Saccharomyces cerevisiae CTF18 and CTF4 Are Required for Sister Chromatid Cohesion

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CTF4 **and** *CTF18* **are required for high-fidelity chromosome segregation. Both exhibit genetic and physical ties to replication fork constituents. We find that absence of either** *CTF4* **or** *CTF18* **causes sister chromatid cohesion failure and leads to a preanaphase accumulation of cells that depends on the spindle assembly checkpoint. The physical and genetic interactions between** *CTF4, CTF18***, and core components of replication fork complexes observed in this study and others suggest that both gene products act in association with the replication fork to facilitate sister chromatid cohesion. We find that Ctf18p, an** *RFC1***-like protein, directly interacts with Rfc2p, Rfc3p, Rfc4p, and Rfc5p. However, Ctf18p is not a component of biochemically purified proliferating cell nuclear antigen loading RF-C, suggesting the presence of a discrete complex containing Ctf18p, Rfc2p, Rfc3p, Rfc4p, and Rfc5p. Recent identification and characterization of the budding yeast polymerase** k**, encoded by** *TRF4***, strongly supports a hypothesis that the DNA replication machinery is required for proper sister chromatid cohesion. Analogous to the polymerase switching role of the bacterial and** human RF-C complexes, we propose that budding yeast RF-C^{CTF18} may be involved in a polymerase switch **event that facilities sister chromatid cohesion. The requirement for** *CTF4* **and** *CTF18* **in robust cohesion identifies novel roles for replication accessory proteins in this process.**

The establishment of sister chromatid cohesion during S phase is a critical step in the series of events leading to highfidelity cell division. By holding sisters together, cohesion proteins enable kinetochores to face opposite poles of the mitotic spindle, facilitating capture by microtubules from opposite poles (99). The sister chromatid association is sufficient to resist the separating force of the mitotic spindle until each kinetochore has been captured, at which time sister chromatid associations are released at the initiation of anaphase (reviewed in references 50, 72, 77, and 88). Because cohesion tightly binds sisters together from their synthesis to their separation, it must be properly established and maintained in a flexible environment supporting chromatin alterations that permit transcription, replication, repair, and condensation of the genome.

Cohesion between sister chromatids is carried out by at least four classes of proteins. The core particle, cohesin, is composed of at least four subunits encoded in budding yeast by the *SMC1, SMC3, MCD1* (*SCC1*), and *SCC3* (*IRR1*) genes (33, 68). Fully assembled cohesin binds chromatin in vitro and in vivo (9, 68, 97, 101). Orthologs of cohesins have been identified in *Xenopus laevis, Drosophila melanogaster, Schizosaccharomyces pombe, Arabidopsis thaliana, Mus musculus*, and *Homo sapiens* (6, 18, 58, 59, 82, 94, 100, 109, 110). Interestingly, although Mcd1p is required for both cohesion and chromosome condensation in budding yeast, these processes are carried out by distinct protein complexes in the *Xenopus* experimental system (33, 40, 41, 58). In addition, Pds5p, which is also re-

quired for the maintenance of sister chromatid cohesion, genetically and physically interacts with the cohesin complex (36, 78, 94). Thus, interactions between the cohesin complex and Pds5p are required to mediate sister chromatid cohesion. A highly conserved mechanism governs sister chromatid separation at anaphase initiation, mediated by the action of a *CDC20* associated form of the anaphase-promoting complex (APC- $CDC20$), which provides a ubiquitin-conjugating activity directing the degradation of the anaphase inhibitor protein Pds1p (reviewed in reference 70). Upon release from a Pds1p-Esp1p complex, active Esp1p promotes the proteolysis of Mcd1p (94, 104, 107). This event is associated with loss of cohesion between sister chromatids and with poleward movement of the chromosomes (reviewed in references 73 and 116).

Scc2p and Scc4p, members of the second class of proteins, direct the binding of cohesin proteins to chromatin (16, 101). *SCC2* and *SCC4* associate with each other in coimmunoprecipitation experiments but are not core components of the cohesin particle (16). Scc2p may mediate cohesin complex interaction with chromatin via associations with Mcd1p and Scc3p. While the localizations of both Scc2 and Scc4 proteins on chromatin spreads are similar to one another, the two proteins seem to occupy different chromosomal loci from Mcd1p (16, 101). Furthermore, both Scc2p and Scc4p associate with chromatin in a nuclease- and salt-resistant manner, suggesting that they are tightly bound in a higher-order chromatin structure. Scc2p and Scc4p are required for establishment of cohesion early in the cell cycle but are not required for maintenance of cohesion in metaphase arrested cells (16). This function appears to be conserved, since a fission yeast homologue of Scc2p (Mis4p) is also required in S phase (26). *SCC2* homologues have also been identified as the *Coprinus RAD9* (83) and *Drosophila* Nipped-B (81) genes.

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A third class of molecules, defined by *CTF7* (*ECO1*) function, is required to render the cohesin complex competent to mediate siser chromatid cohesion during S phase. Budding yeast are inviable in the absence of *CTF7*, and conditional alleles lead to precocious sister separation (87, 101). Although Ctf7p associates with chromatin, it does not stably associate with the core cohesin particle, nor is it requred for cohesin association with chromatin (87, 101). Execution point studies indicate a requirement for *CTF7* in S phase (87, 101). Interestingly, the chromosome instability and temperature-sensitive lethality of *ctf7* alleles are suppressed by high-copy-number expression of *POL30*, encoding budding yeast proliferating-cell nuclear antigen (PCNA) (87). One hypothesis is that Ctf7p is required at the replication fork to activate interactions between cohesins on sister chromatids for a functional "glue" to be formed. Budding yeast Ctf7p is homologous to a C-terminal domain of the fission yeast Eso1 protein, whose N-terminal segment is homologous to the budding yeast DNA repair polymerase RAD30 (96). The $Esot⁺$ gene functions similarly during S phase and is required for sister chromatid cohesion, pointing to conservation of this activity as well as its association with replication of DNA.

Recent studies have identified proteins more directly involved in DNA replication as members of the fourth class of cohesion proteins. This category includes PCNA by genetic interaction with *CTF7* as described above (87) and a new DNA polymerase family, designated polymerase k, exemplified by budding yeast gene TRF4 (108). PCNA forms a homotrimeric ring structure (clamp) which encircles DNA and supports processive DNA replication by associated DNA polymerases δ and ε (reviewed in references 39 and 47). A "clamp-loader," replication factor C (RF-C), is required to facilitate association of PCNA with DNA (reviewed in reference 71). RF-C is composed of five essential subunits that have a common core region of homology that may facilitate interactions with PCNA (1, 42, 65), as well as interact with DNA (103). RF-C may also mediate a switch from polymerase α -directed replication initiation to processive replication by polymerases δ and ε through competitive interactions with PCNA and the budding yeast single-stranded binding protein replication protein A (RPA) (reviewed in reference 19).

TRF4 and its paralog *TRF5* are both members of the β -polymerase superfamily as defined by protein alignment (3). While neither the *TRF4* nor *TRF5* gene is essential, in combination they exhibit synthetic lethality. Recent work has provided direct evidence that Trf4p encodes a novel polymerase and that a *trf4 trf5* double mutant exhibits highly inefficient S-phase DNA replication (108). Like other budding yeast genes that function in cohesion, *TRF4* is required for both chromosome condensation (14) and sister chromatid cohesion (108). The chromosomal defects present in a *trf4* mutant cell influence the maintenance of sister chromatid cohesion under mitotic arrest conditions, probably through an uncharacterized mechanism that operates during DNA replication (108).

In this work, we present an analysis of two genes, *CTF4* and *CTF18*, that exhibit genetic and physical interactions with components of the replication fork and that are required for sister chromatid cohesion. Analysis of these genes provides independent evidence that cohesion-related functions are indeed carried out by proteins associated with the DNA synthesis machinery. Previous genetic analyses have indicated that neither *CTF4* nor *CTF18* is essential; however, absence of either gene increases chromosome instability and mitotic recombination rates and induces a strong preanaphase delay (51, 53, 69). *CTF4* was first identified in concurrent studies as *CTF4* (*CHL15*) (52) and *POB1* (69) and encodes a 104-kDa polypeptide with motifs suggestive of three zinc finger structures, as well as a helix-loop-helix region (35, 51). Ctf4p exhibits highaffinity binding to DNA polymerase α in vitro (69), and *ctf4* mutants show genetic interactions with conditional alleles of DNA polymerase α (encoded by *POL1* [*CDC17*]) and other genes intimately associated with DNA synthesis (24, 112). A $ctf4\Delta$ mutation also causes synthetic lethality with a null allele of *CTF18*, the other gene investigated in this study (24). *CTF18* (*CHL12*) was independently identified in two screens for chromosome loss mutants (52, 91). *CTF18* encodes a predicted 84-kDa polypeptide with homology to all five subunits of the RF-C complex from yeast and other organisms, with the most significant similarity to the large subunit Rfc1p (17, 53).

Here we find that both *CTF4* and *CTF18* are required for sister chromatid cohesion. This, rather than the presence of damaged DNA, is likely to be the major mechanism underlying chromosome loss in both *ctf4* and *ctf18* null mutants. The preanaphase delay exhibited by each of these mutants is due to a spindle assembly checkpoint arrest. Because we and others have observed that *CTF4* and *CTF18* exhibit genetic interactions with genes that function in DNA replication, we suggest that *CTF4* and *CTF18* act in association with the replication fork complex(es) to facilitate the establishment of robust sister chromatid cohesion. We propose a model in which Ctf18p, an Rfc1p paralog, may act within a complex similar to the previously characterized RF-C, consistent with physical and genetic evidence presented in this report.

The replication fork plays numerous roles in the chromosome cycle, including duplication of genomic DNA, detection and repair of lesions (85, 93), and regulation of transcriptional states (20, 89). The properties of *CTF4* and *CTF18* highlight a new function of the replication fork machinery that is essential for high-fidelity segregation of the genome. The characterization of these proteins, which interact with residents of the replication fork and whose loss of function compromises cohesion, presents new opportunities for investigation of this novel role of the replication machinery.

MATERIALS AND METHODS

Strains and media. Yeast transformation (29, 49) and genetic manipulations (34) were performed by published methods. rYPD is rich medium adjusted to pH 4 and supplemented with additional adenine hemisulfate (30 ng/ml), uracil (20 ng/ml), and L-tryptophan (30 ng/ml). Strain sets (Table 1) within each single experiment are composed of laboratory stocks related by transformation or genetic crosses maintaining background isogenicity. Complete oligonucleotide sequences will be provided on request.

All strains were grown to an optical density at 600 nm of 0.4 at 30°C unless otherwise noted. G_1 arrest occurred after 2.5 h in 3 μ M alpha mating pheromone; S-phase arrest was achieved by incubation for 2.5 h in 0.2 M hydroxyurea (HU); metaphase arrest was achieved by incubation for 4 h in 15 μ g of nocadozole, per ml.

The *CTF18* open reading frame (ORF) was disrupted by transforming with a *Spe*I-*Not*I fragment of pJH28::TRP. PCR primers flanking the inserted fragment were used to detect disruption of the *CTF18* (OLFS278 plus OLFS279). The *CTF4* ORF was disrupted using OLFS369 plus OLFS370 (10); verification PCR used OLFS371 plus OLFS372. A $mad2\Delta$ allele was transferred from YPH1238 using primers OLFS365 plus OLFS366; detection was performed using PCR

| Strain | Genotype | Source or reference |
|----------------|---|---------------------|
| YJH17.2 | MATa ade2-1 can1-100 trp1-1 ura3-1 ctf184::TRP1 his 3-11,15::GFP-Lac1-HIS3 $leu2-3,112::lacO-LEU2$ | This study |
| YJH18.3 | MATa ura3-52 lys2-801 ade2-101 his3 Δ 200 trp1 Δ 63 leu2 Δ 1 ctf18 Δ ::LEU2 mad2 Δ ::HIS3 | This study |
| YCB624 | MATa ura3-52 lys2-801 ade2-101 trp1 Δ 63 his2 Δ 200 leu2 Δ 1 rad9 Δ ::TRP1 | 10 |
| YJH35.1 | MATa ura3-52 lys2-801 ade2-101 trp1 Δ 63 his2 Δ 200 leu2 Δ 1 rad9 Δ ::TRP1 ctf18 Δ ::LEU2 | This study |
| YJH37 | MATa ade2-1 can1-100 trp1-1 ura3-1 ctf4 Δ ::TRP1 his3-11,15::GFP-LacI-HIS3 leu2-3,112::lacO-LEU2 | This study |
| YJH38 | $MATa$ ura3-52 lys2-801 ade2-101 trp1 ars1 $\Delta HIS3$ leu2 Δ 1 CTF4::9Myc-klTRP1 | This study |
| YJH40.4 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1∆ HIS3 leu2∆1 CTF18::9Myc-klTRP1 | This study |
| YJH41.1 | MATa ura3-52 lys2-801 ade2-101 leu2 Δ 1 his3 Δ 200 mad2 Δ ::HIS3 ctf4-65 | This study |
| YJH48 | MATa ura3-52 lys2-801 ade2-101 his3 Δ 200 trp Δ 63 leu2 Δ 1 ctf18 Δ ::LEU2 + pJH76.1 (GAL-CTF18- 9mvc | This study |
| YJH62.6 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1 Δ HIS3 leu2 Δ 1 RFC1::9Myc-klTRP1 | This study |
| YJH63.2 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1∆ HIS3 leu2∆1 CTF18::9Myc-klTRP1 RFC2::6HA-KanMX | This study |
| YJH64.1 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1∆ HIS3 leu2∆1 CTF18::9Myc-klTRP1 RFC4::6HA-KanMX | This study |
| YJH65.3 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1∆ HIS3 leu2∆1 CTF18::9Myc-klTRP1 RFC5::6HA-KanMX | This study |
| YJH67.1 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1\, HIS3 leu2\\,1 CTF18::9Myc-klTRP1 RFC3::6HA-KanMX | This study |
| YJH69.1 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1∆ HIS3 leu2∆1 RFC2::6HA-KanMX | This study |
| YJH70.1 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1∆ HIS3 leu2∆1 RFC3::6HA-KanMX | This study |
| YJH71.1 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1 Δ HIS3 leu2 Δ 1 RFC4::6HA-KanMX | This study |
| YJH72.1 | MATa ura3-52 lys2-801 ade2-101 trp1 ars1∆ HIS3 leu2∆1 RFC5::6HA-KanMX | This study |
| AFS173 | MATa ade2-1 can1-100 trp1-1 ura3-1 his3-11,15::GFP-LacI-HIS3 leu2-3,112::lacO-LEU2 | 92 |
| AFS387 | MATa ade2-1 can1-100 trp1-1 ura3-1 mad2-1 his3-11,15::GFP-LacI-HIS3 leu2-3,112::lacO-LEU2 | 92 |
| s65 | MATa ura3-52 lys2-801 ade2-101 leu2 Δ 1 his3 Δ 200 ctf4-65 | 91 |
| YE77 | $MATa/a$ ura3-52/ura3-52 lys2-801/lys2-801 ade2-101/ade2-101 his3 Δ 200/his3 Δ 200 trp1 Δ 63/trp1 Δ 63 | 53 |
| | $leu2\Delta1/leu2\Delta1$ ctf18 $\Delta::LEU2$ /ctf18 $\Delta::LEU2$ | |
| YE105 | MATa ura3-52 lys2-801 ade2-101 his3 Δ 200 trp Δ 63 leu2 Δ 1 ctf18 Δ ::LEU2 | 53 |
| YRD501 | MATa leu2-3,112 ura3-52 trp1-289 cdc28-1 | Li laboratory |
| YRD510 | MATa leu2-3,112 ura3-52 trp1-289 cdc4-1 | Li laboratory |
| YRD543 | MATa leu2-3,112 ura3-52 trp1-289 his3 Δ 200 cdc7-4 | Li laboratory |
| YRD664 | MATa leu2-3,112 ura3-52 trp1-289 cdc34-2 | Li laboratory |
| YJL179 | MATa leu2-3,112 ura3-52 trp1-289 cdc46-1 | Li laboratory |
| YJL338 | MATa leu2-3,112 ura3-52 trp1-289 cdc2-1 | Li laboratory |
| YJL340 | MATa leu2-3,112 ura3-52 trp1-289 ade2 cdc6-1 | Li laboratory |
| YJL353 | MATa leu2-3,112 ura3-52 trp1-289 cdc17-1 | Li laboratory |
| YPH216 | MATa ade2 ade3 his7 leu2 can1 sap1 cdc9-1 | Hartwell laboratory |
| YPH217 | MAT a ade2 ade3 his7 leu2 can1 sap1 cdc13-1 | Hartwell laboratory |
| YPH221 | MATa his7 leu2 ase2 ase3 sap3 gal1 ura1 cdc23-1 | Hartwell laboratory |
| YPH223 | MATa his7 leu2 ase2 ase3 sap3 gal1 cdc14-1 | Hartwell laboratory |
| YPH277 | $MATa$ ura3-52 lys2-801 ade2-101 trp1 $\Delta 63 + CFVII(RAD2.d.YPH277)$ URA3 SUP11 | 91 |
| YPH499 | MATa ura3-52 lys2-801 ade2-101 his3 Δ 200 trp1 Δ 63 leu2 Δ 1 | 85a |
| YPH501 | $MATa/a$ ura3-52/ura3-52 lys2-801/lys2-801 ade2-101/ade2-101 his3 Δ 200/his3 Δ 200 trp1 Δ 63/trp1 Δ 63 $leu2\Delta1/leu2\Delta1$ | 85a |
| YPH1238 | MATa ura3-52 lys2-801 ade2-101 his3 Δ 200 trp1 Δ 63 leu2 Δ 1 mad2 Δ ::HIS3 | Hieter laboratory |
| YDS489 | $MATa$ ura $3-1$ ade $2-1$ his $3-11$ trp $1-1$ rap $1-5$ | 61 |

TABLE 1. Yeast strains used in this study

primers OLFS367 plus OLFS368. Epitope tagging of *CTF18, CTF4, RFC1, RFC2, RFC3, RFC4*, and *RFC5* was performed as described previously (49) using primer pairs OLFS359 plus OLFS360, OLFS373 plus OLFS374, OLFS491 plus OLFS492, OLFS510 plus OLFS511, OLFS506 plus OLFS507, OLFS508 plus OLFS509, and OLFS512 plus OLFS513, respectively; detection of integration used primer pairs OLFS344 plus OLFS345, OLFS371 plus OLFS372, OLFS493 plus OLFS494, OLFS514 plus OLFS515, OLFS437 plus OLFS438, OLFS439 plus OLFS440, and OLFS516 plus OLFS517.

Vector construction. A *Spe*I-*Not*I fragment from BFS66, containing the *CTF18* ORF and flank, was cloned into pBluescript II to give pJH28. pJH28::TRP was created by removal of the internal two-thirds of *CTF18* ORF by digestion with *Nsi*I and *Mlu*I, and its replacement by ligation of a PCR product from primers OLFS264 plus OLFS265, which amplified the *TRP1* gene from pRS404 (10). pJH72.1 was constructed by cloning the entire *CTF18* ORF into pCR2.1 (Invitrogen TOPO-TA) using primers OLFS274 plus OLFS362 and LTI Pfx Taq; the *CTF18* ORF was then released using *Nco*I-*Eco*RI digestion and cloned into the same sites in pAS2-1 (Clontech), which created an in-frame fusion with the GAL4 DNA binding domain (pJH74.4). pJH73.2 was constructed from a PCR product (using LTI Pfx Taq, primers OLFS274 plus OLFS345, and YJH40.4 genomic DNA) digested with *Pvu*I to liberate *CTF18*-9myc from the coamplified TRP1 marker. *Taq* polymerase modified the ends, and the product was TA cloned into pCR2.1. pJH76.1 was constructed by cloning the *CTF18*-9myc allele

on an *Xho*I-*Hin*dIII fragment from pJH73.2 into the same sites in pRS316GU (74).

pJH78 was constructed using primers OLFS437 plus OLFS438 and genomic DNA from YJH40.4; the product was cloned into pCR2.1. pJH79 was constructed in similar fashion using primers OLFS439 plus OLFS440.

p414GEU1/12 was created by cloning a 2,236-bp *Sal*I-*Eco*RI PCR fragment containing *CTF18* into *Sal*I- and *Eco*RI-cut p414GEU1 (54), placing the *CTF18* ORF in frame with the vector ATG and double E1 epitope tag. The resulting plasmid conferred wild-type chromosome stability and growth at low temperature to a *ctf18* Δ strain in galactose-dependent fashion.

Flow cytometry. A 1-ml volume of cells grown in rYPD was harvested and fixed in 500 μ l of 0.2 M Tris (pH 7.5)–70% ethanol. After being washed in 1 ml of 0.2 M Tris (pH 7.5), samples were resuspended in 1 ml of 0.2 M Tris (pH 7.5) for $>$ 30 min, incubated in 100 μ l of 0.2 M Tris (pH 7.5)–3 mg of RNase A per ml for 2.5 h at 37°C, washed with 1 ml of 0.2 M Tris (pH 7.5), and resuspended in 100 ml of 0.05% trypsin (Gibco 25300-054) at 37°C for 5 min. After a 1-ml wash in 0.2 M Tris (pH 7.5), samples were resuspended in 1 ml of 0.2 M Tris (pH 7.5)–9 μ g of propidium iodide per ml.

Sister chromatid cohesion assay. Log-phase cultures were resuspended in SC-HIS medium–40 mM 3-aminotriazole for 40 min to induce green fluorescent protein (GFP)-LacI expression, incubated in YPD-15 µg of nocodazole per ml for 3 h, fixed in 4% paraformaldehyde for 30 min, washed in 1 ml of SK (1 M sorbitol, 50 mM KPO₄, pH 7.5), and resuspended in 50 μ l of SK.

Chromosome spreads. Chromosome spreads were prepared on slides essentially as described in references 48 and 68. Samples on cured slides were washed in phosphate-buffered saline (PBS) for 20 min, preincubated in PBS–1% bovine serum albumin (BSA) for 1 h, and incubated in polyclonal rabbit anti-myc antibody (sc-789; Santa Cruz Biotechnology) in PBS–1% BSA (at a 1:250 dilution for *CTF18*-9myc and a 1:500 dilution for *CTF4*-9myc) for 2 h. Samples were then washed three times with PBS, incubated for 2 h with fluorescein isothiocyanate (FITC)-conjugated anti-rabbit antibody (Sigma) for 2 h (1:1,000 in PBS–1% BSA), washed three times with PBS, and mounted in Fluorsave (Calbiochem)–2 μ g of 4',6-diamidino-2-phenylindole (DAPI) per ml.

Protein analysis. Samples prepared as described previously (49) were subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis (10% polyacrylamide) (SDS-PAGE) and transferred to a nitrocellulose membrane. Polyclonal rabbit anti-myc antibody (sc-789) and a polyclonal goat anti-rabbit antibody–horseradish peroxidase (111-035-144; Jackson ImmunoResearch) were used for detection. Protein loading was determined by standardization to β -tubulin (polyclonal antibody R43; a gift of D. Koshland). Anti-Orc3p monoclonal antibody (SB3) was a gift of B. Stillman (57). Film scans were obtained with a Hewlett-Packard 4c/T scanner and analyzed using NIH Image 1.61.

Antisera against *CTF18***.** A C-terminal 18-kDa peptide of *CTF18* expressed from pRSETA (InVitrogen) in bacterial strain BL21(DE3)/pLysE was affinity purified on a $Ni²⁺$ -containing column (Qiagen), solubilized in 8 M urea, and used to immunize two New Zealand rabbits (Hazelton, Inc.). Antibodies from one (rabbit 10C) were purified against ctf18 Δ protein extract bound to cyanogen bromide-activated Sepharose beads. The Western blot signal from purified bacterial peptide in serial dilution indicated that the 10C antibody was capable of detecting at least 2 ng of Ctf18p in the experiment in Fig. 6.

Yeast two-hybrid. pJH74 (BD-CTF18) does not autoactivate reporters in strain AH109 (Clontech). AH109 containing pJH74 was transformed with a *Saccharomyces cerevisiae* cDNA library (gift of S. Elledge) cloned into pACT2-1. A total of $180,000$ transformants yielded 162 His⁺-positive strains, of which 65 were also Ade⁺. PCR analysis and plasmid isolation indicated the presence of four distinct genes. Two of these, $RFC3$ and $RFC4$, also conferred β -galactosidase activity by plate assay (Clontech).

In vitro immunoprecipitations. Portions (1 mg) of circular pJH73.2 (*CTF18*- 9myc), pJH78 (*RFC3*), or pJH79 (*RFC4*) were used as templates in a 50-µl linked transcription-translation system (Promega). A 24-µl volume of each product was used per immunoprecipitation in a final volume adjusted to 150 μ l with 1× PBS–1% Triton X-100, and the mixtures were incubated for 2 h at 4° C. A 1-µl volume of anti-myc (sc-789) was added for a further 2-h incubation, followed by immune complex precipitation for 1 h using 10 μ l of protein A-agarose beads (sc-2001; Santa Cruz). Following three washes (each with 1 ml of PBS–1% Triton $X-100$), the beads were resuspended in 20 μ l of HU buffer (49) and boiled for 5 min. The entire supernatant was analyzed by SDS-PAGE (10% polyacrylamide).

Whole-cell extract immunoprecipitations. A total of 5×10^8 logarithmically growing cells were washed twice with water and resuspended in 1.5 ml of icechilled buffer B60 (50 mM HEPES-NaOH [pH 7.3], 0.1% Triton X-100, 20 mM β -glycerophosphate, 10% glycerol, 0.5 mM phenylmethylsulfonyl fluoride, 2 μ g of leupeptin per ml, and 2 μ g of pepstatin per ml 2 \times Complete [Roche Biochemicals], 60 mM potassium acetate), and 1.5 g of ice-chilled glass beads (400 to $600 \mu m$ in diametet was added. The tubes were vortexed eight times for 30 s with 30-s intervals on ice. After 10 min on ice, the lysate was decanted into ice-chilled 15-ml Corex tubes and centrifuged for 20 min at $18,000 \times g$ at 4° C.

A 500-µl volume of clarified lysate was incubated with 25μ l of prewashed protein A-agarose beads at 4° C for 1 h. The beads were pelleted, and 450μ l of the lysate was transferred to a tube containing 7.5μ l of anti-myc antibody (Santa Cruz) and incubated at 4° C for 2 h. Then 25 μ l of prewashed protein A-agaroseconjugated beads was added, and the mixture was incubated for 1 h at 4°C. The beads were then washed successively seven times: four times with B60 adjusted to 100 mM potassium acetate and once each with B60 adjusted to 210, 240, or 270 mM potassium acetate. The beads were boiled in HU buffer for 5 min and briefly pelleted at 13,000 rpm in an Eppendorf centrifuge before the supernatant was loaded for electrophoresis.

Identification of candidate *CTF18* **orthologues.** The protein sequence of budding yeast *CTF18* was used as query for a TBLASTN search of the GenBank nonredundant protein database, yielding a significant match to human genomic clone HS321D2. GeneSCAN analysis of the genomic clone predicted three genes, including candidate Hs*CTF18*, as depicted in Fig. 8. Verification between amino acids 356 and 1220 of the predicted protein is provided by sequence from the cDNA clone pJH3 (data not shown). The GenBank accession numbers for Rfc1p homologs and Ctf18p homologs in other species identified through PSI-

BLAST are as follows: *C. elegans* Ctf18p, T23478; *C. elegans* Rfc1p, T20230; *D. melanogaster* Ctf18p, AAF51072.1; *D. melanogaster* Rfc1p, AAB58311.1; *S. pombe* Ctf18p, CAB62096.1; *S. pombe* Rfc1p, CAA18875; *H. sapiens* Rfc1p, NP_002904.1; *S. cerevisiae* Ctf18p, NP_013795.1; *S. cerevisiae* Rfc1p, NP_014860.1.

RESULTS

The G2/M delay of *ctf4* **and** *ctf18* **mutants is dependent on the spindle assembly checkpoint.** Previous work has shown that cells lacking *CTF18* accumulate a large-budded morphology (53). In agreement, we found that analysis of log-phase cultures of *ctf18* Δ cells by flow cytometry revealed a substantial accumulation of cells with $G₂$ DNA content. Time course analysis in synchronous cultures was performed to address the nature of the delay. After synchronization in $G₁$, cells were released into rich medium and samples were taken every 10 min for flow cytometry (Fig. 1A). In comparison with the wild-type control, the progression of *ctf18* cells was not detectably different for the first 80 min. New G_1 cells appeared in wild-type and *ctf18* strains at 90 and 100 min, respectively, indicating that a second cycle occurred with a \sim 10-min delay in the mutant. Moreover, a large proportion of the *ctf18* population remained in the G_2 peak until the end of the experiment (150 min).

The delay indicated by flow cytometry may reflect late execution of any step after the completion of bulk DNA synthesis through daughter separation. The mitotic cycle was further characterized by analysis of asynchronous cultures using bud size and spindle morphology as indicators of cell cycle position. Comparison between *ctf18*∆ and wild-type controls revealed a dramatic increase in the number of cells with an intermediatelength spindle (Fig. 1B, class III), representing 20% of mutant cells versus 1% of wild-type cells. The abundant class III morphology indicates the presence of many cells with altered progression through metaphase or anaphase. We conclude that an early mitotic delay underlies the G_2 accumulation observed by flow cytometry.

Early mitotic delay is often due to activation of either the DNA damage checkpoint or the spindle assembly checkpoint (reviewed in references 86 and 111). To investigate further, we constructed *rad9*D and *mad2*D mutants in a *ctf18*D background to remove the DNA damage and spindle assembly checkpoints, respectively. DNA content analysis of log-phase cultures grown at 30°C was used to monitor the cell cycle distributions in these double mutants (Fig. 1C). The results indicated that the preanaphase delay in the $ctf18\Delta$ mutant was dependent on the spindle assembly checkpoint and not on the DNA damage checkpoint. Similar analysis of $\text{cf18}\Delta$ *mec1* Δ double mutants, in an $\frac{sml1-l}{m}$ mutant background which renders $\frac{mecl}{\Delta}$ mutants viable (119), also indicated that the DNA damage checkpoint was not responsible for the $\text{cf18}\Delta \text{ G}_2$ peak accumulation (data not shown).

Morphological analysis also indicated a *MAD2*-dependent mitotic delay. A nuclear morphology in which the DAPIstained chromosomal mass crossed the neck between mother and bud was present in higher levels in $ctf18\Delta$ cells than in wild-type cells (Fig. 1D). The frequency of this class of cells was reduced to wild-type levels in the $\text{cf18}\Delta \text{ } \text{ } \text{mad2}\Delta \text{ }$ strain. Thus, in cultures grown at 30°C, the absence of *ctf18* protein leads to activation of the spindle assembly checkpoint.

This conclusion suggested that a similar phenomenon could

FIG. 1. Absence of *CTF4* or *CTF18* leads to a MAD2-dependent preanaphase delay. (A) Log-phase cultures grown in rYPD were arrested in α -factor, released into rYPD, and processed for flow cytometry. The strains were *ctf18*D (YE105) and *CTF18* (YPH499). (B) Log-phase cultures of *ctf18* Δ /*ctf18* Δ (YE77) or wild-type (YPH501) cells were fixed in formaldehyde and stained to visualize the DNA (DAPI) and microtubules (b-tubulin indirect immunofluorescence). Class III was defined as cells with a single DNA mass that crossed the neck and a spindle that did not extend beyond the center of either mother or bud. (C) Early-log-phase cells were processed for flow cytometry. Strains were *CTF18* (YPH499), *mad2*D (YPH1238), *rad9*D (YCB624), *ctf18*D (YE105), *ctf18*D *mad*2D (YJH18.3), *ctf18*D *rad9*D $(YJH35.1)$, *ctf4* (s65), and *ctf4 mad2* Δ (YJH41.1). Similar results were obtained in five independent experiments; representative results are shown. (D) Formaldehyde-fixed log phase cells were stained with DAPI. Large-budded cells with a single nucleus located within the bud neck were scored (6% in wild type, $n = 100$). Strains were *CTF18* (YPH499), *ctf18*D (YE105), and *ctf18*D *mad2*D (YJH18.3).

explain a non-*RAD9*-dependent delay observed in *ctf4* mutant cells (69). Cells lacking *CTF18* cannot survive in the absence of *CTF4*, consistent with the idea that these gene products separately contribute to the same essential function. To determine whether *ctf4* mutant delay was dependent on the spindle assembly checkpoint, *ctf4 mad2* Δ double mutants were created and analyzed by flow cytometry. Again, the accumulation of $ctf4$ cells of G_2 DNA content was dependent on $MAD2$ (Fig. 1C).

Sister chromatid cohesion failure in *ctf18* **and** *ctf4* **mutants.** Most inducers of the spindle assembly checkpoint cause preanaphase delay or arrest associated with a short spindle morphology (reviewed in reference 86). However, an appreciable proportion of *ctf18*D cells contain partially elongated spindles and stretched DNA masses. Interestingly, a *MAD2*-dependent $G₂/M$ delay with an intermediate spindle morphology had recently been described for *ctf7* mutants, which also exhibit a defect in sister chromatid cohesion. One hypothesis (87) is that precocious sister separation results in a loss of tension at the kinetochore-microtubule interface (triggering the spindle assembly checkpoint), allowing an increase in pole separation at metaphase arrest due to altered balance between opposing forces within the spindle. Based on the phenotypic similarity to *ctf7*, we were encouraged to assay *ctf18* and *ctf4* mutants for a sister chromatid cohesion defect.

In wild-type cells, the arms of duplicated sister chromatids remain tightly associated (32) until the dissolution of cohesins at the metaphase-anaphase transition. The *CTF4* or *CTF18* gene was deleted in a strain background containing a tandem array of lac operators integrated at *LEU2* on chromosome III and expressing a GFP-LacI fusion (92). This configuration allows the assessment of sister chromatid cohesion on chromosome arms throughout the cell cycle: transition from one GFP signal spot to two indicates separation of sister chromatids. Detection of sister chromatid cohesion proficiency can be enhanced by inducing metaphase arrest using a microtubuledisrupting drug such as nocodazole (33, 68). Under these conditions, wild-type cells arrest with unseparated chromatids whereas mutants defective in cohesion exhibit separation.

Log-phase $ctf4\Delta$ and $ctf18\Delta$ cultures were arrested in nocodazole for 3 h, fixed, and scored for the number of GFP spots per cell. Parallel analysis of wild-type and *mad2-1* strains served as positive and negative controls for cohesion. Compared to wild-type cells, *ctf4* and *ctf18* mutants exhibited a high level of precociously separated sister chromatids (Fig. 2A). The number of cells with two GFP spots was comparable to that seen in *mad2-1* cells, which cannot respond to the loss of spindle integrity and inappropriately proceed to anaphase (92). A time course experiment in which the number of GFP spots was observed through a synchronous cycle indicated that the number of cells containing separated sisters exhibited a steady accumulation over time (Fig. 2B). Wild-type and *ctf18* Δ strains were arrested with α -factor for 2.5 h, released into nocodazolecontaining medium, and sampled every 10 min. The number of cell bodies containing two GFP signals was small for both wild-type and mutant strains early in the cycle, and increased as the cells traversed S phase and entered nocodazole arrest. The gradual, and early, appearance of cohesion failure suggests that mutant cells perform faulty cohesion establishment that results in slow decay of sister association.

The sister chromatid cohesion defect predicts that many mutant cells should not be able to recover after exposure to nocodazole. *ctf18*D cells bearing a galactose-inducible copy of *CTF18* were grown in repressing (glucose) or inducing (galactose) medium to early log phase and subjected to a 3-h nocodazole arrest. The cells were then spotted onto solid medium without nocodazole, and the ability to form microcolonies was scored the following day (Fig. 2C). Approximately 90% of *ctf18*D cells not expressing *CTF18* failed to recover. This result

FIG. 2. *CTF4* and *CTF18* are required for sister chromatid cohesion. (A) Early-log-phase cultures were treated to induce GFP-LacI expression, incubated for 3 h in rYPD containing nocodazole, and fixed in paraformaldehyde. For each strain, 100 cells exhibiting GFP fluorescence were scored within each experiment. The histogram shows the mean and standard deviation for a minimum of three trials. Strains were *CTF18* (AFS173), *ctf18* Δ (YJH17.2), *ctf4* Δ (YJH37), and *mad2-1* (AFS387). Note that approximately 5% of *ctf4* Δ or *ctf18* Δ cells have >1 chromosome III as determined during G1 arrest. (B) Kinetics of sister chromatid separation in *ctf18*D mutants. Log-phase cultures were arrested in a-factor and released into rYPD-nocodazole. After a-factor release, both strains sychronously completed S phase at 60 min postrelease as determined by flow cytometry (data not shown). YJH17.2 data were normalized using the α -factor arrest time point, to remove the contribution of cells containing >1 copy of chromosome III. A total of 100 informative cells were counted at each time point. (C) A *ctf18*Δ strain containing GAL-CTF18 on a plasmid (YJH48) was grown in selective medium containing either galactose or glucose. Log-phase cells were then shifted to rYPD-nocadozole for 4 h and then plated onto SC-URA glucose medium. CFU were counted after 24 h. The experiment was performed twice with similar results (mean values are shown).

is predicted by the data in Fig. 2A. If the GFP-marked chromosome separates with kinetics representative of other chromosomes, the appearance of \sim 30% of cells with two GFP spots predicts that \sim 5 of the 16 sister chromatid pairs in an average cell will have disassociated. Because dissociated sister chromatids are expected to segregate randomly, the mitotic division following drug removal is expected to result in very high inviability.

CTF18 **is required for mitotic condensation of the rDNA array.** In budding yeast, the cohesion defect observed in *mcd1, pds5*, and *trf4* mutants is accompanied by defective chromosome condensation (33, 36, 68, 108). To test whether *CTF18* protein is required for chromosome condensation, wild-type and *ctf18* cells were arrested in nocodazole-containing medium to obtain a uniformly staged culture at the point when condensation is complete. Formaldehyde-fixed cells were subjected to fluorescence in situ hybridization (32) with a digoxigenin-labeled DNA probe against the repetitive rDNA region. Bulk DNA was visualized with propidium iodide, and the rDNA probe was detected with antidigoxigenin and fluorescein isothiocyanate-conjugated secondary antibodies. Wild-type condensation of the rDNA region resulted in a distinct loop structure, as illustrated in the wild-type examples (Fig. 3A). However, 32% of *ctf18* Δ nuclei exhibited a decondensed rDNA staining pattern, indicating a condensation defect (Fig. 3). Thus, like the three other cohesion proteins for which mitotic chromosome condensation has been tested, Ctf18p is required for wild-type condensation of the rDNA as well as for sister chromatid cohesion.

Ctf4p and Ctf18p are stable proteins that associate with chromatin throughout the cell cycle. To determine whether the steady-state levels or chromatin associations of Ctf4p or Ctf18p were regulated, epitope-tagged alleles of both genes were generated by integration of a 9myc epitope in frame after the last codon (Fig. 4A). Both epitope-tagged alleles conferred wildtype stability to a test chromosome, a sensitive indicator of protein function (data not shown).

Analysis of Ctf4p and Ctf18p levels indicate that they do not vary dramatically throughout the cell cycle. To assay the accumulation levels of Ctf18p-9myc and Ctf4p-9myc, cultures were grown to early log phase at 30 $^{\circ}$ C, arrested for 2.5 h in α -factor, released into pheromone-free medium, and sampled every 10 min for 1.5 cell cycles. In a comparison with a β -tubulin standard, Ctf18p abundance varied over a three- to fivefold range, reaching a maximum during S phase and minimum during $G₂/M$ (Fig. 4B). These results are consistent with published mRNA levels (90), which show a shallow peak accumulation of *CTF18* transcript near the G_1/S boundary, and with the presence of two degenerate *Mlu*I cell cycle (MCB) boxes in the promoter region of *CTF18* (53). Arrest release experiments did not reveal significant fluctuations in *CTF4* protein levels in the cell cycle (data not shown).

Previous studies have shown that association between cohesion proteins and chromatin in budding yeast can be visualized by indirect immunofluorescence on spread chromosomes (48, 68). We detected both Ctf4p and Ctf18p association with chromatin in cells arrested in G_1 , S, or M phase (Fig. 5A and B) in this assay. Cells expressing epitope-tagged alleles were spheroplasted, lysed on glass slides in the presence of detergent, and washed to remove loosely adherent cellular contents. The presence and location of Ctf18p-9myc or Ctf4p9-myc proteins was compared with the location of DAPI-stained chromatin. Visible signals were observed in all samples of similar exposure, suggesting that some Ctf4p and Ctf18p is chromosome associated at each of these arrest points.

In a quantitative analysis, the proportion of Ctf4p or Ctf18p

B

FIG. 3. Absence of *CTF18* leads to a condensation defect. (A) Nocodazole-arrested *CTF18* (YPH499) and *ctf18*Δ (YE105) cells were assayed by fluorescence in situ hybridization using an rDNA probe. rDNA is detected by immunofluorescence (green); chromosomal DNA is stained with propidium iodide (red). A nucleus with decondensed rDNA is indicated by the arrow. (B) A total of 100 nuclei were scored for rDNA condensation status in four categories. Data were collected twice (averages are shown).

that remained bound to chromatin in a simple fractionation protocol was analyzed (57). Whole-cell extracts obtained from spheroplast lysis were subjected to centrifugation through a sucrose cushion, which separates a pellet fraction (enriched for chromosome-associated proteins) from a supernatant fraction. For both Ctf4p and Ctf18p, there was a three- to fourfold increase in bound protein between G_1 -and S-phase arrests (Fig. 5C). In contrast, we and others (57) noted a different binding pattern for Orc3p, another known chromatin binding protein. Furthermore, we noted electrophoretic mobility variants for both Ctf4p and Ctf18p in this protocol, whose significance is not yet apparent. The increase in association with the pellet fraction at S-phase arrest (HU) and early M-phase arrest (nocodazole) is consistent with roles for Ctf4p and Ctf18p in DNA replication and/or cohesion.

CTF18 **exhibits genetic interaction with replication mutants.** Genetic interaction between *CTF18* and a subset of cell cycle genes involved in DNA metabolism as well as other aspects of the chromosome cycle was tested by using a synthetic dosage lethality phenotype. In synthetic dosage lethality, a process may be efficiently disrupted if one interacting factor is present

at levels that interfere with complex formation in vivo and the function of another factor is limited by a hypomorphic mutation (25, 54, 55, 115). Since overexpression of *CTF18* from a galactose-inducible promotor (in p414GEU1/12) did not cause a growth defect, it could be used in a synthetic dosage lethality screen.

A subset of cell cycle mutants were chosen that included genes involved in DNA metabolism as well as other aspects of the chromosome cycle. Each mutant was transformed under noninducing conditions with either a GAL-*CTF18*-expressing minichromosome (p414GEU1/12) or with empty vector (p414GEU1). After induction, growth of the transformants with or without overexpression of *CTF18* was compared (Table 2). A temperature series ranging from 25 to 37°C was used to detect changes in maximum permissive growth temperature. A synthetic dosage lethal interaction was found with *cdc2-1, cdc7-4, cdc17-1*, and *cdc46-1*, each of which functions in an aspect of DNA synthesis. *CDC2* and *CDC17* encode catalytic subunits of polymerase δ and polymerase α , respectively (43). *CDC7* encodes a protein-kinase whose activity controls replication initiation, probably through modification of targets in the prereplication complex (64). *CDC46* encodes an essential protein of the prereplication complex, components of which are proposed to remain associated with the replicative fork (2). These genetic interactions extend the list of interactions already identified through traditional synthetic lethality (24, 87) and support the idea that Ctf18p interacts with replication fork proteins in vivo.

Ctf18p can interact with components of the RF-C complex. To search for proteins with which Ctf18p interacts, we screened a yeast cDNA library using the two-hybrid system. The coding region of *CTF18* was cloned in frame downstream of the GAL4 DNA binding domain (GAL4BD-CTF18). This fusion protein complemented the chromosome loss defect of a *ctf18*D mutant (data not shown). The screen identified two interacting genes whose fusion proteins were capable of conferring expression to three different GAL4-driven reporters: *HIS3, ADE2*, and the β-galactosidase gene (data not shown). Sequence analysis of the library plasmids indicated that they contained either *RFC3* or *RFC4*, each present as full-length fusions.

To investigate further, Rfc3p, Rfc4p, and Ctf18p-9myc were produced in vitro using a coupled transcription-translation system. Rfc3p and Rfc4p were radiolabeled using $[^{35}S]$ methionine. Unlabeled Ctf18-9myc product was incubated with each, in the presence of a polyclonal anti-myc antibody. Immune complexes were captured on protein A-agarose beads, washed, and analyzed by SDS-PAGE. Labeled Rfc3p and Rfc4p were enriched specifically and reproducibly through coimmunoprecipitation with Ctf18p (Fig. 6A).

To directly address whether Ctf18p interacts with Rfc3p and Rfc4p in vivo as well as with other members of the RF-C complex, 6HA epitope-tagged alleles were generated of *RFC2, RFC3, RFC4*, and *RFC5* in the presence and absence of the *CTF18*-9myc allele. In coimmunoprecipitation experiments, we found that Ctf18p-9myc precipitated with Rfc2p-6HA, Rfc3p-6HA, Rfc4p-6HA, and Rfc5p-6HA reproducibly (Fig. 6B). These results suggest the presence of an alternative RF-C complex, which we have termed RF-CCTF18 to distinguish it from the canonical, PCNA-loading RF-CRFC1. These results

FIG. 4. Ctf18p throughout the cell cycle. (A) α -factor arrest-release time course for CTF18-9myc cells (YJH40.4). CTF18-9myc was detected using anti-myc antibody and compared with β -tubulin (Tub 2p) on the same blot. (B) (Left) Fluctuation in Ctf18p-9myc accumulation was determined in normalized units by comparison of band intensity relative to b-tubulin. (Right) Separate aliquots from the same time course were prepared for flow cytometry and analyzed for DNA content (data not shown) as well as being scored for nuclear morphologies. These analyses indicated the execution of S phase by the majority of cells between 30 and 60 min and the initiation of M phase between 70 and 80 min.

are consistent with the observed homology between *CTF18* and *RFC1* and the in vitro interactions observed between Ctf18p-9myc, Rfc3p, and Rfc4p. Interestingly, it has been demonstrated that *RAD24*, another *RFC1* homolog, can also form a complex with the small RF-C subunits to facilitate alternative functions (31, 85).

Early purification methods for generating a biochemically

defined RF-C complex (promoting PCNA association with DNA) involved a variable species that migrated near the predicted molecular weight of Ctf18p. Therefore we analyzed an active purified fraction graciously donated by the Stillman laboratory to ascertain whether Ctf18p might be present as a substoichiometric component. Affinity-purified polyclonal antibody (10C) that provided robust detection of 2 ng of Ctf18p

FIG. 5. Ctf4p and Ctf18p are chromatin associated. Cells grown to early log phase in rich medium were arrested in α -factor (G₁), HU (S), or nocadozole (M) for 2.5 h. (A) Ctf18p-9myc (in YJH40.4) was detected using indirect immunofluorescence on chromosome spreads. For comparison, an untagged control strain (YPH277) in log phase is shown in the top row. Drug-induced arrests were verified by flow cytometry as shown. (B) Ctf4p-9myc (YJH38) cells were similarly analyzed. (C) Chromatin pellet fractions from cells containing Ctf18p-9myc (YJH40.4), Ctf4p-9myc (YJH38), and Orc3p (YJH62.6) were generated from arrested cell populations as above and analyzed by Western blotting. Similar results were obtained for Ctf18p-9myc and Ctf4p-9myc in four independent experiments. Results for Orc3p are consistent with previous data from Liang and Stillman (57). In these experiments, a pellet-to-supernatant cell equivalent loading ratio of 4:1 was used.

TABLE 2. Synthetic dosage lethality with CTF18*^a*

| Allele | Description | Phenotype ^b |
|-----------|--|--------------------------|
| $cdc2-1$ | DNA polymerase δ large subunit | Lethal at 28° C |
| $cdc7-4$ | Protein kinase, controls initiation of DNA synthesis | Toxic at 30°C |
| $cdc17-1$ | DNA polymerase α large subunit | Toxic at 34° C |
| $cdc46-1$ | Acts at origins to initiate replication | Toxic at 32°C |
| $rap1-5$ | Repressor/activator protein, affects telomere structure | Lethal at 30° C |
| $cdc9-1$ | DNA ligase | NP. |
| $cdc13-1$ | Binds telomeres | NP. |
| $cdc14-1$ | PTPase required for mitotic exit | NP. |
| $cdc23-1$ | Anaphase promoting complex subunit | NP. |
| $cdc28-1$ | p34CDC2 kinase homolog | NP. |
| $cdc34-2$ | Ubiquitin-conjugating enzyme, SCF subunit | NP. |
| $cdc4-1$ | Ubiquitin-conjugating enzyme, SCF subunit | NP. |
| $cdc6-1$ | Involved in DNA replication initiation | NP |

^a The growth of cells expressing GAL-CTF18 was compared with that of cells containing vector. *^b* The lowest temperature with a GAL-CTF18-induced phenotype is given. NP,

no synthetic phenotype; Lethal, no growth; Toxic, markedly slow growth. Descriptions are adapted from YPD (http://www.proteome.com) or SGD (http: //genome-www.stanford.edu/Saccharomyces).

by Western blot analysis (see Materials and Methods) did not detect a Ctf18p band in 100 ng of purified RF-C complex (Fig. 6C). Therefore we conclude that Ctf18p is not a detectable substoichiometric subunit of the biochemical activity characterized for processive DNA synthesis.

ctf18 **mutant cells contain short telomeres.** A screen for telomere length alteration in a collection of mutants exhibiting increased chromosome loss (91) identified a telomere length defect in all three alleles of *ctf18* (V. Lundblad and F. Spencer, unpublished data). The screen assayed telomere length following cleavage of genomic DNA with *Xho*I, which cuts in both subtelomeric Y' elements, to generate a broad 1.2 -to 1.5 -kb band, as well as in non-Y'-containing termini, to generate bands ranging in size from 2 to >6 kb. Both types of terminal restriction fragments exhibited a moderate reduction in size in the *ctf18* null strain, indicating that a loss of *CTF18* function influenced telomere length (Fig. 7).

Alterations in telomere length maintenance can be a consequence of either reduced telomerase function or defects in other telomere-associated proteins that control chromosome end protection and/or replication (22). Telomerase null mutants die on extended outgrowth, accompanied by progressive telomere shortening. This phenotype is enhanced in *rad52* mutants, which lack a compensatory recombinational pathway that maintains the telomere and hence cell viability (60). Clonal senesence was not observed in *ctf18* or *ctf18 rad52* mutants after 240 generations of outgrowth (data not shown). Therefore it is likely that the short telomere phenotype is not due to an absence of telomerase. Interestingly, the synthetic dosage lethality screen described above revealed a synthetic interaction with *rap1-5* in the presence of excess Ctf18p (Table 2). *RAP1* encodes a transcriptional repressor-activator DNA

FIG. 6. Ctf18p interacts with a subset of RF-C components but is not a component of purified RF-C. (a) Unlabeled Ctf18p-9myc was used to pull down [³⁵S]methionine-labeled Rfc3p or Rfc4p. The slowest-migrating bands are approximately the size expected for full-length products; the faster-migrating bands are consistent with the positions of in-frame translational start sites. Similar results were obtained from four independent trials. (B) Immunoprecipitation experiments were performed in yeast whole-cell extracts from strains containing 6HA epitope-tagged alleles of Rfc2p, Rfc3p, Rfc4p, or Rfc5p in the presence and absence of a CTF18-9myc allele. Duplicate SDS-PAGE gels loaded as shown were transferred and probed with anti-myc and anti-HA antibodies. The slowest-migrating species detected by the anti-myc antibody corresponds to Ctf18p-9myc. Species detected by the α -HA antibody are consistent with the expected migrations for RF-C proteins as indicated. Note that excess immunoprecipitate was loaded in the rightmost lane, accounting for the increased intensity of the background cross-reactive band. (C) A 100-ng portion of purified RF-C complex was probed for the presence of Ctf18p using affinity-purified antibody 10C. The antibody detected a strong band at 85 kDa in protein from wild-type cells (WT extract, from YPH499) that was absent from *ctf18*Δ cells (*ctf18*Δ extract, from YE105). Lanes 1 to 3 show the Western blot, and lane 4 is silver stained SDS-PAGE. Starred species are those identified by Fien and Stillman (23) and Cullmann et al. (17).

FIG. 7. *CTF18* participates in telomere length control. Meiotic segregants from a single tetrad derived from a *ctf18*D/*CTF18* heterozygote are shown. The broad band at 1.2 kb [detected by a radiolabeled poly(CA)_n/(GT)_n probe as in reference 84] contains termini from chromosomes with Y' elements.

binding protein with effects on both gene expression and telomere length; cells bearing the *rap1-5* mutation exhibit a decline in telomere length at semipermissive temperatures (61).

CTF18p homologues are detected in fission yeast and higher eukaryotes. Although Ctf18p has homology to RF-C subunits, the similarity is concentrated within an approximately 250 amino-acid region containing RF-C homology boxes II through VIII and falls off sharply outside of this region. This is clearly seen in a pairwise-alignment diagram with Rfc1p from budding yeast, the most homologous RF-C subunit (Fig. 8A). Interestingly, Ctf18p is also homologous to predicted *C. elegans, D. melanogaster, S. pombe*, and *H. sapiens* proteins. While these predicted proteins have significant homology over six of the eight RF-C boxes, the similarity to Ctf18p extends outward from the central region, indicating that the predicted proteins are more closely related to Ctf18p than to Rfc1p. This is clearly seen in an alignment between *S. cerevisiae* Ctf18p and a predicted *H. sapiens* Ctf18p (Fig. 8B). This observation is supported by cluster analysis (Fig. 8C) of a multiple alignment containing the Ctf18p homologs and Rfc1 proteins from *S. cerevisiae, C. elegans, D. melanogaster, S. pombe*, and *H. sapiens*. The Rfc1 proteins have been experimentally defined (71, 98) with the exception of the *C. elegans* predicted Rfc1p. In cluster analysis, the Ctf18p homologs form a group apart from the Rfc1 proteins. Additional work is required to determine if the candidate Ctf18p homologs are involved in sister chromatid segregation during mitosis.

DISCUSSION

In this study, we have identified a role for *CTF4* and *CTF18* in sister chromatid cohesion. Consistent with a cohesion defect, *ctf4* and *ctf18* mutants induce the spindle assembly checkpoint. An increased fraction of cellular Ctf4p and Ctf18p appears to associate with chromatin during early S- and early M-phase arrests. *ctf18*∆ mutants exhibit a chromosome condensation defect, similar to other cohesion mutants tested to date (33, 36, 108). *CTF18* is putative paralog of *RFC1*, and its encoded protein physically interacts with Rfc2p, Rfc3p, Rfc4p, and Rfc5p. However, it is not a component of the canonical PCNA clamp-loading RF-C complex. We and others have observed that *CTF4* and *CTF18* interact physically and genetically with many genes that function in DNA replication. These results directly address the hypothesis suggested by recent identification of a requirement for budding yeast DNA polymerase κ for the proper establishment of cohesion (108). The requirement for DNA polymerases and accessory factors in

FIG. 8. Candidate *CTF18* orthologues and conserved RF-C boxes. (A and B) Modified output of National Center for Biotechnology Information pairwise BLAST (BLOSUM62) of representative sequences to illustrate the distribution of similarity. Roman numerals indicate RF-C homology boxes, the hatched box represents the ligase homology domain, and black bars mark the relative positions of RF-C boxes. (A) ScCtf18p versus ScRfc1p (e-value = 1e⁻¹⁵). (B) ScCtf18p versus HsCtf18p (e-value = $3e^{-35}$). (C) Proteins from *C. elegans, D. melanogaster, S. pombe*, and *H. sapiens* were identified among the top hits from a PSI-BLAST search using *S. cerevisiae* Ctf18p as the query. Clustal X v1.8 (44) was used to make a multiple alignment of proteins from *C. elegans, D. melanogaster, S. pombe*, and *H. sapiens*. Using the multiple alignment, a bootstrapped neighbor-joining tree was produced. The analysis shows that Ctf18p homologs cluster away from Rfc1p homologs.

cohesion strongly supports the hypothesis that cohesion establishment is a job performed by proteins associated with replication fork complexes.

Checkpoints and cohesion. The *MAD2*-dependent preanaphase delay observed in both *ctf4* and *ctf18* mutants is predicted after cohesion failure, because dissociation of a sister chromatid pair should relieve tension on the kinetochore-microtubule junction, thereby activating the spindle assembly checkpoint. As noted previously (87), cohesion failure is not repaired following activation of the spindle assembly checkpoint but, rather, appears to be augmented during mitotic arrest. In agreement, we have observed a marked reduction in the viability of *ctf18* cells after a 3-h checkpoint-induced arrest. While the classic model of checkpoint function emphasizes the opportunity allowed for repair of an inducing lesion, in this case a checkpoint-induced delay favors removal of damaged cells from the dividing population.

Interestingly, Kouprina et al. (53) noted a *RAD9*-dependent arrest in a *ctf18* null mutant at 11°C. In contrast, at the normal growth temperature (30°C) we detected no appreciable *RAD9* or $MEC1$ dependence of the preanaphase delay in $\text{cf }18\Delta$ cells; therefore we propose that the 11°C observation reveals an additional physiological consequence. Perhaps at low temperature *ctf18*D cells accumulate defects that are detected by the DNA damage checkpoint. Paradoxically, Kouprina et al. (53) found that *ctf18* Δ *rad9* Δ cells exhibit increased (rather than decreased) viability at the nonpermissive temperature. We propose an explanation based on the observed sister chromatid cohesion defect. A delay induced by the *RAD9*-dependent checkpoint will probably augment the frequency of cohesion failure in $\text{cf18}\Delta$ cells, similar to a nocadozole-induced preanaphase delay (Fig. 2B). Thus, removal of the checkpoint may explain the enhanced cell survival.

If checkpoint arrest promotes cell death in cohesion-defective cells, why do viable cohesion mutants exhibit aneuploidy? One potential explanation is that adaptation of the spindle assembly checkpoint allows at least some cells with separated chromosomes to progress through mitosis. Cohesion-defective mutants are frequently isolated in chromosome loss screens. Within a collection of viable chromosome loss mutants encompassing approximately 60 genes (91), alleles of six genes required for cohesion (*CTF4, CTF7, CTF18, SMC1, MCD1*, and *SCC2*) have been identified to date. This observation suggests that cohesion defects may underlie a significant proportion of naturally occurring chromosome instability phenotypes.

CTF18 **as an RFC1 paralog.** Similarity between Ctf18p and Rfc1p has been noted previously (Cullmann et al. 1995), and BLASTP alignment analyses using the entire predicted yeast proteome indicate that these two proteins exhibit the highest degree of homology to one another (e-value = $3e^{-15}$ [Fig. 8]). In a two-hybrid screen, we identified strong interactions of Ctf18p with both Rfc3p and Rfc4p. In vitro immunoprecipitation experiments confirmed the ability of Ctf18p to directly interact with in vitro-generated Rfc3p and Rfc4p. Furthermore, in vivo coimmunoprecipitation studies indicate that Ctf18p interacts with Rfc2p, Rfc3p, Rfc4, and Rfc5p. Independent studies have also identified a complex which contains Ctf18p, Rfc2p, Rfc3, Rfc4p, and Rfc5p protein (M. Mayer and P. Hieter, personal communication). Recent studies indicate that RF-C homology box IV, a highly conserved β -sheet found

in both Rfc1p and Ctf18p, may be required to mediate interactions between Rfc1p and PCNA (1). In addition, the region between RF-C homology boxes IV and VIII has been implicated in RF-CRFC1 subunit interactions (5). However, Ctf18p is not detected in a purified, biochemically active RF-C fraction. Together, these observations support the concept of a novel CTF18-containing RF-C like complex $(RF-C^{CTF18})$ distinct from PCNA-loading RF-CRFC1.

This is not the first reported instance of an alternative RF-C like protein complex. ScRAD24p exhibits limited homology to Rfc1p observed in alignments of multiple family members (62). Rad24p physically associates with Rfc2p, Rfc3p, Rfc4p, and Rfc5p (31, 85). Moreover, *RAD24, RFC2*, and *RFC5* are required for DNA damage checkpoint activity (31, 75, 85), consistent with the formation of a functional $RF-C^{RAD24}$ protein complex. Based on structural and genetic analyses, it has been proposed that the RF-CRAD₂₄ complex may provide a clamploading activity to a PCNA-like trimer (80, 105).

Roles of *CTF4* **and** *CTF18* **in association with DNA replication proteins.** The synthetic dosage lethality results we report here, as well as the physical association of CTF18p with RFC3p and RFC4p, add to the growing list of genetic and physical interactions involving *CTF4, CTF18*, and components of the replication complex (Fig. 9). These interactions strongly suggest that the molecular functions of *CTF4* and *CTF18* are closely related to DNA synthesis. They also support a model in which sister chromatid cohesion defects may be among the consequences of mutations in other proteins that function at DNA replication forks. Although previous studies have suggested that Ctf4p and Ctf18p each play roles associated with DNA metabolism, these roles have not been defined. Intriguingly, *CTF4* was isolated as a DNA polymerase α (Cdc17p) binding protein, and genetically interacts with with *CDC17, RAD27, DNA2*, and *RFC1* (24). *ctf4* mutants exhibited general properties consistent with DNA synthesis defects, including an elevated rate of mitotic recombination and an accumulation of preanaphase cells in log phase (51, 69). However, previous work indicated that the *ctf4*-induced preanaphase delay was not *RAD9* or *MEC1* dependent (69), and therefore it was unlikely that compromised DNA replication caused the observed preanaphase delay. The presence of a cohesion defect offers an explanation for the preanaphase delay and suggests a new direction for elucidation of the molecular role of Ctf4p.

Similarly for *CTF18*, genetic interactions and mutant phenotypes have also supported an uncharacterized role in DNA metabolism. As noted by Kouprina et al. (53), *CTF18* exhibited similarity to RF-C subunits and an elevated rate of mitotic recombination. Formosa and Nittis (24) observed traditional synthetic lethality between *CTF18* and both *DNA2* and *CTF4*. The synthetic dosage lethality reported here, with temperature-sensitive alleles of the replication genes *CDC2, CDC17, CDC46*, and *CDC7*, provide additional indication of a molecular function that influences the DNA synthesis machinery. Physical association with Rfc3p and Rfc4p in vivo and in vitro further supports this concept. However, the new observations of a spindle assembly checkpoint-dependent delay and a sister chromatid cohesion defect provide impetus to the search for a molecular role for Ctf18p in this aspect of chromosome metabolism.

FIG. 9. Physical and genetic interactions among *CTF4, CTF18, CTF7*, and DNA synthesis proteins. The bold circles denote replication-associated proteins affecting sister chromatid cohesion. Physical associations are indicated by symbol overlap, synthetic lethality is indicated by solid lines with double arrows, and synthetic dosage lethality is indicated by dashed lines with single arrows. The interactions are described in this work [1] and references 87 [2], 112 [3], 24 [4], 69 [5], 11 [6], 56 [7], 4 [8], 28 [9], 67 [10], 114 [11], and 27 [12]. The physical interactions among RF-C subunits are described in references 17, 23, 75, and 104.

CTF18 **and telomere structure.** The short telomere length displayed by *ctf18* null mutants, as well as the previously isolated *ctf18* alleles, demonstrates that *CTF18* is required for normal telomere structure. Since a *ctf18* null mutant does not exhibit a telomere replication defect comparable to that of a telomerase null mutant, it is unlikely that this is due to an absence of telomerase. However, proteins important for full telomere replication may play structural rather than enzymatic roles. For example, the *Mre11-Xrs2-Rad50* complex, which has been proposed to play a structural role in DNA repair of sister chromatids, may facilitate telomere replication by presenting chromosome ends to telomerase for replication (76). Ctf18p may also make a structural contribution to telomere length maintenance. Specifically, we suggest that loss of cohesion at chromosome ends in a *ctf18* mutant decreases the efficiency of complete telomere replication. This hypothesis is attractive in light of the proposal that telomerase functions as a dimer at chromosome termini (79).

Cohesion and replication. A temporal linkage between DNA synthesis and sister chromatid cohesion is suggested by the tight association between sister chromatids observed from the time of their generation until anaphase (32). Furthermore, the specificity of sister association must be derived from something other than sequence similarity because the efficiency of pairing between homologous chromosomes in a diploid does not indicate a comparable intimate physical relationship (12, 13). The establishment of sister chromatid cohesion as a part of the chromosomal duplication process provides an attractive model that satisfies the specificity requirement, as well as the time of appearance, of the association between sisters.

Indirect evidence for a relationship between DNA synthesis and sister chromatid cohesion is found in several recent studies, indicating that known cohesion proteins possess S-phase roles. *S. cerevisiae* Mcd1p, Ctf7p, Scc2p, and Scc4p must all be functional during S phase (16, 87, 101, 102), as must fission yeast Mis4p, an Scc2p homologue (26). In addition, *S. pombe* Eso1p includes extensive homology to both Ctf7p and polymerase η (96) in distinct domains. It has been recently proposed that this type of homology configuration, suggestive of an evolutionary gene fusion or splitting event, may indicate that where the two polypeptides are encoded separately, they nonetheless function together in the cell (21, 63). This view suggests that Ctf7p in *S. cerevisiae* acts together with a DNA polymerase activity. Finally, the novel polymerase κ protein family in budding yeast not only has a demonstrated requirement in replication of the genome but also plays a poorly understood role in establishing and/or maintaining robust cohesion between sisters (108). The independent observation that both *CTF4* and *CTF18* possess functions important for sister chromatid cohesion strongly supports the concept that replication forks directly mediate molecular events important for sister chromatid cohesion.

The proposed roles for *CTF4* and *CTF18* proteins in coupling S-phase DNA replication and cohesion do not preclude potential activities that are also important for cohesion maintenance. Several observations are consistent with roles for each of these proteins in cohesion establishment, such as genetic and physical interaction with replication proteins, the early defect in sister association observed for *ctf18*D cells, and the augmented chromatin association of both Ctf4p and Ctf18p during early S phase suggested by cell fractionation. However, a detailed analysis of rapid-response conditional alleles is required to evaluate functional execution points for these genes. Whether replication-associated proteins will contribute solely to the establishment of robust cohesion or will be important for later steps in the chromosome cycle that support the maintenance of cohesion until anaphase remains to be seen. We note that neither *CTF4* nor *CTF18* is an essential gene, although cohesion and replication are essential functions. The synthetic lethality observed for double null cells is consistent with each making a complementary but nonoverlapping molecular contribution to the essential function they share. Whether this represents spatial, temporal, or biochemical differentiation awaits further study.

The identification of polymerase κ , a new class of polymerase with a distinct function, is consistent with the previously observed division of labor among other characterized polymerases in budding yeast (reviewed in references 19 and 43). It has been proposed that cohesion may be established by polymerase κ through a mechanism whereby this specialized replication fork variant is required for replication through particular chromosomal regions (108; reviewed in reference 95). In support of this idea, the distribution of cohesion on chromatin

is observed to be nonrandom (9, 30, 37, 66, 97), with evident areas of enrichment. Interestingly, recent studies have led to an attractive model for polymerase switching necessary for the transition from the primase function of polymerase α to the highly processive extension functions of polymerases δ and ε (106, 117, 118). Under this model, sequential competitions for common binding sites on single-stranded binding protein and the PCNA sliding clamp support the notion that the polymerase switch occurs in two steps: displacement of polymerase α by RF-CRFC1 followed by displacement of RF-CRFC1 by polymerase δ . It is tempting to speculate that RF-CCTF18 may promote a switch from the polymerase δ or ε complex to polymerase κ at sites of cohesion, in a role analogous to that of RF-CRFC¹. This switch might incorporate chromatin association or activation of cohesion accessory factors such as CTF7 (87, 101) or the SCC2-SCC4 complex (16). We note that *CTF18* is not an essential gene, although cohesion and replication are essential functions. Perhaps in a *ctf18* null mutant, Rfc1p partially substitutes for Ctf18p, similar to the proposed functional substitutions between Trf4p and Trf5p (15, 108).

Taking a broad view, it is not yet clear whether a cohesion role is specific to polymerase κ or whether the regulation of sister chromatid association is a general job of replication forks. In favor of the general scenario, the tight association between Ctf4p and polymerase α large subunit (69) is consistent with a cohesion role for polymerase α -primase-mediated replication. Interestingly, bovine *SMC1* and *SMC3* candidate orthologues encode proteins characterized within the recombination protein complex RC-1, which also contains DNA polymerase ε (45, 46). Moreover, *S. cerevisiae* Mcd1p is required for wild-type radiation resistance (38), as is its candidate orthologue *Rad21* in *S. pombe* (7, 8). The involvement of cohesion subunit proteins in DNA recombination and in repair suggests that these processes may incorporate cohesion remodeling activities in vivo. Perhaps the absence of a juxtaposed sister chromatid caused by compromised cohesion may lead to increased utilization of a homolog in recombinational DNA damage repair. This hypothesis may explain increased rates of mitotic recombination in cohesion-deficient mutants. Although not well understood, the association between cohesion and repair-recombination pathway activities strongly suggests a role for sister chromatid cohesion regulation in many aspects of DNA metabolism.

The requirement for *CTF4* and *CTF18* in cohesion identifies a role for these replication accessory proteins in high-fidelity chromosome segregation. The proposed intimate coupling between DNA synthesis and the establishment of sister chromatid cohesion by one or more DNA polymerase complexes provides a mechanism for the specificity of this tight and essential chromatin association. However, it is not yet clear how many distinct steps or molecular complexes operate in cohesion, which must incorporate sufficient flexibility to support both local and global dynamic chromatin restructuring events, including transcriptional responses, DNA repair, DNA replication, and mitotic chromosome condensation. It is to be anticipated that the protein complexes operating within mechanisms governing the establishment, maintenance, and release of sister chromatid cohesion are just coming to light.

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