Centromeres Convert but Don’t Cross

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A long-standing problem in chromosome biology concerns the dynamic nature of centromeres. These chromosomal sites assemble the protein machines called kinetochores that connect chromosomes to the spindle microtubules for segregation to daughter cells during mitosis and meiosis. In multicellular eukaryotes, centromeres are typically composed of highly homogeneous tandem repeats that evolve rapidly despite their highly conserved function [1]. For tandem repeats to evolve, a mutation must spread by some recombinational process, but a persistent dogma is that centromeres do not undergo homologous chromosome recombination (the shuffling of genetic segments between chromosomal pairs). New evidence [2] challenges this dogma and addresses the problem of rapidly evolving centromeres.

The Role of Crossing Over in Meiosis

Centromeres do not act alone in orchestrating chromosome segregation. In order for sister kinetochores to properly disjoin (separate) and segregate chromosomes equally to daughter cells in mitosis, their sister chromatids must be linked so that the pulling forces from the two halves of the spindle generate tension to correctly orient the kinetochores, stabilize kinetochore attachments, and signal that kinetochores are ready to disjoin. Centromeres in multicellular eukaryotes are typically embedded in heterochromatin, the permanently condensed chromatin found around centromeres, in contrast to the euchromatic chromosome arms, which decondense between mitoses. Heterochromatin has been implicated in facilitating cohesin association between sister chromatids around the centromere. This cohesion is mediated by cohesins, proteins that link the sisters together and that are enriched around centromeres [3], and possibly also by catenation (interlocking) of DNA threads observed between sister centromeres [4]. In most eukaryotes, homologs become physically linked during meiosis through the recombinational process of “crossing over”—the breakage and reciprocal reunion of homologous chromatids, resulting in a chiasma, the point where recombinant chromatids cross over each other (Figure 1). Failure to cross over is a major source of non-disjunction (improper segregation) at the first meiotic division in animals [5,6], underscoring the importance of chiasmata for segregation of homologs.

As early as 1930, observations on the distribution of chiasmata along chromosomes led Karl Sax to predict that crossing over (and hence genetic recombination) is reduced around the centromere [7], and this “centromere effect” was verified in the fruitfly Drosophila melanogaster soon afterward [8]. Suppression of crossing over around or in centromeres has since been verified in several animals [9,10], plants [11–14], and fungi [15,16], with estimates of crossover suppression ranging from 3-fold to >200-fold in different organisms.

Why is crossing over suppressed around centromeres? In Drosophila [5], humans [6], and budding yeast (Saccharomyces cerevisiae) [17], non-disjunction events at the second meiotic division are enriched in centromere-proximal crossovers. This suggests that crossovers that are too close to the centromere disrupt pericentric sister chromatid cohesion, leading to premature separation of sister chromatids, which then segregate randomly. Thus, selective pressure to reduce crossing over near the centromere is likely to be strong. Crossing over within the centromere itself could be even more deleterious, leading to attachment of the centromere to both halves of the spindle, resulting in chromosome breakage and loss.

Centromeres, Heterochromatin, and Crossover Suppression

How is crossing over suppressed at centromeres? The location of centromeres in heterochromatin raises the possibility that the crossover suppression seen at the centromere may simply be a property of the surrounding heterochromatin. Early attempts to separate heterochromatin from the centromere utilized inversions of pericentric heterochromatin on the Drosophila X chromosome and suggested that the centromere can suppress recombination independently of its flanking heterochromatin [18]. Subsequent work confirmed that heterochromatin also suppresses crossing over [19], consistent with its proposed role in facilitating cohesion. An increase in crossovers in Drosophila mutants that affect heterochromatin structure support the role of heterochromatin in suppressing pericentric crossovers [20]. Crossover suppression in plants also appears to be a feature of both centromeres and flanking heterochromatin. In Arabidopsis thaliana, crossing over is reduced >200-fold in the 2.3-Mb centromere region of Chromosome 1, and 10–50 fold by the 1-Mb heterochromatic flanking regions [12].

At the molecular level, centromeres are distinguished from both heterochromatin and euchromatin by specialized nucleosomes containing the centromere-specific histone H3 variant known as CENP-A or CenH3, which is necessary to form the kinetochore. Occasionally functional CenH3-containing centromeres can arise on DNA that was previously non-centromeric and be faithfully transmitted (neocentromeres), indicating that centromere inheritance is epigenetic, dependent on the presence of CenH3 nucleosomes, not on specific DNA sequences (reviewed in [1]). Despite the apparent irrelevance of centromeric DNA sequence to kinetochore function, natural centromeres in plants and animals are usually composed of Mb-sized tandem arrays of short

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Abbreviations: CenH3, centromeric histone H3 variant; CRM2, centromere-specific retrotransposon of maize 2; LD, linkage disequilibrium

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Figure 1. Chromosome connections in meiosis. Kinetochores attach homologous chromosomes to opposite halves of the spindle. Homologs are held together by chiasmata, in which recombinant chromatids cross each other. Sisters are held together by cohesins and possibly by catenation of centromeric DNA threads, which have been observed in human mitosis. Cohesion is released in two steps: on chromosome arms to resolve chiasmata and separate homologs in the first meiotic division; and around centromeres to separate sisters in the second meiotic division.

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Figure 2. Unequal exchange in satellite arrays. Identical tandem satellite repeats become diversified over time by mutation. Unequal exchange results in gain or loss of tandem repeats. Repeated exchange can lead to homogenization of satellite repeats (left). If the unit of exchange consists of multiple diverged monomers, higher-order repeats are generated (right).

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(150–180 bp) noncoding “satellite” repeats. These arrays may also be rich in transposon insertions, probably because suppression of crossing over prevents their elimination through recombination. The same or similar repeats comprise the flanking pericentric heterochromatin, underscoring the epigenetic specification of centromeres by CenH3 nucleosomes.

Although both centromeres and pericentric heterochromatin are rich in repetitive elements, repeats per se do not appear to be necessary for crossover suppression. For example, centromere 8 of rice (*Oryza sativa*), which has only very little satellite DNA, lacks detectable crossovers in a 2.3-Mb span around the 750-kb centromere region that contains discontinuous blocks of CenH3-containing nucleosomes. Remarkably, there is little difference in gene activity, transposon composition, or abundance of common histone modifications between this recombination-free region and adjacent recombining regions [21], suggesting that crossover suppression does not depend on DNA sequence but instead is epigenetic.

A clearer separation of centromeric and heterochromatin effects can be found in budding yeast, which is unusual in having “point” centromeres that are only ~120 bp in length [22], harbor a single CenH3 nucleosome [23] and lack surrounding heterochromatin. Suppression of crossing over at yeast centromeres is modest, estimated at only 3–6 fold, and extends over only about 10 kb or less [15,24], although this represents as much as 80 times the length of the centromere itself. This suppression is eliminated by a point mutation in the centromere that renders it unable to assemble a functional kinetochore [25], strongly suggesting that the kinetochore mediates suppression.

**Satellite Arrays and Recombination**

Although crossing over is suppressed around centromeres, the tandem satellite array structure that is typical for most centromeres is best explained by extensive and repeated recombination. The generation of such arrays has been modeled as a recombinational process of random unequal exchange [26]. Unequal exchange can act on variation in the individual satellite monomers due to mutation to lead to expansion of new repeat variants and/or formation of higher-order repeats (Figure 2), as well as eliminating variation in monomers (homogenization). In the human X chromosome, the CenH3-containing chromatin is found centrally in the most recent and most homogeneous higher-order repeats of the human alpha satellite array, whereas the older and more diversified satellite monomers comprise the flanking pericentric heterochromatin [27]. Analysis of the CenO satellites in centromeres of rice revealed segmental duplications, insertions and deletions, inversions, and reshuffling of variant satellite monomers [28]. Unequal exchange occurs at a high frequency between sister centromeres in mitotically cycling mouse (*Mus musculus*) chromosomes and is negatively regulated by DNA methylation, without which loss of repeats occurs [29]. However, it is unknown whether these recombination events can be transmitted through meiosis to the next generation. These observations provide evidence of extensive recombination in centromeres over evolutionary time scales and underscore the instability of repeat arrays to recombination and the necessity of suppressing crossing over in order to maintain centromere structure. How can this evidence for recombination in centromeres be reconciled with crossover suppression?

**Conversion in Centromeres**

In the same year that Sax predicted the centromere effect on crossing over, a new model of recombination, called gene...
conversion, was proposed to explain non-reciprocal recombination events in mosses and basidiomycetes [30]. Gene conversion is now thought to be a normal part of the homologous recombination pathway in which a programmed double-strand break in the DNA is repaired by copying a short (usually ~2 kb or less) stretch of the homologous chromosome. The resulting conversion event may then be resolved into a either a crossover or a noncrossover (Figure 3). Could noncrossover gene conversions contribute to recombination in centromeres in the absence of crossing over?

The localized nature of gene conversion makes it significantly more difficult to detect than crossing over. A key problem is the need for numerous closely spaced unique markers in the highly repetitive sequences of the centromere and pericentromere. Consequently this question has been most thoroughly addressed in budding yeast, which lacks centromeric and pericentromeric repeats. Most studies have concluded that gene conversion is moderately suppressed (4- to 7-fold) at yeast centromeres, along with crossing over [24,25]. However, initiating double-strand breaks are not found within the point centromeres, but rather nearby [31,32]. One study reported that when nearby conversion events were examined, the conversion tract frequently included part or all of the centromere, and concluded that conversion rates at centromeres were not different than in non-centromeric regions [33]. Thus, the small size of yeast centromeres means that the relationship between the kinetochore and suppression of gene conversion has remained ambiguous.

To determine whether gene conversion events can occur within large centromeres and provide the recombination events underlying both satellite homogenization and centromere diversity, a new report by Shi et al. [2] studied events within the centromeres of maize (Zea mays). They developed 238 centromeric markers based on insertion polymorphisms of the centromere-specific transposon CRM2 that map to all ten maize centromeres. To verify their centromeric location, centromeric chromatin was immunoprecipitated with an anti-CenH3 antibody. CenH3 is distributed discontinuously in maize centromeres [34] and only about 30% of CRM sequences can be immunoprecipitated with anti-CenH3 [34–36]. Markers were then assessed in two parental lines and in 94 recombinant inbred lines derived from their progeny. As expected, no crossovers were observed. However, in two cases a single marker from one parent was gained in a centromere with all markers of the other parent, indicating a conversion event. The formal possibility that these events represent double crossovers is unlikely given the failure to find single crossovers.

Shi et al. then proceeded to assess their marker set in 53 highly diverse inbred lines representing the diversity of maize and found widespread evidence for marker recombination since the origin of maize, perhaps 9,000 years ago [37]. They could distinguish between crossovers and noncrossover conversions by determining the linkage disequilibrium (LD), or tendency of markers in a population to occur together on the same chromosome. In crossing over, LD decreases with distance, whereas the short conversion tracts of noncrossovers show no relationship between LD and distance, because the conversion of one marker ordinarily has no effect on the coinheritance of its neighbors. No correlation was found between distance and LD in centromere 2, which has been fully sequenced [36], consistent with noncrossover conversion. Two population genetic methods gave similar estimates of the conversion rate of >1×10⁻³ conversions per marker per generation, a rate not dissimilar to one estimate for the conversion rate on the chromosome arms [39].

These results are significant both for understanding the regulation of recombination in maize and for understanding the evolution of centromeres. Except in yeasts, studying recombination in centromeres has hitherto been largely a matter of inferring the occurrence of ancient events based on present-day sequences. The results of Shi et al. show that it is possible to study centromeric recombination in action in a multicellular eukaryote. They also confirm that such recombination can take place between homologs and not solely between sisters, with implications for the creation and spread of new centromere variants. Meiotic recombination involves complete end-to-end pairing of homologs, whereas a gene conversion event requires only a local homologous interaction, and it is possible that the observed conversion events occurred during mitotic development rather than during meiosis (Figure 3). For example, the mitotic threads seen to connect human sister centromeres [4] might sometimes be resolved via breakage events that initiate repair by homologous recombination. By this scenario, the surprisingly high level of genetic exchange observed by Shi et al. might be a consequence of the many mitoses that occur for each meiotic generation within a maize lineage.

Widespread gene conversion might be a general feature of centromeres of multicellular eukaryotes. Human centromeres are composed of higher-order alpha satellite repeat arrays [27], and evidence for their periodic homogenization suggests an underlying gene conversion mechanism [39]. As is the case for unequal exchange between sisters, which is the most attractive explanation for the large expansions and contractions of alpha satellite repeat arrays, centromeric gene conversion challenges the widely held perception of centromeres as genetically stable regions of the genome. The actions of gene conversion and unequal exchange provide variation that makes possible Darwinian competition of centromeres that may lead to their rapid diversification [40]. Thus the problem of both homogenization and diversification of centromeres in the absence of crossovers can be resolved.

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