HURP permits MTOC sorting for robust meiotic spindle bipolarity, similar to extra centrosome clustering in cancer cells

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Introduction

During meiosis, the assembly of a microtubule (MT)-based bipolar spindle is essential for the formation of a competent egg, capable upon fertilization of generating viable offspring. The spindle ensures accurate chromosome segregation to prevent aneuploidy of the embryo. It also defines the cell division plane, leading to the formation of a large egg and a tiny polar body. In most species, meiotic spindles lack canonical centrosomes. Instead, the formation of a bipolar spindle results from organization of MTs and, in vertebrates, from multiple, discrete MT-organizing centers (MTOCs) around chromosomes (Dumont et al., 2007; Schuh and Ellenberg, 2007). Little is known about the mechanisms that govern the MT-MTOC reorganization into a bipolar structure. This is partly because of the lack of genetic studies on meiotic spindle organization using loss-of-function mutants in vertebrates. To characterize molecular factors involved in meiotic spindle assembly, we have started to analyze spindle assembly factors of knockout mice that are viable but female sterile, potentially caused by essential functions in meiotic divisions. Using this strategy, we have identified hepatoma up-regulated protein (HURP) as a key meiotic factor. HURP is an MT-associated protein and a Ran GTPase effector previously identified for its function in mitotic kinetochore fiber stability. However, it is dispensable for mitosis completion (Koffa et al., 2006; Silljé et al., 2006). Here, we show that HURP, via its local concentration by activity of the Kinesin-5 motor, is required for the assembly of a robust central MT domain, itself allowing the establishment and maintenance of meiotic spindle bipolarity. This domain acts as a scaffold for the accurate sorting of MTOCs to spindle poles and serves to congress chromosomes toward the spindle equator. This new function for HURP sheds light on the complex architecture of the female meiotic spindle. Moreover, by extending our study to cancer cells bearing multiple centrosomes (Godinho et al., 2009), we have

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we present evidence that the genetic requirement for meiosis and the pathways used by some tumors to divide might be evolutionarily shared.

Results and discussion

Hurp deficiency specifically impairs completion of meiotic divisions in oocytes

Knockout mice for Hurp were found to be viable but female sterile, indicating that HURP is dispensable for mitosis (Tsai et al., 2008). We reasoned that the sterility could be caused by abnormal oocyte meiotic divisions giving rise to aneuploid embryos upon fertilization. Therefore, we analyzed meiotic progression of oocytes collected from Hurp−/− females. Although HURP is expressed at a constant level in maturing wild-type (wt) oocytes (Fig. 1 A), Hurp−/− females harbor undetectable levels of the protein, confirming a true knockout (Fig. 1 B). Most wt oocytes extrude their first polar body 9 h after nuclear envelope breakdown (NEBD; Fig. 1 C). In contrast, 12 h after NEBD, only 50% of Hurp−/− oocytes extruded their polar bodies (Fig. 1 C). Associated with this delay, chromosome dynamics were altered. Homologous chromosomes neither congressed properly nor formed a proper metaphase plate (Fig. 1 D, Video 1, and Video 2). More than 60% of Hurp−/− oocytes exhibited lagging chromosomes at anaphase (Fig. 1 D, arrow; Fig. 2 A; Fig. 3 A). This is consistent with the formation of stable K fibers only at the end of MI (Brunet et al., 1999) and suggested that HURP performs additional functions during meiosis, which were not anticipated from studies in mitosis (Silljé et al., 2006; Wong and Fang, 2006).

Hurp deficiency affects the setting up of meiotic spindle bipolarity

Meiotic spindle assembly in the absence of HURP was analyzed by time-lapse spinning-disk microscopy using EB3-GFP, a well-characterized MT plus-end marker, which was previously used in mouse oocytes (Schuh and Ellenberg, 2007). It gives a strong signal throughout the spindle, presumably caused by short MTs of mixed polarity, which are abundant in meiotic spindles (Srayko et al., 2006; Yang et al., 2007; Liang et al., 2009). In wt, at NEBD, multiple MTOCs and MTs formed evenly distributed asters associated with chromosomes (Fig. 2 A), which were sorted into a bipolar structure 3.5 h later (Fig. 2 C). In contrast, in Hurp−/− oocytes, spindle assembly was impaired (Fig. 2 A). After NEBD, MTOCs and MTs rapidly became unevenly distributed, with chromosomes expelled to the periphery of the MT array (Fig. 2 B and Fig. S2 A). The bipolarization of such structures was nevertheless observed with a delay of 1.5 h (Fig. 2 C). Hence, HURP is required for timely bipolarization at early stages of MI.

Hurp deficiency impairs the maintenance of meiotic spindle bipolarity

In wt oocytes, upon bipolarization, chromosomes converged to the spindle equator, and eventually, 9 h after NEBD on average, anaphase and chromosome segregation occurred. The second meiotic spindle then rapidly assembled and remained stable during the MII arrest (Fig. 2 A and Video 4). In 80% of Hurp−/− oocytes, spindles observed in MI were unsteady (Fig. 2 D). In 80% of Hurp−/− oocytes, spindles observed in MI were unsteady (Fig. 2 D). Spindle collapse, irreversible in some cases, was partially reverted in 60% of oocytes within a few hours (Fig. 2 D and Video 6). To quantify this, spindle lengths were scored throughout 17 h of meiotic maturation (Fig. 2 E). In Hurp−/− oocytes, the mean spindle length was significantly reduced compared with wt (18.54 ± 2.62 µm vs. 26.42 ± 3.18 µm, respectively; Fig. 3 A). Remarkably, although size was stable in controls (Fig. 2 E, green curve), it varied over time in Hurp−/− oocytes, even for spindles reaching the expected spindle length (Fig. 2 E, red curves). Therefore, HURP is required for establishment and maintenance of meiotic spindle bipolarity.

HURP localizes to a central domain of the meiotic spindle

HURP localization and dynamics were then analyzed in maturing oocytes. At NEBD, multiple MTOCs and associated MTs surround the chromosomes to organize into a bipolar spindle. At this stage, HURP was enriched on MTs assembled between MTOCs and chromosomes (Fig. 1 F and Fig. S1 A). After spindle bipolarization, HURP was restricted to the central domain of MI and MII spindles (Fig. 1 F). Such localization, lost in Hurp−/− oocytes (Fig. S1 C), was also observed for GFP-tagged HURP (Fig. 1 G, Fig. S1 B, and Video 3). As expected, HURP localization was abolished upon MT depolymerization and expression of a dominant-negative Ran mutant (Fig. S1, D and E). In mitosis, HURP is considered to be a marker of kinetochore fibers (Koffa et al., 2006; Silljé et al., 2006). In oocytes, HURP could be observed on MTs emanating from the kinetochores only at the end of MI and in MII (Fig. 1 H). Strikingly, during MI, HURP was associated with nonkinetochore MTs crossing the spindle equator, reminiscent of intercalar MTs (Fig. 1 H).

HURP promotes MT stability in the central region of the spindle

We then measured relative MT densities using EB3-GFP. At NEBD, both wt and Hurp−/− oocytes exhibited comparable MT densities. However, the density of MTs increased with time in wt but remained steady in the Hurp deficiency (Fig. 3 A). Similarly, 2 h after NEBD, in oocytes treated with monastral to inhibit bipolarization, the density of MTs was lower, and monasters were smaller in Hurp−/− oocytes compared with wt (Fig. 3 B). These differences were not caused by disparities in EB3-GFP expression (Fig. S3, C and D). Using individual MT tracking, we did not observe significant differences in MT growth rates at NEBD (Fig. S3 A) or 2 h
We also analyzed MT density along the spindle axis later in MI. In wt, the density of MTs culminates in the spindle equator where HURP accumulates and where interpolar MTs later upon monastrol treatment, which facilitated MT tracking at this stage (Fig. S3 B). Thus, HURP does not promote MT growth but rather MT stability around chromosomes.

Figure 1. Hurp⁻/⁻ oocytes present late anaphases and lagging chromosomes and spread MII plates. (A) HURP levels are constant during meiotic maturation. Oocytes were collected in prophase I (Prol), 4 h after NEBD (MI), during polar body (PB) extrusion, and during MII. (B) HURP is absent in liver extracts from Hurp⁻/⁻ mice. (C) Percentage of first polar body extrusion over time in wt versus Hurp⁻/⁻ oocytes. (D) Aberrant bivalent separation in Hurp⁻/⁻ oocytes. (top) wt with normal chromosome separation. (bottom) Hurp⁻/⁻ with lagging chromosomes (arrow) and delayed anaphase I, both expressing H2B-RFP. Asterisks, polar bodies. All times are given in hours and minutes after NEBD. (E) Hurp⁻/⁻ oocytes present disorganized MII plates, which were rescued by HURP expression. Chromosomes in MII from wt (top left), Hurp⁻/⁻ (top right), and Hurp⁻/⁻ oocytes expressing GFP-HURP (bottom). (F) HURP localizes in the vicinity of chromosomes in the spindle midzone. Oocytes stained with Hoechst (blue), tubulin (green), and HURP. Asterisk, polar body. (G) GFP-HURP behaves like endogenous protein. Oocytes are shown expressing EB3-mCherry (green) and GFP-HURP. Maximal Z projections are shown. Asterisk, polar body. (H) HURP does not label K fibers in prometaphase I. Oocytes were stained at NEBD + 5 h (prometaphase I), NEBD + 8 h (MI), and MII with Hoechst; CREST for kinetochores; and HURP. Maximum projections or selected stacks are shown. Asterisks, kinetochore MTs not decorated by HURP. Arrows, HURP on interpolar MTs. Bars, 10 µm.
**Hurp is required for MTOC sorting toward spindle poles**

In oocytes lacking centrosomes, spindle assembly depends on the poorly understood self-organization of MTOCs and MTs around chromosomes. We characterized MTOC dynamics during MI. 2 h after NEBD, upon bipolarization, MTOCs were do overlap (Fig. 3 C, top). In contrast, MT density was lowest in this region in Hurp−/− oocytes (Fig. 3 C, bottom). Therefore, after bipolarization, the reduction of MT density is restricted to the central domain of the spindle in these oocytes. Collectively, HURP promotes the organization of MTs into a robust midzone.

**Hurp−/− oocytes are delayed in bipolarization and undergo spindle collapse.** (A) Hurp−/− oocytes present spindle defects. For A, B, and D, oocytes express EB3-GFP (green) and H2B-RFP (red). Asterisks, polar bodies. (B) MTs are not evenly distributed around chromosomes after NEBD in Hurp−/−. Mean projections of 10 Z planes are shown. (C) Setup of spindle bipolarity is delayed in Hurp−/− oocytes. Bipolarity was scored when two poles were distinguishable. It is delayed in Hurp−/− oocytes, starting at 297 ± 101 min instead of 209 ± 74 min (**, P = 0.0334). Lines indicate the mean time for bipolarity setup. (D) Transient spindle collapse in Hurp−/− oocytes during MI. Bipolarity reappears at 14 h and 40 min after NEBD. (E) Unstable spindle length in Hurp−/− oocytes. Length was determined by longitudinal line scans of images from wt [n = 12] and Hurp−/− [n = 14] oocytes. Representative curves are shown. All times are given in hours and minutes after NEBD. Bars, 10 µm.
Figure 3. *Hurp* /− oocytes show reduced MT density and improper MTOC distribution. (A) Reduced MT density early on in *Hurp* /− oocytes. Fluorescence intensity in wt and *Hurp* /− oocytes, expressing EB3-GFP and H2B-RFP, was assessed in the chromosome vicinity. MT density is lower in *Hurp* /− oocytes (*n* = 9) from NEBD + 30 and + 60 min (**, *P* = 0.0125 and 0.0071) compared with wt (*n* = 10). (B) Prometaphase I monasters are smaller, with fewer MTs in *Hurp* /− oocytes. Fluorescence intensity was assessed as in A in monastrol-treated oocytes at NEBD + 2 h. Mean monaster size: 29.19 ± 3.04 µm for *Hurp* /− compared with 32.86 ± 2.22 µm for wt (left; **, *P* = 0.0143). Monaster MT density is lower in *Hurp* /− than wt (right histogram; wt, *n* = 9; and *Hurp* /−, *n* = 7; **, *P* = 0.0228). Error bars represent standard deviation. (C) Low MT density in the spindle center in *Hurp* /− oocytes. Longitudinal line scans from MI spindles of wt (*n* = 6) and *Hurp* /− (*n* = 10). Intensities were normalized to maximum value within the same spindle, and spindle size was interpolated. Light curves, individual spindles. Dark curves, mean values. (D) MTOCs are progressively sorted at spindle poles in wt. (top) MTOCs were labeled using pericentrin (green) and chromosomes with Hoechst (red). (bottom) Quantitative analysis of MTOC distribution along the spindle axis at 2 (*n* = 16), 4 (*n* = 18), 6 (*n* = 22), and 8 h (*n* = 17) after NEBD. The pericentrin signal was binarized to assess the individual MTOC area (a.u., arbitrary units). Coordinates of MTOCs are plotted along the spindle axis (0–1). Median values are shown in red. (E) MTOCs are progressively clustered during MI. Box plot representation of individual MTOC size at selected time points. Median values are shown in red. (F) Abolished MTOC sorting in *Hurp* /− oocytes. wt and *Hurp* /− oocytes were observed at NEBD + 7 h and labeled as in D. (G) Quantification of MTOC distribution along the spindle axis in late MI in wt (*n* = 5) versus *Hurp* /− (*n* = 10). Pericentrin signal was binarized as in D. Individual MTOC size is plotted against its coordinates on the spindle axis (0–1). Bars, 10 µm.
Kinesin-5 activity is required to recruit HURP to the central region of the spindle

Kinesin-5 is a key motor protein for bipolarity setup of mitotic and meiotic spindles (Sawin et al., 1992; Walczak et al., 1998; Mailhes et al., 2004). HURP and Kinesin-5 are part of a large molecular complex (Koffa et al., 2006). We observed Kinesin-5 accumulation close to chromosomes before spindle bipolarization (Fig. S3 E) and on interpolar MTs (Fig. S3 E, red arrows). This indicated that Kinesin-5 and HURP localized similarly, potentially acting synergistically to support bipolarity. To test

Kinesin-5 recruits HURP to the spindle midzone for robust spindle bipolarity. (A) Loss of HURP accumulation in the central region upon Kinesin-5 inhibition. MII oocytes are shown after monastrol addition, expressing EB3-mCherry (left) and GFP-HURP (right). (B) Longitudinal line scan from oocytes observed in A. Intensity was normalized, and spindle length was interpolated (n = 5). (C) Representative spindle length and relative intensity of HURP fluorescence after monastrol addition. (D) HURP overexpression promotes central spindle assembly and perturbs spindle bipolarization and chromosome congression. GFP-HURP overexpression (see Materials and methods) affects spindle formation and MI division (noninjected: 100% polar body, n = 41; and GFP-HURP overexpression: 7% polar body, n = 67). (E) HURP overexpression disrupts MTOC clustering. Oocytes overexpressing HURP were labeled for DNA and MTOCs (pericentrin, green). (right) Quantification as in Fig. 3 G (n = 5). a.u., arbitrary units. All times are given in hours and minutes after NEBD. Bars, 10 µm.

scattered along the entire spindle axis (Fig. 3 D). Subsequently, they were targeted toward spindle extremities forming larger structures (Fig. 3, D and E) and were clustered at spindle poles by 6 h after NEBD (Fig. 3 D). The sorting and clustering were achieved long after a bipolar structure was visible (3.5 h after NEBD; Fig. 2 C). This indicates that robust spindle bipolarization relies on progressive outward sorting of MTOCs. Although at NEBD in Hurp−/− oocytes, MTOC distribution was similar to wt (Fig. S2, A and B), it was impaired later on (Fig. 3, F and G). Thus, HURP allows efficient sorting of MTOCs toward the poles.
HURP in proper meiotic spindle architecture. Furthermore, HURP overexpression also abolished MTOC sorting and clustering to the poles (Fig. 4 E). HURP is required to organize a central array within the meiotic spindle, which permits outward sorting of MTOCs and subsequent organization of spindle poles, as well as proper control of spindle length. All these parameters contribute to the definition of a robust bipolar spindle. The central array seems to favor homologue chromosome congression (Fig. S3 F).

HURP has a critical role in the clustering of extra centrosomes during mitosis in human cancer cells

The process of bipolar spindle assembly in the presence of multiple MTOCs in female meiosis resembles the clustering of extra centrosomes during cancer cell division. We hypothesized that the presence of extra centrosomes could impose an essential role of HURP in cancer cell division, which requires centrosome overexpression. HURP RNAi on August 17, 2017jcb.rupress.orgDownloaded from
clustering. To test this, we characterized mitosis in U2OS cells, in which centrosomes are overduplicated by overexpressing PLK4 (Kleylein-Sohn et al., 2007) after HURP RNAi. In controls, despite extra centrosomes, the majority of cells (~80%) successfully divide because of efficient clustering of extra centrosomes before anaphase (Fig. 5 A, top). The percentage of cells with transient multipolar spindle intermediates is similar in control versus HURP-depleted cells (Fig. 5 A, left). However, HURP knockdown results in a drastic increase in multipolar spindles in postanaphase cells (Fig. 5 A, bottom). 61.2% of HURP-depleted extracentrosomal cells (n = 262) display multipolar ana/telephase figures compared with 21.6% in controls (n = 164). Consistently, following live U2OS cells revealed that HURP knockdown induced an increase in fragmented multipolar anaphases (64.25 ± 2.3% in HURP RNAi vs. 22.93 ± 4.7% in controls; Fig. 5, C and D; Video 8; and Video 9). Our data suggest that HURP has a conserved role in MTOC organization and assembly of bipolar spindles in both oocytes and cancer cells with extra centrosomes.

In mitosis, HURP stabilizes K fibers but is dispensable. In contrast, we show that HURP is an essential meiotic spindle assembly factor. It associates with MTs and is enriched in the vicinity of chromosomes, where it promotes the assembly of a central MT array. This structure in turn allows the outward sorting of MTOCs to the poles. Through an original inside-out mechanism, HURP, possibly via Kinesin-5 activity, controls the formation and maintenance of a robust bipolar acentriolar spindle (Fig. S3 F). Interestingly, this central array is also important for proper chromosome alignment on the spindle equator, in agreement with previous findings that, in meiosis, chromosome congression does not rely on K fibers (Brunet et al., 1999; Wignall and Villeneuve, 2009). Oocytes undergo extremely asymmetric divisions in size. Losing canonical centrosomes seems to be a common strategy to minimize the size of polar bodies: lacking astral MTs, the distance between one spindle pole to the plasma membrane is reduced. One consequence of losing this powerful engine for astral MT assembly is a strong actin-based spindle positioning (Azoury et al., 2008; Schuh and Ellenberg, 2008). Another consequence might be the assembly of a robust central domain, based on chromosome-mediated MT assembly. A prominent central array of MTs has been reported for meiotic spindles assembled in Xenopus laevis egg extracts (Yang et al., 2008) and in Drosophila melanogaster oocytes, where it also depends on the activities of an MT-associated protein, INCENP (inner centromeric protein), and on a kinesin-like protein, Subito (Colombié et al., 2008). In mitotic cells, this meiotic mode of spindle assembly is probably present but dominated by other mechanisms, as previously demonstrated for the Kinesin-14 Ncd/HSET (non-claret disjunctional/human spleen embryo and testes; Kwon et al., 2008). Indeed, we present evidence that bundling extra centrosomes in cancer cells reveals a latent meiotic-like program. The mechanisms of MTOC sorting in mouse oocytes share similarities with properties of some cancer cells, which cluster extra centrosomes before division to ensure bipolarity and thus viability (Kwon et al., 2008). The role of HURP in these two processes is conserved, yet further investigations are needed to demonstrate that the mechanisms at play are shared.

HURP was first identified as being up-regulated in hepatocellular carcinomas (Tsou et al., 2003), solid tumors characterized by the presence of multiple centrosomes and often of clonal origin (Yao and Mishra, 2009). Therefore, it is tempting to speculate that the essential meiotic and oncogenic function of HURP originates from an evolutionarily conserved stem cell–ness.

Materials and methods

Mouse oocyte collection, culture, and microinjection

Collection of oocytes from OF1 and BL129 (Hurp+/− and Hurp−/−; Tsai et al., 2008) mice was performed as previously described (Brunet and Maro, 2007), and prophase I arrest was ensured by adding 1 µM milrinone (Reis et al., 2006) to the M2 + BSA medium. The in vitro synthesized cRNAs were injected into the oocyte cytoplasm using an Eppendorf microinjector (Femtotjet), and oocytes were further kept for 2–3 h in the prophase I arrest to allow expression of fusion proteins. Resumption of meiosis as indicated by NEBD was triggered upon transfer of oocytes into a milrinone-free M2 + BSA medium. Nocodazole and monastrol were used at 10 µM and 100 µM, respectively, in M2 + BSA. All live cultures and imaging were performed at 37°C.

Plasmid construction and in vitro transcription of synthetic RNA

The murine HURP cDNA (provided by A.P. Tsou, National Yang-Ming University, Taipei, Taiwan) was subcloned into pRN3-mCherry and pRN3-GFP. The pRN3 plasmids containing RanWT, Ran124N, RanG69L, and Histone-2BFP have been previously described (Tsuroi et al., 2004; Dumont et al., 2007). The EB3 comes from the human cDNA (gift from F. Nothias, Université Pierre et Marie Curie, Paris, France) and was subcloned into pRN3-mCherry and pRN3-GFP. In vitro synthesis of capped cRNA was performed as previously described (Verlhac et al., 2000). For GFP-HURP overexpression, cRNA were further poly-A tailed using the Poly(A) Tailing kit (Applied Biosystems) according to the manufacturer’s instructions before purification.

Immunofluorescence

Immunofluorescence on mouse oocytes was performed as previously described (Kubiak et al., 1992). Rabbit α-murine HURP (sc-98809; Santa Cruz Biotechnology, Inc.) was used at 1:50. Rat monoclonal antibody against tyrosinated α-tubulin (YL 1/2; Abcam) was used at 1:200. Human β-CREST was used at 1:60. Mouse antipericentrin (BD) was used at 1:400. After primary antibodies were added and subsequent washes were performed, oocytes were labeled with the corresponding secondary antibodies (Jackson Immunoresearch Laboratories, Inc.). Chromatin was labeled either with 5 µg/ml Hoechst (Invitrogen) or 10 µg/ml DAPI (Sigma-Aldrich).

Image acquisition of fixed oocytes was performed on a confocal microscope (SP5/AOBS; Leica) equipped with a Plan Apochromat 63×/1.4 NA oil objective. 30 confocal sections, every 0.5 µm, were taken and then displayed as a maximal Z projection.

Immunofluorescence on U2OS cells was performed as previously described (Kwon et al., 2008), and images were collected by a microscope (Axiovert; Carl Zeiss, Inc.) equipped with a spinning-disk confocal head (CSU-22; Yokogawa) and a Plan Apochromat 100×/1.4 NA oil objective using SlideBook software (Intelligent Imaging Innovations). 3D images were taken with a 0.3-µm step size and then displayed as a maximal Z projection.

Immunoblotting

Immunoblotting of mouse oocytes was performed as described previously (Tetreau et al., 2003a). 50 oocytes were loaded per lane. Protein extracts were made from livers from Hurp+/− and Hurp−/− mice. For U2OS cell immunoblotting, rabbit α-HURP (A3000853A; Bethyl Laboratories, Inc.) and mouse α-mouse α-tubulin (DM1a; Sigma-Aldrich) were used at 1:1,000 and 1:2,000 dilutions, respectively.

Live confocal microscopy

Live images were taken at 37°C in M2 + BSA using a confocal microscope with a Plan Apochromat 63×/1.4 NA objective or on a microscope (DM6000B; Leica) with a Plan Apochromat 40×/1.25 NA objective, both enclosed in a thermostatic chamber (Life Imaging Service) equipped with a charge-coupled device camera (CoolSnap HQ2; Roper Industries) coupled to a filter wheel (Sutter; Roper Industries) and a spinning disk (CSU-X1-M1; Yokogawa). MetaMorph software 7.0 (Universal Imaging) was used to collect and analyze data, and ImageJ (National Institutes of Health) was used.
to analyze and process data. 10 confocal sections, every 3 μm, were taken and then displayed as a maximal 2 projection.

For time-lapse imaging of docyccline-inducible PLK4-overexpressing U2OS cells (provided by E. Nigg, University of Basel, Basel, Switzerland), cells were transfected with control (luciferase) or HURP RNAi, synchronized with thymine for 18–20 h, released to docyccline for 10 h followed by a second thymidine treatment for an additional 18 h, and then released to medium long-term. Live-cell imaging was performed on an automated inverted microscope (TE2000E) equipped with the Perfect Focus system (Nikon) enclosed within a temperature- and CO2-controlled environment using a 20×/0.75 NA Plan Apochromat objective lens.

siRNA

Four different oligos of siRNAs (ON-TARGETplus set of four; LQ-016846-00-0002) against human HURP/DLG7 were purchased from Thermo Fisher Scientific. Oligo target sequences of siRNA for HURP are 5′-AGACUAAAGA-UUGAUAACGA-3′, 5′-GUACAAGUCCGAUGAUU-3′, 5′-GGUCU-AAACUGCGAAUA-3′, and 5′-UAAAGUGGCGUUAAGA-3′. An siRNA duplex targeting luciferase was used as a control. Cells were transfected with Lipofectamine RNAiMAX (Invitrogen) according to the manufacturer’s instructions.

Quantification analysis

The measure of MT density along the spindle axis and monastral-induced loss of HURP fluorescence intensity was performed on oocytes expressing either EB3-GFP alone or GFP-HURP together with EB3-MCherry. Background values were measured within a region of interest outside the cell. After background subtraction, two perpendicular lines were used to measure spindle length and width and were used to perform a line scan. The fluorescence intensities were divided by the maximal value of intensity coming from the corresponding spindle and, therefore, are expressed in arbitrary units. To compare intensities of spindles varying in size, we interpolated the data to identical length intervals.

The total and local MT fluorescence signal intensities (after NEBD and at NEBD + 2 h in monastral) were measured in oocytes expressing EB3-GFP and H2B-RFP (single confocal images). Background values were measured within a region of interest outside the cell and were subtracted before quantification. The total fluorescence intensity was measured inside a circle of a fixed size having the mean diameter of all oocytes. For the measure of local MT fluorescence intensity, a circle was drawn delimiting the area around chromosomes showing MT growth in wt oocytes. The mean size of this region was used to measure the corresponding fluorescence intensity in Hurp−/− oocytes, allowing us not to over- or underestimate this intensity in Hurp−/− oocytes. The same protocol was applied to measure the fluorescence intensities from monasters.

The determination of MT speed was performed using single stacks of oocytes expressing EB3-GFP using the manual tracking plugin from ImageJ. Images were taken every 250 ms over a period of 5 s. Only MT ends in focus were analyzed. Mean values for each individual MT track were processed for statistical analysis.

Normalization, correction, and fitting of the measured fluorescence intensities were performed using Excel software (Microsoft). Statistical analysis was performed using online QuickCalcs software (GraphPad Software, Inc.).

Online supplemental material

Fig. S1 shows that HURP labels MTs in the vicinity of chromosomes and is undetectable in Hurp-deficient oocytes, and its localization depends on the presence of MTs and RanGTP. Fig. S2 shows that MTOC distribution in Hurp-deficient oocytes is comparable with wt at very early steps of meiotic maturation. Fig. S3 shows that MT growth at NEBD is not affected in Hurp-deficient oocytes. Video 1 shows chromosome segregation in a wt Hurp−/− oocyte. Video 2 shows lagging chromosomes at anaphase I in a Hurp−/− oocyte. Video 3 shows dynamic localization of HURP in vivo. Video 4 shows meiotic maturation in a Hurp−/− oocyte. Video 5 shows meiotic maturation in a Hurp−/− oocyte with polar body extrusion. Video 6 shows meiotic maturation in a Hurp−/− oocyte requiring the Eg5 kinesin inhibitor monastrol leads to syntelic orientation of chromosomes and aneuploidy in mouse oocytes.

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