Emergence and melting of active vortex crystals

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Melting of two-dimensional (2D) equilibrium crystals, from superconducting vortex lattices to colloidal structures, is a complex phenomenon characterized by the sequential loss of positional and orientational order. Whereas melting processes in passive systems are typically triggered by external heat injection, active matter crystals can self-assemble and melt into an active fluid by virtue of their intrinsic motility and inherent non-equilibrium stresses. Emergent crystal-like order has been observed in recent experiments on suspensions of swimming sperm cells, fast-moving bacteria, Janus colloids, and in embryonic tissues. Yet, despite recent progress in the theoretical description of such systems, the non-equilibrium physics of active crystallization and melting processes is not well understood. Here, we establish the emergence and investigate the melting of self-organized vortex crystals in 2D active fluids using an experimentally validated generalized Toner-Tu theory. Performing hydrodynamic simulations at an unprecedented scale, we identify two distinctly different melting scenarios: a hysteretic discontinuous phase transition and melting through an intermediary hexatic phase, both of which can be controlled by self-propulsion and active stresses. Our analysis further reveals intriguing transient features of active vortex crystals including meta-stable superstructures of opposite spin polarity. Generally, these results highlight the differences and similarities between crystalline phases in active fluids and their equilibrium counterparts.

Melting of crystal structures is a ubiquitous phase transition phenomenon that typically requires the injection of heat from an external source [1]. An intriguing exception to this rule are active non-equilibrium crystals that can both self-assemble [2] and melt into an active fluid [4–6], owing to the intrinsic motility of their microscopic constituents. The spontaneous emergence and destruction of crystal-like order can be observed in a wide variety of natural and artificial systems [2] [7]–[9]. Striking examples range from suspensions of active colloids [2] [10] [11], bacteria [7] and sperm cells [9] to biological tissues [8] that can solidify and fluidize during embryonic development [12]–[14]. Yet, despite recent experimental advances [7] [9] [12] [14] [15] and important theoretical progress [6] [12] [13] [16] [23], many key aspects of active melting processes remain poorly understood. This may not be a big surprise given that it took several decades to decipher the complex melting scenarios of even the most basic 2D equilibrium crystal structures [1].

In thermal equilibrium, 2D crystals can exhibit long-range orientational order while positional order is suppressed at finite temperatures [20]–[29]. Over the last decades, several competing theories on the type and nature of solid-liquid transitions in 2D equilibrium systems have been proposed [1]. Approaches such as density-functional [30] and grain-boundary theories [31] suggest possible mechanisms for first-order transitions. Seminal work by Kosterlitz, Thouless, Halperin, Nelson, and Young (KTHNY) [32] predicted a two-step continuous melting transition which proceeds through an intermediate hexatic phase characterized by quasi-long-range orientational order and short-ranged positional order. Such a hexatic phase has been observed in experiments on colloidal systems [33] and superconducting lattices [34] as well as in numerical simulations of repulsive disks [35].

The complex melting dynamics of 2D equilibrium crystals raises fundamental questions about liquid-solid transitions in far-from-equilibrium systems [6] [36]–[38]. In particular, it remains an open question to which extent concepts from equilibrium melting processes translate to non-equilibrium melting. Thus far, experimental and numerical studies of particulate active matter draw a complex picture. Whereas Monte Carlo simulations for active particles with inverse-power-law repulsion [4] reported an intermediate hexatic phase consistent with KTHNY scaling prediction, agent-based simulations [5] and active Brownian particle simulations [6] suggest that active crystal structures can melt into a hexatic state without the KTHNY-typical unbinding of topological defect pairs.

Active fluids are another important class of non-equilibrium systems, which show intriguing transitions from active turbulence [39]–[43] to highly ordered vortex arrays. Vortex arrays have been observed in dense suspensions of swimming sperm cells [9] or microtubule [44], and have been have been predicted to form spontaneously by a wide range of generic active fluid models [40] [41] [43] [45] [50]. However, so far it has not been established whether such active systems exhibit true crystalline order, and if so, how active vortex crystals (AVCs) melt.

The systematic study of AVC formation and melting has remained an unsolved challenge owing to the pro-

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FIG. 1. Active vortex crystals and their melting: (a) Our active fluid simulation (solving (1) with \( \lambda = 7 \), \( \alpha = -0.8 \) on a \( 1000 \pi \times 1000 \pi \) domain) exhibits AVC superstructures comprised of meta-stable opposite-polarity domains. The zoom-in shows that these domains are demarcated by an interfacial layer of turbulent active fluid. Panel (b) illustrates the melting of the AVC superstructures shown in panel (a) for \( \lambda = 5.6 \) after time \( t = 500, t = 1000, \) and \( t = 5000 \). (c) Area fraction of the crystal domains as a function of active advection for different times, starting from the superstructure a at time \( t = 0 \). Note that the transient width of the domain boundaries in the superstructures is controlled by the strength of the active advection. The orange line corresponds to the snapshots shown in panel (b).

RESULTS

Our starting point is a generalization of the incompressible Toner-Tu equations \([52-54]\) for the active fluid velocity field \( \mathbf{u} \) and pressure field \( p \) \([39, 40, 51]\):

\[
\partial_t \mathbf{u} + \lambda \mathbf{u} \cdot \nabla \mathbf{u} = -\nabla p - (1 + \Delta)^2 \mathbf{u} - (\alpha + \beta \| \mathbf{u} \|^2) \mathbf{u}, \\
\nabla \cdot \mathbf{u} = 0,
\]

where \( \lambda \) is an active advection parameter which incorporates the effects of active nematic stresses \([40, 55, \text{56}]\). The activity parameter \( \alpha < 0 \) is proportional to the square of self-propulsion velocity, and \( \beta \) sets the velocity relaxation time scale (which can be scaled out). Active flows described by (1) are self-driven through a linear instability induced by the Swift-Hohenberg operator \((1 + \Delta)^2\), which favors periodic flow patterns of wave length \( 2\pi \) \([57, 58]\). \( \alpha \) can be derived from a generic agent-based model \([55, 56]\) that accounts for the particles’ self-propulsion, their hydrodynamic interactions near a surface, and their steric interactions. It has been shown \([59]\) that this minimal continuum theory quantitatively captures essential statistical properties of dense
bacterial suspensions in quasi-2D microfluidic channels. Here, we apply (1) to study AVC dynamics by performing large-scale numerical simulation on a doubly periodic domain of size $L \times L$ with $L$ ranging up to 1000$\pi$ (Fig. 1), using a pseudo-spectral method for the spatial discretization and a fourth-order Runge-Kutta scheme for time stepping (Methods).
Large-scale active vortex crystals.

As an illustration of the AVC emerging in our system, Fig. 1 shows the vorticity field for a simulation on a $1000\pi \times 1000\pi$ domain resolved with $8192^2$ grid points. At this very large system size, AVC superstructures, i.e., crystal domains of opposite polarity with domain boundaries comprised of active turbulence regions, crystallize from an initially turbulent active fluid. The highly dynamic domain boundaries play a prominent role in the melting of AVC superstructures, which can be induced, for example, by decreasing the active advection parameter. As active advection is decreased, the superstructures melt, and the turbulent boundary layers spread in area, destroying the crystal structure (Fig. 1b). Since the AVC superstructures are formed by crystal domains demarcated by an active fluid interfacial layer, a natural order parameter for our analysis is the fraction of area covered by the crystal domain $A_{\text{crystal}}/A_{\text{total}}$ (Methods).

To illustrate this transition, we evaluate the crystal area fraction as a function of active advection (at a fixed activity $\alpha = -0.8$) for different times, which is shown in Fig. 1. Below $\lambda \approx 5.6$, the crystal domains melt completely into a statistically isotropic active fluid. Above a critical value of $\lambda = 6.0$, almost the entire domain is covered by vortex crystals, with the area between the crystals of different polarity occupied by a layer of active fluid. There is a consistent, but slow decrease in the area of this boundary layer as advection is increased. In the following sections, we present a detailed characterization of the crystalline order and its melting.

Non-equilibrium phase diagram.

We performed $\sim 1000$ simulations on smaller domains of size $L = 20\pi$ to map out the $(\alpha, \lambda)$ phase diagram shown in Fig. 2. These parameter scans revealed three distinct phases: The AVC phase Fig. 2b forms for $\alpha \approx -0.8$ and sufficiently large values of the active advection parameter $\lambda$, corresponding to strong extensile stresses (red-colored domain in Fig. 2h). In this phase, vortices of same spontaneously chosen handedness self-organize in a triangular lattice, phenomenologically similar to those observed in dense sperm suspensions [9]. Strikingly, this spontaneous symmetry breaking occurs after an initial turbulent transient [Supplementary Video (2)]. The AVC phase is surrounded by an extended active turbulence (AT) phase (green domain in Fig. 2h), in which transient vortices of either handedness coexist in the fluid (Fig. 2i). Finally, for low active advection $\lambda \ll 1$, corresponding to contractile stresses [10], the system settles into a stationary square flow-lattice state (Fig. 2i), which can be explained with classical pattern formation theory [42].

Unexpectedly, our simulations reveal two distinct AVC-AT transition scenarios, characterized by phase coexistence and hysteresis, respectively. To demonstrate the characteristics of phase coexistence (gray in Fig. 2g), we keep the activity parameter $\alpha = -0.8$ fixed and decrease the active advection parameter $\lambda$ (vertical scan). This transition is characterized by an intermittent switching between active turbulence and AVCs [Supplementary Video (3)]. The energy time series of a corresponding simulation is shown in Fig. 2g. In the AVC phase (red-colored domain in Fig. 2a, we keep the active advection fixed at value $\lambda = 7$ and change the activity parameter $\alpha$ (horizontal scan). In the transition region for low activities, a vortex crystal will not emerge from random initial conditions for these parameters, but the crystal itself is a stable solution. This is illustrated by the AVC time fraction for the horizontal scan shown in Fig. 2a, which clearly exhibits a hysteresis loop. As the activity parameter is further increased, a second transition without hysteresis is observed. Closer to the boundary of the active turbulence region, the vortex arrays start showing a fluid-like arrangement of vortices rather than crystal-like or hexatic. For a systematic characterization, we present results from significantly larger domains in the following sections.

Hexatic phase.

To characterize the crystalline order in this system, we perform simulations that are two orders of magnitude larger than for the phase diagram ($L = 200\pi$), and construct a Voronoi partition to identify 5-fold and 7-fold defects (Methods). Deep inside the crystal regime, we observe a well-ordered structure with bound pairs of dislocations (Fig. 3a: $\lambda = 15, \alpha = -0.8$). Closer to the active turbulence region, the order starts unravelling (Fig. 3b: $\lambda = 7, \alpha = -0.75$, and c: $\lambda = 7, \alpha = -0.7$). In Fig. 3a, we observe the unbinding of dislocation pairs, as well as a few free disclinations. As we move closer to the active turbulence region, Fig. 3b, clusters of defects emerge. As we demonstrate below, Fig. 3b corresponds to a hexatic phase, which points at a connection to equilibrium melting transitions following the KTHNY scenario.

As a quantitative characterization of the crystalline phase we use the dynamic Lindemann parameter, which is defined as the relative displacement of neighboring vor-
Emergence of AVCs.

Next, we characterize the emergence of AVCs as a function of system size. To this end, we have determined the transient time until a uniform AVC is formed for an ensemble of 100 simulations for each system size, covering domain sizes between $L=10\pi$ and $L=160\pi$. Fig. 4a shows the resulting scatter plot, which demonstrates that the lifetime of the transient active fluid depends sensitively on the initial condition and increases considerably with domain size.

For small domains, this is mainly rooted in the fact that the emergence of a uniform AVC occurs after a spontaneous discrete symmetry breaking through a turbulent transient [13], which renders the transient time a random variable. In fact, the PDF of transition times is well captured by

$$P(T) = \frac{\delta}{\tau} \left[1 - e^{-\frac{T}{\tau}}\right]^{\delta-1} e^{-\frac{T}{\tau}},$$

where $\tau$ and $\delta$ depend on the vortex lifetime and domain size, respectively. This expression can be rationalized from the observation that vortex lifetimes in active turbulence have an approximately exponential distribution [12]. A good estimate for the transient time is the time after which the spontaneous symmetry breaking occurs. Its distribution can be obtained from the PDF of the time it takes for one polarity of vortices to decay, which amounts to computing the maximum survival time of a set of like-signed vortices. Assuming statistical independence of the individual decay processes yields the proposed PDF (4). Figure 4b (inset) shows the corresponding fits for the PDFs of the transient durations obtained from $10^4$ simulations for each domain size, demonstrating excellent agreement.

For sufficiently large domain sizes, an additional effect comes into play: crystal domains with both polarity can coexist, leading to metastable AVC superstructures with very long lifetimes (see, e.g., Fig. 1h). The temporal evolution of AVC clusters is illustrated in Fig. 4b, which shows the number of positive and negative vortices as a function of time ($L=160\pi$). In this example, two vortex clusters of approximately equal sizes but opposite polarity coexist for more than 8000 nondimensional time units, before a uniform AVC forms. These metastable
AVC superstructures explain the extreme outliers in the transient duration which are the cause for the sharp increase of the mean transient time for system sizes beyond $L = 120\pi$. This divergence of transient durations with domain size, a supertransient feature, is observed in a variety of dynamical systems such as coupled map lattices and reaction diffusion systems [61–64]. Unlike supertransient chaos, our transient state is characterized initially by a turbulent regime and then a gradually evolving superstructure. An extreme example of such a superstructure is shown in Fig. 4 for a domain of size $L = 1000\pi$. The slowly evolving crystal domains are separated by a highly dynamic interfacial area of active turbulence [Supplementary Video (1)].

**DISCUSSION**

Using large-scale simulations at unprecedented scale, we have determined the comprehensive phase diagram of a generalized Toner-Tu model for active fluids. Our analysis establishes that the self-organized active vortex arrays show crystalline order and predicts a rich melting scenario from active vortex crystals to active turbulence. Depending on the path through the parameter space, we find two distinct transition scenarios. The first transition scenario resembles a KTHNY-like melting. When transitioning from the AVC phase to the AT phase, we first observe the unbinding of defect pairs, which results in the loss of long-range orientational order before melting into an active fluid. In this parameter range, small domains exhibit a highly dynamic melting and re-crystallization. On very large domains, broken-symmetry AVC domains of opposite polarity emerge, whose melting results from the spreading of the turbulent interfacial layers. Intriguingly, in contrast to the KTHNY-like melting, the second transition scenario shows hysteresis between the AVC and the AT state. Generally, our results reveal the roles of nonlinear advection and activity in the complex self-assembly and melting of AVCs, and highlight deep connections between phase transitions in active matter and their classical equilibrium counterparts.

Experimental tests of our predictions could be an exciting direction for future work. Dense suspension of spermatozoa, which show both, active turbulence [65] as well as self-organized regular vortex arrays [9] are arguably the best candidate to test our prediction on AVC-AT transitions. Although previous experiments [9] so far suggest a fluid-like arrangement of vortices, rather than crystal-like or hexatic, it may be possible to achieve long-range crystal order through a careful tuning of experimental conditions [66]. For instance, the type of sperm cells as well as the intracellular ionic concentrations could affect the nature of sperm motility [67]. Furthermore, the analysis of crystalline order in such systems would also require conducting experiments on large domains. If a vortex crystal phase is achieved, the activity can be tuned, for instance, by changing the motility through the ambient temperature [68] to induce a potential melting transition. The preferred handedness of sperm cells on planar surfaces [69] precludes the observation of a spontaneously broken chiral symmetry. This could be alleviated by confinement between two walls. Such experiments will significantly enhance our knowledge of crystalline order, not just in active matter, but in out-of-equilibrium systems in general.
METHODS

Simulation details.

We perform direct numerical simulations of the vorticity field \( \omega = \nabla \times u \) in a periodic domain by using a fully dealiased pseudo-spectral algorithm. The mean velocity \( \langle u \rangle \) is integrated separately. The corresponding evolution equations follow from (1) and take the form:

\[
\partial_t \omega + \lambda u \cdot \nabla \omega = -\left(1 + \Delta\right) \omega - \alpha \omega - \beta \nabla \cdot \left( |u|^2 u \right) \tag{5}
\]
\[
\partial_t \langle u \rangle = -(1 + \alpha) \langle u \rangle - \beta \langle |u|^2 u \rangle. \tag{6}
\]

We solve (5) and (6) with a fourth-order Runge-Kutta method for time stepping combined with an integrating factor for the linear terms. Our code is parallelized using GPUs (graphics processing units) in order to accelerate the computations. For the rest of the text, the parameter values are listed in Table I.

Phase diagram.

The phase diagram (Fig. 2a) is obtained from simulations of 477 different parameter configurations as shown in Fig. 1. For each configuration, we use two different initial conditions: a random initial condition and a vortex crystal.

The different phases shown in Fig. 2a are defined as follows. The square lattice, active turbulence and vortex crystal phases show obvious qualitative differences as noted in the main text and are easily distinguished visually. The hysteresis phase in the marginal stability region is identified as such when the simulations are bistable; the simulations starting with random initial conditions result in an active turbulence phase whereas a vortex crystal initial condition remains stable. The simulations are checked for convergence until a total simulation time of \( T = 2000 \left( 4 \times 10^5 \right) \) steps). The coexistence region is defined by evaluating the PDF of the energy density. If the PDF has two peaks (see, e.g., Fig. 2b), it is defined as a temporally intermittent pattern.

Phase transition.

The phase transition between active turbulence and vortex crystals in small domains (Fig. 2g and h) is characterized as follows. For the transition curves in both increasing and decreasing directions of parameter values, we conduct simulations in the range \( 5.0 \leq \lambda \leq 7.0 \) and \(-1.1 \leq \alpha \leq -0.6 \). For \( \lambda = 5 \) and \( \alpha = -1.1 \), we start our simulation from random initial conditions. For the rest of the simulations, the final snapshot of the previous simulation is used as the initial condition. Once a statistically steady state is reached (after about \( 10^6 \) time steps), we collect data for \( 10^6 \) time steps and evaluate the PDF of the energy density. If the PDF has only one peak, the order parameter \( T_{\text{crystal}}/T_{\text{total}} \) takes the value 0 or 1, depending on the phase. Otherwise, the energy density at the minimum between the two peaks of the PDF, \( E_{\text{min}} \), is evaluated. The order parameter then takes the value of the probability that the energy density \( E > E_{\text{min}} \). This process is repeated five times, and the mean and the standard deviations are used to construct the transition curves and estimate the uncertainties, which are shown in Fig. 2k and l.

To evaluate the phase transition curve in large domains (Fig. 1k), we first identify centers of the strong vortices \( 12 \cdot 70 \). Then an order parameter field is obtained by calculating, for each point \( (x, y) \), the difference between the number of positive and negative vortices within a circle of radius \( r \) centered at \( (x, y) \) (\( r \) is about half the mean distance between the nearest neighbors and next nearest neighbors). The resulting field is then smoothed using a Gaussian filter with standard deviation \( \sigma = 12 \). The original vorticity field and the smoothed field are shown in Fig. 6. The turbulent region is then defined as the area where the absolute value of this smoothed field is less than half the maximum value of the field. Once this turbulent region is defined, the

FIG. 5. Overview of simulation parameters: The crosses show the parameter configuration used to obtain the phase diagram (Fig. 2, domain size \( L = 20\pi \)). Each cross represents simulations with two different initial conditions, as noted in the main text. The PDFs in Fig. 2 are based on simulations with parameters marked in blue (\( L = 20\pi \)). The phase transition curves Fig. 2k and l are based on simulations with parameters marked in green (\( L = 20\pi \)). The parameters used in Fig. 3 are indicated in red (\( L = 200\pi \)). Fig. 3 (\( L = 10\pi - 160\pi \)) and Fig. 4 (\( L = 1000\pi \)) are based on the parameter choice shown in yellow. Fig. 1k (\( L = 1000\pi \)) and Fig. 2k (\( L = 20\pi \)) are based on the parameter choice indicated by a blue circle. See also Table I for the parameter values.
order parameter $A_{\text{crystal}}/A_{\text{total}}$ is calculated by evaluating the fraction of the total area covered by the vortex crystal.

**Table I.** Simulation parameters: Domain size $L$, active advection parameter $\lambda$, activity parameter $\alpha$, number of grid points $N$, time step $\Delta t$. The parameter $\beta$ is set to 0.01 in all simulations.

| Figure | $L$   | $\lambda$ | $\alpha$ | $N$    | $\Delta t$ |
|--------|-------|-----------|----------|--------|-----------|
| 1a     | $1000\pi$ | 7.0       | -0.800   | $8192^2$ | 0.005     |
| 1b     | $1000\pi$ | 5.7       | -0.800   | $8192^2$ | 0.005     |
| 1c     | $1000\pi$ | [5, 6.5]  | -0.800   | $8192^2$ | 0.005     |
| 2a     | $20\pi$  | [0.0, 15.0] | [-1.200, 0.000] | $256^2$ | 0.005     |
| 2b     | $20\pi$  | 8.0       | -0.800   | $1024^2$ | 0.001     |
| 2c     | $20\pi$  | 3.0       | -0.800   | $1024^2$ | 0.001     |
| 2d     | $20\pi$  | 0.1       | -0.200   | $1024^2$ | 0.001     |
| 2e     | $20\pi$  | 5.6       | -0.800   | $256^2$  | 0.005     |
| 2f     | $20\pi$  | 5.0, 5.6, 6.6 | -0.800 | $256^2$  | 0.005     |
| 2g     | $20\pi$  | [5.0, 7.0] | -0.800   | $256^2$  | 0.005     |
| 2h     | $20\pi$  | 7.0       | [-1.100, -0.600] | $256^2$ | 0.005     |
| 3a     | $200\pi$ | 15.0      | -0.800   | $1024^2$ | 0.005     |
| 3b     | $200\pi$ | 7.0       | -0.750   | $1024^2$ | 0.005     |
| 3c     | $200\pi$ | 7.0       | -0.700   | $1024^2$ | 0.005     |
| 3d     | $200\pi$ | 7.0       | -0.750, -0.725, -0.715, -0.700 | $1024^2$ | 0.005     |
| 3e     | $200\pi$ | 15.0      | -0.800   | $1024^2$ | 0.005     |
| 3f     | $200\pi$ | 7.0       | -0.750, -0.725, -0.715, -0.700 | $1024^2$ | 0.005     |
| 3g     | $200\pi$ | 15.0      | -0.800   | $1024^2$ | 0.005     |
| 4a inset| [10\pi, 160\pi] | 7.0 | -0.800 | $256^2$, $512^2$, $1024^2$ | 0.005     |
| 4a     | $20\pi$, $40\pi$ | 7.0 | -0.800 | $256^2$ | 0.005     |
| 4b     | $160\pi$ | 7.0       | -0.800   | $1024^2$ | 0.005     |
| 6      | $1000\pi$ | 5.6       | -0.800   | $8192^2$ | 0.005     |

**FIG. 6.** Analysis of melting of AVC superstructures: Top row shows the vorticity field for $\lambda = 5.6$ at different times (initial condition, at $t = 1000$ and at $t = 2000$). Bottom row shows the corresponding smoothed fields.

**Transient durations.**

To evaluate transient durations (Fig. 4a and b), we conduct simulations starting with random initial conditions until a converged vortex crystal state is reached for each domain size. The convergence is defined as follows. By employing a vortex identification algorithm [42][70], we obtain a time series of the number of strong vortices of both polarity. A converged vortex crystal is obtained when the number of vortices of either sign reaches 93% of the theoretical maximum number of vortices possible in the domain. To obtain the mean and median transient durations in Fig. 4a, the simulations are repeated 100 times for each domain size and the corresponding mean and median durations are calculated. The PDFs (inset Fig. 4h) are obtained by evaluating the transient durations for three different domain sizes from $10^4$ simulations each, starting from random initial conditions. The error bars correspond to the difference between the maximum and the minimum from the 5 bootstrapped PDFs obtained from 2000 simulations each. The theoretical curves are obtained by fitting (4) to the numerical data. The corresponding values of the free parameters $\delta$ and $\tau$ are, respectively, 4.97 and 54.59 for $L = 10\pi$ (blue...
Defects, dynamic Lindemann parameter and orientational correlation.

For each parameter choice in Fig. 3, simulations starting from random initial conditions are run for at least $1.2 \times 10^7$ time steps until a statistically steady state is reached. To identify the defects, centers of strong vortices are first evaluated [42, 70]. By constructing a Voronoi partition (using the Python open source module scipy.spatial.Voronoi) over this vortex core configuration, 5-fold and 7-fold defects are determined [3].

To evaluate the dynamic Lindemann parameter, we obtain 500 snapshots separated by $\Delta t = 0.1$, after the simulations have reached a converged state. For each snapshot, we identify the centers of strong vortices. The trajectory of each vortex core is then tracked. Only vortices which survive the entire duration of the simulation after reaching the statistically steady state are included in the analysis. The dynamic Lindemann parameter is then evaluated following 4 in the main text.

The orientational correlation is evaluated according to 3 in the main text, after identifying the centers of the strong vortices in converged simulations. The results are averaged across 10 snapshots (with $\Delta t = 2000$) each from three simulations with different initial conditions.

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AUTHOR CONTRIBUTIONS

M.J. and M.W. designed the research. D.A.S. wrote the DNS code. M.J. and D.A.S. conducted the analysis. All authors contributed to the interpretation of the results and the writing of the manuscript.

ADDITIONAL INFORMATION

List of supplementary videos.

1. Evolution of AVC superstructures. The inset shows a zoom-in into the turbulent interfacial area.

2. Emergence of a vortex crystal starting from random initial conditions.

3. The active matter system in the marginal stability region between the turbulent phase and the vortex crystal. The right panel shows how the energy density changes with time.

Competing financial interests.

The authors declare no competing financial interests.