A graphical theory of competition on spatial resource gradients

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Resource competition is a fundamental interaction in natural communities. However little is known about competition in spatial environments where organisms are able to regulate resource distributions. Here, we analyze the competition of two consumers for two resources in a one-dimensional habitat in which the resources are supplied from opposite sides. We show that the success of an invading species crucially depends on the slope of the resource gradients shaped by the resident. Our analysis reveals that parameter combinations which lead to coexistence in a uniform environment may favor alternative stable states in a spatial system, and vice versa. Furthermore, differences in growth rate, mortality or dispersal abilities allow a consumer to coexist stationarily with - or even outcompete - a competitor with lower resource requirements. Applying our theory to a phytoplankton model, we explain shifts in the community structure that are induced by environmental changes.

INTRODUCTION

Competition for limiting resources is one of the most important species interactions in ecology and has long been considered as a major driver for shaping the spatial structure of communities and limiting species richness. A graphical theory of resource competition was advanced by MacArthur (1972), León & Tumpson (1975) and Tilman (1980, 1982) and revolves around the assumption that consumers that reduce limiting resources to the lowest level will exclude all other competitors (the $R^*$-rule). Being confirmed experimentally (Miller et al. 2005, but see also Wilson et al. 2007), this theory provides a fundamental framework for interpreting the relationship between organisms and their shared resources in a uniform environment (Grover 1997; Chase & Leibold 2003). Extensions of competition theory to spatial settings (Chesson 2000a, 2000b; Klausmeier & Tilman 2002; Amarasekare 2003) can be grouped into different classes. In spatially homogeneous environments, coexistence can be mediated by trade-offs in life history parameters (Levins & Culver 1971; Levin 1974; Kneitel & Chase 2004). In heterogeneous environments regional coexistence can be mediated by spatial segregation (Tilman 1982; Gross and Cardinale 2007), whereas local coexistence may be promoted via source-sink effects (Mouquet & Loreau 2003; Leibold et al. 2004) or positive correlations between dispersal and competitive abilities (Abrams & Wilson 2004). However, previous theory has not thoroughly explored resource competition in continuous spatially variable habitats.

In a system with resource flows, spatial heterogeneity can be created by biotic interactions, as local resource consumption can modify resource levels over larger ranges (Huston & DeAngelis 1994), possibly establishing resource gradients over the full extent of the habitat. The analysis of competition in such systems is mathematically challenging (Grover 1997), as not only the resource availability, but also the size of favorable areas (Skellam 1951), dispersal rates (Abrams & Wilson 2004) and many other factors become crucial for the survival of a population (Ryabov & Blasius 2008). Despite extensive theoretical studies on unstirred chemostats (Hsu & Waltman 1993; Smith & Waltman 1995; Wu et al. 2004), persistence and competition in streams (Speirs & Gurney 2001; Lutscher et al. 2007), vegetation patterns in water-limited habitats (Klausmeier 1999; von Hardenberg et al. 2001), well-mixed systems with a light gradient (Huisman & Weissing 1994; Diehl 2002), and non-uniform phytoplankton systems (Dutkiewicz et al. 2009; Yoshiyama et al. 2009; Ryabov et al. 2010), translating competition theory to extended systems, where species are able to shape resource distributions, still remains a great challenge.

In this study we develop a general framework for analyzing the competition between two consumers for two limiting resources in a spatially continuous habitat in which resource distributions are regulated by biotic interactions. In extension of the $R^*$-rule for uniform systems, we introduce the notion of an invasion threshold in a spatial system as the maximal resource requirement for a consumer to invade in the presence of a resident species. In this way, the outcome of competition can be interpreted graphically in the resource plane, by comparing the location of the invader’s critical resources with respect to the invasion threshold lines. We derive analytic expressions for the invasibility conditions and show that they can be related to how resources are spatially reduced as a result of resource-use by the resident.

Using this approach we find that environmental heterogeneity increases the likelihood of coexistence. Thereby we identify two forms of stationary coexistence in spa-
tially variable habitats. First, coexistence can be mediated by a trade-off in resource requirements. This is characterized by spatial segregation of the two species and we find that parameter combinations which allow coexistence in a uniform system favor alternative stable states in a spatially extended system and *vice versa.* Second, coexistence can arise from positive correlations of growth rates and resource requirements (gleaner-opportunist trade-off), characterized by a lack of spatial segregation.

To illustrate these ideas, we investigate two phytoplankton species competing for light and a nutrient in a water column. We identify two distinct regimes of bistability in the competition outcome and show that our approach provides a powerful basis for projecting the influence of environmental changes on the community composition.

**MODELLING FRAMEWORK AND ANALYSIS**

**Resource competition in uniform systems**

As a background for the following discussion, we briefly review the theory of resource competition in a uniform environment (MacArthur 1972; León and Tumpson 1975; Tilman 1980, 1982; Grover 1997; Chase & Leibold 2003). Consider two consumer species $i = 1, 2$ growing on two essential resources $N$ and $I$. The dynamics of the population density $P_i$ of each species is determined by the difference between growth $\mu_i(N, I)$ and mortality $m_i$:

$$\dot{P}_i = (\mu_i(N, I) - m_i) P_i,$$

where the growth rate in general can be written as

$$\mu_i(N, I) = \mu_{max,i} \mu \left( \frac{N}{H_{N,i}} \cdot \frac{I}{H_{I,i}} \right).$$

Here $\mu_{max,i}$ is the maximal growth rate of species $i$, $\mu$ is a monotonically increasing function of both arguments with upper bound $\lim_{N, I \to \infty} \mu(N, I) = 1$, and the half-saturation constants $H_{N,i}$ and $H_{I,i}$ define the species’ resource adaptation. In particular, this functional form includes von Liebig’s law of minimum, where the limitation of growth follows the Monod kinetics

$$\mu_i(N, I) = \mu_{max,i} \min \left\{ \frac{N}{N + H_{N,i}}, \frac{I}{I + H_{I,i}} \right\}.$$  

In a uniform environment, levels of system resources can be represented as a point in a two-dimensional resource plane (Fig. 1a). In this plane, states of balance between growth and mortality, $\mu_i(N, I) = m_i$, determine the zero net growth isoclines (ZNGIs), which for each species divide the resource plane into areas of positive and negative population net growth. Assuming eqn 3 for the growth rate, we can calculate the critical values, $I^*_{1} = H_{I,1} m_i / (\mu_{max,i} - m_i)$ and $N^*_{1} = H_{N,1} m_i / (\mu_{max,i} - m_i)$, and the ZNGI takes the form of two orthogonal lines (dashed lines in Fig. 1).

The resource concentrations in the absence of consumers define the coordinates of a resource supply point $S$. A growing population will deplete resources, shifting the system state in the direction of the consumption vector $CV$, starting at the resource supply point $S$, with the zero net growth isocline (ZNGI) and mortality $m_i$, and the ZNGI takes the form of two orthogonal lines (dashed lines in Fig. 1).

Assume that resident species 1 has attained equilib-
rium. Then invasion of a second species 2 is possible if the equilibrium resource concentrations shaped by the resident are sufficient to allow positive growth of the invader. Thus, in resource space a part of the invader’s ZNGI must be located below the equilibrium point E (Fig. [1b]). For the specific form of eqn 3, this condition is fulfilled for all critical resource values of the invader that are located in the rectangular area defined by \( I_2^* < \tilde{I}_1 \) and \( N_2^* < \tilde{N}_1 \). This line defines the invasion threshold \( T \) for species 2 (blue line in Fig. 1b).

Combining the invasion analysis for each of the two species yields the well-known outcomes of resource competition in a uniform environment (Tilman 1982). Without a trade-off in the use of the two resources (i.e. the two ZNGIs do not intersect), the species with the lowest resource requirements wins. In the presence of a trade-off, competition can lead to stable coexistence, competitive exclusion or bistability (i.e. alternative stable states with outcomes depending on initial conditions), depending on whether or not each species has a greater impact on the resource that most limits its own growth (see Fig. S1).

### Resource distributions in a spatial system

The classical approach implicitly assumes the presence of boundaries which confine the system, between which organisms and resources are uniformly distributed. In the following we aim to translate these ideas to an open system where the favorable area of a species is confined by resource availability, and where organisms and resources are not distributed uniformly. Consider a one-dimensional environment, where the population densities \( P_i(z, t) \) depend on a spatial coordinate \( z \). A spatial analog to eqn 1 can be written in the form

\[
\frac{\partial P_i}{\partial t} = (\mu_i(N, I) - m_i) P_i + D_i \frac{\partial^2 P_i}{\partial z^2},
\]

where \( D_i \) characterizes the diffusive dispersal ability of population \( i \). Again the growth rate \( \mu_i(N, I) \) follows the general form of eqn 2 but the resource concentrations \( N(z, t) \) and \( I(z, t) \) are functions of \( z \), so that the two populations are indirectly coupled through the spatial profile of their shared resources.

As a case study, we examine a two-species phytoplankton-nutrient model in an incompletely mixed water column (Radach & Maier-Reimer 1975, Jamart et al. 1977, Klausmeier & Litchman 2001, Huisman et al. 2006, Ryabov et al. 2010). In the model, eqn 4 describes the density \( P_i(z, t) \) of phytoplankton species \( i \) at depth \( z \) and time \( t \), and is complemented by an equation for the nutrient \( N(z, t) \)

\[
\frac{\partial N}{\partial t} = - \sum_{i=1}^{n} \alpha_i \mu_i(N, I) P_i + D_n \frac{\partial^2 N}{\partial z^2},
\]

and an equation for the light intensity \( I(z, t) \), which describes the absorption of light by water and phytoplankton (Kirk 1994)

\[
I(z) = I_{in} \exp \left[ -K_{bg} z - \int_0^z \sum_{i=1}^{n} k_i P_i(\xi, t) d\xi \right].
\]

In these equations, parameter \( D \) obtains the role of the turbulent diffusivity, \( \alpha_i \) is the nutrient content of a phytoplankton cell, \( I_{in} \) is the incident light intensity, \( K_{bg} \) is the water turbidity and \( k_i \) is the attenuation coefficient of phytoplankton cells. As boundary conditions we assumed impenetrable borders at the surface and at the bottom for the phytoplankton biomass and an impenetrable surface and a constant concentration \( N_B \) at the bottom for the nutrient (for model parameters see Table S1).

The model describes a situation in which the two resources, light and nutrient, are supplied from opposite sides of a spatial habitat. This is typical in the ocean where light is supplied from above and many macronutrients from below (Fig. 2a). As shown in Fig. 2b, this gives rise to characteristic resource distributions with inverse gradients, where population growth is maximal at an intermediate position at which both resources are sufficiently available (see Fig. S6 for further typical model solutions).

Within the favorable range of the population, the light and nutrient distributions in Figs. 2a and 2b can be well approximated as straight lines on a logarithmic scale. This means that resource distributions at equilibrium decay (or grow) exponentially in space

\[
\tilde{N}_i(z) \sim e^{c_{N,i} \cdot z}, \quad \tilde{I}_i(z) \sim e^{-c_{l,i} \cdot z}.
\]

This exponential dependence can be derived analytically in the limit of low mortality (see Appendix S1), and it is typical for many other ecosystems (see e.g. Fig. S5). Eqn 7 implies that a population \( i \) in monoculture is “shading” resources by a constant percentage

\[
c_{N,i} = \frac{1}{N_i} \frac{dN_i(z)}{dz}, \quad c_{l,i} = -\frac{1}{\tilde{I}_i} \frac{d\tilde{I}_i(z)}{dz}.
\]

Here, the logarithmic resource gradients \( c_{N,i} \) and \( c_{l,i} \) measure the influence of the population on the resource distribution.

Translated into the resource plane, the resource distributions of the spatially extended system can be compactly represented as a parametric curve, \( (N(z), I(z)) \), (solid line in Fig. 1c). This ‘system state curve’ (SSC) naturally extends the notion of a system state point E of a homogeneous system to a spatially continuous environment (Ryabov et al. 2010). As shown in Fig. 1c, due to the spatial coupling and source-sink effects the system state curve at equilibrium does not settle at the ZNGI. Instead, it intersects the ZNGI at two points which mark
the boundaries of positive net growth. This inner part of the system state curve (marked in bold in Fig. 1c) corresponds to the resident’s favorable range (horizontally dashed lines in Fig. 2b, c). Given the exponential dependence (eqn 7), within the favorable range the system state curve approaches a straight line with slope $\gamma_i = c_{I,i}/c_{N,i}$ in the resource plane with logarithmic axes (see Fig. 1c in the model or Fig. S4c for field data).

**Invasion analysis in a spatial system**

In analogy to competition theory in a uniform system, we may now ask whether it is possible to predict the outcome of two-species competition from knowledge about equilibrium resource distribution which has been shaped by each species alone in the absence of the other species. Assuming that we know the resident’s system state curve at equilibrium $(\tilde{N}_1(z), \tilde{I}_1(z))$, what can we say about the success of an invader (red color in Fig. 2c)?

To address this problem we define the invasion threshold $T$ in the resource plane as the set of critical resources of all invaders of small initial density, which have zero total growth in the presence of the resident (blue solid line in Fig. 1d, obtained by invasion analysis in numerical simulations with 7000 parameter combinations). In this formulation, the success of an invader depends on whether or not its ZNGI intersects with the invasion threshold (cross-hatching in Fig. 1). Thus, invasion analysis can be performed graphically by comparing the location of the invader’s critical resources with the invasion threshold.

By comparison of Figs. 1b and 1d, the invasion threshold has a different shape in a uniform or a spatially continuous system. It is difficult to predict the invasion threshold from general principles. Exact conditions can be expressed in terms of eigenvalues of a corresponding boundary value problem (Hsu & Waltman 1993; Appendix S3), which unfortunately cannot be solved in general. In the following we show that, with the assumption of exponential resource distributions (eqn 7), it is possible to calculate the invasion threshold for species with similar resource requirements.

**Calculation of the invasion threshold**

Consider first the simplest case where resident (species 1) and invader (species 2) differ only in their half saturation constants, $H_{N,i}$ and $H_{I,i}$, but are otherwise identical ($\mu_{max,1} = \mu_{max,2}$, $m_1 = m_2$, $D_1 = D_2$). As the invader density is assumed to be small, and its influence on the resource distributions can thus be neglected, the possibility of invasion depends entirely on the invader’s growth rate in the equilibrium resource distribution shaped by the resident, $\mu_{2}(\tilde{N}_1(z), \tilde{I}_1(z))$. The mathematical iden-
tity

\[ \frac{1}{H_2} e^{cz} = \frac{1}{H_1} e^{c(z - \Delta z)}, \quad \text{with} \quad \Delta z = \frac{1}{c} \ln \frac{H_2}{H_1}, \quad (9) \]

shows that division of an exponential function by different (half-saturation) constants \( H_1 \) and \( H_2 \) is equivalent to a shift \( \Delta z \) in position along the \( z \)-axis. Therefore, assuming that the growth rate follows eqn (2) and the resource distributions are exponential (eqn (7)), it is possible to express the growth rate, \( \mu_2(z) \), of the invader through that of the resident

\[ \mu_2(\tilde{N}_1(z), \tilde{I}_1(z)) = \mu_1(\tilde{N}_1(z - \Delta z N), \tilde{I}_1(z - \Delta z I)). \quad (10) \]

Here the shifts \( \Delta z N = c_{N,1}^{-1} \ln H_{N,2}/H_{N,1} \) and \( \Delta z I = -c_{I,1}^{-1} \ln H_{I,2}/H_{I,1} \) have opposite signs, since the resource gradients are inverse.

Introducing a new position \( z' = z - \Delta z N \), we find that \( \mu_2(z') = \mu_1(\tilde{N}_1(z'), \tilde{I}_1(z' + \Delta)) \), with \( \Delta = \Delta z N - \Delta z I \). The change of variables \( z \rightarrow z' \) corresponds to a shift along the \( z \)-axis (see Fig. 2c) and has no effect on the conditions for survival as long as boundary effects can be neglected. Thus, the difference in growth between invader and resident depends only on the value and sign of the single parameter \( \Delta \). If \( \Delta = 0 \), the distinct resource requirements of the species just result in a parallel translation of the growth rate profile. Since the resident species at equilibrium has zero total growth, the condition for survival as long as boundary effects can be neglected is that of the resident

\[ \mu_2(z) \text{ of the invader through the resident } \mu_1(z). \]

Here the shifts \( \Delta z N = c_{N,1}^{-1} \ln H_{N,2}/H_{N,1} \) and \( \Delta z I = -c_{I,1}^{-1} \ln H_{I,2}/H_{I,1} \) have opposite signs, since the resource gradients are inverse.

\[ \Delta \text{ describes the invasion threshold.} \]

Thus, we can formulate the invasibility criterion in the form \( \Delta < 0 \). The parameter \( \Delta \) describes the invasion threshold and can be interpreted in terms of resource gradients and relative competitive abilities

\[ \Delta = \frac{1}{c_{N,1}} \ln \frac{H_{N,2}}{H_{N,1}} + \frac{1}{c_{I,1}} \ln \frac{H_{I,2}}{H_{I,1}}. \quad (11) \]

If for instance, both species have equal resource requirements for \( N \), \( H_{N,2} = H_{N,1} \), invasion is only possible, \( \Delta < 0 \), if the invading species is a better competitor for \( I \), i.e., \( \ln(H_{I,2}/H_{I,1}) < 0 \).

If the limitation of growth follows von Liebig’s law (eqn [3]), the spatial profiles for light and nutrient limitation are shifted independently and the net shift \( \Delta = \Delta z N - \Delta z I \) measures the difference in the size of the resident and invader’s favorable habitats. This is depicted in Fig. 2c for a situation with \( \Delta z N < \Delta z I \), where the narrowing of the favorable range due to higher \( N \) requirements \( (N_2^* > N_1^*) \) is less than the widening due to better \( I \) adaptation \( (I_2^* < I_1^*) \). Therefore the net growth rate of the invader is greater than that of the resident, allowing for successful invasion (see Fig. 2c). Note that in the example shown in Figs. 1 and 2, the invader, species 2, is a good light competitor, \( I_2^* < I_1^* \), but needs higher nutrient concentrations, \( N_2^* > N_1^* \). Therefore it will have maximal production at a deeper depth (Fig. 2c).

Since the maximal growth and mortality rates are assumed to be equal for both species, the ratio of half-saturation constants equals the ratio of critical resource values (e.g., \( H_{I,2}/H_{I,1} = I_2^*/I_1^* \)), and the invasibility criterion reads

\[ \Delta = \frac{1}{c_{N,1}} \ln \frac{N_2^*}{N_1^*} + \frac{1}{c_{I,1}} \ln \frac{I_2^*}{I_1^*} < 0. \quad (12) \]

This expression has a straightforward geometrical interpretation: a species can invade the spatial habitat if its critical resources \( (N_2^*, I_2^*) \) are located below a straight line, \( \ln I - \ln I_1^* = -\gamma_1 (\ln N - \ln N_1^*) \), with a slope of absolute value \( \gamma_1 = c_{I,1}/c_{N,1} \) passing through the point \( (N_1^*, I_1^*) \) in the double-logarithmic resource plane (blue dashed line in Fig. 1a). Thus, we have derived a first-order approximation for the invasion threshold \( T \).

\[ \text{Competition outcome} \]

In a similar way it is possible to analyze the invasion potential of species 1 in a system with resident species 2. Assume without loss of generality that \( N_2^* > N_1^* \) and \( I_2^* < I_1^* \), and denote by \( \gamma_{cr} \) the slope (taken with opposite sign) of a straight line passing through the two critical resource points, \( \gamma_{cr} = -(\ln I_2^*/\ln I_1^*)/(\ln N_2^*/\ln N_1^*) \). Then, combining eqn (12) and its counterpart for species 2, we obtain four different outcomes of spatial resource competition (Fig. 3a). If the critical resource values for each species are located below the invasion threshold of its competitor, \( \gamma_1 < \gamma_{cr} \) and \( 1/\gamma_2 < 1/\gamma_{cr} \), both species can invade the monoculture of the other, leading to coexistence. In the opposite case, if the critical resource values for every species lie above the invasion threshold of its competitor, \( \gamma_1 > \gamma_{cr} \) and \( 1/\gamma_2 > 1/\gamma_{cr} \), neither of the two species can invade, leading to bistability. Finally, one species may be a superior competitor if only this species can grow in the presence of its competitor.

This analytic conclusion is confirmed by numerical simulations of the phytoplankton model eqns 3-6 in a large parameter range (Fig. 3b). The figure demonstrates that in spite of the high complexity and nonlinearity of the model, the relation between the slope of the invasion threshold lines, \( \gamma_i \), and the location of the critical resource values gives a good prediction of the competition outcome.
Differences in growth, mortality and dispersal

In general, the locations of the invasion thresholds may depend on trait differences between the resident and invader, such as maximal growth rate, mortality and dispersal rates. As shown in Appendix S3, the generalized form of the invasibility criterion reads

$$\delta_{12} = \frac{1}{e_{N,1}} \ln \frac{N_2^*}{N_1^*} + \frac{1}{e_{L,1}} \ln \frac{I_2^*}{I_1^*} < \Delta_{12}$$

where $\Delta_{12}$ is a complicated function of the trait values and resource gradients, but does not depend on the critical resource levels. If all trait differences vanish, we obtain $\Delta_{12} = 0$ and recover the previous result, eqn 12. The geometrical representation of the new term $\Delta_{12}$ is a shift of the invasion threshold line toward larger ($\Delta_{12} > 0$) or smaller ($\Delta_{12} < 0$) resource requirements in the normal direction by $\delta_{12} = \Delta_{12} e_{N,1} e_{L,1}/\sqrt{c_{N,1}^2 + c_{L,1}^2}$. Thus, the slope of the invasion threshold lines remain unchanged, but their location is shifted. Fig. 4a shows an example with a positive shift of the invasion threshold, where species 2 can invade the system despite the fact that it has higher requirements for both resources than the resident species 1.

For an illustration of how specific trait values can change the sign of $\delta_{12}$, consider the particular case that species 2 has higher values of both growth rate, $\mu_{max,2} = \beta \mu_{max,1}$, and mortality, $m_2 = \beta m_1$, with $\beta > 1$. Since the critical resource values $N_2^* > N_1^*$ and $I_2^* > I_1^*$ are independent of $\beta$, this scaling would not affect the competition outcome in a uniform environment and species
In the resource plane this effectively leads to a counter-clockwise rotation of the invasion threshold line, and graphical analysis reveals that by increasing $N_B$ the community composition shifts from dominance of the best nutrient competitor, through a regime of coexistence, to the prevalence of the best light competitor (Fig. S3). This result might explain observed negative correlations of the abundance of high light-adapted species with nutrient concentrations (Johnson et al. 2006).

Similarly we can study the influence of consumption rates. In a well-mixed water column the slope of consumption vectors is given by $\Gamma_i = (k_i z_B)/\alpha_i$ (Huisman & Weissing, 1994; Diehl 2002). Thus, the relative resource consumption scales with the ratio of the light attenuation coefficient to the cell nutrient content, $k_i/\alpha_i$.

In our model, according to Fig. 3, two species coexist if $\gamma_1 < \gamma_{cr} < \gamma_2$. Since the ratio of resource gradients $\gamma_1$ also grows with $k_i/\alpha_i$ (see Fig. S2a and eqn S6), we observe coexistence if the best nutrient competitor (here species 1) has a small $k_1/\alpha_1$ ratio and species 2 a large $k_2/\alpha_2$ ratio. The reverse situation (large $k_1/\alpha_1$ and small $k_2/\alpha_2$) leads to bistability (Fig. S3a). Thus, we obtain a rule which is diametrically opposed to that for uniform systems: coexistence arises if each species reduces its least limiting resource more strongly in relation to the other, whereas stronger reduction of the most limiting resource favors bistability (compare Figs. S3 and S1).

Fig. 5a shows the competition outcome in dependence of the critical resources of species 2 for the case that $k_1/\alpha_1 > k_2/\alpha_2$, i.e. species 1 shades relatively more light than species 2. In accordance with our rule, if species 2, which has a smaller light attenuation coefficient, is more strongly limited by light ($I_2^* > I_1^*$), it may coexist with species 1 (upper blue area), whereas stronger nutrient limitation of species 2 ($N_2^* > N_1^*$) can lead to bistability (white area). However, if the difference in nutrient limitation is sufficiently large, then the two species can again coexist (lower blue area, Appendix S2). The model further exhibits a regime of bistability between coexistence and single-species dominance (gray area), where the invasion of species 1 does not exclude species 2, but species 2 cannot invade if species 1 has already established (see also the vertical profiles in Fig. S6). If the two species differ in their maximal growth and mortality rates, $\beta > 1$ (Fig. 5b), all bifurcation lines are shifted towards larger critical resource values. Then, if species 2 is parameterized to have higher resource requirements (i.e., erial resource levels located above the black dashed lines in Fig. 5b), it can have all possible competition outcomes, even without a trade-off in resource requirements. The corresponding spatial profiles in Fig. S6 reveal two fundamentally different types of coexistence in a spatial system, mediated either by a trade-off in resource requirements (characterized by spatial segregation of the two species) or by a gleaner-opportunist trade-off (lack

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**Application to a phytoplankton model**

Applying this approach to a phytoplankton community, the slopes $\gamma_i$ of the invasion lines can be obtained by numerical simulations (Fig. S2), analytic estimations (Appendix S1), or in field enclosure experiments (Jäger et al. 2008). Tracing the dependence of $\gamma_i$ as a function of system parameters allows us to project shifts in the species composition. Consider, for instance, the effects of increasing the nutrient resource concentration at the bottom of the water column, $N_B$. More nutrients at the bottom give rise to a larger biomass of the resident species, which in turn yields a steeper nutrient decay within the production layer, i.e. a reduction of $\gamma_i$ (see Fig. S2c).
of spatial segregation).

**DISCUSSION**

The extension of resource competition theory to spatially variable environments remains a major challenge for community ecology. Usual approaches to include environmental heterogeneity consider a range of resource supply points, assuming that the system evolves independently for each supply point (Tilman 1982), while other studies have concentrated on patch occupancy models (Levins 1979) or metacommunity models of discrete patches that are connected by dispersal (Levin 1974; Mouquet & Loreau 2003; Abrams & Wilson 2004; Gross & Cardinale 2007). However, links between local patches might not only affect the dispersal of species, but also the spatial flow of matter and resources, leading to metacommunity dynamics (Loreau et al. 2003) which require a distinct theoretical approach (Huston & DeAngelis 1994, Smith & Waltman 1995, Wu et al. 2004).

Our study takes a significant step in this direction by exploring competition in a spatially continuous environment where species interact indirectly by modifying resource availability beyond their local neighborhood. We study the critical resource requirements of a successful invader in such a system and find that the invasion threshold can be approximated by a straight line on a double-logarithmic scale, with a slope that is determined by the ratio of logarithmic resource gradients. Thus, the likelihood of invasion in a two-species community can be understood from the analysis of equilibrium resource distributions in a monoculture of the resident.

Using this approach, we clarify the bases of coexistence in spatially variable habitats for a model that has widespread and important applications (see Table 1). Thereby, we synthesize two previous sets of results concerning coexistence: i) uniform habitat theory, showing that coexistence can arise from trade-offs in the use of two resources (Tilman 1980, 1982), and ii) spatially variable theory showing that coexistence can arise from a growth-based trade-off (Hsu & Waltman 1993; Smith & Waltman 1995; Wu et al. 2004). We show that in a spatially variable habitat with two resources, both forms of coexistence can occur.

Assuming that two species trade off in their resource adaptation, in a uniform system both species can coexist if each species mostly consumes its most limiting resource (León & Tumpson 1975). In contrast, in a spatial environment the species coexist if they each, in monoculture, create a resource distribution with a relatively smaller gradient \((c_1\) or \(c_N\)) of their most limiting resource. In other words, invasion is possible if the resident species does not “shade” its most limiting resource too much. As a consequence, parameter combinations which allow coexistence in a uniform system can lead to alternative stable states in a spatially extended system and *vice versa*.

These striking differences between resource competition in uniform and spatially variable systems can be explained by the observation that in a uniform environment, resources are equally available everywhere, whereas in a spatially extended system consumers compete for locally available resources. Thus in a uniform system, a species can suppress invasions by reducing those resources for which it is the best competitor. In a spatial setting the same strategy may not work because a local reduction of vital resources does not exclude invasions elsewhere in the habitat. Consider for instance the situation shown in Fig. 2c. Here the resident, species 1, is the best nutrient competitor, so that in a uniform environment invasions are prevented if species 1 reduces nutrients more strongly than light (i.e. a small value of \(k_1/\alpha_1\), or \(\Gamma_1\), in Fig. S1). In a spatial situation, however, this strategy (small value of \(k_1/\alpha_1\)) will promote invasion of species 2 (see Fig. 3), because species 1 can only shade nutrients vertically above its favorable range, whereas species 2 invades at a greater depth. Therefore, to prevent invasions, species 1 should reduce light as much as possible (large value of \(k_1/\alpha_1\)) to deteriorate growth conditions in the deep layers.

Applying our theory to a spatial phytoplankton model, we identify two distinct regions of bistability, of either alternative states for each species or bistability between coexistence and a monoculture. Bistability has been theoretically described in phytoplankton communities with differently mixed layers (Yoshiyama et al. 2009, Ryabov et al. 2010). Here we identify bistability in a system with uniform diffusivity, however, unlike in well-mixed systems, the range of bistability is smaller than that of coexistence (see Fig. 5).

Finally, in a spatial system the slope of the invasion threshold changes with the biomass of the resident (see Appendix S1), because species with higher density have a stronger influence on resource distributions. In this way, competitive interactions are intricately linked to productivity, which potentially can give rise to novel causal relations between productivity and diversity (Gross & Cardinale, 2007; Hillebrand & Matthiessen, 2009).

**Extensions of the theory**

An essential part of our analysis is based on the assumption of exponential decay of resources within the favorable range of the resident (eqn 7). This assumption makes spatial invasion analysis analytically tractable, as it allows us to express the growth rate of an invader by that of a resident species, thereby circumventing the complicated problem of actually having to calculate spatial density profiles. Exponential resource distributions correlate with field data and can be observed in aquatic systems (Figs. 2a and S4; Kirk 1994; Karl & Letelier
Spatial system with resource gradient equally likely as coexistence, each species mostly consumes its least limiting resource or resource trade-off essential difference in resource requirements has larger not possible its most limiting resource less likely than coexistence, location depends on differences in curved shape, defined by gradients and resource availability, species with higher resource requirements (opportunist) has larger \( \mu_{max} \), or smaller \( m \) or \( D \) \( \rightarrow \) spatial profiles with separation species with higher resource requirements (opportunist) has larger \( \mu_{max} \), or smaller \( m \) or \( D \) \( \rightarrow \) spatial profiles without separation.

| Competition outcome | Uniform system | Spatial system with resource gradient |
|---------------------|----------------|--------------------------------------|
| Invasion threshold  | simple form defined by resource values, on the ZNGI | curved shape, defined by gradients and resource availability, location depends on differences in \( \mu_{max}, m, \) or \( D \) |
| Coexistence with resource trade-off | each species mostly consumes its most limiting resource | each species mostly consumes its least limiting resource or essential difference in resource requirements |
| Coexistence without resource trade-off | not possible | species with higher resource requirements (opportunist) |
| Bistability | equally likely as coexistence, each species mostly consumes its least limiting resource | less likely than coexistence, each species mostly consumes its most limiting resource, possible also without resource trade-off |

Table I: Comparison of competition outcome in uniform and spatially extended systems with inverse resource gradients

With increasing distance from the favorable range one should expect deviations from exponential profiles (Fig. 2b). Invasion thresholds then become curved (Fig. 1d) and our analytic theory only provides a first-order approximation for competition between biologically closely related species (but note that our general approach of investigating invasion thresholds does not have this limitation). In this sense our theory complements the approach developed by Yoshiyama et al. (2009) for competition between sufficiently different species.

For simplicity we assumed that the favorable ranges of both species are far from the system borders, so that boundary effects on the species survival are negligible. A general approach should include both forms of the invasion threshold, presented in Fig. 1b and 1d, as limit cases, where the range of positive growth is confined either by the boundaries of a well mixed system or by the resource gradients. Our numerical simulations in the phytoplankton model (not shown) demonstrate that alternative stable states replace regimes of coexistence or vice versa, as the favorable range moves from the interior of the water column to the surface.

The analysis can be extended to take other system parameters into account. For example, in Appendix S4 we show how sinking or floating can effectively be interpreted as a change of mortality rates, thus also influencing the position of invasion thresholds. To simplify settings we considered competition between two species under equilibrium conditions. The approach could be generalized to include temporal changes in habitat conditions. Resource fluctuations shift SSCs together with invasion thresholds and can provide different time niches for \( r \) and \( K \) strategists (Grover 1990, 1991). A further interesting perspective would be to consider additional species. The success of a third competitor can be analyzed by including another invasion threshold in the resource plane. However, analysis of three-species competition is more intricate because it involves the study of invasion thresholds for three possible pairs of resident species. In preliminary investigations we have found that the combination of gleaner-opportunist and resource limitation trade-offs may promote stationary coexistence of three species on two resources.

The method developed in this manuscript is independent of the nature of the abiotic resources involved. Situations with spatial profiles of two inverse chemical resource gradients arise naturally whenever resources are provided from two opposite sides, such as interfaces between different environments (D’Hondt et al. 2004). Moreover, our theory can easily be extended to include spatial gradients of other vital factors, independently of whether or not they are actually influenced or consumed by the population. In fact, such a situation is obtained as a special case of our phytoplankton model by setting the light attenuation coefficient \( k_i = 0 \), so that light intensity decays exponentially with depth in the water column, independently of the biomass distributions. Similar spatial variation of growth factors or stress agents is abundant along environmental gradients and in transition zones between different habitat types. Thus, the general approach that we develop should apply to a wide spectrum of systems, ranging from flow reactors, stream ecosystems and transport-limited vegetation systems to marine sediments and many others.

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SUPPORTING INFORMATION

S1. Calculation of the logarithmic resource gradients

In this section we provide analytic estimates for the light and nutrient gradients resulting from a single species population in the phytoplankton model, eqns 3–6. We calculate resource distributions in the production layer, defined by the critical depths $z_N^*$ and $z_I^*$ where either the nutrient or the light intensity reaches a critical value, $N(z_N^*) = N^*$ and $I(z_I^*) = I^*$ (see dashed lines in Figs. 2b and c). We further assume small mortality $m \ll \mu_{\text{max}}$, so that the biomass can diffuse from the favorable layer without essential losses and the phytoplankton density has only small variation within the favorable layer, i.e. $P(z) \approx P_0$ for $z_N^* \leq z \leq z_I^*$.

From eqn 6 within the favorable range the logarithmic gradient of the light intensity equals

$$c_I = -\frac{d \ln I(z)}{dz} = K_{bg} + kP_0,$$  \hfill (S1)

and the light distribution can be written

$$I(z) = I^* e^{c_I(z_I^* - z)}.$$ \hfill (S2)

Regarding the nutrient profile we assume that for small mortality the critical nutrient concentrations are small, $N^* \ll H_N$, so that the growth rate close to the critical point $N \approx N^*$ can be linearized $\mu_N(N) \approx \mu_{\text{max}} \frac{N}{H_N}$. Substituting this expression into eqn 5 we obtain at equilibrium

$$-\alpha \mu_{\text{max}} \frac{N}{H_N} P_0 + D \frac{dN}{dz^2} = 0.$$  

A solution to this equation, that is monotonically increasing with depth, is given by

$$N(z) = N^* e^{c_N(z - z_N^*)},$$  \hfill (S3)

with the logarithmic gradient

$$c_N = \frac{d \ln N(z)}{dz} = \sqrt{\frac{\alpha \mu_{\text{max}} P_0}{DH_N}}.$$  \hfill (S4)

To complete our calculation the phytoplankton density in the production layer is estimated as $P_0 \sim \frac{B}{w}$ where $B$ is the total biomass and the width of the production layer $w = |z_N^* - z_I^*|$ only weakly depends on the model parameters (Beckmann and Hense 2007). If the biomass maximum is located far from the surface, the value of $B$ can be estimated in two limiting cases (Klausmeier and Litchman 2001)

$$B = \begin{cases} \frac{\ln(I_{in}/I^*)}{k} & \text{if } P_0 k \gg K_{bg} \\ \frac{D(N_B - N^*)}{amZ_M} & \text{if } P_0 k \ll K_{bg} \end{cases}$$  \hfill (S5)

where $Z_M = Z_B - \ln(I_{in}/I^*)/K_{bg}$.
These equations can be further simplified because typically \( N_B \gg N^* \). Substituting eqns S5 into eqns S1 and S4, we obtain the ratio of resource gradients \( \gamma = c_1/c_N \) (see also Fig. S2).

\[
\gamma = \begin{cases} 
  \frac{K_{bg} + \ln(I_{in}/I^*)}{w} & \text{if } P_0k \gg K_{bg} \\
  \sqrt{\frac{\alpha \mu_{max} \ln(I_{in}/I^*)}{k \DH_N}} & \text{if } P_0k \ll K_{bg}. 
\end{cases}
\]  

(S6)

**S2. Bifurcation lines in Fig. 5**

In this section we derive analytic estimates for combinations of critical resource values that allow mutual invasibility of a fixed “reference” species 1 with a variable “test” species 2 (bifurcation lines in Fig. 5a main text). Both species differ in the values of consumption rates, \( k_i \) and \( \alpha_i \), have identical maximal growth rate \( \mu_{max} \) and mortality \( m \), but while the critical resource values \( N^*_1 \) and \( I^*_1 \) of the reference species are fixed, \( N^*_2 \) and \( I^*_2 \) are taken from a wide range.

The critical resource values of all test species that can invade in the presence of the reference species are determined by the invasibility criterion, eqn 12

\[
\frac{1}{c_{N,1}} \ln \frac{N^*_2}{N^*_1} + \frac{1}{c_{I,1}} \ln I^*_2 < 0.
\]  

(S7)

The border of this region is shown as a green dashed line in Fig. 5a. Similar, we obtain the critical resource values of the test species, which allow the invasion of the reference species when the test species has reached equilibrium

\[
\frac{1}{c_{N,2}} \ln \frac{N^*_1}{N^*_2} + \frac{1}{c_{I,2}} \ln I^*_1 < 0.
\]  

(S8)

In this case however, to plot the bifurcation line, we need to consider the dependence of \( c_{N,2} \) and \( c_{I,2} \) on the critical resource values, \( N^*_1 \) and \( I^*_1 \), of the test species (see. eqns S1, S4 and S5). For these aims we pick first one special “hybrid” test species, that has the same critical resource values \( N^*_1 \) and \( I^*_1 \) as the reference species, but consumption rates, \( \alpha_2 \) and \( k_2 \). We denote the logarithmic resources gradients of this species as \( c'_{N,2} \) and \( c'_{I,2} \).

To proceed we assume that the critical resource values are small: assuming \( N^*_1 \ll N_B \) we approximate \( N_B - N^*_2 \approx N_B \), and assuming \( I^*_1 \ll I_{in} \) we obtain

\[
\ln I_{in} \approx \ln I_{in}^* - O \left( \frac{I^*_2 - I^*_1}{I_{in}} \right).
\]

Thus, in both limits, \( K_{bg} \gg P_0k \) and \( K_{bg} \ll P_0k \), in eqn S5 the total biomass \( B \), and from eqn S1 also the value of \( c_{I,2} \), is independent of the critical resource values \( N^*_2 \), \( I^*_2 \) of the test species. This means that the logarithmic light gradients are identical for all test species

\[
c_{I,2} = c'_{I,2}.
\]
In a similar way we estimate the logarithmic nutrient gradients, but from eqn S4 the value of \( c_{N,2} \) depends on the half-saturation constant of the test species, \( c_{N,2} \sim \sqrt{1/H_{N,2}} \). Then the logarithmic nutrient gradient of an arbitrary test species can be expressed in terms of the logarithmic nutrient gradient of the “hybrid” species

\[
c_{N,2} = c_{N,2}' \sqrt{H_{N,1}/H_{N,2}},
\]

and using the identity \( H_{N,1}/H_{N,2} = N_1^*/N_2^* \) we obtain

\[
c_{N,2} = c_{N,2}' \sqrt{N_1^*/N_2^*}.
\]

Therefore, inequality S8 takes the form

\[
\frac{1}{c_{N,2}} \sqrt{\frac{N_2^*}{N_1^*}} \ln \frac{N_2^*}{N_1^*} + \frac{1}{c_{I,2}'} \ln \frac{I_2^*}{I_1^*} > 0,
\]

(S9)

where the parameters \( c_{N,2}' \) and \( c_{I,2}' \) are calculated for the “hybrid” species having the critical resource values, \( N_1^* \) and \( I_1^* \), and the consumption rates, \( k_2 \) and \( \alpha_2 \). Note that this expression holds in both limits when \( K_{bg} \gg P_0 k \) and when \( K_{bg} \ll P_0 k \), implying that it might provide a good approximation in general. The border of this region is shown as red dashed line in Fig. 5a.

S3. Invasion thresholds and principal eigenvalues

The dynamics of a spatially extended population \( P(z,t) \) is determined by a reaction-diffusion equation, eqn 4, with certain boundary conditions. The solution to this equation can be presented in the form

\[
P(z,t) = \sum_{n=1}^{\infty} e^{\lambda_n t} \psi_n(z),
\]

(S10)

where \( \psi_n \) and \( \lambda_n \) are the eigenfunctions and eigenvalues of the equation

\[
(\mu(z) - m - \lambda_n)\psi_n + D \frac{d^2\psi(z)}{dz^2} = 0
\]

(S11)

with the same boundary conditions for the functions \( \psi_n(z) \). The eigenvalue \( \lambda^* \) with the largest real part, the so-called principal eigenvalue, is of special interest for determining the dynamics of \( P(z,t) \) because the term with this eigenvalue dominates the sum S10 at large times (see e.g., Ryabov & Blasius 2008). The sign of the real part of the principal eigenvalue determines if the population will grow, and so is able to persist \( \lambda^* > 0 \), or if it declines exponentially on the whole habitat, \( \lambda^* < 0 \). Note, that \( \lambda^* \) depends on the spatial resource profiles and therefore may change in time with depletion of resources by a growing population. If a species has attained equilibrium, it has zero population net growth and its principal eigenvalue equals zero, \( \lambda^* = 0 \).

The principal eigenvalue can be used for invasion analysis in a spatially extended system. We can define a family of invaders which for small initial density in the presence of the resident species have zero net growth, and thus a zero principal eigenvalue \( \lambda^*_{inv} = 0 \). In the space of the invader’s critical resource values, this family defines the invasion threshold (in general, a hypersurface), separating successful and unsuccessful invaders. In our setting, and given that resident and invader have identical maximal growth rate, \( \mu_{max,2} = \mu_{max,1} \), and mortality, \( m_2 = m_1 \), this corresponds to the condition \( \Delta = c_{N,1}^{-1} \ln H_{N,2}/H_{N,1} + c_{I,1}^{-1} \ln H_{I,2}/H_{I,1} = 0 \) (see eqn 11).
Now consider an invader which has the same half saturation constants as the resident species, but different maximal growth rate, $\mu_{\text{max},2}$, mortality, $m_2$, or dispersal ability, $D_2$. It can be shown (Stakgold, 2000) that the eigenvalue spectrum, and in particular the principal eigenvalue, of this species shifts in the positive direction with an increase of the growth rate difference $\mu_{\text{max},2} - \mu_{\text{max},1}$, and in the negative direction with an increase in the difference of destructive factors, such as dispersal rates $D_2 - D_1$ or mortalities $m_2 - m_1$. Thus, in general, the invader’s principal eigenvalue will be positive or negative, even though the constant remains $\Delta = 0$ (since the half saturation constants remain unchanged). Assume without loss of generality that the invader has a positive total growth rate, $\mu_{\text{inv}} > 0$. This means that it should be possible to invade the system with even higher resource requirements. Thus, there should be a ‘border-line’ invader with the same values of $\mu_{\text{max},2}$, $m_2$, and $D_2$, but larger half saturation constants so that its principal eigenvalue equals zero, $\lambda_{\text{inv}}^* = 0$. As the resource requirements of this second invader are higher, its $\Delta$ value is positive. This value, which we denote by $\Delta_{12}^*$, defines the difference in the favorable ranges (according to eqn 11) between this species and the resident. Thus all species with maximal growth rate $\mu_{\text{max},2}$, dispersal rate $D_2$ and mortality $m_2$, and whose half-saturation constants satisfy the inequality

$$\frac{1}{c_{N,1}} \ln \frac{H_{N,2}}{H_{N,1}} + \frac{1}{c_{I,1}} \ln \frac{H_{I,2}}{H_{I,1}} < \Delta_{12}^* \tag{S12}$$

can invade the system. As the boundary problem can not be solved in general, we cannot calculate the value of $\Delta_{12}^*$ but we can make some predictions about its sign. Similar to $\lambda_{\text{inv}}^*$, this value increases with $(\mu_{\text{max},2} - \mu_{\text{max},1})$, decreases with $m_2 - m_1$ and $D_2 - D_1$, equals zero when all these differences vanish, and changes sign together with $\lambda_{\text{inv}}^*$.

Assuming Monod limitation of growth by two essential resources (eqn 3) we can rewrite eqn S12 in terms of the critical resources values

$$\frac{1}{c_{N,1}} \ln \frac{N_2^*}{N_1^*} + \frac{1}{c_{I,1}} \ln \frac{I_2^*}{I_1^*} < \Delta_{12} \tag{S13},$$

where

$$\Delta_{12} = \Delta_{12}^* + \left( \frac{1}{c_{N,1}} + \frac{1}{c_{I,1}} \right) \ln \frac{\mu_{\text{max},1} - m_1 m_2}{(\mu_{\text{max},2} - m_2) m_1}. \tag{S14}$$

Eqns. S13 and S14 define the invasion threshold for the invader, species 2, as a straight line with slope $\gamma_1 = c_{N,1}/c_{I,1}$ in double logarithmic resource space. The location of the invasion threshold depends in a complex way on the resource gradients and on the differences in maximal growth rate, $\mu_{\text{max},i}$, mortality, $m_i$, or dispersal rates, $D_i$. In the resource plane, the invasion threshold can be located above ($\Delta_{12} > 0$) or below ($\Delta_{12} < 0$) the critical resource values of the resident, $(N_I^*, I_I^*)$, see Fig 4a.

Consider a special case when the growth and mortality rates of species 2 are rescaled, $\mu_{\text{max},2} = \beta \mu_{\text{max},1}$ and $m_2 = \beta m_1$. Then the second term in eqn [S14] vanishes, however $\Delta_{12}^*$ can be nonzero because both $\mu_{\text{max},2} - \mu_{\text{max},1}$ and $m_2 - m_1$ are positive. Substituting $\mu_{\text{max},2}$ and $m_2$ into eqn 4 and dividing it by $\beta$, we obtain that in equilibrium the distribution of species 2 should satisfy the equation

$$\mu_1(z) P_2 - m_1 P_2 + \frac{D_1}{\beta} \frac{\partial^2 P_2}{\partial z^2} = 0 .$$

Therefore these changes can be interpreted as a reduction of the diffusivity $D_2 = D_1/\beta$. Since $\Delta_{12}^*$ decreases with $(D_2 - D_1)$ the shift of the invasibility threshold is positive ($\Delta_{12}^* > 0$) if $\beta > 1$ and negative ($\Delta_{12}^* < 0$) if $\beta < 1$. 

S4. Extension to systems with sinking or advection

The dynamics of a population living in a unidirectional flow (e.g. sinking, floating, etc.) can be described by the reaction-diffusion-advection equation (Murray 2002, Ryabov & Blasius 2008)

\[
\frac{\partial P}{\partial t} = (\mu(z) - m)P - v \frac{\partial P}{\partial z} + D \frac{\partial^2 P}{\partial z^2},
\]

(S15)

where \( v \) is the flow or sinking velocity, which is positive when the flow is directed towards larger \( z \) values. Substituting \( P = \tilde{P} \exp \left( \frac{vz}{2D} \right) \) into eqn S15, we obtain

\[
\frac{\partial \tilde{P}}{\partial t} = \mu \tilde{P} - \left( m + \frac{v^2}{4D} \right) \tilde{P} + D \frac{\partial^2 \tilde{P}}{\partial z^2}.
\]

(S16)

Note that \( \partial_t \tilde{P} \) has the same sign as \( \partial_t P \), so that both functions grow and decline simultaneously. Furthermore, they have the same boundary conditions \( \tilde{P}(0) = \tilde{P}(Z_B) = 0 \). Eqn. S16 describes a population \( \tilde{P}(z,t) \) in a system without flow but with an effective mortality

\[
m' = m + \frac{v^2}{4D}.
\]

(S17)

Therefore, the presence of an advective flow which washes out a population from a favorable range can be interpreted as an effective increase of the mortality by the value \( v^2/4D \).

This higher mortality entails an increase of resource requirements. Assuming eqn 3 for the growth rate, we obtain new critical resource values

\[
I_i^{*'} = H_{I,i} \frac{m'}{\mu_{\text{max},i} - m_i}, \quad N_i^{*'} = H_{N,i} \frac{m'}{\mu_{\text{max},i} - m_i}.
\]

(S18)

Since \( m' > m \), the new critical resources values \( I^{*'} \) and \( N^{*'} \) in the presence of a positive flow are larger than in a system without flow. This, in turn shifts the zero net growth isoclines and invasion thresholds towards higher resource values in the resource plane.
Numerical scheme. The model was integrated using a backward difference method, based on the finite volume scheme (Pham Thi et al. 2005). For the numerical solution we have discretized all variables on a grid which consisted of 400 points. Diffusion terms were approximated by a second order central discretization scheme and integration was made via the trapezoidal rule. The resulting system of ordinary differential equations was solved by the CVODE package (http://www.netlib.org/ode). Further we verified that the results remain unchanged if we double the number of points in the grid. See Ryabov et al. (2010) for further details on numerical procedures.

Boundary conditions. As boundary conditions we assumed impenetrable borders at the surface and at the bottom for the phytoplankton biomass and an impenetrable surface and a constant concentration $N_B$ at the bottom for the nutrient.

Initial conditions. We used a linear initial nutrient distribution changing from 0 at the surface to $N_B$ at the bottom. For both phytoplankton species we assigned a uniform distribution of small initial density. We also investigated the influence of different initial conditions, however we did not find any new solutions beside the ones described.

To perform the invasion analysis, in the numerics the growth of the invader species was suppressed during the first 10000 simulation days, to make sure that an equilibrium distribution of the resident species was established. Then we simulated the system for a duration of further 40000 days to obtain the final competition outcome. To test for bistability, this simulation was repeated by a second simulation in which the roles of invader and resident were exchanged.

Parameter values. Default species and model parameters are listed in Table S1.

Figure 1d. Calculation of the invasion threshold is based on 7000 simulations of invasion by species with different half saturation constants $H_N$ and $H_I$. However residents and invaders have the same $\mu_{\text{max}}$, $m$, and $D$, so that the invasion threshold is not shifted with respect to the resident ($N^*, I^*$) values.

Figure 2. Half-saturation constants of the invader, $H_N^{\text{inv}} = 8 \mu\text{mol photons m}^{-2}$ and $H_N^{\text{inv}} = 0.09 \text{mmol nutrient m}^{-3}$.

Figure 3b. The figure shows the results of more than 1000 simulations where parameter values of environmental conditions ($I_{in}$, $K_{bg}$, $D$, $N_B$) and values of consumption rates, $k_i$ and $\alpha_i$, were randomly chosen from a uniform distribution in a wide range (see Table S1). Specific parameters, which do not vary, correspond to species 1 and 2 in Table S1. In particular, the half-saturation constants, maximal growth and mortality rates were fixed, so that the critical values, ($N^*_i$, $I^*_i$), and $\gamma_{cr}$ remain constant. To determine $\gamma_1$ and $\gamma_2$ numerically we performed simulations for each species in monoculture. Typically a change in environmental parameters leads to a simultaneous increase or decrease of $\gamma_1$ and $\gamma_2$. As a result all data points group along one diagonal.

In the case of competitive dominance (where one species wins and the other is excluded) the ratio $\rho = \ln B_1/B_2$ practically approaches positive or negative infinity after our standardized simulation period of 40000 days (see above). To visualize intermediate values of $\rho$ close to the coexistence region, the color scale was truncated at absolute values of $|\rho| \geq 7$, i.e. log-ratios above 7 where set to $\rho = 7$ and log-ratios below -7 where set to $\rho = -7$.

To judge bistability, simulations were repeated two times, with either species 1 taken as resident and species 2 as invader (upward-pointing triangles) or species 2 as resident and species 1 as invader (downward-pointing triangles).

Figure 5. Invasion thresholds are plotted on the basis of invasion analysis for 7000 combinations of species 1 and
test species 2. Parameter values taken as in Table S1. However, for species 2 half saturation constants were varied to achieve the chosen values of critical resources, and maximal growth rate and mortality taken as (a) $\mu_{\text{max},2} = 0.04$ h$^{-1}$ and $m_2 = 0.01$ h$^{-1}$ and (b) $\mu_{\text{max},2} = 0.08$ h$^{-1}$ and $m_2 = 0.02$ h$^{-1}$.

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Figure S1: Two-species resource competition in a uniform system (compare to Fig. 3). The four typical cases for the competition outcome are shown, depending on the mutual slopes of consumption vectors $\Gamma_i$ (values of critical resources and supply points remain fixed). For each case the zero-net-growth isoines (ZNGI’s, dashed lines) of species 1 (green) and species 2 (red) are plotted in the $(N, I)$-resource plane, together with the supply point (black circle) and the consumption vectors (green and red arrows). Each species in monoculture reduces resources to an equilibrium point, corresponding to the intersection of its consumption vector with its ZNGI. Invasion of the other species is possible if its critical resource values are located below this point (see Fig. 1b). Define the critical slope $\Gamma_{cr}$ by the slope of a straight line from the supply point to the intersection of the two ZNGIs (indicated as black dashed line). Species 1 can invade if the consumption vector of species 2 has a slope less than the critical slope ($1/\Gamma_2 > 1/\Gamma_{cr}$, top panel), i.e. species 2 has relatively stronger influence on its most limiting resource $N$ than on resource $I$. In contrast, species 2 can invade if the slope of the consumption vector of species 1 is larger than the critical slope ($\Gamma_1 > \Gamma_{cr}$, right-hand panel), i.e. species 1 has a relatively stronger influence on its most limiting resource $I$. 
Figure S2: Ratio of the logarithmic resource gradients $\gamma_1 = c_{i,1}/c_{N,1}$ in the phytoplankton model as a function of several species specific and environmental parameters. Colorcode indicates the value of \text{arctan} $\gamma_1$ calculated numerically for a monoculture of species 1 in the phytoplankton model (see Table S1 for parameters which do not vary). Since $\gamma_1$ equals the slope of the invasion line in the resource plane, this allows to project shifts in the species composition and to gain insight of how a change of one parameter can compensate for the influence of other parameters. For example, an increase of $\gamma$ corresponds to a clockwise rotation of the invasion threshold lines, thereby favoring the best nutrient competitor.

(a) Dependence of $\gamma$ on species traits. Slope of the invasion threshold $\gamma$ grows with the ratio $k/\alpha$. This dependence is also evident from both limits in eqn S6. Together with Fig. 3, this result confirms our suggestion that two-species coexistence is more probable if each species relatively less consumes its most limiting resource (see main text).

(b) Dependence of $\gamma$ on environmental conditions. Slope of the invasion threshold $\gamma$ grows with turbulent diffusivity $D$ and background turbidity $K_{bg}$. This indicates that a turbid but weakly-mixed environment should result in the same species composition as stronger mixed but clean waters (while the total biomass will, of course, in general be different).

(c) Dependence of $\gamma$ on resource supply. Increase of incident light intensity $I_{in}$ or of bottom nutrient concentration $N_B$ decreases $\gamma$ (thus leading to a counter-clockwise rotation of the invasion thresholds) and creates more favorable conditions for the best light competitor (compare to Fig. S3).
Figure S3: Geometrical method to project the outcome of spatial resource competition in dependence of the ambient nutrient concentration, \( N_B \). Top panel: shift in the competition outcome between species 1 (green) and species 2 (red) in the phytoplankton model caused by an increase of the bottom nutrient concentration, \( N_B \). Bottom panel illustrates this shift as the result of a counter-clockwise rotation of the invasion thresholds. The slopes, \( \gamma_1 \) and \( \gamma_2 \), of the invasion thresholds in the bottom panel were calculated numerically for a monoculture of species 1 and 2. To obtain a relatively small difference between \( \gamma_1 \) and \( \gamma_2 \) we used \( \alpha_1 = 8 \times 10^{-10} \) mmol nutrient cell\(^{-1} \) and \( k_2 = 6 \times 10^{-10} \) m\(^2\) cell\(^{-1}\).
Figure S4: Exponential resource distributions in a vertical water column. Shown are the profiles of photosynthetically active radiation (gray), nitrate (blue) and chlorophyll (green) measured during the HOT program ([http://hahana.soest.hawaii.edu](http://hahana.soest.hawaii.edu), station ALOHA, cruise 114; see also Karl 2010). Resource distributions plotted (a) on linear and (b) on logarithmic scales. As it is commonly supposed, the light intensity can be approximated by an exponential distribution (black dotted lines and $R^2$ values in (a) and (b)). In contrast, the nutrient profile exhibits two distinct regimes: In the area below the production layer the nutrient concentration can be well fit by a linear dependence ($R^2 = 0.91$, black dashed line in (a)); however, within the production layer an exponential distribution gives a better description ($R^2 = 0.99$, black dashed line in (b)). We argue that the regime of exponential decay within the production layer is more crucial for competition, because in this area growth can be nutrient limited, whereas in the linear part of the nutrient profile growth is nutrient saturated. Interestingly, the interval of exponential decay ranges over three orders of nitrate concentrations, making this area crucial for the competition among a wide range of phytoplankton species which can be limited by this nutrient (Litchman et al. 2007). Note, that similar exponential distributions in this interval of depths have been observed in mean nitrate concentrations between 1989-2006 (Karl & Letelier 2008). (c) System state curve (SSC), showing the distributions in the resource plane with logarithmic axes. The solid blue line shows actual cruise data for the light intensity and the nutrient. Since light intensities are not reported below a depth of 140 m, the system state curve was extrapolated, assuming that the light intensity follows the same exponential decay up to 300 m (blue dotted line). Dashed line shows a straight line fit (with $R^2$ value) within the production layer.
Figure S5: Exponential distribution of resource concentrations in deep marine sediments. Inverse concentration profiles (note the logarithmic scale) result from biologically catalyzed reactions, which consume and release metabolites in a complex biogeochemical reaction network deep in the sediment column.

(a) Concentration profiles of sulfate (red) and methane (green) from ODP leg 207, site 1258, Demerara Rise, Equatorial Atlantic (Shipboard Scientific party, 2004; Arndt et al. 2006) show a sulfate-methane transition zone in a sediment depth of 150-300 m. Substrates are supplied by a downward flux of sulfate from the sediment-water interface and by the biogenic production of methane from deeply buried organic matter in black shale sequences at a depth below 400 m. Upward diffusing methane and downward diffusing sulfate are depleted by deep sedimentary microbial communities in the process of anaerobic methane oxidation (AMO). Note that the exponential distribution of methane ranges over several orders of magnitude.

(b) Depth profiles of sulfate (red) and barium (blue) along the sediment column of ODP leg 201, site 1228, eastern Pacific ocean (Shipboard Scientific party, 2004; D’Hondt et al, 2004), exhibiting two reversed zonation patterns. The characteristic sulfate-barium transition zone that extends from the water-sediment interface to greater depth (0 to 40 m) is mirrored by a second reversed succession that extends upward from the basement-sediment interface in depth from 40 to 80 m. Here, sulfate enters the sediment from two directions: from the overlying ocean and from an underlying basaltic aquifer. In each transition zone the depletion of sulfate by microbial activity promotes the remobilization of biogenic barium (Torres et al. 1996), giving rise to high concentrations of dissolved barium in the pore fluids beyond the zone of sulfate depletion.

Data are obtained during the ocean drilling program (ODP, [http://iodp.tamu.edu/]). Dashed and dotted lines show straight line fits to the data on the logarithmic axes. $R^2$ values of the fit are indicated.
Figure S6: Illustration of different competition outcomes in the phytoplankton model. The outcome of two species competition is shown (indicated by color coding), depending on the critical resource values of species 2 (critical resources ($N^*_1, I^*_1$) of species 1 fixed, green circle), when the maximal growth rates and mortalities are identical (top panel), or differ by the factor $\beta = 2$ (bottom panel), similar to Fig. 5 from the main text. Insets (a) - (d) show typical spatial profiles corresponding to the different competition regimes (critical resource values of species 2 indicated as red circle). Plotted are the phytoplankton concentration of species 1 (green) and species 2 (red), and the distribution of light (black dashed line) and nutrient (black solid line) as a function of depth $z$.

(a) and (d): Two fundamentally different coexistence mechanisms in a spatial system. (a) Coexistence due to a resource limitation trade-off, mediated by niche segregation in resource requirements which becomes apparent as a spatial separation of density profiles. (d) Coexistence due to a gleaner-opportunist trade-off. The two species are not spatially separated, but species 1 with smaller resource requirements (gleaner) can utilize a larger favorable range, whereas the high growth rate of the stronger resource limited species 2 (opportunist) allows it to survive on a smaller spatial range.

(b) and (c): Two kinds of bistability in the competition outcome. (b) Alternative stable state when each species cannot grow in the presence of its competitor. (c) Alternative stable states of either coexistence (species 1 can invade the monoculture of species 2, but does not reach a high abundance) or a monoculture of species 1 (species 2 cannot establish in the presence of species 1).

Parameter values of species 1 see in Table S1. Simulation parameters for species 2: (a) $H_{I,2} = 24.7 \mu$mol photons $m^{-2} s^{-1}$, $H_{N,2} = 0.012$ mmol nutrient $m^{-3}$, (b) $H_{I,2} = 12.2 \mu$mol photons $m^{-2} s^{-1}$, $H_{N,2} = 0.123$ mmol nutrient $m^{-3}$, (c) $H_{I,2} = 9 \mu$mol photons $m^{-2} s^{-1}$, $H_{N,2} = 0.22$ mmol nutrient $m^{-3}$, (d) $H_{I,2} = 24.7 \mu$mol photons $m^{-2} s^{-1}$, $H_{N,2} = 0.044$ mmol nutrient $m^{-3}$. 
Table S1: Default model parameters

| Symbol | Interpretation | Units | Value |
|--------|----------------|-------|-------|
| Independent variables | | | |
| $t$ | Time | h | |
| $z$ | Depth | m | |
| Dependent variables | | | |
| $P(z, t)$ | Population density | cells m$^{-3}$ | |
| $I(z, t)$ | Light intensity | $\mu$mol photons m$^{-2}$ s$^{-1}$ | |
| $N(z, t)$ | Nutrient concentration | mmol nutrient m$^{-3}$ | |
| Parameters | | | |
| $I_{in}$ | Incident light intensity | $\mu$mol photons m$^{-2}$ s$^{-1}$ | 600 (100 ... 1000)* |
| $K_{bg}$ | Background turbidity | m$^{-1}$ | 0.1 (0.01 ... 1)* |
| $Z_B$ | Depth of the water column | m | 100 |
| $D$ | Vertical turbulent diffusivity | cm$^2$ s$^{-1}$ | 0.3 (0.01 ... 0.7)* |
| $N_B$ | Nutrient concentration at $Z_B$ | mmol nutrient m$^{-3}$ | 10 (0.1 ... 10)* |
| Species 1 | | | |
| $H_I$ | half-saturation constant for light | $\mu$mol photons m$^{-2}$ s$^{-1}$ | 20 |
| $H_N$ | half-saturation constant for nutrient | mmol nutrient m$^{-3}$ | 0.04 |
| $k$ | light attenuation coefficient | m$^2$ cell$^{-1}$ | $6 \times 10^{-10}$ (1 $\times$ 10$^{-10}$ ... 1 $\times$ 10$^{-9}$)* |
| $\alpha$ | cell nutrient content | mmol nutrient cell$^{-1}$ | $2 \times 10^{-10}$ (1 $\times$ 10$^{-10}$ ... 1 $\times$ 10$^{-9}$)* |
| $\mu_{max}$ | maximal growth rate | h$^{-1}$ | 0.04 |
| $m$ | mortality rate | h$^{-1}$ | 0.01 |
| Species 2 | | | |
| $H_I$ | half-saturation constant for light | $\mu$mol photons m$^{-2}$ s$^{-1}$ | 15 |
| $H_N$ | half-saturation constant for nutrient | mmol nutrient m$^{-3}$ | 0.065 |
| $k$ | light attenuation coefficient | m$^2$ cell$^{-1}$ | $1 \times 10^{-10}$ (1 $\times$ 10$^{-10}$ ... 1 $\times$ 10$^{-9}$)* |
| $\alpha$ | cell nutrient content | mmol nutrient cell$^{-1}$ | $5 \times 10^{-10}$ (1 $\times$ 10$^{-10}$ ... 1 $\times$ 10$^{-9}$)* |
| $\mu_{max}$ | maximal growth rate | h$^{-1}$ | 0.04 |
| $m$ | mortality rate | h$^{-1}$ | 0.01 |

* in Fig. 3b and Fig. S2
Appendix S5 provides further details of simulation parameters specific to certain figures.