Dual pathway spindle assembly increases both the speed and the fidelity of mitosis

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Summary

Roughly half of all animal somatic cell spindles assemble by the classical prophase pathway, in which the centrosomes separate ahead of nuclear envelope breakdown (NEBD). The remainder assemble by the prometaphase pathway, in which the centrosomes separate following NEBD. Why cells use dual pathway spindle assembly is unclear. Here, by examining the timing of NEBD relative to the onset of Eg5-mEGFP loading to centrosomes, we show that a time window of 9.2 ± 2.9 min is available for Eg5-driven prophase centrosome separation ahead of NEBD, and that those cells that succeed in separating their centrosomes within this window subsequently show >3-fold fewer chromosome segregation errors and a somewhat faster mitosis. A longer time window would allow more cells to complete prophase centrosome separation and further reduce segregation errors, but at the expense of a slower mitosis. Our data reveal dual pathway mitosis in a new light, as a substantive strategy that increases both the speed and the fidelity of mitosis.

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Introduction

The mitotic spindle is a dynamic, self-assembling protein machine whose main task in the cell is to segregate sister chromatids accurately. Every animal somatic cell spindle assembly by one of two possible pathways: the classical prophase pathway, in which the centrosomes migrate to opposite sides of the nucleus ahead of nuclear envelope breakdown (NEBD), or the prometaphase pathway, in which the centrosomes separate after NEBD by a more complex process that includes contributions from kinetochore and cortex-mediated mechanisms (Rattner and Bens, 1976; Rosenblatt, 2005; Rosenblatt et al., 2004; Toso et al., 2009; Waters et al., 1993; Whitehead et al., 1996).

The prophase and prometaphase pathways are not arbitrary points on a continuum of spindle assembly pathways, but are topologically and temporally distinct and genetically separable. In the prophase pathway, the centrosomes migrate to opposite sides of the nucleus ahead of NEBD, so that when rupture of the nuclear membrane occurs, the basic bipolarity of the spindle is already set up, with the chromatid pairs sandwiched between the equivalent poles. In the prometaphase pathway, this is not so: the centrosomes are both on the same side of the chromatin at NEBD, and different mechanisms must be used to assemble a bipolar spindle under conditions in which the component microtubules of the spindle are already interacting with the chromosome arms and the kinetochores. Mechanisms specific to the prometaphase pathway include kinetochore-based mechanisms (Toso et al., 2009), actomyosin-dependent pulling of the poles towards the cell cortex (Rosenblatt, 2005) and chromatinduced microtubule nucleation (the Ran pathway) (Caudron et al., 2005).

To classify cells as following the prophase or prometaphase pathways, we used a previously-established criterion (Toso et al., 2009) which asks simply, do the chromosomes lie between the two poles at the moment of NEBD? This simple classification serves our present purpose well, because it establishes a prophase pathway population that can be formally compared with all remaining cells, which we classify collectively as prometaphase pathway. Our classification will if anything underestimate any benefits deriving from prophase centrosome separation (see later), because the population of prometaphase pathway cells contains cells with a broad range of intermediate stages of centrosome separation, including for example those that have not begun to separate their centrosomes, and those in which the centrosomes are well separated but in which both centrosomes are nonetheless on the same side of the chromatin at NEBD.

It is clear that mitotic progression is not entirely dependent on centrosome separation, because mitosis, and indeed full development, still occurs in flies lacking centrosomes (Basto et al., 2006). In mammalian somatic cells also, laser ablation of the centrosomes in prophase has been shown not to inhibit bipolar spindle formation (Khodjakov et al., 2000). Nonetheless, centrosome migration in prophase serves to establish the bipolar geometry of the spindle ahead of nuclear envelope breakdown, and it is important to find out why cells do this. We were therefore keen to explore the question, do mitotic progression and mitotic outcome differ between the prophase and prometaphase pathways? Here we have addressed this question by direct live cell imaging of a large number of individual cells.

Results and Discussion

To compare progression and outcomes along the prophase and prometaphase pathways, we established live-cell imaging of a stable HeLa cell line expressing mCherry-α-tubulin (to mark
microtubules) and transiently transfected it with full-length human Eg5 (a kinesin-5) tagged with monomeric EGFP (mEGFP) under the control of a low-expressing SV40 promoter. Control measurements from live-cell movies established that ectopic expression of tagged Eg5 does not perturb mitotic progression or bipolar spindle assembly and anaphase onset compared to an empty vector control (Fig. S1). This system allows us to visualise both spindle assembly and the dynamics of Eg5. High resolution live cell imaging shows that Eg5-mEGFP is loaded on to the centrosomes (the incipient spindle poles) in early prophase (Fig. 1A). The loading of Eg5-mEGFP to the centrosomes not only equips them to separate by motor-driven microtubule sliding, but also marks a point in mitotic time very close to the start of the mechanical programme of mitotic spindle assembly. Previous studies using immunofluorescence have shown that the loading of myc-tagged human and *Xenopus* Eg5 onto centrosomes requires it to be phosphorylated by Cdk1-CyclinB (Blangy et al., 1995; Sawin and Mitchison, 1995). Treatment with roscovitine, a small molecule inhibitor of Cdk1, confirms that both the loading of Eg5-mEGFP to the centrosome, and its continued presence there, are controlled by a roscovitine-sensitive mitotic kinase (Fig. S2). It follows that the instant at which Eg5-mEGFP begins to load to the centrosomes provides a fiducial point in mitotic time, giving us a defined start-point for the process of Eg5-driven centrosome separation, as well as an opportunity to index the timing of subsequent events in mitotic progression, including NEBD. Time lapse tracking of the Eg5-mEGFP signal, subsequent to this start point, reports both the fractional population of Eg5 at the centrosomes, and the position and velocity of the centrosomes as prophase progresses.

Plots of the distance between centrosomes at the time point immediately before NEBD showed no correlation with NEBD onset time (Fig. 1C), strongly suggesting that the mitotic clock controlling NEBD indeed runs entirely independently of Eg5-driven centrosome separation. As a more direct test, we used EI III, an Eg5-specific small molecule inhibitor (dimethylenastron). EI III specifically inhibits the motor domains of Eg5, driving them, like monastral (Crevel et al., 2004), into a motor. ADP state that binds only weakly to microtubules (Cochran et al., 2005; Crevel et al., 2004; DeBonis et al., 2003; Luo et al., 2004). In wild type cells, and in our stable mCherry-x-tubulin HeLa cell line also, treatment with 1 µM EI III blocks centrosome separation, producing, as expected from previous studies (Gartner et al., 2005; Mayer et al., 1999; Tanenbaum et al., 2008), mono-astral spindles (Fig. 1D,E). Quantitation reveals that whilst 1 µM EI III substantially reduces the total amount of EGFP-Eg5 that loads to the centrosomes (Fig. 1B), it does not affect either the normalized kinetics of loading, or the duration of the interval between the start-point of Eg5 loading and NEBD (Fig. 1F, yellow box). These data confirm that in mitotic Hela cells, the timing of NEBD is uncoupled from the process of Eg5-driven centrosome separation.

Our data indicate that the interlude between the start of Eg5 loading and NEBD is 9.2 ± 2.9 min (mean ± SD; n=82; Fig. 1B). The existence of this time window for kinesin-5-driven centrosome separation suggested to us that cells race to complete centrosome separation ahead of NEBD, and that prophase pathway cells are those that succeed, whilst prometaphase pathway cells are those that fail. To examine which factors determine success or failure, we next asked whether in cells that lose the race to complete prophase centrosome separation (those that go on to use the prometaphase pathway), any individual process in centrosome separation is especially slow and potentially limiting. We measured three parameters, the time-lag between the start of Eg5 loading and the start of centrosome motion (the lag time), the mean duration of centrosome motion (the translocation time), and the mean velocity of centrosome motion.

The initial lag reduces the time available for centrosome translocation. The mean initial lag is not significantly different (p<0.01) between prophase (3.0±2.0 min, Fig. 2A) and prometaphase pathway cells (4.1±2.8 min, Fig. 2D). It is possible that this initial lag represents the time required for Eg5 to establish active crosslinks between microtubules issuing from opposite poles. Imaging between the nucleus and the substrate allows visualisation of the evolving antiparallel array of sliding microtubules and supports this idea, in that Eg5-mEGFP is seen to enrich specifically to the overlapped region (Fig. 3A to D; supplementary video 1). Immunofluorescent imaging with anti-Eg5 antibodies and anti-α-tubulin confirm that endogenous Eg5 shows the same localisation pattern (Fig. 3E and F). The translocation time, the mean duration of centrosome migration in prophase, is significantly longer in prophase pathway cells than in prometaphase pathway cells (Fig. 2B,E, p<0.0001). The mean velocity of centrosome separation is also significantly increased in prophase pathway cells (Fig. 2C,F, p<0.01). Individual cell histories (Fig. 2G) reveal no obvious correlations between the lag period, the translocation time and the centrosome velocity, indicating that no individual process is dominantly rate-limiting, but rather that cells that fail to complete prophase pole separation tend to show a longer lag, a lower velocity of centrosome separation and a shorter translocation time.

Given that centrosome separation in prophase is apparently dispensable, why do cells do it? One possibility is that prophase centrosome separation produces a speed-gain during the subsequent stages of mitosis. Using the initiation of Eg5 loading to the centrosomes as a marker in mitotic time, we find, consistent with previous work (Toso et al., 2009) that prophase pathway cells form bipolar spindles 3–4 min more quickly than prometaphase pathway cells (Fig. 4A). This small time advantage, in a total of ∼30 minutes, might provide a drive towards the prophase pathway. But we suspected a second possibility, that prophase centrosome separation improves the overall fidelity of mitotic chromosome segregation.

Previous work has shown that if cells are blocked in a monopolar state with Eg5 inhibitors, and then released, there is a concurrent increase in the incidence of segregation errors (Mailhes et al., 2004; Thompson and Compton, 2008), seen as lagging chromosomes during anaphase (Bakhoun et al., 2009). This increased error rate is thought to be a consequence of an increased number of merotelic microtubule-kinetochore attachments, that in turn are caused by the monopolar spindle geometry. However, inhibition-release experiments do not test whether the transient monopolar state (bracketed state in Fig. 4A) that occurs during normal, unperturbed, bipolar spindle formation via the prometaphase pathway also causes an increase in anaphase segregation errors. To test this possibility, we followed mitosis by live cell imaging in a large number of cells and determined firstly whether each cell used the prophase pathway or prometaphase pathway and secondly whether each subsequently made a segregation error (seen as one or more
Fig. 1. Eg5 loading to centrosomes indexes the mitotic clock. (A) Successive views of an mCherry-α-tubulin HeLa cell, transiently expressing Eg5-mEGFP. Images were acquired every 2 min. T=0 is assigned as the first frame in which Eg5 loading becomes detectable. Internal consistency was checked by averaging all sequences, and showed that the corresponding frame in the averaged time-course was the first frame in which the Eg5-mEGFP signal was statistically significantly (P<0.01) brighter than that in the preceding frame (asterisks in Figure 1A & 1F). Lag time is that between T=0 and initiation of centrosome separation. Translocation time is that between initiation of centrosome separation and NEBD. NEBD, the moment of nuclear envelope breakdown (NEBD), delineating the end of prophase, is defined as the first frame in which mCherry-α-tubulin fluorescence is apparent inside the nuclear volume. This cell used the prometaphase pathway; see supplementary video 2. (B) Time window between the onset of Eg5-loading and NEBD. The mean is 9.2±2.9 min (n=82). (C) Time window duration and centrosomes interdistance are uncorrelated in both the prophase (filled circles) and prometaphase (open circles) pathways, showing that an NEBD countdown timer operates independently of centrosome separation distance. (D) Eg5 loading to centrosomes in the presence of an Eg5-specific small molecule inhibitor (1 μM EI III; conditions as in (A)); see supplementary video 3. Centrosome separation is blocked. (E) Centrosome separation distance versus time in the absence (filled circles) and in the presence (open squares) of EI III. (F) Normalised Eg5-mEGFP intensity on the centrosomes in the absence (filled circles) and presence (open squares) of EI III. All values are shown as mean ± SD. The time window for prophase centrosome separation (yellow boxes) operates identically in the presence of absence of EI III.
lagging chromosomes) during anaphase. In 3 independent experiments on a total of 1388 mitotic HeLa cells, we found a total of 21 lagging chromosomes (see representative movie frames in Fig. 4C). Of these, a total of 5/685 prophase pathway cells (0.7%) showed a lagging chromosome and a total of 16/703 prometaphase pathway cells (2.3%) showed a lagging chromosome. These data reveal, crucially, that prophase centrosome separation indeed increases the fidelity of chromosome segregation.

Since prophase centrosome separation increases the fidelity of mitosis, and helps thereby to maintain genome stability, why not send all cells along the prophase pathway? Our data (Fig. 4B)
show that prophase centrosome migration completes with roughly exponential kinetics (rate constant $= 0.16 \, \text{min}^{-1}$, half time $= 4.3 \, \text{mins}$) so that for $90\%$ of cells to complete centrosome separation before NEBD would require $\sim 20 \, \text{minutes}$, and for $99\%$ to complete, $\sim 35 \, \text{minutes}$. Clearly, improving the success-rate for prophase centrosome separation is desirable, but to do so, cells would need to spend considerably more time waiting for prophase centrosome separation to complete. We speculate that there is an evolutionary drive tending to minimise the overall time spent in mitosis. If so, cells face a dilemma: they can either allow more time for prophase centrosome separation, thereby reducing the mitotic error-rate, or they can reduce the time spent in prophase centrosome separation, tolerate a slightly higher error rate, and complete mitosis faster. As we have shown, cells solve this potential dilemma by requiring that prophase centrosome separation races against a 9.2 minute countdown clock. By using this mechanism, the cell population is split approximately 50:50 between the prophase and prometaphase pathways and the cells thereby achieve both a substantial overall improvement in the fidelity of chromosome segregation and an appreciable overall acceleration of mitosis (see schematic in Fig. 4A). Were less time to be allocated to prophase centrosome separation, fewer

Fig. 3. Eg5 localizes to anti-parallel microtubules that bridge between the two centrosomes in early prophase. (A to C) Time course of centrosome separation in an mCherry-α-tubulin HeLa cell transiently transfected with mEGFP-Eg5. Images shown are 30s apart. (A) Eg5-mEGFP, (B) mCherry-α-tubulin and (C) merge. The cell used the prophase pathway. (D) Enlarged views of Eg5-mEGFP and mCherry-α-tubulin at 0:30 min; see supplementary video 4. (E and F) Representative images of HeLa cells in early prophase, fixed and stained with anti-Eg5 antibodies, anti-α –tubulin and DAPI.
Fig. 4. (A) Schematic of dual-pathway mitosis. The onset of Eg5 loading marks the opening, and NEBD the closure, of a ~9 minute time window. Prophase pathway cells succeed in completing centrosome separation within this window; prometaphase pathway cells do not. Prophase pathway cells achieve bipolar spindle formation 3-4 minutes faster than prometaphase pathway cells because they avoid the need to resolve a transient monopolar state (brackets). Prophase pathway cells make at least 3-fold fewer segregation errors than prometaphase pathway cells (B) Cumulative completion of centrosome separation. Open symbols: prophase pathway cells. Centrosome separation completes exponentially, with a rate constant of 0.30 ± 0.02 min⁻¹. Filled symbols: all cells. Centrosome separation completes exponentially with a rate constant of 0.16 ± 0.01 min⁻¹. Data at later time points were calculated by extrapolating a time for completion of centrosome separation based on the measured velocity of centrosome separation prior to NEBD (see Methods). A longer time window would allow more cells to complete prophase centrosome separation, and further reduce segregation errors, but would delay mitosis. A shorter window would drive all cells through the prometaphase pathway, producing more errors. (C) Successive frames from live-cell movies of HeLa cells expressing Histone2B–EGFP/mRFP-α-tubulin that follow either the prophase pathway or prometaphase pathway. Schematic in first column indicates the position of centrosomes at the time point before nuclear envelope breakdown (~3 min). Yellow arrows indicate a lagging chromosome in a prometaphase pathway cell (bottom two rows). Scale bar is 10 μm.
cells would succeed in separating their centrosomes in prophase, and the error rate would increase. Were more time to be allocated to prophase centrosome separation, the error rate would decrease, but the overall time spent in mitosis would be increased. Our data reveal dual-pathway mitosis as a substantive biological strategy that improves both the speed and fidelity of mitosis, thereby reducing the risk of cancer and other genetic diseases.

Materials and Methods

Human HeLa cells were grown as described previously (McAinsh et al., 2006). To generate stable cell lines a pIRESpuro2 vector (Clontech Laboratories, Inc) containing mCherry-a-tubulin was constructed (pMC206) and transfected into HeLa cells. Stable clones were selected with 0.4 g/ml puromycin. To generate mEGFP-tagged Eg5, a PCR fragment of full-length human Eg5 was ligated into pcDNA5/FRT in frame with a carboxy-terminal FLAG-mEGFP epitope tag, under the control of a low-expressing SV40 promotor (pMC207). For live-cell imaging experiments, the mCherry-a-tubulin stable cell line was transiently transfected with this vector and imaged after 24 hours. Live-cell imaging was performed in chambers (Lab-Tek II; Thermo Fisher Scientific) with Leibovitz L-15 medium containing mCherry-a-tubulin. We also thank Catarina Samora and Shona Moore for experiments measuring the error rate in the prophase/prometaphase pathways. We thank Frank Kozielski for the kind gift of a human Eg5 cDNA that improves both the speed and fidelity of mitosis, thereby revealing dual-pathway mitosis as a substantive biological strategy.

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Competing Interests

The authors declare no competing interests.

References

Fig. S1. Transiently expressed Eg5-mEGFP does not change the timing of bipolar spindle formation or anaphase onset in a stable line of HeLa cells that express mCherry-α-tubulin. The cell line is transfected with control vector (A) or Eg5-mEGFP expressing vector (B). Images were acquired every 3 min. The averaged times of bipolar spindle formation are 5.5±2.5 min (n=83) and 5.0±2.5 min (n=57), respectively. The averaged times of anaphase onset are 37.9±14.6 min (n=60) and 34.4±17.6 min (n=48), respectively. Data values are mean ± SD.

Fig. S2. Eg5-mEGFP loading onto centrosomes is controlled by a roscovitine-sensitive mitotic kinase. (A) 100 μM roscovitine unloads Eg5-mEGFP from centrosomes to which Eg5 has already loaded. The mCherry-α-tubulin signal also reduces. Following roscovitine washout at 47 min, the Eg5-mEGFP signal and the mCherry-α-tubulin signal re-accumulate on centrosomes. (B & C) Quantitative analysis of Eg5EGFP (B) and mCherry-α-tubulin (C) signal. Each data series represents a single centrosome.
Video 1. Mitosis of a HeLa mCherry-z-tubulin cell transiently expressing human Eg5-mEGFP. Eg5 (green) shows the expected localization to microtubules (red) throughout. This cell uses the prophase pathway. Scale bar is 10 μm. Images were recorded every 3 mins.

Video 2. Eg5 loading to the centrosomes indexes the mitotic clock. The cell shown follows the prometaphase pathway. Still images are shown in Figure 1A. Scale bar is 10 μm. Images were recorded every 2 mins.

Video 3. Eg5 loading to centrosomes in the presence of 1 μM Eg5 inhibitor (EI III). Still images are shown in Figure 1D. Scale bar is 10 μm. Images were recorded every 2 mins.

Video 4. High spatiotemporal resolution live-cell imaging of prophase centrosome separation (Figure 3A–D). The cell shown follows the prophase pathway. Scale bar is 5 μm. Images were recorded every 15 secs.