
Microbial communities drive global biogeochemical cycling (7). Bacteria and archaea, for example, strongly influence marine carbon, nitrogen, and sulfur fluxes, thereby modulating global ocean productivity and climate (2, 3). Elucidating the processes that shape microbial communities over space and time is important for predicting how biogeochemical cycles will change with changing environmental conditions.

Taxonomic microbial community profiling can reveal intriguing, but often unexplained, variation

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Differences in metabolic function among organisms are thought to underlie much of this variation as a result of selection for specific metabolic pathways based on physicochemical conditions ("metabolic niche effects" or "environmental filtering"). However, other factors such as biotic interactions (5–7), limits to spatial dispersal (4), and neutral demographic drift (8) could also affect community composition. Moreover, distantly related microbes can often perform similar metabolic functions, further complicating a mechanistic interpretation of taxonomic community variation (9). Direct estimates of functional potential in terms of community gene content using environmental shotgun sequencing—or metagenomics—have revealed correlations between the distribution of particular metabolic pathways and environmental conditions (2, 10). However, it is not known how metabolic niche effects interact with other community assembly mechanisms, nor how variation in taxonomic composition relates to ecosystem function. Ideally, one would hope to split community variation into distinct components ("axes of variation"), each of which relates to different ecological processes.

Here, we combine taxonomic and functional profiling of more than 100 bacterial and archaeal communities across the global ocean to elucidate the role of environmental filtering, global functional redundancy, and dispersal limitation in shaping marine microbial communities. We generated taxonomic profiles based on shotgun DNA sequences of the 16S ribosomal gene, a standard marker gene in microbial ecology (11). Functional profiles were generated by associating individual organisms with metabolic functions of particular ecological relevance, such as photoautotrophy and nitrate respiration, using an annotation database.
that we created on the basis of experimental literature. We then explored taxonomic composition, functional potential, and taxonomic composition within individual functional groups in relation to environmental conditions and geographical location.

We performed regression modeling of the relative abundances of functional groups, as well as the proportions of various operational taxonomic units (OTUs) within each functional group, using 15 key abiotic environmental variables such as dissolved oxygen, salinity, temperature, and depth (table S1). For each OTU or functional group, we used a stepwise model selection algorithm to choose the appropriate subset of environmental variables that maximized predictive power while avoiding unrelated variables. In general, environmental conditions strongly predicted the functional profiles of microbial communities but weakly predicted the taxonomic composition within each functional group, as measured by the cross-validated coefficient of determination of the models ($R^2_{\text{cv}}$). In particular, the $R^2_{\text{cv}}$ for the relative abundances of almost all functional groups [mean $R^2_{\text{cv}} = 0.48 \pm 0.27$ (SD) over all functions] exceeded the average $R^2_{\text{cv}}$ achieved for the OTU proportions within those groups [mean $R^2_{\text{cv}} = 0.15 \pm 0.09$ over all functions; Fig. 1C]. To further explore the relative importance of individual environmental variables, we performed correlation analyses between each functional group or OTU and each environmental variable. Consistent with our regression modeling, most environmental variables either correlated more strongly with relative functional group abundances than with OTU proportions within functional groups, or did not correlate strongly with either of them (Fig. 1, D and E). These patterns persisted at higher taxonomic levels (e.g., genus, family, or order; figs. S1 and S2).
Hence, our results are not caused by suboptimal choice of taxonomic resolution, but reflect a general lack of environmental effects on the composition within functional groups.

The interaction between environmental conditions and function is only partly reflected in overall taxonomic community structure. Regression models of relative taxon abundances against environmental variables at the whole-community level and at any taxonomic resolution had a lower $R^2$ (Fig. 1A and fig. S3) than achieved for most functional groups (blue bars in Fig. 1C), but comparable to or higher than the mean $R^2$ achieved for the taxon proportions within functional groups (gray bars in Fig. 1C and fig. S2). Similarly, environmental variables were generally less correlated to relative taxon abundances at the community level than to relative functional group abundances (Fig. 1B versus Fig. 1D). Further, the distinction between water column zones based on function is comparable in strength to a distinction based on taxonomy at the whole-community level (Fig. 2, B and C, and fig. S4). Indeed, permutational multivariate analysis of variance (PERMANOVA) tests (11) of dissimilarities in community composition yielded similar pseudo-F statistics (23.8 and 24.4, respectively; $P < 0.001$ in both cases). Similarly, the segregation of functional groups between water column zones (e.g., mesopelagic versus surface) was comparable to or higher than the segregation of taxa (as determined by permutation tests; Fig. 2, E to G). Together, these results support the interpretation that overall community structure results from the superposition of functional structure, which is strongly related to environmental conditions, and composition within functional groups, which is only weakly related to environmental conditions. In particular, an organism's metabolic potential appears to be the main trait selected for by environmental conditions across the global ocean. In support of this interpretation, we found that OTUs sharing a higher number of functions tend to correlate with each other more positively (Fig. 2H), consistent with a strong role for metabolic niche effects (11).

Our results also suggest that energetic and stoichiometric constraints, not the identity of the microbes, drive ocean microbial metabolic function. Indeed, we found that the proportions of individual OTUs within a functional group were generally only weakly correlated to the group's overall abundance (figs. S5 and S6). Similar smaller-scale observations have been reported previously. In a wastewater treatment plant, the ratio of aerobic ammonia-oxidizing bacteria and heterotrophic bacteria remained constant over time, while highly variable taxonomic composition within each functional group was only weakly explained by environmental factors (8). In another study, metatranscriptomics in two distinct ocean regions revealed strongly conserved diurnal succession patterns of community gene expression, despite largely nonoverlapping taxonomic affiliations of transcripts between the two regions (12).

If environmental conditions interact with functional community structure, what drives the taxonomic variation within individual functional groups? The high taxonomic richness and variability within functional groups indicate a high global functional redundancy (Fig. 3 and fig. S6), which may have emerged partly as a result of horizontal gene transfer (1). The variation within functional groups that is not explained by our regression models could be due to limited granularity of our environmental characterizations and functional assignments. For example, unknown physicochemical variables and phenotypic differences within functional groups could drive location-dependent growth differences between competing clades. This may explain why, in some cases, functional groups were dominated by a few OTUs in multiple samples despite a high OTU richness (fig. S6). However, the variables considered here are known to be important predictors for marine microbial community composition (13, 14), and it is unlikely that unconsidered environmental variables alone could explain the apparent randomness seen within so many functional groups (fig. S6).
Fig. 4. Community dissimilarities versus geographical distances in the mesopelagic zone. (A and B) Bray-Curtis dissimilarities between microbial communities, compared with geographical distances (one point per sample pair). Community dissimilarities are calculated in terms of relative functional group abundances (A) and relative OTU abundances (B). Spearman rank correlations (ρ) and statistical significances (P) are shown. Only sample pairs in the mesopelagic zone and within the same ocean region are considered. Least-squares regression lines are shown for reference. (C and D) Similar to (A) and (B), but showing residual Bray-Curtis dissimilarities after accounting for environmental effects. For other taxonomic resolutions, see fig. S8. For dissimilarities within functional groups, see figs. S9 and S11. For surface water samples, see fig. S7.

when compared to the much better predictability of functional community structure (Fig. 1).

Alternatively, spatially limited dispersal may cause distribution patterns not attributable to environmental conditions, although its importance for microorganisms is generally thought to be low and to depend strongly on the type of environment [e.g., lakes versus open ocean (4)]. Marine microorganisms, in particular, can be dispersed at global scales by large ocean currents and were previously found to be recruited from a global marine seed bank (15). To test the effects of dispersal limitation, we compared geographical sample distances with dissimilarities in functional and taxonomic composition at the community level as well as within individual functional groups. To avoid comparisons spanning continental land masses, we only included sample pairs within the same ocean region. When environmental effects were not taken into account, Mantel correlation tests revealed significant positive correlations between geographical distances and community dissimilarities (P < 0.05; Figs. 4, A and B, and Figs. S7, A and B, and fig. S8, A to C), consistent with previous findings at similar scales (14). We also observed weaker distance decay in similarity within functional groups (figs. S9 and S10). When we accounted for environmental effects, residual dissimilarities showed no significant correlations with geographical distance, neither at the community level (Fig. 4, C and D, fig. S7, C and D, and fig. S8, D to F) nor within functional groups (figs. S11 and S12). This suggests that the distance decay in similarity is driven by environmental variation affecting community function (and, to a lesser extent, composition within functional groups) and that dispersal limitation in the open ocean is of limited importance relative to other community assembly mechanisms.

Unexplained variation may be driven by community-level processes such as metabolic interdependencies, chemical signaling and defense, or predation by viruses and eukaryotes (5–7, 16). For example, predation by bacteriophages specialized on specific hosts can cause variation in host community composition that cannot be attributed to environmental variation (16). Functional community structure and taxonomic composition within functional groups thus appear to constitute roughly complementary axes of variation, with the former being affected more strongly by environmental conditions and the latter shaped greatly by community-level processes (5). Other mechanisms such as priority effects (17) or demographic drift (8) may also explain some of the variation within functional groups.

Marker gene sequencing can yield detailed microbial taxonomic community profiles and reveal intriguing variation—for example, down the ocean water column (14). The high dimensionality of these profiles, however, poses a challenge to their mechanistic interpretation. Generic statistical techniques such as principal components analysis (18) are often used to identify important axes of variation, but these axes typically lack an ecological interpretation. Alternatively, detected species may be combined at higher taxonomic levels that resemble the depth at which traits vary across lineages; however, the optimal taxonomic level is highly trait-dependent (9). Further, many metabolic phenotypes are dispersed irregularly across microbial clades (19). Similar functions may be performed in distant clades (Fig. 3, A to C, and fig. S13) and, conversely, members of the same clade can fill separate metabolic niches (9, 19).

For example, none of the 28 metabolic functions considered here appear to be monophyletic (Fig. 3C and fig. S13). Irregular trait distributions are caused by diverse evolutionary processes, including adaptive loss of function (20) and metabolic convergence accelerated by frequent horizontal gene transfer (1). As a result, taxonomic profiles, at any level, often obscure the relationship between community structure and biogeochemical processes.

Here we have shown that binning organisms into functional groups enables the integration of functional, taxonomic, and environmental information to elucidate community assembly processes. Comparison of functional profiles with environmental variables (Fig. 1D) yields insight into the processes driving variation in community composition along geochemical gradients and, reciprocally, gives information about the effects of that variation on ecosystem processes, although specific causal mechanisms need to be experimentally validated. For example, our functional profiles reveal that groups capable of fermentation or nitrate respiration are overrepresented in deeper zones, where oxygen is often limited and nitrate serves as an alternative electron acceptor for respiration (Fig. 2A). The prevalence of alternative metabolic modes at depth leads to increased richness of both functional groups and OTUs, especially in the mesopelagic zone (Fig. 2D). Although an increase of taxonomic richness with depth has been noted before (14), our analysis shows that this richness gradient is related to the number of available metabolic niches.

The zonation of functional groups with depth (Fig. 2A) is reflected in metagenomic profiles of the same samples (fig. S14) (14) and in previous metagenomic studies (2). However, metagenomic profiles do not directly translate into functional potential, because the same or similar genes can be involved in different—in some cases reversed—metabolic pathways, depending on the microorganism (27). For example, variants of the dissimilatory sulfite reductase genes (dsrAB), abundant in the mesopelagic zone (fig. S14), could be involved in either respiratory sulfur reduction or chemolithothrophic sulfur oxidation (22). Such ambiguities are inherent to metagenomics (27) and apply equally to methods that estimate community gene content by projecting marker genes to sequenced genomes (fig. S14) (23). On the other hand, our phenotype profiles suggest that sulfide and sulfate oxidizers are generally more abundant than sulfate respirers (Fig. 2A), indicating that detected dsrAB genes are mainly involved in sulfur
oxygenation. Translation of taxonomic information into phenotype profiles based on experimental evidence can thus facilitate the ecological interpretation of metagenomes. The full potential of taxonomic profiling remains underused because of our current inability to associate many known taxa with any function. For example, a large fraction of the ubiquitous but poorly studied phyllum Thaumarchaeota is potentially involved in ammonia oxidation (24) but had to be excluded from our functional annotations. Similarly, microeukaryotes likely contribute to several metabolic functions, such as photosynthesis or cellulolysis. Future functional profiling should thus include eukaryotic microorganisms and incorporate putative metabolic potential for uncultured clades revealed by single-cell genomics (25).

The bulk of global biogeochemical fluxes is driven by a core set of metabolic pathways that evolved in response to past geochemical conditions (7). Through time, these pathways have spread across microbial clades that compete within metabolic niches, resulting in an enormous microbial diversity characterized by high functional redundancy. As shown here, splitting variation of microbial community composition into variation of functional structure and taxonomic variation within functional groups reveals an intriguing pattern: The functional component in itself captures most of the variation predicted by environmental conditions, whereas the residual component (i.e., variation within functional groups) only weakly relates to environmental conditions. This has implications for the interpretation of differences in community structure across environments and time. Differences in taxonomic composition that do not affect functional composition may have little relevance to ecosystem biochemistry; conversely, physicochemically similar environments could host taxonomically distinct communities (26). Functional (rather than purely taxonomic) descriptions of microbial communities should therefore constitute the baseline for microbial biogeography, particularly across transects where geochemical gradients shape microbial niche distribution (27). The residual variation within functional groups can then be analyzed separately to elucidate additional community assembly mechanisms such as biotic interactions, dispersal limitation, or demographic drift. An incorporation of global microbial functional profiles and their response to potentially changing environmental conditions, into future biogeochemical models will greatly benefit re-constructive and predictive modeling of Earth’s elemental cycles.

REFERENCES AND NOTES

11. See supplementary materials on Science Online.

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SUPPLEMENTARY MATERIALS

www.sciencemag.org/content/353/6305/1272/suppl/DC1
Materials and Methods
Tables S1 to S4
Additional data table S1
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Sensory Biology

Bats perceptually weight prey cues across sensory systems when hunting in noise

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Anthropogenic noise can interfere with environmental information processing and thereby reduce survival and reproduction. Receivers of signals and cues in particular depend on perceptual strategies to adjust to noisy conditions. We found that predators that hunt using prey sounds can reduce the negative impact of noise by making use of prey cues conveyed through additional sensory systems. In the presence of masking noise, but not in its absence, frog-eating bats preferred and were faster in attacking a robotic frog emitting multiple sensory cues. The behavioral changes induced by masking noise was accompanied by an increase in active localization through echolocation. Our findings help to reveal how animals can adapt to anthropogenic noise and have implications for the role of sensory ecology in driving species interactions.

A nthropogenic noise is a globally rising environmental pollutant that has been linked to lower survival and reduced reproductive success of many animal taxa (7–9). Noise can mask environmental cues, making it difficult to hear moving prey or approaching predators, and can interfere with the perception of acoustic communication signals (4–6). Signal producers may be able to reduce the masking impact—for example, by calling louder (7–9)—but such signaling strategy is unavailable to receivers. Some receivers can depend on perceptual strategies to maintain cue detection and thereby adapt to noisy environments (10, 11).

Predators such as bats and owls are highly specialized to hunt prey by ear (12); thus, noise that masks prey sounds severely hampers their foraging success (4, 5). However, predators may be able to adapt to masking levels of anthropogenic noise by actively shifting their attention or emphasis placed on processing cues from different sensory modalities from the same prey (13–16). We refer to this as cross-modal perceptual weighting (17).

We studied the effect of masking noise on the attack behavior of the fringe-lipped bat (Trachops cirrhosus), a neotropical species that is specialized
Decoupling function and taxonomy in the global ocean microbiome
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