De novo centromere formation on chromosome fragments with an inactive centromere in maize (Zea mays)

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Abstract The B chromosome of maize undergoes nondisjunction at the second pollen mitosis as part of its accumulation mechanism. Previous work identified 9-Bic-1 (9-B inactivated centromere-1), which comprises an epigenetically silenced B chromosome centromere that was translocated to the short arm of chromosome 9(9S). This chromosome is stable in isolation, but when normal B chromosomes are added to the genotype, it will attempt to undergo nondisjunction during the second pollen mitosis and usually fractures the chromosome in 9S. These broken chromosomes allow a test of whether the inactive centromere is reactivated or whether a de novo centromere is formed elsewhere on the chromosome to allow recovery of fragments. Breakpoint determination on the B chromosome and chromosome 9 showed that mini chromosome B1104 has the same breakpoint as 9-Bic-1 in the B centromere region and includes a portion of 9S. CENH3 binding was found on the B centromere region and on 9S, suggesting both centromere reactivation and de novo centromere formation. Another mini chromosome, B496, showed evidence of rearrangement, but it also only showed evidence for a de novo centromere. Other mini chromosome fragments recovered were directly derived from the B chromosome with breakpoints concentrated near the centromeric knob region, which suggests that the B chromosome is broken at a low frequency due to the failure of the sister chromatids to separate at the second pollen mitosis. Our results indicate that both reactivation and de novo centromere formation could occur on fragments derived from the progenitor possessing an inactive centromere.

Keywords CENH3 · Centromere reactivation · De novo centromere formation

Abbreviations 9-Bic-1 9-B inactivated centromere-1 9S Short arm of the chromosome 9 BFB Breakage fusion bridge CENH3 Centromeric histone H3
ChiP-seq Chromatin immunoprecipitation-sequencing
CNV Copy number variation
CRM Centromeric retrotransposon of maize
DAPI 4',6-diamidino-2-phenylindole
FISH Fluorescence in situ hybridization

Introduction

Eukaryotes require a functional centromere for faithful transmission of genetic material during cell division. The centromere is the region of DNA underlying the kinetochore, which is a complex structure of protein and RNA that orchestrates chromosome movement during mitosis and meiosis by serving as a scaffold to attach the chromosome to microtubules (Wang and Dawe 2018). Chromosomes contain only one active centromere, as multiple active centromeres could attach to different spindle poles and rip a chromosome apart during cell division, but exceptions do exist (Zhang et al. 2010) and holocentric chromosomes have attachments along the length of the chromosome (for a review, Bures et al. 2012).

Centromeres typically contain highly repetitive, species-specific DNA sequences (for review: Hartley and O’Neill 2019). Many studies have shown that the DNA sequences found in canonical centromeres are neither required, nor sufficient, to establish kinetochore formation (for review: Burrack and Berman 2012). The DNA sequence of maize (Zea mays) centromeres is composed of a 156-bp centromere-specific satellite repeat CentC and Centromeric Retrotransposon of Maize (CRM) family members (Jiang et al. 1996; Ananiev et al. 1998; Wolfgruber et al. 2009). Supernumerary maize B chromosome centromeres have similar DNA components, but CentC and CRM elements are joined by arrays of a B-specific repeat (ZmBs) in the centromeres of the maize B chromosome, which enables easy identification of the B centromere in cytological spreads (Lamb et al. 2005; Blavet et al. 2021).

When specific DNA sequences are unable to form functional centromeres, centromere specification is said to be epigenetic. Several epigenetic marks are associated with active centromeres in plants including the presence of the centromere-specific histone H3 variant CENH3 (CENP-A in mammals) (Zhong et al. 2002), which replaces ~15% of canonical H3 nucleosomes within animal and yeast centromeres (Joglekar et al. 2008; Johnston et al. 2010; Bodor et al. 2014), pericentromeric phosphorylation of histone H3 Ser-10 (Manzanero et al. 2000; Houben et al. 2007a, 2007b; Gao et al. 2011), and histone H2A Thr-133 phosphorylation (Dong and Han 2012; Fu et al. 2013).

The maize B chromosome is a supernumerary chromosome that occurs in only some lines and can vary in copy number amongst individuals that carry it. Presence of a low number of B chromosomes is generally neither advantageous nor deleterious, although plants containing many B chromosomes have reduced fitness and/or mild leaf phenotypes such as striping (Staub 1987). Since B chromosomes provide no selective advantage for their host, they require a drive mechanism to maintain themselves in populations. Maize B chromosomes utilize nondisjunction in the second pollen mitosis, and the process requires trans-acting factors located on the long arm of the B chromosome (Roman 1947; Lin 1978; Lamb et al. 2006). The second aspect of the drive mechanism involves preferential fertilization of the egg versus the central cell by the sperm containing the B chromosomes (Roman 1948).

The B-A translocation stock TB-9Sb contains an intact, functional B centromere attached to the short arm of chromosome 9 (9S; Robertson 1984). Through a series of translocations and rearrangements, the active B centromere of TB-9Sb was transferred to the distal end of an otherwise normal 9S (Han et al. 2006). This dicentric chromosome, 9-B inactivated centromere-1 (9-Bic-1), is stable in mitosis and meiosis because the B centromere was inactivated upon transposition (Han et al. 2006). In the presence of a B chromosome, which supplies the trans-acting factors required for nondisjunction, 9-Bic-1 may attempt to undergo nondisjunction (Han et al. 2007). This finding shows that centromere activity is not needed for nondisjunction. The inactive centromere remains adhered together at the second pollen mitosis in this case and thus its short arm occasionally breaks and releases the inactive B centromere when the chromosome 9 centromeres proceed toward opposite poles (Han et al. 2006, 2007).

We were able to use this property of 9-Bic-1 in conjunction with C1, a dominant allele of the c1 gene required for anthocyanin pigmentation in the embryo and endosperm of the kernel, to identify plants
possessing 9-Bic-1 that had undergone chromosome breakage. Plants containing one copy each of 9-Bic-1 and chromosome 9, each containing C1, were crossed to a c1 tester line. If 9-Bic-1 breaks distal to C1, the chromatid-type breakage-fusion-bridge (BFB) cycle will occur, and a mosaic color pattern will be seen in the endosperm (Fig. 1). If 9-Bic-1 breaks proximal to C1, the resulting BFB cycle will produce a colorless endosperm and colored embryo (Han et al. 2007). The endosperm and embryo will be fully colored if no breakage occurs. The loss or mosaic phenotype of C1 in the endosperm would be associated with embryos that receive the other sperm containing the broken fragment.

Here, we attempted to reactivate the inactive B centromere on 9-Bic-1 by utilizing nondisjunction afforded by the presence of B chromosomes. We identified two heritable mini chromosomes containing released B centromeres. One of the mini chromosome arose from 9-Bic-1 chromosomal breakage, and the second mini chromosome originated via B chromosome breakage. Chromatin immunoprecipitation-sequencing (ChIP-seq) showed that the released centromere in 9-Bic-1 derived mini chromosome regained association with detectable CENH3 together with the establishment of a de novo centromere. For the second mini chromosome, the B centromere originating from the B chromosome lost function, but a de novo centromere arose elsewhere on the chromosome.

**Materials and methods**

**Plant material**

All lines used in this study were in the B73 inbred background. Seedlings were grown in the Ernie and Lottie Sears Plant Growth Facility at the University of Missouri-Columbia under the following conditions: 16 h light, 25 C day/20 C night.

**FISH and immunolocalization**

Fluorescence in situ hybridization (FISH) and immunolocalization in mitosis and meiosis were performed as described (Kato et al. 2004; Han...
et al. 2009; Gaeta et al. 2011). A combination of 5′-labeled synthetic oligonucleotides and nick-translated probes were used. CentC was labeled with 6-FAM (fluorescein; green); telomeric sequences, which hybridize to ZmBs (Alfenito and Birchler 1993; Masonbrink et al. 2012), were labeled with 6-FAM or Texas Red. The signal from the B centromere at the exposure used is readily visible, while the telomeres are not. ZmBs was labeled with Texas Red. Individual loci from 9S were amplified from B73 genomic DNA (see Supplemental Table 2) and labeled with Texas Red-dCTP by nick translation (Kato et al., 2004). Chromosomes were counterstained with 4′,6-diamidino-2-phenylindole (DAPI) and mounted with Vectashield (Vector Laboratories). Images were captured on an Olympus BX61 microscope using Applied Spectral Imaging Software V.4.5-SP3 and a COOL-1300QS digital camera.

Copy number variation analysis

CNV analyses were performed as described previously (Yang et al. 2021). In brief, genomic DNA sequencing data were trimmed using cutadapt (Martin 2011) and filtered with FASTX toolkit (Hannon 2010) using parameter -Q33 -q 20 -p 80. Then, each fastq file was aligned to B73 reference v4 (Jiao et al. 2017) plus mitochondria and chloroplast genomes using Bowtie2 (Langmead and Salzberg 2012) under parameter –phred33 -N 0 –no-unal -k 10. The SAM file was further filtered to keep 0 mismatch reads using a perl script. Only unique mapped reads in the SAM file were kept using a perl script. The read number within each gene region was counted for each mini B, 9-Bic-1 or B73. Ratios were calculated by comparing mini B or 9-Bic-1 to B73 and plotted along each chromosome using R.

B chromosome deficiency mapping

Deficiency mapping to mini chromosomes was performed as described previously (Blavet et al. 2021). In brief, low-quality read filtering and mapping procedures are the same as in the copy number variation analysis. The uniquely mapped reads that have 0 mismatches in the SAM file were regarded as aligned reads on the A chromosomes and were filtered out from the fastq (after filtering low-quality reads) file using a perl script to obtain the unmapped reads. Each fastq file that consists of unmapped reads was aligned to the B chromosome (Blavet et al. 2021) and only reads that mapped to a unique location were kept. The reads were counted in the regions of 1 kb along each sequence and the results were plotted by ggplot2 in R.

ChIP-seq and data analysis

CENH3 ChIP-seq was performed as described (Fu et al. 2013) using ~5.0–7.5 g of 14-day whole seedling tissue. ChIP DNA was sequenced with paired-end 100 nt Illumina. Reads were trimmed and cleaned with TRIMMOMATIC (Bolger et al., 2014) (LEADING:30 TRAILING:29 SLIDINGWINDOW:4:30 MINLEN:29 AVGQUAL:28) prior to mapping.

For testing CENH3 peaks on A chromosomes (from chromosomes 1 to 10), CENH3 ChIP-seq reads were mapped to the B73 version 4 plus B chromosome using bowtie2 with default parameters. Reads were further filtered to 0 mismatches and unique mapped reads using perl scripts. The alignments were converted to BAM files, then sorted and merged using SAMtools (Li et al. 2009). Read coverage and enrichment were displayed after converting BAM files to TDF files using igvtools (Thorvaldsdóttir et al. 2013).

For CENH3 binding on the B centromere region, we first adopted the approach of B chromosome deficiency mapping to identify B chromosome specific CENH3 ChIP-seq reads. The reads were aligned to the B73 genome with nondefault parameters: “–phred33 -N 0 –no-unal -k 10”; the reads recorded in the SAM file were treated as mapped reads on the A genome and thus filtered out of the original FASTQ file using a perl script and the remaining reads were aligned to the maize B genome (chrB + 307 B scaffolds) using Bowtie2 with nondefault parameters: “–phred33 -N 0 –no-unal.” Only reads that mapped to a unique location on the B genome were kept. The alignments were converted to BAM files, then sorted and merged using SAMtools. Read coverage and enrichment were displayed after converting BAM files to TDF files using igvtools.
Results

Identification of mini chromosomes potentially derived from 9-Bic-1

Approximately 10,000 kernels were screened for breakage of 9-Bic-1, of which 1412 kernels exhibited signs of 9-Bic-1 breakage using the C1-based system described above. When 9-Bic-1 is broken, there are four possible fates for the resulting fragment: (i) the B centromere remains inactive, in which case the fragment will be lost (Fig. 2A); (ii) the fragment translocates onto another chromosome that contains an active centromere (Fig. 2B); (iii) centromere mis-division of chromosome 9 centromere of 9-Bic-1 forms an isochromosome that contains an inactive B centromere on both ends (Fig. 2C); or (iv) a broken fragment from either breakage of 9-Bic-1 or the B chromosome becomes heritable (Fig. 2D). FISH was used to examine each of the 1412 kernels that showed signs of chromosome breakage. In the vast majority of cases \( n = 1391 \), no fragment was observed, presumably due to loss of the fragment. However, isochromosomes were observed ten times, a single translocation of the inactive B centromere was observed, and ten mini chromosomes containing a B centromere were recovered (Supplemental Table 1; Supplemental Fig. 1; Blavet et al. 2021).

The B sequences on mini B1104 and mini B496 are different

Previous research determined the breakpoint of six out of ten mini chromosomes from this screening on the B chromosome by using B chromosome deficiency mapping (Blavet et al. 2021). In that study, mini B876 was found to be broken in the B centromere region, which indicates it could be derived from B centromere misdivision that occurs at a low frequency (Blavet et al. 2021; Carlson and Chou 1981; Kaszás and Birchler 1996; 1998). However, its breakpoint in the centromere could alternatively indicate that it results from 9-Bic-1 breakage followed by reactivation. Five other mini chromosomes showed breakpoints in the proximal knob region likely due to B chromosome breakage (Blavet et al. 2021), given that these sequences are not present in 9-Bic-1. Mini B524 and mini B1139 were not subjected to this analysis because mini B524 material could not be further perpetuated and B chromosomes still exist in mini B1139 material, which makes it unsuitable for deficiency mapping. The results showed that mini B1104 has the same breakpoints as 9-Bic-1 in the centromere and chromosome arm 9S with its B portion broken at the centromere region adjacent to the centromeric knob, which suggest that mini B1104 was a 9-Bic-1 derivative (Fig. 3, Supplemental Fig. 2). The mini B496 might have been derived from an intact B chromosome as it is broken at the B centromeric knob region, which is distinct from 9-Bic-1 (Fig. 3, Supplemental Fig. 2).

Both mini B1104 and mini B496 contain a 9S segment

Previous results suggest that 9-Bic-1 lost its chromosome 9 short arm tip because homozygous
9-Bic-1 exhibits an albino phenotype (Han et al. 2007). Plants homozygous for 9-Bic-1 were used to roughly map the translocation point of the inactive B centromere between 1.85 and 3.44 Mb on 9S (Supplemental Fig. 3). To determine the translocation point of 9-Bic-1 accurately, we used genomic DNA sequencing data of a 9-Bic-1 heterozygote and B73 to perform copy number variation (CNV) analysis for chromosome 9. We found the first three Mb on chromosome 9 was approximately at a ratio of 0.5

Fig. 3 Breakpoints of mini B1104 and mini B496 on B chromosome. Eight mini B chromosomes and 9-Bic-1 were subjected to B chromosome deficiency mapping as previously performed (Blavet et al 2021). B73 without B chromosomes was used as a negative control (C_B73). Sequence reads unique to the B chromosome were aligned to the B chromosome reference sequence. The Y axis is the aligned read numbers of log10 of 1 kb regions on the B chromosome. Each black dot equals 1 kb resolution. The X axis is the position along the B reference genome. Mini B1104 has the same breakpoint as 9-Bic-1 at 2.1 Mb, which is marked by the magenta arrow. The mini B496 is broken at 7.5 Mb, which suggests it is not a 9-Bic-1 derivative. The background signals in B73 and the minichromosomes result from cross homology of knob and CentC from the A chromosomes to the B reference (Blavet et al., 2021). C, control. D, deficiency

Fig. 4 Translocation point of 9-Bic-1 and mini chromosomes on chromosome 9. Copy number variation along chromosome 9 in 9-Bic-1, mini B1104, and mini B496. Y axis is log2 of the ratio. X axis is the ratio along chromosome 9. Each red dot is the ratio comparing read numbers of 9-Bic-1, mini B1104, or mini B496 to B73 within a single gene window. The diagram below each plot shows the chromosome constitution for each group. Red oval represents the inactive B centromere. The centromere of chromosome 9 is depicted as a green circle. The 9-Bic-1 heterozygote contains one 9-Bic-1 and one intact chromosome 9. Both mini B1104 and mini B496 contain one mini chromosome and two chromosomes 9. A The first three Mb (box in A) showed lower copy number suggesting the breakpoint of 9-Bic-1 is at ~3 Mb. B A ~500 kb region (box in B) immediately after the breakpoint of 9-Bic-1 was found to have a higher ratio in mini B1104 suggesting it is a 9-Bic-1 derivative. C A ~200 kb region within the ninth Mb in chromosome 9 was found to have a higher ratio in mini B496 (box)
but the ratio reached 1 from the fourth Mb through the remainder of chromosome 9, which suggests that the breakpoint is located between the 3rd and 4th Mb on chromosome 9. Finally, the breakpoint of 9-Bic-1 was further localized between Zm00001d044806 and Zm00001d044808 (2.94 ~ 3.06 Mb) (Fig. 4).

Next, we checked the CNV of mini B1104 and mini B496 on chromosome 9. The chromosome portion between Zm00001d044808 and Zm00001d044815 (3.06 ~ 3.52 Mb) had a higher ratio compared to the remainder of the region when comparing mini B1104 and B73, which suggests that this ~ 500 kb 9S segment is on mini B1104 (Fig. 4). This result also supports the conclusion that the mini B1104 was directly derived from 9-Bic-1.

Although the centromere of mini B496 likely has an origin from the normal B chromosome, it also contains an interstitial 9S region between Zm00001d044995 and Zm00001d044999 (9.51 ~ 9.67 Mb) because this portion of the chromosome has higher CNV when comparing mini B496 to B73 (Fig. 4). FISH with a probe specific to this region confirmed the presence of a 9S segment in mini B496 (Supplemental Fig. 4). These results suggest that mini B496 might have fused with a broken 9-Bic-1, although its precise history can only be hypothesized. The CNV analysis of the other mini chromosomes indicates that they contain no 9S portion (Supplemental Fig. 5).

mini B1104 and mini B496 have functional centromere activity

The two mini chromosomes, mini B496 and mini B1104, have been inherited through at least seven generations, which indicates that they contain a functional centromere. Immunolocalization showed that known epigenetic marks of functional maize centromeres, such as CENH3 (Zhong et al. 2002), H3-Ser10ph (Gao et al. 2011), and H2A-Thr133ph (Dong and Han 2012), are present on both mini B496 and mini B1104 (Fig. 5). Both mini chromosomes also exhibit typical mini chromosome behavior during meiosis (Supplemental Fig. 6; Birchler and Han 2013); they do not pair during pachytene when present in multiple copies (Supplemental Fig. 6A and B), and sister chromatids prematurely separate and lag in anaphase I (Supplemental Fig. 6E and F).

Examination of mini chromosomes reveals de novo centromere formation

In order to determine the DNA sequences associated with the functional centromeres of mini B496 and mini1104, ChIP-seq was performed with a maize anti-CENH3 antibody on three biological replicates each of the inbred B73, B73 with B chromosomes (B73+B), and lines containing TB-9Sb, 9-Bic-1, and mini B496. Five biological replicates, two from generation 6 and three from generation 7, were subjected to ChIP for line mini B1104. The B73 +B and TB-9Sb lines have a functional B centromere, which can be used as positive controls to test CENH3 binding on the B centromere, while 9-Bic-1, which contains an inactive B centromere, and B73 without B chromosomes can serve as negative controls.

The B chromosome sequencing project has identified 575 kb of B chromosome centromeric sequences consisting of six B scaffolds and the 250 kb proximal portion of the short arm of the assembled B reference, which includes a portion of the centromere (Blavet et al. 2021). Because the two mini chromosomes contain intact B centromeric regions, we first checked the CENH3 occupation on these sequences. To this end, the CENH3 CHIP-seq reads were first mapped to B73. The reads that aligned to the B73 genome were filtered and the remaining reads were mapped to the maize B segment. This step is performed to reduce mapping ambiguity due to homology of centromeric repeats between the B chromosome and A chromosomes, such as CentC and CRM (Lamb et al. 2005; Jin et al. 2005). When the CENH3 read pileup was compared across all lines in the 575 kb B centromeric sequences (Blavet et al. 2021), it was found that 9-Bic-1 and mini B496 had no or markedly reduced CENH3 binding compared to TB-9Sb and B73+B that contain active B centromeres (Fig. 6). In contrast, the B1104 centromere sequences showed an association with CENH3 to some degree. These results suggest that reactivation of canonical centromeric chromatin occurred in mini B1104, while for mini B496, whose centromere is derived from the B chromosome, the B centromere became inactive.

To determine whether a region other than the canonical centromeric repeats acquired centromere activity, CENH3 enrichment on the A chromosomes was analyzed. The CENH3 binding regions, denoting the centromeres of chromosomes 1 through 10,
were virtually indistinguishable amongst the four lines studied (Supplemental Fig. 7). However, lines mini B496 and mini B1104 displayed CENH3 binding peaks on 9S that were absent in all of the other lines (Fig. 7A). In mini B1104, a ~300 kb 9S region (from 3.1 to 3.35 Mb) displayed unique CENH3 binding (Fig. 7B). In contrast, for mini B496, the unique CENH3 binding domain stretched for ~160 kb from 9.51 to 9.67 Mb on 9S, which is present on this mini chromosome (Figs. 4 and 7C, Supplemental Fig. 4). Taken together, the CENH3 ChIP-seq results strongly suggest that both mini chromosomes contained de novo formed centromeres.

Discussion

Both mini B1104 and mini B496 have de novo centromere formation

Cases of centromere formation in the last two decades show the commonality of de novo centromere formation. Adding alien chromosomes to wheat or oat resulted in chromosomal mutations such that these alien chromosomes lacked the canonical centromeric sequences but still showed normal transmission, which suggested they had de novo centromeres (Nasuda et al. 2005; Topp et al. 2009). Also, many cases of de novo centromeres have been documented in maize (Fu et al. 2013; Liu
et al. 2015, 2020; Schneider et al. 2016). Our data suggest that a 300 kb region of 9S on mini B1104 gained CENH3 binding. During the formation of mini B496, the active B centromere lost its function but newly active centromeric chromatin formed on non-canonical centromeric sequences of this mini chromosome.

Centromere reactivation occurred on mini B1104

Previous results documented a small, inactivated B chromosome centromere reactivated after being recovered from an unstable dicentric (Han et al. 2009). However, the result could not discriminate whether exactly the same sequences were associated with function in the original active and reactivated centromeres. Our result suggesting CENH3 binding on the B centromere sequences in mini B1104 indicates the same sequences that were previously inactive can regain CENH3 association. Given the small size of mini B1104, it is not possible to determine whether the B centromere, the de novo CENH3 associated sequences, or both, can organize a kinetochore.

Fig. 6 CENH3 binding on B centromere. CENH3 ChIP-seq reads from six samples were mapped to seven B centromeric sequences to determine CENH3 binding. The location of the three most common repeats including ZmBs, CRM2, and CentC are displayed in each scaffold. B73+B and TB-9Sb each contain an active B centromere, while no B centromere is present in B73. 9-Bic-1 comprises inactive B centromeric chromatin. In B73 plus B and TB9-Sb, CENH3 enrichment was found in each scaffold. For the two mini chromosomes, mini B1104 and mini B496, the CENH3 occupation was found to be remarkably reduced in mini B1104 and almost absent in mini B496. Three kinds of B centromeric repeat, CentC, CRM2, and ZmBs were blasted to each sequence to locate each repeat type in the B centromeric scaffolds.

Mini B496 is a secondary derivative of 9-Bic-1

Based on our analysis, mini B496 is probably derived from a normal B chromosome breakage but it also
contains an interstitial fragment from chromosome 9S. It has lost centromere activity on the canonical B centromeric repeats but acquired a de novo formation of a centromere, which allowed it to be inherited. The formation of mini B496 cannot be known with certainty, but might have occurred as follows: (i) a mini B was produced from breakage that happened on a normal B chromosomes during attempted non-disjunction at the second pollen mitosis and its B centromere remained active; (ii) within the same sperm was a fractured chromosome in 9S from an attempted nondisjunction of 9-Bic-1; (iii) the mini B and broken 9-Bic-1 chromosome fused together to form a dicentric chromosome with centromere 9; (iv) the dicentric chromosome fractured on 9S and the B centromeric region of mini B496 lost its function, but de novo centromere formation occurred on the 9S portion (Fig. 8). Previous results have documented the rapid establishment of de novo centromeres in maize (Liu et al. 2020).

**B chromosome breakage during the second pollen mitosis**

During the second pollen mitosis, chromosome breakage can occur during non-disjunction of the B
De novo centromere formation on chromosome fragments with an inactive centromere in maize (Zea…

chromosome, and such breakage could result in the formation of mini B chromosomes. The broken end of a mini B chromosome will be repaired and stabilized by adding a telomere to its end in the embryo after the sperm with mini B chromosome joins with the egg cell (McClintock 1939; McKnight and Shippen 2004). The inactive B centromere on 9-Bic-1 remains adhered together when normal B chromosomes are present at the second pollen mitosis (Han et al. 2007). Chromosome breakage of 9-Bic-1 is indicated when a mosaic endosperm occurs, but this process does not monitor chromosome breakage on the normal B chromosome.

In this study, 10 mini chromosomes with B repeat signal (as observed using FISH) were recovered from a total of 1412 kernels, which exhibited signs of chromosome breakage in the endosperm. From these 10 mini chromosomes, two of them were not subjected to further breakpoint analysis on the B chromosome due to either a failure to perpetuate the line or because a B chromosome was still present with the mini chromosome, which obscures the ability to perform a comparison with the B reference sequence. From the analysis of the remaining eight mini chromosomes, it is ambiguous as to whether mini B876 was derived from a B centromere mis-division or a similar break in 9-Bic-1 followed by reactivation because its breakpoint was found in the B centromere region.

However, mini B1104 has been definitively determined to be derived from 9-Bic-1. The other six mini chromosomes have breakpoints in the centromeric knob region, which suggests that these mini chromosomes are likely from normal B chromosome breakage. In this case, the percentage of B chromosome breakage is at least 0.42% (6/1412) but potentially an underestimate due to the inability to recognize all cases. This result suggests that the normal B chromosome centromere can be attached to the spindle from both poles at the second pollen mitosis at least in some cases, which can fracture the centromere or the adjacent knob, which is surrounded by the B specific repeat that is involved with nondisjunction (Lamb et al. 2005; Blavet et al. 2021) if the B specific repeat region remains adhered between sister chromatids.

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Author contribution J.B., F.H., R.D., H.Y., designed the research; R.D., H.Y., B.Z., performed research; H.Y., C.C., T.J., J.C. contributed analytical/computational tools; H.Y., R.D., C.C., T.J., J.B. analyzed data; R.D., H.Y., J.B. wrote the paper. All authors read, discussed, and approved the final version of the manuscript.

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Data availability Raw data of B73, 9-Bic-1 and six lines with mini B chromosome used for B chromosome deficiency mapping or CNV analysis were downloaded from NCBI-SRA as BioProject PRJNA634743. Genomic sequencing data of mini B496 and B1104 CENH3 and ChIP-seq data are available in NCBI-SRA as BioProject PRJNA736453.

Declarations

Conflict of interest The authors declare no competing interests.

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References

Alfenito MR, Birchler JA (1993) Molecular characterization of a maize B-chromosome centric sequence. Genetics 135:589–597
Ananiev EV, Phillips RL, Rines HW (1998) Chromosome-specific molecular organization of maize (Zea mays L) centromeric regions. Proc Natl Acad Sci USA 95:13073–13078
Birchler JA, Han F (2013) Meiotic behavior or small chromosomes in maize. Front Plant Sci 4:505
Blavet N, Yang H, Su H, Solansky P, Douglas RN, Karafiatova M, Simkova L, Zhang J, Liu Y, Hou J, Shi X, Chen C, El-Walid M, McCaw ME, Albert PS, Gao Zhao C, Ben-Zvi G, Glick L, Kol G, Shi J, Jrana J, Simkova H, Lamb JC, Newton KJ, Dawe RK, Dolezel J, Ji T, Baruch K, Cheng J, Han F, Birchler JA, Bartos J (2021) Sequence of the supernumerary B chromosome of maize provides insight

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into its drive mechanism and evolution. Proc Natl Acad Sci USA 118:e2104254118
Bodor DL, Mata JF, Sergeev M, David AF, Salimian KJ, Panchenko T, Cleveland DW, Black BE, Shah JV, Jansen LE (2014) The quantitative architecture of centromeric chromatin. Elife 3:e02137
Bolger AM, Lohse M, Usadel B (2014) Trimmomatic: a flexible trimmer for Illumina Sequencing Data. Bioinformatics, 31:17
Bures P, Zedek F, Markova M (2012) Holocentric chromosomes. Plant Genome Diversity 2:187–208
Burrack LS, Berman J (2012) Flexibility of centromere and kinetochore structures. Trends Genet 28:204–212
Carlson WR, Chou T-S (1981) B chromosome nondisjunction in corn: control by factors near the centromere. Genetics 97:379–389
Dong Q, Han F (2012) Phosphorylation of histone H2A is associated with centromere function and maintenance in meiosis. Plant J 71:800–809
Fu S, Lv Z, Gao Z, Wu H, Pang J, Zhang B, Dong Q, Guo X, Wang X-J, Bircher JA, Han F (2013) De novo centromere formation on a chromosome fragment in maize. Proc Natl Acad Sci USA 110:6033–6036
Gaeta RT, Danilova T, Zhao C, Masonbrink RE, McCaw ME, Bircher JA (2011) Recovery of a telomere-truncated chromosome via a compensating translocation in maize. Genome 54:184–195
Gao Z, Fu S, Dong Q, Han F, Bircher JA (2011) Inactivation of a centromere during the formation of a translocation in maize. Chromosome Res 19:6033–6036
Han F, Gao Z, Bircher JA (2009) Reactivation of an inactive centromere reveals epigenetic and structural components for centromere specification in maize. Plant Cell 21:1929–1939
Han F, Lamb JC, Bircher JA (2006) High frequency of centromere inactivation resulting in stable dicentric chromosomes of maize. Proc Natl Acad Sci USA 103:3238–3243
Han F, Lamb JC, Yu W, Gao Z, Bircher JA (2007) Centromere function and nondisjunction are independent components of the maize B chromosome accumulation mechanism. Plant Cell 19:524–533
Hannon GJ (2010) FASTX-Toolkit. http://hannonlab.cshl.edu/fastx_toolkit
Hartley G, O’Neill RJ (2019) Centromere repeats: hidden gems of the genome. Genes (basel) 10:223
Houben A, Demidov D, Caperta AD, Karimi R, Augeci F, Vlasenko L (2007a) Phosphorylation of histone H3 in plants – A dynamic affair. Biochim Biophys Acta 1769:308–315
Houben A, Schroeder-Reiter E, Nagaki K, Nasuda S, Wanner G, Murata M, Endo TR, Endo TR (2007b) CENH3 interacts with the centromeric retrotransposon cereba and GC-rich satellites and locates to centromeric substructures in barley. Chromosoma 116:275–283
Jiang J, Nasuda S, Dong F, Scherrer CW, Woo S, Wing RA, Gill BS, Ward DC (1996) A conserved repetitive DNA element located in the centromeres of cereal chromosomes. Proc Natl Acad Sci USA 93:14210–14212
Jiao Y, Peluso P, Shi J, Liang T, Sitzer MC, Wang B, Campbell MS, Stein JC, Wei X, Chin CS et al (2017) Improved maize reference genome with single-molecule technologies. Nature 546:524–527
Jin W, Lamb JC, Vega JM, Dawe RK, Bircher JA, Jiang J (2005) Molecular and functional dissection of the maize B chromosome centromere. Plant Cell 17:1412–1423
Joglekar A, Bouck D, Finley K, Liu X, Wan Y, Berman J, He X, Salmon E, Bloom K (2008) Molecular architecture of the kinetochore-microtubule attachment site is conserved between point and regional centromeres. J Cell Biol 181:587–594
Johnston K, Joglekar A, Horii T, Suzuki A, Fukagawa T, Salmon ED (2010) Vertebrate kinetochore protein architecture: Protein copy number. J Cell Biol 189:937–943
Kaszás E, Bircher JA (1996) Misdivision analysis of centromere structure in maize. EMBO J 15:5246–5255
Kaszás E, Bircher JA (1998) Meiotic transmission rates correlate with physical features of rearranged centromeres in maize. Genetics 150:1683–1692
Kato A, Lamb JC, Bircher JA (2004) Chromosome painting using repetitive DNA sequences as probes for somatic chromosome identification in maize. Proc Natl Acad Sci USA 101:13554–13559
Lamb JC, Han F, Auger DL, Bircher JA (2006) A trans-acting factor required for non-disjunction of the B chromosome is located distal to the TB-4Lb breakpoint on the B chromosome. Maize Genetics Cooperation Newsletter 80:51–54
Lamb JC, Kato A, Bircher JA (2005) Sequences associated with A chromosome centromeres are present throughout the maize B chromosome. Chromosoma 113:337–349
Langmead B, Salzberg S (2012) Fast gapped-read alignment with Bowtie 2. Nat Methods 9:357–359
Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R (2009) The sequence alignment/map format and SAMtools. Bioinformatics 25:2078–2079
Lin BY (1978) Regional control of nondisjunction of the B chromosome in maize. Genetics 90:613–627
Liu Y, Su H, Pang J, Gao Z, Wang X-J, Bircher JA, Han F (2015) Sequential de novo centromere formation and inactivation on a chromosomal fragment in maize. Proc Natl Acad Sci USA 112:E1263-1271
Liu Y, Su H, Zhang J, Shi L, Liu Y, Zhang B, Bai H, Liang S, Gao Z, Bircher JA, Han F (2020) Rapid birth and death of centromeres on fragmented chromosomes in maize. Plant Cell 32:3113–3123
Manzanero S, Arana P, Puertas MJ, Houben A (2000) The chromosomal distribution of phosphorylated histone H3 varies between plants and animals at meiosis. Chromosoma 109:308–317
Martin M (2011) Cutadapt removes adapter sequences from high-throughput sequencing reads. Embnet J 17:10–12
Masonbrink R, Gaeta R, Bircher J (2012) Multiple maize minichromosomes in meiosis. Chromosome Res 20:395–402
McClintock B (1939) The behavior in successive nuclear division of the genome. Genes (basel) 10:223
McKnight TD, Shippen DE (2004) Plant telomere biology. Plant Physiology 134:1925–1933
McKnight TD, Shippen DE (2004) Plant telomere biology. Plant Cell 16:794–803
Nasuda S, Hudakova S, Schubert I, Houben A, Endo TR (2005) Stable barley chromosomes without centromeric repeats. Proc Natl Acad Sci USA 102:9842–9847

Robertson DS (1984) Different frequency in the recovery of crossover products from male and female gametes of plants hypoploid for B-A translocations in maize. Genetics 70:117–130

Roman HL (1947) Mitotic nondisjunction in the case of interchanges involving the B-type chromosome in maize. Genetics 32:391–409

Roman HL (1948) Directed fertilization in maize. Proc Natl Acad Sci USA 34:36–42

Schneider KL, Xie Z, Wolfgruber TK, Presting GG (2016) Inbreeding drives maize centromere evolution. Proc Natl Acad Sci USA 113:E987-996

Staub RW (1987) Leaf striping correlated with the presence of B chromosomes in maize. J Hered 78:71–74

Thorvaldsdóttir H, Robinson JT, Mesirov JP (2013) Integrative Genomics Viewer (IGV): high-performance genomics data visualization and exploration. Brief Bioinform 14:178–192

Topp CN, Okagaki RJ, Melo JR, Kymast RG, Phillips RL, Dawe RK (2009) Identification of a maize neocentromere in an Oat-Maize Addition line. Cytogenetics and Genome Research 124:228–238

Wang N, Dawe RK (2018) Centromere size and its relationship to haploid formation in plants. Mol Plant 11:398–406

Wolfgruber TK, Sharma A, Schneider KL, Albert PS, Koo D, Shi J, Gao Z, Han F, Lee H, Xu R, Allison J, Birchler JA, Jiang J, Dawe RK, Presting GG (2009) Maize Centromere Structure and Evolution: Sequence Analysis of Centromeres 2 and 5 Reveals Dynamic Loci Shaped Primarily by Retrotransposons. PLoS Genet 5:e1000743

Yang H, Shi X, Chen C, Hou J, Ji T, Cheng J, Birchler JA (2021) Predominantly inverse modulation of gene expression in genomically unbalanced disomic haploid maize. The Plant Cell https://doi.org/10.1093/plcell/koab029

Zhang W, Friebe B, Gill BS, Jiang J (2010) Centromere inactivation and epigenetic modifications of a plant chromosome with three functional centromeres. Chromosoma 119:553–563

Zhong CX, Marshall JB, Topp C, Mroczek R, Kato A, Nagaki K, Birchler JA, Jiang J, Dawe RK (2002) Centromeric retroelements and satellites interact with maize kinetochore CENH3. Plant Cell 14:2825–2836

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