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Mechanism of replication fork reversal and protection by human RAD51 and RAD51 paralogs

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ABSTRACT

SMARCAL1, ZRANB3 and HLTF are all required for the remodeling of replication forks upon stress. Using reconstituted reactions, we show that the motor proteins have unequal biochemical capacities, explaining why they have non-redundant functions. Whereas SMARCAL1 uniquely anneals RPA-coated ssDNA, suggesting an initial function in fork reversal, it becomes comparatively inefficient in subsequent branch migration. We also show that low concentrations of RAD51 and the RAD51 paralog complex, RAD51B-RAD51C-RAD51D-XRCC2 (BCDX2), directly stimulate SMARCAL1 and ZRANB3 but not HLTF, providing a mechanism underlying previous cellular data implicating these factors in fork reversal. Upon reversal, RAD51 protects replication forks from degradation by MRE11, DNA2 and EXO1 nucleases. We show that the protective function of RAD51 unexpectedly depends on its binding to double-stranded DNA, and higher RAD51 concentrations are required for DNA protection compared to reversal. Together, we define the non-canonical functions of RAD51 and its paralogs in replication fork reversal and protection.
INTRODUCTION

Homologous recombination (HR) is one of the key cellular pathways required for the maintenance of genome stability. Historically, recombination proteins have been mostly studied in the context of DNA double-strand break (DSB) repair\(^1,2\). The first step in recombinational repair of broken DNA is DNA end resection, where specialized nucleases degrade 5'-terminated DNA strands at the DSB sites, leading to 3' overhangs. Resection is typically initiated by the MRE11 nuclease acting within the MRE11-RAD50-NBS1 (MRN) complex in conjunction with CtIP. The ensemble first nicks 5'-terminated DNA strand near the break end\(^3\). It has the unique capacity to nick past protein blocks, such as stalled topoisomerases or the Ku heterodimer, or past secondary DNA structures, which initiates processing of DNA breaks. The 3'-5' exonuclease of MRN is subsequently believed to degrade DNA between the nick sites and back towards the broken end, resulting in 3' overhangs of limited length\(^4\). The initial short-range resection by MRN-CtIP is in most cases extended by either of two long-range nucleases, EXO1 or DNA2, which functions together with the Bloom (BLM) or Werner (WRN) helicase\(^5\).

Downstream of resection in DSB repair, the ssDNA overhang is first coated by the single-strand DNA (ssDNA) binding protein Replication Protein A (RPA). In the next step, RPA is replaced by RAD51, a key recombinase in eukaryotic cells. The replacement of RPA and loading of RAD51 on the ssDNA overhang is facilitated by a group of proteins termed as recombination mediators, which include the BRCA2 protein, and the RAD51 paralog complexes, either RAD51B-RAD51C-RAD51D-XRCC2 (BCDX2), or RAD51C-XRCC3 (CX3)\(^1,2,6,7,8\). These proteins catalyze the formation of the RAD51 nucleoprotein filament, which has the capacity to identify, pair, and invade homologous double-stranded DNA (dsDNA). In vegetative cells, the homologous DNA is typically the sister chromatid that carries the identical genetic sequence. Once the RAD51 nucleoprotein filament invades homologous DNA, the 3'-terminated end serves as a primer for DNA synthesis that recovers any missing genetic information near the break site. The joint molecule intermediates are subsequently matured and processed to yield two recombined molecules, thus restoring genome integrity in a largely error-free manner. The function of recombination factors in the context of DSB repair is now relatively well understood\(^1,2\).

More recently, it has been discovered that several recombination proteins have separate DSB repair-independent functions in the response to replication stress\(^9,10\). When replicating damaged templates, repetitive DNA sequences or DNA at telomeres, during replication-transcription conflicts or upon the overexpression of oncogenes, the forks can uncouple and the leading strand polymerase may transiently stall\(^11\). Upon prolonged stalling, depending on the cellular context, replication can restart by PRIMPOL-mediated repriming, or the forks can undergo reversal\(^12,13,14\). Fork reversal involves annealing of the two nascent DNA stands yielding a 4-way junction, followed by branch migration\(^15,16,17\). In contrast to DSB repair, the functions of recombination proteins under replication stress remain mostly undefined.
For a long time, fork reversal was thought to be only a pathological process\textsuperscript{18}. More recent data however uncovered that depending on the context and cellular genetic background, fork reversal may be beneficial\textsuperscript{15, 16, 19}. Fork reversal may in fact limit the extent of ssDNA at stalled forks, provide cells time to deal with the respective challenge, and in this way prevent DNA breakage or even enables specific DNA repair. Indeed, several motor proteins with unique capacities to reverse replication forks have been identified, including but not limited to SMARCAL1, ZRANB3 and HLTF\textsuperscript{16, 20}. Depletion of SMARCAL1 and ZRANB3 results in sensitivity to conditions inducing replication stress and enhancing genome instability, indicating that fork reversal catalyzed by these enzymes in wild type cells can be a protective process\textsuperscript{21, 22, 23, 24, 25, 26}.

In certain conditions, uncoupling of leading and lagging strand DNA synthesis may result in the formation of ssDNA gaps, which together with aberrant fork reversal can lead to DNA degradation and genome instability. Gaps of ssDNA on newly synthesized daughter strand or on the apex of the reversed replication fork represent entry points for pathological nucleolytic degradation, unless stabilized by RAD51\textsuperscript{10, 16, 27}. The degradation of nascent DNA can be particularly observed in cells with impaired functions of BRCA1, BRCA2 and a growing number of additional factors, which are believed to recruit and stabilize RAD51 on DNA, or otherwise directly or indirectly inhibit the respective nucleases\textsuperscript{9, 17, 19, 28, 29}. Remarkably, the same pro-DNA end resection nucleases that promote DSB repair, including MRE11 (presumably within the MRN complex), EXO1 and DNA2 were shown to contribute to the pathological nascent DNA degradation in various genetic backgrounds\textsuperscript{28, 30, 31, 32, 33, 34}. Notably, degradation of reversed forks by DNA2 in wild type cells may mediate fork restart\textsuperscript{35}. The function of BRCA2 in fork protection is genetically separable from its function in DSB repair, showing that there are distinct mechanisms at play\textsuperscript{9}. How RAD51 protects nascent DNA remains uncharacterized, but it has been assumed that RAD51 binding to ssDNA on the regressed arm of the reversed forks serves as a barrier against the entry of the resection nucleases\textsuperscript{16, 27}.

Cellular data suggest that the impaired RAD51 function to protect stalled forks in BRCA-deficient cells results in extended degradation of nascent DNA of multiple kilobases in length\textsuperscript{9}. Such DNA degradation is dependent on the proteins implicated in fork reversal, including SMARCAL1, ZRANB3 and HLTF, which suggested that reversed, but not stalled replication forks are primarily subjected to degradation\textsuperscript{9, 27, 30, 31, 36}. However, nascent DNA degradation is not observed upon depletion of RAD51 itself\textsuperscript{17, 31}. This apparently paradoxical observation was explained by a model where RAD51 also promotes fork reversal. Accordingly, in the absence of RAD51, the substrates for DNA degradation, i.e. reversed forks, are not formed. In the absence of the relevant substrate, the function of RAD51 in fork protection becomes irrelevant, explaining why nascent DNA is stable upon depletion of RAD51\textsuperscript{17, 37}. The function of RAD51 in fork reversal, in contrast to DNA protection, is independent of BRCA2\textsuperscript{9, 31}. How mechanistically RAD51 promotes replication fork reversal remains undefined, but it appears to be distinct from its function in canonical recombination\textsuperscript{38}. Similarly as with BRCA2, the function of RAD51 in DSB
repair and fork metabolism/modulation is genetically separable\textsuperscript{38}. Recently, the BCDX2 RAD51 paralog complex was found to promote fork reversal alongside RAD51 in cellular assays\textsuperscript{39}.

The seemingly paradoxical situation where RAD51 may both promote and prevent DNA degradation highlights the difficulties explaining cellular phenotypes and thus warrants the need for a more direct understanding of the underlying mechanisms\textsuperscript{37}. Here we use reconstitution biochemistry, which allows us to study elements of the reactions governing the metabolism of challenged replication forks in isolation. We show that the fork remodelers SMARCAL1, ZRANB3 and HLTF have different substrate preferences, suggesting that they may catalyze different reactions and thus act at distinct steps during fork remodeling, explaining cellular data why these factors have non-redundant functions. We show that RAD51 and the BCDX2 RAD51 paralog complex directly promote the motor-driven strand annealing activity of SMARCAL1 and ZRANB3, which identifies the mechanism by which RAD51 and BCDX2 may facilitate fork reversal. We show that compared to fork reversal, higher RAD51 concentrations are required for DNA protection against MRE11-RAD50 exonuclease, MRN endonuclease, and EXO1 and DNA2 nucleases. In contrast to the current models, we demonstrate that the protective function of RAD51 also involves its capacity to bind dsDNA. Together, our data provide comprehensive insights into the mechanisms underlying the function of RAD51 and RAD51 paralogs in the metabolism of challenged replication forks.
RESULTS

SMARCAL1, ZRANB3 and HLTF have unequal biochemical activities

SMARCAL1, ZRANB3 and HLTF have all been implicated in replication fork reversal in vitro and in vivo\(^\text{22,40,41}\). The loss of either of these enzymes was shown to abolish nascent DNA degradation in BRCA1/2-deficient cells, suggesting that these factors may act in a non-redundant manner to promote fork reversal\(^\text{19}\). To better understand the function of these fork remodelers, we expressed and purified SMARCAL1, ZRANB3 and HLTF from insect Sf9 cells (Fig. 1a). All three translocases hydrolyzed ATP, as expected, with SMARCAL1 showing the highest specific activity, followed by HLTF and ZRANB3 (Supplementary Fig. 1a). We next set out to compare the relative activities of these motor proteins in biochemical assays mimicking elements of fork reversal. We first used oligonucleotide-based DNA substrates resembling stalled replication forks with ssDNA gaps either in the leading or the lagging DNA strand (Fig. 1b). We observed, as reported previously, that SMARCAL1 in the presence of the ssDNA binding protein RPA was more efficient on forks with leading strand gaps, as opposed to ZRANB3, which prefers RPA on lagging strand gaps\(^\text{42}\) (Fig. 1c). Using the leading strand gap substrate, SMARCAL1 was also more efficient than HLTF (Fig. 1c), while the three translocases exhibited similar specific activities on the lagging strand gap substrate (Fig. 1c). In contrast to the activities of SMARCAL1 and ZRANB3 that are regulated by RPA\(^\text{42}\), the function of HLTF was not RPA sensitive (Fig. 1d). During fork reversal, the initial annealing of the nascent DNA strands leads to the formation of a 4-way junction (Holliday junction, HJ), which is further branch migrated by the motor proteins, leading to reversed forks of up to several kilobases in length\(^\text{16}\). Using a mobile HJ substrate to assay for branch migration, we observed that SMARCAL1 was in contrast the least efficient enzyme, essentially incapable of branch migration at physiological (150 mM) salt concentrations. However, under less restrictive conditions in lower salt, the branch migration activity of SMARCAL1 was readily detected (Fig. 1f, right panel). Instead, both ZRANB3 and HLTF were highly and comparably efficient in branch migration at 150 mM salt (Fig. 1e, f).

The activity of the SMARCAL1 and ZRANB3 enzymes was first analyzed by a topoisomerase-coupled assay that monitors the annealing of RPA-coated ssDNA bubbles in plasmid DNA, which can be observed as changes in DNA topology\(^\text{25,43,44}\). Such activity is thought to mimic the initial stages of fork remodeling. Both SMARCAL1 and ZRANB3 were shown to anneal the RPA-coated DNA bubbles as a result of their motor functions, as ATP hydrolysis is required for this reaction. However, the specific activities of SMARCAL1 and ZRANB3 have not been compared. We observed that SMARCAL1 was comparably efficient to ZRANB3, while HLTF showed much smaller capacity to anneal DNA in this assay (Fig. 1g, h).
Taken together, our data indicate that SMARCAL1, ZRANB3 and HLTF possess quite different biochemical activities and substrate preferences. Our results support a model positing that fork reversal is not catalyzed by a single enzyme in a processive manner, but that it is rather a dynamic process that involves the sequential engagement of several factors.

SMARCAL1 specifically anneals RPA-coated ssDNA

Several helicases, such as members of the RecQ family, were reported to anneal two ssDNA molecules, but the reactions were inhibited by RPA\textsuperscript{45,46}. Considering that cellular RPA concentration is thought to be sufficient to coat all ssDNA in most cases, the physiological relevance of these observations remains unclear. In this regard, the reported activities of the RecQ family members differ from the canonical RecO/RAD52 family annealing proteins, which anneal RPA-coated ssDNA to promote homologous recombination\textsuperscript{47,48}. The observation that SMARCAL1 and ZRANB3 are efficient in annealing bubbled DNA coated with RPA prompted us to investigate whether the enzymes can anneal two ssDNA molecules similarly as RAD52. The annealing of the bubbled DNA in the topoisomerase-coupled assays could result from an annealing activity \textit{per se}, or can be a consequence of the dsDNA translocase activity rezipping the bubble from the side. To distinguish between these two possibilities, we set to define the function of the fork remodelers in complementary ssDNA annealing. We observed that SMARCAL1, ZRANB3 and HLTF were all able to anneal free ssDNA, which can be explained by multiple ssDNA binding sites on a single enzyme or by enzyme oligomerization, which can bring multiple ssDNA molecules to close proximity, stimulating their annealing. However, in the presence of RPA, the ssDNA annealing by ZRANB3 and HLTF was strongly reduced, similarly as observed with the RecQ family helicases\textsuperscript{45}. In contrast, ssDNA annealing by SMARCAL1 remained highly proficient in the presence of RPA (Fig. 2a, b, c). Unlike RPA, the ssDNA annealing capacity of SMARCAL1 was abrogated when the ssDNA was pre-coated with mitochondrial SSB, showing that the annealing of ssDNA by SMARCAL1 is allowed in the presence of RPA in a specific manner (Fig. 2d). The annealing activity of SMARCAL1 was also observed in the absence of ATP, or when using the motor-dead SMARCAL1 (D549A, E550A, SMARCAL1-HD) variant showing that this particular activity is not ATPase dependent (Supplementary Fig. 2a), as is the case of RAD52\textsuperscript{48}. Therefore, the annealing of two ssDNA molecules by SMARCAL1 mechanistically differs from the annealing of bubbled DNA in the topoisomerase coupled assays, which largely require ATP hydrolysis\textsuperscript{25,43,44}. The N-terminal region of SMARCAL1 contains a previously defined RPA-binding site, the integrity of which is required for the recruitment of SMARCAL1 to DNA damage sites, and to direct SMARCAL1 to substrates with RPA-coated ssDNA gaps\textsuperscript{21} (Fig. 2e). To test for the requirement for direct interaction between SMARCAL1 and RPA in ssDNA annealing, we expressed and purified SMARCAL1 (ΔN), lacking the RPA interaction domain (Fig. 2f). The truncated SMARCAL1 was fully proficient in DNA
branch migration in the absence of RPA and identical to wild type SMARCAL1 as an ATPase (Supplementary Fig. 2b, c). However, the mutant was inefficient in ssDNA annealing in the presence of RPA, showing that the direct interaction of SMARCAL1 with RPA is essential for this activity (Fig. 2g). Our results reveal that SMARCAL1 possesses a strand annealing activity similar to that of the RAD52 protein family, which also rely on specific interaction with RPA. We suggest that such activity may be employed during the very initial steps of fork reversal, when the daughter ssDNA molecules are separated from the parental strands and anneal with each other. These results further underline the mechanistic differences between the fork remodeling enzymes SMARCAL1, ZRANB3 and HLTF.

**RAD51 and BCDX2 paralogs promote motor-driven strand annealing activity of SMARCAL1 and ZRANB3 but not HLTF**

Challenged DNA replication forks may undergo reversal, and reversed replication forks must be subsequently protected by RAD51 to prevent pathological nascent DNA degradation. However, RAD51, along with the RAD51 paralog BCDX2 complex, were also paradoxically implicated in promoting fork reversal, through a yet unknown mechanism. To elucidate whether the function of RAD51 and the RAD51 paralogs in fork remodeling may be direct, we next expressed and purified RAD51 and the BCDX2 complex (Fig. 3a). The BCDX2 complex was obtained upon co-expression of all subunits, as the preparation of the individual proteins resulted in poor yields and solubility. The BCDX2 complex did not aggregate, bound ssDNA and very weakly hydrolyzed ATP, as observed previously.

We next set out to test whether RAD51 and BCDX2 complex affect the strand annealing and motor activities of SMARCAL1. To this point, we used the established topoisomerase-coupled assay. Strikingly, we observed that low concentrations of RAD51 and the BCDX2 complex promoted bubbled DNA annealing by SMARCAL1, while none of these co-factors had a notable capacity to mediate DNA annealing per se without SMARCAL1, even at much higher concentrations (Fig. 3b, Supplementary Fig. 3e, f). Additionally, controls where no ATP was used or helicase-dead SMARCAL1 variant replaced the wild type protein largely abolished DNA annealing, indicating that a large proportion of the relaxed DNA signal in the assay can be linked to the ATP hydrolysis-driven translocation activity of SMARCAL1 (Fig. 3b, Supplementary Fig. 3g). Similarly to SMARCAL1, we observed that RAD51 and the BCDX2 paralogs promoted the annealing capacity of ZRANB3. As above with SMARCAL1, we observed that RAD51 and BCDX2 could both promote ZRANB3 independently of each other (Fig. 3c). Only limited changes in DNA topology were observed when using helicase-dead ZRANB3 and all co-factors, demonstrating that the majority of the signal in the assay can be linked to the motor activity of ZRANB3 leading to DNA annealing (Supplementary Fig. 3h).
Differently from SMARCAL1 and ZRANB3, HLTF was not stimulated by either RAD51 or BCDX2 (Fig. 3d). The RAD51C-XRCC3 (CX3) complex could also promote DNA annealing with both SMARCAL1 and ZRANB3 (Supplementary Fig. 3i, j, k), despite it has not been found to promote fork reversal in vivo\(^{39,51}\). While the CX3 complex may not be efficiently recruited to physiological substrates in cells, the experiments were informative as they revealed that the critical component of the paralogs may be RAD51C, which is present in both BCDX2 and CX3 complexes. Together, we show that RAD51 and BCDX2 promote the translocation-driven annealing of RPA-coated bubbled DNA by SMARCAL1 and ZRANB3, suggesting that the co-factors may have a direct role in fork reversal to stimulate the DNA translocases.

SMARCAL1 and ZRANB3 physically interact with RAD51 and BCDX2

To test whether the functional interplay between SMARCAL1 and ZRANB3 with RAD51 and BCDX2 may involve direct physical interactions, we immobilized RAD51 and performed pulldown experiments with the co-factors. We observed that BCDX2 interacted with RAD51 (the RAD51B component was detected), as described previously\(^{52}\). Importantly, we found that both SMARCAL1 and ZRANB3 also interacted with RAD51 (Fig. 4a). We next immobilized the BCDX2 complex, and observed a direct interaction with SMARCAL1 and ZRANB3, as detected by Western blotting and silver staining (Fig. 4b, c). These results collectively suggest that the interplay of SMARCAL1 and ZRANB3 with RAD51 and BCDX2 likely involves direct physical interactions.

Point mutation in SMARCAL1 disrupts physical and functional interactions with RAD51

We next set out to define motifs in SMARCAL1 and ZRANB3 that mediate the interactions with RAD51 and BCDX2. We failed to identify an interaction motif with BCDX2, but we found regions in SMARCAL1 mediating the binding to RAD51. Physical and functional interactions between RAD51 and many of its co-factors, such as BRCA2, BARD1, MMS22L, RECQL5, SWSAP1 and FINGL1 are mediated by the FXXA motif\(^{53,54,55,56,57,58}\). We identified such a motif in SMARCAL1, which is conserved in evolution (Fig. 4d, Supplementary Fig. 4a). The FXXA motif is positioned ahead of the conserved αI SNF2 family ATPase domain (Supplementary Fig. 4b). The mutation of phenylalanine 446 into alanine (F446A) in SMARCAL1 disrupted the physical interaction with RAD51 (Fig. 4e, f). In contrast, disruption of F439, which is part of a less conserved FXXA sequence in human SMARCAL1 upstream of F446, did not impair the interaction (Fig. 4f, Supplementary Fig. 4a). We note that SMARCAL1 F446A variant per se was very similar to wild type SMARCAL1 in its fork reversal and ATPase capacities in vitro and retained its physical interaction with the BCDX2 complex (Supplementary Fig. 4c, d and Fig. 4g). ZRANB3 contains a phenylalanine at the analogous position to
SMARCAL1 ahead of the ATPase domain. The phenylalanine however does not conform to the FXXA motif, and the F47A substitution mutant retained its capacity to interact with RAD51 and was impaired in its ATPase activities (Supplementary Fig. 4e-h). We next found that mutation F736A in ZRANB3 disrupted interaction with RAD51, however it is likely that the mutation affected the fold of the substrate recognition domain, as it likewise abolished the biochemical activities of ZRANB3 and may thus not represent a direct interaction motif (Supplementary Fig. 4i-k). Due to the impact of this mutation on the activities of ZRANB3 per se, we could not test for the physiological relevance of the interaction with RAD51.

To investigate the physiological relevance of the disrupted physical interaction between SMARCAL1 and RAD51, we used MCF10A SMARCAL1-KO cells, which were complemented with either wild type SMARCAL1 or the F446A variant (Fig. 4h). Following replication stress induced by hydroxyurea, it was previously demonstrated that SMARCAL1-mediated fork reversal can lead to nascent DNA degradation, as long as the nascent DNA is not protected by RAD51. The nascent DNA degradation is evident in cells lacking BRCA1 or BRCA2, which may be required to recruit, load or stabilize RAD51. In agreement with previous data, we observed extensive nascent DNA degradation in BRCA1-depleted SMARCAL1 KO cells reconstituted with wild type SMARCAL1 (Fig. 4h). In contrast, such extensive DNA degradation was not observed in BRCA1-depleted SMARCAL1 KO cells reconstituted with empty vector (no SMARCAL1), and it was partially attenuated in cells with SMARCAL1 F446A, which was expressed at levels comparable to wild type (Supplementary Fig 4l). Taking into consideration that nascent DNA degradation in BRCA-deficient cells requires the fork reversal activity of SMARCAL1, our results suggest that SMARCAL1 F446A, which does not interact with RAD51, might display a defective fork reversal activity in mammalian cells.

High concentrations of RAD51 protect DNA from degradation by MRE11, EXO1 and DNA2 nucleases

In homologous recombination, the processing of DSBs is initiated by short-range resection that involves the endo- and exonuclease activities of MRN and CtIP, followed by the long-range pathways catalyzed by EXO1 and/or DNA2-BLM/WRN. The same nucleolytic pathways were shown to degrade nascent DNA at stalled replication forks, with relative contributions of the nucleases dependent on conditions and genetic background. To investigate the function of RAD51 in DNA protection, we reconstituted DNA end resection reactions without or with various concentrations of RAD51. We observed that RAD51 inhibited all DNA end resection reactions tested, including the exonuclease activity of MRE11-RAD50, the endonuclease of MRE11-RAD50-NBS1 in conjunction with phosphorylated CtIP (Fig. 5a-c), and the long-range pathways of EXO1 and DNA2-WRN (Fig 5f-k). These experiments allowed us to make several conclusions.
First, higher RAD51 concentrations were in general required for DNA protection compared to motor-driven strand annealing. Approximately 300 nM to low micromolar RAD51 was required for a robust protection against nucleolytic degradation, depending on the substrate and the respective nuclease analyzed. The RAD51 concentrations required for the inhibition of the endonuclease of MRN-CtIP were similar to those required to inhibit the nonspecific NspI endonuclease, which may be used as a read-out for RAD51 filament formation (Fig. 5a, b and Supplementary Fig. 5a). These results suggest that efficient inhibition of DNA end resection occurs under conditions permissive for stable RAD51 filament formation.

Second, the DNA affinity of RAD51 directly corresponds to its efficacy in inhibiting DNA degradation. To this point, we used the RAD51 variants that differ in their capacity to bind DNA. The tightly binding RAD51-KR mutant generally inhibited resection more efficiently than wild type RAD51, while the poorly DNA binding RAD51 variants KA, YA and TP were largely deficient in protection (Fig. 5a-d, h, i). Furthermore, the affinity of RAD51 to DNA depends on the presence of ATP. ATP binding stimulates DNA binding by RAD51, while ATP hydrolysis leads to RAD51 dissociation from DNA. Correspondingly, the highest DNA binding affinity of RAD51 is observed in buffers with the non-hydrolysable ATP analog ATP-γ-S. In accord with the modulation of DNA binding affinity of RAD51 by the nucleotide co-factors, we observed the strongest DNA protection by RAD51 when reactions contained ATP-γ-S, followed by ATP, and the weakest protection in reactions lacking ATP (Fig. 5e-g). These latter experiments could only be performed with resection nucleases that do not require ATP, such as the exonuclease of MR and the exonuclease of EXO1.

Third, we did not find any apparent specific functional interactions between RAD51 and the DNA end resection nucleases. To this point, we compared the capacity of human and yeast RAD51/Rad51 to protect DNA against the human and yeast resection nucleases and nuclease complexes (Supplementary Fig. 5b-h). We observed that human RAD51 was generally more efficient than yeast Rad51, which may suggest some degree of specificity. However, human RAD51 was also more efficient in protecting DNA against non-specific nucleases Exo-III and NspI (Supplementary Fig 5i, j). These experiments suggested that the higher efficacy of human RAD51 compared to yeast Rad51 to protect DNA is not due to specific functional interactions between human RAD51 and the nuclease complexes, but may be rather due to a higher DNA affinity or protection capacity of human RAD51 per se.

The function of RAD51 in DNA protection corresponds to its capacity to bind double-stranded DNA

BRCA2 is required for nascent DNA protection by RAD51. Interestingly, the regions of BRCA2 required for DNA protection are distinct from those needed for homologous recombination and RAD51 loading onto RPA-coated ssDNA. The BRCA2 BRC repeats are needed for homologous
recombination, and in biochemical assays they were shown to enhance the binding of RAD51 to RPA-coated ssDNA, while they reduce the capacity of RAD51 to bind dsDNA\textsuperscript{9, 64, 65}. The BRC repeats are however not involved in DNA protection upon replication stress. Rather, the C-terminal RAD51 binding site is required for DNA protection\textsuperscript{9}. The biochemistry of the C-terminal site is less understood, but it was demonstrated that it facilitates RAD51 binding also to dsDNA\textsuperscript{66}. These results prompted us to investigate whether RAD51 binding to ssDNA overhangs at DNA ends, as indicated in current models\textsuperscript{16}, indeed explains its protection function.

To investigate whether RAD51 binding to ssDNA or dsDNA explains its function in DNA protection, we prepared blunt-ended, 5'-overhanged and 3'-overhanged substrates. The MR exonuclease was inhibited by RAD51 to the same extent irrespectively of the presence of the overhang (Fig. 6a, b). This showed that the overhang is not required to assure protection against MRE11, and infers that the protection is instead dependent on RAD51 binding to the dsDNA part of the substrate. The same result was observed with EXO1. Although EXO1 shows a clear preference to process 5'-recessed strands and is least efficient on 5'-overhangs\textsuperscript{67}, RAD51 inhibited DNA degradation to a similar extent with all structures tested (Fig. 6c, d), suggesting that overhang is not needed for protection. Likewise, the MRN-CtIP endonuclease clips dsDNA, and we showed that a variety of protein blocks promote such activity, as long as the blocks are located at DNA ends or DNA overhangs. Our observation that RAD51 blocks the MRN-CtIP endonuclease (Fig. 5a, b) again suggests that the inhibitory function is caused by RAD51 binding to dsDNA. Together, these results demonstrate that the binding of RAD51 to dsDNA is responsible for its inhibitory effect on the resection nucleases, at least in vitro. We show that the function of RAD51 to protect DNA from nucleolytic degradation is structural, and directly corresponds to the affinity of RAD51 to bind dsDNA. Tightly-bound RAD51 filaments then serve as a non-specific barrier against DNA degradation. Our results challenge the current model suggesting that the binding of RAD51 to ssDNA overhangs at the apex of the reversed fork structure explains its function in protection\textsuperscript{16, 27, 67}. Our data instead suggest that the capacity of RAD51 to bind dsDNA is relevant for its function in DNA protection, which is ultimately responsible for the maintenance of genome stability during replication stress (Fig. 7).

DISCUSSION

Here we used biochemistry to study the function of the replication fork remodelers SMARCAL1, ZRANB3 and HLTF, their regulation by RAD51 and RAD51 paralogs, as well the interplay of RAD51 with nucleases implicated in pathological DNA degradation upon replication stress.

SMARCAL1, ZRANB3 and HLTF have non-redundant activities
Depletion of either SMARCAL1, ZRANB3 or HLTF brings about a profound defect in replication fork reversal, as observed by electron microscopy, or by proxy methods scoring for e.g., nascent DNA degradation in various genetic backgrounds upon stress. To better define the function of fork remodelers, we compared here the specific activities of the fork remodelers on various substrates mimicking elements of fork reversal. We observed that SMARCAL1, but not ZRANB3 or HLTF, has a unique capacity to anneal RPA-coated ssDNA, a function reminiscent of the RAD52 protein family.

The annealing function of SMARCAL1 depends on the RPA interaction motif within the N-terminus of SMARCAL1. We hypothesize that such annealing function might be relevant during the initial annealing of the displaced daughter strands during the early steps of fork reversal. The annealing activity of SMARCAL1, similarly to RAD52, does not involve ATPase activity. Previously, the function of SMARCAL1 and ZRANB3 was monitored in assays scoring for the annealing of bubbled DNA within circular plasmid. Such activity, in contrast to annealing of RPA-coated ssDNA oligonucleotides, is dependent on the motor activities of the remodelers. We show that in contrast to SMARCAL1 and ZRANB3, which had similar activities, HLTF was largely inactive in this assay.

Using oligonucleotide-based substrates mimicking fork reversal, it has been previously reported that SMARCAL1 and ZRANB3 have opposing preferences with respect to whether RPA is located on the leading or the lagging DNA strand. We show that in contrast, the activity of HLTF does not appear to be affected by RPA. In branch migration, using 4-way junction substrates without ssDNA, HLTF and ZRANB3 were the most active enzymes, while SMARCAL1 was the least efficient. Our experiments demonstrated that the fork remodelers possess quite different specific activities with respect to the substrates used. The data support model where fork remodeling is not catalyzed by a single enzyme in a processive manner, but it is rather a process with various remodelers acting in a distributive manner, depending on the nature of the DNA intermediate and the substrate preference of the respective remodeler. Such model would explain the non-redundant relationship of remodeler in fork reversal.

**Novel biochemical functions for RAD51 and the paralog complex BCDX2 in promoting DNA translocases**

Previous cellular data suggested that RAD51 and the BCDX2 complex promote fork reversal, but the underlying mechanism was not clear. The function of RAD51 in fork remodeling was shown to be genetically separable and thus different from its canonical role in homologous recombination. Specifically, the strand exchange function of RAD51 was dispensable, pointing at a potential structural function. We show here that RAD51 and the RAD51 BCDX2 paralog complex stimulate the strand annealing and branch migration activities of SMARCAL1 and ZRANB3, two of the key enzymes implicated in fork reversal. SMARCAL1 and ZRANB3 were stimulated when RAD51 concentration was too low to support a nucleoprotein filament formation.
In accord with a recent cellular study that identified a function of BCDX2 in promoting fork reversal\cite{39}, we find that the paralog complex also directly stimulates SMARCAL1 and ZRANB3. Unexpectedly, RAD51 and BCDX2 stimulated SMARCAL1 and ZRANB3 independently of each other, as we observed mostly additive effects when combined. The function of the RAD51 paralogs, such as the BCDX2 complex, in homologous recombination remains poorly defined. Some reports suggest a joint function for the paralogs and RAD51. Specifically, BCDX2 was shown to have a classical recombination mediator function to load RAD51 on RPA-coated ssDNA\cite{6} to remodel RAD51 filaments for activation\cite{71}, or to make them more resistant against disruption\cite{72}. However, RAD51-independent function of the RAD51 paralogs were also identified in cellular studies, such as in the single-strand annealing pathway of DSB repair\cite{73}, and BCDX2 was also found to physically and functionally associate with the HELQ helicase\cite{74}. The function of BCDX2 to promote SMARCAL1/ZRANB3 described here \textit{in vitro} also does not require RAD51.

The interplay of RAD51 and paralogs in promoting SMARCAL1 and ZRANB3 involves physical interactions

RAD51 and BCDX2 did not stimulate HLTF, a third enzyme shown to catalyze fork reversal, suggesting a specificity in the interplay of SMARCAL1 and ZRANB3 with RAD51 and BCDX2. In accord, we found that RAD51 and BCDX2 physically interact with SMARCAL1 and ZRANB3. We could then map the RAD51 interaction site in SMARCAL1 and constructed a single point mutant (SMARCAL1-F446A) that disrupted the physical interaction with RAD51. The SMARCAL1 mutant was not impaired in its activities \textit{per se}, but was deficient in promoting nascent DNA degradation in BRCA1-deficient cells, a process that requires the fork reversal activity of SMARCAL1, supporting the idea that the identified interplay of SMARCAL1 and RAD51 is physiologically relevant.

The function of RAD51 in DNA protection is largely non-specific

Downstream of fork reversal, RAD51 protects nascent DNA from degradation against nucleases. We reconstituted both the endo- and exonuclease activities of MRE11, as well as EXO1 and DNA2, which were all implicated in nascent DNA degradation, depending on the cellular background. We then tested for the effect of RAD51 on the individual nucleolytic pathways and observed that RAD51 inhibited them all to a similar extent. The level of inhibition was comparable when we used other non-cognate yeast or bacterial nucleases. Interestingly, human RAD51 was somewhat more efficient in DNA protection compared to yeast Rad51, but this difference was observed in conjunction with both the cognate human as well as with the non-specific nucleases, suggesting that there is no apparent functional interaction between RAD51 and the resection nucleases. RAD51 therefore inhibits DNA degradation in a
largely non-specific manner, as a physical barrier on DNA. The concentrations of RAD51 required for DNA protection were higher than those promoting fork remodeling, in accord with cellular data\textsuperscript{77}. DNA protection assays with RAD51 and non-specific nucleases are used as a proxy for RAD51 filament formation. Therefore, we conclude that RAD51 filament formation is likely prerequisite for DNA protection.

**RAD51 protects nascent DNA upon binding to dsDNA**

Contrary to current models, we provide evidence that the binding of RAD51 to dsDNA (as opposed to ssDNA) may be crucial for its function in DNA protection (Fig. 7). Eukaryotic RAD51 has a similar affinity to both ssDNA and dsDNA\textsuperscript{61,76,77}. In homologous recombination, RAD51 needs to bind ssDNA to form an active nucleoprotein filament. The binding of RAD51 to dsDNA is instead inhibitory, and several recombination factors facilitate RAD51 binding specifically to ssDNA\textsuperscript{57,65,77,78}. The physiological relevance of the high affinity RAD51 binding to dsDNA is not known. The strong dsDNA binding capacity of eukaryotic RAD51 is somewhat paradoxical, because *E. coli*’s RecA binds preferentially to ssDNA, demonstrating that stable dsDNA binding is not strictly associated with DNA strand exchange activity\textsuperscript{77,78}. Because of the recombination paradigm, but largely without direct evidence, it has been assumed that RAD51 binding to ssDNA at the apex of the reversed fork is responsible for DNA protection\textsuperscript{16,27}. Our data challenge this model, and instead suggest that RAD51 binding to the dsDNA part of the reversed fork is primarily responsible for its protection capacity.

We showed that the MRE11-RAD50 or EXO1 exonucleases were inhibited by RAD51 to a similar extent when using blunt-ended dsDNA or overhanded substrates, demonstrating directly that the presence of the overhang was not relevant for protection, at least in the reconstituted assay. RAD51 binding to dsDNA also fully explains its protective function against the MRN-CtIP endonuclease ensemble. MRN-CtIP only nicks dsDNA, even on overhanded substrates\textsuperscript{3,79}. Binding of proteins to overhangs, including non-cognate factors such as streptavidin, is not inhibitory for MRN-CtIP\textsuperscript{79}. RAD51 however clearly inhibits the MRN-CtIP endonuclease when using blunt-ended dsDNA, suggesting that dsDNA is the relevant substrate for RAD51 binding with respect to its protective function. We therefore propose that RAD51 inhibits the resection nucleases due to its binding to dsDNA. This model better explains the protective function of RAD51 in vivo, as it applies for reversed forks with any type of DNA structure at the end of the reversed arm, and does not rely on the presence of an overhang to mediate DNA protection. Electron microscopy analyses revealed that the regressed arms of the reversed forks are mostly composed of dsDNA, irrespectively of the conditions tested\textsuperscript{17}. Cell-based assays further indicated that fork degradation can reach multiple kilobases in length, which goes beyond the length of the regressed arm of the reversed fork\textsuperscript{17,31}. Therefore, dsDNA is clearly being degraded at challenged replication forks, and RAD51 is involved in its protection.
One of the best-known co-factors of RAD51 in DNA protection is BRCA2. BRCA2 functions as a recombination mediator in homologous recombination to load RAD51 on RPA-coated ssDNA, and additionally has the capacity to channel RAD51 to ssDNA away from dsDNA. This particular pro-recombination function depends on the BRC repeats of BRCA2, and could be largely recapitulated with only the short BRC4 peptide. However, the BRC repeats of BRCA2, despite being essential for homologous recombination, are dispensable for fork protection. Instead, the protective function of BRCA2 depends on its C-terminal RAD51 binding site. Strikingly, this site was previously demonstrated to strongly facilitate RAD51 binding to dsDNA. How the site affects ssDNA versus dsDNA binding is not known. The capacity of the BRCA2 C-terminus to promote RAD51 binding to dsDNA thus also agrees with our direct observation that RAD51 binding to dsDNA is critical for DNA protection in vitro. A number of additional factors including BRCA1 are also required for DNA protection. Whether they affect RAD51 binding to dsDNA, or facilitate DNA protection because of a different mechanism, such as upon RAD51-independent DNA binding or through a direct inhibition of the nucleases, remains an interesting area of investigation for future studies. Together, our data suggest that the dsDNA binding capacity of RAD51 may have evolved in conjunction with its function to prevent pathological DNA degradation at the replication fork.
Materials and Methods

Cloning, expression and purification of recombinant proteins

Human SMARCAL1, ZRANB3. Recombinant FLAG-SMARCAL1, FLAG-ZRANB3 and their variants were expressed in insect Spodoptera frugiperda 9 (S/9) cells and purified by affinity chromatography, using pFastBac-FLAG-SMARCAL1 and pFastBac-FLAG-ZRANB3 expression constructs. Point mutagenesis of the corresponding DNA sequences was carried out by QuikChange II site-directed mutagenesis kit (Agilent Technologies), and the proteins were expressed and purified similarly as the wild type counterparts. Primers used for cloning and mutagenesis are listed in Table S1.

Human HLTF. The HLTF sequence was codon-optimized for S/9 insect cells and synthesized by Synbio Technologies, and cloned using NheI and XmaI sites (New England Biolabs) into pFB-2xMBP-CtIP-His81 to create pFB-2xMBP-HLTFco-His, replacing the CtIP sequence with that of HLTF. The bacmids and baculoviruses were prepared according to manufacturer’s instructions (Bac-to-Bac system, Life Technologies). S/9 cells were transfected using a Trans-IT insect reagent (Mirus Bio). For protein production, S/9 insect cells were seeded at 0.5 x 10⁶ cells per ml 16 hours before infection. The cells were then infected with respective baculoviruses and incubated for 52 hours at 27 °C with constant agitation. Cells were harvested (500 g, 10 minutes) and washed once with ice cold Phosphate buffer saline (PBS). The cell pellets were snap frozen in liquid nitrogen and stored at -80 °C. All the subsequent steps were carried out on ice or at 4 °C. The pellets were resuspended and incubated for 20 minutes with continuous stirring in 3-volumes of lysis buffer (50 mM Tris-HCl pH 7.5, 1 mM dithiothreitol [DTT], 5 mM beta-mercaptoethanol [β-ME], 1 mM phenylmethylsulfonyl fluoride [PMSF], 1:400 [v/v] protease inhibitor cocktail [Sigma, P8340], 30 μg/ml leupeptin [Merck]). Next, 50% glycerol was added to reach a final concentration of ~16% to the cell extract, followed by 6.5% volume of 5 M NaCl (final concentration 305 mM), and further incubated for 30 minutes with continuous stirring. The cell suspension was centrifuged for 30 minutes at 48,000 g to obtain soluble extract. The supernatant was transferred to tubes containing pre-equilibrated amylose resin (New England Biolabs, 4 ml per liter of S/9 culture) and incubated for 1 hour with continuous rotation. The resin was collected by spinning at 2,000 g for 2 minutes and washed extensively batch-wise and also on a disposable 10 ml column (ThermoFisher) with amylose wash buffer (50 mM Tris-HCl pH 7.5, 1 mM β-ME, 1 mM PMSF, 10% glycerol, 1 M NaCl). The final wash was performed at 300 mM NaCl. Protein was eluted with amylose elution buffer (50 mM Tris-HCl pH 7.5, 5 mM β-ME, 1 mM PMSF, 10% glycerol, 300 mM NaCl, 10 mM maltose [Sigma]) and the total protein concentration was estimated by Bradford assay. To cleave off the maltose-binding protein (MBP) tag, 1/6 (w/w) of PreScission Protease, with respect to total protein concentration in the eluate, was added and incubated for 1 hour at 4 °C. The sample was then
supplemented with 10 mM imidazole and further passed through pre-equilibrated (amylose elution buffer supplemented with 10 mM imidazole) Ni-NTA agarose resin (Qiagen) on a disposable column for 1 hour in flow. The Ni-NTA resin was washed 4-times with Ni-NTA wash buffer (50 mM Tris-HCl pH 7.5, 5 mM β-ME, 1 M NaCl, 10% glycerol, 1 mM PMSF, 40 mM imidazole). Prior to elution, the protein was washed once with the same Ni-NTA wash buffer as above but with 150 mM NaCl. Protein was eluted in the same buffer supplemented with 300 mM imidazole, and subsequently dialyzed (50 mM Tris-HCl pH 7.5, 5 mM β-ME, 100 mM NaCl, 10% glycerol, 0.5 mM PMSF), sub- aliquoted, snap frozen and stored at -80 °C for later use.

**Human RAD51 paralogs BCDX2 and CX3.** Sequences for human RAD51 paralogs (RAD51B, RAD51C, RAD51D, XRCC2 and XRCC3) were codon-optimized for expression in Sf9 cells and synthesised by Synbio Technologies. FLAG-RAD51B and 10xHis-RAD51C were cloned into a pFB dual expression vector (ThermoFisher). The multiple cloning site 1 was utilized for FLAG-RAD51B using BamHI and NotI cloning sites and the multiple cloning site 2 was used for 10xHis-RAD51C employing the XmaI and NheI restriction sites, to create pFB-FLAG-RAD51Bco-10xHis-RAD51Cco. RAD51D and XRCC2 were similarly cloned into the same sites, respectively, to obtain pFB-RAD51Dco-XRCC2co. XRCC3 was synthesized as BamHI-FLAG-XRCC3-NotI and cloned into the above vector to remove the RAD51B sequence to obtain pFB-FLAG-XRCC3co-10xHis-RAD51Cco. Baculoviruses expressing RAD51B-RAD51C, RAD51D-XRCC2 and XRCC3-RAD51C were prepared separately and Sf9 cells were co-infected with optimized ratios for these viruses to express the BCDX2 complex as a heterotetramer and the CX3 complex as a heterodimer. Both the complexes were purified in an identical manner using affinity chromatography. Cells were harvested 52 hours post infection, washed once with cold PBS, and the pellets were frozen in liquid nitrogen and stored at -80 °C until further use.

The subsequent steps were carried out on ice or at 4 °C. The cell pellet was resuspended in lysis buffer (50 mM Tris-HCl pH 7.5, 2 mM β-ME, 1:400 [v/v] protease inhibitor cocktail [Sigma], 1 mM PMSF, 30 μg/ml leupeptin (Merck), 20 mM imidazole) for 20 minutes. Then, 50% glycerol was added to a final concentration of ~16%, followed by 5 M NaCl to a final concentration of 305 mM. The suspension was incubated for additional 30 minutes with gentle agitation. The total cell extract was centrifuged at 48,000 g for 30 minutes to obtain soluble extract. The extract was then bound to Ni-NTA resin (Qiagen) for 1 hour batch wise followed by extensive washing with Ni-NTA wash buffer (50 mM Tris-HCl pH 7.5, 2 mM β-ME, 300 mM NaCl, 10% glycerol, 1 mM PMSF, 10 μg/ml leupeptin, 20 mM imidazole) both batch wise and on a disposable column. The protein complexes were eluted by Ni-NTA elution buffer (Ni-NTA wash buffer containing 300 mM imidazole). The eluates were diluted 1:6 with a dilution buffer (Ni-NTA elution buffer without imidazole and 0.5 mM β-ME) and bound to FLAG resin (Sigma) pre-equilibrated with dilution buffer in flow with a total contact time of ~90 minutes. Protein bound FLAG-resin was washed 3-times with FLAG wash buffer (50 mM Tris-HCl pH 7.5, 0.5 mM β-
**Drosophila Topoisomerase I.** To prepare N-terminally truncated *Drosophila* topoisomerase I (catalytic subunit) with 6xHis tag on its C-terminus, the ND423 plasmid (a kind gift from James T. Kadonaga, University of California, San Diego, USA), was transformed in BL21 (DE3) pLysS cells and protein was purified by nickel affinity chromatography. The cell pellet from 1 liter culture was resuspended and sonicated in lysis buffer (50 mM Tris-HCl pH 7.5, 1 mM PMSF, 500 mM NaCl, 10% glycerol, 2 mM β-ME, 10 μg/ml leupeptin, 20 mM imidazole) and supplemented with 1:400 protease inhibitor cocktail (Sigma). Soluble extract was obtained by centrifugation at 48,000 g for 30 minutes and was incubated with pre-equilibrated Ni-NTA resin for 2 hours at 4 °C. Next, resin was washed 4 times with Ni-NTA wash buffer (50 mM Tris-HCl pH 7.5, 1 mM PMSF, 500 mM NaCl, 10% glycerol, 2 mM β-ME, 20 mM imidazole). Before elution the resin was washed once with the same Ni-NTA wash buffer as above but with 100 mM NaCl. Protein was eluted with elution buffer (50 mM Tris-HCl pH 7.5, 1 mM PMSF, 100 mM NaCl, 10% glycerol, 0.5 mM β-ME, 300 mM imidazole, 10 μg/ml leupeptin). Peak fractions were pooled and diluted 1:5 in dilution buffer (50 mM Tris-HCl pH 7.5, 0.5 mM β-ME, 100 mM NaCl, 10 % glycerol, 1 mM PMSF, 10 μg/ml leupeptin) and the sample was loaded onto pre-equilibrated HiTrap S and HiTrap Heparin columns connected in tandem (GE Healthcare), and washed with 20 ml of dilution buffer. The same buffer with a salt gradient up to 1 M NaCl was used to elute the protein from the HiTrap Heparin column after the HiTrap S column was disconnected. Peak fractions were pooled and dialyzed in dilution buffer for 2 hours. Protein was aliquoted, snap-frozen and stored at -80 °C.

**Yeast Rad51 (yRad51).** The *Rad51* sequence from *S. cerevisiae* was codon-optimized for bacterial expression and was purchased from GenScript. It was cloned into pMALT-P (a kind gift from the Kowalczykowski laboratory, UC Davis, USA) with BamHI and PstI restriction sites. The construct codes for Rad51 protein with a N-terminal MBP-tag separated from Rad51 by a PreScission protease site. The plasmid was transformed into BL21 (DE3) pLysS cells, 0.2% glucose was used to supplement the culture medium. The cell pellet from 2 liters was resuspended and sonicated in Buffer A (50 mM Tris-HCl pH 7.5, 1 mM PMSF, 500 mM NaCl, 1 mM DTT, 10% glycerol), supplemented with 1:500 protease inhibitor cocktail (Sigma). Soluble extract was obtained by centrifugation, and was incubated with amylose resin (New England Biolabs) at 4 °C for 1 hour in Buffer A containing 0.01% NP40. Extensive washes were carried out with Buffer A with 1 M NaCl, and subsequently with Buffer A containing 300 mM NaCl without PMSF. MBP-Rad51 was eluted with 10 mM maltose in Buffer A and 300 mM NaCl. Peak fractions were pooled, incubated with PreScission Protease for 1 hour at 4 °C (1:7; w/w), and
further diluted with 20 mM Tris-HCl pH 7.5 to reduce NaCl concentration to 150 mM NaCl (final).

The sample was loaded onto pre-equilibrated HiTrap Q HP column (GE Healthcare). Buffer R (20 mM Tris-HCl pH 7.5, 1 mM EDTA, 0.5 mM DTT, 10% glycerol) with 150 mM NaCl was used to wash the column. The same buffer R with a salt gradient up to 700 mM NaCl was used to elute the protein. Next, peak fractions were pooled and dialyzed over-night in Buffer R with 100 mM NaCl and without EDTA.

Protein was aliquoted, snap-frozen and stored at -80 °C.

**Human RAD51.** The RAD51 sequence was cloned from pTXB3-RAD51 construct into pMALT-P vector BamHI and PstI restriction sites, yielding N-terminal MBP tag, PreScission protease site and RAD51. The mutants were created by site directed mutagenesis using QuickChange II site-directed mutagenesis kit following manufacturer’s protocol (Agilent). The oligonucleotides used for mutagenesis are listed in supplementary Table S1. The RAD51 protein variants were expressed in BL21 (DE3) pLysS cells. Each culture was supplemented with 0.2% glucose, induced with 0.5 mM IPTG and grown overnight at 18 °C. The cells were then pelleted at 2,500 g for 15 minutes at 4 °C, washed once with STE buffer (10 mM Tris-HCl pH 8, 500 mM NaCl, 1 mM EDTA), snap-frozen and kept in -80 °C until use. The pellets were then resuspended in lysis buffer (50 mM Tris-HCl pH 7.5, 1 mM PMSF, 1 mM DTT, 10% glycerol, 500 mM NaCl, 1:500 protease inhibitor cocktail [Sigma]), sonicated and lysate was clarified by centrifugation at 48,000 g for 30 minutes. Next, the lysate was incubated with amylose resin for 1 hour batch-wise at 4 °C, washed first with wash buffer I (50 mM Tris-HCl pH 7.5, 1 mM DTT, 10% glycerol, 1 M NaCl) and then with wash buffer II (50 mM Tris-HCl pH 7.5, 1 mM DTT, 10% glycerol, 300 mM NaCl) followed by elution with wash buffer II containing 10 mM maltose. To cleave off the MBP tag, PreScission Protease was added to the eluate and incubated overnight at 4 °C (1:5, w/w). Cleaved RAD51 eluate was diluted with 50 mM Tris-HCl pH 7.5 to lower the NaCl concentration to 150 mM. The eluate was then applied to a Hitrap Q column (GE Healthcare). The column was washed sequentially with wash buffer III (20 mM Tris-HCl pH 7.5, 1 mM EDTA, 0.5 mM DTT, 10% glycerol, 150 mM NaCl) and eluted with wash buffer III with 300 mM NaCl. The fractions containing RAD51 were pooled and dialyzed in 20 mM Tris-HCl pH 7.5, 1 mM DTT, 20% glycerol and 100 mM NaCl overnight. The dialyzed protein was aliquoted, snap-frozen and stored at -80 °C. Wild type RAD51 was prepared from 4 liters of culture and all other variants were prepared from 1 liter cultures following the same purification procedure.

**Human RPA.** Recombinant human RPA was expressed from p11d–tRPA construct (a kind gift from M. Wold, University of Iowa) in BL21 (DE3) pLysS cells. Bacterial culture was grown at 37 °C (200 RPM) until O.D. 600 = 0.6, induced with 0.4 mM IPTG, and shaken at 18 °C (200 RPM) overnight. Bacterial pellet was obtained by centrifugation, washed once with SD Buffer (10 mM Tris pH 8.0, 150 mM NaCl, 1 mM EDTA), snap-frozen and stored at -80 °C. Cell lysis, followed by purification using ÄKTA pure (GE Healthcare) using HiTrap Blue HP, HiTrap desalting and HiTrap Q chromatography columns (all GE Healthcare).
Human mitochondrial single-stranded DNA binding protein (SSB). Recombinant mitochondrial SSB was expressed and purified from *E. coli* BL21 cells\(^8^4\).

**Human DNA2 and WRN.** Recombinant DNA2 was expressed in *Sf9* insect cells and purified by affinity chromatography by utilizing the N-terminal 6xHis and the C-terminal FLAG affinity tags\(^8^3\). WRN was purified by exploiting the maltose-binding protein (MBP) tag at the N-terminus and 10xHis tag at the C-terminus. The MBP tag was cleaved off with PreScission Protease during purification. Detailed procedures were described\(^8^3, 8^4\).

**Human MRN, MR, MRE11, phosphorylated CtIP and EXO1.** Recombinant MRN (MRE11-RAD50-NBS1) and MR (MRE11-RAD50) were purified as a complex from *Sf9* insect cells. Individual baculoviruses coding for MRE11-6xHis, RAD50co-FLAG and NBS1co were prepared by standard laboratory procedures according to manufacturer’s instructions (Bac-to-Bac, Life Technologies). Detailed procedures describing purification strategies have been described\(^8^3\). Briefly, the purification involved affinity chromatography using NiNTA agarose (Qiagen) and anti-FLAG affinity resin (Sigma). Recombinant MRE11 alone was prepared using pFB-MBP-MRE11-His vector, using amylose resin, PreScission Protease to remove the MBP tag, followed by affinity chromatography with NiNTA agarose\(^8^5\). Phosphorylated CtIP was purified from *Sf9* cells using amylose and NiNTA resins\(^3\). The recombinant human EXO1 gene was expressed from pFB-EXO1-FLAG in *Sf9* insect cells by using the Bac-to-Bac expression system (Invitrogen), according to manufacturer’s recommendations. Protein purification was performed with anti-FLAG (Sigma) affinity chromatography followed by HiTrap SP HP ion exchange chromatography (GE Healthcare) utilizing ÅKTA pure (GE Healthcare)\(^8^6\).

**Yeast MRX, Mre11, phosphorylated Sac2 and Exo1.** The proteins were prepared from *Sf9* insect cells and purified using affinity and ion exchange chromatography\(^8^7, 8^8, 8^9\). Exo1 was prepared according to an established procedure\(^6^7\).

**Preparation of oligonucleotide-based DNA substrates.**

Oligonucleotides were either 5'-end-labeled with [\(\gamma\)-\(^3^2\)P]-ATP (Perkin Elmer) and T4 polynucleotide kinase (NEB), or 3'-end-labeled with [\(\alpha\)-\(^3^2\)P]-dCTP (Perkin Elmer) and terminal transferase (NEB) enzymes, respectively. The labeled DNA was then purified on a Micro Bio-Spin P-30 Tris chromatography columns (Bio-Rad)\(^9^0\). Sequences for all oligonucleotides used to obtain the DNA substrates are listed in supplementary Table S1.

Branch migration substrate was prepared as described previously\(^9^1\). Briefly, 5'- or 3'-end-labeled 2 \(\mu\)M XO1 was mixed with 2.4 \(\mu\)M XO2 (1:1.2 ratio) in annealing buffer (10 mM Tris-HCl pH 8, 50 mM NaCl, 10 mM MgCl\(_2\)). In parallel, 2 \(\mu\)M each (1:1 ratio) of XO1c.MM2 and XO2c.MM oligonucleotides were similarly combined. The respective mixes were heated for 3 minutes at 95 °C and slowly cooled down to room temperature overnight. The two respective samples were then combined and annealed.
together (37 °C for 30 minutes), followed by gradual cooling down to room temperature (2 hours). Substrate was then stored at -20 °C until further use.

Fork reversal substrates were prepared as described earlier\textsuperscript{42}. Briefly, to create a fork with a leading strand gap, 3’- or 5’-labeled nascent #DC-6 (100 nM final) was annealed with unlabeled parental #DC-2 (120 nM final) in annealing buffer (as above) by heating (3 minutes at 95 °C) and gradually cooled down to room temperature overnight. Similarly, the complementary half comprising of unlabeled parental #DC-1 (180 nM final) and unlabeled nascent #DC-4 (180 nM final) were separately annealed. These two corresponding halves (#DC-6 + #DC-2 and #DC-1 + #DC-4) were then combined and annealed at 37 °C for 45 minutes and then cooled down to room temperature during 2 hours, and stored at -20 °C until further use. To create a fork with a lagging strand gap, 3’- or 5’-labeled nascent #DC-3 (100 nM final) was annealed with unlabeled parental #DC-1 (120 nM final) and the corresponding half containing unlabeled parental #DC-2 (150 nM final) was annealed with unlabeled nascent #DC-5 (150 nM final) oligos. These two halves (#DC-3 + #DC-1 and #DC-2 + #DC-5) were then combined and annealed as above.

To prepare the 5’-labeled 70 bp-long dsDNA substrate, which was quadruple blocked with streptavidin, the oligonucleotides PC210 and PC211 were used\textsuperscript{89}. To prepare 5’-labeled 50 bp-long dsDNA, oligonucleotides X12-3 and X12-4C were annealed. To prepare 5’-overhang substrate, X12-3 and X12-4SC were annealed, creating a structure with 19 nt-long 5’-overhang and 31 base pairs of dsDNA. To prepare 3’-overhang substrate, X12-3SC + X12 -4NC were annealed, creating a structure with 19 nt-long 3’-overhang and 31 base pairs of dsDNA, as described\textsuperscript{92}.

**Topoisomerase-coupled annealing assays**

The bubbled DNA annealing assay was performed as described\textsuperscript{43} with the following modifications. pBluescript II KS (+) plasmid (a kind gift from Marcus Thelen, IRB, Bellinzona, Switzerland) was used as a substrate. 100 ng supercoiled DNA was mixed with 1 µg RPA in TE pH 8.0 in 10 µl volume, and incubated for 45 minutes at 37 °C. Next, 16.5 nM (final) of catalytic domain of *Drosophila* topoisomerase I was added to the reaction mixture and incubated for additional 10 minutes at 37 °C. Next, annealing buffer (20 mM Tris-HCl pH 7.5, 5 mM MgCl\textsubscript{2}, 1 mM DTT, 0.2 mg/ml BSA, 100 mM NaCl), 2.5 mM ATP (final) and corresponding amounts of proteins (as indicated in figures) or protein storage buffer were added. The final volume was adjusted to 20 µl with water and the reactions were incubated for 30 minutes at 37 °C. The reactions were terminated by adding 2.5 µl of 5 M NaCl at room temperature for 2 minutes, followed by 6.5 µl of 2% stop buffer (100 mM Tris-HCl pH 7.5, 150 mM EDTA, 2% SDS [w/v], 30% glycerol, 0.1% bromophenol blue) and 1 µl Proteinase K (20 mg/ml, Roche), and incubated 10 minutes at 37 °C. The mixture was resolved by 1% agarose gel electrophoresis in 1x TAE buffer, and DNA was visualized by post-staining with GelRed (Biotium) according to manufacturer’s
instructions. The gels were then imaged (InGenius3, GeneSys) and quantitated as the fraction of near
or fully relaxed DNA using Image J. Graphs were generated by GraphPad Prism software.

**Fork reversal and branch migration assays**

The assays were carried out in a reaction buffer containing 20 mM Tris-HCl pH 7.5, 5 mM MgCl₂, 1
mM DTT, 0.1 mg/ml BSA, 10% glycerol, 2.5 mM ATP (unless indicated otherwise), and 1 nM (fork
reversal) or 0.5 nM (branch migration) DNA substrate, with 150 mM NaCl (unless indicated otherwise).
Master-mixes were prepared on ice and where indicated, RPA (3 nM) was added to the master-mix for
15 minutes on ice. 13 µl reaction mixture was then dispersed to individual tubes and supplemented with
other recombinant proteins (as indicated) and final volume was adjusted to 15 µl with protein storage
buffer. The reactions were continued for additional 30 minutes at 37 °C and terminated by the addition
of 5 µl stop buffer (100 mM Tris-HCl pH 7.5, 150 mM EDTA, 0.2% SDS [w/v], 30% glycerol, 0.1%
bromophenol blue) and 1 µl Proteinase K (20 mg/ml, Roche) and incubated for 10 minutes at 37 °C.
Samples were loaded onto 8% polyacrylamide (19:1 acrylamide:bisacrylamide) gels in 1x Tris-borate-
EDTA (TBE) (BIO-RAD Mini-PROTEAN system, 1 mm thick) and separated for 60 minutes at 80 V
at room temperature. The gels were dried using a BIO-RAD gel drier on 17 CHR paper (Whatman),
and were exposed to storage phosphor screens and scanned using Typhoon FLA 9500 (GE Healthcare)
phosphor imager. The gels were quantitated with ImageJ. Graphs were generated by GraphPad Prism
software.

**Single-stranded DNA annealing assay**

DNA annealing reactions were carried out at 37 °C for the times indicated using complementary oligo-
nucleotides X12-3 and X12-4C (please see supplementary Table S1 for sequences), 1 nM each. The
X12-3 oligonucleotide was labeled at the 5'-end. Control reactions were supplemented with protein
storage buffer. Reaction master mixes were prepared separately with the respective ssDNA in a buffer
containing 25 mM Tris-acetate pH 7.5, 5 mM magnesium acetate, 1 mM DTT, 1 mM ATP and supple-
mented with 4 nM RPA or SSB where indicated. The two respective master mixes were then incubated
for 5 minutes at 37 °C to allow RPA or SSB binding. The two respective mixes containing complemen-
tary ssDNA were then combined. Motor proteins and cofactors (when indicated) were added immedi-
ately and reaction volume was adjusted with water. The reactions were then incubated at 37 °C; 15 µl
reaction mixture was withdrawn at the indicated time points into tubes containing 5 µl stop buffer (100
mM Tris-HCl pH 7.5, 150 mM EDTA, 0.2% SDS [w/v], 30% glycerol, 0.1% bromophenol blue) and 1
µl Proteinase K (20 mg/ml, Roche). Tubes were kept on ice until the collection of the last time point,
and finally transferred to 37 °C for 10 minutes to achieve deproteination. Samples were then loaded
onto 8% polyacrylamide (19:1 acrylamide:bisacrylamide) gels in 1x TBE (BIO-RAD Mini-PROTEAN system, 1 mm thick), and processed as described above.

**ATPase assays**

ATPase assays were performed in a buffer containing 20 mM Tris-HCl pH 7.5, 5 mM MgCl₂, 1 mM DTT, 0.1 mg/ml BSA, 10% glycerol, 100 mM NaCl (unless otherwise indicated in the figures), 100 nM ATP, 1 nM of [γ-³²P] ATP (Perkin Elmer). 2961 bp long supercoiled pBluescript II KS (+) (7 nM, in molecules) or 5 nM unlabeled fork (#DC-1 + #DC-2) or Holliday junction structures ([XO1+XO2]+[XO1c.MM2 + XO2c.MM]) were used as a substrate. Recombinant proteins were added on ice and the samples were incubated at 37 °C for 60 minutes. Reactions were stopped with 2 µl of 0.5 M EDTA and separated using TLC plates (Merk) and 0.3 M LiCl and 0.3 M formic acid as a mobile phase. Dried plates were exposed to storage phosphor screens (GE Healthcare) and scanned by a Typhoon FLA 9500 phosphorimager (GE Healthcare). Signals were quantified using ImageJ software. Spontaneous ATP hydrolysis signal (Pi) from no protein lanes were removed as a background and the fraction of ATP hydrolysis was obtained as a normalized value. Graphs were generated by GraphPad Prism software.

**Electrophoretic mobility shift assays**

The electrophoretic mobility shift assay (EMSA) to characterize the binding of BCDX2 complex to 70-mer ssDNA (PC210) or dsDNA (PC210 annealed with PC211), 1 nM final; was carried out in 15 µl volume in a binding buffer containing 20 mM Tris-acetate pH 7.5, 1 mM DTT, 1 mM magnesium acetate, 0.1 mg/mL BSA (NEB) and 1 mM ATP. PC210 was labeled at 5' end. Reactions were assembled on ice and supplemented with increasing concentrations of BCDX2, and incubated for 30 minutes at 37 °C. The reactions were mixed with 5 µl loading buffer (50% glycerol, 0.1 % bromophenol blue) and loaded on 8% polyacrylamide (19:1 acrylamide:bisacrylamide) gels in 1x TBE (BIO-RAD Mini-PROTEAN system, 1 mm thick), and separated for 150 minutes at 80 V at 4 °C. The gels were dried using a BIO-RAD gel drier on 17 CHR paper (Whatman), exposed to storage phosphor screens and scanned using Typhoon FLA 9500 (GE Healthcare) phosphor imager.

**Protein-protein interaction assays**

To study the interaction between SMARCAL1 wild type and corresponding SMARCAL1 F→A variants with RAD51 or between ZRANB3 WT, ZRANB3 F→A variants with RAD51, bacterial soluble extract containing MBP-RAD51 was incubated with amylose resin (50 µl, NEB). The resin was washed with wash buffer I (25 mM Tris-HCl pH 7.5, 0.5 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.2 µg/µl
BSA) and incubated with recombinant purified FLAG-SMARCAL1, FLAG-ZRANB3 or the FLAG-BCDX2 complex (all 1 µg) in 150 µl IP buffer (25 mM Tris-HCl pH 7.5, 0.5 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.2 µg/µl BSA, 10% Glycerol) for 1 hour at 4 °C. The resin with bound proteins was washed 5 times with 1 ml wash buffer III (25 mM Tris-HCl pH 7.5, 1 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.05% Triton X-100), and eluted with wash buffer III (70 µl) containing 30 mM maltose and Avidin (0.11 µg/µl) as a stabilizer. Samples were analyzed by Western blotting using anti-MBP primary antibody (MBL, M091-3, 1:1000) against MBP-RAD51 and anti-FLAG primary antibody (Sigma, F3165, 1:1000) against SMARCAL1, ZRANB3 or against RAD51B of the BCDX2 complex, respectively, by standard procedures.

To study interaction between RAD51 and ZRANB3 variants, FLAG-tagged ZRANB3 variants were expressed in Sf9 cells, cells were lysed and soluble extract containing the FLAG-ZRANB3 proteins was bound to M2 anti-FLAG affinity resin (50 µl, Sigma). The resin was washed 3-times with 1 ml wash buffer I (25 mM Tris-HCl pH 7.5, 0.5 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.2 µg/µl BSA) and incubated for 1 hour at 4 °C with recombinant purified RAD51 (1 µg) in 150 µl IP buffer (25 mM Tris-HCl pH 7.5, 0.5 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.2 µg/µl BSA). The resin with bound proteins was washed 5-times with 1 ml wash buffer II (25 mM Tris-HCl pH 7.5, 0.5 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.1% NP40, 0.2 µg/µl BSA), and proteins were eluted with wash buffer II (70 µl) containing 150 ng/µl 3xFLAG peptide (GLP BIO) and Avidin (0.11 µg/µl [Sigma]) as a stabilizer. Samples were analyzed by Western blotting using anti-RAD51 primary antibody (Abcam-133534, 1:1000) or by Ponceau staining to show ZRANB3, using standard laboratory procedures.

To study the interaction between ZRANB3 and the BCDX2 complex, or between SMARCAL1 variants and the BCDX2 complex, 1 µg (1 µl) anti-His primary antibody (MBL-D2913) was mixed with 15 µl Dynabeads Protein G (Invitrogen) slurry in a solution containing 150 µl 1X PBS containing 0.05% Tween 20 (PBS-T). The mixture was incubated for 45 minutes at room temperature with gentle mixing. The cocktail was washed 3 times with 150 µl PBS-T and was further resuspended in 60 µl IP buffer (25 mM Tris-HCl pH 7.5, 0.5 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.2 µg/µl BSA), which was then supplemented with 1 µg recombinant purified BCDX2 complex and incubated for 1 hour at 4 °C with gentle mixing. The beads were washed 3-times with 150 µl wash buffer (50 mM Tris-HCl pH 7.5, 1 mM DTT, 3 mM EDTA, 100 mM NaCl, 0.05% Triton X-100) and again resuspended in IP buffer. Purified recombinant SMARCAL1 or ZRANB3 (1 µg) was added and incubated for 1 hour at 4 °C with gentle mixing, and washed 4-times with wash buffer (50 mM Tris-HCl pH 7.5, 100 mM NaCl, 0.05% Triton X-100). Proteins were eluted by heating the beads for 3 minutes at 95 °C in 60 µl SDS buffer (50 mM Tris-HCl pH 6.8, 1.6 % SDS, 10 % Glycerol, 10% DTT, 0.01 % Bromophenol Blue) and transferred to a new tube containing Avidin as a stabilizer (0.11 µg/µl). Samples were resolved by polyacrylamide gel electrophoresis and protein bands were visualized either by silver staining or by
Western blotting using anti-FLAG to detect SMARCAL1 and ZRANB3, and by anti-His primary antibodies to detect RAD51C of the BCDX2 complex by standard procedures.

**Antibodies.** The antibodies used for Western blotting, immunoprecipitation and DNA fiber assay were used as follows: Mouse anti-RAD51B (Santa Cruz sc-377192; 1:1,000 dilution for WB), Mouse anti-XRCC2 (Santa Cruz sc-365854; 1:1,000 dilution for WB), Mouse anti-XRCC3 (Santa Cruz sc-271714; 1:1,000 dilution for WB), Rabbit anti-RAD51C (Abcam ab95069; 1:1,000 dilution for WB), Rabbit anti-RAD51D (Abcam ab202063; 1:1,000 dilution for WB), Rat anti-BrdU (Abcam ab6326; 1:100 dilution for DNA fiber assay), Mouse anti-BrdU (Becton Dickinson BD347580; 1:100 dilution for DNA fiber assay), Mouse anti-SMARCAL1 (Santa Cruz sc-376377; 1:1,000 dilution for WB), Mouse anti-BRCA1 (Santa Cruz sc-6954; 1:100 dilution for WB), Rat anti-TUBULIN (Abcam ab-6160; 1:50,000 dilution for WB), Goat anti-mouse Alexa Fluor 488 (Thermo Fisher A-11029; 1:300 dilution for DNA fiber assay), Goat anti-rabbit Alexa Fluor 594 (Thermo Fisher A-11008; 1:300 dilution for DNA fiber assay), Mouse anti-his (MBL D291-3; 1 µg for pulldown assay), Mouse anti-FLAG (Sigma F3165; 1:2,000 dilution for WB), Mouse anti-MBP (MBL M091-3; 1:1,000 dilution for WB), Rabbit anti-RAD51 (Abcam ab133534; 1:1,000 dilution for WB).

**Nuclease protection assays**

Endonuclease assays with MRN-pCtIP were performed in a 15 µl volume in a nuclease buffer (25 mM Tris-HCl pH 7.5, 1 mM manganese acetate, 5 mM magnesium acetate, 1 mM DTT, 1 mM ATP, 0.25 mg/mL BSA, 1 mM phosphoenolpyruvate, 80 U/ml pyruvate kinase [Sigma]) with 1 nM biotinylated DNA substrate (in molecules, 70 bp-long, 5'-end-labeled PC210 and PC211) and incubated with streptavidin (15 nM, Sigma) for 5 minutes at room temperature to block the DNA ends. Before the addition of nucleases, increasing amounts of yeast and human RAD51 were added to the reactions and preincubated for 10 minutes at 37 °C. Recombinant MRN and pCtIP were subsequently added, and the reaction was incubated for 2 hours at 37 °C. Endonuclease assays with yeast MRX-pSae2 were performed similarly as MRN-pCtIP, however yeast and human RAD51 assembly (10 minutes incubation) and subsequent reactions were performed at 30 °C for 30 minutes.

Exonuclease assays with human MR were performed in the nuclease buffer as above, but with 5 nM DNA substrate (in molecules, except where mentioned otherwise), which was 50 bp-long dsDNA (X12-3 and X12-4C) or with a 3' (X12-3SC + X12-4NC) or 5' (X12-3 and X12-4SC) overhangs. Yeast and human RAD51 were then preassembled for 10 minutes on DNA at 37 °C and the assays were subsequently incubated for 120 minutes at 37 °C after adding the nucleases. RAD51 assembly and subsequent nuclease assays with yeast MR were performed at 30 °C.
For nuclease assays with MRE11, the reaction buffer contained 25 mM Tris-HCl pH 7.5, 3 mM manganese acetate, 5 mM magnesium acetate, 1 mM DTT, 1 mM ATP, 0.25 mg/ml BSA, 1 mM phosphoenolpyruvate and 80 U/ml pyruvate kinase (Sigma). 1 nM 5'-end-labeled 50 bp-long dsDNA (in molecules, X12-3 and X12-4C) was used as a substrate. Human or yeast RAD51/Rad51 assembly (10 minutes) and further reactions were carried out at 37 °C for 120 minutes.

Nuclease assays with human EXO1 were performed with 1 nM 5'-end-labeled 50 bp-long dsDNA substrate (in molecules, X12-3 and X12-4C) in a buffer with 25 mM Tris-HCl pH 7.5, 2 mM magnesium acetate, 1 mM DTT, 1 mM ATP, 0.1 mg/ml BSA, 1 mM phosphoenolpyruvate and 80 U/ml pyruvate kinase (Sigma). Human or yeast RAD51/Rad51 was assembled on DNA for 10 minutes at 37 °C and the reactions were further incubated at 37 °C for 30 minutes. However, for assays with yeast Exo1, the buffer contained 5 mM magnesium acetate and the reactions were performed at 30 °C.

Endonuclease assays with NspI (NEB) were performed with 1 nM (in molecules) 5'-end-labeled 50 bp-long dsDNA (X12-3 and X12-4C) in CutSmart buffer (NEB) and supplemented with 2.5 mM ATP to aid RAD51 binding to DNA. Substrates were preincubated with increasing amount of human or yeast Rad51 for 10 minutes at 37 °C followed by the addition of 1 unit of NspI (per 15 µl) and further incubation for 30 minutes at 37 °C.

EXO-III mediated Holliday junction degradation assay (0.5 nM 5'-end-labeled substrate, in molecules) was performed in branch migration assay buffer containing 100 mM NaCl. However, in the assays comparing the DNA protection by human and yeast RAD51/Rad51, 1 nM 5'-end-labeled 50 bp-long dsDNA substrate was used in branch migration assay buffer without additional salt. RAD51 assembly (10 minutes) and subsequent reactions (30 minutes) were performed at 37 °C.

All reactions were stopped with 0.5 µl EDTA (0.5 M) and 1 µl proteinase K (20 mg/ml, Roche), and incubated at 50 °C for 30 minutes. An equal volume of formamide dye (95% [v/v] formamide, 20 mM EDTA, 0.1% bromophenol blue) was added and the samples were heated at 95 °C for 4 minutes and separated on 15% denaturing polyacrylamide gels (ratio acrylamide: bisacrylamide 19:1, Bio-Rad). The gels were fixed for 30 minutes at room temperature (40% methanol, 10% acetic acid, 5% glycerol) and dried on a 3 MM CHR paper (Whatman). The dried gels were exposed to storage phosphor screens (GE Healthcare) and scanned by a Typhoon phosphor imager (FLA 9500, GE Healthcare). The gels were quantitated with ImageJ. Graphs were generated by GraphPad Prism software.

For DNA2-catalyzed assays, randomly labeled 2.2-kbp-long substrate was prepared by amplifying the human NBS1 gene by PCR from pFB-MBP-NBS1-his plasmid using Phusion high-fidelity DNA polymerase (New England Biolabs) and the NBS1_F and NBS1_R primers (see Table S1). 66 nM [α-32P]dCTP was added to the PCR reaction together with the standard dNTPs concentration (200 µM each). The PCR reaction product was purified using the QIAquick PCR purification kit (Qiagen) and...
Chroma Spin TE-200 columns (Clontech). Purified DNA was quantitated by comparing the radioactive DNA fragment with known amounts of a cold PCR product on an agarose gel stained with GelRed (Biotium). Nuclease assays with PCR-based DNA substrates<sup>86</sup> were performed in a 15 µl volume in a reaction buffer containing 25 mM Tris-acetate pH 7.5, 2 mM magnesium acetate, 1 mM ATP, 1 mM DTT, 0.1 mg/ml BSA, 1 mM phosphoenolpyruvate (PEP), 80 U/ml pyruvate kinase (Sigma), 50 mM NaCl, RPA (176 nM) and 1 nM substrate (in molecules). Increasing amount of RAD51 (as indicated in the figure) was preincubated with the substrates for 10 minutes at 37°C in the reaction buffer, which was then supplemented with other recombinant proteins and the reaction was continued for 30 minutes at 37 °C. Reactions were stopped with 5 µl 2% stop solution (150 mM EDTA, 2% sodium dodecyl sulfate, 30% glycerol, 0.1% bromophenol blue) and 1 µl of proteinase K (20 mg/ml, Roche) and incubated at 37°C for 10 minutes. Samples were resolved by 1% agarose gel electrophoresis. Gels were dried on DE81 chromatography paper (Whatman), exposed to storage phosphor screens (GE Healthcare) and scanned by a Typhoon 9500 phosphor imager (GE Healthcare). The gels were quantitated with ImageJ. Graphs were generated by GraphPad Prism software.

**Cellular assays**

MCF10A cells were maintained in a 1:1 mixture of DMEM and Ham’s F12 medium (Thermo Fisher Scientific), supplemented with 5% horse serum (Thermo Fisher Scientific), 20 ng/ml human epidermal growth factor (Peprotech), 100 ng/ml cholera toxin, 10 µg/ml insulin and 0.5 µg/ml hydrocortisone (Sigma-Aldrich). The human embryonic kidney fibroblast cell line HEK293T was maintained in DMEM supplemented with 10% Fetalgro bovine growth serum. Gateway LR recombination (Thermo Fisher Scientific) was used to recombine pDONR223-SMARCAL1(F446A) with the lentiviral expression vector pHAGE-C-FLAG-HA-DEST<sup>93</sup>. Recombinant lentiviruses were generated by cotransfecting helper packaging vectors together with lentiviral vectors into HEK293T cells using the TransIT-293 transfection reagent (Mirus Bio). Virus-containing supernatants were collected 48 hours after transfection and utilized to infect MCF10A cells in the presence of 8 µg/ml polybrene. 48 hours after viral addition, MCF10A cells were selected using 1 µg/ml puromycin for 3 days. To perform RNAi treatments, MCF10A SMARCAL1 KO cells complemented with WT and F446A mutant SMARCAL1 were transfected with control or BRCA1 siRNA (GE Dharmacon) using lipofectamine RNAiMAX (Thermo Fisher Scientific) according to manufacturer’s instructions and subjected to DNA fiber assays 3 days after transfection. To analyze cell lysates by Western blotting, cells were collected by trypsinization and lysed in SB lysis buffer (62.5 mM Tris-HCl pH 6.8, 1.25% SDS, 12% glycerol, 0.71 M [5%] β-ME, 0.002% bromophenol blue). Whole cell extracts were sonicated and heated for 5 minutes at 95 °C. Following gel electrophoresis and transfer of cell extracts onto nitrocellulose, membranes were incubated for 1 hour or overnight in blocking buffer (5% milk in TBS + 0.1% Tween20). Membranes were
subsequently incubated with primary antibodies diluted in antibody blocking buffer for 2 hours at room temperature or overnight at 4 °C. Detection was achieved using appropriate horseradish peroxidase (HRP)-conjugated secondary antibodies. Anti-SMARCAL1 (1:1000, Santa Cruz Biotechnology), anti-BRCA1 (1:100, Santa Cruz Biotechnology) and anti-TUBULIN (1:50000, Abcam) antibodies were used in western blot experiments.

**Single-molecule analysis of DNA replication**

Exponentially growing MCF10A cells were pulse-labeled with 30 μM CldU (25 minutes), washed and exposed to 150 μM IdU (35 minutes). After exposure to the second nucleotide analog, the cells were washed again in warm 1x PBS and treated or not for 4 hours with hydroxyurea (HU, 2 mM, Sigma). Labeled cells were trypsinized and resuspended in ice-cold PBS at 4 × 10^5 cells/ml. Two microliters of this suspension were spotted onto a pre-cleaned glass slide and lysed with 10 μl of spreading buffer (0.5% SDS in 200 mM Tris-HCl pH 7.4, 50 mM EDTA). After 6 minutes, the slides were tilted at 15° relative to horizontal, allowing the DNA to spread. Slides were air-dried, fixed in methanol and acetic acid (3:1) for 2 minutes, rehydrated in PBS for 10 minutes and denatured with 2.5 M HCl for 50 minutes at room temperature. Slides were then rinsed in PBS and blocked in PBS + 0.1% Triton X-100 (PBS-T) + 5% BSA for 1 hour at room temperature. Rat anti-BrdU (1:100, Abcam) and mouse anti-BrdU (1:100, BD) were then applied to detect CldU and IdU, respectively. After 1 hour incubation, the slides were washed in PBS and stained with Alexa Fluor 488-labeled goat anti-mouse IgG1 antibody and Alexa Fluor 594-labeled goat anti-rat antibody (1:300 each, Thermo Fisher Scientific). Slides were mounted in Prolong Gold Antifade (Thermo Fisher Scientific) and held at -20 °C. Replication tracks were imaged on a Nikon Eclipse 90i microscope fitted with a PL Apo 40X/0.95 numerical aperture (NA) objective and measured using ImageJ software. In each experiment, 100 or more dual-labeled tracts were measured for fork degradation estimation.

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**DATA AVAILABILITY**

All primary data is available in this manuscript, supplementary information or source data.
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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

AUTHOR CONTRIBUTIONS

S.H. performed most of the experiments, analyzed data, prepared figures. A.S. did some protection assay with human RAD51. L.R. prepared RAD51. A.T. did in vivo studies with mutant SMARCAL1. G.R., I.C., A.A. generated reagents for the study. R.A. performed preliminary protection experiments with overhanded substrates. A.C. supervised A.T. for cellular studies using DNA fiber assay. P.C. conceived, supervised and wrote the manuscript with input from S.H.

FIGURE LEGENDS

Figure 1. SMARCAL1, ZRANB3 and HLTF possess distinct biochemical activities

(a) Recombinant SMARCAL1, ZRANB3, and HLTF were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.
(b) A schematic of replication fork reversal assay (leading and lagging strand gap structure is shown).
(c) Fork reversal assays with SMARCAL1, ZRANB3 and HLTF with RPA (3 nM). Top, quantifications (error bars show SEM of three replicates); bottom, representative experiments.
(d) Fork reversal assays with HLTF without or with RPA (3 nM). Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.
(e) A schematic of Holliday junction branch migration assay.
(f) Holliday junction branch migrations assay with SMARCAL1, ZRANB3 and HLTF. Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.
(g) A schematic of topoisomerase-coupled annealing assay.
(h) Comparison of SMARCAL1, ZRANB3 and HLTF in topoisomerase-coupled annealing assays.
ATP hydrolysis by HLTF is required, as no detectable annealing was observed without ATP.
**Figure 2. SMARCAL1 anneals RPA-coated ssDNA**

(a) A schematic of ssDNA annealing assays.

(b) Annealing of ssDNA by SMARCAL1, ZRANB3 and HLTF without or with RPA (4 nM). Representative experiments are shown.

(c) Quantification of experiment as in (b) at 30 minutes (error bars indicate SEM of three replicates).

(d) Annealing of ssDNA by SMARCAL1 without or with human mitochondrial SSB (4 nM). Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.

(e) Top, a schematic showing domain organization of SMARCAL1. RPA binding domain is located in the N-terminal part of SMARCAL1 (indicated in dark blue). SMARCAL1ΔN lacking RPA binding domain is shown below.

(f) Recombinant SMARCAL1-WT and SMARCAL1ΔN were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.

(g) A comparison of SMARCAL1-WT and SMARCAL1ΔN in ssDNA annealing without or with RPA (4 nM). Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.

**Figure 3. RAD51 and BCDX2 promote SMARCAL1 and ZRANB3 mediated branch migration and DNA annealing**

(a) Recombinant RAD51 and BCDX2 were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.

(b) Annealing helicase assay. RAD51 and BCDX2 separately stimulate SMARCAL1-mediated annealing of RPA-coated ssDNA. Top, quantifications (error bars indicate SEM of four replicates); bottom, representative experiment. Statistical significance; *(P < 0.05), ***(P < 0.001), two-tailed t-test.

(c) RAD51 and BCDX2 separately promote ZRANB3 mediated strand annealing. Top, quantifications (error bars indicate SEM of four replicates); bottom, representative experiment. Statistical significance bars; ns (P > 0.05, not significant); **(P < 0.01); two-tailed t-test.

(d) RAD51 and BCDX2 do not promote HLTF-mediated DNA annealing. Shown is a representative experiment.

**Figure 4. SMARCAL1 and ZRANB3 physically interact with RAD51 and BCDX2**

(a) Soluble extract from *E. coli* containing MBP-RAD51 (bait) was immobilized on amylose resin and incubated with purified recombinant proteins (RAD51 paralogs, ZRANB3, SMARCAL1, [prey]) as indicated. Western blot analyses were performed with anti-MBP and anti-FLAG antibodies.
(b) and (c) Anti-His antibody was coupled to Protein G agarose, bound to the BCDX2 complex (bait) and tested for interaction with ZRANB3 (prey) or SMARCAL1 (prey), respectively. Samples were subjected to either silver staining or Western blot analysis by anti-FLAG and anti-His antibodies.

(d) Multiple sequence alignment showing the presence of a consensus FxxA motif in SMARCAL1 along with previously characterized RAD51 interacting proteins or BRCA2 domains (highlighted in grey with bold red letters).

(e) Recombinant SMARCAL1-WT and SMARCAL1-F446A were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.

(f) SMARCAL1-F446A fails to interact with RAD51. Soluble extract from *E. coli* containing MBP-RAD51 (bait) was immobilized on amylose resin and incubated with purified recombinant SMARCAL1 variants (prey). Ponceau staining shows RAD51. Western blot analysis was performed with anti-FLAG antibody to detect SMARCAL1.

(g) SMARCAL1-F446A, as SMARCAL1-WT, interacts with the BCDX2 complex. Anti-His antibody was immobilized on protein G agarose, bound to BCDX2 complex (bait) and tested for interaction with SMARCAL1 variants (prey). Samples were subjected to silver staining.

(h) DNA fiber assay to monitor SMARCAL1-mediated nascent DNA degradation in BRCA1-deficient cells. Wild type or SMARCAL1-F446A proteins were expressed in SMARCAL1 KO MCF10A cells upon BRCA1 depletion, as indicated. SMARCAL1-deficiency renders BRCA1-depleted cells resistant to replication fork degradation upon hydroxyurea (HU) treatment, as a result of impaired fork reversal. Top: a schematic of the assay: CldU (25 minutes), IdU (35 minutes) pulse-labeling protocol to evaluate fork degradation upon HU treatment. Under wild type condition the ratio of IdU/CldU tract length will remain ~1, however if there is fork degradation this ratio will be < 1. Bottom: graphical representation of IdU/CIdU tract length ratio. The median value of 100 or more IdU and CldU tracts per experimental condition is indicated. Statistical analysis was conducted using Mann-Whitney test (****p < 0.0001). Data are representative of two independent experiments.

**Figure 5. RAD51 protects DNA from degradation by MRE11, EXO1 and DNA2 nucleases**

(a) Top, a schematic of the assay. Bottom, endonuclease activity of MRN-pCtIP on quadruple blocked 5’-end-labeled 70 bp-long dsDNA is efficiently inhibited by RAD51-WT and RAD51-K133R but not by RAD51-K133A, RAD51-Y232A or RAD51-T131P mutants. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis. In the schematic at the top, asterisk indicates the position of the radioactive label.

(b) Quantification of experiments such as in panel (a) (error bars indicate SEM of three replicates). The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 2 in panel a).
(c) Top, a schematic of the assay. Bottom, RAD51-WT and ATP-hydrolysis deficient K133R mutant efficiently protect DNA against the exonuclease activity of MR (MRE11-RAD50). RAD51 variants (RAD51-K133A, RAD51-Y232A or RAD51-T131P) with decreased DNA binding capacity failed to inhibit MR. 5'-end-labeled 50 bp-long dsDNA was used as a substrate. Asterisk indicates the position of the $^{32}$P label. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis.

(d) Quantification of experiments such as shown in panel (c), error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 2 in panel c).

(e) RAD51 efficiently protects against the exonuclease activity of MR on a 5'-end-labeled 50 bp-long dsDNA in presence of ATP. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis. Top, quantifications (error bars indicate SEM of three replicates); bottom, a representative experiment.

(f) ATP-bound RAD51 efficiently protects against EXO1 on a 3'-end-labeled 50 bp-long dsDNA. Calcium (1 mM) traps ATP-bound RAD51 on DNA, which greatly enhances RAD51-mediated protection. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis.

(g) Quantifications of experiments such as in panel (f), error bars indicate SEM of at least three replicates.

(h) Top, a schematic of the assay. Bottom, RAD51-WT and RAD51-K133R, but not RAD51-K133A variant efficiently protect against EXO1 degradation, using a 5'-end-labeled 50 bp-long dsDNA as a substrate. Asterisk indicates the position of the $^{32}$P label. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis.

(i) Quantification of experiments such as shown in panel (h); error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 2 in panel h).

(j) Top, a schematic of the assay. Bottom, DNA degradation by WRN, DNA2 and RPA is inhibited by RAD51. 2.2-kilobase pair (kbp)-long randomly labeled dsDNA was used as the substrate. Reaction products were separated by 1% agarose gel electrophoresis. Red asterisks indicate random DNA labeling.

(k) Quantification of experiments such as shown in panel (j); error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 4 in panel j).

Figure 6. Nucleolytic degradation is prevented by RAD51 upon binding to dsDNA

(a) Nuclease assays with MRE11-RAD50 (MR) and their inhibition by RAD51. The experiments were carried out with blunt-ended, 5'-overhanded or 3'-overhanded DNA. Asterisk indicates the position of
the labelling. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis. A representative experiment is shown.

(b) Quantifications of experiments such as shown in panel (a), error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation for each substrate without RAD51 (lanes 3, 7 and 11 in panel a).

(c) Nuclease assays with EXO1 and its inhibition by RAD51. Experiments were carried out with blunt-ended, 5'-overhanged or 3'-overhanged DNA. Asterisk indicates the position of the $^{32}$P label. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis. Shown is a representative experiment.

(d) Quantifications of experiments such as shown in panel (c), error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation for each substrate without RAD51 (lanes 2, 7 and 12 in panel c).

Figure 7. Model for replication fork reversal and protection.

Fork remodelers have unequal biochemical functions. SMARCAL1 anneals RPA-coated ssDNA and may promote initial steps in fork reversal. ZRANB3 and HLTF are more proficient in branch migration. RAD51 and BCDX2 interact with SMARCAL1 and ZRANB3 and stimulate their activities. Reversed replication forks are prone to pathological degradation, in certain genetic backgrounds, unless protected by RAD51. We show that unexpectedly the dsDNA-binding capacity of RAD51 promotes DNA protection against nucleases.
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Figure 1. SMARCAL1, ZRANB3 and HLTF possess distinct biochemical activities

(a) Recombinant SMARCAL1, ZRANB3, and HLTF were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.

(b) A schematic of replication fork reversal assay (leading and lagging strand gap structure is shown).

(c) Fork reversal assays with SMARCAL1, ZRANB3 and HLTF. Top, quantifications (error bars show SEM of three replicates); bottom, representative experiments.

(d) Fork reversal assays with HLTF without or with RPA. Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.

(e) A schematic of Holliday junction branch migration assay.

(f) Holliday junction branch migrations assay with SMARCAL1, ZRANB3 and HLTF. Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.

(g) A schematic of topoisomerase-coupled annealing assay.

(h) Comparison of SMARCAL1, ZRANB3 and HLTF in topoisomerase-coupled annealing assays. ATP hydrolysis by HLTF is required, as no detectable annealing was observed without ATP.
Figure 2. SMARCAL1 anneals RPA-coated ssDNA

(a) A schematic of ssDNA annealing assays.

(b) Annealing of ssDNA by SMARCAL1, ZRANB3 and HLTF without or with RPA. Representative experiments are shown.

(c) Quantification of (b) at 30 minutes (error bars indicate SEM of three replicates).

(d) Annealing of ssDNA by SMARCAL1 without or with human mitochondrial SSB. Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.

(e) Top, a schematic showing domain organization of SMARCAL1. RPA binding domain is located in the N-terminal part of SMARCAL1 (indicated in dark blue). SMARCAL1ΔN lacking RPA binding domain is shown below.

(f) Recombinant SMARCAL1-WT and SMARCAL1ΔN were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.

(g) A comparison of SMARCAL1-WT and SMARCAL1ΔN in ssDNA annealing without or with RPA. Top, quantifications (error bars indicate SEM of three replicates); bottom, representative experiments.
Figure 3. RAD51 and BCDX2 promote SMARCAL1 and ZRANB3 mediated bubble DNA annealing
(a) Recombinant RAD51 and BCDX2 were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.
(b) Annealing helicase assay. RAD51 and BCDX2 separately stimulate SMARCAL1-mediated annealing of RPA-coated ssDNA. Top, quantifications (error bars indicate SEM of four replicates); bottom, representative experiment. Statistical significance; *(P < 0.05), ***(P < 0.001), two-tailed t-test.
(c) RAD51 and BCDX2 separately promote ZRANB3 mediated strand annealing. Top, quantifications (error bars indicate SEM of four replicates); bottom, representative experiment. Statistical significance bars; ns (P > 0.05, not significant); **(P < 0.01); two-tailed t-test.
(d) RAD51 and BCDX2 do not promote HLTF-mediated DNA annealing. Shown is a representative experiment.
Figure 4. SMARCAL1 and ZRANB3 physically interact with RAD51 and BCDX2

(a) Soluble extract from E. coli containing MBP-RAD51 (bait) was immobilized on amylose resin and incubated with purified recombinant proteins (RAD51 paralogs, ZRANB3, SMARCAL1, [prey]) as indicated. Western blot analyses were performed with anti-MBP and anti-FLAG antibodies.

(b) and (c) Anti-His antibody was coupled to Protein G agarose, bound to the BCDX2 complex (bait) and tested for interaction with ZRANB3 (prey) or SMARCAL1 (prey), respectively. Samples were subjected to either silver staining or Western blot analysis by anti-FLAG and anti-His antibodies.

(d) Multiple sequence alignment showing the presence of a consensus FxxA motif in SMARCAL1 along with previously characterized RAD51 interacting proteins or BRCA2 domains (highlighted in grey with bold red letters).

(e) Recombinant SMARCAL1-WT and SMARCAL1-F446A were analyzed by polyacrylamide gel electrophoresis and stained with Coomassie Brilliant Blue.

(f) SMARCAL1-F446A fails to interact with RAD51. Soluble extract from E. coli containing MBP-RAD51 (bait) was immobilized on amylose resin and incubated with purified recombinant SMARCAL1 variants (prey). Ponceau staining shows RAD51. Western blot analysis was performed with anti-FLAG antibody to detect SMARCAL1.

(g) SMARCAL1-F446A, as SMARCAL1-WT, interacts with the BCDX2 complex. Anti-His antibody was immobilized on protein G agarose, bound to BCDX2 complex (bait) and tested for interaction with SMARCAL1 variants (prey). Samples were subjected to silver staining.

(h) DNA fiber assay to monitor SMARCAL1-mediated nascent DNA degradation in BRCA1-deficient cells. Wild type or SMARCAL1-F446A proteins were expressed in SMARCAL1 KO MCF10A cells upon BRCA1 depletion, as indicated. SMARCAL1-deficiency renders BRCA1-depleted cells resistant to replication fork degradation upon hydroxyurea (HU) treatment, as a result of impaired fork reversal. Top: a schematic of the assay: CldU (25 minutes), IdU (35 minutes) pulse-labeling protocol to evaluate fork degradation upon HU treatment. Under wild type condition the ratio of IdU/CldU tract length will remain ~1, however if there is fork degradation this ratio will be < 1. Bottom: graphical representation of IdU/CldU tract length ratio. The median value of 100 or more IdU and CldU tracts per experimental condition is indicated. Statistical analysis was conducted using Mann-Whitney test (**p < 0.0001). Data are representative of two independent experiments.
Figure 5. RAD51 protects DNA from degradation by MRE11, EXO1 and DNA2 nucleases

(a) Top, a schematic of the assay. Bottom, endonuclease activity of MRN-pCtIP on quadruple blocked 5’-end-labeled 70 bp-long dsDNA is efficiently inhibited by RAD51-WT and RAD51-K133R but not by RAD51-K133A, RAD51-Y232A or RAD51-T131P mutants. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis. In the schematic at the top, an asterisk indicates the position of the radioactive label.

(b) Quantification of experiments such as in panel (a) (error bars indicate SEM of three replicates). The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 2 in panel a).

Continued on next page.
(c) Top, a schematic of the assay. Bottom, RAD51-WT and ATP-hydrolysis deficient K133R mutant efficiently protect DNA against the exonuclease activity of MR (MRE11-RAD50). RAD51 variants (RAD51-K133A, RAD51-Y232A or RAD51-T131P) with decreased DNA binding capacity failed to inhibit MR. 5’-end-labeled 50 bp-long dsDNA was used as a substrate. Asterisk indicates the position of the $^{32}$P label. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis.

(d) Quantification of experiments such as shown in panel (c), error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 2 in panel c).

(e) RAD51 efficiently protects against the exonuclease activity of MR on a 5’-end-labeled 50 bp-long dsDNA in presence of ATP. Top, quantifications (error bars indicate SEM of three replicates); bottom, a representative experiment.

(f) ATP-bound RAD51 efficiently protects against EXO1 on a 5’-end-labeled 50 bp-long dsDNA. Calcium (1 mM) traps ATP-bound RAD51 on DNA, which greatly enhances RAD51-mediated protection.

(g) Quantifications of experiments such as in panel (f), error bars indicate SEM of at least three replicates.

(h) Top, a schematic of the assay. Bottom, RAD51-WT and RAD51-K133R, but not RAD51-K133A variants efficiently protect against EXO1 degradation, using a 5’-end-labeled 50 bp-long dsDNA as a substrate. Asterisk indicates the position of the $^{32}$P label. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis.

(i) Quantification of experiments such as shown in panel (h); error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 2 in panel h).

(j) Top, a schematic of the assay. Bottom, DNA degradation by WRN, DNA2 and RPA is inhibited by RAD51. 2.2-kilobase pair (kbp)-long randomly labeled dsDNA was used as the substrate. Reaction products were separated by 1% agarose gel electrophoresis. Red asterisks indicate random DNA labeling.

(k) Quantification of experiments such as shown in panel (j); error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation without RAD51 (lane 4 in panel j).
Figure 6. Nucleolytic degradation is prevented by RAD51 upon binding to dsDNA

(a) Nuclease assays with MRE11-RAD50 (MR) and their inhibition by RAD51. The experiments were carried out with blunt-ended, 5'-overhanged or 3'-overhanged DNA. Asterisk indicates the position of the labelling. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis. A representative experiment is shown.

(b) Quantifications of experiments such as shown in panel (a), error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation for each substrate without RAD51 (lanes 3, 7 and 11 in panel a).

(c) Nuclease assays with EXO1 and its inhibition by RAD51. Experiments were carried out with blunt-ended, 5'-overhanged or 3'-overhanged DNA. Asterisk indicates the position of the 32P label. Reaction products were separated by 15% denaturing polyacrylamide gel electrophoresis. Shown is a representative experiment.

(d) Quantifications of experiments such as shown in panel (c), error bars indicate SEM of three replicates. The level of DNA protection by RAD51 is presented as a relative value with respect to DNA degradation for each substrate without RAD51 (lanes 2, 7 and 12 in panel c).
Figure 7. Model for replication fork reversal and protection.

Fork remodelers have unequal biochemical functions. SMARCAL1 anneals RPA-coated ssDNA and may promote initial steps in fork reversal. ZRANB3 and HLTF are more proficient in branch migration. RAD51 and BCDX2 interact with SMARCAL1 and ZRANB3 and promote their activities. Reversed replication forks are prone to pathological degradation, in certain genetic backgrounds, unless protected by RAD51. We show that unexpectedly the dsDNA-binding capacity of RAD51 promotes DNA protection against nucleases.
Supplementary Files

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