During prophase of meiosis I, homologous chromosomes interact and undergo recombination. Successful completion of these processes is required in order for the homologous chromosomes to mount the meiotic spindle as a pair. The organization of the chromosomes into pairs ensures orderly segregation to opposite poles of the dividing cell, such that each gamete receives one copy of each chromosome. Chiasmata, the cytological manifestation of crossover products of recombination, physically connect the homologs in pairs, providing a linkage that facilitates their segregation. Consequently, mutations that reduce the level of recombination are invariably associated with increased errors in meiotic chromosome segregation. In this review, we focus on recent biochemical and genetic advances in elucidating the mechanisms of meiotic DNA strand exchange catalyzed by the Dmc1 protein. We also discuss the mode by which two recombination mediators, Hop2 and Mnd1, facilitate rate-limiting steps of DNA strand exchange catalyzed by Dmc1.
features that distinguish them from mitotic recombination products. Several of these features are promoted by the recombinases and their ancillary proteins, which are critical for regulating homologous chromosome behavior during meiosis.

**Meiotic recombination**

The process of DSB repair in mammals appears to utilize pathways similar to those seen in lower eukaryotes, such as *Saccharomyces cerevisiae* (budding yeast) [2,3], and the final products are either crossovers, which involves exchange of flanking DNA markers between the homologs, or non-crossovers, in which the flanking DNA remains unchanged [2]. In meiosis, the initial steps of HR involve introduction of DSBs at multiple chromosomal DNA sites catalyzed by the Spo11 protein [4] (Fig. 2). This topoisomerase-like reaction cuts DNA to generate a covalent protein–DNA linkage to the 5′ DNA ends on either side of the break. After Spo11 is removed from the DNA ends, the process of HR involves exonuclease activity to generate 3′ single-stranded DNA (ssDNA) tails [5,6]. After resection, two eukaryotic members of the RecA protein family, the ubiquitously expressed Rad51 DNA recombinase and the meiosis-specific Dmc1 DNA recombinase, bind the 3′ ssDNA tails to form helical nucleoprotein filaments, which perform a search for intact homologous double-stranded DNA (dsDNA) [7]. Here we use the term homologous to describe DNA sequence similarity. It should be noted that this term is also often used with a different genetic meaning, i.e. homologous pairs of chromatids. Once the homologous sequence is found, the recombinases promote invasion of the ssDNA ends into the homologous duplex DNA (D-loops). After strand exchange, current models propose that HR intermediates are processed by one of two distinct pathways. The initial and relatively unstable strand invasion intermediates

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**Fig. 1.** Homologous chromosome association and dysjunction. (A) Schematic of male mouse spermatogenesis. Note the period of prophase during which HR promotes exchange of genetic information and homologous chromosomes pairwise interactions. (B) The presence of chiasmata ensures that each chromosome of the homologous chromosome pair segregates to opposite poles of the spindle. The homologous chromosome pairs are represented in red and yellow. Each of the homologous chromosomes comprises two sister chromatids represented by two bars of the same color. The green ovals represent cohesins.

**Fig. 2.** The pathway of meiotic recombination. Copies of homologous chromosomes are represented in red and blue. HR proceeds by two pathways: synthesis-dependent strand annealing (SDSA) and double-strand break repair (DSBR). While SDSA only produces non-crossovers, the second DSB end is captured during DSBR. After DNA synthesis and ligation, Holliday junctions are formed whose resolution results in formation of mostly crossovers and a small proportion of non-crossovers. Formation of both crossovers and non-crossovers between homolog chromatids is shown. Although use of the homologous chromatids is favored during meiotic recombination, final genetic products of recombination may also be generated using sister chromatids as DNA repair templates.
may be displaced from the invaded homolog and anneal to the second single-stranded end of the break. This leads to re-joining of the broken chromosome by synthesis-dependent strand annealing (SDSA) to generate non-crossovers (Fig. 2, right branch) [2,8]. In an optional pathway, they are processed by double-strand break repair (DSBR) [8,9], which includes DNA polymerase-dependent heteroduplex extension synthesis facilitated by Hfl1/Mer3, resulting in relatively more stable strand invasion [10,11]. This alternative process is able to perform the second end capture, and leads to formation of double Holliday junctions. During and after the formation of joint molecules and DNA synthesis to restore sequences that were lost or damaged at the site of the original DSB lesion, joint molecules must be resolved to allow chromosome segregation and formation of chiasmata. The structure of the joint molecules dictates whether a DNA helicase, endonuclease, or a combination of both is required for resolution resulting in the formation of crossovers and non-crossovers [12]. Whereas DSBR and SDSA occur both in cells that divide through mitosis and in cells that divide through meiosis, the major pathway for repair DSBs in mitosis appears to be the SDSA pathway, with DSBR primarily occurring in meiosis [13]. During mitotic recombination, the recipient DNA duplex is generally a sister chromatid. In meiosis, however, the situation is more complex, as either the homolog chromatid or the sister chromatid may provide the template for repair (i.e. using either DSBR or SDSA). It has been suggested that the preferred meiotic inter-homolog recombination is promoted by meiosis-specific components that inhibit inter-sister chromatid recombination [14]. Meiotic double Holliday junction intermediates (which are ultimately resolved as crossovers) are essential for the proper segregation of chromosomes. These crossovers also play an important role by shuffling parental genomes, generating genetic diversity.

Dmc1 is at the center of meiotic recombination

DMC1 was first identified in a screen for genes specific to S. cerevisiae meiosis [15], and is present in almost all eukaryotes, including mice and humans [16]. Deletion of DMC1 in budding yeast, plants and mice results in severe abnormalities that reflect an indispensable role of this protein in meiotic recombination [15,17,18]. S. cerevisiae and mouse DMC1 mutants show a near-complete block of recombination [15,17], with the S. cerevisiae Dmc1-deficient strain showing no stable strand invasion or double Holliday junction formation [9,19], accompanied by defective synaptonemal complex formation. It has been suggested that Dmc1 promotes recombination almost exclusively between the homologous chromosomes [20], which is unique to meiosis [19]. However, Dmc1 is not required in all organisms. For example, Drosophila melanogaster and Caenorhabditis elegans have Rad51, but lack Dmc1, and, in Schizosaccharomyces pombe (fission yeast), a mutation of DMC1 does not completely abolish meiotic recombination [21,22].

Dmc1-mediated strand invasion and DNA strand exchange

Whereas in vivo studies have revealed the indispensable role of the Dmc1 protein in meiotic recombination, the biochemical mechanism of action of Dmc1 is better studied using purified systems in vitro. Given its homology to RecA and Rad51, Dmc1 was predicted to exhibit the hallmarks of the reactions by which RecA family members promote the recognition of homology between ssDNA and a duplex DNA template to promote the formation of joint molecules (Fig. 2) [23]. Pioneering work, using purified human and mouse proteins, described essential structural characteristics [24] and enzymatic activities of Dmc1 [25,26]. Purified recombinant enzyme had a DNA-dependent ATPase activity, binds preferentially to ssDNA, and catalyzes formation of D-loops in super-helical DNA (strand invasion).

Pre-synaptic and synaptic events of recombination promoted by Dmc1: a biochemical view

Figure 3A shows the molecular events leading to strand exchange promoted by RecA-related proteins. It illustrates the obligatory phases and experimental approaches used to detect and characterize the molecular intermediates. In trying to understand the mechanism by which Dmc1 and other RecA-like recombinases promote strand exchange, it is useful to distinguish four consecutive steps [27–30]. The first two are nucleoprotein filament assembly and conjoining of DNAs. Formation of these two intermediates requires no DNA homology, but they are a prerequisite to assemble the machinery promoting the subsequent steps of DNA homology search and strand exchange. During nucleoprotein filament assembly (the pre-synaptic phase), Dmc1 assembles on ssDNA, creating a helical nucleoprotein filament in which the RNA is stretched and under-wound relative to the B-form DNA. Nucleation of Dmc1 and Rad51 onto ssDNA is a slow process, which renders the pre-synaptic filament assembly prone to degradation by nucleas-
es [15,19,31] or interference by other single-strand binding proteins, such as amounts of replication protein A (RPA) that are sufficient to saturate available ssDNA [32–34]. Certain recombinase accessory factors facilitate the assembly of the Dmc1 pre-synaptic filament. As such, these recombination mediators are critical for the efficiency of homologous recombination. We expand on the mechanism of action of some of these recombination mediators below. Formation of a ternary complex of the Dmc1–ssDNA and Rad51–ssDNA filaments with dsDNA (conjoining of DNAs) initiates the synaptic phase. Here, we use the term ‘synapsis’ to describe the interaction of homologous DNA molecules. It should be noted that this term is also used with a different cytological meaning, which refers to the formation of the synaptonemal complex and consequently close juxtaposition of the homologous chromosomes. The synaptic phase has been proposed to be of critical importance in the reaction of homologous pairing catalyzed by RecA [35], Rad51 [36] and Dmc1 [30]. Juxtaposition of three DNA strands within the synaptic filament permits rapid
During a homology search, segments of the intact Dmc1 mechanism of homology search are used to study homology-independent conjoining of DNA molecules [30,36]. Several experimental approaches have been developed using purified systems that recapitulate all steps leading to strand exchange. For example, recombinase-dependent pairing of two homologous DNAs or synaptic complex formation (Fig. 3B,C) [30,37,38] may be used to study the mechanism of homology search (see below). In the D-loop assay, a short radiolabeled oligonucleotide is used to detect invasion of ssDNA into negatively supercoiled duplex, displacing the non-complementary strand into a D-loop structure (Fig. 3D) [39]. The ssDNA in the D-loop assay represents the ssDNA tail of the resected DSBs, and the dsDNA plasmid with homology to the ssDNA represents the targeted sequence. In the oligonucleotide-based version of another common assay (DNA strand exchange), one strand of a linear double-stranded oligonucleotide is replaced by a labeled linear single-stranded nucleoprotein complex, creating a labeled duplex and a linear ssDNA (Fig. 3E) [40,41]. Methodologies have been also devised to evaluate less stable conformational intermediates that lead to strand invasion: assembly and stability of nucleoprotein filaments formed by the recombinase and ssDNA may be studied by electron microscopy [24,26,42–44], DNA binding shift assay, surface plasmon resonance (for precise measurement of the kinetics of DNA binding) [30], exonuclease I protection [30,45,46] and recombinase turnover from ssDNA [36], while a dsDNA capture assay may be used to study homology-independent conjoining of DNA molecules [30,36].

**Dmc1 mechanism of homology search**

During a homology search, segments of the intact duplex DNA are bound by the Dmc1–ssDNA nucleoprotein filaments and tested reiteratively until homology is found. While it is still unclear how Dmc1 and other members of the RecA family of proteins perform genome-wide homology searches, several groups have shown that, upon identification of homology in the duplex DNA molecule, the pre-synaptic filament of RecA is able to form a stable synaptic complex consisting of three strands and the recombinase, in which strand exchange has already taken place [47,48]. In this complex, the invading ssDNA is part of the new duplex, and the leaving strand has not yet been released. An important question here relates to the steps that lead to this exchange intermediate? Studies of RecA [48,49], Rad51 [50,51] and Dmc1 [38] have shown that formation of a synaptic complex by any of these recombinases includes the transition through several slower conformational changes, such as the later stages of homology recognition, which involve localized melting (base flipping) and annealing (switching) at A.T-rich regions. In these experiments, a fluorescein-labeled ssDNA oligonucleotide (negative strand) in combination with a rhodamine-labeled homologous dsDNA is used to measure fluorescence resonance energy transfer (Fig. 3C). In synaptic complex formation, a ternary complex (three-strand) intermediate forms, bringing fluorescein and rhodamine into close proximity. In this state, the fluorescent dyes undergo fluorescence resonance energy transfer and fluorescein is quenched. This is detected as a reduction in the fluorescein-sensitized emission, and indicates formation of the first interactions between homologous DNA strands. In a synaptic complex formed with the recombinase (Dmc1, Rad51 or RecA), substitution of inosine for guanine (which destabilizes the duplex DNA that stimulates strand exchange) demonstrated a general effect of helix stability on recognition of homology, and A.T mismatches demonstrated a special role of A.T base pairs in recognition of homology [38]. This suggests that the dynamic structure of the double helix (DNA ‘breathing’) significantly contributes to recognition of homology. In this case, it is possible that the extended conformation of the recombinase–ssDNA pre-synaptic filament allows rotational mobility of the bases, making them available to test interactions through collisions with transiently opened base pairs in the DNA duplex.

**Dmc1 versus Rad51 in meiotic recombination**

What is the functional relationship between Rad51 and Dmc1 during normal meiosis? From genetic experiments in budding yeast, it is known that, when Rad51 and Dmc1 work together, the repair of DSBs is prefer-
entially directed to strand exchange between homologous chromosomes rather than sister chromatids. In addition, a Rad51 deletion leads to a notable reduction of inter-homolog recombination, and inter-sister chromatid repair prevails. In the absence of Dmc1, however, a dramatic reduction of both inter-sister and inter-homolog recombination is observed. In the latter case, the strand exchange activity of budding yeast Rad51 is inhibited by the Hed1 protein and the effector kinase Mek1. Removing this inhibition allows efficient recombination, although inter-homolog crossovers are reduced compared to wild-type. Two recent studies in *S. cerevisiae* reveal that the favored inter-homolog strand exchange activity mediated by Dmc1 requires inhibition of Rad51 strand exchange activity [52,53]. These studies imply that the inhibitory action of Hed1 on Rad51 recombinational activity converts Rad51 from a recombination enzyme to a recombination mediator. In agreement with this idea, it has been shown that, in meiosis, the prominent strand exchange activity is exhibited by Dmc1, not Rad51 [54]. In a series of cleverly designed experiments, the authors used a separation-of-function mutant to show that the ability of Rad51 to interact with DNA and form nucleoprotein filaments, but not its strand exchange activity, is a prerequisite for normal meiotic recombination. In the same study, experiments with purified proteins showed that Rad51 is very efficient in stimulating the strand exchange activity of Dmc1.

What are the intrinsic biochemical properties that functionally distinguish Dmc1 from Rad51? Dmc1 and Rad51 share 54% amino acid identity in humans, 52% in mouse, and 45% in yeast. *In vitro*, both purified recombinant Rad51 and Dmc1 bind ssDNA to form helical nucleoprotein filaments and promote DNA strand exchange. Despite similarities in the general mechanism of recombination, Dmc1 and Rad51 show differences in a number of structural/biochemical properties. For example, DNA unstacking, and consequently reactivity toward chemical modification of thymines of Rad51/Dmc1 nucleoprotein complexes, are notably different [46]. Differences have been also observed between Dmc1–ssDNA and Rad51–ssDNA nucleoprotein filaments, although the reported number of Dmc1 and Rad51 promoters per helical turn varies between reports [42,55,56]. Dissimilarities in the structure of Rad51 versus Dmc1 nucleoprotein complexes may account for the increased resistance of native Dmc1 D–loops compared with Rad51 D–loops to dissociation by DNA translocases, such as bloom syndrome protein and Rad54 [46]. This difference in stability indicates a biochemical distinction between intermediates of recombination catalyzed by Rad51 and Dmc1. In addition to the differences stated above, the rate of ATP binding and hydrolysis, protein polymerization rate and the kinetics of DNA binding and dissociation are critical biochemical properties that have not yet been fully investigated in a comparative fashion and may account for the differences in Dmc1 and Rad51 functions. In an alternative view, unique meiotic functions for Rad51 and Dmc1 are more likely to result from the influence of distinct sets of accessory proteins than intrinsic differences in their biochemical properties.

**Recombination mediators and Dmc1-promoted DNA strand exchange**

Proper function of Dmc1 *in vitro* and *in vivo* requires interactions with several meiotic accessory proteins [7,57,58]. These protein factors determine regulatory mechanisms that direct the choices of Dmc1 DNA repair pathways. Biochemically, they stimulate critical steps of the recombination mechanism catalyzed by Dmc1, such as the dynamics and targeting of filament assembly on ssDNA, the choice of homologous DNA for strand invasion, and protection of Dmc1 removal from DNA by factors such as helicases. Understanding HR requires a detailed understanding of the identities and activities of these accessory proteins. Here, we present an up-to-date overview of how one of these factors, the Hop2–Mnd1 complex, stimulates Dmc1.

**Hop2 and Mnd1 have a dual role in recombinase enhancement**

The *MND1* gene was first described in a screen for genes with meiosis-specific expression in *S. cerevisiae* [59], and in the null mutant strain, cells initiate recombination, but do not form heteroduplex DNA and exhibit hyper-resected DSBs [60]. This suggests that Mnd1 may be involved in strand exchange. Similarly, depletion of *MCP7*, the *S. pombe* ortholog of *MND1*, resulted in cell arrest in meiotic prophase, with a reduction in recombination rates [61]. In higher eukaryotes, the *MND1* gene is required for normal male and female fertility. For example, in mouse [41] and *Arabidopsis thaliana* [62,63], mutation of the *MND1* gene results in normal recombination initiation, but meiotic DSBs are abnormally repaired, accompanied by aberrant chromosome synopsis. Similar to the *MND1* mutant phenotype and consistent with a defect in a Hop2-dependent step during meiotic recombination, *S. cerevisiae* [64] and *A. thaliana* *HOP2* deletion mutants exhibit a profound failure in
meiosis, due to uniform arrest at meiosis I, with chromosomes engaged at synapsis with non-homologous partners [65]. Analysis of mouse HOP2 knockout spermatocytes suggests that DSBs are created and processed, but their repair is abnormal as indicated by the accumulation of Dmc1 and Rad51 proteins at DNA repair sites [66]. In sum, genetic and cellular analysis of deletion mutants in various species suggests that Hop2 and Mnd1 act in the same pathway of recombination, and the proteins have a conserved role in efficient DSB repair and normal homologous chromosome synapsis. The functions of Hop2 and Mnd1 proteins when they act together as a complex have been studied more extensively. The functional interaction between Hop2 and Mnd1 was first suggested in studies of budding yeast, in which MND1 acts as a multi-copy suppressor of a HOP2 mutation defect in viable spore production, and these proteins co-immunoprecipitate from meiotic cell extracts [67]. The cooperation between Hop2/Mnd1 and Dmc1/Rad51 is likely to be crucial in vivo. For example, in mice lacking Hop2 and/or Mnd1, progression of recombination is impaired immediately after Dmc1 and Rad51 are loaded onto the end of DSBs [41,66]. Finally, in all organisms analyzed so far, HOP2 and MND1 only appear in those genomes that carry DMC1.

In vitro, Hop2 shows two distinctive activities. First, when it is incorporated into a Hop2–Mnd1 complex, it stimulates Dmc1/Rad51-promoted recombination. This appears to be a prominent activity of Hop2 that has so far been observed using purified recombinant proteins from mouse [36,68–71], budding yeast [72], fission yeast [73], and A. thaliana [74]. Second, purified mouse Hop2 alone and independently of Dmc1 and Rad51 is capable of catalyzing strand invasion [41,68,69]. Although this intrinsic recombinational activity of Hop2 shares mechanistic signatures characteristic of the mammalian RecA-like recombinases, it shows distinctive characteristics. For example, Hop2-mediated strand exchange does not require ATP, and, in contrast to Dmc1, joint molecules formed by Hop2 are more sensitive to mismatches and are efficiently disassociated by the branch migration protein Rad54 [41]. Recent work in mouse spermatocytes suggested that Hop2 may work alone as a recombinase [41]. The authors reason that Dmc1 and Rad51 are inactive in the absence of the Hop2–Mnd1 complex. Therefore, deletion of Mnd1 in mouse spermatocytes leaves Hop2 as the only protein with recombinase activity. In agreement with this possibility, a proportion of Mnd1 knockout spermatocytes show a significantly high level of DSB repair (monitored by histone γ-H2AX and Dmc1/Rad51) and chromosome synapsis. Although these results indicate that DSB repair catalyzed solely by Hop2 may promote homologous chromosome pairing and synapsis, further evidence is required to demonstrate that Hop2 performs homology search and strand exchange in vivo in the absence of Dmc1 and Rad51 (i.e. analysis of Rad51−/−/Dmc1−/−/Mnd1−/− mutant spermatocytes is required).

In the context of Hop2 functions, a possible mode of action for the Mnd1 protein has been revealed, as interaction of Mnd1 with Hop2 down-regulates the D-loop formation activity of Hop2. Interestingly, Mnd1 inhibits the recombinase activity of Hop2, and, when incorporated into the Hop2–Mnd1 complex, promotes strand invasion mediated by Dmc1 and Rad51. It is proposed that Mnd1 works by producing changes in the biochemical properties and oligomerization state of Hop2. These changes result in a new molecular interface in the Hop2–Mnd1 complex that is responsible for Hop2–Mnd1 interaction and stimulation of the Dmc1 and Rad51 recombinases [68].

What is the molecular mechanism directing Hop2–Mnd1 complex stimulation of recombinase-mediated strand exchange? As stated above, efficient strand exchange promoted by Dmc1 and Rad51 requires formation of pre-synaptic ssDNA–Dmc1 nucleoprotein filaments and adjoining of homologous dsDNA and ssDNA. The Hop2–Mnd1 complex acts in these two critical stages of the recombination reaction [30,36]. First, purified Hop2–Mnd1 binds and stabilizes the pre-synaptic filament formed by Dmc1/Rad51 on ssDNA. In a second reaction, Hop2–Mnd1 enhances the ability of the recombinase–ssDNA nucleoprotein filament to capture the duplex target DNA (Fig. 4). Although Hop2–Mnd1 stimulation of the duplex DNA capture is homology-independent, this step is of paramount importance. Such capture is vital for promoting recombinase-mediated DNA pairing. In this reaction, it is proposed that unaligned DNA molecules transition to an aligned ternary complex that facilitates the homology search. Although strong in vitro evidence exists to support a bipartite action of Hop2 on both stabilization and the ability of a recombinase–ssDNA nucleoprotein filament to capture dsDNA, it remains to be determined whether these Hop2–Mnd1 activities contribute to the function of Dmc1 and Rad51 in vivo.

The underlying molecular basis for this bipartite action of Hop2–Mnd1, i.e. in both stabilization of the Dmc1 pre-synaptic filament and assembly of the synaptic complex, has been further clarified in recent work. One of these reports utilized a combination of structural and biochemical approaches to show that the heterodimeric Hop2–Mnd1 complex is a V-shaped
molecule, and that the dsDNA binding functions of the N-termini of Hop2 and Mnd1 work together to promote synaptic complex assembly, whereas the Hop2 C-terminus, which binds ssDNA, participates in stabilization of the Dmc1–ssDNA filament [71]. The biochemical function of the Hop2 N-terminus is highlighted by the fact that this domain is a typical winged helix DNA-binding domain with specific amino acids involved in dsDNA coordination [75]. The second report provides more mechanistic insights, showing that Hop2–Mnd1 has the ability to stimulate DNA strand exchange by modulating a range of Rad51 basic properties, particularly nucleotide and DNA binding [76]. It has been shown that Hop2–Mnd1 enables Rad51 DNA strand exchange, even in the absence of divalent metal ions required for ATP binding. In addition, Hop2–Mnd1 acts in two steps of the Rad51-mediated recombination mechanism. First, during nucleoprotein formation, Hop2–Mnd1 helps to load Rad51 on ssDNA, restricting its dsDNA binding ability. Second, it promotes dsDNA binding during the homology search, by removing the inhibitory effect of ssDNA.

**Hop2 as a potential tumor suppressor gene**

To date, the number of ovarian and breast cancer susceptibility genes identified accounts for less than half of hereditary breast and ovarian cancers. The finding of germline mutations in BRCA2 and related genes, such as PALB2, RAD51C and RAD51D, suggests that DNA recombination repair pathways are linked to breast and ovarian cancer genes. The recent finding of a family tied to ovarian dysgenesis, carrying a deletion of Glu201 in the C-terminal acidic domain of Hop2, supports the critical role for this protein in ovarian development [77]. Subsequent screening of germline mutations in familial and early-onset breast and ovarian cancers detected several mutations affecting Hop2 [78,79]. Interestingly, some of these mutations resulted in truncated versions of Hop2 with dominant-negative activity. Mutations in the HOP2 gene that de-regulated alternative splicing in cells derived from familial ovarian and breast cancer patients have also been found. Some of these splice variants act as dominant-negative mutants that abolish Rad51 foci formation during radiation-induced DNA damage. Together with results showing that constitutive expression of abnormal splice variants of HOP2 induces tumor growth in nude mice [79], these reports strongly suggest a role for inactivating HOP2 mutations in familial and early-onset breast and ovarian cancers.

**Zebrafish, an emerging model for studies of recombination in meiosis**

Current understanding of the genetic controls involved in meiotic recombination of eukaryotes primarily comes from studies of model organisms such as yeast. However, these studies have some limitations because many genes required for meiosis in higher eukaryotes have no orthologs in yeast, and there are clear meiotic differences between mouse and yeast [80–82]. Alternatively, a mouse spermatocyte system, a model that more closely resembles humans, has proved an excellent model, advancing the field by allowing high-resolution observation of chromosome structures that is not possible in simple systems. A forward genetic screen in the mouse demonstrated that many vertebrate meiosis genes are yet to be found [82]. By screening over 17 000 N-ethyl-N-nitrosourea-mutagenized mice, the Reproductive Genomics Group at Jackson Laboratory identified 44 mutant lines with reduced fertility. While many of the mutated genes from this screen have not yet been identified, this great effort revealed novel meiotic functions for six proteins [82]. The discovery of the first meiosis gene from this screen, Mei1, exemplifies the advantage of screening
for meiosis genes in vertebrates [83]. Although it provided valuable new information and tools, the Jackson Laboratory screen also suggested that it will be very difficult to uncover all genes required for vertebrate meiosis through forward genetics studies in the mouse.

Reverse genetics studies in the mouse have been very effective in identifying factors necessary for vertebrate meiosis. Vertebrate meiosis gene candidates identified through genetic screens in lower organisms, gene expression analyses, or protein interaction studies have been functionally analyzed with great success using engineered mouse mutants [82]. Although the mouse will probably continue to be the model-of-choice for vertebrates in the meiosis field, the development of state-of-the-art genetics tools such as engineered site-specific nucleases should make us reconsider whether other model organisms may be better suited to ‘reverse genetic’ screening for meiosis genes.

We propose that the zebrafish (Danio rerio) may serve as a powerful genetic screening platform for the meiosis field. A productive research community has grown around the use of zebrafish, but, without efficient targeted mutagenesis techniques, the use of zebrafish has predominantly been limited to studying embryonic development. The barrier to efficiently producing targeted mutations has been overcome by the development of engineered nucleases [84]. While forward genetic screens for adult phenotypes in zebrafish requires more labor and space than most laboratories are able to commit, screening dozens or even hundreds of genes using engineered nucleases should be possible with the resources and capabilities of most small laboratories [84]. This approach may also be taken with the mouse, but the zebrafish offers several advantages. First, adult zebrafish may be raised and maintained at higher densities and for less cost than mice. Second, the large clutch sizes, external development and accessibility of zebrafish embryos make it relatively easy for laboratories to quickly introduce nucleases by microinjection. Finally, homozygous gynogenetic diploid offspring may be generated from F1 fish or even highly mutagenized founders using the well-characterized early-pressure technique [85].

The use of immunofluorescence microscopy to visualize antibodies against components of the synaptic complex and cytological markers of recombination (i.e. Mlh1 and RPA) on whole-mount preparations of zebrafish spermatocytes has been shown to be an important resource for providing physical evidence of where and when recombination is occurring at the chromosomal level [86]. Here, we present our results obtained with polyclonal antibodies that specifically detect Dmc1 and Rad51 on chromosome cores (revealed by Sycep3 immunostaining) of zebrafish spermatocyte nuclear spreads (Fig. 5A). Initial studies on Dmc1 and Rad51 localization at recombination sites, in the context of meiotic progression, revealed that loading of both recombinases onto chromosomes is observed in leptotene and zygotene, and some remnant foci may still be detected at pachytene (Fig. 5A).

Studies using N-ethyl-N-nitrosourea mutagenesis to generate zebrafish knockouts for proteins involved in meiotic prophase I [87], the successful use of reverse genetics [88–91], and outstanding imaging studies (this work and [86,87,92–94]) illustrate the versatility of this model for studying the meiotic processes and defects. The meiotic segregation defects in zebrafish appear to have substantial similarities to those observed in mammals [88,90,95], and, importantly, may model mechanisms underlying human miscarriages. We found that homologs of important human meiosis genes are also present in zebrafish, and the protein sequences are highly conserved (Fig. 5B,C). It is likely that additional vertebrate meiosis genes have yet to be discovered. The zebrafish promises to be a powerful model system for discovery and analysis of new vertebrate meiosis genes.

**Perspective**

Here we have reviewed advances regarding the critical activities in rate-limiting steps of early stages of recombination. We specifically focused in recent biochemical and genetic progress to elucidate the mechanisms of synaptic events in meiotic recombination. This information has revealed important mechanistic information regarding the molecular mode of action of recombinases and functional interaction of recombinases with ancillary proteins assisting in several steps of the DNA strand exchange process. However, these studies have also generated a number of hypotheses regarding the interplay between recombinases during the strand exchange reaction, and raised the question of whether certain ancillary proteins have a specific relationship with one of the two recombinases. Other important specific questions remain. What are the intrinsic structural and biochemical differences explaining the specific functions of Dmc1 and Rad51 in meiotic recombination? What is the 3D structure of DNA–protein complexes involved in strand exchange? What are the mechanisms by which ancillary factors channel recombination intermediates into various HR pathways? Future work is required to answer these questions. The answers may provide the information necessary to connect recombination events and chromosome interactions that ensure correct homologous chromosome distribution to create a balanced number of chromosomes in gametes.
**Author contribution**

CLS and RJP planned experiments, performed experiments, analyzed data, and wrote the paper.

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