

Sites of strong Rec12/Spo11 binding in the fission yeast genome are associated with meiotic recombination and with centromeres

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Abstract Meiotic recombination arises from Rec12/Spo11-dependent formation of DNA double-strand breaks (DSBs) and their subsequent repair. We identified Rec12-binding peaks across the *Schizosaccharomyces pombe* genome using chromatin immunoprecipitation after reversible formaldehyde cross-linking combined with whole-genome DNA microarrays. Strong Rec12 binding coincided with previously identified DSBs at the recombination hotspots *ura4A*, *mbs1*, and *mbs2* and correlated with DSB formation at a new site. In addition, Rec12 binding corresponded to eight novel conversion hotspots and correlated with crossover density in segments of chromosome I. Notably, Rec12 binding inversely correlated with guanine–cytosine (GC) content, contrary to findings in *Saccharomyces cerevisiae*. Although both replication origins and Rec12-binding sites preferred AT-rich gene-free regions, they seemed to exclude each other. We also uncovered a connection between binding sites of Rec12 and meiotic cohesin Rec8. Rec12-

binding peaks lay often within 2.5 kb of a Rec8-binding peak. Rec12 binding showed preference for large intergenic regions and was found to bind preferentially near to genes expressed strongly in meiosis. Surprisingly, Rec12 binding was also detected in centromeric core regions, which raises the intriguing possibility that Rec12 plays additional roles in meiotic chromosome dynamics.

Introduction

Homologous recombination occurs in meiosis at high levels to ensure proper chromosome segregation during the first meiotic division and to provide the basis for genetic diversity. Some regions in the genome show more recombination activity than others and are called hotspots. In *Schizosaccharomyces pombe* and *Saccharomyces cerevisiae*, meiotic recombination at hotspots and elsewhere is initiated by DNA double-strand breaks (DSBs) in one of the two homologous chromosomes (Cervantes et al. 2000; Nicolas et al. 1989), usually at a number of positions across a range of 50 bp (Xu and Petes 1996) up to 2 kb (Cromie et al. 2005).

Chromatin structure can modulate the accessibility of a region for DSB formation, as shown for the histone acetylase mutant *gcn5*Δ at the meiotic recombination hotspot *ade6-M26* in *S. pombe* (Yamada et al. 2004). Specific DNA sequences seem to have a minor role, as long as they are not required for DNA binding factors that in turn render the neighboring region accessible (α -hotspots). Despite high overall sequence homology (98.6%) between the human and chimpanzee genomes, recombination hotspots, inferred by analyzing polymorphisms at orthologous loci and linkage disequilibrium, were mapped at different locations in the two species (Winckler et al. 2005). In

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addition, the recombination frequency in human female autosomes is higher than in male autosomes (Kong et al. 2002; Mohrenweiser et al. 1998). Besides the well studied α -hotspots (Baudat and Nicolas 1997; Steiner et al. 2002), other hotspots exist, where alternative determinants render the chromatin amenable to DSB formation, e.g., nucleosome-excluding sequences (β -hotspots; Kirkpatrick et al. 1999).

Meiotic DSBs are formed by the conserved topoisomerase II-related protein Spo11 (Rec12 in *S. pombe*), which works in a complex with several other proteins (Keeney 2001; Maleki et al. 2007). A Spo11 dimer forms phosphodiester links of its active site tyrosines, Y135, with the two 5' ends of the DSB. The subsequent, staggered endonucleolytic incision (Neale et al. 2005), presumably catalyzed by the Mre11/Rad50/Xrs2 complex and other activities, is inhibited in *rad50S* mutants in both yeasts (Alani et al. 1990; Cervantes et al. 2000; Young et al. 2002). This phenotype was employed in several, genome-wide analyses for Spo11-dependent DSBs in *S. cerevisiae* (Borde et al. 2004; Gerton et al. 2000; Mieczkowski et al. 2006). A prominent correlation of Spo11-dependent DSBs was found with high guanine–cytosine (GC) content. Correlation between recombination density and GC content was also found in the human genetic map (Yu et al. 2001), but not every hotspot displays high GC content (de Massy 2003). In accordance, an inverse correlation between preferred AT-rich meiotic cohesin (Rec8) binding sites and DSB formation was reported in *S. cerevisiae*: Spo11 does not form DSBs in cohesin-binding regions (Glynn et al. 2004; Mieczkowski et al. 2006). Similarly, low Spo11 binding was found across budding yeast centromeres and telomeres and in regions flanking the rDNA gene cluster. The absence of Spo11 matches the hypo-recombinogenic character of these specific regions. The transposition of the centromeric region *CEN3* in *S. cerevisiae* leads to a decrease of gene conversion and crossing over at the novel centromere (Lambie and Roeder 1986, 1988). Although meiotic DSBs inferred from covalent Spo11-DNA intermediates in *rad50S* strains concentrated in large chromosomal regions or were absent therein (Borde et al. 2004; Gerton et al. 2000), recent studies of meiotic DSB site distribution by monitoring resected ssDNA after Spo11 removal in *dmc1*-mutant strains could not completely confirm this specific pattern (Blitzblau et al. 2007; Buhler et al. 2007). DSBs were more homogeneously distributed along the chromosomes and also found in pericentromeric and telomeric regions, where they were formerly reported to be absent.

In this work, we have used the chromatin immunoprecipitation (ChIP) method, firstly, to confirm Rec12 association with a meiotic recombination hotspot and, secondly, to globally map regions in the fission yeast genome where Rec12 binds. We could show that Rec12 binding agrees

with recombination activity and DSB formation at a selection of Rec12-binding sites. Besides providing a map of Rec12-binding sites, we have identified a number of correlations between Rec12-binding sites and other chromosomal features.

Materials and methods

Strains, media, and general genetic methods

Tetrad dissection was described by Baur et al. (2005). Standard minimal medium, yeast extract liquid (YEL) and yeast extract agar (YEA) media, malt extract media, and the general genetic methods were described by Moreno et al. (1991). Synthetic medium (Edinburgh minimal medium [EMM]) with 2% glucose and EMM without nitrogen source (EMM-N) with 1% glucose were described in Nurse (1975). Strains used are listed in Table 1. *Rec12* was tagged at the C terminus with 13 *myc* epitopes at its endogenous chromosomal locus by polymerase chain reaction (PCR)-based gene targeting (Bähler et al. 1998). The resulting strain was wild type with respect to spore viability and intragenic recombination at *ura4A*. The open reading frames (ORFs) of *rec6* and *rec14* were replaced by the hygromycin resistance cassette to create the deletion mutants *rec6* Δ ::*hyg*^R and *rec14* Δ ::*hyg*^R, respectively (Hentges et al. 2005). The deletion strains showed spore viabilities less than 32%, and intragenic recombination at *ura4A* was reduced more than 2,000-fold compared to wild type. Tagging and gene disruption constructs were confirmed by PCR and subsequent sequencing. The tagged Rec12 fusion protein was detected by Western analysis in meiotic extracts of a *pat1-114* strain. Beside the auxotrophies *lys4-8* (-95) and *ade10-13*, the following genetic markers (with respective phenotypes) were used for tetrad analysis: *gpd1-E13* (no growth on YEA+1.6 M sorbitol; Ohmiya et al. 1995), *rec12myc::kanR* (growth on YEA+100 mg/ml G418), *cdc13-117*, *clr6-1*, and *cdc3-6* (no growth on YEA at 37°C). All crosses involved two unlinked, heterozygous markers, except for *sme2* Δ ::*ura4*, where *sme2* Δ ::*ura4 ura4-D18* was crossed with *ura4-D18*.

Chromatin immunoprecipitation

Haploid *pat1-114* strains were grown in YEL at 25°C to saturation (OD₅₉₅>5). The preculture was diluted into EMM and grown at 25°C to OD₅₉₅ of 0.4. Cells were harvested by centrifugation, washed once with one volume water, resuspended in one volume of EMM-N, and kept for at least 16 h at 25°C to arrest cells in G1. NH₄Cl was added to 0.5 g/l and the temperature risen to 34°C to start meiosis. At various time points, 1-ml samples were taken for 4'-6-

Table 1 *S. pombe* strains

Strain	Genotype	Derivation
KLY52	<i>h</i> ⁺ <i>ura4A ura4-D18 rec12_{myc}::kanR pat1-114</i>	This study
KLY59	<i>h</i> ⁻ <i>ura4-D18 rec12_{myc}::kanR</i>	This study
KLY118	<i>h</i> ⁺ <i>ura4A ura4-D18 rad50S pat1-114 rec12_{myc}::kanR</i>	This study
KLY108	<i>h</i> ⁺ <i>ura4A-10 ura4-D18 rad50S pat1-114 rec14Δ::hygR rec12_{myc}::kanR</i>	This study
KLY123	<i>h</i> ⁺ <i>ura4A ura4-D18 rad50S pat1-114 rec6Δ::hygR rec12_{myc}::kanR</i>	This study
KLY288	<i>h</i> ⁺ <i>gpd1-E13</i>	H. Aiba
GP3718	<i>h</i> ⁺ <i>ade6-M3049 end1-458 pat1-114 rad50S</i>	G. Smith
145–5768	<i>h</i> ⁻ <i>sod2::ura4 ura4-D18</i>	Bern
144–5755	<i>h</i> ⁺ <i>lys3-37 ura4-D18</i>	Bern
3–97	<i>h</i> ⁻ <i>lys3-97</i>	Bern
4–127	<i>h</i> ⁺ <i>ura1-61</i>	Bern
3–120	<i>h</i> ⁻ <i>pro1-1</i>	Bern
1–15	<i>h</i> ⁺ <i>ade3-58</i>	Bern
3–93	<i>h</i> ⁻ <i>lys1-131</i>	Bern
2–69	<i>h</i> ⁺ <i>his1-102</i>	Bern
152–6045	<i>h</i> ⁻ <i>cdc1-7</i>	Bern
29–1147	<i>h</i> ⁺ <i>cdc1-7</i>	Bern
3–87	<i>h</i> ⁻ <i>leu2-120</i>	Bern
4–150	<i>h</i> ⁺ <i>lys7-2</i>	Bern
51–2020	<i>h</i> ⁺ <i>cdc3-6</i>	Bern
51–2021	<i>h</i> ⁻ <i>cdc3-6</i>	Bern
29–1149	<i>h</i> ⁻ <i>cdc13-117</i>	Bern
3–99	<i>h</i> ⁻ <i>lys4-95</i>	Bern
3–100	<i>h</i> ⁺ <i>lys4-95</i>	Bern
15–588	<i>h</i> ⁻ <i>lys4-8</i>	Bern
15–589	<i>h</i> ⁺ <i>lys4-8</i>	Bern
27–1045	<i>h</i> ⁺ <i>ade10-13</i>	Bern
27–1046	<i>h</i> ⁻ <i>ade10-13</i>	Bern
154–6151	<i>h</i> ⁻ <i>clr6-1</i>	Bern
156–6226	<i>h</i> ⁺ <i>sme2Δ::ura4+ ura4-D18</i>	Bern
62–2444	<i>h</i> ⁻ <i>ade6-469 ura4-D18</i>	Bern

diamidino-2-phenylindole staining and 50-ml samples for ChIP. Formaldehyde solution (33.3%, Fluka 47629) was added to a final concentration of 1%, followed by incubation for 15 min at room temperature on a wheel. Due to massive meiotic protein degradation, we were not able to recover precipitated DNA from unfixed cells even in the presence of multiple, highly concentrated protein inhibitors. Cells were washed with one volume of cold phosphate-buffered saline (PBS; 130 mM NaCl, 10 mM NaPO₄, pH 7.5) and cell pellets frozen in liquid N₂.

Cells were lysed in 400 μl of lysing buffer [50 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) pH 7.5, 140 mM NaCl, 1 mM ethylenediaminetetraacetic acid (EDTA), 1% (w/v) Triton X-100, 0.1% (w/v) Na desoxycholate, 1 mM Pefabloc SC, 0.5% (v/v) Pefabloc SC Protector Solution, 0.5% (v/v) *n*-octanol, and protease inhibitor cocktail from Roche (one big tablet/25 ml buffer)] with acid-washed glass beads using the FastPrepMachine (BIO101) at level 6.5 for three cycles, 40 s each. Tubes were kept on ice between cycles. Crude extract was collected, and the beads were washed with 300 μl of lysing buffer. Pooled extract and wash was sonicated with a

micro-tip (tapered 1/8 in.) in a Branson digital Sonifier 250 for 40 cycles at 35% amplitude (pulse on: 0.9 s, pulse off: 7 s) to fragment the chromatin to an average size of 650 bp.

To obtain whole cell extracts (WCEs), the samples were centrifuged once for 5 min, and the supernatant (SN) again for 15 min at 13,000 rpm in a cold micro-centrifuge. Five hundred microliters of WCE was precipitated for 3 h at 4°C on a rotation wheel with 30 μl of pre-coated magnetic beads per sample (3 × 10⁷ Dynal M-280 beads, Sheep anti-mouse immunoglobulin G). The beads had been coated with 15 μl monoclonal mouse anti-myc antibodies [*c-myc* (9E10):sc-40, from Santa Cruz] overnight at 4°C in PBS buffer containing 5 mg/ml bovine serum albumin. Using the MPC-E magnet (Dynal), the beads were collected, drained from solution, and washed with 1 ml of the following buffers: twice with lysing buffer, twice with washing buffer A (50 mM HEPES pH 7.5, 500 mM NaCl, 1 mM EDTA, 1% [w/v] Triton X-100, 0.1% [w/v] Na desoxycholate, and protease inhibitors), twice with washing buffer B (10 mM Tris-HCl pH 8.0, 250 mM LiCl, 1 mM EDTA, 0.5% [w/v] NP-40, 0.5% [w/v] Na desoxycholate, and protease inhibitors), and once with TE (Tris-EDTA;

10 mM Tris–HCl pH 8.0, 1 mM EDTA). A 50 μ l elution buffer (50 mM Tris–HCl pH 8.0, 10 mM EDTA, 1% [w/v] SDS) was then added to the drained beads, and the precipitates were released at 65°C during a 10-min incubation with occasional vortexing. After a brief centrifugation at 13,000 rpm at room temperature, the SN was transferred to a fresh tube and 150 μ l TE/SDS (10 mM Tris pH 8.0, 1 mM EDTA, 1% [w/v] SDS) was added (ChIP). Furthermore, 5 μ l of retained WCE was added to 195 μ l TE/SDS (input). After incubating both ChIP and input at 65°C overnight for reversal of cross-linking, proteins were digested with 1 mg/ml Proteinase K (Invitrogen) in TE at 50°C for 2 h. After adding glycogen (final concentration of 0.4 mg/ml), the DNA was purified by two phenol/chloroform/isoamylalcohol extractions and one chloroform/isoamylalcohol extraction in PGL light tubes (Eppendorf). The DNA was finely resuspended in 30 μ l TE containing 0.05 mg/ml Rnase A (Dnase-free, QbioGene) and incubated at 37°C for 30 min. DNA concentration was measured with the NanoDrop device.

Real-time PCR

Real-time PCR was carried out with AmpliTaq Gold® DNA Polymerase (TaqMan® Universal PCR Master Mix, Applied Biosystems). TaqMan probes used were RTKL1 (forward primer at a final concentration of 18 μ M: 5'-CAAAGTTATGGATGCTAGAGTATTTCAAAGC-3', reverse primer at a final concentration of 18 μ M: 5'-TCGACCGCGACTGACAAG-3', labeled primer at final concentration of 5 μ M: 5'-FAM-CTTGGCAATGGGATTTT-NFQ3'). The labeled primer localizes approximately 450 bp distal to the mapped DSB site of *ura4A* (Gregan et al. 2005). PCR steps for the ABI prism 7000 were 2 min at 50°C, 10 min at 95°C, followed by 40 cycles of 15 s at 95°C, and 1 min at 60°C. Duplets of 1–5 μ l (50 ng) of immunoprecipitated or input DNA were analyzed per sample. The amount of template in 1 μ l of input DNA was set to 100%. The precipitated templates in the ChIP samples were given in percent of input.

DNA labeling, microarray hybridization, and data processing

The labeling, hybridization procedures, and data processing were described elsewhere (Heichinger et al. 2006). Two hybridizations were carried out simultaneously on both microarrays (ORF and intergenic); another two hybridizations were probed only on the intergenic microarray. In three hybridization experiments, we labeled 300 to 600 ng immunoprecipitated DNA with Cy5 and similar amounts of input DNA with Cy3. In the fourth experiment, we swapped the dye. The mean data of these four hybridization experi-

ments were used for further calculations. Four independent control experiments with untagged Rec12 repeatedly gave no labeling of the mock-immunoprecipitated DNA sample, whereas the input DNA was labeled fine. Information on probes indicated in the text is available from a searchable Microarray Primer Database (Penkett et al. 2006). All raw data will be available from ArrayExpress. In addition, all processed data will be available from http://www.sanger.ac.uk/PostGenomics/S_pombe/. To identify Rec12 binding peaks, the sliding-window approach of the ChIPOTIE software (version 1.0) was used on the log₂ values of the average of two (ORF probes) or four (intergenic probes) Rec12-binding ratios with a sliding window of 1,000 and 250 bp steps. Prism software version 4.0 was used for calculations of correlation tests.

The transformation of the Affimetrix data set (meiotic cohesin binding, see Ding et al. 2006) to our intergenic and ORF chip architecture was done by location of the Affimetrix oligos by sequence (http://chromosomedynamics.bio.titech.ac.jp/download/pombe_chip/pombe_chip.html) and assigning the Rec8 to input ratio from the 250-bp window given in Ding et al. (2006) for the wild type to every oligo. These values, assigned to the sequence, were then used for calculation of the average over the length of our intergenic and ORF probes.

DSB Southern

From a meiotic *pat1-114 rad50S* time course, 5×10^8 cells were harvested at appropriate time points (T0, T2, T3, T4, T5, T6), washed with 50 mM EDTA pH 8.0, and resuspended in 300 μ l SpheroBuffer (0.4 M EDTA, 10 mM NaN₃, 5 mM dithiothreitol, 1 mg/ml zymolyase [Seikagaku 120493], 5 mg/ml lysing enzyme [Sigma L-1412], and embedded in low melting-point agarose to a final concentration of 1%). Cells in agarose plugs were lysed in 1.8-ml SpheroBuffer at 37°C for 1 h, proteins were digested once in 1.8 ml NDS/PK (0.5 M EDTA, 10 mM Tris pH 7.5, 10 mM NaN₃, 1% Na lauryl sarcosine, 1 mg/ml Proteinase K) at 50°C overnight and again in 1.8 ml SDS/PK (0.5 M EDTA, 10 mM Tris pH 7.5, 0.5% SDS, 1 mg/ml Proteinase K) at 50°C overnight. Plugs were extensively washed with TE containing 1 mM phenylmethylsulfonyl fluoride, exchanging the buffer several times, at room temperature and stored afterward in TE at 4°C.

Genomic DNA embedded in agarose plugs was digested in the appropriate buffers with restriction enzymes according to the supplier (NEB). In particular, in 200 μ l final volume, 100 units of *Cla*I were used at 37°C overnight. Melted agarose plugs were loaded on a 0.8% Tris–borate–EDTA (90 mM Tris base, 90 mM boric acid [pH 8.3], 2 mM Na₂EDTA) agarose gel and run overnight in the cold

room. After ethidium bromide staining, the DNA was blotted on GeneScreenPlus membrane (PerkinElmer) with the VacuGeneXL device (Pharmacia), according to the manufacturer. Radioactive probes were made by random priming of gel-purified PCR fragments with [α - 32 P]dCTP using the Ready-To-Go DNA Labeling Beads (Amersham). Hybridization and membrane washing procedures were done according to the membrane supplier. Parental and DSB bands were detected by a phosphoimager and quantified with the AIDA software (Raytest, Germany). DSB bands were related to the parental band (100%) after background subtraction of the same area in the T0 time sample lane.

Results

Rec12 associates with the meiotic DNA break region at the *ura4A* hotspot

Point mutations in the 5' region of the *ura4A* gene displayed gene conversion frequencies of 18%, whereas in the 3' region, only 6% were detected (Baur et al. 2005). This conversion gradient was explained by initiation of recombination in the 5' upstream region of *ura4A* by DSB formation. A meiotic DSB site has been detected and mapped approximately 500 bp upstream of the *ura4A* translation start codon (Gregan et al. 2005; Sakem 2005).

To find out, whether Rec12 associates with sequences in the DSB formation region of *ura4A*, we employed ChIP with a functional, C-terminal myc-tagged Rec12 in a *rad50S* mutant. The experiments were carried out by induction of meiosis in haploid *pat1-114* mutant strains with the advantage of high synchronicity (Beach et al. 1985). *pat1-114* meiosis shows a chromosome segregation defect in the first meiotic division (Yamamoto and Hiraoka 2003), but has been used systematically for the timing of early meiotic events including DSB formation (Cervantes et al. 2000; Young et al. 2002). In addition, our strains carried the *rad50S* mutation in which DSBs remain un-resected (Young et al. 2002). Rec12myc-DNA complexes fixed by formaldehyde were purified from meiotic cells by immunoprecipitation using an anti-myc antibody. Precipitated DNA was tested for *ura4A* sequences by real time PCR (Fig. 1). Coincident with DSB formation, the association of Rec12 with *ura4A* was found to increase from the time point 0 h (1.6%) to 4 h (9%).

Strains with *rec14* Δ or *rec6* Δ deletions showed a strong decrease in meiotic recombination (Evans et al. 1997), and a *rec14* mutant had the same altered linear elements as a *rec12* mutant (Molnar et al. 2003). No more *ura4A* hotspot sequences were precipitated in *rec14* Δ and *rec6* Δ strains than in a *rec*⁺ strain with untagged Rec12 (Fig. 1b),

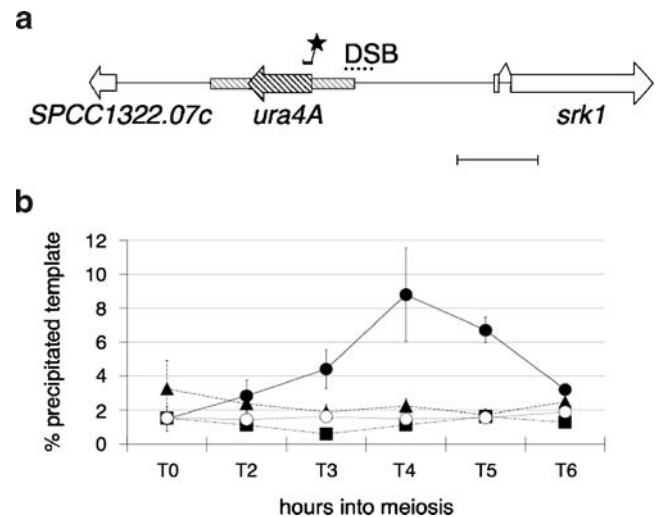


Fig. 1 Rec12 binds to *ura4A* hotspot sequences **a** Map of the *ura4A* insertion used for real-time PCR detection of Rec12-binding sequences. ORFs are given as horizontal arrow boxes. The 1.8-kb *HinD* III *ura4* fragment (Grimm et al. 1988), inserted between SPCC1322.07c and *srk1*, is cross-hatched. Dots indicate the region of DSB formation (Gregan et al. 2005). The location of the *ura4A* TaqMan[®] probe is given by a small bar with an asterisk. The length of the scale bar is 1 kb. **b** Graph showing the ratio of real-time PCR product from immunoprecipitated chromatin versus total chromatin in *rec12myc rad50S* (filled circle), *rec14* Δ *rec12myc rad50S* (filled triangle), *rec6* Δ *rec12myc rad50S* (filled square), and *rad50S rec12+* (open circle) strains, during meiotic progression in a haploid *pat1-114* strain. Independent experiments were performed four times for wild type, three times for *rec14* Δ , and twice for *rec6* Δ . Experiments with the untagged control strain were repeated twice. The y-axis gives the ratio of real time template of immunoprecipitated sample to input sample, normalized to 1- μ l input, which was set to 100% (see “Material and methods”). Error bars indicate the standard error of the mean (SEM)

indicating a requirement of Rec14 and Rec6 for formation of DSBs.

Global analysis of Rec12 binding

The capability of Rec12myc to associate with *ura4A* hotspot sequences prompted us to precipitate Rec12myc-DNA sequences from total chromatin to construct a genomic Rec12-binding map by hybridization to DNA microarrays. The microarrays covered all coding and non-coding regions of the *S. pombe* genome on probes of 180–500 bp (ORF array; Lyne et al. 2003) and 36–2,900 bp (intergenic array; Heichinger et al. 2006). Rec12myc immunoprecipitated DNA as well as input DNA from time point 4 h were differentially labeled and hybridized simultaneously to the microarrays. The Rec12-binding ratio was defined as the coefficient of the amount of immunoprecipitated DNA to input DNA on each probe. Data for the left arm of chromosome II from four independent hybridization experiments are shown in Fig. 2. To single out significant Rec12-binding peaks, the ChIPOTie sliding average algorithm for uniformly arranged data was applied

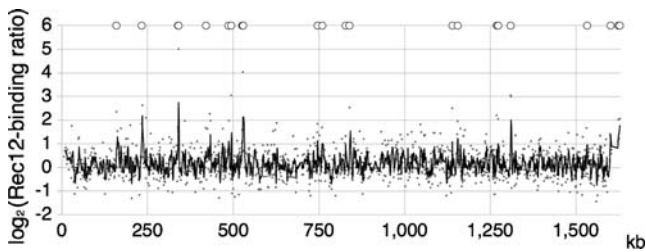


Fig. 2 Mapping of Rec12 binding on the *left arm of chromosome II* on the *x-axis*, the left telomere is given at 0 kb, the centromere at 1,600 kb. Microarray signal ratios of Rec12 enrichment relative to input were plotted for each probe against chromosome position and are given in a \log_2 scale as the mean of 2 (ORF sequences) or 4 (intergenic sequences) independent experiments (*small gray dots*). The *black line* indicates a moving average curve over three values (~ 3.3 kb). The Rec12-binding peaks determined by the CHIPOTie algorithm are indicated above the curve (O)

(Buck et al. 2005). In this way, 144 Rec12-binding peaks were identified genome-wide (Table S1; Fig. 2).

DSB formation near a Rec12-binding peak

To check for DSB formation near a strong Rec12-binding peak, *SPBC1685.13* on chromosome II (Rec12-binding ratio on probe *chr2.187.1>16*, Table S1), *Cla* I digested genomic DNA from time course samples of haploid *pat1-114 rad50S* meiosis was analyzed using Southern hybridization (Fig. 3). Three meiosis-specific bands could be detected at 4 h into meiosis, and the sum was calculated to be 5.6% compared to the parental *Cla* I band. Thus, at sites where meiotic DSBs are formed, high level of Rec12 binding is found on a nearby probe.

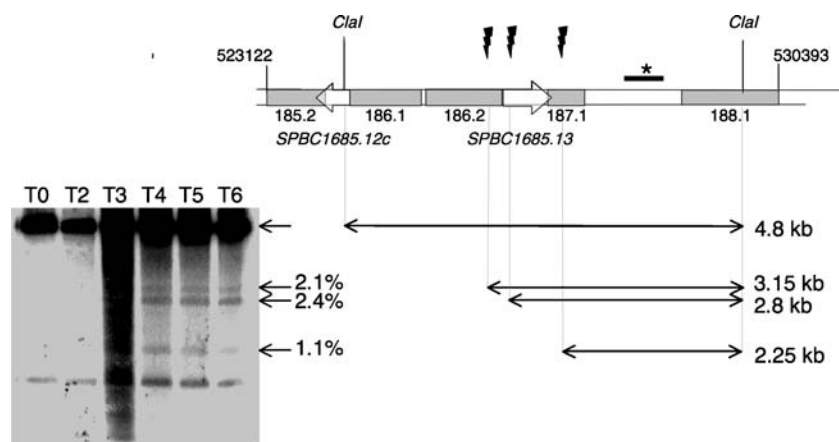


Fig. 3 Meiotic DSBs appear at a Rec12-binding peak. A Southern blot of *Cla* I digested genomic DNA from a *pat1-114* meiotic time course is shown. Above the autoradiogram, hours into meiosis are indicated. The arrows on the right side indicate meiosis-specific DSB fragments and their quantification with respect to the parental *Cla* I fragment. The non-meiosis-specific band below may arise from a cross-hybridization. Such bands were also observed at other loci

Hotspots of recombination localize to sites with high Rec12 binding

To correlate Rec12 binding with recombination activity, probes with various Rec12-binding ratios were checked for nearby genetic markers. Among eight markers with considerably high Rec12-binding ratios on neighboring probes, there were two CHIPOTie Rec12-binding peaks, *chr2.1694.1* and *chr2.103.1* (Table S1). From crosses of various combinations of heteroallelic markers, tetrads were dissected (Table 2) and the spore clones analyzed for the respective phenotypes (“Material and methods”). Gene conversion frequencies ranging from 1% to 16.7% clearly exceeded the average non-Mendelian-segregation (NMS) frequency of 0.28% for all studied genes (Gygax and Thuriaux 1984; Munz 1994), and for six markers, the values were even higher than the value for the well-studied meiotic recombination hotspot *ade6-M26* (5.1%; Gutz 1971).

Rec12 binding and crossover on chromosome I

Genetic distance between two linked markers is a function of the number of crossovers in the interval. As crossover is one way to resolve recombination intermediates initiated at Rec12-dependent DSBs, a positive correlation between Rec12 binding and genetic distance would be expected. We used a set of different markers, which map across chromosome I (Fig. S2), and measured intergenic recombination in the intervals (Table S2). The genetic distance (cM/kb) was compared with the sum of Rec12-binding ratios, which was adjusted to the sum of the probe lengths in the interval. The two sets of data were strongly correlated

(Cromie et al. 2006). The sketch on the right gives the chromosomal location of the three meiosis-specific DSBs on chromosome II (*bolts*). The arrowed (ORF) and gray (intergenic) boxes give the corresponding probes on the microarrays, together with the probe names. The bar with an asterisk indicates the location of the radioactive probe

Table 2 Increased gene conversion frequencies near Rec12-binding sites

Rec12-binding ratio ^a (probe)	Marker	Chromosome	Conversions			Tetrads analyzed	Conversion frequency (%)
			3+:1-	1+:3-	4+:0-		
32.1 (<i>chr2.103.1</i>)	<i>sme2Δ::ura4</i>	II	0	1	0	100	1.0
30.1 (<i>chr2.1694.1</i>)	<i>gpd1-E13</i>	II	11	0	1 ^b	108	12.0
3.6 (<i>chr1.759.1</i>)	<i>rec12_{myc}::kanR</i>	I	13	5	0	108	16.7
3.1 (<i>chr2.139.1</i>)	<i>cdc13-117</i>	II	11	7	0	220	8.2
2.5 (<i>SPBC1105.02c</i>)	<i>lys4-8</i>	II	2	11	0	207	6.3
2.5 (<i>SPBC1105.02c</i>)	<i>lys4-95</i>	II	1	6	0	56	12.5
2.3 (<i>chr2.310.1</i>)	<i>clr6-1</i>	II	3	0	0	37	8.1
2.1 (<i>chr1.1107.1</i>)	<i>cdc3-6</i>	I	8	10	0	141	12.8
1.4 (<i>chr3.475.1</i>)	<i>ade10-13</i>	III	0	2	0	135	1.5

^a Average Rec12-binding ratio of four (intergenic probes) or two (ORF probes) experiments.

^b One tetrad with two conversion events was found and counted twice.

(Pearson coefficient=0.94, p value=0.0002). Figure 4 shows the correlation plot with a simple linear regression fit. The sum of background binding due to unspecific DNA precipitation with the antibody over the long intervals could be responsible for such a positive correlation, although no background binding was detected in control experiments with untagged Rec12 (“Material and methods”). To test whether the correlation depends on background binding, the Rec12-binding ratios were randomly redistributed over chromosome I and added up again in the same intervals (Table S3). Less than 5% of the randomizations showed Pearson correlation p values lower than the observed p value. If the correlation would be a consequence of background binding, more randomizations with p values similar to the observed p value would be expected. We conclude that the amount of Rec12 binding per unit DNA length is not evenly distributed along chromosome I and that high Rec12 binding is associated with increased recombination.

Distribution of Rec12-binding peaks with respect to chromosomes, large intergenic regions, non-protein coding genes, and centromeres

From the 144 CHIPOTIe Rec12-binding peaks (Table S1), 69 were on chromosome I, 44 on chromosome II, and 31 on chromosome III. If the hotspots would be evenly distributed, one would expect less hotspots on chromosomes I and III (64 and 28) but more on chromosome II (52). However, the observed distribution is not significantly different from the expected one ($\chi^2=1.01$, $p=0.6$). Surprisingly, the average Rec12-binding ratio on chromosome II (1.24) is higher than on chromosome I (1.18) and III (1.16, see Table S4). Nevertheless, the ratio of genetic length versus physical length is the same on chromosomes I and II (0.17 cM/kb) and higher on chromosome III (0.22 cM/kb; Munz 1994). The higher average Rec12-binding ratio and the lower numbers of Rec12-binding peaks may reflect more evenly distributed

Rec12 on chromosome II than on the other chromosomes. Extended “cold” or “hot” regions were not detected.

The major meiotic DSB sites *mbs1* and *mbs2* localize to so-called “gene-free regions” (Cromie et al. 2005). These 4- to 8-kb long sequences occur on all three chromosomes (Wood et al. 2002). They display significant deviations from the average nucleotide composition of the whole genome, are sometimes accompanied by repetitions of oligonucleotides (low complexity), and are often associated with $(G-C)/(G+C)$ strand switches (Lobry 1996), where a region with prominent (G) content is followed by a region of similar size with prominent (C) content. Besides the 51 low-complexity gene-free regions, another 21 gene-free regions of similar size lacking low complexity exist. Seventeen out of the 144 CHIPOTIe Rec12-binding peaks fall into gene-free regions (Table S1), among them *mbs2* (measured on probe *chr1.279.2*) with a Rec12-binding ratio of 4. Comparison of the total length of all large gene-free

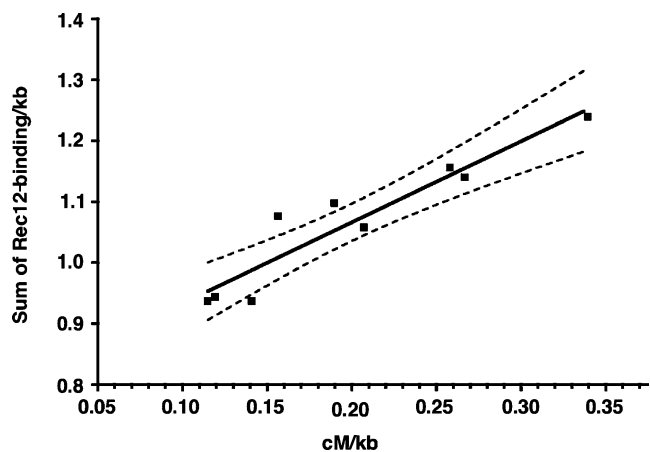


Fig. 4 Rec12 binding correlates with crossovers. Correlation plot of genetic distance in cM/kb measured in intervals along chromosome I (see Table S2) versus the sum of the Rec12-binding ratios in the same intervals divided by the sum of probe lengths. The linear correlation line (solid line) is given together with its 99% confidence limit (dashed lines)

region probes (280.5 kb) with the total length of all measured probes (12,064 kb) leads to an expectation of only 3 hotspots. Thus, Rec12 clearly shows a preference to bind in large gene-free regions. The Rec12-binding ratios on probes within gene-free regions had a higher average value (1.54) compared to the average value of all probes in intergenic regions (1.21, Fig. 5). The effect was most prominent on chromosome I, although two large gene-free regions had low Rec12-binding ratios (data not shown). The same trend was detected on chromosomes II and III, but with overlapping standard errors (Fig. 5).

Thirteen Rec12-binding peaks were found near non-protein coding genes (see Table S1). These genes are transcribed into poly (A)-bearing RNAs but lack long ORFs. Based on the total length of all annotated non-protein coding genes (185 kb), only two Rec12-binding peaks were expected. The average of Rec12 binding to probes covering annotated non-protein coding genes (1.59 ± 0.24) was higher than the average of all probes (1.20 ± 0.01). It has been pointed out, that non-protein coding genes often map to intergenic regions longer than 2.4 kb, and 20% of them are in large gene-free regions (Watanabe et al. 2002). Out of the 17 large gene-free regions with a Rec12-binding peak, only four contain an annotated non-protein coding gene (24%). Thus, the Rec12-binding correlations with large gene-free regions and non-protein coding genes may be independent. Alternatively, there may be so far undetected non-protein coding genes in large gene-free regions. It has been extrapolated that about 300 such genes remain to be discovered (Watanabe et al. 2002).

Recombination across centromere II was reported to be low (Nakaseko et al. 1986). Unexpectedly, strong Rec12 binding was detected at the inner core and inner repeats of all three centromeres (Fig. 6). The ChIPOTIE algorithm captured these sites as Rec12-binding peaks (Table S1).

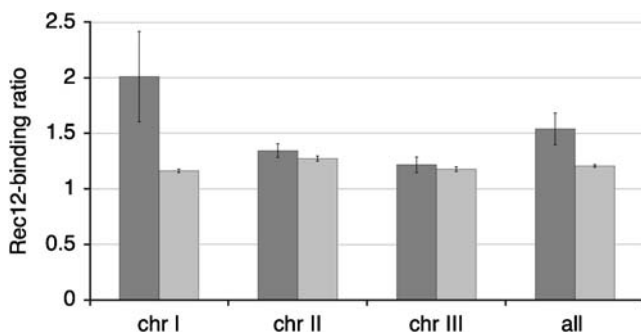


Fig. 5 Rec12 binds to large gene-free regions particularly on chromosome I. The average Rec12-binding ratio for the probes in large gene-free regions (dark bars) versus the average for all probes in intergenic regions (light bars) is shown. The ratios are given for the three chromosomes separately and globally (on the right). The bars indicate the standard error of the mean. Large gene-free regions of the three chromosomes do not vary in length (classes of increasing 1,000 bp steps from 2,000 to 9,000 bp, $\chi^2=20.6$, $p=0.1$ in 95% confidence interval)

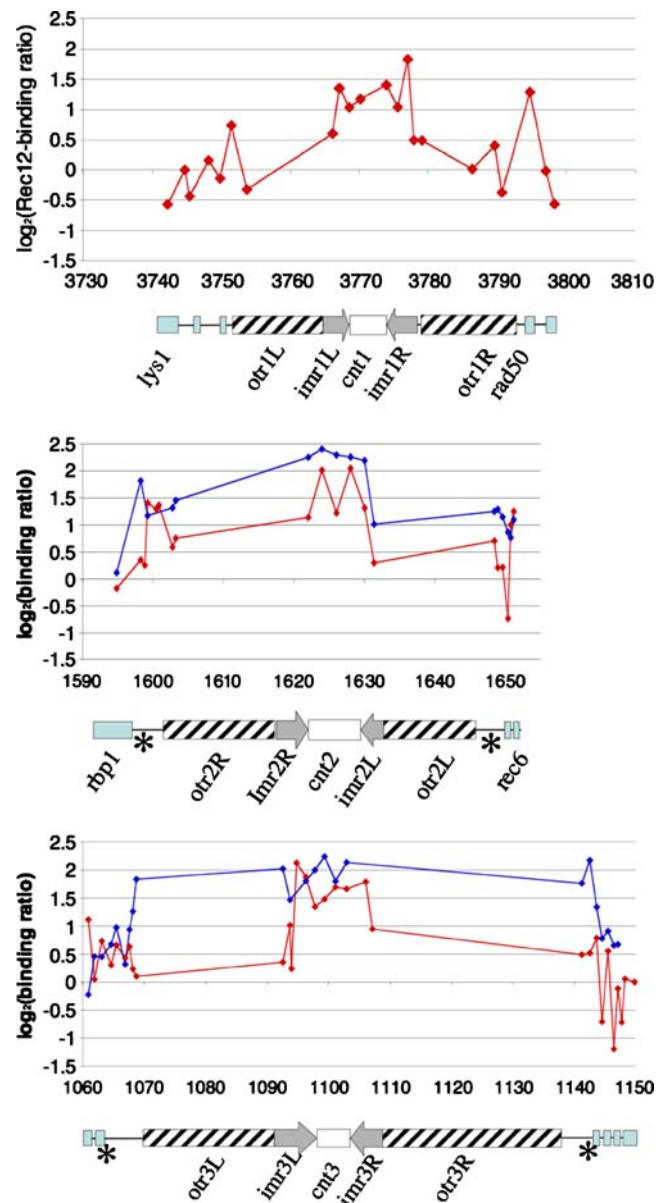


Fig. 6 Rec12- and Rec8 binding in centromere regions. The averaged Rec12-binding ratios (red dots) from four experiments (intergenic probes) or two experiments (ORF probes) and the transformed Rec8-binding ratios (blue dots) from the Hiraoka laboratory (Ding et al. 2006) are plotted against chromosomal location. The units on the x-axis are in kilobase and scaled from the left chromosome ends; the centromere structures are depicted below. Light blue boxes indicate ORF probes, cross-hatched boxes the outer repeats, light gray arrows the inverted inner repeats, asterisks tRNA clusters, and white boxes the core regions. Rec12-binding peaks in centromere I are found in *imr1L* (chr1.1605.1, see Table S1), in *cnt1* (chr1.1605.3), and in *imr1R* (chr1.1607.2); in centromere II in *cnt2* (chr2.651.2 and chr2.651.4); in centromere III in *imr3L* (chr3.535.1), in *cnt3* (chr3.531.1 and chr3.537.2), and in *imr3R* (chr3.538.2)

Inverse correlation of Rec12-binding with GC content and origins of replication

Local high GC content correlates with high Spo11 binding in *S. cerevisiae* on a microarray with ORF probes (Gerton et al. 2000), and the third codon position GC content further increases this correlation (Birdsell 2002). We thus calculated the GC content of every ORF and intergenic probe on the microarrays. A Spearman statistical test showed a significant inverse correlation between Rec12-binding ratio and GC content in both intergenic and ORF sequences (Table 3, Fig. S3a).

The inverse correlation between GC content and Rec12 binding suggests that Rec12 binds preferentially in AT-rich regions. AT-rich islands were proposed to act as origins of replication (Segurado et al. 2003). Early replication forks in meiotic S-phase, stalled by hydroxyurea, which inhibits ribonucleotide reductase and thus depletes the nucleotide pools, were recently mapped on the same ORF and intergenic microarray platforms (Heichinger et al. 2006). A Spearman statistical test revealed a significant inverse correlation between Rec12-binding ratios and meiotic origins of replication (Table 3). Examples of this inverse correlation are shown across a 250-kb region on chromosome II in Fig. S3b.

Rec12-binding correlates with gene expression and cohesin binding

Transcription during meiosis might render chromatin in promoter regions accessible to Rec12. We therefore asked whether genes whose transcription is induced during meiosis were also preferred sites for Rec12 binding. Genome-wide transcription of a *pat1-114* strain at 4 h (Mata et al. 2002) was significantly correlated with the Rec12-binding ratio on the ORF sequences (Table 3). The correlation was most prominent for genes with high induction factors. Out of the ten most strongly induced

genes, seven were also found by the CHIPOTie algorithm as Rec12-binding peaks (*C29A10.02*, *meu10*, *meu3RC*, *dmc1*, *meu4*, *meu19*, and *C22H10.13*).

It was proposed for *S. cerevisiae* that Spo11-binding peaks may represent regions where initiation of recombination was not prevented by cohesin binding (Glynn et al. 2004; Mieczkowski et al. 2006). We therefore compared Rec12 binding with published data on Rec8 binding in *pat1-114* haploid meiosis across chromosome II and a part of chromosome III (Ding et al. 2006). For this purpose, the Rec8-data were transformed to our microarray architecture (see “Material and methods”). Unexpectedly, a significantly positive correlation was evident (Table 3). A 250-kb region of chromosome II is shown in Fig. S3c. Notably, Rec12-binding peaks on chromosome II did not overlap with Rec8-binding peaks on the same probe (Table S1). Moreover, no overlap was detected for probes with a Rec12-binding ratio (\log_2) > 1 (Table S5). But, when the limit was > 0.8, a positive correlation could be detected, suggesting that the Rec8-binding peaks might be at the edges of the Rec12-binding peaks. In fact, of the 43 designated Rec12-binding peaks on chromosome II (excluding peaks in centromere II), 27 had a nearby Rec8-binding peak [Rec8-binding ratio (\log_2) > 0.5] within 2,500 bp on either side. In other words, 62.8% of the Rec12-binding peaks on the arms of chromosome II had a Rec8-binding peak at a distance < 2,500 bp. To test for significance, we calculated how many Rec8-binding peaks fall into random 5,000 bp windows, assuming a Rec8-binding peak width of 3,000 bp and a peak height of 0.5 (Table S6). Under this scenario, only 47.8% of the 5,000 bp windows could expect a hit, less than the 62.8% determined above. The average distance of the 27 Rec12-binding peaks to nearby Rec8-binding peaks was 1.7 kb, with a range of 55 to 2,400 bp (Fig. S4). However, due to different resolutions of the two microarrays (average probe length in Rec12-binding microarrays: 1,100 bp/probe, Rec8-binding microarray: 250 bp/probe), we cannot rule out co-localization for the smaller distances measured. At the centromeres of

Table 3 Correlation of Rec12-binding ratios with specific chromosomal features

Correlation of Rec12 binding with data set, applied probes on microarray (number of pairs analyzed)	<i>p</i> value	Spearman <i>r</i>
GC content, intergenic probes (6020) ^a	<0.0001	−0.10
GC content, ORF probes (4927)	<0.0001	−0.16
GC content, intergenic and ORF probes (10947)	<0.0001	−0.15
Meiotic replication origins, intergenic and ORF probes (8847) ^b	<0.0001	−0.06
Meiotic transcription, ORF probes (4685) ^c	<0.0001	0.09
Rec8 binding, intergenic and ORF probes (5000) ^d	<0.0001	0.18

^a The GC content of each ORF and intergenic probe from the microarray was calculated.

^b Calculated with data from a *pat1-114* strain treated with hydroxyurea at 3 h into meiosis [37]

^c Data from the meiotic mRNA profile (*pat1-114*) at time point 4 h was used [51]

^d Data from two Rec8 CHIP experiments (wild type) on an Affimetrix oligoarray [54] covering the length of chromosome II and part of chromosome III. The data was transformed to the intergenic and ORF array architecture (see “Material and methods”).

chromosome II and III, co-localization of Rec12 binding and Rec8 binding was prominent. Among 20 probes of the inner repeats and the core regions (Fig. 6), we found a high correlation between Rec12- and Rec8 binding (Spearman $r=0.58$, p value <0.0018). No correlation was found in 21 probes of the mating type region.

Discussion

Association of Rec12 with *ura4A* hotspot sequences

To test whether Rec12 associates with DNA sequences of a meiotic recombination hotspot, we checked Rec12 immunoprecipitated DNA from a meiotic time course by real-time PCR to *ura4A* sequences (Fig. 1). Experiments in a *rad50*⁺ strain showed only a slight enrichment of precipitated *ura4A* sequences at 4 h (Fig. S1), which can be explained by removal of Rec12 and subsequent repair of DSBs. In *rad50S* mutant strains, however, the amount of precipitated *ura4A* sequences at 4 h was in the range of the amount of broken DNA (4–9%) at this DSB site measured by Southern blot experiments at 4 h in haploid *pat1-114 rad50S* meiosis (Gregan et al. 2005). The low values at late time points were not expected, as meiotic DSBs persist in *rad50S* mutants. This drop of hotspot sequence recovery may be due to degradation of the Rec12 protein in the extracts. As the immunoprecipitation was done after in vivo formaldehyde cross-linking, the values at 2 and 3 h may indicate that Rec12 already binds before or during DNA replication, although DSB formation is not detected at these times (Gregan et al. 2005). Rec7, the *S. pombe* homolog of the initiation complex member Rec114 in *S. cerevisiae*, and the meiotic cohesin subunit Rec8 are expressed upon nitrogen starvation before the onset of DNA replication (Grishchuk 2003; Molnar et al. 2001; Parisi et al. 1999). In addition, mutants of components of the DSB formation complex advance the start of meiotic S-phase, which suggests a role before DNA synthesis (Doll 2005). Rec12myc was not proficient to precipitate *ura4A* hotspot sequences in the recombination deficient *rec6*Δ or *rec14*Δ mutants. There are no Rec6 sequence homologues in other organisms, but a requirement of the *Sordaria macrospora* Rec14 homologue, Ski8, to localize Spo11 to chromosomes in meiotic prophase was also found (Tesse et al. 2003). In *S. cerevisiae*, however, Spo11 association with hotspot sequences was independent of the Rec14 homolog Ski8 (Prieler et al. 2005).

Rec12-binding peaks and DSB formation

We used a sliding window algorithm to identify peaks of Rec12-binding ratios from the Rec12-Chip on CHIP data, which was not otherwise filtered. The known site of strong

DSB formation *mbs2* (Cromie et al. 2005) was detected as a Rec12-binding peak (probe chr1.279.2, Table S1), but the *mbs1* and *ura4A* hotspots were not among the 144 Rec12-binding peaks due to lack of data from nearby probes. However, at both sites, considerable Rec12 binding was detected (2.2 for probe chr1.316.1 at *mbs1* and 2.5 for probe chr3.40.1 5' to *ura4*). In addition, 8 out of 11 major DSB sites on the 1,105 kb *Not I* fragment of chromosome I (Cromie et al. 2007) coincide with Rec12-binding ratios of more than two, although only two of the CHIPOTIE Rec12-binding peaks were found among them. The criteria set for peak scoring by the CHIPOTIE algorithm were obviously rigorous. We can conclude from this accordance that a good part of Rec12 binding comes from covalent Rec12-DNA interactions.

Rec12 binding correlates with meiotic recombination activity

Repair of meiotic DSBs can engage the sister chromatid or the homolog. If the latter is involved, repair results in a gene conversion event in the vicinity of the DSB site and/or leads to a reciprocal exchange of the involved homologs. As Rec12 is responsible for meiotic DSBs, Rec12-binding peaks are expected to correlate with recombination activity.

High gene conversion frequencies go along with high Rec12 binding, but no quantitative relationship (Table 2) can be deduced from the tetrad analysis results. We neither know the location of DSBs nor the position of the used markers nor the steepness of conversion gradients. However, five of these Rec12-binding sites, *chr2.103.1*, *chr1.759.1*, *chr2.139.1*, *SPBC1105.02c*, and *chr2.310.1*, show considerable Rec12-DNA linkage (Cromie et al. 2007). The remaining two sites have either a low Rec12-binding ratio (*chr3.475.1*) or show a shifted Rec12-DNA linkage peak (*chr2.1694.1*), which might be due to strain differences. The marker *sme2*Δ:*ura4* with the lowest gene conversion frequency but the highest Rec12-binding ratio (32.1) of the eight markers is a peculiar exception. The *sme2* gene encodes a meiosis-essential non-coding RNA, called meiRNA, with whom the Mei2 protein forms a dot-like structure in the nucleus at the *sme2* locus during prophase (Shimada et al. 2003). High Rec12 binding with a considerably low gene conversion frequency at this site may indicate that repair of DSBs involves mostly the sister chromatid at this site. Alternatively, Rec12 binding at *sme2* may serve also for another function than DSB formation (see below paragraph on centromeres).

Genetic distance on chromosome I correlates with Rec12 binding (Fig. 4). Gene conversion events without accompanying crossover (13% to 36% depending on the locus; Cromie et al. 2005) could explain the positive y -intercept of the linear regression line because such conversion events do

not contribute to genetic distance. In addition, Rec12 binding leading to DSB repair by interaction with the sister chromatid may also contribute to the positive y -intercept. The formation of Holliday junction intermediates occurs more frequently between sisters than homologous chromatids in *S. pombe* (Cromie et al. 2006). Such DSBs would be scored in Southern blots, but their repair would not result in crossovers or gene conversions. Overall, a good part of the Rec12 binding reflects recombination activity as observed by correlation with DSB formation, gene conversion, and crossover frequencies. In addition, there are no gross differences between regions with respect to Rec12 binding in *rad50S* strains as it was reported for *S. cerevisiae* Spo11 (Gerton et al. 2000). Recently, alternative studies in *S. cerevisiae* *RAD50* wild-type background indicate, however, that this specific pattern could be attributed to the *rad50S* mutant (Blitzblau et al. 2007; Buhler et al. 2007).

Correlation of strong Rec12 binding with chromosomal features

Previously, meiotic hotspots of DSB formation have been reported in large intergenic regions (Cromie et al. 2005). The high representation of Rec12-binding peaks in these regions confirms this finding. It is not obvious why the Rec12-binding pattern with respect to large gene-free regions differs for chromosome I (Fig. 5). It neither has significantly more large gene-free regions than the other chromosomes ($\chi^2=2.5$, $p=0.3$) nor do they significantly differ in size (see legend to Fig. 5). This finding contrasts the situation in budding yeast and could reflect differences of intergenic region lengths between the two yeast genomes (Wood et al. 2002).

Contrary to the correlation of GC content with covalent Spo11-DNA association in *S. cerevisiae* (Gerton et al. 2000), we find that Rec12 binding shows an inverse correlation to GC content. A similar relationship was reported in *Arabidopsis thaliana* where crossover rates from chromosome 4 inversely correlated with GC content (Drouaud et al. 2006). In addition, in the human MHC class II region, only one out of three crossover clusters corresponded to a region of higher than average GC content (de Massy 2003). Thus, the relationship between GC content and recombination hotspots does not seem to be the same in all eukaryotes. Although replication origins and Rec12-binding sites prefer AT-rich gene-free regions, they exclude each other, which would be consistent with an early loading of the DSB initiation complex before or during S-phase, as discussed above.

Promoters of genes with strong transcription during meiotic prophase are more likely to be targets of Rec12 binding. Whether this correlation is due to specific transcription-factor binding remains to be elucidated.

However, it has been shown that transcription-factor binding can promote recombination activity, but is not sufficient for high recombination at all transcription-factor binding sites in a given region (Kon et al. 1997; Mieczkowski et al. 2006; Steiner and Smith 2005).

Rec12 binding at centromeres

Rec12-binding peaks were detected in all three centromeric core regions (Table S1, Fig. 6). Cromie et al. (2007) showed recently a strong correlation between DSB formation and Rec12 chromatin immunoprecipitated DNA (Rec12 covalently bound to DNA, no formaldehyde cross-linking) in a *rad50S* mutant on a genome-wide scale. They found no Rec12-DNA linkage in the centromeres. Thus, we have to assume that the detected Rec12-binding peaks in the centromere are the result of DSB-independent association. Alternatively, the absence of putative DSBs in the centromere might be an artifact of the *rad50S* mutant. DSBs might be resected (and repaired) through an alternative mechanism, thus escaping the *rad50S* trap. In budding yeast, a role of Spo11 in early chromosome pairing and determination of S-phase length was demonstrated (Cha et al. 2000), independent of DSB and synaptonemal complex formation. Such a dual role was also proposed for Rec12 (Sharif et al. 2002). The *rec12Δ* mutant showed frequent non-disjunction in meiosis I. This phenotype was much milder in the *rec12Y98F* active site mutant, implying that Rec12 has a supporting function on chromosomes segregation in meiosis I, besides its activity in DSB formation. However, contradicting results were reported by Davis and Smith (2003). Additional experiments are required to determine whether Rec12-dependent DSB formation occurs in centromeric core regions or whether Rec12 has another role at the centromeres.

Rec12 binding and cohesin

The correlation between Rec12 binding and Rec8 binding was strong in the centromeric core regions of chromosome II and III (Fig. 6), but was also evident for the remaining regions tested (Table 3). Partially, this correlation could be explained by the mutual preference of AT-rich sequences: A comparison of Rec8 binding and GC content revealed indeed an inverse correlation (Spearman correlation $r=-0.14$, $p<0.0001$). However, the proximity of Rec12-binding peaks to Rec8-binding peaks in more than 60% of all strong Rec12-binding peaks allow us to conclude that a functional relationship between the two complexes may exist. Rec8 could direct Rec12 to susceptible regions for DSB formation. This could explain a long-known riddle: region-specific reduction of recombination in strains carrying mutations in the genes *rec8*, *rec11*, and *rec10*. Specific

point mutations in all three genes, as well as *rec8* and *rec11*, but not *rec10* deletion strains, were found to have strongly reduced conversion and crossover frequencies (up to more than 100-fold) in the central parts of the chromosomes, but only a slight (at most 10-fold) reduction at the chromosome ends (DeVeaux and Smith 1994; Li et al. 1997; Parisi et al. 1999). Regional reduction of DSB frequency in *rec8* Δ and *rec11* Δ (both genes coding for cohesin subunits) has been reported (Ellermeier and Smith 2005). Little reduction of recombination at chromosome ends may be due to the presence of mitotic cohesin complexes there during the early stages of meiosis (Kitajima et al. 2003; Yokobayashi et al. 2003). In mitosis, Rec8 and Rec11 are replaced by their paralogs Rad21 and Psc3 (Kitajima et al. 2003; Yokobayashi et al. 2003). Assistance of mitotic cohesins for the Rec12 complex would then explain the maintenance of higher recombination frequency at chromosome ends in *rec8* and *rec11* mutants. The presence of region-specific recombination reduction in the *rec10-109* point mutant (DeVeaux and Smith 1994), but not in other *rec10* point mutants (Wells et al. 2006), and complete loss of recombination in a *rec10* deletion strain (Ellermeier and Smith 2005; Wells et al. 2006), might indicate that cohesin complexes and Rec12 complexes also interact with the linear element structural protein Rec10 (Lorenz et al. 2006; Molnar et al. 2003) for definition of DSB formation sites.

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