Identification of DSB-1, a Protein Required for Initiation of Meiotic Recombination in Caenorhabditis elegans, Illuminates a Crossover Assurance Checkpoint

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Abstract

Meiotic recombination, an essential aspect of sexual reproduction, is initiated by programmed DNA double-strand breaks (DSBs). DSBs are catalyzed by the widely-conserved Spo11 enzyme; however, the activity of Spo11 is regulated by additional factors that are poorly conserved through evolution. To expand our understanding of meiotic regulation, we have characterized a novel gene, dsb-1, that is specifically required for meiotic DSB formation in the nematode Caenorhabditis elegans. DSB-1 localizes to chromosomes during early meiotic prophase, coincident with the timing of DSB formation. DSB-1 also promotes normal protein levels and chromosome localization of DSB-2, a paralogous protein that plays a related role in initiating recombination. Mutations that disrupt crossover formation result in prolonged DSB-1 association with chromosomes, suggesting that nuclei may remain in a DSB-permissive state. Extended DSB-1 localization is seen in mutants with defects in early recombination steps, including spo-11, suggesting that the absence of crossover precursors triggers the extension. Strikingly, failure to form a crossover precursor on a single chromosome pair is sufficient to extend the localization of DSB-1 on all chromosomes in the same nucleus. Based on these observations we propose a model for crossover assurance that acts through DSB-1 to maintain a DSB-permissive state until all chromosome pairs acquire crossover precursors. This work identifies a novel component of the DSB machinery in C. elegans, and sheds light on an important pathway that regulates DSB formation for crossover assurance.

Citation: Stamper EL, Rodenbusch SE, Rosu S, Ahringer J, Villeneuve AM, et al. (2013) Identification of DSB-1, a Protein Required for Initiation of Meiotic Recombination in Caenorhabditis elegans, Illuminates a Crossover Assurance Checkpoint. PLoS Genet 9(8): e1003679. doi:10.1371/journal.pgen.1003679

Editor: Jeff Sekelsky, The University of North Carolina at Chapel Hill, United States of America

Received March 13, 2013; Accepted June 14, 2013; Published August 8, 2013

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Funding: This study was supported by a Department of Energy Office of Science Graduate Fellowship (DOE SCGF, http://scgf.orau.gov/) to ELS, a National Science Foundation Graduate Research Fellowship (NSFGRF, http://www.nsfgrfp.org/) to SER, and by research funds from the NIH (http://grants.nih.gov/grants/oer.htm) (R01 GM067268 to AMV and R01 GM065591 to AFD) and the Howard Hughes Medical Institute (http://www.hhmi.org/). Some strains were provided by the CGC (http://www.cbs.umn.edu/cgc), which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440). 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

Formation of crossovers between homologous chromosomes is essential for successful execution of the meiotic program in most sexually reproducing organisms. In addition to shuffling genetic information between parental chromosomes, crossovers, together with cohesion between sister chromatids, create physical links between homologous chromosomes that enable their segregation to daughter cells during the first meiotic division [1,2]. Disruption of crossover formation leads to chromosome nondisjunction and the formation of aneuploid gametes, and thereby greatly reduces fertility.

Meiotic recombination is initiated by programmed DNA double-strand breaks (DSBs), a subset of which is repaired to form crossovers between homologous chromosomes (for a review, see [3]). While a minimum number of DSBs is needed to promote the necessary crossovers on all chromosome pairs, excessive DSBs could threaten genome integrity. The number of meiotic DSBs in various organisms appears to be highly regulated, presumably to balance the crossover requirement with the risk of mutation. The timing of programmed DSBs during early meiotic prophase is also tightly controlled to maximize the likelihood of productive repair [4,5]. For example, in Saccharomyces cerevisiae, the activities of cell cycle-regulated kinases involved in DNA replication ensure that DSBs occur only after DNA synthesis is complete [6–10]. DSB formation must also be inactivated during meiotic prophase to allow for repair prior to the meiotic divisions. Mechanisms that terminate DSB formation are not well understood, although recent studies have shown that the ATM/ATR family of DNA damage response kinases is involved in down-regulating the number of DSBs in mice, S. cerevisiae, and Drosophila melanogaster [11–14]. Further investigations are needed to better understand the mechanisms underlying these various aspects of DSB regulation.
Author Summary

For most eukaryotes, recombination between homologous chromosomes during meiosis is an essential aspect of sexual reproduction. Meiotic recombination is initiated by programmed double-strand breaks in DNA, which have the potential to induce mutations if not efficiently repaired. To better understand the mechanisms that govern the initiation of recombination and regulate the formation of double-strand breaks, we use the nematode Caenorhabditis elegans as a model system. Here we describe a new gene, dsb-1, that is required for double-strand break formation in C. elegans. Through analysis of the encoded DSBI protein we illuminate an important regulatory pathway that promotes crossover recombination events on all chromosome pairs to ensure successful meiosis.

Meiotic DSBs are catalyzed by the widely conserved, topoisomerase-related enzyme Spo11 [15,16]. Although Spo11 is essential for DSB formation, it does not function alone. In various organisms— including fungi, plants, and animals— additional proteins required for meiotic DSBs have been identified (for a review, see [17]). Unlike Spo11, other known factors involved in DSB formation are poorly conserved. For example, of five meiosis-specific DSB proteins found in S. cerevisiae, only two (Rec114 and Mei4) have known orthologs in other phyla; and even these two proteins are absent in several species, including Caenorhabditis elegans, D. melanogaster, and Neurospora crassa [18]. Additional DSBI proteins have also been identified in other organisms, but none are ubiquitous among eukaryotes [5,19–22].

The nematode C. elegans has emerged as a valuable model system for molecular analysis of meiosis. As in other eukaryotes, SPO-11 catalyzes the formation of meiotic DSBs [23]. MEJ-11 and RAD-50 are also required for DSB formation [24,25] as in S. cerevisiae [17], but these proteins have other essential roles in DNA metabolism, including in the resection of meiotic DSBs [3,26]. In C. elegans, as in other species, meiosis-specific chromosome architecture contributes to DSBI proficiency. In particular, in the absence of HTP-3, an integral component of chromosome axes, DSBs are abolished or sharply reduced [27]. The related protein HTP-1, which is also associated with the axial elements, may also contribute to DSBI formation, while other axial components appear to be dispensable for DSBs [28–30]. Roles for axis components homologous to HTP-3 and HTP-1 in promoting DSBs have also been demonstrated in other organisms [3,31,32]. Additionally, the meiotic kinase CHK-2, which regulates many key events during early meiotic prophase, is required for programmed DSBs in C. elegans [33]. Several other factors are known to influence meiotic DSBI formation, but their effects may be indirect. These include the chromatin-associated proteins HIM-5, HIM-17, and XND-1, which promote normal levels of meiotic DSBs, but whose functions are pleiotropic and not well understood [34–36]. Apart from SPO-11, no protein that specifically functions in initiating recombination has previously been reported. Some aspects of C. elegans meiosis are unusual among model organisms, including the fact that synopsis between homologous chromosomes is independent of recombination [23]. Thus, analysis of DSBI regulation in C. elegans will likely reveal both conserved aspects of meiosis and how regulatory circuits are remodeled during evolution.

Here we identify a novel gene, dsb-1 (double-strand break factor 1), that is required for meiotic DSB formation in C. elegans. dsb-1 mutants lack meiotic DSBs, and show meiotic defects similar to spo-11 mutants. DSBI localizes to meiotic chromosomes coincident with the time of DSB formation, in a manner dependent on the CHK-2 kinase. We also find that a variety of mutations that disrupt crossover formation on one or more chromosomes extend the chromosomal localization of DSBI, suggesting that the DSBI-permissive state may be prolonged. Based on these observations, we infer the existence of a regulatory circuit in which meiotic checkpoint proteins monitor the recombination status of each chromosome pair and act through DSBI to maintain a DSBI-permissive state until all chromosome pairs have attained crossover-competent recombination intermediates.

Results

Identification of dsb-1, a New Gene Required for Meiotic DSB Formation

In C. elegans, mutations that impair meiotic chromosome segregation result in embryonic lethality and a high incidence of males (XO) among the surviving progeny [37]. The dsb-1(we11) mutant was isolated in a genetic screen for maternal-effect embryonic lethality, and was found to produce a high fraction of males among its few surviving self-progeny. A targeted deletion allele of the affected gene, dsb-1(tm5034), was generated independently (see below), and results in defects identical to dsb-1(we11) based on all assays described here. Whereas self-fertilizing wild-type hermaphrodites produce nearly 100% viable progeny and 0.2% males (Figure 1A, [37]), only 3% of progeny from self-fertilizing dsb-1 mutant hermaphrodites survived to adulthood (n>2000; 12 broods), (Figure 1A, Table 1). Among these survivors, 36–38% were male (Figure 1A, Table 1). The brood size (number of fertilized eggs) of self-fertilizing dsb-1 hermaphrodites was also reduced relative to wild-type animals (Table 1).

Chromosome segregation errors in meiosis often reflect defects in crossover formation between homologs. The levels of embryonic lethality and male progeny observed in dsb-1 mutants are quantitatively similar to several previously characterized mutants that fail to make any crossovers during meiotic prophase, such as spo-11 (Figure 1A, Table 1), msh-5, and cos-1 [25,38,39], suggesting that dsb-1 mutants might also lack crossovers. Visualization of DAPI-stained oocytes at diakinesis provides a simple assay for crossover formation in C. elegans. In wild-type hermaphrodites, 6 DAPI-staining bodies are observed in each oocyte nucleus (average = 5.8, Figure 1B and 1C), corresponding to 6 pairs of homologous chromosomes, each held together by a chiasma [40]. In mutants that fail to make crossovers, oocytes typically display 12 DAPI-staining bodies. The number and morphology of DAPI-staining bodies observed in dsb-1 mutant oocytes was similar to spo-11 mutants (average = 11.6, Figure 1B and 1C), indicating an absence of chiasmata in dsb-1 animals.

We investigated whether the disruption of crossover formation in dsb-1 mutants might reflect a defect in homologous chromosome pairing or synopsis. Pairing was assessed using fluorescence in situ hybridization (Figure 1D). Early pachytene nuclei of both wild-type and dsb-1 animals contained a single focus or closely apposed pair of foci, indicating that homologous chromosomes were paired (Figure 1D). Further, co-staining of the axial element protein HTP-3 and the synaptonemal complex central region protein SYP-1 indicated that chromosomes were fully synapsed by early pachytene in dsb-1 animals (Figure 1E), as in wild-type animals. These results indicate that dsb-1 mutants are proficient for homologous chromosome pairing and synopsis.

To assess whether dsb-1 mutants initiate meiotic recombination, we used antibodies against the DNA strand-exchange protein RAD-51, which binds to single-stranded regions adjacent to
Figure 1. *dsb-1* mutants lack meiotic crossovers but are proficient for homologous chromosome pairing and synapsis. (A) Quantification of viable and male self-progeny for the indicated genotypes is shown. Homozygous *dsb-1* (we11 and tm5034) hermaphrodites produce many inviable and male self-progeny compared to wild-type (WT) animals, similar to *spo-11* hermaphrodites. For each bar, the upper number indicates the percentage, and the lower number in parentheses indicates the total number of fertilized eggs (for viability) or adult progeny (for males) counted. (B) Histogram showing the number of DAPI-staining bodies observed in oocytes at diakinesis for the indicated genotypes. 100 oocytes were counted for each genotype. The numbers in parentheses indicate: fertilized eggs (for viability), adult progeny (for male counts), and broods (for brood size) counted. Each of the two sets of experiments was performed in parallel under identical conditions. (C) Each panel shows a representative DAPI-stained oocyte nucleus at the indicated genotypes. (D) Fluorescence in situ hybridization against the SS rDNA genomic locus (SSrDNA) and a locus on the right end of the X chromosome (Xrt) indicated that pairing is not disrupted in *dsb-1* mutants. (E) Synapsis was assessed by immunofluorescence staining of the axial element protein HTP-3 and central region protein SYP-1, components of the synaptonemal complex. Full colocalization of the two markers indicates fully synapped chromosomes. Scale bars, 5 μm.

Table 1. Progeny viability, incidence of males, and brood size from *dsb-1* and *dsb-2* mutants.

| Genotype                      | % Viability (n) | % Males (n) | Brood Size (n) |
|-------------------------------|----------------|-------------|----------------|
| Wild-type                     | 102 (2933)     | 0.20 (2991) | 285 (10)       |
| *spo-11* (me44)               | 2.5 (4063)     | 37 (103)    | 343 (11)       |
| *dsb-1* (we11)                | 3.2 (2142)     | 38 (68)     | 184 (12)       |
| *dsb-1* (tm5034)              | 3.2 (2134)     | 36 (69)     | 178 (12)       |
| *rol-1*                       | 98 (1741)      | 0.06 (1705) | 290 (6)        |
| *rol-1* *dsb-2* (me96)        | 29 (789)       | 15 (229)    | 263 (3)        |
| *rol-1* *dsb-1* (tm5034)      | 3.7 (782)      | 38 (29)     | 196 (4)        |
| *rol-1* *dsb-2* (me96) *dsb-1* (tm5034) | 3.8 (1500) | 35 (57) | 214 (7) |

Quantification of the viable and male self-progeny from hermaphrodites of indicated genotypes, as well as average brood size (fertilized eggs), is shown. The numbers in parentheses indicate: fertilized eggs (for viability), adult progeny (for male counts), and broods (for brood size) counted. Each of the two sets of experiments was performed in parallel under identical conditions.

doi:10.1371/journal.pgen.1003679.g001
dsb-1 Encodes a Member of a Novel Protein Family

Whole-genome sequencing of backcrossed dsb-1(we11) animals identified several mutations in annotated coding sequences, including a nonsense mutation in the previously uncharacterized gene F08G5.1 (Figure 3A), which encodes a predicted protein of 385 amino acids and seemed a plausible candidate based on its meiosis-enriched expression pattern [45]. We found that knockdown of F08G5.1 expression via transgene-mediated cosuppression [46] caused embryonic lethality and male progeny, as well as strong reduction of chiasmata in oocytes at diakinesis was quantified. Similar to spo-11 and WT, irradiated dsb-1 mutants displayed 6 DAPI-staining bodies. Scale bar, 5 μm. (D) Quantification of the number of DAPI-staining bodies observed for each genotype in untreated and irradiated animals. The top number for each bar indicates the average number of DAPI-staining bodies and the number in parentheses indicates the number of oocytes counted. (E) WT, spo-11, and dsb-1 mutants were gamma-irradiated (10 Gy) at 4–5 hours post L4. Eggs laid 20–30 hours post irradiation were counted for progeny survival. The percent survival of dsb-1 mutants partially rescued progeny survival. As for spo-11, irradiation of dsb-1 mutants partially rescued progeny survival.

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Figure 2. dsb-1 is required to initiate meiotic recombination. (A, B) Immunofluorescence staining of RAD-51 was used as a cytological marker of early recombination intermediates in early pachytene nuclei in (A) untreated and (B) irradiated animals. Scale bar, 5 μm. (A) WT nuclei display abundant RAD-51 foci, however, dsb-1 mutants lack RAD-51 foci. (B) dsb-1 mutants were gamma-irradiated (10 Gy) as young adults, and then assessed 2 hours later for the presence of RAD-51 foci. RAD-51 foci were present on chromosomes in dsb-1 mutants after irradiation. (C, D) WT, spo-11(me44), and dsb-1 mutants were gamma-irradiated (10 Gy) as young adults. After 18 hours, irradiated and control animals were fixed, and the number of DAPI-staining bodies in oocytes at diakinesis was quantified. Each panel shows DAPI-stained chromosomes in a single, representative oocyte at diakinesis for the indicated genotype. Similar to spo-11 and WT, irradiated dsb-1 mutants displayed 6 DAPI-staining bodies. Scale bar, 5 μm. (D) Quantification of the number of DAPI-staining bodies observed for each genotype in untreated and irradiated animals. The top number for each bar indicates the average number of DAPI-staining bodies and the number in parentheses indicates the number of oocytes counted. (E) WT, spo-11, and dsb-1 mutants were gamma-irradiated (10 Gy) at 4–5 hours post L4. Eggs laid 20–30 hours post irradiation were counted for progeny survival. The percent survival of dsb-1 mutants partially rescued progeny survival. As for spo-11, irradiation of dsb-1 mutants partially rescued progeny survival.

dsb-1 introduces a premature stop (1ac = >taa) after lysine 96 (Figure 3A). A targeted deletion allele (tm5034) removes 290 bp from predicted exons 3 and 4 and the intervening intron (Figure 3A), resulting in a frameshift mutation that introduces a glutamine immediately followed by a stop codon after lysine 96. The phenotype of dsb-1(tm5034) mutants is indistinguishable from dsb-1(we11) (Figure 1 and 2, Table 1). Both are predicted to lack functional protein based on the early stop codons, and this conclusion is supported by immunofluorescence and immunoblotting experiments (below). Based on the evidence described above that mutations disrupting F08G5.1 specifically interfere with meiotic double-strand break formation, we designated F08G5.1 as dsb-1, for double-strand break factor 1.

The DSB-1 protein has no apparent homologs outside of the genus Caenorhabditis, including other nematode genera. Interestingly, the genomes of C. elegans and several other Caenorhabditids
DSB-1 Localizes to Chromosomes during Early Meiotic Prophase

To further probe the role of DSB-1 in the formation of meiotic DSBs, we generated an antibody against the full-length protein expressed in *E. coli*. Immunofluorescence staining revealed that DSB-1 is absent from somatic nuclei, and specifically localizes to chromosomes during early meiotic prophase (Figure 4A and 4B), while *dsb-1* mutants showed only background staining (Figure S3). Accumulation of DSB-1 on chromosomes was first observed in nuclei marked by crescent-shaped DAPI-staining morphology, corresponding to the “transition zone” (leptotene/zygotene), and disappeared at mid-pachytene (Figure 4A). Chromosomal localization of DSB-1 preceded the appearance of RAD-51 foci, consistent with an early role for DSB-1 in meiotic recombination (Figure 4A). Thus, the localization of DSB-1 to chromosomes corresponds to the period during which DSBs are likely to be generated.

While most nuclei in the late pachytene region of the germline lacked DSB-1 staining, we consistently observed dispersed nuclei in this region that retained bright fluorescence (Figure 4A and S4). These nuclei also contained abundant RAD-51 foci and frequently displayed compact chromosome morphology resembling that seen in the transition zone, along with evidence of asynapsed chromosomes (Figure S4A and S4B). We tested whether these late DSB-1 positive nuclei might be apoptotic by examining *ced-4* mutants, which lack germ line apoptosis [49,50], and found that they were still present in similar numbers (data not shown). The persistence of RAD-51 foci and asynapsed chromosomes suggest that these nuclei may be delayed in completing synapsis or other prerequisites for crossover formation, a conclusion reinforced by further analysis of DSB-1 regulation, described below.

DSB-1 was distributed as a network of foci and stretches of staining on meiotic chromosomes (Figure 4B). One chromosome consistently showed weaker DSB-1 staining. This seemed likely to be the X chromosome, which has many unique features in the germ line, including distinct chromatin marks [51,52] and genetic requirements for meiotic DSBs [35,36]. Co-staining with antibodies against HIM-8, which specifically mark the X chromosome [53], confirmed that the chromosome pair with weaker DSB-1 staining was the X (Figure 4C).

DSB-1 and RAD-51 both localized to chromosomes during early pachytene (Figure 4A). However, we found that RAD-51 did not colocalize with DSB-1 (Figure 4D). Similar findings have been reported for DSB proteins in mice and *Schizosaccharomyces pombe* [18,54]. This could indicate that DSB-1 does not act directly at DSB sites, or that it is removed from DSB sites prior to RAD-51 loading.

Meiotic chromosomes are believed to be organized as chromatin loops tethered at their bases to the proteinaceous chromosome axis [55,56]. Based on work from *S. cerevisiae*, it has been proposed that DSBs occur at sites within chromatin loops that are recruited to the chromosome axis [57–59]. Most DSB-1 staining was associated with the periphery of DAPI-stained chromosomes rather than axes (Figure 4E), suggesting that DSB-1 is primarily associated with chromatin loops. This localization pattern is similar to what has been observed for several DSB proteins in *S. cerevisiae* [6,60–63].

Chromosome Localization of DSB-1 Is Independent of SPO-11, the MRN Complex, and a Properly Assembled Meiotic Chromosome Axis

We tested whether DSB-1 localization depends on other factors required for DSB formation. DSB-1 localized to meiotic chromosomes in the catalytically dead spo-11(tem44) mutant [25], as well as in spo-11( ok79) mutants, which lack functional protein [23], indicating that DSB-1 localizes to chromosomes independently of DSBs and SPO-11 (Figure 5). DSB-1 localization was also independent of MRE-11 and RAD-50 (Figure 5, data not shown), which are required for DSB formation in *C. elegans* [24,25]. In *htp-3* mutants, which lack an essential axial element component that is important for DSB formation [27], DSB-1 was detected on meiotic chromosomes (Figure 5), but the staining appeared somewhat reduced compared to wild-type nuclei.
Chromosome Localization of DSB-1 is Dependent on the Meiotic Kinase CHK-2

The CHK-2 kinase is essential for several key events during early meiotic prophase in *C. elegans*, including DSB formation and homolog pairing [33]. We found that nuclear staining of DSB-1 was strongly reduced, albeit still detectable, in *chk-2(me64)* mutants (Figure 5). Although the intensity of DSB-1 staining was sharply reduced in *chk-2* mutants, it appears that faint fluorescence observed upon prolonged exposure reflects DSB-1, because the nuclear staining pattern resembles that seen in wild-type animals, and because chromosomal staining is not detected prior to meiotic entry. Western blot analysis revealed that DSB-1 protein is expressed in *chk-2* mutants, although the protein levels appear somewhat reduced compared to wild-type (data not shown).

However, the reduction in DSB-1 protein levels in *chk-2* mutants does not appear to fully account for the sharply diminished chromosomal localization of DSB-1. These data indicate that DSB-1 localization to chromosomes is largely dependent on the CHK-2 kinase, and suggest that DSB-1 may act downstream of CHK-2 to promote DSBs.

Disruption of Crossover Formation Prolongs Chromosome Localization of DSB-1

In testing the genetic requirements for DSB-1 localization, we noticed that the zone of DSB-1 staining in the gonad was extended in mutants that disrupt crossover formation. Previous studies have reported a persistence of RAD-51 foci in numerous mutants that are proficient for DSBs but not for crossovers [43,64,65]. In wild-
of the DSB-1-positive region to the LZP region – was consistent across age-matched animals of the same genotype. In wild-type adult hermaphrodites, DSB-1 positive nuclei comprised about 50% of the length of the LZP region (Figure 6A and 6B). However, in most mutants that disrupt crossover formation on one or more chromosomes, this zone of DSB-1 staining was significantly extended (Figure 6A and 6B). We saw some variability in this extension, which tended to correlate with the nature of the mutation: Mutations affecting late steps in the crossover pathway, including msh-5(me23) [30], cos-1(me13) [39], and zhp-3(jf61) [66], extended the DSB-1 zone to ~75%, of the LZP region (Figure 6A and 6B). Mutations that block earlier steps in homologous recombination, including com-1(t1626) [67], rad-51(kg8701) [42], and rad-54(tm1266) [65], extended the DSB-1 zone even further, to ~90% of the LZP region. Mutations that block crossover formation by disrupting synopsis, including syb-1(mz17) [63] and syb-2(ok307) [43], also showed an extension of DSB-1 staining to ~90% (Figure 6B).

Significantly, mutants that lack meiotic DSBs, including spo-11(ok79 or me44) [23,25], mre-11(ok179) [24], and rad-50(ok197) [25], also showed significant extension of the zone of DSB-1 staining to 69–78% of the LZP region (Figure 6A and 6B). Together these findings indicate that the absence of crossovers or crossover precursors, rather than the presence or persistence of earlier recombination intermediates, triggers extension of the DSB-1 zone.

Also of note, in htp-1 and htp-3 mutants [27–29], in which the axial element is disrupted, the region of DSB-1 staining was shorter than in other crossover-deficient mutants (Figure 6B), despite the fact that no crossovers form in these animals and DSBs are either eliminated or reduced [27–29]. This suggests that axis structure may play a role in detecting or signaling the absence of crossover precursors to prolong DSB-1 localization, consistent with proposed roles for the axis in other species [32,69–71].

We tested whether irradiation could suppress the extension of the DSB-1 zone seen in spo-11 mutants. Young adult hermaphrodites were irradiated, then fixed and stained 8 hours later. As controls, we included mutants (mre-11 and msh-5) in which crossover defects are not rescued by exogenous DSBs [24,30]. Irradiation reduced the zone of DSB-1 staining in spo-11(me44) animals to 56%, compared to 70% for unirradiated controls (Figure 6C). In contrast, the length of the DSB-1 zone in wild-type, mre-11, and msh-5 hermaphrodites was unaffected by irradiation (Figure 6C). These data reinforce the idea that the absence of crossovers or crossover precursors induces prolonged DSB-1 association with chromosomes.

Many mutations that result in extension of the DSB-1 zone also cause elevated oocyte apoptosis, which can be triggered in response to persistent DNA damage or asynapsis [43,50,68,72]. We considered the possibility that apoptosis might mediate the observed extension of DSB-1 staining, since this process primarily culls nuclei from the late pachytene, DSB-1 negative region of the gonad (reviewed in [73]). To test this idea, a representative subset of meiotic mutations, including spo-11(ok79), msh-5, syb-2, him-8, and zim-2 (see below) were combined with ced-4(n1162), which abrogates germline apoptosis [49]. These double mutants displayed extended DSB-1 localization similar to that observed in the corresponding single mutants (Figure 6D). We conclude that apoptosis does not account for the extension of DSB-1 staining observed in crossover-defective mutants, nor can it explain the quantitative differences observed among different mutants.
DSB-1 Illuminates a Meiotic Crossover Checkpoint

A

WT  rad-54
WT  msh-5
spo-11(me44)  zhp-3
spo-11(ok79)  him-8

DAPI  DSB-1

B

axis defect  no DSBs  early defect in recombination  late CO defect  lacking CO on one chromosome

Percent of LCP w/ DSB-1

WT  syp-1  syp-2  hlp-1  hlp-3  spo-11(ok79)  spo-11(me44)  mre-11  rad-50  com-1  rad-51  rad-54  msh-5  cosA  zhp-3  him-8  zim-2  him-5

(17)  (13)  (9)  (7)  (5)  (9)  (16)  (4)  (6)  (6)  (5)  (7)  (10)  (4)  (9)  (7)  (9)

C

control  irradiated

Percent of LCP w/ DSB-1

WT  mre-11  msh-5  spo-11  him-8  him-5

(6)  (9)  (5)  (11)  (5)  (5)  (13)  (6)  (8)  (9)  (17)
Extension of DSB-1 Localization Reflects a Genome-Wide and Nucleus-Autonomous Response

To further characterize the extension of DSB-1 localization that occurs in response to defects in crossover formation, we examined mutant situations in which crossover formation was disrupted on only one chromosome. him-8(n611) and zim-2(tm574) specifically disrupt homolog pairing and thus crossover formation on chromosomes X and V, respectively [53,74]. him-5(k1936) does not impair pairing or synapsis, but abrogates DSBs on the X chromosome [36]. All three of these mutations extended the DSB-1 zone to 83-86% of the LZP region (Figure 6A and 6B). Furthermore, irradiation of him-5 animals, in which the crossover defect can be rescued by exogenous DSBs [36], but not irradiation of him-8, suppressed the extension of DSB-1 localization (Figure 6C). These results indicate that the absence of a crossover precursor on a single chromosome pair is sufficient to prolong DSB-1 association with meiotic chromosomes.

Analysis of mutants with chromosome-specific defects in interhomolog recombination also allowed us to test whether DSB-1 staining is specifically prolonged on crossover-deficient chromosomes. In him-5 and him-8 mutants, the autosomes, but not the X chromosomes, are proficient for crossover formation. X chromosomes can be specifically marked in these mutants using HIM-8 antibodies (in him-5 mutants) or by staining for synaptonemal complex components (in him-8 mutants). In both of these genotypes, we observed persistent DSB-1 staining on all chromosomes throughout the region of extended DSB-1 localization (Figure 7A, 7B, and 7C). As in wild-type nuclei, the X chromosome showed weaker DSB-1 staining than the autosomes (Figure 4C and 7A). These findings indicate that the extension of DSB-1 localization is a genome-wide response affecting all chromosomes within the nucleus.

To test whether the extension of DSB-1 localization is regulated by nuclear-intrinsic or extrinsic signals, we examined animals heterozygous for meDf2, a deficiency of the X chromosome pairing center [40]. In meDf2/+ hermaphrodites, X chromosome pairing and synapsis is partially compromised, such that approximately half the nuclei achieve full pairing and synapsis by the end of the pachytene region [75]. Nuclei with asynapsed X chromosomes can be recognized by their more condensed, transition zone-like chromosome morphology, or by co-staining for axial element and central region proteins of the synaptonemal complex [75]. In the late pachytene region of these animals, we found that DSB-1 staining correlated with the status of individual nuclei: those with asynapsed chromosomes were positive for DSB-1 staining, while fully synapsed nuclei lacked DSB-1 staining (Figure 7D). These results indicate that the extension of DSB-1 localization is a response to a signal intrinsic to individual nuclei, and does not extend to neighboring nuclei within the same region of the gonad. However, as in all mutants examined, DSB-1 disappeared by the end of the pachytene region of the gonad, indicative of an extrinsic, spatially regulated “override” signal that triggers progression to late pachytene and loss of the presumptive DSB-1 localization relative to the length of the LZP region. Numbers in parentheses indicate the number of gonad arms quantified. Error bars indicate standard deviation. Genotypes are color-coded based on the category of meiotic defect that they cause (listed above the graph).
permisive state, even when crossover precursors have not been attained on all chromosomes (see Discussion).

Functional Relationships between DSB-1 and DSB-2

The DSB-1 paralog DSB-2 is also involved in meiotic DSB formation [47]. As reported in the accompanying paper by Rosu et al., the two proteins show very similar localization patterns (Figure 8A and 8B, [47]). Both localize to nuclei from leptotene/zygotene through mid pachytene, although DSB-1 staining appears slightly earlier than DSB-2 staining (Figure 8A). They also disappear simultaneously from meiotic chromosomes, both in wild-type animals and various mutants that disrupt crossover formation (Figure 8A, data not shown). Additionally, both proteins show similar distributions along meiotic chromosomes (Figure 8B). Intriguingly, however, the two proteins do not extensively colocalize, but instead rarely coincide (Figure 8B).

To probe the functional interactions between DSB-1 and DSB-2, we localized each protein in the absence of the other. We found that DSB-1 localized to chromosomes in *dsb-2(me96)* mutants, although the fluorescence intensity was reduced relative to wild-type gonads (Figure 9A and 9B; see also [47]). The DSB-1 positive region of the gonad was also somewhat shorter (Figure 9A), despite the reduction of crossovers in *dsb-2* mutants [47]. This suggests that localization of DSB-1 to meiotic chromosomes does not require, but may be reinforced or stabilized by, DSB-2. By contrast, DSB-2 was not detected on meiotic chromosomes in *dbb-1* mutants (Figure 9B). Immunoblotting of whole-worm lysates revealed that DSB-1 protein levels are moderately reduced in *dbb-2* mutants, while DSB-2 protein levels are severely reduced in *dbb-1* mutants (Figure 9C). This parallels our conclusions from *in situ* localization of these proteins, and suggests that the reduction of staining observed on chromosomes is a consequence of lower protein levels.

We also tested the effect of eliminating both DSB-1 and DSB-2 by constructing a double mutant strain. The phenotypes observed in *dbb-1; dbb-2* mutant animals were indistinguishable from *dbb-1* mutants (Figure 10A and 10B). This result is consistent with the idea that these proteins collaborate in some way to promote DSB formation, and argues against more complex epistasis scenarios.

Discussion

DSB-1 and DSB-2 Mediate Initiation of Meiotic Recombination

We have discovered a novel protein, DSB-1, required for meiotic DSB formation in *C. elegans*. Our data indicate that DSB-1 acts specifically to promote DSBs, and does not play a major role in DNA repair or other meiotic processes. DSB-1 localizes to chromosomes during meiotic prophase, concomitant with the period of DSB formation. It appears more abundant on the

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Figure 8. DSB-1 and DSB-2 show similar patterns of localization, but do not colocalize. (A) Composite projection image of a gonad from a WT animal showing DAPI and immunofluorescence staining for DSB-1 and DSB-2. Both proteins are detected on chromosomes by the transition zone (leptotene/zygotene), and persist until mid pachytene. Scale bar, 30 μm. (B) Projections of early pachytene nuclei showing DSB-1 and DSB-2 immunofluorescence and DAPI staining. Although DSB-1 and DSB-2 have similar staining patterns on meiotic chromosomes, they do not colocalize extensively. Scale bar, 5 μm. doi:10.1371/journal.pgen.1003679.g008
Figure 9. DSB-1 promotes the protein localization and protein level of DSB-2. (A) Composite projection image of a gonad from a dsb-2(me96) animal showing DAPI and immunofluorescence staining for DSB-1 and DSB-2. Scale bar, 30 μm. (B) Immunofluorescence staining of DSB-1 and DSB-2 in early pachytene nuclei from WT, dsb-2(me96), dsb-1(we11), and dsb-1(tm5034) animals. Reduced DSB-1 staining is observed in dsb-2 mutants, while DSB-2 is undetectable on chromosomes in dsb-1 mutants. Scale bar, 5 μm. (C) Immunoblotting of DSB-1 and DSB-2 in WT, dsb-1(tm5034), and dsb-2(me96) animals. Lysate from 50 animals, 24 hours post L4, was used for each lane. DSB-1 protein levels are slightly reduced in dsb-2 mutants, while DSB-2 protein levels are severely reduced in dsb-1 mutants. The loading control (bottom panel) is a non-specific band. Panels were taken from the same blot. * indicates a non-specific background band.

doi:10.1371/journal.pgen.1003679.g009

Figure 10. dsb-2; dsb-1 double mutants fail to make DSBs. (A) Quantification of viable and male self-progeny produced by rol-1, rol-1 dsb-2(me96), dsb-1(tm5034), and rol-1 dsb-2(me96); dsb-1(tm5034) hermaphrodites. Whole broods were counted for each genotype. dsb-2; dsb-1 double mutants display levels of progeny viability and males similar to dsb-1 mutants. For each bar, the upper number indicates the percentage, and the lower number in parentheses indicates the total number of fertilized eggs (for viability) or adult progeny (for males) counted. (B) A representative DAPI-stained oocyte nucleus at diakinesis for rol-1 dsb-2(me96); dsb-1(tm5034) showing 12 DAPI-staining bodies. Scale bar, 5 μm. (C) Diagram summarizing the dependencies of DSB-1 and DSB-2 on each other and CHK-2. CHK-2 promotes the chromosomal localization of DSB-1 and DSB-2. Because DSB-2 localization depends on DSB-1, it is unknown whether CHK-2 promotes the loading of DSB-2 directly and through its effect on DSB-1. DSB-1 and DSB-2 promote or stabilize the protein levels and chromosomal localization of each other, with the effect of DSB-1 on DSB-2 being much greater than the converse. DSB-1 and DSB-2 collaborate to promote DSBs and crossover formation.

doi:10.1371/journal.pgen.1003679.g010

autosomes than the X chromosome. The significance of this finding is unclear, since DSB-1 is clearly required for DSBs on all chromosomes, but it may be related to observations that the X chromosome has distinct chromatin structure and differential
genetic requirements for DSB formation [35,36,51,52].

Both DSB-1 and its paralog DSB-2 are required for normal levels of meiotic DSBs. These proteins show a similar temporal and spatial pattern of localization to meiotic chromosomes. The localization of both proteins is also extended to a similar extent in mutants that disrupt crossover formation. In mutants where the localization of both DSB-1 and DSB-2 was assayed simultaneous-ly, as well as in wild-type animals, the proteins localize to the same subset of meiotic nuclei, except that DSB-1 appears slightly earlier, suggesting that they are co-regulated. However, these proteins seem unlikely to act as a complex, since they show little if any colocalization.

Although DSB-1 and DSB-2 appear to play similar roles in meiotic DSB formation, the severity of their mutant phenotypes are not equivalent. As shown by Rosu et al., DSBs are reduced but not eliminated in young dsb-2 mutant hermaphrodites [47], while dsb-1 mutants lack DSBs regardless of age. The less severe defects observed in young dsb-2 mutants likely reflect the presence of substantial residual DSB-1 protein on meiotic chromosomes in dsb-2 mutants, whereas DSB-2 is not detected on chromosomes in dsb-1 mutants, and protein levels are severely reduced. DSB-1 appears to stabilize DSB-2, perhaps by promoting its association with chromosomes, and to a lesser extent is reciprocally stabilized/reinforced by DSB-2.

The CHK-2 kinase promotes the chromosomal association of DSB-1. CHK-2 is also required for DSB-2 localization on meiotic chromosomes [47], although it is not clear whether CHK-2 promotes DSB-2 loading directly, or indirectly through its role in the loading of DSB-1. Our findings suggest a model in which DSB-1 and DSB-2 mutually promote each other’s expression, stability, and/or localization, with DSB-2 depending more strongly on DSB-1, to promote DSB formation (Figure 10C).

The number of sites of DSB-1 and DSB-2 localization per nucleus – too many to quantify in diffraction-limited images – appears to greatly exceed the number of DSBs, estimates of which have ranged from 12 to 75 per nucleus [65,76,77]. DSB-1 and DSB-2 may each bind to sites of potential DSBs, with only a subset of these sites undergoing DSB formation, perhaps where they happen to coincide. They could also be serving as scaffolds to recruit other factors required for DSB formation to meiotic chromosomes and/or to promote their functional interaction. This idea is currently difficult to test, since we have not yet been able to detect chromosomal association of SPO-11 in C. elegans, and no other proteins specifically required for DSBs have been identified.

Alternatively, these proteins may influence DSB formation by modifying chromosome structure. We did not observe overt changes in chromosome morphology in dsb-1 mutants, but further analysis – e.g., mapping of histone modifications – may be necessary to uncover more subtle changes.

A Crossover Assurance Checkpoint Mechanism That Regulates DSB Formation

DSBs normally occur within a discrete time window during early meiotic prophase. In C. elegans this corresponds to the transition zone and early pachytene, based on RAD-51 localization. As DSB-1 is necessary for DSB formation, and its appearance on meiotic chromosomes coincides with the timing of DSBs, we infer that the chromosomal localization of DSB-1 is indicative of a regulatory state permissive for DSB formation. We observed that when crossover formation is disrupted, this DSB-1-positive region is extended. Rosu et al. report a similar extension of DSB-2 in crossover-defective mutants [47].

Previous work has shown that RAD-51 foci persist longer and accumulate to greater numbers in various mutants that make breaks but not crossovers [43,64,65,67,75]. Extended or elevated RAD-51 staining could reflect an extension of the time that DSBs are made, a greater number of DSBs, or a slower turnover of the RAD-51-bound state. However, persistence of DSB-1 and DSB-2 on meiotic chromosomes in these mutants suggests that the period in which nuclei are competent for DSB formation is extended. In support of this idea, in many crossover-defective mutants defective, the number of RAD-51 foci per nucleus not only reaches a far higher level but also peaks later than in wild-type animals and continues to rise even after RAD-51 is normally cleared from meiotic chromosomes [53,75,77], indicating that breaks continue to be generated after their formation would normally cease. Several mutations that impair crossover formation on a limited number of chromosome pairs result in altered crossover distributions on the crossover-competent chromosomes [64,78], which, particularly in light of the current findings, seems likely to reflect changes in DSB activity. In addition, RAD-51 chromatin immunoprecipitation data from our laboratory (C. V. Kotwaliwale and AFD, unpublished) have indicated that the DSB distribution is altered in kin-6-0 mutants, one of the genotypes that show extended DSB-1 staining.

Taken together, these findings strongly suggest that both the temporal and genomic distribution of DSBs is altered in many situations that perturb crossover formation. A similar phenomenon may also account for the “interchromosomal effect” first observed in Drosophila female meiosis [79]. We note that this raises caveats about previously published estimates of DSB numbers in C. elegans that have been based on quantification of RAD-51 foci in genotypes that do not complete crossovers, such as rad-54 or syp-1 mutants [65,76,77].

Extension of the DSB-1 zone occurs even when nuclei are unable to initiate meiotic recombination due to an absence of DSBs. This suggests that the extension is not due to the persistence of unresolved recombination intermediates, but is instead a response to the absence of a particular crossover-competent recombination intermediate, or crossover precursor. We found that disruption of crossover formation on a single pair of chromosomes is sufficient to prolong DSB-1 localization on all chromosomes. Based on this result we believe that chromosomes lacking a crossover precursor may emanate a signal that sustains a DSB-permissive state within the affected nucleus. Thus, a single chromosome pair lacking a crossover precursor elicits a genome-wide response that results in extension of DSB-1 localization, which may reflect a modulation of the timing, and perhaps the extent, of DSB formation. Such a mechanism would help to ensure formation of an “obligate” crossover on every chromosome pair.

All mutants that we found to extend the localization of DSB-1 cause a disruption in crossover formation, although they have various primary molecular defects. It is possible that extension of DSB-1 localization occurs in response to distinct molecular triggers in different mutant situations. For example, spo-11 mutants may be responding to an absence of DSBs, rad-54 mutants to un repaired DSBs, and syp-1 mutants to asynapsed chromosomes. A similar model in which different unfinished meiotic tasks can elicit delays in meiotic progression was proposed in a recent study [90]. However, we feel that a parsimonious interpretation of our data is that the absence of a crossover precursor on one or more chromosomes is sufficient to prolong DSB-1/2 localization. The varying degree of extension seen in different mutants could reflect the engagement of additional regulatory mechanisms, such as the synopsis checkpoint and/or DNA damage checkpoint, which might converge with a crossover assurance mechanism to modulate regulators of DSB-1.
We propose that an “obligate crossover” checkpoint mediates the extension of DSB-1 localization (Figure 11). Our data suggest that DSB formation is activated during early meiosis and normally persists long enough for most nuclei to attain crossover precursors on all chromosomes (Figure 11). If interhomolog recombination is impaired on one or more chromosome pairs, individual nuclei can prolong the DSB-permissive state in an attempt to generate a crossover on every chromosome. Our observation that a block to crossover formation on a single pair of chromosomes results in persistent DSB-1 throughout the affected nuclei is reminiscent of the spindle assembly checkpoint (SAC), in which failure of a single pair of sister kinetochores to biorient on the mitotic spindle triggers a cell-autonomous delay in anaphase onset that affects cohesion on all chromosomes [81]. Interestingly, a key mediator of the SAC, Mad2, is homologous to the meiotic axis proteins HTP-3 and HTP-1 [28,82], which appear be important for the regulatory circuit that mediates prolonged DSB-1 localization in response to crossover defects.

An alternative model would be a negative feedback circuit in which the acquisition of all necessary crossover-intermediates triggers inactivation of DSB formation. According to this view, the presence of crossover precursors generates a signal to exit the DSB permissive state, rather than the absence of precursors extending this period. Such a model would require a ‘counting’ mechanism that enables exit from the DSB permissive state in response to a threshold number of crossover precursors. This seems less likely based on first principles, and also less consistent with our data.

Our observations also suggest that there is a minimum duration of proficiency for DSB formation that does not depend on how rapidly chromosome pairs attain crossover precursors. We would expect meiotic nuclei to achieve crossover precursors on every chromosome in a stochastic manner. If DSB-1 were removed from chromosomes upon reaching this state, we would likely see a patchwork of DSB-1 positive and negative nuclei in the early pachytene region, but instead we observe homogenous staining in this region, and abrupt disappearance of DSB-1 within a narrow zone of the gonad. Additionally, in mutants that appear to be

Figure 11. Model: Nuclei remain in a DSB-permissive state until a crossover precursor has been attained on each chromosome pair. At the onset of meiotic prophase, DSB-1 and DSB-2 are targeted to chromosomes to mediate DSB formation. A standard duration of the resulting DSB-permissive state is usually sufficient to ensure establishment of crossover (CO) precursors on most chromosome pairs. However, chromosome pairs that fail to form a crossover precursor emit a signal that prolongs the DSB-permissive state within individual nuclei. Once all chromosome pairs within a nucleus attain at least one crossover precursor, DSB-1 and DSB-2 are removed from meiotic chromosomes and DSB formation is thereby inactivated. As nuclei approach the bend region of the gonad, an override signal acts to shut off DSB formation regardless of crossover status. Upon exit from the DSB-permissive state, nuclei progress to late pachytene and complete crossover formation.

doi:10.1371/journal.pgen.1003679.g011
defective in triggering the obligate crossover checkpoint, such as htp-3 and htp-1, a zone of DSB-1-positive nuclei similar in length to that in wild-type animals is observed. Together these observations suggest that there is a preset temporal window for DSB formation that can be extended in individual nuclei but not shortened.

The duration of the DSB-permissive state might be specified by an activity or signal that decays with time and/or distance after meiotic entry. We speculate that the disappearance of DSB-1 may reflect a drop below a threshold level of CHK-2 activity, decay of CHK-2-mediated phosphorylation of DSB-1 or other targets, and/or a rise in an opposing activity—e.g., a phosphatase. Any of these could be inhibited by the putative checkpoint mechanism that prolongs DSB-1 localization in response to impaired crossover formation.

The nature of the recombination intermediate that satisfies the requirement for a crossover precursor on all chromosomes remains unknown. We distinguish “crossover precursors” from “interhomolog recombination intermediates” because components that are specifically required for crossovers, including MSH-5, ZHP-3, and COSA-1 [38,39,66,83], are all required for timely disappearance of DSB-1 from chromosomes. However, cytological markers for crossovers, including foci of ZHP-3 and COSA-1, do not appear until the late pachytene region of the gonad [39,66], after DSB-1 and DSB-2 disappear from meiotic chromosomes [47]. Thus, it seems likely that crossover precursors, rather than mature crossovers, are sufficient to allow exit from the DSB-permissive state.

Genetic and cytological evidence indicate that nuclei eventually cease to make DSBs, even when crossovers fail to be made on one or more chromosomes. As nuclei approach the bend region of the gonad at the end of pachytene, an “override” signal appears to shut off DSB formation (Figure 11). Unlike in mammals, where the mature crossovers, are sufficient to allow exit from the DSB-permissive state.

The Relationship between the Crossover Assurance Mechanism and Meiotic Progression

Numerous studies have documented a phenomenon known as the “extended transition zone” in mutants with defects in homolog pairing and/or synapsis [30,43,53,68]. An extended transition zone has been defined as a longer region of the gonad containing nuclei with crescent-shaped DAPI-staining morphology, multiple patches of the nuclear envelope proteins SUN-1 and ZYG-12, and strong foci of the ZIM proteins [68,74,85]. An extended transition zone appears to be a response to asynapsed chromosomes [43,68]. Previous work from our lab showed that the extension of the transition zone in synapsis-defective animals such as htm-8 hermaphrodites produce both sperm and oocytes in roughly normal numbers even when homolog pairing, synapsis, and/or recombination are severely impaired.

Rapid Divergence among DSB-Promoting Proteins

Proteins with apparent homology to DSB-1 are restricted to the Caenorhabditis lineage. Even within Caenorhabditis, DSB-1, DSB-2 and their homologs are only weakly conserved. This reinforces abundant evidence from other organisms that apart from Spo11 itself and the Rad50-Mre11 complex, proteins that promote DSB formation diverge rapidly during evolution [17,18,86]. This might seem surprising given that meiotic DSB formation is an essential aspect of sexual reproduction in most eukaryotes. However, potent and acute evolutionary pressures act on meiosis. For example, the germline is the site of intense warfare between the host genome and selfish genetic elements, which may contribute to the rapid evolution of meiotic proteins. In addition, the genome-wide distribution of DSBs appears to underlie the strongly biased distribution of crossovers observed in many species [87,88], including C. elegans (C. V. Kotwalwala and AFD, unpublished). The nature of this biased distribution shows interesting variation among species [89,90]. Since crossover number and position have a direct impact on the fidelity of meiotic chromosome segregation, mechanisms governing DSB distribution have likely evolved in concert with changes in chromosome structure and the spindle apparatus to maintain reproductive fitness.

Several features of meiosis in C. elegans distinguish it from other organisms in which DSB-promoting factors have been identified. In particular, DSBs and early recombination steps contribute directly to homolog pairing and synapsis in many species, while in C. elegans homolog pairing and synapsis occur independently of DSBs. Additionally, C. elegans lacks Dmc1, Hop2, and Mnd1, which are thought to function together as an essential meiotic recombination module in most eukaryotes [91]. C. elegans also lacks the DSB proteins Mei4 and Rec114, which are conserved between budding yeast and mice [18]. A correlation between the absence of DMC1/Hop2/Mnd1 and Mei4/Rec114 has been noted in several other lineages, and has been suggested to reflect a functional link between the formation of DSBs and their subsequent repair [18]. Interestingly, Rec114, like DSB-1/2, has several potential target sites for ATM/ATR phosphorylation, and these are important for regulation of DSBs in budding yeast meiosis [14]. Thus, the DSB-1/2 family of proteins may play analogous roles to known mediators of DSB formation in other species, despite their lack of apparent sequence similarity.

Materials and Methods

C. elegans Mutations and Strains

All C. elegans strains were cultured under standard conditions at 20°C. The wild-type strain was N2 Bristol. The nonsense dib-1(ce11) allele was generated by EMS mutagenesis. The dsb-1 deletion allele (tm3054) was generated by the Japanese National BioResource for the Nematode. Both dsb-1 alleles were extensively outcrossed to wild-type (5-6x), and additionally outcrossed in a directed three-point cross to dpy-20 unc-30 to eliminate most linked mutations. Additional mutants analyzed in this study were: spo-11(me44, ok79), nre-11(ck179), rad-50(ck197), chk-2(me64), com-1(t1626), rad-51(ck8701), rad-54(ck1268), msh-5(me23), cosa-1(me13), zhp-3(gf61), htp-3(bn3655), htp-1(gk174), syp-1(me17), syp-
were: guinea pig anti-HTP-3 [75], rat anti-HIM-8 [53], rabbit anti-HTP-3 [75], rat anti-HIM-8 [53], and rabbit anti-DSB-2 [47].

Immunofluorescence was performed as previously described [93]. Briefly, hermaphrodites 24–28 hours post L4 were dissected in egg buffer containing sodium azide and 0.1% Tween 20, fixed for 3 min in 1% formaldehyde in the same buffer between a Histobond slide and coverslip, and frozen on dry ice. The coverslip was removed, and slides were transferred to methanol chilled to −20°C. After 1 min, slides were transferred to PBST (PBS containing 0.1% Tween 20), washed in two further changes of PBST, blocked with Roche blocking agent, and stained with primary antibodies in block for 2 hours at room temperature or overnight at 4°C. Slides were then washed with 3 changes of PBST and stained with secondary antibodies. Secondary antibodies labeled with Alexa 488, Cy3, or Cy5 were purchased from Invitrogen or Jackson ImmunoResearch. Following immunostaining, slides were washed, stained in 0.5 mg/ml DAPI, destained in PBST, and mounted in buffered glycerol-based mounting medium containing 4% n-propyl gallate as an autofading agent.

For quantification of DAPI-staining bodies in oocytes, animals were dissected, fixed, and DAPI-stained as described above, omitting the steps involving immunostaining.

FISH procedures have also been previously described in detail [93]. Probes used in this study included the 5S rDNA repeat [23] and a short repeat associated with the right end of the X chromosome [53].

All images were acquired using a DeltaVision RT microscope (Applied Precision) equipped with a 100× 1.40 oil-immersion objective (Olympus) or (for whole gonad images) a 60× 1.40 oil-immersion objective (Olympus). Image deconvolution and projections were performed with the softWoRx software package (Applied Precision). Image scaling, false coloring, and composite image assembly were performed with Adobe Photoshop. All micrographs presented in the figures are maximum-intensity projections of 3D data stacks.

Immunoblotting

Lysate from 50 young adult hermaphrodites, picked at 24 hours post L4, was used for each lane. Gel electrophoresis was performed using 4–12% Novex NuPage gels (Invitrogen). Proteins were transferred to PVDF membrane. Guinea pig DSB-1 antibodies and rabbit DSB-2 antibodies (see above) were used for immunoblotting, followed by detection with HRP-conjugated secondary antibodies and ECL Western Blotting Substrate (Pierce).

Irradiation Experiments

Young adult worms were irradiated with approximately 10 Gy (1000 rad) from a Cs-137 source. For each experiment, unirradiated controls were treated identically to irradiated animals, other than exposure to radiation. For quantification of DAPI-staining bodies at diakinesis, hermaphrodites were irradiated 4–5 hours post L4 and dissected 18 hours post irradiation. To assess progeny survival, animals were irradiated 4–5 hours post L4, eggs laid 20–30 hours post irradiation were quantified, and surviving progeny were quantified 3 days later. For quantification of DSB-1 localization, animals were irradiated 16 hours post L4 and dissected 8 hours post irradiation. For RAD-51 immunofluorescence, animals were irradiated 24 hours post L4 and dissected 1 hour post irradiation.

Whole Genome Sequencing of we11

1000 homozygous we11 animals were picked from an outcrossed, balanced strain. A genomic DNA library was prepared as described in the genomic DNA library protocol from Illumina.
Libraries were sequenced using 76-bp single-end Illumina sequencing. MAQGene [94] was used to identify mutations present in the unc-119 mutant strain.

**Germline Cosuppression**

A 2.1-kb region of genomic DNA including the dsb-1 coding sequence and promoter was amplified by PCR using the following primers: 5′-CCGCCGTCCGATACCGC-3′ and 5′-GTTGCGCTGGTGAGAAGACG-3′. 100 ng/μl of dsb-1 PCR product was combined with 50 ng/μl of one-119 rescuing plasmid pXM651 [95] and injected into unc-119 animals. Rescued non-Unc F1 animals were picked to individual plates and assayed for embryonic lethality and male progeny. F2 animals were dissected, stained, and observed to quantify the number of DAPI-staining bodies in oocytes at diakinesis.

**Quantitative RT-PCR**

12 young adult animals, 24 hours post L4, were used for each genotype. RNA was purified from animals and reverse transcribed into cDNA with the SuperScript kit from Invitrogen using poly-A primers. spo-11 mRNA levels were compared by real-time PCR analysis with SYBR Green (Kapa Biosystems). act-1 and htp-3 mRNA levels were used as normalization controls. Primers used were as follows: spo-11 (5′-TGAGCCCGGATCTGTAGAAT-3′, 5′-TAGCTTGTTCCTTGCTGGTG-3′), act-1 (5′-CCCCCATCAAACATGAAGTC-3′, 5′-TCTGTITGGAGGTTTGAGG-3′), and htp-3 (5′-CGAGTGTAGACGAGGCTATATC-3′, 5′-TGCAAAGATAAACCGACGGTG-3′).

**Supporting Information**

**Figure S1** Mutation of dsb-1 does not affect spo-11 expression. Real-time PCR was used to measure the levels of spo-11 mRNA in dsb-1 mutants and WT animals. RNA was purified from age-matched young adult hermaphrodites at 24 hours post L4. spo-11 mRNA levels were normalized either to (A) act-1 or (B) htp-3 mRNA levels, both of which gave similar results. (TIF)

**Figure S2** Amino acid alignment of DSB-1 homologs. Global alignment of DSB-1 homologs from C. elegans, C. briggsae, C. remanei, and C. japonica were identified in the genome of each species included here. Alignment was performed using Geneious Pro (Geneious align-ment, Blosum62, default settings). (TIF)

**Figure S3** Validation of DSB-1 antibody specificity. Immunofluorescence staining of DSB-1 in early pachytenne nuclei in dsb-1 mutants. Only faint nonspecific background staining is observed. Scale bar, 5 μm. (TIF)

**Figure S4** DSB-1 positive nuclei in the late pachytenne region display RAD-51 foci and regions of asynapsed chromosomes. (A) Immunofluorescence staining of DSB-1 and RAD-51 in nuclei from the late pachytenne region of the gonad. Nuclei positive for DSB-1 staining also show condensed, transition zone-like DAPI-staining morphology, and have abundant RAD-51 foci. (B) Immunofluorescence staining of DSB-1, HTP-3, and SYP-1 in nuclei from the late pachytenne region of the gonad. Nuclei positive for DSB-1 staining contain asynapsed chromosome regions (HTP-3 positive axes not associated with SYP-1). DSB-1 positive nuclei are outlined with a dotted line. (TIF)

**Figure S5** Extension of DSB-1 staining is correlated with the extension of RAD-51 staining in mutants that disrupt crossover formation. Composite projection image of a gonad from a him-8 hermaphrodite, showing DAPI and immunofluorescence staining for DSB-1 and RAD-51. The disappearance of DSB-1 coincides with the disappearance of RAD-51 foci. (TIF)

**Figure S6** Extension of the DSB-1 region in crossover-deficient mutants is not a consequence of apoptosis. Quantification of the zone of DSB-1 localization, showing the percent, by length, of the Lzp region positive for DSB-1 staining. The genotypes indicated along the x-axis are present either as single mutants in the wild-type col-4 background or as double mutants combined with col-4(n1162). Mutation of col-4 abrogates germine apoptosis, but does not markedly or consistently alter the extended zone of DSB-1 localization to chromosomes. Error bars indicate standard deviations. (TIF)

**Acknowledgments**

We thank the CGC for providing strains. We are grateful to the Mitani lab and the Japanese National BioResource Project for providing the bnc5034 allele, and to David King for synthesizing the HTP-3 peptide. We also thank Barbara Meyer, Doug Koshland, and members of the Dembarg lab for helpful discussions.

**Author Contributions**

Conceived and designed the experiments: ELS AFD. Performed the experiments: ELS SER. Analyzed the data: ELS AFD. Contributed reagents/materials/analysis tools: SR AMV JA. Wrote the paper: ELS AFD.

**References**

1. Page SL, Hawley RS (2003) Chromosome Choreography: The Meiotic Ballet. Science 301: 783–789. doi:10.1126/science.1099605.

2. Petronczki M, Siomos MF, Nasmyth K (2003) Un Me´Nage a` Quatre: The Crossover Backend. Trends Genet 19: 473–480. doi:10.1016/S0168-9525(03)00083-7.

3. Keeney S (2001) Mechanism and control of meiotic recombination initiation. Current Topics in Developmental Biology. Academic Press, Vol. Volume 52, pp. 1–53. Available: http://www.sciencedirect.com/science/article/pii/S0070215301520086. Accessed 29 September 2012.

4. Padmore R, Cao L, Kleckner N (1999) Temporal comparison of recombination and synaptonemal complex formation during meiosis in S. cerevisiae. Cell 96: 1239–1256.

5. Gervantes MD, Farah JA, Smith GR (2000) Meiotic DNA Doses Associated with Recombination in S. pombe. Molecular Cell 5: 883–886. doi:10.1016/S1097-7575(00)80197-7.

6. Henderson KA, Kee K, Maleki S, Santini PA, Keeney S (2006) Cyclin-dependent kinase directly regulates initiation of meiotic recombination. Cell 125: 1321–1332. doi:10.1016/j.cell.2006.04.039.

7. Sasanuma H, Hirota K, Fukuda T, Kakusuo N, Kuogo K, et al. (2008) Cdc7-dependent phosphorylation of Mre2 facilitates initiation of yeast meiotic recombination. Genes Dev 22: 398–410. doi:10.1101/gad.1626608.

8. Wan L, Niu H, Fletcher B, Zhang C, Shokat KM, et al. (2006) Cdc28-Chk5 (CDK-S) and Cdc7-Dbf4 (DDK) collaborate to initiate meiotic recombination in yeast. Genes Dev 20: 396–397. doi:10.1101/gad.1626408.

9. Borde V, Goldman AS, Lichten M (2008) Direct coupling between meiotic DNA replication and recombination initiation. Science 290: 869–870.

10. Murakami H, Borde V, Shihata T, Lichten M, Ohta K (2003) Correlation between premeiotic DNA replication and chromatin transition at yeast recombination initiation sites. Nucleic Acids Res 31: 4085–4090.

11. Lange J, Pan J, Cole F, Thelen MP, Jasim M, et al. (2011) ATM controls meiotic double-strand-break formation. Nature 479: 237–240. doi:10.1038/nature10508.

12. Joyce EF, Pedersen M, Tiong S, White-Brown SK, Paul A, et al. (2011) Drosophila ATM and ATR have distinct activities in the regulation of meiotic DNA damage and repair. J Cell Biol 195: 359–367. doi:10.1083/jcb.201104121.
36. Meneely PM, McGovern OL, Heinis FI, Yanowitz JL (2012) Crossover
32. Daniel K, Lange J, Hached K, Fu J, Anastassiadis K, et al. (2011) Meiotic
28. Couteau F, Zetka M (2005) HTP-1 coordinates synaptonemal complex assembly
24. Chin GM, Villeneuve AM (2001) C. elegans mre-11 is required for meiotic
22. De Muyt A, Pereira L, Vezon D, Chelysheva L, Gendrot G, et al. (2009) A High
20. Libby BJ, Reinholdt LG, Schimenti JC (2003) Positional cloning and
19. Liu H, Jang JK, Kato N, McKim KS (2002) mei-P22 Encodes a Chromosome-
14. Carballo JA, Panizza S, Serrentino ME, Johnson AL, Geymonat M, et al. (2013)
843. doi:10.1038/nature09429.
118: 439–452. doi:10.1016/j.cell.2004.07.026.
Dev 15: 1674–1687. doi:10.1101/gad.902601.
HORMAD1. Nat Cell Biol 13: 599–610. doi:10.1038/ncb2213.
distribution and frequency are regulated by him-5 in Caenorhabditis elegans.
Distribution and Frequency Are Regulated by RPD-1. Genetics 190: 1251–1266. doi:10.1534/genetics.111.137463.
Hodgkin J, Horvitz HR, Brenner S (1979) Nondisjunction Mutants of the Nematode Caenorhabditis Elegans. Genetics 91: 67–94.
38. Kelly KO, Dernburg AF, Stamford GM, Vileenme AM (2000) Caenorhabditis elegans msh-5 Is Required for Both Normal and Radiation-Induced Meiotic Crossing Over but Not for Completion of Meiosis. Genetics 156: 617–630.
37. Maleki S, Neale M, Arora C, Henderson K, Keeney S (2007) Interactions links DSB formation with homolog pairing and crossing over during C. elegans meiosis. Genes Dev 19: 2727–2743. doi:10.1101/gad.1338365.
53. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
35. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
34. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
33. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
32. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
31. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
30. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
29. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
28. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
27. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
26. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
25. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
24. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
23. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
22. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
21. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
20. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
19. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
18. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
17. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
16. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
15. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
14. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
13. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
12. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
11. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
10. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
9. Phillips CM, Wong C, Bhalla N, Carlton PM, Weiser P, et al. (2005) HIM-8 Associated Protein Required for the Initiation of Meiotic Recombination in Caenorhabditis elegans. Journal of Molecular Biology 351: 475–485.
78. Herman RK, Kari CK, Hartman PS (1982) Dominant X-chromosome nondisjunction mutants of Caenorhabditis elegans. Genetics 102: 579–600.

79. Schulte J, Redfield H (1951) Interchromosomal Effects on Crossing Over in Drosophila. Cold Spring Harb Symp Quant Biol 16: 175–197. doi:10.1101/ SQB.1951.016.01.015.

80. Woglar A, Daryabaeiji A, Adamo A, Habacher C, Machacek T, et al. (2013) Matefin/SUN-1 Phosphorylation Is Part of a Surveillance Mechanism to Coordinate Chromosome Synapsis and Recombination with Meiotic Progression and Chromosome Movement. PLoS Genet 9: e1003335. doi:10.1371/journal.pgen.1003335.

81. Saito TT, Mohideen F, Meyer K, Harper JW, Colaiacovo MP (2012) SLX-1 Is Required for Maintaining Genomic Integrity and Promoting Meiotic Nondisjunction in the Caenorhabditis elegans Germline. PLoS Genet 8: e1002888. doi:10.1371/journal.pgen.1002888.

82. Aravind L, Koonin EV (1998) The HORMA domain: a common structural determinant in mitotic checkpoints, chromosome synapsis and DNA repair. Trends in Biochemical Sciences 23: 284–286. doi:10.1016/S0968-9055(98)00127-2.

83. Bailis JM, Smith AV, Roeder GS (2000) Bypass of a meiotic checkpoint byinteraction between the chromosomal proteins Red1p and Hop1p. Mol Cell Biol 20: 6646–6658.

84. Phillips CM, Dernburg AF (2006) A Family of Zinc-Finger Proteins Is Required for Maintaining Genomic Integrity and Promoting Meiotic Noncrossover in the Caenorhabditis elegans Germline. PLoS Genet 2: 360–369. doi:10.1371/journal.pgen.0004238.

85. Penkner AM, Fridkin A, Gloggnitzer J, Baudrimont A, Machacek T, et al. (2009) Meiotic Chromosome Homology Search Involves Modifications of the Nuclear Envelope Protein Matefin/SUN-1. Cell 139: 920–933. doi:10.1016/j.cell.2009.10.045.

86. Penkner AM, Frickin A, Gloggnitzer J, Baudrimont A, Machacek T, et al. (2009) Meiotic Chromosome Homology Search Involves Modifications of the Nuclear Envelope Protein Matefin/SUN-1. Cell 139: 920–933. doi:10.1016/j.cell.2009.10.045.

87. Kauppi L, Jeffreys AJ, Kerney S (2004) Where the crossovers are: recombination distributions in mammals. Nat Rev Genet 5: 413–424. doi:10.1038/nrg1346.

88. Malik S-B, Pightling AW, Stefanik LM, Schurko AM, Logsdon JM Jr (2008) An expanded inventory of conserved meiotic genes provides evidence for sex in Trichomonas vaginalis. PLoS ONE 3: e2079. doi:10.1371/journal.pone.0002079.

89. Bigelow H, Doitsidou M, Sarin S, Hobert O (2009) MAQGene: software to facilitate C. elegans mutant genome sequence analysis. Nat Methods 6: 549. doi:10.1038/nmeth.f.260.

90. Larson-Bogue B, Westhorpe FG, Taylor SS (2012) The Spindle Assembly Checkpoint. Current Biology 22: R966–R980. doi:10.1016/j.cub.2012.10.006.

91. Malik S-B, Pightling AW, Stefanik LM, Schurko AM, Logsdon JM Jr (2008) An expanded inventory of conserved meiotic genes provides evidence for sex in Trichomonas vaginalis. PLoS ONE 3: e2079. doi:10.1371/journal.pone.0002079.

92. Harper NC, Rillo R, Jover-Gil S, Assaf ZJ, Bhalla N, et al. (2011) Pairing and Cell Cycle Arrest in C. elegans. Developmental Cell 21: 934–947. doi:10.1016/j.devcel.2011.12.002.

93. Phillips CM, McDonald KL, Dernburg AF (2009) Cytological analysis of meiosis and cell cycle arrest in C. elegans. Cold Spring Harb Symp Quant Biol 2009: 6646–6658.

94. Bigelow H, Doitsidou M, Sarin S, Hobert O (2009) MAQGene: software to facilitate C. elegans mutant genome sequence analysis. Nat Methods 6: 549. doi:10.1038/nmeth.f.260.

95. Woglar A, Daryabaeiji A, Adamo A, Habacher C, Machacek T, et al. (2013) Matefin/SUN-1 Phosphorylation Is Part of a Surveillance Mechanism to Coordinate Chromosome Synapsis and Recombination with Meiotic Progression and Chromosome Movement. PLoS Genet 9: e1003335. doi:10.1371/journal.pgen.1003335.

96. Daudet MA, Declercq F, Dujon B (2005) Comparative Genomics of Hemiascomycete Yeasts: Genes Involved in DNA Replication, Repair, and Recombination. Mol Biol Evol 22: 1051–1025. doi:10.1093/molbev/msi063.

97. Vázquez LM, Mohideen F, Meyer K, Harper JW, Colaiacovo MP (2012) SLX-1 Is Required for Maintaining Genomic Integrity and Promoting Meiotic Noncrossovers in the Caenorhabditis elegans Germline. PLoS Genet 8: e1002888. doi:10.1371/journal.pgen.1002888.

98. Lara-Gonzalez P, Westhorpe FG, Taylor SS (2012) The Spindle Assembly Checkpoint. Current Biology 22: R966–R980. doi:10.1016/j.cub.2012.10.006.

99. Lichten M, de Massy B (2011) The Impressionistic Landscape of Meiotic Recombination. Cell 147: 267–270. doi:10.1016/j.cell.2011.09.038.

100. Malik S-B, Piptilung AW, Stefanik LM, Schurko AM, Logsdon JM Jr (2008) An expanded inventory of conserved meiotic genes provides evidence for sex in Trichomonas vaginalis. PLoS ONE 3: e2079. doi:10.1371/journal.pone.0002079.

101. Harper NC, Rillo R, Jover-Gil S, Assaf ZJ, Bhalla N, et al. (2011) Pairing and Cell Cycle Arrest in C. elegans. Developmental Cell 21: 934–947. doi:10.1016/j.devcel.2011.12.002.

102. Phillips CM, McDonald KL, Dernburg AF (2009) Cytological analysis of meiosis and cell cycle arrest in C. elegans. Cold Spring Harb Symp Quant Biol 2009: 6646–6658.

103. Daudet MA, Declercq F, Dujon B (2005) Comparative Genomics of Hemiascomycete Yeasts: Genes Involved in DNA Replication, Repair, and Recombination. Mol Biol Evol 22: 1051–1025. doi:10.1093/molbev/msi063.

104. Vázquez LM, Mohideen F, Meyer K, Harper JW, Colaiacovo MP (2012) SLX-1 Is Required for Maintaining Genomic Integrity and Promoting Meiotic Noncrossovers in the Caenorhabditis elegans Germline. PLoS Genet 8: e1002888. doi:10.1371/journal.pgen.1002888.