Separate roles for the DNA damage checkpoint protein kinases in stabilizing DNA replication forks

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The DNA damage checkpoint plays a crucial role in maintaining functional DNA replication forks when cells are exposed to genotoxic agents. In budding yeast, the protein kinases Mec1 (ATR) and Rad53 (Chk2) are especially important in this process. How these kinases act to stabilize DNA replication forks is currently unknown but is likely to have important implications for understanding how genomic instability is generated during oncogenesis and how chemotherapies that interfere with DNA replication could be improved. Here we show that the sensitivity of rad53 mutants to DNA-damaging agents can be almost completely suppressed by deletion of the EXO1 gene, which encodes an enigmatic flap endonuclease. Deletion of EXO1 also suppresses DNA replication fork instability in rad53 mutants. Surprisingly, deletion of EXO1 is completely ineffective in suppressing both the sensitivity and replication fork breakdown in mec1 mutants, indicating that Mec1 has a genetically separable role in replication fork stabilization from Rad53. Finally, our analysis indicates that a second downstream effector kinase, Chk1, can stabilize replication forks in the absence of Rad53. These results reveal previously unappreciated complexity in the downstream targets of the checkpoint kinases and provide a framework for elucidating the mechanisms of DNA replication fork stabilization by these kinases.

[Keywords: Checkpoints; Exo1; Rad53; Mec1; Chk1; DNA replication fork stabilization.]

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Eukaryotic cells require specialized surveillance mechanisms called checkpoints to preserve both viability and genome integrity when confronted with endogenous or exogenous agents that interfere with DNA replication. The loss of checkpoint pathways can lead to genomic instability and thereby promote carcinogenesis [Hartwell and Kastan 1994; Kolodner et al. 2002; Nyberg et al. 2002]. The DNA damage checkpoint involves a protein kinase cascade and, in Saccharomyces cerevisiae, the central checkpoint protein kinases are the ATR homolog Mec1 and its downstream effector, the Chk2 homolog Rad53 [Zhou and Elledge 2000]. A second effector kinase, Chk1, has a major role in metazoan checkpoints but a fairly minor role in budding yeast checkpoints. In response to S-phase perturbations, Mec1 is recruited to stalled DNA replication forks, where it is required to phosphorylate and activate Rad53 [Branzei and Foiani 2006]. When activated, checkpoint pathways regulate many aspects of cell metabolism including cell cycle progression, activation of DNA repair pathways, transcription of DNA damage response genes, DNA replication fork stabilization, and DNA replication origin firing [Santocanale and Diffley 1998; Shirahige et al. 1998; Foiani et al. 2000; Lowndes and Murguia 2000; Tercero and Diffley 2001; Nyberg et al. 2002; Andreassen et al. 2006; Branzei and Foiani 2006].

The role of DNA damage checkpoints in replication fork stabilization appears to be especially important. When DNA replication forks in wild-type cells stall because of deoxynucleoside triphosphate (dNTP) depletion with hydroxyurea (HU), they remain competent to resume replication after removal of HU; however, replication forks are unable to resume replication in rad53 mutants after HU removal [Desany et al. 1998; Lopes et al. 2001; Tercero and Diffley 2001; Tercero et al. 2003]. Similarly, replication forks in rad53 or mec1 mutants arrest irreversibly during replication through alkylated DNA leading to incomplete replication and cell death [Tercero and Diffley 2001; Tercero et al. 2003]. This lethality requires passage through S phase but is not prevented by blocking subsequent mitotic entry, indicating that the lethal event is associated with DNA replication and that the role of checkpoints in restraining mitosis cannot account for the lethality [Tercero and Diffley 2001]. Blocking protein synthesis during S phase in wild-type cells does not render them sensitive to HU, nor does it prevent replication fork resumption.
after HU arrest, arguing that checkpoint-dependent induction of transcription is not critical for fork stabilization or viability (Tercero et al. 2003). A hypomorphic mec1 mutant (mec1-100) that cannot block late origin firing in HU but can stabilize replication forks is much less HU-sensitive than mec1Δ cells, arguing that regulation of late origin firing plays a relatively minor role in maintaining cell viability (Tercero et al. 2003). Thus, a process of elimination has pointed to DNA replication fork stabilization as the critical role of Rad53 and Mec1 for cell viability after DNA damage.

How checkpoints regulate replication forks is currently unclear. Chromatin immunoprecipitation (ChIP) experiments have suggested that replisomes remain at stalled forks in wild-type cells but are depleted from stalled forks in checkpoint mutant cells (Cobb et al. 2003; Lucca et al. 2004). Long patches of single-strand DNA accumulate at stalled forks in checkpoint mutants probably because of DNA degradation (Sogo et al. 2002; Feng et al. 2006), consistent with catastrophic breakdown of replisome function. Although there are correlations between replisome stability and checkpoint function, the molecular mechanisms by which checkpoints preserve replication fork function and viability remain to be determined.

The roles of the individual protein kinases in regulating DNA replication forks are still unclear. Because Mec1 is essential for Rad53 activation (Branzei and Foiani 2006) and mec1 and rad53 mutants share similar phenotypes (Lopes et al. 2001; Tercero et al. 2003), it is possible that Rad53 is the main effector at stalled forks and the primary role of Mec1 in fork stabilization is to activate Rad53. However, Mec1 may have roles at replication forks independent of Rad53. mec1-null mutants are considerably more sensitive and have higher rates of replication fork breakdown than rad53-null mutants in HU or MMS (Gardner et al. 1999; Tercero and Diffley 2001). Moreover, ChIP experiments have indicated that mec1 mutants have defects in replisome stability not seen in rad53 mutants (Bierbaek et al. 2005; Cobb et al. 2005), although other studies have suggested that rad53 mutants are also defective in replisome stabilization (Lopes et al. 2001; Sogo et al. 2002; Cotta-Ramusino et al. 2005).

In this study, we describe a genetic approach to examine the role of checkpoints at stalled replication forks. Our work demonstrates that Mec1 and Rad53 have genetically separable roles in fork stabilization. The primary role of Rad53 is to prevent Exo1-dependent replication fork breakdown. Moreover, we describe experiments indicating a previously unappreciated role for Chk1 in fork stabilization during the intra-S checkpoint.

**Results**

**Exo1 suppresses the sensitivity of rad53Δ cells to genotoxic agents**

Because of its critical downstream function in the checkpoint pathway, we initially focused on the role of Rad53 in DNA replication fork stabilization. We considered three different models for how the stability of replication forks might be regulated by activated Rad53 [Fig. 1A], any or all of which might be involved in preventing fork collapse and cell death. Firstly, Rad53 might regulate fork stabilization by direct phosphorylation of some

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**Figure 1.** EXO1 is a suppressor of rad53 lethality in MMS. (A) Models for how Rad53 might stabilize stalled replication forks. See the text for details. (B) Drop assays on YPD + 0.010%, 0.015%, or 0.025% MMS were performed using 1:5 serial dilutions of exponentially growing cultures of the indicated strains. (C) Cell viability was measured as colony outgrowth from exponentially growing cultures on YPD in the presence of 0.015% MMS. (D) The indicated strains were released from G1 arrest at 30°C in YPD medium containing 0.015% MMS, and samples were taken every 10 min. [Left panel] DNA content was measured by flow cytometry. [Right panel] Cell viability for the three strains used in C was scored during α-factor arrest and at the indicated time points during S phase in YPD in the presence of 0.015% MMS.
replisome component. To address this, we are currently surveying the replisome for Rad53-dependent phosphorylation, which will be described elsewhere. Secondly, Rad53 might positively regulate some activity [X] that promotes fork stabilization, either by maintaining the replisome at stalled/damaged forks, reloading replisome components for replication restart, or promoting some replication-coupled DNA repair process. In this scenario, overproduction of “X” might be expected to suppress the sensitivity of rad53 mutants to genotoxic agents. However, we have thus far been unsuccessful in identifying efficient high-copy suppressors of rad53. And thirdly, activated Rad53 might inhibit some activity [Y] that contributes to irreversible fork collapse. In this study, we explore this third option. We reasoned that if the sensitivity of rad53 mutants to DNA replication stress was due, at least in part, to an inability to protect stalled/damaged forks from “Y,” then mutations inactivating “Y” should increase the viability of rad53 mutant cells in the presence of genotoxic agents. We used Saccharomyces cerevisiae to search for mutations that confer upon rad53 cells increased resistance to various forms of DNA replication stress. Details of this screen will be published elsewhere; however, one suppressor mutant stood out from all others. We found that deletion of the EXO1 gene very significantly suppressed lethality of rad53-null mutants treated with the DNA alkylation agent methyl methanesulphonate (MMS).

To characterize this suppression, we examined the effect of MMS on wild-type and rad53Δ mutant strains in the presence or absence of EXO1 using a serial dilution assay [Fig. 1B]. Wild-type cells grew in the presence of 0.010% and 0.015% MMS, whereas rad53Δ mutants were extremely sensitive. Above 0.015% MMS, even wild-type cells were extremely slow growing [Fig. 1B]. Deletion of EXO1 alone conferred mild sensitivity to MMS. Strikingly, deletion of EXO1 in the rad53Δ-null background almost completely suppressed its sensitivity to MMS.

We quantified the ability of these strains to form colonies during chronic exposure to MMS. We plated a fixed number of cells of each strain on plates containing 0.015% MMS and counted surviving colonies, plotted in Figure 1C, as percent survival. Under these conditions, viability of the rad53Δ EXO1+ strain was <0.1% compared with the RAD53Δ EXO1+ strain. In contrast, viability of the rad53Δexox1Δ strain was at least 300-fold higher than the rad53Δ EXO1+ strain and only threefold lower than the RAD53Δ EXO1+ strain. Supplemental Figure 1 shows that deletion of EXO1 also suppressed the MMS sensitivity of strains containing a hypomorphic allele of RAD53 [sad1] (Allen et al. 1994), indicating that suppression is not specific to the RAD53 deletion and does not require deletion of SML1.

Previously, we showed that rad53Δ mutants lose viability during passage through a single S phase in the presence of MMS [Tercero and Diffley 2001]. To examine the effect of EXO1 deletion on viability in a single cell cycle, cells were synchronized in G1 phase with a-factor mating pheromone and released from the G1 arrest in the presence of 0.015% MMS. DNA replication was examined by flow cytometry [Fig. 1D]. As described previously [Paulovich and Hartwell 1995], S phase was rather slow in RAD53+ cells but was considerably faster in rad53Δ cells. This experiment shows that S phase is similarly rapid in the rad53Δexo1Δ double mutant. However, Figure 1D [right panel] shows that the rad53Δ cells lose viability rapidly upon passage through S phase while the rad53Δexo1Δ cells, like RAD53+ cells, maintain high viability. Thus, deletion of EXO1 suppresses most of the loss of viability of rad53Δ cells but does not suppress the accelerated S phase. Previous work has attributed the accelerated S phase to an inability of rad53Δ cells to inhibit late origin firing after DNA damage [Tercero and Diffley 2001]. The results described here suggest that deletion of EXO1 in the rad53Δ background does not restore the block to late origin firing.

We next asked whether deletion of EXO1 could suppress the hypersensitivity of rad53Δ cells to other genotoxic agents including ultraviolet radiation (UV), ionizing radiation [IR], and hydroxyurea [HU]. Figure 2 shows that the rad53Δ strain was more sensitive than the RAD53+ strain to all of the agents tested. After treatment with UV or IR, the viability of the rad53Δexo1Δ strain was considerably higher than the rad53Δ EXO1+ strain, almost as high as the RAD53+ strain [Fig. 2]. Therefore, the sensitivity of rad53Δ cells to a wide range of DNA-damaging agents requires the presence of EXO1. In contrast to the DNA-damaging agents, deletion of EXO1 did not result in a significant increase in the viability of rad53Δ cells treated with HU.

Exo1 is a relatively large protein [702-amino-acid residues] with a nuclease domain found in the N-terminal one-third of the protein. To test if the Exo1-dependent lethality in rad53Δ cells can be attributed to the nuclease activity of Exo1, we constructed strains expressing either wild-type EXO1 or one of two nuclease-deficient versions of EXO1 with mutations in the essential residues D173 and E150, respectively [Fig. 3A]. It has been previously shown that changing either of these two resi-

Figure 2. EXO1-dependent suppression occurs in MMS, UV, and IR, but not in HU. Serial dilutions [1:5] of exponential cultures of RAD53Δ EXO1+, rad53Δ EXO1+, or rad53Δexo1Δ strains were tested for sensitivity to increasing dosages of MMS, UV, IR, or HU [panels from top to bottom], by using a drop assay on plates.
duplicates lethality in rad53 mutants in MMS. Either wild-type or mutated Exo1 versions were integrated in the genome of a rad53 strain generating the following strains: rad53Δexo1Δ strain generating the following strains: rad53Δexo1Δ → rad53Δexo1Δ. Exponentially growing cultures of the indicated strains were serially diluted (1:5), and 4 µL of each dilution was spotted onto YPD and YPD + 0.015% MMS plates. RAD53+/EXO1+ and rad53Δexo1Δ strains were also included as controls. (C) Exo1-dependent lethality does not depend on interaction with Msh2 and Mlh1. The indicated strains were grown exponentially, serially diluted (1:5), and spotted onto YPD + 0.015% MMS. (D) Exo1-dependent lethality is not mediated by homologous recombination. The indicated strains were grown exponentially, serially diluted (1:5), and spotted onto YPD + 0.010% MMS.

Rescue of replication fork breakdown by EXO1 deletion

We showed previously that DNA replication forks in both RAD53+ and rad53Δ cells proceed very slowly when cells replicate in the presence of MMS [Tercero and Diffley 2001]. Whereas wild-type cells ultimately complete DNA replication, rad53Δ cells were unable to finish. This inability to complete DNA replication is correlated with the S-phase-specific loss of viability [Tercero and Diffley 2001; Tercero et al. 2003]. Given the strong correlation between loss of viability and inability to complete DNA replication, we were interested in determining whether deletion of EXO1 would allow rad53Δ cells to complete DNA replication in MMS.

To investigate this, we used density transfer substitution experiments to quantify replication fork progression [Reynolds et al. 1989; Tercero et al. 2000; Tercero and Diffley 2001]. Briefly, cells were grown for at least seven generations in the presence of 12C glucose and 15N ammonium sulfate, ensuring that both parental DNA strands were fully substituted with heavy isotopes [heavy-heavy; HH]. After synchronization in G1 with α-factor mating pheromone, cells were allowed to pass through S phase in the presence of MMS and light isotopes [14C glucose, 14N ammonium sulfate], which results in the generation of Heavy-Light [HL] DNA. As shown by flow cytometry in Figure 4A, wild-type cells proceeded slowly through S phase in the presence of MMS, whereas rad53Δ cells proceeded much faster. As indicated above, EXO1 deletion did not detectably alter the replication kinetics of the rad53 mutant.

DNA preparations from the indicated time points were digested with the appropriate restriction enzyme, and HH-DNA and HL-DNA were separated based on their different density in CsCl gradients. Replication of each fragment of DNA was prelabeled with 12C glucose, 14N ammonium sulfate, ensuring that both parental DNA strands were fully substituted with heavy isotopes. The indicated fragments were then subcloned into S. cerevisiae plasmids for analysis. Exo1 is a critical DNA recombination factor that deletion of another mismatch repair factor, MLH1, did not suppress the sensitivity of rad53Δ cells to MMS. Figure 3D shows that deletion of RAD52, a critical DNA recombination factor, was also completely ineffective in suppressing the sensitivity of rad53Δ cells to MMS. Therefore, the suppression of the hypersensitivity of rad53Δ cells to genotoxic agents is not due to the loss of mismatch repair or recombination.

Figure 3. EXO1-dependent lethality is linked to its nuclease activity. [A] Schematic of S. cerevisiae Exo1 with functional nuclease domains and Msh2/Mlh2-binding domains are indicated by gray, black, and striped boxes, respectively. Asterisks highlight the highly conserved Asp 173 or Glu 150 residues within the I-nuclease domain. (B) Nuclease-deficient Exo1 prevents lethality in rad53 mutants in MMS. Either wild-type or mutated Exo1 versions were integrated in the genome of a rad53Δexo1Δ strain generating the following strains: rad53Δexo1Δ → rad53Δexo1Δ. Exponentially growing cultures of the indicated strains were serially diluted (1:5), and 4 µL of each dilution was spotted onto YPD and YPD + 0.015% MMS plates. RAD53+/EXO1+ and rad53Δexo1Δ strains were also included as controls. (C) Exo1-dependent lethality does not depend on interaction with Msh2 and Mlh1. The indicated strains were grown exponentially, serially diluted (1:5), and spotted onto YPD + 0.015% MMS. (D) Exo1-dependent lethality is not mediated by homologous recombination. The indicated strains were grown exponentially, serially diluted (1:5), and spotted onto YPD + 0.010% MMS.
Replication from ARS607 had already begun by 30 min after release from G1 arrest. DNA replication in this experiment can be seen to proceed from left to right across the entire replicon because of the sequential transfer of fragments from HH to HL from left to right along the replicon with time. As previously noted, the origin associated with the X element at the end of the chromosome (Chan and Tye 1983) appears to remain inactive because the entire replicon is primarily replicated from left to right. Replication was very slow, but by 240 min, the entire region was completely replicated in almost all cells, as previously shown for a checkpoint-proficient strain in the presence of MMS (Tercero and Diffley 2001). There were no significant differences in replication fork progression between \( \text{RAD53}^+ \text{EXO1}^+ \) cells and \( \text{RAD53}^+ \text{exo1}/H9004 \) cells (Supplemental Fig. 2).

In \( \text{rad53}/H9004 \) mutants (Fig. 4C), the beginning of S phase looked very similar to the pattern shown in Figure 4B. Activation of ARS607 occurred during the first 30 min, and forks proceeded slowly into the adjacent fragments. By 60 min, however, replication forks had also originated from the vicinity of the chromosome end and proceeded from right to left. This can be seen as replication of the +65-kb fragment before the +50-kb fragment and is essentially identical to our previous experiments, consistent with the loss of Rad53-dependent inhibition of late origin firing after DNA damage. Most importantly, as we previously showed, replication forks arrest irreversibly in this mutant. There is no further DNA synthesis detectable after 120–150 min, although significant amounts of DNA persist in the HH peak for the remainder of the experiment, especially in the origin-distal fragments at +20 kb and +40 kb. We call this “terminally unreplicated DNA.”

The \( \text{rad53}/H9004 \text{exo1}/H9004 \) strain also showed an accelerated S phase, apparently due to inappropriate initiation events.
near the terminus of the chromosome similar to the rad53ΔEXO1+ strain. However, in contrast to the rad53ΔEXO1+ strain, the rad53Δexo1Δ strain completed DNA replication as efficiently as a checkpoint proficient strain as shown by the complete transfer of DNA from the HH to the HL peak [Fig. 4D]. Figure 4E shows a quantification of this terminally unreplicated DNA at various positions along the replicon in the rad53ΔEXO1+ and rad53Δexo1Δ strains. These experiments show that the deletion of EXO1 prevents the irreversible arrest of DNA replication forks in rad53Δ cells treated with MMS.

rad53 mutant cells are unable to restart stalled DNA replication forks following transient arrest in HU [Pau-lovich and Hartwell 1995; Desany et al. 1998; Lopes et al. 2001]. It has also been shown that Exo1 specifically affects the stability of replication forks in rad53 mutant cells in HU [Cotta-Ramusino et al. 2005]. However, as shown in Figure 2, EXO1 deletion does not rescue the sensitivity of rad53Δ cells to HU. Thus, we wondered whether deletion of EXO1 would allow replication forks to restart following HU removal.

To examine this, we analyzed replication resumption after HU arrest by measuring DNA content using flow cytometry. The indicated strains were blocked in G1 with α-factor and released into medium containing 0.2 M HU. After 2 h, cells were transferred to medium lacking HU but containing nocodazole to prevent passage through mitosis. As shown in Figure 5A, RAD53ΔEXO1+ cells completed replication 40 min after release form HU, whereas rad53ΔEXO1+ cells were unable to resume significant amounts of DNA synthesis even after 90 min. The replication profile in rad53Δexo1Δ cells showed no significant differences from rad53ΔEXO1+ cells, suggesting that both strains are unable to restart DNA replication from stalled forks after HU treatment.

We next used bromodeoxyuridine [BrdU] pulse-chase experiments to label nascent strands during replication in HU, which allowed us to follow the fate of stalled/damaged replication forks after HU removal in a more
direct way. We used strains previously developed in our laboratory that can incorporate BrdU into DNA when present at low concentrations in the medium. These strains express both a nucleoside transporter (hENT) and a thymidine kinase from Herpes Simplex virus (HSV TK) [Vernis et al. 2003], allowing us to perform “pulse-chase” experiments. Cells were synchronized in G1 with α-factor and released into medium containing HU and BrdU. After labeling the nascent DNA associated with the stalled forks, the BrdU was chased by transferring cells to fresh medium lacking HU and BrdU and containing a high concentration of thymidine. DNA was purified, separated on a denaturing alkaline agarose gel, and transferred to a nylon membrane, and the newly replicated DNA was detected with an anti-BrdU antibody as described previously (Vernis et al. 2003).

Nascent DNA replication intermediates labeled in all of the strains at 25°C appeared as a smear [Fig. 5B], consistent with similar experiments in which nascent DNA was detected by blot hybridization using origin-specific probes [Santocanale and Diffley 1998]. After release, the size of the fragments in RAD53‘EXO1‘ cells increased very quickly, and by 60 min almost all the incorporated BrdU had been chased into the high-molecular-weight fraction, indicating rapid restarting of stalled replication forks. In contrast, in rad53ΔEXO1‘ cells, the majority of the nascent DNA remained at the same position after release, indicating a gross failure of replication forks to resume DNA replication. This pattern was indistinguishable from the one obtained in rad53Δexo1Δmutants, indicating that the inability of rad53Δ cells to restart fork progression is not rescued by EXO1 deletion.

We wanted to determine if inappropriate entry into mitosis in HU caused by checkpoint loss compromises the suppressor effect of EXO1 deletion. To this end, we measured survival rates in RAD53‘EXO1‘, rad53ΔEXO1Δ, and rad53Δexo1Δ strains during HU treatment and after release, in the presence or absence of nocodazole [Fig. 5C]. There was a very slight increase in viability in both rad53ΔEXO1‘ and rad53Δexo1Δ strains in the presence of nocodazole [from 3% to 6%], suggesting that in checkpoint mutants a very small proportion of cells in the population may die because of premature entry into mitosis. However, viability remains below 10% in both mutant strains, indicating that blocking entry into mitosis is not sufficient to prevent lethality in rad53Δexo1Δ cells in HU or after HU removal.

**EXO1 deletion does not rescue a mec1Δ mutant**

As described in the introduction, the role of Mec1 in DNA replication fork stabilization is presently unclear. Mec1 may only be required because of its role in Rad53 activation or it may have an additional, separate role. We reasoned that if the only function of Mec1 is to activate Rad53, EXO1 deletion should also suppress the sensitivity of mec1Δ mutants to genotoxic stress.

We therefore compared sensitivity of mec1ΔEXO1‘ mutants to mec1Δexo1Δ cells in MMS. Figure 6A shows that mec1Δ cells are considerably more sensitive to

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**Figure 6.** **EXO1 deletion does not rescue viability or replication fork breakdown in mec1 mutants.** (A) The specified strains were grown exponentially, serially diluted 1:5, and spotted onto YPD plates containing either 0.001%, 0.002%, or 0.010% MMS. (B) An equal number of cells for each strain were plated for viability onto YPD + 0.008% MMS plates, and numbers of colonies were scored and plotted on the graph. (C) Replication fork progression in a mec1ΔEXO1‘ strain. (D) Replication fork progression in a mec1Δexo1Δ strain. A time course of DNA replication in mec1ΔEXO1‘ and mec1Δexo1Δ strains was analyzed after release from α-factor arrest into medium with 0.033% MMS [Supplemental Fig. 3A]. Time points corresponding to 120 min and 240 min after release from G1 block were analyzed by dense-isotope transfer, as described in Figure 4. The strains used in C and D harbor deletions of SML1 to allow viability in the absence of MEC1. (E) Replication fork breakdown rates. The graph shows the percentage of unreplicated DNA remaining at 240 min in MMS in the mec1ΔEXO1‘ and mec1Δexo1Δ strains (last panel in C and D) for each fragment. Black bars represent the mec1ΔEXO1‘ strain and gray bars represent the mec1Δexo1Δ strain.
Figure 7. **EXO1** deletion does not rescue viability or replication fork breakdown in rad53Δchk1Δ mutants. **A** The viability of the indicated strains in YPD + 0.010% MMS was measured, by performing a drop assay with a 1:5 serial dilution series of exponential cultures of each strain. **B** Sensitivity to MMS for the rad53Δ, mec1Δ, and rad53Δchk1Δ strains in the presence and absence of EXO1 was compared. The indicated strains were grown exponentially, serially diluted (1:5), and spotted onto YPD plates containing either 0.006% or 0.010% MMS. **C** Sensitivity to MMS for the rad53Δ, mec1Δ, and rad53Δchk1Δ strains in the presence and absence of EXO1 was measured, by performing a drop assay with a 1:5 serial dilution series of exponential cultures of each strain. **D** Sensitivity to MMS for the rad53Δ, mec1Δ, and rad53Δchk1Δ strains in the presence and absence of EXO1 was measured, by performing a drop assay with a 1:5 serial dilution series of exponential cultures of each strain. **E** Sensitivity to MMS for the rad53Δ, mec1Δ, and rad53Δchk1Δ strains in the presence and absence of EXO1 was measured, by performing a drop assay with a 1:5 serial dilution series of exponential cultures of each strain.
ing either strain from cell lethality. Moreover, with or without nocodazole, the rad53Δchk1Δexo1Δ strain was slightly more sensitive to MMS than the rad53Δstrain. Taken together, these results suggest that Chk1 has a role in maintaining functional DNA replication forks, at least in the absence of Rad53, independent of its role in restraining mitosis.

Discussion

We argued previously that the most important function of Rad53 in promoting cell survival after DNA damage is to preserve the integrity of DNA replication forks (Tercero and Diffley 2001; Tercero et al. 2003). In this study, we provide additional support for this idea, and we show that the nuclease Exo1 plays a critical, antagonistic role in this process. Our results also revealed Rad53-independent roles for the Mec1 and Chk1 checkpoint kinases in DNA replication fork stabilization, leading us to suggest that the simple model in Figure 8A is incomplete. Instead, we propose the model shown in Figure 8B.

We show here that the loss of DNA replication fork stability in rad53Δ mutants treated with DNA-damaging agents requires Exo1 and that removal of Exo1 is sufficient to allow near wild-type levels of fork stabilization in the absence of Rad53. Previous work from Foiani and colleagues has shown that the collapse of DNA replication forks in HU-treated cells requires Exo1 (Cotta-Ramusino et al. 2005). They showed using 2D gels that replication bubbles degenerate into abnormal structures in HU-treated rad53 mutant cells and that this phenotype is dependent on Exo1. However, they noted that the absence of Exo1 did not increase the viability of rad53 mutants. Our results are consistent with this, we also observed that EXO1 deletion does not suppress rad53 lethality in HU, nor did it promote replication restart. We suggest that Rad53 stabilizes DNA replication forks via an Exo1-dependent and an Exo1-independent pathway, both of which are required to maintain functional replication forks in HU. The Exo1-independent pathway is not simply Rad53-dependent restraint of mitotic entry because the lethality of HU in rad53Δexo1Δ cells is not suppressed by nocodazole. Our results show that, in MMS, UV, and IR, Rad53 works primarily via the Exo1-dependent pathway. Previous work has shown that the hypersensitivity of rad53 mutants to MMS is almost entirely due to problems during DNA replication. The results presented here suggest that at least some of the hypersensitivity of rad53 mutants to IR and UV may also be due to difficulties during DNA replication.

We presently do not know the mechanism by which Exo1 interferes with the integrity of the replication forks. It may be that Rad53 controls the stability of some replisome component at the fork. Loss of this component in the absence of Rad53 would simply expose the replication fork to degradation by Exo1. Alternatively, Exo1 may be directly regulated by Rad53. We favor this possibility for several reasons. Firstly, it would help explain specificity. Exo1 is just one of many Rad2 family S’-flap endonucleases [Lieber 1997] and appears to function redundantly with other nucleases in different biological processes. For example, Exo1 exhibits functional redundancy with Rad27 [Fen1] in Okazaki fragment processing [Tishkoff et al. 1997; Tran et al. 2002] and with Mre11 for resection of mitotic DSBs [Tsubouchi and Ogawa 2000; Moreau et al. 2001]. Yet, we did not find a comparable rad53 suppressor effect by deleting other nucleases [M. Segurado and J.F.X. Diffley, unpubl.]. Secondly, the generation of ssDNA at dysfunctional telomeres by Exo1 has been shown to be negatively regulated by Mec1 and Rad53 [Jia et al. 2004]. And finally, quantitative phosphoproteomic approaches in yeast have shown that Exo1 is phosphorylated in vivo in a Rad53-dependent manner in response to MMS treatment [Smolka et al. 2007]. We are currently analyzing Rad53-dependent phosphorylation of Exo1, and further work is required to determine the effect of that phosphorylation on the regulation of the protein and its influence on the replication forks.

In addition to its role in Rad53 activation, our results indicate that Mec1 has a distinct role in fork stabilization. The sensitivity of mec1Δ cells to MMS is not suppressed at all by EXO1 deletion. This could be because Mec1 is also required for Chk1 activation, however, mec1 mutants are considerably more sensitive to MMS than rad53Δchk1Δ double mutants, indicating that Mec1 has a role not accounted for by Rad53 and Chk1 together. The nature of this role at forks is unclear. ChIP experiments have indicated that checkpoint mutants have defects in maintaining occupancy of replisome components at stalled replication forks [Cobb et al. 2003; Lucca et al. 2005; Tercero et al. 2003].

Figure 8. Mec1/Rad53/Chk1 are important to maintain fork integrity. [A] This model summarizes the current view of checkpoint-dependent fork stabilization. The replication forks activate Mec1, the sensor kinase, which then phosphorylates the downstream effector Rad53 kinase. Rad53 acts on the replication forks and stabilize them, whereas the main role of Chk1 is to prevent entry into mitosis. Rad53 has also been proposed to contribute to inhibition of mitosis. [B] This model summarizes the different pathways implicated in the stability of the replication forks based on our results. Mec1, Rad53, and Chk1 kinases all contribute to fork stabilization. Rad53 works through inhibition of Exo1 and also has additional roles at the replication forks. Mec1 has a Rad53-dependent role at the replication forks, and Chk1 is also involved in fork stabilization in the absence of Rad53. Rad53 and Chk1 both contribute to inhibition of mitosis. Black lines and arrows indicated the additional pathways compared with the model represented in A.
et al. 2004; Cotta-Ramusino et al. 2005), although there are conflicting results regarding the contribution of Mec1 and Rad53 kinases to this stabilization. Cotta-Ramusino et al. (2005) have proposed that Rad53 is the critical kinase because rad53 mutants lose DNA polymerases from stalled forks. However, Cobb et al. (2005) have proposed that Mec1 is important to prevent polymerases–replisome disassociation, but that Rad53 is dispensable. Our results do not resolve this discrepancy. We clearly show genetically separable roles for Mec1 and Rad53 in fork stabilization, at least in MMS. However, our data also demonstrate that rad53Δ cells show irreversible fork breakdown in HU that is not suppressed by deletion of EXO1. Identification of functionally relevant Mec1, Rad53, and Chk1 substrates will be required for further progress in this area. Recent phosphoproteomic analysis looking for ATR and ATM targets in humans cells in response to DNA damage (Matsuoka et al. 2007) has identified several essential replisome proteins including Mcm2–7 subunits, RFC clamp-loader components, and DNA polymerases. Mcm2–7 proteins are especially interesting candidates because loss of Mcm2–7 from stalled forks causes an irreversible arrest similar to that seen in checkpoint mutants (Labib et al. 2000).

In addition to the roles of Rad53 and Mec1, the fact that the Exo1-dependent suppression in rad53 mutants requires Chk1 implies that this kinase can also regulate replisome stability, at least in the absence of Rad53. This result is surprising due to the fact that chk1 mutants are relatively insensitive to HU and MMS (Fig. 7A; Sanchez et al. 1999). Our results indicate that, in the absence of Rad53, Chk1 can prevent irreversible DNA replication fork breakdown. Chk1, however, can only accomplish this in the absence of Exo1. This suggests that Rad53 and Chk1 may act redundantly to promote some aspect of replisome stability or replication restart but only Rad53 can counteract the negative effect of Exo1. The fact that deletion of EXO1 in the rad53Δchk1Δ strain significantly increases sensitivity to MMS suggests that Exo1 plays a positive role in survival in this background. Clearly, more work is required to determine the role of Chk1 at replication forks.

In Schizosaccharomyces pombe, the Rad53 and Chk1 homologs [Cds1 and Chk1, respectively] [Walworth and Bernards 1996; Lindsay et al. 1998] may have similar overlapping functions. Like Rad53, Cds1 kinase is required for cell survival when replication is inhibited by HU or when DNA is damaged during S phase. Chk1 is activated after DNA damage most commonly during late S and G2 phase. However, when cells are treated with HU in the absence of Cds1, the checkpoint remains intact and arrest becomes dependent on Chk1 [Martinho et al. 1998; Brondello et al. 1999; O’Connell et al. 2000]. Although there may be differences in the functions of the individual checkpoint proteins kinases in different organisms [Rhind and Russell 2000], it is possible that this redundancy may represent a conserved backup system for the Rad53 checkpoint. Regardless, this is the first time that Chk1 has been shown to have a role in the stabilization of replication forks in budding yeast.

Exo1 homologs have been identified in other eukaryotes, including yeast, flies, and mammalian cells (Szankasi and Smith 1992, 1995; Digilio et al. 1996; Fiorentini et al. 1997; Tishkoff et al. 1998). Further work will be required to determine if the deleterious effect of Exo1 on replication fork stability and cell viability has been conserved in evolution.

Materials and methods

Strains, plasmids, and media

All strains used are listed in Supplemental Table S1 and are derived from W303-1a [MATa ade2-1 ura3-1 his3-11, 15 trpl-1 leu2-3, 112 can1-100].

pRS304-EXO1 was created as follows. The EXO1 3-kb genomic fragment was amplified by PCR using Phusion DNA polymerase with the oligos 5′-CAACATCACGTTCTGTGC-3′ and 5′-GTATGATCCGATGATTGACC-3′, and cloned into pRS304 vector. The exo1-D173A allele from Ylp-exo1-D173A [Tran et al. 2002] was used to replace the wild-type Spel–BamHI fragment of pRS304-EXO1 to create pRS304-exo1-D173A. pRS304-exo1-E150D was created by replacing the Spel–BamHI fragment of pRS304-EXO1 with the exo1-E150D allele from Ylp-exo1-D173A [Tran et al. 2002]. The three constructs were sequenced with the oligo EXO1-752.S (5′-GAACCTGATTTGGTCTTCCGATG-3′) [Tran et al. 2002], to verify correct substitution of the selected fragments. The exo1-point mutant strains were created by targeting MfeI-digested constructs pRS304-exo1-D173A and pRS304-exo1-E150D into the TRP locus of a rad53Δexo1ΔΔ strain.

Unless otherwise indicated, cells were grown at 30°C in YP medium [1% yeast extract [Difco], 2% bacto peptone [Difco]] supplemented with 2% glucose (YPD).

Drop assays and viability

Drop assays were a 1:5 dilution series of exponentially growing cultures on YPD, YPD + MMS, or YPD + HU plates depending on the experiment. In the panels shown in the second and third rows of Figure 2, cells were spotted onto YPD plates and irradiated with ultraviolet or ionizing radiation, respectively, at the indicated dosages.

The viability of asynchronous cultures was calculated by plating 10⁶ cells for wild-type and 10⁷ cells for checkpoint mutants in duplicate onto YPD + 0.015% MMS plates [Fig. 1C] or YPD + 0.008% MMS [Fig. 6B] and scoring after 3 d at 30°C.

The viability of synchronous cultures was calculated by plating 10⁵ cells for wild-type and 10⁶ cells for checkpoint mutants in duplicate onto YPD plates and scoring after 3 d at 30°C.

Cell cycle synchronization and flow cytometry

Cell growth and cell cycle blocks with α-factor, HU, and nocodazole were as described previously [Difflley et al. 1994; Donovan et al. 1997]. Samples for flow-cytometric analysis [FACS] were collected and processed as described previously [Labib et al. 1999].

DNA replication analysis assays

Density transfer were performed essentially as described [Tercero et al. 2000]. DNA was digested with ClaI and SalI before gradient centrifugation in cesium chloride. DNA probes for slot blot hybridization were amplified by polymerase chain reaction.
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(PCR). Probes corresponding to the six fragments were as follows: probe 1 [ARS607], nucleotides 198,945 ± 199,832, probe 2 [1±10 kb], nucleotides 211,014 ± 211,996; probe 3 [±20 kb], nucleotides 218,011 ± 218,700; probe 4 [±40 kb], nucleotides 240,089 ± 240,679; probe 5 [±50 kb], nucleotides 243,315 ± 244,200; and probe 6 [±65 kb], nucleotides 260,048 ± 261,088. The molecular Dynamics PhosphorImager was used to detect the hybridization signals.

The analysis of the data was performed basically as described in http://fangman-brewer.genetics.washington.edu/density_transfer.html with small differences. Briefly, the hybridization signals were quantified and plotted for each time point by using Image Quant TL software. The resulting graphs are shown in Figures 4, 6, and 7 for the indicated strains. For the calculation of the replication fork breakdown rates shown in Figures 4E, 6E, and 7E, the areas of the HH and HL peaks from each graph at the 240-min time point were measured and quantified with the ImageJ software. The percentage of unreplicated DNA at each position was calculated using the equation:

\[
\text{% Unreplicated DNA} = \frac{HH + HL}{HH + HL} \times 100
\]

This equation takes into account the fact that the HL peak is overrepresented due to the nature of semiconservative DNA replication and so the values obtained from the HL peak should be divided by half to calculate the total signal obtained from replicated plus unreplicated DNA.

Pulse-chase BrdU experiments and immunodetection of BrdU-labeled DNA were performed as described (Vernis et al. 2003).

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