Melting of Colloidal Molecular Crystals on Triangular Lattices

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Abstract

The phase behavior of a two-dimensional colloidal system subject to a commensurate triangular potential is investigated. We consider the integer number of colloids in each potential minimum as rigid composite objects with effective discrete degrees of freedom. It is shown that there is a rich variety of phases including “herring bone” and “Japanese 6 in 1” phases. The ensuing phase diagram and phase transitions are analyzed analytically within variational mean-field theory and supplemented by Monte Carlo simulations. Consequences for experiments are discussed.

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In the last decade it has been realized that soft materials can serve as versatile model systems to study phenomena of condensed matter physics. In particular, two-dimensional (2D) systems of colloidal particles interacting with light sources are ideal to mimic the adsorption of atoms and molecules on atomic surfaces, vortices in superconductors with periodic pinning arrays, and many other related phenomena [1, 2, 3]. Experimental studies on flat substrates [4] have beautifully confirmed the existence of a two-stage melting process mediated by the successive unbinding of dislocations [5] and disclinations [6]. If confined to a 1D periodic potential 2D colloidal systems show even richer behavior with re-entrant melting [7] and novel phases such as the locked floating solid [8, 9]. Results from analytical theories have been complemented by a series of numerical simulations (see e.g. Refs. [10, 11, 12]).

Recent experimental [13, 14] and theoretical [15, 16] investigations have studied the effect of 2D periodic potentials on the phase behavior. In particular, experiments on triangular light lattices [13] with a stochiometry of three colloids per site have motivated our research. One observes that even at rather low potential strength the colloids tend to group as trimers forming an almost equilateral triangle, but with a quite significant number of defects, i.e., groups of two or four particles. Upon increasing the potential strength the number of defects decreases rapidly such that they become unobservable already for moderate intensities. Then, the important low energy excitations are the orientation of the trimers, regarded as rigid composite objects, with respect to the lattice direction. Due to the interaction of the trimers long-range orientational order is expected for sufficiently strong coupling. Alignment of the trimers is observed experimentally [13] as soon as the defect-free regime takes over. Interestingly, the same authors also observed a loss of orientational long-range order at even higher intensities, which was interpreted as re-entrant melting and seems to be confirmed by computer simulations [15].

In this Letter we derive phase diagrams for commensurate colloidal systems in high external fields by analytic methods supplemented by Monte-Carlo (MC) simulations. The key idea is to reduce the problem to the low energy degrees of freedom by considering the integer number of colloidal particles which gather in a single potential minimum as a rigid composite object. Its shape is determined by the symmetry of the lattice and the number of the constituting colloidal particles, and its size by the interplay of interparticle repulsion and external potential. Short-time orientational fluctuations close to the potential minima are
considered to be already averaged out leaving only a discrete set of gross orientations. We will mainly focus on *dimers* on a triangular lattice since they exhibit a rich phase diagram and exhibit a series of intriguing phases. At the end of the letter we will shortly report how the phase diagrams for trimers can be obtained in a rather straightforward manner.

For an isolated dimer there are three equivalent orientational ground states on a triangular lattice, denoted by $\sigma = 1, 2, 3$ (see Fig. 1). The effective interaction between the dimers results from the screened Coulomb interaction between the constituent colloids, which is short-ranged for the experimental conditions in Ref. [13]. Therefore, it is appropriate to restrict the effective dimer–dimer interaction to a nearest neighbors. A pair of dimers can be in $3^2$ configurations corresponding to one of four energy levels, with generic ordering $E_1 < E_2 < E_3 < E_4$; see Fig. 1. The relation between these model parameters and the experimental control parameters like screening length and potential strength can be worked out in detail [17], but is far too involved to be presented here. As a guidance there are the following trends. Increasing the laser potential compresses the dimers and thus by increasing the effective distances between neighboring dimers reduces their interaction energies and hence the values of $E_i$ and the spacings between them. This is the basic mechanism underlying orientational melting. Lowering the screening length effectively corresponds to increasing the potential strength.

The Hamiltonian of our model system reads

$$\mathcal{H} = \sum_{\alpha=1}^{3} \sum_{\langle i,j \rangle_{\alpha}} \mathbf{\sigma}_i \cdot H_{\alpha} \cdot \mathbf{\sigma}_j,$$

(1)

where $\langle i,j \rangle_{\alpha}$ denotes the pair of nearest neighbors whose bond-vector is parallel to the orientation of the dimer state $\alpha$. $\mathbf{\sigma}_i$ is a vector representation of the dimer configuration $\sigma_i$. 

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**FIG. 1:** The model system. (a) The triangular lattice of the external field and the three orientational states of the dimers. (b) Four energy levels for the pair-wise interactions of dimers.
at the site $i$: $1 \rightarrow \sigma = (1, 0, 0)$, $2 \rightarrow \sigma = (0, 1, 0)$, and $3 \rightarrow \sigma = (0, 0, 1)$. For the energy matrices $H_\alpha$ one finds in terms of a permutation matrix

$$H_1 = \begin{pmatrix} E_4 & E_3 & E_3 \\ E_3 & E_1 & E_2 \\ E_3 & E_2 & E_1 \end{pmatrix}, \quad P = \begin{pmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 1 & 0 & 0 \end{pmatrix}$$

$$H_2 = P^\top H_1 P, \quad H_3 = PH_1 P^\top.$$ \hfill (2)

Since a shift of the global energy scale does not affect the phase behavior only the three energy differences in units of $k_B T$ constitute dimensionless parameters. Thus one would be led to expect a three-dimensional phase diagram. However, upon rewriting the Hamiltonian in “spin language” it turns out that the parameter space can actually be reduced to only two dimensions \[17\]. We find, omitting an additive constant

$$H = -K \sum_{\langle i,j \rangle} \delta_{\sigma_i,\sigma_j} - M \sum_{\alpha=1}^3 \sum_{\langle i,j \rangle} \delta_{\sigma_i,\alpha} \delta_{\sigma_j,\alpha},$$ \hfill (3)

where the new energy scales $K, M$ are given in terms of the direct dimer interaction energies by $K = -E_1 + E_2$ and $M = E_1 - 2E_2 + 2E_3 - E_4$. This reduction greatly simplifies the analysis and facilitates comparison with future experiments with the two generic control parameters, screening length and laser intensity \[22\].

For $M = 0$ the Hamiltonian reduces to a 3-state Potts model. The critical properties for the 2D $q$-state Potts model for $q \geq 4$ can be determined exactly for ferromagnetic (FM) interactions, $K > 0$. The critical point is known rigorously for square, triangular, and honeycomb lattices \[18, 19\]. There is sufficient numerical evidence that the $q \geq 4$ solution also holds for $q = 3$ although an analytic proof is still missing. The FM–P (paramagnetic) phase transition of the 3-state Potts model is continuous as suggested by the assumed “exact” solution and corroborated by MC simulations and renormalization group analysis \[19\]. Interestingly, a mean-field description gives qualitatively wrong results, since it predicts a discontinuous transition. For negative exchange coupling $K < 0$ the low temperature phase is an antiferromagnetic (AM) Potts state. The results of renormalization group, series expansions, and MC simulations for the transition on a triangular lattice appear to be controversial as far as the nature of the transition is concerned \[19\].

The phase diagram of dimers in the external potential forming a triangular lattice as prescribed by a generalized Hamiltonian, Eq. \[3\], has a richer topology. Generically $K \neq 0$, \[4\].
FIG. 2: Zero temperature ordered structures of 2D colloidal dimers: (a) ferromagnetic (FM), (b) herring bone (HB), (c) Potts antiferromagnetic (AM), and (d) Japanese 6 in 1 (J6/1), structure. In the latter, the energy of the system does not depend on the orientation of dimers on sites denoted by circles, thus this structure is $3^{N/4}$ degenerate.

and one distinguishes between $K > 0$ and $K < 0$. In both cases there are two dimensionless parameters, the ratio of the exchange couplings $\tilde{M} = M/|K|$ and the normalized temperature $\tilde{T} = k_B T/|K|$.

An exact analytic solution of the whole phase diagram, except for the two Potts points, is difficult to obtain and one has to rely on approximative and/or numerical methods. We start with a mean-field (MF) analysis which allows us to determine the topology of the phase diagram and thereby the symmetry of the order parameter in the various low temperature phases. In particular, we employ a variational MF approach where the full density matrix is approximated as a product of single-site density matrices, $\rho(\{\sigma_i\}) = \prod_i \rho_i(\sigma_i)$. By choosing site-independent density matrices the order parameter of the FM phase has been found by minimizing the variational free energy. Appropriate generalizations have been introduced to obtain AM phases and other ordered phases; details will be presented elsewhere [17].

From the MF analysis we find the equation of state for the respective order parameters, and in particular the phase boundaries. It turns out that the variational free energy for colloidal dimers in the P/FM or AM phase is identical to the corresponding result for the Potts model, provided one substitutes the Potts exchange energy $K$ by the effective coupling $K \rightarrow K_{\text{eff}} = K + M/3$. The critical temperatures for the P–FM and P–AM phase transition are located at $k_B T_c = 3K_{\text{eff}}/(2 \ln 2)$ and $k_B T_c = -3K_{\text{eff}}/(4 \ln 2)$, respectively.

The colloidal dimer Hamiltonian, Eq. (3), allows for new intriguing structures which are not realized within the Potts model. We have investigated for herring bone (HB) and Japanese 6 in 1 (J6/1) structures – a term borrowed from weaving patterns for chainmailles.
The corresponding order parameters can be obtained by (numerically) minimizing the variational free energies and the phase boundaries can be determined accordingly; see Fig. 3. Except for the HB–P transition for \( \tilde{M} < -7/3 \) within MF theory all phase transitions are discontinuous. The HB–P transition has already been studied in the context of N\(_2\) adsorbates on graphite. A continuous anisotropic planar rotor model analogous to our discrete version for \( K = 0 \) yielded similar results both within MF and MC description \([20, 21]\).

In our case, the MF description is not expected to predict the correct order of the transition as can be inferred by specializing to the Potts model (\( M = 0 \)). Nevertheless, the MF results should constitute reasonable approximations for the actual phase behavior. In order to gain further insight we have performed extensive MC simulations of the colloidal dimer Hamiltonian, Eq. (3), using a standard Metropolis procedure. The observables have been averaged over \( \geq 5000 \) statistically independent configurations after the system has been equilibrated. Every 1000 MC cycles a new configuration contributing to the averages is considered. Simulations have been performed on several system sizes, \( N = L \times L \) and \( L = 12, 36, 54, 108 \), and periodic boundary conditions have been employed to mimic bulk-like conditions \([17]\). The order parameter characterizing the various phases has been chosen in
analogy with the Potts model, either defined on the whole system or suitable sublattices,

\[ S = \frac{1}{N} \left[ N_\Sigma - \frac{1}{2} \sum_{\sigma \neq \Sigma} N_\sigma \right]. \tag{4} \]

Here, \( N_\sigma \) is the number of dimers in state \( \sigma \), in particular, \( N_\Sigma \) corresponds to majority orientation, and \( N = \sum_\sigma N_\sigma \) is the total number of the dimers on the whole (sub)lattice. We have performed temperature sweeps for \( K > 0 \) and \( K < 0 \) with a number of choices for \( \tilde{M} \) starting from the ordered structures. We have monitored the average order parameter and energy, as well as their respective variance. The transition temperatures corresponding to melting of the ordered structures have been determined by extrapolating the location of the maxima of the heat capacity and susceptibility to the infinite system size. In order to decide whether the respective phase transition is strongly discontinuous or compatible with a continuous/weakly discontinuous transition scenario we have evaluated Binder’s cumulant and respective probability distributions for the order parameter and energy \[17\]; both scenarios are exemplified in Fig. 4.

**FIG. 4:** Order parameter \( S \) as a function of the reduced temperature for the two FM–P transitions. 
Left: Potts-like continuous transition for \( K, M > 0 \). Right: the \( \tilde{M} \)-term driven first order transition for \( K < 0, \tilde{M} > 5 \). Inset: probability distributions of states with respect to the energy per dimer, \( u = U/N \), below, above, and at the transition temperature.

Figure 3 exhibits the phase diagram for the colloidal dimer Hamiltonian; boundaries resulting from both MF description and MC simulations are included in the Figure. The variational MF free energy reproduces the topology of the phase diagram, although some of the transitions become continuous by fluctuation effects. Quantitatively, MF overestimates the transition temperatures with respect to simulation results by a factor of order unity, a feature already familiar from the Ising model \[18\]. Interestingly, the discontinuous HB–P transition appears to saturate for large negative values of \( \tilde{M} \) in strong contrast to the corresponding MF result \[23\]. For colloidal systems the regime corresponding to \( K > 0 \)
and $\tilde{M} < 0$ should be experimentally accessible. A stringent test of our theory would be to experimentally verify the predicted HB–P phase transition.

Fluctuation effects reduce the critical temperature of the Potts model by a known factor $[19]$. The variational MF free energy of the colloidal dimer Hamiltonian for P/FM and AM structures is equivalent to the MF Potts model, provided that the Potts exchange energy is replaced by the effective coupling $K_{\text{eff}}$. This analogy suggests an empirical improvement of the MF result for these transitions by rescaling the transition temperatures with the same respective correction factors. The result of this procedure is shown as dashed lines in Fig. 3. For $K > 0$ there is excellent agreement with simulation results for temperatures $\tilde{T} > 1$. For $K < 0$ the mapping gives an overall quantitative improvement of the phase diagram. In particular, for $M = 0$ the well known Potts solution is recovered.

We close with reporting our main results for trimers on a triangular lattice. Due to the symmetry of the lattice, trimers can only be in one of two orientational states, represented by Ising spins $\sigma = \pm$. Hence the four configurational states for trimer pairs contribute one out of the three possible interaction energies,

$$E_1 < E_2 < E_3.$$  

In contrast to the Ising model the interaction energies for the $(+-)$ and $(-+)$ configurations are different. The relation between the model parameters and the control parameters of the experiments are similar as for dimers. In particular, an increase of the potential strength again leads to a reduction of the interaction energies between trimers and thus favors orientational melting.

The central questions now are how many relevant energies determine the phase behavior and what is the effective statistical model. Similar considerations as for dimers would suggest that there are two relevant energy scales resulting in a two-dimensional phase diagram. Actually, by a simple rearrangement the Hamiltonian can be cast in the form of a Ising model with a single excitation energy $\Delta E = E_1 + E_3 - 2E_2$ corresponding to single trimer flips, irrespective of the configurations of the neighboring trimers. Depending on the sign of $\Delta E$ the ground state is either “ferromagnetic” with parallel $(++)$ (or equivalently $(--)\) configurations or “antiferromagnetic” with an alternating arrangement of $(+-)$ and
configurations. It is interesting to note that it is the relative orientation of trimer pairs and not the trimers itself which determines the ground state.

For the remainder of the analysis known exact results of the 2D Ising model can now be employed to obtain the phase diagram. Considering the parameters of the experimental set-up in Ref. we can identify the critical laser intensity, , above which the orientational order of trimers is lost. This result is consistent with experimental observations, where the colloidal trimer system was found to be orientationally ordered for but disordered for . It would be interesting to extend the experimental analysis to find the actual transition point and compare it with our prediction.

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Both upon increasing the potential strength and the screening length the ratio $|M|/K$ becomes smaller.

Single-dimer flips of the HB ground state correspond to excitation energies $2K$ and $2|M|$ depending on the two possible new orientations of the dimer. For $-M \gg K$, $M$-type excitations are strongly suppressed. However, the MF approach, in which generically the role of $M$ is to renormalize the energy $K$ in a linear way, misses that.