## **Spontaneous Patterning of Confined Granular Rods**

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Vertically vibrated rod-shaped granular materials confined to quasi-2D containers self-organize into distinct patterns. We find, consistent with theory and simulation, a density dependent isotropic-nematic transition. Along the walls, rods interact sterically to form a wetting layer. For high rod densities, complex patterns emerge as a result of competition between bulk and boundary alignment. A continuum elastic energy accounting for nematic distortion and local wall anchoring reproduces the structures seen experimentally.

DOI: 10.1103/PhysRevLett.96.028002

From nanometer-sized molecules forming liquid crystals to timber floating on a river, rod-shaped materials pervade our everyday existence. Onsager first demonstrated that hard-core steric interactions suffice to order rods in thermal equilibrium [1]. Above a critical concentration, orientational entropy is sacrificed for gains in translational entropy; therefore, crowding alone induces a temperature-independent isotropic-nematic (I-N) transition [2]. In this entropically driven transition, the kinetic motion of the rods serves as a mechanism for adequately sampling phase space. While thermal excitation mixes microscopic particles, macroscopic rods require an externally applied energy for randomization. But collisions between granular particles, in contrast to their classical counterparts, are inelastic. Excited granular materials, therefore, readily violate equipartition of energy and exhibit non-Maxwellian velocity distributions [3]. Such driven dissipative systems are far from equilibrium and need not evolve to states of maximal configurational entropy. For example, prior studies utilizing macroscopic rods focused on jammed or other metastable states [4,5]. Nevertheless, some aspects of granular rod behavior resemble their microscopic counterpart. A density dependent nematic to smecticlike transition was previously observed for granular rods, for both vertical and horizontal vibrations [5]. However, the relation of ordering seen in granular rods to that in liquid crystals remained unknown.

Here we report on finite-sized driven dissipative granular systems of rods in steady state that share many properties with lyotropic liquid crystals at equilibrium. By using excluded volume scaling predicted by mean field theories, our data give a single normalized transition density for all rod aspect (length-to-diameter, L/D) ratios and densities tested. However, excluded volume interactions in the bulk cannot explain all rod behavior because confining boundaries also strongly influence rod ordering. For relatively low rod densities, experimental results for rod alignment and density profiles with respect to the wall strikingly resemble excluded volume-based theoretical predictions [6]. As the rods become dense, rod ordering in the bulk nematic competes with rod ordering along the surrounding walls. This

frustration creates distinct yet easily reproduced patterns. By modeling the system as a liquid crystal undergoing the elastic deformations of splay and bend subject to a simple wall interaction [7–9], we show that the patterns depend on the relative magnitude of wall and bulk energies.

PACS numbers: 45.70.Qj, 61.30.Eb, 61.30.Hn

Rod-shaped granular materials confined to quasi-2D containers were vertically vibrated. Stainless steel wire, 0.08 cm diameter, was cut into rods of L/D=20, 40, and 60. Containers were either circular (radius R=15 cm or R=7 cm) or square (29 cm diagonal). The chamber height to rod diameter ratio was 20, which restricted the rods to lie mainly flat. Due to the quasi-2D nature of these experiments, rod densities were scaled to the area of the container bottom, A, so that the area fraction is  $\phi=NLD/A$ , where N is the number of rods. A monolayer of rods lying flat and fully covering the container bottom corresponds to  $\phi=1$ .

The rods were shaken with a sinusoidal acceleration of 50 Hz and a peak acceleration of 4 times gravity. The magnetic field of the shaker was minimized by distancing the rod chamber from the shaker. Electrostatic forces were reduced by using aluminum chamber components and applying an antistatic coating to nonmetallic surfaces. The apparatus was carefully leveled. To randomize initial conditions, rods were dropped into the containers from above before experiments commenced. Images of the rods in motion were captured by a high speed digital camera. The position and orientation of each rod were found by an image processing algorithm, developed in MATLAB 6.5. The counting accuracy was close to 100% for low density systems and greater than 75% for samples as dense as  $\phi = 0.6$ .

Unlike the infinite bulk modeled by Onsager's mean field theory [1], rigid boundaries confine our experimental system. We first investigate the I-N transition away from these strong wall influences. By measuring the angular correlation of the rods with respect to the nearest wall [wall-rod correlation,  $g_2(r)$ ], we distinguish rods affected by the side boundaries from those that were not. We define an effective "bulk" when  $g_2(r) \approx 0$ , which occurs at distances r/L > 0.5 from the wall. We will later discuss

in detail  $g_2(r)$  and other parameters in relation to ordering along the boundaries.

To measure rod ordering in the bulk, we characterize alignment via the order parameter  $S = \langle \cos(2\theta_i) \rangle$ , where  $\theta_i$ is the angle of the ith rod with the nematic director. In practice, the order parameter and director are found for an arbitrary Cartesian coordinate system via the matrix  $Q_{\alpha\beta}=$  $\langle 2u_{\alpha}(i)u_{\beta}(i) - \delta_{\alpha\beta} \rangle$ , where  $u_{\alpha}(i)$  and  $u_{\beta}(i)$  are the coordinates of the unit vector specifying the direction of the ith rod. The eigenvalues of Q give  $\pm S$ , and the eigenvectors give the director and its perpendicular vector. A perfect nematic corresponds to S = 1. Standard convention denotes an isotropic system as S = 0, but, by the way S is defined, an isotropic system will have S > 0 that approaches 0 only as  $N \to \infty$  [10]. To distinguish rod ordering from this finite-number effect, we evaluated S of finite numbers of noninteracting rods using 2D Monte Carlo (MC) simulations, where the number of rods in the simulations matched those in experiments.

We first tested the dependence of orientational order on the area fraction,  $\phi$ . Typical experimental results are illustrated in Fig. 1, where L/D=40 in the R=7 cm container. In the top of Fig. 1, the rods in the bulk (highlighted) appear to align as  $\phi$  increases. The bottom of Fig. 1 shows S as a function of  $\phi$  as well as S from the MC simulations. For small  $\phi$ , S is relatively low and coincides with S from simulations, indicating that the experimental system is isotropic. As  $\phi$  increases, S rises and deviates from the values derived from simulations, denoting the transition to an ordered nematic state. Conforming to the convention

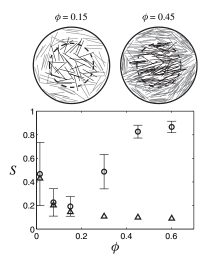


FIG. 1. Dependence of bulk orientational order on rod density for a single rod aspect ratio L/D=40 in the R=7 cm container. Top: Representative images for area fractions  $\phi=0.15$  and 0.45. Dashed lines mark where  $g_2(r)\approx 0$ , and rods with midpoints within these dashed lines were labeled "bulk." Bottom: Order parameter S as a function of  $\phi$  for averaged experimental data  $(\bigcirc)$  with standard deviations. Monte Carlo simulations  $(\triangle)$  of noninteracting isotropically oriented rods are included to distinguish rod ordering from the finite-number effect on S.

that S=0 in an infinite isotropic system, we henceforth subtract the results of the MC simulations from experimental data.

We then fixed  $\phi$  and tested the dependence of orientational order on the rod aspect ratio, L/D. Experimental results for L/D=20 and 60 at  $\phi=0.6$  in the R=15 cm container are shown in Fig. 2 (top). Though the same mass is present in both containers, L/D=60 rods appear more ordered than L/D=20 rods for this value of  $\phi$ . The inset in the bottom of Fig. 2 shows S vs  $\phi$  for all L/D and container sizes and shapes. As L/D increases, the transition to an ordered state shifts to lower  $\phi$ .

By plotting S versus the scaled area fraction  $\rho = \phi L/D$ , the transition curves for different L/D collapse to a single curve [Fig. 2 (bottom)]. This scaling is consistent with Onsager's mean field result, which assumes that the I-N transition stems only from steric interactions between neighboring rods [1]. Within experimental error, the transition appears continuous. The dashed line is a linear least square fitting of the data in the transition region, and, by extrapolating this line to S = 0 [10], we estimate that the critical transition occurs at  $\rho = \rho^*$  between 6 and 9. For comparison, Onsager's mean field model applied to an infinite 2D system of rods predicts a second order I-N transition at  $\rho^* = 3\pi/2 \approx 4.71$  [11], and 2D computer simulations with periodic boundary conditions estimate  $\rho^*$  between 6.5 and 8 [10]. We performed a similar analysis on our data that included both bulk and boundary layer

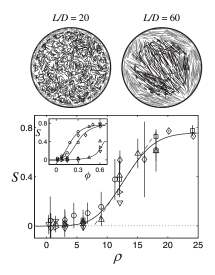


FIG. 2. Dependence of bulk orientational order on rod aspect ratio. Top: Representative images for L/D=20 and 60 at  $\phi=0.3$  in the R=15 cm container. As in Fig. 1, rods within the dashed lines delineate bulk. Bottom: Consolidated plot of S vs scaled area fraction  $\rho=\phi L/D$ , for R=7 cm container: L/D=20 ( $\triangle$ ), 40 ( $\square$ ); R=15 cm container: L/D=20 ( $\triangleright$ ), 60 ( $\bigcirc$ ); square container: L/D=20 ( $\nabla$ ), 40 ( $\bigcirc$ ). The dashed line is a linear least squares fitting of the data in the transition region. By extrapolating this line to S=0, we estimate the transition density between 6 and 9. Inset: Same data plotted as S vs  $\phi$ . All solid lines are guides for the eye.

rods. The measured S for the nematic state is systematically lower (due to cancellation by the alignment of the rods along orthogonal walls), but, within experimental error, the location of  $\rho^*$  remains unchanged.

While the side boundaries do not affect  $\rho^*$ , they profoundly influence rod ordering in other ways. Rods near a wall tend to align parallel along it. We assessed this alignment using the wall-rod angular correlation function  $g_2(r) = \langle \cos[2(\vartheta_w - \vartheta(r))] \rangle$ , where  $\vartheta(r)$  is the angle of a rod located at distance r away from the wall, as described by the shortest line from rod midpoint to wall, and  $\vartheta_w$  denotes the vector locally tangent to the wall.

The left panel in Fig. 3 shows  $g_2(r)$  for L/D = 20 in the R=7 cm container at  $\rho=1.5$  (circles) and  $\rho=9$  (triangles). Experiments show an initially high  $g_2(r)$  that decays as r increases. For comparison, the dashed line represents the MC calculated  $g_2(r)$  for a single rod interacting only via hard-core repulsions with the curved wall. Steric interactions with the wall significantly restrict the possible orientations available for a rod closer than L/2 to the wall. Experimental values of  $g_2(r)$  closely follow the single rod curve for small r; however, these values deviate as r increases. Furthermore, as  $\rho$  increases,  $g_2(r)$  diverges earlier and remains elevated for larger r. We also measured the number density  $\chi(r)$ , normalized with respect to bulk density, of rod midpoints (Fig. 3, right). The measured  $\chi(r)$  is less than 1 for r/L < 0.5, which signifies a depletion of rod midpoints next to the wall. Finally,  $\chi(r)$  obtains a local maximum around  $r/L \approx 0.6$  before returning to bulk values.

Onsager's approximation for excluded volume interactions has been applied to hard spherocylinders in contact with a hard flat wall by Poniewierski [6], and results have been confirmed by simulation [12]. Although this analysis was performed for a 3D system contacting a planar wall, the experimental features in Fig. 3 show striking similarities. The profiles for  $g_2(r)$  and  $\chi(r)$ , however, can be difficult to interpret since geometrical constraints tightly

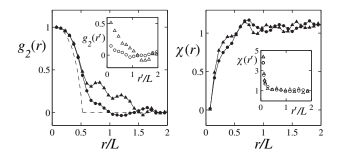


FIG. 3. Wall-rod correlation function  $g_2(r)$  and normalized density profile  $\chi(r)$  versus distance r from wall to rod midpoints for L/D=20 in the R=7 cm container at  $\rho=1.5$  (circles) and  $\rho=9$  (triangles). Data were smoothed by a moving average filter with a span of 3. The dashed line represents calculated  $g_2(r)$  for a single rod interacting with the wall via hard-core repulsions. The insets show  $g_2(r')$  and  $\chi(r')$  versus distance r' from wall to rod ends.

couple rod orientation and position for rod midpoints close to the wall. By analyzing distances r' based on the closest end of each rod to a wall, rod orientation and position are decoupled, thereby simplifying  $g_2(r')$  and  $\chi(r')$  profiles [6]. Figure 3 (left inset) shows a clear  $\rho$  dependent increase in  $g_2(r')$  for rod ends near the wall. And, in contrast to the depletion layer seen for rod midpoints, a strong wetting layer becomes apparent, with an approximate fourfold increase in  $\chi(r')$  for small r' which decays rapidly to bulk values around  $r'/L \approx 0.3$  (Fig. 3, right inset).

Although many similarities exist between the hard-core interactions of rods with straight versus curved boundaries, wall geometry significantly affects  $g_2(r)$ . Rod alignment with the wall does not penetrate as deeply into the sample center for curved boundaries, and, interestingly, the thickness of this layer even regresses at very high  $\rho$  (data not shown). This seemingly unusual behavior can be understood by noting that for  $\rho \gg \rho^*$  the system must accommodate both the orientation of the nematic director as well as the preferred orientation parallel to the surrounding boundaries. This frustration creates competition between the rods in the bulk with those at the boundary, which produces complex patterns, Fig. 4 (left).

Even though the local variations in the experimentally observed patterns are often on the order of rod dimensions,

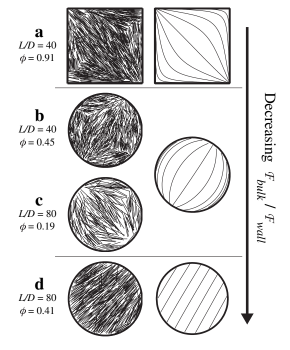


FIG. 4. Elastic deformations of the nematic observed for  $\rho$  above the I-N transition. Experimental images in square and R=15 cm containers (left) and results from numeric minimization of the elastic free energy functional  $\mathcal{F}$  (right) are arranged with respect to relative bulk and wall free energies  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}}$ , with (a)  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}} \gg 1$ , (b),(c)  $\approx 1$ , and (d)  $\ll 1$ . This ratio is modified experimentally by changing L/D or  $\phi$  (see text). A transition can be made from the patterns seen in (b) and (c) to those in (d) by either increasing (b) L/D or (c)  $\phi$ .

we find that these patterns are well approximated by minimizing a continuum free energy functional that describes elastic deformations in ordered liquid crystals [7–9]. Similar methods have been used to study nematic droplets [13]. Consider a total free energy composed of bulk and wall contributions,  $\mathcal{F} = \mathcal{F}_{\text{bulk}} + \mathcal{F}_{\text{wall}}$ . In the bulk, splay and bend deformations contribute to the distortion energy:

$$\mathcal{F}_{\text{bulk}} = \frac{1}{2} \int_{A} K_1(\nabla \cdot \mathbf{n})^2 + K_3(\nabla \times \mathbf{n})^2 dx dy, \qquad (1)$$

where  $\mathbf{n}(x, y) = [\cos \vartheta, \sin \vartheta]$  is the unit vector describing the local nematic director, and  $K_1$ ,  $K_3$  are constants for splay and bend, respectively. Note that contributions from twist deformations are absent in this 2D system. The elastic constants have been linked theoretically to the molecular details of the nematic, generally becoming larger with increasing L/D or  $\phi$  [9]. Often, however, the two constants are comparable in magnitude, and here we shall use the simplifying approximation  $K_1 = K_3 = K$  [8]. To account for interactions favoring rod orientations locally parallel to the wall, we consider the line integral over the container perimeter P:

$$\mathcal{F}_{\text{wall}} = \frac{1}{2}C \int_{P} \sin^{2}(\vartheta_{w} - \vartheta) ds, \tag{2}$$

where C is a positive constant with units of line tension, and  $\vartheta_w$  describes the vector locally tangent to the wall,  $\mathbf{n}_w$ . From Eq. (2), it becomes clear that  $\mathcal{F}_{\text{wall}}$  depends not only on anchoring strength C but also on wall curvature and circumferential length.

Figure 4 (right) shows the results of numeric minimization of the elastic free energy functional  $\mathcal{F}$  using a relaxation method with simulated annealing [14], where the local tangent to each curve represents the orientation of  $\mathbf{n}$  at that point. Similar patterns found in experiment [Fig. 4 (left)] and calculations are grouped and arranged according to the relative ratio of the minimized  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}}$ . Notice that as nematic "stiffness" increases compared to wall anchoring (increasing K/C),  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}}$  decreases, because the system reduces the cost of nematic deformations in the bulk at the expense of misalignment with the wall.

Experimentally, by holding wall geometry constant and varying  $\phi$  and L/D, we changed nematic stiffness K and the ratio K/C, thus modifying  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}}$  [8]. Differing combinations of  $\phi$  and L/D give rise to equivalent patterns, as seen in Figs. 4(b) and 4(c) corresponding to  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}} \approx 1$ . By raising nematic stiffness though increasing L/D [Fig. 4(b)] or  $\phi$  [Fig. 4(c)], we transition to the pattern seen in Fig. 4(d), where  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}} \ll 1$ . At the other extreme, where  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}} \gg 1$ , alignment with the wall is favored at the expense of nematic deformations. By changing wall curvature, we can strengthen wall anchoring and the penalty for wall misalignment, as seen in Fig. 4(a). For a wide range of parameters, we not

only find that the experimental patterns match the theoretical model results but also that  $\mathcal{F}_{\text{bulk}}/\mathcal{F}_{\text{wall}}$  is sufficient to describe the whole range of nematic deformations witnessed in experiments. However, as may be expected for confined nematics [15], line and point defects are visible.

In extending the normal application of liquid crystal continuum theories by using them to describe nonequilibrium, strongly confined systems of rods, we show that the macroscopic granular rods in this steady state system share many properties with thermally equilibrated lyotropic liquid crystals. The benefit of using a granular system, as opposed to one composed of microscopic rods, is that interactions can be purely hard-core, and individual rods are easily visualized. This system may be useful for studying defects and other aspects of lyotropic liquid crystals that might be difficult to probe experimentally.

We thank D. Lathrop for use of the shaker, J. V. Sullivan for designing experimental components, and W. M. Gelbart, M. Schröter, and V. Frolov for helpful comments. This research was supported by the Intramural Research Program of the NIH (NICHD).

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