Modeling microbial metabolic trade-offs in a chemostat

Zhiyuan Li\textsuperscript{1,2,3}, Bo Liu\textsuperscript{4}, Sophia Hsin-Jung Li\textsuperscript{5}, Christopher G. King\textsuperscript{6}, Zemer Gitai\textsuperscript{5}, and Ned S. Wingreen\textsuperscript{5,7}

1. Center for Quantitative Biology, Peking University, Beijing, China.
2. Center for the Physics of Biological Function, Princeton University.
3. Princeton Center for Theoretical Science, Princeton University.
4. Yuanpei College, Peking University, Beijing, China.
5. Department of Molecular Biology, Princeton University.
6. Department of Physics, Princeton University.
7. Lewis-Sigler Institute for Integrative Genomics, Princeton University, Princeton, NJ, USA
ABSTRACT
Microbes face intense competition in the natural world, and so need to wisely allocate their resources to multiple functions, in particular to metabolism. Understanding competition among metabolic strategies that are subject to trade-offs is therefore crucial for deeper insight into the competition, cooperation, and community assembly of microorganisms. In this work, we evaluate competing metabolic strategies within an ecological context by considering not only how the environment influences cell growth, but also how microbes shape their chemical environment. Utilizing chemostat-based resource-competition models, we exhibit a set of intuitive and general procedures for assessing metabolic strategies. Using this framework, we are able to relate and unify multiple metabolic models, and to demonstrate how the fitness landscape of strategies becomes intrinsically dynamic due to species-environment feedback. Such dynamic fitness landscapes produce rich behaviors, and prove to be crucial for ecological and evolutionary stable coexistence in all the models we examined.
INTRODUCTION

The way microbes respond to and shape their local environment influences their community structure (Callahan, Fukami, & Fisher, 2014). Such microbe-environment interactions depend on the allocation strategies of cells, i.e., how a cell allocates its internal resources into various cellular functions, such as transport, assimilation, reproduction, motility, maintenance, etc. (Bachmann, Bruggeman, Molenaar, dos Santos, & Teusink, 2016). Within a microbial cell, energy and biomass are limited, and trade-offs always exist in allocating these valuable internal resources into the various functions required for cell growth. Therefore, the growth rate of cells cannot increase without bound. Rather, evolution acts on cells’ internal resource allocation to optimize growth and survival (S. Goyal, Yuan, Chen, Rabinowitz, & Wingreen, 2010; Liebermeister et al., 2014). To this end, in response to environmental changes, microbes rapidly adjust their metabolic strategies. For example, the yeast *Saccharomyces cerevisiae* switches from fermentation to respiration upon glucose depletion (Zaman, Lippman, Zhao, & Broach, 2008), and *Escherichia coli* exhibits drastic differences in ribosome content between different nutrient conditions (Li et al., 2018; Scott, Gunderson, Mateescu, Zhang, & Hwa, 2010). Moreover, in laboratory long-term evolution studies of microbes, adaptative mutations consistently emerge that reshape metabolism (Bachmann, Molenaar, dos Santos, & Teusink, 2017; Bajic & Sanchez, 2019; Blount, Barrick, Davidson, & Lenski, 2012; Long & Antoniewicz, 2018). Such short-term and long-term adjustments of metabolic strategies presumably confer fitness benefits, and it is important to map metabolic strategies onto these benefits to better understand the regulation and evolution of microbial metabolism.

The convergence towards steady state makes the chemostat an ideal experimental system to culture microorganisms and investigate their metabolic status (Wides & Milo, 2018; Ziv, Brandt, & Gresham, 2013). In a chemostat, fresh nutrients are supplied at a constant rate, while medium with cells is removed at the same rate to maintain constant volume. The metabolite concentrations in the chemostat constitute the chemical environment directly perceived by cells, and determine their growth rates. Importantly, cells also shape this environment through their consumption and secretion of metabolites. One advantage of a chemostat is the automatic convergence of cellular growth rates towards the controlled dilution rate. This convergence occurs through negative feedback between microbes and their environment: the higher the population, the worse the chemical environment, and the slower the growth rate. As a result (provided the nutrient supply allows for faster-than-dilution growth to prevent “washout”), the cells in the chemostat will reach the steady-state population that sustains growth at the dilution rate (De Leenheer, Levin, Sontag, & Klausmeier, 2006; Smith & Waltman, 1995). This stabilization of the cellular growth rate at the controlled dilution rate facilitates precise characterization of cellular physiology in a constant environment. However, it also imposes challenges in quantitatively understanding the advantages and disadvantages of various metabolic strategies: if all metabolic strategies lead to identical growth rates in a chemostat, how should we evaluate whether one strategy is “better” or “worse” than another? If strategies can be compared, are there “best” strategies, and how do these optima shift as the experimental conditions such as nutrient-supply concentrations and dilution rates change?

If we evaluate metabolic strategies by the outcome of competition between species, many insights can be gained from theoretical ecology. For example, resource-competition models have
provided a simple context to explore competition dynamics in chemostat-like ecosystems such as lakes and rivers (Smith & Waltman, 1995). In such models, species interact only indirectly via consumption (and sometimes production) of a common pool of nutrients. A steady state can be reached if the species present can shape the nutrient concentration to support a growth rate equal to their dilution or death rate (Tilman, 1982). Resource-competition models underpin many ecosystem theories including contemporary niche theory as pioneered by MacArthur (MacArthur, 1970), popularized by Tilman (Tilman, 1980, 1982), and extended by Chase and Leibold (Chase & Leibold, 2003). A central component of contemporary niche theory is a graphical approach, generally consisting of three components: zero net growth isoclines (ZNGIs) in chemical space, an impact vector representing a species’ nutrient consumption, and a supply point to describe the external resource supply (Koffel, Daufresne, Massol, & Klausmeier, 2016). This graphical approach is a powerful and intuitive way of assessing the outcome of competition, yet it is not yet commonly utilized in understanding microbial metabolic strategies with trade-offs.

Resource-competition models focusing on various aspects of cellular metabolism vary in their assumptions regarding species-environment interactions, and can lead to diverse results for community structure and population dynamics. In a model where species compete for essential resources, different nutrient requirements can produce intrinsically oscillatory or even chaotic dynamics (Huisman & Weissing, 1999, 2001). Alternatively, cross-feeding (Goldford et al., 2018; Pfeiffer & Bonhoeffer, 2004) can promote stable coexistence, while preferential nutrient utilization (A. Goyal, Dubinkina, & Maslov, 2018) can lead to multistability. With metabolic trade-offs, a model in which growth rate is additive in imported nutrients self-organizes to a state of unlimited stable coexistence (Posfai, Taillefumier, & Wingreen, 2017), while another model with convertible essential nutrients also allows evolutionarily stable coexistence but with a limited number of species (Taillefumier, Posfai, Meir, & Wingreen, 2017). This large variety of models and the richness of possible behaviors raises the question of unification: is there a simple framework that consolidates this diverse group of models into one easily understandable picture?

Continuity of the strategy space adds another layer of complexity in characterizing the “best” metabolic strategy or strategies. With infinite possibilities for allocating cellular resources, how should one pinpoint the optimal ones? Adaptive dynamics in ecological theory, also known as evolutionary invasion analysis, provides valuable guidance (Metz, 2012). This mathematical framework addresses the long-term evolution of traits in asexually reproducing populations by quantifying the fitness of each “trait” as a function of population composition. In this framework, “invasion fitness” is defined as the net-growth rate of a new variant when it is introduced into the resident population in an infinitesimally small amount, and a population allowing only non-positive invasion fitness for any new variant is considered to be “evolutionarily stable” (Doebeli, 2002; Ispolatov, Madhok, & Doebeli, 2016; Rueffler, Van Dooren, & Metz, 2004). Such an evolutionarily stable point is valuable for defining optimal strategies, as a community adopting the most suitable metabolic strategies should not be invasible by any other strategies. As microbes in nature frequently experience environmental fluctuations, it is important to understand whether and how such “optimal metabolic strategies” change with external conditions. Nevertheless, in the standard modeling framework of adaptive dynamics, the environment is implicit, and species directly act on each other without the realistic constraints imposed by competition for resources. Combining the concept of invasion fitness with explicit
competition for resources, under the assumption of metabolic trade-offs, could therefore bring new insights into microbial metabolic strategies and community assemblies.

In this work, we present a mathematical framework for analyzing competition for resources among various metabolic strategies in a chemostat setting. We combine and extend the graphical tools from resource-competition theory and the invasion-fitness approach from adaptive dynamics to relate and unify multiple models for microbial metabolic trade-offs. This combination provides an intuitive scheme to evaluate strategies under various external conditions. The center of the framework is the role of species in creating their own environment. Firstly, the chemical environment shaped by an endogenous species through growth and consumption can be inviting or prohibiting to an invader species, depending on the geometric relationship between the “zero net growth surface” of the invader and the environment created by the endogenous species. This geometry-dependent fitness leads to a general “rule of invasion” to compare pairs of strategies. In evaluating a continuum of strategies, the ensemble of their invasion fitnesses form a landscape, whose shape depends on the chemical environment. This intrinsically dynamic fitness landscape allows for the intransitivity of fitness (Soliveres et al., 2015). We demonstrate how such intransitivity can lead to rich ecosystem dynamics, including mutual invasion, multistability, and oscillations, and how all of these behaviors can be simply related via a graphical representation and the dynamic fitness landscape. Moreover, from the environment-dependent fitness landscape, we can define non-invasible/optimal metabolic strategies – namely, one or more strategies that construct a fitness landscape that places themselves on the top. The mathematical framework we present confers several advantages. Firstly, it separates the effect of changing dilution rate and of varying supply concentrations, facilitating quantitative interpretation as well as predictions for chemostat experiments. Secondly, it establishes an intuitive mapping from various metabolic models to population dynamics. Additionally, it reveals long-term implications, particularly in clarifying the general conditions for coexistence on both ecological and evolutionary time scales.
Results

Metabolic trade-offs and metabolic strategies
As discussed above, microorganisms need to allocate their limited internal resources into different cellular functions. In our models, we use $\alpha_j$ to denote the fraction of internal resources allocated to the $j$-th metabolic function, with $\vec{\alpha} = (\alpha_1, \alpha_2 \ldots)$ representing a metabolic strategy. An exact metabolic trade-off is assumed, such that $\sum_j \alpha_j = 1$. All possible values of $\vec{\alpha}$ define a continuous spectrum of metabolic strategies, which we name the strategy space. One major goal of our work is to construct a general and intuitive framework for evaluating strategies for a broad range of different metabolic models and experimental conditions.

Graphical representation of species creating their own chemical environment
One way to evaluate metabolic strategies is by comparing the competitiveness of “species” with fixed strategies in chemostat-type resource-competition models. In an idealized model of a chemostat (Fig 1A), $p$ types of nutrients are supplied at rate $d$ and concentrations $\vec{c}_{\text{supply}} = (c_1, \text{supply}, c_2, \text{supply}, \ldots, c_p, \text{supply})$, meanwhile cells and medium are diluted at the same rate $d$. The chosen values of $d$ and $\vec{c}_{\text{supply}}$ constitute the “external condition” for a chemostat. Nevertheless, the chemical environment that directly impacts cells is the metabolite concentration inside the chemostat, $\vec{c} = (c_1, c_2, \ldots, c_p)$, which influences the intracellular metabolite concentrations $\vec{q}$ and the growth rate $g_{\sigma}$ of the species $\sigma$. Accordingly, the biomass density $m_{\sigma}$ of species $\sigma$ adopting strategy $\vec{\alpha}_{\sigma}$ in the chemostat obeys:

$$\frac{dm_{\sigma}}{dt} = m_{\sigma} \cdot (g(\vec{c}, \vec{q}_{\sigma}, \vec{\alpha}_{\sigma}) - d). \quad (1)$$

The concentration $c_i$ of the $i$-th nutrient is a variable, influenced by its rate of consumption $l_i$ per cell volume. In a chemostat occupied by a single species $\sigma$, the changing rate of $c_i$ satisfies

$$\frac{dc_i}{dt} = d \cdot (c_{i, \text{supply}} - c_i) - m_{\sigma}/r \cdot l_i(\vec{c}, \vec{q}_{\sigma}, \vec{\alpha}_{\sigma}), \quad (2)$$

where $r$ is a constant representing the biomass per cell volume. (If the volume of the chemostat is $V_{\text{chemostat}}$ and the total volume of cells is $V_{\text{cells}}$, the import flux of the $i$-th nutrient $V_{\text{cells}} \cdot l_i$ implies a rate of change of concentration inside cells of $l_i$ and a corresponding rate of change of the concentration in the chemostat of $V_{\text{cells}} / V_{\text{chemostat}} \cdot l_i = (m \cdot V_{\text{chemostat}}/r) / V_{\text{chemostat}} \cdot l_i = m/r \cdot l_i$). A negative value of $l_i$ corresponds to secretion of the metabolite from cells into the environment.

In this manuscript, we define $\vec{c}$ as the “chemical environment”, and all possible values of $\vec{c}$ constitute the “chemical space”. Within a cell, the concentration of metabolites is influenced by uptake/secretion rates, and influences the growth rate. Different metabolic models assume different forms for such influences, and we use $f(I(\vec{c}, q), \vec{q})$ to represent the rate of change in $\vec{q}$:

$$\frac{d\vec{q}_{\sigma}}{dt} = f(I(\vec{c}, \vec{q}_{\sigma}, \vec{\alpha}_{\sigma}), \vec{q}_{\sigma}). \quad (3)$$

Eqs. (1)-(3) represent a general chemostat model with a single species. The simplicity of the chemostat has inspired many theoretical studies of resource competition. Different model assumptions about how species grow and consume nutrients have produced a variety of intriguing behaviors and conclusions. However, the origins of these differences are not always simple to discern. To facilitate the evaluation of metabolic strategies, we next present a graphical
representation that allows ready visualization of the feedback between species and the
environment in a chemostat.

The steady-state environment created by species $\sigma$ can be obtained by setting Eqs. (1)-(3) to zero. First, when Eq. (3) is equal to zero, $\bar{q}_\sigma$ can be solved as a function of $\hat{c}$, reducing

$g(\hat{c}, \bar{q}_\sigma, \bar{a}_\sigma)$ and $l_1(\hat{c}, \bar{q}_\sigma, \bar{a}_\sigma)$ to functions fully dependent on the variable $\hat{c}$, namely $g(\hat{c}, \bar{a}_\sigma)$ and $l_1(\hat{c}, \bar{a}_\sigma)$. Graphically, the steady state created by a single species can be visualized by the intersection of two nullclines, derived from Eq. (1) and Eq. (2), respectively (details in Methods).

Next, setting Eq. (1) to zero leads to a $p$-dimensional version of the ZNGI, which we name the “growth contour”. For a given metabolic strategy $\bar{a}_\sigma$, the growth-rate function $g(\hat{c}, \bar{a}_\sigma)$ maps different points in the chemical space onto varying growth rates (background color in Fig 1B). At steady state, the relation $dm_\sigma/dt = 0$ (Eq. (1)) requires the growth rate to be exactly equal to the dilution rate (assuming nonzero cell density). Therefore, the contour in chemical space satisfying $g(\hat{c}, \bar{a}_\sigma) = d$ indicates all possible environments that could support a steady state of the strategy $\bar{a}_\sigma$ (red curve in Fig 1B). This contour reflects how the chemical environment determines cell growth.

Thirdly, the nullcline derived from Eq. (2) reflects the impact of cellular metabolism on the chemical environment. At steady state, the nutrient influx should be equal to the summation of dilution and cellular consumption. When Eq. (2) is set to zero, varying values of cell density $m$ lead to different $\hat{c}$ (Eq. (S5)), constituting a one-dimensional “flux-balance curve” in chemical space (purple, cyan, and blue curves in Fig 1B).

Finally, the steady-state chemical environment $\hat{c}_{\sigma,ss}$ created by the species $\sigma$ is the intersection of the growth contour and the flux-balance curve (Fig 1B, red dot). Changes in the chosen conditions $d$ and $\bar{c}_{\text{supply}}$ influence the shapes of the growth contour and the flux-balance curve separately, enabling clear interpretation of chemostat experiments under varied conditions (details in Methods). Sometimes for a given steady-state environment $\hat{c}_{\sigma,ss}$, one would like to derive the supply concentrations that ultimately lead to this environment. Setting Eq. (2) to zero and fixing $\hat{c}_{\sigma,ss}$, varying values of cell density $m$ lead to a straight line in the space of supply concentrations, which we call the “supply line” (see Methods for details). Despite the fact that the supply space and the chemical space are distinct, they share the same units of concentration in each dimension. Therefore, for ease of visualization we typically show supply lines along with other features in the same nutrient space (Eq. (S6), black dashed line in Fig S1).

**Rule of invasion, and the environment-dependent fitness landscape**

We use the outcome of competition between species to evaluate metabolic strategies, assuming each species $\sigma$ adopts a fixed strategy $\bar{a}_\sigma$. We first focus on the outcome of invasion, which corresponds to the “invasion fitness” in adaptive dynamics: here, “invasion” is defined as the introduction of a very small number of an “invader” species to a steady-state chemostat already occupied by a set of “indigenous” species, and the “invasion growth rate” is quantified by the growth rate of the invader species at the moment of introduction. Unlike adaptive dynamics, the invasion growth rate is evaluated with respect to the chemical environment created by the indigenous species, rather than with respect to the population composition of the indigenous species.
In the chemical space, the outcome of an invasion can be summed up by a simple geometric rule, as demonstrated in Fig 1C and D. The growth contour of the invader (species Red) separates the chemical space into two regions: an “invasion zone” where the invader grows faster than dilution (green-colored region in Fig 1C and D), and “no-invasion zone” where the invader has a growth rate lower than dilution. If the steady-state environment constructed by the indigenous species (species Blue) is located within the invasion zone of the invader, the invader will initially grow faster than dilution. Therefore, the invader will expand its population and the invasion will be successful (Fig 1C). By contrast, if the steady-state chemical environment created by the indigenous species lies outside of the invasion zone, the invasion will be unsuccessful (Fig 1D, same species as in Fig 1C but with a different supply condition, and therefore a different steady state). (See Methods for details.)

In the steady-state environment created by the indigenous species, each strategy $\vec{\alpha}$ has an invasion growth rate. We define an environment-dependent “fitness landscape” as the relation between the invasion growth rate and the metabolic strategy of invaders (Eq. (S8)-(S9), see Methods for details). Different indigenous species can create different chemical environments, and thus give rise to different shapes of the fitness landscapes.

**Mutual invasion, a flat fitness landscape, and unlimited coexistence**

The rule of invasion allows for easy assessment of competition dynamics. The emergence of complex dynamics generally requires that competitiveness be non-transitive. For example, if species Red can invade species Blue, that does not mean Blue cannot invade Red. Such mutual invasibility can be observed in substitutable-resource metabolic models, with a simple version illustrated in Fig 2A: two substitutable nutrients $a$ and $b$, such as glucose and galactose, contribute linearly to biomass increase. Since a substantial investment of protein and energy is required for nutrient uptake, the model assumes a trade-off between the allocation of internal resources to import either nutrient. Specifically, a fraction of resources $\alpha_a$ is allocated to import $a$ and a fraction $(1 - \alpha_a)$ to import $b$. As shown in Fig 2B, while the steady-state environment created by Blue is located within the invasion zone of Red, the steady-state environment created by Red is also located within the invasion zone of Blue. According to the rule of invasion, each species can therefore invade the steady-state environment created by the other. In the face of such successful invasions, the only possible stable chemical environment for this system is at the intersection of two growth contours, where the two species can coexist.

The environment-dependent fitness landscape readily explains this coexistence: In the steady-state environment created by Red ($\alpha_a = 0.6$), strategies with smaller $\alpha_a$ have higher fitness (Fig 2C, upper panel). In the steady-state environment created by Blue ($\alpha_a = 0.2$), the fitness landscape changes in shape, and strategies with larger $\alpha_a$ have higher fitness (Fig 2C, middle panel). Therefore, each species creates an environment that is more suitable for its competitor, which leads to coexistence.

Typically in resource-competition models, the number of coexisting species cannot exceed the number of resources (Armstrong & McGehee, 1980; Hardin, 1960; Levin, 1970; McGehee & Armstrong, 1977). This conclusion can be understood intuitively from a graphical approach: The steady-state growth of a species can only be achieved on the growth contour of this species. The
stable coexistence of $N$ species can thus only occur at the intersection of their $N$ corresponding growth contours. Generally, an $N$-dimensional chemical space allows a unique intersection of no more than $N$ surfaces, and the diversity of species is therefore bounded by the number of metabolites in the environment. This theoretical restriction on biodiversity, made formal as the “competitive exclusion principle”, contradicts the tremendous biodiversity manifested in the real world (Friedman, Higgins, & Gore, 2017; Goldford et al., 2018; Maharjan, Seeto, Notley-McRobb, & Ferenci, 2006). There have been a multitude of theoretical efforts to reconcile this contradiction (Huisman & Weissing, 1999, 2001) (Goldford et al., 2018; Pfeiffer & Bonhoeffer, 2004) (Beardmore, Gudelj, Lipson, & Hurst, 2011; Taillefumier et al., 2017).

The mutual invasibility in linear-summation metabolic models with trade-offs presents one possibility resolution of the competitive exclusion paradox. For the environment co-created by Blue and Red (Fig 2B, purple dot), the fitness landscape becomes flat (Fig 2C, bottom panel): species with any metabolic strategy will grow at the same rate as dilution in this environment. Therefore, in this system, once a pair of species with a mutual-invasion relationship construct the chemical environment together, all species become effectively neutral and can coexist indefinitely (see Methods for details). This graphical approach to mutual invasion and the flat fitness landscape provide an intuitive representation of species self-organizing to a state of unlimited coexistence beyond competitive exclusion.

**Rock-paper-scissor invasion loop and oscillation**

Resource-competition models focusing on various aspects of cellular metabolism vary in their assumptions regarding $g(\bar{c}, \bar{x}, \bar{a})$, $I(\bar{c}, \bar{x}, \bar{a})$, and $f(\bar{c}, \bar{x}, \bar{a})$, and can lead to diverse results for community structure and coexistence. However, the above general “rule of invasion” allows us to treat these divergent resource-competition models in a unified framework. In the following example, we utilized a metabolic model slightly different from that of Fig 2, to show that a dynamic fitness landscape is indispensable for coexistence. In the metabolic model shown in Fig 3A, three substitutable nutrients, $a$, $b$, and $c$, contribute additively to cell growth. In this three-dimensional chemical space, the growth contour for each species is a two-dimensional surface (Fig 3B). In addition to requiring enzymes to import the raw forms of these nutrients as in the model of Fig 2A, enzymes are also required to convert the imported raw materials into biomass. In this model, a six-element $\bar{a}$ is required to describe the metabolic strategy, and there is the possibility of “mismatches” in the fraction of internal resources allocated to import and to convert the same nutrient. Such mismatches can produce a “rock-paper-scissor” invasion loop (Fig S3A): In the environment created by species 1, species 2 has a higher fitness but not species 3; therefore species 2 can invade species 1 and establish its own environment; however, this environment lies within the invasion zone of species 3 (Fig 3B) but not of species 1, therefore species 3 subsequently invades; then in turn, species 3 create an environment where species 1 has the highest fitness. Such a loop of invasions leads to oscillatory population dynamics (Fig 3C, upper panel), with an ever-changing fitness landscape (Fig 3C, lower panel).

Oscillation and even chaos in resource-competition model have been demonstrated by Huisman et al. (Huisman, van Oostveen, & Weissing, 1999), and shown to allow dynamical coexistence beyond competitive exclusion. The simple model presented here illustrates how oscillation can be understood as a loop of invasion creating an ever-changing fitness landscape.
**Multistability and chain of invasion**

When species create environments that are more favorable for their competitors, mutual-invasion and oscillations can occur. Can species create environments that are hostile to their competitors, and if so what will be the consequences?

Figure 4A shows a simple metabolic model with two essential nutrients $a$ and $b$, such as nitrogen and phosphorus (see Methods for details). Similar to the model in Fig 2A, the model assumes a trade-off between the allocation of internal resources to import nutrients, so that a resource allocation strategy is fully characterized by the fraction of resources $\alpha_a$ allocated to import nutrient $a$. The growth rate is taken to be the minimum of the two input rates (Odum & Barrett, 1971). As shown in Fig 4B, two species, Red and Blue, each creates a chemical environment outside of the invasion zone of each other. According to the rule of invasion, neither can be invaded by the other. Therefore, the steady state of the community depends on initial conditions – whichever species occupies the chemostat first will dominate indefinitely. It is worth noting that despite the fact that coexistence is excluded in a single chemostat under this metabolic model, the ability of species to create a self-favoring environment allows the spontaneous emergence of spatial heterogeneity and coexistence in an extended system with multiple linked chemostats (Fig S6). In ecology, the spontaneously emergence of spatial heterogeneity has been shown for species with the capacity to construct their own niches (Levin, 1970; Schertzger, Staver, & Levin, 2015), and the chain of chemostats provides a simple model for such spatial coexistence.

From the perspective of the strategy-growth relationship (Fig 4B, inset), species Red ($\alpha_a = 0.65$) creates a fitness landscape where small $\alpha_a$ is disfavored. Symmetrically, species Blue ($\alpha_a = 0.35$) creates a fitness landscape where large $\alpha_a$ is disfavored. However, neither Red nor Blue sits on the top of the fitness landscape each one creates (Fig 4C). In the fitness landscape created by Blue, a slightly larger $\alpha_a$ (green diamond in Fig 4C) has the highest growth rate. Consequently, species adopting the Green strategy can invade Blue. Nevertheless, species Green is not on the top of its own fitness landscape as an even larger $\alpha_a$ (yellow diamond in Fig 4C) maximizes the growth rate in the environment created by Green. A series of replacements by the fastest-growing species in the environment created by the former species creates a chain of invasion.

In this particular model after four steps of replacement, bistability appears. The species with $\alpha_a$ marked by Deep Purple, which is reached by the chain of invasion going from Blue, to Green, to Yellow, to Deep Green, cannot invade the original species Blue. A similar relationship holds between Cyan and Red. This phenomenon highlights the difference between ecological stability and evolutionary stability: Ecologically, as both Blue and Deep Purple create a fitness landscape where the other species grows slower than dilution, they constitute a bistable system. However, evolutionarily, “mutants” with slightly larger $\alpha_a$ can invade Blue, bringing the system towards Deep Purple until bistability collapses.

**Non-invasible strategies**

In this model, with symmetric parameters, the only evolutionarily stable strategy is $\alpha_a = 0.5$ (black diamond in Fig 4C). This is the only strategy that locates itself on the top of the fitness landscape it creates, and therefore cannot be invaded by any other species. This simple model
demonstrates a general definition of optimal (aka evolutionarily stable or non-invasible) strategies: those strategies that create a fitness landscape which places themselves on the top (Eq. (S10)).

A chemical environment defines a fitness landscape, and the steady-state chemical environment created by species is influenced by supply condition, dilution rate, and the details of cell metabolism. Therefore, different chemostat parameters and different metabolic models lead to different optimal strategies. In the following, we described a generally applicable protocol for obtaining the non-invasible strategies, using the metabolic model in Fig 4A as the example (Fig 4D, details in Methods): First, under a chemical environment \( \vec{c} \), the maximal growth rate \( g_{\text{max}}(\vec{c}) \) (background color in Fig 4D) and the corresponding resource allocation strategy \( \vec{\alpha}_{\text{max}}(\vec{c}) \) can be obtained analytically or via numerical search through the strategy space (Eq. (S11)). \( g_{\text{max}}(\vec{c}) \) and \( \vec{\alpha}_{\text{max}}(\vec{c}) \) are independent of the chemostat parameters \( \vec{c}_{\text{supply}} \) and \( d \).

Second, the maximal growth contour for dilution rate \( d \) is defined as all chemical environments \( \vec{c} \) that support a maximal growth rate of \( d \) (Eq. (S12)). Different maximizing strategies \( \vec{\alpha}_{\text{max}}(\vec{c}) \) exist at different points of the maximal growth contour, as shown by the colors of the curve in Fig 4D. By definition, the maximal growth contour envelops the growth contour of any single strategy, and chemical environments on the maximal growth contour are outside of the invasion zone of any strategy. Therefore, if a species is able to create a steady-state environment on the maximal growth contour, it cannot be invaded. Finally, different \( \vec{c}_{\text{supply}} \) form different maximal flux-balance curves (Eq. (S14)), which intersect with the maximal growth contour at one point \( \vec{c}_{\text{opt}} \). Species \( \vec{\alpha}_{\text{max}}(\vec{c}_{\text{opt}}) \) that adopt the maximizing strategy at \( \vec{c}_{\text{opt}} \) create the environment \( \vec{c}_{\text{opt}} \), and are therefore immune to invasion. Under different \( \vec{c}_{\text{supply}} \), different species become non-invasible (orange, green, and blue growth contours in Fig 4D), and the supply lines emanating from different points on the maximal growth contour indicate the supply conditions for which the corresponding strategies are optimal.

**Evolutionarily stable coexistence**

Given \( d \) and \( \vec{c}_{\text{supply}} \), the maximal growth contour and the maximal flux-balance curve are unique, therefore there is only one \( \vec{c}_{\text{opt}} \). Does the uniqueness of \( \vec{c}_{\text{opt}} \) imply a single evolutionarily stable species? Or is coexistence still possible even in the face of evolution? In a recent work (Taillefumier et al., 2017), this question was addressed by modeling a population of microbes competing for steadily supplied resources. Through in-silico evolution and network analysis, the authors found that multiple species with distinct metabolic strategies can coexist as evolutionarily-stable co-optimal consortia, which no other species can invade.

Using a simplified version of Taillefumier et al.’s model (Fig 5A), we employ the graphical approach to help identify the requirements for such evolutionarily-stable coexistence and the role of each species in supporting the consortium. In this model, at the cost of producing the necessary enzymes, cells are not only able to import external nutrients, but can also convert any one of the internal nutrients into any other. Meanwhile, nutrients passively diffuse in and out of the cell. The internal concentrations of nutrient \( a \) and nutrient \( b \) are both essential for cell growth (see Methods for detail). Therefore, metabolic trade-offs in this system have four elements: the fraction of internal resources allocated to import nutrient \( a (\alpha_a) \) or nutrient \( b (\alpha_b) \) and/or convert...
one nutrient into another ($\alpha_{ab}$ converts internal $b$ into $a$, and $\alpha_{ba}$ converts internal $a$ into $b$).

Each species is defined by its internal resource allocation strategy $\mathbf{\alpha} = (\alpha_a, \alpha_b, \alpha_{ab}, \alpha_{ba})$.

Following the general protocol described in the previous section, we first identified the maximal growth rates $g_{\text{max}}(\mathbf{c})$ and the corresponding strategy or strategies $\mathbf{\alpha}_{\text{max}}(\mathbf{c})$ at each point $\mathbf{c}$ in the chemical space, and generated maximal growth contours for different dilution rates (Fig 5B). The maximal growth contours are not smoothly continuous, nor are the corresponding strategies. In chemical space, three distinct sectors of maximizing strategies appear (Fig 5B, Fig S6A):

When nutrient $a$ is very low compared to $b$, the maximizing strategy is a “$b$-$a$ converter” which imports $b$ and converts it into $a$ (blue sector, only $\alpha_b$ and $\alpha_{ab}$ are non-zero). Symmetrically, when $a$ is comparatively high, the optimal strategy is a “$a$-$b$ converter” (green sector, only $\alpha_a$ and $\alpha_{ba}$ are non-zero). Otherwise, the maximizing strategy is an “importer” which imports both nutrients without conversion (red sector, only $\alpha_a$ and $\alpha_b$ are non-zero). On the border between sectors, the maximal growth contour has a discontinuous slope.

Optimal coexistence occurs at these discontinuous points. If an environment point $\mathbf{c}_0$ is located in a continuous region of the maximal growth contour, only one maximizing strategy $\mathbf{\alpha}_{\text{max}}(\mathbf{c}_0)$ exists for that environment (maximizing strategies along the maximal growth contour are indicated by colored squares in Fig 5C). Supply conditions that make $\mathbf{\alpha}_{\text{max}}(\mathbf{c}_0)$ the optimal strategy (i.e. allow $\mathbf{\alpha}_{\text{max}}(\mathbf{c}_0)$ to create the steady-state environment $\mathbf{c}_0$) constitute the supply line for $\mathbf{c}_0$ and $\mathbf{\alpha}_{\text{max}}(\mathbf{c}_0)$. However, at the discontinuous points of the maximal growth contour, where two classes of strategies meet, two different strategies are both maximizing. For example, at the purple dot in Fig 5C a strategy belonging to the “$b$-$a$ converter” class (species Blue) and one belonging to the “importer” class (species Red) are both maximizing strategies. Each strategy derives a supply line from the purple dot (black dashed line, Fig 5C). The two supply lines span a gray region where no supply line from any single strategy enters. Correspondingly, for any supply conditions inside the gray region, no single species can alone create an environment on the maximal growth contour. For example, under the supply condition shown by the black open circle in the gray region, species Blue and species Red both create chemical environments that lie within the maximal growth contour (blue and red dots, Fig 5C), and are thus subject to invasion by other species. Nevertheless, the species-specific growth contours of Blue and of Red intersect at the purple point on the maximal growth contour. Therefore, only when Blue and Red coexist can they co-create an environment on the maximal growth contour, and thus be resistant to invasion from any other species. Indeed, when we simulate multiple species with different maximizing strategies under the supply condition indicated by the open black circle, species Blue and species Red are the only two that survive (Fig 5C, inset).

The optimal coexistence of species Blue and species Red can be understood intuitively from the dynamic fitness landscape. Given a chemical environment, the relation between $\alpha_a$ and growth rate of importer (red curve) or $a$-$b$ converter (green curve), and that between $\alpha_b$ and growth rate of $b$-$a$ converter (green curve) constitute the fitness landscape of species adopting different possible maximizing strategies (Fig 5D). In the environment created by species Blue (blue dot in Fig 5C), not only will some importers grow faster than Blue, species Blue (strategy marked by blue diamond) is not even on the fitness peak of its own class (Fig 5D, upper box). Similarly, in the environment created by species Red, the strategy of Red is not at the top of the fitness landscape (Fig 5D, middle box). By contrast, in the environment co-created by species Blue and
Red (purple dot in Fig 5C), their strategies are at the top of the fitness landscapes of their own classes and at equal height. For all supply conditions in the gray region, species Blue and species Red jointly drive the nutrient concentrations to the discontinuous point of the optimal growth contour, and thereby achieve evolutionarily stable coexistence.

Species creating a new nutrient dimension

One possible solution to the competitive-exclusion paradox is the creation of new nutrient "dimensions" by species secreting metabolites that can be utilized by other species. For example, E. coli secretes acetate as a by-product of glucose metabolism. Accumulation of acetate impedes the growth of E. coli on glucose (Luli & Strohl, 1990), but the acetate can be utilized as a carbon source, e.g. by mutant strains that emerge in long-term evolution experiments (D'Souza et al., 2018; Rosenzweig, Sharp, Treves, & Adams, 1994).

To explore the possibilities of evolutionarily stable coexistence when species create new nutrients, we used a simplified model to represent multi-step energy generation with a dual-role intermediate metabolite (Fig 6A). A single chemical energy source \( S \) is supplied into the chemostat. The pathway for processing \( S \) consists of four relevant reactions driven by designated enzymes: External \( S \) can be imported and converted into intermediate \( I_{\text{int}} \) to generate ATP (with a corresponding fraction of the enzyme budget \( \alpha_{\text{ATP1}} \)). The intermediate has a dual role in energy production: on the one hand, it positively contributes to ATP production via a downstream reaction (with a fraction of the enzyme budget \( \alpha_{\text{ATP2}} \)); on the other hand, it negatively contributes to ATP production through product inhibition of the first energy-producing reaction. To deal with this negative effect of internal intermediate, cells may synthesize transporters (with a fraction of enzyme budget \( \alpha_{\text{exp}} \)) to export intermediate out into environment, where it becomes external intermediate \( I_{\text{ext}} \). By this reaction, cells can increase the dimension of chemical space from one (\( S \)) into two (\( S \) and \( I_{\text{ext}} \)). Cells can also import \( I_{\text{ext}} \) into \( I_{\text{int}} \) (fraction of enzyme budget \( \alpha_{\text{imp}} \)), then use \( I_{\text{int}} \) as an energy source via the second reaction. (See Methods for details.)

The metabolic strategy \( \vec{\alpha} \) in this model has four components: \( \vec{\alpha} = (\alpha_{\text{ATP1}}, \alpha_{\text{ATP2}}, \alpha_{\text{exp}}, \alpha_{\text{imp}}) \). When we examine the maximizing strategies and maximal growth rates in the chemical space, three distinct classes of strategy emerge (Fig 6B). When \( S \) is abundant and \( I_{\text{ext}} \) is low, the maximizing strategies have only two non-zero components, \( \alpha_{\text{ATP1}} \) and \( \alpha_{\text{exp}} \) (Fig S6B), meaning this class of species only imports \( S \), for the first energy-generating reaction, then exports intermediate as waste. Therefore, we call strategies in this class “polluters” (blue section in Fig 6B, Fig S3C). When \( I_{\text{ext}} \) is high while \( S \) is low, the maximizing strategies have two different non-zero components, \( \alpha_{\text{ATP2}} \) and \( \alpha_{\text{imp}} \) (Fig S6B), meaning this class of species relies solely on \( I_{\text{ext}} \) as its energy source. We call these strategies “cleaners” as they clean up \( I_{\text{ext}} \) from the environment, (green section in Fig 6B, Fig S6C). When there are comparable amounts of \( S \) and \( I_{\text{ext}} \) present, a third class of maximizing strategies appears: these cells neither export nor import intermediates, but rather allocate all their enzyme budget to \( \alpha_{\text{ATP1}} \) and \( \alpha_{\text{ATP2}} \) to carry out both energy-producing reactions. We call species in this class “generalists” (red section in Fig 6B, Fig S6C).

As shown in Fig 6B, on the borders between classes of strategies in chemical space, the maximal growth contours turn discontinuously. These points of discontinuity, as in the example in the previous section, are chemical environments corresponding to evolutionarily stable coexistence of species from distinct metabolic classes. The classes of optimally coexisting species change
with dilution rate. When the dilution rate is low \((d = 0.4\), Fig 6C\), at the discontinuous point of
the maximal growth contour, the corresponding two maximizing strategies are one polluter
(species \textit{Blue}) and one cleaner (species \textit{Green}). Their supply lines span a gray region where both
species \textit{Blue} and species \textit{Green} are required to create a steady-state environment on the maximal
growth contour. As by assumption we are only supplying the system with S, the supply condition
always lies on the x-axis of concentration space. For the supply condition shown by the black
open circle in Fig 6C, polluter \textit{Blue} creates a chemical environment (blue dot) far from the
maximal growth contour. When the cleaner \textit{Green} is added to the system, not only does the
biomass of \textit{Blue} increase (inset), but also the steady-state chemical environment moves to the
discontinuous point of the maximal growth contour (cyan dot), where both \textit{Blue} and \textit{Green}
occupy the peaks of their fitness landscapes (Fig 6D). This result is consistent with the long-term
evolution experiment of \textit{E. coli} and also intuitive: polluter \textit{Blue} and cleaner \textit{Green} form a
mutually beneficial relationship by, respectively, providing nutrients and cleaning up waste for
each other, thereby reaching an optimal cooperative coexistence.

A quite different coexistence occurs at higher dilution rate \((d = 0.6\), Fig 6E\). Growth contours at
this dilution rate show two turning points, but neither are between the polluter and the cleaner
class. One discontinuous point is between the cleaner class (green squares) and the generalist
class (red squares), but the gray region spanned by the corresponding supply lines does not cover
the x-axis and so does not represent an attainable coexistence when only S is supplied. The other
discontinuous point is between the generalist class and the polluter class (blue squares). The gray
region spanned by the supply lines of the corresponding two maximizing strategies of generalist
class (species \textit{Red}) and polluter class (species \textit{Blue}) does cover the x-axis. Therefore, a supply
condition with only S within the gray region (e.g., the black open circle) leads to the optimal
coevolution of generalist \textit{Red} and polluter \textit{Blue} on the discontinuous point (purple dot), despite
the fact that they do not directly benefit each other. Indeed, when the generalist \textit{Red} is added to a
system with polluter \textit{Blue} and a cleaner \textit{Green}, the cleaner \textit{Green} goes extinct and the biomass of
the polluter \textit{Blue} decreases (inset). Nevertheless, the steady-state chemical environment is moved
from a cyan dot lying inside the maximal growth contour to the purple dot lying on the maximal
growth contour. In the environment of the cyan dot created by cleaner \textit{Green} and polluter \textit{Blue},
\textit{Blue} is not on the top of the fitness landscape of the polluter class (Fig 6F, upper box). By
contrast, for the fitness landscape created by polluter \textit{Blue} and generalist \textit{Red} (Fig 6F, lower
box), despite being lower in biomass, \textit{Blue} occupies the top of the landscape. Therefore, the
optimal coexistence of this polluter and this generalist does not arise from direct cooperation, but
rather from collaborating to defeat other competitors.
Evaluating microbial metabolic strategies within an ecological context is the major focus of this work. Due to the intensity of competition in the microbial world, it is accepted that natural selection has extensively shaped microorganisms’ internal resource allocation strategies and the regulatory mechanisms controlling these strategies (S. Goyal et al., 2010; Liebermeister et al., 2014). Therefore, a quantitative mapping from metabolic strategies to fitness consequences can further our understanding of both regulation and evolution (Bajic & Sanchez, 2019). Many previous studies of metabolic strategies directly assumed the optimization goal of microbial metabolism to be biomass gain, with the chemical environment acting as a fixed input (Scott, 2014 #86) (Wang & Tang, 2017) (Schuetz, 2012 #84) (Roller, Stoddard, & Schmidt, 2016) (Brophy, 2014 #85), which simplifies the problem into a search for a maximum on a static “fitness landscape”. However, in the natural world where metabolic strategies compete and evolve, the feedback between species and their environment produces an intrinsically dynamic fitness landscape in which the actions of one species can influence the fitness of all species (Bajic, Vila, Blount, & Sanchez, 2018). One profound example is the Great Oxygenation Event, when cyanobacteria created an oxygen-rich atmosphere (Kasting & Siefert, 2002), causing a massive extinction of anaerobic bacteria but also stimulating an explosion of biodiversity (Schirrmeister, de Vos, Antonelli, & Bagheri, 2013). Therefore, metabolic strategies need to be assessed within an ecological context, taking into consideration not only how species respond to the environment, but also how species construct their own environment.

For the past thirty years, researchers have been utilized various mathematical tools to study flexible fitness landscapes that change with space, time, and population composition. The prerequisite for an intrinsically dynamical fitness landscape – that the species composition influences the fitness of all species in the system – takes a particularly simple form in chemostat-type resource-competition models: Microbes shape their local environment by exchanging metabolites within a shared chemical environment, which determines the growth rate of all cells. Focusing on the chemical environment created by microbial metabolism, we exhibited a set of intuitive and general procedures for analyzing strategies within various metabolic models. Namely, we showed that to compare a set of fixed strategies, the geometric relationships between their growth contours and their steady-state chemical environments yield an immediate prediction for the outcome of competitions. In searching for optimality over the continuous family of strategies, the “maximal growth contour” envelope of all growth contours provides candidates, and the supply condition selects the non-invasible strategy or strategies from among these candidates via the flux-balance curve. Such selection of optimal strategies also supports the conclusion that having the fastest growth rate in an environment does not necessarily imply being the most competitive strategy, as this strategy may shift the environment in an unfavorable direction. To be non-invasible, strategies also need to be able to construct the environment for which they are best suited.

From the perspective of chemostat-related experiment, this work demonstrates the subtleties of controlling nutrient limitation in chemostats. The capacity of species to shape their own environment, even in a system as simple as a chemostat, presents challenges to controlling which nutrient or nutrients are limiting. By traditional definition, if increasing a certain nutrient leads to an increase of a cell’s growth rate, that nutrient is considered “limiting”. However, growth rate is
invariant in a chemostat, being set experimentally by the dilution rate, so inferring nutrient limitation requires special attention. For example, if one sees the same cellular responses under different nutrient supplies, what can one conclude? Cells may be creating the same chemical environment out of different supply conditions (cf. Fig S1A), or alternatively cells may be transducing different chemical environments into the same physiological response through mechanisms such as “ratio sensing” (Escalante-Chong et al., 2015). Our graphical approach combined with direct measurements of steady-state nutrient concentrations in the chemostat can precisely define and help guide the control of nutrient limitation (Boer et al., 2010). As described above, changes in supply concentrations shift the flux-balance curve, but do not change the shape of the growth contour. Therefore, by experimentally varying the supply conditions and measuring the chemical environment created by cells, the shape of the growth contour can be obtained. The resulting slope of the growth contour provides information on nutrient limitation even in the absence of detailed knowledge about a cell’s metabolism. For example, in the nutrient $a$ - nutrient $b$ plane, a near-horizontal growth contour indicates $b$-limited growth while a near-vertical growth contour means $a$-limited growth, and an intermediate slope implies that the two nutrients are co-limiting.

Our work also has implications for microbial community assembly. The competitive exclusion principle poses a long-term puzzle in ecology: since species are in constant competition in the natural world, why doesn’t the fittest species outcompete the others and become the sole survivor? Actually, on the basis of simple resource-competition models, it has been argued that the number of stably coexisting species cannot exceed the number of resources, leading to the so-called competitive exclusion principle (Armstrong & McGehee, 1980; Hardin, 1960; Levin, 1970; McGehee & Armstrong, 1977). Nevertheless, tremendous biodiversity manifests in the real world, from environmental surveys to controlled lab experiments (Friedman et al., 2017; Goldford et al., 2018; Maharjan et al., 2006). Our work is not aimed at adding another solution to the paradox of the plankton; rather, our framework helps in summarizing the criteria for coexistence on both ecological and evolutionary timescales. On the ecological scale where a limited number of species with different strategies compete for resources, the intransitivity of competitiveness as a result of fitness landscape deformability is the key to coexistence. Given resource allocation trade-offs, the growth contours of any pair of strategies must intersect, clearly demonstrating why trade-offs prevent a single species from unconditional dominance, allowing various forms of intransitivity under different metabolic models. In the case of substitutable nutrients, the fitness landscape can even be made flat – allowing unlimited coexistence. When the rule of invasion allows non-transitive loops, oscillations and chaos can occur, which have been shown to allow coexistence beyond competitive exclusion (Huisman & Weissing, 1999, 2001). In addition, when multiple nutrients are all essential, the ability of each species to create an environment that favors itself allows for the spontaneous emergence of spatial heterogeneity in an extended system (Fig S5). Meanwhile, on the evolutionary time scale where mutation/adaptation allows searches for the “most suitable” strategies among infinite possibilities, an ongoing threat to diversity is that selection may produce a supreme winner that takes over the habitat. With the dynamic fitness landscape and the maximal growth contour approach, we showed that the condition for evolutionarily stable coexistence is indeed more restricted, occurring only at the discontinuous points of the maximal growth contour. Nevertheless, via the species-environment feedback, a large number of supply conditions can
self-organize to the discontinuous point, where multiple species co-create a non-invasible environment where they jointly locate on the peak of the fitness landscape.

Many future directions can follow this work. From the perspective of experiment, our framework can assist in analyzing and interpreting results of microbial evolution in the lab (Van den Bergh, Swings, Fauwart, & Michiels, 2018), where the continual emergence of new mutants under defined experimental conditions suggests an intrinsically dynamic fitness landscape. From the perspective of theory, we do not yet have a rigorous mathematical theorem concerning the conditions for discontinuity of the maximal growth contour, nor proof that discontinuity necessarily leads to evolutionarily stable coexistence. Theoretical developments paralleling those on the general existence of ecologically stable states (De Leenheer et al., 2006; Marsland III, Cui, & Mehta, 2019) would bring a more comprehensive understanding of evolutionarily optimal states in metabolic models. Besides, the metabolic models considered in this work are highly simplified. Going forward, more detailed and experimentally-based models can be examined using the same graphical and dynamical fitness landscape framework.
Programs for this work are coded in MATLAB R2018a. A repository of all tools used to generate the results can be found at: https://github.com/zhiyuanli1987/Qbiotoolbox.git

**Total RNA and total protein measurements**
The methods for total RNA and protein measurements shown in Figure S2 are described in Li et al., 2018.

### Supplemental Table 1: Symbols

| Chemostat parameters | Nutrient supply. \( c_i, \text{supply} \) is the concentration of the \( i \)-th nutrient in the supply. |
|-----------------------|--------------------------------------------------------------------------------------------------|
| \( \tilde{c}_{\text{supply}} = (c_{1, \text{supply}}, c_{2, \text{supply}}, \ldots, c_p, \text{supply}) \) | |
| \( d \) | Dilution rate (same as supply influx rate to keep volume fixed). |

| Chemostat variables | Chemical environment inside the \( k \)-th chemostat. \( c_{i,k} \) is the concentration of the \( i \)-th nutrient within the medium of the \( k \)-th chemostat. All possible \( \tilde{c} \) constitute the “chemical space”. |
|---------------------|--------------------------------------------------------------------------------------------------|
| \( \tilde{c}_k = (c_{1,k}, c_{2,k}, \ldots, c_{p,k}) \) | |
| \( m_{\sigma,k} \) | Biomass density of species \( \sigma \) in the \( k \)-th chemostat. |

| Species-specific quantities | Resource allocation strategy of species \( \sigma \). \( \alpha_{j,\sigma} \) is the fraction of internal resources allocated to the \( j \)-th cellular function by species \( \sigma \). |
|-----------------------------|--------------------------------------------------------------------------------------------------|
| \( \tilde{\alpha}_\sigma = (\alpha_{1,\sigma}, \alpha_{2,\sigma} \ldots) \) | |
| \( \tilde{q}_\sigma \) | Intracellular concentrations of growth-related metabolites for species \( \sigma \). |
| \( g(\tilde{c}, \tilde{q}, \tilde{\alpha}) \) | Growth rate as a function of \( \tilde{c}, \tilde{q}, \) and \( \tilde{\alpha} \). |
| \( l_i(\tilde{c}, \tilde{q}, \tilde{\alpha}) \) | Intake rate per biomass of the \( i \)-th nutrient as a function of \( \tilde{c}, \tilde{q}, \) and \( \tilde{\alpha} \). \( l_i(\tilde{c}, \tilde{q}, \tilde{\alpha}) \) can be negative to describe cells exporting secondary metabolites. |
| \( r \) | Biomass concentration within an average cell, taken to be a constant that is always equal to 100. |
| \( \tilde{f}(\tilde{I}(\tilde{c}, \tilde{q}, \tilde{\alpha}), \tilde{q}, \tilde{\alpha}) \) | Functions defining the changing rate of intracellular metabolite concentrations \( \tilde{q}, \) as a function of \( \tilde{c}, \tilde{q}, \) and \( \tilde{\alpha} \). |
| \( GC_\sigma \) | Growth-rate contour of species \( \sigma \). |
| \( FB_\sigma \) | Flux-balance curve of species \( \sigma \). |
| \( \tilde{c}_{\sigma,ss} \) | The steady-state environment created by one species \( \sigma \). |
| \( SL_\sigma(\tilde{c}) \) | The supply line for species \( \sigma \) in environment \( \tilde{c} \). |
| \( \{\sigma^*\} \) | A set of species stably surviving in a chemostat. The set can contain one or more species. |
|---------------|----------------------------------------------------------------------------------------------------------------------------------|
| \( \tilde{c}_{\{\sigma^*\}} \)_{ss} | The steady-state environment created by a set of species \( \{\sigma^*\} \). |

**Metabolic model and resource allocation strategy**

In modeling population dynamics in a chemostat, multiple assumptions need to be made concerning how cells sense the environment, import nutrients, export metabolites, utilize resources, and grow in biomass. Different assumptions result in different metabolic models. Some metabolic models focus on trade-offs in resource allocation, as the amount of internal resources “owned” by a cell, including proteins and energy, is limited. Cells need to allocate these limited resources into different cellular functions, such as metabolism, gene expression, reproduction, motility, maintenance, etc. We use \( \alpha_j,\sigma \) to represent the fraction of internal resources allocated to the \( j \)-th cellular function of species \( \sigma \), with \( \vec{\alpha}_\sigma = (\alpha_{1,\sigma},\alpha_{2,\sigma},...) \) representing the resource allocation strategy of species \( \sigma \). For simplicity, we assume each species has a fixed resource allocation strategy.

**Dynamic equations for a single species in a chemostat**

In a chemostat with nutrient supply \( \tilde{c}_{\text{supply}} \), dilution rate \( d \) and a single species \( \sigma \) with fixed strategy \( \vec{a}_\sigma \) and intracellular metabolite concentration \( \vec{q}_\sigma \), the cell biomass density \( m_\sigma \) and the chemostat nutrient concentrations \( \vec{c} \) are generally described by the following equations:

\[
\frac{dm_\sigma}{dt} = m_\sigma \cdot (g(\vec{c},\vec{q}_\sigma,\vec{a}_\sigma) - d),
\]

\[
\frac{d\tilde{c}}{dt} = d \cdot (\tilde{c}_{\text{supply}} - \tilde{c}) - m_\sigma / r \cdot \vec{I}(\vec{c},\vec{q}_\sigma,\vec{a}_\sigma).
\]

In considering the details of cellular metabolism, one may choose to incorporate the dynamics of intracellular metabolites that originate from nutrient import and influence cell growth. We make the assumption that the biomass concentration \( r \), e.g. protein concentration, is constant for cells under all growth conditions. Thus, an increase of total cell mass induces a corresponding increase of total cell volume. \( m_\sigma \) is the cell mass per element of volume in the chemostat, and \( r \) is the cell mass per element of volume within a cell. For a chemostat-to-cell flux of mass \( J \), the concentration of the metabolite in the chemostat decrease by \( J/V_{\text{chemostat}} \) while the concentration in cell increase by \( J/V_{\text{cell}} \). As a result, the metabolites imported into cells are enriched by a factor of \( r \), and metabolites secreted by cells are diluted by \( 1/r \). Also, all intracellular metabolites are diluted by cellular growth, which is generally a slow process compared to metabolic reactions and can be ignored in most cases. We use a function \( \tilde{f}(\vec{I}(\vec{c},\vec{q}_\sigma,\vec{a}_\sigma),\vec{q}_\sigma,\vec{a}_\sigma) \) to represent the rate of change of intracellular metabolite concentration \( \vec{q}_\sigma \):

\[
\frac{d\vec{q}_\sigma}{dt} = \tilde{f}(\vec{I}(\vec{c},\vec{q}_\sigma,\vec{a}_\sigma),\vec{q}_\sigma,\vec{a}_\sigma).
\]

Eq. (S2) represents \( p \) equations for \( p \) types of nutrients, and Eq. (S3) represents \( h \) equations for \( h \) growth-related intracellular metabolites.

**Species-specific steady state**

In the steady state of a chemostat, Eqs. (S1)-(S3) will all be equal to zero.
For intracellular metabolites, as \( f^*(-\hat{c}(\hat{\sigma}, \hat{\alpha}_\sigma), \hat{\sigma}, \hat{\alpha}_\sigma) = 0 \) as a result of Eq. (S3) being equal to zero, given an environment \( \hat{c} \), the steady state of \( \hat{\sigma} \) can be expressed as a function of \( \hat{c} \): \( \hat{\sigma} = f^{-1}(\hat{c}, \hat{\alpha}_\sigma) \).

**Growth contour (GC):** From the perspective of the environment influencing species, at each constant environment, the steady-state growth rate is fully determined by \( \hat{c} \): \( g^*(\hat{c}, \hat{\sigma}) = g(\hat{c}, \hat{f}^{-1}(\hat{c}, \hat{\alpha}_\sigma)) \). If the biomass of a species is non-zero \( (m \neq 0) \), Eq. (S1) requires \( g^* = d \). In the \( p \)-dimensional chemical space, this requirement defines a \((p-1)\)-dimensional surface, constituted by all environments \( \hat{c} \) that support an equal-to-dilution growth rate. This surface reduces to the zero-growth isoclines in contemporary niche theory when the chemical space is two-dimensional and the growth rate \( g \) solely relies on \( \hat{c} \) monotonically, but is not necessarily limited by the nutrient dimension or the form of the growth function. For convenience, we name this surface the “growth contour” (GC) for species \( \sigma \):

\[
G_{C\sigma} := \{ \hat{c} | g(\hat{c}, \hat{f}^{-1}(\hat{c}, \hat{\alpha}_\sigma), \hat{\alpha}_\sigma) = d \}.
\]

(S4)

An example of growth contours is shown in Fig 1D.

**Flux-balance curve (FB):** Eq. (S2) describes how species act on the environment. In steady state, the influx, out-flux, and consumption by species should be balanced for each nutrient, which enables calculation of the biomass density-to-dilution ratio for every \( i \):

\[
\frac{c_\text{supply} - c_i}{I_i(\hat{c}, \hat{f}^{-1}(\hat{c}, \hat{\alpha}_\sigma), \hat{\alpha}_\sigma)} = \frac{m_\sigma}{d \cdot r} \quad \text{AND} \quad c_i < c_\text{supply}.
\]

(S5)

For example, for two nutrients \( a \) and \( b \), the flux balance curve is:

\[
\frac{c_{a, \text{supply}} - c_a}{I_a(\hat{c}, \hat{f}^{-1}(\hat{c}, \hat{\alpha}_\sigma), \hat{\alpha}_\sigma)} - \frac{c_{a, \text{supply}} - c_a}{I_a(\hat{c}, \hat{f}^{-1}(\hat{c}, \hat{\alpha}_\sigma), \hat{\alpha}_\sigma)} = 0,
\]

as demonstrated in Fig 1B and Figure S1.

In chemical space, the steady-state environment \((\hat{c}_{\sigma, ss})\) with non-zero biomass of species \( \sigma \) will be located at the intersection of the growth contour and the flux-balance curve. This environment is constructed by the species \( \sigma \) via its consumption of nutrients. If \( \hat{c}_{\sigma, ss} \) exists, this species can survive in the chemostat. Otherwise, this species will be washed out by dilution even without competition from other species. For the following discussion, we only consider species that can survive when alone in a chemostat.

**Supply line (SL):** The flux-balance curve is determined by the supply condition \( \hat{c}_{\text{supply}} \). In many cases, it is helpful to derive the supply conditions that enable a species \( \sigma \) to construct a steady-state environment \( \hat{c}_{\sigma, ss} \). All possible values of \( \hat{c}_{\text{supply}} \) that can produce a given \( \hat{c}_{\sigma, ss} \), form a straight line in the space of supply concentrations, described by:

\[
SL := \{ \hat{c}_{\text{supply}} | \hat{c}_{\text{supply}} = \frac{m_\sigma}{d \cdot r} \cdot \hat{I}(\hat{c}_{\sigma, ss}, \hat{\sigma}, \hat{\alpha}_\sigma) \}
\]

with varying non-negative values of \( m_\sigma / d \). An example of a supply line is shown in Fig S1A.

**Dynamic equations for multiple species in a chemostat**
In nutrient competition models, multiple species ($\sigma = 1 \ldots n$) each with biomass density $m_\sigma$ compete for resources. They have species-specific growth rates $g(\vec{c}, \vec{q}_\sigma, \vec{a}_\sigma)$ and import rates $\vec{I}(\vec{c}, \vec{q}_\sigma, \vec{a}_\sigma)$, yet all experience the same chemical environment $\vec{c}$. Therefore, Eq. (S1) and Eq. (S3) remain the same for each species, while the rate of change of chemostat nutrient concentrations is influenced by the summed action of all species:

$$\frac{d\vec{c}}{dt} = d \cdot (\vec{c}_{\text{supply}} - \vec{c}) - \sum_{\sigma=1}^{n} m_\sigma/r \cdot \vec{I}(\vec{c}, \vec{q}_\sigma, \vec{a}_\sigma).$$  \hspace{1cm} \text{(S7)}$$

**Multiple species steady state**

Multiple species, even if each alone can survive in chemostat, do not generally coexist when competing together. For a system starting with $n$ different species, the stable steady state contains $n^*$ ($1 \leq n^* \leq n$) species with non-zero biomass. We define these $n^*$ surviving species as a stable species set $\{\sigma^*\}$, and mark the steady-state environment created by this set as $\vec{c}_{\{\sigma^*\},\text{ss}}$.

If $n > 1$, according to Eq. (S1), $\vec{c}_{\{\sigma^*\},\text{ss}}$ must be located at the common intersection of growth contours formed by every species in $\{\sigma^*\}$.

**Invasion**

Invasion is defined as introducing a small number of invaders (with biomass density $m_{\text{inv}}$) to a steady-state chemostat occupied by a set of local species. At the time of introduction, if the invader can increase in biomass ($\frac{dm_{\text{inv}}}{dt} > 0$), the invasion is successful; otherwise if the invader decreases in biomass ($\frac{dm_{\text{inv}}}{dt} < 0$), the invasion is unsuccessful. If the biomass stays constant ($\frac{dm_{\text{inv}}}{dt} = 0$), the species is neutral with respect to the local species.

In evaluating invasion by a species $\sigma$ with strategy $\vec{a}_\sigma$ of any environment $\vec{c}$, we make two assumptions:

1. The biomass of the invader is so small that it does not disturb the environment at the time of introduction.
2. There is a separation of timescales such that the concentrations of intracellular metabolites reach equilibrium instantaneously at the time of introduction of the invader, therefore Eq. (S3) is always equal to zero and $\vec{q}_\sigma = \vec{f}^{-1}(\vec{c}, \vec{a}_\sigma)$ holds.

Therefore, we define the “invasion growth rate” of a species $\sigma$ with strategy $\vec{a}_\sigma$ introduced into environment $\vec{c}$ as:

$$g_{\text{inv}}(\vec{a}|\vec{c}) = g(\vec{c}, \vec{f}^{-1}(\vec{c}, \vec{a}_\sigma), \vec{a}_\sigma).$$  \hspace{1cm} \text{(S8)}$$

**Invasion zone:** By definition, the growth contour of the invader $GC_{\text{inv}}$ divides the chemical space into two regions: an “invasion zone” that includes all environments where the invader has an invasion growth rate higher than dilution, and “no-invasion zone” where the invader has an invasion growth rate lower than dilution. If the steady-state environment constructed by local species $\vec{c}_{\{\text{local}\},\text{ss}}$ is located within the invasion zone of the invader, $g_{\text{inv}}(\vec{a}_{\text{inv}}|\vec{c}_{\{\text{local}\},\text{ss}}) > d$, therefore $\frac{dm_{\text{inv}}}{dt} > 0$ by Eq. (S1), and the invasion is successful; otherwise, if $\vec{c}_{\{\text{local}\},\text{ss}}$ is located outside of the invasion zone of the invader, $g_{\text{inv}}(\vec{a}_{\text{inv}}|\vec{c}_{\{\text{local}\},\text{ss}}) < d$, and the invasion is unsuccessful. If $\vec{c}_{\{\text{local}\},\text{ss}}$ locate exactly on the growth contour, it is neutral.

Two examples of this rule of invasion are presented in Fig 1C and 1D.
If the growth rate monotonically increases with the concentration of each nutrient, it can be proven that the invasion zone is always above the growth contour of the invader (an environment \( \hat{c}_+ \) “above” the growth contour \( G_{\text{inv}} \) is defined as \( \exists \hat{c}_0 \in G_{\text{inv}}, \text{ s. t. } c_{i,\hat{c}_0} \geq c_{i,0} \forall i \). If the growth rate is not monotonically increasing with nutrient concentrations, identifying the invasion zone requires more model-specific analysis.

**Fitness landscape**

We quantified the fitness landscapes in the chemostat via the relationship between metabolic strategies \( \alpha_\sigma \) and the invasion growth rates of an invader adopting strategy \( \alpha_\sigma \) in a given chemical environment \( \hat{c} \). Specifically,

\[
F_{\text{idn}} := g_{\text{inv}}(\hat{a}|\hat{c}). \tag{S9}
\]

Each environment \( \hat{c} \) defines a fitness landscape. A set of species \( \{\sigma^*\} \) constructs a steady-state environment \( \hat{c}_{(\sigma^*),\text{ss}} \) and a corresponding fitness landscape \( g_{\text{inv}}(\hat{a}|\hat{c}_{(\sigma^*),\text{ss}}) \). Some examples of fitness landscapes are shown in Figs 2C, 3C, 4B-C, 5D and 6D-F.

**Non-invasible/optimal/ evolutionarily stable strategies**

A set of species \( \{\sigma^*\}_{\text{opt}} \) is non-invasible, aka optimal or evolutionarily stable, if no other species can invade the steady-state environment constructed by \( \{\sigma^*\}_{\text{opt}} \):

\[
g_{\text{inv}}(\hat{a}_\sigma|\hat{c}_{(\sigma^*),\text{ss}}) < d, \forall \sigma \notin \{\sigma^*\}_{\text{opt}}. \tag{S10}
\]

Equivalently, Eq. (S10) can be expressed as “a set of species \( \{\sigma^*\}_{\text{opt}} \) construct a fitness landscape which places themselves on the top”, according to Eq. (S9).

The steady state constructed by \( \{\sigma^*\} \) is influenced by the supply \( \hat{c}_{\text{supply}} \) and the dilution rate \( d \). For different chemostat parameters, the non-invasible species set \( \{\sigma^*\}_{\text{opt}} \) may be different. In the following steps, we described a generally applicable protocol for obtaining the non-invasible strategies:

1. **Maximal growth rates and maximizing strategies**

   In a metabolic model with trade-offs in resource allocation, the maximizing resource allocation strategy \( \hat{a}_{\text{max}} \) under a given environment \( \hat{c} \) is defined as the strategy that maximizes invasion growth rate:

   \[
g_{\text{max}}(\hat{c}) := \max_{\hat{a}}(g_{\text{inv}}(\hat{a}_\sigma|\hat{c})) \]

   \[
   \hat{a}_{\text{max}}(\hat{c}) := \arg \max_{\hat{a}}(g_{\text{inv}}(\hat{a}_\sigma|\hat{c})). \tag{S11}
   \]

2. **Maximal growth contour**

   For a given dilution rate \( d \), all environments that support a maximal growth rate of \( d \) constitute the “maximal growth contour”:

   \[
   G_{\text{max}} := \{\hat{c}_0 | g_{\text{max}}(\hat{c}_0) = d\}. \tag{S12}
   \]

   \( G_{\text{max}} \) is generally formed by many species, with each species adopting the maximizing strategies \( \hat{a}_{\text{max}}(\hat{c}_0) \) corresponding to one environment \( \hat{c}_0 \) on the maximal growth contour.
\( G_{\text{max}} \) is outside of the invasion zone for any species \( \sigma \). (Otherwise, if a species \( \sigma \) could invade an environment \( \hat{c}_0 \) on \( G_{\text{max}} \), \( g_{\text{inv}}(\hat{a}_\sigma|\hat{c}_0 \in G_{\text{max}}) > d \), this would directly violate the requirement by Eqs. (S11) and (S12) that max \( g_{\text{inv}}(\hat{a}_\sigma|\hat{c}_0) = d \). Therefore, the necessary and sufficient condition for a set of species to be evolutionarily stable, is to construct a steady-state environment on the maximal growth contour:

\[
\hat{c}_{(\sigma^{*})_{\text{opt,ss}}} \in G_{\text{max}}.
\]  

(S13)

Therefore, a strategy belonging to the non-invasible set must be a maximizing strategy.

An example of maximal growth contour is shown in Figs 4D, 5B-C, and 6D

3. Non-invasible strategy

Nevertheless, adopting one of the maximizing strategies along the maximal growth contour does not guarantee that a species will satisfy Eq. (S13) and become non-invasible, as a maximizing strategy for environment \( \hat{c}_1 \) may end up constructing a different environment \( \hat{c}_2 \). To identify a non-invasible species for supply condition \( \hat{c}_{\text{supply}} \), the flux-balance condition needs to be considered, with the strategies maximized at each environment. This requirement forms a “maximal flux-balance curve” in the chemical space:

\[
FB_{\text{max}} := \{ \hat{c} | \frac{c_{\text{l, supply}} - c_{\text{l}}}{l_{\text{i}}(\hat{c}, \hat{f}^{-1}(\hat{c}, \hat{a}_{\text{max}}(\hat{c})), \hat{a}_{\text{max}}(\hat{c}))} = \frac{m}{d \cdot r} \text{ AND } c_{\text{l}} < c_{\text{l, supply}} \}.
\]  

(S14)

If the intersection of the maximal growth contour and the maximal flux-balance curve exists, it is the evolutionarily stable environment under dilution rate \( d \) and supply condition \( \hat{c}_{\text{supply}} \), \( \hat{c}_{\text{opt}} \). The maximizing strategy for this environment, \( \hat{a}_{\text{opt}} = \hat{a}_{\text{max}}(\hat{c}_{\text{opt}}) \), constructs the environment \( \hat{c}_{\text{opt}} \), and is evolutionarily stable.

4. Evolutionarily stable coexistence at the discontinuous points of the maximal growth contour

Inversely, for each environment \( \hat{c}_0 \) on the maximal growth contour, all supply conditions that enable the maximizing strategy of \( \hat{c}_0 \) to become the non-invasible strategy can be calculated from the supply line according to Eq. (S6):

\[
SL(\hat{c}_0) := \{ \hat{c}_{\text{supply}} | \hat{c}_{\text{supply}} = x \cdot \hat{I}(\hat{c}_0, \hat{f}^{-1}(\hat{c}_0, \hat{a}_{\text{max}}(\hat{c}_0)), \hat{a}_{\text{max}}(\hat{c}_0)) + \hat{c}_0 \}.
\]  

(S15)

for any non-zero value of \( x \). Some examples are shown in Fig 4D.

When there are discontinuous points on the maximal growth contour, there can be “gaps” in the nutrient supply space, where no single strategy on the maximal growth contour satisfies Eq. (S14). Under this condition, more than one strategy is required to co-create an environment on a discontinuous point of the maximal growth contour. Therefore, discontinuous points of the maximal growth contour permit evolutionarily stable coexistence, where \( \{ \sigma^{*} \}_{\text{opt}} \) contains more than one species. Two examples of such discontinuities and coexistence are shown in Fig 5 and Fig 6.

Metabolic models

Different assumptions can be made regarding the metabolic models \( \hat{f}(\hat{c}, \hat{x}, \hat{a}) \), \( g(\hat{c}, \hat{x}, \hat{a}) \), and \( \hat{I}(\hat{c}, \hat{x}, \hat{a}) \), focusing on various aspects of cellular growth. Different assumptions lead to distinct classes of metabolic models with various results. Nevertheless, our analysis schemes, including the invasion geometry, fitness landscape, and evolutionary stable strategies, are generally
applicable for various metabolic models. In this work, we used five metabolic models to
illustrate multiple aspects of the species-environment feedback:

1. Metabolic model with two essential nutrients

When two nutrients are both essential for growth, such as nitrogen and phosphorus, and both
require a substantial allocation of resources for import, the system can be abstractly modeled as
shown in Fig 4A. In this metabolic model, we assume an exact trade-off between the allocation
of limited resources to import nutrient \( a \) or nutrient \( b \). The fraction of resources allocated to
import nutrient \( a \) is represented by \( \alpha_a \), thus leaving a fraction \( \alpha_b = 1 - \alpha_a \) to import nutrient \( b \).
The import rate of nutrient \( i \) is assumed to follow the Monod equation as a function of nutrient
concentration, and is proportional to \( \alpha_i \):

\[
I_i(\vec{c}) = \alpha_i \cdot \frac{c_i}{c_i + K_i} \quad \text{for } i = a, b. \quad (S16)
\]

Import of both nutrients is required for cell growth:

\[
g(\vec{c}) = \gamma \cdot \min(I_a(\vec{c}), I_b(\vec{c})). \quad (S17)
\]

For this model, for simplicity we do not explicitly consider intracellular metabolites. Rather,
import directly determines growth.

In this model, a “species” is defined by its value of \( \alpha_a \).

Nutrient limitation can be clearly quantified in this system: if \( I_a(\vec{c}) > I_b(\vec{c}) \), the system is
limited by nutrient \( b \); if \( I_a(\vec{c}) < I_b(\vec{c}) \), the system is limited by nutrient \( a \).

A species with the following parameters was used to generate Fig S1, focusing on how supply
conditions and dilution rate influence nutrient limitation:

| \( K_a \) | \( K_b \) | \( \gamma \) | \( \alpha_a \) |
|--------|--------|--------|--------|
| 0.7    | 1.3    | 10     | 0.3    |

In Fig S1A, to demonstrate how species construct the same environment out of different supply
conditions, the chemostat dilution rate was set to \( d = 1 \), and three supply conditions were used:
\( \vec{c}_{\text{supply}} = [0.6, 0.3546] \) (purple), \( \vec{c}_{\text{supply}} = [0.8, 0.5273] \) (cyan), and \( \vec{c}_{\text{supply}} = [1,0.7] \) (blue).

In Fig S1B, to demonstrate how dilution rates may switch the limiting nutrient, we used the same
supply condition as the blue condition in Fig 1D (\( \vec{c}_{\text{supply}} = [1,0.7] \)), and three dilution rates: 0.5
(yellow), 1 (red), and 1.6 (deep red).

Species with following parameters were used to generate Fig 4B-D:

| \( K_a \) | \( K_b \) | \( \gamma \) |
|--------|--------|--------|
| 0.5    | 0.5    | 10     |

The strategy \( \alpha_a \) varies for different species. In Fig 4B, Species Blue has \( \alpha_a = 0.35 \), species Red
has \( \alpha_a = 0.65 \). In Fig 4C, we started with species Blue and species Red. We then generated the
fitness landscape for each species at the steady-state environment it constructed, then chose the
strategy that maximized invasion growth rate for this fitness landscape to generate a new species,
and iterated this process five times. The species Black has \( \alpha_a = 0.5 \).

In generating Fig 4D, we followed the protocols described in section “Non-invasible /
evolutionarily stable strategies”.

2. Metabolic model with substitutable nutrients
When two nutrients are mutually substitutable for growth, such as glucose and galactose, the system can be described by metabolic model as shown in Fig 2A. The trade-off and import functions are taken to be the same as in Model 1: *metabolic model with two essential nutrients*. However, import of the two nutrients contributes additively toward growth rate:

\[
g(\vec{c}) = \gamma \cdot (I_a(\vec{c}) + I_b(\vec{c})).
\]

(S18)

A species is defined by its value of \(\alpha_a\).

For this model, all growth contours intersect at one point. The growth contour of species \(\sigma\) satisfies the equation:

\[
\alpha_a \cdot \frac{c_a}{c_a + K_a} + (1 - \alpha_a) \cdot \frac{c_b}{c_b + K_b} = \frac{d}{\gamma}.
\]

Regardless of the value of \(\alpha_a\), the environment \([\frac{K_a}{d^{-1}}, \frac{K_b}{d^{-1}}]\) is always on the growth contour. Species with the following parameters were used to generate Fig 1C-D and Figure 2:

\[
\begin{array}{ccc}
K_a & K_b & \gamma \\
1.2 & 0.8 & 3
\end{array}
\]

The strategy \(\alpha_a\) varies for different species. In Fig 1C-D and Fig 2B-C, Species Blue has \(\alpha_a = 0.2\), species Red has \(\alpha_a = 0.6\). Supply conditions are different among the three figures: in Fig 1C, \(\vec{c}_{\text{supply}} = [0.5, 1]\); in Fig 1D, \(\vec{c}_{\text{supply}} = [1, 0.5]\); in Fig 2B, \(\vec{c}_{\text{supply}} = [1, 1]\).

All conditions in Fig 2D are the same as in Fig 2B other than that five additional species are added to the system. Their strategies are indicated by the legend at the right.

3. Metabolic model with substitutable nutrients that require assimilation

In cells, the assimilation of imported raw material, such sugars, into biomass such as proteins, takes multiple steps and enzymes and consumes a considerable amount of energy. When the resources allocated to nutrient assimilation are considered, a cell’s strategy becomes more complex. A mathematical model involving three substitutable nutrients \(a, b, c\) that need assimilation is shown in Fig 3A, with \(\alpha_{i1}\) represents the fraction of resources allocated to importing nutrient \(i\) into internal metabolite, and \(\alpha_{i2}\) represents the fraction of resources allocated to assimilate the internal \(i\) into biomass. In this model, the import rate has a similar form to the previous two models,

\[
I_i(c_i) = V \cdot \alpha_{i1} \cdot \frac{c_i}{c_i + K_i}.
\]

(S19)

The internal metabolite concentration \(c_{i,\text{internal}}\) has an influx of \(r \cdot I_i(c_i)\), meanwhile, it is diluted by cell growth at the rate \(g\). We assume all nutrients are substitutable therefore the internal pools contribute via summation to growth, and are converted into biomass at a rate \(k \cdot \alpha_{i2} \cdot c_{i,\text{internal}}\):

\[
\frac{dc_{i,\text{internal}}}{dt} = I_i(c_i) - g(c_{i,\text{internal}}) \cdot c_{i,\text{internal}} - k \cdot \alpha_{i2} \cdot c_{i,\text{internal}}.
\]

(S20)

Therefore, the mass converted into biomass per unit time per unit volume is: \(\Sigma_i(k \cdot \alpha_{i2} \cdot c_{i,\text{internal}})\), and the growth rate defined as the relative gain of total biomass \(M\) is:

\[
g(c_{\text{internal}}) = \frac{dM}{dt} = \frac{k}{r} \cdot \Sigma_i(\alpha_{i2} \cdot c_{i,\text{internal}}).
\]

(S21)

In generating Fig 3B-C, the chemostat parameters were: \(\vec{c}_{\text{supply}} = [1,1,1]\), and \(d = 1\), and the species parameters were:

| \(V\) | \(K_i\) \((i = a, b, c)\) | \(k\) |
|-------|-----------------|-----|
| 1000  | 0.5             | 11  |

The three species allocate their resources differently:

| Strategies | \(\alpha_{a1}\) | \(\alpha_{a2}\) | \(\alpha_{b1}\) | \(\alpha_{b2}\) | \(\alpha_{c1}\) | \(\alpha_{b2}\) |
To generate the Fig S3, all other parameters are the same, other than \( k = 10 \).

### 4. Metabolic model with essential nutrients that can be interconverted

If two nutrients are both essential for growth, and a cell is able to convert one nutrient into another albeit at a certain cost, as shown in Fig 5A, metabolic trade-offs involve the following four elements of the allocation strategy \( \vec{\alpha} \):

- \( \alpha_a \): Fraction of resources allocated to import nutrient \( a \).
- \( \alpha_b \): Fraction of resources allocated to import nutrient \( b \).
- \( \alpha_{ab} \): Fraction of resources allocated to convert internal \( b \) into \( a \).
- \( \alpha_{ba} \): Fraction of resources allocated to convert internal \( a \) into \( b \).

To implement trade-offs, the sum of elements of \( \vec{\alpha} = (\alpha_a, \alpha_b, \alpha_{ab}, \alpha_{ba}) \) is taken to be equal to 1.

In this metabolic model, cells internalize nutrient \( a \) and nutrient \( b \) from the chemostat to supply internal concentration of nutrients, \( c_{a,\text{internal}} \) and \( c_{b,\text{internal}} \). Meanwhile, the internal nutrients can be converted into each other. Nutrients also diffuse in and out of the cell passively with rate \( \beta \). Cell growth requires both internal nutrients, and depletes them in a fixed proportion.

In this model, the growth rate of a cell is taken to be:

\[
g(\vec{c}_{\text{internal}}) = \frac{\gamma}{\frac{K_a}{c_{a,\text{internal}}} + \frac{K_b}{c_{b,\text{internal}}}.
\]

(S22)

The net import rate, including passive diffusion, is:

\[
I_i = (\alpha_i + \beta) \cdot c_i - \beta \cdot c_{i,\text{internal}}, \quad i = a, b.
\]

(S23)

Therefore, the dynamical equations for the internal nutrients are:

\[
\frac{dc_{a,\text{internal}}}{dt} = I_a + \alpha_{ab} \cdot c_{b,\text{internal}} - \alpha_{ba} \cdot c_{a,\text{internal}} - g/K_a,
\]

(S24)

\[
\frac{dc_{b,\text{internal}}}{dt} = I_b + \alpha_{ba} \cdot c_{a,\text{internal}} - \alpha_{ab} \cdot c_{b,\text{internal}} - g/K_b.
\]

(S25)

A species is defined by its value of \( \vec{\alpha} \).

This metabolic model was used to demonstrate how to obtain locally optimal strategies and cartels, as shown in Fig 5. The parameter values used to generate the plots in Fig 5B-D were:

| \( \gamma \) | \( K_i (i = a, b) \) | \( \beta \) |
|---|---|---|
| 1 | 1 | 0.2 |

In generating Fig 5B, we searched for the maximizing strategies in the chemical space, and classified them by their non-zero values. Maximal growth contours for four dilution rates: 0.1, 0.2, 0.3, 0.4 are shown from black to gray and white colors.

In generating Fig 5C, the chemostat parameters were set to \( \vec{c}_{\text{supply}} = [0.5,1] \), and \( \beta = 0.2 \). The maximal growth contours for \( \beta = 0.2 \) were drawn, along with maximizing strategies along the contour shown as squares with colors corresponding to their sub-classes. At the discontinuous point of the maximal growth contour where the “converter” and the “importer” converge, the
distinct two maximizing strategies are denoted species *Red* and species *Blue*. In generating the competition dynamics in inset, additional to the species *Red* and species *Blue*, ten other maximizing strategies along the maximal growth contours were chosen.

5. Metabolic model with multiple energy generating steps

Cell growth is also tightly coupled with energy production. For example, with a single carbon supply as the energy source, cells employ multi-step reactions to generate multiple ATP molecules. Each step requires dedicated enzymes. The reaction intermediates, such as acetate, usually have dual roles: on the one hand, they positively contribute to ATP production via downstream reactions; on the other hand, they negatively contribute to ATP production by hampering upstream reactions. To deal with the negative effects of intermediates, cells may transport them out into the environment, generally with some metabolic cost for transporters. On the other hand, cells can also uptake such intermediates and use them as an energy source. We abstract such a process by the model shown in Fig 6A. A single chemical energy source *S* is supplied into the chemostat, which can be converted into intermediate *I* by cells. Four reactions are possible in this model, each mediated by a specific enzyme:

1. Import the resource *S* into the cell and convert it into internal intermediate *I* to extract energy (e.g. ATP). The fraction of the model enzyme budget allocated to this reaction is $\alpha_{\text{ATP1}}$. We assume the reaction is reversible, with the concentration of *S* contributing positively to the reaction rate while the concentration of *I* contributes negatively:

\[
J_1 = \alpha_{\text{ATP1}} \cdot V_1 \cdot \frac{[S] - [I]}{K_3} \cdot \frac{1}{K_1 + [S] + \frac{[I]}{K_5}}.
\]  

2. Process *I* via a downstream reaction to obtain more energy. The fraction of enzymes being allocated to this reaction is $\alpha_{\text{ATP2}}$. For this model system, it does not qualitatively influence the final results whether this reaction is product inhibited, so we neglect product inhibition. For simplicity, we assume this reaction has Michaelis–Menten form:

\[
J_2 = \alpha_{\text{ATP2}} \cdot V_2 \cdot \frac{[I]}{K_2 + [I]}.
\]  

3. Export the internal intermediate out into the environment by diffusion, with a fraction of proteins $\alpha_{\text{exp}}$ allocated to channels that allow the excretion of the intermediate into the environment to become external intermediate *I*.

\[
J_3 = \alpha_{\text{exp}} \cdot k \cdot ([I] - [I]).
\]  

4. Import the external intermediate into cells, with a fraction of proteins $\alpha_{\text{imp}}$ allocated to the import process. To reflect the property of the internal intermediate in inhibiting this transport reaction, the rate for this process is also product-inhibited:

\[
J_4 = \alpha_{\text{imp}} \cdot V_4 \cdot \frac{[I] - [I]}{K_6} \cdot \frac{1}{K_4 + [I] + \frac{[I]}{K_7}}.
\]  

Under this model, the rate of change of the concentration of the energy source *S* in the chemostat is:
\[ \frac{d[S]}{dt} = d \cdot ([S_{\text{supply}}] - [S]) - \frac{m}{r} \cdot J_1. \]  

The rate of change of the external intermediate concentration in the chemostat is:

\[ \frac{d[I_{\text{ext}}]}{dt} = d \cdot (-[I_{\text{ext}}]) - \frac{m}{r} \cdot (J_4 - J_3). \]  

The concentration of the intracellular metabolite \( I_{\text{int}} \) follows the equation:

\[ \frac{d[I_{\text{int}}]}{dt} = J_1 - J_2 - J_3 + J_4. \]  

The growth rate is a weighted sum of the ATP produced by \( J_1 \) and \( J_2 \):

\[ g = n_{\text{ATP1}} \cdot J_1 + n_{\text{ATP2}} \cdot J_2. \]  

In generating plots in Fig 6B-F, the species parameters were:

| \( V_1 \) | \( V_2 \) | \( k \) | \( V_4 \) | \( K_1 \) | \( K_2 \) | \( K_3 \) | \( K_4 \) | \( K_5 \) | \( K_6 \) | \( K_7 \) | \( n_{\text{ATP1}} \) | \( n_{\text{ATP2}} \) |
|---|---|---|---|---|---|---|---|---|---|---|---|---|
| 5 | 1 | 8 | 10 | 0.5 | 0.5 | 0.5 | 0.1 | 0.5 | 15 | 10 | 1 | 1 |

Maximal growth contours for dilution rates 0.2, 0.4, and 0.6 are shown in Fig 6B.

For Fig 6C-D, the chemostat parameters are: \( S_{\text{supply}} = 1, d = 0.4 \).

For Fig 6E-F, the chemostat parameters are: \( S_{\text{supply}} = 1.8, d = 0.6 \).

A summary of maximizing strategies in chemical space is shown in Fig S4.

**Dynamic equations for multiple species in a chain of chemostats**

Real ecosystems seldom exist in isolation. We modeled interconnected ecosystems via a chain of chemostats labeled \( k = 1 \) to \( k_{\text{tot}} \) (Fig S5A). Each chemostat exchanges medium and cells at leakage rate \( l \) with its two neighboring chemostats (if \( k = 1 \) or \( k = k_{\text{tot}} \), there is only one neighbor). The chemostat parameters \( c_{\text{supply}} \) and \( d \) are taken to be identical for all chemostats.

For the \( k \)-th chemostat, the dynamical equations for the biomass density of species \( \sigma \) and the concentration of the \( i \)-th nutrient are:

\[ \frac{dm_{\sigma,k}}{dt} = m_{\sigma,k} \cdot (g_\sigma(c_k) - d) + l \cdot (m_{\sigma,k-1} + m_{\sigma,k+1} - 2 \cdot m_{\sigma,k}), \]  

\[ \frac{dc_{i,k}}{dt} = d \cdot (c_{i,\text{supply}} - c_{i,k}) - \sum_{\sigma=1}^{n} m_{\sigma,k} \cdot l_{\sigma}(c_k) + l \cdot (c_{i,k+1} + c_{i,k-1} - 2 \cdot c_{i,k}). \]  

A steady-state solution to these equations is shown in Fig S5, using the same growth and import models and parameters as in Fig 4, with the leakage rate set to be \( l = 1 \).
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COMPETING INTERESTS

The authors declare that they have no conflict of interest.
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Figure 1 - Chemostat behavior represented in chemical space.

A. Schematic diagram of a chemostat occupied by a single microbial species. In the well-mixed medium (pale blue) of a chemostat, cells (orange ellipses) consume nutrients and grow. An influx of nutrients with fixed concentrations (blue and green arrows) is supplied at the same rate as dilution, keeping the medium volume constant.

B. Visual representation of how a species creates its own chemostat environment. Background color indicates the growth rate of cells as a function of metabolite concentrations $c_a$ and $c_b$, with the growth contour shown by the red curve. The flux-balance curve is shown in blue. Black curves with arrows show the time trajectories of chemostat simulation.

C. Example of successful invasion of species Blue by species Red. A small amount of species Red is introduced to a steady-state chemostat of species Blue. Growth contours and steady-state environments of species Blue and species Red are shown as curves and dots in the corresponding colors (colored background indicates the “invasion zone” of Red, and represents the growth rate.
of Red in this zone). The supply condition is marked by a black circle. Black curves with arrows show the time trajectory of the invasion in chemical space. Inset: time course of species biomass in the chemostat during the invasion.

D. Same as (C), except that because the supply condition (black circle) is different, the attempted invasion by species Red is unsuccessful.
Figure 2 – Metabolic models with substitutable nutrients and coexistence in a chemostat.

A. Example of a metabolic model with a trade-off in allocation of internal resources for import of two substitutable nutrients, with both nutrients contributing additively to growth. Species Red and species Blue allocate resources differently (indicated by parameter $\alpha_a$, see Methods).

B. Growth contours and the steady-state environments created by Red or Blue alone, under the supply condition shown by the black circle. Black curve with arrows shows a trajectory in chemical space. Purple dot indicates the steady-state environment created by Red and Blue together. Lower inset: time course of species biomass.

C. From top panel to bottom panel: the fitness landscape created by Red alone (for the red dot in (B)), created by Blue alone (for the blue dot in (B)), and created by both species (for the purple
Diamonds mark the locations of Red and Blue strategies and their corresponding fitness in each fitness landscape.

D. Growth contours and the species-specific steady-state environments for seven different species alone, under the supply condition shown by the black circle. Black curve with arrows shows a trajectory in chemical space. Lower inset: time course of species biomass in the chemostat.
Figure 3 - Rock-paper-scissors oscillations.

A. Example of a metabolic model with a trade-off in allocation of internal resources for import and assimilation of three substitutable nutrients, with all three nutrients contributing additively to growth. Species Red, species Blue, and species Green allocate resources differently (see Methods).
B. Growth contours (surfaces), flux-balance curves (lines), and steady-state nutrient concentrations (dots) for the three species in a three-dimensional chemical space. Black curves with arrows show the system’s limit-cycle trajectory.

C. The top panel shows the time course of species biomass in the chemostat for the limit cycle in (B). The bottom panel shows how the fitness landscape changes with time over one period of the oscillation.
Figure 4 – Multistability, chain of invasion, and non-invasive strategy.

A. Example of bistability for a metabolic model with a trade-off in allocation of resources for import of two essential nutrients, with the lower of the two import rates determining growth rate. Species Red and species Blue allocate resources differently (indicated by parameter $\alpha_a$, see Methods).

B. Bistability of the system in (A) shown in chemical space. Black curves with arrows show the trajectories of simulations with different initial conditions. Inset: the fitness landscape created by species Red or Blue alone, with colors corresponding to the steady-state environments shown by colored dots in the main panel.

C. A chain of invasion. Fitness landscape created by species with different internal resource allocation strategies (marked by diamond shapes). Starting from species Blue, the species having the highest growth rate in the fitness landscape created by the “former” species is chosen. This creates a chain of invasion from Blue to Light Green, Yellow, Deep Green, Deep Purple, all the way (intermediate processes omitted) to the species Black, which places itself on the peak of its own fitness landscape. The same procedure is also performed starting with species Red.
D. Depiction of non-invasible strategies under different supply conditions. Black-white background indicates the maximal growth rate of model in (A) under each environment, and the contour of maximal growth rates contains different strategies (represented by red-to-blue color). Growth contours of three species adopting one of the “maximizing strategies” are colored by their strategies. The supply conditions allowing these strategies to be “non-invasible” are marked by dashed black lines.
A. Metabolic model with a trade-off in allocation of internal resources for import of two nutrients plus their interconversion, with both nutrients necessary for growth.

B. Three subclasses of maximizing metabolic strategies in chemical space are indicated by background color, and circles with arrows illustrate the metabolic strategies of each subclass. The maximal growth contours for four growth rates (0.1, 0.2, 0.3, 0.4) are marked by gray colors.

C. Two maximizing strategies co-creating a non-invasible steady state. At dilution rate 0.2, the maximal growth contour and the corresponding maximizing strategies are shown as colored squares. At a discontinuous point of the growth contour, the supply lines of two distinct metabolic strategies (Red and Blue) span a gray region, where any supply condition (e.g. black...
circle) requires the two maximizing strategies to co-create the environment on the discontinuous point. Red and blue dots mark the environments created by species *Red* and species *Blue* alone, and the purple dot marks the environment co-created by *Red* and *Blue*. Black curve with arrows shows a trajectory in chemical space. Inset: competition dynamics of species *Red* and species *Blue* together with 10 other maximizing species with different strategies. 

D. The fitness landscapes for the three environments in (C) indicated by corresponding box colors. For class Green and Red, the strategy is represented by $\alpha_a$, for class Blue, the strategy is represented by $\alpha_b$. 
Species creating new nutrient dimensions and achieving evolutionarily stable coexistence.

A. Metabolic model with a single supplied nutrient $S$. Cells allocate enzymes to convert $S$ into internal intermediate $I_{\text{int}}$ and produce energy (denoted as “ATP”), export internal intermediate into the chemostat to become $I_{\text{ext}}$, import external intermediate, or consume $I_{\text{int}}$ to produce ATP. The growth rate is the sum of ATP production (see Methods).

B. Three subclasses of maximizing metabolic strategies in chemical space are indicated by background color, and circles with arrows illustrate the metabolic strategies of each subclass. The maximal growth contours for three growth rates (0.2, 0.4, 0.6) are marked by black-to-white colors.

C. At dilution rate 0.4, two maximizing strategies co-create a non-invasible environment. The maximal growth contour and the corresponding maximizing strategies are shown as colored squares. At a discontinuous point of the growth contour, the supply lines of two distinct metabolic strategies (Green and Blue) span a gray region, where any supply condition (e.g. black circle) requires two maximizing strategies to co-create the environment at the discontinuous point. Blue dot marks the environment created by species Blue alone, and the cyan dot marks the environment co-created by Blue and Green. Black curve with arrows shows a trajectory in chemical space. Inset: time course of species biomass, with species Green added to the chemostat at time 100.

D. The fitness landscapes for two environments in (C) indicated by corresponding colors of the boxes, reflecting the relationship between instantaneous growth rate and resource allocation strategy. For class Blue and Red, the strategy is represented by $\alpha_{\text{ATP}}$; for class Green the strategy is represented by $\alpha_{\text{imp}}$.

E. Same as (C), except that the dilution rate is 0.6. Inset: time course of species biomass, starting with Blue and Green, with species Red added to the chemostat at time 100.

F. Same as (D), except corresponding to the two steady-state environments shown in (E).
**SUPPLEMENTAL FIGURES**

Figure S1

A. Various supply concentrations can lead to the same steady-state chemical environment. Background color indicates the growth rate of cells as a function of nutrient concentrations $c_a$ and $c_b$, with the growth contour shown by the red curve. The supply line for the steady-state environment (purple dot) is shown as a dotted black line. Different supply concentrations ($c_{a,\text{supply}}$ and $c_{b,\text{supply}}$) along the supply line are marked by purple, cyan, and blue circles, with the corresponding flux-balance curves shown in the same colors.

B. Dilution rate can flip nutrient limitation. The external supply condition is marked by a blue circle, and the flux-balance curve for this supply is shown in the same color. Three growth contours with increasing dilution rates are shown from yellow to deep red, and the corresponding steady-state environments are shown in colored dots.
Figure S2 - Nutrient supply shifts the relationship between RNA/Protein ratio and growth rate in chemostat.

A. The relationship between ribosome abundance represented by RNA/Protein ratio (y-axis) and growth rate (x-axis) of *E. coli* cultured in chemostats from phosphorus limitation (P-limited, green open circles and dotted line) to nitrogen limitation (N-limited, blue open circles and dotted line). Starting from the P-limited condition, data for decreasing the supply concentration of nitrogen by 2, 5, and 10-fold are shown as solid dots and corresponding best-fit lines. Each measurement was repeated three times and standard errors are shown by bars.

C. Same as (B), but for phosphorus and carbon limitation instead of phosphorus and nitrogen limitation. Starting from the P-limited condition, data for decreasing the supply concentration of carbon by 2, 5, and 10-fold are shown as solid dots and corresponding best-fit lines.
Figure S3

A. The fitness of Species 1, 2, and 3 in the steady-state environment constructed by species 1, 2, and 3 for the model in Fig 3.
B. Growth contours (surfaces), flux-balance curves (lines), and steady-state nutrient concentrations (dots) for three species in a three-dimensional nutrient space, with a different conversion speed ($k = 10$) than in Fig. 3 ($k = 1$) (see Methods). Black curves with arrows show the system’s oscillatory trajectory.

C. Course of species biomass in the chemostat over a long duration for species in (B).
Figure S4

A

Chemostat 1  Chemostat 2  Chemostat h

Leakage

B

Biomass density

800

600

400

200

0

0  5  10  15  20

Chemostat number

C

Nutrient concentrations

0.22

0.2

0.19

0.18

0.17

0.16

0.15

0.14

0.13

0.12

0.11

0.1

5  10  15  20

Chemostat number

D

Strategy (z1, z2)

0.7

0.6

0.5

0.4

0.3

0.3

0.2

0.1

0

0.9

0.8

0.7

0.6

0.5

0.4

0.3

5  10  15  20

Chemostat number
With initial seeding of two species, one at each of the two ends of a chain of chemostats, a steady-state gradient of species biomass density spontaneously emerges accompanied by a gradient of nutrient concentrations, even though the supply conditions and dilution rates are identical for all the chemostats.

A. Schematic of $k_{tot}$ linked chemostats exchanging medium and cells via leakage, described by Eqs. S34-S35. The two species in the chemostats (Blue and Red) are the same bistable pair as in Fig 4B and the leakage rate is $l = 1$.

B. The species composition along 20 linked chemostats for the system in (A). Species colors correspond to those in Fig 4B, with species Blue having $\alpha_a = 0.35$ and species Red having $\alpha_a = 0.65$. The dashed black curve shows the sum of the two biomass densities. The initial condition was cell-free chemostats with a small amount of Blue added to Chemostat 1 and small amount of Red added to Chemostat 20.

C. Concentrations along the 20 chemostats for nutrient $a$ (green) and nutrient $b$ (cyan) for system in (A).

D. The fitness landscape along the chain of chemostats. The $x$-axis is the 20 linked chemostats, and the $y$-axis is the metabolic strategy represented by $\alpha_a$. Color indicates the growth rate of species adopting the given strategy in the $k$-th chemostat.
Figure S5

A

\( c_p \quad c_o \quad 1 \quad 0 \quad 0 \quad 1 \quad 1 \)

\( \alpha_a \quad \alpha_d \quad 0 \quad 0 \quad 0 \quad 0 \quad 0 \)

\( c_p \quad c_o \quad 1 \quad 0 \quad 0 \quad 1 \quad 1 \)

\( \alpha_{ac} \quad \alpha_{ba} \quad 0 \quad 0 \quad 0 \quad 0 \quad 0 \)

B

\( l_{m1} \quad 0.2 \quad 0.5 \quad 1.5 \quad 1 \quad 1 \quad 0 \quad 0 \)

\( \alpha_{ATP1} \quad \alpha_{ATP2} \quad 0 \quad 0 \quad 0 \quad 0 \quad 0 \)

\( l_{m2} \quad 0.2 \quad 0.5 \quad 1.5 \quad 1 \quad 1 \quad 0 \quad 0 \)

\( \alpha_{exp} \quad \alpha_{PP} \quad 0 \quad 0 \quad 0 \quad 0 \quad 0 \)

C

Polluter

Cleaner

Generalist

\( \alpha_{ATP1} \quad \alpha_{ATP2} \quad \alpha_{PP} \quad \alpha_{pp} \quad \alpha_{ATP1} \quad \alpha_{ATP2} \)
Figure S5 – Maximizing strategies in chemical space.

A. For each environment in the chemical space, the maximizing resource allocation strategies that maximize growth rates for the model in Fig 5A. Each strategy is represented by the four elements $[\alpha_a, \alpha_b, \alpha_{ab}, \alpha_{ba}]$, and values for each element are shown by a heatmap. Black-to-white curves are the maximal growth contours for $d = 0.1, 0.2, 0.3, 0.4$.

B. For each environment in the chemical space, the maximizing resource allocation strategies that maximize growth rates for the model in Fig 6A. Each strategy is represented by the four elements $[\alpha_{ATP1}, \alpha_{ATP2}, \alpha_{exp}, \alpha_{imp}]$, and values for each element are shown by a heatmap. Black-to-white curves are the maximal growth contours for $d = 0.2, 0.4, 0.6$.

C. Schematic representations of the three classes of maximizing strategies appearing in (B).