Emergent Hyperuniformity in Periodically Driven Emulsions

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We report the self-organization of microfluidic emulsions into anomalously homogeneous structures. Upon periodic driving confined emulsions undergo a first-order transition from a reversible to an irreversible dynamics. We evidence that this dynamical transition is accompanied by structural changes at all scales yielding macroscopic yet finite hyperuniform structures. Numerical simulations are performed to single out the very ingredients responsible for the suppression of density fluctuations. We show that, as opposed to equilibrium systems, the long-range nature of the hydrodynamic interactions are not required for the formation of hyperuniform patterns, thereby suggesting a robust relation between reversibility and hyperuniformity which should hold in a broad class of periodically driven materials.

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What is the most effective way to homogeneously fill space with an ensemble of particles? At thermal equilibrium, an obvious effective strategy would be to endow the particles with interactions promoting the formation of a crystal. In a $d$-dimensional system, thermal fluctuations would spontaneously organize the particles into an ordered state where the average position of the particles is a perfect lattice. The number fluctuations of these averaged positions would scale as $\Delta N^2 \sim \epsilon^{d-1}$ in boxes of size $\epsilon$; crystals are hyperuniform [1]. At large scales, they are much more homogeneous than a random set of points with number fluctuations of the order of the box volume $\epsilon^d$. However, perfect crystals are not the only patterns being hyperuniform [1,2]. Over the last decade much attention has been devoted to disordered structures displaying miniature density fluctuations. As it turns out, such hyperuniform patterns have been shown to display outstanding optical properties such as complete photonics band gaps [3–5]. Until very recently the only two controlled strategies to engineer hyperuniform materials were based on numerical optimization techniques [3,5,6], or the jamming of athermal hard spheres [7–11]. In 2015, two sets of numerical simulations have demonstrated that hyperuniformity emerges when ensembles of particles driven out of equilibrium approach a critical absorbing phase transition [12,13]. However, these numerical models based on elegant toy models lack a truly analogous physical system, in which hyperuniformity emerges from genuine physical interactions.

In this Letter we demonstrate for the first time an experimental system in which self-organization into hyperuniform patterns occurs away from the jamming point. The system consists of a periodically driven emulsion, in which maximal hyperuniformity is reached at the onset of reversibility of the droplet dynamics, even though the emulsion does not reach a genuine absorbing state. We also identify the minimal ingredients required to produce hyperuniform emulsions by means of numerical simulations. Surprisingly, we show that long-range interactions impair the emergence of hyperuniform structures. We therefore conjecture that hyperuniformity is intimately related to reversibility in periodically driven systems. We believe that this work opens new possibilities to control the self-organization of a broad class of systems (vortices in superconductors, soft glasses, colloidal suspensions, …) into hyperuniform structures and provides potential new avenues to high-yield and controllable design of isotropic band gap materials [3–5].

The experimental setup is the one used in Ref. [14] consisting of flowing a monodisperse emulsion (area fraction: 0.36) in a microfluidic channel (0.5 cm × 5 cm × 27 ± 0.1 μm). The droplets have a diameter $a = 25.5 \pm 0.5 \mu m$ comparable to the height of the channel and therefore undergo two-dimensional motion. The inlet of the channel is connected to a syringe pump that drives the suspension sinusoidally. The emulsion is prepared in a reproducible initial state reached after a sequence of 10 high-amplitude oscillations. Then, a sequence of $10^3$ cycles at the desired amplitude is applied to ensure that the measurements are performed in a statistically stationary state. The mean displacement of the droplets occurs along the main flow direction and is sinusoidal, and its amplitude $\Delta$ is the sole control parameter of the experiments. The data are collected by tracking the instantaneous position of $\sim 3 \times 10^3$ droplets remaining in the field of view throughout the entire flow cycle. Two snapshots of the emulsion are shown in Figs. 1(a) and 1(b) and correspond to $\Delta = 18.3a$ and $\Delta = 43.3a$, respectively.

Following Ref. [15] the macroscopic reversibility of the system is measured by determining the fraction of active...
particles $f_a$, which is the fraction of droplets that behave irreversibly. A droplet is here defined to be active if it does not return at the end of a cycle within the spatial extent of the Voronoi cell it occupied at the start of the cycle. In these microfluidic experiments, the viscous flows are reversible in time [14]. However, above a driving amplitude $\Delta^*/a = 28.1 \pm 0.3$ the dynamics abruptly becomes irreversible. A macroscopic fraction of the droplets remains endlessly active upon periodic driving as illustrated in Fig. 1, and quantified in Fig. 2(a), where $f_a$ is plotted as a function of driving amplitude.

As $f_a$ is constructed from a metric-free criteria, it is both affected by the changes in the dynamics, and in the structure of the emulsion. We now disentangle and elucidate these two concomitant collective phenomena. Let us begin with the dynamical arrest of the strobed dynamics. The reversible states where droplets retrace their steps back to their initial Voronoi cell, $\Delta < \Delta^*$, do not correspond to interaction-free conformations: in the course of the cycles, even the passive particles continuously interact with all the other droplets via hydrodynamic interactions. They also a priori experience a number of weak but irreversible perturbations such as short-range potential interactions, or minute shape deformations which cannot be experimentally measured. As a result, even in the reversible regime, they return only on average to their initial position after a cycle. The transition to an irreversible state where all the particles are active is associated with a discontinuous amplification of the mean-square displacement of the strobed dynamics [16], thereby causing the escape of the droplets from their initial Voronoi cell. We characterize the fluctuations of the strobed dynamics by the distribution of the modulus of the displacements in the flow direction at the end of each cycle in Fig. 2(b). At the onset of irreversibility this distribution is not merely widened, the statistics of the droplet displacements undergoes a sharp qualitative change at $\Delta^*$ thereby confirming the collective nature of the dynamical arrest. The strobed-displacement statistics is Gaussian for high $\Delta$. Unexpectedly, below $\Delta^*$ the displacement statistics converges to a non-Gaussian universal distribution that is much broader. The decay of the distribution is consistent with an exponential tail which captures the existence of large-amplitude jumps typically corresponding to droplets exiting their Voronoi cell. These intermittent displacements are akin to the cage jumps found in glass forming liquids [17]. Importantly, these results imply that the dynamical arrest of the strobed dynamics does not belong to the absorbing-phase-transition scenario reported in Refs. [12,13,15,18].

FIG. 1 (color online). Close-up of the emulsion at the beginning of the 500th cycle. The active droplets are shown as open orange circles, the passive droplets as filled blue circles. (a) For a driving amplitude $\Delta/\Delta^* = 0.65$, the dynamics is reversible. (b) For a driving amplitude $\Delta/\Delta^* = 1.54$, the dynamics is not reversible anymore. Note also the markedly different structure in the two cases: for low amplitudes the structure is more homogeneous.

FIG. 2 (color online). (a) Fraction of active particles $f_a$ in the steady state at various oscillation amplitudes. At $\Delta = \Delta^*$ reversibility abruptly breaks down. Inset: Variations of the mean-squared strobed displacement in the flow direction $\langle \delta x^2 \rangle$ plotted as a function of $\Delta$. (b) Centered and normalized probability density distribution of the strobed displacements of the droplets, $|\delta x|$, in the flow direction for the experiments (top curves) and numerics (bottom curves). The numerical curves have been shifted for the sake of clarity. The color indicates the mean fraction of active particles. See color bar in (a). (c) Structure factor $S(k)$ at various amplitudes [Experiments: thin lines, simulations: Thick lines, Same color code as in (b)]. For sake of clarity the curves corresponding to $\Delta < \Delta^*$ have been shifted down by a constant value. The dashed line is a guide and corresponds to $k^{0.45}$. 

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We now focus on the central results of this Letter and demonstrate the emergence of hyperuniform structures. Having a quick look back at Fig. 1, we see that the configuration in the reversible regime looks much more homogeneous than in the irreversible regime. In order to quantify this apparent structural change, we compute the structure factor $S(k)$ of the emulsion at various amplitudes, Fig. 2(c). The existence of a structural transition at the onset of reversibility is very noticeable from the sudden change of $S(k)$ across all scales. The shift of the high-$k$ peak of the structure factor corresponds to the change of the liquidlike structure reported in Ref. [14]. However, the most striking feature at the transition occurs at low $k$. For high driving amplitudes (upper curves), $S(k)$ decays algebraically as $k$ goes to 0, indicating the absence of long-range order. In contrast, at low amplitudes (lower curves) $S(k)$ decays algebraically as $k$ goes to 0, large-scale density fluctuations are suppressed [1]. The emulsion seems to self-organize into a hyperuniform state.

In order to quantify the degree of hyperuniformity and the extent of the hyperuniform regions, we go back to real space and directly measure the statistics of the droplet number $N_\ell$ in a $\ell \times \ell$ box [1,12]. Figure 3(a) shows the variations of the variance $\Delta N_\ell^2 \equiv \langle N_\ell^2 \rangle - \langle N_\ell \rangle^2$ normalized by $\langle N_\ell \rangle$ as a function of $\ell$. Any ensemble of particles with no spatial correlation whatsoever has a variance that scales as $\Delta N_\ell^2 \sim \langle N_\ell \rangle$. Therefore, $\Delta N_\ell^2 / \langle N_\ell \rangle$ is a decreasing function of $\ell$ for hyperuniform systems. In Fig. 3(a) we quantify the level of hyperuniformity of the emulsions by normalizing the variance $\Delta N_\ell^2$ by that of a random set of points of the same size and density. This procedure is used to minimize any statistical artifacts due to the finite sample size. For sake of clarity we have shifted the curves starting from small driving amplitudes at the top. For large drivings ($\Delta \gg \Delta^*$), $\Delta N_\ell^2 / \langle N_\ell \rangle$ does not show any significant variations. The density fluctuations are normal. Conversely, as the dynamics becomes reversible ($\Delta < \Delta^*$), the emulsion becomes locally hyperuniform. $\Delta N_\ell^2 / \langle N_\ell \rangle$ indeed first decays algebraically with $\ell$ up to a box size $\ell_{HU}$ above which it increases. We expect the variations to then saturate for high values of $\ell$ that are not accessible with our experimental setup. This nonmonotonic behavior is very similar to that reported first in Ref. [12], at the onset of an absorbing phase transition.

More quantitatively, the system homogeneity is quantified by the exponent $\lambda > 0$ defined as $\Delta N_\ell^2 / \langle N_\ell \rangle \propto \ell^{-\lambda}$. For a random set of points $\lambda = 0$, while $\lambda = 1$ for perfect crystals. In Fig. 3(b) the exponent $\lambda$ fitted for small $\ell$ is shown as a function of the driving amplitude $\Delta$. Interestingly, the closer the system to the reversible transition, the more the density fluctuations are suppressed. At $\Delta = \Delta^*$, we find that $\lambda \sim 0.5$, which is again close to the one reported for systems close to an absorbing phase transition ($\lambda \sim 0.45$) in Ref. [12] and in Ref. [13] at intermediate scales.

The extent of the hyperuniform regions is characterized by measuring the length $\ell_{HU}$, where $\Delta N_\ell^2 / \langle N_\ell \rangle$ deviates from a decreasing power law, Fig. 3(c) [16]. This length scale undergoes nonmonotonic variations with the driving amplitude. When increasing $\Delta$, $\ell_{HU}$ decays from ten to five droplet diameters around $\Delta / \Delta^* \sim 0.7$. Then approaching the reversibility transition $\ell_{HU}$ increases again to its maximal value ($\sim 10a$) before dropping down to zero above $\Delta^*$. Two comments are in order. First, we do not see any sign of a divergence of $\ell_{HU}$ as $\Delta$ approaches $\Delta^*$, which means that the emulsion never self-organizes into a fully

![Figure 3](color online). (a), (b), and (c) Normalized density fluctuations for experiments (a), simulations with long-range interactions (b), and short-range interactions (c). The curves are offset for the sake of clarity. At high amplitudes (red curves) the density fluctuations are normal, whereas in the reversible regime, at low amplitudes (blue curves), the density fluctuations are suppressed. (d) Fitted power-law exponents $\lambda$ from the curves plotted in (a), (b), and (c). The lines are guides to the eye. Density fluctuations are only suppressed for $\Delta \lesssim \Delta^*$. (e) Extent of the hyperuniform regions $\ell_{HU}$. Filled symbols: experiments, open symbols: simulations. The lines are guides to the eye. The system is only hyperuniform up to length scales of $\sim 10a$ ($\sim 15a$) in the experiments (in the simulations).
hyperuniform state where the density fluctuations would be suppressed at the entire system scale. However, the typical extent of the hyperuniform regions are much larger than the typical distance below which the droplets display translational order. The pair correlation function of the emulsion decays exponentially over distances that are at most of the order of a couple of particle diameters [16]. Second, the rather complex variations of $\xi_{HU}$ contrasts with that of all the other structural and dynamical quantities which only display a significant change at the transition point $\Delta^*$. A potential explanation is that the intrinsic slowing down of the strobed dynamics below $\Delta^*$ makes the hyperuniform self-organization too slow to be experimentally achieved, although all the other (local) observables have reached a steady state.

What causes this emulsion to self-organize into hyperuniform large-scale structures? The droplets in the experimental system interact through various forces: hydrodynamic forces that are time reversible, but also short-range irreversible forces such as depletion, van der Waals, and electrostatic forces that are specific to the nature of the fluids and surfactants forming the emulsion. To find out which of these ingredients are relevant to achieve hyperuniformity, we perform numerical simulations using a model containing only minimal hydrodynamic interactions and steric repulsion due to the finite size of the droplets. The flow induced by a moving droplet in a geometry as used in the experiment is described by a potential-flow dipole [19]. Each of the 896 droplets is advected by the local flow $u(\mathbf{r})$ with the friction being modeled through a mobility coefficient $0 < \mu < 1$. The equation of motion for droplet $i$ is $\dot{\mathbf{r}}_i = \mu u(\mathbf{r}_i)$. Assuming pairwise additive interactions, the flow at the location of particle $i$ is $u(\mathbf{r}_i)$ is the sum of the contributions from the driving flow $\mathbf{u}_0$, and from the flow induced $\mathbf{u}_j$ by all other particles $i \neq j$ in the system. The full equations of motion are

$$\dot{\mathbf{r}}_i = \mu \left( \mathbf{u}_0(t) + \sum_{j \neq i} \frac{2\mathbf{r}_{ij}\mathbf{r}_{ij} - \mathbf{1}}{2\pi|\mathbf{r}_{ij}|^3} \cdot \mathbf{r}_j \right), \quad (1)$$

where $r_j$ is the dipole vector associated with the particle $j$. The strength of the dipole is proportional to the velocity of this particle relative to the ambient fluid. The boundary conditions in the flow direction are periodic, whereas the flow is bounded by walls in the transverse direction. The infinite number of dipole images that arise due to long-ranged interactions can yield hyperuniform disordered states at thermal equilibrium [21]. A natural question is, therefore, does the emergence of hyperuniformity depend on the long-range nature of the particle-particle interactions in this nonequilibrium system as well? To answer this question, we perform the same simulations as above, but apply a very short-ranged cutoff to the hydrodynamic interactions of the form $\Theta(2a - |\mathbf{r}_{ij}|)$, where $\Theta$ is the Heaviside function, keeping all the other parameters unchanged. We shall stress that this screening preserves the reversible nature of the microscopic dynamics. As is evident from Fig. 2(a), the same reversible-to-irreversible transition, yet smoother, is observed, thereby further demonstrating the robustness of our main findings. Counterintuitively, the extent of the hyperuniform regions is clearly not reduced by screening: it extends up to the entire simulation window. As opposed to equilibrium systems, long-ranged interactions impair hyperuniform self-organization. This observation might be explained by a change in the nature of the reversibility transition from first order to critical when the interactions are short ranged. This hypothesis is supported by the sharp increase of the structural relaxation time at $\Delta^*$ [16], which is not observed for unscreened interactions. In addition, this critical scenario is consistent with that reported in Refs. [12,13,15,22] where short-range interactions are involved. However, a thorough finite-size scaling analysis that goes beyond the scope of this Letter would be required to unambiguously confirm that long-range interactions suppress criticality in our system.

Together with that of Refs. [12,13], our experimental and numerical results strongly suggest that any ensemble of particles at the onset of a reversible-to-irreversible transition should self-organize into anomalously homogeneous patterns. This robust feature could be effectively used to...
assemble a broad class of materials into hyperuniform structures from colloidal suspensions [15,23–25], to soft glasses [22], to shaken grains [26] to vortices in superconductors [27].

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