

Spindle assembly checkpoint insensitivity allows meiosis-II despite chromosomal defects in aged eggs

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Abstract

Chromosome segregation errors in mammalian oocyte meiosis lead to developmentally compromised aneuploid embryos and become more common with advancing maternal age. Known contributors include age-related chromosome cohesion loss and spindle assembly checkpoint (SAC) fallibility in meiosis-I. But how effective the SAC is in meiosis-II and how this might contribute to age-related aneuploidy is unknown. Here, we developed genetic and pharmacological approaches to directly address the function of the SAC in meiosis-II. We show that the SAC is insensitive in meiosis-II oocytes and that as a result misaligned chromosomes are randomly segregated. Whilst SAC ineffectiveness in meiosis-II is not age-related, it becomes most prejudicial in oocytes from older females because chromosomes that prematurely separate by age-related cohesion loss become misaligned in meiosis-II. We show that in the absence of a robust SAC in meiosis-II these age-related misaligned chromatids are missegregated and lead to aneuploidy. Our data demonstrate that the SAC fails to prevent cell division in the presence of misaligned chromosomes in oocyte meiosis-II, which explains how age-related cohesion loss can give rise to aneuploid embryos.

Keywords ageing; aneuploidy; KIF18A; oocyte meiosis; spindle assembly checkpoint

Subject Categories Molecular Biology of Disease

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Introduction

Chromosome segregation is accomplished by the spindle, a barrel-shaped molecular machine made of microtubules and associated molecules (Compton, 2000; Walczak *et al*, 2010; Dumont & Desai, 2012; Heald & Khodjakov, 2015; Bennabi *et al*, 2016). Ensuring correct chromosome segregation during cell division and thus the formation of euploid daughter cells is paramount, and the impact of

errors that cause aneuploidy is highly context-dependent. Whereas in somatic cells aneuploidy is associated with cancer (Potapova & Gorbisky, 2017; Ben-David & Amon, 2020), in germ cells aneuploidy compromises developmental potential of the embryo and thus is a major cause of reproductive failure (Hassold & Hunt, 2001; MacLennan *et al*, 2015).

Chromosome segregation in oocytes occurs over two successive meiotic divisions. Meiosis-I (M-I) comprises germinal vesicle breakdown (GVBD) and ends with the segregation of homologous chromosomes. Meiosis-II (M-II) follows thereafter without an intervening interphase, the oocyte immediately assembles the M-II spindle, and the cell cycle arrests at metaphase (Met-II) as a result of the oocyte specific cell cycle regulator Cytostatic Factor (CSF; Madgwick & Jones, 2007). M-II is ultimately completed upon fertilisation, wherein sister chromatids are separated, and embryonic development is initiated. Chromosome segregation in female meiosis is notoriously error-prone, and the likelihood of segregation errors in oocytes is increased by advancing maternal age, therein causing developmentally compromised aneuploid embryos (Hassold & Hunt, 2001; Greaney *et al*, 2018; Mihajlović & FitzHarris, 2018; Charalambous *et al*, 2022). Landmark work revealed an important clue as to how ageing leads to aneuploidy, that oocytes exhibit a progressive age-related loss of cohesin, the molecular ties that maintain chromosomal cohesion prior to segregation in anaphase (Angell, 1991, 1997; Angell *et al*, 1993; Chiang *et al*, 2010; Lister *et al*, 2010; Duncan *et al*, 2012; Merriman *et al*, 2012; Sakakibara *et al*, 2015). Progressive cohesion loss during ageing leads to premature separation of homologous chromosomes in M-I (Fig EV1), resulting in bioriented sister pairs on the spindle that separate prematurely in anaphase-I (A-I; Sakakibara *et al*, 2015). However, how premature chromosome disjunction in oocytes from ageing females ultimately leads to aneuploidy remains unclear.

A second well-established idiosyncrasy of mammalian oocytes that likely contributes to aneuploidy is that the spindle assembly checkpoint (SAC), which prevents anaphase until all chromosomes achieve proper biorientation and alignment in most cells (Musacchio & Salmon, 2007; Lara-Gonzalez *et al*, 2012; Musacchio, 2015) is weak in oocytes. This has been extensively established in oocyte

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M-I, where treatments and manipulations that cause chromosome misalignment fail to prevent anaphase and completion of M-I (Gui & Homer, 2012; Kolano *et al*, 2012; Lane *et al*, 2012; Sebestova *et al*, 2012; Kyogoku & Kitajima, 2017; Lane & Jones, 2017). However, whether the SAC can function in M-II, and how this could contribute to aneuploidy genesis, has been harder to address due to absence of tools and approaches to specifically interrogate M-II without perturbing other aspects of oogenesis. Here we have developed genetic and small molecule approaches to specifically address the SAC in M-II. Our results show a stark failure of the SAC to effectively police chromosome segregation fidelity in M-II. This failure is not intrinsically oocyte age-dependent but plays a major role in age-related aneuploidy by permitting errors arising from chromosome cohesion loss to translate into embryonic aneuploidy.

Results and Discussion

Establishment of a genetic model to interrogate the spindle assembly checkpoint in oocyte meiosis-II

We first sought to develop a mouse model with chromosome alignment defects specifically in oocyte M-II, thus allowing us to examine SAC function in M-II without having disrupted chromosome segregation in M-I. We decided to generate an oocyte-specific knockout of kinesin-8 family member, KIF18A. In somatic cells, KIF18A ensures correct biorientation and alignment of sister chromatids at the metaphase plate (Stumpff *et al*, 2008, 2012; Janssen *et al*, 2018). Given that chromosome alignment in M-II comprises biorienting sister chromatids similar to mitosis, whereas M-I biorients homologues chromosomes on a bivalent chromosome, we reasoned that KIF18A may play a major role in chromosome alignment in M-II but not M-I. To test this, we created a transgenic mouse line carrying a *Kif18a^{GFP}* allele, wherein endogenous KIF18A is N-terminally tagged with GFP, allowing us to visualise the endogenous protein, but also where the *Kif18a^{GFP}* allele harbours loxP sites to allow tissue-specific deletion (Fig EV2A). Interestingly, GFP-KIF18A was undetectable in GV-stage oocytes, and first observed only in late M-I at the spindle midzone, where it remained throughout anaphase-I/telophase-I (A-I/T-I), and was readily detected in metaphase-II (Met-II; Figs 1A and EV2B). To conditionally delete *Kif18a*, we crossed *Kif18a^{GFP}* mice with mice that express Cre-recombinase under the oocyte-specific *Zp3* promoter during oocyte growth (*Zp3-Cre*), prior to both meiotic divisions (Fig EV2C and D). *Kif18a* deletion in oocytes was confirmed by PCR (Fig EV2E) and a lack of GFP-KIF18A signal (Figs 1B and EV2F; hereafter these oocytes are referred to as KIF18A-cKO). To determine whether KIF18A knock-out affected M-I, we analysed spindles, finding no impact of *Kif18a* deletion upon spindle length (Fig EV3A). Moreover, we found no difference in chromosome alignment (Fig 1B) nor any change in inter-kinetochore distance (Fig EV3A). Furthermore, KIF18A-cKO oocytes underwent GVBD and extruded the first polar body with normal timing (Fig 1C). Importantly, *Kif18a* deletion did not increase the level of aneuploidy in the resulting Met-II eggs (Fig 1C; 2.6% in KIF18A-cKO vs. 3.6% in wild-type [WT], 5.1% in GFP knock-in [GFPⁱⁿ], and 3.8% in heterozygotes [HET]). We therefore conclude that KIF18A is dispensable for spindle formation and chromosome segregation fidelity in M-I.

Next, we assessed the requirement for KIF18A in Met-II eggs. Whilst there was no difference in Met-II spindle length and inter-kinetochore distance between KIF18A-cKO and control groups (Fig EV3B) or microtubule (MT) turnover as measured using photoactivatable-GFP assay (Fig EV3D), the frequency of misaligned chromosomes was significantly increased in KIF18A-cKO Met-II eggs (Fig 2A). In somatic cells, KIF18A depletion causes misalignment by increasing the amplitude of rapid stereotyped oscillations of bioriented sister chromatid pairs about the metaphase plate (typical frequency 1 min⁻¹; Mayr *et al*, 2007; Stumpff *et al*, 2008). In Met-II oocytes however, by performing high temporal resolution centromere live-imaging (8 s intervals), we found that chromosomes do not undergo stereotyped rapid oscillatory movements irrespective of KIF18A presence or absence (Fig 2B). However, using longer term imaging we observed three distinct chromosome behaviours that gave rise to misalignment (Fig 2C). First, the majority of misaligned chromosomes (56%) were initially aligned but became displaced from the metaphase plate ~ 18–33 min prior to anaphase onset. Second, in 22% of cases, misaligned chromosome transiently aligned, only to misalign at the opposite spindle pole. Third, the remaining 22% of chromosomes were misaligned near the same spindle pole throughout the whole movie. Notably, the chromosome misalignment persists until anaphase onset and is thus different from the misalignment that occurs transiently during Met-II spindle formation (Kouznetsova *et al*, 2019). Thus, KIF18A is required for maintaining the equatorial positioning of aligned chromosomes, exclusively in M-II, and so its conditional depletion in oocytes provides a tool for examining the impact of chromosome misalignment in Met-II eggs.

KIF18A-cKO eggs demonstrate SAC fallibility in meiosis-II

In somatic cells, a single misaligned chromosome can activate the spindle assembly checkpoint, the failsafe pathway that prevents M-phase exit, thus preventing anaphase until all chromosomes are aligned. Briefly, unattached kinetochores recruit SAC components including MAD2, which leads to the production of a Mitotic Checkpoint Complex (MCC) that prevents anaphase promoting complex (APC)-mediated Cyclin B and Securin destruction, and thus postpones anaphase until all chromosomes are correctly aligned and attached (Musacchio & Salmon, 2007). To determine the impact of chromosome misalignment on the completion of M-II we parthenogenetically activated KIF18A-cKO eggs and monitored misaligned chromosome behaviour (Fig 2D). Surprisingly, misaligned chromosomes had no impact on the timing of anaphase onset, consistent with a lack of SAC activation (Figs 2D and EV3C). Strikingly, in all cases where the chromosome was sufficiently distanced from the metaphase plate at anaphase onset to allow reliable assessment ($n = 8$), the misaligned chromosome travelled in its entirety to the nearest spindle pole (Fig 2D). Interestingly, kinetochore labelling revealed that misaligned chromosomes in KIF18A-cKO Met-II eggs were always non-bioriented sister-chromatid pairs (Fig 2A), and thus their delivery to only one daughter cell inevitably leads to aneuploidy. Notably, the frequency of anaphase lagging chromosome also increased in KIF18A-cKO Met-II eggs (Fig EV3C; 54.8% in KIF18A-cKO vs. 15.1% in WT), suggesting KIF18A has dual role in preventing chromosome segregation errors in M-II. Thus, KIF18A-cKO eggs undergo anaphase despite misaligned chromosomes following parthenogenetic activation, suggesting an absence of a robust SAC in oocyte M-II.

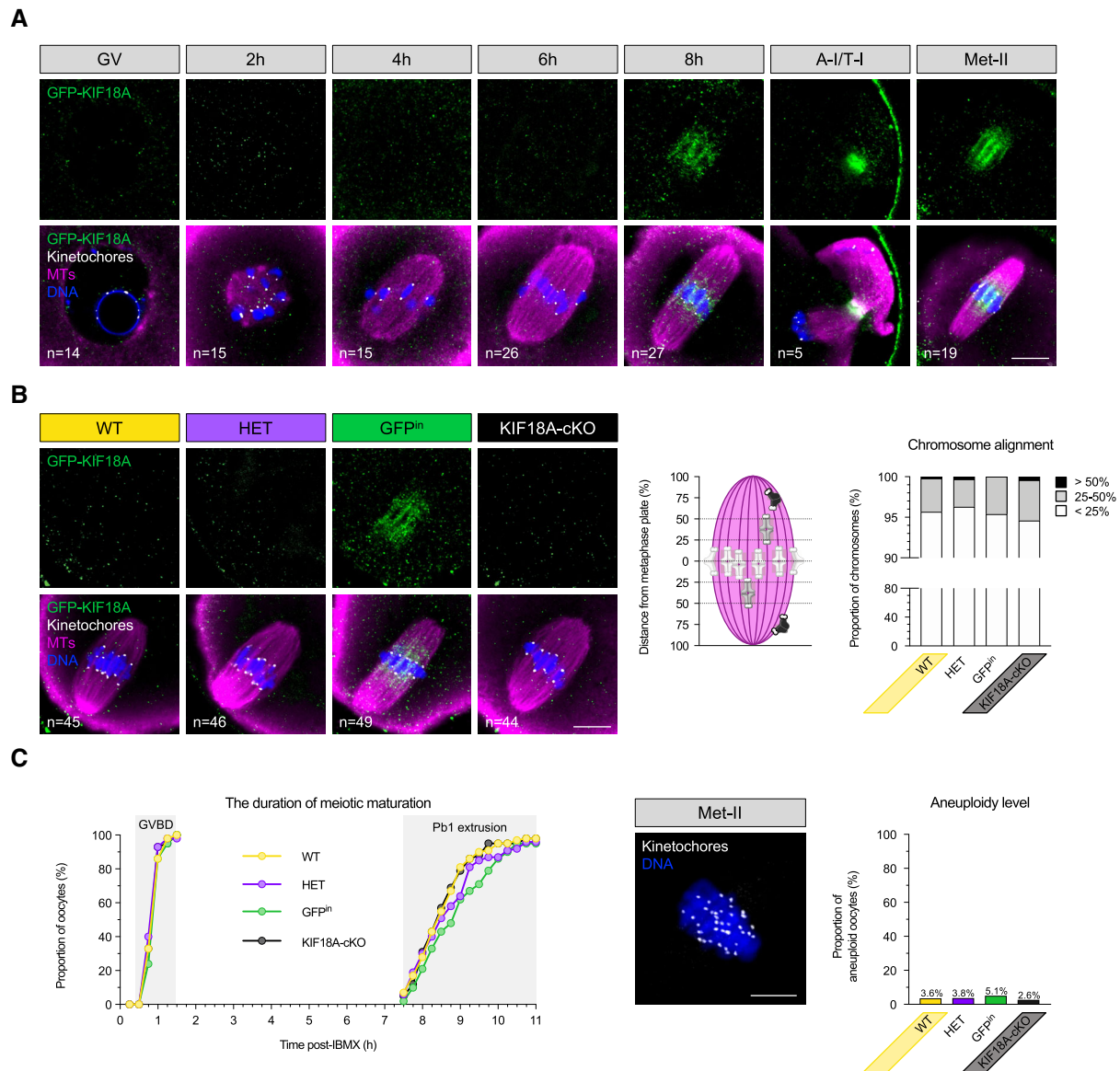


Figure 1. KIF18A removal has no impact on chromosome segregation in meiosis-I.

A, B (A) GFP-KIF18A protein is first detected in late M-I (*Kif18a*^{wt/GFP}) oocytes; GV—germinal vesicle; h—hours post-IBMX release, A-I/T-I—anaphase-I/telophase-I; Met-II—metaphase-II. (B) Late (8 h) M-I stage spindles in WT (*Kif18a*^{wt/wt}), HET (*Kif18a*^{wt/LA}), GFPⁱⁿ (*Kif18a*^{GFP/GFP}) and KIF18A-cKO (*Kif18a*^{LA/LA}) oocytes. Chart shows no difference in chromosome alignment between KIF18A-cKO and controls (χ^2 -test, $P = 0.36$). In (A and B), DNA is labelled with Hoechst33342 (blue), kinetochores with CREST (grey) and GFP-KIF18A and microtubules (MTs) with chicken-GFP (green) and β -Tubulin (magenta) antibodies, respectively. 'n' is the total number of oocytes per group. Scale bars, 10 μ m.

C Charts show no difference in meiotic maturation rate (one-way ANOVA with multiple comparisons) and aneuploidy level (χ^2 -test, $P = 0.33$) between KIF18A-cKO oocytes ($n = 38$) and controls (WT [$n = 55$], HET [$n = 52$], GFPⁱⁿ [$n = 39$]); GVBD—germinal vesicle break-down; Pb1—first polar body. An exemplar WT Met-II oocyte used in chromosome number assessment. DNA is labelled with Hoechst33342 (blue) and kinetochores with HEC1 antibody (grey). Scale bar, 5 μ m.

Source data are available online for this figure.

Small molecule-induced spindle errors fail to activate the SAC in oocyte meiosis-II

To exclude the possibility that the absence of a SAC response to misaligned chromosomes might be unique to KIF18A-cKO Met-II eggs, we also developed pharmacological approach to induce chromosome misalignment in normal (non-transgenic) Met-II eggs using

consecutive monastrol and nocodazole treatments (MonNoc eggs). Whilst neither of the drugs alone produces individually misaligned chromosomes, 200 μ M monastrol-induced spindle collapse followed by monastrol wash-out in a low concentration of nocodazole (10 nM) resulted in bipolar spindles that have misaligned pairs of sister chromatids in 35.7% of cases (Fig 3A; $n = 15/42$ eggs; 1.9 ± 0.3 misaligned chromosomes per egg). To assess the impact

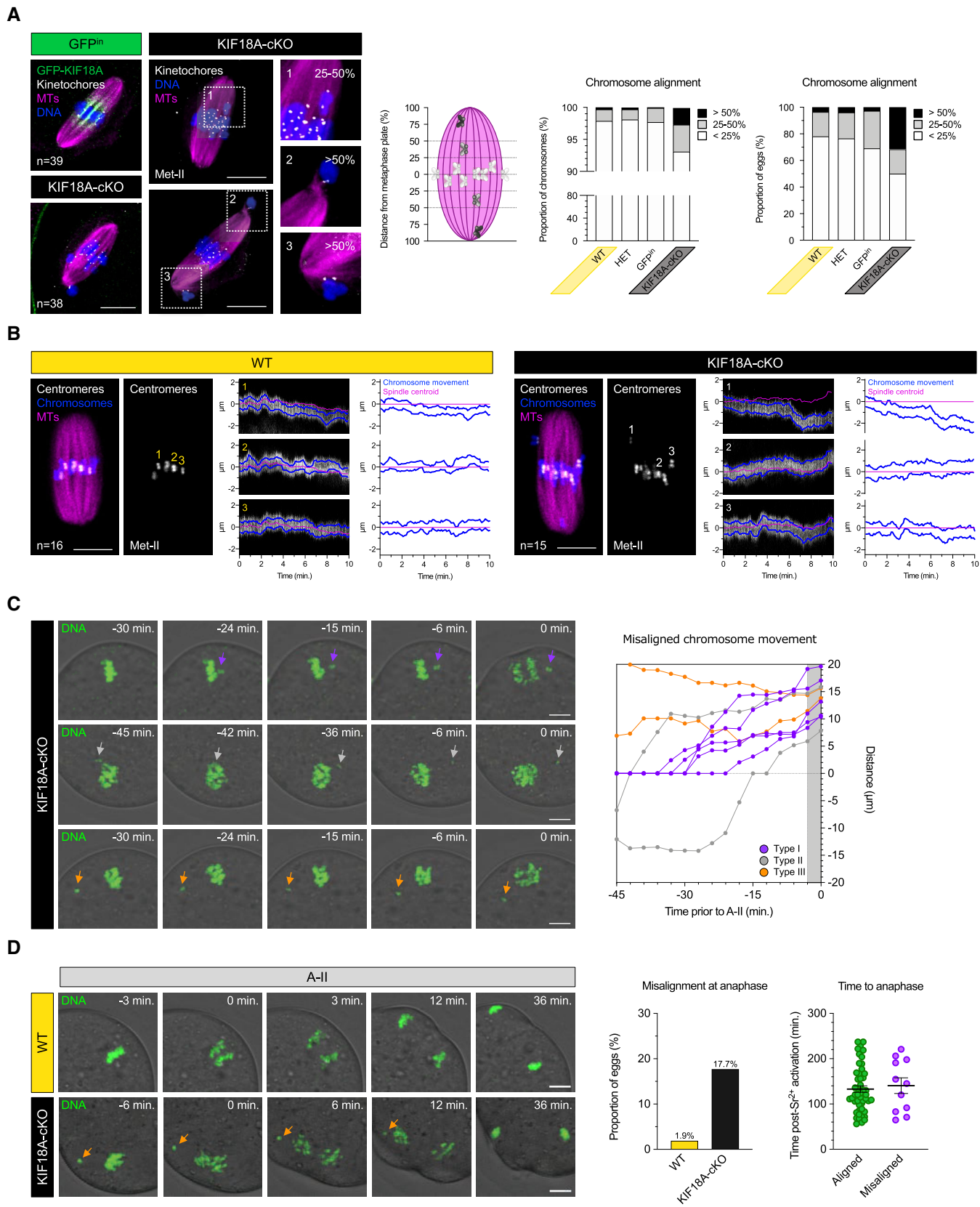


Figure 2.

Figure 2. Misaligned chromosomes fail to delay anaphase onset in KIF18A-cKO eggs.

- A Met-II-spindles in GFPⁱⁿ and KIF18A-cKO oocytes. Insets show mildly (25–50%) and severely (> 50%) misaligned chromosomes in KIF18A-cKO Met-II eggs. Note that misaligned chromosomes are pairs of sister chromatids. DNA is labelled with Hoechst33342 (blue) and GFP-KIF18A, kinetochores and microtubules (MTs) are detected with chicken-GFP (green), HEC1 (grey) and β -Tubulin (magenta) antibodies, respectively. Charts show increased chromosome misalignment in KIF18A-cKO ($n = 38$) vs. control Met-II eggs (WT [$n = 55$], HET [$n = 51$], GFPⁱⁿ [$n = 39$]; χ^2 -test, $P < 0.02$). Scale bars, 10 μ m.
- B High temporal resolution live imaging of chromosome movements in WT and KIF18A-cKO Met-II eggs. Chromosomes are visualised with H2B-RFP (blue), centromeres with Maj.Sat.-mClover (grey) and spindle with SiR-Tubulin (magenta). Kymographs show individual chromosome movement along the vertical spindle axis. Charts show centromere movements (blue) relative to spindle centroid (magenta). 'n' is the total number of Met-II eggs per group. Scale bars, 10 μ m.
- C Time-lapse confocal images depicting three types of misaligned chromosome behaviour in KIF18A-cKO Met-II eggs (left) with graphical representation of their movement (right). Top: aligned chromosome wanders of the metaphase plate shortly prior anaphase (purple; $n = 5/9$). Middle: misaligned chromosome travels to the opposite spindle pole (grey; $n = 2/9$). Bottom: chromosome remains misaligned near the same spindle pole throughout the movie (orange; $n = 2/9$). Purple, grey and orange arrows respectively indicate exemplary misaligned chromosomes for each type of movement.
- D Anaphase-II (A-II) with ostensibly normal chromosome segregation in WT (top) and severely misaligned chromosome (orange arrow) in KIF18A-cKO (bottom) eggs. Charts show significantly increased proportion of KIF18A-cKO ($n = 62$) vs. WT ($n = 53$) eggs with misaligned chromosomes (χ^2 -test, $P < 0.01$) and no anaphase delay in KIF18A-cKO eggs (unpaired t-test, $P = 0.66$). Error bars represent SEM.

Data information: In (C and D), DNA is visualised with SiR-DNA (green) and indicated time (min) is relative to the A-II onset. Scale bars, 10 μ m.

Source data are available online for this figure.

of misaligned chromosomes on meiotic exit, we parthenogenetically activated MonNoc eggs, imaging Cyclin B1-GFP destruction as a means of determining APC activation, and monitored chromosome behaviour in M-II. In MonNoc eggs, chromosomes were segregated even in eggs with clearly misaligned chromosomes at the time of anaphase onset, and misalignments had no impact on anaphase onset timing or Cyclin B1-GFP destruction dynamics (Fig 3A), thus further supporting the observation that misaligned chromosomes do not activate the SAC in eggs. Thus, misaligned chromosomes fail to prevent meiotic exit not only in KIF18A-cKO, but also in normal (non-transgenic) Met-II eggs.

We next wondered why the SAC is insensitive to misaligned chromosomes in oocyte M-II. We examined MAD2 localisation in both KIF18A-cKO and MonNoc Met-II eggs, and observed no MAD2 recruitment at the misaligned kinetochore pairs (Fig 3B). A high concentration of nocodazole (10 μ M) to completely collapse the spindle caused kinetochore-MAD2 recruitment as expected (Fig EV4A and B), and prevented Cyclin B1-GFP destruction at egg activation (Fig EV4A). Thus, a severe spindle insult can activate the SAC to prevent M-II completion suggesting the failure of misaligned chromosomes to recruit MAD2 in KIF18A-cKO or MonNoc eggs is unlikely a simple absence of checkpoint machinery. Taken together, this suggests that although the SAC can be activated in extreme circumstances, small numbers of misaligned chromatid pairs generated by two entirely distinct approaches fail to recruit MAD2, permitting chromosome segregation errors and aneuploidy.

SAC ineffectiveness renders aged eggs error-prone in M-II

Misaligned chromosomes are rare in healthy oocytes from young females (Figs 2A and D and 4A; Pan *et al*, 2008; Shomper *et al*, 2014). However, several groups have shown that the incidence of misalignment is far more substantial in Met-II eggs from aged mice (Liu & Keefe, 2008; Pan *et al*, 2008; Lister *et al*, 2010; Shomper *et al*, 2014), and indeed this mirrors the age-related increase in misalignments in human Met-II eggs (Battaglia *et al*, 1996; Volarcik *et al*, 1998). In mouse, it has been demonstrated that the misaligned chromosomes at Met-II are prematurely individualised sister chromatids (Shomper *et al*, 2014; Yun *et al*, 2014), that result from cohesion loss that causes sisters to erroneously separate in M-I (Sakakibara *et al*, 2015). This led us to hypothesise that whereas a

weak SAC might be relatively inconsequential in young Met-II eggs, SAC ineffectiveness in M-II might render eggs unable to prevent random segregation of the individual sister chromatids that emerge in M-II specifically in aged eggs.

Consistent with previous reports, we observed significant increase in the rate of chromosome misalignment in aged Met-II eggs (from mice aged 14–16 months) compared to young (2–3 months) (Fig 4A; 69.2% of aged vs. 0% of young eggs, Fisher's Exact test; $P < 0.0001$) and confirmed that the misalignments are indeed prematurely individualised sisters in all cases (Fig 4A; 7.8 ± 1.3 misaligned chromatids per egg). Strikingly, the majority of these misaligned individual chromatids had no detectable MAD2 signal at their kinetochores (57/70), whilst others had only a very weak signal (13/70) (Fig 4A). To directly determine whether aged Met-II eggs possessing misaligned individual chromatids can complete M-II, we monitored chromosome behaviour following egg activation (Fig 4B). We found that 78.6% of aged Met-II eggs ($n = 11/14$) segregated chromosomes and divided in the presence of misaligned chromatids, all of which travelled to the nearest spindle pole during anaphase. Given that individualisation of sister chromatid pairs precedes M-II spindle assembly (Yun *et al*, 2014; Sakakibara *et al*, 2015), their attachment to spindle microtubules is established independently, and so their segregation is random. Therefore, SAC insensitivity to misaligned chromosomes in M-II renders aged eggs prone to chromosome segregation errors and puts them at great risk of aneuploidy.

An integrated view of oocyte aneuploidy across M-I and M-II

Previous studies have sought to determine the proportional contribution of chromosome segregation errors in M-I and M-II to the aneuploidy that leads to infertility (Angell, 1997; Hassold & Hunt, 2001; Fragouli *et al*, 2011; Handyside, 2012; Greaney *et al*, 2018). However, our data demonstrate that errors in the two meiotic divisions are inextricably linked. Elegant imaging experiments previously showed that ageing induces premature chromosome separation in oocyte M-I, which not only predisposes oocytes to segregation errors in M-I itself (Sakakibara *et al*, 2015), but also leads to prematurely individualised sister chromatids that have a tendency to misalign on the M-II spindle (Shomper *et al*, 2014; Yun *et al*, 2014), but the fate of these individualised chromatids was

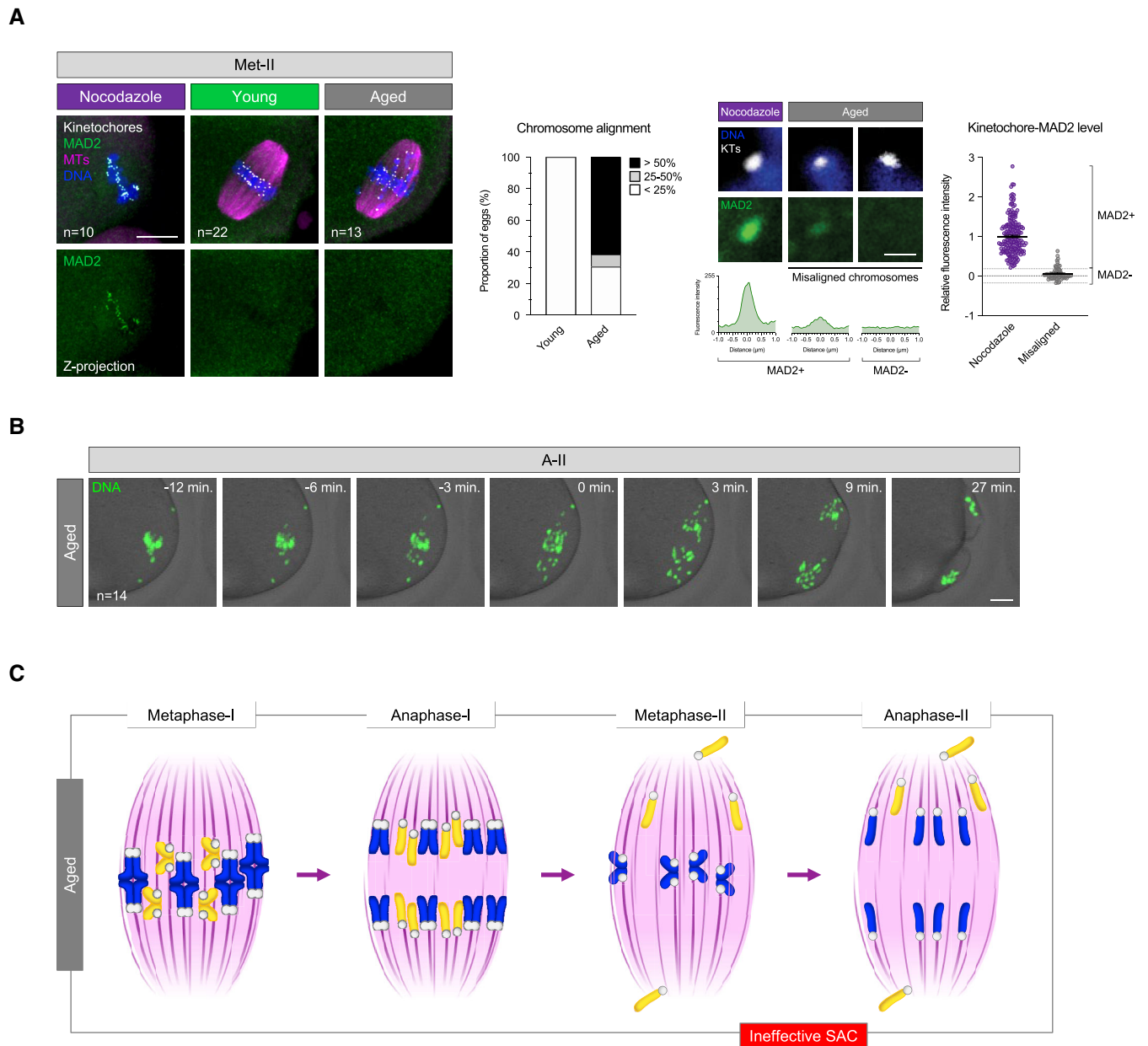


Figure 4. SAC ineffectiveness permits chromosome segregation error at egg activation.

A MAD2 immunofluorescence (green) in young and aged Met-II eggs (nocodazole-treated young Met-II eggs served as a positive control). Graph shows age-related increase in proportion of Met-II eggs with misaligned chromosomes (Fisher's Exact test, $P < 0.0001$). Exemplar weak MAD2⁺ and MAD2⁻ misaligned chromatids are shown. Graph shows significantly lower levels of kinetochores-MAD2 at misaligned chromatids in aged eggs compared to eggs treated with nocodazole alone (Mann Whitney, $P < 0.0001$). Chromosomes are labelled with Hoechst33342 (blue), kinetochores (grey) with CREST and microtubules (magenta) with β -Tubulin antibody. Scale bars, 10 μm (left) and 1 μm (right). Error bars represent SEM. Data from totals of 10, 22 and 13 oocytes across 2 experimental days, for nocodazole, young and aged groups, respectively.

B Time-lapse confocal images show aged Met-II eggs successfully completing M-II following Sr^{2+} activation despite chromosome misalignment. DNA is visualised with SIR-DNA (green) and indicated time (min) is relative to the anaphase onset. Scale bars, 10 μm . 'n' is the total number of aged Met-II eggs.

C Schematic shows a series of events during female meiosis that leads to age-related aneuploidy in embryo. Age-related cohesion loss leads to premature separation of homologue chromosomes and consequent biorientation of sister chromatids in meiosis-I. This causes premature sister chromatid disjunction during anaphase-I and the presence of individualised misaligned chromatids in Met-II eggs. SAC ineffectiveness to detect misaligned chromosomes allows anaphase-II entry and random segregation of individualised chromatids that results in embryo aneuploidy.

Source data are available online for this figure.

SAC fallibility is the norm in eggs and embryos

Whereas the SAC is nearly ubiquitous in somatic cells, and efficient to limit chromosome segregation errors, it has long been known that its function is idiosyncratic in early development. In lower vertebrates such as *X. laevis* and zebrafish, functional SAC is completely absent in oocytes and embryos and becomes acquired during development (Hara *et al*, 1980; Gerhart *et al*, 1984; Shao *et al*, 2013; Zhang *et al*, 2015). In *C. elegans* embryos, the SAC is present but its strength varies with cell size and fate (Galli & Morgan, 2016; Gerhold *et al*, 2018). In mammalian embryos, the SAC is ineffective since anaphase proceeds despite chromosome misalignment even in normal embryos (Vázquez-Diez *et al*, 2019). Similarly, in mammalian oocyte M-I, it is well established that chromosome segregation errors are not robustly prevented by the SAC (Gui & Homer, 2012; Kolano *et al*, 2012; Lane *et al*, 2012; Sebestova *et al*, 2012). This phenomenon remains poorly understood but could relate to a possible re-purposing of the checkpoint to survey DNA damage in M-I (Collins *et al*, 2015; Marangos *et al*, 2015; Lane *et al*, 2017). Our data in M-II bridge the gap, revealing that the SAC is ineffective to prevent aneuploidies throughout all of oogenesis and early embryogenesis in mammals. It is worth underlining that the causes of SAC failure may be different at different stages of development. For example, whereas anaphase takes place despite MAD2 accumulation on the kinetochores of misaligned chromosomes in M-I (Gui & Homer, 2012; Lane *et al*, 2012) and embryos (Vázquez-Diez *et al*, 2019), MAD2 is apparently not recruited in Met-II eggs. In somatic cells, MAD2 recruitment at the kinetochore occurs in response to Aurora B kinase-mediated error correction and their consequent detachment from spindle microtubules (Santaguida *et al*, 2010). Surprisingly, we found that misaligned chromosomes tend to lack stable kinetochore-MT attachments, and phosphorylated Aurora-B targets are detected at misaligned kinetochores (data not shown), suggesting that failure to recruit and activate Aurora kinase is unlikely responsible for SAC activation failure. It is possible that unstable kinetochore attachment to spindle microtubules could satisfy the SAC in eggs and provide a potential explanation for SAC inability to respond to misaligned chromosomes. Nonetheless, we find here that SAC weakness is the norm in early mammalian development. Whether this provides some as-yet unappreciated selective benefit (Ayala & Coluzzi, 2005; Sirard, 2011), or whether manipulations that might in the future reduce the toll of aneuploidy for some patients might increase reproductive success, remains to be seen.

Material and Methods

Animal models

All the animal work was performed in the animal facility of Centre Hospitalier de l'Université de Montréal (CRCHUM) and approved by the Comité Institutionnel de Protection des Animaux du CHUM (CIPA) under the protocol number IP18034GFs. All non-transgenic CD1 mice were bred at either Charles Rivers Laboratories (Crl:CD1 [ICR], Raleigh, NC, USA) or Envigo (Hsd:ICR [CD-1], Indianapolis, IN, USA), delivered to CRCHUM animal facility at 2 or 9 months of age, and used at 2–3 months (young) or 14–16 months (aged),

respectively. The conditional *Kif18a*^{GFP} knock-in mouse reporter line was generated in Dr. Laura Reinholdt's laboratory (JR#27013, 129S-*Kif18a*^{tm1.1Lgr}/Lgr, see below). Zp3-Cre^{T8} transgenic line (C57BL/6-Tg(Zp3-cre)93Kw/J, strain#: 003651) was obtained from The Jackson Laboratory (de Vries *et al*, 2000). All transgenic animals were crossed and bred at CRCHUM animal facility. Animals were maintained in individually ventilated cages (up to 5 animals/cage) at 22 ± 2°C and 40–60% humidity in 12 h light/dark cycle with lights on from 6:30 am to 6:30 pm and *ad libitum* access to food and water.

Generation of *Kif18a*^{GFP} mouse reporter line

To create the conditional GFP N-terminal tag allele, a BAC clone was obtained from TransgeneOmics (PMID: 18391959, PMID: 23417121, mouse *Kif18a*-NFLAP (ENSMUSG00000027115, RP24-307116); MCB_4891). This clone contained a *Kif18a*-NFLAP allele with an N-terminal GFP fusion and a neomycin-kanamycin resistance gene (neo) within an artificial intron flanked by loxP sites. Recombineering was used to retrieve the genomic interval from the BAC clone into a gene-targeting vector for mESCs. The original neo cassette was replaced with a neo cassette flanked by frt sites to create a new single 5' loxP site in the artificial intron and 3' loxP was added to intron 2. The resulting targeting vector was electroporated into CJ7 mESCs (RRID:CVCL_C316, 129S1/SvImJ background), targeted clones were identified by loss of allele assays and confirmed by Southern blotting. Two correctly targeted clones were microinjected into C57BL/6J (RRID:IMSR_JAX:000664) embryos. Because the C57BL/6J genetic background was previously found to reduce penetrance of the *Kif18a* mutant infertility phenotype (PMID: 25824710), the breeding scheme for the chimeras was designed to minimise C57BL/6J. Chimeras were first bred to C57BL/6J females to identify males with germline transmission (GLT) of the targeted mESC genome by coat colour. One confirmed GLT male was then bred to 129S1/SvImJ (RRID:IMSR_JAX:002448) to establish the breeding colony. After two backcrosses to 129S1/SvImJ, N2 heterozygous males were bred to a FLP recombinase-expressing strain from a 129 genetic background (RRID:IMSR_JAX:003946) to remove the frt-neo cassette from the targeted allele. The resulting “GFPin” allele is shown in Fig EV2A and this allele was subsequently maintained through intercrossing and mice from this colony were shipped to CRCHUM for these studies. Sperm from a homozygous male were cryopreserved at The Jackson Laboratory as JR#27013, 129S-*Kif18a*^{tm1.1Lgr}/Lgr. The functionality of the conditional allele was tested by crossing 129S-*Kif18a*^{tm1.1Lgr} males to Sox2-cre (RRID:IMSR_JAX:008454) females to excise the translational start site, as well as the coding portion of exon 2, and resulting heterozygous *Kif18a*^{KO} mice were intercrossed. Heterozygous *Kif18a*^{KO} mice were also crossed to heterozygous *Kif18a*^{gcd2} mice. Homozygous KO mice and compound heterozygous *Kif18a*^{KO}/*Kif18a*^{gcd2} mice exhibited hypogonadism and infertility as described previously, confirming loss of function (PMID: 25824710, PMID: 16822657).

Genotyping

PCR genotyping was performed using MyTaqTM Extract-PCR Kit (Meridian Bioscience, Cat# BIO-21126) and MyTaqTM HS Red Mix (Meridian Bioscience, Cat# BIO-25047) according to manufacturer's

instructions. For animal genotyping, genomic DNA was extracted from the ear clips. For gamete genotyping, 10 GV-stage oocytes retrieved from a single female were pooled together to extract DNA. To detect *Kif18a* alleles, two PCR assays were used with following primers: (i) KIF18A-F: 5'-CCAGTAATACATTTGCTAGGTGGAT-3' and KIF18A-R: 5'-GTAATGGCAATGCCTGGGA -3'; (ii) KIF18A-F: 5'-CCAGTAATACATTTGCTAGGTGGAT-3' and GFP-R: 5'-TCCTTGAAGAAGATGGTGCG-3'. The initial DNA denaturation was performed at 95°C for 3 min, followed by 35 cycles of amplification at 95°C for 15 s, 59°C for 15 s, 72°C for 1 min (for assay 1) or 20 s (for assay 2). In assay 1, the WT allele yields a 715 bp product, the GFPⁱⁿ allele a 1,990 bp product and KIF18A-cKO allele a 627 bp product. In assay 2, GFPⁱⁿ allele yields 682 bp product. Myogenin-specific primers were used as an internal positive control (MYOG-F: 5'-CCTGTAACCCAGGCAGAGAG-3' and MYOG-R: 5'-AGCCTAGGTAGGGACTAAGGG-3'; yields 217 bp product). The following pair of primers was used to detect Cre transgene: Cre-F: 5'-GCGGTCTGGCAGTAAAACTATC-3' and Cre-R: 5'-GTGAAACAGCATTGCTGTCACTT-3'; 100 bp product). The initial DNA denaturation was done at 95°C for 3 min, followed by 35 cycles of amplification at 95°C for 15 s, 57°C for 15 s, 72°C for 20 s. Interleukin-2 precursor-specific primers were used as an internal positive control (IL-2-F: 5'-CTAGGCCACAGAATTGAAAGATCT-3' and IL-2-R: 5'-GTAGGTGGAAATCTAGCATCATCC-3'; 324 bp product).

Fertility trial

Sexually mature females (2 months of age) of three different genotypes: *Zp3-Cre^{Tg/0}; Kif18a^{wt/wt}* (WT oocytes), *Zp3-Cre^{Tg/0}; Kif18a^{wt/GFP}* (HET oocytes) and *Zp3-Cre^{Tg/0}; Kif18a^{GFP/GFP}* (KIF18A-cKO oocytes) were continuously mated with wild-type B6D2F1/J (strain#: 100006) male mice of confirmed fertility (2 months of age; Jackson Laboratories) until 5 litters were born. Cages were checked daily for pups.

Oocytes and egg collection and handling

Fully grown, germinal vesicle (GV)-stage oocytes were collected from 2 to 3 months (young) and 14–16 months (aged) old non-transgenic CD1 or transgenic mice, 44–46 h following pregnant mare's serum gonadotrophin (PMSG; Genway Biotech, Cat. # GWB-2AE30A) hormonal stimulation (5 and 10 IU of PMSG/mouse, respectively). Oocyte handling outside the incubator was done in M2 media (Sigma; Cat# M7167) and in the presence of 200 μM 3-isobutyl-1-methylxanthine (IBMX; Sigma, Cat# I5879) to maintain them at GV stage. The surrounding cumulus cells have been removed and the oocytes washed through M16 media (Cat# M7292, Sigma) with 200 μM IBMX and left in the incubator at 37°C and 5% CO₂ for approximately 1 h to recover prior any further manipulation. To obtain M-II oocytes, females were stimulated with 5 IU of human chorionic gonadotropin (hCG; Sigma, Cat# CG10-1VL) 48 h following PMSG and sacrificed 14 h post-hCG. Cumulus-oocyte complexes were released from the oviduct into M2 media containing 0.3 mg/ml hyaluronidase (Sigma; Cat# H4272) to dissociate the oocytes from cumulus cells. MII-oocytes were then washed through M16 media and left in the incubator at 37°C and 5% CO₂ prior any further manipulation.

Drug treatments

To disassemble the spindle in Figs 4A and EV4, eggs were exposed to 10 μM nocodazole (Cat# 487928, Sigma) either for 20 min prior fixation or during the entire live imaging. To induce chromosome misalignment in Fig 3, ovulated Met-II oocytes (14–16 h post-hCG stimulation) were exposed to consecutive 200 μM monastrol (Cat# 475879, Sigma) and 10 nM nocodazole treatments (each lasting 2 h), and live imaged in the presence of 10 nM nocodazole.

Microinjection

GV-stage oocytes and ovulated eggs were microinjected in M2 media with or without 200 μM IBMX, using a picopump (World Precision Instruments), intracellular electrometer (Warner instruments), and micromanipulators (Narishige), mounted on a Leica DMI4000 inverted microscope. The mRNAs used for injections were *in vitro* synthesised using mMessage mMachine kits (T3 and T7, Cat# AM1348 and AM1344, Ambion), followed by poly-adenylation using Poly(A) Tailing kit (Cat# AM1350, Invitrogen) according to the manufacturer's instructions. The following linearised plasmids were used as templates: pRN3: H2B-RFP (a gift from Alex McDougall, Observatoire Océanologique de Villefranche sur Mer, Villefranche sur Mer, France; Prodon *et al*, 2010), pTALYM3B15 (Maj. Sat.-mClover; Addgene #47878, Miyanari *et al*, 2013) and pCMX: Cyclin B1-GFP (Addgene #26061, Hagting *et al*, 1999). Injected oocytes were left in the incubator for at least 2 h to allow the time for fluorescent protein expression.

Egg activation and live-cell confocal imaging

For live-imaging experiments in Figs 2C and D, and 4B, and EV3C, eggs were incubated in M16 media with 1 μM SiR-DNA alone (Cat# CY-SC007, Cytoskeleton Inc.) for 2 h to label the chromosomes or, in Figs 3A and EV4A, in combination with 1xSPY555-Tubulin (Cat# CY-SC203, Cytoskeleton Inc.) to label the spindle. For live imaging experiment in Fig 2B, eggs were incubated in 300 nM SiR-Tubulin (Cat# CY-SC002, Cytoskeleton Inc.) for 1 h to label the spindle. Eggs were parthenogenetically activated using 10 mM SrCl₂ (Cat# 439665, Sigma) in Ca²⁺-free M2 media and placed in an on-stage incubating chamber (at 37°C) mounted on a Leica SP8 confocal microscope. Z-stacks time-lapse images were acquired with 20× air objective (0.75 NA) using 2 μm optical section and step size at 3 min time-intervals for experiments in Figs 2C and D, and 4B, and EV3C and 5 min intervals in Figs 3A and EV4A. For Fig 2B, single optical sections were taken at 8 s time-intervals. For the experiments in Figs 2B and 3A, and EV4A, the oocytes were illuminated with 488, 552, 638 nm light (using respectively, 0.5, 0.15 and 0.5% power in Fig 2B, and 1, 1 and 0.5% in Figs 3A and EV4A) and in Figs 2C and D, and 4B, and EV3C with 638 nm laser (0.5% power) and fluorescence was detected with a HyD detector.

Fixation and immunofluorescence

Oocytes were fixed using 2% paraformaldehyde (PFA; Cat# P6148, Sigma) in phosphate-buffered saline (PBS; Cat# OXBR0014G, ThermoFisher) for 20 min, followed by permeabilisation step in 0.25% Triton-X 100 (Cat# T9284, Sigma) in PBS for 10 min at ambient

temperature. Blocking was done in 3% BSA (Cat# A7906, Sigma) in PBS for 1 h or overnight at 4°C and was followed by incubation step in antibodies for 1 h at 37°C. Primary antibodies were used at following dilutions: chicken anti-GFP (1:500; Cat# GFP-1020, Aves Labs), mouse anti- β -Tubulin (1:1,000; Cat# T4026, Sigma), rabbit anti-PRC1 (1:200; Cat# 15617-1-AP, Proteintech), rabbit anti-MAD2 (1:300; Cat# 924601, BioLegend), rabbit anti-HEC1 (1:500; a gift from Robert Benezra, Memorial Sloan Kettering Cancer Center, New York, USA) and CREST (1:100; a gift from Marvin J. Fritzler, University of Calgary, Calgary, Canada). Secondary antibodies were used at following dilutions: goat anti-mouse Alexa 488 (1:1,000; Cat# A-11029, ThermoFisher), goat anti-mouse Alexa 568 (1:1,000; Cat# A-11031, ThermoFisher), goat anti-mouse Alexa 633 (1:500; Cat# A-21050, ThermoFisher), goat anti-rabbit Alexa 568 (1:1,000; Cat# A-11011, ThermoFisher), donkey anti-rabbit Alexa 647 (1:1,000; Cat# A-31573, ThermoFisher), donkey anti-chicken Alexa 488 (1:500; Cat# 703-545-155, Jackson ImmunoResearch), donkey anti-chicken Alexa 647 (1:500; Cat# 703-605-155, Jackson ImmunoResearch) and goat anti-human Alexa 546 (1:1,000; Cat# A-21089, ThermoFisher). Chromosomes were labelled with Hoechst 33342 (20 μ g/ml; H1399, Invitrogen) for 20 min at ambient temperature. Imaging of fixed samples was carried out on a Leica SP8 confocal microscope using 63xoil-immersion objective (1.4 NA) using 0.9 μ m optical section and step size.

Microtubule turnover measurement

To measure the microtubule turnover in Fig EV3D, PAGFP-Tubulin was photoactivated with 405 nm laser within a defined rectangular shaped region of interest positioned in between the metaphase plate and one of the spindle poles and live-imaged at 30 s. time intervals for 5 min to assess the dissipation of the fluorescent signal. The fluorescence decay curves and MT half-live ($T_{1/2}$) were obtained by plotting the fluorescence intensity decay obtained for each spindle against time and fitted into a double exponential curve $f(t) = A \times \exp(-k_1t) + B \times \exp(-k_2t)$ using the cftool in MATLAB, wherein t represents the time; A is the non-stable interpolar MT population, B is the stable KT-MT population and k_1 and k_2 are the decay rates of the two MT populations, respectively. The half-life ($T_{1/2}$) of KT-MT population was calculated as $\ln 2/k_2$. Each measurement was corrected for photobleaching, that was measured by imaging oocytes in the presence of 10 μ M MT-stabilising agent, Taxol (Cat# T7402, Sigma), where the MT-turnover rate was minimal.

Image analysis and statistics

All image analysis was performed in ImageJ/Fiji (Schindelin et al, 2012) and Imaris (version 9.6.0, Bitplane). To assess spindle length, interkinetochore distance and chromosome alignment in Figs 1B and 2A, and 4A, and EV3A and B, spatial coordinates of the spindle poles and individual kinetochores were manually determined based on β -Tubulin and CREST/HEC1 immunofluorescence signal, and distances calculated using Pythagoras' theorem. The extent of chromosome alignment was determined by calculating chromosome distance from each spindle pole (expressed as a relative distance from the spindle midpoint) and subdivided into three categories: aligned (< 25%), mildly misaligned (25–50%) and severely misaligned (> 50%). For Fig 1C, the germinal vesicle

breakdown and polar body extrusion timing was determined by inspecting oocytes each 15 min within first 1.5 and 7–11 h following IBMX washout and the aneuploidy level was assessed by counting the number of kinetochores in fixed Met-II eggs. For Fig 2B, chromosomes oscillatory movements were measured in the metaphase of meiosis-II (in the period 12–18 h post-IBMX release) based on centromeric Maj. Sat.-mClover signal and normalised relative to spindle centroid (calculated in ImageJ/Fiji using SiR-Tubulin fluorescence). For Figs 2C and 3A, values of chromosome metaphase displacement were obtained by measuring chromosome distance from the metaphase plate. For Fig 2D, misalignment at anaphase was determined based on the presence of chromosome optically separate from the metaphase plate at the time frame prior to anaphase onset. For Figs 2D and 3A, and EV3C, time to anaphase was calculated from the moment of Sr²⁺ addition to the first time-frame when two main body masses of segregating chromosomes appeared optically separate from each other. For Figs 3B and 4A, and EV4B, kinetochore-MAD2 immunofluorescent signal intensity was quantified by calculating the ratio between the average fluorescent signal (mean grey value) of individual kinetochores and cytoplasmic background (20 kinetochores per egg). For Fig EV2B, PRC1 signal and GFP-KIF18A colocalisation assessment was done by drawing a line across the metaphase plate and plotting the fluorescent signal intensity profile in Fiji. For Fig EV3C, anaphase-II chromosome was considered lagging if it was optically separate from the main body of segregating chromosomes when the spindle was oriented to be planar to the field of view. For Figs 3A and EV4A, CyclinB1-GFP destruction was assessed by measuring the average cytoplasmic fluorescent signal intensity at the Z-plane with the largest oocyte membrane perimeter over time. Statistical analysis was performed in GraphPad Prism8 using χ^2 -test, unpaired or paired t -test, Fisher's Exact test, Mann-Whitney test and one-way ANOVA where appropriate. Error bars represent standard error of the mean (SEM).

Data availability

Our study includes no data deposited in public repositories.

Expanded View for this article is available [online](#).

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Author contributions

Aleksandar I Mihajlović: Conceptualization; data curation; formal analysis; validation; investigation; visualization; writing – original draft; writing –

review and editing. **Candice Byers:** Writing – original draft. **Laura Reinholdt:** Writing – original draft. **Greg FitzHarris:** Conceptualization; supervision; funding acquisition; validation; writing – original draft; writing – review and editing.

Disclosure and competing interests statement

The authors declare that they have no conflict of interest.

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