboratories in which to study the origin of biodiversity and its evolution (Whittaker and Fernández-Palacios, 2007; Warren et al., 2015). Perhaps the best-studied evolutionary phenomenon on oceanic islands has been that of cladogenetic speciation, that is, how from a colonization event a lineage radiates into a collection of species (Gould and Eldredge, 1977).
Radiations can be adaptive or non-adaptive (Gittenberger, 1991; Schluter, 2000; Rundell and Price, 2009). Non-adaptive radiations are those in which diversification is not accompanied by niche differentiation and usually gives rise to ecologically similar allopatriec species (Gittenberger, 1991; Rundell and Price, 2009). On the other hand, in adaptive radiations, ecological speciation is the driver of diversification, and niche differentiation among radiation-originating species allows sympatric patterns (Schluter, 2000). One key point in defining an adaptive radiation is that the traits presented by the species (phenotype) must have a clear correlation with the environment in which the species occur (Schluter, 2000; Givnish, 2015). Therefore, in the study of the adaptive nature of a radiation, it is essential to study the variability of functional traits in the species of the radiation and their correlation with abiotic factors. The best-studied examples of adaptive radiation have focused on charismatic organisms, such as the Darwin finches of the Galápagos Islands (Grant and Grant, 2008), the silversword alliance of Hawai’i (Landis et al., 2018), or the Anolis lizards of the Greater Antilles (Losos, 2009). However, the number of studies that have investigated the adaptive nature on radiations of other groups of less charismatic organisms is still very limited (Gaya et al., 2015; Mao et al., 2018).

Lichens are the symbiotic phenotype of fungi with a specialized type of nutrition that takes sugars from a population of extracellular algae and/or cyanobacteria (hereafter, the photobionts). Thus, for lichen-forming fungi (hereafter, the mycobionts), the identity and abundance of their associated photobionts constitute the trophic dimension of their ecological niche. Although more than 50 genera of algae and cyanobacteria are known to act as lichen photobionts (Friedl and Büdel, 2008; Sanders and Masumoto, 2021), the vast majority of the nearly 20,000 described species of lichen-forming fungi (Lücking et al., 2017) are associated with algae from the class Trebouxiphyceae (Friedl and Büdel, 2008). Among the Trebouxiphyceae, the species of the genus Trebouxia are the most common photobionts (Friedl and Büdel, 2008; Muggia et al., 2020). The role of photobionts as drivers of speciation in lichen-forming fungi is still poorly known. There are several factors that may determine the contribution of photobionts in mycobiont diversification (Singh et al., 2017). On the one hand, it has been proposed that photobiont niches are influenced by abiotic conditions and these preferences may limit lichen distributions (Peksa and Škaloud, 2011). It has been shown that photobiont switches between physiologically compatible photobionts are mechanisms to widen the mycobiont ecological niches and geographic ranges (Fernández-Mendoza et al., 2011; del Campo et al., 2013; Rolshausen et al., 2018, 2020) or for ecological speciation (Ortiz-Alvarez et al., 2015). On the other hand, not all mycobiont species show similar specialization toward their photobionts (Pérez-Ortega et al., 2012; Magain et al., 2017), ranging from narrow specialists (Garrido-Benavent et al., 2017) to generalists (Muggia et al., 2014). In addition, different Trebouxia species have been found to coexist inside a single lichen thallus (Piercey-Normore, 2006; del Campo et al., 2010; Muggia et al., 2013). These species show differences in various aspects of their physiology, such as growth rate (Casano et al., 2011), photosynthetic output (Casano et al., 2011), response to oxidative stress (del Hoyo et al., 2011), and heavy metal tolerance (Álvarez et al., 2015). It is still not clear whether coexistence is a widespread phenomenon or it is reduced to certain species or lineages, and its significance in ecological and evolutionary times.

Ramalina is a well-known genus of lichen-forming fungi which forms usually large fruticose thalli of a more or less light-yellow color. The genus belongs to a clade of fruticose genera in the Ramalinaceae (Spjut et al., 2020), a family dominated by microlichens (Kistenich et al., 2018). The genus is subcosmopolitan and with c. 230 species is one of the largest genera of lichen-forming fungi (Lücking et al., 2017). Most of its diversity is concentrated in five areas: The Andes (Marcano et al., 2021), East Africa (Krog and Swinscow, 1976), Baja California (Bowler and Rundel, 1972, 1973), Australasia (Stevens, 1987; Blanchon et al., 1996) and Macaronesia (Krog and Østhagen, 1980a,b; Krog, 1990). Macaronesia is a group of volcanic archipelagos in the North Atlantic: Azores, Cape Verde, Canary Islands, Madeira, and Selvagens Islands. In spite of the small area, a total of 37 species of Ramalina have been recorded from the area, of which 25 are endemic (68%, most outstanding being the island of Porto Santo which, with an area of 42 km², hosts a total of 6 endemic Ramalina species (Sparrius et al., 2017). Among the endemic species of Ramalina in Macaronesia two groups of saxicolous species stand out, the R. bourgaeana group and the R. decipiens group, which are defined by the presence of characteristic structures, chondroid strands embedded in the medulla in the former and adjacent to the cortex in the later (Krog and Østhagen, 1980a; Spjut et al., 2020). The taxonomy of the Ramalina decipiens group has been recently revisited through an integrative approach recognizing 15 species-level lineages, six of which are newly described and two remain formally undescribed (Blázquez et al., in prep.). Of the fifteen recognized species, all are endemic to Macaronesia, except R. maderensis, which has also been reported from Saint Helena (Aptroot, 2008). In addition, most of them are single-island endemics, but two species (R. decipiens and R. maderensis) are widespread.

In this study, we hypothesize that photobiont-mycobiont associations play an important role in adaptive radiations of lichen-forming fungi. Photobiont differences among the species of a radiating clade of mycobionts, either in identity or abundance, could indicate a segregation of the trophic niche of the clade. We propose the Ramalina decipiens group as a framework to study the role of trophic niche segregation as a key driver in adaptive radiation. In order to explore and test this hypothesis, we are to analyze (i) the diversity of the photobionts associated with the species of the R. decipiens group, (ii) the relative importance of mycobiont identity, island, and macroclimate on photobiont community structure, (iii) the level of specialization of each Ramalina species toward its photobionts and (iv) whether specialization and photobiont diversity are phylogenetically structured. In addition, as algal coexistence has been previously reported in Ramalina species, we also intend (v) to explore to what extent this phenomenon occurs in the group.
**MATERIALS AND METHODS**

**Sampling**
We studied 197 specimens of the *R. decipiens* group representing the 15 species-level lineages (Blázquez et al. in preparation). Seven of those species correspond to previously known species (Krog and Østhagen, 1980a,b; Krog, 1990). Another six represent new species which are going to be described in Blázquez et al. (in preparation). They are referred here with the following provisional names: *R. delicata* nom. Prov., *R. fortunata* nom. Prov., *R. gomerana* nom. Prov., *R. papryacea* nom. Prov., *R. sabinosae* nom. Prov., and *R. sampaiana* nom. Prov. Two further species-level lineages will remain unnamed until further material is available and are referred in the present study as *Ramalina* sp. 1 and *Ramalina* sp. 2. The specimens were collected throughout 66 localities spanning the seven Canary Islands, the islands of Madeira and Porto Santo of the Madeira archipelago, and the islands of Sal and São Vicente of the Cape Verde archipelago (Supplementary Table 1).

**DNA Extraction, Amplification, and Sequencing**
A small thallus fragment from each specimen (c. 10 mm²) was excised from the tip of a single lacinia with the help of razor blade and forceps under a Nikon SMZ800 stereomicroscope. First, we inspected the thalli under a x80 magnification to check for areas free of epiphytic microalgae and fungi. Thallus tips were the only zones free of epiphytes in all samples. The choice of thallus tips is also supported by the finding of Molins et al. (2021), who reported that there were no differences in photobiont composition among different parts of the thallus in specimens of a species of the same genus (*R. farinacea*) from the same biogeographic region as the present study. Samples were washed with acetone to remove secondary metabolites and stored at −80°C. After 1 h of freezing, they were pulverized using TissueLyser II (Qiagen) with two crystal beads. Genomic DNA was extracted using E.Z.N.A.® Forensic DNA Kit (Omega Bio-Tek), following the instructions of the manufacturer. The second region of the ribosomal internal transcriber spacer (ITS2) was used as barcode to prospect photobiont diversity. We used primers FGITS2-f and FGITS2-r (Dal Grande et al., 2018) for amplification. These primers included Fluidigm CS1 and CS2 universal oligomer sequences at their 5’ ends. PCR reactions were carried out in a total volume of 15 μl, containing 3 μl of template DNA, 0.3 μl of each primer (10 μM), 0.6 μl of MyFi DNA Polymerase (Bioline, Sydney, Australia), 3 μl of 5x MyFi Reaction Buffer, and 7.8 μl of distilled water. PCR settings consisted in an initial denaturation at 95°C for 1 min; 35 cycles of 95°C for 15 s, 54°C for 15 s, and 72°C for 15 s; with a final extension at 72°C for 5 min. PCR products were checked in 1% agarose gels stained with SYBR™ Safe DNA Gel Stain (Thermo Fisher Scientific). PCR products were quantified using the Qubit dsDNA HS (High Sensitivity) Assay Kit (Thermo Fisher Scientific) and pooled in equimolar concentrations for sequencing on a single MiSeq run (Illumina, United States) at the RTSF Genomics Core at Michigan State University (East Lansing, Michigan). The Genomics Core completed library preparation by secondary PCR targeting the Fluidigm CS1/CS2 oligos at the ends of the primary PCR products. Primers used for the second PCR included dual indexed, Illumina library compatible sequences. Finished libraries were bulk normalized using Invitrogen SequaPRep DNA Normalization Plates (Thermo Fisher Scientific) and the recovered products pooled. The pool was QC’d and quantified using a combination of Qubit dsDNA HS (Thermo Fisher Scientific), Agilent 4,200 TapeStation HS DNA1000 (Agilent Technologies), and Kapa Illumina Library Quantification qPCR assays (Kapa Biosystems, Inc.). This pool was loaded onto an Illumina MiSeq v2 Standard flow cell and sequencing performed in a 2x250bp paired-end format using a MiSeq v2 500 cycle reagent cartridge. Custom sequencing and index primers complementary to the Fluidigm CS1/CS2 oligos were added to appropriate wells of the reagent cartridge. Base calling was done by Illumina Real-Time Analysis (RTA) v1.18.54, and output of RTA was demultiplexed and converted to FastQ format with Illumina Bcl2fastq v2.19.1.

**Sequence Processing**
Raw sequence data were processed using the DADA2 pipeline (Callahan et al., 2016) in R 4.0.3 (R Core Team, 2020). DADA2 takes a set of demultiplexed paired-end fastq files, filters the sequences based on their length and quality, assembles them into error-corrected amplicon sequence variants (ASVs), and removes chimeric ASVs. We chose this pipeline because ASVs make possible to work at a finer resolution than other approaches based on the creation of operational taxonomic units (OTUs). Besides, ASVs allow for greater reproducibility, reusability, and comprehensiveness than OTUs (Callahan et al., 2017). Following Dal Grande et al. (2018) we considered that ASVs that represented <0.005% (i.e., n = 100) of the total reads were most probably exogenous to the symbiosis and did not include them in downstream analyses. We then assigned taxonomy to the ASVs using all sequences assembled by Muggia et al. (2020) as a taxonomic reference. This was done using the assignTaxonomy function in DADA2. Given that microalgae can show ecophysiological differences even when they are phylogenetically close (Sadowsky et al., 2016) we chose to conduct all analyses in this study at the ASV level. All sequences obtained in this study are available in the SRA (NCBI) under BioProject PRJNA764073.

**Alignment, Phylogenetic Relationships, and ASV Haplotype Networks**
ASVs were aligned using MAFFT v7.308 (Katoh et al., 2002) as implemented in Geneious® v.9.1.8. Sequences of *Asterochloris glomerata* (MW043487, Pino-Bodas and Stenroos, 2021) and *Vulcanochloris guanchorum* (KR952330, Vančurová et al., 2015) were included as outgroup. We inferred phylogenetic relationships among ASVs using Maximum Likelihood using RAxMLHPC2 v. 8.2.4 (Stamatakis, 2014) as implemented in CIPRES Science Gateway (Miller et al., 2011). We selected the GTR+GAMMA substitution model and carried out 1,000 rapid bootstrap
pseudoreplicates to evaluate nodal support. We considered nodes with bootstrap values equal or higher than 70% to be significantly supported. The resulting phylogenetic tree was visualized in FigTree 1.4.4 and edited using Adobe Illustrator CS5 (Adobe Systems Inc., San José, United States). Additionally, we generated haplotype networks for the Trebouxia species represented by more than three ASVs. We used the TCS method (Templeton et al., 1992) as implemented in PopART v.1.7 (Leigh and Bryant, 2015) for network construction. These networks were edited in Adobe Illustrator CS5 to show the prevalence of the different recovered ASV in each sampled Ramalina species and island.

**Differences in Frequency of ASVs in Islands and Species**

To explore whether the frequency of ASVs was different among islands and Ramalina species we adjusted a two-way ANOVA for each ASV. ASV frequency was used as the response variable and island and Ramalina species as explanatory variables. We used Levene’s test to check whether data for each ASV met the normality assumption necessary to carry out a two-way ANOVA. In the cases in which our data did not meet the assumptions, we employed the non-parametric Kruskal-Wallis test. This was done using the functions aov, leveneTest, and kruskal.test in base R 4.0.3 (R Core Team, 2020). ASV’s occurring only in one specimen were not included in this analysis.

**Photobiont Community Structure**

In order to explore the relative importance of mycobiont identity, macroclimate, and island in shaping photobiont communities we performed distance-based redundancy analysis (dbRDA, Legendre and Anderson, 1999). Following Shankar et al. (2017), we calculated an abundance-weighed phylogenetic UniFrac distance-based dissimilarity matrix. This was done with the R package phyloseq (McMurdie and Holmes, 2013). We transformed the ASV table into relative ASV abundances using the function decostand of the vegan package (Oksanen et al., 2012) with the method “total” and combined it with the taxonomy of the ASVs and the RAXML phylogeny previously generated into a phyloseq object using the merge_phyloseq function. The dissimilarity matrix was calculated from this object using the UniFrac function. The weighed UniFrac dissimilarity matrix was used as the response matrix in the dbRDA. We used three explanatory matrices: (1) first matrix corresponded to the Ramalina species, (2) second to the island where samples were collected, (3) and third matrix included five macroclimatic variables downloaded from WorldClim (Fick and Hijmans, 2017): average temperature (°C), precipitation (mm), solar radiation (kJ m⁻² day⁻¹), wind speed (ms⁻¹), and water vapor pressure (kPa). Because lichens are photosynthetic and poikilohydric symbioses, we chose to use these instead of the 19 widely used bioclimatic variables also available in WorldClim, that are derived from temperature and rainfall only. All layers of the five variables (data for each month of the year are available) were downloaded at a resolution of 30 s (~1 km²) and merged into year averages using the calc function of the raster R package (Hijmans et al., 2015). Then, variable values for each sample were obtained through their geographic coordinates using the extract function from the raster package. Before computing the dbRDA we checked for correlation between WorldClim variables in our samples. We calculated a Pearson correlation coefficient matrix and generated a correlation hierarchical cluster plot with the absolute correlation values. Precipitation and water vapor pressure were dropped as they were correlated (absolute correlation >0.5) with solar radiation and average temperature, respectively. Then we inspected the remaining variables to check their normality. To do so we generated a histogram for each variable with the hist function of base R. Solar radiation was found to be not normal and was log-transformed. dbRDA was then calculated with the dbRDA function of the vegan R package (Oksanen et al., 2012). Variance partitioning between the three explanatory matrices was estimated using adjusted R². Adjusted R² significance was assessed for each fraction with a permutation-based ANOVA test with 2000 permutations. This was done in vegan using the functions varpart and anova.cca. Significance of the dbRDA as a whole was also assessed with a permutation-based ANOVA test with 2000 permutations, this time with the function anova. To visualize variance partitioning we generated a Venn diagram plotting the output of the varpart function.

**Algal Co-occurrence Analyses**

Based on the ASVs abundance table we calculated for each Ramalina species: number of studied thalli, percentage of thalli that showed algal coexistence, minimum number of ASVs in one thallus, mean number of ASVs per thallus, and maximum number of ASVs in one thallus. For the species that showed algal coexistence, we also calculated: minimum relative abundance of the main photobiont found in one thallus, mean relative abundance of the main photobiont across all thalli showing algal coexistence and maximum relative abundance of the main photobiont found in one thallus. 

**Bipartite Network, Specialization, and Modularity**

We built two different adjacent matrices i × j, where i represents mycobionts and j ASVs. In the first matrix, interaction frequency between mycobiont i and ASV j was calculated as the sum of reads for ASV j found in all specimens of mycobiont i previously rarefied to the depth of the sample with the lower number of reads (Weiss et al., 2017). In the second matrix, number of read are not taken into account and the interaction frequency was based in the number of specimens of the mycobiont i where ASV j was found. The rarefaction was done with the rarefy_even_depth function of the phyloseq (McMurdie and Holmes, 2013) R package. The presence-absence conversion was done using the function decostand of the vegan package (Oksanen et al., 2012) with the method “pa.” For each network, we calculated the parameter d’ (Blüthgen et al., 2006) to estimate the specialization of the mycobionts toward
the ASVs. This index ranges from 0 (no specialization) to 1 (high specialization). We also calculated species degree, which is the sum of links per species, and ND, which is species degree normalized by the number of possible partners. All analyses were carried out in the R package bipartite (Dormann et al., 2008), using the functions plotweb and specieslevel. Following the approach of Gómez et al. (2015), we used the quantitative modularity (Q) of the bipartite networks to explore the occurrence of different photobiont niches in the species of the Ramalina decipiens group. Optimal modular configuration was calculated using the Beckett algorithm (Beckett, 2016) as implemented in the computeModules function of the bipartite package. The algorithm was run 20 times with 10⁶ MCMC steps and a tolerance level of 10⁻⁶ retaining the iteration with the highest likelihood value as the optimal modular configuration.

In order to check for significance, we built a null model based on 500 random networks calculated using the function nullmodel and the vaznull null model which implements the null model proposed by Vázquez et al. (2007) in which matrix connectance remains the same as in the original network but total marginals change. We calculated modularity for each of the random matrices and calculated a z-score as \( Z = \left( \frac{X_{\text{observed}} - \mu_{\text{null}}}{\sigma_{\text{null}}} \right) \) where \( X_{\text{observed}} \) is the actual value of the parameter, \( \mu_{\text{null}} \) is the mean of the parameter for the population of matrices in the null model, and \( \sigma_{\text{null}} \) their standard deviation. p-values for the z-score were calculated as the number of elements of the null model showing higher or lower values than the observed value divided by the total number of elements in the null model. To explore if the resulting modules were a consequence of the species geographic distribution we compared the geographic distances within and between modules. This was done with PERMANOVA tests using the adonis function of the vegan R package (Oksanen et al., 2012). The geographic distance matrix was obtained with the distm function of the geosphere R package (Hijmans et al., 2017).

Genetic Diversity Statistics
We calculated genetic diversity statistics at both the island and Ramalina species levels. We used the software DnaSP v6.12.01 to calculate two indices: haplotype diversity (HD, Nei, 1987), and nucleotide diversity (\( \pi \), Nei, 1987). \( \pi \) was calculated using the Jukes and Cantor correction (Lynch and Crease, 1990). Instead of using the raw ASVs abundance table as input, we generated DNA alignments for each species and each island, which included a representative sequence of each ASV present in each sample. The two indices were then calculated for each alignment.

Phylogenetic Signal
We used the phylogeny of the Ramalina decipiens group which will be published in a forthcoming paper (Blázquez et al. unpublished) to calculate the phylogenetic signal of the specialization, species degree, ND, haplotype diversity and nucleotide diversity. We employed two different methods to calculate the phylogenetic signal: Pagel's \( \lambda \) (Pagel, 1999) as implemented in the phylosig function of the phytools R package (Revell, 2012) and Blomberg's \( K \) (Blomberg et al., 2003) as implemented in the phylosignal function of the picante R package (Kembel et al., 2010). Pagel's \( \lambda \) can have values between 0 (no phylogenetic signal) and 1 (Brownian motion model, that is, correlation among trait values is proportional to the extent of shared ancestry). Blomberg's \( K \) can be interpreted as follows: if \( K = 1 \) the trait under analysis follows a mode of evolution that is consistent with Brownian motion. If \( K > 1 \) close relatives are more similar than expected under Brownian motion. On the other hand, in \( K \) values <1 close relatives are less similar than expected (Blomberg et al., 2003).
Photobiont Community Structure

The dbRDA analysis returned a $F=3.265$ with a $p=0.001$. The constrained axes explained 0.34 of the variance in ASV community structure (Figure 3). Island was the variable with the greatest simple effect. On the other hand, macroclimate was the variable with the smallest simple effect. However, the main driver of ASV community structure was the mixed effect of island and macroclimate, explaining a greater variance proportion than all other simple and mixed effects combined. Mycobiont identity was the least important variable, taking its simple and mixed effects into account.

Co-occurrence Analyses and Frequency of ASVs

Out of 197 thalli studied, 102 showed coexistence of several ASVs (51.77%). Specimens from the Canary Islands and Cape Verde showed a higher proportion of multiple ASVs within the thallus (56 and 58%, respectively) than specimens from the Madeira archipelago (38%). Algal coexistence was observed in all species (Table 2) except Ramalina sp. 1, which is most likely an artifact given that the species is represented in our dataset by only one specimen. The mean number of ASVs per thallus was 2.19, with a median of 2. Mean ASV count per thallus ranged from 1.30 in Ramalina hamulosa to 4.67 in Ramalina pluviariae. The sample with the highest ASV count was a Ramalina maderensis collected in Lanzarote (SPO 6337) containing 10 algae. However, on average the main photobiont amounted to 94% of the reads in thalli showing algal coexistence. The mean abundance of the main photobiont in thalli showing coexistence ranged from 83% in Ramalina erosa to 99% in Ramalina sp. 2. There were only six thalli, of varied origin and representing four Ramalina species, in which the main photobiont accounted for less than 70% of the reads. In most of them, the main and secondary photobionts were different ASVs belonging to the same Trebouxia species.
Bipartite Network, Specialization, and Modularity

The bipartite interaction networks (Figure 4; Supplementary Figure 1) clearly depict ASVs 1 and 2 as the main photobionts of the *Ramalina decipiens* group. *Ramalina* sp. 1 is the only species not associated with these ASVs. This is most probably due to the poor representation of this species in the dataset. Excluding *Ramalina* sp. 1, whose only thallus associates with another ASV belonging to *Trebouxia* sp. TR9, specialization (d') based on rarefied reads varied from 0.98 in *Ramalina delicata* to 0.51 in *Ramalina maderensis*. Species degree varied from 37 in *Ramalina decipiens* to 4 in *Ramalina erosa* and ND varied from 0.56 in *Ramalina decipiens* to 0.06 in *Ramalina erosa*. Species degree and ND were highest in the two most sampled species (*R. decipiens* and *R. maderensis*). Specialization parameters obtained from the data non considering read counts returned similar results for species degree and ND, but overall lower values of d', that ranged from 0.59 in *R. erosa* to 0.27 in *R. decipiens*. All specialization parameters are summarized in Table 3. The network based on rarefied reads was significantly modular (Q=0.18, z-score=24.75, value of p <0.001), and detected the presence of six modules (Figure 5). The modularity analyses of the second network resulted in a non-modular pattern (Q=0.21, z-score=-0.68, value of p =0.241). The modular structure recovered in the first analysis seems to be a consequence of the mycobiont species distributions inasmuch as the geographic distances between and within modules were significantly different (F=7.98, R² =0.17, p <0.001).

Genetic Diversity Statistics

Nucleotide diversity (π) and haplotype diversity (Hd) were calculated at the island and species levels (Figure 6). At the island level, π ranged from 0.036 in Tenerife to 0.096 in Sal. Hd values ranged from 0.678 in Porto Santo to 0.928 in Fuerteventura. On average, Cape Verde had the highest π value (0.092) followed by Madeira (0.060) and the Canary Islands (0.052). Regarding Hd, Cape Verde again showed the highest value (0.810), followed by the Canary Islands (0.803) and Madeira (0.730). At the species level, π values ranged from 0.014 in *Ramalina hamulosa* to 0.077 in *Ramalina papyracea*. Hd values ranged from 0.709 in *Ramalina erosa* to 0.905 in *Ramalina pluviariae*.

Phylogenetic Signal

The parameters d', species degree, and ND derived from both networks were not phylogenetically structured according to Pagel’s λ and Blomberg’s K. Regarding genetic diversity, neither π nor Hd were phylogenetically structured either. Phylogenetic signal statistics are summarized in Table 4.

DISCUSSION

We explored the role of the photosynthetic partners in the radiation of a clade of symbiotic organisms. According to our knowledge, this is the first time that the trophic niche is analyzed within the evolutionary framework of an island radiation in...
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| TABLE 1 | p-values of two-way ANOVA and Kruskal-Wallis test exploring the effect of island and/or mycobiont species on the frequency of each ASV. |
|---|---|---|---|
| ASV | P species | P island | P species/island |
| ASV 1 | 0.8864 | 0.0028*** | 0.4886 |
| ASV 2 | 0.2556 | 0.4255 | 0.2016 |
| ASV 3 | 0.5121 | 0.0803 | 0.7751 |
| ASV 4 | 0.0831 | 0.9234 | 0.9996 |
| ASV 5 | 0.5794 | 4.73E-14*** | - |
| ASV 6 | 0.9861 | 0.1584 | 0.9857 |
| ASV 7 | 0.6763 | 6.17E-12*** | - |
| ASV 8 | 0.5544 | 0.8430 | 0.6702 |
| ASV 9 | 0.1188 | 0.0007*** | 2.85E-12*** |
| ASV 10 | 0.9892 | 0.4981 | 0.6184 |
| ASV 11 | 0.9162 | 6.26E-08*** | - |
| ASV 12 | 0.9886 | 0.9360 | 0.9964 |
| ASV 13 | 0.5398 | 0.0879 | 0.9068 |
| ASV 21 | 1.25E-03*** | 0.0011** | - |
| ASV 23 | 0.8087 | 0.1121 | 0.9435 |
| ASV 28 | 0.1180 | 0.0988 | 0.9997 |
| ASV 29 | 0.9410 | 0.1843 | 0.9864 |
| ASV 35 | 0.9788 | 0.3024 | 0.9924 |
| ASV 36 | 0.0125*** | 0.6126 | 0.2036 |
| ASV 38 | 0.7826 | 0.2846 | 0.2359 |
| ASV 42 | 1.26E-02*** | 0.8119 | |
| ASV 43 | 0.0909 | 0.0006*** | - |
| ASV 47 | 1.0000 | 0.4689 | 0.9905 |
| ASV 49 | 0.9996 | 0.0105** | 0.9507 |
| ASV 54 | 0.9831 | 0.2656 | 0.5889 |

ASVs lacking the island-species interaction are those who did not meet the assumptions necessary to perform a two-way ANOVA. Bold values are statistically significant. *< 0.05; **< 0.01; ***< 0.001.

lichen-forming fungi. We found no differences in the identity of the photobionts associated with the species of the Ramalina group. The vast majority of specimens analyzed contained as their main photobiont one of the two most abundant ASVs, both belonging to Trebouxia sp. TR9. The frequency of both ASVs does not differ among Ramalina species, and we have indistinctly found both acting as major photobionts in thalli collected from the same locality on multiple occasions. The prevalence of T. sp. TR9 as main photobiont is not exclusive of the Ramalina group. It was found in Macaronesian specimens of Ramalina farinacea using Sanger sequencing (del Campo et al., 2013) and NGS (Molins et al., 2021). BLAST searches of the two predominant ASVs revealed their presence also in other genera. Both appear in Protoparmelia montagnei thalli in southern Spain (Singh et al., 2017) while ASV2 additionally appears in Lecanographa amylacea thalli in Poland and Sweden (Ertz et al., 2018). The fact that T. sp. TR9 is the main photobiont of Ramalina species in Macaronesia is congruent with our current knowledge of its physiology. Casano et al. (2011) showed that T. sp. TR9 grew faster and had better photosynthesis performance in in vitro experiments under relatively high temperatures and irradiances than T. jamesii, the main photobiont of R. farinacea thalli in the Iberian Peninsula (del Campo et al., 2013; Molins et al., 2021). The presence of both ASVs in southern Spain (Singh et al., 2017) also seems to point to the fact that this species can cope with higher temperature and light intensity given the similarity of that locality to those in which the Ramalina group species occur in Macaronesia. Surprisingly, the records from Poland and Sweden occurred in localities with radically different climatic conditions (Ertz et al., 2018). Lichen photobionts can show different physiologies even when they are phylogenetically very close (Sadowsky et al., 2016). The presence of ASV2, but not ASV1, in climatically divergent localities, such as the Białowieża Primeval forest in Poland or the island of Gotland in Sweden (Ertz et al., 2018), might suggest that they have differences in ecophysiological adaptations. Their presence in thalli collected in the same locality could be reflecting climatic differences at the microsite scale, as we have detected differences up to 4°C in two rocky outcrops separated by only 200 m (data not shown). The existence of such physiologcal differences will be explored in a forthcoming study.

We did not find differences in photobiont dominance between mycobiont species either. We detected algal co-occurrence in c. 50% of the studied thalli, a phenomenon reported in different lineages of lichen-forming fungi (Blaha et al., 2006; Piercey-Normore, 2006; del Campo et al., 2010; Muggia et al., 2014; Rolshausen et al., 2018). Casano et al. (2011) showed that the two Trebouxia species detected in R. farinacea apparently displayed complementary physiological behaviors, arguing that algal co-occurrence in this species could have allowed it to expand its geographical range. However, most photobiont co-occurrence reports have been based on Sanger sequencing, so the relative abundance of each photobiont could not be assessed. Recently, Molins et al. (2021) revisited photobiont diversity in R. farinacea with NGS data showing that in c. 70% of all thalli one Trebouxia species was dominant. Paul et al. (2018) reported similar results for Umbilicaria hispanica and U. pustulata, with the main photobiont being represented by at least 80% of the reads in c. 80% of all thalli. In our study, this proportion is even more pronounced, with the main photobiont being dominant in 94%
of the thalli showing algal co-occurrence. The overwhelming dominance of the main photobiont over the rest sheds doubt on the ecological importance of algal co-occurrence. Photobiont co-occurrence has been studied more in-depth in coral-dinoflagellate symbioses. Young corals establish symbiosis with a subset of the photobionts available in their environment (Poland et al., 2013; Yamashita et al., 2013). This first photobiont community is restructured in response to environmental heterogeneity (Rowan, 2004) or other processes, such as re-establishment of photobiont communities after a previous stress-induced loss (Jones et al., 2008). After the restructuration process, adult corals usually end up with a single Symbiodiniaceae lineage dominating the photobiont community (Parkinson and Baums, 2014). This agrees with the results shown by Molins et al. (2021) for Ramalina farinacea in which young and old thalli showed different microalgal communities, more diverse in young thalli.

The community structure of the photobionts associated to species of the R. decipiens group is chiefly explained by the

| Species            | N° thalli | % thalli >1 ASV | Mean ASV count/thallus | Maximum n° ASV/thallus | Minimum % main photobiont | Mean % main photobiont | Max % main photobiont |
|--------------------|-----------|-----------------|------------------------|------------------------|---------------------------|------------------------|-----------------------|
| Ramalina decipiens | 57        | 60%             | 2.46                   | 9                      | 42.93%                    | 93.85%                 | 99.94%                |
| Ramalina delicata  | 3         | 33%             | 2.33                   | 5                      | 95.89%                    | 95.89%                 | 95.89%                |
| Ramalina erosa     | 9         | 22%             | 1.22                   | 2                      | 67.04%                    | 83.50%                 | 99.96%                |
| Ramalina fortunata | 8         | 38%             | 1.38                   | 2                      | 99.59%                    | 99.81%                 | 99.95%                |
| Ramalina gomerana  | 3         | 33%             | 3                      | 7                      | 85.78%                    | 85.78%                 | 85.78%                |
| Ramalina hamulosa  | 10        | 30%             | 1.3                    | 2                      | 95.84%                    | 97.99%                 | 99.81%                |
| Ramalina madrensis | 57        | 53%             | 2.23                   | 10                     | 54.28%                    | 93.74%                 | 99.98%                |
| Ramalina nematodes | 6         | 50%             | 2                      | 5                      | 90.28%                    | 96.63%                 | 99.87%                |
| Ramalina papyracea | 6         | 50%             | 2                      | 4                      | 96.19%                    | 98.33%                 | 99.63%                |
| Ramalina pluviariae| 6         | 100%            | 4.87                   | 7                      | 83.74%                    | 93.73%                 | 99.79%                |
| Ramalina portosantana | 8   | 50%             | 2                      | 5                      | 88.48%                    | 94.70%                 | 99.95%                |
| Ramalina sabinosae | 4         | 100%            | 2.25                   | 3                      | 80.00%                    | 94.22%                 | 99.75%                |
| Ramalina sampaiana| 13        | 38%             | 1.92                   | 6                      | 69.96%                    | 90.89%                 | 98.60%                |
| Ramalina sp. 1    | 1         | 0%              | 1                      | 1                      | -                         | -                      | -                     |
| Ramalina sp. 2    | 6         | 50%             | 1.5                    | 2                      | 99.85%                    | 99.89%                 | 99.96%                |

*Number of studied thalli, percentage of thalli that showed algal coexistence, minimum, mean, and maximum number of ASVs per thallus and minimum, mean, and maximum relative abundance of the main photobiont across thalli showing algal coexistence.*
interaction between island and macroclimate, mycobiont identity being only of marginal importance. This finding agrees with previous studies that have reported macroclimate as a major driver of genetic variation of photobionts (Beiggi and Piercey-Normore, 2007; Nelsen and Gargas, 2009; Fernández-Mendoza et al., 2011; Leavitt et al., 2016; Magain et al., 2017). Other studies, however, have suggested a greater importance of mycobiont identity (Buckley et al., 2014; Leavitt et al., 2015; Vančurová et al., 2018; Jüriado et al., 2019). The relative importance of both factors can be influenced by the scale of the study. For instance, Pino-Bodas and Stenroos (2021) studied the diversity of the photobionts associated with the genus Cladonia by Sanger sequencing of the ITS and actin type I genes and found that the relative importance of the different factors differed between the two genes. However, when the ITS dataset was trimmed to include only the specimens for which actin type I sequences were available (losing mycobiont species in the process) the results for both datasets were similar (Pino-Bodas and Stenroos, 2021). The importance of the island-macroclimate interaction in our study is congruent with the climatic heterogeneity of Macaronesia, as the different islands show marked differences in the vegetation assemblages they harbor (del Arco Aguilar and Rodríguez Delgado, 2018) but at the same time, due to their pronounced orography, close areas can experience radically different climatic conditions (Sánchez-Benítez et al., 2017).

We found no phylogenetic signal in either the specialization parameter $d'$ or the genetic diversity indices tested, suggesting that species specialization toward their photobionts is not an evolutionarily conserved trait. Magain et al. (2017) studied

### TABLE 3 | Summary of specialization parameters of the bipartite networks based on the sum of the rarefied reads of the thalli of each species and the conversion of the abundance of ITS2 reads of the thalli to presence-absence data.

| Mycobiont species | Rarefied reads network | Presence-absence network |
|-------------------|------------------------|--------------------------|
|                   | $d'$  | Species degree | ND | $d'$ | Species degree | ND |
| *Ramalina decipiens* | 0.54 | 37 | 0.56 | 0.28 | 38 | 0.58 |
| *Ramalina delicata* | 0.98 | 5 | 0.08 | 0.46 | 5 | 0.08 |
| *Ramalina erosa* | 0.68 | 4 | 0.06 | 0.59 | 4 | 0.06 |
| *Ramalina fortunata* | 0.72 | 5 | 0.08 | 0.54 | 5 | 0.08 |
| *Ramalina gomerana* | 0.56 | 7 | 0.11 | 0.38 | 7 | 0.11 |
| *Ramalina hamulosa* | 0.70 | 6 | 0.09 | 0.53 | 6 | 0.09 |
| *Ramalina maderensis* | 0.51 | 33 | 0.50 | 0.29 | 33 | 0.50 |
| *Ramalina nematodes* | 0.51 | 9 | 0.14 | 0.53 | 9 | 0.14 |
| *Ramalina papyraceae* | 0.74 | 6 | 0.09 | 0.46 | 6 | 0.09 |
| *Ramalina pluviariae* | 0.54 | 12 | 0.18 | 0.31 | 12 | 0.18 |
| *Ramalina portosantana* | 0.70 | 8 | 0.12 | 0.41 | 8 | 0.12 |
| *Ramalina sabinosae* | 0.58 | 5 | 0.08 | 0.42 | 6 | 0.09 |
| *Ramalina sampaiiana* | 0.51 | 14 | 0.21 | 0.33 | 14 | 0.21 |
| *Ramalina sp. 1* | 0.62 | 6 | 0.09 | 0.44 | 6 | 0.09 |
| *Ramalina sp. 2* | 1.00 | 1 | 0.02 | 1.00 | 1 | 0.02 |

### FIGURE 5 | Modularity plot featuring the six modules identified by the Becket algorithm ($Q=0.179$, $z$-score=$709.31$; value of $p < 0.001$) in the rarefied reads network. Taxa are sorted according to their modular affinity, mycobiont species as rows, and photobiont ASVs as columns. Red boxes delineate the six modules. Color intensity is proportional to the relative abundance of each ASV per mycobiont species.
patterns of mycobiont specialization toward their cyanobionts in the *Peltigera* section *Polychaetosporina*. They found that generalist and specialist species were distributed all over the phylogeny proposing that photobiont specialization by lichen-forming fungi does not equal to an evolutionary dead-end as it has been proposed for other organisms (e.g., Haldane, 1951; Nosil and Mooers, 2005). Similar results of no phylogenetic conservatism in mycobiont specialization toward photobionts have been recently reported by Pino-Bodas and Stenroos (2021) for the genus *Cladonia*. Our results suggest that the *R. decipiens* group species associate with locally adapted photobionts, as do other lichen-forming fungi (Blaha et al., 2006; Muggia et al., 2014; Rolshausen et al., 2018). This is particularly clear in *R. maderensis*, the only species of the group occurring in the Cape Verde archipelago. In Cape Verde, *R. maderensis* has performed a photobiont switch and associates predominantly with ASVs belonging to *Trebutxia*.
C25. *Trebouxia* C25 has only been found previously in *Parmotrema aldabrense* thalli in Kenya, in a locality near the coast (Muggia et al., 2020). Nothing is known about this species physiology but the fact that it occurs mainly in tropical regions as well as all other clade C *Trebouxia* species in Muggia et al. (2020), could be indicative of a physiology adapted to higher temperatures and light intensities than those in which *T*. sp. TR9 thrives.

We have not observed differences neither in identity nor in the abundance of ITS2 reads in the photobionts associating with the different species of the *R. decipiens* group. Because differences in identity can be linked with differentiated growth rates and photosynthetic performances at the species (Casano et al., 2011) and strain (Determeyer-Wiedmann et al., 2019; Aigner et al., 2020) levels, we conclude that a partitioning of the trophic niche has probably not occurred during the *R. decipiens* group radiation. This fact, coupled with the marginal importance of mycobiont species on photobiont community structure and the geographic structure underlying the bipartite network modules, suggests that the photobionts are not a factor driving the *R. decipiens* group diversification. A segregation of the trophic niche is a key driver of adaptive radiation in many other examples, such as the Darwin finches (Grant and Grant, 2008), East African cichlid fishes (Danley and Kocher, 2001), *Tylomelania* freshwater gastropods (Rintelen et al., 2004), *Cyprinodon* pupfishes (Martin and Feinstein, 2014), *Tetragnatha* spiders (Kennedy et al., 2019), and many others (Gillespie et al., 2020). However trophic niche differentiation is not the only factor that can explain adaptive radiation. It has been hypothesized that Hawaiian lobeliads have diversified physiologically across environments differing in a key resource (i.e., light, Montgomery and Givnish, 2008). Recently it has been proposed (Gillespie et al., 2020) that symbiotic interactions with microbes could facilitate adaptive radiation. Physiological speciation in the *R. decipiens* group and a study of the species associated microbiomes will be explored in forthcoming works.

## CONCLUSION

We have found no evidence for trophic niche segregation in the context of the diversification of the Macaronesian endemic species of the *R. decipiens* group. They do not differ neither in the identity nor in the abundance of their photobionts. Thus, we argue that the photobionts are not a key factor driving speciation in the group. Instead, *Ramalina* species appear to associate with locally adapted *Trebouxia*. Algal coexistence was common, but the contribution of the secondary photobionts was marginal in most thalli. This opens new questions about the ecological importance of algal coexistence.

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## DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

## AUTHOR CONTRIBUTIONS

SP-O, FG, IP-V, and MB designed the study. MB, IP-V, and SP-O organized the fieldwork, research permits, and collected the specimens. MB and LH-M performed the laboratory work. MB analyzed the data and wrote the manuscript. All authors contributed to the article and approved the submitted version.

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## SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.784182/full#supplementary-material

**SUPPLEMENTARY FIGURE 1 |** Bipartite network based on presence-absence data showing interactions between *Ramalina* species and *Trebouxia* ASVs. Interactions with ASVs belonging to *Trebouxia* clades A, C, I, and S are depicted in green, orange, red and blue, respectively. Width of the links is proportional to the frequency of the association.
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