Changes in the foliar fungal community between oak leaf flushes along a latitudinal gradient in Europe

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Changes in the foliar fungal community between oak leaf flushes along a latitudinal gradient in Europe

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Abstract
Aim: Leaves support a large diversity of fungi, which are known to cause plant diseases, induce plant defences or influence leaf senescence and decomposition. To advance our understanding of how foliar fungal communities are structured and assembled, we assessed to what extent leaf flush and latitude can explain the within- and among-tree variation in foliar fungal communities.

Location: A latitudinal gradient spanning c. 20 degrees in latitude in Europe.

Taxa: The foliar fungal community associated with a foundation tree species, the pedunculate oak Quercus robur.

Methods: We examined the main and interactive effects of leaf flush and latitude on the foliar fungal community by sampling 20 populations of the pedunculate oak Quercus robur across the tree’s range. We used the ITS region as a target for characterization of fungal communities using DNA metabarcoding.

Results: Species composition, but not species richness, differed between leaf flushes. Across the latitudinal gradient, species richness was highest in the central part of the oak’s distributional range, and foliar fungal community composition shifted along the...
1 | INTRODUCTION

Leaves support a large diversity of fungi, both on their surface and within and between their cells (Jumpponen & Jones, 2009; Lafontaine-Lapointe & Whitaker, 2019; Redford et al., 2010; Turner et al., 2013; U’Ren et al., 2019; Vacher et al., 2016). These fungi are known to cause plant diseases, induce plant defences in response to biotic and abiotic stresses (Arnold et al., 2003; Jaber & Enkerli, 2017) and influence leaf senescence and decomposition (Vacher et al., 2016). While these fungi are crucial for plant health and ecosystem functioning, we lack insights into the patterns and drivers of within- and among-tree variation in the foliar fungal community (Agan et al., 2021; Allen et al., 2020; Millberg et al., 2015; Moler & Aho, 2018; Piepenbring et al., 2015; Sokolski et al., 2017; Terhonen et al., 2011; Unterseher et al., 2018). One factor that might explain within-tree variation in the foliar fungal community is the co-existence of leaves from different leaf flushes within individual trees. Many plant species produce multiple leaf flushes during the growing season, and newly produced leaves might differ from older leaves in chemistry and resistance against plant enemies (Agan et al., 2021; Piepenbring et al., 2015; Unterseher et al., 2018). At larger spatial scales, the foliar fungal community might vary along the distributional range of the host plant. For example, for a single fungal community might be more diverse at the southern range (e.g. latitudinal diversity gradient hypothesis; Mittelbach et al., 2007) or at the centre of the range (centre-periphery hypothesis; Pironon et al., 2017). Since the frequency of multiple leaf flushes is expected to change with global warming, and latitudinal gradients are commonly a proxy for climate, it is important to examine how the production of multiple leaf flushes and latitude jointly shape the structure of foliar fungal communities.

Many plant species produce leaves continuously, or in distinct flushes, during the growing season (Auerbach & Simberloff, 1984; Fuenzalida et al., 2019; Moles & Westoby, 2000; Prado et al., 2014). The production of multiple leaf flushes is linked to many factors, such as climatic conditions, plant age and herbivory (Bobinac et al., 2012; Hilton et al., 1987; Moles & Westoby, 2000). Physical and chemical differences between leaves from different flushes might explain within-tree variation in the foliar fungal community. If the establishment of foliar fungi is limited by dispersal, we might expect that leaves from earlier flushes have a higher diversity of fungi due to the accumulation of species over time. Alternatively, as young and developing leaves are often more susceptible to colonization, and the development of second and subsequent flushes frequently coincides with peaks of spore production, they might have a higher species diversity than leaves from the first leaf flush. As an illustrative example, the severity of oak powdery mildew on Quercus robur has been found to be lower on leaves from the first flush than on leaves from the second flush (Call & St. Clair, 2017). The only two studies that we are aware of that assessed the impact of the production of multiple leaf flushes on the foliar fungal community have found the highest species richness in leaves from the first leaf flush as well as differences in community composition between leaf flushes (Table S1; Agan et al., 2021; Unterseher et al., 2018). Knowing which guilds are more likely to grow in each leaf flush would further help us to understand how foliar fungal communities are distributed and structured within trees. Whereas a previous study focused on differences in the abundance of a single fungal guild (plant pathogens) between leaf flushes (Piepenbring et al., 2015), no study so far has examined differences in the relative abundance of other fungal guilds between leaf flushes.

The foliar fungal community associated with a given plant species might vary across its range (Millberg et al., 2015; Moler & Aho, 2018). Based on the latitudinal diversity gradient hypothesis, we might expect that the diversity of fungi associated with a host plant is highest at the southern part of the distributional range (Mittelbach et al., 2007). As an alternative, the environmental conditions might be suboptimal at the margins of a plant species’ distributional range, in which case we would expect that the largest proportion of the fungal community is present in the core of the distributional range of the host species (Pironon et al., 2017). As another alternative, there might be marginal asymmetry, where the fungal diversity is lower at the northernmost (expanding) margin, corresponding to newly colonized areas, whereas it is the same at the southernmost margin, where the species has been for much longer time. While few studies have examined latitudinal clines...
in the diversity of foliar fungal communities associated with a given host species, the existing reports are contradictory (Table S2), with studies showing increases (Sokolski et al., 2017), decreases (Terhonen et al., 2011) or no changes in the diversity of the foliar fungal community with latitude (Allen et al., 2020; Millberg et al., 2015). Likewise, reports on latitudinal changes in community composition are variable (Table S2). Millberg et al. (2015) and Moler and Aho (2018) showed that foliar fungal communities in needles from Pinus sylvestris and Pinus albicaulis changed with latitude, while Allen et al. (2020) reported that the foliar fungal community of reed (Phragmites australis) did not change with latitude. Importantly, previous studies only focused on restricted parts of the distributional range of the plant species (in particular, the northern or middle part of the range; see Table S2) and did not examine non-linear relationships between latitude and the structure of the foliar fungal community. Likewise, we lack information on how the relative abundance of fungal guilds changes with latitude. The relationship between the foliar fungal community and latitude might also differ between the leaf flushes. For example, we might expect more fungal species linked to secondary leaf flushes in southern Europe, where one can find the highest abundance of plants with multiple leaf flushes. As latitudinal clines are often characterized by large changes in climate, they might inform us about future changes in the climate (i.e. space-for-time substitution; De Frenne et al., 2013). It is thus important to understand the patterns and mechanisms behind latitudinal clines in the diversity and composition of the foliar fungal community across the plant’s distribution.

Differences in community composition between leaves belonging to different flushes and along latitudinal gradients can be due to species turnover or nestedness (Baselga, 2010). These two components are related to different ecological processes influencing fungal community assembly (Baselga, 2012; Hewitt, 1999; Svenning & Skov, 2007). For example, changes in the foliar fungal community composition due to species turnover can reflect species sorting by the environment or dispersal limitation, while species nestedness can reflect priority effects or differences in the level of specialization (Debray et al., 2021; Thébault & Fontaine, 2010; Toju et al., 2015; Weidlich et al., 2021). By knowing the relative contribution of species turnover and nestedness in shaping changes in community composition, we can thus further our understanding of how foliar fungal communities are structured and assembled.

We investigated the main and interactive effects of leaf flush and latitude on the foliar fungal community for a foundation tree species (the pedunculate oak, Quercus robur) along a latitudinal gradient of c. 20 degrees in Europe during a full growing season. For this, we sampled leaves from 20 oak populations from northern Spain to southwestern Finland during the early, mid and late season. More specifically we addressed the following questions:

1. Do species richness, evenness and community composition of the foliar fungal community differ between leaf flushes during the early, mid and late season?
2. How do species richness, evenness and community composition of foliar fungal communities change with latitude across the species’ range? Is this pattern similar for all leaf flushes?
3. Does the relative abundance of functional guilds (plant pathogens, mycoparasites, saprotrophs and symbiotrophs) differ between leaf flushes or along the latitudinal gradient?
4. Are differences in the foliar fungal community composition between leaf flushes and along the latitudinal gradient a product of species turnover or nestedness?

2 | MATERIALS AND METHODS

2.1 | Natural history

The pedunculate oak Quercus robur is a deciduous tree belonging to the Fagaceae that grows in a wide range of climatic conditions from northern Spain to southern Finland (Petit et al., 2002). The leaves of the pedunculate oak harbour a highly diverse fungal community on their surfaces (epiphytes) and within and between their cells (endophytes; Cordier et al., 2012; Jakuschkin et al., 2016). These foliar fungi differ strongly in their functional roles (Faticov et al., 2021; Jumpponen & Jones, 2009). Foliar fungi can be classified into several functional groups (sensu Nguyen et al., 2016), such as plant pathogens, mycoparasites (i.e. species that feed on cells of leaf-associated fungi), saprotrophs (i.e. species that feed on dead or decaying matter) and symbiotrophs (i.e. species that derive their nourishment from a symbiotic relationship with the plant). Notably, there is large variation in our knowledge of fungal species: There are species that are well described and whose ecology is studied extensively, such as pathogen species from the Erysiphe genus (Desprez-Loustau et al., 2011; Faticov et al., 2022), whereas other species are not described yet or their ecology is still unknown.

The pedunculate oak flushes its first leaves from April in the southern part of its distribution area and from the end of May in the northern part of its distribution area. Leaf senescence usually starts in September in the northern part of its distribution area and in October in the southern part of the distribution area (Crawley & Akhteruzzaman, 1988; Ekholm et al., 2019; Wenden et al., 2020). Pedunculate oaks are well known for having multiple leaf flushes during a single growing season (Hilton et al., 1987). The frequency of second and subsequent leaf flushes is strongly dependent on the climate. During the growing season, pedunculate oaks generally have three or four leaf flushes in southern Europe, two or three leaf flushes in central and western Europe, and one or two leaf flushes in northern Europe (Beikircher & Mayr, 2013; Hilton et al., 1987). The production of multiple leaf flushes can also be due to other factors, such as plant age, plant genetics, environmental conditions or as a compensation to herbivory in the first leaf flush (Beikircher & Mayr, 2013; Bobinac et al., 2012).

2.2 | Field sampling and sample preparation

To assess the influence of the production of multiple leaf flushes on the foliar fungal community across a latitudinal gradient in Europe during a full growing season, we selected 20 oak populations...
(avoiding urbanized areas) from seven countries across a latitudinal gradient in Europe with a minimum distance of 20 km among populations (Figure S1). In each population, we selected three adult (reproductive) oaks. From each oak, we randomly collected five fully expanded leaves from the first and (if present) second leaf flush, in both the early (8–12 June), mid (27–31 July) and late season (7–18 September). The five leaves were then placed together in a Ziploc bag with 70 g of silica gel for drying. From each leaf, we took eight 5 mm diameter leaf discs, four from each side of the midrib. Leaf discs were punched in a laminar flow hood with a metal corer, which was sterilized with 95% ethanol and flamed over a Bunsen burner after each sample. Leaf discs for each combination of tree and leaf flush (i.e. \( n = 40 \) leaf discs) were pooled into a single sample.

2.3 Molecular work

We milled the leaf discs using two metal beads in a TissueLyser II (Qiagen) and extracted DNA from 20 mg per sample using the Macherey-Nagel NucleoSpin Plant II kit, following the standard protocol. To characterize the foliar fungal communities, we used the forward primer ITS7 (Ihrmark et al., 2012) and the reverse primer ITS4a (White et al., 1990), which target a 250–450 bp fragment of the ITS2 region (Schoch et al., 2012). For full details on the molecular methods and bioinformatics, see Supplementary Text S1. Our final dataset contained 2,291,469 sequences assigned to 7030 amplicon sequence variants (ASVs). Foliar fungal communities were represented by 261 samples with an average of 8813 reads per sample. Each ASV was identified to species level by comparison with species hypothesizes (SHs) in the UNITE database v8 released on 12/08/2021 (Abarenkov et al., 2010). To calculate species richness (number of ASVs per sample) and Pielou’s evenness (Pielou, 1966), we rarified each sample to 1000 reads, omitting those samples with less reads and retaining 223 samples. From these 223 samples, 146 samples were from the first flush and 77 from the second leaf flush. We calculated species richness and evenness using the function estimate_richness from the R-package Phyloseq, which performs standard alpha diversity estimates by operating on the cumulative population of each sample (McMurdie & Holmes, 2013). We used MetagenomeSeq’s cumulative sum scaling (CSS) in the analyses of community composition as a normalization method to account for uneven sequencing depth (Paulson et al., 2013). Sequencing data have been deposited at SRA NCBI with the accession number (to be added upon manuscript acceptance).

2.4 Statistical analysis

As a general approach, we used linear mixed models and permutation multivariate analysis of variance (PERMANOVAs). For the univariate response variables, we fitted linear mixed models using the function lmer in the R-package lme4 (Bates et al., 2015; R Core Team, 2019), specifying a Gaussian distribution with an identity link, and testing for significance using the function Anova in the R-package car (Weisberg, 2019). We inspected the distribution of the residuals after running each model to check for normality and heteroscedasticity, and no model required transformation of the response variable. We calculated the marginal \( R^2 \) for the models using the function r.squaredGLMM in the R-package MuMIn (Barton, 2009). For the multivariate response variable community composition, we used the function adonis2 in the R-package vegan (Oksanen et al., 2015), with Bray–Curtis dissimilarity for the count data and Sørensen dissimilarity for presence-absence data. Since random factors cannot be specified in adonis2, the described random factors (below) were added as fixed factors in the community composition models. We used the function emmeans from the R-package emmeans (Lenth, 2020) for paired contrasts between leaf flushes or seasons.

To test if species richness, evenness and community composition of the foliar fungal community, as well as the relative abundance of the different guilds (proportion of reads belonging to each guild out of the total number of reads in each sample) differed between leaf flushes and changed with latitude during the growing season, we modelled each response variable as a function of leaf flush identity, latitude and season. As differences between leaf flushes might differ between the early, mid and late season, we included the interaction between leaf flush identity and season. To account for non-linear relationships along the latitudinal gradient, we included the quadratic term of latitude. As differences between leaf flushes might change with latitude, we included the interaction term between leaf flush and latitude, and between leaf flush and the quadratic term of latitude. To account for the hierarchical sampling design, we included the random factor population, and to account for repeated measurements, we included tree as a random factor. When we found a significant interaction between leaf flush and season, we used season-specific models to further investigate differences between leaf flushes in the early, mid and late season, where we included the fixed factor leaf flush identity and the random factors population and tree. To examine which species differed in abundance between leaf flushes, we conducted differential abundance analyses with the extension Deseq2 (Love et al., 2014) in the R-package Phyloseq (McMurdie & Holmes, 2013), adjusting P-values using the Benjamini–Hochberg’s procedure to account for multiple comparisons.

To test to what extent differences in community composition between leaf flushes and along the latitudinal gradient are explained by species turnover or nestedness, we used the R-package betapart (Baselga & Orme, 2012). We analysed differences in community composition between leaf flushes by computing dissimilarity values between the first and second leaf flush, both for the full growing season and separately for the individual seasons. We analysed changes in community composition along the latitudinal gradient by computing dissimilarity values between countries, in a series of pairwise comparisons of each country with its neighboring countries. In both analyses, we partitioned the total \( \beta \)-diversity (Sørensen dissimilarity) into two indices, where \( \beta \)-STU is the turnover component and \( \beta \)-SNE is the species nestedness component (Culp et al., 2019).
3 | RESULTS

3.1 Differences in the foliar fungal community between leaf flushes throughout the growing season

We obtained reads from 7030 ASVs, from which 3676 ASVs were identified up to species level with high confidence and 2576 ASVs could be assigned to functional group (sensu Nguyen et al., 2016). Species richness of the foliar fungal community increased with the progression of the growing season but did not differ between the leaf flushes (Figure 1a, Table 1). Evenness did not differ between either seasons or leaf flushes (Table 1). Differences in community composition between the early, mid and late season explained 2% of the variation in the foliar fungal community, and differences between leaf flushes explained 5%–7% of the variation (Figure 2, Table 1).

The relative abundance of plant pathogens tended to be lower in leaves from the first flush than in leaves from the second flush (Figure 1b, Tables S3 and S4). The relative abundance of mycoparasites was lower in leaves from the first flush than in leaves from the second flush, but only during the late season (Figure 1c, Tables S3 and S4). The relative abundance of saprotrophs and symbiotrophs did not differ between leaf flushes or throughout the growing season (Table S3). The relative abundance of ascomycetes with unknown function was generally higher in leaves from the first than in those from the second leaf flush (Figure 1d, Tables S3 and S4). The relative abundance of basidiomycetes with unknown functions increased from the early to the mid-season and differed between leaf flushes (Figure 1e, Tables S3–S5). The relative abundance of unknown fungi was higher in the early and late season than during the mid-season, and tended to be higher in the first leaf flush (Figure 1f, Tables S3 and S5).

The pathogens Didymella maydis, Microstroma bacarum and Oidiodendron griseum were more abundant in leaves from the second flush than in those from the first flush, even though these differences became non-significant when adjusting p-values for multiple comparisons (Figure S2, Table S6). While mycoparasites were generally more common in leaves from the second flush, the only species that significantly differed between the two leaf flushes was Sporobolomyces phaffii, which was more abundant in leaves from the first flush.

![Figure 1](https://onlinelibrary.wiley.com/doi/10.1111/jbi.14508)
first flush (Figure S2, Table S6). From the ascomycetes with unknown functions, *Aureobasidium subglaciale* and *Gnomoniopsis paraclavulata* were more abundant in leaves from the first flush than in those from the second flush (Figure S2, Table S6).

### 3.2 | Changes in the foliar fungal community across the latitudinal gradient

Species richness was highest in the central part of the latitudinal gradient, a pattern similar for leaves from the first and the second flush (Figure 3a, Table 1). Evenness increased from southern Europe to the central part of the latitudinal gradient and then levelled off at high latitudes (Figure 3b, Table 1). The foliar fungal community composition changed non-linearly with latitude, with a particularly steep change in community composition at higher latitudes, a pattern that was similar for leaves from the first and second leaf flush (Figure 3c, Table 1).

The relative abundance of plant pathogens decreased with latitude, while the relative abundance of basidiomycetes with unknown function and unknown fungi followed the opposite pattern (Figure 4a,c,d, Table S3). The relative abundance of saprotrophs changed non-linearly with latitude, but differently for the two leaf flushes: the relative abundance of saprotrophs in leaves from the first flush was lowest in the central part of the latitudinal gradient, whereas the relative abundance of saprotrophs in leaves from the second flush exhibited the opposite pattern (Figure 4b, Table S3). Mycoparasites, symbiotrophs and ascomycetes with unknown functions did not change in relative abundance along the latitudinal gradient (Table S3).

### 3.3 | Species turnover or nestedness?

Overall, differences in community composition between leaves from the first and second flush were mainly a result of turnover and only to a minor extent a result of nestedness (Figure 5a). The exception

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**TABLE 1** The impact of leaf flush, season, latitude (linear and squared) and its interactions on species richness, evenness and composition of the foliar fungal community of the pedunculate oak (*Quercus robur*). Shown are $\chi^2$ values (models on richness and evenness), $F$-values (models on community composition), degrees of freedom, $p$-values and marginal $R^2$ values. For community composition, results are given for both absolute counts and presence-absence of species. Statistically significant results ($p<0.05$) are shown in bold.

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<th>$p$</th>
<th>$R^2$</th>
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</table>
4 | DISCUSSION

To examine the effects of leaf flush and latitude on the foliar fungal community, we sampled leaves from the first and second flush in 20 oak populations across a latitudinal gradient in Europe during the early, mid and late season. Species composition, but not species richness and evenness, differed between the first and second leaf flush. Species richness was highest in the central part of the latitudinal gradient and the foliar fungal community composition changed with latitude, and these latitudinal patterns were similar for both leaf flushes. For the fungal guilds, the relative abundance of plant pathogens and mycoparasites was lower in leaves from the first flush than in those from the second flush, and the relative abundance of plant pathogens and saprotrophs decreased with latitude. Shifts in community composition between leaf flushes and across the latitudinal gradient were mostly a result of turnover and only to a minor extent a result of species nestedness. Our findings demonstrate that the co-existence of multiple leaf flushes and environmental variables associated with latitude play a key role in shaping small- and large-scale spatial variation in the foliar fungal community of a foundation tree in the temperate forests, with major consequences for plant health, species interactions and ecosystem dynamics.
We found that community composition, but not species richness and evenness, differed between oak leaf flushes, a pattern that was consistent between the early, mid and late season. The lack of a difference in species richness between leaf flushes in our study contrasts with the two previous studies on this topic, which all found a higher species richness on leaves from the first flush than on leaves from the second flush. Unterseher et al. (2018) found a higher species richness of foliar fungi in leaves from the first flush than in leaves from the second flush in tropical plants in Thailand, and Agan et al. (2021) found a higher species richness in Pinus sylvestris-needles from the current year than in needles produced in previous years in Estonia and Norway. While very speculative, this might suggest that there are differences in the effects of the production of multiple leaf flushes on the species richness of the foliar fungal community between tropical and temperate regions, or between coniferous trees (where leaves from different years co-exist) and deciduous trees. For evenness, no comparable studies exist. Our finding of a strong difference in the composition of the foliar fungal community between leaf flushes matches the two previous studies (Agan et al., 2021; Piepenbring et al., 2015; Unterseher et al., 2018). When investigating differences in functional guilds between leaf flushes, we found that plant pathogens and mycoparasites were more abundant in leaves from the second flush, while ascomycetes with unknown functions were more abundant in leaves from the first flush. In contrast to our finding of a higher relative abundance of plant pathogens in leaves from the second flush, Piepenbring et al. (2015) reported that plant pathogens in southwestern Panama were more likely to occur in older leaves than in young leaves. The difference in the relative abundance of plant pathogens between leaf flushes might be due to differences in the effect of leaf flush between tropical and temperate regions, but might also be explained by the identity of the host tree species. For example, in previous studies on oak trees, leaves from the second flush have been shown to be more susceptible to oak powdery mildew (Erysiphe spp.) infection (Gaytán et al., 2022; Marçais et al., 2009; Marçais & Desprez-Loustau, 2014), and this relationship between leaf flush and susceptibility might differ among plant species (Jain et al., 2019).

Our finding of a higher species richness of the foliar fungal community in the central part of the latitudinal range of the pedunculate oak supports the centre-periphery hypothesis, which states that the performance of species decreases from the centre to the margins of their distributional ranges (Pironon et al., 2017). This pattern might emerge either because the range of the oak-associated

FIGURE 4 The relationship between latitude and the foliar fungal community on the pedunculate oak Quercus robur, separately for each leaf flush, across a latitudinal gradient in Europe. Panels show the relationship between latitude in decimal degrees and the relative abundance of (a) plant pathogens, (b) saprotrophs, (c) basidiomycetes with unknown functions, and (d) unknown fungi. Statistically significant p-values are shown in the top-left of each panel. Shown are regression lines with their associated standard error (shaded area), separately for each leaf flush. Dots represent a leaf flush on a single tree. Only guilds with statistically significant results are presented; for the full set of panels, see Figure S5.
microorganisms matches that of the oak trees (e.g. for specialists), or because the lower performance of oaks at their distributional margins indirectly and negatively affects the performance of the associated microorganisms. While previous studies have shown increases (Sokolski et al., 2017), decreases (Terhonen et al., 2011) and no changes in the diversity of the foliar fungal community with latitude (Allen et al., 2020; Millberg et al., 2015), these studies did not target the full distributional range of the focal plant species, and did not examine non-linear latitudinal relationships. Evenness increased from the southern to the central part of the latitudinal range and then levelled off at high latitudes. Since there are no comparable studies for evenness of foliar fungal communities across latitudinal gradients, we cannot assess the generality of these results. Foliar fungal community composition changed with latitude throughout the distributional range, and this change was particularly pronounced at higher latitudes. Similarly, Millberg et al. (2015) and Moler and Aho (2018) reported changes in the foliar fungal communities (on Pinus albicaulis and P. sylvestris) across a latitudinal gradient, but both only focused on the mid and northern part of the host range, thereby corroborating our results on the mid and northern distributional range of oaks. Among functional guilds, the relative abundance of plant pathogens decreased towards higher latitudes, while basidiomycetes with unknown functions and unknown fungi exhibited the opposite pattern. The higher relative abundance of plant pathogens at lower latitudes matches the prediction that climate warming might increase the richness and severity of many plant pathogens at higher latitudes (Liu et al., 2019; Velásquez et al., 2018). We only found an interactive effect of leaf flush and latitude on the foliar fungal community for the guild of saprotrophic fungi: leaves from the first flush had the lowest relative abundance of saprotrophs in the central part of the latitudinal gradient, while saprotrophs in leaves from the second flush exhibited the opposite pattern. The foraging ascomycete hypothesis states that saprotrophs go through an endophytic stage in leaves to be able to colonize other substrates or survive times of resource scarcity (Nelson et al., 2020). Our results then suggest that leaves from the first flush are more likely to support the endophytic stage of saprotrophs in the edges of the distributional range of oaks, while leaves from the second flush are more likely to support the endophytic stage of saprotrophs in the central part of the distributional range of oaks. Changes in community composition between leaf flushes and along the latitudinal gradient were mostly a product of species turnover, and only to a lesser degree the result of species nestedness. The high contribution of species turnover in explaining differences in the foliar fungal community between leaf flushes might indicate that foliar fungi have a relatively narrow (realized) niche, and that the physical and chemical differences between leaves from different leaf flushes are important in the spatial separation of foliar fungi, and particularly so for plant pathogens and mycoparasites. The high contribution of species turnover in explaining the latitudinal change in community composition supports a previous meta-analysis on fungal communities, which showed that fungi are locally diverse, but have a hight spatial turnover at the biogeographic scale (Meiser et al., 2014). Taken together, the high species turnover suggests that both leaf flush and environmental variation related to latitude contribute to the high regional diversity of fungi. Importantly, different mechanisms might cause the same patterns: for example, turnover can be due to the strong dependency of spore dispersal and leaf colonization on environmental and climatic conditions (Ahanger et al., 2013; Linnakoski et al., 2017; Tibpromma et al., 2021), but also be due to strong interactions among fungal species, both of which are well-known for structuring fungal communities (Thébault & Fontaine, 2010; Toju et al., 2015). Overall, we need experimental studies to assess if changes in community composition of foliar fungi are mostly a product of the fundamental or realized niche, and whether this pattern differs among species living in different biogeographic regions with strong differences in mean and variance of climates, such as the tropics, temperate region and the arctic.

Overall, we identified leaf flush and latitude as two important mechanisms in shaping the distribution of foliar fungi at small and large spatial scales, respectively, in a foundation tree species in the temperate region. Using space-for-time substitution (sensu De Frenne et al., 2013), we can expect that the structure of the foliar fungal community on oaks will change with climate warming, with a notable increase in the relative abundance of plant pathogens and mycoparasites at higher latitudes. The increase in the relative abundance of plant pathogens might be even more pronounced due to the increase in the frequency of second and subsequent leaf flushes with climate warming, as at least the second leaf flush is predicted to have higher levels of plant pathogens. While leaf flush and latitude are rarely taken into account in studies on saprotrophic fungi, they are important factors determining the structure of the foliar fungal community on oaks.
account in studies on foliar fungal communities, our findings illustrate that these factors contribute to the maintenance of a high regional diversity in foliar fungi, and are important for our understanding of spatial variation in the foliar fungal community across multiple spatial scales.

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**CONFLICT OF INTEREST**

All authors declare no conflict of interest.

**DATA AVAILABILITY STATEMENT**

Data are available from the Dryad Digital Repository (Gaytán et al., 2022). https://doi.org/10.5061/dryad.931zcrjp.

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BIOSKETCH
Álvaro Gaytán – PhD at Stockholm University, where I study the community of organisms linked to oaks. One of my main interest is to understand how spatial and temporal variation in climate affects the life cycle of a diverse community of herbivores and fungi on oak, and the consequences for species interactions within this food web.

Author contributions: AG, KG and AJMT conceived and designed the experiment. AG, XM, BC, IVH, PDF, CM, BGHT, JPJGTH, PUR, NB, RJ, PP and SS carried out the field work. AG, MF and KP carried out the molecular work. MF and AA conducted the bioinformatic analyses. AG analysed the data with support from MF, AA and AJMT. AG wrote the first draft and all authors contributed to the final manuscript.

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