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SMARCAL1 catalyzes fork regression and Holliday junction migration to maintain genome stability during DNA replication

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SMARCAL1 (SWI/SNF-related, matrix-associated, actin-dependent regulator of chromatin, subfamily A-like1) maintains genome integrity during DNA replication. Here we investigated its mechanism of action. We found that SMARCAL1 travels with elongating replication forks, and its absence leads to MUS81-dependent double-strand break formation. Binding to specific nucleic acid substrates activates SMARCAL1 activity in a reaction that requires its HARP2 (Hep-A-related protein 2) domain. Homology modeling indicates that the HARP domain is similar in structure to the DNA-binding domain of the PUR proteins. Limited proteolysis, small-angle X-ray scattering, and functional assays indicate that the core enzymatic unit consists of the HARP2 and ATPase domains that fold into a stable structure. Surprisingly, SMARCAL1 is capable of binding three-way and four-way Holliday junctions and model replication forks that lack a designed ssDNA region. Furthermore, SMARCAL1 remodels these DNA substrates by promoting branch migration and fork regression. SMARCAL1 mutations that cause Schimke immunoosseous dysplasia or that inactivate the HARP2 domain abrogate these activities. These results suggest that SMARCAL1 continuously surveys replication forks for damage. If damage is present, it remodels the fork to promote repair and restart. Failures in the process lead to activation of an alternative repair mechanism that depends on MUS81-catalyzed cleavage of the damaged fork.

Keywords: DNA repair; HARP; Holliday junction; fork reversal; SIOD; SAXS

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SMARCAL1 [SWI/SNF-related, matrix-associated, actin-dependent regulator of chromatin, subfamily A-like1], also known as DNA-dependent ATPase A and HARP (Hep-A-related protein), is a member of the SNF2 family of ATPases [Flaus et al. 2006]. Many of these proteins use the energy of ATP hydrolysis to translocate along DNA and thereby remodel DNA structures or DNA–protein interactions. They function in many cellular processes, including transcription, DNA replication, and DNA repair.

The mechanism of how SMARCAL1 acts to repair damaged forks remains largely unknown. Biochemically, SMARCAL1 can bind to DNA that contains single- and double-stranded regions such as forks and DNA hairpins (Muthuswami et al. 2000; Yusufzai and Kadonaga 2008). DNA binding activates its ATPase activity, and this activity promotes DNA single-strand annealing even in the presence of RPA (Yusufzai and Kadonaga 2008). The N-terminal RPA-binding domain of SMARCAL1 is not necessary for this DNA strand-annelling activity (Bansbach et al. 2009; Yusufzai et al. 2009), but patient-derived mutants lack this function. The molecular basis for this activity may not be simply translocation along dsDNA, since the related protein RAD54 cannot perform this function despite being a robust translocase (Yusufzai and Kadonaga 2008).

SMARCAL1 is a multidomain protein. The ATPase domain, which lies in the C-terminal half of the protein, is split into two regions of primary amino acid sequence by a 115-amino-acid linker sequence. The N-terminal half of the protein contains a highly sequence conserved RPA-binding domain (Bansbach et al. 2009; Ciccia et al. 2009; Postow et al. 2009; Yuan et al. 2009; Yusufzai et al. 2009), a 200-amino-acid region of low sequence conservation without predicted domain structure, and two HARP domains. The HARP domains are 55 amino acids in length with high sequence similarity but unknown function and structure. They are separated by 40 amino acids, and the second HARP domain is linked to the ATPase domain by an additional 47 amino acids.

Fusing the HARP domains to the ATPase domain of the SNF2 proteins BRG1 or HELLS is sufficient to reconstitute DNA-dependent ATPase and annealing helicase activities, suggesting that the HARP domains are important determinants of the SMARCAL1 enzyme specificity (Ghosal et al. 2011). Paradoxically, the closest homolog of SMARCAL1 in humans, annealing helicase 2 (AH2, also known as ZRANB3), also has annealing helicase activity despite a different domain structure and no unambiguous HARP domains (Yusufzai and Kadonaga 2010).

In this study, we took genetic, biochemical, and biophysical approaches to understand how SMARCAL1 functions to maintain genome integrity. We found that SMARCAL1 travels with at least some elongating replication forks, and the MUS81 structure-specific endonuclease cleaves damaged forks in SMARCAL1-deficient cells. The HARP2 domain is essential for DNA binding, and both biochemical and small-angle X-ray scattering (SAXS) data indicate that the HARP2+SNF2 domains provide the minimal enzymatic unit. The HARP domain resembles the DNA-binding domain of the PUR-α protein and has limited ability to bind DNA on its own. Surprisingly, we found that SMARCAL1 can bind three-way and four-way DNA structures and model replication forks. Furthermore, SMARCAL1 branch-migrates the four-way junction and catalyzes extensive fork regression of model replication forks. These data provide mechanistic insight into how SMARCAL1 functions and suggest that it remodels stalled replication forks through fork regression and branch migration to promote replication fork restart and prevent replication-associated DNA double-strand breaks.

Results

SMARCAL1 is present at DNA replication forks during an unperturbed S phase and prevents MUS81-dependent double-strand breaks

Previous analyses indicated that SMARCAL1 localizes to nuclear foci that colocalize with replisomes in response to agents that induce replication stress (Bansbach et al. 2009; Ciccia et al. 2009; Postow et al. 2009; Yuan et al. 2009; Yusufzai et al. 2009). This localization is dependent on an interaction with the replisome protein RPA. Silencing SMARCAL1 using RNAi causes elevated levels of γ-H2AX in replicating cells (Bansbach et al. 2009; Postow et al. 2009; Yuan et al. 2009). To determine whether SMARCAL1 actually is a component of active replisomes, we used the iPOND procedure (Sirbu et al. 2011) to purify active and stalled replication forks. SMARCAL1 is purified with nascent, 5-ethynyl-2′-deoxyuridin [EdU]-labeled DNA at elongating replication forks even when replication is not perturbed (Fig. 1A). It is not purified with

![Figure 1](image-url)

Figure 1. SMARCAL1 acts at replication forks to prevent MUS81-catalyzed double-strand breaks. (A) Cells were labeled with EdU for 10 min, the EdU was removed, and thymidine was added for 20 min or HU was added for 3 h prior to purifying the nascent DNA–protein complexes using the iPOND procedure. (B) EdU-labeled cells were treated with 2 mM HU for the indicated lengths of time prior to performing iPOND. The “No Clk” controls in A and B are samples treated with EdU only, but no biotin-azide was added during the click reaction. (C) U2OS cells were transfected with siRNAs targeting SMARCAL1 (S), MUS81 (M), or nontargeting (NT) as indicated. Three days after transfection, the cells were either stained with antibodies to γ-H2AX or harvested for immunoblotting with the indicated antibodies. The percentage of cells staining positive for γ-H2AX was determined by immunofluorescent imaging from three independent experiments. Cells with >10 foci were counted as positive. Error bars are the standard deviation (SD, n = 3).
the EdU-labeled DNA once the labeled DNA segment is no longer adjacent to the fork (after a chase in medium lacking EdU), indicating that it travels with at least some moving replisomes. As expected, SMARCAL1 is also found at forks stalled with hydroxyurea (HU) [Fig. 1A,B], and its mobility on SDS-PAGE gels is altered in these circumstances due to phosphorylation by checkpoint kinases [Bansbach et al. 2009].

The MUS81 endonuclease cleaves some blocked and damaged replication forks, generating a double-strand break and initiating recombination-based repair mechanisms [Osman and Whitby 2007]. Therefore, we hypothesized that the high level of γH2AX found in SMARCAL1 silenced cells could be due to double-strand breaks catalyzed by MUS81. To address this question, we measured the proportion of cells containing γH2AX after SMARCAL1 and/or MUS81 depletion. As expected, silencing SMARCAL1 caused an induction of γH2AX, while silencing MUS81 had no effect [Fig. 1C]. MUS81 silencing prevented γH2AX induction in SMARCAL1 silenced cells without significantly altering the efficiency of SMARCAL1 silencing [Fig. 1C]. Thus, γH2AX induction after SMARCAL1 depletion is MUS81-dependent.

**SMARCAL1 binds a wide variety of DNA substrates that combine ssDNA and dsDNA**

Our results suggest that SMARCAL1 either processes or prevents the formation of MUS81 substrates. Little is known about SMARCAL1 substrate specificity other than it prefers to bind DNA with both single- and double-stranded characteristics rather than ssDNA or dsDNA (Supplemental Fig. 1A; Muthuswami et al. 2000; Yusufzai and Kadonaga 2008), and its ATPase activity is activated upon DNA binding. To clarify the DNA determinants that mediate SMARCAL1 DNA binding and activation, we investigated a broad range of possible DNA substrates. We first evaluated how long the ssDNA arms of a fork need to be and found that significant SMARCAL1 binding is observable even with a fork length of only 5 nucleotides [nt] per arm [Supplemental Fig. 1B,C]. Increasing the arm lengths beyond 5 nt increases the binding affinity. We also observe a second DNA–protein complex forming when the ssDNA region is lengthened to 20 nt or more. The second, higher-molecular-weight complex may contain more than one SMARCAL1 molecule.

We next varied the length of one of the ssDNA arms while keeping the other constant and found that the length of the second arm did not influence binding affinity [Supplemental Fig. 1D,E]. In fact, SMARCAL1 bound equivalently to a fork and an ssDNA overhang substrate. Both DNA substrates stimulated SMARCAL1 ATPase activity as well [Supplemental Fig. 1F]. Furthermore, DNA substrates with either a 5’ or 3’ recessed end bind and stimulate SMARCAL1 ATPase activity equivalently (Fig. 2A–C).

At a replication fork, the free 5’ end of the nascent nucleic acid on the lagging strand template would consist of a short RNA primer rather than DNA. To test whether SMARCAL1 can bind and be activated on the lagging strand, we examined a nucleic acid substrate that mimics this chimeric nucleic acid structure. A RNA–DNA primer substrate bound and stimulated SMARCAL1 equivalently to the DNA–DNA substrate (Supplemental Fig. 1G–I).

Next, we assessed how the length of ssDNA alters SMARCAL1-binding affinity. Five nucleotides are suffi-
cient to allow some binding and elicit significant ATPase activity (Fig. 2D–F). SMARCAL1 binding and ATPase activity increase as the length of the ssDNA increases. SMARCAL1 also binds and is stimulated efficiently by a gapped DNA substrate. Maximum binding and activation require only a five single-stranded nucleotide gap, and even a nick can elicit some activity [Fig. 2G–I; Supplemental Fig. 1J,K]. When a bubble replaces the gap, increasing the length of the mismatched nucleotides to 16 significantly increases affinity [Supplemental Fig. 1L,M]. Thus, the length of ssDNA needed for optimal binding and activation of SMARCAL1 is shortest when it is presented in the context of a gap.

We also investigated how the length of dsDNA affects binding and ATPase stimulation of SMARCAL1. Optimal SMARCAL1 binding and ATPase activation requires 20 dsDNA nucleotides [Fig. 2J–L]. Greater dsDNA lengths yield no further improvement in SMARCAL1 affinity [data not shown]. Fifteen nucleotides of dsDNA on either side of a 5-nt gap are sufficient to elicit maximal SMARCAL1 binding [Supplemental Fig. 1N,O].

Taken together, these results show that SMARCAL1 binds and is activated by any nucleic acid structure that contains both single- and double-stranded regions, including an RNA–primer template. The optimal length of ssDNA that elicits binding depends on the structural context of the DNA, with 5 nt being sufficient for a gap and longer lengths promoting better binding to a forked or single-stranded overhang substrate. The optimal length of dsDNA is ~15 nt. Finally, the dsDNA and ssDNA must be within the same molecule, since adding these separately to SMARCAL1 does not elicit any binding [Supplemental Fig. 1A].

**HARP2 but not HARP1 is required for SMARCAL1 DNA-binding, ATPase, and annealing helicase activities**

To understand how SMARCAL1 binds DNA, we examined the affinity of a series of truncated SMARCAL1 proteins for a forked DNA substrate [Fig. 3A]. While deletion of the first 198 and the last 84 amino acids had no effect on SMARCAL1 DNA binding, deletion of the first 424 amino acids containing the HARP domains severely compromises the DNA-binding and ATPase activities of SMARCAL1 [Fig. 3B–E].

These results led us to hypothesize that the HARP domains may be essential for SMARCAL1 DNA binding. To test this hypothesis, we assessed the behavior of a series of HARP domain mutants [Fig. 4A]. SMARCAL1 lacking the first HARP domain (∆HARP1) binds to and is activated by a forked DNA substrate, although with slightly reduced affinity compared with wild-type SMARCAL1 [Fig. 4B,C; Supplemental Fig. 2A]. In contrast, deleting the second HARP domain alone or in combination with the first HARP domain (SMARCAL1-1HARP2 and ∆HARP1+2) severely attenuated both DNA binding and ATPase activation. The effects of the deletions were even more severe when assayed with a 5-nt, single-stranded gap substrate [Fig. 4D,E].

![Figure 3](image-url) The SMARCAL1 N terminus containing the HARP domains is necessary for DNA-binding and ATPase activity. (A) Diagram of the SMARCAL1 proteins used to identify domains required for function. Wild type (WT) in all figures is full-length SMARCAL1. (B) Overexpressed SMARCAL1 proteins were purified from HEK-293T cells and examined on an SDS-PAGE gel by immunoblotting. (C,D) Increasing amounts of purified SMARCAL1 proteins were incubated with the forked DNA substrate to measure DNA binding. (E) Increasing amounts of forked DNA were added to the SMARCAL1 fragments to measure DNA-stimulated ATPase activity. Error bars represent the mean ± SD from three independent experiments. In cases in which no error bars are visible, the SD is smaller than the symbol size. The DNA substrates corresponding to each symbol and line color are the same in D and E.

To confirm the deletion results, we generated point mutants in HARP1 and HARP2. The HARP domains are evolutionarily conserved [Fig. 4F]. We mutated two of the invariant residues within each domain to alanine [HARP1 W277A/F279A and HARP2 W372A/F379A]. These mutants exhibit DNA-binding and ATPase activity similar to the corresponding complete deletion of the domain [Fig. 4G–I]. Interestingly, we found that the decreased DNA-binding and ATPase activity of the HARP1-WF mutant yielded only a slight impairment of SMARCAL1 annealing helicase activity, while mutation of the HARP2 domain completely abolished the ability of SMARCAL1 to anneal an RPA-coated plasmid substrate [Fig. 4J]. The complete deletion of HARP1 also had no effect on the SMARCAL1 annealing helicase activity [Supplemental Fig. 2B]. These results suggest that HARP2 is critical for the DNA-binding, ATPase, and annealing helicase activities of SMARCAL1. HARP1 may have a supporting role in facilitating SMARCAL1 function.

Finally, we asked whether the HARP domains themselves have any DNA-binding activity. We found that a SMARCAL1 fragment encompassing both HARP domains [amino acids 198–425] is sufficient to bind forked DNA, albeit with much lower affinity than the full-length protein [Supplemental Fig. 3A–C]. The HARP domain–DNA complex did not migrate as a discrete band in the electrophoretic mobility shift assay; however, we were able to supershift the DNA–protein...
complex with an antibody that recognizes the recombinant HARP1-2 protein fragment, confirming the complex was not due to a contaminant in the protein purification.

The HARP2-ATPase constitutes a structural core motor domain

Our biochemical results demonstrate the importance of the HARP domain in SMARCAL1 function. To gain mechanistic insight into how SMARCAL1 might use this novel domain, SAXS experiments were performed to determine the spatial arrangement of the HARP2 and ATPase domains in solution (Fig. 5A). Limited proteolysis of the full-length protein purified from insect cells revealed a proteolytically resistant fragment consisting of the HARP2-ATPase regions (Supplemental Fig. 4A). Kratky analysis of SMARCAL1(325–954) revealed parabolic features, suggesting that the protein is globular with distinct domains (Supplemental Fig. 4B,C). The radius of gyration \( r_g \) obtained from the Guinier region was 33.0 ± 0.3 Å (Supplemental Fig. 4D), indicating that the 75-kDa protein is elongated when compared with glucose isomerase, a spherical protein at 173 kDa with a similar \( r_g \) of 32 Å.

SAXS data provide complete structural information and can be used to distinguish between different conformations of a high-resolution model or build a complete atomistic model from known domains (Rambo and Tainer 2010). Therefore, we used the SAXS data of SMARCAL1 and homology models of both the HARP2 and ATPase domains to determine the solution state of the protein. To date, there are no known structural homologs of the HARP domain. However, we discovered by sequence–structure comparison (Shi et al. 2001) that there is good agreement between the predicted secondary structural elements of the HARP domains with tandem PUR repeats observed in the structure of the purine-rich element-binding protein PUR-\( \alpha \) (Supplemental Fig. 5; Graebsch et al. 2009). PUR repeats are ~140-residue motifs consisting of anti-parallel \( \beta \)-\( \beta \)-\( \beta \)-\( \alpha \) topology that bind ssDNA and dsDNA and thus provide a reasonable structural model for the HARP domains. A model of the core ATPase domain was also created based on the crystal structure of Sulfolobus solfataricus (Sso) Rad54, which shares 23% sequence identity and 58% overall similarity with SMARCAL1 (Supplemental Fig. 6).

Preliminary normal mode analysis (Suhre and Sanejouand 2004) was performed on the core ATPase domain to produce a family of alternative conformations. Each conformation was then combined with the HARP2 model for partial ab initio modeling using a simulated annealing search algorithm. The models converged into an elongated structure that was independently validated by the close resemblance to the three-dimensional (3D) molecular envelope generated from the SAXS data using GASBOR [Fig. 5B], and the remarkable agreement between the experimental scattering curve and the theoretical curve calculated from the docking model [Fig. 5C]. The resulting HARP2-ATPase model revealed that the HARP2 and ATPase motifs form one continuous domain in the absence of DNA, suggesting that their association constitutes a structural and functional core domain necessary to drive translocation. To test this idea, we assayed whether the HARP2-ATPase protein is sufficient to
catalyze strand annealing. Indeed, SMARCAL1(325–870) and SMARCAL1(325–890) are both efficient ATP-dependent annealing helicases (Fig. 5D,E).

SMARCAL1 can bind and branch-migrate a four-way junction

The DNA-binding activities of SMARCAL1 characterized thus far suggest that SMARCAL1 may have dsDNA- and ssDNA-binding surfaces. Combined with the energy of ATP hydrolysis, SMARCAL1 may translocate along the DNA in a way that leads to single-strand annealing. To determine whether these properties could yield any other enzymatic consequences, we expanded our search for SMARCAL1 substrates to more complex DNA structures, including three-way and four-way Holliday junctions. Surprisingly, despite lacking any designed ssDNA regions, SMARCAL1 could bind these DNA substrates with only slightly reduced affinity compared with a fork substrate (Fig. 6A,B). Furthermore, both DNA substrates activated the SMARCAL1 ATPase (Fig. 6C).

Given that these structures bind SMARCAL1 and stimulate its ATPase activity, we asked whether SMARCAL1 could also induce branch migration like Rad54 (Bugreev et al. 2006). We prepared a synthetic Holliday junction consisting of two homologous and two heterologous arms, similar to those used in previous branch migration studies (Fig. 6D; Gari et al. 2008b). Indeed, SMARCAL1 catalyzed branch migration in an ATP-dependent manner (Fig. 6E,F). As expected, the SIOD patient-derived ATPase-defective mutant (R764Q) failed to promote branch migration despite having the ability to bind DNA (Fig. 6G,H; Yusufzai and Kadonaga 2008). To test the importance of the HARP domains in this process, we examined the branch migration properties of HARP1-WF and HARP2-WF SMARCAL1 mutants. While the HARP1 mutant was able to branch-migrate the Holliday junction as efficiently as the wild-type protein, the HARP2 mutant had severely attenuated activity (Fig. 6I,J).

SMARCAL1 can bind and branch-migrate a replication fork

Previous studies indicate that SMARCAL1 acts at stalled replication forks but may not have an essential function in homology-directed double-strand break repair. Double-strand breaks are only thought to form in normal cells at persistently stalled forks (Petermann et al. 2010; Sirbu et al. 2011). Thus, we investigated whether SMARCAL1 could bind and process other branched structures that might exist at a transiently stalled fork. Specifically, we compared SMARCAL1 affinity to model forks with no nascent DNA strands, a leading strand, a lagging strand, or both. Strikingly, we found that SMARCAL1 binds to and is activated by each of these structures (Fig. 7A–C). To determine whether SMARCAL1 can catalyze remodeling of these replication fork structures, we prepared a substrate to monitor fork regression (Fig. 7D; Gari et al. 2008b). SMARCAL1 catalyzed displacement of the two “nascent” DNA strands and annealing of the parental strands (Fig. 7E,F). Again, the SIOD patient-derived R764Q mutation eliminated this activity.

SMARCAL1 does not possess any helicase activity (Yusufzai and Kadonaga 2008), so it is unlikely that it could unwind the nascent strands before annealing both parental and both nascent strands together. To confirm that the SMARCAL1 fork reversal activity is coordinated without the formation of ssDNA intermediates, we labeled the model nascent leading strand of the synthetic replication fork and performed a time-course assay. We found that only a double-stranded product consisting of the two nascent strands is formed without the appearance of any ssDNA intermediates (Supplemental Fig. 7A–C). We conclude that SMARCAL1 processes replication fork structures by coupling unwinding and annealing in a concerted manner to yield fork regression. As expected, the SMARCAL1 HARP1-WF mutant is able to regress the replication fork as efficiently as the wild-type protein, whereas mutations in the SMARCAL1 HARP2 domain eliminate fork regression activity (Supplemental Fig. 7D–F). Thus, HARP2 but not HARP1 is critical for SMARCAL1 fork regression activity.
Finally, we tested whether SMARCAL1 can catalyze fork regression and sustained migration on a plasmid-sized substrate that more closely models a stalled replication fork. We created a joint molecule by annealing gapped plasmids (Fig. 7G). This substrate mimics a stalled fork in which the lagging strand is 14 nt longer than the leading strand (Ralf et al. 2006; Blastyak et al. 2007). The extent of fork regression of this substrate was detected by restriction enzyme digestion to liberate a linear 5\[^{32}\text{P}\]-labeled lagging strand. SMARCAL1 efficiently catalyzed remodeling of this substrate, yielding substantial amounts of a regressed fork corresponding to movement of at least 836 base pairs (bp) (Fig. 7H). This reaction is dependent on the amount of SMARCAL1 added to the reaction and requires ATP hydrolysis, since ATP\(\_S\) completely blocked remodeling of the substrate.

**Discussion**

Previous studies by our group and others defined SMARCAL1 as a replication stress response protein that acts to preserve genome integrity during DNA replication (Bansbach et al. 2009; Ciccia et al. 2009; Driscoll and Cimprich 2009; Postow et al. 2009; Yusufzai et al. 2009). Immunofluorescent imaging demonstrated that SMARCAL1 accumulates at damaged replication forks due to its interaction with RPA. We now report that SMARCAL1 associates with active, elongating replisomes, and its absence causes MUS81-dependent DNA damage. Significantly, we found that SMARCAL1 exhibits a much broader range of enzymatic activities than previously recognized, including an ability to promote branch migration of Holliday junctions and fork reversal of model replication forks. Concerted fork regression and branch migration coupled to DNA polymerization provides one mechanism to allow DNA damage bypass and replication restart (Petermann and Helleday 2010). SMARCAL1 depletion does not significantly slow the overall rate of DNA replication but is required for efficient DNA replication restart of stalled or collapsed replication forks (Ciccia et al. 2009). Thus, SMARCAL1 may continuously survey replisomes and promote efficient restart of stalled forks through its fork remodeling activity. In the absence of SMARCAL1, slowed or damaged forks are cleaved by MUS81, perhaps as an alternative mechanism of fork repair.

In addition, our results indicate that all SMARCAL1 activities require the HARP2 and SNF2-like ATPase domains. The HARP2 domain is required for DNA binding, and the HARP2-ATPase domains together form the
In contrast to the HARP2 domain, the HARP1 domain makes a modest contribution to the DNA-binding and ATPase activities of SMARCAL1 and is largely dispensable for its annealing, branch migration, and fork regression functions. While vertebrate SMARCAL1 proteins contain two HARP domains, invertebrate SMARCAL1 proteins contain only a single HARP domain adjacent to the ATPase domain, suggesting that only a single HARP domain is essential for its evolutionarily conserved functions.

Our conclusions about the important function of the HARP2 domain are generally consistent with a recent report that found that the HARP domains are important for the annealing helicase activity of SMARCAL1 (Ghosal et al. 2011). However, the Chen group (Ghosal et al. 2011) reported that deleting both HARP1 and HARP2 together did not impair either DNA-binding or ATPase activity despite eliminating the annealing helicase activity. In contrast, our data with both deletion and point mutants clearly point to a requirement for the HARP2 domain for all SMARCAL1 enzymatic functions. We tested multiple proteins purified from both insect and human cells using several different DNA substrates and always found that the HARP2 domain was critical for DNA binding, ATPase activity, strand annealing, and branch migration. We do not have an explanation for this discrepancy.

The ability of SMARCAL1 to efficiently bind to Holliday junctions and model replication forks that lack ssDNA...
regions was unexpected, since SMARCAL1 has very little affinity to dsDNA compared with the optimal substrates containing at least 15 nt of dsDNA and 5 nt of ssDNA. One possibility is that SMARCAL1 captures a small amount of these structures as the dsDNA regions near the junction or fork breath to expose ssDNA. Our data indicate that only a small amount of ssDNA or even just a nick is necessary for SMARCAL1 DNA binding when it is in the context of a gap. Likewise, only a small amount may be needed in the context of these more complicated structures. We also observed no significant specificity of human SMARCAL1 for 3' or 5' recessed junctions. Furthermore, a 5' recessed junction containing a model RNA–DNA primer, as would be found during lagging strand replication, efficiently binds and activates SMARCAL1. This contrasts with a previous report that found a preference for a 3'-hydroxyl recessed end (Muthuswami et al. 2000). The origin of this difference may be because the previous report used a fragment of bovine SMARCAL1, whereas we used full-length human SMARCAL1 in our studies.

The ATP-dependent activity of SMARCAL1 to remodel Holliday junctions and replication forks and prevent DNA damage during S phase is reminiscent of the activities of other proteins, including FANCN, WRN, RAD5, BLM, and HLTF (Constantinou et al. 2000; Ralf et al. 2006; Blastyak et al. 2007; Franchitto et al. 2008; Gari et al. 2008a,b; Opresko et al. 2009; Achar et al. 2011]. All of these proteins are thought to be recruited to damaged replication forks, but it is unclear whether any travel with active forks like SMARCAL1. In contrast to SMARCAL1, none of these proteins contain a HARP domain or exhibit annealing helicase activity. Instead, most are DNA helicases. Thus, the enzymatic mechanisms by which they remodel replication fork structures are likely to be different. Why there are so many different enzymes that can catalyze similar reactions on DNA is unclear. It is possible that some of these enzymes work coordinately at the same damaged fork. In this regard, it is interesting that the loss of WRN, like SMARCAL1, also causes MUS81-dependent fork cleavage (Franchitto et al. 2008], and we and others have found WRN in SMARCAL1 purifications, suggesting a possible physical interaction (Ciccia et al. 2009; data not shown). Coordination of their enzymatic activities might help remodel damaged forks in cells where many other replisome and repair proteins may be present. However, these proteins must also have distinct functions, since inactivating mutations cause different human diseases.

In summary, our data suggest that SMARCAL1 surveys DNA replication forks. When it detects a problem, it uses its DNA-stimulated ATPase motor to remodel the fork by catalyzing strand annealing, branch migration, and fork reversal to promote efficient fork repair. These activities are encoded within the HARP2-SNF2 ATPase domains, which form a functional enzyme flanked by regulatory sequences. Absence of SMARCAL1 forces the use of alternative fork repair mechanisms that involve MUS81-dependent DNA double-strand breaks.

Materials and methods

Cell culture

HEK-293T and U2OS cells were cultured in DMEM supplemented with 7.5% FBS. Si9 cells were cultured in Insect XPRESS medium with 7.5% FBS at 27°C.

Antibodies

The antibodies used were as follows: Flag-M2 (Sigma), γH2AX and GAPDH (Millipore), RPA [Bethyl Laboratories], H3 (Abcam), and MUS81 (Novus). The SMARCAL1 antibody was described previously [Bansbach et al. 2009].

Detection of γH2AX

γH2AX foci were detected by indirect immunofluorescent imaging of fixed U2OS cells 72 h after transfection with siRNA as previously described [Lovejoy et al. 2009].

iPOND

The iPOND technique was performed as described previously [Sirbu et al. 2011]. Briefly, cells were labeled for 10 min with EdU, then treated with 2 mM HU for increasing amounts of time. Alternatively, after the EdU labeling period, 10 μM thymidine was added to the growth medium for 20 min as a "chase" sample. This concentration of thymidine does not block replication but is sufficient to ensure that no additional EdU is incorporated. After cross-linking with formaldehyde and a click reaction to conjugate biotin to the EdU-labeled nascent DNA, protein–DNA complexes were isolated with streptavidin beads, cross-links were reversed, and the eluted proteins were analyzed by immunoblotting. The "no click" control omitted the biotin-azide during the click reaction.
Protein purification

Flag-SMARCAL1, His-SMARCAL1[325–954], HARP1-WF, and HARP2-WF were purified from baculovirus-infected cells essentially as described previously (Yusufzai and Kadonaga 2008) except that cells were lysed in TNT buffer containing 20 mM Tris [pH 7.5], 150 mM NaCl, 0.1 mM EDTA, 1 mM DTT, 0.2 mM PMSE, 1 mg/mL leupeptin, 1 mg/mL apropin, and 0.1% Triton X-100. Proteins for structural studies were purified by Ni-NTA affinity, ion exchange, and gel filtration chromatography. To purify SMARCAL1 proteins from human cells, HEK-293T cells were transfected with pLPCX-Flag-HA-SMARCAL1 plasmids using Lipofectamine 2000 [Invitrogen]. Seventy-two hours after transfection, the cells were lysed in TNT buffer for 30 min on ice. After high-speed centrifugation, the cleared lysates were incubated with Flag-M2 beads [Sigma] for 3 h at 4°C. The beads were washed three times in wash buffer [TNT buffer containing 0.3 M LiCl] and twice in SMARCAL1 buffer [20 mM HEPES at pH 7.6, 20% glycerol, 0.1 mM KCl, 1.5 mM MgCl₂, 0.2 mM EDTA, 1 mM DTT, 0.2 mM PMSE, 0.01% IGEPA CA-630]. The bound proteins were eluted in SMARCAL1 buffer containing 0.25 mg/mL Flag peptide on ice, flash-frozen, and stored at –80°C.

DNA-binding, annealing helicase, and ATPase assays

The gel mobility shift assays for DNA-binding, annealing helicase, and SMARCAL1 ATPase assays were performed as described previously [Yusufzai and Kadonaga 2008] with the following modifications. For the gel mobility shift assay, increasing concentrations of purified SMARCAL1 [0, 0.5, 1, 2 nM final concentrations] were combined with radiolabeled oligonucleotide probe [1 nM final concentration] in binding buffer supplemented with 0.2% IGEPA CA-630. The samples were loaded into a 5% polyacrylamide 1× TBE gel [82 × 28.5 cm, 1 mm thick], and subjected to electrophoresis in 0.5× TBE for 2 h and 30 min at 50 V at 4°C. The gels were dried and quantified using a Molecular Imager FX [Bio-Rad]. DNA-binding reactions were performed at least twice, and a representative experiment is shown with quantitation. For the annealing helicase assay, the topoisomerase I was purchased [Invitrogen], and pBluescript was used as the plasmid substrate. For the ATPase assay, increasing concentrations of oligonucleotides [0, 2, or 8 nM final concentration] were incubated with purified SMARCAL1 [8 nM final concentration] in a final volume of 10 μL, and the reactions were incubated for 30 min at 30°C. The results are presented as the percent of ATP hydrolyzed to ADP during the reaction. ATPase assays were performed a minimum of three times each, and graphs depict means and standard deviation error bars. All oligonucleotide sequences are described in Supplemental Table 1, and all DNA substrates are described in Supplemental Table 2. All figures show a representative experiment from at least two replicates.

Homology modeling

The HARP repeats were identified as evolutionary structural homologs to PUR-α repeats using the FUGUE sequence-structure homology recognition server [Shi et al. 2001]. The HARP2 (amino acids 325–536) homology model was constructed using the crystal structure of PUR-α [Protein Data Bank [PDB] ID 3K44] [Graeber et al. 2009] residues 41–185 as a template. The ATPase model (SMARCAL1 residues 451–856) was generated from residues 455–891 of the SsoRad54 crystal structure [PDB ID 1Z63] [Durr et al. 2005]. In both cases, the SMARCAL1 sequences were threaded onto the crystal structure using Swiss PDB Viewer, and the model was optimized using Swiss Model [http://swissmodel.expasy.org].

SAXS data collection and model building

SAXS data were collected at the SIBYLS beamline at the Advanced Light Source and prepared as described [Hura et al. 2009]. Specifically, SAXS data were collected on SMARCAL1[325–954] in buffer containing 20 mM HEPES [pH 7.6], 200 mM NaCl, 2 mM MgCl₂, 0.5 mM TCEP, 5% glycerol, and 1% sucrose. The protein sample was prepared for SAXS as described [Kazantsev et al. 2011] using a Shodex KW402.5 size exclusion column. The peak fraction was analyzed for SAXS as a 2/3 dilution series starting from 3 mg/mL. Three exposure times [0.5, 1, and 6 sec] were taken at 25°C and 12 keV. Guinier and Kratky analysis was performed as described [Putnam et al. 2007; Rambo and Tainer 2011]. Linearity of the Guinier region for each exposure demonstrated a lack of radiation damage and aggregation [Supplemental Fig. 4D]. SAXS profiles were overlaid, inspected for concentration-dependent scattering, and merged [Hura et al. 2009]. For modeling, the composite scattering curve was generated from data from 1-sec exposures of 2 and 3 mg/mL samples. The maximum dimension [116 Å] was determined using GNOM [Svergun 1992]. Atomistic-based modeling of the SAXS data was achieved with the program BUNCH [Petoukhov and Svergun 2005] using HARP and ATPase homology models. The models were treated as independent domains in a simulated annealing algorithm to determine their relative spatial arrangements. Missing residues between the HARP and ATPase domains [397–450] were modeled as dummy residues as described [Petoukhov and Svergun 2005]. Ab initio modeling was performed with GASBOR using 630 dummy residues. Ten independent modeling runs were performed and averaged [Volkov and Svergun 2003] to produce a final macromolecular envelope. The final model targeted residues 325–856, consistent with a Porod volume of 91,148 Å³ calculated from the SAXS data. The missing C-terminal 99 residues were not included in the modeling based on proteolytic sensitivity of the C-terminus [Supplemental Fig. 4A].

Branch migration and fork regression assays

Oligonucleotide #48 was end-labeled with [γ-32P]ATP and T4 polynucleotide kinase [New England Biolabs] and purified through a G25 column [GE Healthcare]. To prepare tailed or forked intermediates, 250 nM complementary ssDNA oligonucleotides (#48/#54 and #55/#56 for the branch migration, and #48/#50 and #53/#54 for the fork regression) were annealed in 20 mM Tris at pH 7.5, 50 mM KCl, 5 mM MgCl₂, 100 μM [32P]-labeled and 48 nM nonlabeled 250 nM complementary ssDNA oligonucleotides (#48/#54 and #55/#56 for the branch migration, and #48/#50 and #53/#54 for the fork regression) were annealed in 20 mM Tris at pH 7.5, 50 mM KCl, 5 mM MgCl₂, 100 μM [32P]-labeled and 48 nM nonlabeled DNA intermediates were incubated in reaction buffer [40 mM Tris at pH 7.5, 20 mM KCl, 5 mM MgCl₂, 100 μg/mL BSA, 2 mM ATP, 2 mM DTT] for 30 min at 37°C. The DNA substrates were diluted threefold in reaction buffer and mixed with increasing amounts of SMARCAL1 in a 20-μL reaction volume. The reaction was completed for 20 min at 37°C and 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 loaded into 8% polyacrylamide 1× TBE gels [82 × 28.5 cm, 1 mm thick] and subjected to electrophoresis in 1× TBE for 90 min at 80 V at room temperature. The gels were dried and quantified using a Molecular Imager FX [Bio-Rad].

The plasmid-sized replication fork model substrate was generated and purified as described [Blastyak et al. 2007]. Recombinant SMARCAL1 purified from insect cells was incubated with 0.5 nM substrate for 20 min at 37°C in reaction buffer [20 mM Tris at pH 7.5, 50 mM KCl, 5 mM MgCl₂, 100 μg/mL BSA, 2 mM ATP or ATPγS, 1 mM DTT]. The reaction was quenched by the addition of 10 mM ATPγS and 10 mM MgCl₂. One microliter...
(2–20 U, depending on the enzyme) of the indicated restriction enzymes was added to the reaction and further incubated for 30 min at 37°C. The reaction products were then separated on a 6% polyacrylamide gel. The gel was dried and quantified using a Molecular Imager FX (Bio-Rad).

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