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CHAPTER 10

The World Is Patchy and Heterogeneous! Trade-off and Source-Sink Dynamics in Competitive Metacommunities

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When we discuss communities beyond their most essential attributes as open systems, generality may elude us, except the generality of diversity. WHITTAKER AND LEVIN, 1977

Introduction

(Levin 1992). This chapter reviews conceptual advances in the understanding of Recognition that the world is patchy and heterogeneous has been the basis for many advances in both fundamental and applied ecology over the last thirty years spatial mechanisms of competitive coexistence and places those advances in the context of metacommunity ecology. We discuss work from the past half century through extremely recent results in order to highlight unexpected links and reinerpretations. We show that apparently different mechanisms have common eleetical framework. This framework could lead to a general theory of biological ments and we discuss options for integrating these elements into a broader theodiversity for a natural world increasingly transformed by human activities.

For the sake of generality, ecologists frequently look for simple rules that apply a wide range of taxa. Community ecology has provided one such simple law in ardin 1960) derived from the Lotka-Volterra models (Lotka 1925; Volterra 26) and Gause's experiments (1934). This principle states that the number of form of the competitive exclusion principle (Lack 1944; Hutchinson 1957; hes. It has brought community ecology into a modern synthesis led by the existing species cannot exceed the number of limiting factors or ecological rks of Hutchinson (1957), Hardin (1960), MacArthur (1967, 1972), and oth-In reality, however, for the vast majority of communities, the competitive exsion principle is too restrictive because it focuses only on local limiting factors homogeneous environment. The original theory could not explain high sity found in natural systems with few identified limiting factors, especially etheory reductionism in favor of complexity and entered a new era without gants and aquatic communities (e.g., Hutchinson 1961). Ecology rejected

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a theory of diversity but with a diversity of theories explaining species richness reviewed by Chesson 2000a, Barot and Gignoux 2004)

1958), the idea became central to community ecology only after Levins defined heterogeneity has led ecologists to focus on mesoscale ecology (Roughgarden barriers (e.g., deep water between coral reefs, ponds separated by land, plant factors or historical context. Although early community ecologists recognized the metapopulations (Levins 1969, 1970), and Levins and Culver (1971) applied the concept to competition theory. The natural link between patchiness and regional et al. 1988; Holt 1993; Ricklefs and Schluter 1993) to incorporate more realisti-Patchy environments have spatially discrete elements either because of natural patches separated by a matrix inconducive to plant growth, etc.) or species characteristics (e.g., sessile species with a dispersal stage, territorial organisms with natal dispersal, etc.). Heterogeneity implies differences in the environmental conditions found in two patches (table 1.1), for example due to different limiting discrete nature of habitats (Skellam 1951; Andrewartha and Birch 1954; Huffaker Forty years after Hutchinson discussed paradoxically high plankton diversity (Hutchinson 1961), recognition of the importance of patchiness and heterogeneity has led to more comprehensive and realistic theories of species diversity. Patchiness and heterogeneity are two interconnected forms of environmental variation. cally the dynamics of species diversity in metacommunities.

1970) metapopulation approach of describing dynamics in terms of extinctions and recolonization of patches. These models assume a net separation between local and regional time scales such that coexistence within a patch is impossible, and onization abilities (Levins and Culver 1971; Horn and MacArthur 1972; Hastings ments have opened different perspectives on local species coexistence through The recognition of spatial structure leads to differentiation between local (within patch) and regional (among patch) processes (see table 1.1). Since the work by Skellam (1951) there has been a separation in model formalism and assumptions between these two scales (figure 10.1). For example, models of coexistence in discrete, homogeneous environments have adopted the Levins (1969, they explain regional coexistence given a trade-off between competitive and col-1980). In contrast, models that consider coexistence in heterogeneous environsource-sink or mass effects in metacommunities (see Holyoak et al., chapter 1 for 2003). This divergence has resulted for example, in competition-colonization definitions). These models have shown that local coexistence is possible through regional niche differentiation and dispersal between patches (Levin 1974; Shmida trade-offs and source-sink dynamics being viewed as mutually exclusive mechanisms of coexistence when they should be studied in a common framework releand Ellner 1984; Amarasekare and Nisbet 2001; Mouquet and Loreau 2002, vant to metacommunity ecology (Amarasekare et al. 2004).

mon framework under metacommunity ecology (Wilson 1992; Holt 1993, 1997a; Ecologists have recently begun to synthesize these two approaches into a com-

Spathesis: coexistence in a competitive metacommunity ĵ 1 Taneyhill 2000 Wang et al. 2002 Chesson 2000b. Amarasekare & Nisbet 2001: Mouquet & Lorean 2002 Amarasekare et al, 2004 Regional similarity Pacala & Rees 1998 Yu & Wilson 2001 Levine & Rees 2002 Wilson 1992, Holt 1993, 1997a Mouquet & Loreau 2003 Spatial storage Hybrid models Community patterns 2000 Û Management of the second second May & Nowak 1994 Tilman 1994 Individual 1990 scale Shmida & Ellner 1984 Chesson 1985 Comins & Noble 1985 Roughgarden 1986 Source sink Mass effect Pulliam 1988 Community Hasting 1980 wasa & patterns Slatkin 1974 Hanski 1983 coexistence Transient 1980 1 _ 1 Levins & Culver | Source Sink Levins 1974 Competition-Hom & MacArthur 1972 colonization Homogeneity 1970 Heterogeneity Precursors Skellam 1951 1953

colonization hypothesis (gray arrows) and a third branch that have studied the consequences of bottom of the figure. These papers are reviewed in the text and summarized in table 10.1. This list is Figure 10.1 Representation of the chronology of some important papers that have studied the conditions of coexistence between competing species in patchy habitats. There was rapid separation between studies that considered homogeneous (white region) and heterogeneous (gray region) environments. Three main branches have emerged from the precursor works of Skellam (1951) and Levene (1953), respectively the source-sink hypothesis (dark gray arrows), the competitiontransient local coexistence in patch models (white arrows). We have added a chronological axis at the not exhaustive, but rather represents what we think are the key papers.

Mouquet and Loreau 2002, 2003; Loreau et al. 2003; Leibold et al. 2004; Holyoak proaches (Yu and Wilson 2001; Levine and Rees 2002), a thorough clarification of the ideas underlying spatial coexistence is necessary for further progress in metacommunity ecology. Contemporary ecologists (including the authors of this chapter) have also often failed to recognize insightful contributions of early works ences in focus between more recent metacommunity ecology and the early work ct al., chapter 1). Although we are not the first to make the link between these apsuch as Skellam (1951) and Levene (1953). This oversight is partially due to differon species coexistence in space. For example Levins and Culver (1971), only oriefly considered the general topic of coexistence and species richness. Their foand systems. The diversity of biological systems modeled also contributed to the cus instead was on the role of competition in the dynamics of rare species on is-

lack of early synthesis because the questions addressed in marine versus terrestrial systems were not necessarily similar, and hence generalizations were difficult to

With a view toward clarification, in this chapter we review theory on coexistence in patchy environments, focusing on the competition-colonization (herafter CC) and the source-sink (hereafter SS; also known as the mass effect) hypotheses. We retrace the chronology of these two approaches, highlighting their key results, differences, and similarities. We end with the basic elements of a more general framework that allows ecologists to examine patch dynamics and mass effects simultaneously. Table 10.1 and figure 10.1 provide an overview of these models that might be used as a guideline during the reading of this chapter. We models that might be used as a guideline during the reading of this chapter. We is presented elsewhere (Keddy 1982; Shmida and Ellner 1984; Kadmon and Shmida 1990; Gonzalez et al. 1998; Forbes and Chase 2002; Cottenie et al. 2003; Kneitel and Miller 2003; Urban 2004; Miller and Kneitel, chapter 5; Cottenie and development of these ideas has probably biased our choices, but our goal is to be comprehensive rather than exhaustive.

The Precursors: Skellam (1951) and Levene (1953)

eral ecological niches, with one allele favored in one niche and the other allele favored in another, might increase the possibilities for attainment of equilibrium such conditions, he found that the polymorphism was maintained if the weighted arithmetic mean of the "relative fitnesses" of homozygotes was less than one. This annual plants within the same habitat and showed that coexistence was possible given that "a disadvantage in direct competition may be offset by a superiority in reproductive capacity" (215). He further demonstrated that the advantage in reproductive capacity that an inferior competitor needs to persist depends on the density of the superior competitor (the effect of the superior competitor on the inferior competitor). Levene studied similar coexistence concepts with genes (1953; see also Levins and MacArthur 1966). He examined the conditions for the posed a simple model of allele frequencies considering that "the existence of sevwith both alleles present in substantial proportions" (Levene 1953, 331). Under persal for both local and regional species coexistence. He showed that a population could maintain itself in unfavorable regions by random dispersal from favorable regions. He stated that the stability of such a system depended "on the ability of the population in favorable habitats to make good the decline in the unfavorable ones" (Skellam 1951, 202). He also studied competition between two maintenance of polymorphism without any heterozygote advantage. He pro-The works of Skellam and Levene are seminal to the development of metacommunity theory. Skellam used diffusion theory to study the consequences of dis-

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condition, which is obviously true when the heterozygote has higher fitness, can also be obtained for particular parameter values without the heterozygote being superior to both homozygotes in any single niche.

Both authors, therefore, investigated the role of regional heterogeneity in local coexistence. Skellam's work also examines CC dynamics long before metapopulation theory (Levins 1969, 1970) brought it to mainstream attention (see next section). Skellam's work is seminal to the development of metacommunity theory because it provides the basis for both the competition-colonization trade-off and source-sink dynamics. Levene (1953) makes the fundamental link between niche theory and spatial coexistence. He is, to our knowledge, the first to propose such a regional vision of niche separation allowing local coexistence despite strong local niche overlap. Although neither author extended his results to community ecology or considered the consequences of variation in dispersal ability, these papers are the roots from which metacommunity ecology has grown, and they should be viewed as ecological masterpieces.

Competition, Patchiness and Spatial Heterogeneity The Levins Metapopulation Model

Levins defined a metapopulation as a set of local populations with extinctions and recolonizations, giving us the most common formalism for metapopulation theory (1969, 1970). His work is based on three major assumptions: (1) The environment consists of an infinite number of patches ("localities" in the terminology of table 1.1) with identical environmental conditions, and each patch supports a local population. (2) Local dynamics operate on a faster timescale than regional dynamics, with each patch reaching carrying capacity instantaneously upon colonization. (3) Dispersal is infrequent, serving only to colonize empty patches and having no effect on within-patch population dynamics.

With these assumptions, the dynamics of patch occupation results in a balance between extinction (by stochasticity and disturbance) and recolonization from other patches in the region. Levins modeled the proportion of patches *p* occupied by a species at the regional scale:

$$\frac{dp}{dt} = cpV - ep$$
with $V = 1 - p$,

where c is the recolonization rate of empty patches, and e is the extinction rate of occupied patches. This model assumes that the colonization probability for any patch is proportional to the proportion of patches already colonized. In order to highlight competition for empty habitat, we have modified the original model to isolate V_i the proportion of empty patches in the region. The equilibrium is given by

Table 10.1 Summary of important metacommunity models										
Authors		Org	Comp	М	CCR	CCL	SSD	SST	Notes	
Skellam	1951	TA, P	P	3, 5	No	Yes	Yes	No	Basis for CC and SS hypothesis.	
Levene	1953	All	D	5	No	No	Yes	No	Explains allelic polymorphism without any higher adaptive value for	
									heterozygote.	
Levins and Culver	1971	TA	D	1	Yes	No	No	No	Basis for CC models.	
Horn and MacArthur	1972	All	D	1	Yes	No	No	Yes		
Slatkin	1974	All	D	1	Yes	No	No	No	CC model and proto-SS since a CC trade-off in habitat 1 allows specie persist and maintain itself in habitat 2 via a SS effect. Recolonization from doubly-occupied patches enhances the potential	
Levin	1974	All	D	1, 2, 3	Yes	No	Yes	Yes	species coexistence. Perturbation theorem and first real SST.	
Hastings	1980	MA	D	1	Yes	Yes	No	No	Assumes extinction is a result of disturbance and finds a hump-shapes	
Hanski	1981	TA	D	3	Yes	No	No	No	relationship between species richness and disturbance rate. Spatial variance in the better competitor can be high due to inefficient	
Hanski	1983	TA	D	1	Yes	No	No	No	dispersal. Rescue effect. Coexistence in patchy environment depends on the disti	
Shmida and Ellner	1984	P	P	4	Yes	Yes	Yes	27-	between local and regional time scales.	
Chesson	1985	MA, P	P	4	No	No	res Yes	No	Coined the term mass effect.	
Comins and Noble	1985	MA, P	P	4	No	No	Yes	No No	Spatial and temporal storage effects.	
Iwasa and Roughgarden	1986	MA	P	1,4	No	No	Yes	No	Assuming that the environment is uniformly variable, every coexisting must have its own transient spatial niche. Number of locally coexisting species cannot exceed the number of	
									communities in the region. Regional compensation between larval grorates.	
Kishimoto	1990	All	D	3	No	No	Yes	Yes	An infinite number of engine	
Wilson	1992	All	D	3	No	No	Yes	No	An infinite number of species can exist over two patches. Coined the term metacommunity. Limitation to the founder effect arg	
Holt	1000								proposed by Levin (1974).	
Goldwasser et al.	1993	All	D	3	No	No	Yes	Yes	Mesoscale and metacommunity perspective.	
Tilman	1994	P	D	1	No	Yes	No	No	There is a limit to coexistence with the CC if one considers finite habita	
1 minut	1994	P	D	1	No	Yes	No	No	Applies the Hastings (1980) CC model at the individual scale.	

Holmes and Wilson	1998	P	D	1	Yes	Yes	No	No	If the superior species is rare, it is possible for a long-distance disperse is both an inferior competitor and a bad reproducer, to coexist
Pacala and Rees	1998	P	P, D	1,4	Yes	No	No	No	Model that combines CC and the successional niche.
Chesson	2000b	All	P, D	4, 5	No	No	Yes	No	Extends the concept of spatial storage effects.
Muko and Iwasa	2000	MA	P	4	No	No	Yes	No	Between habitat variation in mortality ratios promotes coexistence, w of reproductive rates does not.
Taneyhill	2000	All	D	1	Yes	No	No	No	Migration from doubly occupied patches is always stabilizing.
Amarasekare and Nisbet	2001	TA	D	3	No	Yes	Yes	Yes	Perturbation theorem with spatial variance in species fitness. Limit to competitive asymmetry.
Yu and Wilson	2001	All	P	1, 4	Yes	Yes	No	Yes	A dispersal fecundity trade-off can partition variation in patch densit creating the conditions for SS based on a priority effect.
Levine and Rees	2002	P	P	5	No	Yes	Yes	No	Provide a hybrid model where coexistence occurs via SS dynamics an facilitated by a CC trade-off.
Mouquet and Loreau	2002	All	P	1, 4	No	No	Yes	No	Definition of the regional similarity constraint. High local diversity a intermediate dispersal and regional species dissimilarity.
Wang et al.	2002	All	D	1	Yes	No	No	No	Stochastic local extinctions make it likely that species with strong con effects on each other can coexist.
Mouquet and Loreau	2003	All	P	1,4	No	No	Yes	No	Community patterns in a source-sink metacommunity.
Amarasekare et al.	2004	All	P, D	1	Yes	Yes	Yes	Yes	Spatial variation in the expression of a life-history trade-off can constrather than promote coexistence.

Notes: Org refers to the type of organisms considered: (All) for all species (or undefined species), (TA) for terrestrial animals, (P) for plants, (MA) for marine animals. Comp refers to t competition, either by dominance (D) or by preemption (P). M refers to the type of model used: (1) patch dynamic, (2) diffusion reaction, (3) Lotka-Volterra, (4) lottery model, and (5) next four columns describe the coexistence mechanism addressed: competition-colonization trade-off at the local (CCL) or regional (CCR) scales as well as source-sink directional (SSD) a old (SST) dynamics as defined in the text. Notes describes the contribution within the paper that motivated our choice to include it in the table. We acknowledge that there are many of community models that we did not include due either to our oversight or because they did not add any fundamentally new result. Some of these papers are also presented in figure 10.1.

$$5 = 1 - \frac{e}{c}$$

$$\hat{\gamma} = \frac{e}{c}.$$
(10.2)

The minimum proportion of available patches to maintain a species in the metapopulation is V_i , and the basic metapopulation growth rate of a species is r = c/e (Fagerström and Westoby 1997; Loreau and Mouquet 1999). This growth rate is the change in the proportion of occupied patches when a species first enters a metapopulation. The persistence of a species requires that r be greater than or equal to 1 (recolonization must compensate for extinction) and that a minimum number of patches be available at the regional scale $(V > \dot{V})$. Note also that there will always be some empty patches at equilibrium because, as \dot{V} tends toward zero, cends toward infinity or e tends toward zero, both of which are biologically unrealistic (Tilman 1994). There is a strong similarity with models of resource competition if one considers available patches as a resource consumed by a species at a rate r. In this context the quantity \dot{V} can be interpreted as Tilman's R^* (1982) since it is the minimum amount of resources (available patches) required for persistence in a metapopulation (Loreau and Mouquet 1999).

One can generalize equation 10.1 to multispecies competition if species differ only in their metapopulation growth rates.

$$\frac{dp_i}{dt} = c_i p_i V - e_i p_i$$

$$V = 1 - \sum_i p_i$$

$$V_i = \frac{c_i}{e_i}$$

$$V_i^* = \frac{1}{r},$$

$$V_i^* = \frac{1}{r},$$

where V_i^* is the minimum proportion of available patches for a particular species takes the form of a lottery (Sale 1977; Chesson and Warner 1981) with an infinite number of patches. At equilibrium, only the species with the highest r_i (i.e., lownow the threshold required for the persistence of other species and thus competitively excludes other species (Loreau and Mouquet 1999). This result matches competitive exclusion by the species with the lowest R^* as proposed by Tilman (1982). When competition takes the form of a simple lottery based on extinction and recolonization, V_i^* , therefore, defines species' competitive abilities.

The Compatition of The Thirty of The

The Competition-Colonization Trade-off Hypothesis

The first models to consider competition in a metapopulation focused on dominance competition. These models are variations of the Levins (1969, 1970) model and all explore coexistence without classical niche partitioning mechanisms but with species differing in the way they use a single resource—space. Levins and Culver (1971) proposed the first such model. They assumed that the competitive effect between species affected their relative extinction or colonization rates. For example, for two competing species the extinction and colonization parameters of species 1 can be written as

$$e_1 = e_{12}p_2 + e_{10}(1 - p_2)$$
 (10.4)

and

$$c_1 = c_{12}p_2 + c_{10}(1 - p_2),$$
 (10.5)

where $e_{1,2}$ $c_{1,2}$ are the respective extinction and colonization rates for species 1 in patches with species 2. Similarly, $e_{1,0}$, $c_{1,0}$ are the respective extinction and colonization rates for species 1 in patches that do not contain species 2. The proportion of patches occupied by species 2 is p_2 . When competition affects extinction (equation 10.4), they found that coexistence is possible if:

$$c_1 > e_{12} - e_{10}, \tag{10.6}$$

which can be interpreted as a requirement that the colonization rate of species 1 must be higher than the competitive effect of species 2 on species 1. They found that when competition affects colonization (equation 10.5), coexistence is possible if

$$\frac{c_{10}}{c_{10} - c_{12}} > 1, \tag{10.7}$$

which is always true, given that $c_{10} \neq 0$ and $c_{12} > 0$. The main result is thus that there must be a trade-off between the colonization and competitive effects for coexistence (equation 10.6) when competition affects species' extinction rates.

Following Levins and Culver (1971), Horn and MacArthur (1972) studied competition in an environment with two kinds of habitat. They used the same model and varied the fraction of habitat where each species outcompeted the other. They found no limit to the number of species that can coexist in such a heterogeneous environment as long as there is a supply of unoccupied patches and the colonization rate increases for each successive new competitor. They also found that the inferior competitor needs only to persist in one habitat to persist in both habitats. While Horn and MacArthur examined equal within- and between-habitat dispersal values, Levin (1974) then examined a similar model in which local (within-habitat) dispersal is much higher than regional (B habitats)

dispersal. This case is relevant when patches tend to be clumped according to types. He showed that coexistence in this system does not require two types of patches if both species have within-habitat dispersal parameters sufficiently high that enough colonists of each species are present to replace lost populations.

single species metapopulation model. He assumed that competition is sufficiently rapid (or patches sufficiently small) that inferior competitors are instantaneously sumed a strict hierarchy of competitive dominance and added the possibility of Hastings (1980) developed a simpler model also adapted from the original that the extreme result of this assumption is the modeling of microsites, or excluded, and no patch contains more than a single species. He acknowledged oatches that hold only a single individual (table 1.1). Hastings (1980) also asextinction by external disturbances. For two species, the model reads

$$\frac{dp_1}{dt} = c_1 p_1 (1 - p_1) - e p_1$$

$$\frac{dp_2}{dt} = c_2 p_2 (1 - p_1 - p_2) - c_1 p_1 p_2 - e p_2.$$
(10.8)

This corresponds to a very strong, unidirectional competitive effect with species extinctions that are independent of competition. One can reformulate Hastings's model in terms of V^* as we did for the simple metapopulation model. The equilibrium is then

$$\hat{p}_{1} = 1 - V_{1}^{*}
\hat{p}_{2} = V_{1}^{*} - V_{2}^{*}
V_{1}^{*} = \frac{e}{c_{1}}
V_{2}^{*} = \frac{c_{1}\hat{p}_{1}}{c_{2}} + \frac{e}{c_{2}}.$$
(10.9)

worse competitor has to be a better colonizer (low V^*). This result is interesting to colonize empty sites and linked to species V*. Hastings (1980) also found a hump-shaped relationship between species richness and disturbance rate. At low For species 2 to coexist at equilibrium with the superior competitor, the equation must satisfy $\hat{p}_2 > 0$, that is $V_2^* < V_1^*$. The competitive advantage of species 1 is then. more than compensated by the ability of species 2 to get free patches, that is the because it helps to differentiate between two aspects of a species' competitive disturbance rates the best competitor excludes all other species; at intermediate disturbance rates coexistence is most likely; but at high disturbance rates coexis Species 1's performance is determined solely by its own life-history characterise ability; dominance competition that is based on direct displacement of one species by a superior competitor and spatial competition that is based on the ability tics, while that of species 2 also depends on species 1's ability to exclude it $(c_1\hat{p}_1/c_2)$.

tence is impossible because species do not have sufficiently high colonization rates Dynamics in Competitive Metacommunities to compensate for extinction.

Hastings's (1980) model opened the way for extremely local, microsite (table 1.1) applications of CC models to plant communities and parasite strains (Nee and May 1992; May and Nowak 1994; Tilman 1994). These more recent models explicitly assume that only one individual can occupy a microsite, that is, that the and the colonization rate is equivalent to the species' reproductive rate. Both of ultimate level of patchiness is the individual itself (Grubb 1986). In this context the extinction rate of Hastings's (1980) model is equivalent to a mortality rate, these new parameters are independent of local competition. As in Hastings's (1980) model, these formulations all assume a strict competitive hierarchy and instantaneous exclusion of inferior competitors. These models predict that a virtually infinite number of species can coexist locally given an appropriate trade-off between competitive ability and fecundity or mortality (but see Adler and Mos-

Some other models yield similar results. Models based on other types of competition yield similar results. For example, Hanski (1981) introduced spatial heterogeneity (spatial variance in abundances) into the Lotka-Volterra competition model in the context of patchy habitats. He studied regional coexistence in cases where local coexistence was not possible. His analysis suggests that an inferior competitor might survive regionally if the spatial variance in abundance of the better competitor is high, for example, due to a low rate of dispersal. Shmida and Ellner (1984) proposed a pure lottery model (Sale 1977; Chesson and Warner 1981) with plant species competing for microsites (table 1.1). They found basigies in terms of competitive versus colonizing abilities explain local coexistence cally the same constraint on species parameters: differences in life-history stratebetween competing species.

Although the CC hypothesis has received much attention from ecologists, it scales. One source of confusion is that colonization subsumes a suite of species has also generated confusion because of its application at both local and regional teristics, particularly fecundity. In actuality, trade-offs between any two of these characteristics (e.g., fecundity, long and short distance dispersal, and colonization rate) and that competition is not always clearly separated from these characfeatures may lead to coexistence (Holmes and Wilson 1998, Yu and Wilson 2001). For example, Yu and Wilson (2001) showed that a trade-off between dispersal and fecundity could enhance the potential for species coexistence. Holmes and Wilson (1998) examined the nature of dispersal and demonstrated that, when the superior competitor is not very abundant, an inferior competitor with low fecundity but long distance dispersal can persist. This confusion about the nature of the trated with Hastings's (1980) model, the notion of competitive superiority can be trade-off is increased by confusion about the nature of competition. As we illusdefined from different perspectives. A fugitive species that is supposed to be a bad

sults from stochasticity or deterministic disturbances (as in Hastings 1980), while at local scales, microsites become available through individual mortality (as in Tilman 1994). Even if these different causes of patch vacancy are necessarily recompetition. An additional source of confusion is that very different factors drive the dynamics of available patches at the different scales, and these dynamics are crucial to coexistence. At the regional scale (table 1.1), availability of localities repetitor for available patches (a species with a low V^\star) in models involving spatial competitor in models of dominance competition can be defined as a good comlated, their interpretations in terms of population dynamics are different.

Transient Local Coexistence

patch states: unoccupied by either species, occupied by species 1 or 2 alone, or co-occupied by both species. He found that migration from co-occupied patches increases the likelihood of coexistence by allowing rare species to colonize patches tence cannot occur via a CC trade-off alone. Therefore, several models have ver's assumption overestimates the number of patches where species co-occur and hence overestimates reciprocal competitive effects. Slatkin instead included four One of the main criticisms of patch dynamic models based on the original Levins 1985; Chesson and Huntly 1997) that when this assumption is abandoned and Slatkin (1974) pioneered this approach with a hybrid between Cohen's model Culver assumed that species were distributed independently. This assumption out that species cannot both sort independently and affect each other's colonizations of dominant competitors (Hanski 1983; Levine and Rees 2002). Yu and Wilson (2001) pointed out (following Chesson and Warner 1981; Comins and Noble seedlings only compete for available patches, as in a lottery system, local coexis-(1970) and Levins and Culver's (1971) model. In their original paper, Levins and allowed simplifications of the mathematics in their model, but Slatkin pointed tion or extinction rates (see also Levin 1974). Slatkin argued that Levins and Cul-(1969, 1970) formulation is the unrealistic assumption that colonization leads instantaneously to exclusion of inferior competitors and carrying capacity populaincluded transient patches in which inferior and superior competitors coexist. while increasing the extinction rates of common species.

portion of sites occupied by that species in the region (but see Nee et al. 1991). He table 1.1). The rescue effect weakens intraspecific competition, thus changing the balance of inter- and intraspecific competition on the regional scale, such that the Hanski (1983) found an exception to Slatkin's result if extinction is a function. of patch occupancy (as colonization is for all patch dynamic models) so that the probability of extinction for a given species is negatively correlated with the procalled this dependence of extinction on patch occupancy "the rescue effect" (cf. species with the wider initial distribution can outcompete the other species.

ing these two previous models into a broader framework. He showed that immi-Taneyhill (2000) elaborated on the effects of co-occupied patches by integrated

gration from co-occupied patches is always stabilizing and that the rescue effect Dynamics in Competitive Metacommunities

(as defined by Hanski 1983) is typically destabilizing for the less widespread competitor. Wang et al. (2000, 2002) extended these results by adding stochastic extinction and considering extinction to be a function of local abundance. They found that stochastic local extinctions make it more likely for strongly competing

occupied patches increases the probability of coexistence between competing species. In these models, doubly-occupied patches introduce an element of spatial heterogeneity (variation in spatial abundances) because they act somewhat as a refuge for the inferior competitor. Co-occupied patches supply colonizers that are The key result to emerge from these studies is that colonization from copetitor. As Chesson has pointed out (2000b), such heterogeneity in propagule already removed from the fraction of patches occupied solely by the inferior comproduction or resource-use contributes to coexistence by creating spatial niches. To simplify his model, Slatkin (1974) also examined species with similar colonization and extinction rates, an approach continued by others examining doubly-occupied patches (Taneyhill 2000; Wang, Zhang, et al. 2000; Wang, Wang, et al. 2002). These models also, therefore, reveal that ecologically similar species that would exclude each other in a uniform environment may coexist in a patchy environment (Slatkin 1974; Taneyhill 2000; Wang, Zhang, et al. 2000; Wang, Wang, et al. 2002). The following section focuses specifically on spatial heterogeneity, which is more commonly associated with SS models.

Regional Heterogeneity and Source-Sink Dynamics

The SS perspective focuses on spatial heterogeneity of the environment rather than variation in species' life-history traits. All SS models share a set of common assumptions: (1) the environment is divided into patches, and each patch has different environmental conditions (regional heterogeneity); (2) local and regional dynamics operate on the same time scale; and (3) immigration is frequent and can change the outcome of local competition (a mass effect sensu Shmida and Whittaker 1981). This idea has been applied to a wide range of taxa and different modes of competition, and investigated using either the Lotka-Volterra model (for both dominance and preemptive competition) or lottery models (for preemptive competition).

LOTKA-VOLTERRA MODELS (LV)

Aside from the initial work of Skellam (1951) and Levene (1953), the first person to apply SS ideas to community ecology was Levin (1974). He developed a twopatch LV model of competition in which founder effects (random differences in initial abundances) allow different species to dominate numerically in different patches (preemptive competition leading to a priority effect). Dispersal from safe areas maintains species in communities from which they would otherwise be ex-

cluded (due to lower initial abundance). Given the perturbation theorem (when view) the coexistence equilibrium is stable when dispersal between communities is low. Levin suggested that dispersal has the potential to increase species richness but that high dispersal rates reduce local species richness below that in isolation. This finding suggests that intermediate dispersal maintains differences between different stable boundary equilibria occur in different places in space, small perturbations can create stable interior equilibria; see Amarasekare 2000 for a rethe localities and communities.

(spatial variance in fitness). They have shown that when competitive asymmetry sal threshold. High dispersal rates undermine the potential for coexistence by since clarified the conditions for species undergoing dominance competition by considering spatial variation in the strength of competition at the regional scale because these species can maintain themselves at low resource levels. The surplus different limiting factors in each locality. Amarasekare and Nisbet (2001) have between species is high, local coexistence is possible only below a critical disper-Subsequent studies focused on the maintenance of spatial variation in fitness and the effects of this variation on diversity. Following Levin, Kishimoto (1990) ciprocal competition coefficients) and dispersal between the two patches is intermediate. The interaction between strong competitors results in a resource surplus becomes open to other less competitive species. As the number of species increases, the resource surplus decreases, and further invasion becomes less likely. dances for species engaged in preemptive competition, or due to differences in competitive ability for species engaged in dominance competition, for example, showed that many species can coexist in a two-patch system if the reciprocal competitive effects between the two dominant species are sufficiently strong (high re-Again, competitive superiority can be due to either differences in initial abunreducing spatial variance in fitness (see also Nichimura and Kishida 2001).

Wilson's (1992) results is at least partly because Wilson does not consider a mass effect that would rescue species from local extinction, regardless of the metacom dispersers for a second metacommunity assembly process. In this context, the community scale. Note, however, that the difference between Levin's (1974) and An important result from Levin (1974) is that spatial variance in fitness can result only from heterogeneity in initial species abundances, provided competition is preemptive. Wilson (1992) found somewhat different results by considering patches colonized at random by individuals from a disperser pool. Species then interact in a closed system; since the outcome is dependent on initial conditions model then pools species at the metacommunity scale to generate a new pool species composition at the regional scale converges toward the species composition the patches end up with different combinations of species. The Wilson (1992) siders extinctions and recolonizations based on species abundances at the meta tion at the local scale. The founder effect does not hold in the long term if one com munity assembly rules.

Dynamics in Competitive Metacommunities

The SS concept has been well-integrated into lottery models (Sale 1977; Chesson and Warner 1981). Because they consider recruitment as central, immigration and emigration can be crucial in these models. For example, some authors (Shmida and Ellner 1984; Kadmon and Shmida 1990; Loreau and Mouquet 1999) have included immigration from a regional pool in the form of a mass effect, which maintains high local species richness in plant communities. In these modber of seeds at a site wins. Given such rules for site attribution, immigration from els, plants compete for available microsites, and the species with the highest numoutside the community can maintain inferior species in the system.

Lottery models with SS dynamics have also been applied to several aquatic systems. These systems are essentially competitive metacommunities with sessile adults and mobile larvae. Planktonic larvae produced in all the local communities enter a common pool and are equally redistributed among communities. Chesson (1985) developed a stochastic model with regional heterogeneity such that each locality favors the adults of a different species (via increased survival). He showed that species can coexist locally with emigration-immigration and higher adult survival in favorable localities. As with the temporal storage effect (Chesson and Warner 1981; Chesson 1983, 1984) in which a population sustains a positive average growth rate in a temporally fluctuating environment if growth rates in good years more than compensate for bad years, here a "spatial storage effect" promotes coexistence because it buffers a species from poor recruitment in patches where it has a negative growth rate (See also Hoopes et al., chapter 2). Iwasa and Roughgarden (1986; see also Muko and Iwasa 2000) found similar results in another model of marine systems.

While the above studies suggest that spatial heterogeneity is important for coexistence, Comins and Noble (1985) showed that spatial heterogeneity does not have to be permanent to promote coexistence. In their model coexistence requires ditions although there are no permanent differences between parts of the habitat. statistical distribution of environmental conditions is the same in all patches gach species to be dominant in site establishment under some environmental con-Rather, the environment is uniformly variable in the sense that the long-term transient niches). They found that every coexisting species must have its own onmental variability in the basic model can be obtained either with a few patches gansient niche. They also found that the stabilizing effect (coexistence) of envind complete mixing or with a large number of patches and little mixing.

stence results from a spatial storage effect, as defined above (see also Hoopes al., chapter 2). Two other mechanisms driven by spatial variation contribute to le storage effect but can also act alone to allow coexistence: spatial variation in Chesson (2000b) recently proposed a generalization of these results where codedegree of nonlinearity in species competitive responses (spatial relative nonDynamics in Competitive Metacommunities

linearity) and localized dispersal that concentrates an invading species in areas favorable to its growth and reproduction (growth-density covariance; see Chesson 2000b and Chesson et al., chapter 12 for more details).

2000b and Chesson et al., Chapter 12 101 inch chapter 18) have recently de-Mouquet and Loreau (2002, 2003; Loreau et al., chapter 18) have recently developed a metacommunity model of lottery competition, which, because it shares veloped a metacommunity model of lottery competition, which, because it shares some characteristics with metapopulation CC models, helps to compare the outsome characteristics with metapopulation CC models, helps to compare the outsome characteristics with metapopulation at the

some characteristics with metapopulation CC models, net poor court comes of SS and CC models. This model is based on equation 10.3 applied at the individual level and assumes that localities receive a constant number of imminidividual level and assumes that metacommunity. The model is given by grants from other localities in the metacommunity. The model is given by

$$\frac{dP_{ik}}{dt} = (I_{ik} + [1 - a]c_{ik}P_{ik})V_k - m_{ik}P_{ik}$$

$$I_{ik} = \frac{a}{N - 1} \sum_{i \neq k}^{N} c_{il}P_{il}$$

$$V_k = 1 - \sum_{i = 1}^{S} P_{ik}.$$
(10.10)

Here *i* refers to species and *k* to localities. The parameter *a* describes the proportion of gration term. At the local scale, *S* species compete for a limited proportion of gration term. At the local scale, *S* species compete for a limited proportion of wacant microsites, V_k . The metacommunity consists of N localities that differ in vacant microsites, V_k . The metacommunity consists of N localities, the species their local conditions. When there is no dispersal between localities, the species with the highest local basic reproductive rate $(r_k = c_{ik}/m_k)$ excludes all other spewith the highest local basic reproductive rate $(r_k = c_{ik}/m_k)$ excludes all other species in the locality because it decreases the proportion of vacant sites, V_k , below cies in the locality because it decreases the proportion of vacant sites, V_k , below persal, and different species dominate in different localities (due to spatial hereogeneity in competitive rankings), local coexistence is possible. Individuals emergence areas prevent competitive exclusion in sink areas (where they igrating from source areas prevent competitive exclusion in sink areas (where they are competitively inferior). At equilibrium each individual of each species must are competitively inferior). At equilibrium each individual that survives somewhere in on average during its lifetime produce one individual that survives somewhere in the metacommunity so that each species' average net reproductive rate at the scale of the metacommunity is:

$$\overline{R_i} = \frac{\sum_{k=1}^{N} \hat{V}_k r_{ik} w_{ik}}{\sum_{N} w_{ik}} = 1.$$
 (10.11)

Here w_{ik} is the total quantity of propagules of species i in community k per unit of

 $w_{ik} = (1-a)c_{ik}\hat{P}_{ik} + \frac{a}{N-1}\sum_{l \neq k}^{N} c_{il}\hat{P}_{il}.$

(10.12)

In this lottery model, a species' competitive ability is directly related to its reproductive rate. Thus equation 10.11 can be interpreted as a constraint of regional competitive similarity (or equivalency) between coexisting species. For instance, each species' basic reproductive rate r_{ik} must be balanced over the metacommunity for equation 10.11 to hold. Local basic reproductive rates can be different locally so that different species will dominate in each locality, but they are equivalent when averaged over the region. Conditions for species coexistence are then a function of the proportion of dispersal between communities. When dispersal is too high, the metacommunity is homogenized, and species richness declines because the species that is the best competitor at the regional scale excludes all other species from the metacommunity.

to coexistence when the assumption of a strict dominance hierarchy was relaxed. If coexistence could occur via spatial heterogeneity and SS dynamics, however, a ate spatial variation in initial species abundances. More recently, Yu and Wilson (2001) incorporated such spatial variation in patch density in what is essentially Huntly 1997) without dominance competition. Yu and Wilson varied the fraction showed that inferior competitors can be rescued from competitive exclusion in communities with high h by emigrating from communities with low h (where the off. Yu and Wilson also showed that including a trade-off between dispersal and preemptive competition, but can allow coexistence via SS dynamics. Levine and Rees (2002) also found that a competition-colonization trade-off could not lead competition-colonization trade-off operating simultaneously can predict the The SS models discussed above assume that extrinsic spatial heterogeneity crogeneity can also arise from founder effects or other random factors that genera CC lottery model at the microsite scale (based on appendix E in Chesson and h of available habitat in each community. In models with complete mixing, they best competitor cannot persist or is at very low density) because of the CC tradefecundity can increase the potential for coexistence via spatial variation in patch density. In other words, CC itself cannot lead to coexistence in a lottery model with drives spatial variance in fitness. However, as shown by Levin (1974), intrinsic hetabundance patterns observed in annual plant communities (Levine and Rees 2002).

SYNTHESIS ON SOURCE-SINK DYNAMICS

Despite the differences in their mathematical formulations, IV and lottery models yield similar results. First, local coexistence in a SS metacommunity results from spatial variation in the strength of competition. Second, coexistence is most likely for intermediate dispersal rates. There is a critical dispersal threshold below which dispersal is too low to prevent local competitive exclusion and a critical dispersal threshold above which coexistence is impossible because spatial variation in fitness is homogenized by immigration. Third, species have limits to regional dissimilarity or competitive asymmetry that are a function of dispersal between communities.

Spatial heterogeneity is essential for coexistence in these models. It arises from spatial variation in abiotic or biotic factors that cause spatial variation in the strength of competition or from spatial variation in species abundances due to founder effects or other random phenomena. As pointed out by Amarasekare tions, and it is likely that increasing dispersal will rapidly decrease species richness (2000), these two kinds of heterogeneity correspond to two different types of local dynamics—threshold and directional. Threshold dynamics occur in systems where the outcome of competition depends on initial abundances (e.g., preemp-(following an initial increase at very low dispersal rates). Directional systems are tive competition) such that coexistence is possible given spatial variation in species abundances. In contrast, systems with directional local dynamics (e.g., dominance competition) are driven by spatial environmental heterogeneity and phenotypic plasticity. Threshold systems are highly sensitive to large perturbaless sensitive to perturbation, and they are more likely to have higher species richness at high dispersal rates. In both systems, however, very high dispersal values tend to homogenize spatial variance and reduce species richness. The results obtained with threshold systems (Levin 1974; Kishimoto 1990; Amarasekare 2000; exist only because the environment is patchy (but otherwise homogeneous), and abundances vary in space. Regional niche differentiation is not needed for species Yu and Wilson 2001; Levine and Rees 2002) are striking because species can co-

its to similarity and regional niche differentiation. For instance, the condition of Because coexistence in SS models is frequently dependent on maintaining garden do not interpret their results in the context of niche theory, but they state spatial variation in fitness, these models point toward generalizations about limregional similarity defined by Mouquet and Loreau (2002) casts coexistence in terms of niche theory and thereby alludes to earlier models. Iwasa and Rough-(in theorem 2) that coexistence requires compensation between larval productivity of species at the regional scale, that is, that species must have similar larval productivity at the scale of the region (1986, 198). This idea of regional compensation is also evident in Chesson's (1985) work on the spatial storage effect (see also Chesson 2000b). Models based on the LV framework produce the same regional similarity constraint. For instance, Amarasekare and Nisbet (2001) define a limit to competitive asymmetry that is equivalent to a constraint of similarity between coexisting species. These different results can be interpreted in the context of niche theory given that each locality has a different limiting factor (resulting in a different species dominating each community). Niche differentiation occurs at the regional scale and registers at the local scale through dispersal. Defining species as similar at particular spatial scales helps to reconcile niche theory with the high species diversity observed in natural systems and helps interpret niche theory in the context of metacommunity ecology. Further work to integrate classic limiting similarity concepts with the idea of regional niche difto coexist in such a metacommunity.

ferentiation will improve our understanding of competitive metacommunities (Chesson 1991).

Toward a General Framework

Although CC and the SS models have typically been regarded as two different approaches to studying spatial coexistence, elements of both classes of mechanisms are likely to operate simultaneously in natural communities. We present several key points of contact between CC- and SS-mediated coexistence and propose the basis for a general framework.

Convergence

ence of an essential nutrient (or toxin) alters the relative strengths of intra- versus regional scale due to spatial heterogeneity in the environment. For instance, if lows spatial coexistence mechanisms such as CC and SS to be interpreted in the spatial niche differences arise because superior competitors can displace inferior competitors from occupied patches, but inferior competitors can only occupy partitioning with inferior competitors, by virtue of their superior dispersal abilities that are better at acquiring empty patches, and superior competitors that because of their superior displacement abilities are better at acquiring occupied patches. In the case of SS models, niche differences between species arise at the spatial variation in abiotic factors such as temperature, humidity, salinity or presinterspecific competition for a limiting resource in different locations of the landscape (for examples see Miller and Kneitel, chapter 5; Cottenie and De Meester, chapter 8; Kolasa and Romanuk, chapter 9), then space itself constitutes a second competition to be stronger than interspecific competition. This requirement alcontext of classical niche theory. In CC models with displacement competition, patches not colonized by the superior competitors. This is equivalent to resource As pointed out by Chesson (2000a), coexistence always requires intraspecific niche axis along which species exhibit differences.

A key issue in integrating CC and SS ideas in a common framework involves identifying the conditions under which both classes of mechanisms can contribute to coexistence. While the role of spatial heterogeneity and source-sink dynamics in allowing coexistence when life-history trade-offs cannot do so has been investigated (e.g., Yu and Wilson 2001; Levene and Rees 2002), how spatial variation influences trade-off mediated coexistence has not received much theoretical attention. For example, spatial heterogeneity in the biotic or abiotic environment can lead to spatial variation in the expression of a life-history trade-off. Contrary to the conventional wisdom that environmental heterogeneity promotes species coexistence, heterogeneity that influences the expression of a life-history trade-off can also constrain opportunities for coexistence (Amarasekare et al. 2004). In such a situation, source-sink dynamics can play a key role in enhancing opportu-

nities for coexistence. For example, consider an interspecific trade-off between recies that allocates more energy to reproduction at the cost of reduced predator defense is competitively superior to the species that employs the opposite allocation strategy. There is spatial variation in predator abundance such that it is present in some localities but not others. In localities where the predator is present, the analysis). This result is due to emigration from localities in which the trade-offis source exploitation and susceptibility to a predator (as in McPeek 1996). The spetrade-off is expressed and local coexistence is possible. In localities where the natural enemy is absent, the inferior competitor is excluded because energy allopetitive disadvantage. If there is dispersal between localities, however, coexistence expressed (sources for the inferior competitor) rescuing the inferior competitor from exclusion in localities in which the trade-off is not expressed (sinks for the cated to predator defense (a net cost now with no benefit) gives it an overall comis possible everywhere (see Amarasekare et al. 2004 for a formal mathematical inferior competitor). Thus, simultaneous operation of trade-offs and source-sink dynamics can increase opportunities for coexistence.

The models we have presented in this review have considered fundamentally different types of competitive interactions that should be carefully considered when testing spatial theory. The key distinction between dominance and preemptive competition is that, because the former involves displacement of inferior competitors by superior competitors, the ability of the inferior competitor to persist is an explicit function of both the superior competition species compete to replace individuals that have died, and no displacement occurs. The ability of any given species to persist is a function solely of its colonization and extinction rates. The key results we have reviewed in this chapter can be summarized as follows. Under dominance competition coexistence can occur either via a CC trade-off in a spatially homogeneous environment, or via SS dynamics in a spatially heterogeneous environment. When competition is preemptive, coexistence cannot occur via a CC trade-off unless there is some spatial heterogeneity and a trade-off between species competitive ability for distinct limiting resources.

Hybrid Models

The fact that CC and SS models have been considered alternative mechanisms has made it difficult to identify conditions under which elements of both mechanisms operate. This underscores the need for a more general framework that accommodates both approaches and allows predictions about the relative importance of competition-colonization and source-sink in mediating spatial coexistence in nature (Amarasekare et al. 2004).

Hybrid models have opened the way for an integrated framework (figure 10.1). The Horn and MacArthur model (1972) is a notable example because it demonstrates that a combination between CC and SS processes can explain coexistence

of Levine and Rees (2002) makes the link with SS for CC models with dominance competition but no strict asymmetry. In all of these examples, a combination of sink in a single framework and compared their operation under different types of competition and dispersal. They showed that counterintuitive properties arise between competing species. More recently Pacala and Rees (1998) proposed a cessional species have a greater ability to exploit resource-rich conditions of reson (2001) makes the link with source-sink for preemptive competition, and that competition-colonization and source-sink can elevate species richness. More recently Amarasekare et al. (2004) combined competition-colonization and sourcedue to interactions between these two different classes of mechanisms. For instance, spatial variation in the expression of a life-history trade-off (spatial hetmodel that combined competition-colonization and the successional niche. They showed that coexistence between species of different successional stages can result from competition-colonization, successional niche partitioning where early succently disturbed habitats, or a combination of the two. The model by Yu and Wilerogeneity) can constrain rather than promote coexistence.

A general framework may be obtained most easily by incorporating hierarchical spatial structure and spatial heterogeneity into the patch occupancy framework. Patch occupancy models (following Levins and Culver 1971) are attractive because of their analytical tractability for large numbers of patches, but they lack real world applications because of the unrealistic assumption of instantaneous local dynamics. This assumption has been important in developing simple metapopulation models but "there is no reason to believe that natural systems fall into two distinct classes with respect to the difference between the local and the regional time scales" (Hanski 1983). Explicit local dynamics will be required to track local abundances and details of local competitive interactions that influence coexistence and give predictions that correspond to what ecologists can measure in the field. Such hybrid models should also be able to vary the relative contribution of CC and SS mechanisms. Because the type of competition can alter the conditions for coexistence, preemptive and dominance competition should both be considered.

Several different avenues for further research have been investigated, but many more still require investigation.

- (1) Classical SS effects at the regional scale with threshold versus directional local dynamics.
- (2) Classical CC trade-offs at regional or local scales as well as a combination at different scales. Consideration of whether competition-colonization can lead to coexistence when dispersal occurs on the same timescale as competition may also be important.
- (3) A combination of CC trade-offs and source-sink at the regional scale as in the successional perspective discussed above. This may extend the conditions

- under which competition-colonization can increase opportunities for coexistence via SS dynamics.
 - ries of species diversity (Hubbell 2001; Mouquet and Loreau 2003; Chase distinct life-history strategies and source-sink between species of the same (4) Regional competition-colonization between groups of species that have very group. This point seems promising to merge the niche and the neutral theoet al., chapter 14).
- Regional source-sink between groups of species that would use very different resources and competition-colonization (either local or regional) between the species with similar resource use. (2)

Conclusions

The world is patchy and heterogeneous, the recognition of which has been central to many theories of species coexistence over the last thirty years. Only recently have ecologists integrated these ideas into the emerging field of metacommunity based either on competition-colonization or source-sink dynamics, and we have coexistence in metacommunities. We have shown that these apparently different oretical framework. We have also emphasized how these mechanisms could be ecology. In this chapter we have focused on the long history of spatial models discussed how these two approaches provide alternative mechanisms for species mechanisms have common elements that could be integrated into a broader thebroaden the niche concept to include regional phenomena. Our focus was some-1997a; Nee et al. 1997; Hassell 2000; Hoopes et al., chapter 2; Holt and Hoopes, chapter 3) or genetic polymorphisms (Karlin and McGregor 1972a, 1972b; De interpreted in the context of niche theory and how this interpretation can help what restrictive in that we concentrated exclusively on competitive metacommunities with no consideration of trophic interactions (Nowak and May 1994; Holt geneous environments would require assembling these other pieces of the same vironments will allow for the development of a general theory of biological diver-Meeus et al. 1993), both of which are integral components of biological diversity in patchy environments. A complete synthesis of coexistence in spatially heteropuzzle. Understanding mechanisms that maintain species diversity in patchy ensity that is needed now more than ever, given the catastrophic loss of biodiversity due to increasing human encroachment.

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CHAPTER 11

Assembly Dynamics in Metacommunities

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Introduction

Much of the theory of community ecology is concerned with dynamics of interacting species at a local level—how interactions among the species determine transient and asymptotic properties of communities. Best known is the generalized Lotka-Volterra model, although there are numerous elaborations on this as well. However, communities are open systems liable to be invaded and changed by species from outside; structure at a local level may depend as much on the pool of species available in the region as on local interactions (Cornell 1985; Ricklefs 1987; Leibold et al. 1997; Shurin et al. 2000). How can we extend the theory of local communities to this larger spatial scale? And how might mechanisms for species interactions at this large scale result in dynamics different from those at the local scale?

One response to the openness of communities has been to develop a theory of assembly dynamics; this invokes an external pool of species that from time to time contributes new species to the local community (Post and Pimm 1983; Rummel and Roughgarden 1985; Mithen and Lawton 1986; Drake 1990; Law and Morton 1996; Lockwood et al. 1997). New species may get established, thereby changing the structure of the local community, and may bring about further change by causing extinction of species in the community. For a finite species pool, an assembly graph can be constructed describing the sets of species that can persist locally and the transitions between these sets caused by arrival of new species (Law and Morton 1996; Warren et al. 2003; Zimmermann et al. 2003). Assembly dynamics then use this graph to gain insight into assembly patterns that emerge through random arrival of species (Law 1999). In the long term assembly leads to uninvasible communities (endpoints) or cyclic sequences of communities, uninvasible by species not present within the sequence (endcycles) (Morton and Law 1997).

In many cases however, species pools are better thought of as the union of sets of species present in the local communities scattered across a region. For the purpose of considering assembly dynamics, communities might be better thought of as embedded in a metacommunity of many local communities linked by dispersal (Leibold et al. 2004). This couples the regional species pool to the local communities so that, in the long term, the species pool contains only those species that