



Extensile to contractile transition in active microtubule–actin composites generates layered asters with programmable lifetimes

John Berezney^a, Bruce L. Goode^b , Seth Fraden^a , and Zvonimir Dogic^{a,c,d,1}

^aDepartment of Physics, Brandeis University, Waltham, MA 02454; ^bDepartment of Biology, Brandeis University, Waltham, MA 02454; ^cDepartment of Physics, University of California, Santa Barbara, CA 93106; and ^dBiomolecular Science and Engineering Program, University of California, Santa Barbara, CA 93106

Edited by Michael Cates, Centre for Mathematical Sciences, University of Cambridge, Cambridge, United Kingdom; received September 9, 2021; accepted December 21, 2021

We study a reconstituted composite system consisting of an active microtubule network interdigitated with a passive network of entangled F-actin filaments. Increasing the concentration of filamentous actin controls the emergent dynamics, inducing a transition from turbulent-like flows to bulk contractions. At intermediate concentrations, where the active stresses change their symmetry from anisotropic extensile to isotropic contracting, the composite separates into layered asters that coexist with the background turbulent fluid. Contracted onion-like asters have a radially extending microtubule-rich cortex that envelops alternating layers of microtubules and F-actin. These self-regulating structures undergo internal reorganization, which appears to minimize the surface area and maintain the ordered layering, even when undergoing aster merging events. Finally, the layered asters are metastable structures. Their lifetime, which ranges from minutes to hours, is encoded in the material properties of the composite. These results challenge the current models of active matter. They demonstrate self-organized dynamical states and patterns evocative of those observed in the cytoskeleton do not require precise biochemical regulation, but can arise from purely mechanical interactions of actively driven filamentous materials.

self-organization | active matter | pattern formation | nonequilibrium driving

Cytoskeletal active stresses, which are collectively generated by thousands of microscopic molecular motors, empower living organisms with the capacity to reshape themselves and their environment. Precise control of these self-organized stresses in both space and time enables diverse biological processes, including embryogenesis, intracellular transport, cell motility, and cytokinesis (1–3). Reconstituting such spatiotemporal dynamics in synthetic active materials is worth pursuing for several reasons. From a fundamental perspective, the development of experimental active matter systems provides an opportunity to test the emerging theories of internally driven nonequilibrium systems (4). In turn, such theoretical frameworks could enable a rigorous description of various dynamical architectures observed in biology (5, 6). From an applications perspective, active matter systems could provide an experimental platform for creating a new class of autonomous shape-changing materials that mimic sought-after properties of living organisms.

The above-described considerations inspired studies of structures and dynamics that emerge in collections of interacting motile agents, which are either chemically synthesized or biologically isolated. Several behaviors have been observed. Self-propelled Brownian particles, which lack structural anisotropy and attractive interactions, form dense condensates that coexist with a dilute gas and are reminiscent of equilibrium gas–liquid phase separation (7, 8). Motile objects with polar symmetry form dense flocks that exhibit directed sinuous motion (9). Anisotropic active agents, such as bacterial swimmers or

molecular motor-powered extensile bundles, remain spatially homogeneous but generate turbulent-like dynamics and autonomous flows (10–12). Finally, cytoskeletal filaments and molecular motors also exhibit bulk contractions into dense solid-like materials (13–17). These advances provide a foundation for introducing biological functionality into materials science. However, several barriers remain. First, predicting the emergent self-organization and the bulk material properties from known microscopic dynamics remains a challenge (18, 19). For example, it is difficult to predict whether a particular combination of motor proteins and cytoskeletal filaments will exhibit extensile turbulent-like dynamics or contractile network collapse (20). Second, living organisms can rapidly assemble large-scale dynamical architectures, disassemble these same structures at a later predetermined time, and then, reassemble the same constituents into other distinct structures. In contrast, engineering synthetic active materials with multiple dynamical states and controlling the switching between these states remain a challenge.

Motivated by these twin challenges, we developed a composite system of cytoskeletal active matter that merges an active microtubule (MT)-based extensile network powered by kinesin-1 motors and a passive network of entangled F-actin filaments.

Significance

Active forces sculpt the forms of living things, generating adaptable and reconfigurable dynamical materials. Creating synthetic materials that exhibit comparable control over internally generated active forces remains a challenge. We demonstrate that active composite networks, collectively driven by the force-generating molecular motors, exhibit complex spatiotemporal patterns similar to those observed in cell biology. Amongst others, we describe robust self-assembly of onion-like layered asters. A self-regulating mechanism ensures the asters' layered structure survives coalescence-like events, while their temporal stability is encoded in the mechanical properties of the network. Our model system elucidates the essential role of passive elasticity in controlling the emergent nonequilibrium dynamics while also establishing a robust experimental platform for engineering lifelike materials.

Author contributions: J.B., B.L.G., S.F., and Z.D. designed research; J.B. performed research; B.L.G. contributed new reagents/analytic tools; J.B. analyzed data; and J.B., S.F., and Z.D. wrote the paper.

The authors declare no competing interest.

This article is a PNAS Direct Submission.

This open access article is distributed under [Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 \(CC BY-NC-ND\)](https://creativecommons.org/licenses/by-nd/4.0/).

¹To whom correspondence may be addressed. Email: zdogic@ucsb.edu.

This article contains supporting information online at <http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.2115895119/-DCSupplemental>.

Published January 27, 2022.

Such composites of active and passive filaments enable independent control of the magnitude of the active stresses, exclusively generated by the active MT component, and the passive viscoelasticity, dominated by the entangled network of F-actin filaments. This additional degree of freedom widens the accessible phase space, whose full exploration reveals diverse dynamical states and complex kinetic pathways. With increasing F-actin concentration, we observe a transition from extensile turbulent-like dynamics to bulk contraction. The transition regime is characterized by the coexistence of an extensile fluid and locally contracted onion-like asters. The finite-lifetime structured asters exhibit a layered spatial structure that survives coalescence. Taking these behaviors together, the active MT and passive actin composites share certain features of seemingly distinct categories of active matter, including extensile and contractile networks, as well as active condensation.

Composite MT–Actin Networks

To prepare a composite network with tunable mechanical properties, we combined active MT bundles with a passive actin network whose viscoelasticity is well controlled. G-actin polymerizes into micrometer-long semiflexible filaments that, at high concentrations, form a viscoelastic entangled network with an elastic plateau at intermediate frequencies (Fig. 1*A*) (21, 22). Over the range of F-actin concentrations studied here, the elastic plateau increases from 0.01 to 2 Pa, which is higher than the elasticity of the MT networks alone (23). Consequently, the composite's viscoelasticity is dominated by the F-actin component, effectively decoupling the network's passive mechanics from the active stresses that are only generated by the MT component. The previously used nonspecific bundling agent, poly(ethylene glycol), induced coassembly of mixed MTs and F-actin bundles that lacked activity (24, 25). To produce a force-generating MT network, we used protein regulator of cytokinesis 1 (PRC1), a cross-linker that specifically bundles antiparallel MTs while still allowing for the kinesin-driven interfilament sliding (26, 27).

Controlling Initial Conditions

The preparation of the composite played a critical role in establishing reproducible dynamics. Both actin polymerization and the reconfiguration of extensile MT networks are adenosine triphosphate (ATP)-dependent processes, but they take place on different timescales. G-actin polymerizes over hours, while the active MT network rearranges on the timescale of seconds (23, 28). Upon simply mixing all the components, the MTs formed an extensile network within minutes, which thereafter, generated substantial flows before G-actin could polymerize into F-actin. The ATP-powered MT-driven flow churned the polymerizing actin into various flow-dependent structures, resulting in ATP-dependent initial conditions that violated our design constraints.

To overcome this challenge, we developed a two-step protocol. First, we prepared samples with the same initial configuration consisting of a fully polymerized F-actin network interdigitated with a passive MT network (Fig. 1*A* and *B*). Subsequently, we activated the MT network to initiate the nonequilibrium dynamics. To accomplish this, we mixed monomeric G-actin with a kinesin–MT–PRC1 bundled network that was frozen in a particular configuration due to the lack of ATP. G-actin was polymerized over hours using a nonhydrolyzable ATP analog, adenylyl-imidodiphosphate (AMP-PNP) (29). Replacing all traces of ATP with AMP-PNP ensured the kinesin–MT network remained quiescent throughout the F-actin polymerization. Subsequently, we initiated nonequilibrium dynamics with a caged ATP precursor (30). Upon illumination with ultraviolet (UV) light, the active composite almost instantaneously transitioned from quiescence to

turbulent-like flows (Movie S1). Intriguingly, the speed of the flows exhibited an initial peak, thereafter settling into a steady state that was sustained by the ATP regeneration systems (Fig. 1*C*). The concentration of (adenosine 5'-triphosphate, P3-(1-(4,5-dimethoxy-2-nitrophenyl)ethyl) ester (DMNPE-caged ATP) determined the stepping speed of the kinesin motors and the overall system dynamics (11, 31). This two-step protocol was essential for generating reproducible dynamics. It provided ATP-independent, uniformly interdigitated F-actin–MT networks from which the nonequilibrium dynamics was initiated.

Extensile Fluid at Low F-Actin Concentrations

In the absence of F-actin, MTs formed a network of bundled filaments to which kinesin motor clusters bound passively. Uncaging the ATP induced motor stepping and MT sliding. At a steady state, the network rapidly reconfigured through repeating cycles of motor-driven bundle extension, buckling, fraying, and reannealing, as previously observed (11). Such network dynamics powered the large-scale flows of the aqueous solution. Previously, motor-driven MT networks were studied either within viscous Newtonian fluids or passive liquid crystals (11, 23, 32). Here, we examined how a passive viscoelastic component influences the MT network dynamics.

At low F-actin concentrations, the kinesin-driven MT bundles generated an extensile fluid whose properties were tuned by the F-actin concentration. Upon initiation of activity, both the MT and F-actin networks yielded, exhibiting turbulent-like flows (Fig. 2*A* and Movie S2). With increasing F-actin concentrations, the flow speed decreased (Fig. 2*B*). The spatial structure of the autonomous flow fields was characterized by an exponentially decaying spatial velocity–velocity correlation. Increasing the F-actin concentration had little effect on the spatial flow structure (Fig. 2*C*). This suggests that the active flow fluidized the F-actin network such that it contributed little to the material dynamical length scale.

The two interpenetrating networks can couple directly through steric interactions and indirectly through the flows, which are generated by the active MT network but influence both components. To investigate such coupling, we labeled both filaments. In the presence of activity, regions with high MT concentration also had higher F-actin concentrations (Movie S3). Correlating the two networks showed activity-dependent spatial colocalization (Fig. 2*D*). When active dynamics ceased due to ATP depletion, the colocalization of the two networks declined. Eventually, they became anticorrelated as the F-actin relaxed away from the static MT bundles, filling the previously empty voids (Fig. 2*D* and *E*).

Aster Formation and Dynamical Coexistence at Intermediate F-Actin Concentrations

With increasing network viscoelasticity, the flow speed decreased and approached zero around \sim 6 μ M F-actin. Beyond this point, different dynamics emerged. In particular, the composite network separated into a high-density contracting phase, which coexisted with a low-density extensile fluid (Fig. 3*A* and Movie S4). Initially, we observed a dense system-spanning network of nodes connected by thin threads (Movie S5). Over time, the threads thinned and ruptured. Subsequently, the ruptured threads contracted toward the nodes and rounded, assuming aster-like shapes that reduced their surface area. Because the motor-condensed filament-rich structures had a layered interior with an outer cortex enriched in radially aligned MTs, we refer to them as “layered asters.” Similar to single-filament asters, layered asters continued to increase in size through coalescence events (15, 33–35).

To quantify the emergence of structure within the active composite, we Fourier transformed the images to calculate the power spectral density (PSD) of the MT fluorescence

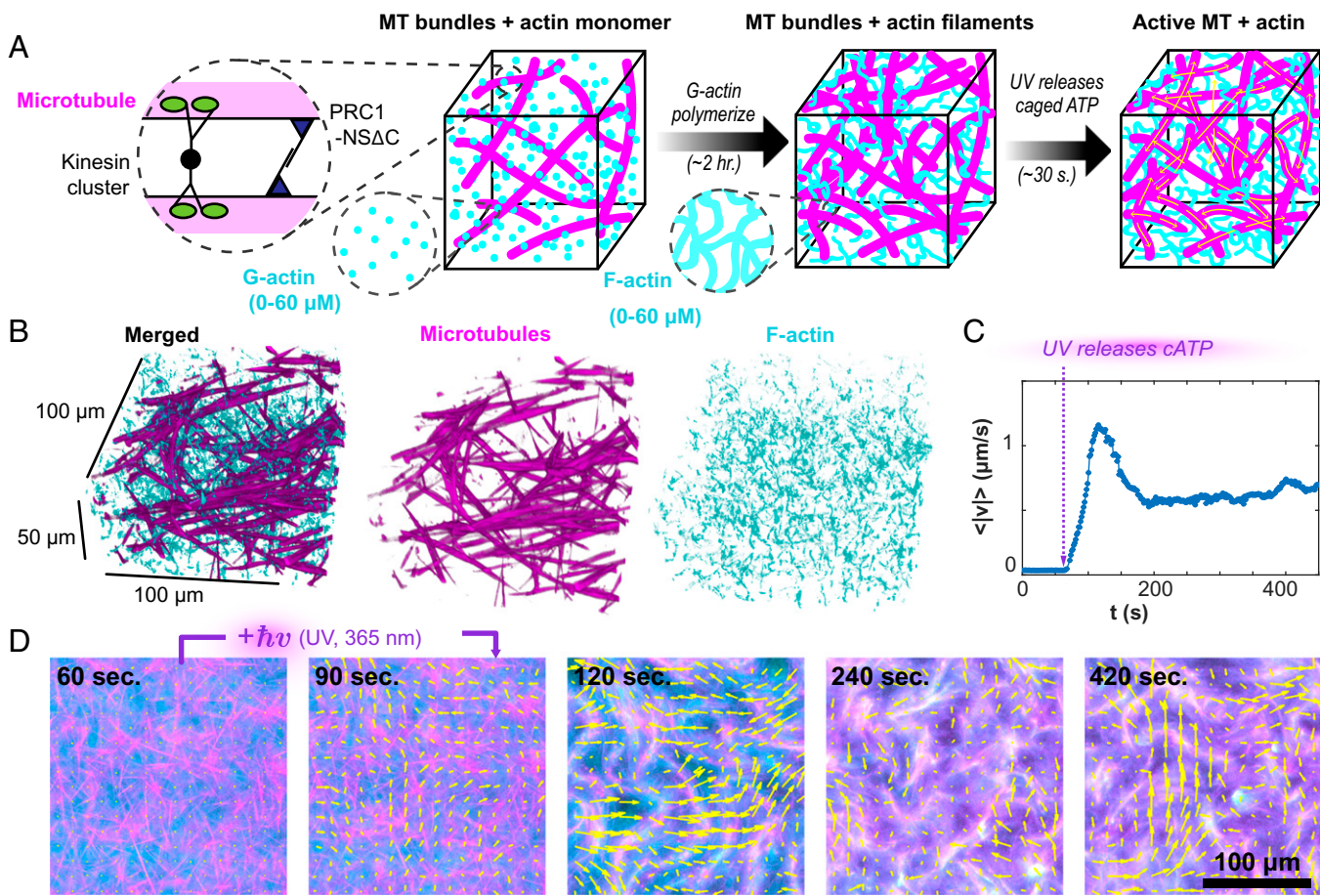


Fig. 1. Light-activated F-actin–MTs composites. (A) GMPCPP-stabilized MTs are bundled by a specific MT cross-linker, PRC1-NS Δ C, and driven by streptavidin-based kinesin motor clusters. AMP-PNP-bound G-actin is added to the MT bundle network and polymerizes in the high-salt buffer. After 2 h, filamentous actin formed a uniform interpenetrating actin network. At that point, DMNP-caged ATP (cATP) was cleaved with UV light, and the kinesin motors generated large-scale reorganization. (B) Confocal microscopy shows the isotropic uniform interpenetrating network of actin and MTs. (C) UV illumination releases caged ATP, generating flows measured by particle image velocimetry. (D) Kinesin-driven flow drives the reorganization of the actin and MT networks (Movie S1). Yellow arrows indicate the velocity field.

micrographs over time. The presence of a distinct peak in the $PSD(q,t)$ is a result of a strong spatial correlation in the MT concentration, characteristic of structures with length scale q_m^{-1} (Fig. 3B). The position of this peak, $q_m(t)$, and its height, $PSD(q_m,t)$, changed with time. One can identify three stages of separation, delineated by the rate of change in the structural length scale. In the first stage (I), the density contrast developed between the filament-rich and filament-poor regions, leading to an increase in the magnitude of $PSD(q_m)$ but little change in the length scale, q_m^{-1} (Fig. 3C and D). In the second stage (II), $q_m^{-1}(t)$ increased concurrently with the peak's magnitude, $PSD(q_m)$, indicating coarsening of the separated domains (Fig. 3C and D). The development of a peak in the PSD and the subsequent decrease in q_m are markers of classical spinodal decomposition (36). However, in active composites, q_m scaled as $\sim t^{-1}$, much faster than equilibrium phase separation of a minority phase (37–39). At the third stage (III) of separation, the rapid increase in both $PSD(q_m)$ and q_m^{-1} halted, and nonmonotonic changes ensued, in contrast to the phase separation of passive components.

We used image segmentation to quantify the formation of filament-rich and filament-poor domains (Fig. 3E and F). At the shortest times, the initially uniform composite structure became mottled with small filament-poor voids, which grew in size and number. N_V is the number of voids per image frame (Fig. 3A, 22 min and E, right axis). Growth in N_V continued in regime II,

where it reached a maximum, thereafter dropping rapidly toward unity. During this transition, the many separate voids joined as the filament-rich threads ruptured. The collapse in N_V captured the structural transition of the filament-rich domains from a continuous phase to a minority phase. There was a concomitant decrease in the area fraction of the contracted phase, φ_C , in regimes I and II (Fig. 3E, left axis). In stage III, φ_C reached a minimum and began to increase, a nonmonotonic behavior suggesting the emergence of new dynamics.

We examined how the material partitioned between the domain types by estimating the total F-actin and tubulin concentration from the fluorescence intensities (Fig. 3F). During regimes I and II, the concentration of the condensed phase, C_C , increased, while the concentration of the background extensile fluid, C_F phase decreased. Subsequently, during stage III, C_C and C_F reached a peak and nadir, respectively, before slowly returning toward their initial values.

Layered Aster Have a Programmable Lifetime. We observed well-defined asters by 50 min (Fig. 4A–C). Beyond this time point, the asters began disassembling from their maximally compacted state. At the lowest F-actin concentration (8 μ M), asters completely disassembled within 100 min (Fig. 4A and Movie S6). Above 10 μ M, aster dissolution occurred over in several stages (Fig. 4B). First, the aster diameter slowly increased through expansion, while its filament density decreased. As

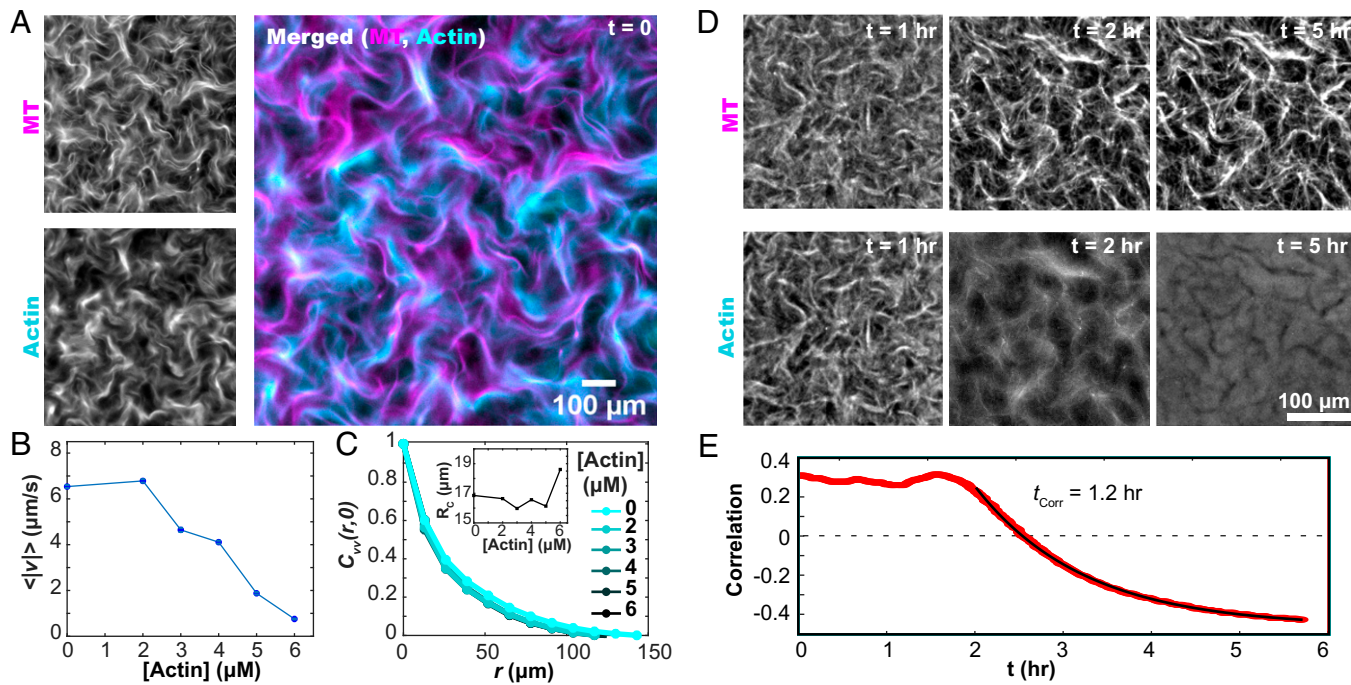


Fig. 2. F-actin controls the dynamics of the extensible MT network. (A) Kinesin motors drive the reorganization of both F-actin and MT networks. (B) The mean speed of the autonomous flows decreases with increasing F-actin concentration. (C) The spatial velocity-velocity correlation function of the autonomous flows is independent of the F-actin concentration. (C, Inset) The correlation lengthscales, extracted from fits to exponential decay, are independent of F-actin concentration. (D) Upon cessation of motor-driven flow, F-actin relaxes away from its out-of-equilibrium configuration while the MT bundles remain fixed (Movie S3). (E) After motor-driven activity halts at 1.8 hours, the spatial correlation between the MT and F-actin concentration fields decreases as a function of time.

asters swelled, the captured material dissociated from the asters, joining the background fluid, which demonstrated that the asters' constituents were biochemically intact, but the asters were no longer stable (Movie S6). F-actin concentration above 30 μM stabilized the contracted asters, such that even after 5 h, little structural change occurred (Fig. 4C and Movie S6).

We quantified the aster disassembly dynamics by plotting the time evolution of the average MT concentrations within the layered asters, $I_{\text{MT}}(t)$. The MT concentration initially increased, consistent with local contractions (left axes in Fig. 4 D–F). The concentration reached a peak and then began to decay as the asters disassembled. The decay timescale from the peak intensity varied with the F-actin concentration. For samples containing 8 μM F-actin, at 180 min, $I_{\text{MT}}(180)/I_{\text{MT}}(0) = 1$, indicating complete disassembly. For 10 μM F-actin samples, this measure of disassembly was reached at 300 min. At higher 30 μM F-actin concentrations, asters never fully disassembled: $I_{\text{MT}}(300)/I_{\text{MT}}(0) = 1.9$. Thus, F-actin concentration governs the disassembly timescale.

The aster's effective size provided additional insight into the disassembly dynamics (right axes in Fig. 4 D–F). At early times, the aster's average effective radius, r_{eff} , grew along with the intensity. In this regime, coalescence primarily drove the increase in size, while contraction drove the increase in concentration. By about 10 to 20 min, r_{eff} reached a maximum and then began to decrease. I_{MT} continued to increase, suggesting the first peak is driven by contraction-dominated dynamics. Subsequently, the radius began to increase with time. Thereafter, expansion occurred as the aster intensity started dropping, a sign of the incipient disassembly. The final decay in r_{eff} was not observed at 30 μM F-actin. Interestingly, the separation of these timescales was small for 8 μM F-actin, and the two peaks in r_{eff} were poorly defined.

Self-Regulating Structure of the Layered Aster. Confocal microscopy revealed that the aster's structure consists of three

concentric layers, which varied both in the composition and the local symmetry (Fig. 5A and Movie S7). The outer cortex, enriched in MTs, consisted of a monolayer of radially aligned bundles, largely excluding F-actin (Fig. 5B). The cortex enclosed an isotropic network of MTs and F-actin. The aster core was enriched in MTs. With increasing radius, the concentration of F-actin increased, while that of MTs decreased, generating an intermediate layer enriched in F-actin.

To understand the development of the self-organized asters, we quantified the spatiotemporal organization in terms of the F-actin, MTs, and kinesin clusters (Fig. 6A). We measured the average intensity, $\langle J_C(t) \rangle$, for each component over time, within 10 μm of the center of each aster. The normalized F-actin concentration remained nearly constant, whereas the concentrations of the kinesin motor clusters and MTs increased (Fig. 6B). Notably, the increase in the motor concentration was greater than the increase in the MT concentration. The accumulation of motors over the entire aster structure was less striking (Fig. 6C). This highlights a preference for the motor clusters to move inward, accumulating at the core. Such inward motion is common in aster-forming systems (35, 40). It suggests that the cortex is polarity sorted, with MT plus ends pointing inward.

Large ($\sim 100\text{-}\mu\text{m}$) aster assemblages were propelled by the surrounding extensible fluid flow (Movie S4). When two asters encountered each other, they coalesced while robustly preserving the aster's intricate layered structure (Fig. 7 and Movie S8). The coalescence initiated as the two cortex layers overlapped. At this point, the portions of the outer cortices proximal to each other disappeared. Presumably, upon the overlap, the inactive polarity-sorted MT cortex monolayers annihilated as their opposite polarity enabled efficient interfilament sliding. Subsequently, the actin-rich cores moved toward each other and joined, forming a spherical droplet-like aster. Finally, the partially disrupted outer cortex reassembled and rounded itself in a process that suggests the presence of effective surface tension.

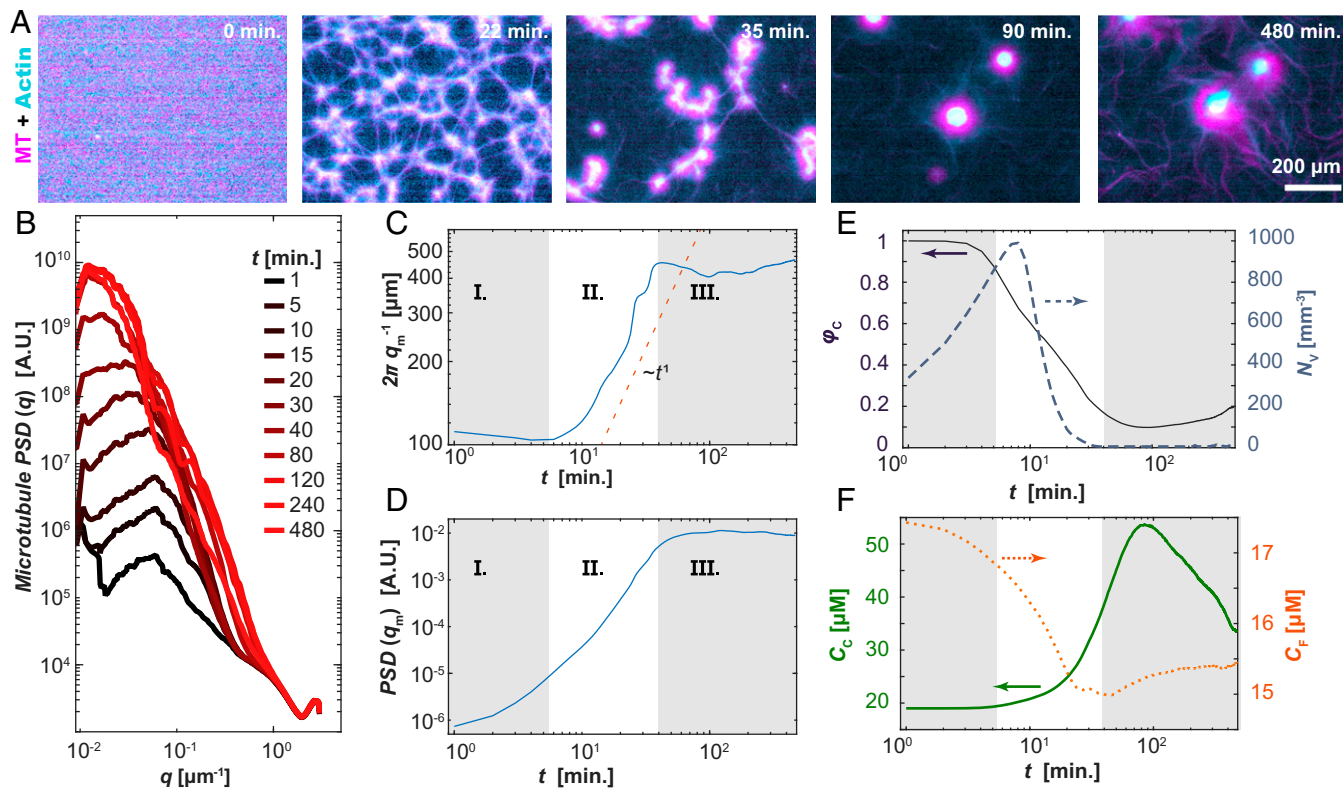


Fig. 3. Structure of self-organized layered asters. (A) Time series showing a uniform composite phase separating into filament-rich aster and filament-poor extensile fluids (Movie S4). (B) The PSD, $PSD(q)$, of the MT fluorescence micrographs shows the emergence of a peak with increasing time. (C) The position of the inverse of the PSD peak density, q_m^{-1} , plotted against time exhibits three stages of phase separation. (D) The magnitude of the PSD peak is plotted as a function of time. (E) Time-dependent fractional area of the contractile domain (solid line; left axis) and the number density of the interstitial extensile fluid domains (dashed line; right axis) extracted from real-space images. (F) The combined F-actin and MT concentration within the contractile aster phase (solid line; left axis) and active fluid phase (dotted line; right axis) as a function of time. Samples contained 100 μ M caged ATP, 1.3 mg/mL MTs, and 3 μ M actin.

Bulk Contraction at High Concentrations of F-Actin

At intermediate concentrations of actin, the viscoelastic network quickly yielded, limiting the large-scale propagation of contractile active stresses and generating the local contraction of structured asters. At still higher F-actin concentrations, the increased network integrity permitted active stresses to percolate across the entire sample, causing the composite to collapse (Fig. 8 and Movie S9). After uncaging the ATP, the bulk contracting MT–actin network pulled inward, separating from the chamber margins while maintaining the internal connectivity (Fig. 8A). The boundaries of the contracting structure remained uniform, while its geometry was defined by the chamber shape.

We measured the fractional width of the contracting composite over time (Fig. 8D). Immediately after ATP uncaging, the network exhibited little change. After a lag time of several minutes, rapid contraction ensued. The width of the contracting network was described by two exponential decays and associated timescales:

$$W(t) = a_1 e^{\frac{(t-b)}{c_1}} + a_2 e^{\frac{(t-b)}{c_2}} + d. \quad [1]$$

The initial lag time and exponential-type decay are reminiscent of previous observations from single-component filament–motor systems (15, 41). However, the double exponential form suggests a more complicated response.

We quantified the changes in the fluorescence profiles of F-actin and MTs across the sample chamber (Fig. 8B). Like the coexistence of structured asters and an extensile fluid, a filament-poor extensile phase coexisted with the contracting network. The MTs strongly partitioned into the contracting phase, while F-actin

separated more evenly. The contracting structure became heterogeneous over time. MT-poor voids developed, which yielded late-stage large variations in the MT density (Fig. 8B). Averaging over the local fluctuations, the density of MTs was higher at the edge of a contracting network, as seen previously (15). Interestingly, F-actin is more evenly distributed throughout the sample. At the edges of the bulk-contracted network, MT bundles extended perpendicular to the contraction direction, locally excluding F-actin (Fig. 8C). This outer layer is reminiscent of the MT-rich cortex of layered asters (Fig. 5A).

We studied how F-actin concentration affects bulk contractions. The MTs partitioned almost entirely into the contracting network, while the F-actin partitioned less strongly (Fig. 9A). Internal fracturing played a more important role at both the lower and higher F-actin concentrations, while intermediate concentrations yielded more uniform contraction (Fig. 9B). MT concentration decreased with increasing distance from the sample edge. The degree to which the F-actin followed this behavior varied, suggesting that the mechanical coupling between the two networks depends on the composite’s composition. For 15 μ M F-actin, the density profiles of both filaments were comparable (Fig. 9B). At 30 μ M F-actin, the profiles between F-actin and MTs were uncorrelated, and the F-actin profile did not increase at the edges. At 50 μ M F-actin, there was an anticorrelation in the F-actin and MT profiles.

The final width of the contracted network exhibited a minimum for 30 μ M F-actin (Fig. 9C). At both the low and high ends of the F-actin, the contracted network started to lose its connectivity. To quantify this internal self-tearing, we used image segmentation to estimate the area fraction of MT-poor

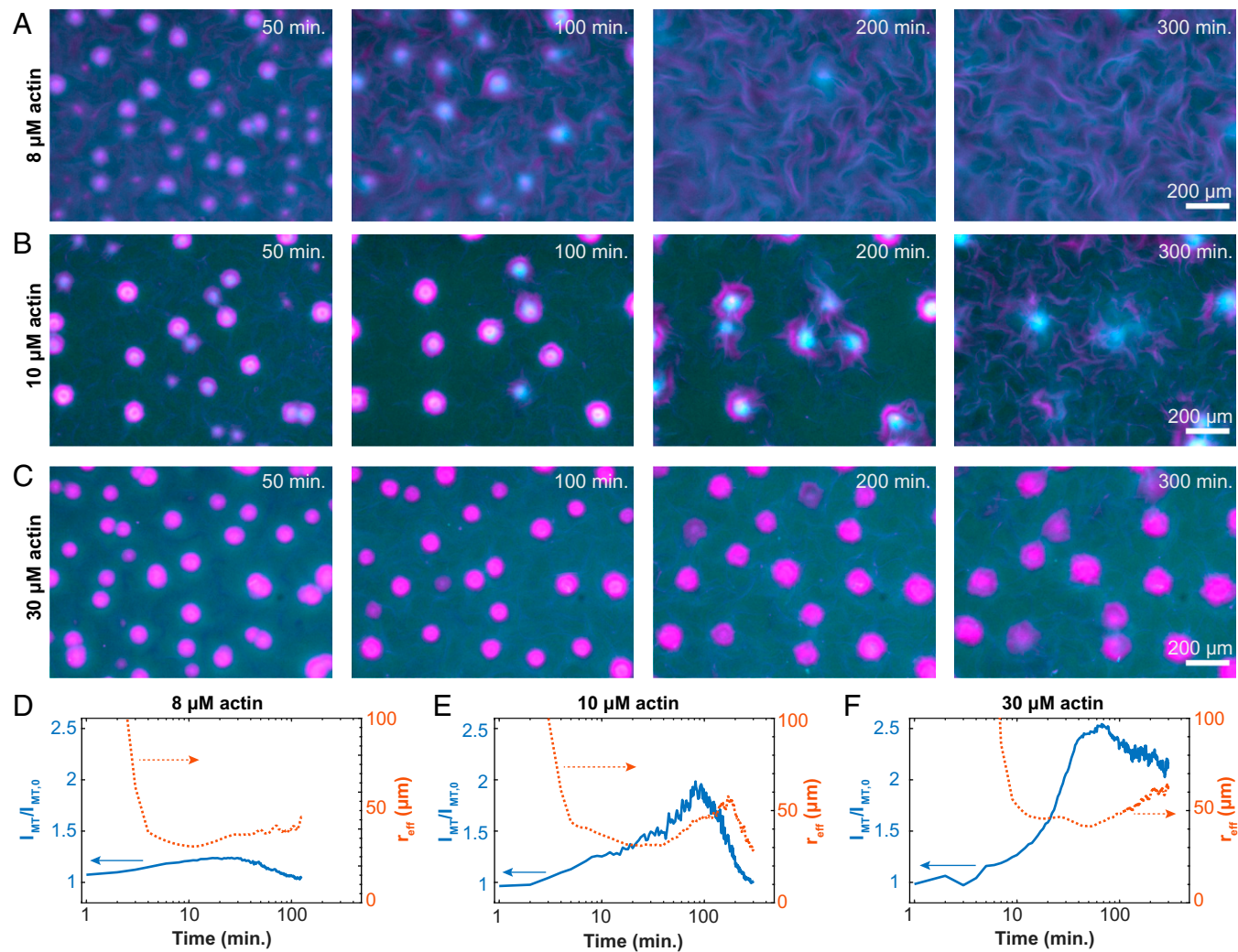


Fig. 4. Programmable finite lifetime of layers asters. (A–C) Time-lapse images show the structural changes of contracted asters embedded in an active fluid at 8, 10, and 30 μM F-actin (Movie S6). (D–F) Normalized MT fluorescence intensity within the contracted aster phase (solid line; left axes) and equivalent spherical radius of aster domains (dashed line; right axes) show how temporal evolution depends on actin concentration (8, 10, and 30 μM actin).

voids within the contracted domain (Fig. 9D). Maximal contractility yielded the smallest void fraction, while F-actin concentrations above and below that value resulted in a larger separation of F-actin and MTs. Maintaining network connectivity is important for generating maximal contractility. Another notable feature was the partitioning of the filaments between the contracting composite network and the enveloping fluid phase. Increasing F-actin concentration monotonically increased the concentration of actin filaments within the contracted network (Fig. 9E). However, the ratio of F-actin inside and outside the contracted network decreased as a function of F-actin concentration and approached unity (Fig. 9E, Inset). These observations suggest that the composite network changes from a low-density limit, where both filaments cocontract, to a high-density limit, where the MT network contracts within a viscoelastic suspension of entangled F-actin.

State Diagram of the Actin–MT Composites

Our work builds on the previous studies of active composites (42, 43). So far, we have described how increasing F-actin concentrations induce transitions from the extensile fluid to layered asters and then, to global contractions. To map out the entire phase diagram, we varied both the F-actin and the ATP concentration. Because the dynamics was time dependent, the diagram

summarizes structures observed after 1 h (Fig. 10). Changing ATP concentrations had a similar effect as F-actin. Low ATP concentrations drove MT network contractions, even in the absence of F-actin. Increasing ATP concentration induced a transition to extensile fluids. This echoes previous results showing how motor activity can fluidize cytoskeletal networks (23, 44–46).

While F-actin concentration determines the composite's viscoelasticity, the ATP concentration plays a dual role; it controls both the active stresses and the sample viscoelasticity. Kinesin-generated active stress is determined by its ATP-dependent stepping rate (47). Between the sequential force-generating steps that kinesin motors take, they remain attached to the MT, acting as passive cross-linkers. Lower ATP concentrations increase the fraction of kinesin clusters that passively cross-link MTs. Thus, decreasing ATP effectively increases the viscoelasticity of the MT network alone (23). Consequently, it is not surprising that ATP concentration, like F-actin, also plays a role in determining the composite's behavior at nonzero F-actin concentrations. Similar to F-actin, increasing ATP concentration induced transitions from the bulk contraction to structured asters and extensile fluids.

Extensile Bundles Drive Bulk Contractions

The microscopic details that underlie the emergence of the above-described dynamics remain unclear. To gain insight, we

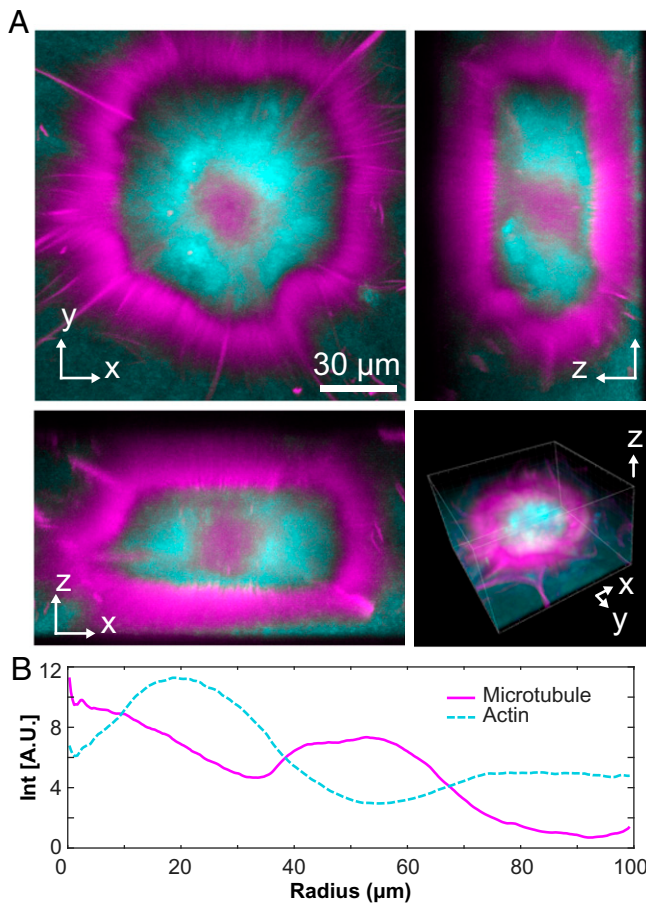


Fig. 5. Structure of layered asters. (A) Two-dimensional projections of an aster imaged with confocal microscopy (Movie S7). (B) The radial profile of MTs and actin reveals a layered organization consisting of an MT-rich core, an actin-rich intermediate layer, and the outer cortex composed of radially aligned MTs.

quantified the behavior of individual MT bundles within the contracting network. Isolated fluorescent MT bundles were doped into a network of unlabeled filaments. Upon initiation of activity, these bundles extended to several times their initial contour length before buckling (Fig. 11). Therefore, these results demonstrate that, despite the long-term contraction, the initial kinesin-driven activity generates MT bundle extension. Such observations provide a glimpse into the microscopic mechanisms by which active networks produce two seemingly separate processes of extensile fluid flows and solid-like contractions.

Discussion

The dynamics of active matter is determined by the balance between active stresses, generated by the motile constituents, and the viscous or elastic reaction stresses that arise due to the deformation of the material. In single-component formulations of cytoskeletal-based active matter, independent control of active and passive stresses can be challenging. For example, in conventional MT-based active fluids, kinesin motors act both as passive cross-linkers and generators of active forces (23, 48). Consequently, decreasing the ATP concentration decreases the magnitude of the active stresses while increasing the elastic modulus of the material. This dual role limits the full exploration of the nonequilibrium phase space while also impeding comparison with theory. The MT–actin active composites overcome these obstacles, allowing one to independently tune both the active stresses that are exclusively generated by the MT component and

the passive stresses that are largely determined by the viscoelastic F-actin network. Separating these two control parameters yielded several unexpected findings. First, increasing the passive viscoelasticity changed the symmetry of the self-organized active stresses, inducing a transition from extensile to contractile dynamics. Second, the transition from extensile to contractile stresses was marked by the coexistence of two dynamically organized structures: filament-rich layered asters and filament-poor extensile fluids. Third, the layered asters exhibited self-regulating spatial organization. Finally, the layered asters were also transients whose lifetimes were encoded in the F-actin concentration. We discuss each of these observations.

Passive Viscoelasticity Controls the Symmetry of Active Stresses.

With notable exceptions from two-dimensional motility assay geometries (49, 50), cytoskeletal active matter typically exhibits either uniform contractile or extensile dynamics, which implies the existence of a single type of self-organized active stress (13, 16, 20, 43, 51–53). It remains a challenge to predict the nature of the mesoscopic active stresses given known microscopic dynamics. Several mechanisms have been proposed, including 1) the spatial arrangement of motors along polar filaments and their enhanced propensity for tip binding, 2) the spatially dependent dynamics of motor stepping along the filaments, and 3) the asymmetry introduced by mechanical rigidity of the constituent filaments (17, 52, 54, 55). Disentangling these mechanisms in a single-component network is challenging as filament spatial organization and mechanics and the dynamics of motors on those filaments are interrelated (56).

We found that a viscoelastic F-actin network can induce a transition from extensile flows to bulk contractions. We also observed the coexistence of an extensile and a contractile state. These dual observations challenge existing theoretical models.

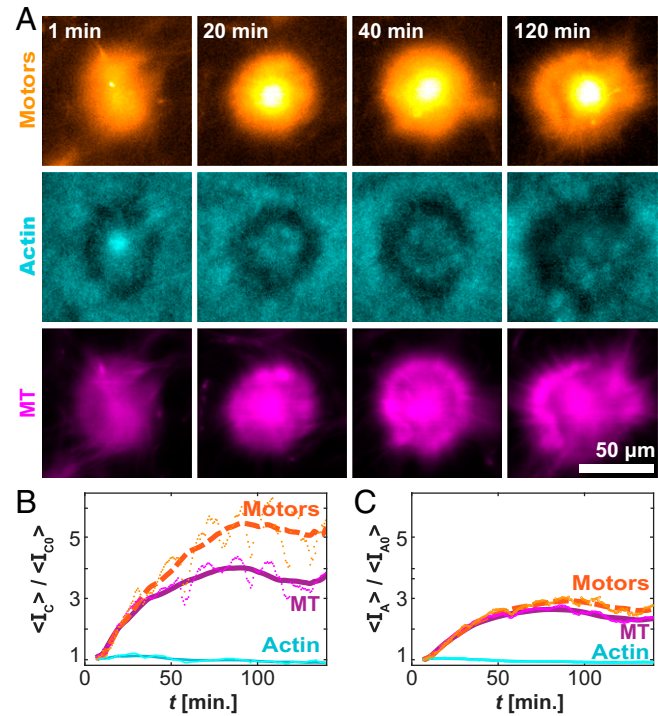


Fig. 6. Temporal evolution of the aster structures. (A) Time-lapse images of a single aster for the motor (Top), actin (Middle), and MT (Bottom) channels. (B) Time evolution of the normalized mean intensity within 6.5 μm of the centroid. (C) Time evolution of the normalized intensity over the entire contracted domain. Lines represent a moving average over 20 min. Data are from 300 μM caged ATP, 6 μM actin, and 100 μM caged ATP.

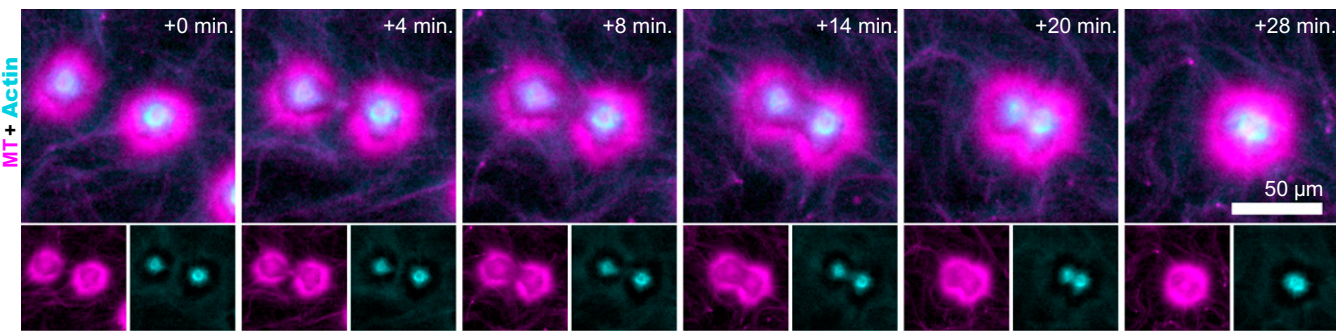


Fig. 7. Coalescence of layered asters. Time-lapse images show the coalescence of two nearby asters (Movie S8). Coalescence events preserve the intricate structure of the layered asters.

They are also different from previous work with myosin-driven MT–actin composites, which showed that the viscoelastic response could be tuned by the addition of a second material component, but the symmetry of the active stress did not change (43). However, they are broadly consistent with previous experimental work in which the deformations of rigid filaments changed from extensile to contractile through the addition of passive cross-linkers (52). Even in contractile conditions, the constituent MT bundles initially exhibited extensile dynamics (Fig. 11). Notably, the rectification of extensile into contractile forces was demonstrated for nonlinear elastic networks (57). The applicability of these descriptions to the viscoelastic active composites studied here requires further investigation.

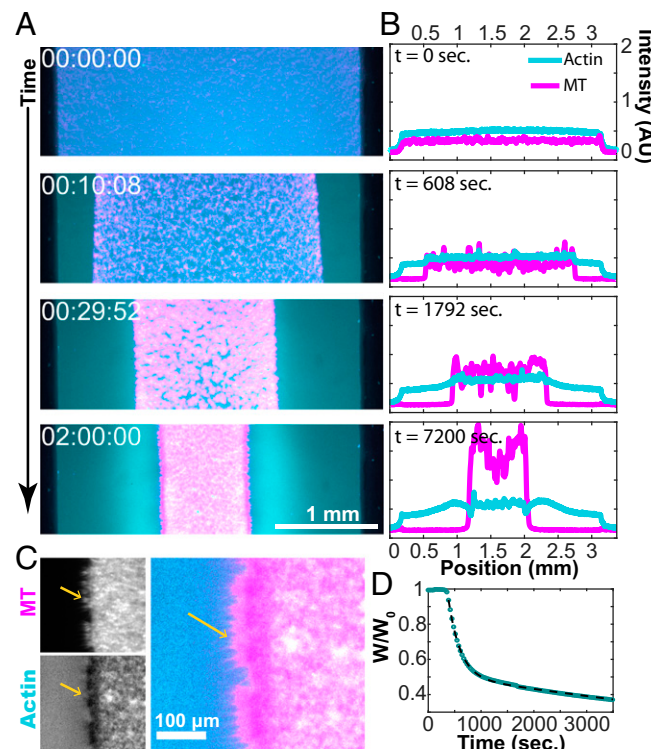


Fig. 8. Bulk contraction at high F-actin concentrations. (A) Dual-color fluorescence micrographs of actin (cyan) and MTs (magenta) show the MT-rich network contract toward the chamber interior (Movie S9). (B) Intensity profiles of fluorescence micrographs show the accumulation of F-actin and MTs during bulk contraction. While nearly all of the MTs are carried in the contracting structure, a significant fraction of F-actin is left behind. (C) MT bundles point outward from the contracted domain. (D) A plot of the normalized width as a function of time. The fit to Eq. 1 yields $c_1 = 203$ s and $c_2 = 3,470$ s, and the lag time $b = 269$ s.

We speculate about a possible mechanism by which the viscoelasticity controls the symmetries of the active stresses. Extensile fluids are driven by a cyclical process of motor-driven MT bundle extension, buckling, fraying, and reannealing (11). The extension requires bundles with mixed polarity, in which the MT plus ends are uniformly distributed and are equally likely to point in both directions. Molecular motors quickly sort an isolated bundle, yielding static spatial domains, within which all MT plus ends point in the same direction. It is likely that the cyclical network reconfiguration continuously mixes the polar domains with the MT bundles, which in turn, sustains the steady-state generation of the extensile stresses and the associated nonequilibrium turbulent dynamics. It is possible that the viscoelastic actin network sterically suppresses the buckling and the reannealing of MT bundles. The suppression of such dynamical modes could quickly generate fragmented and polarity-sorted MTs domains. Such architectures can no longer sustain extensile stresses. However, an MT network could exhibit contractile dynamics through motor end binding or some other mechanisms (15).

Active Contractile Dynamics Generate Layered Aster Composites. We also showed that the extensile to contractile transition is associated with the formation of layered asters. During the initial stages of formation, we observed a system-spanning network of threads that yielded and rearranged to form spherical filament-rich asters. Intriguingly, similar morphological evolution is observed in the passive phase separating viscoelastic polymer mixtures, but in the latter case, each phase is enriched in one polymer type (36, 58). In the active system, both filaments separated into one phase whose composition was not governed by thermodynamics. On longer timescales, we observed the slow differential motion of the various components driving the onion-like layering (Fig. 6).

Single-filament asters have been studied previously (40, 53, 59–66). Their formation and stability are explained by invoking the preference of molecular motors to move toward and reside at a filament tip (67). The size of conventional asters is usually comparable with the filament length, and the radial organization extends from the motor-rich core to their periphery. In contrast, layered asters have an outer cortex that is many times the length of the constituent MTs. Furthermore, the cortex envelops a sizeable region of isotropically organized filaments. The mechanisms that determine the size of the cortex and the asters' actin-rich core remain unknown.

Structured asters and bulk contracted composites share certain structural features. Notably, bulk contractions yield a monolayer of dense MTs, which emerge at the very edge of the contracting structure. This layer is reminiscent of the MT-rich cortex observed in layered asters. Both layers exclude F-actin, while the MT bundles point along the surface normal. Similar edge-bound structures have also been observed in other

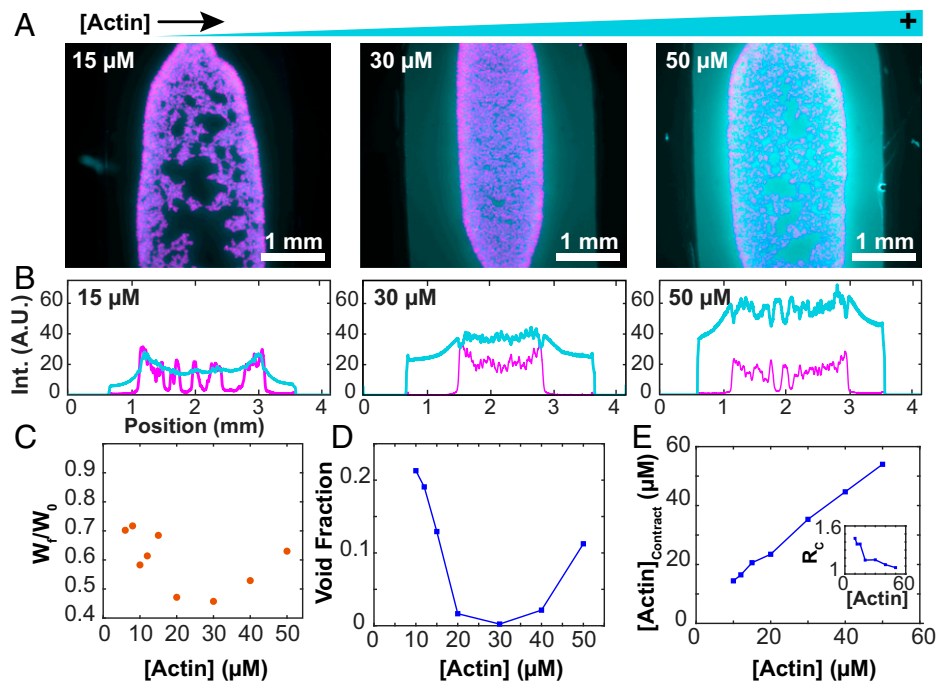


Fig. 9. Bulk contraction changes nonmonotonically with F-actin concentration. (A) Fluorescence micrographs of bulk-contracted domains at various F-actin concentrations. Images represent the densest structures formed. (B) Profiles of the intensity of the actin and MT signals across the sample chamber. (C) The normalized width of the final contracted structures varies nonmonotonically with the F-actin concentration. (D) The void fraction of MTs within the contracted structures depends on the F-actin concentration. (E) Increasing overall F-actin concentration increases the concentration of F-actin within the contracted domains. (Inset) The ratio of F-actin concentration in and outside the contracted domains.

contractile systems (68, 69). Thus, they might be a ubiquitous yet still poorly understood feature of bulk contractions. Understanding the molecular mechanisms that lead to their formation might provide insight into the formation of layered asters.

Structured Asters Have a Finite Lifetime. Another poorly understood feature of layered asters is their finite and programmable lifetime. The intrinsic length scales and timescales of kinesin-1 motors moving on MTs are micrometers and seconds, respectively (47, 70). By comparison, the asters remain stable for tens

of minutes to hours. Imbuing synthetic self-assembly processes with a finite lifetime remains a challenge, especially when compared with cells that rapidly assemble large-scale structures, such as a mitotic spindle or a cytokinetic ring, at a predetermined location for a predetermined time before equally rapidly disassembling them. Finite-lifetime asters have also been observed but not emphasized in single-filament systems, suggesting that their transitory nature might be a ubiquitous phenomenon (35, 71). Understanding the mechanisms that govern the asters' finite lifetimes might reveal more broadly applicable strategies for controlling temporal dynamics.

One can only speculate about a possible mechanism that controls asters lifetime. The MT cortices likely have polar order. Thus, they act as a highway for motor clusters to move inward, accumulating in the aster core. The buildup of motor proteins beyond a critical concentration could destabilize the aster. Furthermore, by linking multiple MTs, the inward-moving motor clusters could contribute to the stability of the outer monolayer cortex and the entire layered aster. Exploring the role of PRC1 cross-linkers, which specifically bundle anti-parallel MTs, in the stability of the asters and the outer MT cortex remains a topic for future exploration.

Liquid–liquid phase separation (LLPS) has emerged as a key organizing principle in cell biology (72, 73). The surface area minimization and coalescence of active composite asters are evocative of liquid-like phenomena. However, there are important differences. Layered aster formation is driven not by thermodynamic forces but by the action of molecular motors. Furthermore, the core of the layered aster is a likely viscoelastic material that will appear as liquid or solid at different timescales (45). Nevertheless, our observations raise intriguing questions at the nexus of the emerging areas of active matter, phase separation in cell biology, and biophysics of cytoskeleton. From the perspective of cell biology, our findings should promote studies that address if and how a combination of cytoskeletal fibers and

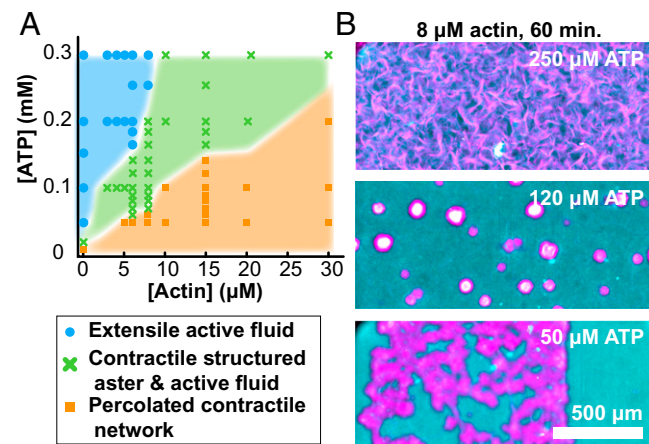


Fig. 10. Dynamics phase diagram of actin–MT active composites. (A) State diagram representing the self-organized structure observed after 1 h of activity as a function of ATP and F-actin concentration. Bulk contraction is characterized by the retraction of the sample edges before self-tearing into localized structures. (B) Representative merged fluorescence micrographs of the active composite demonstrating self-organized structure at fixed actin concentration while varying ATP concentration (MT, magenta; F-actin, cyan).

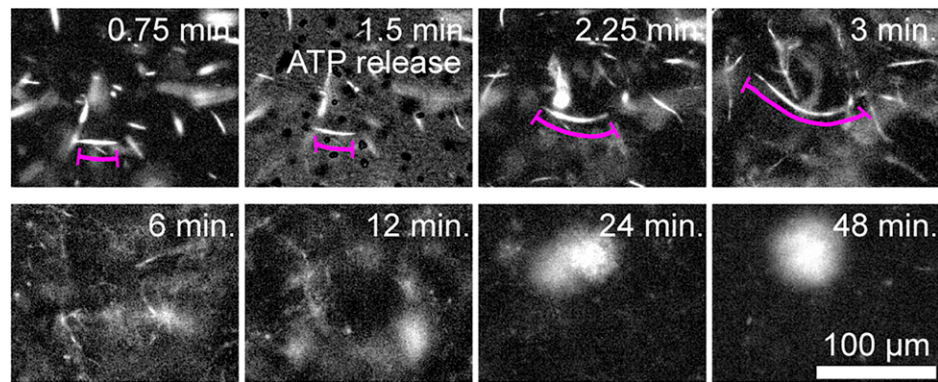


Fig. 11. Bulk contractions are driven by extensile MT bundles. Time-lapse images showing initial extension of fluorescently labeled MTs in a composite network. At longer times, bulk contraction is observed (100 μ M caged ATP and 8 μ M actin).

nonequilibrium fluctuations can promote LLPS (45, 74). From the active matter perspective, some observed phenomenology is reminiscent of viscoelastic phase separation observed in the passive binary polymer mixtures (36). Further experiments are necessary to determine if there are any fundamental similarities between our observations and the classical phase separation and the role active advection could play in controlling timescales of phase separation (75). More broadly, recent advances demonstrated that active swimmers in a viscoelastic background can coordinate their motility into a large-scale oscillatory dynamical state (76). Adding viscoelastic responses to active matter thus is an important direction that can help to address questions about how disordered microscopic stresses can self-organize into desired dynamical structures.

Our work revealed the formation of intricate spatiotemporal patterns through mechanical interactions between actin filaments and MTs alone. Understanding the mechanism governing our simplified systems might give insight into several phenomena in cell biology that involve interplay between MTs and the actin cytoskeleton. Among others, these include MT penetration into filopodia, lamellipodia, and growth cones; MT “tracking” along actin stress fibers; meiotic MT spindles that are surrounded by and embedded in the actin network; MTs growing out of centrosomes, which are also actin-organizing centers; and the intricate interplay of actin and MT cytoskeleton that controls the onset of cytoplasmic streaming in the *Drosophila* oocyte (77–81). In the last example, the onset of cytoplasmic streaming is determined by depolymerization of the viscoelastic actin network. Intriguingly, this is a ubiquitous feature of reconstituted active/MT composites studied here. Finally, we note that the segregation of the actin, MTs, and motors into layered asters with timescales controlled by physical properties has similarities to cell extract systems, which can form membrane-less cell-like compartments whose internal organization is coordinated in time without genetically controlled mechanisms (82).

In summary, we showed that MT–actin composites exhibit distinct organized states that evolve. The coexisting dynamical structures are reminiscent of the multiple physiological states observed in cell biology. Elucidating the governing equation of layered asters and the extensile fluids represents a challenge for existing theoretical formalisms of active matter. Answering this challenge might enable real-time control of coexisting out-of-equilibrium states, a hallmark of dynamical and adaptable biological cells.

Materials and Methods

MTs. Short fluorescently labeled guanyl-(alpha,beta)-methylene-diphosphonate (GMPCPP)-stabilized MTs were generated by established methods (11). Briefly, recycled Alexa-647-labeled and unlabeled tubulin monomers were polymerized in excess GMPCPP to yield short fluorescent MTs with a labeling

efficiency of about 1%. Small aliquots were snap frozen in 80 mM 1,4-piperazinediethanesulfonic acid and 2 mM magnesium chloride, pH 7.4 (M2B), at 8 mg/mL and used throughout the experiment.

Kinesin Motor Clusters. Short biotinylated kinesin constructs were prepared via established protocols (11). Simply, heavy chain kinesin-1 truncated at residue 401 and fused to biotin carboxyl carrier protein was expressed in *Escherichia coli* cells, harvested, and purified via fast protein liquid chromatography. These biotinylated motors were snap frozen in M2B and used throughout the experiment. To generate motor clusters, the biotinylated motors were incubated in a 2:1 ratio with streptavidin tetramers. In the case of fluorescently labeled motor clusters, Dylight 488 neutravidin was used in place of streptavidin.

PRC1-NSΔC. PRC1-NSΔC, a truncated version of the MT cross-linker, protein regulator of cytokinesis 1 (PRC1), was expressed in and purified from *E. coli*. The nonstructured region at the C terminus of the PRC1 protein is deleted, which reduces the interactions between adjacent cross-linkers. These proteins were snap frozen in 50 mM sodium phosphate buffer, pH 7.0, 400 mM imidazole, 150 mM sodium chloride, 5 mM tris(2-carboxyethyl)phosphine, and 2 mM ethylenediaminetetraacetic acid as an aminopolycarboxylic acid as described previously (27).

Actin. Rabbit muscle actin was purified as previously described (83). The nucleotides in the G-actin solution were then replaced by AMP-PNP using previous protocols (84). Briefly, G-actin solution was passed through ion exchange resin to bind free nucleotides directly into buffers prepared with AMP-PNP, a nonhydrolyzable ATP analog. G-actin solutions (~150 μ M) were snap frozen in 5 mM tris(hydroxymethyl)aminomethane-HCl, pH 8.0, 0.2 mM CaCl_2 , 0.3 mM AMP-PNP, and 0.5 mM dithiothreitol.

Chambers. Flow cells in which the experiment took place were created by sandwiching Parafilm between acrylamide brush-coated glass slide and coverslips. The Parafilm is then briefly raised above the melting temperature, creating a sealed chamber with inlets, outlets, and a size of about 10 \times 30 \times 0.1 mm. After loading the experimental solutions, the inlet and outlet are sealed with silicone vacuum grease.

Experiment. A bundled MT network was prepared in solution with actin monomers, loaded into chambers, sealed, and polymerized over 2 h in darkness. Then, UV light was shown on the sample, releasing caged ATP and allowing kinesin activity to commence. The bundled MT network was formed by combining formed GMPCPP MTs, PRC1-NSΔC, kinesin motor clusters, and caged ATP under red light emitting diode light. This solution also contains an ATP regeneration system (21 units/mL pyruvate kinase, 40 mM phosphoenolpyruvate) and antioxidants (18.7 mM glucose, 1.4 μ M glucose oxidase, 0.17 μ M catalase). To this, the appropriate amount of actin is added, mixed, and loaded into the experimental chamber, which is promptly sealed. After 2 h of actin polymerization, activity is initiated by exposing the sample to UV light (365 nm) for 30 s. The experimental slides can hold up to 12 parallel chambers, allowing for experiments to probe the effects of the varied parameter (e.g., actin concentration) on the same experimental preparation.

Microscopes. The samples were observed on a Nikon Ti microscope with 4x, 10x, and 20x objectives and Andor Neo or FLIR Blackfly cameras controlled by Micromanager. Confocal microscopy was performed using a Leica SP8 confocal microscope.

Data Availability. All study data are included in the article and/or supporting information.

ACKNOWLEDGMENTS. We thank Bulbul Chakraborty, Moumita Das, Margaret Gardel, Chase Broedersz, Andreas Bausch, Gijsje Koenderink, Michael

- V. E. Deneke et al., Self-organized nuclear positioning synchronizes the cell cycle in *Drosophila* embryos. *Cell* **177**, 925–941.e17 (2019).
- V. Ruprecht et al., Cortical contractility triggers a stochastic switch to fast amoeboid cell motility. *Cell* **160**, 673–685 (2015).
- J. Sedzinski et al., Polar actomyosin contractility destabilizes the position of the cyto-kinetic furrow. *Nature* **476**, 462–466 (2011).
- M. C. Marchetti et al., Hydrodynamics of soft active matter. *Rev. Mod. Phys.* **85**, 1143 (2013).
- J. Brugués, D. Needleman, Physical basis of spindle self-organization. *Proc. Natl. Acad. Sci. U.S.A.* **111**, 18496–18500 (2014).
- M. Mayer, M. Depken, J. S. Bois, F. Jülicher, S. W. Grill, Anisotropies in cortical tension reveal the physical basis of polarizing cortical flows. *Nature* **467**, 617–621 (2010).
- J. Palacci, S. Sacanna, A. P. Steinberg, D. J. Pine, P. M. Chaikin, Living crystals of light-activated colloidal surfers. *Science* **339**, 936–940 (2013).
- I. Theurkauff, C. Cottin-Bizonne, J. Palacci, C. Ybert, L. Bocquet, Dynamic clustering in active colloidal suspensions with chemical signaling. *Phys. Rev. Lett.* **108**, 268303 (2012).
- V. Schaller, C. Weber, C. Semmrich, E. Frey, A. R. Bausch, Polar patterns of driven filaments. *Nature* **467**, 73–77 (2010).
- C. Dombrowski, L. Cisneros, S. Chatkaew, R. E. Goldstein, J. O. Kessler, Self-concentration and large-scale coherence in bacterial dynamics. *Phys. Rev. Lett.* **93**, 098103 (2004).
- T. Sanchez, D. T. Chen, S. J. DeCamp, M. Heymann, Z. Dogic, Spontaneous motion in hierarchically assembled active matter. *Nature* **491**, 431–434 (2012).
- S. Zhou, A. Sokolov, O. D. Lavrentovich, I. S. Aranson, Living liquid crystals. *Proc. Natl. Acad. Sci. U.S.A.* **111**, 1265–1270 (2014).
- J. Alvarado, M. Sheinman, A. Sharma, F. C. MacKintosh, G. H. Koenderink, Molecular motors robustly drive active gels to a critically connected state. *Nat. Phys.* **9**, 591–597 (2013).
- P. M. Bendaix et al., A quantitative analysis of contractility in active cytoskeletal protein networks. *Biophys. J.* **94**, 3126–3136 (2008).
- P. J. Foster, S. Fürthauer, M. J. Shelley, D. J. Needleman, Active contraction of microtubule networks. *eLife* **4**, e10837 (2015).
- S. Köhler, V. Schaller, A. R. Bausch, Structure formation in active networks. *Nat. Mater.* **10**, 462–468 (2011).
- M. P. Murrell, M. L. Gardel, F-actin buckling coordinates contractility and severing in a biomimetic actomyosin cortex. *Proc. Natl. Acad. Sci. U.S.A.* **109**, 20820–20825 (2012).
- S. Fürthauer, M. J. Shelley, How crosslink numbers shape the large-scale physics of cytoskeletal materials. *arXiv [Preprint]* (2021). <https://arxiv.org/abs/2106.13273> (Accessed 12 January 2022).
- D. Needleman, Z. Dogic, Active matter at the interface between materials science and cell biology. *Nat. Rev. Mater.* **2**, 1–14 (2017).
- J. Roostalu, J. Rickman, C. Thomas, F. Nédélec, T. Surrey, Determinants of polar versus nematic organization in networks of dynamic microtubules and mitotic motors. *Cell* **175**, 796–808.e14 (2018).
- M. L. Gardel, M. T. Valentine, J. C. Crocker, A. R. Bausch, D. A. Weitz, Microrheology of entangled F-actin solutions. *Phys. Rev. Lett.* **91**, 158302 (2003).
- P. A. Janmey et al., The mechanical properties of actin gels. Elastic modulus and filament motions. *J. Biol. Chem.* **269**, 32503–32513 (1994).
- D. A. Gagnon et al., Shear-induced gelation of self-yielding active networks. *Phys. Rev. Lett.* **125**, 178003 (2020).
- S. Asakura, F. Oosawa, On interaction between two bodies immersed in a solution of macromolecules. *J. Chem. Phys.* **22**, 1255–1256 (1954).
- F. Hiltitschi et al., Measuring cohesion between macromolecular filaments one pair at a time: Depletion-induced microtubule bundling. *Phys. Rev. Lett.* **114**, 138102 (2015).
- P. Chandrakar et al., Microtubule-based active fluids with improved lifetime, temporal stability and miscibility with passive soft materials. *arXiv [Preprint]* (2018). <https://arxiv.org/abs/1811.05026> (Accessed 12 January 2022).
- R. Subramanian et al., Insights into antiparallel microtubule crosslinking by PRC1, a conserved nonmotor microtubule binding protein. *Cell* **142**, 433–443 (2010).
- M. Levin et al., Kinetics of actin networks formation measured by time resolved particle-tracking microrheology. *Soft Matter* **16**, 7869–7876 (2020).
- S. Z. Chou, T. D. Pollard, Mechanism of actin polymerization revealed by cryo-EM structures of actin filaments with three different bound nucleotides. *Proc. Natl. Acad. Sci. U.S.A.* **116**, 4265–4274 (2019).
- H. Higuchi, E. Muto, Y. Inoue, T. Yanagida, Kinetics of force generation by single kinesin molecules activated by laser photolysis of caged ATP. *Proc. Natl. Acad. Sci. U.S.A.* **94**, 4395–4400 (1997).
- G. Henkin, S. J. DeCamp, D. T. Chen, T. Sanchez, Z. Dogic, Tunable dynamics of microtubule-based active isotropic gels. *Philos. Trans. A Math. Phys. Eng. Sci.* **372**, 20140142 (2014).
- G. Duclos et al., Topological structure and dynamics of three-dimensional active nematics. *Science* **367**, 1120–1124 (2020).
- M. Soares e Silva et al., Active multistage coarsening of actin networks driven by myosin motors. *Proc. Natl. Acad. Sci.* **108**, 9408–9413 (2011).
- T. D. Ross et al., Controlling organization and forces in active matter through optimally defined boundaries. *Nature* **572**, 224–229 (2019).
- V. Wollrab et al., Polarity sorting drives remodeling of actin-myosin networks. *J. Cell Sci.* **132**, jcs219717 (2018).
- H. Tanaka, Viscoelastic phase separation. *J. Phys. Condens. Matter* **12**, R207 (2000).
- A. J. Bray, Theory of phase-ordering kinetics. *Adv. Phys.* **51**, 481–587 (2002).
- M. E. Cates, E. Tjhung, Theories of binary fluid mixtures: From phase-separation kinetics to active emulsions. *J. Fluid Mech.* **836**, P1 (2018).
- E. D. Siggia, Late stages of spinodal decomposition in binary mixtures. *Phys. Rev. A* **20**, 595 (1979).
- T. Surrey, F. Nédélec, S. Leibler, E. Karsenti, Physical properties determining self-organization of motors and microtubules. *Science* **292**, 1167–1171 (2001).
- J. Alvarado, L. Cipelletti, G. H. Koenderink, Uncovering the dynamic precursors to motor-driven contraction of active gels. *Soft Matter* **15**, 8552–8565 (2019).
- R. Zhang, N. Kumar, J. L. Ross, M. L. Gardel, J. J. de Pablo, Interplay of structure, elasticity, and dynamics in actin-based nematic materials. *Proc. Natl. Acad. Sci. U.S.A.* **115**, E124–E133 (2018).
- G. Lee et al., Myosin-driven actin-microtubule networks exhibit self-organized contractile dynamics. *Sci. Adv.* **7**, eabe4334 (2021).
- T. Strübing et al., Wrinkling instability in 3D active nematics. *Nano Lett.* **20**, 6281–6288 (2020).
- D. Humphrey, C. Duggan, D. Saha, D. Smith, J. Käs, Active fluidization of polymer networks through molecular motors. *Nature* **416**, 413–416 (2002).
- A. Senoussi et al., Tunable corrugated patterns in an active nematic sheet. *Proc. Natl. Acad. Sci. U.S.A.* **116**, 22464–22470 (2019).
- M. J. Schnitzer, S. M. Block, Kinesin hydrolyses one ATP per 8-nm step. *Nature* **388**, 386–390 (1997).
- L. M. Lemma, S. J. DeCamp, Z. You, L. Giomi, Z. Dogic, Statistical properties of autonomous flows in 2D active nematics. *Soft Matter* **15**, 3264–3272 (2019).
- L. Huber, R. Suzuki, T. Krüger, E. Frey, A. R. Bausch, Emergence of coexisting ordered states in active matter systems. *Science* **361**, 255–258 (2018).
- L. Farhadi, C. Fermino Do Rosario, E. P. Debold, A. Baskaran, J. L. Ross, Active self-organization of actin-microtubule composite self-propelled rods. *Front. Phys.* **6**, 75 (2018).
- G. H. Koenderink et al., An active biopolymer network controlled by molecular motors. *Proc. Natl. Acad. Sci. U.S.A.* **106**, 15192–15197 (2009).
- S. Stam et al., Filament rigidity and connectivity tune the deformation modes of active biopolymer networks. *Proc. Natl. Acad. Sci. U.S.A.* **114**, E10037–E10045 (2017).
- C. Hentrich, T. Surrey, Microtubule organization by the antagonistic mitotic motors kinesin-5 and kinesin-14. *J. Cell Biol.* **189**, 465–480 (2010).
- J. M. Belmonte, M. Leptin, F. Nédélec, A theory that predicts behaviors of disordered cytoskeletal networks. *Mol. Syst. Biol.* **13**, 941 (2017).
- M. Lenz, Reversal of contractility as a signature of self-organization in cytoskeletal bundles. *eLife* **9**, e51751 (2020).
- M. Scholz, K. L. Weirich, M. L. Gardel, A. R. Dinner, Tuning molecular motor transport through cytoskeletal filament network organization. *Soft Matter* **16**, 2135–2140 (2020).
- P. Ronceray, C. P. Broedersz, M. Lenz, Fiber networks amplify active stress. *Proc. Natl. Acad. Sci. U.S.A.* **113**, 2827–2832 (2016).
- M. Tateno, H. Tanaka, Power-law coarsening in network-forming phase separation governed by mechanical relaxation. *Nat. Commun.* **12**, 912 (2021).
- F. J. Nédélec, T. Surrey, A. C. Maggs, S. Leibler, Self-organization of microtubules and motors. *Nature* **389**, 305–308 (1997).
- C. J. Miller, D. Harris, R. Weaver, G. B. Ermentrout, L. A. Davidson, Emergent mechanics of actomyosin drive punctuated contractions and shape network morphology in the cell cortex. *PLoS Comput. Biol.* **14**, e1006344 (2018).
- A. Das et al., Stratification relieves constraints from steric hindrance in the generation of compact actomyosin asters at the membrane cortex. *Sci. Adv.* **6**, eay6093 (2020).
- L. Haviv et al., Reconstitution of the transition from lamellipodium to filopodium in a membrane-free system. *Proc. Natl. Acad. Sci. U.S.A.* **103**, 4906–4911 (2006).
- Y. Ideses et al., Spontaneous buckling of contractile poroelastic actomyosin sheets. *Nat. Commun.* **9**, 2461 (2018).
- Y. Ideses, A. Sonn-Segev, Y. Roichman, A. Bernheim-Groswasser, Myosin II does it all: Assembly, remodeling, and disassembly of actin networks are governed by myosin II activity. *Soft Matter* **9**, 7127–7137 (2013).
- T. Torisawa, D. Taniguchi, S. Ishihara, K. Oiwa, Spontaneous formation of a globally connected contractile network in a microtubule-motor system. *Biophys. J.* **111**, 373–385 (2016).
- A. B. Verkhovsky, T. M. Svitkina, G. G. Borisy, Polarity sorting of actin filaments in cytochalasin-treated fibroblasts. *J. Cell Sci.* **110**, 1693–1704 (1997).

67. F. Nédélec, T. Surrey, Dynamics of microtubule aster formation by motor complexes. *C. R. Acad. Sci. Ser. IV Phys. Astrophys.* **2**, 841–847 (2001).

68. B. Lemma, N. P. Mitchell, R. Subramanian, D. J. Needleman, Z. Dogic, Active microphase separation in mixtures of microtubules and tip-accumulating molecular motors. arXiv [Preprint] (2021). <https://arxiv.org/abs/2107.12281> (Accessed 12 January 2022).

69. T. J. Mitchison, P. Nguyen, M. Coughlin, A. C. Groen, Self-organization of stabilized microtubules by both spindle and midzone mechanisms in *Xenopus* egg cytosol. *Mol. Biol. Cell* **24**, 1559–1573 (2013).

70. K. S. Thorn, J. A. Ubersax, R. D. Vale, Engineering the processive run length of the kinesin motor. *J. Cell Biol.* **151**, 1093–1100 (2000).

71. F. Backouche, L. Haviv, D. Groswasser, A. Bernheim-Groswasser, Active gels: Dynamics of patterning and self-organization. *Phys. Biol.* **3**, 264–273 (2006).

72. C. P. Brangwynne, P. Tompa, R. V. Pappu, Polymer physics of intracellular phase transitions. *Nat. Phys.* **11**, 899–904 (2015).

73. A. A. Hyman, C. A. Weber, F. Jülicher, Liquid-liquid phase separation in biology. *Annu. Rev. Cell Dev. Biol.* **30**, 39–58 (2014).

74. T. Wiegand, A. A. Hyman, Drops and fibers: How biomolecular condensates and cytoskeletal filaments influence each other. *Emerg. Top. Life Sci.* **4**, 247–261 (2020).

75. R. Singh, M. E. Cates, Hydrodynamically interrupted droplet growth in scalar active matter. *Phys. Rev. Lett.* **123**, 148005 (2019).

76. S. Liu, S. Shankar, M. C. Marchetti, Y. Wu, Viscoelastic control of spatiotemporal order in bacterial active matter. *Nature* **590**, 80–84 (2021).

77. F. Farina *et al.*, The centrosome is an actin-organizing centre. *Nat. Cell Biol.* **18**, 65–75 (2016).

78. J. Roeles, G. Tsiavaliaris, Actin-microtubule interplay coordinates spindle assembly in human oocytes. *Nat. Commun.* **10**, 4651 (2019).

79. B. Mogessie, M. Schuh, Actin protects mammalian eggs against chromosome segregation errors. *Science* **357**, eaal1647 (2017).

80. M. Dogterom, G. H. Koenderink, Actin-microtubule crosstalk in cell biology. *Nat. Rev. Mol. Cell Biol.* **20**, 38–54 (2019).

81. M. E. Quinlan, Cytoplasmic streaming in the *Drosophila* oocyte. *Annu. Rev. Cell Dev. Biol.* **32**, 173–195 (2016).

82. X. Cheng, J. E. Ferrell Jr., Spontaneous emergence of cell-like organization in *Xenopus* egg extracts. *Science* **366**, 631–637 (2019).

83. S. Shekhar, G. J. Hoeprich, J. Gelles, B. L. Goode, Twinfilin bypasses assembly conditions and actin filament aging to drive barbed end depolymerization. *J. Cell Biol.* **220**, e202006022 (2021).

84. H. I. Balcer *et al.*, Coordinated regulation of actin filament turnover by a high-molecular-weight Srv2/CAP complex, cofilin, profilin, and Aip1. *Curr. Biol.* **13**, 2159–2169 (2003).