Predicting the First Steps of Evolution in Randomly Assembled Communities

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Abstract

Microbial communities can self-assemble into highly diverse states with predictable statistical properties. However, these initial states can be disrupted by rapid evolution of the resident strains. When a new mutation arises, it competes for resources with its parent strain and with the other species in the community. This interplay between ecology and evolution is difficult to capture with existing community assembly theory. Here, we introduce a mathematical framework for predicting the first steps of evolution in randomly assembled communities that compete for substitutable resources. We show how the fitness effects of new mutations and the probability that they coexist with their parent depends on the size of the community, the saturation of its niches, and the metabolic overlap between its members. We find that successful mutations are often able to coexist with their parent strains, even in saturated communities with low niche availability. At the same time, these invading mutants often cause extinctions of metabolically distant species. Our results suggest that even small amounts of evolution can produce distinct genetic signatures in natural microbial communities.

Main Text

Microbes often live in ecologically complex communities containing hundreds of coexisting species (1–4). As residents of these communities compete with each other, they can evolve over time by acquiring mutations (5–7). These evolutionary changes can alter the ecological interactions between species, driving shifts in community composition (8–10). Conversely, the community also creates the context in which organisms evolve, by influencing the structure of the adaptive landscape (11–15). Understanding how the community influences these evolutionary paths (and vice versa) is a crucial step toward predicting and controlling microbial ecosystems.

Longstanding conceptual models suggest that community structure can impact evolution in different ways. Some models predict that the rate of evolution should decline in larger communities, as more niches are filled by other species (7, 16–18). Others have proposed that diverse communities could create new opportunities for local adaptation, by suppress-

ing key competitors or creating new niches through crossfeeding (11–14, 19, 20). These conceptual models also make different predictions about how adaptive mutations will impact their community when they invade. Some mutations will replace their parent strain, while others can encroach on other species (9, 21) or diversify into coexisting lineages (5, 6, 22). Each of these behaviors has been observed empirically, yet it remains challenging to predict which should dominate in a given community.

The source of this challenge lies in the niches inhabited by different species, and how mutations alter or move between them. While much progress has been made in small communities where the relevant niches can be explicitly defined (21–24), it is difficult to extend this approach to larger communities like the gut microbiome, where species can compete for many different combinations of resources. In this high-diversity limit, even basic questions about the effects of community structure remain unresolved: how does the benefit of a mutation depend on the diversity of the community and the metabolic overlap between its members? Do mutations primarily compete with their parent strain, or do they continue to stably diversify, as suggested by recent evidence from the gut microbiome (5, 6, 25) and some *in vitro* communities (26, 27)?

Resource competition models provide a mechanistic framework to address these questions (28–32). In these models, niches are not defined in advance, but emerge organically through differences in resource consumption (33). An extensive body of work has used this framework to investigate the process of community assembly, where species compete to colonize a new environment (28, 34-37). These model communities can recapitulate some large-scale features of natural (38–40) and experimental ecosystems (39–42). By contrast, the evolutionary dynamics that emerge from resource competition are difficult to model with traditional community assembly theory. While some studies have started to explore these effects, previous work has mostly focused on small communities (30, 31, 43) or the long-term states attained over infinite evolutionary time (30, 44). Both approaches are poorly suited for understanding how a focal species evolves in different community backgrounds, which is the scenario most accessible in experiments. To address this gap, we develop a theoretical framework for predicting the initial steps of evolution in an assembled community with many coexisting species. By extending random matrix approaches from community assembly theory, we derive analytical predictions that describe how the fitness benefits and fates of new mutations scale with the diversity and metabolic overlap of the surrounding community, enabling quantitative tests of the conceptual models above.

Modeling first-step mutations in randomly assembled communities

To study the interplay between ecological competition and new mutations in a mathematically tractable setting, we turn to a simple resource competition model (28–32, 36, 45) where microbes compete for $\mathcal{R} \gg 1$ substitutable resources that are continuously supplied by the environment (Fig. 1A). Each strain μ in the community is characterized by a resource utilization vector $\vec{r}_{\mu} = (r_{\mu,1}, \dots, r_{\mu,\mathcal{R}})$, which describes how well it can grow on each of the supplied resources. Following previous work (30, 32), it will be convenient

to further decompose this vector into an overall magnitude $X_{\mu} \equiv \log(\sum_{i} r_{\mu,i})$, which represents a strain's resource consumption budget, and a normalized resource strategy $\vec{\alpha}_{\mu} \equiv \vec{r}_{\mu} / \sum_{i} r_{\mu,i}$, which describes the relative effort devoted to importing each of the resources. For simplicity, we neglect additional factors like regulation (46), crossfeeding (39), and resource sequestration (47), allowing us to focus on the fundamental evolutionary pressures imposed by resource competition alone.

With these assumptions, the relative abundances of the strains (f_{μ}) follow the dynamical equations,

$$\frac{\partial f_{\mu}}{\partial t} = f_{\mu} \left[\sum_{i=1}^{\mathcal{R}} \alpha_{\mu,i} e^{X_{\mu}} h_i(\vec{f}) - 1 \right], \tag{1}$$

where $h_i(\vec{f}) \equiv (\kappa_i / \sum_j \kappa_j) \cdot (\sum_\mu \alpha_{\mu,i} e^{X_\mu} f_\mu)^{-1}$ denotes the local availability of resource i, and κ_i is its external supply rate (SI Section 1.1). Previous work has used this model to study the process of community assembly (28, 36, 37), where S distinct species arrive in a new environment, and form an ecologically stable community containing $S^* \leq \mathcal{R}$ survivors (Fig. 1B, left). The value of S^* depends on the environment and the resource preferences \vec{r}_μ of the S initial strains. While it is difficult to measure these kinetic parameters directly, past research has shown that emergent features of large ecosystems can often be mimicked by randomly drawing the uptake rates from a common statistical distribution (28, 34–36). For concreteness, we initially focus on the binary resource usage model from Ref. (36), in which each strain utilizes a random subset of $\sim \mathcal{R}_0$ resources, with an overall uptake budget X_μ drawn from a Gaussian distribution (SI Section 1.1); we also consider several alternative choices in Figures S2 and S6.

Individual realizations of this model produce assembled communities with similar numbers of surviving species (S^*), which can be predicted using methods from the physics of disordered systems (28, 29, 36; Fig. 2B; SI Section 3). In our analysis below, it will be convenient to treat the expected number of surviving species as an input parameter, and classify the assembled communities as a function of their niche saturation, S^*/\mathcal{R} (SI Section 3.2).

To account for evolution, we model the very first mutational steps that occur in a randomly assembled community (Fig. 1B). This scenario might apply to the initial phases of *in vitro* passaging experiments (12, 26), or recently colonized gut microbiomes (48, 49). Through much of our analysis, we will focus on a particularly simple class of "knock-out" mutations, where the mutant loses its ability to consume one resource ($\alpha_i \rightarrow 0$). We assume that the cell can compensate for this deletion by increasing the uptake of other resources in its repertoire, but this compensation may not be perfect, corresponding to a shift in the effective budget (ΔX). We also consider "knock-in" mutations, where a strain gains the ability to consume a resource (e.g. through horizontal gene transfer; 25), as well as more general changes that influence the uptake rates of multiple resources simultaneously ($\vec{\alpha} \rightarrow \vec{\alpha} + \Delta \vec{\alpha}$). We can classify these mutations by the magnitude of their phenotypic change, $\gamma \equiv \mathcal{R}_0 \sqrt{\sum_i \Delta \alpha_i^2}$, defined such that $\gamma \approx 1$ for knock-out mutants.

While the phenotypes of these mutants are simple, their fitness effects depend on the local



Figure 1: Modeling the first steps of evolution in a randomly assembled community that competes for substitutable resources. (A) Microbial strains compete for \mathcal{R} resources that are continuously supplied by the environment at rates κ_i . Each strain μ has a characteristic set of uptake rates $r_{\mu,i}$ (arrows), which can be altered by further mutations. (B) A local pool of S initial species, whose phenotypes are randomly drawn from a common statistical distribution, self-assembles into an ecological equilibrium with $S^* \leq S$ surviving species (left). A new mutation (M) arises in one of the surviving species (P); if the mutation provides a fitness benefit, its descendants can either replace the parent strain (top right) or stably coexist with the parent, potentially driving another species to extinction (bottom right).

environment, which is shaped by the other resident strains. Using the assembled communities as a backdrop allows us to quantify how this evolutionary landscape varies with the size and composition of the surrounding community.

Distribution of fitness effects of mutations in newly assembled communities

The rate of evolution in a newly assembled community will depend on the supply of beneficial mutations. This landscape is often summarized by the local distribution of fitness effects (DFE), denoted by $\rho_{\mu}(s)$, which gives the relative probability that a mutation in focal strain μ will have an invasion fitness s. The shape and scale of $\rho_{\mu}(s)$ determine the availability of beneficial mutations, and therefore the rate of evolutionary change. Our resource competition framework allows us to ask how these features depend on the composition of the larger community.

For a community at ecological equilibrium, a mutation that arises in a resident strain μ and changes its resource uptake phenotype to a new value $X_{\mu} + \Delta X$ and $\vec{\alpha}_{\mu} + \Delta \vec{\alpha}$ will have an invasion fitness,

$$s_{\rm inv} \approx \Delta X + \sum_i \Delta \alpha_i g_i \,,$$
 (2)

where $g_i \equiv h_i - \overline{h}$ is the excess availability of resource *i* relative to the ecosystem average, $\overline{h} \equiv \frac{1}{\mathcal{R}} \sum_i h_i$ (30) (SI Section 1.2). Equation (2) shows that the invasion fitness of a mutation that only affects the overall uptake budget of a strain ($\Delta \vec{\alpha} = \vec{0}$) is independent of the surrounding community. In contrast, the benefits of mutations that change the resource consumption strategy of a strain will depend on the interactions between species, which are mediated by the values of the resource availabilities, g_i . For example, if a resource has a lower relative availability ($g_i < 0$), then it is not worth devoting energy to consume it, and a knock-out mutation for that resource should be beneficial.

Replica-theoretic calculations similar to those performed in Ref. (36) allow us to predict the joint distribution of g_i as a function of the community assembly parameters (SI Section 3.2). In a large ecosystem, the excess availabilities are well-approximated by a Gaussian distribution,

$$g_i \sim \left(1 - \frac{S^*}{\mathcal{R}}\right) \left[\delta \kappa_i + Z_i \cdot C \cdot \sqrt{\frac{\mathcal{R} - \mathcal{R}_0}{S^* \mathcal{R}_0}}\right],$$
(3)

where $\delta \kappa_i \equiv \kappa_i - 1$ is the external supply of each resource, *C* is an $\mathcal{O}(1)$ factor which depends on $\mathcal{S}/\mathcal{S}^*$ (SI Section 3.2), and the Z_i are uncorrelated standard Gaussians. Eqs. (2) and (3) allow us to calculate the distribution of fitness effects, $\rho_{\mu}(s)$, by aggregating over mutations with different values of $\Delta \vec{\alpha}$ and ΔX .

To understand how the community influences the DFE, it is helpful to consider the simplest case, where the resource supply is uniform ($\delta \vec{\kappa} = \vec{0}$) and mutations have no direct costs or benefits ($\Delta X = 0$). In this case, Eqs. (2) and (3) imply that $\rho_{\mu}(s)$ will also follow a



Figure 2: Distribution of fitness effects of first-step mutations as a function of commu**nity complexity.** (A) As in Fig. 1, a community is assembled from S initial species, leaving S^* alive at equilibrium. The surviving species produce mutations, whose invasion fitness s_{inv} is equal to their initial relative growth rate. (B) Number of surviving species (S^*) as a function of the sampling depth (S) and the inter-species variation in the total uptake budget (ϵ). Curves show theory predictions from SI Section 3.1, while points show means and standard deviations over 10^3 simulation runs with $\mathcal{R} = 200$, $\mathcal{R}_0 = 40$, and $\sigma_{\kappa} = 0$. (C) Distribution of fitness effects of knock-out (and knock-in) mutations with $\Delta X = 0$ in communities with two different levels of niche saturation (S^*/R). Black curve shows the theoretical predictions from Eq. (4), while dots represent a histogram over all possible strategy mutations in 10^3 simulation runs using the same parameters as panel B, with $S^*/S = 0.1$. (D) The width σ_{inv} of the distribution of fitness effects in panel C as a function of niche saturation, for various values of sampling permissivity S^*/S and per-species resource usage \mathcal{R}_0 . Curves show the theoretical predictions from Eq. (4), while the dots show the average over 10^3 simulation runs. (E) Pearson correlation between the fitness effect of a mutation in the community and its fitness effect in monoculture, for different values of niche saturation and scaled variation in resource supply rates, $\operatorname{Var}_{env}/\operatorname{Var}_{comm} \equiv \sigma_{\kappa}^2 \cdot \mathcal{R}_0/(1 - \mathcal{R}_0/\mathcal{R})$. Curves show the theoretical predictions from SI Section 3.5, while points show the average over all mutations in 10^3 simulated communities with parameters the same as panel C.

Gaussian distribution, with mean zero and standard deviation

$$\sigma_{\rm inv} \sim \gamma \cdot C \left(1 - \mathcal{S}^* / \mathcal{R} \right) \sqrt{\frac{\mathcal{R} - \mathcal{R}_0}{\mathcal{S}^* \mathcal{R}_0^3}},\tag{4}$$

where $\gamma \equiv \mathcal{R}_0 \sqrt{\sum_i \Delta \alpha_i^2}$ is the overall magnitude of the phenotypic change produced by the mutation (Fig. 2C-D). Since this result does not explicitly depend on the focal species μ , it implies that the DFEs should be similar for all strains in the community (Fig. S1). Furthermore, since the specific details of the mutations only enter through their overall magnitude γ , this implies that knock-out and knock-in mutations — as well as multi-resource mutations with the same value of $\gamma \approx 1$ — will have statistically similar DFEs (Fig. 2C).

Equation (4) shows that the community influences the DFE primarily through the degree of niche saturation, S^*/\mathcal{R} . The magnitude of the typical fitness effect approaches zero as $S^* \to \mathcal{R}$ (Fig. 2D), which is consistent with the idea that the rate of evolution will be slower in communities where more niches are already filled. However, since the mean of the DFE is still centered at s = 0, the overall fraction of beneficial mutations remains constant as $S^* \to \mathcal{R}$ (Fig. 2C). This implies that surviving organisms are not necessarily at an "evolutionary optimum," where any change to their resource consumption tends to be deleterious.

However, this symmetry between the frequency of beneficial and deleterious mutations critically depends on the assumption of perfect trade-offs ($\Delta X = 0$), which might not hold in practice. For example, a beneficial mutation could halt the production of an enzyme which is used for metabolizing a low-availability resource, while leaving the expression of other enzymes in that now-defunct pathway intact – resulting in a net cost to pure fitness. If all mutations carried such a direct cost ($\Delta X < 0$), then Eq. (2) implies that the entire DFE would shift to the left by a constant amount $-|\Delta X|$. If this shift is larger than the typical spread of the DFE ($|\Delta X| \gg \sigma_{inv}$), then the beneficial tail of $\rho_{\mu}(s)$ will approach an exponential shape, whose height and width will both strongly decline with the degree of niche saturation S^*/\mathcal{R} . Thus, the availability of beneficial mutations can sensitively depend on the genetic architecture underlying resource consumption.

Similar results apply when the resources are supplied at different rates ($0 < |\delta \kappa_i| \ll 1$; SI Section 3.5). In this case, the fitness effect of a mutation will depend on both the community and the external environment, as encoded by the resource availabilities in Eq. (3). The overall magnitude of the environmental contribution declines as niche saturation increases $(\mathcal{S}^*/\mathcal{R} \to 1)$, consistent with previous work showing that larger communities self-organize to "shield" their member species from the external environment (30, 32, 36). Interestingly, however, Eq. (3) shows that the *relative* contribution of $\delta \vec{\kappa}$ to a mutation's fitness effect actually increases with niche saturation, so the external environment can still exert an influence on the overall direction of the fitness landscape. This effect is strikingly illustrated when we compare the fitness effect of each mutation with its expected value in the absence of the community (Fig. 2E). The correlations between the two DFEs can be substantial when $\sigma_{\kappa}^2 \gtrsim \mathcal{R}/\mathcal{S}^*\mathcal{R}_0$, indicating that environmental shielding is often incomplete.

Ecological diversification in large communities

While the invasion fitness describes the initial growth of an adaptive mutation, a successful variant will eventually reach a size where it starts to impact the other members of the community. Some of these mutants will eventually replace their parent strain, due to the principle of competitive exclusion (28). Other mutations can stably coexist with their parent by exploiting a different ecological niche (30). How does the frequency of these ecological diversification events depend on the composition of the surrounding community?

We can analyze the probability of mutant-parent coexistence in our model by recasting it as the outcome of two correlated assembly processes. First, an initial ecosystem E_0 is formed through our standard community assembly process (Fig. 1B, left). Then, a second ecosystem is formed when one of the surviving species in E_0 produces a beneficial mutant M, and the combined community $E_0 + M$ is allowed to reach its new ecological equilibrium (Fig. 1B, right). The latter event requires that the parent strain has a positive relative abundance in the first ecosystem ($f_P^{E_0} > 0$), and that the mutant survives in the second ecosystem ($f_M^{E_0+M} > 0$). The probability that the mutant coexists with its parent can then be expressed as a conditional probability,

$$\mathbb{P}_{\text{coex}} = \mathbb{P}[f_P^{E_0+M} > 0 \mid f_P^{E_0} > 0, \ f_M^{E_0+M} > 0], \tag{5}$$

which averages over the random resource uptake rates in the initial community, as well as the random effect of the adaptive mutation.

The correlated assembly process in Eq. (5) is challenging to analyze (37), since the mutant and parent phenotypes are closely related. Fortunately, we will show that we can often approximate the coexistence probability by considering a third community assembly process, in which the mutant and parent are introduced simultaneously with the other species. This approximation differs from the two-ecosystem model in Eq. (5), since the final community can contain "rescued" species that would not have survived in E_0 before the mutant strain invaded. However, simulations and theory indicate that these differences result in only small corrections to the coexistence probability across a broad range of parameters (SI Section 1.3, Fig. S3), so that the simultaneous assembly approximation is often valid.

When this approximation holds, the coexistence probability can be evaluated by extending the replica-theoretic calculations in Eq. (3) (SI Section 3.4). We find that the coexistence probability can be expressed in terms of quantities from the original ecosystem, before mutant invasion:

$$\mathbb{P}_{\text{coex}} \approx \int_0^1 df_P \, p(f_P) \int_0^{s_{\text{coex}}(f_P)} ds \, \rho(s|f_M > 0) \,, \tag{6}$$

where $p(f_P)$ is the distribution of relative abundance of the parent (conditioned on survival), $\rho(s|f_M > 0) \propto \max(0, s) \cdot \rho_{\mu}(s)$ is the distribution of invasion fitnesses of mutants that survive genetic drift (30), and $s_{\text{coex}}(f_P)$ is defined by

$$s_{\text{coex}}(f_P) \equiv \frac{f_P \mathcal{S}^* \gamma \sigma_{\text{inv}}}{\sqrt{\mathcal{R}_0\left(\frac{\mathcal{S}^*}{\mathcal{R}}\right) \left(1 - \frac{\mathcal{R}_0}{\mathcal{R}}\right)}}.$$
(7)

Since $\mathcal{R}_0 \gg 1$, the integral in Eq. (6) reduces to

$$\mathbb{P}_{\text{coex}} \approx \frac{\rho_{\mu}(0) \cdot \sigma_{\text{inv}}^2}{\int_0^\infty s \rho_{\mu}(s) \, ds} \cdot \frac{\gamma^2}{\mathcal{R}_0(\mathcal{S}^*/\mathcal{R})(1 - \mathcal{R}_0/\mathcal{R})},\tag{8}$$

where σ_{inv} is the width of the DFE from Eq. (2). For a mutation with perfect trade-offs ($\Delta X = 0$), the first factor is an O(1) constant, so the coexistence probability is dominated by the second factor.

Equation (8) shows that the coexistence probability is largest for small values of S^*/\mathcal{R} (Fig. 3A), consistent with the idea that diversification is easier when there are many open niches left to exploit, as in the case of adaptive radiation (20). Interestingly, however, we find that the coexistence probability does not vanish even as communities approach full saturation $(S^*/\mathcal{R} \to 1)$, but instead plateaus at a nonzero value. This is true even in fully saturated communities $(S^* = \mathcal{R})$, where other species must be driven to extinction when the successful mutant invades.

At this point, the coexistence probability is most strongly determined by the phenotypic effect size of the mutation. A mutation that changes the resource consumption strategy infinitesimally ($\gamma \rightarrow 0$) can never coexist with its parent. By contrast, the coexistence probability of a single-resource knockout mutant ($\gamma = 1$) scales inversely with the total number of resources \mathcal{R}_0 utilized by a typical organism, rather than the total number of resources in the environment. This suggests that mutants and parents can coexist even for "large" communities containing many species and resources (Fig. 3B and Fig. S4).

We see that simulation results generally support the theoretical predictions in Eq. (8), but start to exceed this value for communities very close to full saturation. The reason for this deviation comes from our "simultaneous assembly" approximation, which allowed the mutant's invasion to "rescue" species that had previously gone extinct. Allowing extinct species to re-invade flattens the mutant-parent coexistence probability (Fig. S3), suggesting that competition from these rescued species plays a significant role in our theoretical model for extremely high niche saturation. These rescued species could be relevant in some natural ecosystems, where a local species pool is maintained in separate spatial niches, with frequent migration events allowing species to re-invade. Regardless, our theoretical results provide a lower bound on the coexistence probability for large ecosystems, so that diversification could be even more common under some conditions.

In reality, most mutations that change an organism's resource consumption strategy are unlikely to be perfect trade-offs. More generally, we find that mutations with a direct cost or benefit ΔX change the coexistence probability by the factor

$$\frac{\mathbb{P}_{\text{coex}}(\Delta X)}{\mathbb{P}_{\text{coex}}(\Delta X=0)} \sim \begin{cases} \frac{\sigma_{\text{inv}}}{\sqrt{2\pi\Delta X}} e^{-\Delta X^2/2\sigma_{\text{inv}}^2} & \text{if } \Delta X \gg \sigma_{\text{inv}}, \\ 1 - \sqrt{\frac{\pi}{2}} \frac{\Delta X}{\sigma_{\text{inv}}} & \text{if } |\Delta X| \ll \sigma_{\text{inv}}, \\ \Delta X^2/\sigma_{\text{inv}}^2 & \text{if } \Delta X < 0, |\Delta X| \gg \sigma_{\text{inv}}. \end{cases}$$
(9)

This result shows that strategy mutations which carry a direct fitness cost are more likely to coexist with their parent strain, provided that they are still favored to invade in the current

community context. Conversely, strategy mutations with strong pure fitness benefits are likely to drive their parent to extinction (Fig. 3B). This effect arises from the fact that a nonzero ΔX changes the typical fitness of mutations that establish in the population. The coexistence probability is highly sensitive to these changes in invasion fitness, as shown by the bounds of integration in Eq. (6): mutants with a smaller invasion fitness are much more likely to coexist with their parent strain (Fig. 3C, inset). Coexistence is unlikely when invasion fitness is above a characteristic level

$$\bar{s}_{\text{coex}} \sim \gamma^2 \frac{1 - \mathcal{S}^* / \mathcal{R}}{\mathcal{R}_0^2 (\mathcal{S}^* / \mathcal{R})}.$$
 (10)

This "coexistence fitness" \bar{s}_{coex} is smaller than the typical invasion fitness σ_{inv} by a factor of $\mathcal{R}_0^{-1/2}$ – meaning that diversification is driven by a narrow range of mutants with fitness high enough to establish but small enough to coexist.

We can gain some intuition for this effect by considering a loss-of-function mutation for a resource that is slightly overutilized by the population ($g_i < 0$). Equation (1) shows that the growth rate of this variant (relative to its ancestor) is given by $\sim g_i(t)$. As the mutant grows in abundance, it begins to replace its parent, which tends to reduce the overall utilization of resource i ($\partial_t g_i > 0$). If g_i reaches zero before the parent strain goes extinct, then the mutant will coexist with its parent, having lost its initial growth advantage. This coexistence is most likely to happen if the parent strain has high abundance, or if $|g_i|$ (and therefore s_{inv}) is initially small (Fig. S5).

The same argument applies for more complex mutations which affect multiple resources. It also explains why mutations with small phenotypic effects, $\gamma \rightarrow 0$ (e.g. a pure fitness mutation with no strategy change), cannot coexist with their parent. If the mutant and parent have near-identical resource consumption strategies, the mutant's invasion produces very little negative feedback on its growth rate relative to its parent, making it likely the parent strain will be driven to extinction (Fig. S4B). Similar logic applies for extensions of the model discussed in SI Section 3.5, such as non-uniform resource supply or allowing the parent species to use a different number of resources from the rest of the population (Fig. S6). Together, these results suggest that *in situ* diversification could be common even in large and saturated communities, particularly for mutants on abundant backgrounds with lower-than-expected invasion fitnesses.



Figure 3: Ecological diversification in large communities. (A-C) Probability that a successful knock-out mutant coexists with its parent strain as a function of (A) the niche saturation S^*/\mathcal{R} , (B) the typical number of resources used per species \mathcal{R}_0 , and (C) the change in overall uptake budget of the mutant ΔX . Inset shows the dependence on the total invasion fitness s_{inv} . In all panels, curves show the theory predictions from SI Sections 3.3 and 3.4, while points show means and standard errors over 10^4 simulation runs with base parameters $\mathcal{R} = 200$, $\mathcal{R}_0 = 40$, $S^*/S = 0.1$, and $\sigma_{\kappa}^2 = 0$.

Successful mutations drive extinctions in other niches

In addition to displacing their parents, successful mutants can also drive other species in the community to extinction. This is particularly evident in saturated communities $(S^* = R)$, where competitive exclusion implies that the invasion of a new strain must be accompanied by extinction of at least one other. Figure 4B shows that the number of extinctions steadily rises with the degree of niche saturation, exceeding 1% of the initial community (~1-10 species) for many combinations of parameters. Moreover, these extinctions are not completely independent of each other, since they are somewhat overdispersed compared to a simple Poisson expectation (Fig. 4B, inset). Previous work has suggested that even distantly related strains could exhibit correlated dynamics if their resource consumption strategies are anomalously similar to each other (26). Could such hidden metabolic similarity be responsible for driving the extinction events above?

We tested this idea by examining the number of resources that were jointly consumed by the invading and displaced strains (a proxy for their overall metabolic similarity). Interestingly, we found that the number of resources shared with the mutant was not substantially higher for the displaced species, and was comparable to a randomly drawn species from the larger community (Fig. 4C). This suggests that the extinction events in Fig. 4 cannot be explained by traditional measures of niche overlap (33). Rather, successful mutants can displace species outside of their apparent niche, even when they stably coexist with a more metabolically similar parent.

Since extinctions were not well-predicted by their overall metabolic similarity to the mutant, we conjectured that these displaced species may possess other features that render them vulnerable to extinction. For example, low-abundance species may be less welladapted to the current environment, and thus sensitive to perturbations like the invasion of a mutant. Consistent with this hypothesis, we found that the extinction probability for very low-abundance species is much higher than the community average, approaching \sim 50% at the highest levels of niche saturation (Fig. 4D). Furthermore, although the displaced species are metabolically distant from the invading mutant, we find they are more likely to share the mutant's strategy for the resource targeted by the mutation. Displaced species are more likely to use the resource gained by a successful "knock-in" mutation, and are correspondingly less likely to use the resource lost in a successful knock-out strain (Fig. 4E). These results illustrate that in high-dimensional ecosystems, the invasion of a new mutant can have a small impact on a diverse range of metabolic strategies. If a resident strain is maladapted enough to already be on the edge of extinction, the invasion of a mutant can be enough of a perturbation to displace it from the community. Thus, the abundance of an organism might often be a better predictor of its fate than its apparent metabolic niche.



Figure 4: Successful mutations drive extinctions of metabolically distant species. (A) Schematic showing the extinction of an unrelated species (blue) after a beneficial knockout mutation (orange) invades. In this example, the displaced species and the mutant share one common resource, but not the one targeted by the knock-out mutation. (B) Average number of extinctions after the invasion of a successful mutant, as a function of niche saturation $\mathcal{S}^*/\mathcal{R}$; parent strains are excluded from the extinction tally. Inset: full distribution of extinctions for the starred parameters, compared to a Poisson distribution with the same fraction of zero counts. Points denote the averages over 10^4 simulation runs with the same base parameters as Fig. 3. (C) Distribution of the number of resources jointly utilized by the displaced species and the invading mutant (parent strains excluded). Points denote the results of simulations with $S^*/\mathcal{R} = 0.9$. Gray curves show the analogous background distribution between the mutant and all other species in the community, regardless of whether they become extinct. (D) Probability of extinction as a function of initial relative abundance for the starred point in panel B. (E) The fold change in probability that the displaced species uses the same resource targeted by the mutant (i.e., the resource being knocked in or out), relative to the background distribution of resource use in the population. Points denote means and standard errors for knock-in and knock-out mutations as a function of \mathcal{R}_0 for $\mathcal{S}^*/\mathcal{R} = 0.8$, with the the remaining parameters the same as panel B.

Robustness of ecosystem to subsequent mutations

Our analysis above focused on the first wave of mutations arising in a newly assembled community, where the initial states could be predicted using existing community assembly theory (28, 29, 35, 36). However, subsequent waves of evolution could eventually drive the ecosystem away from this well-characterized initial state (30). To assess the robustness of our results under the acquisition of further mutations, we simulated successive waves of mutations using a generalization of the approach in Fig. 3. We considered the simplest case where resident strains could generate knock-out mutations on any of the resources they currently utilized. We also assumed that the dynamics were mutation-limited (30, 31), so that the community relaxes to a well-defined steady state between each successive mutation. We continued this process until one of the surviving strains had accumulated 10 mutations in total, which typically corresponded to 100-200 successful mutations in the larger community.

We first asked how these subsequent waves of mutations altered the genetic structure of their community. While the total number of surviving strains decreased slightly over time (eventually stabilizing at an intermediate value), the number of strains related through *in* situ diversification events increased approximately linearly over the same time window (Fig. 5B). This indicates that the ecological diversification events in Fig. 3 continue to occur at a high rate even after additional mutations have accumulated, more quickly than individual branches of diversified lineages go extinct. Nonetheless, the abundance trajectories in Figure 5A indicate that these extinction events among close relatives are not uncommon. For example, Lineage 1 seeds eight ecological diversification events over the course of the simulation, but only three of these strains are alive at the end of the simulation. On average, we find that newly diversified lineages coexist for \sim 40 mutational steps before one of them goes extinct (Fig. 5C). Nonetheless, longer coexistence is possible: one of the diversification events in Fig. 5A is maintained for >150 mutational steps, enough time for multiple additional mutations to accumulate within each lineage. Thus, coexisting mutant-parent pairs are often maintained through further evolutionary perturbations, even as the total number of species stabilizes over time.

Our first-step analysis showed that mutations in more abundant strains were more likely to coexist with their parent. This prediction is borne out by our multi-step simulations as well: while some highly abundant lineages seeded many *in situ* ecological diversification events (e.g. Lineage 1 in Fig. 5A), the typical surviving lineage experienced no more than one (e.g. Lineage 2), and many species went extinct without diversifying at all (e.g. Lineage 3). Consistent with our earlier predictions, lineages at low abundance were more prone to extinction when a new mutant invaded, while coexistence events tended to happen to high-abundance lineages (Figure 5A, right). The combination of these factors creates a "rich get richer" effect where high-abundance species diversify at the expense of driving low-abundance species to extinction (30).

Finally, we asked how the accumulation of mutations shifts the landscape of beneficial mutations from the initial "assembled" state in Figs. 2 and 3. We found that the distribution of fitness effects does not dramatically differ between the start and the end of the



Figure 5: Mutation and diversification over longer evolutionary timescales. (A) Left: an example simulation showing the step-wise accumulation of ~100 adaptive knock-out mutations in a community with $\mathcal{R} = 100$, $\mathcal{R}_0 = 20$, $\mathcal{S}^*/\mathcal{R} = 0.9$, $\mathcal{S}^*/\mathcal{S} = 0.1$. Solid lines denote the abundance trajectories of 3 example lineages. Large points indicate extinction events in these lineages (red), diversification events (green), and mutation events that displace their parent strain (blue) in the highlighted lineages, while smaller points indicate analogous events for other species in the community. Dashed lines illustrate offshoots of the highlighted lineages that eventually went extinct. Right: Relative abundances of strains when they experienced mutation, diversification and extinction events, respectively. Lines denote histograms aggregated over 10 simulation runs. (B) Left: Total number of surviving strains over time. Grey lines denote replicate simulations for the same parameters in panel A, while their average is shown in black. Right: Total number of strains related to another surviving strain through one or more *in situ* diversification events. (C) The probability of mutant-parent coexistence being maintained (i.e., both lineages surviving) as a function of time since initial divergence, over ten simulations. (D) The distribution of fitness effects at the start of the simulation (red) and after 90 accumulated mutations (blue), compared to the first-step predictions from Fig. 2 (black). (E) The probability that a mutation event leads to stable diversification as a function of time. Black points denote binned values aggregated over ten replicate simulations, while red point denotes the analogous result for 10^4 first-step simulations.

simulation, although its shape changes slightly from a Gaussian distribution toward a twotailed exponential (Fig. 5D). In contrast, the impacts of successful mutations exhibit larger changes: the proportion of mutation events which result in mutant-parent coexistence increases substantially with evolutionary time, rising from about 20% to 80% over the course of the simulation (Fig. 5E). This shift cannot be explained by changes in the overall number of species or the availability of beneficial mutations, but instead deviates entirely from the replica-theoretic predictions that describe our initial state. Thus, while many our qualitative conclusions continue to hold on longer evolutionary timescales, the accumulation of mutations can lead to quantitative differences from our first-step analysis above. In both cases, the evolved communities exhibit distinct genetic signatures compared to their purely assembled counterparts, which could motivate future tests of *in situ* evolution.

Discussion

Large ecological communities apply complex evolutionary pressures to their resident species, leading to controversy about how these organisms' evolutionary trajectories are affected by the background community (17). Here, we address this challenge by developing a theoretical framework for predicting the first steps of evolution in randomly assembled communities that compete for substitutable resources. This provides a quantitative foundation for understanding how the fitness benefits and fates of new mutations should scale with the diversity and metabolic overlap of the surrounding community.

Our results show that the supply of beneficial mutations does not necessarily run out in larger communities – as expected in the simplest models of niche filling – but rather that the benefits of these mutations systematically decline with S^*/R . We also find that the fitness effects of mutations can be broadly correlated with the external environment, even in large communities where internal resource concentrations are shielded from environmental shifts (32, 36). These distributions of fitness effects can be measured in modern experiments, e.g. by performing barcoded fitness assays in communities of different size (12, 48).

Our finding that successful mutants often coexist with their parents is reminiscent of empirical observations from the gut microbiome, where recently diverged strains differing by only a handful of mutations appear to stably coexist within their host (5, 6). While spatial structure could also contribute to this coexistence (27, 50), our model demonstrates that *in situ* diversification can continue to occur even in a well-mixed environment when most niches are already filled. Similar behavior has also been observed in generalized Lotka-Volterra models with spatiotemporal chaos (51). This suggests that ongoing diversification may be a generic feature of large microbial communities, providing an alternative mechanism for the "diversity begets diversity" hypothesis (17, 52, 53).

Beyond diversification, Figure 5 demonstrates that successful mutants can also drive distantly related species to extinction. This behavior is consistent with recent empirical observations in the human gut microbiome (8) and strain-swapping experiments in synthetic gut communities (54). It could also contribute to the extinction events observed in community passaging experiments (55). The fact that the displaced species are metabolically diverged from the invading mutants creates obstacles for inferring these interactions from metabolomics measurements (56) or metabolic reconstructions (57). Our findings suggest that future efforts should instead focus on how the invading mutant impacts low-abundance species more generally, and whether they utilize the specific resource(s) targeted by the mutation. These results align with recent work emphasizing the importance of collective interactions in shaping microbial community dynamics (58).

Here, we focused on the simplest possible resource competition model, neglecting important factors like crossfeeding (59), spatial structure (60), and metabolic regulation (46), which are all thought to play key roles in natural microbial communities. We also assumed a simplified model of evolution that does not account for competition between simultaneously occurring mutations. These clonal interference effects can enhance diversity by allowing strains to temporarily evade competitive exclusion (31). On the other hand, competition between lineages also selects for more strongly beneficial mutations (61), which we predict are less likely to coexist with their parent. Our results provide a baseline for incorporating these effects, which will be crucial for understanding how large microbial communities will evolve.

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Data and materials availability: Source code for community assembly simulations and numerical calculations are available at Github (https://github.com/jdmcenany/First_Step_Muts).

Supplementary Information

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4 Numerical Simulations

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Figure S1: Distribution of invasion fitnesses for knock-out and knock-in strategy mutations of a single organism within a single sampled community. Black curve shows Gaussian theory prediction, while dots are histogram values over all possible strategy mutations. Simulation was run for $S^*/\mathcal{R} = 0.8$, $\mathcal{R} = 200$, $\mathcal{R}_0 = 40$, $S^*/\mathcal{S} = 0.1$, and $\sigma_{\kappa}^2 = 0$.



Figure S2: Our main conclusions about mutation in assembled communities can be brought into qualitative and quantitative accordance with other consumer resource schemes. (A) Distribution of invasion fitnesses for knock-in and knock-out strategy mutations for the Tikhonov consumer resource model (36), which breaks knock-in/knock-out symmetry. Black curves show theoretical prediction, while points show simulation results. (B) Mutant-parent coexistence probability for the Tikhonov CRM as a function of niche saturation, S^*/R . The average fitness benefit (cost) for knock-out (knock-in) mutations effectively gives strategy mutations a nonzero change in pure fitness, which lowers (raises) the theoretically predicted coexistence probability relative to the neutral case. Red and blue curves show these adjusted predictions, while crosses and pluses show simulated results. (C) Distribution of resource availability h_i for an ecosystem assembled with strategy vectors drawn from the Dirichlet distribution as described in SI Section 4. Black curves show theoretical prediction from the binary resource use case studied in this paper, while points show simulated results. (D) Probability of mutant-parent coexistence for knock-out strategy mutations in this Dirichlet assembled community. Black curve shows the theoretical prediction for binary resource use. All simulations run for $S^*/\mathcal{R} = 0.8$, R = 200, $\mathcal{R}_0 = 40$, $\mathcal{S}^*/\mathcal{S} = 0.1$ unless otherwise noted.



Figure S3: Probability of coexistence as a function of niche saturation, with simulations using the "simultaneous assembly" approximation where mutant invasion is modeled using a single assembly process (SI Section 1.3). Red points show the simultaneous assembly approximation, which allows reappearance of extinct species. Black points show simulations where extinct species were disallowed from returning (as in the rest of our study); this change results in coexistence probability increasing at high saturation. Simulations were performed with $\mathcal{R} = 200$, $\mathcal{R}_0 = 40$, and $\mathcal{S}^*/\mathcal{S} = 0.1$ using knock-out mutations.



Figure S4: Coexistence probability for different strategy mutations. Left: Mutant-parent coexistence probability as a function of niche saturation S^*/\mathcal{R} for full knock-out mutations and "global" strategy mutations with $\gamma = 1$ (which have similarly-sized changes to all nonzero entries of $\vec{\alpha}$, as described in the Appendix). Right: Mutant-parent coexistence probability as a function of mutation effect size γ for $S^*/\mathcal{R} = 0.8$, for partial knock-out and global strategy mutations. Black curves show theoretical prediction. All simulations run with $\mathcal{R} = 100$, $\mathcal{R}_0 = 40$, and $S^*/\mathcal{S} = 0.1$ using Dirichlet-distributed resource usage vectors.



Figure S5: Scatterplot showing the invasion fitness of knock-out strategy mutations against the relative abundance of the parent before mutant invasion, with colors indicating whether the mutants coexist with or replace their parent at ecological equilibrium. The dashed line shows the theoretical prediction that should divide replacement events from coexistence events. Simulations run for $\mathcal{R} = 200$, $\mathcal{R}_0 = 40$, $\mathcal{S}^*/\mathcal{S} = 0.1$, and $\mathcal{S}^*/\mathcal{R} = 0.8$.



Figure S6: Coexistence probability of parents and knock-out mutants for generalizations of the community assembly procedure discussed in the main text. Left: Mutant-parent coexistence probability as a function of the number of resources \mathcal{R}_P used by the parent strain, while the background community uses $\mathcal{R}_0 = 40$ resources. Right: Mutant-parent coexistence probability as a function of scaled variance in resource supply σ_{κ}^2 . All simulations run with $\mathcal{R} = 200$, $\mathcal{R}_0 = 40$, $\mathcal{S}^*/\mathcal{S} = 0.1$, and $\mathcal{S}^*/\mathcal{R} = 0.8$.



Figure S7: Checks to ensure that we have consistently identified extinctions numerically. **(A)** Plot of the number of putatively identified survivors for an assembled community, as a function of the numerical extinction threshold. If the resource deficit for an organism is above this threshold times $Std(h_i)$, the organism is labeled extinct. The plateau over multiple orders of magnitude indicates a numerically stable definition of survival vs. extinction; the dashed line shows the threshold used throughout our analysis. **(B)** Scatterplot of numerical resource deficit of organisms before and after invasion of a knock-out strategy mutant, with color indicating whether the resource deficit after invasion passes the extinction threshold, (i.e., whether the species has gone extinct). The fact that most red and gray points are clearly separated shows that most species identified as extinct are insensitive to our precise choice of threshold. Simulations were run with $\mathcal{R} = 200$, $\mathcal{R}_0 = 40$, $\mathcal{S}^*/\mathcal{S} = 0.1$.

1 Community Assembly with Mutations

In this section, we outline the setup that allows us to derive our main theoretical results. First, we describe the consumer resource model we study, and how the task of finding an ecological equilibrium can be described as an optimization problem. Then, we define a convenient fitness gauge which brings our model in line with previous work. Finally, we outline an approximate procedure for introducing mutations into an assembled community, while considering only a single community assembly process.

1.1 Model Outline

As summarized in the Model section, we are interested in the following system: consider an ecosystem formed by combining S species, identically and independently drawn from some underlying distribution, competing for \mathcal{R} resources. We allow this ecosystem to come to ecological equilibrium, at which point $S^* \leq S$ species will survive at nonzero abundance. Then, suppose that one randomly selected surviving species undergoes a mutation to produce a new strain, which can differ from its parent in its ability to consume resources. If the mutant has positive invasion fitness, it will rise to nonzero abundance in the population, and the abundances of other species may change as well. We are interested in analyzing the properties of this new ecosystem with a single mutant, particularly the probability that the mutant and its parent strain are able to coexist at ecological equilibrium.

First, we define our community assembly process and consumer resource model, which generally follows previous work (28–32, 36, 45). We consider a well-mixed community formed from the combination of S species, indexed by $\mu \in \{1, ..., S\}$, and \mathcal{R} resources, indexed by $i \in \{1, ..., \mathcal{R}\}$. We generally consider the large-ecosystem limit, where $S, \mathcal{R} \to \infty$ while maintaining a constant ratio S/\mathcal{R} . Each organism is associated with an "uptake budget" X_{μ} and a "strategy vector" $\vec{\alpha}_{\mu}$, which contains \mathcal{R} non-negative entries that sum to 1. The uptake budget dictates the overall capacity of the organism to consume resources, setting its maximum growth rate under ideal growth conditions; the strategy vector describes how the organism allocates this uptake budget between substitutable resources. The consumer-resource dynamics for the abundance n_{μ} of each species and the concentration c_i of each resource are described by

$$\frac{\partial n_{\mu}}{\partial t} = n_{\mu} \left[\sum_{i=1}^{\mathcal{R}} \alpha_{\mu,i} e^{X_{\mu}} h_i(\vec{c}) - 1 \right],$$
(S1)

$$\frac{\partial c_i}{\partial t} = K_i - h_i(\vec{c}) \left[\sum_{\mu=1}^{\mathcal{S}} n_\mu \alpha_{\mu,i} e^{X_\mu} + 1 \right],$$
(S2)

where K_i is the supply of each resource, and $h_i(\vec{c})$ is the availability of resource *i* to cells, which depends on the concentration of each resource. Note that our model assumes a symmetry between consumption of resources and their contribution to biomass; this neglects the possibility of idiosyncratic effects like resource sequestration (47), where some organ-

isms remove resources from the environment without corresponding growth in abundance. We assume that resource dynamics are fast, and the total biomass of organisms large, so that the resource supply is nearly balanced out by their consumption by organisms. Then, the availability of each resource will be at a local fixed point set by the species abundances:

$$h_i = \frac{K_i}{\sum_{\mu} n_{\mu} \alpha_{\mu,i} e^{X_{\mu}}} \tag{S3}$$

$$=\frac{\kappa_i/\mathcal{R}}{\sum_i f_\mu \alpha_{\mu,i} e^{X_\mu}},\tag{S4}$$

normalizing by total biomass to transform strain abundances n_{μ} into relative abundances f_{μ} . We have defined rescaled resource supplies $\kappa_i \equiv K_i / (\sum_j K_j / \mathcal{R})$, which has units such that $\langle \kappa_i \rangle = 1$ and $\operatorname{Var}(\kappa_i) = \sigma_{\kappa}^2$ (in simulations, we choose $\sigma_{\kappa}^2 = 0$ or normally distributed κ_i). The simplified relative abundance dynamics are

$$\frac{\partial f_{\mu}}{\partial t} = f_{\mu} \left[\sum_{i=1}^{\mathcal{R}} \alpha_{\mu,i} e^{X_{\mu}} h_i(\vec{f}) - 1 \right].$$
(S5)

For any choice of species, these dynamics possess a Lyapunov function and a single stable equilibrium (30, 32). At this equilibrium, none of the surviving organisms can increase their abundance, while none of the extinct species could invade the ecosystem if introduced at low abundance. Thus, at ecological equilibrium, the bracketed term, or "resource surplus," must be non-positive for all μ , and is equal to zero if and only if species μ survives at nonzero abundance.

All that remains is to choose the resource consumption parameters of our competing species, which we draw from a common statistical distribution. For our theoretical analysis, we follow the procedure of Ref. (36), where the uptake budgets X_{μ} are drawn from a normal distribution with variance ϵ^2 , and the resource consumption strategies $\vec{\alpha}_{\mu}$ are normalized binary vectors where each entry has an independent probability $\mathcal{R}_0/\mathcal{R}$ of being nonzero.

1.2 Ecological Equilibrium as Optimization

Conveniently, the task of finding the unique stable equilibrium for our consumer resource model can be recast as a constrained optimization problem (36, 37). Specifically, we treat the resource availabilities h_i (rather than the relative abundances f_{μ}) as free variables to optimize over, and consider the task

optimize
$$\sum_{i} \kappa_{i} \log h_{i}$$

subject to $\sum_{i} \alpha_{\mu,i} h_{i} - e^{-X_{\mu}} \leq 0 \ \forall \mu.$ (S6)

The constraint is the same as that discussed above, requiring that the resource surplus be non-positive for each organism. Framed in this manner, we observe that the model

is symmetric under uptake budget translation, which we can view as a change in "pure fitness" decoupled from the environment. If we change variables into translated fitness $\tilde{X}_{\mu} = X_{\mu} + \chi$, and rescaled $\tilde{h}_i = e^{-\chi}h_i$, then the optimization problem becomes

optimize
$$\sum_{i} \kappa_{i} \left(\log \tilde{h}_{i} + \chi \right)$$

subject to $e^{\chi} \left(\sum_{i} \tilde{h}_{i} \alpha_{\mu i} - e^{-\tilde{X}_{\mu}} \right) \leq 0 \ \forall \mu.$ (S7)

Since χ is just an arbitrary constant, it is clear that these optimization problems are the same, up to the rescaling of the h_i . Suppose that h_i^* solves the first optimization problem. Then, $\tilde{h}_i^* = e^{-\chi} h_i^*$ will solve the gauge-shifted optimization problem

optimize
$$\sum_{i} \kappa_{i} \log \tilde{h}_{i}$$

subject to $\sum_{i} \tilde{h}_{i} \alpha_{\mu i} - e^{-\tilde{X}_{\mu}} \leq 0 \,\forall \mu.$ (S8)

To convince ourselves that these two problems really have the same solution, we return to the equation defining h_i in terms of the relative equilibrium abundances f_{μ} :

$$h_i = \frac{\kappa_i / \mathcal{R}}{\sum_i \alpha_{\mu i} e^{X_\mu} f_\mu}.$$
(S9)

Translating the uptake budgets by χ rescales h_i by $e^{-\chi}$, while leaving f_{μ} unchanged. Therefore, both optimization problems correspond to the same f_{μ} , which is the actual observable we are interested in.

This gauge symmetry allows us to relate our consumer resource model to that analyzed in Refs. (36) and (37), by choosing a gauge such that $\langle h_i \rangle = 1$. Let's define a constant h_0 based on the equilibrium solution in our (arbitrary) original gauge,

$$h_0 = \frac{1}{\mathcal{R}} \sum_i h_i^*,\tag{S10}$$

and consider what happens when we set $\chi = \log h_0$. The rescaled solution is

$$\tilde{h}_i^* = \frac{h_i^*}{h_0},\tag{S11}$$

which by definition has mean 1. It is now natural to define a new variable, $g_i = 1 - h_i$, which has mean zero. The optimization problem is now

optimize
$$\sum_{i} \kappa_{i} \log(1 - g_{i})$$

subject to $1 - e^{-\tilde{X}_{\mu}} - \sum_{i} g_{i} \alpha_{\mu i} \leq 0 \ \forall \mu.$ (S12)

For a large ecosystem, the g_i 's should be small, allowing us to Taylor expand the optimization function to second order. We also recall that for surviving species, the constraint should achieve equality (i.e., resource surplus should be zero). Since the g_i 's are small and have mean zero, the third term on the LHS of the constraint equation should be small. To achieve equality, the first and second terms on the LHS must therefore nearly cancel, implying that the \tilde{X}_{μ} are small for survivors. In other words, choosing a gauge such that all $h_i \approx 1$ is the same as choosing a gauge where the pure fitnesses of surviving organisms are near zero. This observation allows us to Taylor expand the \tilde{X}_{μ} as well. Altogether, the approximate optimization problem is

optimize
$$\sum_{i} \kappa_{i} \left(g_{i} + \frac{g_{i}^{2}}{2} \right)$$

subject to $\tilde{X}_{\mu} - \sum_{i} g_{i} \alpha_{\mu i} \leq 0 \,\forall \mu.$ (S13)

For organisms that go extinct, the Taylor expansion we did of the $e^{-\tilde{X}_{\mu}}$ may not be very accurate, since \tilde{X}_{μ} can be arbitrarily negative. However, since $1 - \tilde{X}_{\mu} \leq e^{-\tilde{X}_{\mu}}$, this approximation will not cause us to erroneously label any extinct species as alive – and for the species which are alive, the approximation should be sufficiently accurate to avoid worry. While it is not yet clear why this approximation is useful, we will see below that remapping the consumer resource problem in this way makes it amenable to replica-theoretic analysis. Note that for numerical simulations, we do not make these approximations, instead solving the original optimization problem in Eq. (S6).

1.3 Single-Ecosystem Approximation of First-Step Invasion

As discussed in the Model section, mutant-parent coexistence can be described as two correlated community assembly processes. Here, we argue that we can reasonably approximate the conditions for mutant-parent coexistence while considering only one community assembly process, which simplifies our theoretical work. First, we describe the two-ecosystem framework. An ecosystem E_1 is formed through standard community assembly: S species are placed into competition, and S^* remain alive at equilibrium. Then, one of the surviving species produces a beneficial mutant offspring. This effectively creates a new community assembly problem for ecosystem E_2 , where the initial S^* species plus one mutant are allowed to reach ecological equilibrium. The mutant-parent coexistence probability can therefore be written as

$$\mathbb{P}_{\text{coex}} = \mathbb{P}\left[\text{parent alive in } E_2 | \text{mutant alive in } E_2, \text{ parent alive in } E_1\right], \quad (S14)$$

where the conditional probability requires that (1) the mutant be beneficial and (2) the parent be alive when it produces a mutant. Standard manipulation of probabilities allows us to rewrite this as

$$\mathbb{P}_{\text{coex}} = \frac{\mathbb{P}\left[\text{P and M in } E_2\right] - \mathbb{P}\left[\text{P and M in } E_2, \text{ P not in } E_1\right]}{\mathbb{P}\left[\text{M in } E_2\right] - \mathbb{P}\left[\text{M in } E_2, \text{ P not in } E_1\right]}.$$
(S15)

First, we argue that $\mathbb{P}[P \text{ and } M \text{ in } E_2, P \text{ not in } E_1] \approx 0$. This probability describes a situation where the parent is unable to survive in the initial ecosystem, but its hypothetical mutant offspring is. Moreover, the invasion of the mutant offspring adjusts the ecosystem in a way where the original parent strain is also able to simultaneously invade. Intuitively, both of these requirements are unlikely: it will be difficult for a small mutation to allow an otherwise unfavored species to invade, and we would naively expect that the presence of the mutant would make it more difficult for the parent species to survive – not easier – due to their shared competition for resources. Making this approximation, we find

$$\mathbb{P}_{\text{coex}} \approx \frac{\mathbb{P}\left[\text{P and M in } E_2\right]}{\mathbb{P}\left[\text{M in } E_2\right] - \mathbb{P}\left[\text{M in } E_2, \text{ P not in } E_1 \text{ or } E_2\right]}.$$
(S16)

The next approximation we make is similar: we assume that if the mutant is able to survive, the parent would likely be able to survive in its absence. Mathematically, $\mathbb{P}[M \text{ in } E_2] \gg \mathbb{P}[M \text{ in } E_2, P \text{ not in } E_1 \text{ or } E_2]$. While it is possible that a small-effect beneficial mutation could push an organism over the survival threshold, we expect that *most* randomly chosen surviving organisms will not be able to be rescued from extinction by a single mutation. Thus, we write

$$\mathbb{P}_{\text{coex}} \approx \frac{\mathbb{P}\left[\text{P and M in } E_2\right]}{\mathbb{P}\left[\text{M in } E_2\right]}.$$
(S17)

Now, our coexistence probability depends only on one ecosystem. But there is still an issue: E_2 is not a traditionally assembled community, since it is formed only from the survivors of an earlier community assembly process plus a mutant, rather than independently sampled organisms. Let's instead consider E_s , an ecosystem obtained from simultaneous community assembly of S independently sampled species, plus one "mutant" species which is closely related to one of the S other species. The key difference between E_2 and E_s is that E_s can potentially contain species which were extinct in E_1 , but are brought "back to life" by the invasion of the mutant. While such returns are possible (37), we conjecture that the invasion of a closely-related mutant strain should be a relatively small perturbation to the ecosystem, and as such these re-introductions should be rare – particularly when the number of resources and species are large. So, as a first approximation, we do not expect them to meaningfully impact \mathbb{P}_{coex} . After these approximations, we now have

$$\mathbb{P}_{\text{coex}} \approx \frac{\mathbb{P}\left[\text{P and M in } E_s\right]}{\mathbb{P}\left[\text{M in } E_s\right]}.$$
(S18)

This approximation allows us to estimate the mutant-parent coexistence probability in an assembled community by only considering one community assembly process, which includes two closely correlated organisms. (Which one we have labeled the "parent" and "mutant" is essentially arbitrary after making these approximations.) As shown in Fig. S3, simulations confirm that these approximations are valid across most of the parameter regime we study, so our one-ecosystem approximation produces results which match the two-ecosystem one. An exception is at very high expected niche saturation, where species "coming back to life" after mutant invasion becomes more relevant. However, there is a narrow range of parameters where this effect is significant (particularly for ecosystems with many resources), and our approximation where these species are allowed to return nonetheless sets a lower bound on the mutant-parent coexistence probability.

2 Replica-Theoretic Analysis of First-Step Invasion

2.1 Partition Function and the Replica Trick

We would now like to argue that our model can be treated similarly to Refs. (36) and (37), meaning that their results describe a baseline assembled community before we add mutations. While we will not reproduce every element of their calculation, we restate key steps and approximations. Since ecological equilibrium can be described as the solution to an optimization problem, we can write a partition function for a system with energy $F(\vec{g}) = \sum_i F_i(g_i) = \sum_i \kappa_i \left(g_i + \frac{g_i^2}{2}\right)$ at temperature β^{-1} . In the zero-temperature limit $\beta \rightarrow \infty$, the system will be exactly at ecological equilibrium; at finite values of β , fluctuations from equilibrium correspond to shot noise in the population. The partition function is

$$Z = \int_{\Omega} \prod_{i} dg_{i} e^{-\beta F(\vec{g})}, \tag{S19}$$

where Ω corresponds to the constrained region in our optimization problem, where resource surplus is non-positive. We can write out this region explicitly using the Heaviside function $\theta(x) = \max(0, x)$ as follows:

$$Z = \int_{-\infty}^{1} \prod_{i} dg_{i} e^{-\beta F_{i}(g_{i})} \prod_{\mu=1}^{S} \theta\left(\sum_{i} g_{i} \alpha_{\mu i} - \tilde{X}_{\mu}\right)$$
(S20)

$$= \int_{-\infty}^{1} \prod_{i} dg_{i} e^{-\beta F_{i}(g_{i})} \prod_{\mu=1}^{S} \int d\Delta_{\mu} \theta(-\Delta_{\mu}) \delta\left(\Delta_{\mu} + \sum_{i} g_{i} \alpha_{\mu i} - \tilde{X}_{\mu}\right)$$
(S21)

$$= \int_{-\infty}^{1} \prod_{i} dg_{i} e^{-\beta F_{i}(g_{i})} \prod_{\mu} \int \frac{d\Delta_{\mu} d\hat{\Delta}_{\mu}}{2\pi} \theta(-\Delta_{\mu}) e^{i\sum_{\mu} \hat{\Delta}_{\mu} \left(\Delta_{\mu} + \sum_{i} g_{i} \alpha_{\mu i} - \tilde{X}_{\mu}\right)}.$$
 (S22)

where X_{μ} is the pure fitness of species μ , in the gauge described in 6.1.2. We have traded our complicated integration region for two additional integration variables for each species: the resource surplus Δ_{μ} and its auxiliary Fourier variable $\hat{\Delta}_{\mu}$.

This partition function is written for a particular choice of *S* sampled species. However, we are interested not in any particular choices of species, but the behavior of the ecosystem when *typical* species are drawn from some random distribution. These typical ecosystems can be analyzed by calculating $\langle \log Z \rangle$, where the average is over our random draws of species. This average can be calculated through use of the replica trick,

$$\langle \log Z \rangle = \lim_{n \to 0} \frac{\langle Z^n \rangle - 1}{n}.$$
 (S23)

In accordance with the replica trick, we consider *n* copies of our system, which share identical resident species but may differ in shot noise (encoded by the temperature β). Then, we treat *n* as a real number and consider the limit $n \rightarrow 0$, comparing our results to simulation to ensure this approximation holds. With *n* copies of our system, the average partition

function is

$$\begin{split} \langle Z^{n} \rangle &= \int \prod_{i,a} \left[dg_{i}^{a} e^{-\beta \sum_{i,a} F_{i}(g_{i}^{a})} \right] \times \\ &\prod_{\mu=1}^{S} \left\{ \prod_{a} \left[\int \frac{d\Delta_{\mu}^{a} d\hat{\Delta}_{\mu}^{a}}{2\pi} \theta(-\Delta_{\mu}^{a}) \right] e^{i \sum_{a} \hat{\Delta}_{\mu}^{a} \Delta_{\mu}^{a}} \left\langle e^{-i \tilde{X}_{\mu} \sum_{a} \hat{\Delta}_{\mu}^{a}} \right\rangle_{\tilde{X}_{\mu}} \times \\ &\prod_{i} \left\langle e^{i \sum_{i,a} \hat{\Delta}_{\mu}^{a} g_{i}^{a} \alpha_{\mu,i}} \right\rangle_{\alpha_{\mu,i}} \right\}, \end{split}$$
(S24)

where *a* runs from 1 to *n* and indexes the replicas of our system. Next, let's average over our random draws of species. Suppose that the pure fitnesses \tilde{X}_{μ} are drawn from a normal distribution with mean χ and variance ϵ^2 . (Recall that χ represents our fitness gauge, and will later be set to ensure that $\langle g_i \rangle = 0$.) The \tilde{X}_{μ} -dependent term then becomes

$$\left\langle e^{-i\tilde{X}_{\mu}\sum_{a}\hat{\Delta}_{\mu}^{a}}\right\rangle_{\tilde{X}_{\mu}} = \exp\left[-i\chi\sum_{a}\hat{\Delta}_{\mu}^{a} - \frac{\epsilon^{2}}{2}\left(\sum_{a}\hat{\Delta}_{\mu}^{a}\right)^{2}\right].$$
(S25)

Now, let's consider the average over the strategy vectors. For binary resource strategies, the $\alpha_{\mu i}$ are defined as

$$\alpha_{\mu i} = \frac{\sigma_{\mu i}}{\sum_{j} \sigma_{\mu j}},\tag{S26}$$

where $\sigma_{\mu i}$ are i.i.d. random variables equal to 1 with probability $\mathcal{R}_0/\mathcal{R}$ and 0 otherwise. Unlike the $\sigma_{\mu i}$, the $\alpha_{\mu i}$ are not independent for the same μ with different *i*'s. However, we can approximate them as independent by separating out the dependent portion, provided that $\mathcal{R}_0 \gg 1$:

$$\alpha_{\mu i} = \frac{\sigma_{\mu i}}{\mathcal{R}_0 + b_\mu \sqrt{\mathcal{R}_0}} \tag{S27}$$

$$\approx \frac{\sigma_{\mu i}}{\mathcal{R}_0} \left(1 - \frac{b_\mu}{\sqrt{\mathcal{R}_0}} \right). \tag{S28}$$

The b_{μ} are mean-zero $\mathcal{O}(1)$ random variables representing the fluctuations in the total number of resources used by each species. While not technically independent of the $\sigma_{\mu i}$, this dependence would contribute to terms with smaller powers of \mathcal{R}_0 and can thus be ignored.

So, we consider their averages separately:

$$\left\langle e^{i\sum_{i,a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}\alpha_{\mu,i}}\right\rangle_{\alpha_{\mu,i}} \approx \left\langle \prod_{i} \left[1 - \frac{\mathcal{R}_{0}}{\mathcal{R}} + \frac{\mathcal{R}_{0}}{\mathcal{R}} e^{i\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}/\mathcal{R}_{0}} e^{-i\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}b_{\mu}/\mathcal{R}_{0}^{3/2}} \right] \right\rangle_{b_{\mu}} \approx \left\langle \prod_{i} \exp\left[i\frac{1}{\mathcal{R}}\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}\left(1 - \frac{b_{\mu}}{\mathcal{R}_{0}}\right) - \frac{(1 - \mathcal{R}_{0}/\mathcal{R})}{2\mathcal{R}_{0}\mathcal{R}} \left(\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}\right)^{2} \right] \right\rangle_{b_{\mu}}$$
(S29)

$$= \left\langle e^{-i\sum_{i,a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}b_{\mu}/\mathcal{R}\sqrt{\mathcal{R}_{0}}}\right\rangle_{b_{\mu}}\prod_{i}\exp\left[i\frac{1}{\mathcal{R}}\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i} - \frac{(1-\mathcal{R}_{0}/\mathcal{R})}{2\mathcal{R}\mathcal{R}_{0}}\left(\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}\right)^{2}\right]$$
(S30)

$$\approx \exp\left[-\frac{\left(\sum_{i,a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}\right)^{2}}{2R^{2}\mathcal{R}_{0}}\right]\prod_{i}\exp\left[i\frac{1}{\mathcal{R}}\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}-\frac{\left(1-\mathcal{R}_{0}/\mathcal{R}\right)}{2\mathcal{R}\mathcal{R}_{0}}\left(\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}\right)^{2}\right]$$
(S31)

$$\approx \prod_{i} \exp\left[i\frac{1}{\mathcal{R}}\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i} - \frac{(1-\mathcal{R}_{0}/\mathcal{R})}{2\mathcal{R}\mathcal{R}_{0}}\left(\sum_{a}\hat{\Delta}^{a}_{\mu}g^{a}_{i}\right)^{2}\right].$$
(S32)

Effectively, this approximation treats $\alpha_{\mu i} \approx \sigma_{\mu_i}/\mathcal{R}_0$. Finally, to bring our answer in line with Refs. (36) and (37), we rescale variables $\epsilon \to \epsilon/\mathcal{R}_0$, $\chi \to \chi/\mathcal{R}_0$, $\Delta^a_\mu \to \Delta^a_\mu/\mathcal{R}_0$, and $\hat{\Delta}^a_\mu \to \mathcal{R}_0 \hat{\Delta}^a_\mu$. The final partition function is

$$\langle Z^n \rangle = \int \prod_{i,a} \left[dg_i^a e^{-\beta \sum_{i,a} F_i(g_i^a)} \right] \prod_{\mu=1}^S \left\{ \prod_a \left[\int \frac{d\Delta_\mu^a d\hat{\Delta}_\mu^a}{2\pi} \theta(-\Delta_\mu^a) \right] \times \exp\left[i \sum_a \hat{\Delta}_\mu^a \left(\Delta_\mu^a + \frac{\mathcal{R}_0}{\mathcal{R}} \sum_i g_i^a - \chi \right) - \frac{\epsilon^2}{2} \left(\sum_a \hat{\Delta}_\mu^a \right)^2 - \frac{\mathcal{R}_0(1 - \mathcal{R}_0/\mathcal{R})}{2\mathcal{R}} \sum_i \left(\sum_a \hat{\Delta}_\mu^a g_i^a \right)^2 \right] \right\}.$$
(S33)

This is the same as the partition function calculated in Refs. (36) and (37), except for the presence of χ . We can ignore χ until later in the calculation (for example, by absorbing it into the definition of Δ^a_{μ} , which causes it to appear only within the Heaviside function), and re-introduce it only when we require that $\langle g_i \rangle = 0$.

2.2 Introducing First-Step Mutations

The above partition function is valid for an ecosystem where all species are sampled randomly. However, in our "simultaneous assembly" approximation of first-step mutation, we would like to include two sampled species which are very related to each other, differing only by a single knock-out mutation. We will index the parent and mutant species P and M; they have the same resource consumption strategy, except that the parent uses resource 1, while the mutant's capacity to use resource 1 is lowered by a factor of γ . (For a full knock-out mutation which preserves our assumption about binary resource use, $\gamma = 1$, but this more general type of mutation will aid us when interpreting our conclusions later.) Their pure fitnesses are also the same, except that the mutant's pure fitness is increased by ΔX (which can be negative to indicate an overall cost to the mutation). With the parentmutant pair accounted for, the partition function becomes

$$\begin{split} \langle Z^{n} \rangle &= \int \prod_{i,a} \left[dg_{i}^{a} e^{-\beta \sum_{i,a} F_{i}(g_{i}^{a})} \right] \prod_{\mu=1}^{S-1} \left\{ \prod_{a} \left[\int \frac{d\Delta_{\mu}^{a} d\hat{\Delta}_{\mu}^{a}}{2\pi} \theta(-\Delta_{\mu}^{a}) \right] \times \\ \exp \left[i \sum_{a} \hat{\Delta}_{\mu}^{a} \left(\Delta_{\mu}^{a} + \frac{\mathcal{R}_{0}}{\mathcal{R}} \sum_{i} g_{i}^{a} - \chi \right) - \frac{\epsilon^{2}}{2} \left(\sum_{a} \hat{\Delta}_{\mu}^{a} \right)^{2} - \frac{\mathcal{R}_{0}(1 - \mathcal{R}_{0}/\mathcal{R})}{2\mathcal{R}} \sum_{i} \left(\sum_{a} \hat{\Delta}_{\mu}^{a} g_{i}^{a} \right)^{2} \right] \right\} \times \\ \prod_{a} \left[\int \frac{d\Delta_{P}^{a} d\hat{\Delta}_{P}^{a}}{2\pi} \frac{d\Delta_{P}^{a} d\hat{\Delta}_{P}^{a}}{2\pi} \theta(-\Delta_{P}^{a}) \theta(-\Delta_{M}^{a}) \right] \\ \exp \left\{ i \sum_{a} \hat{\Delta}_{P}^{a} \left(\Delta_{P}^{a} + \frac{\mathcal{R}_{0}}{\mathcal{R}} \sum_{i} g_{i}^{a} + \left(1 - \frac{\mathcal{R}_{0}}{\mathcal{R}} \right) g_{1}^{a} - \chi \right) + \right. \\ \left. i \sum_{a} \hat{\Delta}_{M}^{a} \left(\Delta_{M}^{a} + \frac{\mathcal{R}_{0}}{\mathcal{R}} \sum_{i} g_{i}^{a} + \left(1 - \gamma - \frac{\mathcal{R}_{0}}{\mathcal{R}} \right) g_{1}^{a} - \chi - \mathcal{R}_{0} \Delta X \right) - \\ \left. \frac{\epsilon^{2}}{2} \left[\sum_{a} \left(\hat{\Delta}_{P}^{a} + \hat{\Delta}_{M}^{a} \right) \right]^{2} - \frac{\mathcal{R}_{0}(1 - \mathcal{R}_{0}/\mathcal{R})}{2\mathcal{R}} \sum_{i} \left[\sum_{a} \left(\hat{\Delta}_{P}^{a} + \hat{\Delta}_{M}^{a} \right) g_{i}^{a} \right]^{2} \right\}. \end{split}$$
(S34)

(Note that the ΔX term has a factor of \mathcal{R}_0 to account for the rescaling we did earlier.) The next few steps of the calculation are a series of approximations which allow us to evaluate this complicated integral, paralleling those in Refs. (36) and (37).

2.3 Saddle Point Approximations

For $\mathcal{R} \gg 1$, the central limit theorem dictates that the mean and variance of the equilibrium g_i (which encode the availabilities of each resource) should approach a deterministic value. Introducing the mutant-parent pair is a relatively small change in the ecosystem, so we expect that these deterministic values should be the same as those calculated in previous work, if the mutant-parent pair were not present. This motivates us to define replica-specific "order parameters"

$$m^a = \sum_i g_i^a,\tag{S35}$$

$$q^{ab} = \sum_{i} g^a_i g^b_i. \tag{S36}$$

As we did with the resource surplus Δ^a_{μ} , we can introduce these order parameters as new integration variables, defined via delta functions in their Fourier representation. This results in an integral

$$\begin{split} \langle Z^n \rangle &= \int \prod_{a \leq b} \left(\frac{dq^{ab} d\hat{q}^{ab}}{2\pi} \right) \prod_a \left(\frac{dm^a d\hat{m}^a}{2\pi} \right) \exp\left[i \left(\sum_{a \leq b} q^{ab} \hat{q}^{ab} + \sum_a m^a \hat{m}^a \right) \right] \times \\ \prod_{i=2}^R \left\{ \int_{-\infty}^1 \prod_a dg_i^a \exp\left[-\sum_a \beta F_i(g_i^a) - i \sum_a \hat{m}^a g_i^a - i \sum_{a \leq b} \hat{q}^{ab} g_i^a g_i^b \right] \right\} \times \\ \sum_{\mu=1}^{S-1} \left\{ \prod_a \left[\int \frac{d\Delta_\mu^a d\hat{\Delta}_\mu^a}{2\pi} \theta(-\Delta_\mu^a) \right] \times \right. \\ \exp\left[i \sum_a \hat{\Delta}_\mu^a \left(\Delta_\mu^a + \frac{\mathcal{R}_0}{\mathcal{R}} m^a - \chi \right) - \frac{1}{2} \sum_{a,b} \left(\frac{\mathcal{R}_0(1 - \mathcal{R}_0/\mathcal{R})}{\mathcal{R}} q^{ab} + \epsilon^2 \right) \hat{\Delta}_\mu^a \hat{\Delta}_\mu^b \right] \right\} \\ \times \prod_a \left[\int \frac{d\Delta_\mu^a d\hat{\Delta}_P^a}{2\pi} \frac{d\Delta_M^a d\hat{\Delta}_M^a}{2\pi} \theta(-\Delta_P^a) \theta(-\Delta_M^a) \int_{-\infty}^1 dg_1^a \right] \exp\left\{ \sum_a \beta F_1(g_1^a) - i \sum_a \hat{m}^a g_1^a - i \sum_{a \leq b} \hat{q}^{ab} g_1^a g_1^b + i \sum_a \hat{\Delta}_P^a \left(\Delta_P^a + \frac{\mathcal{R}_0}{\mathcal{R}} m^a + \left(1 - \frac{\mathcal{R}_0}{\mathcal{R}} \right) g_1^a - \chi \right) \\ + i \sum_a \hat{\Delta}_M^a \left(\Delta_M^a + \frac{\mathcal{R}_0}{\mathcal{R}} m^a + \left(1 - \gamma - \frac{\mathcal{R}_0}{\mathcal{R}} \right) g_R^a - \chi - \mathcal{R}_0 \Delta X \right) \\ - \frac{1}{2} \sum_{a,b} \left[\frac{\mathcal{R}_0(1 - \mathcal{R}_0/\mathcal{R})}{\mathcal{R}} (q^{ab} - g_1^a g_1^b) + \epsilon^2 \right] \left(\hat{\Delta}_P^a \hat{\Delta}_P^b + 2\hat{\Delta}_P^a \hat{\Delta}_M^b + \hat{\Delta}_M^a \hat{\Delta}_M^b \right) \right\}. \end{split}$$

So far, we have made no approximations from our initial partition function. Now, however, we invoke the saddle-point approximation, stating that the probability distribution for the order parameters and their Fourier conjugates is so heavily peaked that their values are equivalent across replicas. In short, we assume

$$m^a = m, (S38)$$

$$q^{aa} = q^D \tag{S39}$$

$$q^{ab} = q^O \text{ if } a \neq b, \tag{S40}$$

and likewise for the conjugate (hatted) order parameters. We recall that we must eventually choose our fitness gauge χ such that m = 0, a deviation from the analysis in Refs. (36) and (37) due to our choice of consumer resource model. Our saddle point approximation allows us to significantly simplify our integral, by eliminating the integrals over the replica-specific order parameters. This change decouples our large integral expression into a product of three *independent* integrals: one over the resource availabilities g_i for i > 1, one over the surpluses Δ^a_{μ} for all species besides the parent and the mutant, and the final integral over g_1 and the surpluses for the mutant and parent strains.

Because we assume a diverse, many-species ecosystem, the species besides the parent and mutant will be most important in setting the saddle-point values of our order parameters and their Fourier conjugates. Thus, we approximate that these saddle-point values are unchanged from the calculation in Refs. (36) and (37), which includes only the first two sets of integrals (i.e., those independent of the parent and mutant). While we will not restate their calculation here, we will summarize their results. If we assume that the values of q^D and q^O are similar –

$$q^D \approx q^O = q,\tag{S41}$$

$$q^D - q^O = \frac{\mathcal{R}}{\beta}x,\tag{S42}$$

then the saddle-point values of the conjugate variables are

$$\hat{m} \approx i\beta,$$
 (S43)

$$\hat{q}^O \approx i \left(\frac{\beta}{x} \sqrt{\frac{q}{\mathcal{R}}}\right)^2,$$
 (S44)

$$\hat{q}^D - \frac{\hat{q}^O}{2} \approx i\beta \frac{x-1}{2x}.$$
(S45)

These equations leave q and x undetermined, which can be solved for implicitly and approximated through numerical equations which we will discuss later.

We also make the approximation $q^O - g_1^a g_1^b \approx q^O$. The order parameter q is formed by adding together $g_i^a g_i^b$ for each of \mathcal{R} resources, and we expect each of these quantities to be positive. So, the scale of $q - g_R^a g_R^b \sim q(\mathcal{R} - 1)/\mathcal{R} \sim q$. After making this approximation, let us also recall that $F_i(g_i^a) = \kappa_i \left[g_i^a + (g_i^a)^2/2\right]$. Knowing that m = 0 and $\hat{m} \approx i\beta$ causes all the terms linearly dependent on the g_i^a to cancel out, besides the mutation-dependent g_1^a terms: thus, before considering the effects of the mutation, our system is invariant under rotation of the $g_i^{a'}$'s. Practically, this means that our initial choice of a knock-out mutation was arbitrary: a knock-in mutation, or any strategy change $\Delta \vec{\alpha}$ with the same overall magnitude, can be redefined as a change in a single element through an appropriate rotation. Our results should therefore be valid for mutations which affect multiple resources simultaneously.

2.4 Joint Distribution of Δ_P and Δ_M

Now that we have decoupled our integrals through the saddle-point approximation, we are nearly ready to analyze the phenomenon of parent-mutant coexistence. To do this, we only care about the part of the partition function which depends on those strains. For simplicity, we will also at this point assume uniform resource supply ($\kappa_i = 1$); we will discuss at the end of our calculation what happens when this assumption is relaxed. With the saddle-point approximation for $\kappa_i = 1$, the portion of the integral in Eq. (S37) which

depends on the parent and mutant strains is

$$Z_{PM}^{n} = \prod_{a} \left[\int \frac{d\Delta_{P}^{a} d\hat{\Delta}_{P}^{a}}{2\pi} \frac{d\Delta_{M}^{a} d\hat{\Delta}_{M}^{a}}{2\pi} \theta(-\Delta_{P}^{a}) \theta(-\Delta_{M}^{a}) \int_{-\infty}^{1} dg_{1}^{a} \right] \times$$

$$\exp\left\{ \sum_{a} \left[-\frac{\beta}{2x} (g_{1}^{a})^{2} \right] + \frac{1}{2} \left(\frac{\beta}{x} \sqrt{\frac{q}{\mathcal{R}}} \sum_{a} g_{1}^{a} \right)^{2} + i \sum_{a} \hat{\Delta}_{P}^{a} \left(\Delta_{P}^{a} + \frac{\mathcal{R}_{0}}{\mathcal{R}} m + \left(1 - \frac{\mathcal{R}_{0}}{\mathcal{R}} \right) g_{1}^{a} - \chi \right) + i \sum_{a} \hat{\Delta}_{M}^{a} \left(\Delta_{M}^{a} + \frac{\mathcal{R}_{0}}{\mathcal{R}} m + \left(1 - \gamma - \frac{\mathcal{R}_{0}}{\mathcal{R}} \right) g_{1}^{a} - \chi - \mathcal{R}_{0} \Delta X \right) - \frac{\mathcal{R}_{0} (1 - \mathcal{R}_{0} / \mathcal{R})}{2\beta} x \sum_{a} \left(\hat{\Delta}_{P}^{a} + \hat{\Delta}_{M}^{a} \right)^{2} - \frac{1}{2} \sum_{a,b} \left[\frac{\mathcal{R}_{0} (1 - \mathcal{R}_{0} / \mathcal{R})}{\mathcal{R}} q + \epsilon^{2} \right] \left(\hat{\Delta}_{P}^{a} \hat{\Delta}_{P}^{b} + 2 \hat{\Delta}_{P}^{a} \hat{\Delta}_{M}^{b} + \hat{\Delta}_{M}^{a} \hat{\Delta}_{M}^{b} \right) \right\}.$$
(S46)

Next, we perform a change of variables to $\hat{\Delta}^a_{\pm} = \hat{\Delta}^a_P \pm \hat{\Delta}^a_M$:

$$Z_{PM}^{n} = \prod_{a} \left[\int \frac{d\Delta_{P}^{a} d\Delta_{M}^{a}}{4\pi} \frac{d\hat{\Delta}_{+}^{a} d\hat{\Delta}_{-}^{a}}{4\pi} \theta(-\Delta_{P}^{a}) \theta(-\Delta_{M}^{a}) \int_{-\infty}^{1} dg_{1}^{a} \right] \times \\ \exp\left\{ \sum_{a} \left[-\frac{\beta}{2x} (g_{1}^{a})^{2} \right] + \frac{1}{2} \left(\frac{\beta}{x} \sqrt{\frac{q}{\mathcal{R}}} \sum_{a} g_{1}^{a} \right)^{2} \\ + i \sum_{a} \hat{\Delta}_{+}^{a} \left(\frac{\Delta_{P}^{a} + \Delta_{M}^{a}}{2} + \frac{\mathcal{R}_{0}}{\mathcal{R}} m + \left(1 - \frac{\gamma}{2} - \frac{\mathcal{R}_{0}}{\mathcal{R}} \right) g_{1}^{a} - \chi - \mathcal{R}_{0} \frac{\Delta X}{2} \right) \\ + i \sum_{a} \hat{\Delta}_{-}^{a} \left(\frac{\Delta_{P}^{a} - \Delta_{M}^{a}}{2} + \frac{\gamma}{2} g_{1}^{a} + \mathcal{R}_{0} \frac{\Delta X}{2} \right) - \frac{\mathcal{R}_{0} (1 - \mathcal{R}_{0} / \mathcal{R})}{2\beta} x \sum_{a} \left(\hat{\Delta}_{+}^{a} \right)^{2} \\ - \frac{1}{2} \sum_{a,b} \left[\frac{\mathcal{R}_{0} (1 - \mathcal{R}_{0} / \mathcal{R})}{\mathcal{R}} q + \epsilon^{2} \right] \left(\sum_{a} \hat{\Delta}_{+}^{a} \right)^{2} \right\}.$$
(S47)

We now aim to decouple the replicas, by eliminating terms which involve more than one replica. All these terms are of the form $(\sum_a f^a)^2$, where f^a is some quantity which depends only on one replica. We can decouple these replicas using the identity

$$\exp\left(\frac{1}{2}(Cx)^2\right) = \int d\omega \frac{e^{-\omega^2/2}}{\sqrt{2\pi}} e^{Cx\omega}.$$
(S48)

Thus, we can decouple the replicas at the cost of introducing an additional Gaussian integration variable per square term. We can now drop the a indices and replace them with a

simple power to *n*:

$$Z_{PM}^{n} = \int d\omega_{1} d\omega_{2} \Biggl\{ \int \frac{d\Delta_{P} d\Delta_{M}}{4\pi} \frac{d\hat{\Delta}_{+} d\hat{\Delta}_{-}}{4\pi} \theta(-\Delta_{P}) \theta(-\Delta_{M}) \int_{-\infty}^{1} dg_{1} \\ \times \exp\left[-\frac{\beta}{2x} g_{1}^{2} + \omega_{1} \frac{\beta}{x} \sqrt{\frac{q}{\mathcal{R}}} g_{1} - \frac{\mathcal{R}_{0}(1-\mathcal{R}_{0}/\mathcal{R})}{2\beta} x \hat{\Delta}_{+}^{2} \\ + i\hat{\Delta}_{+} \left(\frac{\Delta_{P} + \Delta_{M}}{2} + \frac{\mathcal{R}_{0}}{\mathcal{R}} m + \left(1 - \frac{\gamma}{2} - \frac{\mathcal{R}_{0}}{\mathcal{R}} \right) g_{1} + \omega_{2} V_{\text{tot}}^{1/2} - \chi - \mathcal{R}_{0} \frac{\Delta X}{2} \right) \\ + i\hat{\Delta}_{-} \left(\frac{\Delta_{P} - \Delta_{M}}{2} + \frac{\gamma}{2} g_{1} + \mathcal{R}_{0} \frac{\Delta X}{2} \right) \Biggr] \Biggr\}^{n} \frac{e^{-(\omega_{1}^{2} + \omega_{2}^{2})/2}}{2\pi}.$$
(S49)

Here, we have defined $V_{\text{tot}} = q\mathcal{R}_0(1 - \mathcal{R}_0/\mathcal{R})/\mathcal{R} + \epsilon^2$. This parameter, defined as ψ^2 in Refs. Refs. (36) and (37), represents the total variance in fitness among sampled species from both strategy and pure fitness. As we will demonstrate later, the integration variables ω_1 and ω_2 do in fact have physical meaning: ω_1 is related to the invasion fitness of the mutant, while ω_2 is related to the abundance of the parent. Next, we perform the integral over g_1 . To do so, we assume that most of the weight of the integral is concentrated near small g, meaning that we can approximate the integration bounds as $-\infty$ to ∞ rather than $-\infty$ to 1. We also condense all of the resulting terms which do not depend on $\hat{\Delta}_+$ or $\hat{\Delta}_-$ into a constant term c.

We can write the result as

$$Z_{PM}^{n} = \int d\omega_{1} d\omega_{2} \Biggl\{ \int \frac{d\Delta_{P} d\Delta_{M}}{4\pi} \frac{d\hat{\Delta}_{+} d\hat{\Delta}_{-}}{4\pi} \theta(-\Delta_{P}) \theta(-\Delta_{M}) \\ \times \exp\left[-\frac{1}{2}\vec{\Delta}\mathbf{M}\vec{\Delta}^{t} + i\vec{\Delta}\cdot\vec{v} + c\right] \Biggr\}^{n} \frac{e^{-(\omega_{1}^{2} + \omega_{2}^{2})/2}}{2\pi},$$
(S50)

where

$$\vec{\Delta} = \begin{bmatrix} \hat{\Delta}_+ & \hat{\Delta}_- \end{bmatrix},\tag{S51}$$

$$\mathbf{M} = \frac{x}{\beta} \begin{bmatrix} A & B \\ B & C \end{bmatrix},\tag{S52}$$

$$\vec{v} = \begin{bmatrix} \frac{\Delta_P + \Delta_M}{2} + D & \frac{\Delta_P - \Delta_M}{2} + E \end{bmatrix},$$
(S53)

and we define

$$A = \mathcal{R}_0 \left(1 - \frac{\mathcal{R}_0}{\mathcal{R}} \right) + \left(1 - \frac{\gamma}{2} - \frac{\mathcal{R}_0}{\mathcal{R}} \right)^2, \tag{S54}$$

$$B = \frac{\gamma}{2} \left(1 - \frac{\gamma}{2} - \frac{\mathcal{R}_0}{\mathcal{R}} \right), \tag{S55}$$

$$C = \frac{\gamma^2}{4},\tag{S56}$$

$$D = \omega_2 V_{\text{tot}}^{1/2} + \frac{\mathcal{R}_0}{\mathcal{R}} m + \omega_1 \sqrt{\frac{q}{\mathcal{R}}} \left(1 - \frac{\gamma}{2} - \frac{\mathcal{R}_0}{\mathcal{R}}\right) - \chi - \mathcal{R}_0 \frac{\Delta X}{2}, \tag{S57}$$

$$E = \mathcal{R}_0 \frac{\Delta X}{2} + \omega_1 \frac{\gamma}{2} \sqrt{\frac{q}{\mathcal{R}}}.$$
(S58)

We can now perform the integrals over $\hat{\Delta}_+$ and $\hat{\Delta}_-$. This yields

$$Z_{PM}^{n} = \int d\omega_{1} d\omega_{2} \left[\frac{1}{\sqrt{\det \mathbf{M}}} \int \frac{d\Delta_{P} d\Delta_{M}}{4\pi} \theta(-\Delta_{P}) \theta(-\Delta_{M}) \times \exp\left(-\frac{1}{2} \vec{v} \mathbf{M}^{-1} \vec{v}^{t} + c\right) \right]^{n} \frac{e^{-(\omega_{1}^{2} + \omega_{2}^{2})/2}}{2\pi}.$$
(S59)

Now, let's utilize the replica trick limit $n \to 0$. Again following the methodology in Refs. (36) and (37), this can be used to recover the joint probability distribution $\rho(\Delta_P, \Delta_M)$:

$$\lim_{n \to 0} Z_{PM}^n = \int_{-\infty}^0 d\Delta_P d\Delta_M \int d\omega_1 d\omega_2 \frac{e^{-(\omega_1^2 + \omega_2^2)/2}}{2\pi} \times \frac{\exp\left(-\frac{1}{2}\vec{v}\mathbf{M}^{-1}\vec{v}^t\right)}{\int_{-\infty}^0 d\Delta'_P d\Delta'_M \exp\left(-\frac{1}{2}\vec{v}'\mathbf{M}^{-1}\vec{v}'^t\right)},$$
(S60)

$$= \int_{-\infty}^{0} d\Delta_P d\Delta_M \rho(\Delta_P, \Delta_M).$$
(S61)

Recall that $\mathbf{M}^{-1} \sim \beta$, an inverse temperature which scales with the number of cells in our community, reflecting the shot noise in our population at any given time. If we assume that our population is always exactly at equilibrium (i.e., is large enough that shot noise is negligible), we can take the $\beta \rightarrow \infty$ limit, in which case the exponentials involving \mathbf{M}^{-1} will be very strongly peaked. In this limit, we can therefore approximate the ratio in Eq. (S60) as a delta function:

$$\rho(\Delta_P, \Delta_M) = \int d\omega_1 d\omega_2 \frac{e^{-(\omega_1^2 + \omega_2^2)/2}}{2\pi} \delta(\Delta_P - \Delta_P^*) \delta(\Delta_M - \Delta_M^*),$$
(S62)

where $\Delta_P^*(\omega_1, \omega_2)$ and $\Delta_M^*(\omega_1, \omega_2)$ are the non-positive values of Δ_P and Δ_M that minimize $\vec{v} \mathbf{M}^{-1} \vec{v}^t$ for a given ω_1 and ω_2 .

2.5 Parent-Mutant Coexistence Probability

The parent-mutant coexistence probability is encoded in the joint distribution $\rho(\Delta_P, \Delta_M)$:

$$\mathbb{P}[\text{Both survive}] = \int d\omega_1 d\omega_2 \frac{e^{-(\omega_1^2 + \omega_2^2)/2}}{2\pi} \delta(\Delta_P^*(\omega_1, \omega_2)) \delta(\Delta_M^*(\omega_1, \omega_2)).$$
(S63)

So, the next step is to uncover the functional forms of Δ_P^* and Δ_M^* , which represent the values of Δ_P and Δ_M that minimize $\vec{v}\mathbf{M}^{-1}\vec{v}^t$, constrained such that both Δ_P and Δ_M are non-positive. If the global minimum happens to be within the allowed region, these quantities are simple:

$$\Delta_P^*|_{\Delta_P^* < 0, \ \Delta_M^* < 0} = -D - E, \tag{S64}$$

$$\Delta_M^* \big|_{\Delta_P^* < 0, \ \Delta_M^* < 0} = -D + E.$$
(S65)

This unconstrained minimum will be in the allowed region if both of these quantities are negative, which occurs when D > |E|. If both resource surpluses are negative, neither the parent or mutant survive at equilibrium, so

$$\mathbb{P}[Both \operatorname{die}] = \mathbb{P}[D > |E|].$$
(S66)

What if the mutant survives and the parent goes extinct, such that $\Delta_M^* = 0$? Then Δ_P^* will be found at

$$\Delta_P^* \big|_{\Delta_P^* < 0, \ \Delta_M^* = 0} = \frac{2B(D+E) - 2(CD+AE)}{A - 2B + C},\tag{S67}$$

$$\approx -2E + \frac{2}{A}(B - C)(D - E).$$
(S68)

This expansion comes from the fact that $A \sim \mathcal{R}_0$ (provided that \mathcal{R}_0 does not get too close to \mathcal{R}), while $B, C \sim 1$. Similarly, if the mutant goes extinct while the parent survives, we find

$$\Delta_M^* \big|_{\Delta_P^* = 0, \ \Delta_M^* < 0} = \frac{2E(A+B) - 2D(B+C)}{A+2B+C},\tag{S69}$$

$$\approx 2E - \frac{2}{A}(B+C)(D+E).$$
(S70)

Parent-mutant coexistence requires that *none* of these optima are in the allowed region, so that $\Delta_P^* = \Delta_M^* = 0$. This is satisfied when

$$\mathbb{P}[Both survive] \approx \mathbb{P}[BE + CD < -|AE - BD - CE|, \ D < |E|].$$
(S71)

Recalling that $A \sim \mathcal{R}_0 \gg 1$, the *AE* term dominates unless $|E| \ll 1$, meaning that *E* must be small for the inequality to be possible. (Note that the $\mathcal{R}_0 \Delta X$ term is not actually $\mathcal{O}(\mathcal{R}_0)$, because the natural fitness scale for a mutant turns out to be $\Delta X \sim \mathcal{R}_0^{-3/2}$.) When this is the case, the *BE* and *CE* terms will be even smaller, allowing us to safely ignore them:

$$\mathbb{P}[Both survive] \approx \mathbb{P}[CD < -|AE - BD|, \ D < |E|],$$
(S72)

$$= \mathbb{P}[CD < -|AE - BD|]. \tag{S73}$$

We have eliminated the second condition because it is now redundant with the first: since C > 0, the first inequality requires D < 0, trivially satisfying D < |E|. Finally, let's recall that in our simultaneous assembly approximation of mutant invasion (Appendix 1.3), we must condition on the probability that the mutant survives:

$$\mathbb{P}_{\text{coex}} \approx \frac{\mathbb{P}[\text{Both survive}]}{\mathbb{P}[\text{Mutant survives}]},$$
(S74)

$$\approx \frac{\mathbb{P}[CD < -|AE - BD|]}{\mathbb{P}[E > D, E > 0]}.$$
(S75)

In principle, this is a quantity that can be calculated numerically. However, this result is difficult to intuitively interpret in this form, and it still depends on the saddle-point order parameter q. Nonetheless, we can immediately make two meaningful conclusions. If we send the number of resources consumed by a typical organism $\mathcal{R}_0 \to \infty$, then A becomes large and $\mathbb{P}_{\text{coex}} \to 0$. Likewise, if we send the effect size of our mutation $\gamma \to 0$, then $\frac{|CD|}{|AE-BD|} \sim \gamma \to 0$, and thus $\mathbb{P}_{\text{coex}} \to 0$. Since we are considering a mutation whose effect size is comparable to changing one resource's uptake rate by a factor of γ , both of these observations suggest the same thing: if the phenotypic effect of a mutation is too small, mutant-parent coexistence is impossible. This phenomenon is a signature of "mesoscopic" mutation, suggesting that our results may not be captured by models of mutations with infinitesimal phenotypic effects, such as adaptive dynamics approaches (44, 62).

3 Scaling Analysis of First-Step Mutations

In this section, we aim to restore physical intuition to our replica-theoretic results, finding approximate expressions which indicate how the frequency of inter-niche evolution scales with community parameters such as the number of surviving species.

3.1 Sampling Depth and Fitness Gauge

We must utilize the saddle-point results for an assembled community without mutation, largely from Refs. (36) and (37), to calculate our order parameters m and q. However, we opt for a different set of independent variables, defining our assembly process in terms of the number of sampled species S and the typical number of surviving species S^* . The community assembly parameter ϵ , which describes the variation in pure fitness among sampled species, will then be defined implicitly as a function of these parameters. This change of variables is helpful because S^* is a measurable property of an experimental ecosystem, while ϵ describes a property of *sampled* species, which may be very different from typical surviving species and is thus more difficult to access experimentally.

With this set of variables, a natural ratio to consider is the "sampling permissivity" S^*/S , roughly the probability that a new independently sampled species would be able to invade the ecosystem. It turns out that it will be useful to define an effective "sampling depth" λ ,

which is a function of S^*/S :

$$\lambda \equiv \operatorname{erf}^{-1}\left(1 - 2\frac{\mathcal{S}^*}{\mathcal{S}}\right)\sqrt{2}.$$
(S76)

For an undersampled system ($S^*/S \to 1$), the sampling depth $\lambda \to -\infty$; for an oversampled system ($S^*/S \to 0$), we have $\lambda \to \infty$. From the saddle-point results of Refs. (36) and (37) (which we do not re-derive here), we know that

$$m = \frac{\mathcal{R}}{\mathcal{R}_0} \left(V_{\text{tot}}^{1/2} \lambda + \chi \right).$$
(S77)

Recall that χ sets our choice of fitness gauge: the uptake budget of sampled species is drawn with mean χ/\mathcal{R}_0 and standard deviation ϵ/\mathcal{R}_0 . But in our community assembly approximations, we have chosen our gauge such that $\langle g_i \rangle = m/\mathcal{R} = 0$. This implies that

$$\chi = -V_{\rm tot}^{1/2}\lambda.\tag{S78}$$

As we sample more species and λ increases, the uptake budgets or pure fitnesses of surviving species will be further and further into the high-fitness tail of the distribution. Since the typical pure fitness of surviving species is nearly zero in our fitness gauge, χ must become more negative as λ increases to compensate for the increased sampling. In the consumer resource model analyzed in Refs. (36) and (37) (where $\chi = 0$), an increase in sampling depth results in an overall increase in m: as oversampled species become higher and higher in pure fitness, the typical resource becomes less valuable to consume. This effect causes resource knock-out mutations to be more beneficial on average than knock-in mutations in the oversampled limit. In our model, we adjust the "zero" of pure fitness to compensate for this effect, such that m = 0 regardless of λ .

For future sections, it is helpful to define a function $I(\lambda)$ such that

$$I(\lambda) = \frac{1+\lambda^2}{2} \operatorname{erfc}\left(\frac{\lambda}{\sqrt{2}}\right) - \frac{\lambda}{\sqrt{2\pi}} e^{-\lambda^2/2}.$$
(S79)

3.2 Niche Saturation and σ_{inv}

From saddle-point results in Refs. (36) and (37), we can write expressions for V_{tot} , q, and ϵ in terms of quantities we have already defined:

$$V_{\text{tot}}^{1/2} = \left(1 - \frac{\mathcal{R}_0}{\mathcal{R}}\right) \left(\frac{2\left(1 - \frac{S}{\mathcal{R}}I(\lambda)\right)}{-\frac{S}{\mathcal{R}}I'(\lambda)} - \lambda\right),\tag{S80}$$

$$q = V_{\text{tot}} \frac{S}{\mathcal{R}} I(\lambda) \frac{\mathcal{R}}{\mathcal{R}_0 (1 - \mathcal{R}_0 / \mathcal{R})} + \mathcal{R} \sigma_\kappa^2 \left(1 - \frac{\mathcal{S}^*}{\mathcal{R}} \right)^2,$$
(S81)

$$\epsilon^{2} = V_{\text{tot}} \left(1 - \frac{S}{\mathcal{R}} I(\lambda) \right) - \mathcal{R}_{0} \sigma_{\kappa}^{2} \left(1 - \frac{\mathcal{R}_{0}}{\mathcal{R}} \right) \left(1 - \frac{\mathcal{S}^{*}}{\mathcal{R}} \right)^{2}.$$
 (S82)

The order parameter q describes the total variance in resource availability, allowing us to find the distribution of the resource availabilities h_i : a Gaussian with mean 1 and variance q/\mathcal{R} . A mutation changing use of a single resource by a factor γ will have an invasion fitness with magnitude $\gamma |1 - h_i|/\mathcal{R}_0$. This leads us to define the quantity

$$\sigma_{\rm inv} \equiv \frac{\gamma}{\mathcal{R}_0} \sqrt{\frac{q}{\mathcal{R}}},\tag{S83}$$

which describes the typical magnitude of the invasion fitness of a strategy mutant $\vec{\alpha} \rightarrow \vec{\alpha} + \Delta \vec{\alpha}$, where $|\Delta \vec{\alpha}| \approx \gamma / \mathcal{R}_0$. (For example, for a complete knock-out mutant for a system with binary resource use, we have $\gamma = 1$.)

While these equations completely determine all the variables we have defined, they are difficult to interpret physically, having complicated dependence on the ratios S/\mathcal{R} and $\mathcal{R}_0/\mathcal{R}$ as well as the sampling depth λ . We can make progress by considering how these equations depend on the *niche saturation* $S^*/\mathcal{R} \leq 1$, describing how close to maximum capacity we expect our final ecosystem to be. By rewriting $S/\mathcal{R} = (S/S^*)(S^*/\mathcal{R})$, we separate out the parts of these equations that depend on the sampling depth λ (or equivalently, S^*/S) from the niche saturation S^*/\mathcal{R} . (Technically, rather than using the value of S^* for any particular ecosystem as an input parameter, we use its expectation $\langle S^* \rangle$ over random ecosystems with the same underlying parameters. Since these values tend to be close in practice, as shown in Figure 2B, we will simply write S^* in our scaling relationships.)

To simplify our results, it is helpful to expand our expressions in the undersampled and oversampled limits $(\lambda \to \pm \infty)$. It is worth noting that the $\lambda \to -\infty$ limit for fixed S^*/\mathcal{R} does not correspond to a physical community assembly process: the expression for ϵ in Eqn. (S82) must be positive, but $(S/\mathcal{R}) \cdot I(\lambda) \to \infty$ as $\lambda \to -\infty$, making the RHS of that equation negative. Effectively, this means that in order to sample species such that a desired fraction of niches are filled, a minimum fraction of sampled species must go extinct in the final ecosystem (on average). However, many quantities we are interested in are still well-defined in the $\lambda \to -\infty$ limit, and indeed scale similarly to the $\lambda \to \infty$ limit: we use this to demonstrate an overall lack of dependence on sampling depth.

We first quote results for $\sigma_{\kappa}^2 = 0$. From standard expansions of the error function, we obtain

$$\sigma_{\rm inv} \approx \begin{cases} \gamma \left(1 - \frac{S^*}{\mathcal{R}}\right) \sqrt{\frac{\mathcal{R} - \mathcal{R}_0}{S^* \mathcal{R}_0^3}} & \text{when } \lambda \to -\infty, \\ \gamma \sqrt{2} \left(1 - \frac{S^*}{\mathcal{R}}\right) \sqrt{\frac{\mathcal{R} - \mathcal{R}_0}{S^* \mathcal{R}_0^3}} & \text{when } \lambda \to \infty. \end{cases}$$
(S84)

Remarkably, the typical invasion fitness scale of a strategy mutation has very weak dependence on the sampling depth λ , changing only by a factor of $\sqrt{2}$ from the extremely undersampled to the extremely oversampled regimes. This result allows us to effectively drop out λ (and thus S^*/S) as a variable, which was very difficult to see from our saddle-point results above.

3.3 Scaling Analysis of \mathbb{P}_{coex}

We got our expression for σ_{inv} "for free" from the saddle-point value of q, without need to explicitly consider the impact of the parent and mutant on the ecosystem. The same is not true for the mutant-parent coexistence probability: to obtain an interpretable expression for \mathbb{P}_{coex} , we must start from our results in Appendix 2.5, and then undergo a similar procedure of plugging in saddle-point values in the $\lambda \to \pm \infty$ limits. Recall that in our two-ecosystem approximation, we have

$$\mathbb{P}_{\text{coex}} \approx \frac{\mathbb{P}[\text{Both survive}]}{\mathbb{P}[\text{Mutant survives}]}.$$
(S85)

To get some intuition, let's start with the simpler calculation of the denominator of this expression: the probability that the mutant species survives in our assembled ecosystem. From our earlier results, this is

$$\mathbb{P}_{M \text{ lives}} = \mathbb{P}\left[\Delta X + \mathcal{Z}_{1}\sigma_{\text{inv}} > 0, \\ \mathcal{Z}_{2}V_{\text{tot}}^{1/2} + V_{\text{tot}}^{1/2}\lambda + \mathcal{Z}_{1}\frac{\sigma_{\text{inv}}\mathcal{R}_{0}}{\gamma}\left(1 - \gamma - \frac{\mathcal{R}_{0}}{\mathcal{R}}\right) - \mathcal{R}_{0}\Delta X < 0\right],$$
(S86)

where Z_i are independent standard normal random variables. Looking at these inequalities, we can get some intuition for what they mean. The first inequality represents the probability the mutant is beneficial relative to the parent: the chances that the deterministic fitness change ΔX combined with the fitness change due to the strategy mutation $Z_1\sigma_{inv}$, is positive. The second inequality represents the requirement that the mutant is well-adapted enough to survive in the population in the first place: the V_{tot} factors represent total fitness variation across sampled species, while Z_2 represents the randomness in how well-adapted the parent organism is. Indeed, we expect the Gaussian distribution of Z_2 to map directly onto the truncated Gaussian distribution of the parent's relative abundance f_P :

$$f_P \approx \frac{2}{\mathcal{S}I'(\lambda)} (\mathcal{Z}_2 + \lambda)\theta(-\mathcal{Z}_2 - \lambda).$$
 (S87)

The terms which do not scale with $V_{\text{tot}}^{1/2}$ are corrections related to the fitness effect of the mutation. But for a small mutation, we expect these to produce a relatively small change in how well-adapted the organism is. Furthermore, the only circumstance where these corrections are likely to matter is one where the parent would not be able to survive in the population on its own, but the mutant can: a pathological situation for our approximation of \mathbb{P}_{coex} that we have argued is rare. So, it is fair to approximate

$$\mathbb{P}_{\mathrm{M \ lives}} \approx \mathbb{P}\left[\mathcal{Z}_{1} > -\frac{\Delta X}{\sigma_{\mathrm{inv}}}, \ \mathcal{Z}_{2} < -\lambda\right],\tag{S88}$$

$$= \frac{1}{2} \operatorname{erfc}\left(-\frac{\Delta X}{\sigma_{\rm inv}\sqrt{2}}\right) \cdot \frac{1}{2} \operatorname{erfc}\left(\frac{\lambda}{\sqrt{2}}\right),\tag{S89}$$

$$=\frac{\mathcal{S}^*}{2S}\mathrm{erfc}\left(-\frac{\Delta X}{\sigma_{\mathrm{inv}}\sqrt{2}}\right).$$
(S90)

This result should not be too surprising: if the pure fitness benefit ΔX carried by the mutant is zero, then the chance the mutant lives is $(1/2) \cdot (S^*/S)$: there are even odds the mutant is beneficial relative to the parent, and a probability S^*/S that it is one of the S^* well-adapted species which can survive in the community. If the mutant carries a change in pure fitness along with a change in strategy, its survival probability rises or falls accordingly as the deterministic pure fitness change $|\Delta X|$ approaches the same size as the stochastic fitness change $\sim \sigma_{inv}$ due to altered strategy.

Now, let's calculate the numerator of \mathbb{P}_{coex} , the probability that the mutant and parent will both be alive in our simultaneously assembled ecosystem. Again approximating that terms which scale with overall fitness variation $V_{tot}^{1/2}$ dominate terms which scale with mutant fitness change ΔX and σ_{inv} , we find that

$$\mathbb{P}_{\mathrm{P,M \ live}} \approx \mathbb{P}\left[\frac{\gamma V_{\mathrm{tot}}^{1/2}}{\mathcal{R}_0^2}(\mathcal{Z}_2 + \lambda) < \Delta X + \mathcal{Z}_1 \sigma_{\mathrm{inv}} < \frac{\gamma(1 - \gamma - \frac{\mathcal{R}_0}{\mathcal{R}})V_{\mathrm{tot}}^{1/2}}{\mathcal{R}_0^2(1 - \frac{\mathcal{R}_0}{\mathcal{R}})}(\mathcal{Z}_2 + \lambda)\right].$$
(S91)

Can we interpret this probability physically? The outer inequality only holds when $\mathcal{Z}_2 < -\lambda$. We calculated this probability earlier: it should be roughly $\mathcal{S}^*/\mathcal{S}$, the chance that the parent species is well-adapted enough to survive before mutant invasion. The middle term is the leading contribution to the invasion fitness of the mutant, which is $\mathcal{O}(\mathcal{R}_0^{-3/2})$. However, the inequality bounds it between two quantities of size $\mathcal{O}(\mathcal{R}_0^{-2})$. So, we conclude that parent-mutant coexistence is achieved when the invasion fitness of the mutant is sufficiently small.

Since the allowed region for Z_1 is smaller than its typical width by a factor of $1/\sqrt{R_0}$, we can approximate its probability density as constant over the allowed region. This gives us

$$\mathbb{P}_{P,M \text{ live}} \approx \frac{e^{-(\Delta X/\sigma_{inv})^2/2}}{\sqrt{2\pi}} \frac{\gamma^2 V_{tot}^{1/2}}{\sigma_{inv} \mathcal{R}_0^2 (1 - \frac{\mathcal{R}_0}{\mathcal{R}})} \int_{-\infty}^{-\lambda} (-\mathcal{Z}_2 - \lambda) \frac{e^{-\mathcal{Z}_2^2/2}}{\sqrt{2\pi}} d\mathcal{Z}_2$$
(S92)

$$=\frac{e^{-(\Delta X/\sigma_{\rm inv})^2/2}}{\sqrt{2\pi}}\frac{\gamma^2 V_{\rm tot}^{1/2}}{\sigma_{\rm inv} \mathcal{R}_0^2 (1-\frac{\mathcal{R}_0}{\mathcal{R}})}\frac{-I'(\lambda)}{2}.$$
(S93)

Finally, we divide by the probability the mutant lives, and consider the undersampled and oversampled limits when $\Delta X = 0$:

$$\mathbb{P}_{\text{coex}} \approx \begin{cases} \gamma \sqrt{\frac{2\mathcal{R}}{\pi \mathcal{S}^*}} \sqrt{\frac{1}{\mathcal{R}_0(1-\mathcal{R}_0/\mathcal{R})}} & \text{when } \lambda \to -\infty, \\ \gamma \sqrt{\frac{\mathcal{R}}{\pi \mathcal{S}^*}} \sqrt{\frac{1}{\mathcal{R}_0(1-\mathcal{R}_0/\mathcal{R})}} & \text{when } \lambda \to \infty. \end{cases}$$
(S94)

Once again, the dependence on the sampling depth is nearly negligible, contributing only an O(1) numerical factor. Note, however, that this result is different from that in the main text in Equation (8). This is because in our theoretical calculation, we arbitrarily chose the identity of the parent species and the mutation, effectively giving each surviving species and each mutation an equal chance of being selected. In reality, each surviving species produces mutations at a rate proportional to its abundance, and mutations have a probability proportional to their invasion fitness of surviving genetic drift. In the next section, we will discuss how to correct for this difference.

3.4 Dependence of \mathbb{P}_{coex} on Invasion Fitness and Abundance

Let us reframe our expression for \mathbb{P}_{coex} in terms of the abundance f_P of the parent and the invasion fitness s_{inv} of the mutant. We can read off the invasion fitness of a beneficial mutant as the saddle-point value of Δ_M^* when Δ_P is fixed at zero, representing the initial growth rate of the mutant *before* it has a chance to invade:

$$s_{\rm inv} \approx \Delta X + \mathcal{Z}_1 \sigma_{\rm inv} + \frac{\gamma}{\mathcal{R}_0^2} V_{\rm tot}^{1/2} (-\mathcal{Z}_2 - \lambda).$$
 (S95)

We can plug this expression into our earlier probability that both the mutant and parent live to find that

$$\mathbb{P}_{\mathrm{P,M \ live}} \approx \mathbb{P}\left[0 < s_{\mathrm{inv}} < \frac{\gamma^2 V_{\mathrm{tot}}^{1/2}}{\mathcal{R}_0^2 (1 - \mathcal{R}_0 / \mathcal{R})} (-\mathcal{Z}_2 - \lambda)\right].$$
(S96)

Recall that $-Z_2 - \lambda$ is a random variable describing how well-adapted the parent and mutant strains are to the ecosystem; it is proportional to the abundance of the parent, as we calculated earlier. While Z_2 is not independent of s_{inv} , its contribution to s_{inv} is small, so approximating them as independent is reasonable. Plugging in this relationship, we find that

$$\mathbb{P}_{\text{coex}} \approx \mathbb{P}\left[s_{\text{inv}} < -I'(\lambda) \frac{\gamma^2 V_{\text{tot}}^{1/2} \mathcal{S}}{2\mathcal{R}_0^2 (1 - \mathcal{R}_0 / \mathcal{R})} f_P\right],\tag{S97}$$

$$= \mathbb{P}\left[s_{\text{inv}} < \frac{\gamma^2 (1 - \mathcal{S}^* / \mathcal{R})}{\mathcal{R}_0^2 (\mathcal{S}^* / \mathcal{R})} \mathcal{S}^* f_P\right].$$
(S98)

So, our analysis of mutant-parent coexistence has an easily-interpreted meaning: a mutant strain will coexist with its parent when its (positive) invasion fitness is below

$$s_{\text{coex}}(f_P) = \frac{\gamma^2 (1 - \mathcal{S}^* / \mathcal{R})}{\mathcal{R}_0^2 (\mathcal{S}^* / \mathcal{R})} \mathcal{S}^* f_P.$$
(S99)

We can calculate this probability by treating s_{inv} and f_P as independent random variables, accounting for the probability the mutant survives genetic drift. In this case, their probability densities scale as

$$\rho(s_{\rm inv}) \sim s_{\rm inv} \theta(s_{\rm inv}) e^{-(s_{\rm inv} - \Delta X)^2 / 2\sigma_{\rm inv}^2}, \tag{S100}$$

$$\rho(f_P) \sim f_P \theta(f_P) e^{-(f_P + D\lambda)^2/2D^2},$$
(S101)

where $D \equiv -2/SI'(\lambda)$. Finally, \mathbb{P}_{coex} can be calculated using standard integration methods, giving the results quoted in the main text after applying similar scaling analysis to the previous section. We can also fix either s_{inv} or f_P to calculate a conditional coexistence probability. For example, plugging in a "typical" $f_P = 1/S^*$ indicates that $\bar{s}_{coex} = \frac{\gamma^2(1-S^*/\mathcal{R})}{\mathcal{R}_0^2(S^*/\mathcal{R})}$ is the typical value of s_{inv} above which the coexistence probability begins to fall.

3.5 Extensions of Coexistence Scaling

In this section, we will consider somewhat more general formulations of our model: nonuniform resource supply, and a mutant whose background parent species uses a different number of resources than the typical organism in the ecosystem. We will find that our qualitative results remain sound, and we can largely explain these results in terms of the "invasion fitness versus abundance" framework we have discussed so far.

3.5.1 Non-Uniform Resource Supply

First, let's consider non-uniform resource supply (nonzero σ_{κ}^2), as in (37). If each resource has a supply $\kappa_i = 1 + \delta \kappa_i$, then the availability h_i of that resource will have a component which scales with $\delta \kappa_i$ in addition to a community-dependent component. Thus, mutations that knock out a highly-supplied resource will be less beneficial on average. We can determine the strength of this effect by starting with the integrals over the g_i^a in the second line of Eq. (S37), where the resource supply enters through $F_i(g_i) \equiv \sum_i \kappa_i (g_i + g_i^2/2)$. This portion of the partition function was labeled A_i by Ref. (37) (see their Appendix 2, section 1.3). We can follow their calculation to obtain

$$A_{i} = \int \frac{d\omega}{2\pi} e^{-\omega^{2}/2} \left\{ \int_{-\infty}^{1} dg_{i} \exp\beta \left[-\left(\frac{1}{2x} + \frac{\delta\kappa_{i}}{2}\right) g_{i}^{2} - \left(\frac{\omega b}{\sqrt{R}} + \delta\kappa_{i}\right) g_{i} \right] \right\}^{n}, \quad (S102)$$

where *b* is a term independent of $\delta \kappa_i$ whose value does not concern us here. The contents of the brackets are a Gaussian integral over the g_i , centered where the integrand attains its maximum value. Taking non-uniform resource supply, $\delta \kappa_i \neq 0$, will shift this saddle-point value of g_i by approximately $x\delta\kappa_i \ll 1$. The saddle-point value of *x* was calculated in Ref. (36) as $x = 1 - S^*/R$, following our typical substitution of $(S^*/R)(S/S^*)$ for S/R. This shift in the central value of g_i will shift the invasion fitness of a knock-out mutation targeting that resource by a corresponding amount, since the invasion fitness of a knockout mutation is simply g_i/R_0 . Thus, we can write

$$s_{\rm inv} = \Delta X - \frac{\delta \kappa_i}{\mathcal{R}_0} \left(1 - \frac{\mathcal{S}^*}{\mathcal{R}} \right) + \mathcal{Z}\sigma_{\rm inv}(\delta \vec{\kappa} = 0).$$
(S103)

for a mutation knocking out resource *i*. The first term represents the mutation's change in pure fitness, the second term reflects that environmental supply makes some mutations more or less beneficial on average, while the stochastic third term depends on the specific draws of species in the community, as we calculated earlier. We can quantify the relative importance of environment and community by considering the invasion fitness of a mutation in a monoculture community, where invasion fitness *only* depends on the environment:

$$s_{\rm inv}^{\rm mono} \approx \Delta X - \frac{\delta \kappa_i}{\mathcal{R}_0}.$$
 (S104)

The correlation between the invasion fitness of a mutant in monoculture and in the community context illustrates how much the impact of environmental supply is felt through the "shielding" effects of the community: in the oversampled limit $(\lambda \to \infty)$, we find that

$$\rho(s_{\rm inv}^{\rm comm}, s_{\rm inv}^{\rm mono}) \approx \left(1 + \frac{\mathcal{R} - \mathcal{R}_0}{\mathcal{S}^* \mathcal{R}_0 \sigma_\kappa^2}\right)^{-1/2},\tag{S105}$$

assuming that $\delta \kappa_i$ are drawn from a distribution with mean zero and variance σ_{κ}^2 . In general, non-uniform resource supply will increase the variance in invasion fitness σ_{inv} :

$$\sigma_{\rm inv} \approx \begin{cases} \gamma \left(1 - \frac{S^*}{\mathcal{R}}\right) \sqrt{\frac{\mathcal{R} - \mathcal{R}_0}{S^* \mathcal{R}_0^3} + \frac{\sigma_{\kappa}^2}{\mathcal{R}_0^2}} & \text{when } \lambda \to -\infty, \\ \gamma \sqrt{2} \left(1 - \frac{S^*}{\mathcal{R}}\right) \sqrt{\frac{\mathcal{R} - \mathcal{R}_0}{S^* \mathcal{R}_0^3} + \frac{\sigma_{\kappa}^2}{2\mathcal{R}_0^2}} & \text{when } \lambda \to \infty. \end{cases}$$
(S106)

Note that σ_{inv} still scales with an overall factor of $1-S^*/\mathcal{R}$, leaving our previous qualitative results unaffected. Increasing resource supply variation increases σ_{inv} and thus lowers the coexistence probability, by making it less likely that the invasion fitness of an arbitrary beneficial mutation will be small enough to make coexistence possible.

3.5.2 Variation in Number of Metabolized Resources

Now, let's consider what happens if the parent species uses \mathcal{R}_P resources on average, rather than the \mathcal{R}_0 resources that most species in the community use. This extension helps us disentangle what terms in our expressions depend on the typical metabolic overlap between members of the community (encoded by \mathcal{R}_0), and what depends on the specific number of resources used by the parent and mutant species. It can also help determine whether generalist or specialist species are more likely to diversify in a given community.

The more resources a given organism uses, the more its growth rate and overall fitness are averaged out between different independent variables, effectively reducing the variance in its abundance as well as the invasion fitness of any particular mutation. So, if \mathcal{R}_P is larger, the parent species is less likely to have particularly high abundance, but the invasion fitness of a strategy mutation it obtains is likely to be smaller in magnitude. The first effect should lower the coexistence probability, while the second should raise it. Performing the calculation up to our expression for the joint probability of Δ_P and Δ_M gives us

$$A \approx \mathcal{R}_0 \frac{\mathcal{R}_0}{\mathcal{R}_P} \left(1 - \frac{\mathcal{R}_P}{\mathcal{R}} \right), \tag{S107}$$

$$B \approx \frac{\mathcal{R}_0}{\mathcal{R}_P} \frac{\gamma}{2} \left[\frac{\mathcal{R}_0}{\mathcal{R}_P} \left(1 - \frac{\gamma}{2} \right) - \frac{\mathcal{R}_0}{\mathcal{R}} \right], \tag{S108}$$

$$C \approx \frac{\mathcal{R}_0^2}{\mathcal{R}_P^2} \frac{\gamma^2}{4},\tag{S109}$$

$$D \approx \omega_2 V_P^{1/2} + \lambda V_{\text{tot}}^{1/2}, \tag{S110}$$

$$E \approx \mathcal{R}_0 \frac{\Delta X}{2} + \omega_1 \frac{\mathcal{R}_0}{\mathcal{R}_P} \frac{\gamma}{2} \sqrt{\frac{q}{\mathcal{R}}}.$$
(S111)

Here, we have defined $V_P = \frac{\mathcal{R}_0^2(1-\mathcal{R}_P/\mathcal{R})}{\mathcal{R}\mathcal{R}_P}q + \epsilon^2$, reflecting the different variation in overall fitness between the parent and other species in the community. We have once again defined γ such that $\gamma \approx 1$ for a knock-out mutation.

Since most of the community is unaffected by changing \mathcal{R}_P , we use the same saddle-point results we calculated before. Thus, the value of σ_{inv} is unchanged, although the invasion fitness of the mutant has a prefactor of $\mathcal{R}_0/\mathcal{R}_P$ to account for the altered number of resources. The inequality we obtain that predicts mutant-parent coexistence (for $\Delta X = 0$) is now given by

$$\mathbb{P}_{\mathrm{P,M \ live}} \approx \mathbb{P}\left[\mathcal{Z}_1 < \frac{\gamma V_P^{1/2}}{\mathcal{R}_0(1 - \mathcal{R}_P/R)\sqrt{q/R}} \left(-\mathcal{Z}_2 - \lambda \frac{V_{\mathrm{tot}}^{1/2}}{V_P^{1/2}}\right)\right],\tag{S112}$$

where Z_1 is once again proportional to the invasion fitness of the mutant, and the Z_2 dependent factor on the RHS is proportional to the abundance of the parent strain. The calculations otherwise proceed as before, yielding the results shown in Figure S6. For these parameter values, it appears that mutant-parent coexistence is more likely for mutations on a background strain which metabolizes many resources, since these mutations tend to have small invasion fitness.

4 Numerical Simulations

Throughout this work, we have performed numerical simulations of community assembly in order to verify and extend our analytical results. These simulations were performed by randomly sampling species and solving the optimization problem corresponding to ecological equilibrium (described in SI Section 1.2), rather than explicitly solving the abundance dynamics: the final outputs are the resource availabilities h_i and the surpluses Δ_{μ} . The code for these simulations is based on that in Ref. (37).

Much of our analysis relies on identifying whether individual species are present in the population or extinct. Mathematically, these cases correspond to whether Δ_{μ} is zero or negative. Numerically, however, Δ_{μ} is never precisely zero, necessitating the definition of an "extinction threshold." Any species with a surplus Δ_{μ} below the extinction threshold, a small negative number, is classified as extinct. We typically defined the extinction threshold as $-10^{-3} \cdot \text{Std}(h_i)/\mathcal{R}_0$, several orders of magnitude smaller than the typical fitness effect of a knock-in or knock-out mutation.

In Figure S7, we demonstrate that this is a reasonable choice of threshold. In order to be numerically sensible, there should be a clear separation between typical values of Δ_{μ} for surviving and extinct species, and the threshold should be well within this gap. We verified this by plotting the number of putative surviving species as a function of the extinction threshold. In the neighborhood of the threshold we have chosen, the number of putative survivors is independent of the threshold, suggesting that there is a clear separation between alive and extinct species and that our choice of threshold should not heavily impact our results. We also analyzed the values Δ_{μ} for each strain before and after mutant

invasion in order to ensure that we were reliably identifying species that go extinct. A scatterplot of initial vs. final Δ_{μ} values shows that almost all species are clearly separated into a "surviving" cluster and an "extinct" cluster, again suggesting that our results are not due to numerical artifacts.

In addition to our standard simulations, we also verified the robustness of our conclusions using organisms with Dirichlet-distributed resource use (Figs. S2 and S4). In these simulations, each entry of the strategy vectors $\vec{\alpha}_{\mu}$ for each organism is drawn from the gamma distribution $\Gamma(\mathcal{R}_0/\mathcal{R}, 1)$ and then normalized to sum to 1. This procedure results in Dirichlet-distributed entries which roughly retain the sparseness properties of binary resource use: typically, $\sim \mathcal{R}_0$ entries of each vector will be $\mathcal{O}(\mathcal{R}_0)$, while the others will be much smaller. Knock-out mutations were simulated by choosing a random entry of the parent organism's strategy vector with value at least γ/\mathcal{R}_0 , and reducing it by γ/\mathcal{R}_0 . We also used alternate "global-effect" mutations: rather than knocking out one resource, these mutations slightly perturbed each element of the resource use vector. Specifically, a strategy mutation $\Delta \vec{\alpha}$ was constructed by multiplying each element of $\vec{\alpha}$ by an independent $\sum_i \Delta \alpha_i = 0$ and $|\Delta \vec{\alpha}| = \gamma/\mathcal{R}_0$.