A global network of coexisting microbes from environmental and whole-genome sequence data

Samuel Chaffron,1 Hubert Rehrauer,2 Jakob Pernthaler,3 and Christian von Mering1,4

1Institute of Molecular Life Sciences and Swiss Institute of Bioinformatics, University of Zurich, CH-8057 Zürich, Switzerland; 2Functional Genomics Center Zurich, University of Zurich and ETH Zürich, CH-8057 Zürich, Switzerland; 3Limnological Station of the Institute of Plant Biology, University of Zürich, CH-8802 Kilchberg, Switzerland

Microbes are the most abundant and diverse organisms on Earth. In contrast to macroscopic organisms, their environmental preferences and ecological interdependencies remain difficult to assess, requiring laborious molecular surveys at diverse sampling sites. Here, we present a global meta-analysis of previously sampled microbial lineages in the environment. We grouped publicly available 16S ribosomal RNA sequences into operational taxonomic units at various levels of resolution and systematically searched these for co-occurrence across environments. Naturally occurring microbes, indeed, exhibited numerous, significant interlineage associations. These ranged from relatively specific groupings encompassing only a few lineages, to larger assemblages of microbes with shared habitat preferences. Many of the coexisting lineages were phylogenetically closely related, but a significant number of distant associations were observed as well. The increased availability of completely sequenced genomes allowed us, for the first time, to search for genomic correlates of such ecological associations. Genomes from coexisting microbes tended to be more similar than expected by chance, both with respect to pathway content and genome size, and outliers from these trends are discussed. We hypothesize that groupings of lineages are often ancient, and that they may have significantly impacted on genome evolution.

[Supplemental material is available online at http://www.genome.org/]

Symbiosis—as defined in its broadest sense (de Bary 1879; Saffo 1993)—is widespread in nature, ranging from obligate mutualistic partnerships to commensalism to clearly detrimental, parasitic interactions (Paracer and Ahmadian 2000). The phenomenon is not restricted to a particular domain of life, but can occur, for instance, between bacteria, archaea, and protists, which, in turn, can live together inside a specific animal host (Brauman et al. 1992; Tokura et al. 2000). Many instances of symbiosis are known, but they are not always understood mechanistically. The situation may not always be stable either: Symbionts may “cheat,” and/or compete among each other for specific resources (Palmer et al. 2003; Ferriere et al. 2007; Johnstone and Bshary 2008).

Leaving aside macroscopic organisms, symbiosis and local coexistence among single-celled microbes are even less well characterized. The extent, specificity, and stability of microbial associations are difficult to assess systematically in the environment, since elaborate staining procedures and/or molecular sequencing are needed in order to detect and differentiate between microbial lineages in situ. Nevertheless, several close partnerships between microbial species have already been identified. These include consortia of methanogen-oxidizing archaea and sulfate-reducing bacteria (AOM, “anaerobic oxidation of methane”) (Boetius et al. 2000; Caldwell et al. 2008; Knittel and Boetius 2009); consortia of phototrophic green sulfur bacteria surrounding motile beta-proteobacteria (Overmann and Schubert 2002; Wanner et al. 2008); consortia of sulfate reducers, sulfate oxidizers, and other lineages inside marine, gutless oligochaete worms (Dubilier et al. 2001; Woyke et al. 2006; Ruehland et al. 2008); and consortia of extremophilic lineages conducting ferrous iron oxidation in acidic pyrite mine run-offs (Tyson et al. 2004). Such groupings probably do not constitute “symbioses” in a classical sense (Saffo 1993), but they are typically interpreted as syntrophic associations in which one partner consumes metabolites produced by the other. In addition, predatory and parasitic relationships are also known. An example for the latter is Nanoarchaeum equitans, a small archaeon that appears to be an obligate parasite of another archaeal species (Huber et al. 2002; Forterre et al. 2009). Despite such specific findings, the discovery of microbial associations has so far been largely interest-driven (or even fortuitous), meaning that a comprehensive picture of microbial coexistence has yet to emerge.

The notion that microbes in the environment perhaps exist in a less solitary manner than commonly assumed is also supported by the rapidly accumulating knowledge on intra- and interspecies microbial communication (Ryan and Dow 2008; Shank and Kolter 2009). Essential activities of single species such as nutrient uptake, biofilm formation, or cellular differentiation can be organized and synchronized by communication and cooperation (Parsek and Greenberg 2005; Waters and Bassler 2005; Kolter and Greenberg 2006; Gilbs et al. 2008; Ng and Bassler 2009). While it is less clear whether and to what extent microbes may interact with other species via specific communication, some bacteria are known to “eavesdrop” and to even respond to signals that they cannot themselves generate (Visick and Fuqua 2005). In addition, an interspecies relationship has been shown to evolve and quickly deepen in a laboratory evolution experiment (Hansen et al. 2007; Harcombe 2010).

Apart from the few cases of well-described, specific interactions, relatively little is known about how natural microbial assemblages form and how they are structured, if at all (Ruan et al. 2006; Horner-Devine et al. 2007; Fuhrman and Steele 2008; Raes and Bork 2008; Fuhrman 2009). They are often taxonomically highly complex and can encompass hundreds of different species, and at least some aspects of the composition of any given community are thought to be based on historical contingency (Martiny et al. 2006). Moreover, naturally occurring communities are difficult.
to reassemble and/or study under controlled conditions in the laboratory, since most of the constituting lineages are not available in pure cultures (typically more than 95% of species present in a given sample cannot be cultivated) (Staley and Konopka 1985). The difficulties in cultivating microbes have often been linked to slow growth and unknown nutritional requirements, but might also be partly attributed to their syntecology—for example, reflecting a need to coexist within a biofilm or to aggregate together with partner species in order to ameliorate adhesion (Min and Rickard 2009) or to dispose of otherwise inhibitory metabolic products.

Since the establishment of the first comprehensive microbial phylogeny using the 16S rRNA gene (Fox et al. 1980) and the invention of techniques for rapidly generating large blocks of 16S rRNA sequence data (Lane et al. 1985; Giovannoni et al. 1990; Ward et al. 1990), a great variety of environments have been sampled to study microbial diversity in situ. Today, the 16S rRNA gene remains the marker of choice for identifying microbes in their environments, and the size of databases dedicated to this gene is growing exponentially (Desantis et al. 2006; Puersse et al. 2007; Cole et al. 2009). In addition, environmental sequences are increasingly being annotated with contextual information (e.g., geographic position, temperature). An important effort to define and standardize such sequence meta-data has been initiated within the Genomic Standards Consortium (Field et al. 2008a,b), specifically by developing the MIENS standard (minimum information about an environmental sequence). However, the existing annotations of many of the legacy sequences in the databases will have to be migrated to such standards, which requires considerable effort. The results of such efforts are increasingly being made available in integrated resources such as CAMERA (Seshadri et al. 2007), IMG/M (Markowitz et al. 2008), and megDB (Kottmann et al. 2010), but at present only a small minority of 16S rRNA sequences have geo-referencing or other contextual information.

Using 16S RNA sequences in combination with other data, classical ecological questions including species (co-)occurrence and diversity have also been addressed extensively in microbes (Bell et al. 2005; Langenheder et al. 2006; Ruan et al. 2006; Horner-Devine et al. 2007; Smith 2007; Langenheder and Prosser 2008). In doing so, many of the concepts that have originally been developed for macroscopic organisms have been adapted and applied to microbes. However, these studies have mostly focused on one specific environment, or one specific lineage, at a time (e.g., Alonso et al. 2007; Newton et al. 2007; Fuhrman and Steele 2008) (this way, ecological questions can be studied in a more defined setup). What has not been addressed much, so far, is the global partitioning of microbial lineages among all sampled environments. Here, we take a first step in this direction, by systematically studying microbial lineages among all sampled environments.

Results and Discussion

In order to comprehensively characterize the occurrence of microbial lineages in the environment, we first grouped publicly available, full-length 16S rRNA sequences at various levels of sequence identity, thereby creating unsupervised sets of “operational taxonomic units” (OTUs; see Methods for details). Each OTU was assigned a taxonomic annotation that reflected the consensus of its member sequences, and a single sequence was chosen to represent each OTU in subsequent sequence comparisons. Next, we comprehensively compiled environmental “sampling events” of 16S sequences; such an event is defined here as a unique combination of submitting authors, project title, and isolation source, as annotated in the respective database records. We assumed that sequence entries for which all three fields are exactly identical were sampled together, at a given site. Our procedure (Fig. 1) thus resulted in a large matrix that connects OTUs to environmental sampling events (Table 1). Depending on the OTU definition, this matrix contained roughly between 700 and 5000 distinct OTUs, which were mapped to roughly 3000 distinct sampling events (we only retained sampling events that encompassed at least two OTUs, and conversely, only OTUs that were observed in at least three sampling events).

Next, we examined this matrix for any non-random assortment of OTUs to environments, which would manifest itself as groups of OTUs observed together more often than expected by chance. Our underlying null model is that of global, random dispersal of lineages across environments (Harvey et al. 1983; Finlay 2002; Kunin et al. 2008a; Hubert et al. 2009), and essentially corresponds to the first part of Baas Becking’s enigmatic statement, “Everything is everywhere, but, the environment selects” (de Wit and Bouvier 2006). While this null model is clearly not applicable for macroscopic organisms with distinct biogeographic distribution patterns, it does represent the simplest default assumption for microbes, and it is appropriate for the very large geographical and temporal scales that we consider here. By computing the hypergeometric probability of pairwise co-occurrences and correcting for multiple testing, we found that, indeed, a large number of statistically significant associations between OTUs can be observed, irrespective of the precise choice of OTU definition cutoff (Fig. 2; Supplemental Fig. S1; Table 1). A concrete example for such an association is shown in Figure 1B (data from Sorensen et al. 2005; Baati et al. 2008; Isenbarger et al. 2008; Sahl et al. 2008; R Amdouni, E Ammar, H Baati, N Gharsallah, and A Sghir, unpubl.): a well-characterized lineage of *Cyanobacteria* (belonging to the halophilic *Eukhalothece*) (Garcia-Pichel et al. 1998) was observed to be associated with an uncharacterized lineage having no cultivated or named representatives (a monophyletic sister group of the *Psychroflexus* lineage [Bacteroidetes]). This particular association is based on three independent sampling events in which both lineages had been observed together, by three distinct laboratories in three distinct countries. Considering that the OTU definition in this case is relatively narrow (97%) and that this association occurs against a backdrop of about 2800 sampling events covering more than 5000 OTUs, the observation becomes highly significant ($P < 3 \times 10^{-15}$, after multiple testing adjustment). Overall, several thousand of such associations could be identified. To assess the effects of potential biases in the sampling data, and in order to estimate our false discovery rate (FDR) empirically, we performed a conservative randomization of our data—by keeping constant the size distributions of both sampling events and OTUs, but shuffling the connections between OTUs and sampling sites. This resulted in a reduction of the number of reported associations by greater than 99% for most of the OTU definition cutoffs (Table 1), which translates to FDRs of ~1%, except at very broad OTU definitions (i.e., when setting the OTU clustering cutoff to 85% sequence identity).
The few remaining false-positive associations were observed mainly among widely sampled lineages known to occur inside the mammalian digestive tract; this likely reflects the strong study bias toward 16S gene sequences of this habitat (Ley et al. 2008; Hamady and Knight 2009). In addition to assessing the statistical significance, we also computed a “specificity” value (or “association strength”) for all OTU pairs. This value corresponds to the Jaccard similarity; it is 1.0 if a pair of OTUs is always observed together (but never separately), and zero for a pair of OTUs that is always observed in distinct environments, but never together. Remarkably, we found associations at both extremes of specificity (i.e., close to 1.0 or close to zero) (see Supplemental Fig. S3 for the overall distribution). An example for the former is shown in Supplemental Figure S1A: a previously undescribed bacterial OTU (a sublineage of the candidate division JS1) was observed together with a specific Methanococcales lineage in marine sediments, again by three distinct laboratories (once in the Mediterranean, twice in the Gulf of Mexico at distinct sites). However, in this case, the two lineages were never found separately, in any of the 2800 environment samplings we studied. The likelihood of observing such a specific association by chance is again very low ($P = 1.1 \times 10^{-7}$ after correction for multiple testing). An example for a less specific but nevertheless highly significant association is shown in Supplemental Figure S1B: A lineage of gamma-Proteobacteria (nosocomial pathogens from the genus Stenotrophomonas) was frequently observed together with a lineage of Bacilli (genus Staphylococcus). The two lineages were sampled together 10 times—by seven distinct laboratories—in various air samples, skin samples, dust, and on Chinese cabbage. The association is highly significant ($P < 10^{-9}$ after correction for multiple testing), but less specific: Both lineages have also been observed separately (in 32 and 17 sampling sites, respectively). Not all of the latter observations were related to skin samples. Stenotrophomonas, for example, may also form distinct blooms in shallow coastal lagoons (Piccini et al. 2006). While co-occurrence alone cannot offer any mechanistic explanation for lineage associations, the additional information in the specificity of an association does provide a constraint when discussing possible scenarios (obligatory mutualism, for example, would be expected to result in a high association specificity). Barring any additional information, we did choose to interpret our observed associations conservatively, by assuming that they for the most part simply reflect shared or overlapping niche preferences. Instances of undescribed, specific mutualisms and parasitisms are presumably contained within our findings, but additional experimental follow-ups will be required for a detailed characterization of such interactions (Orphan 2009). That notwithstanding, this first part of our analysis already provides an empirical base for discovery and allows us to explore more specific hypotheses about the reasons for the coexistence of sets of uncultured genotypes.

Next, we searched our observed co-occurrence relations for previously known microbial associations (Supplemental Fig. S1).
While we did not recover the known association between the Nanoarchaeum and Ignicoccus lineages, nor the Chlororhabdus consortium, we did find strong evidence for AOM consortia (Supplemental Fig. S1D). We also observed the known association between the lineages Leptospirillum (phylum Nitrospira) and Acidithiobacillus (phylum Proteobacteria), both of which are known to thrive in acidic bioleaching environments. In this case, the association we found was remarkably strong and specific: Out of 21 independent observations of Leptospirillum (by 18 distinct author teams in various settings), all but a single one also included observations of Acidithiobacillus (i.e., 20 out of 21; \( P < 10^{-35} \)) (Supplemental Fig. S1C) (in this case, the OTU clustering distance was 90%). Remarkably, this association appeared to be somewhat asymmetric: Acidithiobacillus did occur occasionally without its partner (in an additional 18 sampling events), suggesting that the mutual dependencies might not be equally strong in both directions. As a further test of our associations, we conducted an independent co-occurrence search of microbial lineages in the published literature (Supplemental Fig. S5). The frequencies of co-mentions of species names in PubMed can, indeed, reveal ecological associations (Freilich et al. 2010), albeit limited to those lineages that are already validly named and for which cultivated type strains typically exist. We find that more than 70 of our lineages are co-occurring very specifically in certain marine sediments; they are from three distinct phyla, and each lineage is entirely uncharacterized. (A closer phylogenetic analysis revealed that the two Planctomycetes OTUs are related to each other, to the exclusion of other Planctomycetes lineages; they have been found also in other marine and freshwater environments, and our co-occurrence analysis thus defines a more restricted home context for these lineages.) Specific modules such as this example are striking and likely provide a first glimpse onto hitherto undescribed microbial consortia.

While our 16S-based OTUs provide fairly objective coverage of phylogenetic lineage space, they do not, in themselves, contain any information about molecular and ecological functions. We therefore attempted to represent each OTU by its best match in the exclusion of other taxa'' of vegetation analysis. One such possible partitioning is shown in Figure 2C; it conveys more information than a simple list of pairwise co-occurrences because it groups specific lineages, at the exclusion of others. Module formation can occur even if the various associations are not all highly significant (e.g., due to undersampling); this is because a certain fraction of missing or poorly scoring associations can be tolerated as long as the overall topology remains that of a tightly linked module. Furthermore, partitioning allows the annotation of keywords that describe the commonalities among the associated sampling sites of the various modules (Fig. 2C; see Methods). Among the modules, we observed intriguing cases where all or the majority of lineages have not been characterized before. An example is shown in Figure 3 (data from Hejlsberg et al. 2008; Harrison et al. 2009; Takeuchi et al. 2009; Ghosh et al. 2010)—five lineages that are co-occurring very specifically in certain marine sediments; they are from three distinct phyla, and each lineage is entirely uncharacterized. (A closer phylogenetic analysis revealed that the two Planctomycetes OTUs are related to each other, to the exclusion of other Planctomycetes lineages; they have been found also in other marine and freshwater environments, and our co-occurrence thus defines a more restricted home context for these lineages.)

The observed associations were not limited to pairwise co-occurrences. When plotting the associations as a graph, a densely connected network of OTUs emerged (Fig. 2A). The topology of that network is clearly nonrandom; it exhibits a high clustering coefficient, short average minimum path length, and a connectivity degree distribution that has no characteristic maximum (i.e., the network is roughly matching the “scale free, small-world” criteria) (Barabasi and Oltvai 2004). This topology suggests that the network can be meaningfully partitioned, and that doing so should reveal modules of densely connected microbial lineages; these might be regarded as the microbial equivalents of the “syn-taxa” of vegetation analysis. One such possible partitioning is shown in Figure 2C; it conveys more information than a simple list of pairwise co-occurrences because it groups specific lineages, at the exclusion of others. Module formation can occur even if the various associations are not all highly significant (e.g., due to undersampling); this is because a certain fraction of missing or poorly scoring associations can be tolerated as long as the overall topology remains that of a tightly linked module. Furthermore, partitioning allows the annotation of keywords that describe the commonalities among the associated sampling sites of the various modules (Fig. 2C; see Methods). Among the modules, we observed intriguing cases where all or the majority of lineages have not been characterized before. An example is shown in Figure 3 (data from Hejlsberg et al. 2008; Harrison et al. 2009; Takeuchi et al. 2009; Ghosh et al. 2010)—five lineages that are co-occurring very specifically in certain marine sediments; they are from three distinct phyla, and each lineage is entirely uncharacterized. (A closer phylogenetic analysis revealed that the two Planctomycetes OTUs are related to each other, to the exclusion of other Planctomycetes lineages; they have been found also in other marine and freshwater environments, and our co-occurrence thus defines a more restricted home context for these lineages.) Specific modules such as this example are striking and likely provide a first glimpse onto hitherto undescribed microbial consortia.
likely abundant (Pedros-Alio 2006) and that can be widely found and easily accessed (each OTU had to be sampled at least three times to be included here). And, second, there seems to be a notable stability of environmental habitat preferences among microbial lineages in general (Von Mering et al. 2007; see also below). This suggests that a sequenced strain may represent other members of its OTU in terms of its genomic content and aspects of its ecology even if it has diverged from them to some degree. We were able to map between 350 and 660 genomes to a subset of our OTUs (this depends on the OTU resolution; notably it also means that a significant fraction of sequenced genomes currently cannot be connected to an OTU that has been repeatedly observed in the environment). This mapping translates to between 200 and 410 significant partnerships for which genomic information is available for both partners, covering a small but significant fraction of all the instances of co-occurrence we detected. To our knowledge, this is the first time that a global, environmentally motivated association network between genomes has been constructed. We used this network to objectively assess potential constraints on genome evolution, which might be a consequence of...
Figure 3. Example of a novel, previously undescribed module of coexisting lineages. (A) Five distinct microbial lineages are shown; they belong to three different phyla and are defined at an OTU-clustering distance of 90% sequence identity at the 16S rRNA gene. The five lineages have been exclusively observed through environmentally sampled sequences and have not been named. (B) The table shows all occurrence counts of these lineages among our sampling data; the P-values indicated have been corrected for multiple testing, against the background of all lineages defined at 90%. Adjusted P-values (FDR; p) and odds ratios (or) are indicated. (*) The samples by Li et al. (2008) have been collected at distinct sites, covering a distance of more than 600 miles; collection was at different water depths (FDR; p) and odds ratios (or) are indicated. No named relatives in candidate phylum OP8.

The association of a genome to its preferred environment and to other lineages in that environment. We observed four highly significant trends among co-occurring genomes: They tend (1) to have more similar genome sizes, (2) to be more similar in GC content (i.e., the fraction of the genome consisting of guanine and cytosine), (3) to be more similar with respect to relative coverage of functional pathways, and (4) to be phylogenetically more closely related than randomly selected pairs of genomes (Fig. 4). The latter trend was also visible from 16S sequences alone (Supplemental Fig. S2). The trend to phylogenetic relatedness is presumably the easiest to rationalize: Pairs of ecologically associated lineages, which are also closely related phylogenetically, would arise naturally assuming that neither lineage had changed their habitat preferences since they split from their last common ancestor. We indeed observe this signal and detect that it extends surprisingly far back in time: Lineages that have diverged up to 10% at the 16S sequence identity level are still clearly enriched among environmentally associated pairs (Fig. 4A; the peak seen at 15% sequence divergence is largely due to a single, well-covered cluster; see Supplemental Figs. S6, S8). In principle, this relatedness signal could also explain our three other observations: Phylogenetically related genomes are known to exhibit similar GC contents, genome sizes, and functional composition. To assess this possibility, we tested these three signals for independence from the phylogenetic signal, by correcting for the underlying correlations as learned from randomly selected genome pairs (Fig. 4). In the case of GC content similarity, we find that the signal can, indeed, be largely explained by phylogenetic relatedness alone—it is not an independent observation. This would argue against environmental selection on GC content, at least at longer time scales, and it gives further support to algorithms that partition environmental sequences based on genomic signatures (McHardy and Rigooutos 2007; Mrazek 2009). In contrast, importantly, we observed that both genome size similarity and functional similarity could not be explained solely by phylogenetic relatedness. For example, while randomly selected pairs of genomes have genome sizes that can vary considerably, environmentally associated genome pairs tend to level off at ~20%–30% genome size difference, on average (Fig. 4E; P < 10⁻¹³). This is remarkable because it suggests that a given environment tends to select for a particular optimal genome size range, even across distinct lineages; furthermore, it suggests that lineages spend sufficient time in their preferred environments to allow for these optimal genome sizes to be selected for and maintained (against a mutational spectrum that is thought to be largely biased toward deletions in bacteria) (Mira et al. 2001; Nilsson et al. 2005). Our observation confirms what has been known anecdotally from a number of environments: Planktonic marine environments, for example, persistently select for small to very small genome sizes (Giovannoni et al. 2005; Ting et al. 2009), whereas soil microbes are often among those with the largest genomes. Our results are also in line with observations indicating different average genome sizes in distinct environments (Raes et al. 2007; Angly et al. 2009). Regarding the functional similarity of genomes, we likewise observe that it is much stronger than what would be expected based on relatedness alone (Fig. 4G). Here again, lineage-environment associations appear to be stable enough to allow selection for similar functional repertoires even in unrelated lineages. However, apart from a phylogenetic signal, functional similarities can also arise due to similarities in genome size (van Nimwegen 2003; Konstantinidis and Tiedje 2004; Ranae et al. 2004). When correcting for the dependency between genome size
and functional content, we again find that co-occurring genomes of identical size are much more similar in functional terms than expected (Fig. 5). In Figure 5, we not only plotted genome size and functional similarity, but also phylogenetic relatedness (by means of a color code). This reveals the expected, combined trends: Environmentally associated lineages that tend to be most similar in functional terms also tend to be those that are both, phylogenetically the most related and also the most similar in terms of genome size. Outliers from these trends should reveal interesting exceptions, inviting speculations on distinct ecological scenarios. We highlight a few of such extremes in Figure 5. The first example represents an outlier case because the two lineages are very closely

Figure 4. Coexisting lineages display similarities in genomic features. Here, we exclusively focus on co-occurring lineages for which completely sequenced genomes could be mapped to both partners (this genome mapping is globally visualized in Supplemental Fig. S6). Properties of such co-occurring genomes are compared, and contrasted against randomly paired genomes. (A) The distribution of 16S sequence divergence scores; shifted to the left for co-occurring genome pairs (i.e. they tend to be related phylogenetically). In panels E, F, and G, we test for independence between phylogenetic relatedness, and observations as shown in panels B, C, and D, respectively. Here, each dot denotes a pairwise genome comparison, and lines correspond to running medians.
related phylogenetically (both are Bacilli), yet they currently hold the record in terms of functional divergence. Note that the various species of Bacilli are very closely related phylogenetically and cannot easily be distinguished based on 16S alone (Vilas-Boas et al. 2007; Kolsto et al. 2009) (our algorithm thus assigned the two genomes arbitrarily among the two lineages, well within 98% sequence identity). Nevertheless, this reflects on the known, large phenotypic and genomic diversification within the so-called Bacillus "group" (Vilas-Boas et al. 2007; Kolsto et al. 2009) (e.g., Bacillus cereus is usually motile and produces a capsule and toxins, whereas B. cereus tends to be motile and to make no capsule). Importantly, our data show that such a high level of phenotypic and genomic plasticity among co-occurring lineages is exceptional, especially when they are so closely related phylogenetically. Perhaps the unusual life cycle of Bacillus (involving a resilient endospore stage) is conducive to unusually large changes in lifestyle and phenotypes, over short time periods. In contrast, the second example describes two lineages that are very distant phylogenetically (one is an Archaeon and the other a Bacterium), and yet they co-occur quite specifically. In our data, these two (Salinibacter and Haloquadratum) are outliers because, despite their distance, they have very similar genome sizes and very similar functional pathway coverage, marking the current record at such a large phylogenetic distance. Perhaps, both lineages have independently entered the same niche (i.e., warm, fully oxygenated brines) (see also Kunin et al. 2008b), and have thus converged toward coarsely similar overall genomic features. The next example again presents two fairly unrelated lineages (related only at the phylum level), which are even more closely matched in terms of genome size and functional genome composition. They occur together very specifically, on acidic human skin, dust, and in filtered air (seven observations by five distinct laboratories; $P = 6 \times 10^{-13}$; note that dust and air samples are related to skin since they may contain small skin-derived particles). Despite apparently occupying the same niche, it is notable that only one of them (Finegoldia) has a tendency for opportunistic pathogenesis (Goto et al. 2008). The last example concerns two lineages that occur together in the open ocean (Prochlorococcus and Alteromonas). They are unrelated phylogenetically, and they were chosen as outliers here because their genomes are unusually distinct in size (Alteromonas is more than twice as large as Prochlorococcus) (Bocap et al. 2003) (note that our analysis has insufficient resolution to specify the exact "ecotype" for either lineage). Since both were sampled in the open water, it is difficult to envisage any mechanism of their association, but this particular pair has been noted before—Alteromonas has been enriched as a co-contaminant in Prochlorococcus cultures, facilitating the growth of the latter (Morris et al. 2008) (perhaps by alleviating oxidative stress). In the ocean, the association is probably rather unspecific, although it is conceivable that Alteromonas, as an opportunistically growing heterotroph, may profit from biomass accumulated by the primary producer Prochlorococcus. These lifestyles are quite distinct and might explain the unusually large differences in genome size.

Overall, however, we find that co-occurring genomes tend to closely match each other's genome sizes and broad functional composition. These results seem to be compatible with a picture of competition (Hibbing et al. 2010), rather than cooperation, among most of the distinct microbial lineages found at any given site: If the majority of lineages were to routinely cooperate by specialization and division of tasks, this would presumably result in genomic features that might become more distinct from each other over time. Of course, the broad view that we take here could easily make us miss cooperation among a subset of lineages, such as syntrophy and other mutual benefits from the juxtaposition of
distinct molecular capabilities. But such interactions are perhaps anyway more fleeting encounters rather than stable mutualisms. Indeed, long-term obligatory mutualism usually requires stable and specific physical contact between the organisms in question (Boucher 1985), a requirement that makes it perhaps less feasible for microbes that are generally dispersed easily (except, of course, when vertically inherited together within a eukaryote) (Vautrin and Vavre 2009).

In the future, statistical approaches like ours stand to benefit greatly from the projected further increases in both microbial genome sequencing (Ahmed 2009; Chain et al. 2009; The NIH HMP Working Group 2009) and 16S rRNA sampling (Tringe and Hugenholtz 2008; Costello et al. 2009). Both types of data will prove particularly valuable when augmented with standardized information about the environments sampled, for example, by following the recommendations of the MIENS standard (http://gensc.org/gc_wiki/index.php/MIENS). Novel and specific microbial assemblages can already be identified using the current data (see Fig. 3; Supplemental material), and more such discoveries can be expected with higher data coverage. Note that our approach does not require prior information about environmental ontologies or hierarchies of sampling sites; instead, groups of biologically related sampling sites are defined by the data themselves (Fig. 2; Supplemental Fig. S1). In general, approaches that integrate sequence data from both strain sequencing and from environmental marker gene sequencing hold great potential, since they connect the molecular information contained in the (pan-)genome of each lineage to the quantitative occurrence pattern of that lineage around the globe.

Methods

Definition of taxonomic units and sampling events

All 298,591 available 16S rRNA sequence records were downloaded from the Greengenes database (Desantis et al. 2006) on January 2009. At Greengenes, these sequences had already been cleaned of potential chimera by the program Bellerophon (Huber et al. 2004). We filtered sequences according to their lengths (≥900 nt for Archaea and ≥1200 nt for Bacteria) and additionally flagged sequences predicted to be chimeric by the program ChimeraSlayer (http://microbiomemutl.sourceforge.net/). We also removed from the analysis all sequences lacking annotations in any of the fields “author,” “title,” or “isolation_source.” This was done in order to be able to define a sampling event for each record. In our study, a “sampling event” is defined as the unique concatenation of these three annotation fields (author + title + isolation_source).

Archaeal and bacterial sequences were aligned separately, using the secondary-structure aware aligner “Infernal” (Nawrocki et al. 2009), together with the corresponding 16S rRNA covariance models of the RDP database (Cole et al. 2009). Before defining OTUs, we removed sequences for which the alignment had not been successful (i.e., Infernal bit-score < 0). OTUs were built for both Archaea and Bacteria by hierarchical clustering (complete linkage), at various distances (from 0.2 to 0.01), using the clustering tool of the RDP pyrosequencing pipeline (Cole et al. 2009; http://pyro.cme.msu.edu/). Because not all 16S sequences reported in databases are necessarily genuine environmental sequences (Tanner et al. 1998; Barton et al. 2006), we assembled a database of potential laboratory contaminants, containing 38 distinct sequences (Tanner et al. 1998; Barton et al. 2006). Homology searches revealed that between 47 and 309 of our OTUs contained such sequences (matching at 97% sequence identity or better). However, these OTUs are rarely involved in significant co-occurrences; for example, in Figure 2 only 17 of the OTUs shown contain potential contaminants, and these are scattered over various smaller clusters (they are flagged in Fig. 2 and in the detailed Supplemental material).

In order to compute sequence divergence values for pairs of OTUs, we first selected a single sequence to represent each OTU. (We chose the sequence that had the minimum sum of squares of distances to all other sequences within that cluster; note that this does not favor short sequences since the distances we used are length-normalized.) We then aligned these representative sequences pairwise (using the program “water” from the EMBOSS package) (Rice et al. 2000) and determined their sequence identity.

Classification of taxonomic units

In order to assign taxonomic classifications to entire OTUs, we first assessed the pre-annotated taxonomies of all individual 16S rRNA sequences in Greengenes (sensu RDP taxonomy). Where these were still annotated as “unclassified,” we re-ran the taxonomy classification using the RDP classifier (Cole et al. 2009). Taxonomy predictions reported there were considered reliable, if supported by a minimum bootstrap value of 80%. To assign taxonomy classifications to OTUs, we then used a simple majority vote: If more than half of the sequences present within a cluster agreed upon a classification, the OTU was annotated as belonging to this taxon. In case of conflicts, we assigned consensus classifications at increasingly higher levels of taxonomy until the majority vote condition was again met.

Co-occurrence analysis

In order to reduce the search space for co-occurrence testing (which encompasses potentially more than 2 billion pairs, for example, in the case of OTUs defined at 99% sequence identity), we limited our analysis to OTUs occurring in at least three distinct sampling sites. Conversely, we only considered sampling events encompassing at least two distinct OTUs. For these “filtered” OTUs and samplings (see also Table 1), we tested the co-occurrence significance for all possible pairs using the Fisher’s exact test. For each test, the four cells in the contingency table denoted the number of samples containing both OTUs, one of the two OTUs only, or none of the two, respectively. Subsequently, we adjusted all P-values for multiple testing using the Benjamini and Hochberg FDR controlling procedure (Benjamini and Hochberg 1995), as implemented in the “multtest” library of the statistical software package R (http://www.r-project.org). We also verified our FDR empirically, by re-computing the associations using randomized input data. For this, we randomly reassigned the various OTUs to the various sampling events, under the constraint that each OTU kept the overall number of samples it mapped to, and each sample kept the overall number of OTUs. This maintained the size distributions of both, samples and OTUs (results are provided in Table 1). To compute the necessary large number of tests in a reasonable time, we used a C-implementation of the test in the Apophenia library for scientific computing (http://apophenia.sourceforge.net), using the python SWIG interface as a wrapper. For selected examples of co-occurring lineages discussed in the text (Figs. 1, 3; Supplemental Fig. S1), we also computed the odds-ratio (“or,” a statistical measure of effect size), in order to assess the strength of the reported associations. Note that our input data, and thus also our predicted associations, likely suffer from under-sampling and probably also from systematic biases in the sampling. Both effects are difficult to quantify, but are likely present due to variable choices of PCR primers (information about primers
is often not available in the sequence records), and also due to experimental biases in DNA extraction protocols. However, while such biases can likely depress the detection of certain lineages, it is less likely that they generate false-positive associations at the level of specificity that we observe here (see Supplemental Fig. S3, and see also the randomizations described above). We also noted that, overall, larger samples contribute more co-occurrence associations than smaller samples, as expected. We quantified this in two ways: by stratifying the input data by sample size, and by randomly down-sampling the larger environmental samples (these often focus on the mammalian gut). The results of both tests are summarized in Supplemental Figure S7; reassuringly, we observe that entirely removing gut-related sequences through keyword searches, while lowering the number of association clusters, still supports the quantitative conclusion that we report in Figures 4 and 5 (see Supplemental Fig. S10).

Network inference
Based on the co-occurrence analysis results, we constructed networks of coexisting microbes for different levels of OTU definitions. For this, the FDR cutoff for each individual edge in the network was 0.001. In order to obtain a simplified view on the results and to identify cohesive modules of coexisting microbial lineages, we clustered our networks using the Markov cluster algorithm (MCL algorithm; http://micans.org/mcl) (Enright et al. 2002). This clustering was performed using as the similarity metric (i.e., edge weights) the normalized co-occurrence similarity between OTUs, defined here as the Jaccard similarity coefficient (i.e., \( \text{cooc_count}/(\text{otu1_count} + \text{otu2_count} - \text{cooc_count}) \)). We set MCL’s “inflation” parameter to 2.0 when running the algorithm. All network images were generated using custom Python scripting and the Python module “NetworkX” (http://networkx.lanl.gov), which provides an interface to the “Graphviz” graph visualization software (http://www.graphviz.org).

Cluster annotation
To annotate clusters in the co-occurrence network with environmental information, we relied on the controlled vocabulary maintained by the Environment Ontology project (EnvO, version 1.51; http://environmentontology.org). In a first step, we assigned EnvO keywords to each OTU in the network; to do so, we scanned all words in the “isolation_source” field from each OTU and assigned ontology terms to that OTU based on exact matches. For many of its terms, EnvO also provides “synonyms”; for cases in which a term could not be matched directly, we also allowed matches via these synonyms, but only for synonyms of the categories “EXACT” or “NARROW” (omitting the categories “RELATED” and “BROAD”). The Fisher’s exact test then allowed us to assign significantly over-represented keywords (FDR = 0.01; \( P \)-value adjusted for multiple testing using the Benjamini and Hochberg procedure) for each given cluster or subnetwork, compared to the background frequency of these terms in the entire network.

Comparative genomics
First, we mapped available, completely sequenced genomes to our OTUs, for various levels of OTU definitions. For this, we extracted the 16S rRNA genes predicted for 881 complete genomes contained in the ReSeq database (ReSeq 35, 05-13-2009), requiring a minimum length of 700 bp. We then compared these sequences against representative 16S sequences from each OTU, using BLAST with the following parameters: “-a 2 -m 8 -p blastn -v 1000 -b 1000 -r 2 -q -3 -G 5 -E 2 -e 0.01.” For genomes that are annotated with more than one predicted 16S RNA gene, we retained the longest copy. For the mapping, we then ranked all sequence matches by bit-score (best score first) and, parsing through the list, assigned each genome to the best-matching OTU (skipping those that were already previously assigned to another genome). In addition, we required that the alignment length for the BLAST hit was at least 800 bp and that the sequence identity of the match was 97% or greater.

We then analyzed co-occurring OTUs by comparing their mapped genomes, using several characteristics: genome size, GC content, and relative coverage of KEGG (Kyoto Encyclopedia of Genes and Genomes) pathways (Kanehisa et al. 2008). To compute genome size ratios, we used the total DNA length of the non-redundant chromosomes and plasmids, expressed in nucleotides; to compute GC content ratios, we used the predetermined values for the complete genomes available at http://www.ncbi.nlm.nih.gov/ genomes/iProks.cgi. In order to compare genomes in terms of their encoded functions, we assessed the relative coverage of pathways as annotated at KEGG, using the KEGG API (http://www.genome.jp/kegg/soap). We computed normalized vectors describing the relative pathway coverage among all annotated genes of a given genome and then compared these vectors by computing their Euclidean distance. In order to exclude potential artifacts arising from occasional annotation errors in KEGG, we repeated this analysis with two additional, independent systems of functional genome annotation, retrieving essentially the same results (Supplemental Fig. S4).

To test for the statistical independence of our observations made for a given distance measure, against another measure (usually against phylogenetic distance) (Fig. 4E–G), we first learned the dependency between the two measures based on randomly selected pairs of genomes (blue dots). This dependency was then described using a running median (blue lines in Fig. 4). Next, we assessed the data of interest (i.e., pairs of co-occurring genomes) by computing for each data point its vertical distance to the (blue) running median, divided by that median itself. This measure has been termed “relative distance to median” (“dm”; see, for example, Newman et al. 2006); it permits us to compare data at a given, fixed setting of a second, potentially confounding variable. From this, we generated a distribution of normalized distance values, which we compared to the corresponding random background distributions, using the non-parametric Kolmogorov-Smirnov test.

Data availability
Raw input data, as well as all computed results of this study (including sequence data, operational taxonomic units, co-occurrence statistics, network clustering, and genome mapping) are available online at http://mblnx-kallisto.uzh.ch:8888/microbial_coexistence/. In addition, a zoomable and clickable version of the network in Figure 2C is available as Supplemental Figure S12, which can be downloaded from the Supplemental materials.

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References


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