RECCQ-like helicases Sgs1 and BLM regulate R-loop–associated genome instability

Emily Yun-Chia Chang,1* Carolina A. Novoa,1* Maria J. Aristizabal,2,6 Yan Coulombe,3,4 Romulo Segovia,1 Richa Chaturvedi,3,4 Yaoqing Shen,5 Christelle Keong,1 Annie S. Tam,1,6 Steven J.M. Jones,5,6 Jean-Yves Masson,3,4 Michael S. Kobor,2,6 and Peter C. Stirling1,6

1Terry Fox Laboratory, British Columbia Cancer Agency, Vancouver, Canada
2Centre for Molecular Medicine and Therapeutics, Vancouver, Canada
3Genome Stability Laboratory, Centre Hospitalier Universitaire de Québec Research Center, Quebec City, Canada
4Department of Molecular Biology, Medical Biochemistry and Pathology, Laval University Cancer Research Center, Quebec City, Canada
5Michael Smith Genome Sciences Centre, Vancouver, Canada
6Department of Medical Genetics, University of British Columbia, Vancouver, Canada

Sgs1, the orthologue of human Bloom’s syndrome helicase BLM, is a yeast DNA helicase functioning in DNA replication and repair. We show that Sgs1 loss increases R-loop accumulation and sensitizes cells to transcription–replication collisions. Yeast lacking SGS1 accumulate R-loops and γ-H2A at sites of Sgs1 binding, replication pausing regions, and long genes. The mutation signature of sgs1Δ reveals copy number changes flanked by repetitive regions with high R-loop–forming potential. Analysis of BLM in Bloom’s syndrome fibroblasts or by depletion of BLM from human cancer cells confirms a role for Sgs1/BLM in suppressing R-loop–associated genome instability across species. In support of a potential direct effect, BLM is found physically proximal to DNA:RNA hybrids in human cells, and can efficiently unwind R-loops in vitro. Together, our data describe a conserved role for Sgs1/BLM in R-loop suppression and support an increasingly broad view of DNA repair and replication fork stabilizing proteins as modulators of R-loop–mediated genome instability.

Introduction

Genome instability is an enabling characteristic of tumor formation because it creates mutational diversity in premalignant cell populations, allowing the necessary mutations in driver genes to occur at a sufficiently high frequency (Stratton et al., 2009). One of the best-understood ways in which genome instability arises in cancer is an acquired defect in a DNA repair pathway. Mutations in homologous recombination (HR), nucleotide excision repair, cross-link repair, and mismatch repair are all clearly linked to an increased risk of cancer (Curtin, 2012). Germline mutations in genes coding for DNA repair proteins dramatically increase cancer risk and can be associated with other symptoms, whereas somatically acquired cancer driver mutations in genes that function in the same repair pathways are found in sporadic cancers.

One such DNA repair protein is the RECCQ-like helicase, BLM, which is involved in resolution of concatenated DNA molecules during HR, at replication forks, and in anaphase (Böhm and Bernstein, 2014). BLM can act on a wide array of substrates in vitro including Holliday junctions, D-loops, G-quadruplexes, DNA:RNA hybrids, and single-stranded overhangs (Popuri et al., 2008; Croteau et al., 2014). Germline BLM mutations lead to Bloom’s syndrome, which is characterized by cancer predisposition, short stature, and other symptoms (de Renty and Ellis, 2017). At the cellular level, defects in BLM are characterized by high levels of sister chromatid exchanges, and DNA replication and mitotic defects (Böhm and Bernstein, 2014). BLM is highly conserved across evolution, and much of what is known about its function was first described for its orthologue in Saccharomyces cerevisiae, Sgs1. Research on Sgs1 has linked its activity to HR, both at the stage of end-resection and double-Holliday junction resolution, restart of stalled DNA replication forks, meiosis, and telomere maintenance (Ashton and Hickson, 2010). Both BLM and Sgs1 have multiple interacting partners that regulate their activity and cooperate with them in catalyzing DNA transactions. BLM/Sgs1 forms a complex with Topoisomerase III and RMI1/2, which work together with BLM to decatenate DNA molecules. The BLM-Top3-Rmi1/2 complex further associates with members of the Fanconi anemia
Recently, defects in DNA repair proteins have been linked to a novel mechanism of genome instability involving the formation of excessive DNA:RNA hybrids on genomic DNA. These hybrids form a structure called an R-loop, in which RNA binds to a complementary DNA strand and exposes the non-template strand as a single-stranded DNA loop (Chan et al., 2014b). R-loops are thought to cause genome instability primarily by interfering with DNA replication. R-loop collision with replication forks leads to fork stalling and an increase in double-strand breaks or error-prone mechanisms of replication (Chan et al., 2014b). The best understood players in R-loop metabolism are those involved in RNA processing, in which normal transcript elongation, termination, splicing, packaging, nuclear export, and RNA degradation have all been shown to suppress R-loop formation (Li and Manley, 2005; Gómez-González et al., 2011; Mischo et al., 2011; Wahba et al., 2011; Stirling et al., 2012). Among these, RNaseH’s have the most direct effects, with RNaseH1 working only on R-loop substrates, and RNaseH2 targeting R-loops and ribonucleotides incorporated into DNA (Cerritelli and Crouch, 2009). The THO complex, made up of Hpr1, Mft1, Tho2, and Tph2 in yeast, is an mRNA export complex whose disruption has been repeatedly associated with R-loop-mediated genome instability (Chávez et al., 2000; Gómez-González et al., 2011). Finally, Sen1, the yeast homologue of senataxin, is a DNA/RNA helicase with multiple functions, including coordination of replication and transcription, likely exploiting a direct R-loop unwinding activity (Mischo et al., 2011).

Interestingly, defects in canonical DNA repair proteins such as HR factors BRC1 and BRC2 (Wahba et al., 2013; Bhatia et al., 2014; Hatchi et al., 2015), nucleotide excision repair proteins XPG and XPF (Sollier et al., 2014), the FA pathway (García-Rubio et al., 2015; Schwab et al., 2015), and the DNA damage response kinase ATM (Tresini et al., 2015) have all been associated with stabilization or signaling involving R-loops. Moreover, R-loops have been shown to contribute to DNA replication stress in these DNA repair mutants, and in some cases, a direct role for the repair protein in R-loop removal has been suggested (e.g., R-loop displacement by the FANC-M helicase; Schwab et al., 2015). Indeed, the BLM protein is known to cooperate with the HR pathway and is critical for the activation of the FA pathway. Moreover, Sgs1 has synthetic phenotypes with RNaseH2 deletions in yeast, suggesting a functional cooperation between the two proteins (Kim and Jinks-Robertson, 2011; Chon et al., 2013).

Here we show that in yeast deleted for SGS1, genome instability is partially transcription-dependent, and that both R-loops and DNA damage accumulate at sites of Sgs1 action in the genome. By physically mapping Sgs1 binding sites in the yeast genome, we observe a strong association between Sgs1 binding and sites that gain R-loops and DNA damage in sgs1Δ yeast. Indeed, unbiased mutation accumulation in sgs1Δ cells identifies increases in structural rearrangements at R-loop–prone sites. Finally, we confirm R-loop–associated genome instability in BLM-depleted human cells and in Bloom’s syndrome fibroblasts, further showing that BLM localizes near R-loops in cells and is capable of resolving R-loops as efficiently as D-loops in vitro. Together these data establish Sgs1/BLM as a regulator of R-loop–coupled genome instability, adding to the growing repertoire of DNA repair proteins with functions in R-loop mitigation, and extending the notion that transcription–replication collisions are one of the drivers of mutagenesis in cancer.

**Results**

**R-loop formation and consequences in sgs1Δ**

Yeast Sgs1 has been ascribed a downstream or cooperative role with RNaseH2 in suppressing genome instability, presumed to be related to shared roles in DNA replication and potential cooperation at sites of transcription-mediated instability (Kim and Jinks-Robertson, 2011; Chon et al., 2013). However, based on recent studies linking replication fork–associated DNA repair proteins to R-loop suppression, we hypothesized that Sgs1 may be involved in mitigating R-loop–associated genome instability. To directly assess if deletion of SGS1 alters the levels of DNA:RNA hybrids, we used chromosome spreads and quantified staining with the S9.6 mAb, which recognizes DNA :RNA hybrids in a manner largely independent of sequence (Hu et al., 2006). This analysis showed that SGS1 deletion increases S9.6 staining compared with WT (Fig. 1A). Ectopic R-loops are known to drive transcription–replication conflicts leading to recombination. To determine the effect of Sgs1 on this process, we tested recombination rates in a plasmid system in which transcription is driven by a promoter that is oriented to be colliding (IN) or codirectional (OUT) with the origin of replication (Fig. 1B). This analysis showed that sgs1Δ-driven hyperrecombination was significantly enhanced when S-phase Histone H4 promoter ([HHF]) transcription is IN with DNA replication but was unaffected in the OUT orientation or by controls in which transcription occurs in G1 (CLB-IN) or G2 (BLB-IN) phase (Fig. 1B). Transcription–recombination conflicts lead to DNA damage, and we found that, whereas deletion of SGS1 leads to a small increase in levels of DNA damage as measured by Rad52-YFP foci, combined loss of RNaseH2A (i.e., sgs1Δrnh201Δ) leads to a synergistic increase in DNA damage (Fig. 1C; Chon et al., 2013). Importantly, this enhanced damage effect was suppressed by overexpression of RNaseH1. Because RNaseH1 only degrades RNA in DNA:RNA hybrids, as opposed to functioning in replication or ribonucleotide excision repair, the synergistic DNA damage in sgs1Δrnh201Δ is a result of R-loops as opposed to the effects of Rnh201 or Sgs1 on replication. Cells lacking SGS1 exhibit hyperrecombination in scenarios independent of R-loops based on Sgs1’s well-established role in HR and replication fork protection. Thus, we propose that a subset of genome instability events in sgs1Δ could be R-loop related.

**Loss of SGS1 exhibits synergistic genome instability with R-loop suppressors**

We next sought to further probe the role for R-loop modulators in sgs1Δ phenotypes. We first assessed the fitness of sgs1Δ in combination with deletions of R-loop regulators in the THO complex (MFT1), RNaseH enzymes (RNH1 and RNH201), and senataxin (SEN1) and observed synergistic fitness defects (Huertas and Aguilera, 2003; Gómez-González et al., 2009; Mischo et al., 2011; Skourtis-Stathaki et al., 2011). Loss of Sgs1 significantly exacerbated fitness defects in rnh201Δ, sen1-1, and mft1Δ cells (Fig. 2A). Similar synergy was seen when measuring chromosome instability using the A-like Faker (ALF) assay for disruption of the MAT locus in chromosome...
III (Fig. 2 B) and a LEU2 plasmid-based direct repeat recombination (Fig. 2 C; Stirling et al., 2011). Double mutants of SGS1 and the THO complex subunit MFT1 caused a dramatic, greater than additive, increase in instability in these assays compared with the single mutants, suggesting a synergistic effect (Fig. 2, B and C). Deletion of known R-loop suppressors with diverse modes of action (i.e., THO complex, Sen1, and RNaseH) all exhibited synergistic chromosome instability phenotypes when combined with sgs1Δ (Fig. S1). These synergies are not surprising because loss of SGS1 is known to promote hyperrecombination through loss of its role in resolution of Holliday junctions (Ashton and Hickson, 2010), although this phenotype would not explain the increases in R-loop staining we see in sgs1Δ (Fig. 1).

R-loop levels increase with transcript frequency and length in direct repeat recombination assays. Therefore, to implicate Sgs1 further in transcription-associated recombination, we assessed the roles of transcript length and frequency with derivatives of the LEU2 direct repeat systems and compared to mft1Δ as a control. Although sgs1Δ had higher recombination frequencies in both assays, comparing the rates of recombination for the indicated strain and plasmid. Fold increases over WT +HHF-out are shown above each bar. 11 or more independent frequencies were measured for each sample. (C) DNA damage synergy in sgs1Δmft1Δ is R-loop-dependent. Quantification of Rad52-YFP foci in the indicated strain with either an empty vector (−) or a Gal-inducible RNH1 (+) construct (n = 3). Error bars are SEM. **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.

Sgs1 binds genomic sites that show increased R-loops and DNA damage upon deletion of SGS1

Given the increase in DNA:RNA hybrid levels in the sgs1Δ mutant and the observed relationship between SGS1 and transcription-associated recombination, we hypothesized that Sgs1 might be recruited to sites of transcription and could impact the R-loop and DNA damage landscape at these sites. To test this, we mapped Sgs1 binding by ChIP-chip, and R-loops and γ-H2A levels in sgs1Δ by DRIP-chip and ChIP-chip (Stirling et al., 2012; Chan et al., 2014a). Sgs1 does bind to ORFs, with a preference for longer and more highly transcribed genes (Fig. 3, A and D; and Fig. S2 A; Sgs1-containing genes had a mean length of 1,586 bp, compared with 1,338 bp for all genes; P < 0.0001). Sgs1 also associated at sites bound by the Rrm3 helicase (Fig. 3 G), which functions at stalled replication forks to promote replisome progression, thus demarcating replication obstacles in the yeast genome (Santos-Pereira et al., 2013). Further underscoring a shared role, previous work
showed that RRM3 is required for robust growth in sgs1Δ cells (Schmidt and Kolodner, 2004). Finally, we observed some but very sparse binding of Sgs1 along telomeres and the ribosomal DNA (rDNA) loci (Fig. S2, B and C). The relatively low levels of Sgs1 association at the rDNA was surprising given Sgs1’s known role in promoting both replication and transcription of the rDNA loci. Given the surprising occupancy of Sgs1 at ORFs, and in particular long genes, we next focused on potential effects of SGS1 deletion on DNA:RNA hybrid and γ-H2A profiles. Consistent with the increase in S9.6 staining and Rad52 foci observed in the sgs1Δ mutant, we also found that loss of SGS1 resulted in increased DNA:RNA hybrids and γ-H2A at a subset of genomic loci. More specifically, we found that loss of SGS1 increased DNA:RNA hybrid and γ-H2A levels at longer genes (Fig. 3, B, C, E, and F), an effect corroborated by our observation of increased recombination at longer versus shorter reporter genes in the sgs1Δ mutant (Fig. 2 D). Analysis of the genes significantly occupied by DNA:RNA hybrids or γ-H2A signal in both replicates of sgs1Δ but not WT confirmed a significant shift toward longer than average genes (i.e., sgs1Δ DNA:RNA hybrid containing genes were 1,745 bp and γ-H2A containing genes were 1,540 bp compared with 1,338 bp for all genes; P < 0.0001 ANOVA with Holm-Sidak correction). The distribution of genes with increased DRIP and γ-H2A signal in sgs1Δ also showed a small bias to subtelomeric regions (Fig. S2 B).

Loss of Sgs1 also increased DNA:RNA hybrid and γ-H2A levels at other sites, namely, regions bound by Rrn3 (Fig. 3 G; DNA replication slow zones), an observation consistent with Sgs1 functioning at replication obstacles (Cobb et al., 2003), and with loss of Sgs1 leading to R-loop stabilization and DNA damage at these sites. Interestingly, our profiles also revealed sites where loss of SGS1 increases DNA:RNA hybrid levels without concomitant effects on γ-H2A: the rDNA loci and a subset of telomeres (Fig. S2). However, we observed higher γ-H2A signal flanking the rDNA, and show that increased rDNA instability in the sgs1Δ mutant could be suppressed to WT levels by ectopic expression of RNaseH1 (Fig. S2 D), suggesting that although these DNA:RNA hybrids were not associated with increased damage at the rDNA loci, as measured by γ-H2A occupancy, they may still contribute to instability.

Figure 2. Cooperative genome maintenance by R-loop regulators and Sgs1. [A] Growth defects in strains lacking Sgs1 and the indicated gene. Quantitative growth curve analysis (n = 4 with technical triplicates in each experiment; error bars are SEM) showed observed fitness values were significantly lower than expected values. (B and C) Synergistic genome instability in SGS1, MFT1 double deletions. (B) Mating frequency as a measure of chromosome instability by the ALF assay. (C) Recombination frequency in the direct repeat plasmid system LNA (Prado et al., 1997). Fold increase over WT is shown above each measurement in the indicated strains. Error bars are SEM. (D and E) Plasmid-based LEU2 recombination frequency as a measure of transcript length (D) and frequency (E). Above each panel is a schematic of the assay that measures recombination frequency, transcript length dependence, and transcript frequency dependence. For D, the length of the intervening sequence between the leu2 repeats [dark bars] is shown at right. The fold increase over WT is shown above each bar. The direction of transcription is indicated by an arrow. For E, Dex, dextrose [low expression]; Gal, galactose [high expression]. For D, n ≥ 14 and for E, n ≥ 6 independent frequencies were measured. Boxplots in D and E plot whiskers to the maximum and minimum values, with the box between the 25th and 75th percentile and the line at the median value.
Normal Sgs1 binding sites define R-loop prone fragile sites in sgs1Δ cells

To probe the interdependence of the genome-wide profiles generated, we focused on Sgs1-bound regions and analyzed DNA:RNA hybrid and γ-H2A levels at these sites. Overall, we found that sites of Sgs1 binding were associated with DNA:RNA hybrids and γ-H2A, the levels of which increased when SGS1 was deleted (Fig. 4 A). Focusing on ORFs, we found that increasing levels of Sgs1 binding signal were associated with increased levels of DNA:RNA hybrids and γ-H2A in the sgs1Δ mutant compared with WT (Fig. 4 B). This trend was stronger for γ-H2A than DNA:RNA hybrids, suggesting that Sgs1 functioned directly to prevent DNA damage at a subset of ORFs, although this was not always through a role in preventing DNA:RNA hybrid accumulation at those sites. To probe this relationship further, we divided ORFs into Sgs1-bound and not-bound groups. Consistent with our observation that Sgs1-bound peaks defined sites of R-loop and γ-H2A occupancy in WT cells (Fig. 4 A), we found that under WT conditions, Sgs1-bound ORFs had higher levels of DNA:RNA hybrids.
and γ-H2A compared with ORFs that were not bound by Sgs1 (Fig. 4, C and D). Importantly, we found that the levels of DNA:RNA hybrids and γ-H2A significantly increased upon deletion of SGS1 only for the group of Sgs1-bound genes, suggesting that the direct association of Sgs1 with these ORFs functioned to mitigate the levels of DNA:RNA hybrids and γ-H2A. To take this analysis a step further, we focused only on ORFs that both were bound by Sgs1 and gained DNA:RNA hybrids in the sgs1Δ mutant compared with WT, and found that upon deletion of SGS1, these genes had a greater increase in γ-H2A levels than genes that were bound by Sgs1 but did not accumulate hybrids upon its loss (Fig. 4 E). Interestingly, this analysis also revealed a set of 155 ORFs that gained DNA:RNA hybrids when SGS1 was deleted but did not pass our binding threshold for Sgs1 binding. Consistent with these sites representing indirect effects of Sgs1 on the DNA:RNA hybrid landscape, they did not show increased γ-H2A levels compared with the rest of ORFs that were not bound by Sgs1 (Fig. 4 E). Together these results suggest that Sgs1 normally binds to fragile, R-loop–prone regions, and that loss of SGS1 activates a subset of these sites to accumulate R-loops and DNA damage.

Genome instability in sgs1Δ yeast occurs at R-loop–prone regions

Our data suggest that Sgs1 reduces R-loops and DNA damage at specific genomic loci. To create an unbiased view of ongoing instability in the genome of sgs1Δ cells, we performed a mutation accumulation and whole-genome sequencing experiment (Stirling et al., 2014). Passaging homozygous sgs1Δ/Δ diploids for ~1,000 generations created a set of 12 mutation accumulation strains that we sequenced at >50× coverage (Fig. 5 A). This analysis revealed a modest approximately twofold increase in single-nucleotide variants (SNVs) for sgs1Δ/Δ compared with WT, which was similar to the rates seen at the CAN1 reporter.

Figure 4. Intersection of Sgs1, R-loops and DNA damage. All profiles were generated in duplicate with quantile normalized and averaged data shown here. (A) Mean DNA:RNA hybrid (top) and γ-H2A (bottom) occupancy across Sgs1-binding sites for the indicated strains. Two-sided Wilcoxon test p-value comparing mean occupancy scores in sgs1Δ versus WT is noted below. (B) Chromatrap plots showing a heat map of the difference in DNA:RNA hybrid (left) and γ-H2A (right) occupancy in sgs1Δ compared with WT. Protein coding genes are sorted by length and Sgs1 occupancy and aligned at the TSS (Hentrich et al., 2012). (C and D) Box plots comparing mean occupancy scores of DNA:RNA hybrids (C) and γ-H2A (D) in WT and the sgs1Δ mutant at Sgs1-bound versus not bound ORFs. *, P < 0.05, for two-tailed Wilcoxon tests. The numbers of genes in each category are shown in brackets. All other comparisons are not significant. (E) Box plot showing increases in γ-H2A levels at genes that are bound by Sgs1 and that gain DNA:RNA hybrids upon loss of SGS1 compared with those that do not gain hybrids. *, P < 0.05, for one-tailed Wilcoxon test. The numbers of genes in each category are shown in brackets. Changes in γ-H2A were not observed when genes that gained hybrids but were not bound to Sgs1 were compared with those that have neither hybrids nor Sgs1 association. The whiskers on boxplots in C–E represent 1.5× the interquartile range.
locus (Segovia et al., 2017). However, sgs1Δ/Δ exhibited a ~12× increase in copy number variants (CNVs) compared with WT (Fig. 5 B). The majority of these changes were segmental, although aneuploidy also increased in sgs1Δ/Δ (Fig. 5 C). Analysis of the predicted breakpoints shows that most originate within or near a TY retrotransposon element (Fig. 5 D and Table S2). Indeed, Sgs1 has been ascribed a role in Ty1 element expansion based on its role in HR (Bryk et al., 2001). Other breakpoints appear to be at telomeres, and at a set of protein coding genes. Breakpoint-associated genes (Table S2) often have paralogs in the yeast genome, and this, along with the TY element enrichment, is consistent with the role for Sgs1 in rejecting HR reactions that could lead to CNVs between repetitive sequences (Myung et al., 2001). Because TY elements and telomeres are known to be hot spots of R-loop formation, we explored the potential correlation of CNV breakpoints within protein coding genes and R-loop occupancy. The R-loop signal was significantly higher than the mean background of WT DRIP peaks (10.2 [n = 19] vs. 4.1 [n = 14,307]; Mann–Whitney test, P < 0.0001; Chan et al., 2014a). Interestingly, as for Sgs1–bound and DRIP- and γ-H2A–associated genes, analysis of genes associated with CNV breakpoints within protein coding genes and R-loop occupancy. The R-loop signal was significantly higher than the mean background of WT DRIP peaks (10.2 [n = 19] vs. 4.1 [n = 14,307]; Mann–Whitney test, P < 0.0001; Chan et al., 2014a). Interestingly, as for Sgs1–bound and DRIP- and γ-H2A–associated genes, analysis of genes associated with CNV breakpoints in sgs1Δ/Δ were significantly longer than the genome mean (1.745 bp compared with 1.338 bp, Mann–Whitney P = 0.0425). Overall, the mutation signature of sgs1Δ/Δ cells supports its known specific role in promoting noncrossover events and rejecting HR (Myung et al., 2001).Because TY elements and telomeres are known to be hot spots of R-loop formation, we explored the potential correlation of CNV breakpoints within protein coding genes and R-loop occupancy. The R-loop signal was significantly higher than the mean background of WT DRIP peaks (10.2 [n = 19] vs. 4.1 [n = 14,307]; Mann–Whitney test, P < 0.0001; Chan et al., 2014a). Interestingly, as for Sgs1–bound and DRIP- and γ-H2A–associated genes, analysis of genes associated with CNV breakpoints in sgs1Δ/Δ were significantly longer than the genome mean (1.745 bp compared with 1.338 bp, Mann–Whitney P = 0.0425). Overall, the mutation signature of sgs1Δ/Δ cells supports its known specific role in promoting noncrossover events and rejecting HR (Myung et al., 2001). In addition, it is consistent with the observed sensitivity of sgs1Δ/Δ cells to transcription-associated recombination and the accumulation of DNA:RNA hybrids and DNA damage in sgs1Δ/Δ cells.

R-loops accumulate and cause DNA damage in BLM-depleted cell lines

The human orthologue of Sgs1 is the Bloom’s syndrome helicase BLM, one of five RECQ-like helicases in humans and the closest sequence orthologue. To determine whether a role of Sgs1 in R-loop metabolism is conserved in mammalian cells, we used siRNA to target BLM in HeLa cells (Fig. S3 A), and measured R-loop levels by immunofluorescence. Knockdown of BLM leads to a significant increase in S9.6 staining, which could be abolished by overexpression of GFP-RNaseH1 (Fig. 6, A and B). Similar staining results were obtained for BLM−/− knockout derivatives of the near diploid HCT116 cell line compared with an isogenic control line (Fig. S3 C). Importantly, this accumulation was also seen in Bloom’s syndrome fibroblasts relative to an isogenic control complemented with WT BLM (Fig. 6, C and D). Moreover, the reduction of R-loop levels was not seen in fibroblasts complemented with the BLM K695T mutant, which abolishes its helicase activity (Fig. 6, C and D; Neff et al., 1999), suggesting a direct role for the BLM helicase activity in removing R-loops.

Like Sgs1 in yeast, BLM depletion increases genome instability phenotypes, and we therefore examined the effects of RNaseH1 overexpression in BLM knockdown cells on genome instability. BLM knockdown caused chromosome instability as measured by increased micronucleus formation after 48 h (Fig. S3, D and E), likely as a result of an inability to resolve anaphase bridges at mitosis (Naim and Rosselli, 2009). Importantly, the increase in micronuclei formation was significantly suppressed by overexpression of GFP-RNaseH1 (Fig. S3, D and E). BLM knockdown also induced DNA breaks as measured by the neutral comet assay and γ-H2AX focus accumulation, with both defects significantly reduced by ectopic expression of GFP-RNaseH1 (Fig. 6, E and F). Indeed, Bloom’s syndrome fibroblasts also showed high levels of γ-H2AX foci, which could be reduced by RNaseH1 expression (Fig. 6 G), whereas fibroblasts complemented with WT BLM showed lower levels of γ-H2AX foci that were not reduced by RNaseH1 expression (Fig. 6 G). These data suggest that a considerable proportion of DNA damage and genome instability in BLM-deficient mammalian cells, and potentially in Bloom’s syndrome, may be a result of a reduced ability of these cells to process R-loops.
Figure 6. R-loop accumulation and DNA damage in BLM-depleted cells. (A and C) Representative images of S9.6 staining in HeLa cells treated with the indicated siRNA targeted for BLM (si-BLM) or a control luciferase (si-Luc; A) or Bloom's syndrome fibroblasts complemented with an empty vector control (BSF), WT BLM (BSF + WT), or helicase dead mutant (BSF + HM; C). Cells were transfected with either a control vector (GFP) or one expressing GFP-RNaseH1. (B and D) Quantification of S9.6 signal intensity for nuclear area in the indicated conditions in HeLa cells (B) or Bloom's syndrome fibroblasts (D). Cell numbers scored across three independent replicates are noted below panel B for HeLa cells. (E) RNaseH-dependent DNA breaks in BLM-deficient HeLa cells. (Left) Representative comet tail images from single-cell electrophoresis. (Right) Quantification of comet tail moment under the indicated conditions. t tests were used for comparisons shown. (F and G) Percentages of cells with ≥10 γH2AX foci in HeLa cells treated with indicated siRNA expressing GFP or GFP-RNaseH1 (F) or Bloom's syndrome fibroblasts expressing GFP or GFP-RNaseH1 (G). *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001.
Assessing potential direct effects of BLM on R-loops

There are several possible mechanisms by which Sgs1 and BLM could impact R-loop–mediated genome instability, including effects on replisome stability at transcription–replication conflicts, or through direct unwinding of R-loops, alone or collaboratively with topoisomerase III or the FA pathway. To support these direct models, and rule out potential indirect effects, for example, through effects on DNA damage signaling pathways or transcriptional effects (Grierson et al., 2012; Tresini et al., 2015), we tested the helicase activity of recombinant BLM and Sgs1 directly on both R- and D-loop substrates. Although BLM has previously been shown to unwind R-loops in vitro (Popuri et al., 2008), how this activity compares to its ability to unwind other structures is unclear, and to our knowledge, whether Sgs1 can unwind R-loops is unknown. We found that either BLM or Sgs1 could unwind R- and D-loop substrates of the same sequence composition with nearly identical efficiencies and that this occurred in an ATP- and concentration-dependent manner (Fig. 7, A and B). Thus, both helicases are highly efficient R-loop resolvases. To determine whether BLM and R-loops are ever proximal in human cells, we performed a proximity ligation assay (PLA) with antibodies targeting BLM and DNA:RNA hybrids. Remarkably, BLM showed a clear and reproducible PLA signal in cells with S9.6 that was significantly higher than in single primary antibody controls, thus showing that BLM comes in close proximity to DNA:RNA hybrids in cells (Fig. 7 C and Fig. S4 A). BLM has many interaction partners and interdependent relationships in DNA damage repair (Suhasini and Brosh, 2012; Chaudhury et al., 2013; Ling et al., 2016). Given the previously reported physical and functional interactions of BLM with the FA pathway, which itself has been implicated in R-loop suppression (García-Rubio et al., 2015; Schwab et al., 2015) we tested potential synergy or epistasis with FA pathway components by siRNA knockdown. Knockdown of FANC D2 or FANC M increased DNA:RNA hybrid staining, but this was epistatic to coincident knockdown of BLM (Fig. 7, D and E). Furthermore, BLM was required for hydroxyurea (HU)-induced increases in FANC D2 foci (Fig. S4, B and C), supporting literature placing these factors in the same pathway during replication stress (Ling et al., 2016; Panneerselvam et al., 2016). Similar epistatic results were found when we performed knockdown of BLM together with topoisomerase III, which forms a complex with BLM to decatenate DNA and to resolve stalled replication forks (Fig. 7 F). Comparison of the effects of knockdown of other RECQ-like helicases, WRN and RECQL5, which have been separately implicated in R-loop biology, again showed S9.6 staining increases. Importantly, double knockdowns of BLM and RECQL5 were not epistatic and led to further enhanced S9.6 staining, whereas knockdown of BLM with WRN showed no additional staining compared with WRN knockdown alone (Fig. 7 G). These data support a model in which local effects of BLM at R-loops occur in the same pathway as those of FANC D2 and FANC M, but not other R-loop suppressors like RECQL5. This provides an important constraint on our model, and shows how R-loop resolution can occur through multiple independent pathways. Together, our data support a conserved mechanism for Sgs1/BLM in suppressing transcription–associated genome instability at R-loop sites (Fig. 8).

Discussion

Our data show that loss of SGS1 or BLM leads to R-loop accumulation across species and that at least some of the associated genome instability is contingent upon transcription–replication conflicts and/or R-loops. These data match with unbiased mutation accumulation analysis, which shows that loss of SGS1 increased CNVs flanked by homologous repeats and regions of high R-loop occupancy. Published mutation accumulation in rnh1Δ rnh201Δ mutants also found increased deletion and duplication events between TY elements, suggesting these could be fragile sites in R-loop–prone mutants (O’Connell et al., 2015). Global profiling of Sgs1 binding sites the in genome in concert with mapping of DNA:RNA hybrids and phospho-H2A-Serine129 in WT and sgs1Δ mutants revealed a remarkably cohesive picture. Sgs1 binding sites define regions that increase in both R-loops and DNA damage when SGS1 is deleted. These sites, many of which are at long genes, overlap with Rrm3 binding sites, indicating that they may be difficult-to-replicate sites of frequent fork stalling. Indeed, long genes have previously been implicated as a binding site for Rrm3 (Santos-Pereira et al., 2013). Moreover, Top2, an interacting partner of Sgs1 that cooperates with Sgs1 and Top3 at rDNA (Mundbjerg et al., 2015), has also been linked to transcription at long protein coding genes in yeast (Joshi et al., 2012). Thus, regions that are sensitive to topological stress may be more likely to form R-loops in sgs1Δ cells. We do not know with high resolution where damage sites occur in BLM-deficient human cells; however, BLM depletion does create ultrafine bridges and DNA damage at common fragile sites (Lukas et al., 2011). Some common fragile sites, in particular those at very long genes, have been linked to transcription–replication conflicts and R-loops (Helmrich et al., 2011), and thus for specific sites of instability in the human genome, we suggest there may be previously unappreciated links between R-loops and BLM.

Recently it was observed that, despite binding them broadly, only a subset of R-loops are degraded by Rnh1 in yeast (Zimmer and Koshland, 2016). This highlights that there may be a poorly understood regulatory distinction between normal and abnormal R-loop formation in cells, and we believe this distinction may be relevant to locus-specific differences seen in DRIP-chip profiles in mutant strains such as sgs1Δ. In the case of the rDNA, Sgs1 is known to have a role in facilitating both replication and transcription (Lee et al., 1999; Versini et al., 2003) and accordingly, we observed that enhanced rDNA instability seen in sgs1Δ cells is completely suppressed by RNaseH1 overexpression. More recently, the chromatin state of histone H3 on DNA-flanking R-loops have been cited as key determinants of whether R-loops will be DNA-damaging (Garcia-Pichardo et al., 2017), and we observed many examples of sites where R-loop occupancy and γ-H2A increase independent of one another. How the replisome and repair proteins like Sgs1/BLM influence transcription–replication conflicts in the context of chromatin is only beginning to be understood.

Possible models of Sgs1/BLM action at R-loops

Overall, we favor a model in which Sgs1/BLM works in proximity to R-loops at sites of transcription–replication conflict, consistent with studies of collaboration between Sgs1 and RNaseH2 (Kim and Jinks-Robertson, 2011; Chon et al.,
Within this local model (Fig. 8), there remain several nonmutually exclusive possibilities: the simplest model is that Sgs1/BLM unwinds R-loops directly, as supported by in vitro experiments (Fig. 7; Popuri et al., 2008). Sgs1/BLM could also unfold G-quadruplexes associated with the nontemplate strand opposite a DNA:RNA hybrid in a so-called G-loop (Duquette et al., 2004). The role of Sgs1/BLM in fork stabilization may also allow time for other factors to resolve R-loop blockages. Finally, Sgs1/BLM could direct the activity of another R-loop helicase. For example, our ChIP data link Sgs1 binding sites to those of Rrm3, and both Rrm3 and its paralog Pif1 have been recently implicated in R-loop resolution at specific loci of the yeast genome (Tran et al., 2017). In humans, BLM is known to bind FANCM and FANCI (Suhasini and Brosh, 2012), and our data suggest that the collaboration between BLM and the FA pathway is likely to be important for mitigating the effects of R-loops in human cells (García-Rubio et al., 2015; Schwab et al., 2015). For example, BLM physically and functionally interacts with FANCM (Ling et al., 2016), potentially coordinating its activity with the activity of the FA pathway, and FANCM can use its strand migration activity to remove R-loops (Schwab et al., 2015). Indeed, FANCD2 also regulates...
BLM stability and assembly at stalled replication forks, and reciprocally, FANCD2 activation requires BLM (Chaudhury et al., 2013; Panneerselvam et al., 2016). Whether these connections also extend to Sgs1 and the distantly related FA pathway of S. cerevisiae is not known. Thus, there are multiple levels of regulation to be explored across systems in the future. Indeed, the scenario may be more complex in human cells, as BLM paralogs WRN and RECQL5 have both been linked to DNA:RNA hybrid metabolism in the test tube or in cells (Chakraborty and Grosse, 2010; Saponaro et al., 2014).

Defective replisome-associated DNA repair proteins shifting the R-loop landscape

Our data add Sgs1/BLM to a growing list of DNA repair proteins that seem to work toward the error-free resolution of R-loops and the prevention of deleterious transcription–replication conflicts (Bhatia et al., 2014; Sollier et al., 2014; García-Rubio et al., 2015; Hatchi et al., 2015; Schwab et al., 2015; Chang and Stirling, 2017). In addition, there is now evidence that DNA damage alone may induce R-loops in mammalian cells (Britton et al., 2014; Tesini et al., 2015). The idea that normal robust DNA replication itself prevents R-loop accumulation is also gaining support—for example, with recent studies showing effects of the MCM helicase or POLD3 in preventing DNA:RNA hybrid metabolism (Tumini et al., 2016; Vijayaraghavan et al., 2016). The abundance of fork-protection factors emerging as R-loop regulators supports a generalized concept that the functional DNA replication machinery is an important way to mitigate deleterious transcription–replication collisions. We speculate that there are common mechanisms at play in cancers experiencing abnormal replication stress and that transcription-mediated genome instability will play a role in tumor mutation accumulation. Recent studies showing that R-loop levels vary among specific cell types in normal primary samples, and that they increase in people carrying cancer risk alleles, raises the hope that the specific role for R-loops in oncogenesis will be elucidated (Zhang et al., 2017).

Materials and methods

Yeast growth and media

Yeast were cultured according to standard conditions in the indicated media at 30°C unless otherwise indicated. Growth curves were conducted in YPD media in 96-well plates using a TECAN M200. The area under the curve was used to compute expected and observed fitness values (Stirling et al., 2011, 2012). A list of yeast strains and plasmids used in this study can be found in Table S1.

Recombination and genome instability assays

Recombination events in L, LYΔNS, LNA, pARSHLB-IN, pARS HLB-OUT, pARSLCB-IN, pARSBLB-IN, L-lacZ, and GL-lacZ systems (gifts of A. Aguilera, CABIMER, Sevilla, Spain) were scored by counting leucine-positive colonies (González-Aguilera et al., 2008; Gómez-González et al., 2009, 2011; Herrera-Moyano et al., 2014). Recombination frequencies and cell viability were obtained from the mean value of three tests performed with 3–9 independent transformants each as described (Stirling et al., 2012). In assays where yeast strains were transformed with a recombination and an overexpression vector, recombination and viability plates maintained both plasmids. To measure rDNA stability, yeast with URA3 inserted into the rDNA locus (a gift of D. Koshland, University of California, Berkeley, Berkeley, CA) was treated as for the recombination assays except that loss of URA3 was measured by the frequency of 5′fluoroorotic acid–resistant colonies (Wahba et al., 2011). Cell viability was measured by growing test strains on SC minus uracil plates. To maintain plasmids, all aspects of the rDNA instability assay were done on media lacking leucine. Finally, frequencies of chromosome III loss were quantified in MAd haploid knockout collection strains using the ALF assay essentially as described (Ang et al., 2016). In brief, overnight cultures of haploid MAd cells were mixed 1:3 with a MATa mating tester strain, pelleted, and spotted in 100 µl of sterile water onto synthetic media (SD) lacking all amino acids, where only prototrophic diploids can grow. Mated ALFs form colonies on the SD plates. Cell viability of the overnight culture was determined by diluting cells 1:100,000, plating 100 µl on YPD, and counting colonies after 48 h. The frequency of mated colonies on the SD plate was expressed as a frequency of the total viable cells plated. Graphing and statistical analyses were done in GraphPad (Prism).

Yeast chromosome spreads and live cell imaging

Chromosome spreads were performed as previously described (Wahba et al., 2011; Chan et al., 2014a). In brief, midlog cells grown at 30°C in YPD were washed and spheroplasted in 1.2 M sorbitol, 0.1 M potassium phosphate, 0.5 M MgCl₂, 10 mM dithiothreitol, and 150 µg/ml Zymolyase 20T, pH 7.0, for 20 min at 37°C (Chan et al., 2014a). Ice-cold stop solution (0.1 M 2-[N-morpholino]ethanesulfonic acid, 1 M sorbitol, 1 mM EDTA, and 0.5 mM MgCl₂, pH 6.4) was added before lysis with 1% vol/vol Liposol and fixation and spreading on glass slides in 4% wt/vol paraformaldehyde and 3.4% wt/vol sucrose. Spreads were incubated with a 1:1,000 dilution of S9.6 antibody (mouse; Kerafast) in blocking buffer (5% BSA and 0.2% skim milk powder in 1X PBS) overnight at 4°C before washing three times in blocking buffer and incubating with a 1:1,000 dilution of Cy3-conjugated goat anti-mouse antibody for 1 h (115-165-003; Jackson Laboratories). Slides were washed three times with blocking buffer and mounted in FluorSave mounting media (Calbiochem) before imaging. For each sample, at least 60 nuclei were visualized, and the nuclear fluorescent signal
was quantified using ImageJ (Schneider et al., 2012). Each mutant was assayed in quadruplicate. For comparison purposes, the S9.6 median fluorescence intensity of the WT strain of each experiment was used for normalization. Mutants were compared with WT by the unpaired t test.

For live cell imaging, cells expressing Rad52-YFP were grown to logarithmic phase before any indicated treatments. Log-phase cells, treated or untreated, were bound to concanavalin-A–coated slides and imaged on a Leica dmi8 inverted fluorescence microscope using the appropriate filter sets (see Microscope image acquisition; Stirling et al., 2012).

Microscope image acquisition
Both yeast and human cell images were acquired using an Objective HCX PL APO 1.40 NA oil immersion 100x objective (Leica) on an inverted DMi8 microscope (Leica) equipped with a motorized Differential Interference Contrast imaging turret and a filter cube set for FITC/ YFP/DAPI/TRITC for multicolor immunofluorescence. All images were captured at room temperature using a scientific complementary metal oxide semiconductor camera (ORCA Flash 4.0 V2; Hamamatsu) and collected using MetaMorph Premier acquisition software and post-processed (including gamma adjustments, counting of cells with/without foci, and intensity measurements) using ImageJ. For all microscopy experiments, the significance of the differences was determined using Prism5 (GraphPad) or R. For intensity measurements, samples were compared with t tests or ANOVA; GraphPad performs F-tests for variance as part of this analysis. For comparisons of proportions, Fisher’s tests were used and p-values were Holm-Bonferroni–corrected in the event of multiple comparisons. Sample sizes were determined post hoc and are listed in the figure legends.

DRIP-chip and ChIP-chip analysis
DRIP- and ChIP-chip were generated and analyzed as described previously (Stirling et al., 2012; Chan et al., 2014a). For γ-H2A profiles, 5 μl of anti–γ-H2A antibody was used (rabbit; ab15083; Abcam), and profiles were normalized to an h2a-S129A mutant. Sgs1-flag profiles were generated using 4.2 μl of antibody flag (mouse; F3165; Sigma-Aldrich) and profiles were normalized to mock immunoprecipitates. Complete datasets can be found at ArrayExpress: E-MTAB-5582. Data were normalized using the rMAT software (Droit et al., 2010). All profiles were generated in duplicate with averaged and quantile normalized data used for plotting and calculating mean enrichment scores. Mean feature scores were generated by averaging all probes whose start sites fell within the start and end positions of the desired genomic feature. The same was done for Rrm3 peaks (coordinates derived from Herrera-Moyano et al., 2014). CHROMATRA plots were generated as described previously with genes aligned by their transcription start site (TSS) and sorted by length (Hentrich et al., 2012). Mean gene profiles were generated by averaging all probes that mapped to the genes of interest. Here, probes mapping to features of interest were split into 40 bins, and probes matching to the 1,500 bp of flanking sequences were split into 20 bins. Gene length mean gene profiles were generated by splitting all genes into gene-length classes and averaging probes in 150-bp increments. Enriched features were determined as those where at least 50% of the probes had values greater than 1.5. Only genes appearing in both replicates were considered. Comparing the length of enriched genes was done using a Mann–Whitney test (GraphPad).

Mutation accumulation and whole genome sequencing
Mutation accumulation experiments were conducted as described (Segovia et al., 2017). Single colonies from passage 40 were grown overnight in YPD to prepare genomic DNA by two rounds of phenol-chloroform extraction (Stirling et al., 2014). Whole genomes were sequenced using the Illumina HiSeq2500 platform, and sequence files were deposited at the NCBI sequence read archive (accession no. SRP094860 for sgs1Δ/LA genomes, and no. SRP091984 for WT genomes). Read quality control, alignment to UCSC saccer3, and variant calling were performed exactly as described (Segovia et al., 2017). CNVs were detected using an in-house version of CNAseq (Jones et al., 2010) and Nexus copy number 7.5.2 (Biodiscovery Inc.). Variants were manually checked for read support using the Integrated Genomics Viewer.

Cell culture and transfection
HeLa cells were cultivated in DMEM (Stemcell Technologies), whereas HCT116 was grown in McCoy’s 5A media, both supplemented with 10% FBS (Life Technologies) in 5% CO2 at 37°C. Immortalized Bloom’s syndrome fibroblast lines (gift from J. Campisi, Buck Institute, Novato, CA) were grown as previously described (Davalos and Campisi, 2003). For RNA interference, cells were transfected with either single siRNA sequences targeting BLM (si-BLM, 5′-GCUAGG AGUCUGCGUCGCCGA-3′), FANC D2 (si-FANC D2, 5′-GGUCAG AGCUGUAUAUCUC-3′; Blackford et al., 2015; Schwab et al., 2015), Luciferase GL3 Duplex as a control (si-Luc, 5′-GUUACGCGAGUACUCCA-3′), or siGENOME-SMARTpool siRNAs from Dharmacon (Non-targeting siRNA Pool 1 as si-Com, si-BLM, si-FANC M, si-TOP3A, si-RECQL5, and si-WRN). Transfections were done with Dharmafect1 transfection reagent (Dharmacon) according to the manufacturer’s protocol and harvested 48 h after the siRNA administration. For experiments with overexpression of GFP or nuclear-targeting GFP-RNaseH1 (gift from R. Crouch, National Institutes of Health, Bethesda, MD), transfections were performed with Lipofectamine 3000 (Invitrogen) according to manufacturer’s instructions 24 h after the siRNA transfections.

Immunofluorescence
For S9.6 staining, cells were grown on coverslips overnight before siRNA transfection and plasmid overexpression. 48 h after siRNA transfection, cells were washed with PBS, fixed with ice-cold methanol for 10 min, and permeabilized with ice-cold acetone for 1 min. After PBS wash, cells were blocked in 3% BSA and 0.1% Tween 20 in 4× SSC buffer for 1 h at room temperature. Cells were then incubated with primary antibody S9.6 (1:500; mouse, ENH001; Kerafast) overnight at 4°C. For HCT116, nucleolin was stained by coincubating with anti-nucleolin (rabbit; ab22758; Abcam) at 1:1,000. Cells were then washed three times in PBS and stained with mouse Alexa-Fluoro-568-conjugated secondary antibody (1:1,000; Life Technologies) for 1 h at room temperature, washed three times in PBS, and stained with DAPI for 5 min. Cells were imaged on LeicaDM18 microscope at 100x, and ImageJ was used for processing and quantification of S9.6 intensity in images. Only GFP-positive cells were counted, and micronuclei were counted in asynchronous cells from the same slides. For γ-H2AX and FANC D2 foci, the immunostaining was performed the same way with the differences of fixation with 4% paraformaldehyde for 15 min and permeabilization with 0.2% Triton X-100 for 5 min on ice. Primary antibodies for γ-H2AX (rabbit; ab81299; Abcam), FANC D2 (rabbit; NB100-182SS; Novus) and rabbit Alexa-Fluoro-568-conjugated secondary antibody were all diluted 1:1,000. Where indicated, cells were treated with DMSO or 2 mM HU (Sigma-Aldrich) for 2 h before fixing.

Neutral comet assay
The neutral comet assay was performed using the CometAssay Reagent kit for Single Cell Gel Electrophoresis Assay ( Trevigen) in accordance with the manufacturer’s instructions. Electrophoresis was performed at 4°C, and slides were stained with PI and imaged on LeicaDM18 microscope at 20x. Comet tail moments were obtained using an ImageJ.
plugin as previously described (Mathew et al., 2014). At least 50 cells per sample were analyzed from each independent experiment.

**Western blotting**

Whole cell lysates were prepared with RIPA buffer containing protease inhibitor (Sigma-Aldrich) and phosphatase inhibitor (Roche Applied Science) cocktail tablets, and the protein concentration were determined by Bio-Rad Protein assay (Bio-Rad). Equivalent amounts of protein were resolved by SDS-PAGE and transferred to polyvinylidene fluoride microporous membrane (Millipore), blocked with 5% skim milk in TBS containing 0.1% Tween 20, and membranes were probed with the following antibodies: BLM (rabbit; ab2179; Abcam), FANCM (rabbit; ab95014; Abcam), GADD45 (mouse; MA5-15738; Thermo Fisher Scientific), FANCD2 (rabbit; NB100-182SS; Novus), RECQL5 (rabbit; A302-520A-T; Bethyl), WRN (rabbit; A300-238A-T; Bethyl), TOP3A (rabbit; 14525–1-AP; Proteintech), and α-tubulin (mouse; 32-2500; Life Technologies). Secondary antibodies were conjugated to HRP, and peroxidase activity was visualized using Chemiluminescent HRP substrate (Thermo Fisher Scientific).

**Proximity ligation assay**

Cells were grown on coverslips, washed with PBS, and fixed with 4% paraformaldehyde for 15 min. After permeabilization with 0.2% Triton X-100 for 5 min, cells were blocked in 3% BSA, 0.1% Tween 20 in 4x SSC for 1 h at room temperature. Cells were then incubated with primary antibody overnight at 4°C (1:500 rabbit BLM antibody; PLA0029; Sigma-Aldrich,); as negative control; 1:200 mouse S9.6 antibody as negative control; 1:1000 rabbit BLM antibody with 1:200 mouse S9.6; or 1:1000 rabbit γ-H2AX (Abcam) with 1:200 mouse S9.6 as positive control. After washing with 1x PBS twice, cells were incubated with premixed PLA probe minus mouse and PLA probe antirabbit plus (Sigma-Aldrich) for 1 h at 37°C. Binding of PLA probes, ligation, and amplification was performed with the reagents from the Duolink In Situ kit (Sigma-Aldrich) according to the manufacturer’s instructions. Slides were mounted in Duolink In Situ Mounting Medium with DAPI and imaged on LeicaDMi8 microscope at 100x.

**In vitro helicase assay**

BLM was tagged at the N-terminus with GST and at the C-terminus with His10 and purified as described for PALB2 (Buisson et al., 2010). BLM was tagged at the N-terminus with GST and at the C-terminus with His10 and purified as described for PALB2 (Buisson et al., 2010). BLM was tagged at the N-terminus with GST and at the C-terminus with His10 and purified as described for PALB2 (Buisson et al., 2010). BLM was tagged at the N-terminus with GST and at the C-terminus with His10 and purified as described for PALB2 (Buisson et al., 2010).

**Unwinding assays** were performed in MOPS buffer (25 mM MOPS, pH 7.0, 60 mM KCl, 0.2% Tween-20, 2 mM DTT, 2 mM ATP, and 2 mM MgCl2). BLM or Sgs1 and labeled R-LOOP or D-LOOP (100 nM) substrates were incubated in MOPS buffer for 20 min at 37°C, followed by deproteinization in one-fifth volume of stop buffer (20 mM Tris-Cl, pH 7.5, and 2 mg/ml proteinase K) for 20 min at 37°C. Reactions were loaded on an 8% acrylamide gel, run at 150 V for 120 min, dried onto filter paper, and autoradiographed.

**Online supplemental material**

Fig. S1 shows additional quantification of ALF phenotypes for yeast mutants. Fig. S2 highlights additional analyses of Sgs1 binding to the yeast genome and the position of R-loops and γ-H2A2 relative to telomeres and rDNA, as well as an rDNA instability assay. Fig. S3 shows BLM levels in different cell types, S9.6 staining data for an additional BLM knockout cell line, and example images and quantification of micronucleus formation in BLM knockdown cells. Fig. S3 also shows Western blots of the various knockdown experiments in Fig. 7. Fig. S4 shows the accumulation of FANCD2 foci in HU-treated cells, and the impact of BLM knockdown on this phenotype. Table S1 is a summary of yeast strains and plasmids used in this study. Table S2 summarizes the positions of mutations, CNVs, and indels detected in the sgs1Δ mutation accumulation experiments.

**Acknowledgments**

We acknowledge Andres Aguilera, Judith Campisi, Peter Cejka, Frederic Chedin, Karlene Cimprich, Robert Crouch, and Doug Kosher for providing reagents, cell lines and methods. We thank Philip Hieter, in whose laboratory some of the early work was conceived.

This research is funded by the Canadian Cancer Society (grant 703263), the Canadian Institutes of Health Research (grant MOP-136982 to P.C. Stirling and project grant 363317 to J.-Y. Masson), and the Natural Sciences and Engineering Research Council of Canada (grant RGPIN 402095-11 to M.S. Kober). P.C. Stirling is a Canadian Institutes of Health Research New Investigator and Michael Smith Foundation for Health Research Scholar. J.-Y.M. is a Fond de Recherche du Québec - Santé Chair in genome stability.

The authors declare no competing financial interests.

Author contributions: E.Y.-C. Chang, C.A. Novoa, M.J. Aristizabal, Y. Coulombe, R. Segovia, R. Chaturvedi, C. Keong, and A.S. Tam performed the experiments and analyzed data. Y. Shen provided formal data analysis. S.J.M. Jones, J.-Y. Masson, M.S. Kober, and P.C. Stirling provided the infrastructure, supervision, and funding. E.Y.-C. Chang, C.A. Novoa, M.J. Aristizabal, J.-Y. Masson, and P.C. Stirling conceived of and designed the experiments. E.Y.-C. Chang, C.A. Novoa, M.J. Aristizabal, and P.C. Stirling wrote the manuscript.

Submitted: 23 March 2017

Revised: 16 August 2017

Accepted: 18 September 2017

**References**

Ang, J.S., S. Duffy, R. Segovia, P.C. Stirling, and P. Hieter. 2016. Dosage Mutator Genes in Saccharomyces cerevisiae: A Novel Mutator Mode-of-action of the Mph1 DNA Helicase. *Genetics*. 204:975–986. https://doi.org/10.1534/genetics.116.192211

Ashton, T.M., and I.D. Hickson. 2010. Yeast as a model system to study RecQ helicase function. *DNA Repair (Amst.).* 9:303–314. https://doi.org/10.1016/j.dnarep.2009.12.007

Bhatia, V.I, S.I. Barroso, M.L. García-Rubio, E. Tumini, E. Herrera-Moyano, and A. Aguilera. 2014. BRCA2 prevents R-loop accumulation and associates
53BP1 nuclear bodies form around DNA lesions generated by mitotic transmission of chromosomes under replication stress. *Nat. Cell Biol.* 13:243–253. https://doi.org/10.1038/ncl2201

Mathew, V. A., L. Pauleau, N. Steffen, A. Bergner, P.B. Becker, and S. Erhardt. 2014. The histone-fold protein CHAC1 influences chromatin composition in response to DNA damage. *Cell Reports.* 7:321–330. https://doi.org/10.1016/j.celrep.2014.03.008

Mischo, H.E., B. Gómez-González, P. Grzechnik, A.G. Rondón, W. Wei, L. Steinmetz, A. Aguilera, and N.J. Proudfoot. 2011. Yeast Sen1 helicase protects the genome from transcription-associated instability. *Mol. Cell.* 41:21–32. https://doi.org/10.1016/j.molcel.2010.12.007

Mundberg, K., S.W. Jørgensen, J. Fredsoe, I. Nielsen, J.M. Pedersen, I.B. Bentsen, M. Lisby, L. Bjerregaard, and A.H. Andersen. 2015. Top2 and Sgs1-Top3 Act Redundantly to Ensure rDNA Replication Termination. *PLoS Genet.* 11:e1005697. https://doi.org/10.1371/journal.pgen.1005697

Myung, K, A. Datta, C. Chen, and R.D. Kolodner. 2001. SGS1, the Saccharomyces cerevisiae homolog of BLM and WRN, suppresses genome instability and homologous recombination. *Nat. Genet.* 27:113–116. https://doi.org/10.1038/sj.nrg.3500736

Naim, V., and F. Rosselli. 2009. The FANC pathway and BLM collaborate during translesion synthesis by Rev3 polymerase in cisplatin-treated yeast. *Nat. Genet.* 41:114:2663–2668. https://doi.org/10.1038/ng.427

Neff, N.F., N.A. Ellis, T.Z. Ye, J. Noonan, K. Huang, M. Sanz, and M. Proytcheva. 2014. The histone-fold protein CHR AC14 influences chromatin transmission of chromosomes under replication stress. *Nat. Cell Biol.* 16:2826–2835. https://doi.org/10.1093/emboj/16.10.2826

Popuri, V., C.Z. Bachrati, L. Muzzolini, G. Mosedale, S. Costantini, E. Giacomini, T.J. Pohl, C.F. Chen, A. Chan, S. Pott, and V.A. Zakian. 2016. PIF1 family DNA helicases suppress R-loop mediated genome instability at tRNA genes. *Nat. Commun.* 8:15025. https://doi.org/10.1038/ncomms15025

Sgs1/Blm suppresses r-loop formation • Chang et al. 4005

Skourti-Stathaki, K., N.J. Proudfoot, and N. Gromak. 2011. Human senataxin resolves RNA/DNA hybrids formed at transcriptional pause sites to promote Xrn2-dependent termination. *Mol. Cell.* 42:794–805. https://doi.org/10.1016/j.molcel.2011.04.026

Sollier, J., C.T. Stork, M.L. García-Rubio, R.D. Paulsen, A. Aguilera, and K.A. Cimprich. 2014. Transcription-coupled nucleotide excision repair factors promote R-loop-induced genome instability. *Mol. Cell.* 56:777–785. https://doi.org/10.1016/j.molcel.2014.10.020

Stirling, P.C., M.S. Bloom, T. Solanki-Patil, S. Smith, P. Sipahimalani, Z. Li, M. Kofoid, S. Ben-Aroya, K. Myung, and P. Hieter. 2011. The complete spectrum of yeast chromosome instability genes identifies candidate CIN cancer genes and functional roles for ASTRA complex components. *PLoS Genet.* 7:e1002057. https://doi.org/10.1371/journal.pgen.1002057

Stirling, P.C., Y.A. Chan, S.W. Minaker, M.J. Aristizabal, I. Barrett, P. Sipahimalani, M.S. Kobor, and P. Hieter. 2012. R-loop-mediated genome instability in mRNA cleavage and polyadenylation mutants. *Genes Dev.* 26:165–175. https://doi.org/10.1101/gad.179721.111

Stirling, F.C., Y. Shen, R. Corbett, S.J. Jones, and P. Hieter. 2014. Genome destabilizing mutator alleles drive specific mutational trajectories in Saccharomyces cerevisiae. *Genetics.* 196:403–412. https://doi.org/10.1534/genetics.113.159806

Stork, C.T., M. Bocek, M.P. Crossley, J. Sollier, L.A. Sanz, F. Chédin, T. Swigut, and K.A. Cimprich. 2016. Co-transcriptional R-loops are the main cause of estrogen-induced DNA damage. *elife.* 5:e17548. https://doi.org/10.17554/elife.17548

Stratton, M.R., P.J. Campbell, and P.A. Futreal. 2009. The cancer genome. *Nature.* 458:719–724. https://doi.org/10.1038/nature07943

Suhasini, A.N., and R.M. Brosh Jr. 2012. Fanconi anemia and Bloom’s syndrome crosstalk through FANCJ-BLM helicase interaction. *Trends Genet.* 28:7–13. https://doi.org/10.1016/j.tig.2011.09.003

Tran, P.L.T., T.J. Pohl, C.F. Chen, A. Chan, S. Pott, and V.A. Zakian. 2017. PIF1 family DNA helicases suppress R-loop mediated genome instability at tRNA genes. *Nat. Commun.* 8:15025. https://doi.org/10.1038/ncomms15025

Tresini, M., D.O. Warmerdam, P. Kolovos, L. Snijder, M.G. Vrouwe, J.A. Demmers, W.F. van IJcken, F.G. Grosvold, R.H. Medema, J.H. Hoeijmakers, et al. 2015. The core splicesome as target and effector of non-canonical ATM signalling. *Nature.* 523:53–58. https://doi.org/10.1038/nature14512

Tumini, E., S. Barrosso, C.P. -Calero, and A. Aguilera. 2016. Roles of human POLD1 and POLD3 in genome stability. *Sci. Rep.* 6:38873. https://doi.org/10.1038/srep38873

Versini, G., I. Comet, M. Wu, L. Hoopes, E. Schwob, and P. Pasero. 2003. The yeast Sgs1 helicase is differentially required for genomic and ribosomal DNA replication. *EMBO J.* 22:1939–1949. https://doi.org/10.1093/emboj/cdg180

Vijayraghavan, S., F.L. Tsai, and A. Schwacha. 2016. A Checkpoint-Related Function of the MCM Replicative Helicase Is Required to Avert Accumulation of RNA:DNA Hybrids during S-phase and Ensuing DSBs during G2/M. *PLoS Genet.* 12:e1006277. https://doi.org/10.1371/journal.pgen.1006277

Wahba, L., J.D. Amon, D. Koshland, and M. Vuica-Ross. 2011. RNase H and multiple RNA biosynthesis factors cooperate to prevent RNA:DNA hybrids from generating genome instability. *Mol. Cell.* 44:978–988. https://doi.org/10.1016/j.molcel.2011.10.017

Wahba, L., S.K. Gore, and D. Koshland. 2013. The homologous recombination machinery modulates the formation of RNA-DNA hybrids and associated chromosome instability. *eLife.* 2:e00505. https://doi.org/10.7554/eLife.00505

Zhang, X., H.C. Chiang, Y. Wang, C. Zhang, S. Smith, X. Zhao, S.J. Nair, J. Michalek, I. Jato, M. Lautner, et al. 2017. Attenuation of RNA polymerase II pausing mitigates BRC1-associated R-loop accumulation and tumorigenesis. *Nat. Commun.* 8:15908. https://doi.org/10.1038/ncomms15908

Zimmer, A.D., and D. Koshland. 2016. Differential roles of the RNases H in preventing chromosome instability. *Proc. Natl. Acad. Sci. USA.* 113:12220–12225. https://doi.org/10.1073/pnas.1613448113