

Features of the *Arabidopsis* recombination landscape resulting from the combined loss of sequence variation and DNA methylation

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The rate of meiotic crossing over (CO) varies considerably along chromosomes, leading to marked distortions between physical and genetic distances. The causes underlying this variation are being unraveled, and DNA sequence and chromatin states have emerged as key factors. However, the extent to which the suppression of COs within the repeat-rich pericentromeric regions of plant and mammalian chromosomes results from their high level of DNA polymorphisms and from their heterochromatic state, notably their dense DNA methylation, remains unknown. Here, we test the combined effect of removing sequence polymorphisms and repeat-associated DNA methylation on the meiotic recombination landscape of an *Arabidopsis* mapping population. To do so, we use genome-wide DNA methylation data from a large panel of isogenic epigenetic recombinant inbred lines (epiRILs) to derive a recombination map based on 126 meiotically stable, differentially methylated regions covering 81.9% of the genome. We demonstrate that the suppression of COs within pericentromeric regions of chromosomes persists in this experimental setting. Moreover, suppression is reinforced within 3-Mb regions flanking pericentromeric boundaries, and this effect appears to be compensated by increased recombination activity in chromosome arms. A direct comparison with 17 classical *Arabidopsis* crosses shows that these recombination changes place the epiRILs at the boundary of the range of natural variation but are not severe enough to transgress that boundary significantly. This level of robustness is remarkable, considering that this population represents an extreme with key recombination barriers having been forced to a minimum.

decrease in DNA methylation | epigenetic inheritance | DNA methylome | epi-haplotype

Meiotic recombination is a fundamental process in genetics whereby maternally and paternally inherited homologous chromosomes exchange material, either nonreciprocally by gene conversion or reciprocally by crossing over (CO). COs are not distributed uniformly along the genome but occur more often in chromosome arms and are strongly suppressed in pericentromeric regions (1–3), partly as a result of sequence and chromatin determinants (1, 4–8). It is commonly believed that in plants and mammals high levels of DNA sequence polymorphisms as well as heterochromatic features associated with repeats, notably dense DNA methylation and transcriptional silencing, play a central role in this suppression (1, 4).

Suppression of COs by dense DNA methylation has been demonstrated experimentally in the fungus *Ascobolus* (7). Specifically, COs were reduced when the recombination interval was methylated on one homolog and were abolished almost completely when methylated on both homologs. In *Arabidopsis*, two recent mapping studies analyzed F₂ progeny derived from crosses between Columbia *ddm1* and *met1* [*Col(ddm1)*,*Col(met1)*] DNA

methylation mutants and wild-type Landsberg [Ler(WT)] accessions and showed that loss of DNA methylation could not alleviate the suppression of COs in pericentromeric regions of chromosomes (9, 10). However, as pointed out by the authors, this experimental design could not rule out an inhibitory effect of sequence divergence between *Col* and *Ler* on COs.

An ideal design would use crosses between isogenic individuals, with one of the crossing partners having decreased DNA methylation levels throughout the genome (9). Melamed-Besudo and Levy (9) implemented such an approach by crossing *Col(ddm1)* mutant to *Col*(WT). Using two fluorescent markers spanning a 16-centimorgan (cM) interval on the arm of chromosome 3, they detected increased CO rates in F₂ plants derived from these parents relative to plants derived from a *Col*(WT)×*Col*(WT) control cross and concluded that COs in euchromatic regions can be up-regulated by loss of DNA methylation. A similar approach at a genome-wide scale and with high mapping resolution, particularly in pericentromeric regions, has not been attempted because of a lack of appropriate molecular and genetic tools. Hence, the combined effect of DNA methylation and sequence variation on COs has not been tested comprehensively in *Arabidopsis* or in any other higher eukaryote.

We previously reported the construction of a large population of epigenetic recombinant inbred lines (epiRILs) in *Arabidopsis* (11, 12), which provides a powerful experimental system to conduct such a test. These epiRILs were obtained by first crossing a fourth-generation plant homozygous for the recessive *ddm1-2* mutation with a near-isogenic WT individual. The *ddm1-2* mutation mostly affects transposable elements (TEs) and other repeats, which lose DNA methylation and become transcriptionally reactivated in a transmissible manner in many instances (11–14). However, transposition events are relatively rare (15).

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Thus, F_1 individuals can be considered homozygous throughout the genome, except at the *DDM1* locus and at the few loci affected by TE mobilization, but have chromosome pairs that differ markedly in their DNA methylation levels and transcriptional activity over TEs and other repeats (11, 16). A single F_1 *DDM1/ddm1* individual was backcrossed to the WT parental line, and after the progeny homozygous for the WT *DDM1* allele were selected, the epiRILs were propagated through seven rounds of selfing. In this design, more than 85% of all informative recombination events occur in the first two inbreeding generations (F_1 and backcross), with fewer informative events being contributed by each subsequent generation (17).

Previous targeted analysis indicated that many of the parental differences in DNA methylation and transcriptional activity of repeats are inherited stably in the epiRILs (11, 12). Regions with segregating DNA methylation states therefore can serve as physical markers to detect the frequency and distribution of recombination events along chromosomes even though the two homologs have nearly identical DNA sequences.

In this study we report the construction of a recombination map using genome-wide DNA methylation data from 123 epiRILs. This map was derived from 126 meiotically stable differentially methylated regions (DMRs) covering 81.9% of the total genome. Estimates of the genetic length for each chromosome revealed that global recombination rates are comparable with those of classical *Arabidopsis* crosses. On a local scale, we demonstrate that suppressed recombination activity within repeat-rich, pericentromeric regions of chromosomes is maintained robustly even after the removal of sequence polymorphisms and repeat-associated DNA methylation. Furthermore, we were able to identify 3-Mb regions flanking pericentromeric boundaries that appear to be subject to additional suppression and show that this effect is accompanied by increased recombination activity in chromosome arms. A direct comparison with 17 classical *Arabidopsis* crosses reveals that these recombination changes place the epiRILs at the

boundary of the range of natural variation but appear not to be severe enough to transgress that boundary significantly.

Results

Construction of a Recombination Map Using Transgenerationally Stable DMRs. To demonstrate that transgenerationally stable DMRs can be used for the construction of a recombination map in an isogenic population, we carried out methylated DNA immunoprecipitation followed by hybridization to a whole genome DNA tiling array (MeDIP-chip) on 123 epiRILs and on the two parental lines (256 array experiments including replicates). The 123 epiRILs originally were chosen using a selective (epi)genotyping strategy for two uncorrelated complex traits, flowering time and root length. We used a three-state Hidden Markov Model (HMM) to classify tiling array signals into three underlying DNA methylation states (18): unmethylated (U), intermediate methylation (I), or methylated (M). Benchmarking of these HMM calls against whole-genome bisulphite sequencing data ($\sim 30\times$) for six epiRILs confirmed that both the MeDIP protocol and the analysis method performed well (*SI Appendix, Fig. S1 and Table S1*). Comparison of the two parental DNA methylomes revealed 2,611 DMRs representing clear instances of methylation loss in *ddm1* (transitions from M to U). These DMRs (median length: 1,211 bp; range: 318–24,624 bp) were distributed throughout the genome but, as expected, were more abundant in pericentromeric regions (Fig. 1*A* and *SI Appendix, Table S2*) (19).

We examined the DNA methylation state at all parental DMRs in each of the 123 epiRILs and inferred their parent of origin (i.e., epigenotypes). Segregation was not compatible with stable inheritance of *ddm1*-induced DNA hypomethylation for 1,744 (66.8%) of the parental DMRs, and in most of these cases our data pointed to fully or partially penetrant reversion to WT DNA methylation. In contrast, 867 (33.2%) of the parental DMRs segregated in the expected 3:1 Mendelian ratio (*SI Appendix, Fig. S2 and Table S3*). Stable DMRs were associated with a comparatively lower abundance of siRNAs in the WT and *ddm1* parental

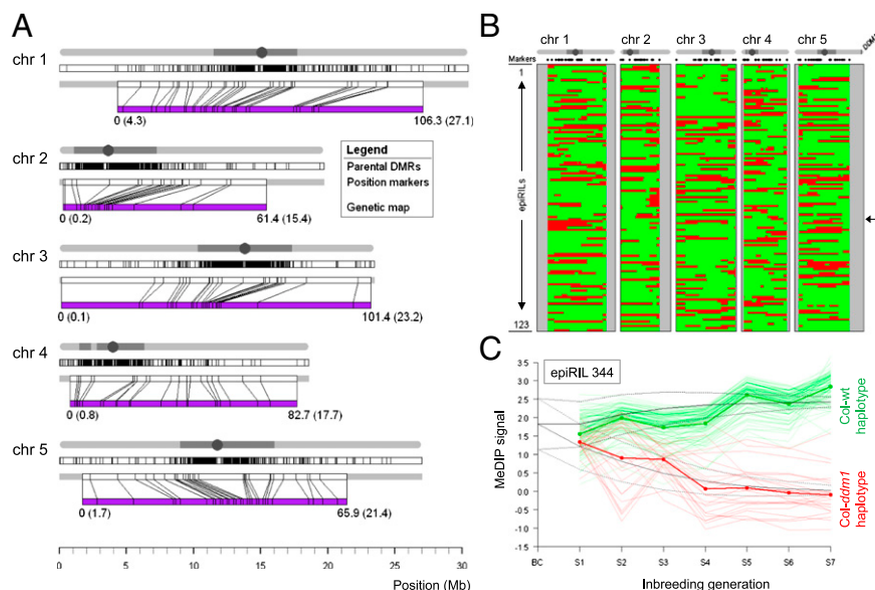


Fig. 1. Recombination map construction. (A) Genome-wide distribution of the 2,611 parental DMRs (Top) and the 126 DMRs (i.e., markers; Middle) retained for construction of the recombination map (purple, Bottom) for each of the five *Arabidopsis* chromosomes. The mapping between physical and genetic positions of markers is shown. (B) Inference of inherited WT (green) and *ddm1* (red) haplotypes along the genome (x-axis) as inferred from the recombination map for each of the 123 epiRILs (y-axis) (*SI Appendix, Table S5*). Chromosome extremities not covered by the genetic map are indicated in gray. The genome of epiRIL 344 is indicated by an arrow. A schematic representation of each chromosome is plotted above the map with the physical location of the *DDM1* gene shown at the end of chromosome 5. (C) Transgenerational methylation data for epiRIL 344. Shown are the average methylation signals for the 126 markers, with regions that are predicted to become fixed for the *ddm1* haplotypes (thin red lines) and the WT haplotypes (thin green lines) after seven selfing generations. The average signals (red and green thick solid lines) are in agreement with Mendelian inbreeding theory (black solid lines).

lines (*SI Appendix, Fig. S3*). These findings are in agreement with previous analyses (11, 12) and indicated that the 867 stable DMRs are not efficient targets of siRNA-mediated DNA remethylation, even after eight rounds of meiosis. These stable DMRs therefore could serve as physical markers in an extension of the Lander–Green algorithm (20) to derive a genetic map. After application of the algorithm and removal of mainly genetically redundant markers (i.e., markers located less than 0.0001 cM apart), 126 of the original 867 markers were retained (Fig. 1 *A* and *B* and *SI Appendix, Fig. S2* and *Table S4*). These 126 markers covered ~81.9% of the total genome (74.7, 77.0, 98.4, 91.1, and 73.0% of chromosomes 1, 2, 3, 4, and 5, respectively).

Many of the 126 markers contained TE sequences, consistent with the targeted effect of *ddm1* on these and other repeats (*SI Appendix, Fig. S4*). However, in a vast majority of cases, markers included only TE relics, which likely have lost their capacity to be mobilized (*SI Appendix, Table S6*). Indeed, both comparative genomic hybridization (*SI Appendix, Fig. S5*) and preliminary whole-genome resequencing suggested that none of the 126 DMRs contain sequences that were mobilized in the parental *ddm1* line or the epiRILs (*SI Appendix, Table S6*). Consistent with this finding, pair-wise recombination fractions between the 126 markers indicated a well-behaved and robust genetic map, reminiscent of those typically seen in classical crosses involving DNA sequence markers, with high correlation among linked loci and virtually no correlations among loci in different chromosomes (*SI Appendix, Fig. S6*). Moreover, all inferred *ddm1*-inherited non-recombinant pericentromeric haplotypes contained significantly less DNA methylation and were more actively transcribed than their WT counterparts (*SI Appendix, Figs. S7* and *S8*).

To test further the transgenerational stability of the 126 markers as well as our inference of the parental epigenotypes at these marker locations, we performed genome-wide DNA methylation analysis for one selected line (epiRIL 344) for each of its seven selfing generations (7×2 replicates = 14 array experiments). Fixation occurred for the predicted parental epigenotype in each case, and the rate of approach toward fixation was consistent with Mendelian inbreeding theory for a backcross-derived RIL (19) (Fig. 1*C*). Taken together, these results rule out any ambiguity in the actual location or DNA methylation state of the stable DMRs used for constructing the genetic map.

Total Genetic Length in the epiRILs Does Not Diverge Significantly from the Natural Range.

One approach for evaluating the

epiRILs recombination map is by comparison with a Col (WT)×Col(WT)-derived reference cross. In this set-up, changes in recombination patterns can be attributed directly to DNA methylation loss. However, tracking recombination events in such a reference is experimentally challenging. It requires a system akin to the fluorescent marker reporters used by Melamed-Besudo and Levy (9), which does not easily scale genome-wide. An alternative approach is to evaluate the epiRILs in the context of natural variation. In terms of DNA sequence and DNA methylome divergence of its founder parental lines, the epiRILs can be viewed as representing an extreme situation with key barriers to recombination having been forced to a minimum. An important question therefore is how genome-wide recombination patterns in this population compare with those seen in crosses derived from different pairs of natural accessions.

We estimated the genetic length for each of the five epiRIL chromosomes using Haldane's map function. The lengths were 106.3, 61.4, 101.4, 82.7, and 65.9 cM for chromosomes 1–5, respectively, and correlated positively with physical chromosome length (*SI Appendix, Fig. S9*). The total length of the genetic map was 417.7 cM, yielding an average marker spacing of ~0.804 Mb (3.45 cM). These estimates are similar to those previously reported for genetic maps based on classical *Arabidopsis* crosses (21–24). The use of other map functions that account for CO interference, such as the Kosambi or Carter and Falconer functions, yielded very similar results (*SI Appendix, Fig. S10*). To perform a more direct comparison between the epiRIL map and those of classical *Arabidopsis* crosses, we reanalyzed recombination data obtained for 17 F₂ populations (24) that were derived from pairs of 18 distinct natural accessions. In total, these populations consisted of 7,045 plants (~410 plants per cross; range: 235–462 plants), which were genotyped at 235 markers on average (range: 215–257 markers) (24). To facilitate a meaningful comparison, we constructed a consensus map using 83 markers that were shared across populations (*SI Appendix, Fig. S11* and *Table S7*). Thorough testing showed that the reduction to 83 markers in the epiRIL and F₂ maps led to no significant loss of information in capturing the linkage structure along chromosomes (*SI Appendix, Figs. S12* and *S13*), and the 83 markers therefore were deemed appropriate for this comparative analysis.

Estimates of the genetic length of each of the five chromosomes revealed substantial natural variation among the F₂ populations (Figs. 2*B* and 3*A*). However, the genetic lengths of the

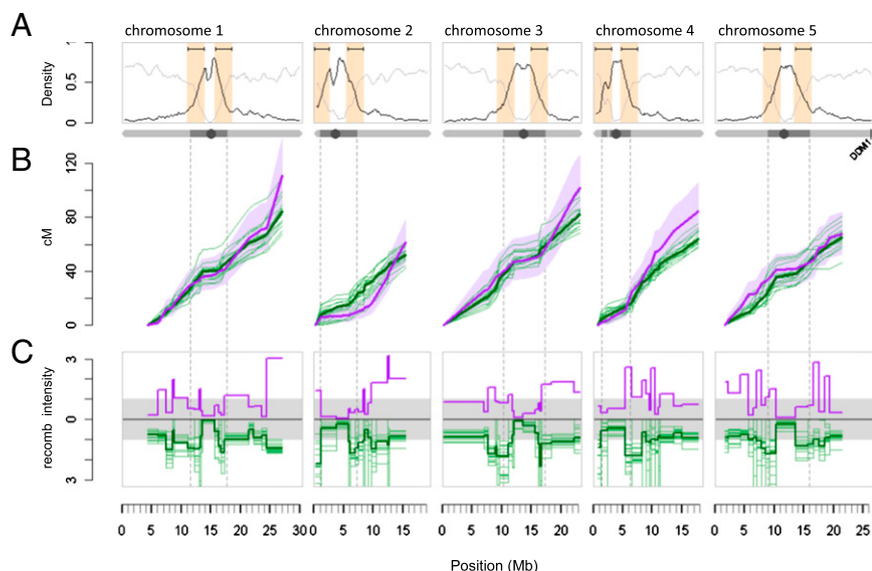


Fig. 2. Comparison of global and local recombination patterns in the epiRILs and the 17 F₂ populations (24). (A) Chromosome-wide gene (light gray line) and transposon (dark line) density distribution. The 3-Mb windows bracketing the intersection points between transposon- and gene-dense regions are indicated in orange. (B) Cumulative cM lengths of the epiRILs (thick purple line) and each of the F₂ populations (thin green lines) using the consensus map. Purple shading shows the $\pm 95\%$ confidence interval (CI). The thick green line denotes the average F₂ cumulative length (in cM). The dotted vertical lines define the pericentromeric regions of each chromosome. (C) The distribution of normalized recombination intensities (cM/Mb of a given marker interval divided by the cM/Mb chromosome average) shows suppression of recombination within pericentromeric regions and elevation at its boundaries. Color coding is as in B.

because of the sparsity of their genetic markers (two to three markers per pericentromeric region). Marker density in the epiRIL map, in contrast, was relatively high within AT zones and even permitted fine mapping of shared and nonshared recombination breakpoints to a resolution as low as 4 kb (*SI Appendix, Figs. S16 and S17 and Tables S9 and S10*).

Furthermore, our data indicate that suppression of recombination within AT zones in the epiRILs is accompanied by increased recombination in chromosome arms (Fig. 3*D* and *SI Appendix, Fig. S18*). This apparent compensatory effect reconciles the enhanced local suppression seen in the epiRILs with the earlier observation that chromosome-wide recombination rates are relatively large compared with the F_2 populations. This effect was most pronounced on epiRIL chromosomes 1, 2, and 5 (the chromosomes with the strongest suppression in the AT zone), with recombination intensities being 1.23, 1.6, and 1.3 times above the chromosomes' average (*SI Appendix, Table S8*). We failed to identify a similar trend in the F_2 populations (*SI Appendix, Fig. S18 and Table S8*), suggesting that this effect is a specific feature of the epiRIL recombination landscape.

Discussion

In this study we demonstrate that stable DNA methylation differences can be used as physical markers to derive genome-wide recombination patterns in a near isogenic population of epiRILs. We find that recombination suppression is maintained robustly in pericentromeric regions of the epiRILs, despite the extensive loss of sequence variation and of DNA methylation and transcriptional silencing over repeats. This observation indicates that these factors do not play a major role in the suppression of pericentromeric COs. This finding is contrary to common belief and is particularly intriguing given the interplay between recombination and transcription observed in yeast and the mouse (26, 27). Whether mechanisms exist in *Arabidopsis* that actively sequester the recombination machinery away from gene-promoter regions or other genomic elements, as in the mouse (27), remains to be determined.

Nonetheless, our results indicate that loss of DNA methylation over repeat sequences can lead to a local reinforcement of recombination suppression in pericentromeric regions and to increased recombination activity along chromosome arms. Similar results were reported by Melamed-Bessudo and Levy (9) and Mirouze et al. (10) using genetically divergent populations. Therefore we conclude that the absence of sequence polymorphisms is insufficient to counteract the enhanced suppressive effects induced by the loss of DNA methylation in pericentromeric regions. On the other hand, the lack of sequence polymorphisms still may be partly responsible for the increased recombination rates observed in chromosome extremities (9).

Melamed-Bessudo and Levy (9) demonstrated that *ddm1*-induced demethylation of only one homolog produces the same recombination changes seen when both homologs are demethylated. Our results and conclusions therefore should be generalizable to the two-homolog situation. However, it has been shown in *Ascobolus* that DNA methylation of a known recombination hotspot inhibits COs more severely when both homologs are methylated (7). Similar localized dosage effects may therefore also be present in *Arabidopsis*.

Our study and those of Melamed-Bessudo and Levy (9) and Mirouze et al. (10) have used well-characterized *ddm1* and *met1* DNA methylation mutants as a tool to perturb genome-wide methylation levels experimentally. Both *ddm1* and *met1* experience a nearly 70% reduction in DNA methylation levels genome-wide. This drastic loss probably sets an upper limit to the amount of demethylation that can be incurred in nature. Indeed, it is difficult to conceive of mechanisms that would elicit similar or more severe changes under natural settings, unless they involve

spontaneous mutations in genes important for DNA methylation control, such as *ddm1* or *met1*. Interestingly, a recent analysis of *Arabidopsis* mutation accumulation lines showed that drastic alterations in the methylome of one outlier line were likely caused by a spontaneous mutation in a methyl-transferase gene (28, 29), which must have arisen during just 30 generations of selfing. This observation suggests that similar events are certainly plausible under natural conditions.

An assessment of whether strong methylation loss can elicit recombination changes at magnitudes that are sufficient to drive genome evolution in this species has been lacking. Our study is an initial step in providing such an assessment. Our analysis of the 17 F_2 populations derived from 18 natural accessions (24) allowed us to quantify the magnitude of the recombination changes observed in the epiRILs in the context of natural variation. Although we find that the epiRILs nearly always are situated at the boundary of the natural range, there is no strong evidence that local and global recombination patterns in this population markedly transgress the natural range. Indeed, in many cases, several of the F_2 populations displayed even more extreme divergence from the F_2 population average than did the epiRILs. These findings lead us to conclude that severe losses of DNA methylation along *Arabidopsis* chromosomes have no drastic implications for recombination-mediated genome evolution. This high level of robustness raises questions concerning the precise mechanisms that have shaped the recombination landscape in this species in the first place.

Of course, severe depletion of DNA methylation can drive other important events, such as large-scale structural rearrangements and polyploidization, which may impact the course of genome evolution. In addition, natural epigenetic variation, such as that associated with differential DNA methylation, can act on complex traits that are under natural selection (30), thereby changing linkage disequilibrium relations within and across chromosomes. However, understanding and documenting the impact of epigenetic variants on complex traits is challenging, mainly because of the technical difficulties in ruling out the confounding effect of DNA sequence polymorphisms (31). Because of this limitation, it has been argued that the epiRILs constitute an ideal system for the study of epigenetic inheritance in *Arabidopsis* (17, 32–34). We and others have shown recently that many adaptive phenotypes, such as plant height, flowering time, and growth rate, are highly heritable in this population (12, 35, 36). Segregating phenotypic effects also have been observed in another epiRIL population which was obtained from a cross between Col(*met1*) and Col(WT) (37).

A logical next step in the analysis of these populations is to map and characterize the epigenetic basis of these complex traits. The linkage map reported here (Fig. 1*B*) can be used in conjunction with classical quantitative trait-locus mapping methods to achieve this characterization in the *ddm1*-derived epiRILs. Ultimately, such efforts should contribute significantly to our understanding of epigenetics in adaptive evolution.

Materials and Methods

Methylome Analysis. MeDIP was carried out as previously described (18) followed by hybridization to a custom NimbleGen tiling array covering the *Arabidopsis* genome at 165 nt resolution (38). Including dye-swaps, we performed a total of 256 array experiments (*SI Appendix, section 1*). For each array, probe signals were classified into three underlying methylation states, methylated (M), intermediate (I), or unmethylated (U), using the HMM presented previously (*SI Appendix, section 2*) (18). These inferred methylation states were cross-validated against whole-genome bisulphite-sequencing data of six epiRILs (*SI Appendix, section 3, Fig. S1, and Table S1*).

Definition of Parental DMRs. We conducted a probe-level comparison of the HMM calls between the *ddm1* and WT parents (*SI Appendix, section 4*). Probe-level methylation calls were denoted as polymorphic when the parents differed (e.g., I in *ddm1* and M in WT) and as nonpolymorphic when they

were identical (e.g., U in *ddm1* and U in WT). Neighboring probes reporting the same polymorphic state were collapsed into single regions. Hence, parental DMRs were defined as regions of at least three consecutive probes that reported the same extreme polymorphic state (i.e., transitions from M in WT to U in *ddm1* or vice versa). We found 2,611 DMRs, all of which were U in the *ddm1*. Detailed summary statistics are given in *SI Appendix, Table S2*.

Calling of Parental Origin of DMRs in the epiRILs. For any given epiRIL the parental origin of each DMR (i.e., epigenotype) was determined using an HMM-based inference method (*SI Appendix, section 4*).

Mendelian Segregation Criterion. Under the assumption that DMRs were stable for eight generations of breeding, both WT- and *ddm1*-like parental states should appear according to Mendelian segregation ratios in the epiRILs. The sampling variation around these ratios was calculated from a binomial distribution taking into account the sample size ($n = 123$), the cross design, and the 8% F_2 contamination previously reported (12). DMRs in the epiRILs showing a percentage of WT-like states between 62.7% and 83.3% (the expected value being 73%) were taken as putative transgenerationally stable markers. In total 867 parental DMRs fulfilled this criterion and were used subsequently as a starting point for map construction (*SI Appendix, section 5, Fig. S2, and Table S3*).

Extension of Lander–Green Algorithm. Derivation of a genetic map using DMRs was carried out through an extension of the Lander–Green algorithm (20), which was designed to accommodate marker and individual specific error rates. Our implementation of this algorithm is detailed in *SI Appendix, section 6*.

Transcriptome Analysis of epiRILs and *ddm1* Seedlings. Whole-genome expression profiling was performed using a custom NimbleGen tiling array (37). For experimental details, see *SI Appendix, section 7*.

Transgenerational Analysis of DMRs. MeDIP-chip was carried out for epiRIL 344 for seven generations of selfing after the backcross, following the protocol described above. At each generation, DNA from five siblings was pooled. The expected signal behavior was derived using a Markov Chain strategy, considering the Mendelian inheritance of the marker probes (*SI Appendix, section 8*).

Construction of the Consensus Map. To facilitate a meaningful comparison of the epiRILs map with those of the 17 different F_2 populations, we constructed a consensus map (*SI Appendix, section 9 and Fig. S11*) by using the epiRILs map as a reference and selecting from each of the F_2 maps the SNPs closest to the reference, allowing a maximum distance of ± 1.39 Mb. The average distance from reference was ± 0.17 Mb, which led to little loss of information in capturing the recombination structure along the genome (*SI Appendix, Figs. S12 and S13*). Markers deemed too distant were not included in the consensus map. This process resulted in 83 markers (*SI Appendix, Table S7*).

Recombination Intensities at Major Annotation Transitions. Fig. 2 A and C shows that the recombination intensity increases rapidly at the pericentromeric boundaries, which also coincide with major transitions in genome content from genes to transposons. To find the area where the recombination intensity is maximal, we implemented a sliding window approach (*SI Appendix, section 10 and Fig. S15*).

Note Added in Proof. During the reviewing process, Yelina et al (Yelina NE, Choi K, Chelysheva L, Macaulay M, de Snoo B, et al. (2012) Epigenetic Remodeling of Meiotic Crossover Frequency in Arabidopsis thaliana DNA Methyltransferase Mutants. *PLoS Genet* 8(8): e1002844. doi:10.1371/journal.pgen.1002844) reported elevated centromere-proximal COs, coincident with pericentromeric decreases and distal increases in met1 mutants. However, total numbers of CO events were found to be similar between wild type and met1. These results support the trends observed in the epiRIL population.

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