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# The genetic and epigenetic landscape of the Arabidopsis centromeres

Short title: Assembly of the Arabidopsis centromeres.

**One-sentence summary:** Long-read sequencing and assembly of the Arabidopsis centromeres reveals their genetic and epigenetic topography.

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#### Abstract:

Centromeres attach chromosomes to spindle microtubules during cell division and, despite this conserved role, show paradoxically rapid evolution and are typified by complex repeats. We used long-read sequencing to generate the Col-CEN *Arabidopsis thaliana* genome assembly that resolves all five centromeres. The centromeres consist of megabase-scale tandemly repeated satellite arrays, which support high CENH3 occupancy and are densely DNA methylated, with satellite variants private to each chromosome. CENH3 preferentially occupies the satellites with least divergence and greatest higher-order repetition. The centromeres are invaded by *ATHILA* retrotransposons, which disrupt genetic and epigenetic organization of the centromeres. Crossover recombination is suppressed within the centromeres, yet low levels of meiotic DSBs occur that are regulated by DNA methylation. We propose that Arabidopsis centromeres are evolving via cycles of satellite homogenization and retrotransposon-driven diversification.

#### Introduction:

Despite their conserved function during chromosome segregation, centromeres show diverse organization between species, ranging from single nucleosomes to megabase-scale tandem repeat arrays (1). Centromere 'satellite' repeat monomers are commonly ~100–200 bp, with each repeat capable of hosting a CENPA/CENH3-variant nucleosome (1, 2). CENPA/CENH3 nucleosomes ultimately assemble the kinetochore and position spindle attachment on the chromosome, allowing segregation during cell division (3). Satellites are highly variable in sequence composition and length when compared between species (2). The library of centromere repeats present within a genome often shows concerted evolution, yet they have the capacity to change rapidly in structure and sequence within and between species (1, 2, 4, 5). However, the genetic and epigenetic features that contribute to centromere evolution are incompletely understood, in large part due to the challenges of centromere sequence assembly and functional genomics of highly repetitive sequences.

Genomic repeats are notoriously difficult to assemble from fragmented sequencing reads. This difficulty is derived from repeat length and their level of similarity, with longer, high-identity repeats being the most challenging to correctly assemble (6). As sequencing reads have become longer, eukaryotic *de novo* genome assemblies have captured an increasingly complete and accurate picture of repetitive elements genome-wide. For example, using long but noisy reads, such as PacBio Continuous Long Reads, or early versions of Oxford Nanopore Technologies (ONT) long reads, researchers have routinely assembled interspersed retrotransposons, such as LINE and SINE elements in the human genome (7) and LTR elements in plant genomes (8). However, due to high error rates, these third-generation sequencing technologies did not enable accurate assembly of large and homogenous repeats, including telomeres, centromeres, segmental duplications and tandemly repeated rRNA genes (9). Owing to improved DNA extraction and library preparation protocols, together with advanced machine learning-based basecalling, ONT long reads have become substantially longer and more accurate (>100 kbp with 95–99% modal accuracy) (*10*). Additionally, PacBio's High-Fidelity (HiFi) reads, while shorter (~15 kbp), are exceptionally accurate (>99%) (*11*). Using these technologies with new computational methods, researchers recently assembled a complete telomere-to-telomere (T2T)

representation of the CHM13hTERT human genome, including centromere satellite arrays, and other sequences that were previously unresolved (5, 9, 12-17). This work revealed that current ONT and HiFi reads are sufficient to span the interspersed unique marker sequences in human centromeres, suggesting that truly complete genome assemblies for diverse eukaryotes are on the horizon.

*Arabidopsis thaliana* is a major model plant species and its genome was first sequenced in 2000, yet the centromeres, telomeres, and ribosomal DNA repeats have remained unassembled, due to their high repetition and similarity (*18*). The Arabidopsis centromeres are known to contain millions of base pairs of the *CEN180* satellite repeat, which support CENH3 loading (*19–23*). We used long-read ONT DNA sequencing, followed by polishing with high-accuracy PacBio HiFi reads, to establish the Col-CEN reference assembly, which wholly resolves all five Arabidopsis centromeres. The assembly contains a library of 66,131 *CEN180* satellites, with each chromosome possessing mostly private satellite variants. Higher-order *CEN180* repetition is prevalent within the centromeres and is also chromosome-specific. We identify *ATHILA* LTR retrotransposons that have invaded the satellite arrays and interrupt genetic and epigenetic organization of the centromeres. By analyzing SPO11-1-oligo data from mutant lines, we demonstrate that DNA methylation epigenetically silences initiation of meiotic DNA double-strand breaks (DSBs) within the centromeres. Together, our data are consistent with satellite homogenization and retrotransposon invasion driving cycles of centromere evolution in Arabidopsis.

#### Complete assembly of the Arabidopsis centromeres

The current *Arabidopsis thaliana* TAIR10 reference genome, representing the Columbia (Col-0) accession, is an exceptionally accurate and complete eukaryotic assembly and an invaluable resource for plant science (*18*). However, TAIR10 fails to represent the telomeres, some rDNAs and the centromere satellite arrays. To resolve these remaining sequences, we collected Col-0 genomic ONT and HiFi sequencing data comprising a total of 73.6 Gbp (~56× coverage of reads >50 kbp) and 14.6 Gbp (111.3× coverage, 15.6 kbp mean read length), respectively. This long-range sequence information, combined with our optimized *de novo* assembly and validation pipeline, yielded a nearly closed and

highly accurate assembly of the Col-0 genome (Col-CEN v1.2). Chromosomes 1, 3 and 5 are wholly resolved from telomere-to-telomere, and chromosomes 2 and 4 are complete apart from the short-arm 45S rDNA clusters and adjacent telomeres (**Fig. 1**).

After telomere patching and repeat-aware polishing with R9 and R10 ONT reads, HiFi reads and short Illumina reads using methods developed for the Human T2T project (24), the Col-CEN assembly is highly accurate with a QV of 45.99 and 51.71 inside and outside of the centromeres, equivalent to approximately one error per 40,000 and 148,000 bases, respectively (**Fig. S1-S2A** and **Table S1**). Additionally, Hi-C and Bionano optical maps validate the large-scale structural accuracy of the assembly (**Fig. S2**). The Col-CEN assembly is highly concordant with TAIR10, showing no large structural differences within the chromosome arms (**Fig. 1B**). 97.5% of Col-0 BAC contigs that align with high coverage and identity (>95%) to TAIR10 also align with high coverage and identity to Col-CEN, and 99.9% of TAIR10 gene annotations (longer than 40 bp) are represented in Col-CEN.

Despite the broad similarity between Col-CEN and TAIR10, we note a number of important distinctions between the assemblies. First, unlike TAIR10, Col-CEN reconstructs all five centromeres spanning 12.6 Mbp of new sequence, 120.0 and 97.6 kbp of *45S* rDNA in the chromosome 2 and 4 nucleolar organizer regions (NORs), and the complete telomeres of the 8 chromosome arms without sub-telomeric NORs (**Fig. 1A–1C, S1–S3**). Aside from these assembly differences, we found several instances of apparently genuine variation between the Col-0 laboratory strains used to generate TAIR10 and Col-CEN (**Fig. S4** and **Tables S2–S3**). For example, we identified a Col-CEN thionin gene cluster deletion relative to TAIR10 (**Fig. S4**). In total, 27 TAIR10 genes are missing from Col-CEN (**Tables S2–S3**). Finally, to comprehensively account for variation between Col-0 strains, we aligned Col-0 ONT, HiFi, and Illumina data to the Col-CEN assembly and called variants, providing a database of potential allelic differences, including heterozygous variants (https://github.com/schatzlab/Col-CEN). Notably, this analysis revealed only 41 and 37 SV calls from ONT and HiFi data genome-wide, respectively, consistent with very low levels of heterozygosity.

We confirmed the order of chromosome landmarks flanking centromere 1 using fluorescent *in situ* hybridization (FISH), which included labeling of a telomeric repeat cluster located adjacent to the centromere (**Fig. 1D** and **S5**). To provide independent validation of centromere structure, we performed *in silico* digestion with *Asc*I and *Not*I and compared the predicted fragments to physical maps generated previously using restriction digestion, pulsed-field gel electrophoresis and Southern blotting, which showed a high degree of agreement with Col-CEN (**Fig. S6**) (*25–27*). Given the relatively low resolution of Southern blotting, we also examined our Bionano optical map data across the centromeres (**Fig. S7**). The optical contigs are consistent with the structure of the assembled *CEN180* arrays, although the low density of labeling sites within the centromeres prevents full resolution by optical fragments alone (**Fig. S7**).

The assembled centromere sequences are characterized by a repeated 178-bp motif (*CEN180*) that is organized into higher-order repeats (HORs) (**Fig. 1D, 2** and **S8**). We validated the structural and baselevel accuracy of the centromeres using techniques from the Human T2T consortium (9, 12). Briefly, we aligned our Col-0 long reads to the assembly and observed even coverage across the centromeres, with few loci showing plausible alternate base signals (**Fig. S1B**). We also observed relatively few 'missing' *k*-mers that are found in the assembly but not in Illumina short reads, which are diagnostic of residual consensus errors that remain after polishing (**Fig. S1B**) (28). We observe that unique 'marker' sequences are relatively frequent, with a maximum distance between consecutive markers in the assembled centromeres of only 41,765 bp, suggesting that our long reads can confidently span several unique markers and thus reliably assemble centromeric loci (**Fig. S1C**). Notably, the five centromeres are relatively distinct at the sequence level, with each exhibiting chromosome-specific repeats (**Fig. 1E, 2** and **Tables S4–S5**). This is consistent with our assembly pipeline unambiguously separating the five centromere sequences. To independently validate the presence of chromosome-specific *CEN180* variants, we designed FISH probes to label high copy repeats that vary in frequency between the centromeres. For example, the *CEN180-a*, *CEN180-y* and *CEN180-b* probes specifically label centromere 1 (**Fig. 1F** and **S5**), providing cytogenetic validation for chromosome-specific satellites predicted by the Col-CEN assembly.

#### The Arabidopsis CEN180 satellite repeat library

We performed *de novo* searches for tandem repeats to define the centromere satellite library (**Table S4**). We identified 66,131 *CEN180* satellites in total, with between 11,848 and 15,613 copies per chromosome (**Fig. 2, S9** and **Table S4**). The *CEN180* repeats form large tandem arrays, with the satellites within each centromere predominantly found on the same strand, with the exception of centromere 3, which is formed of two blocks of satellites on opposing stands (**Fig. 1D** and **S8**). The distribution of repeat monomer length is tightly constrained around 178 bp (**Fig. 2A** and **S9**). We aligned all *CEN180* sequences to derive a genome-wide satellite consensus and calculated nucleotide frequencies at each alignment position to generate a position probability matrix (PPM). Each satellite was compared to the PPM to calculate a variant distance by summation of disagreeing nucleotide probabilities. Substantial sequence variation was observed between satellites and the PPM, with a mean variant distance of 20.2 per *CEN180* (**Fig. 2A**). Each centromere shows essentially private libraries of *CEN180* monomer sequences, with only 0.3% sharing an identical copy on a different chromosome (**Fig. 1E** and **Table S4**). In contrast, there is a high degree of *CEN180* repetition within chromosomes, with 57.1–69.0% showing one or more duplicates (**Table S4**). We also observed a minor class of *CEN160* tandem repeats found on chromosome 1 (1,289 repeats, mean length=158.2 bp) (23).

We aligned CENH3 ChIP-seq data to the Col-CEN assembly and observed on average 12.9-fold log<sub>2</sub>(ChIP/input) enrichment within the *CEN180* arrays, compared to the chromosome arms (**Fig. 1D** and **S8**) (*19*). CENH3 ChIP-seq enrichment is generally highest within the interior of the main *CEN180* arrays (**Fig. 1D** and **S8**). We observed a negative relationship between CENH3 ChIP-seq enrichment and *CEN180* variant distance (**Fig. 2D–2E**), consistent with CENH3 nucleosomes preferring to occupy satellites that are closer to the genome-wide consensus. In this respect, centromere 4 is noteworthy, as it consists of two distinct *CEN180* arrays, with the right array showing both higher variant distances

and lower CENH3 ChIP-seq enrichment (**Fig. 1D, 2D** and **S8**). Together, this is consistent with satellite divergence leading to loss of CENH3 binding, or vice versa.

To define CEN180 higher-order repeats (HORs), monomers were considered the same if they shared five or fewer pairwise variants. Consecutive repeats of at least two monomers below this variant threshold were identified, yielding 2,408,653 HORs (Fig. 2D and Table S5). Like the CEN180 monomer sequences, HORs are almost exclusively chromosome-specific (Table S5). The mean number of CEN180 monomers per HOR was 2.41, equivalent to 429 bp (Fig. 2B and Table S5), and 95.4% of CEN180 were part of at least one HOR. HOR block sizes show a negative exponential distribution, with the largest HOR formed of 60 monomers on chromosome 3, equivalent to 10,689 bp (Fig. 2B). Many HORs are in close proximity (26% are <100 kbp apart), although they are dispersed throughout the length of the centromeres. For example, the average distance between HOR blocks was 380 kbp and the maximum distance was 2,365 kbp (Fig. 2B and Table S5). We also observed that HOR blocks that were a greater distance apart showed a higher level of variants between the blocks (variants/monomer) (Fig. 2F), which is consistent with satellite homogenization being more effective over repeats that are physically closer. The CEN180 groups with highest CENH3 occupancy also show the greatest level of higher-order repetition and higher CG DNA methylation frequency (Fig. 2D-2E and 2G). However, a notable exception to these trends is centromere 5, which harbors 6.8–13.4% of HORs compared to the other centromeres, yet still recruits comparable CENH3 (Fig. 2G and Table S5).

#### Invasion of the Arabidopsis centromeres by ATHILA retrotransposons

In addition to reduced *CEN180* higher-order repetition, centromere 5 is also frequently disrupted by breaks in the satellite array (**Fig. 2G** and **S8**). Genome-wide, within the main satellite arrays, the vast majority of sequence is *CEN180* (92.8%), with only 111 interspersed sequences larger than 1 kbp. Within these 111 gaps, we identified 53 intact and 20 fragmented *ATHILA* LTR retrotransposons of the *GYPSY* superfamily, belonging to the *ATHILA*, *ATHILA2*, *ATHILA4*, *ATHILA5* and *ATHILA6A/6B* 

subfamilies (Fig. 3A-3C and Table S6) (29-31). The intact ATHILA elements have a mean length of 11.05 kbp, and the majority have highly-similar paired LTRs, target site duplications (TSDs), primer binding sites (PBS), polypurine tracts (PPT) and GYPSY superfamily open reading frames (Fig. 3C and Table S6). LTR comparisons indicate that the centromeric ATHILA elements are young, with on average 98.7% LTR sequence identity, which was significantly higher than for ATHILA elements located outside the centromere (96.9% identity, n=58, Wilcox test  $P=4.89\times10^{-8}$ ) (Fig. 3D and S10). We also identified 12 ATHILA solo LTRs, which is consistent with post-integration intra-element homologous recombination (Table S6). Notably, we observed 6 instances where gaps containing intact ATHILA fragments or solo LTRs were duplicated on the same chromosome that were located between 8.9 and 538.5 kbp apart, consistent with transposon sequences being copied post-integration, potentially via the same mechanism that generates CEN180 HORs. For example, a pair of adjacent ATHILA5 and ATHILA6A elements within centromere 5 had been duplicated within a higher-order repeat (Fig. S11). These duplicated elements share target site duplications (TSDs) and flanking sequences, are nearly identical in length (11,345 and 11,346 bp for the ATHILA6A copies) and show high identity between copies (99.5% and 99.6%, respectively) (Fig. S11 and Table S6). In contrast, the surrounding CEN180 sequences display higher divergence and copy number variation between the higher order repeats (94.3– 97.3% identity) (Fig. S11). This indicates an elevated rate of sequence change within the CEN180 compared to the ATHILA, following this higher-order duplication.

We analyzed centromeric *ATHILA* elements for CENH3 ChIP-seq enrichment and observed a decrease relative to the surrounding *CEN180*, yet higher levels than in *ATHILA* elements located outside of the centromere (**Fig. 3E**). The *ATHILA* elements also show greater H3K9me2 enrichment compared to flanking *CEN180* (**Fig. 3E**). We used our ONT reads to profile DNA methylation over the *ATHILA* and observed dense methylation, at a similar level to the surrounding *CEN180*, although with higher CHG-context methylation (**Fig. 3F**). Hence, *ATHILA* elements are distinct from the *CEN180* satellites at the chromatin level. We profiled *CEN180* variants around gaps containing *ATHILA* insertions (n=65) (including intact elements, fragments and solo LTRs) and observed an elevation of *CEN180* satellite divergence in the flanking regions (**Fig. 3G**), which is reminiscent of elevated *PSR* tandem repeat

divergence observed at the junction of a *NATE* retrotransposon in *Nasonia* (*32*). This may indicate that *ATHILA* insertion was mutagenic on the surrounding satellite repeats, or that transposon insertion influenced the subsequent divergence or homogenization of the adjacent *CEN180* repeats. We used FISH to cytogenetically validate the presence of the *ATHILA6A/6B* and *ATHILA2* sub-families within the centromeres (**Fig 3H** and **S5**). Together, this indicates that centromeric *ATHILA* insertions interrupt the genetic organization of the Arabidopsis *CEN180* satellite arrays.

#### Epigenetic organization and meiotic recombination within the centromeres

To assess genetic and epigenetic features of the centromeres, we analyzed all chromosome arms along their telomere–centromere axes using a proportional scale (**Fig. 4A**). Centromere midpoints were defined as the point of maximum CENH3 ChIP-seq enrichment (**Fig. S12**). As expected, *CEN180* satellites are highly enriched in proximity to centromere midpoints (**Fig. 4A**). Gene density drops precipitously as the centromeres are approached, whereas transposons reciprocally increase, until they are replaced by *CEN180* (**Fig. 4A**). Gene and transposon densities are tracked closely by H3K4me3 and H3K9me2 ChIP-seq enrichment, respectively (**Fig. 4A**). H3K9me2 enrichment is observed in the centromere, although there is a reduction in the center coincident with CENH3 enrichment (**Fig. 4A**), consistent with reduced H3 occupancy caused by CENH3 replacement. Interestingly, a slight increase in H3K4me3 enrichment is observed within the centromeres, relative to the flanking pericentromeric regions (**Fig. 4A**). We observed striking biases in base composition over the centromeres, which are relatively GC-rich compared to the AT-rich chromosome arms (**Fig. 4A**).

Using our ONT sequencing data with the DeepSignal-plant algorithm (*33*), we observed dense DNA methylation across the centromeres in CG, CHG and CHH contexts (**Fig. 4A–4B**). However, CHG DNA methylation shows relatively reduced frequency within the centromeres, compared to CG methylation (**Fig. 4A**). This may reflect centromeric depletion of H3K9me2 (**Fig. 4A**), a histone modification that maintains DNA methylation in non-CG contexts (*34*). To further investigate the DNA methylation environment associated with CENH3 deposition, we performed ChIP using either

H3K9me2 or CENH3 antibodies and sequenced the purified DNA with ONT. We analyzed methylation frequency in reads that aligned to the centromeric regions and observed dense CG methylation in both read sets, but depletion of CHG and CHH methylation in the CENH3 read set relative to the H3K9me2 read set (**Fig. S13**). This further supports that replacement of H3 with CENH3 causes a decrease in maintenance of non-CG methylation within the Arabidopsis centromeres.

To investigate genetic control of DNA methylation in the centromeres, we analyzed bisulfite sequencing (BS-seq) data from wild type and eight mutants defective in CG and non-CG DNA methylation maintenance pathways (**Fig. S14**) (*34*, *35*). Centromeric non-CG methylation is eliminated in *drm1 drm2 cmt2 cmt3* mutants, and strongly reduced in *kyp suvh5 suvh6*, whereas CG methylation is intact in these backgrounds (**Fig. S14**) (*34*, *35*). Both CG and non-CG methylation in the centromeres is strongly reduced in both *ddm1* and *met1*, although the non-CG reduction is greater in *ddm1* compared with *met1* (**Fig. S14**) (*35*). Hence, centromeric CG-context methylation is relatively high compared with non-CG, and non-CG methylation shows an unexpected dependence on CG maintenance pathways.

We observed high pericentromeric ChIP-seq enrichment of the heterochromatic chromatin marks H2A.W6, H2A.W7 and H3K27me1, which are relatively depleted within the centromeres (**Fig. 4A**) (*36, 37*). The Polycomb-group modification H3K27me3 is depleted in the centromeres and found largely in the gene-rich chromosome arms (**Fig. 4A**). Enrichment of the euchromatic histone variant H2A.Z is low in the centromeres, but similar to H3K4me3, it shows a slight increase in the centromeres, relative to the pericentromeres (**Fig. 4A**). To independently characterize centromeric chromatin, we performed immunofluorescent staining of Arabidopsis nuclei for CENH3-GFP and euchromatic and heterochromatic histone modifications (**Fig. 4C, S15** and **S16**). Quantification of fluorescence intensity confirmed that heterochromatic marks are relatively depleted where CENH3-GFP is enriched (**Fig. 4C** and **S16**). Hence, the Arabidopsis centromeres show depletion of heterochromatic and enrichment of euchromatic marks, relative to the adjacent pericentromeres, consistent with a hybrid chromatin state.

Meiotic recombination, including unequal crossover and gene conversion, has been proposed to mediate centromere sequence evolution (4, 38). We mapped 2,080 meiotic crossovers from Col×Ler  $F_2$ sequencing data against the Col-CEN assembly that were resolved on average to 1,047 kbp (Fig. S17). As expected, crossovers were potently suppressed in proximity to the centromeres (Fig. 4A–4B and S17). We observed high centromeric ChIP-seq enrichment of REC8-cohesin and the HORMA domain protein ASY1, which are components of the meiotic chromosome axis (Fig. 4A) (39, 40). To investigate the potential for meiotic DSB formation within the centromeres, we aligned SPO11-1-oligo data from wild type (41). Overall, SPO11-1-oligos are low within the centromeres, although we observed an increase relative to the flanking pericentromeric heterochromatin, reminiscent of H3K4me3 and H2A.Z enrichment patterns (Fig. 4A). To investigate the role of DNA methylation, we mapped SPO11-1-oligos sequenced from the CG DNA methylation mutant met1-3 (41), which showed a gain of DSBs in proximity to the centromere (Fig. 4A-4B). To provide cytological evidence of recombination close to the centromeres, we immunostained meiocytes in early prophase I for CENH3 and V5-DMC1, which is a marker of meiotic interhomolog recombination (Fig. 4C and S18-S19). DMC1-V5 foci were observed along the chromosomes and adjacent to the surface of CENH3 foci, but not within them (Fig. 4C). Hence, despite suppression of crossovers, we observe evidence for low levels of meiotic recombination initiation associated with the centromeres, which is influenced by DNA methylation.

Finally, we analyzed chromatin and transcription around *CEN180* and *ATHILA* retrotransposons at the fine scale, and compared wild type and the DNA methylation mutant *met1-3*. CENH3 nucleosomes show a strongly phased pattern of enrichment with the *CEN180* satellites, with relative depletion in spacer regions at the start and end of the satellites (**Fig. 4D**). Interestingly, these CENH3 spacer regions also associate with elevated DNA methylation and *CEN180* variant frequency (**Fig. 4D**), consistent with CENH3-nucleosome occupancy influencing epigenetic modification and genetic divergence of satellites. In *met1*, we observed loss of CG-context DNA methylation in both the *ATHILA* and *CEN180* repeats (**Fig. 4E** and **S20**) (*42*). However, analysis of RNA-seq and siRNA-seq signal shows elevation of *ATHILA* transcripts and siRNAs in *met1*, but not within *CEN180* (**Fig. 4E** and **S20**) (*42*). The strongest RNA and siRNA increases are observed in the internal 3' regions of the *ATHILA* (**Fig. 4E** and

**S20**), which correspond to 'TSI' transcripts and easiRNA populations previously reported (43–45). This further indicates that epigenetic regulation of the *CEN180* satellites and *ATHILA* elements are distinct.

#### DISCUSSION

#### A new resource for plant centromere biology

Arabidopsis thaliana is a powerful model system for basic research and plant science, and the exceptionally accurate TAIR10 reference has served as a foundation for understanding the structure, function and evolution of eukaryotic genomes. Leveraging recent advances in sequencing technology and genome assembly methods, we have generated the Col-CEN reference genome, which matches the accuracy of TAIR10 in protein-coding regions, yet also resolves the large centromeric satellite arrays in each of the five chromosomes. By profiling chromatin and recombination distributions within the centromeres, we demonstrate that Col-CEN enables new biological insights from previously published functional genomics data, across various samples and experimental treatments. Our centromeric DNA methylation analysis also demonstrates how long-reads can resolve epigenetic patterns of repetitive DNA, highlighting the potential of complete reference assemblies for functional genomics of repeats. The Col-0 centromeres contain interspersed unique sequences that facilitate accurate assembly with long and accurate sequencing reads. However, similar to the T2T consortium, the Col-CEN assembly required extensive manual processes to polish and curate repetitive loci (9, 24, 46). We anticipate that as genome assembly becomes more automated and accurate within repeats, researchers will be able to compare complete centromere sequences across large populations and diverse species, ushering in a new era of comparative genomics and revealing how centromere diversity and evolution impact genome function.

#### Arabidopsis centromere evolution via satellite homogenization and ATHILA invasion

The Col-CEN assembly reveals the architecture of the Arabidopsis centromeres, which consist of megabase-scale CEN180 tandem arrays that are variably invaded by ATHILA retrotransposons. Extensive sequence variation is observed between the CEN180 satellites, and the majority of variant monomer sequences are private to each centromere. This is consistent with satellite homogenization occurring primarily within chromosomes. CEN180 that are the least divergent from the genome-wide consensus and with most higher-order repetition show the highest CENH3 occupancy. This suggests that CENH3 chromatin may promote recombination pathways that lead to homogenization, including DSB formation and repair via homologous recombination. For example, interhomolog strand invasion and non-crossover repair during meiosis, using allelic or non-allelic templates, has the potential to cause CEN180 gene conversion and structural change (Fig. S21). Similarly, repair and recombination using a sister chromatid may also contribute to changes to the CEN180, which could occur during either mitosis or meiosis (Fig. S21). We note that CEN180 higher-order repeats show an average length of 432 bp, which is within the range of observed Arabidopsis meiotic gene conversions (47), although we also observe large (10–100 kbp) intra-centromere duplications, for which the origin is less clear. We observe a proximity effect on divergence between CEN180 higher-order repeats, with repeat blocks further apart showing greater sequence differences. These patterns are reminiscent of human alpha-satellite higherorder repeats, although alpha-satellite HORs are longer and occur over greater physical distances (5, 12, 48, 49). As meiotic crossover repair is strongly suppressed within the centromeres, consistent with patterns across eukaryotes (38, 50-52), we do not consider unequal crossover to be a major pathway driving Arabidopsis centromere evolution. However, we propose that an ongoing, recombination-based homogenization process, occurring between allelic or non-allelic locations on the same chromosome, maintains the CEN180 library close to the consensus that is optimal for CENH3 recruitment (Fig. S21).

Aside from homogenizing recombination within the *CEN180*, the centromeres have experienced invasion by *ATHILA* retrotransposons. The ability of *ATHILA* elements and other LTR retrotransposons to insert within the centromeres is likely determined by the integrase protein. For example, chromovirus clade retrotransposons are known to target the centromere via an integrase C-terminal chromodomain (*53*), yet *ATHILA* are phylogenetically distant and our searches using the chromodomain HMM motif

(PF00385) did not identify significant hits. The Tal1 *COPIA* element from *Arabidopsis lyrata* also shows a strong insertion bias into *CEN180* arrays when expressed in *Arabidopsis thaliana* (54), despite satellite sequences varying between these species, indicating that epigenetic information may be important for targeting (55). The majority of the Col-CEN centromeric *ATHILA* elements appear young, based on LTR identity, and possess many features required for transposition, although the centromeres show striking differences in the frequency of *ATHILA* insertions, with centromeres 4 and 5 being the most invaded. *ATHILA* elements show lower CENH3 and higher H3K9me2 and CHG DNA methylation than the surrounding *CEN180*, and are associated with increased satellite divergence in adjacent regions. Therefore, *ATHILA* transposons represent a potentially disruptive influence on the genetic and epigenetic organization of the centromeres. However, transposons are widespread constituents of centromeres in diverse eukaryotes and have been found to contribute to repeat sequence and centromere protein evolution (e.g., mammalian CENP-B is derived from a Pogo DNA transposase) (56). Therefore, *ATHILA* elements may also beneficially contribute to centromere integrity and stability.

The advantage conferred to *ATHILA* elements, or other transposons, by integration within the centromeres is presently unclear, although we speculate that they may be engaged in centromere drive (57). Haig-Grafen scrambling via recombination has been proposed as a defense against such putative drive elements within the centromeres (58). For example, maize meiotic gene conversion was observed to eliminate centromeric *CRM2* retrotransposons (38). Therefore, satellite homogenization pathways may serve as a mechanism to purge *ATHILA* insertions, although in some cases this also results in transposon duplication (**Fig. S22**). A gene conversion-type mechanism may explain the 6 *ATHILA* intra-chromosome duplications that appear to have occurred post-integration. We also note that the presence of *ATHILA* solo LTRs is consistent with homologous recombination acting on the centromeric retrotransposons following integration (**Fig. S22**). Intriguingly, centromere 5 and the diverged *CEN180* array in centromere 4, show both high *ATHILA* density and a striking reduction of *CEN180* higher-order repetition. This indicates that *ATHILA* may inhibit *CEN180* homogenization, or that loss of homogenization facilitates *ATHILA* insertion, or both. We propose that each Arabidopsis centromere represents different stages in cycles of satellite homogenization and *ATHILA*-driven diversification.

These opposing forces provide both a capacity for homeostasis, and a capacity for change, that could be responsible for the paradoxically rapid evolution of centromeres. Assembly of centromeres from multiple Arabidopsis accessions, and closely related species, has the potential to reveal new insights into the wider mechanism of centromere formation and the evolutionary dynamics of *CEN180* and *ATHILA* repeat populations.

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immunocytology. NE generated the DMC1 epitope-tagged line. NH and KC sequenced DNA and contributed to the assembly. LS, JT and KS performed PacBio sequencing. TK and RM provided intellectual input. TM and MAL performed FISH. FB supervised ChIP-seq, immunofluorescence analysis and wrote the manuscript. AB performed *ATHILA* annotation and genome analysis and wrote the manuscript. TM supervised DNA sequencing, genome assembly and analysis and wrote the manuscript. MCS supervised genome assembly, validation, annotation and analysis and wrote the manuscript. IH supervised DNA sequencing, genome assembly, validation, annotation and analysis and wrote the manuscript.

Data availability: The ONT sequencing reads used for assembly are available for download at ArrayExpress accession E-MTAB-10272 (http://www.ebi.ac.uk/arrayexpress/) (Username: Reviewer\_E-MTAB-10272 Password: YVJAaVii). The PacBio HiFi reads are available for download at European Nucleotide Archive accession number **PRJEB46164** (https://www.ebi.ac.uk/ena/browser/view/PRJEB46164). All data, code and materials are available in the manuscript or the supplementary materials and at https://github.com/schatzlab/Col-CEN.



Figure 1. Complete assembly of the Arabidopsis centromeres. A. Genome-wide circos plot of the Col-CEN assembly. Quantitative tracks (c-i) are aggregated in 100-kbp bins and independent y-axis labels are given as (low tick value, mid tick value, high tick value, unit of measurement): (a) chromosome labels with centromeres shown in red; (b) genomic features showing telomeres in blue, 45S rDNA in yellow, 5S rDNA in black, and the chromosome 2 mitochondrial insertion in pink; (c) genes (0, 25, 51, # of genes); (d) transposable elements (0, 84, 167, # of transposable elements); (e) Col×Ler F<sub>2</sub> crossovers (0, 7, 14, # of crossovers); (f) CENH3 (-0.5, 0, 3, log<sub>2</sub>(ChIP/input)); (g) H3K9me2 (-0.6, 0, 2, log<sub>2</sub>(ChIP/input)); (h) CG methylation (0, 47, 95, % methylated); (i) CHG methylation (0, 28, 56, % methylated); (i) CHH methylation (0, 7, 13, % methylated). B. Plot showing syntenic alignments between the TAIR10 and Col-CEN assemblies. C. Col-CEN genome assembly ideogram with annotated chromosome landmarks (not drawn to scale). D. CENH3 log<sub>2</sub>(ChIP/input) (black) plotted over centromeres 1 and 4 (19). CEN180 density per 10 kbp is plotted for forward (red) or reverse (blue) strand orientations. ATHILA retrotransposons are indicated by purple ticks on the xaxis. Beneath are heatmaps showing pairwise % sequence identity values of all non-overlapping 5-kbp regions. An image of a FISH-stained pachytene chromosome 1 is shown above, which has been probed with upper-arm-specific BACs (green), ATHILA (BAC T1J24, purple), CEN180 (pAL1, blue), the telomeric repeat (TTTAGGG)<sub>n</sub> (green) and bottom-arm-specific BACs (yellow). E. Dotplot analysis comparing the five centromere regions, using a search window of 120 or 178 bp. Red and blue shading indicate detection of similarity on the same or opposite strands, respectively. F. Col-0 pachytene-stage chromosomes stained with DAPI (black) and CEN180- $\alpha$  (red) and CEN180- $\beta$  (purple) FISH probes. Chromosome 1 specific BACs (green) were also used for FISH. The location of chromosome 1 is indicated in the last image by the arrow. The scale bar represents  $10 \mu M$ .



**Figure 2. The Arabidopsis** *CEN180* **satellite repeat library. A.** Histograms of *CEN180* monomer lengths (bp), and variant distances relative to the genome-wide consensus. Mean values are shown by the red dotted line. **B.** As for A, but showing widths of *CEN180* higher order repeat (HOR) blocks (monomers, 'mers'), and the distance between HOR blocks (kbp). **C.** Heatmap of a representative

satellite region within centromere 2, shaded according to pairwise variants between *CEN180*. **D.** Circos plot showing: (i) *GYPSY* LTR transposon density; (ii) *CEN180* density; (iii) centromeric *ATHILA* rainfall plot; (iv) *CEN180* density grouped by decreasing CENH3 log<sub>2</sub>(ChIP/input) (red=high; navy=low); (v) *CEN180* density grouped by decreasing higher-order repetition (red=high; navy=low); (vi) *CEN180* density grouped by decreasing higher-order repetition (red=high; navy=low); (vi) *CEN180* grouped by decreasing variant distance (red=high; navy=low); and (vii) CENH3 log<sub>2</sub>(ChIP/input) (purple), across the centromere regions. **E.** *CEN180* were divided into quintiles according to CENH3 log<sub>2</sub>(ChIP/input) and mean values for each group with 95% confidence intervals plotted. The same groups were analyzed for *CEN180* variant distance (red), higher-order repetition (blue) and CG-context DNA methylation (purple). **F.** Plot of the distance between pairs of HOR blocks (kbp) and divergence (variants/monomers) between the HOR block sequences. **G.** Plots of CENH3 log<sub>2</sub>(ChIP/input) (black) across the centromeres, compared to *CEN180* higher-order repetition on forward (red) or reverse (blue) strands. A heatmap is shown beneath that is shaded according to the density of higher-order repeats.



**Figure 3. Invasion of the Arabidopsis centromeres by** *ATHILA* **retrotransposons. A.** Dotplot of centromeric *ATHILA* retrotransposons using a search window of 50 bp. Red and blue indicate forward-

and reverse-strand similarity. The elements assigned to different ATHILA subfamilies and solo LTRs are indicated. B. Maximum likelihood phylogenetic tree of the 111 intact ATHILA elements based on their full-length sequence. Branches are color-coded according to ATHILA subfamily, and stars at the branch tips indicate presence inside (white) or outside (black) the centromeres. C. An annotated map of a representative ATHILA6B element located on chromosome 5, with LTRs (blue) and the core protein domains (red) highlighted. D. Histograms of LTR percent sequence identity for centromeric ATHILA elements (n=53), compared to ATHILA elements outside of the centromeres (n=58). Mean values are indicated by the red dashed lines. E. Meta-profiles of CENH3 (orange) and H3K9me2 (blue) ChIP-seq signals (log<sub>2</sub>(ChIP/input)) around CEN180 (n=66,131), centromeric intact ATHILA (n=53), ATHILA located outside the centromeres (n=58), GYPSY retrotransposons in the genome (n=3,979), and random positions (n=66,131). Shaded ribbons represent 95% confidence intervals for windowed mean values. F. As for E, but analyzing ONT-derived percent DNA methylation in CG (dark blue), CHG (blue) and CHH (light blue) contexts. G. Meta-profiles of the number of CEN180 sequence edits (insertions, deletions and substitutions relative to the CEN180 consensus), normalized by CEN180 presence/absence, in positions surrounding CEN180 gaps that containing ATHILA sequences (n=65), or random positions (n=65). All edits (dark blue) are analyzed, in addition to substitutions (blue), indels (light blue), insertions (light green), deletions (dark green), transitions (pink) and transversions (orange). Shaded ribbons represent 95% confidence intervals for windowed mean values. H. A Col-0 pachytene-stage chromosome spread was stained with DAPI (black) and with an ATHILA6A/6B GAG FISH probe (red) and chromosome 5 specific BACs (green). The scale bar represents 10  $\mu$ M.



**Figure 4. Epigenetic organization and meiotic recombination within the centromeres. A.** Quantification of genomic features plotted along chromosome arms that were proportionally scaled between the telomeres (*TEL*) and centromere midpoint (*CEN*), which was defined by maximum CENH3

ChIP-seq log<sub>2</sub>(ChIP/input) enrichment. Data analyzed were gene, transposon and CEN180 density, CENH3, H3K4me3, H3K9me2, H2A.W6, H2A.W7, H2A.Z, H3K27me1, H3K27me3, REC8 and ASY1 ChIP-seq (log<sub>2</sub>(ChIP/input)), % AT and GC base composition, DNA methylation, SPO11-1oligos (in wild type and *met1*) and crossovers (see **Table S7** for information on data sets). **B.** Plot quantifying crossovers (red), % CG DNA methylation (pink), CENH3 (blue), SPO11-1-oligos in wild type and *met1*, and *CEN180* density along centromere 2 (*CEN2*). C. An Arabidopsis Col-0 interphase nucleus immunostained for the heterochromatic mark H3K9me2 (magenta) and CENH3-GFP (green). The white line indicates the area of the confocal section used for the intensity plot shown on the right. The scale bar represents 5  $\mu$ M. Beneath is shown a male meiocyte in early prophase I immunostained for CENH3 (red) and V5-DMC1 (green). Scale bars are 10 μM (upper row) and 1 μM (lower row). D. Plots of CENH3 ChIP enrichment (grey), DNA methylation in CG (blue), CHG (green) and CHH (red) contexts and CEN180 variants (purple), averaged over windows centered on all CEN180 starts. The red lines show 178-bp increments. E. Meta-profiles of CG-context DNA methylation in wild type (green) or met1 (pink/purple) (42), RNA-seq in wild type (green) and met1 (pink/purple) (42), and siRNA-seq in wild type (green) and met1 (pink/purple) (42) around CEN180 (n=66,131), centromeric intact ATHILA (n=53), ATHILA located outside the centromeres (n=58), GYPSY retrotransposons in the genome (n=3,979) and random positions (n=66,131). Shaded ribbons represent 95% confidence intervals for windowed mean values.

# **Supplementary Materials**

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#### **Materials and Methods**

#### Genomic DNA extraction and ONT and PacBio HiFi sequencing

For genomic DNA extraction associated with ONT sequencing, 3 week old Col-0 seedlings were grown on  $\frac{1}{2}$  MS media and 1% sucrose and kept in the dark for 48 hours prior to harvesting. Approximately 10 g of tissue was used per 200 ml of MPD-Based Extraction Buffer pH 6 (MEB). Tissue was flash frozen and ground tissue in liquid nitrogen, using a pestle and mortar, and resuspended in 200 ml MEB. Ground tissue was thawed in MEB with frequent stirring. The homogenate was forced through 4 layers of miracloth, and then filtering again through 4 layers of fresh miracloth by gravity. 20% Triton x-100 was added to a final concentration of 0.5% on ice, followed by incubation with agitation on ice for 30 minutes. The suspension was centrifuged at 800g for 20 minutes at 4°C. The supernatant was removed and the pellet resuspended using a paintbrush in 10 ml 2-methyl-2,4 pentanediol buffer pH 7.0 (MPDB). The suspension was centrifuged at 650g for 20 minutes at 4°C. The supernatant was removed and the pellet was washed with 10 ml of MPDB. Washing and centrifugation was repeated until the pellet appeared white and was finally resuspended in a minimal volume of MPDB. From this point onwards all transfers were performed using wide bore pipette tips. 5 ml CTAB buffer was added to the nuclei pellet and mixed via gentle inversion, followed by incubation at 60°C until full lysis had occurred, taking between 30 minutes and 2 hours. An equal volume of chloroform was added and incubated on a rocking platform, with a speed of 18 cycles per minute, for 30 minutes, followed by centrifugation at 3000g for 10 minutes. An equal volume of phenol/chloroform/isoamyl alcohol (PCI, 25:24:1) was

added to the lysate, followed by incubation on a rocking platform (18 cycles per minute) for 30 minutes. The lysate was centrifuged at 3000g for 10 minutes and the upper aqueous phase was transferred into a fresh tube. The PCI extraction was then repeated. The extraction was then repeated using only chloroform.  $1/10^{\text{th}}$  volume of 3M Sodium Acetate was added to the lysate and mixed by gentle inversion. Two volumes of ice cold ethanol were added and mixed by inversion. DNA was precipitated at -20°C for 48 hours. The precipitated DNA was removed using a glass hook and washed three times in fresh 70% ethanol. The DNA was dissolved in 120 µl of 10 mM Tris-Cl (pH 8.5).

Approximately 5 µg of DNA was size selected to be >30 kbp, using the BluePippin<sup>TM</sup> Size-Selection System (Sage Science) and the 0.75% DF Marker U1 cassette definition, with Range mode and BP start set at 30,000 bp. Library preparation followed the Nanopore SQK-LSK109 protocol and kit. Approximately 1.2-1.5 µg of size-selected DNA in a volume of 48 µl was used for library preparation. DNA was nic-repaired and end-prepped by the addition of 3.5 µl of NEBNext FFPE Buffer and NEBNext Ultra II End Prep Reaction Buffer, followed by 2 µl of NEBNext DNA Repair Mix and 3 µl NEBNext Ultra II End Prep Enzyme Mix (New England Biolab, E7180S), with incubation for 30 minutes at 20°C, followed by 30 minutes at 65°C. The sample was cleaned using 1×volume AMPure XP beads and eluted in 61 µl of nuclease-free water. Adapters were ligated at room temperature using 25 µl Ligation Buffer, 10 µl NEBNext T4 DNA Ligase and 5 µl Adapter Mix for 2 hours. The library was cleaned with 0.4×volume AMPure XP beads, washed using ONT Long Fragment buffer and eluted in 15 µl elution buffer.

For genomic DNA associated with PacBio HiFi sequencing, Col-0 plants were grown at the Max Planck Institute for Plant Breeding Research, Cologne, Germany. DNA extraction (from an individual plant), library preparation and DNA sequencing was performed at the Max Planck Genome Center, Cologne, Germany. High molecular weight DNA was isolated from 1.5 gram of vegetative material with a NucleoBond HMW DNA kit (Macherey Nagel). Quality was assessed with a FEMTOpulse device (Agilent) and quantity measured by fluorometry Quantus (Promega). A HiFi library was then prepared
according to the manual "Procedure & Checklist - Preparing HiFi SMRTbell® Libraries using SMRTbell Express Template Prep Kit 2.0" with initial DNA fragmentation by g-Tubes (Covaris) and final library size binning by SageELF (Sage Science). Size distribution was again controlled by FEMTOpulse (Agilent). The size-selected library was sequenced on one SMRTcell on a Sequel II device with Binding kit 2.0 and Sequel II Sequencing Kit 2.0 for 30 hours.

#### **Col-CEN** genome assembly

Libraries were sequenced on 6 ONT R9 flow cells and 1 ONT R10 flow cell, and the resulting .fast5 files were basecalled with Guppy (v4.0.15), using the dna\_r9.4.1\_450bps\_hac.cfg and dna\_r10.3\_450bps\_hac.cfg configurations, respectively. This yielded a total of 73.6 Gb of sequence (~613× total coverage). The fastq files of ONT reads used for genome assembly are available for download at ArrayExpress accession E-MTAB-10272 (http://www.ebi.ac.uk/arrayexpress/). We trimmed adapters using Porechop (v0.2.4) and filtered for read lengths greater than 30 kbp and mean read quality scores >90%, using Filtlong (v0.2.0) (https://github.com/rrwick/Filtlong), which yielded 436,146 reads with a mean length of 43.9 kbp (19.15 Gbp), equivalent to  $161 \times$  coverage of the TAIR10 genome with ~55x coverage of ultra-long reads (>50 kbp). Flye (version 2.7) was used to assemble the reads, specifying a minimum read overlap of 10 kbp and a *k*-mer size of 17 (*59*).

#### **Contig screen**

We performed a comprehensive contig screen using methods inspired by the Vertebrate Genomes Project (VGP), though adapted for an inbred plant genome (46). We first aligned Flye contigs to the Columbia reference chloroplast (GenBank accession NC\_000932.1) (60), and mitochondria (GenBank accession NC\_037304.1) (61) genomes with Minimap2 (v2.17-r941, -x asm5) (62). Contigs with at least 50% of their bases covered by alignments were considered to be chloroplast or mitochondria genome sequences and were removed from the assembly.

We next used BLAST to screen for contigs representing bacterial contamination. We first masked the Flye assembly with windowmasker (v1.0.0, -mk\_counts -genome\_size 131405362) (63). We then

aligned the Flye contigs to all RefSeq bacterial genomes (downloaded on 2020/05/21) with megablast (v2.5.0, -outfmt "6 std score"), providing the windowmasker annotations with "-window\_masker\_db" (64). We removed BLAST alignments with an E value greater than or equal to 0.0001, a score less than 500, and a Percent Identity less than 98%, and any contigs (four in total) with remaining alignments were manually inspected. Two of the four contigs were already identified as being chloroplast or mitochondria sequence and the other two were clearly nuclear contigs, so we determined that no contigs were derived from bacterial contaminants.

After removing chloroplast and mitochondria contigs, we performed one final screen to remove contigs with low read support. We aligned ONT reads (>=40 kbp) to the contigs with Minimap2 (v2.17-r941, - x map-ont) and removed any contigs (one in total) with more than 50% of its bases covered by fewer than 15 reads. Though we did not use its standard pipeline, we made use of purge\_dups scripts for this analysis (*65*). After screening, the assembly consisted of 10 contigs with an N50 of 22,078,741 bp.

#### **Contig scaffolding**

Though the five Columbia chromosomes were represented by only 10 contigs, we used homology-based scaffolding to order and orient contigs, assign chromosome labels, and orient pseudomolecules to match the orientation of TAIR10 chromosomes. We ran RagTag (v1.0.1, --debug --aligner=nucmer --nucmer-params='--maxmatch -1 100 -c 500') using TAIR10 as the reference genome, but excluding ChrC and ChrM (-e) (*66*, *67*). Three small contigs (3,200, 90,237 and 8,728 bp) consisting of low complexity sequence were not ordered and oriented and were removed from the assembly. After scaffolding, the 131,388,895 bp assembly was represented in five pseudomolecules corresponding to the five chromosomes of the Columbia genome. Chromosome 1 was gapless, while the other chromosomes contained one to four 100 bp gaps each (9 in total).

#### Initial pseudomolecule polishing and gap filling

We corrected mis-assemblies and filled gaps in the Columbia pseudomolecules with two rounds of Medaka (v1.2.1) ONT polishing (https://github.com/nanoporetech/medaka). For the first round of

polishing, we aligned R9 ONT reads (>=50 kbp) to the pseudomolecules with mini\_align (minimap2 v2.17-r941, -m). To avoid overcorrection in the centromere satellite sequences, we performed "marker-assisted filtering" to remove alignments not anchored in putatively unique sequences (*12, 24*) (https://github.com/malonge/T2T-Polish). We defined "marker" *k*-mers as 21-mers that occurred once in the assembly and between 14 and 46 times (inclusive) in the Illumina reads. The first round of polishing was completed using `medaka consensus` (--model r941\_min\_high\_g360 --batch\_size 200) and `medaka stitch`. The second round of polishing was performed as for the first round, except we aligned all R10 reads instead of R9 reads and the `medaka consensus` model was set to "r103\_min\_high\_g360". As a result of ONT polishing, the assembly improved from a QV of 32.38 to 33.17 and 34.12 after the first and second rounds, respectively (*28*). After medaka polishing, the assembly contained only a single gap on chromosome 2.

Long-read ONT polishing was followed by short-read polishing of non-centromeres with DeepVariant (68). We first aligned Col-0 genomic DNA Illumina reads to the pseudomolecules with bwa mem (v0.7.17-r1198-dirty) and we compressed and sorted alignments with samtools (v1.10) (69, 70). We then created a VCF file of potential polishing edits with DeepVariant (v1.1.0, -- model\_type=WGS),"bcftools view" (v1.11, -e 'type="ref"' -i 'QUAL>1 && (GT="AA" || GT="Aa")' ) and "bcftools norm". To avoid error-prone short-read polishing in the centromeres, we used Bedtools to remove polishing edits within the centromeres and we used BCFtools to derive a final consensus FASTA file (71, 72). Though short-read polishing did not alter the centromeres, it improved the overall assembly QV to 41.4616.

#### **Telomere patching**

We locally re-assembled and patched telomeric sequences for the 8 Columbia telomeres not adjacent to NORs (all but the beginning of chromosomes 2 and 4). We aligned all R9 reads to the TAIR10 reference with Winnowmap (v1.11, k=15, --MD -ax map-ont) and for each telomere, we collected all reads that aligned once to within 50 bp of the chromosome terminus (*16*). Using Bowtie (*73*) (v1.3.0, -S --all -v 0), we counted the occurrences of the telomeric repeat motif ('CCCTAAA') in each read, and the read

with the most occurrences was designated as the "reference" and all other reads were designated as the "query". Local re-assembly was completed by aligning the query reads to the reference read and computing a consensus with `medaka\_consensus` (v1.2.1, -m r941\_min\_high\_g360). To patch these telomere consensus sequences into the Columbia pseudomolecules, we identified the terminal BAC sequences for each of the 8 chromosome arms. For each chromosome arm, we aligned the terminal BAC sequence to the Columbia pseudomolecules and the telomere consensus sequence with Nucmer (v3.1, --maxmatch). Using these alignment coordinates, the consensus sequences were manually patched such that everything after the terminal BAC sequence was replaced with telomere consensus sequence. Telomeres were then manually confirmed to be structurally valid.

#### Assembly curation and preparation

After polishing and telomere patching, we performed final curation steps to correct lingering misassemblies and screen for contamination. First, while it was not straightforward to fill the remaining chromosome 2 gap *de novo*, we were able to replace the gap locus with the corresponding region in TAIR10. We found two BAC sequences flanking the gap locus that aligned concordantly to both the Col-0 pseudomolecules and TAIR10. These BAC contigs were aligned to the pseudomolecules and TAIR10 with Nucmer (v3.1, --maxmatch -1 250 -c 500) and the gap locus between the BAC contigs in the Columbia pseudomolecules was replaced with the corresponding TAIR10 locus between the BAC contigs.

To identify and correct structural mis-assemblies, we aligned Columbia long-reads to the Columbia pseudomolecules and called structural variants (SVs). First, we used Bedtools `random` (v2.29.2, -1 100000 -n 50000 -seed 23) to simulate 50,000 100 kbp exact reads from TAIR10. These reads, along with R9 (>=50 kbp) and R10 Columbia reads were aligned to the Columbia pseudomolecules with Winnowmap (v1.11, k=15, "--MD -ax map-pb" for TAIR10 reads and "--MD -ax map-ont" for ONT reads). After compressing and sorted alignments with samtools (v1.10), Sniffles (v1.0.12, -d 100 -n -1 -s 3) was used to infer SVs from each of the alignments (74). SVs with fewer than 30% of reads supporting the ALT allele were removed and the three resulting VCF files were merged with Jasmine

(v1.0.10, max\_dist=500 spec\_reads=3 --output\_genotypes) (75). There were a total of three variants called by all three read sets, including two deletions and one insertion that we corrected. REF and ALT alleles for these SVs were manually refined and validated, and ALT alleles were incorporated into the pseudomolecules using `bcftools consensus`.

Next, we manually inspected all gaps filled by Medaka and found that a 181 bp region containing a 100 bp gap on chromosome 5 was incorrectly replaced with 103 bp of sequence and we manually replaced the filled sequence with the original gap locus. This ultimately produced the Col-CEN v1.1 assembly. We used VecScreen to do a final contamination screen. We first aligned the Columbia pseudomolecules to the VecScreen database with blastn (v2.5.0, -task blastn -reward 1 -penalty -5 -gapopen 3 -gapextend 3 -dust yes -soft\_masking true -evalue 700 -searchsp 175000000000 -outfmt "6 std score"). The BLAST alignments did not yield any "moderate" or "strong" matches to the database, so we determined that there was no contamination.

#### Additional polishing and generation of the Col-CEN v1.2 assembly

To further polish the Col-CEN v1.1 assembly, we aligned all HiFi reads that were at least 16 kbp long to the Col-CEN v1.1 assembly with Winnowmap2 (v2.0, k=15 greater-than distinct=0.9998 --MD -ax map-pb) and we filtered alignments with Samtools "view" (v1.10, -F 256) (*16*, *70*). We then used "falconc bam-filter-clipped", a part of the IPA package, to remove chimeric read alignments (-t -F 0x104) (https://github.com/PacificBiosciences/pbipa). Using these filtered alignments, we polished the Col-CEN v1.1 assembly with a special branch of Racon that outputs polishing edits in VCF format (v1.6.0, -L -u) (https://github.com/isovic/racon/tree/liftover) (*76*). Polishing edits were then filtered with Merfin, using 21-mers derived from the Col-O Illumina reads (-peak 30) (*77*) and incorporated into the assembly with BCFtools "consensus" (*71*).

To identify and correct putative larger mis-assemblies with a second, independent method, we assembled all HiFi reads at least 16 kbp long with Hifiasm (v0.15-r327, -l0), and aligned the resulting primary contigs to the Racon polished assembly with minimap2 (v2.20-r1061, --cs -cx asm5). We called

variants with paftools "call" and manually inspected all variants larger than 1 kbp in IGV (https://github.com/lh3/minimap2/tree/master/misc) (62). Ultimately, two sequences were inserted into the Racon assembly, ultimately producing the Col-CEN v1.2 assembly. The Col-CEN v1.2 assembly contained five pseudomolecules, two missing telomeres, and partially resolved NOR sequence at the beginning of chromosomes 2 and 4. Chromosomes 1, 3 and 5 were completely sequence resolved from telomere-to-telomere (T2T). The final Col-CEN v1.2 assembly FASTA file includes these 5 pseudomolecules and the Columbia chloroplast and mitochondria reference genomes.

To catalog variation between Col-0 lab strains, heterozygous loci, or potential lingering misassemblies, we aligned Col-0 reads to Col-CEN v1.2 and called variants. To call small variants, we aligned all HiFi reads at least 16 kbp long to the Col-CEN v1.2 assembly with Winnowmap2 (v2.0, k=15 greater-than distinct=0.9998 --MD -ax map-pb) and called variants with DeepVariant (v1.1.0, -model type=PACBIO). The same HiFi alignments were used to call SVs with Sniffles (v1.0.12, -d 50 -n -1 -s 10) and variants with less than 30% of reads supporting the ALT allele were removed. The same process was used to call SVs with ONT data (Winnowmap v2.0) (k=15 greater-than distinct=0.9998 --MD -ax map-ont). The resulting VCF files are available on GitHub (https://github.com/schatzlab/Col-CEN). During analysis, we uncovered two potentially misassembled loci, though plausible corrections apparent. We have listed these loci in an "issues" file on GitHub were not (https://github.com/schatzlab/Col-CEN). These, and potential future issues identified by ourselves or the community, will be considered in future assembly updates.

For assembly validation, we aligned Hi-C reads to Col-CEN with bwa mem (v0.7.17-r1198-dirty) and processed the alignments with the Arima mapping pipeline (https://github.com/ArimaGenomics/mapping\_pipeline) (http://broadinstitute.github.io/picard/) (78). Hi-C heatmaps were made with Cooler and HiGlass (Cooler v0.8.10, 50 kbp resolution) (79) (https://higlass.io/).

#### **Genome annotation**

Genes were lifted-over from TAIR10 with Liftoff (v1.5.1, -copies -a 1 -s 1) (80). Since ChrC and ChrM were directly copied from TAIR10, their lift-over genes were replaced with their original TAIR10 annotations. We inspected every TAIR10 gene that did not lift over to provide an explanation for the discrepancy. All presence/absence variable genes are listed in Table S2 and all missing genes (including for reasons other than genuine biological variation) are documented on GitHub (https://github.com/schatzlab/Col-CEN). We also inspected every gene that lifted over in multiple copies. All copy-number-variable genes are listed in Table S3 and all genes that lifted over in multiple copies (including for reasons other than genuine biological variation) are listed on GitHub (https://github.com/schatzlab/Col-CEN). We used EDTA (v1.9.6, --sensitive 1 --anno 1 --evaluate 1) to perform de novo transposable element (TE) annotation, providing transcripts with "--cds" and the TAIR10 TE library with "--curatedlib" (81, 82). The TE annotation was supplemented with a manual annotation of centromere gaps using dotplot analysis and further manual annotation of the centromeric ATHILA elements (see section below). We used LASTZ to identify regions with similarity to 5S, 45S rDNA and the mitochondrial genome. To generate similarity heatmaps, the centromere region was divided into adjacent 5 kbp regions, which were compared using the pairwiseAlignment (type='global') and pid functions in R, using the Biostrings library. Sequences were compared in forward and reverse directions, and the highest percent sequence identity value kept. These values were then plotted in the heatmap.

#### **CEN180** repeat annotation

To identify repetitive regions, we divided the genome assembly into adjacent 1 kbp windows. In each window, for each position, we defined 12-mers and exactly matched these sequences to the rest of the window. We identified windows where the proportion of non-unique 12-mers was greater than 10%, and merged contiguous windows that were above this threshold. For each region, we generated a histogram of the distances between 12-mers to test for periodic repeats. For example, if a region contains an arrayed tandem repeat of monomer size N, then a histogram of the 12-mer distances will show peaks at values N, N×2, N×3 ... . The N value was obtained for each region, using the most frequent 12-mer distance. Next, 5 sequences of length N were randomly chosen from within the region and matched

back to the sequence using the R function matchPattern (max.mismatch=N/3 with.indels=T). For each set of matches we identified overlapping repeats. If the overlap was less than 10 nucleotides, the overlap was divided at the midpoint between the repeats. If the overlap was 10 nucleotides or greater, the larger repeat was kept. The set of non-overlapping matches with the highest number was kept for further analysis. These sequence matches were aligned using mafft (--retree 2 --inputorder) (*83*), and a consensus repeat monomer was derived from the multiple sequence alignment. This consensus sequence was matched back to the region using matchPattern (max.mismatch=N/3 with.indels=T), and overlaps were treated in the same way.

Our approach identified 66,131 *CEN180* repeats with a mean length of 178 bp. The set of unique *CEN180* sequences (n=22,440) were aligned using mafft (--sparsescore 1000 --inputorder) (83). A consensus sequence was generated from the multiple sequence alignment, which was:

5'-

## AGTATAAGAACTTAAACCGCAACCCGATCTTAAAAAGCCTAAGTAGTGTTTCCTTGTTAGA AGACACAAAGCCAAAGACTCATATGGACTTTGGCTACACCATGAAAGCTTTGAGAAGCA AGAAGAAGGTTGGTTAGTGTTTTGGAGTCGAATATGACTTGATGTCATGTGTATGATTG-

3'. In order to analyze *CEN180* diversity, for each position of the multiple sequence alignment (809 positions), we calculated the proportion of A, T, G, C and gaps. The alignment for each monomer at each position was then compared to these proportions and used to calculate a variant distance for the monomer. For example, if a monomer had an A in the alignment at a given position, and the overall proportion of A at that position was 0.7, the variant distance for that monomer would increase by 1-0.7. This was repeated for each position of the alignment, for each monomer. This 'weighted' variant distance was used to assess how similar a given *CEN180* monomer is to the genome-wide consensus. Alternatively, to compare pairwise differences between two specific monomers, the two sequences were compared along the length of the multiple sequence alignment and each instance of disagreement counted to give a 'pairwise' variant score.

To identify higher order repeats (HORs) in a head-to-tail (tandem) orientation, each monomer was taken in turn and compared to all others using a matrix of pairwise variant scores. If a pair of monomers had a variant score of 5 or less, and were on the same strand, they were considered a match. For each match, monomers were extended by +1 unit in the same direction on the chromosome, and these were again compared for pairwise variants. This process was repeated until the next monomers had a pairwise variant score higher than threshold, or the repeats were on opposite strands, or the end of the array was reached, with these conditions defining the end of the HOR. We also searched for repeats in head-tohead (inverted) orientation, which was identical apart from that repeats must be on opposite strands, and when monomers are extended to search for HORs, one is extended +1 position along the chromosome, whereas the other decreases -1. HORs were defined for each instance of 2 or more consecutive monomer matches. We define each HOR as consisting of block1 and block2 of CEN180 monomers. The size of each block was recorded, in terms of monomers and base pairs, in addition to the distance between the block start coordinates. Cumulative pairwise variants per CEN180 monomer were also calculated between each pair of blocks to provide a 'block' variant score. To measure higher order repetition of each monomer, we summed the HOR block sizes in mers, such that if a monomer was represented in three 5-mer blocks, it would score 15.

#### ATHILA annotation

To resolve the sequence of the centromeric *ATHILA* elements, we used LTRharvest (*84*) to complement the EDTA run that was used for the annotation of all Arabidopsis TEs (see above). We ran LTRharvest three times using 'normal', 'strict' and 'very strict' parameters. The parameters were gradually adjusted to allow us to capture the full-length sequence of the *ATHILA* subfamilies, based on older studies that reported the total and LTR lengths of intact *ATHILA* elements (*30*). These parameters were -maxlenltr 2500 -miniltrlen 400 -mindistltr 2000 -maxdistltr 20000 -similar 75 -mintsd 0 -motif TGCA -motifmis 1 for the 'normal' run; -maxlenltr 2000 -minilenltr 1000 -mindistltr 4000 -maxdistltr 16000 -similar 80 -mintsd 3 -motif TGCA -motifmis 1 for the 'strict' run; and -maxlenltr 2100 -minilenltr 1100 -mindistltr 5000 -maxdistltr 14000 -similar 85 -mintsd 4 -motif TGCA -motifmis 1 -vic 20 for the 'very strict' run. Coordinates of predicted intact elements from EDTA, LTRharvest and the manual dotplot annotation

of centromeric TEs were merged and sequences aligned using mafft (85). Through these steps, we were able to pinpoint with base-pair resolution the external junctions of every ATHILA element, and the internal junctions of the LTRs with the internal domain (5'-LTR with PBS; PPT with 3'-LTR). Overall, we identified 111 intact elements, 53 inside and 58 outside of the centromeres, of which 43 (81%) and 40 (69%) respectively have a detectable target site duplication (TSD), 20 fragmented ATHILA and 12 solo LTRs (10 with a TSD, 83%) (Table S6). We further identified open reading frames (minimum 300 bp) in the internal domain of the intact elements using getorf in EMBOSS (86), and the core domains of the gag and pol genes by running HMMER v3.3.2 (http://hmmer.org/) (-E 0.001 --domE 0.001) and using a collection of Hidden Markov Models (HMMs) downloaded from Pfam (http://pfam.xfam.org/) that describe the genes of GYPSY LTR retrotransposons: PF03732 for gag; PF13650, PF08284, PF13975 and PF09668 for protease; PF00078 for reverse transcriptase; PF17917, PF17919 and PF13456 for RNase-H: PF00665, PF13683, PF17921, PF02022, PF09337 and PF00552 for integrase; PF03078 for an ATHILA-specific domain. Given that many ATHILA subfamilies do not appear to contain the core domains of reverse transcriptase, RNase-H and integrase (Table S4), as these are described by the Pfam models, we used the full-length sequence of the intact elements to examine their phylogenetic relationships. The multiple alignment file was produced using mafft with the G-INS-i parameter (85), and FastTree (-nt) to generate the maximum likelihood tree (87). The tree was visualized and annotated with FigTree (http://tree.bio.ed.ac.uk/software/figtree/).

#### **ONT DNA methylation analysis**

To identify CG, CHG and CHH methylation contexts we used DeepSignal-plant (v. 0.1) (*33*), which uses a deep-learning method based on bidirectional recurrent neural network (BRNN) with long short-term memory (LSTM) units to detect DNA 5mC methylation. R9 reads were filtered for length and accuracy using Filtlong (v0.2.0) (--min\_mean\_q 90, --min\_length 30000. https://github.com/rrwick/Filtlong). Basecalled read sequence was annotated onto corresponding .fast5 files, and re-squiggled using Tombo (v 1.5.1). Methylation prediction for the CG, CHG, and CHH contexts were called using DeepSignal-plant using the respective models: model.dp2.CG.arabnrice2-1\_R9.4plus\_tem.bn13\_sn16.balance.both\_bilstm.b13\_s16\_epoch6.ckpt, model.dp2.CHG.arabnrice2-

1\_R9.4plus\_tem.bn13\_sn16.denoise\_sig1nal\_bilstm.both\_bilstm.b13\_s16\_epoch4.ckpt model.dp2.CHH.arabnrice2-

1\_R9.4plus\_tem.bn13\_sn16.denoise\_signal\_bilstm.both\_bilstm.b13\_s16\_epoch7.ckpt.

The script call\_modification\_frequency.py provided in the DeepSignal-plant package was then used to generate the methylation frequency at each CG, CHG and CHH site.

To identify CG methylation in Nanopore reads we also used Nanopolish (v 0.13.2), which uses a Hidden Markov model on the nanopore current signal to distinguish 5-methylcytosine from unmethylated cytosine. Reads were first filtered for length and accuracy using Filtlong (v0.2.0) (--min\_mean\_q 95, - -min\_length 15000. https://github.com/rrwick/Filtlong). The subset was then indexed to the fast5 files, and aligned to the genome using Winnowmap (v1.11, -ax map-ont). The read fastq, alignment bam, and fast5 files were used as an input to the Nanopolish call-methylation function. The script calculate\_methylation\_frequency.py provided in the Nanopolish package was then used to generate the methylation frequency at each CG containing *k*-mer.

#### **Bionano optical mapping**

DNA was extracted following Bionano's Plant DNA Isolation Kit (#80003) and protocol. Isolated DNA was labeled with Bionano's Direct Label and Stain Kit (DLS #80005) and samples were run on a Saphyr chip and analyzed with BionanoAccess software v1.6, Bionano Tools v1.6 and Bionano Solve v3.6\_09252020. Data generation reached 2,290 Gb equating to roughly 1,523× coverage after quality filtering for molecules containing at least 10 labels per molecule (read). *De novo* assembly of the Bionano data was performed with default assembly settings resulting in 19 contigs for a total assembly length of 132.961 Mbp. Further comparison of the Bionano contig maps was made with the Col-CEN v1.2 genome assembly. Bionano maps and molecules support the Col-CEN genome assembly where Bionano maps are capable of alignment. However, due to a lack of labelling sites, the centromere sequences generally result in breakage of the Bionano maps.

#### Chromatin immunoprecipitation and sequencing (ChIP-seq)

Approximately 12 grams of 2 week old Col-0 seedlings were ground in liquid nitrogen. Nuclei were isolated in nuclei isolation buffer (1 M sucrose, 60 mM HEPES pH 8.0, 0.6% Triton X-100, 5 mM KCl, 5 mM MgCl<sub>2</sub>, 5 mM EDTA, 0.4 mM PMSF, 1 mM pepstatin-A, 1×protease inhibitor cocktail), and crosslinked in 1% formaldehyde at room temperature for 25 minutes. The crosslinking reaction was quenched with 125 mM glycine and incubated at room temperature for a further 25 minutes. The nuclei were purified from cellular debris via two rounds of filtration through one layer of Miracloth and centrifuged at 2,500g for 25 minutes at 4 °C. The nuclei pellet was resuspended in EB2 buffer (0.25 M sucrose, 1% Triton X-100, 10 mM Tris-HCl pH 8.0, 10 mM MgCl<sub>2</sub>, 1 mM EDTA, 5 mM DTT, 0.1 mM PMSF, 1 mM pepstatin-A, 1×protease inhibitor cocktail) and centrifuged at 14,000*g* for 10 minutes at 4 °C.

The nuclei pellet was resuspended in lysis buffer (50 mM Tris-HCl pH 8.0, 1% SDS, 10 mM EDTA, 0.1 mM PMSF, 1 mM pepstatin-A) and chromatin was sonicated using a Covaris E220 evolution with the following settings: power=150V, bursts per cycle=200, duty factor=20%, time=60 seconds. Sonicated chromatin was centrifuged at 14,000g and the supernatant was extracted and diluted with 1×volume of ChIP dilution buffer (1.1% Triton X-100, 20 mM Tris-HCl pH 8.0, 167 mM NaCl, 1.1 mM EDTA, 1mM pepstatin-A, 1×protease inhibitor cocktail). The chromatin was incubated overnight at 4 °C with 50µl Protein A magnetic beads (Dynabeads, Thermo Fisher) pre-bound with either 5µl  $\alpha$ -CENH3 (21), or  $\alpha$ -H3K9me2 antibody (mAbcam 1220). The beads were collected on a magnetic rack and washed twice with low-salt wash buffer (150 mM NaCl, 0.1% SDS, 1% Triton X-100, 20 mM Tris-HCl pH 8.0, 2 mM EDTA, 0.4 mM PMSF, 1 mM pepstatin-A, 1×protease inhibitor cocktail). Immunoprecipitated DNA–protein complexes were eluted from the beads (1% SDS, 0.1 M NaHCO<sub>3</sub>) at 65°C for 15 minutes. Samples were reverse crosslinked by incubating with 0.24 M NaCl at 65°C overnight. Proteins and RNA were digested with Proteinase K treatment, and RNAse A, and DNA was purified with

phenol:chloroform:isoamyl alcohol (25:24:1) extraction and ethanol precipitation. Library preparation followed the Nanopore SQK-LSK109 protocol and kit (as above) and sequenced on separate flongle flowcells.

# **Per-read DNA methylation analysis following CENH3 and H3K9me2 ChIP and ONT sequencing** The resulting .fast5 files were basecalled with Guppy (v5.0.11+2b6dbffa5), using the dna\_r9.4.1\_450bps\_sup.cfg and aligned to the Col-CEN reference with Winnowmap (v1.11, k=15, --MD -ax map-ont). Reads overlapping centromeric positions (Chr1: 14840000-17560000, Chr2: 3823000-6046000, Chr3: 13597000-15734000, Chr4: 4204000-6978000, Chr5: 11784000-1456000) were extracted, providing a set of 5,130 and 11,150 CENH3- or H3K9me2-associated centromeric reads, respectively. The methylation predictions for CG, CHG and CHH methylation contexts were extracted using DeepSignal-plant (v0.1) (*33*) within these read sets. The resulting .tsv files were filtered to remove ambiguous calls (prob\_cf=0.5) and used to calculate the mean methylation state of each context, across individual reads within both data sets. These values were then plotted in R version 4.0.0.

#### ChIP-seq and MNase-seq data alignment and processing

Deduplicated paired-end ChIP-seq and MNase-seq Illumina reads (**Table S7**) were processed with Cutadapt v1.18 to remove adapter sequences and low-quality bases (Phred+33-scaled quality <20) (88). Trimmed reads were aligned to the Col-CEN genome assembly using Bowtie2 v2.3.4.3 with the following settings: --very-sensitive --no-mixed --no-discordant -k 10 (89). Up to 10 valid alignments were reported for each read pair. Read pairs with Bowtie2-assigned MAPQ <10 were discarded using Samtools v1.9 (70). For retained read pairs that aligned to multiple locations, with varying alignment scores, the best alignment was selected. Alignments with more than 2 mismatches or consisting of only one read in a pair were discarded. Single-end SPO11-1-oligo reads were processed and aligned to the Col-CEN assembly using an equivalent pipeline without paired-end options, as described (*41*). For each data set, bins per million mapped reads (BPM; equivalent to transcripts per million, TPM, for RNA-seq data) coverage values were generated in bigWig and bedGraph formats with the bamCoverage tool from

deepTools v3.1.3 (90). Reads that aligned to chloroplast or mitochondrial DNA were excluded from this coverage normalization procedure.

#### **RNA-seq data alignment and processing**

Paired-end RNA-seq Illumina reads (2×100 bp) (**Table S7**) (*42*) were processed with Trimmomatic v0.38 to remove adapter sequences and low-quality bases (Phred+33-scaled quality <3 at the beginning and end of each read, and average quality <15 in 4-base sliding windows) (*41*, *91*). Trimmed reads were aligned to the Col-CEN genome assembly using STAR v2.7.0d with the following settings: -- outFilterMultimapNmax 100 --winAnchorMultimapNmax 100 --outMultimapperOrder Random -- outFilterMismatchNmax 2 --outSAMattributes All --twopassMode Basic --twopass1readsN -1 (*92*). Read pairs with STAR-assigned MAPQ <3 were discarded using Samtools v1.9 (*70*). For retained read pairs that aligned to multiple locations, with varying alignment scores, the best alignment was selected. Alignments with more than 2 mismatches, or consisting of only one read in a pair, were discarded.

#### Small RNA-seq data alignment and processing

Small RNA-seq Illumina reads (**Table S7**) (*42*) were processed with BBDuk from BBMap v38.22 (*93*) to remove ribosomal sequences, and with Cutadapt v1.18 (*88*) to remove adapter sequences and lowquality bases (Phred+33-scaled quality <20). Trimmed reads were aligned to the Col-CEN genome assembly using Bowtie v1.2.2, allowing no mismatches (*73*). For reads that aligned to multiple locations, with varying alignment scores, the best alignment was selected. For each small RNA size class (18–26 nucleotides), TPM values in adjacent genomic windows were calculated based on the total retained alignments (across all size classes) in the library.

#### Bisulfite sequencing data alignment and processing

Paired-end bisulfite sequencing Illumina reads (2×90 bp) (**Table S7**) (42) were processed with Trim Galore v0.6.4 to remove sequencing adapters, low-quality bases (Phred+33-scaled quality <20) and 3 bases from the 5' end of each read (94). Trimmed reads were aligned to the Col-CEN assembly using Bismark v0.20.0 (95). Read pairs that aligned equally well to more than one location and duplicate

alignments were discarded. Methylated cytosine calls in CG, CHG and CHH sequence contexts were extracted and context-specific DNA methylation proportions were generated in bedGraph and bigWig formats using the bismark2bedGraph and UCSC bedGraphToBigWig tools. DNA methylation proportions for cytosines covered by <6 reads were excluded. Single-end bisulfite sequencing reads (50 bp) (**Table S7**) (*34*, *35*) were processed and aligned to the Col-CEN assembly using an equivalent pipeline without paired-end options.

#### Fine-scale profiling around feature sets

Fine-scale profiles around *CEN180* (n=66,131), randomly positioned loci of the same number and width distribution (n=66,131), centromeric intact *ATHILA* elements (n=53), *ATHILA* elements located outside the centromeres (n=58), and *GYPSY* retrotransposons (n=3,979) were calculated for ChIP-seq, RNA-seq, small RNA-seq and bisulfite-seq data sets by providing the above-described bigWig files to the computeMatrix tool from deepTools v3.1.3 in 'scale-regions' mode (*90*). Each feature was divided into non-overlapping, proportionally scaled windows between start and end coordinates, and flanking regions were divided into 10-bp windows. Mean values for each data set were calculated within each window, generating a matrix of profiles in which each row represents a feature with flanking regions and each column a window. Coverage profiles for a ChIP input sequencing library and a gDNA library (**Table S7**) were used in conjunction with those for ChIP-seq and SPO11-1-oligo libraries, respectively, to calculate windowed log<sub>2</sub>([ChIP+1]/[control+1]) coverage ratios for each feature. Meta-profiles (windowed means and 95% confidence intervals) for each group of features were calculated and plotted using the feature profiles in R version 4.0.0.

#### **Crossover mapping**

Total data from 96 Col×Ler genomic DNA  $F_2$  sequencing libraries (2×150 bp) were aligned to the Col-CEN assembly using bowtie2 (default settings). Polymorphisms were identified using the alignment files with samtools mpileup (-vu -f) and bcftools call (-mv -Oz). The resulting polymorphisms were filtered for SNPs (n=522,112), which was used as the 'complete' polymorphism set in TIGER. These SNPs were additionally filtered by, (i) removing SNPs with a quality score less than 200, (ii) removing SNPs where total coverage was greater than 300, or less than 50, (iii) removing SNPs that had reference allele coverage less than 20 or greater than 150, (iv) removing SNPs that had variant allele coverage greater than 130, (v) masking SNPs that overlapped transposon and repeat annotations and (vi) masking SNPs within the main *CEN180* arrays. This resulted in a 'filtered' set of 248,695 SNPs for use in TIGER. DNA sequencing data from 260 wild type Col×Ler  $F_2$  genomic DNA (192 from ArrayExpress E-MTAB-4657 and 68 from E-MTAB-6577) was aligned to the Col-CEN assembly using bowtie2 (default settings) and the alignment analyzed at the previously defined 'complete' SNPs using samtools mpileup (-vu -f) and bcftools call (-m -T). These sites were used as an input to TIGER, which identifies crossover positions by genotype transitions (*96*). A total of 2,080 crossovers were identified with a mean resolution of 1,047 bp.

#### Epitope tagging of V5-DMC1

The DMC1 promoter region was PCR amplified from Col-0 genomic DNA using the Dmc1-PstI-fw and Dmc1-SphI-rev oligonucleotides. The remainder of the DMC1 promoter, gene and terminator were amplified with oligonucleotides Dmc1-SphI-fw and Dmc1-NotI-rev. The resulting PCR fragments were digested with PstI and SphI, or SphI and NotI, respectively, and cloned into PstI-NotI-digested pGreen0029 vector to yield a pGreen-DMC1 construct. To insert 3 N-terminal V5 epitope tags, first two fragments were amplified with DMC1-Nco-F and 3N-V5-R and 3N-V5-F and Dmc1-Spe-rev and then used in an overlap PCR reaction using the DMC1-Nco-F and Dmc1-Spe-rev oligonucleotides. The PCR product resulting from the overlap PCR was digested with NcoI and SpeI and cloned into NcoIand SpeI-digested pGreen-DMC1. The resulting binary vector was used to transform dmc1-3/+ heterozygotes (SAIL\_126\_F07). We used dmc1-seq11 and Dmc1-Spe-rev oligonucleotides to amplify wild type DMC1 allele and Dmc1-Spe-rev and LA27 to amplify the dmc1-3 T-DNA mutant allele. The presence of the V5-DMC1 transgene was detected with N-screen-F and N-screen-R oligonucleotides. This oligonucleotide pair amplifies a 74 bp product in Col and a 203 bp product in V5-DMC1. To identify dmc1-3 homozygotes in the presence of V5-DMC1 transgenes, we used DMC1-genot-compl-F and DMC1-genot-compl-R oligonucleotides, which allowed us to distinguish between the wild type DMC1 gene and V5-DMC1 transgene. All oligonucleotide sequences are provided in Table S8.

#### Cytogenetic and immunocytological analyses

For fluorescence in situ hybridization (FISH), spreads of meiotic chromosomes at pachytene stage of meiosis were prepared from young flower buds fixed in ethanol:acetic acid (3:1) and stored in 70% ethanol until use. Chromosome spreads were prepared as described (97). To identify individual chromosome arms, chromosome-specific A. thaliana BAC clones were arranged into contigs. More specifically, the following BAC contigs were used: five (F10C21/AC051630 - F12K21/AC023279; Fig. 1D, 1F, S5A and S5D), 15 (F13M18/AL087094 – F12K21/AC023279; Fig. S5C and S5E) or 29 (F6F9/AC007797 – F12K21/AC023279; Fig. S5B) chromosome 1 upper-arm-specific BACs; five (F2J6/AC009526 – T2P3/B21868; Fig. 1D and S5A) or 36 (F2J6/AC009526 – T6H22/AC009894; Fig. S5B) chromosome 1 bottom-arm-specific BACs; five (T21B4/AF007271 – T8M17/AF296835; Fig. S5A) or 29 (T2007/AB026660 - T8M17/AF296835; Fig. 3H and S5E) chromosome 5 upper-armspecific BACs; five (F5M8/AL082902 - T31G3/AB026662; Fig. S5A) chromosome 5 bottom-armspecific BACs. The Arabidopsis (TTTAGGG)<sub>n</sub> telomere repeat probe was prepared by PCR, as described (98). All DNA probes were labeled with biotin-dUTP, digoxigenin-dUTP, or Cy3-dUTP by nick translation, then pooled, ethanol-precipitated and pippeted on pepsin-treated and ethanoldehydrated slides containing suitable chromosome spreads. The slides were heated to 80°C for 2 minutes and incubated at 37°C for 12 hours. The hapten-labeled probes were immuno-detected as described (97). BAC contigs and other DNA probes were visualised using fluorescently labeled antibodies against biotin-dUTP (avidin-Texas red, Vector Laboratories, cat. no. A-2006-5) and digoxigenin-dUTP (mouse anti-digoxigenin, Jackson ImmunoResearch, 200-002-156, goat anti-mouse Alexa Fluor 488, Invitrogen, A11001, and goat anti-mouse Alexa Fluor 647, Invitrogen, A21235). Chromosomes were counterstained with DAPI (2 µg/mL) in Vectashield (Vector Laboratories). Fluorescence signals were analyzed and imaged using a Zeiss AxioImager epifluorescence microscope (Carl Zeiss) with a CoolCube camera (MetaSystems). Images were acquired separately using the Isis software (MetaSystems) for all four fluorochromes using appropriate excitation and emission filters (AHF Analysentechnik). The four monochromatic images were pseudocoloured, merged, and cropped using Photoshop CS (Adobe Systems), and chromosome length was measured using ImageJ (National Institutes of Health).

The *CEN180* pAL FISH probe, which labels all centromeres, was amplified using primers ATH\_cen180F and ATH\_cen180R (**Table S8**) (*99*). PCR amplification was performed as follows: initial denaturation at 95°C for 5 minutes; 35 cycles of denaturation at 95°C for 20 seconds, annealing at 46°C for 20 seconds and extension at 72°C for 20 seconds; and a final extension at 72°C for 5 minutes. PCR products were purified using the NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel) and labelled by nick translation. To design *CEN180* probes specific to individual chromosomes or sets of chromosomes, *CEN180* sequences identified in the Col-CEN assembly were aligned using MAFFT (v7.450) and used to identify repeats with high copy number and distributions biased to specific chromosomes. Oligonucleotide FISH probes homologous to specific *CEN180* sequences were designed that were 60 nucleotides in length, with a GC content between 30-50% and selected to minimize self-annealing and formation of hairpin structures, using Geneious (v11.1.5) (**Table S8**). Double-stranded DNA probes were prepared and labelled, as described (*97*).

To design FISH probes against *ATHILA* transposons the sequences encoding the highly variable GAG domains for each sub-family were aligned using MAFFT (v7.450) and consensus sequences were generated. PCR primers were then designed to amplify subfamily GAG domain genes, using Primer3 (v2.3.7) implemented in Geneious (**Table S8**). PCR amplification was performed as follows: initial denaturation at 95°C for 5 minutes; 35 cycles of denaturation at 95°C for 20 seconds, annealing at 58°C for 20 seconds and extension at 72°C for 20 seconds; and a final extension at 72°C for 5 minutes. PCR products were purified using the NucleoSpin Gel and PCR Clean-up kit and subsequently cloned into the pGEM-T Easy Vector System (Promega), using TOP10 competent cells. Positive colonies were screened using SP6/T7 primers and five clones of each *ATHILA-GAG* gene were Sanger sequenced. Subsequently, clones with the highest pairwise sequence similarity to specific *ATHILA* sub-family consensus sequences were used as templates for PCR amplification. Purified PCR products were labelled by nick translation, as described (*97*).

For analysis of chromatin during mitotic interphase, nuclei were isolated from 1 week old seedlings (wild type Col-0 and CENH3-GFP (*100*)) and treated as described (*37*). Primary antibodies were diluted 1:200 while the secondary antibodies Alexa488 and Alexa555 goat anti rabbit or goat anti mouse conjugates (Molecular Probes) were diluted 1:500. The primary antibodies used were anti-GFP (mouse, Roche 11814460001), anti-H3K4me1 (rabbit, Abcam Ab8895), anti-H3K4me3 (rabbit, Abcam Ab8896), anti-H3K4me3 (rabbit, Abcam Ab8896), anti-H3K9me2 (mouse, Abcam Ab1220), anti-H3K27me1 (rabbit, Abcam Ab124688), anti-H3K27me3 (rabbit, Sigma Aldrich 07-449) and anti-K36me3 (rabbit, Abcam Ab9050). To visualize DNA, nuclei were mounted in Vectashield containing DAPI. Images were acquired with the LSM980 Axio Observer with the Airyscan2 detector from Zeiss. Images were Airscan processed using the Zen Black software. Images were further analyzed using Fiji software. To correct for 3D shifts between channels in the Z plane, differences between the channels were estimated by imaging fluorescent beads. The channels were then aligned to correct for this shift. Areas of interest were resliced in Image J to obtain line plots. Intensity plots were then made using the ggplot2 package in R 3.5.1.

To immunocytologically analyse meiosis, fresh buds at floral stage 8 and 9 were dissected to release the anthers that contain male meiocytes (*101*). Chromosome spreads of meiotic and mitotic cells from anthers were performed, followed by immunofluorescent staining of proteins as described (*39*). The antibodies used in this study were:  $\alpha$ -ZYP1 (rabbit, 1/500 dilution) (*102*),  $\alpha$ -H3K9me2 (mouse, 1/200 dilution) (Abcam, ab1220),  $\alpha$ -CENH3 (rabbit, 1/100 dilution) (Abcam, ab72001) and  $\alpha$ -V5 (chicken, 1/200 dilution) (Abcam, ab9113). Chromosomes stained with ZYP1, CENH3 and H3K9me2 were visualized with a DeltaVision Personal DV microscope (Applied Precision/GE Healthcare). Chromosomes stained with DMC1-V5 and CENH3 were visualized with a Leica SP8 confocal microscope. Chromosomes stained with H3K9me2 were visualized with a Stimulated emission depletion nanoscopy mounted on an inverted IX71 Olympus microscope.

**Table S1. Consensus quality (QV) score of the Col-CEN Arabidopsis genome assembly.** Consensus quality scores (QV) were calculated from "missing" 21-mers (k\_asm) present in the Col-CEN assembly, but not present in the short read Illumina library. k\_total shows the total number of 21-mers. QV scores were calculated for Col-CEN individual chromosomes (green), centromeres (blue), chromosome arms (orange), or the whole genome (yellow).

feature_label	chrom	start	end	k_asm	region_len	k_total	error_rate	QV
Chr1	Chr1	0	32540122	491	32540122	32540102	0.00000071 85324858	61.44
Chr2	Chr2	0	22217084	12938	22217084	22217064	0.00002773 841542	45.57
Chr3	Chr3	0	25743512	956	25743512	25743492	0.00000176 8393137	57.52
Chr4	Chr4	0	21578073	4606	21578073	21578053	0.00001016 568227	49.93
Chr5	Chr5	0	29480885	4525	29480885	29480865	0.0000730 9552866	51.36
CEN1	Chr1	14841109	17559778	37	2718669	2718649	0.00000064 80851998	61.88
CEN2	Chr2	3823791	6045243	26	2221452	2221432	0.00000055 73441539	62.54
CEN3	Chr3	13597187	15733925	368	2136738	2136718	0.00000820 1946565	50.86
CEN4	Chr4	4203901	6977949	4308	2774048	2774028	0.00007400 600007	41.31
CEN5	Chr5	11784130	14551809	1934	2767679	2767659	0.00003328 6578	44.78
Chr1-CEN1	NA	NA	NA	454	29821453	29821433	0.00000072 49552477	61.40

Chr2-CEN2	NA	NA	NA	12912	19995632	19995612	0.00003075 906296	45.12
Chr3-CEN3	NA	NA	NA	588	23606774	23606754	0.00000118 61153	59.26
Chr4-CEN4	NA	NA	NA	298	18804025	18804005	0.00000075 46574935	61.22
Chr5-CEN5	NA	NA	NA	2591	26713206	26713186	0.00000461 8941798	53.35
Whole Genome	NA	NA	NA	23516	NA	131559576	0.00000851 2530102	50.70
All CEN	NA	NA	NA	6673	NA	12618486	0.00002518 859647	45.99
Whole Genome - All CEN	NA	NA	NA	16843	NA	118941090	0.00000674 3688897	51.71

### Table S2. TAIR10 gene models that show presence-absence variation (PAV) in Col-CEN. TAIR10

gene models were mapped onto Col-CEN using Liftoff (80). Genes that occurred as presence-absence variants (PAVs), as they did not map to Col-CEN, are listed and classified as loci in the CLUSTER\_ID column.

ID	CHROM	START (1-based)	END	LEN	PLUS_STRAND	NOTE	CLUSTER_ID
AT1G34800	Chr1	12773164	12773442	279	0	protein_coding_gene	PCG_0 (THIONIN)
AT1G34805	Chr1	12776578	12776856	279	0	protein_coding_gene	PCG_0 (THIONIN)
AT1G34830	Chr1	12793536	12794023	488	0	protein_coding_gene	PCG_0 (THIONIN)
AT1G34840	Chr1	12796984	12797247	264	0	protein_coding_gene	PCG_0 (THIONIN)
AT1G38065	Chr1	14289578	14292060	2483	0	protein_coding_gene	PCG_1
AT1G56820	Chr1	21273314	21273395	82	1	tRNA	TRNA_6
AT1G56910	Chr1	21277861	21277942	82	1	tRNA	TRNA_6
AT1G57030	Chr1	21283986	21284067	82	1	tRNA	TRNA_6
AT1G57210	Chr1	21292992	21293073	82	1	tRNA	TRNA_6
AT1G57240	Chr1	21294341	21294422	82	1	tRNA	TRNA_6
AT1G57300	Chr1	21297221	21297302	82	1	tRNA	TRNA_6

AT1G57330	Chr1	21298753	21298834	82	1	tRNA	TRNA_6
AT1G58808	Chr1	21784645	21786869	2225	1	other_RNA	PCG_2
AT1G58848	Chr1	21791783	21797050	5268	1	protein_coding_gene	PCG_2
AT1G58983	Chr1	21806020	21807487	1468	0	protein_coding_gene	PCG_2
AT1G59030	Chr1	21808193	21809509	1317	0	protein_coding_gene	PCG_2
AT1G59077	Chr1	21810644	21813023	2380	0	protein_coding_gene	PCG_2
AT1G59124	Chr1	21816443	21820572	4130	1	protein_coding_gene	PCG_2
AT1G59312	Chr1	21839858	21841972	2115	1	protein_coding_gene	PCG_2
AT5G36670	Chr5	14401491	14406427	4937	1	protein_coding_gene	PCG_3
AT5G36680	Chr5	14406802	14409137	2336	0	protein_coding_gene	PCG_3
AT5G36690	Chr5	14415185	14417288	2104	0	protein_coding_gene	PCG_3
AT5G36700	Chr5	14421576	14424511	2936	0	protein_coding_gene	PCG_3
AT5G36720	Chr5	14429661	14429924	264	0	protein_coding_gene	PCG_3
AT5G36722	Chr5	14431599	14432216	618	0	protein_coding_gene	PCG_3
AT5G36800	Chr5	14484565	14485409	845	0	protein_coding_gene	PCG_3
AT5G36820	Chr5	14495617	14496849	1233	1	protein_coding_gene	PCG_3

**Table S3. TAIR10 gene models that mapped as additional copies to Col-CEN.** TAIR10 gene models are listed that mapped via Liftoff to more than one location in Col-CEN (80). The CLUSTER\_ID column indicates close linkage of the duplicated genes.

TAIR10 ID	TAIR10 CHR	TAIR10 START	TAIR10 LEN	TAIR10 NOTE	COLCEN ID	ColCEN CHR	ColCEN START	Col-CEN LEN	CLUSTER_ID
AT2G1614 5	Chr2	7008520	80	miRNA	AT2G16145	Chr2	9512451	80	miRNA_1
AT2G1614 5	Chr2	7008520	80	miRNA	AT2G16145_1	Chr2	9517494	80	miRNA_1
AT1G2482 2	Chr1	8774997	2886	protein_coding	AT1G24822	Chr1	8780249	2886	PCG_5
AT1G2482 2	Chr1	8774997	2886	protein_coding	AT1G24822_1	Chr1	8848453	2884	PCG_5
AT1G2490 9	Chr1	8785785	2130	protein_coding	AT1G24909	Chr1	8830487	2130	PCG_5
AT1G2490 9	Chr1	8785785	2130	protein_coding	AT1G24909_1	Chr1	8844863	2130	PCG_5
AT1G2514 1	Chr1	8817678	705	protein_coding	AT1G25141_2	Chr1	8802028	705	PCG_5
AT1G2514 1	Chr1	8817678	705	protein_coding	AT1G25141_1	Chr1	8839849	705	PCG_5

AT1G2514 1	Chr1	8817678	705	protein_coding	AT1G25141	Chr1	8854226	705	PCG_5
AT1G2521 0	Chr1	8833018	2095	protein_coding	AT1G25210_1	Chr1	8772607	2094	PCG_5
AT1G2521 0	Chr1	8833018	2095	protein_coding	AT1G25210	Chr1	8840811	2093	PCG_5
AT1G5993 0	Chr1	22061083	399	protein_coding	AT1G59930	Chr1	24161766	399	PCG_6
AT1G5993 0	Chr1	22061083	399	protein_coding	AT1G59930_1	Chr1	24163425	399	PCG_6
AT1G7793 2	Chr1	29302725	795	protein_coding	AT1G77932	Chr1	31405241	795	PCG_7
AT1G7793 2	Chr1	29302725	795	protein_coding	AT1G77932_1	Chr1	31411502	795	PCG_7
AT1G7794 0	Chr1	29303897	1486	protein_coding	AT1G77940	Chr1	31406413	1486	PCG_7
AT1G7794 0	Chr1	29303897	1486	protein_coding	AT1G77940_1	Chr1	31412674	1486	PCG_7
AT5G3915 0	Chr5	15669898	911	protein_coding	AT5G39150	Chr5	18143268	911	PCG_8
AT5G3915 0	Chr5	15669898	911	protein_coding	AT5G39150_1	Chr5	18156831	911	PCG_8
AT5G3917 0	Chr5	15680731	595	protein_coding	AT5G39170_1	Chr5	18154101	595	PCG_8
AT5G3917 0	Chr5	15680731	595	protein_coding	AT5G39170	Chr5	18167662	595	PCG_8
AT5G3919 0	Chr5	15692591	991	protein_coding	AT5G39190_1	Chr5	18152385	991	PCG_8
AT5G3919 0	Chr5	15692591	991	protein_coding	AT5G39190	Chr5	18179541	991	PCG_8
AT5G4091 0	Chr5	16395507	3623	protein_coding	AT5G40910	Chr5	18882480	3623	PCG_9
AT5G4091 0	Chr5	16395507	3623	protein_coding	AT5G40910_1	Chr5	18887652	3623	PCG_9
ATCG0091 0	ChrC	100709	72	tRNA	ATCG00910_1	Chr4	8541426	72	tRNA_1
ATCG0091 0	ChrC	100709	72	tRNA	ATCG00910	ChrC	100709	72	tRNA_1

**Table S4. Unique and repeated** CEN180 monomer sequences within and between chromosomes.CEN180 monomers were compared across the genome to identify unique versus repeated sequences.For repeated sequences we show which chromosomes they occurred on.

Chr	Total	Unique	Repeated	Chr1	Chr2	Chr3	Chr4	Chr5	Chr2,
									Chr4, Chr5
Chr1	13,578	4,174	Chr1	9,372	0	265	0	2	25
Chr2	12,293	3,887	Chr2		8,363	20	20	7	
Chr3	11,848	3,944	Chr3			7,662	0	7	
Chr4	15,613	4,951	Chr4				10,660	0	
Chr5	12,799	5,484	Chr5					7,287	

All	66,131	22,440	Total	9,372	8,363	7,947	10,680	7,303	43,691
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**Table S5.** *CEN180* **higher order repeats.** *CEN180* monomers were classified as being the same if they shared 5 or fewer pairwise variants, and consecutive blocks identified as higher order repeats (HORs). HORs are all in a tandem orientation and are classified as being intra- or inter-chromosome. The mean HOR block size, in monomers and bp, and the mean distance between intra-chromosome HORs (bp) are listed.

Chr	Monomers	Intra- chromosome HORs	Inter- chromosome HORs	Mean HOR monomers	Mean HOR block (bp)	Mean HOR distance (bp)
1	13,578	814,715	24,110	2.41	429	365,291
2	12,293	584,684	13,757	2.35	418	434,776
3	11,848	413,642	2,743	2.50	446	334,277
4	15,613	498,876	611	2.40	427	402,170

5	12,799	55,515	0	2.86	509	167,045
All	66,131	2,367,432	41,221	2.41	429	365,291

**Table S6. Structural and sequence characteristics of centromeric** *ATHILA* **retrotransposons.** Analysis of 111 gaps greater than 1 kbp in the main *CEN180* arrays identified 53 intact and 20 fragmented *ATHILA* retrotransposons, as well as 12 solo LTRs. For each sequence we report the *ATHILA* subfamily class based on the TAIR10 classification and our phylogenetic analysis, and information on element length, strand, target site duplications (TSDs), long terminal repeat (LTR) position and length, and hits with Hidden Markov Models (HMMs) that describe *GYPSY* LTR retrotransposon open reading frames (see Methods). The 'quality' column indicates whether the *ATHILA* is an 'intact' full-length element, i.e. it contains clearly identified LTRs and, possibly, a TSD; a fragment - note that we also included as fragments and not as intact elements, i) *ATHILA* copies with large internal deletions (e.g. the 4872 bp *ATHILA*2 element in centromere 4 has complete and highly similar LTRs but also a ~6 kbp internal deletion), and ii) *ATHILA* copies with a deletion that included the whole LTR plus additional sequence in the internal domain; or a solo LTR. The 'comment' column' includes notes on interesting characteristics for some elements. For example, it highlights the *ATHILA5* duplicates in centromere 5 that contain the internal *CEN180* repeats, and some cases where two intact *ATHILA* of the same subfamily share one LTR (LTR-internal.region-LTR-internal.region-LTR), possibly as a result of post-integration interelement homologous recombination. Given that the LTRs of the *ATHILA6A* and *ATHILA6B* subfamilies appear identical, it was not possible to further allocate solo LTRs of the *ATHILA6* clade into their respective subfamilies. In addition to the *ATHILA* elements, a small number of other TEs were identified but not further analyzed due to their fragmented organization. The majority of these elements occur in centromere 1 and are shown at the end of the Table. Note that for these elements the coordinates refer to the position of the gaps and not the TEs within the gaps. Due to size, Table S6 is attached as a separate file 'Table S6.xlsx'.

Table S7. Summary of short-read Illumina sequencing libraries aligned to the Col-CEN assembly. All data sets were generated from plants in a Col-0 background, with the exception of the Col×Ler  $F_2$  genomic DNA sequencing libraries that were used to identify meiotic crossovers.

Library	Study accession	Run accession	Read length	Tissue	References
CENH3 ChIP-seq	PRJNA349052	SRR4430537	2×100 bp	Seedling	(19)
H3K9me2 ChIP-seq	PRJEB36221	ERR3813867	2×75 bp	Floral bud	(39)
H3K27me1 ChIP-seq	PRJEB36221	ERR3813864	2×75 bp	Floral bud	(39)

H3K4me1	PRJEB36221	ERR3813865	2×75 bp	Floral bud	(39)
ChIP-seq					
H3K4me2	PRJEB36221	ERR3813866	2×75 bp	Floral bud	(39)
ChIP-seq					
H3K4me3	PRJEB15183	ERR1590146	2×150 bp	Floral bud	(41)
ChIP-seq					
1					
H3K27me3	PRJNA252965	SRR1509478	2×100 bp	Floral bud	(103)
ChIP-seq					
enn seg					
H2A.W6	N/A	N/A	50 bp	Seedling	This study
ChIP-seq					
Chin -seq					
H2A.W7	N/A	N/A	50 bp	Seedling	This study
ChID sog			Ĩ	C	·
ChIP-seq					
H2A.Z ChIP-	PRJNA219442	SRR988546	50 bp	Leaf	(37)
			1		~ /
seq					
REC8	PRJEB36221	ERR3813871	2×75 bp	Floral bud	(39)
			F		(57)
ChIP-seq					
ASY1	PRJEB36320	ERR3829803	2×75 bp	Floral bud	(40)
	1132250520	Litti 5027003	2015 op	i ioiui oud	(70)
ChIP-seq					
SPO11-1-oligos	PRJEB15185	ERR1590157	50 bp	Floral bud	(A1)
51 011-1-011g08	I NJED I J 10J	EIXIX1370137	50 op		(41)
MNase-seq	PRJEB15184	ERR1590154	2×100 bp	Floral bud	(41)
gDNA	PRJEB23842	ERR2215865	2×100 bp	Floral bud	(41)
			- L		()

RNA-seq (Col-	PRJEB9919	ERR966157-	2×100 bp	Leaf	(42)
0 and <i>met1-3</i> )		ERR966162			
Bisulfite-seq	PRJEB9919	ERR965674-	2×90 bp	Leaf	(42)
(Col-0 and		ERR965677			
met1-3)					
Bisulfite-seq	PRJNA172021,	SRR534177–	50 bp	Leaf	(34, 35)
(Col-0 and	PRJNA222364	SRR869314,			
mutants)		SRR1005412-			
		SRR1005415			
Small RNA-seq	PRJEB9919	ERR966148-	50 bp	Leaf	(42)
(Col-0 and		ERR966149			
<i>met1-3</i> )					
Col-0×Ler-0	E-MTAB-4657	E-MTAB-4657	2×150 bp	Leaf	(104, 105)
genomic DNA	E-MTAB-6577	E-MTAB-6577			
F <sub>2</sub>					
Hi-C (Col-0)	PRJNA253621	SRR1504819	2×50 bp	Leaf	(106)

 Table S8. Oligonucleotides. The sequence of oligonucleotides used for V5-DMC1 construction and genotyping, and FISH, are listed.

Oligo name	Sequence 5' to 3'	Purpose
Dmc1-PstI-fw	ATATATACTGCAGGATATCAAACATTTACC TGAAAAGA	Cloning 3V5-DMC1
	ΙΟΑΑΑΟΑ	
Dmc1-SphI-rev	ATATATGCATGCTTCTTTTAACTCTTCTCAT	Cloning 3V5-DMC1
Dmc1-SphI-fw	AAAGAAGCATGCTTAAGCCAACAGAG	Cloning 3V5-DMC1
Dmc1-NotI-rev	ATATATATATATGCGGCCGCGAGTTTTGCA	Cloning 3V5-DMC1
	GCAATTATGAAA	

Dmc1-Spe-rev	TATCAAACTAGTGTAAAGTAAACCTTGGTT	Cloning <i>3V5-DMC1</i> , genotyping <i>dmc1-3</i>
DMC1-Nco-F	TTTCTTTCCATGGATTAAAAAAATTTG	Cloning 3V5-DMC1
3N-V5-F	GGTAAACCAATCCCAAACCCACTCCTCGGT	Cloning 3V5-DMC1
	CTCGACTCAACAGGAAAGCCTATTCCTAAT	
	CCTCTTCTTGGACTTGATTCTACTATGATG	
	GCTTCTCTTAAGTAAGTGA	
3N-V5-R	GGGTTTGGGATTGGTTTACCAGTAGAATCA	Cloning 3V5-DMC1
	AGTCCAAGAAGAGGATTAGGAATAGGCTT	
	TCCCATTTTCTCGCTCTAAGAGTCTCTA	
Dmc1-screen-N-fw	CTCTCACTCTTCCAAGCTTA	Genotyping 3V5-DMC1
Dmc1-screen-N-rev	AGAGATCAATCACTTACTTAAGAG	Genotyping 3V5-DMC1
LA27	TAGCATCTGAATTTCATAACCAATCTCGAT	Genotyping <i>dmc1-3</i>
	ACAC	
DMC1-genot-compl-F	CATACATTGACACAGAGGGAACC	Genotyping <i>dmc1-3</i> in the
2.1.01 genet comp. 1		presence of <i>3V5-DMC1</i>
DMC1-genot-compl-R	ATGGAACCCAAAAGAGGAGAC	Genotyping <i>dmc1-3</i> in the
		presence of 3V5-DMC1
ATH_cecen180F	CATATTCGACTCCAAAACACTAACC	Amplification of pAL
		universal CEN180 probe

ATH_cen180R	AGAAGATACAAAGCCAAAGACTCAT	Amplification of pAL universal <i>CEN180</i> probe
CEN180-α	CCGCAACAGGATCTTAAAGGCGTAAGAAT TTTATTCTGTTAAAAGACACAAAGCCAAA GA	CEN180 FISH probe
CEN180-β	ATTGAATCTTTGTTAGAAGATACAAAGAC AAAGACTCATACGGACTTCGACTACACTAT C	CEN180 FISH probe
CEN180-γ	TTAAACTGCAATTGGATCTTAAAGGCGTAA GAATTGTATCCTTGTTAAAAGACACAAAG C	CEN180 FISH probe
CEN180-δ	CGCATCTTATAAGCCTAAGTAGTATTTCCT TGTTAGAATACACAAAGTCAAAGACTCAT A	CEN180 FISH probe
CEN180-ε	TCTTATAAGCCTAAGTAGTGTTTCCTTGTT AGAAGACACAAAAGCCAATGACTCATATCG C	CEN180 FISH probe
ATHILA2_GAG_F	GGATCCACTCGACCACCTTG	Amplification of the <i>ATHILA2</i> FISH probe
ATHILA2_GAG_R	AACCCTTGAAACGCTCCCAT	Amplification of the <i>ATHILA2</i> FISH probe
ATHILA6A6B_GAG_ F	GATCCACTCGATCACCTGGAC	Amplification of the <i>ATHILA6A/6B</i> FISH probe

ATHILA6A6B_GAG_	TCCCATGCTTCGCAGAAAGT	Amplification of the
R		ATHILA6A/6B FISH probe



**Figure S1. Validation of the Col-CEN centromere assembly. A.** Assembly consensus quality (QV) scores of the individual and collective (All) centromeres. **B.** IGV screenshots depicting quantitative tracks across the five centromeres. All coverage tracks are binned via averaging, whereas the marker and missing k-mer tracks are aggregated in 10 kbp windows with no IGV binning. Secondary Allele Coverage tracks depict the coverage of the most covered alternate sequence (if any) indicated by the

alignments at every position. The "marker" and "missing" k-mer tracks are plotted with a y-axis log scale. **C**. Distribution of distances (bp) between consecutive marker 21-mers.


**Figure S2. Telomere assembly and validation of the Col-CEN assembly using Bionano and Hi-C data. A.** IGV screenshot showing the start of Col-CEN chromosome 3, including the assembled telomere. Gene models and mapped TAIR10 BACs are indicated, in addition to matches to the telomeric repeat (orange, CCCTAAA). Also shown in blue are ONT and HiFi read mappings to the Col-CEN assembly. **B.** A Hi-C heatmap generated by aligning Col-0 Hi-C reads to the Col-CEN assembly (*106*). **C.** Bionano *de novo* assembly contigs were mapped to the Col-CEN reference assembly. The green and blue bars represent the expected labeling positions in the ONT reference assembly, where blue bars are expected labeling positions, green regions lack Bionano labels and light brown bars represent predicted labeling positions not linked to a Bionano optical contig. Centromere regions generally lack predicted labeling sequences and therefore Bionano *de novo* assembled contigs are broken.



**Figure S3. Dotplot sequence similarity comparison of TAIR10 and the Col-CEN genome assembly.** A dotplot depicting unique (blue) and repetitive (red) Nucmer alignments (--maxmatch -1 50 -c 250) between TAIR10 and Col-CEN.



**Figure S4.** Genic copy number variation loci between the TAIR10 and Col-CEN assemblies. A. On the left is an IGV screenshot showing a region of chromosome 1 from the Col-CEN assembly that contains a thionin gene cluster that shows a deletion relative to TAIR10 with 4 genes that did not map to Col-CEN (Cluster PCG\_0, see **Table S2**). The screenshot shows alignment of PacBio Hifi reads (upper track). Below, 100 kbp exact WGS reads were simulated from TAIR10 and their alignments are shown (middle track). Finally, TAIR10 BAC contig alignments are shown (lower). Purple marks indicate insertions and additional colors in the coverage tracks indicate substitutions. Uneven TAIR10 simulated read and BAC contig coverage indicates a structural difference between TAIR10 and Col-CEN at this locus, yet uniform HiFi coverage supports Col-CEN assembly accuracy, suggesting that this discrepancy is due to genuine biological variation, rather than misassembly. To the right a dotplot of the PCG\_0 cluster in Col-CEN versus TAIR10 is shown. **B.** As for A., but showing Cluster PCG\_3 on chromosome 5, where 8 TAIR10 genes did not map to Col-CEN (see **Table 2**). **C.** As for A., but showing Cluster PCG\_8 on chromosome 5, where 3 TAIR10 genes mapped with an extra copy to Col-CEN (see **Table 3**).



**Figure S5. Fluorescent** *in situ* hybridization (FISH) analysis of the Arabidopsis centromeres. A. Pachytene-stage meiotic chromosomes were spread and stained with DAPI (white), and FISH performed using probes designed to label all *CEN180* (blue, pAL), pericentromeric *ATHILA* (purple, BAC T1J24), the telomeric repeat (green, *TEL* (TTTAGGG)<sub>n</sub>), chromosome 1 specific BACs (yellow and green) and chromosome 5 specific BACs (red and yellow). The scale bar represents 10 μM. **B.** As

for A., apart from the *CEN180-a* probe (blue) was used for FISH, together with chromosome 1 specific BACs labelled in red and yellow. A blow-up of centromere 1 is shown beneath. **C.** As for A., but labelling with the *CEN180-a* (red), *CEN180-y* (green) and *CEN180-b* (green) FISH probes, together with chromosome 1 specific BACs (yellow). Blow-ups of the centromere 1 region are shown inset. **D.** A cell dividing at metaphase I of meiosis is shown that was stained by DAPI (white), and the *CEN180-* $\varepsilon$  FISH probe (green). **E.** As for A, but labelling with an *ATHILA2* subfamily specific *GAG* probe (green) and chromosome 1 (yellow) and 5 (red) specific BACs.



Figure S6. Comparison of the Col-CEN assembly with physical maps derived from pulsed-field gel electrophoresis and Southern blotting. On the right hand side of the figure published pulsed-field gel electrophoresis and Southern blotting data are shown, where genomic DNA was digested using either AscI or NotI (25–27). The probe used for hybridization is labelled underneath the blots. To the left are physical maps of the Col-CEN assembly that have been virtually digested for AscI (green) or NotI (purple) and site locations indicated relative to chromosome coordinates. The position of plus strand (red) and minus strand (blue) CEN180 are indicated on the x axis. Above each physical map the location of the probes used for Southern blot hybridization are indicated. We further annotate the predicted size of cross-hybridizing fragments following restriction digestion, for comparison with the reproduced data. We note that for *CEN1* the authors interpret probe hybridization as indicating binding to two separate ~4.7 Mbp arrays. However, an incorrect BAC sequence used when designing the (specifically, BAC F8L2 restriction maps sequence: https://www.ncbi.nlm.nih.gov/nuccore/AC087569) predicted an incorrect NotI site, which was inside of the AscI cutting site. However, based on analysis of our assembly the NotI site is in fact outside of the AscI site and thus the probes are binding to the same fragment (25). This region has now also been resolved correctly in the TAIR10 reference assembly.

## Chromosome 1 coordinates (Mbp)

Col-CEN v1.1

contig mapping

Col-CEN v1.1

contig mapping

mappings

Bionano de novo assembly

Raw Bionano molecule

mappings

Bionano de novo assembly

Raw Bionano molecule

## Chromosome 2 coordinates (Mbp)

700 350 70 70 50 50 50 50 50 50 50 50 50 50 50 50 50	an tan tan in in in tan	1999 - 1999 -



**Figure S7. Bionano optical mapping across the Col-0 centromeres.** Bionano *de novo* assembly contigs mapped to the Col-CEN reference assembly. The green and blue bars represent the expected labeling positions in the ONT reference assembly, where blue bars are expected labeling positions,

green regions lack Bionano labels and light brown bars represent predicted labeling positions not linked to a Bionano optical contig. Centromere regions generally lack predicted labeling sequences and therefore Bionano *de novo* assembled contigs are broken. Below the Bionano contigs (blue background with blue bars) are raw molecule mappings to the Bionano contigs at ~1,000× coverage (yellow background with blue dots indicating labelled sites).



**Figure S8. CENH3,** *CEN180* and sequence identity across the Arabidopsis centromeres. CENH3 log2(ChIP/input) (black) (19), plotted over each centromere. *CEN180* density per 10 kbp is plotted showing forward (red) or reverse (blue) strand orientation. The location of *ATHILA* retrotransposons is indicated by purple ticks on the x axis. Beneath the plot are heatmaps indicating pairwise % identity values of all non-overlapping 5 kbp regions.



**Figure S9. The Arabidopsis** *CEN180* **satellite repeat library analysed by chromosome. A.** Histograms of *CEN180* monomer lengths (bp), and variants relative to the genome-wide consensus, shown for each chromosome. Mean values are shown by the red dotted line. **B.** *CEN180* sequence conservation represented by sequence logo plots. The global genome-wide sequence logo is shown first, followed by each individual chromosome. Positions with less than 50% coverage are not shown.



**Figure S10. Dotplot comparison of** *ATHILA* **retrotransposons located inside or outside the main centromeric** *CEN180* **arrays.** Dotplot of centromeric *ATHILA* retrotransposons using a search window of 75 bp. Red and blue indicate forward and reverse strand similarity. The elements assigned to different *ATHILA* subfamilies are indicated, in addition to whether they are located inside or outside the main centromeric *CEN180* arrays.



Figure S11. Higher order duplication of ATHILA elements post-integration. A. Dotplot analysis of a large region that has duplicated within the centromere of chromosome 5, forming higher order repeats (HOR1 and HOR2). The boundaries of each HOR are indicated by the black boxes within the dotplot. Each higher order repeat contains one ATHILA5 and one ATHILA6A element that show high identity (99.5 and 99.6%) between copies. In contrast, the surrounding blocks of CEN180 repeats within each HOR are more variable in size and show lower sequence identity (94.3-97.3%). Additional evidence that this region was duplicated after the insertion of the ATHILA5 and ATHILA6A copies includes, i) their nearly identical lengths (11,345 vs. 11,346 bp for ATHILA6A, and 10,968 vs. 10,961 bp for ATHILA5, ii) the identical target site duplication (TSD) for the ATHILA5 copies (GTAGT), iii) the identical flanking sequences (CCTAAGTAGT for the upstream and GTAGTGTTTC for the downstream region of ATHILA5, and AGACACAAAG for the downstream region of ATHILA6A), and iv) the fact that both ATHILA5 contain internal CEN180 copies in identical positions within their 5'-LTRs (see B). B. Dotplot analysis of one of the duplicated ATHILA5 elements from A, which contains one complete and one partial copy of CEN180, located internally and downstream of the 5'-LTR. We postulate that the CEN180 repeats inserted within the original ATHILA5 copy prior to this region being duplicated.



Figure S12. Mappability within the centromeres and CENH3 ChIP-enrichment compared between the Col-CEN and TAIR10 assemblies. A. Genome mappability was computed based on the uniqueness of *k*-mers for each genomic position, with up to *e* mismatches permitted (zero mismatches were permitted) using GenMap v1.3.0 (*107*, *108*). The uniqueness of *k*-mers, or (*k*,*e*)-mappability, was calculated for each position using 50-, 150-, 200- and 300-mers. (*k*,*e*)-mappability for a given position represents the reciprocal value of the frequency with which the *k*-mer occurs in the genome. Chromosome-scale profiles were generated by calculating mean (*k*,*e*)-mappability values within adjacent 10-kb genomic windows. **B.** CENH3 log<sub>2</sub>(ChIP/Input (purple) plotted along the Col-CEN (upper) or TAIR10 (lower) chromosomes. *CEN180* are indicated as ticks on the x-axis for forward (red) and reverse (blue) strand.



**Figure S13. Profiling DNA methylation of H3K9me2 and CENH3 ChIP DNA using ONT.** We performed ChIP-seq on Col-0 nuclei using H3K9me2 or CENH3 antibodies. The resulting DNA was then sequenced using a ONT Flongle flow cell. Reads were mapped to the Col-CEN assembly and filtered for those aligning within the centromeres. Read IDs were extracted, duplicates removed, and then used to extract fast5 files. The fast5 files were then analysed using DeepSignal-plant in order to calculate the mean methylation value for each context across each read. The boxplot shows mean DNA methylation levels across single reads for the CG, CHG and CHH sequence contexts. We observe that methylation is significantly lower in the CENH3 ChIP reads compared to H3K9me2, and that the difference is strongest for the CHG and CHH sequence contexts. CG context methylation is high in both H3K9me2 or CENH3 ChIP-seq read sets.



**Figure S14. Centromeric DNA methylation in wild type and CG and non-CG context pathway mutants. A.** Plots of CENH3 (black) and H3K9me2 (purple) ChIP-seq enrichment along chromosomes scaled proportionally along the telomere-centromere axes (*19*, *39*). DNA methylation profiles calculated from BS-seq data are plotted for CG (blue), CHG (red) and CHH (green) sequence contexts in the indicated genotypes (*35*, *42*). Comparison of Col-0 and *met1* is shown using independent data sets that were sequenced with either paired-end or single-end reads (*35*, *42*). As a comparison, DNA methylation profiles generated from ONT reads using the DeepSignal-plant and Nanopolish algorithms

are shown to the right. **B.** As for A., but comparing data from *cmt2*, *cmt3*, *drm1 drm2*, *drm1 drm2 cmt2 cmt3*, *kyp suvh5 suvh6* and *ddm1* (34, 35).



**Figure S15. Immunofluorescence analysis of euchromatic marks in isolated nuclei relative to CENH3. A.** Protein sequences of Arabidopsis H3.1, H3.3 and CENH3 were aligned using CLC Main Workbench. H3 N-terminal lysine residues known to be modified and investigated here are highlighted in red. **B.** Arabidopsis nuclei were stained for euchromatic marks (Magenta) and CENH3-GFP (green) and DNA (cyan=DAPI). The white line indicates the area of the confocal section. The confocal section is also depicted at the left bottom of each merged image. The intensity plot for the confocal section is shown on the right. Scale bars are 5μm.



Figure S16. Immunofluorescence analysis of heterochromatic marks in isolated nuclei relative to CENH3. Arabidopsis nuclei were stained for heterochromatic marks (Magenta) and CENH3-GFP (green) and DNA (cyan=DAPI). The white line indicates the area of the confocal section. The confocal section is also depicted at the left bottom of each merged image. The intensity plot for the confocal section is shown on the right. Scale bars are 5  $\mu$ m.



Figure S17. Mapping Col×Ler single nucleotide polymorphisms (SNPs) and crossovers against the Col-0 centromere assembly. A. Histograms showing the frequency of qualities, coverage, reference and variant allele coverages for single nucleotide polymorphisms (SNPs) called against the assembly using data from 260 Col×Ler genomic DNA  $F_2$  sequencing libraries. The red lines indicate thresholds where sites were filtered out of analysis. B. Histogram of crossovers mapped against the

assembly per Col×Ler  $F_2$  plant. The red dotted line indicates the mean value. **C.** Plot of the assembly showing *CEN180* satellite density per 10 kbp for forward (red) and reverse (blue) strands (upper). Beneath, the frequency per 10 kbp of total Col×Ler SNPs (red) are plotted, in addition to SNP frequency filtered for quality and coverage values, as in A (blue), and SNPs following repeat-masking (green). The lower plot shows crossovers per 10 kbp (blue) mapped against the assembly.



**Figure S18. Epitope-tagging and functional complementation of V5-DMC1. A.** Inflorescences of wild type (Col-0), *dmc1-3* and *V5-DMC1 dmc1-3*. Fertility is evident from silique length. **B.** Quantification of seed set per silique in wild type (Col-0), *dmc1-3* and *V5-DMC1 dmc1-3*. **C.** PCR based detection of the N-terminally epitope-tagged *V5-DMC1* transgene, alongside Col-0 and *dmc1-3* null controls. PCR primers flank the *DMC1* ATG translation start site. The expected PCR product sizes are 203 and 74 bp for epitope-tagged and wildtype *DMC1*, respectively. Unincorporated oligonucleotides are seen in 'no DNA' control. **D.**  $\alpha$ -V5 western blot from Col-0 and *V5-DMC1 dmc1-3* protein extracts from closed flower buds. The expected size of *V5-DMC1* is 41.7 kDa.



**Figure S19. Immunocytological staining of the Arabidopsis centromeres. A.** Somatic interphase nucleus immunostained for CENH3 (green), H3K9me2 (red) and stained for DAPI. Scale bar = 5  $\mu$ M. **B.** As for A, but showing an Arabidopsis male meiocyte in pachytene immunostained for CENH3 (green), ZYP1 (green) and H3K9me2 (red), and stained for DAPI (blue). Scale bar=5  $\mu$ M. **C.** Mitotic and meiotic cells immunostained for H3K9me2 and imaged using STED super resolution microscopy. The colour-scale indicates the intensity of staining, with yellow representing the maximum intensity. Scale bars = 5  $\mu$ M.



Figure S20. DNA methylation, RNA and siRNA expression associated with *ATHILA* elements in wild type and *met1*. A. CG, CHG and CHH context DNA methylation in wild type (Col-0, green) or

*met1* (pink/purple) measured using BS-seq (42), over *CEN180* (n=66,131), centromeric *ATHILA* (n=53), non-centromeric *ATHILA* (n=58), all *GYPSY* retrotransposons in the genome (n=3,979) and random positions (n=66,131). Shaded ribbons represent 95% confidence intervals for windowed mean values. **B.** Heatmap analysis of RNA-seq (42), siRNA-seq (42) and DNA methylation (42) data from wild type (Col-0) or *met1*. Each row represents an individual *ATHILA*, ordered according to their location within or outside the main centromeric *CEN180* arrays, and then by subfamily.



**Figure S21. Model for** *CEN180* **sequence evolution in Arabidopsis.** At the top of the diagram a representative array of five *CEN180* monomers (rectangles) is shown. Mutations, including base substitutions and replication slippage, generate monomer sequence variants (red). On the left hand side of the diagram we consider a similar representative region of five *CEN180* passing through meiosis, each of which has a distinct sequence, indicated by color. The 4 chromosomes are shown as two sisters

of each homolog. During meiotic prophase I, one chromosome experiences a DNA double strand break (DSB, red star). The DSB is processed via resection to form single stranded DNA that is bound by RAD51/DMC1, which promote invasion of another chromosome. We show four possible scenarios where the invading strand enters, (i) an allelic location on the sister chromatid, (ii) a non-allelic location on the sister chromatid, (iii) an allelic location on a homolog, or (iv) a non-allelic location on a homolog. Crossover repair, via either the Class I or Class II pathways, are suppressed within the centromere. Therefore, we propose that centromeric strand invasion events are instead repaired via meiotic noncrossover pathways, including synthesis-dependent strand annealing (SDSA), which can result in gene conversion. For simplicity conversion of single CEN180 repeats is indicated, although based on patterns of higher order repetition we propose resection and conversion may involve multiple monomer repeats (up to 60). Recombinant CEN180 arrays generated by these pathways are then subject to selection and genetic drift in populations. On the right hand side of the diagram, we indicate that DSB formation and repair within the CEN180 arrays may also occur outside of meiosis. In this case, repair may proceed via non-homologous end joining (NHEJ), or using intersister homologous recombination in either allelic or non-allelic locations. These pathways may also generate variation in CEN180 arrays that will be subject to selection and genetic drift.





Figure S22. Model for ATHILA integration and sequence evolution within the Arabidopsis centromeres. We consider a representative region of ten CEN180 monomers, with distinct monomers color-coded. The sister and homologous chromosomes are shown. A de novo ATHILA integration event is shown within one of the chromosomes. The paired long terminal repeats (LTRs, red) are shown approximately to scale, but the internal region of the transposon is not represented, but would typically consist of ~8 kbp of sequence. Following integration we consider three potential further changes to the ATHILA insertion. As we observe multiple centromeric ATHILA solo LTRs, we propose that DNA double strand break (DSB) formation and repair may occur within the ATHILA that results in formation of a solo LTR. This pathway may occur during mitosis or meiosis, and the resulting solo LTR would then be subject to selection and/or genetic drift. On the right hand side of the diagram we consider an alternative pathway during meiotic prophase I, showing two potential outcomes. In the left hand branch, a meiotic DSB (red star) forms in a CEN180 linked to the ATHILA insertion (which is hemizygous). The DSB undergoes resection to form single stranded DNA (ssDNA) which is able to invade a homologous chromosome that lacks the ATHILA insertion. Based on the large size (10-100s kbp) of CEN180 higher order repeats that we observe, we propose that an extended form of resection may occur that causes deletion of the ATHILA from the donor chromosome. The invading strand then undergoes template driven DNA synthesis that copies CEN180 sequence from a different chromosome. Following dissolution of strand invasion and non-crossover repair with the parental chromosome, the ATHILA has effectively been eliminated. The resulting chromosomes are then subject to selection and genetic drift. An alternative outcome of this pathway is shown on the right hand side. In this case, a meiotic DSB forms on the homolog that lacks the ATHILA, followed by resection, ssDNA formation and strand invasion of the homolog that carries an ATHILA insertion. In this case, template driven DNA synthesis and non-crossover repair copies and duplicates the ATHILA. We propose that this recombination process represents a mechanism to eliminate the ATHILA, as although in some situations new copies of ATHILA are generated, due to the greater abundance of CEN180 satellites in the centromeres there is a higher chance overall of this pathway eliminating the transposons.