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Article

Keywords:

DOI: <https://doi.org/>

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Additional Declarations: There is **NO** Competing Interest.

Fungal plastiphily and its link to generic virulence traits makes environmental microplastics a global health factor

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Key words: microplastics, plastiphily, pathogenic fungi, plastisphere mycobiome, generic virulence traits, ITS metabarcoding, co-occurrence networks, community assembly, human pathogens, global health factor, fungal communities

Abstract

Fungi comprise significant human pathogens, causing over a billion infections each year. Plastic pollution alters niches of fungi by providing trillions of artificial microhabitats, mostly in the form of microplastics, where pathogens might accumulate, thrive, and evolve. However, interactions between fungi and microplastics in nature are largely unexplored. To address this knowledge gap, we investigated the assembly, architecture, and ecology of mycobiomes in soil (micro)plastispheres near human dwellings in a model- and network-based metagenome study combined with a global-scale meta-analysis. Our results reveal a strong selection of important human pathogens, in an idiosyncratic, otherwise predominantly neutrally assembled plastisphere, which is strongly linked to generic fungal virulence traits. These findings substantiate our niche expansion postulate, demonstrate the emergence of plastiphily among fungal pathogens and imply the existence of a 'plastisphere virulence school', underpinning the need to declare microplastics as a factor of global health.

Introduction

One of humanity's irrevocable legacies for centuries to come is that all life on Earth must interact with plastics. Plastic is a pervasive pollutant in the biosphere, where it affects all levels of biological organisation¹. Elusive microplastic (MP; < 5 mm)² and nanoplastic particles (< 1 µm), primarily derived from fragmentation of larger plastic objects, can invade tissues and organs, and even infiltrate cells, where they can cause inflammatory responses and subcellular dysfunctions³. Organismic interactions with plastic debris, such as ingestion and entanglement, are known to sometimes have severe consequences in the fitness of individuals, including obstruction⁴, asphyxiation⁵ and impairment of life history traits⁶. Plastic debris also impacts populations and trophic networks by introducing allochthonous chemicals⁷, disrupting intraspecific communication⁸ or altering environmental conditions⁹. While direct and indirect impacts have been extensively studied primarily in relation to larger organisms and at local scales, microorganisms are increasingly being studied for cascading impacts of plastic pollution at community level that could affect entire ecosystems^{10,11}.

For microbes, MP pollution is essentially a terraforming event, with trillions of man-made, long-lived, physico-chemically diverse substrates permeating all of Earth's biomes^{12,13} and providing a variety of ecological niches for microbial life¹⁴. It is in these interaction landscapes that MP pollution probably unfolds its greatest ecological disruption potential. Therefore, in recent years, the focus of MP research has shifted to the microbial colonisation of plastic debris in different ecosystems, with the common goal of elucidating the nature of the plastisphere^{10,11,15}. The scientific consensus is that plastispheres harbour multipartite, dynamic microbial

communities that generally differ in structure and composition from biological assemblages in the surrounding compartment or other natural substrates^{11,14}. These altered microbiomes have serious ecological implications, such as the concentration of antibiotic resistance genes¹⁶, the facilitation of horizontal gene transfer¹⁷, and the accumulation and transmission of pathogenic microbes^{10,18}, which affect ecosystem functioning. Therefore, exploring the mechanisms that lead to the formation of epiplastic communities is crucial for assessing the ecological risks posed by the plastisphere microbiota.

Fungal pathogens have a huge impact on plant and animal life, driving species extinctions, causing ecosystem failure, and threatening food security¹⁹⁻²¹. Fungal infections in humans represent a devastating and worsening yet overlooked global health crisis, with over one billion cases and more than 1.6 million deaths annually^{22,23}. A critical factor in the emergence of fungal diseases is anthropogenic alteration of the natural environment, which entails new opportunities for fungal pathogens to thrive, diversify and evolve^{20,24}. Yet, plastisphere mycobiomes, especially in terrestrial environments, have been almost entirely omitted, although plastic pollution is one of the most profound alterations to the Earth's surface²⁵ and fungal pathogens are ecologically predisposed and biologically fine-tuned to benefit from it. Pathogenic fungi are excellent colonisers of hydrophobic substrates such as phyllospheres²⁶ and animal epithelia²⁷ that are also prevalent in all types of extreme and oligotrophic habitats in the natural and built environment²⁸, such as bare rock²⁹, medical devices³⁰ and even dishwashers³¹, where they form highly resistant, polymicrobial biofilms³². This adaptability and niche width is enabled by a spectrum of characteristic traits, most notably virulence factors such as extremotolerance²⁸, melanisation³³, oligotrophy^{32,34} and polymorphism^{35,36}. Preliminary evidence that MPs act as a reservoir for extremotolerant and melanised pathogenic fungi in terrestrial ecosystems¹⁰ suggests that the plastisphere is an ideal 'home' for pathogens and a putative 'school' for virulence, raising fundamental questions about the assembly of plastisphere mycobiomes.

Microbial community assembly is generally driven by two complementary processes: stochastic processes (neutral theory)³⁷, which emphasise the role of probabilistic dispersal and ecological drift, and deterministic processes (niche theory), which emphasise environmental filtering and biological interactions as drivers of community structure³⁸. Adopting niche theory to epiplastic communities would thus imply that certain microbes are positively selected by MPs and can thrive in the plastisphere due to differences in their relative ecological fitness, a phenomenon we call 'plastiphily'. Recent model integrations of these theories, however, have shown that neutral processes dominate the assembly of epiplastic bacteriomes in both aquatic³⁹ and soil environments^{16,40}, indicating that the bulk of these communities arise by chance. On the other hand, network-based studies have highlighted the importance of biotic interactions such as competition and facilitation, as well as so-called keystone taxa in the

formation of microbial biofilms on environmental plastics^{41,42}, and polymer type was considered as a deterministic driver in plastisphere bacteriome development¹¹. Selective factors play a crucial role in structuring mycobiomes during colonisation of extreme habitats⁴³ and the succession of dysbiotic mycobiota⁴⁴, which are inevitably linked to pathogenicity outcomes. Considering their systematic co-occurrence on soil MPs¹⁰ and assuming fitness advantages due to the above-mentioned virulence traits, it is likely that certain fungal pathogens are selected by MPs and thus 'plastiphilic'. To date, reports on the ecological mechanisms underlying the assembly of soil plastisphere mycobiomes are lacking, so at this stage virtually nothing is known about the origin of epiplastic fungal pathogens in terrestrial systems.

We tested the existence of such selection effects of MP by deciphering the ecological processes, biotic interactions and fungal traits that determine the assembly and architecture of soil (micro)plastisphere mycobiomes. We hypothesise that (1) the plastisphere metacommunity (MC) has an idiosyncratic architecture that (2) is structured by an interplay of neutral and deterministic processes, whereby (3) particularly pathogenic fungi act as keystone species and (4) are subject to selection effects. Finally, we hypothesise that (5) eventual selection correlates with the presence of generic virulence traits. Therefore, we applied a neutral community model and co-occurrence network analysis to ITS sequence data obtained by metabarcoding fungal (micro)plastisphere and soil metagenomes from municipal plastic pollution hotspots in Siaya town, Kenya. For appropriate functional and ecological characterisation, the identified phylotypes were pre-annotated with relevant trait data through an extensive database and literature-based meta-analysis. The model results were contextualised with phylogenetic and community ecology data as well as network outputs. Finally, we employed a novel model-derived selection index and correlation analysis to comparatively assess selection effects and disentangle potentially determinant traits. Our findings provide first insights into the origin of epiplastic mycobiomes and pathogenic fungi in the soil environment and describe the fungal plastiphily phenomenon with its alleged link to fungal virulence and global health relevance.

Results

Plastisphere and soil metacommunity. After consolidation of the data and exclusion of singletons, we obtained a total dataset consisting of 96 samples and 4,772,721 high-quality ITS sequences, which were clustered into 2,441 fungal OTUs (2,208 after rarefaction). About 37% of all phylotypes (838 OTUs) were identified at the genus or species level and subjected to meta-analysis (species-level synopsis in Supplementary Data 3). The soil MC was much more species-rich (1991 OTUs) than the plastic MC (927 OTUs), indicating that the plastisphere is the more selective compartment. The five most species-rich classes on MP,

Dothideomycetes, Sordariomycetes, Agaricomycetes, Eurotiomycetes and Tremellomycetes, accounted for a similar proportion of plastisphere (89%) and soil (86%) reads, while they were systematically and significantly more diverse in soil than in the plastisphere (Fig. 1b, Supplementary Table 3). Assignment to guilds resulted in four main guilds (Fig. 1c, g, Supplementary Data 1), of which saprotrophs were the most diverse in the soil ($p < 0.05$), while MP showed similar diversity levels for saprotrophs and pathogens (Fig. 1c). Among abundance fractions, conditionally rare and abundant taxa (CRAT) were similarly diverse between compartments, while all other fractions were significantly more diverse ($p < 0.05$) in the soil (Supplementary Table 9).

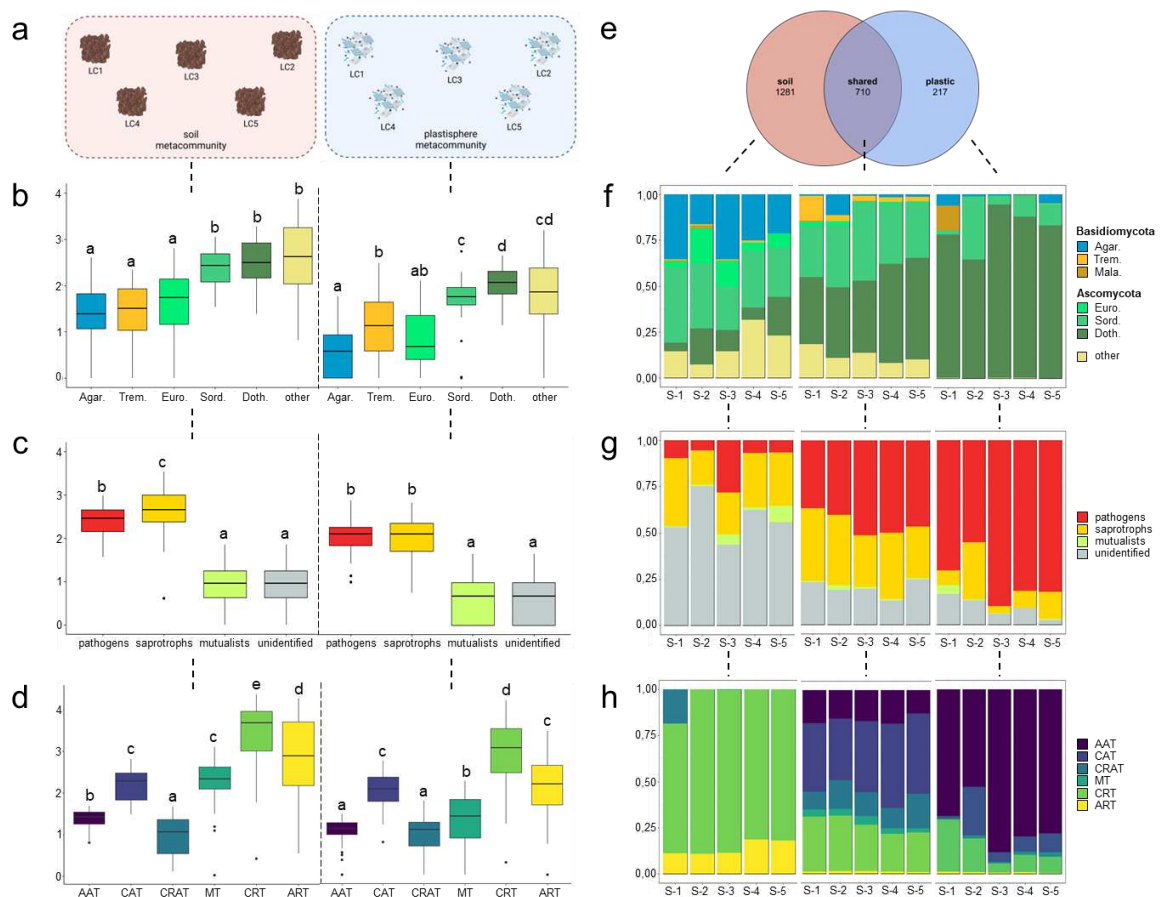


Fig. 1 Nested and idiosyncratic diversity between soil and plastisphere metacommunities. a Schematic illustration of soil and plastisphere metacommunities. LC = local community, corresponding to a soil or MP sample, respectively. Soil MC comprised 49 samples and 1991 OTUs, while plastisphere MC comprised 46 samples and 927 OTUs. **b** Shannon diversity among the most speciose classes found on MP and in the soil, where Agar. = Agaricomycetes, Trem. = Tremellomycetes, Euro. = Eurotiomycetes, Sord. = Sordariomycetes and Doth. = Dothideomycetes. 'other' codes for all remaining identified and unassigned classes of the respective compartments. **c** Shannon diversity among ecological guilds found on MP and in the soil. **d** Shannon diversity among abundance fractions of either compartment. ART, always rare taxa; CRT, conditionally rare taxa; CRAT, conditionally rare and abundant taxa; MT, moderate taxa; CAT, conditionally abundant taxa; AAT, always abundant taxa. The lower-case letters indicate significant differences between groups ($p < 0.05$) via one-way ANOVA. **e** Venn diagram showing the number of shared and specific OTUs of the metacommunities. Compartment-specific and shared abundances of **(f)** speciose classes, including Mala. = Malasseziomycetes, **(g)**

ecological guilds and **(h)** abundance fractions separated by sites (S-1 to S-5). Colour coding is consistent for all panels.

Compartment-specific and shared diversity. Around 64% of the soil-inhabiting and 23% of the plastic-colonising phylotypes were found exclusively in local communities of the respective compartments (specific MC), with 710 OTUs shared between both metacommunities (Fig. 1e). Across all sites, the soil-specific MC was dominated by Sordariomycetes and Agaricomycetes (Fig. 1f left), had large proportions of unidentified guilds, and small and low numbers of pathogens (Fig. 1g left), with most reads assigned to low-abundance fractions (Fig. 1h left). The shared MC comprised predominantly Dothideomycetes and Sordariomycetes (Fig. 1f middle) and between 36 to over 50% pathogens (Fig. 1g middle), with reads assigned to all six abundance fractions similarly across all sites (Fig 1h middle), resembling an 'intermediate state' between soil- and plastic-specific MC. In terms of abundance, the plastic-specific MC was solely dominated by Dothideomycetes and pathogens, which reached proportions of over 90%, respectively (Fig. 1f, g right), with always abundant taxa (AAT) comprising between around 50–90% of reads (Fig. 1h right). These observations demonstrate both the nestedness and strong idiosyncrasy of the metacommunities, as well as the presence and dominance of plastic-specific fungal pathogens.

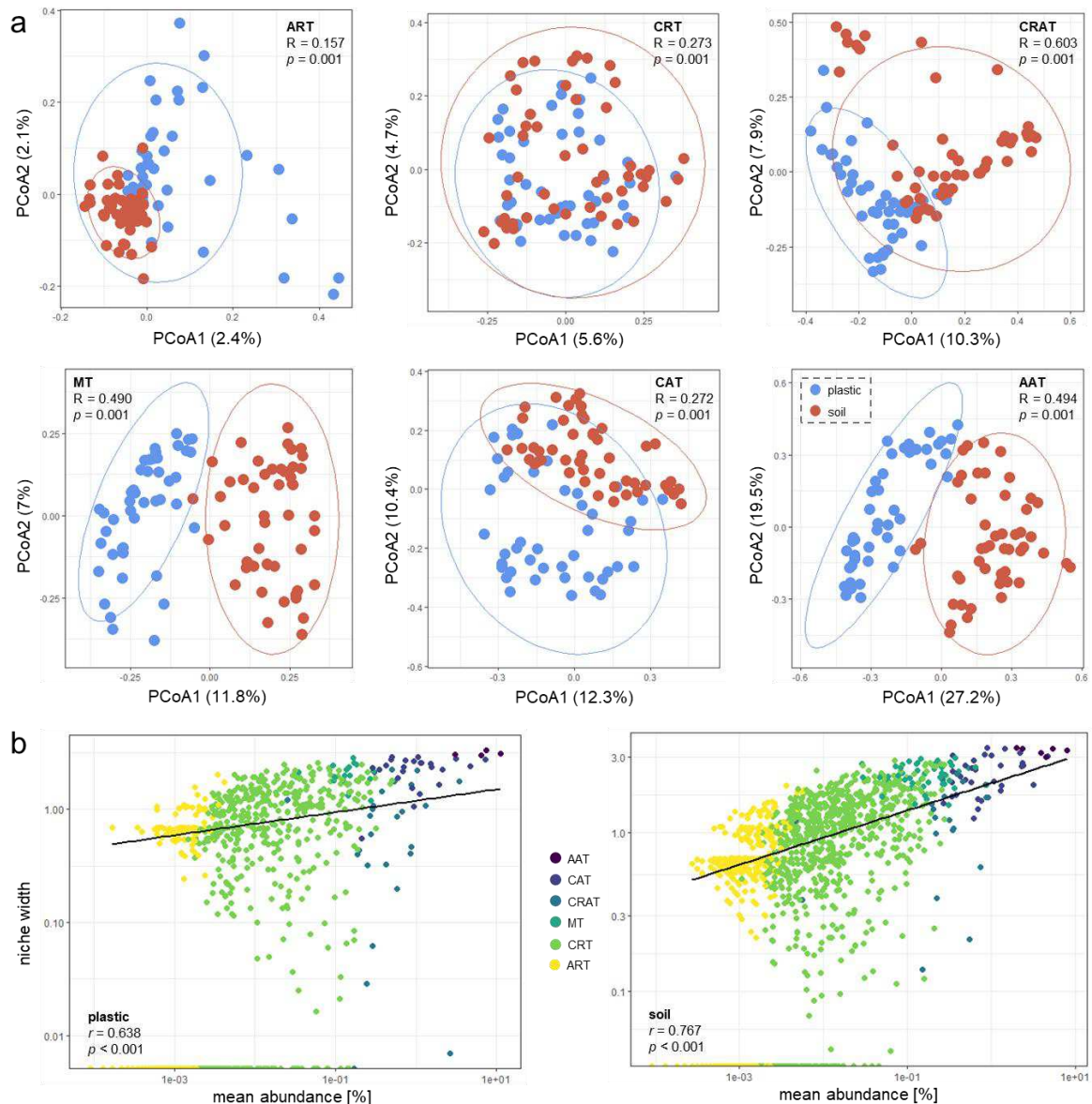


Fig. 2 Variation in mycobiome composition and niche width between soil and plastisphere abundance fractions. **a** Beta diversity between abundance fractions of soil and plastisphere metacommunities based on Bray-Curtis dissimilarity and visualised by PCoA ordination. ART, always rare taxa; CRT, conditionally rare taxa; CRAT, conditionally rare and abundant taxa; MT, moderate taxa; CAT, conditionally abundant taxa; AAT, always abundant taxa. Ellipsoids represent 95% confidence intervals. Similarities (sample statistic 'R') and significance (included p values) were assessed using ANOSIM. **b** Spearman correlation between fungal niche width and mean relative abundance of OTUs in the plastisphere and soil MC. Bubble colours code for the six distinct abundance fractions mentioned above. Correlation coefficient (r) and significance (p) value included.

PCoA ordinations and ANOSIM tests of each fraction (Fig. 1d) revealed a complete compositional separation of soil and plastisphere fungal communities ($p < 0.01$), with the highest differences in beta diversity between CRAT ($R = 0.603$) and AAT fractions ($R = 0.494$, Fig. 2a, Supplementary Table 10). Mean abundance correlated significantly ($p < 0.01$) and strongly ($r > 0.6$) with niche width in both MCs (Fig. 2b), with no difference between average niche width between compartments.

To further characterise the plastic-specific MC, we assessed the phylogenetic distributions of the 84 plastic-specific phylotypes identified to species or genus level (Fig. 3). These fungi made up 9% of OTUs and ca. 13% of reads of the entire plastisphere MC and belonged predominantly (79 taxa) to the subkingdom Dikarya, indicating an overall low evolutionary distance. Ecologically, plastic-specific fungi could be roughly categorised into saprotrophic or mutualistic non-opportunists with low to no virulence score (37 taxa, e.g., Agaricomycetes, Leotiomyces, Mortierellomycetes) and predominantly pathogenic opportunists with high virulence score (34 taxa, e.g., diverse Dothideomycetes, most Tremellomycetes and all Eurotiomycetes). Among fungi explicitly relevant to human health, we found dimorphic, polyextremotolerant black yeasts, such as *Exophiala* and *Knufia*; ubiquitous, hyaline multi-host pathogens like *Fusarium* and *Aspergillus*; several highly clinically relevant, cryptococcal yeasts, including *Papiliotrema*, *Filobasidium* and *Cryptococcus*; and multiple allergenic, toxigenic and pathogenic black fungi, such as *Alternaria*, *Curvularia*, *Chaetomium* and *Stachybotrys*, which were by far the most abundant plastic-specific fungi (ca. 80% of reads).

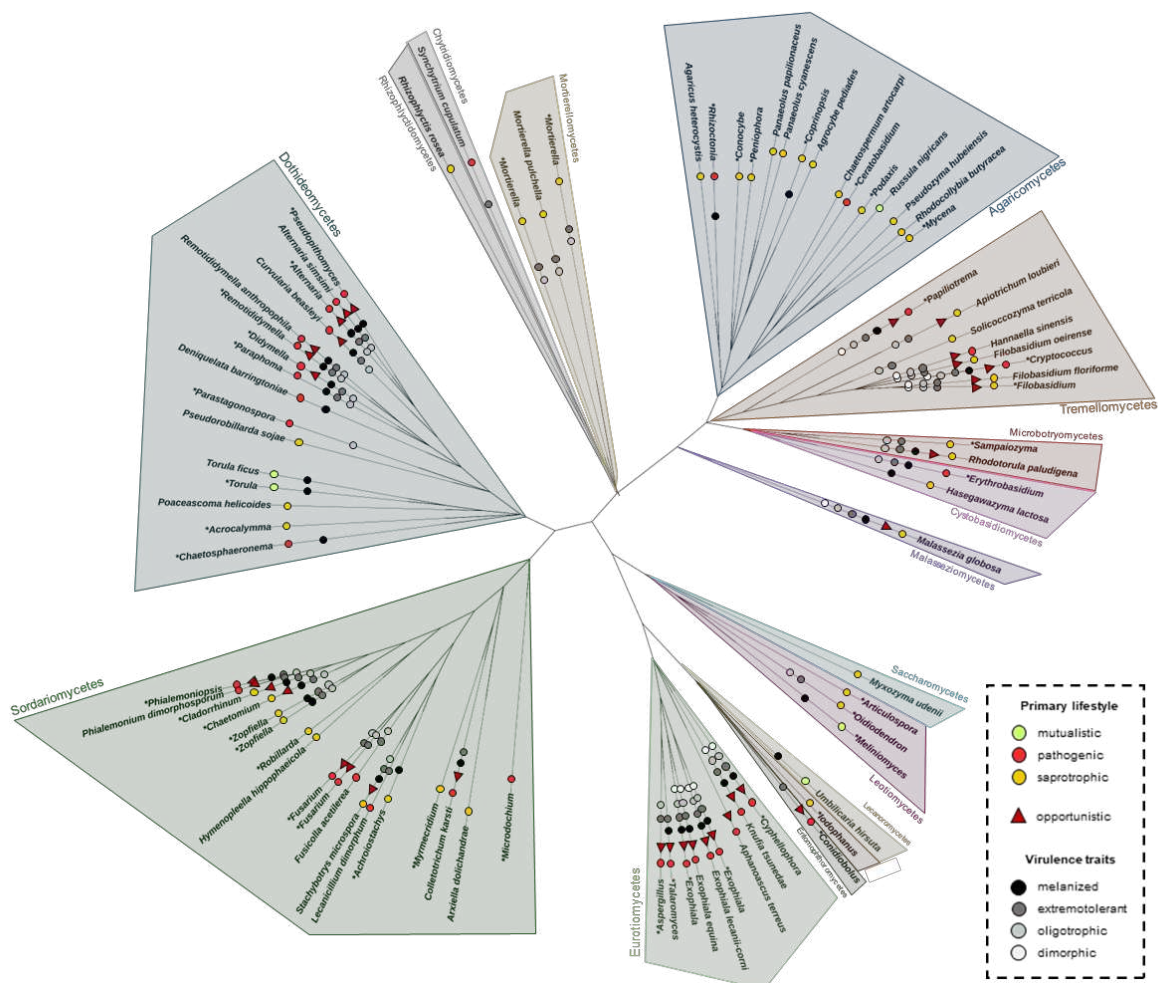


Fig. 3 Phylogeny of plastic-specific fungi. Displayed are those 84 OTUs that have been found exclusively on MP and identified at genus or species level. Clade colours code for fungal classes.

Phylotypes were classified into ecological guilds (green, red and brown bubbles) and annotated with trait data including opportunism (dark red triangle) and the four generic virulence traits, namely melanisation (black bubble), extremotolerance (dark grey), oligotrophy (light grey) and polymorphism (white). Traits were only annotated when present. Pez. = Pezizomycetes. (*) indicates 'unclassified'.

Mycobiome assembly. The neutral community model (NCM) of Sloan et al.⁴⁵ was used to assess the contribution of stochastic and deterministic processes to mycobiome assembly in microplastispheres and the bulk soil (Fig. 4). As indicated by the overall lower Akaike's information criterion (AIC) scores, the NCM outperformed the binomial distribution model for both compartments (Supplementary Table 11 and 12), indicating an influence of passive dispersal and ecological drift beyond purely random sampling from the respective metacommunities⁴⁶. The NCM estimated much of the correlation between occurrence frequency and relative abundance variation of plastisphere and soil OTUs, explaining 62% (plastic) and 67.5% (soil) of the community variance (Fig. 4), indicating the stronger effect of stochastic processes on soil community assembly. Migration rates (m) were estimated to be higher for soil than for plastisphere fungal communities (Supplementary Table 11), suggesting greater dispersal of fungi from the soil MC. In both compartments, there were several phylotypes that occurred more or less frequently than predicted by the NCM given their overall abundance in the MC (Fig. 4 and Supplementary Data 2). Points that are above the prediction (cyan) represent fungi that are more frequent than expected, suggesting that they are actively selected and maintained by the compartment, while points below the prediction frame (dark red) represent fungi that occur less frequently than expected, suggesting that the environmental compartment is either selecting against them or limiting their ability to disperse (Fig. 4). Plastisphere taxa above prediction are therefore considered plastiphilic fungi.

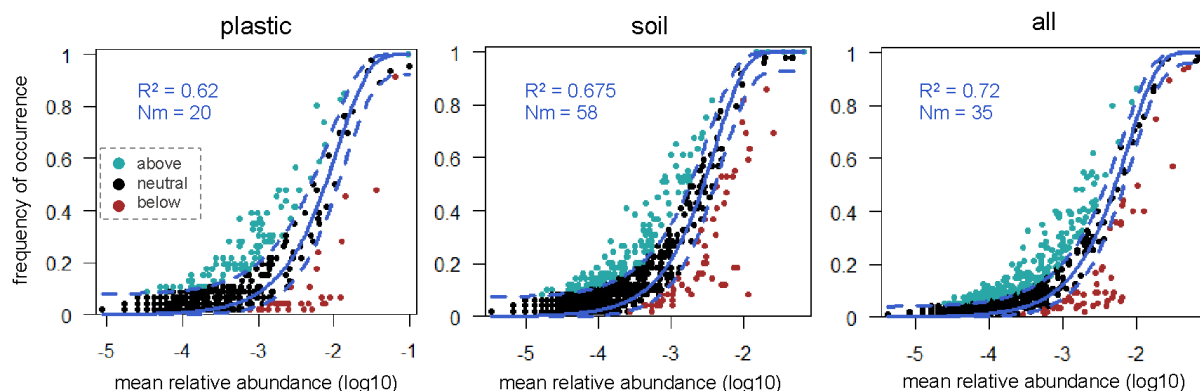


Fig. 4 Fit of the neutral community model for mycobiome assembly. The predicted occurrence frequencies for plastic, soil and all representing fungal communities from MP, soil and both compartments, respectively. Each point represents a fungal OTU and different colours indicate OTUs that occur more (cyan) or less frequently (dark red) than predicted by the neutral model. Solid blue lines indicate the optimum fit to the NCM and dashed blue lines represent 95% confidence intervals around the model prediction. R^2 indicates the fit of the neutral model and Nm equals the MC size times immigration.

Ecological distinction between neutral and non-neutral partitions. To disentangle the effects of selective processes on community structure and ecology, we separated each MC into three partitions comprising the OTUs found above, below, and within the neutral prediction, respectively (Supplementary Data 2). Taxa from the neutral and non-neutral partitions showed different community composition, diversity, and abundance in soil and plastic (Fig. 5). Beta diversity among partitions and compartments were assessed using Hellinger distance based on square-root transformed read counts and visualised by non-metric multidimensional scaling (NMDS). Community structure between partitions differed significantly in plastic (ANOSIM $R = 0.765$), soil ($R = 0.872$) and between compartments ($R = 0.845$) at $p < 0.01$ (Fig. 5a). In both compartments, the non-neutral partitions of MC exhibited higher within-group and between-group heterogeneity than the neutral partitions (Supplementary table 14), as shown by the density of each cluster and the ordination distances, respectively (Fig. 5a). Overall, neutral partitions accounted for the majority of fungal OTUs for plastic (85%) and soil (88%) (Fig. 5b). However, the fitting proportions of abundance differed markedly between compartments, with neutral partitions accounting for 93% of soil reads but only 60% of MP reads (Fig. 5b), indicating the prevalence of dominant taxa within the non-neutral plastisphere. Moreover, the plastisphere above-partition harboured the most even (ANOVA $p < 0.05$) and second most diverse ($p < 0.05$) community among all six partitions (Fig. 5c, Supplementary Table 17 and 18), suggesting the existence of a diverse, relatively equitable plastiphilic mycobiome.

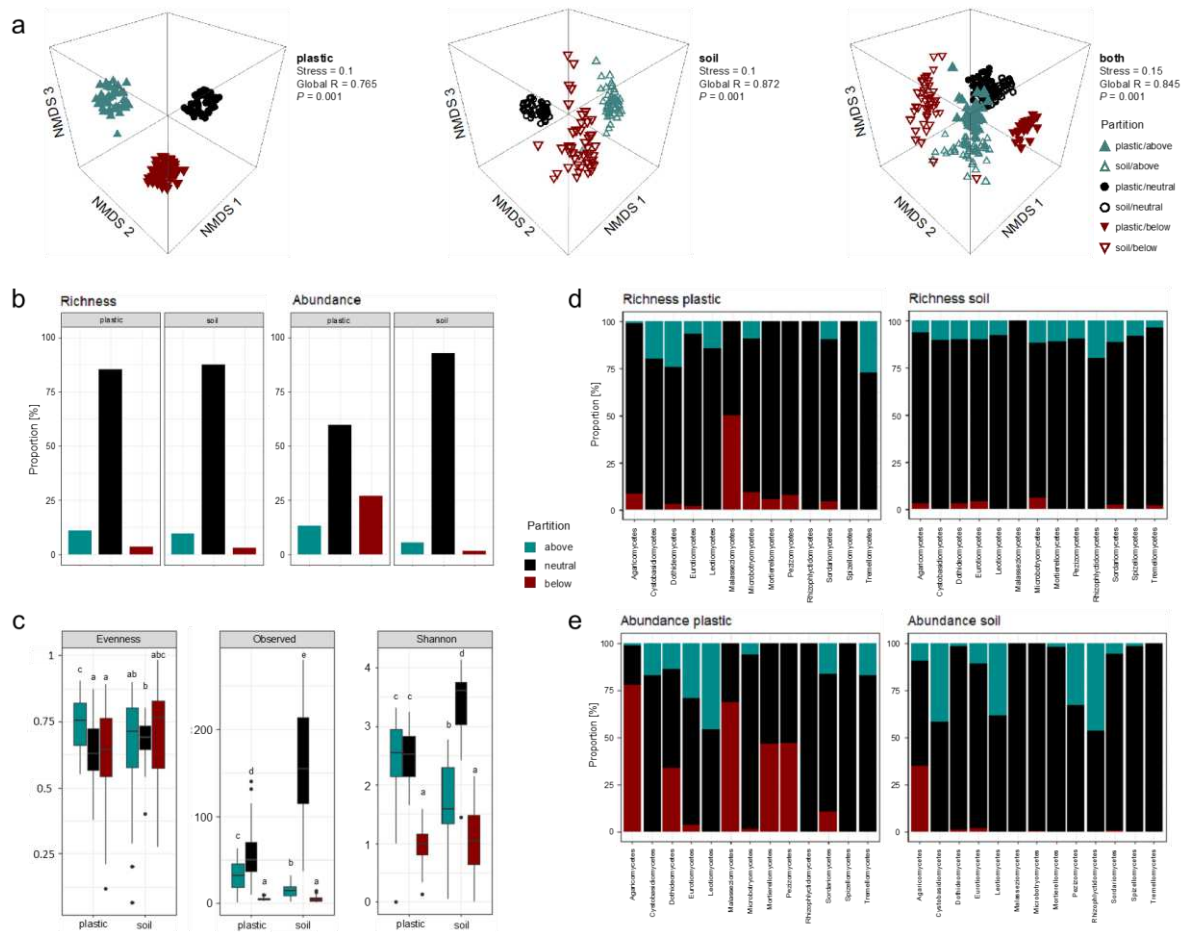


Fig. 5 Ecological variance between neutral and non-neutral partitions. Communities were pooled and OTUs were then divided into separate sample partitions based on their consistence with (black), deviation above (cyan) or below (dark red) the neutral prediction. Colour coding is consistent for all panels. **a** NMDS ordination based on Hellinger distance resemblance matrices of square-root transformed OTU abundances. Significant variation between partitions was tested by PERMANOVA (p -values) and ANOSIM (global R). NMDS ordination stress values included. **b** Fitting proportions of fungal OTU richness and abundance by the Sloan NCM. **c** Inter-partitional alpha diversity metrics according to compartment, including species richness, Shannon diversity and Pielou's species evenness estimated for plastic and soil. **d** Distinct fitting proportions of richness (OTU number) and **e** abundance (sequence number) of plastisphere and soil fungal classes, including only those identified classes comprising at least 0.1% of sequences.

Network properties and fungal keystone taxa. To explore the fungal relationships in the soil and in the plastisphere, co-occurrence network analysis based on the most prevalent OTUs (present in > 10% of respective samples) was performed using only significant and strong Spearman correlations ($r > 0.6$ or < -0.6 , $p < 0.01$). The two networks showed considerable differences in their structure and topological properties (Fig. 6a and Supplementary Table 19). The soil network consisted of 259 nodes and 498 edges (1 negative), with a modularity of 0.77, an average path length of 5.82 and an average degree of 3.85. The plastisphere network was considerably smaller, with 97 nodes and 166 exclusively positive edges, and had lower values for modularity (0.63), path length (3.73) and degree (3.42). However, the plastisphere network had the higher clustering coefficient (0.41) and harboured around twice the proportion of

pathogens compared to the soil network with an overall markedly higher average virulence score (Fig. 6a, Supplementary Table 19). To investigate the keystone taxa behind the assembly of plastisphere and soil MC, the fungi with the highest degree and betweenness centrality (BC) were identified for each network (Supplementary Table 20)^{47,48}. Among the two networks, the OTUs functioning as central hubs (highest degree), gatekeepers (highest BC), and thus also keystones were completely different. The keystone taxa of the plastisphere were mostly pathogenic, mainly plastiphilic Sordariomycetes and Dothideomycetes, with varying frequency and relatively high relative abundance, some of which are also known plastic biodegraders, such as *Fusarium* and *Phoma* (Fig. 6 b–d, Supplementary Table 20).

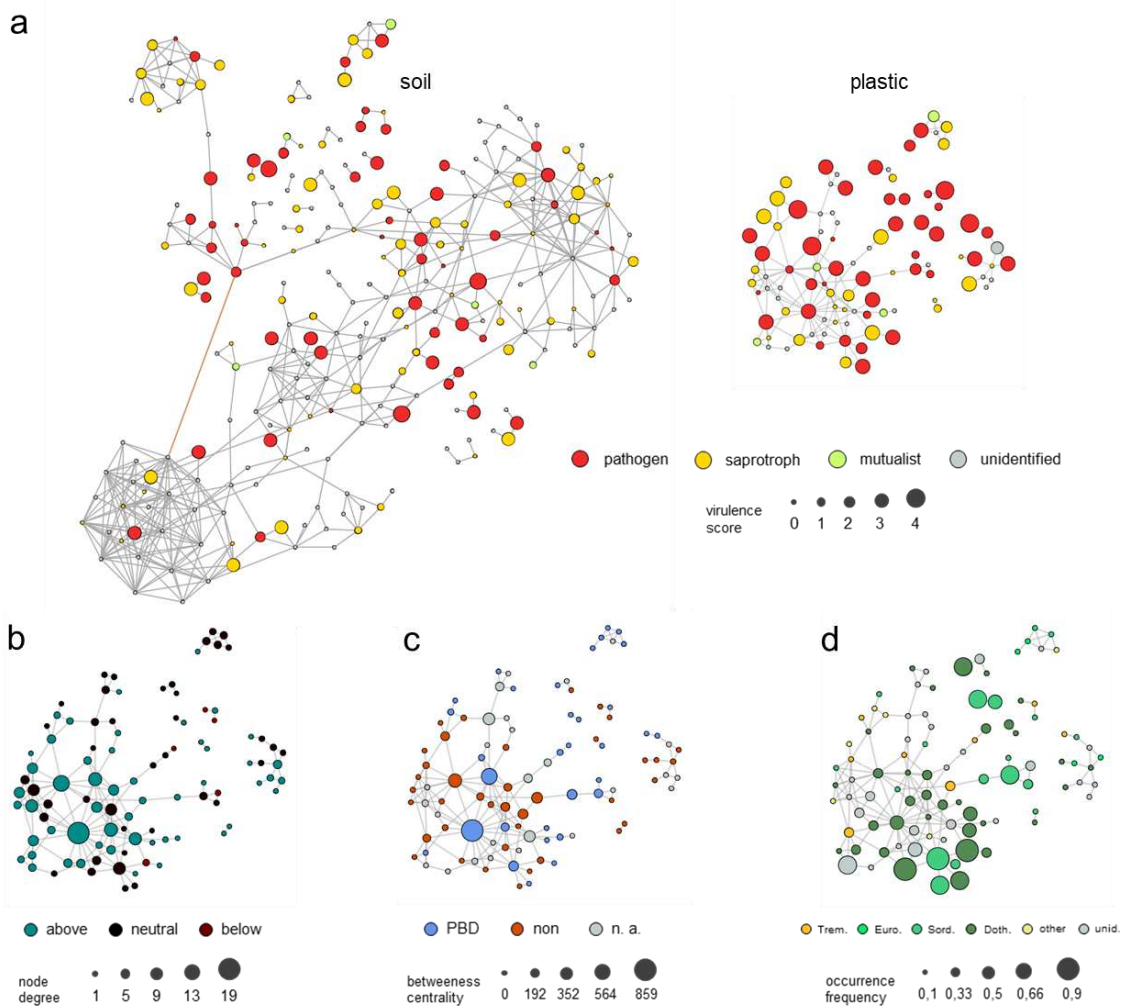


Fig. 6 Co-occurrence networks of soil and plastisphere mycobiomes. Networks only include OTUs with > 10% occurrence frequency across the soil or plastisphere MC, respectively. Nodes represent OTUs and edges represent significant ($p < 0.01$) and strong ($p > 0.6$ or < -0.6) Spearman correlations between nodes, where grey = positive and orange = negative correlations. **a** Comparison between soil (left) and plastisphere network (right). Node colour indicates guild affiliation and node size is proportional to the number of GVTs of each fungal OTU at the genus level (i.e., virulence score). **b** Correlations between plastiphilic and non-plastiphilic OTUs in the plastisphere. Node colour encodes the respective partition of each OTU, with above-partition OTUs (cyan) representing plastiphilic fungi. Node size is proportional to the number of connections of a node (i.e., degree). **c** Co-occurrences of putative plastic biodegraders in the plastisphere. Node colour indicates plastic biodegradability, where PBD = plastic biodegraders, non = non-degraders and n. a. = unidentified fungi without corresponding annotation.

Node size is proportional to the number of shortest paths between any pair of nodes passing through an individual node (i.e., betweenness centrality). **d** Relationships between plastisphere fungi of distinct classes. Node colour indicates class affiliation, where Trem. = Tremellomycetes, Euro. = Eurotiomycetes, Sord. = Sordariomycetes, Doth. = Dothideomycetes, other = all other assigned classes and unid. = unidentified classes. Node size is proportional to the relative occurrence frequency of each OTU across the entire plastisphere MC.

Plastiphily among ubiquitous human pathogens. To test for differential selection of fungal pathogens by soil and MP, different taxonomic and ecological groups of pathogenic fungi were selected (Supplementary Table 21, Supplementary Data 3). These included general ‘pathogens’ and ‘opportunists’ and their subgroups of Dothideomycetes and Tremellomycetes, as these classes were not only relatively the most selected by MP (Fig. 5d) but also characteristic of the plastisphere¹⁰. Furthermore, we classified genera shared between compartments that have a ‘cosmopolitan’ distribution, ‘high’ or ‘extremely high’ adaptability and are ‘opportunistic’ as the ‘CAO’ (cosmopolitan adaptable opportunists) group (21 genera) and the genera *Aspergillus*, *Candida*, and *Cryptococcus* additionally as “hidden killers”¹⁹, to individually assess selection effects associated with globally relevant and the most important fungal pathogens for humans, respectively. Selection was quantified using within-group selection (WGS), i.e., the proportion of above OTUs in a group to the total OTUs in that group, across-metacommunity selection (AMS), i.e., the proportion of above OTUs in a group to the above OTUs in the entire MC, and a selection index (SI) calculated from the product of the two. In addition, we determined the times average selection (TAS) of groups from the ratio of the groups’ WGS and the average selection in the respective MC, i.e., the proportion of the above OTUs in the MC (Supplementary Table 21 and 22).

Over 25% of epiplastic pathogens were in the above-partition (TAS of 2.4), accounting for just under half of all plastiphilic fungi, and showed selection 3 to 22 times stronger than any other guild according to the SI (Supplementary Table 21–23). In contrast, only 9% of all soil pathogens were in the above-partition (AMS of 13%), which showed slightly below-average selection (TAS < 1), and a 2.5-fold lower SI than saprotrophs. WGS and TAS of all eight pathogenic groups investigated were significantly higher on MP than in the soil ($p < 0.001$) (Fig. 7b, Supplementary Table 22). The correlation between AMS and WGS was highly significant and almost perfectly inverse for the pathogenic groups on MP ($r = -0.95$, $p < 0.01$) but positive in soil ($r = 0.76$, $p < 0.05$). According to the SI values, fungal species belonging to opportunists, CAO, hidden killers, dothideomycetous and tremellomycetous opportunists were selected 6, 9, 18, 37 and 275 times more strongly on MP than in soil, respectively (Fig. 6a, Supplementary Table 22). Overall, 19 out of 21 CAO genera were relatively more speciose on MP (compared to 2 in soil), while species of 13 out of 21 CAO genera were more strongly or exclusively selected by MP (compared to 5 in soil) (Fig. 7a). The positive correlation between WGS and AMS was equally strong and significant ($r = 0.86$, $p < 0.01$) for CAO genera on

plastic and in the soil, averaging 27% and 1.6% on MP, and 9% and 0.4% in soil, respectively (Fig. 7c, Supplementary Table 21 and 22). Among these genera, *Curvularia* (about 100 times higher than in the soil), *Hannaella* and *Cryptococcus* (both selected by MP only) had the highest SI in the plastisphere (Fig. 7a, c).

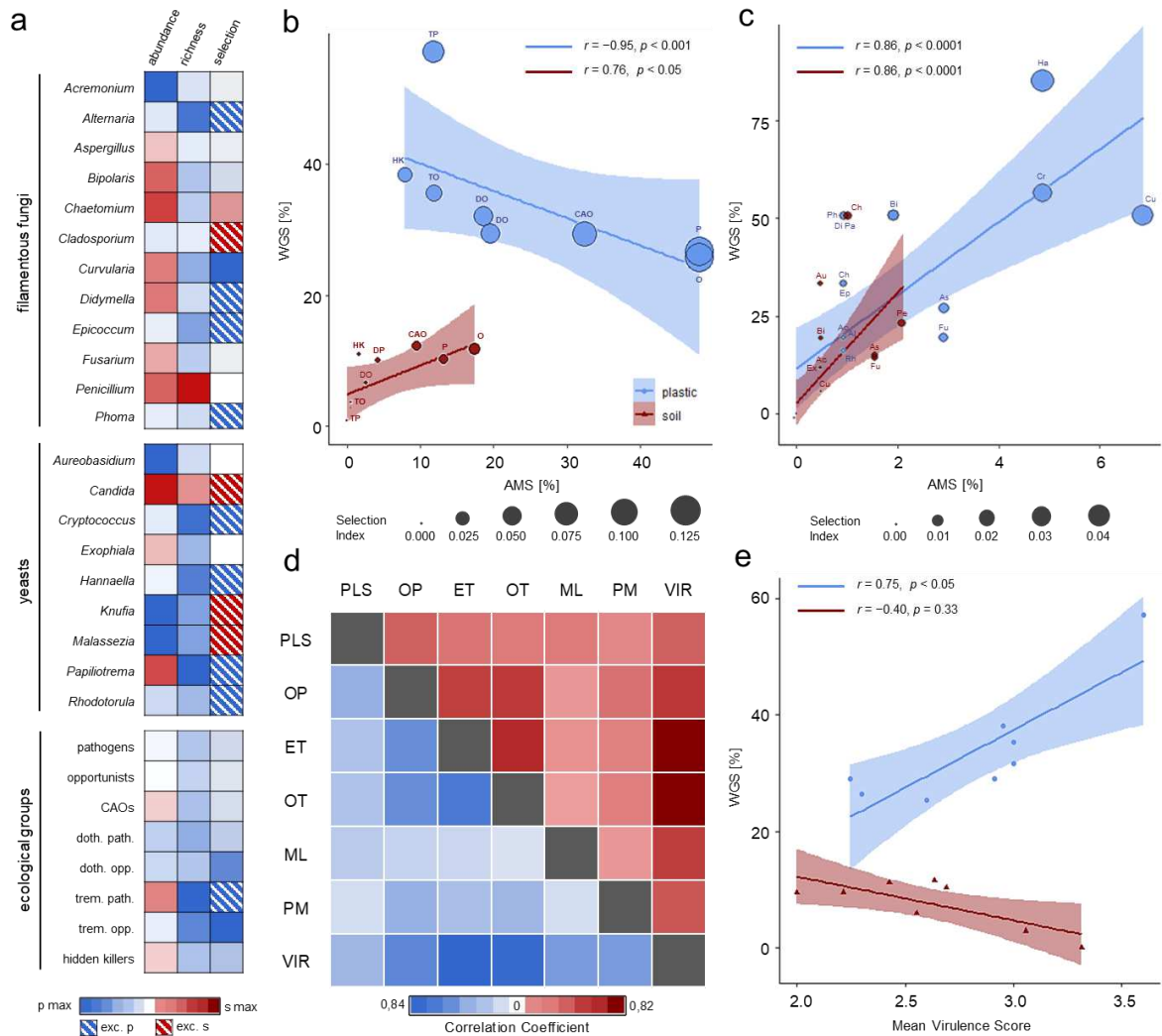


Fig. 7 Differential selection of pathogenic fungi between soil and plastisphere. **a** Ratios of relative abundance, relative richness, and selection of different taxonomic and ecological groups of pathogenic fungi between MP and soil. The first 21 groups listed represent CAO (cosmopolitan adaptable opportunist) genera separated according to growth form into ‘filamentous fungi’ and ‘yeasts’. The ‘ecological groups’ include pathogens, opportunists, taxonomic subgroups of them, the total CAO group and ‘hidden killers’ comprising the genera *Aspergillus*, *Candida*, and *Cryptococcus*. Here, doth. = dothideomycetous, trem. = tremellomycetous, path. = pathogens and opp. = opportunists. Blue colour indicates MP:soil ratio > 1, red colour indicates ratio < 1 and white colour indicates ratio = 1. Darker colours indicate more extreme ratio values. Striped boxes indicate exclusive selection of the group on MP (blue) or soil (red). Selection was assessed using the selection index (SI) introduced in this study, which is calculated from the product of within-group selection (WGS) and across-metacommunity selection (AMS). **b** Correlation between WGS and AMS for pathogenic groups, where P = pathogens, O = opportunists, DP = dothideomycetous pathogens, DO = dothideomycetous opportunists, TP = tremellomycetous pathogens, TO = tremellomycetous opportunists and HK = hidden killers. Bubble size corresponds to SI value. **c** Correlation between WGS and AMS for CAO genera. Genera were abbreviated with the first two letters. Bubble size corresponds to SI value. **d** Correlation heatmap showing Spearman correlations between different combinations of traits. PLS = pathogenic lifestyle, OP = opportunism, ET = extremotolerance, OT = oligotrophy, ML = melanisation, PM = polymorphism and

VIR = virulence score. Darker colours indicate stronger positive correlations. **e** Correlation between WGS and mean virulence score for pathogenic groups. Correlation coefficient (r) and significance (p) value included.

Link between plastiphily and generic virulence traits. Virulence score was significantly ($p < 0.01$) positively correlated with opportunism ($r = 0.65$ and 0.59) and pathogenic lifestyle ($r = 0.4$ and 0.39) in both MCs (Fig. 6d). Opportunism and pathogenic lifestyle were significantly positively correlated with each individual GVT in both compartments, with oligotrophy and extremotolerance each having the strongest correlations (Fig. 6d, Supplementary Table 24). Overall, extremotolerance and oligotrophy were the two GVTs correlated most strongly with each other ($r = 0.73$ and 0.65 , $p < 0.01$). Pathogenic groups had systematically and significantly higher virulence scores than their non-pathogenic or non-opportunistic counterparts for both compartments (Supplementary Fig. 1, Supplementary Table 25). The virulence scores of the pathogenic groups were thereby higher on MP than in soil, although not significantly (Supplementary Table 22). We found a strong and significant positive correlation between WGS and virulence score for pathogenic groups on MP ($r = 0.75$, $p < 0.05$), which was not the case in the soil ($r = -0.44$, $p > 0.05$) (Fig. 6e).

Discussion

The methodological adequacy of the molecular analyses and the strict compliance of the research data management with the FAIR++ guiding principles⁴⁹ were approved in the previous study¹⁰. The variation between soil and MP might be more pronounced than found here, as a complete separation of the compartments is technically not possible at this point⁵⁰, leading to cross-contamination that artificially increases the degree of mycobiome nestedness. Describing ecological phenomena from community barcoding data requires derivation of functional information from taxonomic profiles⁵¹. Fungal metabarcoding studies, however, suffer from a general scarcity of appropriate databases, insufficient taxon coverage, and a limited number of investigated traits^{51,52}. For a comprehensive and accurate assignment of trait data in our study, we collected and harmonised records and information from five relevant databases and hundreds of scientific publications, annotating nine traits for all identified fungal genera at an appropriate granularity (Supplementary Data 1). Virulence is a context-dependant, multifactorial and relative extended phenotype of microorganisms⁵³. Unlike bacteriologists, mycologists have not defined classical virulence factors, even in notorious human pathogens, due to a teleological-holistic view of such traits²⁸ and the largely opportunistic nature of fungal infections⁵³. Here, we used a set of well-described, commonly observed and symplesiomorphic autecological^{28,54} and physiological traits⁵⁵⁻⁵⁷ that contribute

to the pathogenicity and virulence of a wide variety of fungi (Supplementary Data 1), and thus qualify as 'generic virulence traits' (GVTs) to capture the virulence potential of fungi. Overall, we therefore argue that our meticulous implementation, fair documentation, and comprehensive contextualisation allow a clear interpretation of the obtained results in a synecological context and thus inferences and predictions of real-world phenomena.

Our analyses revealed significant compositional, ecological, and phylogenetic differences between soil and plastisphere MCs (Fig. 1–3), indicating the emergence of idiosyncratic mycobiomes on MP. Previous studies have also reported structural differences between fungal communities on MP and in the surrounding medium^{58,59}, but primarily focused on high-abundance taxa¹⁰. Comparing different abundance fractions, we found clear differences in alpha (Fig. 1d) and beta diversity (Fig. 2a) between epiplastic and soil fungal communities in each fraction, including the so-called “rare biosphere”⁶⁰ (ART and CRT). Plastic appears to favour the presence of Dothideomycetes and Tremellomycetes, as well as fungal pathogens in general (Fig. 1b, c and Fig. 2b, c). Saprotrophic fungi dominate fungal biodiversity outside symbioses or on non-living substrates^{61,62}. The observed convergence of saprotroph and pathogen diversity on plastic (Fig. 1c) makes it possibly the only inanimate fungal habitat with such a pattern of pathogen (co-)dominance, reminiscent of dysbiotic mycobiomes during disease^{63,64}. Among the plastic colonisers, we detected for the first time many plastic-specific fungi that were not present in the surrounding soil (Fig. 3). Unlike the rare biosphere inhabiting plastic-specific bacteria from marine MP described by Scales et al.⁶⁵, the plastic-specific fungi here were among the conditionally and always abundant taxa (Fig. 1h), suggesting a high ecological relevance for the plastisphere biome. We hypothesise that these specific fungi were alien species introduced from other habitats by windblown MP or singular disposal events. The discovery of specificity and cryptic diversity in the plastisphere (Fig. 1f, g) demonstrates the potential of MPs as a reservoir of exclusive and unexplored fungi, and a vector for alien species, exposing how little we know about the origin, scope, and complexity of the fungal-plastic reciprocity.

The soil plastisphere in its entirety represents a fragmental environmental compartment embedded in the three-dimensional continuum of the soil matrix. This habitat fragmentation implies community nestedness⁶⁶, while the spatial proximity suggests community coalescence between the two compartments⁶⁷. We found more than 75% of epiplastic OTUs also in the soil, where they accounted for slightly more than one third of the MC (Fig. 1e), indicating that the plastisphere is largely nested in the soil mycobiome. This finding is consistent with the ‘nested subset theory’ according to which communities with lower species richness in habitat patches are nested in species-rich communities of larger habitats⁶⁸. In terms of dominance structure of taxonomic and ecological groups, the shared MC clearly represented an ‘intermediate state’

between specific MCs, as shown by the transition in relative abundance of Dothideomycetes and pathogens (Fig. 1f, g). We interpret these patterns as indicative of intercompartmental coalescence and therefore accept our second hypothesis. While the observed mycobiome nestedness indicates strong community coalescence in form of migration of soil fungi onto the plastisphere, the distinct dominance patterns of pathogens imply a dynamic exchange of pathogens emanating from the plastisphere (Fig. 1c, g). Falling leaves feed global fungal coalescence processes by introducing plant-associated species into the soil in a cyclic equilibrium, thereby influencing planetary nutrient cycling^{67,69}. Assuming a global plastisphere mycobiome, deposited MP could drive the coalescence of fungal communities on a similar scale but with unpredictable dynamics, due to its ubiquity, distinctiveness, persistence, and sheer numbers, especially in open areas such as savannas, deserts, oceans, and parts of the anthroposphere. The coalescence of pathogens may be a consequence of plastic pollution that has the potential to alter the proportion, identity, and diversity of fungal pathogens in virtually any environmental compartment.

The NCM⁴⁵ successfully predicted most fungal distributions across local communities in both compartments (Fig. 4), following the basic trend of abundant taxa being widespread in the MC⁷⁰, consistent with neutral theory^{37,38}. Passive dispersal and ecological drift alone explained a considerable proportion of mycobiome structure on plastic and in soil. This finding is consistent with the results of Sun et al.⁷² and Zhu et al.¹⁶, who, using the same model, found that epiplastic bacteriomes in aquatic and terrestrial systems, respectively, were mainly formed by neutral processes. However, we found several taxa that deviated from neutral predictions (Fig. 5) and are thus likely to be under environmental selection pressure (positive/negative)⁴⁶, which was also the case for certain soil bacteria being impacted by the plastic type¹¹. Therefore, we partially reject our third hypothesis of balanced complementary interplay by remarking that plastisphere mycobiomes are strongly influenced by stochasticity. In general, taxa that occurred less frequently than predicted (below-partition) are likely to be selected against by the respective compartment^{46,71}. In contrast, taxa that were more widespread than expected (above-partition) are likely to be selected by the compartment^{46,71}. Specifically, epiplastic taxa in the above-partition are likely to show plastiphily, i.e., adaptation to the plastic habitat with correspondingly increased ecological fitness. Plastiphilic fungi can thus systematically colonise plastics, form colonies, and successfully reproduce in the plastisphere¹⁰. Importantly, the non-neutral partitions of plastic and soil differed significantly in composition and ecology (Fig. 5), proving compartment-specific selection. Given the intrinsically similar environmental conditions, the differential selection must be due to substrate particularities as previously proposed^{11,16}, with MP artificially widening the fungal niche. Furthermore, plastiphilic fungi formed the most equitable and second most diverse partition

(Fig. 5c), suggesting that plastiphily plays an important role in the emergence of fungal community diversity in terrestrial systems.

Our co-occurrence network analysis revealed numerous distinct correlations among fungi in soil and on MP (Fig. 6a, Table 1), demonstrating the importance of selective forces in the assembly of mycobiomes and implying the emergence of fundamentally different interaction landscapes on MP. The lower overall size and average path length of the plastisphere network indicates a compact network with 'small world' properties, implying rapid material and information transport among co-occurring species^{73,74}, while the higher clustering coefficient suggests a more complex network with stronger interactions than in soil^{47,48}. Overall, however, the degree to which these correlations represent real-world organismic interactions remains to be investigated. Interestingly, plastiphilic fungi were overrepresented in an exclusively positive network (Fig. 6b), suggesting that well-adapted, presumably active fungi¹⁰ interact synergistically in assembling the plastisphere mycobiome. Conversely, the higher geodesic distance and number of neutrally assembled taxa in the soil network could mean that the soil fungi are mainly dormant spores from a historic contingent. However, both networks showed relatively high modularity and average degree, indicating distinct functional groups, and realised niches⁷⁵ in both compartments. As expected, both networks contained predominantly positive edges (Fig. 6a), which is consistent with the stress gradient hypothesis^{43,76}, according to which facilitation and niche sharing prevail over competition in harsher environments such as the investigated low-water, oligotrophic, and UV-irradiated ferralsols⁷⁷. Positive networks, however, are unstable due to possible fitness interdependences, resulting in rapid regime shifts and low resilience of both MCs when perturbed⁷⁶. While the keystone taxa in soil were neutrally assembled saprotrophic or unidentified fungi, plastisphere keystones were abundant, primarily plastiphilic pathogens with the capability to biodegrade plastics (Fig. 6b, c). Apparently, metabolically versatile pathogens such as *Phoma* and *Fusarium*^{78–80} drive plastisphere assembly based on trophic interactions as 'keystone plastic degraders'. By co-opting their repertoire of degradative traits^{78,80} to modify or degrade polymer components (i.e., 'exaptative plastibiome'), these pathogens may facilitate co-colonisation, co-degradation and co-metabolisation of MP by other members of the community, as previously described for other substrates^{81,82}, ultimately enabling plasticlastic commensalism and syntrophy. It has been shown that some microbes can completely degrade, assimilate, and mineralise at least certain plastics under experimental conditions^{83–85}, a feat that is yet to be observed in nature. The co-selection of fungi into the plastisphere based on their biodegradability may indicate that fungal metabolism has already become better adapted to plastic polymers. Given MP in the deep sea, atmosphere, and Arctic ice⁸⁶, the presumed evolving trophic plastiphily of fungi offers a vague but real prospect for future bioremediation of plastic if we ever stop the emission of plastic waste into the environment.

As hypothesised, the indices used revealed that MP selects pathogens more strongly than members of any other guild (Fig. 7). Moreover, selection of pathogens was much stronger in the plastisphere than in the soil, where it did not even exceed MC average (Fig. 7b, c). Remarkably, this relation was observed for all ecological and phylogenetic groups of pathogens considered (Fig. 7a–c, Supplementary Table 22). Plastic is therefore arguably the more suitable habitat for fungal pathogens in the investigated systems compared to soil. Among the plastiphilic fungi, we found numerous opportunistic human pathogens with cosmopolitan distribution and high ecological adaptability (the CAO group), including allergenic, toxigenic, and pathogenic black fungi like *Alternaria*, *Bipolaris*, *Curvularia* and *Epicoccum* (Supplementary Data 1). Melanised opportunists, primarily Dothideomycetes, also dominated the plastic-specific mycobiome (Fig. 3). These versatile pathogens infect crops²⁴, livestock and domestic animals⁸⁷, cause building damage⁸⁸ and a spectrum of opportunistic phaeohyphomycoses in humans⁸⁷. Several polymorphic cryptococcal yeasts, such as *Hannaella* and *Papiliotrema*, which are known to cause a range of potentially fatal cryptococcoses⁸⁹, were in fact only selected by plastic. In addition, polyextremotolerant black yeasts of the genera *Exophiala* and *Knufia*⁹⁰ were exclusive to plastic here. These pleomorphic fungi are common in indoor plastic niches like dishwashers³² and water storage tanks³¹ and able to cause systemic infections in humans^{31,90}. Alarmingly, we found some of the most important human pathogens on plastic waste, namely *Candida*, *Cryptococcus* and *Aspergillus fumigatus*, which are responsible for millions of life-threatening respiratory and cerebral infections each year¹⁹. The observed exclusive selection of many of these primary human pathogens^{19,53} by MP (Fig. 7), however, represents an ecological paradox. According to competitive exclusion theory⁹¹, selection of these fungi outside their primary niches, e.g., the human body^{19,35,53}, is highly unlikely as they will be outcompeted and eliminated by other better adapted species in the environment⁹¹. Therefore, it could be argued that the plastisphere resembles their ancestral habitat so closely that it represents a surrogate primary niche for these pathogens outside the host where they can thrive and evolve, presumably convergently. Considering that most plastiphilic pathogens are not only cosmopolitan but also some of the most abundant soil fungi worldwide⁷⁰, we argue that selection of fungal pathogens may become a global emergent consequence of plastic pollution.

According to the 'exaptation theory'^{92,93}, virulence traits of opportunistic fungi are also present in non-opportunistic species, as they have originally evolved for different purposes, i.e., environmental stress tolerance, and thus inherently exhibit 'dual-use'. Our meta-analysis supports this hypothesis, as we found that all GVTs are present in both opportunistic and non-opportunistic fungi (Supplementary Data 1). Pathogenic and opportunistic groups, however, showed systematically and significantly higher virulence potential than their non-pathogenic or non-opportunistic counterparts in both compartments, with virulence scores of epiplastic

pathogens being tentatively higher than those of soil equivalents (Supplementary Fig. 1, Supplementary Table 22). Although it is axiomatic that pathogens are more virulent than non-pathogens, no inferences about *in vivo* pathogenicity or virulence can be made from the presented data. Virulence traits and potentials are linked to both pathogenic lifestyle and opportunism, with oligotrophy and extremotolerance showing the strongest association (Fig. 7b). This is consistent with previous genomic studies that found a strong link between extremotolerance and opportunistic fungal pathogenicity²⁸. Our results show that plastiphily correlates with generic virulence potentials in fungi (Fig. 7e). Therefore, adaptation to the plastisphere appears to be linked to the number or coincidence of fungal GVTs. The observation that GVTs are strongly linked to both opportunism and to plastiphily supports the hypothesis that phenotypes relevant to fungal pathogenicity are shaped by plastisphere selection pressures. This corresponds to the 'accidental virulence' hypothesis⁹³ and the concept of the 'environmental virulence school'³⁶ according to which exposure to selection pressures in the environment outside the host most likely result in (pre)adaptations with benefits for pathogenesis. Although caution is required when inferring phenomena from patterns in ecology^{51,52}, it is evident that all GVTs considered are associated with both opportunistic pathogenicity and affinity for plastics, which has even been shown using several pathogens found in this study to be selected by or exclusive to plastic^{28,32,34}. Therefore, the plastisphere must be considered a 'school for virulence', where plastiphilic fungi can 'exercise' their exaptations toward higher virulence potentials ('plastiphilic virulence') (Fig. 8).

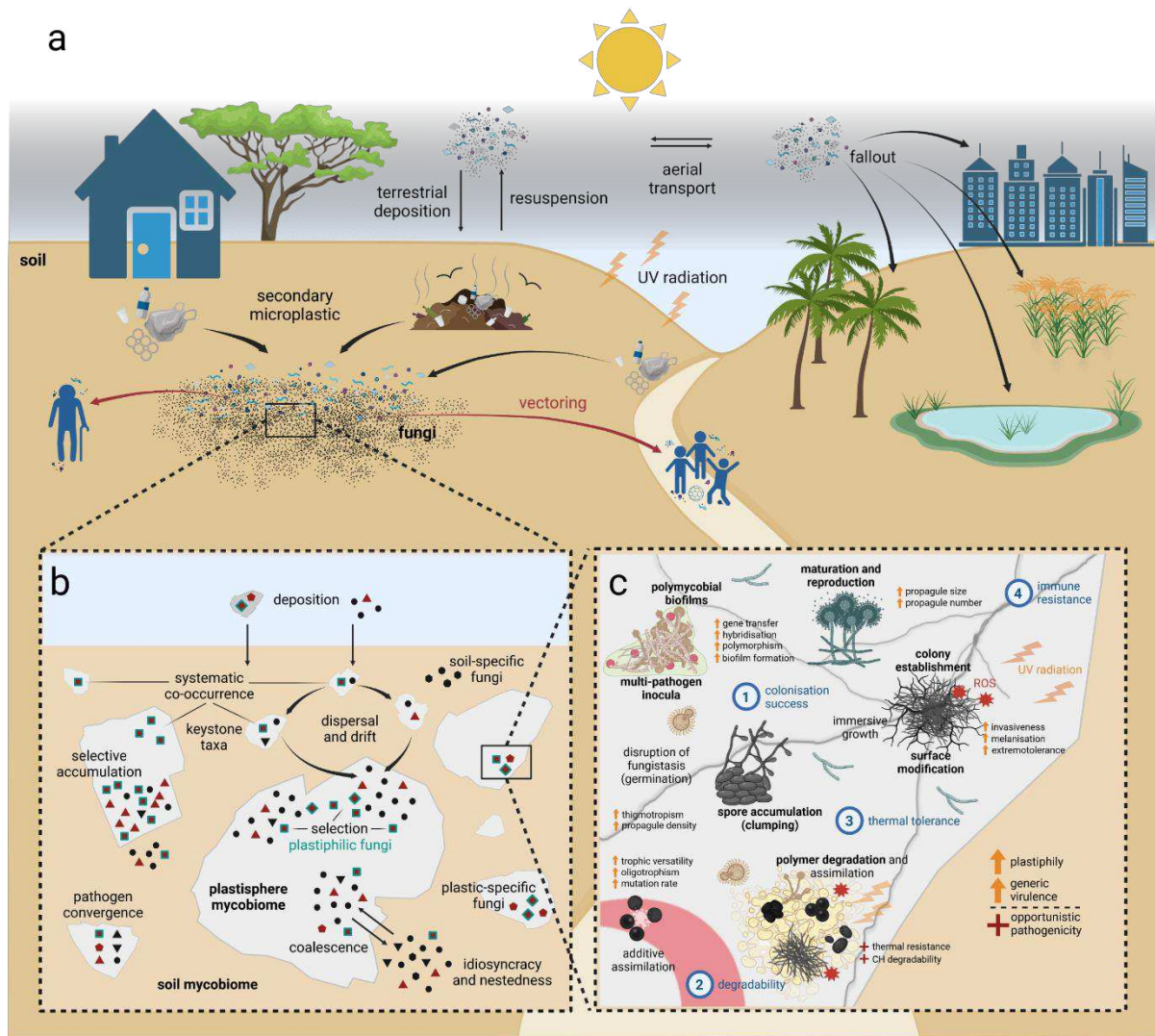


Fig. 8 Microplastic cycle and the ‘fungal niche expansion’ postulate. **a** Extract of the terrestrial microplastic cycle according to the global odyssey of plastic pollution¹⁰⁶ extended by the fungal-plastic reciprocity as observed in this study. Along the cycle, MP is permanently habitat, interaction landscape and vector of associated fungi and therefore important driver of fungal ecology, biogeography, and epidemiology. **b** Origin and ecology of epiplastic mycobiomes in the soil environment according to this study and Gkoutselis et al.¹⁰. Each symbol type represents a distinct fungal species. Plastisphere mycobiomes on deposited microplastics in soils, consisting of non-pathogenic (black symbols), and several pathogenic species (red symbols), assemble predominantly via dispersal and drift (neutral processes) from the MC (symbols without frame = neutrally assembled fungi). Several, mainly pathogenic fungi, however, are selected by MP and are therefore plastiphilic (symbols with cyan frame) or appear to co-select each other into the plastisphere through biotic interactions (systematically co-occurring species, especially keystone taxa) (deterministic processes). The resultant epiplastic communities show a high diversity (pathogen convergence) and selective accumulation of certain pathogens. Despite compositional and ecological idiosyncrasies of the different mycobiomes, with diverse plastic- and soil-specific fungal taxa, a large proportion of the plastic-associated species overlap with the planktonic ones in the soil (nestedness), indicating intercompartmental coalescence. **c** ‘The plasticsphere virulence school’. Fungal plastiphily is directly linked to the incidence of symplesiomorphic traits in fungi that, *inter alia*, (co)determine virulence in humans, such as melanisation, polymorphism, extremotolerance and oligotrophy (exaptations). Adapting to the plastisphere will therefore result in the ‘fine-tuning’ of these and other relevant traits through natural selection, eventually increasing the generic virulence potentials of epiplastic pathogens in humans (plastiphilic virulence). According to the ‘host invasion trait congruency’ theorem, exposure of fungi to the plastisphere micro-niches will convergently favour evolutionary (pre)adaptations to all four criteria relevant for the emergence of opportunistic pathogenicity in humans (1–4)³⁵. Ecological phenomena found in this study and in Gkoutselis et al.¹⁰ is

highlighted in bold. Many other empirically grounded mechanisms have been adapted and proposed. (↑) = increase, (+) = acquisition/emergence and CH = hydrocarbons. Created with biorender.com.

In summary, our results support the notion that plastic pollution is indeed closely linked to the occurrence of fungal infections by providing reservoirs, vectors and an evolutionary 'training ground' for pathogenic fungi in terrestrial systems, which could favour all four criteria relevant for the emergence of fungal pathogenicity³⁵ (Fig. 8c). The assembly of a diversity of pathogens (pathogen convergence) in locally concentrated polymicrobial inocula on MP, together with selective accumulation¹⁰ and pathogen coalescence effects will certainly increase the number and density of infectious agents (propagule size), while particle dispersibility, invasiveness and persistence could increase the range and number of transmission events (propagule number), synergistically enhancing pathogen pressure⁹⁴. This in turn would, according to the principles of biological invasions⁹⁵, increase both reach and colonisation success of adherent pathogens (criterion 1). In oligotrophic soil environments such as savannas, deserts, and Arctic valleys, where plastic debris are 'islands of carbon', plastiphilic fungi may 'learn' to overcome longer periods of nutrient deprivation by utilising trace additives on the particle surface or even modify metabolic pathways to exploit the polymer itself. Potent hydrocarbonoclastic species could evolve into 'zombie degrader fungi' capable of assimilating neurotransmitter and other aromatic compounds of the human host^{54,96} (criterion 2). In the very same environments, MP in topsoil, especially in equatorial regions, are exposed to high UV radiation and yet are becoming more attractive to fungi as carbon residues become more accessible through polymer weathering⁹⁷. Melanised, extremotolerant plastic degraders would thrive under those conditions and quickly adapt to high temperatures by evolving thermal exaptation into thermal tolerance. On these 'warm little polymers', plastiphilic fungi may overcome the thermal exclusion zones imposed by mammal endothermy and homeothermy^{35,98}, and thus acquire the ability to thrive inside the human body (criterion 3). Finally, exposure to immense oxidative stress during polymer weathering and biodegradation⁹⁹ may 'train' plastiphiles to resist reactive oxygen species deployed by immune cells¹⁰⁰, arming them to resist the human immune system (criterion 4) ('host invasion trait congruency' theorem). More than any other hydrophobic habitat, plastic could thus unleash the virulence potential of polyextremotolerance in fungi. Plastisphere life may further promote a pathogenic phenotype by fine-tuning thigmo responses^{36,101}, disrupting fungistasis¹⁰², favouring biofilm formation^{10,103}, and increasing rates of mutation, hybridisation, and gene transfer^{53,104}. Ultimately, plastiphilic fungal pathogens would thus become increasingly virulent to humans, while even previously non-pathogenic species could emerge as 'accidental' opportunists⁹³ once they acquired a critical repertoire of GVTs through their 'passage' of the plastisphere (Fig 8c). As global plastic pollution coincides with global warming^{24,105}, habitat fragmentation⁷⁰, desertification¹⁰⁵ and an unprecedented

number of immunosuppressed individuals^{19,21}, plastiphilic fungal virulence is likely to play an essential role in human health throughout the world in the future.

Given the strong link between virulence potential, opportunistic pathogenicity, and the selection of ubiquitous, extremely abundant, and significant human pathogens by MP, plastic pollution may contribute substantially to the occurrence of fungal infections worldwide and might have been doing so for a long time. Therefore, there is an urgent need to classify microplastics as a global health factor. Taken together, our findings provide first insights into the origins of epiplastic mycobiomes and the mechanisms behind the emergence of fungal plastiphily and demonstrate the important role of plastic debris in the ecology and evolution of pathogenic fungi. This new knowledge is not only crucial for developing strategies to combat both global burdens – fungal infections and plastic pollution – but also forms the basis for a deeper understanding of the biodiversity, ecology, functionality, and adaptability of plastic-colonising soil microbes in an increasingly plasticised world.

Material and Methods

Sampling and selective subsampling. The environmental samples were identical to those used in the previous study by Gkoutselis et al.¹⁰. Briefly, the study design involved the selection of five different sites with high levels of plastic contamination and human activities within the town limits of Siaya (Siaya County, Western Kenya) followed by the random collection of five samples from the contaminated topsoil of each plot (total n = 25). The sites included two open landfills, a roadside, a marketplace, and a courtyard. Further details on the sampling procedure, the contextual data collection and the site descriptions are provided in Gkoutselis et al.¹⁰. All relevant data of the sampling campaign have been deposited in DiversityCollection¹⁰⁷ at the Staatliche Naturwissenschaftliche Sammlungen Bayerns (SNSB), Germany, in accordance with the FAIR++ guiding principles for scientific data management⁴⁹. To double the subsample pool, biological replicates were compiled to each subsample, yielding a total of 50 plastic and soil subsamples, respectively (total n = 100). To ensure comparability, the operational design and methodology of Gkoutselis et al.¹⁰ were strictly followed. The size of the plastic debris ranged from 1 to 28 mm with the majority (84%) of particles < 5 mm. Therefore, all plastic subsamples were considered MP subsamples². As before, MPs were combined into subsamples regardless of shape and polymer type. All subsamples were frozen at -20 °C until further processing.

DNA extraction, PCR and Illumina sequencing. All steps were conducted as previously described¹⁰. Briefly, total metagenomic DNA from soil and plastic subsamples was extracted

using the NucleoSpin® Soil kit (MACHEREY NAGEL, Düren, Germany) according to the manufacturer's instructions after initial adjustments. NanoDrop® ND-1000 UV-Vis spectrophotometer (Thermo Fisher Scientific, Waltham, United States) and 0.8% agarose gel electrophoresis were used to determine the quality and concentration of the isolated DNA. ITS1 and ITS2 regions were amplified by nested PCR based on ITS1F and ITS4 primers, each containing a unique combination of TAG and INDEX sequences, part of the Illumina sequencing primer, and P5 and P7 adapters, respectively, to generate dual-indexed amplicons. An enzymatic purification step (ExoSAP digestion) was interposed to remove or deactivate excess reactants prior to PCR2. Cycling and purification were performed in a thermocycler (BioRad, Hercules, USA). The combination of TAG (PCR1) and INDEX (PCR2) sequences allowed the generation of unique sequence patterns for each sample for multiplexing of amplicons. Quality control and pooling were performed as previously described¹⁰. Gel bands were visualised using Gel Doc™ EZ gel imaging system (BioRad, Hercules, USA). Amplicons were paired-end sequenced on an Illumina MiSeq 3000 sequencer (Illumina, Inc., San Diego, USA) at the Department of Genetics, LMU Munich, Germany.

Bioinformatics. Barcodes (TAG sequences) were removed from the obtained raw reads and the reads were demultiplexed using `extract_barcode.py` and `split_libraries_fastq.py` as implemented in QIIME¹⁰⁸. The demultiplexed reads were then imported to the QIIME2 pipeline¹⁰⁹ and remaining adapter and primer sequences were trimmed, according to the applied sequences motives, using the `cutadapt` tool¹¹⁰. Quality filtering, including removal of chimeric sequences and dereplication of exact sequence variants was performed using the DADA2 plugin under QIIME2¹⁰⁹ based on a maximum expected error of 2 and a minimum fold abundance of parent sequences of 1 over potential chimeras. OTUs were clustered from the amplicon sequence variants (ASVs) from the DADA2 output using VSEARCH and a threshold of 97% sequence similarity. Taxonomy assignment was performed using the feature classifier `classify-sklearn` command implemented in QIIME2 and a Naïve Bayesian classifier previously trained on the UNITE (v.8.0) dynamic database¹¹¹ with a chunk size of 20,000. OTU representative sequences were then taxonomically classified using a threshold of 97% sequence similarity and a minimum confidence score of 70%. Samples containing less than 1,000 sequences¹¹² and singletons⁷¹ were removed from the dataset.

Neutral community model. To determine the processes driving mycobiome assembly, we implemented the neutral community model developed by Sloan et al.⁴⁵, which is an adaptation of neutral theory adjusted to large microbial populations. The model predicts the relationship between frequency with which taxa occur in a set of local communities (fungal communities of individual plastic or soil subsamples) and their abundance across the wider metacommunity (fungal communities of all plastic or soil subsamples). In principle, the NCM predicts that taxa

that are abundant in the MC will be widespread as they are more likely to be randomly dispersed among different sampling sites, whereas rare taxa are more likely to be lost at different sites due to ecological drift (i.e., stochastic loss and replacement of individuals). Unlike many other contemporary neutral models, such as the unified neutral theory of biodiversity³⁷, the NCM used here does not incorporate the process of speciation. Here, dispersal among communities is estimated by Nm , which determines the correlation between observed occurrence frequency of fungi (proportion of local communities comprising each OTU) and their abundance in the MC (mean relative abundance across all local communities), with N describing the MC size and m being the migration rate. The migration rate represents the probability that a random loss of an individual in a local community will be replaced by dispersal from the MC, as opposed to reproduction within the local community. The parameter R^2 indicates the overall fit to the neutral model⁴⁵, assessed by comparing the sum of squares of residuals with the total sum of squares⁴⁶. To test whether incorporating ecological drift and dispersal limitation improve the fit of the model beyond just random sampling from the MC⁴⁵, we compared the NCM fit with the fit of a binomial distribution model. To compare the fit of the neutral and binomial model, the Akaike information criterion (AIC) was calculated and compared for each model^{16,46}. Fitting of the free parameter m was performed in R (R 4.2.0)¹¹³ using non-linear least-squares fitting from the `minpack.lm` package¹¹⁴. Wilson score interval from the `HMisc` package in R¹¹⁵ was used to calculate the binomial 95% confidence intervals. Computation of the AIC was also conducted in R (R 4.2.0)¹¹³. Calculation of 95% confidence intervals around all fitting statistics was done by bootstrapping based on 1,000 replications (Supplementary Data 2) We used the R code of Burns et al.⁴⁶ for all analyses. The NCM was applied separately to soil and plastic as well as to the combined dataset on rarefied OTU matrices. To analyse deviations from the neutral predictions, OTUs from each dataset were separated into three partitions depending on whether they occurred more frequently than (above-partition), less frequently than (below-partition) or as frequently as predicted by the NCM (neutral-partition) based on 95% confidence intervals. Ecologically, fungi above or below prediction are therefore considered as being actively selected for (selection) or against (exclusion) by the target compartment, respectively (deterministic processes), while taxa within prediction are considered randomly assembled (neutral processes). Fungi in the above-partition of the plastisphere were defined as 'plastiphilic' or 'plastiphiles'. Mean relative abundance, as well as observed and predicted occurrence frequency of each OTU of each compartment can be found in Supplementary Data 2.

Meta-analysis. Only identified genera in the combined rarefied dataset were subjected to meta-analysis. For all other phlotypes, 'unidentified' was set as default for any annotation or assignment. For inferences about physiology, ecology, biogeography, and relevance to humans, fungi were annotated with four relevant traits, namely growth form, distribution,

adaptability, and opportunism (Supplementary Data 1). Growth form was annotated using the comprehensive FungalTraits database¹¹⁶, with the categories 'filamentous', 'yeast-like' (= yeasts and dimorphic yeasts) and 'zoosporic' (= any zoosporic group). Distribution was determined using the GlobalFungi database¹¹⁷, with genera classified according to the number of continents on which corresponding specimens have been recorded into 'endemic' (1), 'widespread' (2–5) and 'cosmopolitan' (>5). Adaptability was categorised according to the number of possible lifestyles (FungalTraits)¹¹⁶ or guilds (FUNGuild)¹¹⁸ into 'low' (1 or 2), 'high' (3 or 4) and 'extremely high' (>4). Finally, opportunism (= opportunistic human pathogenicity^{10,28}) was determined by querying FungalTraits¹¹⁶, FUNGuild¹¹⁸, Encyclopedia of Life (EOL)¹¹⁹, Index Fungorum (<http://www.indexfungorum.org>) and all results (up to a maximum of 100) of a PubMed search using the respective genus name. Any relevant record of a specimen as an aetiological agent for an infection or disease in humans qualified the corresponding genus as 'opportunistic', otherwise it was considered 'non-opportunistic'. Generally, in the absence of entries for any trait in the relevant database(s), the appropriate literature was consulted instead. To further elaborate on the functionalities, epidemiology, and relevance to human health, recovered fungi were assigned to three distinct ecological groups, namely guilds, cosmopolitan adaptable opportunists (CAO) and so-called “hidden killers”¹⁹ (Supplementary Data 1). Guilds were assigned by consolidating the entries for 'primary_lifestyle' in FungalTraits¹¹⁶. As we focused primarily on human pathogenicity in this study, we chose to simplify the assignment by using more generic versions of the classical fungal guilds¹¹⁸, where 'pathogen' = any pathogen or parasite, 'saprotroph' = any saprotroph, sooty mold, endophyte, or epiphyte, and 'mutualist' = any mycorrhizal and lichenised group. Missing records in the database(s) were compensated by relevant literature. CAO were those fungi that showed 'cosmopolitan' distribution, 'high' or 'extremely high' adaptability and 'opportunistic' potential according to our trait data annotation. *Hidden killers* were named and defined according to the study of Brown et al.¹⁹ and consisted of only the most significant human pathogens causing opportunistic invasive mycoses, in our study *Aspergillus*, *Candida* and *Cryptococcus*.

To approach fungal virulence potential, we compiled and annotated four well-described, commonly observed and highly conserved autecological and physiological traits that contribute to the emergence of pathogenicity and determine the virulence of a vast diversity of phylogenetically distinct fungi (Supplementary Data 1), namely melanisation⁵⁶, polymorphism⁵⁷, extremotolerance²⁸ and oligotrophy¹²⁰. We therefore term these traits 'generic virulence traits' (GVTs). Data for each GVT were acquired from FungalTraits¹¹⁵, GlobalFungi¹¹⁶, EOL¹¹⁸, and all results (up to a maximum of 100) of a Google Scholar search using the respective genus and trait name (e.g., '*Curvularia*, melanisation'). It is impossible to make a general statement at the genus level regarding the presence of any of these GVTs, as

even plesiomorphic and ancestral traits of a genus could have been lost in individual species¹²¹. At the same time, it is impossible to annotate such traits for each species identified in a metagenome study due to the lack of appropriate databases, publications, and a generally limited number of observed traits^{51,52}. In our study, each GVT was only assigned to a particular genus when at least three different species of that genus were found to have the trait. The four GVTs were approximated as follows: (1) Melanisation: based on the pigmentation of vegetative structures (e.g., hyphae, yeast cells), fungi were categorised into 'hyaline', 'melanised' or 'pigmented' (unspecified or non-melanin pigmentation). (2) Polymorphism: fungi with the ability to shift from a multicellular hyphal to a unicellular yeast growth form⁵⁷ were classified as 'dimorphic', all other ones were considered 'monomorphic'. (3) Extremotolerance: fungi were classified as 'extremotolerant', if they met any of the several criteria proposed by Gostinčar et al.²⁸, including superficial growth in exposed habitats (e.g., rock-inhabiting fungi) subjected to varying temperatures, dryness, and solar irradiation; lichenised lifestyle (mycobiont in an algal-fungus combination); osmotolerant or osmophilic from dry indoor habitats, desert soils, and similar habitats and halotolerant or halophilic fungi from hypersaline and highly contaminated environments; and psychrotolerant or psychrophilic fungi from deep ocean waters and polar regions and similar habitats. All other fungi were classified as 'non-extremotolerant'. (4) Oligotrophy: fungi were classified as 'oligotrophic', if they met any of the criteria described by Gostinčar et al.¹²⁰, including growth in extremely nutrient-poor natural habitats and environments, such as rock surface or subsurface, Arctic dry valleys, deserts, and high mountain areas; growth in (oligotrophic) anthropogenic habitats such as monuments, concrete walls, biofilters and other indoor habitats; growth on silicon, metals, glass and on a variety of more or less durable organic surfaces, including plastic materials and similar polymers; or biodegradation of complex, non-natural compounds, such as phenolic hydrocarbons, TNT, radioactive materials, plastics, etc.^{54,78–80}. All other fungi were classified as 'non-oligotrophic'. To achieve a simple parametrisation, we introduced a 'virulence score' based on the number of GVTs found in each phylotype (0–4). In a real virulence context, this score boasts only a vague informative value, due to the intrinsic methodological imprecision and the complex context-dependency of fungal infections²⁸. Instead, we argue that it constitutes a suitable reductionistic approximation metric for fungal virulence potentials and an ordinal scale for eco-evolutionary projections, as similar presence, identity, and coincidence of the described GVTs translate into similar options regarding ecological niches and positions in the evolutionary landscape^{122,123}.

Abundance fractions. To accurately analyse the architecture of the target mycobiomes, we defined thresholds for relative abundance and frequency of occurrence to classify all OTUs of each MC into six different abundance fractions as previously described⁷¹. Therefore, for each compartment, samples were grouped according to sites to form site-specific (bulk) samples.

(1) always abundant taxa (AAT) were defined as OTUs with an abundance of $\geq 1\%$ in all samples; (2) conditionally abundant taxa (CAT) were defined as taxa with an abundance of $\geq 0.01\%$ in all samples and $\geq 1\%$ at least one sample; (3) moderate taxa (MT) were defined as OTUs with an abundance between 0.01 and 1% in all samples; (4) conditionally rare and abundant taxa (CRAT) were defined as OTUs ranging from rare ($< 0.01\%$) to abundant ($\geq 1\%$); (5) conditionally rare taxa (CRT) were defined as OTUs with an abundance of $> 1\%$ in all samples and $< 0.01\%$ in at least one sample; and (6) always rare taxa (ART) were defined as OTUs with an abundance $< 0.01\%$ in all samples.

Statistical analysis. Beta diversity analysis using Hellinger distance, ANOSIM and PERMANOVA (Fig. 5a) were performed with the Primer 7 software package and the add-on package PERMANOVA+ (PRIMER-e Ltd, Plymouth, United Kingdom)¹²⁴. All other statistical analyses and data visualisations were performed in R (R 4.2.0)¹¹³, using the *vegan*¹²⁵, *phyloseq*¹²⁶, *ggplot*¹²⁷, *microbiome*¹²⁸, *reshape2*¹²⁹, *ggpubr*¹³⁰, *spaa*¹³¹ and *dplyr*¹³² packages. Correlation analyses used in this study were Spearman tests conducted in R (R 4.2.0)¹¹³.

Phylogenetic analysis. To demonstrate the evolutionary distance of plastic-specific taxa, a phylogenetic tree was constructed, including only phylotypes from the rarefied dataset identified at least to the genus level ($n = 84$). Nucleotide sequences were aligned using MUSCLE¹³³ as implemented in MEGA11¹³⁴. Phylogenetic relationships were inferred using the maximum likelihood method, and Tamura-Nei distance¹³⁵ as a substitution model and a bootstrap method based on 1,000 replications as a phylogeny test. The phylogram was calculated in MEGA11¹³⁴ and visualised using iTOL version 6¹³⁶.

Selection indices. Positive selection of fungi was quantified separately for epiplastic and soil fungi based on the results of the NCM using four distinct indices:

- (I) Within-group selection (WGS), i.e., the proportion of above OTUs in a group to the total OTUs in that group:

$$WGS_i = \frac{n_{i\ ap}}{n_i}$$

- (II) Across-metacommunity selection (AMS), i.e., the proportion of above OTUs in a group to the above OTUs in the total MC:

$$AMS_i = \frac{n_{i\ ap}}{n_{j\ ap}}$$

- (III) Selection index (SI) calculated from the product of the WGS and AMS of each group:

$$SI_i = WGS_i \times AMS_i$$

- (IV) Times average selection (TAS), i.e., the ratio of the groups' WGS and the proportion of above OTUs in the MC:

$$TAS_i = \frac{WGS_i}{\frac{n_{j\ ap}}{n_j}}$$

where n is the number of OTUs, i is the group under examination, j is the corresponding MC and ap refers to the respective above-partition. All calculated values can be found in Supplementary Table 21 and 22 and Supplementary Data 2.

Network analysis. For co-occurrence network construction, the most frequent 10% of OTUs of each MC were chosen. The Spearman correlation matrix was calculated using the psych package¹³⁷ in R. To highlight the most important relationships, only strong positive ($\rho > 0.6$) or strong negative ($\rho < -0.6$) and highly significant ($p < 0.01$) correlations were used for network construction^{48,74,82}. Network analysis and visualisation was conducted in Gephi 0.9.7¹³⁸. The network properties, clustering coefficient, modularity, average path length, average normalised degree, and positive ratio were included in the analyses (Table 1). OTUs with maximum degree and betweenness centrality were considered keystone taxa^{48,74}. Phylotypes in the plastisphere network were annotated with the trait of plastic biodegradability (Supplementary Data 4). Relevant data were obtained from the sources mentioned in Ekanayaka et al.⁷⁹ and with the phrases used there extended by the respective genus name.

Data availability

Sequences from community barcoding are linked under BioProject accession number PRJNA705067. Environmental samples are stored in the collection at the Mycology Department, University of Bayreuth. All other relevant data is available upon request.

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Acknowledgements

This study was funded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) – SFB 1357-39197756; subproject C04. We wish to thank Derek Peršoh (Faculty

of Biology, Chemistry and Earth Sciences, University of Bayreuth) for his help in the optimisation of sequence data processing, Stella Franca Rösch (Chair of Marketing and Innovation, University of Bayreuth) for her support in graphic formatting and Luca Söhnle (Dept. of Mycology, University of Bayreuth) for his assistance in the subsampling process. Gratitude is also dedicated to the Ministry of Agriculture and Rural Development Kenya Plant Health Inspectorate Service (KEPHIS) for providing the export permit (Phytosanitary certificate no. 207/2019), the Bayerische Landesanstalt für Landwirtschaft, Institut für Pflanzenschutz for issue of the import authorisation (IPS 4a-7322.451) and the ITCER e.V. (<https://itcer.org>) for on-site support.

Author contributions

G.G. and S.R. carried out the laboratory work and wrote the original draft. A.B. carried out Illumina sequencing. Sequence data processing was done by J.H. G.G., S.R. and J.H. performed the statistical analyses. G.G. and G.R. were responsible for the meta-analysis. The creation of the final version of the manuscript was supported by all authors. M.A.H. and G.R. acquired funding, designed the study, and supervised the project. G.R. collected and provided the environmental samples from Kenya.

Competing interests

The authors declare no competing interests.

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