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Statistically learning the functional landscape of microbial communities

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Microbial consortia exhibit complex functional properties in contexts ranging from soils to bioreactors to human hosts. Understanding how community composition determines function is a major goal of microbial ecology. Here we address this challenge using the concept of community-function landscapes-analogues to fitness landscapes-that capture how changes in community composition alter collective function. Using datasets that represent a broad set of community functions, from production/degradation of specific compounds to biomass generation, we show that statistically inferred landscapes quantitatively predict community functions from knowledge of species presence or absence. Crucially, community-function landscapes allow prediction without explicit knowledge of abundance dynamics or interactions between species and can be accurately trained using measurements from a small subset of all possible community compositions. The success of our approach arises from the fact that empirical community-function landscapes appear to be not rugged, meaning that they largely lack high-order epistatic contributions that would be difficult to fit with limited data. Finally, we show that this observation holds across a wide class of ecological models, suggesting community-function landscapes can be efficiently inferred across a broad range of ecological regimes. Our results open the door to the rational design of consortia without detailed knowledge of abundance dynamics or interactions.

Biology is a science of connecting scales of organization. From proteins to ecosystems, we are faced with the question of how variation at a lower scale of organization gives rise to changes at a higher scale. For example, understanding protein evolution requires learning how variation in the primary amino acid sequence determines fold and function. Similarly, at the level of the organism, genetic variation drives changes in phenotype and fitness. In both cases, interactions between constituent parts give rise to functional system properties. One of the most powerful conceptual frameworks for thinking about how these functional properties emerge from components and their interactions is the notion of a landscape¹, where the height of the landscape encodes a scalar-valued function or fitness and position on the landscape corresponds to a particular configuration of components. Landscape thinking permits us to articulate key properties of the mapping from genotypes to fitness, including the relative fitness of related genotypes², the extent and

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nature of interactions between genes^{3,4} and the dynamics of evolutionary trajectories⁵⁻⁷.

Communities of microbes also exhibit functional properties, from degrading complex substrates^{8,9} to resisting invasions¹⁰⁻¹², that arise from the constituent parts and their interactions (Fig. 1a). It is natural to ask whether the landscape concept can also be useful in these scenarios. Here we take inspiration from methods for understanding landscapes for proteins and organisms to characterize the functional landscapes of microbial communities¹³.

During the past century, it has become routine to infer landscapes in the protein and organismal context statistically¹⁴⁻¹⁷. Given that directly assaying all possible mutation combinations is typically infeasible, the statistical approach aims to approximate the landscape from a smaller number of measurements via regression¹⁸, with the interactions between mutations quantified via nonlinear epistatic terms. It is important to note that this approach explicitly neglects the details of the complex and often dynamical underlying processes that cause the fitness change. For example, a statistical approach does not explicitly account for the complex physical interactions between residues that alter the function of an enzyme. Similarly, at the organismal level, this approach neglects the details of how mutations impact gene expression or life history traits. Despite this dramatic simplification, regression-based statistical approaches have been highly successful in both these contexts^{2,19,20}.

Inspired by these successes, here we take a landscape approach to quantitatively predict functions of interest in microbial communities. From this perspective, the presence and absence of species are analogous to mutations in a protein or genome, and a regression can be formulated to predict community function from species presence and absence alone. This is in contrast to most existing approaches to predicting community function, which almost exclusively seek to understand how species presence impacts abundance dynamics and, consequently, function²¹⁻²⁴. Here we consider the possibility that community function can be understood without the intermediate step of predicting dynamics (Fig. 1b). We note that this approach explicitly ignores priority effects²⁵, multistability²⁶ or any other scenario when presence/ absence information does not uniquely specify the community state. Nevertheless, as we will show, community-function landscapes prove remarkably predictive across a range of ecological contexts. This does not mean that priority effects are absent, merely that, for the examples considered here, their impact on community function is, on average, weak enough that the predictive power remains high.

Implementing this approach requires measurements of community function for sets of synthetic communities constructed from libraries of taxa. Here we utilize six existing datasets of this type, representing a diverse set of functional properties^{21,22,27-29}. For all the datasets we study, we find that the functional landscape is well described by models including only additive and pairwise epistatic terms. Moreover, we find that the ruggedness of these landscapes is surprisingly low, such that the effects of species presence/absence on function are, in fact, dominated by additive terms. We support these observations computationally, showing that a regression approach succeeds in learning the community-function landscape across a large class of ecological models, despite using only additive and pairwise terms and only species presence/absence as input.

Taken together, our results show that, at least in the six examples presented here, learning the properties of communities can be accomplished without a detailed understanding of the interactions between taxa or their abundance dynamics. Our findings enable a powerful conceptual framework for predicting community functional properties, from invasion resistance to biotechnological applications.

Results

Learning community-function landscapes via regression

We first formulated a statistical approach to fitting community-function landscapes using datasets that comprise measured values of microbial

community functions for a set of defined species combinations. In each of these experiments, a defined pool of species was used to construct communities combinatorially. Each community was then incubated, typically for a defined period of time, and then a functional property of interest was assayed. For a pool of *N* total species, there are $2^N - 1$ possible species combinations, and measuring all possible combinations is frequently intractable. The first goal of our investigation was to ask whether we can predict the function of all $2^N - 1$ communities by fitting a statistical model to a small subset ($\ll 2^N - 1$) of all possible consortia. The resulting model would provide a global picture of the community-function landscape.

We formulated this problem as a linear regression of the following form:

$$y = \beta_0 + \sum_i \beta_i x_i + \sum_{i < j} \beta_{ij} x_i x_j + \dots, \qquad (1)$$

where y is the scalar-valued function and x_i represents the presence or absence of species *i* in that community. The coefficient β_i is the additive effect of including species *i* in the community, and β_{ij} is analogous to the effect of pairwise epistasis in genetic fitness landscapes, which measures the impact beyond individual additive effects of adding both species *i* and *j*. The ellipses denote higher-order epistasis terms, for example, three-way epistatic terms captured by third-order polynomials and so on.

We took the convention that $x_i = 1$ if species *i* is present in a community and $x_i = -1$ if that species is absent. We denoted absence using -1 instead of 0 to simplify the interpretation of the regression coefficients (discussion in ref. 30). In brief, using $x_i = \pm 1$ allows us to interpret β_i as the average effect of adding species *i* to community function, where the average is taken over all community compositions. Similarly, the pairwise coefficient β_{ij} captures the average epistatic effect of species *i* and *j* together across many consortia. Moreover, this convention (corresponding to the Fourier expansion of the landscape) has some convenient mathematical properties that make it easier to quantify how much variation in the measured function is captured by additive, pairwise and higher-order terms^{30,31}.

We considered regressions truncated at first, second and third orders. Many of the datasets we utilized sampled a number of community configurations that are comparable to the total number of coefficients to be inferred. To mitigate the risk of overfitting, we employed L_1 -regularized regression (LASSO)³² using a cross-validation procedure to estimate the regularization hyperparameter (Methods). To assess out-of-sample generalization error, we applied an additional leave-one-out cross-validation scheme in which each data point was iteratively left out of sample and the model was fit to all remaining data points, allowing an out-of-sample prediction for each distinct experimental community (Methods).

Community function is predictable from species composition

We compiled six datasets in which synthetic bacterial communities were assembled from a pool of species. These datasets represent a broad spectrum of community functions: Clark et al.²² measured the production of the short-chain fatty acid butyrate (Fig. 2a); Langenheder et al.²⁷ measured a combination of biomass and redox activity on the monosaccharide xylose (comment in Methods); Sanchez-Gorostiaga et al.²¹ measured the breakdown of the polysaccharide starch (Fig. 2b,c); Diaz-Colunga et al.²⁸ measured the total production of iron-scavenging siderophores (Fig. 2d). In addition, we considered biomass-related community functions: work by the Sanchez lab (Methods) measured total community biomass (Fig. 2e), and Kehe et al.²⁹ measured the abundance of a single target species (Fig. 2f). Details about the size and species pools for each dataset are given in Supplementary Table 1.

In each dataset, community function can be defined as a measurable scalar quantity, for example, the concentration of a compound



or a measurement of biomass at a given point in time. Therefore we fit models of the form shown in equation (1). We investigated truncating the model at successively higher orders of epistatic terms to determine what degree of model complexity is needed to accurately predict community function. The quality of the fits are shown in Fig. 2 (bar plots) for increasingly complex models, with shaded bars showing in-sample R^2 and white bars showing out-of-sample R^2 (Methods); scatter plots visualize the quality of second-order out-of-sample model predictions.

Remarkably, across all six datasets studied here, models of first or second order provide high-quality predictions ($R^2 - 0.8$ for second-order models). In most cases, additive models alone (for example, $y = \beta_0 + \sum_i \beta_i x_i$) already have strong predictive power ($R^2 > 0.5$), although the addition of second-order terms yielded an increased quality of fit for all datasets. We note that residuals of observed versus predicted values demonstrate similar patterns of heteroskedasticity across datasets, where communities with higher values of the function relative to the mean tend to be underestimated and those with lower values of the function tend to be overestimated (Fig. 2 and Extended Data Fig. 1). It is possible that these patterns are a consequence of bias induced by regularization¹⁸. We note that errors in our predictions do not correlate with community richness (Extended Data Fig. 2), indicating that our models generalize well to diverse communities.

For the dataset by Clark et al. (Fig. 2a), the authors of the original study predicted butyrate production using a complex model that parameterized interactions and abundance dynamics. The quality of our statistical predictions using species presence/absence are similar to those obtained using a complex dynamical model²², suggesting that detailed dissections of community dynamics are not always necessary to make reasonably good predictions of community function.

Empirical community-function landscapes are not rugged

We demonstrated that statistical models based on species presence/ absence can predict microbial community functions with surprising accuracy. In particular, simple models containing only additive and/or pairwise epistatic terms explained the vast majority of the variation in the data (Fig. 2). Our statistical approach represents a strategy for approximating the empirical community-function landscapes for these datasets. We wanted to gain intuition for why these regressions appeared to be so successful. To do this, we sought to quantify the ruggedness of community-function landscapes.

In an evolutionary context, the ruggedness of a fitness landscape dictates the number of local fitness optima and has important implications for the predictability of evolution⁵. In a community context, very rugged landscapes are expected to be much harder to approximate globally using regression methods such as those used here, simply because ruggedness arises from a substantial number of strong high-order epistatic terms. High-order terms are challenging to learn statistically due to the explosive combinatorial increase in the number of terms as model order increases. Thus, we expect that rugged landscapes will be difficult to learn statistically, while non-rugged ones will be straightforward to approximate using low-order models.

We quantified ruggedness using two complementary approaches: first, by explicitly quantifying the relative contribution of terms of different orders to the total variance, as explained below; and second, by using an established metric of ruggedness denoted r/s ('roughness over slope'). To pursue the first approach, we started with the combinatorially complete dataset from Langenheder et al.²⁷ in which growth on xylose was measured for all $2^6 - 1 = 63$ species combinations that can be formed from a six-species pool. This combinatorially complete dataset allowed us to compute the coefficients of the exact full-order empirical landscape, which is a model of the form equation (1) that includes epistatic terms of all possible orders (that is, up to sixth order). Using this exactly inferred landscape, we generated a Fourier amplitude spectrum (Methods), which is a decomposition that reflects the total variance of the landscape that is captured by terms of each order^{30,31}. This spectrum, shown in Fig. 3a (red line), indicates that ~78% and ~16% of the variance in the landscape is explained by first- and second-order terms, respectively, leaving ~6% of variance remaining for higher-order



Fig. 2 | Community function is predictable from species presence/absence in empirical datasets. For each dataset, regularized linear regressions were performed using models truncated at the first, second and third order (equation (1)). Bar plots show the quality of fit (R^2) for each of these models, either using all experimental data or using a systematic leave-one-out cross-validation approach

(labelled 'LOO-CV'). Scatter plots show out-of-sample prediction values for the second-order regression fits obtained via the leave-one-out cross-validation procedure. **a**–**f**, Analyses are shown for datasets by Clark et al.²² (**a**), Langenheder et al.²⁷ (**b**), Sanchez-Gorostiaga et al.²¹ (**c**), Diaz-Colunga et al.²⁸ (**d**), data from the Sanchez lab (**e**) and Kehe et al.²⁹ (**f**). Extended Data Figs. 1–4 provide more details.

terms. In other words, this exact community-function landscape displays a low degree of ruggedness, as it is dominated by low-order terms and is largely free of consequential higher-order terms.

To assess whether this result reflects a meaningful property of the empirical community-function landscape, we performed a randomization test by computationally shuffling the assignments between function measurements and community compositions. For 100 such randomizations, we inferred the new landscape and computed the Fourier amplitude spectra. The results are plotted in Fig. 3a (black line). For the randomized landscapes, we found that terms of third order were most important and additive terms alone captured only 10% of the variance; the peak at third order arises from the fact that there are combinatorially more terms possible at this order than any other. We concluded that a lack of ruggedness in the true landscape is not spurious but rather a distinctly non-random structural feature of this landscape.

We performed similar analyses for the remaining five datasets to estimate the relative importance of coefficients of different orders. Although these datasets are not combinatorially complete and therefore do not permit the inference of the exact amplitude spectra, we inferred a truncated amplitude spectra via third-order regression. We found that in each case, the dominant coefficients are additive, with pairwise coefficients typically the next most important (Fig. 3b). The large fraction of variance explained by additive and pairwise coefficients across these cases again indicates a low degree of ruggedness.

As a second approach to quantify ruggedness, we computed the roughness/slope ratio (r/s) (refs. 33,34), a commonly used ruggedness metric that quantifies how well a landscape is fit by a purely additive model. Explicitly, roughness r is computed by fitting a model with additive terms only (for example, $y = \beta_0 + \sum_i \beta_i x_i$) and determining a residual to this fit. The roughness (r) is the root-mean-square value of these residuals. Slope s is defined as the mean (absolute) value of the additive model error relative to the typical magnitude of an additive term. Large values of this ratio mean that the approximation afforded by an additive model is poor, indicating a high degree of ruggedness.

To make a well-defined comparison between all datasets, we computed normalized r/s values, defined as the ratio between r/s on the original dataset and r/s for 100 randomized landscapes (Methods). Randomized landscapes served here as a natural high-ruggedness comparison. We found that this ratio was consistently $\ll 1$, indicating



Fig. 3 | **Empirical community-function landscapes are not rugged. a**, Normalized Fourier amplitude spectrum obtained from the combinatorially complete landscape from Langenheder et al.²⁷. Normalized amplitude values correspond to the fraction of total landscape variance that is captured by terms at each order. The empirical amplitude spectrum (red) demonstrates that landscape variance is primarily explained by first-order terms, with higher-order terms explaining decreasing fractions of the variance. This denotes a low degree of ruggedness in the empirical landscape. Black traces show spectra obtained from randomized landscapes (points and error bars indicate the mean and standard deviation, respectively, across 100 randomizations). Unlike the empirical landscapes, the randomized versions are rugged: first-order terms explain a relatively small fraction of total variance. **b**, For each additional dataset that is not combinatorially complete, the fitted coefficients of the regularized

third-order linear regression are used to infer the normalized amplitude spectrum at first through third order. As in panel **a**, first-order terms explain more variance in the landscape than terms at second and third orders, indicating a lack of ruggedness. **c**, For each dataset, normalized *r/s* values (a measure of ruggedness; text) are computed by calculating the ratio of empirical *r/s* values on original landscapes to the *r/s* value of 100 randomized landscapes. The normalized *r/s* values for all datasets are notably smaller than 1, again indicating a low degree of ruggedness in empirical landscapes compared to their randomized counterparts. The boxplots show the median as the centre line, with the boxes corresponding to the upper and lower quartiles, whiskers corresponding to values that lie within 1.5× the interquartile range and individual points indicating outliers.

that true landscapes are much less rugged than comparable random landscapes (Fig. 3c).

We concluded that across a diverse range of microbial community functions, empirical landscapes possess a low degree of ruggedness, corresponding to the dominance of low-order (that is, additive and pairwise epistatic) terms, and a notable absence of higher-order epistatic terms. This enables low-order statistical models to faithfully parameterize empirical landscapes and thereby accurately predict community functions.

Ecological models indicate when landscapes can be learned

We have demonstrated multiple empirical examples where the functional landscape of a microbial community proves to be non-rugged, allowing low-order statistical models to predict the function of interest. To understand the expected generality of this observation beyond the six datasets considered here, we turned to ecological models, generating large ensembles of random communities for which the functional landscapes can be evaluated *in silico*. Our goal was not to fit models to the empirical examples above but to probe the conditions under which a lack of ruggedness is expected to be rare or common and to identify the ecological scenarios under which low-order statistical inference is expected to fail. We first sought to interrogate synthetically generated communityfunction landscapes using the generalized Lotka–Volterra (gLV) model, with total abundance (biomass) serving as the function of interest. The gLV model has many variants that differ in assumptions regarding the structure of randomly generated species interactions, but as recently argued by Barbier et al., a large class of such variants can be reduced to a four-parameter 'reference model⁴³⁵. We adopted this reference model for our analyses (Methods), focusing on a sweep of the two key parameters describing interaction strength (μ) and interaction variability (σ). Further analysis varying all four parameters is presented in Supplementary Fig. 5.

For each combination of μ and σ , we performed 10 trials of generating a random pool of N = 10 species (a pool small enough to evaluate the combinatorially complete set of all possible communities) and computed the exact community-function landscape (mimicking our procedure for the Langenheder et al. data in Fig. 3a). The fraction of variance explained by the first- and second-order terms, equivalent to the predictive power (R^2) of the second-order approximation of the landscape, is shown in Fig. 4a. These values are averaged over the ten trials at each point in the μ - σ plane. In the reference model, positive μ corresponds to interactions that are competitive on average, while larger values of σ correspond to a greater degree of variability in the



Fig. 4 | Ecological models indicate both optimism and caution for inferring community-function landscapes. a, The gLV model was used to generate combinatorially complete synthetic community-function landscapes, taking total abundance as the function of interest. Landscapes were generated using an N = 10 species pool, and two parameters controlling the structure of randomly drawn interactions were varied: an interaction-strength parameter μ and an interaction variability parameter σ , with ten landscape trials for each point in the $\mu - \sigma$ plane. Heat map shows the (trial-averaged) R^2 of the second-order approximation of the exact landscape, that is, the variance explained by first- and second-order terms combined. This value is computed from the exact landscape coefficients inferred as in Fig. 3a (no fitting required). Note that as interaction variability σ becomes large, the model becomes unstable, causing species abundances to diverge. In our simulations, any points in the $\mu - \sigma$ plane that encountered divergences in more than five landscape trials are indicated in grey.

The black curve shows an analytical prediction (computed in the $N \rightarrow \infty$ limit³⁶) for the stability boundary of the gLV model, beyond which species abundances will typically diverge. Supplementary Figs. 5 and 6 detail more. **b**, A consumer resource model (CRM) was used to generate synthetic community-function landscapes with randomly generated cross-feeding networks (schematic). One resource was supplied externally ('source'), while the function of the community was defined to be the concentration of a different resource ('target'). Networks comprising N = 10 species and L = 8 resources were generated so as to vary the length of the shortest path (P) connecting the source resource to the target. Plot shows the variance explained (R^2) by models truncated at first, second and third order, computed using the exact landscape coefficients and shown as a function of P. The R^2 value declines with P, as expected. Solid lines correspond to mean values obtained across trials, with error bands indicating +1 standard deviation.

strength of interactions. Note that as interaction variability σ becomes large, the Lotka–Volterra model becomes unstable, causing species abundances to diverge³⁶. In our simulations, any points in μ – σ plane that encountered divergences in more than five trials are indicated in grey (Fig. 4a). Additional details about these simulations are described in Methods.

We found that across the entire range of parameters for which the dynamics of the model are stable, second-order regression provides excellent fits ($R^2 \ge 0.9$) to gLV community-function landscapes (Fig. 4a; the non-grey region is all yellow). Further analysis shows that R^2 remains similarly high even as we change other model parameters, including interaction asymmetry and the variability of species' carrying capacity (Supplementary Fig. 5). These results indicate that non-rugged functional landscapes can be observed across a wide range of ecological scenarios.

It is important to stress that the lack of ruggedness is not a trivial property of all gLV models. Although the Lotka–Volterra model studied here contains only second-order interactions, these interactions couple species *abundances*, and therefore we do not expect the functional landscape to be well approximated by a low-order regression on species presence/absence alone. For example, rugged gLV landscapes can and do exist (Supplementary Fig. 6). In the ensemble of models described by Barbier et al., such examples are curiously rare (Supplementary Fig. 6), at least for biomass as the property of interest. However, it is likely that for other ensembles (for example, with a more complex correlation structure of interactions), rugged landscapes could be more common. Identifying the ecological mechanisms dictating whether the landscape of a given functional property will be rugged or smooth is an important question for future work.

As an illustration, we present one mechanism that will generically lead to increased ruggedness. Consider a nutrient X_1 that is broken down through a chain of L reactions following a linear pathway $X_1 \xrightarrow{n_1} X_2 \xrightarrow{n_2} \dots \xrightarrow{n_L} X_{L+1}$, where each reaction is performed by a specialized community member n_i . If X_1 is the only nutrient supplied to the consortium, and the function of interest is the concentration of the end product X_{L+1} , then the concentration of X_{L+1} will be non-zero if and only if all species $n_1, ..., n_L$ are present. Mathematically, if presence or absence were denoted as $s_i = 0$ or 1, respectively, such a landscape would be described by a single *L*th-order term: $y = \beta_{1...L} s_1 s_2 ... s_L$. Under the convention used here, where presence or absence is denoted by $x_i = \pm 1$, this corresponds to coefficients at all orders being equally important. (As a simple illustration, for three species $s_1 s_2 s_3 = (x_1 + x_2 + x_3 + x_1 x_2 + x_2 x_3 + x_1 x_3 + x_1 x_2 x_3) / 8$, which is easy to check by a direct substitution $s_i \equiv \frac{x_i+1}{2}$.) In this scenario, regressions exploiting only low-order models will provide poor predictive power and the landscape will be rugged (Fig. 4b).

To build on the intuition from this thought experiment, we used a consumer resource model (CRM) to generate synthetic landscapes with N = 10 species competing for L = 8 nutrients. Only one resource was supplied externally ('source', Fig. 4b, red dot), while the function of the community was defined to be the concentration of one of the other resources ('target', Fig. 4b, blue dot). In each trial, we constructed a random cross-feeding network for a pool of N = 10 species, with each species capable of converting one randomly chosen resource into another. We then computed the complete functional landscape, as above for the gLV, by simulating all possible subsets of this ten-species pool. Simulation details are described in Methods.

On the basis of the intuition from the thought experiment above, we expected the predictive power of low-order approximations to correlate with the length of the shortest path (P) connecting the source resource to the target (schematic, Fig. 4b). The landscapes corresponding to cross-feeding networks with long paths from source to target (that is, large P) are expected to be more rugged. To confirm this, Fig. 4b plots the fraction of variance (R^2) explained by low-order approximations of the exact synthetic CRM landscapes, shown as a function of the shortest chain length *P*. We observe that while successful in cross-feeding networks with small *P*, low-order models were increasingly challenged as *P* increases, with R^2 dropping below 0.5 for second-order models beyond *P* = 4. Increasing model order from additive only to third order substantially improved predictive power, particularly at large *P*, consistent with the idea that increasing *P* increases the prevalence of higher-order terms. These results illustrate that low-order approximations of community-function landscapes may fail to make accurate predictions in ecological scenarios with long chains of trophic dependencies or other situations when function is strongly contingent on the simultaneous presence of multiple species.

Discussion

The key result of our study is the demonstration that functional properties of microbial communities can be predicted by simple statistical models knowing only which species are present or absent. Remarkably, in analogy to fitness landscapes describing proteins and organisms, we showed that regressions can quantitatively describe empirical landscapes for a wide range of community functions. We found that the success of these regressions derives from the fact that the underlying landscapes are not rugged, allowing the majority of variation in function to be captured by additive and pairwise terms in the regression. The predictions did not require specialized knowledge or measurement of microscopic system properties, only a dataset comprising quantitative measurements of a function taken from a collection of defined communities drawn from a fixed pool of species.

Our simulations of generalized Lotka–Volterra and consumer resource models demonstrated that we can expect low-order approximations of landscapes to work well across a range of ecological contexts. We identified a clear exception, however, in functions that are strongly contingent on the simultaneous presence of multiple highly specialized community members. One might therefore expect that the low-order landscape approximations might encounter challenges, for example, in systems with long linear chains of reactions, such as those present in anaerobic digesters³⁷ and Winogradsky columns³⁸. A critical direction to be addressed in future work is a systematic analysis of the ecological mechanisms that either enable or impede the performance of approximations of community-function landscapes. A better understanding of these mechanisms would provide a more principled view of community functional properties amenable to our statistical approach.

Because our approach predicts community function from species presence/absence, it explicitly assumes that replicate communities with the same composition will have the same function. This assumption could fail if communities exhibit alternative stable states^{39,40} with distinct final abundances and functions despite identical presence/ absence compositions. While alternative stable states have been documented in synthetic consortia²⁶, they do not appear to be widespread or have large impacts on function in the communities studied here. Extended Data Fig. 3 shows predictions for replicate communities with identical compositions. We find that most replicates in all six datasets studied are well predicted by our model. We conclude that alternative stable states do not drive substantial functional variation across the consortia studied here. However, none of the datasets studied here systematically varied the initial relative abundances for communities of fixed composition, and this may result in communities not reaching alternative stable states despite their existence. In ecological contexts where alternative stable states are pervasive and drive large functional variation, our predictions would begin to degrade in quality.

Another question is how our results could be extended to contexts with extensive functional redundancy between taxa. Our regression approach would probably struggle in these scenarios. This is easiest to see in the extreme limit of taxa that are perfectly interchangeable: a high-dimensional OR function, where OR is defined as the Boolean logical operator, has Fourier terms of all orders, and any low-order model would be a poor approximation. Thus, the excellent performance of our method on the available datasets is probably aided by the fact that the labour-intensive nature of the combinatorial experiments favours synthetic communities with relatively low redundancy. The experimental cost associated with increasing the number of species discourages the inclusion of taxa highly similar to those already in the pool. This is the opposite regime of the natural communities, which are often phylogenetically under-dispersed^{41,42}.

Whereas the method as presented would probably struggle in communities with extensive redundancy, the argument above indicates how this limitation could be remedied, namely by grouping redundant taxa before performing regressions. In fact, our results suggest that the high predictive power of a simple regression on the variables describing the presence or absence of any member in a group could be taken as the criterion indicating that the grouping was chosen appropriately. This offers a path towards quantitative prediction of complex community functions in the high-diversity regime and echoes recently proposed ideas from multiple groups⁴³⁻⁴⁵.

Our results complement previous studies demonstrating that community-function landscapes follow patterns of global epistasis²⁸. These patterns were first discovered in the context of organismal fitness landscapes⁴⁶⁻⁴⁸. Global epistasis refers not to the impacts of individual epistatic terms but rather to properties (for example, diminishing returns) that emerge from the collective impact of many epistatic contributions. In the context of microbial communities, global epistasis arises as a linear relationship between the impact of adding a specific species on community function and the function of the 'background' community to which the species was added²⁸. It is worth noting that the regression and global epistasis approaches represent distinct strategies for predicting community function. The regression approach presented here attempts to predict the function of an arbitrary community via an ansatz assuming that only low-order epistatic contributions are important, whereas the global epistasis approach attempts to estimate the effect of adding a species to a given background community harnessing the predictability of patterns arising from combined epistatic contributions up to arbitrary order⁴⁷. Connecting the concept of global epistasis with the regression approach to learning functional landscapes remains an important exercise, as the two approaches may provide complementary insights in different ecological scenarios.

Perhaps the greatest downstream impact of our study is the possibility of using statistically inferred landscapes to rationally design communities with desired functional properties. Because community-function landscapes can be approximated by sampling only a subset of all possible species combinations, it is conceivable that even large synthetic consortia with predefined functions can be designed and optimized computationally given only a small number of measurements. Determining the optimal sampling strategy to accurately infer landscapes remains an important avenue for future work. For example, it is unclear whether the sampling should include a mix of simple communities of <3 taxa or whether high-diversity communities are more informative given a restricted number of measurements. However, even without extensive optimization, our models were able to identify the communities with the highest functional output, even when these communities were left out of sample. The simplicity of the approach makes it readily portable across contexts and functions and its performance could offer an appealing advantage relative to alternative design strategies^{22,49}.

Methods

Collection and preprocessing of datasets

Datasets were compiled from six experimental efforts to measure various community functions in defined synthetic microbial consortia. Details about these datasets and references are listed in Supplementary Table 1.

The dataset by Langenheder et al.²⁷ was generated by measuring the activity of synthetic communities with xylose provided as the sole carbon source. Metabolic activity was measured via the absorbance of

a redox dye (tetrazolium violet) at 600 nm over multiple time points. It should be noted that because cell scattering at 600 nm probably also contributed to absorbance, the functional values collected in this study reflect a combination of both redox activity and biomass growth. While this detail complicates the mechanistic interpretation of the data, it does not present a problem for our analysis, as the data still represent a complex functional property for the regression approach to predict. Here the community-function landscape we sought to approximate was constructed from the functional values collected at the endpoint of the experiment (48 h).

The dataset by Sanchez-Gorostiaga et al.²¹ included 53 community configurations out of the possible 63. We note that as we confirmed with the authors of the original study, the potentially ambiguous phrasing in the original manuscript ('every combination of six amylolytic soil bacteria') referred to every *pairwise* combination being included in the list, not that the dataset was combinatorially complete.

Total community biomass data collected by Diaz-Colunga et al. were not included in the original manuscript²⁸. Detailed experimental protocols for isolation, culturing and community assembly are given in refs. 28,50, with the biomass dataset differing from these methodological details in the following ways: (1) the eight isolates used in the biomass dataset were distinct from the siderophore production isolates (though isolated using the same methodology), (2) the medium used in biomass growth experiments lacks a trace mineral supplement, (3) biomass was quantified by measuring the optical density (OD) at 600 nm using 100 µl of endpoint cultures in an AccuSkan FC plate reader (Fisher Scientific) and (4) monoculture measurements for four strains were omitted, resulting in a dataset of 160 unique configurations (instead of 164 as should otherwise result from the methodology in ref. 28). Though isolates used in the biomass dataset were not sequenced or taxonomically identified, colonies either possessed distinct morphologies, possessed distinct colour profiles when grown on chromogenic agar plates (CHROMagar Mastitis GN) and/or were isolated from separate environmental samples and were therefore probably genomically distinct.

Data from Kehe et al.²⁹ were generated using a microwell array approach in which each community was assembled by randomly grouping nanolitre droplets of defined species composition into 2–19 droplet combinations. Due to this stochastic assembly, initial species abundances varied beyond binary presence/absence: for example, a three-droplet combination containing two droplets of species A and one droplet of species B will have different initial abundances than the combination of one droplet of species A and two droplets of species B. Because the formulation of community-function landscapes in equation (1) operates on binary species presence/absence, this variation in initial abundances was ignored and a species was considered to be present in a community if it had positive initial abundance.

In cases where datasets contained experimental replicates, the mean over replicates was taken to obtain a single community-function value for each unique species combination.

Statistical inference of landscapes via regression

Community-function landscapes were approximated for empirical datasets by fitting equations of the form equation (1) truncated at first, second and third order via LASSO regularization³². Tenfold cross validation was used to estimate the regularization hyperparameter. All models were fit using the package glmnet in R version 4.1. 2. For all datasets considered here, the computational demands of fitting our models were modest, and all model fitting and analysis was performed using a personal computer.

To fit the data, two strategies were employed. First, all available data points were used to fit landscape models. The coefficients of determination (R^2) for these fits are shown in grey bars in Fig. 2, and predicted versus observed values are shown in Extended Data Figs. 3 and 4. Second, to obtain an estimate of out-of-sample model accuracy,

Calculation of Fourier amplitude spectrum

Because equation (1) with $x_i \in \{-1, 1\}$ corresponds to the Fourier expansion of a fitness landscape³⁰, the Fourier amplitude equation³¹ at order p can be written simply as

$$A_p = \sum_{i \in p} \beta_i^2.$$
 (2)

It can be shown that sum of amplitudes across all orders is equal to the total variance of the landscape. This permits the calculation of the fraction of total variance explained by terms at order *p* as $A_p / \sum_{p'} A_{p'}$.

Calculation of r/s ruggedness metric

The roughness-slope ratio, *r/s*, is a measure of landscape ruggedness that quantifies how well a landscape is fit by a purely additive model. This quantity was computed by first fitting a linear model of the form:

$$y = \beta_0 + \sum_i \beta_i x_i, \tag{3}$$

where y is the measured community function, and the coefficients β_0 and β_i are obtained ordinary least-squares regression.

The roughness, *r*, is defined as the root-mean-squared-error of the resulting fit, or

$$r = \sqrt{\frac{1}{L}\sum_{i} (y_i - \hat{y}_i)^2},$$
(4)

where *L* is the number of data points in the landscape and \hat{y} is the value of *y* fitted by linear regression. The slope, *s*, is defined as the mean of the absolute value of coefficients β_i

$$y = \frac{1}{n} \sum_{i} |\beta_i|.$$
 (5)

Larger values of *r/s* indicate a greater deviation from linearity, therefore a more rugged landscape. In contrast, an *r/s* value of 0 would correspond to a perfectly additive landscape.

The values of *r/s* depend on landscape size, and sensible comparisons of this quantity between datasets require an appropriate normalization. Because randomized landscapes represent a natural high-ruggedness comparison, *r/s* values computed on randomized landscapes were used as scaling factors, for example

$$r/s_{\text{normalized}} = \frac{r/s_{\text{original}}}{r/s_{\text{randomized}}},$$
 (6)

was computed. Normalized r/s values close to 1 indicate that the empirical landscape is as rugged as the randomized landscape, while values close to zero indicate a low degree of ruggedness in the empirical landscape. One hundred randomizations were performed for each dataset, and values of $r/s_{normalized}$ were computed for each; the distributions of these values are shown in Fig. 3c.

Simulations of the generalized Lotka-Volterra model

Community-function landscapes were synthetically generated using the generalized Lotka–Volterra (gLV) model, with total abundance

serving as the community function of interest. A four-parameter 'reference model' formulated by Barbier et al.³⁵ was used to explore important dimensions of the gLV parameter space. The model can be written as follows:

$$\dot{N}_i = \frac{r_i}{K_i} N_i \left(K_i - N_i - \sum_j \alpha_{ij} N_j \right), \tag{7}$$

where for species *i* in a pool of *N* total species, N_i is the abundance, r_i is the intrinsic growth rate and K_i is the carrying capacity. The parameters α_{ij} are interaction coefficients between species *i* and *j*.

The equilibria of equation (7) are determined by the values of K_i and α_{ij} . For each synthetic landscape, these parameters were randomly generated. The carrying capacities K_i were independently drawn from a gamma distribution with mean 1 and variance ζ^2 , while the interaction coefficients were drawn from a normal distribution with mean μ / N and variance σ^2 / N. These definitions ensure that the full range of distinct qualitative behaviours is spanned by parameter values of order 1 (ref. 51 for details). A broad, ecologically relevant regime of the interaction parameter space was explored in Fig. 4a ($\mu \in [-1, 4]$ and $\sigma \in [0, 1.5]$). The standard deviation of carrying capacities was fixed at $\zeta = 0.3$. The fourth and final parameter is interaction asymmetry, defined as $\gamma = \operatorname{corr}(\alpha_{ij}, \alpha_{ji})$. In Fig. 4a, interactions were set to be symmetric by taking $\gamma = 1$. These parameter values and ranges are identical to those used in Fig. S2 of ref. 35, allowing direct comparison with Fig. 4a. Supplementary Fig. 5 provides a more thorough parameter sweep varying both γ and ζ .

Landscapes were generated over a grid of points in μ - σ space. Each landscape was generated through the following steps:

- 1. A set of carrying capacity parameters K_i and interaction parameters α_{ij} were drawn for a pool of N = 10 species as described above. All r_i were fixed to 1.
- 2. Equation (7) was simulated to equilibrium for all species combinations.
- 3. Total endpoint 'biomass' (sum of abundances $\sum_i N_i(\infty)$) was computed for each simulation.
- 4. Exact, full-order landscapes (equation (1)) were computed, using biomass as the community function.

Initial species abundances were drawn from an exponential distribution with mean 0.1. Numerical integration was performed using ode15s (MATLAB). At each point in the μ - σ space, ten trials of landscapes were generated. Note that for larger values of σ , species abundances in equation (7) are more likely to diverge. Parameter combinations for which divergences were encountered in more than half of trials are indicated by grey values in Fig. 4a.

Simulations of the consumer resource model with random cross-feeding networks

Community-function landscapes were synthetically generated using a consumer resource model (CRM), taking the equilibrium concentration of a terminal waste product as the function of interest. The CRM is given as follows:

$$\begin{split} \dot{N}_{i} &= N_{i}(r_{i} - m_{i}), \\ r_{i} &= \sum_{\alpha} C_{i\alpha} R_{\alpha}, \\ \dot{R}_{\beta} &= \frac{K_{\beta} - R_{\beta}}{\tau_{\beta}} - \sum_{i} C_{i\beta} R_{\beta} N_{i} + \sum_{i} \gamma r_{i} N_{i} D_{i\beta}. \end{split}$$
(8)

Here N_i , r_i and m_i are the abundance, total resource uptake and maintenance costs, respectively, of species *i* in a pool of *N* total species. R_{α} is the concentration of resource α . The matrices $C_{i\alpha}$ and $D_{i\beta}$ describe which resources a species consumes and secretes, respectively, with secretions assumed proportional to the metabolic uptake r_i . An efficiency factor $\gamma < 1$ ensures that energy cannot be gained but only lost. Here γ was set to 0.5. The decay rates τ_{β} and species maintenance costs m_i are all set to 1 for simplicity. The resource carrying capacity for the single externally supplied resource, K_1 , was set to 10^5 , and all remaining resource carrying capacities, K_{β} , were set to 0. The total number of resources was fixed to L + 1, with resource L + 1 representing a terminal waste product that no species can consume.

Random cross-feeding networks were generated to explore how chains of trophic dependencies impact the ruggedness of community-function landscapes. To do this, it was assumed that each species *i* can consume and secrete exactly one resource, denoted in_{*i*} and out_{*i*}, respectively. These were selected through the following steps, performed independently for each species:

- 1. The identity of the consumed resource in, was chosen randomly between 1 and *L* with equal probability.
- The secreted resource out_i can range from 1 to L + 1, where resource L + 1 is the terminal waste product; however it must be distinct from in_i, leaving L possible choices. With probability p, we set out_i = in_i + 1; otherwise out_i was drawn from any of the remaining values at random.

The parameter p thus allows the exploration of a range of network topologies, from long linear pathways with a high degree of trophic dependency (at $p \approx 1$) to random graphs (at $p \approx 1/L$).

After selecting in_i and out_i for all species, we verified that resource L + 1 was 'reachable' through the network from resource 1, that is, whether there existed at least one path from resource 1 to resource L + 1. If this was not the case, the functional landscape (the concentration of resource L + 1 for a given set of species) would be identically zero; such networks were discarded as invalid and the steps above were repeated until a valid circuit was obtained.

For Fig. 4b, random cross-feeding networks were generated by carrying out the steps above across a range of values of p, fixing N = 10 total species and L = 8 total resources. Twenty valid random trials over 15 values of $p \in (1/L, 1)$ were generated to create 300 total random cross-feeding networks. The community-function landscapes for these networks were then computed by setting initial species abundances to 0.1, simulating equation (8) to equilibrium for all species combinations (ode15s, MATLAB), taking the equilibrium concentration of resource L + 1 as the function of interest and using this complete combinatorial landscape to determine the exact coefficients of its Fourier decomposition (equation (1)).

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Data analysed here are either available from the original studies or in the following repository: https://github.com/abbyskwara2/ regression_on_landscapes.

Code availability

Code to run all analyses presented in this paper is available in the following repository: https://github.com/abbyskwara2/regression_on_landscapes.

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Author contributions

S.K., M.T., K.G., M.Y., A. Skwara, A. Sanchez and A.S.R. conceptualized the study. A. Skwara, K.G., M.Y., M.T. and S.K. developed the methodology. A. Skwara, K.G., M.Y., M.T. and S.K. designed and conducted formal analysis. J.D.-C. and A. Sanchez designed experiments and collected experimental data, and J.D.-C. assisted with data analysis and visualization. K.G., S.K., M.T. and A. Skwara wrote the paper. S.K. and M.T. supervised the project and acquired funding.

Competing interests

The authors declare no competing interests.

Additional information

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Extended Data Fig. 1 | **Residuals obtained from second-order regressions.** Data from (A) Clark et al.²², (B) Langenheder et al.²⁷, (C) Sanchez-Gorostiaga et al.²¹, (D) Diaz-Colunga et al.²⁸, (E) data from the Sanchez lab, and (F) Kehe et al.²⁹.



Extended Data Fig. 2 | **Residuals obtained from second-order regressions as a function of community richness.** Data from (A) Clark et al.²², (B) Langenheder et al.²⁷, (C) Sanchez-Gorostiaga et al.²¹, (D) Diaz-Colunga et al.²⁸, (E) data from the Sanchez lab, and (F) Kehe et al.²⁹.



Extended Data Fig. 3 | **Model predictions from second-order regression fits** with all data in-sample and experimental replicates included. Data from (A) Clark et al.²², (B) Diaz-Colunga et al.²⁸, (C) data from the Sanchez lab, and (D) Kehe et al.²⁹. Black points correspond to the mean observed value for each distinct

experimental community, and are identical to the points shown in Extended Data Fig. S1. Gray points correspond to values for distinct experimental replicates. The datasets here include only datasets for which experimental replicates were measured.



Extended Data Fig. 4 | **Model predictions from second-order regression fits with all data in-sample.** Data from (A) Clark et al.²², (B) Langenheder et al.²⁷, (C) Sanchez-Gorostiaga et al.²¹, (D) Diaz-Colunga et al.²⁸, (E) data from the Sanchez lab, and (F) Kehe et al.²⁹.

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		Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information	about <u>availability of computer code</u>
Data collection	For the dataset collected for this study, optical density measurements with an AccuSkan FC plate reader (Fisher Scientific) were made with the standard Skanlt software v5.0. All simulation data was generated using MATLAB R2022b, and all associated scripts are available at https://github.com/abbyskwara2/regression_on_landscapes.
Data analysis	All dataset analyses were performed using R v4.2.1, R package tidyverse v2.0.0, and R package glmnet v4.1-7. All code and scripts necessary to reproduce these analyses are available at https://github.com/abbyskwara2/regression on landscapes.

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Data analyzed in this study is either available from the original studies or in the following repository: https://github.com/abbyskwara2/regression_on_landscapes.

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Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description	This is a method development study which made use of 6 datasets of combinatorial microbial communities, as well as simulated datasets of combinatorial microbial communities.				
Research sample	These datasets considered here were comprised of microbial communities for which a scalar-valued community function was measured for each distinct experimental community. These datasets included one dataset which was collected for this study, and 5 existing datasets from Langenheder et al. 2010, Clark et al. 2021, Sanchez-Gorostiaga et al. 2019, Diaz-Colunga et al. 2023, and Kehe et al. 2019. This study additionally generated simulated datasets of combinatorial microbial communities.				
Sampling strategy	No sample size calculations were performed for the dataset collected in this study. For this dataset, 160 distinct experimental communities were generated, a comparable number to similar studies.				
Data collection	For the dataset collected in this study, all experiments were performed in the Sanchez lab by Juan Diaz-Colunga.				
Timing and spatial scale	For the dataset collected in this study, all experiments were performed in January 2022.				
Data exclusions	No data were excluded from the analyses.				
Reproducibility	For the dataset collected in this study, the majority of experimental communities contained biological replicates. All replicates were included in the analysis, and all replicates are plotted in Supplementary Figure 2C.				
Randomization	The experimental communities for the dataset collected in this study were not randomized, as the experimental setup was designed to ensure that many different community compositions with a range of community richness would be tested.				
Blinding	For the dataset collected in this study, blinding during data collection was not possible as the researcher performing the experiments was required to know the identities of the microbes added to each experimental community.				
Did the study involve fiel	d work? Yes XNo				

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

n/a	Involved in the study		
\boxtimes	Antibodies		
\boxtimes	Eukaryotic cell lines		

X Palaeontology and archaeolog	\boxtimes
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\boxtimes		Animals	and	other	organisms
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- Clinical data
- Dual use research of concern
- Plants

Methods

n/a Involved in the study

- ChIP-seq
- Flow cytometry
- MRI-based neuroimaging