From equator to pole: splitting chromosomes in mitosis and meiosis

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During eukaryotic cell division, chromosomes must be precisely partitioned to daughter cells. This relies on a mechanism to move chromosomes in defined directions within the parental cell. While sister chromatids are segregated from one another in mitosis and meiosis II, specific adaptations enable the segregation of homologous chromosomes during meiosis I to reduce ploidy for gamete production. Many of the factors that drive these directed chromosome movements are known, and their molecular mechanism has started to be uncovered. Here we review the mechanisms of eukaryotic chromosome segregation, with a particular emphasis on the modifications that ensure the segregation of homologous chromosomes during meiosis I.

Segregation machinery and components

The accurate segregation of the genetic material in eukaryotes is guided by three basic principles: [1] Force needs to be generated to power the movement of the DNA. [2] DNA needs to be linked to other cellular structures that will mediate its segregation. [3] The units of DNA to be partitioned need to be held together prior to being segregated. The molecular components that ensure that these requirements are fulfilled are described below.

**Powering chromosome movement (microtubules)**

The most prominent structure in a mitotic cell is the bipolar spindle (made up of microtubules and associated motor proteins), which provides the force to move chromosomes and thereby bring about their segregation. Microtubules are nucleated by the centrosome (called spindle pole body [SPB] in yeasts). Microtubules are assembled from heterodimers of α-tubulin and β-tubulin, which self-assemble in their GTP-bound state into rigid tubes with an ~25-nm outside diameter, the walls of which are built from single layer of tubulins [Fig. 1; Desai and Mitchison 1997]. Microtubules are polar, with their minus end at or near the spindle pole, and the plus end projecting away from the spindle pole [Fig. 1A]. One feature that underpins the biological role of microtubules (see below) is that they are dynamic; i.e., new subunits can be added or removed from either end. They can switch from polymerization to depolymerization (catastrophe) or vice versa (rescue) [Fig. 1B] in response to GTP hydrolysis within the tubulin dimers themselves as well as the activity of associated motor proteins and regulators.

Given their inherent dynamics and the existence of associated motor proteins, microtubules could theoretically contribute to chromosome segregation by acting in two ways: as a ratchet to exert pushing and pulling forces or as tracks along which cellular motors can carry chromosomes as cargo. Although motors play important roles in chromosome segregation [Nicklas 1989; Song and Mandelkow 1993; Endow et al. 1994; Noda et al. 1995; Gaglio et al. 1996; Tytell and Sorger 2006], they are not essential in fungi [Tanaka et al. 2005, 2007; Grishchuk and McIntosh 2006], and their depletion in vertebrates does not completely abolish chromosome motion [Kapoor et al. 2006; Yang et al. 2007]. Additionally, microtubules can support directional motion in the absence of motor function [Koshland et al. 1988; Lombillo et al. 1995; Grishchuk et al. 2005]. Indeed, microtubule depolymerization is thought to provide the primary force that drives chromosome motion: A single depolymerizing microtubule can generate up to 10 times as much force as a motor enzyme [Inoué and Salmon 1995]. Microtubules grow via the addition of GTP-bound tubulin dimers, which hydrolyze GTP after polymerization. The GDP-bound dimer is bent compared with the GTP-bound counterpart [Fig. 1C]. This bend is constrained within the microtubule lattice in such a way that some of the energy released from GTP hydrolysis is stored in the polymer lattice in the form of physical strain. During microtubule depolymerization, this energy is released as the dissociated tubulin dimers adopt their preferred bent conformation. It has been estimated that a single protofilament can generate up to 5 pN during depolymerization, this means that a single microtubule (composed of 13 protofilaments)
can produce a force of 65 pN (Grishchuk et al. 2005). This is far higher than what is required for chromosome segregation—as little as 0.1pN, as predicted by theoretical analyses (Nicklas 1965). In a pioneering study, Nicklas (1983) was able to measure the force exerted by the spindle on a single chromosome. By using a microneedle to apply and measure the force needed to stall a chromosome in grasshopper spermatocytes, he estimated that the spindle could generate up to 700 pN on a single kinetochore with multiple microtubules attached (Nicklas 1983). However, more recent measurements of spindle forces suggest that this might be a large overestimation (Ferraro-Gideon et al. 2013). Thus, it is still uncertain how much force a spindle generates in cells.

**Linking chromosomes to the spindle (kinetochores)**

Harnessing the energy provided by microtubules and converting it into directional and processive chromosome movement require a coupling device that can associate with both chromosomes and microtubules while resisting the force applied on chromosomes. The kinetochore is the structure that achieves this feat. Kinetochores are protein complexes that assemble on centromeres, specific regions of each chromosome specified by the presence of the histone H3 variant CENP-A. Kinetochore function depends on their ability to form persistent load-bearing attachments to the highly dynamic plus ends of microtubules. The persistent attachment is important, since, due to lack of inertia in the cellular environment, force needs to be constantly applied on chromosomes during their separation. Our understanding of the properties and function of the kinetochore has been enhanced by several lines of investigation: from genetics and proteomics to structural biology and single-molecule biophysics.

The kinetochore can be thought of as comprising three parts: the inner kinetochore, which interacts with centromeric chromatin; the outer kinetochore, which directly interacts with spindle microtubules; and the central kinetochore, which connects the two (Fig. 2A; for excellent recent reviews of kinetochore structure, see Biggins 2013; Cheeseman 2014). Kinetochore proteins tend to be well...
conserved in different species, with some exceptions mostly in the inner kinetochore components [see Westermann and Schleiffer 2013 for a summary of homologs]. The microtubule-binding elements of the outer kinetochore capture microtubules by chance: First, they associate with the microtubule lateral surface, which provides a larger contact surface compared with microtubule tips [Fig. 2B; Hayden et al. 1990, Rieder and Alexander 1990, Tanaka et al. 2005, Franco et al. 2007, Magidson et al. 2011]. These initial lateral attachments are aided by the nucleation of additional microtubules at the kinetochore, which become integrated into the spindle [Kitamura et al. 2009]. Additionally, in vertebrate cells, chromosomes modulate the local concentration of Ran-GTP to facilitate microtubule capture by kinetochores [Caudron et al. 2005; Kalab et al. 2006]. Lateral attachments are subsequently converted into the stronger and more processive end-on attachments. The kinetochore also directly modulates microtubule dynamics. Indeed, the recombinant Ndc80 complex favors rescue [the transition from microtubule shortening to growth] by directly stabilizing the tips of disassembling microtubules [Akiyoshi et al. 2010, Umbreit et al. 2012]. Thus, the kinetochore both controls and harnesses the force generated by microtubules to direct chromosome segregation.

How does the kinetochore use the chemical energy of microtubule depolymerization to power chromosome movement? Careful tracking of chromosome movement and microtubule dynamics showed that disassembly of tubulin subunits from kinetochore-bound microtubule plus ends was associated with poleward chromosome movement [Gorbsky et al. 1987; Inoué and Salmon 1995]. This suggested that kinetochores could move chromosomes toward the pole as they maintain their attachment to the disassembling plus end. Two nonmutually exclusive models have attempted to explain the mechanism by which kinetochores maintain the attachment with the disassembling tip to provide directional movement: the biased diffusion model and the conformational wave model [Fig. 1D, Asbury et al. 2011]. In the first model, the kinetochore forms multiple additive and mobile attachments with microtubules [Hill 1985]. Diffusion that increases the contacts with the microtubule favors attachment, thereby providing a biased direction. Lending support to this model, microtubule-binding elements are present at multiple copies in the kinetochore [Joglekar et al. 2006, 2008; Johnston et al. 2010], making kinetochores able to form multivalent attachments to microtubules, as indeed shown by electron microscopy [EM] studies of the budding yeast kinetochore [Gonen et al. 2012]. Furthermore, recombinant Ndc80 and Dam1 complexes diffuse rapidly along the microtubule lattice [Westermann et al. 2005, 2006; Powers et al. 2009; Alushin et al. 2010]. The alternate, conformational wave model postulates that as the microtubule protofilaments bend outward during depolymerization, they push on the kinetochore, pulling it along the microtubule [Koshland et al. 1988]. The conformational wave model relies on a structure that would serve as a hook on which bending microtubules could push during disassembly. A microtubule-encircling ring has been proposed to be a possible mediator. In support of this model, the Dam1 complex in budding yeast forms a ring with 16-fold symmetry around microtubules in vitro [Miranda et al. 2005; Westermann et al. 2005], and EM studies show that budding yeast kinetochore rings often encircle microtubules [Gonen et al. 2012]. However, in vitro studies have shown that the Dam1 complex is capable of tracking disassembling microtubules even in the absence of the ring structure [Gestaut et al. 2008; Grishchuk et al. 2008]. Importantly, a purely conformational wave mechanism would predict that kinetochores would detach more quickly during assembly, when curling protofilaments are much less prominent [Mandelkow and Mandelkow 1985]. However, single-molecule studies suggest that kinetochores actually detach more quickly during disassembly [Akiyoshi et al. 2010]. It is likely that mechanisms and features proposed by both models contribute to the load-bearing attachments.
of kinetochores. Indeed, using the deformation of moving kinetochores as a readout of forces exerted on them, it was found that both active force generation within kinetochores and passive frictional interactions with microtubules contribute to these persistent attachments (Dumont et al. 2012).

**Holding sister DNA molecules together (cohesion)**

Microtubules and kinetochores could, in principle, move chromosomes in any direction. Directionality in chromosome movement requires that sister chromatids be physically linked to provide an opposing force to that of the microtubules. Cohesin is the protein complex that achieves this by entrapping sister chromatids in a ring (for a recent review, see Marston 2014). Condensins, which are related to cohesins and also form a ring, give chromosomes their compact rod-shaped structure that allows their capture and movement during chromosome segregation (for review, see Hirano 2012). The pericentromere, the chromosomal region surrounding the centromere, is the region that experiences the highest levels of force, as evidenced by the separation of sister centromeres, but not arms, during metaphase in yeast (Goshima and Yanagida 2000; He et al. 2000; Tanaka et al. 2000). Both cohesin and condensin are highly enriched at the pericentromere (Blat and Kleckner 1999; Megee et al. 1999; Tanaka et al. 1999; Glynn et al. 2004; Kiburtz et al. 2005; D’Ambrosio et al. 2008; Verrijzer et al. 2014) and are crucial to the architecture of pericentromeric chromatin in both yeast (Yong-Gonzalez et al. 2007; Ng et al. 2009; Stephens et al. 2013) and mammals (Ribeiro et al. 2009). Cohesin and condensin organize pericentromeric chromatin into a spring: Condensin compacts chromatin along the spindle axis, whereas cohesin localizes around the spindle axis and prevents the chromatin from spreading out radially (Stephens et al. 2011). This spatial confinement provides pericentromeric chromatin with the necessary rigidity to counterbalance spindle forces, allowing it to stretch, rather than break, under the force of the spindle. Furthermore, pericentromeres have been proposed to be cross-linked together, a feature that would allow a more efficient tension-based stabilization of multiple attachment sites (Stephens et al. 2013).

**Moving chromosomes in the right direction (orienting kinetochores)**

As described above, kinetochores capture microtubules in an essentially stochastic way. However, faithful chromosome segregation requires that sister kinetochores attach to microtubules emanating from opposite spindle poles; this is termed amphitelic attachment, and the sister kinetochores are said to be bioriented [Fig. 2B]. Sister kinetochores can also attach to microtubules from the same pole (syntelic attachments). Additionally, a single kinetochore can attach to microtubules from opposite poles, giving rise to merotelic attachments [Fig. 2B]. Since both syntelic and merotelic attachments are not compatible with accurate chromosome segregation during mitosis, what mechanisms are in place to ensure that correct attachments are made?

Tension lies at the heart of these mechanisms. The fundamental importance of tension was first made evident by elegant micromanipulation experiments with insect cells that showed that tension is used as a readout of accuracy (Li and Nicklas 1995; Nicklas et al. 1995; Nicklas 1997). When sister kinetochores are bioriented, the pulling force of spindle microtubules is counteracted by the cohesin linkages between sister chromatids, generating tension between sister kinetochores. This is the only mode of attachment that will exert equal force on each sister kinetochore, thereby producing even tension across them. Artificially applying tension on kinetochores both stabilizes and increases the number of microtubule–kinetochore attachments (Nicklas et al. 1995; King and Nicklas 2000). Tension stabilizes bipolar attachments by both direct (mechanical) and indirect (chemical) means. Kinetochores bind strongly to growing microtubules and weakly to shrinking microtubules. Tension suppresses microtubule disassembly, thus favoring the strongly bound state (Akiyoshi et al. 2010). In the indirect regulation, the pulling apart of sister centromeres removes kinetochores from the field of action of the kinase Aurora B, which continuously phosphorylates kinetochore components within its reach to disrupt kinetochore–microtubule attachments (Biggins et al. 1999; Tanaka et al. 2002; Liu et al. 2009; Welburn et al. 2010). Aurora B plays a crucial role in ensuring correct chromosome–microtubule attachments by releasing kinetochores in two ways: It weakens the attachment of the outer kinetochore proteins to the microtubule and directly destabilizes the kinetochore-attached microtubule tip (Lampson et al. 2004; Sarangapani et al. 2013).

Finally, in mitotic chromosomes, sister kinetochores are arranged in a way that favors biorientation. Indeed, sister kinetochores are thought to assume a back-to-back geometry, which favors their capture of microtubules from opposite poles [Figs. 2C, 4C [below]]. Data from budding yeast point to the cohesin- and condensin-dependent pericentromere architecture producing an intrinsic bias of sister kinetochores toward biorientation (Indjian and Murray 2007; Ng et al. 2009; Peplowska et al. 2014; Verrijzer et al. 2014).

Thus, the correct attachment of kinetochores to microtubules depends on both chromosome architecture (dictated by cohesin and condensin) and the generation of tension. Meiosis, the specialized cell division that gives rise to haploid gametes from a diploid progenitor cell, is guided by the same basic principles but with certain modifications to allow for a different chromosome segregation pattern.

**Specialization of the chromosome segregation machinery for meiosis**

In meiosis, two rounds of segregation follow a single round of replication (for a review, see Marston and Amon 2004). In the first meiotic division (meiosis I), homologous chromosomes segregate away from each other, and sister chromatids congregate. The first meiotic division is often called “reductional,” as it is this segregation event that results in the reduction in ploidy. In the second
meiotic division (meiosis II), much like in mitosis, sister chromatids segregate (Fig. 3). Meiosis follows the same principles of chromosome segregation as mitosis; however, three important modifications underpin the first meiotic division: [1] Homologous chromosomes are physically linked together, usually by chiasmata, the products of homologous recombination. [2] Sister kinetochores attach to microtubules emanating from the same spindle pole (they are mono-oriented). [3] Cohesion is lost in a stepwise manner: Cohesion is lost on chromosome arms during the first division but is protected at the pericentromere (Fig. 3B). Importantly, these modifications are specified by properties of the chromosome rather than the spindle (Paluiulis and Nicklas 2000). Nevertheless, the spindle does show meiosis-specific features in some organisms [Yi et al. 2013]. It is likely that these adaptations are important for other developmental aspects of meiosis rather than for dictating the specialized chromosome segregation pattern [see below].

**Linking homologs**

The biorientation of sister chromatids in mitosis relies on the tension created only by correct attachments: The pulling force of microtubules is counteracted by the physical linkages of cohesin between sister chromatids. In the first meiotic division, however, it is homologous chromosomes that must segregate away from each other. Meiosis relies on the same tension-based mechanisms for ensuring correct attachment of chromosomes to the spindle.

To enable these mechanisms to ensure biorientation of homologous chromosomes in meiosis I, physical linkages that can counteract spindle tension are generated between homologs by homologous recombination. Several excellent reviews [Baudat et al. 2013; Boarde and de Massy 2013; de Massy 2013] have summarized recent strides in our understanding of homologous recombination in meiosis and its regulation. In many organisms, recombination is preceded by homologous chromosomes pairing along their lengths; this pairing is stabilized by synapsing through the assembly of a proteinaceous structure known as the synaptonemal complex (SC) [for reviews, see Zickler and Kleckner 1999, Bhalla and Dernburg 2008]. Recombination starts in this context with the action of the endonuclease Spo11, which introduces deliberate and stochastic double-strand breaks along the chromosome [Keeney et al. 1997, 1999; Romanienko and Camerini-Otero 1999]. A subset of these breaks is repaired using the nonhomologous homologous chromosome, creating physical links between homologs called chiasmata. The resulting homologous chromosome pair, called a bivalent, can now orient on the spindle, with interhomolog chiasmata resisting the pulling forces of the spindle. A single chiasma is sufficient to support the tension that is required for the accuracy of chromosome segregation [Hillers and Villeneuve 2003]. It is at present unclear how the counterbalancing resistance provided by centromere-distal chiasmata is transmitted to kinetochores on the centromere. Whether meiotic chromosomes possess spring-like behavior at the point of tension, as observed for pericentromeres in mitosis, and whether increased structural rigidity of chromosome arms is required to allow force transduction along them remain questions for the future.

Once connections between homologs are made, homologous chromosomes need to attach to opposite poles so that they can segregate away from each other in anaphase I. In a manner similar to biorientation of sister chromatids in mitosis, tension and the action of Aurora B kinase play critical roles in achieving correct biorientation of homologs [Monje-Casas et al. 2007; Sakuno et al. 2011, Meyer et al. 2013].

**Protecting linkages between sister chromatids during meiosis I**

Once homologs are bioriented, the links between them need to be severed to allow the poleward movement of chromosomes. In budding yeast, phosphorylation of the Rec8 cohesin subunit targets it for cleavage by separase [Brar et al. 2006; Ishiguro et al. 2010; Katis et al. 2010; Attnet et al. 2013]. Cohesin cleavage on chromosome arms resolves chiasmata, thereby allowing homologous chromosomes to segregate [Buonomo et al. 2000; Kudo et al. 2006]. Cohesin in centromeric regions, however, must be protected from separase activity during meiosis I in order to ensure faithful segregation of sister chromatids in the second meiotic division. Genome-wide screens in budding yeast and fission yeast identified the shugoshin proteins, distant relatives of the fruit fly Mei-S332 protein, as essential factors protecting centromere cohesin at the end of the first division [Kerrebrock et al. 1992, Kitajima et al. 2004; Marston et al. 2004; Rabitsch et al. 2004]. Shugoshin recruits the protein phosphatase PP2A to the pericentromere, which dephosphorylates Rec8, thereby rendering it refractory to separase cleavage [Kitajima et al. 2006; Riedel et al. 2006; Ishiguro et al. 2010; Katis et al. 2010]. Residual pericentromeric cohesin provides the resistance to spindle forces during meiosis II, where sister kinetochores are bioriented in preparation for the mitosis-like segregation of sister chromatids to opposite poles.

**Cosegregation of sister chromatids during meiosis I**

The comigration of sister chromatids in the first division requires that sister kinetochores be mono-oriented; i.e., attach to microtubules that emanate from the same spindle pole. The different arrangement of kinetochores in meiosis compared with mitosis was first reported from EM work in *Drosophila melanogaster* spermatocytes [Goldstein 1981], where sister kinetochores were shown to be very closely associated. This suggested a mechanism for mono-orientation whereby sister kinetochores “fuse”; i.e., they create a single microtubule-binding interface. Recent support for this model came from fluorescence microscopy studies in maize meiocytes. Li and Dawe [2009] found that the MIS12 and NDC80 kinetochore components span across the sister centromeres to form a direct cross-linking bridge between the sister kinetochores. Knockdown of MIS12 by RNAi weakened this link and caused a third of sister chromatids to segregate away from each other [Li and Dawe 2009]. While these
Figure 3. The chromosome segregation program in mitosis (A) and meiosis (B). During DNA replication, cohesin rings (blue) topologically entrap sister DNA molecules to give rise to sister chromatid cohesion. (A) In mitotic metaphase, sister kinetochores (red circles) are bioriented (black triangles): They attach to microtubules (green) emanating from opposite spindle poles. In anaphase, cohesin is cleaved, allowing sister chromatids to separate. (B) In meiotic prophase I, homologous recombination (HR) allows homologous chromosomes to be physically linked via chiasmata. In meiotic metaphase I, sister kinetochores are thought to fuse so as to present as a single microtubule-binding interface. In anaphase I, centromere-distal cohesin is cleaved, allowing homologous chromosomes to separate. Centromere cohesin, however, is protected. In meiosis II, much like in mitosis, sister kinetochores biorient in metaphase II, and the cleavage of centromere cohesin in anaphase II allows sister chromatids to segregate.
studies have been critical for understanding higher eukaryote sister mono-orientation, the bulk of the molecular insights into mono-orientation mechanisms has come from studies in budding and fission yeasts.

Budding yeast and sister kinetochore fusion  The first suggestion for a sister kinetochore fusion model in budding yeast meiosis was inspired by EM studies of the meiosis I spindle, which showed that the number of kinetochore microtubules was not sufficient for each sister kinetochore to be attached independently to the spindle [Winey et al. 2005]. Many studies have since supported the view that in budding yeast, mono-orientation is achieved by the rearrangement of sister kinetochores to create a single microtubule-binding unit [Fig. 4A]. Crucial to this rearrangement is monopolin, a four-protein complex identified by functional genomics and proteomics [Töth et al. 2000; Rabitsch et al. 2003; Petronczki et al. 2006]. Monopolin consists of Mam1, a protein expressed exclusively during the first meiotic division; casein kinase Hrr25; and the proteins Csm1 and Lrs4 [Fig. 4B]. Monopolin associates with kinetochores from late prophase I (when the expression of the MAM1 gene is induced) until the end of the first meiotic division (Töth et al. 2000). At least two additional kinases play important roles in the function of monopolin: polo-like kinase Cdc5 and the Dbf4-dependent kinase (DDK). Cdc5 releases Csm1 and Lrs4 from the nucleolus, where they normally reside [Clyne et al. 2003; Lee and Amon 2003; Rabitsch et al. 2003]; subsequently, Cdc5 and DDK together act to phosphorylate Lrs4 [Matos et al. 2008]. Whether these are the only critical functions that these kinases play in mono-orientation is not yet known. The stable association of monopolin with kinetochores also requires Spo13, a meiosis I-specific protein that regulates many aspects of budding yeast meiosis in poorly understood ways [Shonn et al. 2002; Katis et al. 2004; Lee et al. 2004]. Recently, it was shown that mono-orientation in budding yeast also depends on the temporal regulation of kinetochore–microtubule attachments [Miller et al. 2012]. Kinetochore–microtubule attachments are abolished in prophase of the first meiotic division because Ndc80, which provides the main microtubule-binding activity of the

Figure 4. Mono-orientation of sister kinetochores in meiosis I. (A) Monopolin [pink] cross-links sister kinetochores to create a single microtubule-binding interface. (B) Diagram of the organization of the monopolin complex. Csm1 [dark blue] and Lrs4 [light blue] form a V-shaped complex, with the Csm1 globular heads spaced at 10 nm apart. Csm1 is thought to interact directly with the N terminus of Dsn1 [red] via Csm1’s globular head. The other globular head interacts with Mam1 [yellow], which in turn recruits a copy of casein kinase [Hrr25, purple]. The copy number of each protein in the complex is indicated in brackets. (C) In fission yeast, sister kinetochore orientation is determined by centromeric cohesion: When there is no cohesion at the core centromere, sister kinetochore biorient [left]; cohesion at the core centromere allows for mono-orientation in meiosis I [right]. (D) In C. elegans, a kinetochore sheath forms around the bivalent in meiosis I [left] or sister chromatids in meiosis II [right]. Aurora B kinase [AIR-2, yellow] forms a ring around the mid-bivalent [meiosis I] or sister chromatid interface [meiosis II]. AIR-2 is also thought to mark the site for CLASP-dependent microtubule growth that pushes the dividing chromosomes apart in a kinetochore-independent manner. (E) Kinetochore microtubule-stabilizing protein CLASP promotes microtubule polymerization between chromosomes. This microtubule growth could generate the force required for the segregation of chromosomes.
kinetochore, is destabilized in an Ipl1-dependent manner [Miller et al. 2012; Kim et al. 2013]. If kinetochores are induced to engage with microtubules prematurely, mono-orientation is not established [Miller et al. 2012]. It seems likely that monopolin cannot bind to kinetochores that are already associated with microtubules.

Monopolin is recruited to kinetochores by binding to the N terminus of Dsn1 [Mis13 in other organisms], a component of the MIND complex of the central kinetochore [Fig. 4B; Sarkar et al. 2013; Sarangapani et al. 2014]. Notably, this could also act as the point at which sister kinetochores are cross-linked into a single functional unit. The crystal structure of the Csm1:Lrs4 complex was the first to provide strong evidence for direct cross-linking of sister kinetochores [Corbett et al. 2010; Corbett and Harrison 2012]. Csm1:Lrs4 form a V-shaped complex, with each of the globular “heads” containing a dimer of Csm1 C-terminal domains [Fig. 4B]. This suggested that each “head” could bind to a different sister kinetochore to bring about cross-linking. This has been further supported by data showing that N-terminally truncated Dsn1 dominantly prevents sister kinetochore mono-orientation in meiosis I when produced in the presence of wild-type Dsn1 [Sarkar et al. 2013]. This points to the N terminus of Dsn1 as a potential cross-linking site for kinetochores [Fig. 4B].

More recently, single-molecule techniques have been used to directly assess the role of monopolin in fusing sister kinetochores. Kinetochores purified from meiosis I withstand more force and have more microtubule-binding elements than mitotic kinetochores [Sarangapani et al. 2014]. This is unlikely to be solely due to monopolin-induced rearrangements in the architecture of individual kinetochores: DNA replication and thus the presence of a sister kinetochore are required for this increase in strength [Sarangapani et al. 2014]. Importantly, monopolin was able to increase the strength of mitotic kinetochores in vitro in a manner that was independent of casein kinase activity [Sarangapani et al. 2014]. The latter highlights our lack of understanding of the role of casein kinase in promoting mono-orientation. The kinase activity of Hrr25 is required for mono-orientation but not for the recruitment of monopolin [Petronczki et al. 2006]. Intriguingly, Hrr25 phosphorylates Mam1 in a manner that destabilizes the monopolin complex [Corbett and Harrison 2012]. This suggests that Hrr25 might phosphorylate kinetochore components to either achieve mono-orientation or ensure that the correct sisters are fused. The precise role played by Mam1 is also unclear. Mam1 is unlikely to act simply as a recruitment factor, since both Csm1 and Lrs4 are recruited to kinetochores in mitotic anaphase, when Mam1 is not expressed [Akiyoshi et al. 2009; Brito et al. 2010]. Furthermore, Mam1 binds to Csm1 to occlude one of its kinetochore-binding interfaces [Corbett and Harrison 2012]. In vitro and structural work will be required to elucidate the precise nature of sister kinetochore fusion and the role played therein by each component of monopolin.

Fission yeast and the role of cohesion in mono-orientation In fission yeast, mono-orientation has been shown to be driven by modifications in centromere cohesion [Sakuno et al. 2009]. Fission yeast centromeres are composed of heterochromatic outer repeats and a central core domain containing CENP-A nucleosomes. In mitosis, cohesin is highly enriched at the outer repeats flanking the central domain of the centromere, which is depleted for cohesin; this is thought to favor a back-to-back configuration of sister kinetochores, which in turn promotes their biorientation [Fig. 4C, left panel]. In meiosis, however, the meiosis-specific cohesin subunit Rec8 is enriched also at the central core of the centromere, which is thought to favor a side-by-side arrangement of sister kinetochores to allow their efficient mono-orientation [Fig. 4C, right panel; Watanabe and Nurse 1999; Sakuno et al. 2009]. The meiosis-specific factor Moa1 is recruited to the kinetochore via the inner centromere protein CENP-C and promotes cohesion at the core centromere [Yokobayashi and Watanabe 2005; Tanaka et al. 2009]. Rec8 disappearance from the core centromere before prophase II would allow for the biorientation of sister kinetochores in meiosis II. Here it is important to note that Sgo1, which protects centromere cohesin from being cleaved in anaphase I, localizes only at the pericentromere and not at the core centromere [Sakuno et al. 2009].

Lending further support to the importance of geometry for mono-orientation, an artificial tether between the central domains of sister centromeres restored mono-orientation in meiosis I in the absence of Rec8 [Sakuno et al. 2009]. However, centromeric tethers induced in mitosis led to a modest increase in cosegregation of sister chromatids [Sakuno et al. 2009]. In contrast, ectopic localization of monopolin to the mitotic kinetochore in budding yeast leads to up to 50% mono-orientation of sister kinetochores [Monje-Casas et al. 2007; Miller et al. 2012]. Thus, other factors are very likely to be required for mono-orientation in fission yeast.

Rec8 has been shown to play a role in many other organisms, including mice [Parra et al. 2004], Arabidopsis thaliana (Chelysheva et al. 2005), and Caenorhabditis elegans [Severson et al. 2009]. This is in contrast to budding yeast, where Rec8 does not seem to be required for establishing mono-orientation of sister kinetochores [Tôth et al. 2000; Monje-Casas et al. 2007]. This could stem from the very different nature of the budding yeast centromere and its kinetochore–microtubule attachments. Budding yeast has simple point centromeres of 125 base pairs (bp) with no flanking heterochromatin, which would not allow for accurate differential organizations of centromere cohesion. Furthermore, in budding yeast, one kinetochore (or a pair of sister kinetochores in meiosis I) attaches to a single microtubule [Winey et al. 2005]. In fission yeast, on the other hand, multiple microtubules (two to three) bind a single kinetochore. This might make sister kinetochore fusion a much more efficient mechanism of mono-orientation in budding yeast. The ability to clamp together microtubule-binding elements suggested that monopolin might have a conserved role in other organisms to prevent merotelic (Rabitsch et al. 2003) even though they are dispensable for sister...
kinetochore mono-orientation (Rabitsch et al. 2003). Importantly, unlike budding yeast, where monopolar seems to clamp microtubule-binding sites directly (Corbett et al. 2010), fission yeast Pcs1 and Mde4 have been shown to prevent merotely by recruiting condensin to centromeres (Tada et al. 2011).

Last, the finding that meiosis-specific cohesins are necessary for sister kinetochore mono-orientation in many organisms but are not sufficient (at least in fission yeast) suggests that additional factors are required. This raises the possibility that undiscovered factors that are functionally analogous to budding yeast monopolin cross-link kinetochores in these organisms. Potentially, meiosis-specific cohesins in the centromeric region may be required to place sister kinetochores in sufficient proximity to allow their cross-linking. This would be envisaged to be especially important in organisms with large centromeres that contact multiple microtubules, while in organisms with point centromeres, monopolin may in itself be sufficient.

Other unusual features of meiosis in some organisms

Sister kinetochore mono-orientation in holocentric chromosomes So far, our discussion has been focused on chromosomes where the kinetochore assembles on a single locus; these are called monocentric chromosomes because they have one centromere. In some organisms, however, kinetochores assemble along the entire length of the chromosome; these are termed holocentric chromosomes. Organisms with holocentric chromosomes include the nematode worm C. elegans and also some insects, arachnids, and several plant species (Albertson and Thomson 1993). One important consequence of holocentric chromosomes is that microtubules can attach at several points along them, so the pulling forces of the spindle are not exerted at a single point. Additionally, this imposes certain modifications for meiotic chromosome segregation. In meiosis I, homologous chromosomes are linked by chiasmata to form a cruciform bivalent [Fig. 3B]. Because kinetochores assemble alongside the length of the chromosome, holocentric bivalents could have microtubule-binding surfaces facing in all directions. To resolve this, kinetochores assemble to form a sheath that encapsulates bivalents [Fig. 4D]. Interestingly, and in stark contrast to mitosis, the cup-like assembly of kinetochores on holocentric chromosomes in meiosis does not require CENP-A [Dumont et al. 2010]. The meiosis-specific cohesin Rec8 plays a fundamental role in connecting the sister chromatids so that they can be encapsulated (Severson et al. 2009). The fact that Rec8 directs key aspects of chromosome segregation in organisms whose kinetochores are arranged in dramatically different ways points to an early adaptation of meiotic cohesin in establishing the meiotic chromosome segregation program.

Segregation of homologs without chiasmata Despite the importance of homologous recombination in the segregation of chromosomes in meiosis I, chromosomes can segregate accurately in the absence of visible chiasmata. In budding yeast, for instance, a single nonexchange [achiasmatic] chromosome segregates faithfully in 90% of meioses [Dawson et al. 1986; Mann and Davis 1986; Guacci and Kaback 1991], and in many organisms, sex chromosomes segregate without any recombination at all (for review, see Wolf 1994). Achiasmatic chromosomes often form physical associations by alternative mechanisms, which can allow the generation of tension when they biorient on the meiosis I spindle. In the female silk moth, Bombyx mori, the SC remains associated with the bivalents until anaphase I (Rasmussen 1977), presumably functioning as a substitute for chiasmata. In some mammals, SC proteins [e.g., SYCP3] form links between sex chromosomes [Page et al. 2006; de la Fuente et al. 2007]. This is reminiscent of the role the budding yeast SC component Zip1 plays to pair centromeres in early meiotic prophase, which is thought to promote the biorientation of homologs [Tsukuboshi and Roeder 2005; Gladstone et al. 2009; Newnham et al. 2010]. Together, these data point to an ancient role for SC proteins in maintaining association of homologous achiasmatic chromosomes. In fruit fly oocytes, heterochromatic threads provide a substitute link between the achiasmatic fourth chromosomes [Dernburg et al. 1996; Karpen et al. 1996]. In fruit fly male meiosis, however, there is no SC, and heterochromatic regions only play a role in a subset of chromosomes [Tsai et al. 2011]. Here, the necessary connection is provided by the nucleolar tDNA regions in a manner dependent on male meiosis-specific proteins SNM and MNM [Thomas et al. 2005].

Poleward microtubule flux We have seen how the major force for chromosome motion is provided by the disassembly of microtubules at the kinetochore-bound plus end. However, microtubules also disassemble at their minus end on the spindle pole in a process termed poleward microtubule flux [Fig. 1E]. Microtubule flux has been shown to play important roles in chromosome congression in metaphase [for review, see Ganem and Compton 2006]. However, it can also generate force. Indeed, microtubule flux may be particularly important in meiosis to support chromosome motion. Microtubule-marking experiments have allowed the rate of poleward microtubule flux to be measured. By comparing the rate of flux with the velocity of chromosome motion, one can estimate the contribution of poleward microtubule flux to chromosome movement. Strikingly, in many of the meiotic cells that have been studied, such as Xenopus egg extracts and insect spermatocytes, flux velocity exceeds chromosome velocity [Desai et al. 1998; LaFountain et al. 2004], strongly suggesting that in these systems, poleward microtubule flux is likely to be the primary mechanism driving chromosome segregation. Conversely, in somatic cells that have been studied, such as PtK1, PtK2, LLC-PK1, Newt lung, and HeLa, poleward microtubule flux makes a small contribution to the chromosome movement [Ganem et al. 2005].

Chromosome segregation without kinetochores or centromeres In the “canonical” system of chromosome segregation depicted in Figure 1A, kinetochores capture
microtubules emanating from the spindle pole, the centrosome, to power chromosome motion. However, in many organisms, including fruit flies, *C. elegans*, mice, and humans, female meiosis takes place in the absence of centrosomes. This could be an adaptation that avoids the formation of multipolar spindles once sperm carrying the centrosome enter after fertilization. In these acellorosomal divisions, the spindle is assembled from the self-organization of many alternative microtubule-organizing centers (Schuh and Ellenberg 2007), often including chromatin itself (Heald et al. 1996). A striking example of how acelloral meiosis redefines basic principles of chromosome segregation is provided by *C. elegans*. *C. elegans* oocytes do not rely on kinetochores to power the separation of chromosomes; kinetochores are simply required to orient chromosomes in the spindle (Dumont et al. 2010). Instead, microtubules nucleate between the separating bivalents, and it is this that drives the separation of chromosomes (Fig. 4E). The microtubule dynamics regulator protein CLASP as well as Aurora B kinase and the kinase BUB-1 form a ring at the midbivalent that nucleates microtubules. The directed growth of these bundles of microtubules pushes the homologs apart. Why might such a kinetochore-independent mechanism have arisen? Dumont and Desai (2012) suggested that it might be an adaptation of holocentric chromosomes: Aurora kinase associates with the site of crossover, thereby marking both the site for cohesin loss (Schvarzstein et al. 2010) and the site for microtubule growth that will drive segregation. This ensures that homologs, but not sister chromatids, segregate in the first division. Alternatively, or in addition, this may represent a more general mechanism used in acelloral division. DNA-coated beads separate in mouse oocytes in the absence of kinetochore function (Deng et al. 2009), suggesting that kinetochore-independent mechanisms may be in place to generate the force that powers chromosome segregation in acelloral meiosis of other organisms.

Conclusion

Although a wealth of studies has provided detailed insights into how chromosomes are moved during mitosis, key questions remain about how microtubule-generated force is coupled to chromosomes by the remarkable molecular machine that is the kinetochore. During meiosis, adaptations to both kinetochores and the spindle alter the way that force is generated and used. What is the biological rationale underlying these modifications? How do they effect the specialized pattern of meiotic chromosome segregation at the molecular level? The biochemical reconstitution of many components of the cell division machinery coupled with high-resolution imaging of live cells will allow for a plethora of exciting questions to be answered.

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