



The intra-S phase checkpoint directly regulates replication elongation to preserve the integrity of stalled replisomes

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DNA replication is dramatically slowed down under replication stress. The regulation of replication speed is a conserved response in eukaryotes and, in fission yeast, requires the checkpoint kinases Rad3^{ATR} and Cds1^{Chk2}. However, the underlying mechanism of this checkpoint regulation remains unresolved. Here, we report that the Rad3^{ATR}-Cds1^{Chk2} checkpoint directly targets the Cdc45-MCM-GINS (CMG) replicative helicase under replication stress. When replication forks stall, the Cds1^{Chk2} kinase directly phosphorylates Cdc45 on the S275, S322, and S397 residues, which significantly reduces CMG helicase activity. Furthermore, in *cds1^{Chk2}*-mutated cells, the CMG helicase and DNA polymerases are physically separated, potentially disrupting replisomes and collapsing replication forks. This study demonstrates that the intra-S phase checkpoint directly regulates replication elongation, reduces CMG helicase processivity, prevents CMG helicase delinking from DNA polymerases, and therefore helps preserve the integrity of stalled replisomes and replication forks.

replication fork stability | checkpoint | CMG complex

Eukaryotic DNA replisomes must overcome an extensive array of replication barriers to complete genomic DNA replication (1). These barriers include a great number of potential DNA secondary structures (e.g., G4, hairpins, triplexes), numerous proteins that tightly bind to DNA, various DNA lesions, and collisions with transcription apparatus (2–8). Most cancer cells also need to contend with dysregulated cell cycle progression that results in replication stress (9, 10). The cause of oncogene-induced replication stress is not fully understood but includes altered origin firing, increased replication–transcription clashes (11), misregulation of ribonucleotide reductase, and exhaustion of deoxyribonucleotide triphosphates (dNTPs) (12). All of these can result in the stalling of replication forks, which potentially causes fork instability, gross genomic alterations, and cancer (5, 13–16).

Stalling replication forks are unstable, and the intra-S phase checkpoint is absolutely essential for preventing such stalling forks from collapsing (5, 17–19). However, despite extensive studies in the past two decades, the reasons why stalling forks are unstable has not been fully defined. It is also not known precisely how the checkpoint acts to stabilize stalling forks. Previous studies have shown the formation of reversed forks and reported the dissociation of replication factors from stalled forks in checkpoint-deficient cells (17, 19–21). However, it remains to be clarified whether fork reversal and the dissociation of replication factors from stalling forks are a cause or a consequence of fork collapse. Most likely, fork collapse is a multistep process during which fork reversal and/or replication factor dissociation lead to increasingly severe fork damage until irreversible fork collapse occurs. In contrast to the findings of Cobb and Lucca (20, 21), who reported that DNA polymerases are lost from sites of replication stalling in budding yeast, more recent work from De Piccoli et al. (22) did not detect the dissociation of specific replication factors from

stalling forks. The reasons for these contradictory data are unclear but may be related to the duration for which forks were stalled, how the replication proteins were isolated by chromatin immunoprecipitation (ChIP), and the sensitivity of the assays used to measure the global changes of specific replication factors within replisomes. For example, a 10 to 20% difference in a Western blotting assay may not appear very striking, but a collapse rate of 10 to 20% of the total replication forks would indeed be severe and may possibly be beyond cells capability to fix and thus complete DNA replication.

It has been reported that the Mrc1/Claspin protein forms a fork protection complex with Timeless/Tof1 and Tipin/Csm3 (Swi1 and Swi3 in *Schizosaccharomyces pombe*) and that this plays important roles in stabilizing stalling forks (23, 24). Mrc1 also plays a critical role in checkpoint activation and in the physical association between the replicative helicase and DNA polymerase ϵ (25–27). Thus, the requirement of Mrc1 for the stability of stalling forks is likely to be related to both its role in the checkpoint response and

Significance

The intra-S phase checkpoint is essential to stabilize stalling/stalled replication forks. However, the mechanism underlying checkpoint regulation is poorly understood; two key questions remain to be answered: 1) why are stalling forks unstable when checkpoint is deficient, and 2) which protein(s) is the critical target of checkpoint regulation? Here, we demonstrate that CMG replicative helicase becomes uncoupled from stalled/blocked DNA polymerases in checkpoint-deficient *cds1^{Chk2}* Δ cells, likely reflecting physical separation of the helicase and polymerases. This uncoupling is prevented by the Cds1^{Chk2}-mediated phosphorylation of Cdc45. We propose that replication elongation is directly regulated in response to replication fork stalling via phosphorylation of Cdc45, which significantly reduces CMG helicase processivity, in order to prevent helicase–polymerase uncoupling and promote replication fork stability.

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its function in the physical association of the CMG replicative helicase and DNA polymerase ϵ at replication forks. Besides the fork protection complex, the checkpoint also regulates several enzymes, preventing them from damaging stalling forks (28–30). In addition, in response to replication stress, an important aspect of checkpoint regulation is to inhibit late origin firing (31–35), which will reduce the number of replication forks encountering replication stress, control potential DNA damage, and therefore increase cell survival rate.

Furthermore, it was reported that, in budding yeast, the rate of DNA replication was dramatically slowed down in response to replication stress caused by DNA alkylation, and this was dependent on the checkpoint kinases Mec1 and Rad53 (36), indicating that replication slowing is an active process. In fission yeast checkpoint-deficient cells ($cds1^{Chk2}\Delta$ or $mrc1\Delta$), DNA synthesis is prolonged in the presence of HU (hydroxyurea treatment) and remains dependent on the MCM helicase, and HU causes increasing fork collapse and DNA damage (37). Dna2, an essential flap endonuclease for processing Okazaki fragments in replication forks, is regulated by Cds1^{Chk2} for preventing fork reversal in response to replication fork stalling (17). Taken together, these studies suggest that checkpoint may directly target the replisome and regulate the replication elongation step.

When replication is experimentally impeded by HU (a common tool for studying replication fork stalling that inhibits ribonucleotide reductase, thus impeding DNA polymerization), the ATR-dependent intra-S phase checkpoint stabilizes stalling replication forks such that fork pausing is transient and can be relieved when the HU is removed. Furthermore, when the checkpoint is deficient, stalling forks rapidly collapse, leading to irreparably dysfunctional replication (18, 38), pathological reversed forks (17, 19), DNA breaks (5), and unscheduled recombination (13, 39). However, due to the fact that only a limited number of checkpoint targets have been identified and their regulation mechanisms elucidated, two key questions remain to be answered (14, 17, 40, 41): 1) Why are stalling forks unstable when the checkpoint is deficient, and 2) which protein factor or factors are the critical target of regulation by checkpoint?

In fission yeast, the Cds1^{Chk2} kinase defines the major intra-S phase branch of the Rad3^{ATR} checkpoint. In mammalian cells, this ATR-dependent function is performed mainly by Chk1, a functional homolog of yeast Chk2. Mammalian Chk2 responds to ATM activation in response to DSBs (42). Fission yeast $cds1^{Chk2}\Delta$ cells are hypersensitive to HU due to rapid fork collapse and global replication failure (17, 43). We therefore searched for intra-S phase checkpoint targets by screening for spontaneous genetic mutations that conferred HU resistance to fission yeast $cds1^{Chk2}\Delta$ cells. Our rationale was that, if a mutation located in a checkpoint target partially mimics the intra-S phase checkpoint regulation, it will render checkpoint-deficient cells more HU resistant. We identified an *R319I* mutation in *cdc45* (*cdc45-R319I*) encoding a subunit of the CMG (Cdc45-MCM-GINS) replicative helicase. *cdc45-R319I* rescued the 2.5 mM HU sensitivity of $cds1^{Chk2}\Delta$ cells by ~600- to 3,000-fold, suggesting that Cdc45 may be a crucial target of the checkpoint response to replication stress. Subsequently, we demonstrate that Cdc45 is directly targeted by the Rad3^{ATR}-Cds1^{Chk2} checkpoint pathway: when replication forks stall in the presence of HU, Cds1^{Chk2} directly phosphorylates Cdc45 on S275, S322, and S397. This Cds1^{Chk2}-mediated Cdc45 phosphorylation dramatically reduces CMG helicase activity, resulting in the slowing of replication elongation.

Using a ChIP assay (44), we further demonstrate that the CMG replicative helicase becomes uncoupled from stalled/blocked DNA polymerases in checkpoint-deficient $cds1^{Chk2}\Delta$ cells, likely reflecting the physical separation of the helicase and polymerases. This uncoupling is prevented by the Cds1^{Chk2}-mediated Cdc45 phosphorylation. Thus, we propose that replication elongation is directly regulated in response to replication fork stalling via the phosphorylation of Cdc45, which reduces CMG helicase processivity in order to help prevent helicase-polymerase uncoupling and promote replication fork stability.

Results

Cds1^{Chk2} Targets the Replicative Helicase CMG Complex. To explore how the intra-S phase checkpoint pathway regulates replication forks, we established a genetic screen to identify mutations that

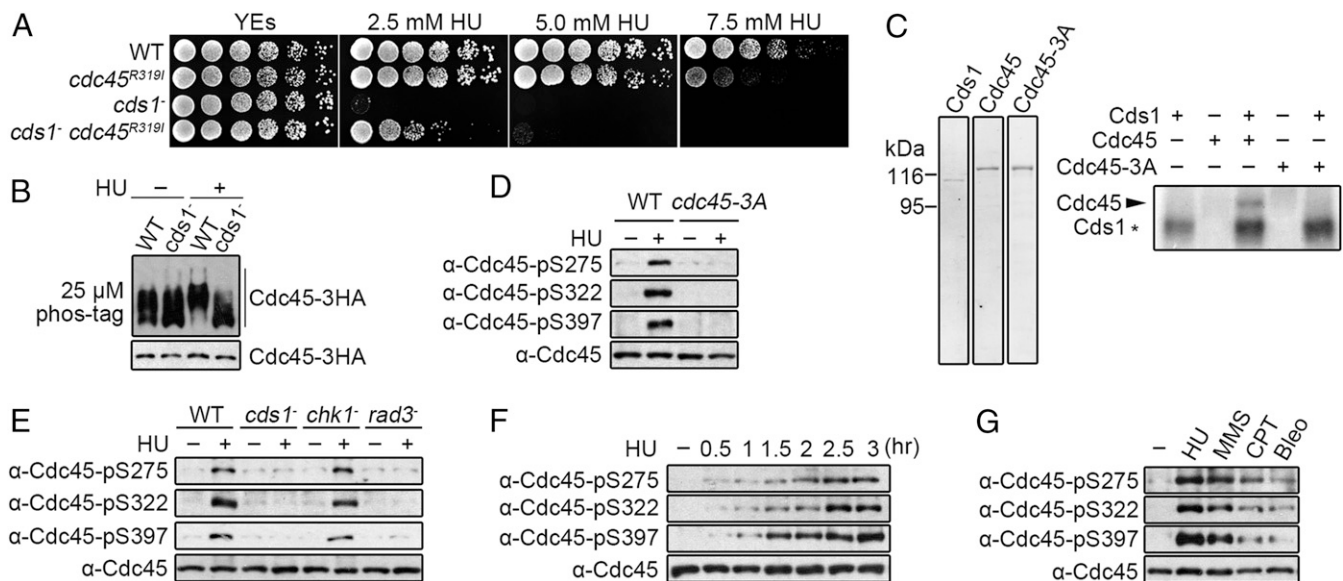


Fig. 1. Cds1^{Chk2} phosphorylates Cdc45 on S275, S322, and S397 when replication forks stall. (A) A fivefold serial dilution of the indicated strains grown under the designated concentration of HU. (B) Cdc45 phosphorylation in response to HU. (Top) Phos-tag sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE). (Bottom) SDS-PAGE. HA-tagged Cdc45 was detected by α -HA. (C, Left) Purified Cds1, Cdc45-MBP-tag, and Cdc45-S275A-S322A-S397A (S3A). (Right) Autoradiograph of purified Cdc45 following in vitro phosphorylation by Cds1^{Chk2}. (D) Detection of Cdc45 phospho-S275, -S322, and -S397 in vivo using phospho-specific antibodies. (E) Phosphorylation of Cdc45 S275, S322, and S397 in the indicated strains. (F) Time course of S275, S322, and S397 phosphorylation following HU treatment. (G) Cdc45 phosphorylation in response to the indicated genotoxic agents.

may mimic checkpoint regulation. We reasoned that mutants which render *cds1^{Chk2}Δ* cells resistance to replication fork stalling by HU treatment (45) may be located in checkpoint targets and may act by mimicking checkpoint regulation. *cds1^{Chk2}Δ* cells were plated on rich media agar containing 5 mM HU, and colonies emerging after several days of incubation (SI Appendix, Fig. S1 A and B) were isolated and confirmed for HU resistance (SI Appendix, Fig. S1C). Genomic DNA was then extracted and subjected to deep sequencing. The mutations identified were individually reintroduced into a *cds1^{Chk2}Δ* background to establish if they restored HU resistance. A *cdc45-R319I* mutation was identified that dramatically suppressed the HU sensitivity of *cds1^{Chk2}Δ* cells (Fig. 1A). When assayed in a *cds1⁺* background, cells harboring the *cdc45-R319I* mutation were not sensitive to 2.5 mM HU and exhibited little sensitivity to 5.0 mM. At higher concentrations, sensitivity became more pronounced (Fig. 1A). The *cdc45-R319I* mutation is recessive; introducing a second wild-type copy of *cdc45* by integrating *cdc45⁺* into the *ura4* locus prevented the rescue of HU sensitivity of *cds1^{Chk2}Δ* cells by *cdc45-R319I* (SI Appendix, Fig. S1D).

Following HU treatment, we observed that Cdc45 displayed a *cds1^{Chk2}*-dependent mobility shift on sodium dodecyl sulphate-polyacrylamide gel electrophoresis gels containing 25 μM Phos-tag, indicating that Cdc45 is phosphorylated in response to fork stalling (Fig. 1B). Mass spectroscopy (MS) determined that the S275, S322, and S397 sites of Cdc45 are phosphorylated (SI Appendix, Fig. S1E). In vitro, Cds1^{Chk2} directly phosphorylated Cdc45-MBP-tag (Fig. 1C), and no Cdc45 phosphorylation was observed when the S275, S322, and S397 residues were mutated to alanine, confirming that S275, S322, and S397 were also in vitro Cds1^{Chk2} targets (Fig. 1C). Together with the analysis of the *cdc45-R319I* mutant, the Cds1^{Chk2}- and replication stress-dependent phosphorylation of Cdc45 strongly suggests that Cdc45 is a checkpoint target in response to fork stalling. We thus developed our subsequent studies to confirm the Cds1^{Chk2}-dependent phosphorylation of Cdc45 and examine its physiological function and mechanism in stabilizing stalling replication forks. The *cdc45-R319I* mutant was not analyzed further since it is a random untargeted mutation, although it did provide the clue that Cdc45 may be regulated in response to replication stress.

Phospho-specific antibodies were prepared against each of the three sites (SI Appendix, Fig. S1F) and used to confirm that the in vivo phosphorylation of Cdc45 on S275, S322, and S397 depended on presence of HU, Cds1^{Chk2}, and Rad3^{ATR} but not on Chk1 (Fig. 1D and E). Furthermore, all three serine residues were

concurrently phosphorylated over time upon HU exposure (Fig. 1F), and the phosphorylation was elicited more avidly by agents that interfere with replicative polymerases (HU or methyl methanesulphonate [MMS]) when compared to agents such as camptothecin (CPT) that predominantly block replicative helicase progression or bleomycin (Bleo) that induces DNA breaks (Fig. 1G). Taken together, these results demonstrate that Cdc45 is phosphorylated on S275, S322, and S397 by Cds1^{Chk2} in response to replication fork stalling by agents that interfere with DNA synthesis.

S275, S322, and S397 are all located within the CID domain where Cdc45 interacts with MCM and GINS. S275 and S322 but not S397 lie within a consensus motif for Cds1^{Chk2} in *S. pombe* (SI Appendix, Fig. S1G). The Cds1^{Chk2} consensus motif is I/LxR/KxxS/T with some flexibility (46, 47). S322 and S397 are highly conserved from yeast to human cells. S275 is not conserved, but human Cdc45 has threonine and serine residues (three and seven residues, respectively) from the site corresponding to S275 of *S. pombe* Cdc45. These sites are conserved from *Xenopus laevis* to *Homo sapiens* and both are within a loose Chk2 consensus motif. Thus, Cds1^{Chk2}-mediated Cdc45 phosphorylation on S275, S322, and S397 may be conserved in eukaryotes.

***cdc45-3A* and *cdc45-3D* Mutations Stabilize Stalling/Stalled Replication Forks in Checkpoint-Deficient Cells.** To examine the role of Cds1^{Chk2}-mediated Cdc45 phosphorylation in stabilizing stalling forks, S275, S322, and S397 were mutated individually or all together, either to alanine (phospho deficient) or aspartic acid (potential phospho mimic). Individual mutations displayed similar growth rates, and HU sensitivities compared to wild-type (wt) cells (Fig. 2A). Mutations at S275 or S322 did not rescue the HU sensitivity of *cds1^{Chk2}Δ* cells, whereas *cdc45-S397A* and *cdc45-S397D* modestly reduced the HU sensitivity only at low HU concentrations (Fig. 2B). However, the triple *cdc45-3A* and *cdc45-3D* mutations significantly rescued the HU sensitivity of *cds1^{Chk2}Δ* cells (Fig. 2C). From these spot-test assays, in which cells are grown under a chronic exposure to HU, we estimated an ~600- to 3,000-fold increase in survival when compared to *cds1^{Chk2}Δ cdc45⁺* cells. A clonogenic survival assay, in which cells were transiently exposed to 12.5 mM HU for 3 h, led to increased cell survival when compared to the *cds1^{Chk2}Δ* single mutant of ~15- to 20-fold (Fig. 2D). Both the *cdc45-3A* and *cdc45-3D* mutants showed a similar level of modest sensitivity to high HU concentration (7.5 mM) in a *cds1⁺* background when compared to *cdc45⁺* cells (Fig. 2C and D). The basically equivalent rescue effect of the

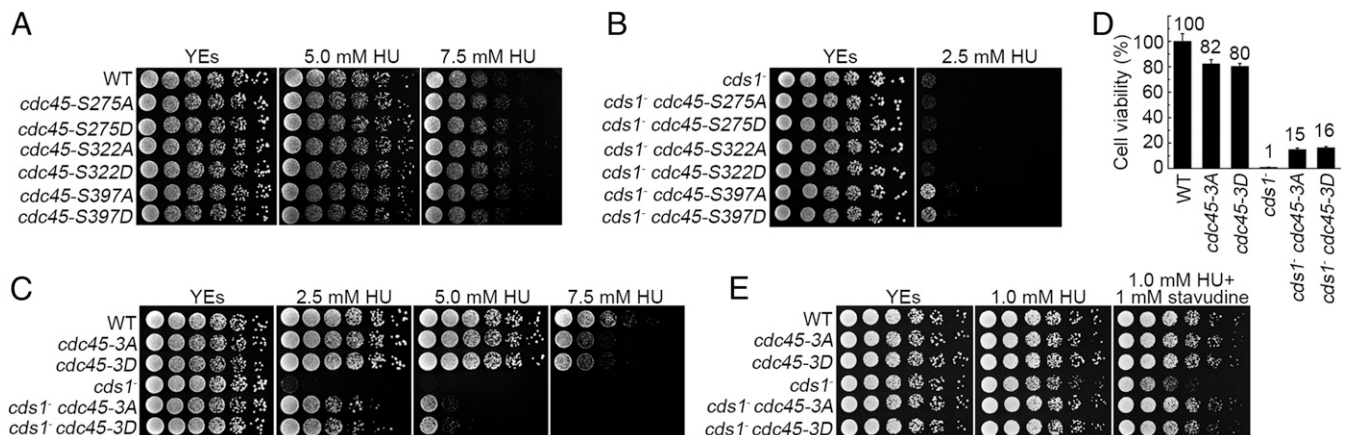


Fig. 2. Cdc45-3A/3D mutations significantly rescue the HU sensitivity of *cds1^{Chk2}Δ* cells. (A) Fivefold serial dilution assays of the indicated single *cdc45* phospho-site mutant strains grown with the indicated HU concentrations. WT = *cdc45⁺* control. (B) Equivalent serial dilution assay in a *cds1^{Chk2}Δ* genetic background. (C) Equivalent serial dilution assay of the indicated triple phospho-site *cdc45* mutations in *cds1⁺* (WT) and *cds1Δ* backgrounds. (D) The relative cell viability of the indicated strains after treatment with 12.5 mM HU for 3 h. The data are presented as the mean ± SD. (E) Serial dilution assay of the indicated triple phospho-site *cdc45* mutations in *cds1⁺* (WT) and *cds1Δ* backgrounds in the presence of the dideoxynucleoside analog stavudine.

cdc45-3A and *cdc45-3D* mutations on the HU sensitivity of *cds1^{Chk2}Δ* cells suggests that both alleles alter the biochemical property of the CMG helicase in a similar way (see Fig. 5).

The critical role of the Cdc45 residues S275, S322, and S397 for the biochemical property of CMG helicase was further examined. These three serine residues were mutated to either glutamate (*cdc45-3E*, bearing a larger residual group compared to aspartate), threonine (*cdc45-3T*, similar to serine except having an extra methyl group), cysteine (*cdc45-3C*, similar to serine but with thiol replacing hydroxyl group), lysine (*cdc45-3K*, positive charge), asparagine (*cdc45-3N*, possessing a chemically polar residual group), glycine (similar to alanine but possessing a smaller residual group), and valine (hydrophobic residual group). These new mutants were all examined for their growth in the presence or absence of HU and for their rescue effect on the HU sensitivity of *cds1^{Chk2}Δ* cells. As shown in *SI Appendix, Fig. S1H*, these mutants, with the exception of *cdc45-3E*, which exhibited a severe growth defect, grew similarly well compared to *cdc45⁺* cells in normal medium (HU absent). The viable mutants, with the exception of *cdc45-3T*, showed apparent growth sensitivity to 5.0 or 7.5 mM HU (*SI Appendix, Fig. S1H*), and the level of the HU sensitivity among the mutants was slightly different, with *cdc45-3A*, *cdc45-3D*, *cdc45-3C*, *cdc45-3K*, and *cdc45-3V* being more sensitive to HU than *cdc45-3N* and *cdc45-3G* (*SI Appendix, Fig. S1H*). The *cdc45-3E* mutant did not grow in the presence of 5.0 or 7.5 mM HU (*SI Appendix, Fig. S1H*). With the exception of *cdc45-3E* (which could not be measured due to a severe growth defect), all the *cdc45* mutants rescued the HU sensitivity of *cds1^{Chk2}Δ* cells. The levels of rescue were different: *cdc45-3A*, *cdc45-3D*, and *cdc45-3C* > *cdc45-3K*, *cdc45-3V* > *cdc45-3N*, and *cdc45-3G* > *cdc45-3T* (*SI Appendix, Figs. S1I and S5D*). These results suggest that the S275, S322, and S397 residues in Cdc45 are critical for preserving the normal activity/function of CMG helicase; a slight difference either in spatial conformation (threonine contains an extra methyl group compared to serine) or in chemical polarity (i.e., cysteine versus serine) alters the properties of the CMG helicase, rendering *cds1^{Chk2}Δ* cells more resistant to HU. The following studies focused on the *cdc45-3A* and *cdc45-3D* mutations in order to gain further insight into the mechanism of how the checkpoint regulates the CMG helicase to stabilize stalling/stalled replication forks.

HU stalls replication forks by limiting the availability of dNTPs, thus limiting the processivity of the replicative polymerases. To establish if Cdc45 phosphorylation site mutants could rescue *cds1^{Chk2}Δ* cell sensitivity to other polymerase stalling agents, we

examined the response to stavudine. Stavudine is a dideoxyribonucleoside analog that is converted to active stavudine triphosphate by cellular kinases in vivo and subsequently incorporated into the leading and lagging strands. The incorporation of stavudine blocks further incorporation of downstream nucleotides and thus stalls replication forks (48). In order to enhance the chance of stavudine incorporation, the cellular concentration of dNTPs was reduced by including 1 mM HU in medium. When treated with 1 mM HU, all strains showed similar growth rates, except *cds1^{Chk2}Δ* cells that showed a very modest sensitivity (Fig. 2 E, Middle). When both 1 mM of stavudine and 1 mM of HU were present, wt, *cdc45-3A*, and *cdc45-3D* strains still displayed a similar growth rate, but *cds1^{Chk2}Δ* cells were ~25-fold more sensitive when compared to wt cells (Fig. 2 E, Right). However, the *cds1^{Chk2}Δ cdc45-3A* and *cds1^{Chk2}Δ cdc45-3D* strains showed an ~25-fold of reduction in stavudine sensitivity when compared to *cds1^{Chk2}Δ* cells. These data further confirm that the *cdc45-3A* and *cdc45-3D* mutations significantly reduce the sensitivity of *cds1^{Chk2}Δ* cells to reagents that stall replication forks.

cdc45-3A and *cdc45-3D* Mutations Reduce Collapsing Rate of Stalled Replication Forks.

To establish how the *cdc45-3A* or *cdc45-3D* mutations influence the stabilization of stalling forks, we used dual-color DNA combing to assess the stability of DNA replication forks upon 5 or 12.5 mM HU treatment (Fig. 3 A, Left). Examples of fibers with asymmetric, stalled, or collapsed and newly started forks are shown (Fig. 3 A, Middle). The percentage of permanently arrested forks (IdU fibers without CldU tracks at one or both ends) increased in *cdc45-3A* and *cdc45-3D* cells when compared to *cdc45⁺* (5 mM HU: 14.5, 14, and 2.5%; 12.5 mM HU: 36, 35, and 9%, respectively). As expected, in *cds1^{Chk2}Δ* (checkpoint-deficient) cells, the percentage of arrested forks unable to resume replication was higher, at ~31 and 65% with 5 or 12.5 mM HU treatment, respectively (Fig. 3 A, Right). Under 5 or 12.5 mM HU treatment for 4 h, both *cdc45-3A* and *cdc45-3D* mutations reduced the frequency of permanently arrested forks in the *cds1^{Chk2}Δ* background by ~34 ([31 – 21]/31) or 25% ([65 – 49]/65) (Fig. 3 A, Right), indicating that even under a short time of HU treatment, a considerable number of replication forks collapsed in *cds1^{Chk2}Δ* cells, but the *cdc45-3A* and *cdc45-3D* mutations significantly reduced the rate of fork collapsing under Cds1^{Chk2} deficiency. In addition, the *cdc45-3A* and *cdc45-3D* cells had a higher rate of fork collapsing than *cdc45⁺* cells, indicating that *cdc45-3A* and *cdc45-3D* mutations do not fully mimic

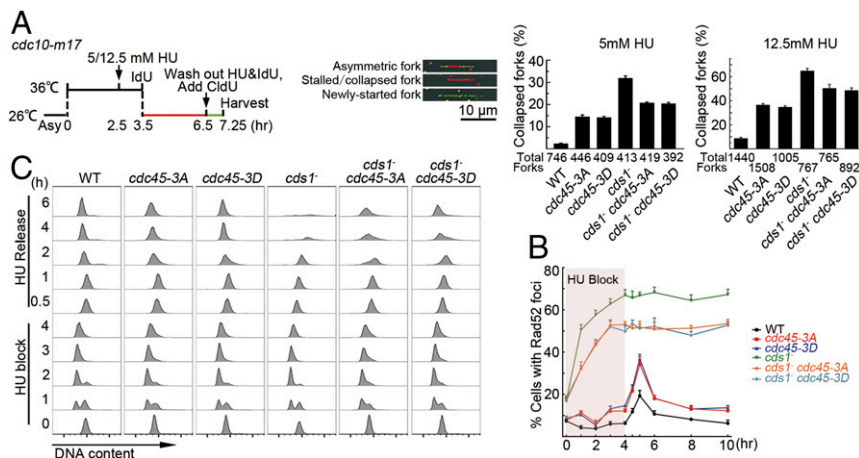


Fig. 3. Cdc45-3A/3D mutations stabilize stalled replication forks. (A) Dual labeling combing to assess the rate of stalled fork collapse. (Left) Experimental scheme. (Middle) Examples of fiber staining. (Right) Percentage of collapsed forks in the indicated genetic backgrounds calculated as the number of IdU-only DNA tracts divided by the total tracts. (B) Time course of the percentage of cells showing Rad52 foci during HU treatment (12.5 mM) for 4 h (pink box) and subsequent release. (C) FACS analysis of cells from B. The data in A and B are presented as the mean ± SD.

$Cds1^{Chk2}$ regulation to Cdc45 (see Fig. 5H, the level of helicase activity).

Fluorescence-activated cell sorting (FACS) analyses were performed to those cells that were analyzed in Fig. 3A. As shown in *SI Appendix, Fig. S2A*, these cells were arrested in the G1 phase after 3.5 h at 36 °C. After the cultures were shifted to a permissive temperature of 26 °C with or without HU, the wt and $cds1^{Chk2} \Delta$ cells progressed nicely to the G2 phase with a DNA content of 2C in the absence of HU (the peaks are slightly skewed to the right 3 h after cell release compared to the 2C peaks before arrest [“0” h] because the cells became larger during arrest) (*SI Appendix, Fig. S2 A, Top and Bottom*). However, a similar level of DNA content of ~1C (the peaks are slightly skewed to the right due to a larger cell size) was detected among wt, $cds1^{Chk2} \Delta$, $cdc45-3A$, $cdc45-3D$, $cdc45-R319I$, $cds1^{Chk2} \Delta cdc45-3A$, $cds1^{Chk2} \Delta cdc45-3D$, and $cds1^{Chk2} \Delta cdc45-R319I$ cells (*SI Appendix, Fig. S2 A, Top and Bottom*). These results suggest that the forks without CldU incorporation represent stalled or collapsed forks rather than resulting from the merging of two neighboring forks.

Fork collapse results in DNA lesions that recruit the Rad52 recombination protein to repair the broken forks. During repair, Rad52 forms a focus. Thus, we monitored Rad52 foci as an indirect readout of fork collapse in cells treated with HU and then released into fresh media (*SI Appendix, Fig. S2B*). Under unperturbed growth condition (–HU) ~8% of wt cells and ~18% of $cds1^{Chk2} \Delta$ cells showed Rad52 foci (Fig. 3B). $cdc45-3A$ and $cdc45-3D$ were similar to wt, whereas the $cds1^{Chk2} \Delta cdc45-3A$ and $cds1^{Chk2} \Delta cdc45-3D$ double mutant strains were similar to the $cds1^{Chk2} \Delta$ single mutant (Fig. 3B). As previously reported (37), during 4 h of HU treatment, the percentage of wt cells exhibiting Rad52 foci did not increase, whereas the percentage of the Rad52 foci positive $cds1^{Chk2} \Delta$ mutant increased to ~67%. Consistent with the $cdc45-3A$ and $cdc45-3D$ mutations partially suppressing the $cds1^{Chk2} \Delta$ mutant sensitivity to HU, the $cds1^{Chk2} \Delta cdc45-3A$ and $cds1^{Chk2} \Delta cdc45-3D$ double mutants displayed a less dramatic increase in Rad52 foci positive cells at ~50%. The single $cdc45-3A$ and $cdc45-3D$ mutants showed only a modest increase (~14%). Upon release from HU, wt and the single $cdc45-3A$ and $cdc45-3D$ mutants showed a transient pulse of Rad52 foci positive cells, whereas the percentage of Rad52 foci positive cells observed after the release from HU arrest did not change in $cds1^{Chk2} \Delta$ or the $cds1^{Chk2} \Delta cdc45-3A$ and $cds1^{Chk2} \Delta cdc45-3D$ double mutant cultures (Fig. 3B). Flow cytometry assays showed that the $cdc45-3A$ and $cdc45-3D$ mutations increased the ability of $cds1^{Chk2} \Delta$ cells to re-enter the cell cycle after 4 h of HU treatment (Fig. 3C). These results are consistent with our interpretation that the $cdc45-3A$ and $cdc45-3D$ mutations reduce the rate of collapse of stalling forks in $cds1^{Chk2} \Delta$ cells.

Checkpoint Regulation and the $cdc45-3A$ and $cdc45-3D$ Alleles Decrease Fork Speed. $cdc45-3A$ and $cdc45-3D$ cultures grow slightly more slowly than $cdc45^+$ (Fig. 4A). Pulse labeling of asynchronous cultures with EdU demonstrated that the number of S phase cells in $cdc45-3A$ and $cdc45-3D$ increased by ~65% compared to $cdc45^+$ cultures (Fig. 4B). Consistent with an extended S phase, the average amount of EdU incorporated per cell in $cdc45-3A$ and $cdc45-3D$ was decreased by ~30% (Fig. 4C). This suggests that the reduced growth rate results from an extended S phase, probably because of slower replication fork progression. To address this, DNA combing was used to examine fork speed in $cdc45^+$ and appropriate mutant cells during the unperturbed S phase, during HU treatment, and after release from HU arrest (schematic, Fig. 4D). *SI Appendix, Fig. S3 A and B* show examples of the labeled DNA fibers and the distributions of their lengths. During the unperturbed S phase (10 min EdU incorporation), the median tract length for $cdc45^+$ cells was 4.1 μ m (Fig. 4E, *Left*), translating to ~14~21 nucleotides per second (nt/s), based on ~2 to 3 kb/per μ m of DNA fiber (49). $cdc45-3A$ and $cdc45-3D$ cells displayed a

reduced fork speed (3.02 and 3.03 μ m, respectively: 10~15 nt/s). This indicates that $cdc45-3A$ and $cdc45-3D$ reduce the activity/processivity of the CMG helicase. Fork speed was also decreased in the $cds1^{Chk2} \Delta$, $cds1^{Chk2} \Delta cdc45-3A$, and $cds1^{Chk2} \Delta cdc45-3D$ cells. While the explanation for a slower fork speed in the $cds1^{Chk2} \Delta$ cells is unknown, it is possible that a basal level of checkpoint regulation is required to promote replication fork movement through difficult-to-replicate loci (5).

To examine fork speed in the presence of HU, the cultures were first incubated with HU for 1 h. Then, EdU was added, and the cultures incubated for an additional 3 h. As shown in Fig. 4E, *Middle*, fork speed in $cdc45^+$ cells dramatically decreased; a tract length of 5 μ m equates to a speed of 1 nt/s. A less dramatic decrease (~2 nt/s) was observed for $cds1^{Chk2} \Delta$ cells. $cdc45-3A$ and $cdc45-3D$ single mutant cells displayed a fork speed appreciably above $cdc45^+$ cells but substantially below that of $cds1^{Chk2} \Delta$ cells. Importantly, the double mutant $cds1^{Chk2} \Delta cdc45-3A$ and the $cds1^{Chk2} \Delta cdc45-3D$ cells exhibited a fork speed that was appreciably below $cds1^{Chk2} \Delta$ cells but still substantially above that of $cdc45^+$ cells. These data are again consistent with $cdc45-3A$ and $cdc45-3D$ reducing the activity/processivity of the CMG helicase, therefore partially mimicking the regulation of Cdc45 by $Cds1^{Chk2}$. Following the release from HU arrest, replication forks quickly recovered in $cdc45^+$ cells. Forks recovered to a much lesser extent in $cds1^{Chk2} \Delta$ cells. As expected, $cdc45-3A$ and $cdc45-3D$ mutants recovered less well than $cdc45^+$ cells but better than $cds1^{Chk2} \Delta$ cells. Importantly, $cds1^{Chk2} \Delta cdc45-3A$ and $cds1^{Chk2} \Delta cdc45-3D$ double mutant cells recovered appreciably better than $cds1^{Chk2} \Delta$ cells, although substantially less well than $cdc45^+$ cells (Fig. 4E, *Right*). Thus, the recovery of replication following HU treatment and the extent of changes in replication fork speed correlate with HU sensitivity, suggesting fork progression is a critical target of the intra-S phase checkpoint regulation in response to replication fork stalling.

Changes in fork speed have been correlated with changes in origin firing (50). To establish if wt, $cds1^{Chk2} \Delta$, $cdc45-3A$, $cdc45-3D$, $cdc45-R319I$, $cds1^{Chk2} \Delta cdc45-3A$, $cds1^{Chk2} \Delta cdc45-3D$, and $cds1^{Chk2} \Delta cdc45-R319I$ cultures display similar efficiencies of DNA replication initiation, the “start” cell cycle control gene, $cdc10^+$, was replaced by the temperature-sensitive $cdc10-m17$ allele. Cells were arrested at the G1 phase and subsequently released into a synchronous S phase in the presence of BrdU for 40 min (–HU) or 3 h (+HU). DNA combing was then performed and interorigin distances estimated from BrdU staining (51). As shown in Fig. 4F, *Bottom*, no matter whether HU was present or absent, the average distance between two neighboring initiation sites among these cells was equivalent. Thus, the efficiency of replication initiation is not affected by the $cds1^{Chk2} \Delta$, $cdc45-3A$, $cdc45-3D$, $cdc45-R319I$, or the cells bearing $cds1^{Chk2} \Delta$ combined with the $cdc45$ mutant. These results are consistent with that the fission yeast *S. pombe* cells have only eight late origins (1 to 2% of total origins) (52) and that only a few origins in *S. pombe* are affected by checkpoint regulation in response to replication stress (53).

$Cds1^{Chk2}$ -Mediated Cdc45 Phosphorylation Inhibits CMG Helicase Activity. To establish if $Cds1^{Chk2}$ -mediated Cdc45 phosphorylation reduces CMG helicase activity, we assayed the helicase activity of CMG isolated from the chromatin of unperturbed wt and mutant cells and from cells experiencing DNA replication stress. Under unperturbed growth conditions, the helicase activities of CMG-3A and CMG-3D were reduced by ~50%. A second potential phosphor-mimic mutant, $cdc45-3E$, grew very poorly (*SI Appendix, Fig. S4 A–C*) and displayed an 85% reduction in helicase activity (Fig. 5A and B). Under replication stress (the presence of HU), the helicase activity of CMG from wt cells was reduced by ~70% but remained unchanged when isolated from $cds1^{Chk2} \Delta$ cells (Fig. 5C and D). In support of this result, the helicase activity of the CMG complex isolated from HU-treated cells was examined in the presence or absence of λ PPase. As shown in Fig. 5E and F,

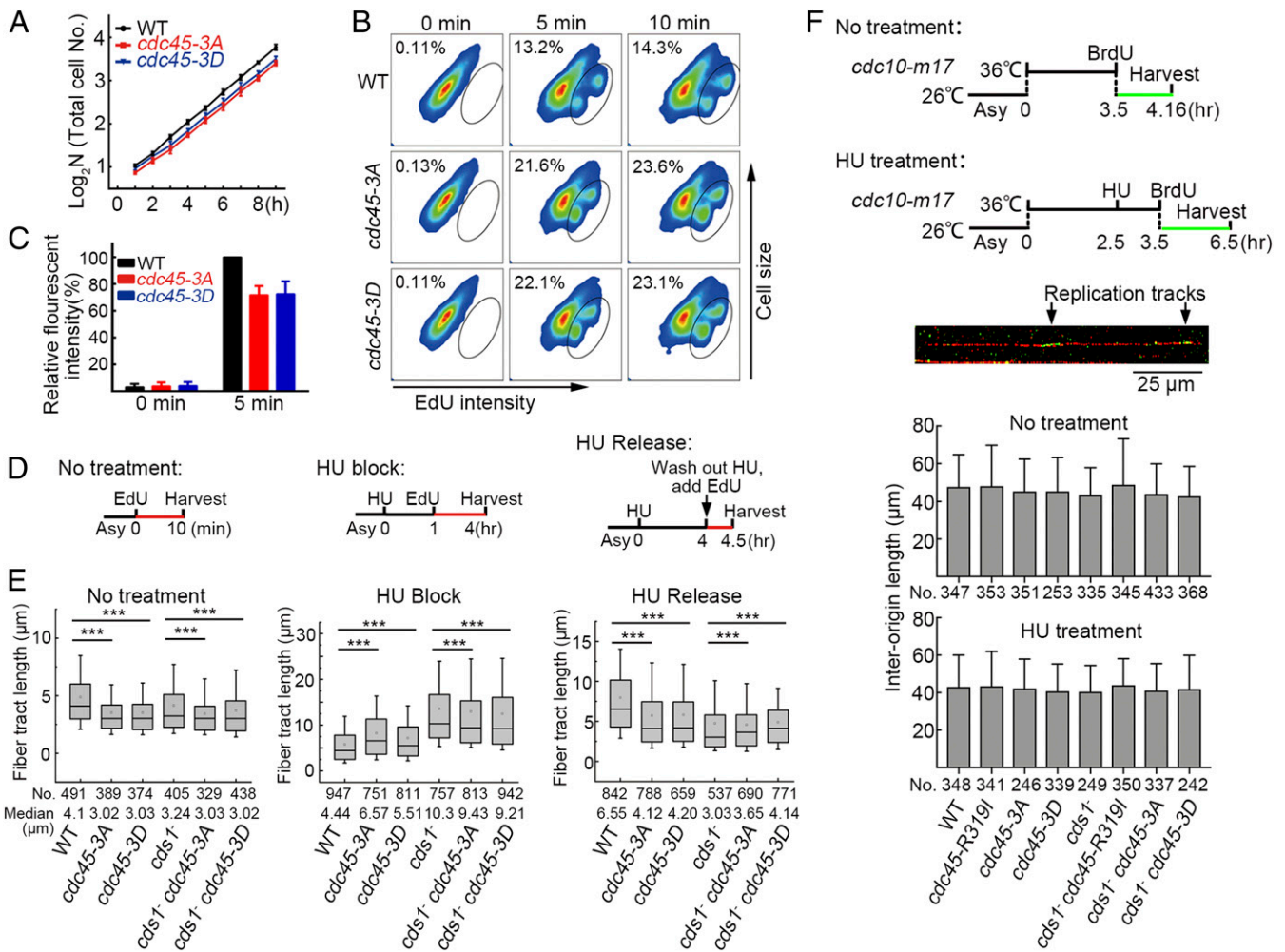


Fig. 4. Fork speed correlates with HU survival. (A) Growth curves of the indicated strains. (B) The percentage of cells in S phase measured after 5 and 10 min EdU pulses by FACS in the indicated strains. EdU-labeled (S phase) cells are encircled. (C) The relative fluorescent intensity of EdU-stained cells. (D) Experimental scheme for measuring fork speed by DNA combing. (E) The median length of EdU-labeled DNA fibers in the WT and indicated mutant strains under normal cell growth conditions in the presence of HU and after release from HU treatment. The numbers of measured DNA fibers are indicated. Representative images are shown in *SI Appendix, Fig. S3A*. (F) The measurement of interorigin distance in wt, *cds1*⁻, *cdc45-3A*, *cdc45-3D*, *cdc45-R319I*, *cds1*⁻ *cdc45-3A*, *cds1*⁻ *cdc45-3D*, and *cds1*⁻ *cdc45-R319I* cells in the presence or absence of HU. (Top) Experimental scheme. (Bottom) Average interorigin distance in the indicated strains. The data in A, C, and F are presented as the mean value \pm SD. The data in E is presented as a box plot with an interquartile range of 25 to 75%, the median value (line), mean value (small hollow square), limited inferior <5%, and limited superior >95%. The Mann-Whitney *U* test was applied for statistical analysis. ****P* < 0.001.

the CMG helicase activity was dramatically reduced in the HU-treated cells, but the helicase activity was nearly recovered to the level of normal CMG helicase activity. This result indicates that the CMG activity is greatly reduced in stalling replisomes, and this reduction appears solely dependent upon *Cds1*^{Chk2} checkpoint regulation. Consistent with *Cds1*^{Chk2} activity inhibiting CMG via phosphorylation of Cdc45 S275, S322, and S397, when CMG and CMG-3A were isolated in parallel from cells either treated, or not, with HU (Fig. 5 *G* and *H*), the helicase activity was reduced by ~75% for CMG, whereas the activity of CMG-3A, which is already compromised, was not significantly further reduced. This is consistent with Cdc45-3A being largely insensitive to *Cds1*^{Chk2} regulation (Fig. 1C) and suggests that there are no further major modifications on Cdc45 or other subunits of CMG complex by the intra-S phase checkpoint that reduce CMG helicase activity.

When CMG or CMG-3A were treated *in vitro* by *Cds1*^{Chk2}, Cdc45, but not Cdc45-3A, was phosphorylated (Fig. 5 *I*, Top). The helicase activity of CMG was reduced by ~50%, whereas there was no significant change in the activity of CMG-3A. The difference in the reduction of helicase activity (~75% *in vivo* versus ~50%

in vitro) is probably because the level of *Cds1*-mediated phosphorylation of Cdc45 in the *in vitro* reaction did not reach to a maximum extent that is reached *in vivo*. No detectable changes in the level of Cdc45, formation of CMG complex, or affinity of CMG to chromatin DNA were observed between *cdc45*⁺ and *cdc45-3A* cells, either with or without HU treatment (*SI Appendix, Fig. S5 A-C*), which is consistent with that the *cdc45-3A* or *cdc45-3D* cells grew relatively normally in the absence of exogenous replication stress. However, the S phase (~10% of the cell division cycle) of these mutant cells is lengthened by ~65% (Fig. 4 *A* and *B*). And this suggests that the conformation of the mutant CMG complex should be changed in some points, which results in a reduced helicase activity (Fig. 5 *A* and *B*). Taken together, our data suggests that *Cds1*^{Chk2}-mediated Cdc45 phosphorylation on S275, S322, and S397 reduces CMG helicase activity and slows fork speed in response of replication fork stalling.

***Cds1*^{Chk2} Prevents DNA Polymerases and CMG Helicase in Stalling Forks from Separation.** The *Cds1*^{Chk2}-mediated inhibition of CMG helicase activity at stalling replication forks implies that a key mechanism in

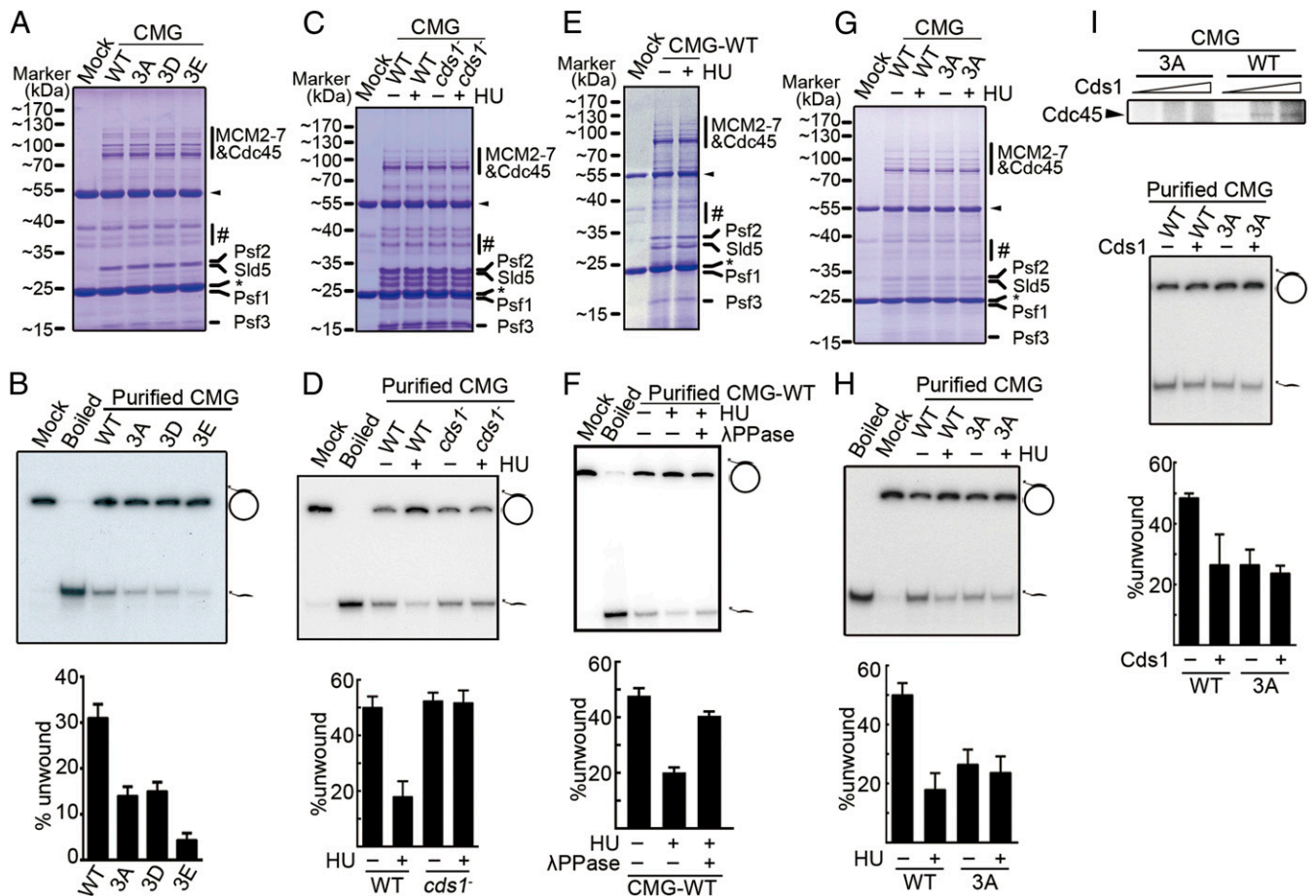


Fig. 5. $Cds1^{Chk2}$ -mediated phosphorylation of Cdc45 regulates CMG helicase activity. (A) Coomassie-stained CMG complexes purified from $cdc45^+$, $cdc45-3A$, $cdc45-3D$, and $cdc45-3E$ (CMG, CMG-3A, CMG-3D, and CMG-3E, respectively) growing without HU treatment. Heavy and light antibody chains are indicated by a black arrow and star, respectively. #; contaminants from HA antibody-conjugated resins. (B, Top) Autoradiograph reporting the helicase activity of CMG and CMG-3A/3D/3E. The position of the substrate (^{32}P -labeled oligonucleotide annealed to a M13 circular ssDNA) and displaced oligonucleotide were indicated. (Bottom) Quantification of three independent experiments. (C, E, and G) Indicated CMG complexes purified from untreated or HU-treated cells as in A. (D, F, and H) Autoradiograph and quantification of the helicase activity of purified CMG complexes shown in C, E, and G (see B for details). The presence of λ PPase is indicated in F. (I, Top) $Cds1$ -dependent phosphorylation of the Cdc45 subunit of the indicated CMG complexes in vitro. (Bottom) Autoradiograph and quantification of the helicase activity of the indicated CMG complexes with and without in vitro $Cds1$ treatment (see B for details). The data in B, D, F, H, and I are presented as the mean \pm SD.

checkpoint regulation to preserve replisome integrity is to maintain the link between strand separation and DNA polymerization. Thus, we examined whether two core components, DNA polymerase α and the CMG helicase, are separated when replication forks are stalling. The levels of DNA polymerase α (Spb70 subunit) and MCM (Mcm7 subunit) were assessed in normal and stalling forks in wt and $cds1^{Chk2}\Delta$ cells. First, we observed that the levels of Spb70 (Pol- α subunit) and Mcm7 (CMG subunit) on chromatin were similar in the presence or absence of HU for wt, $cds1^{Chk2}\Delta$, $cds1^{Chk2}\Delta cdc45-3A$, and $cds1^{Chk2}\Delta cdc45-3D$ cells (Fig. 6A). When replication forks were isolated from the chromatin by immunoprecipitation against DNA Pol- α (Spb70 subunit), the amount of the two subunits was equivalent between the four cultures when replication forks were not stalled (Fig. 6B). However, when the replication forks were isolated from cells treated with HU, the levels of Spb70 remained comparable between the cultures, but the level of Mcm7 decreased by $\sim 27\%$, $\sim 15\%$, and $\sim 16\%$, respectively, in $cds1^{Chk2}\Delta$, $cds1^{Chk2}\Delta cdc45-3A$, and $cds1^{Chk2}\Delta cdc45-3D$ cells (Fig. 6B). A physical separation of replicative helicase and DNA polymerases collapses forks because a normal/functional replication fork requires both replicative helicase and DNA polymerases present in forks, and their actions must be strictly coupled. The above data indicate that $\sim 1/6$ and 7 to 1/4 of

forks collapsed after 3 h of HU treatment in the $cds1^{Chk2}\Delta cdc45-3A$, $cds1^{Chk2}\Delta cdc45-3D$, and $cds1^{Chk2}\Delta$ cells. These results suggest that the inhibition of CMG helicase activity is required to prevent the separation of the CMG replicative helicase and DNA polymerase α .

The separation of CMG and DNA polymerases in the $cds1^{Chk2}\Delta$ cells was further examined in response to replication fork stalling due to MMS treatment (54), with replication forks isolated by immunoprecipitation against either DNA Pol- α , Pol- δ , or Pol- ϵ . The results are shown in Fig. 6C–K. The total amount of Pol- α (Spb70), Pol- δ (Cdc1 and Pol3), Pol- ϵ (Pol2), PCNA, Mcm7, or Cdc45 on chromatin was equivalent between wt and $cds1^{Chk2}\Delta$ cells (Fig. 6C–D, F, G, I, and J). In the isolated forks (immunoprecipitated with polymerase subunits), the amount of Spb70, Cdc1, Pol3, Pol2, or PCNA remained equal, and the amounts of Mcm7 and Cdc45 also remained equal in wt cells (Fig. 6C, F, and I), but the amounts of Mcm7 and Cdc45 were reduced by ~ 30 to 40% in the $cds1^{Chk2}\Delta$ cells, again consistent with the separation of helicase and DNA polymerases in stalling forks (Fig. 6D, G, and J). As in the case of HU stalling replication forks, the $cdc45-3A$ and $cdc45-3D$ mutants, which partially mimic the checkpoint regulation on Cdc45, reduced the level of the CMG helicase and DNA polymerases separation (Fig. 6E, H, and

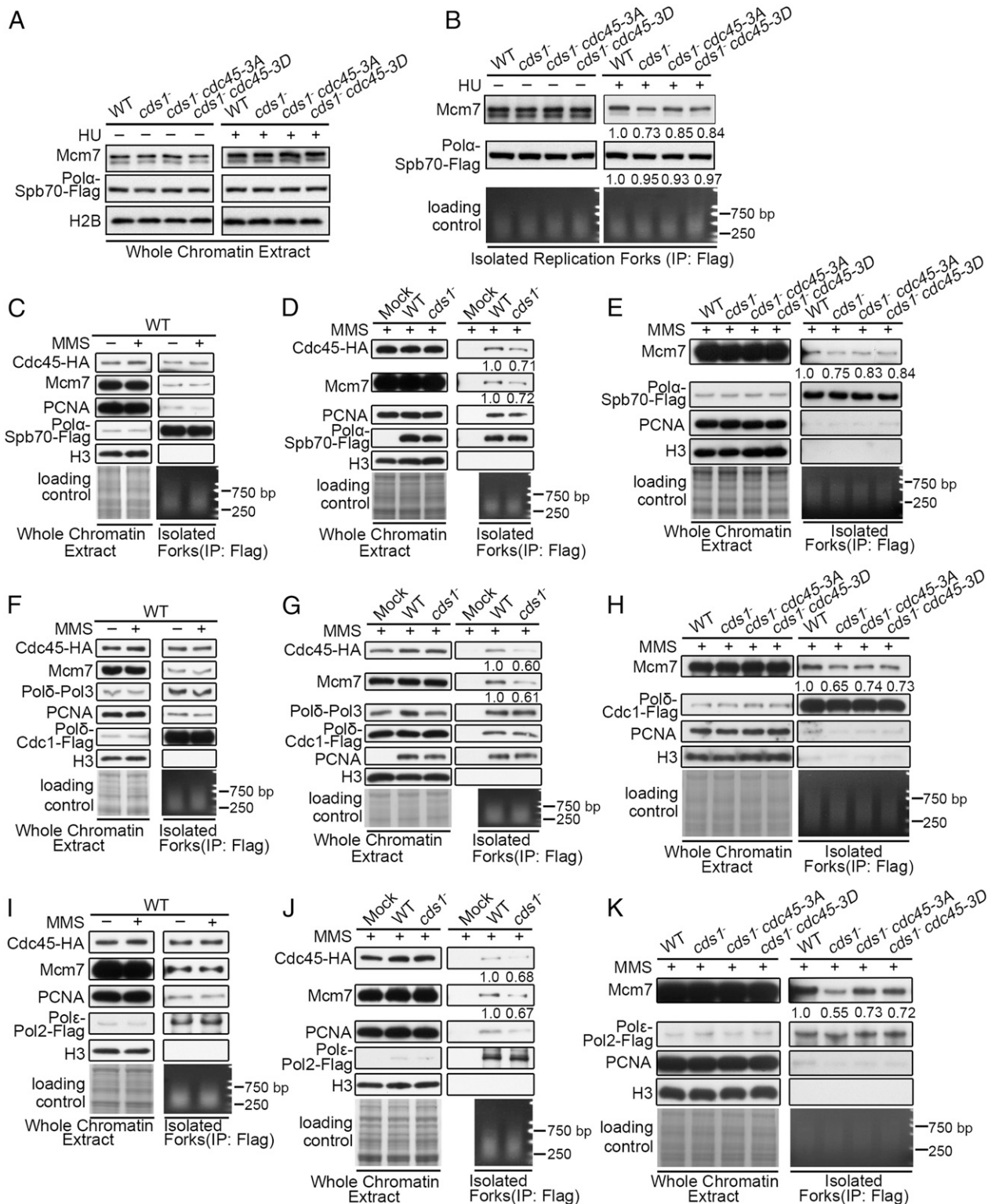


Fig. 6. *Cds1^{Chk2}* prevents separation of DNA polymerases and CMG helicase in stalling replication forks. (A) The levels of Pol- α -Spb70 and Mcm7 on chromatin from the indicated strains in presence or absence of HU. (B) The amount of Mcm7 in stalling forks isolated from the indicated strains. (C–K) The levels of Mcm7, Cdc45, PCNA, Pol- α -Spb70 (C–E), Pol- δ -Cdc1 (F–H), and Pol- ϵ -Pol2 (I–K) on chromatin isolated from unperturbed or MMS-treated (forks stalling) cells of wt, *cds1⁻*, *cds1⁻ cdc45-3A*, and *cds1⁻ cdc45-3D*. Loading controls, Coomassie blue-stained gel, and isolated fork DNA are shown at the bottom of the panels.

K). In addition, the physical separation of the replicative helicase CMG and DNA polymerases helicase is supposed to increase the amount of single-stranded DNA (ssDNA) on chromatin and potentially cause more DNA breaks. By measuring RPA foci, the

cellular level of broken DNA, and the sensitivity of chromatin DNA to the ssDNA-specific endonuclease P1 in the HU-treated wt, *cds1^{Chk2} Δ*, *cds1^{Chk2} Δ cdc45-3A*, and *cds1^{Chk2} Δ cdc45-3D* cells, the results show that RPA foci significantly increased in

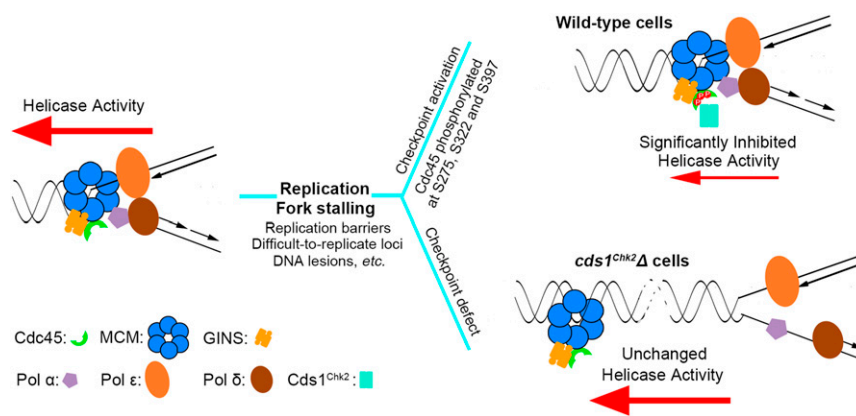


Fig. 7. Cds1^{Chk2} phosphorylates Cdc45 to inhibit CMG helicase activity, reduce fork speed, and prevent physical separation of the CMG helicase and DNA polymerases in stalling/stalled replication forks. This results in the stabilization of stalling replication forks. (Left) As replication forks traverse DNA, the polymerases act coordinately with the CMG helicase to synthesize the leading and lagging strands. When replication polymerases are impeded, a small amount of ssDNA is revealed, which activates the intra-S phase ATR-dependent checkpoint pathway (Top, Right). Activated Cds1^{Chk2} phosphorylates the Cdc45 subunit of the CMG replicative helicase complex on S275, S322, and S397. This results in a severe reduction of CMG helicase activity and prevents the physical uncoupling of strand separation and DNA polymerization, thus stabilizing the stalling replication fork. (Bottom, Right) In checkpoint-deficient cells, CMG helicase activity is not inhibited, which results in DNA unwinding proceeding faster than DNA polymerization, the exposure of excessive ssDNA, and fork collapse.

c-ds1^{Chk2}Δ cells, but Cdc45-3A and Cdc45-3D mutations noticeably reduced RPA foci in the *c-ds1^{Chk2}Δ* cells (SI Appendix, Fig. S6 A, B, and D). Consistent with more RPA foci, an ~28% increase in the level of broken DNA was detected in the *c-ds1^{Chk2}Δ* cells compared with wt, no matter whether chromatin DNA was treated or untreated by nuclease P1 (SI Appendix, Fig. S6C), suggesting that chromatin DNA in the *c-ds1^{Chk2}Δ* cells is more vulnerable, probably due to more ssDNA. Taken together, these results indicate that the CMG helicase moves away from DNA polymerases or replication forks when replication forks are stalling in the absence of the intra-S phase checkpoint kinase Cds1^{Chk2} and that a reduction of CMG helicase activity (in the *c-ds1^{Chk2}Δ* mutants) alleviates this separation.

Discussion

The catalytic centers in replicative DNA polymerases have an ultra-fine structure (55). These centers can accommodate normal A:T and G:C pairs but do not accommodate either A:C or G:T pairs, even though the three-dimensional difference between the A:T and A:C or the G:C and G:T pair is small. An ultra-fine catalytic center greatly increases the fidelity of DNA synthesis, with an error rate of ~10⁻⁴ per base incorporated (56). However, it also means that DNA polymerases can be easily blocked, causing fork stalling. Base modifications, such as pyrimidine dimerization, base loss, base alkylation, and base oxidation, can block DNA polymerases (57, 58). A human cell has about 10⁵ DNA lesions per day, the majority of which are able to block DNA polymerases (59, 60). Furthermore, genomic DNA harbors a great number of sites capable of forming DNA secondary structures which can act as native replication barriers. For example, human genomic DNA contains ~700,000 potential G4 structures, plus a great number of potential hairpin and triplex forming sequences (61–63). Secondary DNA structures can form particularly on the DNA template for lagging strand synthesis because a single-stranded region of ~150 nt or more is available on the lagging strand template due to the time gap between the unwinding of the double-stranded DNA (dsDNA) template and the subsequent initiation of lagging strand DNA synthesis. Thus, the stalling of DNA polymerases is likely a frequent event during DNA replication.

In this study, through a large scale of genetic screening, Cdc45, a subunit of the CMG complex, was identified as a potential checkpoint target: the *c-ds1^{Chk2}Δ* mutated cells had an almost normal cell growth cycle when compared to *c-ds1^{Chk2}Δ* cells, but the

mutation could dramatically reduce the sensitivity of *c-ds1^{Chk2}Δ* cells to 2.5 mM HU (Fig. 1A). Subsequent work demonstrated that, under conditions of fork stalling, Cds1^{Chk2} concurrently phosphorylated Cdc45 on S275, S322, and S397 (Fig. 1 B–G). The phospho-mimic and nonphosphorylation mutations, *c-ds1^{Chk2}Δ* and *c-ds1^{Chk2}Δ*, respectively, both dramatically reduced the sensitivity of *c-ds1^{Chk2}Δ* cells to 2.5 mM HU (~600- to 3,000-fold; Fig. 2C) and to 1.0 mM stavudine (~25-fold; Fig. 2E). The profound reduction in the sensitivity of *c-ds1^{Chk2}Δ* cells to HU or stavudine strongly suggests that Cdc45/CMG is a crucial checkpoint target in the checkpoint-mediated stabilization of stalled forks. The response to stavudine, the incorporation of which completely blocks replication, strongly indicates that *c-ds1^{Chk2}Δ* and *c-ds1^{Chk2}Δ* mutations increase the stability of stalled forks rather than simply reducing the rate of dNTPs consumption during HU-induced fork stalling. Consistent with this, we show that interorigin distances are unchanged in the early S phase in *c-ds1^{Chk2}Δ* and *c-ds1^{Chk2}Δ* cultures when compared to *c-ds1^{Chk2}Δ*, demonstrating there is no systemic initiation defect in these cells.

We next demonstrated that Cds1^{Chk2}-mediated phosphorylation of Cdc45 on S275, S322, and S397 dramatically reduced CMG helicase activity (Fig. 5 D and H) and verified that Cds1^{Chk2} checkpoint kinase-dependent slowing of replication in the presence of HU occurred in *S. pombe* (Fig. 4 E, Middle). The stress-induced slowing of replication fork progression is consistent with the slowing of replication under replication stress that was observed by Paulovich and Hartwell (36). The rate of replication elongation, or fork speed, depends largely on replicative helicase activity and on the degree of chromatin compaction: replication fork speed in eukaryotic cells is ~20 nt/s, but the nt incorporation rates by DNA polymerase δ and ε on naked DNA are ~400 nt/s (64). In a previous study, we showed that the chromatin surrounding stalled replication forks becomes more compact and that this compacted chromatin contributes to preventing the CMG helicase from moving away from DNA polymerases/replication forks (Chromfork control) (44). This current work answers the following question: Does the intra-S phase checkpoint target any other factors to slow down DNA replication under replication stress? While a number of replication components have been reported to be phosphorylated by the checkpoint in response to fork stalling or DNA damage, including RPA, Dna2 (17, 65–67), and MCM subunits (68–70), none of these modifications have been linked directly to the regulation of replication fork speed.

Our identification of the regulation of the CMG helicase by Cdc45 phosphorylation demonstrates that cells use at least two mechanisms to prevent the CMG helicase from delinking from DNA polymerases. One is Chromsfork-mediated chromatin compaction around stalled forks and the other is the checkpoint-mediated reduction of CMG helicase activity.

Unusually for the analysis of phosphorylation site mutants, both the *cdc45-3A* and *cdc45-3D* alleles mimic the checkpoint regulation by stabilizing stalling replication forks. A simple explanation for this is that Cds1^{Chk2}-mediated phosphorylation of Cdc45 on S275, S322, and S397, and *cdc45-3D* and *cdc45-3A* mutations all reduce the CMG helicase activity (Fig. 5). Why does the *cdc45-3A* phosphomimic mutation, like the *cdc45-3D* phospho-mimic, reduce the CMG helicase activity? We propose that the reason pertains to a unique biochemical property of CMG helicase. CMG is composed of Cdc45, MCM, and GINS. MCM is the catalytic component, but it alone does not have detectable ATPase and helicase activity (71). Once associated with Cdc45 and GINS, MCM, or actually CMG, becomes a robust ATPase and helicase (72–76). This demonstrates that robust CMG helicase activity requires proper interactions among Cdc45, MCM, and GINS. Based on recently resolved structures of human Cdc45 and GINS and *Saccharomyces cerevisiae* CMG at a resolution of 3.7 to 4.8 Å (77–79), the S275, S322, and S397 residues of Cdc45 lie in the CID domain that is responsible for interacting with Mcm2, Mcm5, Psf1, and Psf2. In support of a core role for these serine residues in coordinating Cdc45-CMG interactions, the mutants *cdc45-3C*, *cdc45-3K*, *cdc45-3V*, *cdc45-3N*, and *cdc45-3G* also caused otherwise wt cells to become sensitive to HU and significantly rescued the HU sensitivity of *cds1^{Chk2}Δ* cells (SI Appendix, Fig. S1 H and I). Importantly, it has also been observed in other proteins that the phenotype of the phosphor-mimic mutation (serine or threonine to aspartate) is recaptured by the mutation of serine or threonine to alanine (80, 81).

The importance of the S275, S322, and S397 residues for CMG helicase activity is further evidenced by our observation that a *cdc45-3T* allele (three serines changed to threonines) caused slightly increased sensitivity to 5.0 or 7.5 mM HU when compared to *cdc45⁺* and moderately rescued the sensitivity of the *cds1^{Chk2}Δ* cells to 1.5 mM HU (SI Appendix, Figs. S1 H and I and S5D). This implies that the S275, S322, and S397 residues are critical for Cdc45 to interact properly with MCM and GINS for a normal level of CMG helicase activity. This could explain why the phosphorylation of these three sites, or mutation to aspartic acid (*cdc45-3D*), alanine (*cdc45-3A*), and glutamic acid (*cdc45-3E*), all significantly reduced the helicase activity of CMG (Fig. 5 B and D) and rescued the sensitivity of *cds1^{Chk2}Δ* cells to HU (Fig. 2 C and D; note, we were unable to examine *cdc45-3E* in serial dilution assays because the *cdc45-3E* cells grew very poorly [SI Appendix, Figs. S1 H and I and S4 A–C]). Taken together, our data indicate that the intra-S phase checkpoint regulates Cdc45 to decrease the helicase activity of CMG in response to fork stalling. The S275, S322, and S397 sites within the CID domain of Cdc45 provide an ideal mechanism to regulate CMG helicase activity because even a very minor change in these three serines, such as a mutation to threonine, affects the CMG helicase activity.

Why does the stabilization of stalled replication forks require the reduction of replicative helicase activity? The answer to this question should account for the fundamental biochemical

reactions occurring at forks. Normally, the helicase-mediated unwinding of template dsDNA must be strictly coordinated with the synthesis of the leading and lagging strands. This coordination is achieved through protein interactions between the replicative helicase and DNA polymerases. In *Escherichia coli*, this is mediated by the τ subunit of the replisome and in eukaryotes by the Ctf4 and Mrc1 proteins (26, 82, 83). However, when (as stated above) DNA polymerases are blocked during DNA replication, the replicative helicase, powered by ATP hydrolysis, can move forward inappropriately and delink from the DNA polymerases. This would, in effect, collapse forks: once CMG moves away from a fork without being closely followed by DNA polymerization, the two unwound, and complementary DNA strands behind the helicase are vulnerable to nucleases and also can potentially reanneal, allowing nucleosomes to reassemble on the reannealed dsDNA, making the separation of replicative helicase and DNA polymerases irreversible (Fig. 7). Thus, when DNA polymerases are blocked, the intra-S phase checkpoint Cds1^{Chk2} kinase concurrently phosphorylates Cdc45 on the S275, S322, and S397 sites and dramatically reduces the CMG helicase activity. This, along with other responses such as Chromsfork, prevents the CMG helicase delinking from DNA polymerases and contributes to preserving the integrity of replisomes.

Replication stress not only affects genomic stability but also threatens cell survival. Thus, several distinct but closely related mechanisms of the intra-S phase checkpoint regulation have been developed to cope with replication stress. Besides its critical role in stabilizing stalling replication forks, the intra-S phase checkpoint also regulates to block mitosis (84, 85) in order to avoid catastrophe caused by cell division when DNA replication is incomplete. Furthermore, it has been well documented that the intra-S phase checkpoint regulates to inhibit late origin firing when cells suffer replication stress (31–34, 86). The inhibition of late origin firing reduces the number of replication forks encountering stress, thus alleviating the replication stress of cells; preventing the exhaustion of RPA (and possibly some other replication factors as well) is another reason for checkpoint regulation to inhibit late origin firing, thus prohibiting replication catastrophe (34).

Materials and Methods

The information of strains, plasmids, antibody generation or purchased antibodies, the preparation of cell or chromatin extracts, DNA combing assay, the isolation of replication forks, the isolation of the replicative helicase CMG complex from chromatin, expression and purification of *S. pombe* Cds1 and Cdc45, EdU incorporation and detection, and assay of in vitro CMG helicase activity are in SI Appendix.

Data Availability. All study data are included in the article and/or SI Appendix.

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