



Natural variation identifies SNI1, the SMC5/6 component, as a modifier of meiotic crossover in *Arabidopsis*

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The frequency and distribution of meiotic crossovers are tightly controlled; however, variation in this process can be observed both within and between species. Using crosses of two natural *Arabidopsis thaliana* accessions, Col and Ler, we mapped a crossover modifier locus to semidominant polymorphisms in *SUPPRESSOR OF NPR1-1 INDUCIBLE 1 (SNI1)*, which encodes a component of the SMC5/6 complex. The *sni1* mutant exhibits a modified pattern of recombination across the genome with crossovers elevated in chromosome distal regions but reduced in pericentromeres. Mutations in *SNI1* result in reduced crossover interference and can partially restore the fertility of a Class I crossover pathway mutant, which suggests that the protein affects noninterfering crossover repair. Therefore, we tested genetic interactions between *SNI1* and both *RECQ4* and *FANCM* DNA helicases, which showed that additional Class II crossovers observed in the *sni1* mutant are *FANCM* independent. Furthermore, genetic analysis of other SMC5/6 mutants confirms the observations of crossover redistribution made for *SNI1*. The study reveals the importance of the SMC5/6 complex in ensuring the proper progress of meiotic recombination in plants.

meiosis | crossover | SNI1 | SMC5/6 | *Arabidopsis*

Sexual reproduction is widespread in eukaryotes and promotes genetic variation within the populations. At the heart of this process is the meiotic cell division, in which crossover recombination shuffles genetic information between homologous chromosomes (1). Meiotic recombination initiates from the programmed DNA double-strand breaks (DSBs), which, following resection, form single-stranded DNA filaments (2, 3). These filaments can invade the complementary DNA sequences of a homologous chromosome, leading to the formation of interhomolog joint molecules (JMs) (3). During meiosis, the “ZMM” proteins stabilize interhomolog JMs and promote Class I crossover repair (2, 3), which is the major crossover pathway in most species. The Class I pathway exhibits interference, meaning that crossovers are placed further apart on the chromosome than expected at random (2–4). As an alternative to crossover repair, JMs can be repaired as non-crossovers (NCOs), which is promoted via DNA helicases including *BLM/SGS1/RECQ4* (2, 3). In many eukaryotes, the number of DSBs exceeds the final number of crossovers, and DNA helicases play a key role in limiting crossovers by disassembly of meiotic recombination intermediates and promoting their repair as NCOs (5–7). For example, *Arabidopsis thaliana recq4a recq4b* loss-of-function mutants show a twofold crossover increase due to a failure in NCO formation (7, 8). In budding yeast, the STRUCTURAL MAINTENANCE OF THE CHROMOSOME 5/6 (SMC5/6) complex plays a dual role in DNA helicase control by providing SUMO activity via its NSE2/MMS21 subunit, which stimulates the anti-recombination function of *RECQ4*, and also by limiting

heteroduplex extension, thus reducing helicase substrates (9, 10). JMs may also be repaired via structure-specific endonucleases, including *MUS81*, which can result in NCOs or Class II non-interfering crossovers (2–4).

The frequency and chromosomal distribution of crossovers are tightly controlled, as high crossover numbers may break down favorable combinations of alleles, potentially reducing fitness (11). However, significant variation in crossover frequency is also observed between individuals, sexes, populations, and species (12–14). Genetic factors that modify crossover levels or patterns have the potential to profoundly influence selective responses during adaptation (13, 15, 16). For example, high variation across the major histocompatibility complex, which encodes cell surface proteins essential for the human adaptive immune system, is achieved by meiotic recombination hotspots within the locus (17, 18). Similarly, crossover hotspots were observed in a subset of plant *NLR* resistance genes (19).

Significance

Meiotic recombination plays a fundamental role in shaping genetic diversity in eukaryotes. Extensive variation in crossover rate exists between populations and species. The identity of modifier loci and their roles in genome evolution remain incompletely understood. We explored natural variation in *Arabidopsis* crossover and identified *SNI1* as the causal gene underlying a major modifier locus. To date, *SNI1* had no known role in crossover. *SNI1* is a component of the SMC5/6 complex that is closely related to cohesin and condensin. *Arabidopsis sne1* and other SMC5/6 mutants show similar effects on the interference-independent crossover pathway. Hence, our findings demonstrate that the SMC5/6 complex, which is known for its role in DNA damage repair, is also important for control of meiotic crossover.

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Genetic loci that act in trans to control variation in meiotic recombination levels or patterns have been described in plants and animals. For example, the PRDM9 zinc finger KRAB-SET domain protein has been shown to recruit DSB machinery to specific DNA sequence motifs at recombination hotspot in humans and mice (20, 21). Genetic variation in PRDM9 governs crossover hotspot location within and between vertebrate species (22–24). A sequence polymorphism in E3 SUMO and ubiquitin ligases related to human RNF212 and HEI10/CCNB1IP1 has been identified that controls crossover frequency in vertebrates, invertebrates, fungi, and plants (25–31). HEI10 in plants acts in the Class I pathway and is a dosage-sensitive regulator of crossover number (30). Variation in the meiosis-specific α -kleisin cohesin subunit REC8 influences the crossover rate in some mammals and plants, which acts to tether chromatin loops to the chromosome axis during prophase I (27, 28, 32, 33). Genetic

variation in *Arabidopsis* also identified TAF4b as a recombination modifier locus, which encodes a general transcription factor paralog expressed specifically during meiosis (34). In this study, we further investigated variation in crossover frequency in *A. thaliana* and identified SNI1, a component of the SMC5/6 complex, as a factor preventing errors in meiotic recombination in plants.

Results

Genetic Mapping of the Col/Ler Recombination Quantitative Trait Locus rQTL4. Segregation of linked reporters expressing fluorescent proteins in seed or pollen (fluorescent-tagged lines, FTLs) can be used to measure crossover frequency within defined chromosomal intervals (35–38). In our previous work, we used a Col-420×LLCLL F₂ population (in which chromosomes 1, 2, 4, and 5 were Col/Ler segregating) to map quantitative trait loci

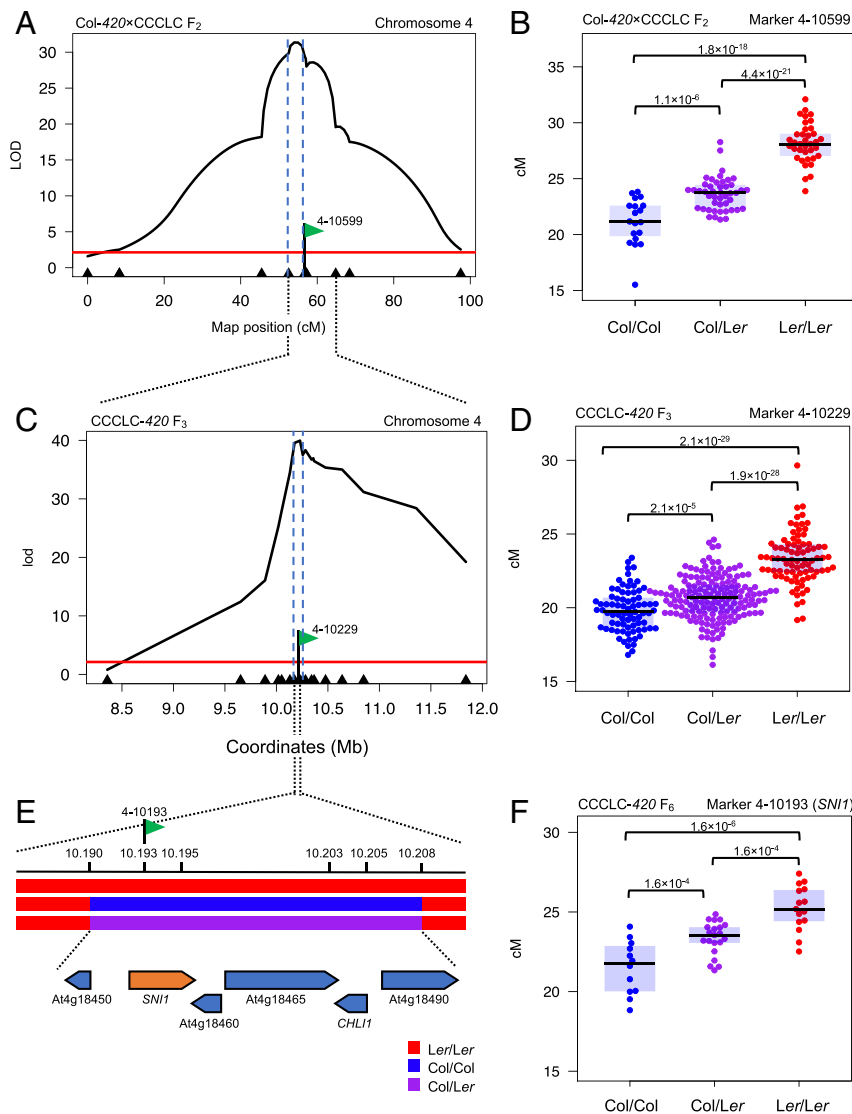


Fig. 1. Fine-mapping of the *rQTL4* recombination modifier locus. (A) LOD scores for markers associated with 420 crossover frequency from *Col-420*×*CCCLC* F₂ population. The x axis corresponds to the chromosome 4 genetic map with markers used for mapping indicated by arrowheads. The red line indicates the 95% significance threshold, and blue dashed lines mark the credible interval. The marker 4-10599 is labeled with a green flag. (B) Effect plots showing 420 crossover frequency for *Col/Col*, *Col/Ler*, and *Ler/Ler* individuals for the marker 4-10599 in the F₂ population. Each dot represents one individual. *P* values were estimated by Welch *t* test. (C) As in A but showing the *Col-420*×*CCCLC* F₃ population. The x axis corresponds to chromosome 4 physical positions (Mb). The marker 4-10229 is labeled with a green flag. (D) As in B but for the marker 4-10229 in the F₃ population. (E) The genotypes of progeny of the H-27 recombinant plant, which has only a 19.5-kb region segregating for *Col/Ler* and the remainder of the genome being fixed. Gene models are shown beneath the genotypes. The marker 4-10193 is labeled with a green flag. (F) As in B but for the marker 4-10193 in the progeny of the H-27 F₅ recombinant.

responsible for variation in crossover frequency between the *Arabidopsis* Col and *Ler* accessions (30, 39). We identified two significant *rQTLs* located on chromosomes 1 and 4, exhibiting logarithm of the (base 10) odds ratio (LOD) scores of 40.1 and 53.5, which explained 23.3% and 33.6% of the variance in crossover frequency, respectively (30). *rQTL1* was identified as *HEI10*, encoding an E3 ubiquitin/SUMO ligase acting in the Class I crossover pathway (30). In this study, we sought to identify the causal locus for *rQTL4*.

We constructed a Col-420×CCCLC (39) F₂ population ($n = 102$) and genotyped all individuals for 11 markers spanning the previously identified *rQTL4* region (30). This revealed a strong *rQTL* association between markers 4-9652 and 4-10366 with an LOD score of 31.72 (Fig. 1A and *SI Appendix, Table S1*). *rQTL4* is semi-dominant with the Col allele associating with reduced 420 crossover frequency (marker 4-10599; Fig. 1B). We selected a single F₂ individual heterozygous over *rQTL4* and fixed for *rQTL1* and generated an F₃ population ($n = 2,280$), which was genotyped at markers 4-8358, 4-10847, and 4-11840 to identify recombinants within the *rQTL4* interval. We identified 325 recombinants, which were measured for 420 crossover frequency and genotyped using 16 markers, which narrowed the credible interval to a 53-kb region containing 26 genes (Fig. 1C and D and *SI Appendix, Table S2*). We developed an F₄ population from an *rQTL4* heterozygous F₃ individual ($n = 152$) (*SI Appendix, Fig. S1A and Table S3*) and genotyped using four internal markers. This allowed us to identify one individual fixed for *Ler* upstream of the interval, which showed 420 crossover frequency equivalent to the Col/*Ler* heterozygous *rQTL4* level (mean = 21.31 cM). We used the progeny of this individual to generate a population of 1,056 F₅ and genotyped them to find recombinants within the *rQTL4* interval. We identified an individual (H-27) that was Col/*Ler* heterozygous for a 19.5 kb encompassing six genes, with the remainder of the region fixed for one parent (Fig. 1E). The progeny of this plant ($n = 48$) showed variation in recombination frequency similar to that observed in the original Col-420×CCCLC F₂ population (Fig. 1F and *SI Appendix, Table S4*). Genotyping within the 19.5-kb interval showed that plants fixed for Col allele present a 420 crossover frequency at 21.52 cM, whereas heterozygotes showed 23.35 cM and *Ler/Ler* homozygotes 25.23 cM on average. Mean 420 crossover frequency was statistically different between the groups, with the greatest difference between the Col/Col and *Ler/Ler* groups (Welch *t* test; $P = 1.57 \times 10^{-6}$). Therefore, we conclude that *rQTL4* corresponds to one of the six identified genes in the 19.5-kb interval.

Variation in *SNII* Underlies *rQTL4*. To determine which of the candidate genes corresponds to *rQTL4*, we performed an allelism test by crossing transfer-DNA mutants in all six genes with F₆ individuals fixed for Col/Col and *Ler/Ler* over the *rQTL4* interval. Mutation of the *rQTL4* causal gene should result in a significant difference in recombination frequency of plants obtained from crosses with F₆^{Col/Col} and F₆^{Ler/Ler} plants. We observed that only a mutant in *SNII* (At4g18470) showed a significant difference in 420 crossover frequency between the two crosses, strongly suggesting that this gene corresponds to *rQTL4* (Fig. 2A and B and *SI Appendix, Table S5*). In addition, *SNII* was the only candidate gene which showed nonsynonymous substitutions in the coding sequence when Col and *Ler* accessions were compared (Fig. 2B).

We measured 420 crossover frequency in independently isolated homozygous mutations in the candidate genes (Fig. 2C). This assay was not possible for *CHL11* (At4g18480), which has a very strong dwarf mutant phenotype (40), and for At4g18490 due to the FTL reporter silencing. From the four remaining genes, a *snii-1* loss-of-function allele (41–43) showed a significant increase in crossovers (29.41 cM, $n = 12$) when compared to the wild type (21.58 cM, $n = 8$) (Fig. 2C and *SI Appendix, Table S6*).

We next performed a complementation experiment in which Col and *Ler* *SNII* alleles were transformed into the *snii-1* mutant background. This showed that both Col and *Ler* *SNII* variants can complement the *snii-1* elevated crossover phenotype, demonstrating that the effect on 420 crossover rate in *snii-1* is due to the nonsense mutation at the *SNII* locus and not a secondary mutation (Fig. 2D and *SI Appendix, Table S7*). We observed that the *SNII*^{Ler} transformants were higher recombining than *SNII*^{Col} transformants, although the difference was not statistically significant.

Next, we measured whether the Col and *Ler* *SNII* alleles lead to differences in messenger RNA (mRNA) expression by performing qRT-PCR from closed flower buds, comparing *rQTL4* F₆ individuals (the progeny of individual H-27). We did not observe a significant difference in expression between the Col and *Ler* *SNII* alleles, indicating that the variation observed most likely does not act via mRNA expression level (*SI Appendix, Fig. S1 B and C and Tables S8 and S9*). Consistently, transformation of additional *SNII* copies did not change 420 crossover rate (*SI Appendix, Fig. S1D and Table S10*).

We investigated how the two alleles might modify recombination rate in different environmental conditions. To this end, we tested the effect of both alleles on recombination at different temperatures, since temperature influences the frequency of recombination (44, 45). We took advantage of the fact that F₆^{Col} and F₆^{Ler} differ only in the 19.5-kb interval containing *SNII*. We observed that the 420 crossover frequency was less responsive to thermal conditions in F₆ carrying *SNII*^{Ler} than in F₆ *SNII*^{Col} plants (Fig. 2E and *SI Appendix, Table S11*). This suggests that variability within the *SNII* gene may be important in modulating meiotic recombination under varying environmental conditions.

Comparison of the Col and *Ler* *SNII* alleles revealed the existence of two nonsynonymous substitutions: L142F and I235V. The first of these substitutions is also present in Ct-1 accession, which was previously used in crosses with Col-420 to map *rQTL* (*SI Appendix, Fig. S2A*) (37). As Ct-1×Col-420 mapping population did not reveal *rQTL* located on chromosome 4, we concluded that the second substitution, I235V, is likely responsible for the *SNII* recombination phenotype. The V235 (*Ler*) variant exists in 379 out of 1,135 *A. thaliana* accessions (46). The I235 (Col) allele was considered as ancestral, as it is conserved across Iberian relict accessions and in *SNII* homologs of *Arabidopsis lyrata*, *Arabis alpina*, and *Camelina sativa* (Fig. 2F and *SI Appendix, Fig. S3*). In general, *SNII* shows high sequence variation across *A. thaliana* accessions, with more than 30 different alleles and 59 nonsynonymous substitution sites (Fig. 2G and *SI Appendix, Fig. S2 B and C*) (42).

Mutation of *SNII* Causes Remodeling of the Meiotic Crossover Landscape. To further investigate crossover frequency in the mutant, we crossed *snii-1* with five different FTL reporter lines positioned in varying chromosomal locations [Col Traffic Lines, CTLs (38)]. In all cases, we observed significant changes in *snii-1* crossover rates, though the extent and direction of those changes varied according to the region (Fig. 3A and B and *SI Appendix, Table S12*). Specifically, we observed crossover increases ranging from 22.6% (interval CTL1.18) to 50.4% (interval CTL3.4) in distal chromosomal regions (Fig. 3A and B). Interestingly, we observed a 25.7% reduction in crossover frequency, from 17.5 to 13.0 cM, in the pericentromeric interval CTL3.9 (Fig. 3A and B). These observations are consistent with a global redistribution of crossover frequency in *snii-1*.

To explore the impact of *snii-1* on crossovers genome-wide, we used genotyping-by-sequencing (GBS) of F₂ individuals. To this end, we backcrossed (6×) *snii-1* into the *Ler* accession and produced Col^{snii-1}×*Ler*^{snii-1} F₁ hybrids. We also used CRISPR-Cas9 mutagenesis to generate a *snii-2* allele de novo in the Ct accession (*SI Appendix, Fig. S2 C and D*) and produced

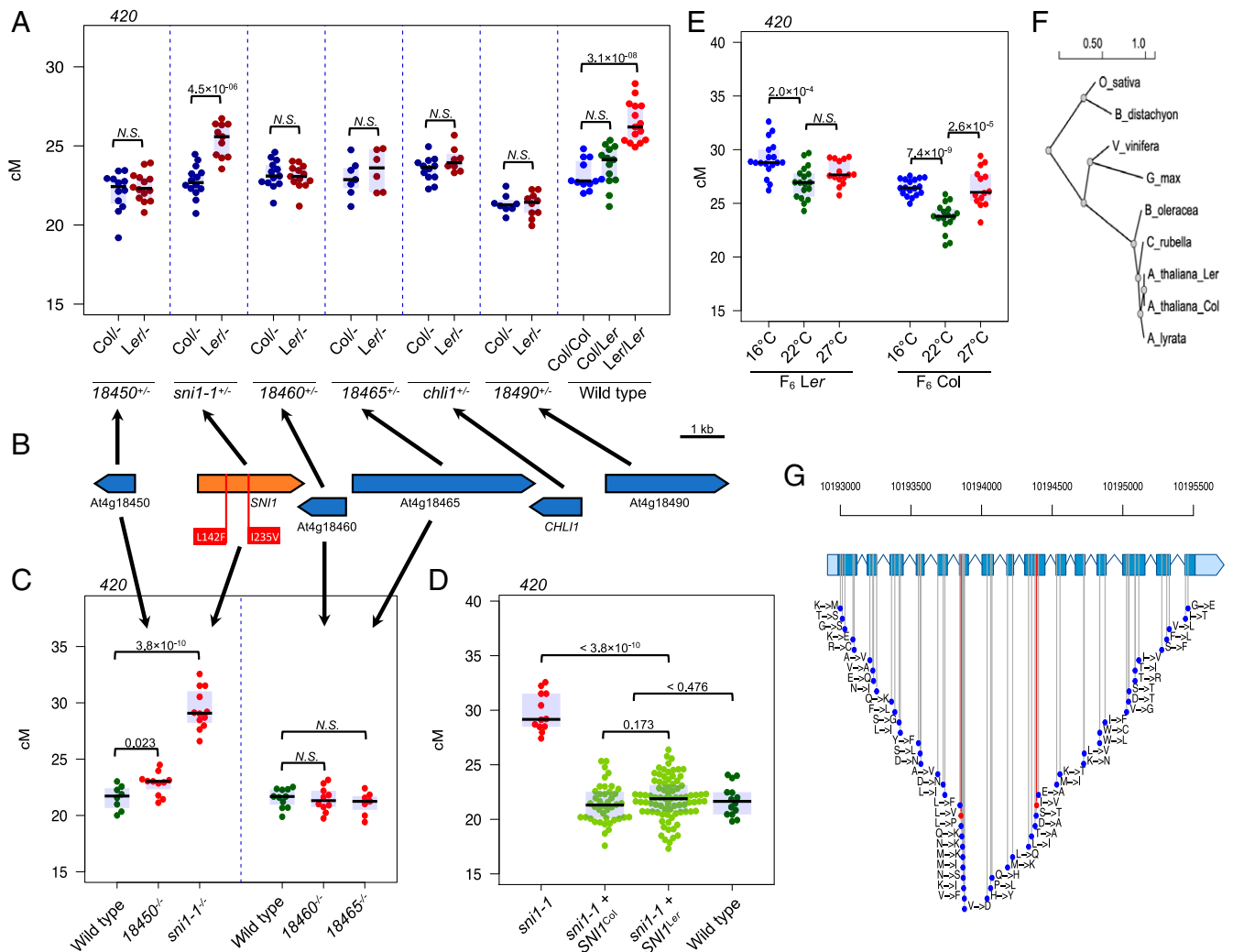


Fig. 2. *SNI1* corresponds to *rQTL4*. (A) 420 crossover frequency for F₁ plants obtained from the cross between mutant lines for the six genes and F₆ individuals fixed for Col/Col and Ler/Ler over the *rQTL4* interval. Measurements for the wild-type F₆ individuals for Col/Col, Col/Ler, and Ler/Ler were used as a control. Each dot represents one individual in A, C, D, and E. Significance in A, C, and D was assessed by Welch *t* test. (B) The genes within the *rQTL4* 19.5-kb Col/Ler segregating interval with non-synonymous substitutions indicated. (C) 420 crossover frequency for the mutants of genes located within *rQTL4* locus. (D) Complementation of the *sni1* mutation with *SNI1^{Col}* and *SNI1^{Ler}* alleles restores wild-type 420 crossover frequency in T₁ generation. (E) F₆ lines carrying *SNI1^{Ler}* are less responsive to temperature stress than the *SNI1^{Col}* as measured in the 420 interval. (G) Location of all non-synonymous substitution sites in the *SNI1* gene observed within 1,135 *A. thaliana* accessions (46). The two substitutions differing Col and Ler alleles are indicated in red. (F) Phylogenetic tree of *SNI1* orthologs identified by basic local alignment search tool (BLAST) searches in selected eukaryotic genomes. The phylogenetic tree was constructed using maximum likelihood method.

Col^{*sni1-1*} × Ct^{*sni1-2*} F₁ hybrids. 420 crossover frequency was elevated in both hybrids in the *sni1* background compared to wild-type controls (53.2% for Col × Ler; 35.4% for Col × Ct), indicating that the effect is not sensitive to interhomolog polymorphisms (Fig. 3C and SI Appendix, Tables S13 and S14). Interestingly, Col × Ler F₁ plants in which only the *SNI1^{Ler}* allele was active showed significantly higher 420 crossovers than wild-type hybrids (15.98 cM versus 13.50 cM), which is consistent with *Ler SNI1* increasing crossover repair. We sequenced genomic DNA from 174 and 229 F₂ individuals from each population and identified 1,194 and 2,260 crossovers per Col^{*sni1-1*} × Ler^{*sni1-1*} and Col^{*sni1-1*} × Ct^{*sni1-2*} population, respectively, using the TIGER pipeline (47). As a control, we compared to crossovers mapped in wild-type F₂ populations generated from Col × Ler and Col × Ct F₁ hybrids (8, 48). To eliminate potential effects of *sni1-1* introgression in the Col^{*sni1-1*} × Ler^{*sni1-1*}, chromosome 4 was removed from the analysis. Total crossover number per individual was

slightly higher in *sni1* than in wild type (9.55 versus 8.18 for Col × Ct, Mann–Whitney–Wilcoxon test $P = 1.41 \times 10^{-10}$ and 6.86 versus 6.25 crossovers per F₂ for Col × Ler, $P = 6.03 \times 10^{-3}$; Fig. 3D). We observed a significant change in the recombination landscape in the *sni1* mutants for both hybrids, with elevated crossovers in the chromosome arms and less in the pericentromeres (Fig. 3 E–G).

To exclude a possibility that the crossover estimates are affected by aneuploidy we tested for evidence of copy number variation in our GBS sequencing data. We calculated sequence coverage separately for each chromosome in 241 Col^{*sni1-1*} × Ct^{*sni1-2*} and 305 Col × Ct F₂ plants. None of the tested samples showed a coverage deviation larger than 10% from the total sample coverage, consistent with minimal aneuploidy present in these populations (SI Appendix, Table S15).

We compared Col × Ler and Col × Ct crossover maps using 500-kb windows and observed significant correlation between

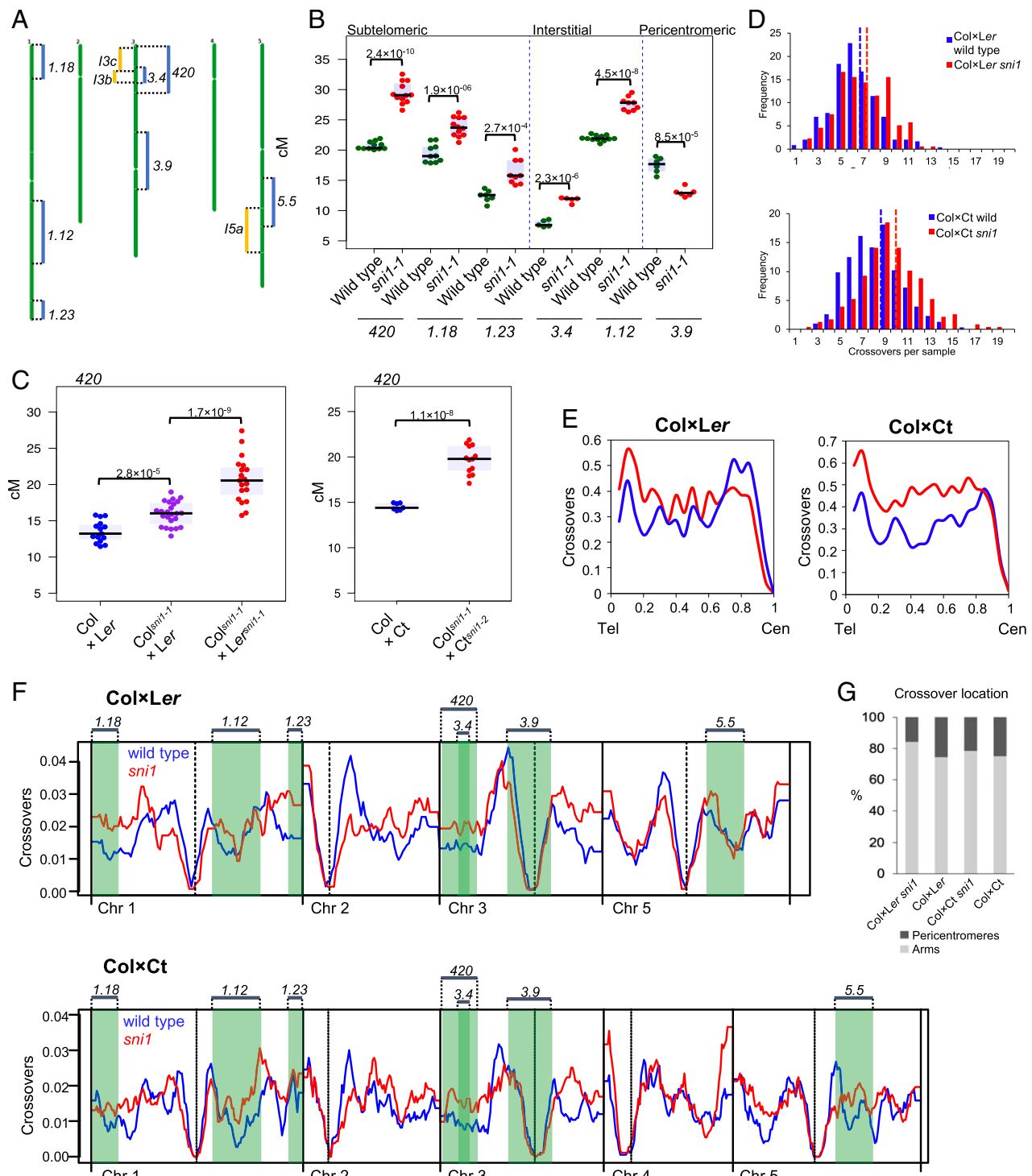


Fig. 3. The *sni1* mutants exhibit elevated crossover levels in chromosome arms and subtelomeric regions but reductions in pericentromeres. (A) Ideogram of *A. thaliana* chromosomes showing the positions of fluorescent reporter intervals for seed- (blue) and pollen-based (yellow) systems. (B) Crossover frequency in the seed-based intervals as measured in wild type (green) and *sni1-1* (red) F₂ siblings. (C) 420 crossover frequency (cM) in Col×Ler, Col^{sni1-1}×Ler, and Col^{sni1-1}×Ler^{sni1-1} F₁ hybrids (Left) and Col×Ct and Col^{sni1-1}×Ct^{sni1-2} F₁ hybrids (Right). Significance in B and C was assessed by Welch t tests; each dot represents one individual. (D) Histograms presenting the number of crossovers per individual in Col×Ler (blue) and Col^{sni1-1}×Ler^{sni1-1} (red) (Top) and Col×Ct (blue) and Col^{sni1-1}×Ct^{sni1-2} (red) (Bottom) F₂ GBS populations. The mean crossover number of each population is denoted by a vertical dashed line. (E) Crossover frequency along the proportional (scaled) length of the chromosomes from telomeres (Tel) to centromeres (Cen) in Col×Ler (blue) and Col^{sni1-1}×Ler^{sni1-1} (red) F₂ populations (Left) and Col×Ct (blue) and Col^{sni1-1}×Ct^{sni1-2} (red) (Right). Chromosome 4 in Col×Ler populations was excluded from the analyses in D through F to avoid potential bias from the 587-kb Col^{sni1-1} introgressed region in the Ler^{sni1-1} parent. (F) Crossover frequency over the chromosomes in Col×Ler (blue) and Col^{sni1-1}×Ler^{sni1-1} (red) F₂ populations (Top) and Col×Ct (blue) and Col^{sni1-1}×Ct^{sni1-2} (red) (Bottom) F₂ populations. Crossovers were tallied into 300-kb windows, divided by the number of F₂ individuals, and a rolling mean plotted along the chromosomes. Cen positions are denoted by vertical dashed lines and TEL positions by vertical solid lines. (G) Crossover location in wild type and *sni1* F₂ GBS populations as divided into chromosome arms and pericentromeres. The pericentromeres were defined as regions with higher-than-average DNA methylation, which surround the centromeres (88).

crossover distributions (Spearman Rho = 0.564; *SI Appendix, Fig. S4*). When the wild-type maps were compared to *sni1*, lower correlations were observed (Rho = 0.508 and Rho = 0.339 for Col×*Ler* and Col×Ct, respectively; *SI Appendix, Fig. S4*). Similarly, Col^{*sni1-1*}×*Ler*^{*sni1-1*} and Col^{*sni1*}×Ct^{*sni1-2*} maps showed low correlation (Rho = 0.355). Therefore, differences in crossover distribution between both hybrids are exacerbated in the *sni1* background.

Mutation of *SNI1* Results in Reduced Crossover Interference. We investigated crossover frequency in *sni1-1* using cytological approaches. DAPI staining and fluorescent in situ hybridization (FISH) of metaphase I chromosome spreads were used to count chiasmata on individual chromosome pairs (49). This indicated a small but significant reduction in chiasma numbers in the *sni1-1* mutant compared to the wild type (10.2 versus 8.9; Fig. 4*A* and *B* and *SI Appendix, Tables S16* and *S17*), which contrasts with the small overall increase observed via sequencing (Fig. 3*D*). This may be due to closely spaced crossovers being underestimated by chiasmata counts. Also, we analyzed Class I crossovers via MLH1 immunostaining on pachytene-stage male meiocytes (50) (Fig. 4*C* and *D* and *SI Appendix, Table S17*). While 7.4 MLH1 foci were detected in Col-420 line ($n = 50$), 6.2 foci were observed in *sni1-1* ($n = 44$), showing a significant reduction. This suggests that Class I/Class II crossover ratio is changed in *sni1-1* and interfering Class I crossovers decrease. Therefore, we tested whether crossover interference is affected in *sni1-1*, by crossing

sni1-1 with the three color FTL *I3bc* line (51). This allowed us to measure male-specific crossover frequency in two adjacent intervals, *I3b* and *I3c*, to measure the number of double crossovers (DCOs) and calculate interference (35, 36). We observed a significant reduction in crossover interference, measured as a coexistence of coincident (CoC), in the *sni1-1* mutant when compared to wild type (Fig. 4*E–G* and *SI Appendix, Table S18*).

For our Col×Ct and Col^{*sni1-1*}×Ct^{*sni1-2*} F₂ GBS data, we have sufficient crossover events (2,478 and 2,260, respectively) to perform cis-DCO analysis by filtering for parental–heterozygous–parental genotype transitions (52, 53) and measure DCO distances. We compared the distribution of DCO distances between the wild type and *sni1* within each F₂ population (Fig. 4*H*) and observed a significantly closer DCOs in *sni1* (Kolmogorov–Smirnov test, $P = 0.028$). Along with the observed decrease in MLH1 foci number, the reduction in crossover interference may be apparent and result from a change in the balance of Class I versus Class II crossover repair in *sni1* mutants.

Meiotic Recombination Phenotypes of *sni1* Are Likely Independent of Systemic Acquired Resistance. *SNI1* was identified in a genetic screen for regulators of the salicylic acid (SA)–mediated defense response (41), and *sni1* mutations have been found to suppress systemic acquired resistance (SAR) (42, 43). To test whether the effect of *sni1* on meiotic recombination is linked to its role in SAR, we investigated how SA, the major trigger in SAR, affects

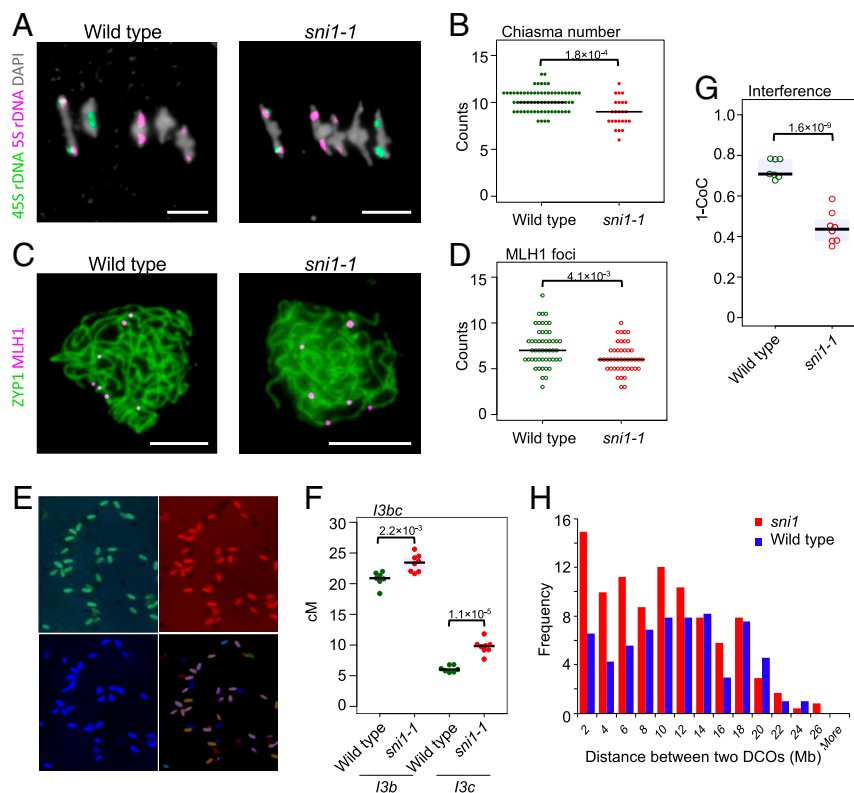


Fig. 4. The *sni1-1* mutant shows weakening of crossover interference. (A) Representative images of DAPI-stained bivalents at metaphase I in wild type (Col) and *sni1-1*. FISH probes against 45S (green) and 5S (red) ribosomal DNA were used to identify chromosomes, allowing for chiasma counting depending on chromosome morphology. (Scale bar, 5 μm.) (B) Quantification of chiasma count data from wild type and *sni1-1*. Significance was assessed by Mann–Whitney–Wilcoxon (MWW) tests. (C) Representative images of pachytene stage male meiocytes from wild type (Col-420) or *sni1-1* stained for MLH1 (red) and ZYP1 (green). (Scale bar, 5 μm.) (D) Quantification of MLH1 foci on pachytene stage meiotic chromosomes in wild type and *sni1-1*. Significance was assessed by MWW tests. (E) Microphotographs of segregating *I3bc* pollen grains as seen in three fluorescent channels. Composite image was also shown. (F) *I3b* and *I3c* genetic distances in wild type and *sni1-1*. Each dot represents measurements from four to 10 pooled individuals. Significance was assessed by Welch *t* tests. (G) Crossover interference (1-CoC) between the *I3b* and *I3c* intervals. (H) Histograms of cis-DCO distances in Col×Ct (blue) and Col^{*sni1-1*}×Ct^{*sni1-2*} (red) calculated in 2-Mb windows. Frequency was scaled to the number of F₂ individuals in each population.

crossover frequency. To this end, we treated flowering Col^{sni1-1-420} and wild-type Col-420 with SA by spraying flowering plants (1 mM of SA solution on a weekly basis). This treatment induced *PR1* expression, indicating successful triggering of SAR (SI Appendix, Fig. S5 A and B and Tables S19 and S20). We did not observe any change in 420 crossover frequency, either in wild type or *sni1-1*. We measured crossover frequency in *I5a* (Col background) and *LTL5.5* (*Ler* background) chromosome intervals in SA-treated plants and also did not observe any change when compared to nontreated control (SI Appendix, Fig. S5 C and D and Tables S21 and S22). Altogether, our data indicate that SA does not affect meiotic recombination in these regions.

Mutations in some DNA repair genes partially suppress the retarded growth of *sni1-1* (42, 54). To see their effect on the meiotic phenotype of *sni1*, we created the double mutants *sni1-1 rad51*, *sni1-1 brca2a*, *sni1-1 brca2b*, and *sni1-1 atr* in the background of the reporter line 420. The *sni1-1* mutation was not able to suppress the sterility phenotype of *rad51* (SI Appendix, Fig. S6A) (however, see ref. 55). We measured 420 crossover frequency in *sni1-1 brca2a* and *sni1-1 brca2b* and observed no suppression of recombination (SI Appendix, Fig. S6B and Table S23). Surprisingly, we observed a further increase of 420 recombination frequencies in the *sni1-1 atr* mutant, which together with no improvement in the fertility of the double mutant compared to *sni1-1* suggests that the meiotic recombination phenotype of *sni1-1* is independent of ATR (SI Appendix, Fig. S6 C and D and Tables S24 and S35).

Recent studies have shown that the immunity-related phenotypes of *sni1* are dependent on *EDS1* (56). In the double-homozygous mutant *sni1 eds1-2*, plant growth is partially restored and transcript accumulation of *PR1* is abrogated (56). Therefore, we crossed the *sni1-1-420* line with *eds1-2* to test whether the meiotic phenotype of *sni1* are dependent on *EDS1*. 420 crossover frequency was not changed in *eds1-2* background when compared to wild type, and *sni1-1 eds1-2* double-homozygous mutants were not statistically different from the *sni1-1* single mutant (SI Appendix, Fig. S5E and Table S25). This indicates that unlike immunity-related phenotypes, the meiotic recombination phenotype of *sni1* is independent of *EDS1*.

Meiotic Recombination Phenotypes of *sni1*. Recent studies have revealed that SNI1 is a component of the SMC5/6 complex and is a homolog of NSE6 (Fig. 5A) (42). The role of SMC5/6 in timely resolution of meiotic recombination intermediates has been described in budding yeast (9, 10, 57). Consistently, we observed anaphase I bridges, chromosome fragments, and micronuclei formation in *sni1-1* meiocytes (Fig. 5B), which was also reported for *Arabidopsis nse4* and *nse2* mutants (58, 59). A single chromosome fragment was observed in one metaphase I cell (out of 24), while all of the analyzed anaphase I cells ($n = 15$) showed some degree of chromosome fragmentation. Dyads were formed in at least 10% of the meiocytes, which led to the production of unreduced gametes (Fig. 5B), which is also typical for SMC5/6 mutants (59). We confirmed these abnormalities also in the *sni1-2* mutant allele (SI Appendix, Fig. S7). Concurrently with the submission of this work, another *sni1* allele has been characterized and displayed a similar but stronger phenotype (55). Unreduced gametes may potentially lead to the formation of polyploid seeds, which in turn would affect the crossover frequency measurements. However, only 3.7% ($n = 2,292$) pollen grains were unreduced in *sni1-1* (SI Appendix, Table S26), and we did not observe larger seeds in *sni1-1* progeny, which is an indicator of polyploidy. Moreover, seed-based crossover measurements repeatedly returned 3:1 Mendelian segregation ratios of fluorescent reporters (green to non-green and red to non-red seeds), as expected for the diploid plants.

We investigated the formation of meiotic DSBs in the *sni1-1* mutant by scoring RAD51 foci at early prophase I chromosomes

and did not observe a significant change when compared to wild type (Fig. 5 C and D and SI Appendix, Table S27). To ensure that *sni1-1* does not produce SPO11-independent DSBs that can be repaired by crossovers, we created a *sni1-1 spo11-1* double mutant. This double mutant was sterile, showing that *sni1-1* mutation is not able to restore fertility in *spo11-1* (SI Appendix, Fig. S8). In prophase I, we found no cells at pachytene stage neither in *spo11-1* ($n = 300$) nor *sni1-1 spo11-1* ($n = 300$) (SI Appendix, Fig. S8), which indicates that there is no synapsis in these mutants. Consequently, 10 univalents were observed for both *spo11-1* and *sni1-1 spo11-1* at metaphase I (SI Appendix, Fig. S8 and Table S28). In two out of 43 metaphase I cells, we observed single bivalents in *sni1-1 spo11*, which, however, did not differ significantly from the results reported for the single *spo11-1* mutant (60). Interestingly, chromosome fragments were detected in 18.6% of *sni1-1 spo11-1* cells ($n = 43$), while this was not observed in *spo11-1*. As immunostaining with RAD51 antibodies did not show an increase in the number of DSBs in the early stages of *sni1-1* mutant prophase (Fig. 5 C and D), these chromosomal fragments likely arose as a consequence of chromosome condensation problems.

Together, these results indicate that DSB formation proceeds normally in *sni1-1*, and the mutant does not produce SPO11-independent DSBs that could be repaired as crossovers. In addition, we analyzed the synaptonemal complex component ZYP1 (ZIP1 homolog) and meiotic cohesin SYN1 (REC8 homolog) in wild-type and *sni1-1* (Fig. 5E). This revealed that synapsis proceeds normally in the *sni1-1* mutant without apparent cohesion failures.

Mutants of Other Components of the SMC5/6 Complex Show Similar Changes in Crossover as *sni1-1*. We next measured crossover frequency in mutants of other SMC5/6 complex components and compared them to *sni1-1* and wild type. In *A. thaliana*, homologs of all SMC5/6 complex subunits have been identified and are either embryo lethal or show strongly disturbed development which results in partial or complete sterility (42, 61–63). *SMC6* is encoded by two genes in the *Arabidopsis* genome, *SMC6A* and *SMC6B*, which are partially functionally redundant (62). *NSE4* is encoded by two functionally non-redundant copies, *NSE4A* and *NSE4B*, in which expression of *NSE4B* is significantly lower than *NSE4A*, and *nse4a* single mutants show fertility defects (63). We crossed mutants of *NSE4A*, *ASAP1* (homolog of *NSE5*), and a combination of mutants for *SMC6A* and *SMC6B* subunits with the Col-420 reporter line to assess crossover frequency. Crossovers were dramatically elevated in the *nse4a* mutant (30.71 cM; Fig. 5F and SI Appendix, Table S29). As the *asap1* is almost sterile and *smc6a smc6b* double mutants are lethal in the homozygous state, which precludes measurement of recombination, we tested the effect of their decreased expression. *ASAP1/asap1* heterozygotes showed a significant crossover increase from 21.06 to 23.22 cM (Fig. 5F and SI Appendix, Table S29). Out of the *smc6* mutants tested, *smc6a* did not show any effect on crossover frequency, while *smc6b* (62) showed significant increases (to 24.13 cM). We also constructed *smc6a smc6b* sesquimutants (homozygous for one mutation and heterozygous for the other) and observed similar increases in 420 crossover frequency as the single *smc6b* mutant (22.16 cM and 23.35 cM, respectively). Thus, mutants of SMC5/6 components show a consistent change in crossover frequency with that observed in the *sni1-1* mutant.

Besides *SNI1*, mutants of *NSE2* are also viable in *A. thaliana* and produce enough seeds to assess crossover frequency using FTL reporters. *NSE2* is located on chromosome 3, within the 420 interval; therefore, it was not possible to test recombination using this FTL. Instead, we crossed *nse2-2* with *CTL1.23* and *CTL3.9*, which represent subtelomeric and pericentromeric intervals, respectively (Fig. 3A). The *nse2* mutant shows significant changes in both intervals when compared to the wild type

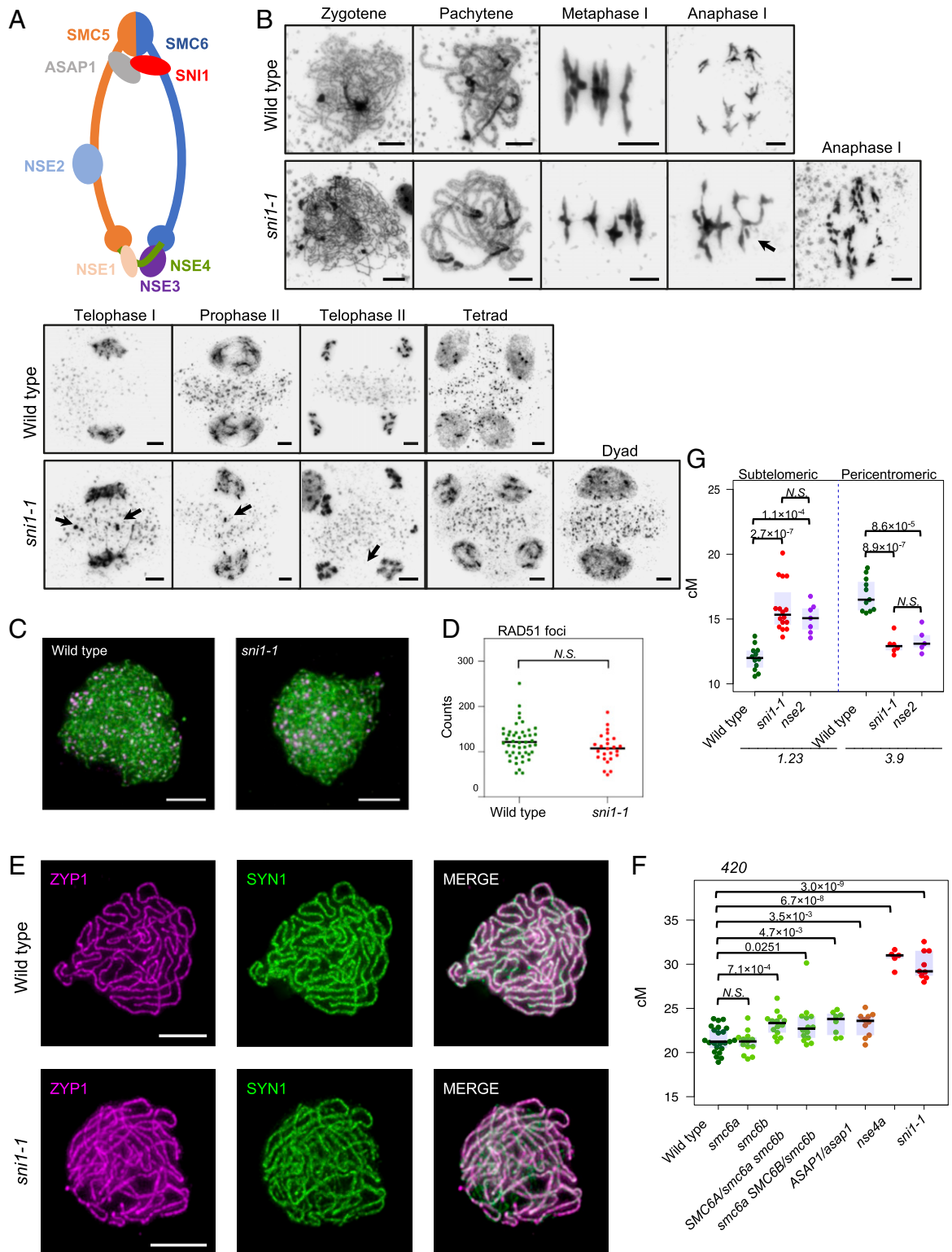


Fig. 5. The meiotic phenotype of the *sni1* mutant is similar to SMC5/6 complex mutants. (A) Schematic representation of SMC5/6 complex. The interface of the protein interaction of the SNI1-ASAP1 subcomplex (gray-red) is unknown in plants. (B) Cytological characterization of the *sni1* mutant in comparison to wild type (Col). The stages of meiotic progression were labeled. Chromosome fragments and micronuclei observed in the *sni1* mutant are indicated by arrows. (Scale bar, 5 μ m.) (C) Representative images of ASY1 (green) and RAD51 (magenta) coimmunostaining on wild type (Col-420) and *sni1-1* male meiocytes at zygonema. (D) Quantification of RAD51 foci number per cell in wild type and *sni1-1*. (E) Representative images of ZYP1 (magenta) and SYN1 (green) coimmunostaining on wild type (Col-420) and *sni1-1* male meiocytes at pachynema. (Scale bar, 5 μ m.) (F) 420 crossover frequency in SMC5/6 complex mutants. (G) *CTL1.23* and *CTL3.9* genetic distances (cM) in wild type, *sni1-1*, and *nse2-2*. Significance in D through F was assessed by Welch *t* test; each dot represents one individual.

(Fig. 5G and *SI Appendix, Table S30*) (*CTL1.23*: wild-type 12.0 cM, *nse2* 15.1 cM; *CTL3.9*: wild-type 16.9 cM, *nse2* 13.4 cM). The *nse2* changes had the same direction and magnitude as observed in *sni1-1* (*CTL1.23*: *sni1-1* 15.9 cM; *CTL3.9*: *sni1-1* 13.0 cM). Altogether, our results are consistent with a common role

of SMC5/6 complex in ensuring proper progress of meiotic crossover.

We also generated double *sni1-1 nse2* and *sni1-1 nse4a* mutants, which both showed severe developmental abnormalities and did not flower (*SI Appendix, Fig. S9*). This indicates that the

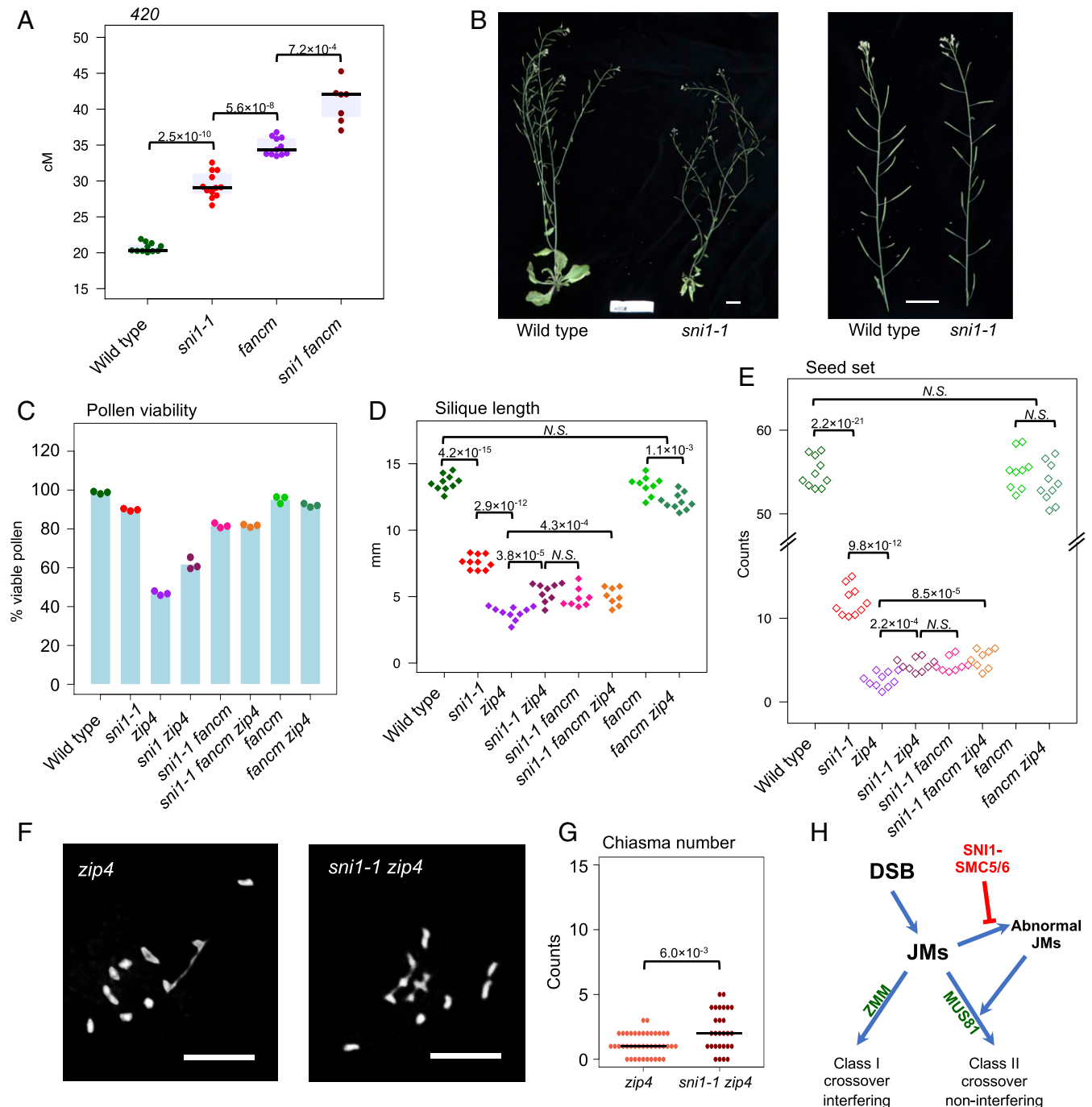


Fig. 6. SNI1 affects Class II crossover repair in FANCM-independent manner. (A) 420 crossover frequency in wild type, *sni1*, *fancm*, and *sni1 fancm* plants. Significance was assessed by Welch *t* test; each dot represents one individual. (B) Reduced size and fertility of *sni1* mutant plant in comparison with wild type. Whole plants (Left) and primary inflorescences (Right) are shown. (Scale bar, 2 cm.) (C–E) Fertility assays in *sni1*, *zip4*, and *fancm* mutants and their combinations as assessed via pollen viability (C), silique length (D), and seed set (E). Significance was assessed by Welch *t* test. (F) Representative images of DAPI-stained chromosomes at metaphase I in *zip4* and *sni1 zip4*. (Scale bar, 5 μ m.) (G) Quantification of chiasma count data from *zip4* and *sni1 zip4*. Significance was assessed by Mann–Whitney–Wilcoxon tests. (H) Model of SNI1 role in meiotic crossover formation. SNI1, together with SMC5/6 complex, prevents formation of abnormal JMs, which require MUS81 endonuclease for their successful repair.

lack of several components of the SMC5/6 complex leads to loss of its function.

Modification of the Crossover Landscape in *sni1-1* Is Independent of *FANCM*. In budding yeast, the *smc5 sgs1* and *nse4 sgs1* double mutants are synthetically sick (10). Similarly, when we combined *sni1* with *recq4a recq4b* (orthologous to *sgs1*), the triple-mutant plants died early after germination (*SI Appendix, Fig. S10A*), and the same was observed for *sni1 RECQ4A/recq4a recq4b* mutant. This suggests that RECQ4 helicases are crucial for somatic DNA repair during replication in the absence of SNI1. As the *sni1-1 recq4a recq4b* triple mutant never enters flowering stage, we were not able to assess crossover frequency. The lethality of *sni1 recq4a recq4b* triple mutants resembles the synthetic lethality of *mus81 recq4a recq4b* triple mutants (7, 64). We therefore crossed *sni1-1* with *mus81* and observed that the resulting double mutant is also synthetically lethal (*SI Appendix, Fig. S10B*). We propose that SMC5/6-SNI1 prevents formation of complex JMs, which if unrepaired in the absence of RECQ4 or MUS81 cause severe defects in DNA replication and cell division.

To test the genetic interactions with *FANCM*, we crossed *fancm* mutants with *sni1-1*. The resulting *sni1-1 fancm* double mutant did not suppress the developmental phenotypes of *sni1-1* (*SI Appendix, Fig. S10A*). In *A. thaliana* inbred lines, the null *fancm* mutation leads to a dramatic increase in crossover frequency via the Class II pathway (65). We compared 420 crossover recombination in *sni1-1* and *fancm* single mutants to the double mutant and observed an additive effect of the two mutations which manifests in further elevation of crossover rate from 29.41 cM in *sni1-1* and 34.78 cM in *fancm* to 40.93 cM in *sni1-1 fancm* (Fig. 6A and *SI Appendix, Table S31*). This shows that the increase in crossovers observed in *sni1-1* mutants is likely not due to mis-regulation of *FANCM*.

The *sni1-1* Mutation Partially Restores Crossover Formation to *zip4* Mutants via Elevated Class II Crossovers. If the increase in 420 crossover frequency in *sni1-1* is due to a higher number of Class II crossovers, then we would expect *sni1-1* to partially restore fertility to Class I pathway mutants (2). Therefore, we crossed *sni1-1* with the *fancm zip4* double mutant and analyzed pollen viability, silique length, and seed set in the resulting double- and triple-mutant lines (Fig. 6 C–E and *SI Appendix, Fig. S10 C and D* and Tables S32–S35). For all the traits analyzed, *sni1-1* exhibits reduced values compared to wild type, showing that the fertility is decreased in this mutant (Fig. 6 B–E and *SI Appendix, Fig. S10 C and D*). However, the double *sni1-1 zip4* mutant showed significantly higher fertility than the *zip4* single mutant, supporting the hypothesis that crossover elevation observed in the *sni1-1* mutant results from additional Class II events (Fig. 6 C–E and *SI Appendix, Tables S32–S35*). Interestingly, we found that the *fancm* mutation is not able to completely restore fertility in the *zip4* mutant when combined with the *sni1-1* mutation (Fig. 6 C–E and *SI Appendix, Fig. S10 C and D*). This is likely the result of other meiotic abnormalities observed in the absence of SNI1, including anaphase I bridges and chromosome fragments (Fig. 5B). Also, the *sni1-1 fancm* double mutant showed similar fertility to *sni1-1 fancm zip4*, which is much lower than that of the wild-type *fancm zip4* but also *sni1-1* (Fig. 6 C–E). To confirm that the partial restoration of fertility in *sni1-1 zip4* is due to higher crossover numbers, we counted bivalents and chiasmata in metaphase I. While the average number of bivalents per cell in *zip4* was 1.2, we observed 2.2 bivalents in *sni1-1 zip4* (Fig. 6 F and G and *SI Appendix, Table S36*). This result is further consistent with the *sni1-1* mutant exhibiting elevated Class II crossovers.

Discussion

Our genetic mapping revealed SNI1, a functional homolog of the NSE6 component of the SMC5/6 complex, as a factor ensuring proper course of meiotic recombination in plants. SMC5/6 plays multiple roles in somatic cells during DNA damage repair, replication fork restarting, and telomere maintenance (66). For these reasons, the complex is essential across eukaryotes with most subunits being highly conserved. However, NSE6, together with NSE5, forms a module which in yeast is not permanently bound to the SMC5/6 complex (67, 68). Therefore, the absence of NSE6 is tolerated in many organisms including fission and budding yeast, vertebrate cells, and plants (42, 67, 69, 70). Moreover, SNI1 is poorly conserved at the sequence level, though it shows structural similarities among different organisms, including several armadillo repeats (42). These features indicate that SNI1/NSE6 is more susceptible to variation, which may explain identification of this gene as a modifier of meiotic crossover. Our data suggest that the I235V substitution in *SNI1* is casual to rQTL4, although it is unclear how this change affects protein functionality as evidenced by altered recombination frequency (Fig. 1) and different responses to environmental changes (Fig. 2E).

Previously, variation in REC8, a meiosis-specific component of cohesin, has been implicated as causative for differences in genome-wide recombination rates in even-toed ungulates (27, 28, 71) and plants (32, 33). Our work reveals that the SMC5/6 complex, similar to the cohesion complex (SMC1/SMC3), plays a role in crossover formation. Cohesin is a component of the chromosome axis, which holds sister chromatids together and organizes them into multiple chromatin loops (72). Anchoring chromatin loops to the axis by REC8-cohesin results in local exclusion of the recombination machinery and promoting of intersister repair of DSBs (73–75). The role of SMC5/6 in meiotic crossover formation remains incompletely characterized; however, it is known that the complex can also affect chromatin structure (76). Similarly to cohesin, the SMC5/6 complex can hold two sister chromatids inside its ring, aligning them and stimulating homologous recombination (77). Epistatic interactions between mutants of SMC5/6 and cohesin suggest that the two complexes act in related pathways in somatic cells (78). Moreover, SMC5/6-dependent SUMOylation of the SCC1 subunit of cohesin, a somatic counterpart of REC8, is required for DNA damage-induced sister chromatid cohesion (79–81). It will be interesting to investigate whether these relationships between the cohesion and SMC5/6 complexes are observed for meiotic recombination. Our data indicate that the SNI1 component of SMC5/6 complex is not required for the proper formation of the meiotic chromosome axis and sister chromatid cohesion (Fig. 5E).

In the *sni1-1* mutant, we observed no additional DSBs that could be repaired as crossovers (Fig. 5 C and D and *SI Appendix, Fig. S8*). Therefore, the increase in crossover number is likely due to the formation of abnormal JMs. The elevation of crossover frequency in chromosome distal regions (Fig. 3), seeming drop of crossover interference (Fig. 4 E–H) and partial restoration of the Class I *zip4* mutant fertility (Fig. 6 C–G), indicate that some of these abnormal JMs are repaired as Class II crossovers. Furthermore, our genetic assays show that these crossovers are largely independent from *FANCM* helicase activity (Fig. 6A) but likely depend on MUS81 endonuclease (*SI Appendix, Fig. S10B*). Together, this is consistent with the SMC5/6-SNI1 complex preventing formation of inappropriate JMs (Fig. 6H).

We propose that natural variation in *SNI1* alters the function of the SMC5/6 complex and may have biological significance in modulation of meiotic recombination under varying environmental conditions (Fig. 2E). SMC5/6-SNI1 may affect meiotic

DSB repair pathway choice, ultimately leading to changes in the crossover landscape.

Materials and Methods

Plant Material. All the lines used in the study are described in *SI Appendix, Table S37*. Genotyping of mutant lines was performed using oligonucleotides as described in *SI Appendix, Table S38*. Plants were grown in controlled environment chambers at 22 °C with long day 16-/8-h light/dark photoperiods, 60% humidity, and 150- μ mol light intensity. Prior to germination seeds were kept for 3 d in the dark at 4 °C to stratify germination.

Measurement of Crossovers Using Seed-Based Systems. Pictures of seed were acquired using epifluorescent stereomicroscope Lumar version 12 (Zeiss) equipped with a charge-coupled device (CCD) camera in brightfield, ultraviolet (UV) through a dsRed filter, and UV through a green fluorescent protein filter. The CellProfiler program was used to identify seed boundaries in micrographs and to assign a dsRed and eGFP fluorescence intensity value to each seed object (37, 82). Histograms of seed fluorescence were used to classify fluorescent and non-fluorescent seeds for each color. The genetic distance is calculated as $cM = 100 \times (1 - (1 - 2(N_G + N_R)/N_T)^{1/2})$, where N_G is the number of green alone seeds, N_R is the number of red alone seeds, and N_T is the total number of seeds analyzed.

rQTL Mapping. Genomic DNA was extracted using cetrimonium bromide and genotyped using PCR amplification of Col/Ler simple sequence length polymorphism, cleaved amplified polymorphic sequences (CAPS), or derived CAPS markers (*SI Appendix, Table S39*). R statistical package rQTL was used to perform one-dimensional QTL mapping (83). The Haley–Knott regression algorithm using 0.1-cM steps for rQTL4 fine mapping was implemented. To fit models with multiple QTLs, the *fitqtl* function with Haley–Knott regression was used. LOD score significance thresholds were established from 10,000 permutations for each mapping population.

Generation of CRISPR-Cas9 Mutant of *SN11* in Ct Background for GBS. To obtain a *sn11* mutant in Ct background (*sn11-2*), a single guide RNA (gRNA) targeted within the first exon of the gene was designed. Agrobacterium-based transformation was performed using a vector containing the gRNA (*SI Appendix, Table S40*) under the *U6* promoter and a *ICU2::Cas9* transgene. Transformants were genotyped in PCR with primers flanking the *SN11* gRNA target site, and Sanger sequencing was performed to detect deletions. Mutants with heritable deletions causing a frame shift in *SN11* were identified and selfed. M₂ plants were screened for individuals not carrying the CRISPR-Cas9 construct. The resulting *sn11-2* mutant is described in *SI Appendix, Fig. S2 C and D*.

Backcrossing of *sn11-1* (Col-0) Mutant to Ler Background for GBS. To obtain *sn11-1* mutant in Ler background, the *sn11-1* (Col) allele was crossed to Ler-0. F₁ plants obtained were backcrossed six times to Ler as recurrent female parent, and each generation was checked to carry *sn11-1* mutation by genotyping (*SI Appendix, Table S30*). The genotype of the B_C line was verified using 39 previously described Col/Ler indel markers (37).

Complementation of *sn11-1* Mutation. A DNA fragment containing *SN11* was amplified from Col or Ler genomic DNA using primers described in *SI*

Appendix, Table S40. The PCR products were cloned into the pFGC-pcoCas9 binary vector using one step cloning kit (Vazyme). These vectors were transformed into Col-420 and *sn11-1-420* hemizygous plants using *Agrobacterium* strain GV3101 and floral dipping.

Cytological Analysis. Chromosome spreads and FISH were conducted following the previously described protocol (84). Immunolocalizations were performed as previously described (85). The primary antibodies were anti-AtZYP1 (rat; 1/500 dilution), anti-MLH1 (rabbit; 1/500 dilution), anti-ASY1 (rabbit, 1/500), anti-RAD51 (rat; 1/500), and anti-SYN1 (rabbit; 1/500). Secondary antibodies were anti-rat IgG FITC-conjugated (Agrisera) and anti-rabbit IgG Alexa Fluor 555 conjugated (Molecular Probes) for immunolocalizations to detect ZYP1+MLH1 and ASY1+RAD51 and anti-rat Alexa Fluor 555-conjugated (Molecular Probes) and anti-rabbit FITC-conjugated (Sigma-Aldrich) for the immunolocalization to detect ZYP1+SYN1. For scoring RAD51 and MLH1 foci, blind analyses were applied as a protection against bias. Slides were examined on an Olympus BX61 epifluorescence microscope equipped with a CCD Olympus DP71 camera and analyze using analySIS software (Soft Imaging System).

Fertility Assays. Pollen viability was assessed from 1,500 pollen grains collected from open flowers of three plants per genotype, using Alexander staining (86). Seed set and silique length were assessed from five fruits, located at positions 6 through 10 of the main stem, in ten plants per genotype.

Additional Experimental Procedures. A detailed description of qRT-PCR analysis, DNA extraction, and library preparation for GBS, GBS bioinformatics analysis, Pollen-based measurements of genetic distance, and crossover interference can be found in *SI Appendix*.

Data Availability. All data are publicly available. Sequencing data for *sn11* ColxCol GBS libraries have been deposited under ArrayExpress accession E-MTAB-9413 (87).

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