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# The *Arabidopsis* Hop1 homolog ASY1 mediates cross-over assurance and interference

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## Abstract

The chromosome axis plays a crucial role in meiotic recombination. Here, we study the function of ASY1, the *Arabidopsis* homolog of the yeast chromosome axis-associated component Hop1. Specifically, we characterized cross-over (CO) distribution in female and male meiosis by deep sequencing of the progeny of an allelic series of *asy1* mutants. Combining data from nearly 1,000 individual plants, we find that reduced ASY1 functionality leads to genomic instability and sometimes drastic genomic rearrangements. We further observed that COs are less frequent and appear in more distal chromosomal regions in plants with no or reduced ASY1 functionality, consistent with previous analyses. However, our sequencing approach revealed that the reduction in CO number is not as dramatic as suggested by cytological analyses. Analysis of double mutants of *asy1* with mutants with three other CO factors, MUS81, MSH4, and MSH5, as well as the determination of foci number of the CO regulator MLH1 demonstrates that the majority of the COs in *asy1*, similar to the situation in the wildtype (WT), largely belong to the class I, which are subject to interference. However, these COs are redistributed in *asy1* mutants and typically appear much closer than in the WT. Hence, ASY1 plays a key role in CO interference that spaces COs along a chromosome. Conversely, since a large proportion of chromosomes do not receive any CO, we conclude that CO assurance, the process that ensures the obligatory assignment of one CO per chromosome, is also affected in *asy1* mutants.

## Significance statement:

The regulation of the number and placement of cross-overs (COs) during meiosis is critical to ensure meiotic fidelity and promote new genetic combinations. Here, we investigated the function of ASY1, which resides on the chromosome axis and plays a key role in CO formation. Our results show that COs in *asy1* mutants are positioned closer to each other than in the wildtype and that, despite a roughly similar number of COs, not every chromosome receives a CO. With this, our results shed light on the mechanisms regulating two important but still poorly understood aspects of meiosis: CO assurance, which safeguards at least one CO per chromosome pair, and CO interference, which prevents two COs from occurring close to each other.

## Introduction

Meiotic cross-overs (CO) are critical to the generation of genetic diversity. Through COs, segments of homologous chromosomes (homologs) are exchanged during the first meiotic division, creating new and unique combinations of alleles. Furthermore, COs are crucial for the equal segregation of chromosomes at the end of meiosis I since they physically link homologs and by that position them so that each cell pole will receive one homolog.

To generate COs, DNA double strand breaks (DSBs) are deliberately introduced into chromosomes in early prophase I. A fraction of these DSBs is then repaired by homologous recombination (HR) between the homologs, resulting in COs (1). HR relies on the resection of DSBs by the MRN complex generating 3'-overhanging single-stranded DNA, the search for sequence homology between

the homologs, and subsequent strand invasion mediated by the RecA-related recombinases radiation sensitive51 (RAD51) and disrupted meiotic cDNA (DMC1). Following DNA synthesis of the invading strand, and the capture of the second end of the initial DSB, a tetrahedral cross-strand intermediate is formed that is known as double Holliday junction (dHJ), which, when resolved, can result in a CO or a nonCO (NCO) (2, 3). NCOs can also be generated without the formation of dHJs (4).

COs are typically grouped into two classes, which can be distinguished by the molecular machinery that catalyzes the resolution of the dHJ. Type I COs rely on the ZMM group of proteins, including the DNA mismatch repair proteins mutl homologue1/3 (MLH1/3) (1). A hallmark of type I COs is that they experience positive interference, i.e., COs are more distantly spaced from each other than

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expected by chance (5). This results in a gamma-shaped distribution (with a shape distribution parameter larger than one) of COs across the genome. In contrast, type II COs are formed by a different pathway, which involves the endonuclease MMS4 AND UV SENSITIVE81 (MUS81) and are not subjected to interference (1).

Of key importance for meiotic recombination is the formation of the chromosome axis, by which a proteinaceous scaffold is formed along the DNA, and from which chromatin loops are thought to extrude (6). The chromosome axis plays a key role in DSB induction in many species and early steps of CO formation (7). Later in meiosis I, this axis becomes the lateral element in the synaptonemal complex (SC) that zips the two homologs together and promotes the maturation of COs in many species.

An important protein, which is associated with the chromosome axis, is Hop1 and its orthologs, such as ASYNAPTIC1 (ASY1) in *Arabidopsis* (8, 9). Earlier work has implicated ASY1/Hop1 in many central aspects of meiotic recombination, such as promoting interhomolog-biased DNA repair, chromosome pairing, SC formation, and CO production (8–14). In addition to these conserved functions, there are also species-specific functions of Hop1-type proteins for meiotic recombination, e.g., Hop1 is strictly required for DSB formation in yeast while DSBs are still formed in *asy1* mutants in *Arabidopsis* (10, 15).

The number and pattern of CO formation varies between species, and can be modulated by environmental conditions (16). The number of COs is also different between male and female meiosis. In *Arabidopsis*, there are usually 10 to 12 COs formed during male meiosis, but typically only 6 to 7 during female meiosis (17). The majority of COs, i.e., approximately 85%, are type I COs, and the 2 to 3 COs per chromosome are typically evenly distributed along the chromosome in male meiocytes (18). In contrast, a single CO is typically observed per chromosome in female meiocytes, and it is more often located in the pericentromeric regions (17). Importantly, all chromosomes typically undergo at least one CO in both sexes. This is insured through a mechanism referred to as CO assurance, which has been detected in many species (19–21).

How CO interference and assurance are brought about is currently not understood. In yeast, several mutants have been identified that affect CO interference, including topoisomerase II (TOPO II) (20). Whether these components function in a similar manner in other organisms remains unclear. One attractive model of how interference could be established is the “beam-film model” (22, 23). This model predicts that COs are formed at sites of high mechanical stress between the engaged homologs. Once a CO is formed, it locally releases this stress, and this relaxation spreads along the chromosomes laterally, preventing the formation of a CO nearby (22, 23). However, how this tension stress and its relaxation are transmitted is not clear. Recent studies suggest an alternative model proposing the diffusion-mediated coarsening of the proCO factor HEI10 (ZHP-3/4 in *Caenorhabditis elegans*), a RING finger-containing E3 ubiquitin ligase, that accounts for the CO positioning and interference (24, 25).

CO site selection appears to involve both epigenetic information, such as DNA methylation (26–29), and genetic factors such as the activity of the cell cycle kinase CYCLIN DEPENDENT KINASE A;1 (CDKA;1) (30, 31), as for instance shown in *Arabidopsis*. Interestingly, the number and position of COs have often been found to be modulated in polyploid plants, possibly to adapt to the challenge of faithfully assorting multiple chromosome sets during meiosis (32). Typically, polyploid species show a reduced CO num-

ber, and the remaining COs are often located at distal positions, a pattern which is also often found in diploid crop species such as maize (33). How this change in pattern is accomplished is not clear. It is also not clear whether allelic variants of the core meiotic machinery genes that appear to be associated with polyploidization, including alleles for ASY1, are functionally associated with this change in CO number and distribution (34).

Here, we have assessed the role of ASY1 in CO formation and distribution, using a genomics approach. To this end, we generated recombination maps of female and male meiosis for nearly 700 *asy1* mutant plants, and characterized their CO patterns in comparison to the WT. First, we observed many instances of full or partial chromosome aneuploidy in the progenies of the *asy1* mutants, confirming the importance of ASY1 for meiotic fidelity. Our work revealed that the overall number of COs is reduced in *asy1* mutants but this trend is less pronounced in the surviving progeny than previously estimated by cytological studies. We show that COs in *asy1* largely belong to the class I of COs and tend to closely localize at the telomeric and subtelomeric regions, highlighting a role for ASY1 in CO interference. Furthermore, the distribution of COs per chromosome is altered in plants with reduced ASY1 functionality and a large percentage of chromosomes being devoid of any CO, suggesting that ASY1 is a crucial factor for CO assurance as well.

## Methods

### Plant materials and growing conditions

The *A. thaliana* accessions Columbia (Col-0) and Landsberg *erecta* (*Ler-1*) were used as the WT references in this study. The *asy1* T-DNA insertion line (SALK\_046,272) in Col-0 background (35) and *mlh1-3* (SK25975) was obtained from the T-DNA mutant collection at the Salk Institute Genomics Analysis Laboratory (SIGNAL, <http://signal.salk.edu/cgi-bin/tdnaexpress>) via NASC (<http://arabidopsis.info/>). The hypomorphic non-phosphorylatable mutant *asy1*<sup>T142V</sup> in Col-0 background was generated previously in Yang et al. (36, 37). The *asy1* mutant in *Ler* background was generated in this study by CRISPR/Cas9 approach (Fig. S1). All plants were grown in growth chambers with a 16 h light/21°C and 8 h dark/18°C cycle at 60% humidity. The generation of genome sequencing populations was described in the main text. For this procedure, homozygous *asy1* mutants (*asy1*<sup>Col-0</sup>, *asy1*<sup>T142V</sup>, and *asy1*<sup>ler-1</sup>) were used.

### Genomic sequencing and raw read processing

For *Arabidopsis* genomic DNA extraction, about 2 cm<sup>2</sup> of leaf was harvested and used for DNA extraction using the SPRI beads-based method (38). Subsequently, genomic DNAs were quantified using a plate fluorometer with SYBR Green I, and normalized. After normalization, the samples were checked on agarose for uniformity. The generation of the sequencing libraries was performed following the Rowan protocol (39). Briefly, samples were fragmented, adapters were added, the fragments were amplified with indexed P5 and P7 primers, and visualization on agarose before being pooled (see Supplementary Material 1 for index and pooling information). Pools then underwent size selection with magnetic beads before measuring DNA concentration using a Qubit (Invitrogen) and verifying size distribution with the Agilent Bioanalyzer. Pools were sequenced on an Illumina HiSeq 4000 sequencer to obtain PE150 reads. Raw reads were checked for sequence quality, trimmed, and demultiplexed using custom Python scripts, as described previously, using `allprep-15.py` (40). After demultiplexing, each individual retained reads corresponding to 1.7x cover-

age of the haploid *A. thaliana* genome. Next, they were aligned to the TAIR10 genomic reference (<https://www.arabidopsis.org>) using BWA and default parameters (41), producing sam files (bwa-doall.py version 0.7.17) appropriate for dosage analysis (see below). For the purpose of genotyping and the creation of genetic maps, the data was also concatenated into a single mpileup.txt file using samtools (42). Finally, this mpileup file was further processed to obtain the percentages of each base call at each position and each sample using the script called mpileup-parser.py (beta-mpileup-parser\_vMEM-O.py, <https://github.com/Comai-Lab/mpileup-tools>).

## Dosage analysis and genotyping

Aneuploidy and dosage variation detection were performed as previously described (43). Briefly, the genome was divided into consecutive nonoverlapping 100 kb bins and reads that mapped to each bin were counted for each sample using a custom python script (bin-by-sam-v7.py, <https://github.com/Comai-Lab/bin-by-sam>). For each bin, read counts were normalized to account for variation in total read counts, as well as compared to the mean values for the whole population, in order to detect variation in dosage relative to the expected value of two in a diploid background. Data were plotted (Fig. 1 for an example), and aneuploidy was detected visually, as a variation in copy number that is consistent across at least three consecutive bins.

## Determination of CO locations and data analysis

A list of Col-0/Ler-1 SNPs was derived from a previously published list of ~870 K SNP positions, ([http://mtweb.cs.ucl.ac.uk/mus/www/19genomes/variants.SDI/ler\\_0.v7c.sdi](http://mtweb.cs.ucl.ac.uk/mus/www/19genomes/variants.SDI/ler_0.v7c.sdi)) (44). We first filtered this list by selecting positions that had been detected by both the IMR and DENOM software, that did not include any ambiguous bases (only A, T, G, and C were allowed), with phred scores >100, and for which the reported consensus base SNP and the high mapping quality consensus base SNP were in agreement. This reduced the SNP set to ~418 K positions. Next, we further filtered this list by only retaining positions that were covered at least once in at least 500/1108 of our samples and for which two both expected alleles were detected in our population (at least 5% of each allele type). The resulting ~125 K positions were used for genotyping. For each sample, the allele calls at each of these SNP positions were recorded. Because we sequenced each sample at low coverage, allele calls were pooled in order to obtain robust and complete genotype calls for each sample. Specifically, allele calls were pooled into a total of 239 consecutive nonoverlapping 500 kb bins, and a per-bin genotype was derived for each individual by calculating the mean percentage of Col-0 allele in each bin, as previously described (45). The resulting genotype information was input into R/qtl (version 1.42 to 8) for the creation of genetic maps for each cross (est.map function), and the detection of COs (countXO function) and the determination of their position (locateXO function) (46).

## CO interference

The testcross design described above was used to assess interference. A Python script was used to divide the genome into nonoverlapping intervals of arbitrary length. Recorded recombination breakpoints were counted, inferring COs per interval. The recombination frequency of any interval was equal to the (number of COs)/(number of scored chromatids). The predicted double COs number was determined by taking the square of the recombination rate. Double COs instances were scored based on COs

pairs. Ten instances of triple or higher number CO's were observed in the 4.5 Mb bin analysis, and scored as individual instances. For example, an interval with three COs would be scored as having two double COs, one between CO1 and CO2, the second between CO2 and CO3. These data are summarized in Supplementary Material 2.

## Cytogenetic analyses

Meiotic chromosome spreads and fluorescence in situ hybridization (FISH) were performed according to procedures previously described (47). The following DNA probes were used: pTa71 [45S ribosomal DNA (rDNA), pTa71 of *Triticum aestivum*; (48)] and pCT4.2 [5S 133 rDNA; (49)].

Immunolocalization procedures were conducted following the method detailed in Armstrong et al. (50), with modifications detailed in Varas and Pradillo (51). The primary antibodies used were kindly provided by Professor Chris Franklin (University of Birmingham): anti-AtASY1 (rat; 1/1000), anti-AtZYP1 (rat; 1/500), anti-MLH1 (rabbit, 1/500), anti-RAD51 (rabbit, 1/300). Secondary antibodies were anti-rat and anti-rabbit IgG FITC conjugated (Agrisera) and anti-rabbit and anti-rat IgG Alexa Fluor 555 conjugated (Molecular Probes).

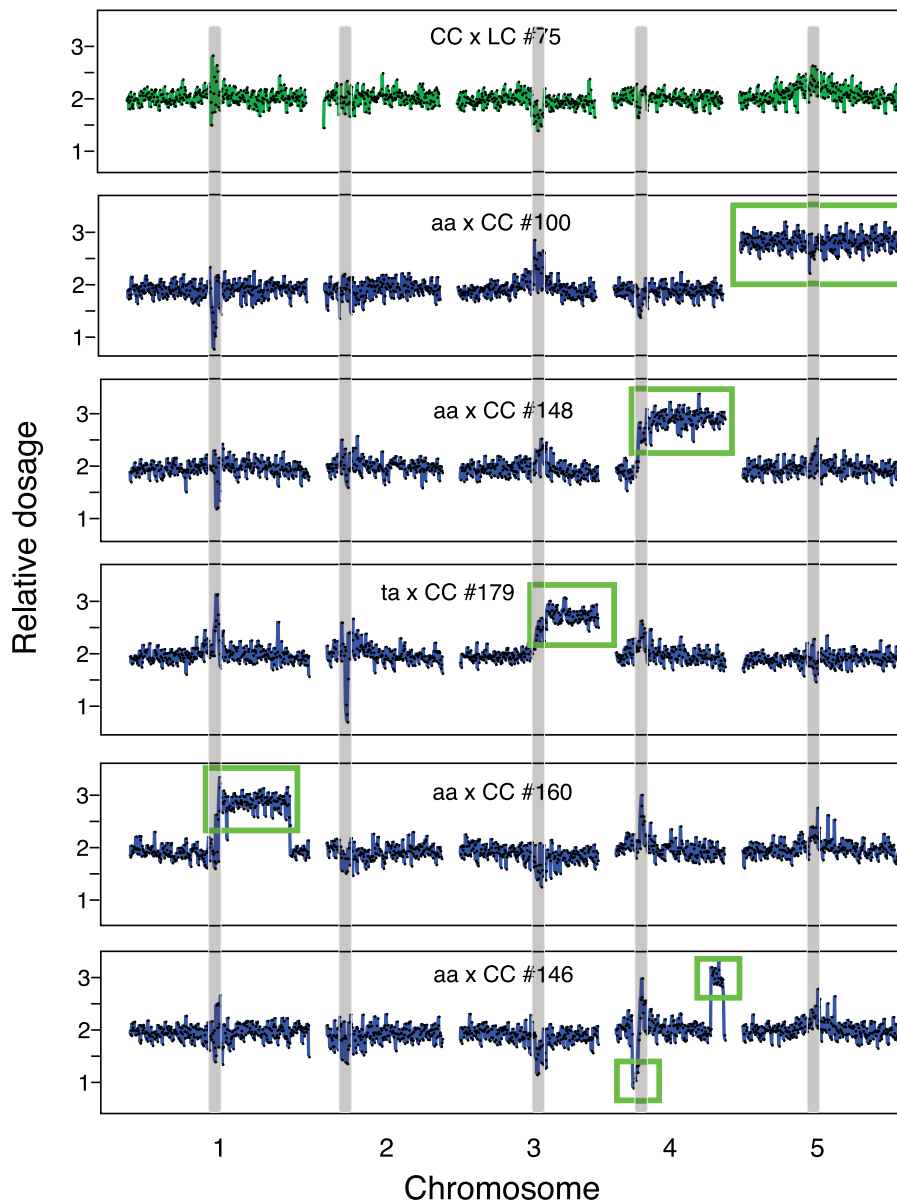
Slides were examined on an Olympus BX61 epifluorescence microscope and imaged with a CCD Olympus DP71 camera and computer using analySIS software (Soft Imaging System). For foci scoring, blind-coded digital images were used. Adobe Photoshop (CC 2018 v.19.0) and ImageJ (version 1.51 s) were applied for image processing.

## Results

### Creation of mapping populations to study ASY1 function in meiotic recombination

To study the effect of ASY1 on meiotic recombination, we sought to dissect CO distribution through a backcross design of the two different *Arabidopsis thaliana* accessions Columbia-0 and Landsberg erecta (*Ler-1*). Therefore, we first generated an *asy1* null mutant using a CRISPR/Cas9 approach in the background of the accession Landsberg erecta (*Ler-1*) (52). Cas9 was targeted to the ASY1 coding region at the first exon-intron junction by a single guide RNA harboring 20 bases (Fig. S1A). A mutant allele (*asy1<sup>Ler-1</sup>*) harboring a guanine (G) insertion at the beginning of the second exon of ASY1 was subsequently identified from the T2 generation that did not harbor the CRISPR/Cas9 T-DNA anymore. This G insertion results in a frameshift and premature stop codon after the translation of the first 13 amino acids. Homozygous *asy1<sup>Ler-1</sup>* mutants showed high levels of pollen mortality, and a high percentage of aborted seeds, matching the mutant phenotype of the previously identified *asy1* null mutant in the Columbia-0 accession (*asy1<sup>Col-0</sup>*) (Fig. S1B to E) (12, 53). That this allele represents a loss-of-function mutant was confirmed by the absence of ASY1 from male meiotic cells by immuno-detection (Fig. S1F). A complementation test using the previously generated functional ASY1 reporter (*PRO<sub>ASY1</sub>: ASY1: GFP*) (36) showed that the fertility defects of *asy1<sup>Ler-1</sup>* mutants were fully rescued by this reporter, corroborating that the meiotic defects of this CRISPR/Cas9-induced allele are due to the loss of ASY1 function (Fig. S1B to E).

Next, the *asy1<sup>Ler-1</sup>* mutant was combined with either an *asy1<sup>Col-0</sup>* null mutant, or a previously generated hypomorphic *asy1* mutant, called *asy1<sup>T142V</sup>* (both in the Col-0 genetic background), in which the threonine residue of the presumptive CDKA;1 phosphorylation site at position 142 is replaced by the nonphosphorylatable amino acid valine (36). *In vitro*, the nonphosphorylatable mutant



**Fig. 1.** Aneuploidy in the progeny of the *asy1* mutants. Dosage variation in six representative individuals from different crosses (CC x LC #75, aa x CC #100, aa x CC #148, ta x CC #179, aa x CC #160, and aa x CC #146). Each panel represents one individual and, for each individual, the five chromosomes are shown. For each individual, relative dosage, e.g.,  $\sim 2$  for normal diploids, is depicted based on short-read sequence coverage variation after being normalized to the whole population. Dots represent consecutive nonoverlapping 100 kb bins. None of the progeny of wild-type control crosses (CC x LC and LC x CC) exhibited aneuploidy (top panel, shown in green). Crosses involving *asy1* or *asy1*<sup>T142V</sup> mutants produce many aneuploid progeny (panel 2 to 6, colored in blue), including full trisomics (panel 2, aa x LC #100), partial trisomics (3rd panel, aa x CC #148), or more complex aneuploids (panels 5 and 6, aa x CC #160 and aa x CC #146). Many of the progeny of the crosses involving the hypomorphic *asy1*<sup>T142V</sup> mutant carried an extra copy of the bottom arm of chromosome 3 (Chr. 3B) (Table S2), suggesting that it originated from the parental line (panel 4, ta x CC #179). The regions that are present in additional or missing copies are indicated by the green rectangles. Centromeric and pericentromeric regions (gray vertical bands) consistently exhibited variation that mimicked dosage variation, but more likely originated from allelic variation between Col-0 and Ler-1 in these repeat-rich regions. These variations were consistently observed in all populations and were not recorded as aneuploidy.

*asy1*<sup>T142V</sup> has reduced binding affinity to ASY3, a key component of the chromosome axis, and shows decreased chromosome association in vivo (36, 54). These combinations then resulted in the following Col-Ler F1 hybrids: the first carries two null alleles of ASY1 (referred to as aa from now on), and the second carries a null allele in Ler-1 and the hypomorphic allele in Col-0 (*asy1*<sup>T142V-Col-0</sup> *asy1*<sup>Ler-1</sup> referred to as ta from now on).

Finally, we produced six mapping populations by backcrossing these hybrids to WT Col-0 plants (Table S1): specifically, the hybrids with reduced ASY1 functionality were crossed as female par-

ents to the Col-0 WT (referred to as CC from now on) to assess the importance of ASY1 in female meiosis. These crosses are designated in the following as aa x CC ( $n = 207$ ) for crosses with the *asy1* null mutant and ta x CC ( $n = 213$ ) for crosses with the hypomorphic mutant *asy1*<sup>T142V</sup>. Reciprocal crosses were performed to monitor male meiosis, designated as CC x aa ( $n = 216$ ) for crosses with the *asy1* null mutant and CC x ta ( $n = 214$ ) for crosses with the hypomorphic mutant *asy1*<sup>T142V</sup>. As a control, we also generated F1 hybrids between the WT parental lines Col-0 and Ler-1, and crossed them reciprocally with a Col-0 tester line giving rise



to two additional populations, named LC x CC ( $n = 117$ ) and CC x LC ( $n = 119$ ). To characterize recombination in *asy1* mutants, the progeny of these six populations were subjected to whole-genome sequencing.

## Reduction of ASY1 functionality results in aneuploid offspring and genome instability

Low recombination frequencies often cause the formation of univalents and hence unequal chromosome distributions and aneuploidy in the resulting spores (11). Previous cytological data (12, 10, 53, 54) suggested that CO numbers might be reduced in *asy1* mutants since not all homologs are able to form bivalents. Consistent with these earlier studies, we also found univalents in meiotic chromosome spread analyses of male meiocytes produced by *ta* plants (Fig. S2A). To investigate the effect of these univalents on the karyotype of the resulting progenies, we screened each of these populations for the presence of aneuploidy. Looking for variation in sequencing read coverage across the whole genome, we found aneuploid plants in the progenies of all four crosses involving *asy1* mutants, while none were found in the control WT crosses (Table S2 and Figs. 1 and 2).

Most of the aneuploids recovered were trisomics, carrying one additional copy of a single chromosome (Fig. 1, second panel). Trisomies of all chromosome types were recovered, except for chromosome 1 (Table S2), perhaps because of more severe selection against gametes carrying additional copies of this particular chromosome, as observed previously (55, 56), or because of the more severe phenotypic consequences associated with additional copies of chromosome 1 in *A. thaliana* (57). Conversely, we did not detect any instance of a missing chromosome, probably because it is lethal or severely detrimental at the gametic stage, consistent with previous work (58–60).

Interestingly, we also found examples of segmental aneuploids, carrying only a part of an additional chromosome (Fig. 1, panels 3 to 6 and Table S2). The karyotypes of some of segmental aneuploids could have originated from a single miss-repaired DNA DSB (Fig. 1, panels 3, 4), in which case only a terminal segment remains. In addition, we find individuals carrying signs of two or more breaks (Fig. 1, panels 5, 6), resulting in more drastically rearranged genomes. Comparing the approximate location of these breaks with the expected position of the centromeres (the *Arabidopsis* genome initiative), suggested that more than half (9/17) of these miss-repaired breaks occurred at or around the centromeres (Table S2, second column). While this pattern possibly suggests a direct or indirect role of ASY1 in the formation and/or repair of DSB near centromeres, breaks at centromeres are also often enriched after gamma radiation (59), in the progeny of aneuploids (56), and after genome elimination (58), suggesting that centromeric regions are in particular sensitive to DNA stresses.

Most of the individuals exhibiting segmental aneuploidy originated from crosses involving the *asy1* mutants as a female (Table S2). This overrepresentation in females is consistent with the idea that segmental aneuploids are less fit than euploids and hence are counter-selected during pollen tube growth. Moreover, segmental aneuploids appeared much more frequently in crosses involving *asy1* null mutants, as we only found one case originating from a cross with the hypomorphic mutant (Table S2), suggesting that genomic instability scales with the level of ASY1 reduction.

Strikingly, one individual (#102) from the *aa* x CC cross exhibited signs of high levels of genomic instability (Fig. S3). Plant #102

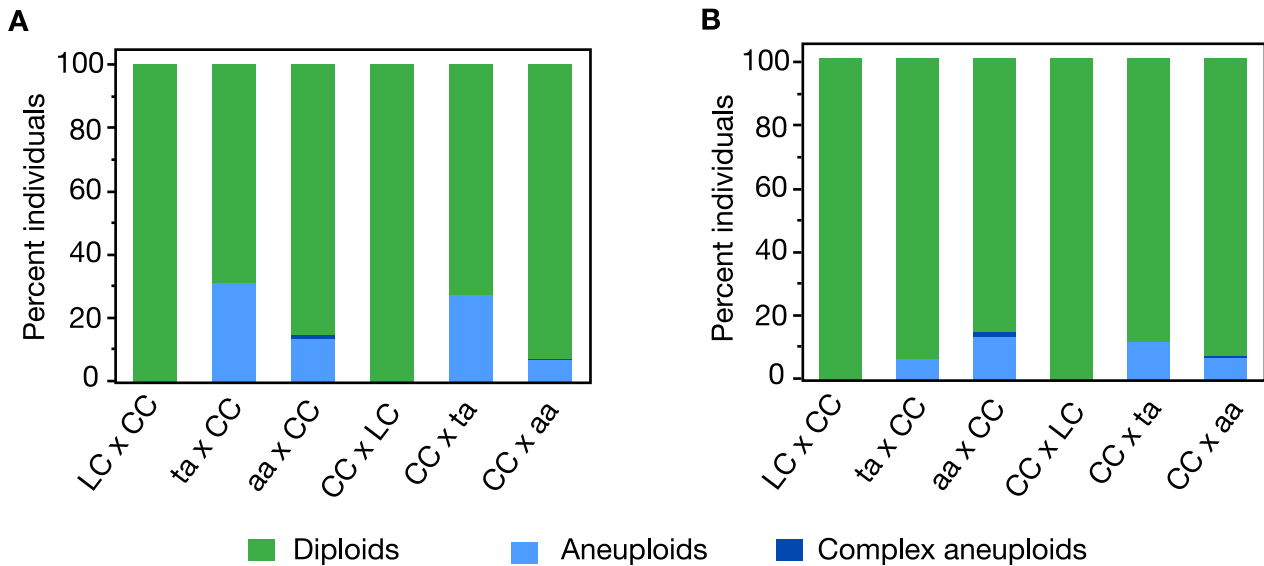
carried an extra copy of the top arm of chromosome 1 but coverage information for that region of the genome was highly variable compared to the diploid individual from the same cross (Fig. S3, panel 1, 3). Comparison with other segmental aneuploids of chromosome 1 confirmed that this level of variability did not originate from methodological artifacts associated with the presence of a third copy of the corresponding sequences (Fig. S3, panel 2). Such a catastrophic genome restructuring event is reminiscent of chromothripsis, when laggard chromosomes are incorporated into micronuclei, and undergo massive DSBs, resulting in restructuring and copy number variation clustered on a single chromosome (61). Chromothripsis is often associated with cancer in humans (61), but has also been observed in *A. thaliana* following CENH3-mediated genome elimination (58). To confirm the variability observed in our low coverage data, we sequenced this individual deeper and confirmed the observed dosage variation (62). Further analysis of the structure of this chromosome confirms the presence of hundreds of novel DNA junctions clustered on the top arm of chromosome 1, consistent with the consequences of chromothripsis (62). It is likely that other plants carrying severely rearranged chromosomes were strongly counter-selected and are not found in our experiments. Thus, this sample provides a unique opportunity to witness a snapshot of the extent of meiotic instability of *asy1* mutants.

Unexpectedly, both of the crosses involving the hypomorphic mutant produced a population of plants enriched in segmental trisomy for chromosome 3B (Fig. 1, individual *ta* x CC #179). The fact that this particular karyotype was clearly overrepresented in those two crosses (56/213 for *ta* x CC and 36/214 for CC x *ta*), but never found in the other four crosses (Table S2), suggests that the hybrid plant carrying the hypomorphic *asy1* mutation itself carried this segmental aneuploidy, and transmitted it to a subset of its progeny. Possibly, this additional copy of chromosome 3B originated from defective meiosis in the parental *asy1*<sup>T142V</sup> or *asy1*<sup>ler-1</sup> mutants that were used to produce the F1 hypomorphic *ta* mutant (see the "Methods" section). Since the additional copy of a chromosomal segment would strongly interfere with the identification of COs on this particular chromatid using the method applied in this study, we discarded all aneuploid progenies from further analysis. The final number of diploid individuals remaining for each population ranged between 117 and 200 (Table S2).

We cannot completely exclude that the likely aneuploidy in the original *ta* line could have affected meiosis beyond producing an over-representation of chromosome 3 aneuploids and hence, the results from this line have to be taken cautiously. However, the comparison between WT and *aa* remains unchanged and notably, the results from the *ta* line are in accordance with the results obtained for *aa* (see below). Thus, we believe that the analysis of the *ta* plants brings additional confidence to our conclusions.

## Number and position of COs are affected by reduced functionality of ASY1

To characterize the number and position of COs in the WT and *asy1* mutant lines, each remaining diploid individual was genotyped at each of 239 nonoverlapping 500 kb marker bins covering the entire *A. thaliana* genome. The data were input into r-qt1 (see the "Methods" section), and separate genetic maps were created for the six populations (Fig. S4). In all cases, the marker order was consistent with the physical order of the bins. We next identified the position of all COs detectable in these samples (Supplementary Material 2).



**Fig. 2.** Percentage of diploid and aneuploid individuals in the various crosses. For each cross-type, the percentage of diploid (green), simple aneuploids (light blue), and complex aneuploids (dark blue) are represented. Complex aneuploids represent individuals with partial chromosomes, suggesting the presence of at least two breaks in a single chromosome. The overall percentage of aneuploid individuals is highest in the crosses involving the *ta* mutant (panel A). In those crosses, extra copies of the bottom arm of chromosome 3 (Chr. 3B) possibly originated from the parental line. Those individuals are discarded in panel B.

The mean number of COs per chromosome varied depending on whether the *asy1* mutant was the male or the female, the chromosome type, and the level of ASY1 functionality (Fig. 3). Specifically, the number of COs per chromosome was not significantly different for the control or mutant crosses for the subtelo-centric chromosomes (chromosomes 2 and 4). On the other hand, there was a significant decrease in CO number for chromosomes 1, 3, and 5 on the male side and chromosomes 1 and 3 on the female side (*t*-test *P* values <0.05 or less, Fig. 3). In most of the cases for female meiosis, the hypomorphic allele exhibited CO numbers that were intermediate between the control crosses and the crosses involving the *asy1* null allele. A similar finding was also obtained for male meiosis in the *asy1* hypomorphs, but only when all chromosomes were combined (Fig. 3). However, it is important to note that, in comparison to the cytological method, a sequencing approach is biased towards viable progenies, especially in mutants where the meiotic process is severely defective. This bias might underestimate the extent of meiotic defects actually present.

Next, we analyzed the chiasma formation in *asy1* mutants by chromosome spreads of male meiocytes at metaphase I. This analysis revealed a significant decrease in the chiasma frequency of *asy1* mutants (*asy1*<sup>Le<sup>r</sup>-1</sup>:  $2.62 \pm 1.39$ , *n* = 47; *asy1*<sup>Col-0</sup>:  $2.78 \pm 1.17$ , *n* = 65; *asy1*<sup>T142V</sup>:  $5.2 \pm 0.3$ , *n* = 41 vs.  $8.7 \pm 0.2$  in the WT, *n* = 47, Fig. 4). To have a resolution at chromosomal level, FISH experiments were performed to distinguish different chromosomes using the 5S and 45S rDNA probes in *ta* mutants in which the occurrence of aneuploids is not so severe as that in *asy1* null mutants. We found that all chromosomes, except for chromosome 2, showed a reduction in CO formation (Fig. 5 and Table S3), and this reduction was higher for the large chromosomes 1 and 5, consistent with earlier work analyzing *asy1* null mutants (12, 53) (Fig. 5C).

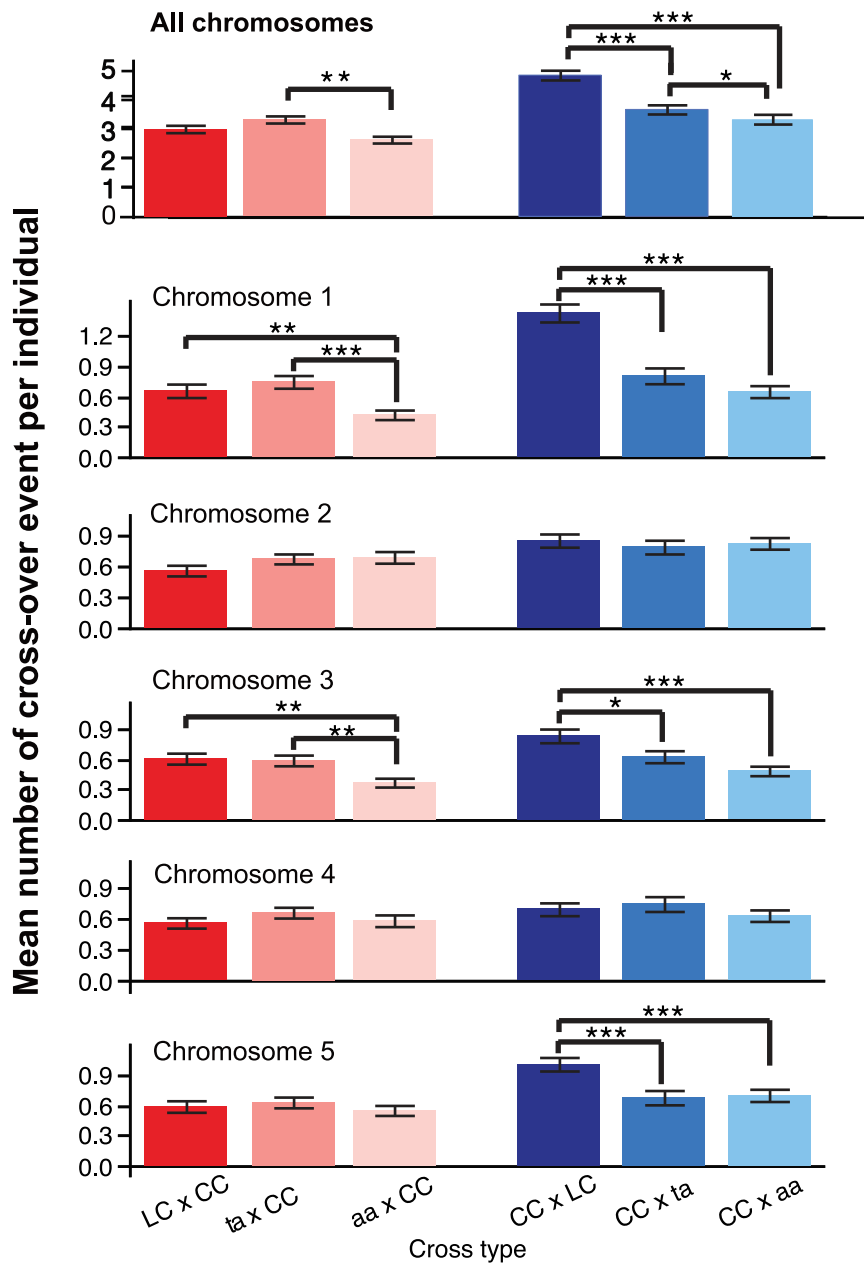
To examine whether the reduction in bivalent formation in *asy1* mutants is due to a reduction in the formation of DSBs, we analyzed RAD51 foci in leptotene/zygotene. We found that RAD51 foci numbers in male meiosis of *aa* ( $136.8 \pm 10.9$ ; *n* = 14) and *ta* ( $134.1 \pm 17.6$ , *n* = 15) were not significantly different from those of the WT LC hybrids ( $134.6 \pm 14.2$ , *n* = 12; *aa* vs. LC *P* = 0.89; *ta*

vs LC *P* = 0.99) (Fig. S5). Therefore, the observed reduction in bivalent number in the *asy1* hybrids does most likely not originate from reduced DSB formation, consistent with previous analysis of *asy1* null mutants (10).

The discrepancy between the observed chiasmata and the COs identified by sequencing, especially for chromosome 4, for which we did not find any significant reduction in CO number in our genetic maps, prompted us to analyze CO distribution within each chromosome (Fig. 6 and Fig. S6). For ease of visualization and comparison, we expressed CO position as the distance to the telomeres, expressed in 0.5 Mb units, and are presenting data from the left and right arms separately to distinguish the effect of the mutations from those of the short arm of the subtelo-centric chromosomes. The distribution of COs was markedly different for the *ta* and *aa* crosses compared to the control crosses in both cross directions (Fig. 6A). Specifically, COs were inclined to localize near the telomeres in the *asy1* mutant crosses, while they were more evenly distributed along the length of the chromosomes in the WT control crosses. The hypomorphic mutant exhibited an intermediate behavior. This trend was not only visible when data for all chromosomes were pooled (Fig. 6A), but also for each individual chromosome type (Fig. 6B), with the exception of the left arms of chromosomes 2 and 4, which are too short to detect changes in distribution. This telomere-proximal accumulation of COs is consistent with previous and our cytological analyses of spread chromosomes, which revealed long-stretched bivalents due to the chiasmata in chromosomal arm ends (60% in *asy1*, 30 out of 50 bivalents) (Fig. 5) (12, 53). These findings also resolved the apparent difference to the cytological work that does not have the resolution to reveal closely spaced COs.

### CO interference is reduced in *asy1* mutants

The observed overaccumulation of COs in distal chromosome positions could be due to two different mechanisms. On the one hand, the number of interference-sensitive type I COs could be reduced at the benefit of additional type II COs, which are not subject to interference. On the other hand, type I COs, which are



**Fig. 3.** The mean number of CO per individual. For each cross-type, the mean number of CO events was calculated for each individual (top panel) and for each chromosome and each individual (all other panels). CO events were only detectable for the hybrid (*Ler*-*Col*) parent, not the *Col*-0 parent, and therefore represent either events that occurred on the paternal side (blue bars) or the maternal side (red/pink bars) only. Since each meiosis produces four gametic cells and each CO event can only be detected from two of the four gametes, the average number of CO per chromosome per individual indicates only half of the CO events formed in each meiosis. For each cross type and chromosome, the mean and SE are represented. Within a cross type and a cross direction, significant differences between the means were tested on a pair-wise basis using Student's t-test. Significant comparisons are indicated as follows: \*\*\*: P value <0.0001, \*\*: P value <0.01, \*: P value <0.05. Lack of significance level between two specific means indicates that the comparison was not significant.

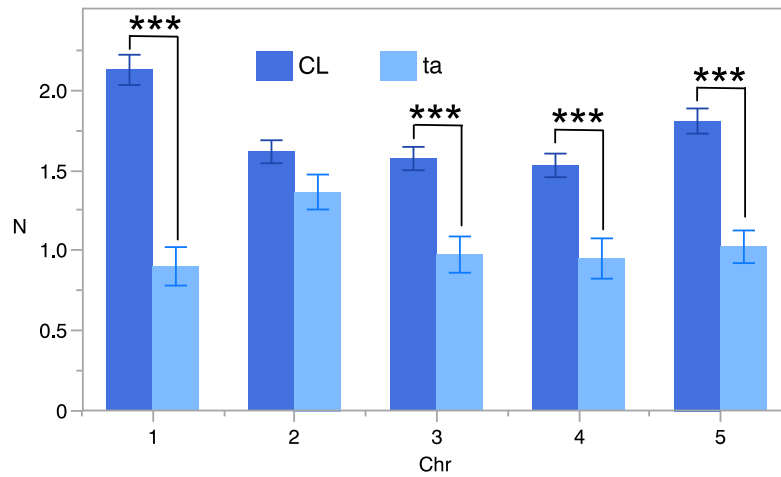
sensitive to interference, might lose their sensitivity in meiocytes with reduced or no ASY1 functionality.

To discriminate between these two possibilities, we generated double mutants of *asy1<sup>Col-0</sup>* with mutants in *MSH4* (also in *Col*-0 background), one key component of type I CO formation, and double mutants of *asy1* with *mus81* (in *Col*-0 as well) in which type II COs are reduced. Depleting *MSH4* in *asy1* caused a drastic reduction of chiasma numbers, i.e., in 25 meiocytes of the *asy1 msh4* double mutants a chiasma frequency of only 0.07 CO per meiocyte was observed, while this number, as shown above, is  $2.78 \pm 1.17$  ( $n = 65$ ) in *asy1* single mutants.

Thus, the majority of the remaining COs in *asy1* belong to the type I class of COs. Consistently, the *asy1 mus81* double mutants did not show a strong reduction of chiasmata i.e., from 57 meiocytes, a chiasmata frequency of  $1.9 \pm 0.75$  per meiocyte was obtained, suggesting that type II COs are present only at a small frequency in *asy1* mutants. Thus, the loss of ASY1 does apparently not grossly alter the relative proportion of type I to type II COs found in the WT.

To complement the genetic analysis, we performed immunolocalization analysis of MLH1, which marks type I COs, in male meiocytes (Fig. 7A and B). In WT-LC hybrids, we observed an aver-





**Fig. 4.** Mean chiasma frequencies per chromosome. The percentages represent the chromosome contribution to the total chiasma frequency. Means and SE values are shown. The mean values were significantly different (t-test  $P$  value  $<0.001$ ) for all chromosomes, except for chromosome 2. The decrease in chiasma frequency was highest for the large chromosomes 1 and 5.

age of  $7.2 \pm 0.3$  MLH1 foci per meicyte ( $n = 48$ ), matching previous analyses. Strikingly, we did not find significant changes in the frequency of MLH1 foci in the *aa* plants ( $7.8 \pm 0.3$ ,  $n = 35$ ;  $P = 0.406$ ). A similar number was also found in the *ta* plants ( $7.8 \pm 0.3$ ,  $n = 52$ ;  $P = 0.405$ ). The discrepancy between the determination of COs in *asy1* mutants by sequencing with the number of MLH1 foci possibly indicates that not all MLH1 foci mature into COs. Notably, MLH1 foci were found on both synapsed and nonsynapsed chromosome regions. However, although the MLH1 antibody and scoring of MLH1 foci have been often used in the past, we could fully exclude technical difficulties and for instance sometimes chromosomes are not clearly stained making it difficult to evaluate the results of the immuno-detection.

Therefore, we sought to complement the MLH1 localization by a previously established live cell imaging setup of meicytes. To achieve this, we first generated a genomic reporter of MLH1 in which the GFP fluorescent tag was inserted immediately before the stop codon ( $PRO_{MLH1}: MLH1: GFP$ , called *MLH1: GFP*), and then transformed this construct into *mlh1* mutants. In contrast to the severe fertility reduction of *mlh1* mutant plants, mutants harboring the *MLH1: GFP* construct were almost fully rescued, as shown by the fully developed siliques and good pollen viability, suggesting that *MLH1: GFP* is functional (Fig. S7A to C).

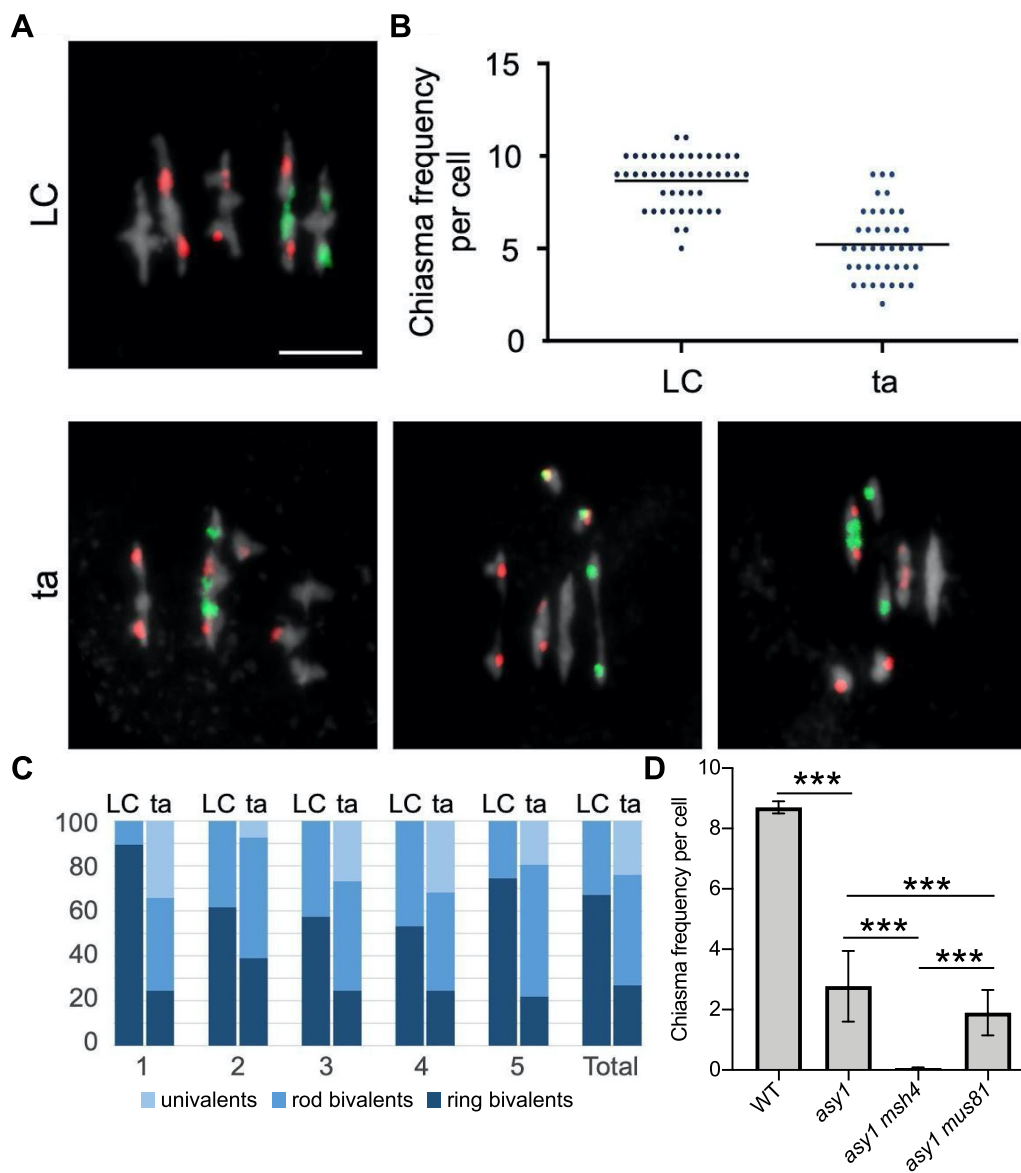
Next, the expression and localization of *MLH1: GFP* at different stages of male meiosis were imaged by laser scanning confocal microscopy. We found that at premeiosis, i.e., S-phase, *MLH1: GFP* was expressed in entire meicytes with many bright foci in the nuclei presumably highlighting DNA mismatch repair sites, which arise during replication (Fig. S7D). Subsequently, from early to mid prophase I, *MLH1: GFP* was only diffusely present in both the cytosol and nuclei of male meicytes, and no particular pattern was observed (Fig. S7D). However, when meicytes reached late prophase I, e.g., late pachytene or diplotene stages, clear foci appeared in the nuclei, presumably marking type I CO intermediates (Fig. S7D).

Next, to check CO distribution, *MLH1: GFP* reporter was introgressed into *asy1* mutant plants and compared to wild-type plants bearing this construct. To cover all MLH1 foci in a meicyte, z-stacks with  $0.8 \mu\text{m}$  spacing intervals were acquired (Supplementary Movies 1 to 4), and the foci were counted using the image analysis software Fiji. Consistent with our immunodetection (Fig. 7B), we found that there was a similar number

of MLH1 foci in *asy1* mutant ( $9.13 \pm 1.52$ ,  $n = 37$ ) when compared to the WT ( $9.42 \pm 1.36$ ,  $n = 34$ ), corroborating not only the presence of type I COs in *asy1* mutants but also that apparently not all MLH1 foci in *asy1* mutants mature into COs. Notably, compared to the WT, closely located MLH1 foci were frequently observed in *asy1* mutants (Fig. 7C, Supplementary Movies 1 to 4).

The preference of telomere-proximal formation of CO in *asy1* mutant, together with the observation that these COs are of type I with largely the same number of MLH1-positive foci suggest that *ASY1* might play a role in CO interference. To further investigate this possibility, we compared the distribution of distances between adjacent COs in the six different crosses (Fig. 8). Inter-CO distance followed a broad distribution that appeared skewed toward short values for the *ta* and *aa* plants. Both female and male distributions in the *aa* mutant were significantly different from that of the corresponding WT crosses (Fig. 8A and B, Kolmogorov-Smirnov adjusted for multiple testing,  $P = 0.03$ ,  $0.02$ , respectively). To further probe this skewness, we divided inter-COs segments into two classes, close and far, arbitrarily changing the dividing line between these two classes and observing the effect on the results (Fig. 8B and C). The female control cross only yielded few instances of double COs (Fig. 8B, LC-CC cross), restricting our ability to observe significant differences between cross types. The *ta*-CC cross was significantly enriched in distant double COs (Fig. 8A and B). Nevertheless, significant enrichment for close events in female meiosis was demonstrated in the *aa*-CC cross when close and far bins were divided at 10 Mb (Fig. 8C, 10 Mb units). The higher recombination frequency in male meiosis provided more cases of syntenic COs than in female meiosis. Thus, the male CC-*aa* cross proved significant for any size bin and the male *ta* mutant displayed strong and significant differences above 1.255 Mb (Fig. 8C).

A direct measurement of interference, such as those obtained by the classical rate of observed vs. expected double COs, was not applicable because the number of individuals used for each cross limited the number of COs and statistical power. Nonetheless, measuring interference on a sliding, nonoverlapping window of 4.5 Mb, suggested reduction of interference at multiple genomic intervals for both mutants, the *aa* null mutation displaying stronger reduction than the *ta* mutation (Fig. S8). The overall distribution of interference values for WT control crosses and



**Fig. 5.** Determination of the chiasmata number in *asy1* mutants using FISH. (A) Representative metaphase I nuclei of LC (up) and *ta* (down) meocytes using FISH to identify the different chromosome types. Red signals (5S rDNA loci) are present in chromosomes 3, 4, and 5, whereas green signals (45S rDNA loci, NORs) appear in the acrocentric chromosomes 2 and 4. DAPI staining appears in gray. Examples of *ta* cells include univalents in different chromosomes: 1 and 3 (left); 4 (middle); 5 (right). Bars represent 5 μm. (B) Quantification of chiasmata in metaphase I cells. Each dot represents an individual cell and bars indicate the mean. (C) Percentage of pairs of univalents, rod bivalents (with chiasmata only in one chromosome arm), and ring bivalents (with at least one chiasma on both chromosome arms) in the WT LC hybrids (left) and the *ta* hybrids (right). (D) Quantification of chiasmata in metaphase I cells in WT, *asy1*, *asy1 msh4*, and *asy1 mus81* mutant plants. \*\*\*  $P < 0.0001$ .

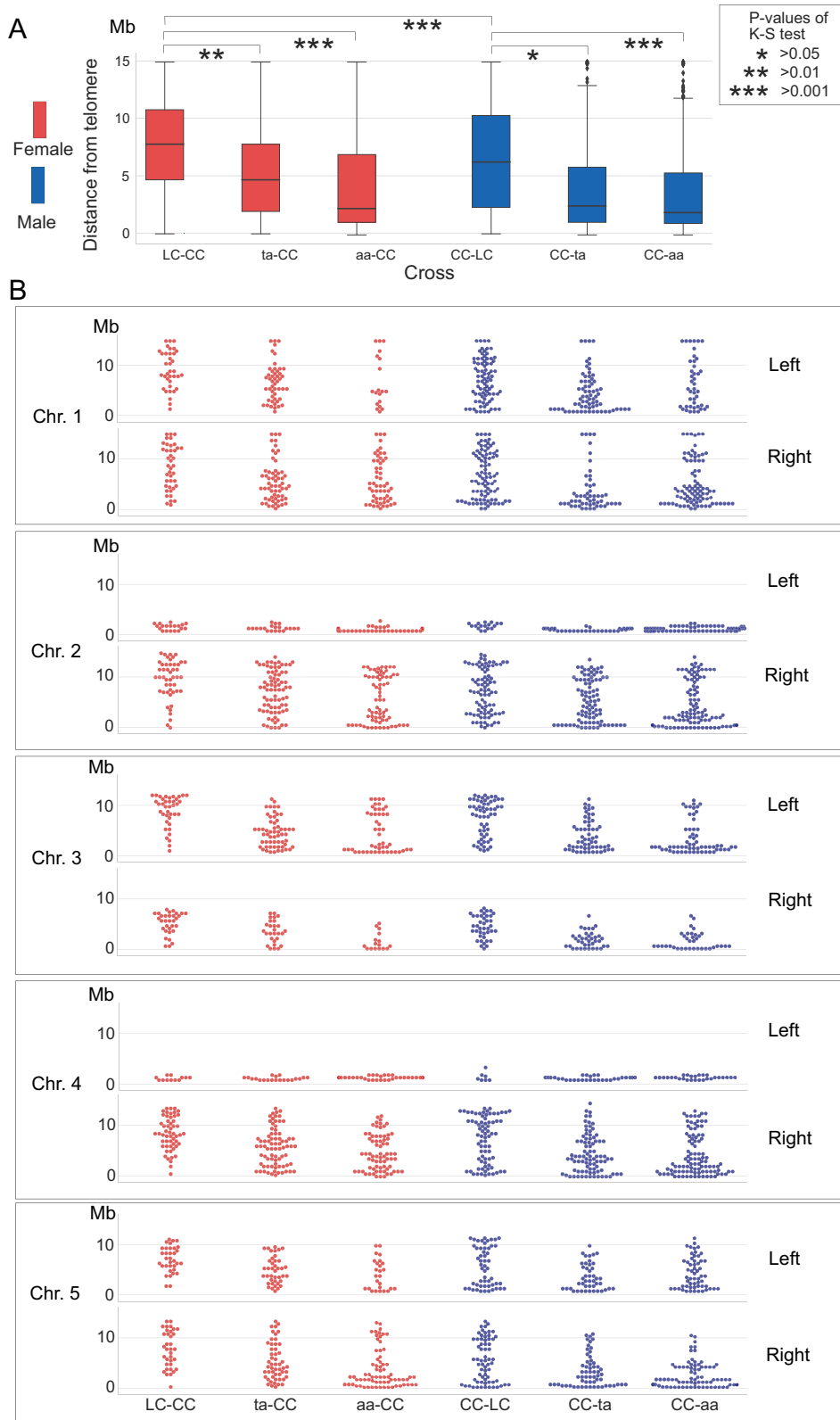
*asy1* mutant crosses was highly significantly different (Fig. S8, Kolmogorov–Smirnov test adjusted for multiple testing,  $P < 0.001$  for all comparisons).

### CO assurance is affected in the ASY1 mutants

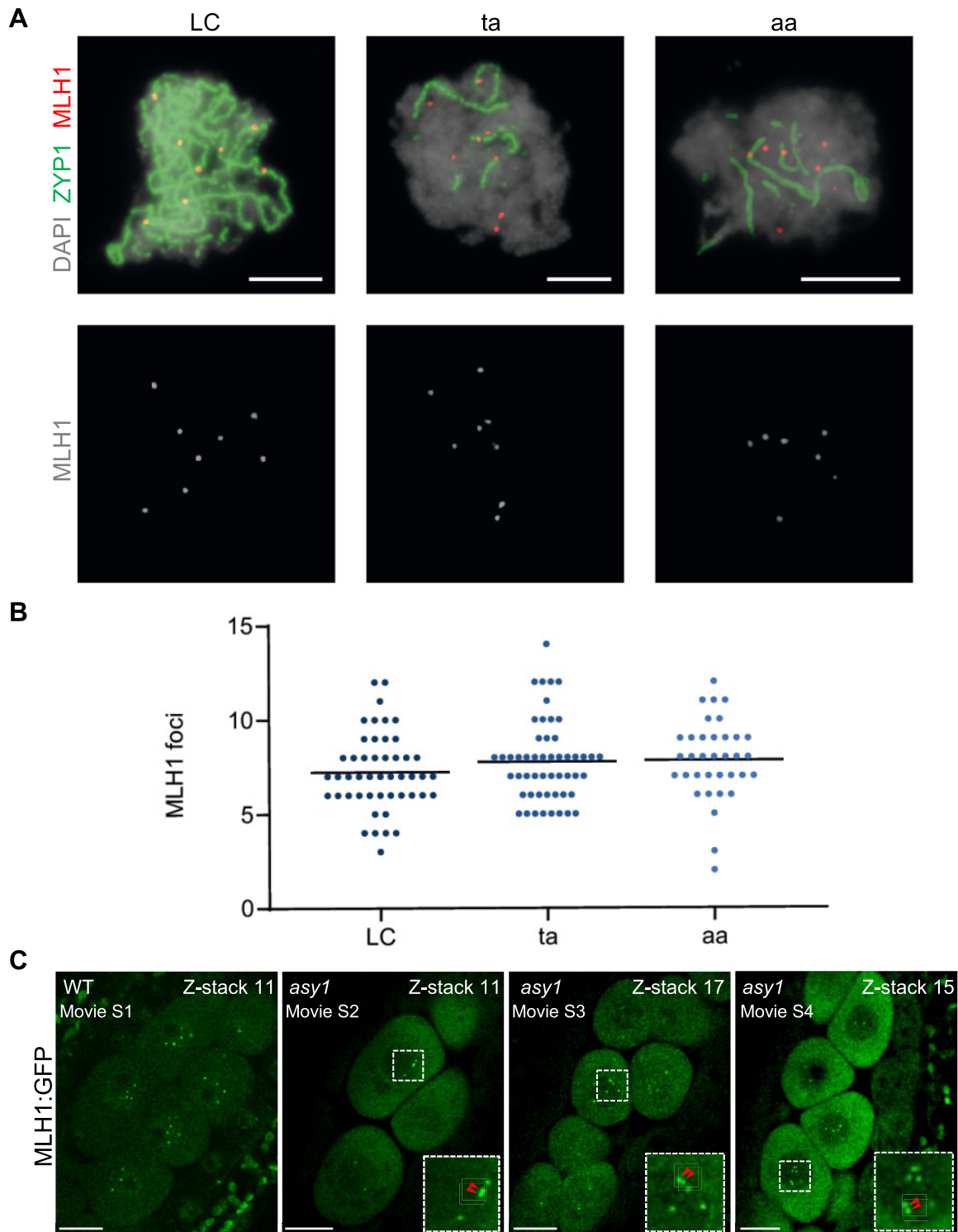
The above data indicated that ASY1 is required for the proper number and distribution of COs along chromosomes. However, a telomeric or subtelomeric CO is sufficient to keep bivalents together, preventing a random distribution of homologs in meiosis I, and subsequent aneuploidy, as seen for instance in many crop species in which COs are usually distally located (33, 63, 64). Likewise, a low number of COs does not necessarily lead to the formation of univalents as exemplified by female meiosis in *Arabidopsis*, during which only about 6 to 7 COs are formed but every chromosome receives at least one CO (17). Thus, according to our se-

quencing data and chiasma counting (Fig. 5 and Table S3), the 5 to 6 COs formed in female meiosis and especially the 6 to 7 COs (reflecting the situation in female WT meiosis) formed in the male meiosis of *asy1* null and hypomorphic mutants could be roughly sufficient to equip every chromosome with a CO obeying CO assurance. However, *asy1* mutants show a high degree of univalent formation (Fig. 5C, Tables S4 and S5), as also reported previously (12, 53). Consistently, we also found significantly more nonrecombined chromosomes in our *asy1* backcross populations compared to the WT control cross (Fig. 9 and Table S6). This suggested impaired CO assurance as well.

The hallmark of CO assurance is the presence of one obligatory CO that causes a nonrandom distribution of COs over the genome, resulting in a deviation from a poisson distribution of COs and an overrepresentation of chromatids exhibiting a single CO (Fig. 9, LC



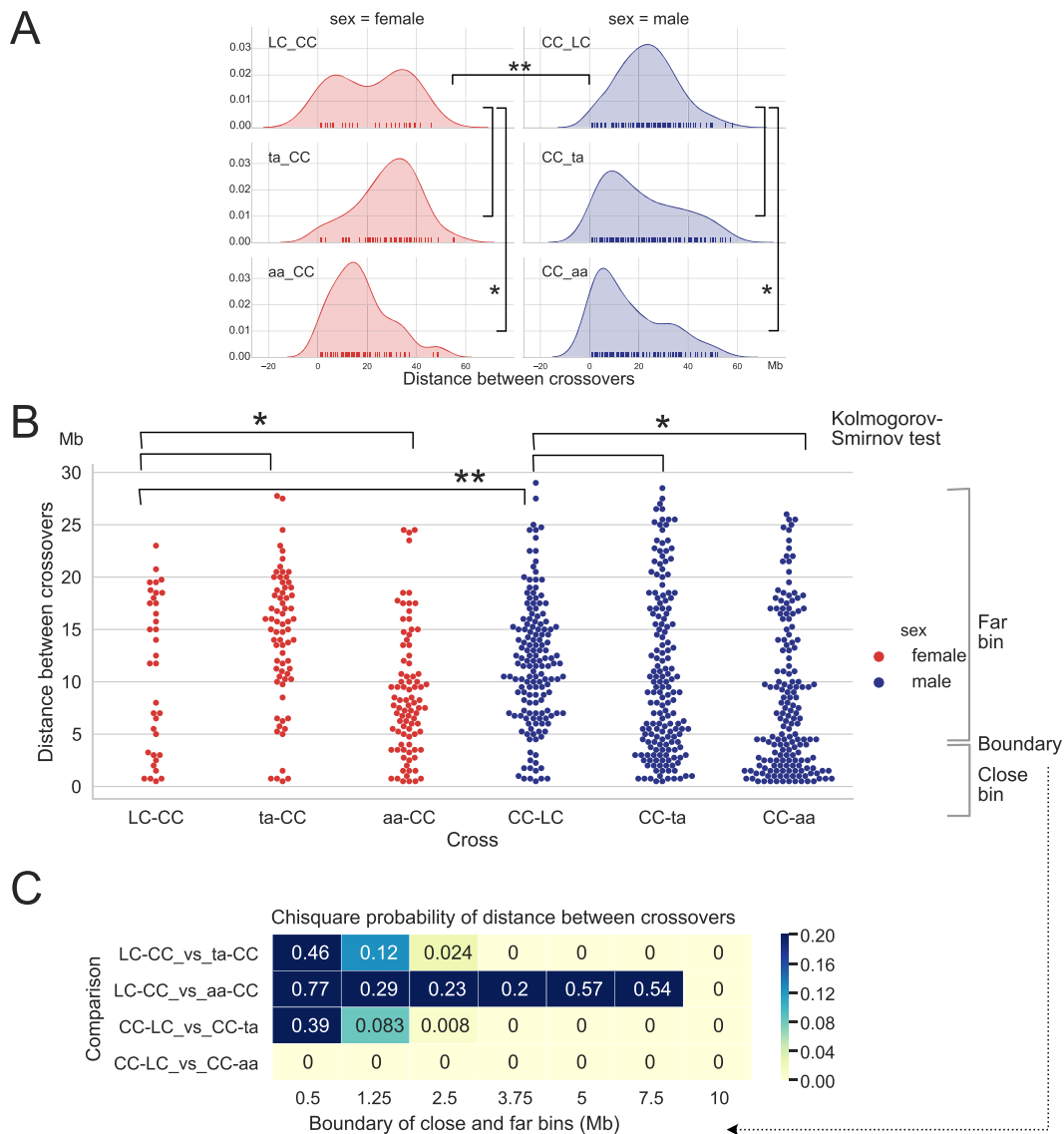
**Fig. 6.** Distribution of CO position in the different crosses. (A) Box plot of the cumulative distributions of CO-telomere distance. The significance of comparisons was tested by the Kolmogorov–Smirnov test (K-S). (B) Swarm plots of CO-telomere distance per chromosome. The distance from the CO to the telomeres is shown and the maximum distance for each arm is at the centromere.



**Fig. 7.** Analysis of MLH1 foci. (A) Representative images showing pachytene (LC) and pachytene-like (ta, aa) meocytes after immunolabeling for detecting the class I CO marker MLH1 (red). The central element protein of the SC ZYP1 is detected in green and DAPI staining appears in gray. Bars represent 5  $\mu$ m. (B) Comparison of the number of MLH1 foci. Each dot represents an individual cell and bars indicate the mean. (C) Presence of MLH1 foci in WT and *asy1* mutant plants. One of the z-stacks was shown from Supplementary Movies 1 to 4. Red arrows indicate the closely located MLH1 foci. Bars represent 10  $\mu$ m.

x CC cross). In contrast, in the ta x CC progenies and to an even larger extent in the aa x CC progenies, we consistently observed distributions of COs that were closer to the expected random distribution (asterisks, significance test in Fig. 9). If the CO distribu-

tion deviates in the *asy1* crosses from a random distribution, it is usually due to an underrepresentation of the 1-CO class, e.g., chromosome 1 in male meiosis (Fig. 9, lower three panels), combined with an overrepresentation of both nonrecombined chromatids



**Fig. 8.** The Frequency of COs and the distance between COs. CO to CO distance distribution displayed as (A) Kernel Density Estimation and (B) swarm plots, using in 1 Mb units. Knock out of ASY1 is associated with reduced distance between COs. C. Chi-square probability for comparison of close vs. far binned CO-CO distances. The respective upper limit of the close bin varies and is shown below each column of P values. Yellow denotes significant difference by the Chi square test. Note that significance for the ta-CC cross was caused by an increase of the distant CO class, as opposed to an increase in the number of close COs, as observed in the aa crosses. Significance thresholds: \* <0.05, \*\* <0.01.

(diamonds, significance test in Fig. 9), and chromatids with more than one CO. Thus, we conclude that another key feature of COs, i.e., CO assurance, is also affected by the loss of ASY1 function, namely, ASY1 is required for the acquisition of one obligatory CO for each homolog pair.

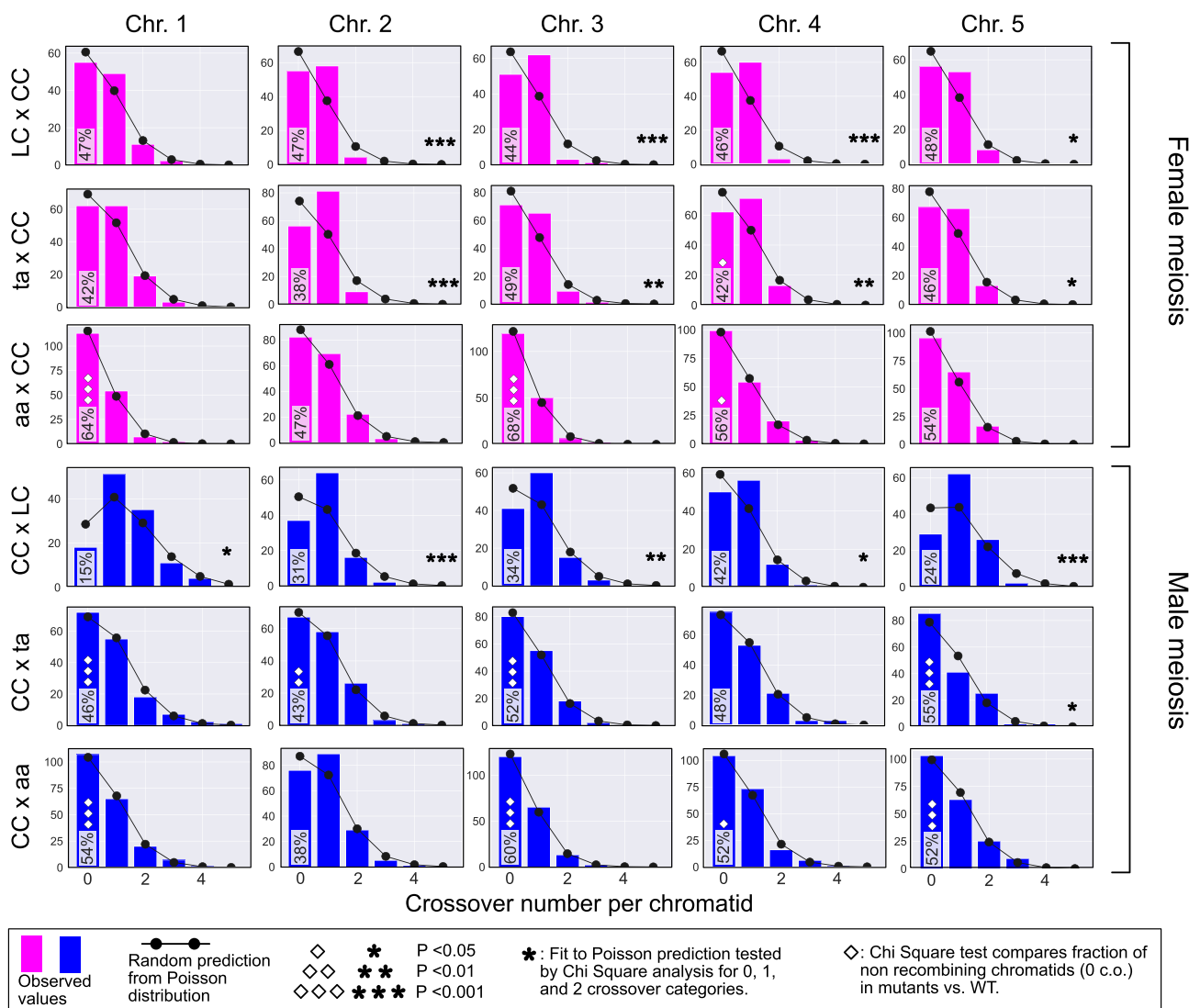
### Discussion

ASY1 and its homolog Hop1 are important for several aspects of meiotic recombination (8–12). Here, we have focused on the role of ASY1 in CO placement in female and male meiosis. Mapping COs in hybrids between two different *Arabidopsis* accessions that are mutant for ASY1 has allowed us to reveal several important and previously not recognized aspects of the function of ASY1, foremost that ASY1 mediates CO interference and plays an important role for CO assurance. Using two *asy1* alleles with different strengths shows that these two aspects are sensitive to the

level of ASY1 functionality. Our findings are consistent with another report by Lambing et al. that appeared while our paper was in revision (65). Lambing et al. demonstrate that ASY1 is involved in CO interference and counteracts telomere-led CO placements in male meiosis.

Interestingly, CO interference was also reduced in yeast cells mutant for the AAA-ATPase PACHYTENE CHECKPOINT2 (PCH2) (66–70). PCH2 has been previously found to be important for the removal of ASY1 and Hop1 to promote assembly of the SC (69, 71, 72). This raises the question of why loss of ASY1 and a failure to remove it can cause similar mutant phenotypes. However, recent work indicated that PCH2, at least in *Arabidopsis*, also plays a role in delivering ASY1 into the nucleus, and in the absence of PCH2 or its cofactor COMET, ASY1 strongly accumulated in the cytoplasm while ASY1 is a solely nuclear localized protein in the WT (36, 73). Thus, it is possible that the timely and correct association of ASY1 with the axis, especially during early phases of axis for-





**Fig. 9.** Comparison of CO numbers per chromatid in WT and *asy1* mutants. The bar plots illustrate the count of chromatid classes for given genotypes and chromosomes. Pink and blue bars are for female and male meiosis, respectively. The random expectation according to the Poisson distribution is displayed by the black line with round black circles. The star markers display significance of the random fit (low P corresponds to significant deviation). Note that almost all chromosomes for both test crosses significantly deviate from a random distribution while all *asy1* null mutants and most of the *asy1* hypomorphic mutant show a pattern that matches a Poisson distribution. The % of the 0 CO chromatid class is displayed inside the 0 CO bar. For this fraction of 0 CO chromatids, the P of the Chi Square comparison with WT values is marked with diamonds (low P corresponds to significant deviation). The exact number of instances in each category can be found in Table S6.

mation, could be compromised in *pch2* reconciling the *pch2* and *asy1* mutant effects with respect to CO interference. Conversely, we postulate that interference should also be affected in *comet* mutants.

Pairing and synapsis usually start in the distal regions of homologs in *Arabidopsis* (74–76). Notably, despite the general failure of chromosome alignment and synapsis in the *asy1* mutant, telomere clustering and the pairing of telomeres of homologs has been found to occur during meiotic interphase and early prophase I (50). Thus, the placement of COs in the distal regions of homologs in *asy1*, as observed here and in previous studies, is consistent with this observation (12, 53). With that respect, it is interesting to note that synapsis occurs in only short chromosomal sections in *Sordaria mer2* mutants and COs tend to cluster in these sections, thus also modulating the strength of interference (77).

We also observed that not all MLH1 foci in *asy1* matured into COs and hence, ASY1 directly or indirectly is involved in CO matu-

ration. Since several MLH1 foci were found outside of synapsed regions, it is an intriguing question to test in the future whether only MLH1 foci on synapsed regions will mature into COs. However, recent studies have shown that loss of ZYP1 in *Arabidopsis* does not abolish COs and hence, synapsis per se is not a prerequisite for CO formation (78–80). Instead, the maturation of MLH1 foci has been previously found to depend on the pairing regulator Ph1 in rye wheat hybrids (81). However, how Ph1 charges MLH1 foci is not clear and whether there is a direct link to ASY1 action at the chromosome axis needs to be resolved in the future. Remarkably, even the hypomorphic *asy1* mutant showed a strong distalization of COs. Therefore, it is plausible that even a slight modulation of the axis and/or the speed by which the axis is formed could restrict COs to distal positions while maintaining a high level of fertility, as seen in the hypomorphic *asy1*<sup>T142</sup> mutant, which is semifertile (36).

In this context, it is interesting to note that many polyploid species show a distalization of COs based on cytological analy-

ses (32, 82). Moreover, mutations in *ASY1* and of the axis component *ASY3* have been found to be specifically associated with tetraploidy in the *Arabidopsis arenosa* species, hinting at an adaptive advantage of these mutations in polyploids (34, 83). While the benefit of distalized COs in polyploids is obscure, CO numbers have often been found to be reduced in polyploids, and this effect would be very beneficial for the faithful segregation of species with more than two homologs, as it would reduce the level of chromosomes that are interconnected by COs and hence prone to missegregation (32). Reduced CO number in polyploids has been speculated to be due to increased interference (32). Our findings offer an alternative explanation, i.e., reduction/slowing down of pairing with concomitant reduction of interference. Moreover, our findings would also explain why COs are distalized in polyploids. However, whether an *asy1*-dependent decrease in interference represses missegregation in polyploids can only be a valid explanation if the remaining COs that form in the distal region of chromosomes are more likely to all concern the same pair of homologs. Such an exclusive CO position could possibly be forced by sterical constraints between different homologs. The determination of CO patterns by sequencing in polyploid species will help resolve this question. In addition, live cell imaging of meiosis, as recently established for *Arabidopsis* (84), and tracking of chromosomes in polyploid species could be a very powerful tool to address this point.

Why interference is reduced in *asy1* mutants remains unclear and can, as far as we can tell, not be explained by the so far known functions of *ASY1* in meiosis, e.g., interhomolog versus intersister repair bias. Applying the beam-film model of interference to our data (22), leads to the speculation that a reduction in functional *ASY1* protein levels would prevent the tension relaxation brought about by a CO to spread along the chromosome, possibly because the chromosome axis is not connected by *ASY1*. Consistently, the *ASY1* homolog *Hop1* has been proposed, based on *in vitro* data, to build long head-to-tail polymers (85, 86). Such a polymer could possibly transmit a relaxation force. However, at least in such a model, it seems unlikely that *ASY1* itself contributes to the tension since, in that case, no COs would be expected in *asy1* mutants in the first place. Hence, more likely under the assumption of the beam-film model, an *ASY1* polymer, if built *in vivo*, could serve as a platform for other proteins that help relaxing the mechanical stress, such as *topo II* (87). Interestingly, *TOPO II* has been coprecipitated with *ASY1* from *Brassica oleracea* meiocytes (88). However, it is currently not clear whether this interaction is direct or whether *TOPO II* in *Brassica* is also involved in interference control.

In the light of the recently proposed model of CO interference by diffusing mediated coarsening of the proCO factor *HEI10* (24), one could speculate that loss of *ASY1* limits *HEI10* to telomeric/subtelomeric regions and by that promotes the formation of type I COs there. However, why this should be the case is not clear as yet and needs further exploration.

Our work showed that *ASY1* regulates not only interference but also CO assurance. The beam-film model predicts that CO assurance is not a function of interference, although the initial obligatory CO drives interference subsequently by providing the starting gun for mechanical stress. While some mutants that affect interference in yeast, were found to still undergo an obligatory CO, such as mutants in *TOPO II*, CO assurance and interference were concomitantly compromised in others mutants, e.g., in mutants of *MutS HOMOLOG4 (MSH4)* (20, 89). We currently do not know whether the effect of *ASY1* on interference and assurance have the same biochemical foundation, or reflect two different functions of *ASY1*.

Moreover, the observation that the class with one CO is often underrepresented in *asy1* mutant populations in combination with closely spaced COs in the telomeric and subtelomeric regions suggests that *asy1* mutants could represent a rare case of negative interference, i.e., the attraction of COs after a first CO is formed (5). Notably, we found that the extent of interference and assurance reduction in *asy1* mutants appears to depend on the individual chromosome for yet unknown reasons. Thus, the exploration of the dynamics and function of the chromosome axis in CO interference and CO assurance, ideally with chromosome-specific resolution, remains an exciting question.

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## Supplementary Material

Supplementary Material is available at [PNAS Nexus](#) online.

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## Authors' Contributions

A.S., I.H., and Lu.C. designed the research; G.P., I.M., C.Y., N.F.J., N.L., L.C., H.T.T., and M.P., performed the experiments. G.P., I.M., C.Y., M.P., Lu.C., and A.S. analyzed the data. I.H., C.Y., M.P., Lu.C., and A.S. wrote the manuscript.

## Data Availability

All material generated in this study is freely available from A.S. Sequence data from this article can be found and is freely accessible in the NCBI SRA database under BioProject number PRAJNA 723,952.

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