Breaking of forward-backward symmetry in driven systems

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The dynamical pair approximation was modified to study the stationary states in a two-dimensional repulsive-lattice-gas model driven far from equilibrium by the application of an external field. This approximation distinguishes between the forward, backward, and transverse directions with respect to the electric field. In the present driven system, the forward-backward symmetry is broken at the level of the pair approximation. The difference between the forward and backward directions is confirmed by Monte Carlo simulations.

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Phase transitions far from equilibrium are extensively studied in very different physical systems. A well known type of nonequilibrium state may be produced by the application of a homogeneous electric field causing persistent particle transport (for a recent review see the paper by Schmittmann [1]). The driven lattice-gas model was first suggested by Katz, Lebowitz, and Spohn [2] as a means of investigating nonequilibrium behavior. They observed that the electric field modifies the particle distribution as well as the ordering transition. This process is accompanied by anisotropic correlations. In the literature [3–5] the transverse and longitudinal directions are distinguished; however, the forward and backward directions are assumed to be equivalent. In fact, the difference between the forward and backward directions has not yet been investigated in these systems except for the one-dimensional lattice gas [6]. In this former model, for example, the probabilities of the (0010) and the reflection-symmetric (0100) configurations at four subsequent sites are not equivalent and the difference is an odd function of the electric field. In this Brief Report we demonstrate that the breaking of forward-backward symmetry may also be observed for a two-dimensional system. For this purpose we analyze the effect of electric field on the particle distribution in a half-filled lattice gas with a repulsive nearest-neighbor interaction. In this model the breaking of the forward-backward symmetry may be observed at the level of pair distribution because the particle distribution is not translation invariant in the ordered phase.

The stochastic dynamics of driven lattice gases is characterized by single-particle jumps to one of the empty neighboring sites (Kawasaki dynamics). The jump rates are biased by a uniform electric field parallel to one of the principal directions [1, 2]. In other words, the jumps are favored along, and suppressed against the field direction. The difference between the forward and backward jump rates results in modified particle distribution in which the forward-backward (reflection) symmetry is broken. As mentioned above the breaking of the forward-backward symmetry may be observed as a difference between the probabilities of some pairs of k-point configurations. In a previous paper [6] we have shown that the probabilities of the (01) and (10) configurations are equivalent if the lattice sites are occupied uniformly. This equivalence is no longer valid if the particle distribution exhibits a long-range order characterized by two sublattices with different occupations. This situation is characteristic of the equilibrium state of a square lattice gas with a repulsive nearest-neighbor interaction.

Adopting the previous notations we choose the particle charge, the coupling, lattice, and Boltzmann constants to be unity. This system exhibits a chessboardlike ordered structure below a critical temperature $T_c = 0.5673$ [7]. In the ordered structure two sublattices (A and B) may be distinguished in which the average occupations are different; $n_A = 1/2 + x$ and $n_B = 1/2 - x$, where $x$ ($|x| \leq 1/2$) is the order parameter. Henceforth we assume that the particles reside dominantly in sublattice A, that is, $x > 0$.

In lattice-gas models the particle jumps are blocked by the occupancy of the target point. Consequently, the particle fixed at a given point causes congestion in the driven system, i.e., the average occupancy is higher at the preceding point than at the subsequent one. In the ordered structure the particle positions are fixed by the long-range order. This phenomenon led us to distinguish the forward and backward directions when analyzing pair correlations in such systems in which the particles prefer to reside in one of the sublattices.

The present driven lattice-gas model has been studied by Leung, Schmittman, and Zia [4] using renormalization-group technique and by Dickman [5] using dynamical pair approximation, respectively. These authors have shown that the critical temperature decreases with the electric field and the phase transition becomes first order above a threshold value ($E_{th} \approx 1.49$ for the Kawasaki hopping rate). Furthermore, the critical temperature becomes zero at $E = 2$ and the ordering
is completely suppressed at zero temperature if $E > 3$
[4].

Our method may be considered as an extension of the
dynamical pair approximation used by Dickman for two-
dimensional driven lattice gases [5, 8]. We have deter-
mined the stationary solution of the equations describing
the time evolution of sublattice occupation and proba-
bility of (1,1) pairs along the mentioned directions (for
details see Refs. [5, 6, 8]). These equations summarize
the loss and gain contributions of elementary jumps for
all the possible configurations on the sites affecting the
jump rates. The probability of the configurations is de-
termined at the level of pair approximation. Instead of
the uniform longitudinal pair distribution function, we
introduce two quantities: $p_f(1,1)$ and $p_b(1,1)$, the for-
ard and backward pair probability functions where the
first argument refers to an occupied site of the preferred
sublattice, the second ones to the subsequent and preced-
ing sites, respectively. It is underlined that the present
distinction between $p_f(1,1)$ and $p_b(1,1)$ is reasonable in
the ordered phase for $E \neq 0$.

The above mentioned equations are solved numeri-
cally for different temperatures and electric fields. The
Kawasaki hopping rate is used to avoid the singular-
arity that comes from the nonanalytic behavior of the
Metropolis rate, namely the probability of jumping from
site $j$ to $k$

$$
\Gamma(j \to k) = \frac{1}{1 + \exp(\beta(AH - E\Delta r))},
$$

where $\Delta H$ is the energy difference between the final and
initial states, $\Delta r$ is the displacement, and $\beta = 1/T$.
The temperature dependence of the order parameter $x(T)$ is
very similar to those found by Dickman [5], thus his main
conclusions remain unchanged. Evidently, our method re-
produces the results obtained by Dickman for $E = 0$ and
$E \geq 2$ where there is no difference between $p_f(1,1)$ and
$p_b(1,1)$. The critical temperature is slightly de-
creased when taking the symmetry breaking into con-
cideration for $0 < E < 2$.

In Fig. 1 the solid line illustrates the calculated differ-
ence between $p_f(1,1)$ and $p_b(1,1)$ as a function of reduced
temperature $T/T_c$ ($T_c$ means the equilibrium critical
temperature) for a fixed electric field $E = 0.4$. This differ-
ence is strongly related to the existence of the order para-
meter $x$; therefore, it disappears for high temperatures.
Our calculation shows that this symmetry breaking is
suppressed by the ordering process at low temperatures.

Monte Carlo simulations have been carried out to check the
prediction of the present dynamical mean-field ap-
proximation. Using the usual techniques [9] the runs were
performed on a half-filled lattice with periodic boundary
conditions for different sizes ($L = 40, 64, 128, 256$). Each
run was started from an ordered state. After a thermal-
ization of 2000 Monte Carlo steps per particle (MCS)
we determined $p_f(1,1)$ and $p_b(1,1)$ by averaging over
$10^4$ MCS. To decrease the statistical error we repeated
this procedure 10 times at a given temperature. By this
means we could avoid the uncertainty that comes from
averaging in a polydomain structure which may appear
for long time simulations below $T_c$. Rigorous analysis of

![FIG. 1. Difference between $p_f(1,1)$ and $p_b(1,1)$ as a func-
tion of reduced temperature at a fixed electric field $E = 0.4$. The
solid line illustrates the theoretical prediction, the bullets represent
Monte Carlo data.](image)

the formation of the polydomain structure goes beyond
the scope of the present paper [10].

Figure 1 demonstrates that the Monte Carlo data show
qualitative agreement with our theoretical calculation in
the single domain state. The size dependence seems to
be irrelevant when measuring the above mentioned pair
distribution functions.

Several simulations have been performed to visualize the
typical elementary processes below $T_c$. Each particle
residing in an ordered surrounding can jump and may
return to the original site at low temperatures. In other
words, one can observe particle-hole (Frenkel) pairs with
a finite lifetime. The contribution of particle-hole pairs
to $p_f(1,1)$ and $p_b(1,1)$ depends on their relative ori-
entation. According to Eq. (1) the probability of forward
(backward) jumps is greater (less) than that of the trans-
verse ones. Consequently, the particle-hole pairs are "po-
larized" and give different contributions to $p_f(1,1)$ and
$p_b(1,1)$.

Theoretical descriptions including the present one as-
ume the existence of an ordered structure with infinite
range of order. The breaking of forward-backward sym-
metry, however, is independent of the presence of long-
range order. On the analogy of the one-dimensional sys-
tem [6] this symmetry breaking is also observable for high
temperatures, i.e., this phenomenon is not related to the
phase transition. This fact is demonstrated by Monte
Carlo simulations in Fig. 2 where the difference between
the probability of the four-point configurations (1100) and
(0011) on four subsequent sites along the field is plot-

![FIG. 2. Difference between the probability of four-point config-
urations (1100) and (0011) vs temperature for $E = 0.4$. The
arrow indicates the equilibrium critical temperature $T_c$.](image)
ted as a function of temperature for a fixed electric field $E = 0.4$. The simulations were carried out on a $200 \times 200$ lattice by averaging over 50,000 MCS. As expected, this type of symmetry breaking is suppressed by the ordering process at low temperatures and by the thermal fluctuations at high temperatures. It is emphasized that the same difference is found between other pairs of four-point configurations [i.e., $(0101)$ and $(1010)$], in agreement with the one-dimensional results [6].

In summary, we have studied the forward and backward symmetry breaking in a half-filled two-dimensional lattice-gas model with a repulsive nearest-neighbor interaction. In this system the symmetry breaking may be investigated by introducing two pair probability functions, $p_f(1,1)$ and $p_b(1,1)$, along the driving field in the ordered (single-domain) state. These quantities were analyzed by using Monte Carlo simulations and by the modification of the dynamical mean-field theory within the framework of pair approximation. In the single-domain state the difference between $p_f(1,1)$ and $p_b(1,1)$ depends on the electric field, its value is strongly related to the order parameter, and it is suppressed by the ordering process in the low-temperature limit. The breaking of forward-backward symmetry, however, is observable between other $k$-point ($k \geq 4$) configurations in agreement with the one-dimensional calculations. Monte Carlo simulations show that the symmetry breaking itself exists in this driven system independently of the presence of long-range order. The value characteristic of this symmetry breaking depends on $E$ and has a maximum below the equilibrium critical temperature.

It is expected that forward-backward symmetry breaking is characteristic of all driven lattice gases and it should be taken into consideration when rigorously analyzing the ordering process.

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[10] The polydomain structure as a consequence of enhanced particle transport along the domain wall will be the subject of a subsequent paper.