

ASY1 acts as a dosage-dependent antagonist of telomere-led recombination and mediates crossover interference in *Arabidopsis*

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During meiosis, interhomolog recombination produces crossovers and noncrossovers to create genetic diversity. Meiotic recombination frequency varies at multiple scales, with high subtelomeric recombination and suppressed centromeric recombination typical in many eukaryotes. During recombination, sister chromatids are tethered as loops to a polymerized chromosome axis, which, in plants, includes the ASY1 HORMA domain protein and REC8-cohesin complexes. Using chromatin immunoprecipitation, we show an ascending telomere-to-centromere gradient of ASY1 enrichment, which correlates strongly with REC8-cohesin ChIP-seq data. We mapped crossovers genome-wide in the absence of ASY1 and observe that telomere-led recombination becomes dominant. Surprisingly, asy1/+ heterozygotes also remodel crossovers toward subtelomeric regions at the expense of the pericentromeres. Telomeric recombination increases in asy1/+ occur in distal regions where ASY1 and REC8 ChIP enrichment are lowest in wild type. In wild type, the majority of crossovers show interference, meaning that they are more widely spaced along the chromosomes than expected by chance. To measure interference, we analyzed double crossover distances, MLH1 foci, and fluorescent pollen tetrads. Interestingly, while crossover interference is normal in asy1/+, it is undetectable in asy1 mutants, indicating that ASY1 is required to mediate crossover interference. Together, this is consistent with ASY1 antagonizing telomere-led recombination and promoting spaced crossover formation along the chromosomes via interference. These findings provide insight into the role of the meiotic axis in patterning recombination frequency within plant genomes.

meiosis | crossover | axis | ASY1 | interference

eiosis is a specialized cell division that increases genetic diversity in populations (1, 2). Meiosis halves the chromosome number to produce haploid gametes via a single round of DNA replication and two rounds of chromosome segregation (1, 3). During prophase I of meiosis, homologous chromosomes undergo DNA double-strand breaks (DSBs) that can be repaired using an interhomolog pathway, which may result in crossovers or non-crossovers (1, 3). In plants, meiotic DSBs are formed via a topoisomerase-VI-like complex containing SPO11-1, SPO11-2, and MTOPVIB (4). Meiotic DSBs are resected to form 3'overhanging single-stranded DNA (ssDNA), which is bound by the RecA homologs RAD51 and DMC1 that promote strand invasion of a homolog (1, 3). A set of pro-crossover factors, termed the ZMM pathway, act to protect interhomolog strand invasion events from antirecombination pathways (3). Class I crossover events generated via the ZMM pathway are more widely spaced along the chromosomes than expected by chance, which is known as interference (5). A minority of crossovers are generated by the Class II repair pathways in wild type, which do not show interference (3).

Homologous chromosomes associate with a specialized axis structure during meiosis, which is conserved across eukaryotes and is required for efficient and accurate interhomolog recombination (6). Following S-phase, replicated sister chromatids are associated via cohesin complexes containing the meiosis-specific kleisin REC8 (7, 8). Immunostaining of REC8 during prophase I reveals a linear axis, to which the chromatin is attached (6). In addition to REC8-cohesin, major components of the plant meiotic chromosome axis include the HORMA domain protein ASY1 and the coiled-coil proteins ASY3 and ASY4 (9-12). In this configuration, coaligned chromatin loops project laterally from the axis, resembling mitotic lampbrush configurations, although with a juxtaposed homolog (6). The tethered-loop axis model proposes that meiotic DSBs are generated on the chromatin loops that become tethered to the axis during interhomolog repair (6, 13). Axis-localized HORMA domain proteins are required during meiosis to promote homolog pairing, DSB repair, and synaptonemal complex (SC) assembly (14–19). However, there are also important differences in the function of meiotic HORMA proteins between species. For example, mouse HORMAD1, budding yeast Hop1, and Caenorhabditis elegans HTP-3, but not Arabidopsis ASY1, are required for meiotic DSB formation (9, 15, 20-22). In late prophase I, the axis is remodeled, which is associated with depletion of HORMA proteins and loading of transverse filament SC proteins, including ZYP1a and ZYP1b (18, 23).

Significance

Meiosis is fundamental to eukaryotic reproduction and shapes patterns of genetic variation. Meiotic recombination is also a vital tool during crop improvement, which allows introgression of wild variation into agricultural strains. Despite this, the levels and distributions of crossovers along chromosomes can limit breeding. For example, many crops show highly skewed crossover distributions toward the telomeres. This can lead to the problem of linkage drag when variation within nonrecombining regions is selected. Our findings demonstrate how gene dosage of key components of the meiotic chromosome axis can be used to remodel the recombination landscape. Therefore, modifying *ASY1* and *ASY3* gene dosage in crop species may provide a strategy to change recombination patterns or levels in order to accelerate strain improvement.

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Data deposition: ASY1 ChIP-seq library data have been deposited in the ArrayExpress database at EMBL-EBI (http://www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-8705. Sequencing data for wild type and asy1 Col×WS GBS libraries have been deposited under ArrayExpress accession E-MTAB-8715, and data for asy1/+ Col×Ler GBS libraries has been deposited under ArrayExpress accession E-MTAB-8725.

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Genome-wide analyses have revealed that meiotic DSB and crossover frequency are highly variable between the telomeres and centromeres of plant chromosomes (24-29). For example, the centromeres and surrounding repetitive sequences (pericentromeric heterochromatin) are frequently suppressed for meiotic recombination (24-29). High meiotic crossover levels are typically observed in distal subtelomeric regions, which also tend to have higher gene density (24-29). However, the factors and mechanisms that shape the meiotic recombination landscape along chromosomes remain incompletely understood. To investigate the role of the axis during meiosis, we mapped ASY1 enrichment throughout the Arabidopsis genome using chromatin immunoprecipitation sequencing (ChIP-seq). We observe an ascending ASY1 gradient from the telomere to the centromere, which correlates positively with REC8 ChIP-seq data (30). We mapped crossovers genome-wide in asy1 mutants and observe that recombination becomes telomere-led, likely reflecting telomere pairing observed early in prophase I (31). We show that asy1/+ heterozygotes maintain crossover numbers but remodel recombination frequency toward the telomeres at the expense of the pericentromeres. The zone of telomere-led recombination in asy1 and asy1/+ corresponds to distal regions of the chromosomes with lowest ASY1 and REC8 ChIP-seq enrichment in wild type. Through analysis of double crossover distances, fluorescent pollen tetrads, and MLH1 foci, we show that crossover interference is normal in asy1/+ heterozygotes, but is undetectable in asy1 homozygotes. Together, our data show that ASY1 exerts a major effect on the crossover landscape via mediating interference and acting as a gene dosage-dependent antagonist of telomere-led recombination.

Results

Telomere–Centromere Gradients of ASY1 and REC8 ChIP-Seq Enrichment. To investigate the genome-wide localization of ASY1, we performed ChIP-seq using a polyclonal rabbit α -ASY1 antibody raised against full-length recombinant protein (12). Immunostaining of anther spreads using the α -ASY1 antibody shows specific detection in meiocytes, and not in adjacent somatic cells (Fig. 1A). Coimmunostaining of ASY1 and REC8-HA showed highly correlated signals during early prophase I (signal intensity correlation r = 0.76-0.85, n = 10; Fig. 1B). We performed ChIP-seq using the α-ASY1 antibody on meioticstage floral buds and obtained two independent biological replicate libraries with 26,488,565 and 39,593,737 mapping read pairs (17.5× and 28.2× genome coverage, respectively; SI Appendix, Table S1). The ChIP-seq replicates are highly correlated at the genome scale ($r_s = 0.91$ using 10-kb adjacent windows; SI Appendix, Table S2). To determine the specificity of ASY1 ChIPseq enrichment, two controls were performed. First, the α -ASY1 antibody was used for ChIP-seq from leaf tissue, where ASY1 is not expressed (12). Second, preimmune serum was used for ChIP-seq from floral tissue. After deduplication, only 0.29% and 0.39% of reads in these libraries mapped to the Arabidopsis genome (SI Appendix, Table S1). In contrast, 90.1% and 93.2% of deduplicated ASY1 ChIP-seq reads were mapped (SI Appendix, Table S1). This demonstrates the low background of reads that map to the Arabidopsis genome obtained from our ChIP protocol, in the absence of the epitope or the α -ASY1 antibody. For further analysis, ASY1 ChIP-seq libraries were normalized using an input chromatin library to generate log₂(ChIP/input) enrichment values across the genome (Fig. 1C).

At the genome scale, we observed highest ASY1 ChIP-seq enrichment over the centromeric and pericentromeric regions (Fig. 1*C* and *SI Appendix*, Fig. S1). An ascending gradient of ASY1 ChIP-seq enrichment was observed from telomeres to centromeres, with the sharpest increase observed as the centromeres are approached (Fig. 1*D*). We observed a striking positive correlation between ASY1 and REC8-HA ChIP-seq enrichment (e.g., $r_s = 0.88-0.93$ at 10-kb scale; Fig. 1C and SI Appendix, Fig. S1 and Table S2) (30), which is consistent with their highly correlated immunostaining patterns (Fig. 1B). We compared ASY1 ChIP-seq enrichment to DSBs using SPO11-1-oligos as a marker (Fig. 1C) (25). At the chromosome scale, the regions in proximity to the centromere where ASY1 is highest have the lowest DSBs (Fig. 1C). However, when considering the chromosome arms alone, ASY1 and SPO11-1-oligos show a weak positive correlation ($r_s = 0.48$ at 10-kb scale; Fig. 1 C and D and SI Appendix, Fig. S1). At the fine scale, SPO11-1-oligos are highest within nucleosome-depleted gene promoters and terminators (Fig. 1E) (25). In contrast, ASY1 and REC8 are highest within nucleosome-enriched gene bodies (Fig. 1E) (30). Variation in ASY1 enrichment within genes correlates positively with REC8 and nucleosome occupancy (MNase-seq), but does not correlate with SPO11-1-oligos in gene promoters or terminators (Fig. 1F). Equally, variation between genes in promoter SPO11-1-oligo levels does not correlate with ASY1 or REC8 ChIP-seq enrichment within gene bodies (Fig. 1F).

Telomere-Led Recombination Dominates in *asy1* **Mutants.** As we observed a gradient of ASY1 ChIP-seq enrichment between the telomeres and centromeres, we sought to investigate crossover patterning along chromosomes in *asy1* mutants. Homozygous *asy1* mutants have low fertility due to reduced chiasmata and a high incidence of univalent chromosomes at metaphase I, which leads to aneuploid gametes (9, 15). Despite this, low numbers of viable progeny can be obtained from *asy1* homozygotes. Therefore, we crossed *asy1*+ individuals in Col [*asy1-4* (15)] and Ws-4 [hereafter Ws; *asy1-3*/+ (32)] backgrounds to generate wild type or *asy1* Col×Ws F₁ plants. The F₁ plants were self-fertilized, and 187 wild type and 169 *asy1* F₂ progeny were generated and used for DNA sequencing. The TIGER pipeline was used to identify crossover locations from the sequencing data (*SI Appendix*, Table S3) (33).

As expected, a significant decrease in crossovers per F_2 was observed in *asy1* (mean = 4.6) compared to wild type [mean = 7.9; Mann–Whitney–Wilcoxon (MWW) test $P = 4.37 \times 10^{-37}$; Fig. 2A and SI Appendix, Table S3]. However, the number of crossovers observed per asy1 F2 individual was higher than predicted from bivalent counts per meiosis in asy1-3 [mean = 1.5 (32)] and asyl-4 [mean = 1.9 (9); Fig. 2A and SI Appendix, Table S3]. This may reflect generation of viable F_2 plants selecting for gametes with at least one crossover per chromosome in order to balance segregation at metaphase I. Alternatively, as chiasmata measurements are made from male meiosis, whereas F2 crossover data reflect both male and female meiosis, this could indicate sex differences in crossover reduction in asy1. A further possibility is that closely spaced crossovers may be counted as single chiasmata in asy1, causing an underestimation of recombination. In wild type, crossover number per chromosome is positively correlated with physical length (r = 0.98, $P = 3.4 \times 10^{-3}$), whereas no significant correlation exists in asy1 (Fig. 2B). Exceptionally, chromosome 2 shows a crossover frequency close to wild type in asy1, with a striking increase on the short, nucleolar organizing region (NOR)-bearing arm (Fig. 2 B and D and SI Appendix, Table S3). This is consistent with chiasmata and fluorescent in situ hybridization (FISH) analysis in asy1 mutants in the Ws accession, where the short arm of chromosome 2 also showed high chiasmata frequency (34). Interestingly, in Col×Ws F₁ hybrids, NOR2 rDNA gene clusters are transcriptionally silenced, whereas NOR4 on chromosome 4 are expressed (35). Nucleolar silencing is known to involve formation of heterochromatin at the transcriptionally repressed NOR (36). Hence, heterochromatin formation at NOR2 could contribute to closer alignment of homologs and thereby promote crossover formation on chromosome 2 in asy1.

When recombination was analyzed along scaled telomere-centromere axes, we observed a strong bias of *asy1* crossover formation toward the subtelomeric regions (Fig. 2 *C*–*E*). Analysis



Fig. 1. The landscape of ASY1 ChIP-seq enrichment throughout the *Arabidopsis* genome. (*A*) Meiotic cells in early prophase I or adjacent somatic cells immunostained for ASY1 (green) and stained for chromatin (DAPI, blue). (*B*) Male meiocytes in early prophase I immunostained for ASY1 (green) and REC8-HA (red) and stained for chromatin (DAPI, blue). (*Inset*) Correlation plot of ASY1 and REC8-HA signal intensity (n = 10 cells). (Scale bars, $10 \,\mu$ M.) (C) ASY1 (blue) and REC8-HA (pink) ChIP-seq enrichment [log₂(ChIP/input)] and SPO11-1-oligos [log₂(SPO11-1-oligos/gDNA), red] along the *Arabidopsis* genome in adjacent 10-kb windows, smoothed using a moving average (25, 30). Vertical solid and dotted lines indicate the telomeres and centromeres, respectively. The pericentromere boundaries are indicated by pale blue ticks on the *x* axis. (*D*) Data as in *C*, but analyzing proportionally scaled chromosome arms from telomeres to centromeres. (*E*) Data as in *C*, but showing between gene transcriptional start (TSS) and termination (TTS) sites and 2-kb flanking regions. The same number of randomly positioned windows of the same widths were analyzed as a control. (*F*) Data as in *C*, but analyzed as heat maps within genes and 2-kb flanking regions. Genes were ranked by ASY1 levels in gene bodies (TSS–TTS; *Upper*) or by promoter SPO11-1-oligos (*Lower*). Shading is equal to defined quantiles of coverage values mapped linearly to a vector of six colors.



Fig. 2. Telomere-led recombination predominates in *asy1*. (A) Histograms of crossovers per F_2 individual for wild type and *asy1*. Red dashed lines indicate mean values. (*B*) crossovers per chromosome per F_2 for wild type (blue) and *asy1* (red) plotted against chromosome length in megabases. (*C*) crossover frequency in wild type (blue) and *asy1* (red), and ASY1 ChIP-seq enrichment [log₂(ChIP/input), black] analyzed along proportionally scaled chromosome arms, oriented from telomeres to centromeres. Col×Ws SNP density is shown by green shading. (*D*) Crossover frequency (crossovers/150 kb per F_2) plotted along the *Arabidopsis* genome for wild type (blue) and *asy1* (red), with Col×Ws SNP density (green) shaded. Telomere and centromere positions are indicated by vertical solid and dotted lines, respectively. (*E*) Comparison of F_2 genetic map lengths (in centimorgans, cM) in wild type (blue) and *asy1* (red). (*F*) Spreads of male meiocytes at metaphase I in wild type (Col) and in *asy1* and *asy3* mutants with DAPI-stained chromatin. Arrows indicate potentially distal chiasmata locations. (Scale bars, 10 μ M.) (*G*) Crossover positions analyzed relative to the closest telomere in wild type (blue) and *asy1* (green) analyzed over the same regions.

of chiasmata in *asy1*, *asy3*, and *asy4* axis mutants has shown a high incidence of rod bivalent configurations (Fig. 2F) (9, 10, 34), which may reflect distal crossover locations. To investigate

recombination in relation to telomere position, we assigned each crossover a distance to its nearest telomere and plotted events on a common axis (Fig. 2G and SI Appendix, Fig. S2). The crossover

counts observed were analyzed in windows relative to the telomere in wild type and *asy1* and used to perform χ^2 tests, with correction for multiple testing (*SI Appendix*, Table S4). We observed that windows in the first megabase of chromosomes show significantly greater crossovers in *asy1* (Fig. 2G and *SI Appendix*, Figs. S2 and S3 and Table S4), which we term the telomere-led zone (TLZ). Interestingly, the TLZ corresponds to distal regions that have relatively low ASY1 and REC8 ChIP-seq enrichment in wild type (Fig. 2G). At the fine scale, we observed that *asy1* crossovers show a preference to form in nucleosome-depleted, AT-rich regions with higher-than-average SPO11-1-oligos, which were similar to wild type crossovers (*SI Appendix*, Fig. S4) (25). Hence, although crossovers are highly distalized in *asy1* mutants, they retain a local bias toward chromatin and sequence features related to elevated DSB levels (25).

Crossovers Remodel toward Telomeric Regions in asy1/+ Heterozygotes.

We next sought to investigate whether asy1/+ heterozygotes associate with remodeled crossover frequency. We self-fertilized asy1-4/+ Col×Ler F₁ hybrids to generate 191 F₂ plants, which were then sequenced (*SI Appendix*, Table S3). These data were compared to crossovers previously mapped in a control Col×Ler wild type F₂ population (37). Crossovers per F₂ were not significantly different between wild type (mean = 7.54) and asy1-4/+ (mean = 7.92) populations (MWW test, P = 0.059; Fig. 3A and SI Appendix, Table S3). A positive correlation exists between number of crossovers per chromosome and physical chromosome length in both asy1/+ ($r = 0.97 P = 6.17 \times 10^{-3}$) and wild type ($r = 0.97 P = 4.83 \times 10^{-3}$) Col×Ler (Fig. 3B). Hence, global crossover numbers are maintained in asy1/+ heterozygotes, relative to wild type.

At the chromosome scale, despite crossover numbers being maintained, we observed that the asy1/+ recombination landscape was remodeled (Fig. 3 C and D). Specifically, crossovers increased in the distal subtelomeric regions in asy1/+ compared to wild type at the expense of the pericentromeric regions (Fig. 3 Cand D). The centromeres remained crossover-suppressed in both wild type and asy1/+ populations (Fig. 3 C and D). We repeated analysis of crossover positions relative to the nearest telomere and compared crossover counts between wild type and asy1/+ using χ^2 tests (Fig. 3E and SI Appendix, Table S4). This identified the first megabase within the subtelomeric regions as showing significantly elevated crossovers in asy1/+ compared to wild type (Fig. 3E and SI Appendix, Fig. S3 and Table S4), which overlaps with the TLZ observed in asy1 (Fig. 2G). As noted, the TLZ contains regions of relatively low ASY1 and REC8 ChIP-seq enrichment in wild type, which may explain the sensitivity of crossovers in these regions to reduced ASY1 gene dosage (Fig. 3E). These findings demonstrate that asyl/+ heterozygotes maintain crossover numbers, but show remodeling of recombination toward distal regions.

Crossovers Are Sensitive to ASY1 and ASY3 Gene Dosage, but Not **REC8.** To further investigate changes to crossover frequency associated with meiotic axis gene dosage, we used fluorescent tagged lines (FTLs) (38, 39). FTL intervals are defined by T-DNA insertions that express different colors of fluorescent protein (green, red, or blue) in pollen (LAT52 promoter) or seed (NapA promoter) (38, 39). When an individual is hemizygous for linked T-DNAs, patterns of fluorescence in pollen or seed can be used to quantify crossover frequency within the intervals flanked by the T-DNAs (40, 41). We crossed the subtelomeric FTL interval 420 on chromosome 3 to asy1/+, asy3/+, and rec8/+, using two independent alleles in each case (Fig. 3F and SI Appendix, Table S5). We observed that all asy1/+ and asy3/+ heterozygotes showed significantly increased 420 crossover frequency compared to wild type (*t* test *P* value range = 2.39×10^{-7} to 2.00×10^{-9} ; Fig. 3G and SI Appendix, Table S5). In contrast, rec8/+ heterozygotes showed no significant difference (t test P = 0.26; Fig. 3G and SI Appendix, Table S5). As 420 is located distally, we observed

a relatively high genetic distance in *asy1* and *asy3* (~10 cM; Fig. 3F and *SI Appendix*, Table S5) despite these backgrounds having reduced crossovers genome-wide (Fig. 2) (9, 11). It is not possible to measure FTLs in *rec8* homozygotes, as they are completely sterile (7, 42, 43).

Due to the remodeling of crossovers along the chromosomes observed in asy1/+ (Fig. 3C), we also measured recombination using the CEN3 FTL, which spans the DNA methylated centromere and pericentromere of chromosome 3 (Fig. 3F and SI Appendix, Table S6). CEN3 showed a significant decrease in crossover frequency in *asy1/+* and *asy3/+* heterozygotes compared to wild type (*t* test $P = 4.60 \times 10^{-9}$ and 2.05×10^{-6} ; Fig. 3G and SI Appendix, Table S6). No significant difference in CEN3 crossover frequency was observed between rec8/+ and wild type (t test P = 0.11; Fig. 3G and SI Appendix, Table S6). In asy1 and asy3 homozygotes, CEN3 crossovers were virtually eliminated compared to wild type (*t* test $P = 5.85 \times 10^{-11}$ and 1.40×10^{-8} ; Fig. 3G and SI Appendix, Table S6), consistent with telomere-led recombination dominating in these backgrounds. These experiments demonstrate remodeling of the crossover landscape toward the telomeres via reduced gene dosage of ASY1 and ASY3, but not REC8.

Cytogenetic Analysis of Meiosis in *asy1/+* and *asy3/+* Heterozygotes.

We next analyzed meiotic progression of asy1/+ and asy3/+heterozygotes using chromosome spreads of pollen mother cells and DAPI staining of chromatin (Fig. 4A). Chromosomes paired normally at pachytene in the asy1/+ and asy3/+ heterozygotes, and heterochromatic regions of dense DAPI staining were visible during prophase I (Fig. 4A). Five bivalents were detected at diakinesis, and no missegregation of chromosomes was observed at anaphase I or meiosis II in asy1/+ and asy3/+ (Fig. 4A). Consistently, no significant decrease in asy1/+ or asy3/+ seed count or pollen viability was observed (SI Appendix, Tables S7 and S8). To further assess chromatin organization, we immunostained male meiocytes for ASY1 and the heterochromatic histone modification H3K9me2 in wild type, asy1/+, asy1, asy3/+, and asy3 (Fig. 4B) (26, 44). H3K9me2 staining on chromosomes was observed in all genotypes, consistent with normal heterochromatin formation (Fig. 4B). ASY1 was undetectable in asy1, and showed an altered punctate staining pattern in asy3, as reported (Fig. 4B) (9).

To cytologically analyze Class I crossovers, we immunostained diakinesis-stage male meiocytes using α -MLH1 antibodies and stained chromatin with DAPI (Fig. 4*C* and *SI Appendix*, Fig. S5 and Table S9). We did not observe significant differences in total MLH1 foci between *asy1/+* and wild type (MWW test *P* = 0.128), but a small yet significant decrease occurred in *asy3/+* (MWW test *P* = 2.26 × 10⁻⁷; Fig. 4 *C* and *D* and *SI Appendix*, Table S9). We quantified MLH1 foci overlapping pericentromeric heterochromatin, defined by DAPI-dense regions, and observed a significant decrease in *asy1/+* and *asy3/+* compared to wild type (MWW test *P* = 4.53 × 10⁻⁵ and *P* = 8.94 × 10⁻⁴; Fig. 4 *C* and *D* and *SI Appendix*, Fig. S5 and Table S9). This is further consistent with distalization of crossovers away from centromere-proximal regions in *asy1/+* and *asy3/+* heterozygotes.

To investigate the effect of asy1/+ heterozygosity on axis loading of ASY1, we quantified α -ASY1 immunostained signal intensity during early prophase I and observed a 21% reduction in asy1/+ compared to wild type (MWW test P = 0.019; Fig. 4 D and E and SI Appendix, Table S10). As ASY3 is required for polymerization of ASY1 during meiosis (Fig. 4B) (9), we also quantified α -ASY1 signal intensity in asy3/+ and observed a 25% reduction compared to wild type (MWW test $P = 1.6 \times 10^{-3}$; Fig. 4 D and E and SI Appendix, Table S10). We immunostained pachytene-stage cells for the synaptonemal complex (SC) transverse filament ZYP1 (45). Continuous ZYP1 signal was observed along chromosomes in asy1/+ and asy3/+ compared to wild type,



Fig. 3. Distal increases in crossover frequency in *asy1/+* and *asy3/+* axis heterozygotes. (*A*) Histograms of crossovers per F_2 individual for wild type and *asy1/+*. Red dashed lines indicate mean values. (*B*) Crossovers per chromosome per F_2 for wild type (blue) and *asy1/+* (red) plotted against chromosome length in megabases. (*C*) Crossover frequency in wild type (blue) and *asy1/+* (red) and ASY1 ChIP-seq enrichment [log₂(ChIP/input), black] analyzed along proportionally scaled chromosome arms, oriented from telomeres to centromeres. Col×Ler SNP density is shown by green shading. (*D*) Crossover frequency (crossovers/150 kb per F_2) plotted along the *Arabidopsis* genome for wild type (blue) and *asy1/+* (red), with Col×Ler SNP density (green) shaded. Telomere and centromere positions are indicated by vertical solid and dotted lines, respectively. (*E*) Crossovers analyzed relative to the closest telomere in wild type (blue) and *asy1/+* (red). The lower plot shows ASY1 (green) and REC8-HA (pink) ChIP-seq enrichment [log₂(ChIP/input)] analyzed over the same regions. (*F*) DNA methylation (CG, CHG, CHH) in wild type (Col) is plotted along chromosome 3, and the positions of the *420* and *CEN3* FTL intervals are indicated. (G) Crossover frequency (in cM) within the *420* and *CEN3* FTL intervals in the indicated genotypes. Black dots represent replicate measurements, and red dots represent mean values. To assess significant differences, *t* tests were performed (n.s., not significant; ***P < 0.01).



Fig. 4. Cytological analysis of meiosis in asy1/+ and asy3/+ heterozygotes. (A) Spreads of wild type, asy1-4/+, and asy3-1/+ male meiocytes at the labeled stages of meiosis, with chromatin stained by DAPI. (B) Male meiocytes immunostained for ASY1 (red) and H3K9me2 (green) and stained for DNA (DAPI, blue) in the indicated genotypes. (C) Pollen mother cells immunostained for MLH1 (red) at diakinesis stage in wild type, asy1-4/+, and asy3-1/+. Chromatin was stained with DAPI (blue). The mean number of MLH1 foci and the subset of foci overlapping heterochromatin are printed for each genotype. (D) Quantification of MLH1 foci per meiocyte, ASY1 immunostaining signal, and ZYP1 immunostaining-derived synaptonemal complex (SC) length (in µM) per cell in the indicated genotypes. Statistical significance was assessed using MWW tests. (E) Male meiocytes immunostained for ASY1 (green) and ZYP1 (red) at early and late prophase I in wild type, asy1-4/+, and asy3-1/+. (Scale bars, 10 µM.)

and no significant differences in SC length were observed (MWW tests P = 0.74 and P = 0.27; Fig. 4 D and E and SI Appendix, Table S11). Hence, although we detect a reduction in ASY1 loading in asy1/+ and asy3/+ heterozygotes, full pairing and synapsis occurs in these backgrounds.

Crossover Interference Is Maintained in *asy1/+* **but Is Absent in** *asy1.* We next investigated crossover interference in *asy1/+* and *asy1* using analysis of double crossover (DCO) events. As we sequenced F_2 individuals, which derive from two independent meioses, in some cases, there is uncertainty about whether an observed DCO occurs in *cis* or *trans (SI Appendix,* Fig. S6) (46). Importantly, only *cis* DCOs, which occurred in the same meiosis, are relevant for measurement of crossover interference (46). In our F_2 data, a subset of *cis* DCOs can be identified from Col–Het–Col, Ler–Het–Ler, or Ws–Het–Ws genotype blocks (*SI Appendix*, Fig. S6) (46, 47). Therefore, we filtered for DCOs following this genotype pattern, which resulted in 118, 86, 98, and 73 DCOs in the Col×Ler wild type and *asy1*/+ and Col×Ws wild type and *asy1* populations, respectively (*SI Appendix*, Table S12). For each population, a matched set of randomly positioned DCOs of the same widths was generated as a control comparison. We analyzed *recq4a recq4b* crossover data in the same way (37), where the interference-insensitive Class II crossover repair pathway is greatly increased (37, 48). In *asy1*, the majority of the remaining crossovers have been shown to be dependent on the Class I pathway (9, 15).

Consistent with the action of crossover interference, distances between observed DCOs were significantly greater than random in both wild type Col×Ler (MWW $P = 5.67 \times 10^{-4}$) and Col×Ws (MWW test $P = 3.79 \times 10^{-4}$; Fig. 5 A and B and SI Appendix, Table S12). The *asy1/+* DCOs were also more widely spaced than random (MWW $P = 4.94 \times 10^{-9}$), which is consistent with normal crossover interference in this background (Fig. 5 A and B and SI Appendix, Table S12). In contrast, the spacing of DCOs in *asy1* was not significantly different from random (MWW P = 0.842; Fig. 5 A and B and SI Appendix, Table S12), showing an absence of detectable crossover interference. Using the same analysis method, DCOs observed in *recq4a recq4b* were not significantly different from random (P = 0.187), as expected due to greatly elevated noninterfering crossover repair (37, 48).

To independently measure crossover interference in wild type, asy1/+, and asy1, we used the distally located three-color FTL interval I3bc, which overlaps the 420 FTL interval on chromosome 3 (Fig. 5C, Table 1, and *SI Appendix*, Fig. S7). We measured crossover frequency between the I3bc T-DNAs in a *qrt1* background, where the four sister haploid cells produced from each male meiosis remain physically attached, allowing tetrad pollen analysis (Table 1) (38). To estimate crossover interference, we calculated I3b crossover frequency with and without a crossover in the adjacent I3c interval (Table 1). These measurements are used to calculate an interference ratio, where values close to 0 indicate stronger interference (38).

In wild type, the more distal interval I3b shows higher crossover frequency than I3c and an interference ratio of 0.34 (Fig. 5D and Table 1). In asy1/+, a significant increase in I3b crossover frequency occurred compared to wild type (Perkins $P = 5.20 \times 10^{-9}$), whereas *I3c* was not significantly changed (Fig. 5D and Table 1). Consistent with our previous observations, no significant difference was observed in the interference ratio between wild type and asy1/+ (Perkins P = 0.997; Fig. 5D and Table 1). In asy1, both genetic intervals showed a significant reduction in crossover frequency compared to wild type (I3b Perkins $P = 3.95 \times 10^{-49}$ and *I3c* Perkins $P = 1.06 \times 10^{-5}$), although the more distal *I3b* interval maintains a higher level of crossovers than I3c (Fig. 5D and Table 1). In contrast to asyl/+, the asyl homozygotes showed an interference ratio of 1.24 that was significantly different than wild type (Perkins $P = 7.20 \times 10^{-6}$; Fig. 5D and Table 1), further consistent with an absence of crossover interference.

Finally, to investigate Class I crossovers in wild type and asy1, we immunostained MLH1 at diakinesis stage and stained DNA with DAPI (Fig. 5E and SI Appendix, Tables S9 and S13). In wild type (Col), no univalent chromosomes are observed, and, on average, 10.4 MLH1 foci occurred on the bivalents (Fig. 5E and SI Appendix, Tables S9 and S13). In asy1, we observed a higher incidence of univalent chromosomes per cell (mean = 5.6) compared to bivalents (mean = 2.6; Fig. 5E and SI Appendix, Table S13). Interestingly, we observed MLH1 foci on both univalents (mean = 5.3) and bivalents (mean = 5) in *asy1* (Fig. 5E and SI Appendix, Table S13). The presence of MLH1 foci on asy1 bivalents is consistent with crossover formation via the Class I pathway. MLH1 foci have been reported on univalent chromosomes in *dmc1* and in haploid meiosis and may represent sites of intersister repair (49). In order to estimate crossover interference, we measured MLH1 interfoci distances on bivalents in wild type and asy1 (SI Appendix, Table S14). We observed that MLH1 foci were significantly closer in asyl compared to wild type (MWW test $P = 2.19 \times 10^{-6}$; SI Appendix, Table S14), which is further consistent with a loss of crossover interference. Therefore, our combined analysis of DCO spacing from sequencing data, fluorescent pollen tetrads, and MLH1 foci show that crossover interference is absent in asy1.

Discussion

Our data inform a model for how ASY1 and the meiotic axis pattern crossover frequency along plant chromosomes (Fig. 6). Previous work has shown that *asy1* mutants undergo normal

telomere clustering, formation of meiotic DSB foci during early leptotene, and polymerization of an axial structure marked by REC8 and ASY3 (Fig. 64) (9, 15, 31). However, DMC1 foci dynamics are altered in asy1, resulting in a failure of interhomolog recombination and depletion of crossovers (9, 15). Using highresolution mapping of crossovers via sequencing F₂ plants, we show that recombination becomes largely restricted to a telomere-led zone (TLZ) in asyl homozygotes (Fig. 6B). We propose that the proximity of telomeres during early prophase in asy1 is responsible for telomere-led recombination (Fig. 6A) (31). Telomere-led recombination is active in wild type, but ASY1 antagonizes this activity to promote crossover formation in interstitial and centromereproximal chromosome regions (Fig. 6B). Using ChIP-seq, we observe a gradient of ASY1 enrichment from the telomeres to the centromeres, which is paralleled by REC8 cohesin enrichment (30). We propose that differential ASY1 enrichment represents a mechanism to distribute recombination more evenly along the chromosome arms. However, as heterochromatin increases in proximity to the centromere, this causes suppression of meiotic DSBs and crossovers, despite high levels of ASY1 and REC8 (25, 26, 50).

We show that plants heterozygous for asy1/+ and asy3/+ mutations undergo remodeling of the crossover landscape, with a shift toward the distal subtelomeres, at the expense of interstitial and pericentromeric regions. Interestingly, the distal regions that undergo crossover increases in asy1/+ overlap the TLZ observed in asy1 and have relatively low levels of ASY1 and REC8 ChIP-seq enrichment in wild type. Using meiotic immunocytology, we quantified a ~21% reduction in ASY1 loading on chromatin in asy1/+. This could indicate a threshold effect over which ASY1 antagonizes telomere-led recombination and promotes crossovers in the chromosome arms, toward the centromere. In asyl/+ heterozygotes, the distal regions would drop below this putative threshold and the strength of telomere-led recombination would increase. As interference remains operational in asy1/+, this would lead to a relative loss of crossovers in the interstitial and pericentromeric regions (Fig. 6B). Alternatively, this may reflect a nonlinear effect of decreased ASY1 expression on recombination along the chromosomes. It is notable that genetic variation in axis components, including ASY1 and ASY3, has been strongly associated with adaptation to tetraploidy in Arabidopsis arenosa, which may include distalization of crossovers (51, 52). Our results show that gene dosage of ASY1 and ASY3 may contribute to these effects, in addition to the influence of specific variants on protein function (51, 52).

Crossover interference is mediated by topoisomerase II and the axis protein Red1 in budding yeast (53), while the SC component SYP-1 has been implicated in Caenorhabditis elegans (54, 55). Using analysis of double crossover distances, MLH1 foci, and fluorescent pollen tetrads, we show that ASY1 is required for detectable crossover interference in Arabidopsis. Interestingly, crossovers in axis mutants are largely dependent on the Class I interfering repair pathway (9, 15). For example, chiasma are eliminated in asy3 msh4 double mutants, and we show that MLH1 foci form on asy1 bivalents (9, 15). Therefore, despite the Class I pathway mediating the majority of crossover formation in asy1, interference signaling between recombination sites is inactive. Crossover interference has been proposed to occur via mechanical stress acting across paired homologous chromosomes, which is transmitted along the axis and relieved at crossover designated sites (13, 53). In this respect, ASY1 may mediate crossover interference via transmission of mechanical stress, when chromatin loops connected to the axis undergo cycles of expansion and contraction during early prophase I (13). In the absence of ASY1, the mechanical properties of the axis may be altered, meaning force can no longer be transmitted and crossover interference is not detected. Alternatively, ASY1 may control sensitivity of interhomolog repair sites to the interference signal or mediate transmission of a biochemical signal along the chromosome axis (56). Our work indicates that axis HORMA domain proteins can play a critical role in mediating crossover interference along chromosomes during meiosis.



Fig. 5. Crossover interference is maintained in *asy1/+* but is absent in *asy1*. (A) Histograms showing the distribution of observed double crossover distances (DCOs, red) in megabases in wild type, *recq4a recq4b* (37), and *asy1/+* Col×Ler F_2 individuals or wild type and *asy1* Col×Ws F_2 individuals. Alongside are identical histograms showing the distribution of matched randomly generated distances (blue). Mann–Whitney–Wilcoxon (MWW) tests were performed to assess significant differences between observed DCOs and random, with *P* values indicated. (*B*) Diagrams showing spacing of identified DCOs along the proportional physical length of chromosomes (as percentages). DCOs are connected via arcs and color-coded proportional to the distance between them (red, greatest; blue, smallest). (C) DNA methylation (CG, CHG, CHH) in wild type plotted along chromosome 3 with the positions of the *I3bc* FTL T-DNAs indicated by vertical lines and colored triangles. (*D*) Crossover frequency (in cM) within *I3b*, contingent on crossover in the adjacent interval *I3c*, in wild type, *asy1*, and *asy1*. Interference ratios calculated from the *I3bc* data are plotted for the same genotypes. (*E*) Representative images of pollen mother cells immunostained for MLH1 (red) at diakinesis stage in wild type and *asy1*. Chromatin was stained with DAPI (blue). White and orange arrows in the merged images indicate MLH1 foci located on bivalents or univalents, respectively. (Scale bars, 10 μ M.)

Table 1.	Pollen tetrad analysis of crossover frequency and
interferer	nce within the <i>I3bc</i> FTL intervals in wild type (Col),
asy1-4/+,	and asy1-4

Wild type					
Tetrad class	(Col)	asy1-4/+	asy1-4		
A – NCO	2,338	2,092	913		
B – SCO <i>I3c</i>	328	317	46		
C – SCO <i>I3b</i>	1,313	1,574	113		
D – SCO I3b and SCO I3c	12	16	6		
E – SCO <i>I3b</i> and SCO <i>I3c</i>	8	17	1		
F – SCO <i>I3b</i> and SCO <i>I3c</i>	13	10	0		
G – SCO I3b and SCO I3c	14	13	1		
H – DCO <i>l3c</i>	3	2	2		
I – DCO <i>I3b</i>	5	8	5		
J – DCO <i>I3c</i> and SCO <i>I3b</i>	0	0	1		
K – SCO <i>I3c</i> and DCO <i>I3b</i>	0	0	1		
L – DCO I3c and DCO I3b	0	0	0		
Total	4,034	4,049	1,089		
Genetic distance					
<i>I3b</i> cM (<i>P</i> value)	17.23	20.72 (5.20 × 10 ⁻⁹)	7.02 (3.95 × 10 ⁻⁴⁹)		
I3c cM (P value)	4.87	4.75 (0.895)	3.35 (1.06 × 10 ⁻⁵)		
<i>I3b</i> cM without adjacent <i>I3c</i> CO	18.37	22.07	6.94		
<i>I3b</i> cM with adjacent <i>I3c</i> CO	6.22	7.47	8.26		
Interference ratio	0.34	0.34	1.24		
P value	_	0.997	$7.20 imes10^{-6}$		

Fluorescent pollen tetrads were classified into one of 12 color patterns: non-crossover (A – NCO), single *I3c* crossover (B – SCO *I3c*), single *I3b* crossover (C – SCO *I3b*), two-strand *I3bc* double crossovers (D – SCO *I3b* and SCO *I3c*), three-strand double crossovers type 1 (E – SCO *I3b* and SCO *I3c*), three-strand double crossovers type 2 (F – SCO *I3b* and SCO *I3c*), four-strand double crossovers (G – SCO *I3b* and SCO *I3c*), four-strand double crossovers (G – SCO *I3b* and SCO *I3c*), four-strand double crossovers (G – SCO *I3b* and SCO *I3c*), four-strand double crossovers within interval *I3c* (H – DCO *I3c*), four-strand double crossovers within interval *I3b* (I – DCO *I3b*), double *I3c* crossover and single *I3b* crossovers (K – SCO *I3c* and DCO *I3b*), and double *I3c* crossovers and double *I3b* crossovers (L – DCO *I3c* and DCO *I3b*). Recombination frequency was measured using the Perkins method (38). Interference ratios and statistical tests were calculated using the Malkova method as described (38). The locations of the *I3bc* FTL T-DNAs are 498,916 (CFP), 3,126,994 (YFP), and 4,319,513 (dsRed2) bp on chromosome 3 (40).

Materials and Methods

Plant Materials. Arabidopsis plants were grown under long day conditions (16 h light/8 h dark) at 20 °C. The following mutant alleles in the Col-0 background were used: *rec8-1* (Salk_091193) (9), *rec8-3* (SAIL_807_B08) (49), *asy1-1* (Salk_144182) (15), *asy1-4* (Salk_046272) (23), *asy3-1* (Salk_143676) (9), and *asy3-2* (SAIL_423_H01) (9). The *asy1-3* allele is in the Ws-4 background (32). The *REC8-HA rec8* line was as reported (30).

ASY1 Chromatin Immunoprecipitation and Sequencing. ASY1 ChIP was performed using 8 g of floral buds or leaf tissue. Nuclei isolation and chromatin recovery were performed as described (25, 57). Chromatin was sheared using a Bioruptor instrument (Diagenode) for 15 min at high power alternating 30 s on and 30 s off, followed by 15 min at high power alternating 30 s on and 1 min off. Chromatin immunoprecipitation was performed using an α -ASY1 antibody (12), or the preimmune serum, at a dilution of 1/160. DNA purification, DNA library preparation, and sequencing were performed as described (25).

ChIP-Seq Data Analysis. Deduplicated paired-end ASY1, REC8-HA, H3K9me2, H3K4me1, H3K4me2, H3K4me3, H3K27me1, and H3K27me3 ChIP-seq reads, paired-end MNase-seq reads, and single-end SPO11-1-oligo, H2A.Z, and H2A.W reads (25, 30, 58, 59) were aligned to the TAIR10 reference genome using Bowtie2 (version 2.2.9) (60) with the following settings: -very sensitive -p 4 -k 10. For paired-end reads, the Bowtie2 options –no-discordant and –no-mixed were also applied. Prior to alignment, single-end SPO11-1-oligo reads were processed as described (25). Up to 10 valid alignments were reported for each

read or read pair. Aligned reads with more than two mismatches were discarded using the SAM optional field "XM:i." Uniquely aligning reads were extracted by removing alignments with the SAM optional field "XS:i" and with Bowtie2-assigned MAPQ scores lower than 42. Alignments consisting of reads that mapped to multiple loci were filtered such that only those with MAPQ scores higher than or equal to 10 remained, from which the alignment with the highest MAPQ score was retained. Where MAPQ scores for multiple valid alignments were equal, one alignment was randomly selected. Alignments consisting of only one read in a pair were discarded. Unique and multiple alignments in BAM format were combined, and coverage was calculated for each coordinate in the genome using Rsamtools (version 1.26.1). Coverage was normalized by the sum of coverage for each library. The log₂ ratio of Chl-P:input coverage was calculated to control for background and variation in mappability across genomic loci. A library for Columbia genomic DNA (gDNA) that had been extracted using CTAB, fragmented using dsDNA shearase, and subjected to paired-end sequencing on an Illumina NextSEq 500 instrument as described (25, 58) was aligned to TAIR10 and used to calculate log₂(MNase/ gDNA) ratios. Additionally, the first gDNA read in each pair was trimmed to 50 bp, aligned, and used to calculate log₂(SPO11-1-oligo/gDNA) ratios.

To generate chromosome-scale profiles, mean coverage values within adjacent 10-kb windows were calculated. Log₂ ratios of windowed ChIP:input coverage were then calculated and smoothed by applying a moving average. Additionally, DNA methylation proportions derived from published bisulfite sequencing reads were used to profile DNA methylation levels at the chromosome scale (61). Spearman's rank-order correlation coefficients were calculated for each pair of profiled data sets and presented in correlation matrices separately for the chromosome arms and pericentromeres. The pericentromeres are defined as the regions surrounding the centromeres with higher-than-average DNA methylation (26). For analysis along telomere–centromere axes, data values were first calculated in 10-kb windows along the chromosomes. Chromosome arms were then oriented such that each began at the telomere and ended at the centromere and divided into windows along their proportional lengths. Data values were then averaged across all chromosome arms and plotted.

Fine-scale coverage profiles around TAIR10 representative gene transcription start and termination sites (TSSs and TTSs) were generated using the normalizeToMatrix function from the Bioconductor package Enriched-Heatmap (version 1.11.1) (62). Each feature was divided into proportionally scaled windows between start and end coordinates, and 2-kb flanking regions were divided into 20-bp windows. For each window along each feature and its flanking regions, an average value was calculated using the "w0" method for ChIP-seq data. The default profile-smoothing method implemented in the normalizeToMatrix function was applied. The resulting matrix of windowed coverage values was used to generate a mean profile, or a heat map in which each row represents a single feature. Mean profiles and heat maps were plotted such that the distance between feature start and end coordinates along the *x*-axis represents the mean feature length.

Crossover Mapping via Genotyping by Sequencing. Wild type and *asy1* Col×Ws and *asy1/+* Col×Ler F₂ plants were grown, and genomic DNA was extracted from leaf tissue using a CTAB protocol as described (37, 50, 63). A total of 150 ng of DNA was used to generate each sequencing library as described (37, 50, 63). A total of 96 libraries were pooled and sequenced on one lane of an Illumina NextSeq500 instrument using a 300-cycle Mid-Output kit (Illumina). Sequencing data analysis and mapping of crossovers were carried out using the TIGER pipeline as described (33, 37, 50, 63).

Measuring Crossover Frequency Using Fluorescent FTL Pollen and Seed. Scoring of fluorescent seeds and measurement of crossover frequency within the 420 genetic interval were performed by microscopy and using CellProfiler as described (39, 64, 65). Scoring of fluorescent pollen grains and measurement of crossover frequency within the *CEN3* FTL genetic interval were performed using an Accuri C2 (BD Biosciences) flow cytometer as described (66). For measurement of crossover interference within the *I3bc* FTL intervals, *qrt1* pollen tetrads were scored using a Leica SP8 confocal microscope. Calculation of crossover frequency and the interference ratio were performed as described (38).

Cytological Analysis of Meiosis. Fixation of *Arabidopsis* inflorescences and chromosome spreads of pollen mother cells (PMCs) were performed as described (67). Immunostaining of ASY1, ZYP1, and MLH1 were prepared on acetic acid chromosome spreads using fixed inflorescences. After chromosome spreading, the slides were incubated in boiling 10-mM Tris-sodium citrate, pH 7.0, for 45 s, followed by incubation in 1× PBS with 0.1% Triton X-100 (PBST) for 5 min. Primary antibodies were diluted in a solution of 1% BSA diluted in PBST that was added onto the slides, followed by incubation for 20 h at 4 °C for ASY1 and ZYP1 immunostaining or 40 h at 4 °C for MLH1 immunostaining. The



Fig. 6. ASY1 acts as a gene dosage-sensitive antagonist of telomere-led recombination and mediates crossover interference. (A) A single pair of homologs is represented at the G2, leptotene, and pachytene stages of meiosis in wild type, asy1/+, and asy1. Centromeres are represented as red dots, telomeres as blue dots; the nucleolus is a pale blue circle, and crossover positions are indicated by black triangles. Telomere-led alignment of homologs is shown with black bars connecting two chromosomes at leptotene stage. (B) A single aligned pair of homologs is shown, with the positions of crossovers indicated by crosses in wild type, asy1/+, and asy1. Crossovers are promoted in proximity to the telomere-led zone (TLZ, blue). ASY1 (purple) loading antagonizes the TLZ and promotes interstitial and proximal crossovers via interference. Note that this model represents metacentric chromosomes such as chromosomes 1, 3, and 5. The acrocentric chromosomes 2 and 4, which bear nucleolar organizing regions (NORs) on their short arms, may differ in crossover patterning.

slides were washed in PBST three times for 5 min each at room temperature. Following this, a solution of secondary antibodies diluted in PBST was added, and the slides were incubated for 30 min at 37 °C. The slides were washed in PBST three times for 5 min each at room temperature, and a solution of DAPI/Vectashield was added and a coverslip added to the slide before imaging. The following antibodies were used for immunostaining: α-ASY1 (rat, 1/500 dilution) (11), α-ZYP1 (rabbit, 1/ 500 dilution) (45), and α-MLH1 (rabbit, 1/200 dilution) (68). Coimmunostaining of ASY1 and H3K9me2 was performed using fresh floral buds. Inflorescences were dissected on damp filter paper under a stereo microscope, and six buds at floral stages 8 to 9 (69) were isolated and transferred to 5 µL of enzyme digestion solution (0.4% cytohelicase, 1.5% sucrose, 1% polyvinylpyrrolidone) on a microscope slide. The buds were dissected to recover the anthers, while the rest of the bud tissue was discarded. The slide was then incubated in a moist box at 37 °C for 1 min, and the anthers were gently opened with a brass rod to release the meiocytes. A total of 5 μ L of enzyme digestion solution was added, and the slide was incubated in a moist box at 37 °C for 2 min. After this, 10 µL of 1% Lipsol was added, and the solution was gently mixed with a needle for 1 min before adding 20 μL of 4% paraformaldehyde. The slides were then left to dry for 4 h. Incubation of slides with antibodies for immunostaining of proteins was performed as described earlier. The following antibodies were used: α-ASY1 (rabbit, 1/500 dilution) and H3K9me2 (mouse, 1/100 dilution; Abcam ab1220).

Microscopy was conducted using a DeltaVision Personal DV microscope (Applied Precision/GE Healthcare) equipped with a CCD CoolSNAP HQ2 camera (Photometrics). Image capture was performed using SoftWoRx software version 5.5 (Applied Precision/GE Healthcare). To analyze colocalization of ASY1 and REC8 immunostaining signal on meiotic cells, the contour of chromatin (stained with DAPI) was marked and signal intensity was quantified for every pixel within the marked area using the package coloc2 from Fiji. Following MLH1 immunostaining of diakinesis cells, heterochromatin was identified and marked based on brighter DAPI signal using Fiji (*SI Appendix*,

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Fig. S5). MLH1 foci were then compared with the marked heterochromatic regions to score overlaps. Synapsed chromosomes immunostained for ZYP1 were marked along their length using the Line Selection Tool of Fiji. Pixel length was recorded and then converted into μ M using the Setting Measurement Scale of Fiji. For quantification of ASY1 signal intensity, all slides were prepared alongside one another, and images were captured using the same exposure time. The contour of each cell was marked, and the intensity within this region measured. Each cell was captured as *Z*-stack of 10 optical sections of 0.2 μ M each, and the maximum intensity projection was reconstructed using ImageJ as described (23, 26). A region adjacent to the cell was also marked, and the intensity was measured and used as mean background intensity to subtract from the within cell intensity.

Data Availability Statement. All data are publically available. ASY1 ChIP-seq library data have been deposited in the ArrayExpress database at EMBL-EBI (http://www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-8705 (70). Sequencing data for wild type and *asy1* Col×Ws GBS libraries have been deposited under ArrayExpress accession E-MTAB-8715 (71), and data for *asy1*/+ Col×Ler GBS libraries has been deposited under ArrayExpress accession E-MTAB-8725 (72).

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