Kinesin and myosin motors compete to drive rich multiphase dynamics in programmable cytoskeletal composites

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Abstract

The cellular cytoskeleton relies on diverse populations of motors, filaments, and binding proteins acting in concert to enable nonequilibrium processes ranging from mitosis to chemotaxis. The cytoskeleton’s versatile reconfigurability, programmed by interactions between its constituents, makes it a foundational active matter platform. However, current active matter endeavors are limited largely to single force-generating components acting on a single substrate—far from the composite cytoskeleton in cells. Here, we engineer actin–microtubule (MT) composites, driven by kinesin and myosin motors and tuned by crosslinkers, to ballistically restructure and flow with speeds that span three orders of magnitude depending on the composite formulation and time relative to the onset of motor activity. Differential dynamic microscopy analyses reveal that kinesin and myosin compete to delay the onset of acceleration and suppress discrete restructuring events, while passive crosslinking of either actin or MTs has an opposite effect. Our minimal advection–diffusion model and spatial correlation analyses correlate these dynamics to structure, with motor antagonism suppressing reconfiguration and demixing, while crosslinking enhances clustering. Despite the rich formulation space and emergent formulation-dependent structures, the nonequilibrium dynamics across all composites and timescales can be organized into three classes—slow isotropic reorientation, fast directional flow, and multimode restructuring. Moreover, our mathematical model demonstrates that diverse structural motifs can arise simply from the interplay between motor-driven advection and frictional drag. These general features of our platform facilitate applicability to other active matter systems and shed light on diverse ways that cytoskeletal components can cooperate or compete to enable wide-ranging cellular processes.

Significance Statement

The cytoskeleton is a paradigmatic active matter system—comprising protein filaments, motors, and crosslinkers—that mediates wide-ranging cellular processes from migration to morphogenesis. The cytoskeleton’s composite nature, conferring versatility and programmability, is one of its hallmarks. Yet, state-of-the-art active matter designs are largely limited to single force-generating components and substrates. Here, we engineer composites of MTs and actin driven by kinesin and myosin motors to restructure, contract, and flow to form structures ranging from interpenetrating scaffolds to phase-separated clusters. Surprisingly, kinesin and myosin compete to delay rapid restructuring and suppress demixing. Our bioinspired nonequilibrium composites not only bring reconstituted systems a critical step closer to mimicking cytoskeletal complexity but are also foundational for diverse material applications from wound healing to soft robotics.

Introduction

The cytoskeleton is a dynamic, nonequilibrium material comprising protein filaments, including actin, microtubules (MTs), and intermediate filaments, as well as motor proteins, such as myosins and kinesins, that actively push and pull on the protein filaments (1–8). Crosslinking proteins also connect and bundle filaments as needed for cellular processes (9–12). This complex composite continues to restructure and reconfigures in response to demands of the cell to enable diverse processes from cytokinesis to mechanosensing (3–5, 7, 8, 13–21). In vitro systems of reconstituted cytoskeletal proteins, which display rich and tunable dynamics, are also intensely studied as model active matter...
platforms to shed light on the nonequilibrium physics underlying force-generating, reconfigurable systems (7, 12, 19, 22–40).

Interacting networks of semiflexible actin filaments and rigid MTs provide tensile and compressive strength to the cytoskeleton while allowing for cell mobility, key to processes such as division and chemotaxis (15, 16, 41–45). Further, recent studies have shown that in vitro actin–MT composites exhibit emergent mechanical properties that are not a simple sum of the single component systems (46–48). For example, composites with comparable concentrations of actin and MTs display both enhanced filament mobility and increased stiffness (46), as well as an emergent nonmonotonic dependence of elasticity on actin crosslinking (47).

More recently, myosin II minifilaments have been incorporated into actin–MT composites, showing that synergistic interactions between actin and MTs prevent disordered flow and rupturing often seen in actomyosin networks without crosslinkers (26–28). These studies have also shown that composites can be tuned to display enhanced mechanical strength (27), coordinated motion of actin and MTs, sustained ballistic contraction, and mesoscale restructuring (26, 28)—all in the absence of crosslinking proteins to chemically connect the filaments.

Microtubule-based active matter systems have also been engineered using clusters of kinesin motors that crosslink and pull on microtubule bundles to create active nematics (23, 24, 30, 31, 34, 35, 49–55). In these systems, kinesins generate long-lasting turbulent flows by cyclically extending, buckling, fracturing, and healing bundles (49). More recently, actin has been incorporated into active microtubule fluids, resulting in turbulent flow, contraction, or formation of layered asters (29).

The distinct dynamics and structures that kinesin-driven and myosin-driven systems display beg the question as to how different active components and substrates cooperate or compete to control cellular processes. While composite active matter is beginning to be developed to introduce more control and tunability over single-substrate systems (26–29, 56), incorporating two active components that act on distinct substrates represents a paradigm shift in active matter. Beyond the cellular relevance, such designs can elucidate general principles for nonequilibrium programmable materials that can reconfigure and generate force and determine how to enhance programmability and expand the dynamical and structural phase space by altering the active and static nature of crosslinkers and the substrates on which they act.

Here, we engineer actin–MT composites that undergo a rich combination of advective flow, contraction, and multimode restructuring driven by kinesin and myosin motors. These dynamics are coupled to distinct time-evolving structures that range from interpenetrating actin–MT scaffolds to microscale phase-separated amorphous clusters. We couple differential dynamic microscopy (DDM) with particle image velocimetry (PIV) to discover that competition between kinesin–MT activity and actomyosin activity delays the onset of rapid restructuring while crosslinking of either actin or MTs accelerates the time evolution of active dynamics. Our advection–diffusion model and spatial correlation analyses correlate the dynamic antagonism that we observe with suppressed demixing of double-motor composites and the crosslinker-mediated acceleration with enhanced restructuring and clustering. Despite these complexities, we find that the broad phase space of active dynamics can be organized into three general classes with distinct types and rates of ballistic motion.

Results and discussion

Active cytoskeleton composite design and formulation–structure phase space

We first describe our design strategy for realizing an active matter system that has two force-generating components that act on two distinct, yet homogeneously co-mixed, substrates. Namely, we engineer composites of co-entangled MTs and actin (46) and incorporate kinesin clusters and myosin II minifilaments that crosslink and push and pull on pairs of MTs and actin, respectively, to generate force and motion (49, 57) (Fig. 1A, Fig. S1). To investigate the extent to which actomyosin and kinesin–MT activities act synergistically or antagonistically to dictate dynamics, we perform experiments with either kinesin (K, Fig. 1B) or both kinesin and myosin (K + M, Fig. 1B). We further characterize the effect of passive crosslinking of the MTs (MT XL, Fig. 1) or actin (Actin XL, Fig. 1) at crosslinker-protein molar ratios R that are high enough to induce measurable changes in the viscoelastic properties but low enough to prevent filament bundling (47). To observe the dynamics of our active cytoskeleton composites, we collect sequential two-color time-series of actin and MTs comprising composites over the ~1-h time course of measurable active dynamics. As shown in Fig. 1B, by simply incorporating or omitting myosin motors and passive crosslinkers, we are able to drive substantial changes in the active restructuring, emergent dynamics, and programmable phase space of non-equilibrium properties (Movies S1–S3).

All composites begin in similar structural states with interpenetrating networks of actin and MTs uniformly distributed throughout the field of view (Fig. 1B, top row). However, each composite formulation reconfigures into distinct structural states over activity times of T ≈ 30 min, where T = 0 is defined as the time at which kinesin is added to the composite. While we do not visualize the motors, the spatially uniform active dynamics that we see at the onset of activity indicate that, like the filaments, the motors are uniformly mixed throughout the composite.

Examining the three kinesin-only composites (no myosin), we find that without passive crosslinkers, composites form loosely connected MT-rich amorphous clusters. Actin filaments first co-localize in the cluster centers but are then squeezed out into the surrounding space as the clusters contract further and disconnect from one another (Fig. 1B, dark blue boxes). Passive actin crosslinking hinders this separation of actin and MTs, enabling the slow uptake of actin into MT-rich clusters, such that the composite becomes a connected network of clusters of co-localized actin and MTs (Fig. 1B, dark green boxes). MT crosslinking leads to similar amorphous MT clustering and actin–MT de-mixing without crosslinking, but these MT-rich regions coalesce over time, resulting in larger-scale phase separation of actin and MTs compared to the non-crosslinked case (Fig. 1B, dark red boxes).

Turning to the double-motor composites that incorporate both kinesin and myosin, we find that the addition of myosin impedes the kinesin-driven de-mixing described above and reduces the degree of restructuring over the course of activity (Fig. 1B, light shaded boxes). This effect can be seen in the images at all time points (rows in Fig. 1B), in which actin and MT networks are more evenly distributed and interpenetrating than composites without myosin. Without passive crosslinkers, composites show little rearrangement (Fig. 1B, light blue boxes), as seen in previous experiments on myosin-driven actin–MT composites (26–28). Crosslinking of actin or MTs enables more restructuring of the double-motor composites, but this reconfiguration and demixing
is still more subdued than that for kinesin-only composites (Fig. 1B, light green and red boxes).

**Actin and MTs exhibit three distinct classes of coordinated ballistic dynamics**

To determine the non-equilibrium dynamics that enable this rich formulation-dependent restructuring, i.e., how the system gets from one structural state to another, we perform DDM on the actin and MT channels of each two-channel (i.e., two-color) video. As we move from one structural state to another, we perform DDM on the actin and MTs to achieve similar distances d between crosslinks (48). We incorporate kinesin clusters and myosin II minifilaments to drive composites out of steady-state. (B) We acquire two-color confocal time-series of actin (green) and MTs (red) to capture motor-driven dynamics and reconfiguration. Each column includes images taken at three different time points, T = 5, 20, and 35 min, during motor activity for composites with kinesin (K, columns 1, 3, and 5), kinesin and myosin (K + M, columns 2, 4, and 6), no crosslinking (No XL, columns 1 and 2), actin crosslinking (Actin XL, columns 3 and 4), and MT crosslinking (MT XL, columns 5 and 6). Below each composite image is a zoom-in of a 25 µm x 25 µm region denoted by a dashed-line box in the main image and single-channel images showing separately the MTs (middle, red) and actin (right, green). The 50-µm scale bar in the top right panels applies to all full-size images.

**Figure 2A** shows two-dimensional (2D) image structure functions \( D(q_x, q_y, \Delta t) \) for the MT and actin channels of each two-color video that represents a different type of dynamic we measure, which we describe below. The plots in the left and right columns correspond to \( D(q_x, q_y) \) for sample “short” and “long” lag times, \( \Delta t = 3 \) and \( \Delta t = 20 \text{ s} \), and the color is set by the value of \( D(q_x, q_y) \), with low (blue) and high (red) values indicating lower and higher correlations, respectively (see Fig. S2 for more \( D(q_x, q_y) \) examples). The first notable feature in Fig. 2A (and Fig. S2) is the similarity in the functional form of \( D(q_x, q_y) \) for actin and MT channels of the same video and lag time, indicating that the actin and MT network dynamics are well coupled despite cases in which we observe large-scale de-mixing (Fig. 1B). The lower magnitudes of D for actin compared to MTs are due to the comparatively lower signal of the actin channel. Moreover, the more uniform \( D(q_x, q_y) \) values seen in the purple-bordered plots labeled “Slow,” as compared to the middle (orange, Fast) and bottom (magenta, Multimode), are indicative of more homogeneous and slow motion, in which fluctuations decorrelate less over a given lag time and over varying length scales (i.e., wave vectors). The modest radial asymmetry seen most clearly in the orange-bordered plots is a sign of anisotropic motion, which we discuss in later sections. Finally, the reduced correlation at \( \Delta t = 20 \text{ s} \) compared to \( \Delta t = 3 \text{ s} \) indicates that the decorrelation timescales are <20 s.

To quantify the dynamics represented in Fig. 2A, we azimuthally average each \( D(q_x, q_y, \Delta t) \) to compute a corresponding one-dimensional (1D) function for each lag time, \( D(q, \Delta t) \). Figure 2B shows sample \( D(q, \Delta t) \) curves for the three videos analyzed in Fig. 2A. We use the distinct functional features of these curves to organize our data for all composite formulations and activity times into three classes: Slow (top), Fast (middle), and Multimode (bottom). Slow curves show a simple slow rise to plateau at large lag times (Fig. 2B, top panel); Fast curves exhibit oscillations in the decorrelation plateau (Fig. 2B, middle panel); and Multimode curves reveal two distinct plateaus at well-separated lag times (Fig. 2B, bottom panel).

These nontrivial functional forms cannot be accurately described by exponential functions typically used in DDM (26, 28, 58, 59), so we instead use a function that assumes Schuls distribution of speeds, as has been used in other ballistic biological applications.
systems such as swimming *Escherichia coli* (60, 61) (see Methods and Supplementary Material). This function captures the oscillatory plateaus seen in the Fast class, and a sum of two Schulz speed distributions accurately captures the two-plateau behavior of the Multimode class.

From the $D(q, \Delta t)$ fits, we extract decay times, $\tau(q)$, which exhibit a power-law dependence on $q$ that further quantifies the type and rate of motion (Fig. 2C). Despite the varied functional forms of the $D(q, \Delta t)$ curves shown in Fig. 2B, the corresponding $\tau(q)$ for each class exhibits power-law scaling of $\tau(q) \sim q^{-1}$, indicative of ballistic motion for both actin and MTs. Similar ballistic-like dynamics have been previously reported for myosin-driven cytoskeleton composites (26, 28). Fitting each $\tau(q)$ curve to the power-law relation $\tau(q) \approx (\langle \mathbf{v} \rangle q)^{-1}$ yields the average speed $\langle \mathbf{v} \rangle$ of each filament type measured over the course of the corresponding video. As listed in Fig. 2C, we find that $\langle \mathbf{v} \rangle$ for the Fast class is $\sim 7\times$ larger than the Slow $\langle \mathbf{v} \rangle$. Fitting the Multimode $D(q, \Delta t)$ data results in two distinct $\tau(q)$ curves with corresponding $\langle \mathbf{v} \rangle$ values that differ 2.4-fold, suggesting that Multimode composites undergo a combination of Slow and Fast dynamics.

In the following sections, we use the distinct $D(q, \Delta t)$ characteristics described above to correlate the Slow, Fast, and Multimode classes of dynamics with composite formulation and activity time. Namely, we define the Slow class as having $D(q, \Delta t)$ curves that exhibit single, steady large-$\Delta t$ plateaus, while Fast curves display single large-$\Delta t$ plateaus but with pronounced oscillations, and the Multimode class exhibits two distinct, steady $D(q, \Delta t)$ plateaus (Fig. 2B).

**Motor competition delays the onset of acceleration and suppresses multimode dynamics**

Having identified quantitative metrics to classify network dynamics, we now determine how the dynamics vary with composite formulation and activity time $T$. We first evaluate the average actin and MT speeds $\langle \mathbf{v} \rangle$ determined from the corresponding $\tau(q)$ for each time series (7–15 per formulation) for each of the six composite formulations. Figure 3A–C shows $T$-dependent effects of crosslinking (different panels) and motors (dark versus light shades), with speeds spanning over three orders of magnitude during motor activity. Notably, as suggested by the 2D image structure functions shown in Fig. 2A, actin and MT speeds are well correlated (comparing open and closed symbols) across all composites and activity times despite the varying degrees to which they co-localize or de-mix (Fig. 1B).

We find that actin and MTs in all composites accelerate and reach a maximum speed $v_{\text{max}}$ at activity time $T(v_{\text{max}})$ (Fig. 3D), after which $\langle \mathbf{v} \rangle$ decreases. By classifying each data point in Fig. 3A as Slow, Fast, or Multimode (Fig. 3E), as described above, we measure the average Slow speed to be $\langle \mathbf{v}_{S} \rangle \approx 0.15 \mu m/s$, which is an order of magnitude slower than the Fast speed of $\langle \mathbf{v}_{F} \rangle \approx 1.8 \mu m/s$. The average low and high speeds for Multimode data are comparable to those of Slow and Fast values, respectively, with $\langle \mathbf{v}_{M} \rangle \approx 0.17 \mu m/s$ and $\langle \mathbf{v}_{M} \rangle \approx 1.7 \mu m/s$.

We next turn to evaluating how composite formulation programs the different dynamical classes and their dependence on activity time $T$. The average filament speed for the un-crosslinked kinesin-driven...
composite (no myosin) increases ~20-fold in the first $T \approx 20$ min, transitioning from Slow to Multimode to Fast dynamics (Fig. 3A), reaching $v_{\text{max}} \approx 7 \mu$m/s. Following this initial acceleratory period, the composite slowly decelerates over the course of ~40 min but never returns to dynamics classified as Slow. Introducing myosin substantially delays the onset of acceleratory dynamics, increasing $T(v_{\text{max}}) \sim 3$-fold, but has little impact on the magnitude of $v_{\text{max}}$ (Fig. 3A and D). Moreover, Slow dynamics dominate over more of the activity time than for the kinesin-only composite, as seen by the higher proportion of light blue versus dark blue squares in Fig. 3A.

These results indicate that Fast dynamics are due primarily to kinesin-driven motion, as there is minimal change in $v_{\text{max}}$ upon the addition of myosin and that myosin activity counteracts kinesin activity to delay the onset of Fast dynamics rather than cooperating synergistically to amplify active dynamics. We can understand this competition as follows.

Keeping in mind that the actin and MTs in the composites form co-entangled interpenetrating networks of comparable mesh sizes, we can assume that every actin filament is sterically interacting with several MTs and other actin filaments and vice versa. Kinesin acts to drive MTs together, which, in turn, attempt to pull co-entangled actin filaments with them, competing with entanglements from other actin filaments that resist kinesin-driven straining. However, because actin filaments are more flexible and relax faster than MTs, they are able to be swept up with the kinesin-driven MT network and then diffuse out of MT-rich clusters to maximize their entropy.

Incorporating myosin into the composites strongly enhances the competition between kinesin-driven pulling of actin and steric entanglements by pulling actin filaments together, which, in turn, attempt to pull interpenetrating MTs with them, counteracting the force of kinesin driving MTs together. The surprising antagonistic interaction between the two motors may also be due to the contractile versus extensile nature of actomyosin and kinesin–MT activity, respectively (62). Namely, kinesin motors are highly processive such that they principally induce nematic bundling, sliding, and extensile motion of rigid MTs, whereas low-duty ratio myosin motors primarily bend, compress, and contract semiflexible actin filaments into asters or foci (49, 57).

We expect this competition to manifest structurally as enhanced actin–MT mixing and interpenetration, as we see in Fig. 1B. In other words, while both filament types are pulled toward like filaments (actin to actin, MTs to MTs) by their respective motors, entanglements with the other filament type resist this motor-driven self-association, thereby facilitating mixing. The net result is reduced clustering and increased actin–MT interpenetration in double-motor composites. While the dynamics eventually mirror those of kinesin-only composites, the structure remains more homogeneous, as shown in Fig. 1B.

The fact that motor antagonism leads to a time delay rather than suppression of active dynamics suggests that eventually kinesin straining beats out myosin straining such that the dynamics mirror kinesin-only composites after being gated by myosin activity. Kinesin–MT straining likely “beats out” actomyosin activity due to the higher density of kinesin clusters compared to myosin minifilaments. As we describe in Methods, in all double-motor composites, there are ~75 force-generating kinesin clusters for every myosin II minifilament, and the average spacing between kinesin clusters connecting a pair of MTs is ~12 nm compared to ~2.6 µm (>200x longer) for myosin minifilaments along actin filaments. This increased density of strain-generating linkers along MTs, as well as their higher duty ratio and processivity, likely causes kinesin–MT force generation to dominate over that of actomyosin.

We now turn to the effect of passive crosslinking on single-motor and double-motor composites. As shown in Fig. 3A, the signatures of motor competition and activity gating seen for unlinked networks are preserved upon crosslinking of both actin (Fig. 3B) and MTs (Fig. 3C). The primary effect of crosslinking is a decrease in the maximum speed $v_{\text{max}}$ and the time over which the composites accelerate to this maximum $T(v_{\text{max}})$ (Fig. 3D). Further, both crosslinking types exhibit Multimode dynamics at the onset of activity (red and green triangles and stars), effectively
eliminating the initial Slow dynamics seen in unlinked composites, likely due to crosslinking reducing the degrees of freedom and increasing the connectivity of the composites, thereby suppressing spatially uncorrelated microscale fluctuations. In other words, large-scale restructuring (attributed to Multimode dynamics) and acceleration to \( v_{\text{max}} \) are facilitated by crosslinking in kinesin-only composites. Conversely, crosslinking of double-motor composites eliminates the initial Multimode dynamics seen for their unlinked counterparts, instead switching directly from Slow to Fast dynamics with minimal structural reconfiguration. This reduced restructuring of crosslinked double-motor composites compared to kinesin-only composites can be seen in Fig. 1.

To further understand the nature of Slow, Fast, and Multimode dynamics and why crosslinking alters the propensity to exhibit each type, we return to our Fig. 1 results, which show that crosslinking leads to larger and denser filament aggregates compared to unlinked cases. The reduced degrees of freedom and enhanced connectivity that crosslinking provides may explain this enhanced mesoscale clustering, which, in turn, suppresses microscale fluctuations available to the more randomly distributed and less bundled filaments that emerge in the unlinked cases. This mechanistic picture suggests that Fast dynamics are dominated by coordinated motion or flow of the composites while uncorrelated microscale fluctuations describe Slow dynamics. Conversely, as we describe above, we expect Multimode dynamics to arise from mesoscale restructuring, bundling, and de-mixing events.

### Fast, slow, and multimode classes correlate with distinct velocity fields and distributions

To corroborate the mechanisms that we postulate underlie the different dynamical classes in the preceding section, we evaluate the directionality and spatiotemporal coordination of the local dynamics that correspond to the sample Fast, Slow, and Multimode data that we analyze in Fig. 2.

We first create temporal color maps, which colorize each frame by the time it occurs during the video t, and overlay all colorized frames (Fig. 4A; Fig. S3). In this way, the maps depict the motion of the composites over the course of each video. Figure 4A shows the color maps for the actin channel, which are nearly indistinguishable from the MT channel of the same video (see Fig. S3), in line with our DDM results that show that actin and MTs within any given composite exhibit similar dynamics (Figs. 2 and 3). The Slow map (top panel) shows small-scale, randomly-oriented motion, while the Fast map shows spatially coordinated and nearly unidirectional motion. The Multimode map displays features of both Fast and Slow dynamics, exhibiting directionality on small scales but largely uncorrelated motion at larger scales.

To quantify the features described above, we perform PIV on the actin and MT channels of the videos analyzed in Fig. 4A. PIV vector fields in Fig. 4B and Fig. S4 show overlaid velocity fields at four equally spaced times t over the course of the videos analyzed in Fig. 4A. Arrow lengths and directions represent the average velocity of features over 20 frames (∼7.5 s) in the surrounding 8 x 8 square-pixel (6.7 \( \mu \text{m} \times 6.7 \ \mu \text{m} \)) region of the field of view.

As shown in Fig. 4B and Fig. S4, Slow fields exhibit motion that is slow (small arrows) and randomly oriented (no preferred arrow direction), while Fast fields show rapid directional motion with large arrows that all point in a similar direction. Multimode fields (Fig. 4B, bottom row) reveal features of both Slow and Fast modes, as seen by the different arrow sizes and directions. Figure 4C, which shows the histograms of speeds computed from PIV analysis of each video, corroborates the dynamics we observe in the sample flow fields (Fig. 4B) as well as in our DDM analysis (Fig. 2C). Namely, the speed distribution for the Fast class (middle row) is shifted substantially to the right of that for the Slow video (top row), and the Multimode distribution (bottom row) shows two distinct peaks that approximately align with Slow and Fast distributions, respectively. To further quantify the speed distributions and compare to our DDM results, we fit each histogram to a Schulz distribution (Fig. 2C, solid lines; Fig. S5), which we likewise used in the fitting function for the corresponding DDM image structure functions (see Supplementary Material Methods).

We find that the Slow and Fast distributions are well described by a single Schulz distribution, while Multimode distributions require a sum of two Schulz distributions. The average speed \( \langle v \rangle \) and standard deviation \( \sigma \) determined from each fit (listed in the corresponding panel) show that the speeds measured in Fourier space using DDM (Fig. 2B) and in real space using PIV are statistically indistinguishable (Table S2), with average values of \( \langle v \rangle \approx 0.3 \mu \text{m/s} \), \( \langle v \rangle \approx 1.7 \mu \text{m/s} \), \( \langle v \rangle \approx 0.2 \mu \text{m/s} \), and \( \langle v \rangle \approx 0.8 \mu \text{m/s} \) for Slow (S), Fast (F), and Multimode (M1, M2) videos, respectively.

Motivated by the apparent class-dependent anisotropy (or lack thereof) in the PIV vector fields, we also evaluate the corresponding velocity orientation distributions (Fig. 4D), which reveal isotropic motion for the Slow class, with no perceptible peak and comparable occurrences of all angles, compared to sustained unidirectional Fast motion, as evidenced by the sharply peaked narrow distribution. The Multimode distribution displays features of both Fast and Slow distributions, with a broader sampling of directions compared to Fast but with more pronounced peaks compared to Slow.

As noted in the previous section, we also see evidence of anisotropic dynamics in our DDM analysis, manifested as radial asymmetry in the \( D(q_x, q_y, \Delta t) \) plots for the Fast class and to a lesser extent in the Multimode plots (Fig. 2A, Fig. S2). To quantify this anisotropy in Fourier space, we evaluate an anisotropy factor \( A_F(q, t) \) by computing weighted azimuthal integrals of the DDM image structure function (detailed in Methods and Supplementary Material (63)). \( A_F \) can assume values between −1 and 1 for x- and y-directed motion, respectively, with \( A_F = 0 \) indicating isotropic motion. Figure 4E shows that the distributions of \( A_F \) values for Slow and Fast classes exhibit distinct peaks at \( A_F \approx 0 \) and \( A_F > 0 \), indicative of isotropic and y-oriented motion, respectively. Conversely, the Multimode distribution is broader with multiple peaks that span from \( A_F < 0 \) to \( A_F > 0 \) and include a significant fraction of near-zero values. Likewise, the Mullirome PIV orientation distribution samples a broad range of angles (isotropic, \( A_F = 0 \)) while also exhibiting distinct peaks (directionality, \( |A_F| > 0 \)).

### Spatiotemporal variations in dynamics are suppressed by motor antagonism

To better elucidate the mechanisms dictating the different dynamical classes and the influence of motor antagonism on said mechanisms, we use both DDM and PIV to resolve variations in the short-time dynamics of the composites, i.e. those that occur within the time t of a given video.

We first evaluate the average speed \( \langle v \rangle (t) \) as a function of time t for the actin and MT channels of each video analyzed in Figs. 2 and 4, which we compute from the corresponding PIV vector fields, where \( \langle v \rangle \) is averaged over all vectors in a single field. As shown in the \( \langle v \rangle (t) \) plots in Fig. 5A and Fig. S6, Slow and Fast dynamics are largely stationary over the course of a given video, with...
nearly constant speeds. In contrast, Multimode traces show discrete and abrupt shifts from intermediate to fast motion to steady slow motion.

Observing the time dependence of the corresponding average velocity orientations \( \langle \theta(t) \rangle \), we find similar trends as for \( \langle v(t) \rangle \), where the directionality of both Fast and Slow examples is nearly independent of \( t \), while the average orientation of Multimode vectors undergoes an abrupt and discrete shift at \( t = 60 \text{ s} \).

To corroborate and better characterize the apparent stationarity of Slow and Fast class dynamics and the non-stationary Multimode dynamics shown in Fig. S5A and B, we compute instantaneous DDM image structure functions \( D(q, \Delta t, t) \), which, unlike the \( D(q, \Delta t) \) curves shown in Fig. 2B, are not averaged over time \( t \) \( (63) \). By evaluating the probability distribution of \( D(q, \Delta t) \) values for all \( t \) in a given video, we can determine the extent to which dynamics are temporally heterogeneous during the acquisition time. Namely, ergodic stationary dynamics are expected to follow a Gaussian distribution of structural correlations, which are quantified by \( D_{\text{S}}(q, \Delta t, t) \). As shown in Fig. 5C, the Fast and Slow distributions are strongly overlapping, with the Slow distribution being well fit to a Gaussian function. Conversely, the Multimode distribution is distinctly non-Gaussian—with no obvious peak, a broad distribution of values, and significant noise—indicative of large intermittent fluctuations in structural correlations \( (63) \).

To quantify the extent to which the temporal \( D \) distributions deviate from Gaussianity, we compute the skewness \( S_{K} \equiv \langle (D_{\text{S}} - D) \rangle^3 / \langle (D_{\text{S}} - D)^2 \rangle^{3/2} \), which is zero for a Gaussian distribution. For reference, the distributions shown in Fig. 5C have skewness values of \( S_{K,M} \approx 0.42 \), \( S_{K,F} \approx 0.66 \), and \( S_{K,S} \approx 0.86 \) for the Slow, Fast, and Multimode classes, respectively. Positive skewness, largest for Multimode distributions, has been reported for colloidal gels that are en route toward arrest and has been interpreted as arising from discrete restructuring processes such as coalescing or rupturing, as well as intermittent fluctuations and rearrangements \( (63) \).

To determine the prevalence of non-stationary dynamics across the formulation phase space and activity times, we compute skewness values for all composite formulations and times \( T \) evaluated in Fig. 3. Figure 5D-I shows stacked confidence ellipse plots comparing skewness \( S_{K} \), average speeds \( \langle v \rangle \), and anisotropy factors \( A_{F} \) colorized by dynamical class and separated into panels for kinesin-driven composites (Fig. 5D, F, and H) and with (Fig. 5E, G, and I) myosin (also see Fig. 57). The individual points correspond to all data points shown in Fig. 3, and the ellipses enclose one standard deviation around the mean. As shown, the Multimode data exhibit the largest skewness values, as seen by the magenta ellipses being furthest to the right in Fig. 5F and H. Fast and Slow \( S_{K} \) values are similar to one another and deviate less from zero. The higher skewness for Multimode data is coupled with relatively fast speeds (Fig. 5F) but low anisotropy (Fig. 5H).
These couplings further support our interpretation that Multimode dynamics arise from large intermittent restructuring events, which we expect to have no preferred directionality but give rise to periods of time—e.g., during a restructuring event—that exhibit fast dynamics.

Comparison of the composites driven by kinesin only (Fig. 5D, F, and H, darker shades) versus two motors (Fig. 5E, G, and I; lighter shades) reveals that the presence of myosin nearly eliminates Multimode dynamics, as evidenced by the lack of magenta ellipses in Fig. 5E, G, and I. Further, the distributions of data points for the double-motor composites generally exhibit smaller skewness values as compared to kinesin-only composites, as seen by the ellipses shifted to the left in Fig. 5G and I compared to Fig. 5F and H. Despite these differences, we also observe that the distributions of speeds for composites with and without myosin are not significantly distinct, as we discussed in the previous section (also see Fig. 3).

Taken together, these results demonstrate that Multimode dynamics arise from discrete and abrupt restructuring events and coarsening, and the presence of myosin suppresses this restructuring such that double-motor composites exhibit very few instances of Multimode dynamics and remain more homogenously mixed at the end of the activity. In the absence of mesoscale discrete restructuring, the double-motor networks take longer to coarsen and switch to Fast coordinated flow.

**Motor competition inhibits composite restructuring and de-mixing enhanced by crosslinking**

To connect the dynamics we measure with various structures and reconformation, we develop a minimal model that aims to capture the key dynamical features of our composites. As described in Methods and Supplementary Material, our model simulates filament motion that arises from motor-driven advection and thermal diffusion and works against steric hindrances and viscous traps due to motor and protein crosslinking (see Fig. S8 and Table S1). We purposefully simplify the model, ignoring details such as filament flexibility and individual motor dynamics that other models incorporate (64–66), to facilitate applications to other systems and identify the key parameters that dictate the experimental phenomena we observe.

Our model simulations show that all composites start as homogeneous interpenetrating networks of actin and MTs at $T = 0$ (Fig. S9), as we see in experiments (Fig. 1B), but subsequently restructure to varying degrees depending on the composite formulation. Figure 6A, which shows sample simulation snapshots of the final states ($T = T_f$) of the six composite formulations, reveals strong suppression of restructuring by motor competition, similar to our experimental observations, with the K + M composites undergoing substantially less restructuring and de-mixing than the kinesin-only composites. Also, in line with experiments, cross linking of either actin or MTs in simulated composites enhances aggregation and clustering compared to composites without cross linkers. This agreement between model predictions and experimental observations suggests that it is the balance between frictional jamming and motor-driven de-mixing that dictates the different formulation-dependent structural regimes.

To quantify the degree of restructuring in simulations, we compute the probability distributions of like filaments ($g_{A,(MT−A)/MT}(r)$) and unlike filaments ($g_{A,(MT−MT)/A}(r)$), a radial distance $r$ from a given actin/microtubule (A/MT) for the initial ($T = 0$) and final ($T = T_f$) states of all composites (see Supplementary Material Methods). For homogeneous well-mixed networks, all distributions should equate to 1 for all $r$ values, which we find to be the case for the initial states of all simulated composites (Fig. S9). The more $g_{A,M}(r)$ or $g_{MT−MT}(r)$ values are above 1, the more clustering of actin or MTs, respectively. Conversely, $g_{A,M}(r) < 1$ or $g_{MT−MT}(r) < 1$ indicates...
Fig. 6. Minimal advection–diffusion modeling characterizes and corroborates expected motor-driven restructuring. (A) Simulation snapshots, each 155 µm x 155 µm, show sample final configurations of MTs (red) and actin (green) in composites with no passive crosslinking (No XL), actin crosslinking (Actin XL), and MT crosslinking (MT XL), subject to active forcing by kinesin (K) or both kinesin and myosin (K + M). Simulation details are provided in Supplementary Material Methods and Table S1. (B–I) The difference between the initial (μ = 0) and final (μ = T) values of various structural correlation functions, Δgμ(r) = gμ(r, T) − gμ(r, 0), versus radial distance ρ between two filaments (B–E) or pixels (F–I) for composites depicted in A. The type of passive crosslinking is color coded according to the border colors in A. (B, C) The difference in like-filament distributions for actin (ΔgA−A), open symbols) and MTs (ΔgMT−MT, filled symbols) in composites with (B) kinesin or (C) both kinesin and myosin. (D, E) Unlike-filament distribution differences for (D) actin (ΔgA−MT, open symbols) and (E) MTs (ΔgMT−A, filled symbols) for the composites analyzed in B and C. (F–I) SIA differences Δgμ for actin (open symbols) and MTs (filled symbols) computed from (F, G) experimental time-series and (H, I) simulation snapshots for composites with (F, H) kinesin or (G, I) both kinesin and myosin. Error bars for simulations are standard errors across three replicates and for experiments are standard errors across 100 images from three time-series.

To understand the underlying mechanisms driving this restructuring more fully, we consider that while kinesin motor activity adds to the advective term for MTs in the model, the processive nature of kinesin also increases the drag on the MTs. Conversely, the addition of non-processive myosin motors increases filament advection with a relatively smaller increase in drag. Thus, kinesin activity acts to collect MTs into locally segregated and de-mixing of actin from MTs or vice versa. Figure 6B–E plots the differences between the final and initial distributions, e.g. ΔgA−A = gA−A(r, T) − gA−A(r, 0), such that values of zero indicate minimal restructuring, while positive “like” distributions and negative “unlike” distributions indicate like-filament clustering and de-mixing of unlike filaments, respectively. As shown, the composites with kinesin and myosin show minimal de-mixing regardless of crosslinking (Δg ≈ 0 in Fig. 6C and E), while all composites without myosin show signatures of clustering and de-mixing, which is generally more pronounced in the crosslinked composites (Fig. 6B and D).

Finally, to directly compare the predicted and experimental restructurung, we perform identical spatial image autocorrelation (SIA) analysis (see Methods) on the initial and final experimental videos and simulation snapshots. SIA computes the correlation in intensities g(r) between two pixels separated by a radial distance r in a given image such that g(r) indicates the lengthscales over which structural features in an image are correlated. Specifically, g(r) values range from 1 for complete correlation (such as when r = 0) to 0 for complete decorrelation, e.g. for r values much larger than the size of structural features. Similar to Fig. 6B–E, we evaluate the differences between the final and initial structural functions Δgμ(r) for actin and MTs in all simulated composites (Fig. 6H and I), which we compare to experimental values (Fig. 6F and G). We find that in both experiments and simulations, the presence of myosin reduces the distance over which structural correlations are enhanced over the time course of motor activity, evidenced as faster decay in Δgμ(r) with increasing r in Fig. 6G and I compared to Fig. 6F and H. This feature is indicative of reduced large-scale clustering and de-mixing of actin and MTs, as is also evident in Figs. 1B and 6A. Moreover, in both experiments and simulations, passive crosslinking generally leads to increased structural correlations (larger Δgμ values) compared to composites without crosslinkers, in particular at larger distances and for actin crosslinking. The increased aggregation with actin crosslinking manifests in experiments as minimal decay and non-monotonic dependence of Δgμ(r) with increasing r for actin and MTs, respectively, indicative of fewer smaller-scale clusters and increased mesoscale (>10 µm) structural correlations. In simulations, increased aggregation can be seen as larger Δgμ values in the presence of actin crosslinkers across all lengthscales.

We note that given the simplicity of our model and the simulated renderings of the composites, as well as the noise in our microscope images, as can be seen in Fig. 1 and Movies S1–S3, we do not expect quantitative agreement between experiments and simulations. Rather, we aim to capture qualitatively similar dependence of structural features on crosslinking and motor competition, as we describe above. Namely, the presence of myosin inhibits restructuring while passive crosslinking enhances it. The generally larger Δgμ values measured in experiments compared to simulations are likely due to the noise and finite depth of the experimental images, which limit the occurrence of “empty space” that are seen in simulated composites, thereby overestimating correlations across lengthscales as compared to simulated images. Moreover, the flexibility of the actin filaments, not accounted for in the model, may also allow for greater restructuring and clustering.
arrested clusters that can either sweep up or squeeze out actin filaments. The addition of passive crosslinking of actin or MTs accelerates this process by facilitating the coalescence of smaller clusters into larger ones. On the other hand, myosin activity allows for filament redistribution within clusters and diffusive migration of filaments out of clustered regions, thereby inhibiting segregation between actin and MTs and increasing the rate at which newly formed clusters can dissolve back into a mixed state. Succinctly stated, motor antagonism can arise from an interplay between competitive motor-driven advection and frictional drag, irrespective of its origin—steric interactions or passive or active crosslinking.

**Conclusion**

The cytoskeleton is a non-equilibrium multifunctional composite comprising diverse protein filaments, motors, and crosslinkers that cooperate and compete to enable diverse cellular structures and processes. As such, the cytoskeleton is one of the primary inspirations to the burgeoning field of active matter, and much of current active matter research seeks to learn from and emulate the cytoskeleton. The composite nature of the cytoskeleton, which confers its signature versatility and programmability, is one of its hallmarks. Yet, current active matter platforms are largely limited to a single force-generating component and/or substrate. We address this gap by engineering co-entangled and crosslinked composites of MTs and actin filaments driven by kinesin and myosin motors—breaking new ground in active matter design by incorporating multiple independently tunable force-generating components and active substrates.

By coupling Fourier-space and real-space analyses (DDM and PIV), we show that composites undergo a combination of Fast adective flow, Slow isotropic fluctuations, and Multimode restructuring that result in structures ranging from interpenetrating actin–MT scaffolds to de-mixed amorphous clusters. Surprisingly, competition between kinesin and myosin driving delays the onset of kinesin-driven acceleration without appreciably changing the range of speeds that the different composites exhibit. Motor antagonism also suppresses mesoscale restructuring events that underlie Multimode dynamics, thereby sustaining mixed networks of actin and MTs. Conversely, passive crosslinking hastens the onset of kinesin-mediated acceleration and subsequent deceleration by enhancing network connectivity and suppressing uncorrelated microscale motion. Importantly, the emergent dynamics and extensive programmable phase space of non-equilibrium properties we reveal are a result of very subtle changes in substrate connectivity and activity.

Our work brings reconstituted cytoskeleton systems an important step closer to mimicking the complexity of the active composite cytoskeleton by integrating two distinct and ubiquitous motor–filament systems, actomyosin and kinesin–MT networks, that have been shown to interact and co-mediate important cellular processes including morphogenesis and exocytosis (filament systems, actomyosin and kinesin–MT networks, that cooperate and compete to enable diverse cellular structures and processes. As such, the cytoskeleton is one of the primary inspirations to the burgeoning field of active matter, and much of current active matter research seeks to learn from and emulate the cytoskeleton. The composite nature of the cytoskeleton, which confers its signature versatility and programmability, is one of its hallmarks. Yet, current active matter platforms are largely limited to a single force-generating component and/or substrate. We address this gap by engineering co-entangled and crosslinked composites of MTs and actin filaments driven by kinesin and myosin motors—breaking new ground in active matter design by incorporating multiple independently tunable force-generating components and active substrates.

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Our work brings reconstituted cytoskeleton systems an important step closer to mimicking the complexity of the active composite cytoskeleton by integrating two distinct and ubiquitous motor–filament systems, actomyosin and kinesin–MT networks, that have been shown to interact and co-mediate important cellular processes including morphogenesis and exocytosis (67, 68), mechanosensation (69), and migration and stiffening (70). Interactions between actomyosin, kinesin, and MTs have also been implicated in wound healing, mitosis, and cytoplasmic streaming (7, 15, 16, 28, 43, 71, 72). Because the motor and filament concentrations in our composites are within physiological ranges (73), our results offer new insight into the macromolecular dynamics and interactions that contribute to these cellular processes. We note that cell-like confinement of in vitro cytoskeletal networks has also been shown to play a key role in recapitulating dynamics and structures seen in cells (71, 72, 74). We plan to build in this layer of complexity in our future work (75).

Finally, the programmability of our composites, with multiple well-controlled tuning knobs—motors, filaments, and crosslinkers—which can each be varied independently while maintaining composite integrity, opens the door for reconfigurable materials that can be programmed to exhibit varying types and rates of motion and restructuring over broad spatiotemporal scales. For example, materials based on our designs could be used as spatially controlled microactuators, responsive filtration and sequestration devices, and self-curing and self-repairing infrastructure technologies. Our minimal advection–diffusion model that recapitulates our experimental trends is broadly applicable to active composite networks and lays the foundation for more complex predictive models that quantitatively capture the structure and dynamics of composite active matter. As such, we anticipate that our double-motor material design, intriguing dynamical results, and corresponding modeling framework will spark a new class of studies that explore the broad parameter space of this platform.

**Methods**

See Supplementary Material Methods for more detailed descriptions of each of the following sections.

**Protein preparation**

Rabbit skeletal actin monomers (Cytoskeleton), biotinylated actin monomers (Cytoskeleton), porcine brain tubulin dimers (Cytoskeleton), biotinylated tubulin dimers (Cytoskeleton), rhodamine-labeled tubulin dimers (Cytoskeleton), and myosin II (Cytoskeleton) are reconstituted and flash-frozen into single-use aliquots according to previously described protocols (28, 48). Biotinylated kinesin-401 is expressed in Rosetta (DE3)pLysS competent E. coli (Thermo Fisher) and purified, as described in Supplementary Material.

For composites that incorporate actin or MT crosslinking, actin–actin or MT–MT crosslinker complexes are prepared according to previously described protocols (48). In brief, biotinylated actin or biotinylated tubulin is combined with NeutrAvidin and free biotin at a ratio of 2:2:1 protein:free biotin:NeutrAvidin.

Immediately prior to experiments, myosin II is purified as previously described (27) and stored at 4°C. Kinesin clusters are formed by incubating dimers at a 2:1 ratio with NeutrAvidin with 4 µM DTT for 30 min at 4°C.

**Active cytoskeleton composite preparation**

Actin-MT composites are formed by polymerizing 2.32 µM unlabeled actin monomers and 3.48 µM tubulin dimers (% rhodamine-labeled) in PEM-100 (100 mM PIPES, 2 mM MgCl₂, and 2 mM EGTA) supplemented with 0.1% Tween, 10 mM ATP, 4 mM GTP, 5 µM Taxol, and 0.47 µM AlexaFluor488–phalloidin (Life Technologies).

For crosslinked composites, a portion of either actin monomers or tubulin dimers is replaced with equivalent crosslinker complexes to achieve the same overall actin and tubulin concentrations and crosslinker:protein ratios of Rₐ = 0.02 for actin or R_MT = 0.005 for MTs. Rₐ and R_MT values are chosen to achieve similar lengths between crosslinkers d along actin filaments and MTs (dₐ ≃ 60 nm, d_MT ≃ 67 nm) (48) and to be high enough to induce measurable changes in the viscoelasticity compared to unlinked networks but low enough to prevent filament bundling.
Composites are polymerized for 30 min at 37°C, after which 1.86 μM phalloidin is added and the composite is incubated for 10 min at room temperature. Following, 50 μM blebbistatin (26), an oxygen scavenging system, 0.47 μM myosin II, and 0.35 μM kinesin are added. Concentrations of actin, tubulin, myosin II, and kinesin are within reported physiological ranges (73, 76, 77).

While myosin activity is controlled by blebbistatin deactivation, kinesin starts to act immediately, so T = 0 for each experiment is set as the time kinesin is added. Each sample is gently flowed into a ~1 mm x 24 mm sample chamber composed of a silanized (78) coverslip and microscope slide fused together by a ~100-μm-thick paraffin spacer and sealed with epoxy, creating an airtight chamber.

**Fluorescence microscopy**

Imaging of AlexaFluor488-labeled actin and rhodamine-labeled MTs comprising composites is performed using a Nikon A1R laser scanning confocal microscope with a 60x 1.4 NA oil immersion objective (Nikon), 488-nm laser with 488/525-nm excitation/emission filters, and 561-nm laser with 565/591-nm excitation/emission filters. 488-nm illumination also locally deactivates blebbistatin (26–28). Time-series (videos) of 256 × 256 square-pixel (213 μm × 213 μm) images are collected at 2.65 fps for a maximum video time of t_{max} = 1000 frames (~377 s ± 6.28 min). Imaging begins 5 min after the addition of kinesin motors (T = 5 min) in the middle of the ~100-μm-thick sample chamber. Each successive video is collected in a different field of view of the same sample until there is no longer any discernible restructuring or motion (T = 60 – 120 min). A total of 7–15 videos are collected for each of the six formulations (no crosslinking, actin crosslinking, and MT crosslinking, with and without myosin). Each video includes two channels that separate the actin and MT signals such that they can be processed separately and compared.

**DDM**

DDM is performed on the actin and MT channels of each video, as described previously (28). Image structure functions are determined by taking the square of 2D Fourier transforms of differences between an image at time t and one at t + Δt. This process yields the instantaneous image structure function D_{ij}(q_x, q_y, Δt, t), where q_x and q_y are x and y components of the wave vector q. As typically done in DDM analysis, we average D_{ij} over all times t (frames) of a given video, and all wave vectors q with the same magnitude q, to determine the 1D image structure function D(q, Δt) that can be fit to various models. We fit D(q, Δt) versus Δt for each wave vector q to a model in which the distributions of speeds are described by one or two Schulz functions (60) (see Supplementary Material Methods), as has been done to describe other ballistic biological systems (60, 61). For Slow and Fast data in which one distribution is sufficient, there are four free parameters (A, β, r_1, Z_1), whereas for Multimode data, there are seven (adding r_2, Z_2, and f) (see Supplementary Material Methods). For each video, we perform fits for 40 different q values in the range q = 0.8–2 μm^{-1} (~3–8 μm), from which we extract r(q) curves for the actin and MT channels. By fitting each r(q) curve to r(q) = (|q|q)^{-1}, we compute the average speed ⟨v⟩ for each channel of each video. We determine the error associated with ⟨v⟩ using two methods. First, we compute v from each individual (q, t) pair (i.e. v = 1/r(q)) and determine the standard error across those values. Second, we use the Schulz parameter Z determined from our D(q, Δt) fits and our measured ⟨v⟩ value to compute the standard deviation σ and corresponding standard error via the relation Z = (2σ)^2 − 1. Error bars shown in Fig. 3 are the larger of the two values for each case.

To determine the degree to which dynamics deviate from radial symmetry, implying directionality, we compute the anisotropy factor A_r of D_{ij}(q_x, q_y, Δt, t) in q-space by computing A_r(q, Δt, t) = \frac{\int D(q_x, q_y, Δt, t) \cos(2θ)dθdΩ}{\int D(q_x, q_y, Δt, t) dθdΩ} and averaging over q, Δt, and t (79, 80). θ is defined relative to the positive y-axis such that A_r > 0 and A_r < 0 correspond to motion along the y- and x-directions, respectively, and A_r = 0 indicates isotropic motion.

To evaluate the time dependence of dynamics over short timescales (within the time t of a single video), we investigate the temporal distribution of instantaneous image structure functions D_{ij}(q_x, q_y, Δt, t) for a given Δt and q. For steady-state dynamics, one would expect this distribution to be Gaussian. Deviations from Gaussianity indicate sporadic events, which cause larger than typical structural decorrelations. We quantify this non-Gaussian behavior by evaluating the skewness, S_k = ((⟨D(x, y)⟩)^3)/⟨(⟨D(x, y)⟩)^2⟩^{3/2}, where the average is over Δt and q.

**PIV**

PIV is performed using the GPU-accelerated version of OpenPIV (81). Interrogation windows of 8 × 8 square pixels, with a 4 × 4 square-pixel overlap, are used to generate 64 × 64 grids of velocities for MT and actin channels of each time series. Average velocities v̄ for each interrogation window are determined from image pairs separated by Δt = 10 frames (~3.77 s). From the measured velocities, we determine the distribution of individual speeds v(t) and velocity orientations θ(t) across each image over the course of a video. To identify and exclude spurious velocities during statistical analysis, we reject those points for which the signal-to-noise ratio was <2. We fit the speed distributions to Schulz distributions by minimizing the mean square difference between the predicted statistical assign distribution to each bin of (width 50 nm/s) for a given choice of parameters and the actual fraction of speeds in each bin. Arrows plotted in Fig. 4B and Fig. S4 represent the local velocity on a regular Cartesian grid, with arrow length proportional to speed. Visualizations at different video times t are superposed, with arrow color representing t.

**SIA**

SIA is performed separately on actin and MT channels of microscope images and simulation snapshots (see below) using custom Python scripts (27, 28, 81). SIA measures the correlation in intensity g_2 of two pixels in an image as a function of separation distance r (82). We generate autocorrelation curves g_2(t) by taking the fast Fourier transform of the image I(r), multiplying by its complex conjugate, applying an inverse Fourier transform F^{-1}, and normalizing by the squared intensity: g_2(t) = \frac{F^{-1} |F(I(r))|^2}{\int |I(r)|^2 dr}. Correlation curves shown in (i) Fig. 6F and G and (ii) Fig. 6H and I are averages across (i) 100 microscope images from three time series and (ii) simulation snapshots from three independent runs (see below). Error bars indicate the standard error.

**Computational model**

To predict motor-driven restructuring, we develop a minimal model that captures the key energetic components of our system, as fully described in Supplementary Material Methods and Table S1. In brief, we allow filaments to interact with neighboring filaments via (i) motor-generated forces that can either pull the interacting filaments toward each other or push them away and (ii) crosslinks that increase the frictional forces on the interacting...
filaments. The movement of a filament center to a neighboring grid point within a small temporal time step is then a stochastic event with a probability that can be calculated by the standard solution to the Fokker–Planck equation given by

\[ p_i(x) \geq 1 - \frac{1}{2} (1 + \text{erf} \left( \frac{x - \mu_i}{\sigma_i} \right)) \]

where \( i \) is the distance to the next grid point in a particular direction, \( \mu_i \) is the average advection-induced displacement in that direction, and \( \sigma_i \) is the diffusion-based rms 1D displacement of the filament along the direction to the specific grid point. The subscript \( i \) represents a specific filament in the model. The movement probability of filament \( i \) to a neighboring grid point that either (i) contains the center of filament \( j \) or (ii) is empty is given by (i) \( p_{ij} = p_i \times p_j \) or (ii) \( p_{ij} = 1 \).

We implement our model on a 150 µm \( \times \) 150 µm hexagonal lattice with 2.5-µm spacing and use numerical values for all model parameters that are based on experimental and literature values (see Table S1). Initially, each lattice point is either empty or occupied by a center of an actin filament or MT using probabilities matching the average volume fraction occupied by these elements. The movement of the filaments is simulated in each iteration by calculating the likelihood of each possible movement \( p_{ij} \) for all grid points \( i \) containing filament centers and randomly picking one of these movements to occur based on these probabilities (Fig. S8) (83). We perform three independent simulation runs for each composite formulation (Fig. S9).

To quantify the degree of clustering and segregation of the different filaments, we compute the probability distributions of filaments that are alike, \( g_{AT} = \frac{N_{AT}}{N} \) or \( g_{MT-MT} = \frac{N_{MT-MT}}{N} \), and unlike, \( g_{AM} = \frac{N_{AM}}{N} \) or \( g_{MT-AT} = \frac{N_{MT-AT}}{N} \), a radial distance \( r \) from a given actin filament (A) or MT (see Supplementary Material Methods). In the above, \( N_{AT} \) is the number of actin/MT neighbors at distance \( r \) from a specific filament, \( f_{AM} \) is the actin/MT volume fraction, and \( N(\cdot) \) is the maximum number of possible neighbors that reside a distance \( r \) from the specific actin/MT. An increase in \( g_{AM-AM} \) above 1 indicates clustering of actin/MTs, while a decrease in \( g_{MT-MT/AT} \) below 1 indicates segregation of unlike filaments. Correlation analysis data shown in Fig. 6B–E are averages over all filaments of the same type over three statistically independent replicates with error bars representing the standard error.

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Supplementary material

Supplementary material is available at PNAS Nexus online.

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Author contributions

R.M.R.-A. conceived the project, guided the experiments, interpreted the data, and wrote the manuscript. R.J.M. guided the experiments, analyzed and interpreted the data, and wrote the paper. K.A.L. purified and assayed the kinesins. C.J.C. performed the experiments and analyzed the data. J.M. and M.S. analyzed the data and prepared the figures. J.L.R. helped conceive the project, guide the experiments, interpret the data, and write the manuscript. P.K. and C.G. designed and performed the modeling and simulations. M.D. helped interpret the data, develop the theoretical models, and write the manuscript. M.J.R. helped guide the experiments and provided useful feedback.

Data availability

All data generated or analyzed during this study are included in this published article and its supplementary information files or available at Zenodo: doi: 10.5281/zenodo.8165779.

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