I. INTRODUCTION

Community ecology aims at shedding light on how competing species assemble and coexist in their hosting habitats \[1\]. This has proven to be a formidable challenge. A main reason is that ecological dynamics spans a wide range of spatial and temporal scales, from those typical of individuals to those characterizing large populations or communities. For a long time, ecologists have empirically characterized biodiversity at the different spatial scales; for example, counting the average number of species hosted in a given area – \( \text{species area relationship (SAR)} \) \[2, 3\] –, or the distribution of their abundances – \( \text{species abundance distribution (SAD)} \) \[4, 5\]. Often, the ecological forces determining these patterns act at a given spatio-temporal scale but can affect others as well. The inverse problem, i.e. linking observed ecological patterns with the causes originating them at different scales, is arguably the central problem in ecology \[6\].

This kind of problem sounds familiar to experts in statistical physics, where large-scale emergent behaviors result from interactions among simple local units. Tools of statistical physics are indeed extremely useful to shed light on this crucial issue in ecology. In particular, a natural approach to such complex problems is to drastically simplify them. To this aim, we consider ecosystems made up of competing non-motile species, such as trees, or having a motility range much smaller than the typical size of the population, such as many communities of microorganisms. In this case, a possible simplifying assumption is that all emergent phenomena originate at the single-individual scale and, more drastically, that differences among individuals, possibly belonging to different species, can be neglected. These simplifying assumptions constitute the basis of the ecological neutral theory proposed by Hubbell \[7\].

Ecological neutral theory \[8\] was built upon theoretical ideas of Kimura’s neutral theory of population genetics \[8\]. In particular, both theories underscore the role of stochastic demographic fluctuations in determining the fate of populations and completely neglect deterministic effects stemming from fitness differences. Ecological neutral theory has elicited heated controversies, as it hinted that classical ecological concepts, such as niches, might play a marginal role in structuring communities of competing species. Despite these contentions, neutral theory had a considerable impact on ecological thinking, owing to its ability to quantitatively predict non-trivial patterns of biodiversity with simple models and very few adjustable parameters \[9–11\].

Spatially implicit neutral models describe well-mixed communities of individuals subject to immigration from a larger reservoir of species where diversity is maintained via speciation. They can be solved analytically \[12–16\], yielding analytical expressions for the SAD. Beside the mathematical appeal, these exact solutions have been extremely helpful for fitting empirical data and therefore testing neutral theory or, at least, promote it as a null-model \[17\]. For an exhaustive survey of ecological neutral theory, we refer the reader to Hubbell’s book \[7\] and reviews \[9–11\].

The focus of this review is on spatially-explicit neutral and near-neutral population models. Explicitly describing space is crucial to address the fundamental ecological questions sketched at the beginning of the introduction.
However, spatially-explicit models are still poorly understood, especially if compared with their well-mixed counterparts. These models are often variants of familiar models in non-equilibrium statistical physics. One of the most studied neutral model is the voter model with speciation, or multi-species voter model, which generalizes the more familiar two-species voter model. The stepping-stone model and the contact process are other examples of spatial models that have been studied in parallel in the physics and population biology literature. We shall discuss how these analogies can be used to advance our understanding of spatial ecology and the main open problems. This review heavily relies on extensive numerical computations of lattice models based on previous works by the authors. This might have biased the choice of some topics and we apologize if some relevant works are not properly discussed.

The review is organized as follows. In Sect. II we introduce the multispecies voter model on a lattice and its dual representation in terms of coalescing random walkers. We then discuss its predictions of macroecological patterns: the SAR, and the SAD. For the latter, we compare two recent analytical approaches with novel computational results. We mainly address the two-dimensional case for its ecological relevance, but at the end of the section we also discuss the one-dimensional case. In Sect. III we discuss other neutral models, where, unlike in the voter model, lattice sites are not necessarily occupied by exactly one individual at all times. In particular, we consider the stepping stone model, where each lattice site hosts a local community of individuals, which is relevant for modeling microorganisms and their associated macroecological patterns. We then consider a multispecies variant of the contact process, where lattice sites can be either empty of occupied by single individual. In Section IV we introduce non-neutral effects on a simplified two-species competition model, where adjusting a single parameter one can tune the departure from neutrality, here modeled as a specific habitat preference. Physically, this habitat preference can be thought as a form of quenched disorder. We discuss how this disorder generically favors species coexistence using the language of statistical mechanics, and also discuss other forms of disorder such as temporal heterogeneity. Finally, Sect. V is devoted to perspectives and conclusions.

II. VOTER MODEL WITH SPECIATION

A. Description of the model

A paradigmatic example of spatial neutral model is the voter model with speciation, sometimes also called multispecies voter model, which is a generalization to more than two species of the voter model. The latter is a widely studied model that has been used, and often proposed independently, in diverse contexts, from population genetics to spatial conflicts, spreading of epidemic diseased, opinion dynamics and linguistics.

The voter model with speciation is defined on a lattice, where each site hosts one individual belonging to some species. At each discrete time step, a lattice site is chosen at random and the residing individual is removed (death event). Then, as illustrated Fig. 1A, the dead individual is replaced:

- With probability $\nu$, by an individual of a new species not present in the system (speciation event). Notice that, because of speciation, the total number of species is not fixed. In population genetics, this type of event is interpreted as a mutation within the same species.

- With complementary probability $(1-\nu)$, by a new individual of an existing species (reproduction event). In this case, the newborn belongs to the same species of another (parent) individual chosen at random in the neighborhood of the vacant site. In the simplest case, the nearest-neighbors (NN) are chosen with uniform probability. More generally, the parent individual is selected according to a probability distribution $P(\vec{r})$ (the dispersal kernel) over the neighbors within a distance $\vec{r}$.

![FIG. 1: Examples of transitions in the 2D voter model with speciation.](image)

Most of this section will be devoted to the ecologically relevant case where the system is a two-dimensional (2D) square lattice. At the end of the section, we will briefly present the main results in 1D for comparison.

B. Duality

The voter model with speciation is dual to a system of coalescing random walkers with an annihilation rate. The dual process is constructed as follows. We start by placing on each lattice site a random walker.

The dynamic of the dual process proceeds backward in time. At each discrete (backward) time step, with probability $1-\nu$, a randomly chosen walker is moved to a new site, where the dispersal kernel $P(\vec{r})$ here plays the role of the distribution of possible displacements. If the...
Duality is a very useful property to understand the physics of the voter model. For example, it immediately stems from duality that the $\nu \to 0$ limit is fundamentally different in $D \leq 2$ and $D > 2$. As a matter of fact, in $D \leq 2$ the random walk is recurrent, meaning that the probability of two randomly chosen individuals to belong to the same species approaches one as $\nu \to 0$; in other words, in the absence of speciation, one expects monodominance of one species in the long term. The same property does not hold in $D > 2$, where random walkers are not recurrent and one expects coexistence of species even in the limit $\nu \to 0$. Interestingly, the ecologically most relevant case, $D = 2$, is the critical dimension of this model. We shall see that this fact is a source of a number of non-trivial features for ecologically relevant quantities.

Duality is also an extremely powerful tool for computational analyses. If one is interested in the static, long-term, properties of the voter model with speciation, it is numerically much more efficient to simulate the dual dynamics than the forward one. In a dual simulation, after all walkers coalesced or were annihilated, species can be assigned to the start site of each walker, obtaining a stationary configuration of the voter model. Beside computational speed, this approach has also the advantage of eliminating finite-size effects induced by the boundary conditions, as the coalescing random walkers can be simulated in a virtually infinite system. For illustrative purposes, in Fig. 2c and 2d we show two configurations of the 2D voter model obtained with the dual dynamics for two different dispersal kernels.

C. $\beta$–diversity

The first ecological pattern we consider is the so-called $\beta$-diversity, which is a measure of how the species composition in an ecosystem varies with the distance. The $\beta$-diversity is defined as the probability, $F(\vec{r})$, that two randomly chosen individuals at a distance $\vec{r}$ are conspecific, i.e. belong to the same species. Mathematically, this quantity can be written in terms of the two-point correlation function $G_{s_i,s_j}(\vec{r}) = \langle n_{s_i}(\vec{x})n_{s_j}(\vec{x} + \vec{r}) \rangle$, where $n_{s_i}(\vec{x})$ denotes the number of individuals of species $s_i$ at location $\vec{x}$. Indeed, one can easily show that

$$F(\vec{r}) = \frac{\sum_i G_{s_i,s_i}(\vec{r})}{\sum_{i,j} G_{s_i,s_j}(\vec{r})},$$

where the sums extend over all species in the ecosystem. Eq. (1) expresses the $\beta$-diversity as the ratio between the couples of conspecific over the total number of couples in the sample.

Let us now study the evolution equation of $F(\vec{r},t)$ for the voter model with speciation and NN dispersal. Although we shall focus on the 2D case, for later convenience it is useful to present the general calculation in $D$ dimensions. Following [11, 41, 42] we write

$$F(\vec{r},t+1) = \left(1 - \frac{2}{N}\right) F(\vec{r},t+1) +$$

$$+ \frac{1 - \nu}{DN} \sum_{k=1}^{D} [F(\vec{r} + \vec{e}_k,t) + F(\vec{r} - \vec{e}_k,t)].$$

The first term in the r.h.s. of Eq. (2) represents the fact that $F$ does not change if two generic individuals at distance $\vec{r}$ survive at a single time step, i.e. they are not selected for updating. The second term represents the events in which one of the two individuals dies (with prob. $2/N$), no speciation occurs (with prob. $1 - \nu$) and the dead individual is replaced by a conspecific from the 2D neighbor sites. Taking the continuous limit ($N \to \infty$ with the lattice spacing $a \to 0$ and the speciation
probability $\nu \to 0$) with $k^2 = 2D\nu/a^2$ finite, one obtains at stationarity the differential equation

$$
\frac{1}{r^{D-1}} \frac{d}{dr} r^{D-1} \frac{dF}{dr} - k^2 F(r) + cD(r) = 0 \tag{3}
$$

where $\delta^D$ is the $D$-dimensional Dirac delta, and because of isotropy the $\beta$-diversity $F(r)$ is now function of $r = |\vec{r}|$ only. The solution of Eq. (3) is $[11]$

$$
F(r) = \frac{\kappa^{D-2}}{(2\pi)^{D/2}} (\kappa r)^{(2-D)/2} K_{(2-D)/2}(\kappa r), \tag{4}
$$

where $K_0$ is the modified Bessel function of the second kind of order $z$ and the constant $c$ is fixed by the requirement $\int_{r_0}^{a} d^D r F(\vec{r}) = 1$. We recall that Eq. (4) is a continuous expression, valid for distances much larger than the lattice spacing $[11]$. Although we derived Eq. (4) for NN dispersal, the same results hold for a general dispersal kernel for distances larger than the kernel range, provided that the kernel range is finite.

For $D = 2$, Eq. (4) implies that $F(r) \propto K_0(\kappa r)$, which is characterized by a slow logarithmic decay, $\sim -\ln(\kappa r)$, up to distances of order $1/\kappa \sim 1/\sqrt{\nu}$, followed by a faster, exponential falloff. Remarkably, the $\beta$-diversity empirically measured in several tropical forests in Central and South America is consistent with a logarithmic decay for large distances $[13]$. We remark that this logarithmic decay is the signature that $D = 2$ is the critical dimension for the voter model. In contrast, in $D = 1$, Eq. (4) becomes $F(r) \propto \sqrt{\pi \nu} K_{1/2}(\kappa r) \sim \exp(-\kappa r)$.

Although the $\beta$-diversity decays exponentially on scales $1/\kappa \sim 1/\sqrt{\nu}$ both in $1D$ and $2D$, there are important differences. Because $2D$ is the critical dimension, a large biodiversity (i.e. a large average number of species) can be sustained by very low values of the speciation rate $\nu$. This implies that in $2D$ there are many species living on scales much smaller than $1/\kappa$, where the correlations decay logarithmically. Conversely, in $1D$ to maintain biodiversity one needs a large value of $\nu$, so that $1/\kappa$ is the only characteristic scale and there is no additional structure on scales smaller than $1/\kappa$. This crucial point will be further elucidated in the rest of the section, where we will discuss other observables in $2D$ (subsections II D and II E) and compare them with their $1D$ counterparts (subsection II E).

D. Species-Area Relationships

We now focus on the SAR, defined as the average number of species, $S$ of a given taxonomic level occupying a given area of size $A$. SARs are widely studied in ecology as a measure of ecosystem richness and quantify how larger habitats support more species than smaller ones $[2]$. Empirical measures of SARs at multiple scales often reveal three different regimes $[2, 3, 7]$. At small areas, the number of species increases rather steeply, nearly linearly, with the sampled areas. A similar steep increase is observed at very large, continental scales. Instead, at intermediate scales, a slower, sublinear growth is often observed. Such a growth is well approximated by a power law $S \sim A^z$, $z < 1$, over a wide range of taxa $[44]$, though a logarithmic behavior $S \approx C \ln A$ has also been proposed. An extensive meta-study by Drakare et al. $[45]$ reconsidered a large body of SAR studies from the literature, revealing that the power law provides a better fit in half of the cases. This study also observed that the exponent $z$ correlates positively with the body size of the considered group of species, so that small microorganisms typically display very shallow SAR curves as compared with larger organisms, see also $[46]$ and Sect. III A.

Simulations of the (dual) voter model with speciation yields SARs, shown in Fig. 3a, qualitatively similar to those obtained from field data. In the voter model, the steep initial regime is mostly determined by the dispersal range $K$. For areas significantly larger than $K^2$, a sublinear growth is observed (see Fig. 3b). The growth becomes progressively more shallow as the speciation rate $\nu$ is decreased. For larger scales, the slope of the SAR curves become steeper again. The area at which this final crossover occurs increases as $\nu$ is decreased.

An interesting question is whether the sublinear growth regime in the voter model can be characterized by a power-law $S \sim A^z$ and, in this case, what is the value of the exponent $z$ as a function of $\nu$. To this aim, we review a classic estimate of $z$ by Durrett and Levin $[20]$ based on the duality of the voter model (see Sect. III B). The speciation rate $\nu$ sets a time scale $1/\nu$ which also corresponds to a characteristic length scale $\xi = 1/\sqrt{\nu}$ because of the diffusive behavior of random walkers. Notice that walkers with an initial separation much larger than $\xi$ are likely to be eliminated by the death rate before coalescence occurs. This observation alone explains the linear scaling of $S(A)$ for areas $A \gg \xi^2 = \nu^{-1}$. At these scales, species are uncorrelated, as can also be inferred from the analysis of $\beta$-diversity in the previous section. For a system of coalescing random walkers in $2D$, the density of occupied sites $\rho(t)$ decays asymptotically as $[47, 48]$

$$
\rho(t) \sim \frac{\ln t}{\pi t}. \tag{5}
$$

The logarithmic coarsening observed in the two-species voter model without speciation can be related to the logarithm appearing in Eq. (5) $[49]$. Assuming $\nu \ll 1$, the annihilation rate at time $t$ in an area $\xi \times \xi$ can be approximated as the annihilation rate per walker $\nu t$ times the number of walkers in the absence of annihilations $\xi^2 \rho(t)$. Integrating over time, we find that the total number of annihilations, i.e. the total number of species, is $[50]$

$$
S(\xi^2) \approx \nu \xi^2 \int_{t_0}^{1/\nu = \xi^2} dt \rho(t) = 
= \frac{\ln^2(\xi^2) - \ln^2(t_0)}{2\pi} \approx \frac{2}{\pi} (\ln \xi)^2, \tag{6}
$$

where $t_0$ is the time at which the asymptotic expression $[5]$ starts to be valid. The upper temporal cut-off is set to
1/ν (with 1/ν = ξ²) because the number of killing events occurring after a time ~ 1/ν is bounded by the number of walkers in the system, which is ξ²ρ(1/ν) ~ ln ξ [32]. Finally, combining Eq. (6), the fact that S(1) = 1 and matching a power law behavior S = A² in the range of scales from A = 1 to A = ξ², one finds [20]

\[
z = \frac{\ln[S(A)]}{\ln(A)} = \frac{2\ln[\ln(1/\sqrt{\nu})] + \ln(2/\pi)}{\ln(1/\nu)}.
\] (7)

Also in this case, the logarithmic dependence of the exponent z on ν derives from the fact that D = 2 is the critical dimension for the voter model.

More recent results disputed the validity of Eq. (7). Scaling arguments hinted that z should approach a finite value z ≈ 0.2 in the limit of vanishing ν (see [42] and Sec. II E 1), while numerical simulations suggested a power law dependence z ~ ν₀.15 [21]. Finally, further numerical simulations [22], based on the dual representation of the voter model with speciation (see Sect. II B) and spanning a very wide range of speciation rates from 10⁻³ to 10⁻¹¹ confirmed the logarithmic behavior predicted by Eq. (7). The exponents measured in such simulations, shown in Fig. 3, are well fitted by a phenomenological expression of the form

\[
z = \frac{1}{q + m \ln(\nu)}
\] (8)

which is consistent with Eq. (7) up to order ln ln ν, see also Fig. 3. However, fitted values of the prefactors q and m are not consistent with Eq. (7). This discrepancy is probably due to pre-asymptotic effects as well as to the approximation of assuming a power-law range between A = 1 and A = ln(1/ν).

We conclude with a brief discussion of the role of the dispersal kernel. The comparison between NN dispersal and a square dispersal kernel of range K = 7, shown in Figs. 3 and 31 demonstrates that the exponent z depends to some extent on the dispersal kernel. However, numerical evidence [21] suggests that when the dispersal kernel range is large enough (approximately K ≥ 5) the exponents are very weakly dependent on K. Moreover, SARs obtained with different values of K can be rescaled onto a universal function of A and ν only via the transformation S = f(A, ν, K) = K²φ(A/K, ν) with a fitted value of χ ≈ 1.97. This small departure from the trivial exponent χ = 2 suggests, also in this case, the presence of logarithmic corrections. To the best of our knowledge, a formal derivation of this scaling law and of the exponent χ is currently an open problem.

E. Species-Abundance Distributions

We now discuss Species-Abundance Distributions (SADs), P(n; A), that measure the relative abundance of species in a given area A. More precisely, denoting S(A) the total number of species sampled in an area A, each composed by n_i (i = 1, ..., S(A)) individuals, P(n; A)dn is the probability that a randomly picked species has an abundance between n and n + dn. While the expression of P(n; A) for well-mixed neutral models is known [14], computing it for spatially explicit models, such as the voter model with speciation, has proven to be a rather hard problem. We first discuss in section II E 1 an approach based on scaling considerations, and underline its limitations. In Sec. II E 2 we discuss how this approach can be generalized at the critical dimension, present numerical results, and discuss a recent attempt to compute P(n; A) exploiting duality.

1. Power-law scaling relation

In the voter model with speciation, the SAD is not only a function of the system size A, but also of the speciation rate ν. Although we are mainly interested in 2D, it is instructive to consider the general case in which A = L², where L is the linear size of the sample. Following [11] [12], we assume a standard scaling form for the SAD

\[
P(n; A, ν) = n^{-β}\Psi(nν^α, Aν^{D/2})
\] (9)

where the exponents α and β remain unspecified for the time being, whereas the exponent D/2 stems from the
diffusive nature of neutral models \( \nu \sim t^{-1} \sim L^{-2} \sim A^{-2/D} \). Note that in models with long-range, non-diffusive dispersal \footnote{\textbf{[50]}} the scaling form might differ. Equation \( \textbf{(10)} \) describes a power-law dependence of \( P \) on \( n \) that, as usual for scaling laws, holds up to a scale that is described by the scaling function \( \Psi \), that depends on dimensionless combinations of the population size \( n \), the speciation rate \( \nu \), and the system size \( A \). To the best of our knowledge, there is no available analytical prediction for the exponent \( \beta \). For the exponent \( \alpha \), we recall that in the dual formulation of the voter model with speciation the population size \( n \) is the number of coalescences that occur before a annihilation \footnote{\textbf{[48, 51]}}. This implies that \( \alpha \) characterizes the temporal decay of the density of coalescing random walkers, \( \rho(t) \sim t^{-\alpha} \). However, as shown in \footnote{\textbf{[48, 51]}} \( \rho(t) \) decays as \( \rho(t) \sim t^{-\min(1,D/2)} \) for \( D \neq 2 \) and \( \rho(t) \sim \log(t)/t \) in \( D = 2 \). Consequently, one should expect the power-law scaling of Eq. \( \textbf{(10)} \) to work in \( D = 1 \) and \( D \geq 3 \), but not at the critical dimension \( D = 2 \), where logarithmic corrections should appear. In the following, we discuss how such corrections can be taken into account in a generalized scaling framework.

2. Generalized scaling relation

In order to allow for logarithmic corrections, Zillio et al. \footnote{\textbf{[30]}} proposed to employ the generalized scaling relation

\[
P(n; A) = g(A) \Psi(n/f(A)).
\]

The dependence on \( \nu \) was omitted as the above scaling law was applied to observational data for which the speciation rate is unknown and assumed to be fixed. The key aspect of Eq. \( \textbf{(10)} \) is that \( f \) and \( g \), are general functions and not necessarily power-laws as in conventional scaling, allowing for the possibility to include logarithms or other functional dependencies. On the contrary, the scaling function \( \Psi(x) \) is still assumed to be a power law

\[
\Psi(x) \sim x^{-\Delta}
\]

for \( x \sim 0 \), where \( \Delta \) is an exponent to be determined. Thus, also Eq. \( \textbf{(10)} \) postulates a power-law dependence on \( n \), but with a more general cut-off for large areas. After specifying the functions \( f \) and \( g \), Eq. \( \textbf{(10)} \) can be tested by plotting \( P(n; A)/g(A) \) versus \( x = n/f(A) \) for a set of different areas and assessing the quality of the data collapse onto a single curve, \( \Psi(x) \).

To determine the functions \( f \) and \( g \), we impose that \( P(n; A) \) needs to be normalized, \( \int_0^\infty dn \ g(A) \ \Psi(n/f(A)) = 1 \), and that its average value is \( \langle n \rangle = \int_0^\infty dn \ n \ g(A) \ \Psi(n/f(A)) \). Substituting the scaling form \footnote{\textbf{[11]}} into these equations, it is possible to derive conditions that the functions \( f \) and \( g \) must obey, depending on the value of \( \Delta \). In particular, as shown in Appendix A, the case \( \Delta = 1 \) is marginal and needs to be treated with care (other values \( \Delta \neq 1 \) are explicitly analyzed in the Appendix). Approaching it as \( \Delta = 1 - \epsilon \) with \( \epsilon \ll 1 \), Eq. \( \textbf{(11)} \) becomes

\[
\Psi(x) = x^{-1+\epsilon} \sim \frac{1}{x} \left[ \exp(\epsilon \ln(x)) \right] \sim \frac{1}{x} [1 + \epsilon \ln(x)]
\]

up to first order in \( \epsilon \). At the same order in \( \epsilon \), the conditions for \( P \) become \( 1 \sim g(A)f(A) \ln(f(A))[1 + \frac{\epsilon}{2} \ln(f(A))] \) and \( \langle n \rangle \sim g(A)f(A)^2 \) from which we finally obtain

\[
f(A) = \langle n \rangle \ln\langle n \rangle \left[ 1 + \frac{\epsilon}{2} \ln\langle n \rangle \right]
\]

\[
g(A) = \frac{1}{\langle n \rangle \ln^2\langle n \rangle \left[ 1 + \frac{\epsilon}{2} \ln\langle n \rangle \right]}
\]

up to first order in \( \epsilon \). Notice that for values of \( \Delta \) close to 1– both functions \( f \) and \( g \) include logarithmic corrections. By means of a similar calculation, one can estimate the \( k \)-th moment \( \langle n^k \rangle \), and verify that all the moment ratios \( \langle n^k \rangle / \langle n^{k-1} \rangle \) scale in the same way, up to a multiplicative constant

\[
\frac{\langle n^k \rangle}{\langle n^{k-1} \rangle} = \frac{\int dn \ n^k \ P(n; A)}{\int dn \ n^{k-1} \ P(n; A)} \propto f(A) \quad k \geq 1.
\]

revealing a highly anomalous scaling.

Zillio et al. \footnote{\textbf{[30]}} showed that this scaling form provides a much better collapse of empirical data from the Barro Colorado tropical forest than a power-law scaling relation such as Eq. \( \textbf{(9)} \). This supports the idea that \( \Delta \) is close to its marginal value 1 in tropical forests.

We tested computationally whether Eqs. \( \textbf{(10)} \) and \( \textbf{(13)} \) provide a good collapse of SADs from the voter model with speciation and whether the relationship between...
FIG. 5: Moment ratios for different values of \( k \). As predicted by Eq. (10), in the case \( \Delta = 1 \) all moment ratios \( \langle n^k \rangle / \langle n^{k-1} \rangle \) scale in the same way with \( f(A) \) up to a multiplicative constant. As in Fig. 4, the fitted value is \( \epsilon = 0.08 \).

the moments (14) holds. In simulations, an additional parameter is the speciation rate \( \nu \). As discussed above, \( \nu \) appears in scaling relationships via the dimensionless combination \( A \nu D/2 \), that in 2D equals \( A \nu \). Thus, although Eqs. (10) and (14) do not include speciation explicitly, we expect these relationships to hold if \( A \nu \) is kept constant. We therefore performed computational analyses fixing \( A \nu = 200 \), although the conclusions are robust against this choice. Results are summarized in Figure 4 which shows plots of the SAR, for systems with different linear size, \( L \) and different speciation rates \( \nu \) (with \( L^2 \nu = A \nu = 200 \)). Observe in Fig. 4 that the smaller the size (or the larger the speciation rate) the smaller the maximal abundance. Figure 4 show the data collapse as given by Eqs. (10) and (13), where \( \langle n \rangle \) is the average number of individuals measured in each area \( A \) and \( \epsilon \) is a free parameter that we fitted obtaining \( \epsilon = 0.08 \) and an almost perfect collapse of 7 different curves. The small value of \( \epsilon \), is consistent with the assumed small deviation from \( \Delta = 1 \). A similar collapse for \( A \nu = 20 \) leads to an even smaller value \( \epsilon \approx 0.069 \) (not shown). We verified that either removing all logarithmic corrections (thus plotting results as a function of \( \langle n \rangle \)) or simply fixing \( \epsilon = 0 \) in Eq. (10) and (13) leads to less convincing collapses, as shown in Fig. 3 and 4, respectively. Clearly, these deviations can pass unnoticed in the presence of statistical fluctuations. Probably, this is the reason why in [52] a simple scaling law was claimed to work for the 2D voter model with speciation. Finally, we also verified that moment ratios scale as \( f(A) \), as predicted by Eq. (14), as illustrated in Fig 5.

In summary, a non-standard scaling form, including logarithmic terms—"a priori expected for scaling theories at the critical dimension—provides an excellent collapse both for empirical data and for a simple 2D neutral model. However it is interesting to remark that the data collapsed in [50], were obtained from tropical forests of different areas \( A \). It is reasonable to assume that the speciation rate \( \nu \) do not vary much among these forests. Therefore, the product \( A \nu \) is not fixed, as in our computational analyses. A possible explanation is that, although the collapse achieved in this way is not perfect, the deviations from perfect scaling are too small to be appreciated in observational data due to the limited sample size. We have verified in simulation that, as a matter of fact, keeping \( \nu \) constant (rather than \( A \nu \)), small deviations from perfect collapse are observed.

We conclude this section mentioning that, remarkably, a heuristic expression for the SAR has been recently derived for the voter model with speciation following a completely different approach. [31, 32]. Let us define \( P(x,t) \) as the distribution of the number of individual of a given species at time \( t \). If we approximate \( x \) as a continuous quantity, we can heuristically write a Fokker-Planck equation for the evolution of \( P(x,t) \)

\[
\partial_t P(x,t) = \nu \partial_x [xP(x,t)] + \partial_x^2 [I(x)P(x,t)]
\]

(15)

where the first term in the right hand side is the negative drift due to speciation, and the second is the fluctuation in population size, where \( I(x) \) is the average number of interfaces of a species of size \( x \). The crucial underlying approximation is to neglect fluctuations of \( I(x) \). This approximation is appropriate if the distribution of the number of interfaces at fixed value of \( x \) is a very peaked function. In this simple framework, all the dependence on the spatial dimension of the voter model is recap into the function \( I(x) \). The steady-state solution of Eq. (15) is

\[
P_{st}(x) = \frac{e^{-\nu \int dx \frac{x}{I(x)}}}{I(x)}.
\]

(16)

From duality considerations [31, 32], the average number of interfaces must scale in 2D as \( I(x) = x/(1 + c \ln x) \) where \( c \) is a non-universal constant. Notice how the expression of \( I(x) \) includes familiar logarithmic terms. Substituting this expression into Eq. (10) leads to an explicit expression for the SAR, which obeys a scaling law with logarithmic corrections similar to Eq. (14), though not identical. A careful comparison between this result and the previous scaling form is an interesting issue, but beyond the scope of this review.

F. One-dimensional case

Many of the non-trivial properties of the voter model with speciation discussed so far are specific to the critical dimension \( D = 2 \). To further illustrate this fact, it is useful to briefly compare with the \( D = 1 \) case, which is also relevant to describe quasi one-dimensional ecosystems, such as river basins [53]. For simplicity, we limit ourselves to the case of NN dispersal.

To the best of our knowledge, even in \( D = 1 \) an exact expression for the SAR, i.e. the average number of species in a segment of length \( L \), is not known. To find
an estimate, we follow an approach similar to that leading to Eq. (9). In $D = 1$, the density of walkers behaves as $\rho(t) \sim 1/\sqrt{t}$, to be contrasted with the 2D expression of Eq. (5). Dimensional arguments then suggest that the average number of species must a function of $L\sqrt{D}$ only, $S(L; \nu) = \Psi(L\sqrt{D})$. This argument is well supported by computational results, see Fig. 6, and inset. The function \( \psi \) is linear for large arguments, with a coefficient around 1.2 and it is nearly constant for arguments $L\sqrt{D} \ll 1$. This means that the non-trivial power-law regime characteristic of 2D SARs is absent in $D = 1$, where species have a characteristic size $1/\sqrt{D}$, as discussed at the end of Sect. II C. In $D = 1$ with NN dispersal, Eq. (8) for the probability that two individuals at distance $r$ are conspecific $\beta$-diversity can be solved (see [14]) even without the continuous approximation, obtaining

\[
F(r) = \exp(-\alpha(\nu)r) \quad \text{with} \quad \alpha(\nu) = \ln \left[ \frac{(1-\nu)}{(1-\sqrt{D}\nu(1-\nu))} \right],
\]

(17)

where $\alpha(\nu) \approx \sqrt{(2\nu)}$ for $\nu \to 0$.

Knowing $F(r)$, it is possible to derive an upper bound to the average number of species. In $D = 1$, individuals are organized in $N_s(L; \nu)$ segments of conspecific individuals on average, so that $S \leq N_s$, with the equality holding if no species is present in more than one segment. We can compute $N_s$ from the probability $P_{i-1,j} = F(|i-j|)$ that two sites $i$ and $j$ are occupied by conspecific individuals [54], so that

\[
N_s(L) = L - \sum_{i=1}^{L-1} P_{i-1,j} = L - (L - 1)F(1) = L - (L - 1)e^{-\alpha(\nu)}
\]

(18)

which for $\nu \to 0$ can be approximated as $N_s \approx 1 + \sqrt{2\nu}(L-1)$.

To obtain a lower bound, let us denote with $n_i(L)$ the frequency of species represented by the individual in site $i \in [0, L - 1]$. It then follows from Jensen’s inequality (see also [54])

\[
S = \sum_i \frac{1}{n_i(L)} \geq \sum_i \frac{1}{\langle n_i(L) \rangle},
\]

(19)

where $(n_i) = \sum_j P_{i,j}$, with $P_{i,j} = F(|i-j|)$ given by Eq. (17), can be easily summed numerically.

In Fig. 6, we compare the numerically obtained SAR with the upper [18] and lower [19] bounds. Notice that the upper bound is very close to the actual SAR, implying that most species are organized in single segments.

A comparison between SARs in 2D and 1D reveals a similar scenario: although there is no explicit solution available, the scaling properties of SARs in 1D are much simpler to the 2D case, because of the lack of logarithmic corrections. In Fig. 7, we show that employing the naive scaling form $P(n; L) \sim n^{-\nu}$, vs. $n/\langle n \rangle$ (derived in Appendix A for the non-marginal case $\Delta \neq 1$), we obtain a perfect collapse, to be contrasted with the deviations observed in Fig 4 for the 2D case.

III. OTHER NEUTRAL MODELS

In the voter model with speciation, the habitat is saturated and each site is always occupied by an individual. In this section, we study neutral spatial models where the number of individuals that can live in a site site is varied. We consider three variations variations: the stepping-stone model with speciation, where each site can host many individuals but the landscape remain saturated; the contact process with speciation, where occupancy is limited to a maximum of one individual per site, but sites can also be empty; and the O'Dwyer-Green model, where occupancy is unbound.

A. Stepping-Stone Model with speciation

In the voter model, each lattice site hosts a single individual, which is appropriate when modeling big sessile species, such as trees, each occupying a well-defined area and exploiting its corresponding local resources. On the other side of the spectrum, microorganisms, such as small eukaryotes or bacteria, are often present in very large numbers on tiny spatial scales, where all individuals share the same resources. For these species, it is more
appropriate to think of the habitat as subdivided into small patches, connected by migration and each hosting a large number of individuals directly competing with each other [53]. Aimed at modeling such ecological cases, in this section we consider the stepping-stone model [25, 26] with speciation, which generalizes the voter model with speciation to the case in which each site can host a fixed number $M$ of individuals.

Similar to the voter model with speciation, at each time step an individual is randomly chosen and killed. With probability $\nu$, it is replaced by an individual of a novel species. With complementary probability $(1-\nu)$, a reproduction event occurs. The parent of the new individual is selected with probability $(1-\mu)$ among the surviving $M-1$ individuals present at the same site, and with probability $\mu$ among the $M$ individuals in a randomly chosen neighboring patch (according to a probability distribution on the neighbors $P(\hat{r})$, similar to the case of the voter model). The particular case of $M = 1$ reduces to the voter model with speciation up to a time rescaling, as at each time step spatial diffusion occurs with probability $\mu$. Like the voter model, the stepping-stone model admits a dual representation in terms of coalescing random walkers with annihilation/death, which can be exploited for efficient numerical simulations. The main difference with the voter model is that, in the dual stepping-stone model, at each step a random walker can either move to another site or stay in the site of origin. Coalescence can happen in both circumstances, corresponding to reproduction of an individual from neighboring sites or from the same site. For full details on the implementation we refer to [27].

As revealed by numerical simulations of the stepping-stone model based on the dual representation, SARs are qualitatively similar to those of the voter model, although the exponents $z$ are, in general, smaller than in the voter model [24]. In particular, the exponent depends not only on $\nu$, but also on the combination of parameters $M \mu$, which determines the regimes of the model. For $M \mu \ll 1$, each local site is likely to contain only one species. In this limit, each site behaves as one individual up to a time rescaling, so one should expect the same exponents as in the voter model with speciation. In the opposite limit $M \mu \gg 1$, there is a large diversity of species at each site. An analytical argument suggests that, in this latter limit, the exponent should be a factor two smaller than in the former limit [27]. Let us study this limit in the dual representation. Since random walkers in the same site have a low probability of coalescence, they will wander for a long time before coalescing. Therefore, we can assume that, asymptotically, they will behave as in the well-mixed case. This implies that their density in an area smaller or equal than $\xi^2$ approximately decays according to the mean-field formula

$$\rho(t) \sim \frac{1}{t}.$$  \hspace{1cm} (20)

Observe that in this case the characteristic length is $\xi = \sqrt{\mu/\nu}$, as random walks diffuse with probability $\mu$ at each time step. Proceeding as in Eq. (3), the average number of species in an area $\xi^2$ can be estimated as

$$S(\xi^2) \sim \nu M \xi^2 \int_{t_0}^{t} \frac{\mu/\nu = \xi^2}{t} = M \mu \ln \left( \frac{\xi^2}{t_0} \right).$$  \hspace{1cm} (21)

To compute $z$, we also need an estimate for $S(1)$, that in this case is not trivially equal to one. As the population is assumed to be well-mixed in an area equal to $\xi^2$ or smaller, the composition of a single site can be thought as a sample of $M$ individuals from this well-mixed population. The probability distribution of the species abundance in such a sample is given by Ewens' sampling formula [54], which is an important result in population genetics. Substituting its expression yields

$$S(1) = \sum_{j=0}^{M-1} \frac{M\mu}{M\mu + j} \approx M\mu \ln(1 + \mu^{-1}).$$  \hspace{1cm} (22)

Combining Eqs. (21) and (22) and assuming a power law in the range from $A = 1$ to $A = \xi^2$, we find an exponent

$$z \sim \frac{\ln \ln(\xi^2)}{\ln \ln(\xi^2)} = \frac{\ln \ln(\nu/\mu)}{\ln(\nu/\mu)}.$$  \hspace{1cm} (23)

which, to the leading order, is a factor 2 smaller than the corresponding estimate for the voter model [7]. The decrease of the exponent $z$ with the combination of parameters $M \mu$ is confirmed in numerical simulation, see Fig. 8 although the asymptotic reduction is less than the factor two predicted by the above estimate.

![FIG. 8: Species-area exponents for the Stepping Stone Model at fixed $\nu = 10^{-6}$, different local population size $M$ and dispersal rate $\mu$, with NN dispersal. The numerical estimate of the exponent $z$ in the voter model for NN-dispersal and the same value of the speciation rate is shown for comparison.](image)

Summarizing, the stepping-stone model at large local community size $M$ yields smaller values of the species-area exponent $z$ than the voter model [27]. This fact is consistent with the ecological observation that microbial communities, characterized by very large local community sizes, typically display very shallow species-area...
relations, and that in general there seems to be a positive correlation between the exponent $z$ and the body size of a taxonomic group $^{40}$. In the stepping-stone model, a decrease in the SAR exponent is observed in the regime $M \mu \gg 1$ where each site hosts a large number of species and therefore provides a buffer for biodiversity $^{27}$. This interpretation is also consistent with the “cosmopolitan” nature of many microbial species, i.e. the fact that relatively small communities of microbes host a biological diversity comparable with that observed in the whole planet $^{55, 57}$. This feature has sometimes been explained invoking the fact that microbes have the possibility of long-range dispersal $^{57}$. However, numerical simulations show that, in the voter-model with speciation, long-range dispersal leads to steeper, rather than shallower SARs $^{50}$.

### B. Contact Process with speciation

In the voter model, every dead individual is instantly replaced by a newborn, leading to a constantly saturated environment. The implicit underlying assumption is that the birth rate is infinite, so that death events are the rate-limiting steps. Such assumption constitutes a good approximation in resource-rich ecosystems. In less rich ecosystems, where the birth rate is finite, the environment is not always saturated and empty gaps can exist $^{58}$.

To explore this latter case, we study here the contact process with speciation, which is the multi-species variant of the well-known contact process $^{19, 28, 29, 58, 59}$. As usual, we consider the model on a 2D square lattice. Sites of the lattice can be occupied by individuals belonging to different species or empty. The model is defined in continuous time; each individual dies at a rate $d$ and reproduces at a rate $b$. In case of a death, the site is simply left vacant. A reproduction event is considered successful only if the individual has at least one vacant neighboring site. In such a case, one of the vacant neighboring sites is chosen at random. With probability $\nu$, the site is occupied by an individual of a new species (speciation event); with complementary probability, $(1 - \nu)$, the newborn is of the same species of the parent.

As in the traditional contact process $^{32, 58}$, the relevant parameter determining the asymptotic density of occupied sites $\rho$ is the dimensionless birth-to-death ration $\eta = b/d$. For $\eta < \eta_c \approx 1.649$ the absorbing state in which all sites are empty is stable. A non-equilibrium phase transition at $\eta = \eta_c$ separates this region from a stable active phase $(\eta > \eta_c)$ characterized by a non-vanishing value of $\rho$ that depends on $\eta$ $^{28, 29}$. For $\eta \to \infty$ one has $\rho \to 1$ and the model is equivalent to the voter model with speciation $^{12}$.

Unfortunately, dual techniques to efficiently simulate the contact process with speciation have not been yet developed. Forward simulations show that the SAR and the corresponding exponents are remarkably similar to the voter model with speciation even at small values of $\eta$, corresponding to very fragmented ecosystems as shown in Fig. 9. For values of $\eta$ very close to $\eta_c$ (but within the active phase) and small values of $\nu$, SAR exponents tend to be smaller than in the voter model, see inset of Fig. 9.

In principle, in a very fragmented ecosystem it would not make sense to sample empty areas, or areas with very few individuals. With this idea in mind, an alternative to the standard definition of SAR used so far is to weight the sample of a given area with their number of individuals, i.e. of occupied sites. Adopting this definition one finds different SARs for small values of $\eta$ $^{27}$. These SARs do not seem to be characterized by a clear power-law range. We refer the reader to Ref. $^{27}$ for a broader discussion of this issue.

### C. O’Dwyer-Green model

We have seen that finding exact results for neutral spatial models constitutes a formidable problem, and even in the simple case of the voter model only asymptotic results are known.

To make progress in this direction, O’Dwyer and Green...
IV. NEAR-NEUTRAL MODELS

So far, we examined neutral ecological models. However, biodiversity patterns that are well predicted by neutral models can also be predicted by richer, non-neutral models \[62\]. A main difficulty in comparing neutral and non-neutral models is the disproportionately large number of possible ecological effects (and corresponding parameters) that typically enter the latters. In this section, with the aim of understanding basic non-neutral effects in a simple setting, we present a minimal model introduced in \[61\], where one can continuously move from a neutral to a non-neutral scenario by varying a single parameter, tuning the amount of spatial disorder. We then discuss generalizations to other types of spatio-temporal disorder.

A. Habitat-preference model

We consider a variant of the voter model where different sites are preferred habitats for each one of the competing species. For the sake of simplicity, we limit ourselves to the case of two species \(A\) and \(B\) with \(N_A\) and \(N_B\) individuals, respectively. We assume habitat saturation, so that the total population is \(N = N_A + N_B = L^2\) where the system is a square lattice of size \(L\) with periodic boundary conditions. Individuals of type \(A\) and \(B\) can also migrate to the system from an infinite reservoir where they are equally represented. Each lattice site can be of type \(A\) or \(B\), i.e. being a preferred habitat for colonization by species \(A\) or \(B\), respectively. After colonization, mortality and dispersal do not depend on being on a preference site. Ecologically, this means that the fitness advantage belongs to the seeds and not to the individuals themselves (see \[67\] for a different choice). The \(a\) vs. \(b\) character of each site is chosen randomly at the beginning and it remains fixed over time – quenched disorder. To maintain the model globally symmetric, we assume equal proportions of \(a\) and \(b\) sites. Since the intensity of the two biases \((a\) favouring \(A\) and \(b\) favouring \(B\)) are identical. The dynamics proceeds as follows. At each discrete time step a lattice site is randomly chosen with uniform probability and the residing individual is killed. The individual is replaced either by an immigrant from the reservoir (with probability \(\mu\)) or by an offspring of an individual residing in one of the four neighboring sites (with probability \((1 - \mu)\)). In both cases, the colonization probability is biased by an additional factor \(\gamma\) for the individuals that have preference for the empty site. In formulas, the probability of colonization of a site \(x = \{a, b\}\) by an individual \(X = \{A, B\}\) \((Y = \{B, A\})\) having (not having) preference for that site is

\[
W_X(n_X, n_Y) = \begin{cases} 
(1 - \mu) \frac{(1+\gamma)n_X}{(1+\gamma)n_X + n_Y} + \mu \frac{1+\gamma}{2+\gamma} 
\end{cases},
\]

\[
W_Y(n_X, n_Y) = \begin{cases} 
(1 - \mu) \frac{n_Y}{(1+\gamma)n_X + n_Y} + \mu \frac{1+\gamma}{2+\gamma},
\end{cases}
\]

respectively, where \(n_X\) \((n_Y)\) denotes the number of individuals of species \(X = \{A, B\}\) \((Y = \{B, A\})\) in the neighborhood of the considered site. Similar models have been proposed also in the context of heterogeneous catalysis \[68\] and social dynamics \[69\]. For \(\gamma = 0\) and \(\mu = 0\), the standard (neutral) voter model with two species is recovered. For \(\gamma = 0\) but \(\mu \neq 0\), it corresponds to the noisy voter model \[71\] .

Although we assumed NN dispersal, one can generalize also this model to an arbitrary dispersal kernel \(P(r)\). Here, we only consider the NN dispersal and global dispersal (GD), i.e. a mean-field version of \[23\]. The latter can be thought as a variant of the two islands model \[72\] of population genetics, where each island host \(N/2\) individuals and is favorable to one of the two species. In the mean-field version, the state of the system is univocally determined by the numbers of individuals \(N_{Aa}\) and \(N_{Bb}\) residing on their island of preference. The numbers of
individuals outside the island of preference can be obtained as \(N_{Ba} = N/2 - N_{Aa}\) and \(N_{Ab} = N/2 - N_{BB}\). The dynamics is thus encoded in the probabilities per elementary steps that \(N_{Xa}\) (with \(X = \{A, B\}\) and \(x = \{a, b\}\)) increases or decreases by a unit:

\[
W_{N_{Xa} \rightarrow N_{Xa}+1} = \left(\frac{1}{2} \frac{N_{Xa}}{N}\right) W^X_{N_a}(N_A, N_B)
\]

\[
W_{N_{Xa} \rightarrow N_{Xa}-1} = \frac{N_{Xa}}{N} W^X_{N_a}(N_A, N_B)
\]

(25)

where \(W^X_{N_a}\) and \(W^X_{N_b}\) are given by eqs. (24) with \(n_X\) and \(n_Y\) replaced by \(N_X = N_{Xa} + N_{Xb}\) and \(N_Y = N_{Ya} + N_{Yb}\), respectively.

B. Extinction times

In the absence of immigration (\(\mu = 0\)) and for finite populations \(N < \infty\), persistent coexistence of the two species is not possible: demographic stochasticity eventually drives one of the species to extinction (the absorbing state) with the fixation (in the jargon of population genetics) of the other species. However, interesting information on the system can be obtained by studying the dynamics toward extinction [63]. Of particular interest is the average extinction time, \(\langle T_{\text{ext}} \rangle\), and its dependence on system properties, such as the deviation from neutrality and the population size.

In the neutral case (\(\gamma = 0\)), as discussed, the system recovers the voter model with NN dispersal and the Moran model [37] in the version with global dispersal. In this limit, the extinction time is set by the population size. In particular, for large \(N\) we have \(\langle T_{\text{ext}} \rangle \sim N\ln N\) for NN-dispersal [73] and \(\langle T_{\text{ext}} \rangle \sim N\) for global dispersal [37, 73]. To inquire the effect of habitat preferences we simulated many realizations of the model [24] starting \(N_A = N_B = N/2\) until the extinction of one of the two species.

Figure 10 shows the average extinction time, measured in generations (i.e. \(N\) elementary steps of the dynamics described in the previous section), as a function of the population size \(N\) for different values of \(\gamma\) in the model with NN-dispersal. For \(\gamma = 0\) we observe the \(N\ln N\) behavior expected in the neutral case. Habitat preference (\(\gamma > 0\)) leads to a dramatic increase of the average extinction time, which becomes exponential in \(N\)

\[
\langle T_{\text{ext}} \rangle \propto \exp(C(\gamma) N),
\]

(26)

for large enough \(N\). The dependence of the constant \(C(\gamma)\) on \(\gamma\), shown in the inset, is well-fitted by a power-law with exponent \(\approx 1.63\). The mean field version of the model presents similar qualitative features with the only difference that \(\langle T_{\text{ext}} \rangle \propto N\) for \(\gamma = 0\) and with some differences in the \(\gamma\) dependence of \(C(\gamma)\), as shown in [63].

The exponential dependence of the average extinction times on \(N\) indicates that habitat preference, as a non-neutral effect, has a stabilizing impact on the population dynamics. Indeed, when \(N\) is large enough, the two species coexist on any realistic time scale. The stabilizing effect of habitat preference reflects also in the probability of fixation, \(P_{\text{fix}}\), namely in the dependence of the probability that the species, say \(A\), gets fixated when initially present as a fraction \(x = N_A/N\) of the population. In the neutral case, standard computation [74] shows that \(P_{\text{fix}}(x) = x\). As shown in [63], when \(\gamma\) is increased, \(P_{\text{fix}}(x)\) develops a much steeper dependence on \(x\) and quickly reaches values \(\approx 1/2\) even for small \(x\), provided \(\gamma\) is large enough. In other words, the stabilization due to habitat preference tends to compensate any initial disproportion between the population of the two species.

C. Coexistence

In the presence of immigration (\(\mu > 0\)), a temporarily extinct species can recolonize, leading to a steady dynamical coexistence between the two species. However, if the typical recolonization time 1/\(\mu\) is large compared to the average extinction time \(\langle T_{\text{ext}} \rangle\), such recovery from extinction is slow and unlikely. Since there is a symmetry between the two species, this implies that most of the time the population is dominated by one of the two species. Therefore, the distribution of the population size of any of the two species, \(P(X)\) \((X = A, B)\) is peaked at 0 and at the population size \(N\), corresponding to dominance of either of the two species. We denote this state of coexistence as monodominance, see Fig. 11a. In the opposite limit \(\langle T_{\text{ext}} \rangle \gg 1/\mu\), temporary extinctions are very un-
likely and the distribution is peaked at \( N_A = N_B = N/2 \) leading to pure coexistence of the two species (Fig. 11). For intermediate values of \( \mu \), temporary extinctions are still possible though the replenishment due to immigration will tend to equilibrate the two populations. In this case of mixed coexistence, the distribution is characterized by three local maxima at \( N_X = 0, N/2, N \) (Fig. 11). Then, the equilibrium distribution \( P(N_A) \) can be computed imposing the detailed-balance condition

\[
\frac{P(N_A + 1)}{P(N_A)} = \frac{W_{N_A \to N_A + 1}}{W_{N_A \to N_A - 1}}, \quad (28)
\]

which must hold at stationarity since the process is one dimensional [73]. To determine \( \mu_c(N) \) for the transition from monodominance to coexistence, it is enough to determine when \( P(N_A) \) passes from being a decreasing function of \( N_A \) (we are assuming \( N_A < N/2 \) and see Fig. 11h) to an increasing one (Fig. 11i). Using (28) with (27) and imposing \( P(N_A + 1) > P(N_A) \) after some algebra, one obtains the inequality \([(2 + N)\mu - 2](N - 2N_A - 1) > 0 \), which is verified whenever \( \mu > 2/(2 + N) \). Notice that, in the case of global dispersal, the distribution is uniform along this line, i.e. for \( \mu = \mu_c \) one finds \( P(N_A) = 1/N \), which is not the case for NN-dispersal.

**D. Generalizations**

A slightly different version of the habitat preference model introduced above was analyzed in detail in [73], by means of a Kramers-Moyal expansion of the master equation in the global dispersal case. It was shown that, neglecting the effect of fluctuations, a non-vanishing local preference corresponds to an effective potential \( V(\delta \rho) \) for the relative difference of densities \( \delta \rho = (N_A - N_B)/N \). The potential \( V(\delta \rho) \) has a minimum at the coexistence state \( \delta \rho = 0 \). When fluctuations are considered, the only possible steady states are the absorbing states \( \delta \rho = \pm 1 \), where \( V(\delta \rho) \) is singular. The minimum at \( \delta \rho \) is separated from the negative singularities by two potential barriers. As \( \gamma \) and/or \( N \) increase, the basin of attraction of the coexistence state becomes larger and deeper and the two symmetric barrier become closer to the absorbing states and progressively higher. Consequently the time needed to escape the coexistence state becomes much longer, therefore unaccessible in computer simulations. Three different regimes can then be identified: the absorbing, intermediate (quasi-active) and active phase, respectively. In the absorbing phase, symmetry is broken and one of the two species reaches extinction with certainty. This regime is equivalent to the monodominance regime in Fig. 11. The active phase is characterized by a coexistence of both species, and survives fluctuations only in the infinite-size limit. This corresponds to the coexistence phase of Fig. 11. Finally, the intermediate state is a mixture of the two previous ones: the absorbing states and the coexistence state are locally stable, thus, the system is tri-stable, and the steady state depends on the initial conditions. This is the mixed state of Fig. 11. These results provide a nice example of how noise can effectively change the shape of a deterministic potential. Still, the presence of absorbing states – with the associated singularities in the steady state distribution – prevent true phase transitions from occurring: the only possible steady state for any finite system is an absorbing
one. Only in the infinite-size limit, noise vanishes and the coexistence state becomes truly stable [76].

Another study scrutinized the case in which there are local habitat preference only at some lattice sites, while all other sites are neutral [77]. An interesting example which has been analyzed in details is that of a square lattice where only the left (resp. right) boundary has a preference for species A (resp. B) ([77], see also 78, 79). The conclusion is that even mild biases at a small fraction of locations induce robust and durable species coexistence, also in regions arbitrarily far apart from the biased locations. As carefully discussed in [77] this result stems from the long range nature of the underlying critical bulk dynamics of the neutral voter model, and is robust to the introduction of non-symmetrical biases — i.e. stronger for one of the species— except for the fact that the state of coexistence is no longer symmetric. These conclusions have a number of potentially important consequences, for example, in conservation ecology as it suggests that constructing local “sanctuaries” for different competing species can result in global increase of stability of their populations, and thus and enhancement of biodiversity, even in regions arbitrarily distant from the protected zones [77].

E. Temporally-dependent habitat preferences

We have just illustrated in various ways that spatial quenched disorder generically fosters species coexistence. An equally important question is what happens when the preference for a species or its competitor (generalizations to more than two species being straightforward [80]) are time dependent, i.e. what happens if neutrality is transiently broken in favor of one of the coexisting species, while maintaining the time-averaged symmetry among species on the long term? This type of question has a long tradition in ecology. Many theoretical studies have tried to explain the impact of environmental fluctuations on population growth and ecosystem stability (see e.g. 67, 68). On one hand, environmental stochasticity enhances fluctuations and extinction rates, that can have a destabilizing effect on the ecological community. On the other hand, it can also foster stability as a result of the differential response of species to environmental variations.

Similarly to the case of spatial disorder, one can devise quasi-neutral models where habitat-preferences for different species are time-dependent. Recently, different works have systematically analyzed these models and shown that adding temporal-dependent habitat preference greatly improves predictions of empirical ecological patterns with respect to purely neutral theories [81, 82, 83].

V. PERSPECTIVES AND CONCLUSIONS

The range of ecological problems we discussed in this review is by no means exhaustive, and we believe there are many directions that still need to be explored or fully understood.

A prominent example is the role of different speciation mechanisms on spatial biodiversity. In the models discussed in this review, speciation events involve a single individual (point speciation mode, in the language of evolutionary ecology). This assumption is convenient from the modeling perspective, but leads to fitted values of the speciation rate that are incompatible with independent estimates [84]. This assumption also tends to generate too many young species which last for a short time and overweight rare species. To address these issues, recently, another mechanism called protracted speciation has been proposed in the context of neutral models [85]. In protracted speciation, the speciation event does not occur at a single generation, but is a gradual event lasting for more generations. Introducing protracted speciation partially solves some of the aforementioned problems [85]. In real ecosystems, even more speciation mechanisms are at play [86]. For example, in parapatric speciation, two spatially-separated population of the same species can diverge and give rise to two different species. This would correspond to a speciation event involving a group of individual rather than a single one. The role of different speciation modes in maintaining biodiversity and in patterning the spatial organization of species is still under discussion and modeling results can provide very useful contributions to this debate.

As mentioned in the Introduction, ecological neutral theory elicited a heated debate which is far from being solved as, in many cases, non-neutral models based on the concept of niche and neutral models yield similar fits of biodiversity patterns [63, 64, 87]. In recent years a new view on this debate has been emerging. In Chase and Leibold’s words: “niche and neutral models are in reality two ends of a continuum with the truth most likely in the middle” [88]. Indeed, the ecological forces underlying niche and neutral models are not mutually exclusive, and demographic stochasticity plays an important role also in non-neutral settings. However, it has been difficult to clarify the importance of different neutral and non-neutral mechanisms, as most non-neutral model are characterized by a large number of parameters. Some progress in this direction has been obtained in simplified settings which, similarly to the model presented in Sect. IV, allow for a controlled departure from neutrality. For instance, Haegeman and Loreau [89] added the main ingredients of neutral theory, demographic stochasticity and immigration, to a LotkaVolterra competition model. Similar problems have been studied in Refs. 90–92. An interesting future direction would be to study similar models in a spatial context, as well as in an eco-evolutionary context, see e.g. 93.

We have seen throughout this review how some observ-
ables measured by ecologists corresponds to well known quantities in statistical physics: for instance, the $\beta$-diversity is closely related with a two-point correlation function. Other observables, such as SARs and SADs, are less common in statistical physics. A potentially fruitful future direction is to consider other observables which are common in statistical mechanics, such as multi-point correlation functions, and measure them in ecosystems. In this direction, it is very interesting the study of species clustering in $\Psi$, depending on the exponent $\Delta$, see Eq.(10).

In summary, we proposed an overview of different stochastic spatial models in population ecology. We have seen that even very simple models are a source of challenging problems in statistical physics. In particular, because of speciation, each species is bound to extinction and is therefore ultimately transient. This feature is in contrast with traditional classical spin system defined on a lattice where, even when in out-of-equilibrium conditions, the number of spin components is fixed from the beginning. Further, ecosystems are typically two-dimensional and, due to the underlying diffusive behavior, $D = 2$ is the critical dimension for these models. We have shown that this fact often leads to logarithmic correction to scaling laws, which have been difficult to analyze both analytically and numerically. Despite these difficulties, remarkable progress has been made in recent years and we believe that cross-fertilization between statistical physics and ecology will still be crucial in the future to deepen our quantitative understanding of how ecosystems are organized.

Appendix: General scaling relationships

In this brief Appendix, we discuss general condition imposed on the functions $f$ and $g$ by the properties of the function $\Psi$, depending on the exponent $\Delta$, see Eq. [10] and Eq. [11]. Let us write the normalization condition for the function $\Psi$

$$g(A)f(A)\left[\int_{c_1}^{c} dx \ x^{-\Delta} + \int_{c}^{\infty} dx \ x^{-\Delta}\right] = 1.$$  \hspace{1cm} (29)

Notice that only the first term depends on $A$. On one hand, the first integral is singular for $\Delta > 1$ and thus

$$1 \sim g(A)f(A) f(A)^{-\Delta} = g(A)f(A)^{\Delta-1}.$$  \hspace{1cm} (30)

On the other hand, if $\Delta < 1$, the integral is weakly dependent on $f(A)$, so that

$$1 = g(A)f(A).$$  \hspace{1cm} (31)

Similarly, the first moment of $\Psi$ is

$$\langle n \rangle = g(A)f^2(A)f(A)^{\Delta-1} = g(A)f(A)^{\Delta+1}$$  \hspace{1cm} (32)

if $\Delta < 2$ and

$$\langle n \rangle = g(A)f^2(A)$$  \hspace{1cm} (33)

otherwise. Combining the expressions above, different regimes emerge as a function of $\Delta$: if $\Delta < 1$, $f(A) = \langle n \rangle$, while for $1 < \Delta < 2$, $f(A) = \langle n \rangle^{1/(2-\Delta)}$, while no specific prediction for $f(A)$ can be made in the case $\Delta \geq 2$. In particular, for $\Delta < 1$ one has a simple scaling form $f(A) = \langle n \rangle$ and $g(A) = 1/\langle n \rangle$ which applies, for example, to the 1D case as described in the main text. The marginal case $\Delta = 1$ is treated in detail in Sec. [15].

Acknowledgments

MAM is grateful to the Spanish-MINECO for financial support (under grant FIS2013-43201-P; FEDER funds), as well as to J. Hidalgo, S. Suweis, A. Maritan, C. Bolírile for a long term collaboration on issues related to the content of this paper.

[1] M. L. Cody and J. M. Diamond, *Ecology and evolution of communities* (Harvard University Press, 1975).
[2] F. Preston, *Ecology* 41, 611 (1960), ISSN 0012-9658.
[3] M. Rosenzweig, *Species diversity in space and time* (Cambridge Univ Pr, 1995), ISBN 0521499526.
[4] R. MacArthur, *Am. Nat.* 94, 25 (1960).
[5] M. Tokeshi, *Adv. Ecol. Res.* 24, 111 (1993).
[6] S. A. Levin, *Ecology* 73, 1943 (1992).
[7] S. Hubbell, *The unified neutral theory of biodiversity and biogeography* Princeton University Press (Princeton, New Jersey, USA, 2001).
[8] M. Kimura, *The neutral theory of molecular evolution* (Cambridge University Press, 1983).
[9] D. Alonso, R. S. Etienne, and A. J. McKane, *Trends Ecol. Evol.* 21, 451 (2006).
[10] J. Rosindell, S. P. Hubbell, and R. S. Etienne, *Trends Ecol. Evol.* 26, 340 (2011).
[11] S. Azaele, S. Suweis, J. Grilli, I. Volkov, J. R. Banavar, and A. Maritan, *Rev. Mod. Phys.* 88, 035003 (2016).
[12] M. Vallade and B. Houchmandzadeh, *Phys. Rev. E* 68, 061902 (2003).
[13] A. J. McKane, D. Alonso, and R. V. Solé, *Theor. Pop. Biol.* 65, 67 (2004).
[14] I. Volkov, J. Banavar, S. Hubbell, and A. Maritan, *Nature* 424, 1035 (2003).
[15] S. Pigolotti, A. Flammini, M. Marsili, and A. Maritan, *Proc. Nat. Acad. Sci.* 102, 15747 (2011).
[16] R. S. Etienne and D. Alonso, *J. Stat. Phys.* 128, 485 (2007).
[17] J. Rosindell, S. P. Hubbell, F. He, L. J. Harmon, and R. S. Etienne, *Trends Ecol. Evol.* 27, 203 (2012).
[18] R. S. Etienne and J. Rosindell, *PloS one* 6, e14717 (2011).
[19] R. Durrett and S. A. Levin, *Philos. Trans. Royal Soc. London B: Biol. Sci.* 343, 329 (1994).
[93] P. Villa Martin, J. Hidalgo, R. Rubio de Casas, and M. Muñoz, Plos Comp. Biol 12, e1005139 (2016).
[94] J. B. Plotkin, J. Chave, and P. S. Ashton, Am. Nat. 160,