Endomembranes promote chromosome missegregation by ensheathing misaligned chromosomes

Nuria Ferrandiz1, Laura Downie1, Georgina P. Starling1, and Stephen J. Royle1

Errors in mitosis that cause chromosome missegregation lead to aneuploidy and micronucleus formation, which are associated with cancer. Accurate segregation requires the alignment of all chromosomes by the mitotic spindle at the metaphase plate, and any misalignment must be corrected before anaphase is triggered. The spindle is situated in a membrane-free “exclusion zone”; beyond this zone, endomembranes (mainly endoplasmic reticulum) are densely packed. We investigated what happens to misaligned chromosomes localized beyond the exclusion zone. Here we show that such chromosomes become ensheathed in multiple layers of endomembranes. Chromosome ensheathing delays mitosis and increases the frequency of chromosome missegregation and micronucleus formation. We use an induced organelle relocalization strategy in live cells to show that clearance of endomembranes allows for the rescue of chromosomes that were destined for missegregation. Our findings indicate that endomembranes promote the missegregation of misaligned chromosomes that are outside the exclusion zone and therefore constitute a risk factor for aneuploidy.

Introduction

Accurate chromosome segregation during mitosis is essential to prevent aneuploidy, a cellular state of abnormal chromosome number (Duijf and Benezra, 2013). Errors in mitosis that lead to aneuploidy can occur via different mechanisms. These mechanisms include mitotic spindle abnormalities (Ghadimi et al., 2000), incorrect kinetochore-microtubule attachments (Cimini et al., 2001), dysfunction of the spindle assembly checkpoint (Kalitsis et al., 2000), defects in cohesion (Daum et al., 2011), and failure of cytokinesis (Fujinawa et al., 2005). Some of these error mechanisms result in the missegregation of whole chromosomes, a process termed chromosomal instability (CIN). The majority of solid tumors are aneuploid, with higher rates of CIN, and so understanding the mechanisms of chromosome missegregation is an important goal of cancer cell biology. In addition, chromosome missegregation is associated with micronucleus formation, which is linked to genomic rearrangements that may drive tumor progression (Crasta et al., 2012; Ly et al., 2017; Liu et al., 2018).

While the mitotic spindle has logically been the focus of efforts to understand chromosome missegregation, there has been less attention on other features of mitotic cells such as intracellular membranes. In eukaryotic cells, entry into mitosis constitutes a large-scale reorganization of intracellular membranes. The nuclear envelope (NE) breaks down, while the ER and Golgi apparatus disperse to varying extents (Hepler and Wolniak, 1984; Warren, 1993). These organelle remnants—collectively termed “endomembranes”—are localized toward the cell periphery, while the mitotic spindle itself is situated in an “exclusion zone” that is largely free of membranes and organelles (Bajer, 1957; Porter and Machado, 1960; Nixon et al., 2017). The endomembranes beyond the exclusion zone are densely packed, although the details of their ultrastructure vary between cell lines (Puhka et al., 2007; Lu et al., 2009, 2011; Puhka et al., 2012; Champion et al., 2017). This arrangement means that, although mitosis is open in mammalian cells, the spindle operates within a partially closed system. Several lines of evidence suggest that endomembranes must be cleared from the exclusion zone for the mitotic spindle to function normally (Vedrenne et al., 2005; Schlaitz et al., 2013; Champion et al., 2019; Kumar et al., 2019; Merta et al., 2021). In addition, it is thought that this arrangement is required to concentrate factors needed for spindle formation (Schweizer et al., 2018).
This study was prompted by a simple question: What happens to misaligned chromosomes that find themselves beyond the exclusion zone? We show that such chromosomes become “enveloped” in multiple layers of endomembranes. Chromosome enveloping delays mitosis and increases the frequency of chromosome missegregation and subsequent formation of micronuclei. Using an induced organelle relocalization strategy, we demonstrate that clearance of endomembranes allows the rescue of chromosomes that were destined for missegregation. Our findings indicate that endomembranes are a risk factor for CIN if the misaligned chromosomes go beyond the exclusion zone boundary during mitosis.

**Results**

**Misaligned chromosomes outside the exclusion zone are ensheathed in endomembranes**

During mitosis, the spindle apparatus is situated in a membrane-free exclusion zone. Outside the exclusion zone, the ER and NE—collectively called endomembranes—surround the mitotic spindle. We investigated the organization of endomembranes in mitotic cells using light microscopy and EM. First, we carried out live-cell imaging of mitotic RPE-1 cells that stably coexpress GFP-Sec61β and Histone H3.2-mCherry with SiR-Tubulin. These images revealed a mitotic spindle-sized exclusion zone from which GFP-Sec16β signal was absent (Fig. 1A). Second, serial block face scanning electron microscopy (SBF-SEM) of mitotic RPE-1 cells showed that the ellipsoid exclusion zone is largely devoid of endomembranes, including mitochondria and other organelles. Outside the exclusion zone, endomembranes are tightly packed, and the border between these two regions is clearly delineated and could be segmented (Fig. 1B).

Misaligned chromosomes are those that fail to attach or lose their attachment to the mitotic spindle. What happens to misaligned chromosomes that end up among the endomembranes beyond the exclusion zone? HeLa cells have high rates of chromosome misalignment, and live-cell imaging showed that misaligned chromosomes could be situated beyond the exclusion zone (Fig. 1C). Reconstruction of SBF-SEM data from HeLa cells showed that three to four layers of endomembranes ensheathe the chromosomes beyond the exclusion zone (Fig. 1D and Videos 1 and 2). We use the term ensheathed to describe how these chromosomes are surrounded by endomembranes but not fully enclosed in any one layer, as though in a vesicle.

To study chromosome ensheathing in diploid cell lines, we needed to artificially increase the frequency of misaligned chromosomes in mitosis. Our main model was RPE-1 cells pretreated with 150 nM GSK923295, a centromere protein E (CENP-E) inhibitor (Wood et al., 2010), before washing out the drug for 1 h (Fig. 2A). In parallel, we also used a system of targeted Y-chromosome spindle detachment in DLD-1 cells (Ly et al., 2017; Fig. S1). Using live-cell imaging in both cell types, we observed that misaligned chromosomes beyond the exclusion zone are submerged in endomembranes (Figs. 2B and S1E). Next, we used an image analysis method to determine the location of kinetochores in 3D space and map these positions relative to the exclusion zone boundary (see Materials and methods; Figs. 2, C and D; and Fig. S1, F and G). Kinetochores of chromosomes that were not aligned at the metaphase plate therefore fell into two categories: those that were surrounded by
GFP-Sec61β signal, termed ensheathed, and those that were not, termed free (Fig. 2, B and D). Spatial analysis revealed that the kinetochores of ensheathed chromosomes were beyond the exclusion zone, whereas kinetochores of free chromosomes lay at the boundary in RPE-1 cells (Fig. 2 D). In DLD-1 cells, the distinction was even more clear, with the kinetochores of free chromosomes positioned inside the exclusion zone S1F. The exclusion zone therefore approximately defines chromosome misalignment, with those chromosomes beyond the exclusion zone likely to be ensheathed by endomembranes. However, imaging GFP-Sec61β was required to verify that a chromosome was fully ensheathed.

We again used SBF-SEM to observe how chromosomes beyond the exclusion zone interact with endomembranes in RPE-1 cells. Cells observed by fluorescence microscopy to have at least one ensheathed chromosome were selected for 3D EM analysis (Fig. 2 E). Segmentation of these datasets confirmed that the chromosome was fully beyond the exclusion zone boundary (Fig. 2 F and Video 3) and was ensheathed in several layers of endomembranes (Fig. 2 E). The observation of ensheathed chromosomes raised immediate questions about their fate and whether ensheathing leads to aberrant mitosis.
ensheathed chromosomes delay mitotic progression

To determine the impact of ensheathed chromosomes on cell division, we first analyzed mitotic progression in RPE-1 cells stably expressing GFP-Sec61β with induction of ensheathed chromosomes using GSK923295 pretreatment. Cells that had at least one ensheathed chromosome showed prolonged mitosis (median NE breakdown [NEB]-to-anaphase timing of 66 min compared with 27 min in GSK923295 pretreated cells in which all chromosomes were aligned). The time to align the majority of chromosomes (NEB-to-metaphase) was delayed for cells with either a free or an ensheathed chromosome, but cells with an ensheathed chromosome had an additional delay to progress to anaphase (Fig. 3 A). Given these delays, we next confirmed that the spindle assembly checkpoint was active in these cells. The amount of Mad2 and Bub1 detected by immunofluorescence at CENP-C-positive kinetochores of free or ensheathed chromosomes was similar and was four-fold higher than at kinetochores of aligned chromosomes (Fig. 3, B and C; and Fig. S2, A and B, for DLD-1 cells). Using live-cell imaging, we found that GFP-Mad2 was recruited to kinetochores of ensheathed chromosomes (Fig. 3, D and E; and Video 4). Semiautomated 4D tracking of chromosomes allowed us to monitor their GFP-Mad2 status over time, relative to anaphase onset. These data revealed that GFP-Mad2 is lost from ensheathed chromosomes with similar kinetics to the signals at misaligned chromosomes that successfully congress to the metaphase plate (Fig. 3 E).

The failure of ensheathed chromosomes to congress is likely due to a lack of microtubule attachment, suggesting that endomembranes inhibit chromosome–microtubule interactions. We confirmed that ensheathed chromosomes have no stable end-on kinetochore-microtubule attachments by detecting colocalization of kinastrin, a marker for stable end-on attachment (Dunsch et al., 2011), with kinetochores of aligned and misaligned chromosomes (Fig. S3, A–C). Live-cell imaging of RPE-1 cells stably coexpressing Histone H3.2-mCherry and GFP-Sec61β, stained with SiR-Tubulin, showed that ensheathed chromosomes that failed to congress had no detectable microtubule contacts; free chromosomes that had microtubule contacts could be rescued and aligned at the metaphase plate, albeit after a delay (Fig. S3, D and E).

These results suggest that ensheathed chromosomes hinder mitotic progression in a spindle assembly checkpoint–dependent manner. Lack of microtubule contact is sensed by the spindle assembly checkpoint, but ultimately, the checkpoint is extinguished in the absence of congression after a long delay. The cells then proceed to anaphase, resulting in missegregation of the ensheathed chromosome.

ensheathed chromosomes promote formation of micronuclei

To understand the fate of cells with an ensheathed chromosome, we next examined mitosis in control or GSK923295-pretreated RPE-1 cells stably expressing GFP-Sec61β using live-cell spinning disc microscopy (Fig. 4 A). In cells with an ensheathed chromosome, we observed the long delay in mitosis relative to control cells, and that mitosis was often resolved by missegregation and formation of a micronucleus (Figs. 4 A and S2 C for DLD-1 cells). These experiments suggested that ensheathed chromosomes are potentially a precursor to micronuclei. We therefore followed the fate of mitotic cells by long-term live-cell imaging to understand the likelihood of mitotic outcomes. Our sample of cells pretreated with GSK923295 included the three metaphase classes: aligned (25.8%), free (5.4%), and ensheathed (65.6%). The most frequent fate of cells with an ensheathed chromosome was micronucleus formation (39%). Of the 47 cells that formed a micronucleus after division in the dataset, 46 were from the ensheathed class (Fig. 4 B). This promotion of micronucleus formation was significant in cells with an ensheathed chromosome compared to free (P = 1.3 × 10−3, Fisher’s exact test). A smaller proportion of cells with an ensheathed chromosome exited mitosis normally, albeit with a delay (34%), with the remainder showing other defects or death (20% or 8%). Cells pretreated with GSK923295, that had aligned all their chromosomes, had similar fates to parental and control cells (Fig. 4 B; and Videos 5 and 6). These fate-mapping experiments suggest that ensheathing of chromosomes by endomembranes promotes the formation of micronuclei.

Micronuclei formed from ensheathed chromosomes have a disrupted NE

Micronuclei can undergo a collapse of their NE, which manifests as ER tubules invading the micronuclear space (Hatch et al., 2013). We therefore asked if micronuclei that formed from ensheathed chromosomes were similarly defective. Using confocal imaging of RPE-1 cells stably coexpressing GFP-Sec61β and either mCherry-BAF or LBR-mCherry that were fixed 8 h after washout of GSK923295 to examine micronucleus integrity, we found that the majority of micronuclei have ER inside the micronucleus (Fig. 5). The fluorescence of GFP-Sec61β was higher at the micronucleus compared with the main nucleus (Fig. 5 B). Moreover, the levels of either mCherry-BAF or LBR-mCherry were correlated with GFP-Sec61β. To confirm that these micronuclei had disrupted NEs, we stained for H3K27Ac, a modification to Histone H3 that is removed by exposure to the cytoplasm (Mammel et al., 2021). Intact micronuclei had H3K27Ac signals similar to those of the corresponding main nucleus, whereas in micronuclei that were disrupted, the signal was lost (Fig. 5 A). The ratio of H3K27Ac signal at the micronucleus compared with the main nucleus was anticorrelated with the ratios of GFP-Sec61β, mCherry-BAF, and LBR-mCherry (Fig. 5 B). Since the majority of micronuclei formed after pretreatment of RPE-1 cells with GSK923295 are derived from ensheathed chromosomes (Fig. 4 B), these data suggest that the ensheathing process may contribute to the formation of defective micronuclear envelope. However, due to the low rates of missegregation of free chromosomes, it was not possible to conclude whether disruption was specific to chromosome ensheathing.

Induced relocalization of ER enables the rescue of ensheathed chromosomes

Does ensheathing of misaligned chromosomes cause chromosome missegregation? To answer this question, we sought a way to clear the mitotic ER and test whether this enabled subsequent rescue of misaligned chromosomes to the metaphase plate. To
clear the mitotic ER, we used an induced relocalization strategy (Fig. 6 A). Induced relocalization of small organelles has been demonstrated for Golgi, intracellular nanovesicles, and endosomes, typically using heterodimerization of FKBP-rapamycin-FRB with the FKBP domain fused to the organelle and the FRB domain at the mitochondria (Dunlop et al., 2017; Hirst et al., 2015; Larocque et al., 2020; van Bergeijk et al., 2015). We reasoned that a large organellar network, such as the ER, may be cleared by inducing its relocalization to the cell boundary. Our strategy therefore comprised an ER-resident hook (FKBP-GFP-Sec61β) and a plasma membrane anchor (stargazin-mCherry-FRB) with application of rapamycin predicted to induce the

Figure 3. Impact of ensheathed chromosomes on cell division. (A) Mitotic timing of RPE-1 cells. Cumulative frequencies for NEB to metaphase (NEB-Meta) and metaphase to anaphase (Meta-Ana) are shown. RPE-1 stably expressing GFP-Sec61β were treated with 150 nM GSK923295 for 3 h before washout. Three classes of metaphase were seen: all chromosomes aligned (Aligned, n = 29), cells with one or more free chromosomes (Free, n = 11), and cells with one or more ensheathed chromosome (Ensheathed, n = 107). Timing of untreated parental (Parental, n = 69) and stable RPE-1 (Control, n = 52) cells is also shown. Inset in Meta-Ana shows same data on an expanded time scale. Comparison of NEB-Meta and Meta-Ana timing distributions for ensheathed vs. control, P = 1.9 × 10^{-57} and 7.8 × 10^{-23}, Kolmogorov–Smirnov test. (B) Micrographs of immunofluorescence experiments to detect Bub1 or Mad2 (SAC, red) at kinetochores (CENP-C, blue) in cells stably expressing GFP-Sec61β (green); DAPI-stained DNA is shown in gray. Scale bars, 10 µm; 2 µm (insets). (C) Quantification of Bub1 and Mad2 immunofluorescence at kinetochores marked by CENP-C. Ensheathed chromosomes were classified using the GFP-Sec61β signal. Dots represent kinetochores, boxes show IQR, bar represents the median, and whiskers show 9th and 91st percentiles (Bub1: nA = 132, nF = 30, nE = 37; Mad2: nA = 103, nF = 20, nE = 31). (D) Stills from live-cell imaging experiments to track Mad2 levels at kinetochores of ensheathed chromosomes. A GSK923295-pretreated RPE-1 cell is shown, stably coexpressing GFP-Mad2 (green) and mCherry-Sec61β (red); DNA is stained using SiR-DNA (blue). Time relative to anaphase is shown in minutes. Insets show 2× zoom of the indicated ROI. Scale bars, 10 µm; 2 µm (insets). (E) Quantification of live Mad2 imaging experiments. Kaplan–Meier plot to show congression times of the last misaligned chromosome to align. Measurement of mCherry-Sec61β (mean ± SD) and GFP-Mad2 is shown for the misaligned that congressed and those that were missegregated (misseg). A linear regression fit with 95% confidence intervals is shown for GFP-Mad2. All plots are shown in time (minutes) relative to anaphase onset. Total cells with misaligned chromosomes, n = 72; cells where all chromosomes congressed, n = 56; and where there was missegregation, n = 16.
relocalization of ER to the plasma membrane (Fig. 6 A). HCT116 cells were used for these experiments, as they are near diploid and easy to transfect and showed a fate and mitotic response to GSK923925 pretreatment similar to those of RPE-1 (Fig. S4).

We found that the clearance of ER in mitotic cells with this strategy was efficient, occurring in 89.2% of HCT116 cells expressing the system after treatment with 200 nM rapamycin. Onset was variable, with a median time to maximum clearance of 15 min (interquartile range [IQR], 12–24 min; Fig. 6 B). Importantly, induced relocalization of FKBP-GFP-Sec61β to the plasma membrane represented the clearance of ER and not the extraction of the protein. First, immunostaining of two other endogenous ER-resident proteins, KDEL and calnexin, also showed relocalization to the plasma membrane (Fig. 6 C). Second, SBF-SEM imaging allowed us to observe the relocalization of ER to the plasma membrane (Fig. 6 D). Here, the expansion of the exclusion zone and the direct attachment of hundreds of ER tubules to the plasma membrane could be unambiguously visualized.

We next tested whether ER clearance could be used as an intervention in cells with ensheathed chromosomes. To do this, HCT116 cells expressing FKBP-GFP-Sec61β and stargazin-mCherry-FRB, pretreated with 150 nM GSK923295 to induce ensheathed chromosomes, were imaged as 200 nM rapamycin was applied to induce clearance of the ER. In control cells where no rapamycin was applied, the cells were arrested in mitosis for prolonged periods. In cells where the ER had been cleared, congression of the ensheathed chromosome was clearly seen after clearance had occurred (Fig. 7 A and Video 7). We used automated image analysis to track the 3D position of the misaligned chromosome over time, in an unbiased manner (Fig. 7, B–C). Congression of the ensheathed chromosome within 80 min was seen in 86.7% of cells with induced ER clearance. In control cells, the majority (66.7%) were unable to resolve the ensheathed chromosome in the same time (Fig. 7, A–C). These data suggest that ER clearance is an effective intervention in cells with ensheathed chromosomes and points to a causal role for endomembranes in chromosome missegregation.

Discussion

This study demonstrates that misaligned chromosomes located beyond the exclusion zone are liable to become ensheathed by endomembranes. The fate of cells with ensheathed chromosomes is biased toward missegregation, aneuploidy, and micronucleus formation. We showed that if the ER was cleared by induced relocalization in live mitotic cells, these chromosomes could be rescued by the mitotic spindle, an intervention which suggests that chromosome ensheathing by endomembranes is a risk factor for chromosome missegregation and subsequent aneuploidy.

Chromosomes can become misaligned during mitosis for a number of reasons, but we show here that those that transit out of the exclusion zone become ensheathed by endomembranes. We demonstrated this with four different cell models: RPE-1 or HCT116 cells pretreated with a CENP-E inhibitor, DLD-1 cells with targeted disconnection of the Y-chromosome, and HeLa cells with spontaneously arising misaligned chromosomes. In each case, misaligned chromosomes beyond the exclusion zone typically became ensheathed in endomembranes. Although the
morphology of mitotic endomembranes varies between cell lines (Puhka et al., 2007; Lu et al., 2009, 2011; Puhka et al., 2012; Champion et al., 2017), all ensheathed chromosomes were draped in several layers of endomembranes. We use the term ensheathed to describe how these chromosomes are surrounded by endomembranes but not fully enclosed in any one layer as though in a vesicle. The ensheathing membrane follows the contours of the chromosome closely. Our SBF-SEM analysis did not uncover any obvious electron-dense connections between the ensheathed chromosome and its surrounding membranes, although a previous report indicated that exogenous DNA clusters may physically interact with mitotic ER (Wang et al., 2016).

A major finding of our work is that ensheathing promotes missegregation and micronucleus formation. Our 3D EM images of ensheathed chromosomes show that microtubules face a difficult task to negotiate several layers of endomembranes to make the contact between kinetochore and spindle that is necessary for rescue and alignment. In cases where contact is made, endomembranes are also likely to impair the congression of the chromosome, as suggested by a recent study in which excess ER was shown to slow chromosome motions (Merta et al., 2021). Since endomembranes are a risk factor for missegregation, their precise organization—e.g., the sheet-to-tubule ratio of the ER—may influence the likelihood for missegregation (Champion et al., 2017). The lack of attachment is sufficient to prolong spindle assembly checkpoint signaling and delay mitosis. Ultimately, the cells progress to anaphase and missegregate, likely due to checkpoint exhaustion after prolonged metaphase (Uetake and Sluder, 2010; Yang et al., 2008). Whatever the mechanism, the role of endomembranes in promoting missegregation may be important for tumor progression. It is possible that in tumor cells that are aneuploid, endomembranes may contribute to the higher rates of CIN observed (Funk et al., 2016; Nicholson and Cimini, 2015). In non-transformed cells, misaligned chromosomes that arise spontaneously are more often of the free class, suggesting that the ensheathing mechanism described here is most relevant in a cancer context.

The fate of cells with ensheathed chromosomes was biased toward missegregation and formation of micronuclei. Interestingly, a previous study found that artificially tethering endomembranes to aligned chromosomes within the exclusion zone caused mitotic errors, although the outcome was dependent on at what stage tethering was induced (Champion et al., 2019). Tethering before mitotic entry resulted in segregation errors and multilobed nuclei, whereas tethering during metaphase had little consequence. Although conceptually similar, the ensheathing process reported here is a natural consequence of a misaligned chromosome becoming entangled in endomembranes. Key differences include the position of the ensheathed chromosome, the lack of microtubule attachments, no direct membrane-chromosome tethering, and multiple vs. single endomembrane layers; these likely explain the different observed mitotic phenomena. We found that the micronuclei that result from ensheathed chromosomes had disrupted envelopes 8 h after release from CENP-E inhibition. Rupture of micronuclei has been shown to lead to DNA damage and activation of innate immune and cell invasion pathways (Ly et al., 2017; Hatch et al., 2013; Mammel et al., 2021; Bakhoum et al., 2018). The presence
of ER in the micronuclear space of disrupted micronuclei indicates that ensheathing may increase the likelihood of rupture. We speculate that this may occur by endomembranes physically interfering with envelope reformation at the micronucleus, although it is possible that ER is present in the micronuclear space as a consequence, rather than a cause, of disruption.

Mitosis in human cells is open, yet we have known for >60 yr that the spindle exists in a membrane-free ellipsoid exclusion zone (Bajer, 1957; Porter and Machado, 1960; Nixon et al., 2017). It seems intuitive that the spindle must operate in a membrane-free area to avoid errors, but recent work suggests that the exclusion zone is actively maintained and that this arrangement is important for concentrating factors for spindle assembly (Schweizer et al., 2015) or for maintenance of spindle structure (Kumar et al., 2019; Schlaitz et al., 2013). We found that ER clearance, via an induced relocalization strategy, could be used as an intervention to improve the outcome for mitotic cells with ensheathed chromosomes. Induced relocalization of small

Figure 6. Inducible relocalization of ER in mitotic cells. (A) Schematic diagram of the ER clearance procedure. Rapamycin induces the heterodimerization of the ER-resident FKBP-GFP-Sec61β and the plasma-membrane localized Stargazin-mCherry-FRB. (B) Cumulative histogram showing the time to detection of ER clearance. An automated segmentation procedure was used to monitor ER localization in mitotic cells. The time at which the largest decrease in ER localization occurred was taken (n = 35–37, see Materials and methods). Random occurrence is shown for comparison. The median (IQR) ER clearance time in rapamycin-treated cells was 15 (12–24) min; rapamycin is applied after the first frame (T = 0). (C) Induced relocalization of FKBP-GFP-Sec61β to the plasma membrane causes ER clearance. Typical immunofluorescence micrographs of mitotic HCT116 cells pretreated with GSK923295, expressing FKBP-GFP-Sec61β (green) and Stargazin-mCherry-FRB (blue), treated or not with rapamycin (200 nM). Cells were stained for ER markers KDEL or Calnexin as indicated (red), DNA was stained with DAPI (gray). Insets are 2× expansions of the ROI shown. Scale bars, 10 µm; 1 µm (insets). (D) SBF-SEM imaging of control or ER-cleared (rapamycin) mitotic HCT116 cells. A single slice is shown with segmentation of ER (green), plasma membrane (yellow), mitochondria (blue), and chromosomes (red). Scale bars, 5 µm; 1 µm (insets). Insets are 2× expansions of the indicated ROI shown without segmentation; green arrowheads indicate ER attachment to the plasma membrane.
organelles has previously been demonstrated (Dunlop et al., 2017; Hirst et al., 2015; Larocque et al., 2020; van Bergeijk et al., 2015), but the movement of a large organellar network by similar means had not been attempted previously. Surprisingly, ER clearance in mitotic cells was efficient, although it was much slower than the relocalization of intracellular nano-vesicles, taking tens of minutes rather than tens of seconds (Larocque et al., 2020). We speculate that the efficiency of clearance is due to cooperativity of relocalization, since the FKBP-GFP-Sec61β molecules are dispersed in the ER, which is interconnected. These experiments were important to show that ensheathing was causal for chromosome missegregation. We note that this method has many future applications: to selectively perturb mitotic structures, at defined times, during cell division. For example, ER clearance and concomitant expansion of the exclusion zone is an ideal manipulation to probe the function of this enigmatic cellular region.

**Materials and methods**

**Molecular biology**

The following plasmids were gifts, available from Addgene, or from previous work as indicated: Histone H3.2-mCherry (A. Bowman, University of Warwick), pAc-GFP-C1-Sec61β (#15108; Addgene), psPAX2 (#12260; Addgene), pMD2.G (#12259; Addgene), pWPT-GFP (#12255; Addgene), Stargazin-GFP-LOVpep (#80406; Addgene), LBR pEGFP-N1 (#61996; Addgene), EGFP-BAF (#101772; Addgene), pMito-mCherry-FRB (#59352; Addgene), Histone H2B-mCherry (Cheeseman et al., 2013), and pFKBP-GFP-C1 (Clarke and Royle, 2018).

To generate a plasmid to express mCherry-Sec61β, EcoRI-BglIII digestion product of pAc-GFP-C1-Sec61β was ligated into pmCherry-C1 vector (made by substituting mCherry for EGFP in pEGFP-C1 [Clontech] by AgeI-XhoI digestion). LBR-mCherry was made by amplifying the LBR insert from LBR in pEGFP-N2 using (5’-AAGCTTGTTACCATGCAAGTAGAATTTGC-3’ and 5’-TCGAGCGGCCGCTGGTAACCGAGGC-3’).
GAGGGATCCGTGATCGATTATGAAATATACGG-3′) and ligating into pmCherry-N1 using KpnI and BamHI. The mCherry-BAF construct was amplified from EGFP-BAF using oligonucleotides (5′-AAGCTTAGCTCTGACAACCTCCTAAAAGC-3′ and 5′-TCGAGGAGTTCGAAAGGATCC-3′) and inserted into pmCherry-CI using BglII and HindIII.

For lentiviral transduction plasmids, constructs for expression (mCherry-BAF, GFP-Mad2, mCherry-Sec61β) were cloned into pWPT-GFP using MluI-SalI sites or MluI-BstBI for LBR-mCherry. Plasmids for ER clearance were generated as follows. For FKBP-GFP-Sec61β, a BglII-EcoRI fragment from pAc-GFP-Cl-Sec61β was ligated into pFKBP-GFP-CI. Stargazin-mCherry-FRB construct was made by PCR of Stargazin encoding region from RPE-1 cells was achieved by lentiviral transduction of cells stably expressing GFP-Sec61β and mCherry, mCherry-BAF, or LBR-mCherry with GFP-Sec61β and validated using a combination of Western blot, FACS, and single cell clones were validated by fluorescence microscopy. Note that the transgenic expression of GFP-Sec61β in RPE-1 cells was achieved by lentiviral transduction of cells stably expressing GFP-Sec61β. For stable expression of GFP-Mad2 with mCherry-Sec61β, dual lentivirus transduction was used. Individual cells positive for GFP and mCherry signal were sorted by FACS, and single cell clones were validated by fluorescence microscopy.

Cell biology

HCT116 (CCL-247; ATCC) and HEK293T (CRL-11268; ATCC) cells were maintained in DMEM supplemented with 10% FBS and 100 U ml⁻¹ penicillin/streptomycin. DLD-1-WT and DLD-1-C-H3 (Ly et al., 2017) cell lines were gifts from Don Cleveland (University of California San Diego, San Diego, CA). These cell lines and their derivatives were maintained in DMEM supplemented with 10% Tetra-Free FBS (D2-118, SLS), 2 mM L-glutamine, 100 U ml⁻¹ penicillin/streptomycin, and 100 μg ml⁻¹ hygromycin. RPE-1 (Horizon Discovery) and derived cell lines were maintained in DMEM/F-12 Ham supplemented with 10% FBS, 2 mM L-glutamine, 100 U ml⁻¹ penicillin/streptomycin, and 0.26% sodium bicarbonate (NaHCO₃). All cell lines were kept in a humidified incubator at 37°C and 5% CO₂. Cells were routinely tested for mycoplasma contamination by a PCR-based method.

RPE-1 GFP-Sec61β stable cell line was generated by FugeneHD (Promega) transfection of pAc-GFP-C1-Sec61β. DLD-1-WT mCherry-Sec61β and DLD-1-C-H3 mCherry-Sec61β stable cell lines were generated by Genecure (Merck Millipore) transfection of mCherry-Sec61β into the respective parental lines. Individual clones were isolated by G418 treatment (500 μg ml⁻¹) and validated using a combination of Western blot, FACS, and fluorescence microscopy. Stable coexpression of Histone H3.2-mCherry, mCherry-BAF, or LBR-mCherry with GFP-Sec61β in RPE-1 cells was achieved by lentiviral transduction of cells stably expressing GFP-Sec61β. For stable expression of GFP-Mad2 with mCherry-Sec61β, dual lentivirus transduction was used. Individual cells positive for GFP and mCherry signal were sorted by FACS, and single cell clones were validated by fluorescence microscopy.

For immunofluorescence, cells were fixed at room temperature using PFA solution (3% formaldehyde and 4% sucrose in PBS) for 15 min and permeabilized at room temperature in 0.5% (vol/vol) Triton X-100 in PBS for 10 min. Cells were blocked in 3% BSA in PBS for 60 min at room temperature. Cells were then incubated for 60 min at room temperature with primary antibody dilutions prepared in 3% BSA in PBS as follows: mouse anti-Bub1 (ab54893, 1:500; Abcam); mouse anti-Mad2 (sc-65492, 1:200; Santa Cruz); rabbit anti-calnexin (ab22595, 1:200; Abcam); guinea pig anti-CENP-C (PD030, 1:2,000; Medical and Biological Labs Company); rabbit anti-H3K27ac (ab4729, 1:100; Invitrogen); and rabbit anti-KDEL (PA1-013, 1:200; Invitrogen). After three PBS washes, coverslips were rinsed and mounted with Vectashield containing DAPI (Vector Laboratories) and sealed. In cases where GFP signal required amplification, cells were incubated with GFP-booster (Alexa Fluor 488, 1:200; Chromotek) at the secondary antibody step. Where amplification of mCherry was required, mouse anti-mCherry (IC5; ab125096, 1:500; Abcam) was used with Alexa Fluor 568-conjugated secondary antibody.

For FISH of DLD-1 WT and DLD-1-C-H3 cells, the degron expression construct was used with Alexa Fluor 647-conjugated secondary antibody. For FISH of DLD-1 WT and DLD-1-C-H3 cells, the degron system was induced, and cells were synchronized by double thymidine (2.5 mM) treatment. Samples were fixed in Carnoy’s fixative (3:1 vol/vol methanol/glacial acetic acid) for 5 min at
room temperature, rinsed in fixative before addition of fresh fixative, and incubated for a further 10 min. Samples were rinsed in distilled water before FISH probe denaturation and hybridization following the manufacturer’s protocol (Xcyteng Centromere Enumeration Probe, XCE Y green, D-0824-050-FI; MetaSystems Probes). To dye chromosomes or microtubules in fixed- or live-cell imaging, cells were incubated for 30 min with 0.5 μM SiR-DNA or SiR-Tubulin (Spirochrome), respectively.

Biochemistry
For Western blot, cells were harvested and lysates were prepared by sonication of cells in UTB buffer (8 M urea, 50 mM Tris, and 150 mM β-mercaptoethanol). Lysates were incubated on ice for 30 min, clarified in a benchtop centrifuge (20,800 g) for 15 min at 4°C, boiled in Laemmli buffer for 10 min, and resolved on a precast 4–15% polyacrylamide gel (Bio-Rad). Proteins were transferred to nitrocellulose using a Trans-Blot Turbo Transfer System (Bio-Rad). Primary antibodies were diluted in 4% BSA in PBS and used as follows: rabbit anti-Sec61β (PA3-015, 1:1,000; Invitrogen); HRP-conjugated mouse anti-β-actin (sc-47778, 1:20,000; Santa Cruz); rabbit anti-mCherry (ab183628, 1:2,000; Abcam); anti-GAPDH (G9545, 1:5,000; Sigma-Aldrich); rabbit anti-CENP-A (2186, 1:100; Cell Signaling); mouse anti-BAF (A-11, 1:500; Santa Cruz); and mouse anti-LBR (SAB1400151, 1:500; Sigma-Aldrich). Secondary antibodies of anti-mouse, anti-rabbit, and anti-rat IgG HRP conjugates were prepared in 5% milk in PBS. For detection, enhanced chemiluminescence detection reagent (GE Healthcare) and manual exposure of Hyperfilm (GE Healthcare) was performed.

Microscopy
For fixed-cell imaging experiments, a Personal DeltaVision microscope system (Applied Precision), based on an IX-71 microscope body (Olympus) was used with a CoolSNAP HQ2 interline charge-coupled device camera (Photometrics) and a 60× oil-immersion 1.42-NA oil PlanApo N objective and equipped with Precision Control microscope incubator, Tokai Hit stage top incubator, and Applied Precision motorized xyz stage. Illumination was via a Lumencor SPECTRA X light engine (DAPI, 395/25; GFP, 470/24; mCherry, 575/25; CY-S, 640/30), dichroics (quad: reflection 381-401:464-492:531-556:619-644; transmission 409-456:500-523:564-611:652-700; GFP/mCh: reflection 464-492:561-590; transmission 500-553:598-617) and filter sets (DAPI: excitation 387/11, emission 457/50; GFP: excitation 470/40, emission 525/50; TRITC: excitation 575/25, emission 597/45; mCherry: excitation 572/28, emission 632/60; and CY-S: excitation 640/14, emission 685/40). Image capture was by softWoRx 5.1.1 (Applied Precision). Images were deconvolved using softWoRx 3.0 with the following settings: conservative ratio, 15 cycles, and high noise filtering.

For live-cell imaging, cells were plated onto fluorodishes (WPI) and imaged in complete medium in an incubated chamber at 37°C and 5% CO2. Most live-cell imaging was done using a Nikon CSU-W1 spinning disc confocal system; SoRa upgrade (Yokogawa) was used with either a Nikon 100×, 1.49 NA, oil, CFI SR HP Apo TIRF or 63×, 1.40 NA, oil, CFI Plan Apo objective with optional 2.3× intermediate magnification and 95B Prime camera (Photometrics). The system has a CSU-W1 (Yokogawa) spinning disk unit with 50 μm and SoRa disks (SoRa disk used), Nikon Perfect Focus autofocus, Okolab microscope incubator, Nikon motorized xy stage, and Nikon 200-μm z-piezo. Excitation was via 405-, 488-, 561-, and 638-nm lasers with 405/488/561/640-nm dichroic and blue, 446/60; green, 525/50; red, 600/52; and FRed, 708/75 emission filters. Acquisition and image capture was via NIH Elements (Nikon).

For mitotic progression and fate experiments, the DeltaVision system described above was used. For live-cell imaging of HeLa cells, a spinning disc confocal system (UltraView VoX; PerkinElmer) with a 60×, 1.40 NA, oil, Plan Apo VC objective (Nikon) was used. Images were captured using an ORCA-R2 digital charge-coupled device camera (Hamamatsu) after excitation with 488- and 561-nm lasers and 405/488/561/640-nm dichroic and 525/50, 615/70 filter sets. Images were captured using Volocity 6.3.1. All microscopy data were stored in an OMERO database in native file formats.

SFB-SEM
To prepare samples for SFB-SEM, RPE-1 GFP-Sec61β cells on gridded dishes were first incubated with 150 nM GSK923295 (Selleckchem) for 3 h to induce misaligned chromosomes, before release of cells from treatment and incubation for ~30 min with 0.5 μM SiR-DNA (Spirochrome) to visualize DNA. HeLa cells on gridded dishes were not treated and were not stained. Using live-cell light microscopy, cells with an ensheathed chromosome were selected for SFB-SEM. Fluorescent and bright-field images of the selected cell were captured, and the coordinate position was recorded. Cells were washed twice with phosphate buffer (PB) before fixing (2.5% glutaraldehyde, 2% paraformaldehyde, 0.1% tannic acid [low molecular weight] in 0.1 M phosphate buffer, pH 7.4) for 1 h at room temperature. Samples were washed three times with PB and then postfixed in 2% reduced osmium (equal volume of 4% OsO4 prepared in water and 3% potassium ferrocyanide in 0.1 M PB solution) for 1 h at room temperature, followed by a further three washes with PB. Cells were then incubated for 5 min at room temperature in 1% (wt/vol) thiocarbohydrazide solution, followed by three PB washes. A second osmium staining step was included, incubating cells in a 2% OsO4 solution prepared in water for 30 min at room temperature, followed by three washes with PB. Cells were then incubated in 1% uranyl acetate solution at 4°C overnight. This was followed by a further three washes with PB. Walton’s lead aspartate was prepared adding 66 mg lead nitrate (TAAB) to 9 ml 0.03 M aspartic acid solution at pH 4.5, and then adjusting to final volume of 10 ml with 0.03 M aspartic acid solution and to pH 5.5 (pH adjustments with KOH). Cells were incubated in Walton’s lead aspartate for 30 min at room temperature and then washed three times in PB. Samples were dehydrated in an ethanol dilution series (30, 50, 70, 90, and 100% ethanol, 5-min incubation in each solution) on ice, and then incubated for a further 10 min in 100% ethanol at room temperature. Finally, samples were embedded in an agar resin (AGAR 100 RI140; Agar Scientific).

SFB-SEM data were segmented using Microscopy Image Browser v2.60, and the resulting 3D model was visualized in

Ferrandiz et al.
Chromosome ensheathing promotes missegregation
Data analysis
Kinetochore position analysis was in two parts. First, the positions of kinetochores and spindle poles in hyperstacks were manually mapped using Cell Counter in Fiji. The kinetochore point sets were classified into three categories: those aligned at the metaphase plate and those that were misaligned, with the latter group subdivided into kinetochores of chromosomes that were ensheathed and those that were not (free). Second, the ER channel of the hyperstack was segmented in Fiji to delineate the exclusion zone. Next, the Cell Counter XML files and their respective binarized ER stacks were read by program written in Igor Pro (WaveMetrics). To analyze the position of points relative to the exclusion zone in each cell, the ratio of two Euclidean distances was calculated (see Eq. 1). Where $C$ is the centroid of all aligned kinetochores, $P_i$ is the position of a kinetochore and $Q_i$ is the point on the path from $C$ through $P_i$ where the exclusion zone/ER boundary intersects with the path.

$$\frac{CF_i}{CQ_i}.$$ (1)

The ratio of these two distances gave a measure of how deep the point was placed inside or outside the exclusion zone (0 being on the boundary and 1 being far outside of the exclusion zone as from the centroid to the boundary, on a log2 scale).

For analysis of live-cell GFP-Mad2 and mCherry-Sec61β imaging, a semiautomated 4D tracking procedure was used. Briefly, the DNA channel from these videos was used for segmentation of chromosomes and metaphase plate as discrete 3D objects over time. The centroid-to-centroid distance was found for each chromosome relative to the plate (congression was taken as the merging of chromosome and plate objects), and the time of anaphase onset was determined. Fluorescence signals were taken from each chromosome object using a 3-pixel expansion of the region of interest (ROI). For mCherry-Sec61β, the mean voxel density was used. For GFP-Mad2, the maximum pixel intensity at each z position was taken from the expanded ROI and averaged per time point; this method gave a more accurate measure of Mad2 recruitment than the mean voxel density. Signals from each channel are expressed as a ratio of chromosome to plate. Mad2 signals were grouped by whether the chromosome congressed, and then measurements from all chromosomes relative to anaphase were used to fit a line by linear regression. Only the last chromosome to congress (or not) was analyzed per cell. Data processing was via Fiji/ImageJ followed by analysis in Igor Pro.

Automated kinetochore-kinastrin colocalization was using a script that located the 3D position of kinetochores (CENP-C) and kinastrin puncta from thresholded images using 3D Object Counter in Fiji. These positions were loaded into Igor, and the Euclidean distance to the nearest kinastrin punctum from each kinetochore was found. ER clearance experiments were quantified using two automated procedures. First, ER, DNA, and plasma membrane were segmented separately, the plasma membrane segments were used to define the cell, and the total area of segmented ER within this region was measured for all z-positions over time using a Fiji macro. Data were read by Igor, and the ER volume over time was calculated. ER clearance manifested as a rapid decrease in ER volume, but the onset was variable. The derivative of ER volume over time was used to find the point of rapid decrease, and this point was used to define the time to ER clearance. Random fluctuations in otherwise constant ER volume over time also resulted in minima that occurred randomly. This process was modeled and plotted for comparison with the control group, where no clearance was seen. Second, the segmented DNA was classified into misaligned chromosome and main chromosome mass by a user blind to the conditions of the experiment. 3D coordinates of these two groups were fed into Igor, where the centroids and boundaries of the chromosome and main chromosome mass were defined. The closest Euclidean distance between the centroid of the chromosome and edge of the main chromosome mass was used as the distance. Misalignment, shown as a colorscale, is this distance normalized to the starting distance. Figures were made with Fiji, R, or Igor Pro and assembled using Adobe Illustrator.

Statistical testing
Comparison of mitotic timing distributions was done using a Kolmogorov-Smirnov test ($P$ values are $P_{n,[\varepsilon]}$). The effect of presence of ensheathed chromosome on mitotic fate (frequency of micronucleus formation) was examined using Fisher’s exact test with no correction. Chromosome congression times were not normally distributed, and so the effect of ER clearance was determined using Wilcoxon rank test. Exact $P$ values for all tests are quoted, rather than using arbitrary levels of significance.

Online supplemental material
Fig. S1 shows ensheathed chromosomes in DLD-1 cells. Fig. S2 shows spindle assembly checkpoint activation and micronucleus formation in DLD-1 cells. Fig. S3 shows lack of microtubule attachments of ensheathed chromosomes. Fig. S4 shows mitotic timing and fate of HCT116 cells pretreated with CENP-E inhibitor. Fig. S5 shows stable transgene expression in RPE1 cells. Video 1 shows a 3D reconstruction of an ensheathed chromosome in a HeLa cell. Video 2 shows a 3D reconstruction of an ensheathed chromosome in a HeLa cell. Video 3 shows a 3D reconstruction of an ensheathed chromosome in an RPE-1 cell. Video 4 shows an example of GFP-Mad2 at an ensheathed chromosome. Video 5 shows an example of mitotic outcome of a cell with aligned chromosomes. Video 6 shows an example of mitotic outcome of a cell with an ensheathed chromosome. Video 7 shows an example of ER clearance and subsequent rescue of an ensheathed chromosome.

Data availability
All code used in the manuscript is available at https://github.com/quantixed/Misseg.

Acknowledgments
We thank Claire Mitchell and Laura Cooper from the Computing and Advanced Microscopy Unit (CAMDU) for their help and
support. Faye Nixon, Alison Beckett and Ian Prior at the Liverpool Biomedical EM Unit provided SBF-SEM imaging. We are grateful to Richard Bayliss, Vishakha Karnawat, Jonathan Millar, Alonso Pardal and James Shelford for feedback on the project and manuscript.

This work was supported by a Pioneer Award and Programme Award from Cancer Research UK (C25425/A24167 and C25425/A27718). L. Downie was supported by a studentship from BBSRC Midlands Integrative Biosciences Training Partnership (BB/M01166X/1).

The authors declare no competing financial interests.

Author contributions: Conceptualization—N. Ferrandiz; investigation and resources—N. Ferrandiz and L. Downie; formal analysis—L. Downie and S.J. Royle; funding acquisition and software—S.J. Royle; supervision and writing—original draft—M. Ferrandiz and S.J. Royle; visualization—G.P. Starling and S.J. Royle; writing—review & editing—N. Ferrandiz, L. Downie and S.J. Royle.

Submitted: 5 March 2022
Revised: 29 March 2022
Accepted: 5 April 2022

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Figure S1. Ensheathed chromosomes in DLD-1 cells after targeted missegregation of Y-chromosome. (A) Schematic diagram after Ly et al. (2017), showing how reexpression of a CENP-A mutant (C-H3) in DLD-1 cells where CENP-A is degraded causes selective misalignment of the Y-chromosome. WT and C-H3 lines were further modified to express mCherry-Sec61β. (B) Western blot of lysates from WT or C-H3 DLD-1 cells treated with doxycycline (Dox) and/or indole-3-acetic acid (IAA) as indicated. Upper blot shows anti-CENP-A detection of endogenous CENP-A fused to EYFP-AID tag (66 kD) and expression of untagged CENP-A (either WT or C-H3). Lower blot shows GAPDH loading control. (C) Typical FISH images locating the Y-chromosome in the main nucleus in control cells and in a micronucleus in cells expressing C-H3 CENP-A. Scale bar, 10 µm. (D) Western blot of lysates from stable cell lines expressing mCherry-Sec61β derived from WT (G2) or C-H3 (G11). Detection of Sec61β or mCherry is shown as indicated with actin loading controls. Migration of Sec61β and mCherry-Sec61β is indicated by black and red arrowheads, respectively. Note that the expression of mCherry-Sec61β downregulates endogenous Sec61β. (E) Deconvolved wide-field microscopy images showing an ensheathed chromosome in G11 cells but not in G2 cells treated with Dox/IAA. Scale bars, 10 µm; 2 µm (insets). (F) Spatially averaged view of all kinetochores in the G11 DLD-1 Dox/IAA dataset (see Materials and methods). Small gray points represent kinetochores at the metaphase plate. Colored points represent misaligned chromosomes that were ensheathed (orange) and those that were not (blue). Spindle poles are shown in black. (G) Box plot to show the relative position of each kinetochore relative to the exclusion zone boundary. Ratio of kinetochores within the exclusion zone are <0 and those within the ER are >0 on a log2 scale. Dots represent kinetochore ratios from 50 DLD-1 cells at metaphase. Boxes show IQR, bar represents the median, and whiskers show 9th and 91st percentiles. Source data are available for this figure: SourceData FS1.
Figure S2. **Spindle assembly checkpoint and micronucleus formation in DLD-1 cells.** (A) Micrographs of immunofluorescence experiments to detect Bub1 or Mad2 (SAC, green) at kinetochores (CENP-C, blue) in cells stably expressing mCherry-Sec61β (red); DAPI-stained DNA is shown in gray. Scale bars, 10 µm; 2 µm (insets). (B) Quantification of Bub1 and Mad2 immunofluorescence at kinetochores marked by CENP-C. Ensheathed chromosomes were classified using the mCherry-Sec61β signal. Dots show kinetochore measurements, boxes show IQR, bar represents the median, and whiskers show 9th and 91st percentiles (Bub1: nA = 52, nF = 49, nE = 52; Mad2: nA = 55, nF = 57, nE = 55). (C) Stills from a video showing an example of ensheathed chromosomes in G11 DLD-1 cells forming micronuclei following Dox/IAA treatment. Scale, 10 µm.
Figure S3. Ensheathed chromosomes do not have stable microtubule-kinetochore attachment. (A) Micrographs of RPE-1 cells stably expressing GFP-Sec61β (gray) pretreated with GSK923295 immunostained for tubulin (red) and CENP-C (green); DNA stained with DAPI. Examples show end-on attachments at aligned kinetochores and potential lateral kinetochore-MT contacts for ensheathed chromosomes. (B) Micrographs of RPE-1 cells stably expressing GFP-Sec61β (gray) pretreated with GSK923295 immunostained for kinastrin (red) and CENP-C (green); DNA stained by DAPI. Scale bars, 10 µm; 2 µm (insets). (C) Frequency distributions of the proximity of the nearest kinastrin punctum to each kinetochore (CENP-C punctum). Kinetochores (n, % with kinastrin <600 nm): aligned (3,124, 26.8%); free (74, 4.1%); ensheathed (227, 6.2%). (D and E) Still images from live-cell imaging experiments of RPE-1 cells stably expressing GFP-Sec61β (green) and Histone H3.2-mCherry (gray), pretreated with 150 nM GSK923295 and stained with SiR-Tubulin (red). Similar results were recorded in 25 cells with free chromosomes and 16 cells with ensheathed chromosomes. Scale bars, 10 µm; 2 µm (insets).
Figure S4. **Mitotic timing and fate of HCT116 cells pretreated with CENP-E inhibitor.** (A) Mitotic timing of HCT116 cells. Cumulative frequencies for NEB to metaphase (NEB-Meta) and metaphase to anaphase (Meta-Ana) are shown. Cells were treated with 150 nM GSK923295 for 3 h before washout for 1 h and subsequent imaging. Control, \( n = 43 \); GSK pretreatment, \( n = 40 \); pooled from three experiments. (B) Sankey diagram to show the fate (right) of cells in each of the three metaphase classes (left). Fates include normal division, micronuclei formation, death, and other defects (lagging chromosome, cytokinesis failure). Note that the fate of cells (and not chromosomes) is tracked.
Figure S5. Stable transgene expression in RPE1 cells and fate of misaligned chromosomes in RPE1 cells stably expressing GFP-Sec61β. (A–D) Western blots to examine expression of proteins in parental RPE1 cells or clonal cells stably expressing GFP-Sec61β alone or with Histone3.2-mCherry, LBR-mCherry, or mCherry-BAF, as indicated. Membranes were probed for GFP, Sec61β, mCherry, LBR, BAF. Actin or tubulin is shown as a loading control. Green or red arrowheads indicate the expected position of GFP- or mCherry-tagged protein; black arrowheads indicate the untagged protein. (E) Mitotic timing of RPE1 cells stably expressing transgenes. Cumulative frequencies for NEB to metaphase, metaphase to anaphase, and NEB to anaphase are shown. Parental, n = 69; GFP-Sec61β alone, n = 52; GFP-Sec61β and LBR-mCherry, n = 66; GFP-Sec61β and mCherry-BAF, n = 51. (F) Sankey diagram to show the fate (right) of RPE1 cells in each of the three metaphase classes (left). Fates include normal division, micronuclei formation, death, and other defects (lagging chromosome, cytokinesis failure). Note that the fate of cells (and not chromosomes) is tracked. LBR-mCherry/GFP-Sec61β, n = 51; mCherry-BAF/GFP-Sec61β, n = 67; pooled from three experiments. (G) Sankey diagram to show the fate (right) of chromosomes in each of the three metaphase classes (left) after GSK923295 pretreatment. Fates include rescue, micronuclei formation, death, and other defects (lagging chromosome, cytokinesis failure). Number of chromosomes: free, 146; ensheathed, 207; lagging, 9. The same dataset was analyzed for the outcome of cells (classified by the final misaligned chromosome) in Fig. 4. Note that ensheathed chromosomes at metaphase that were rescued all became “free” chromosomes before rescue. Source data are available for this figure: SourceData FS5.
Video 1. **3D reconstruction of an ensheathed chromosome in a HeLa cell.** SBF-SEM data from a HeLa cell with spontaneously occurring ensheathed chromosome. The following cellular features are shown (in order of appearance): spindle microtubules (green), centrioles (yellow), DNA (red), mitochondria (multicolored then gold), endomembranes (white), plasma membrane (blue). Playback, 25 fps.

Video 2. **3D reconstruction of an ensheathed chromosome in a HeLa cell.** Same reconstruction but showing only chromosomes (red) and endomembranes (blue). Endomembranes that ensheathe the chromosome of interest are shown in purple. Playback, 25 fps.

Video 3. **3D reconstruction of an ensheathed chromosome in an RPE-1 cell.** A substack from SBF-SEM imaging showing a chromosome (gray) outside the exclusion zone (pink), ensheathed in endomembranes (blue). Three complete rotations are shown with DNA only, DNA plus exclusion zone boundary, finally with endomembranes (ER and mitochondria, brown) added. Scale bar, 2 µm. Playback, 4 fps.

Video 4. **Example of GFP-Mad2 at an ensheathed chromosome.** GSK923295-pretreated RPE-1 cell stably expressing GFP-Mad2 (left) and mCherry-Sec61β (middle) with DNA stained with SiR-DNA (right). Time, hh:mm. Playback, 8 fps.

Video 5. **Example of mitotic outcome of a cell with aligned chromosomes.** Control RPE-1 cell expressing GFP-Sec61β (green) stained with SiR-DNA (red). Cell has all chromosomes aligned and divides normally. Time, hh:mm. Playback, 8 fps.

Video 6. **Example of mitotic outcome of a cell with an ensheathed chromosome.** GSK923295-pretreated RPE-1 cell expressing GFP-Sec61β (green) stained with SiR-DNA (red). Cell has an ensheathed chromosome and missegregates, leading to a micronucleus. Time, hh:mm. Playback, 8 fps.

Video 7. **Example of ER clearance and subsequent rescue of an ensheathed chromosome.** Control (left) and ER clearance (right) in mitotic HCT116 cells expressing FKBP-GFP-Sec61β (green) and Stargazin-mCherry-FRB (red). DNA is stained with SiR-DNA (magenta). Scale bar, 10 µm. Playback, 10 fps.