Microbial ecosystems are dominated by specialist taxa

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Abstract

Abundance and specificity are two key characteristics of species distribution and biodiversity. Theories of species assembly aim to reproduce the empirical joint patterns of specificity and abundance, with the goal to explain patterns of biodiversity across habitats. The specialist-generalist paradigm predicts that specialists should have a local advantage over generalists and thus be more abundant. We developed a specificity index to analyse abundance–specificity relationships in microbial ecosystems. By analysing microbiota spanning 23 habitats from three very different data sets covering a wide range of sequencing depths and environmental conditions, we find that habitats are consistently dominated by specialist taxa, resulting in a strong, positive correlation between abundance and specificity. This finding is consistent over several levels of taxonomic aggregation and robust to errors in abundance measures. The relationship explains why shallow sequencing captures similar β-diversity as deep sequencing, and can be sufficient to capture the habitat-specific functions of microbial communities.

Keywords

Abundance, beta-diversity, communities, microbial ecology, microbiota, specialist taxa, specificity.


INTRODUCTION

The vast majority of microbial ecosystems, are characterised by highly skewed abundance-rank distributions: a few taxa account for the majority of individuals while most taxa are represented by only a few individuals (Connolly et al. 2014). In a spatial setting, the specificity, defined here as a measure for the unevenness of which a taxon occurs in different habitats, becomes important as abundance-rank distributions differ among habitats. The extremes of the specificity continuum are (1) taxa found with equal abundances in many habitats (generalists) and (2) taxa always and only found in one habitat (specialists). Extreme specialists are indicator species, with strong ecological preferences, that are specific to a given habitat (Dufrene & Legendre 1997). A long-standing view in ecology states that the differences between specialists and generalists arise from disparity in their resource utilisation: specialists have a narrow resource utilisation range and high peak performance, whereas generalists have a broader range but lower peak performance (Peers et al. 2012). If high performance is linked to high local competitive ability and migration does not greatly influence local population dynamics, specialists should have higher local abundance than generalists, which are found in several habitats. This pattern has for example been observed in the malaria parasite: species with narrow host range are associated with higher peak parasitaemia than species with a broad one (Garamszegi 2006).

Alternatively, specificity may be linked to abundance and the role of species in ecosystem functioning. In the case of host-associated microbial communities, hosts can recruit, either passively or actively (Kremer et al. 2013), their symbionts based on the functions they provide (Burke et al. 2011a). The impaired development of axenic animals also shows that microbiota provide essential functions to their hosts (Brummel et al. 2004; Rawls et al. 2004; Sison-Mangus et al. 2015).

Recruited microbes are expected to thrive and be locally abundant. If furthermore, the recruited functions are costly to maintain outside of the host, they are likely specific, resulting in a strong, positive correlation between local abundance and specificity.

A positive abundance–specificity relationship has important practical implications for our understanding of biodiversity. The total diversity of a landscape, γ-diversity, is traditionally partitioned between the average within-site diversity (α-diversity) and the among-site diversity (β-diversity). Several quantitative diversity indices accounting for abundance exist for community data (Graham & Fine 2008). They have in common that communities that share most or all of their abundant taxa are less β-diverse than communities that share none or a few. In the context of microbial ecology, where communities are often dominated by a few abundant taxa, we speculate that the β-diversity of less abundant taxa is driven by a positive correlation between abundance and specificity of microbial taxa. As a consequence of this, the β-diversity of these communities should be easy to recover with small sampling effort. In case of microbial data, this means shallow sequencing depth by next generation sequencing approaches.

Previous studies have noted a positive relationship between abundance and occupancy. Habitat specialist taxa tend to be rare when habitats are simply defined by locality (Székely & Langenheder 2014) but become dominant when habitats are defined along an abiotic gradient (Fortunato et al. 2013; Logares et al. 2013), especially at extremes of the gradient. Nemergut et al. (2011) showed that detectable taxa are generally confined to single assemblage but did not report information about their local abundance. Other findings suggest that this relationship may be widespread. For instance, a relationship between specificity and abundance has recently been hinted at in butterfly gut microbiota, where communities at different developmental stages were characterised by different stage-specific abundant taxa (Hammer et al. 2014). The
different stages are characterised by different diets, stressing that habitats can be influenced by diverse biotic and abiotic factors. The same relationship was apparent in sponge microbial communities (Motinho-Silva et al. 2014; Reveillaud et al. 2014), but the trend was not discussed. A combination of biological (transient, dead or dormant taxa) and technical (contamination, clustering methodology) reasons lead to the increasing classification of locally abundant species as unspecific when sampling depth was increased.

Here, we investigate the abundance–specificity relationship of microbial communities, aiming to test if we can generalise it in a wide range of environments. We develop a local specificity index that separates specificity from abundance and test how it correlates with relative abundance in different microbiota habitats, defined by environment type rather than locality. We introduce a permutation test to assess the significance of this relationship. We apply our method to three data sets of microbiota data, one from zooplankton (this study), one from humans (Costello et al. 2009) and the last one from environmental samples (Caporaso et al. 2011). All data sets, chosen to reflect a wide array of environments, habitats, sampling conditions and sequencing depths, revealed strong, positive relationships between abundance and specificity. We discuss the implications of this finding for biodiversity surveys and functional metagenomics as well as mechanisms that can lead to specificity and to high local abundance of specific species.

**MATERIAL AND METHODS**

**Data sets**

Sample collection and DNA sequencing for the zooplankton data set

The *Daphnia*, sediments and freshwater samples were isolated from the Agelsee pond, near Frauenfeld in Switzerland (47°55′ N, 8°86′ E) in June 2012. *Daphnia* are aquatic crustaceans of 0.5–5 mm body length. In our study pond, three species are sympatric: *Daphnia magna*, *D. pulex* and *D. longispina*. But we focused on *D. magna* as it outnumbers the others 10 to 1. *Daphnia* samples were collected using plankton net with 200 μm mesh size and kept in pond water during transport to the laboratory. Within 5 h, *D. magna* specimens were separated from other plankton species, washed in autoclaved medium and individually frozen at −20 °C in buffer for later DNA extraction. Samples from the top layer (5 cm) of the pond sediments were collected with a hand-held dreg, drained of standing water and stored 24 h at 4 °C in the dark before DNA extraction. Water samples were collected at the *Daphnia* sampling site and microbes successively filtered through 52-, 19- and 7-μm filters. The additional 3-, 1- and 0.2-μm filters were used for DNA extraction. Total genomic DNA was extracted using the CTAB method for the *Daphnia* and water samples and the PowerSoil kit (MoBio, Carlsbad, CA, USA), following the manufacturer’s instructions, for the sediment samples.

The V3–V5 region of the 16S rRNA gene (ca. 640 nucleotides) was PCR amplified using the following forward and reverse primers: 5′-ACACGGYCCARACTCTCCTAC-3′ (positions 327–345) and 5′-GTGWTTAATTGCWGCAA-3′ (positions 951–969). Amplicons from the different samples were pooled at equimolar ratio for multiplexing. Amplicon libraries were sequenced on a GS FLX instrument using LibL Titanium chemistry (454 Life Sciences, Roche) at Microsynth (Balgach, Switzerland).

**Bioinformatics pipeline for zooplankton data set**

QIIME 1.6 pipeline (Caporaso et al. 2010a) was used for bioinformatics analyses. Reads were filtered based on their length (> 500, < 580) and average Phred score (> 30). They were then chimera filtered using de novo chimera detection and clustered into operational taxonomic units (OTUs) at 97% sequence identity level using USEARCH (Edgar 2010) with QIIME default parameters. Taxonomic assignment was done using the RDP classifier (Wang et al. 2007) with the Greengenes database, version 12_10 (McDonald et al. 2011). The most abundant sequence of OTU cluster was chosen as its representative and sequences were aligned with PyNAST (Caporaso et al. 2010b). A phylogenetic tree was then built using FastTree (Price et al. 2010) for computation of UniFrac distances (Lozupone et al. 2007). OTUs represented by a single read were filtered out from the count table.

**Human microbiota data set**

The human microbiome data set is described in and was collected by Costello et al. (2009): filtered reads were downloaded from the European Nucleotide Archive (www.ebi.ac.uk/ena/data/view/ERA000159) and OTU picking was performed using the analysis pipeline detailed in Costello et al. (2009). Singletons were filtered out.

**Global patterns data set**

The ‘Global Patterns’ data set of environmental samples was obtained from Caporaso et al. (2011) and is directly available as OTU count tables from the R phyloseq package (McMurdie & Holmes 2013). Singletons were filtered out.

**Specificity index and permutation test**

Here, we consider a quantitative measure for the specificity of a species (or OTU) in a focal habitat, based on a reinterpretation of the indicator values of Dufrene & Legendre (1997). A high value indicates that this species is found predominantly in this focal habitat. We consider a microbiota count table \( M = (a_{ij}) \) where \( a_{ij} \) is the count, or relative abundance, of species \( i \) in sample \( j \). We use OTUs as proxy for species. The microbiota samples originate from \( H \) different habitats (e.g. different locations, different hosts) and there are \( S^h \) samples from habitat \( h \). We note \( S^h \) the number of samples from habitat \( h \) where species \( i \) is present. The local specificity index \( \lambda^h_i \) of species \( i \) in habitat \( h \) is defined as:

\[
\lambda^h_i = A^h_i \times B^h_i
\]

where

\[
A^h_i = \frac{S^h_i}{S^h}, \quad B^h_i = \frac{\langle a_{ij} \rangle^h}{\sum_{h=1}^{H} \langle a_{ij} \rangle^h} \quad \text{and} \quad \langle a_{ij} \rangle^h = \frac{\sum_{i=1}^{S^h} a_{ij}}{S^h}
\]

\( A^h_i \) is the prevalence of species \( i \) in habitat \( h \), i.e. the fraction of samples from habitat \( h \) where species \( i \) was found. In case
of limited sampling, it should be understood as discoverability, especially for rare species. \((a_i)^h\) is the average local abundance of species \(i\) in habitat \(h\) and \(B^h_i\) is the fraction of the share of habitat \(h\) in the total population of species \(i\), i.e., summed across habitats. \(\Lambda_i^h\) ranges from 0 to 1. A value of 0 means that the species is never found in habitat \(h\), whereas a value of 1 means that the species is always and only found in that habitat, i.e. the species is a perfect indicator of that habitat. Unlike strict definitions of specificity, that use only presence/absence information, our index uses the full abundance spectrum and is therefore robust to contamination and differences in sequencing depths. \(\Lambda_i^h\) is the same as INDVAL but for one change: INDVAL keeps only the maximum local specificity across habitats (INDVAL(i) = \(\max_{h} \Lambda_i^h\)) to have one value per species, whereas we keep all values to have one value per pair (species, habitat).

Since \(B^h_i\) requires the computation of the total count of species \(i\), the abundances must be on the same scale in different samples for the index to be meaningful. We use a double normalisation of abundances to avoid the undue effect of individual rich samples and sample rich habitats. Abundances are first transformed to relative abundance (sample-level normalisation) and then averaged within each habitat (habitat-level normalisation) to produce \(\langle a_i\rangle^h\).

The variability of \(\Lambda_i^h\) is assessed by stratified bootstrap. A new data set is created from the original by sampling communities with replacement within each habitat (the strata). The specificity index is computed on this data set and the process is repeated 999 times to estimate the variability of \(\Lambda_i^h\). The significance of \(\Lambda_i^h\) is also assessed by bootstrap resampling: original communities are resampled with replacement and randomly assigned to habitats (habitat permutation). The specificity index is computed on each bootstrapped data set and the process is repeated 999 times. These values represent the distribution of \(\Lambda_i^h\) under the null hypothesis of no association between local abundances and habitats. The traditional way of assessing the significance of \(\Lambda_i^h\) would be to compare it to this distribution. We are however interested in the significance of high specificities and therefore compare \(\Lambda_i^h\) to the null distribution of \(\max_{h} \Lambda_i^h\) rather than \(\Lambda_i^h\). In other words, the observed specificity is compared to the maximum randomised specificity across all habitats. This corrects for the multiplicity of habitats and means that a species is significantly specific only if more specific to a habitat than to any other one. Local specificities, permutation tests and standard error computations were performed using custom R code and the R phylseq package (McMurdie & Holmes 2013).

RESULTS

Zooplankton

This data set includes 32 microbiota samples from one pond in Switzerland, but corresponding to three different habitats: microbiota associated with the planktonic crustacean D. magna (11 samples), microbes in the free water (15 samples) and microbes from the sediment surface of the pond (6 samples). These three habitats are interconnected: D. magna filter water for uptake of food, water and sediments are in close contact and Daphnia browse over the sediment surface to enrich their food (Horton et al. 1979; Ebert 2005). Furthermore, Daphnia resting eggs (euphii) diapause in the sediments.

We used the zooplankton data set to derive abundance-specificity curves (Fig. 1) using our local specificity index. We find that on average the more abundant a taxon is in a habitat, the more specific it is to this habitat. Furthermore, the positive abundance-specificity relationship observed is significantly steeper than also positive random relationship (grey dots and dark grey curves in the three panels of Fig. 1). The latter arises because universal species with heterogeneous abundances across samples have a skewed abundance distribution with many small values and a few high ones. Chance grouping of high values in the same habitat during the randomisation scheme shifts the distribution of randomised specificities upwards compared to species with the same overall abundance but a more even distribution across samples. The upward trend of the specificity curve is nevertheless significantly steeper in the non-randomised than in the randomised data sets (Fig. 1): the most locally abundant species are highly specific to their habitat.

In parallel, we studied the diversity of samples using either Bray–Curtis distances, which account for differences in terms of species composition and abundances, or weighted UniFrac (wUF) distances, which also account for phylogenetic relatedness between species, combined with multidimensional scaling. The results (Fig. 2) show that the samples cluster neatly by habitats for the Bray–Curtis distances (Fig. 2, left) but in a less pronounced way for wUF distances (Fig. 2, right). The first axis distinguishes sediments from water and Daphnia. Daphnia and water samples then form a Daphnia-Water gradient along the second axis.

We evaluated the robustness of our findings with respect to copy number variation in SSU-rRNA genes across bacterial taxa (Vetrovsky & Baldrian 2013). Copy numbers of this gene vary from 1 to 15 in sequenced microbes and thus may influence the calculation of relative frequencies. Vetrovsky & Baldrian (2013) provided a copy number prediction scheme based on the average copy number at a given taxonomic level and estimated the average error to be 38% when predictions are made at the phylum level. We corrected observed species counts using either (1) average phylum-level (class in case of the abundant Proteobacteria) copy number or a (2) random copy number drawn from a truncated Gaussian, approximating the copy number distribution observed within each phylum (class in case of the abundant Proteobacteria). Our results did not change (Fig. S1). A further source of bias of amplicon-based microbiota studies is the classification of species as OTUs, which is built on a similarity threshold (typically 97% similarity). This threshold impacts specificity and abundance of microbial taxa (Youngblut et al. 2013): all things remain unchanged, higher thresholds lead to more OTUs, with lower abundances and higher specificities. However, using thresholds ranging from 91 to 99% identity does not change our finding of a positive relationship between abundance and specificity (Fig. S2).
Human microbiome data set

The human body is host to diverse bacterial communities whose collective population largely outnumbers human cells (Costello et al. 2009). We analyse the publicly available data set built by Costello et al. (2009), who sampled microbial communities across 26 body sites on seven subjects at two time points. Sampling sites were grouped by body habitat as in the original study (Costello et al. 2009): Skin, External Auditory Canal (EAC), Gut, Oral Cavity, Hair, Nostril and Hair. Skin is the most diverse habitat with 18 sites ranging from sole of foot to forehead, while other environments included much less sampling sites (minimum 2). This data set is interesting for our purpose as the habitats are very diverse but in close proximity and therefore allow for exchange of bacteria. Furthermore, the human microbiota has been extensively studied and is well characterised. Previous studies show that body habitats differ in their overall composition, based on ordination plots of UniFrac distances (Costello et al. 2009).

All body habitats exhibit a positive and significant abundance–specificity relationship, with the exception of the skin (Fig. 3). Skin is considered here as a single body habitat as in the original study but really encompasses a variety of habitats with different environmental variables. For example, parts of the skin such as armpits, soles of feet, hand palm and index finger differ in key environmental variables such as humidity, temperature and light exposure. An alternative grouping of skin sampling sites that respect body symmetry (see Supporting information for details), splits skin into six habitats. Using this classification, we find a stronger abundance–specificity relationship (Fig. S3). The finding of the abundance–specificity relationship is also stable across time: the analysis performed on each time point individually shows the same
positive trend (Fig. S4) and the most abundant taxa are conserved across time (Fig. S5).

Global patterns
The global patterns (GP) data set consist of 25 environmental samples gathered by Caporaso et al. (2011) with an average depth of about a million reads per sample. The samples are distributed across eight different habitats, with two to four replicates per habitat: human faeces (n = 3), hand palm skin (n = 3) and dorsal tongue surface (n = 3), faecal samples from human twins (n = 2), soil (n = 3), freshwater and freshwater sediments (n = 5), ocean water (n = 3) and marine sediments (n = 3). We left out the artificially assembled mock communities as they are unlikely to capture natural abundance-specificity relationships.

Like in the previous data sets, all habitats exhibit a strong abundance-specificity relationship: abundant species have on average a high local specificity. In this data set we note that many low frequency (<1e-4) taxa have high local specificity (Fig. 4), but many more have low specificity. A rarefaction analysis of the samples (Fig. S6) confirms that the relationship is preserved at much shallower sampling depths (5000 reads per samples). It also shows that species with high local specificity (>0.9) are easy to capture: 5000 reads capture on average 40% of the specific species accounting for 96% per cent of the specific population.

DISCUSSION
The specificity of species from microbial communities to a given habitat type is positively correlated with their local abundance. This relationship is robust to various technical and biological aspects of microbiota assessment and analysis, such as differences in 16S rRNA gene copy numbers, OTU definition threshold and sampling depth. The diverse data sets used to test this relationship reveal that even when habitats have frequent opportunities for microbial exchange, microbial compositions remain distinct. The rich global pattern data set further shows that abundant taxa are highly specific to their habitat even though habitat-specific taxa are found everywhere across the abundance scale. Our communities differ clearly by habitat in terms of specific and abundant species, even when those habitats interact with each other through known overlaps and migration from a regional pool of taxa, like in the zooplankton data set. Local microbiota may share some taxa, like the planktonic freshwater bacteria from the genus Limnohabitans that accounts on average for 15% of both Daphnia and water samples but we found only one instance of shared abundant taxa. The fraction of shared abundant taxa varies with the differentiation between the

Figure 3 Positive relationship between local abundance and specificity in the human microbiome (Costello et al. 2009). Details of the legend are identical to Figure 1. Note that ‘skin’ encompasses 18 body sites corresponding to very different environmental variables and does not form a coherent habitat.

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Habitats being compared. In the absence of differentiation between habitats, the Neutral Community Model (NCM) predicts that taxa with high mean abundance should have high occupancy and variance inversely proportional to the migration rate (Sloan et al. 2006). This abundance-occupancy trend has been reported in Nemergut et al. (2011) but was not observed in another large-scale soil study (Barberan et al. 2012). However, if habitats are well differentiated, as in Barberan et al. (2012) or this study, NCM does not rule out a mix of (1) habitat specialists with high abundances and occupancies in their habitat but low ones in other habitats and (2) habitat generalists with low abundances and occupancies in all habitats. In this case, differentiation leads to habitat-exclusive abundant taxa (Gravel et al. 2006).

The monotonic increase in the abundance–specificity curves at the habitat level suggests that habitat filtering plays an important role in shaping local communities and that locally abundant taxa are likely to be specific to their habitat. Our finding means that β-diversity is driven by taxa that are both specific and locally abundant. As a consequence, β-diversity can be estimated without a large sampling effort. It explains why wUF and other weighted (semi)metrics (Graham & Fine 2008) or quantitative β-diversity measures are unexpectedly good at separating different habitats even at relatively low sampling depth (~2000 reads per sample) and why low-depth sampling captures the same relationship among samples as does deep sampling, as noted by Caporaso et al. (2011) on the GP data set. Our analysis also revealed that taxa specific to a habitat do not need to be strongly phylogenetically differentiated. Figure 2 shows that water and Daphnia specific taxa are phylogenetically close, but taxonomically different; the difference may lie in their ecological function, as was previously suggested in this setting (Qi et al. 2009) and in other studies (Burke et al. 2011a; Jiang et al. 2012).
Previous studies provide examples of abundance–specificity relationships (although not formally analysed) and suggest diverse mechanisms through which this relationship may come about. For example, Barberán et al. (2012) report that specialists, especially in desert and Antarctic soils, are more abundant than predicted based on their persistence across habitats. Species with active hydrocarbon degradation capabilities are not detectable in sediments, but become very abundant (6–18%) after oil spill (Mason et al. 2014). Habitat specialists are locally abundant at the extremes of salinity gradients (Logares et al. 2013) and coastal marine environment are reproducibly dominated by taxa with limited distributions (Fortunato et al. 2013). In these examples, specificity may be achieved by high growth rates of specialists. Host-associated communities also provide interesting examples: sponge species share most of their OTUs but harbour species-specific members that make up to 30% of the communities (Reveillaud et al. 2014). Moitinho-Silva et al. (2014) suggest that ‘dominant OTUs do account for sponge communities’. Likewise resident communities of Daphnia (this study) and algal surfaces (Burke et al. 2011b) are substantially made up of bacteria that are rare in the surrounding waters. In those examples, specificity may result from species sorting along a nutrient gradient but specificity can also result from active selection from the host as in the squid-Vibrio system (Kremer et al. 2013). Finally, chemical-physical barriers may limit competition, prevent invasion and therefore favour resident taxa. Jones & McMahon (2009) show that immigration of atmospheric taxa into lakes that are otherwise closed systems is essentially erased by habitat filtering: the studied lakes are dominated by abundant and specific species. Similarly, bile production is thought to protect the human gut from foreign microbe species. Alterations caused by liver cirrhosis can lead to invasion and displacement of up to 40% of the resident community by a few commensal oral species that are absent from healthy communities (Qin et al. 2014).

Community assembly

Community assembly involves four distinct kinds of processes: selection, drift, speciation and migration (Vellend 2010). They all have been shown to play a role in microbial community assembly (Costello et al. 2012). Our findings suggest that even when habitats are in close spatial proximity and species migration is possible, selection plays a comparatively central role in determining the composition of local communities, as suggested by the Baas-Becking hypothesis, and may mitigate the effects of migration (Gravel et al. 2006; Jones & McMahon 2009). It is also well known that assemblages are typically dominated by a few abundant taxa while most taxa are rare and that neutral assembly can lead to such a skew (Sloan et al. 2006). Our observations may be a consequence of (1) habitat differentiation, (2) selection for specialist taxa across habitats (for example in the form of habitat filtering or high growth rate in the favoured habitat of a taxon), and (3) neutral assembly processes within habitat. Selection implies that habitat-level relative abundances would be skewed in favour of specialists that would then be abundant through the neutral assembly of individual communities. The skew should increase with increased migration and increased differentiation between habitats, as observed in this study.

Deterministic components of assembly processes have been suggested elsewhere (Vellend 2010; Stegen et al. 2012). Connolly et al. (2014) deduced, from studying abundance-rank curves of over 1000 communities from the marine biosphere, that lognormal abundances are a better fit than the neutral Poisson-gamma distribution. They suggest differences between species (e.g. in terms of growth rates, niche sizes, etc.) as mechanisms leading to lognormal abundances. Recent work (Harris et al. 2014) in statistical ecology looking at the fit of neutral models (Hubbell 2001) to human gut microbiota reach similar conclusions: neutrality is rejected from abundance data at all but low taxonomic ranks (genus). Overall, habitat filtering selects specialist taxa, probably based on markedly different metabolic and functional roles (Fierer et al. 2007; Philippot et al. 2010) and among those, some taxa with selective advantage become dominant in local communities.

Limits and potential confounding factors

Quantitative metagenomics is a powerful approach to study microbial communities because it enables us to obtain data sets with very deep sampling in short time periods and high spatial resolution. However, it is subject to a variety of biases that may influence the inference obtained from the analysis. Standard methodology assumes that each microbial taxon has the same number of 16S rDNA genes, although it is known that species differ from 1 to 15 copies of this gene (Vetrovsky & Baldrian 2013). Our analyses suggest that our results do not suffer from a serious bias caused by copy number variation. Assumptions about the definition of the OTU as a surrogate for a species definition may also influence the interpretation of microbiota studies because it affects both the species classification and their abundance. Again, our results are robust with regard to this assumption.

A further bias arises from the complex interplay between niche and habitat. Niche is defined by a volume in a space of mostly unknown biotic and abiotic environmental variables (Hutchinson 1957). Habitats may be comprised of a single niche, or may include multiple niches. For example, the human skin is diverse and covers many environmental variables. Not surprisingly, the skin habitat does not show a strong abundance–specificity relationship. After a custom classification of skin sampling sites based on hierarchical clustering, body geography and symmetry, we recovered more narrowly defined habitats and stronger abundance–specificity relationships. In conducting a custom classification we took care to avoid circularity and to artificially increase the abundance–specificity trend by clustering the samples using only presence/absence data, not abundance or prevalence data. Additionally, the skin habitat in the GP data set consists only of hand palms and exhibits a strong association from the start. The same observation applies for the butterfly gut microbiota study of Hammer et al. (2014). The gut at different developmental stages corresponds to different diets. It is dominated by different taxa, suggesting that host diet is a key environmental variable for niches of gut microbiota (see also Delsuc et al. 2014).
Our local specificity index captures relative specificity rather than endemism. Our double normalisation, standard in microbiota studies, assumes (1) that each habitat has a carrying capacity shared by all species and (2) that all habitats contribute equally to the regional pool of taxa. When these assumptions are violated we need to be careful with the interpretation of the results. For example, in a continent-island scenario, a species can be locally very specific to a small habitat (an island) and yet be mostly found in the much larger habitat (the continent) (see Supporting information). Our index therefore works only for relative specificity rather than endemism. An important consequence is that the abundance–specificity relationship is influenced by all habitats included in the study; it is stronger if habitats are very distinct, like in the GPs, and weaker but still significant for more similar habitats, like in the Human Microbiome.

Finally, our method counts all zeroes as true absence when computing prevalence in a given habitat. Replacing zeroes by probability of absence, estimated from zero-inflated distributions such as Gaussian or Negative Binomial (McMurdie & Holmes 2014) should alleviate this problem. This limitation induces a downward bias of specificity values for rare species but does not weaken the abundance–specificity relationship, as confirmed by the rarefaction analysis of the GP data set (Fig. S6). Furthermore, the high depths of the GP data set almost guarantee discovery of taxa with even low frequency (> 1e-5) and confirm that the relationship is not an artefact of rare taxa having low specificity simply because of discoverability issues.

CONCLUSION

The main finding of our analysis is the on average high specificity of abundant microbe species in microbiota samples from well differentiated habitats. It means that (1) shallow sampling is sufficient to recover compositional differences and (2) locally abundant species, many of which are habitat specific, are likely to perform habitat-specific ecological functions. This finding has implications for other studies. By reducing the sampling effort per site, it will be possible to increase the replication effort and the number of habitats studied. For example, we suggest that in studies of $\beta$-diversity surveys, one should aim for moderate sequencing depth, but for a large sampling width. On the other hand, the existence of specific species across the abundance scale means that $\alpha$-diversity surveys capture most specific species with shallow sampling, but cannot span deep sequencing as some species are both rare and specific.

Furthermore, our study suggests that abundant species are on average good predictors of environmental conditions and that the search for microbial indicators should start with abundant species. An exciting extension in this direction would be possible if transcriptional activity is positively correlated with abundance or skewed towards abundant taxa (Motininho-Silva et al. 2014). In this case, shallow sampling might be sufficient to identify the genes of key biological processes provided by microbial ecosystems (Burke et al. 2011a; Jiang et al. 2012; Qin et al. 2014).

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AUTHORSHIP

MM, SP and DE designed the study. SP collected the data. MM performed the modelling work and analysed the data. MM wrote the first draft of the manuscript, and all authors contributed substantially to revisions.

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