ATR phosphorylates SMARCAL1 to prevent replication fork collapse

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The DNA damage response kinase ataxia telangiectasia and Rad3-related (ATR) coordinates much of the cellular response to replication stress. The exact mechanisms by which ATR regulates DNA synthesis in conditions of replication stress are largely unknown, but this activity is critical for the viability and proliferation of cancer cells, making ATR a potential therapeutic target. Here we use selective ATR inhibitors to demonstrate that acute inhibition of ATR kinase activity yields rapid cell lethality, disrupts the timing of replication initiation, slows replication elongation, and induces fork collapse. We define the mechanism of this fork collapse, which includes SLX4-dependent cleavage yielding double-strand breaks and CtIP-dependent resection generating excess single-stranded template and nascent DNA strands. Our data suggest that the DNA substrates of these nucleases are generated at least in part by the SMARCAL1 DNA translocase. Properly regulated SMARCAL1 promotes stalled fork repair and restart; however, unregulated SMARCAL1 contributes to fork collapse when ATR is inactivated in both mammalian and Xenopus systems. ATR phosphorylates SMARCAL1 on S652, thereby limiting its fork regression activities and preventing aberrant fork processing. Thus, phosphorylation of SMARCAL1 is one mechanism by which ATR prevents fork collapse, promotes the completion of DNA replication, and maintains genome integrity.

Keywords: ATR, SMARCAL1; DNA damage response; DNA replication; cell cycle checkpoint; HARP

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Every cell division cycle, cells must accurately and completely replicate their genome. Errors in replication result in mutations and chromosomal rearrangements that contribute to tumorigenesis. Replication stress caused by DNA lesions, insufficient nucleotides, or even collisions of replication and transcriptional machineries increases the chance of errors. However, replication stress also activates a DNA damage response [DDR] that slows cell cycle progression and promotes DNA repair to ensure accurate duplication of the genome.

The DDR kinase ataxia telangiectasia and Rad3-related (ATR), a member of the phosphoinositol 3-kinase-like kinase (PIKK) family, coordinates much of the cellular response to replication stress [Cimprich and Cortez 2008; Nam and Cortez 2011]. ATR is activated upon replication fork stalling and uncoupling of the replicative helicase and polymerase [Byun et al. 2005]. Once activated, ATR phosphorylates hundreds of substrates to induce the replication checkpoint and promote fork repair.

ATR is essential for cell viability, and hypomorphic ATR mutations cause the rare disease Seckel syndrome, characterized by growth retardation, microcephaly, and other developmental problems [O'Driscoll et al. 2003]. ATR is thought to be a good drug target for cancer therapy because its function is especially critical in replicating tumor cells, which have elevated levels of replication stress due to activated oncogenes and frequent loss of the G1 checkpoint [Reaper et al. 2011; Toledo et al. 2011b; Schoppy et al. 2012]. The mechanism by which ATR-selective inhibitors kill cells is unknown but is likely linked to the replication fork stabilization and repair activities of ATR instead of its G2 checkpoint function [Nam et al. 2011; Toledo et al. 2011a]. Defining these mechanisms is important for the development of ATR pathway inhibitors for cancer treatment.

Replication fork repair is a complex process that can proceed through multiple pathways depending on the cause, persistence, and genomic context of the replication stress. These mechanisms include fork stabilization to

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allow completion of replication by a converging replication fork, lesion bypass, template switching through recombination or fork reversal, and double-strand break (DSB)–mediated restart (Branzei and Foiani 2010). Many enzymes participate in these activities, including helicases, DNA topoisomerases, nucleases, and specialized polymerases. ATR can phosphorylate many of these enzymes; however, the mechanisms by which it promotes fork stabilization and repair and cell viability remain largely unknown.

One ATR substrate that acts at stalled forks is SMARCAL1 (also known as HARP) (Bansbach et al. 2009; Postow et al. 2009). SMARCAL1 binds branched DNA structures and can catalyze DNA annealing, branch migration, fork regression, and fork restoration (Yusufzai and Kadonaga 2008; Betous et al. 2012, 2013; Ciccia et al. 2012). SMARCAL1 is recruited to stalled forks through an interaction with replication protein A (RPA) (Bansbach et al. 2009; Ciccia et al. 2009; Yuan et al. 2009; Yusufzai et al. 2009), which directs it to regress stalled forks with a leading strand gap and restore a normal fork structure (Betous et al. 2013). Both overexpression and siRNA silencing of SMARCAL1 cause replication-associated DNA damage (Bansbach et al. 2009). Furthermore, loss-of-function mutations in SMARCAL1 cause the human disease Schimke immunosseous dysplasia, which is characterized by growth defects, renal failure, immune deficiencies, and predisposition to cancer (Boerkoel et al. 2002; Baradaran-Heravi et al. 2012; Carroll et al. 2013). How ATR phosphorylation of SMARCAL1 regulates its genome maintenance functions at a damaged fork has not been investigated.

Using a selective ATR inhibitor (ATRi), we demonstrate that acute inhibition of ATR kinase activity perturbs the timing of replication initiation, impairs fork elongation rates, and causes rapid lethality in S-phase cells experiencing replication stress. Stalled forks collapse when ATR is inhibited due to SLX4-dependent endonuclease cleavage, which yields DSBs and the CtIP-dependent appearance of single-stranded template and nascent DNA strands. Excessive SMARCAL1 activity is partly responsible for this aberrant fork processing. ATR phosphorylation of a conserved SMARCAL1 serine regulates SMARCAL1 and is one mechanism by which ATR maintains genome integrity during DNA replication. Thus, our results provide a mechanistic description of fork collapse in mammalian cells and define a specific enzymatic pathway responsible for this collapse. They also explain why both too much and too little SMARCAL1 causes replication-associated DNA damage, emphasizing the need to properly regulate this replication fork repair enzyme. Finally, these data provide insights into the mechanism of action of ATRis that are being developed to treat cancers with high levels of dependency on this replication stress response pathway.

Results

Acute ATR inhibition causes rapid cell death in cells experiencing replication stress

Conditional deletion of ATR in dividing mouse or human cells causes cell death as the mRNA and protein levels decay over time (Brown and Baltimore 2000; de Klein et al. 2000; Cortez et al. 2001). The gradual nature of these genetic loss-of-function experiments precludes an analysis of the immediate effects of ATR deficiency. To overcome this technical challenge, we used a selective ATR kinase inhibitor to examine how cells respond to acute and transient ATR inhibition. In this analysis, 39% of asynchronous cultured U2OS cells were no longer able to form colonies after a 10-h treatment with a concentration of ATRi sufficient to block phosphorylation of CHK1, and 71% of cells were inviable after 20 h of treatment (Fig. 1A).

Acute treatment with the combination of ATRi and replication stress induced by the ribonucleotide reductase inhibitor hydroxyurea (HU) caused a greater reduction in
viability than either drug alone [Fig. 1A,B]. Notably, 5 h of treatment with HU and ATRi caused a 55% reduction in viability. This percentage corresponds approximately to the proportion of cells in S phase at the onset of treatment plus the cells that enter S phase during the 5-h treatment. ATR inhibition is synergistic with HU: At the 20-h time point, the combination of HU+ATRi gave an experimental response (99% decrease in viability) greater than the additive effect predicted from HU alone and ATRi alone (89%).

To determine how rapidly the cell lethality occurs, we treated U2OS cells with or without HU and ATRi for up to 5 h. While HU alone or ATRi alone had only modest effects at earlier time points, the combination of HU and ATRi caused a 30% decrease in colony-forming ability within 45 min of treatment [Fig. 1C,D]. Since HU treatment is expected to affect primarily S-phase cells, we tested whether these cells could recover from this treatment and complete DNA replication. By flow cytometry, at least 20% of cells treated with HU and ATRi for 5 h failed to complete DNA synthesis 24 h after release into fresh growth medium compared with only 7% of controls (Fig. 1E). After a 16-h HU treatment, this fraction of lagging cells increased to at least 65% of ATR-inhibited cells compared with only 15% of controls [Fig. 1F]. Thus, ATR is required during an acute replication stress challenge to ensure that cells can recover, complete DNA synthesis, and retain long-term viability.

**ATR inhibition deregulates replication timing control and causes replication fork collapse**

To understand the cause of the rapid lethality when ATR is inhibited, we used fiber labeling to examine DNA replication. Cells were labeled with IdU [green] for 20 min and then with ClidU for 20 min in the presence of DMSO (red bars) or 5 mM ATRi [blue bars] during both labeling periods before harvesting for fiber labeling. [A] Representative replication tracks and quantification of the length of ClidU (red) tracks and newly synthesized DNA strands after fork stalling, [B] Origin initiation was scored as the percentage of red-only tracks. [C–F] RPE-hTERT cells were labeled with IdU for 20 min, treated with 2 mM HU for 2 h in the presence of DMSO (red bars) or 5 mM ATRi [blue bars], and then labeled with ClidU for 20 min before harvesting for fiber staining. Representative images and quantification of ClidU [C] and IdU [F] track lengths in dual-labeled fibers are shown. Percentage of collapsed forks [green-only tracks] [D] and newly initiated origins [red-only tracks] [E] were quantitated. In all experiments, data was collected from several experimental samples with high-quality DNA fibers. Error bars are SEM.

To examine what is happening to the stalled forks when ATR is inhibited, we first examined the levels of DNA damage in ATRi-treated cells. H2AX phosphorylation (γH2AX) provides an indirect marker for DNA damage, since it is phosphorylated at stalled forks and at DSBs. As expected, HU treatment alone causes low but detectable levels of γH2AX, which increase over time [Fig. 3A,B]. In the presence of ATRi, γH2AX is absent at the earliest time points of HU treatment, indicating that ATR is required to phosphorylate H2AX near stalled replication forks, consistent with previous observations [Sirbu et al. 2011]. However, by 1 h in HU, the γH2AX in ATRi-inhibited cells exceeds that in controls. At this time point, a few cells display large, poorly defined γH2AX foci, but most contain a pan-nuclear staining pattern without discernible foci, suggesting rapid spreading through the chromatin. These data suggest the activation of the DSB-sensing kinases ATM and DNA-PKcs between 30 and 60 min of HU treatment when ATR is inhibited. Indeed,
we detected elevated RPA32 S4/S8 phosphorylation [a DNA-PK-dependent phosphorylation site] and CHK2 T68 phosphorylation [an ATM-dependent phosphorylation site] in ATR-inhibited cells compared with controls with kinetics similar to γH2AX [Fig. 3C]. Thus, a DSB may form at the stalled fork, generating the ATM- and DNA-PKcs-activating signal.

To provide direct evidence of DSB formation, we used a neutral COMET assay. The combination of HU and ATRi treatment caused an increase in COMET tail moment compared with controls, similar to what is caused by high-dose camptothecin (CPT) treatment [Fig. 3D], confirming DSB formation and replication fork collapse in ATRi-treated cells.

**ATR inhibition causes nascent-strand ssDNA formation**

To confirm that the γH2AX and RPA32 hyperphosphorylation that happens in cells treated with HU and the ATRi occur at replication forks, we used iPOND [Sirbu et al. 2011, 2012]. iPOND uses click chemistry to conjugate biotin to a nucleoside analog (EdU) incorporated in newly synthesized DNA, thereby permitting a single-step purification of proteins near the replication fork. As expected, the γH2AX is associated with the newly synthesized DNA strands [Fig. 4A]. Additionally, we observed a striking accumulation of hyperphosphorylated RPA purified with the nascent DNA when ATR is inhibited [Fig. 4A]. Quantitative mass spectrometry following the iPOND purification also confirmed an increase in total RPA in the HU+ATRi conditions [B. Sirbu and D. Cortez, unpubl.]. iPOND only purifies RPA bound to or immediately adjacent to the EdU-labeled nascent strand and therefore does not detect an increase in RPA in cells treated with HU alone [Sirbu et al. 2011].
Thus, the increased abundance of RPA in the iPOND samples may suggest that the EdU-labeled nascent strand becomes single-stranded and bound to RPA when ATR is inhibited.

To directly test whether the nascent strand becomes single-stranded at stalled forks in the absence of ATR activity, we developed an assay to selectively detect nascent-strand ssDNA using a short (10 min) BrdU incubation immediately before stalling forks with HU. Under nondenaturing conditions, the BrdU antibody selectively recognizes ssDNA [Fig. 4B]. Treatment with HU and DMSO vehicle resulted in little BrdU staining [Fig. 4C, see “4h, DMSO” image]. Conversely, treatment with HU and ATRi resulted in robust BrdU staining, indicating that the nascent strand becomes ssDNA [Fig. 4C,D]. The nascent-strand ssDNA became detectable after 1 h of treatment and increased dramatically after 2 and 4 h (Fig. 5C,D). The timing of nascent-strand ssDNA formation correlates with the cellular lethality and γH2AX hyper-phosphorylation caused by ATR inhibition.

We also tested whether ATR inhibition caused the appearance of parental-strand ssDNA at the replication fork. After labeling overnight with BrdU, we released cells into fresh growth medium for 2 h before treating with HU with or without ATRi. As expected, some parental-strand ssDNA formed during HU treatment alone compared with untreated; however, much more parental-strand ssDNA formed in the ATR-inhibited condition, indicating that ATR inhibition results in increased nascent- and parental-strand ssDNA [Fig. 4E]. Thus, fork collapse when ATR is inactivated is characterized by DSBs and a large increase in ssDNA, consisting of both the template and newly synthesized DNA strands.

**ATR prevents SLX4- and CtIP-dependent formation of DSBs and nascent-strand ssDNA at stalled forks**

We reasoned that the nascent-strand ssDNA could be generated from template-strand resection after DSB formation, branch migration of the stalled fork to yield a reversed fork structure with unequal length nascent DNA strands, or both [Fig. 4B]. The MUS81 structure-specific endonuclease generates DSBs during persistent exposure to replication stress agents or in response to silencing or inhibition of replication fork repair proteins such as WRN, SMARCAL1, and CHK1 [Hanada et al. 2007; Franchitto et al. 2008; Forment et al. 2011; Betous et al. 2012]. Therefore, we tested whether siRNA depletion of MUS81 affects DSB formation in HU/ATRi-treated cells. Surprisingly, MUS81-depleted cells showed no difference in γH2AX or DSB formation compared with controls [Fig. 5A; data not shown]. We also failed to observe a change in HU/ATRi-induced ssDNA formation with MUS81 knockdown or in MUS81-deficient HCT116 cells [Supplemental Fig. S1].

Multiple other structure-specific nucleases exist in mammalian cells. SLX4 is a molecular scaffold that functions as the structural subunit of the SLX4−SLX1 Holliday junction resolvase and coordinates the SLX1, XPF, and MUS81 nucleases [Fekairi et al. 2009; Svendsen et al. 2009]. In contrast to MUS81, silencing SLX4 by siRNA largely abrogated DSB formation in HU+ATRi-treated cells [Fig. 5B]. Furthermore, SLX4 depletion also resulted in a large decrease in nascent-strand ssDNA formation [Fig. 5C,D]. This result is not due to a change in the S-phase population [Fig. 5E]. Furthermore, the response in the nascent-strand ssDNA assay correlates with SLX4 knockdown efficiency, indicating that it is not an off-target effect of the siRNA [Fig. 5F]. Silencing SLX1, XPF, MUS81, or the GEN1 nuclease individually had no consistent effect on nascent-strand ssDNA formation [Supplemental Figs. S1–S4]. Thus, it is likely that multiple SLX4-coordinated nucleases function redundantly to process HU-stalled replication forks when ATR is inhibited.

Once the fork is cleaved, ssDNA could be generated by DNA end resection. CtIP promotes resection at DSBs through multiple exonucleases [Paull 2010]. Silencing CtIP significantly reduced the amount of nascent-strand ssDNA with three of four siRNAs, and the fourth siRNA decreased nascent-strand ssDNA in all experiments but did not reach statistical significance [Fig. 5G,H]. CtIP protein levels decreased to nearly undetectable levels with all four siRNAs, and the change in nascent-strand ssDNA intensity was not due to a change in the S-phase population of these cells [Supplemental Fig. S5]. These results suggest that at least some of the ssDNA that accumulates at stalled forks when ATR function is inhibited is generated by end resection. Knockdown of individual exonucleases (including EXO1 and DNA2) alone did not result in any change in nascent-strand ssDNA, consistent with redundancy in end resection mechanisms [Supplemental Fig. S6].

**Nascent-strand ssDNA formation involves replication fork remodeling**

We next asked whether any fork processing steps occur upstream of SLX4-dependent cleavage. Specifically, we hypothesized that fork remodeling enzymes may be needed to generate a reversed replication fork structure that looks like a Holliday junction (or “chicken foot”), which is a preferred substrate for SLX4-dependent endonucleases. Therefore, we tested whether nascent-strand ssDNA formation depends on enzymes known to catalyze replication fork regression. SMARCAL1 is a DNA translocase that can anneal DNA strands and perform replication fork regression at stalled forks with leading strand gaps [Betous et al. 2012, 2013]. Silencing SMARCAL1 with four siRNAs decreases nascent-strand ssDNA-positive cells compared with controls [Fig. 6A–C; Supplemental Fig. S7]. Silencing other enzymes capable of catalyzing fork regression, including FANCM, HILT, ZRANB3, and BLM, did not decrease nascent-strand ssDNA [Supplemental Figs. S8, S9].

To further confirm the requirement of SMARCAL1 in the fork collapse associated with ATR inactivation, we examined whether a similar phenotype could be observed in the Xenopus cell-free replication system. In this system, replication of sperm chromatin in the presence of the
DNA polymerase inhibitor aphidicolin causes ATR activation due to polymerase and helicase uncoupling (Byun et al. 2005). Prelabeling the newly synthesized DNA strands with BrdU allowed us to observe the appearance of nascent-strand ssDNA when either CPT was added to promote fork collapse or ATR was inhibited in the presence of aphidicolin [Fig. 6D–F]. Depleting xSMARCAL1 from the replicating extracts prevented the nascent-strand ssDNA generation only in the context of ATR inhibition with aphidicolin (Fig. 6D–F). xSMARCAL1 depletion had no effect on CPT-induced ssDNA or the activation of the ATR checkpoint (Fig. 6G). Thus, just as in mammalian systems only yielded a partial suppression of the ssDNA phenotype, indicating that other mechanisms may also be important.

Replication stress induces SMARCAL1 phosphorylation after it accumulates at stalled forks and binds DNA

One mechanism to explain these results is that ATR prevents aberrant fork processing by directly regulating SMARCAL1. The lack of effect of xSMARCAL1 depletion on CPT-induced ssDNA formation would be consistent with this mechanism since ATR is active in these circumstances. ATR can phosphorylate SMARCAL1 (Bansbach et al. 2009), but the functional consequences have not been described. Therefore, we investigated whether SMARCAL1 phosphorylation is a mechanism by which ATR stabilizes stalled replication forks.

SMARCAL1 phosphorylation in response to replication stress can be monitored by a pronounced gel mobility shift of the endogenous protein [Fig. 7A; Bansbach et al. 2009]. GFP-SMARCAL1 protein expressed at near endogenous levels allows a robust signal and is sufficient for the phosphorylation stress can be monitored by a pronounced gel mobility shift of the endogenous protein (Fig. 7A; Bansbach et al. 2009), but the functional consequences have not been described. Therefore, we investigated whether SMARCAL1 phosphorylation is a mechanism by which ATR stabilizes stalled replication forks.
enous levels in HEK293T cells also is phosphorylated in response to HU (Fig. 7A). SMARCAL1 localizes to stalled forks via an interaction between its N terminus and RPA2 [Bansbach et al. 2009; Ciccia et al. 2009; Yuan et al. 2009; Yusufzai et al. 2009]. However, abrogation of stalled fork localization with a mutant that does not bind RPA (ΔN)
caused a loss of damage-dependent phosphorylation, indicating that SMARCAL1 must be in proximity to the stalled fork to be phosphorylated (Fig. 7B). Deletion or point mutations in the SMARCAL1 HARP1 domain decrease but do not eliminate DNA binding (Betous et al. 2012) and decrease damage-dependent SMARCAL1 phosphorylation (Fig. 7B). Deletion or mutation of the HARP2 domain eliminates DNA binding (Betous et al. 2012) and eliminates phosphorylation (Fig. 7B). These HARP mutants retain the ability to localize to stalled forks [Supplemental Fig. S10], indicating that both localization and DNA binding are required for damage-dependent phosphorylation. Thus, phosphorylation occurs after SMARCAL1 is positioned to catalyze fork remodeling. However, the ability of SMARCAL1 to translocate on DNA is not a prerequisite for phosphorylation, since the ATPase-deficient R764Q mutant exhibits the same mobility shift as the wild-type protein in response to HU (Fig. 7B).

**S652 phosphorylation reduces SMARCAL1 activity**

To test the functional significance of damage-dependent phosphorylation, we isolated SMARCAL1 before and after exposure to HU. Phosphorylated SMARCAL1 isolated from HU-treated cells retains only 50% activity compared with SMARCAL1 isolated from untreated cells (Fig. 7C). This suggests that HU-dependent post-translational modifications inactivate SMARCAL1. This difference in activity is not due to a difference in cell cycle stage induced by HU, since the cells were synchronized and released into S phase prior to the experiment.

Next, we used mass spectrometry and phosphopeptide mapping to identify damage-induced phosphorylation sites. Mass spectrometry of SMARCAL1 purified from HU-treated cells identified phosphorylation of S652, which lies within a consensus sequence [SQ] for the ATR kinase [Supplemental Fig. S11] on a linker between the two lobes of the SMARCAL1 ATPase domain [Fig. 7D]. We identified two additional SQ phosphorylation sites [S173 and S919] through phosphopeptide mapping (Supplemental Fig. S11). These phosphorylation sites are within poorly conserved, unstructured regions near the N and C termini, respectively (Fig. 7D).

Since S652 is highly conserved, we raised phosphopeptide-specific antibodies to this phosphorylation site. These antibodies show that S652 phosphorylation is increased in response to HU [Fig. 7E]. Phosphorylation is largely ATR-dependent in cells [Fig. 7F], and purified ATR can phosphorylate S652 in an in vitro kinase assay [Fig. 7G].

To determine whether these phosphorylation sites are functionally important, we tested whether phosphorylation alters SMARCAL1 localization or biochemical functions. Phosphorylation of S173, S652, or S919 does not regulate SMARCAL1 localization to stalled replication forks, since neither nonphosphorylatable nor phosphomimetic mutations in these residues alter their ability to colocalize with RPA foci in response to DNA damage [Supplemental Fig. S12]. Mutations of S173 or S919 also did not affect the DNA-stimulated ATPase activity of SMARCAL1 [Fig. 8A,B]. In contrast, the phosphomimetic S652D mutant is a much less active DNA-stimulated ATPase compared with the wild type [Fig. 8C,D], even though it has indistinguishable DNA-binding activity [Supplemental Fig. S13]. The extent of ATPase inhibition caused by the phosphomimetic mutation is similar to the inhibition of SMARCAL1 observed after HU-treatment [Fig. 7C], suggesting that S652 phosphorylation is a major mechanism by which SMARCAL1 is inhibited in cells exposed to persistent replication stress. Combining the S173D, S652D, or S919D mutations did not further alter SMARCAL1 activity [data not shown], and mutation of S652 to alanine had no effect on SMARCAL1 activity, indicating that the serine itself is not required for function.

Consistent with its reduced ATPase activity, we also observed a significant reduction in the ability of the phosphomimetic S652D SMARCAL1 to catalyze fork regression compared with the S652A or wild-type proteins, indicating that SMARCAL1 phosphorylation at S652 limits its ability to perform branch migration [Fig. 8E,F]. Combined with the observation that SMARCAL1
must first bind DNA before it can be phosphorylated, these data are consistent with a model in which SMARCAL1 initially is recruited to a damaged fork, binds the DNA to perform a fork remodeling activity, and then is phosphorylated on S652 to regulate this activity.

**Too much SMARCAL1 activity in cells phenocopies the effect of ATR inhibition on fork collapse**

To confirm that ATR-dependent SMARCAL1 phosphorylation also inhibits SMARCAL1 activity in cells at stalled replication forks, we used an assay that measures the consequences of uncontrolled SMARCAL1 activity. Expression of wild-type SMARCAL1 from a strong promoter causes the induction of γH2AX throughout the nucleus in a pan-nuclear pattern (Fig. 9A–C; Bansbach et al. 2009) similar to what is observed when ATR is inhibited in HU-treated cells (Fig. 3A), supporting a functional link between ATR and SMARCAL1. The induction of pan-nuclear γH2AX by overexpressed SMARCAL1 is dependent on DNA replication, the ability of SMARCAL1 to localize to stalled forks, and its enzymatic activity (Bansbach et al. 2009). Like ATR inhibition, overexpression of SMARCAL1 causes increased levels of ssDNA at replication forks [Fig. 9D,E]. Furthermore, just as is the case with ATR inhibition, knockdown of SLX4, but not MUS81, resulted in a decrease in ssDNA formation in cells expressing SMARCAL1 [Fig. 9E]. Taken together, these data indicate that the phenotype of cells overexpressing SMARCAL1 is similar to that induced when ATR is inhibited, as predicted if ATR phosphorylates SMARCAL1 as a mechanism of limiting its activity and preventing fork collapse.

If ATR-catalyzed S652 phosphorylation decreases SMARCAL1 activity to prevent aberrant fork processing, then the S652D phosphomimetic mutation should yield a protein that is less capable of inducing the pan-nuclear γH2AX phenotype. Indeed, overexpression of GFP-SMARCAL1 S652D induces significantly less pan-nuclear H2AX phosphorylation compared with the wild-type protein [Fig. 9A,B]. We did observe some cells with pan-nuclear γH2AX when S652D is overexpressed; however, those cells contained, on average, twice the amount of S652D protein overexpression compared with the level of the wild-type or S652A proteins needed to induce pan-nuclear γH2AX [Fig. 9C].

Finally, consistent with our model, we also found that overexpression of the S652D SMARCAL1 protein yielded less ssDNA than the S652A mutant, although it does remain slightly more active than a SMARCAL1 ΔN protein that cannot localize to stalled forks because it lacks an RPA-binding domain [Fig. 9D]. These data suggest that S652 of SMARCAL1 is one of the ATR phosphorylation targets necessary to prevent fork collapse and emphasize the need for properly regulating SMARCAL1 to achieve successful stalled fork stabilization and repair.

**Discussion**

Our data are consistent with a specific model of how ATR prevents fork collapse in the context of replication stress.

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**Figure 9.** Phosphorylation of SMARCAL1 at S652 decreases its activity at DNA replication forks in cells. (A–C) GFP-SMARCAL1 wild-type and mutant proteins were overexpressed in U2OS cells. Cells were stained with DAPI to mark the nucleus and antibodies to γH2AX. Images were acquired using an Opera automated confocal microscope, and the levels of GFP-SMARCAL1 and γH2AX levels were quantitated in each nucleus using Columbus software. (A) Representative images. (B) Data represent the percentage of cells expressing between 500 and 2500 arbitrary units of GFP-SMARCAL1 that contain a mean γH2AX intensity of >1000 arbitrary units. Error bars represent SEM from three independent experiments. (*) \( P = 0.0007; \) (**) \( P = 0.023. (C) The expression level of GFP-SMARCAL1 (as measured by GFP intensity) in each cell that had a γH2AX intensity of >1000 arbitrary units is plotted in box and whisker format, significantly higher GFP-SMARCAL1 S652D protein levels were needed to induce γH2AX than either wild-type or S652A protein \( P < 0.0001. (D) GFP-SMARCAL1 proteins with the indicated mutations were expressed in U2OS cells. BrdU was added to the culture medium 16 h prior to fixation and staining in non-denaturing conditions to measure the total level of ssDNA. The mean intensity (arbitrary units) of BrdU staining per GFP-SMARCAL1 S652D protein levels were needed to induce γH2AX than either wild-type or S652A protein \( P = 0.013). (E) GFP-SMARCAL1 proteins were expressed in U2OS. These cells were then transfected with control, MUS81, or SLX4 siRNA. BrdU was added to the culture medium 16 h prior to fixation and staining in non-denaturing conditions to measure the total level of ssDNA. The mean intensity (arbitrary units) of BrdU staining per cell is graphed. The line indicates the mean value in each population [Mann-Whitney test, S652A vs. S652D, \( P = 0.013); (F) Model for nascent-strand ssDNA generation at stalled forks. Black and red lines represent template and nascent strands, respectively. HU causes uncoupling of the replicative helicase and polymerases, resulting in template-strand ssDNA at the replication fork. ATR prevents aberrant fork remodeling by the SMARCAL1 enzyme. In the absence of ATR-dependent SMARCAL1 S652 phosphorylation, a Holliday junction-like structure may persist at the fork and is cleaved by SLX4-dependent nucleases, generating a DSB. CtIP-dependent nucleases then resect the break, yielding nascent-strand ssDNA. CtIP may also process a reversed fork structure prior to SLX4 cleavage, which could contribute to the nascent-strand ssDNA formation.
tion checkpoint-deficient yeast is linked to its requirement to complete DNA replication and separable from its G2 checkpoint activity [Nam et al. 2011]. Defining how ATR promotes the completion of DNA replication is essential to understanding the major pathway that controls genome integrity in S phase and identifying the mechanism of action of ATR-directed drugs. Our data indicate that acute ATR inhibition results in a rapid decrease in the rate of fork elongation. This effect likely results from loss of checkpoint control of origin firing, leading to a decrease in elongation rates due to depletion of essential replication factors such as nucleotides or replication proteins. The deregulation of origin timing is unlikely to be cell-lethal by itself, since it happens within 20 min of adding the ATRi, but loss of cell viability requires several hours without additional genotoxic agents. Furthermore, suppression of origin firing by the addition of CDK2 or CDC7 inhibitors did not improve the viability of ATRi-treated cells [Supplemental Fig. S14]. More likely the essential function of ATR is to stabilize or repair stalled replication forks, which collapse when ATR is inhibited. Deregulated origin firing would exacerbate this problem by creating additional stalled forks. Consistent with this interpretation, viability is lost within 45 min in cells treated with both ATRi and HU. The increase in irreversible replication fork collapse, concomitant formation of DSBs, and excess ssDNA when ATR is inactivated suggest improper enzymatic processing of the stalled DNA. These fork processing events may be largely unsuccessful attempts to repair the damaged fork.

Aberrant stalled fork processing when ATR is inactivated

Previous studies implicated the MUS81 endonuclease as an enzyme that cleaves persistently stalled replication forks after several hours of HU treatment, leading to the appearance of high levels of γH2AX [Hanada et al. 2007]. MUS81 was also reported to generate DNA damage in cells treated with a CHK1 inhibitor [Forment et al. 2011]. However, we found no effect of MUS81 on the generation of DSBs in cells treated with the ATRi. Instead, we observed a strong dependency on SLX4. Furthermore, we found that SLX4 is required for nascent-strand ssDNA formation. SLX4 is a scaffolding protein for several endonucleases, including MUS81, SLX1, and XPF [Fekairi et al. 2009; Svendsen et al. 2009]. Since we did not observe a significant effect of silencing any of these nucleases individually, our data suggest that more than one of them may function redundantly to cleave stalled forks when ATR is inhibited. CtIP-dependent exonucleases then process the cleaved fork to generate nascent-strand ssDNA. SLX4 depletion yields a complete loss of DSBs as measured by the COMET assay but only a partial suppression of the nascent-strand ssDNA. Therefore, it is possible that CtIP also directs resection of a stalled, repressed fork when ATR is inhibited prior to DSB formation, thereby generating a nascent-strand ssDNA overhang with or without SLX4-dependent cleavage.

A preferred substrate of SLX4-dependent endonucleases resembles a Holliday junction, which can form at a stalled fork due to either torsional stress or enzymatic processing. Our data are consistent with enzymatic processing of the fork by the SMARCAL1 fork-regressing enzyme to generate a preferred substrate for SLX4-dependent cleavage when ATR is inhibited. ATR phosphorylates SMARCAL1, and previous studies indicate that excess SMARCAL1 activity causes replication fork problems [Bansbach et al. 2009]. Thus, the fork remodeling activities of SMARCAL1 promotes fork repair in some cases but can threaten genome stability if not properly regulated. SMARCAL1 is not needed for fork collapse in all circumstances. For example, our data indicate that CPT treatment induces fork collapse and ssDNA formation independently of xSMARCAL1. Since ATR would be active in CPT-treated cells, SMARCAL1 would be appropriately regulated.

Consistent with our model, we found that phosphorylated SMARCAL1 purified from cells treated with HU is significantly less active than SMARCAL1 purified from untreated cells. We identified three damage-induced phosphorylation sites on SMARCAL1, including S652, which is phosphorylated by ATR. Phosphomimetic mutations in S652 yield a protein that is significantly less active in catalyzing ATP hydrolysis and fork reversal in vitro and also less active in cells. S652 lies within the linker between the two RecA lobes of the ATPase domain. Flexibility of this linker is needed for conformational changes required for catalysis [Durr et al. 2005; Sprouse et al. 2006; Lewis et al. 2008]. Since S652 phosphorylation does not impair SMARCAL1 localization or DNA-binding...
activity, we suspect that phosphorylation of the linker reduces the ability of SMARCAL1 to undergo this conformational change and thereby inhibits its translocation activity. SMARCAL1 phosphorylation requires it to first localize to the stalled fork and bind DNA. Thus, SMARCAL1 likely acts at damaged forks prior to ATR-dependent phosphorylation, which serves as a mechanism of limiting its activity.

Overexpression of active SMARCAL1 causes pan-nuclear γH2AX staining. This effect requires localization of SMARCAL1 to replication forks and its enzymatic activity. The exact source of this staining pattern is not clear, although it forms well before there is any evidence of apoptosis, is chromatin-associated [data not shown], and is also observed in cells treated with ATRi and HU. Thus, our data suggest that it is linked to the aberrant fork processing and collapse that occurs when ATR cannot regulate SMARCAL1. Consistent with this interpretation, the phosphomimetic S652D SMARCAL1 protein is less capable of inducing this phenotype and is less active as a fork regression enzyme. Furthermore, like ATR inhibition, SMARCAL1 overexpression also induces SLX4-dependent excess ssDNA formation at stalled forks, and this effect is alleviated by a phosphomimetic mutation in S652.

Another prediction of our model is that the S652A mutant should retain too much activity in HU-treated cells, perhaps generating some fork collapse events even when ATR is not inhibited [assuming S652 phosphorylation is not completely redundant with other mechanisms]. Unfortunately, since SMARCAL1 overexpression causes significant perturbations to DNA replication on its own and the S652 site does not appear to be regulated properly on the exogenously expressed SMARCAL1 wild-type protein [Supplemental Fig. 11], we were unable to do effective genetic complementation experiments to examine the effect of the S652A mutant on fork collapse and ssDNA generation. Future experiments—either knocking in the mutation into the genome or using expression systems with native promoter and gene structures—will be needed.

Neither SMARCAL1 nor SLX4 silencing completely prevents nascent-strand ssDNA formation caused by ATR inhibition, suggesting that additional aberrant fork processing events remain to be identified. The lack of complete suppression by SMARCAL1 silencing may also be due to the competing increase in ssDNA generated due to the need for some properly regulated SMARCAL1 protein to protect stalled forks. In addition, it seems likely that additional mechanisms regulate the activity of SMARCAL1. Our phosphopeptide maps indicate that S652, S173, and S919 are the major HU-induced phosphorylation sites on SMARCAL1; however, mutation of all three serines to alanines does not eliminate the HU-induced mobility shift of the protein on SDS-PAGE gels [data not shown]. Thus, there must be additional regulatory post-translational modifications that could also contribute to SMARCAL1 regulation. Finally, the ATRi may not completely block all ATR activity, so the results observed may not be equivalent to a complete loss of ATR function.

Conclusions

ATR signaling has long been known to regulate DNA replication and prevent fork collapse, based largely on studies in yeast. However, the molecular mechanisms of what ATR does to prevent fork collapse and even what fork collapse is in mammalian cells are not well understood. Our data define an ATR-dependent replication fork protection mechanism in human cells that is critical for genome maintenance and cell viability. These studies also help define the mechanism of action of ATR targeted therapeutic agents. Specifically, combinations of selective ATR pathway inhibitors with either intrinsic or added replication stress provide a rapid mechanism of cancer cell killing due at least in part to aberrant processing of stalled replication forks.

Materials and methods

Cell culture, clonogenic survival, and ATRi

U2OS cells were maintained in DMEM with 7.5% FBS. RPE-hTERT cells were maintained in DMEM/F12 medium with 10% FBS, 0.348% sodium bicarbonate, and 2 mM L-glutamine. MUS81−/− HCT116 cells [Shimura et al. 2008] were maintained in McCoy’s 5A medium with 10% FBS. For clonogenic assays, U2OS cells were seeded onto 60-mm cell culture plates at 200–5000 cells per plate in triplicate. The following day, cells were treated with drug, washed, and released into fresh growth medium for 10–14 d prior to staining with methylene blue. The VE-821 ATR-selective inhibitor [Reaper et al. 2011] was synthesized by the Vanderbilt Institute for Chemical Biology Chemical Synthesis facility and used in most experiments. ATR inhibition in Xenopus extracts was achieved with 64 μM 4-[4-[[3R]-3-methylmorpholin-4-yl]-6-[4-(methylylsonyl)piperidin-4-yl]pyrimidin-2-yl]-1H-indole [Foote et al. 2010], synthesized by CheminPharma. The minimum concentration needed to achieve a near-complete block of ATR-dependent CHK1 phosphorylation was determined and used in all experiments. This concentration did not inhibit ATM, DNA-PKcs, or mTOR kinases [Reaper et al. 2011; data not shown]. siRNA transfections were performed using either HiPerfect (Qiagen) or Dharmafect 1 [Dharmacon] at a final siRNA concentration of 10 nM. siRNAs were purchased from Dharmacon or Ambion.

Fiber labeling

htERT-RPE cells were labeled with 20 μM EdU for 20 min, rinsed twice with equilibrated HEPES-buffered saline, treated with 2 mM HU or 3 μM ATRi as indicated, and then labeled with 100 μM CdU (Sigma) for 20 min. Cells were harvested and resuspended in ice-cold PBS. Two microliters of cell suspension was deposited on a microscope slide, and 10 μL of spreading buffer [200 mM Tris-HCl at pH 7.4, 0.5% SDS, 50 mM EDTA] was added to the cells on the slide for 6 min. Next, the slides were tilted to 15° to stretch the DNA fibers. Following fixation in a 3:1 solution of methanol/acetone acid, the DNA was denatured with 2.5N HCl, blocked with PBS containing 10% goat serum and 0.1% Triton X-100, and stained with rat anti-CdU [Abcam] and mouse anti-EdU [Becton Dickinson] diluted in PBS containing 10% goat serum and 0.1% Triton X-100 followed by secondary antibodies [Alexa-594 goat anti-rat IgG and Alexa-488 goat anti-mouse; Invitrogen].
ATR regulates SMARCAL1

ATPase and fork reversal assays

ATPase assays were performed as described previously with a splayed-arm DNA substrate [Betous et al. 2012]. Fork reversal assays were completed with 3 nM gel-purified, labeled fork reversal DNA substrate containing a leading strand gap and 3 nM SMARCAL1 protein in reaction buffer (40 mM Tris at pH 7.5, 100 mM KCl, 5 mM MgCl₂, 100 μg/mL BSA, 2 mM ATP, 2 mM DTT). Reactions were terminated by the addition of 3× stop buffer (0.9% SDS, 50 mM EDTA, 40% glycerol, 0.1% bromophenol blue, 0.1% xylene cyanol). Samples were separated on 8% polyacrylamide (19:1) × TBE gels. The gels were dried and quantified using a Molecular Imager FX (Bio-Rad). Sequences of oligonucleotides used to make substrates are listed in Supplemental Table S1.

Immunoblotting and antibodies

Rabbit polyclonal SMARCAL1 909 antibody was described previously (Banbach et al. 2009). The pS652 phosphopeptide-specific antibody was ordered from Bethyl Laboratories and made using the following peptide antigen: KSDVL(pS)QLPAK. Specificity was confirmed using phosphatase-treated lysates and a S652A mutant [data not shown]. Rabbit polyclonal antibody against the C-terminal 890 amino acids of xSMARCAL1 was raised at Josman, LLC. Additional antibodies used included RPA2 [clone 9H8, Abcam]; γH2AX [clone JBW301, Upstate Biotechnology]; Flag M2 [Sigma]; CHK1 pS317, CHK1 pS345, and CHK2 pT68 [Cell Signaling]; CHK1 and GAPDH (G4, Santa Cruz Biotechnology); RPA32 pS4-S8, RPA32, and SLX4 (Bethyl Laboratories); H3 and CtIP [Abcam]; and HA (Covance). Quantitative immunoblotting was performed using an Odyssey instrument.

Pan-nuclear γH2AX immunofluorescent assay

U2OS cells were transfected with GFP-SMARCAL1 encoding vectors [pLEGFP-C1, Clontech] using Fugene HD (Promega) according to the manufacturer’s instructions. Twenty-four hours after transfection, the cells were seeded into 96-well CellCarrier plates [Perkin Elmer]. Forty-eight hours after transfection, the cells were washed once with PBS, fixed with 3% paraformaldehyde solution, and permeabilized with 0.5% Triton X-100 for 10 min at 4°C. Fixed cells were then incubated with mouse anti-BrdU antibody [BD Pharmingen] for 60 min at 37°C followed by Alexa-488 goat anti-mouse [Invitrogen] secondary antibody. Images were collected using a Zeiss camera at a constant exposure time and quantitated with Cell Profiler (http://www.cellprofiler.org).

Neutral COMET assays and iPOND

The neutral COMET assays were performed in accordance with the manufacturer’s [Trevigen] instructions. The iPOND procedure was carried out as described previously [Sirbu et al. 2011, 2012].

ssDNA immunofluorescent assays

To detect parental-strand ssDNA, the cells were labeled for 20 h with 10 μM BrdU and then released into fresh growth medium for 2 h prior to addition of drugs. To detect nascent-strand ssDNA, the cells were labeled for 10 min with 10 μM BrdU immediately prior to addition of drugs. Three millimolar HU was added with 5 μM ATRi. Next, the cells were washed once with PBS, permeabilized with 0.5% Triton X-100 for 10 min at 4°C, fixed with 3% paraformaldehyde/2% sucrose solution for 10 min, and blocked for 15 min in 3% BSA-PBS. Fixed cells were then incubated with mouse anti-BrdU antibody [BD Pharmingen] for 60 min at 37°C followed by Alexa-488 goat anti-mouse [Invitrogen] secondary antibody. Images were collected using a Zeiss camera at a constant exposure time and quantitated with Cell Profiler (http://www.cellprofiler.org).

Xenopus cell-free replication experiments

Xenopus low-speed extract was prepared as described [Lebofsky et al. 2009]. Immunodepletion of xSMARCAL1 was carried out by incubation of serum with nProtein A-Sepharose Fast Flow beads [GE Healthcare] at a 1:1 ratio for 1.5 h at 4°C, and the resulting beads were mixed with extract at a 3:1 extract:bead ratio for 1 h at 4°C. Rabbit IgG (10 mg/mL; Sigma-Aldrich) was incubated with nProtein A–Beads for mock depletions at a 1:10 ratio for 1 h at 4°C. For nuclei spin-downs, extracts (20 μL) were transferred to 1 mL of dilution buffer (30% glycerol, 150 mM KCl, 0.5% Triton X-100, 80 mM PIPES at pH 6.8, 1 mM MgCl₂, 1 mM EGTA), and 1 mL of fixation buffer (30% glycerol, 0.5% Triton X-100, 80 mM PIPES at pH 6.8, 1 mM MgCl₂, 1 mM EGTA, 4% formaldehyde) was added. The resulting 2 mL was then layered on top of a glycerol cushion (40% glycerol, 80 mM PIPES at pH 6.8, 1 mM MgCl₂, 1 mM EGTA) and spun at 3500 rpm for 15 min in a JS-4.2 rotor. Nuclei were washed with TBS and blocked in TBS + 1% BSA overnight. BrdU antibody [il55, Santa Cruz Biotechnology] was added for 5 h, followed by Alexa-594 goat anti-mouse [Invitrogen]. Mean BrdU signal intensity was quantitated using ImageJ.

Protein purification

Flag-SMARCAL1 was purified from HEK293T cells following transient transfection or from insect cells after baculovirus infection as described previously [Betous et al. 2012]. Endogenous SMARCAL1 was purified from HEK293T cells using SMARCAL1 909 antibodies bound to protein A-conjugated magnetic beads using the same procedure.

In vitro kinase assay

GST-S652 protein contained amino acids 645–661 of SMARCAL1 inserted into pBG101. GST-His protein was produced from the pBG101 vector without insertion of SMARCAL1 amino acids. It was larger than GST-S652, since there was no stop codon to prevent translation through the polylinker. Both proteins were purified from Escherichia coli using glutathione sepharose. Kinase reactions were performed with hyperactive ATR–ATRIP (ATR-interacting protein) purified from human cells and incubated with the TOPBP1 AAD protein as previously described [Mordes et al. 2008].

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References


Boeroekel CF, Takashima H, John J, Yan J, Stankiewicz P, Brown EJ, Baltimore D. 2000. ATR disruption leads to chromo-

catalyzes fork regression and Holliday junction migration to maintain genome stability during DNA replication. Genes Dev 14: 397–402.


O’Driscoll M, Ruiz-Perez VL, Woods CG, Jeggo PA, Goodship JA. 2003. A splicing mutation affecting expression of ataxia-


ATR phosphorylates SMARCAL1 to prevent replication fork collapse

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