Spontaneous Formation of a Globally Connected Contractile Network in a Microtubule-Motor System

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ABSTRACT Microtubule (MT) networks play key roles in cell division, intracellular transport, and cell motility. These functions of MT networks occur through interactions between MTs and various associated proteins, notably motor proteins that bundle and slide MTs. Our objective in this study was to address the question of how motors determine the nature of MT networks. We conducted in vitro assays using homotetrameric kinesin Eg5, a motor protein involved in the formation and maintenance of the mitotic spindle. The mixing of Eg5 and MTs produced a range of spatiotemporal dynamics depending on the motor/filament ratio. Low motor/filament ratios produced globally connected static MT networks with sparsely distributed contractile active nodes (motor-accumulating points with radially extending MTs). Increasing the motor/filament ratio facilitated the linking of contractile active nodes and led to a global contraction of the network. When the motor/filament ratio was further increased, densely distributed active nodes formed local clusters and segmented the network into pieces with their strong contractile forces. Altering the properties of the motor through the use of chimeric Eg5, which has kinesin-1 heads, resulted in the generation of many isolated asters. These results suggest that the spatial distribution of contractile active nodes determines the dynamics of MT-motor networks. We then developed a coarse-grained model of MT-motor networks and identified two essential features for reproducing the experimentally observed patterns: an accumulation of motors that form the active nodes necessary to generate contractile forces, and a nonlinear dependency of contractile force on motor densities. Our model also enabled us to characterize the mechanical properties of the contractile network. Our study provides insight into how local motor-MT interactions generate the spatiotemporal dynamics of macroscopic network structures.

INTRODUCTION

Living cells continuously change their state to adapt to external environments (1). This dynamic behavior relies on the continual reorganization of the cytoskeleton (2,3). Microtubules (MTs) are a major component of the cytoskeleton, interacting with various associated proteins to organize hierarchical structures spanning tens of micrometers to millimeters. The structures vary dynamically during the cell cycle according to the physiological roles of the cell (4–6), including mitotic spindle assembly (7), intracellular transport (8), neural pathfinding (9,10), plant cell morphogenesis (11–13), and regulation of protein synthesis (14).

In most of these phenomena, the centrosome plays central roles in nucleating and organizing MT structures (15–19). In addition to these centrosome-associated MTs, acentrosomal MTs form bundles and network structures (20) and contribute to various cellular processes, including cell migration (21), neuron morphology (22), and the formation and maintenance of plant cell shapes (23,24). In both cases, a number of associated proteins are known to be involved in structure formation, including PRC1 in mitotic spindle assembly (25–27), and MAP2 and tau in neural morphogenesis (28).

Although many kinds of elements are required for the formation of MT-related structures in cellular contexts, previous in vitro studies have found that characteristic structures such as asters, vortices, and networks thereof can emerge from a mixture of only a few components (29–33). These experimental findings have sparked a wealth of theoretical research (34,35). An important finding from these in vitro studies is that the bundling and sliding of cytoskeletal filaments play a crucial role in pattern formation. However, it is not yet understood how the bundling and sliding of MTs generate various pattern dynamics.

To address this question, we used Eg5 as a motor protein with the ability to both bundle and slide MTs. Eg5 is an evolutionarily conserved kinesin that is known to contribute to the formation and maintenance of the mitotic spindle and...
the morphogenesis of axons and dendrites (36–40). A striking feature of Eg5 is that it has two motor units (each of which has two motor heads) linked through a coiled-coil region, so that each Eg5 molecule has the defined and minimum number of motor units to bundle and slide MTs. Using this eminently suitable motor protein, we investigated the temporal evolution of the motor-MT mixture under a wide range of motor/filament ratios.

Although a previous study using native Eg5 reported that it did not form asters (32), we found that Eg5, under some conditions, can form asters and aster networks. The MTs associated with the asters have defined polarity, generating a flux of MTs toward the center of the aster that results in local contractions between two connected asters. We observed that these local contractile dynamics generated diverse spatiotemporal dynamics in macroscopic MT networks depending on the spatial distribution of the asters. These dynamics are characterized by the connectivity and contractility of the MT network, which is categorized as a static network, active network, or aggregation. The contractile dynamics of global MT networks was also observed in a recent study (published after the original submission of this work) using Xenopus egg extract (41,42). We also found that the modulation of motor properties, such as velocity and affinity for MTs, induced a different kind of dynamics that led to the formation of uniformly sized, isolated asters.

Using these findings, we developed a coarse-grained model based on the local interactions of multiple asters to reproduce our experimentally observed patterns. This model revealed two features essential to the active patterning of the MT-motor system: an accumulation of motors that forms the active nodes needed to exert contractile forces, and a nonlinear dependency of contractile force on motor densities. With this model, we calculated the mechanical energy stored in MT networks, which enabled us to recharacterize our experimentally observed patterns in terms of the mechanical behaviors of MT-motor systems.

Our study raises the possibility that living cells may utilize diverse macroscopic MT patterns through the regulation of geometrical network hub configurations (43,44). These configurations may be controlled by changing the local interactions, densities, and activities of a few basic cytoskeletal elements (45) in the absence of an elaborate network-organizing apparatus.

**MATERIALS AND METHODS**

**Cloning and plasmid construction**

The human Eg5 construct (pcDNA3.4/HsEg5-EGFP-FLAG-8xHis) used in this work was a kind gift from Ken’ya Furuta (National Institute of Information and Communications Technology, Tokyo, Japan). To generate the chimeric KIF5Bhead-Eg5tail construct, the motor domain sequence of kinesin family member 5B (KIF5B, NM_004521.2; amino acid residues 5–333) was amplified from cDNA from HEK293 cell total RNA, which was isolated from 2 × 10⁶ cells using the SV Total RNA Isolation System (Z3101; Promega, Madison, WI). The amplified sequence was replaced with the Eg5 motor domain sequence (1–368) using the ligation-independent In-Fusion (Clontech, Palo Alto, CA) method.

**Protein expression and purification**

To purify the kinesin constructs, Exp293F cells (Invitrogen) were used as previously described (46). Cells were collected by centrifugation at 700 g for 10 min, washed three times with phosphate-buffered saline (pH 7.4), and stored at −80°C. The frozen cells were suspended in lysis buffer (20 mM Tris-HCl at pH 7.5, 250 mM NaCl, 1 mM MgSO₄, 100 μM ATP, 10% (w/v) sucrose, 10 mM imidazole-HCl at pH 7.5, and 0.5 mM dithiothreitol) containing 1 mM phenylmethyleneisulfonfonyl fluoride and a protease inhibitor cocktail (Nacalai Tesque, Kyoto, Japan). The resuspended cells were then disrupted by a sonicator (Sonifier 250; Branson, Danbury, CT). After ultracentrifugation at 21,000 g for 45 min at 4°C, the supernatant was loaded onto Ni-IMAC resin (Bio-Rad, Hercules, CA) and eluted with 20 mM Tris-HCl at pH 7.5, 250 mM NaCl, 1 mM MgSO₄, 100 μM ATP, 10% (w/v) sucrose, 250 mM imidazole-HCl at pH 7.5, and 0.5 mM dithiothreitol. The eluate was next loaded onto anti-FLAG agarose (Sigma, St. Louis, MO) and eluted with lysis buffer supplemented with 350 μg/mL 3×FLAG peptide (Sigma). The concentration of the products was determined by using the Bradford method with the Coomassie Plus protein assay reagent (Thermo Fisher Scientific, Waltham, MA).

Tubulin was purified from porcine brain tissue through two successive cycles of polymerization and depolymerization. A phosphocellulose column (Whatman, Maidstone, UK) or high-molarity PIPES buffer (47) was used to remove contaminating MT-associated proteins. Tubulins purified with high-molarity PIPES buffer were used to polymerize guanosine-5′-[γ-(oxo/2)-methylene]triphosphate (GMPCPP) MTs. To prepare fluorescently labeled MTs, tubulin was labeled with ATTO647N (ATTO-TEC) or ATTO565 (ATTO-TEC). ATTO647N- and ATTO565-labeled MTs were polymerized by copolymerizing and labeled and unlabeled tubulin for 30 min at 37°C, and stabilized with 40 μM paclitaxel (Sigma). The labeling ratios were 5% and 10% for ATTO647N-MT and ATTO565-MT, respectively.

**Observation of the spatiotemporal dynamics of the MT-kinesin network**

Time-lapse confocal images of fluorescently labeled MTs and green fluorescence protein (GFP)-tagged kinesins were collected using a Nikon A1 confocal laser-scanning microscope system with a Plan Apo VC 20X DIC N2 objective lens (NA = 0.75, Nikon) and diode lasers (Sapphire 488-20, Sapphire 641-20, and Cube 640-40C; Coherent, Santa Clara, CA). The flow chamber was composed of two coverslips (32 mm × 24 mm and 18 mm × 24 mm) that were coated with Teflon (K. Furuta, personal communication) and spaced with double-coated tape (80 μm thick, W-12; 3M, St. Paul, MN). To prepare the Teflon-coated coverslips, the uncoated coverslips were first cleaned with 2% Hellmanex II (Hellma, Müllheim, Germany) in Milli-Q water for 20 min with heat (2510; Branson), followed by a 1:1 mixture of ammonium hydroxide and hydrogen peroxide for 20 min with heat. Then, the surface was activated with oxygen plasma for 1 min (PDC-32G; Harrick, Pleasantville, NY), followed by fluorosilanization. Finally, the coverslips were coated with 0.5% Teflon AF 1601 (Dupont, Wilmington, DE) in FC-40 (3M). The chamber was first passivated using 1% (w/v) Pluronic F127 dissolved in BRB80 (80 mM PIPES-KOH, 1 mM MgSO₄, 1 mM EGTA) buffer and then incubated for 5 min to prevent nonspecific adsorption of proteins. After the chamber was washed with BRB80 buffer supplemented with 10 μM taxol, a mixed solution of fluorescently labeled MTs and GFP-fused kinesin constructs in assay buffer (80 mM PIPES-KOH pH 6.8, 2 mM MgSO₄, 1 mM EGTA, 2 mM dithiothreitol, 70 units/mL creatine kinase (Roche, Basel, Switzerland), 5 mM creatine phosphate, 42.5 units/mL glucose oxidase (Sigma), 42.5 units/mL catalase, 25 mM glucose, and 2 mM ATP) was allowed to flow into the chamber. Most images were
recorded at time intervals of 1 min. The observations were conducted in a region ~30–40 μm away from the glass surface.

Measurement of the velocities of nodes

Eg5 tends to first accumulate at the intersection of the MTs and then move toward the plus end, forming motor-accumulated nodes in MT networks. To measure the velocities of these nodes, the drift in the movies was first corrected with the use of an ImageJ software plugin (StackReg, http://bigwww.epfl.ch/thevenaz/stackreg/). Then, the motor channel was split and filtered using a median filter. The detection of nodes in the images was based on finding local maxima using the Find Maxima function of ImageJ. The position of nodes at each time point was determined by two-dimensional Gaussian fitting, using custom software (48).

Characterization of the spatiotemporal dynamics of the node velocities

We quantified the randomness and persistence of the node velocities by calculating the spatial correlation function of the node velocity vectors, \( \mathbf{v} = (v_x, v_y) \). To obtain the correlation function, we first calculated the pairwise node velocity correlations, \( v_x(r)(r + d) \) and \( v_y(r)(r + d) \), for all distance vectors, \( d \), between any two velocity vectors separated at the node-node distance, \( |d| \). Next, we averaged these pairwise velocity correlations over all nodes located around the distance \( |d| \pm 10 \mu m \) to obtain the correlation functions \( C_{x}(\|d\|) \) and \( C_{y}(\|d\|) \) for the \( x \) and \( y \) directions, respectively. We calculated the \( x \) and \( y \) components separately because the chambers we used in our experiments were highly elongated in the \( x \) direction compared with the \( y \) direction. We performed these calculations at each time point to obtain the temporal changes that occurred in the spatial correlation function of the node velocity.

Characterization of the spatiotemporal dynamics of the MT network

We characterized the spatiotemporal dynamics of the MT networks by measuring the time evolution of the number and size of the clusters, which we defined as connected structures in the MT images, and calculating the connectivity and contractility of the network. To determine these values, we first performed an image analysis combining binarization and morphology operations. Time-series images of MT channel were median-filtered with a radius of 20 pixels to reduce the background noise. The resultant images were binarized with a threshold value that was set at 0.85 times the mean gray-scale value at the initial time point. The binarized images were morphologically closed using a 20-pixel-radius disk as a structuring element to merge components that were smaller than the structuring element. The factor 0.85 and the closing operation were needed to ensure the mean grayscale value at the initial time point. The binarized images were then filtered with a radius of 20 pixels to reduce the background noise. The resultant images processed by the closing operation usually contained many smaller elements. The factor 0.85 and the closing operation were needed to ensure that plus-ended motors accumulate in the center of the aster, i.e., into nodes (Fig. S12 B). We attributed this accumulation to the facts that 1) a single aster has a radial array of filaments with their plus ends pointing inward, and 2) plus-end-directed motors move along the filaments toward the center of the aster according to their polarity. This motor accumulation process can be implemented by a constant flux, \( f \), of motor concentration per unit of time, during which the motors can travel through adjacent lattice nodes, \( i \) and \( j \), against the motor concentration gradient:

\[
\frac{dC_i}{dt} = f(C_j) - f(C_i) \tag{1}
\]

Here, we set the maximum motor concentration, \( C_{\text{max}} \), at all nodes to take into account the saturation of the motors at each aster. The initial condition, \( C_i \), is randomly drawn from the uniform distribution between zero and \( C_{\text{max}} \). The parameter \( C_{\text{max}} \) determines the motor concentration in the whole system, and served as the main control parameter during the simulations.

Because filaments between two adjacent asters are arranged mostly in an antiparallel orientation, a link connecting two adjacent nodes contracts through the sliding action of the motors (Fig. S12 C). When the motors are abundant in both of the adjacent nodes, the link between the two nodes contracts as the nodes pull each other. However, when the motors are scarce at either of the adjacent nodes, there is only a small degree of contraction because of the lack of strong sliding activity. These contraction processes can be described by a dipole interaction (56–59) between nodes \( i \) and \( j \) with a magnitude \( K_{ij} \) and a node-node distance \( L_{ij} \), both of which are dependent on the motor concentrations \( C_i \) and \( C_j \). Because the force dipoles acting on the \( i \)th and \( j \)th node are equal and opposite to each other, \( K_{ij} \) and \( L_{ij} \) are given by, to the lowest order, \( \propto C_i + C_j \) and \( \propto -(C_i + C_j) \), respectively. We generalize these relationships into the following equations:

\[
K_{ij} = \frac{1}{4} (K - k) \left[ 1 + \tanh \left( \frac{C_i - C_j}{\epsilon} \right) \right] \times \left[ 1 + \tanh \left( \frac{C_j - C_i}{\epsilon} \right) \right] + k \tag{2}
\]
FIGURE 1  Self-organized structures emerged from the mixture of MTs and Eg5. (A) Time-series images showing the formation of bundled MT structures under low MT and Eg5 concentrations ([tubulin] = 10 nM, [Eg5] = 2.97 nM). The images for ATTO647N-labeled MT (magenta) and GFP-fused Eg5 (cyan) are merged. The times displayed in the images represent the time that elapsed after initial mixing. The arrowhead in the bottom image (60 min) indicates a jointed point of bundled MTs. (B) General image of bundled MT structures. The yellow box denotes the MT structure shown in (A). (C) Time-series images show the time-dependent evolutionary dynamics of astral structures in the crowded MTs ([tubulin] = 1 μM, [Eg5] = 8.91 nM). The times displayed in the images represent the time that elapsed after mixing. The arrowheads in the bottom image (60 min) indicate the formation of astral MT structures, with Eg5 accumulated at their centers. (D) Enlarged images of astral MT structures shown by the arrowheads in (C). (E) Vector field representation of node velocities in the static network shown in (C). The velocity of each node is shown by the length of the arrows and the color scale. (F) Temporal evolution pattern of the mean node velocity in the static network shown in (C). The mean value and standard deviation (SD) are shown. The total trace number of nodes is 216.

\[
L_{ij} = \frac{1}{4} (l - L) \left[ 1 - \tanh \left( \frac{C_j - \tilde{C}}{\epsilon} \right) \right] \\
\times \left[ 1 - \tanh \left( \frac{C_i - \tilde{C}}{\epsilon} \right) \right] + L, 
\]

where \( K \) and \( k \) represent the maximum and minimum contractile strengths, respectively. \( K \) corresponds to the stall force of the motors, whereas \( k \) originates from the elastic restorative forces of MT filaments against their deformations. \( L \) represents the maximum compression length, which takes into account the limitation of compression due to the finite size effect of the MT filaments. \( C \) and \( \epsilon \) are phenomenological parameters that control the onset and saturation of contractility. The hyperbolic tangent profiles used in Eqs. 2 and 3 are some of the simplest forms that can be used to describe the nonlinearities in the onset and saturation of contractility, as well as to stabilize numerical calculations for high motor concentrations. The saturation effect originates from the fact that the space on which motors work becomes restrictive at high motor concentrations. In contrast, the nonlinear onset of the contractile force \( K_{ij} \) can be attributed to the existence of positive feedback between the motor binding/unbinding kinetics and filaments: a higher density of motors recruits more filaments, which promotes motor binding to filaments and prevents the motors from escaping far away from the filaments, due to confinement by the filaments.

In this study, we assumed that the motion of the nodes was overdamped due to solvent friction. Furthermore, because we focused only on motor-driven filament rearrangements, we did not account for thermal fluctuations and restricted our model to deterministic motions. With these simplifications, the equation describing the motion of the position vector \( r_i \) of the \( i \)th node can be written in the following form:

\[
\gamma \frac{dr_i}{dt} = - \sum_j K_{ij} \left( \left| r_i - r_j \right| - L_{ij} \right) \frac{r_i - r_j}{\left| r_i - r_j \right|}. 
\]

where \( \gamma \) is a friction coefficient and the summation runs over the nodes \( j \) that possess a link to node \( i \). The total elastic energy, \( E \), stored in the system is calculated by the following equation:

\[
E = \sum_{ij} \left( K_{ij} \left( \left| r_i - r_j \right| - L_{ij} \right)^2 \right).
\]
The nodes started linking together through MTs in solution referred to as active nodes, which can exert active contractile force. In the early stage of the dynamics, the formation of a globally connected MT network occurred. Due to the increased Eg5 concentration, both the number and size of active nodes in the network increased compared with those observed in the static network. After a considerable amount of time had elapsed (~20 min), the linking of contractile active nodes brought about a global contraction of the network. The velocity field of the nodes indicated an immotility of active nodes during network formation and rapid movements during the contraction step (Fig. 2 B; Movie S4). The temporal evolution of the mean node velocity and the velocity correlation function clearly showed the existence of a waiting period before contraction started (Figs. 2 C and S3 B). Since the temporal evolution of the total network size and that of the largest cluster size behaved almost identically (Fig. 2 D), one can conclude that the network contracted as a whole. This dynamic pattern with global connectivity and contractility is referred to as an active network (Table S1). In addition to the global contraction, occasional self-rupturing of the network was also observed (Fig. S3 C). Active-network dynamics was observed under several Eg5 concentrations (18, 22.5, and 29.7 nM; Fig. S4 and Table S1). Interestingly, under some Eg5 concentrations, both static and active networks were observed (Fig. S3 D), probably because of the initial distribution of node positions (Fig. S5).

When the Eg5 concentration was further increased ([Eg5] = 45 nM), MT-Eg5 clusters of various sizes formed even before the observations began (Fig. 3 A, 5 min). Densely distributed active nodes in the clusters caused an immediate contraction of the network and separated it into aggregates of different sizes and shapes (Figs. 3 A and S6 A; Movie S4). In addition to the aggregates of various sizes, several spherical asters of similar size were segregated during the contraction (Fig. S6 B). The time-evolution pattern of the node velocity field and the mean node velocity showed that contraction started at the early stage of the dynamics and was localized (Figs. 3, B and C, and S6 C). Based on the large difference between the time series for
the total network size and the largest cluster size, one can conclude that network segmentation occurred at the early stage of the dynamics (Fig. 3D). This dynamic pattern with local connectivity and strong contractility is called aggregation (Fig. S6 D and Table S1).

**Effects of MT properties on network dynamics**

We next investigated the effects of MT properties on spatiotemporal dynamics by using MTs shortened by shearing and GMPCPP-MTs instead of taxol-stabilized (normal) GTP-MTs. We confirmed that the sheared MTs and GMPCPP-MTs were shorter in length than the GTP-MTs (Fig. S7, A and B). Before observing the spatiotemporal dynamics of the MT network, we observed Eg5 moving on both GMPCPP-MTs and taxol-stabilized GTP-MTs (Fig. S8). The mean velocities were 19.0 ± 0.2 nm/s (GMPCPP-MTs) and 9.4 ± 0.1 nm/s (taxol-MTs), respectively. Although the velocity of Eg5 was higher on GMPCPP-MTs than on GTP-MTs, a mixture of Eg5 and both kinds of shortened MTs displayed all three of the distinct patterns observed in the system composed of Eg5 and normal GTP-MTs (Fig. S9 and Table S1), suggesting that modulation of MT properties does not have a critical effect on the types of spatiotemporal patterns. However, the correspondence between the observed patterns and the concentration range of Eg5 was different (Fig. S9 and Table S1). Since the density of the MTs did not change with the use of shortened MTs (Fig. S7 C), this shift in the transition concentration of Eg5 probably resulted from differences in the lengths of the MTs.

**Effects of motor properties on network dynamics**

We next investigated how the spatiotemporal dynamics of MT networks can be affected by the activity of a motile cross-linker. To address this question, we altered the motor properties by replacing the motor head of Eg5 with that of KIF5B (Fig. S10, A and B), which is similar to a construct used in a previous study (32). This chimeric motor (KIF5Bhead-Eg5tail) exhibited processivity along MTs with a 10-fold higher velocity than the wild-type Eg5 protein, and
with less affinity for MTs (Fig. S10, C–F). The mixture of MT and KIF5B head-Eg5 tail showed temporal evolution patterns distinct from those of the MT-Eg5 system (Fig. 4), suggesting that the activity of the motile cross-linker critically affected the MT network dynamics. The eventual steady-state patterns of the system revealed many isolated astral structures with an isotropic shape, regardless of the motor concentration used ([KIF5B head-Eg5 tail] = 6.25–37.5 nM).

FIGURE 3 The strong contraction force caused by a motile cross-linker results in MT network fragmentation. (A) Time-series images showing the formation of a globally connected contractile MT network ([tubulin] = 1 μM, [Eg5] = 45 nM). Images of ATTO647N-labeled MTs (magenta), ATTO565-labeled MTs (yellow), and GFP-fused Eg5 (cyan) are merged. The times displayed in the images represent the time that elapsed after mixing. (B) Vector field representation of node velocities in the aggregation shown in (A). The velocity of each node is indicated by the length of the arrows and the color scale. (C) Temporal evolution pattern of the mean node velocities in the aggregation shown in (A). (D) Temporal evolution pattern of the size of the total cluster and the largest cluster.

FIGURE 4 Spatiotemporal dynamics of the MT network driven by KIF5B head-Eg5 tail. Time-lapse images show the temporal evolution of isolated asters. Images of taxol-stabilized ATTO647N-labeled MTs (magenta), ATTO565-labeled MTs (yellow), and KIF5B head-Eg5 tail (cyan) are merged. (A–C) The KIF5B head-Eg5 tail concentration was 6.25 nM (A), 12.5 nM (B), or 37.5 nM (C).
These results are consistent with a previous study that used a similar chimeric construct (32) and artificially oligomerized kinesins (29,30). This dynamic pattern including the formation of such astral clusters is referred to as isolated asters. Although the spatial patterns of the final state were similar, the intermediate temporal patterns were dependent on the motor concentration. When the motor concentration was low ([KIF5Bhead-Eg5tail] = 6.25 nM), the motors localized to form asters. However, these asters were distributed so sparsely that a globally connected network did not form (Fig. 4 A; Movie S7). At an intermediate concentration ([KIF5Bhead-Eg5tail] = 12.5 nM), the motors accumulated to form numerous active nodes, which were interconnected through bundled MTs to form a global network. The formed network immediately started to contract, but the links between the active nodes broke readily upon contraction, such that the network structure collapsed into isolated asters (Fig. 4 B; Movie S6). At a high motor concentration ([KIF5Bhead-Eg5tail] = 37.5 nM), the MTs assembled on separate asters even before the start of observations (Fig. 4 C).

Coarse-grained modeling of the spatiotemporal patterns observed experimentally

To understand the mechanism underlying the spatiotemporal dynamics of the MT-motor system and identify essential factors of the system, we developed a coarse-grained model for the contractile MT networks (Figs. 5 A and S12; see also the “Coarse-grained model” section in Materials and Methods) and examined the temporal evolution of MT networks under various motor concentrations with fixed motor properties (Movies S6, S7, and S8).

In our model, MT networks were described by a graphical representation with a set of nodes and links (Figs. 5 A and S12, A and B). Since almost all experimentally observed patterns included astral structures that exerted contractile forces, we regarded each network node as a contractile unit with radially extending links (an active node). Each link represents bundles of MTs interconnecting the active nodes.

Motor concentrations were assigned to each node, and links were elastically extended depending on the motor concentration at each node (Eqs. 2 and 3). Because plus-end-directed motors move toward the centers of asters and tend to be localized (Fig. 1, A and B), we assumed that the motors move against the gradient of motor concentration with a constant flux $f$ to accumulate in regions of higher concentrations. Furthermore, because neighboring filaments between adjacent nodes are mostly in an antiparallel configuration (Fig. S12 C), we modeled a link between two adjacent nodes to contract through the motors’ sliding activity, which induces overlapping of neighboring filaments. The links were broken when they extended beyond a critical length, $L_c$, in a manner similar to the bond-dilution process that is widely used to investigate biopolymer networks (54).

Fig. 5 B shows representative pattern evolutions with different initial motor concentrations, $C_{ub}$. At a low motor concentration of $C_{ub} = 0.1$, the motors accumulated on a few nodes, and these nodes then attracted surrounding nodes (Fig. 5 B, Static Network, Node Velocity; Movie S8), corresponding to the local contractile dynamics observed in our experiments (Fig. 1 D). Because these active nodes were too sparsely distributed to link to each other, the system remained static, corresponding to the experimentally observed static network (Figs. 1 and S2). As the motor concentration was increased to $C_{ub} = 0.4$, many active nodes became interconnected to form a network that contracted as an entire domain (Fig. 5 B, Active Network, Node Velocity; Movie S9). This global contraction corresponded to the active networks observed in the experiments (Figs. 2 and S3). A slight increase of the motor concentration around $C_{ub} = 0.6$ caused some active nodes to exhibit strong local contracting forces, leading to the breakage of several links. Link breakage, in turn, caused local stress concentrations, resulting in occasional rupturing of the entire domain (Fig. S13 B). At a higher motor concentration of $C_{ub} = 0.7$, the active nodes pulled their surrounding nodes so rapidly upon the mixing of motors with MTs that the network no longer maintained global connectivity. The resultant structures contracted rapidly to form chain-like aggregates (Fig. 5 B, Aggregation, Node Velocity; Movie S10). These results are similar to the experimental patterns observed with the high motor concentration (Figs. 3 and S6). Changes in the total area and the cluster number at a simulation time of $T = 500$, as the motor concentration was varied, are summarized in Fig. 5 C.

Modulation of model parameters reproduced the isolated-asters pattern

Increasing the strength, $K$, of the contraction between adjacent active nodes and the motor flux, $f$, which mimics the elevated sliding activity of the motors, resulted in the formation of multiple asters of small filament-motor aggregates (Fig. 5 B, Isolated Asters, Node Velocity). These isolated asters were created through rapid, strong contractions followed by many link breaks. This dynamics reproduced the observation of isolated asters with KIF5Bhead-Eg5tail (Figs. 4 and S11).

The elastic energy storage in the MT network calculated by coarse-grained modeling recharacterized static networks, active networks, and aggregation

To compare the mechanical behaviors of the modeled MT networks, we calculated the elastic energy stored within them. We found three distinct patterns in the temporal
evolution of the total elastic energy, $E$, stored in the system (Fig. 6). The energy in the static network remained small, whereas the energy in the active network increased to a maximum during the contraction phase and was released through dissipation via link breakage. In the aggregation phase, the network immediately collapsed under self-generated mechanical stress, whereas the energy decreased continuously (Fig. 6 A). Fig. 6 B depicts the time at which the energy peaked, $T_p$ (Fig. 6 A), as the motor concentration was varied. Thus, three distinct phases were clearly identified as a static network ($T_p >>$ observation time), an active network ($T_p$ remaining finite), and aggregation ($T_p \rightarrow 0$).

**DISCUSSION**

In this study, we demonstrated that a simple system composed of MTs and cross-linking motors exhibited a variety of spatiotemporal dynamics (Figs. 1, 2, 3, and 4; Table S1). The patterns were categorized as static networks, active networks, aggregation, and isolated asters, and characterized by the connectivity and contractility of the networks.
The static and active networks were similar in that the MTs were globally connected. However, after network formation, the dynamics of the static network showed a lack of global contractility, whereas the active network displayed contraction throughout the entire structure. Aggregations and active networks shared contractile dynamics, but aggregations showed a lack of global connectivity and only local contractions. In isolated asters, the strong contractility or weak connectivity that resulted from modulation of the motor properties caused connectivity breaks in the network, resulting in localized contraction, even when a globally connected network was formed. Regarding the connectivity and contractility of networks, the elemental structure is the active node, the motor-accumulated point from which MTs extend radially. The active nodes are thought to be formed via reorganization of MTs and motors (Fig. S13 C), based on the cross-linking and sliding activities of motor proteins, and have MTs with uniform polarity surrounding around them. Since the MTs in the active nodes have a defined polarity, contraction forces emerge between the adjacent active nodes linked through MT bundles, and this contractile interaction between active nodes is the origin of network contractility.

The geometric configuration of active nodes may play some role in the temporal evolution of the network dynamics. In the static network, the active nodes were distributed so sparsely that links between contractile active nodes rarely formed. In the active network, connected links of active nodes formed and rendered the network capable of undergoing global contractility. During aggregation, densely distributed active nodes formed local clusters and exerted strong contractile forces to break linkages among the clusters. The importance of the geometrical configuration of active nodes is further supported by evidence that differences in the initial spatial distribution of motors correlated with the subsequent temporal evolution pattern (Fig. S5). However, it is difficult to control the initial distribution of active nodes in the system used here. The relationship between the geometry of the node positions and the temporal evolution patterns of the networks will be addressed in more detail in future work.

Furthermore, the spatiotemporal dynamics of the MT-motor network were affected by modulation of the MT and motor properties (Figs. 4 and S9). The results obtained using GMPCPP-MTs and shortened MTs suggested that the experimental modulation of MT properties had little effect on the types of patterns observed (Fig. S9), although shifts in the transition concentration of the motor protein occurred. In contrast, the results obtained with the chimeric KIF5B<sub>head</sub>-Eg5<sub>tail</sub> protein indicated that modulation of the

**FIGURE 6** Mechanical properties of the model MT networks. (A) Temporal changes in the total elastic energy (E) stored in the system for the static network (C<sub>ub</sub> = 0.1), active network (C<sub>ub</sub> = 0.59), and aggregation (C<sub>ub</sub> = 1.0) patterns. (B) Relationship between the time at which E reaches a local maximum, T<sub>p</sub>, and the motor concentration, C<sub>ub</sub>. Bars reflect the standard deviation of the mean value for 100 repeated simulations, starting with randomized initial distributions of the motor concentration. The model parameters are the same as those in Fig. 5 B.

**FIGURE 7** Contractility and connectivity of MT networks. (A) Dependence of contractility and connectivity on the Eg5 concentration in the experiments. The mean and standard deviation are shown. Number of observed chambers: one (4.5 nM), one (8.9 nM), five (14.9 nM), three (19.8 nM), five (22.5 nM), one (29.7 nM), and five (45 nM), respectively. (B) Dependence of contractility and connectivity on the motor concentration, C<sub>ub</sub>, in the simulations. Average values are shown, with bars reflecting the SD of the mean value for 100 repeated simulations, starting with randomized initial distributions of the motor concentration.
motor properties critically affected the spatiotemporal dynamics, which led to a different type of pattern, namely, isolated asters.

On the basis of the spatiotemporal dynamics of the MT-kinesin network, we developed a simple model of network contraction that reproduced our experimentally observed patterns. To construct a simple model of the macroscopic contractile patterns of filament-motor networks on a scale of several millimeters over ~2 h, we employed a coarse-grained description in which the MTs were represented by a set of extensible links without buckling, based on our experimental observations that showed no apparent buckling of the bundles around active nodes (Fig. S13 A). Our model suggests that two essential features account for our experimental findings. First, motors must move between adjacent nodes to accumulate and form active nodes; otherwise, they fail to exert significant contractile forces. Second, the magnitude of contraction between nodes must depend on the motor concentration in a threshold-dependent manner (Eqs. 2 and 3). Without this nonlinear dependency, the model network would experience excessive link breakages during active node formation. The typical spatiotemporal dynamics of the modified model, where the contraction strength depends linearly on the motor concentration, is shown in Fig. S13 C. The modified model predicted the formation of large holes in the network during contraction over nearly all concentrations of the initial motor density, and eliminated the aggregation phase at a high motor concentration (Fig. S13 D). These results conflicted with our experimental observations, suggesting that nonlinearity is necessary to maintain network connectivity in global contractile dynamics. Furthermore, recent theoretical work on the actomyosin network (61) also provides support for our assumption of nonlinear dependency, showing that the net contractile force in the actomyosin system emerges in a threshold-dependent manner.

In addition, our model showed an energy-storage capacity of active networks (Fig. 6). During the temporal evolution of an active network, the interconnection of active nodes prevents local link breakages and enables the storage of elastic energy in the entire network, except when occasional ruptures occur (Figs. S3 C and S13 B). From a biological point of view, contraction of the MT network and energy storage might contribute to increased mechanical stability of cells when the network periphery is cramped, so that it cannot contract, but rather can span tightly throughout the whole cell surface.

Although contraction principles for actin-myosin networks have been widely studied experimentally (62–65) and theoretically (66–71), surprisingly few studies of the contraction mechanism of MT networks have been published. Foster et al. (41) recently demonstrated the contractile dynamics of large-scale MT networks in Xenopus extracts and modeled them using active hydrodynamic theory, providing insight into the macroscopic cytoskeletal organization (42). The in vitro motor-filament system that we studied here is composed of simpler and well-characterized components, i.e., MTs and a single kind of active cross-linker. This system exhibits not only similar contractile dynamics but also network ruptures, which can be understood by using an elastic network model.

Previous studies on MTs focused mainly on the role of MTs in establishing the mitotic spindle and intracellular transport. Recent cell biology studies, however, have provided evidence that network-like MT structures with multiple organizing centers can form, which can be viewed as network nodes in myocytes (43), muscle cells (44), and plants (72), suggesting possible biological functions of MT networks. It is known that in these networks, many MT organizing centers without any obvious organizing apparatus can play an important role in establishing network structures. This study of the MT-motor network with self-organized contractile active nodes will provide a new, to our knowledge, basic framework for understanding the spatiotemporal dynamics of MT networks with multiple acentrosome-organizing centers.

SUPPORTING MATERIAL

Supporting Materials and Methods, 13 figures, one table, and 10 movies are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(16)30446-5.

AUTHOR CONTRIBUTIONS

T.T., D.T., and K.O. designed the study. T.T. designed and performed the experiments. T.T. and D.T. performed the image analysis. D.T. designed and performed the simulations. All of the authors contributed to the writing of the manuscript.

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REFERENCES

1. Alberts, B., A. Johnson, ...., P. Walter. 2002. Molecular Biology of the Cell, 4th ed. Garland Science, New York.
2. Karsenti, E. 2008. Self-organization in cell biology: a brief history. Nat. Rev. Mol. Cell Biol. 9:255–262.
3. Huber, F., J. Schnaüß, ...., J. Käs. 2013. Emergent complexity of the cytoskeleton: from single filaments to tissue. Adv. Phys. 62:1–112.
4. Mimori-Kiyosue, Y. 2011. Shaping microtubules into diverse patterns: molecular connections for setting up both ends. Cytoskeleton (Hoboken). 68:603–618.
5. de Forges, H., A. Bouissou, and F. Perez. 2012. Interplay between microtubule dynamics and intracellular organization. Int. J. Biochem. Cell Biol. 44:266–274.
6. Vignaud, T., L. Blanchoin, and M. Théry. 2012. Directed cytoskeleton self-organization. Trends Cell Biol. 22:671–682.
7. Walczak, C. E., and R. Heald. 2008. Mechanisms of mitotic spindle assembly and function. Int. Rev. Cytol. 265:111–158.
8. Hirokawa, N., Y. Noda, ..., S. Niwa. 2009. Kinesin superfamily motor proteins and intracellular transport. Nat. Rev. Mol. Cell Biol. 10:682–696.
9. Dent, E. W., and K. Kilil. 2001. Axon branching requires interactions between dynamic microtubules and actin filaments. J. Neurosci. 21:9757–9769.
10. Kilil, K., and E. W. Dent. 2005. Touch and go: guidance cues signal to the growth cone cytoskeleton. Curr. Opin. Neurobiol. 15:521–526.
11. Shaw, S. L., R. Kamnyar, and D. W. Ehrhardt. 2003. Sustained microtubule treadmill in Arabidopsis cortical arrays. Science. 300:1715–1718.
12. Dixit, R., and R. Cyr. 2004. The cortical microtubule array: from dynamics to organization. Plant Cell. 16:2546–2552.
13. Lindebloom, J. J., M. Nakamura, ..., D. W. Ehrhardt. 2013. A mechanism for reorientation of cortical microtubule arrays driven by microtubule severing. Science. 342:1245533.
14. Kim, S., and P. A. Coulombe. 2010. Emerging role for the cytoskeleton as an organizer and regulator of translation. Nat. Rev. Mol. Cell Biol. 11:75–81.
15. Karsenti, E., and I. Vernos. 2001. The mitotic spindle: a self-made machine. Science. 294:543–547.
16. Wittmann, T., A. Hyman, and A. Desai. 2001. The spindle: a dynamic assembly of microtubules and motors. Nat. Rev. Cell Biol. 3:E28–E34.
17. Gadde, S., and R. Heald. 2004. Mechanisms and molecules of the mitotic spindle. Curr. Biol. 14:R797–R805.
18. Goshima, G., and J. M. Scholey. 2010. Control of mitotic spindle length. Annu. Rev. Cell Dev. Biol. 26:21–57.
19. Subramanian, R., and T. M. Kapoor. 2012. Building complexity: insights into self-organized assembly of microtubule-based architectures. Dev. Cell. 23:874–885.
20. Keating, T. J., and G. G. Borisy. 1999. Centrosomal and non-centrosomal microtubules. Biol. Cell. 91:321–329.
21. Efimov, A., A. Khartonov, ..., I. Kaverina. 2007. Asymmetric CLASP-dependent nucleation of noncentrosomal microtubules at the trans-Golgi network. Dev. Cell. 12:917–930.
22. Kapitein, L. C., and C. C. Hoogenraad. 2015. Building the neuronal microtubule cytoskeleton. Neuron. 87:492–506.
23. Paradze, A., A. Wright, and D. W. Ehrhardt. 2006. Microtubule cortical array organization and plant cell morphogenesis. Curr. Opin. Plant Biol. 9:571–578.
24. Oda, Y. 2015. Cortical microtubule rearrangements and cell wall patterning. Front. Plant Sci. 6:236.
25. Jiang, W., G. Jimenez, ..., R. Fukunaga. 1998. PRC1: a human mitotic spindle-associated CDK substrate protein required for cytokinesis. Mol. Cell. 2:877–885.
26. Mollinari, C., J. P. Kleman, ..., R. L. Margolis. 2002. PRC1 is a microtubule-binding and bundling protein essential to maintain the mitotic spindle midzone. J. Cell Biol. 157:1175–1186.
27. Subramanian, R., E. M. Wilson-Kubalek, ..., T. M. Kapoor. 2010. Insights into antiparallel microtubule crosslinking by PRC1, a conserved nonmotor microtubule binding protein. Cell. 142:433–443.
28. Dehmelt, L., and S. Halpain. 2005. The MAP2/Tau family of microtubule-associated proteins. Genome Biol. 6:204.
29. Nédélec, F. J., T. Surrey, ..., S. Leibler. 1997. Self-organization of microtubules and motors. Nature. 389:305–308.
30. Surrey, T., F. Nédélec, ..., E. Karsenti. 2001. Physical properties determining self-organization of motors and microtubules. Science. 292:1167–1171.
31. Sanchez, T., D. T. Chen, ..., Z. Dogic. 2012. Spontaneous motion in hierarchically assembled active matter. Nature. 491:431–434.
32. Hentrich, C., and T. Surrey. 2010. Microtubule organization by the antagonistic mitotic motors kinesin-5 and kinesin-14. J. Cell Biol. 189:465–480.
33. Sumino, Y., K. H. Nagai, ..., K. Oiwa. 2012. Large-scale vortex lattice emerging from collectively moving microtubules. Nature. 483:448–452.
34. Karsenti, E., F. Nédélec, and T. Surrey. 2006. Modelling microtubule patterns. Nat. Cell Biol. 8:1204–1211.
35. Marchetti, M. C., J. F. Joanny, ..., A. R. Simha. 2013. Hydrodynamics of soft active matter. Rev. Mod. Phys. 85:1143.
36. Wojcik, E. J., R. S. Buckley, ..., S. Kim. 2013. Kinesin-5: cross-bridging mechanism to targeted clinical therapy. Gene. 531:133–149.
37. Nadar, V. C., A. Ketschek, ..., P. W. Baas. 2008. Kinesin-5 is essential for growth-cone turning. Curr. Biol. 18:1972–1977.
38. Nadar, V. C., S. Lin, and P. W. Baas. 2012. Microtubule redistribution in growth cones elicited by focal inactivation of kinesin-5. J. Neurosci. 32:5783–5794.
39. Falnikar, A., S. Tole, and P. W. Baas. 2011. Kinesin-5, a mitotic microtubule-associated motor protein, modulates neuronal migration. Mol. Biol. Cell. 22:1561–1574.
40. Kahn, O. I., V. Sharma, ..., P. W. Baas. 2015. Effects of kinesin-5 inhibition on dendritic architecture and microtubule organization. Mol. Biol. Cell. 26:66–77.
41. Foster, P. J., S. Fürthauer, ..., D. J. Needleman. 2015. Active contraction of microtubule networks. eLife. 4:4.
42. Belmonte, J. M., and F. Nédélec. 2016. Large-scale microtubule networks contract quite well. eLife. 5:5.
43. White, E. 2011. Mechanical modulation of cardiac microtubules. Pflugers Arch. 462:177–184.
44. Oddoux, S., K. J. Zaal, ..., E. Ralston. 2013. Microtubules that form the stationary lattice of muscle fibers are dynamic and nucleated at Golgi elements. J. Cell Biol. 203:205–213.
45. Malikov, V., E. N. Cytrynbaum, ..., V. Rodionov. 2005. Centering of a radial microtubule array by translocation along microtubules spontaneously nucleated in the cytoplasm. Nat. Cell Biol. 7:1213–1218.
46. Torisawa, T., M. Ichikawa, ..., K. Furuta. 2014. Autoinhibition and cooperative activation mechanisms of cytoplasmic dynein. Nat. Cell Biol. 16:1118–1124.
47. Castoldi, M., and A. V. Popov. 2003. Purification of brain tubulin through two cycles of polymerization-depolymerization in a high-molarity buffer. Protein Expr. Purif. 32:83–88.
48. Furuta, K., and Y. Y. Toyota. 2008. Minus-end-directed motor Ncd exhibits processive movement that is enhanced by microtubule bundling in vitro. Curr. Biol. 18:152–157.
49. Sharp, D. J., K. L. McDonald, ..., J. M. Scholey. 1999. The bipolar kinesin, KLP61F, cross-links microtubules within interpolar microtubule bundles of Drosophila embryonic mitotic spindles. J. Cell Biol. 144:125–138.
50. Kapitein, L. C., E. J. Peterman, ..., C. F. Schmidt. 2005. The bipolar mitotic kinesin Eg5 moves on both microtubules that it crosses. Nature. 435:114–118.
51. Kim, T., W. Hwang, and R. D. Kamm. 2009. Computational analysis of a cross-linked actin-like network. Exp. Mech. 49:91–104.
52. Kim, T., W. Hwang, ..., R. D. Kamm. 2009. Computational analysis of viscoelastic properties of crosslinked actin networks. PLOS Comput. Biol. 5:e1000439.
53. Borau, C., T. Kim, …. R. D. Kamm. 2012. Dynamic mechanisms of cell rigidity sensing: insights from a computational model of actomyosin networks. PLoS One. 7:e49174.
54. Sankararaman, S., G. I. Menon, and P. B. S. Kumar. 2004. Self-organized pattern formation in motor-microtubule mixtures. Phys. Rev. E Stat. Nonlin. Soft Matter Phys. 70:031905.
55. Gowrishankar, K., S. Ghosh, …. M. Rao. 2012. Active remodeling of cortical actin regulates spatiotemporal organization of cell surface molecules. Cell. 149:1353–1367.
56. Broedersz, C. P., and F. C. MacKintosh. 2011. Molecular motors stiffen non-affine semiflexible polymer networks. Soft Matter. 7:3186–3191.
57. Head, D. A., G. Gompper, and W. J. Briels. 2011. Microscopic basis for pattern formation and anomalous transport in two-dimensional active gels. Soft Matter. 7:3116–3126.
58. Levine, A. J., and F. C. MacKintosh. 2009. The mechanics and fluctuation spectrum of active gels. J. Phys. Chem. B. 113:3820–3830.
59. MacKintosh, F. C., and A. J. Levine. 2008. Nonequilibrium mechanics and dynamics of motor-activated gels. Phys. Rev. Lett. 100:018104.
60. Kapitein, L. C., B. H. Kwok, …. E. J. Peterman. 2008. Microtubule cross-linking triggers the directional motility of kinesin-5. J. Cell Biol. 182:421–428.
61. Hiraiwa, T., and G. Salbreux. 2016. Role of turnover in active stress generation in a filament network. Phys. Rev. Lett. 116:188101.
62. Köhler, S., V. Schaller, and A. R. Bausch. 2011. Structure formation in active networks. Nat. Mater. 10:462–468.
63. Koenderink, G. H., Z. Dogic, …. D. A. Weitz. 2009. An active biopolymer network controlled by molecular motors. Proc. Natl. Acad. Sci. USA. 106:15192–15197.
64. Gardel, M. L., J. H. Shin, …. D. A. Weitz. 2004. Elastic behavior of cross-linked and bundled actin networks. Science. 304:1301–1305.
65. Murrell, M. P., and M. L. Gardel. 2012. F-actin buckling coordinates contractility and severing in a biomimetic actomyosin cortex. Proc. Natl. Acad. Sci. USA. 109:20820–20825.
66. Lenz, M., M. L. Gardel, and A. R. Dinner. 2012. Requirements for contractility in disordered cytoskeletal bundles. New J. Phys. 14:14.
67. Lenz, M., T. Thoresen, …. A. R. Dinner. 2012. Contractile units in disordered actomyosin bundles arise from F-actin buckling. Phys. Rev. Lett. 108:238107.
68. Lenz, M. 2014. Geometrical origins of contractility in disordered actomyosin networks. Phys. Rev. X. 4:041002.
69. Jülicher, F., K. Kruse, J. Prost, and J.-F. Joanny. 2007. Active behavior of the cytoskeleton. Phys. Rep. 449:3–28.
70. Voituriez, R., J. F. Joanny, and J. Prost. 2006. Generic phase diagram of active polar films. Phys. Rev. Lett. 96:028102.
71. Kruse, K., J. F. Joanny, …. K. Sekimoto. 2004. Asters, vortices, and rotating spirals in active gels of polar filaments. Phys. Rev. Lett. 93:099902.
72. Ehrhardt, D. W. 2008. Straighten up and fly right: microtubule dynamics and organization of non-centrosomal arrays in higher plants. Curr. Opin. Cell Biol. 20:107–116.