The relationship between land cover and microbial community composition in European lakes

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Abstract

Microbes such as bacteria, archaea, and protists are essential for element cycling and ecosystem functioning, but many questions central to the understanding of the role of microbes in ecology are still open. Here, we analyze the relationship between lake microbiomes and the land cover surrounding the lakes. By applying machine learning methods, we quantify the covariance between land cover categories and the microbial community composition recorded in the largest amplicon sequencing dataset of European lakes available to date. We identify microbial bioindicators for these land cover categories. Combining land cover and physico-chemical bioindicators identified from the same amplicon sequencing dataset, we develop two novel similarity metrics that facilitate insights into the ecology of the lake microbiome. We show that the bioindicator network, i.e., the graph linking OTUs indicative of the same environmental parameters, corresponds to microbial co-occurrence patterns. Taken together, we demonstrate the strength
of machine learning approaches to identify correlations between microbial diversity and environmental 

factors, potentially opening new approaches to integrate environmental molecular diversity into monitor-

ing and water quality assessments.

**Keywords:** microbial ecology, machine learning, bioindicators, lake ecology

## 1 Introduction

Ecosystems are governed by processes at very different scales, ranging from the level of chemical reactions that 

shape the metabolism of single organisms to processes at the landscape level \[\text{[Allen et al., 2014; Dobzhansky,} \]

\[\text{1966; Odum, 1997]. Furthermore, the different scales are interlinked. For example, lakes and other freshwater} \]

habitats accumulate water from their catchment, and with it, nutrients, stressors, and pollutants. Because 

a lake’s water quality depends on its watershed’s ability to collect and purify water, lakes can be considered 

sentinels of environmental change of the landscape they are part of \[\text{[O’Neill et al., 1997; Williamson et al.,} \]

\[\text{2008]. Microbes, in turn, play an essential role in the functioning and the stability of ecosystems and can} \]

be regarded as “first responders” to environmental change \[\text{[Colwell, 1997; Docherty and Gutknecht, 2011;} \]

\[\text{Webster et al., 2018]. Indeed, because of their sensitivity to changes in the physico-chemical makeup of} \]

their environment, microorganisms are increasingly being used as bioindicators for ecosystem integrity in 

monitoring schemes \[\text{[Birk et al., 2012; Cordier et al., 2018; Hering et al., 2010; Kermarrec et al., 2014].} \]

While anthropogenic land cover change is one of the drivers of ecosystem quality decline \[\text{[Intergovern-

mental Platform on Biodiversity and Ecosystem Services (IPBES), 2019; Song et al., 2018; Steinbauer et al.,} \]

\[\text{2018]}, it is almost impossible to study its effects on freshwater ecosystems in well-controlled, experimen-

tal settings. Furthermore, the complex nature of ecosystems undermines the study of the interconnection 

between land cover and microbial community composition in field experiments and observational settings 

\[\text{[Levin, 1998]. The response of an organism to an environmental signal can be modulated by the presence} \]

and abundance of the other organisms in the ecosystem \[\text{[Green and Sadedin, 2005; Levins and Lewontin,} \]

\[\text{1980; Wang et al., 2020]. Because the interdependencies of environmental parameters induce confounding} \]

effects into our statistical analyses, they, furthermore, undermine our ability to detect direct and causal links. 

To make matters worse, in a fixed-size landscape, the areas of different land cover classes are not statisti-

cally independent from each other because an increase in one necessarily leads to decreases in others \[\text{[King} \]

\[\text{et al., 2005]. Because of this, bioindicators (species identified by the indicator value function) can be used} \]
as apparent proxy measurement for the ecological variables they are indicative for (and therefore of use for biomonitoring schemes), but are not necessarily in a biologically relevant relationship with it (Landres et al. 1988, Simberloff 1998).

While not being able to identify causal effects, with the required statistical and interpretative caution, insights into how freshwater microbiomes might be affected by land cover and land cover changes can still be gained. For example, recent studies have shown that microbial communities in waterways and lakes that are surrounded by different land cover types are significantly dissimilar (Kraemer et al. 2020, Saxena et al. 2015) and that land conversion for agricultural or urban uses influence the microbial communities of nearby stream sediments (Martin et al. 2020). Furthermore, it is well-known that the introgression of nitrates and other nutrients can lead to eutrophication and has a strong and characteristic effect on the microbial community composition (Gatti et al. 2018, Han et al. 2019, Sagova-Mareckova et al. 2021). Nevertheless, a systematic examination of the interrelation between land cover and microbial community composition for lakes in Europe is lacking for now.

In a prior publication, we developed the covariation framework, a statistical framework for the study of environmental microbiomes in observational settings such as environmental monitoring (Sperlea et al. 2021). The central idea behind this approach is to find a projection of the high-dimensional microbial community composition into the one-dimensional space of a target environmental parameter; calculating the $R^2$ between this projection and the measured values of the environmental parameter in question corresponds to the amount of variation in the latter explained by the variation in the former and can be considered a measure of covariation. The covariation framework circumvents many of the obstacles described above: First, by estimating the covariation using machine learning methods that can model non-linear dependencies in non-independent data, like Random Forests (Breiman 2001), as projection function, it handles the interdependencies between microbial species in the dataset. Second, it explicitly avoids any association with direct interaction, correlation, let alone causal relationships between the microbiome and the environmental parameters in question.

In this paper, we use the covariation framework as well as other machine learning-based approaches to study the relationship between land cover surrounding a lake and the microbiome of the lake. To this end, we analyze the largest amplicon sequencing dataset of European lakes available to date in concert with land cover data from the OpenStreetMap (OSM) project as well as the CORINE Land Cover (CLC) dataset from the Copernicus Land Monitoring Service (European Union 2012, OpenStreetMap contributors 2017). The former of the two data sources provides an open, community-driven, and, thus, rather detailed but
potentially incomplete land cover categorization. In contrast, the latter dataset is based on high-resolution satellite imagery and contains a hierarchical categorization of land cover in 44 classes. Based on these analyses, we identify multi-target bioindicators, i.e., species indicative of multiple environmental parameters, as well as environmental parameters that might act as drivers of microbial community composition. Furthermore, we propose a novel data abstraction, the response map, that clusters environmental parameters in terms of the response they engender in the microbiome. Aside from providing targets for future experimental investigation, the results presented here highlight that aggregation of variables when studying ecosystems can obscure real relationships. Finally, this study provides a thorough analysis of the response of the lake microbiome with regard to a wide range of land cover and physico-chemical parameters.

2 Materials and methods

2.1 Amplicon sequencing

Sampling was part of a pan-European study conducted in August 2012 (eukaryotic sequences are published in (Boenigk et al., 2018); NCBI Bioproject PRJNA414052, prokaryotic sequences are published and described in (Nuy et al., 2020) and (Bock et al., 2020); NCBI Bioproject PRJNA559862). Methods for data collection, extraction, sequencing, and amplicon processing are described in detail in these studies (Bock et al., 2020, Boenigk et al., 2018, Nuy et al., 2020) and will be briefly outlined below.

To analyze bacterial and protistan freshwater communities on a large scale, 280 lakes were sampled throughout Europe. Sampling details and information on measured physico-chemical and geographical parameters can be found in (Boenigk et al., 2018). For DNA analyses, filtered water samples were air-dried and frozen in liquid nitrogen. Genomic DNA was extracted using the my-Budget DNA Mini Kit (Bio-Budget Technologies GmbH, Krefeld, Germany) with modifications after (Boenigk et al., 2018). Amplicon sequencing targeted the V2-V3 region of the 16S rRNA gene for bacteria, the V9 region of the 18S, and the ITS1 gene for eukaryotes. Samples were commercially sequenced (Fasteris, Geneva, Switzerland) on an Illumina HiSeq 2500 sequencer generating 300 bp long paired-end reads. Adapter removal, quality trimming, and demultiplexing were performed by the sequencing company.

Sequence processing was performed using a provisional version of the Natrix pipeline (Welzel et al., 2020). If not stated otherwise, all software versions and parameters were used as described in (Welzel et al., 2020). The main steps included quality checks using FASTQC (Andrews, 2010) and PRINSEQ (Schmieder and Edwards, 2011), assembly of paired-end reads with PANDASeq (Masella et al., 2012) and dereplication and chimera
removal using UCHIME (usearch v7.0.1090 with default parameters) \cite{Edgar2011}. AmpliconDuo \cite{Lange2015} was used to discard sequences that were not found in both technical replicates. The remaining sequences were clustered using SWARM \cite{Mahé2014} and further aggregated to identical V9 sequences. This aggregation served as the basis for the OTU tables. The taxonomic assignment of the eukaryotic sequences was performed by a BLAST search \cite{Altschul1990} against the NCBI nt database (from Dec 5, 2017) and for the prokaryotic sequences against SILVA SSURef 132 \cite{Quast2012}. For all downward analyses, we combined the prokaryotic and eukaryotic OTU tables.

### 2.2 Land cover data

Two different land cover datasets were used in this study. For both, we accessed data for the year 2012 because this was also the year the lake samples were collected. The CLC dataset was downloaded from the official website of Copernicus Earth Observation program (CLC 20212, v.2020_20u1, 100m raster GeoTiff) \cite{EuropeanUnion2012}. The relative areas of the land cover classes were extracted from the dataset for circular areas around the sampling points with different radii using QGIS 3.16 \cite{QGISDevelopmentTeam2020}. Areas were aggregated to higher-level land cover classes according to the hierarchical CLC class model.

OSM land cover data was extracted from the OSM planet file from September 2012 archived at archive.org. This file was loaded in a PostgreSQL database and queried using a routine adapted from SEDE-GPS \cite{Sperlea2018} to retrieve the map tiles surrounding the sampling position, to fuse these, and to extract a circular area of a given radius. Map tiles were rendered using the default mapnik map style, which was adjusted to (i) merge pixels of land use sub-categories with the respective main category (such as “tertiary road” with “road”) and (ii) remove signs, labels, and point of interest markers. The pixel-areas summarised per unique category of the resulting image were read out and translated back to meters.

Outlier land cover values in all subsets (concerning both the radius as well as the land cover category) of these two datasets were detected using the function `boxplot.stats` in R 4.0.3 for the different radii and land cover categories, separately. Samples containing an outlier or a value of zero for a given land cover category at the given radius were discarded for the analysis of the respective land cover category and radius.

Additional physico-chemical parameters were taken from \cite{Sperlea2021}.

### 2.3 Microbial biodiversity and land cover

A straightforward way of determining whether land cover changes impact lake microbiomes is to assess whether the distribution of land cover types surrounding the lake is predictive of the lake’s microbial biodi-
versity. To this end, we extracted the relative area covered by different land cover categories in circular areas around the sampling sites of the European lake dataset from both the CORINE land cover (CLC) dataset as well as the OpenStreetMap (OSM) project (see Methods). These two datasets differ in the way they were generated and their categorization of land cover. While the former is derived from satellite data, the latter is annotated in a community-driven manner, based on landscape features observed “on the ground”. To distinguish between effects present at shorter or longer geographic ranges, we extracted and analyzed areas surrounding the sampling points within radii ranging from 1 km to 10 km, in steps of 1 km, as well as 25 km and 50 km. We then assessed the degree to which the relative sub-areas of the land cover categories contained in the extracted areas can be used to predict a set of biodiversity metrics calculated for the microbial communities of the sampled lakes using Random Forest models.

To calculate biodiversity metrics, the OTU table was rarefied using the \texttt{rrarefy} function from the R package \texttt{vegan} (v2.5-6, ref. Oksanen et al., 2019). Biodiversity metrics were calculated from rarefied OTU tables using the \textit{diversity} (Shannon index, Simpson diversity, inverse Simpson diversity), and \textit{renyi} (Renyi entropy) functions from the R package \texttt{vegan}, except for species richness, which is the total number of OTUs present, and Pielou’s evenness, which was calculated by dividing the sample’s Shannon index by the log of the sample’s richness (Chao and Jost, 2015, Daly et al., 2018). Renyi entropy metrics were calculated for $\alpha = \{0.25, 0.5, 1, 2, 4, 8, 16, 32, 64, \infty\}$, as different values for this parameter drastically change the metrics sensitivity to relative species abundance (Chao and Jost, 2015). For the prediction of biodiversity metrics based on land cover categories, Random Forests from the R package \texttt{caret} (version 6.0.86, ref. Kuhn, 2008) were trained using 10-fold cross-validation without further feature selection.

### 2.4 Covariation framework

Covariation between the lake microbiomes and land cover areas was quantified using the covariation framework presented in ref. Sperlea et al., 2021. Methodologically, the framework is a straightforward machine learning approach, training a machine learning model to predict the values of an environmental parameter after feature selection. However, the model’s prediction is interpreted as the projection of the microbial community composition to the space of the target variable while leveraging the model’s potential to model non-linear interdependencies in the microbiome. This way, the coefficient of determination, $R^2$, can be interpreted as a measure of covariation between the microbiome as a whole and the target variable. As feature selection method, the \texttt{multipatt} function (with the parameter “indval”) from the \texttt{indicspecies} R package (v1.7.9, ref. Cáceres and Legendre, 2009) was used after Hellinger transformation (Legendre and Gallagher, 2009).
of the OTU counts to identify bioindicator OTUs for tertiles of the respective land cover category.

Random Forest models from the R package `caret` (version 6.0.86, ref. [Kuhn, 2008]) were trained in a 10-fold cross-validation scheme with the OTU tables as independent and the relative area of a single land cover category as dependent variables, with both being centered and log-ratio transformed. Because of statistical limitations of the Random Forest model, some combinations of area size and land cover category could not be used for model training.

Confidence intervals for the model evaluations were estimated based on resampling of predicted and measured dependent variable pairs with replacement with thousand repetitions. Statistical significance of relevant models was asserted by comparing the $R^2$ value with results gathered by thousand repetitions of training models with the same hyper-parameter setting on resampled biodiversity data in a Student’s t-test as implemented in the `t.test` function in R 4.0.3.

### 2.5 Bioindicator analysis

Bioindicator OTUs for land cover categories were identified using the indicator species method as implemented in the `multipatt` function in the `indicspecies` R package (v1.7.9) [Cáceres and Legendre, 2009], [Dufrêne and Legendre, 1997]. A significance level of $\alpha = 0.05$ was applied after Benjamini-Hochberg correction for the total number of land cover categories analyzed in this study.

The similarity of two environmental parameters was calculated in terms of the microbiome’s response to changes in them by calculating the Jaccard similarity between the lists of OTUs indicative of the two parameters. The resulting similarity matrix was visualized as a force embedded network using the function `qgraph` from the package `qgraph` (v1.6.9, ref. [Epskamp et al., 2012]). Furthermore, a dissimilarity matrix was derived from the similarity matrix by inversion after the addition of a random number in the order of $10^{-8}$ in order to avoid the division by zero. This dissimilarity matrix was visualized as a dendrogram using `upgma` and `ggdendrogram` from the packages `phangorn` (v2.5.5., ref. [Schliep, 2011]) and `ggdendro` (v0.1.22, ref. [de Vries and Ripley, 2020]), respectively. For the ordination of the environmental parameters, the `metaMDS` function from the R package `vegan` was used.

For the bioindicator network, an edge was created between all pairs of bioindicator OTUs that are indicative of at least one common environmental parameter. Null-hypothesis networks were created the same way based on resampled indicator lists; for these, each lake parameter is assigned the same number of randomly selected OTUs as pseudo-indicators in such a way that the distribution of cardinalities is the same as that of the real OTUs. Node properties (degree, closeness centrality, eigenvector centrality, page rank, and authority
score) were calculated using the **igraph** package in R 4.0.3 (v1.2.6, ref. Csardi and Nepusz 2006).

### 2.6 Network inference methods

In this study, we apply several methods for the inference of network structures from OTU tables. Most of these employ similarity measures as edge weights calculated between all pairs of OTUs, which act as nodes in the network. Simple co-occurrence, checkerboard score (Connor and Simberloff 1979; Stone and Roberts 1990), Bray-Curtis similarity (Bray and Curtis 1957), Kullback-Leibler divergence (Kullback and Leibler 1951), Pearson and Spearman correlation were used as similarity metrics. The co-occurrence metric was defined as the number of samples in which the two OTUs in question had non-zero occurrence. Pearson and Spearman correlations were calculated using the `cor` function in R and results below the significance level \( \alpha = 0.05 \) were discarded. For the calculation of the Kullback-Leibler divergence, zeroes in the OTU table were replaced by \( 10^{-8} \) to avoid infinities created by the logarithm. Additionally, the method SparCC (Friedman and Alm 2012) was used as implemented in the `sparcc` function from the `SpiecEasi` package (v1.1.0, ref. Kurtz et al. 2015) with default parameters. For all networks, OTUs that have zero counts for 25 or more sampling sites were excluded from the analysis to avoid statistical artifacts that are based on the rarity of the OTUs in question (Berry and Widder 2014).

All figures were generated using the R packages `ggplot2` (v.3.3.2, ref. Wickham 2016), unless otherwise noted, and following the guidelines laid out in ref. [Hattab et al. 2020](#).

### 3 Results

#### 3.1 Microbial biodiversity in lakes only marginally reflects differences in surrounding land cover

A straightforward way of determining whether land cover changes impact lake microbiomes is to assess whether the distribution of land cover types surrounding the lake is predictive of the lake’s microbial biodiversity. To this end, we extracted the relative area covered by different land cover categories in circular areas around the sampling sites of the European lake dataset from both the CORINE land cover (CLC) dataset as well as the OpenStreetMap (OSM) project (see Methods). To be able to distinguish between effects present at shorter or longer geographic ranges, we extracted and analyzed areas surrounding the sampling points within radii ranging from 1 km to 10 km, in steps of 1 km, as well as 25 km and 50 km. We then assessed the degree to which the variation in the relative sub-areas of the land cover categories contained in the extracted...
Figure 1: Evaluation of Random Forest models trained to predict biodiversity metrics from land cover. A. Results for the land cover in a radius of 3 km around the sampling site and all biodiversity metrics surveyed in this study. Lines represent confidence intervals estimated based on resampling (see methods). For results of other radii, and results for Renyi entropy with other values for $\alpha$, see supplementary figures 1 and 2. B. Results for the full range of radii as well as 25 and 50 km for Renyi entropy with $\alpha = 0.5$. Grey areas and error bars represent confidence intervals; results for 25 and 50 km are visually separated to underline that these radii are not in the range of the other radii.

areas explain the variation in microbial alpha diversity of the sampled lakes using Random Forest models. A set of alpha diversity metrics was used instead of a single metric because they capture different mathematical characteristics of biodiversity [Daly et al. 2018].

Our results show that there is, at best, a marginal relationship between land cover and microbial biodiversity, as no combination of radius, biodiversity metric, and dataset results in an $R^2 > 0.2$ (see figure 1A, supplementary figure 1 and 2). For all radii studied here, the lake microbiome’s Renyi entropy is most predictable from land cover, followed by species richness. Furthermore, we found no significant difference between $R^2$ values obtained for the same biodiversity metric at different radii (see figure 1B), indicating that the results presented here are most likely due to statistical artifacts rather than processes that shape microbial biodiversity based on surrounding land cover. In general, the land cover data collected from the OSM dataset is less predictive for microbial biodiversity than the CLC datasets (see figures 1A and B). Therefore, we focus our further analysis on the CLC dataset. Taken together, these results suggest that if land cover has a structuring effect on lake microbiomes, this relationship is not reflected in alpha diversity metrics.
Figure 2: Covariation between land cover surrounding lakes and the lake’s microbiome. Numbers in brackets (A and C) and in x-axis labels (B) refer to the CLC category number code (see table 1). For ease of display, full-length labels of the CLC categories have been shortened in some cases. Vertical lines represent confidence intervals estimated from resampling (see methods) and dots represent covariation as obtained in model evaluation. A. Results for all high-level land cover categories in the CLC dataset. B. Results for low-level land cover categories; for each land cover category, only the results for the radius with the highest R^2 are shown. C. Results for all mid-level land cover categories in the CLC dataset. Vertical lines between facets separate groups of categories that are subcategories of the categories in A. For all results, see supplementary table 1.

3.2 Microbial community structure covaries with specific land cover categories

While not visible at the level of alpha diversity, we hypothesized that there must be an apparent relationship between land cover surrounding the lakes and the lake microbiome when analysing it at the fine-grained level of microbial community composition. To investigate this hypothesis, we applied the covariation framework to the OTU tables and the relative areas of the land cover categories present in the CLC datasets. Higher R^2 values indicate a higher degree of covariation between the microbiome as a whole and the target parameter, i.e., the relative area of the land use category in question. On level 1 of CLC category hierarchy, we observe covariation of R^2 > 0.05 for “artificial surfaces (1)” and “agricultural areas (2)” at very low radii as well as increasing covariation for “forest and semi-natural areas (3)” with increasing radii (see figure 2A). In
Table 1: Numbers of microbial indicators for land cover categories (with respective radius, left) and physico-chemical parameters (right) and the respective $R^2$ resulting from the covariation framework (results for physico-chemical parameters taken from Sperlea et al. (2021)).

| Parameter Indicators | Parameter | Radius (km) | $R^2$ |
|----------------------|-----------|-------------|-------|
| Herbaceous vegetation (32) | 1056 | 7 | 0.42 |
| Forest, semi-natural areas (3) | 703 | 10 | 0.36 |
| Arable land (21) | 445 | 9 | 0.36 |
| Non-irrigated arable land (211) | 416 | 9 | 0.34 |
| Discontinuous urban fabric (112) | 276 | 8 | 0.29 |
| Agricultural areas (2) | 272 | - | - |
| Natural grasslands (321) | 265 | 10 | 0.29 |
| Urban fabric (11) | 247 | 7 | 0.29 |
| Artificial surfaces (1) | 177 | - | - |
| Forests (31) | 140 | 7 | 0.15 |
| Inland waters (51) | 77 | 6 | 0.18 |
| Industrial or commercial units (121) | 77 | 10 | 0.17 |
| Heterogeneous agricultural areas (24) | 69 | - | - |
| Transitional woodland-shrub (324) | 61 | 10 | 0.09 |
| Open spaces with little vegetation (33) | 52 | 9 | 0.42 |
| Rare Rocks (332) | 34 | 9 | 0.32 |
| Water bodies (5) | 32 | - | - |
| Pastures (23) | 21 | - | - |
| Sparsely vegetated areas (333) | 15 | 2 | 0.38 |
| Permanent crops (22) | 6 | - | - |
| Fruit trees (222) | 3 | 8 | 0.40 |
| Inland wetlands (41) | 2 | 4 | 0.20 |
| Artificial, non-agricultural vegetated areas (14) | 2 | - | - |
| Industrial, commercial and transport units (12) | 2 | - | - |
| Vineyards (221) | 1 | 6 | 0.18 |
| Mineral extraction sites (131) | 1 | 9 | 0.18 |
| Mine, dump and construction sites (13) | 1 | 9 | 0.18 |
| Dump sites (132) | 1 | 8 | 0.22 |
| Construction sites (133) | 1 | 10 | 0.57 |

In contrast, the anthropogenic effects that are expected with built surfaces and agriculturally used land (Gatt et al., 2018, Martin et al., 2020) act on rather short ranges.

The covariation observed between the lake microbiome and land cover categories at level 2 of the CLC hierarchy paints a more nuanced picture (see figure 2C). For example, we observe increasing covariation with the lake microbiomes at increasing radii for the land cover categories “arable land (21)” and “scrub and/or herbaceous vegetation associations (32)”. In contrast, for “urban fabric (11)” and “inland waters (51)”, we observe a peak in covariation at radii of 7 km and 6 km, respectively, with lower $R^2$ for the other radii. For “forests (31)” and “open spaces with little or no vegetation (33)”, the covariation for different radii stays within the respective confidence intervals of the covariation at the 1 km radius. Notably, the covariation between the microbiome and sub-categories of a CLC category often deviate strongly from the covariation between the microbiome and the respective super-category. The same applies to covariations observed at level 3 of the CLC category hierarchy (see figure 2B). Taken together, these results show that a broad array of land cover categories have an impact on the lake’s microbial community composition at the OTU level and they do so at different radii. This can be interpreted as a reflection of different mechanisms being at play for the influence of lake ecology of different land cover categories. Furthermore, the more specific land cover categories at levels 2 and 3 of the CLC category hierarchy generally show higher levels of covariation with the lake microbiome, indicating that aggregation of land cover classes can hide relationships apparent at more fine-grained levels of analysis.

To identify general spatial trends, we separately calculated the mean covariation of all land cover categories...
for each radius and land cover hierarchy level. For all but one radius-hierarchy level combination, the average $R^2$ value is below 0.15 (see supplementary figure 3) and throughout all combinations, the relative standard deviation is close to or higher than 100%. This shows that there are neither general spatial trends nor a generally higher covariation at higher levels of the CLC hierarchy. Instead, to observe specific, radius-dependent effects of land cover on the lake microbiome, it is important to differentiate between land cover categories.

3.3 Microbial lake bioindicators for surrounding land cover categories

Using the indicator value method that is part of the covariation framework, we identified 2,354 OTUs that act as bioindicators for the land cover categories in a total of 4,453 indicator-parameter pairs (for a complete list, see supplementary table 2). Among the land cover categories studied in this paper, for “scrub and/or herbaceous vegetation associations (32)” and “forest and semi-natural areas (3)” we identify the highest number of indicator OTUs with 1056 and 703 OTUs, respectively (see table 1). Most of the indicator OTUs are Bacteria (87%) from the phyla Proteobacteria (29%), in particular Alphaproteobacteria (13%), Bacteroidetes (28%), in particular Flavobacteraea (16%), or Cyanobacteria (15%). Furthermore, most of the OTUs obtained are indicative for more than one land cover parameter (fig. 3A). All OTUs indicative for more than seven land cover parameters are bacteria (see tables 2 and 3). Taken together, these results support the notion that bacteria are more sensitive to environmental changes or respond to environmental signals in a different manner than microbial eukaryotes (Bock et al., 2020, Logares et al., 2018).

In a previous paper, we identified bioindicator OTUs for physico-chemical parameters while working with the same amplicon sequencing dataset as analyzed here Sperlea et al. (2021). Comparing the results of this analysis with those in the prior publication, we observed that almost all bioindicators indicative for at least eight land cover categories are also indicative of the lake’s altitude (see table 2). This underlines the central role the geographic location of a site plays in its ecology (see discussion) (Bock et al., 2020, Karlsson et al., 2005). However, a literature search for either the multi-target bioindicators on the Genus or Species level of taxonomy did not result in any information supporting or calling our results into question.

3.4 Structural insights into the lake microbiome from multitask bioindicators

To further elucidate the relationship between the lake microbiome and environmental parameters, we combined the bioindicator OTUs identified for land cover parameters with those for physico-chemical parameters identified in Sperlea et al. (2021). This way, we obtained a data structure that can be described as a set...
of maps between a set of OTUs and a set of environmental parameters. From this, we derived two distinct similarity matrices: One stating the similarity of OTUs in terms of the number of environmental parameters they are indicative of, and one that displays the similarity of the parameters in terms of the OTUs assigned to them.

The former can be turned into a bioindicator network as follows. Each bioindicator OTU is assigned to a node and edges are drawn between nodes representing OTUs that are indicative for at least one common environmental parameter (see supplementary table 3 for the entire network). We noticed correlations between the cardinality (i.e., the number of occurrences of an OTU in all bioindicator lists) of the nodes of the

Figure 3: (Caption on the following page.)
bioindicator network and the respective nodes’ degree, closeness centrality, eigenvector centrality, page rank, and authority score (see supplementary figure 4). Because this result could be due to basic graph properties, we compared the square of the Pearson correlation coefficient, $R^2$, resulting from the correlation of cardinality with node properties of the bioindicator network with those gained from resampled networks (see Methods for details). We find that the nodes in the bioindicator network have statistically significant properties (see figure 3B), which suggests a biological relevance of the bioindicator network’s structure.

Furthermore, we compared the bioindicator network to networks inferred from the original OTU table. More specifically, we asked whether the node properties generated using network inference methods correlate with the node properties of the bioindicator network. We chose this approach to comparing the two network structures as it can capture relative differences between node properties and might thus be robust with regard to global effects of a network method, e.g., a consistently lower degree. Our results show that applying a high cut-off to co-occurrence and checkerboard score similarity matrices results in networks similar to the bioindicator network. In contrast, neither correlation-based nor compositionality-aware methods do so (see figure 3C).

The second data structure presents the similarity of pairs of environmental parameters in terms of the Jaccard similarity of the list of bioindicator OTUs identified for them. A high Jaccard similarity between two environmental parameters suggests that the microbiome responds to changes in the parameters in a similar manner. Along these lines, a visualization of this similarity matrix can be seen as a response map of the microbiome with regard to environmental changes. We attempted to visualize the resulting similarity matrix using non-metric dimensional scaling with up to 10 dimensions but were unable to receive stress values < 0.05, suggesting that the responses of the microbiome to environmental change are non-trivial. Nevertheless, the visualization of the similarity matrix as a UPGMA-derived dendrogram and an undirected graph (see figures 4A and B, respectively) results in multiple distinct clusters of highly similar environmental
Figure 4: Clustering of physico-chemical and land cover parameters using the Jaccard similarity of the features’ lists of bioindicator OTUs. Visualization of the similarity matrix as A. dendrogram using UPGMA and B. force embedded graph, in which edge size represents a higher Jaccard similarity. Node coloring in B represents clustering of parameters in A. Numbers represent CLC land cover categories according to the CLC legend (see table 1). Other abbreviations: A.G/Alk.Gran – alkalinity, Alt – altitude, Ann – anions, CatSum/CtS – sum of all cation concentrations, COND/CON – conductivity, Ctn – cations, DOC – dissolved organic carbon, DRSi/DRS – dissolved reactive silica, LF – conductivity (measured in the field, SumIons/SmI – sum of all ion concentrations, TP – total phosphorus.

The largest one of these comprises the concentration of magnesium, potassium, anions, cations, phosphorus, as well as temperature, pH, altitude, and a wide range of land cover categories. Most of the clusters identified here can be explained with regard to physico-chemical and landscape ecological processes in lakes (see Discussion). A further notable result is the relatively high distances of many of the same CLC categories’ subcategories in both the dendrogram and the graph. This underscores our prior finding that lake microbiomes react to different land cover categories in different ways (see figure 2).

4 Discussion

The development of scalable Next-Generation Sequencing (NGS) methods has dramatically furthered the study of environmental microbiomes (Boughner and Singh, 2016, Snyder et al., 2008, Tan et al., 2015). However, technical and theoretical obstacles interfere with analyzing the wealth of data generated using NGS methods. For one, the dimensionality of microbiome datasets (i.e., the number of microbial species...
in ecosystems) is usually many orders of magnitude larger than the number of samples. For example, the
dataset analyzed in this paper is, in terms of the number of samples, the biggest amplicon sequencing dataset
for European lake microbiomes published to date, but still contains ~1000 times more OTUs than samples.
Together with the sparsity of OTU tables, this places the analyses of microbial communities at the edge
of statistical feasibility, as, in such a domain, regression is ill-defined (Carr et al., 2019 | Weiss et al. 2017
2015). Furthermore, the counts of different OTUs in the OTU tables are not independent in at least two
regards. Firstly, they are compositional because of technical details of the sequencing procedure (McGeoch
and Chown 1998 | Quinn and Erb 2020). Secondly, in the environment, different populations interact,
through which the size of one population can, directly or indirectly, influence the size of another (Schaffer
1981). The independence of features is, however, necessary for many statistical approaches.

In this paper, we study the relationship between lake microbiomes and the land cover surrounding the
lakes while taking into account the aforementioned statistical obstacles. It is important to stress that such
an analysis cannot uncover direct or functional connections without experimental validation; instead, the
results presented here point to apparent relations that are present in lake ecosystems and that structure their
functioning. The central analytical tool in our study is the covariation framework, in which Random Forest
models are trained to approximate a projection of the microbial community composition to a one-dimensional
space defined by one environmental parameter (Sperlea et al. 2021). As, in environmental, observational
settings, we cannot exclude the possibility of the presence of confounding factors, we denote the relationship
between the projected microbiome and the environmental parameter of interest as covariation instead of
correlation to emphasize that we are not quantifying direct effects.

Our results show that the microbiome covaries to a considerable extent ($R^2 > 0.3$) with the areas covered
with forest-like vegetation (but not forest areas themselves), arable land, open spaces, plantations, and
constructed environments such as urban areas and roads (see table 1, figure 2 and supplementary table 1).
These results are in agreement with other recent studies of land cover and lake microbiomes (Kraemer et al.
2020 | Marmen et al. 2020 | Sperlea et al. 2018). On the other hand, the low covariation of the microbiome
with, e.g., areas of land covered with pastures or mine and construction sites (CLC categories 23 and 13,
respectively) indicate that changes in these categories of land cover are not reflected in the microbiome.
On a more speculative level, these results could indicate that there are more ecological processes linking
the lake microbiome to areas from the former group of land cover categories than to areas from the latter.
More notable, the covariation between the microbiome and land cover categories are, generally, lower than
those between the microbiome and physico-chemical parameters reported by Sperlea et al. (2021). This
result is not surprising as changes in land cover do not impact lake microbiomes directly but relayed via physico-chemical parameters [Marmen et al., 2020, Saxena et al., 2015]. A notable exception would be direct pollution generated through land use as, e.g., tire wear on roads and urban areas, but we do not find evidence for this in our analyses. In addition, areas that are too small to appear in the CLC dataset but that have a strong impact on lake ecology, such as a small group of trees at the shore of a lake, are not represented in our analysis.

Aggregation of similar parameters is a necessary step for the analysis of ecological processes but comes with the danger of obscuring ecologically relevant heterogeneity and, by that, losing explanatory power (Greenland and Morgenstern, 1989, King and Baker, 2010, Ulanowicz, 1986, Yodzis, 1988). For example, it has been shown that aggregation of OTUs to higher-order taxonomic groups reduces predictive power in biomonitoring settings, although this is in part due to the incompleteness of taxonomic reference databases (Chen et al., 2013, Janßen et al., 2021, Kermarrec et al., 2014, Sagova-Mareckova et al., 2021, Sperlea et al., 2021). In our results, both the aggregation of OTU tables to alpha diversity metrics (see figure 1 and supplementary figure 3) and the aggregation of land cover classes to higher-order CLC categories (see figure 2 and supplementary table 1) lead to lower $R^2$ values. This underlines the importance of heterogeneity and non-linear processes in modelling the relationship between lake microbiomes and their habitat.

Just as the OTU counts, the environmental parameters studied here are not statistically independent but compositional and spatially autocorrelated (King et al., 2005). While, in most experimental settings, one would attempt to identify the effect of a single parameter on the study object by controlling for confounding effects via partial correlations or regression on residuals. However, spatial autocorrelation and indirect effects are significant for ecosystem functioning and integral for understanding it. They should not be considered “noise” that needs to be removed in the analysis (Freckleton, 2002, Legendre, 1993, Levins and Lewontin, 1980, Yodzis, 1988).

For example, the interdependence of ecological parameters leads to the emergence of apparent drivers of the microbial community composition, i.e., parameters that covary with the microbiome but are not necessarily directly and causally linked to it. In the context of this study, drivers appear most directly as parameters for which high numbers of microbial bioindicators have been identified. Our analysis suggests that the main driver of microbial community composition is the altitude of the lake (see table 1), which is in line with altitude being one of the determining factors of vegetation and ecosystem composition. The other parameters with high (> 500) numbers of indicators have been described as drivers of lake ecology themselves (e.g., temperature) and/or are known to be strongly correlated with altitude (as, e.g., herbaceous and forest
vegetation, temperature, pollution, nutrient load), or with correlates of it (as water conductivity and pH is dependent on water temperature) \cite{Forster2021, Urban2002}. These results validate the use of microbial bioindicators for the analysis of ecosystem structure.

To gain further insights into how the microbiome responds to its environment that go beyond the concept of ecological drivers, we compared the lists of bioindicators identified for the ecological parameters studied here and in a recent publication \cite{Sperlea2021} and visualized the results in a response map (figures 4A and B). Intuitively, environmental parameters that are close or connected with high-weighted edges in the response map elicit similar responses from the microbiome. Instead of attempting to identify direct or causal relationships, this approach allows us to analyse the structure of the ecosystem “through the eyes of the microbiome”. In principle, such an observer-based methodology allows for the integrated analysis of the effects of heterogeneous environmental parameters as long as the microbiome is affected by them.

We identify four clusters of ecological parameters in the response map (figures 4A and B), the largest of which contains the altitude, but also a wide range of land cover parameters and ion concentrations. This cluster is probably due to altitude acting as a major ecological driver. Furthermore, we find one cluster that contains the land cover parameters “Open spaces with little vegetation (33)”, “Bare Rocks (332)”, and “Sparsely vegetated areas (333)”, one cluster comprising the concentration of bicarbonate and calcium and the alkalinity, and a last one that connects the concentration of dissolved organic carbon with “ Transitional woodland-shrub (324)”. The first of these clusters suggests that the sub-categorization of category 33 as present in the CLC categorization is not reflected in the lake microbiome. The second cluster might be explained by the existence of a calcium-bicarbonate equilibrium in freshwater ecosystems \cite{Kopacek2020} and a subpopulation of the lake microbiome responding to deviations from it. The third and last cluster points towards the fact that plant matter is the principal source of dissolved organic carbon in soils \cite{Mayer2020}, and therefore, again, the functional connection between freshwater ecosystems and the landscape surrounding them.

Like environmental parameters that act as drivers of ecosystem processes, bioindicators indicative of a high number of environmental parameters, i.e., multi-target bioindicators, might be particularly sensitive with regard to environmental change. The taxonomic distribution of the multitask bioindicators for land cover categories identified here (see table 3) deserves a few words of discussion. First, the absence of Eukaryotes among the high-ranking multitask bioindicators suggests that bacterial niches can be more specific with respect to environmental factors, making Bacteria presumably more potent and more sensitive indicators for ecosystem health. Second, the relatively low number of Alpha- and Betaproteobacteria among the multitask
indicators contrasts their high abundance in a broad range of freshwater ecosystems (Šimek et al., 2013) as well as the environment-specific abundances of certain taxa among these classes (Nuy et al., 2020). Both of these findings point towards the difference between fidelity and specificity as defined in the context of the IndVal method (Cáceres and Legendre, 2009; De Cáceres et al., 2010; Dufrène and Legendre, 1997). Third and last, the attribution of ecological functionality to these OTUs is not possible, in part because only a tiny minority of microbes have been cultured and studied to a sufficient degree (Thomas and Segata, 2019). Note that the question of how to attribute ecological function to microorganisms goes beyond the scope of this study, which is focused on the identification of microbial bioindicators.

Taken together, in this study, we analyse the relationship between land cover and lake microbiomes using machine learning methods. We propose novel methods for the analysis of relationships between ecological parameters that take the complexity of lake ecosystems into account and show that their results can be explained with regard to known ecological processes. While this study is centered around the largest lake microbiome dataset published to-date and the systematic analysis of the link between land cover and microbial ecology, it still has some limitations. For example, because the samples analyzed in this study were taken over a few days, seasonal effects are not addressed in this study. Furthermore, the fact that the underlying dataset is not longitudinal might obscure the diversity of European lakes. Therefore, further studies that adopt this methodology for large-scale microbial biomonitoring datasets are required to confirm our findings.

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Conflict of Interest
The authors declare that they have no competing financial interests.

Data Accessibility
Raw sequencing data are available under the NCBI BioProject IDs PRJNA414052 and PRJNA559862.

Author’s contributions
TS designed and performed all computational analyses, JPS provided the OSM dataset, HD helped with the network analyses, DB performed the amplicon sequence analysis with Natrix, DB and JB provided the sequencing datasets, JB, GH, and DH supervised the study. All authors discussed the results and wrote and revised the manuscript.
Table 2: Multitask OTUs identified in this study for more than 6 land cover categories and the physico-chemical parameters they have been identified as indicators for in Sperlea et al. [2021]. Numbers represent CLC land cover categories (see table 1). For taxonomic annotation of these OTUs, see table 3.

| OTU  | Occ. | 1  | 11 | 112 | 121 | 2  | 21 | 211 | 23 | 24 | 3  | 31 | 32 | 321 | 324 | 33 | 332 | Phys.-chem. |
|------|------|----|----|-----|-----|----|----|-----|----|----|----|----|----|-----|-----|----|----|-----------|
| N115 | 10   | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF T |
| N213 | 10   | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF Mg pH T |
| N35  | 10   | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF pH T |
| N47  | 10   | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF Mg |
| N74  | 10   | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF TP |
| N1077| 9    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude Cations K LF Mg Na pH SO4 SumIons T |
| N178 | 9    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF pH T TP |
| N395 | 9    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF |
| N469 | 9    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude LF |
| N1276| 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N156 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N166 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N177 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N1844| 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N243 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N470 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N471 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N513 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N519 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N533 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N563 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N689 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N785 | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| N98  | 8    | X  | X  | X   | X   | X  | X  | X   | X  | X  | X   | X  | X  | X   | Altitude CI DOC T |
| Domain  | Phylum   | Class         | Order           | Family         | Genus        | Species                                      |
|---------|----------|---------------|-----------------|----------------|--------------|----------------------------------------------|
| N115    | Bacteria | Proteobacteria| Alphaproteobacteria | Sphingomonadales | Sphingomonadaceae | Synkogonema                                  |
| N213    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Sphingomonadaceae | Sphingomonas sp. CCGE4131                   |
| N35     | Bacteria | Proteobacteria| Alphaproteobacteria | Rhodospirillales | Acetobacteraceae | Rosemonas                                    |
| N427    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Cryomorphaceae | Fluviicola                                   |
| N74     | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Acetobacteraceae | Fluviicola                                  |
| N178    | Bacteria | Bacteroidetes | Cytophaga        | Cytophagales    | Cyclobacteriaceae | Fluviicola                                   |
| N505    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | N99 marine group | Fluviicola                                   |
| N499    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Cryomorphaceae | Fluviicola                                   |
| N1276   | Bacteria | Proteobacteria| Betaproteobacteria | Burkholderiales | Comamonadaceae | Fluviicola                                   |
| N136    | Bacteria | Cyanobacteria | Cyanobacteria    | Subsection 1    | Family 1     | Synechococcus                                |
| N215    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Cryomorphaceae | Fluviicola                                   |
| N577    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | N99 marine group | Fluviicola                                   |
| N1384   | Bacteria | Firmicutes    | Clostridia       | Clostridiales   | 1            | Clostridium sp.                              |
| N243    | Bacteria | Proteobacteria| Alphaproteobacteria | Rhodobacteriales | Rhodobacteriaceae | Rhodobacter                                |
| N470    | Bacteria | Proteobacteria| Gamma proteobacteria | Xanthomonadales | Xanthomonadaceae Incertae Sedis | Acidibacter                                                      |
| N471    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Flavobacteriaceae | Flavobacterium sp.                                   |
| N513    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Flavobacteriaceae | Acidibacter                                              |
| N519    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Flavobacteriaceae | Acidibacter                                              |
| N553    | Bacteria | Proteobacteria| Alphaproteobacteria | Sphingomonadales | Sphingomonadaceae | Nocorophaleum                                            |
| N563    | Bacteria | Firmicutes    | Clostridia       | Clostridiales   | 1            | Clostridium sp.                              |
| N649    | Bacteria | Bacteroidetes | Sphingobacteria  | Sphingobacteriales | NS11-12 marine group | un cultured Bacteroidetes bacterium |
| N785    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | N99 marine group | un cultured soil bacterium                      |
| N998    | Bacteria | Bacteroidetes | Flavobacteria    | Flavobacteriales | Flavobacteriaceae | Flavobacterium                                       |