RESEARCH ARTICLE

Site and land-use associations of soil bacteria and fungi define core and indicative taxa

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One sentence summary: Soil bacterial and fungal communities at thirty sites were yearly assessed over five consecutive years and revealed temporally stable as well as land-use- and site-specific core communities.

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ABSTRACT

Soil microbial diversity has major influences on ecosystem functions and services. However, due to its complexity and uneven distribution of abundant and rare taxa, quantification of soil microbial diversity remains challenging and thereby impeding its integration into long-term monitoring programs. Using metabarcoding, we analyzed soil bacterial and fungal communities at 30 long-term soil monitoring sites from the three land-use types arable land, permanent grassland, and forest with a yearly sampling between snowmelt and first fertilization over five years. Unlike soil microbial biomass and alpha-diversity, microbial community compositions and structures were site- and land-use-specific with CAP reclassification success rates of 100%. The temporally stable site core communities included 38.5% of bacterial and 33.1% of fungal OTUs covering 95.9% and 93.2% of relative abundances. We characterized bacterial and fungal core communities and their land-use associations at the family-level. In general, fungal families revealed stronger land-use associations as...

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compared to bacteria. This is likely due to a stronger vegetation effect on fungal core taxa, while bacterial core taxa were stronger related to soil properties. The assessment of core communities can be used to form cultivation-independent reference lists of microbial taxa, which may facilitate the development of microbial indicators for soil quality and the use of soil microbiota for long-term soil biomonitoring.

**Keywords:** amplicon sequencing; soil microbial diversity; core taxa; core communities; environmental drivers; temporal stability

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## INTRODUCTION

Soil microorganisms constitute the majority of soil biodiversity (Bardgett and van der Putten 2014) and are main drivers of many soil processes (Costa et al. 2018; Hallin et al. 2018). A detailed understanding of belowground microbial diversity and of its influencing factors is the basis for a holistic view and understanding of ecosystem processes in terrestrial environments. However, a census of soil microorganisms remains largely incomplete, due to the enormous diversity and range of abundances of soil microorganisms. High microbial diversities have been observed at different scales ranging from aggregate (Hemkemeyer et al. 2018; Hemkemeyer et al. 2019), to landscape (Karimi et al. 2018), and global assessments (Bahram et al. 2018; Větrovský et al. 2019).

At the landscape scale, soil bacterial and fungal diversities are strongly correlated to soil pH (Lauber et al. 2009; Griffiths et al. 2011), which is caused by direct effects but also by indirect effects such as changing the availability of nutrients (Glassman et al. 2017; Lammel et al. 2018). The number of bacterial taxa in soils depends on the pH and has been reported to reach its maximum at pH values between 6 and 7 (Lauber et al. 2009). Furthermore, community structures of soil bacteria change with pH, because specific bacterial taxa reveal distinct pH preferences. For instance, within the phylum Acidobacteria, taxa belonging to the class Acidobacteria are in general negatively correlated to soil pH, while taxa belonging to Acidobacteria Subgroup 6 commonly reveal a positive correlation to soil pH (Kielak et al. 2016). Further drivers of bacterial community structures depend on the system studied and include factors such as soil texture, climate, and plant communities (Griffiths et al. 2016; Bahram et al. 2018; Karimi et al. 2018; Leff et al. 2018). In comparison to soil bacterial diversity, soil fungal diversity has been shown to be geographically more structured (Taibot et al. 2014; Bahram et al. 2018). In a global meta-analysis that covered 742 sites, Větrovský et al. (2019) identified climate factors as main drivers of soil fungal communities, followed by soil properties, and vegetation parameters. Finally, factors related to land management, such as agricultural intensity (Barnerjee et al. 2019), tillage (Degrune et al. 2017; Babin et al. 2019), fertilization (Hartmann et al. 2015; Piazza et al. 2019), or compaction (Hartmann et al. 2014) may influence diversity of soil bacteria and fungi. While the major environmental determinants of soil bacterial and fungal communities are largely known, less is known about common components of these communities, their taxonomic representatives, and their diversities.

Surveys of soil bacterial and fungal communities usually reveal a large number of unknown taxa. Delgado-Baquerizo (2019) has reported that in a global survey 99% of bacterial and 63% of fungal OTUs remained unclassified at the species-level, and that the number of unclassified bacterial or fungal OTUs at the phylum-level in a sample has ranged between 1.4% and 9.4%. In a meta-analysis on the global diversity of soil fungi, an average of only 53% of the sequences per sample could be assigned to entries in the UNITE reference database, which notably includes sequences from environmental samples (Větrovský et al. 2019). High ratios of unclassified sequences at the species level may be due to a lack of resolution of the used DNA barcodes (e.g. Gschwend et al. 2021a), or due to missing reference sequences. To elucidate the unknown microbial diversity and describe consistently occurring OTUs, several attempts have been made to identify the most common taxa, which could constitute a core of soil microbial communities (Delgado-Baquerizo et al. 2018; Egidi et al. 2019). OTUs contributing to the global bacterial soil core community were assigned in descending order of relative abundance to the phyla Proteobacteria, Actinobacteria, Planctomycetes, Chloroflexi, Verrucomicrobia, Bacteroidetes, Gemmatimonadetes, Firmicutes, Armatimonadetes, Saccharibacteria, and candidate division WS2 (Delgado-Baquerizo et al. 2018). Five of these phyla, i.e. Proteobacteria, Actinobacteria, Planctomycetes, Bacteroidetes, and Firmicutes, have also been reported among those with an average relative abundance of at least 5% in a soil bacterial survey across France (Karimi et al. 2018), which has identified Acidobacteria as an additional dominant phylum. Dominant soil bacterial phyla have revealed distinct ecological preferences such as Alphaproteobacteria and Verrucomicrobia that were more abundant in forest and permanent grassland as compared to arable and vineyard soils, while the inverse was found for Chloroflexi and Gemmatimonadetes (Karimi et al. 2018). However, diverse habitat associations are often detected for taxa assigned to the same phylum. For instance within the phylum Chloroflexi, the family Anaerolineaceae were associated to soils with pH above 5, while Ktedonobacteraceae were associated to a lower soil pH (Mayerhofer et al. 2021). For soil fungi, a global survey of 365 sites has revealed Ascomycota, Basidiomycota, Mortierellomycota, and Mucoromycota as dominant fungal phyla in soils (Tedersoo et al. 2014), which has been largely confirmed, although the high abundance of Mortierellomycota has been questioned (Větrovský et al. 2019; Egidi et al. 2019) have proposed that globally dominant soil fungal OTUs almost exclusively derived from Ascomycota with 80 of 83 dominant fungal OTUs classified to this phylum. Despite the recent interest in taxonomic surveys of soil bacterial (Delgado-Baquerizo et al. 2018; Karimi et al. 2018; Walsh et al. 2019) and fungal diversity (Tedersoo et al. 2017; Egidi et al. 2019), habitat associations of soil bacteria and fungi at lower taxonomic levels are still largely lacking.

In a previous study, 30 long-term monitoring sites of the Swiss Soil Monitoring Network (NABO) have been surveyed over five years to define and assess long-term stability of abundant, rare, and scarce soil bacterial and fungal community components (Gschwend et al. 2021b). Soil bacterial and fungal communities of different sites sampled early in the vegetation period remained temporally stable and structurally distinct over five years. However, that study has not provided detailed analyses of environmental drivers of community structures among land-use types, and individual sites. Furthermore, it has focused on community structures and considered OTUs as anonymous entities without assessing their taxonomy and distributions.
To develop specific microbial indicators for assessing biological soil quality, information on habitat associations of bacteria and fungi at high taxonomic resolution is needed.

Here, we assessed soil bacterial and fungal diversity at the 30 sites of the NABO with a yearly sampling between snowmelt and first fertilization over five years previously described and studied by Gschwend et al. (2021b). Our research goals were to (i) assess site- and land-use-specific soil microbial community measures; (ii) identify taxa, which were consistently detectable (core OTUs) and taxa, which were associated to environmental factors (indictive OTUs); (iii) assess the main environmental factors structuring core communities; (iv) describe diversity and identity of core OTUs as well as their distribution among land-use types.

MATERIAL AND METHODS

Sampling design, DNA extraction and microbial biomass measurement

Samples were taken during five years, from 2012 to 2016, at thirty sites (Fig. S1, Supporting Information) of the Swiss Soil Monitoring Network (NABO) in early spring after snowmelt and before fertilization. Three land-use types, i.e. arable land, permanent grassland, and forests were sampled with ten sites each. Arable sites were managed with crop rotations, which included three to six different crops, and with one exception they were conventionally tilled. Forest sites included four coniferous, two mixed, and four deciduous forests. At each site, three composite samples composed of 25 soil cores of 20 cm depth and 2.5 cm diameter were taken from a 10 m by 10 m plot according to the standardized sampling protocol of the Swiss Soil Monitoring Network (Gubler et al. 2019). Samples were immediately stored at 4 °C after sampling and processed within 48 hours. Homogenized soil was mixed with DNA extraction buffer ([2% hexadecyl trimethyl ammonium chloride (CTAB); 20 mM EDTA pH 8; 2 M NaCl; 100 mM tris hydroxymethylaminomethane pH 8; 2% polyvinylpyrrolidone (PVP-40)], Lazzaro et al. 2006).

Quantitative DNA extraction was achieved by extracting DNA three times from each sample following Bürghmann et al. (2001) with the modifications by Hartmann et al. (2005). DNA quantity was determined using PicoGreen (Invitrogen, Carlsbad, CA) on a Cary Eclipse fluorescence spectrophotometer (Varian, Inc. Palo Alto, CA) and cross-validated using Qubit 1.0 (Life Technologies, Carlsbad, CA, USA). DNA was cleaned using the NucleoSpin® gDNA clean-up kit (Machery-Nagel, Düren, Germany) according to the manufacturer’s instruction. Microbial biomass carbon (Cmic) was assessed using chlorof orm fumigation-extraction according to Vance et al. (1987) with a kcc value of 0.45 (Joergensen 1996). Measurements of soil physicochemical properties, i.e. soil pH, total and organic carbon, total nitrogen, C/N-ratio, bulk density, soil texture and gravimetric water content, have been described in Gschwend et al. (2021b).

Barcode amplification, sequencing and sequence analysis

Bacterial variable region 3 and 4 of the small sub-unit of the ribosomal RNA gene (16S rRNA) were amplified using primers 341F (5′ CCTAYGGDGBCGWSCAG 3′) and 806R (5′ GGACTACNVGGGT HTCTAAT 3′) (Frey et al. 2016). Fungal internal transcribed spacer 2 (ITS2) was amplified using primers ITS3 (5′ CAGCAGTGAAG AACGYRG 3′) and ITS4 (5′ TCCCTGCTTTATTGATATGC 3′) (Teder soo et al. 2014). Four reactions using the GoTaq® Hot Start Polymerase (Promega) were done for each sample using 20 ng of DNA for each reaction. Reactions were performed according to Mayrhofer et al. (2017) with two modifications, which were an initial denaturation at 95 °C for two minutes, as well as 35 PCR cycles for the bacterial and fungal markers. Production of sequencing libraries and paired-end sequencing on an Illumina MiSeq v3 were performed at the Génome Québec Innovation Center at the McGill University (Montréal, Canada).

Raw sequences, (NCBI SRA: PRJNA660320) were quality filtered using a custom sequence analysis pipeline largely based on USEARCH version 9 (Edgar 2010; Frey et al. 2016) and is described in greater detail in Gschwend et al. (2021b). Only sequences occurring in at least two samples were allowed to form OTU centroids. Sequences were clustered into OTUs based on a 97% sequence identity threshold. This threshold was chosen to obtain a conservative estimate of soil microbial diversity and because diversity patterns between OTUs and sequence variants based approaches are highly correlated (Glassman and Martiny 2018). Taxonomic assignment was obtained using the RDP classifier implemented in mothur version 1.36.1 (Schloss et al. 2009) and a minimum bootstrap value of 80% with the SILVA 132 database (Quast et al. 2012) as reference for bacterial sequences. Eukaryotic sequences were classified with the same approach to a Genbank database (Frey et al. 2016) to discriminate between fungal and other eukaryote sequences. Fungal sequences were subsequently compared to the UNITE v 7.2 reference database (Nilsson et al. 2018).

Statistics

All analyses unless stated otherwise, were performed in R (RStudio 2015; R Core Team 2016). Mean values of environmental factors were calculated for samples taken at the same time point to avoid pseudo-replication. Similarly, calculations of alpha- and beta-diversity values were based on median values of OTUs per sampling time point. Spearman correlations were used to link univariate responses to environmental factors. Multivariate responses of communities were assessed by PERMDISP (Anderson et al. 2006) to evaluate homogeneity of dispersions between groups and permutational analysis of variance (PERMANOVA, Anderson 2001) to analyse between group differences. PRIMER7 (Clarke and Warwick 2001; Anderson et al. 2008) was used for PERMANOVA. To accommodate PERMANOVA for the repeated measurements design we created a nested PERMANOVA design according to the PERMANOVA manual (Anderson et al. 2008) and included land-use types as a fixed factor, sites as random factor nested within land-use type, and year as a random factor. Effects on community structures were expressed as square root of component of variation (√CV), which are in the unit of the original community dissimilarity, i.e. Bray–Curtis dissimilarity. The order of covariates in sequential PERMANOVA tests were selected based on the model selection algorithm implemented in distance-based linear model (DISTLM, McArdle and Anderson 2001) within PRIMER7, where AICc was chosen as model selection criterion. P-values of multiple tests were adjusted using Benjamini–Hochberg procedure (Benjamini and Hochberg 1995).

Site specificity was further assessed by leave-one-out cross-validation based on linear discriminant analysis (LDA) for univariate and based on canonical analysis of principal coordinates (CAP, Anderson and Willis 2003) for community structures. LDA and CAP were calculated within R using the functions ‘train’ of the package caret (Kuhn 2008) and ‘CAPdiscrim’ of the package ‘BiodiversityR’ (Kindt and Coe 2005), respectively. For the pairwise comparisons of similarities between land-use types, median values of OTU abundances were obtained for each site followed by determining Jaccard and Bray–Curtis similarities.
RESULTS

Increasing resolution from microbial biomass to community structures

Thirty sites from three land-use types, i.e. 10 each from arable land, permanent grassland, and forest, were surveyed with yearly samplings during five years, which yielded 450 samples. Soil microbial communities were assessed using three different approaches, which were (i) soil microbial biomass, i.e. based on soil microbial carbon (Cmic) content determined with chloroform fumigation extraction, and soil DNA content, that correlated (r = 0.79, P < 0.0001), (ii) alpha-diversity based on OTU richness, Simpson evenness and inverse Simpson index and (iii) beta-diversity based on Jaccard similarities and Bray-Curtis dissimilarities (Table 2, see also supplementary results for a summary of the sequencing data). Microbial biomass and alpha-diversity revealed no site- (reclassification ≤ 4.7%), and low land-use-specificity (reclassification ≤ 61.3%, Table 2). Values of both microbial biomass measures were significantly reduced in arable land (Tukey HSD, P < 0.0007; Table S1, Supporting Information), while bacterial alpha-diversity was increased in arable land (Tukey HSD, P = 0.0096; Table S1, Supporting Information). Fungal alpha-diversity with the exception of fungal OTU richness were significantly lower in forest soils (Tukey HSD, P < 0.01; Table S1, Supporting Information). Community compositions (Jaccard similarity) and structures (Bray-Curtis dissimilarity) were land-use- (Fig. 1) and site-specific with reclassification success rates of 100% for bacteria and fungi (Table 2). To resolve different drivers of soil bacteria and fungi, information on community compositions or structures was needed rather than bulk parameters such as microbial biomass or alpha-diversity.

Partitioning of OTUs into core and indicative groups

The high site-specificity of soil bacterial and fungal community structures, which was maintained over five years, also reflected a high temporal stability. Temporally stable core taxa, i.e. site-core (sc) OTUs and land-use-core (lc) OTUs were defined as outlined in Table 1. Of the 18140 bacterial OTUs (bOTUs) 6979 (38.5%), which covered 95.9% relative abundance were classified as sc-OTUs and 1136 of these sc-OTUs (covering 69.1% relative abundance) were also classified as lc-OTUs (Table 3). A similar proportion of the 8477 fungal OTUs (fOTUs), i.e. 2802 fOTUs (33.1%) and covering 93.2% relative abundance, was classified as sc-OTUs, but only 103 of them (29.4% relative abundance) were also classified as lc-OTUs. In addition to these core taxa, we defined indicative OTUs, i.e. OTUs that structured communities according to environmental conditions. More specifically, we distinguished three categories of indicative OTUs, i.e. OTUs correlated to environmental factor, as well as OTUs indicative for land-use types and OTUs indicative of a given site (see Table 1 for definitions). Most strikingly, the number and particularly the abundance of site-indicative OTUs was higher for fungi (1445 fOTUs, 29.9% relative abundance), as compared to bacteria (1146 bOTUs, 3.1% relative abundance). The vast majority of indicative OTUs were also classified as sc-OTUs (95% for bacteria, 90% for fungi, Fig. S2, Supporting Information). Communities composed of only sc-OTUs, i.e. core communities, were almost perfectly correlated (r = 0.97) to the entire communities, both in terms of alpha- and beta-diversity (Table S2, Supporting Information). Consequently, soil microbial core communities are representative of the respective entire communities. The following analyses were therefore based on these core communities.

Environmental factors driving structures of core communities

Soil habitats of different land-use types were characterized by distinct environmental factors. Arable sites were characterized by increased soil pH, and bulk density, and forest sites were characterized by increased carbon contents and C/N-ratios (Fig. 1), while grassland sites generally revealed intermediate levels of the assessed environmental factors (Table S1, Supporting Information). Soil bacterial and fungal core communities were mainly structured by soil pH and the C/N-ratio (Table 4). In addition to the environmental factors considered, land-use type and site significantly explained variance of soil bacterial (\( CV_{\text{Land-use type}} = 0.23 \), \( CV_{\text{Site}} = 0.31 \)) and fungal (\( CV_{\text{Land-use type}} = 0.31 \), \( CV_{\text{Site}} = 0.49 \)) community structures. Soil pH was the strongest driver for bacterial community structures overall and within each land-use type (Table S3, Supporting Information). The second strongest environmental factor in the overall analysis was the C/N-ratio, but it had no or minimal effects on the community structures within land-use types (Tables S3 and S4, Supporting Information). This may be due to the clear difference in C/N-ratio between forest and the other two land-use types (Table S1, Supporting Information), indicating that a high C/N-ratio represented a proxy for forest soils in the overall analysis. The separate analysis of arable sites also allowed to consider crop as an additional factor shaping microbial communities (Tables S3 and S4, Supporting Information), which was more strongly affecting fungal (\( CV = 0.16 \)) as compared to bacterial (\( CV = 0.06 \)) core communities. In line with the data on core community structures, the strongest correlations of individual OTUs to environmental factors were detected with soil pH for bacterial OTUs (Table S5, Supporting Information) and with soil pH, C/N-ratio, and organic carbon for fungal OTUs (Table S6, Supporting Information).
Table 1. Definitions of OTU groups and subgroups (see also Table 3).

| OTU group | Definition |
|-----------|------------|
| core OTUs | site-core OTUs (sc-OTUs) occur in at least 12 of the 15 samples from a site |
| land-use-core OTUs are a sc-OTU in at least 8 of the 10 sites of a land-use type |
| indicative OTUs environmental-factor-indicative OTUs | correlated to an environmental factor (Spearman rho > 0.4, P < 0.05) |
| site-indicative OTUs | indicative of an individual site (IndVal > 0.8, P < 0.05) |
| land-use-indicative OTUs | indicative of individual or combinations of land-use types (IndVal > 0.8, P < 0.05) |

1Environmental factors are summarized in Table S1 (Supporting Information).

Table 2. Site and land-use specific soil microbial communities at different analytical levels. Site and land-use type specificity was calculated using a leave-one-out reclassification test based on linear discriminant analysis for univariate and canonical analysis of principal coordinates (CAP) with 9999 permutations for community compositions and structures.

| Community parameter | Taxon       | LUT1 | P-value | Site |
|---------------------|-------------|------|---------|------|
|                      |             | Reclass² | P-value | Reclass² | P-value |
| Organic Carbon      |             | 60.7% | 6.95 * 10^-12 | 4.7% | 0.235 |
| Microbial biomass   |             | 60.0% | 2.13 * 10^-11 | 4.0% | 0.384 |
| DNA                 |             | 61.3% | 2.20 * 10^-12 | 2.0% | 0.880 |
| Alpha diversity     |             |       |         |       |
| OTU richness        | Bacteria    | 50.7% | 8.82 * 10^-6 | 0.7% | 0.994 |
| Simpson evenness    | Bacteria    | 54.0% | 1.57 * 10^-7 | 4.0% | 0.384 |
| Inverse Simpson     | Bacteria    | 57.3% | 1.45 * 10^-9 | 0.7% | 0.994 |
| OTU richness        | Fungi       | 28.0% | 0.931     | 4.7% | 0.235 |
| Simpson evenness    | Fungi       | 42.0% | 0.016     | 0.0% | 1.000 |
| Inverse Simpson     | Fungi       | 40.7% | 0.036     | 0.0% | 1.000 |
| Beta diversity      |             |       |         |       |
| Jaccard similarity  | Bacteria    | 100%  | 0.0001   | 100% | 0.0001 |
| Bray–Curtis dissimilarity | Bacteria | 100%  | 0.0001   | 100% | 0.0001 |
| Jaccard similarity  | Fungi       | 100%  | 0.0001   | 100% | 0.0001 |
| Bray–Curtis dissimilarity | Fungi | 100%  | 0.0001   | 100% | 0.0001 |

1LUT = Land-use type
2Reclass.: Reclassification success of leave-one-out tests.
3Cmic: carbon content based on chloroform fumigation extraction.

Table 3. Summary of OTU partitioning into core and indicative groups and subgroups. Core and indicative OTUs were defined at the site and the land-use type level (see Table 1 for definitions of OTU groups and subgroups). LUT = land-use type.

| OTU group | Subgroup | OTUs [N] | Abundance [%] | Correlation1 [rho] | Phyla [N] | Families [N] |
|-----------|----------|----------|---------------|---------------------|-----------|--------------|
| Bacteria  | Core     | ±         | 6 979          | 95.9                | ±          | 31           | 215          |
|           | indicative | ±         | ±              | ±                   | ±          | ±            | ±            |
| Fungi     | Core     | ±         | 2 802          | 93.2                | ±          | ±            | ±            |
|           | indicative | ±         | ±              | ±                   | ±          | ±            | ±            |

1Spearman correlation to entire community (Mantel test).
Figure 1. Separation of bacterial (A) and fungal (B) communities by land-use and correlated environmental factors. Three land-use types, i.e. arable land (blue), permanent grassland (green), and forest (brown), were sampled with 10 sites each. Per site, 15 samples were obtained with yearly triplicates during five years. Average communities for yearly replicates are shown (N = 150). Ordinations are based on canonical analyses of principal coordinates (CAP) constrained by land-use types. Axes show linear discriminants (LD). Arrows indicate significant correlations of communities to environmental factors, i.e. bulk density (BD), clay, silt, sand, pH, mean annual temperature (MAT), mean annual precipitation (MAP), ratio of C/N (C/N), total carbon (C) and nitrogen (N), organic carbon (OC), and elevation.

Table 4. Effects of environmental factors on bacterial (A) and fungal (B) communities as assessed by PERMANOVA. Factors are sorted by their position in the PERMANOVA model with environmental factors as covariates. Year and site were random factors with site being nested within land-use type. Factors below the line are categorical. Significance codes: *** P < 0.001, ** P < 0.01, * P < 0.05

| (A) Bacteria | Env. factor | Pseudo-F | √CV $^2$ | P-value | Env. factor | Pseudo-F | √CV $^2$ | P-value |
|--------------|-------------|-----------|----------|---------|-------------|-----------|----------|---------|
| pH | 18.6 | 0.25 | 0.0001 | *** | C/N-ratio | 5.5 | 0.19 | 0.0001 | *** |
| C/N-ratio | 5.4 | 0.15 | 0.0001 | *** | pH | 2.6 | 0.14 | 0.0001 | *** |
| MAP$^3$ | 2.2 | 0.07 | 0.0001 | *** | Elevation | 1.5 | 0.09 | 0.0086 | ** |
| Clay | 2.0 | 0.06 | 0.0001 | *** | Sand | 1.3 | 0.06 | 0.0330 | * |
| Elevation | 1.4 | 0.05 | 0.0119 | * | Clay | 1.2 | 0.06 | 0.1399 | |
| Corg | 1.5 | 0.07 | 0.0130 | * | MAT$^4$ | 1.2 | 0.06 | 0.1012 | |
| Sand | 1.3 | 0.06 | 0.0338 | * | Corg | 1.2 | 0.06 | 0.1226 | |
| MAT$^4$ | 1.2 | 0.04 | 0.1418 | * | Bulk density | 1.3 | 0.08 | 0.1661 | |
| Year | 4.2 | 0.05 | 0.0001 | *** | Year | 2.2 | 0.06 | 0.0001 | *** |
| LUT$^5$ | 2.9 | 0.23 | 0.0017 | ** | LUT$^5$ | 2.5 | 0.31 | 0.0001 | *** |
| Site | 19.4 | 0.31 | 0.0001 | *** | Site | 14.4 | 0.49 | 0.0001 | *** |
| LUT$^5$ x Year | 1.5 | 0.04 | 0.0001 | *** | LUT$^5$ x Year | 1.3 | 0.05 | 0.0001 | *** |
| Residuals | 0.15 | | | | Residuals | 0.28 | | |

1 Env. factor: environmental factor;
2 √CV: square root of component of variation, expressed as Bray–Curtis dissimilarity;
3 MAP: mean annual precipitation
4 MAT: mean annual temperature
5 LUT: Land-use type

Association of bacterial and fungal core OTUs to land-use types

The similarities of bacterial and fungal communities among land-use types were highest between arable and permanent grassland soils, while they were lowest between arable and forest soils (AG and AF in Fig. 2). The similarity between communities from forest and permanent grassland sites was higher for bacteria than for fungi, which was particularly striking, when relative abundances were considered as accounted for in Bray Curtis similarities (FG in Fig. 2C and D). To assess these differences in greater detail, the distribution of core taxa among the land-use types were analyzed using ternary plots, which depict the abundance of sc-OTUs in each land-use type and in all combinations (Fig. 3). The ternary plots clearly revealed different distributions of bacterial and fungal sc-OTUs among land-use types. On the one hand, bacterial sc-OTUs were distributed among the land-use types and all their combinations except for the combination of ‘arable land and forest’, for which only two lc-OTUs were detected (Fig. 3A). Eighty-seven bacterial sc-OTUs
were unclassified at the family-level, while these numbers were 78.0% for bacterial and 60.3% for fungal OTUs at the genus-level. In order to analyze associations of families to land-use types, we extracted sc-OTUs that were predominantly associated to a single or combinations of land-use types based on the ternary plot (Fig. 4A). The ten most abundant families in each of the seven areas specified in the ternary plot, i.e. triangles A, G, F, AG, GF, AF and AGF, were extracted. They covered in the selected areas 18.7% and 49.2% of the overall relative abundance of bacterial and fungal sc-OTUs, respectively, (Fig. 4B, dark grey area) and resulted in a list of 39 bacterial and 38 fungal families (Fig. 4C and D). Cluster analysis was used to group these families according to their distribution patterns in the land-use types, which yielded seven bacterial and five fungal clusters (Fig. 4C and D). All clusters composed of at least three families included families of several phyla with the exception of the fungal cluster II that was exclusively composed of families from the Ascomycota, but which were not closely related to each other (Fig. S3, Supporting Information). Furthermore, all phyla represented by at least three families were detected in at least two clusters with the exception of Myxococccota (Fig. 4). All three families from the Myxococcota, e.g. Birh141, Haliangiaceae and Sandaracinaceae, were included in bacterial cluster V and occurred in all three land-use types, but showed a preference for ‘arable land and grassland’ (Fig. 4C). More homogenous representations of land-use types within the clusters were found for fungi as compared to bacteria. Most strikingly, fungal cluster V, which was composed of families such as Myxotrichiaceae, Inocybaceae, and Russulaceae, occurred most strongly and almost exclusively in forest soils. Clusters predominantly associated to permanent grassland included only one bacterial family, the Ktedonobacteraceae (cluster IV, Fig. 4C), but eight fungal families, e.g. Mortierellaceae and Chaetothyriaceae (cluster IV, Fig. 4D). Within the clusters, also groupings with more resolved land-use type associations were revealed. For instance, within fungal cluster IV the fungal families Mortierellaceae, Clavariaceae and Herpotrichiellaceae were all most abundant in permanent grassland but revealed a complex occurrence pattern in many land-use types, while the fungal family Chaetothyriaceae was exclusively detected in permanent grassland soils. Similarly, within fungal cluster III, which was mainly associated to arable land, some families such as Lasiosphaeriaceae and Nectriaceae were also prominently detected in the combination ‘arable land and permanent grassland’ while the Bulleribasidiaceae, as an exemption in cluster III, were more abundant in the combination ‘arable land and permanent grassland’ but comparably abundant in ‘arable land’. For bacteria, such clear clustering was less pronounced. Cluster VI exclusively associated to ‘arable land’ but for instance in cluster VII only 11 of the 13 families were most abundant in forest soils. Within cluster VII, families such as Pedosphaeraceae or the candidate WD2101 soil group were also commonly detected in arable and permanent grassland soils. The strongest forest associations were observed for families Acidobacteriaceae Subgroup 1 as well as Acetobacteraceae, Methylocyclaceae, Acidothermaceae and Micropepsaceae. Therefore, stronger associations to land-use types or their combinations were detected for fungi as compared to bacteria. This was further supported by the number of families with their highest abundance in a single land-use type (A, G or F), which was lower for bacteria (20, Fig. 4C) as compared to fungi (30, Fig. 4D).

To detect families, which showed the strongest and most consistent associations to land-use types, we compared core and indicative OTUs. More specifically, we first selected OTUs, which were core and indicative of the same land-use type or land-use type combinations and aggregated these OTUs at the

**Figure 2.** Pairwise comparisons of bacterial (A, C) and fungal (B, D) communities composed of core OTUs for a site, i.e. OTUs that occurred in at least 12 of the 15 samples from a site. Boxplots showing Jaccard (A, B) and Bray-Curtis (C, D) similarities between two sites depending on their land-use type. The Jaccard similarity corresponds to the ratio of shared OTUs between two sites, while the Bray-Curtis similarity takes also the relative abundance of each OTU into account. Sites of three land-use types, i.e. arable land (A), grassland (G), and forest (F), were assessed in pairwise combinations of the same land-use type (AA, GG and FF) as well as between different land-use types (AG, FG and AF).

**Distribution of bacterial and fungal families among land-use types**

For taxonomic characterization of core communities, we focused on the family level, since the classified OTUs can be more reliably assigned at this level and since the number of unclassified OTUs increased at lower taxonomic levels. For instance, 50.7% of the bacterial and 47.1% of the fungal OTUs were unclassified at the family-level, while these numbers were
family-level. This yielded 304 bacterial (Table S7, Supporting Information) and 58 fungal OTUs (Table S8, Supporting Information). Then, we selected families, which included at least four (Bacteria) or two (Fungi) OTUs that were both core and indicative of the same land-uses (Table 5). This resulted in 16 bacterial and 9 fungal families (Table 5), which were also among the families described in Fig. 4, with the exception of bacterial candidate groups SC-I-84 and AKYH767, as well as the fungal family Phaeosphaeriaceae. Two bacterial families, Anaerolinaceae and Pyrinomonadaceae included arable core and indicative OTUs and a single bacterial family, Acidobacteriaceae Subgroup 1, included only forest core and indicative OTUs. No bacterial family included only OTUs that were core and indicative of permanent grassland soils. Among fungi Chaetomiaceae and Myxotrichaceae included only OTUs that were core and indicative of a single land-use type, i.e. arable land and forest, respectively. No fungal family included exclusively OTUs that were core and indicative of permanent grassland soils. Furthermore, no bacterial and fungal OTUs were core and indicative of the combination ‘arable land and forest’ and only bacterial but no fungal families included OTUs that were core and indicative of ‘permanent grassland and forest’. The lack of such OTUs is consistent with the few sc-OTUs detected in the corresponding areas of the ternary plots (Fig. 3), as well as with low similarities of bacterial and fungal communities among arable and forest sites, and equally low similarities among fungal communities of permanent grassland and forest sites (Fig. 2).

DISCUSSION
Land-use-specificity of soil bacterial and fungal communities

Soil bacterial and fungal communities were surveyed during 5 years at 30 sites of the Swiss Soil Monitoring Network including three different land-use types, i.e. arable land, permanent grassland, and forest. This revealed communities that were highly specific to land-use types and sites, and which were stable over five years. A detailed analysis on the long-term stability of these communities early in the vegetation period without considering intra-annual variability has already been described (Gschwend et al. 2021b). Here, we focused on the environmental drivers that shape this land-use- and site-specificity of soil bacterial and fungal communities, as well as on their taxonomic compositions.

Each land-use type was characterized by differences in the combinations of soil properties, management, and vegetation (Table S1, Supporting Information). In arable soils, pH and bulk density were increased, while carbon contents were equal or lower than in permanent grassland and forest soils. Furthermore, management of arable soils included crop rotations, tillage (except one site), mineral and organic fertilization, as well as plant protection, which are known to influence soil bacterial and fungal communities (Hartmann et al. 2015; Rivera-Becerril et al. 2017; Peralta et al. 2018). Microbial biomass was significantly reduced in arable soils as compared to permanent grassland and forest soils (Table S1, Supporting Information), which confirms earlier findings (Dequiedt et al. 2011). Bacterial communities in arable soils were characterized by families such as Anaerolineaceae, Pyrinomonadaceae and Gemmatimonadaceae. Anaerolineaceae are widely distributed in soils, and particularly prevalent in low-oxygen environments, e.g. in compacted soils (Hartmann et al. 2014) or paddy fields (Jiao et al. 2019). As they may act as indicators for soil oxygen depletion (Gschwend et al. 2020) and as they have been recently detected to be associated with soil compaction in arable fields (Longepierre et al. 2021), their high abundance in arable soils may be a sign of soil compaction in arable land due to common management practices with heavy machinery. The families Halangiaceae, Sandaracinaceae and Btii41 revealed similar distributions among land-use types with the highest abundance in arable and grassland soils (Fig. 4C). This is in agreement with findings by Karimi et al. (2018), who...
Figure 4. Distribution of most abundant bacterial and fungal families among land-use types. Based on the ternary plots (Fig. 3) site-core OTUs (sc-OTUs) were selected from seven areas (A) corresponding to sc-OTUs with at least 80% of their abundance in a single land-use type (A, G, F), with at least 40% in each of two land-use type (AG, GF, AF), or with at least 20% in each land-use types (AGF). The proportions of relative abundances covered by the selected sc-OTUs, and their assignment at the family level, is shown in panel (B). Panels (C) and (D) show the relative abundances of the ten most abundant bacterial (C) and fungal (D) families of each area of the ternary plots. Light blue indicates low, dark blue middle, and brown high relative abundances. White areas represent absences of families in an area of the ternary plot. The area in which a family has its highest abundance is indicated by the following color code (Maximal abund.): blue (A), green (G), brown (F), yellow (AG), light green (GF), pink (AF), and orange (AGF). Highest abundances in a single land-use type are indicated by black hatching. Dendrograms show clustering of normalized relative abundances of families in the land-use types and their combinations using average clustering (UPGMA). Red boxes highlight clusters of families with similar distributions among land-use types.
detected the genus *Haliangium* in all samples and who found its highest relative abundance in arable and grassland soils. All three families are classified within the candidate phylum Mycococcota, which regroups many predatory bacterial species (Waite et al. 2020), and represented the only phylum in which all selected families revealed similar habitat preferences in our survey (Fig. S3a, Supporting Information). Fungal communities in arable soils were for instance characterized by Lasiosphaeriaceae, Plectosphaerellaceae, Chaetomiaceae and Mrakiaceae. With the exception of the basidiomycetous yeasts Mrakiaceae and Cystofilobasidiaceae (Liu et al. 2015), fungal families associated to arable soils also occurred in permanent grassland soils (Fig. 4). For instance, Plectosphaerellaceae that include important soil-borne plant pathogens such as Verticillium (Giraldo and Crous 2019) had two lc-OTUs that were also indicative for arable land, as well as one that was indicative for ‘arable land and permanent grassland’ (Table 5). In these cases, OTUs assigned to the same family have distinct land-use type associations, which may for instance be driven by species-specific host plant preferences (Klosterman et al. 2009).

Permanent grassland soils were characterized by soil property values, which lay between those of arable and forest soils (Table S1, Supporting Information). Their management included fertilization, mowing, and grazing, which may change soil bacterial and fungal community structures (Kaiser et al. 2020; Gilmullina et al. 2020). A single bacterial family, the Ktedonobacteraceae (phylum Chloroflexi) had their highest abundance in arable and permanent grassland soils (Fig. 4C). Ktedonobacter racemifer (Ktedonobacteraceae) was isolated from soil of a black locust forest in Italy (Cavaletti et al. 2006). Metabarcoding of bacterial communities from 2173 soil samples across France revealed sequences assigned to Ktedonobacter in 80% of all samples (Karimi et al. 2018). Families that characterized fungal communities in permanent grasslands included for instance the grassland-specific Chaetothyriaceae (Fig. 4D). Chaetothyriaceae include mainly epiphytic species living on plants (Quan et al. 2020) suggesting that their distribution may depend on host

### Table 5. Number of OTUs, which were indicative (IndVal >0.8, P < 0.05) and core for the same land-use types from selected bacterial and fungal families. Families were selected if at least four (bacteria) or two (fungi) OTUs were indicative and core for the same land-use type or land-use type combination. All families are shown in Tables S7 (Bacteria) and S8 (Fungi), Supporting Information. Associations of families to fungal families. Families were selected if at least four (bacteria) or two (fungi) OTUs were indicative and core for the same land-use type or land-use type combination. All families are shown in Tables S7 (Bacteria) and S8 (Fungi), Supporting Information. Associations of families to fungal families. Families were selected if at least four (bacteria) or two (fungi) OTUs were indicative and core for the same land-use type or land-use type combination.

| Family             | A     | AG    | G     | AF    | GF    | F     | Cluster | Main abund. 2 |
|--------------------|-------|-------|-------|-------|-------|-------|---------|----------------|
| **Bacteria**       |       |       |       |       |       |       |         |                |
| Chthoniobacteraceae| 1(1)  | 4(12) | 0(1)  | 0(0)  | 2(5)  | 3(4)  | III     | GF             |
| Pirellulaceae      | 4(5)  | 1(8)  | 0(0)  | 0(0)  | 0(2)  | 0(0)  | V       | A*            |
| Chitinophagaceae   | 4(10) | 3(12) | 2(2)  | 0(0)  | 1(4)  | 1(3)  | V       | GF            |
| Gemmatimonadaceae  | 5(9)  | 3(5)  | 0(0)  | 0(0)  | 0(2)  | 1(1)  | VI      | A*            |
| Anaerolineaceae    | 4(4)  | 0(3)  | 0(0)  | 0(0)  | 0(1)  | 0(0)  | VI      | A*            |
| Pyrinomonadaceae   | 4(4)  | 0(0)  | 0(0)  | 0(0)  | 0(1)  | 0(0)  | VI      | A*            |
| Burkholderiaceae   | 1(2)  | 4(7)  | 0(0)  | 0(0)  | 0(1)  | 3(4)  | VII     | AGF           |
| Pedosphaeraceae    | 4(5)  | 7(11) | 3(3)  | 0(0)  | 1(12) | 3(5)  | VII     | F             |
| WD2101 soil group  | 2(5)  | 9(18) | 1(2)  | 0(0)  | 1(4)  | 1(5)  | VII     | F             |
| Acidobacteriaceae Sg 1 | 0(0) | 0(0)  | 0(0)  | 0(0)  | 0(1)  | 6(8)  | VII     | F*            |
| Acetobacteraceae   | 0(0)  | 0(0)  | 0(0)  | 0(0)  | 2(2)  | 4(6)  | VII     | F*            |
| Caulobacteraceae   | 0(0)  | 1(1)  | 0(0)  | 0(0)  | 0(0)  | 3(5)  | VII     | F*            |
| Acidothermaceae    | 0(0)  | 0(0)  | 0(0)  | 0(0)  | 2(3)  | 2(4)  | VII     | F*            |
| Solibacteraceae Sg 3 | 0(0) | 0(2)  | 0(0)  | 0(0)  | 3(12) | 1(3)  | VII     | GF            |
| SC-1-84            | 1(1)  | 2(4)  | 2(2)  | 0(0)  | 0(3)  | 0(0)  |          |                |
| Others (incl. unclassified) | 47(75) | 53(195) | 12(21) | 0(4)  | 28(110) | 17(41) |          |                |
| **Fungi**          |       |       |       |       |       |       |         |                |
| Pseudoeurotiaceae  | 0(0)  | 1(1)  | 0(0)  | 0(0)  | 0(0)  | 1(1)  | II      | AG*           |
| Lasiopsphaeraceae   | 3(4)  | 2(7)  | 0(1)  | 0(0)  | 0(0)  | 0(0)  | III     | A*            |
| Plectosphaerellaceae| 2(3)  | 1(1)  | 0(0)  | 0(0)  | 0(0)  | 0(0)  | III     | A*            |
| Chaetomiaceae      | 2(2)  | 0(0)  | 0(1)  | 0(0)  | 0(0)  | 0(0)  | III     | A*            |
| Nectriaceae        | 1(1)  | 3(9)  | 0(0)  | 0(0)  | 0(1)  | 0(0)  | III     | A*            |
| Helotiaceae        | 0(2)  | 3(4)  | 0(0)  | 0(0)  | 0(0)  | 0(1)  | III     | A*            |
| Mortierellaceae    | 0(0)  | 2(5)  | 1(1)  | 0(0)  | 0(0)  | 1(1)  | IV      | G             |
| Myxotrichaceae     | 0(0)  | 0(0)  | 0(0)  | 0(0)  | 0(0)  | 2(4)  | V       | F*            |
| Phaeosphaeraceae   | 1(1)  | 1(1)  | 0(0)  | 0(0)  | 0(0)  | 0(0)  |          |                |
| Others (incl. unclassified) | 7(35) | 12(53) | 5(13) | 0(2)  | 0(0)  | 7(20) |          |                |
| **All**            | 16(47) | 25(79) | 6(16) | 0(2)  | 0(1)  | 11(26) |          |                |

1 indicative and lc-OTUs: OTUs, which are indicative and core of the same land-use type(s), note that OTUs cannot be indicative of all sites, i.e. the combination AGF;
2Main abund.: Main abundance in arable land (A), permanent grassland (G), forest (F) or their combinations, according to Fig. 4.
plants. However, in a survey of switchgrass-associated fungal communities, OTUs attributed to this family have also been detected associated to the switchgrass roots and adjacent soils, but not on plant leaves (Lee and Hawkes 2020), indicating that Chaetothyriaceae also include soil fungi.

Forest soils were characterized by relatively high contents of carbon, higher C/N-ratios, and lower soil pH as compared to the arable soils (Table S1, Supporting Information). Bacterial families associated to forest soils included Acidobacteria Subgroup 1, Acetobacteraceae, Acidothermaceae, as well as the more widely distributed WD2101 soil group, and Pedosphaeraceae (Fig. 4C, Table 5). Acidobacteria Subgroup 1 have been repeatedly reported to negatively correlate with soil pH (Kielak et al. 2016) and revealed increased abundances in soils with a pH below 6.5 (Jones et al. 2009). Acetobacteraceae have also been reported to strongly and negatively correlate with soil pH and to have higher abundances in forest as compared to grassland soils (Nacke et al. 2011). Therefore, soil pH, which is well known to be a major driver of soil bacterial communities (e.g. Lauber et al. 2009; Karimi et al. 2018), was the main factor determining forest associated bacterial taxa. Fungal communities in forest soils were mainly composed of ectomycorrhizal families such as Russulaceae, Inocybaceae, and Clavulinaceae, which is in agreement with previous findings (e.g. Frey et al. 2021). Thirteen fungal families were strongly associated to forest (Cluster V, Fig. 4D), but only one of these, the Myxotrichaceae, included indicative OTUs of forest soils (Table S8, Supporting Information). This is likely explained by the different forest ecosystems including deciduous, mixed and coniferous forests that have been sampled. As ectomycorrhizal fungi depend on their host tree species (Bahnmann et al. 2018), none of these families occurred at eight or more forest sites and were thus not generally indicative for forest soils. Myxotrichaceae included for instance Oidiodendron spp., which were repeatedly detected among the abundant soil fungal in metabarcoding surveys of Swiss forest soils (Hartmann et al. 2017; Frey et al. 2020), and which are common saprobes in acid soils but some of which also form ericoid mycorrhiza (Rice and Currah 2005). Therefore, their widespread and indicative distribution in various forest soil ecosystem may relate to a dependence on understory vegetation, or on the general preference for acidic soils.

Similarities of soil bacterial and fungal communities among land-use types
The similarities among soil bacterial communities from different land-use types were lowest for the combination of arable land and forest (Fig. 2), which was also the only land-use type combination for which no bacterial lc-OTU was indicative (Table 5). Similarities between soil bacterial communities from arable and permanent grassland soils corresponded to values observed between permanent grassland and forest soils (Fig. 2). This suggests that soil bacterial communities represented a sequential order following the soil properties and the land-use intensity from arable land, to permanent grassland and forest. For fungi, similarities from communities of permanent grassland and forest soils were equally low as among communities of arable and forest soils (Fig. 2). Furthermore, no fungal OTUs was found that was indicative and land-use core for the combination ‘permanent grassland and forest’ or the combination ‘arable land and forest’ (Table 5). Therefore, soil fungal, unlike bacterial, communities revealed little overlap (Bray–Curtis < 0.10, Fig. 2) between permanent grassland and forest soils. Considering dissimilarities among communities as proxies for the transfer of soil microorganisms among sites allows describing the structure of their metacommunities (Beck et al. 2019; Wisnosi and Lennon 2021). In this view, soil bacterial communities of arable, permanent grassland, and forest soils formed a single metacommunity, which was characterized by a continuous change from arable land, to permanent grassland and forest. Soil fungal communities, however, formed two metacommunities, one created by fungal communities of arable and permanent grassland soils and the other by fungal communities of forest soils.

The distinct structures of soil bacterial and fungal metacommunities can be explained by different factors influencing their community assembly. On the one hand, bacterial communities were more strongly structured by soil properties and climatic factors as compared to soil fungal communities (Table 4). On the other hand, soil fungal communities were more strongly structured by vegetation as compared to soil bacterial communities. For instance, acidophilic bacterial families predominantly occurred in forest soils (Fig. 4C), while ectomycorrhizal fungal families dominated soil fungal communities in forest soils (Fig. 4D). Confirming our results Frey et al. (2021) reported stronger effects of tree species on fungal as compared to bacterial community structures. Stronger vegetation effects on soil fungal as compared to bacterial communities were also revealed in the other land-use types, as crops had a stronger effect on soil fungal as compared to bacterial community structures (Tables S3 and S4, Supporting Information), which is in agreement with the findings of Ai et al. (2018). Stronger legacy effects of different grassland mixtures on soil fungal as compared to soil bacterial communities have been described in a grassland field experiment (Fox et al. 2020), which further supports the stronger impact of plants on soil fungal as compared to bacterial communities.

Potential use of sc-OTUs to provide a temporally stable, cultivation-independent reference list of dominant taxa
Site core OTUs accounted for 38.5% of bacterial and 33.1% of fungal OTUs, but covered 95.9% and 93.2% of relative abundance (Table 3). As sc-OTUs occurred in at least four of the five years, the large majority of retrieved sequences, could be attributed to temporally stable OTUs. These sc-OTUs not only were temporally stable but also included 95% of bacterial and 90% of fungal indicative OTUs (Fig. S2, Supporting Information) and were representative of the diversities of entire communities (Table S2, Supporting Information). Furthermore, OTUs largely restricted to a single site were classified as site-indicative OTUs, which was the case of 14.5% of bacterial and 44.8% of fungal sc-OTUs. Therefore, the majority of sc-OTUs were consistently detectable at several sites revealing their potential to act as reference set for the analysis of soil microbial diversity in soil habitats similar to those assessed in this study. Such reference sets are of particular interest for predictive modelling of soil bacterial and fungal diversity and distribution (Jiao et al. 2019), and may also be used as reference values for long-term soil quality monitoring (Gschwend et al. 2021b), although their intra-annual variability remains to be assessed. Furthermore, they may constitute a set of taxa that can be screened to find robust biodeicators for specific soil functions, such as plant pathogen suppression (Trivedi et al. 2017). Currently, long-term monitoring systems of soil biodiversity are largely lacking (Leeuwen et al. 2017; Guerra et al. 2020), which is particularly concerning given the ongoing environmental changes and the central role of soil biodiversity.
for global ecosystem processes. Finally, sc-OTUs provide support to establish lists of the most characteristic soil microorganisms, for which cultivation strategies or whole-genome sequencing are particularly valuable (Carini 2019). Currently, still too few dominating soil bacterial and fungal taxa have cultured representatives or available genome sequences, which would enable more detailed insight into their functions in the ecosystem (Delgado-Baquerizo et al. 2018; Egidi et al. 2019; Steen et al. 2019).

**CONCLUSIONS**

While microbial biomass and alpha-diversity measures at thirty long-term monitoring sites revealed only few differences among land-use types and sites, community compositions (Jaccard similarity) and structures (Bray–Curtis dissimilarity) yielded characteristic descriptors for each land-use type and site. Therefore, resolution obtained by metabarcoding were necessary to accurately describe soil bacterial and fungal communities. Temporally stable core OTUs accounted for 95.9% of bacterial and 93.2% of fungal sequences. These core OTUs were representative of entire communities and showed responses to distinct habitats. In total 4184 indicative bacterial and 1968 indicative fungal OTUs, of which 95% and 90% were also temporally stable core OTUs, were identified. These yield promising targets for the development of microbial indicators for robust soil quality analyses. Bacterial and fungal families were identified that revealed strong associations to one or more land-use types. In general, fungal families revealed stronger associations to land-use types, which may be explained by the stronger influences of vegetation on fungi as compared to bacteria, whereas bacteria were more strongly correlated with soil properties. Consequently, metacommunities of soil bacteria and fungi were differently structured. On the one hand, bacterial communities represented a sequential order following soil properties and land-use intensity from arable land, to permanent grassland and forest. On the other hand, fungal communities of forest sites showed only minor similarities to those from arable land and permanent grassland sites. The robustly assessed and temporally stable core OTUs may serve as references for future surveys of soil bacterial and fungal diversity. This may facilitate long-term soil quality monitoring by detecting disturbances of the characteristic habitat associated core communities, and it may also enable the development of predictive modelling for metabarcoding based soil quality analyses.

**SUPPLEMENTARY DATA**

Supplementary data are available at FEMS online.

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**REFERENCES**

Ai C, Zhang S, Zhang X et al. Distinct responses of soil bacterial and fungal communities to changes in fertilization regime and crop rotation. Geoderma 2018;319:156–66.

Anderson M, Gorley RN, Clarke K. PERMANOVA+ for primer: Guide to software and statistical methods. Plymouth: Primer-E, 2008.

Anderson MJ, Ellingsen KE, McArdle BH. Multivariate dispersion as a measure of beta diversity. Ecology Letters 2006;9:683–93.

Anderson MJ, Willis TJ. Canonical analysis of principal coordinates: a useful method of constrained ordination for ecology. Ecology 2003;84:511–25.

Anderson MJ. A new method for non-parametric multivariate analysis of variance. Austral Ecol 2001;26:32–46.

Babin D, Deubel A, Jacquierd S et al. Impact of long-term agricultural management practices on soil prokaryotic communities. Soil Biol Biochem 2019;129:17–28.

Bahnmann B, Mašinová T, Halvorsen R et al. Effects of oak, beech and spruce on the distribution and community structure of fungi in litter and soils across a temperate forest. Soil Biol Biochem 2018;119:162–73.

Bahram M, Hildebrand F, Foslund SK et al. Structure and function of the global topsoil microbiome. Nature 2018;560:233–7.

Banerjee S, Walder F, Bühli I et al. Agricultural intensification reduces microbial network complexity and the abundance of keystone taxa in roots. ISME Journ 2019;13:1722–36.

Bardgett RD, van der Putten WH. Belowground biodiversity and ecosystem functioning. Nature 2014;515:505.

Beck S, Anderson IC, Drigo B et al. A soil fungal metacommunity perspective reveals stronger and more localised interactions above the tree line of an alpine/subalpine ecotone. Soil Biol Biochem 2019;135:1–9.

Benjamini Y, Hochberg Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. Journal of the Royal Statistical Society: Series B (Methodological) 1995;57:289–300.

Bürgmann B, Mäschinová T, Halvorsen R et al. A strategy for optimizing quality and quantity of DNA extracted from soil. J Microbiol Methods 2001;45:7–20.

Carini P. A “Cultural” renaissance: genomics breathes new life into an old craft. mSystems 2019;4:e00092–19.

Cavaletti L, Monciardini P, Bamonte R et al. New lineage of filamentous, spore-forming, gram-positive bacteria from soil. Appl Environ Microbiol 2006;72:4360–9.

Clarke KR, Warwick RM. Change in marine communities: an approach to statistical analysis and interpretation, 2nd edition. PRIMER-E: Plymouth, 2001.

Costa OYA, Raajmakers JM, Kuramae EE. Microbial extracellular polymeric substances: ecological function and impact on soil aggregation. Frontiers in Microbiology 2018;9.

Cui H, Sun W, Delgado-Baquerizo M et al. The effects of mowing and multi-level nitrogen fertilization on soil bacterial and fungal communities in a semiarid grassland are year-dependent. Soil Biol Biochem 2020;151:108040.

De Cáceres M, Legendre P. Associations between species and groups of sites: indices and statistical inference. Ecology 2009;90:3566–74.

Degruene F, Theodorakopoulos N, Colinet G et al. Temporal dynamics of soil microbial communities below the seedbed under two contrasting tillage regimes. Frontiers in Microbiology 2017;8.

Delgado-Baquerizo M, Oliverio AM, Brewer TE et al. A global atlas of the dominant bacteria found in soil. Science 2018;359:320–5.
Delgado-Baquerizo M. Obscure soil microbes and where to find them. The ISME Journal 2019;13:2120–4.
Dequeidt S, Saby NPA, Lelièvre M et al. Biogeographical patterns of soil molecular microbial biomass as influenced by soil characteristics and management. Global Ecol Biogeogr 2011;20:641–52.
Edgar RC. Search and clustering orders of magnitude faster than BLAST. Bioinformatics 2010;26:2460–1.
Egidi E, Delgado-Baquerizo M, Plett JM et al. A few ascomycota taxa dominate soil fungal communities worldwide. Nat Commun 2019;10:2369.
Fox A, Lüscher A, Widmer F. Plant species identity drives soil microbial community structures that persist under a following crop. Ecology and Evolution 2020;10:8652–68.
Frey B, Carnol M, Dharmarajah A et al. Only minor changes in the soil microbiome of a sub-alpine forest after 20 years of moderately increased nitrogen loads. Frontiers in Forests and Global Change 2020;3.
Frey B, Rime T, Phillips M et al. Microbial diversity in european alpine permafrost and active layers. FEMS Microbiol Ecol 2016;92.
Frey B, Walthert L, Perez-Mon C et al. Deep soil layers of drought-exposed forests harbor poorly known bacterial and fungal communities. Frontiers in Microbiology 2021;12:1061.
Gilmullina A, Rumpel C, Blagodatskaya E et al. Management of grasslands by mowing versus grazing – impacts on soil organic matter quality and microbial functioning. Applied Soil Ecology 2020;156:103701.
Giraldo A, Crous PW. Inside plectosphaerellaceae. Studies in Mycology 2019;92:227–86.
Glassman SI, Martiny JBH. Broadscale ecological patterns are robust to use of exact sequence variants versus operational taxonomic units. mSphere 2018;3:e00148–18.
Glassman SI, Wang JJ, Bruns TD. Environmental filtering by pH and soil nutrients drives community assembly in fungi at fine spatial scales. Mol Ecol 2017;26:6960–73.
Griffiths RI, Thomson BC, James P et al. The bacterial biogeography of british soils. Environ Microbiol 2011;13:1642–54.
Griffiths RI, Thomson BC, Plassart P et al. Mapping and validating predictions of soil bacterial biodiversity using european and national scale datasets. Applied Soil Ecology 2016;97:61–8.
Gschwend F, Aregger K, Gramlich A et al. Periodic waterlogging transforms the soil microbiome of a sub-alpine forest after 20 years of moderately increased nitrogen loads. Frontiers in Forests and Global Change 2020;3.
Girasoli A, Crous PW. Inside plectosphaerellaceae. Studies in Mycology 2019;92:227–86.
Gilmullina A, Rumpel C, Blagodatskaya E et al. Management of grasslands by mowing versus grazing – impacts on soil organic matter quality and microbial functioning. Applied Soil Ecology 2020;156:103701.
Giraldo A, Crous PW. Inside plectosphaerellaceae. Studies in Mycology 2019;92:227–86.
Gilmullina A, Rumpel C, Blagodatskaya E et al. Management of grasslands by mowing versus grazing – impacts on soil organic matter quality and microbial functioning. Applied Soil Ecology 2020;156:103701.
Giraldo A, Crous PW. Inside plectosphaerellaceae. Studies in Mycology 2019;92:227–86.
Guerra CA, Heintz-Buschart A, Sikorski J et al. Blind spots in global soil biodiversity and ecosystem function research. Nat Commun 2020;11:3870.
Hallin S, Philippot L, Löffler FE et al. Genomics and ecology of novel N2O-reducing microorganisms. Trends Microbiol 2018;26:43–55.
Hamilton NE, Ferry M. ggtern: ternary diagrams using ggplot2. Journal of Statistical Software 2018;87:1–17.
Hartmann M, Brunner I, Hagedorn F et al. A decade of irrigation transforms the soil microbiome of a semi-arid pine forest. Mol Ecol 2017;26:1190–206.
Hartmann M, Frey B, Kölliker R et al. Semi-automated genetic analyses of soil microbial communities: comparison of T-RFLP and RISA based on descriptive and discriminative statistical approaches. J Microbiol Methods 2005;61:349–60.
Hartmann M, Frey B, Mayer J et al. Distinct soil microbial diversity under long-term organic and conventional farming. The ISME Journal 2015;9:1177–94.
Hartmann M, Niklaus PA, Zimmermann S et al. Resistance and resilience of the forest soil microbiome to logging-associated compaction. The ISME Journal 2014;8:226.
Hemkemeyer M, Christensen BT, Tebbe CC et al. Taxon-specific fungal preference for distinct soil particle size fractions. Eur J Soil Biol 2019;94:103103.
Hemkemeyer M, Dohrmann AB, Christensen BT et al. Bacterial preferences for specific soil particle size fractions revealed by community analyses. Frontiers in Microbiology 2018;9.
Jiao S, Xu Y, Zhang J et al. Core microbiota in agricultural soils and their potential associations with nutrient cycling. mSystems 2019;4:e00313–18.
Joergensen RG. The fumigation-extraction method to estimate soil microbial biomass: calibration of the kEC value. Soil Biol Biochem 1996;28:25–31.
Jones RT, Robeson MS, Lauber CL et al. A comprehensive survey of soil acidobacterial diversity using pyrosequencing and clone library analyses. The ISME Journal 2009;3:442–53.
Kaiser K, Wemheuer B, Korolkov V et al. Driving forces of soil bacterial community structure, diversity, and function in temperate grasslands and forests. Sci Rep 2016;6:33696.
Karimi B, Terrat S, Dequiedt S et al. Biogeography of soil bacteria and archaea across france. Sci Adv 2018;4:eaa1808.
Kielak AM, Barreto CC, Kowalchuk GA et al. The ecology of acido-bacteria: moving beyond genes and genomes. Frontiers in Microbiology 2016;7.
Kindt R, Coe R. Tree diversity analysis: a manual and software for common statistical methods for ecological and biodiversity studies. Nairobi: World Agroforestry Centre, 2005.
Klosterman SJ, Atallah ZK, Vallad GE et al. Diversity, pathogenicity, and management of verticillium species. Annu Rev Phytopathol 2009;47:39–62.
Kuhn M. Building predictive models in r using the caret package. Journal of statistical software 2008;28:1–26.
Lammel DR, Barth G, Ovaskainen O et al. Direct and indirect effects of a pH gradient bring insights into the mechanisms driving prokaryotic community structures. Microbiome 2018;6:106.
Lauber CL, Hamady M, Knight R et al. Pyrosequencing-based assessment of soil pH as a predictor of soil bacterial community structure at the continental scale. Appl Environ Microbiol 2009;75:5111–20.
Lazzaro A, Hartmann M, Blaser P et al. Bacterial community structure and activity in different Cd-treated forest soils. FEMS Microbiol Ecol 2006;58:278–92.
Lee MR, Hawkes CV. Plant and soil drivers of whole-plant microbiomes: variation in switchgrass fungi from coastal to mountain sites. Phytobiomes Journal 2020, DOI 10.1094/PHIOMES-07-20-0056-FI.
LeeuwENP, Saby NPA, Jones A et al. Gap assessment in current soil monitoring networks across europe for measuring soil functions. Environ Res Lett 2017;12:124007.

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Leff JW, Bardgett RD, Wilkinson A et al. Predicting the structure of soil communities from plant community taxonomy, phylogeny, and traits. The ISME journal 2018;12:1794–805.

Liu XZ, Wang QM, Göker M et al. Towards an integrated phylogenetic classification of the tremellomycetes. Studies in Mycology 2015;81:85–147.

Longepierre M, Widmer F, Keller T et al. Limited resilience of the soil microbiome to mechanical compaction within four growing seasons of agricultural management. ISME Communications 2021;1:1–13.

Mayerhofer J, Eckard S, Hartmann M et al. Assessing effects of the entomopathogenic fungus metarhizium brunneum on soil microbial communities in agriotes spp. biological pest control. FEMS Microbiol Ecol 2017;93.

Mayerhofer J, Wächter D, Calanca P et al. Environmental and anthropogenic factors shape major bacterial community types across the complex mountain landscape of switzerland. Frontiers in microbiology 2021;12:500.

McArdle BH, Anderson MJ. Fitting multivariate models to community data: a comment on distance-based redundancy analysis. Ecology 2001;82:290–7.

Nacke H, Thürmer A, Wollherr D et al. Pyrosequencing-based assessment of bacterial community structure along different management types in german forest and grassland soils. PLoS One 2011;6:e17000.

Nilsson RH, Larsson K-H, Taylor AFS et al. The UNITE database for molecular identification of fungi: handling dark taxa and parallel taxonomic classifications. Nucleic Acids Res 2018;47:D259–D64.

Peralta AL, Sun Y, McDaniel MD et al. Crop rotational diversity increases disease suppressive capacity of soil microorganisms. Ecosphere 2018;9:e02235.

Piazza G, Ercoli L, Nuti M et al. Interaction between conservation tillage and nitrogen gertilization shapes prokaryotic and fungal diversity at different soil depths: evidence from a 23-year field experiment in the mediterranean area. Frontiers in microbiology 2019;10.

Quan Y, Muggia L, Moreno LF et al. A re-evaluation of the chaetothyriales using criteria of comparative biology. Fungal Diversity 2020;103:47–85.

Quast C, Pruesse E, Yilmaz P et al. The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. Nucleic Acids Res 2012;41:D590–D6.

R Core Team. R: A language and environment for statistical computing [Computer software manual]. Vienna, Austria, 2016. https://www.R-project.org/.

Rice AV, Currah RS. Oidiodendron: a survey of the named species and related anamorphs of myxotrichum. Studies in Mycology 2005;53:83–120.

Rivera-Becerril F, van Tuinen D, Chatagner O et al. Impact of a pesticide cocktail (fenhexamid, folpel, deltamethrin) on the abundance of glomeromycota in two agricultural soils. Sci Total Environ 2017;577:84–93.

RStudio. RStudio: integrated development for R. Boston, MA: RStudio, Inc, 2015; 42:14. http://www.rstudio.com.

Schloss PD, Westcott SL, Ryabin T et al. Introducing mothur: open-source, platform-independent, community-supported software for describing and comparing microbial communities. Appl Environ Microbiol 2009;75:7537–41.

Steen AD, Crits-Christoph A, Carini P et al. High proportions of bacteria and archaea across most biomes remain uncultured. The ISME Journal 2019;13:3126–30.

Tedersoo L, Bahram M, Põlme S et al. Global diversity and geography of soil fungi. Science 2014;346:1256688.

Tedersoo L, Bahram M, Puusepp R et al. Novel soil-inhabiting clades fill gaps in the fungal tree of life. Microbiome 2017;5:42.

Trivedi P, Delgado-Baquerizo M, Trivedi C et al. Keystone microbial taxa regulate the invasion of a fungal pathogen in agro-ecosystems. Soil Biology and Biochemistry 2017;111:10–4.

Vance ED, Brookes PC, Jenkinson DS. An extraction method for measuring soil microbial biomass C. Soil Biol Biochem 1987;19:703–7.

Větrovský T, Kohout P, Kopecký M et al. A meta-analysis of global fungal distribution reveals climate-driven patterns. Nat Commun 2019;10:5142.

Waite DW, Chuvochina M, Pelikan et al. Proposal to reclassify the proteobacterial classes deltaproteobacteria and oligoflexia, and the phylum thermodesulfobacteria into four phyla reflecting major functional capabilities. Int J Syst Evol Microbiol 2020;70:5972–6016.

Walsch CM, Gebert MJ, Delgado-Baquerizo M et al. A global survey of mycobacterial diversity in soil. Appl Environ Microbiol 2019;85:e01180–19.

Wisnioski NI, Lennon JT. Microbial community assembly in a multi-layer dendritic metacommunity. Oecologia 2021;195:13–24.