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Establishment of Cohesion at the Pericentromere by the Ctf19 Kinetochore Subcomplex and the Replication Fork-Associated Factor, Csm3

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Abstract

The cohesin complex holds sister chromatids together from the time of their duplication in S phase until their separation during mitosis. Although cohesin is found along the length of chromosomes, it is most abundant at the centromere and surrounding region, the pericentromere. We show here that the budding yeast Ctf19 kinetochore subcomplex and the replication fork-associated factor, Csm3, are both important mediators of pericentromeric cohesion, but they act through distinct mechanisms. We show that components of the Ctf19 complex direct the increased association of cohesin with the pericentromere. In contrast, Csm3 is dispensable for cohesin enrichment in the pericentromere but is essential in ensuring its functionality in holding sister centromeres together. Consistently, cells lacking Csm3 show additive cohesion defects in combination with mutants in the Ctf19 complex. Furthermore, delaying DNA replication rescues the cohesion defect observed in cells lacking Ctf19 complex components, but not Csm3. We propose that the Ctf19 complex ensures additional loading of cohesin at centromeres prior to passage of the replication fork, thereby ensuring its incorporation into functional linkages through a process requiring Csm3.

Analysis of cohesin distribution along both mitotic and meiotic chromosomes of budding yeast has revealed that the highest levels of cohesin are found in a ~50 kb domain surrounding the ~120 bp centromere sequence, called the pericentromere [3,12–14]. In fission yeast, pericentromeric heterochromatin is important for cohesin association with the pericentromere during mitosis and meiosis [15–18]. Budding yeast lacks pericentromeric heterochromatin but a functional kinetochore is required for pericentric cohesin enrichment [13,19]. The high levels of cohesin in the pericentromere raised a paradox because sister centromeres are known to separate under tension over an approximately 20 kb domain without cohesin cleavage, a phenomenon known as “centromere breathing” [20–22]. A possible solution to the paradox was provided by the observation that exertion of tension across sister kinetochores causes a dramatic reduction in the amount of cohesin associated with this 20 kb domain where breathing is observed [19,23]. However, this region of the pericentromere is thought to form an intramolecular loop, so cohesin cannot be fulfilled by arm cohesins [19], it exhibits unique behavior, the functional relevance of which is unclear.

A clear role for pericentromeric cohesin has, however, emerged in meiosis. In contrast to mitosis, where cohesins are removed along the length of chromosomes simultaneously, meiosis requires that loss of arm cohesins and pericentric cohesins are temporally separated (reviewed in [25]). The Shugoshin protein (Sgo1) localizes to the ~50 kb pericentromeric domain of enriched

Introduction

The accurate transmission of the eukaryotic genome requires that the two copies of each chromosome are held together following their synthesis in S phase until the time of their segregation in mitosis. This chromatid cohesion, which facilitates the biorientation of sister chromatids on the mitotic spindle, is achieved by a multi-subunit complex known as cohesin [reviewed in [1]]. Once proper bipolar attachment is achieved, a protease, separase, cleaves the Scc1/Mcd1 subunit of cohesin and destroys the linkages, thereby triggering the movement of sister chromatids to opposite poles [1].

The establishment of cohesion between sister chromatids is coupled to their replication in S phase. In budding yeast, cohesin is loaded onto chromosomes before DNA replication in a manner dependent on, and at the binding sites of, the cohesin-loading complex Scc2/Scc4 [2,3]. Subsequently, cohesin is thought to translocate from these sites as a result of passage of the transcriptional apparatus [3]. Transformation of this loaded cohesin into functional linkages between sister chromatids requires a second step that takes place during S phase. Scc1 produced after S phase associates with chromosomes but fails to generate cohesion [4]. Several proteins that travel with the replication fork function in this second step. Among the replication fork-associated factors that have been implicated in cohesion function is the Tof1-Csm3 complex which is required for replication fork pausing at replication barriers [5–11]. These observations suggest a tight coupling between cohesion establishment and passage of the replication fork.

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Author Summary

During cell division, chromosomes must be distributed accurately to daughter cells to protect against aneuploidy, a state in which cells have too few or too many chromosomes, and which is associated with diseases such as cancer and birth defects. This process begins with the generation of an exact copy of each chromosome and the establishment of tight linkages that hold the newly duplicated sister chromosomes together. These linkages, generated by the cohesin complex, are essential to resist the pulling forces of the spindle, which will pull the sister chromosomes apart into the two new daughter cells. Here we examine the establishment of cohesin at the pericentromere, the region surrounding the site of spindle attachment and where its forces are strongest. We find that a dedicated pathway promotes cohesin establishment in this region through a two-step mechanism. In the first step, a group of proteins, known as the Ctf19 complex, promote the association of cohesin with this region. In the second step, the Csm3 protein, which is coupled to the DNA replication machinery, ensures its conversion into functional linkages. We demonstrate the importance of this process for accurate chromosome segregation during cell division.

cohesin and protects it from separase-dependent cleavage [14,26–29]. This pericentromeric cohesin is essential for the accurate segregation of sister chromatids during meiosis II. Two kinetochore proteins, Iml3 and Chl4, are important for preventing non-disjunction of sister chromatids during meiosis II [27,30] and for proper localization of Sgo1, a role shared with cohesin [14]. Iml3 and Chl4 are components of a conserved kinetochore subcomplex, known as the Ctf19 complex (see [31] for review) (Figure 1A). Ctf19 is required for the enhancement of cohesin in the pericentromere [19], but whether this function is shared with other members of the complex was not known. Similarly, Csm3, and its fission yeast counterpart, Swi3, have been implicated in pericentromeric cohesin through screens in both meiosis and mitosis [9–11,32].

Here we further investigate the role of Csm3 and the Ctf19 complex in the establishment of cohesin at the pericentromere. We show that two of the more peripheral Ctf19 complex proteins, Iml3 and Chl4, directly increase cohesin association with the pericentromeric region during both meiosis and mitosis. In the absence of Iml3 and Chl4, cohesin binding at the pericentromere is reduced and this has important implications for sister chromatid cohesion and chromosome segregation. Conversely, we find that the replication-fork associated protein, Csm3, is not required for cohesin association with the pericentromere, but plays a role in cohesion establishment at the pericentromere that is non-overlapping with Iml3 and Chl4. Our results indicate that Iml3 and Chl4 ensure the association of cohesin with the pericentromeric region, to facilitate its subsequent incorporation into functional linkages through a replication-coupled step, which requires Csm3.

Results

Iml3 and Chl4 direct increased Scc1 binding at the pericentromere during mitosis

Iml3 and Chl4 are peripheral components of the Ctf19 kinetochore subcomplex [33,34] (Figure 1A). We asked if Iml3 and Chl4 share the reported role of Ctf19 [19] in promoting the enhancement of cohesin in the pericentromere. We examined the localization of the cohesin subunit, Scc1, at 11 sites on chromosome IV (Figure 1B) by chromatin immunoprecipitation (ChIP) followed by real-time quantitative PCR (qPCR). For the purposes of this paper “centromere” refers to the ~120 bp region where the kinetochore assembles, “pericentromere” we define as the ~50 kb enriched cohesin domain and “inner pericentromere” describes the domain in which “breathing” and cohesin removal is observed under tension. We first treated cells with the microtubule-depolymerizing drugs, nocodazole and benomyl, to generate a metaphase arrest where sister kinetochores are not under tension. This has been previously shown to result in high levels of cohesin association with centromeric and pericentromeric regions in wild type cells [19,23]. In wild type cells (Figure 1C, Scc1–6HA, black bars), Scc1 levels were high at cohesin-enriched chromosomal arm sites (A1, A3), but low at the cohesin-poor A2 site. Scc1 association with all pericentromeric sites was increased in wild type cells over the no tag control, although we observed particularly high enrichment at the centromere (C1, C2) and pericentromeric sites P2–4 (Figure 1C). In iml3A and chl4A mutants, Scc1 levels at chromosomal arm sites (A1–3) were comparable to that seen in wild type cells (Figure 1C), indicating that Iml3 and Chl4 do not affect cohesin association with chromosome arms. In contrast, Scc1 was reduced at centromeric (C1, C2) and pericentromeric (P2–4) sites and the enrichment over chromosome arms was lost. Similar results were obtained for chromosome V (Figure S1). Therefore, the role of Ctf19 in promoting assembly of a cohesin-rich pericentromeric domain is shared with other components of the complex.

To investigate the effect of tension across sister kinetochores on cohesin association with the pericentromere in cells lacking Iml3 or CHL4, we arrested cells by depletion of CDC20 (under control of the methionine-repressible promoter, MET-CDC20; Figure 1D). This generates a metaphase arrest where sister kinetochores attach to opposite poles and are under tension, resulting in a reduction in cohesin levels within the ~20 kb inner pericentromere [19,23]. Again Scc1 association on chromosomal arm sites was unchanged in iml3A or chl4A mutants (Figure 1D, A1–3). In contrast to the situation in nocodazole (Figure 1C), however, we observed little enrichment of Scc1 at the centromere (C1, C2) or at the majority of pericentromeric sites (P3–6), even in wild type cells (note the scales in Figure 1C and 1D are different). Outside this region, cohesin is retained upon the exertion of tension in wild type cells (site P2). However, we observed a clear reduction in Scc1 association with this site in iml3A and chl4A mutants in the presence of tension. This indicates that Iml3 and Chl4 may also restrict the region in the pericentromere where cohesin shows tension-dependent association.

Defective pericentromeric cohesin in Ctf19 complex mutants

We investigated the importance of pericentromeric Scc1 for sister chromatid cohesion during mitosis by inducing a metaphase arrest in strains with tetO arrays integrated 2.4 kb from the centromere of chromosome IV and which express TrcR-GFP ( +2.4CEV4-GFP; [21]). Previous reports have indicated that microtubule forces cause sister centromeres to separate specifically at inner pericentromeric regions during metaphase, so that 2 GFP dots are observed in a fraction of wild type cells [20–22]. Wild type, iml3A and chl4A cells were released from a G1 block into a metaphase arrest by depletion of CDC20, either in the presence of nocodazole (in DMSO) to depolymerize microtubules or DMSO alone. Depletion of CDC20 caused a metaphase arrest in all three strains, as judged by spindle morphology in the strains treated with DMSO (Figure S2A) or nuclear morphology in the nocodazole-treated strains (not shown). We scored separation of sister chromatids at CEN4 by monitoring...
Figure 1. Analysis of Sccl localization at the pericentromere and the effect of tension in Ctf19 complex mutants. (A) Schematic diagram of the Ctf19 kinetochore subcomplex adapted from [31,76]. (B) Schematic diagram showing locations of primers used for qPCR analysis of ChIP samples. The hatched area represents the core centromere, and the black, chromosome arms. The ~50 kb pericentromere is split into white, representing the ~20 kb region in which cohesin is displaced and sister chromatids separate under tension. The remainder of the pericentromere in which cohesin is retained under tension is shown in grey. (C) Analysis of Sccl-6HA association in cells arrested in metaphase of mitosis in the absence of microtubules. Strains AM1145 (Sccl-6HA), AM3441 (iml3Δ Sccl-6HA), AM3442 (chl4Δ Sccl-6HA), and AM1176 (no tag) were arrested in medium containing nocodazole and benomyl for 3 h to depolymerize microtubules and induce a metaphase arrest. (D) Analysis of Sccl-6HA association in cells arrested in metaphase of mitosis with sister kinetochores under tension. Strains AM1105 (Sccl-6HA), AM3948 (iml3Δ Sccl-6HA), AM3950 (chl4Δ Sccl-6HA), and AM2508 (no tag control) all carrying MET-CDC20 were arrested in G1 using alpha factor and released into medium containing methionine to deplete CDC20. Cells were harvested 2 hours after release from G1 and metaphase arrest confirmed. The mean values from three independent experiments are shown with error bars indicating standard deviation in (C) and (D).

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the numbers of cells in which 2 GFP dots were visible (Figure 2A). Consistent with previous reports [20–22], we observed separation of sister +2.4CEN4-GFP in ~30% of wild type cells in a manner dependent on microtubules (Figure 2A). This fraction was increased to ~60% in iml3A and chl4A mutants and was also microtubule-dependent (Figure 2A). This indicates that Iml3 and Chl4 are required to generate cohesion at the pericentromere that is effective in resisting microtubule pulling forces.

Ctf19 complex mutants exhibit non-equivalent chromosome loss rates but similar cohesion defects

Ctf19 complex components are required for accurate chromosome transmission during mitosis [33,35–40]. We used a colony-sectoring assay [41] to measure chromosome loss in mutants lacking Ctf19 complex components or Csm3. All mutants tested, with the exception of <i>ubp1A</i> and <i>ubp2A</i>, exhibited chromosome loss rates that were elevated compared to the wild type, however the frequencies were not equivalent (Table S1). The highest chromosome loss rates were observed in mem21A and ctf19A mutants, with other mutants exhibiting lower, but non-equivalent rates of loss.

We compared cohesion defects in Ctf19 complex mutants by examining the separation of +2.4CEN4-GFP foci in cells arrested in metaphase by CDC20 depletion. Figure 2C shows that 120 mins after release from G1, GFP foci separation was similarly increased over wild type in all single and double mutant combinations tested (see also representative time courses Figure S3), despite their different chromosome loss rates. Previous analyses have revealed that the association of Iml3, Chl4, Ctf3, Mcm16 and Mmm22 with kinetochores depends on Ctf19, but not <i>niecens</i> [33,39]. We found that Chl4 also fails to associate with CEN4 in a mem21A mutant and its levels are reduced in both mem16A and iml3A mutants (Figure S4). These findings suggest that Ctf19 and Mcm21 could mediate cohesion establishment through recruitment of the more peripheral components of the sub-complex.

Increased distance between sister centromeres in Ctf19 complex mutants

We next asked if the distance between the separated centromeres is increased in <i>iml3A</i>, <i>chl4A</i> and <i>ctf3A</i> mutants arrested in metaphase by deletion of <i>CDC20</i>. An <i>SPC42-Tomato</i> construct was used to visualize spindle pole bodies (SPBs) together with +2.4CEN4-GFP (Figure 2B) and directly monitor the accumulation of cells in metaphase (Figure S2B) alongside sister centromere separation (Figure S2C). Measurement of distances between separated +2.4CEN4-GFP foci from the 60–120 min time points revealed that the majority of separated sister centromeres were greater than 0 but less than 1 μm apart in wild type cells, consistent with previously published observations (Figure 2D) [21]. However, in <i>iml3A</i>, <i>chl4A</i> and <i>ctf3A</i> mutants, most of the separated sister centromeres were greater than 1 but less than 4 μm apart. Perhaps as a consequence of this increased centromere stretching, we found that the distance between SPBs was also increased in <i>iml3A</i>, <i>chl4A</i> and <i>ctf3A</i> mutants compared to wild type cells (Figure 2E). We conclude that both the frequency and distance of sister centromere separation is increased at metaphase in the absence of Ctf19 complex components.

The Ctf19 complex restricts the domain around the centromere at which sister chromatids separate at metaphase

To ask if the cohesion defects observed in Ctf19 complex mutants at CEN4 also apply to other centromeres we used GFP labels in which <i>tetO</i> arrays are integrated 4.5 kb from CEN6 (Figure 2F) or 1.4 kb from CEN5 (Figure 2G; [22]). Figure 2F shows that CEN6-GFP behaves in a manner reminiscent of +2.4CEN4-GFP in both wild type and <i>iml3A</i> cells. That is, sister centromeres separated at an appreciable frequency in wild type cells (~50%), but with a greater occurrence (~75%) in the <i>iml3A</i> mutant cells. CEN5-GFP, however, exhibited a different behavior, because sister centromere separation was nearly complete (~90%) in wild type metaphase-arrested cells, perhaps as a result of the close proximity of this label to the centromere [2] and no increase in CEN5-GFP separation was observed in <i>iml3A</i>, <i>chl4A</i> or <i>ctf3A</i> mutants (Figure 2G). Because the Ctf19 complex is required for normal levels of the cohesin Scc1 at the pericentromere of chromosome V during mitosis (Figure S1) we reasoned that cohesion defects might be apparent at loci more distant from CEN5. Indeed, two intermediate sites in the pericentromere, at ~12.6 kb and ~17.8 kb to the left of CEN5 (~17.8CEN5-GFP and ~12.6CEN5-GFP, respectively), which split rarely in wild type cells [21], showed an increased separation in the absence of <i>IML3</i> or <i>CHL4</i> (Figure 2H and 2I). However, cohesion at a site just outside the pericentromere (URA3-GFP, 38.4 kb from CEN5 [42]) was virtually unaffected in <i>iml3A</i> and <i>chl4A</i> mutants (Figure 2J) and splitting of GFP signals at a telomeric locus on chromosome V (TEL5-GFP, was not observed in any of the strains (Figure 2K).

This is consistent with our ChIP results and indicates that arm cohesion is intact in <i>iml3A</i> and <i>chl4A</i> mutants. Our results suggest that the Ctf19 complex directs cohesion establishment at the pericentromere of budding yeast chromosomes and that this is essential to restrict the region where sister chromatids separate when their kinetochores are under tension.

The Ctf19 complex also directs cohesion establishment during meiosis

Pericentromeric cohesion is particularly important during meiosis. Following the segregation of homologous chromosomes during meiosis I, sister chromatids lack all arm cohesion and are solely reliant on pericentromeric cohesion. This property of meiosis exposes weak general cohesion defects [43] and also led us to identify <i>IML3</i> and <i>CHL4</i> as important mediators of pericentromeric cohesion [27]. To assess the importance of other Ctf19 complex components for meiosis II, we examined the fate of a GFP label at URA3 (38.4 kb from CEN5) on either one or both copies of chromosome V (heterozygous or homozygous GFP dots, Figure 3A and 3B, respectively) after meiosis. Examination of the fate of heterozygous dots after meiosis II revealed that sister GFP labels failed to segregate into different nuclei in approximately 20% of tetranucleate cells of all Ctf19 complex mutants tested, with the exception of <i>ubp1A</i> and <i>ubp2A</i> (Figure 3C). Meiosis II non-disjunction in Ctf19 complex mutants with homozygous GFP dots was also evident from the high frequency of tetranucleate cells with a GFP dot in just 3 of the 4 spores, which was comparable also in <i>iml3A chl4A</i> or <i>chl4A ctf3A</i> double mutant combinations (Figure 3D). Although most of the mutants with homozygous GFP dots showed only the expected modest number of tetranucleate cells with GFP label in just two spores, <i>ctf3A</i> and <i>mem21A</i> mutants were a notable exception (Figure 3E). Examination of GFP dot segregation in binucleate cells (Figure S5) revealed that Ctf19 and Mcm21, but not other Ctf19 complex components, are also required for accurate segregation during meiosis I. These findings are consistent with our conclusion from the mitotic chromosome loss data (Table S1) that Ctf19 and Mcm21 promote accurate chromosome segregation also in ways other than cohesion establishment.

To address whether meiosis II non-disjunction in Ctf19 complex mutants could be due to a failure to enrich cohesin in
Figure 2. Ctf19 complex mutants exhibit pericentromeric cohesion defects in mitosis. (A) Sister centromeres (+2.4CEN4-GFP; 2.4 kb to the right of CEN4) are more frequently separated in a metaphase arrest in iml3Δ and chl4Δ mutants and this depends on microtubules. Strains of wild type (AM914), iml3Δ (AM3522), and chl4Δ (AM3501), all carrying +2.4CEN4-GFP (2.4 kb to the right of CEN4) and MET-CDC20 were arrested in alpha factor and then released into medium containing methionine to deplete CDC20 and either DMSO (control) or nocodazole (to depolymerize microtubules). Samples were taken at the indicated times after release from G1 for anti-tubulin immunofluorescence (Figure S2) and to determine the number of GFP foci per nucleus. Percentages of cells with two GFP dots are shown for wild type (black squares), chl4Δ (blue circles), and iml3Δ (red triangles) treated with either DMSO (open symbols) or nocodazole (closed symbols). 200 cells were scored at each time point. (B) Cohesion defects were
examined after releasing cells carrying +2.4CEN4-GFP, SPC42-tdTomato (to label spindle pole bodies) and MET-CDC20 from a G1 block, as a result of alpha factor treatment, into medium containing methionine to deplete CDC20. A snapshot of cells at the 120 min time point (metaphase arrest) is shown as an example together with a schematic diagram of chromosome IV with sister centromeres under tension (left). SPBs are shown in red and +2.4CEN4-GFP is shown in green. The black and white arrowheads indicate cohesed and separated +2.4CEN4-GFP foci, respectively. (C) Summary of frequency of +2.4CEN4-GFP separation at metaphase in Ctf19 complex mutants. Values are the mean of all experiments where percentages of separated sister centromeres were determined after scoring 200 cells. 120 min after release from G1, in a metaphase arrest. Error bars indicate standard error. Strains, and number of independent repeats (given in italics), were: wild type (AM4643); 14, chl4D (AM4644); 10, iml3D (AM4647); 7, ctf3D (AM4683); 3, mcm16D (AM6158), 2, mcm22D (AM1660); 2, ctf19D (AM5786); 3, mcm21D (AM5788); 3, chl4D mcm16D (AM6195); 1, chl4D mcm22D (AM6192); 1, and csm3D (AM4717); 6. (D,E) Sister centromeres and SPBs are further apart at metaphase in iml3D, chl4D, and ctf3D mutants. The distance between +2.4CEN4-GFP (D) and SPC42-tdTomato (E) foci was determined for all separated foci from the 60–120 minute time points. For wild type (AM4643), chl4D (AM4644), and mcm16D (AM4647), the average of two independent experiments with error bars representing standard deviation is shown. For ctf3D (AM4683), measurements are from a single experiment. (D) The percentage of +2.4CEN4-GFP foci that were >0–1 μm (blue) or >1–4 μm (red) apart. n = 115/82 (wild type); 202/139 (chl4D); 184/167 (iml3D); 199 (ctf3D). (E) The percentage of SPC42-tdTomato foci that were >0–3 μm (blue) or >3–6 μm (red) apart is shown. n = 338/305 (wild type); 548/244 (chl4D); 331/272 (iml3D); 313 (ctf3D). The percentages of separated spindle pole bodies or +2.4CEN4-GFP foci shown in Figure S2 for this experiment. (F–K) The cohesion defect in iml3D and chl4D mutants is not chromosome-specific but is restricted to centromere-proximal regions. Cells of the indicated genotypes carrying MET-CDC20, SPC42-tdTomato and with GFP labels at various loci were arrested in G1 with alpha factor and released into medium containing methionine and the percentages of separated GFP foci were scored at the indicated timepoints. Tet operators are integrated at: (F) +4.5CEN6-GFP (4.5 kb to right of CEN6) in wild type (AM5329) and iml3D (AM5330); (G) +1.4CEN6-GFP (1.4 kb to right of CEN6) in wild type (AM5189), iml3D (AM5249), chl4D (AM5251), and ctf3D (AM5188); (H) +1.4CEN6-GFP (1.26 kb to left of CEN6) in wild type (AM5345), iml3D (AM5342), and chl4D (AM5560); (I) +1.7CEN5-GFP (1.78 kb to left of CEN5) in wild type (AM5333), iml3D (AM5337) and chl4D (AM5551); (J) URA3-GFP (38.4 kb to left of CEN6) in wild type (AM1081) and iml3D (AM3541) and chl4D (AM3519). (K) -3ORTELS (~30 kb from telomere on right arm of chromosome V) in wild type (AM2511) and iml3D (AM3887) and chl4D (AM3942). Note that the URA3-GFP and TELS-GFP strains did not carry SPC42-tdTomato. Distances indicate the start of the tetO array from the centromere.

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the pericentromere, as in mitosis, we examined localization of the meiosis-specific counterpart of Scc1, Rec8, during meiosis. Wild type, iml3D and chl4D cells were arrested in metaphase I (by depletion of CDC20 [44]), and the localization of the meiotic cohesin, Rec8, was examined at 10 sites on chromosome IV (Figure 3E) by ChIP followed by qPCR. Because sister kinetochores are uniquely mono-oriented during meiosis I (Figure 3F), this leads to a situation where they are not under tension, enabling the retention of high levels of cohesin in the centromere and pericentromere. Consistent with previous reports, Rec8 localization closely resembled that of the mitotic cohesin, Scc1, in wild type cells [12,14]. As in mitosis, deletion of IML3 or CHL4 caused a reduction in Rec8 levels at the centromere (G1–5) and pericentromeric sites (PI–6), but not at chromosomal arm sites (A1–3). We also observed a similar requirement for Ctf19 in localizing Iml3 and Chl4 during meiosis, and a partial dependence on Mcm16 and Mcm22 (Figure S6). Furthermore, we found that Iml3 and Chl4 are localized specifically at the core centromere, despite their ability to influence cohesion in the surrounding pericentromere (Figure S6). We conclude that the Ctf19 complex plays similar roles in pericentromeric cohesion establishment during mitosis and meiosis.

Non-overlapping roles of Csm3 and the Ctf19 complex in cohesion

To understand more about cohesion establishment in the pericentromere, we extended our analysis to Csm3, which is required for proper chromosome segregation in meiosis [10] and has been implicated in cohesion establishment during mitosis [9,11]. Furthermore, Swi3, the fission yeast homolog of Csm3, was shown to have a role in cohesion establishment [32]. Examination of +2.4CEN4-GFP separation in csm3D cells arrested in metaphase by CDC20 depletion confirmed that csm3D mutants exhibit a cohesion defect that is more severe than that of Ctf19 complex mutants (Figures 2C and 4A). Consistently, csm3D cells showed a more pronounced cohesion defect than chl4D cells at loci 12.6 and 17.8 kb away from CEN5 (+12.6CEN5-GFP and +17.8CEN5-GFP; Figure 4B and 4G, respectively). Furthermore, in contrast to chl4D mutants and in agreement with previous reports [9,11], cells lacking CMS3 exhibited a small defect in cohesion at the URA3-GFP locus (Figure 4E). Examination of URA3-GFP separation in csm3D iml3D double mutants, revealed an additive effect (Figure 4E and 4F). Similarly, we observed additive effects on meiotic chromosome segregation in csm3D iml3D mutants (Figure S7). These results indicate that Csm3 plays a role in cohesion establishment that is non-overlapping with the Ctf19 complex and which may not be restricted to the pericentromere.

Csm3 acts after cohesin loading and may couple cohesion establishment to replication fork progression

To ask if the cohesion defects of csm3D mutants could, like that of Ctf19 complex mutants, be due to failure to recruit proper levels of cohesin to chromosomes, we examined the localization of Scc1 in nocodazole-arrested csm3D mutants (Figure 4G and 4H). However, we found that Scc1 levels in the csm3D mutant were comparable to wild type at all sites tested, unlike a Ctf19 complex mutant control, ctf3D (Figure 4H). This indicates that Csm3 does not promote cohesion establishment by directing cohesin association with chromosomes.

Csm3 is part of a complex with Tof1 that travels with the replication fork and is required for fork stalling at DNA-protein linkers, including centromeres [5–8]. Csm3 is thought to achieve fork stalling by counteracting the RRm3 helicase and deletion of RRm3 restores fork stalling in csm3D mutants at replication termination sites [45]. If Csm3 facilitates cohesion establishment by promoting fork stalling, we reasoned that the cohesion defect might also be rescued in csm3D rm3D mutants. Indeed, deletion of RRm3 in csm3D mutants reduced the frequency of +2.4CEN4-GFP separation at metaphase (Figure 4I and 4J), although not to wild type levels. Interestingly, the rm3D single mutant exhibited a lower frequency of GFP dot separation that wild type (Figure 4I and 4J). Since rm3D mutants exhibit increased fork stalling, particularly at hard to replicate sites such as centromeres [46], these results are consistent with the idea that fork stalling facilitates cohesion establishment.

Delayed and inefficient loading of cohesin at centromeres in chl4D mutants

Next we investigated the role of the Ctf19 complex in pericentromeric cohesin recruitment in more detail. Our results show that while cohesin is reduced at the pericentromere of iml3D A
Figure 3. The Ctf19 complex promotes cohesion establishment in the pericentromere during meiosis. Schematic diagram showing the segregation of chromosome V with either heterozygous (A) or homozygous (B) GFP labels in wild type cells. Homologs are shown in white and grey and the GFP dot is shown as a black circle. Sister chromatid cohesion is represented by black lines between sister chromatids. Arrows on kinetochores indicate the direction of attachment to the spindle. (C,D) Ctf19 complex mutations lead to errors in chromosome segregation during meiosis. Cells of the indicated genotypes in which either one copy (C; heterozygous) or both copies (D; homozygous) of chromosome V were marked with GFP (tetR-GFP::LEU2 URA3::tetOx224) at URA3 (38.4 kb from the centromere) were induced to sporulate at 30°C. The percentage of tetrancleate cells with the patterns of GFP dot segregation shown was determined after examining 200 cells, 8 hours after transfer into sporulation medium. Strains used were AM107 (wild type), AM1904 (iml3Δ), AM1902 (chl4Δ), AM1903 (iml3Δ, chl4Δ), AM1905 (chl4Δ), AM3684 (mcm22Δ), AM3685 (mcm16Δ), AM3799 (mcm16Δ), AM3661 (mcm22Δ), AM3798 (ctf19Δ), AM347 (chl4Δ, mcm16Δ), and AM5809 (chl4Δ, mcm22Δ, ctf19Δ) (D). Values shown are the mean from two independent experiments, with error bars representing standard deviation, with the exception of mcm22Δ, nkp1Δ, nkp2Δ (C) and mcm22Δ and chl4Δ ctf19Δ (D), where results are shown from a single experiment. (E,F) Reduced levels of the meiotic cohesin, Rec8, at the pericentromere in Ctf19 complex mutants. (E) Schematic diagram of the primer sets used for qPCR on chromosome IV. (F) Analysis of Rec8-3HA association in cells arrested in metaphase I in meiosis. Strains AM3560 (no tag control), AM3375 (REC8-3HA), AM3422 (chl4Δ REC8-3HA) and AM3377 (iml3Δ REC8-3HA) carrying the pCLB2-3HA-CDC20 allele were harvested for ChIP 8 h after transfer into sporulation medium. The schematic diagram shows the configuration of the chromosomes. qPCR analysis of anti-HA ChIP samples was performed using primers shown in (E).

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Figure 4. Csm3 promotes cohesion establishment at the pericentromere but is not required for cohesin enrichment in this region. (A–E) csm3Δ mutants show cohesion defects in mitosis that are additive with iml3Δ. Strains carrying the indicated GFP labels and MET-CDC20 were released from G1 into a metaphase arrest by CDC20 depletion, and the number of GFP foci per nucleus was determined for 200 cells. Strains with +2.4CEN4-GFP, -12.6CEN5-GFP, and -17.8CEN5-GFP also carried SPC42-tdTomato. (A) Separation of +2.4CEN4-GFP foci in wild type (AM4643), csm3Δ (AM4717), and ctf3Δ (AM4683). (B) Separation of -12.6CEN5-GFP foci in wild type (AM5545), csm3Δ (AM5569), and chl4Δ (AM5560). (C) Separation of -17.8CEN5-GFP foci in wild type (AM5533), csm3Δ (AM5564), and chl4Δ (AM5551). (D) Separation of URA3-GFP foci in wild type (AM1081), chl4Δ (AM5319), and csm3Δ (AM5312). (E) Separation of URA3-GFP foci in wild type (AM1081), iml3Δ (AM3541), csm3Δ (AM5312), and csm3Δ iml3Δ (AM5796). (F) Summary of frequency of URA3-GFP foci separation at the 120 min timepoint. Values are the mean from 3 (wild type, iml3Δ, csm3Δ) or 2 (iml3Δ, csm3Δ) experiments with error bars indicating standard error. For each experiment 200 cells were scored. (G,H) Csm3 is not required for cohesin enrichment in the pericentromere. Strains AM1145 (SCC1-6HA), AM4927 (ctf3Δ SCC1-6HA), AM3757 (csm3Δ SCC1-6HA), and AM1176 (no tag) were arrested in medium containing nocodazole and benomyl for 3 h to depolymerize microtubules and induce a metaphase arrest. Primers at locations depicted in (G) were used for analysis of Scc1-6HA association by qPCR after ChIP (H). (I) Deletion of RRM3 can partially rescue the cohesion defect of csm3Δ mutants. Separation of +2.4CEN4-GFP foci in wild type (AM4643), csm3Δ (AM4717), rmr3Δ (AM6068), and csm3Δ rmr3Δ (AM6066) in a representative experiment (I) and mean values at 120 mins from 3 independent experiments with error bars indicating standard error (J).

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Chl4 directs cohesin enrichment at the pericentromere during metaphase

Previous observations have indicated that a pathway that promotes cohesin loading at the centromere might be particularly active in mitotic cells, although the significance of this is unknown [19,23]. Furthermore, the removal of microtubule forces in metaphase-arrested cells allows cohesin to re-associate with the pericentromere apparently independently of the Scc2/4 complex that is normally responsible for the loading of cohesin at sites throughout the genome [23]. To ask if the Ctf19 complex directs the accumulation of cohesin at the pericentromere during metaphase, we arrested wild type and chl4Δ cells carrying MET-CDC20 and SCC1-6HA were released from a G1 arrest into methionine and nocodazole at 18°C and samples were taken at 15 min intervals for analysis of Scc1 localization at 4 sites on chromosome IV (Figure 5A). FACS analysis indicated that both strains entered into S phase after 60 min under these conditions (Figure S8). In wild type cells, cohesin associated with the two centromeric (C1 and C2) sites after only 30 min (Figure 5B and 5C). Cohesin began to be recruited to the chromosome arm (A1) and pericentromeric (P2) sites a little later, at 45 min, in wild type cells (Figure 5D and 5E). The early loading of cohesin at centromeres is consistent with a specialized mechanism for cohesin recruitment operating in this region. In chl4Δ cells, cohesin loaded onto the chromosome arm (A1) site with the same kinetics as in wild type cells (Figure 5E), but was barely detectable at the two centromeric (C1 and C2) sites, even after 75 min (Figure 5B and 5C). Cohesin recruitment was also both delayed and inefficient at the pericentromeric (P2) site in chl4Δ mutants (Figure 5D). These findings suggest that Chl4 is required for the loading of cohesin at the centromere. Furthermore, the almost complete absence of cohesin at centromeres as chl4Δ cells enter S phase explains why cohesion establishment fails, given that centromeres replicate early in S phase [47]. It follows that the low, but appreciable, levels of pericentromeric cohesin observed in nocodazole-arrested chl4Δ mutants (Figure 1C) must arrive after passage of the replication fork and therefore may not be functional in holding sister chromatids together.

Figure 5. Chl4 directs cohesin association with the pericentromere throughout the cell cycle. (A–E) Time course of cohesin loading in the cell cycle. Wild type (AM4226) and chl4Δ (AM4291) strains carrying MET-CDC20 and SCC1-6HA were arrested in G1 with alpha factor then released into medium containing methionine and nocodazole at 18°C. Samples were taken at the indicated timepoints and SCC1-6HA association was analyzed by ChIP and qPCR. (A) Schematic diagram showing the experimental setup. (G,H) Strains AM4084 (pGAL-SCC1-3HA) and AM5974 (chl4Δ pGAL-SCC1-3HA) were treated as described in (G,H) except that cells were arrested in metaphase by CDC20 depletion in raffinose, rather than glucose medium, and galactose was added to half of the culture, together with nocodazole, to induce SCC1-3HA expression. (I) qPCR analysis after ChIP prior to nocodazole and galactose addition. (J) qPCR analysis after ChIP following microtubule depolymerization and SCC1-3HA induction.
with nocodazole to depolymerize microtubules, thereby eliminating tension, and re-examined cohesin association with these sites (Figure 5H). As expected, cohesin levels did not increase greatly over background upon the eradication of tension at the chromosomal arm site (A1) or the pericentromeric site (P2) in either wild type or chl4A mutants (Figure 5H). However, cohesin levels increased substantially at both centromeric sites (C1 and C2) in wild type cells after microtubule depolymerization (Figure 5H). In contrast, cohesin accumulation at the centromere during metaphase was more modest in chl4A mutants and did not show enrichment over the chromosomal arm site (Figure 5H).

To distinguish between de novo loading of cohesin at the pericentromere and redistribution of cohesin from other chromosomal sites, we restricted our analysis to cohesin that was produced under control of the galactose promoter (pGAL-SCC1-3HA) but with untagged endogenous cohesin. Cells were released from G1 in the presence of raffinose (to prevent pGAL induction) and methionine (to deplete CDC20) to allow cells to accumulate in metaphase with endogenous (untagged) cohesin and sister kinetochores under tension. After 90 min, anti-HA ChIP was performed to determine the background signal in the absence of SCC1-3HA expression (Figure 5I). Subsequently, nocodazole and galactose were added to depolymerize microtubules and induce expression of SCC1-3HA. After a further 60 min, association of Sccl-3HA at 3 chromosomal sites was analyzed. Figure 5J shows that, in wild type cells, only a low level of metaphase-produced cohesin associated with a chromosomal arm site, whereas levels at pericentromeric and centromeric sites were elevated well above background. This is consistent with a previous report [23] showing that de novo produced cohesin preferentially associates with the pericentromere during metaphase. In the chl4A mutant, however, metaphase-produced cohesin at the C2 and P2 sites was hardly elevated above its levels at the A1 arm site (Figure 5J). These results indicate that ChIl4 contributes to the mechanism that drives cohesin loading at the pericentromere during metaphase, although the purpose and functionality of this cohesin remains unknown.

An S phase delay restores cohesin in iml3A and chl4A mutants, but not csm3A mutants

Next we investigated the relationship between Ctf19 complex-dependent recruitment of cohesin to the pericentromere and DNA replication. Our findings indicate that cohesin is barely above background at the centromere prior to the initiation of DNA replication in chl4A mutants (Figure 5B and 5C), however, during a metaphase arrest without microtubules a low, but appreciable, level of cohesin accumulates in this region (Figure 1C). If cohesion establishment at the pericentromere occurs through the normal replication-coupled mechanism in S phase, then the late arrival of cohesin to this region in chl4A cells would exclude its conversion into functional inter-sister linkages. We reasoned that delaying replication fork passage in Ctf19 mutants might allow this late-arriving cohesin to be converted into functional cohesion. The experimental set-up is shown in Figure 6A. We released wild type, chl4A, iml3A, ctf3A and csm3A cells carrying MET-CDC20 from a G1 arrest into medium containing hydroxyurea (HU) to inhibit DNA replication and containing 8 mM methionine to deplete CDC20. After 90 min in the presence of HU, the drug was washed out and cells were allowed to accumulate in metaphase (as a result of CDC20 depletion). FACS analysis confirmed an approximately 90 min delay in bulk DNA replication in all strains (Figure S9). As expected, strains activated the DNA damage checkpoint kinase, Rad53 in HU [48], as evidenced by its hyperphosphorylation (Figure S10A), but the slowest migrating forms disappeared rapidly upon release in all strains, although in the case of the csm3A there was a slight delay (Figure S10A). Nevertheless, none of the mutants showed sensitivity to HU (Figure S10B), indicating that they are able to respond to and recover from the damage caused. In control cells that were not treated with HU, we observed a cohesion defect at CEN4, in chl4A, iml3A, ctf3A and csm3A mutants, as before (Figure 6B). However, remarkably, delaying cells in S phase by HU treatment reduced the separation of sister +2.4CEN4-GFP foci to near wild type levels in iml3A, chl4A and ctf3A mutants, but not csm3A mutants (Figure 6C). In the HU-treated cells, all mutants separated SPBs with approximately the same kinetics (note that SPB duplication is not prevented by HU treatment) (Figure 6D). Therefore, a HU-induced S phase delay rescues the pericentromeric cohesion defect of iml3A and chl4A and ctf3A mutants, but not csm3A mutants.

We examined the association of Scc1 with chromosomes in wild type, chl4A and csm3A cells arrested in metaphase in the absence of microtubules, following a HU-induced S phase delay. Consistent with our finding that Csm3 does not influence cohesin levels in the pericentromere at metaphase in the absence of an S phase delay (Figure 4H), we observed similar levels of cohesin in csm3A and wild type cells (Figure 6G) after a HU delay. In the absence of CHL4, after the HU delay, cohesin levels were, however, reduced in the centromere and pericentromere to a similar extent as without a HU delay (compare Figure 6G with Figure 1C). We also did not observe increased levels of pericentromeric cohesin in iml3A or chl4A mutants arrested in metaphase with sister kinetochores under tension after HU treatment (Figure 6H). These results rule out the possibility that a HU delay rescues cohesion at the pericentromeres of Ctf19 complex mutants by reversing the cohesin recruitment defect in these mutants. Rather, delaying S phase in Ctf19 complex mutants appears to overcome the failure of the late-arriving cohesin to become cohesive.

Improved cohesion in chl4A and iml3A mutants in the clb5A clb6A background

As a further test of the ability of a replication delay to allow pericentromeric cohesion establishment in Ctf19 complex mutants, we examined the separation of +2.4CEN4-GFP dots in metaphase-arrested cells lacking the S phase cyclins, CLB5 and CLB6, which are known to delay origin firing [49,50] (Figure 7A–7D). FACS analysis revealed a 45 min delay in completion of DNA replication in cells lacking the S phase cyclins compared to wild type, iml3A or chl4A mutants (Figure S11). This delay in DNA replication was increased to 60 min and 75 min in clb5A clb6A mutants lacking IML3 or CHL4, respectively (Figure S11), the reasons for which are unclear. We also observed a corresponding delay in spindle pole body separation in the clb5A clb6A background (Figure 7B). As before, chl4A and iml3A mutants separated +2.4CEN4-GFP foci rapidly and with a greater frequency than wild type (Figure 7A). As expected, due to the SPB separation delay, +2.4CEN4-GFP separation was delayed in cells lacking the S phase cyclins, however, deletion of CHL4 or IML3 did not increase the rate or frequency of separation (Figure 7A). This finding has to be interpreted with caution because clb5A clb6A iml3A and clb5A clb6A chl4A cells show an increased delay in spindle pole body separation (Figure 7B) as compared to clb5A clb6A cells, which could account for slower +2.4CEN4-GFP separation in these cells. However, taking the difference in the timing of SPB duplication into consideration, +2.4CEN4-GFP separation in chl4A or iml3A cells was reproducibly lower in the absence of the S phase cyclins, even though clb5A clb6A cells showed more separation than wild type, using these...
Figure 6. Delaying DNA replication by hydroxyurea treatment rescues the cohesion defect of iml3Δ and chl4Δ, but not csm3Δ, mutants. The experimental setup is shown in (A). Cells were released from G1 into medium containing 8 mM methionine in the presence of HU. After 1.5 h, the HU was washed out and a metaphase arrest was induced either by depletion of CDC20 or by nocodazole treatment. (B–E) Analysis of +2.4CEN4-GFP foci separation after the HU-induced replication delay. Wild type (AM4643), chl4Δ (AM4644), iml3Δ (AM4647), ctf3Δ (AM4683), and csm3Δ (AM4717) strains were arrested in G1 with alpha factor, the cultures were split and released into 8 mM methionine either in the absence (B) or presence (C,D) of HU. Samples were taken at the indicated timepoints after release from G1 for scoring of +2.4CEN4-GFP (B,C) or SPC42-tdTomato (D) separation. FACS analysis was used to confirm a delay in bulk DNA replication in the HU-treated strains (Figure S9). (E) Mean GFP dot separation of 3 experiments at the 120 min time point after washing out HU. Error bars represent standard error. (F–H) Analysis of Scc1-6HA association with chromosome IV during metaphase with sister kinetochores not under tension, after an S phase delay. (F) Location of primer sets used for qPCR analysis. (G) Localization of Scc1 in the absence of microtubules. Strains used were AM1145 (SCC1-6HA), AM 3442 (chl4Δ SCC1-6HA), AM3757 (csm3Δ SCC1-6HA), and AM1176 (no tag). (H) Localization of Scc1 when sister kinetochores are under tension. qPCR analysis of Scc1-6HA ChIP from cells harvested 2 h after HU wash-out. Strains used all carried MET-CDC20 and were AM1105 (SCC1-6HA), AM3948 (iml3Δ SCC1-6HA), AM3950 (chl4Δ SCC1-6HA), and AM2508 (no tag control).

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Figure 7. Replication timing affects cohesion establishment and plasmid maintenance in chl4Δ and iml3Δ mutants. (A–D) The frequency of +2.4CEN4-GFP dot separation in clb5Δ clb6Δ cells is not greatly increased by deletion of IML3 or CHL4. Wild type (AM4643), chl4Δ (AM4644), iml3Δ (AM4647), clb5Δ clb6Δ (AM5351), and clb5Δ clb6Δ chl4Δ (AM5426) carrying MET-CDC20, +2.4CEN4-GFP, and SPC42-tdTomato were arrested in G1 with alpha factor and released into medium containing methionine. Percentages of separated GFP foci (A) and SPBs (B) are shown at the indicated time points for a representative experiment. After 120 min there were 5% (wild type), 3% (chl4Δ), and 1% (iml3Δ) anaphase cells, after which they escape from the metaphase arrest. Escape is delayed until 180 min in the clb5Δ clb6Δ cells, when anaphase cells account for 9% (clb5Δ clb6Δ), 3% (chl4Δ), and 9% (iml3Δ), respectively. Mean +2.4CEN4-GFP (C) and SPB (D) separation is shown from three (120, 135 min) or two (150, 165 min) experiments at the indicated timepoints with error bars representing standard error. (E,F) A plasmid with a late-replicating origin (p12) shows improved stability over a plasmid with an early-replicating origin (p306.10) in chl4Δ and iml3Δ cells. (E) Experimental outline is shown. (F) Ratio of colonies growing on selective medium that depends on the presence of the plasmid (-URA) compared to colonies growing on rich medium for wild type (AM1176), iml3Δ (AM3313), chl4Δ (AM3314), and csm3Δ (AM3194) strains transformed with the indicated plasmids.

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criteria (Figure 7C and 7D, compare time points with asterisks). Therefore, removal of the S phase cyclins appears to at least partially rescue the cohesion defect of iml3A and chl4A cells.

A late firing origin rescues the stability of a centromeric plasmid in iml3A and chl4A cells

How does a replication delay allow the inefficiently loaded cohesin in Ctf19 complex mutants to become cohesive? As hypothesized above, it could restore the normal order of cohesin loading and replication fork passage, to allow cohesion generation through the usual replication-fork coupled mechanism in S phase. However, an alternative mechanism of global cohesion establishment has been described in G2 cells subjected to DNA damage [51–53]. Because HU is a DNA damaging agent and clb5A clb6A mutants have also been found to activate the DNA damage checkpoint [54], cohesion could be restored in iml3A and chl4A mutants through DNA damage-dependent cohesion establishment. To distinguish between the replication delay and DNA damage hypotheses we took advantage of the fact that some DNA replication origins are known to replicate early in S phase, whereas others replicate late. We used centromeric (CENS) plasmids that carry a single origin that fires early (p306.10; ARS306) or late (p12; ARS412) [55,56]. Both plasmids also carry the URA3 selectable marker, which allows growth on medium lacking uracil, enabling us to examine their ability to propagate in wild type, chl4A, iml3A and csm3A cells. Strains containing each of the plasmids were allowed to grow for 2.5 h in non-selective medium before plating onto selective (lacking uracil) and non-selective (rich) medium in parallel (Figure 7E). The ratio of colonies that are able to grow on medium lacking uracil compared to rich medium are given in Figure 7F. Although both plasmids were well maintained in wild type cells, they showed reduced stability in all three mutants. In the case of csm3A cells, we observed an equivalent decrease in stability of both the early and late-replicating plasmid, suggesting that replicating timing does not affect their ability to propagate in this mutant. Replication timing did however influence plasmid stability in chl4A and iml3A cells because the early-replicating plasmid displayed a greatly reduced stability compared to the late-replicating plasmid in these cells. Although we have not directly tested whether the decreased stability of the p306.10 plasmid in iml3A and chl4A mutants is the result of defective cohesion, these findings are consistent with the idea that pericentromeric cohesion establishment fails in cells lacking Ctf19 complex components because cohesin is not recruited prior to the passage of the replication fork.

Discussion

The Ctf19 complex and establishment of pericentromeric cohesion

In budding yeast, an approximately 50 kb region around the centromere, the pericentromere, is enriched for cohesin binding. Enhanced binding of cohesin in the pericentromere requires the ~120 bp centromere sequence and is dependent on a functional kinetochore [13,57]. Previous findings identified Ctf19 as being an important component of the kinetochore in this process [19]. We and others [58] have extended these findings and shown that the Ctf19 kinetochore subcomplex is an important mediator of cohesion establishment at the pericentromere. Since all single and double Ctf19 complex mutants show similar cohesion defects, the critical role of the Ctf19 and Mcm21 components in pericentromeric cohesion establishment may be to recruit the more peripheral components. However, our findings also indicate that Ctf19 complex components may have additional functions in chromosome segregation. In particular, Ctf19 and Mcm21 are unique in their requirement for chromosome segregation also during meiosis I. Whether this reflects a greater general requirement for these proteins in chromosome segregation (Table S1) or a specific role in, for example, generating linkages between homologs is unknown. However, Ctf19 components do not appear to play a major role in mediating kinetochore-microtubule attachments, since sister centromeres are pulled further apart in these cells. In addition, visualization of all kinetochores in Ctf19 complex mutants using Mtw1-GFP revealed no obvious attachment defect [JE and AM, unpublished observations]. We note that human Mcm21 and Ctf19 proteins promote bipolar spindle assembly and chromosome compression, respectively [59] and that yeast Chl4 has been found to play a role in converting naked centromere DNA into an established centromere that is heritable [30]. Whether these functions of the Ctf19 complex are mediated solely through a role in cohesion establishment will be important to investigate in the future.

Mechanism by which the Ctf19 complex promotes cohesin enrichment

How does the Ctf19 complex promote cohesin enrichment within the pericentromere? The simplest explanation is that the Ctf19 complex promotes the loading of cohesin at the centromere, which then spreads bidirectionally into the surrounding pericentromere. This could explain how the Ctf19 complex, which is localized within the ~125 bp core centromere, is able to influence cohesin association over a much greater region. Support for this model comes from our finding that cohesion is detected at pericentromeric sites earlier in the cell cycle than at a pericentromeric site (Figure 3B–3D) and the observation that the Scc2 cohesin loader component is reduced at the centromere, but not other pericentromeric sites in the absence of MCM21 [58]. However, Chl4, at least, may be able to influence cohesin association with the pericentromere independently of Scc2/4. We found that Chl4 is required for high levels of newly-synthesized cohesin to associate with the pericentromere during metaphase after the eradication of tension, a process that appears to be independent of Scc2 [29]. Although the functional relevance of metaphase-loaded cohesin remains unknown, these data provide evidence that the Ctf19 complex promotes cohesin association with the pericentromere throughout the cell cycle.

Csm3 acts at a step after cohesin loading in pericentromeric cohesion establishment

We compared the contribution of Iml3-Chl4 to pericentromeric cohesion establishment with that of Csm3. The finding that cells lacking IML3 and CSM3 show synthetic cohesion defects suggests that these genes promote cohesion establishment through different pathways. Indeed, we observed no defect in cohesin association with the pericentromere in the absence of Csm3. This suggests that Csm3 facilitates cohesion establishment in a step after cohesin loading. Interestingly, Csm3 and its binding partner, Tof1, travel with the replication fork, and the Tof1-Csm3 complex is required for stable fork pausing at protein-DNA barriers including those at centromeres [5,6,45,60–63]. Like Csm3, Tof1 has been implicated in cohesion [9,11]. Tof1-Csm3 is conserved, being homologous to the Timeless-Tipin complex in humans and the Swi1-Swi3 complex in fission yeast [8,9,64]. Furthermore, the fission yeast Swi1-Swi3 complex is also required for the stabilization of stalled forks and efficient cohesion generation [8,32]. Taken together, these observations indicate that the stalling and stabilization of replication forks may be the critical function of Csm3 in cohesion establish-
m. Indeed, we found that deletion of the helicase RRM3, which restores fork stalling to csm3Δ mutants [45], partially rescued the cohesion defect. Perhaps Tol1-Csm3 is required to maintain the association of essential cohesion establishment factors with the replisome upon encountering protein-DNA barriers. Such a hypothesis could explain why cells lacking CSM3 show chromosome segregation defects in meiosis II, but not meiosis I. Perhaps there is a greater requirement for Csm3 in generating cohesion at protein-DNA barriers, such as centromeres, where kinetochore binding could impede replication fork progression. Alternatively, this could simply reflect the susceptibility of meiosis II to general cohesion defects due to the absence of arm cohesion [43].

Cohesion establishment and DNA replication

Our analysis of cohesin association in a synchronized cell cycle showed that cohesin loads at centromeres early and in a Chl4-dependent manner. Given that centromeres replicate early in S phase [47], one important function of kinetochore-driven cohesin loading could be to ensure that cohesin is in place at centromeric regions prior to passage of the replication fork, thereby ensuring its incorporation into functional cohesion. In support of this model, allowing more time for cohesin loading by delaying DNA replication, either by HU treatment or deletion of the S phase cyclins, in iml3Δ and chl4Δ mutants improved pericentromeric cohesion. We cannot at present rule out that DNA damage-induced cohesion [51–53] is responsible for the restoration of pericentromeric cohesion in these cells. However, evidence that replication timing in the absence of DNA damage does play a role in centromere function in iml3Δ and chl4Δ cells, came from our observation that a CEN3-containing late-replicating plasmid is more stable than a similar early-replicating plasmid in iml3Δ and chl4Δ cells. Preliminary observations indicate that early and late replicating, but otherwise identical, CEN4 plasmids behave in a similar manner [JF and AM, unpublished observations]. It is interesting to note that while all centromeres replicate early in S phase, among centromeres, CEN4 and CEN6 replicate relatively late in their normal chromosomal context, whereas CEN5 replicates early [47]. In wild type cells we saw the highest frequency of GFP dot separation with a probe close to CEN5. Whether this is due to the increased distance of the CEN4 and CEN6 probes from the centromere (2.4 kb and 4.5 kb, respectively, compared to 1.4 kb for CEN5), or an effect of their replication timing needs further investigation.

A dedicated pathway for cohesion establishment at the pericentromere

A crucial role for pericentromeric cohesion has emerged during meiosis, where its protection during meiosis ensures the fidelity of chromosome segregation during meiosis I [25]. However, it is becoming increasingly clear that pericentromeric cohesion plays specialized roles in mitosis too [19]. Being proximal to the site of microtubule attachment, it is required to resist spindle forces (Figure 2). Recent findings in fission and budding yeast have also demonstrated its importance in specifying the geometry of kinetochore-microtubule attachment [58,63]. However, generation of cohesion at the centromere represents a considerable challenge, since kinetochore assembly generates a protein-DNA barrier to the replication fork whose passage is intimately linked to cohesion establishment. Furthermore, centromeres are known to replicate early during S phase [47], so that a mechanism must exist to ensure that cohesin is in place prior to passage of the replication fork. We propose that a two-step dedicated pathway of cohesion establishment involving the Ctf19 complex and Csm3 overcome these challenges to ensure the reinforcement of cohesion at the pericentromere. In the first step, the Ctf19 complex, through its Inm3 subunit, directs cohesin loading at the centromere, which subsequently spreads throughout the pericentromeric region. This dedicated pathway of cohesin loading is essential to ensure that cohesin is in place prior to passage of the replication fork early in S phase. Csm3 is crucial in this second step to ensure the integrity of the replisome and its association with cohesion establishment factors as it traverses the protein-DNA barrier that surrounds the centromere. The resultant enhanced cohesion in the vicinity of the centromere safeguards the fidelity of chromosome segregation. The Ctf19 complex and Csm3 are highly conserved. Whether such a mechanism of kinetochore-driven cohesion establishment operates in organisms where pericentromeric heterochromatin is known to attract cohesin [15–18,66] is an important question for the future.

Materials and Methods

Yeast strains and plasmids

All strains used for meiotic experiments were derivatives of SK1 and all strains used for mitotic experiments were derivatives of W303. The iml3Δ::kanMX6 and chl4Δ::kanMX6 deletions were described in [27]. The pCLB2-3HA-cdc20 fusion was described in [44]. The RYG3-3HA and SCC1-6HA tags were described in [67] and [57], respectively. pGAL-SCC1-3HA was described in [4]. The CHL4-6HA and IML3-6HA tags were created using a PCR-directed method [68]. Other deletions were generated using genomic DNA from the yeast deletion collection [69] as template for PCR, followed by transformation. The pMET3-cdc20 construct was constructed using a PCR based method as described by [70]. To create the CEN6-GFP strain, a 1 kb region to the right of CEN6 was cloned into pRS306tetO1112 [42] and integrated into a strain carrying tetR-GFP, resulting in the generation of a GFP label 4.5 kb to the right of CEN6. The Spec42-ttdTomato construct was described in [71]. Other GFP chromosome labels were described previously [21,22,42,72]. Strains are listed in Table S2. Plasmids p12 and p306.10 were described in [56] and [53], respectively.

Growth conditions

Growth conditions for individual experiments are given in the figure legends. All cultures were grown at room temperature unless otherwise stated. Meiosis was performed at 30°C as described in [73]. Drugs and alpha factor were removed by filtration, washing with at least 5 volumes of medium lacking sugar. Methionine was used at 8 mM and readded to 4 mM every hour. To depolymerise microtubules a mixture of benomyl (30 μg/mL) and nocodazole (15 μg/mL) were used and nocodazole (7.5 μg/mL) was readded every hour. Hydroxyurea was used at 10 mg/mL in liquid medium and 100 mM in plates.

ChiP assay and quantification

ChiP was carried out as described by [14]. qPCR was performed in a 2 μl SYBR Green reaction using a BioRad iCycler machine. To calculate ChiP enrichment/input, ΔCT was calculated according to: ΔCT = (CTinput − CTChIP) − logE(1 + dilution factor)) where E represents the specific primer efficiency value. Enrichment/input value was obtained from the following formula: EΔ−ΔCT. The average of 3 independent experiments for which qPCR was performed in triplicate are shown with error bars indicating standard deviation. We designed primers corresponding to cohesin-enriched (A1, A3) or cohesin-poor (A2) sites on chromosome arms as well as a centromeric site (C1) and several pericentromeric sites (P1–6), based on published ChiP data after

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microarray hybridization for the mitotic cohesin, Scc1 [23]. We were not able to analyze site C2 during meiosis since this primer set did not give a product in the SK1 strain background, presumably due to divergence from the published S. cerevisiae sequence. Sequences of primers are available on request. For the ChIP shown in Figure S1, primer sets were as described in [14] and Image J software was used to quantify ethidium bromide-stained gels.

**Chromosome loss assay and plasmid loss assay**

To measure chromosome loss or plasmid loss, strains carrying the SUP11 artificial chromosome, or carrying the early or late replicating plasmids (p306.10 and p12), were grown overnight in minimal media lacking uracil (SD/-ura) before transferring to rich media (YPD) for 2.5–3 h. For the chromosome loss assay, approximately 2000–5000 cells were then plated out on YPD media. Loss of the artificial chromosome causes colonies to appear red. The percentage of sectored colonies that were at least half red, indicating loss in the first division after plating, were scored. To avoid amplification of earlier loss events, entirely red colonies were excluded from the analysis. For the plasmid loss assay, equal culture volumes totaling 1000–2000 cells were plated onto each of YPDA and SD/-URA and the percentage of colonies able to grow on SD/-URA, indicating retention of the plasmid, was calculated.

**Microscopy**

Fixing cells for visualization of GFP-labeled chromosomes was performed as described by [67]. Indirect immunofluorescence methods were as previously described [74]. Microscopy was performed on a Zeiss Axioplan 2 microscope and for measurements of inter-CEN and –SPB distance, images were grabbed using a Hamamatsu ORCA-ER camera and analyzed using Zeiss Axiovision software. In the cohesin assay for mitosis, GFP dots were scored as separated if two dots were clearly visible in the same cell. A total of 200 cells were scored and all cells in the field were scored. To confirm metaphase arrest in each experiment, spindle pole body separation was analyzed either using Spc42-dTomato or after anti-tubulin immunofluorescence or parallel samples. The leakiness of the MET-CDC20 construct allowed spindle elongation in a few cells (usually<5% and never more than 10%) only at later timepoints (120 min) and was comparable in all strains in a given experiment. To score GFP dots in meiosis, cells were co-stained with DAPI to visualize nuclear morphology and 200 binucleate or tetranucleate cells were counted from the 5 and 8 h timepoints, respectively.

**FACS analysis**

For flow cytometry, cells were fixed in 70% ethanol at 4°C over night. The cells were then treated with RNase over night, then digested with pepsin (Sigma). The cells were finally treated with propidium iodide (Sigma) to stain the DNA. Samples were briefly sonicated before analysis. FACS analysis was performed according to the manufacturer’s instructions (BD FACS Calibur). FACS data were analyzed using CellQuest software.

**Western blotting**

Samples for immunoblot analysis were prepared after TCA fixation as described by [75]. A goat Rad53 antibody (γC-19; Santa Cruz Biotechnology, inc.) was used at a dilution of 1:1000.

**Supporting Information**

**Figure S1**  Iml3, Chl4, and Ctf3 are required for proper cohesion in the centromere. (A) Percentages of metaphase (open symbols) and anaphase (closed symbols) spindles are shown for wild type (black squares), chl4A (blue circles), and iml3A (red triangles) strains treated with DMSO for the experiment in Figure 2A after anti-tubulin immunofluorescence. (B) The percentages of separated spindle pole bodies (B) or +2.4CE4-GFP foci (C) are shown at the indicated times after release from G1 for wild type (black squares), chl4A (blue circles), iml3A (red triangles), and ef3A (grey diamonds) for 100 cells at each time point for the experiment shown in Figure 2D and 2E. Found at: doi:10.1371/journal.pgen.1000629.s002 (0.95 MB EPS)

**Figure S2**  Iml3, Chl4, and Ctf3 are required for proper cohesion in the centromere. (A) Percentages of metaphase (open symbols) and anaphase (closed symbols) spindles are shown for wild type (black squares), chl4A (blue circles), and iml3A (red triangles) strains treated with DMSO for the experiment in Figure 2A after anti-tubulin immunofluorescence. (B) The percentages of separated spindle pole bodies (B) or +2.4CE4-GFP foci (C) are shown at the indicated times after release from G1 for wild type (black squares), chl4A (blue circles), iml3A (red triangles), and ef3A (grey diamonds) for 100 cells at each time point for the experiment shown in Figure 2D and 2E. Found at: doi:10.1371/journal.pgen.1000629.s003 (1.30 MB EPS)

**Figure S3**  Similar cohesion defects during mitosis in Ctf19 complex mutants. (A–D) Similar frequency of +2.4CE4-GFP separation in Ctf19 complex single and double mutants. Strains of wild type (AM4643), chl4A (AM4644), mcm16A (AM6148), mcm22A (AM6160), chl4A mcm16A (AM6195), chl4A mcm22A (AM6193), ef3A (AM5786), and men21A (AM5788) mutants carrying MET-CDC20, SPC42-dTomato, and +2.4CE4-GFP were released from a G1 arrest into medium containing methionine to induce a metaphase arrest. The percentages of cells with separated spindle pole bodies (A,C) or +2.4CE4-GFP foci (B,D) were scored at the indicated time points. (E,F) Ctf19 complex mutants exhibit a negligible cohesion defect at the URA3-GFP locus. Strains of wild type (AM1081) iml3A (AM3541), ef3A (AM5814), and men21A (AM5812) mutants carrying MET-CDC20 and URA3-GFP were treated as in (A–D). Spindle morphology after tubulin immunofluorescence (E) and separation of URA3-GFP foci (F) was scored at the indicated time points.

Found at: doi:10.1371/journal.pgen.1000629.s004 (0.48 MB EPS)

**Figure S4**  Reduced levels of Chl4 at the centromere in Ctf19 complex mutants during mitosis. Cycling cells of strains AM1176 (no tag), AM3276 (CHL4-6HA), AM4251 (ctf19A CHL4-6HA), AM4360 (iml3A CHL4-6HA), AM4544 (mcm16A CHL4-6HA), and AM5447 (mcm21A CHL4-6HA) were harvested at 6 h after transfer into sporulation medium. These samples contained cells representative of all meiotic stages. (A) Heterozygous dots; (B) Homozygous dots. Found at: doi:10.1371/journal.pgen.1000629.s005 (0.66 MB EPS)

**Figure S5**  Ctf19 and Mcm21 are required for accurate meiosis II chromosome segregation. The percentage of binucleate cells with the patterns of GFP dot segregation shown was determined after scoring 200 cells in the experiment shown in Figure 3A–3D. (A) Heterozygous dots; (B) Homozygous dots. Found at: doi:10.1371/journal.pgen.1000629.s006 (0.73 MB EPS)

**Figure S6**  Iml3 and Chl4 are localized at the centromere in a Ctf19-dependent manner during meiosis. (A) Locations of primer sets along chromosome III used for PCR analysis of ChIP samples. (B) Chl4 localizes specifically to the core centromere in meiosis. Ethidium bromide-stained gels after PCR analysis of ChIP samples from diploid wild type strains carrying CHL4-6HA and otherwise wild type (AM3446), mcm16A (AM3803), mcm22A (AM3801), ef3A (AM3773), or lacking an HA tag (strain AM1835) harvested 6 h after transfer into sporulation medium. These samples contained cells representative of all meiotic stages. (C) Iml3 localizes specifically to the core centromere during meiosis. Ethidium bromide-stained gels after PCR analysis of ChIP samples from diploid wild type strains carrying IMI3-6HA and otherwise wild type (AM3441), mcm16A (AM3802), mcm22A...
(AM3800), ctf19A (AM3760), or lacking an HA tag (strain AM1035) treated as described in (B). (D) Quantification of ChIP PCR in (B). In each case, the signals were quantified and the ratio of ChIP to input (1:500) PCR was calculated and shown as binding ratio. (E) Quantification of ChIP PCR in (C) performed as described in (D).

Found at: doi:10.1371/journal.pgen.1000629.s006 (3.29 MB EPS)

Figure S7 Csm3 is required for accurate chromosome segregation during meiosis and plays a non-overlapping role with Iml3. Cells of the indicated genotypes and in which both copies ([A];B]; homozygous) or one copy ([C];D]; heterozygous) of chromosome V marked with URA3-GFP were analyzed as described in Figure 1. Strains used in (A) and (B) were AM1603 (wild type), AM1903 (iml3Δ), AM3252 (csm3Δ), and AM3774 (iml3Δ csm3Δ). Strains used in (C) and (D) were AM107 (wild type), AM1904 (iml3Δ), AM3379 (csm3Δ), and AM3788 (iml3Δ csm3Δ). Found at: doi:10.1371/journal.pgen.1000629.s007 (0.76 MB EPS)

Figure S8 Relationship between cohesion loading and DNA replication. FACS analysis of the experiment shown in Figure 3A–3E. Cells were analysed by flow cytometry to determine when the cells had replicated their DNA. For each sample, 15,000 cells were analysed. Found at: doi:10.1371/journal.pgen.1000629.s008 (0.67 MB EPS)

Figure S9 DNA replication delay induced by HU treatment. FACS analysis of the experiment shown in Figure 6 (B–D). Cells were analysed by flow cytometry to determine when the cells had replicated their DNA. For each sample, 15,000 cells were analysed. (A) The cells were released from G1 arrest directly into media containing methionine. The red stars indicate the time after G1 release at which most cells have fully replicated genomes (2n). The numbers indicate the time after G1 release, (B) The cells were delayed in S-phase for 90 min by HU treatment, then released into media containing methionine. The blue star indicates the timepoint at which most cells showed fully replicated (2N) genomes. The numbers indicate the time following the release from the 90 min treatment of 10 mg/ml HU. Found at: doi:10.1371/journal.pgen.1000629.s009 (2.98 MB EPS)

Figure S10 DNA damage checkpoint arrest and recovery after HU treatment. (A) Western blot showing Rad53 hyperphosphorylation upon HU treatment and restoration of faster migrating forms after release. Experiment was performed and strains used were as described in Figure 6B–6E, except samples were taken for Western blotting at the indicated timepoints and probed with anti-Rad53 antibody. (B) Sensitivity of wild type (AM1176), iml3Δ (AM3315), csm3A (AM3314), ctf19A (AM3958), csm3Δ (AM1914), and rad53A iml3Δ (AM6154) to HU. Serial dilutions of the induced strains were spotted onto YPDA and plates containing 100 mM HU in parallel and incubated for 2 days at 30°C. Found at: doi:10.1371/journal.pgen.1000629.s010 (7.24 MB EPS)

Table S1 Chromosome loss rates. Found at: doi:10.1371/journal.pgen.1000629.s011 (2.31 MB EPS)

Table S2 Yeast strains. Found at: doi:10.1371/journal.pgen.1000629.s012 (0.06 MB DOC)

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Author Contributions
Conceived and designed the experiments: JF ALM. Performed the experiments: JF ALM. Analyzed the data: JF ALM. Contributed reagents/materials/analysis tools: JF ALM. Wrote the paper: JF ALM.


