Relationship between soil fungal diversity and temperature in the maritime Antarctic

Published online as a letter by MacMillan Publishers in *Nature Climate Change* on 28th September 2015

DOI: 10.1038/NCLIMATE2806

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Soil fungi have pivotal ecological roles as decomposers, pathogens and symbionts^{1,2}. Alterations to their diversity arising from climate change could thus have substantial effects on ecosystems, particularly those undergoing rapid warming that contain few species^{3,4}. Here, we report a study using pyrosequencing to assess fungal diversity in 29 soils sampled from along a 1,650 km climatic gradient through the maritime Antarctic, the most rapidly-warming region in the southern hemisphere^{5,6}. Using a 'space-for-time' substitution approach, we show that soil fungal diversity is higher in warmer habitats, with increases of 4.7 (observed) and 11.3 (predicted) fungal taxa per degree Celsius rise in surface air temperature along the transect. Among 22 measured predictor variables, air temperature was the most consistent and strongest predictor of diversity. We propose that the current rapid warming in the maritime Antarctic (0.34 °C per decade⁶) will facilitate the colonization of soil by a wider diversity of fungi than at present, with data from regression models suggesting 20–27% increases in soil fungal species richness in the southernmost soils by 2100. Such increases in diversity, which provide a sentinel for changes at lower latitudes, are likely to have substantial impacts on nutrient cycling and, ultimately, net productivity in the species-poor soils of maritime Antarctica.

The maritime Antarctic is undergoing rapid climate change. Surface air temperatures in the region, which broadly encompasses the Antarctic Peninsula and islands of the Scotia Arc, have risen by up to 2.8 °C over the last 50 years, at rates several times that of the global mean^{5,6}. Rising temperatures in the region have led to changes to the physical environment, including ice shelf collapses and glacier retreats⁵. However, in recent years, biological responses to warming have also become apparent

across the region⁷. These include order of magnitude increases in the population sizes of the two native angiosperms, increased moss growth rates, and the establishment of non-native plant species^{8–10}. The range expansions of native plant populations and the establishment of non-native species in the typically unvegetated soils of the region are thought to be associated with new areas of land becoming exposed following glacial retreat, enhanced plant growth and reproduction, and accelerated soil nutrient cycling^{7,10,11}.

Although climate change effects on the maritime Antarctic flora have recently become apparent, far less is known of soil microbial responses to warming in the region. Artificial warming experiments in the natural environment have shown relatively minor changes to the composition of bacterial communities in response to increased soil temperatures (0.5-2 °C annual means), which is not surprising as the experiments have only lasted for one to three years^{12,13}. However, the responses of soil fungi to climate warming in the maritime Antarctic have yet to receive attention. Despite their pivotal importance in terrestrial ecosystems as decomposers, pathogens and symbionts^{1,2}, the majority of fungi are filamentous in form and grow slowly in the natural environment¹⁴, hampering assessments of their responses to warming treatments. For instance, in the low Arctic, substantial changes to root symbiotic fungal communities in response to warming only become apparent after 17 years of treatment¹⁵. Here, in order to circumvent the problem of detecting the responses of these slow-growing microbes to warming manipulations, we studied fungi in soil sampled from along a natural climatic gradient through the maritime Antarctic. Using a similar approach to previous 'spacefor-time' substitution studies^{16,17}, we employ the gradient as a proxy to predict changes to soil fungal communities arising from climate warming in the region. We show that surface air temperature is a significant factor shaping the diversity and composition of soil fungal communities. On the basis of our observations, we predict that future warming in the region will lead to 20-27% increases in the numbers of fungal species present in the southernmost soils of the region by the end of the century, and that this will have consequent effects on biological productivity.

We studied 29 soils sampled during the 2007–2008 austral spring and summer from along a 1,650 km gradient between 72°S and 60°S (Fig. 1; Supplementary Table S1). The soils were free of vegetation (Supplementary Fig. S1), and were hence representative of the barren soils that are frequent in maritime Antarctic terrestrial ecosystems. Data from the Regional Atmospheric Climate Model¹⁸ indicated a significant increase in mean annual surface air temperature (MASAT) between south-eastern Alexander Island (72 °S; MASAT -11 °C) and Signy Island in the South Orkney Islands (60°S; MASAT -4 °C), with a 0.62 °C increase in air temperature for each degree decrease in latitude (Fig. 1, *upper inset*; Supplementary Table S2). To determine whether other abiotic parameters altered along the transect, we analysed soils for a suite of 20 physicochemical parameters (including pH, electrical conductivity and the concentrations of 11 elements and five dissolved ions). Soil C:N ratio declined significantly at lower latitudes (R^2 =33%, P=0.001; Fig. 1, *lower inset*; Supplementary Table S2) and was negatively associated with MASAT (R^2 =31%, P=0.002). This is consistent with previous observations that soils in cold ecosystems have higher C:N ratios than those in warmer ecosystems¹⁹, most likely because of slow organic matter decay¹. The strong influence of latitude on MASAT, and its weaker effect on soil C:N ratio, were also confirmed by structural equation modelling (Supplementary

Fig. S2). None of the other parameters that we measured, including the altitude from which the samples were taken, altered significantly with latitude (Supplementary Table S2).

We measured the numbers of fungal species present in soil along the climatic gradient by PCR amplifying and pyrosequencing internal transcribed spacer 2 regions of fungal ribosomal RNA operons. Each fungal taxon was represented by an operational taxonomic unit (OTU), which differed by >3% in sequence identity from other OTUs. In contrast with studies on other continents, where woody plant species are present and ectomycorrhiza-forming basidiomycetes are consequently frequent in soil², we found relatively few basidiomycetes, with ascomycetes dominating the soil fungal communities (Supplementary Fig. S3). When stepwise multiple regression analyses were used to assess how the numbers of observed and predicted (Chao 1) OTUs responded to the 21 predictors along the gradient, MASAT was consistently the best predictor for both measures of soil fungal diversity (Table 1), explaining 23% and 39% of variation in the numbers of observed and predicted OTUs, respectively. Univariate regression analyses indicated significant positive associations between MASAT and the numbers of fungal OTUs, with mean increases of 4.7 observed OTUs and 11.3 predicted OTUs per degree Celsius rise in air temperature along the transect (Fig. 2a, b). Soil potassium (K) concentration, which is known to be positively associated with the frequencies of fungi²⁰, was also included in stepwise regression models, and explained 12–14% of variation in the numbers of OTUs (Table 1). C:N ratio, also previously shown to influence soil fungal abundances^{2,20}, explained a small (4.3%) but significant amount of variation in the predicted number of OTUs (Table 1).

Redundancy analysis (RDA) was then used to determine how the frequencies of the fungal OTUs varied with the 22 predictors across the gradient. These analyses indicated that changes to soil fungal community composition were best predicted by MASAT (R^2 =6.1%, P<0.001), with soil pH, manganese (Mn) concentration and C:N ratio predicting significant, but smaller, amounts of variation in composition (R^2 =4.8%–3.8%, all P<0.03). MASAT was the best predictor for three OTUs closely matching mycobionts of the lichen genus Verrucaria or the family Verrucariaceae (Supplementary Fig. S3), which each increased in frequency in warmer soils (Fig. 3a-c). Verrucaria is typically a rockinhabiting genus in the Antarctic²¹ and it is hence likely that we detected the fungus in its free-living state (or as spores or other propagules) in soil. RDA based on presence-absence data indicated that changes to the frequencies of Verrucaria largely accounted for the increase in fungal diversity in warmer soils, corroborating the previous observation that the numbers of maritime Antarctic lichen taxa increase sharply between 72°S and 60°S²². The yeast *Rhodotorula* and a member of the Helotiales, representatives of which are known to be present in Antarctica (Supplementary Table S3)²³, also accounted for increases in fungal diversity in warmer soils (Fig. 3d, e). In contrast, we found that the frequencies of three other OTUs were best predicted by MASAT (Supplementary Fig. S3), but that these fungi, which matched closely with a Verrucaria, an unclassified ascomycete and Cladosporium, decreased in frequency in warmer soils (Fig. 3f-h). As C:N ratio was associated with air temperature (Fig. 1, lower inset), and there was a priori evidence indicating that this parameter is associated with climate^{1,19}, we also included the ratio in the RDA analyses. Soil C:N ratio was found to be positively associated with the abundances of OTUs matching closely with the lichen-forming

ascomycete *Polysporina* and the free-living ascomycetes *Cladosporium*, *Pseudoeurotium* and *Penicillium* (Fig. 4a–d; Supplementary Fig. S3), confirming the direct influence of this parameter on soil fungal community composition^{2,20}. Associations between OTU frequencies and soil pH, K and Mn concentrations, which influenced soil fungal diversity but did not alter along the gradient, are summarised in Supplementary Fig. S3.

Our observations show that surface air temperature is an important factor shaping the diversity and composition of maritime Antarctic soil fungal communities. Because liquid water availability is tightly coupled with air temperature in maritime Antarctic environments²⁴, it is likely that part of the observed pattern of increasing fungal diversity in warmer habitats is owing to improved access to water, which, in combination with higher temperatures, will enhance metabolic activity, extend the period for which fungi are active each year, and enable a switch from survival to growth and dispersal strategies. Our observations thus extend into the Antarctic the recent finding that climate has an important influence on soil fungal diversity on the Earth's other continents². They suggest that declines in soil C:N ratio arising from climate warming in the maritime Antarctic might result in reductions in the frequencies of several ascomycetes in soil, including the widespread saprotrophs Penicillium and Cladosporium, which are known to be of importance in soil decomposition processes¹. However, the overriding impact of warming is likely to be an increase in the number of fungal taxa present in the soils of the region. The data here indicate that increases are likely in the frequencies of mycobionts of the lichen-forming genus Verrucaria, which are known to influence rock weathering processes, the yeast genus Rhodotorula, and the order Helotiales, some members of which positively affect the in vitro growth of the angiosperms spreading throughout the region²⁵.

The rate at which fungal diversity will increase in maritime Antarctic soils will be influenced not only by rising air temperatures but also by precipitation, which is likely to increase as the region warms⁶, and by vectors such as air currents and human activities, including tourism⁹. However, assuming a current rate of warming over land in the maritime Antarctic of 0.34 °C per decade⁶, and increases of 4.7 observed species of soil fungi per degree Celsius rise in air temperature, our analyses (Table 1) suggest up to a 20% increase in the numbers of fungal species present in the southernmost soils of the region by 2100. The estimates for the predicted numbers of fungal species are similar, with multiple regression models - taking into account both increases in air temperature and decreases in soil C:N ratio (Table 1) - indicating a 27% rise in the numbers of species of soil fungi as decomposers, symbionts and pathogens^{1,2,20,26} and the demonstrable positive effects of increasing soil fungal diversity on productivity in species-poor soils^{3,4}, our data suggest substantial impacts of such increases on the ecology of maritime Antarctic terrestrial ecosystems, and on other low diversity soil fungal communities at lower latitudes.

Acknowledgements

Logistical support was provided by the British Antarctic Survey and the Royal Navy (HMS *Endurance*). Vito Armando Laudicina, Victoria Ord, Phil Coates, Mike Dunn, Paul Torode, Matt

Jobson, Adam Clark, James Wake, Dickie Hall, Gareth Marshall, Magda Biszczuk and Kate Bazeley provided technical support. This work was funded by a UK Natural Environment Research Council Antarctic Funding Initiative grant (NE/D00893X/1; AFI 7/05) led by D.W.H. and a University of Queensland Early Career Researcher Award to P.G.D. All are gratefully acknowledged.

Author contributions

K.K.N., P.G.D., S.P.R., A.G.O. and D.W.H. conceived this study. P.G.D., K.K.N. and D.W.H. collected soil samples, L.C.C. performed DNA extractions and PCRs. P.G.D. processed sequence data and P.G.D. and K.K.N. performed statistical analyses. P.T.F. provided geospatial data derived from the Regional Atmospheric Climate Model. All authors discussed the results and contributed to the preparation of the manuscript.

Accession codes

The amplicon sequences associated with this study have been deposited in the NCBI SRA under accession PRJNA282894.

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Table 1 Data from stepwise multiple regression models using observed and predicted (Chao
1) numbers of OTUs as response variables.

response variable	R ² (%)	predictor variables	slope	<i>F</i> value	P value
observed no. OTUs	35	MASAT* soil K concentration	4.7 4.4 × 10 ⁻³	9.15 4.83	0.006 0.037
predicted no. OTUs	57	MASAT* soil K concentration soil C:N ratio	14.8 9.2 × 10 ⁻³ 4.8	24.55 8.12 4.46	<0.001 0.009 0.045

*mean annual surface air temperature. Error degrees of freedom in all analyses were 26. MASAT and K concentration were expressed in degrees Celsius and mg kg⁻¹ in these analyses, respectively, and C:N ratio was dimensionless. Intercepts in multiple regression models for observed and predicted numbers of OTUs were 112 and 199, respectively.



Figure 1 | Locations of sampling sites along the climatic gradient. Site names, latitudes, longitudes and altitudes are shown in Supplementary Table S1. *Upper inset* shows mean annual surface air temperature at each sampling location for 2007, derived from the Regional Atmospheric Climate Model¹⁸, as a function of latitude. *Lower inset* shows soil C:N ratio as a function of latitude.



Figure 2 | The influence of mean annual surface air temperature on the numbers of soil fungal taxa along the climatic gradient. Data shown are the numbers of (a) observed and (b) predicted fungal taxa recorded in the 29 soils as a function of mean annual surface air temperature. Taxa were grouped at 97% similarity. The numbers of observed and predicted operational taxonomic units in each soil are shown in Supplementary Table S1.



Figure 3 | The influence of mean annual surface air temperature on the frequencies of eight fungal taxa. Lines of best fit are either linear or polynomial functions. Note that *x*-axes are all identically scaled.



Figure 4 | The influence of soil C:N ratio on the frequencies of four fungal taxa. Lines of best fit are either linear or polynomial functions. Note that *x*-axes are all identically scaled.

Methods

Soil sampling. To eliminate any effects of the presence of plants on soil fungal diversity, we sampled only soils without plant cover from along the climatic gradient. The uppermost five centimetres of soil was collected in 50 ml DNA/RNAase-treated plastic tubes (30 mm diam.) from each of five locations at each site and was bulked. The soil was then immediately snap-frozen by immersion in a mixture of dry ice and ethanol (*c*. -80 °C). Samples were maintained at -80 °C from the time of sampling until they were processed.

Air temperature data. The inaccessible nature of most of the sites studied precluded the measurement of soil temperature, and so mean annual surface air temperature (MASAT) data, derived from the Regional Atmospheric Climate Model over Antarctica¹⁸, gridded at a horizontal resolution of 55×55 km, were used as predictors of soil fungal diversity. MASAT values for the year 2007 in the grids in which each of the sampling sites occurred were used in statistical analyses (see below).

DNA extraction, PCR amplification and 454 pyrosequencing. Total DNA was extracted under sterile conditions from 10 g of soil using a PowerMax[®] Soil DNA isolation kit (MO BIO Laboratories, Inc., Carlsbad, CA, USA) as per the manufacturer's instructions. The internal transcribed spacer 2 (ITS2) region of the ribosomal RNA encoding genes was amplified by polymerase chain reaction (PCR) using the primers gITS7 (5' GTGARTCATCGARTCTTTG²⁷) and ITS4 (5' TCCTCCGCTTATTGATATGC²⁸), which target sites in the 5.8S gene and ribosomal large subunit, respectively. The gITS7 primer was 5'-labelled with the 454 FLX sequencing primer adapter B sequence and the ITS4 primer was 5'-labelled with a sample specific barcode sequence and the 454 FLX sequencing primer adapter A sequence. PCRs were performed in duplicate 50 µl reactions, each containing 5 ng template DNA, 1X Phusion[®] High Fidelity PCR Buffer (New England Biolabs Inc.), 0.2 mM of each of the dNTPs (Invitrogen), 0.3 µM of the ITS4 primer, 0.5 µM of the gITS7 primer, and 1U of 1X Phusion[®] High Fidelity DNA Polymerase (New England Biolabs Inc.). Thermocycling conditions were as follows: 98 °C for 30 s, 35 cycles of 98 °C for 10 s, 56 °C for 30 s, 72 °C for 15 s and a final extension at 72 °C for 7 min. Negative controls, consisting of sterile water in place of template DNA, did not yield amplicons. Amplicons were purified using a Wizard® SV Gel and PCR Clean-Up System (Promega), quantified with a Qubit fluorometer with a Quant-iT dsDNA HF assay kit and then 72 ng of each sample was pooled. The pooled sample was purified again using a QIAquick PCR Purification Kit (Qiagen), and then sent to Macrogen (Seoul, Korea) for 454 pyrosequencing²⁹.

Processing of sequence data. Sequences were quality filtered and dereplicated using the QIIME script split_libraries.py with the homopolymer filter deactivated³⁰. Homopolymer errors were corrected using Acacia v. 1.48³¹ and fungal ITS2 sequences were then extracted using ITSx v. 1.0.9³² and checked for chimeras against ITS2 sequences in UNITE v. 6³³ using UCHIME v. 3.0.617³⁴. At least 1,435 non-chimeric quality-filtered ITS2 sequences were derived from each soil sample. The sequences were clustered at 97% similarity using UCLUST v. 1.2.22. UNITE v. 6³⁵ taxonomy was

assigned to representative OTU sequences using BLAST v. 2.2.30. Tables containing the abundances of different OTUs and their taxonomic assignments in each sample were generated and the number of reads was rarefied to 1400 per sample. The mean number of observed OTUs and the estimated total number of OTUs (Chao 1) were calculated using QIIME.

Soil physicochemistry. Soil pH and electrical conductivity were measured in 1:2.5 and 1:5 soil:water (vol:vol) slurries, respectively. Total nitrogen and organic carbon concentrations were determined using an Exeter Analytical CE440 Elemental Analyzer (EAI, Coventry, UK) following desiccation at 105 °C and treatment with HCl to remove inorganic carbon. Concentrations of Ca, Cu, Fe, K, Mg, Mn, Ni, P and Zn (mg kg⁻¹ dry soil) were determined using ICP-MS following reverse aqua regia digests. Water extractable $PO_4^{3^-}$, $SO_4^{2^-}$ and Cl⁻ concentrations (mg kg⁻¹ dry soil) were determined using Dionex ion chromatography. Soil NH₄⁺-N, NO₃⁻-N/NO₂⁻-N and dissolved organic carbon (DOC) concentrations (mg kg⁻¹ dry soil) were determined in 1M KCl (1:4 dry soil equivalent to solution ratio) followed by automated flow injection analysis (Skalar Analytical B.V., The Netherlands). The moisture content of each soil sample was determined gravimetrically. All analyses were based on soil passed through a 4 mm sieve.

Statistical analyses. The influence of MASAT and soil physicochemical parameters on changes in the composition of soil fungal communities between sites (beta diversity) was assessed using Redundancy Analysis (RDA) with Monte Carlo permutation tests to assess the significance of the constraints. Parameters were chosen for inclusion in the final RDA model by forward selection based on Akaike's Information Criterion (AIC). When the inclusion of a new parameter led to other terms becoming insignificant (P > 0.05), it was dropped, leaving the final minimally adequate model. All models were built for Hellinger transformed OTU abundances. All analyses were implemented in R. Associations between soil fungal diversity, latitude, MASAT and soil physicochemical parameters were examined using polynomial and multivariate regressions. For the univariate models we adopted a principle of parsimony, accepting the model with the least number of coefficients commensurate with the greatest amount of variation (R^2 value) explained. Since some of the covariates were themselves related and causally linked (e.g. latitude and MASAT), we hypothesised that each could have both direct effects on the response variables and indirect effects through their direct effects on other predictors. For this reason we used structural equation modelling (SEM) to investigate the direct and indirect effects of latitude, MASAT and soil physicochemical parameters on soil fungal richness. Based on the output from our univariate models, we created a full SEM model to test the hypotheses that latitude directly affected MASAT, soil C:N ratio and fungal richness, that MASAT directly affected fungal richness, that soil K concentration directly affected fungal richness and that MASAT directly affected soil C:N ratio. The hypothetical full model was then challenged with the data and goodness of fit assessed using Chi-squared tests and root mean square error of approximation (RMSEA) values. Variables and associated relationships were then removed progressively if they were not significant. The Chi-squared goodness of fit tested the difference between the hypothetical model pathway and

the observed data. RMSEA values of < 0.05 for the model were considered a good fit to the data. SEM analyses were performed in Amos version 22 (SPSS Software, Chicago).

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Supplementary Information

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				No. C)TUs
Sampling site		Latitude and longitude	Altitude (metres a.s.l.)	Observed*	Predicted (Chao1)*
1	Mars Oasis, Alexander Island	71.878° S, 68.248° W	15	72.85	110.41
2	Mars Oasis, Alexander Island	71.878° S, 68.248° W	15	65.55	118.10
3	Mars Oasis, Alexander Island	71.878° S, 68.248° W	15	49.85	78.02
4	Mars Oasis, Alexander Island	71.878° S, 68.248° W	15	78.40	113.36
5	Mount Holt, Alexander Island	69.408° S, 71.665° W	70	85.05	121.22
6	Hopkins Ridge, Alexander Island	69.366° S, 71.842° W	62	106.80	136.48
7	Hopkins Ridge, Alexander Island	69.366° S, 71.844° W	62	58.75	139.73
8	Hopkins Ridge, Alexander Island	69.367° S, 71.844° W	66	75.65	186.62
9	Hopkins Ridge, Alexander Island	69.368° S, 71.844° W	501	100.75	149.46
10	Jenny Island	67.731° S, 68.365° W	12	100.05	188.23
11	Lagoon Island	67.594° S, 68.247° W	20	106.00	201.55
12	Rothera Point, Adelaide island	67.568° S, 68.114° W	6	43.95	84.36
13	Blaiklock Island	67.543° S, 67.198° W	7	64.50	101.72
14	Detaille Island	66.790° S, 66.869° W	21	96.30	148.67
15	Cape Evenson, Antarctic Peninsula	66.145° S, 65.717° W	59	81.75	122.21
16	Yelcho Station (CHL), Wiencke Island	64.894° S, 63.553° W	6	94.70	170.42
17	Port Lockroy, Goudier Island	64.817° S, 63.483° W	10	76.25	125.57
18	Marambio Station (ARG), Seymour Island	64.236° S, 56.626° W	160	71.55	101.65
19	Alectoria Island	63.977° S, 58.640° W	50	104.50	171.40
20	Alectoria Island	63.977° S, 58.640° W	25	90.50	137.69
21	Spert Island	63.844° S, 60.945° W	92	131.60	245.89
22	Cape Lachman, James Ross Island	63.782° S, 57.783° W	81	158.20	261.64
23	nunatak, unnamed	63.559° S, 57.823° W	617	96.50	140.20
24	Whalers Bay, Deception Island	62.977° S, 60.552° W	91	114.55	201.86
25	South Beaches, Byers Peninsula, Livingston Island	62.655° S, 61.090° W	64	85.15	172.41

Supplementary Table S1 | Site locations and alpha diversity metrics.

26	Edwards Point, Robert Island	62.460° S, 59.509° W	26	147.80	276.18
27	Orión Point, Greenwich Island	62.447° S, 59.736° W	13	113.55	223.13
28	Keller Peninsula, King George	62.086° S, 58.399° W	65	85.15	140.61
	Island				
29	Wynn Knolls, Signy Island	60.701° S, 45.661° W	199	107.00	160.90

*Data are based on sequences clustered at 97% similarity and rarefied to 1,400 reads per soil. Abbreviations: CHL, Chile; ARG, Argentina

Parameter	R ² value (%)	P value
MASAT* (° C)	62.30	<0.001***
Altitude (metres a.s.l.)	2.5	0.413
pH value	0.01	0.966
Electrical conductivity (µS cm ⁻¹)	1.42	0.539
Organic C concentration (mg kg ⁻¹ dry soil)	0.64	0.679
N concentration (mg kg ⁻¹ dry soil)	0.29	0.780
C:N ratio	32.60	0.001**
Ca concentration (mg kg ⁻¹ dry soil)	0.40	0.744
Cu concentration (mg kg ⁻¹ dry soil)	3.14	0.358
Fe concentration (mg kg ⁻¹ dry soil)	6.17	0.194
K concentration (mg kg ⁻¹ dry soil)	0.16	0.835
Mg concentration (mg kg ⁻¹ dry soil)	10.49	0.099
Mn concentration (mg kg ⁻¹ dry soil)	2.97	0.371
Ni concentration (mg kg ⁻¹ dry soil)	7.20	0.176
P concentration (mg kg ⁻¹ dry soil)	0.31	0.775
Zn concentration (mg kg ⁻¹ dry soil)	1.39	0.543
Dissolved organic C concentration (mg kg ⁻¹ dry soil)	0.73	0.658
Dissolved NO_3^N/NO_2^N concentration (mg kg ⁻¹ dry soil)	0.01	0.953
Dissolved NH_4^+ -N concentration (mg kg ⁻¹ dry soil)	0.10	0.869
Dissolved SO ₄ ²⁻ concentration (mg kg ⁻¹ dry soil)	1.37	0.545
Dissolved PO ₄ ³⁻ concentration (mg kg ⁻¹ dry soil)	0.25	0.796
Dissolved Cl ⁻ concentration (mg kg ⁻¹ dry soil)	1.22	0.569

Supplementary Table S2 | R^2 and P values from univariate linear regressions between latitude and mean annual surface air temperature, altitude and soil physicochemical parameters.

*mean annual surface air temperature. Error degrees of freedom in analyses were 26–28. Note that four outlying points were removed from the data matrix (consisting of 667 observations) prior to analyses.

		Closest UNITE	Closest BLAST hit	
OTU	Taxonomic assignment	species hypothesis	NCBI accession no.	Geographical origin
1	Megasporaceae sp.	SH038307.06FU	FR682345	Finland
2	Davidiellaceae; Cladosporium sp.	SH196750.06FU	KJ598781	China
3	Davidiellaceae; <i>Cladosporium</i> sp.	SH196750.06FU	KJ411576	India
4	Pseudoeurotiaceae; Pseudoeurotium sp.	SH209178.06FU	HM589327	Antarctic Peninsula
5	Pseudoeurotiaceae; Pseudogymnoascus sp.	SH236509.06FU	KC457091	King George Island, Antarctica
6	Eurotiomycetes sp.	SH030591.06FU	HQ634132	Antarctica
7	Eurotiomycetes sp.	SH030588.06FU	KC965283	Canadian Arctic
8	Herpotrichiellaceae; Capronia sp.	SH209735.06FU	FR682430	Finland
9	Herpotrichiellaceae sp.	SH228285.06FU	KF636404	Léonie Island, Antarctica
10	Trichocomaceae; Penicillium sp.	SH231333.06FU	KF597019	Himalayas
11	Verrucariaceae sp.	SH022709.06FU	KC966268	Alaskan Arctic
12	Verrucariaceae sp.	SH024484.06FU	KF297206	Canadian Arctic
13	Verrucariaceae sp.	SH221269.06FU	KC965575	Canadian Arctic
14	Verrucariaceae; Verrucaria sp.	SH022729.06FU	FJ667941	United Kingdom
15	Verrucariaceae; Verrucaria sp.	SH022737.06FU	KF296796	Canadian Arctic
16	Verrucariaceae; Verrucaria sp.	SH197325.06FU	FJ667941	United Kingdom
17	Verrucariaceae; Verrucaria sp.	SH022766.06FU	KC965347	Canadian Arctic
18	Verrucariaceae; Verrucaria sp.	SH221271.06FU	FJ664852	United Kingdom
19	Verrucariaceae; Verrucaria sp.	SH022730.06FU	FJ664870	United Kingdom
20	Verrucariaceae; Verrucaria sp.	SH022730.06FU	FJ664872	Iceland
21	Acarosporaceae; Acarospora sp.	SH228160.06FU	EU870668	United Kingdom
22	Acarosporaceae; Polysporina sp.	SH032519.06FU	JX036093	Dry Valleys, Antarctica
23	Ramalinaceae; <i>Waynea</i> sp.	SH026019.06FU	AF282099	USA

Supplementary Table S3 | Closest BLAST hits of the top 50 operational taxonomic units* and their associated UNITE species hypotheses.

24	Lecideaceae; <i>Lecidea</i> sp.	SH224611.06FU	JX036045	Dry Valleys, Antarctica
25	Physciaceae; <i>Buellia</i> sp.	SH028904.06FU	HQ650628	USA
26	Leotiomycetes sp.	SH018537.06FU	GU067676	Austria
27	Helotiales sp.	SH216429.06FU	KC785555	Mount Erebus, Antarctica
28	Helotiales sp.	SH220881.06FU	JX001631	Tibetan Plateau
29	Helotiales sp.	SH207339.06FU	JX852366	Ardley Island, Antarctica
30	Helotiales sp.	SH197325.06FU	EU292671	Alaskan Arctic
31	Helotiales sp.	SH220883.06FU	X88771	Ötztal Alps, Austria
32	Leotiaceae; Alatospora sp.	SH207345.06FU	KF617795	Alaskan Arctic
33	Pseudeurotiaceae sp.	SH209227.06FU	KC455895	Mount Blanc, Italian Alps
34	Pseudeurotiaceae; Leuconeurospora sp.	SH209227.06FU	JQ857041	King George Island, Antarctica
35	Thelebolaceae; Antarctomyces sp.	SH222305.06FU	KC514843	Deception Island, Antarctica
36	Thelebolaceae; <i>Thelebolus</i> sp.	SH195163.06FU	KC457177	King George Island, Antarctica
37	Pezizomycetes sp.	SH194645.06FU	EF635708	Austrian Alps
38	Nectriaceae; Fusarium sp.	SH217297.06FU	KJ655442	China
39	Ascomycota sp.	SH021267.06FU	KC965667	Alaskan Arctic
40	Ascomycota sp.	SH021267.06FU	KC965667	Alaskan Arctic
41	Ascomycota sp.	m.d.	KJ523042	Hawaii
42	Kriegeriaceae; Rhodotorula sp.	SH216408.06FU	JQ857032	King George Island, Antarctica
43	Kriegeriaceae; <i>Rhodotorula</i> sp.	SH228918.06FU	KC206490	Lake Vostok, Antarctica
44	Basidiomycota sp.	SH028275.06FU	KC966210	Alaskan Arctic
45	Unidentified	SH241742.06FU	EF635742	Austrian Alps
46	Unidentified	SH228811.06FU	KC966059	Canadian Arctic
47	Unidentified	SH023732.06FU	KC966243	Canadian Arctic
48	Mortierellaceae; Mortierella sp.	SH217216.06FU	JQ670951	King George Island, Antarctica
49	Mortierellaceae; Mortierella sp.	SH221068.06FU	HM589301	Antarctic Peninsula

50	Mortierellaceae; Mortierella sp.	SH212358.06FU	JQ670948	King George Island, Antarctica
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m.d.; missing data

*OTU numbers correspond to those shown in Supplementary Figure S3.

Supplementary Figures



Supplementary Figure S1 | Images of eight of the sampling sites. The numbers correspond to the site numbers in Fig. 1 and Supplementary Table S1.



Supplementary Figure S2 | Structural equation model showing the influences of latitude, mean annual surface air temperature, soil C:N ratio and soil potassium concentration on the richness of maritime Antarctic soil fungal communities.



Supplementary Figure S3 | Heatmap showing the frequencies of the 50 most abundant taxa of soil fungi along the climatic gradient. R^2 values shown in black and red indicate positive and negative associations, respectively. UNITE species hypothesis accessions for each taxon are shown in Supplementary Table S3.