Traveling and resting crystals in active systems

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A microscopic field theory for crystallization in active systems is proposed which unifies the phase-field-crystal model of freezing with the Toner-Tu theory for self-propelled particles. A wealth of different active crystalline states are predicted and characterized. In particular, for increasing strength of self-propulsion, a transition from a resting crystal to a traveling crystalline state is found where the particles migrate collectively while keeping their crystalline order. Our predictions, which are verifiable in experiments and in particle-resolved computer simulations, provide a starting point for the design of new active materials.

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Self-propelled particles \[1\] exhibit fascinating collective phenomena like swarming, swirling and laning which have been intensely explored by theory, simulation and experiment, for recent reviews see \[2,4\]. In marked contrast to passive particles, self-propelled “active” particles have an internal motor of propulsion, dissipate energy and are therefore intrinsically in nonequilibrium. Examples of active particles include living systems, like bacteria and microbes \[5\], as well as man-made microswimmers, catalytically driven colloids \[6,7\] and granular hoppers \[8\].

If, at high densities, the particle interaction dominates the propulsion, crystallization in an active system is conceivable. It is expected that such “active crystals” have structural and dynamical properties largely different from equilibrium crystals due to the intrinsic drive. In fact, there is experimental evidence for active crystals, both from observations of hexagonal structures for catalytically-driven colloids \[9\] and honeycomb-like textures for flagellated marine bacteria \[10,11\]. Moreover, recent computer simulations have confirmed crystallization \[12,14\] and proved that melting of active crystals differs from its equilibrium counterpart. However, though field-theoretical modelling of active systems has been widely applied to orientational ordering phenomena \[2,13\], there is no such theory for translational ordering of active crystals nor has a systematic classification of active crystals been achieved.

Here we present a microscopic field-theoretical approach to crystallization in active systems and we propose a minimal model which has the necessary ingredients for both, crystallization and activity. In doing so, we combine the phase-field crystal model of freezing \[10\] with the Toner-Tu model for active systems \[17\] using the concept of dynamical density functional theory \[18,19\]. On the one hand, the phase-field-crystal (PFC) model as originally introduced by Elder and coworkers \[16,20\] describes crystallization of passive particles on microscopic length and diffusive time scales. When brought into connection with dynamical density functional theory \[21,23\], the PFC model represents in principle a microscopic theory for crystallization, and it has been successfully applied to a plethora of solidification phenomena \[16,20,25,29\]. On the other hand, Toner and Tu \[17\] investigated the onset of collective motion in self-propelled systems from a general hydrodynamic point of view. Phenomenological coupling parameters of this model can in principle be justified by dynamical density functional theory \[20\], too, but it does not describe crystallization.

In our PFC model for active systems, we find a wealth of different crystallization phenomena. First, we identify two different types of active crystals which we call “resting” and “traveling” depending on their averaged drift velocity. A resting crystal possesses vanishing net particle flux whereas a traveling crystal is migrating with a nonzero velocity while keeping its periodicity. Starting from a disordered initial state, a traveling crystal is typically formed by a coarse-graining process of domains. The threshold in the driving strength upon which traveling crystals are formed depends on the spontaneous local orientational order (as prescribed by the coupling parameters of the bare Toner-Tu model): if there is no such order, the threshold is finite, while there is no such threshold in the presence of spontaneous orientational order. We further identify a transition from a hexagonal to a rhombic traveling crystal if the drive is increased further and find also resting and traveling lamellar phases with one-dimensional periodic ordering. Finally the occurrence of honeycomb-like structures can be explained as well within our model. The knowledge and control over these crystalline states provides an attractive starting point for the design of novel active materials since active crystals possess unique structural, phononic, and rheological properties.

In the following, we first describe our model and then discuss numerical and analytical results. Our dynamical equations are for the local one-particle density field $\psi_1(\mathbf{r},t)$ which is a conserved scalar order parameter and basically describes the reduced density modulation around a fixed averaged density $\bar{\psi}$ as in the traditional
PFC model \[16, 20\], and for a polarization vector field \(\mathbf{P}(r, t)\) which describes the local polar ordering. Activity enters into the equations via a nonzero self-propulsion velocity \(v_0\). In suitably scaled units of time, length and energy, our basic dynamic equations read

\[
\partial_t \psi_1 = \nabla^2 \frac{\delta \mathcal{F}}{\delta \psi_1} - v_0 \nabla \cdot \mathbf{P},
\]

(1)

\[
\partial_t \mathbf{P} = \nabla^2 \frac{\delta \mathcal{F}}{\delta \mathbf{P}} - D_r \frac{\delta \mathcal{F}}{\delta \mathbf{P}} - v_0 \nabla \psi_1.
\]

(2)

Here, \(D_r\) is the rotational diffusion coefficient of the particles and \(\mathcal{F}\) is a free energy functional of \(\psi_1\) and \(\mathbf{P}\) gained from density functional theory. The equations (1) and (2) are consistent with phenomenological symmetry arguments and involve the simplest nontrivial coupling between the two order parameter fields \(\psi_1\) and \(\mathbf{P}\). As outlined in the supplemental material \[31\], they can also be derived from microscopic dynamical density functional theory within an appropriate gradient and Taylor expansion of the order parameter fields \[19, 32\], see also \[33\]. In the sequel, we shall consider two spatial dimensions only.

We now further specify the free energy functional \(\mathcal{F}\) to \(\mathcal{F} = \mathcal{F}_{pfc} + \mathcal{F}_\mathbf{P}\) where

\[
\mathcal{F}_{pfc} = \int d^2 r \left\{ \frac{1}{2} \psi_1 \left[ \varepsilon + (1 + \nabla^2)^2 \right] \psi_1 + \frac{1}{4} \psi^4 \right\}
\]

(3)

is the traditional PFC functional \[16, 20\] describing the tendency of the material to form periodic structures. Here, \(\varepsilon\) sets the temperature \[16, 20\], and the order parameter \(\psi\) corresponds to the total density \(\psi = \bar{\psi} + \psi_1\). The polarization-dependent part

\[
\mathcal{F}_\mathbf{P} = \int d^2 r \left\{ \frac{1}{2} C_1 \mathbf{P}^2 + \frac{1}{4} C_4 (\mathbf{P}^2)^2 \right\}
\]

(4)

describes local orientational ordering due to the active driving following the approach by Toner and Tu \[17\] for neglected convection. The functional possesses two coupling parameters \(C_1\) and \(C_4\) which govern the local orientational ordering due to the drive. If \(C_1 = C_4 = 0\), only gradients in the density \(\psi_1\) can induce local polar order \(\mathbf{P}\) of the active driving. For \(C_1 > 0\) (\(C_4 = 0\)) diffusion tends to reduce the polar order generated by the density gradients. In the third case, \(C_1 < 0\) and \(C_4 > 0\), a net local driving spontaneously emerges already in the absence of density gradients.

Clearly, on the one hand, for vanishing self-propulsion \(v_0 = 0\), Eqs. (1) and (2) decouple and the density equation reduces to the usual phase field crystal model \[16, 20\]. On the other hand, if \(\mathcal{F}_{pfc}\) is neglected, the remaining terms are contained in the model by Toner et al. \[17, 33\], except for the higher-order term in \(\mathbf{P}\) that contributes to translational diffusion. Summarizing, Eqs. (1)–(4) form a minimal approach to characterize crystallization in actively driven systems.

We numerically determined the phase diagram by scanning the \(\psi\)-\(\varepsilon\) plane while keeping the parameters \(C_1\), \(C_4\), and \(v_0\) fixed. As for any numerical result reported subsequently, we proceeded in the following way. For each set of parameter values \((\psi, \varepsilon, C_1, C_4, v_0)\) we started from random initial conditions and then iterated Eqs. (1)–(4) forward in time. Numerical measurements were carried out after equilibration, and a systematic finite size study was performed to test the validity of our results.

For the decoupled case \(v_0 = 0\), the equilibrium phase diagram \[20\] corresponding to the energy functional Eq. (3) is shown in Fig. 1(a). For nonzero active drive \(v_0\), we will first report on the case \(C_1 > 0\).

When we moderately increase \(v_0\) from zero for \(C_1 > 0\), the phase boundaries undergo a temperature shift \(\Delta \varepsilon\) to lower temperatures. An example is depicted in Fig. 1(b). Comparison to Fig. 1(a) shows that switching on the active drive melts crystals and lamellae close to the liquid phase boundary. The patterns still remain at rest, however. For this case, a linear stability analysis and derived amplitude equations for \(\psi_1\) and \(\mathbf{P}\) predict \(\Delta \varepsilon \propto v_0^2/C_1\), which was also verified numerically.

We present an example snapshot of the resting crystalline phase in Fig. 2(a). The peaks of the density distribution \(\bar{\psi}\) form a hexagonal lattice as dictated by the PFC energy functional. \(\mathbf{P}\) points down the density gradients. Consequently the polarization field forms “+1”-defects centered at the density peaks. Since \(\mathbf{P}\) describes the local direction of active drive, density is convected out of the peaks by the active propulsion \(v_0\). This mech-
FIG. 2. Snapshots of the order parameter fields for different phases that are observed when increasing the active drive \( v_0 \) at \( (\bar{\psi}, \varepsilon, C_1, C_4) = (-0.4, -0.98, 0.2, 0) \): (a) resting hexagonal, \( v_0 = 0.1 \), (b) traveling hexagonal, \( v_0 = 0.5 \), (c) traveling quadratic, \( v_0 = 1 \), (d) traveling lamellar, \( v_0 = 1.9 \). The phases are depicted by plotting the density field \( \psi \); bright color corresponds to higher densities. Thin bright needles illustrate the polarization field \( P \) that points from the thick to the thin ends. In panels (b)–(d) the predominant direction of motion is indicated by the bright arrows. Only a fraction of the numerical calculation box is shown.

FIG. 3. Sample-averaged magnitude \( v_m \) of the crystal peak velocities (left scale) and polar order parameter \( p_v \) of the crystal peak velocity vectors (right scale) as a function of \( v_0 \) for \( (\bar{\psi}, \varepsilon, C_1, C_4) = (-0.4, -0.98, 0.2, 0) \). The threshold corresponds to the onset of collective crystalline motion. Thick arrows mark the positions where the snapshots of Fig. 2 were taken; the black stars indicate the intersection points with the phase diagrams in Figs. 1(b) and 1(c). The region above threshold where regular swinging motion could be observed is marked in gray. Inset: peak trajectories illustrating a state of regular swinging motion in a hexagonal crystal; different colors correspond to different peaks; only trajectories of a horizontal row of density peaks are shown that started at the bottom and were traveling to the top of the picture while tracking was performed.

This reduced symmetry induces active propulsion: a net orientation of the polarization field emerges when averaged over the area of a single density peak. The consequence is an active convection of each density peak, originating from the PFC density modulation. These results are in agreement with a linear stability analysis of Eqs. (1) and (2) which predicts that propagating modes appear above a threshold value \( v_{0,c} \).

With further increasing \( v_0 \) the hexagonal pattern is deformed to a rhombic one. In the end, we observe a nearly quadratic structure as depicted in Fig. 2(c). This structural hexagonal–rhombic–quadratic transition appears to be smooth and continuous.

Finally, we observe that the traveling crystal can be melted into a traveling lamellar state if \( v_0 \) is increased to still higher values. An example snapshot of such traveling lamellae is shown in Fig. 2(d). In contrast to the hexagonal–rhombic–quadratic distortion of the traveling crystalline lattices, the traveling crystalline–lamellar transition occurs rather abruptly. The transition is also evident when we compare the two phase diagrams in Figs. 1(b) and 1(c). There, with increasing \( v_0 \), the traveling lamellar regions grow into the traveling crystalline regions.

To quantify the scenario further, we tracked the motion of each density peak. We determined the individual peak velocities \( v_i \), where \( i = 1, ..., N_p \), and \( N_p \) denotes the number of peaks. Samples of up to 1000 density peaks were investigated. The sample-averaged peak velocity magnitude follows as \( v_m = \frac{\sum_{i=1}^{N_p} ||v_i||}{N_p} \). In addition, we calculated the degree of polar orientational order of the normalized peak velocities \( p_v = \frac{||(\sum_{i=1}^{N_p} v_i / ||v_i||)||}{N_p} \). This order parameter detects whether the peaks move coherently (collectively) into the same direction.

For \( C_1 > 0 \), Fig. 3 clearly illustrates the existence of a threshold value \( v_{0,c} \) at which propagation starts. As indicated in the inset, we observed a regular swinging motion of the hexagonal crystals close to the threshold. With in-
Such textures were observed for flagellated marine bac-
villus textures evolved for all tested nonzero values of v. The Toner-Tu model [17, 34]. For this scenario we never ob-
0served a finite threshold value of v. Propagating structures evolved for all tested nonzero values of v. Again, a transition from hexagonal to rhombic to quadratic and then to lamellar structures was observed with increasing v. Furthermore, we note that our equations of motion are invariant under the transformation $\psi \rightarrow -\psi$, $\psi_1 \rightarrow -\psi_1$, $\mathbf{P} \rightarrow -\mathbf{P}$. Because of these symmetry rel-
tions, our analysis equally applies for the investigation of active honeycomb textures that follow from $\psi > 0$. Such textures were observed for flagellated marine bac-
teria [10, 11].

In summary, we extended the phase field crystal model [10, 20] to active systems by combining it with the approach of Toner et al. [17, 34]. As a result, the active drive favors the liquid and lamellar states in the PFC phase diagram and induces a wealth of new active crystalline states of hexagonal, honeycomb, rhombic, and quadratic texture. The global motion of all these structured states is either “resting” or “traveling”. The transition from “resting” to “traveling” involves a complex intermediate swinging motion. When prepared from an initially disordered state, traveling crystals emerge through coarse-graining from a multidomain texture.

Our model can be extended from two to three spatial dimensions where more crystalline lattice structures become stable [24] and to binary mixtures of driven and undriven particles promising a rich variety of mixed active crystals. In principle, our predictions are verifiable in experiments on self-propelled particles and in particle-resolved computer simulations at high density [12–14]. Very recently, traveling crystals have in fact been found in such simulations [35]. Since the new traveling crystalline structures show a nontrivial dynamical response, they may serve as a building block for a new class of active matter with unusual rheological, phononic, and possibly also photonic properties.

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FIG. 4. Coarse-graining in a sample of about 1000 traveling density peaks. The sample is in the traveling hexagonal crystalline state. Depicted are instant pieces of all peak trajectories, drawn in different colors. These pieces connect to lines of equal orientation within collectively moving domains. Over time, some domains of equal velocity orientation grow on the cost of others. Parameter values correspond to Fig. 2(b). Dimensionless times [31]: (a) 4500, (b) 50000, (c) 70000.
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