Fungi as mediators linking organisms and ecosystems

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One sentence summary: The unique set of traits that fungi exhibit, and the versatility of these traits facilitate the mediating role that fungi play in host and ecosystem functioning through the establishment of dynamic evolutionary and ecological interactions.

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ABSTRACT

Fungi form a major and diverse component of most ecosystems on Earth. They are both micro and macroorganisms with high and varying functional diversity as well as great variation in dispersal modes. With our growing knowledge of microbial biogeography, it has become increasingly clear that fungal assembly patterns and processes differ from other microorganisms such as bacteria, but also from macroorganisms such as plants. The success of fungi as organisms and their influence on the environment lies in their ability to span multiple dimensions of time, space, and biological interactions, that is not rivalled by other organism groups. There is also growing evidence that fungi mediate links between different organisms and ecosystems, with the potential to affect the macroecology and evolution of those organisms. This suggests that fungal interactions are an ecological driving force, interconnecting different levels of biological and ecological organisation of their hosts, competitors, and antagonists with the environment and ecosystem functioning. Here we review these emerging lines of evidence by focusing on the dynamics of fungal interactions with other organism groups across various ecosystems. We conclude that the mediating role of fungi through their complex and dynamic ecological interactions underlie their importance and ubiquity across Earth’s ecosystems.

Keywords: biotic interactions; fungal biogeography; symbiosis; mycovirus; mycobiota; omics

INTRODUCTION

Fungi are ubiquitous and form diverse communities temporally and spatially spanning multiple scales across many ecosystems (Fig. 1). We have only begun to scratch the surface of the importance of fungi in our lives and environment owing to their great diversity and challenges involved in studying fungal biology and ecology. Of an estimated 1.5–5 million fungal species only around 120,000 have been described (Hawksworth and Lücking 2017), largely due to the difficulty of isolating and culturing most fungi (Köljalg et al. 2013), which has also been a barrier to assigning them functional roles. Furthermore, it is difficult to distinguish where an individual of a fungal species begins and ends. This is highlighted by the tight association of fungi with other organisms e.g. through lichen, mycorrhizal, endophytic and mixed biofilm associations, but also by potentially frequent cross-kingdom horizontal gene transfer (HGT) between fungi and other organisms (Venice et al. 2020b), and in the heterogeneity of genetic and phenotypic intraspecific and intracellular variation that occurs in fungi (Ehinger et al. 2012; Taylor et al. 2017). With recent advances in molecular methods for gene-based identification of microbes over the last 20 years, some of these challenges have been overcome, leading to a rapid growth in our understanding about the importance of fungi in various ecosystems (reviewed in Peay, Kennedy and Talbot 2016; Nilsson...
Figure 1. Fungi form diverse entities that are distributed across different aspects and scales of space and time. Fungal diversity can be viewed across different levels of biological organization from the gene level to the level of complex communities, as well as from a taxonomic, phylogenetic or functional perspective. Even at the individual level, fungi exhibit different life cycle stages, that all have their unique biogeographical patterns e.g. the growth phase of fungi (yeast or filamentous) is embedded within or on a substrate, whereas the reproductive and dormant phases of fungi usually bridge different substrates/habitats. This is exemplified by a soil dwelling fungus from the family Agaricaceae that exists as a mycelium (growth phase) within the soil matrix, which then produces a fruitbody in the form of a mushroom that bridges habitats between the soil and the atmosphere. Finally, the fungus releases spores that may make their way into the atmosphere before landing in a new patch of soil, each growth phase is subjected to unique environmental factors, biological interactions, and other processes that lead to distinct biogeographic patterns. As fungi bridge the gap between micro- and macro-organisms, they also exhibit spatial patterns at the microhabitat to macro habitat level, including across different depths of a substrate, between different substrates, between different organs of a host, through to macroecological scales such as across landscapes to biomes and the global level. Fungi also exhibit distinct biogeographic patterns across different temporal scales from days to seasons, through to years and geologic time.

et al. 2019). In terrestrial habitats, while relying on their hosts for carbon resources, mutualistic and pathogenic fungi play key roles in driving the physicochemical and biological properties of their environment as well as host population dynamics (Alan Pounds et al. 2006; Clemmensen et al. 2013; Bagchi et al. 2014; Chen et al. 2019; Tedersoo, Bahram and Zobel 2020). Meanwhile saprotrophic fungi are the key decay agents of organic material, playing a central role in carbon and nutrient cycling in many ecosystems (Averill, Turner and Finzi 2014; Fernandez and Kennedy 2016; Netherway et al. 2021). In aquatic habitats, fungi can regulate food-web dynamics and biogeochemical cycling by rendering resources more available to higher order consumers and increasing the efficiency of trophic transfer through aquatic ecosystems (reviewed in Grossart et al. 2019).

With large and varying phylogenetic and functional diversity as well as great variation in dispersal modes, fungi can act as important pathogens, commensals, and mutualists of macro-eukaryotic hosts, and interact with and influence the activity and functions of microbial prokaryotes and eukaryotes in most ecosystems (Fig. 2). They bridge the divide between micro- and macroorganisms, from single celled yeasts to complex macro-fungi that produce structures rivalling plants and animals in terms of size. Recent studies have provided unprecedented insights into the ecology and biogeography of fungi. There is growing evidence pointing towards commonalities and differences in the community assembly patterns and processes of fungi compared to those traditionally considered for microorganisms such as bacteria, but also from macroorganisms such as plants and animals (Tedersoo et al. 2014; Davison et al. 2015; Tisthammer, Cobian and Amend 2016; Kivlin et al. 2017; Bahram et al. 2018; Cameron et al. 2019). The characteristics of fungi and their ecology, including their associations with other organisms, may have enabled fungi to become central agents of ecosystem, ecological, and evolutionary processes (Fig. 2), and to span multiple scales and dimensions of time, space, and biological interactions (Figs 1 and 3). The combined and increasing evidence suggests that fungi can mediate links between different organisms and ecosystems, and potentially alter ecological, evolutionary and biogeographic relationships of the organisms they interact with. Here we review this evidence by exploring the current knowledge around the dynamics of fungal ecological interactions and underlying mechanisms in various ecosystems and discuss how these affect the evolution, ecological fitness and distribution of their associated organisms. We further present a perspective for embracing the importance of fungal interactions in ecosystems.
THE ECOLOGICAL VERSATILITY OF FUNGI

Fungi exhibit diverse lifestyles (nutritional modes), reproductive and dispersal strategies and physicochemical traits, which could play important roles in their success (Fig. 2). Nutrient, carbon, and water acquisition is the central function of all organisms, and the heterogeneous distribution of these resources spatially and temporally is the driving force of evolutionary diversification and adaptation. Fungal hyphae with their indefinite growth habit and the formation of highly interconnected mycelial networks represent a key evolutionary adaptation to terrestrial life that facilitates the foraging of unevenly distributed resources and maximizing the efficient assimilation of these resources between distal patches with incredible plasticity (Fricker et al. 2017; Kiss et al. 2019).

Ecophysiological traits

Recent work suggests that we have likely just scratched the surface in terms of understanding the ecophysiological capabilities of fungi. The well-developed and diverse secretome (i.e. the total set of molecules secreted by cells) of fungi contributes to their metabolic versatility by establishing saprotrophic, mutualistic and pathogenic interactions (Bouws, Wattenberg and Zorn 2008; Choi et al. 2010; Girard et al. 2013). Fungal secretomes overlap greatly between mutualistic, saprotrophic and pathogenic lifestyles (Plett and Martin 2015; Hess et al. 2018), thereby potentially facilitating lifestyle transitions in order to evolve and adapt to varying environmental conditions (Girard et al. 2013). These secretomes enable fungi to extract nutrients from differing and complex substrates, but also protect fungi from hostile environmental conditions (Girard et al. 2013). Symbiotic and free-living fungi can inhabit extreme but vast areas of drylands and play an important role in rock weathering and thereby soil formation and create microhabitats for other organism groups (Coleine et al. 2021). Combined with their hyphal forming habit, perhaps as their key characteristics, the enzymatic arsenal of fungi has facilitated their dominance in inhabiting and decomposing plant derived substrates. The high diversity of carbohydrate-active enzymes (CAZymes) of pathogenic and saprotrophic fungi gives them the flexibility to adapt to different plant species and their resources, and the diversification of fungal nutritional modes is evolutionarily linked to the diversification of angiosperms and gymnosperms (Janusz et al. 2017). A recent study suggests that fungal species with the highest capacity and broadest diversity of enzyme substrates have unexpected life-forms, such as dark septate endophytes and endolithic black
Figure 3. Examples of mediating role of fungi in different ecosystems. Fungi through their unique morphological and ecophysiological properties act as mutualists, commensalists, and antagonists with plants, animals, microbes and other fungi, mediate the health, performance, population dynamics and biogeography of these organisms. Meanwhile, through their affinity for and ability to break down complex substrates (notably plant derived) and even contribute to mineral weathering, fungi mediate carbon and nutrient cycles in both terrestrial and aquatic ecosystems, and through their enormous production and release of spores into the atmosphere fungi may even mediate rainfall. For example, in terrestrial systems mycorrhizal fungi mediate nutrient acquisition of plants as well as their interactions with antagonists such as plant pathogens and herbivores, while saprotrophic fungi mediate the cycling of complex plant derived substrates, and fungal pathogens mediate the population dynamics of eukaryotic hosts. In aquatic systems fungi mediate food web dynamics by controlling resource fluxes to higher order consumers in the process also mediating the efficiency of transfer across trophic levels. In human habitats such as skin and gut, the mycobiome, through complex interactions with other microbes, can play a key mediating role in human health and dysbiosis.

The diversity of secondary metabolites produced by fungi, which are central to both offensive and defensive responses (Rohlfs and Churchill 2011), facilitate associations with hosts (Keller 2019), while (positively or negatively) interacting with and influencing bacteria in establishing interactions with hosts (Bahram et al. 2018; Durán et al. 2018; Pierce et al. 2021). In addition, fungal secondary metabolites play a role in spore germination, either inhibiting spores from competing species or promoting their own (Becker et al. 2012; Khalid et al. 2018). Though underexplored, at least some key fungal secondary metabolites—including aflatoxins, melanin and β-lactam antibiotics—appear to be highly compartmentalized in vesicles that aid in their storage and transport under various conditions such as light and nitrogen limitation (Chanda et al. 2009; Upadhyay et al. 2016; Martin 2020). Fungi can regulate their secondary metabolite production based on environmental conditions, in particular light and temperature via both transcriptional and epigenetic regulations (Keller 2019).

As another important trait in lifestyle flexibility, fungal melanisation (which occurs in most fungal clades) allows fungi to resist environmental stressors (such as high temperatures, free radicals) and offers photoprotection (Cordero and Casadevall 2017), such as against UV radiation. Fungal melanin can also act as a virulence factor (reviewed in Jacobson 2000) and plays an important function in fungal infection across diverse hosts, supported by evidence that highly melanized fungi (belonging mainly to the Dothideomycetes class) are more capable of infecting organisms across different kingdoms, and switching and sharing hosts including between plants and humans.
(Gauthier and Keller 2013) as well as having broad geographical distributions. There have been several human disease outbreaks caused by highly melanized fungi which have likely been directly sourced from the environment, and it appears that the breakdown of physical barriers protecting against direct infection of fungi and their abilities to utilize host nutrients play an important role in the onset of fungal outbreaks (reviewed in Gauthier and Keller 2013). A recent example of an outbreak caused by a highly melanized fungus is the ‘black fungus’ epidemic infecting Covid-19 patients with weakened immune systems in India (Dyer 2021). While it remains unclear to what extent melanin contributes to fungal virulence, overall melanized fungi appear more capable to resist host defences compared to non-melanized fungi.

**Lifestyle versatility**

Another striking property of some fungi is their ability to transition between different lifestyles, which has enabled them to associate with a wide variety of hosts in various environmental conditions and adapt to spatially and temporally heterogeneous resources. While fungi have evolved different lifestyles, certain groups have retained many of the ancestral genes that enable transition between different nutritional modes (Hess et al. 2018). This capability together with retaining genomic potential to associate with a wide variety of hosts while adapting to new conditions (Genre et al. 2020), facilitated by HGT events (Wang et al. 2021) or by exchange of viruses via horizontal viral transfer (HVT) (Marquez et al. 2007), could contribute to host range expansion of mutualistic fungi, which is reflected in the relatively low host specificity of mutualistic in contrast to pathogenic fungi (Polme et al. 2018). The frequent HGT and HVT events between fungi and their hosts (Bian et al. 2020; Wang et al. 2021) may also have enhanced their adaptive capacity with their hosts, including terrestrialization (Wang et al. 2021). A recent study provides direct evidence that fungal viruses can contribute to fungal lifestyle versatility by mediating transitions from pathogenic to non-pathogenic phenotypes (Zhou et al. 2021). There is also evidence that HGT events occur more frequently between different fungal species, compared to other eukaryotic groups (Khalid et al. 2008), which can further contribute to their ecological versatility.

A particularly interesting case of evolutionary fungal lifestyle transitions is represented by the ectomycorrhizal (EcM) lifestyle, a convergent lifestyle that has arisen independently around 80 times or more from ecologically diverse saprotrophic ancestors (Miyauchi et al. 2020). While not true EcM fungi, a diverse variety of saprotrophic fungi can facultatively colonize the roots of plants; however, the functional significance of this relationship is unknown, though it has the potential to represent a transition stage between saprotrophy and an EcM symbiosis (Smith et al. 2017). This convergent evolution is in stark contrast to the arbuscular mycorrhizal (AM) symbiosis which has arisen once and remained relatively unchanged over hundreds of millions of years, although the AM symbiosis can perhaps act along a mutualism-parasitism continuum depending on host and environmental conditions (Johnson et al. 2015). The versatile boundaries between fungal lifestyles appears to be the norm rather than the exception. One piece of evidence supporting this view is that most of the fungi capable of infecting both humans and plants are soil saprotrophs that can be opportunistic pathogens (Gauthier and Keller 2013). Dimorphic switching between a filamentous form and yeast form is a particularly powerful trait that facilitates the transition between lifestyles based on environmental conditions and habitats. This is exemplified by thermally dimorphic fungal pathogens that have evolved the ability to switch from a hyphal form in the environment to a single celled yeast form when infecting an animal host, a phenomenon that is triggered by elevated temperatures (Li and Nielsen 2017).

Such versatility in switching between various symbiotic and free-living lifestyles may give fungi a unique capability to survive under unfavourable host and environmental conditions.

**Reproductive traits**

Reproduction is at the centre of life, with both sexual and asexual reproduction incurring both costs and benefits. Fungi appear to exhibit diverse and often unique and cryptic modes of reproduction, ranging from asexual to sexual, and often both strategies are employed to some degree (Nieuwenhuis and James 2016). It is often assumed that sexual recombination should be most advantageous in terms of accelerating evolutionary adaptation through natural selection and eliminating deleterious mutations (Aanen and Hoekstra 2007). Nevertheless, around 20% of fungal species may be solely asexual (Dyer and Kück 2017), including many evolutionary and ecologically successful fungi such as members of the Glomeromycota that form AM associations, although the presence of strictly asexual reproduction in the Glomeromycota is debated with growing evidence for the contrary (Yildirir et al. 2020). In addition, a sexual cycle has recently been observed in other fungal species previously thought to be solely asexual (Dyer and Kück 2017). Interestingly several fungi exhibit an alternative mechanism to sexual recombination in the form of parasexual recombination (Pontecorvo 1946), which is unique to the fungal kingdom and facilitates genetic transfer without meiosis and the production of sexual structures (Steensels, Gallone and Verstrepen 2021). Parasexuality in fungi may play a key role in pathogenesis in both plant and animal hosts (Caio, Billmyre and Heitman 2013), exemplified by Candida albicans in humans, and the rice blast fungus Magnaporthe oryzae in plants (Steensels, Gallone and Verstrepen 2021). Thus, the diversity and flexibility of reproductive strategies in fungi appears to play a central role in their evolutionary and ecological success. These reproductive strategies play a central role in the adaptation and dispersal of fungi to new environments, as discussed below.

**Dispersal traits**

Together with their reproduction strategies, the unique dispersal traits of fungi play a key role in their ability to span multiple scales of space, time and biological interactions (Fig. 1). Sporulation is a successful and phylogenetically widespread strategy for survival and dispersal that is employed by species of bacteria, protists, fungi and plants, yet fungi may represent most of the spore-forming organisms of the biosphere (Huang and Hull 2017). Sporulation contributes to the ease of fungal dispersal via different vectors. One of these vectors is air currents, which can carry a large proportion of biogenic aerosols comprising fungal spores, as evidenced by a high relative abundance and diversity of fungi in air (Gusareva et al. 2019). Another important dispersal vector for fungi is animals, which range from passive to tightly evolved symbioses and aid in both short and long-distance dispersal (Golan and Pringle 2017; Nuske et al. 2019; Vašutová et al. 2019). Animal associated fungi that are adapted to their host environment may show limited distribution following their hosts, such as fungus-growing ants and termites
with limited distribution ranges (Biedermann and Vega 2020). Another example is the association between truffle-like fungal taxa which—due to forming underground fruiting bodies—rely on mycorrhagous mammals (Nuske et al. 2019) and birds (Caiafa et al. 2021) for dispersal, while the distribution of fungus-dependent animals can be heavily influenced by the production and distribution of fungal fruiting bodies (Claridge 2002). Also, some fungal individuals such as members of the genus Armilaria can disperse over large distances via vegetative growth, forming extensive mycelial networks that may form some of the largest organisms on Earth (Anderson et al. 2018). Overall, there appears to be large variation in the reproductive and dispersal abilities of different fungal taxa that may enable them to occupy complementary niches across various ecosystems.

**Genomic traits**

Last, but not least and connected in some respect to all the above-mentioned traits, one of the main features of fungi is their relatively small and dynamic genomes, which could lead to their greater ecological plasticity compared to other eukaryotes. This is supported by recent evidence suggesting that among AM fungi, those with large genomes show less versatility in host and habitat preferences, as shown in the case of Gigaspora margarita possessing by far the largest annotated fungal genome (Venice et al. 2020b). Also, parasitic fungi associate with a broader range of host species, compared to animal or protozoan parasites with much larger genomes compared to fungi (Ogburn 2019). In addition, the large variability of genome size among fungi can be mostly ascribed to the abundance of transposable elements, which in turn has been shown to be related to the evolution of symbiosis as well as the versatility of fungal lifestyles and genomic plasticity (Hess et al. 2014; Castanera et al. 2016; Faino et al. 2016; Frantzekakis et al. 2018; Miyachi et al. 2020). This is thought to be of particular importance for plant pathogenic fungi in agricultural landscapes that can support large and stable populations of pathogens that can rapidly adapt and evolve new virulence traits in response to host resistance factors even with low effective population sizes, in part due to the accumulation of and variability in the presence of transposable elements within species and between closely related species (Möller and Stukenbrock 2017). The adaptive evolution of fungi in their host environment may extend beyond their genomes, as evidenced by a recent study pointing to the role of epigenetic factors in plant and human fungal pathogens in developing resistance against antifungal agents (Torres-García et al. 2020). Fungal secondary metabolism can also be highly flexible and manipulated by epigenetic chromatin regulators (Pfannenstiel and Keller 2019); notably methylation patterns in fungi appear to be maintained by natural selection not acting on the DNA sequence as the substrate, hinting at the inheritance of phenotypic traits (Madhani 2021).

Taken together, the ecological versatility of fungi seems unrivalled by other organism groups, which is facilitated by their unique dispersal, genomic, morphological, and ecophysiological attributes that are often plastic, thus considering them within a larger framework of traits including biotic interactions (Fig. 2), provides a more integrated understanding of fungal ecology and their central role in ecosystem and host functioning.

**EVOLUTIONARY AND BIOGEOGRAPHICAL PERSPECTIVES**

The early diverging fungi appeared to be endosymbiotic of eukaryotic hosts (Bonneville et al. 2020) and relied on their hosts for primary metabolism (Berbee, James and Strullu-Derrien 2017; Strullu-Derrien et al. 2018). These early fungi were zoosporic, which diverged more than 700 million years ago into non-flagellated fungi, followed by the most diverse fungal lineage of Dikarya around 642 Ma (Tedesco et al. 2018). These diversification events were associated with fungal land colonization, which coincided with the diversification and colonization of land by plants and animals (Berbee, James and Strullu-Derrien 2017; Minter et al. 2017; Lutzoni et al. 2018).

**Plant-associated fungi**

It has been hypothesized that mycorrhizal-like fungi, may have facilitated the successful colonization of land by plants through acting as their primary nutrient uptake mechanism in a nutrient-limited environment (Pirozynski and Malloch 1975), which is plausible given the wide range of extant plants associating with mycorrhizal fungi (>90% of plant families). A growing body of evidence supports this hypothesis. For example, fossil records show the emergence of fungi prior to plants as well as the presence of arbuscular mycorrhizal (AM) fungus-like remnants in embryophytes (i.e. earliest diverging land plants (Remy et al. 1994; Redecker 2000; Bonniveille et al. 2020; Gan et al. 2021), while phylogenetic analyses show the diversification of terrestrial fungi predates that of embryophytes (Lutzoni et al. 2018). Furthermore, experimental results show orthologous genetic pathways are involved in the symbiosis of AM fungi with embryophytes and vascular plants (Rich et al. 2021) and CO$_2$-rich conditions during early plant evolution likely enhanced the net benefit of the AM symbiosis (Humphrey et al. 2018). Recent studies also provide evidence that the evolution of mycorrhizal symbioses has led to the diversification of EcM fungi and their host plants, based on comparing the centre of diversification and origin of hosts with their fungal partners (Looney et al. 2016; Brunnett and Tedersoo 2018). A few studies based on molecular phylogenies show that while some clades may originate from the tropics (e.g. Clavulina (Kennedy et al. 2012)), several major EcM clades appear to show a higher diversification rate in temperate regions (Kennedy et al. 2012; Looney et al. 2016). The diversification centres of at least some EcM fungal clades may have coincided with the distribution range expansion of their host plants, which is greater in temperate regions (Steidinger et al. 2019). This is further supported by shared evolutionary histories of plants and EcM fungi like those belonging to the genus Amanita (Sánchez-Ramírez et al. 2015). In terms of contemporary plant biogeography, due to their heavy reliance on fungal partners in their earlier life stages, the distribution ranges of orchids can be limited by the presence of suitable mycorrhizal fungi (McCormick, Whigham and Canciani-Viruet 2018). There is also evidence suggesting that the limited distribution of mycorrhizal trees beyond the tree line might be at least partly due to a lack of suitable fungal partners (Shemesh et al. 2020). Plant-associated fungi such as mycorrhizal fungi clearly play a central role the in macroecology of plant communities owing to their tightly intertwined evolutionary histories (Tedesco, Bahram and Zobel 2020).

**Animal-associated fungi**

Much less is known about the evolution of animal-associated fungi, despite being phylogenetically more closely related, compared to fungi and plants (Wainright et al. 1993). Based on fossil evidence, the earliest animal-associated fungi are likely parasites that shifted their interactions from plants to insects (Lepidoptera), and these fungi show a Cretaceous diversification like that of their insect and plant hosts (Sung, Poinar and...
et al. (2012), and these associations date back to the mid Paleogene based on fossil records of ant feeding behaviour driven by the fungus (Hughes, Wappler and Labandeira 2011). This is supported by phylogeographic analysis of a closely related species (O. sinensis) and its host insects, which showed that both interacting partners have influenced the evolution and diversification of each other (Zhang et al. 2014).

Biogeographic patterns of host-fungal interactions

Large-scale distribution patterns of fungi also support co-diversifications and similar evolutionary histories of symbiotic fungi and their hosts. Many fungal groups show similar diversity distribution patterns as their hosts. In particular, the diversity of animals and fungal animal parasites and pathogens follow a latitudinal diversity gradient, i.e. they are highest in tropical regions relative to other biomes (Willig, Kaufman and Stevens 2003; Tedersoo et al. 2014), suggesting a potential causal link between diversity of parasites and animal hosts. Plant-associated fungi, such as foliar endophytes (Arnold and Lutzen 2007), AM fungi (Davison et al. 2015) and plant pathogens (Tedersoo et al. 2014), appear to also show an increasing diversity pattern towards the equator (Arnold and Lutzen 2007; Tedersoo et al. 2014; Davison et al. 2015), which follows that of their host plants (Francis and Currie 2003; Hillebrand 2004; Kreft and Jetz 2007). Nevertheless, compared to other groups of symbiotic fungi, ECM fungi also exhibit a contrasting latitudinal diversity gradient, being highest in temperate and boreal forests compared to tropical forests (Tedersoo et al. 2012). This has been ascribed to the greater cover of ECM trees in temperate and boreal forests compared with tropical forests, which is consistent with the species-area relationship, as also reflected in higher species diversification rates of different ECM fungal clades (Kennedy et al. 2012; Tedersoo et al. 2012).

Multipartite evolutionary dynamics

In addition to their historical interactions with animals and plants, fungi have likely been exposed to gene exchanges with bacteria through HGT events—because of their shared niches and constant physical contact—and this may have enabled fungi to more efficiently use carbon sources and interact with other eukaryotes including plants (Naumann, Schüßler and Bonfante 2010; Bruto et al. 2014; Mares et al. 2015; Torres-Cortés et al. 2015; Venice et al. 2020b). Such tripartite interactions may be ancestral, as shown in the case of the AM symbiosis (Naumann, Schüßler and Bonfante 2010). A recent study suggests that by strengthening each other’s fitness (i.e. fitness alignment), interactions of AM fungi and bacteria (particularly nitrogen-fixing) with the same host plant may synergistically shape the evolutionary trajectory of the three interacting partners (Afkhami, Friesen and Stinchcombe 2021). Although the underlying mechanism remains little explored, it is likely that HGT events among these mutualistic partners contribute to their fitness alignment, as implied by evidence of cross-kingdom gene transfer among AM fungi, bacteria, and plants (Li et al. 2018; Venice et al. 2020a, b). Besides increased adaptations to new soil environments and hosts as a result of HGT events, fungi may have also facilitated the transfer of bacterial genes—in addition to those from fungi—to plants, with potential implications for plant-fungal co-evolution and their co-diversification (Sinn and Barrett 2020). This is plausible given that fungi could be intermediate links between bacteria and plants, which is supported by evidence for ancient HGT events between mycorrhizal fungi with both bacteria and plants (Venice et al. 2020b). It remains to be seen whether HGT events have been more frequent between plants and their specific symbiotic fungi, e.g. between ECM plants and fungi. Fungal hyphae are also known to act as an anchor for attachment and colonization of bacteria inside plant tissues, as well as the formation of mixed biofilms that act as HGT hotspots and enable microbes to better cope with unfavourable conditions on host plants (reviewed in Overbeek and Saikkonen 2016).

Another interesting evolutionary and ecological interaction between fungi and their eukaryotic hosts comes in the form of viruses. Fungi and their viruses can contribute to plant adaptation to extreme environmental conditions. This has been shown in the case of a tropical panic grass, where a fungal endophyte confers heat resistance to the plant but only when the fungus is infected with a specific virus, suggesting a tripartite plant-fungus-virus symbiosis (Márquez et al. 2007). Fungal viruses may also play a key role in determining the expression of various traits of endophytic and phytopathogenic fungi including growth rate, morphology and pathogenicity (Xie and Jiang 2014; Zhou et al. 2021). Viruses are common in both plants and fungi, and—as eukaryotes—fungi and plants share viral families with closely related taxa that are genomically integrated in both hosts (Liu et al. 2010). Yet, the same does not appear to be as common between fungi and animals, which are more closely related than fungi and plants, once again highlighting the strong evolutionary and ecological link between fungi and plants since splitting from animals (Roossinck 2019). Thus, HGT events via shared or closely related viruses are perhaps a key mechanism that facilitates the intimate relationships fungi have with plants as endophytes, pathogens and mycorrhizal fungi. In support of this hypothesis, there is a growing number of studies pointing to compatibility and cross-kingdom HGT between fungi and plants (Andika et al. 2017; Nervo et al. 2017; Bian et al. 2020; Sutela et al. 2020), which could have contributed to the evolution of their viromes. Evidence also suggests that fungi can facilitate the spread of plant viruses to other plant hosts, e.g. via fungal spores as a vector (Bian et al. 2020). Viruses may exploit mycoparasitic insects to transmit between plant-pathogenic fungal hosts and form a complex quadripartite virus-fungus-insect-plant interaction with ecological and evolutionary implications (Liu et al. 2016). How exactly fungi and viruses co-evolve remains to be established, but it is plausible that their interactions contribute to fungal success in many ecosystems as discussed in the following sections.

Overall, evidence points to the role of fungi in the evolutionary ecology of their hosts and suggest that land colonization might have been a breakpoint in fungal diversification and evolutionary trajectories, owing to the formation of tight associations and interactions with other organism groups (Naranjo-Ortiz and Gabaldón 2019).

**Fungal Interactions in Host Systems**

As fungi are a major component of the microbiota in many ecosystems (Bahram et al. 2021), their interactions with hosts and other microbes are extremely relevant to consider in understanding host and ecosystem functioning. Although the underlying mechanisms remain unclear, there is a growing body of knowledge pointing to the importance and dynamics of fungal interactions and the role of fungi in mediating interactions between other organisms across ecosystems as discussed below (Fig. 3).
Fungal-plant interactions

The association between plants and fungi is one of the most well studied interactions. Plants as sessile organisms have evolved to live in close association with diverse communities of microbes—of which fungi are a major component—in order to meet their nutritional requirements for growth, survival and reproduction. Thus, they have developed multiple strategies to modulate the composition of microbial communities to their advantage in order to adapt to changing biotic and abiotic conditions (Bulgarelli et al. 2012). These strategies involve the ability to favour or select for a beneficial microbiome while avoiding pathogens from the surrounding environment (Bulgarelli et al. 2012; Bahram et al. 2021). There is also evidence that plants and mycorrhizal fungi prefer associating with partners that provide greater benefits for both partners (Bever et al. 2009; Kiers et al. 2011; Bogar et al. 2019), thereby maintaining their interactions under various environmental conditions. Such selective forces by both partners are reflected in the strong association between host identity or genotype and fungal community structure (Gehring et al. 2014; Martínez-García et al. 2015; Küngas, Bahram and Pöldmaa 2020; Leopold and Busby 2020). Host specificity of both pathogenic and mutualistic fungi can be reflected in their genomes (Baroncelli et al. 2016; Lofgren et al. 2020). Nevertheless, the strength of these interactions can depend on the presence of a third interacting partner (Lehtonen et al. 2005) as well as environmental conditions and resource availability (Johnson et al. 2015). For example, positive feedback mechanisms in plant–microbial systems may be enhanced by mutualistic interactions of mycorrhizal fungi and N-fixing bacteria (Palakurty, Stinchcombe and Afkhami 2018), which can synergistically increase the fitness of all interacting partners (Afkhami, Friesen and Stinchcombe 2020). The specificity of both pathogenic and mutualistic fungal taxa can be reflected in their genomes (Baroncelli et al. 2016; Lofgren et al. 2020). Nevertheless, the strength of these interactions can depend on the presence of a third interacting partner (Lehtonen et al. 2005) as well as environmental conditions and resource availability (Johnson et al. 2015). For example, positive feedback mechanisms in plant–microbial systems may be enhanced by mutualistic interactions of mycorrhizal fungi and N-fixing bacteria (Palakurty, Stinchcombe and Afkhami 2018), which can synergistically increase the fitness of all interacting partners (Afkhami, Friesen and Stinchcombe 2020). Due to the integrated nature of many of these associations between plants and microbes, notably mutualistic fungi, the plant holobiont has been suggested to be a unit of biological organization and selection (Vandenbrocknuyse et al. 2015); however, it is an open question whether such a level of organization can be considered through a macroecological perspective, i.e. can a holobiont disperse as a unit or is it determined by environmental selection following independent dispersal processes of the macro and microorganisms involved? Yet both antagonisms, mutualisms, as well as commensalisms, between land plants and fungi are known to contribute to fungal and plant biogeography (Berbee, James and Strullu-Derrien 2017; Lutzoni et al. 2018; Delavaux et al. 2019), and these interactions strongly impact terrestrial biogeochmical cycles and community dynamics (Tedersoo, Bahram and Zobel 2020).

Fungal-animal interactions

Although the interactions of fungi with animal hosts remain relatively less known, compared to plants, there is also evidence revealing interesting dynamics. For example, animal associated fungal pathogens are one of the most generalist groups of pathogens in terms of host and habitat requirements, highlighted by evidence for the exchange of functional fungi between ecosystems in the case of the transfer of pathogens from human to plant and vice versa (Gauthier and Keller 2013). Such transfers may be facilitated by vectors such as mosquitoes, which are known to carry human opportunistic fungal pathogens existing in non-human environments (Bozic et al. 2017). In addition, some fungal lineages show evolutionary evidence of frequent host switching between plants and insects (Sung, Poinar and Spatafora 2008) and between different insect lineages (e.g. between beetles and ants Araújo and Hughes 2019). There is also evidence that a pathogenic fungus (Erysiphe alphitoides) can interact with various insect herbivores, with a wide range of positive to negative ecological outcomes for the interacting insect depending on species identity, which can affect the structure of insect communities and their interactions with plants at the landscape scale (Tack, Gripenberg and Roslin 2012). A more recent study provides further evidence that pathogenic fungi can mediate the interaction between insect herbivores from competition to facilitation (Klutsch et al. 2016). While the exact mechanisms of the mediating role of fungi remains unknown, besides their direct interactions, fungi may also indirectly interact with insect herbivores by inducing defence signalling pathways and altering the quantity and quality of resources and volatiles in the host plant (Rostáš et al. 2006; Klutsch et al. 2016; Bastías et al. 2018; Eberl et al. 2018; Bu’thner et al. 2021). Also, fungal tissues can provide a richer source of nitrogen and amino acids compared to plant tissues and may thus attract herbivores (Biedermann and Vega 2020; Eberl et al. 2020). Indeed insect–fungus mutualisms—where fungi provide a nutritional source for insects, such as ambrosia beetles, attentive ants and termites, and in turn fungi receive protection and dispersal services from insects—is widespread across insect and fungal lineages, with important roles in ecosystem functions (reviewed in Biedermann and Vega 2020). The interaction of plant–insect–fungus also extends to mycorrhizal fungi, where mycorrhizal associations can modify the production and composition of terpenoids, which may contribute significantly to the plant defence response against herbivore attacks (Sharma, Anand and Kapoor 2017). Another example of fungal–insect–host interactions is hyperparasitism, where fungi parasitize parasitic insects, e.g. Laboulbeniales, associated with bat flies (Jensen et al. 2019).

In aquatic ecosystems, perhaps the most well-known groups are notorious aquatic fungal pathogens and parasites, namely chytrids, which are the dominant fungal group in aquatic environments (Bahram et al. 2018) and are the key agent behind the worldwide amphibian decline (Alan Pounds et al. 2006). These fungi can have a broad ecological impact, as they alter aquatic ecosystem stability and functions by acting as a trophic link by driving energy flows to plankton and can regulate their outbreaks (Grami et al. 2011; Rasconi, Niqlund and Sime-Ngando 2012; Haraldsson et al. 2018). The zoospores of the parasitic chytrid associated with amphibians, Batrachochytrium dendrobatidis (Bd), are a rich food source for aquatic microfauna and protist grazers, who can control pathogen abundance and infection intensity (Schmeller et al. 2014). There is evidence that Bd can disperse from aquatic to terrestrial systems and retain similar functions (Kolby et al. 2015), and that non-amphibian animals can host and act as a reservoir for Bd (e.g. reptiles (Kilburn, Ibáñez and Green 2011), nematodes (Garmyn et al. 2012) and larval fish (Liew et al. 2017)). Furthermore, the hyperparasitism of parasitic chytrids by other fungi together with fungal-parasitized algae can transfer carbon to the detritus pool, where it can be grazed by protists and metazoans, leading to increased food-web stability in aquatic habitats (reviewed in Gleason et al. 2014; Grossart et al. 2019). Much remains to be discovered about the ecology and biogeography of aquatic fungi given the recent discovery of diverse clades (Jones et al. 2011; Grossart et al. 2019).

Fungal-microbial interactions

In addition to direct interactions with plant and animal hosts, fungi can indirectly interact with their hosts and affect ecosystem functions by altering microbial communities, owing to the often-shared resources and thereby interactions of fungi and
bacteria. Such cross-kingdom interactions have been proposed to greatly affect the structure and function of microbiomes with implications for host functioning and health (McFrederick, Mueller and James 2014; Bahram et al. 2018; Deveau et al. 2018; Durán et al. 2018; Bonfante, Venice and Lanfranco 2019; Netherway et al. 2023). For instance, interactions with bacteria may disproportionately affect the growth rate and ecosystem functioning of different fungal guilds (Hiscox et al. 2015; Bahram et al. 2018; Sterkenburg et al. 2018). Competition for nitrogen between EcM fungi and decomposers may lead to greater carbon storage (Averill, Turner and Finzi 2014; Sterkenburg et al. 2018), while stimulation of decomposers via root and mycorrhizal carbon rich exudates may lead to lower carbon storage (Lang et al. 2021). In turn, bacteria can utilize fungal-derived substrates following the decomposition of the more recalcitrant organic matter such as cellulose and lignin (Boer et al. 2005). Based on mapping metabolic interactions using advanced omics methods (e.g. metabolomics; the comprehensive quantitative analysis of all metabolites in a biological system) on the spatiotemporal dynamics of the kefir community, a recent study suggests that metabolic cooperation (i.e. cross-feeding) and spatiotemporal niche partitioning between yeasts and bacteria promote their coexistence in using resources (Blasche et al. 2021). Also, fungal hyphae can act as a dispersal vector, as well as a source of water and nutrients for bacteria (Warminik, Nazir and Elzas 2009; Furuno et al. 2010; Worrich et al. 2017), and thus potentially affect the composition and diversity of bacterial communities. Via transfer of water and nutrients across their mycelial networks, filamentous fungi may be able to better cope with environmental changes than bacteria (Whiteside et al. 2019) and thereby contribute to the stability of bacterial communities. There is also growing evidence that fungi can alter soil bacterial communities through direct competition over resources or production of secondary metabolites (McFrederick, Mueller and James 2014; Bahram et al. 2018; Pierce et al. 2021). In the gut environment—despite acting as a pathogen under certain circumstances—Candida albicans has been shown to act as a mutualistic fungus by inducing antibody production against other fungal infections (Doron et al. 2021). Fungal-associated bacteria may, in turn, have several beneficial effects on fungi and potentially their host plants or animals. Endosymbiotic bacteria in plant symbiotic fungi (Bonfante and Desiro 2017) could contribute to increased fitness of the fungus (Salvioli et al. 2016). These bacteria play an important role in the co-evolution and functioning of host-fungal systems by facilitating the establishment of symbiotic relationships (e.g. as mycorrhizal helpers) as well as the exchange of genetic material and nutrients between them (reviewed in Bonfante, Venice and Lanfranco 2019). A recent study shows that by using anthelmintic metabolites, endosymbiotic bacteria (Mycocevidus) of a plant growth–promoting fungal species (Mortierella sp.) protect the fungus against nematode attack (Bu’ttner et al. 2021). Insect-fungal mutualistic interactions may also involve bacteria as a third partner. For example, in a beetle-fungal mutualism where a fungus (Entomocorticium sp.) provides food resources for southern pine beetles (Dendroctonus frontalis), bacteria protect the beetle’s mutualistic fungus from an antagonistic fungus (Ophiostoma minus) by producing antimicrobials (Scott et al. 2008).

Fungal-viral interactions

Although less characterized, there is also some evidence for fungal–viral interactions in non-fungal hosts. For instance, pulmonary fungal pathogens can impair immune responses to viral infection in humans (Seelbinder et al. 2020), while they can suppress a viral disease (tomato yellow leaf curl virus) in tomato plants (Sun et al. 2021). Other studies also report correlations between the severity or clinical symptoms of viral diseases and fungal diversity, such as in the gut of chronic hepatits B patients (Chen et al. 2011) as well as the gut and oral samples of COVID-19 patients (Lv et al. 2021; Soffrìtti et al. 2021). Furthermore, viruses may use fungal mycelial networks to transfer across host systems, as evidenced by the existence of mycoviruses in various tissues of saprotrophic/pathogenic Armillaria species (Linnakoski et al. 2021), between neighbouring EcM fungal clones (Sutela and Vainio 2020), and possibly between different fungal species from different ecological guilds, i.e. saprotrophic, pathogenic and mycorrhizal, within the same forest stand (Vainio et al. 2017). Given the strong interactions between bacteria and fungi in various habitats and through endophytic associations, it is also plausible that fungi interact with bacteriophages. This has at least been shown between the commonly co-occurring human pathogens Pseudomonas aeruginosa (a bacterium) and Aspergillus fumigatus (a fungus), where a bacteriophage that infects P. aeruginosa was found to attach to the surface of and inhibit the metabolic activity of A. fumigatus—through the bacteriophages ability to sequester iron and render it unavailable to the fungus—thus potentially reducing the co-infection of P. aeruginosa and A. fumigatus (Penner et al. 2016). Clearly the interaction of fungi and viruses is an exciting and emerging avenue of research that will provide novel insights into the functioning of biological and ecological systems.

Finally, given that fungi frequently interact with one another, exemplified by the competition and antagonism between different fungal guilds (Rodriguez Estrada et al. 2012; Mukherjee et al. 2014; Fernandez and Kennedy 2016; Thakur et al. 2019) as well as within and between species from the same guild (Yan et al. 2015), there could be multiple fungal species simultaneously involved and interacting in symbiosis or parasitism with a host. Evidence for this comes from the lichen symbiosis where two unrelated fungi ubiquitously associate with algae (Spiribille et al. 2016), or from trees capable of hosting both AM and EcM fungal symbionts on the same section of roots (Teste, Jones and Dickie 2020). There is also evidence that pathogenic fungi facilitate each other in infecting hosts, e.g. by providing an adhesion site and promoting co-colonization of vaginal and oral tissues by different Candida species (Alves et al. 2014; Tati et al. 2016). Another example is hyperbiotrophy in fungi, where parasitic fungi of fungi (mycoparasites) boost the virulence of pathogenic fungi to exploit them as a conduit to extract nutrients from their common hosts (Laur et al. 2018).

Taken together, there appears to be a dynamic involvement of multiple species in the pathogenic and symbiotic interactions of fungi with other organisms, with important implications for the health of host animals and plant, as well as ecosystem functioning.

Fungal Interactions in a Changing Environment

Given the great versatility of biotic interactions involving fungi as discussed above, it is unclear exactly how global change factors such as warming and altered precipitation may influence the functioning of these interactions. It is probable that such environmental changes may have important consequences for ecosystem functions such as carbon storage (Clemmensen et al. 2013; Averill and Hawkes 2016) as well as plant responses to
biotic stress, which can be contingent on their associated fungi (Bennett and Bever 2007). For example, through mycorrhizal networks, plants can alert their neighbouring plants about herbivory attacks (Babikova et al. 2013), possibly resulting in a less severe response to herbivore attacks with increasing mycorrhizal colonization (Bacht et al. 2019; Sveen et al. 2021). Another example is the tripartite symbiosis between plant-fungus-virus, where plant-fungal symbioses mediated by fungal viruses can confer heat resistance in plants and enable both interacting partners to grow under elevated temperatures (Márquez et al. 2007). Climate warming has been shown to accelerate fungal infection rates, which may alter the dynamics of the food-web structure in aquatic environments (Frenken et al. 2016).

Both climatic and human-induced disturbances may facilitate the emergence of invasive fungi, partly owing to the versatility of fungal interactions and ecology. Fungal invasions can cause major economic losses, with important ecological consequences. This is exemplified by recent shifts in the distributional ranges of several pathogenic fungi towards higher latitudes, posing a threat to production systems in colder climates (Bebber, Ramotowski and Gurr 2013). In addition, given that mycorrhizal fungi probably helped plants to colonize the land, it is not surprising that they still assist, if not play a primary role in facilitating plants in colonizing novel geographic areas (Pringle et al. 2009). For instance, the spread of introduced EcM tree species out of production landscapes and into natural landscapes is associated with both their interaction with co-introduced EcM fungi and through forming novel EcM associations with native EcM fungi (Vlk et al. 2020). Nevertheless, when introduced to new habitats, the lack of co-evolutionary histories may lead to contrasting interaction outcomes for some fungi. A notable example of this is Geomyces destructans that is a widespread commensal fungus in hibernating bats in its native range (Europe), while it causes white-nose syndrome in bats where the fungus is invasive (N. America) (Wibbelt et al. 2010). Furthermore, the versatility of fungal interactions may result in various ecological outcomes from a fungal invasion ranging from increased to reduced fitness, as has been shown in the case of the interaction involving powdery mildew Erysiphe alphioides with local insect herbivores (Tack, Gripenberg and Roslin 2012). Even when the same symbiosis type persists, the interaction efficiency may be altered when novel interactions are formed with invasive fungi. This is exemplified by the lowered diversity of AM fungi when an exotic AM fungus is introduced to the host plant (Koch et al. 2011). The ecological implications of newly formed interactions by invasive fungi warrants further investigation.

Moving to the health of humans, there is evidence that changes in gut fungal and bacterial communities can affect one another, likely due to their direct interactions. For example, the overgrowth of commensal Candida post-antibiotic treatment (Dollive et al. 2013) can prevent the recovery of bacterial communities resulting in long-term compositional shifts (Mason et al. 2012). Both fungal-bacterial and fungal-fungal interactions in healthy individuals are important to prevent transition of the commensal Candida to a pathogenic state (Limon, Skalski and Underhill 2017). Carbohydrate substrates of fungi play an important role in their fitness in the gut, as evidenced by increased abundance of Candida following the consumption of carbohydrate rich foods, whereas they respond negatively to saturated fatty acids (Hoffmann et al. 2013). In addition, by providing a food source (glycan) and a binding/attachment site, Candida have been suggested to facilitate recolonization of the gut Bacteroidetes during antibiotic recovery (Mason et al. 2012). Finally, given that human pathogens from clinical settings can exist in the environment as soil (Bensasson et al. 2019; Opulente et al. 2019; O’Brien et al. 2021), it remains to be seen how changes in the environment can affect the evolution of their virulence and resistance traits as well as transmission to humans. By considering a wider range of fungal traits and their role in biotic interactions (Fig. 2), we may be able to better address such questions.

Overall, evidence suggests that while certain fungal interactions may shift in response to environmental changes, leading to altered host and ecosystem functions, some fungi may play a mediating role for their hosts and ecosystems in coping with or succumbing to environmental stress.

CONCLUSIONS AND FUTURE DIRECTIONS

Unique combinations of fungal traits such as diverse dispersal mechanisms as well as versatile genomic, lifestyle and ecophysiological traits could have contributed to their success in spanning multiple scales and dimensions of time, space, and biological interactions, that is not rivalled by other organism groups (Figs. 1, 2). In addition, fungi ubiquitously form complex multikingdom and multispecies interactions, with important implications for ecosystem and host functioning. Several lines of evidence also suggest that fungi act as mediating links between a wide range of organisms and ecosystems, and that both symbioses, parasitism, and indirect interactions have strongly influenced the evolution and distribution patterns of fungi and their interacting taxa (Fig. 3). We conclude that the versatility, complex interactions, and mediating role of fungi further underscores their importance within and across various ecosystems. Yet, the exact underlying mechanisms of the role of fungi in many systems warrant further investigation. In this respect, the key questions that need to be addressed are: How are interactions between fungi with other organisms established? How does evolutionary history influence these interactions and what functional traits facilitate them? How do fungal interactions with other organisms persist over time and space and what conditions facilitate these interactions? What are the relative roles of bacteria and viruses—including HGT and HVT—in fungal interactions with eukaryotic hosts? Further field and controlled experiments are needed to examine to what extent the environment and spatiotemporal variation in fungal communities contributes to the dynamics of fungal interactions. Deciphering links between fungal diversity, fungal and host functional traits and their interactions will not only improve our understanding of the resilience of ecosystems against environmental perturbation, but also help predict future responses to global change including potential range expansions of both fungi and their interacting partners.

Studies of fungal ecology and biogeography, and inferred interactions, have been greatly facilitated by recently developed tools and databases for assigning fungal traits (Nguyen et al. 2016; Pölme et al. 2020; Zanne et al. 2020). Yet, we still need tools and analytical approaches that account for the flexibility in fungal lifestyles and interactions, embrace the complexity of fungi and their influence in ecosystems, and which can reveal realized interactions in mixed fungal communities (Fig. 2). Recent developments in single cell microbial genomics and droplet microfluidics hold great promise for unravelling microbe–microbe and microbe–environment interactions, including communication, competition, cooperation, and food-web dynamics and formation in constructed and controlled micro ecosystems (Nagy et al. 2018). Although these methods are not currently well-suited for
studying filamentous fungi and macro eukaryotes due to technical limitations (Millet et al. 2019), there is great potential to adapt these approaches to build custom made systems that mimic small scale heterogeneous soil environments and also microcosms involving seedlings and root symbionts (Aleklett et al. 2018). Recent studies have made exciting progress in this regard, where co-culturing of a plant-associated fungus and bacterium isolated from a host was combined with a novel microfluidics system, comparative metabolomics and analytical methods to identify events of metabolic exchange that initiate a symbiosis between these organisms (Uehling et al. 2019; Büttner et al. 2021).

Genomics data combined with transcriptomics and epigenomics approaches (to study all expressed genes and epigenetic factors in environmental samples, respectively) can provide unique opportunities in the next few years for studying fungal interactions at an unprecedented scale and resolution. Yet, perhaps of key interest to study fungal activity and interactions with hosts, viruses, microbes and ecosystems is metabolomics. In this respect, unravelling metabolic interactions by integrating different omics methods (metagenomics, transcriptomics and metabolomics) and mathematical modelling may be a promising approach to relate species-species interactions, genotypes and phenotypes and to gain mechanistic insights into fungal interactions (Blasche et al. 2021). Increasing genetic, transcriptomic and genomic studies on molecular host-fungus interactions and biomolecule degradation pathways provide comprehensive characterisation of functional gene families and activity centres of enzymes, transporters, signalling molecules and transposable elements.

By moving towards the study of the phenotypic expression of fungi, host systems and their interactions, and ecosystems, can we begin to collect cause and effect information on fungal interactions and functional outcomes. By exploring fungi beyond traditional disciplinary and ecosystem boundaries, can we further unravel how communities of fungi and their interacting organisms may shift in response to changes in environmental conditions and how this may affect host and ecosystem functioning and health.

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