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Role for the Fission Yeast RecQ Helicase in DNA Repair in G2

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Members of the RecQ helicase subfamily are mutated in several human genomic instability syndromes, such as Bloom, Werner, and Rothmund-Thomson syndromes. We show that Rqh1, the single Schizosaccharomyces pombe homologue, is a 3′-to-5′ helicase and exists with Top3 in a high-molecular-weight complex. top3 deletion is inviable, and this is suppressed by concomitant loss of rqh1 helicase activity or loss of recombination functions. This is consistent with RecQ helicases in other systems. By using epistasis analysis of the UV radiation sensitivity and by analyzing the kinetics of Rhp51 (Rad51 homologue), Rqh1, and Top3 focus formation in response to UV in synchronized cells, we identify the first evidence of a function for Rqh1 and Top3 in the repair of UV-induced DNA damage in G2. Our data provide evidence that Rqh1 functions after Rad51 focus formation during DNA repair. We also identify a function for Rqh1 upstream of recombination in an Rhp18-dependent (Rad18 homologue) pathway. The model that these data allow us to propose helps to reconcile different interpretations of RecQ family helicase function that have arisen between work based on the S. pombe system and models based on studies of Saccharomyces cerevisiae SGS1 suggesting that RecQ helicases act before Rad51.

The RecQ family of helicases is a subfamily of the DExH box containing DNA helicases. Homologues of the prototype bacterial RecQ helicases have been identified in a wide range of organisms spanning the prokaryotic kingdom up to humans. RecQ family helicases share extensive sequence homology within the seven helicase domains, but conservation in the remainder of the protein is generally limited to structural features such as acid-rich domains (10). Five RecQ family proteins have been identified in humans (37), and three syndromes represented by these proteins have been identified as defective in human genetic disorders: Bloom syndrome, Werner syndrome, and a subset of Rothmund-Thomson syndrome (RTS). Each syndrome presents with very different phenotypes: Bloom syndrome, which is associated with a defect in the BLM protein, is characterized by stunted growth, developmental problems, immunodeficiency, male infertility, and, at the cellular level, a slowing of S-phase progression and a dramatic increase in sister chromatid exchange and genomic instability (21). Werner syndrome, linked to a defect in the WRN protein, is a premature-ageing syndrome, and this is reflected at the cellular level by a decreased population-doubling potential (55). RTS (RECCQ defective) comprises poikiloderma and growth deficiency, and RTS patients also exhibit aspects of premature ageing. All three syndromes are characterized by an increased predisposition to cancer, which is consistent with the increased chromosomal aberrations and hypermutability observed in cultured cells (31). These data suggest that each of these helicases contributes to maintaining genetic stability.

The prototype Escherichia coli recQ mutant is defective in the initiation of recombinational bypass of DNA damage during postreplication repair (PRR). PRR can be defined as the ability of the replication fork to reform and/or bypass unrepaird DNA damage by using recombination proteins. Purified E. coli RecQ helicase is capable of acting in concert with RecA during branch migration and drives the reaction in the direction opposite to that of RecBC, i.e., towards the original unrecombined substrates (24). In addition, when acting in conjunction with topoisomerase III (Top3), RecQ modifies the activity of Top3 in vitro to enable it to act as a type II topoisomerase, promoting plasmid catenation and/or decatenation (25). E. coli topoisomerase III, on its own, acts like a type I topoisomerase, changing the linking number of DNA in steps of one (16).

Both budding and fission yeasts have a single RecQ-like helicase. These are structurally most reminiscent of the human BLM protein. Saccharomyces cerevisiae SGS1 was originally identified because its loss acted as a slow-growth suppressor of top3 mutants. Subsequently, the Sgs1 protein was found to interact with Top3 and to act as a 3′-to-5′ helicase (3, 20). By use of the two-hybrid system, a potential interaction between Sgs1 and Top2 has also been demonstrated (49). In the fission yeast Schizosaccharomyces pombe, the RecQ homologue, Rqh1, was independently identified in screens for radiation-sensitive (rad) and hydroxyurea-sensitive (hus) mutants. Like SGS1, Rqh1 is not an essential gene and mutants lacking rqh1 (rqh1-d) are significantly sensitive to DNA damage. rqh1-d mutants also display an increase in recombination when replication is inhibited (42, 45). Genetic analysis of the sensitivity of S. pombe rqh1-d mutants to UV-induced DNA damage defined a function for Rqh1 in the DNA “damage tolerance” pathway, which also requires recombination and checkpoint functions. This led to the suggestion that rqh1 mutants are defective in the recombinational bypass of UV-induced DNA damage during the S phase (42).

The human BLM and the S. cerevisiae Sgs1 proteins have
been shown to be hexameric helicases (4, 28, 29), and both BLM and Sgs1 have been demonstrated to associate with Top3 directly (3, 20, 50), suggesting that complexes formed by this family of helicases are conserved through evolution. Because defects in RecQ family helicases affect recombination-dependent pathways in all systems studied, it is generally accepted that these proteins regulate recombination. We have investigated the genetic and biochemical interactions of S. pombe Rqh1 with Top3. Our data support a role for Rqh1 in recombination and lead us to propose a new role for Rqh1-Top3 in the repair of UV-induced DNA damage in G2 cells via recombination-mediated repair.

MATERIALS AND METHODS

Genetics and cell biology techniques. Strains were constructed by using standard genetic techniques (38). The protocols for checkpoint measurements, cell scoring, and irradiation have previously been described (18). Indirect immunofluorescence microscopy was performed according to a protocol previously described (8). Brefeldin-a, cells were fixed in 3.7% (vol/vol) paraformaldehyde for 10 min and were stained with primary antibody (9E10 at 1:100, anti-Rqh1 antibody at 1:100, and anti-Rad51 and secondary antibody [fluorescein isothiocyanate-conjugated anti-mouse (Dako)] at 1:150). Synchronized cells were prepared by lactose gradients (2) or by elutriation of 4 liters of mid-log-phase cells (JE-50; Beckman Coulter).

PCR-based gene deletion and tagging. By use of a PCR-based method, the entire top3 open reading frame (ORF) was replaced with either the S. pombe ura4+ gene or the S. cerevisiae LEU2+ gene (top3-d:ura4, top3-d:LEU2). A separate construct, where the Rqh1 helicase domain (bp 1527 to 2517, amino acids 526 to 869) was replaced by the untagged rhq1 (rhq1-hd:ura4), was also created. The top3 and rhq1 genes were tagged with sequences encoding either a C-terminal three-hemagglutinin (HA) or 13-MYC (MYC) epitope as previously described (1). Sequences encoding either an N-terminal 3HA or N-terminal 2xMYC-six-His epitope were introduced at the N terminus of the ORF at the top3 locus by first replacing the ATG with the ura4+ gene and then by using fluoroorotic acid gene replacement to introduce the engineered fragment. The final construct, where the native promoter drives the tagged ORF, was checked by PCR and characterized by Western blot analysis.

Cloning of top3 cDNA. Total RNA was prepared from S. pombe by using the Qiagen RNaseasy minikit, and cDNA was produced with the Gibco BRL SuperScript II protocol. The top3 cDNA was amplified by using PCR primers designed from the top35' sequence (Sanger Centre) and was cloned into pREP1 and pREP4/HVA vectors (15) behind a modified thiamine-repressible promoter (npr1), which fuses it to a sequence encoding three HA epitopes at the 5' end. Overexpression of HA-Top3 resulted in a protein at a size corresponding to ~78 kDa, consistent with the predicted molecular mass of 71.5 kDa for the untagged protein. The lethality of the top3-d:ura4 strain could be complemented by ectopic expression of the cloned cDNA.

Rqh1 mutant alleles. The rhq1-K547A and rhq1-K547R mutations were introduced into the cDNA by site-directed mutagenesis with the USE mutagenesis kit (Amersham Pharmacia). The rhq1-T543I allele was PCR amplified from the genome, and the BamHI-EcoNI fragment from this was used to replace 356 bp (1272 to 1628) of the rhq1 ORF. The sequence of the cDNAs was confirmed by sequencing. To create genomic mutations of rhq1-K547A and rhq1-K547R, the cDNA was used to replace the ura4+ gene in rhq1-hd:ura4 at the rhq1 genomic locus. Correct integration was selected for by fluoroorotic acid resistance and was confirmed by PCR and Western blotting.

Helicase assays. The rhq1 wild-type and mutant alleles were cloned (NdeI-Sall) into pPEPEx-HA and were in vitro translated by using the T7 quick-coupled Tnt kit (Promega). Expression was checked by Western blot analysis. The in vitro translated proteins were then tested for their ability to release an [alpha-32P]-labeled 40-mer oligonucleotide annealed to a single-stranded plasmid, M13mp18 (NEB). Some 40-mer (3.5 pmol) was hybridized to 3.5 pmol of plasmid by heating to 85°C and cooling slowly to room temperature. Two micro liters of [alpha-32P]-dATP and 1.5 μl of Sequenase was added, the reaction materials were incubated at 30°C for 15 min, and the substrate was purified by a G-50 column. Helicase reactions (10 μl) containing 0.1 μl of labeled substrate (approximately 3.5 pmol) were performed in helicase buffer (25 mM Tris-HCl, pH 7.5, 1 mM dithiothreitol, 50 μg of bovine serum albumin/ml, 2 mM MgCl2, and 2 mM ATP) for 30 min at 30°C. For each protein, three reactions were made containing increasing concentrations of in vitro translated protein: 2, 4, and 8 μl, the volume being made to a standard 8 μl with an "empty vector" in vitro translation mix. The reactions were stopped by the addition of 0.5% proteinase K, 100 mM Tris-HCl, pH 7.5, 200 mM EDTA, and 2.5% sodium dodecyl sulfate (SDS) and incubation at 37°C for 10 min. Products were analyzed by electrophoresis in a 15% nondenaturing polyacrylamide gel and were subsequently autoradiographed.

Cell extracts, immunoprecipitation, and size fractionation. The protocols for total cell extracts, soluble extracts, size fractionation by using a Superdex 200 HR 10/30 (Pharmacia) column, and immunoprecipitation experiments have all been previously described (8). Soluble and/or insoluble fractionation was performed as previously described (12).

Antibodies. An rhq1 N-terminal 1.085-kb fragment encoding amino acids 1 to 361 was PCR amplified by using primers incorporating NdeI and BamHI sites and was cloned into pET16b (Novagen). The His-tagged protein, purified under denaturing conditions according to the manufacturer's instructions, was used for immunization of rabbits. The anti-Rqh1 antibody recognized a single band of approximately 175 kDa in trichloroacetic acid extracts of wild-type cells that was absent in rhq1 null mutants and increased in size to ~190 kDa in the rhq1-MYC strain (Fig. 2f). The antibody was affinity purified against the antigen by using AminoLink Plus Coupling gel (Pierce). Rqh1-MYC or Top3-MYC was detected with anti-MYC monoclonal antibody (9E10; PharMingen), and Top3-HA was detected with anti-HA monoclonal antibody (Babco, Richmond, Calif.).

RESULTS

Rqh1 is a 3'-to-5' helicase, and mutations in helicase domain 1 abolish helicase activity. The RecQ family has been demonstrated to have helicase activity in vitro. Since the Rqh1 helicase domains are highly conserved, we anticipated that Rqh1 would share this activity. To demonstrate this, we performed an in vitro helicase assay by using recombinant protein produced by coupled transcription-translation (Promega) in rabbit reticulocyte extract. When primed with either wild-type or mutant rhq1 cDNAs, extracts were found to express a protein of the expected size (175 kDa) upon analysis by SDS-polyacrylamide gel electrophoresis (PAGE) (Fig. 1a). Extracts expressing wild-type Rqh1 were able to displace a radiolabeled 40-base oligonucleotide that had been previously annealed to single-stranded M13mp18 phage DNA (Fig. 1b). Extracts primed with empty vector were not active in this assay, demonstrating dependence on Rqh1. In a similar reaction, Rqh1 was also able to displace a 90-base oligonucleotide, although with lower activity (data not shown). We also demonstrated that, as expected, Rqh1 acts as a 3'-to-5' helicase. In this experiment (Fig. 1c), an oligonucleotide (25 bases) was released from an 85-base oligonucleotide only when annealed to the 5' end. An oligonucleotide (35 bases) annealed at the 3' of the 85-base oligonucleotide was not a substrate for Rqh1-dependent displacement.

The original rad12 mutant in the rhq1 gene results in a T543I substitution in Rhq1 (42). rhq1-T543I mutant cells exhibit ionizing radiation (IR) and UV radiation sensitivities that are intermediate between those of wild-type cells and rhq1-d cells (Fig. 1d). The T543I residue is conserved in all RecQ family members and lies within the highly conserved 1a helicase domain, which contains the ATP binding site. To test if Rhq1-T543I is active as a helicase in vitro, we engineered the T543I mutation into the cDNA. Two further ATP binding site mutations, previously demonstrated in other systems to encode helicase-dead proteins, were also engineered as controls. The corresponding recombinant proteins were expressed in vitro by using the transcription-translation system (Fig. 1a). Neither the Rhq1-T543I mutant protein nor either of the two helicase-dead ATP site mutant control proteins (Rqh1-K547R and
FIG. 1. Rqh1 acts as a helicase and mutations in helicase domain 1 abolish in vitro activity. (a) In vitro translation of Rqh1 alleles. Empty pEPEX-HA vector (v), vector containing wild-type rqh1 (wt), or rqh1 mutants rqh1-T543I (T/I), rqh1-K547A (K/A), and rqh1-K547R (K/R) were used to prime TNT reticulocyte lysates, and proteins were identified following SDS-PAGE and Western blotting with anti-Rqh1 antibody. (b) Helicase assays. These were performed by using increasing amounts of primed lysates (as described for panel a above). The substrate, a 40-base \(^{32}\)P-labeled oligonucleotide annealed to single-stranded M13, is shown diagrammatically to the left. Rqh1 has robust activity, while Rqh1-K547R (K/R), Rqh1-K547A (K/A), and Rqh1-T543I (T/I) have no activity. (c) Directionality of the Rqh1 helicase. Different primed lysates (as described for panel a above) were used. The substrate, an 85-base oligonucleotide annealed at each end to a \(^{32}\)P-labeled 25-mer or 35-mer, is diagrammed below. Rqh1 displaced the 25-mer but not the 35-mer, showing it to have 3'-to-5' helicase activity. Neither of the oligonucleotides was displaced by the helicase-dead protein Rqh1-K547R (K/R). (d) UV survival analysis of genomic mutants in rqh1. Symbols: ×, wild type; ■, rqh1-K547R (K/R); ▲, rqh1-K547A (K/A); ○, rqh1-T543I (T/I); ◯, rqh1-h2 (h2); and □, rqh1-d. (e) Western blots show the relative quantities of different Rqh1 proteins in total cell extracts (i) and their fractionation into soluble (S) and insoluble (P) extracts (ii). deg, degradation products, n, nonspecific band recognized by anti-Rqh1.
Rqh1-K547A) showed helicase activity in the in vitro assay (Fig. 1b).

Because the *rqh1-T543I* mutation leads to a phenotype that is less severe than that of the *rqh1-d* null mutant (42) we examined the phenotypes of the *rqh1-K547R* and *rqh1-K547A* ATP site helicase-dead mutants. Genomic mutants encoding these changes were created (see Materials and Methods). These mutants are at the *rqh1* locus and under the control of the native promoter. The response of both mutants to DNA-damaging agents (UV and ionizing radiation) and to inhibition of DNA synthesis (hydroxyurea) was compared to those of wild-type cells (*rqh1*+) and of the *rqh1-T543I* strain. The two helicase-dead mutants are more sensitive than *rqh1-T543I*, approaching the *rqh1-d* null mutant in all assays (Fig. 1d and data not shown). This suggests that the helicase activity of Rqh1 is required for the majority of its biological functions, but the slight difference in survival compared to *rqh1-d* may reflect a function for Rqh1 that is independent of its helicase activity. Since *rqh1-T543I* is less sensitive, it is likely that *rqh1-T543I* is a hypomorphic mutant. However, it remains possible that Rqh1-T543I is helicase dead in vivo as well as in vitro. To investigate this further, we compared total protein level and distribution into soluble and insoluble fractions for Rqh1, Rqh1-T543I, Rqh1-K547R, and Rqh1-K547A proteins.

Cell extracts were prepared from log-phase cultures and fractionated into soluble and insoluble components as described in Materials and Methods. The insoluble fraction should contain the chromatin, along with membranes and organelles. Under the conditions used, Rqh1 was found to be largely insoluble, which would be consistent with Rqh1 being predominantly associated with chromatin or nuclear structures (Fig. 1e). The Rqh1-T543I, Rqh1-K547A, and Rqh1-K547R proteins were present at approximately wild-type levels (Fig. 1e, panel i), but virtually no soluble fraction was detected (Fig. 1e, panel ii). The Rqh1-h2 protein was found to partition equally between the soluble and insoluble fractions. The *rqh1-h2* (initially known as *hus2*) mutant causes a truncation in helicase domain 5 (*Q789*; stop codon) resulting in a null phenotype (Fig. 1d and reference 42). This increase in solubility is most likely to be due to the cytoplasmic localization of the mutant protein, which we observed by indirect immunofluorescence (data not shown).

**Top3 is essential in the presence of an active Rqh1 helicase.** Since RecQ family helicases interact both genetically and biochemically with Top3 in different model systems, we created a *top3* null allele (*top3-d*) by gene replacement in a diploid *S. pombe* strain (Fig. 2a). Consistent with the published data (22, 33), tetrad analysis demonstrated that *top3* is an essential gene in the presence of a wild-type *rqh1* gene but that, when *rqh1* is deleted, deletion of *top3* could be tolerated (Fig. 2b). It is generally considered that, in the absence of Top3, Rqh1 helicase activity processes an unknown DNA structure in a manner that causes lethality. Thus, consistent with the previously published data, we also observe that *top3-d* is tolerated in *rqh1-h2*, the helicase-dead mutants, *rqh1-K547R* and *rqh1-K547A*, and the hypomorphic *rqh1-T543I* mutant (Fig. 2b). Thus, our data are consistent with Top3 being required downstream of the Rqh1 helicase.

Because deletion of Top3 is lethal in the presence of an active helicase, we examined the consequences of overexpression of Top3. The *top3* cDNA was PCR amplified (see Materials and Methods) and was cloned into the expression vector pREP1. In addition, an active site mutant construct (Y330F in the conserved motif SYPTET) was created by site-directed mutagenesis. Overexpression of Top3 led to the phenotype of slightly elongated cells but had no effect on UV survival of wild-type cells (Fig. 2c and data not shown). However, overexpression of the active site mutant, Top3-Y330F, led to extensive cell death (Fig. 2c). The terminal phenotype was elongated cells with a range of defects, including fragmented nuclei, septa bisecting the nucleus (cut cells), and multiple septa (Fig. 2d). These data are consistent with the mutant protein interfering with the ability of the native protein to carry out its normal function (although it is still able to interact with Rqh1 [data not shown]). This dominant-negative phenotype was suppressed by deletion of *rqh1* (Fig. 2c). These data are again consistent with Top3 being required downstream of the Rqh1 helicase and the belief that, in the absence of Rqh1, a DNA structure that requires resolution by Top3 is not generated.

We next examined if loss of *top3* resulted in increased DNA damage sensitivity compared to that of the *rqh1-d* or *rqh1-T543I* single mutant. The double mutant *top3-d rqh1-d* showed no increase in sensitivity to UV or ionizing radiation when compared to *rqh1-d* (Fig. 2e), consistent with Top3 acting in the same pathway as Rqh1. However, the *top3-d rqh1-T543I* double-mutant sensitivity was intermediate between the sensitivities of *rqh1-T543I* and *rqh1-d* single mutants, suggesting that Top3 is required for the DNA damage response (Fig. 2e). However, analysis of the level of Rqh1-T543I protein in the *top3-d* background demonstrated that the protein level is significantly reduced when compared to a *top3*+ background (Fig. 2f). Thus, the absence of Top3 appears to destabilize Rqh1-T543I, possibly because a direct protein association is disrupted or because the accumulation of suppressor mutations leads to a decrease in Rqh1 expression. A reduction in Rqh1-T543I protein level could account for the intermediate phenotype of the *rqh1-T543I top3-d* double mutant: if Rqh1-T543I is a partially functional protein (the result of a hypomorphic mutation), a reduced expression level might compromise the remaining function.

**A physical interaction between Top3 and Rqh1.** To examine the potential physical interaction between Rqh1 and Top3, we integrated sequences encoding 3×HA epitopes either 3' or 5' of the ORF at the *top3* locus. Similarly, we integrated a 2×MYC–six-His encoding sequence 5' of the ORF (see Materials and Methods). The *top3-HA* strain was slightly UV sensitive, but neither the HA-*top3* nor MYC–six-H–*top3* strains showed any phenotype. In both cases, a single band could be detected by SDS-PAGE followed by Western blotting at a molecular weight consistent with the predicted size of Top3 plus the N-terminal tag (Fig. 3a). We performed reciprocal coimmunoprecipitation experiments for Rqh1 and HA-Top3. Immunoprecipitation with anti-Rqh1 antibodies coprecipitated the majority of HA-Top3 from *rqh1*+ but not *rqh1-d* extracts (Fig. 3b). This indicates that Rqh1 and Top3 are present in the same complex in vivo. Likewise, immunoprecipitation with anti-HA coprecipitated Rqh1.

To further verify that Rqh1 and Top3 exist in a common complex, we fractionated cell extracts by using size exclusion
FIG. 2. Characterization of Top3. (a) Diagrammatic representation of the top3 gene, showing the position of the exons and the active-site tyrosine 330 (Y) and the deletion strategy. (b) top3-d is viable in combination with rqh1-d, rqh1-h2, rqh1-K547R, rqh1-K547R, and rqh1-T543I. Cells, derived from tetrad analysis, were streaked to single colonies on rich media and were incubated for 3 days at 30°C. +, top3+; −, top3-d; wt, wild type. (c) Overexpression of the Top3 active-site mutant, top3-Y330F, is lethal in wild-type cells but viable in rqh1-d. Cells were streaked to single colonies on minimal media with thiamine (+Thiamine) and were then replica plated to minimal media without thiamine (−Thiamine) to induce the expression of Top3-Y330F. (d) Wild-type cells overexpressing Top3-Y330F die elongated, with a range of defects, including degraded DNA, cuts, and multiple septa (inset A), compared to rqh1-d cells overexpressing the same construct (inset B) and wild-type cells overexpressing Top3, which are slightly elongated (inset C). (e) UV and IR sensitivity of rqh1 top3 double mutants. Symbols: ×, wild type; □, rqh1-d; ■, rqh1-d top3-d; ○, rqh1-T543I; and ●, rqh1-T543I top3-d. (f) Western blot of total cell extracts showing Rqh1 protein levels in top3+ and top3-d backgrounds. Rqh1 levels are decreased in the top3-d background (rqh1-T543I top3-d compared to rqh1-T543I). No change was observed in wild-type cells after UV irradiation (UV + wt). Log-phase cells were irradiated with 100 J/m² and allowed to recover for 1 h before total cell extracts were prepared by trichloroacetic acid precipitation. Rqh1-MYC migrates slightly higher than the native protein. n is a nonspecific band recognized by anti-Rqh1 shown as a loading control.
Rqh1 acts with recombination proteins. During the unperturbed cell cycle, intrinsic DNA damage—most probably occurring during DNA replication—is processed by recombination proteins. The loss of recombination functions in S. pombe results in severe chromosomal instability and prolonged cell cycle delay (40). Presumably, during this delay, the aberrant DNA structures are repaired by alternative mechanisms. The Rqh1-dependent lethality of the top3-d mutation reflects the fact that Rqh1 activity creates lethal (or irreparable) DNA structures from this intrinsic DNA damage. We can thus use the lethality of top3-d as an assay to identify genetic backgrounds that do not generate the substrate that Rqh1 processes. In this way, we should be able to order the DNA-processing events that act on intrinsically produced aberrant DNA structures.

We first tested DNA integrity checkpoint mutants because it was previously found that rqh1-d caused synthetic lethality in rad3-d and rad26-d mutants but not in other checkpoint mutants (42). top3-d was not viable in rad1-d, rad3-d, and rad26-d mutants (Table 1). In addition, we tested top3-d rqh1-d rad3-d and top3-d rqh1-d rad26-d triple mutants for viability and found that neither was viable. These data indicate that the synthetic lethality previously observed between rqh1-d and rad3-d or rad26-d is not due to inappropriate regulation of Top3 activity. We have also previously demonstrated that Rqh1 acts with the Cds1 kinase in the same S-phase DNA damage tolerance response, which requires the recombination apparatus (42). top3-d cds1-d double mutants are inviable, indicating that Rqh1 is still able to act as a helicase in the absence of Cds1.

Recent work in S. pombe suggests that Rqh1 and Top3 function during the later stages of IR-induced double-strand break (DSB) repair by homologous recombination (9). If Rqh1 acts during recombination in unperturbed S phase, loss of recombination genes should prevent top3-d lethality because it would prevent the formation of the DNA structures that result in Rqh1-dependent top3-d inviability. We thus created double mutants between top3-d and a number of recombination mutants including rhp51-d (recA/RAD51 homologue), rhp54-d (RAD54 homologue), rhp55-d (RAD51 paralogue), and rad22-d (RAD52 homologue) (Table 1). Each of the four double mutants was viable but slow growing (Fig. 4a, panel i, left, and Table 1), indicating that Rqh1 indeed acts in recombination during normal cell growth.

The suppression of the top3-d lethality by loss of recombination genes is consistent with Rqh1 acting downstream of recombination in the processing of intrinsic DNA damage. In this scenario, which is consistent with the analysis of Top3 function in IR-induced DSB repair (9), loss of recombination genes prevents the formation of the DNA structures that result in Rqh1-dependent top3-d inviability. An alternative explanation of these data is that Rqh1 acts upstream of recombination proteins in the processing of intrinsic lesions. In this model, loss of top3-d results in Rqh1 generating a lesion that becomes a substrate for an inappropriate and lethal attempt at recombination.

Overexpression of Top3 with an active-site tyrosine-to-phenylalanine mutation in rhp51-d cells leads to cell death with highly elongated cells (Fig. 2c and d), similar to a terminal phenotype seen in spores germinated with a top3 deletion (reference 22 and data not shown). Because this dominant-
negative phenotype was suppressed in rqh1-d cells (Fig. 2c), we investigated whether it could also be suppressed by deletion of recombination proteins. In the recombination mutant strains rhp51-d, rhp54-d, rhp55-d, and rad22-d, overexpression of the active-site mutant was not lethal. This shows that the dominant-negative phenotype is suppressed by either deletion of rqh1 or deletion of recombination genes (Fig. 4b) and that the suppression of lethality in the corresponding double mutants is not the result of a rapid accumulation of suppressor mutations.

**TABLE 1. List of mutants**

<table>
<thead>
<tr>
<th>Strain description</th>
<th>Viability</th>
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<tr>
<td>rqh1-d</td>
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<tr>
<td>rad3-d</td>
<td>+</td>
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<tr>
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<td>–</td>
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<tr>
<td>rad3-d top3-d</td>
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*+, severe growth defect; **, most double mutants not viable; viable colonies have severe growth defect.

**UV sensitivity of rqh1-d, top3-d, and recombination mutants.**

We next looked at clonogenic survival following UV and ionizing radiation in the rqh1-d, top3-d, and recombination mutants. After ionizing radiation, like the previously characterized double mutants between rqh1-d and recombination gene deletions (42), double mutants between top3-d and recombination gene deletions and the triple rqh1-d top3-d recombination mutant were as sensitive as but not more sensitive than the single recombination mutant (Fig. 4c and data not shown). However, in response to UV irradiation, the sensitivity of rqh1-d and rqh1-d top3-d double mutants was rescued by concomitant loss of rhp51 or rhp55 (Fig. 4a, panels i and ii). Concomitant loss of rhp54 did not result in an obvious rescue. Concomitant loss of rad22 did rescue the UV sensitivity of rqh1-d top3-d, but the double rad22-d rqh1-d and rad22-d top3-d mutants have a severe growth defect, which makes the epistasis analysis difficult to interpret.

These data support the model in which Rqh1 and Top3 act within the recombination response to UV-induced DNA damage. UV survival of cells lacking Rqh1 and Top3 functions is increased in the absence of recombination. This suggests that, in the absence of recombination, DNA damage is not processed into a form that requires processing by Rqh1 and Top3. If the original damage is not a substrate for recombination (i.e., in rhp51-d or rhp55-d mutants), it can be repaired by an alternative pathway. If it is a substrate of recombination (i.e., rhp54-d strains) the damage is processed and a DNA structure is formed that must either be processed by Rqh1-Top3 or result in cell death. Interestingly, this could possibly explain the lack of rescue by rhp54-d: Rhp54 is thought to act later in recombination, whereas Rhp55 and Rhp51 are thought to act to initiate recombination. Once recombination is initiated, loss of a later step (i.e., rhp54-d) would not be expected to rescue the UV sensitivity because an alternative pathway is no longer possible.

Rqh1 and Top3 have been implicated in recombination repair of IR-induced DSBs in G2 (9). To ascertain if there is a role for Rqh1 in the repair of G2-induced UV damage and to verify the restoration of UV resistance by concomitant deletion of recombination genes, we treated G2-synchronized wild-type, rqh1-d, rhp51-d, and rqh1-d rhp51-d cells with UV and assayed survival. rqh1-d cells are sensitive to low doses of UV in G2, and this sensitivity is reversed by concomitant loss of rhp51. This provides evidence of a role of Rqh1 in G2 repair of UV-induced DNA damage by recombination and confirms the rescue of rqh1-d sensitivity by loss of rhp51 in a discrete cell cycle compartment.

**Rqh1 and Top3 form foci following UV irradiation in G2.** The genetic evidence presented above indicates that Rqh1 and Top3 function in recombination after the Rhp51-dependent initiation step. This prompted us to search for further independent supporting evidence by examining protein localization before and after UV-induced DNA damage: first, we synchronized wild-type cells by centrifugal elutriation and examined the localization of Rqh1. Rqh1 localized to the nucleus throughout the cell cycle. During the majority of the cell cycle, Rqh1-dependent fluorescence overlapped with the DNA but was predominantly nucleolar. However, during and immediately after mitosis nuclei appeared diffusely stained and fluorescence was clearly present in areas of the nucleus that did not contain 4′,6-diamidino-2-phenylindole (DAPI)-staining material (Fig. 5a).

Next, we examined Rqh1, MYC–six-His–Top3, and Rhp51...
FIG. 4. top3-d mutants are viable in recombination mutant backgrounds. (a) Panel i, spot assays to show UV survival of recombination mutants rhp51-d, rhp54-d, rad22A-d, and rhp55-d with rqh1-d and top3-d. Strains are spotted in the order of the key. Panel ii, UV survival of recombination mutants rhp51-d, rhp54-d, rad22A-d, and rhp55-d with rqh1-d and top3-d. ‡ indicates that the suppression of the UV sensitivity of rqh1-d top3-d is seen by concomitant deletion of rhp51, rhp55, and rad22 (only the triple mutant). (b) Top3-Y330F overexpression is viable in combination with rhp51-d, rhp54-d, rhp55-d, and rad22-d. Cells were streaked to single colonies on minimal media with thiamine (+Thiamine) and were then replica plated to minimal media without thiamine (~Thiamine) to induce the expression of Top3-Y330F. (c) IR survival of rhp51-d, rqh1-d, and top3-d mutants. (d) UV survival of wild-type and rhp51-d, rqh1-d, and rhp51-d rqh1-d mutants in G2, showing that rhp51-d rescues the UV sensitivity of rqh1-d in G2. Survival curves for the triple mutant lie under those for the single recombination mutant and thus are obscure in most panels.
localization following treatment of cells with UV. After irradiation with 100 J of UV light/m², a significant proportion of cells exhibited distinct foci of Rqh1 in the DAPI-staining area of the nucleus (data not shown). In contrast, Rhp51 and MYC–six-His–Top3 were not detectable in unirradiated cells, but foci were observed following after UV irradiation (data not shown). To determine if the formation of these foci required passage through S phase, we first synchronized cells in G₂ and then irradiated with 100 J of UV light/m². By quantifying Rhp51, Rqh1, and MYC–six-His–Top3 foci, we observed an accumulation of foci following UV treatment of G₂ cells (Fig. 5b and d).

Treating G₂-synchronized cells with UV light leads to cell cycle arrest because of activation of the DNA damage checkpoints; thus, these cells remain in G₂ for some time before proceeding into mitosis. In order to determine whether the G₂ checkpoint was intact in rqh1-d cells, wild-type and rqh1-d cells were synchronized in G₂ by lactose gradients and were irradiated with 100 J of UV light/m². Both strains delayed mitosis for ~90 min at 29°C following irradiation, indicating an intact G₂-M checkpoint (Fig. 5c, left), but the rqh1-d strain lost viability, which correlated with the accumulation of aberrant postmitosis structures as the cells proceeded through mitosis (Fig. 5c, right).

A comparison of the temporal kinetics of Rqh1 and MYC–six-His–Top3 focus formation with the formation of Rhp51 foci (Fig. 5d) and comparison with the progression of the cells into mitosis (Fig. 5e) proved to be informative: Rqh1 foci are observed more rapidly than are Rhp51 foci (10 min after irradiation). However, the number of cells with foci increases more slowly than the number of cells exhibiting Rhp51 foci. Rhp51 foci are maximal approximately 1 h following treatment. At this point, the proportion of cells with Rqh1 foci is still increasing. There are fewer cells with MYC–six-H–Top3 foci, and fewer MYC–six-H–Top3 foci per cell (this may be an artifact of the detection), and these appear in elongated G₂ cells at later times. By the 180-min time point >50% of cells have undergone mitosis, but only elongated G₂ cells with a single nucleus (i.e., arrested, premitotic cells) were scored.

Several interesting observations arise from this analysis: we see that, in the absence of rqh1, no MYC–six-H–Top3 foci accumulate, indicating (as expected from the biochemistry) that Top3 foci are totally dependent on Rqh1. However, Rhp51 foci still appear in the absence of rqh1 but with altered kinetics (Fig. 5d). In rqh1-d cells, Rhp51 foci are increased in the absence of irradiation (unirradiated and 10-min time points) and following irradiation the proportion of cells with Rhp51 foci increases more slowly. However, by 180 min the numbers are similar to those for rqh1+d cells. Since rqh1-d cells and rqh1+d cells enter mitosis with approximately the same kinetics following irradiation, while rqh1-d cells accumulate gross postmitotic aberrations (Fig. 5c), this suggests that recombination intermediates form but may not have been correctly processed or resolved. Again, this would be consistent with recombination occurring upstream of Rqh1 and Top3.

If the formation of Rqh1 and Top3 foci was always dependent on the initiation of recombination, we would anticipate that Rqh1 and MYC–six-H–Top3 foci would not form in rhp51-d mutant cells in G₂. However, both Rqh1 and MYC–six-H–Top3 foci are seen in rhp51-d cells following UV irradiation (Fig. 5d). The Rqh1 foci were slightly reduced, but the MYC–six-H–Top3 foci are increased. In order to understand and possibly reconcile these observations, we examined the UV response pathways in which Rqh1 and Rhp51 might act. Fission yeast contains two separate excision repair pathways that act to remove UV-induced DNA damage: the canonical nucleotide excision repair (NER) pathway and the UVDE pathway (54). The UVDE pathway is restricted to S. pombe, some bacteria, and several other eukaryotic microorganisms. In S. pombe it is known that the Uve1 enzyme nicks DNA immediately 5’ of the DNA lesion (30). Following this, the nick is processed by one of two separate pathways, which involve the FEN1 homologue Rad2 and the recombination machinery (53). In addition to the physical removal of UV-induced DNA damage, mechanisms exist to bypass the lesions by translesion synthesis or recombination. These are known as damage tolerance or PRR mechanisms. In the budding yeast S. cerevisiae, PRR depends on RAD6, encoding an E2 ubiquitin-conjugating enzyme, and RAD18, encoding a RING finger protein (6). It has been previously demonstrated that the fission yeast homologue of Rad18, Rhp18, is required for the survival of UV damage in G₂ (48).

If Rqh1 and Top3 are acting in a recombination repair pathway following UV-induced DNA damage in G₂, it is likely to be part of the Uve1-dependent repair process or the damage tolerance (PRR) pathway. In the absence of the UVDE pathway (uve1-d mutant), UV-induced DNA damage is repaired by the NER pathway, which is not expected to involve recombination. Indeed, we find that, in uve1-d mutants, Rhp51 foci are reduced following irradiation of G₂ cells (Fig. 5e). In this analysis cells were only scored up to 60 min postirradiation to ensure that the results were not biased by erroneously scoring...
cells passing through mitosis into the following S phase). However, Rqh1 foci were not decreased, indicating that these can be formed independently of Uve1 and Rhp51.

We next looked at the role of the Rhp18-dependent pathway and found that Rhp51 foci were decreased in rhp18-d mutant cells following irradiation. This suggests that both the uve1-dependent and the rhp18-dependent pathways contribute to Rhp51 focus formation (Fig. 5e). Unexpectedly, however, following irradiation of a rhp18-d uve1-d double mutant, Rhp51 foci were increased compared to the foci of the respective single mutants (Fig. 5e). One possible explanation of this would be that loss of both pathways results in a deregulation of repair that leads to the accumulation of DSBs, which activates alternative recombination pathways for repair. Indeed, when we analyzed the other double mutant, uve1-d rhp1-d, we also saw an increase in Rhp51 foci compared to foci of each single mutant (Fig. 5e). Thus, while we can draw some conclusions from these data, it is not possible to use the appearance of Rhp51 foci to define the pathways in which Rqh1 is involved.

Rqh1 foci, like Rhp51 foci, were significantly decreased in an rhp18-d mutant following UV irradiation (Fig. 5e). This suggests that Rqh1 is required for Rhp18-dependent processes in G2. However, in the uve1-d rhp18-d double mutant, Rqh1 foci were reduced following irradiation. This is in contrast to the increase seen for Rhp51 foci. This suggests that the increased Rhp51 foci are independent of Rqh1. Since Rqh1 foci were also moderately decreased in the rhp1-d strain, we examined Rqh1 focus formation in an rhp18-d rhp1-d double mutant. In this case Rqh1 foci were also reduced following irradiation (Fig. 5e). Thus, Rqh1 foci result from Rqh1 functions that are downstream of both Rhp18 and Rhp51. This leads us to propose a model where Rqh1 and Top3 are required both upstream and downstream of Rhp18-dependent recombination (Fig. 6): Rqh1 and Top3 are required downstream of Rhp18 in an Rhp18-dependent pathway and are also required downstream of Rhp51. Since rqh1 and rhp1 are epistatic to each other, while neither is epistatic to rhp18 or uve1 (Fig. 5f and references 42 and 48), we have to propose that both Rqh1 and Rhp51 are required for the recombination-dependent subpathways of Rhp18-dependent repair and Uve1-dependent excision repair. This leads to the prediction that Rqh1 acts upstream of Rhp51 in the Rhp18-dependent pathway, since, if it were acting solely downstream of Rhp51, a deletion of rhp1 would abolish Rqh1 focus formation. This is consistent with the observed increase in MYC–six-H–Top3 foci in an rhp51-d strain, as they would no longer be removed by recombination in the Rhp18 pathway.

FIG. 6. Model of the roles of Rqh1 and Top3 in the repair of UV damage in S. pombe. UV damage is repaired by the classic NER and UVDE pathways. Damage can also be bypassed by translesion synthesis and recombination in PRR pathways mediated by Rhp6 and Rhp18 (homologues of budding yeast Rad6 and Rad18 proteins). Rhp51 has been reported to act in the recombination subpathway of UVDE (35) but is not epistatic to uve1 (Fig. 5f), which is consistent with it also having a role in recombinational bypass in PRR, as has been shown for Rad51 (6). Rhp51 is also required for the repair of DSBs that can occur by a variety of means, including the repair of UV lesions on opposing strands (thin arrow). Rhp51 foci are reduced in either uve1-d or rhp18-d, supporting its roles in the UVDE and Rhp18-dependent pathways. Rqh1 foci are dependent on Rhp18 and Rhp51; therefore, Rqh1 is downstream of Rhp18 in PRR and downstream of Rhp51. The requirement for Rhp51 downstream of Rqh1 in PRR (dashed line) is based on epistasis, since rqh1 and rhp51 are epistatic to each other while neither is epistatic to uve1 or rhp18.

**DISCUSSION**

Two conclusions arise from our analysis of Rqh1 and Top3: (i) using genetics and cell biology, we provide evidence that Rqh1 and Top3 function downstream of Rhp51-mediated strand invasion and downstream of an Rhp18-dependent pathway in response to UV-induced DNA damage in G2, and (ii) we identify a role for Rqh1 and, by association, Top3 in the repair of UV damage during G2.

Previously published data, discussed below, are consistent with, and strengthened by, our ordering of Rqh1-Top3 and Rhp51 functions during recombination. Our identification of a role for Rqh1-Top3 function during the response to UV-induced damage in G2, which is additional to the previously demonstrated role in DNA damage tolerance, PRR during S phase, and IR-induced DSBs, extends the range of processes in which RecQ family helicases are involved.

Characterization of Rqh1 and Top3 in S. pombe. The function of RecQ family helicases is always closely linked to Top3 and associated with recombination (26, 52). However, it remains unclear where RecQ helicases act during these complex processes. In recent years, a role for recombination in processing aberrant DNA structures that arise either spontaneously or at sites of DNA damage during DNA replication has become apparent (14, 34, 36). In E. coli, RecQ functions in a RecF-dependent process that allows the recovery of DNA replication following DNA damage. This is proposed to involve the processing of nascent DNA at blocked replication forks prior to the resumption of DNA synthesis (13). In the absence of RecQ, E. coli cells become hyperrecombinogenic, a result of the persistence of recombinogenic DNA ends. A similar role
for eukaryotic RecQ helicases in the restoration of replication forks has been proposed (27).

Both *S. pombe* and *S. cerevisiae* have a single RecQ-related helicase, Rqh1 and Sgs1, respectively. *S. pombe rqh1-d* mutant cells are highly DNA damage sensitive. In part, this results from an inability to tolerate DNA damage in S phase (42). This is consistent with the bacterial model and also with the observation that Rqh1 loss results in elevated spontaneous and induced recombination (17, 45). The relatively dramatic DNA damage sensitivity of *rqh1-d* mutants (when compared to *S. cerevisiae sgs1 null mutants*) makes *S. pombe* an attractive organism for the study of eukaryotic RecQ helicases. A further advantage of *S. pombe* is the fact that rapidly growing cells spend the majority of the cell cycle in G2, making analysis of DNA damage response in this part of the cell cycle relatively simple. To further characterize Rqh1 function, we deleted the *top3* gene. *top3* is essential in the presence of *rqh1* but becomes inessential when *rqh1* is deleted. This observation, which is consistent with reports from other groups (22, 33) published while this work was ongoing, allows access to a simple assay: we reasoned that mutants in which *top3* deletion was compatible with cell proliferation would identify situations where the DNA structures that are usually processed to lethal events by Rqh1 (in the absence of Top3) do not arise. This should help identify at which stage during recombination Rqh1 functions.

**Rqh1 acts both downstream and upstream of the RecA homologue Rhp51.** We find that *top3* loss of function does not cause inactivation in *rhp51-d, rhp54-d, rhp55-d,* or *rad22-d* (RAD52 homologue) backgrounds. Rhp51 is the *S. pombe* RecA homologue (39) and, by comparison with *E. coli* and *S. cerevisiae,* is assumed to form the nucleoprotein filaments required to initiate recombination (44). Rhp55 acts to mediate Rhp51 filament formation, and Rhp54 is thought to act later, after heteroduplex formation (44). The role of the *S. pombe* Rad52 homologue, Rad22, is ambiguous (41). It is probably involved in initiating and mediating Rhp51 filament formation. There are two straightforward explanations for the fact that, in all four of these mutants, Rqh1-dependent *top3* lethality is avoided: the first is that, in the processing of aberrant DNA structures arising spontaneously during DNA replication, Rqh1-Top3 acts after the formation of Rhp51 filaments. A second is that, in the absence of Top3, Rqh1 initiates a recombination event that is lethal. This could not occur when recombination is abolished.

In addition to using *top3-d* inviability as an assay, we have measured the survival of combinations of *rqh1, top3* and recombination mutants to UV-induced DNA damage. This provides evidence that Rqh1 and Top3 act after Rhp51 in DNA repair. Recombination mutants combined with either *top3-d* or *rqh1-d top3-d* did not show an increase in UV sensitivity, formally placing Rqh1-Top3 function within the recombination-dependent repair response (Fig. 4 and reference 42). Importantly, however, in the cases of *rhp51-d* and *rhp55-d* (in both cases recombination cannot occur due to the inability to form heteroduplex DNA), concomitant deletion of *rqh1, top3,* or both *rqh1 and top3* resulted in a rescue of *rqh1-d top3-d* sensitivity at low doses of UV-induced DNA damage in both asynchronous and G2-synchronized cells. (At higher doses, *rhp51-d* is more sensitive than *rqh1-d,* which may mask the effect.) This observation implies that, during the response to induced DNA damage, Rqh1-Top3 acts either downstream of Rhp51 (the same logic as given in hypothesis one above) or that Rqh1-Top3 acts before recombination (the same logic as given in hypothesis two).

Unlike what we found concerning response to intrinsic DNA damage (lethality), in the case of induced DNA damage, we were able to correlate the rescue data with observations based on cell biology: we find that Top3 foci are dependent on Rqh1 and that Rqh1 foci are partially dependent on both Rhp51 and Rhp18. Thus, the function of Rqh1 and Top3 is complicated, and we suggest that these data provide support for both hypotheses: i.e., that, if DNA damage is not processed by Rhp51 into recombination intermediates, it does not become an abortive substrate for Rqh1, which can act downstream. (Presumably, this damage can then be repaired by an alternative pathway, accounting for the rescue of *rqh1-d* sensitivity by concomitant *rhp51* or *rhp55* deletion.) In addition, based on epistasis analysis and the induction of foci, Rqh1 can act before Rhp51 (hypothesis two) in the Rhp18 UV response to DNA damage in G2 cells.

Because Rqh1 appears to act both upstream and downstream of recombination, depending on the DNA damage response pathway where it is functioning, we cannot precisely define a single linear pathway and thus suggest the model presented in Fig. 6 to account for these data. Furthermore, we cannot use the kinetics and dependency of focus formation to distinguish between the two hypotheses (upstream or downstream) during the processing of spontaneous lesions. However, we strongly favor a similar order of events. To extrapolate from the analysis of induced DNA damage, if the Rqh1-dependent *top3-d* lethality occurs downstream of recombination, in *top3-d* cells the Rqh1 helicase would act on recombination intermediates in such a way that cells entered mitosis with unresolved recombination structures. Thus, removing the helicase or recombination would suppress the lethality. Alternatively, the lethality of *top3-d* cells could be due to inappropriate processing by recombination proteins of DNA structures arising from Rhp18-mediated damage tolerance. This recombination event would then require Rqh1-Top3 for resolution and so would be lethal. In either case then, it would be the events downstream of recombination that are important for viability. This is supported by the fact that deletion of *rhp18,* although significantly reducing Rqh1 foci in response to UV damage, does not suppress the lethality of *top3-d* (Table 1).

**A role for Rqh1 and Top3 during recombination repair in G2.** It has been established in several systems that eukaryotic RecA homologues form foci following DNA damage (23, 32, 47). However, RecA homologues are found at replication sites during the unperturbed S phase along with (in some instances) RecQ homologues (46, 51). Because recombination proteins and RecQ family helicases are involved in DNA metabolism during replication, it is important to establish if foci visualized following UV-induced DNA damage represent persisting replication forks or if they arise de novo as part of the DNA damage repair response. Consistent with data from other organisms, Rhp51 formed clear de novo foci when G2 cells were irradiated with UV. In *S. pombe,* this correlated to the activity of the *uve1*-dependent repair pathway that is known to require recombination functions to repair UV-induced lesions.
In somewhat of a surprise to us, we observed that both Rqh1 and Top3 also formed de novo foci in a significant proportion of UV-damaged G2 cells. Rqh1 clearly acts in multiple aspects and Top3 also formed de novo foci in a significant manner. This most probably reflects an involvement in several recombination-based mechanisms used to bypass DNA damage. The observation that loss of rqh1 increases induced recombination following UV treatment probably reflects this function. Our observations that Rqh1 and Top3 form de novo foci provides evidence that Rqh1 also functions in the response to UV-induced DNA damage in G2 cells. Several reports in the literature are consistent with RecQ family proteins acting in DNA repair outside S phase; Wu et al. (51) reported a physical interaction between BLM and RAD51 in humans and Sgs1 and Rad51 in S. cerevisiae and colocalization of BLM and RAD51 following DNA damage. Bischof et al. (5) also observed BLM focus formation after DNA damage, in this case in late S-G2 cells. Caspari et al. (9) recently demonstrated that IR-induced DSBs in G2 cells resulted in the formation of Rqh1 and Top3 foci and presented genetic evidence that suggests a function for Top3 in resolving Holliday junctions (reviewed in reference 7). Our data, which correlate with both clonogenic survival analysis of G2 cells and the appearance of Rqh1 and Top3 foci in an Rhp51- and Rhp18-dependent manner, provide the direct evidence for a function of a RecQ helicase family member outside DNA replication and the first evidence that Rqh1-Top3 functions in G2 in response to UV-induced DNA damage.

**Top3 may act independently of Rqh1.** It is intriguing to know if Rqh1 always acts in association with Top3. Certainly, it appears that Top3 is required to resolve RecQ family helicase-generated DNA structures in many systems, but it is not possible to ascertain directly if Rqh1 has roles in which it acts independently of Top3. Interestingly, although *rqh1-d* mutant cells were extremely slow growing when combined with *rad22-d*, a triple mutant of *rad22-d rqh1-d top3-d* was significantly more viable (see Fig. 4b, panel i, and Table 1). This suggests that, in this case, increased lethality is due to inappropriate Top3 activity. In other words, the rescue of the growth defect of *rqh1-d rad22-d* by deletion of *top3* is evidence that Top3 functions independently of Rqh1 but not that Rqh1 functions independently of Top3. This is consistent with work by Onodera et al. (43), who reported a functional and physical interaction between Sgs1 and Top3 and an Sgs1-independent function of Top3 in DNA recombination repair (43). In all our experiments we have not been able to separate Rqh1 function from Top3 function. Biochemical analysis showed that not all Top3 was coprecipitated with Rqh1. However, we also find that the majority of Rqh1 is insoluble and that the small percentage of soluble Rqh1 in an asynchronous cell extract most likely arises from the M-G1 cells (data not shown). This precludes a thorough biochemical analysis by protein association.

**Summary.** Our data provide strong evidence that, where Rqh1 functions in a recombination-based pathway, it functions after Rhp51 filament formation. This would be consistent with models that predict that RecQ helicases help determine the correct substrate for appropriate strand invasion by reversing inappropriate heteroduplex intermediates. The fact that we also find evidence for Rqh1 acting before recombination proteins in certain circumstances may ultimately help to reconcile the differences in interpretation between work in *S. pombe* and models based on observations from *S. cerevisiae* that have suggested that RecQ helicases act before Rad51 (11).

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**REFERENCES**


Topoisomerase III is essential for accurate nuclear division in *Schizosaccharomyces pombe*. Nucleic Acids Res. 27:4050–4058.


