

This is a self-archived version of an original article. This version may differ from the original in pagination and typographic details.

Author(s): Aguilar-Trigueros, Carlos A.; Krah, Franz-Sebastian; Cornwell, William K.; Zanne, Amy E.; Abrego, Nerea; Anderson, Ian C.; Andrew, Carrie J.; Baldrian, Petr; Bässler, Claus; Bissett, Andrew; Chaudhary, V. Bala; Chen, Baodong; Chen, Yongliang; Delgado-Baquerizo, Manuel; Deveautour, Coline; Egidi, Eleonora; Flores-Moreno, Habacuc; Golan, Jacob; Heilmann-Clausen, Jacob; Hempel,

Title: Symbiotic status alters fungal eco-evolutionary offspring trajectories

Year: 2023

Version: Published version

Copyright: © 2023 the Authors

Rights: CC BY-NC 4.0

Rights url: https://creativecommons.org/licenses/by-nc/4.0/

Please cite the original version:

Aguilar-Trigueros, C. A., Krah, F., Cornwell, W. K., Zanne, A. E., Abrego, N., Anderson, I. C., Andrew, C. J., Baldrian, P., Bässler, C., Bissett, A., Chaudhary, V. B., Chen, B., Chen, Y., Delgado-Baquerizo, M., Deveautour, C., Egidi, E., Flores-Moreno, H., Golan, J., Heilmann-Clausen, J., . . . Powell, J. R. (2023). Symbiotic status alters fungal eco-evolutionary offspring trajectories. Ecology Letters, 26(9), 1523-1534. https://doi.org/10.1111/ele.14271

LETTER



Symbiotic status alters fungal eco-evolutionary offspring trajectories

Carlos A. Aguilar-Trigueros^{1,2,3,4} | Franz-Sebastian Krah⁵ | William K. Cornwell⁶ |

Amy E. Zanne⁷ | Nerea Abrego^{3,8} | Ian C. Anderson⁴ | Carrie J. Andrew⁹ |

Petr Baldrian¹⁰ | Claus Bässler⁵ | Andrew Bissett¹¹ | V. Bala Chaudhary¹² |

Baodong Chen^{13,14} | Yongliang Chen¹⁵ | Manuel Delgado-Baquerizo^{16,17} | |

Coline Deveautour¹⁸ | Eleonora Egidi⁴ | Habacuc Flores-Moreno⁷ | Jacob Golan¹⁹ |

Jacob Heilmann-Clausen²⁰ | Stefan Hempel¹ | Yajun Hu²¹ | Håvard Kauserud²² |

Stephanie N. Kivlin²³ | Petr Kohout¹⁰ | Daniel R. Lammel¹ | Fernando T. Maestre^{24,25} |

Anne Pringle¹⁹ | Jenna Purhonen^{3,26,27} | Brajesh K. Singh^{4,28} | Stavros D. Veresoglou¹ | Tomáš Větrovský¹⁰ | Haiyang Zhang^{4,29} | Matthias C. Rillig^{1,2} | Jeff R. Powell⁴ |

Matthias C. Rillig and Jeff R. Powell joint senior authorship.

This is an open access article under the terms of the Creative Commons Attribution-NonCommercial License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited and is not used for commercial purposes.

© 2023 The Authors. *Ecology Letters* published by John Wiley & Sons Ltd.

Ecology Letters. 2023;00:1–12. wileyonlinelibrary.com/journal/ele 1

¹Institute of Biology, Freie Universität Berlin, Berlin, Germany

²Berlin-Brandenburg Institute of Advanced Biodiversity Research (BBIB), Berlin, Germany

³Department of Biological and Environmental Science, University of Jyväskylä, Jyvaskyla, Finland

⁴Hawkesbury Institute for the Environment, Western Sydney University, Penrith, New South Wales, Australia

⁵Faculty of Biological Sciences, Department of Conservation Biology, Institute for Ecology, Evolution and Diversity, Goethe University Frankfurt, Frankfurt am Main, Germany

⁶Evolution & Ecology Research Center, School of Biological, Earth, and Environmental Sciences, University of New South Wales, Sydney, New South Wales, Australia

⁷Department of Biology, University of Miami, Coral Gables, Florida, USA

⁸Department of Agricultural Sciences, University of Helsinki, Helsinki, Finland

⁹Biology Department, Oberlin College & Conservatory, Oberlin, Ohio, USA

¹⁰Laboratory of Environmental Microbiology, Institute of Microbiology of the Czech Academy of Sciences, Praha 4, Czech Republic

¹¹Oceans and Atmosphere, CSIRO, Hobart, Tasmania, Australia

¹²Department of Environmental Studies, Dartmouth College, Hanover, New Hampshire, USA

¹³State Key Laboratory of Urban and Regional Ecology, Research Center for Eco-Environmental Sciences, Chinese Academy of Sciences, Beijing, People's Republic of China

¹⁴University of Chinese Academy of Sciences, Beijing, People's Republic of China

¹⁵College of Resources and Environmental Sciences, China Agricultural University, Beijing, People's Republic of China

¹⁶ Laboratorio de Biodiversidad y Funcionamiento Ecosistémico. Instituto de Recursos Naturales y Agrobiología de Sevilla (IRNAS), CSIC, Sevilla, Spain

¹⁷Unidad Asociada CSIC-UPO (BioFun). Universidad Pablo de Olavide, Sevilla, Spain

¹⁸AGHYLE Research Unit, Institut Polytechnique UniLaSalle, Mont-Saint-Aignan, France

¹⁹Departments of Botany and Bacteriology, University of Wisconsin-Madison, Madison, Wisconsin, USA

²⁰Center for Macroecology, Evolution and Climate, GLOBE Institute, University of Copenhagen, Copenhagen, Denmark

²¹Key Laboratory of Agro-ecological Processes in Subtropical Region & Changsha Research Station for Agricultural and Environmental Monitoring, Institute of Subtropical Agriculture, Chinese Academy of Sciences, Hunan, China

²²Evogene, Department of Biosciences, University of Oslo, Oslo, Norway

²³Department of Ecology and Evolution, University of Tennessee, Knoxville, Tennessee, USA

²⁴Instituto Multidisciplinar para el Estudio del Medio "Ramon Margalef", Universidad de Alicante, Carretera de San Vicente del Raspeig s/n, Alicante, Spain

²⁵Departamento de Ecología, Universidad de Alicante, Carretera de San Vicente del Raspeig s/n, Alicante, Spain

Correspondence

Carlos A. Aguilar-Trigueros, Institute of Biology, Freie Universität Berlin, Altensteinstrasse 6, 14195 Berlin, Germany. Email: calgit@gmail.com

Funding information

Alexander von Humboldt-Stiftung, Grant/Award Number: Feodor-Lynen Fellowship; Australian Research Council, Grant/Award Number: DP190103714 and FT0100590; Bundesministerium für Bildung und Forschung, Grant/ Award Number: 01LC1501A: Deutsche Forschungsgemeinschaft, Grant/ Award Number: HE6183: Deutscher Akademischer Austauschdienst; Division of Environmental Biology, Grant/Award Number: 1623040 and 1655759; Grantová Agentura České Republiky, Grant/Award Number: 21-17749S; H2020 European Research Council, Grant/Award Number: 647038 and 694368: Universities Australia

Editor: Dustin John Marshall

Abstract

Despite host-fungal symbiotic interactions being ubiquitous in all ecosystems, understanding how symbiosis has shaped the ecology and evolution of fungal spores that are involved in dispersal and colonization of their hosts has been ignored in life-history studies. We assembled a spore morphology database covering over 26,000 species of free-living to symbiotic fungi of plants, insects and humans and found more than eight orders of variation in spore size. Evolutionary transitions in symbiotic status correlated with shifts in spore size, but the strength of this effect varied widely among phyla. Symbiotic status explained more variation than climatic variables in the current distribution of spore sizes of plant-associated fungi at a global scale while the dispersal potential of their spores is more restricted compared to free-living fungi. Our work advances life-history theory by highlighting how the interaction between symbiosis and offspring morphology shapes the reproductive and dispersal strategies among living forms.

KEYWORDS

functional ecology, fungi, life-history, offspring size, symbiosis

INTRODUCTION

In life-history theory, the ecology and evolution of offspring size is linked to environmental factors that species encounter during reproduction, dispersal and earlycolonization, as well as physiological constraints during their development (Stearns, 1992). By providing a common framework where disparate offspring structures such as plant seeds (Moles et al., 2005), animal ovules (Neuheimer et al., 2015), avian eggs (Stoddard et al., 2017) and mammal size at weaning (Falster et al., 2008) can be compared, life-history theory aims at discovering general principles behind the drivers shaping the ecology and evolution of species at earlier stages of their life cycle. According to life-history theory, alterations in the size of offspring structures of sessile organisms can influence their potential for both dispersal (moving to a new habitat) and colonization (establishing a new individual) (Falster et al., 2008; Kavanagh & Burns, 2014). The size of the offspring structure could impact how far

it can disperse and its capacity to withstand environmental conditions during dispersal. Conversely, change in size alter the amount of resources that can be packed into offspring structures, resources which can facilitate germination and earlier stages of development that ultimately lead to successful colonization and the formation of a new functional individual.

However, most life forms that have been used to develop this knowledge are free-living macroorganisms, ignoring the large diversity of microbial forms. Conspicuously absent is the Kingdom Fungi, which, with 136,000 described species and an estimated diversity of 3–10 million species (Hawksworth & Lücking, 2017), is a large portion of the tree of life. This dearth represents a fundamental knowledge gap because, as we report here (Figure 1), variation in fungal offspring size (up to eight orders of magnitude) is as high or higher than that of macroorganisms whose comparative offspring studies (e.g. plant seeds and avian eggs) dominate the life-history theory (Figure 1). Here, we use fungal spores as offspring

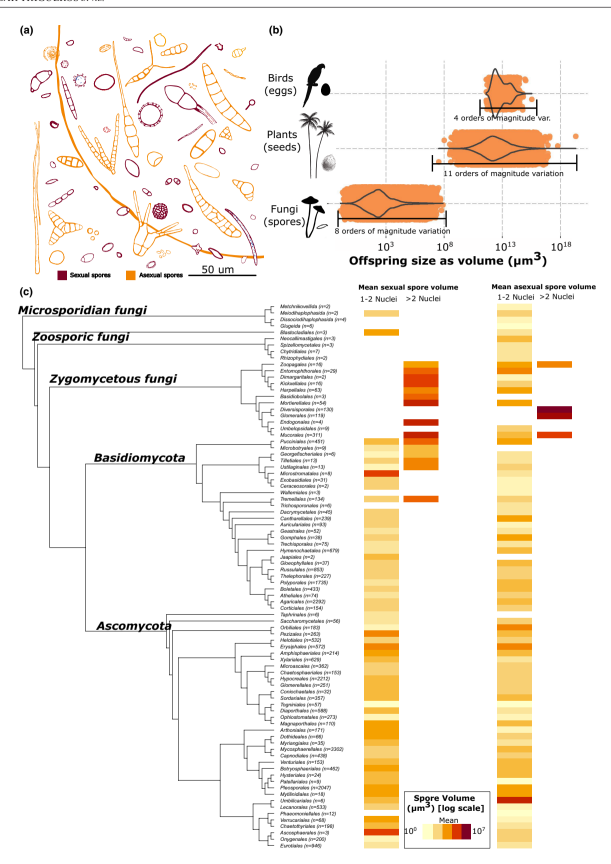
FIGURE 1 Interspecific variation in spore size and symbiotic lifestyles across the fungal kingdom: (a) Illustration of the diversity of shapes and sizes among all fungal sexual and asexual spore types. (b) Interspecific spore size variation is more than eight orders of magnitude across the kingdom, ranging from the mitospores of *Phoma muscivora* of $9.0 \times 10^{-2} \, \mu m^3$ to multinucleate spores of the mycorrhizal fungus *Scutellospora scutata* of $7.8 \times 10^7 \, \mu m^3$. This variation is comparable to that of other offspring structures such as angiosperm seeds and bird eggs (to aid comparison, all offspring structures are presented on the same scale [μm^3]). (c) Phylogenetic tree with terminal branches representing orders (the number of species per order for which we collected spore data is given in parenthesis). The corresponding heatmap displays order averages (in logarithmic scale) of spore size as volume in yellow-to-red colour scale for sexual and asexual spores separating spores types based on the number of nuclei, which is a major distinction in spore types for fungi (see main text and supplementary material for a detailed explanation on descriptions of the biology of these distinct spores). Fungal spores (n=26, 134 species), avian egg data (n=1395 species) were obtained from n=1000, while seed data (n=34,390 species) were obtained from the seed database of Kew Botanical Garden (http://data.kew.org/sid/?_ga=2.73581 714.1287366807.1501084977-1309187973.1501084964).

²⁶Department of Music, Art and Culture Studies, University of Jyväskylä, Jyvaskyla, Finland

²⁷School of Resource Wisdom, University of Jyväskylä, Jyvaskyla, Finland

²⁸Global Centre for Land Based Innovation, Western Sydney University, Penrith, New South Wales, Australia

²⁹College of Life Sciences, Hebei University, Baoding, China



structures (Figure 1b) because they represent reproductive output units produced by a mature mycelium (the 'parent' fungus) that function as dispersal propagules to colonize novel habitats that are usually distantly located

from the parental fungus. Each spore has the potential to develop into a new mycelium, which is independent from the parental one in the new habitat. Spore traits, such as total size, are hypothesized to determine the

likelihood of colonization based on spore interactions with their environment during their release, movement, attachment/landing, dormancy and germination (Golan & Pringle, 2017; Halbwachs, 2015; Kivlin, 2020; Rockett & Kramer, 1974). Thus, spores are functionally analogous to dispersal offspring propagules of other sessile modular organisms like plant seeds (Moles et al., 2005) or marine invertebrate eggs (Neuheimer et al., 2015).

We hypothesize that symbiotic status of fungal species may explain this large variation in spore size because of the contrasting conditions that spores encounter when germinating and colonizing dead organic matter versus living hosts. That is, most extant fungi can be placed along a symbiotic spectrum spanning from asymbiotic species whose spores start the colonization of organic matter substrates, to a variety of symbiotic interactions whose spores initiate the colonization (i.e. infection) in hosts in almost all major domains of life (Lutzoni et al., 2018; Naranjo-Ortiz & Gabaldon, 2019) (Figure 2; Figures S3 and S4). Furthermore, shifts between freeliving to symbiotic life styles have been a major driver in trait evolution in the fungal kingdom (Lutzoni et al., 2018; Naranjo-Ortiz & Gabaldon, 2019) raising the question of how transitions in symbiotic status influence the ecology and evolution of spores. Here, we use a definition of symbiosis that is common in evolutionary biology: the intimate physical living together of distinct species (usually distantly related), whether mutualistic, parasitic, or commensal, including macrobe-microbe interactions, where the former is considered the 'host' and the latter the 'symbiont' (Chomicki et al., 2020).

Specifically, to understand the importance of symbiotic status in explaining differences in the size and function of spores across the fungal kingdom, we asked three questions. First, are transitions in symbiotic status correlated with shifts in spore size? To answer this question, we used linear phylogenetic regression to test whether the spore size of symbiotic groups (e.g. insect pathogens, plant pathogens, ectomycorrhizal) shift in size (i.e. increase or decrease) compared to asymbiotic fungi across all major fungal phyla. We then focused on plant-associated fungi in the Dikarya clade to test whether symbiotic groups of obligate lifestyles have larger offspring than symbiotic groups with facultative lifestyles. We focus on fungi associated to plants in this clade because plants are by far the host type with the largest diversification of symbiotic lifestyles (Lutzoni et al., 2018). In addition, this hypothesis has been repeatedly used to explain why the spores of some obligate plant pathogenic and mutualistic fungi are so large. This hypothesis posits that obligate symbionts may benefit from the greater resources present in large spores, since these resources represent the only means of surviving during dispersal and initial colonization (i.e. infection) of new hosts until resources can be exchanged with the host plant (Garrett, 1970; McLaughlin & Spatafora, 2014). Second, we asked whether spore size distribution across fungal communities at a global scale

can be explained by climate variables, regardless of symbiotic status. We hypothesized that climate may be more important than symbiotic status in explaining spore size distributions given the worldwide distribution of fungi spanning diverse climatic zones. As both asymbiotic and plant-associated fungal species release and disperse their spores into the abiotic environment, climate may act as a key driver of spore size variation (Kendrick, 2017). In addition, based on predictions from life-history theory, we expect species with larger offspring sizes to be associated with limiting environmental regimes (Moles & Westoby, 2004). Third, we asked whether the dispersal potential of spores differs between asymbiotic and plantassociated fungi. Specifically, we tested whether species with smaller spores have a broader geographic distribution (i.e. higher extent of occurrence) and whether this relationship varies between asymbiotic and plantassociated fungi. One of the main ecological functions of offspring is dispersal and, for several fungal groups, it has been proposed that small offspring should travel farther than large offspring, increasing the dispersal potential of species (Norros et al., 2014). However, if plantassociated fungi require large spore sizes, they may have more-limited distributions than asymbiotic fungi.

MATERIALS AND METHODS

Assembly of spore database

Unlike macroorganisms such as plants and animals, no databases of offspring morphology for fungi exist. Therefore, to answer our questions, we created and populated a new database by text mining nearly 100,000 taxonomic descriptions deposited in Mycobank (Robert et al., 2013) (http://www.mycobank.org/; see Supplementary material for further details). In total, we collected information on spore width and length dimensions for >26,000 accepted species (based on taxonomy from the Catalogue of Life; https://www.catalogueoflife. org/), representing 20% of all described fungal species (Figure S1). This database includes spore-dimension data from both sexually and asexually produced spores across major fungal lineages at different stages of fungal life cycles. However, we restricted the analysis described below to sexual spore types described as 'ascospores' and 'basidiospores' (henceforth referred to as 'sexual spores') and asexual spore types described as 'conidia' and 'sporangiospores' (henceforth referred to as 'asexual spores') because they represent the most frequently occurring types of spores in our dataset and thus can be compared across several fungal lineages and symbiotic groups (Figure 1c; Figure S2). We also excluded spores of glomeromycete fungi for our main analyses because their extreme large size may bias the results (see Figure S2 and supplementary material for specific spore definitions and nomenclature used in the analysis; Kendrick, 2017). We

AGUILAR-TRIGUEROS ET AL.

then calculated spore volume using width and length as a proxy for spore size following the formula for a prolate spheroid (Aguilar-Trigueros et al., 2019). We used volume as a proxy for size because it captures the 3D structure of fungal spores and, based on allometric theory, volume scales with other measurements of size, such as weight. Indeed, volume has been used in life-history research as a proxy for offspring size across several large clades (Stearns, 1992). Using this approach, we found that spore size across species varied by more than eight orders of magnitude (Figure 1b).

Assembly of symbiotic status data

We also assembled a symbiotic status database (by mining and crosschecking different functional databases) where fungal species were categorized as asymbiotic (i.e. saprotrophic species that have only been reported as free-living during their whole life cycle) or symbiotic with a wide diversity of host and types of interactions as follows: (1) Asymbiotic saprotrophs; (2) Insect pathogens; (3) Lichen fungi; (4) Plant endophytes; (5) Plant pathogenic necrotrophs; (6) Plant pathogenic biotrophs; (7) Ectomycorrhizal fungi; (8) Arbuscular Mycorrhizal fungi; (10) Human pathogens. For plant symbiotic fungi, we further classify the level of specialization as either facultative symbiosis (species that are reported to alternate between a free-living and symbiotic phase) or obligate symbiosis (species that have been exclusively reported as symbiotic to complete their life cycle) based on the biology of their respective symbiotic group (see supplementary material for data sources and details on the criteria used to define symbiotic groups).

Phylogenetic regression of shifts in spore size and evolutionary transitions in symbiotic groups

Because evolutionary history shapes how and where species are today, the role of this history can be examined by testing how traits shift across the tree of life. For fungal spores, recent reviews and anecdotal evidence suggest that spore size is expected to differ more widely in some fungal clades than others (Aguilar-Trigueros et al., 2019; Ingold, 2001). Thus, we used two phylogenies to test whether transitions in symbiotic status correlate with shifts in fungal spore size. The first phylogeny consists of 1644 fungal species whose genome has been fully sequenced as recently published in (Li et al., 2021). Focusing our analysis on these species allowed us to incorporate the most robust, species-level phylogenetic tree available to date for fungi (as this tree is based on whole genome data) that captures the entire kingdom (i.e. it is not specific to only a subset of fungal clades). However, because this tree only includes a limited number of species, we

also used a taxonomy-based phylogeny consisting of 23,000 species from which we obtained taxonomic data from phylum-to-species level using the function *as.phylo* from ape package (Paradis & Schliep, 2018) (see further details in the supplementary material for the construction of this tree).

Independently for each phylum or fungal group (i.e. Ascomycota, Basidiomycota, zygomycetous fungi, zoosporic fungi and microsporidan fungi), we conducted phylogenetic linear regression models where the logarithm of spore volume was the response variable, symbiotic status (based on the 10 symbiotic guilds classified here) was the explanatory variable and either the genome-based phylogenetic tree from (Li et al., 2021) or the taxonomy derived cladogram was used to account for phylogenetic relatedness. These phylogenetic regressions were conducted on sexual spores of the Ascomycota and Basidiomycota (i.e. ascospores and basidiospores) and asexual spores of all phyla and fungal groups (as defined above, we only include asexual spores referred as 'conidia' or 'sporangiospores') separately as they represent two separate traits under different selection. These phylogenetic linear regression models were conducted using the function phylolm from the phylolm package (Ho et al., 2016).

We conducted additional phylogenetic regression models testing whether spore size is bigger for obligate symbionts compared to facultative symbionts for sexual spores of plant-associated fungi in the Ascomycota and Basidiomycota and asexual spores of the Ascomycota. As above, this phylogenetic regressions were performed the genome-based phylogenetic tree (Li et al., 2021) or the taxonomy-based cladogram.

Relative importance of symbiotic status against climate variables in explaining spore size variation across communities

To obtain climatic information, we first mapped the geographic distributions of fungal species observed in several large-scale, high-throughput DNA-sequencing studies of fungal communities from soil and plant samples covering an extensive breadth of biomes and occurring on all seven continents (Figure S5; see supplementary material section for details on how species annotations were performed). Then, we collected climatic data associated with the locations where those species were found, estimated mean values for each species, and compared the ability of those climate variables and each species' symbiotic status to explain variation in spore size.

For this analysis, we focused on species that in our database are reported to produce only one spore type because it is not possible to determine the spore type associated with environmental DNA sequences. We assessed the importance of fungal symbiotic status (i.e. whether

fungi are free-living saprotroph or plant-associated) in explaining interspecific variation in spore size relative to other drivers, including spore type (i.e. sexual and asexual) and climate across communities worldwide. Phylogenetic linear regression models were fit using the following predictors: spore type (categorical variable), climate (averages of mean annual temperature and precipitation, temperature and precipitation seasonality, maximum solar radiation and minimum water-vapour pressure calculated across locations in which each species was detected; as continuous variables) and symbiotic status (as a categorical variable—free-living or plantassociated). As before, we conducted this analysis using two phylogenetic regression (one using the genome-based phylogenetic tree from (Li et al., 2021) and the other one using the taxonomy-based tree).

Differences between saprotrophic and plantassociated fungi in the relationship between spore size and geographic spread

We assessed the role that fungal lifestyle plays in determining the relationship between geographic range and spore size. As with the previous analysis, we focused on species that in our database are reported to produce only one spore type because it is not possible to determine the spore type associated with environmental DNA sequences. To do this, we estimated species' geographic ranges from their mapped distributions in environmental DNA-sequencing studies. Specifically, geographic range for each species was estimated in two ways: (1) as the maximum distance in meters between samples, in which the species was detected using the ellipsoid method (Vincenty, 1975) calculated with the distVincentyEllipsoid function from the 'geosphere' package in R (Hijmans et al., 2019); and (2) as the range area in square meters using alpha-hull-derived measures (Edelsbrunner et al., 1983) incorporating all samples in which the species was detected using the getDynamicAlphaHull function from the 'rangeBuilder' package in R (Davis Rabosky et al., 2016). Each estimate of range size was then used as a response variable in linear models to estimate slopes representing the strength of the relationship between geographic range and spore volume for fungi with saprotrophic lifestyles (free-living) and those from plant-associated lifestyles (symbiotic). These models included random intercepts representing the taxonomic order (to account for non-independence among fungal species) and the primer set used to amplify fungal DNA (to account for biases among primer sets in their ability to detect fungal species). Because point-estimates can be sensitive to unbalanced sampling designs and, therefore, are unreliable, we used functions in the 'lme4' (Bates et al., 2015) and 'brms' (Bürkner, 2017) packages in R to fit Bayesian models and estimate posterior distributions of the slope parameters and calculated 95%-credible

intervals from four MCMC chains (each 2000 iterations with a 1000-iteration burn-in) to assess differences among fungal lifestyles. To assess relationships within individual orders, separate linear mixed-effects models were also fitted for each combination of taxonomic order and fungal lifestyle for which a minimum of five species with spore volume and geographic extent were available.

All statistical analyses were performed using R version 4.0.1 (Team, 2020). Spore volume was log10-transformed prior to statistical analyses.

RESULTS

Strong differences in spore size among fungal clades

Spore size variation among sexual and asexual spores was strongly structured by species' evolutionary history (Figure 1c, Tables S1 and S2). For instance, the asexual spores from the glomeromycetes are the largest in the kingdom, from 1.5-to-4 orders of magnitude larger compared to other spores (either sexual or asexual) from other groups and this difference shows strong phylogenetic structure (Pagel's lambda ~0.7 depending on the comparison, see Table S1). These spores of glomeromycetous fungi, however, are unique among other fungi because they contain hundreds of nuclei (an unparalleled feature in the kingdom (Kokkoris et al., 2020)), which might partly explain their extremely large size (Aguilar-Trigueros et al., 2019). Further, we found that sexual spores of basidiomycetes are on average 6 µm³ smaller than ascomycetes across the tree (Pagel's lambda=0.8, see Table S2). While this pattern alone cannot determine the mechanisms behind this size difference, it is consistent with the hypothesis that sexual spores of basidiomycetes are smaller than those of the ascomycetes because the Basidiomycota, as a whole, evolved a spore launching mechanism ('the surface tension catapult') that depends on spore size. In contrast, the launching mechanism of ascomycetes does not (Ingold, 2001; Roper & Seminara, 2019). This potential mechanism suggests that the size of the spore is dependent on the anatomy and morphology of the reproductive structure of the parental fungus. Such parent-to-offspring regulation has also been observed in other taxa, such as placental mammals, for whom size at birth depends on the anatomical constraints of the reproductive structure where the offspring develops (Stearns, 1992).

Correlation between transitions in symbiotic status and shifts in spore size vary among fungal clades

We found support for our hypothesis that evolutionary transitions in symbiotic status correlate with shifts

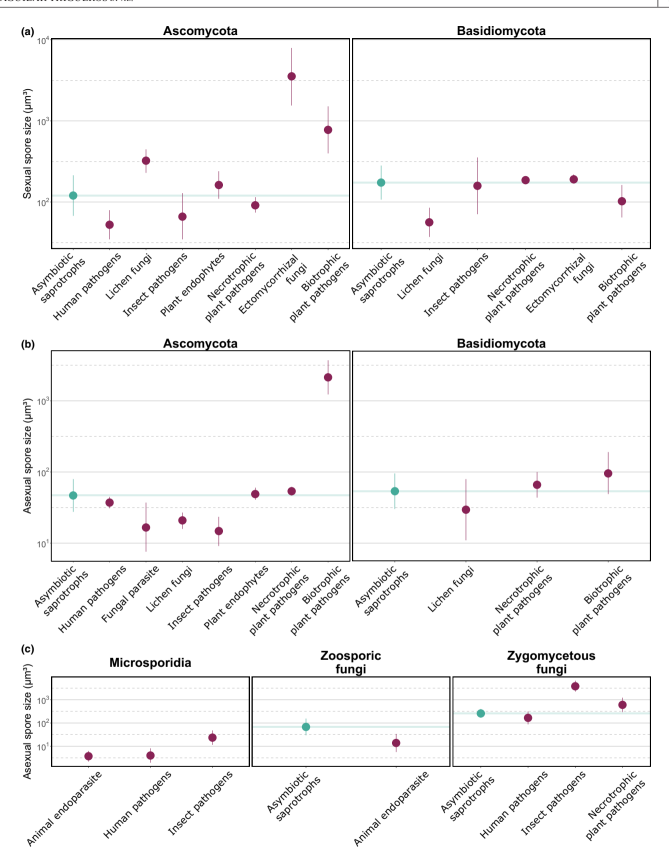


FIGURE 2 Shifts in size of sexual and asexual spores between asymbiotic fungi (blue points) and symbiotic groups (red points) across Dikarya (a, b) and non-Dikarya fungi (c). Point ranges show the predicted mean values (points) and associated standard errors (ranges) for each symbiotic group based on phylogenetic linear regression models using a taxonomy-based phylogeny. The horizontal blue lines are added to help comparison of asymbiotic and symbiotic fungi (except for Microsporidan fungi that have not recorded asymbiotic species). Similar results were obtained in models using a genome-based phylogeny (see Figure S2).

in the size of both sexual and asexual spores (Tables S3) and S4; Figure 2; Figure S3). However, the direction and strength of this correlation highly depended on the symbiotic group, spore type and phylum considered. We found that shifts in sexual spore size during transitions from saprotrophic to symbiotic groups were stronger in the Ascomycota compared to the Basidiomycota, specifically, we found shifts to larger spore sizes among insect pathogens, ectomycorrhizal, lichen and mildew fungi in the Ascomycota (although statistical support for the last two groups was found on only one phylogenetic regression; see Table S3). For asexual spores, we also observed stronger shifts of size and symbiotic status among groups in the Ascomycota compared to the Basidiomyota, although shifts in asexual spore sizes were more heterogeneous: shifts to larger asexual spores were associated with biotrophic and necrotrophic plant pathogens, while shifts to smaller asexual spore sizes were associated with lichen and insect pathogenic fungi (Table S4). Finally, we also detected shifts towards larger asexual spore sizes among insect and necrotrophic pathogens of zygomycetous fungi (Table S4) and for insect pathogens in the Microsporidia.

Among plant-associated fungi, we found a global trend towards increased sexual spore size in fungi with more obligate symbioses in the Ascomycota only (i.e. we found no statistical support for this hypothesis with plant-associated groups in the Basidiomycota). For sexual spores, plant obligate symbionts in the Ascomycota were about 29 µm³ larger than spores of facultative symbionts counterparts, while for asexual spores, obligate symbionts were up to 59 µm³ larger than spores of facultative symbionts (all p-values <0.001; Figure S4; Table S5). A possible mechanism behind large spores being associated with these groups is that spore reserves or thickening of spore cell walls increase chances of survival when dispersing to a host, overwintering and/or overcoming initial host resistance (e.g. penetration of the hard cuticle or the epidermal tissue) (Kemen & Jones, 2012; Wang & Wang, 2017).

Our results are congruent with previous research reporting small differences in spore size across functional groups in Basidiomycota fungi, particularly when comparing the sexual spores of ectomycorrhizal and saprotrophic fungi suggesting than other reproductive traits, such as sporocarp size and shape, might be more functional (Bässler et al., 2014; Calhim et al., 2018; Halbwachs et al., 2017; Kauserud et al., 2008). As we show here, this small difference might be due to the already small size of sexual spores of basidiomycete fungi relative to ascomycete fungi, which prompts the hypothesis that for the Basidiomycota the demand for small spores for the launching platform leaves little room for differentiation during evolution of the symbiotic lifestyle. In the case of necrotrophic pathogens or plant endophytes, the overlap in spore sizes with asymbiotic fungi and their relative large variation in sizes (Figure 2; Figure S3) may reflect differences in the level of symbiotic specialization

(Mengiste, 2012) that is not captured with the current classification. Plant pathologists have long speculated that larger spores may provide the necessary resources for highly host-specialized necrotrophs to overcome host defences and infect healthy host tissue, while such resources may be less important among less specialized necrotrophic pathogens that can only infect weakened plants (Garrett, 1970). We also found large variation in spore size across asymbiotic saprotrophic fungi (for any group or spore type; Figure 2; Figure S3). This variation suggest the existence of different niches filled by saprotrophic species, such as during decay of different substrates or in different successional stages (Purhonen et al., 2020). Finally, we also included in a separate analysis the peculiar case of fungi that cause disease in humans due to their importance. Most of these fungi are described as opportunistic (i.e. causing disease in immuno-compromised individuals (Kendrick, 2017)) and are commonly found growing as free-living in nature; these fungi are, thus, generally considered asymbiotic rather than symbiotic in the mycological literature (Moore et al., 2011). Our results, however, show that such fungi, despite their expected asymbiotic nature, have on average smaller sizes than other asymbiotic fungi (a pattern that holds across the phylogeny in some of our models, Tables S3 and S4, Figure 2). While it is not possible to pinpoint mechanisms, we hypothesize that smaller spores for these fungi may enhance the likelihood to be passively inhaled or ingested (Moore et al., 2011).

Relative importance of species' symbiotic status in explaining offspring size variation

Symbiotic status was also more important for explaining interspecific variation in spore size than climate variables associated with the distributions of fungal species (Table 1). After symbiotic status, mean annual temperature was the second most important variable explaining spore size variation across communities. This is congruent with previous research highlighting that in some species of mushroom-forming fungi, thicker spore walls have higher resistance to UV light exposure and freezing temperatures than species with smaller and lighter spores (Norros et al., 2015). Possibly, for symbiotic fungi, environmental microclimate plays a minor role as the host will buffer these variables (e.g. fungal symbionts of warm-blooded fungal symbionts will be buffered against changes in environmental temperature).

Relationship between offspring size and species' geographic distributions depends on symbiotic status

Finally, we tested the relationships between spore size and geographic distributions for asymbiotic and

AGUILAR-TRIGUEROS ET AL.

TABLE 1 Relative importance of symbiotic lifestyle versus climatic variables in explaining interspecific spore size variation. The fit of two phylogenetic linear regression models with lifestyle and six climatic variables as explanatory factors is compared to the fit of models in which one of these predictors was removed (indicated in the respective row). The first model uses the phylogenetic tree based on whole genome sequences as provided in (Li et al., 2021), which includes 281 species from which we collected climatic data (referred to as the 'genome tree model'). The second model uses a taxonomy-based cladogram for species based on their taxonomy from kingdom to species level (referred to as the 'taxonomy tree model'), which includes 1137 species from which we collected climatic data. AIC, Akaike's Information Criterion; dAIC, delta AIC (difference between the AIC of each model and the one containing all terms). A dAIC>10 indicates no support for dropping that term from the model because it results in a large decline in model fit; dAIC between 4 and 7 indicate considerably low support for dropping that term; and dAIC<2 indicates strong support for dropping that term as it improves model fit (Burnham & Anderson, 2002).

Phylogenetic regression model	Adjusted r^2	Loglike	AIC	dAIC	Phylogeny used
All variables	0.21	-336.72	699.43		Genome tree
	0.09	-1328.2	2684.39		Taxonomy tree
(–) Symbiotic lifestyle	0.18	-341.41	706.82	7.39	Genome tree
	0.07	-1333.42	2690.85	6.45	Taxonomy tree
(–) mean annual temperature	0.21	-337.97	699.94	0.51	Genome tree
	0.09	-1328.23	2682.45	-1.94	Taxonomy tree
(-) mean annual precipitation	0.21	-337.25	698.49	-0.94	Genome tree
	0.09	-1328.34	2682.68	-1.71	Taxonomy tree
(–) Temperature seasonality	0.22	-337.02	698.04	-1.39	Genome tree
	0.09	-1328.23	2682.47	-1.93	Taxonomy tree
(-) Precipitation seasonality	0.21	-337	698	-1.43	Genome tree
	0.09	-1329.66	2685.32	0.93	Taxonomy tree
(-) Maximum solar radiation	0.18	-341.29	706.59	7.15	Genome tree
	0.08	-1330.52	2687.05	2.66	Taxonomy tree
(-) Minimum vapour pressure	0.22	-336.72	697.43	-2	Genome tree
	0.09	-1328.25	2682.49	-1.9	Taxonomy tree

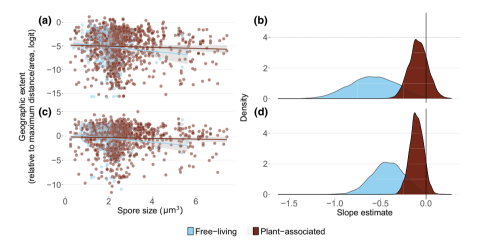


FIGURE 3 Asymbiotic fungal species exhibit a negative relationship between spore size and geographic distributions, while species plant-associated fungi do not. (a, c). Relationship between spore size and geographic distribution (based on polygon area [a] and the maximum distance between samples in which species were detected [c]) for asymbiotic fungal species and fungal species exhibiting varying degrees of host association to plants. Fungal species were detected in global surveys of environmental DNA from soil and plant material. (b, d) Bayesian models were fitted to estimate posterior distributions of the slope parameters representing the strength of the relationship between geographic extent and spore volume. The density plot represents the likelihood that a value associated with the slope estimate was present in the posterior distribution. These models included random intercepts representing the taxonomic order and spore type, as well as the primer set used to amplify fungal DNA (Tables S6–S8). Only species producing a single spore type were used in this analysis.

plant-associated fungal species, which we expect to be negative if smaller offspring size facilitates spread of propagules. Spore size was negatively correlated with the geographic range of free-living fungal species (95% credible interval for slope of maximum geographic distance: -0.71 to -0.11; 95% credible interval for slope of range area: -1.01 to -0.14; Figure 3) but not for symbiotic groups

(95% credible intervals for slope of maximum distance: -0.23 to 0.01; 95% credible intervals for slope of range area: -0.25 to 0.09; Figure 3). In asymbiotic fungi, species with larger spores had a more-limited geographic range compared to species with smaller spores, which may move more easily to new environments. Conversely, geographic range was unrelated to spore size for symbiotic

species, for which host-related factors (including the geographic spread of the host itself) may offset any difference in dispersal due to spore size. For example, smaller spore sizes might actually reduce the chances of 'landing' on a suitable host because smaller spores remain more easily aloft (Norros et al., 2014). The role of other spore traits (such as appendage morphology or spore wall ornamentation) must be assessed to fully understand the dispersal of symbiotic fungi (Gareth Jones, 2006).

DISCUSSION

In this study, we uncover massive variation in spore size in the fungal kingdom whose ecology and evolution is partly explained by transitions in symbiotic status of the species. However, we also found that the direction of this effect (i.e. shifts to smaller or larger spore sizes in symbiotic fungi) and its importance varies widely among symbiotic groups and phyla. For plant-associated fungi in the Ascomycota, our results provide support to the hypothesis that as symbiosis transitions from facultative to obligate, large spore reserves become more important for their survival during dispersal to and assist colonization of the plant. The results emphasize the critical role of symbiotic relationships in driving the evolution of life-history traits, especially in Fungi and suggest two avenues for further research. First, determining the mechanisms behind correlations between shits in spore size along transition to symbiosis; and second, determining why in some symbiotic groups and clades, spore size does not change along symbiotic gradients. For example, shifts to smaller spore sizes in symbiotic groups could be explained by host transmission dynamics. In other symbiotic interactions, such as parasitic animals, it has been proposed that small-sized offspring propagules are produced to increase the chances of transmission when hosts are hard to locate (Poulin, 2011). Exploring the influence of host transmission dynamics on fungal reproductive ecology can reveal intricate life-history strategies in the Fungi beyond spore structures. Direct transmission between hosts, for instance, may reduce reliance on spore dispersal and instead utilize alternative structures like hyphal extensions (as observed in mycorrhizal fungi (Bielčik et al., 2019)) or yeast phenotypes (as it is commonly seen in most insect gut endosymbionts (Gibson & Hunter, 2010)). In addition, as host-symbiont specialization is a long-term evolutionary process (Chomicki et al., 2020), the age of symbiosis might be a predictor of reproductive trait changes (for both host and symbionts). Thus, we propose that the reliance of fungi in other ways of transmission other than spores might explain weaker correlation and symbiotic state we found in some clades. In those cases, variation in spore size might be driven by neutral processes such as drift (which we did not test). In order to test these hypothesis it would be necessary to include more species across different symbiotic lifestyles in phylogenetic studies (James et al., 2020) and the need to populate databases with fungal reproductive traits. Such

data would allow tests of even the most fundamental tenets in life-history for the fungal branch of the tree of life, such as the existence of trade-offs in offspring output-offspring size or allometric scaling relationships between parent size and offspring size.

Finally, information on the diversity of dispersal and colonization strategies among asymbiotic and symbiotic fungi will be useful to forecast the impact of global change on ecosystem functions provided by fungi. For example, disease risk caused by fungal plant pathogens is forecasted to change with increasing global temperature (Chaloner et al., 2020). Such changes are likely due to direct effects on survival of spores during dispersal, and indirect effects of changing habitat quality (e.g. host susceptibility). Information on fungal dispersal strategies for symbiotic groups will refine forecasts of pathogen expansions and likelihood of pathogen spillover from natural ecosystems to croplands. Considering that fungi represent the main cause of crop yield losses and are a main threat to animal health (Fones et al., 2017), such refinements in forecasting are particularly relevant to maintain food security and ecosystem health.

In summary, expanding the realm of life-history analysis beyond plants and animals to other diverse and important clades such as fungi highlights symbiosis as a key biotic driver influencing the ecology and evolution of offspring-size variation. Life-history frameworks are biased towards free-living organisms (Falster et al., 2008; Stearns, 1992) with relatively limited inclusion of parasitic animals (Poulin, 2011). Yet, symbiosis is pervasive through the entire tree of life (including animals and plants) and, as we show here, it explains offspring variation among major clades in the fungal kingdom. Including symbiosis as a life-history parameter creates the need for new theoretical frameworks to determine, for instance, how much the host controls the offspring traits of the symbionts (as in fungi, and possibly bacteria and protists) and how much the symbionts control the offspring traits of their macroorganism hosts.

AUTHOR CONTRIBUTIONS

CAAT conceived the study together with FSK, WKC, JRP and MCR. WKC downloaded data from Mycobank. FSK developed the text mining algorithm. CAAT, JRP, CD and HZ mined the text data and cleaned spore data entries. CAAT digitized manually the spore size data not present in Mycobank, managed the spore database and assembled the fungal functional database. JRP managed and assembled climatic and geographic data. CAAT and JRP performed statistical analysis with input from WKC and FSK. CAAT wrote the first draft, and all authors contributed to the writing of the paper.

ACKNOWLEDGEMENTS

We thank J. Antonovics and Tessa Camenzind for helpful comments. Noa Terracina helped sort out plant disease information. We thank Louis Weiss for providing information on the natural history of microsporidian AGUILAR-TRIGUEROS ET AL.

fungi. We also thank anonymous reviewers for helpful comments to this manuscript. Funding. This research was supported by funding from the Federal Ministry of Education and Research (BMBF) within the collaborative Project 'Bridging in Biodiversity Science (BIBS)' (funding number 01LC1501A) to MCR. CAAT was supported by a Feodor Lynen Fellowship from the Humboldt Foundation. MCR acknowledges support from an ERC Advanced Grant (694368). CAAT, ICA, CD, HZ, MCR and JRP were supported by the Australia-Germany Joint Research Cooperation Scheme, an initiative of Universities Australia (UA) and the Deutscher Akademischer Austauschdienst (DAAD), for the project: 'A new tool of the trade: Trait-based approaches in fungal ecology'. JRP acknowledges support from the Australian Research Council (FT0100590). We acknowledge the contribution of the Biomes of Australian Soil Environments (BASE) consortium in the generation of data used in this publication. The BASE project was supported by funding from Bioplatforms Australia through the Australian Government National Collaborative Research Infrastructure Strategy (NCRIS). TV and PK were supported by the Czech Science Foundation (grant 21-17749S to T. Vetrovsky). Research on microbial distribution and colonization in the BKS laboratory is funded by the Australian Research Council (DP190103714). SH acknowledges funding from the German Science Foundation (grant HE6183). SNK was supported by start-up funds from the University of Tennessee, Knoxville. FTM acknowledges support from the European Research Council (ERC Grant Agreement 647038 [BIODESERT]) and Generalitat Valenciana (CIDEGENT/2018/041). AEZ acknowledges support from the National Science Foundation (DEB: 1623040, 'MacroMycoFunc - Forming an integrated understanding of function across fungi' and DEB: 1655759; 'Collaborative Research: NSFDEB-NERC: Tropical deadwood carbon fluxes: Improving carbon models by incorporating termites and microbes'). Open access funding enabled and organized by ProjektDEAL. Open Access funding enabled and organized by Projekt DEAL.

FUNDING INFORMATION

Alexander von Humboldt-Stiftung, Grant/Award Number: Feodor-Lynen Fellowship: Australian Research Council, Grant/Award Number: DP190103714FT0100590; Bundesministerium für Bildung und Forschung, Grant/Award Number: 01LC1501A; Deutsche Forschungsgemeinschaft, Grant/ Award Number: HE6183; Deutscher Akademischer Austauschdienst; Division of Environmental Biology, Grant/Award Number: 16230401655759; Grantová Agentura České Republiky, Grant/Award Number: 21-17749S; H2020 European Research Council, Grant/ Award Number: 647038694368: Universities Australia

CONFLICT OF INTEREST STATEMENT

The authors confirm that there are no competing interests.

PEER REVIEW

The peer review history for this article is available at https://www.webofscience.com/api/gateway/wos/peerreview/10.1111/ele.14271.

DATA AVAILABILITY STATEMENT

We confirm that all data and code used for this paper have been deposited in the database Fun Fun: https://github.com/traitecoevo/fungaltrait as well as in a Zenodo (10.5281/zenodo.7953831) and GitHub repository (https://github.com/aguilart/Symbiotic-status-and-fungal-spore-size).

ORCID

Carlos A. Aguilar-Trigueros https://orcid.

org/0000-0003-0512-9500

Franz-Sebastian Krah https://orcid.

org/0000-0001-7866-7508

Amy E. Zanne https://orcid.org/0000-0001-6379-9452

Nerea Abrego https://orcid.org/0000-0001-6347-6127

Petr Baldrian https://orcid.org/0000-0002-8983-2721

Claus Bässler https://orcid.org/0000-0001-8177-8997

Andrew Bissett https://orcid.org/0000-0001-7396-1484

Baodong Chen https://orcid.org/0000-0002-1790-7800

Manuel Delgado-Baquerizo https://orcid.

org/0000-0002-6499-576X

Stavros D. Veresoglou https://orcid.

org/0000-0001-6387-4109

Haiyang Zhang https://orcid.org/0000-0001-7951-0502

Jeff R. Powell https://orcid.org/0000-0003-1091-2452

REFERENCES

Aguilar-Trigueros, C.A., Hempel, S., Powell, J.R., Cornwell, W.K. & Rillig, M.C. (2019) Bridging reproductive and microbial ecology: a case study in arbuscular mycorrhizal fungi. *The ISME Journal*, 13 873–884

Bässler, C., Heilmann-Clausen, J., Karasch, P., Brandl, R. & Halbwachs, H. (2014) Ectomycorrhizal fungi have larger fruit bodies than saprotrophic fungi. *Fungal Ecology*, 17, 205–212.

Bates, D., Mächler, M., Bolker, B. & Walker, S. (2015) Fitting linear mixed-effects models using {lme4}. *Journal of Statistical Software*, 67, 1–48.

Bielčik, M., Aguilar-Trigueros, C.A., Lakovic, M., Jeltsch, F. & Rillig, M.C. (2019) The role of active movement in fungal ecology and community assembly. *Movement Ecology*, 7, 36.

Bürkner, P.-C. (2017) brms: An R Package for Bayesian Multilevel Models Using Stan. 2017, 80, 28.

Burnham, K.P. & Anderson, D.R. (2002) Edition 2. Model selection and multimodel inference: a practical information-theoretic approach. New York: Springer.

Calhim, S., Halme, P., Petersen, J.H., Laessoe, T., Bassler, C. & Heilmann-Clausen, J. (2018) Fungal spore diversity reflects substrate-specific deposition challenges. Scientific Reports, 8, 5356.

Chaloner, T.M., Gurr, S.J. & Bebber, D.P. (2020) Geometry and evolution of the ecological niche in plant-associated microbes. *Nature Communications*, 11, 2955.

Chomicki, G., Kiers, E.T. & Renner, S.S. (2020) The evolution of mutualistic dependence. *Annual Review of Ecology, Evolution, and Systematics*, 51, 409–432.

Davis Rabosky, A.R., Cox, C.L., Rabosky, D.L., Title, P.O., Holmes, I.A., Feldman, A. et al. (2016) Coral snakes predict the evolution of mimicry across New World snakes. *Nature Communications*, 7, 11484.

- Edelsbrunner, H., Kirkpatrick, D. & Seidel, R. (1983) On the shape of a set of points in the plane. *IEEE Transactions on Information Theory*, 29, 551–559.
- Falster, D.S., Moles, A.T. & Westoby, M. (2008) A general model for the scaling of offspring size and adult size. *The American Naturalist*, 172, 299–317.
- Fones, H.N., Fisher, M.C. & Gurr, S.J. (2017) Emerging fungal threats to plants and animals challenge agriculture and ecosystem resilience. In: *The Fungal Kingdom*. Washington, DC: ASM Press, pp. 787–809.
- Gareth Jones, E.B. (2006) Form and function of fungal spore appendages. *Mycoscience*, 47, 167–183.
- Garrett, S.D. (1970) Pathogenic root-infecting fungi. Cambridge: Cambridge University Press.
- Gibson, C.M. & Hunter, M.S. (2010) Extraordinarily widespread and fantastically complex: comparative biology of endosymbiotic bacterial and fungal mutualists of insects. *Ecology Letters*, 13, 223–234.
- Golan, J. & Pringle, A. (2017) Long-distance dispersal of fungi. In: The Fungal Kingdom. Washington, DC: ASM Press, pp. 309–333.
- Halbwachs, H. (2015) Gone with the wind—a review on basidiospores of lamellate agarics. *Mycosphere*, 6, 78–112.
- Halbwachs, H., Heilmann-Clausen, J. & Bässler, C. (2017) Mean spore size and shape in ectomycorrhizal and saprotrophic assemblages show strong responses under resource constraints. *Fungal Ecology*, 26, 59–64.
- Hawksworth, D.L. & Lücking, R. (2017) Fungal diversity revisited: 2.2 to 3.8 million species. *Microbiology Spectrum*, 5(4).
- Hijmans, R.J., Williams, E., Vennes, C. & Hijmans, M.R.J. (2019) Package 'geosphere'. Spherical Trigonometry, 1, 7.
- Ho, L.S.T., Ane, C., Lachlan, R., Tarpinian, K., Feldman, R., Yu, Q. et al. (2016) Package 'phylolm'. See http://cran.r-project.org/web/packages/phylolm/index.html (accessed February 2018)
- Ingold, C.T. (2001) Range in size and form of basidiospores and ascospores. *Mycologist*, 4, 165–166.
- James, T.Y., Stajich, J.E., Hittinger, C.T. & Rokas, A. (2020) Toward a fully resolved fungal tree of life. *Annual Review of Microbiology*, 74, 291–313.
- Kauserud, H., Colman, J.E. & Ryvarden, L. (2008) Relationship between basidiospore size, shape and life history characteristics: a comparison of polypores. *Fungal Ecology*, 1, 19–23.
- Kavanagh, P.H. & Burns, K.C. (2014) The repeated evolution of large seeds on islands. Proceedings of the Royal Society B: Biological Sciences, 281, 20140675.
- Kemen, E. & Jones, J.D. (2012) Obligate biotroph parasitism: can we link genomes to lifestyles? *Trends in Plant Science*, 17, 448–457.
- Kendrick, B. (2017) *The fifth kingdom*. Indianapolis: Hackett Publishing.
- Kivlin, S.N. (2020) Global mycorrhizal fungal range sizes vary within and among mycorrhizal guilds but are not correlated with dispersal traits. *Journal of Biogeography*, 47, 1994–2001.
- Kokkoris, V., Stefani, F., Dalpe, Y., Dettman, J. & Corradi, N. (2020) Nuclear dynamics in the arbuscular mycorrhizal fungi. *Trends in Plant Science*, 25, 765–778.
- Li, Y., Steenwyk, J.L., Chang, Y., Wang, Y., James, T.Y., Stajich, J.E. et al. (2021) A genome-scale phylogeny of the kingdom fungi. *Current Biology*, 31(8), 1653–1665.
- Lutzoni, F., Nowak, M.D., Alfaro, M.E., Reeb, V., Miadlikowska, J., Krug, M. et al. (2018) Contemporaneous radiations of fungi and plants linked to symbiosis. *Nature Communications*, 9, 5451.
- McLaughlin, D.J. & Spatafora, J.W. (2014) *Systematics and evolution:* part a. Berlin: Springer-Verlag.
- Mengiste, T. (2012) Plant immunity to necrotrophs. Annual Review of Phytopathology, 50, 267–294.
- Moles, A.T., Ackerly, D.D., Webb, C.O., Tweddle, J.C., Dickie, J.B. & Westoby, M. (2005) A brief history of seed size. *Science*, 307, 576–580.

- Moles, A.T. & Westoby, M. (2004) Seedling survival and seed size: a synthesis of the literature. *Journal of Ecology*, 92, 372–383.
- Moore, D., Robson, G.D. & Trinci, A.P. (2011) 21st century guidebook to fungi. Cambridge: Cambridge University Press.
- Naranjo-Ortiz, M.A. & Gabaldon, T. (2019) Fungal evolution: major ecological adaptations and evolutionary transitions. Biological Reviews of the Cambridge Philosophical Society, 94, 1443–1476.
- Neuheimer, A.B., Hartvig, M., Heuschele, J., Hylander, S., Kiørboe, T., Olsson, K.H. et al. (2015) Adult and offspring size in the ocean over 17 orders of magnitude follows two life history strategies. *Ecology*, 96, 3303–3311.
- Norros, V., Karhu, E., Norden, J., Vahatalo, A.V. & Ovaskainen, O. (2015) Spore sensitivity to sunlight and freezing can restrict dispersal in wood-decay fungi. *Ecology and Evolution*, 5, 3312–3326.
- Norros, V., Rannik, Ü., Hussein, T., Petäjä, T., Vesala, T. & Ovaskainen, O. (2014) Do small spores disperse further than large spores? *Ecology*, 95, 1612–1621.
- Paradis, E. & Schliep, K. (2018) Ape 5.0: an environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics*, 35, 526–528.
- Poulin, R. (2011) *Evolutionary ecology of parasites*. Princeton: Princeton university press.
- Purhonen, J., Ovaskainen, O., Halme, P., Komonen, A., Huhtinen, S., Kotiranta, H. et al. (2020) Morphological traits predict host-tree specialization in wood-inhabiting fungal communities. *Fungal Ecology*, 46, 100863.
- Robert, V., Vu, D., Amor, A.B.H., van de Wiele, N., Brouwer, C., Jabas, B. et al. (2013) MycoBank gearing up for new horizons. *IMA Fungus*, 4, 371–379.
- Rockett, T.R. & Kramer, C.L. (1974) Periodicity and Total spore production by Lignicolous basidiomycetes. *Mycologia*, 66, 817–829.
- Roper, M. & Seminara, A. (2019) Mycofluidics: the fluid mechanics of fungal adaptation. *Annual Review of Fluid Mechanics*, 51, 511–538.
- Stearns, S. (1992) The evolution of life histories, Oxford Univ. Press. Oxford. SteamsThe Evolution of Life histories1992.
- Stoddard, M.C., Yong, E.H., Akkaynak, D., Sheard, C., Tobias, J.A. & Mahadevan, L. (2017) Avian egg shape: form, function, and evolution. *Science*, 356, 1249–1254.
- Team, R.C. (2020) R: a language and environment for statistical computing. Vienna: R Foundation for Statistical Computing Vienna.
- Vincenty, T. (1975) Direct and inverse solutions of geodesics on the ellipsoid with application of nested equations. *Survey Review*, 23, 98, 03
- Wang, C. & Wang, S. (2017) Insect pathogenic fungi: genomics, molecular interactions, and genetic improvements. *Annual Review of Entomology*, 62, 73–90.

SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Aguilar-Trigueros, C.A., Krah, F.-S., Cornwell, W.K., Zanne, A.E., Abrego, N., Anderson, I.C. et al. (2023) Symbiotic status alters fungal eco-evolutionary offspring trajectories. *Ecology Letters*, 00, 1–12. Available from: https://doi.org/10.1111/ele.14271