Driven Disordered Periodic Media with an Underlying Structural Phase Transition

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We investigate the driven states of a two-dimensional crystal whose ground state can be tuned through a square-triangular transition. The depinning of such a system from a quenched random background potential occurs via a complex sequence of dynamical states, which include plastic flow states, hexatic, dynamically stabilized triangle and square phases and intermediate regimes of phase coexistence. These results are relevant to transport experiments in the mixed phase of several superconductors which exhibit such structural transitions as well as to driven colloidal systems whose interactions can be tuned via surface modifications.

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The description of phases and phase transitions in driven steady states is a central theme in the statistical mechanics of non-equilibrium systems [1]. A variety of such states are obtained in the depinning and flow of randomly pinned, periodic media, such as charge-density wave systems and Abrikosov flux-line lattices in the mixed state of type-II superconductors [2]. Many such superconductors exhibit structural transitions in the mixed state, typically between flux-line lattices with triangular and rectangular symmetry [3]. Colloidal systems such as PMMA spheres coated with a low-molecular-weight polymer undergo a remarkable variety of solid-solid transformations in an external field [4]. While applying a sufficiently large current depins the flux lines from the quenched random disorder present in all real materials, the possibility of driving colloidal particles in two dimensions across a disordered substrate has also been raised [5]. What links these diverse systems is the generic problem of understanding the competition between an underlying structural phase transition in a pure periodic system as modified by disorder, and the non-equilibrium effects of an external drive. This Letter proposes and studies a simple model which describes this physics.

Our model system is two-dimensional and consists of particles with two and three-body interactions [6]. The three-body interaction, parametrized through a single parameter $v_3$, tunes the system across a square-triangular phase transition. Our central result, the sequence of steady states obtained as a function of increasing force $F$ for various values of $v_3$, is summarized in the dynamical “phase” diagram of Fig. 1. We obtain a variety of phases: pinned states which may have dominantly triangular or square correlations, a moving liquid/glass phase, a moving hexatic glassy phase, flowing triangular and square states ordered over the size of our simulation cell and a dynamic coexistence regime between these ordered phases. We discuss our characterization of these states and the applicability of simple dynamical criteria for non-equilibrium phase transitions between them.

The model: Particles interact in two dimensions through the interaction potential $1/2 \sum_{i\neq j} V_2(r_{ij}) + 1/6 \sum_{i\neq j\neq k} V_3(r_{ij}, r_{jk}, r_{ik})$, where $r_i$ is the position vector of particle $i$, $r_{ij} \equiv |r_j - r_i|$, $V_2(r_{ij}) = v_2 (\frac{\sigma_0}{r_{ij}})^2$ and $V_3(r_i, r_j, r_k) = v_3 [ f_{ij} \sin^2(\theta_{ijk}) + \text{permutations} ]$. The function $f_{ij} = f(r_{ij}) = (r_{ij} - \sigma_0)^2$ for $r_{ij} < 1.8 \sigma_0$ and 0 otherwise and $\theta_{ijk}$ is the angle between $r_{ij}$ and $r_{jk}$. The two-body (three-body) interaction favours a triangular (square) ground state. Energy and length scales are set using $v_2 = 1$ and $\sigma_0 = 1$. Particles also interact with a quenched random background modeled as a Gaussian random potential $V_d(r)$ with zero mean and exponentially decaying (short-range) correlations. The disorder variance is set to $v_2^2 = 1$ and its spatial correlation length is $\xi = 0.12$. The system evolves through standard Langevin dynamics: $\dot{r}_i = v_i$ and $\dot{v}_i = f^{\text{int}}_i - \alpha v_i + F + \eta_i(t)$. Here $v_i$ is the velocity, $f^{\text{int}}_i$ the total interaction force, and $\eta_i(t)$ the random force acting on particle $i$. A constant force $F = \{F_x, 0\}$ drives the system and the zero-mean thermal noise $\eta_i(t)$ is specified by $\langle \eta_i(t) \eta_j(t') \rangle = 2 T \delta_{ij} \delta(t - t')$ with $T = 0.1$, well below the equilibrium melting temperature of the system. The unit of time $\tau = \alpha \sigma_0^2 / v_2$, with $\alpha = 1$ the viscosity.

Simulation details: Our system consists of 1600 particles in a square box at number density $\rho = 1.1$. At $T = 0.1$, the pure system remains triangular upto $v_3 = 1.5$. For larger $v_3$, a square phase is obtained. Larkin length estimates yield $L_d/a \sim 100$, with $a = 1/\rho^{1/2}$ the lattice parameter, somewhat larger than our system size.

Equilibration in the disorder potential is achieved through a simulated annealing procedure in which the disorder potential is varied by increasing its strength from zero to unity in small steps. The system is stabi-
lized for a minimum of $2 \times 10^5$ Monte Carlo steps at each disorder strength. Such annealed configurations are our initial inputs to the Langevin simulations. We evolve the system using a time step of $10^{-4} \tau$. The external force $F$ is ramped up from a starting value of 0, with the system maintained at up to $10^8$ steps at each $F$.

**Observables:** We monitor structural observables, such as the static structure factor $S(q) = \sum_{ij} \exp(-i\mathbf{q} \cdot \mathbf{r}_{ij})$. Delaunay triangulations yield the probability distributions $P(n)$ of $n = 4, 5, 6$ and 7 coordinated particles ($\sum_n P(n) = 1$). We define order parameters $\psi = (P(4) - P(6))/(P(4) + P(6))$ to distinguish between square and triangular phases, $\psi_\Delta = (P(6) - P(5) - P(7))/(P(6) + P(5) + P(7))$ to distinguish between liquid (disordered) and triangular crystals and $\psi_\Theta = (P(4) - P(5) - P(7))/(P(4) + P(5) + P(7))$ to distinguish between liquid and square crystals. In addition, we compute the hexatic order parameter $\psi_6 = \sum_{ij} \exp(-i\theta_{ij})$ and its correlations, where $\theta$ is the bond angle measured with respect to an arbitrary external axis. The dynamical variables we study include the center of mass velocity $v_{cm}$, the particle flux and its statistics and the Koshelev-Vinokur (KV) “shaking temperature” $T_s$; appropriate to the drive and transverse directions and obtained from $T_s = \langle [v_{\alpha} - v_{\alpha_{cm}}]^2 \rangle / 2$, $\alpha = x, y.$

For small $F$ the solid is pinned. A disorder broadened version of the equilibrium triangle (A) to square (B) transition results as $v_3$ is varied across the $T = 0$ transition value at small $F$; here and below alphabets in brackets refer to the states labelled in Fig. 1. The triangular (A) phase is favoured at non-zero $F$. Upon further increasing $F$, the system undergoes a discontinuous depinning transition which exhibits prominent hysteresis behaviour (Fig. 1 inset). Such a depinned state is inhomogeneous and undergoes plastic flow...
consistent with earlier numerical work. For larger $F$ the velocity approaches the asymptotic behaviour $v_{CM} = F$.

The structure factor $S(q)$ of the plastically moving phase (C) obtained just above the depinning transition consists of liquid-like isotropic rings (Fig. 1(i)). Upon increasing $F$, the circular ring in $S(q)$ concentrates into six smeared peaks which we associate with a hexatic glass (D) [14]. Fig. 2(b) shows the evolution of the hexatic correlation function $g_{6}(r) = \langle \psi_{0}(0)\psi_{0}(r) \rangle$, as $F$ is varied across (C) $\rightarrow$ (D) $\rightarrow$ (E). Note the sharp exponential decay of hexatic correlations in (C), the power-law or quasi-long-range order (QLRO) decay in (D) and the saturation (LRO) of this correlation function in regime (E). The plastic flow regime (C) expands at larger $v_{3}$ [17]. On further increasing $F$, the hexatic glass recrystallizes into a structure which depends on the value of $v_{3}$: for low $v_{3}$ the final crystal is triangular (E) whereas for large $v_{3}$ it is square (F). For intermediate $v_{3}$ the system first freezes into a triangular structure but subsequently transforms into the square via an intervening “coexistence” regime (G) best described as a mosaic of dynamically fluctuating square and triangular regions.

The phase diagram of the pure system in thermal equilibrium accommodates fluid, triangular solid and square solid phases [13]. In the driven system, as $F$ is increased, analogous phases appear in approximately the inverse order to the sequence obtained in the pure case as $T$ is increased. This observation agrees roughly with the KV proposal [13], which identifies the shaking temperature $T_{s}$ with $T$. The shaking temperatures are predicted to fall as $\sim 1/v^{2}$ and as $\sim 1/v$ in the drive and the transverse directions respectively, consistent with our observations in Fig. 2(a). We find that $T_{s}^{\alpha}$ is nearly independent of $v_{3}$. Importantly, within the putative coexistence regime, $T_{s}^{\alpha}$ behaves non-monotonically, implying a breakdown of the KV prediction (see Fig. 2(a)). Typically, for a particular disorder configuration and for $5.5 < v_{3} < 8.5$, $T_{s}^{\alpha}$ appears to increase sharply at a well defined $F$, signifying the start of coexistence. Within G, $T_{s}^{\alpha}$ remains high but drops sharply at the upper limit of G, to continue to follow the interrupted KV behavior. The limits of the coexistence region, though sharp for any typical disorder realization, vary considerably between realizations.

Within the coexistence region the probability of obtaining triangular (square) regions appears to decrease (increase) roughly linearly with $F$; see Fig. 2(c). Real space configurations (Fig. 3) exhibit islands of square and triangular coordination connected by interfacial regions with predominately 5 coordinated particles. Particles with coordination 7 are typically associated with dislocations which are scattered randomly in the interface. This configuration, in the co-moving frame, is extremely dynamic, with the islands rapidly interconverting between square and triangle. This interconversion has complex temporal properties: the power spectrum of coordination number fluctuations shows a prominent $1/f$ fall off over several decades. In addition, particle current fluctuations are enhanced by an order of magnitude, also displaying a regime of $1/f$ behavior (Fig. 4), although over a restricted range as a consequence of the proximity to the wall frequency. This result would be hard to interpret in the absence of a genuine coexistence phase: above the depinning transition, increasing the driving force would naively be expected to reduce current noise monotonically, as observed in all previous simulation work on related models [12, 13, 14].

Renormalization group arguments suggest that neither translational LRO nor QLRO survive in the disordered moving state in two dimensions at any finite drive [15, 19, 20], their closest analog being the moving Bragg glass state argued to be stable in three dimensions and higher [20]. The ordered square (F) and triangular (E) states we obtain at large drives are then to be understood as at a finite size effect arising from the restricted size of our simulation box, although the crossover length scales can be very large at weak disorder [19, 20]. The possibility of alternative dynamically stabilized states with reduced levels of ordering, such as driven transverse smectics, is attractive [15, 21]. In contrast to some previous work [11, 13, 14], we see no evidence for smectic order and flow in weakly coupled channels at large drives – our channels always remain strongly coupled – but note that moving states in which channels transverse to the drive direction are effectively decoupled may be stabilized at higher levels of thermal noise or randomness [14].

No general arguments seem to rule out QLRO in the “moving hexatic glass” phase in two dimensions and our simulations support this possibility; see also Ref. [10].
The drive-induced stabilization of the triangular lattice state we obtain, even well into regimes where $v_3$ would favour a square, is an unusual result. Our finding of a distinct coexistence regime (G) separate from plastic flow states, hexatic and crystal, is novel. Such coexistence occurs in a narrow and reproducible regime in parameter space and is characterized by a variety of dynamical anomalies, including enhanced noise signals with $1/f$ character. The coexistence state appears to be a genuine non-equilibrium state, separated from other regimes through sharp non-equilibrium transitions.

Theoretical work has, so far, neglected the possibility of such dynamic phase coexistence in the non-equilibrium steady states of driven disordered crystals [21]. A non-disordered but frustrated system closely related to the one considered here has been proposed recently as a model for the dynamical heterogeneity seen in the glassy state [22]. In this model, fluctuating regions of crystalline ordering within a liquid background are argued to be responsible for the anomalous dynamical behaviour and slow relaxation in the glassy state, a physical picture which shares some similarities to our ideas regarding the coexistence regime.

In conclusion, we have shown here that the competition between structural phase transitions in a pure system as modified by disorder, coupled to the non-equilibrium effects of an external drive, can have a variety of non-trivial consequences. The ubiquity of structural phase transitions in the vortex state of a large number of superconductors which have been studied recently, as well as the relative ease with which the vortex state can be driven, suggests experimental situations in which the ideas here should find application. Functionalized colloidal particles driven over random substrates constitute a novel system on which our proposals can be tested [18, 23]. Such systems have the further advantage that interparticle interactions can be tuned both through surface modifications and through the application of external fields [22].

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