

Cohesins Determine the Attachment Manner of Kinetochores to Spindle Microtubules at Meiosis I in Fission Yeast

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During mitosis, sister kinetochores attach to microtubules that extend to opposite spindle poles (bipolar attachment) and pull the chromatids apart at anaphase (equational segregation). A multisubunit complex called cohesin, including Rad21/Scc1, plays a crucial role in sister chromatid cohesion and equational segregation at mitosis. Meiosis I differs from mitosis in having a reductional pattern of chromosome segregation, in which sister kinetochores are attached to the same spindle (monopolar attachment). During meiosis, Rad21/Scc1 is largely replaced by its meiotic counterpart, Rec8. If Rec8 is inactivated in fission yeast, meiosis I is shifted from reductional to equational division. However, the reason *rec8Δ* cells undergo equational rather than random division has not been clarified; therefore, it has been unclear whether equational segregation is due to a loss of cohesin in general or to a loss of a specific requirement for Rec8. We report here that the equational segregation at meiosis I depends on substitutive Rad21, which relocates to the centromeres if Rec8 is absent. Moreover, we demonstrate that even if sufficient amounts of Rad21 are transferred to the centromeres at meiosis I, thereby establishing cohesion at the centromeres, *rec8Δ* cells never recover monopolar attachment but instead secure bipolar attachment. Thus, Rec8 and Rad21 define monopolar and bipolar attachment, respectively, at meiosis I. We conclude that cohesin is a crucial determinant of the attachment manner of kinetochores to the spindle microtubules at meiosis I in fission yeast.

Sister chromatid cohesion is established during S phase and maintained throughout G₂ until M phase in eukaryotic cells. This cohesion is mediated by a conserved multisubunit complex, cohesin (4, 5, 11, 16, 29). In fission yeast, the cohesin complex in mitosis comprises at least four subunits: Rad21, Psc3, Psm1, and Psm3 (equivalent to Scc1, Scc3, Smc1, and Smc3, respectively, in budding yeast) (25, 26). Chromosome segregation during mitosis is triggered by the dissolution of sister chromatid cohesion along the whole chromosome length.

During meiosis, cohesin subunit Rad21/Scc1 is largely replaced by its meiotic counterpart, Rec8 (9, 18, 19, 30). There are three crucial features of the chromosome at meiosis I that are not observed in mitosis (13, 26, 32). First, pairing and recombination occurs between homologous chromosomes in meiotic prophase, resulting in the formation of chiasmata. Second, sister kinetochores always attach to microtubules from same spindle pole, which is called monopolar attachment and is known to ensure that sister chromatids segregate to the same pole at meiosis I. Third, cohesion between sister chromatids in the vicinity of centromeres persists until the onset of the second meiotic division. Remarkably, all of these features of meiotic chromosomes are disrupted in *Schizosaccharomyces pombe rec8Δ* cells, suggesting that a meiosis-specific cohesin complex plays a central role in the assembly of meiotic chromosomes (12, 30). In particular, because Rec8 preferentially locates at centromeres and *rec8Δ* cells undergo equational chromosome segregation at meiosis I, it has been proposed that Rec8 is required at centromeres to construct kinetochores that orient

to the same pole (30, 31). However, it has been shown in *Saccharomyces cerevisiae* that if the Scc1/Rad21 cohesin subunit, but not Rec8, is expressed by the *REC8* promoter, then monopolar attachment is established, although sister chromatid cohesion at the centromeres is disrupted during anaphase I (27). Thus, at least in *S. cerevisiae*, Scc1/Rad21 can allow proper orientation and monopolar attachment of sister kinetochores at meiosis I. Therefore, there is an argument that the equational segregation observed in *S. pombe rec8Δ* cells may be due to a loss of cohesin in general rather than a specific requirement for the Rec8 subunit of cohesin (10).

Here we address the open question presented above. First, we examined why *S. pombe rec8Δ* cells undergo equational rather than random division. We show that if Rec8 is absent during meiosis, Rad21 relocates to the centromeres to back up the cohesion of sister kinetochores. Thus, substitutive Rad21 is the reason for the occurrence of equational division. Moreover, we demonstrate that even though enough Rad21 is transferred to the centromeres by an ectopic expression, *rec8Δ* cells never undergo reductional division but instead secure equational division. From these results and others we conclude that Rec8 and Rad21 determine the manner of kinetochore attachment to microtubules at meiosis I, at least in fission yeast.

MATERIALS AND METHODS

Media and fission yeast strains. The complete medium YE or minimal medium (SD, MM, or SSA) was used for the routine culture of *S. pombe* strains. Sporulation agar plate (SPA) was used to induce meiosis and microscopic examination. MM and MM-N (lacking nitrogen) containing 1% glucose were used for synchronous meiosis. All strains used in the present study are listed in Table 1.

Construction of a strain carrying the *rad21* allele fused to GFP. A strain carrying the C-terminally green fluorescent protein (GFP)-tagged *rad21*⁺ allele was constructed by direct chromosomal integration of a PCR-generated frag-

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TABLE 1. Fission yeast strains used in this study

Strain	Genotype
JY333	<i>h⁻ ade6-M216 leu1</i>
JY450	<i>h⁹⁰ ade6-M216 leu1</i>
JZ804	<i>h⁺ mei4::ura4⁺ leu1 ura4-D18 ade6-M210</i>
PY182	<i>h⁹⁰ rec8::kan^r ade6-M216 leu1</i>
PY221	<i>h⁺/h⁺ pat1-114/pat1-114 cdc2-L7/cdc2-L7 rec8⁺-GFP-kan^r/rec8⁺-GFP-kan^r ade6-M210/ade6-M216 leu2/+ +/lys1</i>
PY322	<i>h⁺ rec8::kan^r lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 ade6-M216</i>
PY445	<i>h⁹⁰ mei4::ura4⁺ rec8::kan^r rad21-K1<<ura4⁺ lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 ade6-M216</i>
PY446	<i>h⁹⁰ mei4::ura4⁺ rec8::kan^r lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 ade6-M216</i>
PY447	<i>h⁹⁰ mei4::ura4⁺ lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 ade6-M216</i>
PY448	<i>h⁹⁰ mei4::ura4⁺ rad21-K1<<ura4⁺ lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 ade6-M216</i>
PY545	<i>h⁻ rec8::kan^r leu1</i>
PY594	<i>h⁹⁰ mei4::ura4⁺ ade3::lacO-ura4⁺-kan^r his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 lys1 ade6</i>
PY595	<i>h⁹⁰ mei4::ura4⁺ ade3::lacO-ura4⁺-kan^r his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 lys1 ade6</i>
PY741	<i>h⁺/h⁻ mei4::ura4⁺/mei4::ura4⁺ rec8⁺-GFP-kan^r/rec8⁺-GFP-kan^r ade6-M216/ade6-M210</i>
PY748	<i>h⁻ rec8Δ::P_{adh1}-rad21⁺-FLAG ade6-M210</i>
PY749	<i>h⁻ rec8Δ::P_{rec8}-rad21⁺-FLAG ade6-M210</i>
PY754	<i>h⁺/h⁺ pat1-114/pat1-114 cdc2-L7/cdc2-L7 rec8Δ::P_{adh1}-rad21⁺-GFP/rec8Δ::P_{adh1}-rad21⁺-GFP leu2/+ +/lys1 ade6-M216/ade6-M210</i>
PY770	<i>h⁺/h⁺ pat1-114/pat1-114 cdc2-L7/cdc2-L7 rec8Δ::P_{rec8}-rad21⁺-GFP/rec8Δ::P_{rec8}-rad21⁺-GFP leu2/+ +/lys1 ade6-M216/ade6-M210</i>
PY783	<i>h⁻ rad21-K1<<ura4⁺ rec8::kan^r leu1 ade6-M216</i>
PY785	<i>h⁺ rad21-K1<<ura4⁺ rec8::kan^r lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS ade6-M216 leu1</i>
PY786	<i>h⁻ rad21-K1<<ura4⁺ ade6-M216</i>
PY787	<i>h⁺ rad21⁺-GFP-kan^r mei4::ura4⁺ leu1 ura4-D18 ade6-M210</i>
PY801	<i>h⁹⁰ rad21⁺-GFP-kan^r leu1 ade6-M210</i>
PY878	<i>h⁹⁰ rec8::ura4⁺ rad21⁺-GFP-kan^r ade6-M216</i>
PY893	<i>h⁺/h⁺ pat1-114/pat1-114 cdc2-L7/cdc2-L7 rec8::kan^r/rec8::kan^r rad21⁺-GFP-kan^r/rad21⁺-GFP-kan^r leu2/+ +/lys1 ade6-M216/ade6-M210</i>
PZ207	<i>h⁹⁰ rec8Δ::P_{adh1}-rec8⁺-3HA lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 ura4-D18 ade6</i>
PZ208	<i>h⁹⁰ rec8Δ::P_{adh1}-rec8⁺-3HA rad21::ura4⁺ lys1⁺<<lacO his7⁺<<P_{dis1}-GFP-lac1-NLS leu1 ura4-D18 ade6</i>

ment as described previously (1). The PCR products were purified and transformed into JZ804 cells by using the lithium acetate method. Stable G418-resistant (*kan^r* genotype) transformants were selected and analyzed by PCR to verify their correct integration. The obtained *rad21⁺-GFP* cells grew normally, indicating that Rad21-GFP is functional.

Construction of plasmids carrying cyan-GFP variant (CFP)-tagged Gar1 or Mis6 protein. We constructed pREP1-*gar1⁺-CFP* and pREP1-*mis6⁺-CFP*. The *gar1⁺* open reading frame was amplified by PCR with wild-type genomic DNA as a template and forward primer 5'-GCCGGTGCACAATGAGTTTTAGAGGCGGTCCG-3' (the *SalI* site is underlined) and reverse primer 5'-GGGAGGGATCCAAGAATCTGCCACGGAACAC-3' (the *BamHI* site is underlined). The *mis6⁺* open reading frame was amplified with forward primer 5'-CGGCCAGATCTTAATGGAAAGCTTTGAGA-3' (the *BglII* site is underlined) and reverse primer 5'-CCCCGCGGCCGCAAACGATTTAATGTTGA AA-3' (the *NotI* site is underlined). These PCR products were cloned into pGFT81 (7), and then the C-terminal *GFP* was replaced with the cloned *CFP* fragment.

Construction of strains overexpressing Rad21 from the *adh1* or *rec8* promoter. To construct pREP-P_{rec8}-*rad21⁺-GFP*, the *rec8* promoter region (ca. -108 to -1), the *rad21⁺-GFP* fusion gene, the *ura4⁺* cassette, and the 3'-untranslated region of *rec8* (~600 bp) were cloned between the *PstI* and *SacI* sites of pREP1. To construct pREP-P_{adh1}-*rad21⁺-GFP*, an *adh1* promoter fragment was additionally inserted into the 5' site of the *rad21* sequence. These plasmid constructs were linearized by *SacI* and *Clal* to cut out the P_{rec8}-*rad21⁺-GFP* and P_{adh1}-*rad21⁺-GFP* regions and then transformed into *rec8::kan^r* cells to replace the chromosomal *rec8::kan^r* locus with these constructs by homologous recombination at the *rec8* promoter region and 3' untranslated region. For construction of P_{rec8}-*rad21⁺-FLAG* and P_{adh1}-*rad21⁺-FLAG* alleles on the chromosomes, the *GFP* sequences of the preceding constructs were replaced by a *FLAG* fragment.

Preparation of synchronous meiotic cells and observation of chromosomes marked with GFP. Cells were cultured to log phase, collected by centrifugation, suspended in 20 g of leucine liter⁻¹, and then spotted onto a SPA. When we marked only one chromosome by GFP, we cultured opposite mating-type cells, one marked with GFP and the other not, and mixed them prior to spotting them onto SPA. This allows subsequent synchronous conjugation and meiosis. When cells undergoing meiosis I and II or cells arrested at late prophase I by *mei4Δ* became abundant in the population, they were observed without fixation or after being fixed with cold methanol; zygotes were monitored for GFP and DAPI (4',6-diamidino-2-phenylindole). Images were obtained by using a microscope (Axioplan 2; Zeiss) equipped with a cooled charge-coupled device camera

(Quantix; Photometrics) and Metamorph software (Universal Imaging Corporation). Seven Z sections for GFP signals were converted into single two-dimensional images by taking the maximum signal at each pixel position in the images.

Synchronous meiosis. We induced synchronous meiosis with a temperature-sensitive *pat1-114* allele as described previously (31). Cells were cultured in MM-N at 25°C for 16 h and then shifted to 32°C, and 0.1 g of NH₄Cl liter⁻¹ was added. The cells were harvested and fixed with 1% formaldehyde (Wako) for chromatin immunoprecipitation (ChIP) assay and with methanol for microscopy and fluorescence-activated cell-sorting analysis. For microscopy, fixed cells were washed and suspended in PEMS buffer (100 mM PIPES, pH 6.9; 1 mM EGTA; 1 mM MgSO₄; 1.2 M sorbitol) with DAPI.

ChIP analysis. ChIP assays were carried out essentially as described previously (21). Anti-GFP polyclonal antibodies (Living Colors Full-length A.v. Polyclonal Antibody; Clontech) were used for immunoprecipitation. DNA prepared from whole-cell extracts or immunoprecipitated fractions was analyzed by quantitative PCR with a LightCycler and a LightCycler-DNA Master SYBR Green I kit (Roche). The following primers were used for PCR: forward primer for *cnt*, 5'-ATCTCATTTGCTATTTGGCGAC-3'; reverse primer for *cnt*, 5'-GCGTTTC TTCGGCGAAATGC-3'; forward primer for *imr*, 5'-CACATACCAAAAAGT CTGGC-3'; reverse primer for *imr*, 5'-GCTGAGGCTAAGTATCTGTT-3'; forward primer for *dg*, 5'-TTTTTCAGCGAGACATGTACC-3'; reverse primer for *dg*, 5'-TCATAAAGCAACTGGGTG-3'; forward primer for *dh*, 5'-GTAAG TATGAGCAACTGGCG-3'; reverse primer for *dh*, 5'-GGAACAAATCAGG AAACCGAG-3'; forward primer for *lys1*, 5'-ATTTTCGCATCCAACGCTGC-3'; reverse primer for *lys1*, 5'-ACAACAAAGGCTCTGGGCTT-3'; forward primer for *mes1*, 5'-CGTACATTCAGACTGTTGAAC-3'; forward primer for *rDNA1*, 5'-ATGATTCCTCAGTAACGGCGAG-3'; reverse primer for *rDNA1*, 5'-CAC TCTACTTGTTCGCTATCGGT-3'; forward primer for *rDNA2*, 5'-CCTCTAA CGATAGTTACCTGGTTG-3'; reverse primer for *rDNA2*, 5'-CAGAAATTTG AATGAACCATCGCC-3'; forward primer for *TAS*, 5'-CGTTACCAATTA TTATGAC-3'; and reverse primer for *TAS*, 5'-CGGTGATGTAGTGTACTAT A-3'.

RESULTS

Rad21 is dispensable for normal meiosis, as long as Rec8 function is intact. During meiosis, Rad21 is largely replaced by its meiotic counterpart, Rec8, which plays an essential role in

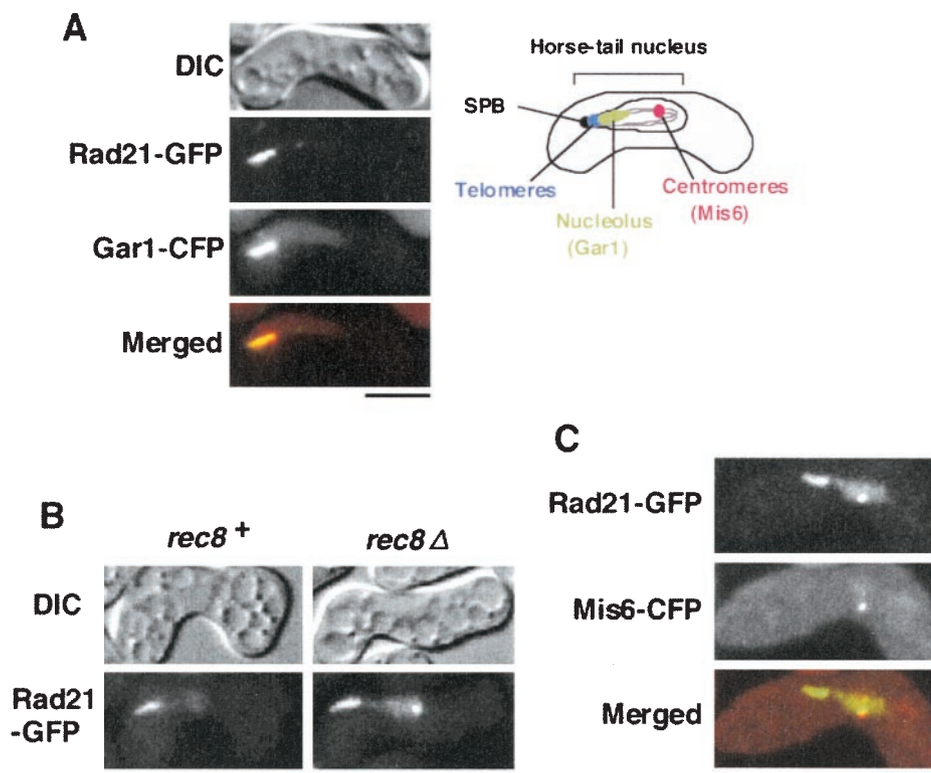


FIG. 1. Rad21 relocates to the centromeres in *rec8Δ* cells. (A) *h⁹⁰ rad21⁺-GFP* cells (PY801) carrying pREP81-*gar1⁺-CFP* during the horsetail period were examined by fluorescence microscopy. In the merged figure, Rad21-GFP and Gar1-CFP are represented by green and red, respectively. The positions of the spindle pole body (SPB), nucleolus, telomere, and centromere are shown in the drawing of the horsetail nucleus. (B) *h⁹⁰ rad21⁺-GFP* cells of *rec8⁺* and *rec8Δ* (PY801 and PY878) were induced to meiosis and examined for GFP fluorescence during the horsetail period. (C) The localization of Rad21-GFP and Mis6-CFP in *h⁹⁰ rad21⁺-GFP rec8Δ* cells (PY878) carrying pREP81-*mis6⁺-CFP*. In the merged figure, Rad21-GFP and Mis6-CFP are represented by green and red, respectively. Bars, 5 μ m.

establishing reductional chromosome segregation at meiosis I. Because Rad21 has been detected in meiotic nuclei at a residual level in several organisms (9, 20, 30), we first sought to determine whether Rad21 plays some role in meiosis in fission yeast. In normal meiosis, Rad21 is detected near the leading edge of the horsetail nucleus, where the telomeres are clustered (30) (Fig. 1A). ChIP assay indicated that Rad21 associates with ribosomal DNA (rDNA) rather than telomere-associated sequences (see below). Moreover, fluorescence microscopy revealed that Rad21 tagged with GFP colocalizes mostly with a nucleolar protein, Gar1 (3), tagged with CFP in the horsetail nucleus (Fig. 1A). These results suggest that Rad21 is enriched in the nucleolus, rather than telomere-adjacent DNA sequences, although residual association with other chromosomal regions might also occur. Given that Rad21 is essential for cell proliferation but *rad21Δ* cells become viable by expressing Rec8 with a constitutive *adh1* promoter (30), we used *rad21Δ P_{adh1}-rec8⁺* cells to inspect the requirement of Rad21 for meiosis. We first assayed meiotic recombination in *rad21*-deleted cells but did not detect any decrease in recombination when tested between the *lys3* and *ura1* loci (located near the telomere of chromosome I) and between the *leu1* and *mat1* loci (located near the centromere of chromosome II) (data not shown). We next examined the requirement of Rad21 for meiotic chromosome segregation. Chromosome segregation was analyzed by viewing the LacI-GFP fusion protein, bound to

LacO DNA repeats that were integrated at the *lys1* locus (*cen1*-GFP dot), ca. 30 kb from the centromere of chromosome I (14). The normal segregation pattern of *cen1*-GFP, a single GFP dot in every nucleus of the tetranucleated cells, was prevalent in the *rad21Δ P_{adh1}-rec8⁺* cells (PZ208) (94%, $n = 543$), as well as in its *rad21⁺* version (PZ207) (92%, $n = 515$). Although we do not know the significance of Rad21 enrichment in the nucleolus, these analyses suggest that Rad21 is virtually dispensable for normal meiotic chromosome segregation and recombination as long as Rec8 function is intact.

Rad21 relocates to the centromeres if Rec8 is absent in meiosis. We have previously shown that *S. pombe rec8Δ* cells undergo mostly equational division at meiosis I. If the Rec8 complex is the sole cohesin in meiosis, *rec8Δ* cells will lose most sister chromatid cohesion during prophase and then undergo random, rather than equational, chromosome segregation. Indeed, the budding yeast *rec8* mutant shows such a phenotype in 70% of cells (9). One explanation for the phenotype of fission yeast *rec8Δ* is that residual Rad21 may substitute for centromeric cohesion, because Rad21 is required at the centromeres for equational segregation in mitosis (2, 17, 28). To investigate this possibility, we determined the localization of Rad21 in *rec8Δ* cells. Although Rad21 is enriched in the nucleolus of *rec8⁺* cells, an additional distinct dot of Rad21 appears in *rec8Δ* cells on the opposite side of the leading edge of the horsetail nucleus (Fig. 1B). To determine the position of the

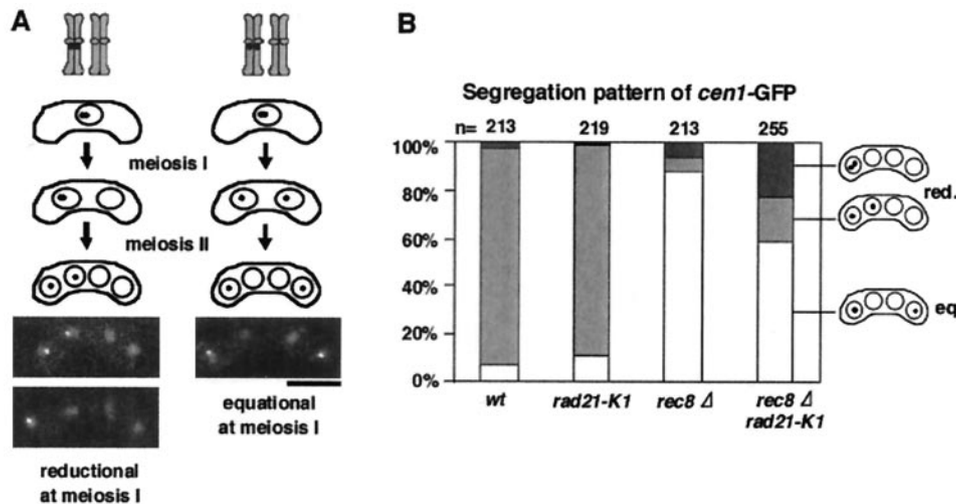


FIG. 2. A mutation in *rad21* impairs equational division at meiosis I in *rec8Δ* cells. (A) The segregation patterns of *cen1*-GFP during meiosis are illustrated with examples. Bar, 5 μ m. (B) Segregation of *cen1*-GFP during meiosis was examined in the indicated cells. A *rec8Δ rad21-K1* strain tagged with *cen1*-GFP (PY785) was crossed with the indicated strain without the tag (wt, JY333; *rad21-K1*, PY786; *rec8Δ*, PY545; *rec8Δ rad21-K1*, PY783) and induced to meiosis at a semipermissive temperature (32°C). Reductional (red.) and equational (eq.) segregation patterns at meiosis I are shown. Note that nondisjunction at meiosis II could be measured only in cells that underwent reductional meiosis I.

Rad21-GFP dot, we coexpressed a kinetochore protein, Mis6 (21), tagged with CFP. Fluorescence microscopy showed the Rad21-GFP dot colocalizing with the Mis6-CFP dot (Fig. 1C), indicating that a significant amount of Rad21 relocates to the centromeres if *Rec8* is depleted. These results raise the possibility that substitutive Rad21 may play a role in ensuring equational chromosome segregation at meiosis I in *rec8Δ* cells.

Rad21 is required for equational division during meiosis I in *rec8Δ* cells. We then decided to introduce a temperature-sensitive mutation, *rad21-K1* (24), into *rec8Δ* and examine chromosome segregation at meiosis I. By crossing the *cen1*-GFP strain with an untagged strain, we monitored the segregation pattern of one pair of sister chromatids during meiosis. Because the two nuclei on either side of the tetrad are derived from the same meiosis I nucleus in fission yeast, reductional or equational segregation in meiosis I can be determined by examining the distribution of the two *cen*-GFP dots in the zygotes (Fig. 2A). Wild-type zygotes represented mostly reductional segregation patterns: $+/+$, $-/-$ (Fig. 2B). The occurrence of some $+/-$, $+/-$ patterns can be explained by recombination between the GFP-associating sequences and the centromere (appearance of recombinants is expected to be 8%). As reported previously (30), *rec8Δ* cells mostly displayed equational segregation patterns: $+/-$, $+/-$ (Fig. 2B). The *rad21-K1* mutation by itself showed no meiotic defect. However, when *rec8Δ* was combined with *rad21-K1*, the equational segregation at meiosis I was partly disrupted with the reductional population increasing to \sim 40% (Fig. 2B). Notably, the *rec8Δ rad21-K1* cells with reductional division at meiosis I underwent random second division (Fig. 2B), suggesting that the “reductional” segregation was erroneous. If equational division is disrupted completely, sister *cen1*-GFP dots will move randomly, producing a complete mixture of equational and reductional patterns (each 50%). Therefore, the observed change would be close to this case. We conclude that Rad21 plays a role in ensuring equational segregation during meiosis I in *rec8Δ* cells.

Rad21 plays a role in cohesion at centromeres during meiosis I in *rec8Δ* cells. For the faithful processing of equational chromosome segregation, centromere cohesion must be preserved at least until the spindle microtubules capture kinetochores. Therefore, centromeric cohesion of *rec8Δ* cells would be intact. This was examined by monitoring *cen1*-GFP marking near the centromere of chromosome I. We utilized a mutation of *mei4Δ* to synchronize the meiotic cells at late prophase I (6). In wild-type cells, the *cen1*-GFP dots were mostly doublet or single at this stage, indicating that sister chromatid cohesion at the centromere was intact and that there was some pairing between homologs (Fig. 3A). The *rad21-K1* cells also displayed no defect in sister chromatid cohesion during meiosis, although there were severe defects during mitosis (25; data not shown). Interestingly, *rec8Δ* cells preserved centromeric cohesion at the *mei4Δ* arrest point in meiotic prophase, although they showed substantially impaired arm cohesion (Fig. 3). Remarkably, the residual centromeric cohesion of *rec8Δ* cells was substantially disrupted by the introduction of the *rad21-K1* mutation. Taken together, these results strongly suggest that if *Rec8* is depleted, Rad21 has a concealed role in preserving sister centromere cohesion at meiosis I, thereby ensuring subsequent equational division.

Sufficient amounts of Rad21 are transferred to the centromeres in meiosis by ectopic expression. In budding yeast, endogenous Scc1 (Rad21 homolog) likely plays little part in centromeric cohesion in *rec8Δ* cells, which undergo rather random chromosome segregation at meiosis I (9). The fact that the *S. pombe* Rad21 complex has an affinity for heterochromatin, which covers a vast region of the centromeres (2, 17), whereas *S. cerevisiae* has no centromeric heterochromatin, might explain why substantial amounts of Rad21/Scc1 relocate to centromeres in fission yeast but not in budding yeast. When Scc1, but not *Rec8*, is expressed by the *REC8* promoter, however, monopolar attachment at meiosis I is mostly established in budding yeast (27). This observation contradicts our current

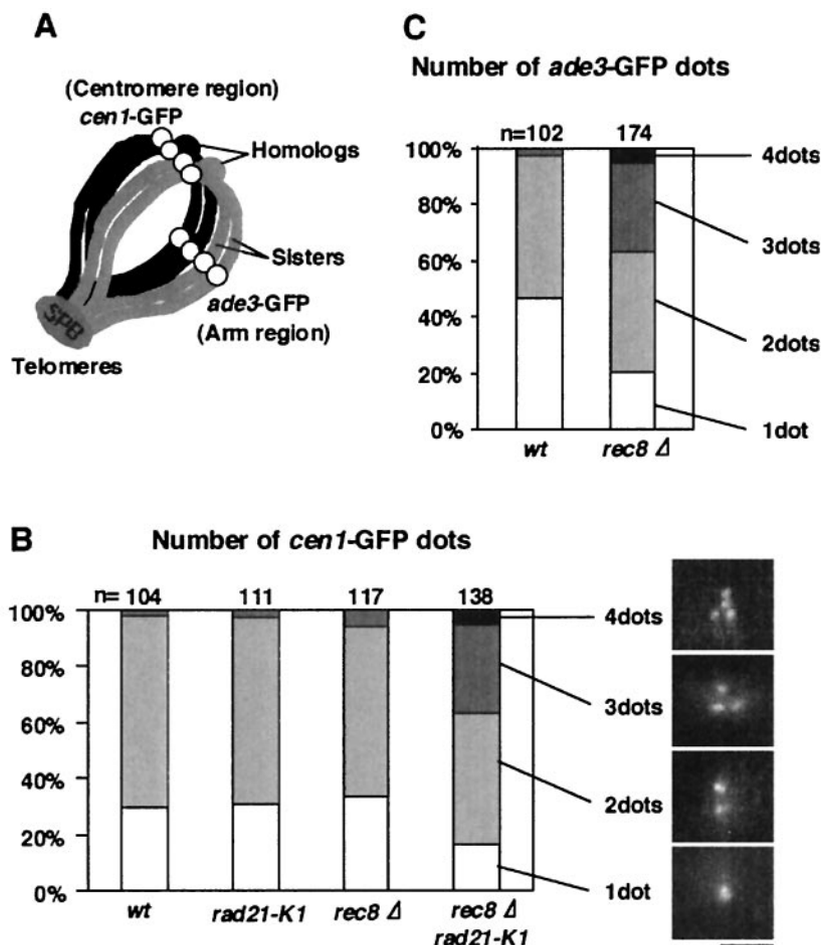


FIG. 3. Sister chromatid cohesion at the centromere is intact in *rec8Δ* cells but impaired by the introduction of a *rad21-K1* mutation. (A) Chromosomes at the horsetail period are depicted. (B) *h⁹⁰ cen1-GFP mei4Δ* cells carrying the indicated mutations (*wt*, PY447; *rad21-K1*, PY448; *rec8Δ*, PY446; *rec8Δ rad21-K1*, PY445) were induced to meiosis at the permissive temperature of 25°C for 8 h and then shifted to 32°C for 4 h. The cells arrested at prophase I (horsetail period) were observed for GFP dots. The number of dots per nucleus is shown with photos of representative nuclei. Bar, 2 μm. (C) Arm cohesion was similarly assayed in wild-type (*wt*) PY594 and *rec8Δ* PY595 cells by observing *ade3-GFP*, which locates in the middle of the left arm of chromosome I.

findings that substitutive Rad21 ensures equational division at meiosis I in fission yeast *rec8Δ* cells. One may argue that the amount of substitutive Rad21 at the centromeres in fission yeast is not sufficient to provide a full level of “cohesin function.” To address this issue, we expressed Rad21-GFP (carrying full function) from the *rec8* promoter or the more robust *adh1* promoter in *rec8Δ* cells and monitored their meioses. Fluorescence microscopy of Rad21-GFP indicated that the centromere signal is indeed stronger in these strains, especially in *rec8Δ::P_{adh1}-rad21⁺-GFP* cells (Fig. 4A). To demonstrate further the association of Rad21 with centromere DNA, we performed a ChIP assay. Extracts were prepared from *rec8Δ::P_{adh1}-rad21⁺-GFP* cells under synchronous meiosis and mostly within prophase I. In accordance with the cytological observations, the ChIP assay indicated that Rad21-GFP associates mainly with the centromere and rDNA sequences (Fig. 4A and B). *S. pombe* centromeres are organized into two distinct domains: a heterochromatic outer repeat region and a central core region associated with histone H3 variant CENP-A (Fig. 4A). Endogenous Rad21 associates with the

outer centromere regions but less with the central core during mitosis (31) (Fig. 4C) and also during meiosis (Fig. 4B), as has been similarly observed in meiotic *rec8Δ::P_{adh1}-rad21⁺-GFP* cells, but the association with chromatin was several times greater in the latter cells (compare Fig. 4A with C). Notably, the increased amount of Rad21 association with the outer centromeric repeat sequences (*dg* or *dh*) exceeds that of endogenous Rec8 association during meiosis (Fig. 4A and D). Similar results were obtained with meiotic *rec8Δ::P_{rec8}-rad21⁺-GFP* cells, although the amount of Rad21 associated with the outer centromere was similar to that of endogenous Rec8 (data not shown).

Rad21 ectopically localized at the centromeres secures bipolar attachment at meiosis I. To examine chromosome segregation in *rad21⁺* overproducers at meiosis I, we monitored the segregation patterns of two centromere-linked markers, *lys1* and *ade6* (16 centimorgans from the centromere of chromosome III), both of which are carried as heterozygous (+/−) in diploids. For this assay, we used a *cdc2* temperature-sensitive mutation that blocks meiosis II so that cells form two

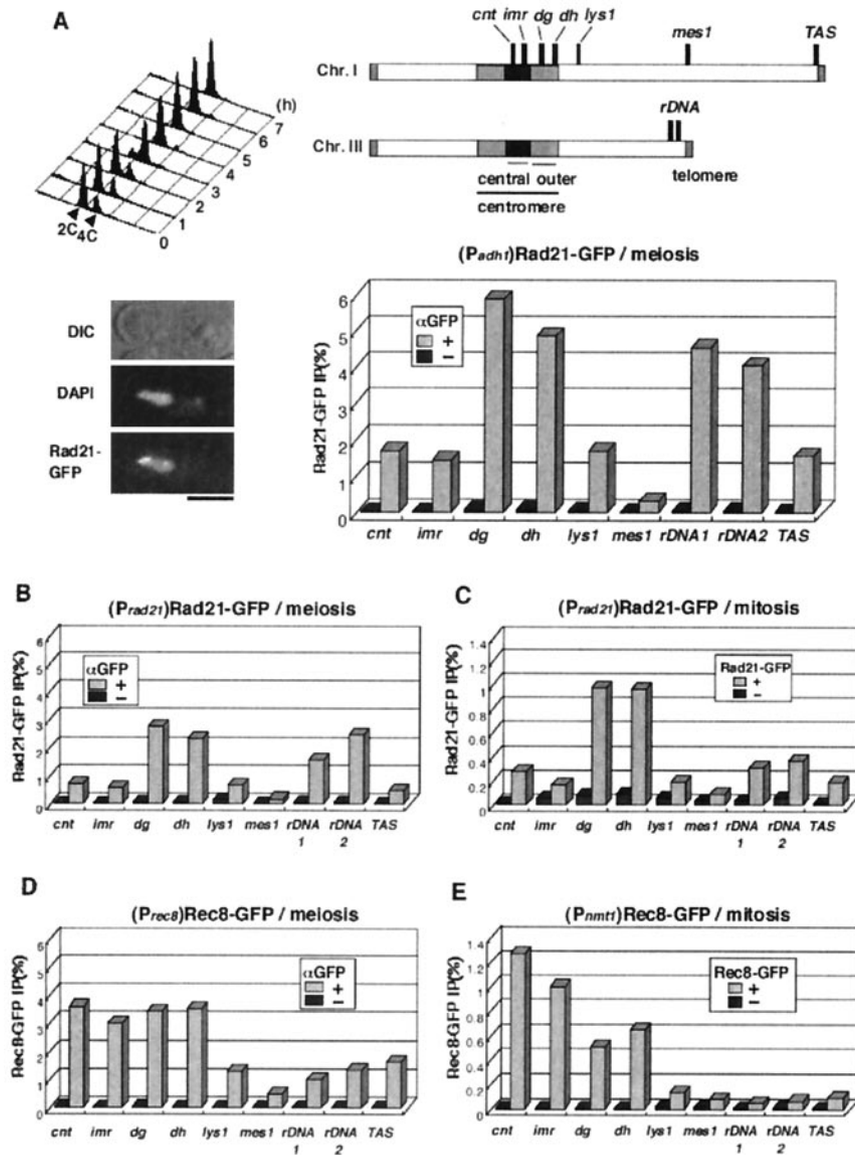


FIG. 4. Rad21 overexpressed in *rec8Δ* cells associates fully with the outer centromere regions in meiosis. (A) Diploid h^+/h^+ *pat1-114 cdc2-L7 rec8Δ::P_{adh1}-rad21⁺-GFP* cells (PY754) were induced to synchronous meiosis and examined for the association of Rad21-GFP at the indicated regions by ChIP assay. Flow cytometric determination of the cellular DNA content indicates that DNA replication was complete at 3 h (full-level association of cohesin was expected to occur at around this time). Therefore, we used this time point for ChIP assay. Microscopic images of the cell at 3 h are shown. Bar, 5 μ m. Schematic representation of *S. pombe* chromosomes I and III, and the primers (*cnt*, *imr*, *dg*, *dh*, *lys1*, *mes1*, *rDNA1*, *rDNA2*, and *TAS*) used for ChIP assay. DNA prepared from whole-cell extracts or immunoprecipitated fractions was analyzed by quantitative PCR. +, sample immunoprecipitated with anti-GFP antibodies; -, negative control without antibodies. (B) ChIP assay was performed in *rec8Δ* cells expressing Rad21-GFP from the native *rad21* promoter (PY893) as described in panel A. (C) Proliferating *rad21⁺-GFP* cells (PY787) were analyzed by ChIP assay with anti-GFP antibodies. We used untagged *rad21⁺* cells (JY333) as a negative control, in this case with anti-GFP antibodies. (D) Meiotic *rec8⁺-GFP* cells (PY741) arrested at prophase by *mei4Δ* were analyzed by ChIP assay with or without anti-GFP antibodies. (E) Proliferating *rec8Δ* cells (PY182) expressing Rec8-GFP from the *nmt1* promoter (pREP81 plasmid) were analyzed by ChIP assay. We used wild-type cells (JY450) as a negative control.

spored dyads after the completion of meiosis I. Therefore, cells undergoing meiotic division in a reductional or equational manner will generate dyads with $+/+$, $-/-$ or $+/-$, $+/-$ genotypes, respectively. By performing random spore analysis, we calculated the percentages of reductional and equational segregation occurring in the cells (Fig. 5A [see the legend for the calculation]). Although wild-type cells showed a reductional segregation pattern (some occurrence of equational

segregation can be explained by recombination between centromeres and GFP-associated sequences), both *rec8Δ::P_{rec8}-rad21⁺-GFP* and *rec8Δ::P_{adh1}-rad21⁺-GFP* cells predominantly displayed an equational pattern (Fig. 5A). Moreover, we monitored meiotic chromosome segregation more directly by observing the segregation pattern of *cen1*-GFP marked on one pair of sister chromatids. The results were consistent; both *rec8Δ::P_{rec8}-rad21⁺-FLAG* and *rec8Δ::P_{adh1}-rad21⁺-FLAG*

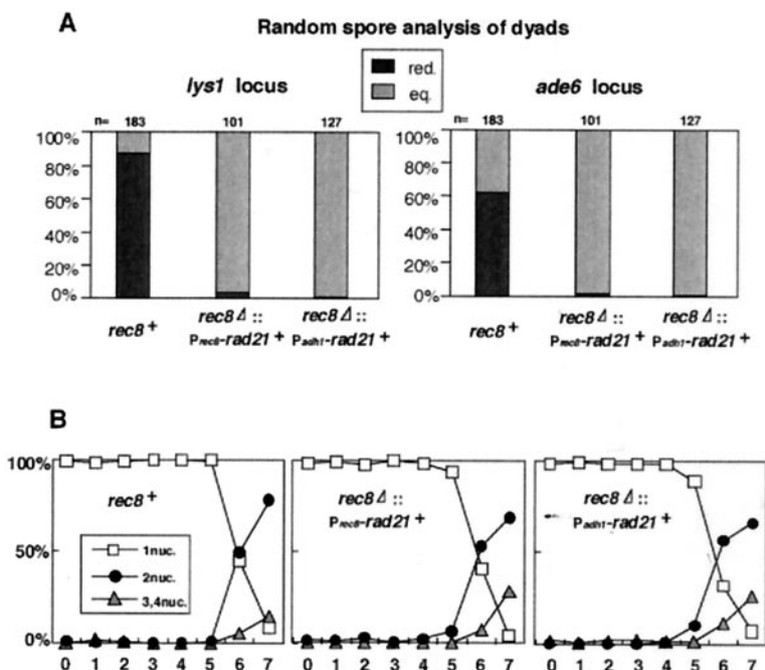


FIG. 5. *rec8Δ* cells ectopically expressing *rad21+* undergo equational chromosome segregation at meiosis I. (A) Diploid h^+/h^+ *pat1-114 cdc2-L7 rec8+*, h^+/h^+ *pat1-114 cdc2-L7 rec8Δ::P_{rec8}-rad21+*-GFP, and h^+/h^+ *pat1-114 cdc2-L7 rec8Δ::P_{adh1}-rad21+*-GFP cells carrying heterozygous centromere-linked markers (*lys1*, *ade6*) (PY221, PY754, and PY770) were induced to synchronous meiosis. Because of the *cdc2-L7* mutation, the cells mostly produced dyads. These were treated with glusulase to obtain free spores, which were plated on YE media containing Magdala Red at 26.5°C for 7 days. Diploid colonies were selected as dark red staining and tested for auxotrophy. Cells undergoing reductional division generate dyads with +/+, -/- genotypes ([+], [-] phenotypes) whereas cells undergoing equational division generate dyads with +/-, +/- genotypes ([+], [+], [-], [-] phenotypes). The number of reductional divisions was calculated by $n[\text{red.}] = n[+] - n[-]$ and that of equational divisions was calculated by $n[\text{eq.}] = (n[+] + n[-])/2$. (B) The nuclear divisions during synchronous meiosis in panel A were monitored by DAPI staining. Note that only some cells underwent meiosis II because of the *cdc2-L7* mutation.

cells underwent equational division at meiosis I (Fig. 6), as in the case of *rec8Δ* cells (Fig. 2B). These results indicate that even if sufficient amounts of Rad21 are loaded into centromeres in meiosis I, *rec8Δ* cells never reestablish monopolar attachment but instead establish bipolar attachment. Finally, we examined the kinetics of nuclear division of *rec8+*, *rec8Δ::P_{rec8}-rad21+*-FLAG, and *rec8Δ::P_{adh1}-rad21+*-FLAG cells in synchronous meiosis. We found that the first and second meiotic divisions progressed with similar kinetics in all

strains (Fig. 5B). Given that the *rec8Δ rad21+* overproducers undergo equational segregation at the first meiotic division, these results imply that the cohesion between sister kinetochores established by Rad21 is not protected during meiosis I, in contrast to the cohesion mediated by Rec8. Taken together, these findings led us to conclude that Rec8 bears specific roles at meiosis I in establishing monopolar attachment and also in maintaining cohesion between sister centromeres and that Rad21 cannot substitute for either of these functions even if it is fully localized to the centromeres during meiosis.

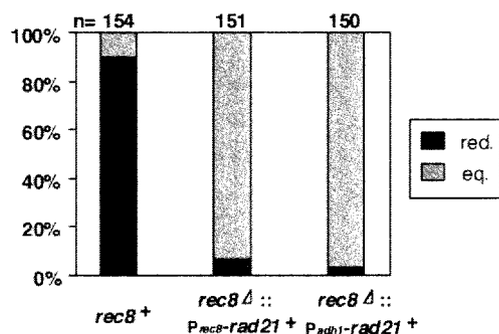


FIG. 6. The segregation pattern of *cen1*-GFP during meiosis I was examined in *rec8+*, *rec8Δ::P_{rec8}-rad21+*-FLAG, and *rec8Δ::P_{adh1}-rad21+*-FLAG cells. A *rec8Δ* strain tagged with *cen1*-GFP (PY322) was crossed with the indicated strain without tag (*rec8+*; *rec8Δ::P_{rec8}-rad21+*-FLAG, PY749; *rec8Δ::P_{adh1}-rad21+*-FLAG, PY748), and the zygotes that formed were analyzed as in Fig. 2A.

DISCUSSION

In the present study, we illuminated the inherent difference of the kinetochore roles of two cohesin subunits, Rad21 and Rec8, the former mitotic and the latter meiotic. Regarding the centromere location of cohesin, large amounts of Rec8 locate at both the outer and the central regions of the centromere in meiosis; however, whereas much Rad21 is located at the outer region, less is present at the central core in mitosis (31) (Fig. 4C and D). We found here that ectopically expressed Rad21 in meiosis displays a location pattern similar to that seen in mitosis, especially at the centromere (Fig. 4A). In excellent contrast, Rec8 ectopically expressed in mitosis displays a location pattern similar to that seen in meiosis (Fig. 4D and E). Thus, the centromeric location pattern of Rec8 or Rad21 does not depend on the developmental state, whether mitotic or mei-

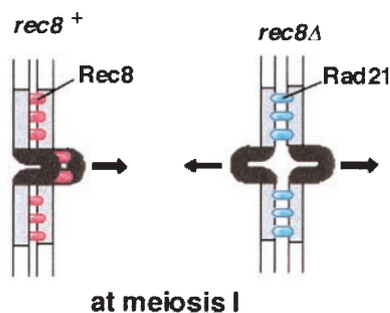


FIG. 7. Model for the regulation of kinetochore orientation by cohesin Rec8 and Rad21 during meiosis I. Rec8 interacts with both the central core and outer repeat regions of the centromere, leading to the monopolar orientation of sister kinetochores at meiosis I. Rad21 associates less with the central core but more with outer repeat regions, thereby ensuring sister kinetochore cohesion and bipolar attachment at meiosis I in *rec8Δ* cells.

otic, but is determined by intrinsic properties of the Rec8 and Rad21 molecules. Central core regions of the centromere are likely to assume an inverted configuration, protruding from other regions of the centromeres, and would provide the site for microtubule attachment (15, 23). Based on the distinct localization pattern of Rec8 and Rad21 at the centromeres, we speculate that only Rec8 may possess the competency to establish cohesion of the sequences along the central region, which would lead to the mono-oriented configuration of sister kinetochores (31). This model fits well with the present finding that, even though much Rad21 locates at the centromeres (especially at the outer regions) and establishes centromeric cohesion, it cannot establish the monopolar orientation of sister kinetochores but instead establishes bipolar attachment (Fig. 7). Thus, Rec8 and Rad21 are apparently the determinants for the orientation of sister kinetochores during meiosis I in fission yeast. The forced expression of Rec8 during mitosis, however, does not lead to the full occurrence of reductional chromosome segregation by itself (30). This inability to establish monopolar attachment in mitosis is probably not due to the lack of Rec8 at the central core regions of the centromere (Fig. 4E), although we do not exclude the possibility that the amount of Rec8 at centromere during mitosis is not sufficient to fulfill its function. Moreover, even in the normal meiotic cell cycle, residual Rec8 at the centromere after meiosis I does not promote monopolar attachment any more at meiosis II. These facts rationalize the assumption of a meiosis I-specific factor that must cooperate with Rec8, but not Rad21, at the kinetochore to establish monopolar attachment.

That Rec8 can support monopolar attachment during meiosis I is not the only meiosis-specific centromere feature that Rad21 cannot support. Rec8 is removed from chromosome in a two-step manner—from chromosome arms in meiosis I but from centromeres only in meiosis II—thus preventing the sister chromatids separating in meiosis I. Rad21, however, is apparently removed from all along the chromosome arms at meiosis I because *rec8Δ::P_{rec8}-rad21⁺* cells undergo the first meiotic divisions with the same kinetics as wild-type cells. In budding yeast *rec8Δ::P_{rec8}-Sccl⁺/rad21⁺* cells, cohesion between sister centromeres mediated by *Sccl*/Rad21 is also destroyed at the onset of the first meiotic division (27). Thus, the

instability of Rad21/*Sccl* regarding this aspect of meiosis I is conserved between budding and fission yeast.

Earlier work in budding yeast identified a meiosis-specific kinetochore factor, Mam1, that functions in establishing monopolar attachment, and this role can be executed only in the presence of cohesin (either Rec8 or *Sccl* complexes) and only during meiosis (27). Another meiotic factor, Spo13, is also involved in the regulation of monopolar attachment in budding yeast (8, 22). So far, Mam1 or Spo13 homologs have not been found in other organisms, including fission yeast, but it may be a conserved feature that a meiosis I-specific factor(s) cooperates with cohesin to establish monopolar attachment at meiosis I. The definitive difference between the two yeasts is that cohesin specificity plays a crucial role in establishing monopolar attachment in fission yeast but not in budding yeast. Studies of the distribution of Rec8 at centromeres in comparison with that of Rad21/*Sccl* in other organisms will be interesting, as will further examinations to determine whether cohesins are determinants of monopolar or bipolar attachment at meiosis I in other eukaryotes.

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