Cell cycle–regulated cortical dynein/dynactin promotes symmetric cell division by differential pole motion in anaphase

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ABSTRACT In cultured mammalian cells, how dynein/dynactin contributes to spindle positioning is poorly understood. To assess the role of cortical dynein/dynactin in this process, we generated mammalian cell lines expressing localization and affinity purification (LAP)–tagged dynein/dynactin subunits from bacterial artificial chromosomes and observed asymmetric cortical localization of dynein and dynactin during mitosis. In cells with asymmetrically positioned spindles, dynein and dynactin were both enriched at the cortex distal to the spindle. NuMA, an upstream targeting factor, localized asymmetrically along the cell cortex in a manner similar to dynein and dynactin. During spindle motion toward the distal cortex, dynein and dynactin were locally diminished and subsequently enriched at the new distal cortex. At anaphase onset, we observed a transient increase in cortical dynein, followed by a reduction in telophase. Spindle motion frequently resulted in cells entering anaphase with an asymmetrically positioned spindle. These cells gave rise to symmetric daughter cells by dynein-dependent differential spindle pole motion in anaphase. Our results demonstrate that cortical dynein and dynactin dynamically associate with the cell cortex in a cell cycle–regulated manner and are required to correct spindle mispositioning in LLC-Pk1 epithelial cells.

INTRODUCTION

The position of the mitotic spindle dictates the location of cytokinesis in diverse cells, thus regulating the size of the two daughter cells (Rappaport, 1996). Although most somatic cells divide symmetrically to generate two equivalent daughter cells, asymmetric cell division is an important process for cell fate determination and morphogenesis during development (Gillies and Cabernard, 2011; Morin and Bellaiche, 2011; Siller and Doe, 2009). In both symmetric and asymmetric divisions, spindle positioning is accomplished by the interactions of dynamic astral microtubules with cytoplasmic or cortical force generators (FGs; Carminati and Stearns, 1997; Dujardín and Vallee, 2002; McCarthy and Goldstein, 2006; Markus et al., 2009; Markus and Lee, 2011; Wuhr et al., 2010; Kimura and Kimura, 2011). Intrinsic signals, such as cortical marks, or extrinsic cues, such as signaling from neighboring cells, influence these interactions by regulating the recruitment or activation of FGs in a spatial and temporal manner, thereby controlling spindle placement in the dividing cell (Morrison and Kimble, 2006; Knoblich, 2010).

A particularly well-studied example of asymmetric spindle positioning is the first mitotic division in the Caenorhabditis elegans embryo, in which the spindle moves toward, and is maintained at, the posterior cortex, giving rise to a larger anterior and a smaller posterior cell (Cowan and Hyman, 2004; Nguyen-Ngoc et al., 2007). Laser ablation experiments show that FGs are present along the cortex of the embryo, with higher FG activity at the posterior cortex resulting in a larger net pulling force acting on the posterior spindle pole (Grill et al., 2001, 2003; Labbe et al., 2004; Krueger et al., 2010). FGs at the posterior cortex also contribute to the characteristic spindle rocking and flattening of the posterior pole observed during movements of the pole toward the posterior cortex (Schneider and Bowerman, 2003). The minus end-directed microtubule motor...
cytoplasmic dynein has been a prime candidate for the cortical FG. However, the mechanism by which dynein generates asymmetric force during metaphase/anaphase is unclear, as components of the dynein complex are not enriched at the posterior cortex of the C. elegans embryo (Couwenbergs et al., 2007; Nguyen-Ngoc et al., 2007). Similarly, during asymmetric divisions of Drosophila neuroblast cells, dynein is believed to exert force to orient the mitotic spindle along the apical–basal cell axis, but the dynein complex, and its regulator dynactin, are not enriched at the neuroblast apical cortex (Siller et al., 2005; Siller and Doe, 2008).

Emerging data suggest that a highly conserved pathway regulates recruitment of dynein and dynactin to the cell cortex for asymmetric spindle positioning. This pathway consists of the noncanonical Gαi protein, which interacts with LGN (Du and Macara, 2004). LGN can simultaneously interact with Gαi at the membrane and with NuMA, a microtubule-associated protein originally identified for its contribution to spindle pole formation (Merdes et al., 1996). NuMA can bind microtubules and has been shown to form a complex with dynein and dynactin (Merdes et al., 1996). Because NuMA cannot bind LGN and microtubules simultaneously due to overlapping binding sites (Du et al., 2002), a favored hypothesis is that NuMA tethers the spindle by transiently associating with astral microtubules and by promoting dynein and dynactin recruitment to the cortical sites. In addition, overexpression of either Gαi or LGN in mammalian cells results in spindle rocking (Du and Macara, 2004), reminiscent of spindle behavior in C. elegans.

In contrast to the well-studied asymmetric cell divisions that occur in worms and flies, most mammalian cells in culture divide symmetrically: the mitotic spindle is positioned at or near the geometric center of the cell, parallel to the substrate (O’Connell and Wang, 2000). Micromanipulation experiments show that normal rat kidney (NRK) cells contain a sensory mechanism that continuously monitors spindle position and maintains spindle orientation with respect to the long axis of the cell (O’Connell and Wang, 2000). Dynein/dynactin appears to be required for this surveillance mechanism and has been localized by immunofluorescence to the cell cortex and the plus ends of astral microtubules (Busson et al., 1998; Faulkner et al., 2000; O’Connell and Wang, 2000; Kobayashi and Murayama, 2009).

Accordingly, treatments that alter astral microtubule dynamics, misregulate dynein, or reduce end-on contacts of astral microtubules with the cortex have been shown to result in spindles that are no longer positioned at the cell center (Faulkner et al., 2000; O’Connell and Wang, 2000; Rankin and Wordeman, 2010; Bruning-Richardson et al., 2011; Samora et al., 2011).

Our previous work in LLC-Pk1 cells showed that microtubule fragments released from the spindle pole during anaphase are transported along the cell cortex, with the plus ends leading; in metaphase cells, however, release and transport of microtubules was rarely observed (Rusan and Wadsworth, 2005). These data suggest the existence of cortically localized, minus end–directed motor proteins and further demonstrate that the localization and/or activity of these motors are regulated by the cell cycle. In the study described here, we found that dynein, dynactin, and the upstream targeting factor NuMA are asymmetrically localized to the cell cortex of mitotic cells, consistent with recent work in HeLa cells (Kiyomitsu and Cheeseman, 2012). We provide evidence that cortical dynein and dynactin are regulated by astral microtubules, polo kinase (Plk1) activity, and the cell cycle. Spindles in LLC-Pk1 cells frequently move along the long axis of the cell, and many cells enter anaphase with an asymmetrically positioned spindle. Of interest, cells with asymmetrically positioned spindles generate symmetric daughter cells via a dynein-dependent differential spindle pole motion during anaphase.

RESULTS

Asymmetric distribution of dynein and dynactin at the cortex in mitotic LLC-Pk1 cells

To gain insight into how dynein and dynactin contribute to spindle positioning in mammalian cells, we generated clonal cell lines expressing localization and affinity purification (LAP)–tagged proteins from bacterial artificial chromosomes (BACs; Cheeseman and Desai, 2005; Poser et al., 2008). With this approach the native regulatory elements are present, and the gene of interest is expressed from the endogenous promoter. For the experiments reported here we used two cell lines, one expressing LAP-tagged dynein heavy chain (DHC) and the other expressing LAP-tagged p50 subunit of dynactin; we refer to the tagged DHC and p50 as dynein and dynactin, respectively. Using the S-tag (in LAP) to retrieve proteins from mitotic cell lysates, we confirmed that dynein and dynactin subunits expressed from BACs were present in native complexes that could interact (Supplemental Figure S1, A–C), consistent with prior results (Poser et al., 2008). Spindle length and mitotic index were similar to those of parental cells (Supplemental Figure S1, D and E), and no abnormalities in mitosis were detected (unpublished data). In addition, because overexpression of p50 is known to disrupt dynactin integrity (Echeverri et al., 1996; Burkhardt et al., 1997; LaMonte et al., 2002), we selected a cell line that exhibited a lower expression level than the endogenous p50 for the following studies (Supplemental Figure S1F).

In mitotic LLC-Pk1 epithelial cells, dynein and dynactin were observed at centrosomes, along spindle microtubules, at kinetochores, and at the cell cortex, consistent with previous reports and supporting the functionality of the BAC approach used here (Figure 1, A and B; Busson et al., 1998; Faulkner et al., 2000; Kobayashi and Murayama, 2009; Ligon et al., 2001; Woodard et al., 2010; Kiyomitsu and Cheeseman, 2012). By total internal reflection fluorescence microscopy, we observed DHC and p50 at the microtubule plus ends with rates consistent with tip tracking (Supplemental Figure S1G). We did not detect any differences in the distribution of dynein and dynactin throughout mitosis (Figure 1, C and D).

We next focused on cortical dynein and dynactin and their role in spindle positioning. In prometaphase and metaphase cells with centrally positioned spindles (symmetrically near the cell center), dynein and dynactin were either absent from the cortex (Figure 1, C and D) or were observed at each end of the cell in a crescent-shaped distribution (Figure 1A, top, C and D, and Supplemental Movies S1 and S2). Cortical dynein and dynactin were not detected at the equatorial region. In cells with asymmetrically positioned spindles, bright patches of dynein and dynactin were observed at the cortex furthest from the spindle (Figure 1, A and B, bottom rows, and Supplemental Movies S3–S5). Time-lapse imaging showed that the distribution of cortical dynein and dynactin changed as the metaphase spindle moved within the cell, as reported previously (Kiyomitsu and Cheeseman, 2012). In the example shown in Figure 1B (top), cortical patches of dynactin were detected at regions distal to the spindle at 0 min. Their fluorescence diminished as the spindle moved toward them at 21 min; new patches subsequently appeared at the new distal cortex after the spindle movement (red arrowheads at 21 min). These results demonstrate that during metaphase the level of cortical dynein and dynactin changes, depending on the proximity of the spindle.

At anaphase onset, we noted that the intensity of cortical dynein and dynactin increased and subsequently decreased as cells progressed into cytokinesis (Supplemental Movies S1–S5). At anaphase, cortical dynein and dynactin appeared relatively smooth and uniform along the polar cortex and, similar to metaphase, were not
FIGURE 1: Dynamics of dynein and dynactin at the mitotic cortex of LLC-Pk1 cells. Time-lapse images of mitotic LLC-Pk1 cells expressing (A) DHC-LAP and (B) p50-LAP. (A) Top, a cell with centrally positioned spindle. Bottom, a cell with displaced mitotic spindle. (B) Top, a metaphase spindle becoming displaced before anaphase onset. Bottom, a cell with displaced mitotic spindle. Arrowheads, DHC-LAP signal at the proximal cortex (A, bottom) or dynamic p50-LAP signal (B, top). Arrows, enriched and uniform DHC-LAP and p50-LAP signal. Scale bars, 10 μm. (C, D) Quantitation of cortical dynein and dynactin localization throughout mitosis. (C) Percentage of cells exhibiting cortical dynactin and dynactin localization. (D) Percentage of cells with indicated localization of cortical dynactin and dynein localization. Error bars in C and D represent SE of proportion; n = 167 (dynein) and 194 (dynactin) cells.
detected at the equatorial region. For centrally located spindles, the observed increase in intensity occurred either at the onset of anaphase or as anaphase progressed (Figure 1A, top; compare 6 with 15 or 17 min, red arrows). For asymmetrically positioned spindles, the increase was more pronounced at the distal cortex (Figure 1, A and B, bottom, red arrows) but could sometimes be detected at the proximal cortex (Figure 1A, bottom, 12 min, arrowhead). The intensity of dynein and dynactin at the centrosome of neighboring interphase cells was unchanged throughout the image sequence (Supplemental Figure S2), indicating that the reduction in dynein or dynactin at the cortex in telophase was not due to photobleaching. These observations demonstrate that 1) cortical dynein and dynactin dynamically associate with the cell cortex, 2) during early mitosis, cortical dynein and dynactin are enriched in regions distal to the spindle and are diminished in regions near the spindle, and 3) the levels of cortical dynein and dynactin increase at anaphase onset and diminish as cells progress into cytokinesis.

**LLC-Pk1 spindle movement parallel to the plane of the cell sheet**

The observation that dynein and dynactin accumulate at the cortex distal to the spindle is counterintuitive: the prevailing view holds that dynein is required for spindle positioning, so one would expect a high concentration of dynein at the cortex proximal to the spindle. This observation led us to hypothesize that dynein/dynactin might be involved in a spindle position correction mechanism and that the levels of dynein/dynactin are reduced after interactions with astral microtubules.

To test this model, we performed time-lapse, phase contrast imaging of LLC-Pk1 cells from spindle assembly to cytokinesis (Figure 2A). The results showed that in 29% of metaphase cells, the spindle assembled at or near the geometric center of the cell (determined as the midpoint of the long axis of the cell) and remained in this location throughout mitosis. We call these cells class I. In 13% of imaged cells, spindle rotation was observed, resulting in the alignment of the spindle along the long axis of the cell, as previously observed for NRK cells (O’Connell and Wang, 2000). We call these cells class II. Strikingly, in 59% of recorded cells, the metaphase spindle was observed to move along the long axis of the cell and, in some cases, translocate from one end of the cell to the other. We call these class III. In these cells, the spindle was often located at one end of the cell at the onset of anaphase. The degree of spindle displacement at anaphase onset, measured as the distance of the spindle equator from the cell midplane (Figure 2B), was 0.27 ± 0.02 of the half-cell length. The extent of spindle displacement correlated with cell length (Figure 2C), indicating that longer cells have a greater tendency to enter anaphase with a mispositioned spindle. Nonetheless, we did not detect a delay in anaphase onset for class III LLC-Pk1 cells (unpublished data), suggesting that mitotic progression was not affected. Of interest, despite having a displaced spindle during metaphase, most class III cells proceeded through anaphase and cytokinesis to produce two daughter cells of equal size (the...
dle elongation, the spindle midzone and cleavage furrow formed anaphase B is responsible for correcting the spindle displacement. Associated with the leading and lagging poles were equivalent toward the opposite cell end. The lengths of the kinetochore fibers remained relatively stationary, and the leading pole moved to the movement of one pole. The lagging pole (top pole in Figure 3C) remained relatively stationary, and the leading pole moved to anaphase A (chromosome-to-cytokinesis) or anaphase B (pole–pole separation) motion, we examined both sets of chromosomes resulted in the correction of the spindle position without an apparent increase in the duration of anaphase.

**Differential pole movements during anaphase**

Recent work showed that anaphase spindle elongation can compensate for spindle displacement in mammalian cells (Xiao et al., 2012). To determine how asymmetrically positioned metaphase spindles could result in symmetric cell divisions in LLC-Pk1 cells, we analyzed the movements of chromosomes and spindle poles during anaphase in class III cells (Figure 3A). We found that during anaphase, the two sets of chromosomes showed differential movement toward the opposite cell ends. Kymographs revealed that the leading set of chromosomes, defined as the chromosomes farthest from the cell cortex, moved a greater distance and at a twofold higher velocity than the lagging set of chromosomes (Figure 3B; 3.36 ± 0.36 vs. 1.56 ± 0.24 μm/min). Kymographs of class I cells showed that both sets of chromosomes moved the same distance at a similar rate (Supplemental Figure S3). The increased velocity of the leading set of chromosomes resulted in the correction of the spindle position without an apparent increase in the duration of anaphase (anaphase onset to furrow initiation was 11.5 ± 0.6 min for class III cells and 10.6 ± 0.6 min for class I cells).

To determine whether the differential movement of chromosomes was contributed by differential anaphase A (chromosome-to-pole) or anaphase B (pole–pole separation) motion, we examined class III cells expressing green fluorescent protein (GFP)–α-tubulin. Time-lapse images (Figure 3C) showed that, upon anaphase onset in a class III cell, the spindle elongated primarily due to the movement of one pole. The lagging pole (top pole in Figure 3C) remained relatively stationary, and the leading pole moved toward the opposite cell end. The lengths of the kinetochore fibers associated with the leading and lagging poles were equivalent throughout anaphase A, demonstrating that spindle pole motion in anaphase B is responsible for correcting the spindle displacement.

As a result of asymmetric spindle pole movements during spindle elongation, the spindle midzone and cleavage furrow formed near the geometric center of the cell. We found that the average displacement of the cleavage furrow from the cell center was significantly reduced from the average displacement of the spindle equator at metaphase (from 0.27 ± 0.02 to 0.04 ± 0.01; Figure 2B). These data show that differential anaphase spindle pole motions result in symmetric divisions in class III LLC-Pk1 cells that had asymmetrically positioned spindles at anaphase onset.

**Dynein activity is required for differential pole movement**

To determine whether dynein activity contributes to differential pole movement in class III cells, we microinjected cells with the CC1 fragment of the p150 subunit of dynactin, an approach used effectively to interfere with the interaction between dynein and dynactin and to inhibit dynein-dependent processes in mammalian cells (Quintyne et al., 1999; Gaetz and Kapoor, 2004; Ferenz and Wadsworth, 2007; Ma et al., 2010). Cells with displaced spindles were microinjected with CC1 immediately after anaphase onset. The results showed that the differential movement of the leading pole was inhibited after microinjection of CC1 (Figure 4A, top), and the cleavage furrow was subsequently initiated away from the cell midpoint (Figure 4B). Control cells microinjected with buffer did not show displacement of the cleavage furrow (Figure 4A, bottom, and B). In addition, we observed that the spindle became disordered after microinjection of CC1, consistent with inhibition of dynein function (Gaetz and Kapoor, 2004). Similarly, we found that treatment of class III cells at the onset of anaphase with a low dose of nocodazole, 12.5 nM, which inhibits microtubule dynamics (Vasquez et al., 1997), prevented the differential movement of the leading pole (Figure 4, C and D). These results demonstrate that dynein function and dynamic microtubules are required for differential pole motion during anaphase B.

**Asymmetric cortical distribution of NuMA**

We next examined the localization of NuMA, a member of the tripartite complex NuMA-LGN-γt3, which is required for the cortical association of dynein (Du and Macara, 2004; Kiyomitsu and Cheeseman, 2012). LLC-Pk1 cells were double stained with antibodies against dynactin p150 and NuMA to compare the distributions of the two proteins at the cortex of mitotic cells (Figure 5A). We found that the distribution of NuMA was strikingly similar to that of p150, with both proteins either absent from the cortex or showing the same distribution (Figure 5B). For example, in metaphase cells with a centrally located spindle, NuMA and p150 were co-localized at both (57.6%) or one (36.4%) end in the majority of cells (Figure 5B, bottom). In metaphase cells with a displaced spindle, both proteins were localized at one end in >90% of cells (Figure 5B, bottom). A small percentage of cells (<16% in all cases) exhibited different cortical distributions of the two proteins (Figure 5B, top, blue bars); of these, 85.7% showed that NuMA was present but p150 was absent, consistent with the role of NuMA as an upstream targeting factor of dynein/dynactin. The observed asymmetric distribution of cortical NuMA is distinct from the symmetric distribution observed in HeLa cells as previously reported (Kiyomitsu and Cheeseman, 2012; see Discussion).

![Figure 3: Asymmetric anaphase chromosome and pole movements result in symmetric division.](image-url)
Astral microtubules regulate accumulation of cortical dynein and dynactin

From our live-cell imaging, we observed that cortical enrichment of dynein and dynactin in metaphase cells was most pronounced in areas of the cortex that were furthest from the spindle, leading us to hypothesize that astral microtubules regulate the distribution of cortical dynein/dynactin. To test this hypothesis, we examined the distribution of microtubules and dynactin in fixed metaphase cells. We found that in cells with a displaced spindle, many astral microtubules reached the proximal cortex but few extended all the way to the distal cortex (Figure 6A, top). Similarly, we found that in metaphase cells that exhibited an irregular cell shape, cortical dynactin frequently accumulated as patches along membrane protrusions that were not directly accessible to astral microtubules (Figure 6A, middle). Conversely, among all of the cells that did not exhibit any detectable cortical dynactin staining, the spindle was generally positioned at the cell center, and astral microtubules extended to all regions of the cortex (unpublished data).

In anaphase cells, we noted that astral microtubules appeared longer than those in metaphase cells. We quantified the average

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**FIGURE 5:** Codistribution of cortical NuMA and the p150 subunit of dynactin. (A) Immunofluorescence images of LLC-Pk1 cells stained with anti-p150 (left; green in merge) and anti-NuMA (middle; red in merge) antibodies; merged images to the right. Arrowheads, codistribution of NuMA and p150 at the cortex. Scale bar, 10 μm. (B) Top, quantitation of cortical NuMA and p150 distribution (n = 231 cells). Error bars represent SE of proportion. Bottom, patterns of NuMA and p150 codistribution observed in cells scored as same (red bars in B; n = 121 cells).
astral microtubule length by measuring the distance from the cen
tosome to the astral microtubule plus end and found that the aver-
age length increased significantly as cells progressed from meta-
phase to anaphase (Figure 6B). Significantly, in anaphase, we noted 
that many cells retained cortical dynactin, despite the presence of 
longer microtubules (Figure 6A, bottom). In summary, we found that 
in metaphase, dynein and dynactin are enriched in regions with few, 
and diminished in regions with many, astral microtubules; whereas in 
anaphase the effects of astral microtubules on cortical dynein/dynac-
tin appear to be dampened.

To test our hypothesis that astral microtubules regulate cortical 
dynein in metaphase, we treated cells with nocodazole at concen-
trations that disassemble microtubules (Figure 6C; Jordan et al., 
1992) and monitored the distribution of cortical dynein. In five of 
eight nocodazole-treated metaphase cells, dynein was observed to 
appear at cortical regions previously devoid of detectable dynein 
(Figure 6C, red arrows). In addition, dynein and dynactin accumu-
lated at kinetochores in nocodazole-treated cells. These results indi-
cate that astral microtubules negatively regulate cortical dynein and 
dynactin.

**Plk1 regulates accumulation of cortical dynein and dynactin**

Plk1 has recently been shown to regulate the association of dynein 
with the cell cortex in HeLa cells (Kiyomitsu and Cheeseman, 2012).

To examine the potential contribution of Plk1 to the cortical dis-
tribution of dynein and dynactin in LLC-Pk1 cells, we arrested cells in 
metaphase with MG132 and inhibited Plk1 activity with BI2536 
(Figure 7, A and B). Addition of BI2536 resulted in an enhancement 
of cortical dynein fluorescence in 62.5% of cells (n = 24 cells). Most 
cells showed an increase in the fluorescence of existing cortical 
patches, as well as the appearance of dynein at cortical sites previ-
ously lacking detectable dynein (Figure 7A, arrowheads and arrows, respectively). In 
cells with a displaced spindle, the majority showed an increase in fluorescence of exist-
ing cortical dynein at the distal cortex; only a minority of cells showed new accumula-
tions at the distal cortex (Figure 7B, Displaced (Distal); compare yellow and blue bars). In contrast, at the proximal cortex, we observed new accumulations of dynein, as 
well as increase in the fluorescence of existing 
cortical patches (Figure 7B, Displaced (Proximal)).

**DISCUSSION**

**Cortical dynein and dynactin are dynamic and regulated 
by the cell cycle**

Determining the dynamic localization of dy-
nein and dynactin complexes in mammalian 
cells has been challenging due in part to the 
lack of suitable probes. Previous work using 
fixed cells demonstrated that dynein and 
dynactin localize to the mitotic cell cortex, 
appearing as a continuous cortical belt or in 
a patchy distribution (Busson et al., 1998; 
Faulkner et al., 2000). In live cells expressing 
GFP-tagged dynein IC, cortical dynein was 
detected, but its dynamics was not de-
scribed (Kobayashi and Murayama, 2009). 
Recently, based on the use of LAP-tagged 
subunits of dynein and dynactin expressed from BACs, a dynamic 
cortical localization of dynein was reported (Kiyomitsu and Cheese-
man, 2012). Consistent with this finding, we report dynamic cortical 
dynein and dynactin in LLC-Pk1 cells. We did not detect a continu-
ous belt of dynein or dynactin as described. This discrepancy might 
result because the signal from the GFP tag on dynein or dynactin 
might be too weak to detect a low concentration of these com-
plexes at the equatorial cortex. Alternatively, different cell types 
might have distinct distributions of dynein and dynactin.

The results of our experiments demonstrate that both dynein 
and dynactin dynamically associate with the mitotic cell cortex in 
mammalian cells. Using live-cell imaging, we show that in cells 
with a centrally located spindle, both complexes are detected at 
cortical regions distal to the spindle and are excluded from the 
equatorial region. In cells with asymetrically localized spindles, 
dynein and dynactin were enriched at cortical regions distant 
distant from the spindle. The motion of the spindle within the cell was 
directed toward regions with enriched cortical dynein and dynac-
tin, and both complexes were diminished after repositioning of 
the spindle in early mitosis. Because both dynein and dynactin 
subunits were tagged with GFP, we could not colocalize them in 
the same cell. However, both complexes exhibited a similar patch-
like distribution in early mitosis and a more uniform distribution 
later in mitosis. Based on these observations, our data support 
the view that these complexes colocalize at the mitotic cortex in 
mammalian cells.

Our results demonstrate that the distribution of dynein and dy-
nectin is regulated by the cell cycle. As mitosis progressed, the per-
centage of cells with detectable cortical dynein increased, with a 
peak in anaphase, and decreased as cells proceeded to cytokinesis. 
Enhanced cortical dynein and dynactin in anaphase could result
from increased association of the complexes at the cortex (higher on-rate), decreased dissociation (lower off-rate), or a combination of both (Figure 7C). Although cortical dynein and dynactin increased in anaphase, we noted that dynein at the cortex proximal to the aster was dimmer than that at the far cortex. Our results and those of Kiyomitsu and Cheeseman (2012) show that inhibition of Plk1 resulted in increased cortical dynein. One possibility is that astral microtubules convey an inhibitory spindle pole–derived Plk1 signal to the cortical cortex. The cell cycle regulation of cortical dynein and dynactin could be directly responsible for the inward transport of peripheral microtubules to the spindle region (Rusan et al., 2002), whereas during anaphase, microtubules move outward to repopulate the cell periphery (Rusan and Wadsworth, 2005). The cell cycle–dependent enrichment of cortical dynein and dynactin in anaphase could power the outward motion of microtubules and centrosome fragments in anaphase cells and the motion of spindle poles that we document here.

**NuMA is dynamically localized to the cortex of LLC-Pk1 cells**

Our observations of cortical dynein and dynactin dynamics are consistent with recent work using HeLa cells (Kiyomitsu and Cheeseman, 2012): in both cell types, the levels of cortical dynein and dynactin were reduced at the proximal and increased at the distal cortex; neither protein was detected at the equatorial region. In HeLa cells, the distribution of NuMA and LGN, members of the tripartite complex NuMA-LGN-Gαi, were distinct from that of dynein and dynactin, showing a symmetric distribution (Kiyomitsu and Cheeseman, 2012). In contrast, in LLC-Pk1 cells we found that the distribution of NuMA was asymmetric, suggesting that regulation of cortical dynein targeting in these cells is upstream of NuMA-LGN-Gαi complex. Consistent with this idea, inhibition of Plk1 in LLC-Pk1 cells did not result in a uniform cortical enhancement of dynein as observed in HeLa cells (Kiyomitsu and Cheeseman, 2012): cortical locations where dynein was present showed an enhancement of dynein, whereas regions lacking dynein were less likely to show a similar enhancement. This could result if cortical regions lacking dynein also lacked NuMA-LGN-Gαi, preventing the increase in dynein upon Plk1 inhibition. This finding indicates that the inhibitory effect of Plk1 on the recruitment of dynein to the cortex, as reported (Kiyomitsu and Cheeseman, 2012), acts at the level of the interaction between dynein/dynactin and the NuMA-LGN-Gαi complex (Figure 7C).

**The spindle correction mechanism requires dynein and dynactin**

Recent work has shown that artificial targeting of Gαi to the apical cortex resulted in spindle displacement in three-dimensional cultured Madin–Darby canine kidney (MDCK) cells (Xiao et al., 2012). Anaphase spindle elongation was found to compensate for the spindle displacement, giving rise to equal-sized daughter cells. Our results show that in unperturbed LLC-Pk1 cells, mitotic spindles frequently moved along the long axis of the cell and as a result were frequently mispositioned at anaphase onset. In these cells, we found that differential pole motion during anaphase onset results in the formation of equal-sized daughter cells. Our results further show that dynein and dynactin and dynamic astral microtubules are required to correct spindle displacement. Although previous studies showed that NRK cells delay the onset of anaphase upon externally induced spindle misalignment (O’Connell and Wang, 2000), we did not detect any delay in the onset of anaphase for LLC-Pk1 cells exhibiting a displaced spindle, suggesting that there might be an inherent difference in the surveillance mechanism between the two.
cell lines. Typically, asymmetric cell division results from a displaced spindle (Siller and Doe, 2009; Gillies and Cabernard, 2011; Morin and Bellaiche, 2011), suggesting that distinct cell types might override the correction mechanism so as to generate unequal-sized daughter cells.

The dynamic and cell cycle-regulated distribution of dynein and dynactin that we report here are consistent with the following model for spindle positioning. During early mitosis, astral microtubules interact with dynein and dynactin at the cell cortex; an imbalance of forces on the asters results in motion of the spindle toward cortical regions of higher dynein density (Grill et al., 2001, 2003). Spindles were often observed to reside near the cortex for several minutes, suggesting that residual dynein and dynactin might tether the spindle to the cortex (Laan et al., 2012). With time, cortical dynein was reduced in regions near astral microtubules. This could result if dynein and dynactin are removed from the cortex by walking toward the microtubule minus end. One possibility is that NuMA is removed from the cortex as a cargo of dynein (Merdes et al., 2000). Alternatively, dynein and dynactin could be released from the cortex directly to the cytoplasm (Figure 7C). At cortical regions distant from the aster and pole, dynein and dynactin accumulate; stochastic interactions of astral microtubules with these regions can result in repositioning of the metaphase spindle within the cell, concomitant with removal of dynein, potentially resulting in multiple motions of the spindle within the cell.

We propose that differential spindle pole motion in anaphase results from dynamic astral microtubules extending to the distal polar cortex, where they engage cortical dynein and dynactin. Our model predicts that the change in microtubule dynamics at the metaphase–anaphase transition (Rusan et al., 2001) and the observed enhancement of dynein and dynactin at the anaphase cortex would increase the probability of a dynamic microtubule encountering cortical dynein/dynactin. Furthermore, as previously proposed (O’Connell and Wang, 2000; Tolic-Norrelykke, 2010), longer astral microtubules might associate with a greater number of motors, and thus it is conceivable that astral microtubules extending from the leading pole might experience a higher pulling force that results in moving the displaced spindle. Although we did not detect dynein or dynactin at the dorsal or ventral cortex, recent work demonstrated that motors in the cytoplasm can generate forces and could contribute to spindle positioning (Kimura and Kimura, 2011; Wuhr et al., 2010).

In summary, we provide evidence for dynamic cortical association of dynein and dynactin in mammalian cells and its regulation by Plk1, astral microtubules, and the cell cycle. The asymmetric spindle positioning in LLC-Pk1 cells and its correction by dynein and dynactin provide a new system for analysis of spindle position and symmetric cell division.

**MATERIALS AND METHODS**

**Cell culture**

LLC-Pk1 cells expressing GFP–α-tubulin (LLC-Pk1-α), DHC-LAP, or p50-LAP were cultured in a 1:1 mixture of Opti-MEM and F-10 media supplemented with 7.5% fetal bovine serum and antibiotics. Cells were grown in a 5% CO₂ atmosphere at 37°C. For time-lapse imaging experiments, cells were plated onto glass coverslips 48–72 h before imaging.

**Immunofluorescence**

Cells were rinsed in 37°C phosphate-buffered saline (PBS) and fixed in either −20°C methanol or 3.2% paraformaldehyde, 0.1% glutaraldehyde, and 0.05% Triton in PBS at 37°C for 20 min, followed by postfixation in methanol at −20°C for 6 min, and rehydrated in PBS containing 0.1% Tween-20 and 0.02% sodium azide (PBS-Tw-Az) for 5 min at room temperature. Cells were then blocked with 3% bovine serum albumin and 0.5% Tween in PBS-Tw-Az for 1 h at room temperature. Primary antibody staining was performed with anti-p150 at 1:100 (BD Transduction Laboratories, Lexington, KY), anti-tubulin at 1:100 (Accurate Chemical and Scientific Corporation, Westbury, NY), or anti-NuMA at 1:5000 (a kind gift from Duane Compton, Dartmouth College, Hanover, NH) for 1 h at room temperature. Secondary antibody staining was performed with Cy3-labeled anti-mouse antibodies at 1:400 (Jackson ImmunoResearch Laboratories, West Grove, PA) or fluorescein isothiocyanate–labeled anti-rat antibodies at 1:128 (Sigma-Aldrich, St. Louis, MO) for 8 h at 4°C. Coverslips were mounted in Vectashield (Vector Laboratories, Burlingame, CA) and sealed with nail polish.

**Inhibitors**

For metaphase arrest, LLC-Pk1 cells were incubated for 1 h with 5 μM MG132 before imaging. For Plk1 inhibition, cells were treated with 10 μM BI2536. Nocodazole was used at 12.5 nM to inhibit microtubule dynamics or at 16 or 33 μM to disassemble microtubules.

**BAC cell line generation and characterization**

The DHC and p50 subunits were labeled at the C-terminus with a LAP tag (Poser et al., 2008). The LAP tag comprises an S-peptide tag and a GFP tag at the C-terminus of the target protein (Cheeseman and Desai, 2005). Mouse DHC and p50 BACs were kind gifts from Anthony Hyman (European Molecular Biology Laboratory, Heidelberg, Germany). Bacteria containing the BAC were grown on LB plates conditioned with chloramphenicol. Subsequent cloning steps to insert the LAP tag were performed according to Gene Bridges-Counter Selection BAC Modification Kit, version 3.1 (Gene Bridges, Heidelberg, Germany). Bacteria were prepared for electroporation at 4°C and electroporated at 1.8 kV. The LAP-tagged BAC was purified using Nucleobond BAC 100 Maxi Prep protocol (Clontech, Mountain View, CA).

To create the p50-LAP cell line, we plated LLC-Pk1 cells at a density of 1.0 × 10⁵ and transfected them 48 h later with LAP-tagged BAC using an Amaza Nucleofector according to the protocol recommended by the manufacturer (Lonza, Walkersville, MD). To create the DHC-LAP cell line, we plated LLC-Pk1 cells to a density of 2.0 × 10⁵ and transfected them 24 h later using Effectene Transfection Reagent according to the protocol recommended by the manufacturer (Qiagen, Valencia, CA). Cells were selected in 400 μg/ml G418 for 2 wk. Cell lines stably expressing LAP-tagged p50 and DHC were prepared using established protocols (Wadsworth et al., 2005).

**Preparation of whole-cell extracts and pull down with S-agarose beads**

LLC-Pk1 cells expressing p50-LAP and DHC-LAP were treated with 100 nM nocodazole 20 h before preparation of cell extracts. Cells were lysed in 50 mM 4-(2-hydroxyethyl)1-piperazineethanesulfonic acid, 150 mM NaCl, 1% NP-40, and protease inhibitors (0.04 mg/ml aprotinin, 0.02 μg/ml leupeptin, and 0.2 mg/ml Pefabloc). Lysates were incubated on ice for 10 min with agitation every 5 min and spun at 14,100 × g in a microcentrifuge for 30 min at 4°C. S-agarose beads were washed with lysis buffer, and the lysate was added to the washed S-agarose beads. The mixture was incubated for 1 h at 4°C with rotation. After incubation, the beads were spun down and washed with lysis buffer. Beads with bound proteins were resuspended in SDS sample buffer and boiled for 3 min before SDS–PAGE.
Western blotting and detection

Protein samples were run on a 10% polyacrylamide gel and transferred onto Amersham Hybond-P membrane (GE Healthcare, Waukesha, WI). Blots were probed with mouse anti-p150 antibody (used at 1:1000; BD Transduction Laboratories), mouse 74.1 anti-dynein IC antibody (used at 1:1000; Chemicon, Temecula, CA), or mouse anti-p50 antibody (used at 1:1000; BD Transduction Laboratories) for 1 h at room temperature in 5% nonfat dry milk dissolved in Tris-buffered saline containing 0.2% Tween-20 (TBS-Tween). The blots were then probed with goat anti-mouse horseradish peroxidase-conjugated secondary antibody (1:5000) for 1 h at room temperature in 5% nonfat dry milk dissolved in TBS-Tween and were detected using chemiluminescence.

Microinjection

To inhibit dynein function, we injected the CC1 fragment of p150 (Ferenz and Wadsworth, 2007) at 15 μM in 50 mM potassium glutamate and 1 mM MgCl₂, pH 7.0, containing 1 mg/ml 10-kDa rhodamine-labeled dextran (Molecular Probes, Eugene, OR) in order to track the microinjected cell.

Image acquisition and analysis

We acquired images using spinning disk confocal microscopy. We used two different microscopes: an Eclipse TE300 microscope (Nikon, Melville, NY) equipped with a CSU 10 spinning disk confocal scan head (PerkinElmer, Wellesley, MA) and an Orca-ER cooled charge-coupled device (CCD) camera (Hamamatsu, Bridgewater, NJ), or a Nikon TiE microscope with a CSU-x1 Yokogawa spinning disk confocal scan head (PerkinElmer) and an Andor iXon+ electron-multiplying CCD camera (Andor, Belfast, Northern Ireland). Both microscope systems were controlled by MetaMorph software (Molecular Devices, Sunnyvale, CA). We used a 100×/numerical aperture (NA) 1.4 objective for fluorescence imaging and a 20×/NA 0.5 objective for phase contrast imaging. For live-cell imaging, exposures were adjusted without saturating the camera’s pixels; typical exposures were 50–1000 ms. For total internal reflection fluorescence microscopy we used a customized system exactly as described previously (Gable et al., 2012). For drug treatments, cells in Rose chambers were imaged, and then nocodazole at 16 or 33 μM (to disassemble microtubules), nocodazole at 12.5 nM (to inhibit microtubules dynamics), or B12536 at 5 μM (to inhibit Ptk1 activity) in non-CO₂ tissue culture medium was added to the dish. Then the top of the chamber was replaced, and imaging resumed. Cells were maintained at ∼37°C during the experiment, but the heater was turned off during changing of the medium. We measured the spindle displacement along the long axis of the cell at anaphase onset as [0.5 × cell length − (distance between chromosome bundle and the proximal cortex)]/[0.5 × cell length]. Similarly, the displacement of the cleavage furrow was calculated as [distance between the cleavage furrow and the proximal cortex]/[0.5 × cell length]. Spindle pole velocity was calculated from kymographs generated from time-lapse phase contrast images using chromosome bundles as reference point. Intensity measurements, daughter cell area measurements, and kymograph constructions were performed using ImageJ software (National Institutes of Health, Bethesda, MD). For cells treated with B12536 or nocodazole, increase in preexisting dynein intensity was measured using movie frames from pretreatment and ∼30 min after treatment; an increase in intensity of 10% or greater was scored as positive. New accumulations of dynein at the cortex were visually scored by two independent observers. To determine statistical significance, Student’s t test was performed.

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