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Historical biome distribution and recent human disturbance shape the diversity of arbuscular mycorrhizal fungi

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- 2 arbuscular mycorrhizal fungi

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Supplementary Information (2 figures, 9 tables)

37	Summary
38	The availability of global microbial diversity data, collected using standardized metabarcoding
39	techniques, makes microorganisms promising models for investigating the role of regional and
40	local factors in driving biodiversity.
41	We modelled the global diversity of symbiotic arbuscular mycorrhizal (AM) fungi using
42	currently available data on AM fungal molecular diversity (SSU-rRNA gene sequences) in field
43	samples. To differentiate between regional and local effects, we estimated species pools (sets of
44	potentially suitable taxa) for each site, which are expected to reflect regional processes. We then
45	calculated community completeness, an index showing the fraction of the species pool present,
46	which is expected to reflect local processes.
47	We found significant spatial variation, globally in species pool size, as well as in local and dark
48	diversity (absent members of the species pool). Species pool size was larger close to areas
49	containing tropical grasslands during the last glacial maximum, which are possible centres of
50	diversification. Community completeness was larger in regions of high wilderness (remoteness
51	from human disturbance). Local diversity was correlated with wilderness and current
52	connectivity to mountain grasslands.
53	Applying the species pool concept to symbiotic fungi facilitated a better understanding of how
54	biodiversity can be jointly shaped by large-scale historical processes and recent human
55	disturbance.
56	Keywords
57	Biodiversity, Dark diversity, Ice Age, Mycorrhizae, Quaternary, Species pool, Tropical grassy
58	biome, Wilderness
59	
60	Introduction
61	Global diversity patterns have frequently been described for macroorganisms, including vascular
62	plants and vertebrates (Gaston, 2000, Orme et al., 2005, Kreft & Jetz, 2007). Yet, understanding
63	the relative roles of different processes in shaping diversity patterns is an ongoing challenge
64	(Pärtel et al., 2016). Local diversity patterns in any group of taxa are expected to emerge as a

65	consequence of simultaneous, and potentially confounding, effects of regional (evolutionary
66	changes, historical dispersal) and local processes (dispersal in contemporary landscapes, local
67	biotic and abiotic filters, natural and anthropogenic disturbances; Huston, 1994; Ricklefs, 2004,
68	2007; Zobel, 2016). Distinguishing between regional and local processes requires diversity data
69	that are comparable and replicated over large spatial scales. Molecular identification of microbial
70	taxa from environmental samples might provide data that are much closer to meeting this
71	requirement than traditional sampling of macroorganisms. However, macroecology of microbes
72	is a recent field (Hanson et al., 2012; Wardle & Lindahl, 2014) and descriptions of global
73	diversity patterns and their potential underlying drivers are largely lacking.
74	Identifying species pools – sets of potentially available species that are able to inhabit and
75	reproduce under particular habitat conditions in given sites (Cornell & Harrison, 2014) – is a
76	useful starting point for distinguishing regional and local processes acting on diversity. Species
77	pools develop via speciation under particular habitat conditions, as well as via historical
78	migrations between regions with similar conditions (Zobel 2016; Pärtel et al. 2016). Hence, one
79	may expect that species pools are shaped mainly by regional factors. Species pools can be
80	partitioned into locally present and locally absent fractions; the latter has been referred to as dark
81	diversity (Pärtel et al., 2011). From these two pieces of information, community completeness –
82	an index characterizing the share of the species pool present at a given site (Pärtel et al., 2013) –
83	can be calculated as the log-transformed ratio of local and dark diversity. Community
84	completeness indicates how easily potentially suitable species reach and establish in local
85	communities, but also how well local populations persist. Hence it can be expected that
86	community completeness is mainly driven by local factors.
87	There is only limited empirical support for the theoretical expectations stemming from the
88	species pool concept (see Lessard et al., 2012 and Zobel, 2016 for review). Empirical species
89	pool studies have hitherto addressed vertebrates, insects and plants, but large scale
90	generalizations have been limited due to the multitude of methods and scales used to assess
91	diversity and the hugely variable depth of diversity data from different parts of the globe.
92	Consequently, local diversity estimates used in large-scale comparisons have often been derived
93	from coarse grid-based distributions, or even from distribution range maps, and have therefore
94	lacked information about actual diversity in local communities. A more suitable approach to

95	disentangling the relative roles of regional and local factors in driving large-scale patterns of
96	biodiversity is to use local community data that are collected in a comparable manner throughout
97	an area of interest and take proper account of species pools.
98	The paucity of current data also poses challenges for dark diversity estimation (Pärtel et al.,
99	2016). For well-studied organisms, expert opinion has been used to estimate dark diversity,
100	either by linking species to habitat types or giving indicator scores along the main environmental
101	gradients (de Bello et al., 2016). Current developments in mathematical dark diversity methods
102	based on species co-occurrences or species distribution modelling provide a promising
103	alternative (Lewis et al., 2016; Ronk et al., 2016). These techniques assume that co-occurring
104	taxa share similar ecological preferences and possibly also joint biogeographic history. Such an
105	assumption is probably valid for stable ecosystems but should be applied with caution to
106	successional ecosystems where many species are not in equilibrium with environmental
107	conditions.
108	Perhaps surprisingly, suitable data for exploring global biodiversity patterns and processes may
109	already be available in the form of microbial community data. Microbial diversity estimates are
110	frequently derived using fairly standardized metabarcoding approaches and thus seem to more
111	easily satisfy criteria of comparability than existing macro-organism data sets (Taberlet et al.,
112	2012; Ficetola et al., 2015). Although microbes had until recently received little attention in
113	macroecology (Wardle & Lindahl, 2014), new information is accumulating rapidly (e.g. Põlme et
114	al. 2013; Tedersoo et al., 2014; Pärtel et al., 2017; Maestre et al., 2015; Louca et al., 2016),
115	providing suitable data for dark diversity calculations using species co-occurrences without
116	relying on empirical expert opinion about habitat preferences.
117	A potentially suitable target for studying regional and local effects on diversity are the
118	microscopic arbuscular mycorrhizal (AM) fungi (subphylum Glomeromycotina; Spatafora et al.,
119	2016). AM fungi live in symbiosis with the roots of about 80% of terrestrial plant species (Smith
120	& Read, 2008) and provide nutrients (mainly P and N) to their host plants in exchange for plant-
121	assimilated carbon. AM fungi alleviate plant abiotic stress and are able to increase plant
122	resistance to pathogens (Smith & Read, 2008; Pozo et al., 2015). There is accumulating
123	information about the geographic distribution of these fungi (Öpik et al., 2010, 2013; Kivlin et
124	al., 2011; Yang et al., 2012; Tedersoo et al., 2014). Most recently, Davison et al. (2015)

125	analysed AM fungal diversity in plant roots based on systematic sampling of 67 sites globally
126	and found little endemism at the continental scale. At the same time, the diversity of AM fungal
127	communities varied in relation to environmental variables (precipitation, soil organic C content
128	and pH), and spatial distance. The species pool concept promises a more powerful approach for
129	disentangling possible large- and small-scale factors determining AM fungal diversity, such as
130	proximity to centres of evolutionary diversification and the effect of contemporary human
131	influence.
132	AM fungi have several advantages as a model group for studying global diversity patterns and
133	underlying processes. Standardised methodologies for delineating AM fungal taxa (Öpik et al.,
134	2014; Öpik & Davison, 2016) and processing environmental samples exist and are widely used
135	(Hart et al., 2015). DNA-based species delimitation is challenging due to the scarcity of
136	sequences from morphologically described species (Öpik & Davison, 2016), so phylogenetically-
137	delimited sequence groups (phylogroups) are often used (groupings of taxa based on 97%
138	similarity of the target gene sequence; Öpik et al., 2010, 2014). Furthermore, the global diversity
139	of such approximately species-level phylogroups of AM fungi is fairly low (< 2000 groups
140	globally; Öpik et al., 2014; Öpik & Davison, 2016).
141	As well as addressing theoretical challenges concerning the roles of regional and local factors in
142	driving observed diversity patterns, the study of global AM fungal diversity can provide
143	additional specific information about the role of historical factors in shaping the global
144	distribution patterns of these fungi. While Beck et al. (2012) emphasized the significance of
145	integrating past environmental conditions into macroecological analyses, little is known about
146	the effect of historical factors on global microbial diversity. Davison et al. (2015) recorded only
147	a minor effect of continental paleogeographic history on AM fungal community composition.
148	The more recent past, however, might have left an important imprint. For example, during the
149	Quaternary period, glacial periods have been more common than warmer conditions, such as the
150	current interglacial, and biodiversity might be better described by conditions during the most
151	recent glaciation (e.g., the Last Glacial Maximum or LGM) than by contemporary factors
152	(Weigelt et al., 2016). Biomes associated with large species pools might indicate regions where
153	AM fungi have diversified.

Here, we use the framework of the species pool concept to study the effects of regional and local drivers on the diversity of AM fungal communities. We used the MaarjAM database (Öpik et al., 2010) to compile data from all available studies addressing AM fungal molecular (SSU rRNA gene sequence) diversity in environmental samples. The specific objectives of the study were: (1) to quantify and map global patterns in the species pools, local diversity, dark diversity and community completeness of AM fungi; and (2) to link these AM fungal diversity measures to various regional and local drivers, including latitude, current and past (LGM) biome distribution, current and past climate, wilderness index (remoteness from human influence) and local vegetation type. Our results show that species pools, local diversity and dark diversity exhibited significant spatial structure at the global scale. Species pool and dark diversity were related to regional factors (LGM biome configuration and climate), community completeness to local factors (wilderness), and local diversity was jointly associated with regional and local factors (wilderness and current biome configuration).

Materials and Methods

We used the Maarj*AM* database (cf. Öpik *et al.*, 2010; updated in November 2016) as a source of AM fungal distribution data. Maarj*AM* is a curated repository containing AM fungal sequence-based records from published studies, each including information about Virtual Taxa (VT) in a specific geographical location. VT are SSU rRNA gene sequence-based approximately species-level phylogroups of AM fungi, which are phylogenetically delimited on the basis of sequence similarity and clade support (Öpik *et al.*, 2010, 2014). A record in the Maarj*AM* database represents the presence of a VT in a plant species at a site in the case of individual plant root-based records, or the presence of a VT at a site in the case of soil samples or mixed-root samples. The database includes records from both Sanger and 454 sequencing platforms and incorporates 2-3 representative sequences per VT per site or per plant species per site from each study (see Öpik *et al.*, 2010 for details). The Maarj*AM* database currently contains c. 24 000 SSU rRNA gene sequence records associated with c. 400 VT. We associated all records of VT to unique geographical coordinates (sites). We also used information about vegetation type recorded for

183	each site: woodland vegetation (forest, woodland, shrubland) or grassland (both natural and
184	semi-natural). Records from disturbed successional habitats were excluded.
185	For further analysis, we selected only sites that were associated with at least 20 records, since
186	very low numbers of records might not allow precise extrapolations of local diversity. This
187	resulted in a total of 128 sites and 361 VT (Fig. 1a, Table S1).
188	We calculated four related diversity measures: i) species pool size, ii) local diversity, iii) dark
189	diversity (the locally absent fraction of the species pool), and iv) community completeness (the
190	ratio of local and dark diversity). Natural logarithm transformation was used for all these
191	measures to express relative differences. On a log scale, differences indicate how many times
192	diversity values differ, e.g. on a log scale the difference between 5 and 10 VT is equivalent to the
193	difference between 50 and 100 VT rather than the difference between 50 and 55 VT. It should be
194	noted that several of these diversity measures are inherently related (e.g. local and dark diversity
195	are additive components of the species pool), and patterns from these measures are expected to
196	covary. At the same time, the pairs local - dark diversity, and species pool size - community
197	completeness are mathematically independent (Pärtel et al. 2013).
198	In order to estimate species pool size (we use this term for the number of AM fungal VT in the
199	pool for simplicity), it is necessary to sum local diversity and dark diversity. Local diversity was
200	determined from observations at individual sites. The number of records per site ranged from 20
201	to 815 (mean 125). To account for differences in sampling intensity between sites, we used the
202	Shannon index-based effective number of species and extrapolation to an asymptote
203	implemented in the iNEXT software (Hsieh et al., 2016). The asymptotic diversity equates to
204	expected local diversity at full sample coverage sensu Hsieh et al. (2016). This technique made it
205	possible to maximise use of the information in the original data, which would have been lost
206	with rarefying approaches whereby many observations are removed (Chao et al., 2016).
207	Supporting Information Figure S1 shows rarefaction and extrapolation curves for each site. On
208	average, extrapolated local diversity was 1.3 times larger than observed local diversity. The ratio
209	of extrapolated / observed local diversity was not related to sequencing platform and was not
210	strongly spatially clustered (Fig S1b).
211	Dark diversity was estimated using species co-occurrence patterns (Lewis et al., 2016). This
	3-F

otherwise frequently co-occur with those species present at the site. Thus, species that are locally present are used as indicators for absent species: if there are frequent co-occurrences, it is assumed that the species share similar ecological requirements. A co-occurrence index, also known as Beals index, was calculated for each VT in each site. Threshold values for assigning VT to the dark diversity were determined on a VT-by-VT basis since the co-occurrence index depends on species frequency (De Cáceres & Legendre, 2008). For each VT, we examined co-occurrence index values for all sites where it was present and recorded the minimum. Then, if the VT was absent from a site, but its co-occurrence index exceeded the minimum observed in sites where it was present, the VT was considered part of the dark diversity. See Lewis *et al.* (2016) for methodological details and working examples. Community completeness was calculated as the log-ratio of local and dark diversity (Pärtel *et al.*, 2013). Species pool size and community completeness were calculated on the assumption that local and dark diversity estimates represent distinct sets of taxa, i.e. without many overlapping taxa.

Geographical distribution

We predicted the global distribution of the four different diversity measures using Generalized Additive Models (GAMs) and the spline-over-the-sphere algorithm in R package mgcv, with the method 'sos.smooth' and the default arguments except k=30 (Wood, 2003). This model can predict smooth variation in diversity values over the globe without producing edges. For each model, we recorded its estimated degrees of freedom (*edf*), *F* and *P* values, and amount of variation described. We measured the predictive power of the model using cross-validation by dividing locations into random 20% bins and estimating values for bins using the rest of the data (Franklin, 2010). We then calculated the correlation between observed and predicted values. We present only prediction maps when predicted values were significantly correlated with observed values. As a measure of uncertainty in our predictions, we mapped the standard deviation of 100 global predictions using random subsets of 80% of sites.

AM fungal diversity drivers

241	In order to relate diversity values to possible drivers, we obtained measures of the following
242	parameters for each site: (1) latitude, (2) current connectivity to biomes, (3) connectivity to
243	biomes during the LGM, (4) major bioclimatic variables describing current conditions and (5)
244	those during the LGM, (6) wilderness index (remoteness from human influence), and (7) local
245	vegetation type.
246	We measured latitude as distance from the equator (km). Although latitude is not a
247	biogeographic gradient per se and climate and biomes are expected to be more directly related to
248	biodiversity, latitude has been often used in previous studies and we included it to permit
249	comparison.
250	We used the current biome vector map from Olson et al. (2001) and the LGM (ca 21,000 yrs
251	before present) biome vector map from Ray & Adams (2001). The current biome map defines 14
252	biomes, while the original LGM biome map defines 24 biomes. Therefore, we regrouped LGM
253	biomes to match the current classifications (Supporting Information Table S2; Fig. 1b,c). To
254	calculate connectivity to biomes, we constructed a grid of points equally distributed across the
255	globe by using centroids of the ISEA3H geodesic discrete global grid system (Sahr et al., 2003).
256	We used R package 'dggridR' to obtain 65,612 points. We determined biome identity for each
257	point and applied Hanski's connectivity index (Hanski, 1994; Moilanen & Nieminen, 2002):
258	Connectivity = $\sum \exp(-d/a)$; where <i>d</i> is the distance from the site to all terrestrial points of a
259	biome. The parameter a defines the influence of distance in the exponential distribution and can
260	be seen as the average influence distance. We used a values 500, 1000 and 2000 km. To improve
261	its distribution, connectivity was In-transformed for modelling.
262	For each site, we compiled 19 bioclimatic variables (Supporting Information Table S3) (Hijmans
263	et al., 2005) to describe both current conditions and the conditions predicted for the LGM
264	according to the Community Climate System Model (Braconnot et al., 2007). The current
265	climate map had resolution of 5' and the LGM climate map had resolution of 10'. Precipitation
266	measures were ln-transformed. We collapsed the 19 variables to 4 principal components using
267	correlation matrices. The four principal components described >90% of total variation. The first
268	axis was strongly correlated with annual mean and winter temperature (r>0.9), the second axis
269	with precipitation during the dry period (r>0.9). The third axis was more related to precipitation
270	during the warm period $(r>0.6)$ and the fourth axis to modern maximum temperature $(r=0.5)$ or

271	diurnal temperature range during the LGM (r>0.6). See Supporting Information Table S3 for the
272	full correlation table.
273	Wilderness can be defined as a continuous index quantifying remoteness and the level of
274	disturbance by modern technological society (Carver & Fritz, 2016). This synthetic variable was
275	first elaborated for Australia (Lesslie & Taylor, 1985), but later applied globally by UNEP-
276	WCMC (http://www.unep-wcmc.org/resources-and-data/global-wilderness). Available data have
277	a resolution of ca 1.4', and for each site we calculated the mean index value for radiuses of 5, 10
278	and 20 km. It should be noted that we had already excluded disturbed sites, so high wilderness
279	index values were indicative of low human impact in the vicinity of sample sites.
280	We obtained information from original publications about local vegetation type for each site
281	from the MaarjAM database and classified each site broadly as grassland (both natural and semi-
282	natural) or woodland (forest and shrublands). Unfortunately, information about other potential
283	local drivers (e.g. geological and soil characteristics, host plants) was not available for all studied
284	sites.
285	We used an information theoretical approach and compared models using Akaike Information
286	Criterion corrected for sample size (AICc, Burnham & Anderson, 2002). We first standardized
287	all our variables to have equal inputs of mean ±1 standard deviation using the R package 'arm'
288	(Gelman 2008). This allows direct comparisons between model coefficients of both continuous
289	and binary variables. Then we modelled each of the driver types separately. If there were several
290	variables available for a driver type (e.g. connectivity to different biomes, wilderness within
291	different radiuses, Supporting Information Tables S4, S5) we selected the variable for which the
292	model resulted in the lowest AICc values. For latitude, principal components of climate and
293	wilderness, we investigated both linear and quadratic relationships, since unimodal patterns are
294	theoretically possible, and selected the model with the lower AICc value. For connectivity to
295	biomes, we only considered linear models where diversity was positively related to connectivity.
296	In a second step, we examined 29 models: (1) the full model with seven variables, (2) seven
297	univariate models, addressing each driver type in isolation, (3) and all pairwise variable
298	combinations to examine pairs of regional and local drivers in combination. Model assumptions
299	were verified by plotting residuals versus fitted values and each independent variable. We
300	calculated the importance of each driver as the sum of Akaike weights from models where the

301	driver was included. Then we took the top-ranked models ($\Delta AICc < 4$) and used full model
302	averaging to identify the most important variables (Grueber et al., 2011). Several of the
303	independent variables were correlated (e.g. latitude with climate and biomes, or past and current
304	climate; see Supporting Information Table S6 for a correlation matrix). Model averaging,
305	however, is relatively insensitive to such correlations (Freckleton, 2011). Details of the top-
306	ranked model are given in Supporting Information Table S7, of model averaging in Table S8,
307	and a summary of all initial models can be found in Table S9. The R package 'MuMIn' was used
308	for multi-model inference (Bartón, 2016).
309	
310	Results
311	
312	AM fungal local diversity, species pool size, community completeness and dark diversity
313	Average richness was estimated to 60 VT per site (Shannon effective number of taxa), with
314	values ranging between 6 and 216. Species pool size per site was on average 132 VT (range: 46
315	to 285) and dark diversity was on average 71 VT (range: 29 to 145). Relationships between local
316	or dark diversity and species pool size are shown in Fig. 2. As expected, AM fungal local
317	diversity co-varied with AM fungal species pool size but variation in dark diversity introduced
318	considerable variation into this relationship. Local and dark diversity were negatively correlated,
319	although not tightly (Fig. 2c). Average community completeness was slightly negative (-0.37),
320	showing that dark diversity estimates often exceeded local diversity at sites. Variation in
321	community completeness was, however, large (range: -2.7 to 1.3).
322	
323	Global distribution of AM fungal diversity measures
324	AM fungal species pool size and local and dark diversity were non-randomly distributed across
325	the globe. Spatial GAM models accounted for 34% of the variation in AM fungal species pool
326	size (Fig. 1e; $edf=14.1$, $F=1.6$, $P<0.0001$), 12% of the variation in AM fungal local diversity
327	(Fig. 1f; edf =4.8, F =0.4, P =0.016), and 45% of the variation in AM fungal dark diversity (Fig.
328	1g; edf=20.8, F=2.5, P<0.001). Large AM fungal species pools were found in southeastern

329	Africa and eastern South America. Small species pools occurred at higher latitudes of the
330	Northern Hemisphere, especially in North America. Higher local AM fungal diversity values
331	were found in southern South America and southern Africa. North America was characterized by
332	low values. Higher AM fungal dark diversity was found close to the equator, in eastern North
333	America, eastern Australia and New Zealand. Low dark diversity was found in northeastern
334	Asia, western North America and southern South America. Cross-validation revealed moderate
335	correlation between actual and predicted values for the species pool size (r=0.41, P<0.001) and
336	dark diversity (r=0.39, P<0.001), while the correlation between actual and predicted local
337	diversity was indicative of lower predictive power (r=0.20, P=0.025). All predictions for North
338	America (and for New Zealand's dark diversity) were associated with high uncertainty
339	(Supporting Information Fig. S2).
340	The spatial GAM for AM fungal community completeness was non-significant (<i>edf</i> =5.5, <i>F</i> =0.4,
341	P=0.052) and cross-validation showed that actual and predicted values of AM fungal community
342	completeness were not significantly related (r=0.08, P=0.367). Thus, community completeness
343	had no identifiable geographical pattern and is more likely linked to local factors. Therefore, we
344	cannot present a prediction map and present instead a map showing observed values for AM
345	fungal community completeness (Fig. 1h); sites with low and high completeness are frequently
346	found in close proximity.
347	
348	found in close proximity. Relationships with tested regional and local drivers
349	According to driver importance and model averaging, AM fungal species pool size was best
350	described by connectivity to Last Glacial Maximum (LGM) tropical grasslands and savannas
351	(Fig. 3a,b). No other driver had comparable importance or significance (Table S8). For AM
352	fungal local diversity, wilderness around the sample site and current connectivity to mountain
353	grasslands had higher importance (Fig. 3c). Wilderness was significant in model averaging (Fig.
354	3d, Table S8), but current connectivity to mountain grasslands was not (P=0.184, but still
355	significant in the univariate model, Table S8, coef.= 0.23, P=0.009). No clearly important driver
356	of AM fungal dark diversity emerged (Fig. 3e). In the averaged model, AM dark diversity was
357	significantly related to current temperature (PC1, Fig 3f, Table S8). Sites with higher annual or
358	winter temperatures exhibited significantly higher dark diversity estimates.

The degree of wilderness in the surrounding area was important in describing AM fungal community completeness (Fig. 3g) and in the averaged model the relationship was close to significant (P=0.08, Table S8). Wilderness significantly explained community completeness in the model where it was the sole explanatory variable (Fig 3h, Table S9). In bivariate plots, local diversity and community completeness formed triangular-shaped relationships with wilderness (Fig 3e,h): both high and low values of diversity or community completeness were recorded at low wilderness, while only high values were recorded at high wilderness.

Discussion

Here we show that application of the species pool concept to AM fungi can reveal previously undescribed global biodiversity patterns and disentangle the effects of potential underlying drivers. Our results support theoretical expectations that the species pool size is linked to regional (and historical) factors, community completeness is linked to local (and contemporary) factors, and local diversity is a result of both. Using a global data set, we found that the species pool, local diversity and dark diversity of AM fungi showed nonrandom global patterns, with distinct regions of high and low magnitude. By contrast, community completeness did not show significant global structure. AM fungal species pool size was larger in regions that were well connected to tropical grasslands during the Last Glacial Maximum (LGM) *c.* 21,000 y ago. Community completeness was higher at sites with lower human impact in the vicinity (larger wilderness). Local diversity was associated jointly with wilderness around the study site and current connectivity to mountain grasslands. Dark diversity was higher (i.e. a greater number of potentially suitable taxa were absent) in currently warm conditions.

Species pool size is related to historical biome distribution

The largest AM fungal species pools were identified in eastern and southern Africa and to a certain extent in eastern South America. These areas are dominated by tropical grasslands, which, together with sparse dry forests, form a distinct and diverse system called the tropical grassy biome (Parr *et al.*, 2014). We found that AM fungal species pool size was primarily associated with the connectivity to areas of tropical grasslands during the LGM (Ray & Adams,

388	2001). During the LGM, tropical grasslands covered ca 21 million km ² (currently ca 20 million
389	km ²), of which 7 million km ² have remained tropical grassland throughout the past 21000 years
390	and constitute refugia. In fact, parts of the same areas have probably been covered by grasslands
391	since the Miocene (Micheels, 2007). Given that glacial conditions have been more common than
392	interglacials during the Quaternary (Weigelt et al., 2016), biome distribution during the LGM is
393	representative of the predominant environmental configuration through much of recent
394	evolutionary time.
395	The phylogenetic analysis by Davison et al. (2015) suggested that the diversification of the
396	majority of current AM fungal VT occurred approximately within the period of 4-30 million
397	years ago, a timing that is corroborated by other molecular clock estimates for particular AM
398	fungal speciation events (reviewed by Öpik & Davison, 2016). This coincides with the
399	appearance and expansion of grasslands (Strömberg, 2011; Strömberg et al., 2013; Parr et al.,
400	2014). High diversity of macroorganisms in particular habitats has often been associated with
401	high availability of that habitats area in space and through time (Mittelbach et al., 2007). It is
402	possible that developing grasslands created new and spatially (and temporally) very abundant (or
403	'voluminous', since roots occupy the three-dimensional space) habitat for AM fungi. Although
404	the relative area of grasslands in global vegetation has never been very high, these habitats may
405	be particularly relevant for AM fungi due to the high density and large total abundance of host
406	plant roots. For instance, contemporary grasslands contribute about 68% of the global fine root
407	surface area and 78% of global fine root length (Jackson et al., 1997). The difference between
408	forests and grasslands is also evident at small scales: average live fine root length is 4.1 km/m² in
409	tropical evergreen forests but 60.4 km/m ² in tropical grasslands (Jackson et al., 1997). The
410	appearance of this vast new grassland habitat may have led to higher diversification rates of AM
411	fungi due to spatial effects (e.g. isolation by distance in a complex three-dimensional habitat),
412	new niches due to the proliferation and spread of grassland plant species, or other mechanisms.
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414	Local diversity is linked both to regional and local factors
415	In contrast to species pool size, local diversity was most strongly associated with wilderness
416	around study sites. Wilderness is a synthetic measure that is inversely related to human impact
417	(Carver & Fritz, 2016). It incorporates remoteness from modern human infrastructure such as

roads, buildings etc., and a lack of strong human influence such as high-input urban and agricultural areas. In this study, we *a priori* omitted sites that were heavily disturbed, but the wilderness index was calculated within radiuses of 5-20 kilometers around study sites. Thus, our measure of wilderness probably reflected human influence on habitat patches neighbouring the local sites under investigation. In this context, the results indicate that human influence can harm meta-community systems and cause loss of taxa in unaffected patches (Lekberg *et al.*, 2007). Recent overviews show a significant decline in global wilderness (Watson *et al.*, 2016), which may constitute a threat to local AM fungal diversity. Connectivity to current mountain grasslands also had a positive effect on local diversity. The most plausible explanation for this is that it also reflects relatively low human impact in mountainous areas (Sandel & Svenning, 2013).

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Higher dark diversity is recorded in warmer climates

High dark diversity of AM fungi was found at lower latitudes: Central America, Sub-Saharan Africa, eastern Asia and eastern Australia. Modelling also identified current annual temperature as the best predictor of dark diversity. Why a greater share of otherwise suitable taxa should be absent in warm areas is not easy to explain, but indicates either more restricted dispersal or more frequent local extinctions. The sites with high dark diversity were often (sub)tropical moist or dry forests, and dark diversity was higher in woodlands compared to grasslands (although this model had low weight compared with the climate model). Woody vegetation in general hinders wind dispersal of plants (Nathan et al., 2008) and the same might be true for AM fungi. Indeed, forests exhibited higher spatial turnover of AM fungal communities compared to grasslands in a recent global survey of AM fungal communities, and there was also a trend of decreasing forest beta diversity along a latitudinal gradient (Davison et al., 2015). It is conceivable that high spatial heterogeneity in (sub)tropical forests might explain why sampling sites towards the equator lacked a larger number of suitable taxa and dark diversity was consequently higher. However, to properly test this hypothesis we require further empirical studies of spatial structure in AM fungal communities, in particular those inhabiting warmer biomes, such as tropical and subtropical habitats.

447 Community completeness as an indicator of local processes Community completeness of AM fungi varied among study sites but did not exhibit geographic 448 structure. In contrast to species pool size and to a certain extent also to local diversity, variation 449 in community completeness is not expected to contain the footprint of biogeographic history; 450 451 rather it is expected to reflect local factors, such as barriers to dispersal, biotic interactions, or disturbances (Pärtel et al., 2013; Ronk et al., 2015). In our models the best descriptor of AM 452 fungal community completeness was the degree of wilderness around study sites: completeness 453 was high when wilderness was high nearby. Indeed, an adverse impact of intensive land use on 454 AM fungi has been noted in earlier studies (Lopez-Garcia et al., 2013; Moora et al., 2014). 455 However, further specific case studies are needed to disentangle the types of interaction and 456 disturbance that might be responsible for low completeness of AM fungal communities in 457 particular sites. There is evidence that AM fungal taxa with specific traits (ruderal, measured as 458 459 ease of sporulation) are more common in anthropogenic habitats (Ohsowski et al., 2014), 460 possibly caused by differences in tolerance to anthropogenic disturbance (Hart & Reader, 2004; Säle et al. 2015). Alternatively, low wilderness may have a cascading effect through loss of 461 functioning meta-communities within highly human-modified areas. 462 463 Methodological assumptions and potential limitations 464 465 Our findings rest on several methodological assumptions. To identify AM fungi we used phylogroups, in the form of 18S rRNA gene-defined VT, and not traditional taxonomically-466 467 defined species. VT are known to merge closely related morphospecies in some, but not all lineages of AM fungi, and across most of the Glomeromycotina phylogeny there is limited 468 469 information about species boundaries with which to assess the exact taxonomic rank of VT (Öpik et al. 2014; Thiéry et al. 2016). Nonetheless, the rank of VT has been shown to capture 470 471 ecologically-relevant responses to environmental gradients (Powell et al. 2011), suggesting that 472 VT-based estimates of local diversity are meaningful even if precise species boundaries are 473 unknown. For dark diversity estimates obtained using co-occurrence techniques, we assume that VT have similar ecological properties in distant parts of the globe. We are unaware of published 474 evidence with which to assess this assumption. However, we excluded all successional sites 475 where taxa might not be in equilibrium with their environment. We also assume that our local 476

and dark diversity measures can be used in parallel. Theoretically, our estimates of extrapolated local and dark diversity might include taxa present at sites but not recorded. In this case, the species pool size would be overestimated and community completeness would be underestimated. However, we do not expect over- or underestimation to be large. Present but unrecorded species are likely to occur at low abundance, and such species would contribute relatively little to local diversity estimates since the Shannon index counts taxa in proportion to their abundance (Chao *et al.*, 2016). However, we excluded sites for which we expected the sampling effort to be seriously limited. Furthermore, rare taxa often have too few co-occurrences to be included in dark diversity calculations (Ronk *et al.*, 2016). Using observed rather than extrapolated diversity decreased average species pools from 132 to 112 and increased average community completeness from -0.76 to -0.37. Observed and extrapolated estimates of the species pool size and community completeness were strongly correlated (r=0.89, r=0.97, respectively). We anticipate that the accumulation of highly standardised local sampling data using high-throughput methods will further avoid uncertainty related to sampling adequacy and estimation of local and dark diversity.

Conclusions

Community theory predicts that regional drivers are primarily responsible for shaping species pool size, local drivers shape community completeness, and local diversity contains the footprint of both regional and local drivers (Pärtel *et al.*, 2013; Cornell & Harrison, 2014; Zobel, 2016). Nevertheless, comprehensive empirical support for these predictions has been scarce. This study of global diversity patterns in AM fungi provides one of the first large-scale, empirical confirmations of the theory. Furthermore, this study found that the historical distribution of biomes during the LGM was the most important tested regional driver, whereas the degree of wilderness in the vicinity of a study site constituted the most important tested local driver of AM fungal diversity patterns.

Tropical grasslands and savannas harbored the largest species pool of AM fungal species and may thus represent evolutionary hotspots and important refugia. Remoteness from human influence was associated with higher local diversity and greater completeness of AM fungal communities. This is a warning signal that anthropogenic factors have shaped and will continue

507	to shape AM fungal communities to a significant extent. Although human impact on microbial
508	communities has been reported elsewhere, our study provides the first evidence of potential
509	global impacts.
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525	All authors discussed the topic during the 16 th New Phytologist Workshop and following e-mail
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527	analyses. MZ coordinated writing of the paper. All authors discussed results and contributed to
528	writing.
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530	References
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532	Barton K. 2016. MuMIn: Multi-Model Inference. R package version 1.15.6. [WWW document]
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- 718 Figure legends:
- Fig. 1. (a) Sampling locations of AM fungal communities from the MaarjAM database. We
- excluded sites where the number of recorded sequences was <20. Locations are slightly jittered
- to show overlapping points. (b, c) Current (Olson et al., 2001) and Last Glacial Maximum
- 722 (LGM, ca 21000 yrs before present; Ray & Adams, 2001) distribution of biomes: 1: Tropical &
- Subtropical Moist Broadleaf Forests; 2: Tropical & Subtropical Dry Broadleaf Forests; 3:
- 724 Tropical & Subtropical Coniferous Forests; 4: Temperate Broadleaf & Mixed Forests; 5:
- 725 Temperate Conifer Forests; 6: Boreal Forests/Taiga; 7: Tropical & Subtropical Grasslands,
- Savannas & Shrublands; 8: Temperate Grasslands, Savannas & Shrublands; 9: Flooded
- 727 Grasslands & Savannas; 10: Montane Grasslands & Shrublands; 11: Tundra; 12: Mediterranean
- Forests, Woodlands & Scrub; 13: Deserts & Xeric Shrublands; 14: Mangroves; 15: Not
- vegetated. (d) Wilderness (the degree to which a place is remote from and undisturbed by the
- 730 influences of modern technological society; UNEP-WCMC). (e, f, g) Global smoothed maps of
- AM fungal species pool size (GAM, $R^2 = 0.34$), local diversity ($R^2 = 0.12$) and dark diversity (R^2
- 732 = 0.45). (h) Distribution of AM fungal community completeness across study sites. A smoothed
- prediction of is not presented because the predictive power of the corresponding model was low.
- Locations are slightly jittered to distinguish immediately neighbouring points. Colours indicate
- 735 quantiles (e h).
- Fig. 2. Relationships between AM fungal local (a, c), dark diversity (b, c), and species pool size
- 737 (a, b) at 128 sites worldwide. Local diversity was estimated as the asymptotic Shannon index-
- based effective number of taxa using coverage-based rarefaction and extrapolation from site
- records. Dark diversity was estimated based on VT co-occurrences globally (absent VT which
- generally co-occur with locally present VT and therefore likely fit local ecological conditions).
- AM fungal species pool (the theoretical set of VT that can inhabit a study site) is calculated by
- summing AM fungal local and dark diversity. Lines indicate the 1:1 relationship, i.e. the upper
- 743 limit that local or dark diversity can have. Semi-transparent symbols are used to show
- overlapping values. The two outliers with large species pools originate from tropical rainforest in
- French Guiana, and temperate beech forest in Georgia. Local and dark diversity are negatively
- correlated (c, Spearman r = -0.45, P<0.001). Local vegetation type is shown (grasslands or
- 747 woodlands).

Fig.3. Importance of potential drivers (sum of Akaike weights in models where the driver was
included) determining AM fungal species pool size, local and dark diversity, and community
completeness (a, c, e, g). Details on the best supported models are presented in Table S7. Scatter
plots show relationships with the most significant drivers from model averaging (Table S8).
Species pool size is related to the connectivity of LGM tropical grasslands (b, bivariate
relationship: R2=0.17, P=<0.001), local diversity is related to wilderness in the vicinity (d,
R2=0.08, P=0.002), dark diversity is related to current temperature (f, R2=0.14, P<0.001),
community completeness is related to wilderness in the vicinity (h, R2=0.07, P=0.004). Species
pool size, local and dark diversity are ln-transformed, completeness is the logratio of local vs.
dark diversity. Connectivity, wilderness and climate PC1 have relative values without units.

Table S1. Summary of data used in analyses. Geographical coordinates, local vegetation type, number of
 records (representative sequences from a sampling unit), number of Virtual Taxa (VT), primers and
 sequencing platform used, and sources.

No.	Lat.	Lon.	Veg. type	rec	VT	Primers	Seq. Platform	Source			
1	69.8	27.2	woodland	101	57	F: NS31 R: AML2	454 sequencing &	Davison et al. 2015 Science & Opik et			
							Sanger	al. 2013 Mycorrhiza			
2	69.8	27.1	woodland	129	61	F: NS31 R: AML2	454 sequencing &	Davison et al. 2015 Science & Opik et			
							Sanger	al. 2013 Mycorrhiza			
3	61.3	73.1	woodland	75	44	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
4	61.3	73.2	woodland	200	76	F: NS31 R: AML2	454 sequencing &	Davison et al. 2015 Science & Opik et			
							Sanger	al. 2013 Mycorrhiza			
5	59.8	18.0	grassland	61	23	F: NS31 R: AM1	Sanger	Santos-Gonzalez et al. 2007 Applied			
						& F: NS31 R:		and Environmental Microbiology &			
						AM1+AM2+AM3		Santos et al. 2006 New Phytologist			
6	59.2	10.4	woodland	28	11	F: NS31 R: AM1	454 sequencing	Moora et al. 2011 Journal of			
								Biogeography			
7	59.0	26.1	woodland	263	40	F: NS31 R: AM1	Sanger	Davison et al. 2011 FEMS			
			1					Microbiology Ecology & Opik et al.			
								2008 New Phytologist			
8	58.6	23.6	grassland	135	58	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
9	58.6	23.6	grassland	142	87	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
10	58.6	23.6	grassland	88	21	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza			
11	58.4	25.3	woodland	27	11	F: NS31 R: AM1	Sanger	Opik et al. 2003 New Phytologist			
12	58.3	27.3	woodland	78	25	F: NS31 R: AML2	454 sequencing	Davison et al. 2012 PLoS ONE			
13	58.2	26.6	grassland	28	14	F: NS31 R: AM1	Sanger	Opik et al. 2003 New Phytologist			
14	56.1	159.9	woodland	94	56	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
15	56.1	159.9	woodland	102	58	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
16	56.1	159.9	woodland	40	15	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza			
17	55.5	-2.2	grassland	57	29	F: NS31 R: AM1	Sanger	Vandenkoornhuyse et al. 2007			
								Proceedings of the National Academy			
								of Sciences of the United States of			
								America			
18	54.1	-0.9	-0.9 woodland 79 33 F: NS31 R: AM1 Sanger		Sanger	Helgason et al. 1998 Nature &					
								Helgason et al. 1999 Molecular			
								Ecology & Helgason et al. 2002 Journal			
								of Ecology & Helgason et al. 2007			
								Journal of Ecology			
19	53.9	-1.4	grassland	36	26	F: NS31 R: AM1	Sanger	Dumbrell et al. 2010 Journal of			
								Ecology			
20	53.0	158.7	woodland	54	32	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
21	53.0	158.7	woodland	77	41	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
22	53.0	158.7	woodland	55	14	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza			
23	52.7	4.7	grassland	36	16	F: NS31 R: AM1	Sanger	Scheublin et al. 2004 Applied and			
	1							Environmental Microbiology			
24	50.8	-104.6	grassland	509	115	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science & Opik et			
_	1							al. 2013 Mycorrhiza			
25	48.5	-79.3	woodland	24	11	F: NS31 R: AM1	Sanger	DeBellis & Widden 2006 FEMS			
	1	_						Microbiology Ecology			
26	47.8	107.1	grassland	206	67	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
27	47.8	107.1	grassland	261	93	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
28	47.5	10.1	grassland	106	63	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
29	47.5	10.1	grassland	101	60	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
30	46.6	16.0	grassland	20	16	F: NS31 R: AM1	Sanger	Macek et al. 2011 Applied and			
								Environmental Microbiology			
31	44.8	-0.4	woodland	175	69	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			

No.	Lat.	Lon.	Veg. type	rec	VT	Primers	Seq. Platform	Source		
32	43.6	-1.2			F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
33	43.5	104.1	grassland	239	78	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
34	43.0	104.1	grassland	179	69	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
35	42.0	116.3	grassland	27	20	F: NS31 R: AML2	Sanger	Chen et al. 2014 Soil Biology and		
								Biochemistry		
36	41.9	43.4	woodland	68	41	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
37	41.9	43.4	woodland	53	21	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza		
38	41.9	43.4	woodland	73	58	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
39	41.6	-79.5	woodland	25	7	F: NS31 R: AM1	Sanger	Burke 2008 American Journal of		
	40.0			-				Botany		
40	40.2	-111.1	grassland	22	8	F: VANS1 or	Sanger	Winther & Friedman 2007 American		
						GEOA2 or GEO11		Journal of Botany		
						R: GLOM1311R				
44	20.2	06.0		00	40	or SS1492	454	D : 1 2045 C :		
41	39.2	-86.2	woodland	90	49	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
42	39.2	-86.2	woodland	95	56	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
43	39.1	-96.6	grassland	37	15	F: NS31 R: AM1	Sanger	Jumpponen et al. 2005 Biology and Fertility of Soils		
44	39.0	-123.1	grassland	35	14	F: NS31 R: AM1	Sanger	Hausmann & Hawkes 2009 New		
		1			•			Phytologist		
45	38.7	140.7	grassland	51	30	F: AMV4.5NF R: AMV4.5NR	Sanger	Saito et al. 2004 Mycorrhiza		
46	38.7	-0.9	woodland	76	29	F: NS31 R: AM1	Sanger	Alguacil et al. 2009 Environmental		
						& F: NS31 R:		Microbiology & Alguacil et al. 2009		
						AM1+AM2+AM3		Microbial Ecology		
47	38.2	-1.2	woodland	150	32	F: AML1 R: AML2	Sanger	Alguacil et al. 2011 Science of the		
								Total Environment & Alguacil et al.		
								2011 Soil Biology and Biochemistry &		
								Torrecillas et al. 2012 Applied and		
								Environmental Microbiology		
48	38.2	-1.8	woodland	25	10	F: NS31 R:	Sanger	Alguacil et al. 2009 Applied and		
						AM1+AM2+AM3		Environmental Microbiology		
49	37.7	-1.7	woodland	71	21	F: AML1 R: AML2	Sanger	Alguacil et al. 2012 Soil Biology and		
								Biochemistry		
50	37.4	-2.8	woodland	726	71	F: NS31 R: AM1	454 sequencing &	Palenzuela et al. 2012 Journal of Arid		
						& F: NS31 R:	Sanger	Environments & Sanchez-Castro et al		
						AML2		2012 Mycorrhiza & Varela-Cervero et		
								al. 2015 Environmental Microbiology		
51	36.0	101.9	grassland	146	39	F: NS31 R: AML2	Sanger	Liu et al. 2012 New Phytologist		
52	35.6	-116.2	grassland	61	24	F: NS31 R: AM1	Sanger	Schechter, S. P.; Bruns, T. D. 2013		
								PLoS ONE & Schechter, S.P.; Bruns,		
								T.D. 2008 Molecular Ecology		
53	35.2	135.4	woodland	29	8	F: NS31 R: AM1	Sanger	Yamato & Iwase 2005 Mycoscience		
54	35.0	102.9	grassland	47	23	F: NS31 R: AML2	Sanger	Shi et al. 2014 PLoS ONE		
55	33.7	101.9	grassland	68	33	F: NS31 R: AML2	Sanger	Shi et al. 2014 PLoS ONE		
56	30.6	34.7	woodland	96	67	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
57	30.6	34.7	woodland	95	57	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
58	30.6	34.7	woodland	66	35	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza		
59	29.5	118.1	woodland	42	18	F: NS31 R: AM1	454 sequencing	Moora et al. 2011 Journal of		
		1				& F: NS31 R:		Biogeography & Opik et al. 2013		
		<u> </u>				AML2		Mycorrhiza		
60	29.5	118.1	woodland	47	20	F: NS31 R: AM1	454 sequencing	Moora et al. 2011 Journal of		
		1				& F: NS31 R:		Biogeography & Opik et al. 2013		
		1				AML2		Mycorrhiza		
61	29.4	79.6	woodland	153	72	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		
62	29.4	79.6	woodland	162	77	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science		

No.	Lat.	Lon.	Veg. type	rec	VT	Primers	Seq. Platform	Source				
63	29.4	118.2	woodland	63	28	F: NS31 R: AM1	454 sequencing	Moora et al. 2011 Journal of				
						& F: NS31 R:		Biogeography & Opik et al. 2013				
						AML2		Mycorrhiza				
64	28.7	77.2	woodland	27	12	F: NS31 R: AM1	Sanger	Deepika & Kothamasi 2015				
								Mycorrhiza				
65	22.4	81.9	woodland	158	83	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
66	22.4	81.9	woodland	169	76	F: NS31 R: AML2 454 sequencing		Davison et al. 2015 Science				
67	20.1	-75.1	grassland	28	8	F: AML1 R: AML2	Sanger	Alguacil et al. 2012 PLoS ONE				
68	16.9	100.5	woodland	215	99	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
69	16.9	100.5	woodland	77	28	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza				
70	15.2	-23.7	woodland	61	21	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza				
71	14.6	-17.0	grassland	136	81	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
72	14.6	-17.0	grassland	137	74	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
73	9.2	-79.9	woodland	63	34	F: NS31 R: AM1	Sanger	Husband et al. 2002 Molecular				
								Ecology & Husband et al. 2002 FEMS				
								Microbiology Ecology				
74	9.0	38.6	woodland	23	12	F: GlomerWT0 R:	Sanger	Wubet et al. 2006 Canadian Journal of				
						one of either		Botany & Wubet et al. 2006				
						GlomerWT1, GlomerWT2,		Mycological Research				
						GlomerWT3, or						
						GlomerWT4						
75	5.3	-52.9	woodland	34	27	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
76	5.3	-52.9	woodland	65	57	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
77	5.3	-52.9	woodland	61	25	F: NS31 R: AML2	454 sequencing	Opik et al. 2013 Mycorrhiza				
78	5.0	9.6	woodland	23	9	F: NS1 R: ITS4 &	Sanger	Franke et al. 2006 Mycological				
'	3.0	3.0	Woodiana			F: NS31 R: AM1	Sanger	Progress & Merckx & Bidartondo 2008				
								Proceedings of The Royal Society B				
79	4.6	-52.2	woodland	44	34	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
80	4.6	-52.2	woodland	55	44	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
81	4.6	-52.2	woodland	66	32	F: NS31 R: AML2	454 sequencing	Opik et al. 2013 Mycorrhiza				
82	0.6	10.4	woodland	297	82	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science & Opik et				
								al. 2013 Mycorrhiza				
83	0.6	10.4	woodland	249	93	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science & Opik et				
								al. 2013 Mycorrhiza				
84	-1.8	35.2	grassland	46	34	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
85	-1.8	35.2	grassland	75	60	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
86	-2.1	35.0	grassland	86	64	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
87	-2.3	34.5	grassland	90	59	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
88	-2.6	35.1	grassland	75	53	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
89	-2.7	35.1	grassland	141	68	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science				
90	-5.9	145.1	woodland	37	21	F: SSU817F R:	454 sequencing	Tedersoo et al. 2015 Science				
						SSU1196ngs						
91	-7.3	147.1	woodland	92	47	F: SSU817F R:	454 sequencing	Tedersoo et al. 2015 Science				
						SSU1196ngs						
92	-9.4	147.4	woodland	127	65	F: SSU817F R:	454 sequencing	Tedersoo et al. 2015 Science				
						SSU1196ngs						
93	-18.9	34.4	grassland	27	15	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New				
								Phytologist				
94	-18.9	34.4	grassland	54	27	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New				
								Phytologist				
95	-18.9	34.5	grassland	37	17	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New				
								Phytologist				
96	-18.9	34.5	grassland	57	28	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New				
	1	1	<u> </u>					Phytologist				
97	-18.9	34.5	grassland	33	19	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New				

No.	Lat.	Lon.	Veg. type	rec	VT	Primers	Seq. Platform	Source			
							-	Phytologist			
98	-18.9	34.4	grassland	71	34	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New			
								Phytologist			
99	-18.9	34.5	grassland	95	42	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New			
								Phytologist			
100	-18.9	34.4	grassland	119	52	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New			
								Phytologist			
101	-19.0	34.4	grassland	67	44	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New			
								Phytologist			
102	-19.0	34.4	grassland	180	84	F: NS31 R: AML2	ML2 454 sequencing Rodriguez-Echeverria et a				
								Phytologist			
103	-19.0	34.2	grassland	150	74	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 Nev			
								Phytologist			
104	-19.0	34.2	grassland	181	94	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 Nev			
								Phytologist			
105	-19.0	34.2	grassland	122	66	F: NS31 R: AML2	454 sequencing	Rodriguez-Echeverria et al. 2017 New			
								Phytologist			
106	-23.8	133.9	woodland	58	14	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza			
107	-23.8	133.9	woodland	156	70	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
108	-23.8	133.9	woodland	157	82	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
109	-24.7	28.7	grassland	222	76	F: NS31 R: AML2	454 sequencing &	Davison et al. 2015 Science & Opik et			
							Sanger	al. 2013 Mycorrhiza			
110	-24.8	28.6	grassland	234	100	F: NS31 R: AML2	454 sequencing &	Davison et al. 2015 Science & Opik et			
							Sanger	al. 2013 Mycorrhiza			
111	-28.6	-51.6	grassland	298	76	F: NS31 R: AML2	454 sequencing	Zobel et al., in prep.			
112	-30.1	-51.7	grassland	487	103	F: NS31 R: AML2	454 sequencing	Zobel et al., in prep.			
113	-31.2	-64.3	woodland	100	49	F: NS31 R: AML2	454 sequencing	Grilli et al. 2015 Environmental			
								Microbiology			
114	-32.8	-64.9	grassland	261	85	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science & Opik et			
								al. 2013 Mycorrhiza			
115	-32.8	-64.9	grassland	287	84	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science & Opik et			
								al. 2013 Mycorrhiza			
116	-33.7	151.2	woodland	42	12	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza			
117	-33.7	151.2	woodland	55	38	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
118	-33.7	151.2	woodland	34	23	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
119	-34.0	19.0	woodland	108	44	F: NS31 R: AML2	454 sequencing &	Davison et al. 2015 Science & Opik et			
							Sanger	al. 2013 Mycorrhiza			
120	-34.0	19.0	woodland	100	41	F: NS31 R: AML2	454 sequencing &	Davison et al. 2015 Science & Opik et			
							Sanger	al. 2013 Mycorrhiza			
121	-35.1	138.7	woodland	85	32	F: NS31 R: AML2	454 sequencing	Opik et al. 2013 Mycorrhiza			
122	-35.1	138.7	woodland	227	86	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
123	-37.3	142.2	grassland	71	21	F: NS31 R: AML2	Sanger	Opik et al. 2013 Mycorrhiza			
124	-37.3	142.2	grassland	271	71	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
125	-39.0	-71.4	woodland	778	75	F: NS31 R: AML2	454 sequencing	Gazol et al. 2016 FEMS Microbiology			
					<u> </u>			Ecology			
126	-39.0	-71.4	woodland	815	81	F: NS31 R: AML2	454 sequencing	Gazol et al. 2016 FEMS Microbiology			
		<u>L</u>	<u> </u>					Ecology			
127	-52.1	-71.4	grassland	190	79	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			
128	-52.1	-71.4	grassland	223	75	F: NS31 R: AML2	454 sequencing	Davison et al. 2015 Science			

Fig. S1. (a) Shannon index based effective number of species for sites with varying numbers of records (number of representative sequences from a sampling unit in a site). Red lines show rarefaction and blue lines extrapolations. We used estimated local diversity extrapolated to the asymptote, i.e. full

768 769 sample coverage *sensu* Hsieh *et al.* (2016). (b) Increase due to extrapolation (extrapolated / observed local diversity) and sequencing platform within study sites. Locations are slightly jittered to show overlapping points.

Table S2. Homogenization of biome classifications between current and Last Glacial Maximum (LGM)
 maps.

ID	Current	LGM
1	Tropical & Subtropical Moist Broadleaf Forests	Tropical rainforest
2	Tropical & Subtropical Dry Broadleaf Forests	Tropical woodland Monsoon or dry forest Tropical thorn scrub and scrub woodland
3	Tropical & Subtropical Coniferous Forests	Montane tropical forest
4	Temperate Broadleaf & Mixed Forests	Broadleaved temperate evergreen forest
5	Temperate Conifer Forests	
6	Boreal Forests/Taiga	Open boreal woodlands Main Taiga
7	Tropical & Subtropical Grasslands, Savannas & Shrublands	Tropical grassland Savanna
8	Temperate Grasslands, Savannas & Shrublands	Temperate steppe grassland Forest steppe Dry steppe
9	Flooded Grasslands & Savannas	
10	Montane Grasslands & Shrublands	Alpine tundra Montane Mosaic Subalpine parkland
11	Tundra	Tundra Steppe-tundra Polar and alpine desert
12	Mediterranean Forests, Woodlands & Scrub	Semi-arid temperate woodland or scrub
13	Deserts & Xeric Shrublands	Tropical semi-desert Tropical extreme desert Temperate desert Temperate semi-desert
14	Mangroves	
15	Not vegetated	Not vegetated

Table S3. Correlation matrix of Bioclimatic PCA from current and Last Glacial Maximum predictions (LGM). Very high correlations r>0.9 are indicated by coloured backgrounds.

	Currer	nt climat	e		LGM climate			
Climatic parameter	PC1	PC2	PC3	PC4	PC1	PC2	PC3	PC4
BIO1 = Annual Mean Temperature	0.94	-0.26	-0.09	0.15	0.95	-0.23	-0.14	0.05
BIO2 = Mean Diurnal Range	0.11	-0.68	0.13	0.18	-0.45	0.24	0.43	0.66
(Mean of monthly (max temp - min								
temp))								
BIO3 = Isothermality (BIO2/BIO7)	0.85	-0.09	-0.15	- 0.25	0.6	0.28	0.23	0.65
BIO4 = Temperature Seasonality	-	-0.07	0.30	0.30	-0.84	-0.2	0.01	-0.35
(standard deviation *100)	0.88							
BIO5 = Max Temperature of Warmest	0.68	-0.50	0.05	0.50	0.82	-0.32	-0.10	0.13
Month								
BIO6 = Min Temperature of Coldest	0.96	-0.04	-0.25	-	0.97	-0.13	-0.20	0.04
Month				0.02				
BIO7 = Temperature Annual Range (BIO5-	-0.8	-0.27	0.35	0.35	-0.87	-0.05	0.23	0.04
BIO6)								
BIO8 = Mean Temperature of Wettest	0.72	-0.30	0.40	0.30	0.85	-0.37	0.00	-0.17
Quarter								
BIO9 = Mean Temperature of Driest	0.86	-0.16	-0.42	0.01	0.92	-0.11	-0.28	0.16
Quarter								
BIO10 = Mean Temperature of Warmest	0.76	-0.42	0.06	0.45	0.87	-0.36	-0.18	-0.07
Quarter								
BIO11 = Mean Temperature of Coldest	0.97	-0.14	-0.19	0.00	0.97	-0.13	-0.12	0.13
Quarter								
BIO12 = Annual Precipitation	0.63	0.68	0.30	-	0.73	0.58	0.27	-0.13
				0.05				
BIO13 = Precipitation of Wettest Month	0.72	0.38	0.49	_	0.83	0.25	0.41	-0.17
				0.20				
BIO14 = Precipitation of Driest Month	0.07	0.92	-0.09	0.29	0.09	0.94	-0.17	-0.09
BIO15 = Precipitation Seasonality	0.31	-0.72	0.42		0.37	-0.78	0.39	-0.07
(Coefficient of Variation)				0.36				
BIO16 = Precipitation of Wettest Quarter	0.73	0.40	0.47	-	0.82	0.29	0.40	-0.17
				0.17				
BIO17 = Precipitation of Driest Quarter	0.14	0.91	0.01	0.32	0.19	0.94	-0.17	-0.12
BIO18 = Precipitation of Warmest Quarter	0.35	0.43	0.69	0.00	0.51	0.27	0.62	-0.33
BIO19 = Precipitation of Coldest Quarter	0.24	0.79	-0.35	0.19	0.27	0.84	-0.33	-0.01

778	
779 780 781	Table S4. Correlation between connectivity of biomes using different distances of influence (500, 1000 and 2000 km). We show only connectivity of biomes that had high importance: cur.10 – current mountain grasslands and shrublands, lgm.7 – Last Glacial Maximum tropical grasslands and savannas.
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785 786	Table S5. Correlation between wilderness measures using different radiuses (5, 10 and 20 km) around study sites.
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789 790 791 792 793 794 795	Table S6. Correlations between independent variables used in models: absolute latitude (abs.lat), connectivity to current and Last Glacial Maximum (LGM) biomes (cur# and lgm#, respectively: see numerical codes of biomes in Fig 1 or Table S1), four current and LGM climate principal components (PC#, PC#lgm, see Table S2 for numerical codes), wilderness and local vegetation type (grassland vs. woodland). For connectivity of biomes we included only the mean distance of influence 1000 km; other distances were highly correlated (see Table S4). For Wilderness we included here only radius of 10 km; other radiuses gave highly correlated values (see Table S5).
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Table S7. Top-ranked models (delta AICc < 4). All variables were standardized with 2 sd values. Polynomial fits are indicted by "+". See model averaging and details about variables in Table S8.

Study variable	Absolute latitude	Conn. current biomes	Conn. LGM biomes	Current climate	LGM climate	Wilderness	Vegetation type = grassland	adjR²	df	logLik	AICc	Delta AICc	Akaike weight
Species pool size	+		0.43					0.27	5	-77.9	166.3	0.00	0.23
3pecies poor 3ize	·		0.35	+				0.26	5	-78.1	166.7	0.38	0.19
			0.31		+			0.26	5	-78.2	166.9	0.56	0.17
			0.41					0.22	3	-80.5	167.1	0.80	0.16
			0.42				-0.1	0.23	4	-80.1	168.5	2.22	0.08
			0.38			0.07		0.23	4	-80.2	168.7	2.36	0.07
		0.0	0.42					0.22	4	-80.5	169.2	2.92	0.05
					+			0.22	4	-80.6	169.5	3.21	0.05
Local diversity				0.22		0.24		0.16	4	-83.9	176.2	0.00	0.73
			0.18			0.20		0.13	4	-85.5	179.3	3.07	0.16
				-0.1		0.25		0.13	4	-85.9	180.1	3.86	0.11
						4							
Dark diversity	-0.1	-0.4	0.28	0.57	+	-0.2	-0.1	0.38	10	-70.8	163.4	0.00	0.77
				0.44		-0.2		0.24	4	-79.4	167.2	3.76	0.12
				0.36			-0.2	0.24	4	-79.5	167.3	3.92	0.11
Community completeness			0.21			0.22		0.14	4	-85.1	178.5	0.00	0.25
		0.2				0.23		0.14	4	-85.2	178.7	0.22	0.23
	0.19	0.09	0.19	-0.1	-0.1		0.07	0.23	9	-80.1	179.7	1.21	0.14
					-0.2	0.26		0.12	4	-86.1	180.5	1.94	0.10
			0.00	-0.1		0.22		0.11	4	-86.6	181.5	2.97	0.06
		0.1-	0.22	-0.2				0.11	4	-86.7	181.8	3.30	0.05
		0.17	0.19			0.00	0.11	0.11	4	-86.7	181.8	3.30	0.05
			0.22			0.23	0.14	0.11	4	-86.8	181.9	3.37	0.05
			0.22			0.55	0.16	0.11	4	-86.8	182.0	3.49	0.04
						0.26		0.09	3	-88.1	182.3	3.81	0.04

Table S8. Averaged models (full average) from top-ranked models (delta AICc<4, see Table S7). All variables were standardized with 2 sd values. Variables with P<0.1 are marked by bold font.

Study variable	Predictors	Coef.	Adj. SE	z value	Р
Species pool size	Connectivity to LGM tropical grasslands	0.37	0.16	2.29	0.022
	Absolute latitude	0.01	0.44	0.02	0.982
	Absolute latitude ²	0.24	0.48	0.49	0.626
	Current climate PC1 (temperature)	0.08	0.38	0.20	0.845
	Current climate PC1 (temperature) ²	0.18	0.43	0.43	0.667
	LGM climate PC1 (temperature)	0.21	0.58	0.35	0.725
	LGM climate PC1 (temperature) ²	0.22	0.46	0.47	0.640
	Vegetation type (grassland)	-0.01	0.03	0.18	0.859
	Wilderness	0.01	0.03	0.16	0.873
	Connectivity to current tropical moist forests	0.00	0.03	0.02	0.988
Local diversity	Connectivity to current mountain grasslands	0.16	0.12	1.33	0.184
	Wilderness	0.23	0.09	2.63	0.009
	Connectivity to LGM tropical grasslands	0.03	0.08	0.38	0.706
	Current climate PC4 (temp. warm periods)	-0.02	0.05	0.29	0.770
Dark diversity	Absolute latitude	-0.11	0.24	0.45	0.650
	Current climate PC1 (temperature)	0.53	0.27	2.00	0.046
	Connectivity to current mangroves	-0.28	0.21	1.32	0.188
	Connectivity to LGM tropical dry forests	0.20	0.15	1.51	0.130
	LGM climate PC1 (temperature)	-0.39	1.42	0.28	0.781
	LGM climate PC1 (temperature) ²	0.71	0.64	1.11	0.268
	Vegetation type (grassland)	-0.13	0.09	1.37	0.170
	Wilderness	-0.18	0.12	1.54	0.124
Community completeness	Connectivity to LGM deserts	0.11	0.12	0.90	0.368
	Wilderness	0.22	0.12	1.73	0.083
	Connectivity to current mountain grasslands	0.07	0.10	0.64	0.519
	Absolute latitude	0.03	0.08	0.35	0.727
	Current climate PC4 (temp. warm periods)	-0.03	0.07	0.40	0.687
	LGM climate PC4 (prec. dry periods)	-0.03	0.07	0.40	0.693
	Vegetation type (grassland)	0.02	0.06	0.37	0.712

Table S9. Details all models tested. Four dependent diversity measures (AM fungal species pool size, local diversity, dark diversity, and community completeness) are related to seven driver types: absolute latitude, connectivity to current and LGM biomes (see biome numbers from Tables S1, three distance of influence are used, 500 km, 1000 km and 2000 km, models with coefficient >0 are given since the negative connectivity has no biological meaning here), current and LGM climate (four principal components, PC1...PC4), wilderness index (mean value in radiuses 5 km 10 km and 20 km) and local vegetation type (grassland vs. woodland). For latitude, climate and wilderness both linear and polynomial models have been considered. Coefficients are comparable since all variables were standardized with 2 sd.

Study variable	Driver type	predictors	Coef	SE	t value	Р	AICc	R ²
sp.pool.size	abs.lat	abs.lat	-0.37	0.08	-4.4	<0.001	172.4	0.14
sp.pool.size	abs.lat	poly(abs.lat, 2)1	-2.07	0.46	-4.5	<0.001	171.3	0.16
sp.pool.size	abs.lat	poly(abs.lat, 2)2	0.82	0.46	1.8	0.077	171.3	0.16
sp.pool.size	cur.biomes	cur.13.500	0.00	0.09	0.0	0.983	191.0	0.00
sp.pool.size	cur.biomes	cur.2.500	0.26	0.09	3.1	0.003	181.7	0.07
sp.pool.size	cur.biomes	cur.2.1000	0.23	0.09	2.6	0.011	184.3	0.05
sp.pool.size	cur.biomes	cur.2.2000	0.14	0.09	1.6	0.108	188.4	0.02
sp.pool.size	cur.biomes	cur.14.500	0.29	0.09	3.4	0.001	179.8	0.08
sp.pool.size	cur.biomes	cur.14.1000	0.27	0.09	3.2	0.002	181.3	0.07
sp.pool.size	cur.biomes	cur.14.2000	0.23	0.09	2.6	0.010	184.2	0.05
sp.pool.size	cur.biomes	cur.7.500	0.31	0.08	3.7	< 0.001	177.8	0.10
sp.pool.size	cur.biomes	cur.7.1000	0.31	0.08	3.7	<0.001	177.8	0.10
sp.pool.size	cur.biomes	cur.7.2000	0.31	0.08	3.6	<0.001	178.2	0.10
sp.pool.size	cur.biomes	cur.1.500	0.34	0.08	4.1	< 0.001	175.3	0.12
sp.pool.size	cur.biomes	cur.1.1000	0.33	0.08	4.0	<0.001	176.0	0.11
sp.pool.size	cur.biomes	cur.1.2000	0.30	0.09	3.5	0.001	179.2	0.09
sp.pool.size	cur.biomes	cur.10.500	0.27	0.09	3.1	0.002	181.6	0.07
sp.pool.size	cur.biomes	cur.10.1000	0.23	0.09	2.6	0.010	184.2	0.05
sp.pool.size	cur.biomes	cur.10.2000	0.12	0.09	1.3	0.186	189.2	0.01
sp.pool.size	cur.biomes	cur.9.500	0.02	0.09	0.2	0.866	191.0	0.00
sp.pool.size	cur.biomes	cur.9.1000	0.05	0.09	0.6	0.573	190.7	0.00
sp.pool.size	cur.biomes	cur.9.2000	0.08	0.09	1.0	0.343	190.1	0.01
sp.pool.size	Igm.biomes	lgm.12.500	0.02	0.09	0.2	0.833	191.0	0.00
sp.pool.size	lgm.biomes	lgm.13.500	0.14	0.09	1.5	0.128	188.6	0.02
sp.pool.size	lgm.biomes	lgm.13.1000	0.16	0.09	1.8	0.080	187.9	0.02
sp.pool.size	lgm.biomes	lgm.13.2000	0.16	0.09	1.8	0.073	187.7	0.03
sp.pool.size	lgm.biomes	lgm.2.500	0.05	0.09	0.5	0.603	190.7	0.00
sp.pool.size	lgm.biomes	lgm.2.1000	0.09	0.09	1.0	0.314	190.0	0.01
sp.pool.size	lgm.biomes	lgm.2.2000	0.11	0.09	1.2	0.234	189.5	0.01
sp.pool.size	lgm.biomes	lgm.1.500	0.32	0.08	3.7	<0.001	177.6	0.10
sp.pool.size	Igm.biomes	lgm.1.1000	0.27	0.09	3.1	0.002	181.3	0.07
sp.pool.size	lgm.biomes	lgm.1.2000	0.17	0.09	2.0	0.050	187.1	0.03
sp.pool.size	lgm.biomes	lgm.7.500	0.38	0.08	4.6	< 0.001	170.8	0.15
sp.pool.size	lgm.biomes	lgm.7.1000	0.41	0.08	5.1	< 0.001	167.1	0.17
sp.pool.size	lgm.biomes	lgm.7.2000	0.40	0.08	4.9	<0.001	169.1	0.16
sp.pool.size	lgm.biomes	lgm.3.500	0.24	0.09	2.7	0.007	183.6	0.06
sp.pool.size	lgm.biomes	lgm.3.1000	0.18	0.09	2.0	0.047	187.0	0.03
sp.pool.size	lgm.biomes	lgm.3.2000	0.13	0.09	1.5	0.138	188.8	0.02
sp.pool.size	lgm.biomes	lgm.4.500	0.08	0.09	0.9	0.383	190.2	0.01
sp.pool.size	lgm.biomes	lgm.4.1000	0.08	0.09	0.9	0.382	190.2	0.01

Study variable	Driver type	predictors	Coef	SE	t value	Р	AICc	R ²
sp.pool.size	lgm.biomes	lgm.4.2000	0.08	0.09	0.9	0.382	190.2	0.01
sp.pool.size	cur.climate	PC1	0.36	0.08	4.3	< 0.001	173.4	0.13
sp.pool.size	cur.climate	poly(PC1, 2)1	2.02	0.46	4.4	< 0.001	170.9	0.16
sp.pool.size	cur.climate	poly(PC1, 2)2	0.99	0.46	2.2	0.034	170.9	0.16
sp.pool.size	cur.climate	PC2	0.00	0.09	-0.1	0.963	191.0	0.00
sp.pool.size	cur.climate	poly(PC2, 2)1	-0.02	0.50	-0.1	0.963	193.1	0.00
sp.pool.size	cur.climate	poly(PC2, 2)2	-0.02	0.50	0.0	0.973	193.1	0.00
sp.pool.size	cur.climate	PC3	0.14	0.09	1.6	0.114	188.5	0.02
sp.pool.size	cur.climate	poly(PC3, 2)1	0.79	0.50	1.6	0.114	189.3	0.03
sp.pool.size	cur.climate	poly(PC3, 2)2	-0.55	0.50	-1.1	0.267	189.3	0.03
sp.pool.size	cur.climate	PC4	-0.10	0.09	-1.2	0.242	189.6	0.01
sp.pool.size	cur.climate	poly(PC4, 2)1	-0.59	0.50	-1.2	0.242	190.4	0.02
sp.pool.size	cur.climate	poly(PC4, 2)2	-0.57	0.50	-1.2	0.252	190.4	0.02
sp.pool.size	lgm.climate	PC1	0.36	0.08	4.3	< 0.001	173.2	0.13
sp.pool.size	lgm.climate	poly(PC1, 2)1	2.03	0.46	4.4	< 0.001	169.5	0.17
sp.pool.size	lgm.climate	poly(PC1, 2)2	1.10	0.46	2.4	0.018	169.5	0.17
sp.pool.size	lgm.climate	PC2	-0.03	0.09	-0.4	0.720	190.9	0.00
sp.pool.size	lgm.climate	poly(PC2, 2)1	-0.18	0.50	-0.4	0.722	193.0	0.00
sp.pool.size	lgm.climate	poly(PC2, 2)2	-0.01	0.50	0.0	0.990	193.0	0.00
sp.pool.size	lgm.climate	PC3	0.07	0.09	0.8	0.400	190.3	0.01
sp.pool.size	lgm.climate	poly(PC3, 2)1	0.42	0.50	0.9	0.400	191.0	0.02
sp.pool.size	lgm.climate	poly(PC3, 2)2	-0.58	0.50	-1.2	0.248	191.0	0.02
sp.pool.size	lgm.climate	PC4	-0.11	0.09	-1.3	0.212	189.4	0.01
sp.pool.size	lgm.climate	poly(PC4, 2)1	-0.63	0.50	-1.3	0.212	190.3	0.02
sp.pool.size	lgm.climate	poly(PC4, 2)2	-0.56	0.50	-1.1	0.263	190.3	0.02
sp.pool.size	wild	wild.5	0.19	0.09	2.2	0.028	186.1	0.04
sp.pool.size	wild	poly(wild.5, 2)1	1.09	0.49	2.2	0.029	188.2	0.04
sp.pool.size	wild	poly(wild.5, 2)2	-0.03	0.49	-0.1	0.945	188.2	0.04
sp.pool.size	wild	wild.10	0.20	0.09	2.2	0.027	186.0	0.04
sp.pool.size	wild	poly(wild.10, 2)1	1.10	0.49	2.2	0.028	188.0	0.04
sp.pool.size	wild	poly(wild.10, 2)2	-0.22	0.49	-0.4	0.663	188.0	0.04
sp.pool.size	wild	wild.20	0.23	0.09	2.7	0.009	184.0	0.05
sp.pool.size	wild	poly(wild.20, 2)1	1.30	0.49	2.7	0.009	185.9	0.05
sp.pool.size	wild	poly(wild.20, 2)2	-0.24	0.49	-0.5	0.629	185.9	0.05
sp.pool.size	veg.type	veg.type = grassl.	-0.02	0.09	-0.3	0.792	190.9	0.00
local.diversity	abs.lat	abs.lat	-0.16	0.09	-1.8	0.080	187.9	0.02
local.diversity	abs.lat	poly(abs.lat, 2)1	-0.87	0.49	-1.8	0.079	187.8	0.04
local.diversity	abs.lat	poly(abs.lat, 2)2	0.72	0.49	1.5	0.146	187.8	0.04
local.diversity	cur.biomes	cur.13.500	0.05	0.09	0.6	0.561	190.7	0.00
local.diversity	cur.biomes	cur.13.1000	0.02	0.09	0.3	0.789	190.9	0.00
local.diversity	cur.biomes	cur.13.2000	0.02	0.09	0.3	0.784	190.9	0.00
local.diversity	cur.biomes	cur.12.500	0.03	0.09	0.3	0.771	190.9	0.00
local.diversity	cur.biomes	cur.12.1000	0.02	0.09	0.2	0.822	190.9	0.00
local.diversity	cur.biomes	cur.12.2000	0.01	0.09	0.1	0.890	191.0	0.00
local.diversity	cur.biomes	cur.2.500	0.11	0.09	1.3	0.212	189.4	0.01
local.diversity	cur.biomes	cur.2.1000	0.12	0.09	1.3	0.191	189.3	0.01
local.diversity	cur.biomes	cur.2.2000	0.10	0.09	1.2	0.240	189.6	0.01
local.diversity	cur.biomes	cur.14.500	0.10	0.09	1.1	0.257	189.7	0.01
local.diversity	cur.biomes	cur.14.1000	0.07	0.09	0.8	0.409	190.3	0.01
local.diversity	cur.biomes	cur.14.2000	0.05	0.09	0.6	0.581	190.7	0.00
local.diversity	cur.biomes	cur.7.500	0.20	0.09	2.3	0.026	186.0	0.04
local.diversity	cur.biomes	cur.7.1000	0.20	0.09	2.3	0.021	185.5	0.04
local.diversity	cur.biomes	cur.7.2000	0.23	0.09	2.7	0.008	183.8	0.05

Study variable	Driver type	predictors	Coef	SE	t value	Р	AICc	R ²
local.diversity	cur.biomes	cur.1.500	0.18	0.09	2.1	0.041	186.7	0.03
local.diversity	cur.biomes	cur.1.1000	0.19	0.09	2.2	0.028	186.1	0.04
local.diversity	cur.biomes	cur.1.2000	0.20	0.09	2.3	0.024	185.8	0.04
local.diversity	cur.biomes	cur.10.500	0.25	0.09	2.9	0.004	182.5	0.06
local.diversity	cur.biomes	cur.10.1000	0.26	0.09	3.0	0.003	182.0	0.07
local.diversity	cur.biomes	cur.10.2000	0.23	0.09	2.6	0.010	184.3	0.05
local.diversity	cur.biomes	cur.9.500	0.13	0.09	1.5	0.131	188.7	0.02
local.diversity	cur.biomes	cur.9.1000	0.17	0.09	1.9	0.058	187.3	0.03
local.diversity	cur.biomes	cur.9.2000	0.20	0.09	2.3	0.022	185.7	0.04
local.diversity	Igm.biomes	lgm.13.500	0.21	0.09	2.4	0.017	185.2	0.04
local.diversity	lgm.biomes	lgm.13.1000	0.24	0.09	2.8	0.006	183.3	0.06
local.diversity	Igm.biomes	lgm.13.2000	0.25	0.09	2.9	0.005	183.0	0.06
local.diversity	lgm.biomes	lgm.1.500	0.11	0.09	1.2	0.217	189.4	0.01
local.diversity	Igm.biomes	lgm.1.1000	0.09	0.09	1.0	0.301	189.9	0.01
local.diversity	lgm.biomes	lgm.1.2000	0.05	0.09	0.6	0.584	190.7	0.00
local.diversity	lgm.biomes	lgm.7.500	0.23	0.09	2.7	0.008	183.8	0.05
local.diversity	lgm.biomes	lgm.7.1000	0.25	0.09	2.9	0.004	182.5	0.06
local.diversity	lgm.biomes	lgm.7.2000	0.27	0.09	3.1	0.002	181.6	0.07
local.diversity	lgm.biomes	Igm.3.500	0.05	0.09	0.5	0.602	190.7	0.00
local.diversity	lgm.biomes	lgm.3.1000	0.01	0.09	0.1	0.946	191.0	0.00
local.diversity	lgm.biomes	lgm.4.500	0.12	0.09	1.3	0.183	189.2	0.01
local.diversity	lgm.biomes	lgm.4.1000	0.12	0.09	1.3	0.183	189.2	0.01
local.diversity	lgm.biomes	lgm.4.2000	0.12	0.09	1.3	0.183	189.2	0.01
local.diversity	cur.climate	PC1	0.09	0.09	1.1	0.296	189.9	0.01
local.diversity	cur.climate	poly(PC1, 2)1	0.52	0.50	1.1	0.292	188.8	0.03
local.diversity	cur.climate	poly(PC1, 2)2	0.88	0.50	1.8	0.079	188.8	0.03
local.diversity	cur.climate	PC2	-0.12	0.09	-1.3	0.192	189.3	0.01
local.diversity	cur.climate	poly(PC2, 2)1	-0.65	0.50	-1.3	0.193	191.3	0.01
local.diversity	cur.climate	poly(PC2, 2)2	0.13	0.50	0.3	0.791	191.3	0.01
local.diversity	cur.climate	PC3	0.02	0.09	0.3	0.794	190.9	0.00
local.diversity	cur.climate	poly(PC3, 2)1	0.13	0.50	0.3	0.794	192.5	0.01
local.diversity	cur.climate	poly(PC3, 2)2	0.38	0.50	0.8	0.454	192.5	0.01
local.diversity	cur.climate	PC4	-0.20	0.09	-2.3	0.025	185.9	0.04
local.diversity	cur.climate	poly(PC4, 2)1	-1.12	0.49	-2.3	0.025	186.6	0.05
local.diversity	cur.climate	poly(PC4, 2)2	-0.58	0.49	-1.2	0.236	186.6	0.05
local.diversity	lgm.climate	PC1	0.14	0.09	1.6	0.115	188.5	0.02
local.diversity	lgm.climate	poly(PC1, 2)1	0.79	0.50	1.6	0.116	190.1	0.02
local.diversity	lgm.climate	poly(PC1, 2)2	0.35	0.50	0.7	0.486	190.1	0.02
local.diversity	lgm.climate	PC2	-0.12	0.09	-1.4	0.163	189.0	0.02
local.diversity	lgm.climate	poly(PC2, 2)1	-0.70	0.50	-1.4	0.164	190.5	0.02
local.diversity	lgm.climate	poly(PC2, 2)2	0.38	0.50	0.8	0.444	190.5	0.02
local.diversity	lgm.climate	PC3	0.07	0.09	0.8	0.404	190.3	0.01
local.diversity	lgm.climate	poly(PC3, 2)1	0.42	0.50	0.8	0.404	191.6	0.01
local.diversity	lgm.climate	poly(PC3, 2)2	0.45	0.50	0.9	0.370	191.6	0.01
local.diversity	lgm.climate	PC4	0.08	0.09	0.9	0.386	190.2	0.01
local.diversity	lgm.climate	poly(PC4, 2)1	0.44	0.50	0.9	0.385	190.9	0.02
local.diversity	lgm.climate	poly(PC4, 2)2	-0.60	0.50	-1.2	0.229	190.9	0.02
local.diversity	wild	wild.5	0.25	0.09	3.0	0.004	182.5	0.06
local.diversity	wild	poly(wild.5, 2)1	1.43	0.49	2.9	0.004	184.6	0.06
local.diversity	wild	poly(wild.5, 2)2	-0.08	0.49	-0.2	0.866	184.6	0.06
local.diversity	wild	wild.10	0.28	0.09	3.2	0.002	180.8	0.08
local.diversity	wild	poly(wild.10, 2)1	1.56	0.48	3.2	0.002	181.9	0.08
local.diversity	wild	poly(wild.10, 2)2	-0.50	0.48	-1.0	0.306	181.9	0.08

Study variable	Driver type	predictors	Coef	SE	t value	Р	AICc	R ²
local.diversity	wild	wild.20	0.25	0.09	2.9	0.004	182.5	0.06
local.diversity	wild	poly(wild.20, 2)1	1.43	0.49	2.9	0.004	184.4	0.07
local.diversity	wild	poly(wild.20, 2)2	-0.22	0.49	-0.4	0.660	184.4	0.07
local.diversity	veg.type	veg.type = grassl.	0.13	0.09	1.5	0.137	188.7	0.02
dark.diversity	abs.lat	abs.lat	-0.27	0.09	-3.2	0.002	181.0	0.08
dark.diversity	abs.lat	poly(abs.lat, 2)1	-1.54	0.48	-3.2	0.002	182.8	0.08
dark.diversity	abs.lat	poly(abs.lat, 2)2	0.28	0.48	0.6	0.568	182.8	0.08
dark.diversity	cur.biomes	cur.2.500	0.18	0.09	2.0	0.045	186.9	0.03
dark.diversity	cur.biomes	cur.2.1000	0.12	0.09	1.4	0.165	189.0	0.02
dark.diversity	cur.biomes	cur.2.2000	0.03	0.09	0.3	0.745	190.9	0.00
dark.diversity	cur.biomes	cur.14.500	0.23	0.09	2.6	0.010	184.3	0.05
dark.diversity	cur.biomes	cur.14.1000	0.23	0.09	2.7	0.009	184.0	0.05
dark.diversity	cur.biomes	cur.14.2000	0.20	0.09	2.3	0.023	185.7	0.04
dark.diversity	cur.biomes	cur.7.500	0.14	0.09	1.6	0.110	188.4	0.02
dark.diversity	cur.biomes	cur.7.1000	0.15	0.09	1.7	0.095	188.2	0.02
dark.diversity	cur.biomes	cur.7.2000	0.11	0.09	1.2	0.225	189.5	0.01
dark.diversity	cur.biomes	cur.1.500	0.19	0.09	2.1	0.034	186.4	0.04
dark.diversity	cur.biomes	cur.1.1000	0.16	0.09	1.8	0.072	187.7	0.03
dark.diversity	cur.biomes	cur.1.2000	0.11	0.09	1.2	0.237	189.6	0.01
dark.diversity	cur.biomes	cur.10.500	0.03	0.09	0.4	0.720	190.9	0.00
dark.diversity	lgm.biomes	lgm.12.500	0.12	0.09	1.4	0.162	189.0	0.02
dark.diversity	Igm.biomes	lgm.12.1000	0.10	0.09	1.2	0.253	189.7	0.01
dark.diversity	lgm.biomes	lgm.12.2000	0.05	0.09	0.5	0.600	190.7	0.00
dark.diversity	lgm.biomes	lgm.2.500	0.21	0.09	2.4	0.019	185.4	0.04
dark.diversity	Igm.biomes	lgm.2.1000	0.24	0.09	2.8	0.007	183.4	0.06
dark.diversity	lgm.biomes	lgm.2.2000	0.25	0.09	2.9	0.004	182.6	0.06
dark.diversity	lgm.biomes	lgm.1.500	0.25	0.09	2.8	0.005	183.1	0.06
dark.diversity	Igm.biomes	lgm.1.1000	0.20	0.09	2.3	0.022	185.7	0.04
dark.diversity	Igm.biomes	lgm.1.2000	0.13	0.09	1.5	0.137	188.7	0.02
dark.diversity	Igm.biomes	lgm.7.500	0.20	0.09	2.3	0.026	185.9	0.04
dark.diversity	lgm.biomes	lgm.7.1000	0.21	0.09	2.4	0.017	185.2	0.04
dark.diversity	Igm.biomes	lgm.7.2000	0.18	0.09	2.0	0.046	186.9	0.03
dark.diversity	Igm.biomes	lgm.3.500	0.22	0.09	2.5	0.015	184.9	0.05
dark.diversity	Igm.biomes	lgm.3.1000	0.19	0.09	2.2	0.033	186.4	0.04
dark.diversity	lgm.biomes	lgm.3.2000	0.17	0.09	2.0	0.050	187.1	0.03
dark.diversity	cur.climate	PC1	0.38	0.08	4.5	< 0.001	171.6	0.14
dark.diversity	cur.climate	poly(PC1, 2)1	2.11	0.47	4.5	<0.001	173.7	0.14
dark.diversity	cur.climate	poly(PC1, 2)2	0.07	0.47	0.1	0.888	173.7	0.14
dark.diversity	cur.climate	PC2	0.20	0.09	2.4	0.020	185.5	0.04
dark.diversity	cur.climate	poly(PC2, 2)1	1.15	0.49	2.3	0.021	187.6	0.04
dark.diversity	cur.climate	poly(PC2, 2)2	-0.13	0.49	-0.3	0.796	187.6	0.04
dark.diversity	cur.climate	PC3	0.14	0.09	1.5	0.125	188.6	0.02
dark.diversity	cur.climate	poly(PC3, 2)1	0.77	0.48	1.6	0.114	181.5	0.09
dark.diversity	cur.climate	poly(PC3, 2)2	-1.47	0.48	-3.1	0.003	181.5	0.09
dark.diversity	cur.climate	PC4	0.10	0.09	1.1	0.265	189.7	0.01
dark.diversity	cur.climate	poly(PC4, 2)1	0.56	0.50	1.1	0.267	191.8	0.01
dark.diversity	cur.climate	poly(PC4, 2)2	-0.11	0.50	-0.2	0.826	191.8	0.01
dark.diversity	lgm.climate	PC1	0.33	0.08	3.9	<0.001	176.5	0.11
dark.diversity	lgm.climate	poly(PC1, 2)1	1.84	0.47	3.9	<0.001	174.5	0.14
dark.diversity	lgm.climate	poly(PC1, 2)2	0.95	0.47	2.0	0.045	174.5	0.14
dark.diversity	lgm.climate	PC2	0.20	0.09	2.3	0.023	185.7	0.04
dark.diversity	lgm.climate	poly(PC2, 2)1	1.13	0.49	2.3	0.023	186.3	0.05
dark.diversity	lgm.climate	poly(PC2, 2)2	-0.60	0.49	-1.2	0.226	186.3	0.05

Study variable	Driver type	predictors	Coef	SE	t value	Р	AICc	R ²
dark.diversity	lgm.climate	PC3	0.02	0.09	0.3	0.783	190.9	0.00
dark.diversity	lgm.climate	poly(PC3, 2)1	0.14	0.48	0.3	0.775	182.0	0.08
dark.diversity	lgm.climate	poly(PC3, 2)2	-1.62	0.48	-3.4	0.001	182.0	0.08
dark.diversity	lgm.climate	PC4	-0.26	0.09	-3.0	0.003	182.3	0.07
dark.diversity	lgm.climate	poly(PC4, 2)1	-1.45	0.49	-3.0	0.004	184.4	0.07
dark.diversity	lgm.climate	poly(PC4, 2)2	0.03	0.49	0.1	0.948	184.4	0.07
dark.diversity	wild	wild.5	-0.07	0.09	-0.8	0.428	190.4	0.00
dark.diversity	wild	poly(wild.5, 2)1	-0.40	0.50	-0.8	0.430	192.4	0.01
dark.diversity	wild	poly(wild.5, 2)2	0.12	0.50	0.2	0.809	192.4	0.01
dark.diversity	wild	wild.10	-0.09	0.09	-1.0	0.325	190.0	0.01
dark.diversity	wild	poly(wild.10, 2)1	-0.49	0.50	-1.0	0.326	191.8	0.01
dark.diversity	wild	poly(wild.10, 2)2	0.27	0.50	0.5	0.595	191.8	0.01
dark.diversity	wild	wild.20	-0.01	0.09	-0.1	0.937	191.0	0.00
dark.diversity	wild	poly(wild.20, 2)1	-0.04	0.50	-0.1	0.937	193.1	0.00
dark.diversity	wild	poly(wild.20, 2)2	-0.12	0.50	-0.2	0.819	193.1	0.00
dark.diversity	veg.type	veg.type = grassl.	-0.22	0.09	-2.6	0.011	184.4	0.05
comm.compl.	abs.lat	abs.lat	-0.03	0.09	-0.4	0.723	190.9	0.00
comm.compl.	abs.lat	poly(abs.lat, 2)1	-0.18	0.50	-0.4	0.723	192.0	0.01
comm.compl.	abs.lat	poly(abs.lat, 2)2	0.49	0.50	1.0	0.328	192.0	0.01
comm.compl.	cur.biomes	cur.8.500	0.00	0.09	0.0	0.992	191.0	0.00
comm.compl.	cur.biomes	cur.8.1000	0.00	0.09	0.0	0.987	191.0	0.00
comm.compl.	cur.biomes	cur.8.2000	0.01	0.09	0.1	0.890	191.0	0.00
comm.compl.	cur.biomes	cur.13.500	0.07	0.09	0.8	0.437	190.4	0.00
comm.compl.	cur.biomes	cur.13.1000	0.05	0.09	0.5	0.599	190.7	0.00
comm.compl.	cur.biomes	cur.13.2000	0.05	0.09	0.5	0.587	190.7	0.00
comm.compl.	cur.biomes	cur.12.500	0.06	0.09	0.6	0.529	190.6	0.00
comm.compl.	cur.biomes	cur.12.1000	0.06	0.09	0.7	0.497	190.5	0.00
comm.compl.	cur.biomes	cur.12.2000	0.05	0.09	0.6	0.553	190.6	0.00
comm.compl.	cur.biomes	cur.2.500	0.03	0.09	0.3	0.745	190.9	0.00
comm.compl.	cur.biomes	cur.2.1000	0.05	0.09	0.6	0.559	190.6	0.00
comm.compl.	cur.biomes	cur.2.2000	0.08	0.09	0.9	0.398	190.3	0.01
comm.compl.	cur.biomes	cur.14.500	0.00	0.09	0.1	0.963	191.0	0.00
comm.compl.	cur.biomes	cur.7.500	0.11	0.09	1.3	0.212	189.4	0.01
comm.compl.	cur.biomes	cur.7.1000	0.12	0.09	1.3	0.195	189.3	0.01
comm.compl.	cur.biomes	cur.7.2000	0.15	0.09	1.8	0.083	187.9	0.02
comm.compl.	cur.biomes	cur.1.500	0.08	0.09	0.9	0.354	190.1	0.01
comm.compl.	cur.biomes	cur.1.1000	0.10	0.09	1.2	0.247	189.6	0.01
comm.compl.	cur.biomes	cur.1.2000	0.13	0.09	1.4	0.154	188.9	0.02
comm.compl.	cur.biomes	cur.10.500	0.20	0.09	2.2	0.027	186.0	0.04
comm.compl.	cur.biomes	cur.10.1000	0.22	0.09	2.5	0.012	184.6	0.05
comm.compl.	cur.biomes	cur.10.2000	0.23	0.09	2.6	0.009	184.1	0.05
comm.compl.	cur.biomes	cur.9.500	0.17	0.09	1.9	0.058	187.3	0.03
comm.compl.	cur.biomes	cur.9.1000	0.19	0.09	2.2	0.030	186.2	0.04
comm.compl.	cur.biomes	cur.9.2000	0.22	0.09	2.5	0.014	184.8	0.05
comm.compl.	lgm.biomes	lgm.11.500	0.02	0.09	0.2	0.821	190.9	0.00
comm.compl.	lgm.biomes	lgm.11.1000	0.04	0.09	0.4	0.688	190.8	0.00
comm.compl.	lgm.biomes	Igm.8.500	0.05	0.09	0.6	0.560	190.7	0.00
comm.compl.	lgm.biomes	lgm.8.1000	0.03	0.09	0.4	0.696	190.8	0.00
comm.compl.	lgm.biomes	Igm.8.2000	0.02	0.09	0.2	0.861	191.0	0.00
comm.compl.	lgm.biomes	lgm.13.500	0.22	0.09	2.5	0.014	184.9	0.05
comm.compl.	lgm.biomes	lgm.13.1000	0.24	0.09	2.8	0.006	183.1	0.06
comm.compl.	lgm.biomes	lgm.13.2000	0.24	0.09	2.8	0.006	183.2	0.06
comm.compl.	lgm.biomes	lgm.1.500	0.00	0.09	0.1	0.960	191.0	0.00

Study variable	Driver type	predictors	Coef	SE	t value	Р	AICc	R ²
comm.compl.	Igm.biomes	lgm.1.1000	0.01	0.09	0.1	0.953	191.0	0.00
comm.compl.	Igm.biomes	lgm.7.500	0.12	0.09	1.4	0.170	189.1	0.01
comm.compl.	Igm.biomes	lgm.7.1000	0.13	0.09	1.5	0.132	188.7	0.02
comm.compl.	lgm.biomes	lgm.7.2000	0.16	0.09	1.8	0.079	187.8	0.02
comm.compl.	Igm.biomes	lgm.4.500	0.12	0.09	1.3	0.187	189.2	0.01
comm.compl.	Igm.biomes	lgm.4.1000	0.12	0.09	1.3	0.187	189.2	0.01
comm.compl.	lgm.biomes	lgm.4.2000	0.12	0.09	1.3	0.188	189.2	0.01
comm.compl.	cur.climate	PC1	-0.05	0.09	-0.6	0.544	190.6	0.00
comm.compl.	cur.climate	poly(PC1, 2)1	-0.30	0.50	-0.6	0.543	190.8	0.02
comm.compl.	cur.climate	poly(PC1, 2)2	0.69	0.50	1.4	0.167	190.8	0.02
comm.compl.	cur.climate	PC2	-0.17	0.09	-1.9	0.061	187.4	0.03
comm.compl.	cur.climate	poly(PC2, 2)1	-0.93	0.50	-1.9	0.062	189.5	0.03
comm.compl.	cur.climate	poly(PC2, 2)2	0.15	0.50	0.3	0.759	189.5	0.03
comm.compl.	cur.climate	PC3	-0.03	0.09	-0.3	0.752	190.9	0.00
comm.compl.	cur.climate	poly(PC3, 2)1	-0.16	0.50	-0.3	0.751	190.3	0.02
comm.compl.	cur.climate	poly(PC3, 2)2	0.82	0.50	1.6	0.103	190.3	0.02
comm.compl.	cur.climate	PC4	-0.20	0.09	-2.2	0.027	186.0	0.04
comm.compl.	cur.climate	poly(PC4, 2)1	-1.10	0.49	-2.2	0.027	187.3	0.04
comm.compl.	cur.climate	poly(PC4, 2)2	-0.44	0.49	-0.9	0.375	187.3	0.04
comm.compl.	lgm.climate	PC1	0.00	0.09	0.0	0.994	191.0	0.00
comm.compl.	lgm.climate	poly(PC1, 2)1	0.00	0.50	0.0	0.994	193.1	0.00
comm.compl.	lgm.climate	poly(PC1, 2)2	-0.05	0.50	-0.1	0.928	193.1	0.00
comm.compl.	lgm.climate	PC2	-0.17	0.09	-2.0	0.054	187.2	0.03
comm.compl.	lgm.climate	poly(PC2, 2)1	-0.96	0.49	-2.0	0.054	188.2	0.04
comm.compl.	lgm.climate	poly(PC2, 2)2	0.52	0.49	1.1	0.295	188.2	0.04
comm.compl.	lgm.climate	PC3	0.05	0.09	0.6	0.558	190.6	0.00
comm.compl.	lgm.climate	poly(PC3, 2)1	0.29	0.50	0.6	0.554	189.2	0.03
comm.compl.	lgm.climate	poly(PC3, 2)2	0.93	0.50	1.9	0.063	189.2	0.03
comm.compl.	lgm.climate	PC4	0.15	0.09	1.7	0.087	188.0	0.02
comm.compl.	lgm.climate	poly(PC4, 2)1	0.86	0.50	1.7	0.087	189.1	0.03
comm.compl.	lgm.climate	poly(PC4, 2)2	-0.50	0.50	-1.0	0.312	189.1	0.03
comm.compl.	wild	wild.5	0.23	0.09	2.7	0.009	183.9	0.05
comm.compl.	wild	poly(wild.5, 2)1	1.31	0.49	2.7	0.009	186.0	0.05
comm.compl.	wild	poly(wild.5, 2)2	-0.11	0.49	-0.2	0.823	186.0	0.05
comm.compl.	wild	wild.10	0.26	0.09	3.0	0.004	182.3	0.07
comm.compl.	wild	poly(wild.10, 2)1	1.44	0.49	3.0	0.004	183.4	0.07
comm.compl.	wild	poly(wild.10, 2)2	-0.50	0.49	-1.0	0.307	183.4	0.07
comm.compl.	wild	wild.20	0.21	0.09	2.4	0.018	185.3	0.04
comm.compl.	wild	poly(wild.20, 2)1	1.18	0.49	2.4	0.018	187.3	0.04
comm.compl.	wild	poly(wild.20, 2)2	-0.14	0.49	-0.3	0.784	187.3	0.04
comm.compl.	veg.type	veg.type = grassl.	0.19	0.09	2.1	0.036	186.5	0.03

Fig. S2. Uncertainty maps for predictions of AM fungal species pool size, local and dark diversity. Global predictions were made using random 80% subsets of the full data. This was repeated 100 times and uncertainty was calculated as the standard deviation of estimates derived from the different iterations.

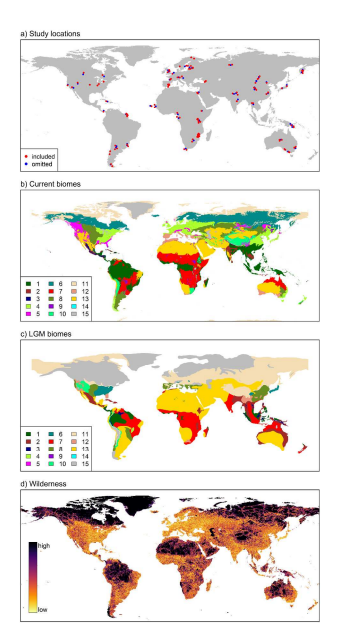


Fig 1 a, b, c, d

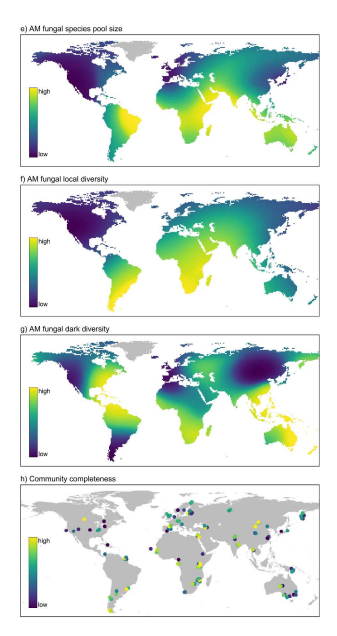
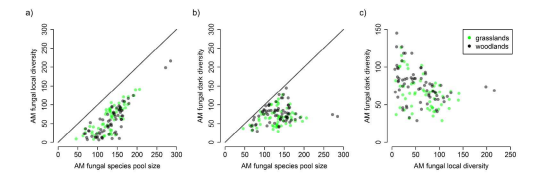


Fig 1 e, f, g, h



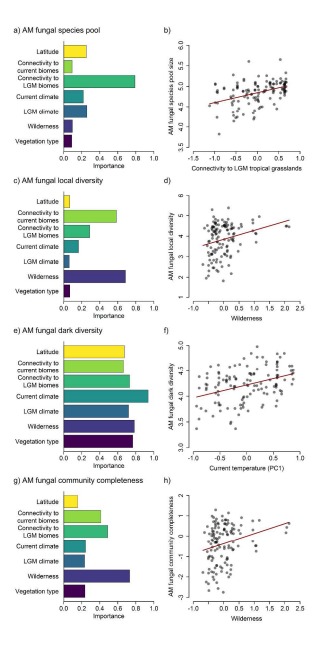


Fig 3

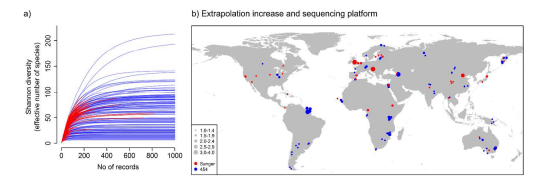


Fig. S1

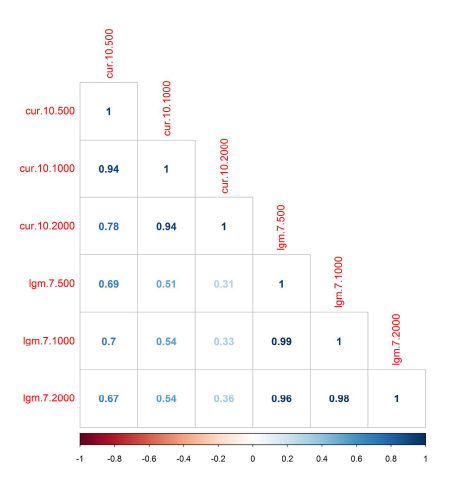
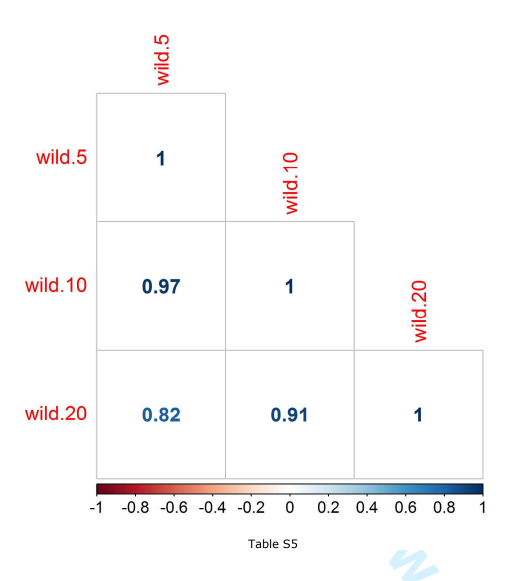


Table S4





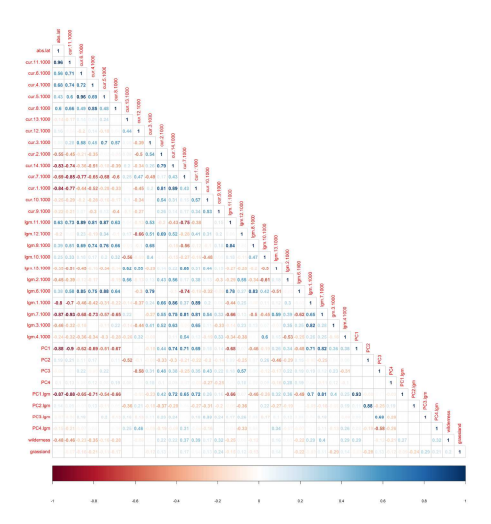


Table S6



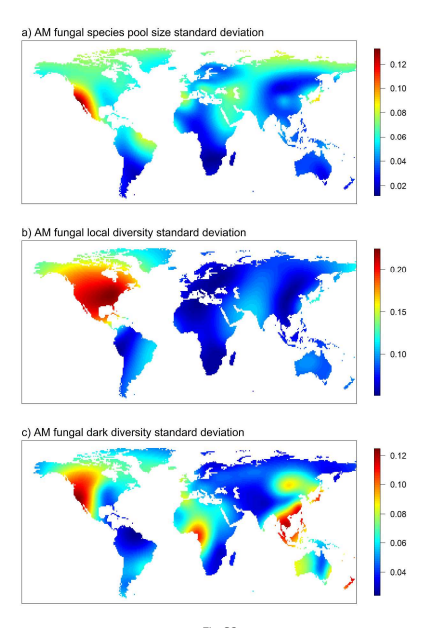


Fig S2