Catastrophic chromosomal restructuring during genome elimination in plants

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Genome instability is associated with mitotic errors and cancer. This phenomenon can lead to deleterious rearrangements, but also genetic novelty, and many questions regarding its genesis, fate and evolutionary role remain unanswered. Here, we describe extreme chromosomal restructuring during genome elimination, a process resulting from hybridization of *Arabidopsis* plants expressing different centromere histones H3. Shattered chromosomes are formed from the genome of the haploid inducer, consistent with genomic catastrophes affecting a single, laggard chromosome compartmentalized within a micronucleus. Analysis of breakpoint junctions implicates breaks followed by repair through non-homologous end joining (NHEJ) or stalled fork repair. Furthermore, mutation of required NHEJ factor DNA Ligase 4 results in enhanced haploid recovery. Lastly, heritability and stability of a rearranged chromosome suggest a potential for enduring genomic novelty. These findings provide a tractable, natural system towards investigating the causes and mechanisms of complex genomic rearrangements similar to those associated with several human disorders.

Nucleosomes containing variant histone CENH3\(^1\) (also known as CENP-A) determine centromeres. In the absence of the endogenous CENH3, *A. thaliana* mitotic and meiotic functions can be complemented by chimeric CENH3\(^2,3\) or CENH3 from diverged plant species\(^4\), but crossing these strains to wild-type individuals results in frequent loss of the chromosomes marked by the variant CENH3. Following stochastic genome elimination in the early mitotic divisions, the progeny can be haploid, aneuploid or diploid\(^2,5\). In nature, similar phenomena involve defective CENH3 loading\(^6\). Thus, mating of individuals that express diverged CENH3s, can lead to mitotic catastrophe.
The consequences of mitotic malfunction on genome integrity can be dire⁷,⁸. Missegregated chromosomes can lead to aneuploidy⁹, but also to extensive and catastrophic restructuring resulting in, sequentially, chromosome sequestration in micronuclei, endonucleolytic damage, defective repair, and finally rescue¹⁰-¹². The resulting structurally rearranged chromosomes may contribute to cancer or developmental syndromes¹³-¹⁶. Nevertheless, chromosomal rearrangements are not necessarily deleterious: some may influence fitness by altering recombination or gene dosage¹⁷. It is possible that pathways leading to disease and to diversity share a common mechanistic basis¹². Genome elimination in Arabidopsis provides a previously lacking organismal system to investigate genome instability during mitotic catastrophes, connected mechanisms, and consequences.

We used the GFP-tailswap haploid inducer²,⁵ in the experimental setup illustrated in Figure 1. This strain is in the Col-0 background and carries a homozygous CENH3 null mutation whose function is partially complemented by a chimeric CENH3 in which an N-terminal GFP fused to the H3.3 N-terminal tail replaces the native CENH3 N-terminal tail. We crossed this strain to polymorphic accession Ler gl1-1 to track haplotypes in the F1 progeny and obtained the expected haploid induction frequency²,⁵ (Figure 1). The recessive gl1-1 mutation confers trichomeless leaves in paternal Ler gl1-1 haploids while it is masked in Col/Ler diploid hybrids. We sequenced 10 of the phenotypically diploid Col/Ler individuals with wild-type phenotype, performed dosage plot and SNP analysis and found that 100% of these were diploid with 50% Col and Ler genomes respectively (Figure 1–figure supplement 1). Plants from the aneuploid class exhibited multiple pleiotropic and morphological defects and had trichomes, except in the
rare exception when the GL1 locus was lost. The five recognizable primary trisomic (2n + 1) phenotypes were represented\textsuperscript{18,19}: Chromosome 1 (Chr1) trisomics have dark green, serrated leaves and are dwarfed, Chr2 trisomics exhibit round leaves and are late flowering, Chr3 trisomics have narrow, yellow green leaves, Chr4 trisomics display narrow and smaller flat leaves, and Chr5 trisomics display light green and narrow leaves. However, aneuploid plants with more severe or unusual phenotypes were also observed, suggestive of other chromosomal combinations or more serious chromosomal aberrations. Chromosome dosage analysis based on whole genome sequencing\textsuperscript{20} (Supplementary file 1) distinguished three chromosomal alteration types in aneuploids (Figure 2). Similar outcomes were obtained using independently derived haploid inducers, either expressing GFP-tailswap (Figure 2E and Figure 2–figure supplement 1) or CENH3 from other plant species\textsuperscript{4}. The most common type, numerical aneuploids, display whole chromosome aneuploidy such as in the classical primary trisomics (Figure 2B shows an example for a numerical Chr3). In our dataset, single primary trisomics (2n + 1) account for 75% of the numerical class. Other individuals from the numerical class with two or more extra whole chromosomes included 16% double primary trisomics (2n + 1 + 1), 2% triple primary trisomics (2n + 1 + 1 + 1) and 3% quadruple primary trisomics (2n + 1 + 1 + 1 + 1). Additionally, we obtained disomic Chr4 haploids (n + 1, a type of numerical aneuploidy that were not included in this analysis) as well as Chr2 or Chr3 monosomic diploids (2n – 1) at 4% frequency (Figure 2–figure supplement 2). These have never been described in Arabidopsis before, possibly because, if they were to arise from meiotic defects, they would result from nullisomic gametes, which are not viable\textsuperscript{21}. Aneuploids resulting from mitotic failure do not have those constraints.
The second alteration type is defined by simple truncations and repair of at most two double stranded DNA breaks per chromosome (Figure 2C shows an example of truncated Chr3). This truncated class was found to occur in 22% of the aneuploid population. In the third class, a single chromosome exhibited many oscillations in copy number state, as if shattered and subsequently rearranged (Figure 2D shows an example of shattered Chr3). This shattered class was found to occur in 11% of the aneuploid population. Additionally, some of the aneuploids exhibited a combination of numerical, truncated and shattered chromosome types (Figure 2E). Alteration of copy number for Chr1, 2 and 3 are represented at similar frequencies based on the average copy number alteration of all five chromosomes, with Chr4 and 5 alterations being, respectively, over- and under-represented (Figure 2F). This may be explained by the uneven distribution between chromosomes of few, selected genes that are highly dosage-sensitive. According to this hypothesis, Chr4 would be selectively depleted for such genes.

Chromosomal truncations have been reported from a selfed trisomic22. To assess whether truncated and shattered aneuploid types could be produced from meiotic missegregation, we sequenced 96 individuals produced by a selfed Col-0 triploid. Because of the irregular meiosis, most gametes produced by triploids are aneuploid23. Dosage analysis revealed that all were numerical aneuploids (Supplementary file 2 and Figure 2–figure supplement 3). To assess whether truncation and shattering could be the result of meiotic defects in the GFP-tailswap line, we sequenced 96 individuals from selfed GFP-tailswap and observed that 98% (n = 94) of the progeny were diploid while two individuals carried single primary trisomies of Chr2 and Chr3 respectively, representing only the numerical class of aneuploids (Supplementary file 3 and Figure
Based on these results, we believe that truncated and shattered aneuploid classes from our crosses reflect genomic instability associated with mitotic errors in the early embryo.

Shattered chromosomes can be recovered from all five *Arabidopsis thaliana* chromosomes (Figure 3A). In some cases, shattering appears to extend to two chromosomes (top panel of Figure 3A) only because the haploid inducer used carries a reciprocal Chr1/Chr4 translocation originating from the integration of GFP-tailswap T-DNAs. SNP analysis demonstrates that all duplicated (copy number 3) and triplicated (copy number 4) regions originated from the haploid inducer (Figure 3B). Single-copy regions displaying loss of heterozygosity carry Ler alleles (i.e. wild-type), consistent with the loss of the haploid inducer haplotype.

Although aneuploids from the shattered class were often sterile, line FRAG00062 was partially fertile and allowed us to investigate the inheritance and stability of the variant DNA. We sequenced sixteen F2 progeny from FRAG00062 and obtained two individuals with precisely the same shattered pattern as the F1 parent and fourteen that appeared diploid (Figure 4A). Meiotic co-inheritance of all dosage variant segments is consistent with a single, stable chromosomal unit that was formed after a catastrophe. To confirm this hypothesis, we used DNA fluorescence *in situ* hybridization (FISH) to visualize the FRAG00062 chromosomes using Col-0 derived BAC painting probes specific for Chr1 and Chr4 (Figure 4B). Mitotic cells contained 11 chromosomes (Figure 4D). FRAG00062 came from a cross using GFP-tailswap line #11, which carried a reciprocal Chr1/4 translocation (Figure 2–figure supplement 1). This allowed us to distinguish the haploid inducer Chr1, the Ler Chr1, and a third Chr1 with rearranged
signals, which we interpret as the shattered extranumerary chromosome (Figure 4C and Figure 4–figure supplement 1). During meiotic Metaphase I (Figure 4D) or other meiotic stages observed from male meiocytes (Figure 4–figure supplement 2), the shattered chromosome does not pair with the parental Chr1s.

Next, we sought to investigate why shattering is restricted to a single chromosome. During genome elimination crosses in other plant species, micronuclei are commonly observed\textsuperscript{24,25}. We dissected embryos from a genome elimination cross and observed one to four micronuclei per cell (Figure 4E–F) in 81% of the embryos ($n = 110$), but none in embryos from control crosses ($n = 21, p < 0.001$). The presence of micronuclei suggests that sub-compartmentalized lagging chromosomes can be shattered by double stranded DNA breaks, reassembled haphazardly by NHEJ, and finally restituted into the main nucleus\textsuperscript{10}.

In order to reconstruct breakpoint junctions, we sequenced FRAG00062 to 100x coverage, extracted read pairs from the ends of duplicated and triplicated blocks and performed \textit{de novo} assembly. Thirty-eight such junctions were assembled (Supplementary file 4) and a random subset of 12/12 were confirmed by PCR (Figure 5–figure supplement 1) followed by Sanger sequencing to demonstrate the accuracy of the \textit{de novo} assembly. All reconstructed junctions were consistent with NHEJ with either microhomology, observed as 2–15 bp of sequence overlap\textsuperscript{16}, blunt fusions, or unidentified sequence insertions (Figure 5B). We also observed inversions (fragments that join in head to head or tail to tail orientation) in 47% of our breakpoint junctions (Supplementary file 4). The size distributions of microhomology tracts and insertions are indicated in Figure 5–figure supplement 2.
Overall, triplicated block sizes (n = 23) from FRAG00062 were significantly smaller than duplicated blocks (n = 23, with p < 0.001, Figure 5C) and these triplications cannot be easily explained from a missegregated chromosome. Duplicated and triplicated blocks could therefore, have different origins. To address this question, we asked whether breakpoint junctions of the two different copy number states display differential association to various genomic and chromatin features such as genes and repeated elements, DNA replication origins, DNase I hypersensitive sites and nine non-overlapping chromatin states that partition the Arabidopsis genome (Supplementary file 5). When analyzing windows of 1,000 bp centered around the breakpoints of duplicated blocks, we observed an enrichment in genic DNA (from 53% background level to 70%, p < 0.01, Figure 5D and 5F). A subtler, but still significant, increase was observed when using larger windows (10,000 bp, from 53% background level to 62%, p < 0.01, Figure 5F). Consistently, 42% of breakpoint junctions from FRAG00062 are predicted to generate chimeric gene products (Supplementary file 4). In the same analysis, we noted that the breakpoint regions of duplicated and triplicated blocks contained some genomic features that differed in frequency. In particular, replication origins, which occupy less than 1% of 10,000 bp windows around the borders of duplicated blocks, are present in almost 8% around the borders of triplicated blocks (compared to a genome average of 3.5%, p < 0.05, Figure 5E and 5G). The association of the breakpoints flanking duplicated DNA to genic DNA and of those flanking triplicated DNA to replication origins suggests the contribution of two distinct mechanisms to restructuring of the same chromosome (Figure 6). The first,
chromothripsis acting through breakage and ligation\textsuperscript{15,30}. The second, chromoananasynthesis, via replication fork collapse and template switching\textsuperscript{14,16,31}.

Our \textit{in silico} reconstruction suggests that NHEJ is involved in repairing breaks that occurred on the shattered chromosomes. To test this explanation, we created a haploid inducer carrying a homozygous null mutation in \textit{LIG4} (DNA Ligase IV), a conserved component of the canonical NHEJ pathway. Pollinating it with wild-type \textit{LIG4}/\textit{LIG4} pollen (from \textit{Ler gl1-1}) resulted in normal haploid induction frequencies. However, when mutant \textit{lig4-2}/\textit{lig4-2} pollen was used, the frequency of haploids doubled at the expense of aneuploids and diploids (\textbf{Table 1} and \textbf{Figure 7}). This effect was still observed when the seed parent carried the WT allele (\textbf{Table 1}). It is possible that parental-specific haploinsufficiency results from early loss of the wild-type \textit{LIG4} allele located on the chromosome targeted for elimination, which in this case is the maternal chromosome. This result indicates that NHEJ contributes to formation or persistence of aneuploid and diploid progeny and that unpaired double-stranded DNA breaks increase elimination of the haploid inducer genome, similar to observations in mouse-human hybrid genome elimination\textsuperscript{32}. We hypothesize that missegregated chromosomes enter a degradative pathway initiated by endonucleolytic breaks. Occasionally, such chromosomes are rescued (i.e. restituted to a haploid or diploid nucleus) through a pathway requiring NHEJ, resulting in aneuploidy. Therefore, more haploids are produced when the NHEJ pathway is impaired (\textbf{Figure 6}).

Taken together, our results provide evidence for the occurrence of chromosome restructuring\textsuperscript{33,34} when diverged individuals hybridize, identifying a centromere-based mechanism for genomic instability. This phenomenon studied here depends on chimeric
CENH3, but a similar effect was observed when the haploid inducer strain expresses CENH3 of a close species\textsuperscript{4}, indicating the effectiveness of natural and artificial variation. While the genesis and fate of restructured chromosomes is difficult to study in humans, their formation, effects, and even transmission in \textit{Arabidopsis} are within experimental reach, as demonstrated by the enhancing effect of NHEJ mutants on haploid induction. The range of phenotypes, the formation of copy variants and of chimeric genes at junctions, and their occasional meiotic transmission, suggest that catastrophic chromosomal restructuring, could contribute to heritable genetic variation.
Any Supplementary data is available in the online version of the paper. All datasets have been deposited to Sequence Read Archive under SRP051357 with the Bioproject ID PRJNA269994.

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Author Contributions S.W.L.C., M.R., I.M.H., L.C. and E.H.T. contributed to conception and overall experimental design. E.H.T. performed all genetic and genomic analyses with assistance from I.M.H. and M.R.. T.M. and M.A.L. planned and performed the chromosome painting experiments, K.B. performed the bioinformatics breakpoint analysis, I.K. advised on bioinformatics. M.P.A.M. performed embryo imaging experiments. E.H.T., I.M.H. and L.C. wrote the paper. All authors discussed results and
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FIGURES LEGENDS

Figure 1. Aneuploid types resulting from centromere-mediated uniparental genome elimination.

The altered CENH3 “GFP-tailswap” strain was hybridized to the recessive glabrous1-1 mutant. Mean percentages of haploid, diploid an aneuploid progeny obtained from crosses to three independent GFP-tailswap lines are indicated, as determined after phenotypic characterization. Individuals belonging to the aneuploid class were sequenced and subjected to chromosome dosage and SNP analysis as indicated by the arrow.

Figure 2. Characterization of the three distinct aneuploid types from GFP-tailswap haploid induction crosses.

(A–D). Dosage plots from all five Arabidopsis chromosomes in consecutive non-overlapping 100 kbp bins and the corresponding SNP plot for the % haploid inducer genome (Col-0) present in each sample. A diploid Col/Ler hybrid (A), an individual with primary Chr3 trisomy from the numerical aneuploid class (B), an individual with a truncated trisomic Chr3 (C) and an individual with shattered Chr3 (D) are shown here. Centromere positions are indicated by red diamonds.

(E). Percentages of the different aneuploid types obtained from three different GFP-tailswap haploid inducer lines.

(F). For each chromosome, the percentage of aneuploid individuals exhibiting altered dosage for that particular chromosome is plotted. All aneuploids characterized in this study are included. Chr4 is overrepresented (** Student’s t-test, p < 0.01) while Chr5 is underrepresented (* Student’s t-test, p < 0.05).
**Figure 3.** Shattered chromosomes are confined to a single chromosome originating from the haploid inducer.

(A). Chromosome dosage plots based on non-overlapping 25 kbp bins across each chromosome for five aneuploid individuals with shattered chromosomes. The GFP-tailswap transgene insertion event that resulted in a reciprocal translocation between Chr1 and Chr4 in one of the haploid inducer parent (GFP-tailswap #11) is indicated with black arrowheads. The translocation is only visible in individuals for which chromosomes 1 and 4 are not balanced with each other. Duplications (copy number 3), triplications (copy number 4) as well as deletions accompanied with loss of heterozygosity (copy number 1) were observed from dosage plots.

(B). Box plots of the percentage of haploid inducer genome present at each copy number state, as determined by the SNP analysis. Mean and standard errors are shown.

**Figure 4.** Stable inheritance and chromosome painting of a shattered aneuploid chromosome.

(A). Dosage analysis from sixteen F2 individuals from a selfed FRAG00062 individual. Progeny individuals either inherited the shattered chromosome intact (n = 2) or appeared diploid (n = 14).

(B). Cartoon of the different versions of chromosomes 1 and 4 expected to be present in FRAG00062. Chromosome painting probes and corresponding chromosome positions used for c and d are shown. Black triangles indicate the position of the reciprocal Chr1/Chr4 translocation present in the haploid inducer line, whereas black circles indicate centromere positions.
(C). A mitotic cell from FRAG00062 with 11 chromosomes, including four painted chromosomes. Scale bar = 5μm.

(D). The shattered Chr1 from FRAG00062 remains unpaired at meiosis as shown here at Metaphase I. Enlargements of the shattered Chr1 and paired Chr1 are shown on the right. Scale bar = 5μm.

(E). Nuclei from a two-cell stage embryo from a wild-type cross (left panel) and from an embryo undergoing uniparental genome elimination (right panel). Nuclei are visualized using CFP-tagged histone H2B from the pollen parent superimposed with an image of the embryo visualized under light microscopy. Note the presence of micronuclei from the embryo undergoing genome elimination (right panel). Scale bar = 5μm.

(F). Percentage of micronuclei observed in wild-type crosses and genome elimination crosses. The different percentages of micronuclei per cell are indicated.

Figure 5. Breakpoint junctions and enriched features surrounding breakpoints of duplicated and triplicated blocks.

(A). Plot of all chromosomes and corresponding dosage plots (blue line) from FRAG00062. Black curves depict novel junctions identified from genomic reconstruction of the shattered haploid inducer Chr1. Black triangles indicate the reciprocal translocation between Chr1 and Chr4 from the haploid inducer genome.

(B). Percentage of junctions with 2–15 bp of microhomology, blunt junctions or junctions with unidentified sequence insertion observed from FRAG00062.

(C). Plot of duplicated (n = 23) and triplicated (n = 23) block sizes from FRAG00062. (** Student’s t-test p < 0.001)
Occupancy of genes and replication origins around breakpoints regions, calculated using windows of 1,000 bp or 10,000 bp centered on each breakpoint. Dashed horizontal lines indicate the genome-wide occupancy of each feature. Error bars indicate standard error.

Enrichment ratio of genes and replication origins (See methods for description of enrichment ratio). Genes (F) are significantly enriched surrounding duplicated breakpoints regardless of window size (1,000 or 10,000 bp, ** p < 0.01). For windows of 10,000 bp, replication origins (G) are significantly enriched at triplicated breakpoints (* p < 0.05) while significantly under-represented (‡ p < 0.05) at duplicated breakpoints.

Figure 6. The process of genome elimination and connected models for chromosomal rearrangements.

Genome elimination ensues when a haploid inducer expressing a variant CENH3 protein mates with the wild type. In many cases, the chromosomes marked by the variant CENH3 missegregate in the embryo and are compartmentalized in micronuclei. DNA damage, NHEJ repair and restitution of the micronucleus to the euploid pole nucleus can result in aneuploidy or diploidy. Alternatively, shattered chromosomes result from chromothripsis and chromoanasynthesis. The former involves fragmentation and random ligation, the latter replication fork collapse and microhomology-mediated strand switching. As a consequence, the pulverized and reassembled chromosome forms a single unit and can be meiotically inherited. The schematics for chromothripsis and chromoanasynthesis are shown sequentially for convenience, but their order has not been determined. In addition, our results obtained using DNA ligase4-2 mutants suggest that the NHEJ pathway plays
an important role in the repair of the haploid inducer chromosomes that contribute to diploid and aneuploid progeny, such that when NHEJ is inhibited, haploid induction frequency increased.

**Figure 7. Dosage plots for lig4-2 haploids isolated from a haploid induction cross using diploid lig4-2 as the male donor.**

Dosage plots of lig4-2 haploids based on 150 kbp non-overlapping bins across all five *Arabidopsis* chromosomes. Euploid chromosome dosage plots for lig4-2 haploids have the appearance of having a copy number of 2 only because euploid chromosome dosage was calculated with the value of 2 in this analysis. Centromere positions are indicated by red diamonds.
MATERIALS AND METHODS

Plant material and growth conditions.

All plants were grown in Sunshine Professional Mix Peat-Lite Mix 4 (SunGro Horticulture) under 16h/8h light/dark photoperiod in a growth room set at 21°C. F1 seeds from GFP-tailswap crosses were germinated on MS agar plates and 2-week old seedlings were transplanted into soil. The lig4-2 (SAIL_597_D10) line used is in the Col-0 background. Genotyping primers (5’ to 3’) used to are lig4-2/LP2: GATATGACAAGCCTTGGCATGAATGT, lig4-2/RP: AAAGTGGATGACATCTCGCTG and LB1: GCCTTTTCAGAAATGGATAAATAGCCTTGCTTCC for the left border of the SAIL T-DNA insertion.

Genomic DNA preparation, sequencing and read processing. All DNA samples were extracted from leaves using Nucleon Phytopure kits (GE Healthcare). 1.5 μg of DNA were used for a PCR-free library preparation using the NEBNext DNA Library reagents with Nextflex-96 indexes (Bioo Scientific) using a PCR-free protocol. 2 μl of each 96-barcoded libraries were pooled and sequenced using the 50 bp protocol on a single lane of Hiseq 2000 at the Vincent J. Coates Genomics Sequencing Laboratory at UC Berkeley. Demultiplexing was performed by the same facility and resulting raw reads were processed with a custom Python script (Filter_N_Adapter_Trim_Batchmode.py – available from GitHub repository: https://github.com/KorfLab/FRAG_project) that removes the filtered reads from Cassava 1.8, adapter sequences, reads that contain Ns and trims reads for quality.
Chromosome dosage analysis.

For dosage plot analyses, 50 bp single reads were mapped to the TAIR10 *A. thaliana* reference genome sequence using BWA$^{35}$ and default parameters. Dosage variation was detected as previously described$^{20}$. The genomic reference chromosomes were partitioned into consecutive non-overlapping bins of 100,000 bp and the percentage of reads mapping to each bin from each sample was recorded. Relative coverage was calculated by dividing the percentage obtained for each bin by either the corresponding mean percentage for all individuals or the corresponding percentage for the control individual. The relative coverage was set at 2 to represent the diploid background copy value.

Single Nucleotide Polymorphism (SNP) analysis

Positions polymorphic between Col-0 and Ler were identified using sequencing reads from a diploid Col/Ler hybrid control, a Ler plant and a Col-0 plant using custom python scripts. Specifically, polymorphic positions were first identified if they were covered at least 25 times in the hybrid reads and contained two alleles, each representing at least 40% of the allelic calls. Reads from the Col-0 and Ler parents were then used to assign alleles to the two parents. Positions were only retained if they were homozygous in both parents (represented at least 97% of the allelic calls) and covered at least 6 times in the Col-0 library and at least once in the Ler library. This process resulted in the identification of 107,640 SNP positions (Supplementary file 6). Next, reads from each of the samples were mined for allele calls at these positions and each read was assigned to one or the other parent based on the parental information. If the read did not match
either allele, the genotype was reported as “na”. Finally, genotype information was pooled by consecutive, non-overlapping bins of 1 Mb to derive a percentage of Ler allele per bin for each sample. Using this measure, the Col-0/Ler diploid hybrid is expected to exhibit 50% Col-0 across the genome.

Cytogenetic analysis

All analyses were carried out using chromosome spreads from young anthers. BAC contigs specific for *A. thaliana* chromosomes 1 and 4 were used as painting probes. BAC DNA was labeled with biotin-, digoxigenin- or Cy3- deoxyuridine triphosphate by nick translation as previously described. Labeled DNA probes were pooled, hybridized to suitable chromosome spreads and visualized using fluorescent microscopy. See Supplementary file 7 for the list of BAC clones used as painting probes.

Breakpoint assembly

Breakpoints from FRAG00062 were identified using a high-density 500 bp bin-size dosage plot produced using 50 bp reads extracted from 100 bp paired-end sequencing reads of the FRAG00062 library obtained from an Illumina HiSeq 2000 instrument. Blocks of duplicated or triplicated dosage were defined by eye. A custom script (batch-specific-junction-search.py – available from GitHub repository: https://github.com/KorfLab/FRAG_project) was used to extract the sequencing reads mapping within a 2,000 bp region around each breakpoint. These sequences were then assembled using the PRICE genome assembler using the standard paired-end assembly setting. Resulting contigs were aligned to the *Arabidopsis* reference genome by NCBI-
BLASTN and characteristic breakpoint junctions were identified when two halves of a contig mapped disconcordantly to the reference genome. Primers flanking 12 randomly selected breakpoint junctions were designed using Primer3 based on their respective de novo assembled contigs. Standard PCR procedures were used for amplification using oligo pairs (Supplementary file 8) and GoTaq Green Mastermix (Promega) on 1 ng DNA from FRAG00062 and FRAG00080 (a diploid sibling control) followed by Sanger sequencing.

**Breakpoint analysis**

The *A. thaliana* TAIR10 genome annotation includes genomic locations for various features in Generic Feature Format Version 3 (GFF). Files specifying genes, transposon, satellite repeats, and replication origins were downloaded from the TAIR FTP site (ftp://ftp.arabidopsis.org//Maps/gbrowse_data/TAIR10/). The GFF file containing the location of mapped replication origins was available from a study by Costas et al. These GFF files were combined with results about mapped DNase I Hypersensitive Sites (DHS) and details from the recent work by Sequeira-Mendes et al., which combined various published epigenomic studies to partition the entire genome into nine different chromatin states. Perl scripts were used to convert the DHS and chromatin state information into GFF format, and these scripts, along with the resulting combined GFF file are available from a GitHub repository: [https://github.com/KorfLab/FRAG_project](https://github.com/KorfLab/FRAG_project). The set of genome features in the combined GFF file were compared to the annotated set of duplicated and triplicated blocks. Various Perl scripts available from the above GitHub repository, along with a GFF representation of all blocks were used to assess the
enrichment of genomic features at the breakpoint regions of duplicated/triplicated blocks.

Specifically, window sizes of either 1,000 or 10,000 bp were centered on each breakpoint coordinate, and the number of bp contributed by each feature of interest were summed across all windows. We also calculated the number of bp contributed by each feature outside those windows.

Enrichment ratios were then calculated using the percentage of bases occupied by each feature across all windows at breakpoints compared to percentage of the same features that occupy the remaining fraction of the aneuploid chromosome. The p-values were determined by shuffling experiments in which the locations of the breakpoints were randomized 1000 times, with the resulting shuffled ratios compared to the ratios observed in the real data (Supplementary file 5).
Figure 1—figure supplement 1. Dosage plots and SNP analysis of diploids from GFP-tailswap haploid induction crosses.

(A). Dosage plots from all five Arabidopsis chromosomes in consecutive non-overlapping 100 kbp bins.

(B). The corresponding SNP plot for the % haploid inducer genome (Col-0) present in each sample.

Figure 2—figure supplement 1. Copy number and T-DNA positions of the GFP-tailswap transgene in three independently derived transgenic lines.

(A). The three lines used in this study and the insertion sites for each GFP-tailswap transgene are shown. Lines #11 and #17 carry four copies of the transgene and the transformation event resulted in a reciprocal translocation of Chr1 and Chr4 for #11 and Chr1 and Chr2 for #17. Line #22 carries a single copy insertion on Chr5.

(B–C). Exact positions of the insertion sites for line #11 (B) and line #17 (C). T-DNA left borders are denoted by ‘LB’.

Figure 2—figure supplement 2. Dosage plots and SNP analysis of atypical aneuploids from GFP-tailswap haploid induction crosses.

(A–D). Dosage plots from all five Arabidopsis chromosomes in consecutive non-overlapping 100 kbp bins and the corresponding SNP plot for the % haploid inducer genome (Col-0) present in each sample. A monosomic Chr2 and Chr3 individual with 2n
10 chromosomes (A, B), a disomic Chr4 individual with 2n = 5 + 1 chromosomes (C) and a haploid individual with a Chr1 minichromosome (truncated class) derived from the haploid inducer genome (D) are shown. The unusually high dosage spike around the centromere of Chr2 in FRA00870 (A) results from monosomy Chr2. Ler sequences are mapped to the Col-0 genomic reference followed by normalization to our diploid control F1 hybrid Col/Ler. Euploid chromosome dosage plots for haploids with disomic chromosomes or minichromosomes (C, D) have the appearance of having a copy number of 2 only because euploid chromosome dosage was calculated with the value of 2 in this analysis. Centromere positions are indicated by red diamonds.

**Figure 2—figure supplement 3. Diversity of primary trisomic aneuploids derived from a selfed triploid (Col-0 ecotype).**

(A). Primary trisomics (2n = 10 + 1) for all five Arabidopsis chromosomes.

(B–D) Higher order primary trisomics from our analysis. We observed (B) double primary trisomics (2n = 10 + 1 + 1), (C) triple primary trisomics (2n = 10 + 1 + 1 + 1) as well as (D) quadruple primary trisomics (2n = 10 + 1 + 1 + 1 + 1). ‘N/A’ indicates the karyotypes that were not observed in this population.

**Figure 2—figure supplement 4. Representative dosage plots from 96 individuals from a selfed GFP-tailswap haploid inducer.**

Representative dosage plot of diploid individuals (94/96 = 98%) showing a euploid dosage from all five *Arabidopsis* chromosomes. We detected two selfed individuals with
numerical whole chromosome primary trisomies (2n + 1) for Chr2 and Chr3, which are shown here.

Figure 4–figure supplement 1. Analysis of duplicated and triplicated blocks from FRAG00062.

(A). Cartoon of duplicated and triplicated blocks on the shattered aneuploid chromosome from FRAG00062.

(B). Rearranged pattern from DNA FISH with the corresponding *de novo* assembled contigs.

Figure 4–figure supplement 2. DNA fluorescence *in situ* hybridization (FISH) on the shattered aneuploid chromosome from FRAG00062.

(A). Cartoon of the regions used for chromosome painting experiments.

(B–D). Meiotic spreads from male meiocytes with DNA painting probes shown in a. Scale bars = 5μm. The unpaired shattered chromosome is visible in Pachytene (B) and Diakinesis (C) and separate precociously at Anaphase I (D).

Figure 5–figure supplement 1. PCR confirmation of breakpoint junctions for FRAG00062.

DNA gel showing PCR assays performed using oligo pairs from a control region and from breakpoint junctions specific to FRAG00062 using DNA from a diploid sibling (top panel) or FRAG00062 (bottom panel).
Figure 5–figure supplement 2. Breakpoint junctions types from FRAG00062.

(A–B). Top: Junction sequence from direct sequencing of breakpoint junctions. Middle and bottom: Reference *A. thaliana* sequence. Sequences in common with the junction sequence are in blue and red. (+) denote sequences that align to the forward strand of the reference genome while (−) denote sequences that align to the reverse strand. Regions of microhomology are highlighted in yellow (A). Inserted sequence is highlighted in green (B).

(C). Distribution of the sizes of microhomology tracts from breakpoints junctions.

(D). Distribution of the sizes of unidentified inserted DNA sequences from breakpoint junctions grouped in 25 bp bins.

Supplementary file 1. List of the aneuploid individuals from *GFP*-tailswap x *Ler glI-1* crosses used in this study.

Unique identifiers (FRAG IDs) are listed for each individual, along with the *GFP*-tailswap parental lines as well as their corresponding aneuploid chromosome types. The final sixteen columns indicate whether individuals exhibited the numerical, truncated or shattered aneuploid chromosome type, and which specific chromosomes displayed these features.

Supplementary file 2. List of aneuploid individuals obtained from selfed Col-0 triploids.
Unique identifiers (FRAG IDs) are listed for each individual and indicate which specific chromosomes displayed the numerical aneuploid chromosome type.

**Supplementary file 3. List of diploid and aneuploid individuals obtained from selfed GFP-tailswap.**

Unique identifiers (FRAG IDs) are listed for each individual and indicate which specific chromosomes displayed the numerical aneuploid chromosome type.

**Supplementary file 4. Breakpoint junctions from FRAG00062.**

The positions and types of repair for each breakpoint junction are indicated, along with their orientation as well as possible outcomes for a novel gene product.

**Supplementary file 5. Enrichment ratios of genomic features surrounding breakpoints from duplicated or triplicated blocks.**

The ratios of genomic features listed in column 1 are calculated using 1,000 bp and 10,000 bp windows surrounding breakpoints from duplicated or triplicated blocks. Regions with enriched ratios has a value of > 1 while regions that are under-represented have values of < 1. Light orange boxes are indicated for $p < 0.05$ and light red boxes for $p < 0.01$.

**Supplementary file 6. Single Nucleotide Polymorphisms between Col-0 and Ler.**

A list of SNPs from Col-0 and Ler used in this study and their chromosomal positions.
Supplementary file 7. BAC clones used for chromosome painting.
Listing of the corresponding chromosome, BAC clone and associated GenBank accession used.

Supplementary file 8. Oligonucleotides sequences for analysis of breakpoint junctions from FRAG00062.

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<th>Haploid Inducer ♀</th>
<th>♂</th>
<th>Total</th>
<th>Aneuploid</th>
<th>Diploid</th>
<th>Haploid</th>
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<tr>
<td>GFP-tailswap</td>
<td>Ler gl1</td>
<td>606</td>
<td>33%</td>
<td>28%</td>
<td>39%</td>
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<td>83% *</td>
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<td>29%</td>
<td>31%</td>
<td>40%</td>
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<tr>
<td>lig4-2 GFP-tailswap</td>
<td>lig4-2</td>
<td>159</td>
<td>14%</td>
<td>5%</td>
<td>81% *</td>
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Table 1 | Haploid induction frequency from genome elimination crosses using lig4-2 mutants.
Haploid inducer lines or haploid inducer line with the lig4-2 mutation were crossed with wildtype Ler gl1 or lig4-2 mutant pollen. (* Student’s t-test, p < 0.001)
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<th>Volume</th>
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<td>Science</td>
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<td>(1984)</td>
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<td>Genetics</td>
<td>48</td>
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<td>(1963)</td>
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</table>
Haploid Inducer
\textit{cenh3-1} \textit{-/-
GFP-tailswap}
(Col-0 ecotype)

\textbf{X}

\textit{gl1-1} \textit{-/-
(Ler ecotype)}

Uniparental Genome Elimination

Aneuploid Class

\begin{itemize}
  \item 37\% Aneuploid Col/Ler \textit{gl1-1} \textit{-/+}
  \item 25\% Diploid Col/Ler \textit{gl1-1} \textit{-/+}
  \item 39\% Haploid Ler \textit{gl1-1} \textit{-}
\end{itemize}

Euploid Classes

Whole Genome Sequencing + Chromosome Dosage & SNP Analysis