Viscoelastic and elastomeric active matter: Linear instability and nonlinear dynamics

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We consider a continuum model of active viscoelastic matter, whereby an active nematic liquid crystal is coupled to a minimal model of polymer dynamics with a viscoelastic relaxation time τ_C . To explore the resulting interplay between active and polymeric dynamics, we first generalize a linear stability analysis (from earlier studies without polymer) to derive criteria for the onset of spontaneous heterogeneous flows (strain rate) and/or deformations (strain). We find two modes of instability. The first is a viscous mode, associated with strain rate perturbations. It dominates for relatively small values of τ_C and is a simple generalization of the instability known previously without polymer. The second is an elastomeric mode, associated with strain perturbations, which dominates at large τ_C and persists even as $\tau_C \to \infty$. We explore the dynamical states to which these instabilities lead by means of direct numerical simulations. These reveal oscillatory shear-banded states in one dimension and activity-driven turbulence in two dimensions even in the elastomeric limit $\tau_C \to \infty$. Adding polymer can also have calming effects, increasing the net throughput of spontaneous flow along a channel in a type of drag reduction. The effect of including strong antagonistic coupling between the nematic and polymer is examined numerically, revealing a rich array of spontaneously flowing states.

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I. INTRODUCTION

Examples of active matter include bacterial swarms, the cellular cytoskeleton, and *in vitro* cell extracts that comprise only polymeric filaments, molecular motors, and a fuel supply [1–4]. Such materials not only are of direct biophysical significance, but also represent a wider class of systems in statistical physics in which strong deviations from thermal equilibrium emerge at the collective macroscopic scale due to the underlying active dynamics of the system's microscopic subunits (rotating bacterial flagella, marching molecular motors that bridge cytoskeletal filaments, etc.). Arising at the level of the microscopic subunits, this driving is distinct from the macroscopic boundary driving of, for example, an imposed shear flow.

For an active particle subject to no externally imposed force, the simplest perturbation it can exert on the local fluid environment is that of a force dipole. Depending on the sign of this dipole, the activity is classified as extensile (where the two forces act from the center of mass of the dipole outward towards the fluid) or contractile (where the opposite applies). Collectively, a fluid of these active particles can exhibit nonequilibrium emergent phenomena on macroscopic length scales that greatly exceed the particle size or spacing. These include activity-induced ordering and bulk fluid flows that arise spontaneously even in the absence of external driving. Depending on the strength of the activity, these flows may remain steady and laminar at the scale of the system, show oscillatory limit cycles at that scale or below, or exhibit spatiotemporal chaos. The latter effect closely resembles conventional inertial turbulence in a passive Newtonian fluid and is accordingly often termed active turbulence or bacterial turbulence [2,5-9]. However, its mechanism is distinct from that of inertial turbulence: It stems from a balance between active stress and orientational relaxation, rather than between inertia and viscosity.

Depending on their symmetry, ordered phases of active fluids can be described by either a polar [10,11] or a nematic

order parameter [1,12,13]. In this work we consider the latter case and throughout denote the nematic order parameter by **Q**. Indeed, many theoretical descriptions of active matter are based on simple continuum models for the hydrodynamics of a suspension of rodlike objects, originally developed to describe a passive liquid crystal [14,15]. To describe an active material, such a model is then augmented by the leading-order terms characterizing the violations of time-reversal symmetry that arise from activity. In particular, this gives an additional active contribution $\Sigma_A = -\zeta \mathbf{Q}$ to the fluid's stress tensor, where the activity parameter ζ is positive for extensile systems and negative for contractile ones.

Even without detailed knowledge of the value of ζ , the approach just described is capable of robust predictions. For example, beyond a critical threshold level of activity an initially quiescent fluid is generically predicted to become unstable to the formation of a spontaneously flowing state [12]. This threshold, for a finite system size, depends on whether the system is extensile or contractile. However, in both cases it decreases with increasing system size [10], tending to zero in the limit of a bulk sample. Any level of activity, however small, can then trigger the formation of spontaneous flows. Consistent with these analytical predictions, numerical solutions of active nematic continuum models [5,7-9] have indeed revealed a host of spontaneously flowing states resembling experimental observations in bacterial swarms [2] and microtubule-based cell extracts [4]. Both of these are extensile nematics, to which we restrict ourselves in our numerical study below [11].

Active nematic fluids are often referred to informally as active gels [5,10], particularly in a biological context. However, while all liquid crystals are somewhat viscoelastic, for example, due to the slow motion of topological defects, most existing models of active matter assume fast local relaxations and so fail to model gels in the conventional sense of the term, as understood by polymer or colloid physicists [16,17]. Certainly the standard continuum models of active matter [1,13] do not capture the types of slow viscoelastic dynamics that one might expect in the cytoskeleton, which contains long-chain flexible polymers and other cytoplasmic components that are expected to have long intrinsic relaxation times (or even divergent ones in the case of a cross-linked network). This is a major shortcoming, because this slow viscoelastic dynamics would be expected to couple strongly to the active liquid-crystalline dynamics and thereby potentially radically modify the effects of activity.

The effect of a viscoelastic polymeric background is also likely to play an important role in modifying active flows and diffusion [18] at a supracellular level. Indeed, long-chain molecules are present in mucus, saliva, and many other viscoelastic fluids both inside and outside the body that are susceptible to colonization by swarms of motile bacteria. Moreover, many bacteria secrete their own polymers [19], particularly during biofilm formation (which, however, is not the topic of this paper). This suggests an evolutionary advantage for bacteria in controlling the viscoelasticity of their surroundings, supporting the view that viscoelasticity and active motion are coupled in a nontrivial way.

With this motivation, the aim of this paper is to study in greater depth the predictions of a model first presented in Ref. [20], which addresses at a continuum level the interplay between active liquid-crystalline dynamics and slow polymeric modes of relaxation. This approach is distinct from, but complements, recent studies of individual swimmers in viscoelastic fluids [21–25], which show that swimming speeds can be either enhanced or suppressed relative to those in a Newtonian solvent, depending on details of the swimming mechanism and particle geometry. Such details do not enter our continuum picture, however, which focuses on emergent and potentially universal many-body behavior at larger length scales.

In a biological context, active matter is often found in confined geometries such as the interior of a cell. Indeed, it has been argued that the confinement of subcellular active matter may in part be responsible for cytoplasmic streaming [26,27], an important process whereby coherent fluid flows facilitate the circulation of nutrients and organelles within the cell [28]. At a larger scale, a recent study of cell migration in artificial channels observed increases in mean cell velocity and flow coherence as the channel was narrowed [29]. Also, suspensions of *B. subtilis* were observed to form stable spiral structures when confined in a droplet [30]. In view of these observations, it is important that any numerical study should consider carefully the effects of system size. In what follows we choose a rectangular channel geometry of fixed aspect ratio bounded by a pair of parallel hard walls. The size of this channel is notionally fixed in simulation units, but we study finite-size effects by then varying instead the microscopic length scales in the problem.

As we will explore in detail below, the interplay of active and polymeric dynamics leads to a host of exotic spontaneous flow states. These include oscillatory shear-banded states in one dimension, while in two dimensions we find activitydriven turbulence even in the limit of infinite viscoelastic relaxation time in which our model describes an active elastomer. In other regimes we find that adding polymer can have calming effects, increasing the net throughput of spontaneous flow along a channel in a type of drag reduction. The paper is structured as follows. In Sec. II we review the equations of motion that describe the coupled dynamics of an active nematic with a viscoelastic polymer, as derived in Ref. [20]. In Sec. III we discuss the simulated sample geometry and give details of our numerical methods. In Sec. IV we perform a linear stability analysis to derive the threshold of instability to spontaneous flow. The results of this linear calculation then provide a route map for performing full nonlinear simulations in Secs. V and VI. In Sec. V we restrict those simulations to one spatial dimension for simplicity, while in Sec. VI we perform full two-dimensional (2D) simulations. Section VII contains a summary of our results and the outlook for future work.

II. MODEL EQUATIONS

In this section we remind the reader of the model equations, as developed originally in Ref. [20]. The state of liquid-crystal (nematic) ordering is denoted by a traceless symmetric tensor $\mathbf{Q} = q \langle \hat{\mathbf{n}} \hat{\mathbf{n}} - \frac{1}{2} \mathbf{I} \rangle$, where $\hat{\mathbf{n}}$ is the nematic director and q is the degree of ordering. The polymeric conformation is similarly denoted by the symmetric tensor $\mathbf{C} = \langle \mathbf{rr} \rangle$, where **r** is the endto-end vector of a chain (or subchain, depending on the level of description used), normalized so that $\mathbf{C} = \mathbf{I}$ in equilibrium. We adopt a single-fluid description in which the concentration fields of polymer and liquid crystal are assumed to remain uniform (in contrast to a two-fluid approach as taken by, for example, Ref. [31]). Accordingly, all three componentsliquid crystal, polymer, and solvent-share the same centerof-mass velocity v. The symmetric and antisymmetric parts of the velocity-gradient tensor $(\nabla \mathbf{v})_{ij} \equiv \partial_i v_j$ are denoted by **D** and Ω , respectively. For any other tensors the symmetric, antisymmetric, and traceless parts carry superscripts S, A, and T.

We introduce a free-energy density $f = f_Q(\mathbf{Q}, \nabla \mathbf{Q}) + f_C(\mathbf{C}) + f_{QC}(\mathbf{Q}, \mathbf{C})$, where f_Q and f_C are the standard forms for nematics [14] and dumbbell polymers [32], respectively. Accordingly we have

$$f_{\mathcal{Q}} = G_{\mathcal{Q}} \left[\frac{(1 - \gamma/3)}{2} \operatorname{Tr} \mathbf{Q}^2 - \frac{\gamma}{3} \operatorname{Tr} \mathbf{Q}^3 + \frac{\gamma}{4} (\operatorname{Tr} \mathbf{Q}^2)^2 \right] + \frac{K}{2} (\nabla_i \mathcal{Q}_{jk})^2, \qquad (1)$$

in which G_Q sets the scale of the bulk free-energy density, *K* is the nematic elastic constant, and γ is a control parameter for the isotropic-nematic transition. Likewise

$$f_C = \frac{G_C}{2} (\text{Tr}\mathbf{C} - \ln \det \mathbf{C}), \qquad (2)$$

where G_C is the polymer elastic modulus.

The lowest-order passive coupling between Q and C is

$$f_{\rm QC} = \kappa \operatorname{Tr}(\mathbf{C} - \mathbf{I})\operatorname{Tr}(\mathbf{Q}^2) + 2\chi \operatorname{Tr}(\mathbf{C}\mathbf{Q}), \qquad (3)$$

where both terms vanish for undeformed polymers ($\mathbf{C} = \mathbf{I}$). Here κ controls how the polymer pressure shifts the isotropicnematic transition and for simplicity we set $\kappa = 0$ throughout. The second term makes it energetically preferable for \mathbf{Q} and \mathbf{C} to align with major axes parallel for $\chi < 0$ and perpendicular for $\chi > 0$. From the volume-integrated free energy $F = \int f \, dV$, the nematic molecular field $\mathbf{H} \equiv -[\delta F / \delta \mathbf{Q}]^{\text{ST}}$ follows as

$$\mathbf{H} = -G_{\mathcal{Q}} \left[\left(1 - \frac{\gamma}{3} \right) \mathbf{Q} - \gamma \mathbf{Q}^{2} + \gamma \mathbf{Q}^{3} \right] - G_{\mathcal{Q}} \gamma \frac{\mathbf{I}}{3} \operatorname{Tr}(\mathbf{Q}^{2}) + K \nabla^{2} \mathbf{Q} - 2\kappa \operatorname{Tr}(\mathbf{C} - \mathbf{I}) \mathbf{Q} - 2\chi \mathbf{C}^{\mathrm{T}}.$$
(4)

The corresponding molecular field $\mathbf{B} \equiv -[\delta F/\delta \mathbf{C}]^{\mathrm{S}}$ for the polymer obeys

$$\mathbf{B} = -G_C(\mathbf{I} - \mathbf{C}^{-1})/2 - \kappa \mathbf{I} \operatorname{Tr}(\mathbf{Q}^2) - 2\chi \mathbf{Q}.$$
 (5)

Using these molecular fields, we follow Ref. [20] in developing minimally coupled equations of motion for Q and C that respectively reduce to the Beris-Edwards liquid-crystal theory and the Johnson-Segalman (JS) polymer model in appropriate limits [14]. We then allow for conformational diffusion in the polymer sector [33], which adds a gradient term in C of kinetic origin [20]. (Alternatively, one can incorporate a nonlocal term in the polymer free energy, though this then produces a more complex form for the polymer stress [34].) To this is added a minimal set of active terms [12], supposing for simplicity that the polymers are not themselves active, but that the origin of the activity resides entirely in **Q**. This is enough to capture, for example, the effect of adding polymer to a cell extract or the collective dynamics of bacterial suspensions in mucus. (A contrasting approach is to build a system of polymers directly from active elements [35].) There remain two active terms linear in \mathbf{Q} . One of these can be absorbed into f_Q and the other is the familiar active deviatoric stress $\Sigma_A = -\zeta \mathbf{Q} \ [\mathbf{12}].$

The resulting equations of motion for Q and C are

$$(\partial_t + \mathbf{v} \cdot \nabla)\mathbf{Q} = \mathbf{Q}\mathbf{\Omega} - \mathbf{\Omega}\mathbf{Q} + \frac{2\xi}{3}\mathbf{D} + 2\xi[\mathbf{Q}\mathbf{D}]^{\text{ST}} - 2\xi\mathbf{Q}\text{Tr}(\mathbf{Q}\mathbf{D}) + \tau_Q^{-1}\mathbf{H}/G_Q, \qquad (6)$$

$$(\partial_t + \mathbf{v} \cdot \nabla)\mathbf{C} = \mathbf{C}\mathbf{\Omega} - \mathbf{\Omega}\mathbf{C} + 2a[\mathbf{C}\mathbf{D}]^{\mathrm{S}} + \tau_C^{-1}(2[\mathbf{B}\mathbf{C}]^{\mathrm{S}}/G_C + \ell_C^2\nabla^2\mathbf{C}).$$
(7)

Here ξ is the flow-alignment parameter of the nematic [36] and *a* is the slip parameter of the JS model. Setting *a* = 1 recovers the Oldroyd-B model [37]. Each controls the relative tendency of molecules to align with streamlines versus rotating with the local vorticity. The parameters τ_Q and τ_C are intrinsic local relaxation times for the nematic and polymer, while ℓ_C governs conformational diffusion in the JS sector [33].

The velocity field \mathbf{v} obeys the Navier-Stokes equation for an incompressible fluid

$$\rho(\partial_t + v_\beta \partial_\beta) v_\alpha = \partial_\beta \Sigma_{\alpha\beta},\tag{8}$$

$$\partial_{\alpha} v_{\alpha} = 0. \tag{9}$$

Here the total stress $\Sigma = -P\mathbf{I} + 2\eta\mathbf{D} + \Sigma_A + \Sigma_Q + \Sigma_C$ combines an isotropic pressure *P*, a contribution from a Newtonian solvent of viscosity η , two reactive stresses

$$\Sigma_{Q} = -K(\nabla \mathbf{Q}) : (\nabla \mathbf{Q}) + 2[\mathbf{Q}\mathbf{H}]^{\mathrm{A}} - \frac{2\xi}{3}\mathbf{H} - 2\xi[\mathbf{Q}\mathbf{H}]^{\mathrm{ST}} + 2\xi\mathbf{Q}\mathrm{Tr}(\mathbf{Q}\mathbf{H}), \qquad (10)$$

$$\boldsymbol{\Sigma}_C = -2a[\mathbf{CB}]^{\mathrm{S}} + 2[\mathbf{CB}]^{\mathrm{A}}, \qquad (11)$$

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and the active stress $\Sigma_A = -\zeta \mathbf{Q}$. The colon in Eq. (10) denotes contraction over the second and third Cartesian indices. In what follows we assume inertialess (creeping) flow and set $\rho = 0$ in Eq. (8).

Having set out this model in which the dynamics of the polymer and liquid crystal are fully coupled, we now consider certain limits in which the coupling between them is diminished. Clearly, setting the constants χ and κ to zero in Eq. (3) eliminates any thermodynamic coupling between **Q** and **C**. It is important to understand, however, that even in this case of zero thermodynamic coupling, **Q** and **C** remain nontrivially coupled in a purely kinematic way: any changes in **C** or **Q** perturb the stress Σ , which in turn perturbs the flow field **v**, which in turn drives both **Q** and **C**. Accordingly, even without any thermodynamic coupling, we find in what follows a rich array of regimes in which the dynamics of **Q** and **C** show strong coupling. We therefore defer the case of true thermodynamic coupling to Sec. VIC.

As already noted, in the limit in which we simply remove the polymeric and active components ($G_C = \kappa = \chi = \zeta = 0$), the model reduces to the Beris-Edwards theory of liquid crystals. Removing instead the liquid-crystalline component ($G_Q = K = \kappa = \chi = 0$) recovers the Johnson-Segalman model of polymeric fluids. These separate models have been studied comprehensively in the earlier literature and we do not consider them further here.

Note finally, and importantly, that in the limit $\tau_C \rightarrow \infty$ our polymer behaves as an elastomer, without any local mechanism for stress relaxation. At zero activity, our model then describes a passive nematic elastomer [38].

III. SIMULATION DETAILS

A. Geometry and boundary conditions

We consider a 2D slab of active viscoelastic matter confined between parallel plates separated by a distance L_y in the y direction and of length L_x in the x direction. At the plates we choose boundary conditions of no slip and no permeation for the fluid velocity **v**,

$$v_{\alpha} = 0$$
 at $y = \{0, L_{\gamma}\} \forall \alpha$,

and zero-gradient boundary conditions for Q and C,

$$\partial_{y} Q_{\alpha\beta} = \partial_{y} C_{\alpha\beta} = 0$$
 at $y = \{0, L_{y}\} \forall \alpha, \beta$.

In the direction x parallel to the plates we adopt periodic boundary conditions for all variables.

The adoption of a zero-gradient boundary condition for the polymeric conformation tensor C is standard practice in the literature on shear banding, though it remains to be justified microscopically. A zero-gradient boundary condition on the nematic order parameter, allowing free rotation of Q at the wall, has been used in some previous studies of active materials [39,40], whereas other studies have adopted anchoring boundary conditions [41]. Recent work comparing both these boundary conditions has shown the essential physics to be qualitatively unchanged by the choice made [5], particularly at large activities where flowing states develop structure on a length scale much smaller than the system size.

Despite this, we attach the following note of caution. In Sec. IV below we calculate the critical threshold activity

required for the onset spontaneous flow, starting from an initial base state of no flow and a perfectly ordered director field. Crucially, we will find this threshold to depend strongly on the orientation of the director in that initial state: States where the director is initially aligned in the x and y directions become separately unstable at activities an order of magnitude apart. That observation in turn has important implications for the boundary condition on \mathbf{Q} , because a boundary condition that favors anchoring of the director parallel (perpendicular) to the walls would be likely to favor an initial condition in which the director is aligned in the x(y) direction, leading to different threshold activities in each case. This suggests that, in some instances, the choice of boundary conditions can have an important effect on the system's behavior. Indeed, recent experiments demonstrate that living liquid crystals, in which conventional liquid crystals are made active by the addition of swimming bacteria, can be influenced by anchoring boundary conditions [42].

On the other hand, the adoption of a zero-gradient boundary condition considerably simplifies our linear stability analysis below, because it permits a spatially homogeneous base state. This in turn means that heterogeneous perturbations to that base state can easily be expressed in a cosine basis. In contrast, anchoring boundary conditions could lead to inhomogeneous base states, complicating the analysis. Accordingly, we work with zero-gradient boundary conditions throughout.

We restrict the fluid velocities to lie in the *x*-*y* plane, setting $v_z = 0$, and assume translational invariance of all variables along *z*. In some calculations we further assume translational invariance along *x*, restricting our states to vary only in one dimension, along *y*. However, we work throughout with 3D tensors for both **Q** and **C**, allowing nonzero components $Q_{\alpha\beta}$ and $C_{\alpha\beta}$ for $\alpha,\beta = x,y,z$. In principle, this could allow the principal axis (director) of **Q** and/or **C** to point out of the *x*-*y* plane of the simulation. Analyzing our numerical results, however, we find in practice that in most of our simulations **Q** and **C** do remain confined to the *x*-*y* plane [43,44]. We therefore expect the key physics to be robust to the dimensionality of the order parameter tensors [45].

B. Units and parameters

We work in units of length such that the gap size $L_y = 1$, of time such that the liquid crystal's local relaxation time $\tau_Q = 1$ and of mass such that its characteristic modulus $G_Q = 1$. The value of the parameter $\eta_Q = G_Q \tau_Q$ that controls the passive nematic viscosity is then also equal to unity. We work in the creeping flow limit of zero Reynolds number, setting $\rho \rightarrow 0$. We use a cell aspect ratio $L_x/L_y = 4$.

To enable a direct comparison with previous numerical studies of active matter without polymer present [5], we fix the solvent viscosity $\eta = 0.567$ throughout. In the **Q** sector of the dynamics we set the isotropic-nematic control parameter $\gamma = 3$ so that the passive liquid crystal lies well within its nematic phase.¹ We set the flow-alignment parameter at $\xi = 0.7$, which lies within the flow-aligning regime.

TABLE I. List of key dimensionless parameters. Hereafter we drop the tildes for clarity.

Symbol	Values explored	Parameter
$\tilde{\zeta} = \frac{\zeta}{G_O}$	$0.001 \rightarrow 10$	extensile activity
$\tilde{\Delta} \equiv \frac{\tilde{K}}{G_O L_y^2}$	$10^{-5} \rightarrow 6.4 \times 10^{-4}$	diffusion constant
$\tilde{\tau}_C \equiv \frac{\tilde{\tau_C}}{\tau_O}$	$10^{-2} ightarrow 10^6 ightarrow \infty$	ratio of relaxation times
$\tilde{G}_C \equiv \frac{\tilde{G}_C}{\tilde{G}_O}$	$\eta_C/\tau_C = 10^{-6} \rightarrow 10^2$	polymer modulus
$\tilde{\chi} \equiv \frac{\chi}{G_Q}$	$0 \leqslant \chi \ll G_Q, G_C$	explicit coupling strength

In the polymer sector we set the slip parameter a = 1 such that the Johnson-Segalman model reduces to the Oldroyd-B model. This eliminates any possibility of a shear-banding instability originating purely from the polymer dynamics. Theoretical studies suggest that this value, which is widely adopted for flexible polymers, may also be reasonable for dense cross-linked filaments [46,47].

With many of the model's parameters having been fixed as just described, there remain only four to be explored numerically: the activity ζ , a diffusivity parameter Δ (defined below) that governs relaxation of spatial gradients in the system, the polymer relaxation time τ_C and its modulus G_C , and (when nonzero) the coupling constant χ . In our chosen units, these are adimensionalized as summarized in Table I. The tildes denote adimensional quantities, but our units have been chosen to make the tildes redundant and we now drop them.

Table I also shows the representative ranges that we explore numerically for these parameters. While these are chosen on the basis of order-of-magnitude estimates at best, they might reasonably describe the recent experiments of Sanchez *et al.* on an extensile kinesin-microtubule mixture akin to a cytoskeletal gel [4], where the level of activity was varied via the concentration of ATP. In Ref. [48], the modulus of a (contractile) actin gel was found to increase by a factor of 10 compared to the passive case in the presence of myosin motor activity, suggesting an upper bound to the dimensionless activity parameter of $\zeta/G_Q \approx 10$.

In our model, the polymer could in principle describe a range of viscoelastic behaviors within the cell, including the effects of the cytosol, which comprises entangled protein filaments and organelles [49]. To encompass this diversity, we vary the polymer relaxation time from $\tau_C = 10^{-2}$, for which the polymeric dynamics are rapid and only contribute extra viscosity to the fluid as a whole, to $\tau_C \rightarrow \infty$, where the polymer effectively acts as an elastomeric solid.

Throughout most of what follows we assume a constant value of $\eta_C \equiv G_C \tau_C$, which controls the polymer viscosity, setting $\eta_C = 1$. To maintain this between runs in which the polymer relaxation time τ_C changes, the polymer elastic modulus G_C must change in inverse proportion to τ_C . This assumption of a constant polymer viscosity, however, is relaxed in those sections where we consider the "true" elastomeric limit $\tau_C \rightarrow \infty$ (at a fixed value of G_C) of an active nematic in a cross-linked polymeric background.

The crossover length scale at which elastic distortions in a passive nematic compete with its bulk free energy is $\ell_Q \equiv$

¹This value in fact corresponds to the spinodal stability limit of the isotropic phase.

 $\sqrt{K/G_Q}$. Recasting the **Q** constitutive equation (6) in this notation, the diffusive term in the molecular field then becomes $\frac{\ell_Q^2}{\tau_Q} \nabla^2 \mathbf{Q}$. The diffusive JS model prescribes an analogous term in the polymer sector, with coefficient ℓ_C^2/τ_C [33]. In most of our numerical runs we assume these to be equal, setting

$$\frac{K}{\eta_O} \equiv \frac{\ell_Q^2}{\tau_O} = \frac{\ell_C^2}{\tau_C} \equiv \Delta.$$
(12)

Note that taking the elastomeric limit $\tau_C \to \infty$ at a fixed value of the diffusivity parameter Δ means taking $l_C^2 \to \infty$ in tandem. In some places below we instead consider fixed $l_C = l_Q$. Taking the elastomeric limit in that case then eliminates the diffusive term $\Delta_C = l_C^2/\tau_C$ from the polymer dynamics; setting $l_C = l_Q$ is then the same as setting $l_C = 0$ in this limit. However, many of our 2D simulations then show an unphysical instability that leads to structure on the scale of the numerical grid. The retention of finite $\Delta_C = \Delta$ as $\tau_C \to \infty$ avoids this problem; we discuss the issue further in Sec. VI B.

Values for Δ are then selected so that the associated length scale l_Q obeys $\Delta x < \ell_Q \ll L_y$, where Δx is the numerical grid size (for which $L_x/1024$ or $L_x/2048$ is a reasonable lower bound, constrained by a maximum feasible number of grid points $N_x = 1024$ or 2048). In this way, the spatial structures predicted by the model are well resolved on the scale of the numerical mesh and also fit comfortably within the simulation box. The velocity correlation length for kinesin microtubule mixtures has been estimated at $l_v \approx 100 \ \mu m$ [4] and if we assume that our length scale ℓ_Q is an order of magnitude smaller than this [45], the range of values of l_Q that we explore numerically would imply a channel width in the range 400 μ m–3 mm, as might describe a typical microfluidic device.

As noted above, throughout most of what follows we disable explicit thermodynamic coupling between **Q** and **C** by setting $\chi = 0$ and $\kappa = 0$. The only remaining coupling between **Q** and **C** is then purely kinematic, caused by the two components sharing a common fluid velocity **v**. In Sec. VIC we move finally to address the case of explicit thermodynamic coupling with $\chi \neq 0$.

C. Numerical details

In our numerical scheme we evaluate spatially local terms in both constitutive equations using an explicit Euler timestepping scheme. To solve the Stokes equation we use a stream function formulation, with a hybrid numerical scheme in which gradient terms in the x and y directions are computed using Fourier and Crank-Nicolson methods, respectively [50,51]. Spatially diffusive terms in the constitutive equations are likewise solved using this hybrid approach. Advective terms are computed using third-order upwind differencing.

We performed the following checks of our numerics. For each run we verified the results to be robust against halving either the spatial mesh size or the time step. Note, however, that because of the erratic (and quite possibly chaotic) nature of many of the dynamical states explored, converging individual trajectories is not a realistic prospect. We instead performed convergence tests on time-averaged quantities and on the overall characteristics of states in the phase diagram. Finally, in our nonlinear runs we have carefully checked the behavior in the early stages of instability for consistency with the growth rates found from our linear stability analysis.

IV. LINEAR STABILITY ANALYSIS

In this section we perform a linear stability analysis of an initially homogeneous, nonflowing base state to determine the regions of parameter space in which such a state is linearly unstable to the formation of a heterogeneous state with spontaneous flows. Doing so enables us to understand the key instabilities present in the model and to form a helpful roadmap in selecting parameter values for the much more computationally expensive nonlinear runs in later sections. We set the explicit thermodynamic couplings to zero throughout this section: $\kappa = \chi = 0$.

A. General procedure

We consider an initially homogeneous, quiescent, uniaxial base state with

$$\overline{Q}_{\alpha\beta} = q(n_{\alpha}n_{\beta} - \delta_{\alpha\beta}), \qquad (13a)$$

$$\overline{C}_{\alpha\beta} = \delta_{\alpha\beta},\tag{13b}$$

$$\partial_{\alpha}\overline{v}_{\beta} = 0.$$
 (13c)

Here q is the magnitude of the nematic order parameter and n_{α} its director. We consider two alternative choices for the initial director field: $\hat{\mathbf{n}} = (1,0,0)$ and (0,1,0), oriented parallel and perpendicular to the walls, respectively. Note that, although we have specified the $Q_{\alpha\beta}$ of this initial base state to be uniaxial, our calculations for departures from the base state do not impose that restriction. The polymer is initially undeformed, consistent with the absence of velocity gradients; with our boundary conditions this absence also implies $\overline{v}_{\alpha} = 0$ in the base state such that the fluid is initially at rest.

We now examine the linear stability of this base state to perturbations that are heterogeneous in one spatial dimension, with wave vector in the flow gradient direction y. Within this 1D assumption, the condition of fluid incompressibility further demands that the only element of the velocity gradient tensor that might become nonzero is $\dot{\gamma} \equiv \partial_y v_x$. To simplify the notation, we collect all the relevant variables into the quantity $\boldsymbol{\phi} = (\mathbf{Q}, \mathbf{C}, \dot{\gamma})$. To the base state $\boldsymbol{\phi} = (\mathbf{Q}, \mathbf{C}, \dot{\gamma})$ prescribed in Eq. (13) above, we then add Fourier-mode perturbations

$$\boldsymbol{\phi} = \overline{\boldsymbol{\phi}} + \sum_{k} \boldsymbol{\phi}^{k}(t) \cos(ky), \qquad (14)$$

with small amplitudes $\phi^k = (\mathbf{Q}^k, \mathbf{C}^k, \dot{\gamma}^k)$. Our boundary conditions at $y = 0, L_y$ require that the wave vector k takes discrete values $k = \pi m/L_y$, with integer m.

Substituting Eq. (14) into the model equations derived in Sec. II and expanding in powers of the mode amplitudes gives, at first order, a linearized equation set for the dynamics of the perturbations:

$$\partial_t \mathbf{p}^k = \mathbf{M}^k \cdot \mathbf{p}^k, \tag{15}$$

where $\mathbf{p}^k \equiv (Q_{xx}^k, Q_{xy}^k, Q_{yy}^k, C_{xx}^k, C_{xy}^k, C_{yy}^k)^T$. Note that \mathbf{p}^k contains only a subset of the full list of variables originally specified in $\boldsymbol{\phi}^k$. The reasons for this are twofold. First, the

Stokes equation requires $\nabla \cdot \Sigma = 0$, with Σ the total stress tensor. This enables us to express $\dot{\gamma}^k$ directly in terms of \mathbf{Q}^k and \mathbf{C}^k as

$$\dot{\gamma}^{k} = -rac{1}{\eta} ig(oldsymbol{\Sigma}_{A}^{k} + oldsymbol{\Sigma}_{Q}^{k} + oldsymbol{\Sigma}_{C}^{k} ig)_{xy}$$

in an obvious notation. This eliminates $\dot{\gamma}^k$ as a dynamical variable: Physically, the shear rate is enslaved to the viscoelastic stress components by the requirement of instantaneous force balance in the limit of creeping flow.

Second, we find that the set of components $Q_{\alpha z}^{k}$ and $C_{\alpha z}^{k}$ evolve independently from the *xx*, *xy*, and *yy* components contained in **p**; this *z*-related set furthermore always has negative eigenvalues, rendering them stable. Accordingly, the only dynamical variables relevant to us are those listed in **p** above.

The eigenvalues of the matrix \mathbf{M}^k then determine whether the initial base state is stable or unstable to the growth of perturbations. Any eigenvalue with a positive real part corresponds to an unstable mode that grows exponentially (in this linear regime), taking the system away from the initial homogeneous nonflowing base state and towards a heterogeneous state with a spontaneous flow.

In what follows we seek a threshold value of the activity at which the base state first becomes linearly unstable, with the values of the other model parameters held fixed. Sweeping the activity level upward, the critical threshold ζ_c at which instability first sets in is the value at which

$$\operatorname{Re}(\omega^+) = 0, \tag{16}$$

where ω^+ is the eigenvalue with the largest real part.

It is possible to show that the only nontrivial (potentially unstable) modes of \mathbf{M}^k have eigenvalues ω^{\pm} given by

$$2\omega^{\pm} = -b \pm \sqrt{b^2 - 4c},$$
 (17)

where

$$b = \frac{a^2 G_C}{\eta} + \frac{\Lambda}{\eta} \left(2G_Q \Lambda \ell_Q^2 k^2 - \zeta \right) + \frac{1 + \ell_C^2 k^2}{\tau_C} + \frac{\ell_Q^2 k^2}{\tau_Q}$$
(18)

and

$$c = \frac{\ell_Q^2 k^2}{\tau_Q} \frac{a^2 G_C}{\eta} + \frac{\Lambda}{\eta} \frac{1 + \ell_C^2 k^2}{\tau_C} \left(2G_Q \ell_Q^2 k^2 \Lambda - \zeta \right) + \frac{\ell_Q^2 k^2}{\tau_Q} \frac{1 + \ell_C^2 k^2}{\tau_C},$$
(19)

with

$$\Lambda = \begin{cases} (5\xi - 3)/12 & \text{for } \hat{\mathbf{n}} = (1,0,0) \\ (5\xi + 3)/12 & \text{for } \hat{\mathbf{n}} = (0,1,0). \end{cases}$$
(20)

For each value of Λ , then, two different modes of instability are possible as determined by Eq. (17). We term these the viscous and elastomeric instabilities and explore them in the next two sections, respectively. For each such mode of instability, the activity threshold depends on Λ . This in turn depends on whether the initial director obeys $\hat{\mathbf{n}} = (0,1,0)$, for which the growing mode involves a bend deformation, or obeys $\hat{\mathbf{n}} = (1,0,0)$, for which it involves splay.

B. Viscous instability

For the first mode of instability, the discriminant $b^2 - 4c$ in Eq. (17) remains positive and the eigenvalues of \mathbf{M}^k are purely real. The condition $\operatorname{Re}(\omega^+) > 0$ for the onset of instability then simplifies to the condition c < 0. Predividing c by the final term on the right-hand side of Eq. (19) and multiplying by η then gives the conditions for instability as

$$\frac{a^2 G_C \tau_C}{1 + \ell_C^2 k^2} + \frac{\Lambda}{\ell_Q^2 k^2 / \tau_Q} \left(2 G_Q \ell_Q^2 k^2 \Lambda - \zeta \right) + \eta < 0.$$
(21)

We identify the four terms on the left-hand side as the wavevector-dependent zero-shear viscosities associated with the polymer, the passive nematic stress, the active nematic stress, and the solvent, respectively. We can therefore rewrite this criterion as

$$\eta_{\text{total}}^{k} = \eta_{C}^{k} + \eta_{Q,\text{passive}}^{k} + \eta_{Q,\text{active}}^{k} + \eta^{k} < 0, \qquad (22)$$

where in fact $\eta_{Q,\text{passive}}^k$ and η^k are independent of k. Each term in Eq. (22) is positive, apart from the term $\eta_{Q,\text{active}}^k$. A simplified analysis, which casts the full tensorial nematic order parameter **Q** in a simplified form in terms of a director **n̂** with a fixed degree of nematic ordering [52], then suggests that

$$\eta_{Q,\text{active}}^{k} = -\zeta \times \frac{\Lambda}{\ell_{Q}^{2}k^{2}/\tau_{Q}} = \frac{\partial \Sigma_{Q,\text{active}}^{k}}{\partial n^{k}} \times \frac{\partial n^{k}}{\partial \dot{\gamma}^{k}}, \quad (23)$$

where the multiplication symbols serve to emphasize the correspondence of terms between the two expressions. In this equation, an initial base state director orientation along $\hat{\mathbf{x}}$ is linked to a director perturbation δn in the y direction, corresponding to a splay instability, whereas an initial base state director along $\hat{\mathbf{y}}$ is linked to a perturbation δn in the $\hat{\mathbf{x}}$ direction, corresponding to a bend instability. Recall that the perturbation wave vector **k** is in the $\hat{\mathbf{y}}$ direction in all these 1D calculations. [Note that all other analytical and numerical results employ the full **Q** formulation, with the exception of Eq. (27), where we make the analogous argument for the elastomeric instability.] Within Eqs. (22) and (23) we recognize the mechanism that drives this instability as follows: A perturbation in the shear rate causes a perturbation in the director orientation, which in turn causes a perturbation in the active contribution to the stress field, which must provide a counterbalancing contribution from the other stress components to maintain force balance. The two shear rate perturbations in this loop are in the same sense, signifying positive feedback. Rearranging Eq. (21), we find the critical activity for the onset of instability to be

$$\begin{aligned} \xi_c^{\text{visc}} &= \frac{\ell_Q^2 k^2 / \tau_Q}{\Lambda} \bigg[\frac{a^2 G_C \tau_C}{1 + \ell_C^2 k^2} + 2G_Q \tau_Q \Lambda^2 + \eta \bigg] \\ &= \frac{r_Q}{\Lambda} \big[\eta_C^k + \eta_{Q,\text{passive}}^k + \eta^k \big], \end{aligned}$$
(24)

where we recognize $r_Q \equiv \ell_Q^2 k^2 / \tau_Q$ as the on-diagonal rate of relaxation of a perturbation to the director field at wavelength *k*.

. . .

For values of the activity just above threshold, the perturbations grow very slowly. They furthermore do so without any oscillatory component because the eigenvalues are real in this case. Relative to this slow growth, the fluid's intrinsic relaxation time scales τ_C and τ_Q become infinitely fast as one approaches the threshold. Accordingly, the fluid's resistance to this instability appears only via its zero-frequency viscosities η_C and η_Q : The relaxation times τ_C and τ_Q cannot appear in Eq. (22) independently of these viscosities. The effect of the polymer is therefore simply to increase the fluid's overall viscosity, delaying the onset of instability. The dependence of the threshold on $\eta_C = G_C \tau_C$ is thus independent of whether τ_C is varied at fixed G_C or G_C at fixed τ_C . Accordingly, we call this mode the viscous instability in what follows.

In view of the relatively trivial role of the polymer in determining the onset of this viscous instability, we expect the result just obtained to match the one in the earlier literature for a purely nematic active fluid, without polymer. Indeed, substituting $K \equiv \ell_Q^2 G_Q$ recovers the criterion in the form originally derived in Refs. [10,40], with an additional term to account for the extra viscosity contributed by the polymer.

In the limit of infinite system size, $k \to 0$ in Eq. (24), we further recover the prediction $\zeta_c^{\text{visc}} \to 0$, implying that a bulk active nematic will be unstable for any level of activity, however small [10,12]. One important prediction of the present work is that this generic instability of a bulk active nematic persists even with polymer present.

C. Elastomeric instability

An alternative mode of instability arises when the discriminant $b^2 - 4c$ in Eq. (17) is negative and the eigenvalues of \mathbf{M}^k have an imaginary part, suggesting a Hopf bifurcation with oscillatory dynamics [53]. The criterion $\operatorname{Re}(\omega^+) = 0$ for the onset of instability then simplifies to the condition

$$b = \frac{a^2 G_C}{\eta} + \frac{\Lambda}{\eta} \left(2G_Q \Lambda \ell_Q^2 k^2 - \zeta \right) + \frac{1 + \ell_C^2 k^2}{\tau_C} + \frac{\ell_Q^2 k^2}{\tau_Q} < 0.$$
(25)

At a given wave vector k, we identity the five terms in b as being respectively proportional to (i) the rate of change of polymer stress during an impulsive strain, (ii) the rate of change of passive nematic stress during an impulsive strain, (iii) the rate of change of active nematic stress during an impulsive strain, (iii) the rate of change of active nematic stress during an impulsive strain, (iv) the on-diagonal rate of relaxation $r_c \equiv (1 + \ell_c^2 k^2)/\tau_c$ for a perturbation in the polymer conformation, and (v) the on-diagonal rate of relaxation r_Q for a perturbation in the director field.

Therefore, we write the criterion for the onset of instability as

$$\frac{1}{\eta}\frac{d\Sigma_{C}^{k}}{d\gamma} + \frac{1}{\eta}\frac{d\Sigma_{Q,\text{passive}}^{k}}{d\gamma} + \frac{1}{\eta}\frac{d\Sigma_{Q,\text{active}}^{k}}{d\gamma} + r_{C} + r_{Q} < 0.$$
(26)

Among the terms on the left-hand side, each is always positive apart from the third. Following the approach described previously for the viscous instability, in which we rewrite the derivative in terms of the director field *n*, the third term reads

$$\frac{\partial \Sigma_{Q,\text{active}}^{k}}{\partial n^{k}} \frac{\partial n^{k}}{\partial \gamma^{k}}.$$
(27)

We recognize the mechanism that drives this instability as follows: A perturbation in the shear strain field causes a perturbation in the director orientation, which in turn causes a perturbation in the shear stress field, which in turn must cause a perturbation in the strain field to maintain force balance. The two shear strain perturbations in this loop are in the same sense, signifying positive feedback. Indeed, this is essentially the same mechanism as for the viscous instability above, but with the strain rate replaced by the strain. Accordingly, we call this mode the elastomeric instability.

Rearranging Eq. (25), we get an expression for the critical activity for the onset of instability as

$$\begin{aligned} \zeta_c^{\text{elast}} &= \frac{\eta}{\Lambda} \Bigg[\frac{a^2 G_C}{\eta} + 2G_Q \frac{\Lambda^2}{\eta} l_Q^2 k^2 + \frac{1 + l_C^2 k^2}{\tau_C} + \frac{l_Q^2 k^2}{\tau_Q} \Bigg] \\ &= \frac{1}{\Lambda} \Bigg[\frac{d\Sigma_C^k}{d\gamma} + \frac{d\Sigma_{Q,\text{passive}}^k}{d\gamma} + \eta (r_C + r_Q) \Bigg]. \end{aligned}$$
(28)

In contrast to the viscous instability discussed above, at the threshold of this new instability the perturbations grow subject to a superposed oscillatory component that has a time period prescribed by the imaginary part of the eigenvalue $\tau_{\omega} = 2\pi/\text{Im}(\omega^+)$. The presence of this time scale means that the relative value of the polymer's relaxation time is relevant to this instability, in contrast to its unimportance to the viscous one: τ_C enters Eq. (28) directly rather than merely via the viscosity $\tau_C G_C$. Put differently, the polymer's *viscoelasticity* is important to this new instability, in contrast to the viscous instability, which depends only on its viscosity. Two distinct thresholds again arise for the elastomeric instability according to whether the deformation corresponds to a bend or a splay in the director field, which is in turn determined by whether the initial director lies in the *y* or *x* direction as set by Λ [see Eq. (20)].

D. Viscous-elastomeric crossover

As described in the previous two sections, viscoelastic active matter is predicted to exhibit two distinct modes of instability: a viscous mode and an elastomeric one. The activity thresholds for these two modes are compared in the phase diagrams of Figs. 1 and 2.

As can be seen, the thresholds for the two instabilities display qualitatively different dependences on the polymer relaxation time τ_C and on the system size. Indeed, taking the elastomeric limit $\tau_C \to \infty$ for any fixed G_C and fixed $\ell_C = \ell_Q$ raises indefinitely the threshold for the viscous instability: $\zeta_c^{\text{visc}} \to \infty$. The fluid's viscosity approaches infinity in this limit, switching off the viscous instability entirely. In contrast, the threshold for the onset of the elastomeric instability remains finite even in the purely elastomeric limit $\tau_C \to \infty$:

$$\lim_{\substack{\tau_C \to \infty, \\ k \to 0}} \zeta_c^{\text{elast}} = \frac{a^2 G_C}{\Lambda}$$
(29)

(where we have also taken the infinite system size limit for clarity of expression). Crucially, then, this second instability can arise even in the limit of an elastomeric solid, because it involves perturbations in strain rather than strain rate.

Now consider instead fixing τ_C, G_C , so that the polymer viscosity remains finite, and taking the limit $\ell_C = \ell_Q \rightarrow 0$. This is equivalent to taking the limit of infinite system size



FIG. 1. Phase diagram showing the threshold activity for the onset of the splay mode of the viscous (solid lines) and elastomeric (dashed lines) instabilities, for $k = \pi/L_y$. The unstable region is to the right of the threshold curve in each case. The dotted lines enclose the regime in which the eigenvalues have an imaginary contribution, signifying oscillatory dynamics. Sets of curves from left to right have $\ell_C^2 = \ell_Q^2 = 10^n$ with n = -5, -4, -3, -2. For all curves the polymer modulus $G_C = 0.1$.

 $L_y \rightarrow \infty$, for which the threshold for the viscous instability tends to zero: For a bulk system, any level of activity, however small, triggers this instability. In contrast, the activity threshold for the elastic instability remains finite even in this bulk limit. These two facts can be seen in Fig. 1.

The physical origin of this important difference in systemsize dependence can be understood by comparing the driving terms for the two modes. The term $\partial n/\partial \dot{\gamma}$ in the viscous criterion [recall Eq. (23)] scales as $1/k^2$, because an infinitesimal shear rate perturbation can cause finite rotation of the director field in the limit of $k \rightarrow 0$. (This stems from the Goldstone



FIG. 2. Phase diagram showing the threshold activity for the onset of the viscous (solid lines) and elastomeric (dashed lines) instabilities, for $k = \pi/L_y$. Leftmost (black lines) and rightmost (red lines) curves correspond to the bend and splay modes of instability, respectively. The unstable region is to the right of the threshold curve in each case. The dotted lines enclose the regime in which the eigenvalues have an imaginary contribution, signifying oscillatory dynamics. For all curves $G_C = 0.1$ and $l_C^2 = l_Q^2 = 10^{-4}$.

mode associated with the spontaneously broken rotational symmetry. A nematic phase exerts no restoring force against global rotations of a state of uniform **Q**, implying by continuity that the force likewise vanishes in the limit $k \rightarrow 0$ [54].) In contrast, the term $\partial n/\partial \gamma$ in the elastic criterion [recall Eq. (27)] is finite at low k, because small strain perturbations cause only small changes in director, even in the limit of zero wave vector.

With these remarks in mind, we now see that dominance of the viscous instability over the elastomeric one (or vice versa) will depend on the ratio τ_C/τ_Q of the polymer and nematic relaxation times and also on the system size, measured in relation to the lengths l_Q, l_C (as set by the gradient terms in the equations of motion). For any fixed system size, below a critical value of the polymer relaxation time $\tau_C = \tau_C^*$ the viscous instability threshold ζ_c^{visc} lies below the elastic one: The viscous instability arises first on increasing the activity and ζ_c^{visc} sets the stability criterion for our base state. In contrast, for $\tau_C > \tau_C^*$ the elastomeric instability arises before the viscous one and ζ_c^{elast} sets the relevant criterion.

This crossover value τ_C^* can easily be shown to obey

$$\tau_C^* \approx \left(1 + \ell_C^2 k^2\right) \frac{\tau_Q}{\ell_Q^2 k^2},\tag{30}$$

which for large system sizes simplifies to $\tau_c^* \approx \tau_Q / \ell_Q^2 k^2$. This form can be understood as follows. The quantity $\tau_Q / \ell_Q^2 k^2$ is the orientational relaxation time for long-wavelength distortions in the nematic phase, which diverges as $k \to 0$. (Recall that this is a Goldstone mode, with no penalty for global rotations in the director field.) For polymer relaxation times faster than this $\tau_C < \tau_C^*$, the polymer behaves in a viscous way relative to the nematic mode and the viscous instability dominates. In the opposite case the polymer behaves elastically and the elastomeric instability dominates.

In the limit of infinite system size the crossover value diverges: $\tau_c^* \to \infty$. In this bulk limit the viscous instability is triggered first (at zero activity threshold) for all values of τ_c , whereas the threshold of the elastomeric instability remains finite even for a bulk sample. As noted in the Introduction, however, many biological environments are strongly confined. This motivates the use of a finite box size in our simulations and a correspondingly careful exploration of both the viscous and elastomeric instabilities. For our typical choice of parameter values given in Table I, the crossover between the two instabilities occurs at $\tau_c^* \approx \eta_Q/Kk^2 = O(10^3 \tau_Q)$.

V. NONLINEAR DYNAMICS (ONE DIMENSION)

In the previous section we derived the activity threshold for linear instability to a state of spontaneous flow in an active nematic with added polymer. We now use those results as a roadmap to explore the model's full nonlinear dynamics. We shall restrict ourselves in this section to 1D calculations, with heterogeneity in the *y* direction only, deferring a study of 2D effects to Sec. VI below. In each run we initialize the system in the homogeneous base state given by Eq. (13), with the director $\hat{\mathbf{n}}$ aligned in the *x* direction. Accordingly, any instabilities that arise below originate from splay deformations. We subject this base state to small added perturbations at the start of the run of the form $\delta \tilde{\gamma} = \sum_{m=1}^{N} \delta A_m \cos(\pi m y/L_y)$, where $\delta \sim O(10^{-10})$, A_m is a random number drawn from a uniform distribution on [-1,1], and N = 32. In this way, the early-time regime of these nonlinear runs acts as an independent cross-check of the linear stability calculations in the previous section.

Our model has three competing local relaxation times: the liquid-crystal relaxation time τ_Q , the polymer relaxation time τ_C , and the active forcing time scale $\tau_a = \eta/\zeta$, which was identified in Ref. [55] by balancing the active modulus ζ against the solvent viscosity η . As discussed previously, the first of these, τ_Q , is our unit of time. We now examine the competition between the second and third time scales by producing phase diagrams in the (ζ, τ_C) plane. Related to τ_a is the active time scale $\bar{\tau}_a = \eta/(\zeta - \zeta_c)$, with the denominator shifted to account for the fact that in a finite system the critical threshold for the onset of activity is nonzero. (Note that $\bar{\tau}_a \to \tau_a$ as $L_y \to \infty$ or $\zeta \to \infty$.)

Whenever the active time scale is shorter than the two relaxation times, we can expect oscillatory dynamics [55]. We will confirm this expectation: first (in Sec. V A) for a material of fixed polymer viscosity $\eta_C = 1$, where we find the dominant period of oscillation τ_{osc} to be set by τ_C , and then (in Sec. V B) for an elastomeric material with $\tau_C \rightarrow \infty$, where instead we find $\tau_{osc} \propto \bar{\tau}_a$. In a biological context, oscillatory states have been observed in fibroblast cells [56] with a period of oscillation proportional to the myosin motor activity. Similar states have also been studied in *Drosophila* embryos [57], where oscillations are thought to play a key role in cell shape formation (morphogenesis).

A. Phase diagram: Fixed viscosity

We begin by considering the case of a fixed polymer viscosity $\eta_C = G_C \tau_C = 1$. (Recall that the passive nematic viscosity $\eta_Q = 1$ by our choice of units and the solvent viscosity $\eta = 0.567$ for consistency with Ref. [5].) Between runs we vary the polymer relaxation time τ_C and in tandem therefore also vary $G_C = 1/\tau_C$. This will allow us to confirm the direct role of the polymer relaxation time τ_C in the dynamics, while removing any potential complications arising from variations in the overall viscosity. In varying τ_C in this section we choose to keep $\ell_C = \ell_Q$ fixed. Note that this effectively eliminates the diffusivity $\Delta_C = \ell_C^2/\tau_C$ from the polymer dynamics in the elastomeric limit $\tau_C \to \infty$. The predictions of our linear stability analysis in this case are shown in the (ζ, τ_C) plane in Fig. 3, with four sets of curves corresponding to different values of $\ell_C = \ell_Q$.

The results of our nonlinear simulations at fixed $\ell_Q = \ell_C = 0.004$ are shown in Fig. 4, with the linear stability thresholds also repeated on the same axes for comparison. The green shaded region indicates when oscillatory instabilities are expected from the linear stability analysis at any point in the discrete spectrum of modes $k = m\pi/L_y$ (cf. Fig. 3 where oscillatory instabilities are also marked with shaded regions, but where we instead fix $k = \pi/L_y$ and plot multiple values of $\ell_Q = \ell_C$). As can be seen, the nonlinear results are consistent with those of the linear stability analysis. For values of the activity below $\zeta_c = \min(\zeta_c^{\text{visc}}, \zeta_c^{\text{elast}})$ we observe quiescent states (black crosses) in our nonlinear simulations, consistent with the analytical prediction of a stable base state



FIG. 3. Phase diagram showing the threshold activity for the onset of the viscous (solid lines) and elastomeric (dashed lines) instabilities in the splay mode, for $k = \pi/L_y$. The unstable region is to the right of the threshold curve in each case. The dotted curves enclose the regime in which the eigenvalues have an imaginary contribution, signifying oscillatory dynamics (shaded regions are both oscillatory and unstable for $k = \pi/L_y$). Sets of curves from left to right have $l_C^2 = l_Q^2 = 10^{-5,-4,-3,-2}$. For all curves the polymer viscosity $\eta_C = G_C \tau_C = 1$.

in this regime. For values of the activity above ζ_c we observe spontaneously flowing shear-banded states (black circles).

These spontaneously flowing states may be categorized according to whether they approach a steady state at long times



FIG. 4. Phase diagram showing the competing effects of the activity ζ and the polymer's viscoelastic relaxation time scale τ_C , for fully nonlinear runs in one dimension with fixed polymer viscosity $\eta_C = 1$. Marked are the critical activities ζ_c^{visc} [Eq. (22)] (black line) and ζ_c^{elast} [Eq. (28)] (red line) predicted by the linear stability analysis. Nonlinear states are denoted by symbols: quiescent (crosses), static bands (open circles), oscillating bands (striped circles), or oscillating and flipping bands (closed circles) where the flow switches direction. In the green shaded regions our linear stability analysis predicts unstable oscillatory growth, at some wave vector *k*. Examples of the states highlighted with the triangle and square are given in Fig. 5 (left and right, respectively). Oscillations are observed for $\tau_C/\bar{\tau}_a > 200$ (blue curve). The parameters are $\eta_C = 1$ and $\ell_Q = \ell_C = 0.004$.



FIG. 5. Space-time plots corresponding to the blue triangle and red square in the phase diagram of Fig. 4. The color scale represents $\eta \dot{\gamma}$ (top row) and Σ_{xy}^{C} (bottom row). The left column shows the state with oscillating interface ($\zeta = 3.2$ and $\tau_{C} = 10^{3}$) and the right column the state with oscillating interface and switching flow direction ($\zeta = 3.2$ and $\tau_{C} = 10^{6}$).

or instead oscillate. In particular, for activity values exceeding the threshold for the viscous instability, but below that for the elastomeric instability, we observe steady shear-banded states (open circles), as seen in earlier studies of the model in the absence of polymer [5]. In contrast, for values of the activity that also exceed the threshold for the elastomeric instability and roughly correspond to $\tau_C/\bar{\tau}_a > 200$ (blue line in Fig. 4), we find time-dependent states in which the shear bands oscillate with a period set by the viscoelastic relaxation time scale τ_C . No counterpart to these oscillatory states was observed in the earlier study without polymer [5]. This confirms our view that the polymer relaxation time is integral to their dynamics.

The separation of time scales (large τ_C/τ_a) required to see these oscillations is reminiscent of earlier work addressing rheological chaos [58]. Intuitively, if the active time scale $\tau_a = \eta/\zeta$ is sufficiently long, then both polymer and liquid crystal can relax any activity-driven deformation and the resulting shear-banded steady state is time independent. Conversely, if the polymer cannot relax, the activity-induced stress quickly enough, the polymer dynamics lag behind, resulting in oscillations. This is also somewhat analogous to the mechanism described in Ref. [55], although in that case the coupling was to a slowly diffusing concentration field rather than a slowly relaxing polymer.

To gain further insight into these oscillatory shear-banded states, we now explore in detail two particular examples denoted, respectively, by the blue triangle and red square in the phase diagram of Fig. 4.

(i) Oscillatory interface with a fixed flow direction. The first example, denoted by the blue triangle in Fig. 4, corresponds to a time-dependent state in which the position of the interface between two shear bands oscillates as a function of time. See Fig. 5 for a space-time plot of the shear rate $\dot{\gamma}$ (top left) and of the polymer shear stress Σ_{xy}^{C} (bottom left). The time period is set by the polymer relaxation time τ_{C} , making it clear that these oscillations are governed by the viscoelastic dynamics of the polymer. As the interface position deviates from the center, large polymeric stresses are seen to develop in the narrower band until eventually the interface turns around and returns in the opposite direction, relaxing the polymer somewhat. The total shear stress and integrated throughput $\Psi = \int_0^{L_y} v_x dy$ also have time-dependent signals (not shown) with a period set by τ_C .

In this example the sign of the throughput, i.e., the overall direction of the flow, remains constant in time. This can be seen by noting that the region of negative shear rate remains always below the one of positive shear rate in Fig. 5 (top left), meaning that the velocity profile is always overall nose shaped and pointing to the left. (Note that a state with sustained throughput to the right could equally well have developed had we made small changes in the initial conditions.)

(ii) Oscillatory interface with a switching flow direction. The second example is denoted by the red square in Fig. 4, with space-time plots of $\dot{\gamma}$ and \sum_{xy}^{C} in Fig. 5 (right). Here two distinct kinds of oscillation are seen in the course of any run. On a slow time scale $\approx 2\tau_C$ the flow periodically reverses direction, as evidenced by the up-down switching between the bands of positive and negative shear rate. Another, more rapid, oscillation is evident: The throughput direction remains the same, but the position of the interface fluctuates with a mechanism resembling the one discussed in the previous example above.

As already noted above, in this section we have kept the values of $\ell_C = \ell_Q$ fixed as the polymer relaxation time changes. Had we instead chosen to keep $\ell_Q^2/\tau_Q = \ell_C^2/\tau_C \equiv \Delta$ fixed as τ_C varied, the initial linear instability would not be oscillatory. While a oscillatory linear instability does not necessarily imply oscillatory nonlinear dynamics, our results in this section demonstrate a reasonable correlation between the two [compare the green region in Fig. 4 with oscillatory banded states (shaded and closed circles)].

B. Spontaneous flow in elastomeric solids

The case of an active elastomeric solid can be approached within our formalism by more than one route, yielding slightly different limiting descriptions that may be appropriate to different physical cases. One choice concerns the way in which the infinite viscosity limit is taken; a second and independent



FIG. 6. Phase diagram showing the threshold activity for the onset of the viscous (solid lines) and elastomeric (dashed lines) instabilities in the splay mode, for $k = \pi/L_y$. The unstable region is to the right of the threshold curve in each case. Sets of curves from left to right have $\Delta_C = \Delta_Q = 10^{-5, -4, -3, -2}$. For all curves the polymer modulus $G_C = 0.1$.

choice concerns the behavior of the polymer stress diffusivity Δ_C . In the previous section we studied the dependence of the model's behavior on the polymer relaxation time τ_C at fixed polymer viscosity $\eta_C = G_C \tau_C = 1$. We also fixed the values of the microscopic length scales $\ell_C = \ell_Q$, in relation to the macroscopic system size. In the regime of large polymer relaxation times ($\tau_C \simeq 10^6$) we found sustained oscillations in the system's long-time response. Such large values of τ_C correspond for all practical purposes to an elastomeric solid, but with a tiny modulus G_C to maintain the overall constraint of fixed polymer viscosity $\eta_C = G_C \tau_C = 1$.

In this section we consider instead taking the "true" limit of an elastomeric solid $\tau_C \to \infty$ at a fixed value of the polymer modulus G_C so that the polymer viscosity $\eta_C = G_C \tau_C$ now also diverges as $\tau_C \to \infty$. We also fix the value of the diffusivity parameter $\ell_Q^2/\tau_Q = \ell_C^2/\tau_C = \Delta$. In this way, taking the limit $\tau_C \to \infty$ means taking the limit $l_C \to \infty$ also.

Our choice of nonzero polymer stress diffusivity $\Delta_C = \Delta > 0$ in the elastomeric limit is not obvious, since physically there is no reason for l_C to diverge in that limit. However, we have found that in our 2D nonlinear work setting $\Delta_C = 0$ generally leads to a short-scale numerical instability unless prohibitively small time- and spatial-step sizes are used. We explore this issue further in Sec. VIB, but even there set $\Delta_C = \Delta$ to retain direct comparability of parameters between the 1D and 2D numerics.

The predictions of our linear stability analysis in this case are shown in Fig. 6 in the (ζ, τ_C) plane, for fixed $G_C = 0.1$ and various fixed values of Δ . In this figure, $\tau_C \rightarrow \infty$ is approached by taking the ordinate to infinity. Note how Fig. 6, which has fixed G_C and Δ_C , differs from Fig. 1, which had fixed G_C and ℓ_C . In particular, the viscous instability is not eliminated when the elastomeric limit is taken at fixed Δ_C in Fig. 6, apparently because the polymer stress can still redistribute itself spatially by diffusion despite having a divergent local viscoelastic relaxation time τ_C . (The latter ensures that the volume-averaged polymer stress cannot decay;



FIG. 7. Phase diagram showing the threshold activity for the onset of the viscous (solid lines) and elastomeric (dashed lines) instabilities in the splay mode, for $k = \pi/L_y$. The unstable region is to the right of the threshold curve in each case. The dotted line shows the locus where $b^2 - 4c = 0$; when $\ell_Q^2/\tau_Q = \ell_C^2/\tau_C = \Delta$ this quantity never becomes negative and therefore the linear instability is never oscillatory. Sets of curves from left to right have $\Delta_C = \Delta_Q = 10^{-5, -4, -3, -2}$. For all curves the polymer relaxation time $\tau_C = \infty$.

see Sec. VI B for further discussion.) Furthermore, the regime in which an oscillatory instability is predicted has been eliminated in Fig. 6 compared to its strong presence in Fig. 1. Linear stability results calculated in the limit $\tau_C = \infty$, again at various fixed $\Delta_Q = \Delta_C \equiv \Delta$, are shown in the plane of polymer modulus and activity (ζ , G_C) in Fig. 7.

With these linear stability results in mind, we now perform nonlinear runs in the elastomeric limit $\tau_C = \infty$ for the particular case of $G_C = 0.1$ and $\Delta = 10^{-4}$. For these parameters the critical activity threshold is $\zeta_c = 2.41$. Accordingly, we simulate a sequence of activity values $\zeta = 2-3.2$ straddling this threshold. The results are shown in Fig. 8. Figure 8(a) shows the time evolution of the (largest mode of the) perturbation to the base state, clearly showing exponential decay (linear stability) for activity values $\zeta < \zeta_c$ and early time exponential growth (linear instability) for $\zeta > \zeta_c$, giving way to saturation at long times due to nonlinear effects. We have checked that the slopes of the lines in Fig. 8(a) correspond to the eigenvalues predicted by the linear stability analysis.

For the particular run $\zeta = 3.2$ we now examine the system's behavior at long times, once it has settled to the ultimate nonlinear attractor of the dynamics. As before, we find states that are shear banded. Note, however, that conventional static shear bands are forbidden by the elastomeric nature of the polymer: A static band with a fixed nonzero shear rate would indefinitely load the polymer, leading to divergent polymer stresses. Indeed, inspecting the shear-rate profiles $\dot{\gamma}(y,t)$ in Fig. 8(b), we see instead traveling bands of local shear rate $\dot{\gamma} = \pm 0.15$ such that the fluid at any location in the flow cell is alternately sheared forward and then backward. Corresponding to this, the shear component C_{xy} of the polymer stress [Fig. 8(b)] also alternates in sign. These results are reminiscent of traveling density bands seen experimentally in cytoskeletal extracts [59]. Note that the initial linear instability is not oscillatory, consistent with the prediction in



FIG. 8. One-dimensional simulations with infinite polymer relaxation time. (a) Evolution of shear-rate perturbations for a range of activities ζ , showing growth for $\zeta > \zeta_c$ (solid lines) and decay for $\zeta < \zeta_c$ (dashed lines). (b) and (c) Space time plots for $\zeta = 3.2 > \zeta_c$. The color scale is (b) shear rate $\partial_y v_x = \dot{\gamma}$ and (c) C_{xy} . The parameters are $\tau_C \to \infty$, $G_C = 0.1$, and $\Delta = 10^{-4}$.

our linear stability analysis that eigenvalues are purely real for $\ell_Q^2/\tau_Q = \ell_C^2/\tau_C = \Delta$; recall Figs. 6 and 7.

Because the polymer relaxation time scale τ_C is infinite in these runs, the period of oscillation τ_{osc} must now be set by a separate time scale. To identify this time scale we performed simulations for a range of values of the activity ζ and solvent viscosity η . In each, we measured τ_{osc} via the largest peak in the Fourier series of the throughput as a function of time (see Fig. 9 inset). Collecting the data from all these runs in Fig. 9, we find reasonable evidence for the scaling

$$\tau_{\rm osc} \propto \bar{\tau}_a \equiv \eta / (\zeta - \zeta_c).$$
 (31)

Physically $\bar{\tau}_a$ is related to the time scale of active forcing identified in Ref. [55]. As discussed in that study, if the active forcing is faster than the relaxation time scales $\bar{\tau}_a < \tau_Q \ll \tau_C$, both the nematic and polymer lag behind, resulting in oscillatory behavior. Note, however, that for large values of the activity the flow becomes increasingly aperiodic and $\tau_{\rm osc}$ becomes less clearly defined, resulting in minor deviations from the suggested scaling law.

To summarize, we have shown both by linear stability analysis and also by full nonlinear simulations (so far in



FIG. 9. Dominant period of oscillation τ_{osc} versus the active time scale $\bar{\tau}_a$ [Eq. (31)]. The inset shows the throughput time series for $\zeta = 6.4$ and $\eta = 0.567$, with τ_{osc} marked. The parameters are $\eta = 0.284$ (red circles), 0.567 (green squares), 1.134 (blue diamonds), $\tau_C \rightarrow \infty$, $G_C = 0.1$, and $\Delta = 10^{-4}$.

one spatial dimension) that an elastomeric active nematic generically undergoes instability towards a spontaneously flowing state that must be oscillatory, otherwise the polymers would suffer infinite loading.

VI. NONLINEAR DYNAMICS (TWO DIMENSIONS)

In the previous section we saw a rich array of dynamical states with heterogeneity in one spatial dimension. These would clearly have been forbidden in any description that constrained the system to remain homogeneous in all directions (zero dimensions). We now increase the dimensionality further, to consider spatiotemporal dynamics in two dimensions. Two features are immediately anticipated that were forbidden in one dimension. One is the presence of $\pm \frac{1}{2}$ point defects in the director field [8,9,45]. The other is the possibility of extensional flow, which was forbidden by the constraint of fluid incompressibility in a 1D flow field of the form $\mathbf{v} = v(y)\hat{\mathbf{x}}$.

Consistent with these expectations, an earlier study of active nematics (without polymer) indeed found a much richer spectrum of phase behavior in two dimensions than in one dimension [5]. We now briefly remind the reader of those earlier findings, as a starting point from which to understand the effects of polymer below. Accordingly, the model's phase behavior without polymer is summarized in Fig. 10 (top) in the plane of (ζ, Δ) , where $\Delta \equiv \ell_Q^2 / \tau_Q$. The solid line shows the linear instability threshold (22) for the onset of the 1D viscous bending instability, given a homogeneous initial state with the director $\hat{\mathbf{n}}$ along y. The dotted line is the counterpart for the 1D viscous splay mode, given an initial director along x. As expected, the instability manifests itself at high ζ and low Δ , consistent with the fact that smaller system sizes (large Δ) tend to be stabilizing.

In the same diagram, the results of numerical runs performed without polymer in two dimensions symbols. are shown by Runs are initialized with ĥ along y, with a perturbation $\delta Q_{\alpha\beta} =$ $\sum_{n=1}^{32} \sum_{m=1}^{32} \delta A_{\alpha\beta}^{mn} \cos\left(\pi m y/L_{y}\right) \cos\left(2\pi n x/L_{x}+\theta_{\alpha\beta}^{n}\right), \text{ where }$



FIG. 10. Phase diagrams in the plane of diffusivity Δ and activity ζ , without polymer (top panel) and with polymer of relaxation time $\tau_C = 4$ and viscosity $\eta_C = 1$ (bottom panel). The solid line is the threshold for the onset of a 1D bending instability starting from a homogeneous initial state with the director $\hat{\mathbf{n}}$ along y. The dotted line is the counterpart for a 1D splay instability given an initial director $\hat{\mathbf{n}}$ of 2D numerical runs with the director $\hat{\mathbf{n}}$ initially aligned along y (subject to perturbations as described in the text). Triangles show 1D banded states, squares oscillatory states, and circles unsteady or chaotic states. The dashed line is the numerically observed crossover line ζ_c^{2D} beyond which 1D states always destabilize to the formation of 2D flows. Open and closed symbols denote states with zero and nonzero net throughput, respectively.

 $\delta = 10^{-10}$, $A_{\alpha\beta}^{mn}$ is a randomly selected magnitude drawn from a uniform distribution on [-1,1], and $\theta_{\alpha\beta}^n \in [0,2\pi)$ is a randomly selected phase (analogous perturbations are also added to $C_{\alpha\beta}$ when present). In the region where no linear instability is predicted the system remains homogeneous and quiescent (no spontaneous flow), even in these 2D runs. In contrast, beyond the threshold of the (bend) instability we find spontaneously flowing states, as expected. For activities only just above threshold, the ultimate solution corresponds to a steady 1D shear-banded state (triangular symbols), even though these runs in principle allow heterogeneity in two dimensions. In contrast, further into the unstable regime we find fully 2D spontaneous flow states. At intermediate activities we observe a range of oscillatory states, including states in which defect pairs undulate along the channel; these we denote by squares in Fig. 10. Spatially, these states exhibit repeating structures at a length scale roughly set by the channel width L_{v} . As the activity is increased (or Δ decreased), these oscillatory states gain additional frequency components when plotting a given scalar observable (e.g., throughput) against time. When there is no longer any discernible periodicity in this signal, we term the state aperiodic. In this limit the characteristic length scale of the resulting nematic texture decreases [45,60] and the order parameter fields lose any obvious spatial periodicity. The numerically observed crossover line ζ_c^{2D} separating 1D from 2D states is shown as a dashed line.

In each run, once the system has attained its ultimate attractor we measure the net throughput of fluid along the channel in the main flow direction x. The criterion that

we adopt to represent significant throughput is discussed in the Appendix. States meeting this throughput criterion are represented in Fig. 10 by closed symbols and those without significant throughput are shown by open symbols. As might be expected, the 1D laminar banded states show significant throughput. In contrast, most of the 2D states, with just a few exceptions, lack any coherent throughput: The more complicated velocity rolls associated with them have no overall flow direction.

With the above discussion in mind, we now consider the effects of adding polymer. We start in Sec. VI A by considering a viscoelastic polymer with a fixed viscosity $\eta_C = G_C \tau_C$. We then turn to the elastomeric limit $\tau_C \rightarrow \infty$ in Sec. VI B. In Secs. VI A and VI B we restrict ourselves for simplicity to the case of zero thermodynamic coupling between the nematic and polymer, $\chi = \kappa = 0$. (As discussed previously, a strong kinematic coupling is however still present.) Finally, in Sec. VI C we consider the effects of including explicit thermodynamic coupling.

A. Viscoelastic active matter

In this section we consider a polymer with a finite viscoelastic relaxation time scale τ_C . To avoid any complications associated with changing the overall viscosity, we consider fixed $\eta_C = G_C \tau_C$. (Accordingly, between runs in which τ_C progressively increases, G_C is progressively decreased in tandem.) We first explore the effects of polymer on the 2D nonlinear dynamics by revisiting the (ζ, Δ) phase diagram discussed above.

1. Long-time behavior: Active drag reduction

The bottom panel of Fig. 10 shows the phase diagram of the model, with polymer now included, for a polymer relaxation time $\tau_C = 4.0$. Compared to the polymer-free case (top panel), a dramatic difference is immediately apparent. Whereas without polymer the regime of significant throughput is confined predominantly to that of laminar 1D banded states, with added polymer even the majority of the more complicated 2D time-dependent flow states show a significant throughput. Three representative snapshot states with a significant throughput are shown in Fig. 11.

To explore this further, we pick a single set of parameters $\zeta = 3.2$ and $\Delta = 2 \times 10^{-5}$ representative of the regime of fully developed active turbulence in Fig. 10 (for both $\tau_C = 0.0$ and $\tau_C = 4.0$) and perform a separate series of runs across which we vary the value of the polymeric relaxation time τ_C , exploring the full range from the Newtonian limit (for the polymer) $\tau_C \to 0$ to the elastomeric limit $\tau_C \to \infty$. (Recall that the polymer modulus G_C must decrease in inverse proportion to τ_C , given our fixed value of $\eta_C = G_C \tau_C = 1$; likewise ℓ_C^2 must increase in proportion to τ_C to keep a fixed value of Δ .)

For this series of runs, we plot in Fig. 12 the root-meansquare fluid velocity $v_{\text{rms}} \equiv \sqrt{\langle |\mathbf{v}|^2 \rangle_{\mathbf{x}}}$, the mean throughput, and the mean defect density [61]. (Each such quantity is timeaveraged across a run, after discarding the initial transient.) As expected, in the regime of small τ_C where the polymer acts simply as an additional solvent, none of these quantities change with τ_C .



FIG. 11. Representative snapshot states with net throughput, from the $\tau_C = 4$ phase diagram (Fig. 10 bottom panel): banded ($\zeta = 0.025$ and $\Delta = 10^{-5}$), oscillatory ($\zeta = 0.6$ and $\Delta = 6.4 \times 10^{-4}$), and chaotic ($\zeta = 1.75$ and $\Delta = 8 \times 10^{-5}$); the color scale indicates $(n_x n_y)^2$. Defects of topological charge $\pm 1/2$ are identified by green dots (+) and red squares (-).

In contrast, once $\tau_C > \tau_Q$ the active stress has to work against an increasingly elastic fluid and $v_{\rm rms}$ decreases with increasing polymer relaxation time. Indeed, for $\tau_C \to \infty$ the polymer effectively arrests the flow altogether. (This holds only for the final behavior, after discarding the initial transient; we show below that this transient can in fact be long lived and have its own rich dynamics.)

Despite this monotonic decrease of the root-mean-square fluid velocity, the mean throughput has a nonmonotonic dependence on τ_C . For small τ_C , the throughput is essentially zero, consistent with the model's limiting behavior in the absence



FIG. 12. (a) Root-mean-square velocity $v_{\rm rms} = \sqrt{\langle |\mathbf{v}|^2 \rangle_{\mathbf{x}}}$ and mean throughput Ψ . (b) Mean defect density *n*. Each quantity is time averaged across a run, after discarding the run's initial transient, with the vertical bars denoting the standard deviation of the time series. The parameters are $\zeta = 3.2$, $\Delta = 2 \times 10^{-5}$, and $\eta_C = 1$.

of polymer discussed above. As τ_C increases to become comparable with τ_Q the throughput increases dramatically, suggesting a more coherent flow state capable of sustaining a net flow in one direction. (Indeed, not only does the mean throughput increase but the fluctuations of the throughput about this mean decrease.) Finally, for large τ_C the throughput again decreases as the overall flow arrests.

This gradual transition with increasing τ_C to a state of significant throughput, followed by arrest, suggests a progressive increase in the spatial coherence of the flow with increasing τ_C . Consistent with this, we see in Fig. 12(b) that the defect density *n* in the nematic's director field markedly falls with increasing τ_C , with the onset of this decline roughly coinciding with the peak in throughput. A strong correlation between $v_{\rm rms}$ and defect density has been noted previously in Refs. [45,60,62].

We have demonstrated, then, that adding polymer to a fluid showing fully developed active turbulence calms the short-scale structure of the spontaneous active flow, decreasing the nematic defect density and thereby increasing the flow correlation length to give a more organized flow state, often with a sustained net throughput [20,63].

For a passive Newtonian fluid displaying conventional inertial turbulence in a pressure-driven pipe flow at high Reynolds number [64], the addition of a small amount of long-chain polymer is known to calm the flow and reduce the ratio of imposed pressure gradient to throughput [65], in an effect known as polymer drag reduction. The phenomenon just described for our active fluid, whereby (zero-Reynolds-number) active turbulence is calmed by the addition of polymer, is strongly reminiscent of that effect and we term it active drag reduction accordingly. The persistent coherent motion to which it leads may have a biophysical analog in cytoplasmic streaming, where it is believed that active cytoskeletal materials such as actomyosin play a role in generating coherent flows, facilitating the transport of nutrients and organelles within the cell [26,27].

2. Transient dynamics: Extensional catastrophe

In the previous section we discussed the long-time behavior of viscoelastic active matter in two dimensions, ignoring the initial transient en route to the ultimate attractor of the dynamics. We now turn to consider that initial transient.

Again we pick a single set of parameters ($\zeta = 2.2$ and $\Delta = 4 \times 10^{-5}$), representative of the regime of fully developed active turbulence in the phase diagram of Fig. 10. We then perform a series of runs, varying the polymer relaxation time τ_C across the full range from $\tau_C \rightarrow 0$ to the elastomeric limit $\tau_C \rightarrow \infty$. We again fix both the overall polymer viscosity ($\eta_C = 1$) and the diffusivity Δ .

In each run we initialize the system as usual in a state that is homogeneous and quiescent, with the polymer undeformed, $\mathbf{C} = \mathbf{I}$. For the given parameters we expect perturbations to grow exponentially until nonlinear effects become important; this is confirmed in the early time growth dynamics in Fig. 13. (Note that the time signals in Fig. 13 do not appear chaotic until the turbulent state becomes fully developed, i.e., at later times in the run.)



FIG. 13. (a) Spatially averaged Deborah number. Marked as a dashed line is the threshold $De_c = \frac{1}{2}$ above which we might expect nonlinear exponential growth in the polymer strain. (b) Spatial average of Tr(**C** – **I**), quantifying the mean polymer strain. When $\langle De \rangle > De_c$ in (a), we indeed observe nonlinear exponential loading of the polymer in (b). Upward curves have τ_C from 10^{-2} (red line) to 10^4 (blue line) with logarithmic spacing. The parameters are $\zeta = 2.2$, $\Delta = 4 \times 10^{-5}$, $\eta_C = 1$.

The degree to which the polymer becomes deformed by this instability, as measured by the growth in Tr(C – I), increases with increasing τ_C . This is to be expected: Polymeric strains build up at a rate set by the velocity gradients in the fluid and relax at a rate set by $1/\tau_C$. Indeed, a polymer obeying the Oldroyd-B equation will, if subject to an indefinitely sustained simple shear flow of rate $\dot{\gamma}$, acquire a shear strain $\dot{\gamma}\tau_C$ and, if subject to an indefinitely sustained strain extensional strain $\dot{\epsilon}\tau_C/(1 - 2\dot{\epsilon}\tau_C)$ [66]. The latter is an infamous result, predicting the polymers to suffer unbounded deformation for any sustained extensional flow of rate $\dot{\epsilon} > 1/2\tau_C$. (In reality this growth would ultimately be cut off by finite chain extensibility or chain rupture, neither of which is captured by the Oldroyd-B model.)

Although the actively turbulent flow states that develop here do not subject any given fluid element to an indefinitely sustained extensional flow, they do nonetheless include regions with significant extension rates. To characterize these, we define a local extensional Deborah number $De = \tau_C \sqrt{\frac{1}{2}(\mathbf{D} : \mathbf{D})}$ and take its spatial average $\langle De \rangle$. In view of our above discussion of the Oldroyd-B model's extensional catastrophe, we might then reasonably expect a dramatic growth of polymer strain in any regime where $\langle De \rangle > De_c \approx \frac{1}{2}$.

In confirmation of this, the time evolution of $\langle De \rangle$ is plotted in Fig. 13(a) for each in our series of runs. For small τ_C (warm colors), $\langle De \rangle$ remains small and the spatially average polymer strain $\langle Tr(\mathbf{C} - \mathbf{I}) \rangle$ likewise remains modest. In contrast, for larger τ_C (cold colors) $\langle De \rangle$ exceeds $De_c \approx \frac{1}{2}$, leading to a dramatic effect on the polymer conformation: Following the initial linear instability, we observe a second period of exponential chain stretching that generates huge polymer strains [see Fig. 13(b)].

Any growth in the polymer strain gives a corresponding growth in the polymeric contribution $\Sigma_C = G_C(\mathbf{C} - \mathbf{I})$ to the stress. Recalling that $G_C = 1/\tau_C$ for this series of runs, the large values of τ_C for which dramatic chain stretching occurs have correspondingly small values of G_C . Accordingly, as long as the polymer strain remains O(1), the polymeric contribution to the stress will remain small. In contrast, in the regime of dramatic chain stretch the polymer strains become huge and the polymer stress then becomes comparable to the nematic and Newtonian ones, despite the small polymer modulus G_C . This in turn feeds back on the flow field, mitigating the extensional flows and halting the divergence of the polymer stress.

3. Limit cycle

In the previous two sections we considered state points (ζ, Δ) deep in the active-turbulent region of the phase diagram in Fig. 10 (which has $\tau_C = 4.0$). While the transient dynamics of corresponding runs at much larger τ_C revealed that polymer deformation proliferates due to the appreciable extensional component of the turbulent activity-driven flow field, it is unclear how this phenomenon manifests in the long-time dynamics. Therefore, we consider a point (ζ, Δ) in Fig. 10 with a more moderate activity (which is less deeply turbulent) and again perform a corresponding run at a much larger $\tau_C = 1000$ (once again at fixed $\eta_C = 1$), where we now run the simulation until long times $t_{\text{max}} \sim O(10\tau_C)$.

The results are shown in Fig. 14. In this case the system ultimately settles to an oscillatory state whose period is set by the polymer relaxation time τ_C . Over the course of one cycle, the system switches from a quasi-1D banded state dominated by bending in the director field (see the snapshot at time t_1) to a different quasi-1D banded state dominated by splay (at time t_2), via a fully 2D intermediate state with velocity rolls. A representative snapshot of such a 2D intermediate state is shown at time t_3 .

Closer study of the full run reveals that the transition from bend to splay initiates via the director, at a single value of x midway between the plates, rotating by $\pi/2$. This disturbance then propagates along the interface until the whole system is splayed. (Destabilization of the bend state to form the splay state has been seen previously in Ref. [67].) The splay-banded state then apparently remains stable for a time of order τ_C , before developing a roll-like instability around $t_3 = 4400$ characterized by alternative pairs of $\pm \frac{1}{2}$ topological defects [54] in the nematic director field.

The degree of inhomogeneity during these transitions can be monitored quantitatively through the power spectrum $P(k_x,t)$ of the spatial Fourier transform of the nonzero-order parameters $\phi = (Q_{xx}, Q_{xy}, Q_{yy}, C_{xx}, C_{xy}, C_{yy})$:

$$P(k_x) = \sum_{i=1}^{6} \int_0^{L_y} dy |\phi_i(k_x, y)|^2.$$
 (32)

Any purely 1D state would have $P(k_x,t) = 0$ for $k_x > 0$. As can be seen, the quasi-1D bend and splay states indeed have small *P*. The 2D roll-like states that arise during the transition

10

10

10

10

1500

1000

500

(b)

 $\mathbf{H}_{\mathbf{k}}$

(Tr(C - I))





FIG. 14. Representative state that continuously oscillates between bend and splay quasi-1D shear-banded states via an intermediate roll state. Plotted are (a) the power spectrum $P(k_x)$ for the first 20 Fourier modes $k_x = 1$ (red) $\rightarrow k_x = 20$ (blue) and (b) spatially averaged Tr(**C** – **I**) against time. The transitions between states are two dimensional in nature, as shown by the periodic spikes in $P(k_x)$. Shown on the bottom are snapshots for the times marked in (a) and (b) that show $(n_x n_y)^2$ (color map), director $\hat{\mathbf{n}}$ (red lines), and defects (symbols). The parameters are $\zeta = 1.24$, $\Delta = 1.6 \times 10^{-4}$, $\tau_C = 1000$, and $\eta_C = 1$.

between these quasi-1D states have, in contrast, a much larger P = O(1) [see Fig. 14(a)].

This oscillating effective dimensionality of the flow state directly influences the polymeric stretching, which we monitor as before with a scalar extension measure $\langle \text{Tr}(\mathbf{C} - \mathbf{I}) \rangle$ [see Fig. 14(b)]. The constraint of fluid incompressibility means that an extensional component to the flow field can only arise in the 2D states. Because of this, during times when the system occupies a quasi-1D bend or splay state the polymer extension relaxes on a time scale τ_C . In contrast, the intervening 2D roll state contains regions of extensional flow leading to significant polymer stretching.

It is worth noting that the nonlinear oscillatory state reported here would not have been predicted from our 1D linear stability analysis, for which the corresponding linear instability is not oscillatory. However, a 2D linear stability calculation that linearizes about one of the inhomogeneous banded states (shown in Fig. 14) may well predict this behavior. Note that oscillatory dynamics in active systems is widely seen in a biophysical context. For example, shape oscillations in developing cells are believed to be driven by actomyosin networks [68], which have been described theoretically using an elastic model [57].

B. Elastomeric active matter

In the previous section we saw that adding a polymer of even modest relaxation time has a dramatic effect on the phase behavior of confined active nematics. For example, it gives rise to an unusual drag-reduction effect with strong enhancements in the flow field's coherence and net throughput, for a fixed level of activity. We also explored the effect of changing the polymeric relaxation time τ_C , moving towards the elastomeric limit of large τ_C . We did so at a fixed overall polymer viscosity $\eta_C = 1$, with G_C decreasing in inverse proportion to τ_C .

In this section we consider the "true" elastomeric limit, in which $\tau_C \to \infty$ is taken at finite G_C so that the polymeric viscosity diverges. Physically, this could be realized by increasing the cross-link density and/or chain length in an entangled polymer of fixed concentration: Our model might then describe an actomyosin cell extract in a background of lightly cross-linked polymer gel. To implement this limit in our simulations we simply remove the local relaxation term prefactored by $1/\tau_C$ in the polymer equation of motion. As mentioned already for the 1D case in Sec. V B, we do however retain finite stress diffusivity $\Delta_C = \ell_C^2/\tau_C$, meaning that $\ell_C \to \infty$ as the elastomeric limit is taken. We discuss this choice further at the end of this section.

The predictions of our linear instability analysis in this limit were shown in Fig. 7. We now explore the instability more fully by performing nonlinear 2D simulations, at infinite τ_C , for a range of polymer moduli $G_C = 10^{-8} - 10^{-1}$. We choose values of the activity ($\zeta = 3.2$) and diffusivity ($\Delta = 8 \times 10^{-5}$) for which the initially homogeneous state is predicted to be unstable.

In each run, we find that the initial homogeneous state indeed destabilizes to form a heterogeneous complex liquidcrystalline texture with a high density *n* of defects in the director field [see Figs. 15(a) and 16 (top)]. Associated with this buildup of defects is a complex elastically turbulent deformation field with regions of (transient) extensional flow, which result in exponential stretching of the polymer [see Fig. 15(b)]. Because in these runs the polymer modulus obeys $G_C < G_Q$, appreciable polymer strain can develop [inset of Fig. 15(b)] during this early-time regime before the resulting polymeric contribution to the stress becomes comparable with solvent and nematic stresses, which are O(1) in our units. (A comparable phenomenon is seen in some models of polymeric glasses [69].)

Once the polymer stress does become comparable with the other stresses, the elastic turbulence arrests into a complex but almost frozen defect pattern. Thereafter, the defect density shown in Fig. 15(a) slowly coarsens over time, roughly as t^{-1} , which is the same as the classical result for coarsening in a passive nematic [70]. See Fig. 16 for snapshots of the



FIG. 15. Results from 2D runs with $\tau_C \to \infty$. We vary G_C from 10^{-8} (blue lines) to 10^{-1} (red lines) with logarithmic spacing. Snapshots for $G_C = 10^{-2}$ are given in Fig. 16. (a) Areal defect density n against time. Steps arise because n is discrete. (b) Scalar measure of polymer stress $G_C \langle \text{Tr}(\mathbf{C} - \mathbf{I}) \rangle$ [the inset shows the same data but not scaled by G_C , i.e., we plot $\langle \text{Tr}(\mathbf{C} - \mathbf{I}) \rangle$]. The parameters are $\zeta = 3.2$, $\Delta = 8 \times 10^{-5}$, and $\tau_C \to \infty$.

nematic texture during this process. Although a defect-free state $n \rightarrow 0$ might arise in the true steady-state limit $t \rightarrow \infty$, the coarsening process that leads to it is sufficiently slow that the strain pattern created by the arrested active turbulence might easily be mistaken for a final steady state. Note that the mean polymer extension continues to grow in this coarsening regime, albeit slowly (roughly logarithmically at intermediate times) with some suggestion of eventual saturation seen in the numerics. This elastomeric arrest of the active turbulent flow field, where strong polymer stretching in extensional flow regions creates strong stresses that oppose the flow,



FIG. 16. Snapshots for the points marked by triangles in Fig. 15(a) at times t = 20,100,700. Images show $(n_x n_y)^2$ (color map) and defects (symbols). The parameters are $\zeta = 3.2$, $\Delta = 8 \times 10^{-5}$, $\tau_C \to \infty$, and $G_C = 10^{-2}$.

is mechanistically reminiscent of the drag-reduction effects reported earlier. Note that the quasihomogeneous state that develops at long times is drastically different from the base state in our earlier linear stability calculation (which assumes that the polymer is undeformed) and is therefore not subject to the same linear instability as found there.

We now return to an issue raised previously in this section and earlier in Sec. V B, which is our choice to retain finite polymer stress diffusivity $\Delta_C = \Delta > 0$ even in the elastomeric limit of $\tau_C \rightarrow \infty$. Since $\Delta_C = l_C^2/\tau_C$, this requires matching divergence of the length l_C . (The physical meaning of this length is ambiguous, but it is often assumed to be of order the polymer coil size.)

Our reason for this choice is ultimately pragmatic: We have found that our 2D numerical simulations become highly unstable, at the discretization scale, if performed with $\Delta_C = 0$. A credible explanation of this behavior is as follows. We have seen that polymers encounter regions of strong elongation in 2D active flows. These convert globular initial stress fluctuations into threadlike ones, creating fine structure in the transverse direction as set by the local compression axis. Once elongation is strong, numerical instability is inevitable unless there is a restoring term to iron out these short-scale fluctuations. However, in the absence of thermodynamic couplings (see the next section), the only term in the polymer dynamics that can achieve this is Δ_C . Thus we retain finite Δ_C in order to capture the short-length-scale physics of local stress redistribution. This mechanism, whatever its details, should not disappear for lightly cross-linked elastomers, since their physics essentially merges with that of free chains at short length scales (below the cross-link separation).

Note that in an elastomer of finite Δ_C , although the mean stress caused by an applied deformation can never relax (which is the defining feature of a solid), any inhomogeneous polymer stress will relax to a uniform one eventually, via the stress diffusion term. This may seem a bit artificial, although a similar effect can arise in many elastomeric systems by other pathways. One of these, although absent from the present model, is solvent permeation flow [32]. Interestingly, one might expect that thermodynamic coupling to the **Q** field, whose stress diffusivity does not vanish as $\tau_C \rightarrow \infty$, could also provide enough stress redistribution to stabilize the numerics. Indeed, we find this to be the case, as we now discuss.

C. Explicit thermodynamic coupling

Throughout the preceding results sections we set $\kappa = \chi = 0$ in Eq. (3), disabling the direct thermodynamic coupling between **Q** and **C**. In that case, the only coupling between **Q** and **C** is purely kinematic, arising because they share a common fluid velocity **v** that is influenced by both sources of stress. In this section we study the effects of a direct coupling.

The κ term in Eq. (3) controls how the polymer pressure shifts the isotropic-nematic transition. Our interest in this paper lies far from the transition region, deep within the nematic phase, so for simplicity we set $\kappa = 0$. The remaining coupling parameter χ governs the energetics of relative orientations of **Q** and **C**. For $\chi < 0$, it is favorable for **Q** and **C** to align. Indeed, experiments (in the passive limit) suggest that single semiflexible polymers can couple to the nematic director field



FIG. 17. Plot of tracer trajectories for parameters characteristic of the turbulent phase for $t = 0 \rightarrow 200\tau_Q$. Polymer diffusion has been disabled by setting $\Delta_C = \ell_C^2/\tau_C = 0$. Trajectories are shown in (a) the no-polymer limit $\tau_C \rightarrow 0$ and in (b) the elastomeric limit $\tau_C \rightarrow \infty$. Colors are used to distinguish the separate tracer paths. The parameters are $\zeta = 3.2$, $\Delta_Q = 10^{-4}$, $\Delta_C = 0$, $G_C = 0.1$, and $\chi = +0.032$.

in this fashion [71]. However, such coalignment tends to arise kinematically even when $\chi = 0$, since **Q** and **C** both have similar alignment tendencies in relation to the spontaneous velocity field (at least for our choices of the flow-alignment and slip parameters ξ and *a*). We therefore do not expect substantially new physics to arise in this case. Therefore, we focus here on antagonistic coupling $\chi > 0$, where **Q** and **C** now prefer to orient with their major axes perpendicular. We do not perform an exhaustive search of parameter space but instead present a selection of the intriguing states observed.

1. Persistent elastomeric turbulence

We have shown previously (in one dimension) that persistent oscillatory states can develop in the elastomeric limit $\tau_C \rightarrow \infty$ provided we keep ℓ_C fixed. Our 2D simulations produce significantly more complex flow fields characterized by strong extensional components, which, as previously discussed, can be numerically problematic. Without any polymer diffusion we found that our numerics required impractically severe convergence criteria; with polymer diffusion retained (by imposing $\Delta_C = \ell_C^2 / \tau_C = \Delta_Q$) the simulations are stabilized but the initial turbulent solid state slowly coarsens away.

However, we might anticipate a similar stabilization to result also from a small thermodynamic coupling between C and Q, the latter of which retains spatial gradients even in the limit $\tau_C \rightarrow \infty$. Indeed, with this coupling we have successfully performed runs at zero polymer diffusivity, i.e., $\Delta_C = 0$, albeit using a significantly smaller space and time steps.² Intriguingly, we find that the transient elastomeric turbulence discussed previously now continues indefinitely, producing an unusual oscillatory state. This can be readily visualized by plotting the paths of massless tracer particles as they are advected by the flow, as shown in Fig. 17. Without polymer present, these tracers perform a random walk as they are advected by the turbulent flow. However, in the elastomeric limit, the tracer trajectories remain confined to periodic orbits for the duration of the simulation. In this limit, the activity-driven turbulent flow field stretches and distorts the elastic polymer background, which, at large enough strains, reacts by producing a restoring force that undoes any displacement, eventually returning displaced material points to their initial location (or nearby). We also plot the defect density n, throughput Ψ , and rms velocity in Fig. 18. These all apparently reach quasi-steady-state values after a short time and show no signs of the coarsening discussed previously for $\Delta_C \neq 0$ in Fig. 15. Also note that the throughput oscillates about zero, consistent with our expectation that an

²If $\Delta_C = 0$, the time step Δt needs to be an order of magnitude smaller than for $\Delta_C \neq 0$ and the spatial step size Δx must also be at least halved.



FIG. 18. Plot of (a) defect density *n* and (b) throughput Ψ (lower blue line) and $v_{\rm rms}$ (upper green line) in the elastomeric limit $\tau_C \to \infty$. Polymer diffusion has been disabled by setting $\Delta_C = \ell_C^2/\tau_C = 0$. The parameters are $\zeta = 3.2$, $\Delta_Q = 10^{-4}$, $\Delta_C = 0$, $G_C = 0.1$, $\chi = +0.032$, and $\tau_C \to \infty$.

elastic solid should undergo no net displacement. The severe numerical requirements when $\Delta_C = 0$ restrict our discussion to the qualitative aspects of the state; a more sophisticated numerical implementation would be required to fully quantify the elastomeric turbulence seen here.

2. Shear bands with interfacial defects

For the remaining examples we restore $\Delta_C = \Delta_Q = \Delta$ and fix a finite value of τ_C comparable to τ_Q . The first of these (which has $\chi = 0.002$) demonstrates an intriguing polymerdriven disorder-order transition. Choosing parameters characteristic of the chaotic state (when $\chi = 0$), at early stages of the run we find the defect-rich disordered state observed without polymer [Fig. 19(c)]. However, as the simulation progresses, ordered regions of nearly uniform director $\hat{\mathbf{n}}$ spread in from the walls towards the center of the channel, forming an increasingly shear-banded-like state [Fig. 19(d)]. Eventually,



FIG. 19. Example where an initially chaotic state organizes into a coherent shear-banded state with defects, which advect with the flow, embedded in the interface. Shown is the time evolution of (a) the power spectrum $P(k_x,t)$, averaged over y [for $k_x = 1$ (red lines) $\rightarrow k_x = 20$ (blue lines)] and (b) throughput Ψ . (c)–(e) Snapshots of $(n_x n_y)^2$ for the states marked left at times (c) t = 50, (d) t = 700, and (e) t = 3800. The parameters are $\zeta = 3.2$, $\Delta = 4 \times 10^{-5}$, $\tau_C = 10$, and $\chi = 0.002$.

the only remaining evidence of the earlier chaotic state are pairs of defects embedded in the interface [Fig. 19(e)].

This transition can be seen quantitatively by examining the power spectrum $P(k_x,t)$ in Fig. 19(a). During the initial chaotic phase, the first 20 wave vectors (plotted) contribute significantly to $P(k_x,t)$, indicating significant spatial structure in the *x* direction, with dynamics aperiodic in time. At long times, once the banded state forms, all amplitudes $P(k_x,t)$ are attenuated, particularly at large k_x . This (admittedly extreme) example is consistent with our drag-reduction argument, whereby polymer calms short-scale structure. Snapshots of the evolution of this state in Figs. 19(c)–19(e) demonstrate a clear transition from a disordered to an almost ordered state.

Correlated with this suppression of short-scale structure is a dramatic increase in the throughput [Fig. 19(b)]. While the chaotic state at early times has zero-mean throughput, the latter banded state develops a strong net flow in a spontaneously chosen direction, in this example towards the right.

Interestingly, during the intermediate phase between chaotic and banded states, we occasionally find transient rotating spiral structures, which when viewed macroscopically³ possess an integer topological charge +1 [see Fig. 19(d)]. Such structures are commonly found in polar active materials [59,72–74], but not in apolar nematics, which much more commonly display $\pm \frac{1}{2}$ defects [4], as were found above. In general, integer defects in passive nematics tend to dissociate [54], a tendency maintained in the active case without polymer. (One exception is in highly confined cylindrical geometries, as studied in Ref. [26] where the authors found a single +1 defect at low activities, which split into a pair of $\pm \frac{1}{2}$ defects only at higher activity values.) It appears that antagonistic coupling can help promote integer defects, but the detailed mechanism for this remains unclear.

3. Shuffling state

By increasing the activity with other parameters fixed, we can disrupt the above shear-banded state. The result [Fig. 20 (upper snapshot)] is a state that shuffles back and forth as a whole, with defects traveling along regions in which q, the largest eigenvalue of \mathbf{Q} , is small. This mechanism is similar to that reported in Ref. [67] (without polymer), where defect motion in walls (regions of high local distortion) was observed. The throughput time series in Fig. 20 reveals that the flow switches direction periodically on a time scale of the order of the viscoelastic relaxation time scale τ_C , once again confirming the direct influence of polymer on the dynamics.

4. Order-disorder coexistence

With a larger coupling constant ($\chi = 0.004$), even more exotic states can develop. In Fig. 20 (lower snapshot) we show a state exhibiting coexistence between chaotic and oscillatory regions (where the director lies in the *x*-*y* plane shown) and pseudoquiescent regions (where the director points along *z*). The active regions travel back and forth, trapped between

³Microscopically, at the heart of the structure, we find a pair of $+\frac{1}{2}$ defects, though these are close enough that the effective director field forms a spiral pattern of topological charge +1.



FIG. 20. Shown on top is an exotic oscillatory state that coherently shuffles alternatively left and right on a time scale $O(\tau_C)$. In the middle is the coexistence of bubbling active domains and regions where the director is out of plane (black). The bottom shows the throughput time series for both states (black line, $\chi = 0.002$; red line, $\chi = 0.004$). The parameters are $\zeta = 6$, $\Delta = 10^{-4}$, $\tau_C = 10$, $\chi = 0.002$ (upper snapshot), and 0.004 (lower snapshot).

the bounding walls. Repeating the simulations a number of times with different seeds for the 2D perturbation, we find that these pseudoquiescent regions sometimes grow to envelop the whole system, which then remains quiescent indefinitely. Such quiescent regions are likely made possible by our assumption of translational invariance along the vorticity axis z. (Recall that in our simulations we consider 3D order parameter tensors **Q** and **C** residing in a 2D space.) It may be that fully 3D simulations are required to fully resolve the dynamical behavior of this state, which could easily then show structure also in the z direction. We leave such 3D investigations to future work.

VII. CONCLUSION

We have studied in depth the linear stability and nonlinear dynamics of the minimally coupled model for viscoelastic active matter derived in Ref. [20]. While treating 3D structural tensors for both the active nematic and the polymer sectors, our studies assumed either 2D flow states (translationally invariant along the vorticity axis z) or 1D states (invariant also along the periodic axis x).

For appreciable activities, the phase diagram without polymer is dominated by chaotic states with no net throughput [5]. Adding polymer results in an active drag-reduction effect whereby the short-scale flow structure is calmed, resulting in a reduced density of nematic point defects. The increased flow correlation length is associated with net material transport around the periodic boundary conditions of our simulation. In fully confined geometries this would likely also help promote long-lived steady circulation as opposed to a mixing flow [27]. While adding polymer can delay the onset of spontaneous flow, our results predict a critical activity that does not diverge in the elastomeric limit $\tau_C \rightarrow \infty$ and indeed vanishes for large enough systems just as it does without polymer [10]. Thus activity-driven flows can occur within active materials that are ultimately solid: These might be called true active gels. Numerical simulation of such materials in one dimension indeed reveals oscillatory shear-banded states where the flow direction switches periodically as the polymer stress is loaded. A range of complex states involving nontrivial interplay of viscoelasticity and activity were seen in systems of large but finite polymer relaxation time τ_C .

As the relaxation time of the polymer was increased, the effect of activity-driven extensional flows was shown to be important. When the spatially averaged Deborah number (which describes the ratio of polymer and extensional time scales) exceeds a critical value, we found that the initial linear instability is followed by a period of rapid exponential deformation of the polymer. For sufficiently large τ_C , this resulted in oscillatory states that cycled between rapid extensional deformation of the polymer and slow stress relaxation on a time scale τ_C .

In the elastomeric limit $\tau_C \rightarrow \infty$ the ultimate fate of the oscillatory and/or turbulent states that we discovered remains sensitive to numerical details of the model. Retention of finite stress diffusivity (which greatly improve numerical stability) in this limit causes ultimate relaxation to a state of uniform stress; this precludes any permanent state of oscillation or elastic turbulence. However, numerically we were able to switch off the stress diffusivity once thermodynamic coupling between nematic and polymeric degrees of freedom was included. In this case we found that the system could be truly a turbulent solid even in steady state. That is, the coupled system shows chaotic activity-induced velocity fields that persist indefinitely but that reverse sign often enough locally that the elastic strain remains bounded, as a solid requires.

More generally, we found a wide range of complex and interesting spontaneous flow states when **Q** and **C** are antagonistically coupled at the free-energy level. Depending on the strength of the coupling, our simulations show that polymer can drive a transition from active turbulence to near-laminar banded flow, result in states with oscillatory dynamics on a time scale τ_C , and create regions where director is oriented out of the plane. Fully 3D simulations should provide useful insight, particularly for the last of these.

Our prediction of active turbulence in soft solid materials $(\tau_Q \rightarrow \infty)$ arises when $G_C/G_Q \lesssim 0.1$. This looks experimentally feasible for subcellular active matter (though probably not swarms of bacteria) within a lightly cross-linked polymer gel. The symmetry breaking characterized by net throughputs, which we observe at moderate polymer relaxation times $(\tau_C/\tau_Q \approx 10)$, could have particular relevance for studies of cytoplasmic streaming, where coherent flow is crucial for material transport within the cell. The same physics may also apply, albeit at larger scales, to cell migration in confined geometries [29]. Our results with explicit coupling that show oscillatory shuffling states may also be relevant in describing the cell shape oscillations of Ref. [56].

We hope our work will promote experiments on these and other forms of active viscoelastic matter; while the effects of viscoelasticity on individual swimmers have been studied previously [75,76], we are not aware of an equivalent study for bulk orientationally ordered phases. One related study did however consider the linear stability of the bulk *isotropic* phase in an Oldroyd-B fluid [18]. While the specifics of that model differ from the one presented here,⁴ both studies find (i) that increasing η_C at fixed τ_C will always eventually suppress the spontaneous flow instability, (ii) that at small τ_C the polymer simply renormalizes the solvent viscosity (this is our viscous limit), and (iii) a region at large τ_C where unstable oscillatory behavior is predicted (this is our elastomeric limit).

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APPENDIX: THROUGHPUT CRITERION

At time *t*, the throughput of the system is defined as

$$\Psi(t) = \frac{1}{L_y} \int_0^{L_y} v_x(t) dy = \langle v_x(t) \rangle_y, \qquad (A1)$$

which is independent of x because of fluid incompressibility. Because this quantity generally exhibits significant fluctuations in time, particularly in the chaotic regime, we define our criterion for significant or net throughput as being when the mean of the throughput time series μ_{Ψ} exceeds the standard deviation σ_{Ψ} :

$$\mu_{\Psi} > \sigma_{\Psi}, \tag{A2}$$



FIG. 21. Method for determining throughput Ψ . Shown on the left is a state with net throughput, in which the throughput direction intermittently switches between right and left. The red bins show the normalized histogram of $\Psi(t)$ and the solid black line is a fit using two Gaussian functions centered at $\pm \mu_{\Psi}$. In this example, the positive throughput state lasted for a shorter time, hence the difference in heights (means and standard deviations are the same). Both peaks will tend to the same height in the limit $t \to \infty$. Here $\zeta = 5$, $\Delta = 3.2 \times 10^{-4}$, and $\tau_C = 1$. On the right is a state with no net throughput for comparison, with $\zeta = 5$, $\Delta = 10^{-5}$, and $\tau_C = 1$. The insets show examples of throughput time series for each run.

where

$$\mu_{\Psi} = \frac{1}{t} \int_0^t \Psi(t') dt' \tag{A3}$$

and

$$\sigma_{\Psi} = \sqrt{\frac{1}{t} \int_0^t [\Psi(t') - \mu_{\Psi}]^2 dt'}, \qquad (A4)$$

which converge to constant values as $t \to \infty$. Note that under this definition, states that have nonzero mean throughput will fail the criterion if this mean is less than the standard deviation of the time series.

In practice, we find that the flow direction can intermittently switch (see, for example, Fig. 21, left). If naively averaged as above, this would clearly produce zero-mean throughput, at least in the limit $t \rightarrow \infty$. Therefore, we instead perform a least-squares fit, fitting the throughput histogram with two Gaussian functions of width σ_{Ψ} , centered at $\pm \mu_{\Psi}$. We have explicitly checked that our results are robust to the number of histogram bins used. Examples of both throughput and nonthroughput states are shown in Fig. 21. We mark states satisfying criterion (A2) in the phase diagrams with closed symbols and those failing it with open symbols.

- M. C. Marchetti, J.-F. Joanny, S. Ramaswamy, T. B. Liverpool, J. Prost, M. Rao, and R. A. Simha, Rev. Mod. Phys. 85, 1143 (2013).
- [2] C. Dombrowski, L. Cisneros, S. Chatkaew, R. E. Goldstein, and J. O. Kessler, Phys. Rev. Lett. 93, 098103 (2004).
- [3] F. Nédélec, T. Surrey, A. C. Maggs, and S. Leibler, Nature (London) 389, 305 (1997).
- [4] T. Sanchez, D. T. N. Chen, S. J. DeCamp, M. Heymann, and Z. Dogic, Nature (London) 491, 431 (2012).
- [5] S. M. Fielding, D. Marenduzzo, and M. E. Cates, Phys. Rev. E 83, 041910 (2011).
- [6] H. H. Wensink, J. Dunkel, S. Heidenreich, K. Drescher, R. E. Goldstein, H. Löwen, and J. M. Yeomans, Proc. Natl. Acad. Sci. USA 109, 14308 (2012).
- [7] L. Giomi, M. J. Bowick, X. Ma, and M. C. Marchetti, Phys. Rev. Lett. 110, 228101 (2013).
- [8] L. Giomi, M. J. Bowick, P. Mishra, R. Sknepnek, and M. C. Marchetti, Philos. Trans. R. Soc. London Ser. A 372, 16 (2014).

⁴There are several differences between our study and Ref. [18]. In that work (i) the stability analysis is about an isotropic base state, (ii) liquid-crystalline stresses are not included, (iii) the active particles are self-propelled, and (iv) the concentration field can vary in space (we assume a homogeneous concentration).

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- [9] S. P. Thampi, R. Golestanian, and J. M. Yeomans, Phys. Rev. Lett. 111, 118101 (2013).
- [10] R. Voituriez, J.-F. Joanny, and J. Prost, Europhys. Lett. 70, 404 (2005).
- [11] F. Jülicher, K. Kruse, J. Prost, and J.-F. Joanny, Phys. Rep. 449, 3 (2007).
- [12] R. A. Simha and S. Ramaswamy, Phys. Rev. Lett. 89, 058101 (2002).
- [13] S. Ramaswamy, Annu. Rev. Condens. Matter Phys. 1, 323 (2010).
- [14] A. N. Beris and B. J. Edwards, *Thermodynamics of Flowing Systems* (Oxford University Press, Oxford, 1994).
- [15] P. G. de Gennes and J. Prost, *The Physics of Liquid Crystals* (Clarendon, Oxford, 1995).
- [16] A. C. Callan-Jones and F. Jülicher, New J. Phys. 13, 093027 (2011).
- [17] J.-F. Joanny and J. Prost, Sém. Poincaré XII, 1 (2009).
- [18] Y. Bozorgi and P. T. Underhill, J. Rheol. **57**, 511 (2013).
- [19] A. W. Decho, Ocean. Mar. Biol. Annu. Rev. 28, 73 (1990).
- [20] E. J. Hemingway, A. Maitra, S. Banerjee, M. C. Marchetti, S. Ramaswamy, S. M. Fielding, and M. E. Cates, Phys. Rev. Lett. 114, 098302 (2015).
- [21] E. Lauga, Phys. Fluids 19, 083104 (2007).
- [22] J. M. Teran, L. Fauci, and M. J. Shelley, Phys. Rev. Lett. 104, 038101 (2010).
- [23] L. Zhu, E. Lauga, and L. Brandt, Phys. Fluids 24, 051902 (2012).
- [24] S. E. Spagnolie, B. Liu, and T. R. Powers, Phys. Rev. Lett. 111, 068101 (2013).
- [25] E. E. Riley and E. Lauga, Europhys. Lett. 108, 34003 (2014).
- [26] F. G. Woodhouse and R. E. Goldstein, Phys. Rev. Lett. 109, 168105 (2012).
- [27] R. E. Goldstein and J.-W. van de Meent, Interface Focus 5, 20150030 (2015).
- [28] L. R. Serbus, B.-J. Cha, W. E. Theurkauf, and W. M. Saxton, Development 132, 3743 (2005).
- [29] S. R. K. Vedula, M. C. Leong, T. L. Lai, P. Hersen, A. J. Kabla, C. T. Lim, and B. Ladoux, Proc. Natl. Acad. Sci. USA 109, 12974 (2012).
- [30] H. Wioland, F. G. Woodhouse, J. Dunkel, J. O. Kessler, and R. E. Goldstein, Phys. Rev. Lett. **110**, 268102 (2013).
- [31] S. Ramaswamy, R. A. Simha, and J. Toner, Europhys. Lett. 62, 196 (2003).
- [32] S. T. Milner, Phys. Rev. E 48, 3674 (1993).
- [33] P. D. Olmsted, O. Radulescu, and C.-Y. D. Lu, J. Rheol. 44, 257 (2000).
- [34] P. Olmsted (private communication).
- [35] G. Jayaraman, S. Ramachandran, S. Ghose, A. Laskar, M. S. Bhamla, P. B. Sunil Kumar, and R. Adhikari, Phys. Rev. Lett. 109, 158302 (2012).
- [36] H. Stark and T. C. Lubensky, Phys. Rev. E 67, 061709 (2003).
- [37] R. G. Larson, Constitutive Equations for Polymer Melts and Solutions (Butterworths, Boston, 1988).
- [38] M. Warner and E. M. Terentjev, *Liquid Crystal Elastomers* (Oxford University Press, Oxford, 2003).
- [39] M. E. Cates, S. M. Fielding, D. Marenduzzo, E. Orlandini, and J. M. Yeomans, Phys. Rev. Lett. **101**, 068102 (2008).
- [40] S. A. Edwards and J. M. Yeomans, Europhys. Lett. 85, 18008 (2009).

- [41] D. Marenduzzo, E. Orlandini, M. E. Cates, and J. M. Yeomans, J. Non-Newtonian Fluid Mech. 149, 56 (2008).
- [42] S. Zhou, A. Sokolov, O. D. Lavrentovich, and I. S. Aranson, Proc. Natl. Acad. Sci. USA 111, 1265 (2014).
- [43] H. See, M. Doi, and R. G. Larson, J. Chem. Phys. 92, 792 (1990).
- [44] P. D. Olmsted and S. T. Milner, Macromolecules 27, 6648 (1994).
- [45] E. J. Hemingway, P. M. Mishra, S. M. Fielding, and M. C. Marchetti (unpublished).
- [46] D. A. Head, A. J. Levine, and F. C. MacKintosh, Phys. Rev. Lett. 91, 108102 (2003).
- [47] C. Storm, J. J. Pastore, F. C. MacKintosh, T. C. Lubensky, and P. A. Janmey, Nature (London) 435, 191 (2005).
- [48] G. H. Koenderink, Z. Dogic, F. Nakamura, P. M. Bendix, F. C. MacKintosh, J. H. Hartwig, T. P. Stossel, and D. A. Weitz, Proc. Natl. Acad. Sci. USA 106, 15192 (2009).
- [49] M. R. Mofrad, Annu. Rev. Fluid Mech. 41, 433 (2009).
- [50] W. H. Press, S. A. Teukolsky, W. T. Vetterling, and B. P. Flannery, *Numerical Recipes in C: The Art of Scientific Computing*, 2nd ed. (Cambridge University Press, New York, 1992).
- [51] C. Pozrikidis, Fluid Dynamics: Theory, Computation, and Numerical Simulation, 2nd ed. (Springer, New York, 2009).
- [52] D. Marenduzzo, E. Orlandini, M. E. Cates, and J. M. Yeomans, Phys. Rev. E 76, 031921 (2007).
- [53] S. H. Strogatz, Nonlinear Dynamics and Chaos: With Applications to Physics, Biology, Chemistry, and Engineering (Westview, Boulder, 2008).
- [54] P. M. Chaikin and T. C. Lubensky, *Principles of Condensed Matter Physics* (Cambridge University Press, Cambridge, 2000).
- [55] L. Giomi, L. Mahadevan, B. Chakraborty, and M. F. Hagan, Nonlinearity 25, 2245 (2012).
- [56] G. Salbreux, J.-F. Joanny, J. Prost, and P. Pullarkat, Phys. Biol. 4, 268 (2007).
- [57] K. Dierkes, A. Sumi, J. Solon, and G. Salbreux, Phys. Rev. Lett. 113, 148102 (2014).
- [58] A. Aradian and M. E. Cates, Europhys. Lett. 70, 397 (2005).
- [59] V. Schaller, C. Weber, C. Semmrich, E. Frey, and A. R. Bausch, Nature (London) 467, 73 (2010).
- [60] S. P. Thampi, R. Golestanian, and J. M. Yeomans, Philos. Trans. R. Soc. London Ser. A 372, 14 (2014).
- [61] D. Huterer and T. Vachaspati, Phys. Rev. D 72, 043004 (2005).
- [62] L. Giomi, Phys. Rev. X 5, 031003 (2015).
- [63] Y. Bozorgi and P. T. Underhill, J. Non-Newtonian Fluid Mech. 214, 69 (2014).
- [64] O. Reynolds, Philos. Trans. R. Soc. London 174, 935 (1883).
- [65] C. M. White and M. G. Mungal, Annu. Rev. Fluid Mech. 40, 235 (2008).
- [66] J. Rallison and E. Hinch, J. Non-Newtonian Fluid Mech. 29, 37 (1988).
- [67] S. P. Thampi, R. Golestanian, and J. M. Yeomans, Europhys. Lett. 105, 18001 (2014).
- [68] L. He, X. Wang, H. L. Tang, and D. J. Montell, Nat. Cell Biol. 12, 1133 (2010).
- [69] S. M. Fielding, R. G. Larson, and M. E. Cates, Phys. Rev. Lett. 108, 048301 (2012).
- [70] A. Bray, Adv. Phys. 43, 357 (1994).
- [71] Z. Dogic, J. Zhang, A. W. C. Lau, H. Aranda-Espinoza, P. Dalhaimer, D. E. Discher, P. A. Janmey, R. D. Kamien,

PHYSICAL REVIEW E 93, 032702 (2016)

T. C. Lubensky, and A. G. Yodh, Phys. Rev. Lett. **92**, 125503 (2004).

- [72] K. Kruse, J.-F. Joanny, F. Jülicher, J. Prost, and K. Sekimoto, Phys. Rev. Lett. 92, 078101 (2004).
- [73] J. Elgeti, M. E. Cates, and D. Marenduzzo, Soft Matter 7, 3177 (2011).
- [74] X. Yang, D. Marenduzzo, and M. C. Marchetti, Phys. Rev. E 89, 012711 (2014).
- [75] X. N. Shen and P. E. Arratia, Phys. Rev. Lett. 106, 208101 (2011).
- [76] D. A. Gagnon, N. C. Keim, X. Shen, and P. E. Arratia, Phys. Fluids 26, 103101 (2014).