Critical phenomena of nonequilibrium dynamical systems with two absorbing states

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We study nonequilibrium dynamical models with two absorbing states: interacting monomer-dimer models, probabilistic cellular automata models, nonequilibrium kinetic Ising models. These models exhibit a continuous phase transition from an active phase into an absorbing phase which belongs to the universality class of the models with the parity conservation. However, when we break the symmetry between the absorbing states by introducing a symmetry-breaking field, Monte Carlo simulations show that the system goes back to the conventional directed percolation universality class. In terms of domain wall language, the parity conservation is not affected by the presence of the symmetry-breaking field. So the symmetry between the absorbing states rather than the conservation laws plays an essential role in determining the universality class. We also perform Monte Carlo simulations for the various interface dynamics between different absorbing states, which yield new universal dynamic exponents. With the symmetry-breaking field, the interface moves, on average, with a constant velocity in the direction of the unpreferred absorbing state and the dynamic scaling exponents apparently assume trivial values. However, we find that the hyperscaling relation for the directed percolation universality class is restored if one focuses on the dynamics of the interface on the side of the preferred absorbing state only.

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I. INTRODUCTION

Many nonequilibrium dynamical models show continuous phase transitions similar to ordinary equilibrium models. In fact, nonequilibrium models can supply much richer critical behavior because their evolving dynamics do not require the detailed balance. So the universality classes of nonequilibrium critical phenomena would be much more diverse and would be governed by various symmetry properties of the evolution dynamics.

An interesting example of nonequilibrium phase transitions is the absorbing phase transition. In this case, there exist some absorbing states in the configurational phase space. If the system gets into one of the absorbing states by the evolution dynamics, then the system is trapped inside of the absorbing states and no further dynamics occur to escape out of the absorbing states. By controlling an external parameter, one can observe a continuous phase transition from an active steady-state phase into an inactive absorbing phase. Recently, various kinds of nonequilibrium models exhibiting such an absorbing phase transition have been studied extensively [1]. Most of the models investigated are found to belong to the directed percolation (DP) universality class [2] [12]. A common feature of these models is that the absorbing phase consists of a single absorbing state.

Only a few models have been studied that are not in the DP universality class. Those are the models A and B of probabilistic cellular automata (PCA) [13] [14], nonequilibrium kinetic Ising models with two different dynamics (NKI) [15] [16], and interacting monomer-dimer models (IMD) [18] [19]. Numerical investigations show that critical behaviors of these models are different from DP but form a non-DP universality class. These models share a common property that the absorbing phase consists of two equivalent absorbing states. By the analogy to the equilibrium Ising model which has two equivalent ground states, we call this non-DP universality class as the directed Ising (DI) universality class.

Recently, the branching annihilating random walks (BAW) with offsprings have been studied intensively [20] [21] [22]. Even though the BAW model has a single absorbing state (vacuum), its critical behavior depends on the parity of the number of offsprings. It has been shown numerically that the BAW models with an odd number of offsprings (BAWo) belong to the DP class, while the BAW models with an even number of offsprings (BAWe) belong to the...
II. DYNAMIC CRITICAL BEHAVIOR OF THE IMD MODEL

The interacting monomer-dimer (IMD) model is a generalization of the simple monomer-dimer model on a catalytic surface, in which particles of the same species have nearest-neighbor repulsive interactions [18]. Here we consider the one-dimensional IMD model with infinitely strong repulsions between the same species. A monomer ($A$) cannot adsorb at a nearest-neighbor site of an already-occupied monomer (restricted vacancy) but adsorbs at a free vacant site with no adjacent monomer-occupied sites. Similarly, a dimer ($B_2$) cannot adsorb at a pair of restricted vacancies ($B$ in nearest-neighbor sites) but adsorbs at a pair of free vacancies. There are no nearest-neighbor restrictions in adsorbing particles of different species. Only the adsorption-limited reactions are considered. Adsorbed dimers dissociate and a nearest neighbor of the adsorbed $A$ and $B$ particles reacts, forms the $AB$ product, and desorbs the catalytic surface immediately. Whenever there is an $A$ adsorption attempt at a vacant site between an adsorbed $A$ and an adsorbed $B$, we allow the $A$ to adsorb and react immediately with the neighboring $B$, thus forming the $AB$ product and desorbing the surface. If this process is not allowed, the IMD model possesses infinitely many absorbing states and is found to be always absorbing [24].

The system has no fully saturated phases of monomers or dimers, but instead two equivalent half-filled absorbing states. These states are comprised of only the monomers at the odd- or even-numbered lattice sites, i.e. $(A0A0\cdots)$ and $(0A0A\cdots)$ where “$A$” represents a monomer-occupied site and “0” a vacant site. These two states are probabilistically equivalent, unless we introduce a symmetry-breaking field discriminating the dynamics at the odd- and even-numbered sites. In this section, we consider the IMD model without a symmetry-breaking field. This model can be parametrized by the monomer adsorption-attempt probability $p$. The dimer adsorption-attempt probability is then given by $1 - p$.

In our previous stationary and dynamic Monte Carlo simulations [18], it was found that the system undergoes a continuous phase transition from a reactive phase into an absorbing phase, which belongs to the DI universality class. The kink representation of the IMD model is complicated due to its multi-component nature. Three types of kinks can be defined between lattice sites occupied by a dimer and a dimer, by a dimer and a vacancy, and by a vacancy and a vacancy. No conservation law is associated with each type of kinks, but the total number of kinks is conserved modulo 2. So, in a broad sense, one can say that the IMD model evolves by the parity-conserving dynamics like the BAWc model. In this section, we discuss the dynamic critical behavior of the IMD model via Monte Carlo simulations. Some of the results reported previously [19] are much improved using new efficient algorithms and some
new results for various interface dynamics are presented.

A. Defect dynamics

We start with a lattice occupied by monomers at alternating sites except at the central vacant site, i.e. \( (\cdots A0A0A0A0A0 \cdots) \) where 0 represents a defect at the central site. Then the system evolves along the dynamic rule of the model. After one adsorption attempt on the average per lattice site (one Monte Carlo time step), the time is incremented by one unit. A number of independent runs, typically \( 5 \times 10^3 \), are made up to \( 3 \times 10^4 \) time steps for various values of \( p \) near the critical probability \( p_c \). Most runs, however, stop earlier because the system enters into one of the absorbing states. We measure the survival probability \( P(t) \) (the probability that the system is still active at time \( t \)), the number of dimers \( N(t) \) averaged over all runs, and the mean square distance of spreading of the active region \( R^2(t) \) averaged over surviving runs. At criticality, the values of these quantities scale algebraically in the long time limit \( 2 \),

\[
\begin{align*}
P(t) &\sim t^{-\delta} \\
N(t) &\sim t^\eta \\
R^2(t) &\sim t^z
\end{align*}
\]

and double-logarithmic plots of these values against time show straight lines at criticality. Off criticality, these plots show curvatures. More precise estimates for the dynamic scaling exponents can be obtained by examining the local slopes of the curves. The effective exponent \( \delta(t) \) is defined as

\[
-\delta(t) = \frac{\log[P(t)/P(t/b)]}{\log(b)}
\]

and similar for \( \eta(t) \) and \( z(t) \). In this paper, we plot the effective exponents against \( 10/t \) with \( b = 10 \). Off criticality, these plots show positive or negative curvatures. The scaling exponents can be extracted by taking the asymptotic values of the effective exponents at criticality.

Our estimates for the critical probability and the dynamic scaling exponents are \( p_c = 0.5322(3) \), \( \delta = 0.290(5) \), \( \eta = 0.00(1) \), and \( z = 1.135(5) \) (see Fig. 1). Note that the estimate for \( z \) is much improved compared with our previous result; \( z = 1.34(20) \) \([19]\). These values are in excellent agreement with those of the DI universality class such as the BAW model \([26]\).

B. Interface dynamics

For the interface dynamics, we start with a pair of vacancies placed at the central sites of a lattice and with monomers occupied at alternating sites, i.e. \( (\cdots A0A0A0A \cdots) \) where the interface between two different absorbing states is placed in the middle of two central vacancies \( 00 \). In this case, the system never enters an absorbing state, so that the survival probability is always equal to 1 and the exponent \( \delta = 0 \). Even though the values of \( \delta \) and \( \eta \) vary with the types of dynamics, their sum \( \delta + \eta \) which is responsible for the growth of the number of kinks (or dimers) in surviving runs is known to be universal \([18,19]\). This guarantees that the generalized hyperscaling relation is always satisfied: \( \beta/\nu_\parallel + \delta + \eta = dz/2 \) where \( d \) is the spatial dimension and \( \beta \) and \( \nu_\parallel \) are steady-state exponents explained in the next section. As \( \delta = 0 \) in this type of interface dynamics, it does not supply any new information about dynamic critical behavior of the system.

In this section, we introduce three different types of interface dynamics which may give nontrivial scaling of the survival probability \( P(t) \). For convenience, the ordinary interface dynamics as above is called as type-A interface dynamics. Our previous study for the type-A interface dynamics found that \( \eta = 0.285(20) \) and \( z = 1.14(2) \) as expected \([19]\).

In the type-B interface dynamics, we stop the evolution if the interface collapses to its initial configuration, i.e. two vacant sites between the absorbing states. Then this run is treated as a dead one. This dynamics is originally introduced by Bassler and Browne for a three species monomer-monomer model \([30]\). At criticality, we measure \( P(t) \), \( N(t) \), and \( R^2(t) \). \( P(t) \) now represents not a true survival probability but a probability of avoiding a collapse. In Fig. 2, we plot the effective exponents against \( 10/t \). Our estimates for the dynamic scaling exponents are \( \delta = 0.73(1) \), \( \eta = -0.41(1) \) and \( z = 1.16(2) \). These values satisfy the generalized hyperscaling relation of the DI universality class and are in excellent accord with those reported by Bassler and Browne \([30]\). To see how much these exponents are robust, we change the criteria for stopping the evolution from two active sites to four and six active sites. We find
that there is no essential change in the values of the exponents. Therefore the nontrivial value of $\delta$ in the type-B dynamics is believed to be universal.

In the type-C interface dynamics, we focus only on the profile between the central site and the leftmost site (the left front) of the active region. We stop the evolution when the left front of the active region comes back to the center (initial position) and treat this run as a dead one (see Fig. 3). So $P(t)$ represents a probability of avoiding a collapse of the active region in the left side with respect to the initial location of the interface. We measure $N(t)$ as the number of dimers only in the left side of the center and $R^2(t)$ as the mean square spreading of the active region in the left side of the center. The type-C interface dynamics is useful when one needs to distinguish the behavior of the left and right fronts of the interface. So it is especially important to consider this type of dynamics for the interface between unconnected absorbing states (see Sec. III). The effective exponents against $10/t$ at criticality are plotted in Fig. 4. Our estimates are $\delta = 0.395(5)$, $\eta = -0.10(1)$ and $z = 1.150(5)$, which also satisfy the generalized hyperscaling relation of the DP universality class. The value of $\delta$ is different from those for the defect and other interface dynamics. It would be useful to check that this exponent is also universal for other models in the DP universality class.

Finally, we introduce the type-D interface dynamics. This dynamics is similar to the type-C dynamics in measuring physical quantities. But we do not stop the evolution when the left front of the active region hits the center. However, while its left front wanders in the right side of the center, we treat this run as a dead one temporarily, and set $N(t)$ and $R^2(t)$ to be zero. When it comes back to the left side of the center, we treat this run as a surviving run again and measure $N(t)$ and $R^2(t)$ as usual in the type-C dynamics. $P(t)$ represents a probability that the active region covers the left side of the center and is expected to converge to a nonzero constant less than 1. The effective exponents against $10/t$ at criticality are plotted in Fig. 5. Our estimates are $\delta = 0.010(5)$, $\eta = 0.29(1)$ and $z = 1.15(1)$. As expected, $\delta$ is nearly zero and $P(t)$ converges to $0.69(1)$ (see Fig. 6). So the type-D interface dynamics yields the same exponents as in the type-A (ordinary) dynamics. The type-A dynamics does not yield the correct scaling exponents for the interface between unequal absorbing states, so in this case the type-D dynamics can be employed instead (see Sec. III-C).

III. THE IMD MODEL WITH A SYMMETRY-BREAKING FIELD

We introduce a symmetry-breaking field which makes the system to prefer one absorbing state to the other [29]. This can be done by differentiating the monomer adsorption-attempt probability $p$ at an even-numbered vacant site and at an even-numbered one. If a monomer attempts to adsorb on an even-numbered free vacant site, the adsorption attempt is rejected with probability $h$ ($0 \leq h \leq 1$). The case $h = 0$ corresponds to the ordinary IMD model discussed in the previous section. For finite $h$, the monomers tend to adsorb more on an odd-numbered site than an even-numbered one. So the absorbing state with odd-numbered sites occupied by monomers is probabilistically preferable to the other absorbing state. However, the kink dynamics of this model still conserves the parity in terms of the total number of kinks. In this section, we show that the symmetry-breaking field forces the system back to the conventional DP universality class via stationary and dynamic simulations. Therefore one can conclude that not the parity conservation but the symmetry between the absorbing states is essential in determining the universality class of the absorbing phase transitions.

A. Stationary simulations

We run stationary Monte Carlo simulations starting with an empty lattice with size $L$. Then the system evolves along the dynamic rule with the symmetry-breaking field using periodic boundary conditions. We set the value of the symmetry-breaking field $h = 0.5$ for convenience. After a sufficiently long time, the system reaches a quasisteady state first and stays for a reasonably long time before finally entering into an absorbing state. We measure the concentration of dimers in the quasisteady state and average over many independent runs which have not yet entered into an absorbing state. The number of independent runs varies from $5 \times 10^3$ for the system size $L = 32$ to $3 \times 10^3$ for $L = 512$.

Elementary scaling theory combined with the finite-size scaling theory [30] predicts that the average concentration of dimers $\bar{\rho}$ at criticality in the steady state scales with system size $L$ as

$$\bar{\rho}(L) \sim L^{-\beta/\nu_{\perp}},$$

(3)

where $\beta$ is the order parameter exponent and $\nu_{\perp}$ is the correlation length exponent in the spatial direction. In the reactive phase ($p < p_c$), the concentration $\bar{\rho}$ remains finite in the limit $L \to \infty$, but it should vanish exponentially with system size in the absorbing phase ($p > p_c$).
At \( p_c \), we expect the ratio of the concentrations of dimers for two successive system sizes \( \tilde{\rho}(L/2)/\tilde{\rho}(L) = 2^{\beta/\nu_\perp} \), ignoring corrections to scaling. This ratio converges to 1 for \( p < p_c \) and approaches 2 for \( p > p_c \) in the limit \( L \to \infty \). We plot the logarithm of this ratio divided by \( \log 2 \) as a function of \( p \) for \( L = 64, 128, 256 \) and 512 in Fig. 7. The crossing points between lines for two successive sizes converge to the point at \( p_c \approx 0.413(1) \) and \( \beta/\nu_\perp \approx 0.23(2) \). The critical probability can be more accurately estimated from the defect dynamics simulations; \( p_c = 0.4138(3) \) (see the next subsection). We run stationary simulations at this value of \( p \) and find the better estimate for \( \beta/\nu_\perp = 0.243(8) \) (see Fig. 8). This value is consistent with the standard DP value of 0.2524(5) \([23]\).

By analyzing the decay characteristics of dimer concentrations at criticality, we can extract information about the relaxation time exponent \( \nu_\parallel \). From the elementary scaling theory, one can expect the short time behavior of the dimer concentration at criticality as

\[
\tilde{\rho}(t) \sim t^{-\beta/\nu_\parallel}.
\]

The characteristic (relaxation) time \( \tau \) for a finite system is defined as the elapsing time for a finite system to enter into the quasisteady state. Then one can find \( \tau \) scales at criticality as

\[
\tau \sim L^{\nu_\parallel/\nu_\perp}.
\]

In Fig. 9, we plot \( \tilde{\rho} \) at \( p_c \) versus time for various system sizes. Investigating the slopes in this double-logarithmic plot, we estimate \( \beta/\nu_\parallel = 0.165(10) \). The double-logarithmic plot (Fig. 10) for the characteristic time \( \tau \) versus the system size \( L \) shows a straight line from which we obtain \( \nu_\parallel/\nu_\perp = 1.45(10) \) which is consistent with the above results and agrees reasonably well with the standard DP value of 1.580(2).

Similar results are obtained for the system with a weaker symmetry-breaking field. Therefore we conclude that the symmetry-breaking field in the system with two absorbing states is relevant in determining the universality class and makes the system to behave like having a single absorbing state. In fact, the number of stationary runs falling into the unpreferred absorbing state compared with the number of those into the preferred state vanishes exponentially in system size (see Sec. IV-A). The preferred absorbing state behaves as a unique absorbing state of the system.

### B. Defect dynamics

When the symmetry between the absorbing states is broken, the defect dynamics are sensitive to the initial configurations. One can start with a lattice with a defect, either in the preferred absorbing state or in the unpreferred absorbing state. The latter case does not show critical spreading of the active region. The domain of the preferred absorbing state grows at the center of the active region while the region of the unpreferred absorbing state recedes with a constant speed. This dynamics is much like two interface dynamics between the preferred and unpreferred absorbing states where two interfaces move in the opposite direction. We will discuss the interface dynamics in the next subsection.

We choose the initial configuration with a defect at the center in the preferred absorbing state. We set \( h = 0.5 \) for convenience. Our estimates for the critical probability and the dynamic scaling exponents are \( p_c = 0.4138(3), \delta = 0.163(5), \eta = 0.315(5), \) and \( z = 1.265(5) \) (see Fig. 11). These values are also improved compared with our previous results \([23]\) and are in excellent accord with the standard DP values; \( \delta = 0.1596(4), \eta = 0.3137(10), \) and \( z = 1.2660(14) \).

### C. Interface dynamics

As the two absorbing states are not probabilistically equivalent, the dynamics of the interface between the preferred and unpreferred absorbing states shows a completely different behavior from the symmetric case. In Fig. 12, evolutions of the interface below, at, and above criticality are shown. For \( p < p_c \), the active region grows linearly in time in both directions. Of course, the left interface front near the preferred absorbing state (we call it \( P\)-interface front) moves slower than the right interface front (\( U\)-interface front) near the unpreferred absorbing state due to the symmetry-breaking field. At the critical point, the \( P\)-interface front behaves like a critical interface (zero velocity with nontrivial temporal scaling) as in the critical defect dynamics, but the \( U\)-interface front still moves with a finite velocity. As we increase \( p \) further into the absorbing phase, the active region cannot grow and the preferred absorbing state dominates. The \( P\)-interface front moves with the same velocity and the same direction as the \( U\)-interface front. The width of the active region is finite, in contrast to its diffusive behavior in the absorbing phase for the symmetric case. Here, we consider the critical case only.
In the symmetric case, four different interface dynamics are introduced. In the ordinary (type-A) interface dynamics, both of the exponents $\eta$ and $z/2$ converge to a trivial value of unity, due to the ballistic nature of the U-interface front even at criticality. The type-B interface dynamics also involve the dynamics of the U-interface front, so it does not yield nontrivial values of the exponents. The type-C and type-D interface dynamics focus only on the profile between the initial position of the interface (the center) and the P-interface front which behaves in a critical fashion.

For the type-C dynamics, we run $2 \times 10^4$ independent samples up to $3 \times 10^4$ time steps at $p_c = 0.4138$ with $h = 0.5$. Our estimates are $\delta = 0.37(1)$, $\eta = 0.11(1)$, and $z = 1.27(1)$ (Fig. 13). As expected, the value of $z$ agrees well with the DP value and these exponents satisfy the generalized hyperscaling relation for the DP universality class, i.e. $\delta + \eta = 0.48(2)$ is in excellent accord with the standard DP value of 0.473(1).

Similarly, we find $\delta = 0.02(2)$, $\eta = 0.45(1)$, and $z = 1.23(2)$ for the type-D dynamics (Fig. 14). Again these values agree well with the DP values. The concept of concentrating only on the relevant interface front may be applied to other types of models in which many types of interfaces coexist and some of them are not equivalent.

### IV. OTHER MODELS

#### A. Probabilistic cellular automata

Grassberger, Krause, and von der Twer [13,14] studied two models of probabilistic cellular automata (PCA), namely models A and B about ten years ago. The A model evolves with rule number 94 in the notation of Wolfram, except 110 and 011 configurations, where the central spin 1 flips to 0 with probability $p$ and remains unflipped with probability $1 - p$. The B model evolves with rule number 50 except 110 and 011 configurations, where the central spin 1 flips to 0 with probability $1 - p$ and remains unflipped with probability $p$. These are the first models investigated which are not in the DP universality class. Both models have two equivalent absorbing states, i.e. (1010· · ·) and (0101· · ·), and exhibit an absorbing phase transition which belongs to the DI universality class. But these models behave differently in the absorbing phase. Once the system enters into one of the two absorbing states, it remains in that state forever in model A but oscillates between these states in model B. In spite of the discriminating behavior, these models belong to the same DI universality class. In the kink representation, the total number of kinks are conserved modulo 2 in the dynamics.

First, consider the A model. We can introduce the symmetry-breaking field which makes the system to prefer (1010· · ·) to (0101· · ·). The system must go through a 111 configuration right before entering into an absorbing state, i.e. (· · ·010111010· · ·) → (· · ·010101010· · ·). If the flipping probability of the central spin in the 111 configuration depends on the even- or oddness of the position of the central spin, the symmetry between two absorbing states can be broken. With probability $h$, we reject the flipping attempt of the central spin in the 111 configuration when it is at an even-numbered site. Then the absorbing state with 1’s at the odd-numbered sites is probabilistically preferable to the other one. Again the parity in the total number of kinks is conserved even with the symmetry-breaking field $h$.

We run the defect and type-C interface dynamics with $h = 0.1$. Our estimates for the defect dynamics are $p_c = 0.2435(4)$, $\delta = 0.1625(25)$, $\eta = 0.310(5)$, and $z = 1.245(5)$ (Fig. 15). These values agree very well with the standard DP values. For the type-C interface dynamics at criticality, we find $\delta = 0.372(2)$, $\eta = 0.115(5)$, and $z = 1.275(5)$ (Fig. 16), which agree with the results for the IMD model with the symmetry-breaking field (see Sec. III-C). So the value of $\delta$ in the type-C dynamics seems to be universal. In order to check whether the value of $\delta$ in the type-C dynamics is universal in the symmetric case, we run the type-C interface dynamics when $h = 0$ and find $\delta = 0.395(5)$, $\eta = -0.08(2)$, and $z = 1.17(2)$ (Fig. 17), which is consistent with those found in the IMD model without the symmetry-breaking field (see Sec. II-B).

We also run stationary simulations at criticality with size $L = 32$ up to 512 with $h = 0.1$. Our estimates for the steady-state exponents are $\beta/\nu_L = 0.245(5)$, $\beta/\nu_\parallel = 0.155(5)$, and $\nu_\parallel/\nu_L = 1.63(5)$ (Fig. 18) which also agree reasonably well with the DP values. We measure the number of stationary runs falling into the unpreferred and preferred absorbing states respectively, i.e. $N_u$ and $N_p$ at criticality in the long time limit. The ratio $R = N_u/N_p$ versus system size $L$ is plotted in Fig. 19. This ratio vanishes exponentially in system size: $R \sim \exp(-L/L_0)$ with $L_0 \approx 13$. It means that the chance of entering into the unpreferred absorbing state is negligible so the system behaves like having a single absorbing state of the preferred one.

For the B model, the situation is quite different. The system oscillates between the two absorbing states; (1010· · ·) ↔ (0101· · ·). In this model, the system must go through a 000 configuration right before entering into an absorbing state, i.e. (· · ·101000101· · ·) → (· · ·010101010· · ·). We can introduce a rejection probability discriminating the even- and odd-numbered sites similar to that in the A model, but it cannot make the two absorbing states probabilistically inequivalent due to the oscillatory nature in the dynamics. In fact, it is meaningless to distinguish the two absorbing states without any dynamic barrier. In the A model, there exists a dynamic barrier between the two absorbing
states, which makes the system to take an infinitely long time to hop from near one absorbing state to near the other absorbing state. This dynamic barrier is similar to the free energy barrier between two ground states in the equilibrium Ising model. Without this barrier, there is no way to distinguish the two absorbing states. Therefore it seems impossible to find the crossover from the DI to the DP universality class in the B model.

B. Nonequilibrium kinetic Ising model

Nonequilibrium kinetic Ising model (NKI) recently introduced by Menyhárd [15–17] evolves with the competing effect of spin flips at zero temperature ($T = 0$) and nearest-neighbor spin exchanges at $T = \infty$. The spin-flip dynamics occurs with probability $p$ and spin-exchange dynamics with $1 - p$. The competition between the two different dynamics at different temperatures drives the system into a nonequilibrium steady state and there is a continuous phase transition as the competition parameter $p$ varies.

In the $T = 0$ spin-flip dynamics, the system evolves trying to lower the energy. A spin is allowed to flip only if the flip lowers the energy of the system or leaves it unchanged. We use the parameter $r$ to distinguish the cases when the energy is lowered or unchanged. We flip a spin with probability $r$ in the former case and flip a spin freely in the latter case. Here, we set $r = 0.5$. In the $T = \infty$ spin-exchange dynamics, nearest-neighbor spins are freely exchanged regardless of the energy change. Any up-down pair of spins can flip to the down-up pair of spins if they are in the nearest neighbor.

The absorbing phase consists of two completely ferromagnetically ordered states which are equivalent. One can not flip a spin in these absorbing states because it increases the energy. These absorbing states are the same as the two degenerate ground states of the equilibrium ferromagnetic Ising model. The absorbing transition of the NKI model belongs to the DI universality class. In terms of ordinary domain-wall language, a domain wall between different spins is interpreted as a walker in the BAW model. Then the NKI model can be mapped exactly to the BAW model with two offsprings with a control parameter for the two walker annihilation process [27][38].

We introduce a symmetry-breaking field which prefers up spins over down spins. So this field plays like an external magnetic field in the equilibrium Ising model. For convenience, we define the symmetry-breaking field $h$ as a probability of not allowing a up spin to flip. Therefore the absorbing state with all spins up becomes the preferred absorbing state. Again the parity in the total number of domain walls is still conserved even with the symmetry-breaking field.

We run the defect dynamics and stationary simulations with $h = 0.1$. Our estimates are $\rho_c = 0.190(5)$, $\delta = 0.17(1)$, $\eta = 0.32(1)$, and $z = 1.25(5)$ (Fig. 20), which agree well with the standard DP value. Stationary simulations at criticality yield $\beta/\nu_\perp = 0.24(1)$, $\beta/\nu_\parallel = 0.155(5)$, and $\nu_\parallel/\nu_\perp = 1.50(5)$ (Fig. 21) which also agree reasonably well with the DP values.

V. SUMMARY AND DISCUSSION

All models studied in this paper (IMD, PCA, NKI) preserve the parity of the total number of kinks but cross over from the directed Ising (DI) to directed percolation (DP) universality class when the parity-conserving symmetry-breaking field is introduced. As we argued in our preliminary paper [4], the essential factor which determines the universality class of a nonequilibrium absorbing phase transition is not the conservation laws in dynamics but the symmetry between absorbing states.

We take a careful look at various kinds of kinks in these models. First, consider the NKI model which is the simplest one in the domain wall (or kink) representation. In the NKI model, a kink is assigned between two neighboring spins in the opposite direction. Only one type of kinks exists in the NKI model and there is a two-to-one mapping between spin configurations and kink configurations. The two absorbing states correspond to the vacuum configuration in the kink representation. The evolution dynamics conserves the total number of kinks modulo 2. By identifying a kink as a walker in the BAW model, the NKI model can be exactly mapped to the BAW model with two offsprings with a control parameter for the two walker annihilation process [27][38]. Numerical results [22][26] and recent field theoretical works for the BAW models [3][30] suggest that the parity conservation is responsible for the DI universality class. The symmetry-breaking field (magnetic field) in the NKI model cannot be represented by a local kink operator in the field theory language, similar to the magnetic field in the equilibrium Ising model in the Bloch wall representation. Therefore the recent field theoretical results by Cardy and Täuber [3][40] do not apply when the symmetry-breaking field is introduced. In order to consider the symmetry-breaking field, one should include the long-range string operator, i.e. the global product of the number operators of kinks in the quantum Hamiltonian, which becomes a highly nontrivial problem. Our numerical results suggest that this long-range string operator is relevant and makes the system to leave a DI fixed point and flow into a DP fixed point by the renormalization group transformations.
The PCA models are similar to the NKI models even though there seems no trivial mapping into the BAW models. These models contain two types of kinks which are assigned between two neighboring 1’s and 0’s respectively. There is no kink between 1 and 0, so the two absorbing states correspond to the vacuum in the kink representation. Due to the parallel updating procedure, it is not easy to examine whether these two types of kinks can be represented by one kink operator in the field theory. But the parity of the total number of kinks is conserved during the evolution. The symmetry-breaking field which discriminates the even-numbered and odd-numbered sites should be also represented by a long-range string operator like that in the NKI model. However, the B model does not have any dynamic barrier between two absorbing states due to the oscillatory nature, so is not affected by the symmetry-breaking field. Except that, we can draw the same conclusion for the PCA models as in the NKI model.

The IMD model has a more complex kink representation due to its multi-component nature. Three different types of kinks are found between $B$ and $B$, $B$ and 0, 0 and 0, where $B$ is a site occupied by a dimer atom and 0 is a vacancy. The parity of the total number of kinks is conserved and the symmetry-breaking field is similar to that in the PCA models. Our numerical results show that the IMD model exhibits the same critical behavior as in the NKI and PCA model. It may imply that the differences between various kinks in the IMD and PCA models are just irrelevant details which do not affect the universal behavior. It may be interesting to study these differences in the field theoretical models.

Recently, a few other models have been introduced with equivalent absorbing states. Those are generalized monomer-monomer models studied by Bassler and Browne [30–32], and generalized Domany-Kinzel models and generalized contact processes studied by Hinrichsen [33]. These models are multi-component models so there are many types of kinks. Unlike the IMD model, there appears no explicit parity conservation law in the kink representation of these models, even though they all have two equivalent absorbing states. Numerical simulations showed that these models belong to the DI universality class. By introducing a symmetry-breaking field, these models cross over from the DI to DP universality class as in the models studied in this paper. These results strongly support our conclusion that the symmetry between absorbing states, not the conservation law, is the essential property to determine the universality class of the absorbing phase transitions. However, we do not exclude the possibility of the hidden conservation law in the kink dynamics of these models. In their absorbing phase, numerical simulations show that large domains of two different absorbing states are formed and active regions between two different domains (domain walls with finite width) survive and diffuse until they annihilate pairwise [33]. It implies that there may be an effective parity conservation law in domain walls, even though there is no parity conservation in microscopic kinks. With the symmetry-breaking field, large domains of the unpreferred absorbing state completely disappear and domain walls (active regions) annihilate by themselves.

In this paper, we also introduce and investigate various interface dynamics. Without the symmetry-breaking field, we find new universal exponents for $\delta$ in the type-B and type-C interface dynamics, but the hyperscaling relation for the DI universality class is always intact. With the symmetry-breaking field, the interface moves, in average, with a constant velocity in the direction of the unpreferred absorbing state. By focusing only on the side of the preferred absorbing state, we find new exponents for $\delta$ for the type-C and type-D dynamics and the hyperscaling relation for the DP universality class is obtained. These new exponents are also shown to be universal. These types of interface dynamics should be useful in studying some other models with many but unequivalent absorbing states.

It is interesting to compare the two dimensional equilibrium Ising model with the one dimensional NKI model. It is well known that two dimensional equilibrium models are related to one dimensional kinetic models via transfer matrix formalism [1]. An extra space dimension is interpreted as the time dimension in one dimensional kinetic models. One can write down the evolution operator of the kinetic model corresponding to a given two dimensional equilibrium model. This evolution operator is Hermitian for the equilibrium model. General nonequilibrium kinetic models in one dimension can be obtained by modifying the above Hermitian evolution operator in a non-Hermitian form, i.e. breaking the detailed balance. Then equilibrium models in $d$ dimensions and nonequilibrium kinetic models in $d - 1$ dimensions can be directly compared.

The NKI model is a special case of general nonequilibrium kinetic Ising models. In the NKI model, the time reversal symmetry is broken completely and its dynamics favors one time direction over the other. Therefore, by adding a directional sense in the time direction to the equilibrium Ising model, one can see the crossover from the equilibrium Ising universality class to the nonequilibrium DI universality class.

Similar things happen for the models with a single absorbing state which belong to the DP universality class. The percolation problem is equivalent to the $q \rightarrow 1$ limit of the $q$-state Potts model [12]. The DP problem is defined by adding a directional sense to the percolation problem. Similarly, one can define nonequilibrium models with $q$ equivalent absorbing states by adding a directional sense to the $q$-state Potts model. Both models have the permutation symmetry between $q$ ground (or absorbing) states, so these nonequilibrium models may be called as the $q$-state directed Potts model. In this sense, the NKI model may be called as the directed Ising model and the NKI universality class as the directed Ising (DI) universality class. It will be interesting to set up and investigate the directed Potts models with $q \geq 3$ [33]. Of course, other types of generalized models with the rotational or cyclic
symmetry are also interesting.

There is a difference in connecting models with different values of \( q \). Crossover from the DI to the DP universality class is obtained by introducing a symmetry breaking field. But the Ising phase transition disappears when the magnetic field is applied to the two dimensional equilibrium Ising model. The percolation universality class appears only through the random cluster formulation of the Potts model \[3\]. Therefore the analogy between the \( q \)-state Potts model and the \( q \)-state directed Potts model is not complete. These similarities and differences should be further investigated in future.

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FIG. 1. Plots of the effective exponents against $10/t$ for the defect dynamics of the symmetric IMD model. Three curves from top to bottom in each panel correspond to $p = 0.5315$, 0.5322, and 0.5329.

FIG. 2. Plots of the effective exponents against $10/t$ for the type-B interface dynamics of the symmetric IMD model at criticality ($p_c = 0.5322$).

FIG. 3. The type-C interface dynamics in the IMD model. Grey, black, and white dots represent monomers, dimers, and vacancies, respectively.

FIG. 4. Plots of the effective exponents against $10/t$ for the type-C interface dynamics of the symmetric IMD model at criticality.

FIG. 5. Plots of the effective exponents against $10/t$ for the type-D interface dynamics of the symmetric IMD model at criticality.

FIG. 6. $P(t)$ versus $10/t$ for the type-D interface dynamics of the symmetric IMD model.

FIG. 7. Plots of $\log\left(\bar{\rho}(L/2)/\bar{\rho}(L)\right)/\log(2)$ versus $p$ for the asymmetric IMD model with the symmetry-breaking field $h = 0.5$ for various system sizes $L = 64, 128, 256$, and $512$.

FIG. 8. The average concentration of dimers $\bar{\rho}$ at criticality in the quasisteady state against the system size $L$ in a double-logarithmic plots for various system sizes $L = 32 – 512$ for the asymmetric IMD model with $h = 0.5$. The solid line is of slope $-0.243$.

FIG. 9. Time dependence of the average concentration of dimers at criticality for various system sizes $L = 32 – 512$ for the asymmetric IMD model with $h = 0.5$. The solid line is of slope $-0.165$.

FIG. 10. Size dependence of the characteristic time at criticality for various system sizes $L = 32 – 512$ for the asymmetric IMD model with $h = 0.5$. The solid line is of slope $1.45$.

FIG. 11. Plots of the effective exponents against $10/t$ for the defect dynamics of the asymmetric IMD model with $h = 0.5$. Three curves from top to bottom in each panel correspond to $p = 0.4132$, 0.4138, and 0.4144.

FIG. 12. Evolutions of the asymmetric IMD interface dynamics for (a) $p < p_c$, (b) $p = p_c$, and (c) $p > p_c$. The region of the preferred (unpreferred) absorbing state is shown in black (grey). The active sites are represented by white pixels.

FIG. 13. Plots of the effective exponents against $10/t$ for the type-C interface dynamics of the asymmetric IMD model for $h = 0.5$ at criticality.

FIG. 14. Plots of the effective exponents against $10/t$ for the type-D interface dynamics of the asymmetric IMD model for $h = 0.5$ at criticality.
FIG. 15. Plots of the effective exponents against $10/t$ for the defect dynamics of the asymmetric PCA A model with $h = 0.1$. Three curves from top to bottom in each panel correspond to $p = 0.2443$, $0.2435$, and $0.2427$.

FIG. 16. Plots of the effective exponents against $10/t$ for the type-C interface dynamics of the asymmetric PCA A model with $h = 0.1$ at criticality.

FIG. 17. Plots of the effective exponents against $10/t$ for the type-C interface dynamics of the symmetric PCA A model at criticality.

FIG. 18. The kink density in the quasisteady state against the system size, time dependence of the kink density at $L = 512$, and size dependence of the characteristic time are plotted for the asymmetric PCA A model with $h = 0.1$ at criticality. The solid lines are of slope $-0.245$, $-0.155$, and $1.63$ from top to bottom.

FIG. 19. The semi-logarithmic plot for the ratio $R$ against system size $L$. The solid is $R = 1.44 \exp(-L/12.9)$.

FIG. 20. Plots of the effective exponents against $10/t$ for the defect dynamics of the asymmetric NKI model with $h = 0.1$. Three curves from top to bottom in each panel correspond to $p = 0.18$, $0.19$, and $0.20$.

FIG. 21. The kink density in the quasisteady state against the system size, time dependence of the kink density at $L = 512$, and size dependence of the characteristic time are plotted for the asymmetric NKI model with $h = 0.1$ at criticality. The solid lines are of slope $-0.24$, $-0.155$, and $1.50$ from top to bottom.
fig 1
Fig 2
fig 4
fig 5
fig 7

\[ \log \left( \frac{\rho(L/2)}{\rho(L)} \right) / \log 2 \]

- ■ 64
- ○ 128
- ▲ 256
- ▼ 512

\( p \)

Y-axis: \( \log \left( \frac{\rho(L/2)}{\rho(L)} \right) / \log 2 \)

X-axis: \( p \)
Fig 8

The graph shows a plot of \( \log \rho \) against \( \log L \). The data points are connected by a straight line, indicating a linear relationship between the two variables.
fig 9

\[ \log \rho(t) \]

\[ \log t \]
fig 15
fig 17
fig 18
fig 21