Host-parasite models on graphs

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(Received 26 October 2004; published 25 October 2005)

The behavior of two interacting populations “hosts” and “parasites” is investigated on Cayley trees and scale-free networks. In the former case analytical and numerical arguments elucidate a phase diagram for the susceptible-infected-susceptible model, whose most interesting feature is the absence of a tricritical point as a function of the two independent spreading parameters. For scale-free graphs, the parasite population can be described effectively by its dynamics in a host background. This is shown both by considering the appropriate dynamical equations and by numerical simulations on Barabási-Albert networks with the major implication that in the thermodynamic limit the critical parasite spreading parameter vanishes. Some implications and generalizations are discussed.

DOI: 10.1103/PhysRevE.72.046134 PACS number(s): 89.75.Hc, 87.23.Cc, 02.50.–r, 05.70.Ln

I. INTRODUCTION

Population models, or reaction-diffusion systems, have attracted enormous interest both in the statistical physics community and as abstract versions of real biological dynamics. One particular aspect is the presence of phase transitions and the contact process or directed percolation in various disguises (see below, [1,2]).

Host-parasite or predator-prey systems are a natural extension of single species models. By their classical results Lotka and Volterra were able to explain the nature of abundance oscillations in interacting species [3,4]. In regular landscapes or lattices, with a finite spreading rate of the species, these oscillations appear as traveling waves, which can be regular or chaotic, depending on the interplay of time scales in population dynamics and spreading, though it is not clear if the phenomenon survives in the thermodynamic limit [5–11]. In nature they have been observed in different systems, to name two extreme cases, e.g., in vole populations [12] and for human diseases such as measles [13]. In the case of measles in a population living on a landscape of nontrivial island structure, power law fluctuations are found instead [14].

Much of these ideas have recently been generalized in the context of small-world or in particular “scale-free” graphs [15–18]. For the latter, a perfectly valid example is given by epidemics of viruses in the Internet since it has as a graph a fat-tailed probability distribution of the number of nearest neighbors, \( P(k) \). Recently, various models have been studied as the particulars of the structure—like the so-called degree distribution \( \gamma \) in \( P(k) \sim k^{-\gamma} \)—are varied. A fundamental discovery concerning disease spreading is an absence of epidemic threshold in the limit of infinite graphs and the finite-size effective “critical point” obeys an unusual scaling as \( L \), the graph size, is varied [19–21].

This closely relates to the present work where we study the influence of a network or graph like structure of the underlying landscape on host-parasite or predator-prey dynamics. The main findings are (i) the absence of oscillations, (ii) the absence of an infection threshold in the limit of an infinite scale-free graph, and (iii) the existence of two separate transitions in the case of Bethe lattices with finite coordination number \( z \) (“empty” \( \rightarrow \) “hosts only,” “hosts only” \( \rightarrow \) “hosts plus parasites,” but no transition “empty” \( \rightarrow \) “hosts plus parasites”). The structure of the rest of the paper is as follows. Section II contains the necessary definitions, and the two following ones analytical considerations and, to compare, numerical simulations of the models. Finally, Sec. V finishes the paper with a discussion.

II. MODEL FORMULATION

A. States and rates

A basic model for epidemiological applications is the contact process, or the so-called susceptible-infected-susceptible (SIS) model. Here one considers individuals living on the nodes of an underlying graph which are either infected (\( I \)) or susceptible (\( S \)) to an infection. An infected individual may spread the disease to a susceptible one if both are in contact, i.e., if they live on neighboring nodes of the graph. Infected individuals recover with a certain rate and in this simple version immediately become susceptible for a new infection. So the dynamics of the SIS model is defined by the rates

\[
\begin{align*}
    r_{S \rightarrow I} = \lambda, & \quad \text{if any neighbor is infected,} \\
    r_{I \rightarrow S} = 1. & \quad \text{(1)}
\end{align*}
\]

In this work we generalize the SIS model to a system with hosts and parasites (HP). In other words we consider infections of a second kind only able to spread onto sites with infections of first kind. So each node in the graph can be in three possible states: Empty (\( e \)), or populated by a healthy host (\( h \)) or a host with parasite (\( p \)). Between three states there are six possible transitions so the dynamics are defined by the following rates (Fig. 1):
graphs used in our simulations and calculations.

As in the SIS model defined above the decay of the host or first kind of infection sets the time scale \( r_{h-e}=1 \). In biological systems \( \alpha>1 \) (even \( \gg 1 \)) if the parasite affects the health of the host. A benefit would mean \( \alpha<1 \). We shall consider cases in which the parasite virtually does not die “on its own” but only when the host is killed, i.e., the case \( 0=e<1, \alpha, \mu, \alpha \).

In Sec. III we present approximate analytical solutions following [19] to the the model of Eqs. (3) which are compared to Monte Carlo simulations in Sec. IV. Particular interest lies in parasite extinction and its dependence on the parasite spreading rate \( \alpha \mu \). But first we define the types of graphs used in our simulations and calculations.

B. Graphs

We study the population dynamics of the HP model on two types of graphs, on Bethe lattices and on scale-free Barabási-Albert (BA) graphs, in their standard version [21]. A Bethe lattice of coordination number \( z \) is an infinite tree, where each node has \( z \) neighbors. When constructing a finite lattice, or Cayley tree, starting from a central node with \( z \) neighbors and adding \( z-1 \) new nodes to each boundary node, the number of boundary nodes grows exponentially. It therefore remains a finite fraction of the total number of nodes in the finite tree, which makes this construction unsuitable for Monte Carlo simulations.

This difficulty can be overcome by a slight modification [22] where a sparse homogeneous graph that closely approximates the Bethe lattice without any boundary nodes is constructed. Take \( L \) nodes and label them by integers from 1 to \( L \). Connect node \( i \) to node \( (i+1) \) for each \( i \) and connect node 1 to node \( L \). Construct \( (z-2) \) independent random pairings of the nodes (an easy way to construct pairings is to sort the nodes randomly and pair the first node of this new order with the second one etc.) of the nodes and add an edge for each pair. By this procedure, we get a graph in which each node is of degree \( z \). For large enough graphs, the loops are negligible [22] and this is a sufficient approximation of a Bethe lattice.

Here, we also use the standard version of Barabási-Albert graphs [21]. These are constructed as usual. New nodes are added one by one connecting them with \( m=3 \) links to the previous ones. From these, the neighbors are chosen with a probability proportional to their respective number of links (preferential attachment). By this construction highly linked nodes are likely to obtain even more neighbors as the graph grows, which results in a fat-tail distribution of probabilities for a node to have coordination number \( k \), \( P(k) \sim k^{- \gamma} \) [21]. The BA graphs have very weak degree correlations, i.e., the conditional probability for a node of degree \( k \) to have a neighbor with \( k' \) is rather trivial [15] compared to many other models and real networks.

III. MEAN FIELD AND DOUBLET APPROXIMATION

A. Bethe lattice

1. Singlet (mean field) approach

In this subsection we extend the known solution for the SIS model on a Bethe lattice [19] to the HP model. \( \rho_h \) and \( \rho_p \) denote the density of hosts and parasites, respectively. For simplicity we consider the limit \( e=0 \), so parasitized patches do not supply host individuals to neighboring empty patches. The rate equations for the densities can be written as

\[
\begin{align*}
\dot{\rho}_h &= -\rho_h + \lambda(1-\rho_p-\rho_h)\Theta - \alpha \mu \rho_p \Phi \\
\dot{\rho}_p &= -\alpha \rho_p + \alpha \mu \rho_h \Phi,
\end{align*}
\]

with \( \Phi=1-(1-\rho_p)^z \) and \( \Theta=1-(1-\rho_h)^z \).

In the absence of parasites the host population follows the dynamics of a SIS model. The trivial state \( \rho_h=0 \) is stable for \( \lambda < 1/z \) and unstable otherwise. In other words, the host population can survive only for \( \lambda > 1/z \).

Similarly, the pure host phase is stable if parasites cannot invade, i.e., if the growth rate of a small parasite population is smaller than its death rate,
application to a spatially uniform insect host-parasitoid model
approximation is widely used, we want to emphasize its ap-
proximation which explicitly treats the joint probabili-
ty of finding a host-parasitoid coexistence state can be
seen. The phase diagram is drawn in which a direct transition from the absorbed state to the co-
existence state can be seen. The phase plane is drawn in
Fig. 2.

2. Doublet approach

The singlet approach neglects occupancy correlations be-
tween adjacent nodes. The next logical step is a pair or dou-
blet approximation which explicitly treats the joint probabili-
ties to find two unparasitized hosts next to each other (\(P_{hh}\)), a healthy host next to a parasitized one (\(P_{hp}\)), and two para-
sitized next to each other (\(P_{pp}\)) in addition to \(p_h\) and \(p_p\). This
approximation is widely used; we want to emphasize its ap-
plication to a spatially uniform insect host-parasitoid model [23,24], to the contact process in a one-dimensional chain [2] and in general over a wide class of models [25].

The approximation uses the probabilities \(P_{\sigma\sigma'}\) to find the nodes adjacent to a randomly picked bond in states \(\sigma\) and \(\sigma' \in \{e, h, p\} \), as well as the conditional probabilities \(\rho_{\sigma\sigma'}\) to find a randomly chosen nearest neighbor of a \(\sigma'\) node in state \(\sigma\). Three-point and higher correlations are neglected, so the conditional probabilities to find a \(\sigma\) node next to a \(\sigma'\) node which is itself linked to a third node with state \(\sigma''\) are approximated by

\[
\rho_{\sigma\sigma'\sigma''} = \rho_{\sigma\sigma'} \quad \forall \quad \sigma''.
\]

From there one obtains the rate equations

\[
\begin{align*}
\partial_t p_h &= (-1 - z\alpha \mu \rho_{ph} + z\lambda \rho_{eh})p_h, \\
\partial_t p_p &= (-\alpha + z\alpha \mu \rho_{hp})p_p,
\end{align*}
\]

\[
\partial_t P_{hh} = -[2 + 2(z - 1)\alpha \mu \rho_{ph}]P_{hh} + \lambda[1 + (z - 1)\rho_{he}]P_{he},
\]

\[
\begin{align*}
\partial_t P_{hp} &= -(1 + \alpha + \alpha \mu[1 + (z - 1)\rho_{ph}])P_{hp} + (z - 1)\lambda \rho_{he}P_{pe} + 2(z - 1)\alpha \mu \rho_{ph}P_{hh}, \\
\partial_t P_{pp} &= -2\alpha P_{pp} + \alpha \mu[1 + (z - 1)\rho_{ph}]P_{hp}, \\
\partial_t P_{he} &= -(1 + (z - 1)\alpha \mu \rho_{ph} + \lambda[1 + (z - 1)\rho_{he}])P_{he} + 2(z - 1)\lambda \rho_{he}P_{ee} + 2P_{hh} + \alpha P_{hp}, \\
\partial_t P_{pe} &= -(1 + (z - 1)\rho_{he}P_{ee} + (z - 1)\alpha \mu \rho_{ph}P_{pe} + P_{hp}, \\
\partial_t P_{ee} &= -2(z - 1)\lambda \rho_{he}P_{ee} + P_{he} + \alpha P_{pe}.
\end{align*}
\]

The joint probabilities \(P_{\sigma\sigma'}\) can be expressed in terms of the conditional probabilities as

\[
P_{\sigma\sigma'} = \rho_{\sigma e} \rho_{\sigma' e} (2 - \delta_{\sigma,\sigma'}),
\]

where \(\delta_{\sigma,\sigma'}\) is the Kronecker symbol. The factor 2 for \(\sigma \neq \sigma'\) reflects the two possible choices, because \(\sigma\) can be on either end of the bond.

There are some subtleties in Eqs. (6)–(13) that might not
be immediately obvious. In Eq. (8), for instance, there is a factor of 2 in the first term. That term describes a process
where an edge connecting two host sites turns due to a death of a host into an edge connecting an empty site to a host-
carrying site. The prefactor comes from the fact that this can happen in two ways, i.e., either of the two hosts can die. For similar reasons, a prefactor of 2 can also be found in
the second term of Eq. (8). However, the rest of the terms in the
equation do not have these prefactors since a similar symme-
try does not exist.

In principle Eqs. (6)–(13) are solvable in the steady state. Consider first the SIS model, i.e., the case without any para-
sites. Setting \(\mu=0\) and looking at the steady state of Eq. (6) immediately yields

\[
\rho_{eh} = \frac{1}{z\lambda}.
\]

Similarly, setting \(P_{he}=0\) in Eq. (13) and using the identities

\[
P_{ee} = \rho_{e e} \rho_{e e} \quad \text{and} \quad P_{he} = 2\rho_{e h} \rho_{h e}
\]

gives

\[
\rho_{e e} = \frac{1}{(z - 1)\lambda}.
\]

Expressing \(p_h\) as

\[
p_h = \rho_{e h} \rho_{e h} = (1 - p_h) \rho_{h e} \rho_{e h},
\]

using the identity \(\rho_{e e} + \rho_{h e} = 1\), and plugging in Eqs. (15) and
(16) finally gives

\[
\partial_t p_h = (-1 - z\alpha \mu \rho_{ph} + z\lambda \rho_{eh})p_h,
\]

\[
\begin{align*}
\partial_t P_{hh} &= -[2 + 2(z - 1)\alpha \mu \rho_{ph}]P_{hh} + \lambda[1 + (z - 1)\rho_{he}]P_{he}, \\
\partial_t P_{hp} &= -(1 + \alpha + \alpha \mu[1 + (z - 1)\rho_{ph}])P_{hp} + (z - 1)\lambda \rho_{he}P_{pe} + 2(z - 1)\alpha \mu \rho_{ph}P_{hh}, \\
\partial_t P_{pp} &= -2\alpha P_{pp} + \alpha \mu[1 + (z - 1)\rho_{ph}]P_{hp}, \\
\partial_t P_{he} &= -(1 + (z - 1)\alpha \mu \rho_{ph} + \lambda[1 + (z - 1)\rho_{he}])P_{he} + 2(z - 1)\lambda \rho_{he}P_{ee} + 2P_{hh} + \alpha P_{hp}, \\
\partial_t P_{pe} &= -(1 + (z - 1)\rho_{he}P_{ee} + (z - 1)\alpha \mu \rho_{ph}P_{pe} + P_{hp}, \\
\partial_t P_{ee} &= -2(z - 1)\lambda \rho_{he}P_{ee} + P_{he} + \alpha P_{pe}.
\end{align*}
\]

\[
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\]
\[ \rho_h = \frac{(z-1)\lambda - 1}{(z-1)\lambda - 1/z}. \]  

(18)

from the numerator of which the critical point follows:

\[ \lambda_c^D = \frac{1}{z-1}. \]  

(19)

Note that this is different from the mean field result \( \lambda_c^{MF} = 1/z \). It is also worth noting that rigorous mathematical results of the contact process [26] give bounds on the critical point as

\[ \frac{1}{z} \leq \lambda_c \leq \frac{1}{z-1}. \]  

(20)

Next consider the boundary between the parasite-absorbing and the coexistence phases. Here, hosts live well while parasites are near extinction. Expanding the steady state solution in the limit of small parasite population we derive an equation for the phase boundary. Define two auxiliary quantities \( \Lambda = \rho_{hp} \) and \( B = \rho_{hp} + \rho_{pp} \). Form an equation for \( \partial A \) and set it to vanish since we are looking at the steady state

\[ \frac{\partial A}{\partial t} = \frac{\partial}{\partial t} \rho_{hp} - \frac{1}{2\rho_p} \left( \frac{\partial}{\partial t} \rho_{hp} - \rho_{hp} M_p \right) = 0 \]  

(21)

where the rate equation (7) has been used, and \( M_p \) is the Malthusian parameter or growth rate at low densities of the parasites, i.e.,

\[ M_p = -\alpha + \alpha \mu z \rho_{hp} = -\alpha + \alpha \mu z A. \]  

(22)

Plugging Eq. (9) into Eq. (21), using Eq. (14) and the results of Eqs. (15) and (16) at vanishing parasite population we arrive at

\[ 2(z-1)(1-B)\lambda^2 + 2\alpha z \mu A(1-A) + 2\alpha [B(1-A - 1 - \alpha) + 2\alpha]\lambda - 2(z-1)A\alpha \mu = 0. \]  

(23)

Similarly, starting at the rate equation for \( B \),

\[ \frac{\partial B}{\partial t} = -\frac{\partial \rho_{pp}}{\partial t} \left( \rho_{pp} M_p - \rho_{pp} \frac{\partial M_p}{\partial t} \right), \]  

(24)

using Eqs. (12) and Eq. (14) together with the results of Eqs. (15) and (16), one gets

\[ 2(z-1)(1-B)\lambda^2 + 2\alpha z \mu A(1-B)\lambda + 2\alpha [(B-A)(1-\alpha) - 1] \lambda - 2(z-1)A\alpha \mu = 0 \]  

(25)

given that \( \lambda \neq 0. \)

Now, solve for \( A \) in the steady state version of Eq. (7), substitute this in Eqs. (23) and (25), and eliminate \( B \) from the resulting two equations to get

\[ \mu = \frac{z(z-1)\lambda^2 + z\alpha \lambda}{z(z-1)^2\lambda^2 + z\lambda^2 (\lambda - 1 - \alpha) + \alpha(1-z)}. \]  

(26)

for the phase boundary between parasite-absorbing and coexistence phases in the \( (\lambda, \mu) \) plane. Note that, contrary to the mean field approximation, the phase boundary defined by this equation does not meet that defined by Eq. (19) at \( \mu \rightarrow \infty \) since \( \lambda = \lambda_c^D \) is not a zero of the denominator of Eq. (26). It also holds that \( \mu_c \rightarrow 1/(z-1) \) as \( \lambda \rightarrow \infty \) so that in this limit the parasites always find hosts on all nodes and therefore behave as the SIS model does.

In addition to the solution above we linearize the doublet rate equations around the previously obtained fixed point with a host population and no parasites, i.e., \( \rho_h = P_{pp} = P_{hp} = P_{ep} = 0 \). Replacing the conditional probabilities \( P_{\sigma \sigma'} \) by joint probabilities \( P_{\sigma \sigma'} \) as in Eq. (14) we get a matrix which is of the form

\[ M = \begin{pmatrix} M_h & M_{hp} \\ 0 & M_p \end{pmatrix}, \]  

(27)

where \( M_h \) governs the stability of the “host only” solution, \( M_{hp} \) the effect of a small parasite population on the hosts, and \( M_p \), the growth of parasites at low densities. The block in the lower left corner is zero since the state without parasites is an absorbing one, i.e., a perturbation in the host density cannot reintroduce parasite population into the system.

The eigenvalues of a matrix with the structure of Eq. (27) are just those of \( M_h \) and \( M_p \), irrespective of \( M_{hp} \). The stability of the host population has been discussed above, so we are only interested in the (real parts of the) eigenvalues of the matrix in the following equation,

\[ \frac{d}{dt} \begin{pmatrix} \rho_p \\ P_{hp} \\ P_{pp} \\ P_{ep} \end{pmatrix} = \begin{pmatrix} -\alpha & \alpha \mu & 0 & 0 \\ 0 & B & 0 & \tilde{\lambda} \\ 0 & \alpha \mu & -2\alpha & 0 \\ 0 & C & 2\alpha - \alpha - \tilde{\lambda} \end{pmatrix} \begin{pmatrix} \rho_p \\ P_{hp} \\ P_{pp} \\ P_{ep} \end{pmatrix}, \]  

(28)

with

\[ B = (z-1)\alpha \mu (z\lambda - 1)/(z\lambda) - 1 - \alpha - \alpha \mu, \]  

(29)

\[ C = (z-1)\alpha \mu (z\lambda - 1)/(z\lambda) + 1, \]  

(30)

\[ \tilde{\lambda} = (z-1)\lambda - 1. \]  

(31)

Note that \( \tilde{\lambda} \) is proportional to the excess over the critical host growth rate, \( \lambda - \lambda_c \). The left column of the matrix in Eq. (28) is empty except for the diagonal element, which gives the first eigenvalue \( -\alpha \). We therefore restrict ourselves to the remaining \( 3 \times 3 \) matrix. It is straightforward to calculate its eigenvalues explicitly, from which the phase boundary can be deduced as follows. For each fixed \( \lambda \), we consider the real part of the largest eigenvalue of the \( 3 \times 3 \) matrix as a function of \( \mu \), and find its zero numerically, leading to a point \( \mu(\lambda) \) that lies at the phase boundary. The results are shown in Fig. 2 for the case \( \alpha = 1.2 \) and \( z = 4 \).

The absence of the tricritical point can be seen easily. As \( \lambda \searrow \lambda_c \), the excess growth rate \( \tilde{\lambda} \searrow 0 \), and the matrix becomes lower triangular. All three diagonal elements yield negative
eigenvalues, in particular in this limit $B \to -\alpha \mu/z - \alpha - 1 < 0$. In particular, none of the eigenvalues approaches zero as $\mu \to \infty$, which leads again to the conclusion that the two phase boundaries do not meet at this limit.

In comparison to these results the mean field approximation underestimates the critical values for the spreading parameters. It does not take into account the clustering of populations, i.e., the fact that next to a populated site there is likely another one, which cannot be invaded any more. So the possibility for growth is overestimated.

The phase diagram of the HP model in the $(\mu, \lambda)$ plane obtained from both theoretical approaches and from a stochastic simulation using graph approximation discussed in Sec. II B is drawn in Fig. 2. In the simulations, rough estimates for the phase boundaries were obtained by performing a series of simulations with different $\lambda$ for each fixed $\mu$ and observing when the population died out. The largest value of $\lambda$ at which the population dies out is then defined to be the estimate for the position of the phase boundary. From the figure we see that both analytical solutions are in qualitative agreement with each other and with the numerical results. A property worth noting of the phase diagram is the lack of a “tricritical point” and thus the phase boundary between empty and coexistence phases. Consider also that the singlet approach does reproduce the features of the phase diagram in the Bethe lattice case.

B. Scale-free graph

1. Singlet approach

On graphs with nonconstant degrees the occupancy of a node depends on its coordination number. In general, the higher the degree of a node, the greater is its tendency to be occupied. Following Ref. [19] the rate equations for the occupancies $p^k_\mu$ and $\bar{p}^k_\mu$ on nodes of degree $k$ can be written as

$$\frac{\partial}{\partial t} p^k_\mu(t) = -\bar{p}^k_\mu(t) + \mu k [1 - p^k_\mu(t) - \bar{p}^k_\mu(t)] \Theta(\lambda, \mu),$$

$$- \mu ak p^k_\mu(t) \Phi(\lambda, \mu),$$

$$\frac{\partial}{\partial t} \bar{p}^k_\mu(t) = -\alpha p^k_\mu(t) + \mu ak \bar{p}^k_\mu(t) \Phi(\lambda, \mu),$$

where $\Theta(\lambda, \mu)$ and $\Phi(\lambda, \mu)$ are the probabilities that a given link points to an infected or a parasitized node, respectively. In Eq. (32) the first term on the right-hand side (RHS) corresponds to the death of the hosts, the second one to the host spreading and the third one to parasite spreading, diminishing the number of sites that carry host but no parasite. In Eq. (33) the first term on the RHS describes the death of the parasites while the second one encompasses the spreading. It is known [19] that there is no epidemic threshold if the distribution of node degrees is fat tailed.

The critical behavior of the HP model as obtained from the mean field equations above turns out to be incorrect and is in contradiction to the numerical findings. To see this, consider the rate equations (32) and (33) in the limit of small $\rho$, i.e., by a Taylor expansion in $\rho$. The interaction term $\mu ak p^k_\mu \Phi(\lambda, \mu)$ is quadratic in $\rho$ since $\Phi(\lambda, \mu) \sim \rho$ and drops out from the expansion to first order. This, in turn, means that in this limit the host population behaves as in the SOS model and the parasite population dies out since its equation only has exponentially decaying solutions. Furthermore, this rules out the possibility of a zero epidemic threshold for the parasites, since when the spreading rate approaches zero also the prevalence does so. This leads to the aforementioned contradiction. The corresponding numerical results are presented in Fig. 4 below.

2. Singlet approach with a substantial host population

Next, we use the singlet approach to look at the behavior of the parasites when the host population is well established. The calculation presented here is a straightforward generalization of that in Ref. [27].

The rate equation of the parasites in a Markovian correlated graph in the singlet approach can be written in the limit of small prevalences as

$$\frac{\partial}{\partial t} \bar{p}^k_\mu = -\bar{p}^k_\mu + \alpha \mu p^k_\mu \Phi(\lambda, \mu),$$

where $\Phi(\lambda, \mu)$ is the distribution of parasites when the host population is well established. The calculation presented here is a straightforward generalization of that in Ref. [27].

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where $\Phi(\lambda, \mu)$ is the distribution of parasites when the host population is well established. The calculation presented here is a straightforward generalization of that in Ref. [27].
\[ \Lambda_M \geq \min_k \left( \frac{1}{\psi(k)} \sum_{k'} A_{kk'} \psi(k') \right), \]  

where \( \psi(k) \) is an arbitrary positive vector. Now, set \( \psi(k) = k \rho_h^k \) and \( A = C^2 \) to get

\[ \Rightarrow \Lambda_M \geq \min_k \left( \frac{\sum_{k'} \sum_{l} \rho_h^k P(l|k) \rho_h^l P(k'|l) \rho_h^{k'} \nu_{kl}}{k \rho_h^k} \right) \]

\[ = \min_k \left( \sum_l \lambda \rho_h^l P(l|k) \sum_{k'} k' \rho_h^{k'} P(k'|l) \right). \]

(39)

Above \( \bar{K}_{mn}^{(l)}(l,k) \) denotes the average nearest neighbor degree of such neighbors that carry a host, conditioned that we are looking at a node of degree \( l \). Since the average nearest neighbor degree of all neighbors \( \bar{K}_{mn}^{(l)}(l,k) = \sum_{k'} k' P(k'|l) \) diverges \[29\] as \( k \to \infty \) and \( \rho_h \) necessarily saturates to a constant value \( \bar{\rho}_h^{\text{const}} \leq 1 \) with large \( k \), \( \bar{K}_{mn}^{(l)}(l,k) \) must also diverge at the same limit. Thus the RHS of Eq. (39) diverges, giving \( \Lambda_M \to \infty \) and

\[ \mu_{\text{critical}} \to 0 \]

(40)

at the thermodynamic limit.

### 3. Doublet approach

Next, we formulate rate equations for a graph with a given degree distribution and degree-degree correlations using the doublet approach or pair approximation. The correlations are included in the treatment since their use is natural in the context of pair approximations. The correlated network contains its uncorrelated counterpart as a special case.

The notation is as follows. \( P_{\sigma \sigma'}^{kk'} \) is the probability that a randomly chosen edge that connects nodes with connectivities \( k \) and \( k' \) is \( \sigma (\sigma') \). possible states being \( e, h, \) or \( p \). \( Q_{\sigma \sigma'}^{kk'} \) is the conditional probability that a randomly chosen edge that connects nodes with connectivities \( k \) and \( k' \) is \( \sigma' \) such that the state of the node with connectivity \( k' \) is \( \sigma' \) conditioned that the state of the node with connectivity \( k \) is \( \sigma \). Let \( \Delta_{kk'} \) be as above.

Using the notation above, the rate equations for the SIS model needed for the present treatment can be written as follows:

\[ \partial_t \rho_h^k = -\rho_h^k + \lambda \sum_{k'} k \Delta_{kk'} P_{eh}^{kk'}; \]

(41)

\[ \partial_t P_{hh}^{kk'} = -2 P_{hh}^{kk'} + \lambda P_{hh}^{kk'} + \lambda \sum_{k''} \Delta_{kk'} P_{hh}^{kk''}(k'-1) Q_{eh}^{kk''}, \]

(42)

where in Eq. (42) the first term on the right hand side denotes the process where an infected node gets cured, the second the process where a node of degree \( k \) infects a node of degree \( k' \) and the third the process where a node of degree \( k'' \) infects a node of degree \( k' \), which in turn has another neighbor of degree \( k \) that is infected, turning the edge between the latter two into an edge connecting two infected nodes.

For the HP model, only one rate equation is needed for the present treatment, namely that of the parasite prevalence

\[ \partial_t P_p^k = -\alpha P_p^k + \mu \lambda \sum_{k'} k \Delta_{kk'} P_{hp}^{kk'}. \]

(43)

Now consider the steady state in the SIS model. Multiply Eq. (41) by \( P(k) \) and sum over all \( k \) to get

\[ \rho_h = \lambda P_{eh}. \]

(44)

\( P_{eh} \) is the fraction of all edges in the network that connect an empty node to one with host and \( \rho_h \) is the average host prevalence in the whole network.

The last term on the right hand side of Eq. (42) is positive. Thus in the steady state we can write, leaving out the said term,

\[ P_{hh}^{kk'} = \frac{\lambda}{2} P_{eh}. \]

(45)

Multiplying this by \( k P(k) \Delta_{kk'} \), summing over all \( k \) and \( k' \), and combining with Eq. (44) we get

\[ P_{hh} = \frac{1}{2} \rho_h, \]

(46)

which implies for the relative density of host-host nearest-neighbor pairs that

\[ \frac{P_{hh}}{(\rho_h)^2} \geq \frac{1}{2} \; \Rightarrow \; \frac{1}{2} \rho_h \to \infty \; \text{as} \; \rho_h \to 0. \]

That is, in the limit of small population, the relative density of host-host pairs is enormous. Thus the prevalence correlations in nearest-neighbor nodes are also huge. Since the singlet approach neglects these correlations, this gives reasons to expect that it is not able to capture the properties in the HP model correctly, even though it is known that in SIS model it does [27].

Consider Eq. (43) in the steady state. Multiplying by \( P(k) \) and summing over all \( k \) gives

\[ \rho_p = \mu P_{hp}, \]

(47)

where \( \mu \to 0 \) as \( \rho_p \to 0 \).

Equation (47) tells us that the number of edges through which the parasite population can spread is proportional to the parasite prevalence (instead of the product of parasite and host prevalences). This, in turn, tells that the dynamics of the parasites is similar to the dynamics of the hosts in the SIS model (since in the SIS model the number of edges that can
spread the population is proportional to the population density in the steady state) and serves as an explanation to the zero threshold of the parasites.

IV. MONTE CARLO SIMULATIONS

For a numerical comparison we have simulated the host-parasite-model in Barabási-Albert networks of sizes \( L = 2^{13}, \ldots, 2^{21} \) under the conditions in which \( \rho_h = 0.30 \) and \( \rho_p \ll \rho_h \), i.e., with a stable host population and parasites close to extinction. The simulations are always started with random initial conditions by giving 25% of the nodes the status host and 5% of the nodes the status parasitized independently. Then the simulation is run for a given saturation period of 1000 Monte Carlo (MC) steps during which even the largest system reaches a stationary state. Quantities of interest are then averaged over another 1000 MC steps, where one MC step refers to the simultaneous (parallel) update event of the state variables of the nodes. The used transition probabilities \( p_{\sigma \sigma'} \) from state \( \sigma \) to state \( \sigma' \) in a single time step are \( p_{eh} = 0.012, p_{he} = 0.05, p_{pe} = 0.25 \), and \( p_{hp} \) is varied in the range from 0.02 to 0.2 to produce the variation in the critical point in the thermodynamic limit.

Figure 3 shows how \( \rho_p \) decays as a function of a host’s parasitization probability parameter \( \mu \). Below a size dependent critical value \( \mu_c(L) \) the parasites become extinct resulting in a left-alone host population obeying dynamics defined by the SIS model. For instance when \( L = 2^{13} \) one may estimate that \( \mu_c = 0.26 \). The inset in Fig. 3 strongly suggests that the relationship \( \rho_p \sim \exp-[(\text{const})/\mu] \) is established as in the SIS model [19].

To track \( \mu_c(L) \) more accurately we have studied the extinction probability \( P_{\text{ext}}(\mu_c, L) \) of the parasites during 2000 MC steps from different realizations of BA graphs. The critical point is then determined to be the highest value of \( \mu \) below which the population dies away in a typical realization of a BA graph, and the sizes of the error bars in \( \mu_c \) are estimated from the width of the window in which \( P_{\text{ext}}(\mu_c, L) \) decays from 1 to 0. Figure 4 shows a scaling \( \mu_c(L) \sim 1/\ln(L) \) in the region \( 2^{21} \geq L \geq 2^{16} \), which again compares to the finite size scaling of the critical threshold in the SIS model [30].

Since the probability for a node to become infected depends on its degree we next take a look at the parasite prevalence of nodes of degree \( k \). Figure 5 and the average degree of a site occupied by a parasite \( \langle k \rho_p \rangle \) in Fig. 6. Figure 5 shows that when approaching \( \mu_c \), the relationship \( \rho_p^k \sim k \) begins to hold better and better whereas \( \rho_p^k \) does not change remarkably since the host population is large. In fact, we have noted that, in analogy with the SIS model, the scaling of \( \rho_p^k \) is not just a matter of coincidence but reflects the more general presence of the factor \( \rho_p^k = 1/[1 + (\text{const})/k] \) which is proportional to \( k \) at small \( \mu_c \), or for large values of the constant. Generally, this behavior implies that the largest connected component of hosts serves as a “scalefree” graph for the parasites thus partly explaining the absence of a critical point in the thermodynamic limit.

As \( \mu \rightarrow \mu_c \), survival of the parasite population becomes more and more difficult. Figure 6 shows a consequence of
to the nonregular nature of the scale-free graphs we have not
seen any indications of, e.g., periodic or chaotic oscillations
that arise in many similar models on regular lattices [10,11].
Another possible angle would be to study contact-process-
like models [2], where the spreading rate out of a graph
vertex to a neighbor would depend on the degree of the out-
vertex, for both parasites and hosts. The phase diagram of
such model would be the same for the Bethe case, but for a
scale-free network one would, in analogy with the contact
process itself [31], expect a finite threshold instead of the
vanishing one for the SIS model. We have confirmed this,
analytically, but obviously numerical studies would be of
interest.

The results have implications, less for Bethe lattices
which serve as an analytically tractable special case, but pos-
sibly for dynamical processes on real scale-free graphs. Ex-
amples can be found from ecology (metapopulation dynam-
ics), where similar multispecies scenarios have already been
studied. Parasitoids do play a crucial role for the population
dynamics of the endangered butterfly species Melitaea cinxia
in its fragmented habitat on the Åland islands in the Baltic
Sea, which fit less well to single species models [33,34]. Due
to the distribution of patch sizes and distances between them,
the corresponding network model has a large tail degree dis-
bution [35]. Whether a given patch is populated by hosts
only or also by parasitoids depends on its local connected-
ness. At least qualitatively the observations agree with those
in Fig. 6 and more systematic studies can be envisioned. The
spreading rate depends on distance between and sizes of
patches, in a nontrivial way [32]. If one translates the under-
lying landscape to a network model, the resulting spreading
rates may depend strongly or weakly on the degree of the
emitting node, i.e., lie somewhere in the range between a
generalized contact process and SIS-type models. Thus the
limit considered by [31] may well be relevant in certain eco-
logical systems.

Another field of examples is epidemiology and vaccina-
tion strategies. Knowledge of nontrivial network structures
in disease transmission can be used for vaccination (see, e.g.,
[36]) or outbreak prediction, e.g., [37], and also the impor-
tance of superinfections has been documented (see, e.g., a
seminal work in an evolutionary context [38]). Our ansatz is
an attempt to combine both points of view. From the scale-
free network viewpoint the fundamental idea of concentrat-
ing the effort on nodes with a high \( k \) is valid here as well
[39,40]; consider in particular the “escape” of parasites close
to extinction mentioned above. To fight parasites one needs,
as well, to avoid random immunization. In this context an-
other paradigmatic model is the susceptible-infected-
removed (SIR) model which is a variant of ordinary perco-
lation. By taking in the HP model the right combination of
limits for the parameters (essentially, disallowing recovery to
the empty state from the \( H \) and \( P \) states), one obtains a
variant of the SIR model which resembles in such language
“bootstrap percolation” since the \( R \) (\( P \)) sites are created only
via contact with a neighbor in \( R \). One should thus take note of
possible generalizations of the HP model using similar
recipes as can be applied to the SIR-style ones [41].

In the case of the SIS model, the crossover (or the time-
dependent picture) to the steady state turns out to be inter-

\begin{figure}
\centering
\includegraphics[width=0.5\textwidth]{fig6}
\caption{(Color online) The expectation value of the degree of a
site occupied by the parasites. A scaling form for the average degree
of parasitized nodes, \( \langle k \mid p \rangle \), is found for small \( \mu \). The straight line in
the inset is a guide to the eye.}
\end{figure}
esting, which might be worth looking at here as well [42]. Another practical case related to this might be, say, viruses spreading as attachments to emails on the Internet [43], where again one is confronted with a dynamical graph (of email connections) on top of a larger one (Internet). Finally, we would like to point out that our work could be extended to other similar multispecies models. An example would be a hierarchy of contact processes \( A \rightarrow B, B \rightarrow C, \ldots \) [44,45].

ACKNOWLEDGMENTS

We thank S. Zapperi and J. Lohi for stimulating discussions. This work was supported by the Academy of Finland through the Centre of Excellence program (M.A., M.P., V.V.) and Deutsche Forschungsgemeinschaft via SFB 611 (M.R.). M.R. thanks Helsinki University of Technology for kind hospitality.