

A Natural Meiotic DNA Break Site in *Schizosaccharomyces pombe* Is a Hotspot of Gene Conversion, Highly Associated With Crossing Over

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Manuscript received October 5, 2004

Accepted for publication November 16, 2004

ABSTRACT

In *Schizosaccharomyces pombe*, meiosis-specific DNA breaks that initiate recombination are observed at prominent but widely separated sites. We investigated the relationship between breakage and recombination at one of these sites, the *mbs1* locus on chromosome I. Breaks corresponding to 10% of chromatids were mapped to four clusters spread over a 2.1-kb region. Gene conversion of markers within the clusters occurred in 11% of tetrads (3% of meiotic chromatids), making *mbs1* a conversion hotspot when compared to other fission yeast markers. Approximately 80% of these conversions were associated with crossing over of flanking markers, suggesting a strong bias in meiotic break repair toward the generation of crossovers. This bias was observed in conversion events at three other loci, *ade6*, *ade7*, and *ura1*. A total of 50–80% of all crossovers seen in a 90-kb region flanking *mbs1* occurred in a 4.8-kb interval containing the break sites. Thus, *mbs1* is also a hotspot of crossing over, with breakage at *mbs1* generating most of the crossovers in the 90-kb interval. Neither Rec12 (Spo11 ortholog) nor I-SceI-induced breakage at *mbs1* was significantly associated with crossing over in an apparently break-free interval >25 kb away. Possible mechanisms for generating crossovers in such break-free intervals are discussed.

IN eukaryotic cells, DNA double-strand breaks (DSBs) are a dangerous form of DNA damage and yet are also crucial for the process of meiosis. During vegetative growth, DSBs are a threat because, left unrepaired, they cause chromosomal fragmentation. In addition, repair of DSBs can promote genomic instability, including DNA rearrangements resulting from homologous recombination. In contrast, the deliberate generation of DSBs appears to be a conserved feature of meiosis precisely because of their ability to promote homologous recombination and generate crossovers. Crossovers, in turn, are critical for correct meiotic chromosome segregation and the promotion of genetic diversity among the products of meiosis. The importance of DSB formation in generating meiotic crossovers has been well established in *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe* and inferred in other organisms (SUN *et al.* 1989; CAO *et al.* 1990; CERVANTES *et al.* 2000; KEENEY 2001). In contrast, the mechanistic relationship between the location and frequency of DSBs and the location and frequency of crossovers is less well understood.

It is commonly held that in *S. cerevisiae* meiotic breaks and crossovers are coincident, but detailed examination of the location of DSBs and crossovers in *S. pombe* revealed a different pattern. DSBs are distributed in pre-

ferred sites at widely separated physical distances (~50–100 kb on average). In contrast, crossovers are observed at a relatively constant frequency per unit of physical distance (YOUNG *et al.* 2002); *i.e.*, crossovers are not consistently more frequent in intervals where strong breaks are observed than in apparently break-free intervals. It was proposed that the DSB and crossover patterns could be reconciled if meiotic DSBs are able to generate crossovers over large distances (tens of kilobases) around their positions (YOUNG *et al.* 2002).

Two factors control how the frequency of DSBs relates to the frequency of crossovers. First, DSBs can be repaired using either sister chromatids or homologous chromosomes (homologs), but genetically observable crossovers can arise only from interhomolog events. The relative frequency of intersister and interhomolog events will therefore affect the proportion of DSBs that generate observable crossovers. Second, interhomolog recombination does not always generate crossovers. More precisely, gene conversions (which indicate interhomolog recombination events) can occur with or without associated crossovers. For example, at the *S. cerevisiae* *ARG4* locus only ~30% of gene conversions are accompanied by a crossover (GILBERTSON and STAHL 1996), while at the *S. pombe* *ade6* locus ~60% are accompanied by a crossover (GRIMM *et al.* 1994). In other organisms, values as low as 15% and as high as 75% have been reported (WHITEHOUSE 1982).

It was long thought that gene conversions with crossovers and those without crossovers resulted from random cutting of a single kind of recombination interme-

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diate, the four-strand Holliday junction. Such random cutting should produce equal numbers of crossovers and noncrossovers. Recently, however, models that generate crossover and noncrossover products via two separate pathways have been proposed, only one of which involves resolution of Holliday junctions, with the second proceeding by synthesis-dependent strand annealing (SDSA). Holliday junction resolution may always result in crossovers, with noncrossovers arising solely from the second pathway (ALLERS and LICHTEN 2001; CROMIE *et al.* 2001; HUNTER and KLECKNER 2001; OSMAN *et al.* 2003; SMITH *et al.* 2003).

In many organisms the relationship between the frequency of meiotic DNA breakage and crossing over is further complicated by the phenomenon of interference, where the occurrence of one crossover inhibits additional crossovers in a surrounding region of variable size. However, while the budding yeast *S. cerevisiae* exhibits interference, some other fungi, including the fission yeast *S. pombe*, do not (STRICKLAND 1958; MUNZ 1994). It should be simpler to study the relationship between DSB and crossover frequencies in organisms lacking interference.

We set out to examine the location and frequency of recombination events associated with *mbs1* (meiotic break site 1), a naturally occurring meiotic break hotspot in *S. pombe* (YOUNG *et al.* 2002). We wished to know if *mbs1* is a conversion hotspot and, in addition, how important it is for generating nearby and distant crossovers. We also wished to measure how often crossovers are associated with gene conversions at *mbs1* and at other loci as a measure of the proportion of interhomolog recombination events that give crossovers. Values consistently higher or lower than 50% would indicate that crossovers and noncrossovers are not generated by random Holliday junction resolution in *S. pombe*.

MATERIALS AND METHODS

Yeast strains and genetic techniques: The *S. pombe* strains used in this study, and their genotypes, are listed in Table 1. Meiotic crosses were carried out by suspending single yeast colonies in 5 ml of supplemented yeast extract liquid medium + adenine (100 µg/ml; GUTZ *et al.* 1974) and growing at 30° until saturated. For each cross, aliquots of 100 µl from two saturated cultures were mixed, washed with water, and spotted on sporulation agar plates (SPA; GUTZ *et al.* 1974). After 2 days incubation at 25°, the cell-ascus mixture from each spot was suspended in 1 ml of water and treated with glucosylase and ethanol to kill vegetative cells, essentially as described by DEVEAUX *et al.* (1992). Tetrad dissection is described in GUTZ *et al.* (1974).

Scoring of the restriction site markers *l-4*, *L*, and *R* (see Figure 4) was done by colony PCR followed by digestion with the appropriate restriction enzyme and gel electrophoresis. Auxotrophic markers were scored by replica plating onto appropriately supplemented nitrogen-base minimal agar (PONTICELLI and SMITH 1989). Replica plating onto yeast extract agar + phloxin B plates (MORENO *et al.* 1991) at 37° was used to score *tps16*. In the case of the cross B tetrad analysis (Figure 2), *ura4* was scored by FOA^R. The *ura1* marker was scored directly among *ura4*⁺ colonies. In *ura4*⁻ spore colonies, the

TABLE 1

S. pombe strains used in this study

Strains	Genotype ^a
GP789	<i>h</i> ⁺ <i>ade6-M26 ura1-171</i>
GP877	<i>h</i> ⁻ <i>ade6-52 ura1-61</i>
GP1252	<i>h</i> ⁻ <i>ade6-52 ura4-D18 ura4⁺-aim tps16-23^b</i>
GP3362	<i>h</i> ⁺ <i>ade6-M26 his3-D1 ura4-D18 mbs1-1::ura4⁺</i>
GP3398	<i>h</i> ⁺ <i>ade6-216 leu2-32 ura4-D18 lys1-37 rec12-171::ura4⁺</i>
GP3410	<i>h</i> ⁻ <i>ade6-3008 ura1-61 rqh1-h2 pom1::3HA.kan</i>
GP3477	<i>h</i> ⁺ <i>mbs1-2 rqh1-h2</i>
GP3672	<i>h</i> ⁺ <i>ade6-M26 mbs1-2 rec12-172</i>
GP3673	<i>h</i> ⁺ <i>ade6-M26 rec12-172</i>
GP3674	<i>h</i> ⁻ <i>ade6-M26 mbs1-2 rec12-172</i>
GP3677	<i>h</i> ⁻ <i>ade6-M26 mbs1-2 rec12-173</i>
GP3696	<i>h</i> ⁻ <i>lys3-37 ura1-61 rqh1-h2</i>
GP3718	<i>h</i> ⁺ <i>ade6-3049 rad50S pat1-114 end1-458</i>
GP3731	<i>h</i> ⁺ <i>ade6-M26 mbs1-102 his3-D1</i>
GP3785	<i>h</i> ⁺ <i>ade6-3034 ura4-D18 arg1-14 leu1-32</i>
GP3801	<i>h</i> ⁺ <i>ade6-52 lys3-37 ura1-61 mbs1-2 rqh1-h2 rec12-172</i>
GP3802	<i>h</i> ⁻ <i>ade6-52 lys3-37 ura1-61 mbs1-2 rqh1-h2 rec12-172</i>
GP3805	<i>h</i> ⁻ <i>ade6-52 lys3-37 ura1-61 rqh1-h2 rec12-172</i>
GP3806	<i>h</i> ⁺ <i>ade6-52 lys3-37 ura1-61 mbs1-2 rqh1-h2 rec12-173</i>
GP3839	<i>h</i> ⁺ <i>ade6-M26 pat1-114 rad50S</i>
GP4282	<i>h</i> ⁺ <i>ade6-210 ura4-D18 mbs1-22::ura4⁺</i>
GP4514	<i>h</i> ⁺ <i>ade6-216 ura1-61 mbs1-102</i>
GP4515	<i>h</i> ⁻ <i>ade6-3049 mbs1-103 rqh1-h2</i>
GP5091	<i>h</i> ⁺ <i>ade7-C8 ura5-294</i>
GP5125	<i>h</i> ⁻ <i>swi5-201::kanMX6 ade7-50</i>
GP5145	<i>h</i> ⁺ <i>ade6-M26 his3-D1 ura4-D18 mbs1-102</i>

^a Mutations other than commonly used auxotrophies and mating-type alleles are described in the following references: *mbs1* alleles (this work), *pat1-114* (INO and YAMAMOTO 1985), *pom1::3HA.kan* (BAHLER and PRINGLE 1998), *rad50S* (FAHAR *et al.* 2002), *rec12-172* and *173* (this work), *rqh1-h2* (ENOCH *et al.* 1992; STEWART *et al.* 1997), *swi5-201::kanMX6* (ELLERMEIER *et al.* 2004), *tps16-23* (GYGAX and THURIAUX 1984), and *ura4⁺-aim* (ZAHN-ZABAL *et al.* 1995).

^b *tps16* mutations map to the *ags1* gene (F. HOCHSTENBACH, personal communication).

ura1 genotype was inferred from the genotypes of the *ura4*⁺ colonies (*i.e.*, if two colonies were *ura4⁺ ura1⁺*, it was assumed that the other two colonies were *ura4⁻ ura1⁻*) or scored on the basis of the configuration of flanking markers (*i.e.*, if the flanking markers *lys3* and *rqh1* were in the parental configuration, then it was assumed that *ura1* was also). This left only one ambiguous tetrad.

In the I-ScdI experiments (Table 2), potential diploids were identified as apparent double crossovers between *lys3-ura1* and *ura1-rqh1*, giving a wild-type phenotype for all of these markers. They were confirmed as diploids by iodine staining on SPA plates (*i.e.*, *h*⁺/*h*⁻) or by DNA content determined by flow cytometry.

Measurement of crossovers associated with gene conversions: Crossovers of flanking markers accompanying gene conversion at *mbs1* were measured in the same tetrads used to assess *mbs1* conversion frequencies. For other loci, gene convertants were selected as intragenic recombinants (prototrophs). The frequency of crossing over of flanking markers was then measured in these spore colonies. In all cases the frequency of crossovers in the total spore population was also measured and was assumed to represent the frequency of incidental crossovers in the convertant spore population. Observed crossovers arise when an associated crossover occurs

without an incidental crossover or when an incidental crossover occurs without an associated crossover; *i.e.*, $C_0 = C_A(1 - C_1) + C_1(1 - C_A)$, where C_0 is the observed frequency of crossovers among convertants, C_A is the frequency of crossovers associated with gene conversion, and C_1 is the frequency of incidental crossovers. The frequency of crossovers associated with gene conversion is then given by $C_A = (C_0 - C_1)/(1 - 2C_1)$. When $C_0 < 0.5$, then $C_A < C_0$, and when $C_0 > 0.5$, then $C_A > C_0$ for all values of C_1 . The flanking markers used with *mbs1* conversions were *ura1* and *rqh1*; with *ade6*, they were *ura4⁺-aim* and *tps16*; with *ura1*, they were *lys3* and *rqh1*; and with *ade7*, they were *swi5* and *ura5*.

Engineering restriction site markers at *mbs1*: The restriction site markers 1–4 are inactivations of *HaeII*, *BstBI*, *NciI*, and *BstBI* sites, respectively, by mutations at positions 6850 (G to A), 7564 (C to T), 8061 (G to A), and 8798 (T to C) using the numbering of cosmid SPAC4G8 (GenBank accession no. Z56276.2). These were generated by subcloning the 3.8-kb *SpeI* fragment containing these sites into a plasmid followed by site-directed mutagenesis using the Stratagene QuikChange kit. The modified *SpeI* fragment was then excised by restriction and used to transform strain GP3362 to FOA^R, resulting in strain GP5145. The markers *L* and *R* are creations of *PmlI* and *XbaI* sites, respectively. Marker *L* is a G-to-C mutation at position 5213 of SPAC4G8, and marker *R* is a T-to-A mutation at position 9977. These changes were made by transforming strain GP4282 with linear DNA carrying the appropriate point mutations (introduced by PCR primer mutagenesis) at the *L* and *R* sites to create *mbs1-103*. The marker *mbs1-22* in strain GP4282 is a replacement of the SPAC4G8 sequence 5208–9981 with that of a 1.8-kb *HindIII* fragment containing *ura4⁺* (GRIMM *et al.* 1988; GenBank accession no. X13976.1). The marker *mbs1-1* in strain GP3362 is a replacement of the SPAC4G8 sequence 6486–9213 with the same 1.8-kb *HindIII* fragment.

Construction of the inducible I-*SceI* cutting system: The *mbs1-2* allele creates an I-*SceI* recognition site at the *mbs1* locus (SPAC4G8 positions 8045–8062; Figure 3). The *mbs1-2* sequence was generated by PCR primer mutagenesis, and a modified 3.8-kb chromosomal *SpeI* fragment containing this mutation was cloned into a plasmid. The *SpeI* fragment was then excised and used to transform strain GP3362 to FOA^R, resulting in a chromosomal *mbs1-2* allele.

The *rec12-172* allele is a replacement of the *rec12* ORF with that of I-*SceI* while the *rec12-173* allele has the same replacement, but with the I-*SceI* ORF in the inverted orientation. A linear fragment containing the I-*SceI* ORF was produced by PCR from the plasmid pSCM525 (PERRIN *et al.* 1993) and used to replace the *rec12* ORF on pJF20 (J. FARAH, unpublished results) in both the correct and the inverted orientations. Linear DNA containing the two modified *rec12* loci was excised, producing linear DNA with the I-*SceI* sequence and *rec12* flanking DNA. The two kinds of linear DNA were used to transform GP3398 to FOA^R, resulting in chromosomal *rec12-172* and *rec12-173* alleles.

Analysis of meiotic DNA breaks: Meiotic inductions, flow cytometry, and preparation of DNA in agarose plugs were performed as described (YOUNG *et al.* 2002). The agarose-embedded DNA was digested with restriction enzymes, separated by gel electrophoresis, Southern blotted, and hybridized with the probes described. Probe c227 is described in YOUNG *et al.* (2002). Probe Spe1A is identical to the sequence 5972–6330 of cosmid SPAC4G8.

RESULTS

The *mbs1* locus is a 2.1-kb cluster of break sites: The *mbs1* locus is a prominent meiotic break site (~10% breakage) visible upon Southern blot analysis of the

501-kb *NotI* “J” fragment of chromosome I (YOUNG *et al.* 2002; Figure 1A). Analysis of a *PmeI* digest (YOUNG *et al.* 2002) localized *mbs1* to a 6-kb gene-free region (interval 5305–11,360 of cosmid SPAC4G8; GenBank accession no. Z56276.2). Finer resolution mapping was then undertaken by probing smaller DNA fragments generated by *SpeI* and *HpaI* digestion (Figure 1B; data not shown). This analysis revealed the *mbs1* locus to consist of four clusters of break sites spread over a 2.1-kb region (Figure 1C; data not shown). The fraction of DNA broken at these four break clusters ranged from 3.3% to 0.9% (Figure 1C), showing a gradient, with the strongest breakage toward the left end as drawn.

The *mbs1* locus is a hotspot for gene conversion: In *S. cerevisiae*, markers <1 kb from DSB sites show gene conversion at high frequency (NICOLAS *et al.* 1989; SUN *et al.* 1989; DETLOFF *et al.* 1992; FAN *et al.* 1995). In general, markers in *S. pombe* convert at low frequency except where mutation or engineering has resulted in high levels of DNA breakage nearby (GUTZ 1971; ZAHN-ZABAL *et al.* 1995; STEINER *et al.* 2002; YOUNG *et al.* 2002, p. 258). To test the natural *mbs1* break cluster for gene conversion, we mutated four restriction sites, each within one of the four clusters of break sites, by single-base-pair transitions (Figure 1C and Figure 2A; see MATERIALS AND METHODS). Strains carrying this modified *mbs1* locus (designated *mbs1-102*) were crossed with *mbs1⁺* strains. The asci produced from these crosses were dissected, and DNA from the spore colonies was assayed for the four restriction sites by PCR and restriction digestion.

The results of this analysis were as follows. Among the 144 tetrads analyzed, 15 showed conversion of at least one of the four restriction sites. One of these 15 tetrads showed two gene conversion events, giving a frequency of 11% conversion per tetrad (supplemental Figure 1 at <http://www.genetics.org/supplemental/>). Considering frequencies as events per chromatid, this represents 16 conversion events among 576 chromatids (2.8%). The four markers converted at comparable frequencies from 1.4% to 1.7% of chromatids (Figure 2B and supplemental Figure 1 at <http://www.genetics.org/supplemental/>). Gene conversion in *S. pombe* is generally infrequent (0.25% of tetrads per locus on average; YOUNG *et al.* 2002, p. 258), making the *mbs1* cluster of sites the most frequently converting markers reported in this organism, with the exception of previously described hotspot mutations. Co-conversion of adjacent markers was common, accounting for 10 of the 16 conversion events observed (63%; Figure 2B; supplemental Figure 1 at <http://www.genetics.org/supplemental/>). As three adjacent markers often co-converted, it appears that conversion tracts are frequently >1.2 kb in length (Figure 2B; supplemental Figure 1 at <http://www.genetics.org/supplemental/>; see DISCUSSION). Co-converting markers always converted to the same parental genotype. No discontinuous conversion tracts were observed, consistent with data from *S. pombe*

and other organisms that suggest that such events are rare (BORTS and HABER 1989; GRIMM *et al.* 1994).

Crossing over in a 90-kb interval containing *mbs1* is highly associated with *mbs1* gene conversion: We expected that the breaks observed at *mbs1* would generate crossovers in the 90-kb interval *ura1-rqh1* (in which *mbs1* is located; Figure 2A). To test this, we examined tetrads that could be scored for both *mbs1* gene conversions and crossovers between the flanking markers *ura1* and

rqh1. Gene conversions were used as a signal that recombinational repair of a break at *mbs1* had occurred. If *ura1-rqh1* crossovers were elevated among *mbs1* convertant spores compared to total spores, it would indicate that *mbs1* breakage frequently generated crossovers in that interval.

The tetrads we analyzed previously for conversion were scored for crossing over between *ura1* and *rqh1*. No conversions of the *ura1* or *rqh1* markers were seen among the 144 tetrads, consistent with the typical low frequency of gene conversion in *S. pombe*. Among the 576 spore colonies analyzed above, 62 showed crossing over in the *ura1-rqh1* interval, *i.e.*, 10.8% of chromatids (Figure 2C; supplemental Figure 1 at <http://www.genetics.org/supplemental/>). This is very similar to the value of 9.7% from random spore analysis (YOUNG *et al.* 2002). Of the 16 conversion events (see above), 13 (81%) showed an accompanying *ura1-rqh1* crossover (Figure 2C). Despite the small number of conversions, the frequency of *ura1-rqh1* crossovers among convertant spores (81%) is significantly higher ($\chi^2 = 69.9$; d.f. = 1; $P < 0.0001$) than that seen in the total spore population (10.8%) and demonstrates a high association between conversion (and presumably breakage) at *mbs1* and crossing over in the *ura1-rqh1* interval (see DISCUSSION).

To estimate how important breakage at *mbs1* is for crossing over in the *ura1-rqh1* interval, we considered the proportion of such crossovers having an associated *mbs1* conversion. As a single crossover produces two crossover spore colonies, the 62 crossover colonies observed represent 31 *ura1-rqh1* crossovers. Among this

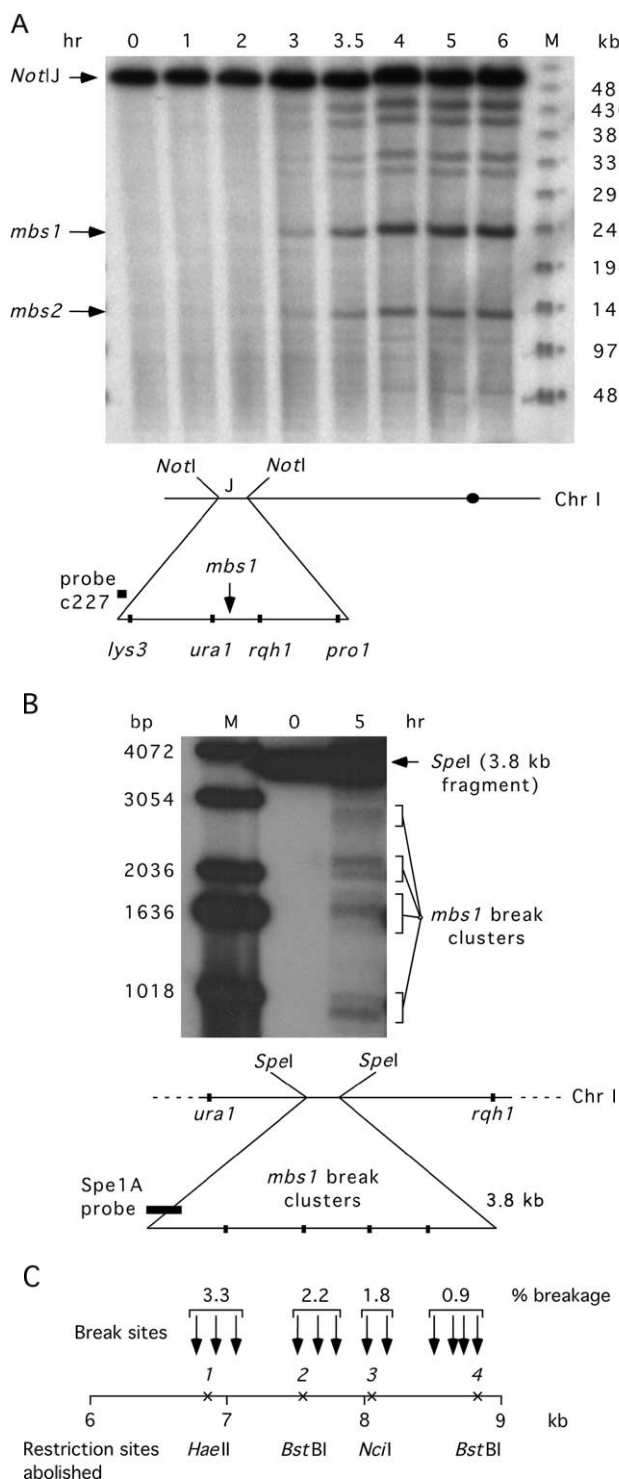


FIGURE 1.—Mapping DNA breakage at *mbs1*. (A) The *mbs1* locus is visible as an intense meiosis-specific break site ($\sim 10\%$ breakage) in the 501-kb *NotI* “J” fragment of chromosome I. DNA from the (break-accumulating) *rad50S* haploid strain GP3718 was harvested at the times indicated after the beginning of *pat1-114* (temperature-inducible) meiosis. The DNA was digested with *NotI*, electrophoresed, and probed as shown. Other meiotic breaks, including *mbs2*, are visible. M, phage- λ marker ladder with sizes in kilobases at the right. The approximate positions of the *NotI* “J” fragment and *mbs1* on chromosome I are indicated. (B) Finer resolution mapping of a *SpeI* digest of meiotic GP3839 DNA (probed as shown) revealed that the *mbs1* break locus consists of four break clusters. The 0- and 5-hr time points of a meiotic induction are shown. M, 1-kb marker ladder (Invitrogen, San Diego) with sizes in base pairs at the left. (C) The break sites were mapped by probing *HpaI* and *SpeI* digests of meiotic GP3839 DNA from each end (additional data not shown). In the first cluster, breaks mapped to positions 6780, 6920, and 7075 with respect to the SPAC4G8 cosmid (numbered telomere-centromere). In the second cluster, they mapped to 7525, 7670, and 7800; in the third to 8020 and 8190, and in the fourth to 8520, 8660, 8740, and 8850. These positions were derived as averages from the four probes, and we estimate their uncertainty to be ~ 50 bp. The intensity of breakage at the four clusters was quantified from the *SpeI* blots. Single-base-pair transitions were introduced to abolish four restriction sites (1, *HaeII*; 2, *BstBI*; 3, *NciI*; 4, *BstBI*) within each of the four break clusters (see MATERIALS AND METHODS). Shown to scale.

total population of 31 crossovers, 42% (13/31) were associated with a conversion event at *mbs1* (the crossover involved the converting chromatid). This suggests that at least 40% of all *ura1-rqh1* crossovers are initiated by *mbs1* breakage. We also noted that only one simple crossover was observed between the four *mbs1* restriction markers in the absence of conversion of any of those markers (supplemental Figure 1 at <http://www.genetics.org/supplemental/>). This has implications for how efficiently conversion of our markers accompanies inter-homolog DSB repair at *mbs1* (see DISCUSSION).

Crossovers >25 kb from *mbs1* are not associated with *mbs1* gene conversion: YOUNG *et al.* (2002) showed that there are genetic intervals in the 501-kb *NotI* “J” fragment containing *mbs1* that lack detectable DSBs but have substantial levels of crossing over. The model of

SZOSTAK *et al.* (1983) predicts that meiotic crossing over is caused by meiotic DSBs, and this is supported by the existence of multiple *rec* gene mutations that abolish both meiotic crossing over and DSBs in *S. pombe*. Therefore, YOUNG *et al.* (2002) hypothesized that breakage at the observed sites, such as *mbs1*, in the *NotI* “J” fragment could generate crossovers not only in the intervals containing the breaks but also in the apparently break-free intervals. One mechanism by which this could happen is by the migration of double Holliday junctions from the initiating break to the crossover position, where junction resolution would occur. As an example, in the interval *lys3-rqh1* there are only two observed break sites, *mbs1* (~10% breakage) and *mbs2* (~3% breakage; YOUNG *et al.* 2002), but several subintervals that do not include these break sites still experience crossovers (e.g., ~3.7% crossing over in *res2-ura1*; Figure 2A). The hypothesis of YOUNG *et al.* (2002) predicts that *mbs1* (with ~80% of all *lys3-rqh1* breakage) would be important for crossing over in much of the *lys3-ura1* interval (including subintervals like *res2-ura1*) and not just in the immediate subinterval (*ura1-rqh1*) that contains the *mbs1* locus.

To test this hypothesis, the association of gene conversion at *mbs1* and crossing over in the interval *lys3-ura1* (or the similar interval *pom1-ura1*) was examined as done previously for the interval *ura1-rqh1*. Again, gene conversions were used as a signal that recombinational repair of a break at *mbs1* had occurred. If crossovers in the *lys3/pom1-ura1* interval were elevated among *mbs1* convertant spores compared to total spores, it would indicate that *mbs1* breakage was responsible for crossovers in that interval.

The same set of tetrads that was studied above was

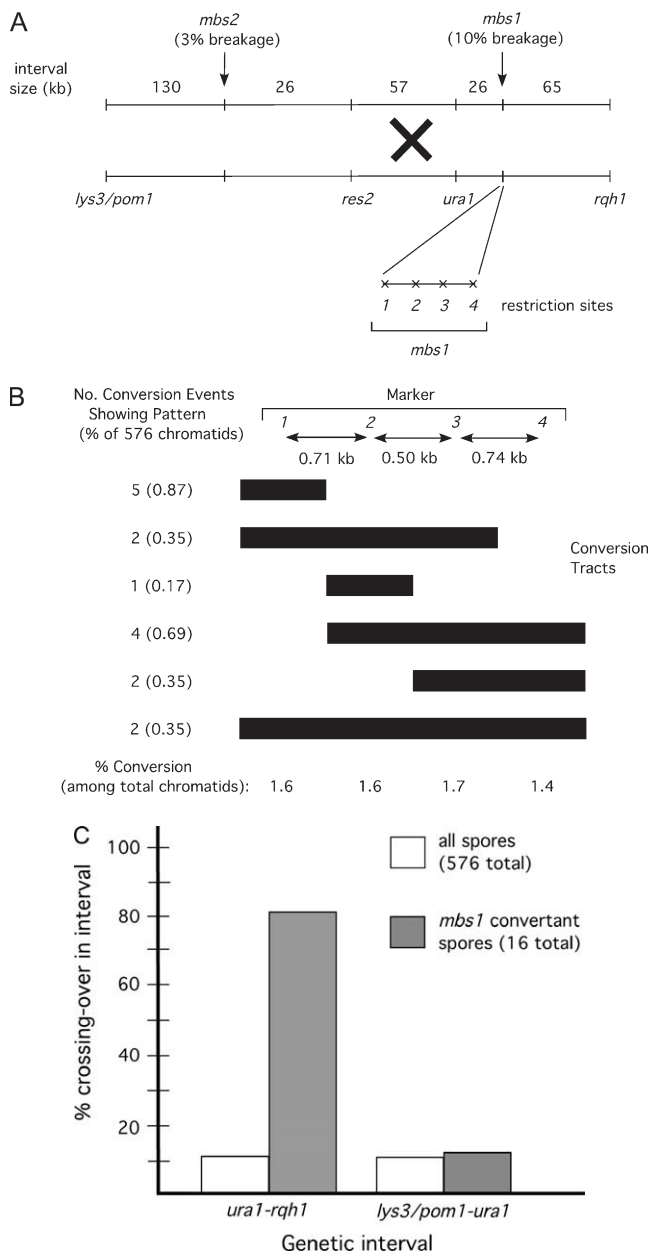


FIGURE 2.—The *mbs1* locus is a hotspot for gene conversion, highly associated with *ura1-rqh1* crossovers. (A) Map of *mbs1* and surrounding region showing markers used to measure crossover frequencies and the four restriction sites used to monitor gene conversion at *mbs1*. The physical sizes of the genetic intervals are shown along with observed meiotic break frequencies at sites *mbs1* and *mbs2*. The restriction sites (1–4) in Figure 1C were scored by PCR and restriction. (B) Conversion patterns of markers 1–4 observed at *mbs1* from 144 tetrads. Tetrads from two crosses (cross A—GP3696 × GP3731, 94 tetrads; cross B—GP3410 × GP5145, 50 tetrads) were scored for restriction sites to identify conversion frequency and distribution (see supplemental Figure 1 at <http://www.genetics.org/supplemental/>). Ten conversion tetrads were identified in cross A (including one apparent double-conversion tetrad) and five in cross B. Bars extend under the markers converted; endpoints are arbitrarily placed at midpoints between the markers scored. (C) Crossovers in the interval *ura1-rqh1*, but not in the interval *lys3/pom1-ura1*, are associated with *mbs1* conversions. The frequencies of crossovers in the intervals *lys3/pom1-ura1* and *ura1-rqh1* are shown among total spores and among *mbs1* conversion spores. The marker *lys3* was used in cross A and the marker *pom1* in cross B; these markers are only ~4 kb apart but are ~210 kb from *ura1* so the data for the crosses were pooled.

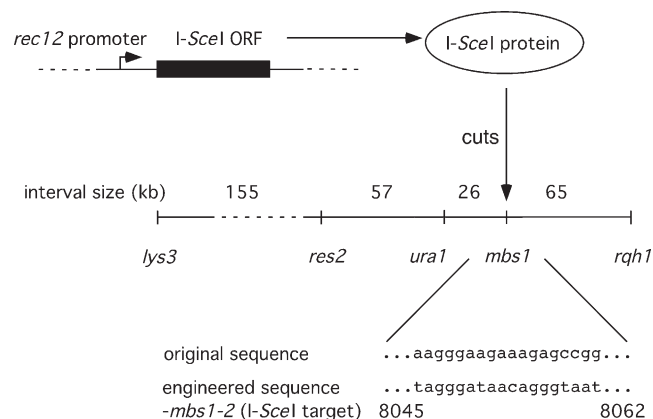


FIGURE 3.—System for generating a single meiotic DSB, located at the *mbs1* region of chromosome I. An exact replacement of the *rec12* ORF by that of I-SceI was engineered (see MATERIALS AND METHODS). A target site for I-SceI was made by mutating 9 bp in the *mbs1* region (numbering refers to the SPAC4G8 cosmid). In controls, the I-SceI ORF was placed in inverted orientation or the target site was absent. The markers used to score crossovers in various intervals are shown, along with the physical sizes of those intervals. Not to scale.

analyzed for *lys3/pom1-ura1* crossovers. The *ura1* marker lies ~26 kb from *mbs1* (Figure 2A). Among the 576 spore colonies analyzed above, 64 had crossovers in the *lys3/pom1-ura1* interval, i.e., 11.1% of chromatids (Figure 2C and supplemental Figure 1 at <http://www.genetics.org/supplemental/>). This is somewhat lower than the frequency previously reported for random spore data (~17%; YOUNG *et al.* 2002). In contrast to the strong linkage between gene conversion at *mbs1* and crossing over in the *ura1-rqh1* interval, only 2 of 16 conversions at *mbs1* were associated with a *lys3/pom1-ura1* crossover (12.5%; Figure 2C; supplemental Figure 1 at <http://www.genetics.org/supplemental/>). This is not significantly different from the level of *lys3/pom1-ura1* crossing over in the total population (11.1%) and does not support the hypothesis of YOUNG *et al.* (2002; see DISCUSSION).

An artificial meiotic DSB at *mbs1* does not frequently stimulate crossovers >25 kb away: As a more direct test of the hypothesis of YOUNG *et al.* (2002), we reduced meiotic DNA breakage to a single genomic site at the *mbs1* locus and observed where this breakage stimulated crossing over. This was done by replacing the coding sequence for Rec12, the *S. pombe* Spo11 ortholog that generates normal meiotic DSBs, with that of the I-SceI endonuclease (CARROLL 2004). As *rec12* is meiotically induced (LIN and SMITH 1994), this should give meiosis-specific I-SceI expression. The 18-bp I-SceI recognition site (COLLEAUX *et al.* 1988), which is absent from the *S. pombe* genomic sequence (WOOD *et al.* 2002), was introduced into the *mbs1* region (Figure 3) so that a single meiotic DSB could be generated there. Control strains were constructed with inactive I-SceI (an inversion of the I-SceI coding sequence at the *rec12* locus) or

TABLE 2

Crossover stimulation near and far from an artificial *mbs1* break

Cross ^a	% crossing over in interval		No. of colonies analyzed
	<i>lys3-ura1</i>	<i>ura1-rqh1</i>	
Active (cross C)	0.33	2.4	2123 ^c
No site (cross D)	<0.25 ^b	0.25	1193 ^d
Inverted ORF (cross E)	0.084	0.34	1187 ^e

^a The frequency of crossing over in the *lys3-ura1* and *ura1-rqh1* intervals was measured in a meiotic cross with I-SceI cutting at *mbs1* (Figure 3) or control versions of that cross. Cross C (GP3674 × GP3801 or GP3672 × GP3802) expressed I-SceI (*rec12-172*) and possessed the I-SceI target site at *mbs1* (*mbs1-2*) as shown in Figure 3. Cross D (GP3673 × GP3805) expressed I-SceI but lacked the I-SceI target site, and cross E (GP3677 × GP3806) had the I-SceI target site (*mbs1-2*) but an inverted (inactive) I-SceI ORF (*rec12-173*).

^b No colonies were observed; the value is the upper 95% confidence limit based on the Poisson distribution (where three colonies are assumed).

^c Sum of five experiments (GP3674 × GP3801, three experiments; GP3672 × GP3802, two experiments). Data from the two types of cross were not statistically different.

^d Sum of three experiments.

^e Sum of two experiments.

with no I-SceI recognition site at *mbs1*. Standard meiotic crosses were carried out and the frequency of crossing over among random spores was measured for the *lys3-ura1* and *ura1-rqh1* intervals (Figure 3). Complementing diploids, which occur in *rec12* meioses (DAVIS and SMITH 2003), would be scored as crossovers in both intervals. These were identified and removed from the analysis (see MATERIALS AND METHODS).

The results of this study are shown in Table 2. Breakage at *mbs1* specifically stimulated crossing over in the *ura1-rqh1* interval 10-fold compared to the controls ($\chi^2 = 40.3$; d.f. = 1; $P < 0.0001$; contingency χ^2 test with controls pooled). Statistically significant ($\chi^2 = 4.5$; d.f. = 1; $P < 0.05$) stimulation was also seen in the *lys3-ura1* interval but the effect was very small. Therefore, it appears that breakage at *mbs1* does not frequently generate crossovers in the *lys3-ura1* interval, i.e., >25 kb from *mbs1* (see DISCUSSION).

***mbs1* crossovers are concentrated in a 4.8-kb region around *mbs1*:** The results presented so far indicate that breakage at *mbs1* is strongly associated with crossing over in the *ura1-rqh1* interval and infrequently, if at all, with crossing over in intervals >25 kb away. This led us to investigate the distribution of crossovers within the *ura1-rqh1* interval. If crossovers are located close to the position of their initiating DSBs, then the *mbs1* locus should be a hotspot for crossovers and the rest of the *ura1-rqh1* interval should be cold. Distinguishing crossover position was not possible using the *I-4* restriction sites

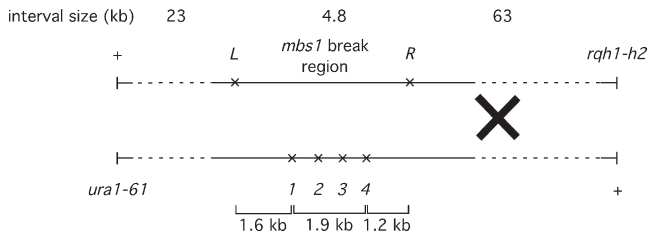


FIGURE 4.—Additional markers at *mbs1* used to identify the position of *ura1-rqh1* crossovers. *Pml*I (*L*) and *Xba*I (*R*) recognition sites were generated as shown by single-base-pair mutations (at positions 5213 and 9977 with respect to the SPAC4G8 cosmid). Cross F is shown (GP4514 × GP4515).

at *mbs1*, because frequent conversion of these markers generally made the location of crossovers ambiguous. Therefore, we introduced two additional restriction sites flanking *mbs1*, denoted “*L*” and “*R*” (Figure 4). Both were single-base-pair changes (see MATERIALS AND METHODS). These two markers define a 4.8-kb region containing the *mbs1* break sites (Figure 4) and, as they are 1.2 and 1.6 kb from the nearest observed break sites, were expected to convert sufficiently infrequently to reveal unambiguously if crossovers lay between them.

A meiotic cross heterozygous for these two new markers and the original four restriction markers was carried out (Figure 4). Tetrads with *ura1-rqh1* crossovers were identified, and analysis of the six markers was restricted to these. Only two-strand double-crossover tetrads, expected to make up ~5% of all *ura1-rqh1* crossovers tetrads, would be missed by study of this subpopulation. In addition, the data already presented showed that the great majority (~80%) of *mbs1* conversion events were associated with *ura1-rqh1* crossovers so that examination of crossover tetrads should also detect most *mbs1* conversion events.

In the 89 tetrads examined, 20 showed *ura1-rqh1* crossovers (one four-strand double, *i.e.*, 11.8% of chromatids, consistent with previous results). Among these 20 crossover tetrads, 5 showed a conversion of either the *L* or *R* marker, or both (four *R* conversions and one tetrad showing independent conversion of *L* and *R*; supplemental Figure 2 at <http://www.genetics.org/supplemental/>). In four of these five, the crossover position could not be located unambiguously. Thus, crossover position could be assigned in 16 tetrads, and among these were three double-crossover tetrads (one four-strand double and two three-strand doubles). This gave 19 crossovers, of which 15 lay between *L* and *R* (79%), three lay between *ura1* and *L* (16%), and one lay between *R* and *rqh1* (5%; Table 3; supplemental Figure 2 at <http://www.genetics.org/supplemental/>). From this analysis the 4.8-kb region containing *mbs1* contains about three-quarters of the crossovers found in the whole 90-kb *ura1-rqh1* interval. When all crossovers were considered, 83% (19/23) either fell between *L* and *R* or were associated with conversion of *L* or *R*.

TABLE 3

Crossovers in the 90-kb *ura1-rqh1* interval are concentrated in a 4.8-kb region around *mbs1*

Crossover position	Tetrad analysis	Random spore analysis
<i>ura1-L</i>	3 (16%)	22 (21%)
<i>L-R (mbs1)</i>	15 (79%)	57 (54%)
<i>R-rqh1</i>	1 (5%)	26 (25%)
Total	19	105

Crossovers between *ura1* and *rqh1* were identified either among 89 tetrads (supplemental Figure 2 at <http://www.genetics.org/supplemental/>) or among 1200 random spores. In the tetrad analysis crossover events were counted, while in the random spore analysis crossover spores were counted. The positions of the crossovers were localized to one of three regions within the *ura1-rqh1* interval (*ura1-L*, *L-R*, and *R-rqh1*; Figure 4). In the tetrad study only *ura1-rqh1* crossover tetrads without *L* or *R* conversions were scored (to avoid ambiguity in crossover position). Crosses using the strains GP4514 and GP4515 (cross F) were used for both analyses. Two independent crosses, giving similar results, were pooled for the random spore analysis.

The frequency with which *mbs1* conversions accompanied crossovers in this tetrad analysis was similar to that reported earlier. Among the 23 total *ura1-rqh1* crossovers, 11 were associated with conversion of one or more of the central four *mbs1* markers (supplemental Figure 2 at <http://www.genetics.org/supplemental/>), consistent with the earlier tetrad analysis (13 of 28). When all six *mbs1* markers were considered (including *L* and *R*), 13/23 crossovers were associated with a gene conversion (supplemental Figure 2 at <http://www.genetics.org/supplemental/>). This further supports the conclusion that 40% or more of *ura1-rqh1* crossovers are caused by *mbs1* breaks. No significant disparity was observed in the direction of conversion, as would be expected if the single-base-pair mutations did not influence recombination (supplemental Figure 2 at <http://www.genetics.org/supplemental/>).

The analysis of crossover position was extended by carrying out the same cross as described above, but analyzing random spores for *ura1-rqh1* crossovers and then scoring these for crossover position with respect to *L* and *R*. The frequency of *ura1-rqh1* crossovers was 105/1200 (8.8%), close to that seen in previous random spore analyses (see above and YOUNG *et al.* 2002). Compared to the tetrad analysis, a somewhat lower proportion (54%) of *ura1-rqh1* crossovers lay in the 4.8-kb interval defined by the *L* and *R* markers, 21% lay in the *ura1-L* interval, and 25% lay in the *R-rqh1* interval (Table 3). However, the two data sets were not significantly different ($\chi^2 = 4.7$; d.f. = 2; $0.1 > P > 0.05$). Random spore analysis does not allow crossovers with ambiguous locations to be identified and discounted. Such events can happen when a crossover is accompanied by conversion of *L* or *R*. In random spore analysis these events

result in two spores, each with an apparent crossover on opposite sides of the converting marker. This would tend to elevate the frequency of apparent *R-rqh1* crossovers, as these are otherwise rare. This effect may explain the somewhat higher level of *R-rqh1* crossovers in the random spore data compared to the tetrad analysis. Nevertheless, it can be seen that 55–79% of *ura1-rqh1* crossovers are located in a 4.8-kb region around *mbs1* and only 22–45% in the remaining 85 kb of this interval (Table 3).

Association of gene conversion with crossing over at other loci: As discussed previously, gene conversion at *mbs1* was accompanied at high frequency (~80%) by crossing over between *ura1* and *rqh1*, markers flanking the *mbs1* site (see Figure 2C). As this value is considerably higher than that reported for most tested loci in *S. cerevisiae* (WHITEHOUSE 1982; GILBERTSON and STAHL 1996), we investigated whether crossovers were highly associated with conversion at other loci in *S. pombe*.

To examine conversions at loci converting less frequently than *mbs1*, we selected for conversion events as intragenic recombinants. For closely spaced intragenic markers in *ade6*, with separation similar to those involved in our crosses, all observed recombinants arose from gene conversion (GUTZ 1971). Among the intragenic recombinants, the frequency of flanking marker exchange was measured and corrected for incidental events to give the frequency of crossovers associated with the conversion (see MATERIALS AND METHODS). The same correction was carried out to calculate the frequency of crossovers associated with *mbs1* conversion.

As can be seen from Figure 5, at all of the loci examined the crossovers were associated with gene conversion at high frequency (64–87%). The value derived for *ade6* is consistent with published results (GRIMM *et al.* 1994). The high association consistently observed implies bias in *S. pombe* meiotic recombination toward the production of crossovers (see DISCUSSION).

DISCUSSION

We set out to characterize the natural meiotic break hotspot *mbs1* and to analyze the frequency and location of recombination events generated by *mbs1* breaks. This is the first study of a natural meiotic break hotspot in *S. pombe*, rather than one generated by point mutations or large insertions (GUTZ 1971; ZAHN-ZABAL *et al.* 1995). We observed that the *mbs1* locus is a hotspot for meiotic gene conversion and crossing over as well as DNA breakage. The determination of conversion frequencies at *mbs1*, showing it to be a conversion hotspot, allowed us to compare break and conversion rates. This comparison suggested that our converting markers are able to detect a substantial proportion of *mbs1* break-repair events and that at least 30% of meiotic breaks are repaired against homologous chromosomes rather than sister chromatids. The use of multiple markers at *mbs1* in our conver-

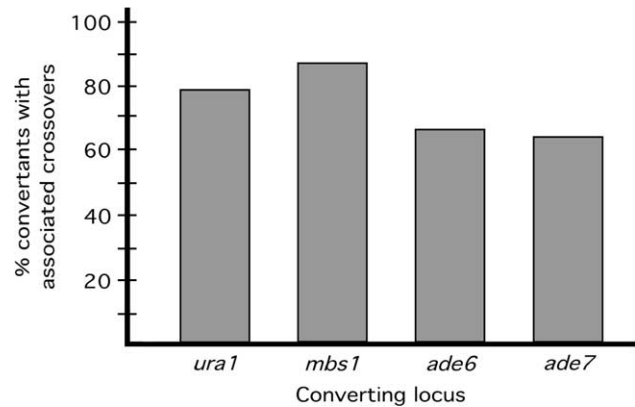


FIGURE 5.—Crossovers are highly associated with gene conversion at four loci in *S. pombe*. Conversion events were identified by tetrad analysis (*mbs1*) or selection for intragenic recombinants (*ura1*, *ade6*, and *ade7*). Crossovers between flanking markers were counted among convertants and among unselected spores, and the frequency of associated crossovers was determined as described in MATERIALS AND METHODS. The *ura1* (cross H, GP789 × GP3696), *ade6* (cross I, GP1252 × GP3785), and *ade7* (cross J, GP5091 × GP5125) crosses were each carried out twice and the results pooled. The *mbs1* data come from tetrad analysis of crosses A and B (Figure 2C) and cross G (GP877 × GP3477) and measurement of crossover frequencies among *mbs1* convertant spores. Uncorrected (observed) crossover frequencies for the flanking markers at each locus (see MATERIALS AND METHODS) were 80% (*mbs1*), 66% (*ura1*), 62% (*ade6*), and 60% (*ade7*). The total number of convertant spore colonies assayed was 30 (*mbs1*), 359 (*ura1*), 1140 (*ade6*), and 359 (*ade7*). The frequency of incidental crossovers, in unselected spores, for each interval (see MATERIALS AND METHODS) was 10% (*mbs1*), 22% (*ura1*), 16% (*ade6*), and 15% (*ade7*).

sion study also allowed us to estimate the size of the conversion tracts, which were often >1 kb in length.

As well as a hotspot for gene conversion, we showed that a 4.8-kb region including *mbs1* is a hotspot for crossovers, which occur in this interval ~10 times more frequently (per kilobase) than the *S. pombe* average. We also observed that crossovers accompany *mbs1* conversions well over 50% of the time, a pattern seen at three other loci, suggesting that recombination is biased to produce crossovers. In contrast to the ability of *mbs1* to stimulate crossovers in the immediate vicinity of the breaks, no substantial stimulation of crossovers >25 kb away was seen. This observation does not support the hypothesis that meiotic DSBs generate crossovers many kilobases away from the location of the breaks, a possibility raised to explain the difference in the distribution of meiotic breaks and crossovers observed in *S. pombe* (YOUNG *et al.* 2002). This leaves the disparity in DSB and crossover distributions unresolved and leads us to suggest that a proportion of meiotic crossovers may result from recombination initiated by single-strand lesions.

More than 30% of *mbs1* breaks can be detected as gene conversions: Our tetrad analysis not only indicated that *mbs1* was a hotspot for gene conversion but also

suggested that the markers we used were able to detect a significant proportion (>30%) of all *mbs1* break-repair events as conversions. Southern blot analysis of meiotic DNA showed that the *mbs1* site undergoes breakage at a frequency of ~10% (Figure 1; YOUNG *et al.* 2002). We observed that 3% of chromatids showed conversion of at least one of the four central *mbs1* restriction markers but there was only one double conversion tetrad (supplemental Figure 1 at <http://www.genetics.org/supplemental/>); *i.e.*, the conversions arose from separate break events. This indicates that 3% of chromatids would need to be broken to generate the observed conversions, representing 30% of all *mbs1* breaks. Some of the remaining 70% of *mbs1* breaks may be repaired using a sister chromatid rather than a homologous chromosome. Therefore, it is likely that our markers detect >30% of inter-homolog DSB repair events at *mbs1*.

A single example was seen of an apparent two-strand double-crossover event within the marker 1–4 constellation (supplementary Figure 1 at <http://www.genetics.org/supplemental/>). This event could have arisen from mismatch repair acting on a region of symmetrical heteroduplex DNA. The existence of this kind of heteroduplex has been inferred in *S. pombe* (FLECK *et al.* 1999). Alternatively, the apparent two-strand double crossover could have arisen from two independent conversions. We conclude that events suggestive of symmetric heteroduplex at *mbs1* do occur, but they are rare.

Size of conversion tracts at *mbs1*: The use of multiple markers at *mbs1* allowed us to estimate the length of conversion tracts in our full tetrad analysis (Figure 2B). Of 16 conversions, 10 (63%) involved two or more markers; of these, 10 were >0.75 kb, 8 were >1.2 kb, and 2 were >2 kb. In addition, 5 of the 6 conversions affecting only one marker involved marker 1 (*i.e.*, the extent of the conversion on one side could not be determined). Analysis of markers at the *ade6* locus also indicated that conversion tracts >1 kb are common (GRIMM *et al.* 1994). These *S. pombe* values are also comparable to an average tract length of 1.5 kb reported for *S. cerevisiae* (BORTS and HABER 1989). Additional *mbs1* conversions including the *L* and *R* markers were seen among the tetrads selected for *ura1-rqh1* crossovers (supplemental Figure 2 at <http://www.genetics.org/supplemental/>). One of these conversion tracts was >3.5 kb (an outside marker converted, allowing no maximum to be set); at *ade6* the longest reported conversion tract was >3.6 kb (including conversion of an outside marker). Gene conversions can arise from either asymmetric or symmetric heteroduplex DNA. According to current models of recombination, asymmetric heteroduplexes correspond to regions of single-strand resection at DSBs, whereas symmetric heteroduplexes occur when strand exchange progresses beyond the resected region of DNA. There is evidence for both symmetric and asymmetric heteroduplex formation during *S. pombe* meiotic recombination (FLECK *et al.* 1999). Therefore, while

longer gene conversion tracts might indicate longer regions of resection, it is difficult to relate the conversion tract and break-resection lengths at *mbs1*.

***mbs1* is a hotspot of crossing over:** Our results indicate that breaks at *mbs1* not only cause crossovers in the 90-kb interval *ura1-rqh1* but also are responsible for at least 40% of all crossovers in this interval. Crossovers in the interval *ura1-rqh1* are greatly elevated among convertants at *mbs1*, compared to the total population. If it is assumed that conversions at *mbs1* indicate repair of *mbs1* breaks, then these breaks generate *ura1-rqh1* crossovers. In addition, 55–79% of *ura1-rqh1* crossovers lie in the 4.8-kb region around *mbs1* (Table 3), and ~40% of *ura1-rqh1* crossovers are associated with a conversion event at *mbs1* (supplemental Figures 1 and 2 at <http://www.genetics.org/supplemental/>). This implies that most *ura1-rqh1* crossovers are caused by *mbs1* breaks and occur close by.

The frequent occurrence of crossovers within a 4.8-kb region around the *mbs1* locus means that *mbs1* forms a hotspot for crossovers (12–18 cM/10 kb) as well as a hotspot for gene conversion. In contrast, the break-free regions to the left of marker *L* and especially to the right of marker *R* are recombinationally much colder (0.7–0.9 cM/10 kb and 0.08–0.4 cM/10 kb). Recombination intensities in these two intervals are also lower than the genome average (1.6 cM/10 kb) or the average of 1.1 cM/10 kb (range 0.6–1.6 cM/kb) reported for the whole *lys3-pro1* region (YOUNG *et al.* 2002). Therefore, in the 90-kb interval *ura1-rqh1* the observed double-strand breakage and the observed crossing over are qualitatively consistent with each other. However, as with analysis of the whole *lys3-pro1* region (YOUNG *et al.* 2002), crossovers still occur in break-free subintervals of *ura1-rqh1*, albeit at lower frequency than the *S. pombe* average.

Crossovers in apparently break-free intervals: In contrast to what was seen for the interval *ura1-rqh1*, our results show no significant enrichment for *lys3/pom1-ura1* crossovers among *mbs1* convertants. Similarly, an artificial break at *mbs1* stimulated frequent *ura1-rqh1* crossing over, but little *lys3-ura1* crossing over (Table 2). It appears that few *mbs1*-initiated recombination events lead to crossovers between *lys3* and *ura1*. This is despite *mbs1* accounting for ~80% of all breakage observed in the interval *lys3-rqh1* and the existence of apparently break-free subintervals that show normal levels of crossovers and for which *mbs1* is the closest observed break site (*e.g.*, *res2-ura1*; YOUNG *et al.* 2002).

Reconstruction experiments demonstrate that we can detect DSB levels lower than that required to explain the crossovers in break-free intervals even if they are spread over many sites rather than concentrated in hotspots (data not shown). We are therefore left with the conclusion that, for these intervals, the closest observed DSBs are not responsible for the observed crossovers but neither is there sufficient double-strand breakage seen within the intervals to account for the crossovers.

It is possible that in crosses to measure recombination genetically DSBs do occur in these intervals, but that these specific DSBs either are absent or do not accumulate under the conditions used to measure DSBs physically. Alternatively, crossovers in these apparently DSB-free intervals might be generated by something other than DSBs. Such initiating events could be single-strand lesions. Single-strand gaps appear to be recombinogenic in bacteria (WANG and SMITH 1984) and recent work suggests that single-strand lesions can be recombinogenic in eukaryotes also. Certain mutant RAG proteins [catalyzing DNA cleavage in V(D)J recombination for mammalian immunoglobulin gene rearrangement] generate nicks rather than DSBs but stimulate homologous recombination even more frequently than do wild-type RAG proteins (LEE *et al.* 2004). Similarly, certain *S. cerevisiae rad52* mutants are proficient for spontaneous mitotic recombination but defective in the repair of DSBs generated by ionizing radiation (MORTENSEN *et al.* 2002). Expression of the site-specific single-strand-nicking protein encoded by ϕ 1 bacteriophage gene II in *S. cerevisiae* also stimulates mitotic recombination (STRATHERN *et al.* 1991). In *S. pombe* the lesion at the *matI* locus that initiates recombination for mating-type switching is a single-strand nick (ARCANGIOLI 1998), although this may be converted into a DSB by replication before recombination (KAYKOV *et al.* 2004). Thus, single-strand lesions may stimulate eukaryotic homologous recombination, although direct evidence is lacking.

High frequency of crossing over associated with gene conversion in *S. pombe*: Our results demonstrate that gene conversions in *S. pombe* are frequently accompanied by crossovers. At the four loci that we investigated, 64–87% of gene conversions had an associated crossover. We assume that interhomolog recombination events that cause gene conversion are typical of all interhomolog recombination events. If so, then recombination in *S. pombe* is biased toward the production of crossovers. This could be explained in two ways. First, there could be a single pathway of recombination that produces crossover and noncrossover outcomes by different resolution of the same Holliday junction intermediates. In *S. pombe* this resolution would have to be biased toward the production of crossovers. Random resolution would lead to equal frequencies of crossovers and non-crossovers. Second, as has been recently suggested, there could be two pathways of recombination, one progressing via resolution of Holliday junctions or related structures and one via SDSA (ALLERS and LICHTEN 2001; CROMIE *et al.* 2001; HUNTER and KLECKNER 2001; OSMAN *et al.* 2003; SMITH *et al.* 2003). The SDSA pathway would produce only noncrossovers and the Holliday junction resolution pathway could give only crossovers. In support of this idea, a mutation in *mus81*, whose product appears to resolve Holliday junctions or related structures in *S. pombe*, does not reduce the frequency

of gene conversions that lack associated crossovers but does greatly reduce the frequency of gene conversions with associated crossovers (OSMAN *et al.* 2003; SMITH *et al.* 2003). The activity of purified Mus81-Eme1 on nicked double Holliday junctions is also consistent with this model (GAILLARD *et al.* 2003; OSMAN *et al.* 2003). The observed bias toward crossovers in *S. pombe* would then suggest that the Holliday junction pathway predominates in that organism.

The observed frequency with which gene conversions have accompanying crossovers in *S. pombe* is higher than that reported for *S. cerevisiae*. The higher frequency of noncrossovers in *S. cerevisiae* may be caused by crossover interference; *i.e.*, any recombination event in the region of interference around a crossover site would be channeled toward a noncrossover. Therefore, in mutants that abolish interference, gene conversions would be more frequently accompanied by crossovers. If so, meiotic recombination in *S. cerevisiae* and other organisms with interference might proceed as in *S. pombe* but with the addition of the interference pathway. This would make *S. pombe* a particularly attractive model organism for studying the fundamental processes of meiotic recombination.

We thank Jürg Kohli and Steve Goldman for *S. pombe* strains; Bernard Dujon for plasmid pSCM525; and Sue Amundsen, Luther Davis, and Andrew Taylor for critical reading of the manuscript. G.A.C. was supported by a long-term European Molecular Biology Organization fellowship (no. ALTF 123-2201). This work was supported by research grant GM31693 from the National Institutes of Health to G.R.S.

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Communicating editor: M. LICHTEN

