

Stability of vacuum in coupled directed percolation processes

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We study the absorbing phase transitions in coupled directed percolation (DP) processes with N -species particles in one dimension. The interspecies coupling is linear, bidirectional, and excitatory. We find that the presence of a spontaneous annihilation process $A \rightarrow 0$ is essential in stabilizing the absorbing phase (vacuum). In the coupled contact processes, the vacuum is stable and the system exhibits DP type transitions, regardless of the coupling strength, for all N . However, in the coupled branching annihilation random walks with one offspring (BAW), where particle annihilations occur only through binary diffusion processes $A+A \rightarrow 0$, the vacuum becomes unstable with respect to an arbitrarily small branching rate in a sufficiently strong coupling regime for $N \geq 3$. The $N=2$ BAW exhibits the DP type transition for any coupling strength, but the inclusion of interspecies hard core (HC) interaction makes the vacuum unstable again and the system is always active in a strong coupling regime. Critical behavior near the zero branching point is characterized by the mean-field scaling exponents, $\beta = \nu_{\perp} = 1/2$ and $\nu_{\parallel} = 1$, regardless of the presence of HC interaction. We also discuss the effects of the asymmetric coupling.

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INTRODUCTION

Nonequilibrium systems with absorbing (trapped) states have been extensively studied in recent years because of their wide applicability to various phenomena in natural science as well as in social and economical science [1,2]. The absorbing transition between an active phase into an absorbing phase can be regarded as one of the simplest and natural extensions of the well-established equilibrium phase transition to nonequilibrium systems.

Critical behaviors near an absorbing transition are categorized into a few universality classes characterized by the symmetry between absorbing states and/or the conservation in dynamics [3–5]. The most prominent universality class is the directed percolation (DP) class [6], which involves typically a single absorbing state without any conservation in dynamics. Most of absorbing-type nonequilibrium models belong to the DP class. Only a small number of models form different classes, such as the directed Ising (DI) class involving two equivalent classes of absorbing states [3,4,7] and the parity-conserving (PC) class with mod(2) conservation of the total particle number [8]. These two classes coincide in one dimension by identifying a domain wall in the DI-type models as a particle in the PC-type models. In higher dimensions, both types of models are always active (no absorbing phase) except at a trivial point (annihilation fixed point) and their critical behavior is described by the mean field theory. None of the models studied so far with higher symmetries than the Ising symmetry (for example, Potts symmetry) stabilizes an absorbing phase even in one dimension.

Recently, a variety of coupled systems have been investigated extensively. Janssen [9] studied coupled DP processes with bilinear and bidirectional interspecies couplings in the

framework of bosonic field theory, where no other critical phenomena were found than the DP. Täuber *et al.* [10] studied linearly and unidirectionally coupled DP processes, where a series of new multicritical phenomena was observed. Coupled PC (DI) processes have been also studied [11], where the absorbing phase become unstable with respect to an arbitrarily small branching rate even in one dimension. More interestingly, the critical behavior near the annihilation fixed point depends on details of particle dynamics, such as the presence of an interspecies hard core (HC) interaction and the branching method [12,13].

Stochastic models for linearly and bidirectionally coupled DP processes have been also studied through two-species branching annihilating random walks with one offspring (BAW₁) in one dimension [14,15]. It was found that the HC interaction is crucial in a strong coupling regime, where the absorbing phase (vacuum) becomes unstable and the system is always active except at the annihilation fixed point of zero branching rate. Critical behavior near the annihilation fixed point was conjectured via an analytic argument and confirmed by numerical simulations [14].

In this paper we studied the N -species BAW₁ models and also the N -species contact processes (CP) with and without interspecies HC interaction. The single-species BAW₁ and CP both belong to the DP universality class, but perturbative renormalization group (RG) calculations and numerical simulations suggest that the dimensional threshold for the vacuum stability is different for these two models [8,11]. For the BAW₁, the vacuum becomes unstable for $d > d_{\text{th}} = 2$, while $d_{\text{th}} = \infty$ for the CP. The essential difference is the absence of spontaneous annihilation process $A \rightarrow \emptyset$ in the BAW₁, where particle annihilations occur only in pairs through binary diffusion processes $A+A \rightarrow \emptyset$. The $A \rightarrow \emptyset$ process may be effectively generated by combinations of branching and hopping processes. However, the generated processes are too weak to stabilize the vacuum for $d > 2$ [16].

In the coupled systems, one can expect a similar scenario, where the stability of the vacuum also depends crucially on the presence of the spontaneous annihilation process. Fur-

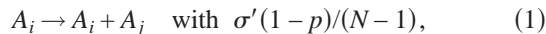
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thermore, as N increases, the vacuum becomes more unstable in the N -BAW₁, because the annihilation process of only the same species of particles is allowed. A sufficiently strong interspecies coupling may wash away completely the absorbing phase for large enough N even in one dimension, even without any HC interaction. In this paper, we address this vacuum stability question in the strong coupling regime and measure the threshold value of N with and without HC interaction.

The outline of this paper is as follows. In Sec. II the N -species CP models are introduced. We find that, for any N , the vacuum is stable for a low branching rate and the DP-type phase transition is observed regardless of the presence of HC interaction. In Sec. III we study the N -species BAW₁ models by numerical simulations. We find that for $N \geq 3$, the vacuum becomes unstable with respect to an arbitrarily small branching rate in a sufficiently strong coupling regime. The critical behavior near the zero branching point is explored in terms of the scaling exponents. In a weak coupling regime, the conventional DP transition into a stable vacuum is observed. The interspecies HC interaction only shifts the DP critical points and the threshold value of the interspecies coupling for the complete disappearance of a stable vacuum. The $N=2$ case is special. Without HC interaction, one can always find the DP transition into a stable vacuum even at the full coupling strength, such as in the N -species CP models. However, the inclusion of the HC interaction makes the vacuum unstable again in a strong coupling regime. We also discuss the effect of the directional asymmetry in the bidirectional couplings on the phase diagram. Finally, we conclude in Sec. IV.

II. COUPLED CONTACT PROCESSES

The N -species coupled contact process (N -CP) is defined by the following evolution rules: (1) Each particle annihilates spontaneously with probability p or (2) creates a particle of the same species with probability $\sigma(1-p)$ or a particle of the different species with probability $\sigma'(1-p)/(N-1)$ in its neighborhood (branching process) as



where $\sigma' = 1 - \sigma$, $i \neq j$, and $i = 1, \dots, N$. Any branching attempt is rejected if it would result in a multiple occupation by the same species particles at a site. With interspecies HC interaction, a multiple occupation by different species particles is also forbidden. Each species is symmetrically coupled with other species by the interspecies coupling strength σ' . For $\sigma' = 0$, all species are completely decoupled.

It is trivial to show that the N -CP models with HC interaction become identical to the single-species CP model. Each site can be occupied by only one particle, regardless of its species. Any configuration and any dynamic process in the N -CP can be exactly mapped on its corresponding configu-

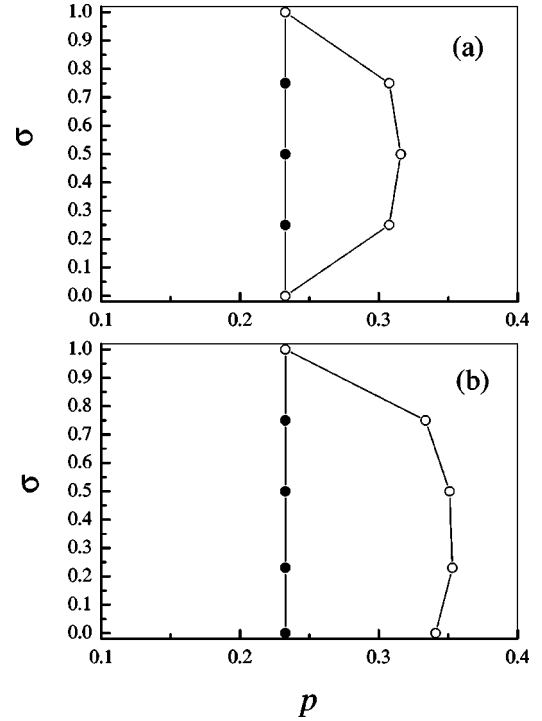


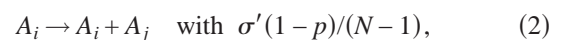
FIG. 1. The σ - p phase diagram for the N -CP, (a) $N=2$ and (b) $N=3$. Filled and open circles correspond to critical points with and without HC interactions, respectively. Lines between data points are guides to the eyes only.

ration and dynamic process of the single-species CP by simply ignoring the particle species.

Without HC interaction, a multiple occupation by different species particles is allowed. One can easily expect that the active phase expands until the multiple occupation is maximized. For $N=2$, there exists a simple duality between two species ($\sigma \leftrightarrow \sigma'$) and the maximum point is located at $\sigma = \sigma' = 1/2$, see Fig. 1(a). For $N \geq 3$, the maximum point is expected to be at $\sigma = 1/N$, where the branching symmetry is perfect and the maximum mixing is expected, see Fig. 1(b). From this reasoning, we expect that complete disappearance of the vacuum does not occur at any coupling strength σ for any finite N . Of course, all absorbing critical phenomena should belong to the DP class.

III. COUPLED BAW₁

The N -species coupled branching annihilating random walks with one offspring (N -BAW₁), is defined by the following evolution rules: (1) Each particle hops to a nearest-neighbor site with probability p or (2) creates a particle of the same species with probability $\sigma(1-p)$ or a particle of the different species with probability $\sigma'(1-p)/(N-1)$ in its neighborhood (branching process) as



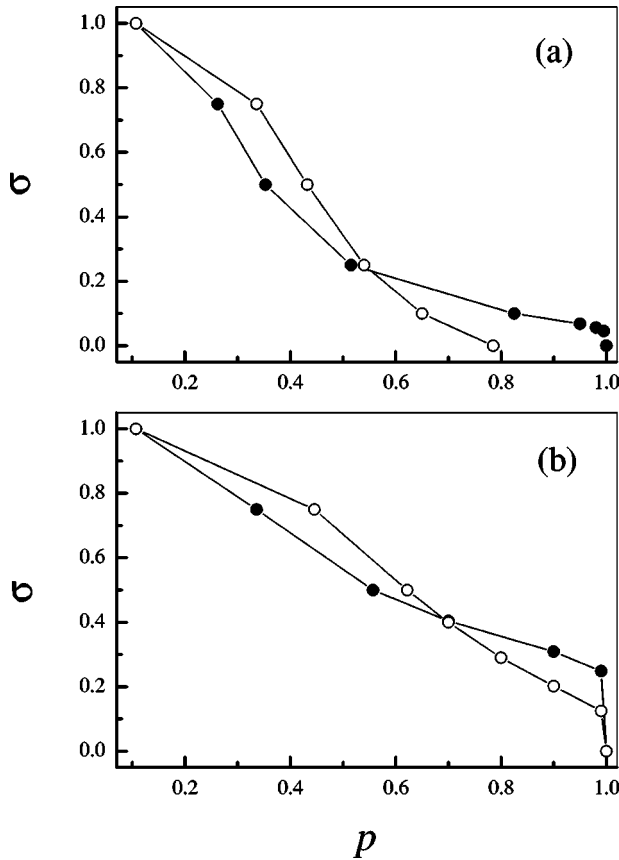


FIG. 2. The σ - p phase diagram of the N -BAW₁, (a) $N=2$ and (b) $N=3$. Filled and open circles correspond to critical points with and without HC interactions, respectively. Lines between data points are guides to the eyes only.

where $\sigma' = 1 - \sigma$ and $i \neq j$. If two identical particles happen to be on the same site, both particles immediately annihilate each other ($A_i + A_i \rightarrow \emptyset$). With interspecies HC interaction, any hopping or branching attempt is rejected if it would result in a multiple occupation of particles, regardless of their species. The above model has a permutational symmetry between species. Later, we will also study an asymmetrically coupled system which breaks the permutational symmetry.

A. $N=2$

The two-species BAW₁ model has been investigated previously for general σ [14]. In this paper we present the numerically improved phase diagram in Fig. 2(a) and briefly summarize the results of [14] for comparison to those for $N \geq 3$ and also for self-containment. Without HC interaction, the system always exhibits the DP-type absorbing transition for all σ , including the case of the maximum coupling strength ($\sigma' = 1$ or $\sigma = 0$). However, with HC interaction, there exists a strong coupling regime ($\sigma' > \sigma'^*$) where the vacuum becomes unstable with respect to an arbitrarily small branching rate ($p = 1^-$) and completely disappears in the phase diagram except at the annihilation fixed point ($p = 1$). The threshold value for the strong coupling regime is numerically estimated as $\sigma'^* \approx 0.95$. In the weak coupling re-

gime ($\sigma' < \sigma'^*$), we observe the conventional DP transition into the vacuum.

It is not surprising to see that the active phase expands as σ' increases, because a collision chance of the same species particles decreases. However, the effect of HC interaction is rather tricky. For small σ' , the system tends to form large domains of the same species particles. The HC interaction induces an effective diffusion bias directed to the domain center, which accelerates the pair annihilation process. Therefore the system becomes less active with HC interaction. For large σ' , the situation is reversed. The system prefers locally heterogeneous configurations and the HC interaction reduces a chance of binary collision of the same species particles. Thus, in this case, the system becomes more active with HC interaction. This explains why two critical lines with and without HC interaction should cross each other as in Fig. 2.

Consider the $\sigma = 0$ line, where a particle cannot create a particle of the same species. As a result, a single particle cannot be annihilated by a single branching and diffusion process: $A \rightarrow A + A \rightarrow \emptyset$. It needs at least three branching processes such as $A \rightarrow AB \rightarrow ABA \rightarrow ABAB$ or $ABBA$. One of the four particle states ($ABAB$) can turn into vacuum via successive pair annihilations only if diffusion across a different species particle is allowed (no HC interaction). Hence, the vacuum can be stable in a low branching (high diffusion) regime without HC interaction. However, in the presence of HC interaction, the ordered AB pairs ($ABAB$) cannot be annihilated by diffusions. Therefore, a single particle has a non-zero probability to survive asymptotically and the vacuum is unstable with respect to an arbitrarily small branching rate.

The critical behavior near the annihilation fixed point in the strong coupling regime is characterized by a set of the mean-field scaling exponents

$$\beta = 1/2, \quad \nu_{\perp} = 1/2, \quad \nu_{\parallel} = 1, \quad (3)$$

where the exponents β , ν_{\perp} , and ν_{\parallel} characterize the scaling behavior of the steady-state particle density ρ_s , the correlation length ξ , and the relaxation time τ , respectively.

The exponent ratios, $\beta/\nu_{\perp} = 1$ and $z = \nu_{\parallel}/\nu_{\perp} = 2$, originate from the ordinary diffusion nature at the annihilation fixed point in one dimension [12]. There is only one independent exponent β , of which the value can be extracted by a simple argument [14]. Consider a particle A created by a particle B . This branching process increases the particle density with the time scale $\tau_b \sim (1-p)^{-1}$. Near the annihilation fixed point of the zero branching rate, the offspring A would be annihilated by colliding with an independent A via diffusion. The time scale for this process is governed by ordinary diffusion: $\tau_d \sim \ell^2$ where the mean distance between particles ℓ is order of the inverse of the particle density ρ^{-1} . Balancing these two time scales, we can expect the steady-state particle density to scale as $\rho_s \sim (1-p)^{\beta}$ with $\beta = 1/2$.

This argument is quite general, so it should apply to many other models exhibiting a critical behavior near the annihilation fixed point (zero branching point). Moreover, the HC interaction does not matter in this argument. So, one can expect that the N -species BAW₁ should belong to the same

class ($\beta=1/2$) for any $N \geq 2$, regardless of the presence of HC interaction, which will be confirmed numerically in next subsection. Moreover, the N -species BAW₂ also belongs to the same class when the HC interaction is present and the branching method is *static*: Two offspring are divided by their parent, $\emptyset B \emptyset \rightarrow ABA$. It is clear that our argument applies to this model, due to the HC interaction. When the branching method is *dynamic* or there is no HC interaction, N -BAW₂ belongs to different universality classes [12].

B. $N \geq 3$

First, consider the $N=3$ case without HC interaction. To map out the phase diagram, we perform the defect-type dynamic Monte Carlo simulations, starting with a single particle. We measure the survival probability $P(t)$, the number of particles $N(t)$, the mean distance of spreading $R(t)$. At criticality, these quantities scale algebraically in the long time limit as $P(t) \sim t^{-\delta}$, $N(t) \sim t^\eta$ and $R(t) \sim t^{1/z}$ [17].

By inspecting the curvature of effective exponents defined as

$$-\delta(t) = \ln[P(t)/P(t/m)]/\ln m, \quad (4)$$

with arbitrary m , and similarly for $\eta(t)$ and $1/z(t)$, we estimate the values of the critical hopping probability p_c and the dynamic exponents δ , η , and z for various values of σ . The σ - p phase diagram is shown in Fig. 2(b). For small σ , the absorbing phase (vacuum) completely disappears. This result is rather unexpected because, for $N=2$, the vacuum is always stable for a low branching rate without HC interaction. Before going into detailed discussion on this vacuum instability without HC interaction, we present numerical results. The threshold value of the vacuum instability is estimated as $\sigma^* \approx 0.125$ ($\sigma'^* \approx 0.875$). For $\sigma > \sigma^*$, we observe that the system undergoes an absorbing transition into vacuum. As expected, we find the DP critical exponents along both paths of constant p and constant σ lines.

To identify the scaling behavior near the annihilation fixed point ($p=1$) for $\sigma < \sigma^*$, we analyze the finite-size effects on the steady-state particle density ρ_s . Using the finite-size scaling theory on ρ_s [18]

$$\rho_s(\Delta, L) = L^{-\beta/\nu_\perp} F(\Delta L^{1/\nu_\perp}) \quad (5)$$

with $\Delta = p_c - p$ and system size L , the value of ν_\perp is determined by collapsing the data of ρ_s with $\beta/\nu_\perp = 1$. We measure ρ_s in the steady state, averaged over $5 \times 10^3 \sim 5 \times 10^4$ samples for several values of Δ ($5 \times 10^{-4} \sim 0.05$) and L ($2^5 \sim 2^9$).

We estimate $\nu_\perp = 0.50(4)$ at $\sigma = 0$. Figure 3(a) shows very good data collapse of ρ_s with $\nu_\perp = 0.50$. These numerical results confirm our previous argument for the universality class near the annihilation fixed point.

To understand the vacuum instability for small σ for $N=3$ without HC interaction, we again consider the $\sigma=0$ line, where a particle can create a particle of different species only. As explained before, there needs to be a sequence of branching processes for a single particle annihilation. For $N=2$ without HC interaction, the process of $A \rightarrow AB \rightarrow ABA$

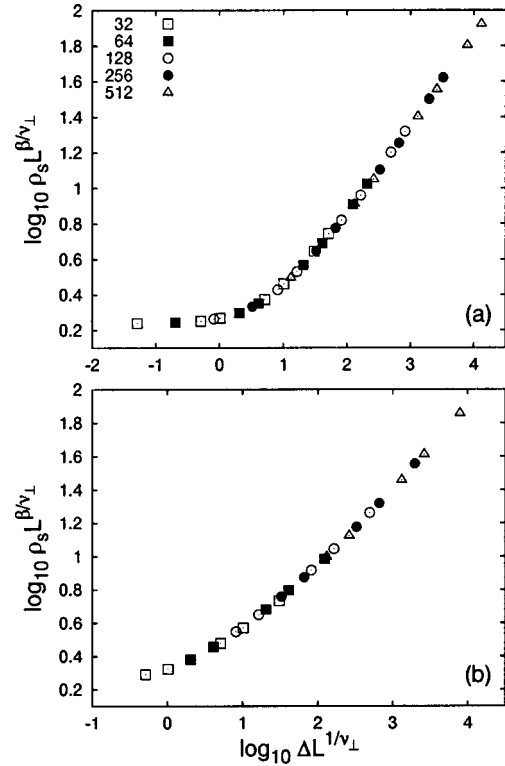


FIG. 3. Data collapse of $\rho_s L^{\beta/\nu_\perp}$ against $\Delta L^{1/\nu_\perp}$ with $\beta/\nu_\perp = 1$ and $\nu_\perp = 1/2$ for system size $L = 2^5, \dots, 2^9$ for $N=3$ (a) without HC interaction at $\sigma=0$ (b) with HC interaction at $\sigma=0.2$.

$\rightarrow ABAB \rightarrow AA \rightarrow \emptyset$ is possible. In this case, AA or BB pairs will quickly go away at a very high diffusion rate, so one may consider only AB pairs, which cannot be annihilated by themselves. The last two reaction events can be regarded as a collision and annihilation event of two AB pairs and the previous reaction events as a creation event of a AB pair out of a parent AB pair. This process is exactly equivalent to the $N=1$ BAW₁ model by identifying an AB pair as a particle. Therefore the $N=2$ case without HC interaction can have a stable vacuum like in the $N=1$ model. The $N=3$ case is different. Three different pairs (AB , BC , CA) are possible. Any pair can branch any other pair, e.g., $AB \rightarrow ACBA \rightarrow CB \rightarrow CBBA \rightarrow CA$. So, the $N=3$ case cannot reduce to the $N=1$ model, in contrast to the $N=2$ case. It seems that this multispecies character is the crucial element for vacuum instability in BAW-type binary diffusion-annihilation models.

Interspecies HC interaction should destabilize the vacuum more easily. We expect that the threshold of vacuum instability should be lowered with HC interaction. In Fig. 2(b), the phase diagram for $N=3$ with HC interaction is presented. We estimate $\sigma^* \approx 0.25$. To estimate ν_\perp for $\sigma < \sigma^*$, we also try the data collapse of ρ_s . In Fig. 3(b), we present the data at $\sigma=0.2$ with the exponent values of $\nu_\perp = 0.50$ and $\beta/\nu_\perp = 1$. Our estimation is $\nu_\perp = 0.50(2)$. Again, we find that our simple argument also applies to this case.

We also perform defect-type Monte Carlo simulations for the $N=4$ case. We estimate $\sigma^* = 0.22(2)$ without HC interaction and $\sigma^* = 0.328(2)$ with HC interaction. The exponent value is estimated as $\nu_\perp \approx 0.52(2)$ at $\sigma=0.15$ without HC interaction and $\nu_\perp \approx 0.52(2)$ at $\sigma=0.3$ with HC interaction.

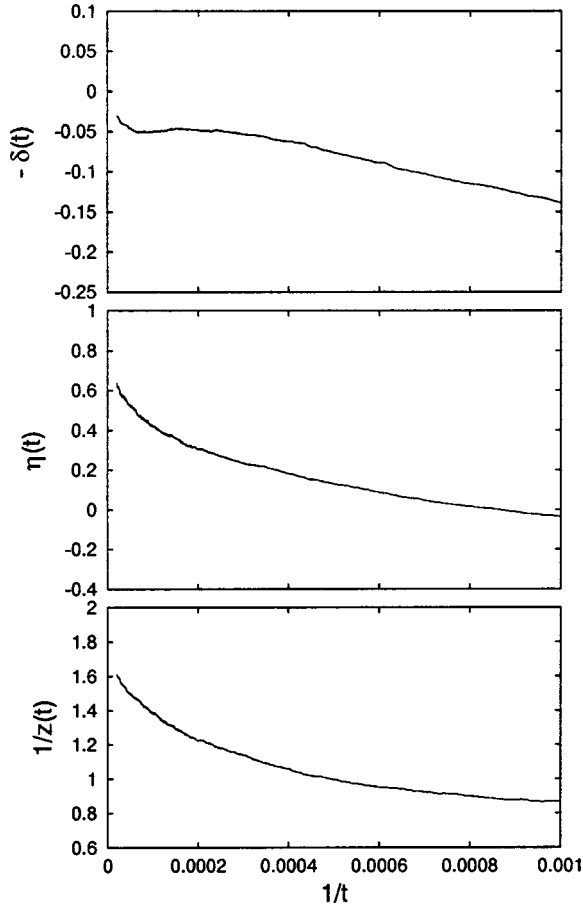
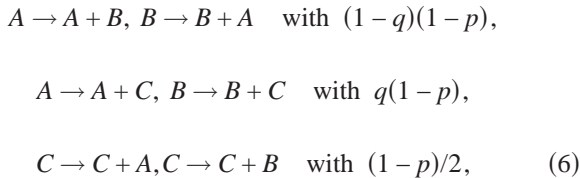


FIG. 4. Plots of the effective exponents against $1/t$ at $p=0.999$ for $q=0.1$. Upward curvature of each exponent indicates that the system is still in the active phase at $p=0.999$.

C. Asymmetric coupling

We study the effect of broken permutational symmetry on the phase diagram. For convenience, we consider the $N=3$ case at $\sigma=0$ ($\sigma'=1$) without HC interaction only. The branching process in Eq. (2) is modified as



where $0 \leq q \leq 1$.

At $q=0$, the species C is completely suppressed and the model becomes identical to the $N=2$ symmetric one at $\sigma=0$. At $q=1/2$, all three species are equivalent and the $N=3$ symmetric model is recovered. Our results on the symmetric N -BAW₁ models in previous subsections indicate that the vacuum is stable in high diffusion regime at $q=0$, but becomes unstable completely at $q=1/2$. For $q < 1/2$, the species C is suppressed in comparison to the other two species A and B . Therefore, the density of the third species C should be proportional to q . Here, we try to locate the threshold value of q for the complete vacuum instability.

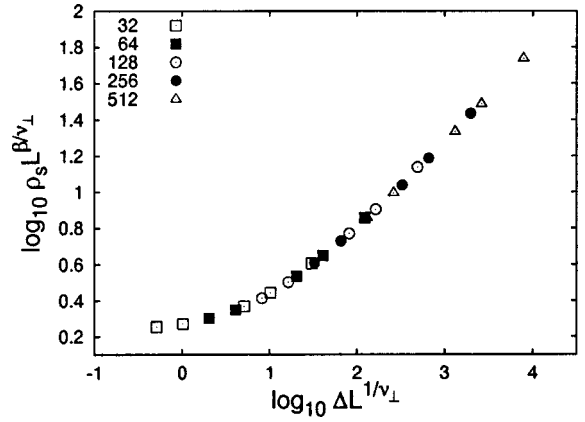


FIG. 5. Data collapse of $\rho_s L^{\beta/\nu_\perp}$ against $\Delta L^{1/\nu_\perp}$ with $\beta/\nu_\perp=1$ and $\nu_\perp=1/2$ for $q=0.1$. The steady state density ρ_s is measured for various system size L ($2^5 \sim 2^9$) and Δ ($5 \times 10^{-4} \sim 3 \times 10^{-2}$).

We perform the defect-type dynamic simulations for several q values ($10^{-1} \sim 1.0$) to locate p_c . In Fig. 4 we present simulation results for $q=0.1$ at $p=0.999$. Effective exponents $\delta(t)$, $\eta(t)$ and $1/z(t)$ show upward curvatures, which imply that the system is still active even at $p=0.999$. It suggests that the criticality is located at $p=1.000(1)$ and the vacuum is completely unstable. For other nonzero q values, we also find the similar results to those for $q=0.1$. We conclude that, for any $q \neq 0$, the system is always active and only critical at $p_c=1.0$. As discussed in the previous subsection, this result again confirms that the multispecies character is relevant (not the symmetry) to vacuum instability. We also check the C -dominant regime for $q > 1/2$ and find a similar result.

To identify the critical behavior, we estimate the exponent ν_\perp by collapsing the particle density data with $\beta/\nu_\perp=1$. Figure 5 shows the scaling plot for $q=0.1$ with $\nu_\perp=0.5$. We estimate $\nu_\perp=0.50(5)$ for $q=0.1$. For other nonzero q values, we also estimate $\nu_\perp \approx 0.5$. Although we do not consider the incomplete coupling case ($\sigma \neq 0$), we expect a nonzero threshold value of σ^* , below which the system is always active.

IV. CONCLUSION

In this paper we study the stability of vacuum in N -coupled DP systems. The interspecies coupling is linear, bidirectional, and excitatory. In the coupled contact processes, the vacuum is always stable at a sufficiently low branching rate for all N , regardless of the coupling strength σ' , and the system undergoes a DP type absorbing transition into the vacuum. On the other hand, in the coupled BAW with one offspring, the vacuum stability is quite fragile for $N \geq 2$ in a strong coupling regime. The absence of a spontaneous annihilation process $A \rightarrow \emptyset$ is crucial for vacuum instability.

We find that the vacuum is unstable with respect to an arbitrarily small branching rate in a sufficiently strong coupling regime ($\sigma' > \sigma'^*$) for $N \geq 3$. The multispecies character is the key element responsible for this vacuum instability and the asymmetry in the interspecies coupling is shown to be

irrelevant. The $N=2$ case is special. The vacuum is always stable as in the coupled contact processes, but the HC interaction is relevant to vacuum instability in a strong coupling regime. We show that the $N=2$ BAW₁ model without HC interaction can reduce to the $N=1$ model even at the full coupling strength, which explains the speciality at $N=2$.

Critical behavior near the annihilation fixed point in a strong coupling regime can be conjectured by a simple argument of balancing two time scales of branching and annihilating random walks. Numerical investigations confirm our conjecture of $\beta=1/2$, $\nu_{\perp}=1/2$, and $\nu_{\parallel}=1$, which also applies to N -BAW₂ with static branching and HC interaction [12].

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