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# Soil-foraging animals alter the composition and co-occurrence of microbial communities in a desert shrubland

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Animals that modify their physical environment by foraging in the soil can have dramatic effects on ecosystem functions and processes. We compared bacterial and fungal communities in the foraging pits created by bilbies and burrowing bettongs with undisturbed surface soils dominated by biocrusts. Bacterial communities were characterized by Actinobacteria and Alphaproteobacteria, and fungal communities by Lecanoromycetes and Archaeosporomycetes. The composition of bacterial or fungal communities was not observed to vary between loamy or sandy soils. There were no differences in richness of either bacterial or fungal operational taxonomic units (OTUs) in the soil of young or old foraging pits, or undisturbed soils. Although the bacterial assemblage did not vary among the three microsites, the composition of fungi in undisturbed soils was significantly different from that in old or young foraging pits. Network analysis indicated that a greater number of correlations between bacterial OTUs occurred in undisturbed soils and old pits, whereas a greater number of correlations between fungal OTUs occurred in undisturbed soils. Our study suggests that digging by soil-disturbing animals is likely to create successional shifts in soil microbial and fungal communities, leading to functional shifts associated with the decomposition of organic matter and the fixation of nitrogen. Given the primacy of organic matter decomposition in arid and semi-arid environments, the loss of native soil-foraging animals is likely to impair the ability of these systems to maintain key ecosystem processes such as the mineralization of nitrogen and the breakdown of organic matter, and to recover from disturbance.

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## Introduction

**ORIGINAL ARTICLE** 

Australia has suffered one of the highest rates of global mammal extinctions over the past 200 years since European settlement (Woinarski *et al.*, 2012). Losses have been most pronounced in the critical weight range (35-5500 g) mammals, which were once common over large areas of continental Australia (Johnson, 2006). The loss of these animals, or the contraction of their ranges, has been attributed to multiple causes associated with European settlement and pastoral practices such as altered fire regimes, overgrazing by livestock, competition

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with exotic pests including the European rabbit (Oryctolagus cuniculus) and predation by introduced species such as the domestic cat (*Felis catus*) and the red fox (Vulpes vulpes) (Johnson, 2006). Two species that have suffered substantial range restrictions are the greater bilby (*Macrotis lagotis*) and the burrowing bettong (Bettongia lesueur). Recent attempts have been made to reintroduce these animals into predator-proof exclosures within their former range in an effort to re-establish viable populations (James and Eldridge, 2007).

Many of Australia's locally extinct animals forage extensively in the soil for seeds, bulbs, invertebrates and fungi (Robley et al., 2001; James et al., 2011; Eldridge et al., 2012). Foraging disturbs the soil surface and breaks up the surface crust (biocrust), altering rates of water infiltration, and creating small pits and depressions that trap water, soil, organic matter and seed (James et al., 2009). These pits develop into patches of higher nutrients, with greater concentrations of plant-available nitrogen and carbon than the surrounding soil matrix (James *et al.*, 2010) and often a different vegetation community (Lavelle *et al.*, 2006). Studies worldwide have shown that modification of the abiotic environment by these animals, a process referred to as ecosystem engineering (*sensu* Jones *et al.*, 1994), alters energy flows and resource availability, increases species richness, diversity and productivity, through niche construction, ultimately controlling the availability and distribution of resources to other organisms (for example, Whitford and Kay, 1999; Jones *et al.*, 2010).

An important process moderated by soildisturbing animals in arid environments is the decomposition of organic matter. Litter and organic matter in these systems are spatially and temporally variable, and is often concentrated within the foraging pits of animals (James and Eldridge, 2007). Litter is a source of carbon, nitrogen and other trace elements, and provides habitat for a range of microand macro-invertebrates involved in the decomposition of organic matter (Haslem et al., 2011). Litter falling into pits comes into close contact with soil, where it is held in situ more effectively than if it remained on the soil surface where it is subject to removal by wind and water (Whitford, 2002). Together with reduced evaporation resulting from lower temperatures in the pits than the undisturbed surface (Eldridge and Mensinga, 2007), this increases the time period over which soil moisture is optimum for microbial decomposition and nutrient mineralization (Steinberger and Whitford, 1983; Jacobsen and Jacobsen, 1998; Whitford, 2002). Litter remaining on the surface, however, is subject to photodegradation (Austin and Vivanco, 2006), potentially reducing the return of carbon to the soil organic pool.

Soil-disturbing animals therefore play an important role in bringing surface-resident organic matter into contact with soil microorganisms. The accumulation of litter in the pits is also likely to exert a strong selective pressure on microorganisms essential for the decomposition process. Given the marked differences in the biotic (litter cover and composition) and abiotic (for example, surface temperature, soil moisture) environments between pits and undisturbed soils, that is, those soils undisturbed by animal activity (for example, Vossbrinck et al., 1979; Wallwork et al., 1985; Eldridge and Mensinga, 2007), we expected that the pits would differ in the composition of soil microorganisms. For example, studies of foraging disturbances constructed by the short-beaked echidna (Tachyglossus aculeatus) indicate a greater diversity and abundance of microarthropods and higher rates of microbial respiration in the pits than undisturbed soil (Eldridge and Mensinga, 2007), suggesting that there are differences in the abundance or structure of microbial communities. Over time, pits collect organic matter, and research has shown that pits over about 12 months old have high levels of organic carbon. This compares with young pits (<3 months old), which have relatively low levels of litter and organic matter (D J Eldridge, unpublished data). We would expect pit age to influence microbial community structure, as these old pits (~12 months) would have more time to establish seedlings and accumulate litter and microorganisms that are present on adjacent, undisturbed surfaces. Furthermore, older pits could provide a greater range of different environments, with differences in depth, shape and orientation, and therefore different soil chemistry and organic matter at varying stages of decomposition.

We compared the community structure of soil microbial communities in old and young pits with the undisturbed surface soil on two soil types in an area where bilbies and bettongs have been reintroduced into their former range. Both bilbies and bettongs construct pits while foraging for buried seed, invertebrates and plant roots. The pits of these two species are indistinguishable, and range from cylindrical-shaped excavations about 15-cm wide and up to 20-cm deep to shallow basin-like structures (Eldridge *et al.*, 2012). Pits are constructed only once, and unlike cache pits of heteromyid rodents (Geluso, 2005), are rarely reworked. Because pits vary in depth and shape, and are constructed in soils of different texture, they provide a range of different physical environments that influences the trapping and retention of litter and the breakdown of organic matter.

We hypothesized that the microbial community in pit soils would support more microorganisms commonly associated with decomposing litter. Conversely, we expected that the microbial community composition in undisturbed soils would support a community dominated by cyanobacteria, given the extensive cover of biocrusts on the soil surface. We used microbial network analysis to examine the structure of microbial communities, particularly in relation to resilience and reactivity (Ruiz-Moreno et al., 2006; Bissett et al., 2013). Examination of microbial networks improves our understanding of why undisturbed soils might be resistant to nutrient amendment, how microbial community structure is altered following pit construction, and how digging promotes nutrient enrichment within these microsites (James et al., 2009).

## Materials and methods

#### The study area

Our study was undertaken within the Australian Wildlife Conservancy's Scotia Sanctuary in southwestern, New South Wales, Australia (33°43'S, 143°02'E) where locally extinct bilbies and bettongs have been released into predator-proof exclosures. Soil samples were collected from two systems; (i) mallee (*Eucalyptus* spp.) west-east-trending dunes of Quaternary alluvium characterized by calcareous and siliceous sands (Rudosols) and (ii) the inter-dunal swales and plains extending to these dunes, which are up to 500-m wide, comprised mainly of loamy, calcareous soils (Calcarosols). The vegetation on the dunes is moderately dense mallee (Eucalyptus socialis, E. dumosa) and the plains are dominated by open mallee woodland with scattered belah (*Casuarina pauper*) and sugarwood (Myoporum platycarpum), and a variable cover of shrubs such as punty bush (Senna artemisioides), hopbush (Dodonaea viscosa), turpentine (Eremophila sturtii), pinbush wattle (Acacia burkittii) and assorted bluebushes (Maireana spp.), depending on whether trees had been removed. Shrubs covered about 50% of the area of the plains. The climate in the area is semi-arid, with cool winters (mean  $\leq 17$  °C) and hot summers (mean 30 °C). Rainfall is highly spatially and temporally variable and averages 243 mm per year. Rainfall is evenly distributed between the six warmer months and the six cooler months.

#### Field sampling

The location, size, depth and age of all foraging pits constructed by bilbies and bettongs have been monitored at 36 large sites at the Scotia Sanctuary since 2007. Because sites were visited every 3 months, we were able to calculate the relative age of particular pits. In October 2009, we collected soil samples from six sites: three on sandy dunes and three on loamy plains. At each of the six sites, we sampled three microsites: (i) young foraging pits, that is, pits constructed since the previous measurements (<3 months old), (ii) old foraging pits, that is, pits older than 12 months and (iii) undisturbed non-pit surface soils at least 3 m from any pit. At each of the six sites, we sampled each microsite at 10 locations. For the young pits, soil was removed from the uppermost 10 mm layer of the soil surface or from the base of the pits after removing any existing organic material. Biocrust was not removed from the soil prior to sampling. Approximately 5 g of soil was collected with a sterilized spatula. The material from the 10 locations was then bulked and stored on ice before being transported back to the laboratory. The same procedure was used to collect samples from old pits and undisturbed surfaces. This resulted in a database of 18 bulked samples (3 replicate sites of 2 soil types  $\times$  3 microsites).

#### Molecular analysis

Environmental DNA was isolated from 500 mg of soil using the FASTDNA Spin Kit for Soil (MP Bio Laboratories, Inc., Carlsbad, CA, USA) according to the manufacturer's instructions and stored at -80 °C until use. DNA was quantified using a NanoDrop ND-1000 Spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA) and the quality checked by PCR amplification of the 16 S rRNA gene using the primer pair 27 f/519r (Weisburg, 1991). Bacterial and fungal specific tag-encoded FLX amplicon pyrosequencing of each sample was carried out using the primers 27 f/519r and funSSUF/funSSUR respectively (Lucero, 2011) on a Roche GS-FLX Titanium at the Research and Testing Laboratory (Lubbock, TX, USA). Sequence reads were analyzed using MOTHUR v1.22 (www.mothur.org) software package (Schloss et al., 2009). Initial quality processing of 454 sequence reads was performed using the mothur implementation of PyroNoise (Quince et al., 2011) using default settings. Sequences containing <200 bp, containing ambiguous bases and homopolymers longer than 8 bp in length were removed. The remaining sequences were aligned to either the bacterial or fungal alignments of the SILVA release 102 reference alignment. Chimeric sequences were identified and removed using the mothur implementation of uchime (Edgar *et al.*, 2011). The taxonomic identity of each unique sequence was determined by comparison against the SILVA release 102 reference database. Taxonomic assignment was made at each level, given a bootstrap value greater than 80, using the RDP classifier (Wang et al., 2007). Sequences that failed to be classified at the phylum level or were classified as either Mitochondria, Archaea or Eukaryota/Prokaryota in the respective datasets, were removed. Sub-sampling was performed at a level of 400 sequences per sample for the bacterial dataset and 1300 sequences per sample for the fungal dataset. Implementation of this process resulted in the exclusion of a bacterial young loam soil sample and bacterial young sand soil sample, as these samples contained fewer than the 400 sequences required. To ensure a balanced design across the bacterial dataset, the corresponding samples were subsequently excluded from the bacterial old pit soil and bacterial surface pit soil sets (2 replicate sites of 2 soil types  $\times$  3 microsites. Uncorrected pairwise distances were calculated between sequence reads with the final clustering of operational taxonomic units (OTUs) performed at a 0.03 distance threshold using the average neighbor algorithm (Schloss and Westcott, 2011). The identity of each OTU defined at 0.03 distance threshold was obtained from the consensus of each sequence within that OTU at a confidence threshold of 80. From these data, two individual data matrices were generated, one for bacteria and one for fungi, each matrix containing every OTU and the number of reads assigned to it from each sample. In this instance, the relative proportion of each OTU was used as a proxy for abundance, as absolute abundance measures were not obtained.

#### Statistical analysis

We used permutation multivariate analysis of variance (PERMANOVA; Anderson *et al.*, 2008) to examine differences in the composition of a data matrix of 2500 bacterial OTUs, defined at 0.03

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distance threshold, and a data matrix of 5895 fungal OTUs, defined at 0.03 distance threshold, in relation to microsite (undisturbed soils, young pits, old pits) and soil type (loam, sand). Relative abundance data were used after resampling, to ensure an equivalent number of sequences. The first stratum of this analysis considered soil type and the second stratum microsite and its interaction with soil type. Pairwise a posteriori comparisons were made, where necessary, using a multivariate analogue of the *t* statistic, the probability levels being obtained by permutation. We tested for differences in richness and diversity of taxa with a mixed-model ANOVA with the same structure as the PERMANOVA analysis. Richness and diversity data were checked for homogeneity of variance (Levene's test) and normality using diagnostic tests but no transformations were needed. For all analyses, significant differences between means were examined using Fisher's Protected Least Significant Difference test. The procedure was repeated for the fungal data.

The degree of association of OTUs with respect to microsite was measured with Indicator Species Analysis in R (De Cáceres *et al.*, 2012) using a data matrix consisting of 2500 bacterial OTUs and 5895 fungal OTUs. Indicator values combine information on relative abundance and frequency of species, and the indicator value is maximal (IV = 100%) when all individuals of a given species are restricted to a particular microsite (for example, old pit), and all samples from the particular microsite contain an occurrence of that species. Data (at the OTU level) were randomized among the treatments and a Monte Carlo randomization procedure performed with 1000 iterations to determine the statistical significance of the indicator values.

The degree of association of OTUs with respect to one another within each microsite was measured using the Pearson's correlation coefficient (r). Bacterial and fungal OTU tables, defined at 0.03 distance threshold, were separated on the basis of microsite, and then reduced by removing any OTUs that did not occur across at least 75% of available samples. A Pearson's r score and P-value were calculated pairwise for each bacterial OTU using the rcor.test algorithm, available from the ltm package (available from https://cran.r-project.org/ package=ltm) as implemented in R version 3.0.2. For each correlation, P-values were generated and the false discovery rate was maintained below 5% using the Benjamini-Hochberg procedure (Benjamini and Hochberg, 1995). Visualization of these interactions, incorporating taxonomic, abundance and microsite occurrence information, was made with the freely available Cytoscape package version 2.8.3 (available at: www.cytoscape.org). For each network, topological metrics of connectivity and density were calculated using the network analysis plug-in (Assenov et al., 2008). Networks pre-embedded with sample and OTU-specific information are provided in the Supplementary Material.

## **Results**

### Richness of bacterial and fungal taxa

Most bacterial and fungal OTUs occurred at very low abundances, with a substantial number of abundances equal to one. Of the original 2500 bacterial OTUs after resampling, 320 (14%) contributed 50% of total OTU abundances. For fungal OTUs, 525 (9%) of the 5895 OTUs contributed 50% of total fungal abundances. There were no differences in bacterial OTU richness (that is, different number of OTUs) among the different soils (P=0.24; range 238–332 OTUs) or among the three microsites (P=0.47). Similarly, fungal richness did not vary with soil texture (P=0.81; range 397–873) or among the three microsites (P=0.17).

Community composition of bacterial and fungal taxa Bacterial communities were observed to contain a high proportion of Actinobacteria, and to a lesser extent, Alphaproteobacteria and Acidobacteria (Figure 1a). Cyanobacteria appeared to constitute a large proportion of the bacterial community, particularly in undisturbed soils. Fungal communities were observed to contain a high proportion of Lecanoromycetes, and to a lesser extent, Archaeosporomycetes (Figure 1b).

There was no significant difference in the composition of either bacterial or fungal OTUs assemblages between loamy and clay soils (P > 0.30). The composition of the bacterial assemblage did not vary among the three microsites (P=0.21; Figure 2a), but there was a significant effect for fungi (Pseudo  $F_{2,8}=3.08$ , P(Perm)=0.003). The composition of fungi in undisturbed soils was significantly different from that in old (pairwise t=2.14, P=0.029) or young (t=2.02, P=0.02) pits, but there was no significant differences between old and young pits (P=0.47; Figure 2b).

#### Microsite indicators

Six cvanobacterial OTUs (Gp I (3 OTUs), Gp X, Gp VII and an unclassified OTU) were indicators of undisturbed pits, as were the single Asanoa OTU (Actinobacteria), a Segetibacter OTU (Sphingobacteria) and an unclassified alphaproteobacterial OTU. A single Hylangium OTU (Myxobacteria, Deltaproteobacteria), Microvirga OTU (Rhizobiales, Alphaproteobacteria) and a Gp IV actinobacterial OTU were indicators of old pits. A single Rubrobacter OTU (Actinobacteria), Ammoniphilus OTU (Bacilli, Firmicutes) and Actinaurispora OTU (Actinobacteria) were indicators of young pits (Table 1). Overall, fungal taxa were better discriminators of the three microsites, with 20 orders containing 170 OTUs, with indicator values >0.70, and almost exclusively from sub-phylum Pezizomycotina. These included orders Dothideales (genera Columnosphaeria, Delphinella), Chaetothyriales (genus Glyphium), Lecanorales (genera Sphaerophoraceae, Cladoniaceae),

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a	Acidobacteria	0	0	0	0	0	0
	Actinobacteria	$\bigcirc$	$\bigcirc$	$\bigcirc$	()	$\bigcirc$	$\bigcirc$
	Chloroflexi	0	0	)。	)。	•	0
	Cyanobacteria	$\bigcirc$	0	•	٠	0	0
	Firmicutes	٥	0	0	0	۰	•
e	iemmatimonadetes	۰	0	۰	۰	۰	•
	Sphingobacteria	0	0	۰	۰	0	0
A	Iphaproteobacteria	$\bigcirc$	$\bigcirc$	$\circ$	$\circ$	$\bigcirc$	$\bigcirc$
	Betaproteobacteria	•	0	0	•	0	0
0	Deltaproteobacteria	0	0	0	•	•	0
0	ther Proteobacteria	۰	•	0	۰	۰	٠
	Others	0	0	0	0	0	0
b	Agaricomycetes	٠	•	۰	۰	0	•
Arc	chaeosporomycetes	$\bigcirc$	$\circ$	0	$\circ$	0	$\circ$
	Dothideomycetes	0	0	0	0	0	0
	Eurotiomycetes	0	0	0	0	0	0
	Glomeromycetes	0	0	0	0	0	0
	Lecanoromycetes	$\bigcirc$	$\bigcirc$	$\bigcirc$	$\bigcirc$	$\bigcirc$	$\bigcirc$
	Leotiomycetes	0	0	0	0	0	0
	M-Ascomycota	0	0	$\bigcirc$	0	0	0
	Pezizomycetes	0	0	•	0	0	0
	Pezizomycotina	•	•	•	0	·	•
	Sordariomycetes	0	0	•	•	•	•
	Spirotrichea	۰	0	۰	•	۰	۰
	Others	0	0	0	0	0	0
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		Sar	-oa	Sar	-oa	Sar	oa.
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Figure 1 Relative abundance of major (a) bacterial and (b) fungal taxa within each microsite. Larger circles indicate greater abundance.

Myxotrichaceae (genus Geomyces), Mycocaliciales (Sphinctrina) and Pleosporales (genera Leptosphaeria, Trematosphaeria, Phaeosphaeria). Ten fungal genera (particularly Eupenicillium, Hamigera, Bionectriaceae and an unclassified taxon from the family Bulgaria) were highly indicative (IV > 0.81) of young pits. Old pits contained a wide range of different OTUs, with the orders Chaetothyriales, Dothideales, Hypocreales, Lecanorales, Mycocaliciales and Pleosporales having a large number of OTUs that were strongly indicative (IV > 69%) of older pits (Table 2).

#### Network analysis

Within the bacterial networks, the mean number of correlations between OTUs was greater in old pit soils (3.45) than either undisturbed (2.516) or young pit (1.294) soils, consistent with a larger number of OTUs co-occurring across the samples (Table 3). The majority of the associations present in young pit soils were between a small number of alphaproteobacterial and actinobacterial OTUs. Young pit soils returned the lowest values for network metrics of clustering (0), density (0.081) and centralization (0.050). Undisturbed soils and old pit soils were similar in relation to clustering (undisturbed = 0.566, old pits = 0.547), density (undisturbed = 0.084, old

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Figure 2 Multi-dimensional scaling biplot of the first two dimensions of an ordination of a reduced matrix of (a) 280 bacterial OTUs and (b) 135 fungal OTUs. Note the clustering of undisturbed samples for both bacteria and fungi.

pits = 0.088) and centralization (undisturbed = 0.089, old pits = 0.096) (Table 3).

Within the fungal networks, the highest mean number of correlations between OTUs (20.497) was observed in undisturbed soils, where many more OTUs (1814 OTUs) were present across multiple samples than in young (321) or old (485) pit soils. Similar to the bacterial networks, young pit soils returned the lowest values for density (0.009), but were also the most centralized (0.116). Old pit soils were the least clustered (0.472) and the least centralized (0.067). Undisturbed soils were similar to young pit soils in terms of clustering (undisturbed = 0.652, young pits = 0.647), whereas fungal young and old pit soils were only similar in relation to the mean number of correlations between OTUs.

#### Discussion

Soil foraging by semi-fossorial animals in arid areas disrupts surface crusts, alters rates of water infiltration, and creates small pits and depressions that trap water, soil, organic matter and seed (James and Eldridge, 2007). We expected to detect substantial differences in the soil microbial community between Table 1 Bacterial taxa, to the level of genus, that are significantly associated with different microsites using Indicator Species Analysis

Order	Family	Genus	Microsite	IV	P-value	No of OTUs	
Cyanobacteria	Family I	Group I	Undisturbed	0.866	0.047	3	
Cyanobacteria	Family X	Group X	Undisturbed	0.866	0.046	1	
Cyanobacteria	Unclassified	Unclassified	Undisturbed	0.866	0.049	1	
Actinomycetales	Micromonosporaceae	Asanoa	Undisturbed	0.866	0.046	1	
Cyanobacteria	Family VIII	Group VIII	Undisturbed	0.866	0.047	1	
Sphingobacteriales	Chitinophagaceae	Segetibacter	Undisturbed	0.866	0.049	1	
Alphaproteobacteria	Alphaproteobacteria	Alphaproteobacteria	Undisturbed	0.866	0.047	1	
Myxococcales	Cystobacteraceae	Hyalangium	Old	1.000	0.008	1	
Acidobacteria	Acidobacteria	Group IV	Old	0.913	0.030	1	
Rhizobiales	Methylobacteriaceae	Microvirga	Old	0.812	0.028	1	
Rubrobacterales	Rubrobacteraceae	Rubrobacter	Young	0.905	0.012	1	
Bacillales	Paenibacillaceae	Ammoniphilus	Young	0.866	0.040	1	
Actinomycetales	Micromonosporaceae	Actinaurispora	Young	0.866	0.040	1	

Abbreviations: IV, indicator value; OTUs, operational taxonomic units.

intact undisturbed soils and recently excavated or older, more established pits in response to differences in plant and litter cover, organic matter decomposition and soil nutrient concentrations. Although we detected some significant differences in the fungal community composition between the soil surface and the pits (described below), there were no discernible differences in the bacterial community and in the fungal community between young and old pits, largely because of the high variability among microsites (Figure 1). Consequently, we undertook an analysis that would test whether the physical variability that is observed in pit soils, in regard to moisture and nutrient trapping, influenced the occurrence of individual species or the manner in which individual species exhibited correlations with one another. Indicator species analysis was implemented to identify specific OTUs that were more strongly associated with a particular microsite type. Critically, indicator species analysis has been previously shown to be suitable for identifying variable taxa where there was no prior assessment, or no significant variation, in the larger community composition (De Cáceres and Legendre, 2009; De Cáceres and Legendre, 2009; De Cáceres et al., 2012). That it were possible to identify species that were statistically indicative of particular microsites when the multivariate (PERMANOVA) analysis was insignificant highlights the fact that there is substantial heterogeneity within microsites, and suggests a level of functional redundancy within microbial taxa that prevents large-scale perturbation of the community despite the loss of species. On the basis of the indicator species (De Cáceres and Legendre, 2009; De Cáceres et al., 2012) and microbial network (Chaffron, 2010; Bissett et al., 2013) analyses, there is sufficient evidence to suggest that pits may be associated with a reduction in autotrophic groups (Figure 1, Tables 1 and 2) that are compensated for by an emergence of taxa capable of decomposing organic material (Tables 1 and 2) and reduced resilience in the microbial communities (Table 3).

Compositional differences between pit and undisturbed soils

Consistent with information from arid soils worldwide, the bacterial community contained high proportions of Actinobacteria and Alphaproteobacteria (Figure 1a) (Yeager et al., 2004; Kuske, 2012). At the community level, we detected no significant differences in bacterial community composition between pits and undisturbed soils (Figure 2). However, consistent with our first hypothesis, filamentous diazotrophic (Cyanobacteria GpI), baeocystous (Cyanobacteria GpVIII) and unicellular (Cyanobacteria GpX) cyanobacteria were found to be indicators of undisturbed soils (Table 1) with a reduction in the observed abundance of cvanobacterial sequence reads when soils were disturbed (Figure 1, Table 1). Cyanobacteria were present in undisturbed soils, however, the presence of these taxa as indicators was reflective of both a decrease in the abundance of cyanobacterial groups and a shift within the morphological and physiological nature of cyanobacteria between undisturbed and pit soils. Among the heterotrophic population, actinobacterial members of the Rubrobacteridae that are pioneers in biological crust formation (Yeager et al., 2004) dominated both undisturbed and pit soils, with a single *Rubrobacter* OTU, an indicator of young pit soils. In addition to cyanobacterial groups, the Sphingobacterial genus Segetibacter has been previously affiliated with the decomposition of cyanobacteria- and plant-derived phytodetritus (Li et al., 2011).

Fungal communities in undisturbed and pit soils comprised a wide range of saprotrophs, with *Lecanoromycetes*, the largest class of lichenized fungi, and to a lesser extent, *Archaeosporomycetes*, comprising about 80% of sequences across the three microsites (Figure 1b). Along with *Pezizomycotina*, these fungal taxa perform a diverse array of ecological functions including wood and litter decomposition, mycorrhizal associations and lichen symbioses, animal and plant pathogens (Spatafora *et al.*, 2006). Evidence for active recession, or at least competitive inhibition, of microbial groups from the old pits was Table 2 Fungal taxa, to the level of genus, that are significantly associated with different microsites using Indicator Species Analysis

Eurotianycetidae Eurotiales Trichocomaceae Leptosphaeria Calegosphaeria Unclassified Undisturbed 0.94 0.002 1 Pleosporomycetidae Pleosporales Melanommataceae Melanommataceae Skeletonemataceae Skeletonema Unclassified Unclassified Unclassified Undisturbed 0.91 0.006 4 Skeletonemataceae Navicula Unclassified Unclassified Unclassified Undisturbed 0.91 0.002 1 Dothideomycetidae Pleosporales Phaeosphaeriaceae Pleosporaceae Pleosporaceae Pleosporaceae Pleosporaceae Pleosporaceae Pleosporales Phaeosphaeria Unclassified Undisturbed 0.88 0.013 6 Pleosporomycetidae Pleosporales Phaeosphaeriaceae Plaeosphaeria Undisturbed 0.88 0.011 4 Dothideomycetidae Pleosporales Phaeotrichaceae Pleosporaceae Pleosporaceae Pleosporales Phaeotrichaceae Sphinctrina Undisturbed 0.86 0.012 19 Pleosporomycetidae Pleosporales Phaeotrichaceae Sphinctrina Undisturbed 0.86 0.011 2 Pleosporomycetidae Mycocaliciales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 19 Pleosporatorycetidae Hateridae Mycocaliciales Sphinctrinaceae Geomyces Undisturbed 0.86 0.012 10 Dothideomycetida Pleosporales Phaeotrichaceae Geomyces Undisturbed 0.86 0.012 10 Dothideomycetida Pleosporales Distributed Distributed 0.86 0.012 10 Dothideomycetida Pleosporales Phaeotrichaceae Geomyces Undisturbed 0.88 0.012 10 Dothideomycetida Pleosporales Distributed Distributed 0.86 0.012 10 Dothideomycetida Pleosporales Distributed Distributed 0.86 0.012 10 Dothideomycetida Examorales Distributed Distributed 0.88 0.012 10 Dothideomycetidae Kylariales Xylariaceae Halperia Unclassified Undisturbed 0.82 0.016 1 Leconomycetidae Ecanorales Dothideales	Subclass	Order	Family	Genus	Microsite	IV	Р	No of OTUs
Pleospornivycetidae Pleosporales Leptosphaeria Laptosphaeria Undisturbed 0.91 0.006 8 Skeletonemataceae Skeletonema Unclassified Unclassified Undisturbed 0.91 0.007 1 Dothideomycetidae Dothideales Dothioraceae Delphinella Undisturbed 0.90 0.003 3 Naviculaceae Navicula Unclassified Unclassified Undisturbed 0.90 0.003 4 Pleospormycetidae Pleosporales Phaeosphaeriaceae Pleospora Undisturbed 0.88 0.011 4 Dothideomycetidae Pleosporales Phaeosphaeriaceae Pleosporare Undisturbed 0.88 0.012 17 Chaetothyriomycetidae Pleosporales Phaeosphaeriae Undisturbed 0.86 0.012 17 Chaetothyriomycetidae Chaetothyriales Chaetothyriales Glyphium Undisturbed 0.86 0.012 17 Chaetothyriomycetidae Chaetothyriales Chaetothyriales Sphinctrina Undisturbed 0.86 0.012 19 Detothideomycetidae Chaetothyriales Chaetothyriales Sphinctrina Undisturbed 0.86 0.012 10 Dothideomycetidae Halteria Unclassified Undisturbed 0.86 0.012 10 Dothideomycetidae Halteria Unclassified Undisturbed 0.86 0.012 10 Dothideomycetidae Halteriae Halteria Unclassified Undisturbed 0.86 0.012 10 Dothideomycetidae Halteriae Halteria Unclassified Undisturbed 0.86 0.012 10 Dothideomycetes Myxotrichaceae Letendraea Unclassified Undisturbed 0.86 0.012 10 Dothideomycetes Myxotrichaceae Hyxotrichaceae Hyxotrichaceae Unclassified Undisturbed 0.82 0.013 1 Dothideomycetidae Dothideales Dothideales Hortaea Undisturbed 0.82 0.013 1 Locanoromycetidae Xylariales Arginaceae Hyxotyola Undisturbed 0.82 0.013 1 Dothideomycetidae Xylariales Arginaceae Hyxotyola Undisturbed 0.82 0.013 1 Locanoromycetidae Xylariales Arginaceae Hyxotyola Undisturbed 0.82 0.013 1 Dothideomycetidae Xylariales Arginaceae Pesuaosphaeria Undisturbed 0.82 0.013 1 Xylariomycetidae Xylariales Arginaceae Hyxotyola Undisturbed 0.82 0.013 1 Dothideomycetidae Xylariales Arginaceae Hyxotyola Undisturbed 0.82 0.013 1 Dothideomycetidae Xylariales Arginaceae Hyxotyola Undisturbed 0.82 0.014 1 Dothideomycetidae Kaenorales Lecanoraleae Sphancorhoraceae Undisturbed 0.82 0.015 1 Dothideomycetidae Dothideales Dothicaceae Sphancorhorace	Eurotiomycetidae	Eurotiales	Trichocomaceae	Unclassified	Undisturbed	0.94	0.002	1
Pleosporomycetidae Pleosporales Melanommataceae Trematosphaeria Undisturbed 0.91 0.006 4 Strombidiidae Strombidium Unclassified Unclassified Undisturbed 0.91 0.007 1 Dothideomycetidae Dothideales Dothioraceae Delphinella Undisturbed 0.90 0.003 3 Pleosporomycetidae Pleosporales Pheosphaeriaceae Pheosphaeria Undisturbed 0.88 0.013 4 Pleosporomycetidae Pleosporales Pleosporaceae Pleospora Undisturbed 0.88 0.011 4 Dothideomycetidae Dothideales Dothioraceae Columnosphaeria Undisturbed 0.86 0.012 17 Chaetothyrionycetidae Pleosporales Pheosphaeriaceae Pheosphaerica Undisturbed 0.86 0.012 19 Pleosporomycetidae Pleosporales Pheotyphaeriae Columnosphaeria Undisturbed 0.86 0.012 19 Pleosporomycetidae Pleosporales Phaeotrichaceae Pheotrichum Undisturbed 0.86 0.011 2 Mycocaliciomycetidae Mycocalicales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 19 Pleosporomycetida Mycocalicales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 10 Leotiomycetes Mycorichaceae Letendraea Unclassified Undisturbed 0.86 0.012 10 Lothideomycetes Myxotrichaceae Myxotrichaceae Geomyces Undisturbed 0.86 0.012 10 Lothidomycetidae Halteriidae Halteria Unclassified Undisturbed 0.86 0.012 10 Lothideomycetidae Lecanorales Lecanorineae Cladoniaceae Undisturbed 0.82 0.016 1 Lecanoromycetidae Xylariales Mylariaceae Hypoxylan Undisturbed 0.82 0.016 1 Lecanoromycetidae Xylariales Aylariaceae Hypoxylan Undisturbed 0.82 0.016 1 Lecanoromycetidae Xylariales Mylariaceae Pseudophaeria Undisturbed 0.82 0.016 1 Lecanoromycetidae Xylariales Mylariaceae Pseudophaeria Undisturbed 0.82 0.012 1 Pleibideomycetes Kirschseiniothelia Unclassified Unclassified Undisturbed 0.82 0.012 1 Peizzales Peizzaceae Peizz Unclassified Unclassified Undisturbed 0.82 0.013 1 Lecanoromycetidae Kagangorthaeea Sphaeria Unclassified Undisturbed 0.82 0.013 1 Lecanoromycetidae Magnaporthaeea Sphaeria Unclassified Undisturbed 0.82 0.013 1 Lecanoromycetidae Magnaporthaeea Sphaeriae Unclassified Undisturbed 0.82 0.015 1 Peizzales Peizzaceae Peizza Unclassified Unclas	Pleosporomycetidae	Pleosporales	Leptosphaeriaceae	Leptosphaeria	Undisturbed	0.91	0.006	8
SkeletonemiatoceaeSkeletonemiaUnclassifiedUnclassifiedUnclassifiedUndisturbed0.910.0021DothideomycetidaeDothidealesDothioraceaeDelphinellaUnclassifiedUndisturbed0.900.0033NaviculaceaeNaviculaUnclassifiedUnclassifiedUndisturbed0.880.0134PleosporomycetidaePleosporalesPlaeosphaeriaceaePleosporatesPlaeosphaeriaUndisturbed0.880.0114DothideomycetidaeOchidealesDothioraceaeColumnosphaeriaUndisturbed0.860.01219PleosporaycetidaeChaetothyrialesChaetothyrialesSpractichumUndisturbed0.860.0112ChaetothyrialesChaetothyrialesSpractichumUndisturbed0.860.01210DothideomycetesTubeufiaceaeLetondraeaUnclassifiedUndisturbed0.860.01210DothideomycetesMycocalicialesSpraticrinaceaeBotryosphaeriaUndisturbed0.860.01210DothideomycetesHalteriidaeHalteriaUnclassifiedUndisturbed0.820.0161LocanormycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0121DothideomyceteaXylarialesXylariaceaeHypxylonUndisturbed0.820.0121LocanormycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0121<	Pleosporomycetidae	Pleosporales	Melanommataceae	Trematosphaeria	Undisturbed	0.91	0.006	4
Strombidiidae Strombidium Unclassified Unclassified Unclasturbed 0.91 0.007 1 Dothideomycetidae Dothideales Dothioraceae Delphinella Undisturbed 0.88 0.013 6 Pleosporomycetidae Pleosporales Pleosopraceae Plaeosphaeria Undisturbed 0.88 0.011 4 Dothideomycetidae Dothideales Dothioraceae Columnosphaeria Undisturbed 0.88 0.011 4 Dothideomycetidae Dothideales Dothioraceae Columnosphaeria Undisturbed 0.86 0.012 17 Chaetothyrionycetidae Pleosporales Phaeosphaeriaceae Plaeosporar Undisturbed 0.86 0.012 19 Pleosporomycetidae Pleosporales Phaeotrichaceae Phaeotrichum Undisturbed 0.86 0.011 2 Pleosporomycetidae Mycocaliciales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 11 Dothideomycetes Mycotrichaceae Letendreae Unclassified Undisturbed 0.86 0.012 10 Leotionycetes Mycotrichaceae Hateria Unclassified Undisturbed 0.86 0.012 10 Leotionycetes Myxotrichaceae Hyxotrichaceae Geomyces Undisturbed 0.86 0.012 6 Dothideomycetidae Dothideales Dothideales Hortaea Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Lecanorales Lecanorineae Cladoniaceae Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Xylariaceae Hypoxylon Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Xylariaceae Pestalosphaeria Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Xylariaceae Pestalosphaeria Undisturbed 0.82 0.012 1 Peizzales Peziza Unclassified Undisturbed 0.82 0.012 1 Peizzales Peziza Unclassified Undisturbed 0.82 0.013 1 Dothideomycetidae Xylariales Xylariaceae Pseudoshaeria Undisturbed 0.82 0.016 1 Dothideomycetidae Kagariceae Hypoxylon Undisturbed 0.82 0.016 1 Dothideomycetidae Kagariceae Preziza Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Kagariceae Preziza Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Agaricales Lycoperdaceae Pseudoshaeria Undisturbed 0.82 0.016 1 Dothideomycetidae Agaricales Lycoperdaceae Desphaeria Undisturbed 0.82 0.016 1 Dothideomycetidae Agaricales Trichocomaceae Cheromophearia Undisturbed 0.82 0.025 1 Dothideomycetidae Agaricales Dothioraceae Claunonsphaeria Undi 0.02 0.025 1	Skeletonemataceae	Skeletonema	Unclassified	Unclassified	Undisturbed	0.91	0.002	1
DothideomycetidaeDothidealesDothioraceaeDelphinellaUndisturbed0.900.0033PleosporamycetidaePleosporalesPhaeosphaeriaceaePhaeosphaeriaUndisturbed0.880.0136PleosporavycetidaePleosporalesPhaeosphaeriaceaePhaeosphaeriaUndisturbed0.880.0114DothideomycetidaeOchidealesDothioraceaeColumnosphaeriaUndisturbed0.860.01219PleosporavycetidaeChaetothyrialesChaetothyrialesChaetothyrialesSpharcintriaUndisturbed0.860.0112ChaetothyrionycetidaeChaetothyrialesSphinctrinaceaeSphinctrinaUndisturbed0.860.01210LeotionycetesTubeuflaceaeLetendraeaUnclassifiedUndisturbed0.860.01210LeotionycetesMyxotrichaceaeGeonycesUndisturbed0.860.01210DothideonycetesBotryosphaerialesLecanorineaeCladoniaceaeUndisturbed0.820.0161DothideonycetidaeVylarialesXylariaceaeHypaylonUndisturbed0.820.0121DothideonycetidaeVylarialesXylariaceaeHypaylonUndisturbed0.820.0121ZylarionycetidaeYylarialesAmphisphaeriaceaePseudonectriaUndisturbed0.820.0121DothideonycetidaeMagnaporthaceaeLeganorineaeSphaerophoraceaeUndisturbed0.820.0121	Strombidiidae	Strombidium	Unclassified	Unclassified	Undisturbed	0.91	0.007	1
NaviculaceáeNaviculaUnclassifiedUnclassifiedUnclassifiedUndisturbed0.900.0034PleosporomycetidaePleosporalesPleosporaceaePleosporaUndisturbed0.880.0114DothideomycetidaeDothidealesDothioraceaeColumnosphariaUndisturbed0.860.01217ChaetothyrialesChaetothyrialesChaetothyrialesChyphumUndisturbed0.860.01219PleosporomycetidaePleosporalesPhaeotrichaceaePhaeotrichaceaeUndisturbed0.860.01211DothideomycetesTubeufiaceaeSphinctrinaceaeSphinctrinaUndisturbed0.860.01210LeotionycetesMyxotrichaceaeMyxotrichaceaeGeorycesUndisturbed0.860.01261DothideomycetesBotryosphaerialesBotryosphaeriaceaeBotryosphaeriaUndisturbed0.820.0161DothideomycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0161DothideomycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0161DothideomycetidaeSylarialesXylariaceaeHypoxylonUndisturbed0.820.0121RylariomycetidaeKrschsteiniotheliaUnclassifiedUndisturbed0.820.0121DothideomycetidaeKagaraceaeBulgariaUnclassifiedUndisturbed0.820.0121SordariomycetidaeM	Dothideomycetidae	Dothideales	Dothioraceae	Delphinella	Undisturbed	0.90	0.003	3
Pleosporomycetidae Pleosporales Pleosporace Phacosphaeria Undisturbed 0.88 0.013 6 Pleosporomycetidae Dothideales Dothioraceae Columnosphaeria Undisturbed 0.86 0.012 17 Chaetothyriomycetidae Chaetothyriales Chaetothyriales Glyphium Undisturbed 0.86 0.012 17 Pleosporomycetidae Chaetothyriales Chaetothyriales Chaetothyriomycetidae Chaetothyriales Chaetothyriales Sphinctrinaceae Phacotrichum Undisturbed 0.86 0.012 11 Dothideomycetidae Mycocaliciales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 11 Dothideomycetes Tubeufiaceae Itelendreae Unclassified Undisturbed 0.86 0.012 10 Leatothyriomycetidae Halteria Unclassified Undisturbed 0.86 0.012 10 Leatothyriomycetidae Ecanorineae Ecanorineae Cadoniaceae Undisturbed 0.86 0.012 6 Sporadotrichida Halteriae Botryosphaeriaceae Botryosphaeria Undisturbed 0.85 0.014 7 Dothideomycetidae Lecanorales Lecanorineae Cadoniaceae Undisturbed 0.82 0.016 1 Dothideaes Vylariaceae Hypoxylon Undisturbed 0.82 0.016 1 Dothideaes Dothideales Hortaea Undisturbed 0.82 0.016 1 Dothideaes Sphinceitae Unclassified Undisturbed 0.82 0.016 1 Dothideaes Bulgariaceae Hypoxylon Undisturbed 0.82 0.012 1 Helotiales Bulgariaceae Bulgaria Unclassified Undisturbed 0.82 0.012 1 Dothideomycetidae Xylariales Amphisphaeriaceae Pseudohalonectria Undisturbed 0.82 0.016 1 Sordariomycetidae Kirschsteiniothelia Unclassified Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Kagaronales Lecanoraleae Pseudohalonectria Undisturbed 0.82 0.016 1 Sordariomycetidae Dothideales Dothioraceae Pseudohalonectria Undisturbed 0.82 0.016 1 Dothideomycetidae Chaetothyriales Chaetothyriales Glyphium Old 0.82 0.016 1 Dothideomycetidae Dothideales Dothioraceae Chromocleista Old 0.82 0.016 1 Dothideomycetidae Chaetothyriales Chaetothyriales Glyphium Old 0.82 0.025 1 Chaetothyriomycetidae Kagarcales Lycoperdaceae Lycoperdona Undisturbed 0.81 0.021 3 Sporadotrichida Halteria Halteria Unclassified Old 0.80 0.025 1 Chaetothyriomycetidae Kagarcales Lycoperdaceae Chromocleista Old 0.79 0.030 4 Hypocreaels Mytorichaceae Ecanorales	Naviculaceae	Navicula	Unclassified	Unclassified	Undisturbed	0.90	0.003	4
Pleosporam_vectidae Pleosporales Pleosporacae Pleospora Undisturbed 0.88 0.011 4 Dothideomycetidae Dothideales Dothioraceae Columnosphaeria Undisturbed 0.86 0.012 17 Chaetothyrionycetidae Pleosporales Chaetothyriales Glyphium Undisturbed 0.86 0.012 19 Pleosporomycetidae Pleosporales Chaetothyriales Sarcinomyces Undisturbed 0.86 0.012 10 Dothideomycetes Mycocialiciales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 10 Dothideomycetes Mycotrichaceae Letendraea Unclassified Undisturbed 0.86 0.012 10 Dothideomycetes Mycotrichaceae Halteria Unclassified Undisturbed 0.86 0.012 10 Dothideomycetes Dothideales Dothideales Hortaea Unclassified Undisturbed 0.86 0.012 10 Dothideomycetes Dothideales Dothideales Hortaea Unclassified Undisturbed 0.85 0.012 6 Sporadotrichida Halteriidae Halteria Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Dothideales Dothideales Hortaea Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Dothideales Dothideales Hortaea Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Axylariaceae Hstaleosphaeria Undisturbed 0.82 0.012 10 Nylariomycetidae Xylariales Axylariaceae Pestaleosphaeria Undisturbed 0.82 0.012 1 Helotiales Bulgariaceae Bulgaria Unclassified Undisturbed 0.82 0.012 1 Dothideomycetes Kirschsteiniothelia Unclassified Unclassified Undisturbed 0.82 0.012 1 Dothideomycetidae Agarciaels Dothicaece Pseudohalonceria Undisturbed 0.82 0.016 1 Sordariomycetidae Agarciaels Dothicaceae Lycoperdon Undisturbed 0.82 0.016 1 Dothideomycetidae Agarciaels Lycoperdaceae Lycoperdon Undisturbed 0.82 0.016 1 Dothideomycetidae Dothideales Dothicaceae Delphinella Old 0.82 0.016 1 Dothideomycetidae Dothideales Dothicaceae Chromocleista Old 0.82 0.015 1 Chaetothyriales Chaetothyriales Chaetothyriales Glyphium Old 0.82 0.025 1 Chaetothyriales Chaetothyriales	Pleosporomycetidae	Pleosporales	Phaeosphaeriaceae	Phaeosphaeria	Undisturbed	0.88	0.013	6
DothifeomycetidaeDothiferaceaeColumnosphaeriaUndisturbed0.860.01217ChaetothyriorycetidaeChaetothyrialesChaetothyrialesGlyphiumUndisturbed0.860.0112ChaetothyriorycetidaeChaetothyrialesChaetothyrialesSarcinomycesUndisturbed0.860.01211QueccalicionycetidaeMycocalicialesSphinctrinaceaeSphinctrinaUndisturbed0.860.01211DothideomycetesMyxotrichaceaeMyxotrichaceaeGeomycesUndisturbed0.850.01216SporadotrichidaHalteriaUnclassifiedUndisturbed0.850.0147DothideomycetidaeBotryosphaeriaceaeBotryosphaeriaUndisturbed0.820.0161DothideomycetidaeDothidealesDothidealesBotryosphaeriaceaeHalteriaUndisturbed0.820.0121NylariomycetidaeXylarialesAmphisphaeriaceaeHypoxylonUndisturbed0.820.0121XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaUndisturbed0.820.0121LecanoromycetidaeKirschsteiniotheliaUnclassifiedUndisturbed0.820.0121ValariomycetidaeMagnaporthalesMagnaporthaceaePestalosphaeriaUndisturbed0.820.0121SordariomycetidaeAgariaceaeLecanorineaeSphaerophoraceaeUndisturbed0.810.0213LecanoromycetidaeA	Pleosporomycetidae	Pleosporales	Pleosporaceae	Pleospora	Undisturbed	0.88	0.011	4
Chaetothyriomycetidae Chaetothyriales Chaetothyriales Claetothyriomycetidae Pleosporomycetidae Pleosporales Phaeotrichaceae Phaeotrichum Undisturbed 0.86 0.011 2 Chaetothyriomycetidae Chaetothyriales Chaetothyriales Sarcinomyces Undisturbed 0.86 0.011 2 Dothideomycetes Mycocaliciales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 10 Leotomycetes Mycotrichaceae Lendraea Unclassified Undisturbed 0.85 0.012 10 Lotomycetes Botryosphaeriales Botryosphaeriaceae Botryosphaeria Undisturbed 0.85 0.014 7 Dothideomycetidae Lecanorales Botryosphaeriaceae Botryosphaeria Undisturbed 0.82 0.016 1 Leotomycetidae Lacanorales Lecanorineae Cladoniaceae Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Aylariaceae Hypoxylon Undisturbed 0.82 0.016 1 Leotomycetidae Xylariales Aylariaceae Hypoxylon Undisturbed 0.82 0.016 1 Dothideomycetidae Sylariales Aylariaceae Hypoxylon Undisturbed 0.82 0.012 1 Helotiales Bulgaria Unclassified Undisturbed 0.82 0.012 1 Helotiales Bulgariaceae Bulgaria Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Amphisphaeriaceae Pseudohalonectria Undisturbed 0.82 0.016 1 Sordariomycetidae Kirschsteiniothelia Unclassified Unclassified Undisturbed 0.82 0.016 1 Sordariomycetidae Agaricales Lycoperdaceae Delphinella Old 0.82 0.016 1 Sordariomycetidae Agaricales Lycoperdaceae Delphinella Old 0.82 0.016 1 Dothideomycetidae Cheotohyriales Chaetothyriales Chaetohyriales Old 0.82 0.015 1 Chaetothyriomycetidae Cheotohyriales Chaetothyriales Chaetohyriales Old 0.82 0.015 1 Chaetothyriales Chaetohyriales Chaetohyriales Sariomyces Old 0.82 0.025 1 Mycocaliciales Sphinctrinaceae Epiphinella Old 0.82 0.025 1 Chaetothyriales Chaetohyriales Chaetohyriales Sariomyces Old 0.80 0.027 3 Sporadorichida Halteria Anteriae Unclassified Old 0.82 0.025 1 Chaetothyriales Chaetohyriales Chaetohyriales Sariomyces Old 0.80 0.025 1 Dothideomycetidae Halteria Anteriae Botricaeae Epiphinelia Old 0.79 0.030 4 Hypocrealorycetidae Halteriae Anteriae Epiphaeria Old 0.79 0.030 4 Hypocrealorycetidae Hypocreales Hypocreae Co	Dothideomycetidae	Dothideales	Dothioraceae	Columnosphaeria	Undisturbed	0.86	0.012	17
Pleosporomycetidae Pleosporales Phaeotrichaceae Phaeotrichaceae Phaeotrichaceae Undisturbed 0.86 0.011 2 Chaetothyriales Chaetothyriales Sarcinomyces Undisturbed 0.86 0.012 11 Dothideomycetes Tubeufiaceae Letendraea Unclassified Undisturbed 0.86 0.012 10 Loctiomycetes Myxotrichaceae Geomyces Undisturbed 0.85 0.014 7 Dothideomycetes Botryosphaeriales Botryosphaeriaceae Botryosphaeria Undisturbed 0.82 0.016 1 Locanoromycetidae Dothideales Dothideales Hortaea Undisturbed 0.82 0.016 1 Dothideomycetidae Dothideales Dothideales Hortaea Undisturbed 0.82 0.012 1 Dothideomycetidae Xylariales Xylariaceae Hypoxylon Undisturbed 0.82 0.012 1 Dothideomycetidae Xylariales Vylariaceae Hypoxylon Undisturbed 0.82 0.012 1 Dothideomycetidae Xylariales Xylariaceae Hypoxylon Undisturbed 0.82 0.012 1 Dothideomycetidae Xylariales Xylariaceae Hypoxylon Undisturbed 0.82 0.012 1 Dothideomycetidae Xylariales Xylariaceae Bulgaria Unclassified Undisturbed 0.82 0.012 1 Dothideomycetidae Xirschsteiniothelia Unclassified Undisturbed 0.82 0.016 1 Pezizales Pezizaceae Peziza Unclassified Undisturbed 0.82 0.016 1 Dothideomycetidae Agaricales Lecanorineae Sphaerophoraceae Undisturbed 0.82 0.016 1 Dothideomycetidae Agaricales Lecanorineae Sphaerophoraceae Undisturbed 0.81 0.021 3 Lecanoromycetidae Agaricales Lecanorineae Sphaerophoraceae Undisturbed 0.81 0.021 3 Lecanoromycetidae Agaricales Lecanorineae Sphaerophoraceae Undisturbed 0.81 0.021 1 Dothideomycetidae Agaricales Dothioraceae Delphinella Old 0.91 0.004 1 Eurotiomycetidae Agaricales Chaetothyriales Graetohyriales G	Chaetothyriomycetidae	Chaetothyriales	Chaetothyriales	Glyphium	Undisturbed	0.86	0.012	19
Chaetothyriomycetidae Chaetothyriales Chaetothyriales Sphinctrinaceae Undisturbed 0.86 0.011 2 Mycocaliciomycetidae Mycocaliciales Sphinctrinaceae Sphinctrina Undisturbed 0.86 0.012 11 Dothideomycetes Myxotrichaceae Myxotrichaceae Geomyces Undisturbed 0.85 0.012 6 Sporadotrichida Halteriidae Halteria Unclassified Undisturbed 0.85 0.014 7 Dothideomycetiae Lecanorales Lecanorineae Cladoniaceae Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Botryosphaeriaceae Botryosphaeria Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Arylariaceae Hypoxylon Undisturbed 0.82 0.012 1 Xylariomycetidae Xylariales Amphisphaeriaceae Hypoxylon Undisturbed 0.82 0.016 1 Dothideomycetidae Xylariales Amphisphaeriaceae Botryosphaeria Undisturbed 0.82 0.012 1 Helotiales Dothideales Outlideae Hortaea Undisturbed 0.82 0.012 1 Sordariomycetidae Xylariales Amphisphaeriaceae Pestalosphaeria Undisturbed 0.82 0.016 2 Pezizales Pezizaceae Peziza Unclassified Undisturbed 0.82 0.016 1 Sordariomycetidae Magnaporthales Magnaporthaceae Pseudohalonectria Undisturbed 0.82 0.016 1 Sordariomycetidae Lecanorales Lecanorineae Sphaerophoraceae Undisturbed 0.81 0.021 3 Lecanorales Lecanorineae Sphaerophoraceae Undisturbed 0.81 0.021 3 Lecanorales Lecanorineae Sphaerophoraceae Undisturbed 0.81 0.021 3 Lecanoralycetidae Dothideales Dothideales Dothioraceae Chromocleista Old 0.82 0.025 1 Dothideomycetidae Chaetothyriales Chaetothyriales Glyphium Old 0.82 0.025 1 Dothideomycetidae Chaetothyriales Sphinctrinae Sphaerophoraceae Old 0.80 0.027 3 Sporadotrichide Halteriide Halteria Unclassified Old 0.80 0.027 1 Sporadotrichide Halteria Unclassified Old 0.80 0.025 1 Mycocaliciales Sphinctrinaceae Sphinctrina Old 0.79 0.030 4 Hypocreanycetidae Halteria Unclassified Old 0.79 0.030 4 Hypocreales Hypocreae Columnosphaeria Old 0.79 0.031 2 Dothideomycetidae Halteria Unclassified Old 0.79 0.030 4 Hypocreales Hypocreaee Hamigera Young 0.86 0.013 1 Eurotiomycetidae Eurotiales Trichocomaceae Hamigera Young 0.86 0.013 1 Eurotiomycetidae Eur	Pleosporomycetidae	Pleosporales	Phaeotrichaceae	Phaeotrichum	Undisturbed	0.86	0.011	2
MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaUndisturbed0.860.01211DothideomycetesTubeufiaceaeLetendraeaUnclassifiedUndisturbed0.860.01210SporadotrichidaHalteridaeHalteriaUnclassifiedUndisturbed0.850.0126SporadotrichidaHalteriaUnclassifiedUndisturbed0.820.0161DothideomycetisBotryosphaeriaceaeBotryosphaeriaceaeUndisturbed0.820.0161DothideomycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0121XylariomycetidaeXylarialesAmphisphaeriaceaeHypoxylonUndisturbed0.820.0121SordariomycetidaeBulgariaceaeBulgariaUnclassifiedUnclassifiedUndisturbed0.820.0121DothideomycetesKirschsteiniotheliaMagnaporthaceaePezizaeUnclassifiedUndisturbed0.820.0162PezizalesPeziacaeaMagnaporthaceaeDelphinellaOld0.820.0161DothideomycetidaeAgaricalesLycoperdaceaeDelphinellaOld0.820.0161SordariomycetidaeAgaricalesLycoperdaceaeDelphinellaOld0.820.0161DothideomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251ChaetothyriomycetidaeChaetothyrialesChaetothyria	Chaetothyriomycetidae	Chaetothyriales	Chaetothyriales	Sarcinomyces	Undisturbed	0.86	0.011	2
DothideomycetesTubeufiaceaeLetendraeaUnclassifiedUndisturbed0.860.01210LeotiomycetesMyxotrichaceaeMyxotrichaceaeGeomycesUndisturbed0.850.0126DothideomycetesBotryosphaerialesBotryosphaeriaceaeBotryosphaeriaUndisturbed0.820.0161LecanoromycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0152XylariomycetidaeXylarialesXylariaceaeHypoxylonUndisturbed0.820.0121DothideomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaUndisturbed0.820.0121DothideomycetidaeSugariaceaeBulgariaceaeBulgariaUnclassifiedUndisturbed0.820.0121DothideomycetidaeKirschsteiniotheliaUnclassifiedUnclassifiedUndisturbed0.820.0121SordariomycetidaeMagnaporthaceaePezizaUnclassifiedUndisturbed0.820.0161SordariomycetidaeAgaricalesLycoperdanUndisturbed0.820.0161LecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.820.0161LotideomycetidaeAgaricalesLycoperdanUndisturbed0.820.0161LecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.820.0123LecanoralesLecanorineaeChaetothyrialesChaetothyrialesOld <td>Mycocaliciomycetidae</td> <td>Mycocaliciales</td> <td>Sphinctrinaceae</td> <td>Sphinctrina</td> <td>Undisturbed</td> <td>0.86</td> <td>0.012</td> <td>11</td>	Mycocaliciomycetidae	Mycocaliciales	Sphinctrinaceae	Sphinctrina	Undisturbed	0.86	0.012	11
LectomycetesMyxotrichaceaeMyxotrichaceaeGeomycesUndisturbed0.850.0126SporadotrichidaHalteriaHalteriaUnclassifiedUndisturbed0.850.0147DothideomycetesBotryosphaeriaceaBotryosphaeriaUndisturbed0.820.0161LecanoromycetidaeLecanoralesLecanorineaeCladoniaceaeUndisturbed0.820.0161DothideomycetidaeXylarialesXylariaceaeHypoxylonUndisturbed0.820.0121XylariomycetidaeXylarialesXylariaceaePestalosphaeriaUndisturbed0.820.0121DothideomycetesKirschsteiniotheliaUnclassifiedUndisturbed0.820.0121DothideomycetidaeMagnaporthaceaePezizaUnclassifiedUndisturbed0.820.0162PezizalesPezizaceaePezizUnclassifiedUndisturbed0.820.0161SordariomycetidaeAgaricalesLycoperdaceaeLycoperdonUndisturbed0.810.0185AgaricomycetidaeAgaricalesLycoperdaceaeLycoperdonUndisturbed0.820.0161DothideonycetidaeDothidealesDothidealesGlyphinnOld0.820.0161EurotomycetidaeAgaricalesLycoperdaceaeLycoperdonUndisturbed0.820.0161DothideonycetidaeChaetothyrialesChaetothyrialesGlyphinnOld0.82	Dothideomycetes	Tubeufiaceae	Letendraea	Unclassified	Undisturbed	0.86	0.012	10
SporadotrichidaHalteriidaeHalteriaUnclassifiedUndisturbed0.820.0147DothideomycetesBotryosphaerialesBotryosphaeriaceeBotryosphaeriaUndisturbed0.820.0161DothideomycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0161DothideomycetidaeXylarialesXylariaceaeHyotaeaUndisturbed0.820.0121XylariomycetidaeXylarialesXylariaceaeHyotaeaUnclassifiedUndisturbed0.820.0121HelotialesBulgariaUnclassifiedUnclassifiedUndisturbed0.820.0121DothideomycetidaeMagnaporthalesMagnaporthaceaePseudohalonectriaUnclassifiedUndisturbed0.820.0161SordariomycetidaeMagnaporthalesMagnaporthaceaeSpaerophoraceaeUndisturbed0.810.0213LecanoromycetidaeLecanoralesLecanorineaeSpherophoraceaeUndisturbed0.810.0185AgaricomycetidaeDothidealesDothioraceaeDelphinellaOld0.820.0251DothideomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251DothideomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0242DothideomycetidaeDothidealesDothioraceaeChaetothyrialesGlyphiumOld0.800.0242 <tr< td=""><td>Leotiomycetes</td><td>Myxotrichaceae</td><td>Myxotrichaceae</td><td>Geomyces</td><td>Undisturbed</td><td>0.85</td><td>0.012</td><td>6</td></tr<>	Leotiomycetes	Myxotrichaceae	Myxotrichaceae	Geomyces	Undisturbed	0.85	0.012	6
DothideomycetesBotryosphaerialesBotryosphaeriaceaeUndisturbed0.820.0161LecanoronycetidaeDothidealesLecanorineaeCladoniaceaeUndisturbed0.820.0161DothideomycetidaeXylarialesXylariaceaeHortaeaUndisturbed0.820.0152XylariomycetidaeXylarialesAmphisphaeriaceaeHypoxylonUndisturbed0.820.0121HelotialesBulgariaceaeBulgariaUnclassifiedUndisturbed0.820.0161DothideomycetesKirschsteiniotheliaUnclassifiedUnclassifiedUndisturbed0.820.0161SordariomycetidaeMagnaporthalesLecanorineaePseudohalonectriaUndisturbed0.820.0161SordariomycetidaeAgaricalesLycoperdaceaeLycoperdonceaeUndisturbed0.810.0213LecanoroycetidaeDothidealesDothicaceaeDelphinellaOld0.820.0151DothideomycetidaeEurotialesTrichocomaceaeChaetothyrialesGlyphiumOld0.820.0251ChaetothyrialesChaetothyrialesChaetothyrialesSphinctrinaOld0.800.0273SporadotrichidaHalteridaeHalteriaUnclassifiedOld0.800.0251ChaetothyriomycetidaeDothidealesDothioraceaeSphinctrinaOld0.800.0251ChaetothyriomycetidaeHalteriidaeHalteria <td>Sporadotrichida</td> <td>Halteriidae</td> <td>Halteria</td> <td>Unclassified</td> <td>Undisturbed</td> <td>0.85</td> <td>0.014</td> <td>7</td>	Sporadotrichida	Halteriidae	Halteria	Unclassified	Undisturbed	0.85	0.014	7
LecanoromycetidaeLecanorineaeCladoniaceaeUndisturbed0.820.0161DothideomycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0152XylarionycetidaeXylarialesXylariaceaeHypoxylonUndisturbed0.820.0121BulgariaBulgariaceaeBulgariaUnclassifiedUndisturbed0.820.0121DothideomycetidaeKirschsteiniotheliaUnclassifiedUnclassifiedUndisturbed0.820.0161PezizalesPezizaceaePezizaUnclassifiedUndisturbed0.820.0161SordariomycetidaeAgaricalesLeconorineaeSphaerophoraceaeUndisturbed0.810.0213LecanoronycetidaeAgaricalesLycoperdaceaeLycoperdonUndisturbed0.820.0151DothideomycetidaeDothidealesDothioraceaeChoronceleistaOld0.820.0251ChaetothyrialesChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251ChaetothyrialesChaetothyrialesChaetothyrialesSarionmycesOld0.800.0251ChaetothyrialesChaetothyrialesShinctrinaceaeSphinctrinaceaeSphinctrinaceaeSphinctrinaceaeSphinctrinaceaeNold0.800.0251SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.02555PleosporalesHypocr	Dothideomycetes	Botryosphaeriales	Botryosphaeriaceae	Botryosphaeria	Undisturbed	0.82	0.016	1
DothideomycetidaeDothidealesDothidealesHortaeaUndisturbed0.820.0152XylariomycetidaeXylarialesXylariaceaeHypoxylonUndisturbed0.820.0231HelotialesBulgariaceaeBulgariaAmphisphaeriaceaePestalosphaeriaUndisturbed0.820.0121BulgariaceaeBulgariaUnclassifiedUnclassifiedUndisturbed0.820.0162PezizaceaePezizaUnclassifiedUnclassifiedUndisturbed0.820.0161SordariomycetidaeLecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.810.0213LecanorowycetidaeAgaricalesLycoperdaceaeDelphinellaOld0.820.0151DothideomycetidaeDothidealesDothicaceaeDelphinellaOld0.820.0251DothideomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0251ChaetothyriomycetidaeChaetothyrialesChaetothyrialesSarcinomyces0.0251ChaetothyrialesChaetothyrialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0251MycocaliciomycetidaeDothidealesSphinctrinaceaeSphinctrinaOld0.800.0251MycocaliciamycetidaePleosporalesLeptosphaeriaceaeColumnosphaeriaOld </td <td>Lecanoromycetidae</td> <td>Lecanorales</td> <td>Lecanorineae</td> <td>Cladoniaceae</td> <td>Undisturbed</td> <td>0.82</td> <td>0.016</td> <td>1</td>	Lecanoromycetidae	Lecanorales	Lecanorineae	Cladoniaceae	Undisturbed	0.82	0.016	1
XylariomycetidaeXylarialesXylariaceaeHypoxylonUndisturbed0.820.0231XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaUndisturbed0.820.0121DothideomycetesKirschsteiniotheliaUnclassifiedUnclassifiedUndisturbed0.820.0162PezizalesPezizaceaePezizaUnclassifiedUndisturbed0.820.0161SordariomycetidaeMagnaporthalesMagnaporthaceaeSphaerophoraceaeUndisturbed0.810.0213LecanoromycetidaeLecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.810.0161DothideomycetidaeDothidealesDothicraceaeDelphinellaOld0.820.0151DothideomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0256DothideomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0251MycocalicionycetidaeDoyccalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0273SporadotrichidaHalteriideeHalteriaUnclassifiedOld0.800.0273SporadotrichidaHalteriideeHalteriaUnclassified <t< td=""><td>Dothideomycetidae</td><td>Dothideales</td><td>Dothideales</td><td>Hortaea</td><td>Undisturbed</td><td>0.82</td><td>0.015</td><td>2</td></t<>	Dothideomycetidae	Dothideales	Dothideales	Hortaea	Undisturbed	0.82	0.015	2
XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaUndisturbed0.820.0121HelotialesBulgariaceaeBulgariaUnclassifiedUndisturbed0.820.0121DothideomycetesKirschsteiniotheliaUnclassifiedUnclassifiedUndisturbed0.820.0162PezizalesPezizaceaePezizaUnclassifiedUndisturbed0.820.0161SordariomycetidaeMagnaporthalesMagnaporthaceaePseudohalonectriaUndisturbed0.810.0213LecanoromycetidaeLecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.810.0185AgaricomycetidaeDothidealesDothioraceaeDelphinellaOld0.760.0391DothideomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0151ChaetothyrialesChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251ChaetothyriomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0251MypocreeidaeLypocrealesHypocrealesEuposphaeriaceaeLeptosphaeriaeOld0.800.0251PleosporomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0342NylarionycetidaeHypocrealesHypocrea	Xylariomycetidae	Xylariales	Xylariaceae	Hypoxylon	Undisturbed	0.82	0.023	1
HelotialesBulgariaceaeBulgariaUnclassifiedUnclassifiedUndisturbed0.820.0121DothideomycetesKirschsteiniotheliaUnclassifiedUnclassifiedUndisturbed0.820.0162PezizalesPezizaceaePezizaUnclassifiedUndisturbed0.820.0161SordariomycetidaeMagnaporthalesMagnaporthaceaePseudohalonectriaUndisturbed0.810.0213LecanoromycetidaeLecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.810.0041DothideomycetidaeDothidealesDothioraceaeDelphinellaOld0.820.0151ChaetothyriomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0256DothideomycetesTubeufiaceaeLetendraeaUnclassifiedOld0.820.0251ChaetothyriomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0251MycocaliciomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0251HypocrealesHypocreaceaeHypocreaeColumnosphaeriaOld0.790.0304HypocrealesHypocreaceaeHypocreaePestosphaeria<	Xylariomycetidae	Xylariales	Amphisphaeriaceae	Pestalosphaeria	Undisturbed	0.82	0.012	1
DothideomycetesKirschsteiniotheliaUnclassifiedUnclassifiedUnclassifiedUndisturbed0.820.0162PezizaceaePezizaUnclassifiedUnclassifiedUndisturbed0.820.0161SordariomycetidaeMagnaporthalesMagnaporthaceaePseudohalonectriaUndisturbed0.810.0213LecanoromycetidaeLecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.810.0185AgaricomycetidaeDothidealesDothioraceaeDelphinellaOld0.910.0041DothideomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0256DothideomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251ChaetothyriomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0251MycocalicialesMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0251SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0251PloporeomycetidaePloporealesLeptosphaeriaceaeLeptosphaeriaOld0.800.0255PloporeomycetidaePloporealesHypocreaceaHypocreaOld0.790.0304HypocrealesHypocreaceaHypocreaceaHypocreaceaYoung0.860.0132NylariomycetidaeKiralesTrichocomaceaeBio	Helotiales	Bulgariaceae	Bulgaria	Unclassified	Undisturbed	0.82	0.012	1
PezizalesPezizaceaePezizaUnclassifiedUndisturbed0.820.0161SordariomycetidaeMagnaporthalesLecanorineaePseudohalonectriaUndisturbed0.810.0213LecanoromycetidaeAgaricalesLycoperdaceaeSphaerophoraceaeUndisturbed0.760.0391DothideomycetidaeDothidealesDothioraceaeDelphinellaOld0.910.0041EurotiomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0251OthideomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251OthideomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0251OthideomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273MycocaliciomycetidaeHalteriidaeHalteriaUnclassifiedOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothioaceaeColumnosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.770.0342NyalariomycetidaeHypocrealesHypocreaceaeHypocreaOld0.770.0314HypocreomycetidaeHypocrealesBionectriaceaeHamigeraYoung0.860.0131 <td>Dothideomycetes</td> <td>Kirschsteiniothelia</td> <td>Unclassified</td> <td>Unclassified</td> <td>Undisturbed</td> <td>0.82</td> <td>0.016</td> <td>2</td>	Dothideomycetes	Kirschsteiniothelia	Unclassified	Unclassified	Undisturbed	0.82	0.016	2
SordariomycetidaeMagnaporthalesMagnaporthaceaePseudohalonectriaUndisturbed0.810.0213LecanoromycetidaeLecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.810.0185AgaricomycetidaeAgaricalesLycoperdaceaeLycoperdonUndisturbed0.760.0391DothidealesDothioraceaeDelphinellaOld0.910.0041EurotiomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0256DothideomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251ChaetothyriomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273MycocaliciomycetidaeHalteriidaeHalteriaUnclassifiedOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothioraceaeColumnosphaeriaOld0.790.0304HypocrealesHypocreaceaeHypocreacOld0.790.0342XylariomycetidaeEurotialesTrichocomaceaeHaingeraYoung0.860.0132LeotomycetidaeEurotialesTrichocomaceaeHaingeraYoung0.860.0132LyppcreendycetidaeEurotialesTrichocomaceaeHaingeraYoung0.860.0132Lopothideomycetidae <td>Pezizales</td> <td>Pezizaceae</td> <td>Peziza</td> <td>Unclassified</td> <td>Undisturbed</td> <td>0.82</td> <td>0.016</td> <td>1</td>	Pezizales	Pezizaceae	Peziza	Unclassified	Undisturbed	0.82	0.016	1
LecanoromycetidaeLecanoralesLecanorineaeSphaerophoraceaeUndisturbed0.810.0185AgaricomycetidaeAgaricalesLycoperdaceaeLycoperdonUndisturbed0.760.0391DothideomycetidaeDothidealesDothioraceaeDelphinellaOld0.910.0041EurotiomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0151ChaetothyriomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0251ChaetothyriomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0251MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaePleosporalesLeptosphaeriaceaeColumnosphaeriaOld0.760.0342PleosporomycetidaeHypocrealesHypocreaceaeHypocreaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.740.0451HypocrealesHypocreaceaeHampisrhaeriaceaePestalosphaeriaOld0.740.0451HypocrealesBionectriaceaeBionectriaceaeYoung0.860.0131EurotiomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.86 </td <td>Sordariomycetidae</td> <td>Magnaporthales</td> <td>Magnaporthaceae</td> <td>Pseudohalonectria</td> <td>Undisturbed</td> <td>0.81</td> <td>0.021</td> <td>3</td>	Sordariomycetidae	Magnaporthales	Magnaporthaceae	Pseudohalonectria	Undisturbed	0.81	0.021	3
AgaricomycetidaeAgaricalesLycoperdaceaeLycoperdonUndisturbed0.760.0391DothideomycetidaeDothidealesDothioraceaeDelphinellaOld0.910.0041EurotiomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0151ChaetothyriomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0256DothideomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.820.0251MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0251MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.740.0451KylariomycetidaeHypocrealesTrichocomaceaeHamigeraYoung0.860.0132KylariomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeHypocrealesBionectriaceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeHypocrealesBionectriaceaeBionectriaceaeYo	Lecanoromycetidae	Lecanorales	Lecanorineae	Sphaerophoraceae	Undisturbed	0.81	0.018	5
DothideomycetidaeDothidealesDothioraceaeDelphinellaOld0.910.0041EurotiomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0151ChaetothyriomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0256DothideomycetesTubeufiaceaeLetendraeaUnclassifiedOld0.820.0251MycocaliciomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0273MycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0242DothideomycetidaeHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0304HypocreomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaOld0.750.0342XylariomycetidaeHypocrealesHypocreaeePestalosphaeriaOld0.740.0451HypocreomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0132LorotiomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0131EurotiomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.82	Agaricomycetidae	Agaricales	Lycoperdaceae	Lycoperdon	Undisturbed	0.76	0.039	1
EurotiomycetidaeEurotialesTrichocomaceaeChromocleistaOld0.820.0151ChaetothyriomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0256DothideomycetesTubeufiaceaeLetendraeaUnclassifiedOld0.820.0251ChaetothyriomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0251MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0255PleosporomycetidaeHypocrealesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.740.0451EurotiomycetidaeKylarialesAmphisphaeriaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeHypocrealesBionectriaceaeEupenicilliumYoung0.840.0165LeotiomycetidaeDothidealesDothioraceaeEupenicilliumY	Dothideomycetidae	Dothideales	Dothioraceae	Delphinella	Old	0.91	0.004	1
ChaetothyriomycetidaeChaetothyrialesChaetothyrialesGlyphiumOld0.820.0256DothideomycetesTubeufiaceaeLetendraeaUnclassifiedOld0.820.0251ChaetothyriomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0251MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0255PleosporomycetidaeHypocrealesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.860.0132XylariomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0132DothideomycetidaeHypocrealesBionectriaceaeEupenicilliumYoung0.840.0165LeotomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.820.0211HypocrealesMyxotrichaceaeMyxotrichaceaeGeomycesYoung<	Eurotiomycetidae	Eurotiales	Trichocomaceae	Chromocleista	Old	0.82	0.015	1
DothideomycetesTubeufiaceaeLetendraeaUnclassifiedOld0.820.0251ChaetothyriomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0251MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0255PleosporomycetidaeHypocrealesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaePestalosphaeriaOld0.740.0451HypocreomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131HypocrealesBionectriaceaeBionectriaceaeYoung0.840.0165LeotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.820.0211MycocalicialesMyxotrichaceaeGeomycesYoung0.820.0211MycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0211Hypocreales	Chaetothyriomycetidae	Chaetothyriales	Chaetothyriales	Glyphium	Old	0.82	0.025	6
ChaetothyriomycetidaeChaetothyrialesChaetothyrialesSarcinomycesOld0.800.0251MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0255PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreacycetidaeHypocreaceaeHypocreaceaeHypocreaOld0.750.0342XylariomycetidaeEurotialesTrichocomaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0132DothideomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetidaeMycorcilicalesSphinctrinaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycorcilicalesSphinctrinaceaeSphinctrinaYoung0.820.0121PleosporomycetidaePleosporalesLeotosphaeriaceaeSphinc	Dothideomycetes	Tubeufiaceae	Letendraea	Unclassified	Old	0.82	0.025	1
MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaOld0.800.0273SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0255PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreomycetidaeHypocreaceaeHypocreaceaeHypocreaOld0.750.0342XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0132DothideomycetidaeEurotialesTrichocomaceaeBionectriaceaeYoung0.860.0131EurotiomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotomycetesMyxotrichaceaeMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaePleosporonelesLeotosphaeriaceaeSphinctrinaYoung0.820.0121	Chaetothyriomycetidae	Chaetothyriales	Chaetothyriales	Sarcinomyces	Old	0.80	0.025	1
SporadotrichidaHalteriidaeHalteriaUnclassifiedOld0.800.0242DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0255PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.750.0342XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.860.0132DothideomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetidaeEurotialesTrichocomaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMyxotrichaceaeMyxotrichaceaeGeomycesYoung0.820.0121PleosporomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121PleosporalesLeptosphaeriaceaeLeptosphaeriaYoung0.820.0121	Mycocaliciomycetidae	Mycocaliciales	Sphinctrinaceae	Sphinctrina	Old	0.80	0.027	3
DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaOld0.790.0255PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.750.0342XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0132DothideomycetidaeEurotialesTrichocomaceaeBionectriaceaeYoung0.860.0131EurotiomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotomycetesMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121PleosporomycetidaePleosporalesLeotosphaeriaceaeLeotosphaeriaYoung0.790.0412	Sporadotrichida	Halteriidae	Halteria	Unclassified	Old	0.80	0.024	2
PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaOld0.790.0304HypocreomycetidaeHypocrealesHypocreaceaeHypocreaOld0.750.0342XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.870.0114HypocreomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetidaeMyxotrichaceaeMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121	Dothideomycetidae	Dothideales	Dothioraceae	Columnosphaeria	Old	0.79	0.025	5
HypocrealesHypocrealesHypocreaceaeHypocreaOld0.750.0342XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.870.0114HypocreomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotomycetesMyxotrichaceaeMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaePleosporalesLeotosphaeriaceaeSphinctrinaYoung0.790.0412	Pleosporomycetidae	Pleosporales	Leptosphaeriaceae	Leptosphaeria	Old	0.79	0.030	4
XylariomycetidaeXylarialesAmphisphaeriaceaePestalosphaeriaOld0.740.0451EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.870.0114HypocreomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetidaeMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121	Hypocreomycetidae	Hypocreales	Hypocreaceae	Hypocrea	Old	0.75	0.034	2
EurotiomycetidaeEurotialesTrichocomaceaeHamigeraYoung0.870.0114HypocreomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetidaeMyxotrichaceaeGeomycesYoung0.820.0211MycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121PleosporomycetidaePleosporalesLeotosphaeriaceaeLeotosphaeriaYoung0.790.0412	Xylariomycetidae	Xylariales	Amphisphaeriaceae	Pestalosphaeria	Old	0.74	0.045	1
HypocreomycetidaeHypocrealesBionectriaceaeBionectriaceaeYoung0.860.0132DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetidaeMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121	Eurotiomycetidae	Eurotiales	Trichocomaceae	Hamigera	Young	0.87	0.011	4
DothideomycetidaeDothidealesDothioraceaeColumnosphaeriaYoung0.860.0131EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetesMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaYoung0.790.0412	Hypocreomycetidae	Hypocreales	Bionectriaceae	Bionectriaceae	Young	0.86	0.013	2
EurotiomycetidaeEurotialesTrichocomaceaeEupenicilliumYoung0.840.0165LeotiomycetesMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaYoung0.790.0412	Dothideomycetidae	Dothideales	Dothioraceae	Columnosphaeria	Young	0.86	0.013	1
LeotiomycetesMyxotrichaceaeGeomycesYoung0.820.0211MycocaliciomycetidaeMycocalicialesSphinctrinaceaeSphinctrinaYoung0.820.0121PleosporomycetidaePleosporalesLeptosphaeriaceaeLeptosphaeriaYoung0.790.0412	Eurotiomycetidae	Eurotiales	Trichocomaceae	Eupenicillium	Young	0.84	0.016	5
Mycocaliciomycetidae Mycocaliciales Sphinctrinaceae Sphinctrina Young 0.82 0.012 1 Pleosporomycetidae Pleosporales Leptosphaeriaceae Leptosphaeria Young 0.79 0.041 2	Leotiomycetes	Myxotrichaceae	Myxotrichaceae	Geomyces	Young	0.82	0.021	1
Pleosporomycetidae Pleosporales Leptosphaeriaceae Leptosphaeria Young 0.79 0.041 2	Mycocaliciomycetidae	Mycocaliciales	Sphinctrinaceae	Sphinctrina	Young	0.82	0.012	1
	Pleosporomycetidae	Pleosporales	Leptosphaeriaceae	Leptosphaeria	Young	0.79	0.041	2
Dothideomycetes Tubeufiaceae Letendraea Unclassified Young 0.79 0.026 2	Dothideomycetes	Tubeufiaceae	Letendraea	Unclassified	Young	0.79	0.026	2

Abbreviations: IV, indicator value; OTUs, operational taxonomic units. Only taxa with an IV > 0.75 are shown.

Table 3 Metrics obtained from analysis of scale-free microbial	l networks of bacterial and fungal microbial communities
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Taxon and microsite	OTUs	Edges	Mean number of neighbours	Clustering coefficient	Density	Centralization	
Bacteria					,		
Old	40	70	3.450	0.547	0.088	0.096	
Undisturbed	31	39	2.516	0.566	0.084	0.089	
Young	17	11	1.294	0	0.081	0.050	
Fungi							
Old	135	485	7.185	0.472	0.054	0.067	
Undisturbed	177	1814	20.497	0.652	0.166	0.095	
Young	81	321	7.926	0.647	0.009	0.116	

Abbreviation: OTUs, operational taxonomic units. Edges represent the number of significant positive and negative Pearson correlation coefficients identified following implementation of the Benjamini-Hochberg procedure at a minimum false discovery rate of 5%.

found, with the insect and plant pathogenic fungi, *Delphinella, Leptosphaeria, Trematosphaeria* and *Columnosphaeria*, found almost exclusively in undisturbed and young pit soils. *Glomeromycetes*, which comprise arbuscular mycorrhizal species, represented about 3% of sequences in young pits and 2% of sequences in old pit and undisturbed soils.

## Community development with pit age

Rubrobacter, Ammoniphilus and Actinaurispora were the only bacterial indicators of young pits and likely represent remnants of the sub-surface community. Rubrobacter is a cosmopolitan and abundant taxon in arid zone soils (Yeager et al., 2004). The presence of Ammoniphilus and Actinaurispora in young pit soils is likely due to the deposition of plant material. Amminophilus has been reported as a strictly aerobic oxalotroph utilizing plant- and algae-derived oxalic acid as a sole carbon. Actinaurispora are known plant endophytes, inhabiting Camptotheca acuminata species (Zhu et al., 2012). The family Micromonosporacaeae, to which Actinaurispora belongs, however, has been tentatively correlated with increasing moisture content in arid and semi-arid soils (Bachar et al., 2010), which may contribute to the presence of this species as an indicator of young soils. Trichocomaceae species were the primary fungal indicators of young pits. A single Trichocomaceae species was a key fungal indicator of undisturbed soils, suggesting that fungal communities of young pits contain residual surface taxa prior to the colonization and diversification of fungal communities observed in older pits. Trichocomaceae species are predominantly saprotrophic, have aggressive colonization strategies and a high tolerance to extreme environmental conditions such as soil drying, high temperature and metal toxicity (Houbraken and Samson, 2011). Their presence in young pits could indicate opportunistic colonization of recently disturbed soil.

On the basis of the criteria used to select the microsites, progression of the microbial community from young to old pits occurs over a period of 9-12 months. Over this time, although little change occurred within the microbial community composition between pit stages, a discernable difference was observed between the undisturbed and pit bacterial and fungal communities, irrespective of their age. Microbial richness among microsites, however, remained unchanged. Spore propagule density and arbuscular mycorrhizal fungi diversity are known to decline with increasing tillage associated with agriculture (Brito et al., 2010; Schalamuk et al., 2013). However, this was not reflected in our fungal species richness, which remained unchanged over time. The progressive accumulation of fungal species attached to organic matter and seed in the pits is consistent with the presence of several lichenized lecanoralean genera including Parmeliaceae and

Myrangium (Smith, 1948) and the epiphytic melanized taxon Sarcinomyces (Wollenzien et al., 1997). The Lecanorales are predominantly lichen-forming fungi that are mycobionts of the genera Xanthoparmelia, Parmotrema and Xanthoria, which are common corticolous lichens of Callitris glaucophylla trees that occur in the study area (Filson and Rogers, 1979). These taxa are typically found on the soil surface or in the pits on detached plant material. At some sites, we also recorded the vagant lichen Chondropsis semiviridis from within the pits. This lichen, which has no attachment to the soil, moves freely along the surface by wind action (Eldridge and Leys, 1999). Similarly, Cladonia spp., another common soil lichen genus, was found on undisturbed surfaces. Along with the lichen genera Endocarpon and Placidium, it is one of the most common lichens forming biocrusts on stable soils in arid and semi-arid areas (Eldridge and Koen, 1998).

Despite our inability to discriminate between the bacterial community of old and young pits, we recorded three indicator species, Hyalangium and Microvirga, and a Gp IV Acidobacteria. The two proteobacterial species were indicative of the presence of established vascular plants. Hyalangium, belongs to the group of Myxobacteria that uses plant lignin and produces small bioactive molecules. *Microvirga* has been implicated in nodule formation, facilitating nitrogen-fixing processes within the rhizosphere (Ardley et al., 2012). The occurrence of these groups in old pit soils is likely to enhance nitrogen fixation, presumably to levels greater than those in the undisturbed and young pit soils, and support the growth of vascular plants occurring in these microsites.

#### Microbial co-occurrence in pit and undisturbed soils

Our analyses thus far indicate that initial disturbance reduces the abundance of key photoautotrophic groups, and that over extended periods of time, capture of organic matter leads to changes in the abundance of some taxa, with increases in those taxa likely reflecting an increased capacity for the assimilation of organic carbon and nitrogen matter.

Resilience is the ability of a system to recover from large disturbances, typically over short time frames. Reactivity, however, is the capacity of a system to respond to small perturbations over extended periods. Under such circumstances, the apparent equilibrium may appear stable, despite moving to a new steady state over long time periods (Neubert et al., 2009). Modularity, defined by the number and size of groups of highly interconnected nodes within a network, is positively correlated with reactivity, and negatively correlated with resilience (Ruiz-Moreno et al., 2006). Analysis of both bacterial and fungal microbial networks revealed stark differences in modularity, reflected in the values of clustering, density and centralization, of microbial co-occurrence networks between undisturbed soils and pit soils at

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different developmental stages (Table 3). Clustering coefficients and density (network connectivity) scores tending towards a value of 1 indicate a highly modular system whereas those tending towards zero represent the opposite (Bissett *et al.*, 2013). Low values of clustering and density associated with microbial communities from contaminated and reference estuarine sediments indicate historical community 'stress' contributing to functional redundancy and reduced correlations among species (Sun *et al.*, 2013). This was reinforced by marginally lower values, for each of these metrics, in contaminated sediments, with the suggestion that this anthropogenic perturbation has contributed an additional stress.

In the present study, bacterial species-species correlations within the young pit soils were almost nonexistent. A clustering coefficient of zero and a slightly lower density value were consistent with reduced modularity, and an increase in functional redundancy associated with a recent external stress (Sun et al., 2013). In contrast, undisturbed and old pit soils were more consistent with increased modularity, suggesting a lack of functional redundancy, with greater species-species correlations, and increased clustering and density. This suggests to us that the bacterial community present in undisturbed soils and old pit soils are more reactive and less resilient than young pit soils. Within the fungal communities, the number of correlations among species, clustering coefficients and density, and hence modularity, were highest in surface soils and lowest in the old pit soils, suggesting that fungal communities within old pits are less reactive and more resilient. In contrast, the young pit soils exhibit reduced modularity, and increased resilience, suggesting that they are likely to respond to nutrient amendments over the short-term, thereby driving large and dramatic structural changes. This is largely because of the high degree of physical disturbance created when foraging pits are established. Within the old pit soils, the bacterial community has largely regained the modularity observed within the undisturbed soils. The fungal community, however, is apparently more resistant at this stage than in the undisturbed soils, suggesting it is able to continue to drive structural changes in response to events such as litter deposition.

A high level of centralization, as a consequence of the high frequency of centralized nodes, was observed among the fungal community in young pit soils (Bulgariaceae, Myxotrichaeae, Trichocomaceae, Tubeufiaceae) and among the bacterial community in undisturbed (Rubrobacteraceae, Geodematophiliaceae, Bradyrhizobiaceae) and old pit (Rhodobacteraceae, Bradyrhizobiaceae) and old pit (Rhodobacteraceae, Bradyrhizobiaceae, Geodermatophiliaceae, Beijerinckiaceae, Comamonadaceae, Methylobacteriaceae) soil (Supplementary Information). Centralized nodes have been proposed to represent keystone species, exhibiting a large influence of the 'information' transfer throughout the community (Bisssett *et al.*, 2013). It has been speculated that these nodes represent critical control points in the cycling of nutrients within the system (Ruiz-Moreno et al., 2006; Bissett et al., 2013). Thus, it is realistic to suggest that these centralized taxa act to stabilize the microbial community. It should be highlighted that these observations were made in the context of a small number of samples defining each microsite, as well as few sequence reads being available to identify species-species correlations. Our observations between the bacterial and fungal datasets suggest that these metrics are susceptible to sequence depth, and pre-treatment of the data by retaining only semi-ubiquitous (occurring across at least 75% of samples) OTUs. These values may also be influenced by the level of heterogeneity within microsites. Despite this, our analyses of network metrics from the bacterial communities suggested that the community structure of old pit soils reflect that of undisturbed soils. Over the long term, this would tend towards decreased responses to nutrient inputs into these soils. This, however, may be partially offset by frequent deposition of plant matter due to the establishment and growth of vascular plants within old pit soils, and subsequent assimilation of this matter by saprotrophic fungi.

# Conclusions

Our study suggests that digging by soil-disturbing animals is likely to create successional shifts in soil microbial and fungal communities, which could account for increases in organic matter of nitrogen in old foraging pits (James et al. 2009). The observed richness of fungal and bacterial OTUs, in undisturbed soils, and young and old pits did not differ, though fewer correlations, and hence an increased resilience, were observed between bacterial OTUs in young pits, and fungal OTUs in young and old pits. This suggests that these communities are more likely to respond over the short term to nutrient amendment, thus promoting nutrient enrichment and contributing to a form of patchiness that is critical for the functioning of arid systems. The action of soildisturbing animals therefore leads to the development of a mosaic of different patches with a varying complement of microorganisms. Given the wide variety in pit size, depth, substrate and spatial configuration, this differential microbial activity will likely lead to the creation of a mosaic of patches of differing resource availability, analogous to larger surface-resident biotic patches such as hummocks and debris mounds. Our work suggests that microbial community composition and co-occurrence change with physical disturbance during the formation of foraging pits. Given the primacy of organic matter decomposition in arid and semi-arid environments, the loss of native soil-foraging animals from these systems may well impair the ability of these systems to maintain key ecosystem processes and to recover from disturbance.

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## **Conflict of Interest**

The authors declare no conflict of interest.

## Acknowledgements

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