

## 1 **Towards understanding diversity, endemicy and global change vulnerability of soil fungi**

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117

## 118 **Summary**

119 Fungi play pivotal roles in ecosystem functioning, but little is known about their global patterns of  
120 diversity, endemism, vulnerability to global change drivers and conservation priority areas. We  
121 applied the high-resolution PacBio sequencing technique to identify fungi based on a long DNA  
122 marker that revealed a high proportion of hitherto unknown fungal taxa. We used a Global Soil  
123 Mycobiome consortium dataset to test relative performance of various sequencing depth  
124 standardization methods (calculation of residuals, exclusion of singletons, traditional and SRS  
125 rarefaction, use of Shannon index of diversity) to find optimal protocols for statistical analyses.  
126 Altogether, we used six global surveys to infer these patterns for soil-inhabiting fungi and their  
127 functional groups. We found that residuals of log-transformed richness (including singletons) against  
128 log-transformed sequencing depth yields significantly better model estimates compared with most  
129 other standardization methods. With respect to global patterns, fungal functional groups differed in  
130 the patterns of diversity, endemism and vulnerability to main global change predictors. Unlike  $\alpha$ -  
131 diversity, endemism and global-change vulnerability of fungi and most functional groups were  
132 greatest in the tropics. Fungi are vulnerable mostly to drought, heat, and land cover change. Fungal  
133 conservation areas of highest priority include wetlands and moist tropical ecosystems.

134

## 135 **Introduction**

136 Human activities affect nearly all habitats through changes in climate and land-use, which in turn alter  
137 vegetation cover and composition. These changes negatively impact many species that have narrow  
138 environmental tolerances and limited dispersal capacity across anthropogenic landscapes (Schulte to  
139 Bühne et al. 2020). Anthropogenic impacts most strongly affect endemic species – i.e., taxa with small  
140 distribution ranges and narrow ecological niches (Brook et al. 2008). Diversity of endemic plants and  
141 animals is higher in areas characterized by historical stability, high precipitation, environmental  
142 heterogeneity, and insularity. Unfortunately, these areas usually coincide with major human  
143 degradations of the environment (Kier et al. 2009; Stein et al. 2014; Sandel et al. 2020).

144 Unlike the situation with plants and animals, global patterns of fungal diversity, endemism and  
145 vulnerability to environmental change remain virtually unknown (Cameron et al. 2019; Guerra et al.  
146 2021b; but see Talbot et al. 2014; Davison et al. 2015). This is alarming, given the fundamental roles  
147 that fungi play in carbon and nutrient cycling processes (Wardle & Lindahl 2014; Crowther et al. 2019).  
148 Comparative studies have indicated that aboveground and belowground biodiversity are driven by  
149 different environmental predictors at local and global scales (Cameron et al. 2019; Le Provost et al.  
150 2021). This suggests differential responses of macro- and microorganisms to land use and climate  
151 changes (Guerra et al. 2021b). As for plants and animals, soil fungal communities are likely vulnerable  
152 to global change drivers. For instance, high-temperature stress (Malcolm et al. 2008; Barcnas-Moreno  
153 et al. 2009; Morgado et al. 2015; Misiak et al. 2021) and prolonged drought (Schmidt et al. 2017; de  
154 Vries et al. 2018) can alter fungal growth, functionality and community composition. Likewise, changes  
155 in land use that result in habitat fragmentation may lead to shifts in prevalence of pathogenic,  
156 mutualistic, and free-living fungal groups (Brinkmann et al. 2019; Makiola et al. 2019; Le Provost et  
157 al. 2021; Rodriguez-Ramos et al. 2021).

158 While thousands of plant and animal species are listed as threatened on the IUCN global Red List, only  
159 262 out of an estimated 2.2-3.8 million fungal species (Hawksworth & Lücking 2017) have been listed  
160 as such. The majority of these are from high-income countries in temperate regions (IUCN 2021) and  
161 are from fungal groups that make conspicuous macroscopic fruiting bodies (Cui et al. 2021). However,  
162 the vast majority of fungi produce no or inconspicuous fruiting bodies and are therefore hard to survey,  
163 which has hampered their conservation assessment (Gonçalves et al. 2021).

164 Here we used the most advanced high-resolution sequencing technology to globally survey soil fungal  
165 diversity and assess their endemism and vulnerability to global change. We hypothesized that i) the  
166 endemism of fungi is relatively higher in the tropics due to greater regional climatic stability; and ii)  
167 vulnerability of fungi to global change is highest in habitats experiencing the strongest global warming  
168 effects (polar regions) and intensive land use (dry tropics). We predicted that because of their intimate  
169 associations with other organisms, endemism and vulnerability patterns are more evident for  
170 macrofungi and biotrophic groups compared with saprotrophic microfungal groups. We then propose  
171 global conservation priorities for these ecologically pivotal fungi.

172

## 173 **Results and Discussion**

### 174 *Fungal diversity*

175 We used the recently generated Global Soil Mycobiome consortium dataset (GSMc; 3,200 plots,  
176 Tedersoo et al. 2021b) along with data from five other global soil surveys (**Fig. 1**; see methods) and  
177 international nucleotide sequence databases to determine the diversity and endemism of fungal

178 functional groups – *viz.* arbuscular mycorrhizal (AM) fungi, ectomycorrhizal (EcM) fungi, non-EcM  
179 Agaricomycetes (mostly saprotrophic macrofungi), molds, pathogens, opportunistic human parasites  
180 (OHPs, mostly thermophilic saprotrophs), early-diverging unicellular lineages (mostly chytrids,  
181 aphelids, and rozellids), and yeasts. Compared to previous meta-analytical approaches (e.g. Vetrovsky  
182 et al. 2019), our cumulative data comprise the largest available globally standardized database based  
183 on directly comparable soil sampling and long-read molecular analysis protocols. Collectively, all  
184 datasets yielded 20,182,427 fungal reads composed of 905,841 ‘species’ – operational taxonomic  
185 units (OTUs), each defined as <98% sequence similarity of the rRNA ITS barcode from all other  
186 OTUs. The genera *Tomentella* (Basidiomycota), *Penicillium* (Ascomycota), and *Mortierella*  
187 (*Mortierellomycota*) were the most species-rich (**Fig. 1**).

188 We combined machine-learning and general linear modeling (GLM) approaches to find the best  
189 predictors of fungal species richness and the Shannon index of diversity for settling the contrasting  
190 results obtained from previous global studies (Tedersoo et al. 2014; Egidi et al. 2019; Vetrovsky et al.  
191 2019). At the site scale ( $\alpha$ -diversity), the best supported results were obtained for residuals of  
192 logarithm-transformed richness accounting for sequencing depth (**Fig. 2**). Since the datasets were  
193 retrieved using different sampling design and therefore differed strongly in the inferred richness (**Fig.**  
194 **3**), we focused mainly on analyses of the largest, GSMc dataset. Total fungal richness had a broadly  
195 unimodal relationship with soil pH ( $R^2_{\text{adj}}=0.133$ ) and responded positively to vegetation age  
196 ( $R^2_{\text{adj}}=0.045$ ; **Fig. 4**). Deserts and Antarctic habitats supported the lowest richness among all biomes  
197 (**Fig. 4**). We validated the results using several datasets, in which fungal richness had a unimodal  
198 relationship with soil pH and positive response to mean annual precipitation (MAP)(**Fig. 5**). Across  
199 datasets, fungal  $\gamma$ -diversity at the ecoregion level was best explained by average MAP ( $R^2_{\text{adj}}=0.179$ ;  
200 **Fig. 6**). The differences in richness trends between  $\alpha$ -diversity and  $\gamma$ -diversity indicate that high  
201 precipitation favors niche differentiation at the regional scale, as reflected by higher turnover between  
202 sites (i.e. increasing  $\alpha$ - to  $\gamma$ -diversity).

203 Our results thus update previous patterns of  $\alpha$ -diversity decrease (Tedersoo et al. 2014) or increase  
204 (Vetrovsky et al. 2019) at high latitudes and confirm relatively lower fungal diversity in Antarctica.  
205 The latter pattern has been/can be ascribed to low plant diversity and coverage (Newsham et al. 2016).  
206 The more prominent latitudinal gradient in  $\gamma$ -diversity reflects a greater positive effect of MAP on the  
207 regional fungal species pool. Disregarding Antarctica, the lack of a global  $\alpha$ -diversity latitudinal  
208 gradient in fungi is unique among terrestrial organisms (Kinlock et al. 2018). By comparison, the  $\gamma$ -  
209 diversity patterns detected resemble those found for soil fauna (Aslani et al. 2022; Potapov et al.  
210 2022) and protistan parasites (Oliverio et al. 2020), all which show slight richness peaks in tropical  
211 latitudes. The distinctly weaker latitudinal diversity gradients of soil organisms compared with most  
212 aquatic and terrestrial macro-organisms may be related to indirect effects of temperature-related  
213 climatic variables as well as soil pH and C/N ratio as main drivers of soil habitat quality. The

214 differences may also be related to higher dispersal capacity of soil organisms who have microscopic  
215 body sizes or dispersal propagules (Soininen et al. 2013; Aslani et al. 2022).

216

### 217 *Fungal endemism*

218 To estimate relative endemism among the world's ecoregions (**Fig. 7; Table 1; see methods**), we  
219 combined indices of community similarity, uniqueness, and species ranges into an overall endemism  
220 index (see Methods). Five metrics were combined, including the number and proportion of endemic  
221 species, mean maximum geographical range of species, Jaccard index, and beta-sim index (**Box 1**).  
222 We found that endemism of all fungi peaked in moist tropical biomes and it was positively related to  
223 mean annual air temperature (MAT;  $R^2_{\text{adj}}=0.277$ ; **Fig. 8**) and soil acidity ( $R^2_{\text{adj}}=0.108$ ; **Table 2**).

224 While endemism patterns of non-EcM Agaricomycetes and AM fungi were similar to those shown  
225 for all fungi, different patterns were found for other functional groups. Endemism of EcM fungi was  
226 related to high mean annual precipitation (MAP) ( $R^2_{\text{adj}}=0.147$ ). Molds, pathogens and yeasts showed  
227 multiple endemism hotspots. Molds ( $R^2_{\text{adj}}=0.199$ ) and pathogens ( $R^2_{\text{adj}}=0.105$ ) had relatively greater  
228 endemism in strongly acidic or alkaline soils, indicating that extreme soil conditions may support  
229 unique soil biota, with limited effective dispersal across edaphically extreme habitats. Human  
230 footprint (see Methods) had a weak negative effect on endemism of all fungi ( $R^2_{\text{adj}}=0.018$ ),  
231 pathogens ( $R^2_{\text{adj}}=0.015$ ), and OHPs ( $R^2_{\text{adj}}=0.056$ ), suggesting that anthropogenic habitat loss or  
232 homogenization may affect endemic species (Finderup Nielsen et al. 2019). European ecoregions had  
233 the lowest endemism for all fungi ( $R^2_{\text{adj}}=0.065$ ), pathogens ( $R^2_{\text{adj}}=0.086$ ) and unicellular fungi  
234 ( $R^2_{\text{adj}}=0.035$ ) compared with those of other areas. Averaged current aerial bioclimatic variables better  
235 explained endemism compared with the ranges of those variables or bioclimatic variables of soil and  
236 last glacial maximum (LGM). Climate change since the LGM had a weak positive effect on  
237 endemism of molds (mean diurnal range and overall climate change:  $R^2_{\text{adj}}=0.073$ ) and OHPs  
238 (isothermality and mean diurnal range:  $R^2_{\text{adj}}=0.056$ ) but not other groups.

239 We found that patterns in fungal endemism were relatively consistent among the five individual  
240 endemism indices and that they resemble endemism patterns of vascular plants and animals, which  
241 exhibit major hotspots in wet tropical habitats (Kier et al. 2009; Barlow et al. 2018). However,  
242 endemism patterns in fungi were somewhat weaker, which may reflect the greater long-distance  
243 dispersal capacity of fungal spores relative to propagules of plants and animals (Golan & Pringle 2017).  
244 In terms of the greater macroorganism richness and endemism found in the tropics, the literature  
245 abounds with hypotheses, including narrower niche breadth, more asymmetric interactions (i.e., greater  
246 specialization), climatic stability, and more rapid evolution due to environmental energy (Vazquez &  
247 Stevens 2004; Brown 2013) in the tropics than elsewhere. Negligible effects of the LGM suggest that

248 climatic stability is not an important driver of fungal endemism, a pattern that contrasts with those of  
249 plants and animals (Rosauer & Jetz 2015). The greater phylogenetic diversity of fungi noted for the  
250 tropics (e.g. Tedersoo et al. 2018) may in part reflect tropical origins for many lineages, as well as the  
251 radiation and rapid speciation of a limited number of EcM fungal genera into higher latitude areas  
252 (Kennedy et al. 2012; Sanchez-Ramirez et al. 2015). On a global scale, plant diversity does not appear  
253 to be causally related to fungal diversity (Tedersoo et al. 2014), but there is some evidence for stronger  
254 mutualistic plant-fungal interactions related to high rainfall (Pöhlme et al. 2018). Pathogenic interactions  
255 warrant further research in this respect, given their major importance as regulators of plant diversity  
256 (Chen et al. 2019). Tropical soil fungi have relatively greater dispersal limitations (Bahram et al. 2013)  
257 and narrower distribution ranges (Tedersoo et al. 2014), suggesting that high local diversity may  
258 contribute to greater regional-scale endemism.

259

### 260 *Vulnerability of fungi to global change drivers*

261 Communities with many species at their environmental niche limits may be particularly vulnerable to  
262 local extinctions (Watson et al. 2013; Smith et al. 2020b). Thus, we evaluated the relative vulnerability  
263 of soil fungal functional groups by estimating the percentage of species occurring at their upper niche  
264 limits to three major global change drivers – land use (land cover change), heat (maximum monthly  
265 temperature), and drought (lowest quarterly precipitation). We projected to the year 2070 relative to the  
266 2015 baseline, using the average vulnerability index (Smith et al. 2020b), land use extrapolations of the  
267 LUH2 global dataset (Hurtt et al. 2020), and climatic extrapolations based on the CCS8.5 scenario  
268 (Karger et al. 2021). For all fungi taken together, predicted vulnerability to heat (best predictor:  
269 maximum monthly temperature;  $R^2_{\text{adj}}=0.583$ ) and drought (precipitation seasonality;  $R^2_{\text{adj}}=0.456$ ) were  
270 the greatest in the tropical and subtropical latitudes. Vulnerability to land use change (isothermality;  
271  $R^2_{\text{adj}}=0.145$ ) peaked in the tropics. The overall additive global change vulnerability was thus the highest  
272 in densely populated tropical and subtropical regions. Fungal functional groups had similar  
273 vulnerability patterns, which were mostly related to temperature. Among fungal groups, average  
274 vulnerability scores were highest for AM and EcM symbionts and unicellular fungi, but these scores  
275 differed only slightly across the global change drivers (**Fig. 9**). The actual vulnerability is probably  
276 underestimated for biotrophic pathogens and EcM fungi, because these groups associate with a limited  
277 number of plant species and are sometimes host-specific (Kennedy et al. 2015). Therefore, the loss of  
278 one of the few key symbiotic partners may greatly reduce the biotic niche of specialist fungi.

279 Patterns of vulnerability in fungi are somewhat similar to those of terrestrial plants and animals,  
280 where vulnerability peaks in drylands prone to desertification (Warren et al. 2013), arctic/alpine areas  
281 (cold-adapted species), and regions with dense human populations (Watson et al. 2013). The  
282 relatively low vulnerability to heat in tundra-inhabiting fungi can be explained by their relatively high



283 temperature optima (Maynard et al. 2019; but see Misiak et al. 2021), acclimation (Romero-Olivares  
284 et al. 2017), and poleward migration potential, despite relatively greater predicted warming in Arctic  
285 ecosystems. Above certain tolerance thresholds, soil organisms may be physiologically constrained by  
286 increasing soil temperature and evaporation, lower soil water potentials, and loss of oxygen due to  
287 greater respiration and faster decomposition, which result in hampered soil functioning and ecosystem  
288 multifunctionality (Delgado-Baquerizo et al. 2017). Open areas are predicted to increase due to  
289 climate change and human activities. This will further expose soil to solar radiation and result in the  
290 loss of fungal plant hosts. While here we calculated average vulnerabilities by adding up the effects of  
291 individual drivers, global change impacts tend to be synergistic (Rillig et al. 2019), so actual  
292 vulnerabilities may be much higher.

293

#### 294 *Implications for conservation*

295 Most fungi and soil organisms do not enjoy the protection and conservation measures that are  
296 afforded to more “charismatic” animals and plants (Ducarme et al. 2013). Nonetheless, fungi and  
297 other soil biota are pivotal to soil health, nutrient cycling, water storage, food security, and many  
298 other ecosystem services. Their biodiversity should hence be brought to the center stage of global  
299 sustainability thinking and conservation planning. For example, these organisms should be factored in  
300 when selecting protected areas otherwise based on plant and animal conservation (Guerra et al.  
301 2021a). The fact that many EcM and plant pathogenic fungal species are associated with specific host  
302 plants indicates that on the local scale it is not only the narrowly-distributed species but also unique  
303 biotic associations that require focused conservation measures. From the fungal perspective, it is  
304 particularly important to protect plant species that act as hubs in modules of biotic interaction  
305 networks, because these hub species typically associate with multiple, distinct fungal partners (Pölme  
306 et al. 2018). In other cases, certain unique plant species or higher taxonomic groups should be  
307 prioritized. For example, in southern South America, the drought-sensitive tree family Nothofagaceae  
308 is the only group known to support EcM fungi that are endemic to this area (Godoy & Marin 2019)

309 Although the vulnerability to environmental change differed among fungal groups, their overall global  
310 patterns were similar. This suggests that broad habitat conservation measures may work for most  
311 fungal groups, including macroscopic non-EcM Agaricomycetes and EcM fungi as well as more  
312 cryptic pathogens and other groups. To accomplish this, fungi need to be incorporated into  
313 conservation frameworks (Gonçalves et al. 2021). Actions to fill existing information gaps at the local  
314 and global levels must also be taken, and global-scale surveys should take into account the soil  
315 biodiversity assessments, complementing the traditional collections-based assessment with  
316 metabarcoding of environmental DNA. This applies to national conservation evaluation programs and  
317 engagement in global policy-making initiatives, such as the System of Environmental Economic

318 Accounting of the United Nations, World Biodiversity Forum, and Post-2020 Global Biodiversity  
319 Framework. Furthermore, promoting the red-listing of endangered fungal species at the national and  
320 global levels is critical (FAO 2020; IUCN 2021). Fungi need active and specific inclusion in national  
321 and global conservation policies and strategies, not just passive and implicit protection.

322 Our study provides evidence that soil fungi may be highly vulnerable to global change, which needs  
323 to be considered when planning how to preserve these key organisms in a changing world. As with  
324 plants and animals, fungi appear to be environmentally sensitive due to the strong impacts that land  
325 cover change, low moisture, and high temperatures have on taxonomic and functional composition  
326 (Brinkmann et al. 2019; Makiola et al. 2019; this study). The endemism of fungi is highest in tropical  
327 forest biomes (Kier et al. 2009; this study), so conservation measures advocated for tropical plants and  
328 animals (Brooks et al. 2006; Barlow et al. 2018) are likely to conserve fungi. Tropical forests are  
329 under continued threat from deforestation and degradation driven by expanding agriculture, extractive  
330 industries, and infrastructural projects (Bebbington et al. 2018). Conservation of herbaceous wetlands,  
331 tropical rainforests and tropical woodlands is supported by our global fungal conservation priority  
332 map that accounts for endemism, vulnerability, and  $\gamma$ -diversity (**Fig. 10**). Additionally, given the  
333 importance of soil pH for soil microbial diversity and composition, it is essential to prioritize areas  
334 with high pedodiversity or mixed landscapes including bogs, various forest types, and grasslands. As  
335 a crucial measure, desertification and loss of soil organic matter needs to be controlled by reducing  
336 the conversion of primary forest to crops and pasture (Smith et al. 2020a). This is important not only  
337 to prevent land degradation processes from impairing bacterial and fungal diversity, but also to sustain  
338 the capacity of drylands to provide essential functions and services, such as soil fertility, carbon  
339 storage, and food production for more than one billion people (Sivakumar 2007; Delgado-Baquerizo  
340 et al. 2018).

341

## 342 **Conclusions**

343 In conclusion, soil fungi show strong endemism patterns, which differ by functional groups and are  
344 driven by both climatic and edaphic factors. Fungal groups also differ strongly in their relative  
345 vulnerability scores to global change, which peak in heavily populated tropical dryland areas.  
346 Unfortunately, these are the very areas most prone to further land degradation and desertification.  
347 Fungal endemism and vulnerability patterns only partly mirror those of vascular plants and animals,  
348 which may be ascribed to their more efficient dispersal mechanisms. Global conservation efforts  
349 should include fungal biodiversity surveys alongside assessments of soil health, below-aboveground  
350 feedbacks, and areas of highest conservation priority, to secure the protection of habitats. Even more,  
351 they should include the monitoring of regional fungal communities over time, to pick relevant

352 changes and to provide early warning signals of impending change. What we do not know, we cannot  
353 efficiently manage or protect.

354

## 355 **Methods**

### 356 *Datasets*

357 To study fungal endemism and vulnerability to global change, we combined data from the Global  
358 Soil Mycobiome consortium (GSMc) open dataset (Tedersoo et al. 2021b) with materials from five  
359 other global soil biological surveys (**Fig. 1**) – BIODESERT (Maestre et al. 2022), MUSGONET  
360 (including the natural sites in Delgado-Baquerizo et al. 2021), CLIMIFUN (Bastida et al. 2021),  
361 GlobalAM (Davison et al. 2021), GlobalWetlands (Bahram et al. 2022) as well as Sanger sequence  
362 data from soil-inhabiting fungi obtained from the UNITE database (Nilsson et al. 2019) covering  
363 GenBank. We obtained the DNA from all five surveys and performed new DNA metabarcoding  
364 analyses following the protocols outlined for the GSMc dataset (Tedersoo et al. 2021b).

365 All datasets comprised information on geographical coordinates and soil pH. Based on geographical  
366 coordinates, we assigned the following climatic and land cover metadata to the samples: i) CHELSA  
367 v2.1 bioclimatic variables for the period 1981-2010 (Karger et al., 2020), ii) CHELSA-TraCE21k  
368 v1.0. for the LGM (Karger et al. 2021), and iii) CHELSA v2.1 climate extrapolations for the year  
369 2070 following the RCP8.5 global warming scenario with SSP5 socioeconomic conditions and the  
370 GFDL-ESM4 global circulation model (Karger et al., 2020); iv) normalized difference vegetation  
371 index (NDVI; Filipponi et al. 2018); v) SoilGrids v.2 soil pH from 0-5 cm depth (Poggio et al., 2021);  
372 vi) land cover type using Copernicus classification v.3 (Buchhorn et al. 2020) for the year 2015; and  
373 vii) human footprint index based on the Land-Use Harmonization (LUH2; Hurtt et al., 2020) or the  
374 year 2015 and 2070 extrapolation. Based on original descriptions of vegetation (age, cover, relative  
375 abundance of species, fire history) or remote sensing data (Google Maps), samples were assigned to  
376 biomes (Olson et al. 2001) and land cover types. Based on Z-transformed differences in present and  
377 LGM bioclimatic variables, we calculated for each sample an averaged LGM climate change index.  
378 Further, for each sample we estimated the human footprint index as the cumulative sum of land-use  
379 state transitions, with the year 1960 used as a baseline.

380

### 381 *Bioinformatics*

382 To infer fungal species and taxonomy, we used a long-read sequencing approach involving the  
383 ribosomal RNA 18S gene V9 subregion, ITS1 spacer, 5.8S gene, and ITS2 spacer to enhance

384 taxonomic resolution and accuracy. We used degenerate, universal eukaryotic primers to cover as  
385 many divergent taxa within the fungi and micro-eukaryotes as possible (Tedersoo et al. 2021a). The  
386 amplicon samples were prepared in 82 PacBio SMRTbell sequencing libraries and sequenced on 48  
387 PacBio Sequel 8M SMRT cells. The obtained reads were quality-filtered, demultiplexed to samples,  
388 trimmed to include only the full-length ITS region, and clustered to operational taxonomic units  
389 (conditionally termed as species) at 98% sequence similarity, which roughly corresponds to species-  
390 level divergence. Taxonomy was assigned based on information from the 10 best BLASTn matches  
391 against the UNITE 9.1 beta dataset (<https://doi.org/10.15156/BIO/1444285>). The resulting species-by-  
392 sample matrices were manually checked library-wise for external and cross-contamination and rates  
393 of index switching artifacts. We excluded several samples for which we suspected contamination, and  
394 removed rare occurrences of dominant species using the following thresholds: abundances = 1 for  
395 species with total abundance of >99 and abundances = 2 for species with total abundance of >999.

396 Based on FungalTraits 1.3 (Pöhlme et al. 2020), species belonging to the kingdom Fungi were assigned  
397 to functional groups based on ecological or physiological characters: i) AM fungi (including all  
398 Glomeromycota but excluding all Endogonomycetes, because there is not enough information to  
399 distinguish AM species from free-living species); ii) EcM fungi (excluding dubious lineages); iii)  
400 non-EcM Agaricomycetes (mostly saprotrophic fungi with macroscopic fruiting bodies; iv) molds  
401 (including Mortierellales, Mucorales, Umbelopsidales, and Aspergillaceae and Trichocomaceae of  
402 Eurotiales and Trichoderma of Hypocreales); v) putative pathogens (including plant, animal and  
403 fungal pathogens as primary or secondary lifestyles); vi) OHPs (excluding Mortierellales); vii) yeasts  
404 (excluding dimorphic yeasts); and viii) other unicellular (non-yeast) fungi (including chytrids, aphids,  
405 rozellids, and other early-diverging fungal lineages). Other groups such as lichen-forming fungi were  
406 not considered, owing to their relative infrequency in soil across samples and ecoregions. Among  
407 these groups, mostly non-EcM Agaricomycetes and EcM fungi comprise many red-listed species of  
408 conspicuous conch-shaped, resupinate, or stipitate fruiting bodies and are hence considered to be of  
409 relatively higher conservation interest (Cao et al. 2021; IUCN 2021).

410

#### 411 *Fungal diversity*

412 To assess patterns in global fungal  $\alpha$ -diversity, we first calculated the residuals of logarithmically-  
413 transformed fungal richness and richness of major functional groups by performing linear regression  
414 against the logarithm of sequencing depth. We also compared other approaches such as residuals from  
415 untransformed richness against square-root-transformed and log-transformed sequencing depth  
416 (Tedersoo et al. 2014), exclusion of singletons, Shannon index of diversity, traditional rarefaction  
417 (depth, 500 reads), and SRS-rarefaction (Beule & Karlovsky 2020) to 500 (minimum) or 3894  
418 (median) reads (**Fig. 2**). Because the approach including singletons and log-log transformation for

419 selecting residuals resulted in best-supported models (**Fig. 2**), we chose this approach for further  
420 analyses. Because the sampling protocols differed in sampling area size, number of subsamples and  
421 DNA extraction protocols, we included only the GSMc dataset in analyses of fungal richness and  
422 composition. The GSMc dataset furthermore featured information about vegetation (age, proportion  
423 of dominant plant taxa and mycorrhiza types, fire history) soil properties (C, N, P, K, Ca, Mg  
424 concentration), and sampling date. Based on geographical coordinates and sampling dates, we  
425 calculated geographic and temporal eigenvectors using the *adespatial* package of R (Dray et al. 2018).  
426 Phylogenetic eigenvectors were calculated for woody plant species composition by mapping the taxa  
427 to a multigene vascular plant phylogram (Qian & Jin 2016).

428 Because of metadata availability and comparability, we exclusively used the GSMc dataset (3200  
429 composite samples by 722,682 species) to estimate the best predictors of fungal  $\alpha$ -diversity and  
430 composition and to update global fungal diversity distribution maps. We first calculated the residuals  
431 of logarithmically-transformed fungal richness and richness of major functional groups by performing  
432 linear regression against the logarithm of sequencing depth. Besides richness residuals and Shannon  
433 index of diversity, we calculated fungal phylogenetic diversity (PD), mean phylogenetic distance  
434 (MPD), and mean neighbor taxonomic distance (MNTD) indices based on a classification tree  
435 (Tedersoo et al. 2018) using the *PhyloMeasures* package of R (Tsirogianis & Sandel 2016). For  
436 predictors, we included biome and continent (dummy variables), bioclimatic, edaphic, and vegetation-  
437 related variables, as well as eigenvectors of woody plant phylogenetic composition, spatial and  
438 temporal distance. Using random forest, we retrieved 20 candidate variables for GLM modeling. In  
439 GLM modeling, we included quadratic terms to account for non-linearity. To avoid an excessive  
440 number of predictors, only significant variables ( $P < 0.001$ ;  $R^2 > 0.020$ ) were kept in the final models.  
441 We also performed additional correlation analyses to illustrate the latitudinal gradient of  $\alpha$ -diversity  
442 and  $\gamma$ -diversity. For generating fungal diversity maps, we performed additional analyses using  
443 bioclimatic variables, database  $pH_{H2O(0-5\text{ cm})}$ , human footprint index and Copernicus land use categories  
444 and their interactions with continuous predictors as described for vulnerability maps. The GSMc  
445 dataset-based  $\alpha$ -diversity analyses were validated by performing similar analyses using other datasets.  
446 Richness residuals were calculated separately for these data and all datasets were subjected to model  
447 selection for soil pH and bioclimatic variables, because information about other predictors was  
448 inconsistent. We also modeled various versions of soil pH including the original measurements of  
449  $pH_{KCl}$  as well as  $pH_{H2O}$  extrapolations for soil depths of 0-5 cm, 0-30 cm, and 15-30 cm. Since the  
450  $pH_{KCl}$  fit significantly better into global models (**Fig. 2B**) and  $pH_{H2O}$  extrapolations were missing for  
451 smaller islands and Antarctic habitats, we used  $pH_{KCl}$  in subsequent analyses.

452 For  $\gamma$ -diversity, logarithmically-transformed cumulative ecoregion (see below) species richness was  
453 subjected to model selection against the logarithmically-transformed number of samples and  
454 sequencing depth to calculate residual richness. Residual richness and endemism indices of all fungi

455 and functional groups were subjected to random forest machine learning analysis to pre-select ten  
456 most important variables for GLM.

457

#### 458 *Endemicity*

459 To infer endemicity patterns in fungi, samples (including data obtained from UNITE, covering  
460 International Nucleotide Sequence Databases entries) were assigned to ecological regions (Olson et al.  
461 2001) based on their geographical coordinates, allowing a 10-km of buffer zone between a terrestrial  
462 ecological region and water (due to low resolution of the map layers in shore areas). Based on  
463 climatic and floristic similarities, the ecological regions were further aggregated into larger areas or  
464 split into smaller, geographically distinct units, which we refer to as ecoregions (**Fig. 7**). Each of 174  
465 ecoregions comprised 1 to 45 soil samples, with surplus samples excluded randomly. Five indices of  
466 endemism – viz., the number of endemic species (weight = 16.7%), proportion of endemic species  
467 (weight = 16.7%), mean maximum geographical range of taxa (weight = 33.3%), Jaccard index  
468 (weight = 16.7%), and beta-sim index (weight = 16.7%) – were selected for calculating the averaged  
469 endemicity index based on the community matrix (Crisp et al. 2001; Villéger & Brosse 2012; **Box 1**).  
470 The former two and latter two indices reflect similar aspects of endemicity and were therefore  
471 downweighted. To account for differences in sampling intensity, we calculated residuals for the  
472 numbers of all species and endemic species by regressing these against the logarithmically-  
473 transformed number of samples and sequencing depth.

474 Of the indices used, only the number and proportion of endemic taxa were significantly positively  
475 correlated with species richness (all fungi:  $r=0.707$  and  $r=0.212$ , respectively), whereas others had no  
476 significant correlation. Furthermore, species richness was not included among the best predictors of  
477 averaged endemicity, indicating that these metrics are independent. Endemicity indices were  
478 calculated using the betapart package v.1.5.4 (Baselga & Orme 2012) of R v.4.1.10 (R Core Team  
479 2022). Endemicity indices of all fungi and functional groups were subjected to random forest machine  
480 learning analysis to pre-select the ten most important variables for GLM. We used the variance (as  
481 coefficient of variation) and averaged values of bioclimatic variables, area, latitude, longitude,  
482 altitude, and soil pH as well as continents (dummy variables) to explain endemicity. GLMs were fitted  
483 using second-order polynomial terms for continuous variables. Only significant variables ( $P<0.050$ ;  
484  $r^2>0.020$ ) were kept in the final models. Based on the predictions revealed by GLMs, endemicity  
485 maps were constructed using the sf v.1.0-5 (Pebesma 2018) package of R.

486

487

488 *Global change vulnerability*

489 The vulnerability of soil fungal groups was estimated relative to three global change drivers – heat  
490 (maximum monthly temperature), drought (negative of inverse hyperbolic sine-transformed  
491 precipitation in the driest quarter), and land cover change – for the year 2070 (relative to 2015  
492 baseline) using the community-mean percentile vulnerability index ( $V_2$ ; Smith et al. 2020b). This  
493 index is based on averaging percentiles of all species at a given global change driver value.

$$V_{2i} = \left( \frac{\sum_{j=1}^n a_{ij} F_j(x_i)}{\sum_{j=1}^n a_{ij}} \right) \times 100$$

494

495 where  $a_{ij}$  is the presence (0 or 1) of species  $j$  in site  $i$ ,  $F_j(x_i)$  is the percentile of species  $j$  given site  
496 parameter value  $x_i$ ,  $x_i$  is the parameter value of site  $i$ , and  $n$  is the total number of species observed.

497 Precipitation in the driest quarter was selected as a proxy for drought, because bioclimatic variables  
498 cover larger areas (including islands) and offer greater resolution compared with other measures of  
499 soil water content and indicators of drought. The vulnerability scores were calculated for each soil  
500 sample using `vuln v.0.0.05` (Smith et al. 2020b) package of R. We also constructed the average  
501 vulnerability score by equally weighting all components. The vulnerability scores were unrelated to  
502 sequencing depth and sample size. We performed a similar random forest and GLM modeling  
503 exercise for determining the main predictors of vulnerability as described above, but allowed  
504 interaction terms between categorical and continuous predictors and used a more relaxed threshold for  
505 keeping variables in the model ( $P < 0.001$ ;  $R^2 > 0.01$ ) due to greater sample size. To construct  
506 vulnerability maps we used a regression-kriging approach (Hengl & MacMillan 2019). To predict  
507 vulnerability scores for each global driver and estimate their prediction uncertainty, thin plate splines  
508 (basis dimensionality = 3) were fitted using a generalized additive model (GAM) with `mgcv v.1.8-38`  
509 (Wood 2011) package. To incorporate the spatial autocorrelation signal, we calculated residuals at the  
510 sampling sites and used inverse distance weighting (IDW) to interpolate residuals beyond the  
511 sampling sites. To obtain final vulnerability predictions, interpolated residuals were added to the  
512 results based on the predicted regression part. By using the relative vulnerability values, we also  
513 prepared the map of fungal vulnerability ascribed to each of the three components. Vulnerability maps  
514 were visualized using the `raster v.3.5-9` (Hijmans 2021) package of R.

515 The maps for conservation priorities were calculated for all fungi using sampling points used in  
516 vulnerability analyses, except points corresponding to cropland and urban and village land cover. For  
517 each sampling point, the respective average endemism,  $\gamma$ -diversity, and vulnerability scores were z-  
518 transformed, followed by adding a constant (5, to exclude negative values), multiplied (to downweigh

519 areas with any low values), and used in a regression-kriging approach (**Table 3**) as described for  
520 vulnerability.

521 Based on methods comparison, we conclude that inclusion of singletons may increase richness model  
522 performance at relatively low sequencing depth in high-quality, third-generation sequencing datasets.  
523 Furthermore, residuals of log-transformed richness against log-transformed sequencing depth yields  
524 significantly better model estimates compared with most other standardization methods. Richness data  
525 from studies with different sampling designs must not be pooled in a common analysis unless the  
526 factor “study” or sampling attributes (e.g. number of subsamples, volume of samples) are accounted  
527 for. Use of original soil pH data strongly outperforms extrapolated data; therefore, soil pH  
528 extrapolations should not be used for testing relative effects of edaphic and climatic and other  
529 properties. This probably applies to other edaphic and vegetation-related values that greatly vary on a  
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531

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540 PlutoF data repository (GSMc data: <https://doi.org/10.15156/BIO/2263453>). Representative sequences  
541 of identical reads per sample are available from the UNITE database. The scripts used for the  
542 bioinformatic analysis are available at GitHub: [https://github.com/Mycology-Microbiology-  
543 Center/GSMc](https://github.com/Mycology-Microbiology-Center/GSMc)

544

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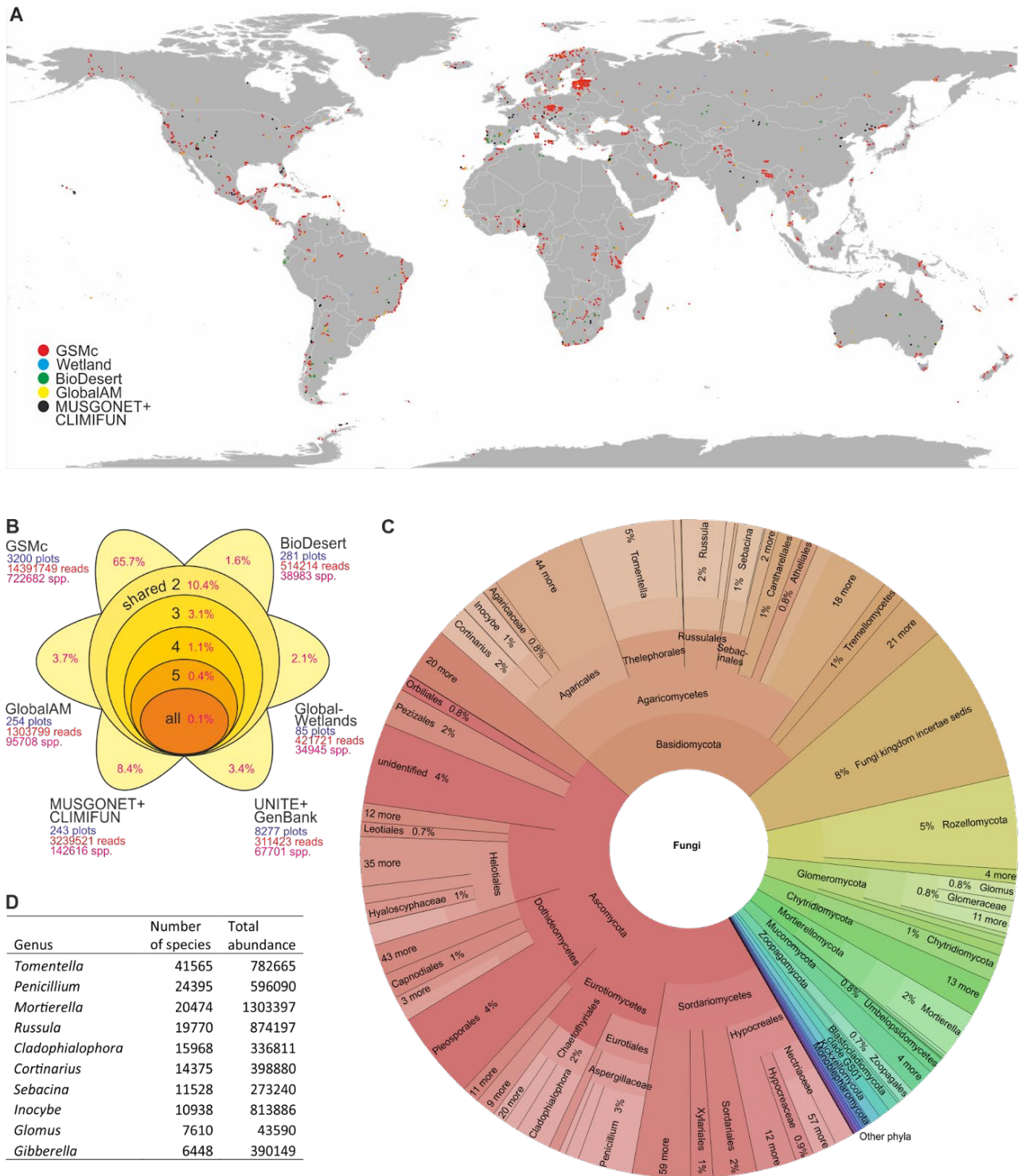
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784 **Figures**

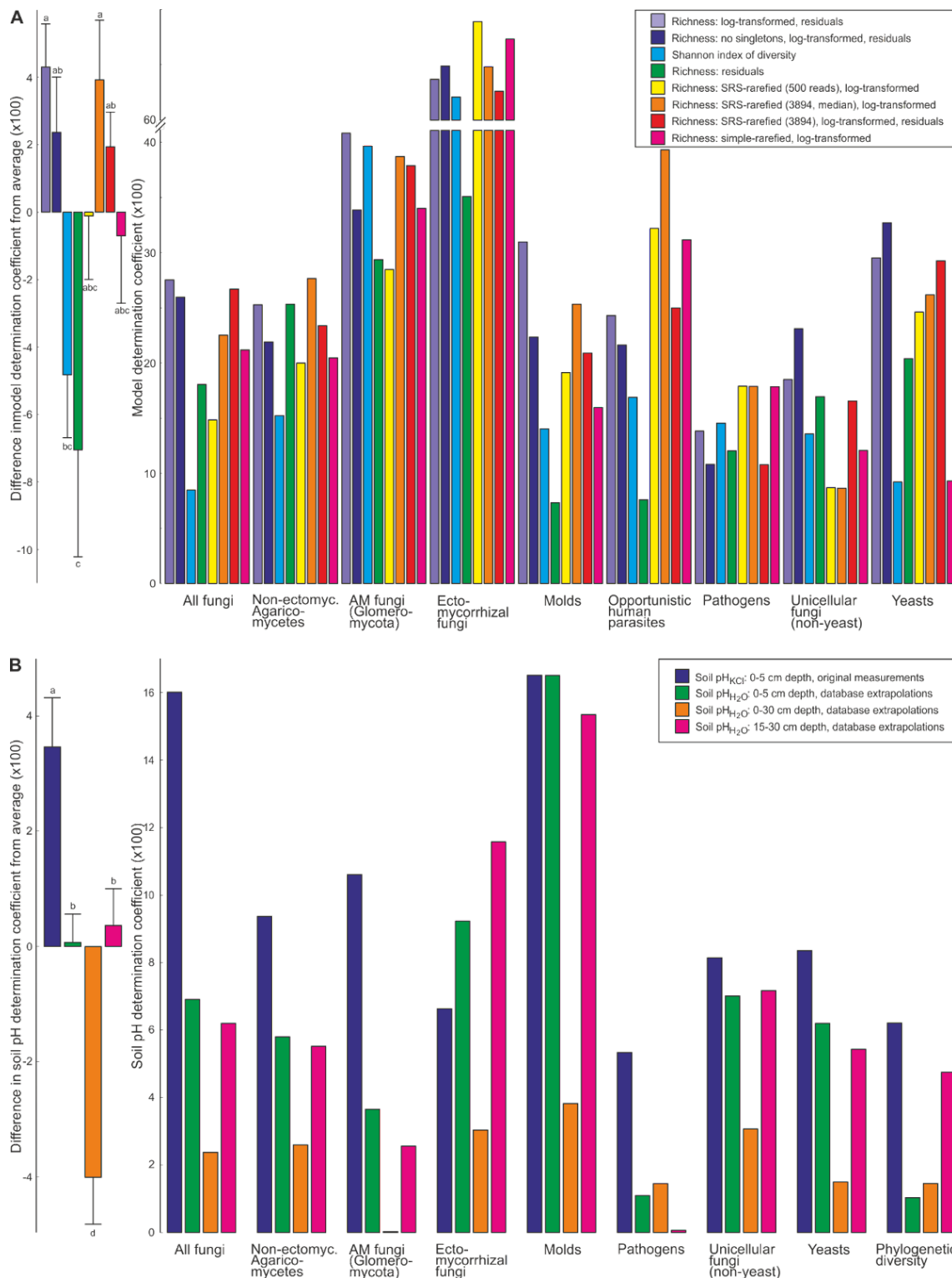
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788 **Fig. 1. Distribution of samples and fungal species across datasets.** (A) Global sampling map, with different  
 789 symbols representing different datasets; (B) species distribution of fungi among datasets, with the proportion of  
 790 unique and shared species indicated in the diagram; (C) Krona chart indicating taxonomic distribution of fungal  
 791 species (interactive chart can be browsed at <https://plutof.ut.ee/#/doi/10.15156/BIO/2483900>); (D) species  
 792 richness and total read abundance of the top 10 most diverse fungal genera.

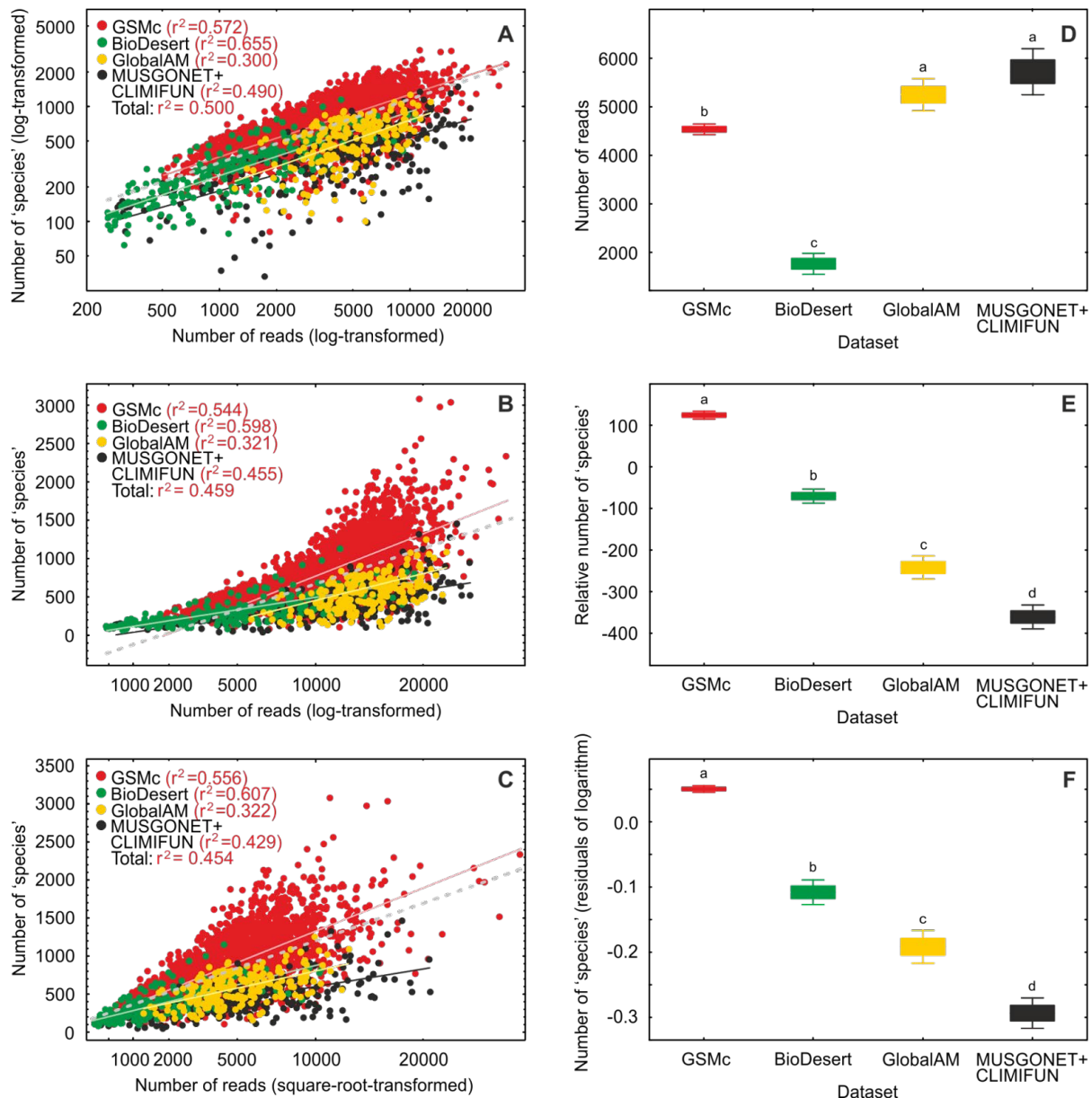


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795 **Fig. 2. Comparison of (A) richness proxies (use of log-transformation, residuals of sequencing depth, SRS**  
 796 **or simple rarefaction) and (B) measures of soil pH on analytical performance.** Relative goodness was  
 797 estimated based on the determination coefficients of the best models (A) or pH-only models (B). In left panels,  
 798 significant among-group differences are indicated with different letters based on Tukey Posthoc tests; bars,  
 799 means; whiskers, SE. Soil pH<sub>KCl</sub> were determined experimentally, whereas pH<sub>H<sub>2</sub>O</sub> were obtained from Poggio et  
 800 al. (2021).

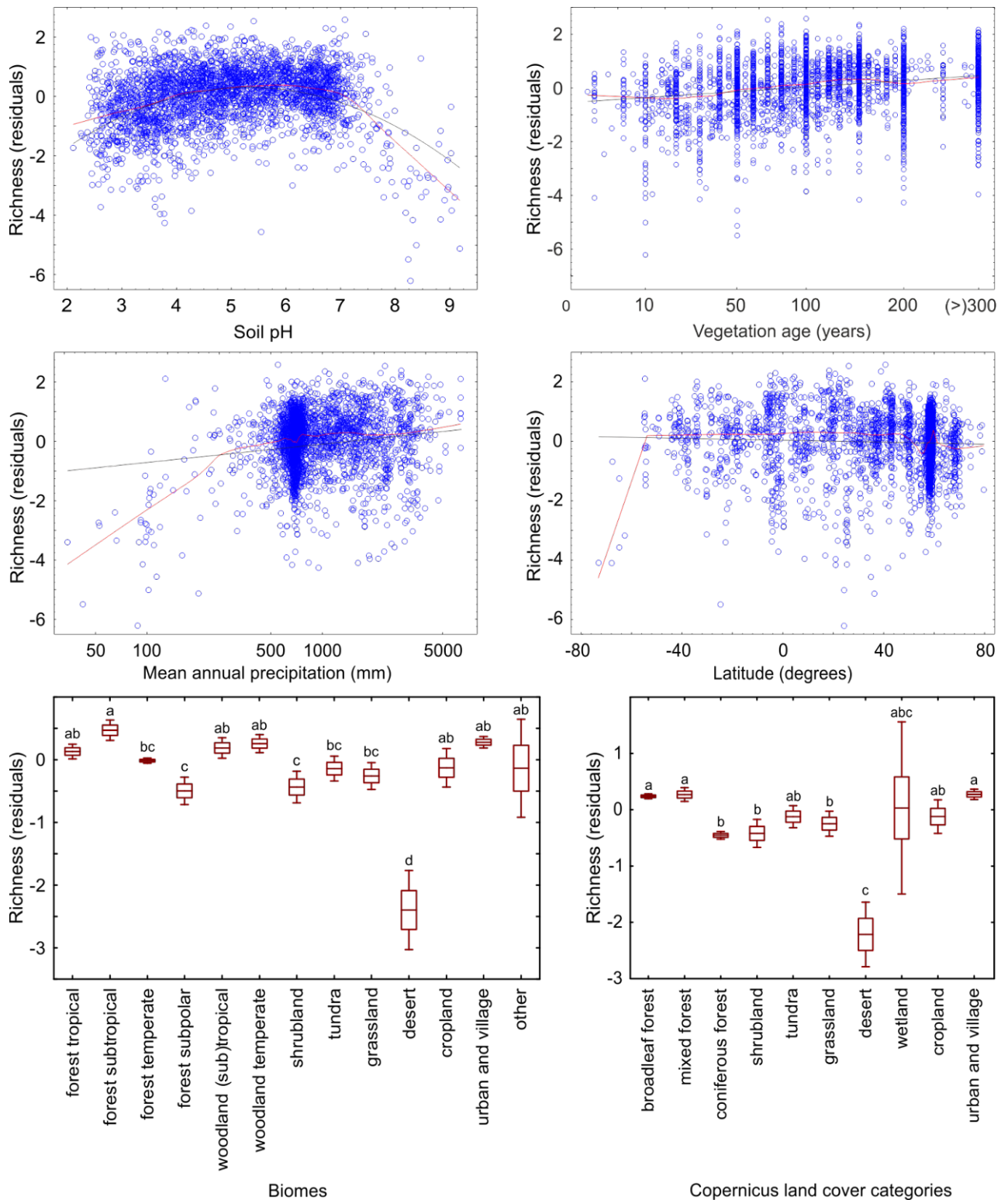
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804 **Fig. 3. Relative 'species' accumulation curves (A-C), sequencing depth (D) and 'species' richness (E-F)**  
 805 **across four datasets.** (A) The log-log relationship between the number of reads and 'species' richness that was  
 806 used for calculation of residuals and further analyses; (B-C) Relatively lower performance of log-linear  
 807 relationships of log-transformed and square-root-transformed sequencing depth; (D) Initial differences in  
 808 sequencing depth among datasets; (E-F) Fungal 'species' richness differences relative to the average in the raw  
 809 data (E) and residuals of the log-log regression analysis (F). In D-F, boxes indicate standard errors around the  
 810 mean and whiskers indicate 95% confidence intervals; letters above whiskers indicate statistically significant  
 811 differences among datasets (using log-transformed data for D-E). These analyses indicate that the log-log  
 812 transformation for calculating residuals is relatively more robust compared with other methods and that richness  
 813 estimates from studies with different methods cannot be directly compared.



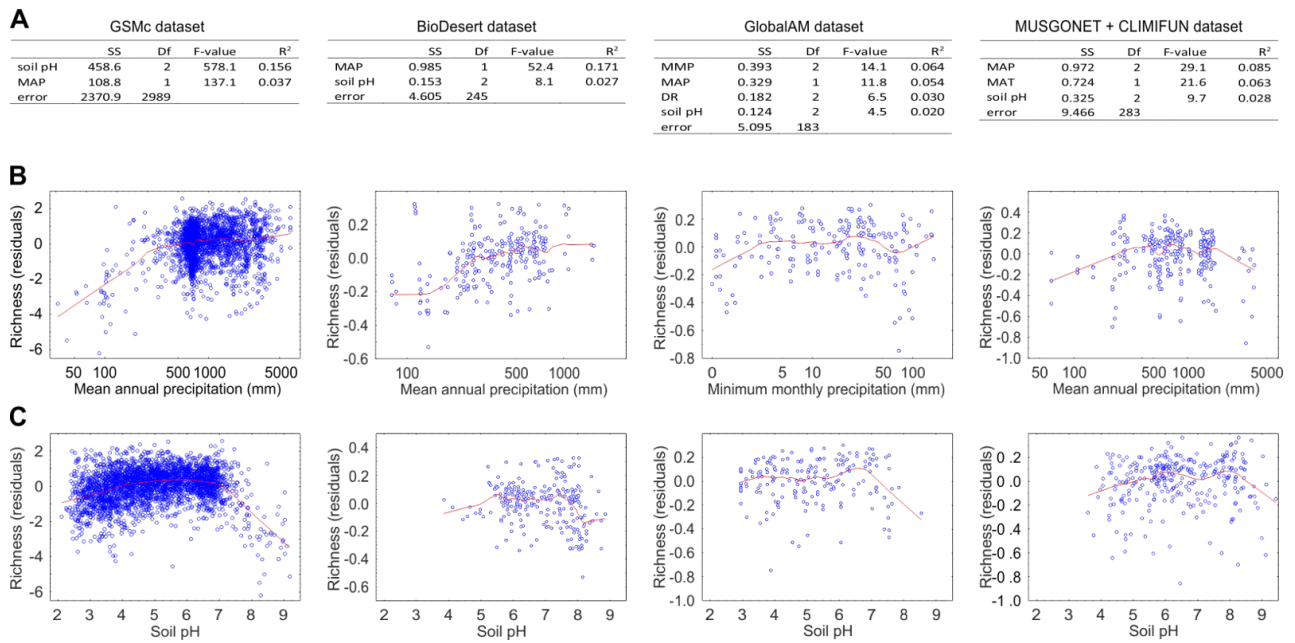
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816 **Fig. 4. Response of  $\alpha$ -diversity of all fungi to soil pH, vegetation age, mean annual precipitation, latitude,**  
 817 **biomes, and land cover categories.** For continuous predictors, black lines indicate linear and polynomial fits  
 818 and red lines indicate lowess fits. For categorical predictors, boxes represent standard error around the mean  
 819 (central line), whiskers depict 95% CI and letters above boxes indicate statistically significant different groups  
 820 ( $P < 0.001$ ).

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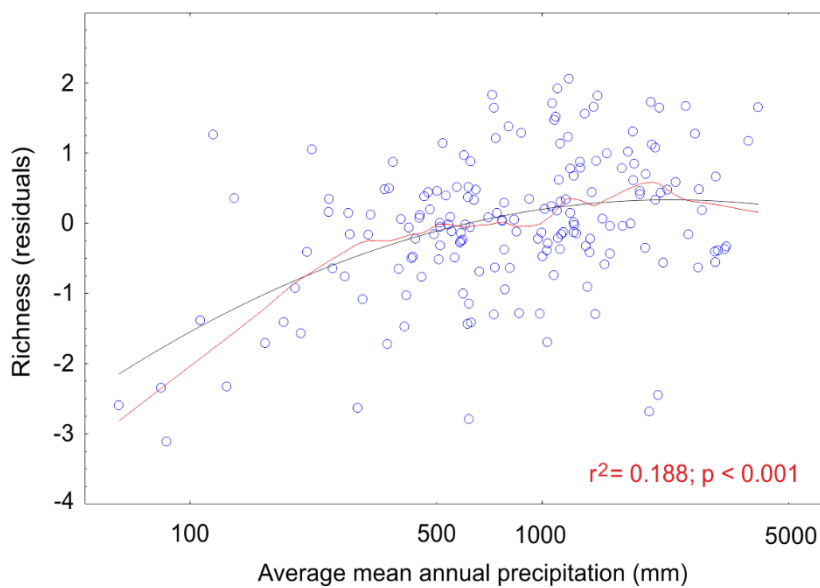


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824 **Fig. 5. Comparison of  $\alpha$ -diversity patterns in all fungi across four most inclusive datasets: (A) best models**  
 825 **(only bioclimatic variables and soil pH were included in model selection), (B) lowess regression curves for**  
 826 **the best-fitting climatic predictor, and (C) lowess regression curves for soil pH.**

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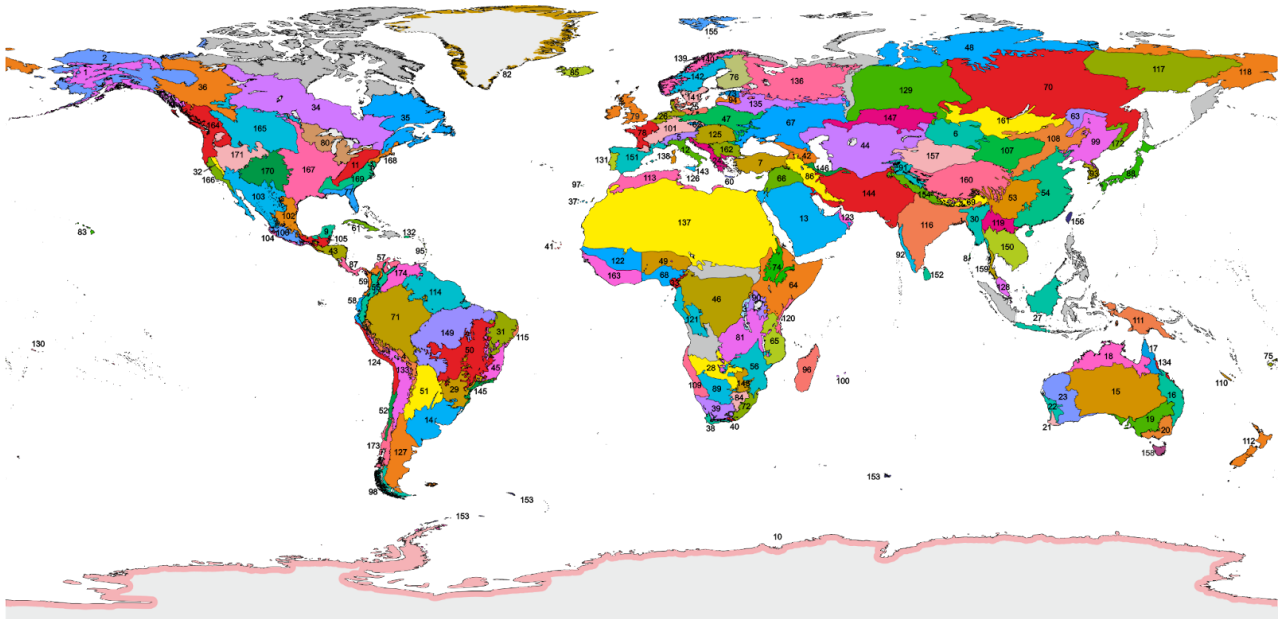
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830 **Fig. 6. The effect of average mean annual precipitation on  $\gamma$ -diversity of fungi at the ecoregion scale. Black**  
 831 **line, best quadratic fit; red line, lowess curve.**

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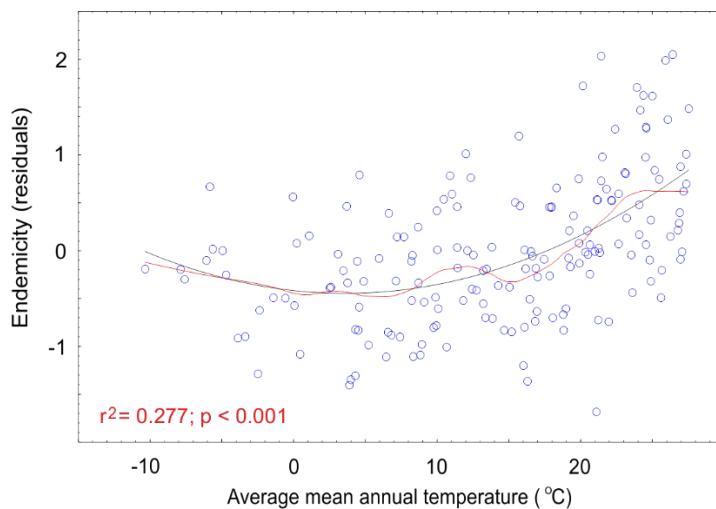
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836 **Fig. 7. Distribution of 174 ecoregions used in endemicity analyses.** Ecoregions excluded from the analyses  
837 due to the lack of data are indicated in gray. Their explanation is given in Table 1.

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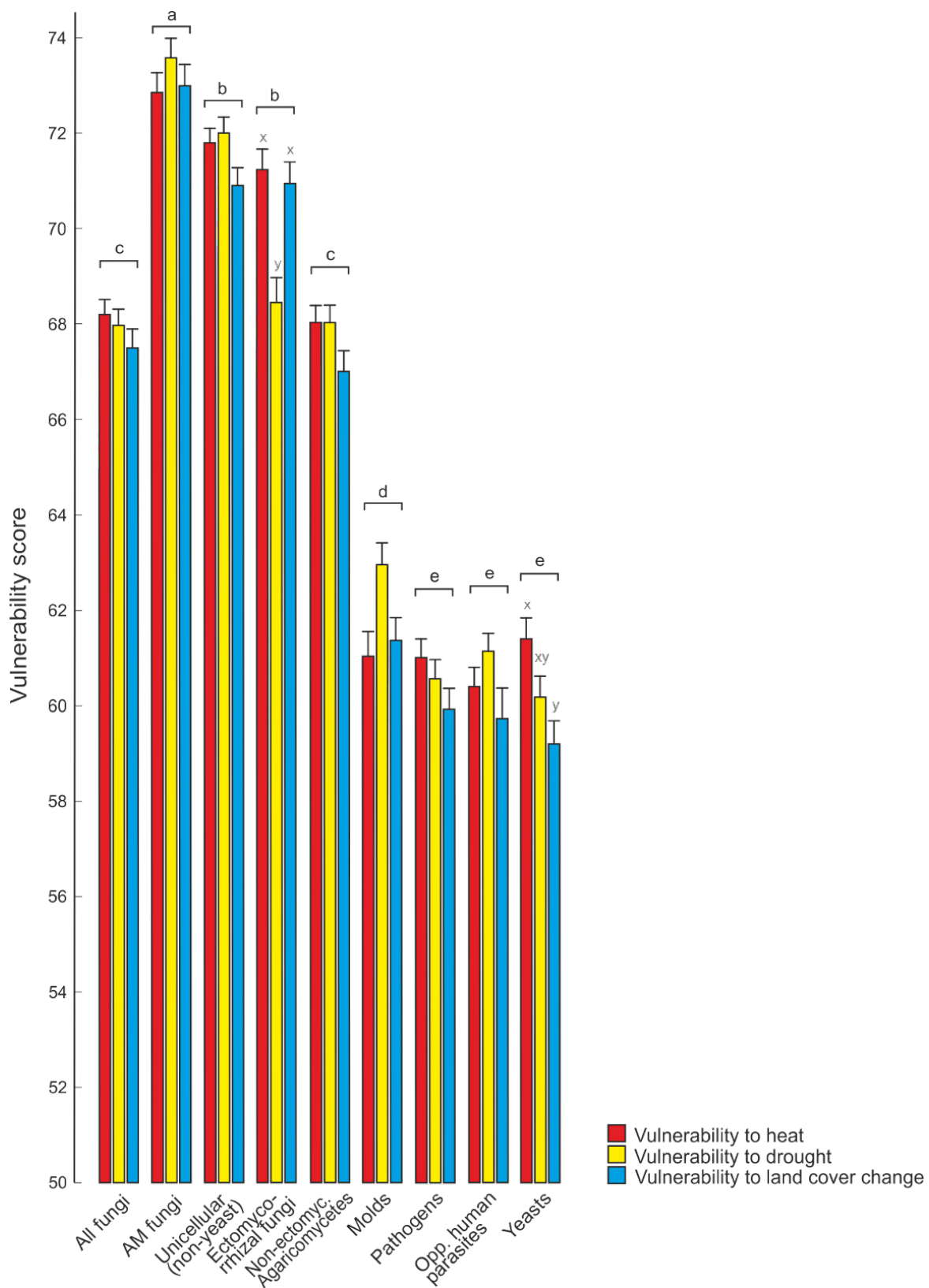
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842 **Fig. 8. The effect of average mean annual precipitation on endemicity of fungi at the ecoregion scale.**  
843 Black line, best quadratic fit; red line, lowess curve.



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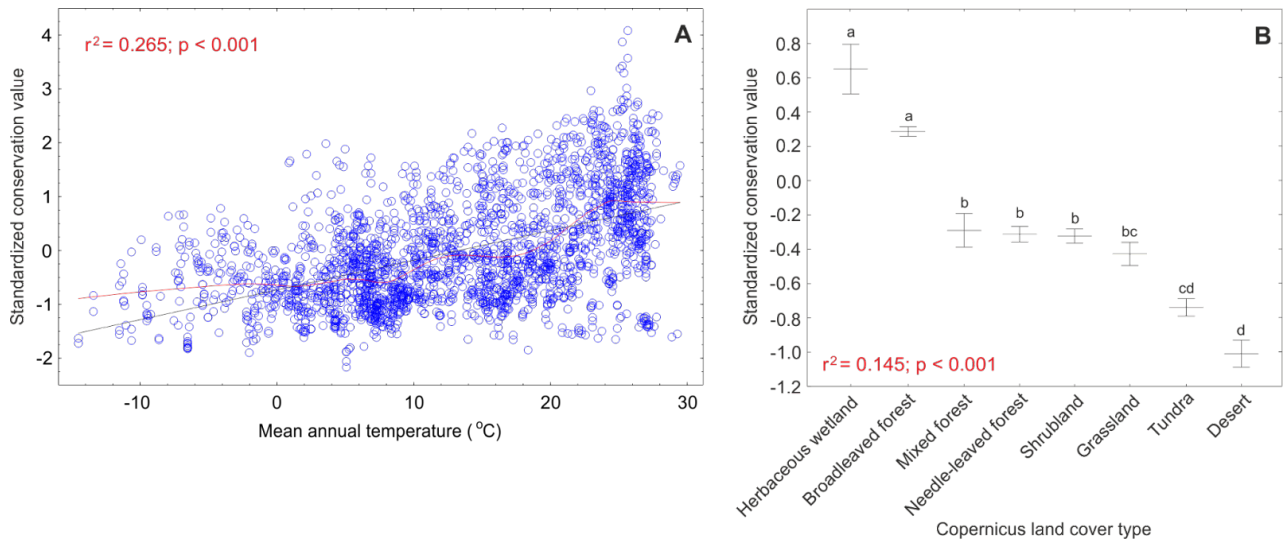
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846 **Fig. 9. Vulnerability of fungi and functional groups to global change drivers.** Different letters indicate  
847 statistically significant ( $P < 0.001$ ) differences among functional groups (a-e) and among global change drivers  
848 within functional groups (x-z).

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854 **Fig. 10. Relationships between conservation priority areas with mean annual temperature (A) and**  
855 **Copernicus land cover types (B).** In A, black and red lines indicate best-fitting linear and lowess functions,  
856 respectively. In B, central lines and whiskers indicate mean and standard errors, respectively; letters above  
857 whiskers indicate statistically significant differences among land cover types.

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## Calculation of endemism indices

To estimate endemism at the ecoregion level, we calculated five endemism indices ( $E_1$ - $E_5$ ). Weighted endemism indices were calculated based on these five indices using z-transformation of residuals in regression analyses against sequencing and sampling depth.

### Five endemism indices

$E_1$ Number of endemic species	$E_1$	
$E_2$ Proportion of endemic species	$E_2 = \frac{E_1}{S}$	$S$ , total species richness in ecoregion
$E_3$ Mean maximum range of species (average weighted endemism; Crisp et al. 2001)	$E_3 = \frac{\sum_i d_i}{S}$	$d_i$ , range of species $i$
$E_4$ Jaccard index of similarity (Jaccard 1912)	$E_4(C_1, C_2) = \frac{C_1 \cap C_2}{C_1 \cup C_2}$	$C$ , ecoregion communities
$E_5$ $\beta_{SIM}$ index of similarity (multiple-site Simpson-based community similarity index, Baselga et al. 2010)	$E_5 = \frac{\sum_{i < j} \min(b_{ij}, b_{ji})}{\sum_i (S_i - S_i) + \sum_{i < j} \min(b_{ij}, b_{ji})}$	$S_i$ , number of species in site $i$ $S_j$ , total number of species $b_{ij}$ and $b_{ji}$ , number of species exclusive to sites $i$ and $j$ , respectively

### Accounting for sampling and sequencing depth for endemism indices

$E_{nR}$ residuals of $n$ endemism index	$E_{nR} = E_n - \hat{E}_n$ $\hat{E}_n = \beta_1 \text{Log}(L_1 + L_2) + \beta_2 \text{Log}(R_1 + R_2) + \beta_3 \text{Log}(L_1) + \beta_4 \text{Log}(L_2) + \beta_5 \text{Log}(R_1) + \beta_6 \text{Log}(R_2) + \beta_7(L_1 + L_2) + \beta_8(R_1 + R_2) + \beta_9 L_1 + \beta_{10} L_2 + \beta_{11} R_1 + \beta_{12} R_2 + \varepsilon$	$L_1$ , number of soil sampling localities $L_2$ , number of UNITE localities $R_1$ , number of reads of soil samples $R_2$ , number of UNITE Sanger reads
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Residuals are calculated based on the best-fitting model.

### Standardizing residuals of endemism indices

$E_{nRz}$ Z-transformed $E_{nR}$	$E_{nRz} = \frac{E_{nR} - \mu}{\sigma}$	$\mu$ , mean $E_{nR}$ $\sigma$ , standard deviation of $E_{nR}$ .
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### Weighting standardized endemism indices

$E_{ave}$ weighted $E_{nRz}$	$E_{ave} = \frac{E_{1Rz} + E_{2Rz} + 2E_{3Rz} + E_{4Rz} + E_{5Rz}}{6}$
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$E_3$  receives double weight, because  $E_1$  and  $E_2$  as well as  $E_3$  and  $E_4$  are calculated on a similar basis.

### Correlation matrix of endemism indices

	$E_{ave}$	$E_{1Rz}$	$E_{2Rz}$	$E_{3Rz}$	$E_{4Rz}$	$E_{5Rz}$
$E_{ave}$	1.000	0.767	0.805	0.554	0.754	0.816
$E_{1Rz}$	0.767	1.000	0.806	0.188	0.452	0.540
$E_{2Rz}$	0.805	0.806	1.000	0.072	0.788	0.745
$E_{3Rz}$	0.554	0.188	0.072	1.000	0.169	0.270
$E_{4Rz}$	0.754	0.452	0.788	0.169	1.000	0.782
$E_{5Rz}$	0.816	0.540	0.745	0.270	0.782	1.000

### Correlation matrix of average endemism of fungal groups

	1	2	3	4	5	6	7	8	9
all fungi (1)	1.000	0.797	0.591	0.517	0.568	0.628	0.675	0.619	0.523
non-EcM Agaricomycetes (2)	0.797	1.000	0.790	0.398	0.461	0.470	0.460	0.595	0.532
AM fungi (3)	0.591	0.790	1.000	0.280	0.375	0.232	0.233	0.578	0.461
EcM fungi (4)	0.517	0.398	0.280	1.000	0.234	0.268	0.250	0.270	0.176
molds (5)	0.568	0.461	0.375	0.234	1.000	0.600	0.413	0.364	0.513
OHP (6)	0.628	0.470	0.232	0.268	0.600	1.000	0.809	0.159	0.354
pathogens (7)	0.675	0.460	0.233	0.250	0.413	0.809	1.000	0.264	0.331
unicellular fungi (8)	0.619	0.595	0.578	0.270	0.364	0.159	0.264	1.000	0.463
yeasts (9)	0.523	0.532	0.461	0.176	0.513	0.354	0.331	0.463	1.000

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861 **Box 1. Calculation of endemism indices and correlation among standardized indices for all fungi and**  
862 **among functional groups.**

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864 **Table 1. The ecoregions used in endemicity analyses.**

Ecoregion number	Putative name
1	Alaskan forests
2	Alaskan tundra
3	Albertine Rift montane forests
4	Yunga
5	Alps conifer and mixed forests
6	Altai steppes and semideserts
7	Anatolian forests
8	Andaman Islands rain forests
9	Yucatan forests
10	Antarctic desert
11	Appalachian mixed forests
12	Appennine forests
13	Arabic drylands
14	Argentina savanna
15	Australia Central drylands
16	Australia E subtropical woodlands
17	Australia NE savannas
18	Australia NW savannas
19	Australia SE shrublands and savannas
20	Australia SE temperate forests
21	Australia SW woodlands
22	Australia W savannas
23	Australia W shrublands
24	Balkan forests
25	Baltic mixed forests
26	Benelux Atlantic mixed forests
27	Borneo-Java rain forests
28	Botswana woodlands
29	Brazil S forests
30	Burman forests
31	Caatinga
32	California Sierra Nevada forests
33	Cameroon W forests
34	Canada N boreal forests
35	Canada NE boreal forests
36	Canada NW boreal forests
37	Canary Islands dry woodlands and forests
38	Cape fynbos
39	Cape karoo
40	Cape thickets
41	Cape Verde Islands dry forests
42	Caucasia
43	Central American forests
44	Central Asian drylands

- 45 Central Atlantic rain forests
- 46 Central Congolian forests
- 47 Central European mixed forests
- 48 Central Siberian tundra
- 49 Central Sudanian savanna
- 50 Cerrado
- 51 Chaco
- 52 Chilean matorral
- 53 China Central forests
- 54 China E forests
- 55 Colombia montane forests
- 56 Zimbabwe-Mozambique woodlands
- 57 Colombia N forests
- 58 Colombia SW dry forests
- 59 Colombia W forests
- 60 Crete Mediterranean forests
- 61 Cuba forests
- 62 Czech mixed forests
- 63 Da Hinggan-Dzhagdy Mountains conifer forests
- 64 East African shrublands
- 65 East African woodlands
- 66 East Asian drylands
- 67 East European forest steppe
- 68 East Guinean forests
- 69 East Himalayan forests
- 70 East Siberia forest and mountain tundra
- 71 West Amazon forests
- 72 Eastern South African woodlands
- 73 Estonian Sarmatic mixed forests
- 74 Ethiopian montane woodlands
- 75 Fiji forests
- 76 Finland taiga
- 77 Florida forests
- 78 France Atlantic mixed forests
- 79 Great Britain Forests
- 80 Great Lakes forests
- 81 Zambia woodlands
- 82 Greenland tundra
- 83 Hawaii forests
- 84 Highveld grasslands
- 85 Iceland boreal birch forests and alpine tundra
- 86 Zagros Mountains forest steppe
- 87 Isthmian forests
- 88 Japan forests
- 89 Kalahari xeric savanna
- 90 Victoria Basin forest-savanna mosaic
- 91 Karakoram drylands

- 92 Western Ghats forests
- 93 Korean forests
- 94 Latvian Sarmatic mixed forests
- 95 Lesser Antilles forests
- 96 Madagascar woodlands
- 97 Madeira evergreen forests
- 98 Magellanic subpolar forests
- 99 Manchurian forests
- 100 Mascarene forests
- 101 Western European broadleaf forests
- 102 Mexico NE shrublands and forests
- 103 Mexico NW deserts and montane forests
- 104 Mexico S dry forests
- 105 Mexico SW lowland forests
- 106 Mexico SW montane forests
- 107 Mongolian deserts
- 108 Mongolian steppe
- 109 Namib drylands
- 110 New Caledonia forests
- 111 New Guinea forests
- 112 New Zealand forests
- 113 North African woodlands
- 114 North Amazon forests
- 115 North Atlantic rain forests
- 116 North India moist forests
- 117 Northeast Siberian taiga
- 118 Northeast Siberian tundra
- 119 Northern Indochina subtropical forests
- 120 Northern Zanzibar-Inhambane coastal forest mosaic
- 121 Western Congolian forests
- 122 West Sudanian savanna
- 123 Oman drylands
- 124 Pacific deserts
- 125 Pannonian forests
- 126 Pantelleria-Lampedusa forests
- 127 Patagonian steppe
- 128 Peninsular Malaysian rain forests
- 129 West Siberian forests
- 130 Polynesian forests
- 131 Portugal woodlands
- 132 Puerto Rico forests
- 133 Puna
- 134 Queensland tropical rain forests
- 135 Russian Sarmatic mixed forests
- 136 Russian taiga
- 137 Sahelian drylands
- 138 Sardinian-Corsican forests



- 139 Scandinavian coastal conifer forests
  - 140 Scandinavian Montane Birch forest and grasslands
  - 141 Scandinavian Sarmatic mixed forests
  - 142 Scandinavian taiga
  - 143 Sicilian forests
  - 144 South Asian drylands
  - 145 South Atlantic rain forests
  - 146 South Caspian forests
  - 147 South Siberian steppe
  - 148 Southeast African bushveld
  - 149 Southeast Amazon forests
  - 150 Southern Indochina tropical forests
  - 151 Spain woodlands
  - 152 Sri Lanka forests
  - 153 Subantarctic islands tundra
  - 154 West Himalayan forests
  - 155 Svalbard Arctic desert
  - 156 Taiwan forests
  - 157 Tarim basin drylands
  - 158 Tasmanian forests
  - 159 Tenasserim-South Thailand semi-evergreen rain forests
  - 160 Tibetan steppe
  - 161 Trans-Baikal forests
  - 162 Transylvanian forests
  - 163 West Guinean forests
  - 164 USA California chaparral
  - 165 USA Central forests and grasslands
  - 166 USA NE hemiboreal forests
  - 167 USA SE forests
  - 168 USA Southern Rockies and steppe
  - 169 USA western steppe and montane forests
  - 170 USA-Canada Pacific forests and grasslands
  - 171 USA-Canada Rockies and central forests and grasslands
  - 172 Ussuri broadleaf and mixed forests
  - 173 Valdivian temperate forests
  - 174 Venezuela woodlands
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867 **Table 2. The best predictors of endemism indices in ecoregions for all fungi.**

	DF	Sum of squares	Mean squares	F-value	R <sup>2</sup> <sub>adj</sub>	P-value	Trend
MAT <sub>mean</sub>	2	27.2	13.6	97.4	0.277	<0.001	U-shaped
soil pH <sub>mean</sub>	1	10.6	10.6	38.2	0.108	<0.001	negative
subcontinent: Europe	1	6.5	6.5	23.5	0.065	<0.001	negative
human footprint index	1	2.1	2.1	7.5	0.018	0.007	negative
error	168	48.3					

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869 **Table 3. The best models of conservation priority co-kriging maps for fungi. All P-values are <0.001.**

	Df	Sum of squares	Mean squares	F-value	R <sup>2</sup> <sub>adj</sub>	Trend
air MAT <sup>1</sup>	2	2133	1066	1426	0.266	positive
MPWM	1	992	992	663	0.134	positive
climate isothermality	1	169	169	113	0.023	positive
MAP	2	167	84	112	0.022	positive
Land cover type <sup>2</sup>	7	128	18	12	0.015	
Land cover type x MAP	7	121	17	12	0.014	
error	2477	3705				

<sup>1</sup>Abbreviations: MAP, mean annual precipitation; MAT, mean annual temperature; MPWM, mean precipitation of wettest month;

<sup>2</sup>Copernicus land cover types: broadleaf forest, mixed forest, coniferous forest, shrubland, tundra, grassland, desert, wetland (note that cropland and urban and village biomes were excluded).

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