Influence of long-range interactions on charge ordering phenomena on a square lattice

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I. INTRODUCTION

The formation of ordered structures is one of the main topics in the field of condensed matter physics. Starting from the relative straightforward crystalline order a wide variety of increasingly complex ordering phenomena has been observed and proposed such as stripes [1–8], charge density waves [9], incommensurate phases [10], and so forth.

These complex ordering patterns usually arise as a result of competing interactions. For example, the kinetic energy of holes competes with the tendency towards antiferromagnetic order in cuprates thus forming stripes. In anisotropic next-nearest-neighbor Ising (ANNNI) models, the next-nearest-neighbor Ising coupling has the opposite sign as the nearest neighbor coupling. The question immediately arises whether higher-order commensurate or incommensurate phases can appear in systems with only one type of interaction.

Of course this is the case. In the continuum the sole presence of long-range interactions will cause particles to form a Wigner crystal. When a fixed number of particles are placed on an underlying lattice the desired Wigner crystalline order may be incommensurate with the lattice, thus leading to frustration.

We have investigated the influence of long-range interactions on charge ordering phenomena on a square lattice. Expanding the results of Refs. [11, 12] we explored the full range of particle densities $0 \leq \rho \leq 1$ and types of long-range interactions $V = 1/r^p$. Our main result is summarized in Figs. 1 (zero temperature) and 2 (finite temperature), where we depict phase diagrams of usual charge ordered patterns. At low densities the competition between the continuum triangular Wigner lattice and the underlying square lattice indeed leads to a plethora of “generalized Wigner” crystals. At higher densities, this leads to variations of the checkerboard pattern which is well-known at half-filling. Stripe phases appear as they are rooted in the topological defects of the checkerboard order.

We do not claim that the phase diagrams we derived are the exact phase diagrams. As is often the case for frustrated systems, a large set of metastable states persists down to zero temperature. Unbiased numerical computation of the energy for a large ensemble of configurations gives us a strong indication that indeed the phase diagram of Fig. 1 is correct; however, these indeed be metastable states incorrectly recognized as ground states.

The layout of this paper is as follows. In Sec. II we introduce the lattice gas model, which is the model describing interacting classical particles on a lattice. In the two subsequent sections we discuss qualitatively the ordered structures at low densities (Sec. III) and at densities close to half-filling Sec. IV. We have performed a Monte Carlo simulation in Sec. V to derive the zero temperature phase diagram of Fig. 1. In Sec. VI we extend these results to finite temperatures using mean field theory; see Fig. 2.

II. LONG-RANGE LATTICE GAS MODELS

The lattice gas model can be defined on any kind of lattice, but we focus only on the square lattice. On each of the $N$ lattice sites there can be a particle or not, denoted by $n_i = 1$ or 0 respectively. These particles interact via some general potential $V_{ij}$. The corresponding Hamiltonian is then

$$H_L = \sum_{i \neq j} V_{ij}(n_i - \rho)(n_j - \rho) - \mu \sum_i n_i.$$  (1)

We subtract the average particle density $\rho$ to prevent divergent energies. In the grand-canonical ensemble, the chemical potential $\mu$ tunes the average particle density $\rho N = \sum_i n_i$. The model (1) is in fact equivalent to the Ising model [13]. Under the replacement $\sigma_i^z = 2n_i - 1$ and considering only a nonzero nearest neighbor interaction $\frac{1}{2} V_{ij} = J$ one finds

$$H_I = J \sum_{(ij)} \sigma_i^z \sigma_j^z - B \sum_i \sigma_i^z.$$  (2)

The chemical potential maps onto an external magnetic field $B = \frac{1}{2} \mu - V_{ij}$, while the particle density maps onto the mean magnetization.

For the ferromagnetic Ising model the ground state is completely magnetically ordered, which amounts to either a full or empty lattice in the lattice gas parlance. In addition, a model with antiferromagnetic coupling will be half-filled with particles if the external magnetic field is small, $|B| < 2J$. 

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FIG. 1. (Color online) The approximate ground state phase diagram of the long-range lattice gas model on a square lattice, based on variational methods as discussed in Sec. V. On the vertical axis the type $p$ of the long-range interaction $V(r) = \frac{1}{r^p}$ is given, together with a logarithmic decaying interaction. The horizontal axis represents the particle density. From low to high densities we identify the following phases: The area without name depicts the “checkerboard-in-a-checkerboard” 1/9 glassy phase described by Ref. [12], the 1/4 Wigner crystal, the “checkerboard-in-a-checkerboard” 1/4’ phase, stripe phases (with a plateau for the 1/3 stripe phase and “C” denotes the channelled stripes as described by Ref. [11]), and finally the checkerboard phase. Phases below 1/4 filling are discussed in Sec. III, above 1/4 filling are discussed in Sec. IV. Below the phase diagram typical particle configurations in six phases are shown enlarged.

Therefore, using the standard grand canonical ensemble will in general not enable us to investigate all possible particle densities: a canonical ensemble, fixed particle number, is required.

We argue that most physical realizations of lattice gas models are in fact at a fixed particle number, and not at fixed chemical potential. One particular example is the oxygen ordering in YBCO planes, where it is beyond doubt that the number of oxygen ions in the lattice is fixed [14]. The patterns in which the oxygen ions align themselves are quasi-one-dimensional, in a manner similar to the expected electronic ordering in TTF-TCNQ salts [15]. While studying the latter, Hubbard has developed a general solution for the ground state of a lattice gas model with long-range interactions at any particle density in one dimension. Hubbard’s solution requires only the interaction energy as a function of distance to be convex.

Among two-dimensional realizations of lattice gas models are for example the ordering of ad-atoms on a surface [10,16,17], $XY$ systems [18], higher order commensurate magnetic phases [10,19,20], or stripe order in high-temperature superconductors [21–23]. Systems with anisotropic short-ranged interactions or competing short- and long-range interactions [24,25] have especially acquired considerable attention over the years; therefore we wish to focus here on the case of long-range isotropic interactions [26–29].

Most studies of lattice gas models in two dimensions, however, restrict their attention to half-filled, empty and full lattices, due to the aforementioned grand-canonical reasons. There are two notable exceptions: the stripe order discussed in Ref. [11] between 1/3 and 1/2 filling and the glassy dynamics at 1/6 filling [12]. These results were obtained for a “quasi-logarithmic” repulsive interaction, which is a solution of Poisson’s equation on a lattice, $V^2V_{ij} = -2\pi \delta_{ij}$. Given the nontrivial ordering patterns discovered there, as a follow-up we present here a systematic study of the ground state orderings at all densities between 0 and 1/2, for general repulsive interactions

$$V_{ij} = \frac{1}{|r_{ij}|^p} > 0.$$  

(3)

In the next two sections we will first discuss qualitatively such long-range lattice gas models at fixed densities, while in the remaining sections the picture will be further quantified using numerical simulations and mean field theory.

III. DILUTE DENSITIES: GENERALIZED WIGNER CRYSTALS

In the previous section we introduced the lattice gas model, which we will now study at fixed densities on the square lattice with long-range repulsive interaction of the form (3). In the limit of very low particle density, the underlying square lattice becomes irrelevant compared to the average interparticle distance,

$$\ell_p \gg a,$$  

(4)
of a Wigner crystalline state of the particles is [16] in the Wigner crystal. The distance between particles ρ is the particle density, and the summation runs over the density. Therefore the energy in the low density limit scales as

\[ E \propto J N \rho^{p/2+1}. \]

The presence of the underlying lattice is now a source of frustration. Reference [16] considers an underlying triangular lattice, leading to frustration only if the density is not of the form 1/p^2. For example, when ρ = 1/9 a perfect triangular Wigner crystal can be formed. On top of a square lattice, however, it is not possible to form a triangular Wigner crystal because \( \sqrt{3} \) is irrational. One can nevertheless construct “almost perfect” triangular crystals. As an example, consider the density ρ = 1/9; see Fig. 3. The lowest energy state is there also a triangular crystal of particles, but not equilateral as for a perfect Wigner crystal. In principle such “almost perfect” Wigner crystals could exist at densities

\[ \rho_{pq} = \frac{1}{pq} \]

with p,q integers while p ≤ q, such that the following equilateral triangle relation is approximated by

\[ p \sim \frac{1}{2} \sqrt{3} q. \]

For example, the densities ρ = 1/9, 1/12, 1/16, etc. would allow such “almost perfect” triangular crystal. Following the work of Hubbard [15] in one-dimensional systems, we will call these particle orderings “generalized Wigner crystals.” From this qualitative reasoning we argue that such Wigner crystals might exist. Note, however, that one has to resort to a numerical computation to find whether such crystals have indeed the lowest energy at a given density.

So far we considered only densities of the form \( \rho_{pq} = 1/pq \). When the density of a lattice gas is between such densities, we suspect that it is favorable to maintain a generalized Wigner crystal structure. The deviation from the \( \rho_{pq} \) density can be accommodated by a superlattices of crystal defects or interstitial vacancies. If such a superlattice of defects forms, the original \( \rho_{pq} \) order is still visible in, for example, the Fourier transformed particle density, where each crystal type has its own specific Fourier peaks. We expect therefore a “plateau” at densities in the vicinity of each specific \( \rho_{pq} \), where the associated crystal structure remains intact modulo the interstitial superlattice. As we will describe in more detail later when discussing the numerical results, indeed a “plateau” is observed for the 1/4, 1/6, and 1/9 states. Using mean field theory, we went as far as the 1/25 crystal phase, as is shown in Fig. 4. We have therefore strong indications the plateau structure exists all the way to \( \rho \to 0 \), yielding an infinite staircase of plateaus. This structure is reminiscent of the devil’s staircase, as exists in the case of the one-dimensional lattice gas in the grand-canonical ensemble [31], where specific charge orderings are stable for a finite window of chemical potential.

Notice that starting at 1/4 filling the generalized Wigner crystal picture certainly fails. If one adds one single particle to the 1/4 crystal, it will be necessarily next to another particle. Since the nearest-neighbor repulsion is the strongest, and nearest-neighbor occupancy can be avoided for any density below half-filling, the 1/4 crystal will be quickly destroyed upon adding particles. For densities above 1/4 it is necessary to start reasoning from the ordering occurring at the half-filled lattice.

IV. DOMAIN WALLS AND STRIPES

Exactly at half-filling the ground state is “checkerboard” like, or antiferromagnetic in the Ising language. This means that one sublattice is exactly filled and the other sublattice is completely empty. Densities slightly less than half-filling
can be obtained by removing particles from the checkerboard pattern, a process we call hole doping. The density of holes $\rho_h$ is defined as follows:

$$\rho_h = 1 - 2 \rho,$$

where $\rho$ is the total particle density. The same scaling arguments for the dilute particle limit $\rho \ll 1$ can be applied to the dilute hole limit $\rho_h \ll 1$, so close to half-filling the energy scales as

$$E \propto E_{1/2} (1 - 2 \rho_h) + J N c h \rho_h^{p/2+1},$$

where $E_{1/2}$ is the energy of the half-filled checkerboard configuration and $c_h$ is some proportionality constant.

This scaling argument assumes that all particles will remain on the filled sublattice of the checkerboard phase. However, the checkerboard phase is a broken sublattice symmetry phase, and therefore domain walls and topological defects can exist between regions where the checkerboard phase is realized on different sublattices. Though the ground state at densities slightly less than half-filled might be unrelated to the checkerboard phase, we discuss in this section possible structures that are related to the checkerboard. Thus the smallest example, which is neglected in the scaling arguments for the dilute particle limit $\rho \ll 1$ can be applied to avoid getting stuck in a local energy minimum. Upon reducing the temperature to zero, was therefore also used to obtain whether the ground state is a Hubbard-type stripe pattern or a "doped" stripe pattern. Lee et al. [11] call these doped stripe patterns "partially filled diagonal channels." Finally, we must emphasize that the discussion in this section does in no way whatsoever constitute a proof of existence of stripe ordered or doped checkerboard phases. The only way to find the state with the lowest energy is a tedious numerical computation.

FIG. 5. Some examples of domain walls between checkerboard phases. (a) A particle on the “wrong” sublattice surrounded by “holes,” which forms the smallest possible domain wall loop. (b) A horizontal domain wall. (c) Instability of a horizontal domain wall, by moving one particle to the other domain. (d) Stable, diagonal domain walls can exist for all long-range interactions.

FIG. 6. Some examples of stripe phases. (a) The $\rho = 1/3$ state. (b) The $\rho = 2/5$ state. (c) The $\rho = 3/7$ state, but now doped.

macroscopic number of domain walls. Such a periodic array of domain walls constitutes a “stripe phase,” similar to the ones discussed in cuprates [1,2,6], nickelates [3], manganites [4–6], or cobaltates [7,8]. Examples can be seen in Fig. 6.

If the ground state of a long-range lattice gas model is perfectly stripy, then the system is effectively reduced to a one-dimensional system. The arguments of Hubbard [15] then apply, and one can thus find the specific stripe ordering, as is shown in Fig. 6(a) and 6(b).

But again, for some densities it may pay off not to form perfect stripes but rather to dope a stripe structure as in Fig. 6(c). It is then a matter of numerical computation to find out whether the ground state is a Hubbard-type stripe pattern or a “doped” stripe pattern. Lee et al. [11] call these doped stripe patterns “partially filled diagonal channels.” Finally, we must emphasize that the discussion in this section does in no way whatsoever constitute a proof of existence of stripe ordered or hole doped checkerboard phases. The only way to find the state with the lowest energy is a tedious numerical computation.

V. SIMULATIONS AND CHARACTERISATION OF THE PHASES

Let us now describe the numerical algorithm that was used to find the lowest energy configurations. First, notice that for each form of order there is symmetry breaking and hence a degeneracy in the ground state. As the simplest example, observe that the checkerboard configuration at half-filling is twofold degenerate. However, for the purpose of finding the specific type of charge order, we do not need to worry about ground state degeneracy.

In our algorithm we took various different types of initial configurations, such as the generalized Wigner crystals of Sec. III, stripe structures of Sec. IV, Wigner crystal structures of defects in the checkerboard, variations of the checkerboard phase, suggestions from literature, and a large ensemble of random configurations. We then swapped filled and empty sites randomly. A swap is accepted if it lowers the total energy of the system. The long-range nature of the interaction was taken into account by summing the interaction over all mirror charges as in an Ewald summation [33]. For the quasi-logarithmic interaction we followed the method of Ref. [11].

The major issue is that one cannot know for sure whether this algorithm leads to the global ground state or that one gets stuck in a local energy minimum. Indeed, for a frustrated system we expect to find a large number of metastable states. The method of simulated annealing, by which we mean slowly reducing the temperature to zero, was therefore also used to avoid getting stuck in a local energy minimum. Upon comparing the energies of configurations obtained from the
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Our work was performed on a square lattice with 90 \times 90 and 154 \times 154 sites. These lattice sizes were chosen such that the linear dimension \( L \) is divisible by the prime numbers 2, 3, 5 (for \( L = 90 \)) and 7 and 11 (for \( L = 154 \)). We looked at all densities that were a multiple of the linear dimension, hence \( \rho = n/90 \) and \( \rho = n/154 \) for all integers \( n = 1, \ldots, L/2 \).

After obtaining the ground state configuration, a Fourier transform of the particle density was taken:

\[
 n(k) = \frac{1}{N} \sum_{i} e^{-i k \cdot r_i} n_i. \tag{12}
\]

The peaks in the Fourier spectrum were used to identify the specific orderings at each density, as can be seen in Fig. 7. The ground state energy as a function of density is shown in Fig. 8, rescaled such that the energy at half-filling equals \( E = 1 \). Indeed, the general scaling behavior close to zero and half-filling, as described in the previous two sections, is retrieved. This is most explicit in the limit of \( p \to \infty \), the energy becomes constant between \( \rho = 0 \) and \( \rho = 1/4 \), and linear between \( \rho = 1/4 \) and \( \rho = 1/2 \). Notice an extra kink in the energy around \( \rho = 1/3 \), which signals the onset of the stripe order.

With the caution that the resulting configurations might be in fact metastable states incredibly close to the actual ground state, we constructed the zero temperature phase diagram in Fig. 1. At low densities the finite lattice size used in our numerical simulations form a limitation with regard to the precision of the results. The first unequivocal observed charge ordering state is the 1/9 generalized Wigner crystal phase, which is stable for a considerable range of densities around \( \rho = 1/9 \). Interestingly, the presence of this phase is remarkably independent of the interaction range \( p \).

The 1/9 phase is followed by the 1/6 phase, which is extensively discussed in the work of Ref. [12]. There the 1/6 phase is characterized as a glassy phase, with infinite ground state degeneracy due to the infinite ways one can tile unit cells of \( 2 \times 3 \) lattice sites. For further details we refer to Ref. [12].

Directly below \( \rho = 1/4 \) densities the 1/4 generalized Wigner crystal phase is stable. As described before, introduction of particles to densities higher than \( \rho = 1/4 \) leads to a superlattice of interstitials which can be called a “checkerboard-in-a-checkerboard 1/4’ phase,” which seems to be absent in the case of logarithmic interactions.

Here the logarithmic interactions seem to play a special role, in that for a much larger density regime than for algebraic interactions the stripe phases seem to be stable. We find, contradictory to the results of Ref. [11], only stripe order of the Hubbard kind and no channeled stripes in the region between 1/4 and 4/9; see Fig. 9. Only the 1/3-stripe order seemed to be present for a larger region of densities. We have found “partially filled diagonal channels,” or equivalently doped stripe orders, only for interaction types \( p = 1 \) and 2 in a very narrow density range. Our approach differs from Ref. [11] in two aspects: we have looked at the ground state, while they looked at low temperature results obtained by simulated annealing only, and we considered nonlocal particle swaps in the configuration while they restricted the system to local swaps only. We have compared the energy of the ground states we found with the explicit ground states of Ref. [11], taking

\[
 \frac{E(p=1/2)}{E(p=1/4)} \approx 9 \times 10^{-5}\% \quad \text{lower in energy than the perfect stripe order.}
\]

For logarithmic interactions the perfect stripe order is 9 \times 10^{-5}\% lower in energy than the channeled stripe order, in contrast to Ref. [11] which finds channeled stripes here. For \( p = 1 \) interactions the channeled stripes are 0.0014\% lower in energy than the perfect stripes.
into account the specific commensurability with the finite lattice size of their doped stripe orders. Figure 9 gives an example of this energy comparison at \( n = \frac{29}{47} \). For all densities that we checked, it was found that our ground state energies are lower, be it only by a very small amount.

Notice that for algebraic interactions, the stripe phase shifts to higher densities when \( p \) increases. For large \( p \), the stripe phase seems to dominate in the whole region between 1/3 and 1/2 fillings.

We also computed the ground state phase diagram for interactions with a finite screening length of the form

\[
V(r) = \frac{e^{-r/\lambda}}{r^p}
\]

turning into a Bessel function \( K_0(r/\lambda) \) for the “screened logarithmic” interaction. We find that the corresponding phase diagram for screened interactions is largely similar to Fig. 1, with only small quantitative differences that do grow with decreasing screening length \( \lambda \).

VI. FINITE TEMPERATURE

The phases described in the previous section survive at finite temperatures, because we are dealing with a system with discrete symmetry in two dimensions. We have computed a finite temperature mean field phase diagram using standard mean field theory [34], for the 1/r interaction (\( p = 1 \)).

It is of course highly questionable whether mean field theory correctly describes the competition between various phases. For example, the rich phase diagram of the ANNNI model is constructed using mean field theory [10], but corrections beyond mean field are shown to tip the delicate balance between different phases and reduce the critical temperature [35]. At the same time Monte Carlo simulations of the ANNNI miss incommensurate phases that are clearly present in analytical extensions of mean field theory [10]. Results from the ANNNI model thus suggests that mean field theory acts as a qualitatively reliable first approximation towards the understanding of complex ordering patterns.

Let us now briefly summarize the quintessence of our mean field theory. For the model Hamiltonian (1) we postulate an ansatz for the density,

\[
\langle n_i \rangle = \bar{n} + \sum_m m_a \cos(Q_a \cdot r_i) \tag{14}
\]

where there can be as many ordering wave vectors \( Q_a \) as one needs to correctly describe the specific charge order. For example, the 1/9 order has ordering wave vectors \( Q_1 = (0, \frac{2\pi}{9}) \), \( Q_2 = (\frac{2\pi}{9}, \frac{4\pi}{9}) \) and all linear combinations inside the first Brillouin zone. When \( m_a \neq 0 \) the translational invariance is spontaneously broken. Using the ansatz (14) one constructs a mean field Hamiltonian

\[
H_0 = \sum_i \left[ -\mu + \sum_a m_a V_{a} \cos(Q_a \cdot r_i) \right] n_i \tag{15}
\]

We then minimize the thermodynamic potential

\[
\Phi = F_0 + \langle H - H_0 \rangle \tag{16}
\]

with respect to the mean field parameters \( m_a \), where \( F_0 = -\frac{1}{p} \log \text{Tr} e^{-\beta H_0} \) and \( \langle \cdot \cdot \cdot \rangle_0 \) implies a thermal average with respect to the mean field Hamiltonian \( H_0 \). Every charge order we found at zero temperature acts as a possible ansatz, and we numerically minimize at each temperature and density the thermodynamic potential \( \Phi \).

Since mean field theory gives only a qualitative phase diagram, and because the zero-temperature phase diagram of Fig. 1 suggests little qualitative difference between various interaction ranges, we restrict ourselves to the Coulomb interaction \( V = 1/r \). The resulting phase diagram is shown in Fig. 2. In addition to the 1/9 Wigner crystal phase we also considered 1/12, 1/16, 1/20, and 1/25 crystals. As for the stripe phases, we studied only the 1/3 and the 3/7 “channeled” state. The phase diagram we thus find indeed matches the zero-temperature phase diagram obtained by numerical simulations of the previous section. We emphasize that further studies are needed to understand the possible incommensurate stripe phases in between 1/3 and 4/7 filling.

The transition to the checkerboard phase and the similar “checkerboard-in-a-checkerboard” 1/4’ phase is of the second order type within the Ising universality class [27]. The thermodynamic potential for some temperatures around \( T_c \), with its typical second order transition behavior, is shown in Fig. 10(a). The question arises whether this 1/4’ “phase” is an artefact of the mean field theory. As is known at half-filling, a liquid-like state with local checkerboard order exists in the presence of long-range interactions [36]. Such correlated liquid phase can also be present away from half-filling, but will be beyond the scope of standard mean field theory.

The phase transitions to more complicated orders are always of the first order kind, an example of which is shown in Fig. 10(b) at 1/9 filling. This is a natural result because transitions from solid to liquid phases are usually discontinuous [37]. The existence of such first order transition implies that one can supercool the high-temperature (or 1/4’) state [38]. At \( \rho = 1/3 \), for example, the 1/4’ phase remains a local minimum of the free energy even though its energy is higher than that of the stripe phase; see Fig. 10(c). This suggests that it might be hard to actually trap the system in the lowest energy state.

![FIG. 10. (Color online) Thermodynamic potential relative to the disordered state \( \Delta \Phi \), in arbitrary units, plotted around various phase transitions. (a) At \( n = 0.27 \) there is a second order phase transition into the 1/4’ state, clearly visible by the Mexican hat potential. The second order phase transition is common for transitions in the Ising universality class. (b) At \( n = 1/9 \) there is a first order transition toward the Wigner crystal, as is customary for solidification transitions. (c) At \( n = 1/3 \) there are two transitions: a second order transition into the 1/4’ state followed by a first order transition towards the stripe phase. The 1/4’ and the stripe phases are locally stable however for a longer range of temperatures. This implies the possibility of supercooling the 1/4’ phase.](image)
In combination with our earlier observation that a correlated liquid-like state at intermediate temperatures might exist, we then also expect glassy physics upon supercooling [39]. With “glassy” we mean that there are macroscopically many local minima of the free energy, leading to slow relaxation rates. There is a difference however with the glassy physics found at $\rho = 1/6$ densities [12]. There the glassy nature is a ground state property, where glassy physics around a first order transition vanishes if the temperature is low enough.

The mean field theory shows the possibility of supercooling, the consequences thereof such as possible glass-like behavior needs to be addressed differently. Finite temperature numerical simulations however have the great disadvantage that they get easily stuck in such a complicated free energy landscape. It remains thus an open challenge to quantitatively describe the consequences thereof such as possible glass-like behavior. It is these domain walls that cause the formation of stripes.

The finite temperature phase diagram has been obtained using mean field theory, yielding only a qualitative description. Numerical and/or analytical extensions of the classical mean field theory will increase the accuracy of the finite temperature phase diagram. Thereby one can address the possibility of supercooling and glassy physics.

A possible next step is to include the kinetic energy of the particles present. This also allows for an extension to quantum particles [40] or $O(n)$ spin variables [41] instead of the classical particles we have considered thus far.

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other structure has been found with a lower energy, the Wigner crystallization into a triangular lattice is not rigorously proven. For classical particles studied in this publication the exact energy is equal to the “quantum” second order perturbation result since that amounts to the expectation value of the interaction energy. For logarithmic interactions we refer to studies of vortex lattices that indicate that triangular lattices are in that case the lowest energy configurations [32]. For $p > 1$ interactions we compared the energy of the triangular lattice with the square lattice, which again favors the triangular lattice.