Centriole and PCM cooperatively recruit CEP192 to spindle poles to promote bipolar spindle assembly

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The pericentriolar material (PCM) that accumulates around the centriole expands during mitosis and nucleates microtubules. While centrosomes facilitate bipolar spindle formation, the individual functions of the centriole and PCM in mitosis remain elusive. Herein, we show the redundant roles of the centriole and PCM in bipolar spindle formation in human cells. Upon depletion of the PCM scaffold components, pericentrin and CDK5RAP2, centrioles remained able to recruit CEP192 onto their walls, which was sufficient for bipolar spindle formation. In contrast, through centriole removal, we found that pericentrin and CDK5RAP2 recruited CEP192 at the acentriolar spindle pole and facilitated bipolar spindle formation in mitotic cells with one centrosome. Furthermore, the chemical perturbation of polo-like kinase 1, a critical kinase for PCM assembly, efficiently suppressed the proliferation of various cancer cell lines from which centrioles were removed. Overall, these data suggest that the centriole and PCM cooperatively recruit CEP192 to spindle poles and facilitate bipolar spindle formation in human cells.
Introduction

Centrosomes nucleate and anchor microtubules, thereby facilitating efficient spindle formation and chromosome segregation during mitosis (Moritz et al., 1995; Kollman et al., 2011; Woodruff et al., 2017). The microtubule-organizing activity of centrosomes depends on the pericentriolar material (PCM) that surrounds one or two centrioles (Hyman, 2014). Abnormalities in centrosome organization and function lead to chromosomal segregation errors; several mutations in centrosomal proteins have also been implicated in the development of diseases such as cancer (Nigg and Raff, 2009; Gönczy, 2015). In addition, PCM disorganization directly causes chromosome mis-segregation (Watanabe et al., 2019; Cosenza et al., 2017). Therefore, elucidating the function and organization of centrosome in mitosis will contribute to a better understanding of the mechanisms through which centrosomes dictate the spindle structure and support accurate chromosome segregation.

PCM contains a large number of proteins, such as the γ-tubulin ring complex (γ-TuRC), CDK5RAP2, CEP192, and pericentrin. During the G2/M transition, CEP192 recruits Aurora A and polo-like kinase 1 (PLK1) to centrosomes in a pericentrin-dependent manner; subsequently, CEP192 activates these kinases to promote microtubule nucleation and centrosome separation (Joukov et al., 2014). CEP192 also supports the organization of other PCM components for efficient bipolar spindle assembly (Gomez-Ferreria et al., 2007). PLK1 phosphorylates pericentrin to further recruit other PCM components to centrosomes, thereby increasing the microtubule nucleation activity of the centrosome during mitosis (Lee and Rhee, 2011). Microtubule nucleation activity depends
on γ-TuRC (Zheng et al., 1995; Wieczorek et al., 2020; Liu et al., 2020; Consolati et al., 2020; Moritz et al., 1995; Kollman et al., 2011), the activity of which is upregulated by the binding of CDK5RAP2 to γ-TuRC (Choi et al., 2010; Hanafusa et al., 2015). In addition to their functions in microtubule nucleation, previous studies have described pericentrin and CDK5RAP2 regulating spindle pole focusing and spindle orientation through the regulation of motor proteins or other spindle pole proteins (Lee and Rhee, 2010; Chavali et al., 2016; Tungadi et al., 2017; Chen et al., 2014).

During the G2/M phase, PCM expands around the pair of centrioles that form the structural core of the centrosome, and increases its ability to nucleate microtubules. In Drosophila and Caenorhabditis elegans, it has been reported that the centrioles regulate the architecture and dynamics of PCM (Kirkham et al., 2003; Conduit et al., 2010; Erpf et al., 2019; Cabral et al., 2019; Sir et al., 2013; Alvarez-Rodrigo et al., 2019; Conduit et al., 2014). In addition, it has been shown that PCM disorganization causes precocious centriole disengagement during mitosis (Seo et al., 2015; Kim et al., 2015, 2019; Watanabe et al., 2019), which can result in impairment of spindle pole integrity (Watanabe et al., 2019). This cross-reactive interplay between centrioles and PCM complicates the analysis of the individual function of PCM at spindle poles independent from the involvement of centriolar machinery. The centriole-independent functions of PCM have been partially characterized in the acentriolar meiotic spindles of mouse oocytes. During meiotic spindle formation in mice, acentriolar microtubule-organizing centers are formed and merge into two equal spindle poles (Clift and Schuh, 2015; Schuh and Ellenberg, 2007). Conditional knockout of pericentrin induces spindle
instability and severe meiotic errors that lead to pronounced female subfertility in mouse oocytes. These findings suggest that pericentrin assists in organizing functional spindle poles to achieve faithful chromosome segregation (Baumann et al., 2017). However, as the system of meiosis is particularly unique compared with that of mitosis, it is unclear whether acentrosomal spindle formation pathways can be directly compared between oocytes and somatic cells.

To evaluate the distinct functions of PCM in human somatic cells independently of centrioles, it is important to utilize an assay system that enables the analysis of mitotic spindles that lack centrioles. As centriole duplication requires PLK4 (Habedanck et al., 2005; Bettencourt-Dias et al., 2005), its specific inhibitor centrinone can be used to remove centrioles (Wong et al., 2015). Treatment with centrinone leads to progressive loss of centrioles and generates mitotic spindles with one or zero centrosomes. Using this strategy, we have previously shown the critical roles of NuMA in the spindle bipolarization in early mitosis of cells without centrosomes (Chinen et al., 2020). Similarly, by using mitotic cells with one centrosome, Dudka et al. recently reported that centrosomes regulate the length of K-fibers, and thereby alter their dynamics in HURP-dependent manner (Dudka et al., 2019).

In this study, we show the redundant roles of the centriole and PCM in bipolar spindle formation in human cells. When PCM assembly was inhibited by depletion of the PCM scaffold proteins pericentrin and CDK5RAP2, we found that CEP192 remained at the centriole wall, where it presumably promoted bipolar spindle formation. Furthermore, we induced the formation of mitotic spindles with only one centrosome by treating human
cells with centrinone. We found that the one-centrosome cells formed a bipolar spindle that accumulated PCM components, including CEP192, at the acentriolar pole. In such cells, depletion of pericentrin or CDK5RAP2 compromised the formation of the acentriolar pole and significantly prolonged mitotic progression. In contrast, the artificial accumulation of PCM components at the acentriolar pole accelerated the mitotic progression in one-centrosome cells. These results demonstrate that the centriole and PCM cooperatively assemble CEP192 at the spindle poles and facilitate bipolar spindle formation in human cells.
Results

CEP192 at the centriolar wall is sufficient for organizing mitotic spindle poles

To understand the functions of PCM in bipolar spindle formation in human cells, we depleted the main components of PCM, such as CEP192, pericentrin, and CDK5RAP2, and observed mitotic progression in HeLa cells. As previously described, the depletion of CEP192 caused severe defects in bipolar spindle formation and prolonged mitotic duration (Fig. 1A, B). On the other hand, double depletion of pericentrin and CDK5RAP2 or their individual depletion had a limited effect on mitotic duration (Fig. 1A, B, 2C, and S1G). These results suggest that CEP192, but not pericentrin or CDK5RAP2, is critical for mitotic progression. It has been suggested that pericentrin and CDK5RAP2 cooperatively recruit PCM components, including CEP192, at centrosomes (Kim and Rhee, 2014). Therefore, we observed the amount and localization of CEP192 at centrosomes upon depletion of pericentrin and CDK5RAP2. We found that a certain quantity of CEP192 remained at centrosomes in pericentrin/CDK5RAP2 double-depleted cells (Fig. 1C and D). To further understand this mechanism, we used gated stimulated emission depletion (STED) microscopy to analyze the detailed localization pattern of CEP192 at centrosomes in pericentrin/CDK5RAP2 double-depleted cells (Fig. 1E–G). Centrioles were marked by poly-glutamylated centriolar microtubules. In control cells, CEP192 was detectable in the PCM clouds that surrounded mother centrioles (Fig. 1E–G). In contrast, in pericentrin/CDK5RAP2 double-depleted cells, the reduced quantity of CEP192 was detectable only on centriole walls. These results raise the possibility that
CEP192 at the centriolar wall, rather than in the PCM cloud, is crucial for the microtubule-organizing center function of centrosomes.

Cells with one centrosome form a bipolar spindle that accumulates PCM components at the acentriolar pole

To understand the functions of PCM independently of centrioles in human cells, we next induced the formation of mitotic spindles with one or zero centrosomes by treating HeLa cells with the PLK4 inhibitors centrinone or centrinone B (Fig. 2A and B). Centrosomes were marked by polyglutamylated centriolar microtubules or centrin to determine their number. We depleted PCM components CEP192, pericentrin, and CDK5RAP2 in one- or zero-centrosome cells, and observed their mitotic progression using live cell imaging. As described above, the depletion of CEP192, but not pericentrin or CDK5RAP2, prolonged mitosis in cells with two centrosomes (Figs. 2C, S1A–G). On the other hand, interestingly, we found that depletion of pericentrin or CDK5RAP2, as well as CEP192, significantly prolonged mitotic duration in one-centrosome cells (Figs. 2D, S1H). In contrast, we found that depletion of pericentrin, CDK5RAP2, or CEP192 had a limited effect on mitotic progression in zero-centrosome cells (Figs. 2E, S1I). These results suggest that pericentrin and CDK5RAP2 are important for mitotic progression in one-centrosome cells, but not in two- or zero-centrosome cells.

We further analyzed the localization patterns of PCM proteins at spindle poles using immunofluorescence microscopy (Figs. 2F, G, S2A–D). We found that the acentriolar spindle poles of one-centrosome cells incorporate a detectable amount of PCM.
components, such as pericentrin, CDK5RAP2, CEP192, and γ-tubulin (Figs. 2F, G, S2A) but not CEP152 or CPAP (Figs. 2G and S2A). In this study, we termed the acentriolar spindle pole that contains PCM the ‘PCM-pole’. In contrast, most spindle poles of zero-centrosome cells lacked PCM components, as previously described (Figs. 2F, G, S2A) (Chinen et al., 2020). PCM components were consistently detectable at the acentriolar spindle poles in one-centrosome cells of various human cell lines (Fig. S2B). Furthermore, the PCM-pole was similarly observed in one-centrosome cells induced by SAS6-depletion using the auxin-inducible degron (AID) system (Fig. S2C, D)(Yoshiba et al., 2019), suggesting that this phenotype was not a specific result of PLK4 inhibition.

We next examined whether the PCM-pole nucleates microtubules using a microtubule regrowth analysis. For this analysis, we immunostained the microtubule end binding protein 1 (EB1), which marks growing microtubule plus ends. When restarting the microtubule nucleation, the EB1 signals started developing around both centriolar and PCM-poles, with PCM-poles nucleating less microtubules (Fig. S2E). Thus, PCM-poles possess microtubule nucleation activity, although this activity appears slightly lower than that of centriolar poles. Collectively, these results suggest that one-centrosome cells assemble PCM at the acentriolar spindle pole, which harbors microtubule nucleation activity (Fig. 2H).

The PCM-pole is formed by either split of the PCM from the centriolar pole or accumulation of PCM.
To understand the mechanism of PCM recruitment to the acentriolar pole in one-centrosome cells, we used time-lapse fluorescence microscopy to track the dynamics of endogenous pericentrin or CDK5RAP2 tagged with mCherry as markers of PCM. This strategy revealed that, at first, pericentrin accumulated at centriolar poles in early mitosis. Subsequently, one-centrosome cells formed pericentrin-positive PCM-poles by either splitting of the PCM from the centriolar pole or de novo accumulation of PCM (38.5% and 51.9%, respectively) (Fig. 3A, B, D). These PCM-poles disappeared after cytokinesis (Fig. 3A, B, E), consistent with the observation that PCM proteins are disassembled after mitotic exit (Woodruff et al., 2014). On the other hand, a detectable amount of pericentrin did not accumulate at the acentriolar spindle poles in most zero-centrosome cells (Fig. 3C, D). Taken together, these observations suggest that one-centrosome cells initially accumulate PCM proteins around centrioles and subsequently generate the acentriolar pole by splitting and/or by recruiting PCM components on the opposite side for bipolar spindle formation (Fig. 3F).

**CDK5RAP2 and pericentrin are crucial for the bipolar spindle formation in one-centrosome cells**

We next analyzed the specific role of PCM in cell division in one-centrosome cells. However, it is difficult to analyze the specific roles of some PCM-pole components. Among those appeared to localize at PCM-poles (Fig. 2F, G, S2A), for example, γ-tubulin also localizes along the whole spindle and regulates several pathways of microtubule nucleation in mitosis (Lecland and Lüders, 2014; Teixido-Travesa et al., 2012). In
addition, CEP192 is required for bipolar spindle formation in cells with two centrosomes (Zhu et al., 2008; Joukov et al., 2014). On the other hand, depletion of the PCM scaffold proteins CDK5RAP2 and pericentrin are known to have little effect on spindle formation in two- or zero-centrosome cells (Fig. 2C-E, S1G-I). Therefore, we selected CDK5RAP2 and pericentrin for further analysis of PCM-poles in one-centrosome cells.

We found that depletion of CDK5RAP2 or pericentrin caused arrest of one-centrosome cells in mitosis with monopolar spindles; however, this effect was not observed in two-centrosome cells (Fig. 4A, B). These results indicate that CDK5RAP2 and pericentrin play an important role in bipolar spindle formation specifically in one-centrosome cells, but not in two-centrosome cells. To further investigate this process, we tracked the dynamics of spindle poles in one-centrosome cells using time-lapse observation of NuMA tagged with mCherry. Upon depletion of CDK5RAP2 or pericentrin, the separation of two NuMA foci was normally detectable in early mitosis (Fig. 4C, D), while the time from nuclear envelope breakdown to cytokinesis was prolonged (Fig. 4C–F). In addition, immunofluorescence analysis revealed that, in the pericentrin or CDK5RAP2-depleted HeLa cells with one-centrosome, the degree of CEP192 localization at the acentrosomal spindle poles was reduced (Fig. 4G). These results indicate that PCM scaffold proteins CDK5RAP2 and pericentrin are crucial for the recruitment of CEP192 at the acentrosomal spindle pole and bipolar spindle formation, but are likely dispensable for the early step of spindle pole generation in one-centrosome cells.
To verify whether pericentrin and CDK5RAP2 are important for bipolar spindle formation in other human cell lines with one centrosome, we observed the spindle structure of RPE1 and A549 cells upon depletion of pericentrin or CDK5RAP2. Through immunostaining, we found that in cells with one centrosome, depletion of pericentrin or CDK5RAP2 induced the formation of monopolar spindles (Fig. S3A–D). These results further support the conclusion that PCM proteins are required for bipolar spindle formation in one-centrosome cells.

Depletion of CEP57 promotes accumulation of PCM components at PCM-poles and facilitates bipolar spindle formation in one-centrosome cells

Next, we sought to further analyze the importance of PCM components at PCM-poles for cell division in one-centrosome cells. Since siRNA-mediated depletion reduces the total expression level of CDK5RAP2 and pericentrin, it is difficult to analyze the function of the PCM components specifically at PCM-poles (Fig. 4). Therefore, we used another approach to indirectly manipulate the amount of PCM components at PCM-poles: depleting CEP57. CEP57 provides a critical interface between the centriole and PCM, and depletion of CEP57 induces the fragmentation of PCM proteins in early mitosis of human cells (Watanabe et al., 2019). Given that 38.5% of one-centrosome cells assembled PCM-poles by splitting PCM from the centrosome (Fig. 3A, D), we hypothesized that, upon CEP57 depletion, the PCM that is dissociated from the centrosome could be incorporated into the acentriolar pole in one-centrosome cells. As expected, the amount of pericentrin at PCM-poles was significantly increased, presumably due to the increased
PCM fragmentation at centriolar poles after CEP57 depletion (Fig. 5A–C). Subsequently, to analyze the effect of CEP57 depletion on the mitotic processes of one-centrosome cells, we performed time-lapse imaging of NuMA and microtubules. We found that depletion of CEP57 promoted bipolar spindle formation more efficiently than in control cells, and thereby shortened the mitotic duration (Fig. 5D–F). Under this condition, CEP57-depleted cells with one centrosome formed two separate NuMA foci, similar to siControl-treated one-centrosome cells, but established a bipolar spindle formation more efficiently (Fig. 5E, G). Overall, these results suggest that accumulation of PCM components at PCM-poles facilitates the bipolar spindle formation in one-centrosome cells.

Pericentrin is crucial for bipolar spindle elongation in cells with two centrosomes

Although pericentrin and CDK5RAP2 are dispensable for efficient mitotic progression in cells with two centrosomes (Fig. 1A, B, 2C, S1G), the detailed functions of these PCM components in bipolar spindle formation have not been carefully examined. We subsequently analyzed the spindle length upon depletion of pericentrin or CDK5RAP2 in HeLa cells. We found that depletion of pericentrin significantly reduced the spindle length compared with that of control cells, whereas depletion of CDK5RAP2 had a limited effect on the spindle length (Fig. 6A, B). To further investigate this defect upon depletion of pericentrin, we performed live cell imaging of mitotic spindle formation in HeLa and HCT116 cells. Depletion of pericentrin delayed the elongation of two spindle poles (Fig. 6E, F, S4A, B). These results suggest that pericentrin supports spindle elongation. In line with this result, immunofluorescence analysis revealed that, in the
pericentrin-depleted HeLa cells, the degree of CEP192 localization at the spindle poles was reduced; however, this was not observed in CDK5RAP2-depleted cells (Fig. 6C, D). This observation implies that pericentrin more efficiently recruits CEP192 to centrosomes, thereby facilitating spindle elongation.

Furthermore, we tested the effect of depletion of pericentrin on the spindle elongation of various cell types. The spindle length in pericentrin-depleted A549, U2OS, A431, and PANC1 cells was significantly shorter than that noted in control cells (Fig. 6G, S4C, F–H). On the other hand, in some cell types (RPE1, GI1, SKOV3), the spindle length upon depletion of pericentrin was not altered compared with that observed in control cells (Fig. S4C–F). These results suggest that pericentrin is required for efficient spindle elongation in certain cell lines with two centrosomes.

The activity of PLK1 is crucial for PCM-pole assembly and bipolar spindle formation in one-centrosome cells

The accumulation of PCM components at centrosomes in mitosis is regulated by PLK1 activity (Haren et al., 2009; Lee and Rhee, 2011; Joukov et al., 2014). However, we found that PLK1 and phosphorylated PLK1 were not detected at most PCM-poles in one-centrosome cells (Fig. 7A–D). To determine if PLK1 was required for PCM-pole assembly and subsequent bipolar spindle formation, we treated cells with a low dose of the PLK1 inhibitor BI 2536 (1 nM) and observed the amount of pericentrin at the centriolar pole and the spindle structure. Treatment of two-centrosome cells with a low dose of the PLK1 inhibitor caused chromosome congression errors and a slight reduction
of pericentrin at centrosomes, but did not affect bipolar spindle formation (Fig. 7E–G).

In contrast, in one-centrosome cells, PLK1 inhibition prevented PCM-pole formation and led to the formation of monopolar spindles (Fig. 7E, F). In addition, PLK1 inhibition greatly reduced the level of pericentrin at the centriolar pole compared with the level recorded in two-centrosome cells (Fig. 7G). Together, these results suggest that PLK1 activity is crucial for PCM-pole assembly and subsequent bipolar spindle formation in one-centrosome cells.

**Dual inhibition of PLK1 and PLK4 prevents cell growth in a wide variety of cancer cell lines**

Mitotic spindle formation is a common target of anti-cancer drugs (Dumontet and Jordan, 2010; Tischer and Gergely, 2019; Henriques et al., 2019). Since the dual inhibition of PLK1 and PLK4 (PLK1+4i) efficiently prevented bipolar spindle formation in one-centrosome cells (Fig. 7E, F), we further tested the potential of PLK1+4i as an anticancer strategy. PLK1+4i efficiently prevented HeLa cell growth (Fig. 8A). The half maximal inhibitory concentration (IC$_{50}$) value of the PLK1 inhibitor against HeLa cells was 1.1 nM (Table S2). Using both PLK1 and PLK4 inhibitors decreased the IC$_{50}$ value to 0.6 nM (Table S2). Therefore, inhibition of both PLK1 and PLK4 has an additive effect on the growth suppression of HeLa cells. Under these conditions, after depletion of centrosomes, most of HeLa cells started to die due to prolonged mitosis (Fig. 6B, C, S6). The results suggest that the toxicity of PLK1+4i may be caused by inhibition of PCM-pole formation. Furthermore, PLK1+4i-treated cells showed cleavage of the apoptosis
marker poly(ADP-ribose)-polymerase 1 (PARP1), suggesting that this drug combination induces apoptosis in mitosis (Fig. 6D). In addition, we assessed the effect of PLK1+4i in 19 cancer cell lines. The PLK1 inhibitor suppressed the proliferation of cancer cell lines to different extents (IC_{50} values in Table S1). U2OS, K562, and HMV-II cells showed approximately 10-fold higher resistance against the PLK1 inhibitor compared with HeLa cells. However, PLK1+4i efficiently prevented the growth of various cancer cell lines, including PLK1 inhibitor-resistant cell lines such as U2OS and K562 (15 cell lines: IC_{50}[−centrinone B]/IC_{50}[+centrinone B] ≥ 1.5) (Fig. 6E and Table 1). Overall, these results suggest that dual inhibition of PLK1 and PLK4 would be an effective drug target against cancer proliferation.
Discussion

In this study, we show that the centriole and PCM cooperate to recruit CEP192 at the spindle pole to facilitate bipolar spindle formation in human cells. We found that, even in cells in which PCM assembly was suppressed, CEP192 at the centriole wall efficiently promoted bipolar spindle assembly (Fig. 1). Furthermore, cells with one centrosome formed a bipolar spindle with a PCM-pole, which accumulates PCM proteins (including CEP192) at the opposite side of the centriolar spindle pole (Fig. 2–3). Consistently, the PCM-pole assembly is critical for cell division in one-centrosome cells (Fig. 4–5).

Overall, the findings in this study illustrate that the centriole and PCM cooperatively promote bipolar spindle assembly through recruitment of CEP192 to the spindle pole in human somatic cells (Fig. 9). In addition, based on this evidence, we propose that dual inhibition of centriole duplication and a critical mitotic kinase PLK1 would be an attractive target for anti-cancer strategies (Fig. 7–8).

In interphase cells, CEP192 localizes at the centrioles and regulates the microtubule nucleation activity of centrosomes (O’Rourke et al., 2014). In the G2/M phase, CEP192 is further recruited to PCM clouds by pericentrin (Joukov et al., 2014), promoting mitotic spindle formation. In pericentrin/CDK5RAP2 double-depleted cells, although the CEP192 localization was restricted on the centriolar wall, these cells efficiently completed mitosis (Fig. 1C–F). These results suggest that a fraction of CEP192 at the centriolar wall is sufficient for its function in mitosis. A previous study suggested that CEP192 supports the sequential activation of PLK1 and aurora kinase A (AURKA) at centrosomes (Joukov et al., 2014). Moreover, it has been shown that phosphorylated
AURKA interacts with TPX2 and promotes spindle assembly (Joukov and De Nicolo, 2018). It is therefore possible that CEP192 at the centriole wall sufficiently activates the PLK1-AURKA pathway, thereby facilitating bipolar spindle formation.

We found that one-centrosome cells efficiently assembled PCM-poles (Fig. 2F, G, S2A-D). On the other hand, intriguingly, most zero-centrosome cells failed to assemble PCM proteins at the acentriolar poles (Chinen et al., 2020). How does this difference occur? In one-centrosome cells, PLK1 was localized only at centriolar poles, but not at PCM-poles (Fig. 7A–D). However, the PLK1 kinase activity is somehow necessary for the assembly of PCM-poles. It is possible that phosphorylation events at the centriole driven by the activity of PLK1 may provide a pool of PCM for the generation of the PCM-pole. In contrast, zero-centrosome cells do not have the platform components (e.g., centrioles) for PCM assembly. Previous research indicated that, in zero-centrosome cells, the activity of PLK1 in the cytoplasm was significantly increased (Takeda et al., 2020). However, in such cells, the PCM-pole was not assembled at the spindle poles (Takeda et al., 2020). Together, these observations suggest that the centriole itself is important for PCM assembly in human cells.

Knockdown experiments further revealed that CDK5RAP2 and pericentrin are crucial for cell division in one-centrosome cells. In addition, depletion of CEP57 augmented the assembly of PCM-poles, and facilitated mitotic progression in one-centrosome cells. These results indicate that the PCM proteins are required for PCM-pole formation in one-centrosome cells, and also raised the possibility that the balance of PCM quantities between two spindle poles may be a critical factor for proper mitotic...
progression in human cells. In line with this notion, it has been shown that in primary human malignancies, centrosome abnormalities such as centriole rosettes are frequently observed (Cosenza et al., 2017). These extra centrioles could lead to a greater accumulation of PCM proteins at the one centrosome, thereby increasing the nucleation of microtubules at this spindle pole and resulting in chromosome missegregation and aneuploidy. Our assay system may be useful for analyzing the balance of PCM quantities and the resulting microtubule nucleation between two spindle poles.

Based on the vulnerabilities of one-centrosome cells described above, our study also highlights the potential of dual inhibition of centriole duplication and PCM assembly as an attractive drug target for cancer therapies. The PLK1 inhibitor efficiently suppressed both PCM maturation and subsequent PCM-pole formation in one-centrosome cells. In this way, a low dose of PLK1 inhibitor efficiently suppressed cell division in one-centrosome cells, but not in two-centrosome cells (Fig. 8B, C). Previous clinical trials of PLK1 inhibitors have not been successful. Therefore, several studies have been performed to improve PLK1 inhibitor toxicity through combination with other inhibitors, such as α/β-tubulin inhibitors (Stehle et al., 2015; Weiß et al., 2015). In this study, the dual inhibition strategy, which inhibits both PLK1 and PLK4, provided an alternative approach to targeting PLK1 in the development of anticancer drugs. Interestingly, treatment with centrinone did not strongly alter the toxicity of microtubule inhibitors (Table S2). This result suggests that the strong toxicity caused by the dual inhibition of PLK1 and PLK4 was not merely due to an additive effect in mitosis, but rather the specific inhibition of both centrosomal and acentrosomal spindle assembly machinery. In addition,
recently, it was suggested that decreased centrosome numbers are associated with poorer response to chemotherapy and an increased invasive capacity of tumor cells in ovarian cancer (Morretton et al., 2019). Therefore, our strategy to suppress PCM assembly in centrosome-reduced cells may be an attractive method for targeting ovarian cancer cells that have a reduced number of centrosomes.
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Author contributions

T.C. and D.K. designed the study; T.C., K.Y., K.F., K.W., Y.T., and Y.N. performed the experiments; and T.C., K.Y., K.H., S.Y., Y.T., and D.K. designed the experiments. T.C., K.Y., Y.T., and D.T. analyzed the data; T.C., K.Y., and D.K. wrote the manuscript, which was reviewed by all authors.

Competing financial interests

The authors declare no competing financial interests.
Methods

Cell culture and transfection

HeLa and U2OS cells were obtained from the ECACC (European Collection of Authenticated Cell Cultures). These cell lines were authenticated by Short Tandem Repeat (STR) profiling at the ECACC. HeLa cells stably expressing EGFP-centrin1 have been previously described (Tsuchiya et al., 2016). HeLa cells expressing pericentrin endogenously tagged with mCherry were generated using the CRISPR/Cas9 system, as previously described, with slight modifications (Natsume et al., 2016). GuideRNA oligos (Pericentrin_gRNA_F: CACCGCTGTATTAATCGGGTGGC and Pericentrin_gRNA_R: AAACGCCACCCGATGATTAACGAGC) were hybridized and cloned into the BbsI site of pX330 (Addgene). To construct the donor plasmid for homology-directed repair, the homology arms of the Pericentrin locus (chr21:47864730-47865813) were amplified (pBS2_Pericentrin_C-ter_InsF: GGTATCGATAAGCTTACCAGGTATGCAGTCCTCCTGCG and pBS2_Pericentrin_C-ter_InsR: CGCTCTAGAATCTTAGAATGCTCCGGGTCCACTGA) from the genomic DNA of HeLa cells and cloned into pBluescript using the Infusion Cloning kit (Takara). A BamHI sequence with a silent mutation to prevent re-cutting was generated in the middle of the homology arm domain by mutagenesis PCR (Pericentrin C-ter silent BamHI_F: TACTTCAAAAGAATCTTGGCCACCCGATGATTAACAGGGATCCATAAAATG and TCATGGCTCTTCTGCGA, Pericentrin C-ter silent BamHI_R: TACCGCTGTATTAATCGGGTGGC and Pericentrin_gRNA_R: AAACGCCACCCGATGATTAACGAGC).
GCCATGACATTTTATGGATCCCTGTTTAATCATCGGGTGAGG). The mCherry cassette containing a hygromycin-resistant gene was introduced into the BamHI site in the middle of the homology arms. The plasmids were introduced into the HeLa cell line stably expressing EGFP-centrin1 (Tsuchiya et al., 2016) and isolated using the limited dilution method with hygromycin. Saos-2, A549, DAUDI, RAJI, HL60, K562, MKN45, MKN1, DU145, PC-3, PANC-1, GI-1, A431, HMV-II, and MCF-7 were obtained from the RIKEN BioResource Research Center. These cell lines were authenticated by STR profiling at the RIKEN BioResource Research Center. HCT116 cells were obtained from the American Type Culture Collection (ATCC CCL-247). HCT116 CMVOS-TIR1 HsSAS6-AID have been previously described (Yoshiba et al., 2019). HCT116 cell lines were cultured in McCoy’s 5A medium (Thermo Fisher Scientific) supplemented with 10% fetal bovine serum, 2 mM L-glutamine, 100 U/ml penicillin, and 100 μg/ml streptomycin. SKOV-3 was provided Dr. Yoko Nagumo. HeLa, U2OS, A549, GI-1, and A431 cells were cultured in Dulbecco’s modified Eagle’s medium containing 10% fetal bovine serum, 100 U/ml penicillin and 100 μg/ml streptomycin at 37 °C in a 5% CO₂ atmosphere. Saos-2, DAUDI, RAJI, HL60, MKN45, MKN1, DU145, PC-3, PANC-1, and SKOV-3 cells were cultured in RPMI1680 medium containing 10% fetal bovine serum, 100 U/ml penicillin, and 100 μg/ml streptomycin at 37 °C in a 5% CO₂ atmosphere. K562 and HMV-II cells were cultured in Ham’s F12 medium containing 10% fetal bovine serum, 100 U/ml penicillin, and 100 μg/ml streptomycin at 37 °C in a 5% CO₂ atmosphere. MCF-7 cells were cultured in MEM medium containing
10% fetal bovine serum, 1 mM sodium pyruvate, 100 U/ml penicillin, and 100 μg/ml streptomycin at 37 °C in a 5% CO₂ atmosphere. Transfection of siRNA constructs was conducted using Lipofectamine RNAiMAX (Life Technologies). Unless otherwise noted, the transfected cells were analyzed 48 h after transfection with siRNA.

**RNA interference**

The following siRNAs were used: Silencer Select siRNA (Life Technologies) against CEP57 (s18692), CEP192 (s226819), CDK5RAP2 (s31429, s31430), pericentrin (s10136, s10138), and negative control #1 (4390843).

**Chemicals**

The following chemicals were used in this study: Centrinone (PLK4 inhibitor, MedChem Express, HY-18682), Centrinone B (PLK4 inhibitor, gift from Dr. Andrew Shiau and Dr. Karen Oegema), BI2536 (PLK1 inhibitor, A10134; AdooQ), proTAME (APC/C inhibitor, I-440, Boston Biochem), Cytochalasin B (actin inhibitor, Wako, 036-17553), paclitaxel (α/β-tubulin stabilizer: Wako, 163-18614), SiR-Tubulin (Microtubule probe, CY-SC002; SPIROCHROME).

**Antibodies**

The following primary antibodies were used in this study: rabbit polyclonal antibodies
against CDK5RAP2 (IHC00063; immunofluorescence [IF] 1:500, Western Blotting [WB] 1:1000; Bethyl Laboratories), Cep192 (A302–324A, IF 1:1,000, WB 1:1000; Bethyl Laboratories), Cep152 (A302–480A, IF 1:1,000; Bethyl Laboratories), CP110 (12780–1-AP, IF 1:1,000; Proteintech), CPAP/CENP-J (11517-1-AP, IF 1:100; Proteintech), and α-tubulin (PM054, IF 1:300; MBL); mouse monoclonal antibodies against γ-tubulin (GTU88) (T6557, IF 1:1,000; Sigma–Aldrich), Polyglutamylation Modification (GT335, mAb) (AG-20B-0020-C100, IF 1:1,000; AdipoGen), α-tubulin (T5168, IF 1:1,000; Sigma–Aldrich), HSP90 (610419, WB 1:5000; BD Biosciences) pericentrin (ab4448, WB 1:1,000; Abcam) and pericentrin (ab28144, IF 1:1,000; Abcam). Fluorescein isothiocyanate-labeled anti-GFP was purchased from Abcam (ab6662, IF 1:250 or 1:500). Alexa 488-labeled Cep152 (A302–480A, IF 1:250; Bethyl Laboratories) was generated with Alexa Fluor labeling kits (Life Technologies) and used for three color staining. The following secondary antibodies were used: Alexa Fluor 488 goat anti-mouse IgG (H+L) (A-11001, 1:500; Molecular Probes), Alexa Fluor 555 goat anti-rabbit IgG (H+L) (A-21428, 1:500; Molecular Probes), and Alexa Fluor 647 goat anti-mouse IgG (ab150115, 1:500; Abcam).

Sample preparations for immunostaining

Cells were treated with 100 nM of centrinone or 500 nM of centrinone B for 1–3 days to induce acentrosomal cells (Fig. 2A, B, F, G, S2A, B). To observe the microtubule nucleation from PCM-poles (Fig. S2E), firstly, HeLa cells expressing pericentrin-
mCherry and EGFP-centrin1 were treated with 100 nM of centrinone for two days. Then, cells were arrested in metaphase through treatment with 20 μM of proTAME for 4 h and incubated on ice for 1 h to depolymerize microtubules. Subsequently, cells were incubated at 25 °C for 5 min. For the Sas-6 depletion experiments using the AID system (Fig. S2C, D), the cells were incubated with 50 μM of indole-3-acetic acid (IAA) for two days. To observe CEP192 localization at acentriolar poles (Fig. 4G), cells were treated with 100 nM of centrinone and siRNA for two days. Then, cells were arrested in metaphase through treatment with 20 μM of proTAME for 6 h. To examine the effect of siRNA on acentriolar cells (Fig. 4A, B, 5A, B, G, S3), cells were treated with 100 nM of centrinone or 500 nM of centrinone B and siRNA for two days. For the chemical perturbation experiments (Fig. 7E–G, 8D), cells were treated with 100 nM of centrinone and BI 2536 for two days.

### Western blotting

For preparation of total cell lysates, cells were lysed in 1× SDS sample buffer. SDS–PAGE was performed using 6 or 10% polyacrylamide gels, followed by transfer on Immobilon-P membrane (Millipore Corporation). Blocking was performed in 2.5% skim milk in PBS containing 0.02% Tween (PBS-T) for 30 min at room temperature. The membrane was probed with the primary antibodies for 12-18 h at 4°C, washed with PBS-T three times. After that, the membrane was incubated with HRP-conjugated secondary antibodies for 1 h at room temperature and washed with PBS-T three times. The signals
were detected with ECL Prime/Select reagents (GE Healthcare) or Chemi-Lumi One 
Ultra (Nacalai Tesque) via the ChemiDoc XRSp system (Bio-Rad).

Sample preparations for live-cell imaging

For live cell imaging, HeLa cells stably expressing EGFP-centrin1, HeLa cells expressing 
pericentrin-mCherry and EGFP-centrin1, HeLa cells expressing CDK5RAP2-mCherry 
and EGFP-centrin1, HeLa cells expressing mCherry–NuMA and EGFP–centrin1 were 
cultured in 35-mm glass-bottom dishes (#627870; Greiner Bio-One) or 24-well 
SENSOPLATE (#662892; Greiner Bio-One) at 37 °C in a 5% CO₂ atmosphere.

To observe the dynamics of PCM (Fig. 3A–E) in one-centrosome or zero-
centrosome cells, cells were treated with 500 nM of centrinone B. To test the effect of 
depletion of PCM proteins or CEP57 on one-centrosome cells (Fig. 4C–F, 5D–F), cells 
were treated with siRNA with 100 nM of centrinone for two days. To test the effect of 
drug combinations on mitotic cell fate (Fig. 8B, C, S5), the cell cycle progression of HeLa 
cells expressing pericentrin-mCherry and EGFP-centrin1 was observed in the presence 
of 100 nM centrinone and 1 nM BI 2536. To simultaneously observe one-centrosome 
cells and zero-centrosome cells (Fig. S2C-E, S1G-I), after two days of treatment with 
0.1% DMSO or 100 nM centrinone (to enrich zero-centrosome cells), HeLa cells 
expressing EGFP-centrin1 were treated with siRNA with 0.1% DMSO or 100 nM of 
centrinone. Prior to imaging, cells were incubated with 100 nM of SiR-Tubulin for 4 h to 
visualize the microtubules.
Microscopy for immunofluorescence analyses

For immunofluorescence analyses, the cells cultured on coverslips (No. 1; Matsunami) were fixed using methanol at −20 °C for 7 min and washed with PBS. The cells were permeabilized after fixation with PBS/0.05% TritonX-100 (PBSX) for 5 min, and blocked in 1% BSA in PBSX for 30 min at room temperature. The cells were then incubated with primary antibodies for 7–24 h at 4 °C, washed thrice with PBSX, and incubated with secondary antibodies and 0.2 μg ml⁻¹ Hoechst 33258 (DOJINDO) for 45–60 min at room temperature. The cells were washed thrice with PBSX and mounted onto glass slides.

We counted the number of spindle patterns using a DeltaVision Personal DV-SoftWoRx system (Applied Precision) or an Axioplan2 fluorescence microscope (Carl Zeiss). Confocal microscopy images were captured by the Leica TCS SP8 system. For deconvolution for confocal microscopy images, Huygens essential software (Scientific Volume Imaging) was used.

STED images were taken using a Leica TCS SP8 STED 3X system with a Leica HCPL APO 100 × /1.40 oil STED WHITE and a 660 nm laser line for depletion. Scan speed was set to 100 Hz with 5× line averaging in a 512 × 80 px format (pixel size 15–20 nm). The Z interval was set to 180 nm. The STED images were processed by deconvolution using the Huygens Professional software (SVI).
Maximum intensity z-projections of a representative picture for each condition were generated using the FIJI distribution of the ImageJ software. The number and step sizes of z-planes are described in the figure legends.

Microscopy for live imaging

A Confocal Scanner Box, Cell Voyager CV1000 (Yokogawa Electric Corp.) equipped with a 60× oil immersion objective or CQ1 Benchtop High-Content Analysis System equipped with a 40× objective was used for live cell imaging. Imaging was initiated after the addition of drugs or 24–48 h after transfection, and images were acquired every 10 min for 24–48 h. Maximum intensity z-projections of representative images for each condition were generated using the FIJI distribution of the ImageJ software. The number and step sizes of z-planes are described in the figure legends.

Cell viability

Cell viability was determined using the WST-8 assay. Exponentially growing cells (1 × 10^3 cells/well in a 96-well plate) were treated with compounds for three (Fig. 8A, Table S2) or four (Fig. 8E, Table S1) days. WST-8 assay reagents (DOJINDO) were added to the culture. After several hours of incubation, the absorbance at 450 nm was measured with FilterMax F3 & F5 Multi-Mode Microplate Readers (Molecular Devices). After subtracting the background (blank well), cell viability (control %) was determined.
Statistical analysis

Statistical analyses were performed using the GraphPad Prism 7 software. Except for Fig. 6B, p-values were determined by non-parametric methods (Mann–Whitney U-test or Kruskal–Wallis test). P-value in Fig. 6B was determined by One-way ANOVA with Tukey’s multiple comparisons test. Details are described in the figure legends.

Data availability

The data supporting the findings of this study are available from the corresponding authors upon request.
Figure 1. CEP192 at centriolar wall, but not the expanded PCM mediated by pericentrin and CDK5RAP2, is sufficient for bipolar spindle formation

(A) Time-lapse observation of the structure of microtubules upon siRNA treatment against the indicated proteins. HeLa cells expressing EGFP-centrin1 and pericentrin-mCherry were observed with a 40× objective. Gray represent SiR-tubulin, respectively. Z-projections: 10 planes, 2.2 µm apart. Scale bar, 10 µm. Time zero corresponds to nuclear envelope break down (NEBD). (B) Mitotic duration, the time required from NEBD to cytokinesis, in (A). Line and error bars represent the mean and SD (N ≥ 50 cells from two independent experiments). Kruskal–Wallis test was used to determine the significance of the difference. *P < 0.05. (C) The localization of PCM proteins in mitotic spindles of the cells in which the indicated protein was depleted. Red and green represent PCM proteins (CDK5RAP2, Cep192 or PCNT) and GT335, respectively. Z-projections of 10 sections, every 0.3 µm. Scale bar, 1 µm. (D) The signal intensity of PCM proteins on mitotic centrosomes of fixed HeLa cells was analyzed (N>45 for each condition). Line and error bars represent median with interquartile range. Kruskal–Wallis test was used to determine the significance of the difference. *P < 0.05, **P < 0.0001. (E) STED images showing centriolar distribution of Cep192 in PCNT/CDK5RAP2 double-depleted cells. HeLa cells were treated with control siRNA or PCNT/CDK5RAP2 siRNA for 48 h and stained with the indicated antibodies. Scale bar, 1 µm. (F, G) Representative line intensity profiles (F) and measured diameters (G) of GT335 and Cep192. The line profiles were measured along the dotted lines in (E). The profiles were fitted with double Gaussian
curves and the distances between the half-maximal intensity points at the far ends were measured as the diameters (schematically indicated with dotted lines and arrows in the profiles; fitted curves are not shown). Horizontal bars and error bars in the plots for the diameters represent median and interquartile range. N = 18 (for siControl) or 22 (for siPCNT/CDK5RAP2) centrosomes; data from two independent experiments were pooled. Mann–Whitney U-test was used to determine the significance of the difference. *P < 0.0001.
Figure 2. Cells with one centrosome can organize bipolar spindles in mitosis by forming a PCM-positive acentriolar spindle pole (PCM-pole).

(A) Schematic illustration of centrinone-induced removal of centriole. (B) DMSO-treated control mitotic spindles (two centrosomes) and centrinone B-treated centrosome-depleted spindles (one or zero centrosomes). Green, red, and blue represent GT335 (polyglutamylated centriole microtubules), α-tubulin, and DNA, respectively. Z-projections: 12 planes, each 0.13 μm apart. Scale bar, 5 μm. (C-E) Mitotic duration, the time required from NEBD to cytokinesis, in DMSO-treated two-centrosome (C), centrinone-treated one-centrosome (D) and zero-centrosome (E) cells, in Fig (S1G-I). Line and error bars represent the mean and SD (N ≥ 20 cells from two independent experiments). Kruskal–Wallis test was used to determine the significance of the difference. *P < 0.05, **P < 0.005, *** P < 0.0001. (F) Distribution of centrosomal factors in centriolar and acentriolar spindle poles. DMSO-treated control mitotic spindles (two centrosomes) and centrinone-treated mitotic spindles (one or zero centrosome) in HeLa cells. Green, red, and blue represent GT335, the protein of interest (Pericentrin, CDK5RAP2, or CEP192), and DNA, respectively. Z-projections: 21 sections, every 1
μm. Scale bar, 10 μm. (G) Quantification of pole patterns in (F). Values are presented as mean percentages from two independent experiments (N = 25 for each experiment). (H) Schematic illustration of PCM localization at spindle poles in two-, one- or zero-centrosome cells.
Figure 3. The PCM-pole is formed by splitting PCM from the centriolar pole or by accumulation of PCM components.

(A–D) HeLa cells expressing EGFP-centrin1 and pericentrin-mCherry were observed with a 60x objective. Magenta and green represent pericentrin and centrin, respectively. Z projections: 20 planes, 1.2 μm apart. Scale bar, 10 μm. Time zero corresponds to the beginning of mitotic cell rounding. (A) Splitting of the PCM components from the centriolar pole in one-centrosome cells. Arrowheads indicate the PCM at the acentriolar spindle pole. (B) PCM accumulation in one-centrosome cells. Arrowheads indicate the accumulation of PCM at acentriolar spindle poles. (C) Cell division in zero-centrosome cells without accumulation of PCM. (D) Quantification of patterns of PCM dynamics in (A–C). Values are percentages of the total cells from 52 (for one-centrosome cells) or 24 (for zero-centrosome cells) cells from two independent experiments. (E) Averaged time courses of pericentrin-mCherry or CDK5RAP2-mCherry signals at the centriolar spindle pole and PCM-pole of 10 cells. Time course data were aligned at PCM-pole formation (0 h). Error bars, SD; A.U., arbitrary units. (F) Schematic illustration of PCM-pole
formation by splitting PCM from the centriolar spindle pole or by accumulation of PCM components.
Figure 4. Pericentrin and CDK5RAP2 are crucial for spindle elongation and spindle bipolarization of one-centrosome cells.

(A) Mitotic spindle structures upon siRNA treatment with or without 500 nM of CentB. Green, red, and blue represent GT335, α-tubulin, and DNA, respectively. Z-projections: 5 planes, 0.3 μm apart. Scale bar, 5 μm. (B) Frequency of mitotic spindle structures after siRNA treatment against the indicated proteins in (A). Values are presented as mean percentages. N > 86, data from two independent experiments were pooled. (C) Time-lapse observation of the structure of NuMA and microtubules upon siRNA treatment against the indicated proteins. Centrinone-treated one-centrosome HeLa cells expressing EGFP-centrin1 and pericentrin-mCherry were observed with a 40× objective. Red, green, and gray represent NuMA, centrin, and, SiR-tubulin, respectively. Z-projections: 10 planes, 2.2 μm apart. Scale bar, 10 μm. Time zero corresponds to nuclear envelope break down (NEBD). Arrowheads indicate the separated two NuMA foci. (D) The time required for the initial establishment of two poles of NuMA in (C). Line and error bars represent the mean and SD (N ≥ 60 cells from three independent experiments). Kruskal–Wallis test was used to determine the significance of the difference. n.s., not significantly
different \((P > 0.05)\). (E) Mitotic duration, the time required from NEBD to cytokinesis, in (C). Line and error bars represent the mean and SD \((N \geq 60\) cells from three independent experiments). Kruskal–Wallis test was used to determine the significance of the difference. \(*P < 0.01, **P < 0.001\). (F) Table of the times from nuclear envelope break down (NEBD) to cytokinesis in (E). (G) The signal intensity of CEP192 on acentrosomal spindle poles. Line and error bars represent the mean and SD \((N \geq 46\) cells from two independent experiments). Kruskal–Wallis test was used to determine the significance of the difference. \(*P < 0.0001\).
Figure 5. CEP57 depletion leads to an increase of PCM at the acentriolar pole and facilitates spindle bipolarization in one-centrosome cells.

(A) Mitotic spindle pole structures of one-centrosome cells upon CEP57 depletion. Green, red, and blue represent centrin, pericentrin, and DNA, respectively. Z-projections: 20 planes, 0.5 μm apart. Scale bar, 5 μm. (B) The signal intensity of pericentrin on centrosomes or PCM-poles in (A). Line and error bars represent the mean and SD (N ≥ 50 cells from two independent experiments). Kruskal–Wallis test was used to determine the significance of the difference. *P < 0.01. (C) Schematic illustration of CEP57-depletion-induced pericentrin accumulation at the PCM-pole. (D) Time-lapse observation of NuMA structures and microtubules upon CEP57 depletion. Centrinoone-treated one-centrosome HeLa cells expressing EGFP-centrin1 and pericentrin-mCherry were observed with a 40× objective. Red, green, and gray represent NuMA, centrin, and SiR-tubulin, respectively. Z-projections: 10 planes, 2.2 μm apart. Scale bar, 10 μm. Time zero corresponds to mitotic onset. (E) The time required for the initial establishment of two poles of NuMA in (D). Line and error bars represent the mean and SD (N ≥ 50 cells from two independent experiments). The Mann–Whitney U-test (two-tailed) was used to obtain a P-value. n.s., not significantly different (P > 0.05). (F) Mitotic duration, the time
required from nuclear envelope break down (NEBD) to cytokinesis, in (D). Line and error bars represent the mean and SD (N ≥ 50 cells from two independent experiments). The Mann–Whitney U-test (two-tailed) was used to obtain a P-value. *P < 0.0001. (G)

Frequency of mitotic spindle structures upon CEP57 depletion. Values are presented as mean percentages ± SD (N = 6, triplicates, two independent experiments, at least 29 spindles in each assay).
Figure 6. Pericentrin is crucial for the bipolar spindle elongation in cells with two centrosomes.

(A) Mitotic spindle structures upon treatment with siRNA in cells with two centrosomes. Red and blue represent α-tubulin and DNA, respectively. Z-projections: 21 planes, 1 μm apart. Scale bar, 5 μm. (B) Quantification spindle length of HeLa cells (N > 14, triplicates, from two independent experiments). Line and error bars represent the mean and SD. One-way ANOVA with Tukey’s multiple comparisons test was used to determine the significance of the difference. *P < 0.005. (C) CEP192 observed in two-centrosome cells. Red, and blue represent GT335, CEP192, and DNA, respectively. Z-projections: 20 planes, 0.5 μm apart. Scale bar, 5 μm. (D) The signal intensity of CEP192 on centrosomes in (C). Line and error bars represent the mean and SD (N ≥ 50 cells from two independent experiments). Kruskal–Wallis test was used to determine the significance of the difference. *P < 0.0001. n.s., not significantly different. (E) Time-lapse observation of the structure of microtubules upon depletion of pericentrin in HeLa cells were observed with a 40× objective. Gray represent SiR-tubulin, respectively. Z-projections: 10 planes, 2.2 μm apart. Scale bar, 10 μm. Time zero corresponds to nuclear envelope break down (NEBD). (F) Averaged time courses of the pole length at each time point in (E). The
length between two poles of spindle was measured from 40 cells from two independent experiments. Time course data were aligned at the time of the NEBD (0 min). Error bars, SD. (G) Mitotic spindle structures of A549 and U2OS cells. Red, and blue represent α-tubulin, and DNA, respectively. Z-projections: 31 planes, 0.5 μm apart. Scale bar, 10 μm. Quantification spindle length of A549 (H) and U2OS (I) cells (N > 40 from two independent experiments). Line and error bars represent the mean and SD. The Mann–Whitney U-test (two-tailed) was used to obtain a P-value. *P < 0.005, **P < 0.0001.
Figure 7. PLK1 is crucial for PCM-pole formation and bipolar spindle formation in one-centrosome cells.

(A, B) PLK1 and phosphorylated PLK1 observed in one-centrosome cells. (A) Red, green, and blue represent PLK1, GT335, and DNA, respectively. Z-projections: 20 planes, 1 µm apart. Scale bar, 10 µm. (B) Frequency of localization of PLK1 in (A). Values are presented as mean percentages from two independent experiments (N > 40 from two experiments). (C) Red, green, and blue represent phosphorylated PLK1, centrin, and DNA, respectively. Z-projections of 20 sections, every 1 µm. Scale bar, 5 µm. (D) Frequency of localization of phosphorylated PLK1 in (A). Values are presented as mean percentages from triplicates (N > 40 from two experiments). (E) Mitotic spindle structures upon PLK1 inhibition with or without 100 nM of centrinone. HeLa cells expressing EGFP-centrin1 and Pericentrin-mCherry were observed with a 63× objective. Green, red, gray, and blue represent GFP (centrin1), RFP (Pericentrin), α-tubulin, and DNA, respectively. Z-projections: 10 planes, 0.3 µm apart. Scale bar, 5 µm. (F) Frequency of mitotic spindle structures in (A). Values are mean percentages from two independent experiments (N = 50 for each experiment). (G) The signal intensity of
Pericentrin on centrin foci of fixed mitotic HeLa cells expressing EGFP-centrin1 and Pericentrin-mCherry (N > 45 for each condition). Line and error bars represent median with interquartile range. Kruskal–Wallis test was used to determine the significance of the difference. *P < 0.05, **P < 0.0001.
Figure 8. Dual inhibition of PLK1 and PLK4 prevents cancer cell proliferation.

(A) Dual inhibition of PLK1 and PLK4 induced cell death in HeLa cells. Cell viability (% of DMSO or CentB mono-treatment) was determined after treatment with CentB and various concentrations of PLK1 inhibitor. (B) Mitotic cell fate of individual cells upon PLK1 and PLK4 dual inhibition. After centrosome depletion, treatment with a low dose of PLK1 inhibitor induced cell death. The mitotic cell fate of HeLa cells expressing EGFP-centrin1 and pericentrin-mCherry after treatment with 100 nM of centrinone, 1 nM of BI2536, or 100 nM of centrinone + 1 nM of BI2536 was observed compared to DMSO alone (solvent control). Cells were observed with a 60× objective. Z-projections: 30 planes, 1 μm apart. Scale bar, 5 μm. Time zero corresponds to the beginning of mitotic cell rounding. (C) Frequency of mitotic events in (B). Values are presented as percentages. N = 205 (for DMSO control, two centrosomes), 211 (for BI, two centrosomes), 107 (for centrinone, two centrosomes), 92 (for centrinone + BI, two centrosomes), 205 (for centrinone, one centrosome), 60 (for centrinone + BI, one centrosome) cells; data from two independent experiments were pooled. (D) Dual inhibition of PLK1 and PLK4 induced the cleavage of PARP. HeLa cells were treated with the drugs for 48 h and PARP cleavage was analyzed by western blotting. The concentrations of centrinone and BI 2536
were 100 nM and 1 nM, respectively. (E) Dual inhibition of both PLK1 and PLK4 in various cancer cell lines. After four days of treatment with 500 nM of centrinone B and various concentrations of BI2536 (0, 0.1, 0.5, 1, 2, 5, 10, or 20 nM), cell viability (% of DMSO control) was determined and is shown as a heat map. The ratios between IC_{50} values (± centrinone B) are shown below the heat map. Exact IC_{50} values are provided in Table 1.
Figure 9. The Centriole and PCM cooperate to recruit CEP192 to the spindle pole to facilitate bipolar spindle formation.

Schematic illustration of the assembly of CEP192 at the spindle pole by the centriole and PCM in human cells. For details, see the Discussion section.
Supplementary Figure 1. Depletion of PCM components in two-, one-, and zero-centrosome cells.

(A-C) Western blot analysis of the efficiency of protein depletion of CEP192 (A), CDK5RAP2 (B) and pericentrin (C) after 48 h of siRNA transfection in HeLa cells. (D-F) Quantification of depleted centrosomal CEP192 (D), CDK5RAP2 (E) and pericentrin. The Mann–Whitney U-test (two-tailed) was used to obtain a P-value. (P < 0.0001). (G-I). Time-lapse observation of the structure of microtubules upon siRNA treatment against the indicated proteins. DMSO-treated two-centrosome (G), centrinone-treated one-centrosome (H), and centrinone-treated zero-centrosome (I) HeLa cells expressing EGFP-centrin1 and pericentrin-mCherry were observed with a 40x objective. Gray represent SiR-tubulin, respectively. Z-projections: 10 planes, 2.2 μm apart. Scale bar, 10 μm. Time zero corresponds to nuclear envelope break down (NEBD).
Supplementary Figure 2. Distribution of centrosomal factors in centriolar and acentriolar spindle poles.

(A–D) Distribution of centrosomal factors in centriolar and acentriolar spindle poles. (A) DMSO-treated control mitotic spindles (two centrosomes) and centrinone-treated spindles (one or zero centrosomes) of HeLa cells. Green, red, and blue represent GT335, protein of interest (γ-tubulin, CEP152, or CPAP), and DNA, respectively. Z-projections: 21 planes, 1 μm apart. Scale bar, 10 μm. (B) PCM-poles were observed in various cells. Magenta, cyan, and blue represent CEP192, GT335, and DNA, respectively. Z-projections: 40 planes, 0.3 μm apart. Scale bar, 5 μm. (C) PCM-poles were observed in one-centrosome spindles induced by SAS6 depletion. Red, green, and blue represent centrin, pericentrin, and DNA, respectively. Scale bar, 5 μm. (D) Quantification of pole patterns in (C). Values are presented as mean percentages from three independent experiments (N = 20 for each experiment). (E) Microtubule nucleation from the PCM-pole. Following treatment with ice, microtubule nucleation (5 min at 25°C) was observed in one-centrosome cells. Gray, red, green, and blue in the merged image represent EB1,
pericentrin, centrin, and DNA, respectively. Z-projections: 21 planes, 1 µm apart. Scale bar, 5 µm.
Supplementary Figure 3. Pericentrin and CDK5RAP2 are crucial for the cell division of one-centrosome cells.

(A-D) Mitotic spindle structures in siPCM-treated RPE1 (A) and A549 (C) cells. After a 2-day siRNA treatment with or without 100 nM of centrinone, mitotic spindle structures were observed with a 63× objective. Green, red, and blue represent GT335, α-tubulin, and DNA, respectively. Z projections: 31 planes, 0.5 μm apart. Scale bar, 10 μm.

Frequency of mitotic spindle structures in RPE1 (B) and A549 (D) cells. Values are mean percentages from two independent experiments (N > 20 for each experiment).
Supplementary Figure 4. Pericentrin is critical for proper spindle elongation in two-centrosome cells.

(A) Time-lapse observation of the structure of microtubules upon siRNA treatment against the indicated proteins. HCT116 cells were observed with a 40× objective. Gray represent SiR-tubulin, respectively. Z-projections: 10 planes, 2.2 μm apart. Scale bar, 10 μm. Time zero corresponds to nuclear envelope break down (NEBD). (B) Averaged time courses of the pole length at each time point in (A). The length between two poles of spindle was measured from 40 cells from two independent experiments. Time course data were aligned at the time of the NEBD (0 min). Error bars, SD. (C) The time required for the establishment of two spindle poles in (A). Line and error bars represent the mean and SD (N ≥ 75 cells from two independent experiments). The Mann–Whitney U-test (two-tailed) was used to obtain a P-value. (P < 0.05). (D) Mitotic spindle structures of RPE1, GI1, SKOV3, A431 and PANC1 cells. Red, and blue represent α-tubulin, and DNA, respectively. Z-projections: 31 planes, 0.5 μm apart. Scale bar, 10 μm. (E-I) Quantification spindle length (N > 40 from two independent experiments) in (D). Line
and error bars represent the mean and SD. The Mann–Whitney \( U \)-test (two-tailed) was used to obtain a \( P \)-value. \( P < 0.0005 \). n.s., not significantly different.
Supplementary Figure 5. Dual inhibition of PLK1 and PLK4 prolongs mitosis and induces cell death in one-centrosome HeLa cells.

Each plot shows the cumulative percentage of cell division or cell death in mitosis at each time point of Fig. 6B, C. N = 205 (for DMSO control, two centrosomes), 211 (for BI, two centrosomes), 107 (for centrinone, two centrosomes), 92 (for centrinone + BI, two centrosomes), 205 (for centrinone, one centrosome), 60 (for centrinone + BI, one centrosome) cells; data from two independent experiments were pooled.
| Cell line | Type of cancer | $IC_{50}$ values of BI2536 (nM) (−CentB) | $IC_{50}$ values of BI2536 (nM) (+CentB) | Ratio of $IC_{50}$ values (−CentB/+CentB) |
|-----------|----------------|----------------------------------------|----------------------------------------|----------------------------------------|
| HeLa      | Cervical cancer | 1.1 ± 0.1                              | 0.4 ± 0.2                              | 2.8                                    |
| U2OS      | Osteosarcoma    | 11.0 ± 3.9                              | 3.6 ± 0.6                              | 3.1                                    |
| Saos-2    | Osteosarcoma    | 2.1 ± 0.4                              | 0.8 ± 0.3                              | 2.6                                    |
| A549      | Lung carcinoma  | 2.3 ± 0.6                              | 1.1 ± 0.2                              | 2.1                                    |
| DAUDI     | Burkitt lymphoma| 3.5 ± 0.6                              | 1.4 ± 0.3                              | 2.5                                    |
| RAJI      | Burkitt lymphoma| 4.5 ± 0.9                              | 1.0 ± 0.1                              | 4.5                                    |
| HL60      | Leukemia        | Not determined                          | Not determined                          | Not determined                         |
| K562      | Leukemia        | 13.4 ± 0.5                             | 2.1 ± 0.6                              | 6.4                                    |
| MKN45     | Gastric cancer  | 4.7 ± 1.3                              | 2.4 ± 0.3                              | 2.0                                    |
| MKN1      | Gastric cancer  | 4.1 ± 0.9                              | 3.4 ± 1.3                              | 1.2                                    |
| DU145     | Prostate carcinoma brain metastasis | 3.1 ± 0.7 | 1.2 ± 0.4 | 2.6 |
| PC-3      | Prostate carcinoma bone metastasis | 3.7 ± 1.8 | 2.0 ± 0.9 | 1.9 |
| PANC-1    | Pancreatic carcinoma | 3.3 ± 0.8 | 2.2 ± 0.6 | 1.5 |
| GI-1      | Glioma          | 3.5 ± 0.2                              | 2.4 ± 0.4                              | 1.5                                    |
| HCT116    | Colon cancer    | 5.7 ± 4.0                              | 3.2 ± 1.6                              | 1.8                                    |
| SKOV-3    | Ovarian cancer  | 2.9 ± 0.6                              | 1.7 ± 0.3                              | 1.7                                    |
| A431      | Epidermoid carcinoma | 2.5 ± 0.5 | 2.0 ± 0.9 | 1.3 |
| HMV-II    | Melanoma        | 10.6 ± 3.0                             | 7.6 ± 4.3                              | 1.4                                    |
| MCF-7     | Breast adenocarcinoma | 6.4 ± 1.6 | 4.3 ± 1.3 | 1.5 |
Various cancer cells were treated with BI 2536 with or without 500 nM of centrinone B for four days and their viabilities were determined using the WST-8 assay. IC$_{50}$ values and the ratio between IC$_{50}$ values (±CentB) are shown. Related to Fig. 6E.

Table S2. IC$_{50}$ values of mitotic inhibitors against the growth of HeLa cells with or without centrinone B (CentB).

| Compounds (Targets) | IC$_{50}$ values (-CentB) | IC$_{50}$ values (+CentB) | Ratio of IC$_{50}$ values (-CentB/+CentB) |
|---------------------|---------------------------|---------------------------|------------------------------------------|
| BI2536 (PLK1)       | 1.1 nM                    | 0.6 nM                    | 1.8                                      |
| Cytochalasin B (β-actin) | 1.5 μM                  | 1.6 μM                    | 0.9                                      |
| Paclitaxel (α/β-tubulin) | 1.8 nM                   | 1.7 nM                    | 1.1                                      |

HeLa cells were treated with several compounds with or without 500 nM of centrinone B for 3 days and their viabilities were determined by WST-8 assay. IC$_{50}$ values (average from at least two independent assays) and the ratio between IC$_{50}$ values (±CentB) were shown.
Figure 1, Chinen, Yamazaki et al

A 2-centrosome (min)

B Time from NEBD to cytokinesis (min)

C SiR-Tubulin

D

E MERGE GT335 CEP192

F

G GT335 n.s.
Figure 2, Chinen, Yamazaki et al.
Figure 3, Chinen, Yamazaki et al

A 1-centrosome (min): Split

B 1-centrosome (min): Accumulation

C 0-centrosome (min): No accumulation

D E 1-centrosome

F Accumulation of PCM
Figure 4, Chinen, Yamazaki et al

A

B

C

D

E

F

G

1-centrosome (min)

siControl siCDK5RAP2 siPericentrin

GT335 α-Tubulin DNA

Bipolar

2-centrosome

Monopolar

Monopolar

1-centrosome

Bipolar

Bipolar

Bipolar

siCDK5RAP2

siPericentrin

siControl

siCDK5RAP2

siPericentrin

siControl

siCDK5RAP2

siPericentrin

B

100

80

60

40

20

0

Cells (%)

siControl siCDK5RAP2 siPericentrin

siControl siCDK5RAP2 siPericentrin

siControl siCDK5RAP2 siPericentrin

2

1

Centrosomes

10

50

170

SiR

80

130

240

mCherry

0

100

360

1

-10

0

10

50

80

150

240

150

200

100

170

260

320

NuMA two foci

EGFP-Centrin mCherry-NuMA SiR-Tubulin

D

E

F

siControl

siCDK5RAP2

siPericentrin

n.s.

n.s.

1-centrosome (min)

Time from NEBD to establishment of NuMA two foci

Sample

siControl

siCDK5RAP2

siPericentrin

Time from NEBD to cytokinesis (min)

290.0 ± 169.9 min

437.2 ± 296.4 min

395.1 ± 246.6 min

G

CEP92 at acentrosolar pole (normalized to siControl signal)

1-centrosome

*
**Figure 5, Chinen, Yamazaki et al**

(A) 1-centrosome

- siControl
- siCEP57

Centrin Pericentrin DNA

(B) 1-centrosome

![Diagram showing Pericentrin intensity](image)

Pericentrin intensity (control ratio)

- Centriolar pole
- PCM-pole

siControl siCEP57

PCM accumulation at PCM-pole

(C) PCM and Centriole

- siControl
- siCEP57

(D) 1-centrosome (min)

- siControl
- siCEP57

-10 0 30 80 110 160 220

EGFP-Centrin mCherry-NuMA SiR-Tubulin

(E) Time from NEBD to establishment of NuMA two foci (min)

- siControl
- siCEP57

(F) Time from NEBD to cytokinesis (min)

- siControl
- siCEP57

(G) Cells (%)

- Bipolar
- Monopolar-like
- Others
Figure 6, Chinen, Yamazaki et al

A) HeLa 2-centrosome

siControl | siCDK5RAP2 | siPericentrin

B) 2-centrosome HeLa

Graph showing spindle length (µm) for siControl, siCDK5RAP2, and siPericentrin.

C) HeLa 2-centrosome

GT35 CEP192 DNA

D) 2-centrosome HeLa

Graph showing CEP192 intensity (control ratio) for siControl, siCDK5RAP2, and siPericentrin.

E) 2-centrosome HeLa (min)

Graph showing spindle length (µm) with time from NEBD (min).

F) 2-centrosome HeLa

Graph showing spindle length (µm) for siControl and siPericentrin.

G) 2-centrosome

A549 | U2OS

siControl | siPericentrin

α-Tubulin DNA

H) 2-centrosome A549

Graph showing spindle length (µm) for siControl and siPericentrin.

I) 2-centrosome U2OS

Graph showing spindle length (µm) for siControl and siPericentrin.
Figure 7, Chinen, Yamazaki et al

A 1-centrosome

B 120

PLK1 at acentriolar pole

C 1-centrosome

D 120

pPLK1 at acentriolar pole

E DMSO

BI 2536 1 nM

Centrinone 100 nM

BI 2536 + Centrinone

DNA Centrin Pericentrin α-Tubulin

F

Bipolar

Bipolar with chromosome congression error

Monopolar

Others

G

Pericentrin intensity at centriolar pole (control ratio)

BI 2536

Centrinone

Centrosomes
Figure 8, Chinen, Yamazaki et al

A. Cell growth (control%) as a function of BI 2536 concentration (nM).

B. Images showing cell division at different time points (min) under the conditions of Centrinone 100 nM + BI 2536 1 nM.

C. Graph depicting Mitotic cell death and Cell division with bars for different conditions of BI 2536 and Centrinone.

D. Western blot analysis showing PARP and Cleaved PARP, and HSP90 under different conditions of BI 2536 and Centrinone.

E. Heatmap representing the Growth (%), BI 2536 concentration, and Centrinone B sensitivity for various cell lines.

Legend:
- Green: Centrinone B -
- Orange: Centrinone B +
- Black: BI 2536 -
- White: BI 2536 +
- Yellow: Centrinone -
- Red: Centrinone +

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Figure 9, Chinen, Yamazaki et al

A model for bipolar spindle formation redundantly supported by centriole and PCM

2-centrosome cell

Centriole recruits CEP192 to centriolar wall

Pericentrin and CDK5RAP2 recruit CEP192 to PCM

PCM (Pericentrin, CDK5RAP2)
Centriole
CEP192

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Supporting figure 1, Related to Fig 2.

A. siCEP192: - +

B. siCDK5RAP2: - +

C. siPericentrin: - +

D. CEP192 at centrosomes (normalized to siControl signal)

E. CDK5RAP2 at centrosomes (normalized to siControl signal)

F. Pericentrin at centrosomes (normalized to siControl signal)

G. 2-centrosome HeLa (min)

H. 1-centrosome HeLa (min)

I. 0-centrosome HeLa (min)
Supporting figure 2, Related to Fig 2.

A

| Centrosomes | 2 | 1 | 0 |
|-------------|---|---|---|
| γ-Tubulin   |   |   |   |
| CEP152      |   |   |   |
| CPAP        |   |   |   |

GT335 DNA

B

| 1-centrosome | A431 | PC3 | DU145 |
|--------------|------|-----|-------|
|               | PANC-1 | MCF-7 | A549 |

GT335 CEP192 DNA

C

HCT116
SAS6-AID
1-centrosome

Centrin DNA
Pericentrin

D

| Cells (%) | 120 | 100 | 80 | 60 | 40 | 20 | 0 |
|-----------|-----|-----|----|----|----|----|---|
| +         |     |     |    |    |    |    |   |
| -         |     |     |    |    |    |    |   |

Pericentrin at acentriolar pole

E

| 1-centrosome | On ice | MT regrowth 5 min |
|--------------|--------|-------------------|

Centrin Pericentrin EB1 DNA
Supporting figure 3, Related to Fig 4.

**A** RPE1 Centrosomes

|      | siControl | siCEP192 | siCDK5 | siPericentrin |
|------|-----------|----------|--------|--------------|
| 2    |           |          |        |              |
| 1    |           |          |        |              |

GT335 α-Tubulin DNA

**B** RPE1 2-centrosome

- siControl
- siCEP192
- siCDK5RAP2
- siPericentrin

Cells (%)

- Bipolar
- Monopolar-like
- Others

**C** A549 Centrosomes

|      | siControl | siCEP192 | siCDK5 | siPericentrin |
|------|-----------|----------|--------|--------------|
| 2    |           |          |        |              |
| 1    |           |          |        |              |

GT335 α-Tubulin DNA

**D** A549 2-centrosome

- siControl
- siCEP192
- siCDK5RAP2
- siPericentrin

Cells (%)

- Bipolar
- Monopolar-like
- Others

A549 1-centrosome

- siControl
- siCEP192
- siCDK5RAP2
- siPericentrin

Cells (%)

- Bipolar
- Monopolar-like
- Others
Supporting figure 4, Related to Fig 6.

A. 2-centrosome HCT116 (min)

B. 2-centrosome HCT116

C. 2-centrosome

D. RPE1

E. GI1

F. SKOV3

G. A431

H. PANC1

- n.s.

- *
Supporting figure 5, Related to Fig 8.

A

2-centrosome

| Treatment          | Event in mitosis (%) |
|--------------------|----------------------|
| DMSO               | ![Graph](image)      |
| BI 2536            | ![Graph](image)      |
| Centrinone         | ![Graph](image)      |
| Centrinone + BI 2536 | ![Graph](image) |

Time after mitotic entry (min)

B

1-centrosome

| Treatment          | Event in mitosis (%) |
|--------------------|----------------------|
| Centrinone         | ![Graph](image)      |
| Centrinone + BI 2536 | ![Graph](image) |

Event in mitosis (%)

Time after mitotic entry (min)

- Cell division
- Mitotic cell death