The Main Role of Srs2 in DNA Repair Depends on Its Helicase Activity, Rather than on Its Interactions with PCNA or Rad51

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ABSTRACT  Homologous recombination (HR) is a mechanism that repairs a variety of DNA lesions. Under certain circumstances, however, HR can generate intermediates that can interfere with other cellular processes such as DNA transcription or replication. Cells have therefore developed pathways that abolish undesirable HR intermediates. The Saccharomyces cerevisiae yeast Srs2 helicase has a major role in one of these pathways. Srs2 also works during DNA replication and interacts with the clamp PCNA. The relative importance of Srs2's helicase activity, Rad51 removal function, and PCNA interaction in genome stability remains unclear. We created a new SRS2 allele [srs2(1-850)] that lacks the whole C terminus, containing the interaction site for Rad51 and PCNA and interactions with many other proteins. Thus, the new allele encodes an Srs2 protein bearing only the activity of the DNA helicase. We find that the interactions of Srs2 with Rad51 and PCNA are dispensable for the main role of Srs2 in the repair of DNA damage in vegetative cells and for proper completion of meiosis. On the other hand, it has been shown that in cells impaired for the DNA damage tolerance (DDT) pathways, Srs2 generates toxic intermediates that lead to DNA damage sensitivity; we show that this negative Srs2 activity requires the C terminus of Srs2. Dissection of the genetic interactions of the srs2(1-850) allele suggest a role for Srs2's helicase activity in sister chromatid cohesion. Our results also indicate that Srs2's function becomes more central in diploid cells.

IMPORTANCE  Homologous recombination (HR) is a key mechanism that repairs damaged DNA. However, this process has to be tightly regulated; failure to regulate it can lead to genome instability. The Srs2 helicase is considered a regulator of HR; it was shown to be able to evict the recombinase Rad51 from DNA. Cells lacking Srs2 exhibit sensitivity to DNA-damaging agents, and in some cases, they display defects in DNA replication. The relative roles of the helicase and Rad51 removal activities of Srs2 in genome stability remain unclear. To address this question, we created a new Srs2 mutant which has only the DNA helicase domain. Our study shows that only the DNA helicase domain is needed to deal with DNA damage and assist in DNA replication during vegetative growth and in meiosis. Thus, our findings shift the view on the role of Srs2 in the maintenance of genome integrity.

KEYWORDS  DNA recombination, DNA repair, PCNA, Rad51, Srs2, genome stability, yeasts

Homologous recombination (HR) is important for maintaining the stability of the genome; it helps repair double-strand breaks (DSBs) and participates in the recovery of damaged replication forks. However, HR mechanisms can generate intermediates that may block replication forks, or nucleoprotein complexes that can lead to cell cycle arrest and even cause cell death in certain genetic backgrounds (1). That is why HR must be tightly regulated to prevent untimely events that could interfere with other DNA replication or repair mechanisms.

The yeast Saccharomyces cerevisiae is an excellent model to isolate and study
mutants that shed light on the processes that maintain genome stability (2, 3). The Srs2 helicase has a major role in HR regulation; it is generally thought that its role is to suppress HR events at an early stage by dismantling the Rad51-presynaptic filament (4, 5). This “antirecombinase” role of Srs2 was first inferred from genetic studies: srs2 mutants show a hyperrecombination phenotype believed to be caused by an inappropriate channeling of the lesions into the homologous recombination pathway (6–9). The Srs2 protein exhibits single-stranded DNA (ssDNA)-dependent ATPase activity that unwinds DNA with 3′→5′ polarity with a $k_{cat}$ of ≈3,000 min$^{-1}$ (10, 11), and the Walker A motif is absolutely required for both ATPase and helicase activities (12). It can unwind a variety of substrates, including those containing forks, flaps, D-loops, 3′ and 5′ single-stranded DNA overhangs, blunt-end double-stranded DNA (dsDNA) substrates, and Holliday junctions (11, 13). Biochemical and electron microscopy analysis revealed that Srs2 can efficiently dismantle the presynaptic filament formed by Rad51, an early HR intermediate (4, 5). It seems that the helicase activity is not responsible for the dissociation, but rather Srs2’s ATP hydrolysis fuels a translocase activity: mutants that cannot bind or hydrolyze ATP fail to disrupt Rad51-presynaptic filaments (12). ATPase mutants show the same sensitivities to genotoxic agents, hyperrecombination phenotype, and genetic interactions as the srs2 deletion mutant (12). Some studies suggest that Srs2 is guided to the Rad51 filament through a physical interaction with Rad51 (5). Rad51 that cannot interact with Srs2 is resistant to Srs2 antirecombinase activity (14, 15). Other studies suggested that the direct interaction between Srs2 and Rad51 not only targets Srs2 to the HR intermediates but also triggers ATP hydrolysis within the Rad51 filament, causing Rad51 to dissociate from DNA (16). It seems therefore that Srs2 dismantles Rad51 by ATP-driven motor activities of Srs2 that can dissociate both DNA structures and protein-DNA complexes. Recently, it was also shown that Srs2 is able to disrupt extended D-loops created by the activity of polymerase δ (17). Moreover, in vitro experiments have shown that Srs2 can unwind structures that resemble D-loops (recombination intermediates) and that this activity is stimulated by Rad51 bound to dsDNA (18).

Srs2 is also needed in the restart of collapsed replication forks together with other members of the Rad6 epistasis group in a process called DNA damage tolerance (DDT). In fact, SRS2 was first identified because mutations in the gene could suppress the DNA damage sensitivity of both rad6 and rad18 mutants (suppressor of BAD six mutant 2), and this suppression requires functional HR (19–24). The main function of the error-free DDT pathway (which includes the Rad6, Rad18, Rad5, Ubc13, and Mms2 proteins) is to ubiquitinate PCNA at its lysine at position 164. If this step is not accomplished, Srs2 is recruited to the replication forks through its binding to SUMOylated PCNA (mediated by adjacent SIM-SUMO-interacting and PIM-PCNA-interacting motifs, which reside at the very end of the protein), and this recruitment seems to prevent HR (23, 25, 26). Thus, when the DDT pathway is impaired, Srs2 activity prevents a possible alternative rescue, resulting in sensitivity to DNA damage. Mutations in SRS2 seem to open the path for HR and thus suppress the sensitivity of DDT mutants (23, 24).

Although initially Srs2 was considered an inhibitor of HR, later work showed additional roles for Srs2 that favor HR. Srs2 was shown to be required for the repair of DSBs. Cells deleted for SRS2 show low survival when a single DSB is created, and it appears to act during HR repair, possibly by unwinding the invading strand from the D-loop to allow reannealing with the other broken chromosomal arm (27). Accordingly, Srs2 acts in the promotion of synthesis-dependent strand annealing (SSA) and inhibition of crossover events (28–30), as well as in additional forms of HR, such as single-strand annealing (SSA), break-induced replication (BIR), as well as in nonhomologous end joining (NHEJ) (31–35).

In recent years, new roles of Srs2 were identified. Srs2 association with SUMOylated PCNA was shown to limit the DNA synthesis by detaching polymerases δ and η from PCNA; this function is independent of the interaction with Rad51 (36). Moreover, Srs2 helicase activity can unwind triplet repeat hairpins at the replication fork; this activity is also independent of Rad51 and plays a critical role in maintaining normal replication without expansion or contraction of repeats (37). Another role for Srs2 is in preventing mutations as
a result of Top1 topoisomerase activity on misincorporated ribonucleotides. Srs2 can process the nick after Top1 activity and promote resection by enhancing Exo1 activity. Again, this role of Srs2 was shown to be Rad51 independent (38). Recently, in vitro experiments showed that Srs2 can remove replication protein A (RPA) and Rad52-RPA complex from ssDNA, although the in vivo significance of these findings is still unclear (39).

Altogether, Srs2 functions as a multifunctional tool that acts in replication, recombination, and DNA repair. By creating a new Srs2 allele that lacks the whole C terminus \[srs2(1-850)\], we show here that the DNA helicase domain alone is sufficient to deal with various types of DNA damage, to complete efficient DSB repair, and to promote meiosis. The synthetic lethality (SL) of \(\Delta srs2\) with other deletion mutations is also largely dependent on the DNA helicase activity. The analysis of the SL screen suggests that Srs2 might be involved in sister chromatin cohesion (SCC). Last, we show that the ploidy state of the cell dictates the importance of Srs2’s activity, and diploids rely more on the helicase’s C terminus in order to maintain genome stability.

RESULTS

The helicase domain of Srs2 is the major player in dealing with DNA damage. Our recent study provided evidence that Srs2 has functions that are independent of its role in the eviction of Rad51 and of its interaction with PCNA. Neither the Rad51 interaction region, PIM (PCNA-interacting motif), nor SIM (SUMO-interacting motif), which allow recruitment of Srs2 to SUMOylated PCNA, are required to deal with DNA damage caused by methyl methanesulfonate (MMS) (34).

We were interested in further investigating the importance of the helicase domain for genome stability. To answer this question, we created a new truncation mutation of Srs2 that lacks 324 amino acids (aa) from the C terminus (out of 1,174 aa). This mutation \[srs2(1-850)\] lacks all the known interaction sites of Srs2, such as those needed to interact with PCNA, Rad51, Nej1, Mre11, Sgs1, Esc2, Ubc9, Siz1, Siz2, Mus81, Rad5, and Rad18 (23, 40–44). Figure S1 in the supplemental material shows that the \(srs2(1-850)\) strain produces protein, which is expressed from its natural promoter at a slightly higher level than the level produced from the wild type (wt). Importantly, this N-terminal region of Srs2 has been shown to lack the ability to bind Rad51 (15).

We first tested how strains carrying this allele handle DNA damage. Figure 1 shows that, surprisingly, the mutants are as proficient as a wt strain when it comes to handling different kinds of DNA damage (MMS [DNA alkylation], hydroxyurea [deoxynucleoside triphosphate {dNTP} depletion], camptothecin [topoisomerase poison], and zeocin [DSBs]). These results imply that Srs2 deals with DNA damage through its helicase region and that the interactions with other proteins are dispensable for its main DNA repair activity. \(\Delta srs2\) mutants are more sensitive to DNA damage as diploid cells than as haploid cells. Consistent with a more central role of HR repair in diploids (34, 45), \(\Delta srs2\) diploids are more sensitive to the DSB-forming agent zeocin. We show that, in contrast, a diploid strain homozygous for the \(srs2(1-850)\) allele is as proficient as the wt parent for growth in the presence of DNA-damaging agents, with the possible exception of MMS, where a barely detected defect can be seen (Fig. 1).

The activity of Srs2 creates toxic intermediates in strains impaired in the DDT pathway and sensitizes them to DNA damage. Deletion of \(SRS2\) suppresses the DNA damage sensitivity (24). This suppression was shown to be caused by mutations in the C terminus of Srs2 (the SIM and PIM motifs) (23, 42, 46, 47). As expected, the \(srs2(1-850)\) allele (also lacking these motifs) suppressed the MMS sensitivity of \(pol30-K164R, \Delta rad18,\) and \(\Delta rad5\) mutants that are impaired in the DDT pathway (Fig. 1E). Thus, cells with the \(srs2(1-850)\) allele act like the wt when the cells are confronted with external insults to their DNA, but when the DDT pathway is inactivated, it behaves like a mutant with the whole \(SRS2\) gene deleted. The helicase part of Srs2 is important for dealing with DNA damage; however, when there are no modifications on lysine 164 of PCNA, Srs2 exerts its negative effects through its C terminus, probably via its interactions with PCNA.

The C terminus of Srs2 is dispensable for Srs2’s role in DNA repair during replication and DSB repair. Truncation of the C terminus of Srs2 does not sensitize the
cells to DNA damage. As Srs2 is involved in several repair pathways, we characterized
the repair capacity of the \textit{srs2}(1-850) mutant. First, we measured the ability of cells
carrying the allele to carry out homologous recombination (48). Strain MK166 allows
the measurement of the rates of ectopic gene conversion (GC) and direct-repeat
recombination (DRR) during normal cell division (Fig. S2). Relative to the wt, a strain
deleted for \textit{SRS2} showed elevated rates of DRR and GC of about 1.5- to 2-fold (49).
In contrast, an isogenic strain with the new allele showed levels of both GC and DRR
similar to those of the wt (Fig. 2A).

Next, we tested whether the \textit{srs2}(1-850) mutant is proficient for the repair of a single
DSB. Since diploid \textit{srs2}(1-850) strains showed some minor sensitivity to MMS, we tested the
proficiency of the allele in the repair of an inducible DSB in both haploid and diploid strains.

In the strains used, a single, defined DSB break is created by an inducible HO
endonuclease; in the haploid strain, the cells can then repair the damage by an ectopic
gene conversion and thus survive and form a colony. Two different diploid strains were
used: in the first strain (allelic), the DSB can be repaired by a gene conversion event in
which the donor sequence originates at the homologous chromosome. In the second
strain (ectopic), the two copies of chromosome V undergo DSBs, and they can be
repaired only by recombination with the ectopic donor (Fig. S3). By comparing the
number of colonies created when cells are plated on galactose-containing media
(continuous HO expression) versus glucose-containing media (no DSB creation), it is
possible to calculate the efficiency of repair. Figure 2B shows that wt diploids exhibit an
efficiency of repair close to 100% in the presence of an allelic donor. Haploids and
diploids that can repair the broken chromosome only by ectopic recombination show
about 60% survival. \textit{SRS2} is essential for DSB repair: \textit{Δ}srs2 strains exhibit very low repair
efficiency in haploid and diploid strains; they are defective for both allelic and ectopic
recombination. In contrast, the \textit{srs2}(1-850) mutant did not show any significant difference
from the wt in any of the three systems tested (Fig. 2B).

\textbf{FIG 1} The \textit{srs2}(1-850) mutant is fully proficient in dealing with DNA damage. (A to D) The \textit{srs2}(1-850) mutant is as resistant as the wt to MMS (A), hydroxyurea
(HU) (B), camptothecin (CPT) (C), and zeocin (D). (E) The \textit{srs2}(1-850) mutant suppresses the DNA damage sensitivity of impaired DDT mutants.
Since srs2 does not seem to affect interchromosomal gene conversion, we tested whether it might have an effect on intrachromosomal recombination. Using strain NA3 (Fig. S4), we measured the abilities of the various strains to repair the DSB by intrachromosomal (Rad51 dependent) gene conversion or by single-strand annealing (SSA), which is Rad51 independent. Again, the efficiency of repair is assessed by comparing the number of cells able to form colonies on galactose- versus glucose-based medium. The wt strain showed a repair efficiency of about 85%, whereas the Δsrs2 mutant had less than 2% repair (Fig. 2C). These results confirm that SRS2 is also required for intrachromosomal recombination initiated by a DSB. In contrast, the srs2(1-850) mutant exhibited a repair efficiency similar to that of the wt; furthermore, the distribution between intrachromosomal GC and SSA was similar to that of the wt (Fig. 2D). Taken together, our results show that only the helicase activity of SRS2 is needed for the repair of DSBs by all types of recombination tested, whereas the C terminus is dispensable.

The helicase domain of Srs2 is sufficient to undergo proficient meiosis. The Srs2 protein has a pivotal role in meiotic progression (51). Δsrs2 diploid cells are unable to undergo a proper meiosis and form few ascis. Moreover, most of these ascis give rise to dead spores. These defects are caused by the need of Srs2 for efficient homologous recombination in meiosis, which is essential for proper chromosomal segregation during the meiotic divisions (51).

To test the srs2(1-850) allele for possible meiotic defects, we subjected wt, Δsrs2/srs2, and srs2(1-850)/srs2(1-850) diploids to meiosis. Diploid cells were allowed to
sporulate for 6 days, and the percentage of cells that completed meiosis to form asci was determined. \( \Delta srs2 \) homozygotes formed very few asci, and most of the spores from these asci were unable to form colonies (Fig. 3A). Homozygous \( srs2(1-850) \) diploids, in contrast, showed no impairment in meiosis. Tetrads dissection showed that, contrary to what is seen in \( \Delta srs2/\Delta srs2 \) strains, viability of the \( srs2(1-850)/srs2(1-850) \) spores was as high as that of the wt spores (Fig. 3B). We conclude that \( srs2(1-850) \) does not have meiotic defects, again pointing to Srs2’s helicase domain as the main need for Srs2’s activity during meiosis.

Genetic interactions of the \( \Delta srs2 \) and \( srs2(1-850) \) alleles. The data we showed so far provide strong evidence for the hypothesis that the main role of Srs2 in genome stability is carried out by the helicase activity present at its N terminus and does not require interactions with additional repair proteins or with PCNA. Systematic screens (52–54) have shown that there is a large number of genes that, when deleted, show a dependence on Srs2 function for survival. For most of the genes, it is unknown why, if mutated, they are synthetic sick or lethal with \( \Delta srs2 \). We saw an opportunity to identify the regions of Srs2 that are responsible for the impaired growth in these genetic backgrounds. We chose genes involved in DNA metabolism that, when deleted, show the most severe negative interactions with the \( \Delta srs2 \) allele. Diploid strains heterozygous for various deletion allele and for \( srs2(1-850) \) or \( \Delta srs2 \) were subjected to meiosis, and tetrads were dissected. The viability and growth rate of double mutant spores defective for each of the chosen genes and carrying the \( srs2(1-850) \) allele was compared to that of the double mutant with \( \Delta srs2 \). The results (Table 1) are grouped according to their phenotypes. We could distinguish three categories.

1. The first set of deleted genes showed synthetic lethality with \( \Delta srs2 \) but grew normally when combined with \( srs2(1-850) \) (Table 1A and Fig. 4A).
   - **Ctf18.** Ctf18 is part of a replication factor C (RFC)-like complex that moves with the replication fork and participates in sister chromatid cohesion and checkpoint response (55).
   - **Csm3.** Csm3, together with Tof1, is located at the replication fork, where it contributes to fork stability by inhibiting fork rotation caused by topological stress of unwinding the dsDNA ahead of the polymerases (56).
   - **Mrc1.** Mrc1 is involved in replication checkpoint activation; it mediates phosphorylation of Rad53 by Mec1 and is active through S phase. It is also required for proper DNA replication (57, 58). The synthetic lethality between \( \Delta mrc1 \) and \( \Delta srs2 \) was found to be independent of the checkpoint activity of \( MRC1 \); the \( mrc1-AQ \)
allele, defective in the checkpoint function of MRC1, does not show synthetic sickness or lethality with \( \Delta srs2 \) (53). Site-specific mutations of MRC1 were found to impair specifically its role at the replication fork but not in DNA damage signaling (59). We crossed one such MRC1 mutant (mrc1-C14) with \( \Delta srs2 \) and observed synthetic lethality (SL). However, when combined with mrc-C14, the srs2(1-850) allele led to normal cell growth (Table 1A). We thus conclude that the DNA replication functions of Mrc1 and the helicase activity of Srs2 are synthetic lethal, and not the other functions of these proteins.

- **Rad9.** Rad9 is a checkpoint adapter, which is required throughout the cell cycle (60–62).

- **Ctf4 and Chl1.** The Ctf4 and Chl1 proteins are involved in sister chromatid cohesion and genome integrity and interact with Ctf18 (63, 64).

- **Rrm3.** Rrm3 is another helicase that assists in the replication of regions of the genome with secondary structures or bound proteins (65, 66).

### TABLE 1 Summary of synthetic interactions with \( \Delta srs2 \) and srs2(1-850)\(^\text{a}\)

| Synthetic lethal with \( \Delta srs2 \) | Normal with srs2(1-850) |
|-----------------|-----------------|
| Acf18           |                 |
| Acsm3           |                 |
| Atof1           |                 |
| Amrc1           |                 |
| Amrc1-C14       |                 |
| Arad9           |                 |

| Synthetic sick with \( \Delta srs2 \) | Normal with srs2(1-850) |
|-----------------|-----------------|
| Acfl4           |                 |
| Achl1           |                 |
| Arrm3           |                 |
| Amrc1           |                 |
| Axrs2           |                 |
| Aasf1           |                 |
| Art109          |                 |
| Amms22          |                 |
| Art107          |                 |
| Ahst3           |                 |
| Aslx5           |                 |
| Avid22          |                 |
| Aelg1           |                 |

| Synthetic lethal with \( \Delta srs2 \) | Synthetic sick with srs2(1-850) |
|-----------------|-----------------|
| Apo12           |                 |
| Arad54          |                 |
| Asgs1           |                 |
| Atop3           |                 |

\(^{a}\)A synthetic sick phenotype is observed when double mutant spores generate colonies smaller than single mutants; synthetic lethality is when spores are unable to generate colonies at all. At least 14 tetrads where dissected for each mutation combination.

![FIG 4 Genetic analysis of genes which are synthetic sick with \( \Delta srs2 \) but grow normally with srs2(1-850). (A to C) Examples of different phenotypes observed when crossed with the SRS2 allele.](image-url)
● **Mre11 and Xrs2.** The Mre11 nuclease and Xrs2, together with Rad50, form the MRX complex, which is important for end resection after DSB creation and for replication fork stability (67).

● **Histone acetylation proteins Asf1, Rtt109, Mms22, and Rtt107.** *ASF1, RTT109, MMS22 and RTT107* encode proteins that are involved in the acetylation of newly deposited histones (marked by an acetyl group in the lysine 56 of histone H3 [H3K56]) (54). This acetylation is important for proper DNA replication and DNA damage response (68).

● **Hst3.** Hst3 is the histone deacetylase that removes the acetyl groups from H3K56 (69). Surprisingly, both lack of acetylation and too much acetylation have a similar synthetic sick phenotype in the absence of Srs2 helicase activity.

● **Slx5.** Slx5 forms with Slx8 a SUMO-targeted ubiquitin (Ub) ligase (STUbL) complex that attaches ubiquitin to poly-SUMOylated proteins, supposedly in order to send them for degradation during replication and DNA repair (70).

● **Vid22.** Vid22 acts as a chromatin remodeler and removes nucleosome from DNA damage repair site. This in turn allows the recruitment of the MRX complex, which initiate the repair (71).

● **Elg1.** The RFC-like complex composed of Elg1 and four of the small subunits of RFC unloads PCNA during DNA replication and repair (72–74).

3. A third group of mutants were synthetic lethal with Δ*srs2* and showed synthetic sickness with the *srs2(1-850)* allele (Table 1C and Fig. 4C). This implies that both Srs2's helicase activity and the C terminus of Srs2 are required for proper cell activity in these genetic backgrounds.

● **Pol32.** Pol32 is a subunit of polymerase α required for efficient DNA synthesis and BIR (break-induced replication) repair (75, 76).

● **Rad54.** Rad54 is a chromatin remodeling factor that is needed for DSB repair; it participates in D-loop formation, extension, and resolution (77).

● **Sgs1, Rmi1, and Top3.** The Sgs1, Rmi1, and Top3 proteins form a complex required for many different aspects of genome stability and DNA repair, including DNA resection, the resolution of Holiday junction intermediates, and the relaxation of supercoiled DNA (78).

Taken together, the results point to the fact that in the absence of Srs2 function, histone deposition, checkpoint activation, and sister chromatid cohesion become impaired (see Discussion).

**Ploidy dictates Srs2 activity.** Previous results have shown that in certain genetic backgrounds (for example, in the absence of the Elg1 RFC-like subunit), deletion of SRS2 has little effect in haploids, but diploids fail to form colonies (72). We therefore tested the double mutants that showed normal growth as haploids for their phenotype as diploids. Double mutants were mated, and the homozygous diploid zygotes (>24 per strain) were manipulated to predetermined locations on rich medium plates. All mutants that showed synthetic sickness with Δ*srs2* were not viable as diploids and generated no, or only a few, viable colonies. Thus, all the genetic interactions of Δ*srs2* are stronger in diploids than in haploids. In contrast, most of the combinations of the *srs2(1-850)* allele and various deletions that showed normal growth as haploids were able to form viable diploids (although in some cases only 2/3 of the zygotes grew). The striking exceptions to this rule were diploids homozygous for the *srs2(1-850)* allele and for Δ*rad54, Δpol32, Δsgs1, or Δtop3*. Whereas these double mutant strains grew slowly as haploids, they failed to form diploid colonies, similarly to the double mutants with Δ*srs2*. This illustrates that in certain situations, the C terminus of Srs2 becomes important in diploids (see Discussion).

In conclusion, Srs2 has various roles in DNA replication and chromosome maintenance, which depend on the genome state of the cell. In haploids and diploids, the DNA helicase activity of *SRS2* is required for supporting proper DNA replication and chromosome...
segregation. However, in some genetic backgrounds (such as in the absence of Pol32, Rad54, or Sgs1), the C terminus is also important for reliable DNA replication in diploids.

**DISCUSSION**

The *srs2* mutant was originally isolated as a suppressor of the DNA damage sensitivity of mutants with an impaired DDT pathway; genetic evidence suggested that this suppression depends on Rad51 (6, 8, 24). *In vitro* experiments showed that Srs2 is able to disrupt Rad51 nucleofilaments (4, 5) and inhibit recombination at D-loops and replication forks by binding to PCNA (4, 5, 23). After these convincing biochemical experiments, it was widely assumed that the role of Srs2 in the maintenance of genomic integrity is to inhibit recombination by removing Rad51 from the DNA. Thus, all phenotypes of Δsrs2 were interpreted in light of this activity.

Although many times described as an “antirecombinase,” Srs2 is essential for DSB repair by HR (27); thus, its activity is both pro- and antirecombinational (44). Moreover, even Srs2 alleles that lack the region required for interactions with Rad51 or PCNA (or with any of the proven Srs2 interactors) are still proficient in promoting synthesis-dependent strand annealing (SDSA) over crossover resolution (79) and perfectly complement the sensitivity of Δsrs2 mutants to DNA damage (34, 37, 38, 80 this work). The new *srs2* allele [*srs2(1-850)*], which lacks the entire C terminus and has only the DNA helicase domain (15), is unable to interact with Rad51 (15) or with any of the known partners of Srs2 (PCNA, Rad51, Nej1, Sgs1, Esc2, Ubc9, Siz1, Siz2, Mus81, Rad5, and Rad18 [23, 40–44]). We showed that *in vivo*, the helicase of Srs2 was enough to fully deal with various DNA-damaging agents and with HO-induced DSB in both haploids and diploids. Mitotic recombination, meiosis progression, and spore survival were also unaffected. Only when the DDT pathway was impaired was the helicase domain insufficient to enable viability. Thus, binding to PCNA through the C-terminal PIM and SIM motifs becomes essential in the absence of the PCNA ubiquitination that allows DDT pathways to work. Our work thus defines two separate domains of the Srs2 protein with different biological relevance.

**Srs2 supports sister chromatid cohesion.** Srs2 plays an important role during DNA replication and chromosome segregation, as evidenced by strong negative genetic interactions with mutants defective in these processes. Our analysis showed that the *srs2(1-850)* allele is less affected than the Δsrs2 allele to inactivation of additional DNA processing functions. This implies that the helicase activity of Srs2 is sufficient for normal growth in most of the mutant backgrounds.

Csm3, Tof1, Ctf18, Ctf4, Mrc1, Skx5, and Elg1 have many diverse roles in keeping genome stability. Analyzing the common role between these proteins revealed that they all have a function in sister chromatid cohesion (SCC) (53, 55, 64, 81–84). In addition to the DNA helicases (Chl1, Sgs1, and Rrm3), Mre11 and Srs2 are also involved in SCC (85, 86). Interestingly, Δsrs2 and Δmre11 synthetic lethality was not dependent on the nuclease activity of the MRX complex (which is necessary for MRX’s role in end resection) or on active HR (87, 88). These results suggest that Srs2 and Mre11 are required for proper DNA replication, but not in their classical role of repairing the DNA damage during replication. Thus, their alternative role in DNA replication could also be in SCC, as the MRX complex has been shown to affect this process (89).

Asf1, Rtt109, Rtt107, Hst3, and Mms22 also have a role in SCC, as histone acetylation metabolism was found to act in the regulation of SCC (90–92). It seems that proper regulation of histone H3 acetylation is important for chromosome cohesion and segregation.

Two nonessential pathways were proposed to promote SCC. The first pathway is composed of Tof1, Csm3, Ctf4, and Chl1, and the second pathway is composed of Mrc1, the Ctf18 RFC-like complex, and the Sgs1-Top3-Rmi1 complex (82, 93). Srs2 does not seem to belong specifically to one of the nonessential SCC pathways. Rather, it has a supporting role for the two SCC pathways. This is evident also by the supporting role of Srs2 in histone acetylation metabolism during SCC. The role of helicases in SCC is unknown; it was suggested that helicases might prepare the DNA for targeting of new
cohesin rings by removing old cohesin units left on the DNA and by stimulating the loading of new cohesins during replication (94, 95). These functions could be executed by Srs2’s DNA helicase and translocase capabilities.

The C terminus of Srs2 is required for specific functions during DNA replication to promote genome integrity. In contrast to the previous lack of synthetic phenotypes, the srs2(1-850) allele was synthetic sick when combined with Δgs1, Δpol32, or Δrad54.

Both Pol32 and Srs2 are implicated in BIR (76). BIR is divided into two pathways: a Rad51-dependent branch and a Rad31-independent branch (96). In the absence of POL32, the C terminus of Srs2 is partly required for cell viability. The C terminus contains the Rad51-interacting motif, a fact that may implicate Srs2 in the Rad51-dependent BIR. Alternatively, it may be the interaction of Srs2 to PCNA that is required. When the Srs2 pathway is disrupted, cells become completely dependent on the Pol32-mediated repair pathway (97).

The synthetic lethality between Δsrs2 and Δrad54 is more complex. It has been proposed that Srs2 and Rad54 actually act in the same pathway, and the SL interaction is due to the generation of toxic intermediates that are trapped and making the cells unable to proceed with the repair without Rad54, but the generated intermediates also cannot recede to an alternative repair pathway due to lack of Srs2 antirecombinase activity (98). This is consistent with our results showing that the C terminus (Rad51 interaction region) of Srs2 is required for proper DNA replication in Δsrs2 Δrad54 cells.

A similar explanation could also be applied to the synthetic sickness of the srs2(1-850) allele in the absence of a functional Sgs1 helicase. Sgs1’s activity affects many stages of the HR process, from resection to resolution, as well as having a role in SCC (99).

Ploidy regulates a wider range of Srs2’s activities. All of the double mutants with Δsrs2 that are synthetic sick as haploid cells become essential in diploid cells. The DNA helicase of Srs2 is, however, sufficient to suppress the SL phenotype in diploids. This implies that the DNA helicase part of Srs2 becomes more central in diploids and is crucial for cell viability in the absence of other factors (72). In certain genetic backgrounds, the C terminus of Srs2 also becomes important in diploids. Mutants that are synthetic sick with srs2(1-850) as haploids (Δrad54, Δpol32, and Δgs1) are inviable as diploids. Ploidy seems to affect the fundamental regulation of the pathways involved in dealing with DNA replication stress. Haploids rely more on the DDT pathways, whereas diploids seem to rely more on HR (34, 45). It seems that PCNA and its modifications affect the regulation of DNA repair during replication, depending on the ploidy of the cell. The fact that diploids rely more on HR to deal with DNA damage is consistent with our finding that Srs2 C terminus and probably its antirecombinase activity is important in diploids, when other factors of HR are unavailable.

In conclusion, we show that the helicase activity of Srs2, and not its physical interactions with Rad51 or PCNA, plays a major role in genome maintenance. PCNA interaction becomes important only in the absence of the DDT pathway. We also show that Srs2 plays a role in SCC and that its helicase activity becomes more important in diploid cells.

MATERIALS AND METHODS

Yeast strains. Saccharomyces cerevisiae strains used in this study are listed in Table 2. Unless otherwise stated, strains used were of one of these backgrounds.

- **MK1166:** MATA lys2::Ty1Sup ade2-1(o) can1-100(o) ura3-52 leu2-3,112 his3del200 trpldel901 HIS3:: lys2::ura3 his4::TRP1::his4 (48).
- **MK203:** MATA::inc ura3::HOcs (V) lys2::ura3::HOcs inc (1.2 kb) ade3::GALHO leu2-3,112 his3-11,15 trpl-1 ade2-1 can1-100 (100). This strain is based on W303 (27).
- **NA3::** MK203 carrying pM53 ([URA3 + [1.2 kb] TRP1-1 [1.4 kb]]) integrated into ura3::HOcs and an additional donor in lys2. The genotype of strain NA3 is MATA::inc ade2 ade3::GALHO ura3::HOcs ---TRP1-1::URA3 (1.2 kb) leu2-3,112 his3-11,13 trpl-1 lys2::ura3::HOcs::inc (50).
- **Sch2::** MATA/MATa ura3::HOcs/URA3 (V) lys2::ura3::HOcs::incRB (1.2 kb) ade3::GALHO leu2-3,112 his3-11,13 trpl-1 ade2-1. (This strain is based on strain W303.)
- **Sch4::** MATA/MATa ura3::HOcs (V) lys2::ura3::HOcs::incRB (1.2 kb) ade3::GALHO leu2-3,112 his3-11,15 trpl-1 ade2-1 can1-100. (This strain is based on strain W303.)

Standard yeast molecular genetic techniques were used to delete individual genes.
### TABLE 2 Yeast strains used in this study

| Strain          | Relevant genotype                  | Reference or source |
|-----------------|------------------------------------|---------------------|
| MK166 diploid   | MATa/MATα                          | 48                  |
| AB101           | MK166 MATa                         | 48                  |
| AB217           | MK166 MATa mrc1::natR              | This study          |
| AB91            | MK166 MATa rad9::natR              | This study          |
| op883           | MK166 MATa srs2::kanMX             | 49                  |
| AB270           | MK166 MATa po30-K164::kanMX srs2::kanMX | This study          |
| op710           | MK166 MATa elg1::HygMX             | Lab stock           |
| op952           | MK166 MATa po30-K164::LEU2         | Lab stock           |
| AB106           | MK166 MATa po30-K164::LEU2 srs2::kanMX | This study          |
| AB365           | MK166 diploid srs2::kanMX          | This study          |
| AB366           | MK166 diploid srs2(1-850)::HygMX   | This study          |
| AB298           | MK166 MATa srs2(1-850)::HygMX      | This study          |
| OP1122          | MK166 MATa rad18::LEU2             | 49                  |
| op890           | MK166 MATa rad5::kanMX             | 49                  |
| OP1125          | MK166 MATa rad18::LEU2 srs2::kanMX | 101                 |
| AB234           | MK166 MATa rad5::kanMX srs2::kanMX | 101                 |
| AB353           | MK166 MATa srs2(1-850)::HygMX rad18::LEU2 | This study          |
| AB339           | MK166 MATa srs2(1-850)::HygMX po30-K164::kanMX | This study          |
| AB341           | MK166 MATa srs2(1-850)::HygMX rad5::kanMX | This study          |
| MK203           | MATa                                | Lab stock           |
| MK15514         | MK203 srs2(1-850)::HygMX           | This study          |
| S JB16          | MK203 srs2::kanMX                   | Lab stock           |
| NA3             | MATa                                | Lab stock           |
| S JB30          | NA3 srs2::LEU2                      | This study          |
| MK15519         | NA3 srs2(1-850)::HygMX             | This study          |
| Sch2            | MATa/MATα                           | Lab stock           |
| Sch 4           | MATα/MATα                           | Lab stock           |
| MK11208B        | Sch2 srs2::LEU2                     | Lab stock           |
| MK15575         | Sch2 srs2(1-850)::HygMX            | This study          |
| MK13120         | Sch4 srs2::LEU2                     | This study          |
| MK15576         | Sch4 srs2(1-850)::HygMX            | This study          |
| MK17285         | MK166 MATa csm3::kanMX              | This study          |
| MK17297         | MK166 MATa csm3::kanMX srs2(1-850)::HygMX | This study          |
| MK17298         | MK166 MATa csm3::kanMX srs2(1-850)::HygMX | This study          |
| MK17323         | MK166 MATa mrc1::natR srs2(1-850)::HygMX | This study          |
| MK17325         | MK166 MATa mrc1::natR srs2(1-850)::HygMX | This study          |
| AB297           | MK166 MATa srs2(1-850)::HygMX      | This study          |
| AB331           | MK166 MATa srs2(1-850)::HygMX      | This study          |
| MK44252         | MK166 MATa ctf18::HygMX             | Lab stock           |
| AB386           | MK166 MATα ctf18::Hyg srs2::kanMX   | This study          |
| AB388           | MK166 MATa ctf18::HygMX srs2::kanMX | This study          |
| AB390           | MK166 MATa ctf18::HygMX srs2(1-850)::HygMX | This study          |
| AB392           | MK166 MATa ctf18::HygMX srs2(1-850)::HygMX | This study          |
| AB367           | MK166 MATa srs2::kanMX elg1::HygMX  | This study          |
| AB368           | MK166 MATa srs2::kanMX elg1::HygMX  | This study          |
| AB369           | MK166 MATa srs2(1-850)::HygMX elg1::HygMX | This study          |
| AB3670          | MK166 MATa srs2(1-850)::HygMX elg1::HygMX | This study          |
| MK7232          | MK166 MATa rrm3::kanMX              | Lab stock           |
| AB379           | MK166 MATα rrm3::kanMX srs2::HygMX  | This study          |
| AB381           | MK166 MATα rrm3::kanMX srs2::HygMX  | This study          |
| AB382           | MK166 MATα rrm3::kanMX srs2(1-850)::HygMX | This study          |
| AB384           | MK166 MATα rrm3::kanMX srs2(1-850)::HygMX | This study          |
| op1149          | MK166 MATa ctfH::kanMX              | Lab stock           |
| AB394           | MK166 MATa ctfH::kanMX srs2::HygMX  | This study          |
| AB396           | MK166 MATa ctfH::kanMX srs2::HygMX  | This study          |
| AB398           | MK166 MATa ctfH::canMX srs2(1-850)::HygMX | This study          |
| AB400           | MK166 MATa ctfH::kanMX srs2(1-850)::HygMX | This study          |
| AB417           | MK166 MATa rtt109::kanMX            | This study          |
| AB424           | MK166 MATa rtt109::kanMX srs2::HygMX | This study          |
| AB426           | MK166 MATa rtt109::kanMX srs2::HygMX | This study          |
| AB428           | MK166 MATa rtt109::kanMX srs2(1-850)::HygMX | This study          |
| AB430           | MK166 MATa rtt109::kanMX srs2(1-850)::HygMX | This study          |
| AB421           | MK166 MATa xrs2::kanMX              | This study          |
| AB457           | MK166 MATa xrs2::kanMX srs2(1-850)::HygMX | This study          |

(Continued on next page)
| Strain     | Relevant genotype                              | Reference or source |
|------------|-----------------------------------------------|---------------------|
| AB459      | MK166 MATα xrs2:KanMX srs2(1-850):HygMX       | This study          |
| AB461      | MK166 MATα xrs2:KanMX srs2::HygMX             | This study          |
| AB463      | MK166 MATα xrs2:KanMX srs2::HygMX             | This study          |
| MK4097     | MK166 MATα mre11::KanMX                       | This study          |
| AB432      | MK166 MATα mre11::KanMX srs2::HygMX           | This study          |
| AB434      | MK166 MATα mre11::KanMX srs2::HygMX           | This study          |
| AB436      | MK166 MATα mre11::KanMX srs2(1-850)::HygMX     | This study          |
| AB438      | MK166 MATα mre11::KanMX srs2(1-850)::HygMX     | This study          |
| MK12598    | MK166 MATα chl1::KanMX                        | Lab stock           |
| AB440      | MK166 MATα chl1::KanMX srs2(1-850)::HygMX     | This study          |
| AB442      | MK166 MATα chl1::KanMX srs2(1-850)::HygMX     | This study          |
| MK7267     | MK166 MATα asf1::KanMX                        | Lab stock           |
| AB444      | MK166 MATα asf1::KanMX srs2::HygMX            | This study          |
| AB446      | MK166 MATα asf1::KanMX srs2::HygMX            | This study          |
| AB448      | MK166 MATα asf1::KanMX srs2(1-850)::HygMX      | This study          |
| AB450      | MK166 MATα asf1::KanMX srs2(1-850)::HygMX      | This study          |
| AB371      | MK166 MATα srs2::HygMX                        | This study          |
| AB372      | MK166 MATα srs2::HygMX                        | This study          |
| MK7781     | MK203 MATα vid22::NatR                        | Lab stock           |
| AB475      | MK203 MATα vid22::NatR srs2::LEU2             | This study          |
| AB477      | MK203 MATα vid22::NatR srs2::LEU2             | This study          |
| AB479      | MK203 MATα vid22::NatR srs2(1-850)::HygMX     | This study          |
| AB481      | MK203 MATα vid22::NatR srs2(1-850)::HygMX     | This study          |
| AB423      | MK166 MATα sxs5::KanMX                        | This study          |
| AB503      | MK166 MATα sxs5::KanMX srs2::HygMX            | This study          |
| AB505      | MK166 MATα sxs5::KanMX srs2::HygMX            | This study          |
| AB507      | MK166 MATα sxs5::KanMX srs2(1-850)::HygMX      | This study          |
| AB509      | MK166 MATα sxs5::KanMX srs2(1-850)::HygMX      | This study          |
| MK14408    | MK166 MATα pol32::KanMX                       | Lab stock           |
| AB411      | MK166 MATα srs2(1-850)::HygMX pol32::KanMX    | This study          |
| AB413      | MK166 MATα srs2(1-850)::HygMX pol32::KanMX    | This study          |
| AB134      | MK166 MATα rad54::KanMX                       | This study          |
| AB401      | MK166 MATα rad54::KanMX srs2(1-850)::HygMX    | This study          |
| AB403      | MK166 MATα rad54::KanMX srs2(1-850)::HygMX    | This study          |
| AB405      | MK166 MATα sgs1::KanMX srs2(1-850)::HygMX      | This study          |
| AB407      | MK166 MATα sgs1::KanMX srs2(1-850)::HygMX      | This study          |
| MK4137     | MK166 MATα sgs1::KanMX                        | Lab stock           |
| 17371      | MK166 MATα mrc1-C14::KanMX                    | This study          |
| 17376      | MK166 MATα mrc1-C14::KanMX srs2(1-850)::HygMX  | This study          |
| 17377      | MK166 MATα mrc1-C14::KanMX srs2(1-850)::HygMX  | This study          |
| AB491      | MK166 MATα top3::LEU2                        | This study          |
| 17396      | MK166 MATα hst3::KanMX                        | This study          |
| 17420      | MK166 MATα hst3::KanMX srs2::HygMX            | This study          |
| 17421      | MK166 MATα hst3::KanMX srs2::HygMX            | This study          |
| 17424      | MK166 MATα hst3::KanMX srs2(1-850)::HygMX      | This study          |
| 17425      | MK166 MATα hst3::KanMX srs2(1-850)::HygMX      | This study          |
| 17428      | MK166 MATα rad9::KanMX srs2::HygMX            | This study          |
| 17432      | MK166 MATα rad9::KanMX srs2::HygMX            | This study          |
| 17433      | MK166 MATα rad9::KanMX srs2(1-850)::HygMX      | This study          |
| 17388      | MK166 MATα rtt107::KanMX                      | This study          |
| 17436      | MK166 MATα rtt107::KanMX srs2::HygMX          | This study          |
| 17437      | MK166 MATα rtt107::KanMX srs2::HygMX          | This study          |
| 17440      | MK166 MATα rtt107::KanMX srs2(1-850)::HygMX    | This study          |
| 17441      | MK166 MATα rtt107::KanMX srs2(1-850)::HygMX    | This study          |

**Determination of recombination rates.** Strain MK166 carries substrates that allow easy scoring of direct-repeat recombination (DRR) (His+ colonies) and ectopic gene conversion (GC) (Lys- colonies). Colonies isolated from plates with various concentrations of methyl methanesulfonate (MMS) were subjected to fluctuation tests, and the rates were calculated as described previously (48). The MMS concentrations used were low and did not cause cell death in the wild-type (wt) strain.

**Repair efficiency measurement.** NA3, MK203, Sch2, and Sch4 strain derivatives were streaked onto yeast extract-peptone-dextrose (YPD) plates. Individual colonies were resuspended in water, appropriately diluted, and plated on YPD and yeast extract-peptone-galactose (YPGal) plates. The colonies were counted after 3 days of incubation at 30°C (27, 50).
SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.01192-18.

FIG S1, TIF file, 0.2 MB.
FIG S2, TIF file, 0.1 MB.
FIG S3, TIF file, 0.5 MB.
FIG S4, TIF file, 0.5 MB.

ACKNOWLEDGMENTS

This work was supported by grants from the Israel Science Foundation (ISF), the Israel Cancer Research Fund, and the Volkswagen Fund to M.K.

We thank present and past members of the Kupiec research group for encouragement, ideas, and support.

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