Destabilization of the replication fork protection complex disrupts meiotic chromosome segregation

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ABSTRACT The replication fork protection complex (FPC) coordinates multiple processes that are crucial for unimpeded passage of the replisome through various barriers and difficult to replicate areas of the genome. We examine the function of Swi1 and Swi3, fission yeast’s primary FPC components, to elucidate how replication fork stability contributes to DNA integrity in meiosis. We report that destabilization of the FPC results in reduced spore viability, delayed replication, changes in recombination, and chromosome missegregation in meiosis I and meiosis II. These phenotypes are linked to accumulation and persistence of DNA damage markers in meiosis and to problems with cohesion stability at the centromere. These findings reveal an important connection between meiotic replication fork stability and chromosome segregation, two processes with major implications to human reproductive health.

INTRODUCTION Meiosis is a conserved cell differentiation process that reduces ploidy and creates genetic diversity through recombination in sexually reproducing organisms. In contrast to mitosis, wherein progenitor and daughter cells carry the same genetic information, meiosis produces daughters with half the number of parental chromosomes. This is accomplished by two consecutive nuclear divisions that follow one round of DNA replication (reviewed in Zickler and Kleckner, 1999; Cavalier-Smith, 2002; Hochwagen, 2008; Ohkura, 2015). Another characteristic that distinguishes meiosis from mitosis is the requirement for programmed double-strand breaks (prDSBs) following DNA synthesis. These are repaired by recombination between homologous chromosomes. The physical connections established in this process are vital for reciprocal exchange of genetic information and for proper chromosome segregation in most organisms (Keeney et al., 1997; Sharif et al., 2002; Davis and Smith, 2003). Lack of prDSBs is associated with chromosome nondisjunction, which contributes to different forms of aneuploidy (Hirose et al., 2011; Sakuno et al., 2011).

Mono-orientation of sister kinetochores in the reductional meiosis I division (MI) is another key feature of meiosis (Kitajima et al., 2003a; Brar et al., 2006; Sakuno et al., 2009, 2011). Separation of homologues to opposite poles of the cell after anaphase I requires that sister kinetochores are mono-oriented, so that they migrate in the same direction on the meiotic spindle. At the onset of anaphase II, as is the case in mitosis, bioriented kinetochores enable sister chromatids to segregate to opposite poles (Yokobayashi and Watanabe, 2005; Hauf et al., 2007; Sakuno et al., 2009; Kim et al., 2015).

Finally, regulation of sister chromatid cohesion is essential for the different chromosome segregation pathways. Gradual loss of cohesion, first on chromosome arms and then at the centromere, facilitates the onset of MI and MII divisions, respectively (Kitajima et al., 2003a; Rabitsch et al., 2003a; Brar et al., 2006). Before MI, the meiosis-specific χ-kleisin cohesin subunit Rec8 is targeted for degradation by the endopeptidase separase everywhere along the chromosome except the centromere, where Rec8 is protected by shugoshin-PP2A (Kitajima et al., 2004; Marston et al., 2004; Ishiguro et al., 2010). This stepwise process promotes chiasma resolution (necessary for the separation of homologues in MI), while keeping sister chromatids tethered at the centromere until MII (required for reductional division). When cells near MII, Rec8 is entirely removed.
from the centromeres, allowing the spindle to pull sister chromatids toward opposite poles (Kitajima et al., 2003a; Brar et al., 2006; Kats et al., 2010).

Replication instability, characterized by the dissociation of replication components that normally ensure efficient resumption of replication following fork stalling or DNA repair (Errico and Costanzo, 2012), is deleterious to genomic integrity and is associated with cancer in humans (Durkin et al., 2008; Allera-Moreau et al., 2012; Coschi et al., 2014; van der Crabben et al., 2016). When it occurs in meiosis, it can reduce fitness of gametes and negatively affect reproductive health (Baudat et al., 2000; Nudell et al., 2000; Baarends et al., 2001; Gu et al., 2007). Although the effects of abnormal replication have been widely studied in mitosis, less is known about the consequences it has in meiosis. Recent work on replication stability genes in fission and budding yeast, however, has shown a link between abnormal replication and meiotic defects (Dolan et al., 2010; Le et al., 2013; Mastro and Forsburg, 2014; Murakami and Keeney, 2014; Wu and Nurse, 2014).

The replication fork protection complex (FPC) is crucial for stabilizing the replisome during replication in vegetative cells and is particularly important in difficult to replicate areas of the genome or during replication stress (reviewed in Leman and Noguchi, 2012). The FPC is also necessary for preventing fork collapse when replication is arrested or when the replisome encounters severe DNA lesions (Noguchi et al., 2003; Shimamoto et al., 2009). The FPC’s primary components are Swi1/Tof1 (human Tipin) and Swi3/Csm3 (human Tipin), which form a heterodimeric complex with interdependent regulation (Noguchi et al., 2004; Chou and Elledge, 2006; Gotter et al., 2007; Ünsal-Kaçmaz et al., 2007).

Although Tof1 (Swi1’s orthologue) was initially identified through its role in fission yeast mating-type switching (Egel et al., 1984), work on budding yeast found that it interacts with topoisomerase 1, suggesting an additional function of the FPC in safeguarding genome integrity (Park and Stenglantz 1999). Further examination of the FPC components in various organisms, including humans, revealed the FPC’s involvement in activating the replication and DNA damage checkpoints upon exposure to genotoxic agents (Murakami and Okayama, 1995; Foss, 2001; Noguchi et al., 2003, 2004; Chou and Elledge, 2006; Gotter et al., 2007; Ünsal-Kaçmaz et al., 2007; Yoshizawa-Sugata and Masai, 2007; Smith et al., 2009; Leman et al., 2010).

Destabilization of the FPC by removing either of its components generates substantial DNA damage. Such damage is thought to arise from the uncoupling of the MCM helicase and the DNA polymerases, which generates large ssDNA gaps that are prone to de-etherios rearrangements and physical breakage (Noguchi et al., 2003, 2004; Tournière et al., 2005; Ünsal-Kaçmaz et al., 2007; Dolan et al., 2010). These observations are consistent with a model in which the FPC acts as a sensor of DNA damage (reviewed in Leman and Noguchi, 2012). In this view, Swi1 stabilizes Swi3 at the fork (Chou and Elledge, 2006; Yoshizawa-Sugata and Masai, 2007). Swi3 senses RPA bound to excess ssDNA (Witosch et al., 2014) and communicates with Mrcl (Claspin), which relays the message to checkpoint effector kinases (Kumagai and Dunphy, 2000; Chini and Chen, 2003). This process allows cells to maintain genome integrity by properly addressing sources of replication instability and DNA damage.

Significantly, the FPC also functions in the regulation of sister chromatid cohesion, a process intimately linked to DNA replication (Ansbach et al., 2008; Leman et al., 2010). In both yeasts, the FPC genetically interacts or indirectly affects Chl1 (the yeast orthologue of human Chlr1), a helicase important for cohesion (Mayer et al., 2001, 2004; Warren et al., 2004; Xu et al., 2007; Ansbach et al., 2008). Loss of FPC components in frog and fission yeast results in dysfunctional chromosome pairing and premature sister chromatid separation, respectively (Errico et al., 2009; Tanaka et al., 2009). Human cell lines depleted of individual FPC components also show cohesion defects, which is consistent with the observation that the FPC physically interacts with Smc1/3 (Leman et al., 2010).

In this work, we examine how replication fork stability facilitates the proper execution of meiosis. We used cells lacking Swi1 and Swi3 (the FPC components) and examined the outcome of their meioses by various means. We observed that fork destabilization did not halt meiotic progression but resulted in defects in recombination. Moreover, lack of each FPC component resulted in abnormal chromosome segregation in both meiotic divisions, which may explain the substantial reduction in spore viability in these strains. These observations support a model in which the FPC contributes to genome stability in meiosis by suppressing DNA damage and promoting proper centromeric cohesion.

RESULTS

FPC mutants have reduced spore viability

In fission yeast, meiosis is coupled to sporulation (McLeod et al., 1987). Thus defective spores indicate underlying meiotic defects. We crossed heterothallic swi1Δ and swi3Δ strains (FPC mutants) to examine their relative spore viability by random spore analyses. Compared with wild-type cells, the FPC mutants showed a substantial drop in spore viability (Figure 1A). These reductions were similar to those of the swi1Δ swi3Δ and rec12Δ mutants. Intriguingly, loss of Rec12 moderately alleviates the spore viability defects of both swi1Δ and swi3Δ cells. Additionally, plating efficiency revealed no substantial decrease in viability in the FPC and Rec12 mutants compared with wild-type vegetative cells (Figure 1A and Supplemental Figure 1A). Therefore the reduction in spore viability likely reflects meiotic defects.

FPC mutants do not impede DNA replication

Because the FPC is necessary for proper replication fork progression (Noguchi et al., 2003, 2004), we asked whether the observed meiotic defects in the swi1Δ and swi3Δ strains were related to defects in meiotic S phase (meiS phase). To explore this, we induced synchronous meiosis by trapping cells in G1 through nitrogen starvation and by heat inactivation of Pat1, a negative meiotic regulator. Increasing the temperature to 34°C causes either haploid or diploid cells carrying the pat1-114 allele to enter meiosis. To avoid any confounding effects associated with meiotic induction of haploid cells (Yamamoto and Hiraoka, 2003; Pankratz and Forsburg, 2005), we generated stable h+/mat2-102 pat1-114/pat1-114 diploids in swi1Δ and swi3Δ backgrounds that could enter meiosis only by temperature shift (Pankratz and Forsburg, 2005; Le et al., 2013). We harvested cells over the course of 8 h and examined replication dynamics, DSB formation and repair, and meiotic progression. We monitored completion of the meiS phase by measuring DNA content changes with fluorescence-activated cell sorting (FACS) (reviewed in Sabatinos and Forsburg, 2009).

Wild-type cells finished bulk DNA replication 3 h after meiotic induction, while the FPC mutants exhibited a broadened peak at that time point, which indicates a modest delay in DNA synthesis (Figure 1B). After 4 h, both FPC mutants finished replication similarly to the wild-type diploid. Although rec12Δ cells also showed a slight delay in replication by hour 3, the rec12Δ swi1Δ and rec12Δ swi3Δ mutants did not entirely finish meiS by hour 6 (Supplemental Figure 1B). These observations indicate that the FPC is mostly dispensable
For replication progression in an unperturbed meiotic phase, except when Rec12 is removed, in which case DNA synthesis is substantially delayed.

Even though fork destabilization did not impede meiotic phase progression, it is possible that programmed DSB formation and repair, which follow replication, could be affected. To address this, we carried out pulse-field gel electrophoresis (PFGE) to separate chromosomes and the signature smears of DSB generation by size (Young et al., 2004). In wild-type cells and the FPC mutants, DSBs were initiated 3 h after induction. By hour 4, the majority of DSBs were created, as evidenced by a strong signal beneath chromosome III. DSB resolution occurred in hour 5 with a concomitant return of chromosome signals. By hour 6, all DSBs were repaired, which is consistent with full meiotic entry (Figure 1C and Supplemental Figure 2A). In the absence of Rec12, cells predictably failed to create any DSBs (Cervantes et al., 2000; Ogino et al., 2006). By contrast, rec12Δ swi1Δ and rec12Δ swi3Δ cells exhibited fast-migrating smears at all time points. This agrees with a constitutive level of DNA damage or DNA breaks and corroborates previous observations involving rec12Δ and other replication stability mutants (Dolan et al., 2010; Le et al., 2013) (Supplemental Figures 1C and 2B). These results imply that the FPC is not required for proper induction and repair of prDSBs during meiosis.

Because fork destabilization did not significantly disrupt DSB dynamics (Figure 1C and Supplemental Figure 2A), we asked whether the same was true for meiotic progression. We stained nuclei with 4′,6-diamidino-2-phenylindole (DAPI) and scored for changes in the number of DAPI-stained nuclei over the course of meiosis (Pankratz and Forsburg, 2005; Le et al., 2013). Wild-type cells and the FPC mutants showed a single DAPI mass until hour 4 and began to show two nuclei by hour 5, when cells enter MI. From hour 6 onward, cells continued to divide their nuclei, until most cells showed three or more DAPI masses, indicating completion of MII (Figure 1D). However, relative to the wild-type strain, swi3Δ and swi1Δ cells correspondingly showed delayed-MI and early-MII entry. Thus, although swi1Δ, but not swi3Δ, cells show similar kinetics to the wild-type strain during metaphase I, both FPC mutants also progressively impede meiotic phase progression, except when Rec12 is removed, in which case DNA synthesis is substantially delayed.

**FIGURE 1:** Spore viability and replication dynamics in FPC mutants. (A) Bulk spore germination of heterothallic cells homozygous for the presence or absence of the FPC and Rec12. Viability was determined by dividing microscopic counts of a spore suspension by the resulting colony counts for each strain. Percent figures indicate viability for each genotype and are presented relative to those of the wild-type strain. At least six trials were conducted per genotype (n ≥ 6000). Significance was determined using chi-squared analysis followed by false discovery rate correction for multiple sample comparisons. p values are reported as follows: ***, p < 0.0001; ****, p < 0.0001; *****, p <<< 0.0001. Error bars represent 95% confidence intervals. Synchronous meiosis in mat2-102/pat1-114/pat1-114 diploids was performed for 8 h. Cells were harvested every hour and examined as indicated in B–D. (B) FACS profiles showing progression of replication through meiotic induction. DNA doubling is denoted as the change from 2C to 4C DNA content. Images are representative of three independent trials. For B–D, hour 0 denotes the time when cells were switched from 25°C to 34°C to elicit meiotic induction. (C) PFGE used to separate whole chromosomes by size and to assess formation and repair of meiotic DSBs. Smears migrating faster than chromosome III represent DSBs. Images are ethidium bromide–stained agarose gels representative of three different trials. (D) DAPI-stained nuclei were counted for each time point to ascertain progression through meiotic divisions, which are reported as follows: black stands for 1 nucleus, dark gray for 2 nuclei, and light gray for 3 or more nuclei (3+); 2 and 3+ nuclei indicate onset of MI and MII, respectively. n ≥ 900 cells per genotype. Significance was determined using a chi-squared test for trend. p values are reported as follows: ns, not significant; *, p < 0.05; ****, p < 0.0001. Error bars are 95% confidence intervals.
while both FPC rec12Δ mutants entered MI late (Supplemental Figure 1D). This reduced efficiency in MI dynamics is consistent with similar observations in other replication mutants (Davis and Smith, 2003). These results suggest that fission yeast cells complete meiosis even when fork integrity is compromised, albeit inefficiently in the absence of either FPC component with or without Rec12. Taken together, these observations indicate that destabilizing the fork by eliminating Swi1 or Swi3 is associated with modest defects in meiotic prophase I that may disproportionately affect downstream events.

FPC mutants have persistent DNA damage

In vegetative cells, absence of the FPC components results in DNA damage accumulation following replication (Noguchi et al., 2004). We asked whether the same was true in meI phase, and whether such damage persisted through meiosis. We used cells in which histone H3, RPA, and Rad52 (fission yeast’s Hht1, Rad11, and Rad22, respectively) were fluorescently tagged and followed their meioses by live-cell microscopy (Libby et al., 2004; Sabatinos et al., 2012; Mastro and Forsburg, 2014) (Figure 2, A and B).

Wild-type cells exhibited RPA signals for most of horse tailing (HT), while a third of cells showed Rad52 signals in late HT, where nuclear oscillation ceases and metaphase I nuclear contraction begins (Figure 2, A–D). This agrees with the timing of chromosome pairing and recombination that ensues after DNA synthesis. When cells reached metaphase I, RPA signals began to decrease, but Rad52 signals showed a transient increase (Figure 2, A–D). This is an expected outcome during the period when most recombination is finalized. As cells entered MI, RPA and Rad52 signals disappeared in most cells and were mainly absent by the time MI was completed (Figure 2, A–D).

By contrast, we observed in the FPC mutants a higher number of cells with DNA damage signals that persisted through meiosis (Figure 2, A–D). In the swi1Δ and swi3Δ strains, the proportion of cells showing RPA foci remained relatively high through the end of MI (Figure 2, A–D). During metaphase I, there was an increase in cells with Rad52 foci, but this fraction gradually decreased in MI and MI. However, compared with the wild-type strain, the FPC mutants showed higher proportions of cells with Rad52 foci at the end of meiosis (Figure 2, A–D). While persistence of DNA damage signals was similar in both FPC mutants, the effect was more pronounced in swi1Δ cells. Three important differences could thus be appreciated between the FPC mutants and wild-type cells: the number of DNA damage signals was greater, more persistent, and at more loci, as evidenced by the larger group of cells showing multiple RPA and Rad52 foci during and after metaphase I. These results reveal that, upon fork destabilization, DNA damage accumulates and remains mostly unrepaired through meiosis, which may negatively affect events important for meiotic segregation.

FPC mutants alter homologous recombination

Rad52 is required to facilitate inter- and intrasister recombination between direct repeats (Muris et al., 1997; van den Bosch et al., 2001; Octobre et al., 2008). Because we observed abnormal Rad52 signals before and during meiosis upon fork destabilization, we asked whether normal recombination dynamics between homologues were affected. To address this question, we made crosses between parents carrying a pair of linked nutritional markers in coupling (+/+→−) or in repulsion (+/−→−). We assessed homologous recombination by scoring colony formation on solid media that selected for each tested nutritional outcome. Relative to wild-type cells, the FPC mutants showed 1.2- to 2.8-fold overall decrease in recombination at four distinct intergenic intervals (Table 1). We also assessed intragenic recombination at an ade6 locus containing a recombination hot spot (Schuchert and Kohli, 1988; Szankasi and Smith, 1995; Fleck et al., 1999). In this assay, a similar reduction in recombination was observed, but was especially pronounced in the absence of Swi3.

Furthermore, because Rad52 antagonizes Dmc1, a RecA-like protein important for homologous recombination (October et al., 2008; Murayama et al., 2013), we examined whether dmc1Δ cells recapitulated the spore viability and recombination outcomes of both FPC mutants. While reduction in spore viability was moderate compared with wild-type cells, the dmc1Δ mutant showed a 1.5-fold decrease in recombination at the his4-lys4 intergenic interval similar to that of the swi1Δ mutant (Supplemental Table 1).

This observation prompted us to ask whether fork destabilization shifted recombination dynamics to favor sister chromatids over homologues as recombination templates. For this, we set up a cross that produced Ade+ offspring only if one of the parent cells performed intra- or interister recombination during meiosis (Osman et al., 1996; Catlett and Forsburg, 2003; Mastro and Forsburg, 2014) (Table 1). Interestingly, in the absence of Swi1 and Swi3, there was a noticeable two- to threefold increase in sister chromatid exchange (Table 1). These results indicate that fork destabilization alters normal recombination dynamics, potentially reducing chiasma and thereby affecting meiotic division.

FPC mutants disrupt chromosome segregation

Chromosome segregation is sensitive to changes in recombination (Watanabe and Nurse, 1999; Murakami and Nurse, 2001; Sakuno et al., 2011; Mastro and Forsburg 2014). Thus we asked whether fork destabilization resulted in abnormal nuclear division in MI and MII. We used cells with fluorescently tagged histones (Hht1-mRFP [monomeric red fluorescent protein]) and followed the outcome of meiosis by live-cell microscopy (Figure 3, A and B) (Cooper et al., 1998; Chikashige and Hiraoka, 2001; Klutstein et al., 2015). Whereas few wild-type cells showed abnormal nuclear masses after MI and MII (Figure 3, C and D), the FPC mutants revealed three- to fourfold increases in cells containing asymmetric nuclei and nuclei with extra DNA signal (Figure 3, C and D). Moreover, in the absence of Swi1 and Swi3, cells exhibited two- to fourfold increases in lagging chromosomes following MI and MII (Supplemental Figure 3, A–C). Although lagging chromosomes do not always result in abnormally sized nuclei, increases in this phenotype may indicate underlying division defects (Yokobayashi and Watanabe, 2005). These observations therefore suggest that the FPC is important for proper chromosome segregation in meiosis.

Premature sister chromatid separation or chromosome nondisjunction could explain the missegregation phenotypes observed in the FPC mutants (Yokobayashi and Watanabe, 2005; Hauf et al., 2007; Dudas et al., 2011). To address the first possibility, we crossed cells heterozygous for a green fluorescent protein (GFP) marker integrated at the lys1 locus (near the centromere of chromosome I) and carrying histones fluorescently tagged as indicated earlier. If MI division occurs normally (i.e., reductional division), GFP signal is distributed to one of the two daughter nuclei. However, if equal division occurs (i.e., abnormal MI division), both nuclear spots show GFP signals (Yokobayashi and Watanabe, 2005; Hauf et al., 2007; Hirose et al., 2011) (Figure 4, A and B). Compared with the wild-type strain, swi1Δ, but not swi3Δ, cells showed a threefold increase in equal division (Figure 4C). This result implies that Swi1 helps to prevent precarious separation of sister chromatids, possibly by contributing to correct monopolar attachment, as was reported previously for cells lacking both Mrc1 (a secondary FPC factor) and Rec12 (Hirose et al., 2011).
FIGURE 2: Persistence of DNA damage in FPC mutants. (A) Live-cell images of meiotic cells carrying fluorescently tagged histone3 (Hht1-mRFP), RPA (Rad11-CFP), and Rad52 (Rad22-YFP). Panels are shown for individual fluorescent channels. Bright-field panels are omitted due to space limitations. A false-color image is rendered to show merged signals. Dotted cell outlines are overlaid on panels for easier visualization of meiotic cells. Images that show the characteristics of each reported mutant phenotype were used. Selected time frames were chosen for optimal representation of nuclear dynamics. The analyzed meiotic time window encompasses HT, metaphase (MT), MI, and MII. Upright scale bars: 5 µm. (B) Cartoon depicting DNA damage persistence through meiosis in FPC mutants. Gray and black dots represent RPA and Rad52, respectively. (C, D) Quantification of RPA and Rad52 signals in cells from microscopy pictures shown in A. More than 150 cells were scored from at least two independent movies for each genotype. Chi-squared analysis was used to determine significance. p values are reported thus: **, p < 0.01; ***, p < 0.001; ****, p << 0.0001. Error bars represent 95% confidence intervals. RPA and Rad52 signals are binned as follows: light-gray for 1 focus, white for 2 foci, and black for more than 3 foci.

To examine whether fork destabilization is associated with chromosome nondisjunction, we used a similar strategy, but in a cross homozygous for the GFP marker at lys1. In each meiotic division, if GFP dots are distributed symmetrically, normal division is inferred to have occurred, whereas asymmetric distribution indicates abnormal division (Yokobayashi and Watanabe, 2005; Hauf et al., 2007;
**TABLE 1:** Recombination dynamics in FPC mutants with intergenic and intragenic recombination data given in the top panels and sister chromatid recombination data given in the bottom panel.

| Intergenic interval | Chr | Genomic distance (cM)a | Fold reductionb | p valueb |
|---------------------|-----|------------------------|----------------|----------|
|                     |     | WT | swi1Δ | swi3Δ | swi1Δ | swi3Δ | swi1Δ | swi3Δ | swi1Δ | swi3Δ | swi1Δ | swi3Δ | swi1Δ | swi3Δ |
| his4-lys4Δ          | 2   | 6.81 ± 0.92 | 4.81 ± 0.78 | 2.43 ± 0.83 | 1.42 | 2.80 | 0.0039 | p << 1E-04 |
| leu2-ura2Δ          | 1   | 0.87 ± 0.13 | 0.67 ± 0.20 | 0.40 ± 0.24 | 1.30 | 2.18 | 0.1164 | 0.0051 |
| leu1-his7Δ          | 2   | 2.28 ± 0.41 | 1.08 ± 0.34 | 1.21 ± 0.20 | 2.11 | 1.88 | 0.0004 | 0.0005 |
| his3-leu1Δ          | 2   | 1.13 ± 0.22 | 0.55 ± 0.08 | 0.92 ± 0.08 | 2.05 | 1.23 | 0.0008 | 0.1078 |

| Intragenic intervalf | Chr | Frequency (x10^-3) | Fold reductionb | p valueb |
|----------------------|-----|--------------------|----------------|----------|
| ade6-M26-52          | 3   | 3.34 ± 0.08 (60/17987) | 1.70 ± 0.05 (52/31129) | 0.60 ± 0.04 (7/11838) | 1.96 | 5.57 | 0.0009 | P << 1E-04 |

| SCE intervalf | Chr | Frequency (x10^-3) | Fold increaseb | p valueb |
|---------------|-----|--------------------|----------------|----------|
| ade6-L469-his3+-ade6-M375/ade6-M375-M210 | 3 | 5.00 ± 0.08 (167/33465) | 16.81 ± 0.24 (186/11064) | 9.13 ± 0.14 (155/16982) | 3.36 | 1.83 | P << 1E-04 | p << 1E-04 |

Significance for genetic distance was determined using a two-tailed Student’s t test. Chi-squared analysis was employed for frequencies. At least five independent trials were carried out per genotype. Error margins represent 95% confidence intervals. Fold change values are relative to wild-type outcomes. At least five independent trials were performed per genotype. Chr, chromosome; WT, wild type.

aHaldane’s formula cM = −50 ln(1−2R) was used to convert recombinant frequencies (R) into centimorgans (cM).

bWild-type cM or frequency values were divided by those of mutants to determine reduction in genetic distance or recombination frequency.

cSignificance calculated by chi-squared test.

dSignificance calculated by chi-squared analysis.

eFrequency values were calculated by dividing number of Ade+ by total number of colonies screened (shown in parentheses).

fFrequency values of mutants were divided by those of the wild type to determine increase in recombination frequency.

gSignificance calculated by chi-squared analysis.

Hirose et al., 2011) (Figure 5, A and B). While only swi3Δ cells show elevated nondisjunction in MI (Figure 5C), both FPC mutants show two- to fourfold increases in nondisjunction of sister chromatids in MI (Figure 5D). These results suggest that Swi3 is necessary for proper homologue segregation in MI, while both FPC components contribute to normal sister chromatid separation in MI.

**FPC mutants destabilize centromeric cohesion**

Some of the missegregation phenotypes associated with fork destabilization can be explained by the effects of reduced homologous recombination (Dudas et al., 2011). However, problems with centromeric cohesion are also associated with abnormal meiotic divisions, as shown previously in Rec8 mutants (Watanabe and Nurse, 1999). We examined dynamics associated with meiotic cohesion using live-cell microscopy. Wild-type and mutant cells carried fluorescently tagged histones and either Rec27-GFP (a component of the linear elements), Sgo1-GFP (shugoshin, a centromeric cohesion protector), or Rec8-GFP (the α-kleisin subunit of meiotic cohesin). Each of these proteins is associated with different aspects of cohesin function and regulation (Watanabe and Nurse, 1999; Kitajima et al., 2004, 2006; Davis et al., 2008).

We reasoned that any changes to the temporal dynamics of Rec27, Sgo1, and Rec8 upon fork destabilization would indicate cohesin-related sources of chromosome missegregation. Casein kinase 1 phosphorylation of the cohesin subunit Rec11 (SA3) facilitates loading of the linear element proteins Rec10 and Rec27 along chromosomal axes, which in turn promotes the alignment of homologous chromosomes, Rec12-dependent DNA breakage, and recombination (Ellermeier and Smith, 2005; Davis et al., 2008; Phadnis et al., 2015; Sakuno and Watanabe, 2015). For the examination of Rec27, Sgo1, and Rec8, we measured GFP fluorescence intensity and scored for the disappearance of GFP foci. This complementary approach provides information about GFP signal abundance and GFP organization as cells proceed from late prophase I to just before MI (designated as time point 0) or from late metaphase I to MI in the case of Rec8. The FPC mutants showed 20–30% lower Rec27-GFP intensity before meiosis, and Rec27-GFP foci elimination was 5–7 min faster and occurred 5 min later in metaphase I than in wild-type cells (Figure 6, A–C, and Supplemental Figure 4, A and B). These Rec7 defects are suggestive of changes to the structural integrity of chromosome axes that may affect homologue pairing and recombination. They may also indicate changes to cohesin stability in meiosis.

In fission yeast, Sgo1 and PP2A protect centromeric cohesin from separate degradation before anaphase I (Kitajima et al., 2004; Riedel et al., 2006; Ishiguro et al., 2010). Given that swi1Δ cells show increased equational division in MI, which is linked to issues with centromeric cohesin stability (Yokobayashi and Watanabe, 2005; Hauf et al., 2007), we asked whether fork destabilization affects the temporal dynamics of Sgo1 (Figure 7, A–B). Relative Sgo1-GFP signal intensity was similar across genotypes. Intriguingly, we observed that swi3Δ, but not the swi1Δ mutant, exhibited a 3-min reduction in the duration of Sgo1-GFP foci compared with wild-type cells (Figure 7, A–D). However, without Swi1, Sgo1-GFP dissipates 2 min later than was observed in the wild-type strain (Supplemental Figure 4, C and D). These results reveal the FPC is necessary for temporal stability of Sgo1 at the centromere and further highlight the separate functions of the FPC components.

Rec8 is a subunit of cohesin that helps keep sister chromatids together following meiS phase (Watanabe and Nurse, 1999; Watanabe et al., 2001). During anaphase I, Rec8 is removed from chromosome arms, which facilitates chiasma resolution and segregation of homologous chromosomes (Bueno et al., 2000;
Kitajima et al., 2003b; Brar et al., 2006). Nevertheless, Rec8 is retained at the centromere until right before anaphase II to ensure correct segregation of sister chromatids (Kitajima et al., 2003b, 2004; Riedel et al., 2006). In addition, Rec8 and Moa1 facilitate mono-orientation of kinetochores in MI, thus promoting segregation of homologous chromosomes to opposite poles of the cell (Yokobayashi and Watanabe, 2005; Hauf et al., 2007). Because lack of Swi1 and Swi3 is associated with abnormal MI division and anomalous Sgo1 timing before anaphase I, we asked whether fork destabilization would disrupt the temporal dynamics of centromeric cohesion. For this, we used live-cell imaging to follow Rec8-GFP from late metaphase I to MI (Figure 8, A and B, and Supplemental Figure 4G).
In wild-type cells, Rec8-GFP signal goes from pan-nuclear (observed in late metaphase I to early anaphase) to a pair of foci that localize to the centromeres and generally persist for more than half an hour (Figure 8, A, B, and E). This signal pattern following MI corresponds to Rec8 elimination at chromosome arms, persistence at the centromere, and degradation before MII (Watanabe and Nurse, 1999; Kitajima et al., 2003b; Le et al., 2013) (Figure 8B). In the FPC mutants, Rec8-GFP signal intensity was higher, but persistence of centromeric cohesion was 8–10 min shorter, and Rec8-GFP foci loss occurred 12–15 min earlier than in wild-type cells (Figure 8, C–E, and Supplemental Figure 4, F and G). This result agrees with the general observation 10 min after MI that Rec8-GFP signal remains as centromeric foci in the wild-type strain but dissolves throughout the nucleus in the FPC mutants (Figure 8, A–E, and Supplemental Figure 4, F and G).

We next investigated whether Rec8 localization at the centromere is affected over the period that encompasses the MI to MII transition, when centromeric Rec8 is protected to keep sister chromatids together until anaphase II. Cells were synchronized in G1 phase of the cell cycle by nitrogen starvation, and meiosis was induced by Pat1 inactivation. We then performed chromatin immunoprecipitation followed by quantitative PCR (ChIP-qPCR). Although the FPC mutants and wild-type cells were statistically similar, fork destabilization consistently led to modest reductions in centromeric Rec8 before and after MI (Supplemental Figure 4E). These results, along with the observation that sister chromatids in the FPC mutants are left untethered at the centromeres longer than in wild-type cells (Supplemental Figure 4, F and G), indicate that fork destabilization accelerates elimination of centromeric cohesion, which not only precipitates early separation of sister chromatids in MI but also contributes to their abnormal segregation in MII.

Swi6 (the fission yeast homologue of human HP1) is essential for heterochromatin formation and for the recruitment of cohesin and Sgo1 to the centromere (Ekwall et al., 1995; Nonaka et al., 2002; Kitajima et al., 2003b; Kawashima et al., 2007; Yamagishi et al., 2008). To further corroborate the centromeric cohesion defects thus far observed, we asked whether fork destabilization impacts centromeric localization of Swi6. We used live-cell microscopy to monitor colocalization of Cnp1-mCherry (the fission yeast homologue of mammalian CENP-A, a centromere-specific histone variant; Takaishi et al., 2000) and Swi6-GFP following MI and MII (Figure 9, A and B). We reasoned that colocalization dynamics of these proteins after each division should reflect the overall ability of cells to mobilize heterochromatin back to the centromere. We observed that swi1Δ, but not swi3Δ or wild-type cells, had a significant three- to 10-fold increase in cells where Cnp1 and Swi6 did not colocalize after each meiotic division (Figure 9, C and D). Although the effect is statistically modest, it nonetheless suggests Swi1 is necessary for correct recruitment of heterochromatin to the centromere.

**Discussion**

The FPC travels with the replisome and is important for coupling the MCM helicase with the DNA polymerases (Tourrière et al., 2005; Ünsal-Kaçmaz et al., 2007; Leman et al., 2010; Leman and Noguchi, 2012). This function helps prevent excessive unwinding and ssDNA accumulation during replication stress. Areas with extended ssDNA segments are prone to fork reversal and unregulated genomic

**FIGURE 4**: MI equational division in FPC mutants. (A) Live-cell images of meiotic cells carrying fluorescently tagged histone3 (Hht1-mRFP) and LacI-GFP on a LacO repeat at lys1 near the centromere of chromosome I. Panels are divided into individual fluorescent channels. Bright-field panels are omitted due to space limitations. A false-color image was generated to show merged signals. Dotted cell outlines are overlaid on panels for easier visualization of meiotic cells. Images that show the characteristics of each reported mutant phenotype were used. Displayed time frames were chosen for optimal representation of nuclear dynamics. The meiotic phases shown are HT, metaphase (MT), MI, and MII. Minute 0 (0′) denotes the last nuclear mass contraction in metaphase I before homologous chromosomes separate in anaphase I. Upright scale bars: 5 µm. (B) Cartoon depicting a pair of homologous chromosomes where one is marked with GFP near the centromeres. Symmetrical distribution of GFP indicates equational division. (C) Quantification of cells showing equational division in MI. More than 150 cells were scored from at least two independent movies for each genotype. Chi-squared analysis was used to determine significance. *p* values are reported as follows: ***, **, and *p < 0.001. Error bars represent 95% confidence intervals.
DNA-binding homeobox-containing proteins and different transcription and chromatin-remodeling factors (Noguchi et al., 2012). This domain is also important for recruiting Swi3 (Noguchi et al., 2012). In fission yeast, the FPC's primary components are Swi1 and Swi3 (Noguchi et al., 2004). Swi1 associates with chromatin through its DDT domain, found in recombination (Noguchi et al., 2003, 2004). In fission yeast, the FPC's primary components are Swi1 and Swi3 (Noguchi et al., 2004). Swi1 associates with chromatin through its DDT domain, found in

FIGURE 5: Nondisjunction in MI and MII in FPC mutants. (A) Live-cell images of meiotic cells carrying fluorescently tagged histone3 (Hht1-mRFP) and LacI-GFP on a LacO repeat at lys1 near the centromere of chromosome I. Panels are divided into individual fluorescent channels. Bright-field panels are omitted due to space limitations. A false-color image is provided to show merged signals. Dotted cell outlines are overlaid on panels for easier visualization of meiotic cells. Images that show the characteristics of each reported mutant phenotype were used. Displayed time frames were chosen for optimal representation of nuclear dynamics. The meiotic phases shown are HT, metaphase (MT), MI, and MII. Minute 0 (0′) denotes the last nuclear mass contraction in metaphase I before homologous chromosomes separate in anaphase I (10′). Upright scale bars: 5 µm. (B) Cartoon depicting a pair of homologous chromosomes marked with GFP near the centromeres. Asymmetric distribution of GFP indicates nondisjunction events. (C, D) Quantification of cells showing chromosome nondisjunction in MI and MII. More than 150 cells were scored from at least two independent movies for each genotype. Chi-squared analysis was used to determine significance. p values are reported as follows: *, p < 0.05. Error bars represent 95% confidence intervals.
FIGURE 6: Elimination of the linear elements in FPC mutants. (A) Live-cell images of meiotic cells carrying fluorescently tagged histone3 (Hht1-mRFP) and Rec27-GFP. Panels are divided into individual fluorescent channels. Bright-field panels are omitted due to space limitations. A false-color image was rendered to show merged signals. Dotted cell outlines are overlaid on panels for easier visualization of meiotic cells. Images that show the characteristics of each reported mutant phenotype were used. Displayed time frames were chosen for optimal representation of nuclear dynamics. The meiotic phases shown are metaphase (MT) and MI. Time preceding the last nuclear mass contraction in metaphase I (0′) before homologous chromosomes separate in anaphase I (10′) is denoted with a negative sign (e.g., −10′). Time 0 is omitted due to space limitations. Upright scale bars: 5 µm. White arrows represent average span of Rec27-GFP elimination. (B) Cartoon depicting elimination of Rec27-GFP during metaphase and before MI. (C) Quantification of Rec27-GFP fluorescence. (D) Quantification Rec27-GFP foci elimination. (E) Duration of Rec27-GFP foci elimination. More than 150 cells were scored from at least two independent movies for each genotype. One-way ANOVA followed by Tukey family-wise comparison was used to determine significance in C and E. Chi-squared analysis was used for D. p values are reported as follows: *, p < 0.05; ***, p < 0.001; ****, p << 0.0001. Error bars represent 95% confidence intervals.
FIGURE 7: Duration of Sgo1 in metaphase in FPC mutants. (A) Live-cell images of meiotic cells carrying fluorescently tagged histone3 (Hht1-mRFP) and Sgo1-GFP. Panels are divided into individual fluorescent channels. Bright-field panels are omitted due to space limitations. A false-color image was generated to show merged signals. Dotted cell outlines are overlaid on panels for easier visualization of meiotic cells. Images that show the characteristics of each reported mutant phenotype were used. Displayed time frames were chosen for optimal representation of nuclear dynamics. The meiotic phases shown are HT, metaphase (MT), and MI. Time preceding the last nuclear mass contraction in metaphase I (0') before homologous chromosomes separate in anaphase I (10') is denoted with a negative sign (e.g., −10'). Upright scale bars: 5 µm. White arrows indicate average span of Sgo1-GFP signal. (B) Cartoon depicting duration of Sgo1-GFP signal during metaphase and before MI. (C) Quantification of Sgo1-GFP fluorescence. (D) Quantification of Sgo1-GFP foci in metaphase I. (E) Duration of Sgo1-GFP foci. More than 150 cells were scored from at least two independent movies for each genotype. One-way ANOVA followed by Tukey family-wise comparison was used to determine significance in C and E. Chi-squared analysis was used for D. p values are reported as follows: *, p < 0.05. Error bars represent 95% confidence intervals.
Stable association of Swi1 and Swi3 facilitates binding to the fork of Mrcl, a secondary FPC component necessary for replication processivity and activation of the replication checkpoint (Shimmoto et al., 2009; Matsumoto et al., 2010). Together these proteins act as sensors of replication stress that protect genomic integrity during DNA duplication. Furthermore, Swi1 and Swi3 are also involved in restarting the fork after exposure to S-phase stressors and play a role in establishing and maintaining proper sister chromatid cohesion in vegetative cells (Noguchi et al., 2004; Matsumoto et al., 2005; Ansbach et al., 2008; Rapp et al., 2010).

In fission yeast, meiS phase takes longer to complete than S phase in vegetative conditions (Wu and Nurse, 2014). DNA damage induced at this stage does not elicit checkpoint arrest and can be repaired through meiotic recombination. In checkpoint mutants, however, excess damage may persist until at least the first meiotic division (Pankratz and Forsburg, 2005). Perturbations to meiS phase are associated with recombination and segregation problems downstream (Watanabe et al., 2001; Wu and Nurse, 2014). Because the FPC contributes to replication processivity and the response to DNA damage, we asked whether absence of its components would disrupt any aspect of the meiotic program.

Loss of spore viability

We observed that the FPC mutants showed decreased spore viability (Figure 1A), which is consistent with previous reports of similar outcomes in other replication-stability mutants (Dolan et al., 2010; Le et al., 2013; Mastro and Forsburg, 2014; Murakami and Keeney, 2014; Wu and Nurse, 2014). The reduction seen in the swi1Δ swi3Δ strain is similar to that of swi1Δ cells and agrees with Swi1's function of stabilizing Swi3 at the fork (Leman et al., 2010; Leman and Noguchi, 2012; Noguchi et al., 2012). Interestingly, combining rec12Δ with the swi1Δ or swi3Δ deletions partially suppressed 33–45% spore viability observed in the single mutants (Figure 1A). This observation suggests that the damage-induced recombination intermediates previously seen during vegetative growth in the FPC mutants (Noguchi et al., 2003, 2004) may also be present during meiotic prophase I and may serve as recombination sources, albeit suboptimal ones, when prDSBs are abolished (Cervantes et al., 2000; Pankratz and Forsburg, 2005).

Meiotic replication and DSB formation

Absence of the FPC components did not halt meiotic progression but was associated with moderately delayed replication (Figure 1, B–D). It is possible that replisome progression was affected by unstable DNA structures like ssDNA gaps or rearranged forks that result from the uncoupling of DNA helicase and the DNA polymerases (Noguchi et al., 2003, 2004; Leman et al., 2010). In that context, physical impediments to fork progression would be alleviated by homologous recombination, which relies on Rec12 for DSB formation. Therefore Rec12 may assist fork progression by stimulating the removal of DNA barriers generated by unstable forks. This scenario is consistent with the observations that swi1Δ and swi3Δ cells repair DSBs as well as the wild-type strain (Figure 1C and Supplemental Figure 2A). Also, when lacking Rec12, the FPC mutants exhibit substantial delay in replication and generate low-migrating PFGE smears indicative of constitutive DNA damage and incomplete DSB repair (Supplemental Figures 1C and 2B). However, because these meiotic induction experiments were carried out in pat1-114 genetic backgrounds, in which chromosome segregation is not entirely normal (Cipak et al., 2012), we cannot eliminate the possibility that the observed results are pat1-114 specific and thus may be different from wild-type meiosis.

Persistent DNA damage

Deletion of swi1 and swi3 generates increased RPA and Rad52 foci in vegetative cells (Noguchi et al., 2004). In the FPC mutants, we observed elevated signals for both markers that intensified in late HT and persisted until MII. This result is consistent with an FPC role in surveilling excess ssDNA levels and in maintaining stability at the fork, and with failure of meiotic cells to respond to meiS phase damage (Pankratz and Forsburg, 2005). A secondary consequence of accumulating large pools of Rad52 is the impact this may have on recombination, because its dissociation is crucial for Rad51 binding (New et al., 1998; Shinohara and Ogawa, 1998). We observed normal formation of prDSBs by PFGE, suggesting that the fork protection complex is not associated with activation of Rec12. We analyzed four distinct intergenic loci and one intragenic locus to assess recombination activity in the FPC mutants (Table 1). We observed a general approximately twofold decrease in homologous recombination similar to that of dmc1Δ cells at the his4-lys4 intergenic locus (Supplemental Table 1), in which exchange between homologues is reduced. This prompted us to ask whether a decrease was also true for sister chromatid exchange. Deletion of the FPC components in meiosis led to increased recombination between sister chromatids (Table 1). This alteration to recombination dynamics, while potentially ensuring resolution of surplus recombination substrates, may compromise downstream meiotic activity, which depends on chiasma formation and resolution for proper chromosome segregation (Dudas et al., 2011; Hirose et al., 2011).

Chromosome missegregation

We used live-cell imaging to follow the FPC mutants through meiosis (Figure 3, A–D, and Supplemental Figure 3, A–C). We observed that, without Swi1 and Swi3, cells have increased lagging chromosomes and chromosome missegregation in both meiotic divisions. We asked whether the observed division anomalies resulted from premature sister separation or chromosome nondisjunction (Figure 4, A–C). Interestingly, we found separate functions of Swi1 and Swi3 during MI. Swi1 appears to facilitate accurate monopolar attachment, because its absence is associated with an increase in equational division. By contrast, Swi3 seems to mediate correct disjunction of homologous chromosomes in MI. In MII, both FPC mutants exhibited increased nondisjunction of sister chromatids (Figure 5, A–D). Thus the missegregation phenotypes of the FPC mutants may be partially related to their reduced homologous recombination.

Cohesin regulators

Another possibility is that these phenotypes result from defects in centromeric cohesion (Watanabe and Nurse, 1999; Kitajima et al., 2003a; Yokobayashi and Watanabe, 2005). We addressed this by examining dynamics of cohesion-associated markers. We looked at the temporal dynamics of Rec27 because it is important for homologue pairing and its elimination coincides with DSB repair and chiasma resolution (Figure 6, A–C, and Supplemental Figure 4, A and B). We found that fork destabilization is associated with reduced Rec27-GFP signal and with accelerated Rec27-GFP foci elimination, which may contribute to recombination anomalies and suggests possible cohesion issues (Phadnis et al., 2015; Sakuno and Watanabe, 2015). Because Sgo1 protects centromeric cohesion before the onset of anaphase I (Kitajima et al., 2004; Ishiguro et al., 2010), we also examined its temporal dynamics in metaphase I (Figure 7, A–C, and Supplemental Figure 3, C and D). For this, we used Sgo1-GFP strains that did not exhibit centromere localization issues, as was
FIGURE 8: Persistence of centromeric cohesion in FPC mutants. (A) Live-cell images of meiotic cells carrying fluorescently tagged histone3 (Hht1-mRFP) and Rec8-GFP. Panels are divided into individual fluorescent channels. Bright-field panels are omitted due to space limitations. A false-color image was generated using ImageJ’s Fire LUT to better show intensity of GFP signals. Dotted cell outlines are overlaid on panels for easier visualization of meiotic cells. Images that show the characteristics of each reported mutant phenotype were used. Displayed time frames were chosen for optimal representation of nuclear dynamics. The meiotic phases shown are metaphase (MT), MI, and MII. Minute 0 (0’) denotes the last nuclear mass contraction in metaphase I before homologous chromosomes separate in anaphase I (10’). Upright scale bars: 5 µm. (B) Cartoon depicting gradual elimination of Rec8, first at chromosome arms and then at the centromere. Rec8 dynamics are shown in the context of MI and MII division. (C) Quantification of Rec8-GFP
fluorescence from metaphase I to MII. (D) Quantification of Rec8-GFP foci loss before MII. (E) Duration of Rec8-GFP foci loss. More than 150 cells were scored from at least two independent movies for each genotype. One-way ANOVA followed by Tukey family-wise comparison was used to determine significance in C and E. Chi-squared analysis was used for D. p values are reported as follows: *, p < 0.05; ****, p << 0.0001. Error bars represent 95% confidence intervals.
FIGURE 10: Model of FPC contribution to meiotic segregation. Proposed model explaining the missegregation phenotypes of FPC mutants. Swi1 stabilizes Swi3 at the fork. Both Swi1 and Swi3 help to suppress generation of large ssDNA gaps bound by excess RPA. Normal RPA levels facilitate proper binding of Rad52 to nucleoprotein filament, which in turn promotes Rad51 binding. Both Rad51 and Dmc1 promote normal homologous recombination, which guarantees proper disjunction of homologous chromosomes in MI. Swi1 is necessary for normal recruitment of Swi6 to the centromere. Centromeric Swi6 helps recruit Sgo1, which stabilizes Rec8 at the centromere. Stable centromeric cohesion ensures monopolar attachment of sister kinetochores, which is necessary for correct reductional segregation in MI. Suppression of DNA damage and maintenance of cohesion stability at the centromere ensures accurate separation of sister chromatids. DDK physically interacts with the FPC components and Swi6. It is also involved in the regulation of Rec8. Thus DDK is shown as having a potential role in modulating cohesion stability at the centromere by a yet unknown mechanism (hence the dashed lines on the side).

Previously reported for a Sgo1-GFP construct lacking a 3′ untranscribed region sequence (Rabitsch et al., 2004). We observed that deletion of Swi3, but not Swi1, decreases the timing of Sgo1 signal. This result implies that Swi3 is important for normal protection of centromeric cohesion and that this function may be independent of its association with Swi1 at the fork. Interestingly, however, absence of Swi1 shifts the timing of Sgo1 centromeric dissociation to a later stage in metaphase I (Supplemental Figure 4, C and D), thus also affecting Sgo1 regulation of Rec8.

Cohesin instability at the centromere

Rec8 is the cohesin subunit whose separase-dependent degradation ensures proper segregation of sister chromatids in MI (Kitajima et al., 2003a; Ishiguro et al., 2010). Because both FPC mutants showed abnormal MI division, we used live-cell imaging to examine the duration of centromeric Rec8 from MI to MII (Figure 8, A–D, and Supplemental Figure 4, F and G) (Le et al., 2013). In the FPC mutants, relative Rec8-GFP intensity was higher than in wild-type cells before and during meiosis. On the basis of this result, we surmise that lack of the FPC components may interfere with Rec8-GFP dissociation dynamics, respectively leading to delayed and early Rec8-GFP elimination at chromosome arms and the centromere. Indeed, we observed that, in swi1Δ and swi3Δ cells, most Rec8-GFP foci significantly disappeared within the first 30 min of meiosis. This meant that for the remaining 20–30 min before MII, cells lacking either FPC component were mostly untethered at the centromere (Figure 8, A–D, and Supplemental Figure 4, F and G). This may explain the high proportion of cells that exhibited nondisjunction issues in MII.

To corroborate this finding, we examined the localization of Rec8 at the centromere during the period when it is actively protected. Relative to wild-type cells, the FPC mutants consistently showed modest reductions of centromeric Rec8 (Supplemental Figure 4E). Although this result is insufficient to entirely explain the FPC mutants’ missegregation phenotype, it suggests that subtle changes in centromere dynamics may impair the function of other cohesin regulators. Indeed, because the heterochromatin protein Swi6 helps to recruit both cohesin and Sgo1 to the centromere (Yamagishi et al., 2008), we asked whether Swi6 recruitment to centromeres is affected in the FPC mutants during MI and MII (Figure 9, A–D). We found that, without Swi1, cells mislocalize Swi6 relative to Cnp1 (a centromeric histone). This observation confirms that the FPC is necessary to maintain cohesion stability at the centromere, a result that has been previously reported for vegetative cells (Ansbach et al., 2008).
homologue nondisjunction issues observed in both FPC mutants, but particularly in swi3∆ cells.

In the absence of Swi1, we observed problems with Swi6 recruitment to the centromere. As shown previously, this effect may disrupt centromeric Rec8 by preventing efficient cohesin and Sgo1 association (Yamagishi et al., 2008). Moreover, swi1A and swi3∆ cells showed defects in the timing of Sgo1 association, an outcome that may further compromise cohesin stability (Figure 10 and Supplemental Figure S5). Thus we propose that Swi1 is important for monopolar attachment and reductional division in MI by maintaining cohesion stability at the centromere. This connection with Rec8 dynamics may also confirm the FPC mutants’ problems with recombination via disruption of Rec27-GFP loci elimination. Given that the FPC physically interacts with Hsk1-Dfp1, fission yeast’s Dbf4-dependent kinase (DDK), and DDK has a role in regulating meiotic replication, DSB formation, and cohesin (Matsumoto et al., 2005, 2010; Matos et al., 2008; Kats et al., 2010; Le et al., 2013), our model predicts DDK or a yet unknown DDK substrate as the link that mechanistically connects replication with meiotic segregation, which would confirm similar observations made in budding yeast (Murakami and Keeney, 2014).

**Significance to human reproductive health**

Mammalian meiosis is characterized by a long-lasting prophase I (Reichman et al., 2017). Insults to recombination and segregation can severely compromise reproductive outcomes (Herbert et al., 2015). Increasingly, it has become apparent that factors involved in replication during cell proliferation can also regulate important aspects of the meiotic program (Mastro and Forsburg, 2014; Murakami and Keeney, 2014; Wu and Nurse, 2014; Nguyen et al., 2017). In this work, we observed that the FPC components are required for proper cohesion dynamics. When the FPC is destabilized, cells exhibit increased missegregation in both meiotic divisions with concomitant persistence of DNA damage markers. In fission yeast, thus lack of either FPC component significantly reduces the fitness of meiotic products. Given that FPC functions are conserved from yeast to human (Leman and Noguchi, 2012), our findings have major implications for reproductive health. It is possible that many cohesinopathies (van der Lelij et al., 2010) and aneuploidies known to result in embryonic lethality and arrested development may arise from replication-related processes. Further examination of additional prophase I–related events will reveal commonalities and differences that define the functional trajectories of the mitotic and meiotic cell division programs.

**MATERIALS AND METHODS**

**Cell growth and culture**

Supplemental Table 2 shows the strains used in this work. Detailed descriptions of general fission yeast culture conditions, media, and standard techniques are found in Sabatinos and Forsburg (2010). Liquid and solid yeast extract plus supplements (YES) was used for the construction and maintenance of all strains, except when starved for 16.5–17 h in EMM minus nitrogen. Starved cultures were refed EMM containing a 1:2 dilution of supplements (minus adenine) plus NH4Cl and shifted from 25°C to 34°C for meiotic induction. Subsequently cells were harvested every hour (FACS samples were also taken), treated with sodium azide, and washed in phosphate-buffered saline (PBS) and citrate-phosphate sorbitol EDTA. A lysing enzymes solution containing 0.2 mg/ml 100T Zymolyase (Nacalai Tesque, Kyoto, Japan) and 0.45 mg/ml Trichoderma harzianum–lysing enzymes (Sigma, St. Louis, MO) was used to digest the cell wall. Following the first time point, the lysing solution was titrated to 50% and 25% in the second and all subsequent time points, respectively, as performed by Cervantes et al. (2000). Lysate from cells was mixed with 2% CleanCut Agarose (Bio-Rad, Hercules, CA) to generate pulse-field gel plugs, which

**PFGE**

Stable diploids were created using ade6-M210/M216 complementation in parent strains carrying the mat2-102 and pat1-114 alleles to perform synchronous meiosis as described by Catlett and Forsburg (2003) with a few adjustments. Briefly, cultures were grown in Edinburgh minimal medium (EMM) plus supplements, except adenine, to an OD650 = 0.7–1.0, after which cells were starved for 16.5–17 h in EMM minus nitrogen. Starved cultures were refed EMM containing a 1:2 dilution of supplements (minus adenine) plus NH4Cl and shifted from 25°C to 34°C for meiotic induction. Subsequently cells were harvested every hour (FACS samples were also taken), treated with sodium azide, and washed in phosphate-buffered saline (PBS) and citrate-phosphate sorbitol EDTA. A lysing enzymes solution containing 0.2 mg/ml 100T Zymolyase (Nacalai Tesque, Kyoto, Japan) and 0.45 mg/ml Trichoderma harzianum–lysing enzymes (Sigma, St. Louis, MO) was used to digest the cell wall. Following the first time point, the lysing solution was titrated to 50% and 25% in the second and all subsequent time points, respectively, as performed by Cervantes et al. (2000). Lysate from cells was mixed with 2% CleanCut Agarose (Bio-Rad, Hercules, CA) to generate pulse-field gel plugs, which
were then incubated at 55°C in a protease K (Bioline Reagents, Taunton, MA)/sarkosyl-EDTA solution for 48 h, with an intermediate change of protease solution after 24 h. The DNA plugs were then washed in Tris-EDTA (TE) and equilibrated in Tris-acetate-EDTA (TAE). A Bio-Rad Chef II Pulse Field Machine was employed to separate DNA by size using a 1% Mega Base Agarose (Bio-Rad, Hercules, CA)/TAE gel. The pulse-field apparatus was run for 48 h at 2 V/cm, with an 1800-s switch time and a 106° angle. The gel was stained in ethidium bromide and visualized in a ChemiDoc Imager (Bio-Rad, Hercules, CA). ImageJ (National Institutes of Health, Bethesda, MD) was used to quantify DSB formation and repair as performed previously (Borde et al., 2000; Mastro and Forsburg, 2014). Briefly, the ethidium bromide signal beneath chromosome III was measured and divided by the summed chromosome signal values of each time point. Significance was computed using one-way analysis of variance (ANOVA) followed by Tukey’s family-wise comparisons.

**Live-cell imaging**

For live-cell imaging, heterothallic strains were grown in PMG with appropriate supplements at 25°C or 32°C to late log phase (OD280 = −0.8). Cells were concentrated and washed in liquid ME and incubated at 25°C for 12–20 h in an air shaker. Starved cells were pelleted in a microfuge and spread on pads made with 2% agarose in liquid sporulation media on top of glass microscope slides. Cover lids were carefully placed on prepared pads and moved clockwise with index finger for two rotations to generate a cell monolayer. Once cell matter was properly distributed, pads were seeded with 1:1 Vaseline/lanolin/paraffin (at indicated ratios by weight).

Imaging was carried out as described in Mastro and Forsburg (2014) and Sabatinos et al. (2015) with a few modifications. Cells were staged at 25°C using a DeltaVision microscope with soft-WorRx version 4.1 (GE Healthcare; Issaquah, WA) equipped with a 60 x/1.4 NA Plan-Apo lens, solid-state illuminator, and 12-bit charge-coupled device (CCD) camera. Fluorescent proteins were excited and detected with the following filter sets and exposure times: GFP: excitation (ex)475/28, emission (em)525/50, 0.15 s; DrsRed or mCherry or red fluorescent protein (RFP): (ex)575/25, (em)632/60, 0.08 s; cerulean or cyan fluorescent protein (CFP): (ex)438/24, (em)470/24, 0.15 s; yellow fluorescent protein (YFP): (ex)513/17, (em)559/38, 0.15 s. The following polyacrylamide gels were used: GFP/mCherry Chroma ET C125705 approximately: 520/50–630/80; Semrock CFP/YFP/DsRed 61008 bs approximately: 415/20–462/32–535/5–635/74. Long-term time-lapse videos used 13 0.5-µm z-sections. Images were deconvolved and maximum-intensity projected (softWoRx). Projected fluorescence images were combined with transmitted light images. Images were contrast adjusted using an equivalent histogram stretch on all samples.

GFP focus discrimination was performed by applying a signal threshold twice as large as the average nuclear background. When examining the spatial and temporal dynamics of fluorescently tagged proteins, an event was determined to have ceased if it failed to occur again after three consecutive frames. Frames were taken every 10 min. Measurement of GFP intensity was carried out using ImageJ. Circles of constant size were overlaid on undivided or divided nuclei to measure mean GFP intensity. Background signal was further removed to calculate net GFP signal intensity using the formula: net intensity = gross intensity − [mean background intensity × nuclear area]. At least two independent experiments and five fields per experiment were assessed for each sample. Cell counts were pooled and presented as proportions ±95% confidence interval, which was calculated using sum-squared error rules. Significance was evaluated using a chi-squared test or Fisher’s exact test, depending on sample size. For continuous GFP-intensity data in Figures 6–8, significance was determined by calculating the area under the curve of each line and then using one-way ANOVA with subsequent Tukey’s family-wise comparisons for all genotypes. Similarly, but for GFP-foci loss data in Figures 6–8, a chi-squared test for trend was employed.

**Flow cytometry and microscopy**

Whole-cell FACS was carried out as described in detail in Sabatinos and Forsburg (2009, 2015) and Sabatinos et al. (2015) with minor changes. To perform cell cycle analysis or microscopy, cells were fixed in 70% ethanol. Following rehydration in 50 mM sodium citrate, cells were treated with 1 µM SytoxGreen (Invitrogen, Carlsbad, CA) plus 10 µg/ml RNase A and incubated at 36°C for 1–2 h. Samples were then sonicated before being analyzed on a FACSScan machine (BD Biosciences, San Jose, CA). DAPI staining was done by first rehydrating fixed cells in PBS and placing them to dry on positively charged glass slides. Mount solution (50% glycerol and 1 µg/ml DAPI) was then applied before cells were photographed on a Leica DMR wide-field epifluorescence microscope equipped with a 63x/1.6 NA Plan-Apo objective lens, 100-W Hg arc lamp for excitation, and a 12-bit Hamamatsu ORCA-100 CCD camera. Images were acquired using OpenLab version 3.1.7 (ImproVision, Lexington, MA) software and analyzed with ImageJ.

**Chromatin immunoprecipitation**

ChiP was performed as described in Rougemaille et al. (2008) with minor changes. Briefly, synchronous meiosis was achieved as mentioned above. Cells were harvested every hour, treated with 37% formaldehyde (1% final volume) followed by quenching with 0.25 M glycine, and washed in Tris-buffered saline (TBS) before being stored at −80°C. Pelleted cells were lysed by bead-beating in a FastPrep machine (MP Biomedicals, Santa Ana, CA), and the resulting lysate was sonicated four times for 15 s (15% duty cycle) with a Branson Sonifier 450 microtip sonicator. Protein concentration was determined using the Bradford method. Some of the lysate was stored as input, while the rest was treated with 1:200 mouse anti-GFP antibody (Abcam 290) (Abcam, Cambridge, MA) and later combined with protein A Dynabeads (Invitrogen, Carlsbad, CA). After several washes (in lysis buffer, wash buffer, and Tris-EDTA), the protein–DNA portion was separated from the Dynabeads using TE and 1% SDS for 30 min, followed by centrifugation. To reverse the formaldehyde cross-link, the resulting elution was then incubated at 65°C for at least 12 h (input included). Afterward, 2 mg/ml proteinase K (Bioline Reagents, Taunton, MA) was used for 2 h at 37°C to eliminate any remnant protein from the samples. DNA was isolated with a Qiagen PCR Purification Kit and analyzed by Q-PCR using a CFX Connect Real-Time System (Biorad, Hercules, CA) software. Fold values were calculated using the 2−ΔΔCt (Livak) method (Livak and Schmittgen, 2001). To determine fold change in cohesin, the input and the immunoprecipitated fractions were used as the calibrator and test parameters, respectively, while act1 was used as the reference gene (testing for Rec8-GFP localization at the euchromatin) and the centromeric dg locus as the target locus (testing for Rec8-GFP localization at the heterochromatin-rich centromere). Significance was determined using a Wilcoxon signed-rank test.
ACKNOWLEDGMENTS

We thank all current members of the Forsburg lab, Amanda Jensen, Ruben Petreca, Nimra Ranatunga, Oscar Aparicio, Sean Curran, and Matthew Michael for technical support and early review of this article. Our appreciation also goes to Tara Mastro, Marc Green, and Sarah Sabatinos for their assistance with PFGE, ChIP, live-cell imaging, and statistical analysis. We also extend our gratitude to Gerald Smith and Yoshinori Watanabe for providing us with strains used to analyze cohesin regulation. This work was made possible by National Institutes of Health grants R35 GM118109 and R01 GM081418 to S.L.F.

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