Nonrandom Distribution of Interhomolog Recombination Events Induced by Breakage of a Dicentric Chromosome in Saccharomyces cerevisiae

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ABSTRACT Dicentric chromosomes undergo breakage in mitosis, resulting in chromosome deletions, duplications, and translocations. In this study, we map chromosome break sites of dicentrics in Saccharomyces cerevisiae by a mitotic recombination assay. The assay uses a diploid strain in which one homolog has a conditional centromere in addition to a wild-type centromere, and the other homolog has only the wild-type centromere; the conditional centromere is inactive when cells are grown in galactose and is activated when the cells are switched to glucose. In addition, the two homologs are distinguishable by multiple single-nucleotide polymorphisms (SNPs). Under conditions in which the conditional centromere is activated, the functionally dicentric chromosome undergoes double-stranded DNA breaks (DSBs) that can be repaired by mitotic recombination with the homolog. Such recombination events often lead to loss of heterozygosity (LOH) of SNPs that are centromere distal to the crossover. Using a PCR-based assay, we determined the position of LOH in multiple independent recombination events to a resolution of ~4 kb. This analysis shows that dicentric chromosomes have recombination breakpoints that are broadly distributed between the two centromeres, although there is a clustering of breakpoints within 10 kb of the conditional centromere.

The cells of solid tumors often have numerous chromosome alterations, both changes in chromosome number and structural alterations including deletions, duplications, and translocations (Cimini 2008). One mechanism that can contribute to the formation of chromosome alterations is the formation of dicentric chromosomes, followed by the breakage of the dicentric leading to secondary chromosome alterations (Stimpson et al. 2012).

In Saccharomyces cerevisiae, as well as in other eukaryotes, dicentric chromosomes can be formed in several ways. In yeast strains with certain telomeric defects, fusions between telomeres of different chromosomes or fusions between the telomere of one chromosome and an internal break on another chromosome can produce dicentric chromosomes (Myung et al. 2001; Craven et al. 2002; Mieczkowski et al. 2003; Pardo and Marcand 2005). Dicentrics also arise as a consequence of ectopic homologous recombination between retrotransposons on nonhomologous chromosomes (Mieczkowski et al. 2006) or as a consequence of nonhomologous end joining between two broken chromosomes (Myung et al. 2001; Craven et al. 2002). Intrachromosomal dicentrics can also be generated by processing or replication of inverted repeats (Lobachev et al., 2002; Narayanan et al. 2006; VanHulle et al. 2007). Lastly, dicentric plasmids and chromosomes have been generated by in vitro manipulations, followed by transformation of the resulting constructions into yeast (Mann and Davis 1983; Koshland et al. 1987).

Regardless of the mechanism by which they are formed, dicentric plasmids and chromosomes are unstable in S. cerevisiae, undergoing various types of structural rearrangements to generate monocentric plasmids or chromosomes (Mann and Davis 1983; Koshland et al. 1987; Hill and Bloom 1987).
1989; Kramer et al. 1994; Narayanan et al. 2006; Pennaneach and Kolodner 2009). The instability is initiated when the two centromeres of a dicentric chromosome are pulled to different daughter cells during anaphase, resulting in the stretching of chromosomal sequences located between the two centromeres (Thrower and Bloom 2001). Although the subsequent double-stranded DNA breaks (DSBs) could simply reflect the mechanical forces applied by the spindle to the dicentric chromosome, breakage of the dicentric is observed when the chromosome experiences forces of ~1 piconewton (pN) (Fisher et al. 2009), much less than the force required to mechanically break a double-stranded DNA molecule (~480 pN; Bensimon et al. 1995). Based on these considerations and the observation that stretching and breakage of dicentric yeast chromosomes occur in anaphase (Thrower and Bloom 2001), it is likely that dicentric breakage requires the enzymatic production of DSBs in stretched chromatin. The enzymes involved in dicentric breakage, presumably endonucleases, have not yet been identified. Alternatively, or in addition, dicentric breakage could occur during nuclear fission or cytokinesis (Quevedo et al. 2012). For dicentric yeast chromosomes formed by telomere–telomere fusions, breakage often occurs at the telomere fusion junction, possibly involving resolution by enzymes that process cruciforms (Pobiega and Marcand 2010). Finally, it should be pointed out that the instability of dicentric chromosomes is not universal, since some dicentrics in human cells are relatively stable (Stimpson et al. 2012).

Following dicentric breakage, a stable karyotype in yeast requires the resulting DSBs to be repaired to generate telomere-capped monocentric chromosomes. There are a variety of mechanisms that can produce these products, depending on whether the strain is a haploid or diploid, the availability of repetitive sequences (for example, transposons) located near the break, the location of essential genes near the break, and other factors. One of the simplest mechanisms for repairing broken chromosome ends is by telomere addition (Haber and Thorburn 1984; Pennaneach and Kolodner 2009). Another common mechanism of generating a monocentric from a dicentric is a deletion that removes one of the two centromeres. Such deletions can occur by intrachromosomal homologous recombination between repeated genes that flank one of the two centromeres (Brock and Bloom 1994; Lemoine et al. 2005) or by non-homologous end joining (NHEJ) following processing of the broken DNA ends (Kramer et al. 1994).

Alternatively, the repair process may involve recombination with a homolog or a nonhomologous chromosome. For example, a DSB resulting from breakage of a dicentric in a diploid strain could be repaired by homologous recombination with a homolog (Haber and Thorburn 1984), either by a crossover (Figure 1A) or by a break-induced replication (BIR) event (Figure 1B). If a broken chromosome contains a repetitive element (such as the retrotransposon Ty), it can also acquire a telomere by homologous recombination with a repetitive element on a nonhomologous chromosome; such events often involve BIRs (Umézu et al. 2002; Lemoine et al. 2005; Argueso et al. 2008; Vernon et al. 2008; Pennaneach and Kolodner 2009; Chan and Kolodner 2011). Alternatively, the broken chromosomes can be joined to the broken ends of nonhomologous chromosomes or to telomeres of other chromosomes by NHEJ, although homologous recombination events involving repeats are more common (Pennaneach and Kolodner 2009). Finally, it should be noted that repair of the broken chromosome may regenerate a dicentric chromosome, leading to additional chromosome rearrangements before a stable karyotype is generated (Admire et al. 2006; Narayanan et al. 2006; Pennaneach and Kolodner 2009).

Most studies of the behavior of dicentric yeast chromosomes focus on the chromosome aberrations generated during the repair of chromosome breaks. Since these aberrations are usually a consequence of homologous recombination between nonallelic Ty elements (as described above), these studies select for dicentric breaks that occur within a Ty element or breaks that occur near a Ty element that are processed to generate a Ty-containing broken end (Hoang et al. 2010). In our study, we examine the location of dicentric chromosome breaks by mapping homologous recombination events between a dicentric chromosome with one conditional centromere and a homologous monocentric chromosome. Our analysis shows that dicentric chromosomes break at sites distributed throughout the region between the two centromeres, but the region near the conditional centromere is particularly prone to breakage.

Materials and Methods

Genetic analysis and media

Standard yeast procedures were used for mating, sporulation, and tetrad dissection (Guthrie and Fink 1991). Rich growth medium, yeast extract, peptone, and dextrose (YPD), and omission media were made following standard protocols (Guthrie and Fink 1991; Lee et al. 2009). The growth media YPGal was identical to YPD, except 2% galactose was substituted for glucose. To select canavanine-resistant transformants, we used solid omission medium lacking arginine (SD −Arg) containing 120 μg/ml of canavanine.

Strains

The genotypes of all strains in this study are given in Supporting Information, Table S1. Three diploids were constructed (details in File S1). All diploids were isogenic except for the location of the heterozygous conditional centromere. The locations of the conditional centromeres in the three strains were on chromosome III near HIS4 (WS49), on chromosome V near CAN1 (WS83), and on chromosome V near coordinate 80 kb (WS92). In addition, the diploids were generated by crosses of one haploid parent isogenic with J178-1d (related to S288c; Brock and Bloom 1994) and one haploid isogenic with PSL5 (isogenic with YJM789; Lee et al. 2009). Since YJM789 has ~60,000 SNPs distinguishing
its genome from S288c (Wei et al. 2007), the diploids used in our study are heterozygous for markers distributed throughout the genome. The primers used in strain constructions are listed in Table S2.

The S288c strain in the Saccharomyces Genome Database (SGD) has a single Ty2 element on the left arm of chromosome III. By PCR and Southern analysis (described in File S1), we found that the left arm of chromosome III in J178-1d contains a Ty1 element closely linked to the Ty2 element. In addition, YJM789 lacks both the Ty1 and the Ty2 elements on the left arm. The primers used to diagnose the location of Ty elements are in Table S2.

Analysis of loss of heterozygosity

In yeast strains that are heterozygous for markers, mitotic crossovers or break-induced replication events can generate loss of heterozygosity (LOH) of markers centromere distal to the recombination event (Figure 1; Lee et al. 2009; St. Charles et al. 2012). We mapped the transition between heterozygous markers and homozygous markers using two techniques: a PCR-based method and an approach utilizing oligonucleotide-containing microarrays. Primers used for the PCR-based approach are listed in Table S3. The details of these methods are described in the Results section and in File S1.

Identification of strains with recombination events induced by dicentric chromosome breakage

The GAL-CEN3 conditional centromere is inactive in cells grown in medium containing galactose and active in cells grown in glucose (Hill and Bloom 1987). In our experiments, all diploid strains were grown from single cells to...
colonies on solid medium containing galactose (YPGal) at 30° for 2 days. Individual colonies were selected from these plates, and restreaked on plates containing glucose (YPD) and incubated at 30° for 2 days. The resulting colonies were then replica plated to solid galactose-containing omission medium lacking uracil (SGal–Ura) to identify derivatives that had lost the URA3 marker adjacent to the conditional centromere.

Ura– derivatives could be generated by recombination (crossovers or BIR; Figure 1A and 1B), gene conversion events unassociated with crossovers or intrachromosomal deletions that remove the GAL-CEN3/URA3 cassette (Figure 1C) or chromosome loss (Figure 1D). Since chromosome loss would result in LOH for markers on both centromeric arms (Figure 1D), we examined Ura– derivatives using a PCR-based analysis to detect LOH for markers on both the right and left arms of chromosomes III or V (details in File S1). Similarly, since strains with conversion events unassociated with crossovers or intrachromosomal deletions of the cassette containing the conditional centromere would retain heterozygosity for markers located near the end of the chromosome (Figure 1C), we excluded such events by a PCR-based analysis to examine heterozygosity for markers near the left telomeres of chromosomes III and V (details in File S1). The remaining Ura– derivatives were analyzed for the location of recombination breakpoints.

**Statistical analysis**

We performed two types of statistical tests. First, we determined whether the distributions of chromosome break sites were significantly different from a random distribution. Second, we looked for associations between recombination breakpoints and various chromosome elements (replication origins, tRNA genes, etc.). The details of these analyses are given in File S1.

**Results**

**Experimental rationale**

The diploid strain used to map breaks induced in a dicentric chromosome has two important features: (1) it is heterozygous for a conditional centromere and (2) it has multiple single-nucleotide polymorphisms (SNPs) that distinguish the homologs. The conditional centromere employed in our analysis is described by Hill and Bloom (1987). The activity of the centromere is regulated by the galactose-inducible GAL1 promoter located ~100 bp from CEN3 sequences. When cells are grown on galactose-containing medium, transcription initiated at the GAL1 promoter inactivates the centromere; in glucose-containing medium, transcription is shut off and CEN3 is functional (Hill and Bloom 1987). Hill and Bloom (1989) inserted a cassette containing GAL-CEN3 and URA3 within the HIS4 gene on the left arm of chromosome III located ~50 kb from the natural centromere. In strains with this insertion, chromosome III is unstable or stable depending on whether the strain is grown in glucose- or galactose-containing media, respectively. In conditions in which both centromeres are active, the intercentromeric region is stretched and, subsequently, broken (Thrower and Bloom 2001; Fisher et al. 2009). In haploid strains, these breaks result in chromosomes with deletions of either the conditional centromere or the natural centromere (Hill and Bloom 1989; Kramer et al. 1994).

Diploid cells have an option for the repair of DSBs that is unavailable to haploids, the use of the intact homologous chromosome as a template. Two of the expected pathways of repair that would result in loss of the conditional centromere are reciprocal crossovers (Figure 1A) and break-induced replication (Figure 1B). For both of these pathways, loss of the conditional centromere (detected as loss of the URA3 gene) would result in LOH of markers located centromere distal to the breakpoint. In experiments in which recombination events reflect DSBs formed at trinucleotide repeats (Tang et al. 2011) or at meganuclease-recognition sequences (Nickoloff et al. 1999), the site of the DSB maps at or near the transition to LOH. Thus, in our experiments, we mapped the transition between heterozygous and homozygous markers as a method of mapping dicentric breaks (discussed in the Mapping of LOH events section of Results below). It should be noted that our mapping of DSBs is limited to those DSBs that are repaired by recombination with the homolog. DSBs that are repaired by sister-chromatid exchange (Kadyk and Hartwell 1992) do not lead to LOH and are undetectable by our analysis.

Breakage of the dicentric chromosome was induced by restreaking cells from individual colonies grown on galactose-containing medium (inactive GAL-CEN) to glucose-containing medium (active GAL-CEN). The resulting colonies were then replica plated to medium lacking uracil. We expected most of the Ura– derivatives of the starting strain could be grouped into three classifications. In class 1 strains (Figure 1, A and B), a DSB resulting from breakage of the dicentric is repaired by a reciprocal crossover or a BIR event. Such events will result in LOH of all markers centromere distal to the position of the recombination event with retention of heterozygosity for markers located on the opposite chromosome arm. Since we analyze only the Ura– product, crossovers and BIR events are not distinguishable; in wild-type strains, however, crossovers are more common than BIR events (McMurray and Gottschling 2003; Ho et al. 2010). In class 2 strains (Figure 1C), a DSB located near the conditional centromere could be repaired to generate a gene conversion event unassociated with a crossover or could be repaired to generate an intrachromosomal deletion; both of these events result in an interstitial LOH event. For class 2 strains, we expect LOH for markers located near the conditional centromere and maintenance of heterozygosity for markers flanking the conversion event. Class 3 strains (Figure 1D) reflect chromosome loss and result in LOH for markers located on both arms of the chromosome.

To distinguish among these classes, our initial characterization of the strains utilized multiple markers located in the
region between the conditional centromere and the native centromere, a marker located near the telomere, and a marker located on the opposite arm of the chromosome (details about coordinates of all markers in File S1). Three different dicentric strains were examined in our study: WS49 (conditional centromere located at SGD coordinate 67 kb of chromosome III), WS83 (conditional centromere located at SGD coordinate 32 kb of chromosome V), and WS92 (conditional centromere located at SGD coordinate 80 kb of chromosome V). For WS49, the telomere-associated marker was located at SGD coordinate 33 kb, ~34 kb centromere distal to the conditional centromere. Since the median size of mitotic gene conversion tracts is 6–8 kb (Lee et al. 2009; St. Charles et al. 2012), most of the conversion tracts that include the conditional centromere should not include marker 33. The opposite-arm marker was at coordinate 116 kb. For the strains WS83 and WS92, the telomere-associated marker was at coordinate 7 kb, located ~25 kb from the conditional centromere in WS83 and 73 kb from the conditional centromere in WS92; the markers on the opposite arm were located at coordinates 561 kb (WS83) and 152 kb (WS92). In summary, for class 1 events, we see a continuous region of LOH beginning at a marker located within the intercentromeric region and extending through the marker located near the telomere; the marker on the opposite chromosome arm retains heterozygosity. For class 2 events, we observe LOH for markers near the conditional centromere but retention of heterozygosity for the marker near the telomere and the marker on the opposite side of the centromere. Class 3 events are defined by strains in which LOH is observed for all markers tested, including the marker on the opposite side of the centromere. Some of the Ura⁻ strains (~10%) could not be classified into any of these three classes. In most of these strains (defined as class 4; complex events), there was more than one transition between heterozygous and homozygous markers. Such events may reflect the “patchy” repair of mismatches within a heteroduplex (St. Charles et al. 2012) or multiple repair events.

From the strains WS49, WS83, and WS92, we isolated 80, 61, and 62 Ura⁻ derivatives, respectively. The percentages of each class of event in the four strains were: WS49 (class 1, 34%; class 2, 14%; class 3, 44%; and class 4, 9%), WS83 (class 1, 41%; class 2, 36%; class 3, 20%; and class 4, 3%), and WS92 (class 1, 39%; class 2, 39%; class 3, 11%; and class 4, 11%).

Mapping of LOH events

In all diploids used in our experiment, the conditional dicentric homolog had many SNPs distinguishing it from its monocentric homolog. Most of the diploids were generated by a cross of haploids isogenic with J178-1d (Hill and Bloom 1989) with a haploid (PSL5) isogenic with YJM789 (Wei et al. 2007). The J178-1d strain has a mixed pedigree, but most of the SNPs that we examined were identical to those of S288c. Two methods were used to diagnose LOH. The first method (described in detail in File S1) was that used in our previous study (Lee et al. 2009). In brief, by BLAST comparisons of the DNA sequences of S288c and YJM789 in the intercentromeric region, we identified SNPs that altered a restriction enzyme recognition site. For each such SNP, we designed primers flanking the SNP. Following PCR amplification of genomic DNA, the resulting DNA fragment was treated with the diagnostic restriction enzyme and analyzed by gel electrophoresis to determine whether the diploid with the recombination event was heterozygous or homozygous for the SNP.

An example of this analysis is shown in Figure 2. In this example, we examined the Ura⁻ derivative WS49-8 for LOH at a SNP located at SGD coordinate 91322 (marker 91 in Table S3). At this position, the haploid parental strain PSL5 (isogenic with YJM789) had a HaeIII site that was lacking in DNA from the other parental strain (J178-#7-20). The primers used to amplify the region containing this SNP resulted in a fragment of ~500 bp. As shown in Figure 2, the fragment derived from the J178-#7-20 strain was not cut by HaeIII, whereas the fragment derived from PSL5 was cut into two fragments of ~170 and 320 bp. The control diploid WS49, following treatment of the 500-bp fragment with HaeIII, had three bands of the appropriate size. The Ura⁻ derivative WS49-8 had the pattern indicating LOH in favor of the PSL5-derived SNP. For WS49, we examined LOH in the intercentromeric region using 14 markers (listed in Table S3). Similar methods were used to map recombination events on chromosome V in diploids WS83 and WS92. For these two strains, we employed the same 34 SNPs previously described (Lee et al. 2009). Detailed mapping of recombination events was restricted to class 1 strains and our subsequent conclusions about dicentric breakpoints are based on only these strains.

In addition to the PCR-based mapping method described above, seven of the events were also analyzed using oligonucleotide-containing microarrays (SNP arrays). Using 25-base oligonucleotides that are identical to PSL5- or J178-1d-derived SNPs, we detected LOH events in the intercentromeric regions as described in our previous studies (St. Charles et al. 2012). The results of the microarray analysis, in general, were in good agreement with those obtained by the PCR-based methods and will be described in more detail below.

Mapping break sites in a strain (WS49) with a dicentric chromosome III

In the diploid WS49, the dicentric homolog has an insertion of the conditional centromere within the HIS4 locus at SGD coordinate 67 kb (Hill and Bloom 1989). In addition to SNPs that distinguish the two homologs, the J178-#7-20-derived homolog has a pair of Ty elements (one Ty1 and one Ty2) located centromere distal to LEU2 that are not present in the PSL5-derived homolog (File S1). We mapped recombination events in 27 class 1 Ura⁻ derivatives (Figure 3). In this figure, each horizontal line denotes the distance between the markers at the transition between heterozygosity and homozygosity.
Although it is evident that the dicentric-induced breaks map throughout the intercentromeric region, the distribution of breaks is nonrandom. To examine the distribution, we divided the intercentromeric region into four intervals of similar size (markers at the ends of intervals shown in parentheses): interval 1 (68–81), interval 2 (81–91 plus 6 kb Ty element), interval 3 (91–102), and interval 4 (102–115). We compared the observed distribution of events in these intervals with the expected distribution if the events were random (details in File S1). The distribution was significantly different from random by chi-square analysis (P = 0.03). We also tested each of the four intervals separately and determined that interval 1 (near the conditional centromere) had a significant (P = 0.01) excess of breakpoints relative to the sum of the other intervals. This value remained significant (P < 0.05) after correcting for multiple comparisons using the method of Benjamini and Hochberg (1995).

**Mapping break sites in strains (WS83 and WS92) that have a dicentric chromosome V**

The elevated frequency of breakpoints near the conditional centromere in WS49 could reflect a property of the conditional centromere. Alternatively, this region of chromosome III could be intrinsically susceptible to chromosome breaks. To distinguish between these two possibilities, we constructed two diploid strains isogenic with WS49 where the conditional centromere was located on chromosome V instead of III. In WS83, the conditional centromere was inserted at the \textit{CAN1} locus \~120 kb from the natural \textit{CEN5}, and in WS92, the conditional centromere was located on the same chromosome arm \~72 kb from \textit{CEN5}.

We previously mapped breakpoints for spontaneous mitotic recombination events between \textit{CAN1} and \textit{CEN5} using the PCR-based method (Lee et al. 2009). The mapping of recombination events reflecting dicentric breaks in WS83 was performed using the same 34 markers employed in our previous study. As shown in Figure 4A, the breakpoints cluster near the conditional centromere. For the statistical analysis, we divided the intercentromeric region into eight intervals: interval 1 (33–49), interval 2 (49–64), interval 3 (64–80), interval 4 (80–94), interval 5 (94–112), interval 6 (112–126), interval 7 (126–141), and interval 8 (141–152). The distribution of events in WS83 was significantly (P = 0.01) different from random. In addition, the number of events in interval 1 was significantly elevated relative to the sum of the events in the other intervals (P < 0.0001, corrected for multiple comparisons).

In our previous analysis of spontaneous mitotic recombination events, the region near \textit{CAN1} had elevated frequencies of recombination (Lee et al. 2009). To determine whether the DNA sequences near the conditional centromere had elevated recombination in regions that were not intrinsically prone to exchange, we constructed the diploid WS92 in which the conditional centromere was inserted near SGD coordinate 80 kb. This region does not have elevated spontaneous mitotic recombination (Lee et al. 2009) and has a low frequency of dicentric breakpoints in WS83 (Figure 4A). As shown in Figure 4B, in WS92, the breakpoints of this new dicentric chromosome are clustered near the conditional centromere. The distribution is significantly (P < 0.0001) nonrandom, and the interval closest to the conditional centromere (interval 4) is significantly enriched for breakpoints relative to the sum of the other intervals (P = <0.01, corrected for multiple comparisons). In summary, these results demonstrate that the region near the conditional centromere is particularly susceptible to breakage in dicentric chromosomes.
Analysis of recombination breakpoints for overrepresentation of elements of chromosome structure or sequence

A number of factors or chromosome elements in yeast have been suggested to be associated with elevated levels of DNA breakage or mitotic recombination (Aguilera et al. 2000 and references in File S1). Using statistical tests described in File S1, we looked for significant overrepresentation of such elements within the dicentric recombination breakpoints including palindromic sequences (≥16 bp), tandem repeats (repeats between 2 and 213 bp with a minimum repeat tract of 24 bp), G4 DNA (four tracts of three G’s separated by spacers <25 bp), tRNA genes, ARS elements, triplet repeats (≥8 repeats), long terminal repeats, peaks of gamma-H2AX, Rrm3p pause sites, replication-termination regions, and highly transcribed genes. For this analysis, we excluded the 10-kb region adjacent to the conditional centromere, since this region has an elevated frequency of recombination breakpoints by a mechanism likely to be specific for the conditional centromere. After corrections for multiple comparisons, no significant correlations with any of these elements were observed in any of the three strains or in the combined data of all three strains.

Mapping of dicentric breakpoints using SNP microarrays

All of the events shown in Figures 3 and 4 were mapped using the PCR-based approach. We also used SNP microarrays to map seven events (WS49-16, WS49-37, WS49-39, WS83-12, WS83-30, WS83-39, and WS83-40) also examined by the PCR-based approach. This alternative method confirmed our mapping procedure and, in addition, mapped the events to higher resolution. We used oligonucleotide-containing microarrays (St. Charles et al. 2012) capable of distinguishing SNP heterozygosity and homozygosity on chromosomes III and V (details in File S1). In Figure 5, A and B, we show low- and high-resolution analysis of a recombination event on chromosome V (WS83-40). In this figure, the blue lines and squares indicate the normalized hybridization ratio (experimental to control) to the YJM789-specific oligonucleotides, and the red lines and diamonds show the normalized hybridization ratio to the J178-1d-specific oligonucleotides. The transition between heterozygosity and homozygosity for markers is between SNPs located at coordinates 141,779 and 144,265, in good agreement with our previous PCR-based mapping of the event between markers 141 and 144 kb. As expected, since the recombination is initiated by breakage of the dicentric chromosome, the homozygous region is derived from the YJM789-related parental strain PSL5. The “spike” of increased hybridization for the J178-1d-specific SNPs near SGD coordinate 30,000 in Figure 5A results from deletion of the YJM789 sequences associated with the insertion of the SUP4-o gene into the YJM789-derived chromosome. Similar patterns of hybridization, confirming our PCR-based mapping, were also observed for strains WS49-16, WS49-37, and WS49-39.

The strain WS83-12 had a different pattern of hybridization than most of the other samples. By the PCR-based mapping method, this Ura− strain had a homozygous–heterozygous transition between markers 94 and 99. Marker 25, however, located centromere distal to the conditional centromere retained heterozygosity. The microarray analysis showed that this strain has two transitions, one located between SGD coordinates 96,550–97,221 and another located distal to the conditional centromere between coordinates 33,332 and 33,770 (Figure 5C). This pattern of hybridization suggests that WS83-12 resulted from a very long (60 kb) gene conversion event unassociated with a crossover, or a double crossover. Since the microarray analysis showed that SNPs derived from the YJM789-related homolog were duplicated, this hybridization profile is not consistent with a large heterozygous deletion. The breakpoints for WS83-12 or other strains that retained heterozygosity distal to the conditional centromere were not used in subsequent analyses.

In the strain WS83-30, the pattern of hybridization indicates that two events occurred. First, there was a crossover or BIR event near SGD coordinate 125 kb; high-resolution analysis

Figure 3 Mapping of dicentric chromosome break sites on chromosome III. Recombination events in 27 class 1 Ura− derivatives of WS49 are shown. Using the PCR-based method described in Figure 2, we mapped recombination events between the conditional centromere located at SGD coordinate 68,096 and CEN3, a distance of ~53 kb (including the heterozygous Ty elements). Each horizontal black line represents an independent event. The length of the line shows the distance separating the closest centromere-proximal heterozygous markers and the closest centromere-distal homozygous markers for each derivative. The J178-1d-derived homolog (shown in black) has two Ty elements (blue arrows) that are missing in the PSL5-derived homolog. Dotted lines show the intervals (I1–I4) used in the statistical analysis described in the text.
shows a breakpoint between SNPs located at 127,038 and 128,941. Following the recombination event, the recombinant chromosome was lost in about half of the cells. A similar hybridization pattern was observed for WS83-39. Elevated frequencies of nondisjunction leading to loss (Campbell and Fogel 1977) or gain (Chua and Jinks-Robertson 1991) of recombinant chromosomes has been previously reported.

In summary, the mapping of recombination events by microarrays supports our previous analysis using the PCR-based method. These results demonstrate that mitotic homologous recombination is an important mechanism for the repair of DSBs generated by breakage of a dicentric chromosome. In diploid strains that are unable to repair DSBs by homologous recombination, the broken chromosome can be “capped” by addition of telomeric repeats (Kramer and Haber 1993), resulting in a large terminal heterozygous deletion. Since this process is much less efficient than homologous recombination (Kramer and Haber 1993), we did not expect it to contribute to the Ura' events in our experiments. The SNP arrays in Figure 5 are consistent with LOH resulting from mitotic recombination rather than deletion formation because the LOH regions have an elevated level of hybridization of the YJM789-specific SNPs associated with a reduced level of hybridization to the J178-1d-specific SNPs. A heterozygous deletion would have a reduced level of hybridization to the J178-1d-specific SNPs without an accompanying increase in the level of hybridization to the YJM789-specific SNPs. Our results, therefore, indicate most of our LOH events reflect mitotic recombination between homologs.

Discussion

Our analysis of recombination events associated with breakage of a dicentric chromosome shows two types of events. About half of the events are located near the conditional centromere, and the other half are randomly distributed in the intercentromeric region. Below, we discuss the interpretation of our mapping results in the context of previous studies of mitotic recombination and chromosome fragility.

Mitotic recombination and LOH

In S. cerevisiae, most DSBs are repaired by homologous recombination. In diploid cells in G2 of the cell cycle, the preferred substrate for the repair of a DSB is the sister chromatid, although the homologous chromosome is also used (Kadyk and Hartwell 1992). Of these two pathways, only exchange involving the homologs leads to LOH of markers distal to the event (Figure 1). In yeast, genetic evidence indicates that spontaneous recombination events are often initiated by a DSB in unreplicated chromosomes and that such breaks are preferentially repaired using the homolog as a substrate (Esposito 1978; Lee et al. 2009; Lee and Petes 2010, St. Charles et al. 2012). In the current experiments, although the DSBs are likely formed in anaphase, it is
Unclear whether dicentric-associated DSBs occur on one chromatid (as shown in Figure 1) or both chromatids simultaneously. If a DSB was induced in only one dicentric chromatid that was repaired using the sister chromatid as a template, the repaired chromosome would retain two centromeres and, therefore, would be likely to break in the subsequent cell cycle. The Ura\textsuperscript{2} strains in our study are enriched for events in which the broken dicentric chromosome recombines with the intact monocentric homolog.

The repair of a DSB in one homolog using the intact homolog as a substrate can occur through a number of different pathways (Heyer \textit{et al.} 2010). In the synthesis-dependent strand-annealing (SDSA) pathway, the broken DNA ends invade the intact homolog, synthesize DNA sequences that span the DNA break, dissociate from the template, and reanneal (Andersen and Sekelsky 2010). The net result of this process is a gene conversion event, the nonreciprocal transfer of sequences between the two chromosomes. This process would result in an interstitial region of LOH (Lee \textit{et al.} 2009; St. Charles \textit{et al.} 2012) rather than an LOH region extending to the telomere. Unless the conversion event removed the conditional centromere, it would not generate a Ura\textsuperscript{2} derivative in our experiments and would not be detectable. Figure 5C shows an example of a possible gene conversion event.

Most of the Ura\textsuperscript{2} strains examined in our experiments have the pattern of LOH that is consistent with repair of the dicentric by crossover or BIR events (Figure 1). We cannot distinguish between these two events in our analysis because we do not recover both daughter cells resulting from dicentric breakage. Since BIR events are less frequent than crossovers in wild-type strains (McMurray and Gottschling 2003; Barbera and Petes 2006; Ho \textit{et al.} 2010), however, it is likely that most of the events represent crossovers.

An important issue is whether our observed recombination breakpoints accurately map the positions of the recombination-initiating DSBs. One possibility is that the broken ends undergo substantial nucleolytic degradation, forming large double-stranded gaps. By this mechanism, the position of the initiating DSB would be located telomere
proximal to the location of the mapped LOH event. This scenario is unlikely for two reasons. First, degradation of the broken ends usually involves loss of one of the two DNA strands rather than formation of a double-stranded DNA gap (Symington and Gautier 2011). Second, events associated with a mitotic recombination hotspot (Tang et al. 2011) or an HO-induced DSB (Nickoloff et al. 1999) map close to the site of the DNA lesion. Another factor that affects the relationship between the site of the initiating DSB and LOH breakpoint is gene conversion. Most mitotic crossovers are associated with an adjacent tract of gene conversion (Pâques and Haber 1999; Lee et al. 2009; St. Charles et al. 2012). Since the conversion tracts have a median size of 6–8 kb, our mapping of the DSB sites is limited to that level of resolution. In addition, our analysis of LOH is also limited by the number of markers analyzed, with an average of ~4 kb between markers on chromosomes III and V. Thus, our study yields only an approximate map position of the DSBs associated with breakage of a dicentric.

**Clustering of DSBs near the conditional centromere**

In all three dicentric strains examined, about half of the recombination breakpoints are within 10 kb of the conditional centromere. The comparison of the breakpoints in strains WS83 and WS92 demonstrate that clustering of events is a consequence of a property of the conditional centromere, since the region of chromosome V that has few events in WS83 (the interval between markers 81 and 91) becomes a hotspot for recombination when the conditional centromere is inserted nearby in WS92 (Figure 4, compare A and B). The mapping of breakpoints also suggests that the effect of the conditional centromere on DSBs extends at least 10 kb, since more than one of the marked intervals have elevated levels of events.

There are two related features of chromosome structure that extend about 10 kb from the yeast centromeres. First, cohesins are preferentially associated with yeast centromeres in a region extending 10–20 kb from the centromere (Blat and Kleckner 1999; Glynn et al. 2004). Second, the pericentric cohesins are bound intramolecularly, forming loops (Yeh et al. 2008). Several studies in *Schizosaccharomyces pombe* and *S. cerevisiae* have shown that the pericentric regions are more “stretchable” in preanaphase than other chromosomal regions (summarized by Thrower and Bloom 2001), although it is unclear whether this property is related to cohesin binding.

One explanation of our results is that the pericentric region located near the conditional centromere in the dicentric is more highly extended than other regions of the intercentromeric region, including the pericentric region located near the natural centromere. Why should the two pericentric regions behave differently? There is evidence that yeast centromere function is dependent on chromosome context. Megee et al. (1999) showed that centromere-associated loading of cohesins was more extensive in plasmids in which the centromeres were flanked by regions of low GC content. We measured the GC content in a 10-kb window centered on the natural and conditional centromeres in our strains. These percentages are: 34.6% (*CEN3*), 35.6% (*CEN5*), 45.1% (conditional *CEN3* in WS49), 38.7% (conditional *CEN3* in WS83), and 40.6% (conditional *CEN3* in WS92). In all three dicentric strains, therefore, the conditional centromere is flanked by sequences that have significantly (*P < 0.0001 by chi-square analysis*) higher GC content than the natural centromere on the dicentric chromosome.

As discussed previously, although breakage of the dicentric chromosome requires the tension established by stretching the chromosome between different spindle poles, the mechanical force exerted by this tension is insufficient to break double-stranded DNA. One scenario is that the preferential extension of the chromosome near the conditional centromere results in increased access to cellular nucleases that generate recombinogenic DSBs. A related possibility is that stretching of the chromosome near the conditional chromosome increases the probability of DSBs formed by nucleases preloaded in the pericentric region. Top2p accumulates to high levels near the centromere during the S period (Bermejo et al. 2009). Although most Top2p binding is lost by G2/M, a small number of persisting Top2p molecules could be sufficient to generate DSBs in the pericentric region.

Another mechanism for chromosome breakage in the dicentric is scission of the chromosome during nuclear fission or cytokinesis (Quevedo et al. 2012). DNA breaks could be a direct consequence of the physical forces exerted during cell division or, perhaps more likely, an indirect consequence of stretching of the chromosome, followed by endonucleolytic cleavage. By this model, the conditional centromere would be located more closely to the cell cleavage plane than the natural centromere. Whatever the explanation for the elevated level of recombination breakpoints near the conditional centromere, our results argue that the conditional centromere has some properties that are different from the natural centromere.

Other evidence that the region near the conditional centromere is more susceptible to breakage than the region near the natural centromere is based on analysis of deletions associated with rad52 dicentric haploid strains (Kramer et al. 1994). In such strains, chromosomes become stabilized by deletion of one of the two centromeres. Kramer et al. (1994) found that deletion of the conditional centromere was about four times more common than deletion of the natural centromere. Since most of these deletions are likely to be initiated by a DSB, these results argue that the region near the conditional centromere is prone to breakage. We cannot, however, rule out the possibility that the preferential recovery of chromosomes with the natural centromere in the experiments of Kramer et al. (1994) was a consequence of a lower rate of nondisjunction of chromosomes with the wild-type centromere compared to chromosomes with only the conditional centromere.

In addition, studies in mammalian cells and plants have suggested that, in dicentric chromosomes, one centromere...
can be more prone to inactivation than the other (Sullivan and Schwartz 1995; Lamb et al. 2008; Stimpson et al. 2012); centromere inactivation in mammals involves both intracentromeric structural changes (deletions) and epigenetic mechanisms (Stimpson et al. 2010).

**Dicentric breakpoints that are not associated with the conditional centromere**

In addition to the recombination breakpoints located near the conditional centromere, we observed breakpoints that are widely distributed throughout the intercentromeric region. We were unable to associate these breakpoints with any single specific chromosome element. It should be pointed out, however, that the resolution of mapping the events and the relatively small number of mapped events makes the statistical analysis challenging. The wide distribution of events argues that the dicentric breaks occur at some common element of chromosome structure (for example, a promoter or other nucleosome-free region) or that the breaks are associated with multiple different types of chromosome elements. We also do not know the enzyme or enzymes associated with breakage of the dicentric, although the topoisomerases are obvious candidates. It is also important to mention that the pattern of recombination breakpoints produced in our system with one conditional centromere and one natural centromere may be different than recombination events induced in a chromosome with two natural centromeres.

Pobiega and Marcand (2010) created dicentric chromosomes with one conditional centromere and one natural centromere in which the dicentric was generated by a fusion between two telomeres. Upon activation of the conditional centromere, DSBs were observed. Although ~40% of these breaks occurred within or near the telomere–telomere fusion (Pobiega and Marcand 2010), DSBs were also detected near the centromeres (S. Marcand, personal communication). The preference for breakage near the telomeric fusion may reflect a particular property of telomeric chromatin or the palindromic nature of telomere–telomere fusions.

In summary, there appear to be three types of chromosome breaks associated with dicentric chromosomes in yeast: breakage near (within 10 kb of) conditional centromeres, breakage at the junction of telomere–telomere fusions (Pobiega and Marcand 2010), and breaks that occur quasirandomly in the intercentromeric region.

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Nonrandom Distribution of Interhomolog Recombination Events Induced by Breakage of a Dicentric Chromosome in *Saccharomyces cerevisiae*

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**Strain Construction:** The genotypes of all strains in this study are given in Table S1. In all of the diploids used in our mapping studies, one of the haploid parental strains was derived by transformation of the haploid J178-1d (Brock and Bloom, 1994). J178-1d was created by a complex series of crosses, at least one of which involved the haploid strain S288c. In one derivative of J178-1d (J178#7-20), a conditional centromere (GAL-CEN3) and a URA3 gene were inserted within the HIS4 locus (SGD coordinates 65934-68333) of chromosome III (his4::GAL CEN3::URA3) (Brock and Bloom, 1994). The orientation of the conditional centromere is the same as that of the wild-type CEN3.

In another haploid derivative of J178-1d (MG42), the conditional centromere and URA3 gene replaced the CAN1 locus (SGD coordinates 31694-33466) on the left arm of chromosome V (can1-∆::GAL-CEN3::URA3). This strain was constructed by transforming J178-1d using a PCR fragment obtained by amplifying plasmid pR285 #7/pR285-GALCEN3#7 DNA (Brock and Bloom, 1994) with the primers GALCEN3/CAN F and GALCEN3/CAN R; the sequences of all primers used in strain constructions are in Table S2. The resulting PCR fragment contains the GAL-CEN3 URA3 cassette with sequences derived from the CAN1 locus at the ends of the fragment. We selected Ura’ transformants and screened those transformants for resistance to canavanine. We also confirmed the location of cassette by PCR using primer pairs CANupF with URA3R, and CANdnR with pBR322Ftest. We confirmed that the CAN1 gene was deleted by using the primer pair CANF and CANR. The orientation of the conditional centromere inserted on chromosome V is opposite to that of the conditional centromere inserted on chromosome III in J178#7-20.

In the haploid strain MG48, the conditional centromere and the URA3 marker were located on the left arm of chromosome V replacing the region between Saccharomyces Genome Database (SGD) coordinates 80163 and 80362 (V80163-80362∆::URA3::GAL-CEN3). This strain was constructed by transforming J178-1d using a PCR fragment obtained by amplifying plasmid pR285 #7/pR285-GALCEN3#7 DNA with the primers GALCEN3/80k F and GALCEN3/80k R, and selecting Ura’ transformants. The insertion of the conditional centromere at the correct site was confirmed using three sets of primer pairs: 80kupF with URA3R, 80kdnR with pBR322Ftest, and 80kupF with 80kdnR.

We also used a haploid strain (PSL5) derived from the sequenced clinical isolate YJM789 (Wei et al., 2007); PSL5 (MATα ade2-1 ura3 can1Δ::SUP4-o gal2 ho::hisG) has been described previously (Lee et al., 2009). The diploids used in our study were constructed by the following crosses: WS49 (J178#7-20 x PSL5); WS83 (MG42 x PSL5); WS92 (MG48 x PSL5). All strains are
heterozygous for SNPs located throughout the genome (St. Charles et al., 2012). WS49, WS83, and WS92 are isogenic except for the location of the conditional centromere.

**Analysis of structural differences between the chromosome III homologs derived from J178-1d and YJM789**

In addition to SNPs that distinguish the chromosome III homologs derived from J178-#7-20 and PSL5, there were also insertions of Ty elements and other structural changes. It is have been noted previously (Hill and Bloom, 1989; Wicksteed et al., 1994) that chromosome III from J178-1d is larger than observed in most other yeast strains. Using ORF-containing microarrays and procedures described in McCulley et al. (2010), we found that J178-#7-20 had a duplication on the right arm of chromosome III that included ORFs between YCR019W and YCR027C. The location of this duplication suggests that it was generated by unequal crossing-over between two previously-mapped pairs of Ty elements termed FS1 and FS2 (Umezlu et al., 2002; Lemoine et al., 2005).

We also examined Ty elements located on the left arm of chromosome III in the two strains by Southern analysis and a series of PCR reactions. The S288c-related strain described in SGD has a single Ty2 element located between KCC4 and LEU2, although some other strains contain both Ty1 and Ty2 between these two genes. We isolated genomic DNA from MS71 (a control wild-type strain; Sia et al., 1997), PSL5, and J178-#7-20, and treated the samples with AclI, a restriction enzyme that does not cut within Ty elements. The resulting fragments were examined by standard Southern analysis, using a probe containing KCC4 sequences; this probe was prepared using genomic DNA and primers RKCC4-1 and KCC4-1 (Table S2). The observed sizes of the AclI restriction fragment hybridizing to the probe were 6.5 kb, 11 kb, and 18 kb for the strains PSL5, MS71, and J178-#7-20, respectively. These sizes suggest that PSL5 has no Ty element near LEU2, MS71 has one Ty element, and J178-#7-20 has two Ty elements. This conclusion was confirmed by analyzing fragments generated by double digests of genomic DNA with Ncol and Nsil. The observed sizes of fragments hybridizing to the KCC4 probe were 8 kb (PSL5), 5 kb (J178-#7-20), and 14 kb (MS71). Since Nsil has three recognition sites in Ty1, but not in Ty2, these results support the conclusion that PSL5 lacks Ty elements near LEU2, MS71 has one Ty2 element, and J178-#7-20 has closely-linked Ty1 and Ty2 elements.

These conclusions were further supported by PCR analysis. Using primers KCC4F2 82211 and Ty2R 85162 (a primer with homology to both Ty1 and Ty2), we observed no amplification with PSL5 genomic DNA, an 800 bp fragment with J178-#7-20 DNA, and a 3 kb fragment with MS71 DNA. Using primers KCC4F2 82211 and Ty2R 85553 (a primer that is Ty2-specific), we observed a low level of a fragment of 1.6 kb with PSL5, a 10 kb fragment with J178-#7-20 DNA, and a 3.3 kb fragment with MS71 DNA. Assuming that the low level of the 1.6 kb fragment observed with PSL5 is non-specific, these results argue that PSL5 lacks a Ty2 element, J178-#7-20 has a centromere-distal Ty1 element and a centromere-proximal Ty2 element, and MS71 has only a Ty2 element. We confirmed that J178-#7-20 has a Ty1 element in the Watson orientation using two PCR primer pairs:
Ty1F with KCC4 F2 82211, and Ty1R with KCC4 F2 82211. Only the PCR reaction of Ty1R and KCC4 F2 82211 yielded the 800 bp fragment as expected if the Ty1 element was in Watson orientation.

**Identification of strains with recombination events induced by dicentric chromosome breakage**

The GAL-CEN3 conditional centromere is inactive in cells grown in medium containing galactose and active in cells grown in glucose (Hill and Bloom, 1987). In our experiments, all diploid strains were grown from single cells to colonies on solid medium containing galactose (YPGal) at 30°C for two days. Individual colonies were selected from these plates, and re-streaked on plates containing glucose (YPD) and incubated at 30°C for two days. The resulting colonies were then replica-plated to YPGal medium lacking uracil to identify derivatives that had lost the URA3 marker adjacent to the conditional centromere. The percentages of colonies that were either Ura− or sectored Ura+/Ura− in cells grown on YPD-containing plates (median value of five independent cultures) were 78% (WS49), 94% (WS83), and 93% (WS92). The median percentages of Ura− or sectored Ura+/Ura− colonies (median value of five independent cultures) in cells grown on galactose-containing plates (inactive conditional centromere) and then plated on galactose-containing plates were 4% (WS49), 2% (WS83), and 2% (WS92).

**Analysis of loss of heterozygosity (LOH) using restriction digests of PCR fragments**

In yeast strains that are heterozygous for markers, mitotic crossovers can generate loss of heterozygosity of markers centromere-distal to the crossover (Lee et al., 2009; St. Charles et al., 2012). The transition between heterozygous markers and homozygous markers, therefore, locates the position of the crossover. We looked for LOH using two procedures, an approach in which LOH was detected by a PCR-based approach (describe below), and an approach utilizing oligonucleotide-containing microarrays. For the first approach, we used genomic DNA sequence information to identify SNPs that distinguished the two haploid strains in the region between the conditional centromere and the natural centromere on chromosome III. We then determined which of these SNPs altered a restriction site, and designed primers that would amplify a region of several hundred bp flanking the SNP. For example, there is a SNP at SGD coordinate 70426 that results in an Hpy166II site in the PSL5 strain that is absent in the J178-#7-20 strain. We designed primers flanking this site that produce a fragment of about 500 bp. If we treat this fragment produced by PCR amplification from the heterozygous diploid strain with Hpy166II and analyze the fragments by gel electrophoresis, we obtain three fragments: about 500 bp (representing the SNP derived from the J178-#7-20 strain), and about 360 and 140 bp (representing the SNP derived from the PSL5 strain). In diploids that undergo LOH for a SNP at this position, we observe either one 500 bp fragment or two fragments of 360 and 140 bp. The location of the SNPs, the primers used to produce the restriction fragments, and the diagnostic restriction enzyme are shown in Table S3.
In the WS49 strain, which has the conditional centromere on chromosome III, we first examined polymorphic markers located centromere-distal to the conditional centromere (33 and 58), within the intercentromeric region (68, 81, 102, 103, 111, 113), and located on the opposite chromosome arm (116) for LOH; the names of the SNPs reflect their approximate SGD coordinates in kb (Table S3). As explained in the Main Text, based on the patterns of LOH observed with these markers, we classified the Ura’ derivatives of WS49 as Class 1, 2, 3, or 4. We then performed more detailed mapping of recombination breakpoints in Class 1 strains using other markers (70, 74, 78, 91, 96, 98, 106, and 107). The conditional centromere on III is inserted near SGD coordinate 67 kb, and the natural centromere is near coordinate 114 kb.

By a similar approach, we mapped crossovers in strains with the conditional centromere on chromosome V. For these experiments, we used the primers and restriction enzymes described in Lee et al. (2009). For the preliminary mapping of strain WS83, we used the chromosome V markers 7, 25, 41, 70, 112, 133, and 561; the conditional centromere is inserted near SGD coordinate 32 kb, and CEN5 is located at SGD coordinate 152 kb. The other markers used for mapping are: 35, 43, 44, 46, 49, 52, 55, 56, 57, 60, 64, 76, 80, 83, 87, 92, 94, 99, 104, 108, 114, 115, 117, 119, 122, 126, 141, 144, 147, and 151. For the preliminary mapping of strain WS92, we used the chromosome V markers 7, 52, 76, 83, and 152. The other markers used for mapping are: 87, 92, 94, 99, 104, 108, 112, 119, 133, 141, 144, 147, and 151.

The primers and restriction enzymes used to analyze heterozygous SNPs on chromosome V are in Table S2 of Lee et al. (2009) with the exception of markers 7, 25, 152, and 561. Markers 7 and 25 are located centromere-distal to the conditional centromeres which are located at 32 kb in MS83 and 80 kb in WS92. Marker 7 is located at SGD coordinate 7005. For this polymorphism, we amplified genomic DNA with the forward primer ATCCTCATTCCAGCTACCT (starting coordinate at 6857) and reverse primer AGTAGTACCTGTTTTAATGGG (starting coordinate at 7249). The diagnostic restriction enzyme was DraI, which cuts the genomic DNA of MG42 and MG48, but not that of PSL5; both MG42 and MG48 are derived from J178-1d. The PCR fragment used to check marker 25 was generated with the primers 5’ CACCTGAGCCACGCATCTG and 5’ GCAACGTGGGAAGAAAACG. We tested Ura’ strains derived from WS83 and WS92 for LOH of a telomere-associated SNP using a polymorphism located on chromosome V at SGD coordinate 24903 (marker 25). The YJM789-derived homolog has a HindIII site at this position that is absent in J178-1d derivatives. The primers used to generate the PCR fragment with this polymorphism were 5’ CACCTGAGCCACGCATCTG and 5’ GCAACGTGGGAAGAAAACG.

For strains WS83 and WS92, we used different markers located on the chromosome V arm opposite the conditional centromere. For WS83, we used a marker (561) located at 560715. For this polymorphism, we amplified genomic DNA with the
forward primer TTCTCAGCCGTACAATCATGC (starting coordinate at 560490) and reverse primer AAACTCTTCTCAAGGCTCTGG (starting coordinate at 560980). The diagnostic restriction enzyme was EcoRI, which cuts the genomic DNA of PSL5, but not that of MG42. For WS92, we used a marker (152) located at 152163. For this polymorphism, we amplified genomic DNA with the forward primer TTGGTAAACAAAGGCAAGC (starting coordinate at 151849) and reverse primer ATGTGGCGTTTGTGCAG (starting coordinate at 152295). The diagnostic restriction enzyme was Cac8I, which cuts the genomic DNA of MG48, but not that of PSL5.

**Analysis of loss of heterozygosity (LOH) using SNP microarrays**

Three recombination events derived from WS49 and four events derived from WS83 were mapped by both the PCR/restriction enzyme method described above and by oligonucleotide-containing microarrays. Because the stability of short duplexes is sensitive to mismatches, it is possible to design an oligonucleotide-containing microarray that can distinguish whether SNPs are heterozygous or homozygous (Gresham et al., 2010). Previously, we designed Agilent SNP arrays to look for LOH throughout the genome in a diploid formed by a cross of W303a (closely related to S288c) and YJM789 (St. Charles et al., 2012). For each SNP analyzed (about 15,000 distributed throughout the genome), four 25-base oligonucleotides were used, two identical to the Watson and Crick strands of the W303a-specific SNP and two identical to the Watson and Crick strands of the YJM789-specific SNP; the polymorphism was the central base of the oligonucleotide. The sequences of the oligonucleotides used in the construction of the microarray are given in St. Charles et al. (2012).

To look for LOH in strains with a recombination event, we labeled genomic DNA from the control strain with a Cy5-tagged nucleotide and DNA from a control heterozygous strain with Cy3-tagged nucleotide. The labeled samples were mixed and hybridized to the microarray. Following hybridization, the arrays were scanned and the ratio of hybridization to the two samples was determined as described previously (St. Charles et al., 2012). Ratios of hybridization for each oligonucleotide were normalized to the Cy5/Cy3 ratio of all of the oligonucleotides on the microarray. In general, the recombination breakpoints determined by the PCR-based method were in good agreement with those determined by microarrays (discussed further in Results).
Statistical Analysis

We performed two types of statistical tests. First, we determined whether the distributions of chromosome break sites in the diploid strain were significantly different from a random distribution. For this analysis, we divided the region between the conditional centromere and the natural centromere into approximately equal-sized intervals. Based on the number of recombination events mapped for each diploid, we calculated the expected number of events in each interval and these numbers were compared to the observed distribution by chi-square analyses; these chi-square tests were done the VassarStats Website (http://faculty.vassar.edu/lowry/VassarStats.html). For WS49, the intervals used (markers shown in parentheses) were: Interval 1 (68 to 81); Interval 2 (81-91), Interval 3 (91-102), and Interval 4 (102-115). For WS83, the physical intervals were: Interval 1 (33-49), Interval 2 (49-64), Interval 3 (64-80), Interval 4 (80-94), Interval 5 (94-112), Interval 6 (112-126), Interval 7 (126-141), and Interval 8 (141-152). For the diploid WS92, we used Intervals 4 to 8 as specified for WS83.

As will be described in the Results section, for all three diplods, we found that the region located approximately 10 kb centromere-proximal to the conditional centromere had an elevated frequency of recombination events. Regions located outside of this “hotspot” were examined to determine if various chromosome elements (replication origins, palindromic sequences, and other elements described below) were over-represented at the recombination breakpoints. For purposes of this calculation, we excluded the intercentromeric region located near the conditional centromere. The lengths of the mapped regions (MRs) examined and the number of events for each strain were WS49 (33538 kb, 14), WS83 (103088 kb, 15), and WS92 (57657 kb, 8). These MRs were calculated by subtracting the SGD coordinate corresponding to the most centromere-distal SNP used in the mapping (excluding the hotspot) from the SGD coordinate representing the boundary of the “natural” centromere. The sum of the MRs for each strain is equivalent to the MR multiplied by the number of events. Thus, the sums of the MRs are 469532 kb (WS49), 1546320 kb (WS83), and 461256 kb (WS92). We define the recombination breakpoints (RBs) as the distance separating the last heterozygous site from the first homozygous site. The analysis was performed in several steps. First, for each strain, we summed the lengths of the RBs over all of the mapped events within the MR; the sum of the RBs for each strain were 72651 kb (WS49), 68900 kb (WS83), and 35469 kb (WS92). Second, we determined the lengths of sequences that are not RBs (NRBs) for all of the mapped events (equivalent to the sum of the RBs subtracted from the sum of the MRs for each strain); the NRBs were 396881 kb (WS49), 1477420 kb (WS83), and 425787 kb (WS92). Third, for each element, we calculated its density within the MR (the number of elements divided by the MR distance). Fourth, we calculated the expected number of elements within the summed RBs by multiplying the density of the chromosomal element by the summed RBs; we calculated the
expected number of elements in the NRBs in a similar way. We then compared the expected numbers of elements within and outside of the RBs with the observed numbers within and outside of the RBs by chi-square analysis.

To illustrate the method, we will describe our analysis to determine whether palindromic sequences are over-represented in recombination breakpoints (RBs) in the WS49 data. To determine the location of the palindromic sequences (> 16 base pairs), we used the data in Lisnic et al. (2005). There are four palindromic sequences located in the 33538 kb MR between SGD coordinates 80845 and 114383, a density of 0.00012/kb. Since we had 14 events, the sum of palindromes for all the events is 56. In WS49, we observed that five palindromes were in the RB regions and 51 palindromes were in the NRB regions. The expected number of palindromes in the RB regions, assuming a random distribution, is 0.00012/kb x 72651 kb or 8.7; the expected number in the NRB regions is 0.00012 x 396881 or 47.3. When the observed and expected numbers were compared by chi-square test, the p value was 0.24, indicating that palindromes are not significantly enriched at the recombination breakpoints.

A similar analysis was done for each strain with the following elements (descriptions of the element and references in parentheses): tandem repeats (repeats between 2 and 213 bp with a minimum repeat tract of 24 bp; Gelfand et al., 2007), G4 DNA (four tracts of 3 G’s separated by spacers <25 bp; Capra et al., 2010), tRNA genes (SGD), ARS elements (SGD), triplet repeats (> 8 repeats; Gelfand et al., 2007), long terminal repeats (SGD), peaks of gamma-H2AX (Szilard et al., 2010), Rrm3p pause sites (Azvolinsky et al., 2009), and replication-termination regions (Fachinetti et al., 2010). We also looked for correlations with highly-transcribed genes. For this analysis, we determined the number of highly-transcribed genes in the MRs for all three strains, defining a highly-transcribed gene as a gene ranking in the top 20%, using the database of Nagalakshmi et al. (2008). For each of these genes, we calculated the midpoint of the transcript, and then determined whether these midpoints were over-represented in the RBs.

For the three individual strains, none of the examined elements had a significant over-representation at the breakpoints when corrections were performed for multiple comparisons. Since the number of events per strain was small, we also examined each element summed over all three strains. For this analysis, we added the numbers of observed and expected events for each category of element, and performed a chi-square analysis on the totals. None of the elements was significantly over-represented in this analysis.
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Table S1  Strain genotypes

| Strain | Genotypes                                      |
|--------|------------------------------------------------|
| J178-7-20 | MATa ade1 met14 ura3-52 leu2-3,112 his3 his4::GAL-CEN3/URA3 rad52 |
| PSL5 | MATa ade2-1 ura3 can1Δ::SUP4-o gal2 ho::hisG |
|       | MATa/MATa ade1/ADE1 ADE2/ade2-1 met14/MET14 ura3-52/ura3 leu2-3,112/LEU2 his3/HIS3 his4::GAL-CEN3/URA3/HIS4 rad52/RAD52 CAN1/can1Δ::SUP4-o |
| WS49 | GAL2/gal2                                    |
| J178-1d | MATa ade1 met14 ura3-52 leu2-3,112 his3 |
| MG42 | MATa ade1 met14 ura3-52 leu2-3,112 his3 can1Δ::GAL-CEN3/URA3 |
| WS83 | MATa/MATa ade1/ADE1 ADE2/ade2-1 met14/MET14 ura3-52/ura3 leu2-3,112/LEU2, his3/HIS3, can1Δ::GAL-CEN3/URA3/can1Δ::SUP4-o GAL2/gal2 |
| MG48 | MATa ade1 met14 ura3-52 leu2-3,112 his3 V80412-80162Δ::GAL-CEN3/URA3 |
|       | MATa/MATa ade1/ADE1 ADE2/ade2-1 met14/MET14 ura3-52/ura3 leu2-3,112/LEU2 his3/HIS3, V80412-80162Δ::GAL-CEN3/URA3/V80412-80162 |
| WS92 | CAN1/can1Δ::SUP4-o GAL2/gal2 |
### Table S2  Primers used in strain constructions and in the mapping of Ty elements

| Primer name     | Sequence (5’ to 3’)                                      |
|-----------------|----------------------------------------------------------|
| GALCEN3/CAN F   | ATGACAAATCCAAAGAAGCCGACATAGAGGAGAAGCATATGTTTCTCATGTTTGCAGCTTA |
| GALCEN3/CAN R   | CTATGCTACAACATTCAAAAATTTTGCTCCTCCTACATGTTTCTGACTGGCGATATAGGCG |
| CAN1F           | AGGCCGGACGAAAGCTAAAC                                        |
| CAN1R           | TACATGGAGACATCTACTGGG                                      |
| pBR322Ftest     | TCGCTACTTGGAGCCACTATC                                      |
| URA3R           | AGCAACAGGACTAGGATGAG                                       |
| CANupF          | AATCTGTCGTAATCGAAAG                                        |
| CANdnR          | TTATACATTAGATTTAGCGT                                        |
| GALCEN3/80k F   | TCAATTCCACTCACACCGTCATGTTGATTTATTTATCATCCTCCTTTTCTATGTTGACAGCTTA |
| GALCEN3/80k R   | TTCTCGAGCTACACACGAGATATATATATGTTGTGTGCTTTTCTTGTGACTGGCGATATAGGCG |
| 80kupF         | TGGTATACGAGCTAGCAGGAC                                       |
| 80kdnR         | AGAGAACTATTCTGACTTAC                                        |
| KCC4 F2 82211  | ACGCTGACCAAAGCTGTACTC                                        |
| Ty2R 85162     | AACCAGACATAGGCGCTAC                                           |
| Ty2R 85553     | AGCTTGAGGTACTTCAGAGGAG                                       |
| RKCC4-1        | CGTTATCTTTCTTGACTATCCTTCTTTC                                |
| KCC4-1         | GCGAAGTCCCCAAATAGCGAGAGATC                                    |
| Ty1F           | TGCAGACGTAAAGCCCAAACCTC                                       |
| Ty1R           | TGTGGACTTCTTGAAGTTAACC                                       |
Table S3  Primers used in PCR-based analysis of polymorphic markers on left arm of chromosome III

| SGD coordinates for polymorphic site (marker name) | F primer sequence (5’ to 3’) | R primer sequence (5’ to 3’) | S288c SGD coordinates of amplification (5’ of F primer) | S288c SGD coordinates of amplification (5’ end of R primer) | Diagnostic restriction enzyme | Strain with restriction enzyme site |
|----------------------------------------------------|------------------------------|------------------------------|--------------------------------------------------------|----------------------------------------------------------|-----------------------------|-----------------------------------|
| 33032 (33) GGTATTGAAGCCGCAAGTT | TGG                           | TCTCCTTCGACACCTTCATCG        | 32634                                                  | 33031                                                     | BbvI                         | PSL5                              |
| 57640 (58) TTGCAAGTGTGAGAGGCC | AGGATGGTTTTGCTCCATTG          | 57355                                                                 | 57814                                                  | HpyCH4IV                                                   | J178-#7-20                   |
| 68096 (68) TCTGTACGTACTTCACC | TGGTTTTGCGGATT CTAC           | 67897                                                                 | 68331                                                  | NheI                                                      | J178-#7-20                   |
| 70426 (70) ATCGCAGGATAGCTATGG | AGAGCTTGCCATAGAAGGCC         | 70294                                                                 | 70788                                                  | Hpy166II                                                   | PSL5                         |
| 74203 (74) G                           | TCGCGTTACAAAAACAGATG          | 73929                                                                 | 74425                                                  | BstBI                                                      | PSL5                         |
| 77646 (78) ACCCATAATAGCGTAACC     | TCGCAATGTCGTCGTGC            | 77434                                                                 | 77926                                                  | BanII                                                      | J178-#7-20                   |
| 80845 (81) T C                       | AGGCCCTAGTATATGACGAAAG       | 80589                                                                 | 81025                                                  | HhaI                                                      | PSL5                         |
| 91332 (91) GCTATCGCACAAGAATCA     | AGTGGAACACCTGTAAGC          | 91011                                                                 | 91496                                                  | HaeIII                                                     | PSL5                         |
| 96056 (96) AC                        | ATCATCAAACAATGAATGACC        | 95913                                                                 | 96412                                                  | BsrBI                                                      | PSL5                         |
| 97697 (98) TGCAGCATTACTATCCTTA     | TGGTTTGATCCCATCTGAAGTGA      | 97313                                                                 | 97859                                                  | BtgI                                                       | PSL5                         |

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| Position  | Sequence 1 | Sequence 2 | Enzyme  | Restriction Site |
|-----------|------------|------------|---------|-----------------|
| 102361    | TGTGGTTTCAATTCCGG | TAGAATTGGAATTCAACGG | HpyCH4V | J178-#7-20 |
| 103500    | TTGATCTACTCCATATTTCC | TGAAG TACTC AAAGT CGTAGC | Hpy99I | J178-#7-20 |
| 105855    | ATGGCAGTTCGCGATGCG | ATACAGAGAGATGGAGAC | BstNI | J178-#7-20 |
| 106777    | AGCAGTGGAGTATGGCGA | ATTCGTTGGACCTTGTTC | Hpy99I | J178-#7-20 |
| 111457    | TGGAGTAATGGAAATGCC | TTGTCTGCAAGGCTTGCC | Rsol | J178-#7-20 |
| 112761    | ATTCAACGAAACATTCG | TGAAGTCTATTGTGCCAC | BstBI | J178-#7-20 |
| 116012    | TCCCAACTCCCTGGCTAATG | ATGGCGAGGTAGCTGATTG | BstI | J178-#7-20 |