Initiation of Meiotic Homologous Recombination: Flexibility, Impact of Histone Modifications, and Chromatin Remodeling

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Meiotic recombination is initiated by the formation of DNA double-strand breaks (DSBs) catalyzed by the evolutionary conserved Spo11 protein and accessory factors. DSBs are nonrandomly distributed along the chromosomes displaying a significant (~400-fold) variation of frequencies, which ultimately establishes local and long-range “hot” and “cold” domains for recombination initiation. This remarkable patterning is set up within the chromatin context, involving multiple layers of biochemical activity. Predisposed chromatin accessibility, but also a range of transcription factors, chromatin remodelers, and histone modifiers likely promote local recruitment of DSB proteins, as well as mobilization, sliding, and eviction of nucleosomes before and after the occurrence of meiotic DSBs. Here, we assess our understanding of meiotic DSB formation and methods to change its patterning. We also synthesize current heterogeneous knowledge on how histone modifications and chromatin remodeling may impact this decisive step in meiotic recombination.

Sexual reproduction depends on halving the genome content of germ line cells and faithful chromosome transmission during meiosis to yield viable gametes. Meiosis comprises one round of DNA replication and two successive rounds of chromosome segregation, allowing the reduction of a diploid genome to produce haploid gametes (Fig. 1A).

Central to meiosis is the process of recombination between the paternal and maternal chromosomes (interhomolog recombination), which is crucial to enhance the genetic diversity of the gametes, but also for providing physical connections among homologs. These connections (i.e., chiasmata) ensure proper alignment of homologous chromosome pairs on the spindle, promoting proper reductional segregation following the regulated release of the sister chromatid cohesion among the duplicated chromosomal arms (Watanabe 2012). Defective meiotic recombination is a source of de novo germline mutations, abnormal genome content in gametes (the source of Down’s syndrome), and infertility. Not surprisingly, cells have developed a variety of mechanisms and tight controls to ensure sufficient and well-dis-
Figure 1. The stages and mechanisms of meiosis. (A) Diploid yeast cells initiate meiosis on nutrient depletion and in the presence of a nonfermentable carbon source. In mammals, the process is started by endocrine/paracrine/juxtacrine stimuli from the surrounding cell and tissue environment. Relevant molecular stages are indicated: S, meiotic replication; DSB, double-strand breaks; CO, crossover; SC, synaptonemal complex; MI, and MII (first and second meiotic divisions, respectively). Colors: parental homologous chromosomes (green and black), sister chromatid cohesion (red), SC (pink). (B) The mechanism of meiotic recombination. Stage 1: Initiation. DSBs are introduced by the Spo11 protein. Stage 2: DSB processing. Strand resection initiates to yield 3' single-stranded DNA (ssDNA) overhangs. One of the 3' ssDNA tails engages in strand invasion and a homology search of the homologous chromosome, resulting in single-end invasion (SEI) and D-loop intermediates. Stage 3: DSB repair. (Legend continues on following page.)
tributed meiotic recombination events within their genomes.

At the DNA level, meiotic recombination can be divided in four successive stages (Fig. 1B): (1) initiation, which consists in the formation of programmed DNA DSBs; (2) DSB processing, which yields the recombinogenic single-strand tails; (3) homologous DSB repair, which involves the homologous recombination pathway and several meiosis-specific differentiation modulations, which facilitate interhomolog interactions; and (4) intermediate resolution and dissolution, which allow the formation of local (up to a few kb in length) noncrossover (NCO) gene conversion events and reciprocal crossovers (COs).

Recently, local and genome-wide studies of normal and mutant cells have uncovered a remarkable variability in the number and positioning of recombination events per chromosome and cell, which reveals an impressive level of flexibility (Szekvölgyi and Nicolas 2010). In this review, we outline our understanding of the control of the initiation events, and how histone modifications and chromatin remodeling impact this initial step of meiotic recombination. The process of meiotic recombination and its relationship to change in chromosome structures and movements, allowing homolog alignment, pairing, and synapsis, is also reviewed in Zickler and Kleckner (2015) and Lam and Keeney (2015).

MEIOTIC DSBs AND RECOMBINATION EVENTS ARE NONRANDOMLY DISTRIBUTED

In all organisms, meiotic recombination is initiated by the formation of a large number of programmed DNA DSBs per cell, which are repaired primarily via recombination between nonsister chromatids (recombination initiation occurs after replication) to generate NCO and CO recombinant products (Fig. 1B). These DSBs are catalyzed by the evolutionarily conserved Spo11 protein (Bergerat et al. 1997). Besides Spo11, a number of accessory proteins are required for DSB formation. In Saccharomyces cerevisiae, 10 DSB proteins are known and functional, but sometimes sequence-divergent orthologs are gradually being identified in other organisms, including mammals. Yeast two-hybrid and co-immunoprecipitation (IP) studies identified several multiprotein subcomplexes, which are successively recruited to the meiotic chromosomes before DSB formation in triggered (Arora et al. 2004; Panizza et al. 2011). The biochemical/structural functions of these DSB protein complexes remains poorly understood. They play a role to select the potential DSB regions along the chromosomes, and ultimately contribute to the recruitment of Spo11 and trigger cleavage.

Spo11 is orthologous to the topoVI family of topoisomerase discovered in archaea, and consistently introduces DSBs by coupled trans-esterification reactions to form covalent tyrosyl-DNA linkages at the 5' termini of the broken DNA. Spo11 is then removed by endonucleolytic cleavage (Neale et al. 2005), liberating short Spo11-DNA oligonucleotide complexes and resected strands, which are further extended to generate recombinogenic 3' single-stranded tails. Over the years, meiotic DSBs have been mapped and quantified in yeast genomic DNA using a variety of approaches, including Southern blot analysis of chromosomal fragments.
or full-length chromosomes, PCR, chromatin immunoprecipitation (ChIP)-Chip, and ChIP-seq analyses of enriched single-stranded DNA (ssDNA) (Buhler et al. 2007) and, most recently, via the purification and sequencing of Spo11-associated oligos, providing exquisite resolution both in terms of mapping and quantification (Pan et al. 2011). The most amenable method to estimate the number of DSBs on a per-cell basis is to count the number of Rad51/Dmc1 immunostaining foci on spread meiotic nuclei (Bishop 1994). On a broad scale, these methods have provided consistent general conclusions. They show the evolutionary conservation of the mechanism of DSB formation and the essential role of interhomolog recombination for the production of viable, euploid meiotic products. Depending on the organism and mutant defect, a lack or reduction in DSB frequency can lead to arrest of meiotic progression and apoptosis, or progression and formation of inviable aneuploid gametes.

To address whether Spo11-catalyzed DSBs play a unique role in meiosis, the functionality of DSBs introduced by other nucleases and clastogenic agents has been examined. Interhomolog recombination was promoted by ionizing radiation (Bowring et al. 2006), indicating that Spo11-catalyzed DSBs are not unique in their ability to promote recombination during meiosis. Other studies used the expression of a heterologous endonuclease, such as HO (Kolodkin et al. 1986; Malkova et al. 2000), VDE (Hodgson et al. 2011), or I-SceI (Farah et al. 2009) in yeast, and the Mos1 transposase in *Caenorhabditis elegans* (Robert and Bessereau 2007). In all cases, interhomolog recombination was induced at the break sites. However, in several respects, the DSBs and recombinants induced by these systems are different than those resulting from Spo11. These site-specific DSBs are not bona fide Spo11 breaks because of (1) the uncontrolled timing of DSB induction, (2) the abnormal and dangerous cleavage of both sister chromatids, and (3) alteration in recombination efficiencies when assayed in the wild-type or Spo11-deficient strain background. This relates to the role of the Spo11 DSBs in the pairing of the homologs. The uniqueness of the Spo11 breaks might reside in its mode of cleavage that allows, like in other site-specific recombination processes, to intimately link break formation and processing, and also to avoid extensive DSB signaling associated with accidental DSBs (Borde et al. 2004). Furthermore, Spo11 break formation is intimately linked to the process of chromosome pairing that enforces interhomolog rather than intersister recombinational repair.

Extensive studies with *S. cerevisiae* (Baudat and Nicolas 1997; Gerton et al. 2000) and later with *Schizosaccharomyces pombe* (Cromie et al. 2007), *Arabidopsis* (Drouaud et al. 2013), mice (Smagulova et al. 2011), chimpanzee, and humans (Myers et al. 2010; Auton et al. 2012) showed that the distribution of meiotic DSBs is not random along chromosomes, explaining the longtime noted discrepancy between physical and genetic map distances. Nowadays, genetic distances are measured by high-throughput microarray or genome-wide sequencing analyses of meiotic progenies, starting from diploid heterozygous strains carrying thousands of single-nucleotide polymorphisms (SNPs). In a pioneer study in *S. cerevisiae*, Mancera et al. (2008) genotyped the four spores of several tetrads to reconstitute the meiotic recombination events per meiotic cell and, on average, found ~90 COs and 46 NCOs (66 after correction for undetected NCO) per meiosis. Consistently, these recombination events are located in correlation with the preferred sites of DSBs and, in a number, suggesting that a large majority of the DSBs (estimated to ~150 per meiosis) are repaired on the homologous chromosome. This is consistent with the single-site genetic and physical analysis of recombination intermediates at the most studied artificial *HIS4::LEU2* hot-spot, indicating that ~80% of the meiotic DSBs are repaired using a nonsister chromatid rather than the sister chromatid as template (Lao et al. 2013). Genome-wide analysis of meiotic products in other organisms confirms that most DSBs are repaired by interhomolog recombination (Cole et al. 2010), but the large excess of DSBs relative to COs—identified directly or by counting the Rad51 and Dmc1 foci on meiotic prophase spreads over the final recombination products—can be explained also if meiotic DSB
repair frequently occurs among sister chromatids. This raises the key question of how and to what extent the interhomolog bias is implemented and, globally, to what extent the meiotic versus mitotic differentiation of DSB repair template choice is organism specific.

**MULTIPLE LAYERS OF CONTROL SHAPE THE DSB DISTRIBUTION: CLUSTERING, INTERFERENCE, AND REDISTRIBUTION**

If recombination-initiation sites were randomly distributed, DSBs would be equally likely to occur at any location along the chromosomes and would not influence one another (Fig. 2A). This is clearly not the case because (1) DSB frequencies vary greatly from site to site (at least 400-fold in *S. cerevisiae*, *S. pombe*, and *Arabidopsis*) (Pan et al. 2011; Choi et al. 2013; Fowler et al. 2014). (2) DSB sites within a hotspot region are strongly localized. In *S. cerevisiae*, they are preferentially localized in intergenic promoter-containing intervals, but depleted from other regions (Baudat and Nicolas 1997). In *S. pombe* and mice, enhanced DSB formation occurs in various regions, but also alternate with regions showing low DSB activity. (3) Over longer distances, DSB-prone and -repressed regions are clustered in subchromosomal domains. The cold regions are found in interstitial coding regions, showing no specific pattern along the chromosomes, but DSB formation is always strongly suppressed in the telomere- and centromere-proximal regions, as well as within the recombinant DNA (rDNA). These quantitative and region-specific variations are indeed strong evidence for the nonrandom nature of the DSB distribution. The contribution of rare and, possibly, random (stochastic) DSBs is not excluded, but remains difficult to measure and map.

Figure 2A illustrates two hypothetical nonrandom modes of DSB distribution: (1) a uniform distribution, when every DSB falls as far from its neighbors as possible, and (2) clustered, when most DSBs are concentrated close together and large regions contain very few, if any, breaks. The observed DSB pattern in *S. cerevisiae* (Borde et al. 2009) illustrated for chromosome III falls between these two extremes; with some deviation, it falls closer to the clustered arrangement, indicating that the contribution of stochasticity might be significant. It is noteworthy that the current genome-wide DSB-mapping techniques (Spo11-oligo sequencing, Dmc1/Rpa/ssDNA ChIP) obscure cell-to-cell variations in DSB formation (as do all mass-biochemical methods that investigate cell populations).

Other strong driver elements that shape the nonrandom distribution of DSBs have been documented in *S. cerevisiae*, but also remain poorly understood. These phenomena are referred to as “DSB interference” and appear to drive even spacing of DSBs and limit simultaneous breakage at allelic positions. First, the occurrence of DSBs reduces the probability of nearby DSBs to occur (Fig. 2B). This is a “cis-inhibition” process, in which strong DSB hotspots suppress the activity of nearby recombination-initiation sites on the same homolog (Xu and Kleckner 1995; Fan et al. 1997). This inhibitory effect of natural or targeted DSBs on adjacent hotspots is known to spread over significant distances along the chromosomes (up to 25–100 kbp in *S. cerevisiae*) (Robine et al. 2007). The second interference phenomenon is that DSB formation on one homolog decreases DSB formation on the other homolog (trans-inhibition) at the cognate allelic position (Rocco et al. 1992; Zhang et al. 2011a).

Whether these cis- and trans-regulatory events occur independently remains to be elucidated, but, importantly, the trans-modulation appears to be genetically controlled, involving the two signal transduction kinases Mec1 (ATR) and Tel1 (ATM) acting as potential direct effectors of the trans-DSB interference (Zhang et al. 2011a; Blitzblau and Hochwagen 2013; Gray et al. 2013). The benefit to locally regulate DSB formation between the sister and nonsister chromatids is to prevent simultaneous DSB formation at the same place on more than one chromatid. This would generate poorly interacting broken molecules, perturb the efficiency of DSB repair, and, finally, facilitate mutagenic end-joining and out-of-register interactions between repeats, as well as genome rearrangements on ectopic interactions. Rare but naturally occurring events of this type may contribute to the arising of de novo germline mutations.

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Figure 2. Spatial distribution of meiotic DNA double-strand breaks (DSBs) along the chromosomes. (A) Hypothetical (blue) and real (black) DSB patterns, representing random and nonrandom distributions. Real DSBs (Borde et al. 2009) establish “cold” and “hot” domains, which alternate along the chromosomes length. (B) DSB interference establishes even spacing among interhomolog crossovers (COs). Interference nucleates at activated DSB sites (yellow rectangles) cleaved by Spo11 (red star) and spreads over significant distances along the sister/nonsister chromatids or homologous chromosome (trans-inhibition, blue triangles), or along the same sister chromatid (cis-inhibition, green triangles). DSBs form sequentially as meiosis progresses such that “late” DSBs fill in the gaps among “early” DSBs. A subset of DSBs matures into COs (blue crosses). (C) DSB redistribution can shuffle recombination-initiation events to reinforce the “obligatory” number of COs per chromosome. Mutant 1 can be, for example, an upstream activating sequence (UAS)_{GAL80} mutant, in which the Gal4 binding site was inactivated to prevent DNA cleavage by the Gal4BD-Spo11 fusion protein. New DSBs readily appear nearby the UAS. Mutant 2 corresponds to, for instance, a set1\(\Delta\) mutant, in which the disappearance of “canonical” hotspots often results in the activation of latent hotspots.
source of human diseases (Campbell and Eichler 2013). Also, cells with broken chromosomes may progress further into the meiotic prophase, and, unless they are arrested by the DNA damage checkpoint before MI, the chromosomes would segregate abnormally, yielding unbalanced genomes.

Another parameter that can result in DSB modulation is the unresolved question of the synchrony/ asynchrony of DSB formation in individual meioses. Time-course experiments in synchronously sporulating yeast cell populations and mutant cells that accumulate unrepaired DSBs (rad50S or sae2Δ) show that a majority of DSBs (~150 in S. cerevisiae) tend to occur within a 30- to 60-minute window, but the level of synchrony in these experiments remains insufficient to conclude whether the DSBs take place simultaneously or progressively in individual cells. This issue concomitantly raises the question of how the recruitment and biochemical cleavage activity of Spo11 are functionally and temporally regulated. Although Spo11’s covalent attachment to the cleaved strand is a suicide mechanism preventing the turnover of the protein, a regulatory pathway must also exist to locally and/or globally terminate the window for DSB formation. How this happens is not understood. The capacity of certain pachytene arrested mutants (e.g., ndt80Δ) to continue DSB formation (Xu and Kleckner 1995), and also the late persistence of Spo11 and other DSB proteins along the chromosomes after DSB formation (Arora et al. 2004; Cole et al. 2010; Gray et al. 2013), are puzzling observations that call to further integrate the connection of DSB formation and repair with changes in chromosome structures and movements and explore the underlying signaling pathway(s) in connection to the meiotic cell-cycle progression.

Another important DSB control can be called “DSB compensation” to denote the homeostatic process that is able to redistribute DSBs along the chromosomes (Fig. 2C). When strong DSB sites become suppressed, locally or genome-wide, as a result of a mutation, a vast number of dormant DSB hotspots get activated to compensate for the loss of DSBs. Consequently, the original DSB distribution can be dramatically remodeled. For instance, when the GAL4BD-Spo11 fusion protein (when the DNA-binding domain of the Gal4 transcription factor is fused with Spo11) (Pecina et al. 2002) targets DSBs near the UASGAL DNA-binding motives, mutating the UASGAL site at the GAL80 locus causes a severe drop of Gal4BD-Spo11-targeted DSB levels (from 10% to 3%) at the GAL80 promoter, but compensatory “weak” DSBs readily appear elsewhere, within the same intergenic region (Fig. 2C, middle track) (Robine et al. 2007). Also, null mutations of certain histone-modifying enzymes (e.g., set1Δ, sir2Δ, Prdm9−/−) significantly suppress the activity of canonical recombination hotspots, but a number of compensatory DSBs appear elsewhere (Fig. 2C, lower track) (Borde et al. 2009; Brick et al. 2012). For example, in Prdm9−/− knockout mice, lacking the sequence-targeted meiosis-specific histone-K4 methylase PRDM9, DSB sites are redirected to different genomic regions, which are flanked by PRDM9-independent H3K4me3 marks (Brick et al. 2012). In this way, “DSB compensation” can, at least in part, buffer the absence of natural (“frequent”) recombination sites to assure the minimal level of DSBs required for homolog pairing and reinforce the “obligatory” COs per chromosome (i.e., ≥1 CO/chromosome arm). It is noteworthy that canonical DSBs can be successfully compensated for in the absence of Set1, giving rise to viable spores (Acquaviva et al. 2013). On the contrary, Prdm9 mutants show severe defects in the progress of meiosis (Hayashi et al. 2005; Brick et al. 2012), indicating that these delocalized DSBs are, in some respects, different from naturally occurring DSBs. The differences between the physiological versus pathological outcomes for “relocalized” and “compensatory” DSBs need to be better understood, especially if they used to modify the natural landscape of meiotic recombination.

It is noteworthy that the total number of recombination events (overall recombination rate) and locations of COs (related to hotspot usage) are two distinct features, which are both determined genetically (Kong et al. 2014). For instance, single-nucleotide variants of PRDM9 affect both recombination rate and location.
Other genetic variants have been identified that alter the genetic map in humans: (1) CTCFL, the testis-specific paralog of CTCF, a Zn-finger protein organizing chromatin loops with cohesins (Sleutels et al. 2012), (2) the meiosis-specific cohesin RAD21L, which plays a role in the formation of meiotic chromosome axis and synapsis, (3) the SUMO ligase RNF212, ubiquitin ligase HEI10 (CCNB1IP1), and Mut5 homolog MSH4, all related to members of the ZMM class of pro-CO factors identified in S. cerevisiae, although with different synapsis defects (Reynolds et al. 2013; Kong et al. 2014; Qiao et al. 2014). Strikingly, significant differences are seen among these genetic variants, as those having a large impact on recombination rates have no effect on hotspot usage. This implies that, in most cases, genetic variants affect the CO or NCO decision rather than directly touching on DSB formation. Accordingly, the yeast Zip3 protein preferentially localizes to DSB hotspots that tend to be repaired as COs (whereas Zip3 is depleted from NCO-biased hotspots) (Serrentino et al. 2013), pointing to the existence of distinct types of DSB sites with regard to CO and NCO formation.

**Histone Modifications**

The chromatin-flanking DSB hotspots have a clear spatial organization of histone marks (Fig. 3A) such that H3K9ac falls closest to hotspots, followed by H3K4me3 concentrated at the +1, +2, +3 nucleosomes, whereas H3K4me1/me2, H3K36me3, H3K79me2, and H3R2me/as are mainly present inside or at the 3’ ends of open reading frames (ORFs) (Zhang et al. 2011b). Histone H3K56 acetylation is a transient mark enriched on nucleosomes that show rapid turnover kinetics (Rufiange et al. 2007; Watanabe et al. 2013). These chemical labels can impact meiotic recombination (1) by affecting the structure and mobility of nucleosome core particles, and (2) by interacting with chromatin-signaling proteins that use histone modifications as docking sites. Mishaps in the proper writing, reading, and erasing of these biochemical tags can interfere with recombination, leading to the formation of pathological diseases, such as infertility or carcinogenesis (Schwartzentruber et al. 2012). Here, we mention a few remarkable examples from various model organisms that exemplify the association between histone modifications and meiotic DSBs. (1) In C. elegans, acetylation of histone H2A at lysine 5 (H2AK5ac) seems to play a crucial role in meiotic DSB and CO formation (Wagner et al. 2010). Deletion of the XND-1 gene (X nondisjunction factor 1) disrupts H2AK5 acetylation and induces a significant change in the meiotic DSB and CO landscape, with most COs occurring abnormally within the gene-rich regions of autosomes. (2) In fission yeast, H3K9ac is specifically enriched at recombination hotspots, but the “prototypical” DSB mark H3K4me3 is less relevant (Yamada et al. 2013). Mutating the H3K9 residue mildly reduced levels of Rec12 binding (homologue of Spo11) and DSBs, indicating that H3K9ac may facilitate recombination initiation by stabilizing the contact between
Figure 3. Predominant positions of histone modifications and chromatin remodelers relative to nucleosome depleted regions (NDRs). (A) Histone modifications, and (B) chromatin remodeling factors flanking NDRs are shown. Arrowheads point to the net directionality of nucleosome movement performed by the corresponding remodeler protein. Ino80 spreads across many positions, but the Arp5 subunit is enriched at the +1 position. Rsc8 and Snf2 are enriched predominantly at the first three genic nucleosomes, but Snf2 is largely depleted at the −1 position and enriched at −2. Isw2 maps mainly to the +1 position and the regulatory subunit loc3 (ISW1a) is particularly enriched at the +1 position, whereas loc4 (ISW1b) is enriched at positions +2, +3, and +4.
Rec12 and DSB sites. (3) In budding yeast, deletion of the Sir2 histone deacetylase causes variable, but genome-wide changes in DSB formation, reducing or elevating DSB levels at ≈12% of hotspots (Mieczkowski et al. 2007). Sir2 could actively repress DSBs within naturally cold regions (such as the rDNA cluster, telomeres, centromeres) to prevent nonallelic homologous recombination. Interestingly, even in the presence of active Sir2, the edges of the rDNA array remain exceptionally susceptible to meiotic DSBs (Vader et al. 2011). It has turned out that a secondary border-specific system, involving the meiotic ATPase Pch2 and Orc1, operates at heterochromatin–euchromatin junctions to shield the edges of the rDNA array. Pch2 is an evolutionarily conserved AAA-ATPase protein, which appears to influence the initiation of recombination and timely meiotic progression (Borner et al. 2008).

A number of studies indicate that histone H3K4me is a fundamental mark of meiotic DSB formation, which is conserved from yeast to human: (1) deletion of the H3K4 methyltransferase Set1 or point mutation of the modifiable histone H3 lysine 4 residue severely reduce meiotic DSB levels in S. cerevisiae (Acquaviva et al. 2013; Sommermeyer et al. 2013); (2) absence of Set1 in the distantly related fission yeast partially reduces DSB formation at various loci, in which Set1 and H3K9 acetylation redundantly regulated meiotic DSB formation (Yamada et al. 2013); (3) deletion of RAD6, as well as the substitution of the ubiquitylation site on histone H2B, both of which affect H3K4 methylation, decrease DSB frequencies at various hotspots (Yamashita et al. 2004); (4) H3K4me3 levels were significantly enriched at several yeast (Robine et al. 2007) and mouse (Smagulova et al. 2011) DSB hotspots, as well as at Arabidopsis thaliana CO hotspots (Choi et al. 2013); and (5) from mice to human, hotspot activity is largely dependent on Prdm9, the sequence-targeted meiosis-specific H3K4 methyltransferase (Grey et al. 2011).

Our latest work (Acquaviva et al. 2013) has revealed a causative and unexpected link between the presence of histone H3 lysine 4 methylation and DSB formation. A genetically engineered Set1 histone methyltransferase (Gal4BD-Set1) targeted to recombination cold regions readily induces DSB formation at these sites (Fig. 4A). The DSB-inducing effect of Gal4BD-Set1 depends on the presence of the modifiable lysine 4 residue (because DSBs at these sites were abolished in the H3K4R mutant), revealing a cause–effect relationship between the presence of histone H3K4me and meiotic DSB formation. Unexpectedly, although Gal4BD-Set1-induced DSBs were strongly dependent on Spp1, the PHD-finger subunit of COMPASS (Fig. 4A), the reverse was not true: tethering of Spp1 to recombination cold spots (Gal4BD-Spp1) strongly induced DSB formation, but these breaks were largely independent of Set1 and histone H3K4me (Fig. 4B). Moreover, the Gal4BD-Spp1-induced DSBs were maintained in both set1Δ and H3K4R mutants, indicating that Spp1 on its own is able to initiate meiotic recombination when recruited to the chromosomes. The DSB-promoting effect of Spp1 is mediated via its PHD finger domain, binding to H3K4 trimethylated nucleosomes, and by its physical interaction with the “core” DSB protein Mer2 (Acquaviva et al. 2013; Sommermeyer et al. 2013). Collectively, these findings suggest that Spp1 makes a contact bridge between DSB hotspots and the chromosomal axis via contacts with Mer2 and histone H3K4me3, and contrast the situation in fission yeast, which uses a meiosis-specific bridge protein (liaisonin) to mediate axis–DSB hotspot interaction (Miyoshi et al. 2012).

It should also be noted that a subclass of budding yeast DSB cold spots, localized proximal to chromosomal axes, are associated with lower histone H3K4me3. In budding yeast, binding of Rec8, the kleisin subunit of meiotic cohesin, along chromosome axes plays a critical role in determining the canonical distribution of meiotic DSB sites (Kugou et al. 2009). Regions spanning ±0.8 kbp around axial Rec8 binding sites show lower Spo11-oligo frequency (Ito et al. 2014). Moreover, Spo11 fused with the Gal4 DNA-binding domain (Gal4BD-Spo11) cannot form meiotic DSBs efficiently when targeted to sites adjacent to Rec8 binding sites. In addition, H3K4me3 levels are remarkably lower in Rec8
binding sites. It is, thus, suggested that reduced histone H3K4me3 down-regulates Spo11 activity on sequences proximal to the axes. One possible mechanism is that the absence of H3K4me3 around the axis hampers formation of tethered axis-loop complexes resulting in local inhibition of DSBs.

Collectively, these recent studies strongly implicate COMPASS, H3K4me3, and Mer2 in the determination of DSB sites. However, there are several key unknowns to understand the relationship among meiotic DSB sites, histone modifications, and higher-order genome architecture. For example, (1) existence of axis-loop contacts and their role in DSB formation have not been proven at the molecular levels (physical evidence is missing); (2) it is not known which chromatin proteins and histone modifications, in particular, COMPASS, Mer2, and H3K4me3, make or participate in mediating these contacts (causality is missing); and (3) mechanisms for DSB site selection in the absence of H3K4 methylation and repression of Spo11 activity in Rec8 binding sites with lower H3K4me3 remain unknown. Even if not absolutely required, the broadly localized H3K4me3 modification has the virtue of permitting the initiation of recombination at numerous places of the genome, a molecular strategy that provides flexibility and ensures a large diversity of recombinant haplotypes to be transmitted by the gametes.

**CHROMATIN REMODELERS AND HISTONE MODIFICATIONS**

Chromatin remodeling factors (Table 1) can significantly accelerate the dynamics of nucleo-
somes to allow for more rapid and localized access of Spo11 to meiotic DSB sites. For instance, the fission yeast SWI/SNF-type ATP-dependent remodeler SNF22 and CHD-1-type ATP-dependent remodeler Hrp3 activate the M26 recombination hotspot before DSB formation (Yamada et al. 2004). During DSB repair in mitotically cycling cells, chromatin remodelers exhaustively participate in disrupting and mobilizing nucleosomes (Mellor and Morillon 2004; Seeber et al. 2013). (1) Physical tethering of INO80 (LexA-Ino80) to lacO-tagged DSBs enhances the mobility of the breaks, causing increased gene conversion rates at ectopic donor sites (Neumann et al. 2012); also, INO80 is recruited to DSBs by carboxy-terminally phosphorylated H2AX, in which it evicts/remodels the H2A.Z variant nucleosomes (Htz1 in yeast) to facilitate end resection. (2) Fun30 mediates the process of end resection after DSB induction (Chen et al. 2012). (3) Rad54 mediates strand-exchange reactions during Holliday junction formation and resolution (Nimonkar et al. 2012). Whether the same remodeling factors function in meiotic recombination remains unclear.

A number of observation points toward a functional link between chromatin remodelers and histone modifications (however, these relationships can be rather complex as the latter affects the sites of chromatin remodeling, and vice versa). (1) On meiotic DSB formation, there is the rapid appearance of open chromatin—showing increased DNaseI and MNase sensitivity at hotspots—promptly followed by γH2AX phosphorylation and histone H4 acetylation spreading over hundreds of kbps away from breaks (Ohta et al. 1994; Downs et al. 2004). (2) Histone H3K56 acetylation (H3K56ac) and SWR1 are mechanistically coupled as acetylated H3K56 modulates the specificity of SWR1 to remove the histone variant H2A.Z from gene regulatory regions (Watanabe et al. 2013). It is also known that A. thaliana meiotic DSB sites significantly overlap with H2A.Z nucleosomes and H3K4me3 (similarly to S. cerevisiae) and the SWR1-deposited H2A.Z nucleosomes promote meiotic DSB formation and repair (Choi et al. 2013). (3) Members of the SWI/SNF family have carboxy-terminal bromodomains, which interact with acetylated histones, to preferentially target the acetylated nucleosomes for eviction (Yodh 2013). (4) The CHD remodelers have two tandem chromodomains, recognizing methylated histone H3 tails.

Bai et al. 2011 has screened ~6000 S. cerevisiae nucleosome-depleted regions (NDRs) for consensus binding sites of known nucleosome-depleting factors (NDFs), including remodelers (e.g., Rsc3) and transcription factors (e.g., Abf1, Rap1, Reb1). A significant fraction of NDRs (30%) contained at least one NDF binding site. Similar observations were published in S. pombe (De et al. 2012), which together suggest that different NDRs depend on the modular binding of NDFs (remodelers, transcription factors, and histone modifiers). As not all NDRs constitute a functional recombination hotspot (Pan et al. 2011), it is easy to envisage that only a combinatorial association of these factors makes an NDR competent for recruiting the Spo11 machinery to initiate homologous recombination.

### Table 1. Chromatin remodeling factors

<table>
<thead>
<tr>
<th>Yeast</th>
<th>SWI/SNF</th>
<th>ISWI</th>
<th>CHD</th>
<th>INO80/SWR1</th>
</tr>
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<tbody>
<tr>
<td><em>Saccharomyces cerevisiae</em></td>
<td>SWI/SNF</td>
<td>ISWI1a</td>
<td>CHD1</td>
<td>INO80/WR1</td>
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<td></td>
<td>RSC</td>
<td>ISWI1b</td>
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<td>ISW2</td>
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<td></td>
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<td>ACF/WCFR</td>
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<tr>
<td><em>Homo sapiens</em></td>
<td>SWI/SNF</td>
<td>CHRAC</td>
<td>NuRD</td>
<td>INO80/SRCAP</td>
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<td>WICH</td>
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*Yeast and human chromatin remodeling complexes classified by their ATPase subunits. CHD, chromo-helicase/ATPase DNA binding.*
In a systematic analysis, Yen et al. (2013) has revealed that remodelers bind to DNA in a nucleosome-position and orientation-specific manner. Nucleosomes are mobilized with a predefined net directionality relative to NDRs (toward or away from them), such that (1) remodelers bind predominantly to the +1 nucleosome, or (2) they are located at multiple positions along the ORF (Fig. 3B). For example, Ino80 spreads across many positions, but the Arp5 subunit of the same INO80 complex is enriched at the +1 position, moving this nucleosome 5′ to 3′. In line with the above, the RSC (remodels the structure of chromatin) remodeling complex plays a highly specific role in the precise positioning of nucleosomes over yeast promoter, a function that cannot be replaced by other closely related remodeling enzymes (Wippo et al. 2011). All of the above collectively suggest that +1 nucleosome-flanking NDRs are differentially processed by remodelers such that the initial positioning of this particular nucleosome may automatically cause the positioning of adjacent nucleosomes. Therefore, NDRs, in which most meiotic DSBs fall in S. cerevisiae and Arabidopsis, might be the active organizing centers (not just simple bystanders) of DNA cleavage in meiosis.

CONCLUDING REMARKS

Numerous studies have suggested that post-translational histone modifications do not represent a “code” (Sims and Reinberg 2008; Lee et al. 2010), but that the majority of these chemical tags mobilize or immobilize, rather than mark, the nucleosomes. Because of the stringent spatial order of histone modifications that flank meiotic DSB sites, many aspects of meiotic DSB control, including hotspot intensity, clustering/interference, and compensation, might operate at the level of the well-positioned +1 nucleosomes bordering DSB hotspots. Indeed, binding of Prdm9 to DSB sites actively reorganizes the flanking +1 nucleosomes, creating a symmetrical pattern of NDRs centered over the 13-mer consensus binding motif of the PRDM9 zinc-finger array (Baker et al. 2014). In this way, H3K4 methylated nucleosomes establish a permissible chromatin structure for meiotic DSB formation, depending on the activity of PRDM9.

In conclusion, the +1 nucleosome-flanking meiotic DSB hotspots seem to be critical for meiotic recombination initiation and repair. We propose that a combinatorial association of histone modifications and nucleosome remodelers positively (or negatively) regulates the turnover/mobility of the critical +1 nucleosome. This, in turn, automatically positions the adjacent nucleosomes to establish a permissive (or restrictive) chromatin context for recombination. The resultant effect of these activities is expected to modulate the efficiency of Spo11-mediated DNA cleavage at meiotic recombination hotspots.

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Lóránt Székvölgyi, Kunihiro Ohta and Alain Nicolas

Cold Spring Harb Perspect Biol 2015; doi: 10.1101/cshperspect.a016527

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