

A mechanism for Rad53 to couple leading- and lagging-strand DNA synthesis under replication stress in budding yeast

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In response to DNA replication stress, DNA replication checkpoint kinase Mec1 phosphorylates Mrc1, which in turn activates Rad53 to prevent the generation of deleterious single-stranded DNA, a process that remains poorly understood. We previously reported that laggingstrand DNA synthesis proceeds farther than leading strand in rad53-1 mutant cells defective in replication checkpoint under replication stress, resulting in the exposure of long stretches of the leadingstrand templates. Here, we show that asymmetric DNA synthesis is also observed in mec1-100 and mrc1-AQ cells defective in replication checkpoint but, surprisingly, not in mrc11 cells in which both DNA replication and checkpoint functions of Mrc1 are missing. Furthermore, depletion of either Mrc1 or its partner, Tof1, suppresses the asymmetric DNA synthesis in rad53-1 mutant cells. Thus, the DNA replication checkpoint pathway couples leading- and lagging-strand DNA synthesis by attenuating the replication function of Mrc1-Tof1 under replication stress.

Rad53 | Mrc1 | replication stress | deleterious ssDNA | asymmetric DNA synthesis

Replication stress, broadly defined as the events that impede induced by a variety of internal and external means (1–3). For instance, DNA lesions, deoxynucleotide triphosphate (dNTP) depletion, oncogene activation, and transcription–replication conflicts all lead to DNA replication stress (2). If left undealt with, replication stress will promote genome instability (2, 4) and drive tumorigenesis (5, 6). To meet these challenges, eukaryotic cells have developed the DNA replication checkpoint pathway to deal with various forms of replication stress, and inactivating genes in this pathway leads to genome instability in yeast and mammalian cells (3, 7–10).

At the molecular level, the checkpoint kinase Mec1 (ATR in mammals) is activated in response to DNA replication stress (3). Activated Mec1 phosphorylates the adaptor protein Mrc1 (Claspin in mammals) (11), which in turn promotes Rad53 (Chk1 in mammals) phosphorylation and activation. In the absence of Mrc1, Rad9, which in general serves as the adaptor protein for Rad53 activation in response to DNA damage, can also activate Rad53 during replication stress (12). Activated Rad53 performs multiple tasks, including 1) inhibition of late replication origin firing (13–15), 2) up-regulation of dNTP levels (16–19), and 3) prevention of replication fork collapse (20-22). Rad53 phosphorylates distinct proteins to regulate dNTP synthesis and late replication origin under replication stress. For instance, Rad53 phosphorylates Dun1, which in turn phosphorylates Sml1 and promotes Sml1 degradation (16). Sml1 is an inhibitor for ribonucleotide reductase (RNR) involved in the rate-limiting step of conversion of NTPs to dNTPs (16). Furthermore, Rad53 phosphorylates multiples sites on Sld3 and Dbf4, two proteins involved in DNA replication initiation,

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to inhibit late replication origin firing (14, 15, 23). Genetic studies indicate that the essential function of Mec1 (ATR) and Rad53 (Chk1) is to prevent the collapse of replication forks under replication stress (20–23), a process that lacks molecular insights. Based on the known mechanisms whereby Rad53 up-regulates dNTP levels and inhibits late replication origin firing, it is likely that Rad53 phosphorylates several proteins to prevent fork collapse under replication stress.

Recent studies indicate that Mec1 and Rad53 prevent fork collapse through inhibition of the generation of deleterious singledstranded DNA (ssDNA) under replication stress. While short ssDNA coated by ssDNA-binding protein (RPA) is important for checkpoint activation (8, 24, 25), long stretches of ssDNAs are detected in replication checkpoint-deficient cells under replication stress in both yeast and human cells (19, 20, 26, 27). In human cells, inhibition of ATR kinase using ATR inhibitors results in excessive ssDNAs, which depletes ssDNA-binding protein RPA and leads to fork collapse and genome instability (26). Using bromodeoxyuridine (BrdU)-immunoprecipitation and strand specific sequencing (BrdU-IP-ssSeq), which can measure the relative amount of leading- and lagging-strand DNA

Significance

During the S phase of the cell cycle, the DNA replication machinery accurately duplicates the genome in spite of numerous hurdles that cause replication stress. The replication checkpoint pathway deals with the replication stress, which, if left undealt with, can lead to fork collapse and genome instability, a process that remains poorly understood. Here, we show that the DNA checkpoint pathway couples leading- and lagging-strand DNA synthesis to prevent the generation of deleterious singlestranded DNA under replication stress by attenuating the replication function of Mrc1, a protein involved in both DNA replication and replication checkpoint, providing a mechanistic insight into how eukaryotic cells overcome replication stress.

The authors declare no competing interest.

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synthesis in budding yeast (28), we observed that lagging-strand DNA synthesis proceeds much farther than leading strand, which leads to the exposure of long stretches of single-stranded leading-strand templates in replication checkpoint deficient *rad53-1* mutant cells under replication stress. Generation of long stretches of ssDNA at the leading-strand template is not due to nucleolytic processing of DNA by 5'-3' Exo1 exonuclease or 3'-5' exonuclease activity of Pol ε , Pol δ , or Mre11. Based on these results, we proposed that Rad53 couples leading- and lagging-strand DNA synthesis under replication stress. However, it is not known whether the upstream kinase Mec1 and the adaptor protein Mrc1 also function similarly to Rad53 to prevent the generation of excessive ssDNA under replication stress.

In addition to serving as an adaptor protein for Rad53 activation under replication stress, Mrc1 also functions in DNA replication under normal growth. Mrc1 is a component of the replication progression complex including Cdc45, the Mrc1-Tof1-Csm3 complex, and the histone chaperone FACT (29). Moreover, Mrc1 interacts with both Cdc45 (30) and leadingstrand DNA polymerase ε (31). Mrc1 and Tof1 are recruited to DNA replication forks. In vitro, Mrc1 and Tof1-Csm3 are required for efficient DNA replication in the reconstituted DNAreplication system using purified proteins (32). Moreover, it has been shown that Mrc1 can stimulate the CMG helicase activity as well as the DNA polymerase activity (33, 34). Studies in both yeast Mrc1 and Xenopus Claspin indicate that the function of Mrc1/ Claspin in DNA-replication checkpoint and DNA replication is separable (35, 36). For instance, deletion of Mrc1 or depletion of its mammalian homolog Claspin results in impaired S phase in a role independent of checkpoint function (35, 36). Moreover, the mrc1-AQ mutant containing mutations at all possible Mec1 S/TQ phosphorylation sites, while showing defects in checkpoint functions such as firing of late replication origins under replication stress, progresses through S phase normally, indicating that this mutation does not affect the replication function of Mrc1. Under replication stress, the Mrc1-Pole interaction is altered (31), and Mrc1 and Tof1 promote the formation of a stable replicationpausing complex at replication forks (30). These results suggest that the role of Mrc1 in DNA-replication checkpoint and DNA replication, while genetically separable, may be connected.

Here, we show that lagging-strand DNA synthesis also proceeds much farther than leading strands in mec1-100 and mrc1-AQ mutant cells defective in DNA-replication checkpoint. Thus, coupling leading- and lagging-strand DNA synthesis under replication stress is an inherent and previously not-well-understood function of the DNA-replication checkpoint pathway. Surprisingly, the asymmetric DNA synthesis is not detected in $mrc1\Delta$ mutant cells in which both of the replication and checkpoint functions of Mrc1 are impaired. Furthermore, depletion of Mrc1 or Tof1 suppresses the asymmetric DNA synthesis observed in rad53-1 mutant cells under replication stress. Thus, the replication function of Mrc1 contributes to the asymmetric DNA synthesis observed in both rad53-1 and mrc1-AQ mutant cells under replication stress. Therefore, we propose that in response to DNA-replication stress, once Rad53 is activated by Mrc1, activated Rad53 will attenuate the function of Mrc1-Tof1 in DNA replication to couple leading- and laggingstrand DNA synthesis.

Results

DNA Synthesis at Lagging Strand Progresses Much Farther than Leading Strand in the *mec1-100* Mutant Cells under Replication Stress. We have shown previously that Rad53 couples leadingand lagging-strand DNA synthesis under replication stress (28). When cells with the *rad53-1* mutation, which impairs Rad53's DNA-replication checkpoint function, were treated with hydroxyurea (HU) that depletes dNTP pools and induces replication stress, lagging-strand DNA synthesis proceeds much farther than leading strand. Mec1 functions upstream of Rad53 and activates Rad53 in response to replication stress. To test whether Mec1 also functions similarly to Rad53 in regulation of DNA synthesis under replication stress, we analyzed DNA synthesis in mec1-100 cells under replication stress using BrdU-IP-ssSeq. It is known that the mec1-100 mutant cells are defective in the DNAreplication checkpoint but maintain the G2/M checkpoint (37, 38). Briefly, wild-type (WT) and mec1-100 cells were arrested at G1 and then released into early S phase in the presence of HU and BrdU, a nucleotide analog incorporated into newly synthesized DNA. We then collected cells for DNA isolation and subsequent immunoprecipitation of newly synthesized DNA using antibodies against BrdU. The resulting DNA was subjected to BrdU-IP-ssSeq (Fig. 1A). Consistent with previous publications (28), WT cells treated with HU, while displaying no apparent effect on initiation of DNA replication from early replication origins, showed inhibition of firing of late replication origins (Fig. 1 B-D). Under the same conditions, late replication origins fired in both rad53-1 (28) and mec1-100 cells under replication stress (Fig. 1 B-D). Moreover, BrdU-IP-ssSeq peaks in WT cells showed a slight bias toward the leading strand (Fig. 1 C and E). In contrast, like rad53-1 mutant cells (28), BrdU-IP-ssSeq peaks in mec1-100 cells showed a strong laggingstrand bias at both early and late replication origins (Fig. 1 C, E, and F), indicating that DNA synthesis at lagging strands proceeds much farther than that of leading strands under replication stress. These results indicate that leading- and lagging-strand DNA synthesis in mec1-100 cells are uncoupled under replication stress and that Mec1 also functions like Rad53 to regulate DNA synthesis in response to DNA replication stress.

Excessive Single-Stranded Leading-Strand Template Is Detected in mec1-100 Mutant Cells under Replication Stress. If DNA synthesis in mec1-100 mutant cells proceeds farther along lagging strands than leading strands, one would expect that long stretches of leading-strand template are exposed in mec1-100 mutant cells (Fig. 2A). To test this idea, we analyzed the distribution of Rfa1 at replication forks in mec1-100 cells using Rfa1 immunoprecipitation and strand-specific sequencing (ChIP-ssSeq) (39, 40). ChIP-ssSeq detects both template and newly synthesized DNA, whereas BrdU-IP-ssSeq is used to detect new synthesized DNA. Therefore, we used (+/-)bias to describe Rfa1 ChIP-ssSeq peaks, in contrast to the leading- and lagging-strand bias for BrdU-IP-ssSeq (39). Rfa1 is the large subunit of the ssDNA-binding protein RPA. Rfa1 was detected at early, but not late, replication origins in WT cells. In contrast, Rfa1 was detected in both early and late replication origins in *mec1-100* and *rad53-1* cells treated with HU (Fig. 2 B-D) (28), consistent with the idea that late origins fire in mec1-100 cells. As reported previously, Rfa1 ChIP-ssSeq in WT cells showed (+) strand bias, indicating that RPA is enriched at lagging template strands in WT cells (Fig. 2 A, B, and E) (39). In contrast, Rfa1 ChIP-ssSeq peaks in mec1-100 cells, like those in rad53-1 cells (28), showed an opposite, but strong (-) bias at both early and late replication origins (Fig. 2 B, E, and F), indicating that RPA is enriched at leading-strand templates in mec1-100 cells under replication stress. These results support the idea that excessive leading-strand templates compared to lagging template strand are exposed and coated with RPA in mec1-100 cells under replication stress. Notably, these results do not exclude the possibility that single-stranded lagging-strand templates are also present in mec1-100 cells.

Deletion of SML1 in mec1-100 Cells Suppresses Uncoupled DNA Synthesis at Early, but Not Late, Replication Origins. We have previously shown that deletion of SML1, which encodes an inhibitor for RNR, in rad53-1 cells leads to the suppression of asymmetric DNA synthesis at early replication origins and, to a lesser extent, at late replication origins (28). We therefore analyzed the effect of SML1 deletion in mec1-100 cells on DNA synthesis and Rfa1 distribution at replication forks under replication stress. As shown in Fig. 3A, deletion of SML1 had no apparent effects on



Fig. 1. The *mec1-100* mutation results in asymmetric DNA synthesis under replication stress. (A) Schematic representation of the experimental procedures to analyze DNA synthesis using BrdU-IP-ssSeq and to analyze the association of RPA at replication forks using ChIP-ssSeq under replication stress. (B) Binary representation of the firing status of DNA-replication origins in WT, *rad53-1*, or *mec1-100* cells released into S phase for 45 min in the presence of 0.2 M HU. The firing status of each DNA-replication origin is defined by the density of BrdU-IP-ssSeq. (C) Snapshot of BrdU-IP-ssSeq signals surrounding an early (*Left*) or late (*Right*) replication origins in cells treated with HU with their relevant genotype marked in the middle. (*D*) Normalized read density of BrdU-IP-ssSeq signals within 20 kb of late replication origins in cells treated with HU. Normalized reads per kilobase per million of mapped reads (RPKM) surrounding replication origins was calculated using a 200-bp sliding window. (*E* and *F*) The average bias of BrdU-IP-ssSeq peaks around early (*E*) or late (*F*) replication origins under HU-induced replication stress. The average log₂ ratio of sequencing reads from the Watson over Crick strands of BrdU-IP-ssSeq was calculated using a 200-bp sliding window.

the firing of late replication origins in mec1-100 cells. However, the BrdU track length in $mec1-100 \ sml1\Delta$ cells appeared to be broader than mec1-100 cells at early replication origins, likely due to elevation of dNTP levels trigged by SML1 deletion (Fig. 3 B and C). In contrast to the strong lagging strand bias in mec1-100 cells, BrdU-IP-ssSeq peaks at early replication origins in $mec1-100 \ sml1\Delta$ cells showed no apparent bias. Furthermore, the strong lagging-strand bias at late replication in mec1-100 $sml1\Delta$ cells was also detected, albeit at a reduced amplitude compared to $mec1-100 \ cells$ (Fig. 3 D and E). These results indicate that deletion of SML1 suppresses the asymmetric DNA synthesis at early, but not late, replication origins.

Analysis of the distribution of RPA using Rfa1 ChIP-ssSeq indicates that deletion of *SML1* in *mec1-100* cells did not affect the overall peak width at both early and late replication origins compared to *mec1-100* cells (Fig. 4*A* and *B*). However, deletion of *SML1* in *mec1-100* cells completely suppressed the Rfa1 ChIP-ssSeq bias at early replication origins while having little effects on Rfa1 ChIP-ssSeq peak bias at late replication origins (Fig. 4*C* and *D*). These results indicate that *SML1* deletion can suppress uncoupled DNA synthesis at early, but not late, replication origins in *mec1-100* cells under replication stress and that an increase in dNTP levels can help couple leading- and lagging-strand DNA synthesis at forks originated from early replication origins.

Uncoupled DNA Synthesis Is Detected in *mrc1-AQ*, but Not *mrc1* Δ Cells, under Replication Stress. It is known that Mec1 phosphorylates Mrc1, which in turn activates Rad53 under replication stress (3, 35). We therefore analyzed whether deletion of *MRC1* affects DNA synthesis and ssDNA generation in a manner similar



Fig. 2. Impairment of Mec1 replication checkpoint function leads to RPA accumulation on the leading-strand template under replication stress. (A) Schematic representation of RPA binding to single-stranded template DNA during DNA replication. RPA is enriched at lagging strands in WT cells, whereas many more RPA proteins bind to long stretches of leading template strand in *mec1-100* or *rad53-1* mutant defective in replication checkpoint. (B) Snapshots of Rfa1 ChIP-ssSeq signal around an early (*Left*) or late (*Right*) replication origin in cells treated with HU. (C and D) Normalized read density of Rfa1 ChIP-ssSeq signal within 20 kb of early (C) or late (*D*) replication origins in cells treated with HU. Reads were normalized to reads per kilobase per million of mapped reads (RPKM) and calculated using a 200-bp sliding window. (*E* and *F*) Average strand bias of Rfa1 ChIP-ssSeq peaks at early (*E*) or late (*F*) replication origins. The average log₂ ratio of Rfa1 ChIP-ssSeq sequence reads from the Watson over Crick strands was calculated using a 200-bp sliding window.

to rad53-1 and mec1-100 mutations. Late replication origins fired in the presence of HU in mrc1 Δ cells (Fig. 5 A–C), consistent with published studies showing that Mrc1 is important to inhibit firing of late replication origins in response to DNA-replication stress (30). The reduced BrdU density in mrc1 Δ cells at early replication origins (Fig. 5B) is likely due to impaired DNA synthesis in these cells (32, 35). Surprisingly, BrdU-IP-ssSeq peaks in mrc1 Δ cells did not show any bias toward lagging strands (Fig. 5 D–E), indicating that leading- and lagging-strand DNA synthesis is not asymmetrically affected in mrc1 Δ cells, which is in contrast to mec1-100 or rad53-1 cells. Because dNTP levels in mrc1 Δ were not elevated compared to rad53-1 or mec1-100 cells (41), the distinct effect of mrc1 Δ on DNA synthesis compared to mcc1-100 and rad53-1 mutations is likely not due to an increase in dNTP levels in mrc1 Δ cells.

It has been reported that Rad9 mediates Rad53 activation under replication stress in the absence of Mrc1 (12, 42). Therefore, it is possible that Rad9 functions to couple leadingand lagging-strand DNA synthesis in the absence of Mrc1. To test this idea, we made the *rad9-aid* degron in both WT and *mrc1* Δ cells and induced its degradation 30 min before cells were released from G1 phase into S phase. Rad9-auxin-inducible degron (AID) was efficiently degraded with the addition of indoleacetic acid (IAA), the plant pheromone for induction of degradation of AID-tagged proteins (*SI Appendix*, Fig. S1A), and Rad9 depletion had no apparent effects on the firing of late replication origins (Fig. 5A and *SI Appendix*, Fig. S1 B and C). BrdU-IP-ssSeq peaks from $mrc1\Delta$ rad9-aid cells did not show any significant bias compared to $mrc1\Delta$ or rad9-aid mutants alone (*SI Appendix*, Fig. S1 D and E), suggesting that uncoupled DNA synthesis was not detectable even in cells lacking both adaptors for Rad53 activation under replication stress.

In addition to serving as an adaptor protein for Rad53 activation during replication stress, Mrc1 also has a role in DNA replication (12, 36). The *mrc1-AQ* allele is a separation of function mutant that is defective in Mec1-mediated Rad53 activation but retains the replication function of Mrc1 (35). Therefore, we analyzed the impact of this mutant on DNA synthesis under replication stress using BrdU-IP-ssSeq. Like with *mrc1A* treated with HU, late replication origins also fired in *mrc1-AQ* mutant cells (Fig. 5A), consistent with a defect in DNA replication checkpoint in these mutant cells. In contrast to *mrc1A*, BrdU-IP-ssSeq peaks in *mrc1-AQ* cells showed a bias toward lagging strands (Fig. 5 D and E). This bias pattern was similar to *rad53-1* and *mc1-100* mutant cells and was opposite to that of WT and *mrc1A* cells. We noted that the bias amplitude of the BrdU-IP-ssSeq peaks in *mrc1-AQ* cells was smaller than that in *rad53-1* and *mec1-1* mutant



Fig. 3. Deletion of *SML1* in *mec1-100* cells suppresses the asymmetric DNA synthesis at early replication origins under replication stress. (*A*) Binary representation of the firing status of DNA-replication origins in *mec1-100*, *sml1* Δ , or *mec1-100 sml1* Δ cells treated with 0.2 M HU for 45 min. (*B* and *C*) Normalized read density of BrdU-IP-ssSeq within 20 kb of early (*B*) or late (*C*) replication origins in cells treated with HU. (*D* and *E*) The average bias of BrdU-IP-ssSeq peaks at early (*D*) or late (*E*) replication origins under replication stress by HU.

cells (compare Fig. 5 D and E to Fig. 1 E and F), suggesting that DNA synthesis at leading and lagging strands is also uncoupled in mrc1-AQ cells but to a lesser degree than in rad53-1 and mec1-100 cells.

We also asked whether depletion of Rad9 could increase the BrdU-IP-ssSeq bias in *mrc1-AQ* cells. We found that induced degradation of Rad9-AID by IAA in *mrc1-AQ* cells did not affect BrdU density at both early and late replication origins (*SI Appendix*, Fig. S2 *A*–*C*). Moreover, Rad9 depletion slightly increased the bias

of BrdU-IP-ssSeq peaks at both early and late replication origins. These results suggest that Rad9 plays a minor role in coupling leading- and lagging-strand DNA synthesis in *mrc1-AQ* cells under replication stress.

Finally, we also analyzed the effect of *SML1* deletion on the asymmetric DNA synthesis in *mrc1-AQ* cells. We observed that deletion of *SML1* suppressed the asymmetric DNA synthesis at both early and late replication origins in *mrc1-AQ* cells based on



Fig. 4. Deletion of *SML1* in *mec1-100* cells suppresses the enrichment of RPA at leading-strand templates at early but not late replication origins. (*A* and *B*) Normalized read density of Rfa1 ChIP-ssSeq signal within 20 kb of early (*A*) or late (*B*) replication origins in cells treated with HU. (*C* and *D*) The average bias of Rfa1 ChIP-ssSeq at early (*C*) or late (*D*) replication origins.



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Fig. 5. The *mrc1-AQ* mutation, but not *MRC1* deletion, leads to the generation of ssDNA under replication stress. (*A*) Binary representation of early- and latefiring replication origins in *mrc1* Δ , *rad9-aid*, *mrc1* Δ *rad9-aid*, or *mrc1-AQ* cells treated with 0.2 M HU for 45 min. *rad9-aid* and *mrc1* Δ *rad9-aid* cells were treated with IAA for Rad9-AID degradation before used for BrdU-IP-ssSeq. (*B* and *C*) Normalized read density of BrdU-IP-ssSeq within 20 kb at early (*B*) or late (*C*) replication origins in cells treated with HU. (*D* and *E*) The average bias of BrdU-IP-ssSeq peaks at early (*D*) or late (*E*) replication origins in different mutant cells under HU-induced replication stress.

analysis of BrdU-IP-ssSeq bias (*SI Appendix*, Fig. S3A and B) and Rfa1 ChIP-ssSeq bias (*SI Appendix*, Fig. S3 C and D) in mrc1-AQ sml1 Δ cells. The differential effects of SML1 deletion on DNA synthesis at late replication origins in mrc1-AQ cells compared to mec1-100 (Fig. 3E) and rad53-1 cells (28) are currently unclear and warrant further investigation.

Taken together, these results indicate that while late replication origins fire in both mrc1-AQ and $mrc1\Delta$ cells under replication stress, leading- and lagging-strand DNA synthesis is uncoupled only in mrc1-AQ cells defective in replication checkpoint but not in $mrc1\Delta$ mutant cells in which both the replication and checkpoint functions of Mrc1 are impaired.

Depletion of Mrc1 or Tof1 in rad53-1 Mutant Cells Suppresses the Asymmetric DNA Synthesis under Replication Stress. While both DNA replication and checkpoint functions are destroyed in $mrc1\Delta$ cells, the DNA-replication function remains intact in mrc1-AQ mutant cells. Therefore, we hypothesize that an intact replication function of Mrc1 contributes to the asymmetric DNA synthesis in mrc1-AQ cells. To explore this idea, we tested whether the replication function of Mrc1 is also needed for the observed asymmetric DNA synthesis in rad53-1 mutant cells. We generated Mrc1-AID degron mutant in rad53-1 mutant cells and started to deplete Mrc1 in G1 phase 30 min prior to release into S phase in the presence of HU and BrdU for analysis of DNA synthesis by BrdU-IP-ssSeq. Mrc1 was efficiently depleted with the addition of IAA (Fig. 6A). Moreover, depletion of Mrc1 in rad53-1 mutant cells resulted in a slight delay in S-phase progression under normal growth conditions (SI Appendix, Fig. S4). Next, we analyzed how depletion of Mrc1 in rad53-1 mutant cells affected DNA synthesis under replication stress. Depletion of Mrc1-AID in rad53-1 cells had no apparent effect on the firing of both early and late replication origins (SI Appendix, Fig. S5A). Moreover, BrdU-ssSeq peak height was also largely unaffected upon depletion of Mrc1-AID (SI Appendix, Fig. S5 B and C). Remarkably, depletion of Mrc1-AID in rad53-1 mutant cells completely suppressed the asymmetric DNA synthesis at both early

and late replication origins in *rad53-1* mutant cells (Fig. 6 *B–D*). These results indicate that an intact replication function of Mrc1 also contributes to the asymmetric DNA synthesis in *rad53-1* cells under replication stress.

Mrc1 forms a complex with Tof1-Csm3 to regulate DNA replication under normal and replication stress conditions (30, 35, 43). Therefore, we tested whether deletion of Tof1 also suppresses the asymmetric DNA synthesis observed in *rad53-1* mutant cells under replication stress. Like Mrc1 depletion, Tof1 depletion in *rad53-1* mutant cells completely suppressed the asymmetric DNA synthesis at both early and late replication origins (Fig. 6 *B–D*). These results support the idea that an intact replication function of the Mrc1-Tof1 complex contributes to the asymmetric DNA synthesis and the generation of deleterious ssDNA in *rad53-1* mutant cells under replication stress. All together, these results suggest that Rad53, once activated by Mrc1 under replication stress, must attenuate the replication function of Mrc1-Tof1 to couple leading- and lagging-strand DNA synthesis under replication stress (Fig. 6*F* and *Discussion*).

rad53-1 mutant cells are highly sensitive to HU. Therefore, we tested whether Mrc1 or Tof1 depletion also suppressed HU sensitivity. Interestingly, we observed that depletion of Mrc1 or Tof1 in *rad53-1* mutant cells led to increased HU sensitivity compared to *rad53-1* mutant cells (Fig. 6E). These results indicate that the HU sensitivity detected in *rad53-1* mutant cells is not exclusively linked to the asymmetric DNA synthesis observed in these mutant cells treated with HU.

Mcm2 Moves ahead of DNA Synthesis in mrc1-AQ, but Not mrc1 Δ , Cells Compared to WT Cells under Replication Stress. We have shown previously that both Mcm6 and Pol2 move ahead of DNA synthesis in rad53-1 cells under replication stress (28). Others also reported that DNA-replication proteins, including Cdc45 and Pol2, move beyond DNA synthesis in mrc1 Δ cells (30). We therefore analyzed the distribution of Mcm2, a subunit of the CMG helicase, in WT, mrc1 Δ , and mrc1-AQ cells when these cells were released from G1 block into early S phase in the



presence HU and BrdU. In WT cells, we observed that Mcm2 chromatin immunoprecipitation and sequencing (ChIP-Seq) peaks displayed a valley at replication origins surrounded by two peaks, suggesting that Mcm2 travels with two forks and moves away from the site of replication initiation under this time course (*SI Appendix*, Fig. S64). However, the valley of Mcm2 ChIP-Seq peaks in $mrc1\Delta$ and mrc1-AQ cells was not detectable. The functional implication of these changes in peak shape in these mutant cells compared to WT cells is not clear.

To determine whether Mcm2 moves ahead of DNA synthesis, we calculated the average peak length of Mcm2 ChIP-Seq peaks normalized against the corresponding BrdU-IP-Seq peak length in WT, $mrc1\Delta$, and mrc1-AQ cells (SI Appendix, Fig. S6 B and C). We observed that the average normalized Mcm2 peak length in $mrc1\Delta$ was similar to that of WT cells, whereas the average Mcm2 ChIP-Seq peak length in mrc1-AQ cells increased compared to WT cells (SI Appendix, Fig. S6C). If we assume that Mcm2 travels the same distance as DNA synthesis (BrdU track length) in WT cells, these results indicate that Mcm2, and likely the CMG helicase, travels ahead of DNA synthesis only in mrc1-AQ, but not in mrc1 Δ cells. Currently, we do not have an explanation for the discrepancy of the effect of $mrc1\Delta$ on the distribution of MCM proteins at replication forks described here and those reported by Katou et al. (30). One possibility is that different methods were used to analyze the distribution of replication proteins. Katou et al. utilized a ChIP-chip method based on an oligonucleotide array to analyze the distribution of replication proteins at chromosome VI with a 300-bp resolution. In contrast, we used ChIP-Seq to analyze the Mcm2 distribution genome wide at single-base-pair resolution. Nonetheless, our results are consistent with our observations that the asymmetric DNA synthesis is detected in mrc1-AQ but not mrc1 Δ cells under replication stress and are also consistent with the observation that Mrc1 stimulates the helicase activity of the CMG helicase in vitro (33) (Discussion).

Discussion

Coupling Leading- and Lagging-Strand DNA Synthesis under Replication Stress Is an Inherent Function of the DNA-Replication Checkpoint Pathway. Previously, we reported that lagging-strand DNA synthesis proceeds much farther than leading strand synthesis in rad53-1 cells under replication stress, resulting in the exposure of a long stretch of leading-strand template coated with RPA (28). Rad53 is an effector kinase activated by Mec1 via the adaptor protein Mrc1 in response to replication stress (3). Therefore, we tested whether mutations at MEC1 and MRC1 also show the same asymmetric DNA synthesis phenotypes observed in rad53-1 mutant cells under replication stress. We found that the asymmetric DNA synthesis was also detected in mec1-100 and mrc1-AQ mutant cells defective in replication checkpoint. Therefore, coupling leading- and lagging-strand DNA synthesis under replication stress is an inherent function of the DNA-replication checkpoint pathway. In vitro, leading- and lagging-strand polymerases function autonomously within a replisome (44), indicating a lack of inherent coordination between replisome components involved in leading- and lagging-strand DNA synthesis. We suggest that the lack of coordination between replisome components in the synthesis of leading and lagging strands may not affect DNA replication and cell fitness under normal growth conditions. However, under replication stress, this lack of coordination between replisome components will lead to the generation of deleterious ssDNA. Consistent with this idea, it has been shown that in the presence of a block on the leading strand, lagging-strand DNA synthesis proceeds further in Escherichia coli (45). We suggest that the DNA-replication checkpoint pathway in eukaryotic cells has evolved to regulate replisome components so that leading- and lagging-strand DNA synthesis is coupled under replication stress.

Mechanistically, we observed that deletion of *SML1*, which leads to an increase in dNTP concentrations, in *mec1-100, mrc1-AQ*, and *rad53-1* mutant cells completely suppresses the

asymmetric DNA synthesis at early replication origins, while its effects on the asymmetric DNA synthesis at forks originated from late replication origins vary among checkpoint mutants analyzed with minor effects in mec1-100 and dramatic effects in mrc1-AQ cells. Furthermore, depletion of Mrc1 or its partner Tof1 in DNA replication completely suppresses the asymmetric DNA synthesis in rad53-1 mutant cells at both early and late replication origins, suggesting that the replication function of Mrc1 and Tof1, if left unchecked, contributes to the asymmetric DNA synthesis in rad53-1 cells. Consistent with this interpretation, we detect asymmetric DNA synthesis in mrc1-AQ cells defective in only the checkpoint function but not in $mrc1\Delta$ cells in which both DNA replication and checkpoint functions are impaired. Therefore, we propose that in response to replication stress, once activated by Mec1 via adaptor protein Mrc1, Rad53 performs at least two functions to couple leading- and lagging-strand DNA synthesis: up-regulation of dNTP levels and attenuation of the replication function of Mrc1-Tof1 (Fig. 6F and SI Appendix, Fig. S7).

Why Does Deletion of SML1 Have Differential Effects on Replication Forks Originated from Early and Late Replication Origins in mec1-100 Mutant Cells under Replication Stress? Deletion of SML1 in mec1-100, mrc1-AQ, and rad53-1 mutant cells completely suppresses the asymmetric DNA synthesis of forks originated from early replication origins. In contrast, deletion of SML1 had minor effects on the asymmetric DNA synthesis at late replication origins in mec1-100 cells as detected both by BrdU-IP-ssSeq and RPA ChIPssSeq (Figs. 3 and 4). SML1 deletion largely, but not completely, suppresses the asymmetric DNA synthesis at late replication origins in rad53-1 cells as detected by BrdU-IP-ssSeq (28) and completely suppressed asymmetric DNA synthesis in mrc1-AQ cells. One likely explanation for the differential effects of SML1 deletion on the asymmetric DNA synthesis at late replication origins in rad53-1, mrc1-AQ, and mec1-100 cells is that these mutant alleles affect the replication checkpoint pathway to different extents. Consistent with this idea, the bias of BrdU-IP-ssSeq peaks in mrc1-AQ cells is the smallest. Nonetheless, these studies indicate that in both rad53-1 and *mec1-100* mutant cells, the effect of *SML1* deletion on the asymmetric DNA synthesis at forks from early replication origins is different from that on late replication origins.

Replication at different chromosome regions in both yeast and mammalian cells occurs in a well-defined spatiotemporal program (46). In general, euchromatin regions replicate early in S phase and heterochromatin regions replicate late in S phase. In budding yeast, the majority of chromatin is euchromatic and replication origins are "site specific" and efficient. However, it is known that the chromatin environment can influence when an origin fires in the S phase of the cell cycle. For instance, when the early replication origin ARS1 is placed near a telomere, initiation of replication from telomeric ARS1 is delayed due to the formation of SIR-dependent heterochromatin (47). Moreover, deletion of Rpd3, a histone deacetylase, in budding yeast results in early activation of late replication origins (48). Furthermore, nucleosomes at early replication origins are different from late replication origins (49) and are a determinant factor for origin selection and functions (50). Finally, local chromatin environment can influence the loading of MCM helicase (51). Therefore, it is possible that differences in chromatin environment at early and late replication origins will impact the composition of the replisome at early and late replication origins, which in turn contributes to differential dependence of replication forks originated from early and late replication origins on dNTP levels. Future studies are needed to test this and other hypotheses.

How Does Mrc1 Couple Leading- and Lagging-Strand DNA Synthesis under Replication Stress? Mrc1 and its mammalian homolog Claspin were first discovered as the primary adaptor protein for activation of effector kinase Rad53 in yeast and Chk1 in

mammalian cells (11, 12, 36). Later on, studies from various groups indicate that Mrc1/Claspin also function in DNA replication. Moreover, the function of Mrc1/Claspin in DNA replication and DNA-replication checkpoint are separable (32, 35, 36, 52). We presented two lines of evidence supporting the idea that the DNA replication function of Mrc1 and Tof1 contributes to asymmetric DNA synthesis in checkpoint mutant cells under replication stress. First, we observed asymmetric DNA synthesis in mrc1-AQ mutant cells defective only in checkpoint function but not in mrc1 Δ cells in which both replication and checkpoint functions of Mrc1 are compromised. Second, depletion of Mrc1 or Tof1 completely suppresses the asymmetric DNA synthesis in rad53-1 mutant cells treated with HU. These results indicate that the asymmetric DNA synthesis observed in rad53-1 mutant cells depends on the replication function of Mrc1 and Tof1. It is possible that depletion of Mrc1 or Tof1 in rad53-1 mutant cells results in a reduced fork speed, which helps couple leading- and laggingstrand DNA synthesis under replication stress. Two observations suggest that this is unlikely the only explanation for the suppression of asymmetric DNA synthesis in rad53-1 mutant cells. First, in the in vitro reconstituted DNA-replication reactions, Mrc1 and Tof1 stimulate DNA synthesis to different extents (32), and yet depletion of Mrc1 or Tof1 has similar/identical effects on the suppression of the asymmetric DNA synthesis in rad53-1 mutant cells. Second, depletion of Mrc1 or Tof1 in rad53-1 mutant cells did not affect the BrdU track length (SI Appendix, Fig. S5 B and C), which reflects replication-fork moving distance, to a detectable degree. Therefore, we suggest that under replication stress, the function of Mrc1 and Tof1 in DNA replication is attenuated to couple leading- and lagging-strand DNA synthesis.

Previously, it has been shown that Mrc1 interacts with several proteins involved in DNA replication. First, Mrc1 is a component of the replication progression complex (29) consisting of Cdc45, Mrc1-Tof1-Csm3, and histone chaperone FACT that is essential for chromatin replication. Cross-linking mass spectrometry analysis of Mrc1-Tof1-Csm3 in complex with the CMG helicase indicates that Mrc1 makes extensive contact with the CMG helicase (53). Second, Mrc1 also interacts with leading-strand DNA polymerase ε (31). Moreover, it has been shown that Mrc1 can stimulate the enzymatic activity of Pol ε (34). While this manuscript was under review, McClure and Diffley reported in BioRxiv that Mrc1 can stimulate the helicase activity of the CMG helicase. Furthermore, Mrc1 is phosphorylated by Rad53 (12, 35, 54), and Rad53-mediated phosphorylation of Mrc1 inactivates its ability to stimulate the helicase activity of the CMG helicase and DNAsynthesis rate (33). We have shown that Mcm2 moves ahead of DNA synthesis in mrc1-AQ with an intact ability to stimulate the helicase activity but not in $mrc1\Delta$ cells in which the ability of Mrc1 to stimulate the CMG activity is missing. Based on these results, we propose the following working model for how Mrc1 couples leading- and lagging-strand DNA synthesis under replication stress. Under normal replication conditions, Mrc1 and its partners Tof1 and Csm3 promote DNA synthesis by stimulating the activities of the CMG helicase and, possibly, Pol ε , which in turn enhance DNA synthesis. In response to replication stress, the temporary uncoupling of CMG helicase and DNA polymerases generates short ssDNA, which will be coated by RPA and serves as the platform for Mec1 activation (3, 8). Activated Mec1 phosphorylates Mrc1, which brings Rad53 closer to Mec1 for phosphorylation and activation. Activated Rad53 phosphorylates Mrc1, and this phosphorylation attenuates the ability of Mrc1 to stimulate the CMG helicase activity and thereby slows down DNA unwinding under replication stress. In this way, DNA synthesis at leading and lagging strand is coupled. Future studies are needed to test this and other models.

Materials and Methods

Yeast Strains. All yeast strains used in this study are listed in *SI Appendix*, Table S1 and were derived from W303 (*leu2-3,112 ura3-1 his3-11, trp1-1,* ade2-1 can1-100). Gene deletions were generated using a one-step replacement with marker cassettes. The *mec1-100* mutation was introduced by crossing with the strain YLL750 (38). For protein depletion, Rad9, Mrc1, or Tof1 was carboxyl-terminally tagged with AID*-9Myc using a one-step insertion into the strain YNK54 (55) with a marker cassette amplified from the plasmid pKan-AID*-9myc or pHIS-AID*-9myc (56). The BrdU-Inc (BrdU-Incorporating) vector containing the genes necessary for BrdU incorporation was introduced by crossing with strains CVy43 or CVy63 (57).

Culture Growth and Cell-Cycle Synchronization. Yeast cells grown to optical density OD₆₀₀ = 0.4 to 0.5 in yeast extract peptone + 2% dextrose (YPD) were synchronized with α -factor (5 μ g/mL, synthesized by EZBiolab) for 1.5 h at 25 °C, adding an additional 5 μ g/mL α -factor after 1.5 h. G1-arrested cells were released into YPD medium at 30 °C containing 400 μ g/mL BrdU (Sigma-Aldrich B5002) and 0.2 M HU (Chem-Impex 24533). A total of 45 min after release into S phase at 30 °C, cell fixation was performed with 1% paraformaldehyde for 20 min at room temperature. For protein depletion experiments, IAA (Sigma-Aldrich I2886) or ethanol was added at G1, 30 min before release into S phase, at a concentration of 1 mM from a 0.5-M stock in ethanol. After cell washing at 30 °C, the same concentration of IAA was also added to medium before release.

ChIP-ssSeq and BrdU-IP-ssSeq. ChIP-ssSeq and BrdU-IP-ssSeq experiments were performed as described previously (28). Briefly, fixed cells were lysed with the glass beads beating method. Chromatin was pelleted, washed, and sheared by sonication using a Bioruptor Pico (Diagenode) to an average fragment size of 200 to 400 bp. Sheared chromatin was cleared by centrifugation followed by immunoprecipitation with anti-Rfa1 antibody (gift from Steven Brill, Rutgers University, Piscataway, NJ) or anti-Mcm2 antibody (gift from Bruce Stillman, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY) and protein G Sepharose beads (GE Healthcare). DNA was recovered from both input and ChIP samples following the Chelex-100 protocol (58) and was purified with the MinElute PCR kit (QIAGEN), and strand-specific sequence libraries were prepared using the Accel-NGS 15 Plus DNA library kit (Swift BioSciences).

Input DNA obtained from the Chelex-100 extraction was used for BrdU-IP. Briefly, DNA was denatured at 95 °C for 5 min followed by incubation in an ice-water bath for 5 min. Samples were then diluted 10-fold with BrdU-IP buffer solution (1× phosphate-buffered saline), 0.0625% Triton X-100 [vol/ vol], 6.7 µg/mL *E. coli* transfer ribonucleic acid, 0.40 µL/mL BrdU antibody [BD Bioscience]). After a 2-h incubation at 4 °C, 20 µL protein G Sepharose beads (GE Healthcare) was added followed by an additional 1-h incubation at 4 °C. Protein G beads were extensively washed, and DNA was eluted with 100 µL Tris-ethylenediaminetetraacetic acid buffer containing 1% sodium dodecyl sulfate (SDS) at 65 °C for 15 min. The eluted DNA was purified with Minelute PCR kit (QIAGEN) and sequenced after ssDNA library preparation using the Accel-NGS 1S Plus DNA library kit (Swift BioSciences).

Yeast Protein Extraction, SDS-PAGE (Polyacrylamide Gel Electrophoresis), and Western Blotting. Whole-cell protein extracts were prepared from 5 mL yeast culture right before fixation. Cells were collected by centrifugation, washed with cold water, and resuspended in 50 μ L cold Tris-buffered saline buffer containing 1 mM dithiothreitol and 1mM phenylmethylsulfonyl fluoride. An equal amount of glass beads was added to each tube, and cells were lysed by bead beating. Lysates were transferred to new tubes, mixed with 50 μ L 2x SDS-PAGE loading buffer, and boiled for 3 min. Alternatively, after washing with cold water, cells were resuspended in 300 μ L 20% trichloroacetic acid and lysed by bead beating with 300 μ L glass beads. Lysates were transferred to new tubes, and proteins were pelleted, resuspended in high-pH 1x SDS-PAGE loading buffer, and boiled for 4 min. For immunodetection of proteins tagged with AID*-9myc, 5 μ L protein extract was resolved in an 8% SDS-PAGE, transferred onto a nitrocellulose membrane, and detected using antimyc antibody (9E10).

HU Sensitivity Assay. Yeast strains were grown overnight at 30 °C in YPD. Cells were diluted to 6×10^6 cells/mL and four additional 10-fold serial dilutions and spotted onto media containing the indicated concentrations of HU and IAA (1 mM) or the same volume of ethanol. Plates were incubated at 30 °C for 3 d before pictures were taken.

Analysis of Cell-Cycle Progression. Yeast cells grown to OD₆₀₀ = 0.4 to 0.5 in YPD were synchronized at G1 with α -factor as described in *Culture Growth* and Cell-Cycle Synchronization. To deplete Mrc1-AID or Tof1-AID, 1 mM IAA or the same volume of ethanol was added at G1 30 min before release into S phase. These cells were then collected, washed with water, and released into

YPD medium at 30 °C with or without IAA. At each indicated time point, 1 mL cell culture was harvested by centrifugation, washed with water, and fixed with 70% ethanol. Samples were further washed with 50 mM Na-citrate pH 7.4 and resuspended in the same buffer containing 250 μ g/mL RNase A (Sigma-Aldrich R6513). Cells were briefly sonicated (three 1-s pulses at 30% amplitude, Sonics Vibra Cell VCX-500) and incubated at 50 °C for 1 h. Samples were then treated with 1 mg/mL Proteinase K (Invitrogen 25530015) for 1 h at 50 °C and stained with 20 μ g/mL propidium iodide (Sigma-Aldrich P4170). DNA content was measured using an Attune Nxt Flow Cytometer (Thermo Fisher Scientific).

Sequence Mapping and Data Analysis. The sequence reads were mapped to the yeast genome (sacCer3) using Bowtie2 software (59). The genome-wide read coverage of Watson and Crick strands was calculated using BEDTools (60) and in-house Perl programs. To determine whether an origin fired in cells treated with HU, we used BrdU-IP-ssSeq data. Briefly, BrdU-IP-ssSeq peaks were identified by merging the sequencing reads of both Watson and Crick strands and using them for peak calling with SICER software (61) with the cutoff false discovery rate (FDR) of 0.01. If there is a detectable peak using these conditions, the origin is classified as "fired." This binary classification does not imply that an origin classified as "fired" fires in each individual cell; it means that it fired in enough cells to identify its BrdU-IP peak. Moreover, all origins used in the analysis are located at the centers of their corresponding BrdU peaks, thereby removing many origins localized at telomeres from the analysis. The majority of origins classified by this approach overlap with origins identified using other means (40). The DNAreplication origins dataset used is listed in Dataset S1. Origins were

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classified as "early" if there was a detectable BrdU-IP-ssSeq peak in WT cells released into S phase for 45 min in the presence of HU. Otherwise, the origin would be classified as "late." To represent the firing state of each origin in Fig. 1*B*, origins were ranked based on the cluster of the matrix of the binary value. The same ranking was used in all subsequent representations of origin firing state. To quantify the bias pattern, the log₂ ratios of sequence reads of the Watson strand over the Crick strand surrounding each DNA replication origin were calculated using a 200-bp sliding window. The ratios were then normalized against the corresponding input to obtain the average bias pattern.

Quantification and Statistical Analysis. For peak calling, the FDR was calculated by SICER software (61), and the cutoff value was set to 0.01.

Data Availability. All raw and analyzed sequencing data have been deposited in the National Center for Biotechnology Information under the Gene Expression Omnibus accession number GSE172093 (62).

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