REPORT

Torques within and outside the human spindle balance twist at anaphase

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At each cell division, nanometer-scale motors and microtubules give rise to the micron-scale spindle. Many mitotic motors step helically around microtubules in vitro, and most are predicted to twist the spindle in a left-handed direction. However, the human spindle exhibits only slight global twist, raising the question of how these molecular torques are balanced. Here, we find that anaphase spindles in the epithelial cell line MCF10A have a high baseline twist, and we identify factors that both increase and decrease this twist. The midzone motors KIF4A and MKLP1 are together required for left-handed twist at anaphase, and we show that KIF4A generates left-handed torque in vitro. The actin cytoskeleton also contributes to left-handed twist, but dynein and its cortical recruitment factor LGN counteract it. Together, our work demonstrates that force generators regulate twist in opposite directions from both within and outside the spindle, preventing strong spindle twist during chromosome segregation.

Introduction

At each cell division, the micron-scale spindle self-organizes from nanometer-scale molecular components to divide the genome. While the identities of nearly all these building blocks are known (Neumann et al., 2010), many questions remain about how they together give rise to the architecture, mechanics, and function of the spindle as an ensemble. Mitotic motors, over a dozen species of which are present in the human spindle, illustrate this gap: although the motility and force-generating capacity of many motors have been closely studied in vitro (Canty et al., 2021; Cross and McAinsh, 2014), it remains poorly understood how motors cooperate in the dense microtubule network of the spindle to give rise to larger-scale microtubule architecture.

Several motors have been found to generate net torque on microtubules in vitro, resulting in helical motility around the microtubule track. Mitotic motors have rotational pitches ranging from ~0.3 to 2.3 µm (Walker et al., 1990; Yajima et al., 2008), far more extreme than the supertwist of 13-protofilament microtubules (no supertwist in 13-protofilament microtubules) (Ray et al., 1993). Importantly, many mitotic motors crosslink and slide microtubule pairs, rather than stepping along isolated microtubules. In vitro experiments that recreate bridged microtubule conformations have demonstrated that torques generated by mitotic kinesins are sufficiently strong to twist and coil two microtubules around each other (Meißner et al., 2024; Mitra et al., 2020), converting nanoscale torque generation into twisted microtubule bundle shapes at length scales of many microns. The plus-end-directed yeast kinesin-8 Kip3 (Bormuth et al., 2012; Mitra et al., 2018) and Caenorhabditis elegans kinesin-6 ZEN-4 (Maruyama et al., 2021) both have a left-handed stepping bias, as does the kinesin-5 Eg5 (Yajima et al., 2008), although this motor’s directional preference has recently been called into question (Meißner et al., 2024). By contrast, the minus-end-directed kinesin-14 Ncd (Mitra et al., 2020; Nitschke et al., 2016; Walker et al., 1990) and cytoplasmic dynein have right-handed stepping biases (Can et al., 2014; Elshenawy et al., 2019). These torques would be expected to...
additively twist microtubule pairs in the same direction (Mitra et al., 2020), and by extension, additively promote left-handed spindle twist. However, the human spindle exhibits only a weak left-handed twist on average (Neahring et al., 2021; Novak et al., 2018; Trupinic et al., 2022). It is not known how molecular-scale torques are balanced in the spindle to produce a relatively achiral structure from chiral motors.

The spindle’s left-handed twist was first quantified in metaphase HeLa and U2OS cells (Novak et al., 2018). The twist has been proposed to allow the metaphase spindle to accommodate mechanical load along the pole-to-pole axis (Trupinic et al., 2022), although its functional importance for chromosome segregation remains to be studied. Twist ranges from −0.2°/µm to 2°/µm at metaphase, and the spindle exhibits only slight global twist on average (Neahring et al., 2021; Novak et al., 2018; Trupinic et al., 2022). It is not known how molecular-scale torques are balanced in the spindle to produce a relatively achiral structure from chiral motors.

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Results and discussion

MCF10A spindles exhibit high baseline twist that peaks in late metaphase and anaphase

To study torque regulation in the spindle, we sought to identify a cell line in which spindles exhibited higher baseline twist than that observed in previously characterized cell lines. We reasoned that because twist differs between the human cell lines RPE1, HeLa, and U2OS (Neahring et al., 2021; Novak et al., 2018; Trupinic et al., 2022), other human cell lines may exhibit stronger twist, which would allow us greater dynamic range to study factors that both increase and decrease twist. We quantified twist using a previously published optical flow method (Trupinic et al., 2022) in which we live-imaged full spindle volumes, computationally rotated the images to view the spindle along the pole-to-pole axis, and calculated the displacement of pixel intensities between successive frames from 20 to 70% of the pole-to-pole axis (Fig. 1A). We selected this region of the spindle to avoid the higher microtubule density near spindle poles and for consistency with previous studies (Novak et al., 2018; Trupinic et al., 2022). These flow vectors were converted to polar coordinates and averaged to produce a single helicity value for each spindle. Compared with other methods of quantifying twist, such as manual bundle tracing (Neahring et al., 2021; Novak et al., 2018; Trupinic et al., 2022) or quantification based on bundle angles, the optical flow method is more strongly influenced by image noise but is more easily applied to large numbers of cells (Trupinic et al., 2022). Furthermore, despite the noisiness of the optical flow method, its results were correlated with the results of the other two methods across a range of helicities, indicating that it detects true biological variability in spindle twist (Fig. S1, A and B).

Focusing on anaphase, when we previously observed that spindle twist is differentially regulated (Neahring et al., 2021), we found that the non-transformed mammary epithelial cell line MCF10A (Soule et al., 1990) exhibited strong, visually apparent left-handed twist in unperturbed cells, while anaphase RPE1 and U2OS spindles were not significantly twisted (Fig. 1B). Early- to mid-anaphase MCF10A cells labeled with either overexpressed GFP-tubulin or SiR-tubulin had significant left-handed twist (negative helicity, −0.92 ± 0.88°/µm and −0.76 ± 0.79°/µm, respectively, mean ± SD), and the twist did not significantly differ between the two methods of labeling microtubules (P = 0.702, one-way ANOVA with Tukey’s post-hoc test).

Our comparison between different human cell lines was performed by live confocal imaging of a single timepoint per cell, leading us to wonder how twist changes during mitotic progression in MCF10A spindles. To image dividing cells volumetrically at high time resolution, we used lattice light sheet microscopy (Fig. 1C and D; and Video 1). This imaging modality allowed us to obtain near-isotropic resolution with minimal phototoxicity, ideal for studying three-dimensional spindle architecture over time (Pamula et al., 2019). Comparing the temporal profiles of 12 spindles revealed several insights into both spindle twist and our analysis methods. Helicity values over time were noisy, suggesting that the optical flow quantification method is limited by its sensitivity to image noise. However, even in this dataset where we obtained multiple measurements of each spindle over time, there was considerable variability from cell to cell, with peak helicities ranging from −2.6 to −2.7°/µm. The average twist across all cells in the first 4 min of anaphase, −1.25 ± 0.62°/µm, was stronger than the average value from the confocal dataset, consistent with the improved signal-
to-noise ratio in these images compared with our confocal data. This highlights the fact that helicity magnitudes are not directly comparable between different imaging modalities or tubulin labeling methods, although twist is comparable within a given dataset. On average, the left-handed twist became stronger in the final minutes of metaphase, consistent with previous findings in HeLa and RPE1 cells (Trupinic et al., 2022). Twist was maintained throughout early and mid-anaphase for ~3 min, before dissipating in late anaphase, typically coinciding with a slower phase of spindle elongation (Fig. 1, C and D). Given the high baseline twist in MCF10A spindles, and given the sustained period of stronger twist in the first few minutes of anaphase, we focused thereafter on anaphase MCF10A cells to study how torques are generated and resisted in the spindle.

Figure 1. MCF10A spindles exhibit high baseline twist that peaks in late metaphase and anaphase. (A) Schematic diagram of spindle twist quantification, using the method developed by Trupinic et al. (2022). Three-dimensional image stacks were rotated to view the spindle along the pole-to-pole axis. Optical flow was computed between successive frames, and flow vectors were converted to polar coordinates and averaged for each spindle (see Materials and methods). (B) Spindle helicity (average degrees rotated around the pole-to-pole axis per µm displacement along the pole-to-pole axis) at anaphase in three human epithelial (RPE1, MCF10A) or epithelial-like (U2OS) cell lines, calculated from GFP-α-tubulin or SiR-tubulin intensity. Negative values represent left-handed helicity, and positive values represent right-handed helicity. Black lines represent mean ± SD. $n = 19, 27, 50$, and 51 spindle pools from $N = 2, 4, 5$, and 5 independent experiments for RPE1 GFP-tub, U2OS GFP-tub, MCF10A GFP-tub, and MCF10A SiR-tub, respectively. n.s. not significant, ****$P = 1.50 \times 10^{-9}$ (MCF10A GFP-tub) and $P = 1.11 \times 10^{-8}$ (MCF10A SiR-tub), one-sample t tests comparing each sample to a mean of 0. (C) Lattice light sheet images of the same MCF10A cell, labeled with SiR-tubulin, at four different timepoints (related to Video 1). The xy view (center) shows maximum intensity projections of the entire spindle region. The yz view (right) shows maximum intensity projections between 30% and 70% of the pole-to-pole axis for the same image volumes after rotating them by 90°. Colors indicate directions of Farneback optical flow vectors, according to the color legend shown in the top image. Scale bars = 3 µm. (D) Length (upper panel) and helicity (lower panel) over time of 12 MCF10A spindles, calculated from time-lapse lattice light sheet images. The center line and shaded region represent the mean and 95% confidence interval.

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Twist in the anaphase spindle

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The midzone motors KIF4A and MKLP1 contribute to the anaphase spindle’s left-handed twist

We next asked what factors give rise to the spindle’s left-handed twist at anaphase. Although the motors Eg5 and KIF18A have been shown to promote left-handed twist at metaphase (Novak et al., 2018; Trupinic et al., 2022), many mitotic motors undergo changes in localization and function at anaphase, and the molecular basis of anaphase spindle twist has not been studied. Anaphase spindle elongation in human cells is powered by several kinesins that localize to antiparallel microtubule overlaps in midzone bundles, where they slide microtubules apart (Vukusic et al., 2017; Yu et al., 2019). KIF4A, Eg5, and the kinesin-6 motors MKLP1 and MKLP2 all redundantly contribute to spindle elongation (Fig. 2 A) (Vukusic et al., 2017, 2021), and we hypothesized that these kinesins may generate anaphase-specific left-handed torques.

Eg5 and the C. elegans homolog of MKLP1 have been shown to step in a left-handed fashion in vitro (Maruyama et al., 2021; Yajima et al., 2008), but the chirality of KIF4A has not been studied. Due to its ability to slide antiparallel microtubules apart (Hannabuss et al., 2019; Wijeratne and Subramanian, 2018), its anaphase-specific localization to midzone microtubule overlaps (Kurasawa et al., 2004), and its contribution to anaphase spindle elongation (Vukusic et al., 2021), KIF4A is a good candidate for a left-handed torque generator in the anaphase spindle. Thus, we characterized the torque generation of KIF4A on suspended microtubule bridges between 2 μm-diameter beads immobilized on a coverslip (Can et al., 2014) (Fig. 2 B). Smaller-sized (0.51 μm-diameter) cargo beads were decorated by multiple kinesin motors, brought close to the microtubule bridge with an optical trap, and their motility was tracked in three dimensions using brightfield microscopy (Fig. S2 A). We first used a truncated version of human kinesin-1 (KIF5B, amino acids 1–560, referred to as K560) to validate our experimental approach since K560 is known to follow a single protofilament on its microtubule track (Can et al., 2014; Ray et al., 1993; Yajima and Cross, 2005). We observed that cargo beads coated with K560 exhibited a combination of left-handed (6.6 ± 0.4 μm pitch; five beads, five rotations), right-handed (4.7 ± 1.4 μm pitch; three beads, three rotations), and straight (15 beads) movements (Fig. 2 C), consistent with the reported pitch lengths of microtubules with 14, 12, or 13 protofilaments, respectively (Hyman et al., 1995; Ray et al., 1993). Unlike kinesin-1, all KIF4A-driven beads exhibited left-handed motility with a shorter pitch of 2.1 ± 0.7 μm (10 beads, 14 rotations; two-tailed t test, P = 10−4; Fig. 2 D), demonstrating that KIF4A generates left-handed torque on microtubules.

We next sought to test whether the motors powering anaphase elongation contribute to left-handed spindle twist in their cellular context. We first tested the individual contributions of these motors by depleting KIF4A, MKLP1, or MKLP2 or inhibiting Eg5 with S-trityl-L-cysteine (STLC). We confirmed motor depletion via Western blot and by scoring individual cells for the expected late anaphase and cytokinesis phenotypes (Fig. S2, B–H). None of these perturbations significantly affected anaphase spindle twist (Fig. 2 E; and Fig. S2, I and J). When we codepleted KIF4A and MKLP1, however, spindles were significantly less twisted with a mean helicity of −0.34 ± 0.54°/μm (Fig. 2, E and F; Fig. S2 D, and Video 2). This suggests that similarly to their roles in elongating the anaphase spindle, the midzone motors KIF4A and MKLP1 together generate left-handed torques to twist the anaphase spindle.

It is possible that spindle motors could modulate spindle twist by altering spindle shape or other microtubule properties, in addition to or instead of directly exerting torques on spindle microtubules. Indeed, a previous study noted that rounder HeLa spindles tended to exhibit stronger twist, although this correlation did not extend to the RPE1 cell line (Trupinic et al., 2022). We quantified anaphase spindle shape after depleting the motors KIF4A or MKLP1 and found that KIF4A knockdown was associated with slight changes in spindle shape: siKIF4A spindles were slightly wider and had a lower length-to-width ratio, on average (Fig. S1, C–E). However, spindle helicity was not significantly correlated with spindle length, width, or aspect ratio (Fig. S1, F–H). These results suggest that KIF4A does not regulate spindle twist simply by changing the shape of the anaphase spindle, consistent with a functional role for its capacity to generate torque on microtubules.

Actin promotes left-handed spindle twist at anaphase

Motor localization in the spindle midzone is only one of several changes that occurs at the metaphase-to-anaphase transition. Interactions between the spindle and the actin cytoskeleton also become more pronounced: actin regulates the anaphase cortical enrichment of the spindle positioning machinery (Kotak et al., 2013, 2014), the spindle midzone provides a spatial cue for actin assembly at the site of the future contractile ring (Pollard and O’Shaughnessy, 2019), and actin has recently been reported to accumulate at centrosomes and between separating chromosomes at anaphase onset (Farina et al., 2019; Kita et al., 2019). Intriguingly, the actin cytoskeleton is intrinsically chiral in cultured human fibroblasts (Tee et al., 2015). Thus, we wondered whether the actin cytoskeleton could contribute to the human spindle’s chiral twist at anaphase.

We disrupted the actin cytoskeleton by treating MCF10A cells with latrunculin A (LatA) and found that this abrogated the anaphase spindle’s left-handed twist (−0.24 ± 0.84°/μm compared with −0.78 ± 0.63°/μm after DMSO treatment). By contrast, treating cells with the myosin II inhibitor blebbistatin or the ROCK inhibitor Y27632 did not affect the spindle twist (Fig. 3, A and B; and Video 3). We confirmed that LatA and blebbistatin disrupted actomyosin contractility by imaging cells later in anaphase, when they inhibited cytokinetic forw...
The midzone motors KIF4A and MKLP1 contribute to the anaphase spindle’s left-handed twist. (A) Schematic diagram of the midzone motors KIF4A (green), MKLP1 (blue), and Eg5 (purple) that cooperate to drive anaphase spindle elongation (gray “F” and arrows). (B) Schematic diagram of the experimental geometry of the in vitro microtubule bridge assay (see Materials and methods, not to scale). A fluorescently labeled microtubule was suspended between two beads of 2 µm diameter. A 0.51-µm diameter cargo bead was densely coated by multiple kinesins and brought onto the microtubule bridge by an optical trap (not shown) and bead motility was imaged using brightfield illumination. (C) Left: Example 3D trajectory of a kinesin-1 (K560)-coated cargo bead.
(Fig. 3 D). Although LatA has been reported to have no effect on the twist of metaphase HeLa spindles (Novak et al., 2018), our results in MCF10A and RPE1 cells indicate that the actin cytoskeleton contributes to left-handed spindle twist in anaphase.

Dynein counteracts left-handed twist in the anaphase spindle

The motor-generated torques characterized to date are expected to additively twist the spindle, and the effects of the actin cytoskeleton described above contribute to twist in the same left-handed direction (Fig. 3). Thus, we sought to understand what factors are required to oppose left-handed torques so that the spindle exhibits only slight global twist. We previously found that NuMA and Eg5 coinhibition in RPE1 spindles increases the spindle’s left-handed twist (Fig. 3 D; Neahring et al., 2021). Since inhibiting Eg5 alone had no effect in that system, we hypothesized that NuMA and its interactors dynactin and dynein (Fig. 4 A) play key roles in resisting spindle twist. Consistent with this hypothesis, siRNA-mediated depletion of dynein heavy chain indeed increased left-handed twist in anaphase MCF10A spindles (Fig. 4, D and E; and Fig. S3, B and D).

We next probed the mechanism by which dynein counteracts left-handed twist. At mitosis, NuMA recruits dynactin and dynein to microtubule minus ends, where they act as a complex to cluster minus ends at spindle poles (Gaglio et al., 1996; Heald et al., 1996; Hueschen et al., 2017; Merdes et al., 1996; Verde et al., 1991). NuMA–dynactin–dynein complexes also localize to the cell cortex, where they generate pulling forces on astral microtubules to position the spindle (Kotak et al., 2012) (Fig. 4 A). Based on mammalian dynein’s in vitro stepping behavior with other adaptors (Elshenawy et al., 2019), its torque generation inside the spindle would be predicted to augment the spindle’s left-handed twist. Because we observed the opposite phenotype upon dynein depletion, we reasoned that dynein likely does not regulate spindle twist by generating torques between spindle microtubule pairs. Instead, we considered the possibility that the cortical pool of NuMA–dynactin–dynein could counteract the spindle’s left-handed twist, either by passively resisting or actively opposing it.

To test the cortical pool of dynein, we depleted LGN, one of several factors that recruits NuMA–dynactin–dynein complexes to the cortex during anaphase (Fig. S3 C) (Du and Macara, 2004; Kiyomitsu and Cheeseman, 2013; Kotak et al., 2014; Seldin et al., 2013). We confirmed that upon LGN depletion, NuMA’s cortical localization was reduced in anaphase MCF10A cells (Fig. 4, B and C). LGN knockdown increased the spindle’s left-handed twist to an average of $-1.20 \pm 0.92^\circ/\mu\text{m}$, significantly stronger than that of control-depleted cells and almost as strong as that of dynein-depleted cells (Fig. 4, D and E; and Video 4). To ask whether this effect was anaphase-specific, we assessed the twist in LGN-depleted and dynein-depleted metaphase spindles (analyzing those with mild or absent pole focusing defects) and found that neither perturbation significantly altered spindle twist at metaphase (Fig. 4 F). Because astral microtubules elongate and dynein is enriched at the cell cortex during anaphase (Collins et al., 2012; Kiyomitsu and Cheeseman, 2013; Kotak et al., 2013), this finding provides further evidence that the cortical pool of dynein is required to counteract left-handed spindle twist.

In this study, we investigated how the spindle attains its relatively untwisted architecture despite being built by chiral force-generators. Focusing on anaphase, when we find that the spindle’s strongest twist is sustained for several minutes (Fig. 1), we identified factors that promote left-handed twist (Figs. 2 and 3) and that counteract it (Fig. 4). Unexpectedly, we find that twist is regulated not just by motors internal to the spindle but also by spindle positioning factors at the cell periphery. Together, our results demonstrate that spindle twist is an emergent phenomenon that integrates inputs from the spindle’s broader cellular environment.

Motors that crosslink, slide, and twist microtubules are abundant in the spindle and have been the focus of most work on spindle twist to date. We find that the midzone motors KIF4A and MKLP1 contribute to left-handed twist additively or redundantly, similar to their overlapping contributions to spindle elongation (Vukusic et al., 2021), reflecting a common design principle in the anaphase spindle. Both KIF4A and MKLP1 can slide antiparallel microtubules apart in vitro (Hannabuss et al., 2019; Nislow et al., 1992; Wijeratne and Subramanian, 2018), both step in a left-handed direction around microtubules (Fig. 2) (Maruyama et al., 2021), and both localize and concentrate at the midzone at anaphase onset (Kurasawa et al., 2004; Matuliene and Kuriyama, 2002), suggesting that they may increase twist at anaphase by directly exerting torques on overlapping antiparallel microtubules. However, we cannot exclude that KIF4A and/or MKLP1 could regulate twist indirectly via effects on the midzone’s microtubule dynamics or organization, potentially altering its material properties or the localization of downstream factors. Directly linking motor-generated torques to global spindle twist awaits the development of mutant motors with altered torque-generating capacities, an exciting future direction.

Opposing the effects of KIF4A and MKLP1, dynein and its targeting factors NuMA and LGN are each required to restrain the anaphase spindle’s left-handed twist. Although dynein inhibition does not increase twist in metaphase spindles (Fig. 4 F,
Trupinic et al., 2022), we observe strong phenotypes after dynein depletion in anaphase MCF10A cells or NuMA knockout in RPE1 cells. LGN–NuMA–dynein complexes at the cortex could reduce twist by exerting active right-handed torques on astral microtubules (Fig. 4 G, Model 1) by pulling outwards on astral microtubules and spindle poles (Model 2) or by increasing astral microtubules’ anchorage in the cortex (Model 3). Although we cannot rule out that pools of NuMA and dynein within the spindle also regulate spindle twist, they do not appear to play a major role since dynein depletion does not affect metaphase twist and since twist in dynein-depleted anaphase spindles does not significantly differ from that of LGN-depleted anaphase spindles (Fig. 4). Dynein and its cofactors illustrate that motors can regulate spindle twist not just by twisting spindle microtubules around each other, but also by mediating interactions between the spindle and cell periphery.

Finally, we find that the actin cytoskeleton promotes the anaphase spindle’s left-handed twist, both in MCF10A cells and in NuMA- and Eg5-inhibited RPE1 cells (Fig. 3). This effect may arise from the cortical pool of actin, although it would act independently from the LGN–NuMA–dynein cortical force-generating machinery since they influence spindle twist in

**Figure 3.** Actin promotes left-handed spindle twist at anaphase. (A) Confocal images of live MCF10A cells treated with 0.1% DMSO, 500 nM latrunculin A, or 25 µM blebbistatin labeled with SiR-tubulin (see also Video 3). Maximum intensity projections of a 2-µm thick low region (magenta) and a 2-µm thick high region (green) relative to the spindle midplane are overlaid. Dashed lines highlight individual microtubule bundles in each region. The helicity of each spindle is indicated in the top right. Positions of spindle poles are indicated by white circles. Scale bars = 3 µm. (B) Helicity of anaphase spindles calculated from SiR-tubulin intensity. Black lines represent mean ± SD. n = 54, 43, 36, and 32 spindles pooled from N = 6, 5, 4, and 3 independent experiments for DMSO, LatA, blebbistatin, and Y27632, respectively. n.s. not significant, **P = 0.0039, one-way ANOVA with Tukey’s post-hoc test. (C) Confocal images of late anaphase MCF10A cells. 500 nM LatA and 25 µM blebbistatin treatment each inhibit cytokinetic furrow progression. Brightfield images (upper row) represent a single z-plane and SiR-tubulin images (lower row) are maximum intensity projections of 10 µm z-stacks. Scale bars = 5 µm. (D) Helicity of anaphase RPE1 spindles, synchronized with RO-3306, calculated from GFP-tubulin intensity. The control and NuMA-KO+STLC conditions are the same cells included in a previous publication (Neahring et al., 2021), reanalyzed using the optical flow method. Black lines represent mean ± SD. n = 22, 18, and 7 spindles pooled from N = 4, 5, and 3 independent experiments for control, NuMA-KO+STLC, and NuMA-KO+STLC+LatA, respectively. **P = 9.99 × 10⁻⁴, ****P = 4.32 × 10⁻¹⁶ (control versus NuMA-KO+STLC) and ****P = 5.93 × 10⁻⁶ (control versus NuMA-KO+STLC+LatA), one-way ANOVA with Tukey’s post-hoc test.
Figure 4. Dynein counteracts left-handed twist in the anaphase spindle. (A) Schematic diagram of LGN, NuMA, and dynein localization in anaphase cells. Dynein and NuMA cluster microtubule minus ends and localize to spindle poles, while LGN–NuMA–dynein complexes localize to cortical crescents where they exert pulling forces on astral microtubules. The purple arrow indicates the direction of dynein stepping, and the gray arrow indicates the direction of force on astral microtubules. (B) Immunofluorescence images (single z-planes) of anaphase MCF10A cells stained for NuMA (magenta) and DNA (blue). Scale bars = 5 µm. (C) Quantification of NuMA enrichment at the cell cortex relative to NuMA intensity in the cytoplasm (see Materials and methods), in immunofluorescence images of MCF10A cells transfected with siRNA targeting luciferase (control) or LGN. The dashed line indicates a cortex/cytoplasm ratio of 1, i.e., no cortical enrichment. Black lines represent mean ± SD. n = 22 cells from 2 independent days in each condition. ****P = 1.55 × 10^{-10}, two-sample t-test. (D) Confocal images of live MCF10A cells transfected with siRNA targeting luciferase (siCtrl), dynein heavy chain, or LGN labeled with SiR-tubulin (see also Video 4). Maximum intensity projections of a 2-µm thick low region (magenta) and a 2-µm thick high region (green) relative to the spindle midplane are overlaid. Dashed
lines highlight individual microtubule bundles in each region. The helicity of each spindle is indicated in the top right. Positions of spindle poles are indicated by white circles. Scale bars = 3 µm. (E) Helicity of anaphase spindles calculated from SiR-tubulin intensity. Black lines represent mean ± SD. n = 65, 45, and 78 spindles pooled from N = 6, 5, and 7 independent experiments for siCtrl, siDHC, and siLGN, respectively. ***P = 9.43 × 10^{-5}, **P = 9.03 × 10^{-4}, one-way ANOVA with Tukey’s post-hoc test. (F) Helicity of metaphase spindles calculated from SiR-tubulin intensity. Black lines represent mean ± SD. n = 39, 38, and 35 spindles pooled from N = 4, 4, and 2 independent experiments for siCtrl, siDHC, and siLGN, respectively. n.s., not significant, one-way ANOVA with Tukey’s post-hoc test. (G) Proposed models for twist regulation in the anaphase spindle. Motors within the spindle midzone generate left-handed torques (blue arrows), which are partially counteracted by LGN-NuMA-dynein complexes at the cell cortex. These cortical complexes could actively generate right-handed torques on astral microtubules (magenta arrows, Model 1), exert outward pulling on spindle poles (Model 2), or serve to anchor astral microtubules in the cortex (Model 3). Together, these active and passive torques establish weak global left-handed twist in the anaphase spindle.

opposite directions. Alternatively, pools of actin within and around the anaphase spindle could regulate its twist. In the future, further molecular dissection of actin-related proteins will provide insight into the role of actin uncovered here, and many myosins, formins, and actin filament themselves are intrinsically chiral (Ali et al., 2002; Depue and Rice, 1965; Lebreton et al., 2018; Mizuno et al., 2011). Although the underlying mechanisms are not yet clear, our finding that the actin cytoskeleton affects twist is exciting because it reveals that multiple cytoskeletal systems coordinately regulate twist.

In conclusion, our study shows that the human anaphase spindle’s weak left-handed twist requires both left-handed torque generators and factors that oppose them. The study of spindle twist is a recent area of inquiry, and many open questions remain. It is unclear why twist differs between cell types and species (Velle et al., 2022), or whether twist is modulated by other mechanisms such as microtubule crosslinking, the spindle’s material properties (Forth and Kapoor, 2017), the distribution of parallel and antiparallel microtubules, or the turnover rates of microtubules and microtubule-associated proteins (Asthana et al., 2021; do Rosario et al., 2023). Finally, it will be interesting to explore the potential functions of spindle twist: are there adverse consequences for chromosome segregation if the spindle is too twisted, or not twisted enough? Twist could provide a mechanical advantage to the anaphase spindle as it separates sister chromosomes, but conversely, our previous work has shown that strongly twisted NuMA- and Eg5-inhibited RPE1 spindles have high rates of chromosome segregation errors (Nehring et al., 2021). More broadly, our work motivates the study of how other cellular structures built from chiral molecular components either co-opt this chirality for their physiological function (for example, chiral actin flows in left-right symmetry breaking) (Naganathan et al., 2016) or balance chiral elements to restrain asymmetry.

Materials and methods

Cell culture

U2OS cells (female human osteosarcoma cells) were a gift from Samara Reck-Peterson (University of California San Diego, San Diego, CA, USA), and hTERT–RPE1 cells (female human retinal epithelial cells) were a gift from Bo Huang (University of California San Francisco, San Francisco, CA, USA). Both cell lines were cultured in DMEM/F12 (11320; Thermo Fisher Scientific) supplemented with 10% fetal bovine serum (10438026; Gibco). MCF10A cells (female human mammary epithelial cells) were purchased from ATCC (CRL-10317) and cultured as recommended by ATCC in MEGM (CC-3150; Lonza) supplemented with bovine pituitary extract, insulin, hydrocortisone, and human epidermal growth factor according to the manufacturer’s instructions, and 100 ng/ml cholera toxin (CS052; Sigma-Aldrich). Inducible NuMA-KO RPE1 cells stably expressing GFP-tubulin and mCherry-H2B (Nehring et al., 2021) were grown in DMEM/F12 supplemented with 10% tetracycline-screened FBS (PS-FB2; Peak Serum) and 5 μg/ml puromycin to select for the sgRNA targeting NuMA. SpCas9 expression was induced by adding 1 μg/ml doxycycline hyclate (D9891; Sigma-Aldrich) 4 days before each experiment and refreshed after 24 and 48 h. All cells were maintained at 37° and 5% CO2.

Transfection, dyes, and small molecule treatments

For siRNA knockdowns, cells were transfected with siRNA targeting luciferase as a negative control (5'-GUAGCGGAAUACUUGA-3', 50 pmol), LGN (Dharmacon ON-TARGETplus SMARTpool, L-004092-00-0005, 100 pmol), dynein heavy chain (5'-AAG GATCAACATGACGGAAT-3', 50 pmol) (Draviam et al., 2006; Tanenbaum et al., 2008), MKL1 (pool of three sequences, sc-35936; Santa Cruz Biotechnology, 50 pmol), MKL2 (5'-ACA CAGGCCUUGAUGAG-3', Dharmacon J-004957-06-0002, 50 pmol), or siKIF4A (pool of three sequences, sc-60888; Santa Cruz Biotechnology, 50 pmol) for 48 h using Lipofectamine RNAiMAX (13778075; Thermo Fisher Scientific) according to the manufacturer’s recommendations.

GFP-α-tubulin was expressed in RPE1, U2OS, and MCF10A cells by infection with BacMam virus. The GFP-α-tubulin coding sequence was cloned into the pEG BacMam vector (plasmid #160451, Addgene; a gift from Eric Gouaux, Oregon Health & Science University, Portland, OR, USA), recombinant bacmid DNA was generated in DH10Bac cells (1036102; Thermo Fisher Scientific), and isolated bacmid DNA was transfected into Si9 cells (a gift from Yifan Cheng, University of California San Francisco, San Francisco, CA, USA) using Cellfectin II (10362100; Thermo Fisher Scientific) for production and amplification of BacMam virus according to a previously described protocol (Goehring et al., 2014). P2 BacMam virus was added to cells 2 days prior to imaging. Alternatively, tubulin was labeled by adding 100 nM SiR-tubulin and 10 μM verapamil for 30–60 min prior to imaging (Inc. CY-SC002; Cytoskeleton). F-actin was labeled by transfecting MCF10A cells with 0.5 μg mEmerald-Lifeact-7 (plasmid #54148, Addgene; a gift from Michael Davidson, Florida State University, Tallahassee, FL, USA) for 3 days using Viafect (E4981; Promega) according to the manufacturer’s recommendations.
For acute drug treatments, latrunculin A (L12370; Invitrogen) was added to a final concentration of 500 nM for 20 min prior to imaging. (-)-blebbistatin (203391; Sigma-Aldrich) was added to a final concentration of 25 µM for 30 min prior to imaging. Y27632 (688001; Sigma-Aldrich) was added to a final concentration of 10 µM for 30 min prior to imaging. STLC (164739; Sigma-Aldrich) was added to a final concentration of 10 µM for 15 min prior to imaging, and DMSO was added to a final concentration of 0.1% (vol/vol) for 30 min prior to imaging.

For experiments in the RPE1 inducible NuMA-KO cell line (Fig. 3 D), cells were synchronized at the G2/M checkpoint by incubation overnight in 9 µM of the CDK1 inhibitor RO-3306 (SML0569; Sigma-Aldrich). Cells were released into mitosis by washing 4X in warm media and were imaged from prometaphase (~30 min in controls) or after reaching the turbulent state (to confirm NuMA knockout; ~60 min in doxycycline NuMA-KO cells). STLC and latrunculin A in the indicated conditions were added 1 h after RO-3306 washout to final concentrations of 5 µM and 500 nM, respectively. Cells that entered anaphase within 90 min of drug addition were used for analysis.

Western blotting
Cells grown in six-well plates were lysed and protein extracts were collected after centrifugation at 4°C for 30 min. Protein concentrations were measured using a Bradford assay kit (Bio-Rad) and equal concentrations of each sample were separated on 4–12% Bis-Tris gels (NP0321; Invitrogen) by SDS-PAGE and transferred to a nitrocellulose membrane. Membranes were blocked with 4% milk in TBST (tris-buffered saline + 0.1% Tween 20) for 1 h, incubated with primary antibodies overnight at 4°C, and incubated with HRP-conjugated secondary antibodies for 45 min. Proteins were detected using SuperSignal West Pico or Femto chemiluminescent substrates and imaged using a ChemiDoc Manager v2.0.0.

Immunofluorescence
Cells were plated on round 25 mm coverslips (CG15XH; Thorlabs) coated with 200 nm fluorescent beads (Invitrogen FluoSpheres objective, 1.0 NA, 421452-9800-000) were immersed in media. Cells were imaged on a spinning disk (CSU-X1, Yokogawa) confocal inverted microscope (Eclipse Ti-E, Nikon Instruments) with the following components: 100× 1.45 NA, 561 nm (100 mW), and 642 nm (100 mW) diode lasers; ET455/50M, ET525/50M, ET630/75M, and ET690/50M emission filters (Chroma Technology); and a Zyla 4.2 sCMOS camera (Andor Technology). Z-stacks consisting of 21 planes spaced 0.5 µm apart were acquired for each cell with Micro-Manager v2.0.0.

Lattice light sheet microscopy
The LLSM was a modified version of the microscope described in Liu et al. (2018) and was controlled with custom LabVIEW software licensed from Janelia Research Campus, HHMI. Cells were plated on round 25 mm coverslips (CG15XH; Thorlabs) coated with 200 nm fluorescent beads (Invitrogen Fluospheres Carboxylate-Modified Microspheres, Ex/Em 660/680, F8807) to measure point spread functions for deconvolution and to align the lattice light sheet. The coverslip, excitation objective (Thorlabs water dipping objective, 0.60 NA, TL20×-MPL), and detection objective (Zeiss Plan-Apochromat 20× water dipping objective, 1.0 NA, 421452-9800-000) were immersed in ~50 ml of phenol red-free MCF10A culture medium maintained at 37°C and 5% CO₂. Cells were labeled with 100 nM SiR-tubulin and 10 µM verapamil and imaged using a 642 nm laser (MPB Communications Inc 2RU-VFL-P-2000-642-BIR) operating with 200 µW input power at the back pupil of the excitation objective. The 3D volumes consisting of 100 planes were acquired with 50 ms exposure per plane and iterating every 30 s by scanning the sample stage (SmarAct MLS-3252 Electromagnetic Direct-Drive) with a 400 nm step (corresponding to ~215 nm along the optical z-axis). A dithered harmonic-balanced hexagonal lattice light sheet (LLS) pattern with a numerical aperture of 0.35 and a sigma value of 0.09 was used (Liu et al., 2023). This LLS yields a 3D theoretical resolution of 340 × 340 × 570 nm at
the 680 nm emission, which ensures that visualization and analysis of 3D spindle dynamics are not compromised by the axis of the poorest resolution. A mask with an aperture of Na_{min} 0.3/Na_{max} 0.4 was used to block the unmodulated light and the higher-order diffractions. Emission light was filtered by a Semrock FF02-685/40-25 bandpass filter and captured by a Hamamatsu ORCA-Fusion sCMOS camera. Images were deconvolved, deskewed, and rotated on a high-performance computing cluster using code available at https://github.com/abcuckerberkeley/LLSMSDTools/.

**Suspended microtubule bridge assay**

**Protein preparation**

The truncated human kinesin-1 (KIFSB, amino acids 1–560) was expressed and purified using the baculovirus expression system in insect cells. GFP was fused to the C-terminus of kinesin to link motors to the antibody-coated beads. The recombinant full-length human KIF4A-GFP was expressed and purified in Sf9 cells as described (Subramanian et al., 2013). Microtubules were polymerized in BRB80 buffer using a mixture of LB655-labeled tubulin and unlabeled pig brain tubulin at a 1:4 ratio in the presence of 100 µM taxol.

**Labeling cargo beads with antibodies**

Carboxyl latex beads (Invitrogen) with a diameter of 0.51 µm were coated with anti-GFP antibodies as previously described (Belyy et al., 2014). The beads were initially pelleted and resuspended in activation buffer (10 mM MES, 100 mM NaCl, pH 6.0). The carboxyl groups on the bead surfaces were functionalized with amine-reactive groups via N-ethyl-N'-(3-(dimethylamino)propyl)carbodiimide (EDC) and sulfo-N-hydroxysuccinimide (sulfo-NHS, Thermo Fisher Scientific) crosslinking for 20 min at 4°C. Extra crosslinking reagents were then removed by centrifugation, and beads were resuspended in PBS at pH 7.4. Anti-GFP polyclonal antibodies (CA 5314/15; Covance, custom produced) were added to the beads and incubated at room temperature for 20 min. The surface of the beads was passivated by incubation with 5 mg/ml bovine serum albumin (BSA) overnight at 4°C. Excess antibodies were removed by centrifugation. Beads were then resuspended in PBS with 0.1% azide for storage at 4°C.

**Preparation of flow chambers**

Custom PEG–biotin coverslips were prepared as previously described (Zhao et al., 2023). Streptavidin-coated beads (2 µm diameter, Spherotech) were introduced into the chamber and incubated for 3 min to allow the beads to settle on the biotinylated coverslip surface. The chamber was washed with 15 µl MB. Equal volumes of anti-GFP-coated cargo beads and 0.5 µM GFP-tagged motor proteins were mixed and incubated for 5–10 min on ice. 10 µl of MB with 2 mM ATP was added to the bead-motor mixture and flowed into the chamber. The cargo beads moved unidirectionally along the microtubule long axis, demonstrating that the microtubule bridges were formed by a single microtubule.

**Data collection**

Experiments were performed with a custom-built optical trapping microscope equipped with a Nikon TiE microscope body, Nikon 100× 1.49 NA Plan-Apochromat objective, and Orca Flash 2.0 CMOS camera (Hamamatsu, Japan). LB655-labeled microtubules were excited with a 632 nm laser beam (Melles Griot) in epifluorescence mode. The fluorescence signal was detected by the camera with an effective pixel size of 43.3 nm after magnification. Movies were recorded at 3 Hz. The coverslip surface was scanned to identify a microtubule bridge that was stable and oriented parallel to the imaging plane between two 2-µm-diameter beads. The fluorescence image of the microtubule was brought into focus to capture the full 3D motion of the bead around the circumference of the microtubule. A 0.51-µm-diameter cargo bead freely diffusing in solution was trapped by a focused 1,064 nm beam (IPG Photonics) and brought close to the microtubule. When the cargo bead exhibited unidirectional motility on the microtubule, the trapping laser was turned off, and the cargo bead was released to determine the polarity of the microtubule bridge. The cargo bead was then placed at the minus end of the microtubule to track its plus-end-directed motility throughout the entire length of the bridge. Bead movement was recorded using brightfield microscopy.

**Analysis of bead trajectories**

The center positions of the cargo beads were determined using two-dimensional Gaussian fitting in ImageJ. Z-positions of the beads were calibrated by decorating the coverslip surface with 0.51-µm-diameter beads and moving the microscope objective ±255 nm in the z-direction using a PIFOC objective scanner (Physik Instrumente) with 30-nm increments. Intensities were recorded at each z-position for all selected beads to obtain a calibration curve (Fig. S2 A). Z-positions of the cargo beads in motility assays were then calculated from the calibration curve based on the intensities of the cargo beads. The 3D trajectory of each cargo bead was plotted using custom-written software in MATLAB (MathWorks) and the helical pitch and handedness of each trajectory were calculated manually.

**Quantification of spindle helicity**

Spindles in early–to-mid-anaphase, with two clearly separated chromosome masses but before the onset of furrowing, were chosen for analysis. Spindles were rotated so that the pole-to-pole axis was horizontal and cropped using the rectangle tool in FIJI (Schindelin et al., 2012). The positions of the two poles were manually assigned based on tubulin intensity. Spindles were resliced along the pole-to-pole axis by permuting the [x,y,z] coordinates to [y,z,x] in MATLAB.
Helicity was quantified using a previously published optical flow method (Trupinic et al., 2022). Instead of using the optical flow pipeline’s Laplacian of Gaussian filter, we did not filter the deconvolved lattice light sheet images, and filtered confocal images by subtracting an image blurred with a Gaussian filter of standard deviation 30 pixels (scipy.ndimage.gaussian_filter1d) followed by despeckling with a three-pixel median filter (scipy.ndimage.median_filter). Briefly, Farnebäck optical flow (Farnebäck, 2003) was calculated between each pair of successive frames lying between 30% and 70% of the pole-to-pole axis. Flow vectors were converted to polar coordinates, weighted by pixel intensities using the “All pixels weighted helicities” method, and averaged for each spindle.

To compare helicity quantification methods in Fig. S1, the bundle tracing method was performed as previously described (Neahring et al., 2021) (a modified version of the method first published by Novak et al. [2018]). For the bundle angle method, the lowest and highest planes of a z-stack were identified that contained clearly visible microtubule bundles. In these two planes, the angles of three bundles were manually measured using the line tool in FIJI and averaged. Helicity was calculated using the formula:

$$\frac{\theta_{\text{bottom}} - \theta_{\text{top}}}{d},$$

where d is the diameter of the spindle in μm and θ is the average angle between midzone bundles and the pole-to-pole axis.

Quantification of spindle shape and cortical fluorescence intensity

Spindle length was determined by calculating the distance between the manually assigned pole positions used for helicity analysis. Width was manually measured at the spindle equator from maximum intensity projections of the tubulin channel. The aspect ratio was calculated as spindle length divided by spindle width.

NuMA fluorescence intensity at the cell cortex was calculated from the single z-plane where cortical intensity appeared brightest near each spindle pole. In FIJI, short line segments 20 px (1.16 μm) wide were drawn across the cortex, through the cytoplasm, and outside the cell (background) near each spindle pole, a total of six line segments per cell. Using the “plot profile” function in FIJI, the maximum intensity was extracted from each line segment. The corresponding values from each pole were averaged, the background value was subtracted from the cortex and cytoplasm values, and the cortex/cytoplasm intensity ratio was calculated.

Statistical analysis

Parametric tests were used based on the assumption that spindle helicity and shape are approximately normally distributed, with approximately equal variance between experimental conditions, although these assumptions were not formally tested. In Fig. 1B, distributions were assessed for a significant difference from 0 helicity with one-sample two-tailed t tests using the ttest function in MATLAB. Helicities, cortical intensities, and spindle shape parameters between experimental conditions were compared using one-way ANOVA with post-hoc Tukey-Kramer tests, using the anova1 and multcompare functions in MATLAB, or using two-sample two-tailed t tests with the ttest2 function in MATLAB when only two groups were compared. We used P < 0.05 as a threshold for statistical significance. The number of cells, the number of independent experiments, and P values are provided in figure legends.

Online supplemental material

Fig. S1 compares three methods of quantifying spindle twist and describes spindle shape across the molecular perturbations used in the study. Fig. S2 contains experiments to validate the depletion of KIF4A, MKLP1, and MKLP2 based on Western blotting and spindle phenotypes, and provides quantification of spindle twist after perturbation of the additional kinesins MKLP2 and Eg5. Fig. S3 contains additional imaging and Western blot data to validate Latrunculin A and siRNA depletions of dynein heavy chain and LGN. The online supplemental material also includes four videos showing an example spindle imaged with lattice light sheet microscopy (Video 1), and top-down and end-on views of spindles imaged with spinning disk confocal microscopy (Videos 2, 3, and 4). Video 2 includes control and siMKLP1+siKIF4A spindles, Video 3 includes DMSO-, latrunculin A-, and blebbistatin-treated spindles, and Video 4 shows control-, dynein-, and LGN-depleted spindles.

Data availability

The data underlying all plots have been deposited at Mendeley Data, at https://doi.org/10.17632/tm2r6vfwm.1, and are publicly available as of the date of this publication.

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Methodology, Validation, Visualization, Writing - review & editing, J. Fernandes: Resources, Writing - review & editing, C.J. Rux: Investigation, Writing - review & editing, K. Nakos: Resources, Writing - review & editing, R. Subramanian: Resources, Supervision, Writing - review & editing, S. Upadhyayula: Funding acquisition, ESI, Supervision, Writing - review & editing, A. Yildiz: Conceptualization, Funding acquisition, Supervision, Writing - review & editing, A. Komar: Conceptualization, Funding acquisition, Supervision, Writing - review & editing.

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Figure S1. **Comparison of helicity quantification methods and comparison of spindle shape across molecular perturbations.** (A) Schematic diagrams illustrating three methods of quantifying spindle helicity. The optical flow method is calculated from all pixels in an end-on (XZ) view of the spindle, the bundle tracing method is calculated from ∼10 bundles per cell manually traced from an end-on (XZ) view of the spindle, and the bundle angle method is calculated from six bundles in a top-down (XY) view of the spindle. (B) Scatterplots comparing the helicity of the same 18 siCtrl (gray) and 14 siLGN (black) cells analyzed by each of the three quantification methods. Pearson’s correlation coefficients are shown on each plot. (C–E) Violin plots with internal boxplots of spindle length (C), width (D), and aspect ratio (length/width, E) for spindles subjected to molecular perturbations. n.s. not significant, *P < 0.05, **P < 0.005, ***P < 0.0005, ****P < 0.00005, one-way ANOVA with Tukey’s post-hoc test. (F–H) Scatterplots of spindle length (F), width (G), and aspect ratio (H) versus spindle helicity, averaged for each experimental condition. Vertical and horizontal gray lines indicate standard deviation. Helicity is not significantly correlated with spindle length, width, or aspect ratio (two-sided Pearson’s correlation test). Pearson’s correlation coefficients are provided in the lower right of each plot.
Figure S2. Validation of midzone kinesin experiments. (A) In the suspended microtubule bridge assay, the z-position of the cargo bead was calibrated by measuring the average intensity of 17 surface-immobilized cargo beads by scanning the microscope objective from −255 to 255 nm relative to the imaging plane. Error bars represent SD. The intensity profile was fitted to a third-order polynomial (red curve, $R^2 = 0.9995$) to obtain the calibration curve. (B–E) Western blots of KIF4A (B), MKLP1 (C), KIF4A and MKLP1 (D), or MKLP2 (E) levels in MCF10A cells transfected with siRNA targeting luciferase (control), KIF4A, MKLP1, or MKLP2 for 48 h, as indicated. GAPDH is shown as a loading control. (F) Live confocal images of the late anaphase phenotype after KIF4A knockdown. In late anaphase, siKIF4A spindles over-elongate and have poorly organized midzone bundles. Scale bars = 5 µm. Brightfield (upper row) and SiR-tubulin (lower row) images represent single z-planes. (G) Spindle length in late anaphase, after the cleavage furrow is visible. Black lines represent mean ± SD. $n = 14$ siCtrl and 11 siKIF4A spindles. ***$P = 1.34 \times 10^{-4}$, two-sample t test. (H) Live confocal images of the telophase phenotypes after KIF4A or MKLP1 knockdown. The midbody is extended (siKIF4A) and cells have cytokinesis defects (siMKLP1). Scale bars = 5 µm. Brightfield (upper row) and SiR-tubulin (lower row) images represent single z-planes. (I) Helicity of MCF10A anaphase spindles calculated from SiR-tubulin intensity. Black lines represent mean ± SD. $n = 34$ and 35 spindles pooled from $N = 4$ independent experiments each for siCtrl and siMKLP2. n.s., not significant, two-sample t test. (J) Helicity of anaphase spindles in MCF10A cells treated with 0.1% DMSO or 10 µM STLC, calculated from SiR-tubulin intensity. Black lines represent mean ± SD. $n = 42$ STLC-treated spindles pooled from $N = 4$ independent experiments, and the same 54 DMSO-treated spindles shown in Fig. 3B. n.s., not significant, two-sample t test. Source data are available for this figure: SourceData FS2.
Video 1. **Volumetric imaging of an MCF10A spindle.** Full spindle volumes were acquired every 30 s at near-isotropic resolution using lattice light sheet microscopy. Microtubules are labeled with SiR-tubulin. Time (in seconds) is shown relative to anaphase onset (t = 0). At four timepoints, the viewing perspective is rotated to an end-on view and panned along the pole-to-pole axis to visualize twist. Scale bar = 3 µm. See also Fig. 1.

Video 2. **Top-down and end-on views of control and siMKLP1+siKIF4A MCF10A spindles.** Microtubules are labeled with SiR-tubulin. Top-down views (top row) pan from the lowest to the highest planes of a z-stack acquired with spinning disk confocal microscopy, with each plane spaced 0.5 µm apart. End-on views (lower row) show the same z-stacks after 90° rotation and image processing (see Materials and methods), panning toward the viewer (144 fps). See also Fig. 2.
Video 3. **Top-down and end-on views of MCF10A spindles treated with DMSO, latrunculin A, and blebbistatin.** Microtubules are labeled with SiR-tubulin. Top-down views (top row) pan from the lowest to the highest planes of a z-stack acquired with spinning disk confocal microscopy, with each plane spaced 0.5 µm apart. End-on views (lower row) show the same z-stacks after 90° rotation and image processing (see Materials and methods), panning toward the viewer (144 fps). See also Fig. 3.

Video 4. **Top-down and end-on views of MCF10A control-, dynein, and LGN-depleted spindles.** Microtubules are labeled with SiR-tubulin. Top-down views (top row) pan from the lowest to the highest planes of a z-stack acquired with spinning disk confocal microscopy, with each plane spaced 0.5 µm apart. End-on views (lower row) show the same z-stacks after 90° rotation and image processing (see Materials and methods), panning toward the viewer (144 fps). See also Fig. 4.