Avoidance of rNMP-induced mutations via RNaseH2 and Srs2-Exo1 dependent mechanisms

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Abstract

Srs2 helicase is known to dismantle nucleofilaments of the Rad51 recombinase to prevent spurious recombination events¹–³, 4, 5, 6 and unwind trinucleotide sequences that are prone to hairpin formation⁷. Here we document a new, unexpected genome maintenance role of Srs2 in the suppression of mutations arising from misinsertion of rNMPs during DNA replication. In cells lacking RNaseH2, Srs2 unwinds DNA from the 5′ side of a nick generated by DNA topoisomerase I⁸ at a rNMP residue. In addition, Srs2 interacts with and enhances the activity of the nuclease Exo1, to generate a DNA gap in preparation for repair. Srs2-Exo1 thus functions in a novel pathway of nick processing-gap filling that mediates tolerance of rNMPs in the genome. Our results have implications for understanding the basis of Aicardi-Goutieres Syndrome, which stems from inactivation of the human RNaseH2 complex⁹.

Mutants of SRS2 (hpr5) and RNH202 (hpr4), which encode a 3′ to 5′ helicase and a subunit of the RNaseH2 complex, respectively, are represented in our collection of hyper-rec mutants¹⁰. These mutants exhibit similar hyper-recombination levels¹⁰, DNA damage sensitivity, and interactions with DNA replication and damage repair genes (Extended Data Fig. 1). The rnh202Δ srs2Δ double mutant was tested to determine if RNaseH2 and Srs2 might act together. A truncation mutant srs2-860¹ that has helicase activity but lacks Rad51 interaction was used (Fig. 1a), as we wished to distinguish between DNA helicase and Rad51 removal activities in vivo. We disabled the ATP hydrolysis activity of srs2-860 through the K41A mutation (srs2-KA860)¹¹. The rnh202Δ srs2-KA860 cells grew slowly (Fig. 1b), revealing that Srs2 helicase activity is important for growth in rnh202Δ. Similar results were obtained with the rnh201Δ srs2Δ mutant, deficient in the catalytic subunit of RNaseH2 (data not shown). We note that the srs2-KA mutation in the full-length protein¹¹ also engenders a slow growth phenotype with rnh202Δ (Extended Data Fig. 2 a,b).
Cells deficient in RNH202, but not SRS2, accumulate spontaneous mutations\textsuperscript{10}. Interestingly, the rnh202 srs2 double mutant shows a much stronger mutator phenotype (Fig. 1c), indicating that SRS2 prevents mutations in rnh202 cells. The srs2-860 allele possesses this mutation avoidance function, but srs2-KA860 is defective in this regard. The srs2-860 mutant has a hyper-rec phenotype because its encoded protein lacks anti-Rad51 activity\textsuperscript{1}, and the rnh202 srs2-860 double mutant shows an additive increase in recombination that is partially dependent on the Srs2 helicase activity (Fig. 1d). Consistent with this finding, the rnh202 srs2 mutant exhibits increased Rad52 foci (Extended Data Fig. 3a), indicative of spontaneous DNA double strand breaks\textsuperscript{20}. The slow growth phenotype of the rnh202 srs2 mutant (Fig. 1a) is due to an increase in G2 cells and can be partially relieved by inactivating the DNA damage checkpoints (Extended Data Fig. 3b–c). Lastly, the DNA damage checkpoints are largely intact in these cells as shown by Rad53 phosphorylation following hydroxyurea (HU) treatment (Extended Data Fig. 3d), which elevates the rNTP:dNTP ratio. However, combining rnh202\textDelta with either srs2\textDelta or srs2-KA860 leads to a synergistic sensitivity to HU (Fig. 1e).

Ribonucleotide misincorporation becomes elevated in cells that harbor the pol2644G mutation in the replicative DNA polymerase POL\textepsilon\textsuperscript{24}, and the combination of rnh202 with pol2644G increases the ribonucleotide load further\textsuperscript{23}. Importantly, the rnh202 srs2 pol2644G triple mutation is lethal (Fig. 1f), and the lethality is rescued by srs2-860 but not srs2-KA860 (data not shown).

Loss of RNaseH2 causes slippage mutations in simple repeats without affecting base substitutions\textsuperscript{8,23}. We sequenced can1 alleles to determine the mutation signatures of various strains. As expected, rnh202\textDelta cells accumulate slippage mutations, including single nucleotide deletions in a mononucleotide run and dinucleotide deletions in dinucleotide repeats (Fig. 1g, Extended Data Table 1a). The srs2\textDelta mutant shows a slight increase in these mutation types, consistent with the role of Srs2 in preventing slippage in long trinucleotide repeats\textsuperscript{7}. The slippage mutations are increased in the rnh202 srs2 double mutant (Fig. 1g, Extended Data Table 1a). To further characterize this mutator phenotype, a reporter that detects slippages in AG repeats was used\textsuperscript{8}. With this reporter, srs2\textDelta is indistinguishable from wild type, while rnh202\textDelta shows a 90-fold increase. This phenotype of rnh202\textDelta is exacerbated by removing Srs2 or inactivating its helicase activity (Fig. 1h, Extended Data Fig. 2c). Altogether, the results show that Srs2 helicase guards against mutations induced by rNMPs in DNA, and they reveal an overlap in mutation sequences between rnh202 and rnh202 srs2 with novel slippage sites of ≥ 3 nucleotides in the double mutant (Extended Data Fig. 4, Extended Data Table 1a).

Mutations that arise in RNaseH2 deficient cells are dependent on topoisomerase I (Topo I), encoded by the TOP1 gene, which cleaves DNA at rNMP residues\textsuperscript{8,26}. We tested top1\textDelta to ask whether it is relevant for rnh202 srs2. Can\textsuperscript{r} mutations arise from different events, including gross chromosome rearrangements\textsuperscript{14}. In our test, top1\textDelta is epistatic to rnh202 (Fig. 2a). Importantly, rnh202 top1 shows a specific loss of slippage events of 2 nucleotides or greater (Fig. 2b, Extended Data Table 1a). The increased mutation rate in rnh202 srs2 is also suppressed by top1\textDelta (Fig. 2a), including a complete loss of slippage events of ≥ 2 nucleotides and a reduction in single nucleotide slippage (Fig. 2b, Extended Data Table 1a).
The HU hypersensitivity (Fig. 2c), the hyper-rec phenotype (Fig. 2d), and the lethality with pol2M644G (Fig. 2e) of rnh202 srs2 are also overcome by top1Δ. Thus, when topoisomerase I is present, the tolerance of rNMPs needs RNaseH2 and Srs2, and recombinogenic DSBs arise when RNaseH2 is absent.

Topo I-mediated cleavage of a rNMP results in unligatable 5′ hydroxyl and 3′ cyclic phosphate termini\(^26\). We reasoned that, in mutation avoidance, Srs2 unwinds DNA from the nick generated by Topo I. To test this, we constructed linear duplex DNA substrates whose ends are occluded by a Streptavidin-biotin complex. As expected, the intact substrate was resistant to unwinding by Srs2 (Extended Data Fig. 5). Importantly, when a nick bearing a 5′ OH group was introduced, ATP-dependent displacement of the strand containing this group occurred (Extended Data Fig. 5). In contrast, little unwinding of the 3′ OH bearing strand was seen (Extended Data Fig. 5).

To examine whether Srs2 unwinds DNA from a nick introduced by Topo I at an embedded ribonucleotide, a DNA substrate with a single UMP residue and blocked ends was prepared. As expected\(^26\), calf thymus Topo I, but not E. coli Topo I, cleaved the DNA/UMP hybrid strand, but not an equivalent substrate without the ribonucleotide (Extended Data Fig. 6a and 6b). Incubation of the Topo I-nicked DNA (Extended Data Fig. 6c) with Srs2 led to the ATP-dependent displacement of the strand with the 5′ OH group, but not that contains the 3′ cyclic phosphate (Fig. 3a). Mutations in MPH1, which encodes a 3′-5′ helicase, also lead to a mutator phenotype\(^25\). However, Mph1 protein does not act on the Topo I-nicked substrate (Extended Data Fig. 7a). Consistent with this, the rnh202Δ mph1Δ double mutant shows the same dinucleotide slippage rate as rnh202Δ (Extended Data Fig. 7b).

We surmised that Srs2-mediated unwinding of the 5′-strand at the Topo I-induced DNA nick prevents religation that is associated with DNA deletion and serves to initiate lesion removal by creating a DNA gap (see discussions later). If this were the case, one would expect a 5′ endo- or exo-nuclease to be involved in the removal of the displaced DNA strand. One strong candidate is Exo1, which has 5′ to 3′ exonuclease and 5′ FLAP endonuclease activities and plays a major role in DNA repair, including DNA break end resection during homologous recombination and DNA digestion during mismatch repair. Importantly, with the dinucleotide slippage reporter, we found that Exo1 has an antimutator activity against rnh202, and that it acts in an epistatic manner with Srs2 (Fig. 3b).

Being prompted by the genetic results, we next tested possible functional interactions between Srs2 and Exo1. For this, DNA substrates blocked at their ends and harbor a 5′ FLAP or a 5′ FLAP adjoining a gap were constructed; the latter substrate resembles the intermediate generated by Srs2-mediated DNA unwinding from a nick. We found that Exo1 acts endonucleolytically on these substrates, incising the single stranded arm of the FLAP structure (Extended Data Fig. 7c); little exonuclease activity was detected with these substrates or with an oligo dT substrate (Extended Data Fig. 7c). Importantly, Srs2 enhanced the endonucleolytic cleavage of the gapped 5′-FLAP DNA in an ATP-independent manner (Fig. 4a). Interestingly, by affinity pulldown, a physical interaction between Srs2 and Exo1 was detected (Fig. 4b). In contrast, we did not detect any enhancement of Exo1 activity by Mph1 (Extended Data Fig. 8a). Moreover, although human EXO1 (hEXO1) digests the 5′
FLAP substrate similarly to yeast Exo1, addition of Srs2 led to inhibition of the reaction (Extended Data Fig. 8b). We also tested the effect of Srs2 on 5′ FLAP cleavage mediated by yeast Dna2 and human FEN1 and found that Srs2 attenuates the cleavage efficiency in both cases (data not shown). These results support the premise that Srs2 and Exo1 co-operate, in a specific fashion, in the downstream processing of the DNA nick originating from Topo I-mediated cleavage at a rNMP.

hEXO1 can digest DNA exonucleolytically with 5′–3′ polarity at a nick17, and we verified that yeast Exo1 has the same activity (Extended Data Fig. 7c). To ask how Exo1 processes different DNA nick types, we prepared substrates that harbor a Topo I-induced nick, a nick with 3′ hydroxyl and 5′ phosphate groups, or a nick with 3′ and 5′ hydroxyl groups. As shown in Fig. 4c, Exo1 can process the Topo I-induced nick, although less efficiently than it does a clean DNA nick. Interestingly, Srs2 enhanced digestion of the Topo I-induced nick, by as much as four fold (Fig. 4c (i) and (iv)). A stimulatory effect of Srs2, although less pronounced, on Exo1 was also observed with the nicked substrate harboring a 5′-OH group (Fig. 4c (ii) and (iv)), but little enhancement occurred at the nick that harbors 3′ OH and 5′ phosphate groups (Fig. 4c (iii) and (iv)). Again, ATP is dispensable for the observed stimulation (Fig. 4c), which suggests Srs2 facilitates the delivery of Exo1 to the cleaved rNMP site. We found that Mph1 is unable to stimulate Exo1-catalyzed digestion of the Topo I-induced nick (Extended Data Fig. 8c), and that digestion by hEXO1 is attenuated by Srs2 (Extended Data Fig. 8d).

The genetic and biochemical data presented above support the model (Extended Data Fig. 9) where Srs2 unwinds DNA from the Topo I-generated nick and recruits Exo1, which utilizes its 5′ exonuclease or 5′ FLAP endonuclease activity to generate a DNA gap to be filled by a DNA polymerase. In the absence of Srs2, unwinding of DNA from the 5′ OH end is not possible and its processing by Exo1 also becomes inefficient. In this case, Topo I could act again at the 3′ cyclic phosphate, cleaving two nucleotides and religating the ends to generate a two-nucleotide deletion15.

Recent evidence has suggested a beneficial role of DNA nicks stemming from ribonucleotide excision repair (RER) in the facilitation of DNA mismatch removal from the newly replicated DNA18,21. However, rNMPs in DNA can be harmful, as they cause slippage and other mutations. To avoid such mutations, RNaseH2 initiates rNMP elimination via RER28. We have shown that in the absence of RNaseH2, the major mechanism to remove embedded rNMPs is cleavage by Topo I and nick processing by Srs2/Exo1, followed by gap repair. rNMPs cleaved by Topo I may also be converted into a recombinogenic DSB. Our work thus uncovers a novel function of Srs2 helicase and Exo1 in mutation avoidance and rNMP lesion repair. Since the Srs2 helicase activity is required for the avoidance of rNMP-induced mutations, the major repair mechanism in cells may entail Srs2-mediated DNA strand separation from the Topo I-induced nick, to create a 5′ FLAP for Exo1 to incise. The physical interaction between Srs2 and Exo1 likely serves to recruit the latter to the FLAP structure.
Methods Summary

Yeast strains are listed in Extended Data Table 1b

Mutation rates were determined by the Lea Coulson fluctuation method. Independent mutants were sequenced to identify mutations. Recombination assays were performed using the leu2-ecoRI::URA3::leu2-bstEII system as described\(^\text{10}\). For the (AG)\(_4\) reporter, rates were determined as described\(^\text{8,29}\). At least 18 independent cultures (minimum 2 isolates/genotype) were used to determine rates, and 95% confidence intervals were determined as described\(^\text{29}\).

In vitro assays

Srs2 and Exo1 were expressed in *E. coli* and Hi5 insect cells, respectively, and purified as described\(^\text{12,27}\). Purified hEXO1 was a gift from Guo-Min Li (University of Kentucky). Nuclease and helicase assays were conducted as described elsewhere; see Methods. Reactions were analyzed by gel electrophoresis and phosphorimaging.

Methods

*In vivo* assays

Yeast strains are listed in Extended Data Table 1b.

To determine mutation rates, *CAN1*\(^+\) strains were grown in 5 mL YPD overnight, washed and resuspended in 1 mL \(d\)H\(_2\)O. Cells were plated onto SC-arginine + 60 µg/mL canavanine to select for *can1* mutants and onto SC medium for total cell number. Plates were incubated at 30°C for 3–4 days. At least 18 independent cultures with a minimum of 2 different isolates per genotype were used to determine rates, calculated by the Lea and Coulson method of the median. 95% confidence intervals were determined as in\(^\text{29}\). Independent mutants were sequenced at the *CAN1* locus to identify mutations. Recombination assays were performed using the leu2-ecoRI::URA3::leu2-bstEII system as described\(^\text{10}\). Briefly, colonies were struck out onto YPD medium and grown for 3 days at 30°C. Whole single colonies were resuspended in 1 mL \(d\)H\(_2\)O. Cells were plated to SC-leucine, -uracil to select for recombinants and onto SC medium for total cell number. Plates were incubated at 30°C for 3–4 days. At least 18 independent cultures with a minimum of 2 different isolates per genotype were used to determine rates, calculated by the Lea and Coulson method of the median. 95% confidence intervals were determined as in\(^\text{29}\).

For the (AG)\(_4\) reporter, rates were determined as described\(^\text{8,29}\). Briefly, single colonies were inoculated into 5 mL YPD and grown for 2 days at 30°C, washed and resuspended in 1 mL \(d\)H\(_2\)O. Cells were plated onto SC-lysine medium to select for slippage mutants and onto SC medium for total cell number. Plates were incubated at 30°C for 3 days. At least 18 independent cultures with a minimum of 2 different isolates per genotype were used to determine rates, and 95% confidence intervals were determined as described\(^\text{29}\).
**DNA substrates**

To construct the DNA substrates for helicase and nuclease assays, as diagramed in Extended Data Table 1c, selected oligonucleotides were first either 5′ end-labeled with \(^{\gamma-32P}\) ATP using T4 polynucleotide kinase (New England Biolab) or 3′ end-labeled with \(^{\alpha-32P}\) Cordycepin 5′-triphosphate (PerkinElmer) and terminal deoxytransferase (Roche). The end-labeled oligonucleotides were then annealed to their partner oligonucleotides in buffer A (50 mM Tris-HCl, pH 7.5, 10 mM MgCl\(_2\), 100 mM NaCl). The hybridized DNA substrates were further purified in a 10% native polyacrylamide gel followed by electroelution at 4°C. To introduce a nick at the rNMP site, the DNA duplex that contains a unique rUMP hybrid (50 pmol) was incubated with (60 units) calf thymus Topo I (Invitrogen) in the reaction buffer (50 mM Tris-HCl, pH 8.0, 50 mM KCl, 1 mM DTT) for 30 min at 37°C. After an incubation with SDS (0.2%) and proteinase K (0.5 mg/ml) for 10 min at 37°C, the DNA substrate was purified using a PCR cleaning kit (Qiagen).

**Nuclease and helicase assays**

For nuclease assays, the DNA substrate (5 nM) was incubated with the indicated amount of Exo1 or hEXO1 in the reaction buffer (20 mM Tris-HCl, pH 7.5, 2 mM MgCl\(_2\), 1 mM DTT, 200 ng / mL BSA, 50 mM KCl) for 10 min at 30°C or 37°C respectively. The reactions were fractionated in an 18% denaturing polyacrylamide gel followed by phosphorimaging analysis of the dried gel.

For helicase assays, the DNA substrate (5 nM) was incubated with the indicated amount of Srs2 or Mph1 in the reaction buffer (20 mM Tris-HCl, pH 7.5, 2 mM MgCl\(_2\), 1 mM DTT, 200 ng / mL BSA, 50 mM KCl) with 1 mM ATP and the ATP regenerating system for 10 min at 30°C. The reactions were fractionated in a 12% native polyacrylamide gel followed by phosphorimaging analysis of the dried gel.
Extended Data

Extended Data Figure 1. Genetic interactions of the RNH201/202/203 genes and SRS2
Lists of genes were obtained from the Saccharomyces Genome Database. Genes in grey are synthetically sick or synthetically lethal with either RNH201/RNH202/RNH203 or with SRS2. Genes in black are synthetically sick/lethal with both RNH201/RNH202/RNH203 and SRS2.

Extended Data Figure 2. Full-length srs2-KA displays the same double mutant phenotype with rnh202Δ as do srs2Δ and srs2-KA860
a. Strains harboring the helicase null K41A variant at the chromosomal SRS2 locus were constructed. b. Synthetic interaction of rnh202Δ and srs2-KA. c. Dinucleotide slippage rates; plotted are median rates with error bars representing 95% confidence intervals; n=18.

Extended Data Figure 3. Increased DNA damage load in the rnh202Δ srs2Δ double mutant
a. Left, percentage cells with spontaneous Rad52-Yfp foci formation, with at least 10 fields of 50–200 cells counted per strain. Right, representative DIC and YFP images of a Rad52-Yfp focus. b. Log phase cells were analyzed by flow cytometry to determine DNA content and data were quantified using FlowJo software. c. Percentage of large doublet cells in mid-log phase cultures. Total number of cells and number of large doublet cells were determined for each strain by examining at least 10 fields of 50–200 cells per strain. d. Wild type and rnh202Δ srs2Δ double mutant cells were grown in YPD media, washed and treated for 1 hour with 100 mM HU. Lysates were extracted and subjected to immunoblotting with an anti-Rad53 antibody. The higher species represents phosphorylated Rad53. Plotted in (a) and (c) are the mean values±s.d, n=3.
Extended Data Figure 4. Mutation spectra at the CAN1 locus for the rnh202Δ single and rnh202Δ srs2Δ double mutant cells

Independent mutations in the rnh202Δ (in blue) and rnh202Δ srs2Δ (in red) CanR mutant strains. Base substitutions are written above the wild type sequence, deletions are represented by “=”, and insertions are represented by “^N^”.
Extended Data Figure 5. Unwinding of nicked duplex DNA by Srs2

Intact duplex DNA (a) and nicked duplex DNA labeled at either the 5′ end (b) or the 3′ end (c) of the nicked strand were incubated with Srs2 for 10 min at 30°C with or without ATP. Reactions were run in a polyacrylamide gel and analyzed by phosphorimaging analysis. The results were quantified in (d). Plotted are the mean values±s.d, n=3.
Extended Data Figure 6. Topo I-catalyzed DNA cleavage at the rUMP site
Duplex DNA with or without an embedded rUMP residue (5 nM) (a) were incubated with either E. coli or Calf thymus Topo I for 30 min at 37°C. Reactions were analyzed by electrophoresis in a polyacrylamide gel followed by phosphorimaging analysis (b). A schematic for the construction of the DNA substrate that harbors a unique nick induced by calf thymus Topo I at a rUMP residue (c).
Extended Data Figure 7. Mph1 does not act at the Topo I–induced nick

a. DNA substrate (5 nM) that harbors a Topo I-induced nick and radiolabeled on the 3′ side was incubated with Srs2 or Mph1 (20 or 40 nM) in the presence or absence of ATP at 30°C for 10 min. The reactions were analyzed as before.

b. (AG)$_4$ dinucleotide slippage rates; plotted are median rates with error bars representing 95% confidence intervals; n=18

c. Exo1 displays 5′ endo- and exo- nuclease activities on the 5′-FLAP and the nicked duplex DNA, respectively. Radiolabeled 5′-FLAP without or with a gap region, nicked duplex DNA, or oligo dT (5 nM each) was incubated with Exo1 for 10 min at 30°C. The reactions were analyzed as before and the products indicated by the arrows.
Extended Data Figure 8. Stimulation of Exo1-catalyzed digestion of a 5′-FLAP and a Topo I-induced nick by Srs2

a. Radiolabeled 5′-FLAP substrate (5 nM) was incubated with Exo1 (0.1 nM) in the absence or presence of Mph1 for 10 min at 30°C. b. The radiolabeled 5′-FLAP substrate (5 nM) was incubated with hEXO1 (4 nM) in the absence or presence of Srs2 (0.4, 2 and 10 nM) for 10 min at 37°C. c. Radiolabeled DNA substrate harboring a Topo I-induced nick (5 nM) was incubated with Exo1 (0.1 nM) in the absence or presence of Srs2 or Mph1 for 10 min at 30°C. d. The radiolabeled substrate harboring a Topo I-induced nick (5 nM) was incubated with increasing amounts of hEXO1 (0.25, 0.5 and 1 nM) in the absence or presence of Srs2 (10 nM) for 10 min at 37°C. The reactions in (a), (b), (c) and (d) were analyzed as before. Plotted in (b) and (d) are the mean values±s.d, n=3.
Extended Data Figure 9. Model of Srs2/Exo1-mediated processing of a DNA nick induced by Top1 at an embedded ribonucleotide

RNase H2 normally removes misincorporated rNMPs from DNA (not shown). In the absence of RNase H2, topoisomerase 1 can cleave at the embedded ribonucleotide, leaving 3′ cyclic phosphate and 5′ OH ends that cannot be ligated. Repair commences with unwinding of DNA from the 5′ OH side by Srs2, followed by DNA digestion by Exo1 via its exonuclease or 5′ FLAP endonuclease activity to result in a DNA gap, to be filled by a DNA polymerase. Our results show that Srs2 enhances the nuclease activities of Exo1. In the absence of Srs2, religation of ends processed by unknown nuclease activities leads to a deletion mutation.
Extended Data Table 1

a. Can^r mutation spectra. b. Yeast Strains c. Sequences of oligo nucleotides used.

|           | Total | Base change | 1 nt repeat | b2 nt repeat |
|-----------|-------|-------------|-------------|-------------|
| WT        | 52    | 52          | 0           | 0           |
| rnh20Δ    | 105   | 41          | 33          | 32          |
| srs2Δ     | 61    | 48          | 8           | 5           |
| rnh20Δ srs2Δ | 102   | 16          | 27          | 59          |
| rnh20Δ top1Δ | 54    | 40          | 14          | 0           |
| srs2Δ top1Δ | 50    | 33          | 17          | 0           |

b

| Strain | Yeast Strain | Reference |
|--------|--------------|-----------|
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Figure 1. Genome stability maintenance role of Srs2 helicase in rnh202Δ cells

a. Strains harboring either the srs2-860 allele or its helicase null K41A variant at the chromosomal SRS2 locus were constructed. b. Doubling times. c. Spontaneous mutation rates. d. Recombination rates. e. Sensitivity to hydroxyurea (HU). f. Synthetic interaction of the pol2M644G mutation in the rnh202Δ srs2Δ background. g. Mutation spectra of CanR mutants obtained in (c). h. Dinucleotide slippage rates. Plotted in (b) are the mean values ±s.d., n=3. Plotted in (c), (d) and (h) are median rates with error bars representing 95% confidence intervals; n=18.
Figure 2. Causation of genome instability in \textit{rnh202\Delta srs2\Delta} mutant by Top1
\textbf{a.} Spontaneous mutation rates. \textbf{b.} Mutation spectra of Can\textsuperscript{r} mutants obtained in (a). \textbf{c.} Sensitivity to hydroxyurea (HU). \textbf{d.} Recombination rates. \textbf{e.} Synthetic interaction of the pol2M644G mutation in the \textit{rnh202\Delta srs2\Delta top1\Delta} background. Plotted in (a) and (d) are median rates with error bars representing 95% confidence intervals\textsuperscript{29}; n=18.
Figure 3. a. Srs2-mediated displacement of the 5′-strand at a Topo I-induced DNA nick
DNA substrates containing a Topo I-induced nick and labeled at either the 3′ or 5′ end were incubated with Srs2 with or without ATP. The reactions were analyzed on a polyacrylamide gel (i); the results were quantified (ii). Heat-denatured (HD) DNA was analyzed in lanes 2 and 7. b. Effects of Exo1 deletion on slippage mutations in *rnh202Δ* and *rnh202Δ srs2Δ* mutant cells. Dinucleotide slippage rates. Plotted in (a) are the mean values±s.d., n=3; in (b), median rates with error bars representing 95% confidence intervals29; n=18.
Figure 4. Interactions of Exo1 with Srs2

a. Endonucleolytic cleavage of a gapped 5′-FLAP DNA by Exo1 (0.1 nM) was examined with and without Srs2 or ATP. The reactions were analyzed on a polyacrylamide gel (i) and the results were quantified (ii). b. Affinity pulldown through the (His)$_9$ tag on Srs2 was used to analyze its interaction with Exo1. The supernatant (S), wash (W) and SDS eluate (E) fractions were analyzed by 4–20% SDS-PAGE. c. Exonucleolytic cleavage of a (i) Topo I-induced nick, (ii) nick bearing a 5′-OH group, and (iii) clean nick bearing a 5′-phosphate by Exo1 was examined with and without Srs2 or ATP. The reactions were analyzed as above (iv). Plotted in (a) and (c) are the mean values±s.d., n=3.