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Heterologous synapsis in *C. elegans* is regulated by meiotic double-strand breaks and crossovers

Hanwenheng Liu^{1,2} • Spencer G. Gordon ¹ • Ofer Rog ¹ •

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Abstract

Alignment of the parental chromosomes during meiotic prophase is key to the formation of genetic exchanges, or crossovers, and consequently to the successful production of gametes. In almost all studied organisms, alignment involves synapsis: the assembly of a conserved inter-chromosomal interface called the synaptonemal complex (SC). While the SC usually synapses homologous sequences, it can assemble between heterologous sequences. However, little is known about the regulation of heterologous synapsis. Here, we study the dynamics of heterologous synapsis in the nematode *C. elegans*. We characterize two experimental scenarios: SC assembly onto a folded-back chromosome that cannot pair with its homologous partner; and synapsis of pseudo-homologs, a fusion chromosome partnering with an unfused chromosome half its size. We observed elevated levels of heterologous synapsis when the number of meiotic double-strand breaks or crossovers were reduced, indicating that the promiscuity of synapsis is regulated by break formation or repair. In addition, our data suggests the existence of both chromosome-specific and nucleus-wide regulation on heterologous synapsis.

Keywords C. elegans · Heterologous synapsis · Synaptonemal complex · Double-strand breaks · Crossover

Introduction

Meiosis maintains the karyotypic stability of a species while generating genetic diversity through exchanges, or crossovers, between the parental genomes. Crossover formation, however, is inherently mutagenic and could disrupt genome integrity unless it integrates local sequence homology with chromosome contiguity. Failed crossover regulation—such as crossovers between homologous sequences that are on different chromosomes—leads to aneuploid gametes, causing inviability, sterility, or congenital conditions in the offspring.

The synaptonemal complex (SC) is a conserved chromosomal interface that plays key roles in regulating crossovers and safeguarding a successful meiosis (Moses 1956, 1958; Fawcett 1956; Page and Hawley 2004; Rog and Dernburg

☐ Ofer Rog ofer.rog@utah.edu

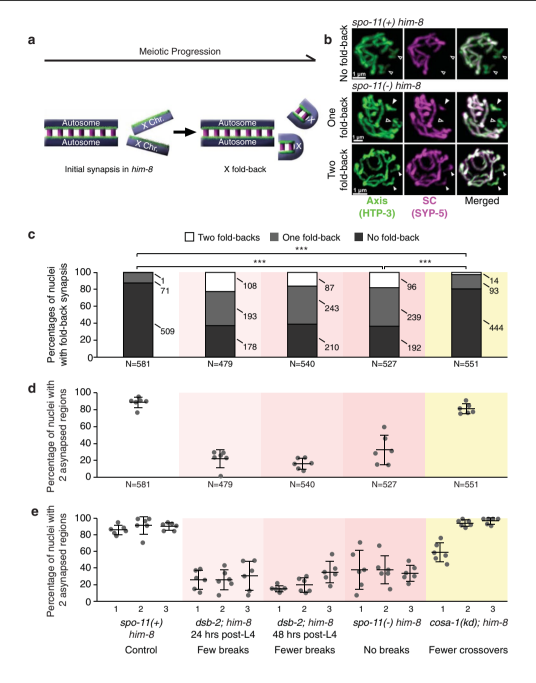
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- School of Biological Sciences and Center for Cell and Genome Sciences, University of Utah, 257 South 1400 East, Salt Lake City, UT 84112-0840, USA
- Present Address: The Division of Biology & Biomedical Sciences, Washington University in St. Louis, 660 South Euclid Avenue, Missouri 63110, USA

2013). The SC assembles between (or synapses) the parental homologous chromosomes (homologs) during meiotic prophase, bringing them into close juxtaposition and aligning them from end to end. In C. elegans, homologs are first brought into close proximity through specialized regions on each chromosome called Pairing Centers (Villeneuve 1994; MacQueen et al. 2005), which attach to the nuclear envelope and interact with cognate protein components (Phillips et al. 2005; Phillips and Dernburg 2006; Penkner et al. 2007; Sato et al. 2009). The SC starts assembling between the homologs at Pairing Centers (MacQueen et al. 2005; Rog and Dernburg 2015), stabilizing pairing interactions and extending them along the length of the chromosome (MacQueen et al. 2002). The physical tethering of the homologs helps ensure recombination occurs between homologous sequences on syntenic locations (Goldman and Lichten 2000).

The SC also ensures crossovers occur only between homologs by regulating crossover-promoting factors. Upon meiotic entry, many double-strand breaks (DSBs) are introduced (Keeney et al. 1997), a subset of which is repaired as crossovers. Crossover formation requires engagement with a homologous repair template, as well as recruitment of a suite of crossover-promoting factors to the





inter-homolog interface by the SC (Woglar and Villeneuve 2018; Li et al. 2018; Cahoon et al. 2019).

While the SC safeguards crossover formation, its mode of assembly also leaves meiosis vulnerable to illegitimate crossovers. The extension of synapsis along the chromosome is sequence-independent and relies on contiguity (MacQueen et al. 2005). This property allows the SC to traverse small regions of heterozygosity, such as insertions and deletions, while maintaining overall register between homologs and in that way allowing crossovers to form (Hammarlund et al. 2005). However, the capacity of the SC for heterologous synapsis carries the risk of bringing

together non-homologous chromosomes, thereby promoting illegitimate crossovers.

Two conditions that result in large-scale, cytologically discernable heterologous synapsis are fold-back synapsis and synaptic adjustment. Fold-back synapsis occurs when a chromosome that fails to find its partner heterologously synapses its left and right halves. While unpaired X chromosomes remain mostly asynapsed in hermaphrodites (Phillips et al. 2005), fold-back synapsis is prevalent upon deletion of all Pairing Center proteins (Harper et al. 2011), likely due to the abundance of unassembled SC subunits. SC associated with unpaired chromosomes was also documented in X chromosome triploids (Mlynarczyk-Evans et al. 2013), and



∢Fig. 1 DSBs and crossovers affect fold-back synapsis. a Schematic of the experimental systems to assess fold-back synapsis in standalone X chromosomes. Initially, only the autosomes synapse, while pairing and synapsis of the X chromosomes are prevented by the absence of HIM-8. As meiosis progresses, some of the unpaired X chromosomes fold back and self-synapse (in the case shown, both X chromosomes). **b** Co-immunofluorescence shows asynapsed regions. Top row: a nucleus from spo-11(+) him-8 lacking fold-back. Two asynapsed regions (axis staining unaccompanied by SC staining) can be seen, indicating no fold-back synapsis. Middle and bottom row: nuclei with no DSBs from spo-11(-) him-8 with one or no asynapsed region visible (middle and bottom rows, respectively), indicating fold-back synapsis of one or both unpaired X chromosomes. Green, HTP-3 (axis). Magenta, SYP-1 (SC). Hollow arrowheads, X chromosomes. White arrowheads, folded-back X chromosomes. Scale bars=1 µm. c Percentage of nuclei with two, one, or zero X chromosomes that underwent fold-back synapsis in each strain. Numbers of nuclei in each category are shown on the right of each stack (all pairwise comparisons p<0.0005, Pearson's chi-squared test). N represents total number of nuclei analyzed. Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers. d Percentage of nuclei with two asynapsed regions, indicating fold-back synapsis on both unpaired X chromosomes. Number of nuclei with fold-back synapsis increases sharply upon perturbation of DSBs, but only slightly when crossover level is lowered (p=0.05764, Student's t-test). N represents the number of nuclei from the 6 gonads assessed for each genotype. Each dot represents one gonad. Bars show mean \pm standard deviation (SD). Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers. e Dynamics of fold-back synapsis are affected by the levels of DSBs and crossovers. Perturbations of DSBs raised the amount of fold-back synapsis but did not reveal significant trends throughout pachytene. The numbers 1, 2, and 3 refer to sub-stages of pachytene (see Fig S1). Bars show mean \pm SD. N=6 gonads for each bin. Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers.

in as much as a third of nuclei in *C. elegans* males, which harbor a single, partner-less X chromosome (Checchi et al. 2014). Chromosomes can also undergo synaptic adjustment: a post-SC-assembly response to heterozygous translocations, inversions, deletions, insertions, or fusions, which has been documented in many eukaryotes (Moses and Poorman 1981; Moses et al. 1982; Zickler and Kleckner 1999). Following initial synapsis based on local sequence homology, asynapsed overhangs, junctions, and loops are minimized at the expense of heterologous synapsis.

Despite the prevalence of heterologous synapsis, and its potentially deleterious consequences, its regulation is poorly understood. One likely mode of regulation is modulation of SC dynamics. Contrary to what its highly ordered appearance in electron micrographs might suggest (Colaiácovo et al. 2003; Page and Hawley 2004), the SC displays liquid-like properties (Voelkel-Meiman et al. 2012; Rog et al. 2017). SC fluidity is modulated in response to meiotic progression, and specifically to the repair of DSBs and formation of crossovers: upon crossover formation,

the SC transitions into a less dynamic state (Pattabiraman et al. 2017; Nadarajan et al. 2017). This effect occurs at least partly *in cis* and can differentiate homolog pairs with and without crossovers (Machovina et al. 2016). However, whether DSBs or crossovers regulate heterologous synapsis has so far not been addressed.

Here, we quantified heterologous synapsis in *C. elegans* worms carrying chromosomes that can undergo fold-back synapsis and synaptic adjustment. We tested the effect of genetic perturbations that alter the levels of DSBs or crossovers and found increased heterologous synapsis in these two conditions. We calculated the frequency of fold-back synapsis for unpaired chromosomes, and the effect of fusing an unpaired chromosome to paired homologs. Finally, by relying on the spatio-temporal organization of the *C. elegans* gonad, we document the dynamics of heterologous synapsis.

Results

Analyzing fold-back synapsis in C. elegans

To quantify the extent of fold-back synapsis, we used worms lacking HIM-8, a protein necessary for pairing, and subsequent synapsis, of the X chromosomes (Phillips et al. 2005). Consequently, the DSBs that are formed on the unpaired X chromosomes are repaired without forming crossovers.

We inferred fold-back synapsis from the absence of asynapsed X chromosomes. We labeled the chromosomal axis protein HTP-3 (Goodyer et al. 2008) and the central region of the SC (referred to throughout simply as "the SC") using antibodies against SYP-1 (MacQueen et al. 2002) or SYP-5 (Hurlock et al. 2020; Zhang et al. 2020). Initial synapsis yields overlapping axis and SC staining for the autosomes, while the two X chromosomes are easily distinguished as asynapsed: axial structures that lack SC staining (Fig 1a, left and Fig 1b, top). If either or both X chromosomes are folded back and self-synapsed, we would see one or zero asynapsed regions, respectively (Fig 1b, middle and bottom). The progression of fold-back synapsis was easy to assess in the C. elegans gonad due to its spatial-temporal organization. We quantitated progression by dividing the pachytene region of the gonad, when crossovers mature, into three bins of equal length, roughly spanning early-, mid-, and late-pachytene (Fig S1; see "Methods"; Schedl 1997).

We first analyzed fold-back synapsis in *spo-11::aid him-8* animals grown on NGM plates (referred to throughout as *spo-11(+)*), which do not perturb the formation of DSBs and crossovers (Zhang et al. 2018; Almanzar et al. 2021) and therefore serve as our control. Consistent with previous analyses (Phillips et al. 2005), without HIM-8, only a minimal number of X chromosomes were synapsed: 71 of 581 nuclei contained one folded-back X chromosome and



only one of 581 nuclei had both X chromosomes folded-back (Fig 1c).

DSBs inhibit fold-back synapsis

DSB and crossover formation are tightly regulated by multiple surveillance mechanisms that impact chromatin and the SC in both chromosome-specific and nuclear-wide fashion (Rosu et al. 2013; Stamper et al. 2013; Machovina et al. 2016; Pattabiraman et al. 2017). We wondered whether the same surveillance mechanisms might also regulate heterologous synapsis.

We analyzed the effects of DSBs on fold-back synapsis by comparing three conditions with progressively reduced DSB number. Worms lacking the DSB-promoting factor DSB-2 exhibit progressive reduction in DSB number with age (Rosu et al. 2013), resulting in an average of 0.92 and 0.31 DSBs per chromosome pair in worms 24 and 48 h post-L4 stage, respectively (see "Methods" for calculations). These numbers are dramatically lower than the estimated 2-5 DSBs per homolog pair in wild-type animals (Colaiácovo et al. 2003; Mets and Meyer 2009; Rosu et al. 2013; Woglar and Villeneuve 2018) and entail that many chromosomes in dsb-2 animals do not receive any DSBs. To eliminate DSBs altogether, we degraded SPO-11, which catalyzes meiotic DSBs (Keeney et al. 1997; Dernburg et al. 1998), using the auxin-degradable allele *spo-11::aid* (Zhang et al. 2015, 2018; Almanzar et al. 2021). When raised on auxin plates, spo-11::aid him-8 worms lacked DSBs and crossovers altogether and are referred to hereon as spo-11(-).

dsb-2; him-8 worms (hereafter dsb-2) at 24 h post-L4 exhibited an increase of nuclei with one or zero asynapsed regions compared to spo-11(+) controls (Fig 1c, one foldedback chromosomes: 193 out of 479; two folded-back chromosomes: 178 out of 479; versus spo-11(+): p<0.0005, Pearson's chi-squared test). Fold-back synapsis was also elevated in dsb-2, 48 h post-L4 (Fig 1c; 243 one-fold-back and 87 two-fold-back chromosomes out 540 total) and in spo-11(-) (Fig 1c; 239 one-fold-back and 96 two-fold-backs out of 527). While the increased frequency of synapsis upon DSB perturbation could have potentially been caused by homologous synapsis of the X chromosomes, we find this unlikely. First, we observe many nuclei with only one asynapsed region (Fig 1c). Second, closer examination of nuclei with no asynapsed regions found most of them to have 7 rather than 6 SC threads (29/32 nuclei with zero asynapsed regions had 7 SC threads in *spo-11(-) him-8* worms). Since C. elegans has 6 chromosome pairs, this observation is consistent with two fold-back synapsis events.

Taken together, we find that DSBs are crucial for suppressing fold-back synapsis. Furthermore, the similar increase of fold-back synapsis in spo-11(-) (no breaks) and dsb-2, 24 h (0.46 DSBs per X chromosome) suggests that the

mechanism underlying the suppression of fold-back synapsis does not rely on the folded-back chromosome undergoing at least one DSB. If that were the case, we would have expected different fold-back frequencies between these two conditions.

Crossovers inhibit fold-back synapsis

The absence of sequence homology between the left and right halves of the X chromosome would prevent formation of crossovers whether or not fold-back synapsis has occurred. However, an orchestrated nucleus-wide response to chromosomes lacking crossovers is thought to help ensure each chromosome receives a crossover (Rosu et al. 2013; Stamper et al. 2013; Machovina et al. 2016). Such a response will be activated in the mutant scenarios analyzed above since not enough DSBs are made to generate at least one crossover on each homolog pair. (This would take place in conjunction with the response to the unpaired X chromosomes (Harper et al. 2011).) Furthermore, the SC responds to crossover formation by locally expanding (Libuda et al. 2013; Woglar and Villeneuve 2018). We therefore examined the role of crossovers in regulating fold-back synapsis.

COSA-1 is essential for the maturation of DSBs to crossovers (Yokoo et al. 2012). We attached aid degron to the N terminus of cosa-1 using CRISPR/Cas9 (Fig S2; Zhang et al. 2015). Our initial construct—aid::cosa-1 with the plant TIR-1 E3 ligase driven by the sun-1 promotor (sun-*1p::tir-1*)—had no discernable effect on COSA-1 function in the absence of auxin, and 70% reduction in viable selfprogeny in the presence of auxin (p=0.0014, Welch's t-test, α =0.05, same below). This reduction in viable progeny did not reach the level of spo-11(-) (Fig S3a) or a cosa-1 null mutant (Yokoo et al. 2012), suggesting that the degradation of AID::COSA-1 was incomplete. Upon switching the TIR-1 promotor from *sun-1p* to *gld-1p* (Chen et al. 2020), we observed 87% reduction in viable self-progeny (Fig S3a, p=0.0475 versus sun-1p, p=0.0009 versus gld-1p on NGM, Welch's t-test), suggesting it was more efficient in degrading AID::COSA-1. We refer to these worms, when grown on auxin, as cosa-1(kd), for COSA-1 knockdown.

To quantify the number of crossovers in *cosa-1(kd)* worms, we quantified the number of joined homologs immediately prior to the first meiotic division (DAPI bodies). Wild-type animals form six DAPI bodies, one for each of the six homolog pairs. A failure to form crossovers results in 7–12 DAPI bodies. *cosa-1(kd)*; *him-8* worms exhibited an average of 9.0 DAPI bodies, similar to *zim-2 zim3 him-8* triple mutant, where only 2 of the 6 homologs (chromosomes I and II) pair and form crossovers (Fig S3b and S3c; p=0.3976, Student's t-test; Phillips and Dernburg 2006). cosa-1(kd) therefore reduces crossovers number so that only about 1/3 of the homolog pairs form crossovers.



We observed only a slight increase in fold-back synapsis in cosa-1(kd); him-8 (hereafter cosa-1(kd)) worms relative to spo-11(+) animals, with 93 out of 551 nuclei that contained only one folded-back X chromosome, and 14 out of 551 nuclei that contained two folded-back X chromosomes (Fig 1c; p=0.0001 versus spo-11(+), Pearson's chi-square test). Notably, cosa-1(kd) worms exhibited a dramatically different fold-back synapsis distribution from dsb-2, 24 h post-L4 (p<0.0005, Pearson's chi-squared test) despite similar levels of crossovers in these two conditions (3.0 and 3.6 crossovers per nucleus in cosa-1(kd) and dsb-2, 24 h, respectively; Fig S3c; Rosu et al. 2013). Since cosa-1(kd) causes a reduction of crossovers despite high levels of DSBs (Yokoo et al. 2012) while in dsb-2, both DSBs and crossovers are reduced (Rosu et al. 2013), the difference we observed suggests that it is DSBs and not crossovers that play a key role in suppressing fold-back synapsis.

To highlight the differences, we focused on the percentage of nuclei with two asynapsed regions, where no fold-back synapsis occurred (Fig 1d). This metric facilitated tracking of fold-back synapsis along meiotic progression. The percentage of nuclei with two asynapsed regions stayed at a high level in *spo-11(+)* throughout pachytene (Fig 1e; bin 1 to bin 3 averages, 85%, 91%, and 90%, respectively). These frequencies lowered by three quarters in both *dsb-2*, 24 and 48 h post-L4, and by two-thirds in *spo-11(-)* (Fig 1e). While most conditions did not reveal significant trends throughout pachytene, an exception was *cosa-1(kd)*, where fold-back synapsis was higher in early pachytene but decreased with meiotic progression (Fig 1e).

DSBs and crossovers affect heterologous synapsis between pseudo-homologs

We next tested another condition resulting in heterologous synapsis: a pair of pseudo-homologs of unequal sizes. We constructed worm strains carrying a single copy of a fusion chromosome, ypT27, in which the left end of chromosome V is joined to the right end of the X chromosome (Lowden et al. 2008). We also deleted him-8, which prevented pairing of the two X chromosomes. In animals heterozygous for the fusion chromosome, the pseudo-homologs initially pair and synapse through chromosome V homology, allowing for crossover formation on that chromosome; the fused X chromosome overhangs at the terminus, and the standalone X chromosome is left unpaired and asynapsed (Fig 2a, left). Subsequently, the pseudo-homologs can be brought into full synapsis through various forms of heterologous synapsis. These included synaptic adjustment (Fig 2a, right, first and second rows), which has been documented in male worms carrying a similar pair of pseudo-homologs (Henzel et al. 2011); folding back of the fused X chromosome (Fig 2a, right, third row); or a combination of the two (Fig 2a, right,

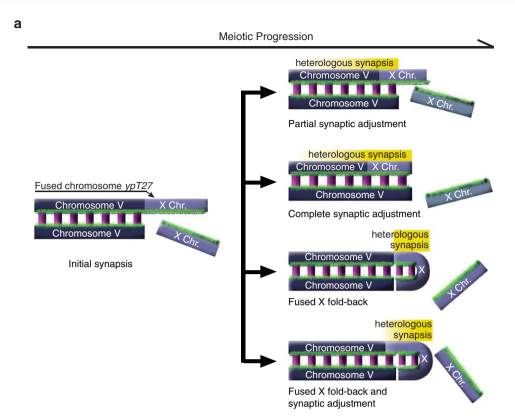
fourth row). We visualized heterologous synapsis in the pseudo-homologs as we did fold-back synapsis and inferred heterologous synapsis from the absence of asynapsed regions, which would occur upon complete heterologous synapsis of the pseudo-homologs or by complete fold-back synapsis of the standalone X chromosome (Fig 2b).

Once again, spo-11(+) served as our control. In a total of 778 nuclei we scored, a great majority—594 nuclei—contained two asynapsed regions: 165 and 19 nuclei had only one or zero asynapsed region, respectively (Fig 3a). This is in line with our fold-back synapsis data above, and with observations on a similar pair of pseudo-homologs in males (Henzel et al. 2011). This observation reiterated the notion that heterologous synapsis is uncommon when other meiotic processes are intact. Notably, however, we observed more heterologous synapsis in nuclei with the fusion chromosome compared to those with the two standalone chromosomes (Figs 1c and 3a; percentage of nuclei with two asynapsed regions, with:without fusion chromosome=73%:88%; p<0.0005, Pearson's chi-squared test). Since the difference between these two experimental scenarios is the fusion of X chromosome on the pseudo-homolog, our data suggest that fusion to a synapsed chromosome regulates heterologous synapsis.

As we experimentally diminished the number of DSBs and crossovers in ypT27/+; him-8 worms, asynapsis decreased. In dsb-2, 24 h post-L4, around half of the total nuclei (377 out of 762) contained two asynapsed regions, and around 1/3 contained one asynapsed region (298 nuclei). In dsb-2 animals at 48 h post-L4, with fewer DSBs, most X chromosomes underwent heterologous synapsis (440 and 148 nuclei out of 840 with one and zero asynapsed regions, respectively; Fig 3a), as was the case in spo-11(-), where no breaks form (Fig 3a; p=0.29, dsb-2, 48 h post-L4 versus spo-11(-), Pearson's chi-square test). Reduced crossovers in cosa-1(kd) also increased heterologous synapsis, with 434 out of 844 nuclei containing two asynapsed regions, and 285 and 125 nuclei bearing one or zero asynapsed regions, respectively (Fig 3a; p < 0.0005 versus spo-11(+), Pearson's chi-square test).

As above, we focused on the percentage of nuclei with two asynapsed regions to compare the different conditions. As pachytene progressed, the percentage of nuclei with two asynapsed regions increased in our *spo-11(+)* control, from a 66% average in bin 1 to 75% in bin 2 to 90% in bin 3 (Fig 3b), indicating a decrease in heterologous synapsis. This came as a surprising contrast with previous observations in *C. elegans* males (Henzel et al. 2011) and in other systems (Moses and Poorman 1981; Moses et al. 1982; Bojko 1990; Torgasheva et al. 2013), where heterologous synapsis increased with meiotic progression. Interestingly, the dynamics of heterologous synapsis differed between perturbations of DSBs and crossovers: while the trend of





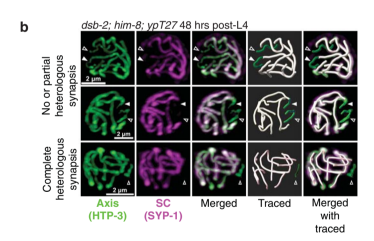


Fig. 2 Heterologous synapsis of pseudo-homologs. a Experimental system to detect outcomes of heterologous synapsis between a chromosome pair of unequal lengths. One of the chromosomes is a standalone chromosome V (dark blue), and the other is a fused chromosome of V and X (light blue). Left: initial synapsis between the fused chromosome and chromosome V leaves an X chromosome overhang. Right: as meiosis progresses, the two pseudo-homologs heterologously synapse. Partial synaptic adjustment (first row) leaves part of the overhang of the X chromosome. Complete synaptic adjustment (second row) means that the fused chromosome and the standalone chromosome V are adjusted to equal lengths. In both cases, much of the right side of chromosome V is synapsed with a heterologous part-

ner (yellow shading). Fold-back synapsis (third row) and a combination of fold-back synapsis and synaptic adjustment (fourth row) can also happen on the fused X chromosome. **b** Co-immunofluorescence shows asynapsed regions for heterologous synapsis in *ypT27+/-; dsb-2; him-8* worms, 48 h post-L4. Top and middle rows: two asynapsed regions are visible in each nucleus, indicating no or partial heterologous synapsis. Bottom row: only one asynapsed region is visible, indicating complete heterologous synapsis. Green, HTP-3 (axis). Magenta, SYP-1 (SC). White skeletons in traced images, synapsed region. Green skeletons in traced images, asynapsed regions. Hollow arrowheads, standalone X chromosome. Filled arrowheads, X chromosome overhang on the fused chromosome. Scale bars=2 μm.



heterologous synapsis in *cosa-1(kd)* mirrored that of the *spo-11(+)* control, it was reversed in the three conditions of perturbed DSBs, exhibiting increased heterologous synapsis with meiotic progression (Fig 3c). These opposite dynamics suggest that the underlying mechanism of heterologous synapsis suppression by DSBs and crossovers might not be the same.

Finally, to assess the precision of our methodology, we also analyzed asynapsed regions using maximum-intensity projections (2D analysis). This approach allows for a higher throughput while compromising precision. 2D analysis led to an apparent decrease in nuclei with two asynapsed regions in spo-11(+) (Fig 3d; two asynapsed regions, 2D versus 3D=49% versus 76%). This is likely because some asynapsed regions were obstructed by overlapping chromosomes. Nonetheless, when comparing spo-11(+) and spo-11(-) worms, we observed significantly fewer nuclei with two asynapsed regions in spo-11(-) worms (9%; p<0.005 compared to spo-11(+), Pearson's chi-squared test), recapitulating the trend we observed in the 3-dimensional analysis above.

Calculating heterologous synapsis frequencies for the standalone and fused X chromosomes

We used the distribution of nuclei with different numbers of asynapsed regions to estimate the frequency of complete fold-back synapsis for an unpaired X chromosome, $R_{\rm S}$ (for standalone; Fig 1c), and the frequency of complete heterologous synapsis for a fused X chromosome, $R_{\rm F}$ (for fused; Fig 3a). First, we fitted the fold-back synapsis dataset for each condition to a binomial distribution and solved for the best-fit R_S (Fig 4; see "Methods"). In most conditions, we observed a good fit to a binomial distribution (e.g., p=0.66029 in spo-11(+), Chi-squared goodness-of-fit test), consistent with the idea that, within each nucleus, the two asynapsed chromosomes undergo heterologous synapsis independently, each subjected to its own stochastic process. Subsequently, we calculated $R_{\rm F}$ for the pseudo-homologs: we used our calculated R_S —assuming it describes well the behavior of the standalone X—to solve for the best-fit $R_{\rm F}$ based on the distribution of asynapsed regions in the pseudohomologs datasets (Fig 4). These, too, exhibited mostly good fits to binomial distribution with two different rates, again consistent with independent stochastic processes.

With these frequencies in hand, we directly compared the effect of chromosome fusion on complete heterologous synapsis. In spo-11(+) controls, both $R_{\rm S}$ and $R_{\rm F}$ remained relatively low. However, the frequency of complete heterologous synapsis was threefold higher when fused to an autosome rather than standalone (Fig 4; average $R_{\rm S}=6.1\%$, $R_{\rm F}=19.9$; p=0.0007, Welch's t-test). The same trend was observed in cosa-1(kd), although both frequencies were correspondingly

higher (Fig 4; compared to spo-11(+), $R_S=9.7\%$, p=0.0987, Student's t-test; $R_F=38.3\%$; p<0.0005, Student's t-test). Interestingly, the trend reversed when DSBs were perturbed. The standalone X chromosome was more likely to engage in heterologous synapsis than the fused X chromosome. As DSBs were gradually depleted, the gap between R_S and R_F gradually diminishes (Fig 4).

Discussion

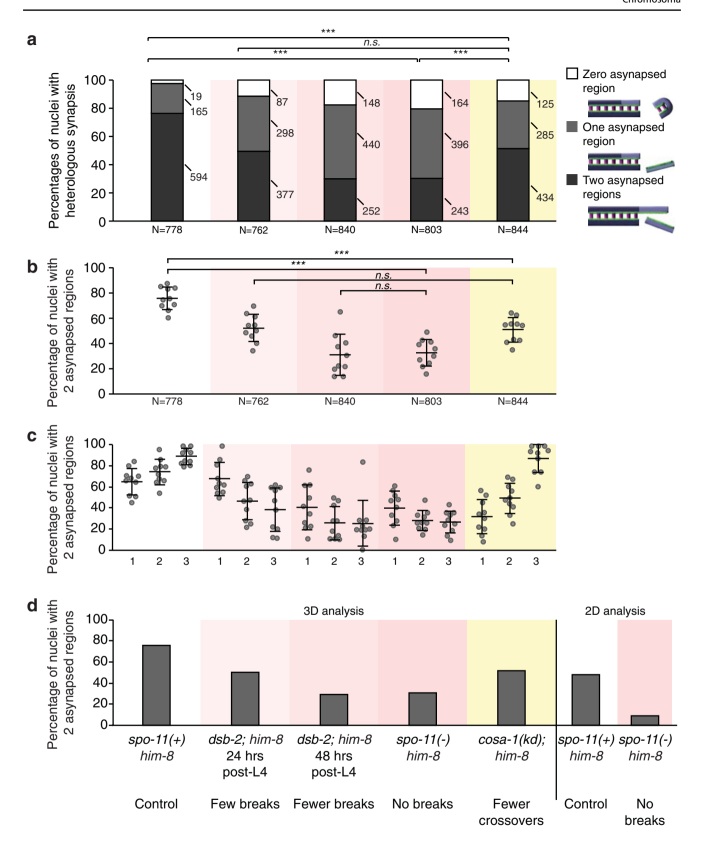
In this study, we perturbed the levels of DSBs and crossovers to examine their effect on heterologous synapsis. We found that heterologous synapsis is an uncommon event when DSB and crossover levels are unperturbed. However, decreased number of DSBs increased heterologous synapsis and also changed its dynamics, causing more heterologous synapsis as meiosis progressed. Decreasing crossover number also increased heterologous synapsis, but to a lesser extent and without affecting synapsis dynamics. Finally, by comparing worms with two different karyotypes, we were able to deduce how attachment to a synapsed chromosome affects heterologous synapsis.

Heterologous synapsis is uncommon

Successful meiosis maintains karyotypic stability by initiating synapsis using homology-based mechanisms. Across organisms, these include both DSB-dependent and -independent mechanisms (Zickler and Kleckner 1999). Worms utilize one of the best characterized DSB-independent mechanisms, where synapsis initiation relies on interactions between Pairing Centers (Villeneuve 1994; MacQueen et al. 2005), attachment to the nuclear envelope (Penkner et al. 2007), and force generation by dynein (Sato et al. 2009). Consistent with the robustness of these mechanisms, we find low levels of heterologous synapsis in worms lacking the X chromosome Pairing Center protein HIM-8, as long as DSBs and crossovers are unperturbed (Fig 1). In worms where the unpaired X chromosome was fused to a synapsed autosome, the frequency of heterologous synapsis was significantly higher but still represented a small subset of chromosomes (Figs 2 and 3). Since the SC helps recruit crossover-promoting factors (Woglar and Villeneuve 2018; Li et al. 2018; Cahoon et al. 2019), such low levels of heterologous synapsis will help limit illegitimate recombination: exchanges between homologous sequences that are not syntenic.

Interestingly, our data shows that heterologous synapsis does not increase with meiotic progression. We observed relatively constant levels of fold-back synapsis of the unpaired X chromosomes (Fig 1e) and decreasing levels of heterologous synapsis for worms with a fused X chromosome (Fig 3c). These findings contrast with earlier studies





of synaptic adjustment in *C. elegans* males (Henzel et al. 2011), and in other systems (Moses and Poorman 1981; Moses et al. 1982; Bojko 1990; Torgasheva et al. 2013),

which observed progressively more adjustment with meiotic progression. One potential explanation for the apparent discrepancy is that by scoring exposed axes, we might have



∢Fig. 3 DSBs and crossovers affect heterologous synapsis in pseudohomologs. a Percentage of nuclei with zero, one, or two asynapsed regions in each strain. Numbers of nuclei in each category are shown on the right of each stack. Nuclei with zero or one asynapsed region (implying complete heterologous synapsis) are rare in spo-11(+) him-8, but common when DSB and crossover levels are perturbed (all pairwise comparisons p < 0.0005, Pearson's chi-squared test). N represents total number of nuclei analyzed. Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers. b Heterologous synapsis is affected by the levels of DSBs and crossovers. Nuclei with two asynapsed regions decrease with lower levels of DSBs or crossovers (p<0.0005 for all comparisons with spo-11(+), Student's t-test). N represents total number of nuclei from the 10 gonads assessed for each genotype. Each dot represents one gonad. Bars show mean ± SD. All worms contain vpT27/+ pseudo-homologs. Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers. c Dynamics of heterologous synapsis are affected by the levels of DSBs and crossovers. Percentage of nuclei with two asynapsed regions, indicating partial or no heterologous synapsis, is shown. As DSBs are progressively eliminated, the levels of heterologous synapsis increase. The numbers 1, 2, and 3 refer to sub-stages of pachytene (see Fig S1). Bars show mean ± SD. N=10 gonads for each bin. All worms contain ypT27/+ pseudohomologs. Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers. d The effects of eliminating DSBs are robust, regardless of the method of scoring synaptic adjustment. Each bar represents total nuclei scored. Three-dimensional (3D) analysis (left) was performed by analyzing reconstructions of confocal z-stacks. Two-dimensional (2D) analysis (right) was performed on maximum-intensity orthogonal projections of independent datasets. N values for 3D analysis are the same as in Fig 3b. N=427 (spo-11(+)) and N=458 (spo-11(-)) for 2D analysis. All worms contain ypT27/+ pseudo-homologs. Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers.

conflated heterologous synapsis with desynapsis. Desynapsis occurs at the end of pachytene (MacQueen et al. 2002) and is particularly prevalent on chromosomes lacking crossovers (Machovina et al. 2016) or experiencing many DSBs (Couteau and Zetka 2011). We find this explanation unlikely since our region of interest in the gonads ended prior to regions of prevalent desynapsis. Also, the gradual decrease in heterologous synapsis started already in bin 2 (Figs 1e and 3c), arguing that late-pachytene events cannot explain our observations.

The propensity of the SC to synapse the chromosome from end to end, as well as the abovementioned studies of synaptic adjustment, have led to the suggestion that synapsed chromosomes represent a low energy state. However, our data suggests that with meiotic progression, the SC has a smaller propensity for heterologous synapsis, potentially representing a proofreading-like mechanism. Interestingly, association of SC proteins with the lone X chromosome in males has been observed specifically in early pachytene

(Checchi et al. 2014). The decreasing propensity for heterologous synapsis could be dependent on modulated biophysical properties of the SC, as previously suggested (Pattabiraman et al. 2017; Nadarajan et al. 2017). Finally, reduced heterologous synapsis in mid-pachytene—a meiotic stage where homologous synapsis is stable in *C. elegans*—suggests a role for sequence homology or a process that depends on homology (such as homologous recombination intermediates) in stabilizing the SC specifically between homologous chromosomes.

In mutants with reduced levels of DSBs, but not with reduced crossovers, the dynamics of heterologous synapsis for the fused X is reversed, with increased heterologous synapsis with meiotic progression (Fig 3c). This result points to a role for repair intermediates in regulating the SC. The similar dynamics between conditions with reduced DSBs (0.3–0.9 per chromosome) and complete lack of DSBs suggests that heterologous synapsis is not restricted by the presence of a DSB (or repair intermediate) *in cis*, i.e., on the chromosome that undergoes heterologous synapsis. But rather that the global level of DSBs, or perhaps a threshold number of DSBs per nucleus, transitions the SC to a restrictive state that limits heterologous synapsis.

Different effects of breaks and crossovers on heterologous synapsis

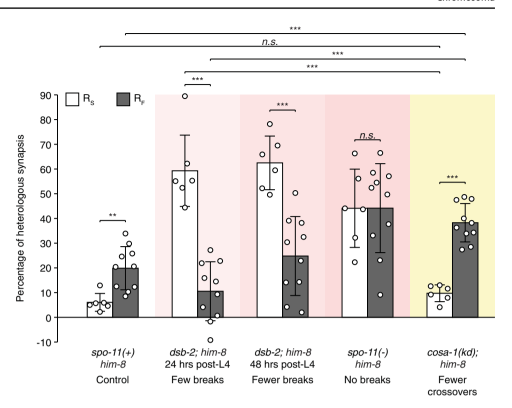
A major finding of our work is that almost all conditions where DSB or crossover levels are perturbed resulted in increased heterologous synapsis. In addition, comparing these conditions is telling.

While both *dsb*-2, 24 h post-L4 and *cosa-1(kd)* animals form a similar number of crossovers—2–3 per nucleus, or ~40% of chromosomes undergoing a crossover—only the former exhibited increased fold-back synapsis (Figs 1 and 4). Since the key difference between the two conditions is DSB level (0.9 per chromosome versus unperturbed levels, likely 2–5 DSBs per chromosome), our data suggests that DSBs or repair intermediates suppress fold-back synapsis. Strong effect of DSBs compared to crossovers on fold-back synapsis makes mechanistic sense as well, considering that the unpaired X chromosomes do not undergo crossovers (Phillips et al. 2005).

Crossovers also play a role in suppressing heterologous synapsis. This is particularly apparent on the fused X chromosome, where reduction of crossovers leads to doubling of $R_{\rm F}$ from 19 to 38%. While our methodology does not allow us to distinguish between different modes of heterologous synapsis—i.e., fold-back synapsis versus synaptic adjustment—an attractive possibility is that crossovers, by virtue of linking the homologs, impose a physical barrier that limits the compression of the chromosomes, thereby preventing complete synaptic adjustment. Consistent with



Fig. 4 Heterologous synapsis frequencies for standalone and fused X chromosomes. R_S and $R_{\rm E}$ in the different experimental conditions. In spo-11(+) and cosa-1(kd), $R_{\rm E}$ is larger than $R_{\rm S}$. Perturbation of DSBs causes generally larger R_S and R_F values but reverses the trend: minimally reducing DSB mostly affect $R_{\rm F}$, but the two values are equalized upon complete depletion of DSBs. Bars show average $R_{\rm F}$ and $R_{\rm S}$. N=6 gonads per condition for R_S . N=10 gonads per condition for $R_{\rm F}$, represented by dots. Error bars show mean ± SD. Relevant genotypes and their effects are indicated at the bottom. Increasingly darker pink shading indicates progressively perturbed DSBs; yellow shading indicates perturbed crossovers.



a *cis* model for suppression is the dose-dependent response of the fused X to reduction in DSBs (Fig 4), which entails a gradual decrease in the chance a crossover would form between the pseudo-homologs. This result is also consistent with the results of Henzel et al. (2011) in *C. elegans* males, where despite significant synaptic adjustment, complete adjustment of the pseudo-homologs is rarely achieved; and with the results of Torgasheva et al. (2013), which identified a role for crossovers in determining the extent of synaptic adjustment in mice.

Regulating heterologous synapsis through attachment to the nuclear envelope?

We document dramatic differences between heterologous synapsis on the standalone and the fused X chromosomes. This holds true upon perturbation of DSBs and crossovers, and also in worms where those pathways are not perturbed (Fig 4). Structurally, the two karyotypes we examined differ by a single phosphodiester bond between the telomeres of chromosomes X and V. However, this change could have multiple mechanistic implications on synapsis.

One possible explanation for the difference is the ability of the fused chromosome to undergo synaptic adjustment in addition to fold-back synapsis. This scenario implies that synaptic adjustment in worms can fully align the pseudohomologs, entailing axial compression or expansion by a factor of two. Alternatively, the difference in heterologous synapsis between the two karyotypes stems from increased propensity of the fused X chromosome to undergo fold-back synapsis. Since extension of synapsis is sequence-independent (MacQueen et al. 2005) and not rate-limiting for completion of synapsis (Rog and Dernburg 2015), it is likely that initiation of fold-back synapsis is impacted by fusion.

In worms, synapsis initiation between homologous chromosomes occurs at Pairing Centers (MacQueen et al. 2005; Rog and Dernburg 2015) and is regulated by local homology and by the attachment of chromosomes to force-generating machinery in the cytoplasm (Sato et al. 2009). Initiation of fold-back synapsis, as examined here, occurred in the absence of sequence homology or Pairing Center proteins. This unconventional initiation likely explains its delayed kinetics: it occurred much later in meiosis compared with synapsis initiation at homologous Pairing Centers (pachytene versus leptotene/zygotene; Fig S1 and MacQueen et al. 2002). The delayed kinetics likely reflects a higher energetic barrier that has to be overcome, and potentially a requirement for higher concentration of SC subunits (Harper et al. 2011), which accumulate with meiotic progression (Pattabiraman et al. 2017).

However, the standalone X chromosomes we analyzed lacked not only the ability to assess local homology at the synapsis initiation site, but also attachment to force-generating machinery in the cytoplasm (Penkner et al. 2007; Sato et al. 2009). In addition to facilitating homology search (Wynne et al. 2012), dynein-driven chromosomal



movements promote synapsis initiation (Sato et al. 2009). The same effect might also promote initiation of fold-back synapsis when the folded-back chromosome is fused to an autosome, as is the case for the pseudo-homologs. Pushing and pulling the chromosome may modify the axis—chemically or physically—which then primes it for association with the SC. A non-mutually exclusive hypothesis is that fold-back synapsis is promoted by the proximity to a synapsed chromosome, perhaps by increasing the local concentration of SC subunits.

Outlook

Heterologous synapsis is a prevalent feature of the meiotic program (Zickler and Kleckner 1999) and may be as widespread evolutionarily as the SC itself. While heterologous synapsis has the potential to promote illegitimate recombination, an important future avenue of research will be to characterize various kinds of illegitimate crossovers (e.g., León-Ortiz et al. 2018), and the degree by which heterologous synapsis promotes them. Recent work from our lab suggests another potential danger posed by heterologous synapsis: elevated meiotic sister-chromatid exchanges (Almanzar et al. 2021). Our finding of heterologous synapsis suppression by DSBs and crossovers opens the door to future studies of this potentially important pathway of preserving genome integrity during sexual reproduction.

Methods

Worm strains, growth conditions, and quantification of brood sizes

Worms were maintained according to standard protocols (Brenner 1974). All worm strains are listed in the Reagent List. Auxin plates contained a final concentration of 1 mM auxin (VWR Cat#AAA10556-36) prepared from a 500 mM stock in ethanol (Zhang et al. 2015). For auxin depletion, only animals that grew on auxin from hatching were analyzed. For analysis of the pseudo-homologs, males lacking the fusion chromosome were mated to hermaphrodites homozygous for the *ypT27* fusion (both animals had identical genotypes otherwise). F1 cross-progeny hermaphrodites were analyzed. For brood counts, four age-matched L4s were picked onto individual NGM or auxin plates and transferred onto new plates every 24 h for three consecutive days. Adult progeny was counted on all plates and summed.

CRISPR/Cas9 construction and injection

aid::cosa-1 repair template consisted of an aid degron in-frame with the open reading frame of cosa-1 through

a six-amino acid linker of glycines and serines, GSGSSG (Zhang et al. 2015). The PAM sequence CGG was mutated to CGC to prevent re-cutting. The template was ordered as two Ultramers from IDT, with a 35 bases overlap between them (Paix et al. 2016). Guide RNA was also ordered from IDT (Fig S2). The resulting strain was verified through PCR and Sanger sequencing (see Reagent List for primer pairs). CRISPR/Cas9 injections were carried out on Zeiss Axio Vert.A1 using Eppendorf FemtoJet 4i and Eppendorf TransferMan 4r.

Immunofluorescence staining

Worm gonad dissection and immunofluorescence staining were carried out following (Phillips et al. 2009) with a slightly modified version of egg buffer (25 mM HEPES pH 7.3, 118 mM NaCl, 48 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂; Zhang and Kuhn 2013). The primary antibodies used were goat anti-SYP-1 (gift of Abby Dernburg; MacQueen et al. 2002), rabbit anti-SYP-5 (gift of Yumi Kim; Hurlock et al. 2020), and guinea pig anti-HTP-3 (Hurlock et al. 2020). These were used at 1:400 dilution. Secondary antibodies include donkey anti-rabbit, donkey anti-guinea pig, and donkey anti-goat from Jackson ImmunoResearch, conjugated with either Alexa Fluor 647 or Alexa Fluor 488. These were used at 1:100 dilution.

Imaging and image analysis

Confocal images were acquired through Zen Black 2.3 on a Zeiss LSM 880 equipped with an Airyscan and a Zeiss Plan-APOCHROME 63x/1.4 NA oil objective. Z-stacks were taken with 0.159 μ m intervals. Pixel size was 0.0353 μ m. Images were Airyscan-processed in Zen 2.3 Blue and then quantified through 3D reconstructions in Imaris 9.5.1 or 9.7.2 (Oxford Instruments), where they were resampled to 0.0353 μ m intervals along the z axis.

For DAPI body quantifications, adult hermaphrodites that were hatched and grown on auxin were dissected and stained with anti-HTP-3 antibodies and DAPI. DAPI bodies in diakinesis nuclei were counted in *z*-stack 3D reconstructions using Imaris. HTP-3 was used to help distinguish between bivalents and univalents.

For analysis of fused X chromosomes, age-matched ypT27/+ worms (24 or 48 h post-L4) were grown on either NGM (for dsb-2 and spo-11(+)) or auxin plates (for spo-11(-) and cosa-1(kd)) at 20°C before dissection. After acquiring confocal z-stacks, maximum intensity projections were generated for each gonad. 2D analyses were carried out directly from the projections, scoring nuclei in pachytene. For 3D analyses, the projections were used to find the regions of interest (ROIs). ROI was defined as nuclei between the end of transition zone (based on DAPI staining;



MacQueen et al. 2002; Colaiácovo et al. 2003; Harper et al. 2011) and the start of diplotene (first row of nuclei with discernable SC disassembly). Each nucleus within the ROI was marked. Then, in Imaris, 3D reconstructions were resampled along the z axis from 0.159 to 0.0353 μ m to create a cubic voxel size, and if necessary, the channels were aligned. Chromosomes lacking an SC (HTP-3-positive and SYP-1-or SYP-5-negative) were scored and verified by tilting the nuclei. After scoring the nuclei, the ROI was divided into three bins of equal size based on the number of rows. Foldback quantification followed the same process, except lacking the ypT27 fusion.

Analysis of DSB perturbation in dsb-2 animals

From previously published results in Fig. 1D by Rosu et al. 2013, we calculated the average number of crossovers in any given nucleus at 24 and 48 h post-L4. The average number of DSBs was calculated from the number of crossovers using the equation in Yokoo et al. 2012.

Calculation of R_S and R_F

We calculated the frequencies of complete fold-back synapsis on the standalone X chromosomes (R_s) and complete heterologous synapsis on the fused chromosomes $(R_{\rm E})$ by finding the best-fit to a binomial distribution. Binomial distributions for probabilities with 0.001 increments of frequencies were calculated for each condition, i.e., nuclei with 0, 1, or 2 asynapsed regions. R_S corresponds to the frequency with the smallest sum of squares of difference to the observation. $R_{\rm F}$ was calculated by testing potential heterologous synapsis frequencies of the fused overhang that increased in increments of 0.001, with R_S for the other chromosome derived from the fold-back analysis above. Results were calculated against observed values, and $R_{\rm F}$ corresponds to the probability with the smallest sum of squares of differences. Averages of all gonads in each condition were used as the final R_S and R_F . All calculations were conducted in Excel (Microsoft Corporation) using BINOM.DIST (number, 2, probability, false) function.

Statistical analysis

Raw data were compiled in Excel (Microsoft Corporation). All statistical analyses were performed in PAST 4 (Hammer et al. 2001), with the exception of goodness-of-fit chi-square test for binomial distribution, which was performed manually in Excel.

The statistical significance value α =0.05 was used for all analyses. In all figures, significant p values were designated as follow: (*) for $0.05 \ge p > 0.005$, (**) for $0.005 \ge p > 0.0005$, and (***) for $p \le 0.0005$. Normality tests were performed

across all datasets prior to further statistical analysis. Datasets where the majority of data was normally distributed were analyzed through Student's *t*-test (equal variance) or Welch's *t*-test (unequal variance); Mann-Whitney *U* test was used otherwise.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00412-021-00763-y.

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Author contribution HL carried out most experimental work. SGG performed the 2D analysis (Fig 3d) and made the initial observation regarding *spo-11(-)* animals. HL and OR analyzed the data and wrote the manuscript, with input from all authors.

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Declarations

Conflict of interest The authors declare no competing interests.

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