

MCM8 Is Required for a Pathway of Meiotic Double-Strand Break Repair Independent of DMC1 in *Arabidopsis thaliana*

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Abstract

Mini-chromosome maintenance (MCM) 2–9 proteins are related helicases. The first six, MCM2–7, are essential for DNA replication in all eukaryotes. In contrast, MCM8 is not always conserved in eukaryotes but is present in *Arabidopsis thaliana*. MCM8 is required for 95% of meiotic crossovers (COs) in *Drosophila* and is essential for meiosis completion in mouse, prompting us to study this gene in *Arabidopsis* meiosis. Three allelic *Atmcm8* mutants showed a limited level of chromosome fragmentation at meiosis. This defect was dependent on programmed meiotic double-strand break (DSB) formation, revealing a role for *AtMCM8* in meiotic DSB repair. In contrast, CO formation was not affected, as shown both genetically and cytologically. The *Atmcm8* DSB repair defect was greatly amplified in the absence of the DMC1 recombinase or in mutants affected in DMC1 dynamics (*sds*, *asy1*). The *Atmcm8* fragmentation defect was also amplified in plants heterozygous for a mutation in either recombinase, *DMC1* or *RAD51*. Finally, in the context of absence of homologous chromosomes (i.e. haploid), mutation of *AtMCM8* also provoked a low level of chromosome fragmentation. This fragmentation was amplified by the absence of *DMC1* showing that both MCM8 and DMC1 can promote repair on the sister chromatid in *Arabidopsis* haploids. Altogether, this establishes a role for *AtMCM8* in meiotic DSB repair, in parallel to DMC1. We propose that MCM8 is involved with *RAD51* in a backup pathway that repairs meiotic DSB without giving CO when the major pathway, which relies on DMC1, fails.

Citation: Crismani W, Portemer V, Froger N, Chelysheva L, Horlow C, et al. (2013) MCM8 Is Required for a Pathway of Meiotic Double-Strand Break Repair Independent of DMC1 in *Arabidopsis thaliana*. PLoS Genet 9(1): e1003165. doi:10.1371/journal.pgen.1003165

Editor: Holger Puchta, Karlsruhe Institute of Technology, Germany

Received: April 13, 2012; **Accepted:** October 28, 2012; **Published:** January 3, 2013

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Funding: RM thanks the EU-FP7 programme (Meiosys-KBBE-2009-222883, <http://www.meiosys.org/>) for financial support. VP thanks Rijk Zwaan for financial support. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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Introduction

Meiosis is a process that occurs in the germlines of sexually reproducing organisms. Two successive rounds of chromosome segregation (meiosis I and II) follow a single round of DNA replication (S phase). The resulting four cells each contain half the genetic content of the pre-meiotic mother cell. The genetic complement of these gametes is a mosaic of the paternal and maternal DNA due to meiotic recombination that occurs between S phase and the first meiotic division [1].

Meiotic recombination begins with programmed DSBs that are dependent on SPO11 and multiple cofactors, including PRD1 in plants [2,3]. DSBs are subsequently resected to yield 3' overhangs that invade the homologous chromosome. At this step, two recombinases co-operate to achieve efficient strand exchange with the homolog, *RAD51* and *DMC1* [4]. *RAD51* is a recombinase involved both at mitosis and meiosis while *DMC1* is specific to meiosis. Importantly, it has been recently shown in *S. cerevisiae* that only the strand exchange activity of *DMC1*, and not of *RAD51*, is required for meiotic crossover formation [5]. *RAD51* appears thus to be an accessory factor of *DMC1* for meiotic homologous crossover formation, but may also serve as a backup to repair breaks

when *DMC1* fails [5]. In *Arabidopsis thaliana*, *RAD51* is indispensable for repair of meiotic DSBs as shown by the extensive meiotic chromosome fragmentation which occurs at meiosis in *Atrad51* mutants [6,7]. *AtDMC1* is required for CO formation but not meiotic DSB repair. Indeed, in *Atdmc1* mutant, meiotic DSBs are repaired in a *AtRAD51*-dependent manner which does not promote chromosome pairing and does not yield COs between homologs, likely using the sister chromatid as a template [7,8]. In addition, consistent with a role of *RAD51* in helping *DMC1* in wild type, the number of *DMC1* foci is severely decreased in a *Atrad51* mutant [7,9], while *RAD51* foci are unaffected in *Atdmc1* [9]. Thus two meiotic functions of *RAD51* emerge, helping *DMC1* to promote COs and promoting DSB repair on the sister without *DMC1*.

Two other *Arabidopsis* mutants, *sds* and *asy1*, have phenotypes reminiscent of *Atdmc1*, repairing breaks using *AtRAD51* but exhibiting major homologous chromosome pairing defects and making no or few COs [10–12]. Both *sds* and *asy1* show localization defects of *AtDMC1* but not of *AtRAD51*, suggesting that they work with *DMC1* to promote interhomolog recombination [12,13]. Based on its amino acid sequence, *SDS* is a cyclin-like protein and *ASY1* is a HORMA domain protein making it the likely functional homologue of *S. cerevisiae* Hop1.

Author Summary

Species that reproduce sexually have two copies of each chromosome, inherited from their father and mother. During a special cell division called meiosis, these two chromosomes are mixed by homologous recombination to give genetically unique chromosomes that will be transmitted to the next generation. This recombination process is initiated by DNA breaks that must be repaired efficiently to maintain fertility. Using the model plant *Arabidopsis thaliana* we revealed here that the gene *AtMCM8* is required to repair a subset of these DNA breaks. However MCM8 appears to not be required for recombination with the homologous chromosome. Instead MCM8 appears to be involved in a safety system that operates to repair DNA breaks that have not been used for homologous recombination. Interestingly the equivalent gene also has an essential meiotic function in the fly and the mouse. However the three species require *MCM8* for different aspects of meiosis.

DSB repair events form intermediates that are resolved as either crossovers (COs) or non-crossovers (NCOs) (gene conversion). COs are required for accurate segregation of chromosomes during meiosis I and can arise from at least two independent pathways known as class I and class II COs. These two pathways coexist in budding yeast, mammals and *Arabidopsis* [1,14–17]. Class I COs are subject to a phenomenon known as interference, whereby the occurrence of a CO significantly reduces the probability of a CO occurring at an adjacent locus, in a distance dependent manner. This pathway is dependent on the ZMM proteins (defined as ZIP1, ZIP2/SHOC1, ZIP3, ZIP4, MSH4, MSH5, MER3) and, in most eukaryotes, is responsible for the majority of COs during meiosis. Class II COs, that do not display interference, require MUS81 [1,14–17].

Here we addressed the meiotic function of MCM8. MCM8 is a member of the eight MCM family proteins (MCM2–9), that all share a well conserved helicase domain. Together MCM2–7, as a hexamer, form a well characterized DNA helicase, which is essential for replication in all eukaryotes [18]. In contrast, MCM8–9 is not present in all eukaryotes [19], being notably missing in *S. cerevisiae*, *S. pombe* and *C. elegans*, but existing in vertebrates and plants. A study in *Xenopus* showed that MCM8 functions during DNA replication at the elongation stage but it is not required for replication licensing. The *Xenopus* MCM8 protein is the only MCM8 representative for which helicase activity has been demonstrated *in vitro* [20]. MCM8 is also involved in, but not essential for the assembly of the pre-replicative complex in human [21]. Very recently, MCM8 and MCM9 has been shown to be involved in homologous recombination-mediated DNA repair in mouse and chicken somatic cells [22,23]. MCM8 has also been shown to be involved in meiosis. In the fruit fly (*Drosophila melanogaster*), in which MCM9 has not been identified, *MCM8* (also known as *REC*) is required for 95% of meiotic COs. In contrast to COs, the frequency of NCOs increases in the absence of *Dmrec* [24]. Finally, a very recent study pointed out a role for MCM8, but not MCM9, in meiotic recombination in mouse [22]. Indeed meiocytes in the mouse *mcm8* mutant accumulate DMCI foci, display synapsis defects and go into apoptosis, consistent with a defect in meiotic DSB repair. The meiotic function of MCM8 has been analyzed only in *Drosophila* and mouse, with contrasting conclusions. This raises the question of the conservation of this function in eukaryotes. The aim of the present study was to further

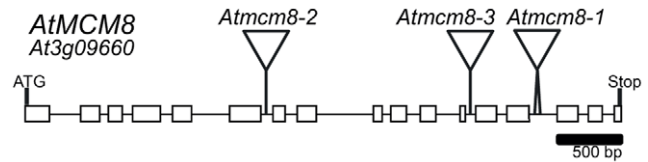


Figure 1. *AtMCM8* gene structure. Exons are represented as black boxes and T-DNA insertions in *Atmcm8-1*, *Atmcm8-2* and *Atmcm8-3* alleles are indicated by triangles.
doi:10.1371/journal.pgen.1003165.g001

explore the meiotic function of MCM8 by deciphering its role in the model plant *Arabidopsis*.

Results

Identification of the *AtMCM8* gene and *Atmcm8* mutations

Phylogenetic analyses of the MCM family [19,24], showed that the *Arabidopsis* genome contains one clear homolog for each MCM2–9, At3g09660 being the MCM8 homolog. We sequenced the At3g09660 CDS using RT-PCR on mRNA from *Arabidopsis* inflorescences. Because of some differences in splicing sites, the At3g09660 CDS slightly differed from the predicted sequence found in the genebank (NM_111800), measured 2,406 bp and contained 17 exons (Figure 1) (genebank BankIt1577803 MCM8 KC109786). We nonetheless confirmed by reciprocal BLAST

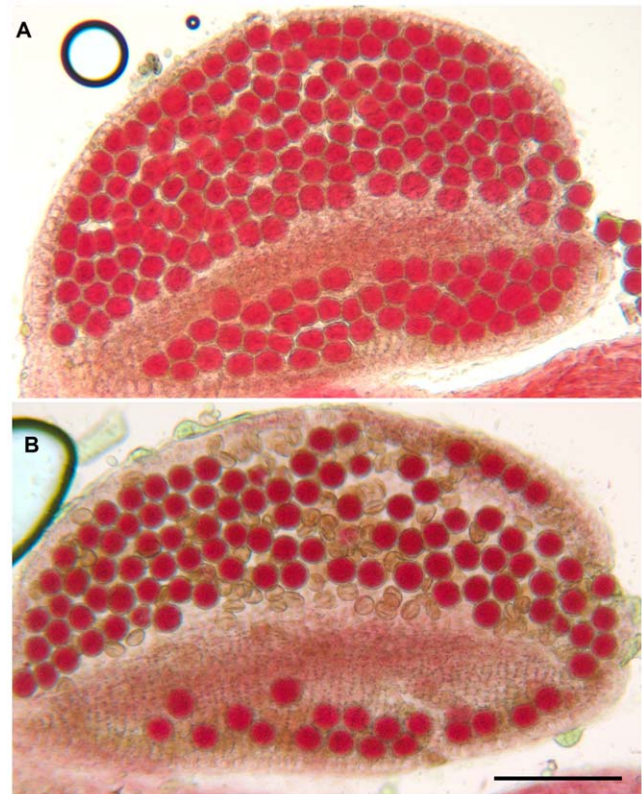


Figure 2. Alexander staining. (A) A wild type anther containing pollen grains that are all viable, as indicated by their red staining and round shape. (B) An *Atmcm8-1* anther containing viable and dead pollen grains as indicated by their abnormal shapes and green coloration. Bar, 100 μ m.
doi:10.1371/journal.pgen.1003165.g002

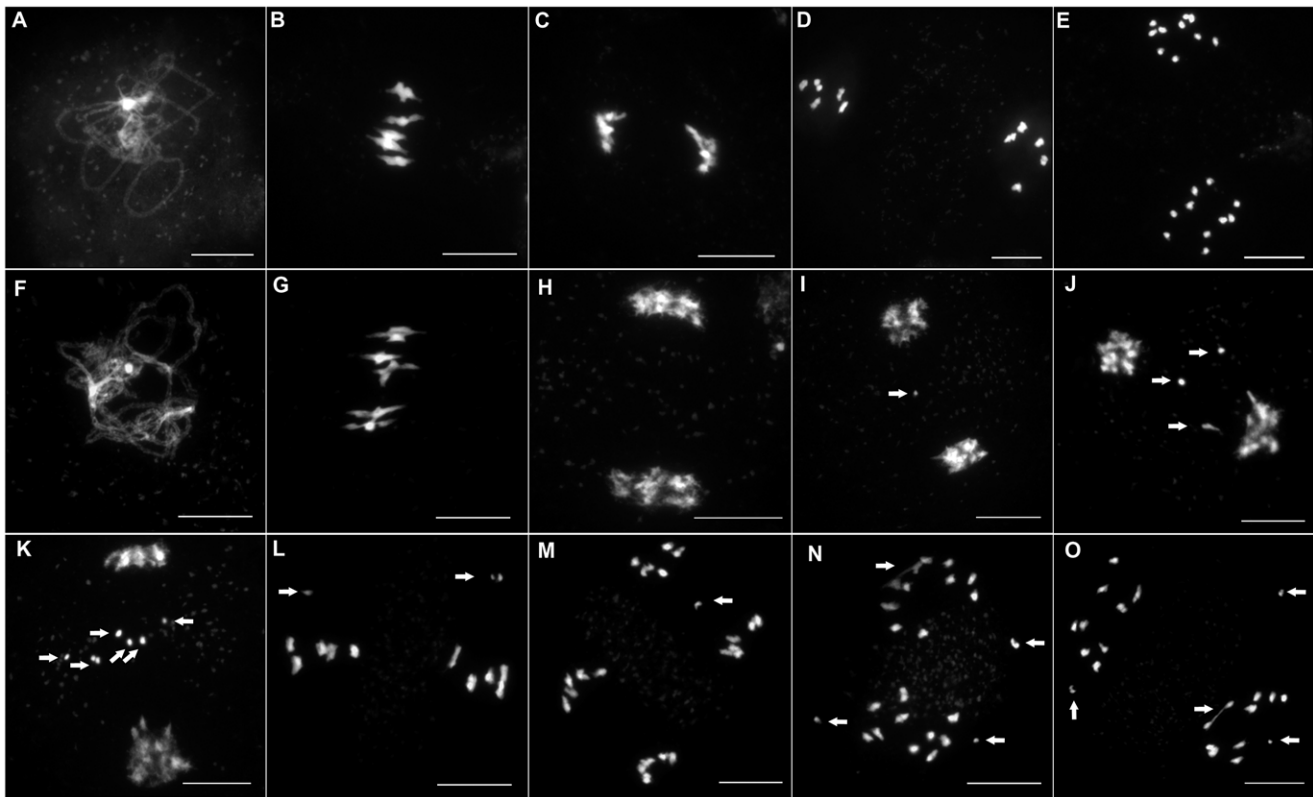


Figure 3. Male meiosis in wild-type and in *Atmcm8*. Male meiosis is shown (A–E) in wild type and (F–O) in *Atmcm8*. Chromosome spreads at (A and F) pachytene, (B and G) metaphase I, (C and H–K) end of anaphase I, (D and L) metaphase II, (E and M–O) anaphase II, using DAPI staining. Fragments and chromosome bridges are indicated with arrows. Bar, 10 μ m.
doi:10.1371/journal.pgen.1003165.g003

analysis and multiple protein alignment that At3g09660 encodes the *Arabidopsis* MCM8 homolog (Figure S1 and [24]).

We identified three T-DNA insertions from the public collections within the *AtMCM8* gene: *Atmcm8-1*, *Atmcm8-2* and

Atmcm8-3 (Figure 1). Plants homozygous for the insertions showed normal vegetative growth but reduced fertility as shown by Alexander staining of pollen (Figure 2). This phenotype (and others described below) was detected only in homozygotes of each mutant. Moreover seed counts showed that *Atmcm8-1* has significantly less seeds than wild type (44.8 ± 5.2 ($n = 41$) compared to 52.4 ± 5.8 ($n = 77$), Z test $p < 10^{-13}$). Allelism tests showed that the meiotic defects observed (see below) were due to the insertions in *Atmcm8*.

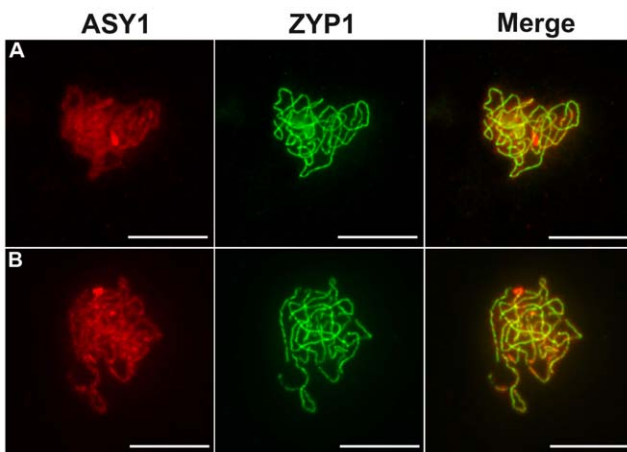


Figure 4. Coimmunolocalization of ASY1 and ZYP1. ASY1 (red), ZYP1 (green) are shown as well as the overlay of both signals (merge) at pachytene in (A) wild type and in (B) *Atmcm8* mutant. In both wild type and mutant the polymerization of the synaptonemal complex, revealed by ZYP1, is completed at pachytene. The ASY1 signal is largely depleted from the chromosomes as the synaptonemal complex forms. Bar, 10 μ m.
doi:10.1371/journal.pgen.1003165.g004

Atmcm8 chromosomes fragment during meiosis

To investigate if this reduction in fertility was linked to a meiotic defect, we analyzed meiotic progression by DAPI (4',6-diamidino-2-phenylindole) staining of meiotic chromosome spreads in all three mutant alleles. In wild type meiosis (Figure 3A–3E), chromosomes condense at leptotene. Then, synapsis is initiated at zygotene until its completion in pachytene when the two homologous chromosomes are connected along their entire length by a proteinous structure called the synaptonemal complex [25] (Figure 3A and Figure 4A). Desynapsis occurs at diplotene and further condensation of the chromosomes occurs. Five bivalents continue to condense and become visible at diakinesis. At metaphase I, the five bivalents align on the metaphase I plate (Figure 3B). At anaphase I homologous chromosomes segregate to opposite poles (Figure 3C). At telophase I the two groups of five recombinant chromosomes begin to decondense. At prometaphase II chromosomes recondense and align on the two metaphase II plates (Figure 3D). At anaphase II each of the ten chromosomes segregate their two sister chromatids to opposite poles resulting in four balanced groups of five chromatids (Figure 3E).

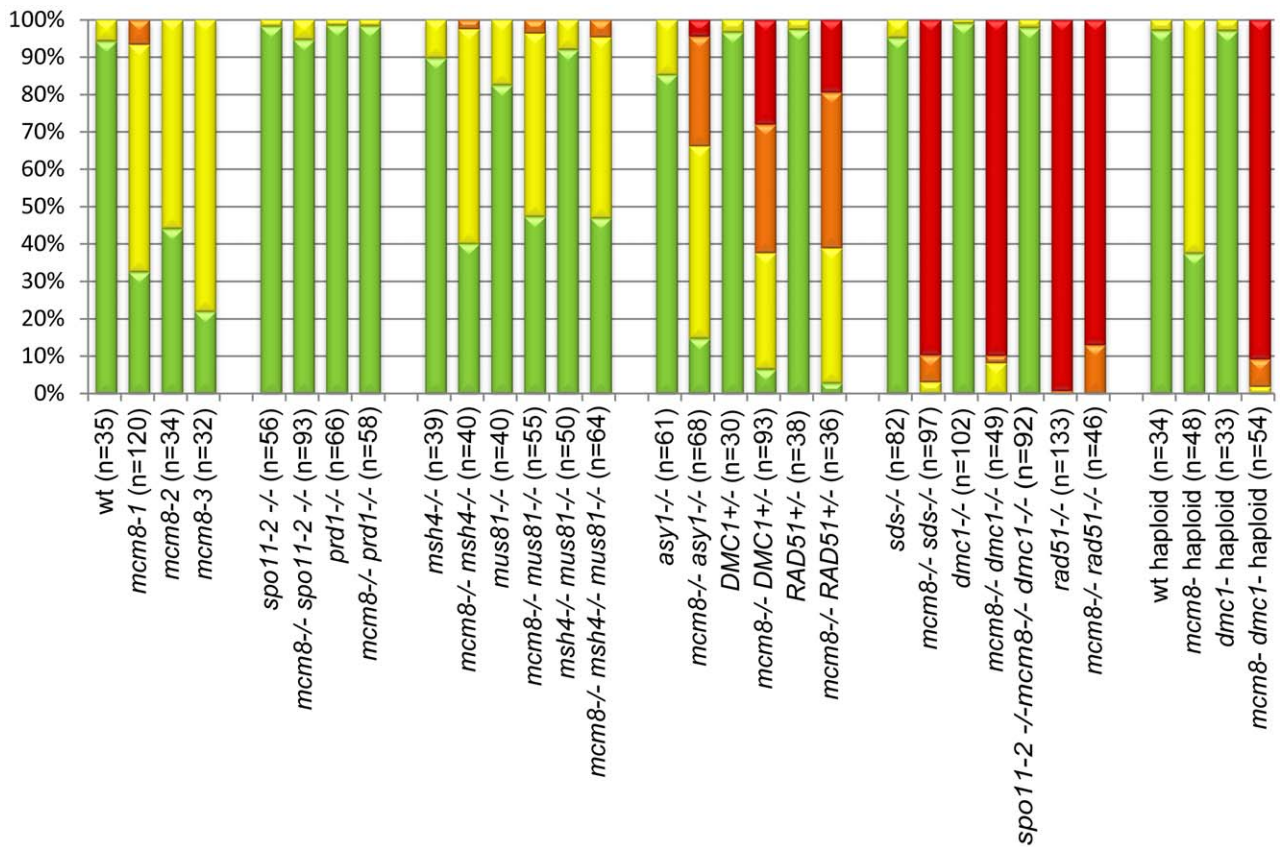


Figure 5. Quantification of chromosome fragmentation levels. For each genotype, a number (indicated in brackets) of late anaphase I/ telophase I cells were observed after chromosome spreads and DAPI staining. DAPI-stained bodies observed above the expected number of chromosomes were counted as fragments, and cells were classified as having 0 (green), 1–5 (yellow), 6–10 (orange), or more than 10 fragments (red). The percentage of each class is shown. doi:10.1371/journal.pgen.1003165.g005

In all three *Atmcm8* alleles, meiosis appeared to progress normally from leptotene through to pachytene (Figure 3F) where chromosomes condensed, aligned and fully synapsed like wild type. The completion of synapsis in *Atmcm8* was confirmed by immunolabeling meiotic chromosomes with antibodies against ASY1 and *AtZYP1* (Figure 4B), that are components of the axial elements and of the transverse filament of the synaptonemal complex, respectively [26,27]. Chromosomes desynapsed normally during diplotene and we observed five bivalents as condensation progressed during diakinesis, revealing the presence of chiasmata (the cytological manifestation of CO). At metaphase I, five bivalents were systematically observed in all mutant alleles, showing that at least one CO is formed per pair of homologous chromosomes (Figure 3G). Anaphase I proceeded, however chromosome fragmentation was observed in all three *Atmcm8* alleles (Figure 3H–3K), with 1 to 10 chromosome fragments detected in 60 to 80% of the cells (Figure 5). Chromosomes aligned on the metaphase II plate, with fragments dispersed throughout the cell (Figure 3L). Anaphase II proceeded but additional chromosome fragments appeared (Figure 3M–3O). This fragmentation persists at telophase II. We also observed fragmentation in female meiosis showing that *Atmcm8* mutation also affects female meiosis (data not shown).

Chromosome fragmentation in *Atmcm8* is dependent on meiotic DSB formation

In *Atspo11-2* and *Atprd1*, no meiotic DSBs are formed and therefore recombination does not occur [3,28]. Thus at metaphase

I, ten univalents are observed and segregate randomly (Figure 6A–6B and 6E–6F). To test whether the chromosome fragmentation seen in *Atmcm8* mutants are dependent on DSB formation or not, we introduced the *Atspo11-2* and *Atprd1* mutations independently into *Atmcm8*. At meiosis, we observed ten univalents at metaphase I in the *Atmcm8/Atspo11-2* or *Atmcm8/Atprd1* and, importantly, the chromosome fragmentation was abolished (Figure 6C–6D and 6G–6H, Figure 5). Therefore, the fragmentation defect of *Atmcm8* is dependent on *AtSPO11-2* and *AtPRD1*. Thus, *AtMCM8* is required for efficient repair of the DSBs that initiate meiotic recombination.

Atmcm8 does not affect CO frequency

We then tested if the *Atmcm8* fragmentation phenotype is dependent on the presence of any of the known pathways of CO formation, using epistasis tests. We used *Atmsh4* and *Atzip4* that are both required for class I CO formation and *Atmus81* that is required for class II CO formation. In the *Atmcm8/Atmsh4*, *Atmcm8/Atzip4*, *Atmcm8/Atmus81* double mutants and the *Atmcm8/Atmsh4/Atmus81* triple mutant, we still observed a chromosome fragmentation defect as in the *Atmcm8* single mutant (Figure 5 and Figure 7, data not shown for *Atmcm8/Atzip4*). Thus the *Atmcm8* fragmentation phenotype is independent of MSH4, ZIP4 and MUS81.

In *Atmcm8* and *Atmcm8/Atmus81* we invariably observed five bivalents at metaphase I, suggesting that the formation of class I COs, which account for most of the CO in wild type, is not grossly

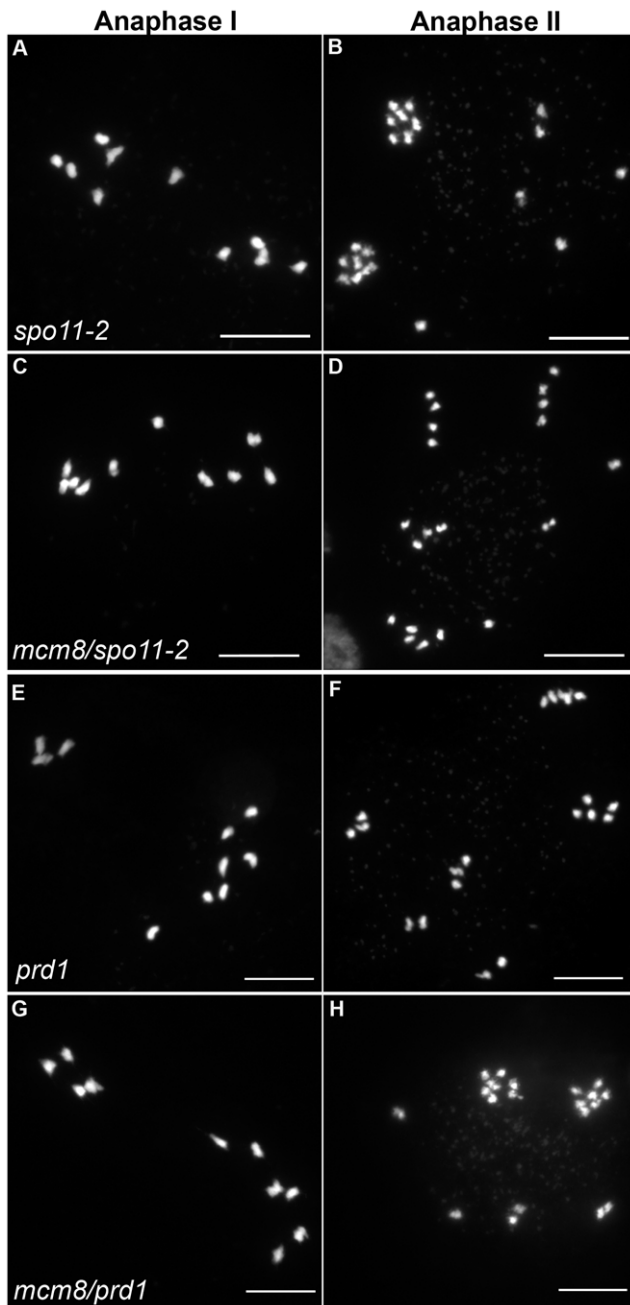


Figure 6. Epistasis tests between *Atmcm8* and two mutants affected in DSB formation. Meiotic spreads with (A–B) *Atspo11-2*, (C–D) *Atmcm8/Atspo11-2*, (E–F) *prd1*, (G–H) *Atmcm8/Atspo11-2* using DAPI staining at anaphase I and anaphase II. Bar, 10 μ m. doi:10.1371/journal.pgen.1003165.g006

affected by the *Atmcm8* mutation. This was further supported by counts of *AtMLH1* foci, a marker of class I COs at late prophase of meiosis I [29,30] (Figure S2), that revealed no significant differences between wild type (10.1 ± 1.4 per cell; $n = 81$) and the *Atmcm8* mutant (10.3 ± 1.9 ; $n = 86$ ($Z p = 0.55$)). In *Atmcm8/Atmsh4* (Figure 6), the residual number of bivalents at metaphase I was unchanged compared to the single *Atmsh4* mutant (1.5 ± 1 ; $n = 91$ vs 1.3 ± 1.1 ; $n = 91$ ($Z p = 0.94$)), strongly suggesting that class II CO formation is not affected neither by *Atmcm8* mutation. We then measured recombination frequency and crossover interference genetically in *Atmcm8*. This was achieved using tetrad analysis

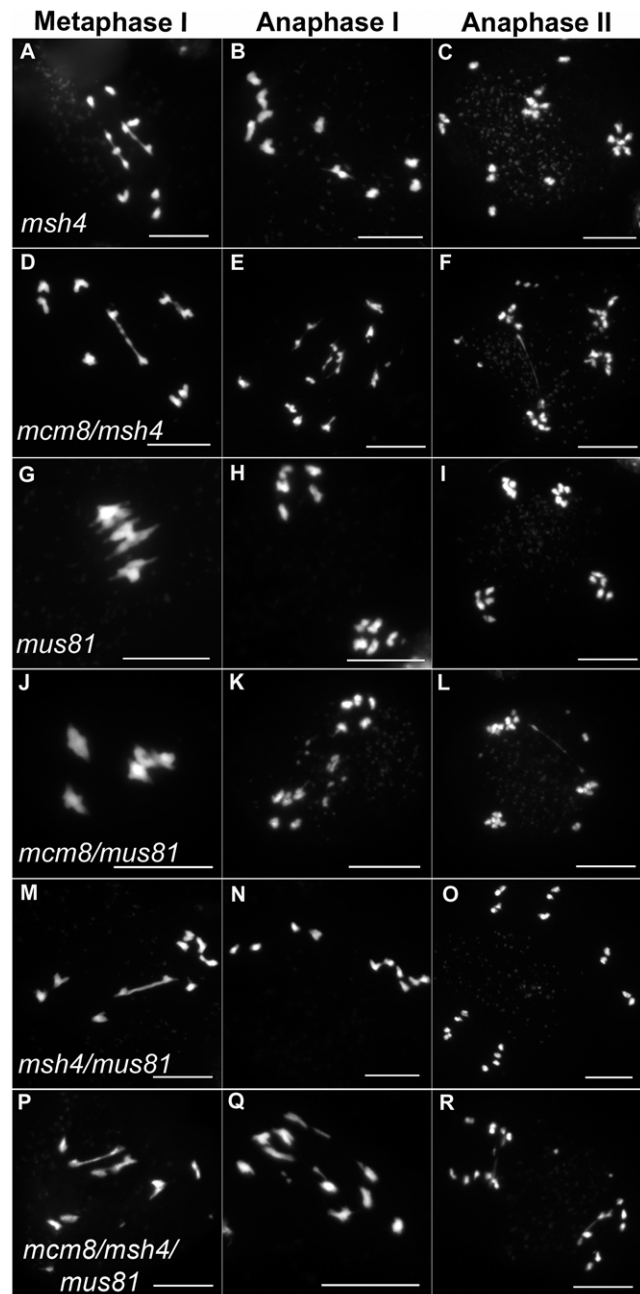


Figure 7. Epistasis tests between *Atmcm8* and mutants affected in crossover formation. Meiotic spreads with (A–C) *Atmsh4*, (D–F) *Atmcm8/Atmsh4*, (G–I) *Atmus81*, (J–L) *Atmcm8/Atmus81*, (M–O) *Atmsh4/Atmus81*, (P–R) *Atmcm8/Atmsh4/Atmus81* using DAPI staining at metaphase I, anaphase I and anaphase II. Bar, 10 μ m. doi:10.1371/journal.pgen.1003165.g007

(Fluorescent-Tagged Lines, FTL) which is a visual pollen assay allowing the measurement of multiple COs simultaneously with access to all four chromatids from the same meiosis [31]. Two different sets of adjacent intervals on chromosome 5 have been analyzed, (I5aI5b and I5cI5d), representing four intervals in total. We did not detect any difference in recombination frequency between the *Atmcm8* and wild type for any of these intervals (Table 1, Genetic Distance), consistent with the cytological data. Also, interference, that affects the distribution of crossovers, was unchanged compared to wild type for both sets of adjacent

Table 1. Genetic distances and interference in *Atmcm8* using FTLs.

	Genetic distance (cM)			Interference ratio**		
	Wild type	<i>Atmcm8</i>	<i>P</i> value*	Wild type	<i>Atmcm8</i>	<i>P</i> value*
I5a	24.2±0.8	22.3±1	0.12	0.27 (chi ² p<10 ⁻³⁰)	0.34 (chi ² p<10 ⁻³⁰)	0.21
I5b	14.4±0.6	16.1±0.9	0.15			
I5c	5.9±0.4	7.5±0.8	0.09	0.43 (chi ² p<10 ⁻⁵)	0.30 (chi ² p<10 ⁻⁴)	0.50
I5d	5.7±0.4	6.3±0.7	0.44			

Values are means ± Standard Error. Number of tetrads: Wild type I5aI5b n = 1986, *Atmcm8* I5aI5b n = 1022, wild type I5cI5d n = 1860, *Atmcm8* I5cI5d n = 646.

*Z-test between wild type and *Atmcm8*.

**The interference ratio is defined as the ratio of genetic distance of I5a with a CO in I5b by the genetic distance of I5a without a CO in I5b. The same was done for the interference ratio between I5c and I5d. Absence of interference would give a ratio of 1 that would tend to 0 with increased interference [57]. The chi square test shows a deviation from 1, and thus the presence of interference [31].

doi:10.1371/journal.pgen.1003165.t001

intervals (Table 1, Interference Ratio). Taken together these data suggest that *AtMCM8* is not involved in CO formation. This contrasts from the observation that the absence of MCM8 reduces COs frequency by 95% in *Drosophila* [24].

Mei9/Rad1 is another gene required for the formation of more than 90% of the COs in *Drosophila* [32]. Given the major difference in MCM8 function between *Arabidopsis* and *Drosophila*, we tested the role of *AtRAD1* [33–35] in crossover formation in *Arabidopsis*. Cytological analysis showed that the single *Atrad1* mutant has no obvious defect in CO formation. We then analyzed if *AtRAD1* has a minor effect. To achieve this, we constructed a *shoc1/Atrad1* double mutant and a *Atmus81/shoc1/Atrad1* triple mutant to be able to detect a weak reduction in CO formation, in a sensitive context where there are no class I and class II COs. However, this triple mutant was not different from *Atmus81/shoc1* (0.99±0.84 (n = 74) versus 1.15±1.28 (n = 75), χ^2 p = 0.36) and neither was *shoc1/Atrad1* different from *shoc1* (1.47±1.07 (n = 51) versus 1.56±0.86 (n = 32), χ^2 p = 0.67). These genes, MCM8 and MEI9/RAD1, are essential for CO formation in *Drosophila* but not in *Arabidopsis* showing divergent functions. However, contrary to RAD1, MCM8 has conserved a meiotic function in *Arabidopsis*.

The *Atmcm8* DSB repair defect is amplified by *DMC1* mutation

DMC1 is involved at the strand invasion stage of meiotic recombination and *Atdmc1* mutants fail to synapse and to make COs (Figure 8A–8B, 8G–8H). However, DSBs are repaired in *Atdmc1*, in an *AtRAD51*-dependent manner, without CO formation, suggesting that the DSBs are repaired on sister chromatids in these mutants [8,12]. In the *Atmcm8/Atdmc1* double mutant, from metaphase I to the end of the meiosis we observed extensive chromosome fragmentation in all cells, which was much more intense than in the single *Atmcm8* mutant (compare Figure 8C–8D to Figure 3I–3K and see quantification in Figure 5). Consistently, the *Atmcm8/Atdmc1* double mutant was completely sterile whereas *Atmcm8* has moderate fertility reduction and *Atdmc1* produce some residual seeds [8,12] (Table 2). Mutating *SPO11-2* in this *Atmcm8/Atdmc1* double mutant abolished the chromosome fragmentation (Figure 8E–8F, Figure 5), demonstrating that MCM8 and *DMC1* act in parallel pathways of meiotic DSB repair.

Furthermore in the *Atmcm8* mutant context, we observed a more drastic meiotic chromosome fragmentation in plants heterozygous for *DMC1* (*Atmcm8*^{-/-}*AtDMC1*^{+/-}) than wild type for *DMC1* (*Atmcm8*^{-/-}*AtDMC1*^{+/+}) (compare Figure 8I–8J to Figure 3I–3K,

quantification on Figure 5), accompanied by a strong reduction of fertility (Table 2). However, the fragmentation observed in *Atmcm8*^{-/-}*AtDMC1*^{+/-} was less dramatic than in the double mutant (*Atmcm8*^{-/-}*Atdmc1*^{-/-}) (Figure 5), which is also supported by the fertility levels (Table 2). This is despite the *AtDMC1* mutation being recessive (in an *AtMCM8*^{+/+} or *AtMCM8*^{+/-} context). Thus, in the absence of *Atmcm8*, the mutation of one of the two copies of *DMC1* was enough to enhance fragmentation, which is even more drastic when both *DMC1* alleles are disrupted.

The *Atmcm8* DSB repair defect is amplified by mutation of *ASY1*, *SDS*, or one copy of *RAD51*

Therefore we tested the relationship of *AtMCM8* with *ASY1* and *SDS*, two proteins that are required for normal *DMC1* localization [3,13]. In the *sds* and *asy1* single mutants, COs are greatly reduced (Figure 9E–9F) [10,11]. In the *Atmcm8/asy1* and *Atmcm8/sds* double mutant, we observed chromosome fragmentation from anaphase I onwards, which was much greater than that seen in the *Atmcm8*^{-/-} single mutant (compare Figure 9G–9H with Figure 3I–3K, quantification on Figure 5). Thus, mutation of *SDS* or *ASY1* amplified the fragmentation phenotype of *Atmcm8*. Finally, both the single *Atrad51* mutant and the double *Atmcm8/Atrad51* mutant show intense chromosome fragmentation (Figure 10). Interestingly, while *AtRAD51*^{+/-} does not show chromosome fragmentation, *Atmcm8*^{-/-}/*AtRAD51*^{+/-} showed more chromosome fragmentation than *Atmcm8* (Figure 10, Figure 5). Thus, in the absence of *Atmcm8*, the mutation of one of the two copies of *RAD51* was enough to enhance fragmentation.

AtDMC1 foci number is unaffected in *Atmcm8*

Given the relationship between *DMC1* functional gene copy number and the degree of *Atmcm8*-dependent fragmentation, we looked at *DMC1* behavior in *Atmcm8*. No significant difference in *DMC1* foci shape or number was observed in *Atmcm8*^{-/-} compared to wild type (Table 2). Similarly, we did not detect any differences in number or shape of *DMC1* foci in *Atmcm8*^{-/-}*AtDMC1*^{+/-} or *Atmcm8*^{-/-}*AtRAD51*^{+/-} compared to either wild type or *Atmcm8*^{-/-} (Figure S3, Table 2). In the *Atmcm8/Atrad51* double mutant, we observed a marked decrease of *DMC1* foci number, which was however similar to what was previously observed in a single *Atrad51* mutant [7] (Table 2). It is intriguing that *Atmcm8*^{-/-}*AtDMC1*^{+/-} and *Atmcm8*^{-/-}*AtRAD51*^{+/-} exhibit a more drastic meiotic defect than *Atmcm8*^{-/-}*AtDMC1*^{+/+}, while *DMC1* foci number and shape appear similar. However, it is

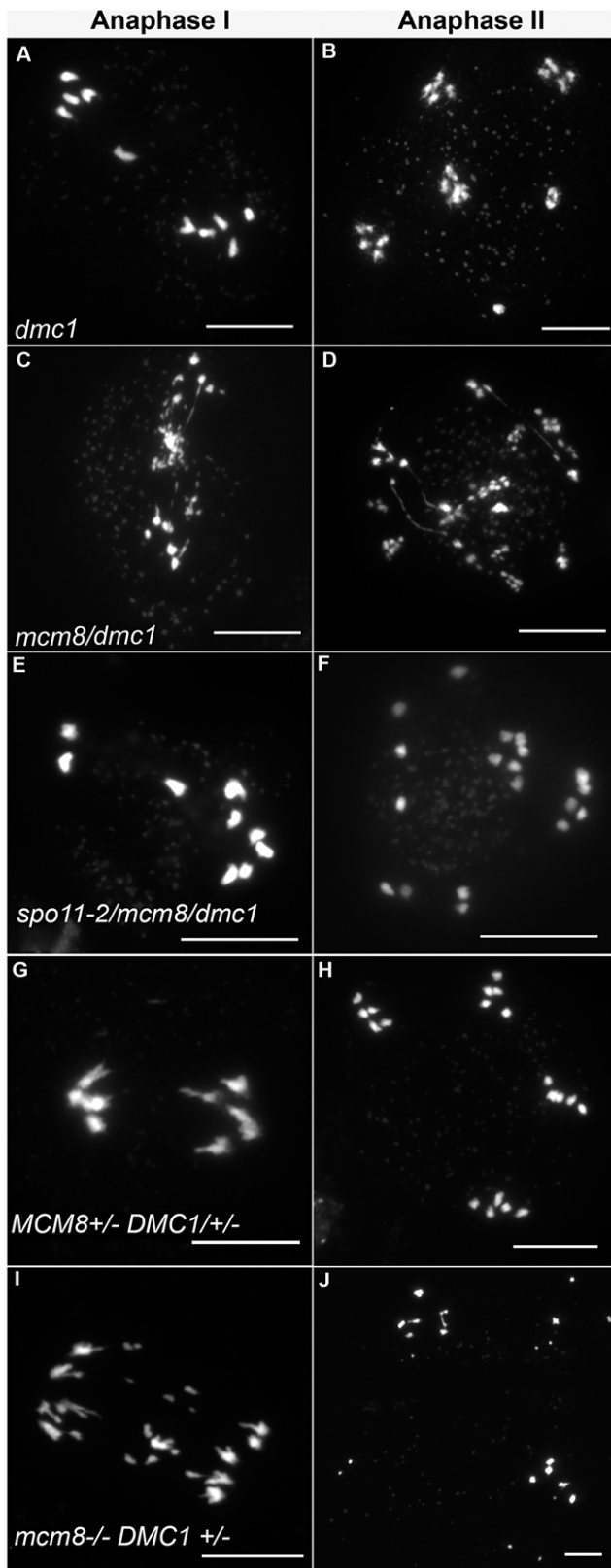


Figure 8. Epistasis tests between *Atmcm8* and *Atdmc1*. Meiotic spreads with (A–B) *Atdmc1*^{-/-}, (C–D) *Atmcm8*^{-/-}/*Atdmc1*^{-/-}, (E–F) *Atmcm8*^{-/-}/*Atdmc1*^{-/-}/*Atspo11*^{-/-}, (G–H) *Atmcm8*^{+/-}/*AtDMC1*^{+/-}, (I–J) *Atmcm8*^{-/-}/*AtDMC1*^{+/-}, using DAPI staining at anaphase I and anaphase II. Bar, 10 μm.

doi:10.1371/journal.pgen.1003165.g008

possible that immunolocalization fails to detect subtle differences in DMC1 protein quantity or dynamics.

In the absence of homologous chromosomes, DSBs fail to be repaired in the absence of both MCM8 and DMC1

Next we explored the functional relationship between MCM8 and DMC1, in haploid plants, where homologous chromosomes are not present. Thus, the only template available for meiotic DSB repair is the sister chromatid. Meiotic chromosome spreads, in a wild-type haploid, showed that the five chromosomes were intact and segregated randomly at anaphase I [36] (Figure 11A–11B), suggesting that DSBs are efficiently repaired. The haploid *Atmcm8* mutant had a limited fragmentation defect (Figure 11C–11D), similar to the defect in the diploid *Atmcm8* mutant (Figure 5 for quantification). The *Atdmc1* haploid had no fragmentation (Figure 11C–11F). In clear contrast, in the double *Atmcm8/Atdmc1* haploid, we observed extensive meiotic chromosome fragmentation (Figure 11G–11H, see Figure 5 for quantification). This shows that in a haploid context, DSB repair is efficient in wild type and *Atdmc1*, only slightly affected in *Atmcm8*, but ineffective in the *Atmcm8/Atdmc1* double mutant. This suggests that in the absence of a homologous template, *AtMCM8* and *AtDMC1* catalyze DSB repair on the sister chromatid in a redundant manner.

Discussion

Here *AtMCM8* was shown to be involved in meiotic DSB repair but not CO formation. This study thus revealed a pathway for DNA DSB repair that does not yield COs. This pathway depends on *AtMCM8* and acts in parallel to the *AtDMC1* pathway from which COs originate.

AtMCM8 is required for efficient meiotic DSB repair but not for CO formation

Arabidopsis MCM8 is required for effective meiotic DSB repair as all *Atmcm8* mutant alleles had a clear, albeit limited, chromosome fragmentation defect at meiosis. The fragmentation is dependent on meiotic DSB formation as it disappears when *AtSPO11-2* or *AtPRD1* is absent. However, in contrast to *Drosophila rec* (*mcm8*) mutants, genetic and cytological data strongly support that CO formation is not affected by *AtMCM8* mutation: (1) In the absence of *AtMSH4* or *AtZIP4* (class I COs) or *AtMUS81* (class II COs) fragmentation still occurred and the number of bivalents was unchanged. (2) MLH1 foci numbers, a marker of class I COs, were unchanged in *Atmcm8*. (3) The genetic analysis using FTLs revealed no difference in terms of genetic distance and the strength of interference. These data showed that *AtMCM8* acts in a pathway which repairs a subset of meiotic DSB and does not lead to CO formation.

Two pathways for DSB repair: one dependent on *AtMCM8* and one on *AtDMC1*

A striking finding was that *AtMCM8* becomes crucial when the DMC1 pathway was affected. Indeed, we observed a drastic amplification of the *Atmcm8* mutant chromosome fragmentation defect when one of the two allelic copies of DMC1 was mutated, which was even more drastic when both DMC1 copies were mutated. This extensive fragmentation defect reflects a failure of DSB repair, as it is abolished by *SPO11-2* mutation. Further, this extensive fragmentation was consistently confirmed in the absence of *AtMCM8* and *SDS*, or *AtMCM8* and *ASY1*. *SDS* and *ASY1* are essential for *AtDMC1* loading/stability [12,37]. Extensive fragmentation was also observed when one copy of *RAD51* was

Table 2. Seed per fruit and fragmentation levels in different combinations of double mutants.

Genotype	Seeds per fruit	Bivalent	Fragmentation	Number of DMC1 foci
wt	46.5±1.8 (46) ^a	yes	none	221.6±10.8 (25)
MCM8 ^{+/+} DMC1 ^{+/-}	47.8±2.2 (51) ^a	yes	none	187.3±10.6 (20)
MCM8 ^{+/-} DMC1 ^{+/-}	49.0±1.8 (44) ^a	yes	none	ND
MCM8 ^{+/-} DMC1 ^{+/+}	50.1±1.4 (48) ^a	yes	none	ND
mcm8 ^{-/-} DMC1 ^{+/+}	30.5±2.2 (69) ^b	yes	+	227.8±10.9 (50)
mcm8 ^{-/-} DMC1 ^{+/-}	7.0±1.2 (85) ^c	yes	++	212.9±4.8 (87)
MCM8 ^{+/+} dmc1 ^{-/-}	1.9±0.2 (76) ^{de}	no	none	ND
MCM8 ^{+/-} dmc1 ^{-/-}	1.3±0.2 (65) ^{de}	no	none	ND
mcm8 ^{-/-} dmc1 ^{-/-}	0.0±0.0 (50) ^e	no	+++	ND
mcm8 ^{-/-} sds ^{-/-}	0.0±0.0 (50) ^e	no	+++	ND
mcm8 ^{-/-} asy1 ^{-/-}	ND	few	++	ND
Genotype	Seeds per fruit	Bivalent	Fragmentation	Number of DMC1 foci
wt	58.4±1.1 (48) ^a	yes	none	221.6±10.8 (25)
MCM8 ^{+/+} RAD51 ^{+/-}	56.8±1.6 (52) ^a	yes	none	202.7±6.3 (43)
MCM8 ^{+/-} RAD51 ^{+/-}	57.4±1.3 (50) ^a	yes	none	ND
MCM8 ^{+/-} RAD51 ^{+/+}	55.8±1.8 (49) ^a	yes	none	ND
mcm8 ^{-/-} RAD51 ^{+/+}	35.4±3.0 (49) ^b	yes	+	ND
mcm8 ^{-/-} RAD51 ^{+/-}	10.3±1.6 (87) ^c	yes	++	182.5±8.0 (29)
MCM8 ^{+/+} rad51 ^{-/-}	0.0±0.0 (50) ^d	no	+++	50±11 (21) [7]
MCM8 ^{+/-} rad51 ^{-/-}	0.0±0.0 (50) ^d	no	+++	ND
mcm8 ^{-/-} rad51 ^{-/-}	0.0±0.0 (50) ^d	no	+++	51.2±7.0 (8)

Values are means ± Standard Error. The number of fruit or cells counted is indicated in brackets. ND: not determined, a–e: indicates significant differences among groups (Newman Keuls test, $p > 0.05$). Number of crosses indicates fragmentation levels, based on Figure 5.
doi:10.1371/journal.pgen.1003165.t002

mutated in the *Atmcm8* mutant. A function of RAD51 as a cofactor of DMC1 has been recently identified in yeast [5], and consistently DMC1 foci number is drastically reduced in the *Arabidopsis rad51* mutant [7,9]. We thus propose that two pathways of DSB repair coexist, one dependent on *AtMCM8* and the other one on *AtDMC1*. In the absence of *AtDMC1*, efficient DSB repair occurs without CO formation. This repair depends on *AtRAD51* [7,8,12] and on *AtMCM8* (this study). Such RAD51-mediated, DMC1-independent, repair also exists in *S. cerevisiae* but is normally inhibited by RAD51 regulators [38–42]. Consequently, we suggest that, in the *Atdmc1* context, *AtMCM8* and *AtRAD51* can cooperate to repair DSBs using the sister as a template. In addition to this function, *AtRAD51* is required for the *AtDMC1*-dependent pathway (possibly as an accessory factor for the DMC1 strand-exchange activity as shown in yeast [5]) as repair is completely defective in the single *Atrad51* mutant [6], like in the double *Atmcm8/Atdmc1* mutant.

The fact that the fragmentation defect is limited in the single *Atmcm8* mutant, suggests that the *AtMCM8/AtRAD51* pathway would be essential for a limited number of events in wild type, when DMC1 fails. The repair events promoted by *AtMCM8* are likely not intended to become a CO, as CO formation was not affected in *Atmcm8*, leaving sister chromatid repair or NCOs as the only other known possibilities. The absence of synapsis in *Atdmc1* [7,8], in which the *AtMCM8/AtRAD51* pathway must be active, favors the hypothesis of sister chromatid repair. In contrast, the DMC1 pathway promotes CO formation. However, DMC1 foci in wild type, outnumber COs by approximately 25 to 1 [7,43]. This suggests that repair of many DSBs catalyzed by DMC1 do not become CO, but NCO (that involve the homologous

chromosome) or sister chromatid exchange (SCE). In *Arabidopsis*, the genome-wide frequency of NCOs and SCEs is currently unknown. We favor the hypothesis that DMC1 promotes NCOs, as DMC1 promotes synapsis. However, it should be noted that DMC1 is also able to promote SCE, notably in the haploid *mcm8* context. Indeed, only the simultaneous mutation of *AtDMC1* and *AtMCM8* in haploids led to extensive chromosome fragmentation. The capacity of DMC1 to promote inter-sister repair was previously shown in other mutant background in both *Arabidopsis* [9] and yeast [44].

In summary we suggest that two pathways of DSB repair exist in wild type meiosis: The first pathway relies on the strand exchange activity of DMC1, and is also promoted by ASY1, SDS and RAD51 as a co-factor of DMC1 [5]. This pathway generates the COs, but also NCOs and SCEs in a ratio that remains to be determined. The second pathway of the model, which may be viewed as a backup pathway in case of failure of DMC1, relies on the strand exchange activity of RAD51 and the helicase activity of MCM8, and uses the sister chromatid as a template.

MCM8 function varies among eukaryotes

The function of MCM8 appears to differ markedly in *Arabidopsis* and in *Drosophila*. Interestingly, DMC1 and MCM8 appear to be partially redundant in *Arabidopsis* while the *Drosophila* genome seems devoid of a DMC1 homolog [45]. Thus CO formation in *Drosophila* appears to rely on a RAD51/MCM8 pathway, which has only a minor role in wild type meiotic DSB repair in *Arabidopsis*. The CO pathways appear to differ considerably in the two species, mainly using ZMMs in *Arabidopsis* but not RAD1, and

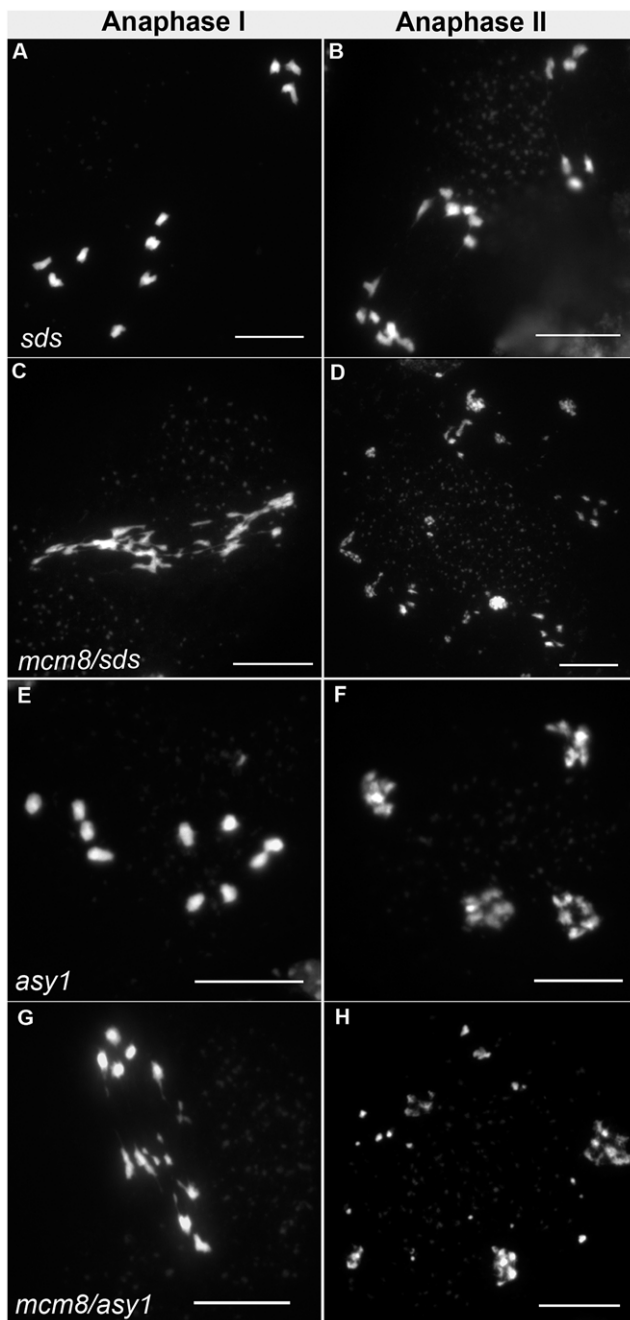


Figure 9. Epistasis tests between *Atmcm8* and *sds* or *asy1*. Meiotic spreads with (A–B) *sds*^{-/-}, (C–D) *Atmcm8*^{-/-}/*sds*^{-/-}, (E–F) *asy1*^{-/-}, (G–H) *Atmcm8*^{-/-}/*asy1*^{-/-}, using DAPI staining at anaphase I and anaphase II. Bar, 10 μ m.
doi:10.1371/journal.pgen.1003165.g009

the reverse in *Drosophila*, i.e. RAD1 but not ZMMs (that are absent from the *Drosophila* genome). *Drosophila* appears to be unique, as in distant species like *S. cerevisiae*, mammals and *C. elegans* CO formation depends mainly on ZMM. Adding to the complexity, MCM8 exists in mammals but not in *S. cerevisiae* and *C. elegans* [19,24]. In mouse, *MCM8* mutation leads to a meiotic arrest, with defects in homologous synapsis and over-accumulation of DMC1 foci before apoptosis, suggestive of defects in DSB repair [22]. We would like to suggest that these defects may be compatible with

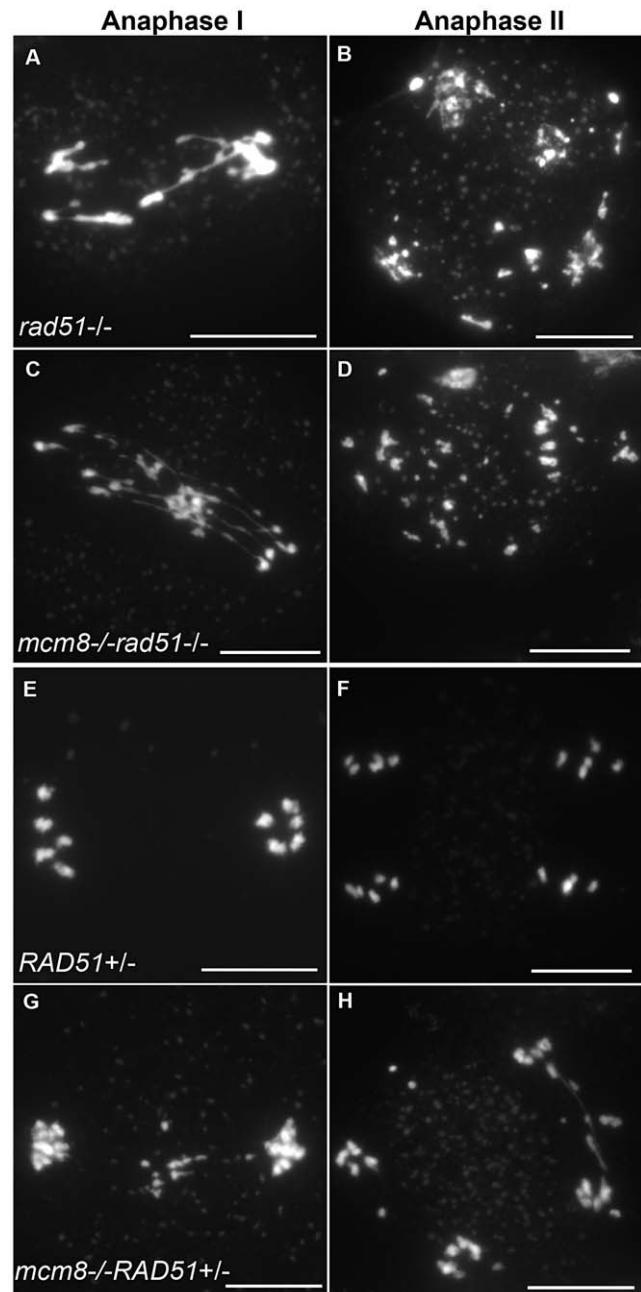


Figure 10. Epistasis tests between *Atmcm8* and *Atrad51*. Meiotic spreads with (A–B) *Atrad51*^{-/-}, (C–D) *Atmcm8*^{-/-}/*Atrad51*^{-/-}, (E–F) *Atrad51*^{+/-}, (G–H) *Atmcm8*^{-/-}/*Atrad51*^{+/-}, using DAPI staining at anaphase I and anaphase II. Bar, 10 μ m.
doi:10.1371/journal.pgen.1003165.g010

MCM8 being required for a backup pathway in the case of failure of DMC1 to repair breaks, like in *Arabidopsis*. The lack of the backup pathway may lead to the accumulation of DMC1 foci, and a failure to repair a subset of breaks, triggering apoptosis (it is noteworthy that DSB repair defects do not trigger meiotic arrest or apoptosis in *Arabidopsis*). This illustrates the variety of mechanisms that arose in the course of evolution to fulfill the conserved outcome of meiotic DSB repair and CO formation.

In conclusion, our data reveals the meiotic function of MCM8 in *Arabidopsis*. Cytological and genetic analyses showed that *AtMCM8* is involved in DSB repair but it is not a determinant

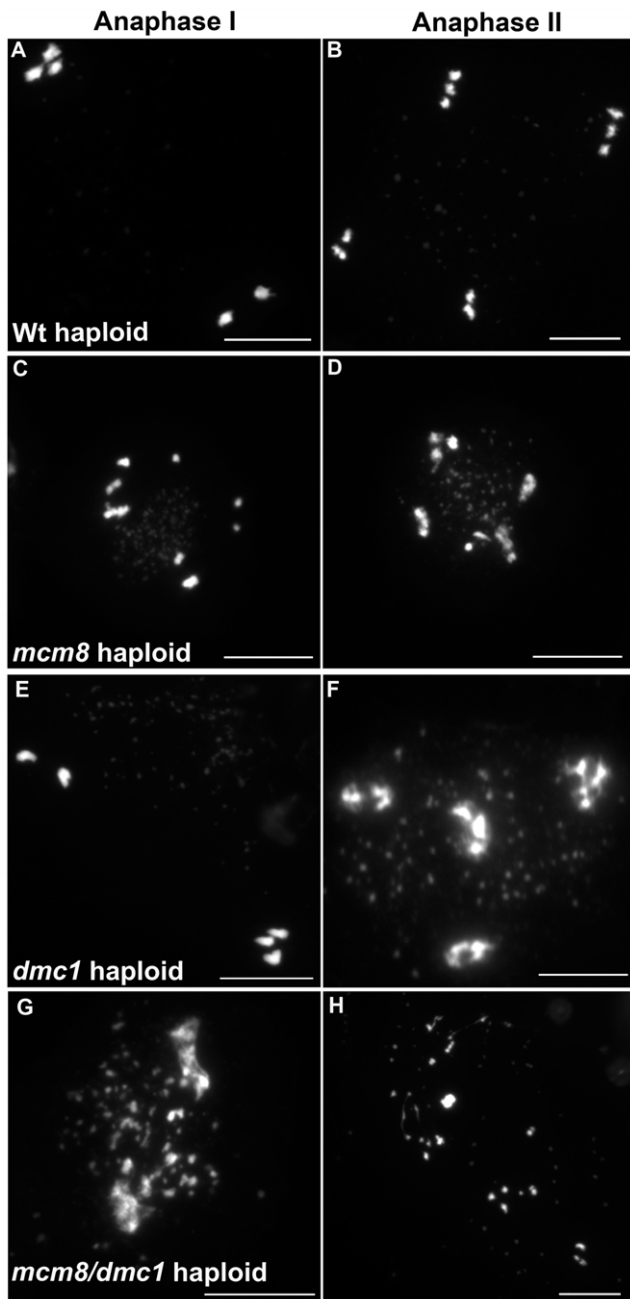


Figure 11. *Atmcm8* haploids during anaphase I and anaphase II. Meiotic spreads with (A–B) wild type *haploid*, (C–D) *Atmcm8* *haploid*, (E–F) *Atdmc1* *haploid*, (G–H) *Atmcm8/Atdmc1* *haploid* using DAPI staining. Bar, 10 μ m.

doi:10.1371/journal.pgen.1003165.g011

for CO formation. This study identified a new pathway of meiotic DSB repair independent of *AtDMC1*.

Materials and Methods

Plant material

A. thaliana accession Columbia (Col-0) was the wild type reference. *Atmcm8-1* (Salk_032764, N532764), *Atmcm8-2* (Salk_104007, N604007) *Atmcm8-3* (Salk_099327, N599327) were obtained from the collection of T-DNA mutants at the Salk Institute Genomic Analysis Laboratory (SIGnAL, <http://signal.salk.edu/cgi-bin/>

tdnaexpress) [46] via NASC (<http://nasc.nott.ac.uk/>). Other mutants used in this study were *Atspo11-2* (*Gabi_749C12*, N359272) [47], *Atprd1* (Salk_024703, N524703) [3], *Atdmc1-3* (Sail_170_F08, N871769) [48], *Atrad51* (*Atrad51-I*) [6], *asy1-4* (Salk_046272, N546272), *sds-2* (Sail_129_F09, N806294) [12], *Atzip4-2* (Salk_068052, N568052) [43], *Atmsh4* (Salk_136296, N636296) [49], *mus81-2* (Salk_107515, N607515), *mus81-3* (Salk_002761, N502761) [50,51], and *shoc1-1* (Salk_057589, N557589). *rad1-1* (*wh1-1*) has a EMS (ethyl methanesulfonate) mutation [33,34] and was provided by C. White.

Growth conditions

Plants were cultivated in greenhouse or growth chamber with a 16 h/day and 8 h/night photoperiod, at 20°C and 70% humidity.

Genetic analysis

Allelism tests were performed by crossing *Atmcm8-1*^{+/-} with *Atmcm8-2*^{+/-} and selecting F1 plants hemizygous for both alleles and likewise for *Atmcm8-2*^{+/-} with *Atmcm8-3*^{+/-}. Double mutants were obtained by crossing heterozygous plants for each mutation and selfing the double heterozygous F1 plants. *Atmcm8/Atmsh4/Atmus81* triple mutant was identified by crossing *Atmcm8/Atmsh4* double heterozygous with *Atmus81* single mutant. As *Atmsh4* and *Atmus81* are linked, a plant heterozygous for *Atmcm8/Atmsh4* was self-fertilized and homozygous for *Atmus81* to identify the triple mutant in the offspring. Haploid *Atmcm8* and *Atmcm8/Atdmc1* were obtained by crossing a heterozygous plant for *Atmcm8* or *Atmcm8/Atdmc1* mutations as male and the GEM line as female [36,52]. In F1, haploid plants of the desired genotype were selected.

Oligonucleotides for PCR genotyping

Plants of interest were selected by PCR genotyping using diagnostic primer sets. The three *AtMCM8* insertions were genotyped by PCR using following primer combinations to amplify genomic DNA flanking the T-DNA insertions. *Atmcm8-1*: left borders (LB) with LBsalk2 (5'-GCTTTCCTCCCTTCCTTTC-3')/N532764L (5'-AGCGCCATTAGCAAAATGTC-3') or with LBsalk2/N532764U (5'-GCAGCTTCATTCTGCAAGTG-3'). Wild type allele with N532764U/N532764L. *Atmcm8-2* LB with LBsalk2/N604007L (5'-TCAC-TACAGCAACGGTGAGC-3'), right border (RB) with RBsalk1 (5'-TCA GAG CAG CCG ATT GTC-3')/N604007U (5'-GCTGATGGAAGACCTTGTGG-3'). Wild type allele with N604007U/N604007L. *Atmcm8-3* LB with LBsalk2/N599327L (5'-TGGTGTGGAATCAGCAGATG-3') or with Lbsalk2/N599327U (5'-TGTGTCTCTGTTGCAAAGGC-3'). Wild type allele with N599327U/N599327L. T-DNA right and left borders were analyzed by sequencing PCR products. *ASPO11-2* wild type allele was amplified using primers 749C12U (5'-GAGCGA-GAATTTTTGGTTGG-3') and 749C12L (5'-CCACAAGG-TCAATTCTTCAAC-3') and mutant allele using N524703L and LBgabil (5'-CCCATTGGACGTGAATGTAGACAC-3'). *AtPRD1* wild type allele was amplified using primers N524703U (5'-AAGTCTGCCCATGGTCACGATTCTCTCTG-3') and N524703L (5'-GCCTGCTCAAAGGGTCCAGC-3') and mutant allele using N524703L and LbSalk2. *AtDMC1* wild type allele was amplified using primers N871769U (5'-TTTTTAATTGTT-TACAGAGGAAATCAG-3') and N871769L (5'-TCCACTCG-GAATAAAGCAATG-3') and mutant allele using N871769L and Lb3sail (5'-TAGCATCTGAATTTTCATAACCAATCTCGATACAC-3'). *AtRAD51* wild type allele was amplified using primers RAD51-1U (5'-ATGCCAAGGTTGACAAGATTG-3') and RAD51-1L (5'-CTCCCCTTCCAGAGAAATCTG-3') and mutant allele using RAD51-1U and LBgabil (5'-CCCATTG-

GACGTGAATGTAGACAC-3'). We amplified *SDS* wild type allele using primers N806294U (5'-CTGCTCCCTGATTA-CAAGCAG-3') and N806294L (5'-CTTAACGCATTCAGG-CAACTC-3') and mutant allele using N806294U and Lb3sail. *AMSH4* wild type allele was amplified using primers N636296U (5'-CTTCTTGACAGTTGTGTTT-3') and N636296L (5'-GCCAGCTGTTTTGTTGTC-3') and mutant allele using N636296L and LbSalk2. *AMUS81A* wild type allele was amplified for Salk_107515 using primers N607515U (5'-CATGCTGACAGTTGAAGGTC-3') and N607515L (5'-CCTCAAACGTTTCTCCAAAT-3') and mutant allele using N607515L and LbSalk2. *AMUS81A* wild type allele was amplified for Salk_002176 using primers N502176U (5'-CACATACGTTTTTGGTCCC-3') and N502176L (5'-AGTGTC-CAAGTCCTGCTTTC-3') and mutant allele using N607515L and LbSalk2. *AtZIP4* wild type allele was amplified using primers N568052U (5'-TCCTTCCCACACCTTGACCC-3') and N568052L (5'-GACTGCTGGAGCAGAACT-3') and mutant allele using N568052L and LbSalk2. *ASY1* wild type allele was amplified using primers N546272U (5'-TCTATGTTTGT-TACGCGTTAATCAG-3') and N546272L (5'-AGTGGCT-CGTAATCTGGTGGCTGC-3') and mutant allele using N546272L and LbSalk2. *SHOC1* wild type allele was amplified using primers N557589U (5'-TTACCGGAGTTTGAACCCG-3') and N557589L (5'-GGCAAAGACTTGAAGGCATC-3') and mutant allele using N557589L and LbSalk2. *AtRAD1* was amplified using primers o629 (5'-CTGGTGAAGAACATTTGG-TAG-3') and o630 (5'-CTCTTATGGCTGCTGCGTCTTC-3'). Polymorphism between wild type and mutant alleles was revealed with DdeI digestion.

Fluorescent tagged lines

FTL lines were obtained from G.P. Copenhaver. For this study, we used two couple of adjacent intervals: I5aI5b and I5cI5d [31]. The procedure to create plants of interest and to collect data was described in [31,53]. Statistical analysis was performed as described in [31].

Cytology, immunolocalization, and antibodies

Alexander staining for pollen viability was performed as described [54]. The protocol described by [55] was used to observe the female meiosis and the protocol described by [29] for male meiotic spreads. Immunolocalization of *AtMLH1* was made as described by [29]. Immunolocalization of *AtZYP1* and *AtDMC1* was performed according to [56] with the modifications described in [43]. The anti-ASY1 polyclonal [56] and anti-ZYP1 polyclonal [49] antibodies were used at a dilution of 1:250. The anti-MLH1 antibody [29] was used at a dilution of 1:200. The anti-DMC1 antibody was described in [43] and the purified serum was used at 1:20.

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Microscopy

For male meiotic spreads, observations were made with a Leica DM RXA2 epifluorescence microscope using an oil PL APO 100X/1.40 objective (Leica). Photographs were taken using a CoolSNAP HQ (Roper Scientific) camera driven by Open-LAB 4.0.4 software (Improvision). For immunocytology and FTLs analyzes, observations were made using a Zeiss Axio Imager2 microscope. We analyzed FTLs using the automatic slide-scanner function of the ZEISS AxioObserver DIC FISH Apotome and its workbench. Photographs were taken using an AxioCam MRm (Zeiss) camera driven by Open-LAB 4.0.4 software AxioVision 4.8. All pictures were processed with AdobePhotoshop 7.0 (Adobe Systems Inc.).

Supporting Information

Figure S1 A clustalW multiple alignment of the MCM8 protein family representatives. Black and grey shading indicate amino acid identical or similar, respectively (BLOSUM62) in at least 50% of the proteins. *At* (*Arabidopsis thaliana* MCM8), *Hs* (*Homo sapiens* MCM8 NP_115874.3), *Dm* (*Drosophila melanogaster* REC NP_732072.1), *Es* (*Entamoeba histolytica* MCM8 EAL48818.1), *Lm* (*Leishmania major* MCM8 CAB89596.2), *Pf* (*Plasmodium falciparum* MCM8 NP_701477.1). (PDF)

Figure S2 Coimmunolocalization of ASY1 and *AtMLH1*. ASY1 (red), *AtMLH1* (green) are shown as well as the overlay of both signals (merge) at diplotene in (A) wild type and in (B) *Atmcm8* mutant. Bar, 10 μ m. (TIF)

Figure S3 DMC1 immunolocalization. DNA (DAPI, blue) and *AtDMC1* (green) are shown as well as the overlay of both signals (merge) at zygotene in (A) wild type and in (B) *Atmcm8* mutant. Bar, 10 μ m. (TIF)

Acknowledgments

We thank Christine Mézard, Mathilde Grelon, and Arnaud Demuyt for critical reading of the manuscript and helpful discussions. We thank Gregory P. Copenhaver and Charles White for generously providing the fluorescent tagged lines and the *Atrad1* mutant, respectively.

Author Contributions

Conceived and designed the experiments: WC VP RM. Performed the experiments: WC VP NF LC CH NV RM. Analyzed the data: WC VP RM. Wrote the paper: WC VP RM.

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