Mcm10 has potent strand-annealing activity and limits translocase-mediated fork regression

Ryan Mayle*, Lance Langston*,b, Kelly R. Molloy*, Dan Zhang*, Brian T. Chait*,1,2, and Michael E. O’Donnell*a,b,1,2

*Laboratory of DNA Replication, The Rockefeller University, New York, NY 10065; bHoward Hughes Medical Institute, The Rockefeller University, New York, NY 10065; and 1Laboratory of Mass Spectrometry and Gaseous Ion Chemistry, The Rockefeller University, New York, NY 10065

Contributed by Michael E. O’Donnell, November 19, 2018 (sent for review November 8, 2018; reviewed by Zvi Kelman and R. Stephen Lloyd)

The 11-subunit eukaryotic replicative helicase CMG (Cdc45, Mcm2-7, GINS) tightly binds Mcm10, an essential replication protein in all eukaryotes. Here we show that Mcm10 has a potent strand-annealing activity both alone and in complex with CMG. CMG-Mcm10 unwinds and then reanneals single strands soon after they have been unwound in vitro. Given the DNA damage and replisome instability associated with loss of Mcm10 function, we examined the effect of Mcm10 on fork regression. Fork regression requires the unwinding and pairing of newly synthesized strands, performed by a specialized class of ATP-dependent DNA translocases. We show here that Mcm10 inhibits fork regression by the well-known fork reversal enzyme SMARCAL1. We propose that Mcm10 inhibits the unwinding of nascent strands to prevent fork regression at normal unperturbed replication forks, either by binding the fork junction to form a block to SMARCAL1 or by reannealing unwound nascent strands to their parental template. Analysis of the CMG-Mcm10 complex residues 557–558 (15–19) identifies Mcm10 interactions with six CMG subunits, with the DNA-binding region of Mcm10 on the N-face of CMG. This position on CMG places Mcm10 at the fork junction, consistent with a role in regulating fork regression.

DNA replication | fork regression | CMG helicase | SMARCAL1 | Mcm10

DNA replication is performed by a replisome composed of numerous proteins that work together to separate and duplicate DNA strands to form two new daughter duplexes (1–5). The replication machinery is conserved in all eukaryotes, and consists of an 11-subunit helicase known as CMG (Cdc45, Mcm2-7, GINS) (6); DNA polymerase Pol α-primase, which primes both strands; and Pols ε and δ, which duplicate the leading and lagging strands, facilitated by proliferating cell nuclear antigen (PCNA) clamps and the clamp loader, replication factor C. Numerous other factors move with replisomes, the functions of which are only now being elucidated. Most replisome accessory factors are nonessential but have roles in genome stability; however, the Mcm10 replisome accessory factor is essential in all eukaryotes tested thus far (7, 8).

Structural studies have shown that Mcm10 contains a globular “internal domain” within which are two DNA-binding elements: an oligonucleotide/oligosaccharide-binding fold and a zinc finger (9, 10). We refer to this globular internal domain as the DNA-binding region. The DNA-binding region of Mcm10 is flanked by N- and C-terminal regions. The N-terminal region of Mcm10 contains a sequence that forms a triple strand α-helical coiled coil and is proposed to mediate Mcm10 oligomerization and/or facilitate interactions with other proteins (11). The C-terminal region of Mcm10 varies in length and complexity across organisms (reviewed in ref. 12). Homolog alignments indicate that the Saccharomyces cerevisiae Mcm10 N-region comprises residues 1–149, the globular DNA-binding region comprises residues 150–356, and the C-region comprises residues 1,585–1,615 (16).

Mcm10 was recently shown to tightly bind CMG, forming a CMG-Mcm10 complex (CMGM), which stimulates both CMG helicase activity and the rate of the replisome (13, 14). Mcm10 is required for CMG activation at origins during replication initiation (15–19). Along with binding CMG, Mcm10 is documented to bind Pol α and PCNA (9, 20). Studies of Mcm10 depletion or loss of function using genetics, cell biology, and cell extracts have identified Mcm10 functions in replisome stability, fork progression, and DNA repair (21–25). Despite significant advances in the understanding of Mcm10’s functions, mechanistic in vitro studies of Mcm10 in replisome and repair reactions are lacking.

The present study demonstrates that Mcm10 on its own rapidly anneals cDNA strands even in the presence of the single-strand (ss) DNA-binding protein RPA, a property previously associated with the recombination protein Rad52 (26). While CMG does not perform strand annealing, CMGM unwinds DNA and then rapidly rewinds the unwound products. Several helicases demonstrate both DNA unwinding and annealing (reviewed in ref. 27). The combined functions are proposed to be involved in such processes as double-strand (ds) break repair, fork regression, stabilization of stalled forks, transcription, and telomere metabolism (27). Roles for Mcm10 in some of these processes have been suggested by studies involving mutation or depletion of Mcm10 (21–25).

To more precisely understand the role of Mcm10 strand annealing activity, we determined the location of the DNA-binding region of Mcm10 on CMG by cross-linking mass spectrometry (CX-MS). This analysis showed that Mcm10 is in close proximity to 6 of the 11 CMG subunits and extends from the N-terminal to the C-terminal domains of CMG. The DNA-binding region of Mcm10 localizes to the N-face of CMG, which places DNA binding by Mcm10 within a moving replisome at the fork junction (28). This location suggests a possible role of Mcm10 in fork regression,

Significance

Fork regression is a way of circumventing or dealing with DNA lesions and is important to genome integrity. Fork regression is performed by double-strand DNA ATPases that initially cause newly synthesized strands to unpair from the parental strands, followed by pairing of the new strands and reversal of the fork. This study shows that Mcm10, an essential replication factor, efficiently anneals complementary strands and also inhibits fork regression by SMARCAL1. Moreover, the study localizes the Mcm10 DNA-binding domain to the N-terminal domains of the replicative CMG helicase at the forked nexus. Thus, forks that are unimpeded would contain Mcm10 at a strategic position where its DNA-binding and/or annealing function may block fork regression enzymes and thereby protect active forks from becoming reversed.

Author contributions: R.M., L.L., K.R.M., B.T.C., and M.E.O. designed research; R.M., L.L., K.R.M., and D.Z. performed research; R.M., L.L., K.R.M., B.T.C., and M.E.O. analyzed data; and R.M., L.L., and M.E.O. wrote the paper.

Reviewers: Z.K., University of Maryland; and R.S.L., Oregon Health & Science University.

The authors declare no conflict of interest.

This open access article is distributed under Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND).

1B.T.C. and M.E.O. contributed equally to this work.
2To whom correspondence may be addressed. Email: Brian.Chait@rockefeller.edu or odonnell@rockefeller.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1819107116/-/DCSupplemental.

Published online December 31, 2018.
important to genomic stability (29–31), and is consistent with the findings of studies of Mcm10 in metazoans (22).

Fork regression is catalyzed by a specialized class of ATP helicases that reverse forks rather than unwind them, most notably SMARCAL1, ZRANB3, and HLTF (32). In the present study, we examined the effect of S. cerevisiae Mcm10 on SMARCAL1 using previously established DNA substrates for this enzyme (33). We found that Mcm10 completely inhibits SMARCAL1 on several forked DNA structures tested here. We propose that Mcm10 is held in proximity to the fork by its known tight interaction with CMG (13). Mcm10-mediated inhibition of SMARCAL1 could be due to nonspecific binding of Mcm10 to ssDNA, or by Mcm10’s annealing activity, either of which could prevent fork regression. We propose a possible regulatory mechanism that prevents the reversal of actively replicating forks and targets fork regression only to those forks that have DNA damage.

Results
CX-MS of CMG-Mcm10. Mcm10 tightly binds CMG and greatly enhances helicase activity (13). To better understand how Mcm10 functions with CMG, we determined the location of Mcm10 on CMG. To delineate the position of Mcm10 on CMG, we reconstituted and purified CMGM then performed CX-MS on the complex. MS analysis of CMGM treated with a bifunctional lysine cross-linking agent revealed an extensive network of intermolecular and intramolecular cross-links between Mcm10 and six subunits of CMG (a total of >1,000 cross-links) (Fig. 1, SI Appendix, Fig. S1, Movie S1, and Dataset S1). We observed cross-links between the N region of Mcm10 and CMG, specifically with Mcm2, Mcm5, Mcm6, Cdc45, Psf1, and Psf2 (Fig. 1, Movie S1, and Dataset S1). The results indicate a central location of Mcm10 on the N-face of CMG near the interface between the Mcm2-7 ring and its accessory factors. In addition, some interactions extend around the outer perimeter of CMG all of the way to the C-face. These CX-MS results greatly extend earlier studies demonstrating the interaction of Mcm10 with the N-terminal regions of Mcm2 and Mcm6 (14, 34).

The globular DNA-binding region of Mcm10 contains eight residues that cross-link to CMG, and all but one of these 31 cross-links are located on the N-face of CMG (Dataset S1). The cryoEM structure of CMG on a forked DNA with ATP shows CMG tracks on DNA N-face first (28), and thus the Mcm10 DNA-binding region is located on the front edge of CMG, directly facing the fork junction. Interestingly, there are many cross-links between CMG and the N- and C-regions of Mcm10 but few cross-links to the Mcm10 DNA-binding region, suggesting that the N- and C-regions of Mcm10 may establish the main affinity of Mcm10 to CMG. This leads us to predict that Mcm10’s DNA-binding domain may be mobile on CMG. We note that most Mcm subunits contain long N-tails that are not visible in cryoEM structures, indicating mobility in these regions (35, 36), and Mcm10 may possibly dock onto these mobile Mcm N-tails.

Fig. 1. CX-MS proximity assays of CMG-Mcm10. Cross-links between Mcm10 and CMG on performing CX-MS on the reconstituted CMG-Mcm10 complex are colored and encircled in dark purple. Four different views of the CMG are shown: N-face (A); C-face (B); Cdc45, side view (C); and GINS, side view (D). More information is provided in Dataset S1 and Movie S1.
The CX-MS results of CMG-Mcm10 revealed extensive cross-links between the C-region of Mcm10 and CMG, consistent with a previous study reporting an interaction of Mcm2 with a deletion mutant of Mcm10 that contained only the C-region (37). Our detailed CX-MS study unexpectedly revealed that the C-region of Mcm10 wraps around the entire outside surface of Mcm6 from the N-face to the C-face of CMG (Fig. 1C and Dataset S1), suggesting that the C-region of Mcm10 may be highly elongated. A few Mcm10 cross-links also extend to the C-face of CMG on the Psf1 subunit (Fig. 1D). In addition, we observed three intramolecular Mcm10 crosslinks between identical residues, consistent with an oligomeric state of Mcm10 mediated by the N-region coiled-coil domain (Dataset S1) (11).

**CMG-Mcm10 Reanneals Unwound Single Strands**. Our previous CMG-Mcm10 helicase studies were performed in the presence of a trap oligonucleotide, a standard addition to unwinding assays (13). We performed unwinding reactions using CMG with and without Mcm10 in the presence of a trap oligo (Fig. 2A). CMG helicase is greatly stimulated by Mcm10, as we reported previously (13). However, when the trap oligo is omitted from helicase assays (Fig. 2B), unwound products appear to be reannealed (Fig. 2C). CMG unwinding is initially stimulated by Mcm10, but the separated strands are rapidly converted back into the duplex substrate. We also tested purified CMGM complex compared with CMG without Mcm10 and again observed unwinding followed by reannealing when Mcm10 is bound to CMG (Fig. 2D).

**Mcm10 Is a Potent DNA Strand-Annealing Protein**. To directly test Mcm10 for annealing activity, we mixed the two individual oligos that compose the mini fork—a 3′-P-5′ end-labeled 5′ flap lagging strand oligo and a 3′ flap leading strand oligo at 20 nM—and incubated them with or without Mcm10 in the absence of ATP and CMG. The addition of a 20-fold molar excess of unlabeled lagging strand oligo was used to quench reactions (SI Appendix, Fig. S2). Reactions were analyzed by native PAGE to assess the rate at which complementary strands anneal with and without Mcm10 (Fig. 3A). Spontaneous annealing at 30 °C without Mcm10 was nearly undetectable, but the addition of Mcm10 resulted in rapid and nearly complete annealing under the same conditions (Fig. 3 B and C). The concentration dependence of Mcm10 annealing is shown in SI Appendix, Fig. S3. CMG alone did not demonstrate annealing activity (SI Appendix, Fig. S4). We also tested fully complementary single strands and observed comparable Mcm10 annealing activity. These experiments demonstrate a previously unreported biochemical activity of Mcm10 in which Mcm10 rapidly anneals cDNA strands.

While Mcm10 has robust strand-annealing activity, it seems paradoxical that Mcm10 also stimulates CMG in DNA unwinding. To explore whether CMG affects Mcm10 annealing activity, we compared Mcm10 with and without the addition of a twofold molar excess of CMG in the annealing assay (Fig. 4A and B). The results show no substantial effect of CMG on the rate of Mcm10 annealing, as the difference in proportion of substrate annealed is within the error of the triplicate experiments (Fig. 4C). These results are consistent with the observation that reconstituted CMGM complex also displays reannealing in the unwinding assay (Fig. 2D) and indicate that Mcm10 annealing remains active while in complex with CMG. We note that at an unperturbed replication fork in vivo, the leading strand template is readily duplicated by Pol ε, so the unwound strands would not be single-stranded and thus would not be expected to reanneal.

**Mcm10 Catalyzes Strand Annealing in the Presence of RPA**. RPA binds ssDNA tightly and is presumed to prevent complementary single strands from reannealing. Several proteins have been identified that promote annealing of RPA-coated ssDNA, most notably Rad52, which is involved in homologous recombination in yeast (26). To study the capability of Mcm10 to reanneal ssDNA in the presence of RPA, we titrated RPA into one strand and empirically determined that 40 nM RPA saturates the 5′ flap lagging strand oligo as determined by electrophoretic mobility shift assay (EMSA) gel shift in a native gel (Fig. 5A). We then performed annealing assays by adding 40 nM RPA to both the leading and lagging strand oligos and then mixing them to initiate the annealing reaction. Reactions were quenched using SDS to remove protein along with a 20-fold excess of unlabeled oligo trap. Spontaneous strand annealing of the RPA-coated
SMARCAL1, ZRANB3, S. cerevisiae

January 15, 2019

Fig. 4. CMG does not inhibit Mcm10 reannealing. (A) Schematic of the reaction. (B) Mcm10 (60 nM) was added with or without CMG (120 nM) to standard annealing assays, and timed aliquots were analyzed by native PAGE. The 0 s lane is a mixture of the two oligonucleotides without protein and stopped within 30 s. (C) Quantitation of reactions. Values are the average of three experiments. Error bars represent 1 SD.

oligos showed very little annealed duplex (Fig. 5B). However, Mcm10 efficiently annealed the RPA coated strands (Fig. 5B), with only a 25% decrease in annealed product compared with Mcm10 activity with no RPA (Fig. 5C).

Mcm10 Inhibits Fork Regression. Studies focusing on Mcm10 loss of function indicate that Mcm10 is important for DNA repair as well as for fork progression and stability of the replisome (21–25). Our present CX-MS findings show that the DNA-binding region of Mcm10 binds the N-terminal face of CMG (Fig. 1). In addition, our earlier cryoEM structure of CMG at a fork shows that the N-terminal face of CMG is directed at the forked junction (28).

Taken together, these two structural observations indicate that Mcm10 is located at or near the fork junction. This places Mcm10 in a position where it could play a role in regulating fork regression. Recent cellular studies have demonstrated that fork regression is a primary response to replication stress or damage (23–31). A subset of SWI/SNF family dsDNA translocases—SMARCAL1, ZRANB3, and HLTF—has been shown to facilitate reversal of stalled or damaged forks to maintain genomic integrity (29–31). Fork regression involves the unwinding and subsequent pairing of nascent strands to form a four-arm Holliday junction that can be migrated to further reverse the fork.

Given the phenotypic consequences of Mcm10 mutation or depletion, and considering that Mcm10 binds to ssDNA (Kd = 50 nM; SI Appendix, Fig. S5) and catalyzes strand annealing in the presence of RPA (Fig. 5), we explored the impact of Mcm10 on fork regression. To do so, we used the well-characterized fork reversal enzyme SMARCAL1 and DNA substrates used previously to study SMARCAL1: forked substrates with a single-strand gap on either the leading or the lagging strand arm (33).

The fork regression substrates contain two mismatches at the base of the fork that prevent spontaneous branch migration/fork reversal. SMARCAL1 catalyzes branch migration through the mismatched region, resulting in two duplex products: a duplex of the nascent strands and a duplex of the parental strands (Fig. 6A).

We expressed and purified human SMARCAL1 in a yeast expression system (SI Appendix, Fig. S6) and tested the impact of S. cerevisiae Mcm10 on fork reversal. SMARCAL1 titrations and time courses showed that both substrates were reversed by SMARCAL1 (SI Appendix, Fig. S7), with the lagging gap substrate reversed more efficiently compared with the leading gap substrate. At 2 nM SMARCAL1, we observed >50% reversal in 2 min for the lagging gapped DNA (Fig. 6A and SI Appendix, Fig. S7). The leading gapped fork DNA required 5 nM SMARCAL1 for 5 min to yield comparable results (Fig. 6B and SI Appendix, Fig. S8).

Titration of Mcm10 into SMARCAL1 fork reversal reactions showed strong inhibition of SMARCAL1 on both substrates. We also formed a DNA fork with no ssDNA gap and found that Mcm10 still inhibited fork reversal by SMARCAL1 (SI Appendix, Fig. S9). These results are consistent with studies of Xenopus laevis Mcm10 demonstrating that Mcm10 binds ssDNA and dsDNA with similar affinities (38). Two conceivable ways in which Mcm10 may inhibit fork regression are (i) the use of Mcm10 annealing activity in which Mcm10 reverses the initial SMARCAL1-induced unwinding of the nascent strands from the parental strands, a necessary first step in forming the four-way junction needed for fork reversal, or (ii) nonspecific binding of Mcm10 to ssDNA or dsDNA, blocking fork regression. We examine these processes in more detail below.

Discussion

The robust strand-annealing activity of Mcm10 demonstrated in this study seems to run counter to the fact that Mcm10 stimulates CMG unwinding activity (13, 14). However, during normal replication, the two daughter strands are replicated to form two duplexes, leaving no complementary single strand for Mcm10 to anneal. As has been proposed for other helicases that both unwind and anneal DNA, CMG-Mcm10 annealing activity may be involved in ds break repair, fork reversal, transcription, and/or telomere metabolism (27). In fact, Mcm10 loss of function studies indicate that Mcm10 is involved in DNA repair and replisome stability in higher eukaryotes (21–25). Below we examine a hypothesis to possibly explain how Mcm10 may regulate fork reversal, known to be required for genome integrity. We also propose a possible consequence of Mcm10’s annealing activity at origins that may necessitate the use of Pol δ in leading strand synthesis.

Mcm10 Inhibition of Fork Regression. Our current data demonstrate that S. cerevisiae Mcm10 inhibits fork regression catalyzed by the fork reversal enzyme SMARCAL1 (human). Fork regression is thought to occur after DNA damage when the leading polymerase uncouples from the CMG helicase (29–31). Fork regression is likely regulated and prevented at active replisomes. Based on our present results demonstrating Mcm10 inhibition of
fork reversal, we propose that Mcm10 may inhibit regression of active forks, helping specify fork regression to damaged forks. The physiological link of Mcm10 to fork regression is suggested by several studies correlating Mcm10 mutation or depletion to replication stall stability and DNA repair (21–25) and link Mcm10 to dsDNA repair enzymes that are proposed to act during fork regression repair (25, 29–31).

Our CX-MS analysis reveals that the DNA-binding region of Mcm10 is located at the N-face of CMG (Fig. 1, Movie S1, and Dataset S1). Combined with our earlier finding that CMG tracks first on the DNA N-face (28), this places Mcm10’s binding region at the front of CMG, facing the fork junction (Fig. 7A). This location is consistent with at least two possible mechanisms for Mcm10 inhibition of fork regression: (i) Mcm10 nonspecifically binds the ssDNA of a forked DNA that is stalled by a lesion on the leading strand, thereby inhibiting fork reversal, as observed in an earlier study using *Escherichia coli* ssDNA-binding protein (33), and (ii) Mcm10 prevents fork reversal by its strand-annealing activity, inhibiting the initial unpairing step in fork reversal in which newly synthesized DNA strands must unpair from their parental DNA strands before pairing to form the fourth arm. The present study demonstrating Mcm10’s robust strand-annealing activity reveals the potential for the second possibility, but we cannot distinguish between the two processes (or any others) at this time.

One may question how fork regression is targeted to DNA-damaged forks but not actively replicating forks. We found that *S. cerevisiae* Mcm10 binds to ssDNA rather weakly (K$_D$ = 50 nM; SI Appendix, Fig. S5), similar to the 120 nM K$_D$ value for the more extensively characterized *X. laevis* Mcm10 (38). This relatively weak interaction with DNA might be harnessed for regulation. We hypothesize that when held to the fork by CMG during active replication, Mcm10 would have a high effective concentration due to proximity effects. Thus, Mcm10 could be an even more potent inhibitor of fork regression when part of an active replication fork than was observed in this study using isolated Mcm10 (Fig. 7A).

On encountering a leading strand lesion, CMG and Pol δ are thought to uncouple, which may lead to dissociation of CMG from DNA. CMG is not highly processive on its own (6, 39), and *S. cerevisiae* CMG–Mcm10 unwinding is 10-fold slower than when coupled to active replication forks (13). CMG dissociation would nullify the proximity effect of Mcm10 at a fork and possibly, given the low affinity of Mcm10 to DNA, no longer prevent fork regression (Fig. 7B). Experiments that enable simultaneous CMG-mediated replisome function and SMARCAL-mediated fork regression are needed to explore this hypothesis.

**Fig. 6.** Mcm10 inhibits SMARCAL1-catalyzed fork regression. Fork regression assays of SMARCAL1 with increasing Mcm10 added. (A) 2 nM SMARCAL1 on a lagging single-strand gap substrate at 2 min of reaction. (B) 5 nM SMARCAL1 on a leading single-strand gap substrate at 5 min of reaction. The two arms of the forked DNA are complementary, and substantial spontaneous regression is prevented by two mismatches at the fork junction, indicated by a kink in the duplex. Quantitation is the average of triplicate experiments. Error bars represent 1 SD.
knowledge about Mcm10–CMG interactions. Moreover, CX-MS analysis places the DNA-binding region of Mcm10 on the N-face of CMG near the DNA fork junction, where it is poised to counteract fork regression at an active replication fork. The present study provides a conceptual framework for future studies exploring the regulatory balance of replication fork progression and fork regression.

Materials and Methods

CX-MS. Reconstituted CMG-Mcm10 complex was cross-linked using disuccinimidyl suberate (DSS) in reactions containing –12 μg of CMG-Mcm10, 1 mM DSS, and 5% DMSO. Reactions were incubated for 25 min at room temperature with agitation at 1,200 rpm, then quenched with 50 mM (final concentration) NH₄CO₃ and methanol-precipitated. Following reduction and alklylation of cysteines, the sample was separated by SDS/PAGE in a 3–8% Tris-acetate gradient mini gel. Cross-linked products were then digested and analyzed by MS. Further details are provided in SI Appendix.

Helicase Assays. The unwinding substrate was a fork DNA with a 3pS- end-labeled lagging strand. Unwinding was assayed by incubating 0.5 nM substrate with 30 or 50 nM CMG with and without 60 nM Mcm10 (as indicated in the figure legends) in the presence of 1 mM ATP at 30 °C. Timed aliquots were withdrawn, quenched, and then analyzed by native PAGE. More information is provided in SI Appendix.

Annealing Assays. Partially complementary oligos forming a fork were combined on ice at a ratio of 10 nM leading strand to 20 nM 3pS- end-labeled lagging strand. Mcm10 was added immediately after combining the two oligos, and reactions were incubated at 30 °C. Reactions were stopped by adding a 10-fold excess of unlabelled lagging strand and then flash-freezing. Reactions were then analyzed by native PAGE. The product of fork regression is observed as a faster-migrating band on the gel that forms when the “nascent” strands are ejected and the 3pS-labeled “parental” gel bands remain as a ddDNA product. More details are provided in SI Appendix.

Fork Regression Assays. A substrate composed of four oligos was constructed by first annealing the two leading strand and two lagging strand oligos to form leading and lagging halves, which were subsequently annealed for 20 min at 30 °C to form the full substrate. Two mismatches at the fork junction prevented exponential spontaneous reversal. Fork regression activity was assayed by incubating SMARCA1 and/or Mcm10 with 0.5 nM substrate at 37 °C with 2 mM ATP. Reactions were quenched, flash-frozen, and then analyzed by native PAGE. The product of fork regression is observed as a faster-migrating band on the gel.

ACKNOWLEDGMENTS. We appreciate the help of Dr. Grant Schauer of the M.E.C. laboratory for determining the Kd value of Mcm10 to DNA. We thank Drs. Huilin Li for suggesting the possibility that Mcm10 may bind the mobile N-tails of Mcms. This research was funded by National Institutes of Health (NIH) Grant R01 GM115809 and the Howard Hughes Medical Institute (to M.E.O.), NIH Grant R01 GM103314 (to B.T.C.), and NIH Grant T32 CA09673 (to R.M.).

1. Bai L, et al. (2017) Architecture of the Saccharomyces cerevisiae replisome. Adv Exp Med Biol 1042:207–228.
2. Bell SP, Labib K (2016) Chromosome duplication in Saccharomyces cerevisiae. Genetics 203:1027–1067.
3. Burgers PMJ, Kunkel TA (2017) Eukaryotic DNA replication fork. Annu Rev Biochem 86:417–438.
4. Li H, O’Donnell ME (2018) The eukaryotic CMG helicase at the replication fork: Emerging architecture reveals an unexpected mechanism. Bioessays 40:98. 5. O’Donnell ME, Li H (2018) The ring-shaped hexameric helicases that function at DNA replication forks. Nat Struct Mol Biol 25:122–130.
6. Moyer SE, Lewis PW, Botchan MR (2006) Isolation of the Cdc45/Mcm2-7-GINS (CMG) complex, a candidate for the eukaryotic DNA replication fork helicase. Proc Natl Acad Sci USA 103:10236–10241.
7. Bielsinska AK (2016) Mcm10: The glue at replication forks. Cell Cycle 15:3024–3025.
8. Thu YM, Bielsinska AK (2014) McM10: One tool for all—integrity, maintenance, and damage control. Semin Cell Dev Biol 39:120–131.
9. Das-Bradso D, Ricke RM, Bielinsky AK (2006) Interaction between PCNA and diubiquitinated histones H2A and H2B: Implications for the regulation of PCNA-dependent processes. Mol Cell 21:1892–1901.
10. Du W, Stauffer ME, Eichman BF (2012) Structural biology of replication initiation factors Mcm10. Structure 20:1932–1941.
11. Du W, Stauffer ME, Eichman BF (2012) Structural biology of replication initiation factor Mcm10. Subcell Biochem 62:197–216.
12. Langton LD, et al. (2017) Mcm10 promotes rapid isomerization of CMG-DNA for replisome bypass of lagging strand DNA blocks. eLife 6:e29918.
13. Lóoke M, Maloney MF, Bell SP (2017) Mcm10 regulates DNA replication elongation by stimulating the CMG replicative helicase. Genes Dev 31:391–305.
14. Douglas ME, Ali FA, Costa A, Difffey JX (2018) The mechanism of eukaryotic CMG helicase activation. Nature 555:265–268.
15. Heller RC, et al. (2011) Eukaryotic origin-dependent DNA replication in vitro reveals sequential action of DDK and S-CDK kinases. Cell 146:80–91.
16. Kanke M, Kodama Y, Takahashi T, Nakagawa T, Masukata H (2012) Mcm10 plays an essential role in origin DNA unwinding after loading of the CMG components. EMBO J 31:2182–2194.
17. van Deursen F, Sengupta S, De Piccoli G, Sanchez-Diaz A, Labib K (2012) Mcm10 associates with the loaded DNA helicase at replication origins and defines a novel step in its activation. EMBO J 31:2195–2206.
18. Watase G, Taksaki H, Kanemaki MT (2012) Mcm10 plays a role in functioning of the eukaryotic replicative DNA helicase, Cdc45/Mcm-GINS. Curr Biol 22:343–349.
19. Warren EM, Huang H, Fanning E, Chazin WJ, Eichman BF (2009) Physical interactions between Mcm10, DNA, and RNA polymerase alpha. J Biol Chem 284:24662–24672.
20. Alver RC, et al. (2014) The N-terminus of Mcm10 is important for interaction with the 9-11 clamp and in resistance to DNA damage. Nucleic Acids Res 42:8388–8404.
21. Chadhia GS, Gamba A, Gillespie P, Brown JJ (2016) Xenopus Mcm10 is a CDK-substrate required for replication fork stability. Cell Cycle 15:2183–2195.
22. Chattopadhyay S, Bielsinska AK (2007) Human Mcm10 regulates the catalytic subunit of DNA polymerase-alpha and prevents DNA damage during replication. Mol Biol Cell 18:4085–4095.
23. Park JH, Kang SW, Jeon Y, Kang S, Hwang DS (2008) Knockdown of human Mcm10 exhibits delayed and incomplete chromosome replication. Biochem Biophys Res Commun 365:575–582.