Role of *Schizosaccharomyces pombe* RecQ Homolog, Recombination, and Checkpoint Genes in UV Damage Tolerance

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The cellular responses to DNA damage are complex and include direct DNA repair pathways that remove the damage and indirect damage responses which allow cells to survive DNA damage that has not been, or cannot be, removed. We have identified the gene mutated in the rad12.502 strain as a *Schizosaccharomyces pombe recQ* homolog. The same gene (designated rqh1) is also mutated in the *hus2.22* mutant. We show that Rqh1 is involved in a DNA damage survival mechanism which prevents cell death when UV-induced DNA damage cannot be removed. This pathway also requires the correct functioning of the recombination machinery and the six checkpoint *rad* gene products plus the Cds1 kinase. Our data suggest that Rqh1 operates during S phase as part of a mechanism which prevents DNA damage causing cell lethality. This process may involve the bypass of DNA damage sites by the replication fork. Finally, in contrast with the reported literature, we do not find that rqh1 (*rad12*) mutant cells are defective in UV dimer endonuclease activity.

Cellular responses to DNA damage can be defined as either direct DNA repair responses, which result in the removal and repair of the damage, or indirect responses (such as the G_2 damage checkpoint) which do not directly repair or remove damage but allow the cell to survive such damage (8). The relationships between the various DNA damage responses are complex and difficult to study in mammalian cells. Since there is good evidence that most of the damage response pathways are highly structurally and functionally conserved (5, 13, 28, 33, 51, 53), we are studying these pathways in the fission yeast model system.

In fission yeast *Schizosaccharomyces pombe*, UV-induced DNA damage is removed by one of two pathways, either the conserved nucleotide excision repair (NER) pathway (10) or a novel pathway initiated by the UV dimer endonuclease (UVDE) (63). In addition to DNA repair, *S. pombe* has several damage responses that do not result in the direct removal of photoproducts but allow the cells to survive DNA damage more effectively. These are controlled by the checkpoint pathway(s) and include G_2 delay (1, 49), transcriptional activation (16, 57), and the control of progress through S phase following DNA damage or exposure to hydroxyurea (2, 15).

In *S. pombe*, the DNA structure checkpoint responses all require a group of six proteins commonly referred to as the checkpoint Rad proteins (7). Mutations in any one of these result in very similar phenotypes, including sensitivity both to DNA damage and to S-phase arrest by hydroxyurea. It has been proposed that the checkpoint Rad proteins form a complex that is capable of monitoring various different aspects of DNA metabolism, including DNA damage/repair and DNA replication (7).

In response to DNA damage, the checkpoint Rad complex activates two downstream kinases, Chk1 and Cds1. Chk1 activation is required for G_2 arrest in response to DNA damage

and can be monitored by a phosphorylation event (58). Cds1 activation is S-phase specific and does not occur in G_1 or G_2 (34). As anticipated, *cds1* deletion cells are sensitive to DNA damage but do not exhibit a defect in mitotic arrest following DNA damage. Cds1 defines a checkpoint-related response specific to S phase. An interesting allele of one of the checkpoint *rad* genes, *rad26.T12*, that is defective specifically in the Cds1 response but not in the Chk1 response has been described (2, 34). *rad26.T12* cells are sensitive to DNA damage during S phase, further indicating that the Cds1-dependent response is important specifically during S phase.

At the beginning of this work, we identify the S. pombe RecQ homolog, Rqh1, by virtue of its involvement in the DNA damage response. RecQ in Escherichia coli is a DNA helicase involved in the RecF recombination pathway (38). RecQ homologs have been identified in eukaryotes, but these have not previously been shown to be involved in the response to DNA damage. In Saccharomyces cerevisiae, a single RecQ homolog, Sgs1, has been identified. This protein appears to associate with DNA topoisomerases (20, 61), although the precise function is not known. There is no radiation sensitivity associated with loss of Sgs1 function (60). In human cells, three RecQrelated proteins, RecQL, WRN, and BLM, have been identified (14, 48, 64). Two of these are associated with genetic diseases: the WRN gene is defective in patients with the premature aging disorder Werner's syndrome (64), and the BLM gene is defective in those with Bloom's syndrome (14), who suffer from sun sensitivity and developmental abnormalities (21)

Cells from Bloom's syndrome patients are not particularly sensitive to DNA-damaging agents and have not been reported to manifest checkpoint deficiencies. However, these patients exhibit a 300-fold increase in neoplasia and have elevated in vivo mutant frequencies in their lymphocytes (31). The most dramatic cellular phenotypes associated with Bloom's syndrome are elevated levels of sister chromatid exchange (11) and a high frequency of chromosome abnormalities (22). These also appear to be associated with a slowdown of the progression of the replication fork (23, 27). While a defect in ligase

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function has been reported for Bloom's syndrome cells (12, 62), the genetic defect is not associated with a DNA ligase.

The work reported here links the phenotypes associated with mutation of the *S. pombe* RecQ homolog (Rqh1) to the correct operation of the checkpoint proteins and Cds1. Our work suggests that RecQ functions in a checkpoint protein-dependent DNA damage response during S phase. This response allows cells to survive damage through a process which also involves the recombination repair proteins. We discuss these results in the context of the Bloom's syndrome phenotypes and suggest that *S. pombe* may prove to be a useful organism with which to study the RecQ helicase family in eukaryotes and its involvement in DNA metabolism.

MATERIALS AND METHODS

Strains, media, and physiological tests. All strains except that with the null mutation at the *rqh1* locus, which was a gift from T. Enoch, have been described previously (see Table 2) or are derived from such strains by classical genetic methodologies (25). Media for cell culture have been described elsewhere (39). UV and gamma survival measurements were performed as previously described (2, 43).

Cosmid clones and P1 clones. Cosmid and P1 clones were supplied by the resource center/primary database of the German human genome project (Max Planck Institute for Molecular Genetics) and are described in reference 29. To tag the cosmids and P1 phage with a selectable marker, the transposon tags developed by Morgan et al. (40) were used. Briefly, the *ars1 ura4* transposon was introduced at random sites by transformation of the cosmid and P1 clones into the *E. coli* donor strain MH1831, followed by transformed into *S. pombe* by using standard techniques (39). To avoid the possibility that the transposon inserted into the *rad12* locus and inactivated the gene, DNA was prepared from a mixed population of tagged DNAs for each cosmid or P1 phage. Survival was assayed by comparative strip tests (9).

Sequencing of rqh1.r12 (rad12.502) and rqh1.h2 (hus2.22). The entire open reading frame (ORF) of the rqh1 gene was amplified in three overlapping PCRs and cloned into pGem plasmids for the preparation of single-stranded DNA. Two independent isolates from each mutant were sequenced side by side and compared for mutations. In each case, a single change was seen in both isolates. These changes were subsequently confirmed by cloning of a number of further isolates of the relevant fragments followed by sequencing with the relevant primer.

UVDE activity assays. UVDE (also known as S. pombe dimer endonuclease) activity was measured in extracts as described previously (54). Two hundred milliliters of culture at a cell density of 107/ml was split in two. Extracts were prepared directly from one sample, while the other was irradiated with UV light (100 J/m²) in water in a Stratalinker. Following a 1-h recovery in fresh medium, cells were broken by dismembranation, using glass beads in 20 mM Tris-HCl (pH 7.9)–1 mM EDTA–10 mM MgCl₂–0.3 M ammonium sulfate, 0.5 mM dithiothreitol (DTT)-10% glycerol-1 mM phenylmethylsulfonyl fluoride. Supernatants were spun at 40,000 rpm for 1 h at 4°C, dialyzed three times against 20 mM HEPES (pH 7.6)-10 mM MgSO₄-10 mM EGTA-5 mM DTT-20% glycerol-1 mM phenylmethylsulfonyl fluoride, and frozen at -70°C in aliquots. Reactions were carried out with 0 to 30 μ g of protein in 25 μ l in 9 mM HEPES-14 mM KCl-1.5 mM MgCl₂-0.08 mM EDTA-0.7% glycerol (pH 7.8)-0.9 mM DTT-0.02 mM deoxynucleoside triphosphates with 100 ng of DNA (irradiated or unirradiated) for 1 h at 37°C. Samples were then treated with RNase and proteinase K (9) and analyzed by electrophoresis.

Cdc10 synchronization. Double mutants carrying *cdc10.V50* and *rqh1.d* or *cds1.d* were created. Logarithmic cultures of *cdc10.V50*, *cdc10.V50 cds1.d*, and *cdc10.V50 rqh1.d* cells were shifted to 37° C for 3 h to accumulate cells at the G₁/S boundary. Cells were then shifted to 26° C and irradiated with 250 Gy 35 min later. Samples were taken for analysis of Cds1 kinase activity following 40 min of recovery. At the time of irradiation, appropriately diluted samples (10^{4} cells per ml) of *cdc10.V50* and *cdc10.V50 rqh1.d* cells were divided in two, and one half of each sample was irradiated with 100 Gy. Irradiated and mock-irradiated samples were in S phase at the time of irradiation. At 140 min, before the peak in septation and after S phase (G₂ cells), similar dilutions were divided in two, and one half was irradiated to assay survival in G₂.

Chk1 phosphorylation and Cds1 kinase assays. Chk1 phosphorylation in rqh1 null cells was monitored by mobility shift after Western blotting of cell extracts made from a strain containing an integrated triple-hemagglutinin-tagged *chk1* locus (58). Cds1 activity was monitored by immunoprecipitation-kinase (IP-kinase) assays using anti-Cds1 serum (34). Extracts were prepared from cells washed in phosphate-buffered saline and then lysis buffer [50 mM Tris (pH 7.5), 80 mM β -glycerophosphate, 250 mM NaCl, 15 mM nitrophenylphosphate, 50 mM NaF, 5 mM EDTA, 1 mM DTT, 0.1% Nonidet P-40, 4-(2-aminoethyl)-benzenesulfonyl fluoride (AEBSF), leupeptin, aprotinin, pepstatin, bestatin,

E-64 (the latter six at 10 µg/ml)]. Cells were disrupted with glass beads (BDH) in a Ribolyser (Hybaid) for 20 s. For Cds IP-kinase assays, 1 mg of total protein in 500 µl of lysis buffer was incubated with affinity-purified Cds1 antibody at a dilution of 1:250 at 4°C for 2 h. Immunocomplexes were collected with protein A-agarose (Sigma), washed three times in lysis buffer, and then washed three times in kinase buffer (10 mM HEPES [pH 7.5], 75 mM KCl, 5 mM MgCl₂, 0.5 mM EDTA, 1 mM DTT). Twenty microliters (50% slurry) of bead pellet was incubated with 10 μ l of 2× kinase buffer-5 μ Ci of [γ -³²P]ATP (ICN)-1 μ l of 2 mM ATP-5 µl of myelin basic protein (MBP; 1-mg/ml stock) at 30°C for 15 min. The reaction was stopped by the addition of 20 μ l of 2× sodium dodecyl sulfate (SDS) sample buffer. After boiling for 3 min, samples were run on 15% polyacrylamide gels, fixed in 40% methanol-10% acetic acid, and dried before exposure to film (Hyperfilm; Amersham). For immunodetection of Chk1, 100 µg of total protein was boiled in SDS sample buffer and loaded onto an SDS-10% polyacrylamide gel. Proteins were transferred to nitrocellulose; nonspecific sites were blocked with BLOTTO (phosphate-buffered saline, 1% fat-free milk powder, 0.05% Tween 20) and incubated in BLOTTO plus antihemagglutinin monoclonal antibody (1:1,000 dilution; BaBco). After being washed in BLOTTO, filters were incubated with peroxidase-conjugated secondary antibody (1:5,000 dilution; DAKO). Chemiluminescence detection of horseradish peroxidase-conjugated secondary antibodies was carried out by mixing 10 ml of 100 mM Tris (pH 8.5)-5.4 mM H₂O₂ with 25 µl of 90 mM p-coumaric acid (Sigma) and 50 µl of 250 mM luminol (Fluka). Blots were incubated with this mixture for 60 s before excess solution was removed and the filter was exposed to Hyperfilm-MP (Amersham).

RESULTS

Identification of a candidate ORF for rad12. rad12.502 strain is the only S. pombe radiation-sensitive mutant of 24 (each defining a separate complementation group) from which we and others have failed to clone a gene by complementation of the radiation sensitivity (4). We therefore mapped the genetic location of rad12.502 relative to known markers along the chromosomes. rad12.502 mapped to within 7 centimorgans of rad4 on the top arm of chromosome I, between rad15 and rad4. Correlation with the physical map (29) allowed us to identify cosmids 2g11, 26b6, 27a5, 16h3, and 11a10 and two P1 clones, 32e5 and 34c8, as potentially encoding the *rad12* gene. These cosmids and phage were marked with the selectable marker $ura4^+$ and with the autonomously replicating sequence ars1 by transposon tagging (40). The tagged constructs were prepared and transformed into rad12.502 ura4.D18 mutant cells. One cosmid, 2g11, was able to restore radiation resistance to rad12.502 mutants. This cosmid has been sequenced by the S. pombe genome project, and the sequence data is available from the Sanger Center (http://www.sanger.ac.uk/pombe/pombe.htm1). Cosmid 2g11 contains an ORF with homology to genes encoding the RecQ family of helicases and to the human BLM gene (Fig. 1). This ORF has previously been identified by complementation of the hus2.22 mutant, which is defective in the recovery from treatment with the DNA synthesis inhibitor hydroxyurea and exhibits radiation sensitivity (15, 55).

Identification of the mutations in *rad12.502* and *hus2.22*. As anticipated, crosses between *rad12.502* and *hus2.22* mutants yielded no wild-type segregants, confirming allelism. Since *rad12.502* and *hus2.22* mutants have distinct phenotypes, we cloned by PCR amplification the entire *recQ*-like ORF from both *rad12.502* and *hus2.22* cells in order to identify the causative mutations. Single base changes were found for both *rad12.502* and *hus2.22*-derived DNAs, which were present in several independent PCR isolates.

The *hus2.22* mutation introduces a stop codon at amino acid position 790, effectively truncating the remaining region of 538 amino acids, which contains the last two conserved helicase motifs (Fig. 1). *hus2.22* is therefore likely to have a null phenotype. This has been confirmed by deletion of the gene (55). The *rad12.502* mutation changes a threonine to an isoleucine at amino acid position 543, within the highly conserved putative ATP binding site (Fig. 1). *rad12.502* clearly does not confer a null phenotype, since *rad12.502* cells are not ionizing radia-



FIG. 1. Relationship between Rqh1, Sgs1, Blm, Wrn, RecQL, and RecQ. (A) Phylogenetic tree showing the relatedness of the six helicases. The region containing the helicase motifs (approximately 340 amino acids) was aligned by using DNAstar software. (B) Bar diagram indicating relatedness and structure of the six proteins. The black bar indicates the helicase region used to generate the phylogenetic tree. This region contains the six domains typical of many helicases. The shaded box indicates a short region with additional homology found in four of the six proteins. The unshaded regions show the extensions for each protein N and C terminal to the conserved regions. These show no homology. (C) Alignments of the two helicase domains containing mutations in *rqh1.r12 (rad12.502)* and *rqh1.h2 (hus2.22)*, respectively. *rqh1.r12* cells contain a mutation that changes a conserved threonine to an isoleucine in helicase domain I, within the putative nucleotide binding site. rqh1.h2 cells contain a mutation which introduces a stop codon at the start of helicase domain V. Alignments were generated by using DNAstar software. Numbers indicate amino acid position on the Rqh1 protein. Changes are indicated in circles above the sequence.

tion sensitive, a phenotype associated with hus2.22 cells (15), and these cells are not as sensitive to UV as hus2.22 or the deletion mutant cells.

To reduce confusion, we have renamed the *hus2* locus *rqh1* (*recQ* homolog), with the following allele designations: *rqh1.h2* (*hus2.22*) and *rqh1.r12* (*rad12.502*).

The rqh1-dependent UV response requires the recombination proteins. rqh1 null mutants are sensitive to S-phase arrest by hydroxyurea, and the original rqh1.h2 and rqh1.r12 alleles have been reported to confer sensitivity to UV light. We have investigated in more detail the relationship between rqh1 and the known DNA repair mechanisms. Two pathways that remove UV-induced DNA damage have been identified in S. pombe (8): NER pathway and the UVDE pathway, which removes dimers in a parallel pathway to NER. Interestingly, recombination repair mutations in S. pombe, such as rhp51 and *rhp54* (the equivalents of *RAD51* and *RAD54*, respectively, of *S. cerevisiae*), also confer significant sensitivity to UV light, demonstrating a role for recombination repair in the response to this form of DNA damage (41, 42).

Double mutants carrying an rqh1 mutation and either a NER mutation (such as rad13 [Fig. 2A]) or a mutation in the structural gene for the UVDE enzyme (designated *uvde* [Fig. 2B]) show significantly more sensitivity to UV than the equivalent single mutants. This result indicates that Rqh1 acts in a pathway separate from either of these repair mechanisms. In contrast, double mutants between rqh1 and representatives of the recombination repair pathway (such as rhp51 [Fig. 2C]) did not show greater sensitivity to UV light than the respective single mutants, which suggests that Rqh1 and recombination repair proteins are involved in a common response to UV radiation damage.



FIG. 2. Survival analysis after UV irradiation. (A) Rqh1 does not act in the NER pathway. wt, wild-type cells; r13, *rad13.d* mutant cells; rqh1, *rqh1.d* mutant cells; rrl1 rqh1.*d* mutant cells; rqh1, *rqh1.d* mutant cells; rqh1 uvde, *rqh1.d* uvde double-null mutant cells. (C) Rqh1 function requires recombination repair proteins. wt, wild-type cells; rqh1, *rqh1.d* mutant cells; 51, *rhp51.d* null mutant cells; rqh1 51, *rqh1.d* mutant cells.



FIG. 3. Relationship between UVDE activity and rqh1. (A) UVDE activity was measured in wild-type cells, uvde null mutant cells, rqh1.r12 (rad12.502) cells, rqh1.d null mutant cells, rad3 null mutant cells, and rad9 null mutant cells (lanes 1 to 6, respectively). The assay measures conversion of a UV damage-containing supercoiled plasmid to the nicked circular form (54). The responsible nuclease activity is dependent on the UVDE activity, since extracts prepared from uvde null mutant cells (lanes 2, induced and uninduced) do not show this activity. For the strains corresponding to lanes 1 to 6, extracts were prepared as described in Materials and Methods either with (induced) or without (uninduced) prior treatment of the cells with UV light. Following incubation of these extracts with equal amounts of UV-irradiated plasmid, DNA was assayed by electrophoresis for conversion into nicked circle form. Two control lanes, UV damage-containing plasmid without any extract (plasmid) and damage-free plasmid in wild-type cell extract (No uv), establish that the activity is dependent on cell extract and plasmid damage. (B) Quantification of the conversion from supercoiled to nicked plasmid compared to control.

UVDE and *rgh1* **mutants.** *rgh1* has previously been linked to the UVDE-dependent second excision repair pathway of S. pombe (19), and the rqh1.r12 (rad12.502) mutant has been reported (19) to be deficient in UVDE activity. However, the structural gene for the UVDE enzyme has recently been identified and is independent of the rgh1 loci (63). Therefore, the published data suggest that Rqh1 has a role in regulating the activity of the UVDE enzyme. To verify this proposition, and to determine the effect of a null mutation of rgh1 on UVDE activity, we prepared cell extracts from logarithmically growing rgh1.r12 and rgh1.d (null) cells. As controls, we also included uvde (null), rad3.d, rad9.d, and wild-type cells. Extracts were prepared both before and 1 h after UV irradiation. Examination of UVDE activity (Fig. 3) by using a plasmid nicking assay indicates that significant activity can be detected in all cells except uvde null mutants and that in all of these cases, significant induction is seen following irradiation. While this nicking assay is not quantitative, these results are reproducible and clearly show that, contrary to the reported literature, UVDE activity does not require Rqh1 function in logarithmically growing S. pombe cells.

Rqh1 is required to tolerate DNA damage that is not removed by NER or the UVDE pathway. When both the UVDE and NER pathways are abolished in *S. pombe*, photoproducts are not repaired at all (63). Rqh1 does not function in either of these pathways, which suggests that Rqh1 is involved in tolerating rather than removing DNA damage from cells. In support of this possibility, we find that *uvde rad13* (NER) *rqh1* triple mutants are more sensitive to UV than the *uvde rad13* double mutant (Fig. 4A). Since all excision activity is lost in the *uvde rad13* double mutant (63), the increased sensitivity of the triple mutant argues for an additional role for Rqh1 independent of direct damage removal.

Rqh1 functions in a pathway that also involves the checkpoint Rad proteins. The S. pombe checkpoint rad mutants rad1, rad3, rad9, rad17, rad26, and hus1 are all very sensitive to UV and ionizing radiation and to exposure to hydroxyurea (8). We have verified (data not shown) that these six mutants define a single epistasis group. We have previously established that the radiation sensitivity of the checkpoint rad mutants results from at least two identifiable defects: the inability to prevent passage through mitosis when the DNA is damaged (dependent on the Chk1 kinase) and a second defect (in a pathway that requires activation of the Cds1 kinase) which results in cell death during S phase when DNA is damaged or S phase arrested with hydroxyurea (2, 15, 34). Within the checkpoint Rad protein-dependent damage response, these two defects can be genetically defined: mutants defective in the Chk1 kinase are defective specifically in the mitotic arrest pathway (2, 59). In contrast, mutants defective in Cds1 kinase or the specific checkpoint rad allele rad26.T12 have normal mitotic arrest following activation of the checkpoint rad pathway (34) but are specifically sensitive to DNA damage during S phase (2).

To address the relationship between *rqh1* and the checkpoint genes within the DNA damage response, we constructed double mutants carrying *rqh1.r12* or *rqh1.d* and each of the checkpoint *rad* mutants. We have compared the UV sensitivities of the double mutants and the respective single mutants. We find that, for example, the *rqh1.d rad17.d* double mutant is no more sensitive than the *rad17.d* single mutant. This finding indicates that Rqh1 acts in a pathway that requires the correct operation of the checkpoint Rad group of proteins (Fig. 4B).



FIG. 4. Rqh1 and checkpoint proteins are required to survive unrepaired damage. (A) Rqh1 is required to survive damage in strains which cannot repair. wt, wild-type cells; uvde, *uvde* null mutant; r13, strain with null mutation of the *rad13* gene, required for NER; rqh1, *rqh1.d* null mutant cells. Double and triple mutants are also indicated (e.g., rqh1 r13 uvde is an *rqh1 rad13 uvde* triple-null mutant). (B) Epistasis between *rqh1* and checkpoint mutants. wt, wild-type cells; rqh1, *rqh1.d* mutant cells; rqh1 r17, *rqh1.d* rad17 double-mutant cells. (C) Rqh1 is specific to the checkpoint metant cells; r26.T12, *rad26 T12* mutant cells; rqh1, *rqh1.d* mutant cells; r26.T12, *rad26 T12* double-mutant cells. (D) Rqh1 is not in the same pathways as Chk1. wt, wild-type cells; rqh1, *rqh1.d* nutant cells; r26, *rad26* null mutant cells; rqh1, *rqh1.d*, *rqh1.d*,



FIG. 5. Chk1 and Cds1 are activated by DNA damage in rqh1 null cells. (A) The phosphorylation status of Chk1 in wild-type and rqh1.d mutant cells 1 h after irradiation (Irrad.) (+) or mock irradiation (-). Chk1 migrates as a single band in unirradiated cells of approximately 56 kDa. After irradiation it becomes phosphorylated, which causes an apparent increase in molecular weight. (B) Cds1 kinase activity is not dependent on Rgh1. Cells were synchronized by temperature shift in a cdc10.V50 background for 3 h and released into S phase before irradiation (Irrad.) (+) or mock irradiation (-). Immunoprecipitation with anti-Cds1 antibodies followed by a kinase assay against MBP measures the activity of Cds1. cdc10.V50 cells, irradiated 35 min after release from 37 to 26°C, activate Cds1 following irradiation. IP-kinase assays from cdc10.V50 cds1.d (c10 cds1) double-mutant cells do not show significant activity against MBP. Results of IP-kinase assays for cdc10.V50 rqh1.d (c10 rqh) double-mutant cells are identical to those for cdc10.V50 single-mutant cells. (C) Clonogenic survival of the same cultures of rqh1⁺ (wild-type [wt]) and rqh.d (rqh) cells following irradiation in S phase (35 min after release from cdc10 arrest) or in G2 (150 min after release from cdc10 arrest).

Since the checkpoint Rad proteins are required for at least two distinct damage responses, Chk1-dependent G₂ arrest and the Cds1-dependent recovery pathway, we constructed rqh1 chk1, rgh1 cds1, and rgh1 rad26.T12 double mutants and compared their sensitivities to those of the respective single mutants and the rad26 null phenotype strain. Combining the rgh1 mutation (which does not confer a mitotic arrest defect) with the chk1.d mutation yields a strain more sensitive than the chk1.d or rqh1.d single mutant. This sensitivity approaches that of the checkpoint rad null mutants such as rad3.d and rad26.d (Fig. 4D). Strains which combine the *rqh1* mutation with either the rad26.T12 or cds1 mutation did not show additional sensitivity, being no more sensitive than the most sensitive single mutants (Fig. 4C and data not shown). The classical interpretation of epistasis data in DNA repair studies is as follows: mutations which, when combined, do not lead to increased radiation sensitivity compared to the most sensitive single mutation act in the same pathway. Mutations which do increase radiation sensitivity when combined act in different pathways. Applying this interpretation, our data show that Rqh1 functions in the Cds1-mediated checkpoint rad-dependent recovery pathway but is not required for Chk1-mediated mitotic arrest.

To address whether Rqh1 functions upstream or downstream of Cds1, and to verify that the Chk1 response is not affected by rgh1 deletion, we assayed the ability of the rgh1 null mutant to activate Cds1 and Chk1 kinases in response to DNA damage. In rqh1.d cells, the Chk1 kinase is phosphorylated to an extent similar to that seen in $rqh1^+$ cells (Fig. 5A), which is consistent with our genetic analysis of rqh1. The activation of Cds1 by DNA damage is specific to S phase (34). Unfortunately, rqh1.d cells are not uniform in size at mitosis. Although this defect is subtle (data not shown), it prevents us from synchronizing rgh1 cells by elutriation. To examine Cds1 activation during S phase, we therefore synchronized cells by a cdc10 arrest-and-release strategy. Cds1 is activated to similar extents in $rqh1^+$ and rqh1.d cells after irradiation during S phase (Fig. 5B). Furthermore, while wild-type cells were approximately 10-fold more sensitive to irradiation in S phase than in G₂, rgh1.d mutant cells were approximately 50 times

more sensitive in S phase than in G_2 (Fig. 5C). Together with the genetical analysis, these data indicate that Rqh1 is not required to generate or propagate the checkpoint-dependent signal and formally place Rqh1 downstream of the checkpoint Rad proteins and Cds1.

Rgh1 is required for viability in cells with defects in DNA synthesis. While studying the genetics of the rgh1.r12 allele, we observed that correct Rqh1 function is required for viability in cells with defects in DNA synthesis that affected chain elongation. We had constructed double mutants containing rgh1.r12 and a selection of cdc mutations. Table 1 shows the detailed results of these analyses. In brief, the rgh1.r12 mutation did not reduce the growth rate or restrictive temperature for cell elongation and inviability when combined with mutations that are involved in the initiation of S phase, cdc18, cdc19, and cdc21 (17, 30, 37). However, the rgh1.r12 mutation did reduce both the growth rate and the restrictive temperature for cell elongation and inviability in mutants of DNA polymerase subunits, DNA polymerase-associated proteins and ligase, cdc1, cdc6, cdc17, cdc20, and cdc27 and showed a synthetic interaction with rad2 which affects DNA chain elongation (18, 36, 44-46). For the remaining *cdc* mutations tested, *cdc10* (G_1 arrest [3]) and cdc25 (G₂ arrest [50]), rgh1.r12 did not have any effect.

DISCUSSION

The rgh1.r12 (rad12.502) mutant is sensitive to UV radiation but is not significantly sensitive to ionizing radiation (47). Efforts to clone the gene mutated in rgh1.r12 cells by complementation of the radiation sensitivity have not been successful (4). By physical mapping, we have identified the mutation responsible for the rqh1.r12 phenotype as a single base change that results in substitution of a threonine residue for isoleucine within the conserved nucleotide binding site of a RecQ homolog. This same ORF is mutated (by a stop codon within the conserved helicase region) in the rgh1.h2 (hus2.22) mutant, which confers both UV and ionizing radiation sensitivity and is required for recovery from hydroxyurea treatment (15). rqh1 null mutants show a phenotype indistinguishable from that of rgh1.h2 mutants (55). The rgh1.r12 mutation separates an aspect of the UV response from other functions of the Rgh1 protein. The helicase activity of the S. cerevisiae homolog of Rqh1, Sgs1, is not required for all of its functions (35), suggesting that biochemical helicase activity is not the only role for this class of proteins. In this context, it is interesting that the rgh1.r12 mutation is within the putative ATP binding motif of

TABLE 1. Summary of cdc-rqh1.r12 interactions

cdc mutation	Function	Interaction	Comment
cdc18	Initiation	No	
cdc19	Initiation (MCM ^a)	No	
cdc21	Initiation (MCM)	No	
cdc1	Polymerase subunit	Yes ^b	Slow growing at 27°C
cdc6	Polymerase subunit	Yes ^b	Slow growing at 27°C
cdc20	Polymerase subunit	Yes ^b	0 0
cdc27	Coprecipitates with Cdc6	Yes ^b	
cdc17	DNA ligase	Yes ^b	Slow growing at 27°C
cdc23	Late S	Yes ^b	0 0
cdc25	G ₂	No	
rad2.d	FEN1/MF-1	Yes	Slow growing at 27°C

^a MCM, minichromosome maintenance.

^b Mutant cannot form colonies at the semipermissive temperature, at which the *cdc* single mutant can form colonies. These double mutants also showed an increased level of cell death during short incubations at the restrictive temperature but do not show a cut (premature mitosis) phenotype.



FIG. 6. A model for the UV response in S. pombe. UV light causes mainly 6-4 and cyclobutane dimers. This damage is repaired either by the classical NER pathway or by a novel repair pathway initiated by UVDE. The processing of sites incised by UVDE appears to follow one of two pathways. The first requires the rad2 and rad18 genes. The second is not characterized (?) but may require recombination functions. In addition to the removal of DNA damage, S. pombe responds to irradiation by activating DNA damage checkpoint genes. This results in several responses, including the activation of the Chk1 kinase (which in turn causes G2 arrest) and activation of Cds1 during S phase. Rqh1 function in the DNA damage response is dependent on the activation of the Cds1-dependent DNA damage tolerance pathway. This pathway also involves the products of the rad18 and recombination repair genes. However, unlike the direct repair pathways (top three arrows), this pathway does not result in the removal of DNA lesions. We speculate that the Rqh1-dependent pathway may involve bypass synthesis, allowing cells to replicate damaged templates. The fact that Rqh1 is required to prevent increased recombination in cells arrested in S phase by hydroxyurea is consistent with a role for Rqh1 in coordinating S phase and recombination at sites of DNA damage.

Rqh1, and it will be interesting to correlate the phenotypes with the biochemical properties of the mutant proteins.

rqh1.r12 and *rqh1* deletion mutants have UVDE activity. Freyer et al. have reported that *rad12* (*rqh1*) defines the second excision repair pathway in *S. pombe*, which is dependent on the activity of the UVDE enzyme (19). However, the *UVDE* gene is distinct from *rqh1* (63). The published data thus indicate that *rqh1* regulates UVDE activity. Our work does not agree with the published report, since we see essentially similar levels of UVDE activity in the *rqh1.r12 rqh1* null mutant and wild-type cell extracts. Furthermore, we do not see a requirement for other aspects of the checkpoint mechanism for UVDE activity since *rad3* and *rad9* mutant cells also exhibit UVDE-dependent nuclease activity. Since the UVDE enzyme has now been cloned (56), it should be possible to study any subtle relationship between Rqh1, checkpoint proteins, and UVDE activity in more detail.

A model for UV-induced damage repair and Rqh1-dependent damage tolerance in *S. pombe*. The radiation-sensitive phenotype associated with the rqh1 mutants is a clear phenotypic marker that allows us to delineate the relationship between the different DNA damage responses and a homolog of the *BLM* and *recQ* genes. We have integrated all of our observations and those reported previously into a model that describes the response to UV-induced DNA damage in fission yeast (Fig. 6). The data on which the model is based are presented in Table 2.

UV light produces primarily cyclobutane pyrimidine dimers and 6-4 photoproducts. These can be repaired or tolerated by a number of defined pathways, as follows. The *rad16*, *rad13*, *rad15*, and *swi10* gene products act to excise photoproducts by the classical NER mechanism. Damage not repaired by NER can be repaired by the UVDE pathway. Once a nick is intro-

TABLE 2. Summary of UV sensitivity dataused to construct model^a

Strain genotype	Sensitivity compared to single mutants, comment	Refer- ence
rqh1.d UVDE	More sensitive	
rqh1.r12 rad13.d	More sensitive	
rad13.d UVDE	More sensitive, no excision of	
nah 1 n12 nad 12 d	UV photoproducts	
UVDF	double mutants	
rah1.d UVDE	More sensitive	
rgh1.d rad13.d	More sensitive	
rqh1.d rad13.d	More sensitive than relevant	
UVDE	double mutants	
rad1.d rad3.d	No additional sensitivity, epistatic	
rad1.d rad9.d	No additional sensitivity, epistatic	
rad1.d rad17.d	No additional sensitivity, epistatic	
raa1.a raa20.a wad1 d huw1 d	No additional sensitivity, epistatic	
rad3 d rad9 d	No additional sensitivity, epistatic	
rad3 d rad17 d	No additional sensitivity, epistatic	
rad3.d rad26.d	No additional sensitivity, epistatic	
rad3.d hus1.d	No additional sensitivity, epistatic	
rad9.d rad17.d	No additional sensitivity, epistatic	
rad9.d rad26.d	No additional sensitivity, epistatic	
rad9.d hus1.d	No additional sensitivity, epistatic	
rad17.d rad26.d	No additional sensitivity, epistatic	
rad17.d hus1.d	No additional sensitivity, epistatic	
raa20.a nus1.a rad1 d rah1 r12	No additional sensitivity, epistatic	
ruu1.u rqn1.r12	<i>rad1.d</i> strain	
rad3.d rqh1.r12	No additional sensitivity, epistatic	
rad9.d rqh1.r12	No additional sensitivity, epistatic	
rad17.d rqh1.r12	<i>rad17.d</i> strain	
rad26.d rqh1.r12	No additional sensitivity, epistatic	
hus1.d rqh1.r12	Very slightly more sensitive than hus1.d strain	
rad1.d rqh1.d	Viable, very slightly more sensitive than <i>rad1.d</i> strain	
rad3.d rqh1.d	Synthetic lethality	
rad9.d rqh1.d	Viable, no additional sensitivity	
rad17.d rqh1.d	Viable, very slightly more sensitive than <i>rad17.d</i> strain	
rad26.d rqh1.d	Synthetic lethality	
hus1.d rqh1.d	Viable, very slightly more sensitive than <i>hus1.d</i> strain	
rad26.T12 rqh1.r12	No additional sensitivity, epistatic	
chk1.d rqh1.r12	More sensitive, similar to rad26.d strain	
rad26.T12 rqh1.d	No additional sensitivity, epistatic	
chk1.d rqh1.d	More sensitive	
chk1.d rad26.T12	More sensitive, similar to <i>rad26.d</i> strain	2
rnp51.a rqn1.r12	No additional sensitivity, epistatic	
mp54.a rqn1.r12 rhp51 d rah1 d	No additional sensitivity, epistatic	
rhp51.a rqn1.a rhp54.d rah1.d	No additional sensitivity, epistatic	
rad18.X rah1.r12	No additional sensitivity, epistatic	
rhp51.d UVDE	More sensitive	
rad18.X UVDE	More sensitive	
rad18.X rad2.d	No additional sensitivity, epistatic	32
rqh1.r12 rad2.d	More sensitive	
rhp51.d rad13.d	More sensitive	
rad9.d rad13.d	More sensitive	24
rua1/.a rad13.d rad18 V rad12 d	More sensitive	24
rado d LIVDE	More sensitive	32
	More consitive	

^{*a*} Where data have been previously published, a reference is given; otherwise, data were generated during this study. In classical epistasis analysis, when double mutants are not more sensitive than the most sensitive single mutant, the relevant proteins are thought to operate in the same damage response. When sensitivity is increased, some aspect of one protein must act in a response that does not involve the second protein.

duced by the UVDE enzyme (6), it is processed in one of two ways. One is not defined (Fig. 6) but may involve recombination proteins, while the other involves further processing of the lesion by Rad2 (the FEN1 homolog) and subsequent repair by a Rad18-dependent mechanism.

In addition to repair, UV damage causes the activation of the checkpoint proteins which mediate cell cycle arrest. The checkpoint proteins activate kinases which define subpathways: a Chk1-dependent mitotic arrest pathway and a Cds1-dependent pathway that results in the survival of DNA damage during S phase. This Cds1-dependent response requires both Rqh1 and recombination repair functions. The ability to survive small amounts of DNA damage that cannot be or have not been removed is most clearly seen when one compares the effects of loss of Rqh1 function in cells that cannot remove UV photoproducts (Fig. 4A).

It is interesting to speculate that this S-phase-specific survival pathway may be related to the bypass of damaged sites by the replication apparatus, a process that is known to require recombination functions in *E. coli*. The relationship between Rqh1-dependent damage survival and recombination is intriguing, since it has recently been demonstrated that mitotic recombination is elevated in Rqh1 mutants (55). This increase may reflect a direct role for Rqh1 in suppressing recombination, or it could be due to an increased level of spontaneous DNA damage in rqh1 mutants (perhaps associated with a role for Rqh1 during normal S phase) which provides appropriate substrates to initiate recombination. Further work will clearly be required to clarify this point.

Rqh1 as a model for RecQ-related helicases in eukaryotic replication and damage responses. The E. coli RecQ helicase is involved in both DNA recombination and repair (38) and has been proposed to regulate recombination (26). Human homologs of the *recQ* gene include the Bloom's syndrome gene BLM (14), the Werner's syndrome gene WRN (64), and a third potential helicase gene known as the RECQL (48) or helicase Q1 (52), which encodes the major DNA-dependent ATPase activity in human cells. Phylogenetic analysis indicates that the Rqh1 protein is most closely related to BLM and RECQL. A single S. cerevisiae member of the RecQ family of proteins, Sgs1p, has been identified as interacting with topoisomerases (20, 61) but does not apparently impart significant radiation or hydroxyurea sensitivity to cells (60). The association of a radiation-sensitive phenotype with mutation of the S. pombe recQhomolog rqh1 provides an ideal phenotypic marker with which we can investigate the mechanism of action of this class of helicases in DNA replication and repair.

Our work demonstrates, for the first time, a link between Rqh1 function, DNA damage checkpoint proteins, and recombination functions. The requirement for Rqh1 when DNA synthesis is compromised by mutations affecting chain elongation (but not its initiation) further suggests a relationship between these Rqh1-dependent responses and DNA replication. Stewart et al. demonstrated enhanced mitotic recombination associated with loss of rgh1, a phenotypic link between the rgh1 null mutant and Bloom's syndrome cells (55). Taken together, these data suggest that the RecQ-related helicase in fission yeast, and by extension the BLM function in human cells, acts to affect recombination functions during S-phase arrest. When S phase is arrested by hydroxyurea (which depletes nucleotide pools), recombination is repressed by the Rqh1 function. We suggest that in response to unrepaired DNA damage, recombination is channeled into a postreplication repair-like pathway that allows the bypass of otherwise toxic lesions during DNA synthesis. Insight into the relationship between BLM function, the checkpoint proteins, and the effects on genomic

stability and cancer susceptibility should thus be gained by studying the RecQ homolog in fission yeast.

REFERENCES

- Al-Khodairy, F., and A. M. Carr. 1992. DNA repair mutants defining G2 checkpoint pathways in *Schizosaccharomyces pombe*. EMBO J. 11:1343– 1350.
- Al-Khodairy, F., E. Fotou, K. S. Sheldrick, D. J. F. Griffiths, A. R. Lehmann, and A. M. Carr. 1994. Identification and characterisation of new elements involved in checkpoints and feedback controls in fission yeast. Mol. Biol. Cell 5:147–160.
- Aves, S. J., B. W. Durkacz, A. Carr, and P. Nurse. 1985. Cloning, sequencing and transcriptional control of the *Schizosaccharomyces pombe cdc10* "start" gene. EMBO J. 4:457–463.
- Barbet, N. C., W. J. Muriel, and A. M. Carr. 1992. Versatile shuttle vectors and genomic libraries for use with *Schizosaccharomyces pombe*. Gene 114: 59–66.
- Bentley, N. J., D. A. Holtzman, G. Flaggs, K. S. Keegan, A. DeMaggio, J. C. Ford, M. Hoekstra, and A. M. Carr. 1996. The S. pombe rad3 checkpoint gene. EMBO J. 15:6641–6651.
- Bowman, K. K., K. Sidik, C. A. Smith, J.-S. Taylor, P. W. Doetsch, and G. A. Freyer. 1994. A new ATP-independent DNA endonuclease from *Schizosaccharomyces pombe* that recognizes cyclobutane pyrimidine dimers and 6-4 photoproducts. Nucleic Acids Res. 22:3026–3032.
- Carr, A. M. 1997. Control of cell cycle arrest by the Mec1^{sc}/Rad3^{sp} DNA structure checkpoint pathway. Curr. Opin. Genet. Dev. 7:93–98.
- Carr, A. M., and M. F. Hoekstra. 1995. The cellular responses to DNA damage. Trends Cell Biol. 5:32–40.
- Carr, A. M., and J. M. Murray. 1996. DNA repair and checkpoint controls in fission yeast: a practical guide, p. 133–147. *In* K. W. Adolph (ed.), Microbial genome methods. CRC Press, Boca Raton, Fla.
- Carr, A. M., K. S. Sheldrick, J. M. Murray, R. Al-Harithy, F. Z. Watts, and A. R. Lehmann. 1993. Evolutionary conservation of excision repair in *Schizo-saccharomyces pombe*: evidence for a family of sequences related to the *Saccharomyces cerevisiae RAD2* gene. Nucleic Acids Res. 21:1345–1349.
- Chaganti, R. S. K., S. Schonberg, and J. German. 1974. A many fold increase in sister-chromatid exchanges in Bloom's syndrome lymphocytes. Proc. Natl. Acad. Sci. USA 71:4508–4512.
- Chan, J. Y. H., F. F. Becker, J. German, and J. H. Ray. 1987. Altered DNA ligase I activity in Bloom's syndrome cells. Nature 325:357–359.
- Cimprich, K. A., T. B. Shin, C. T. Keith, and S. L. Schreiber. 1996. cDNA cloning and gene mapping of a candidate human cell cycle checkpoint protein. Proc. Natl. Acad. Sci. USA 93:2850–2855.
- Ellis, N. A., J. Groden, T.-Z. Ye, J. Straughen, D. J. Lennon, S. Ciocci, M. Proytcheva, and J. German. 1995. The Bloom's syndrome gene product is homologous to RecQ helicases. Cell 83:655–666.
- Enoch, T., A. M. Carr, and P. Nurse. 1992. Fission yeast genes involved in coupling mitosis to completion of DNA replication. Genes Dev. 6:2035– 2046.
- Fernandez-Sarabia, M. J., C. McInerny, P. Harris, C. Gordon, and P. Fantes. 1993. The cell cycle genes cdc22+ and suc22+ of fission yeast Schizosaccharomyces pombe encode the large and small subunits of ribonucleotide reductase. Mol. Genet. 238:241–251.
- Forsburg, S. L., and P. Nurse. 1994. The fission yeast cdc19+ gene encodes a member of the MCM family of replication proteins. J. Cell Sci. 107:2779– 2788.
- Francesconi, S., A. M. De Recondo, and G. Baldacci. 1995. DNA polymerase delta is required for the replication feedback control of cell cycle progression in *Schizosaccharomyces pombe*. Mol. Gen. Genet. 246:561–569.
- Freyer, G. A., S. Davey, J. V. Ferrer, A. M. Martin, D. Beach, and P. W. Doetsch. 1995. An alternative eukaryotic DNA excision repair pathway. Mol. Cell. Biol. 15:4572–4577.
- Gangloff, S., J. P. McDonald, C. Bendixen, L. Arthur, and R. Rothstein. 1994. The yeast type I topoisomerase Top3 interacts with Sgs1, a DNA helicase homolog: a potential eukaryotic reverse gyrase. Mol. Cell. Biol. 14:8391–8398.
- German, J., D. Bloom, and E. Passarge. 1979. Bloom's syndrome. VII. Progress report for 1978. Clin. Genet. 15:361–367.
- German, J., L. P. Crippa, and D. Bloom. 1974. Bloom's syndrome. IV. Analysis of 101 chromatid exchange figures. Chromosoma 48:361–366.
- Gianelli, F., P. F. Benson, S. A. Pawsey, and P. E. Polani. 1977. Ultraviolet light sensitivity and delayed DNA-chain maturation in Bloom's syndrome fibroblasts. Nature 265:466–469.
- 24. Griffiths, D. J. F., N. C. Barbet, S. McCready, A. R. Lehmann, and A. M. Carr. 1995. Fission yeast *rad17*: a homologue of budding yeast *RAD24* that shares regions of sequence similarity with DNA polymerase accessory proteins. EMBO J. 14:5812–5823.
- Gutz, H., H. Heslot, U. Leupold, and N. Loprieno. 1974. Schizosaccharomyces pombe, p. 395–446. *In* R. C. King (ed.), Handbook of genetics, vol. 1. Plenum Press, New York, N.Y.
- Hanada, K., T. Ukita, Y. Kohno, K. Saito, J. I. Kato, and H. Ikeda. 1997. RecQ DNA helicase is a suppressor of illegitimate recombination in *Esch*-

erichia coli. Proc. Natl. Acad. Sci. USA 94:3860-3865.

- Hand, R., and J. German. 1975. A retarded rate of DNA chain growth in Bloom's syndrome. Proc. Natl. Acad. Sci. USA 72:758–762.
- Hoeijmakers, J. H. J., and A. R. Lehmann. 1994. Nucleotide excision-repair among species, p. 57–82. *In* R. G. Tardiff, P. H. M. Lohman, and G. N. Wogan (ed.), Methods to assess DNA damage and repair: interspecies comparisons. John Wiley & Sons Ltd., Chichester, England.
- Hoheisel, J. D., E. Maier, R. Mott, L. McCarthy, A. V. Grigoriev, L. C. Schalkwyk, D. Nizetic, F. Francis, and H. Lehrach. 1993. High resolution cosmid and P1 maps spanning the 14 Mb genome of the fission yeast S. nombe. Cell 73:109–120.
- Kelly, T. J., S. Martin, S. L. Forsburg, R. J. Stephen, A. Russo, and P. Nurse. 1993. The fission yeast *cdc18+* gene product couples S phase to start and mitosis. Cell 74:371–382.
- Langlois, R. G., W. L. Bigbee, R. H. Jensen, and J. German. 1989. Evidence for increased in vivo mutation and somatic recombination in Bloom's syndrome. Proc. Natl. Acad. Sci. USA 86:670–674.
- Lehmann, A. R., M. Walicka, D. J. F. Griffiths, J. M. Murray, F. Z. Watts, S. McCready, and A. M. Carr. 1995. The *rad18* gene of *Schizosaccharomyces pombe* defines a new subgroup of the SMC superfamily involved in DNA repair. Mol. Cell. Biol. 15:7067–7080.
- 33. Lieberman, H. B., K. M. Hopkins, M. Nass, D. Demetrick, and S. Davey. 1996. A human homologue of the *Schizosaccharomyces pombe rad9+* checkpoint control gene. Proc. Natl. Acad. Sci. USA 93:13890–13895.
- 34. Lindsay, H. D., D. J. F. Griffiths, R. Edwards, J. M. Murray, P. U. Christensen, N. Walworth, and A. M. Carr. S-phase specific activation of Cds1 kinase defines a subpathway of the checkpoint response in S. pombe. Genes Dev., in press.
- Lu, J., J. R. Mullen, S. J. Brill, S. Kleff, A. M. Romeo, and R. Sternglanz. 1996. Human homologues of yeast helicase. Nature 383:678–679.
- 36. MacNeill, S. A., S. Moreno, N. Reynolds, P. Nurse, and P. A. Fantes. 1996. The fission yeast Cdc1 protein, a homologue of the small subunit of DNA polymerase δ, binds to Pol3 and Cdc27. EMBO J. 15:4613–4628.
- Mairorano, D., G. B. Van Assendelft, and S. E. Kearsey. 1996. Fission yeast cdc21, a member of the MCM protein family, is required for onset of S phase and is located in the nucleus throughout the cell cycle. EMBO J. 15:863–872.
- Mendonca, V. M., H. D. Klepin, and S. W. Matson. 1995. DNA helicases in recombination and repair: construction of a ΔuvrD ΔhelD ΔrecQ mutant deficient in recombination and repair. J. Bacteriol. 177:1326–1335.
- Moreno, S., A. Klar, and P. Nurse. 1991. Molecular genetic analysis of fission yeast *Schizosaccharomyces pombe*. Methods Enzymol. 194:795–826.
- Morgan, B. A., F. L. Conlon, M. Manzanares, J. B. Millar, N. Kanuga, J. Sharpe, R. Krumlauf, J. C. Smith, and S. G. Sedgwick. 1996. Transposon tools for recombinant DNA manipulation: characterization of transcriptional regulators from yeast. Proc. Natl. Acad. Sci. USA 93:2801–2806.
- Muris, D. F. R., K. Vreeden, A. M. Carr, B. C. Broughton, A. R. Lehmann, P. H. M. Lohman, and A. Pastink. 1993. Cloning the *RAD51* homologue of *Schizosaccharomyces pombe*. Nucleic Acids Res. 21:4586–4591.
- 42. Muris, D. F. R., K. Vreeken, A. M. Carr, C. Smidt, P. H. M. Lohman, and A. Pastink. 1996. Isolation of the *Schizosaccharomyces pombe RAD54* homolog, *rhp54*⁺, a gene involved in the repair of radiation damage and replication fidelity. J. Cell Sci. 109:73–81.
- 43. Murray, J. M., C. Doe, P. Schenk, A. M. Carr, A. R. Lehmann, and F. Z. Watts. 1992. Cloning and characterisation of the *S. pombe rad15* gene, a homologue to the *S. cerevisiae RAD3* and human *ERCC2* genes. Nucleic Acids Res. 20:2673–2678.
- 44. Murray, J. M., M. Tavassoli, R. Al-Harithy, K. S. Sheldrick, A. R. Lehmann, A. M. Carr, and F. Z. Watts. 1994. Structural and functional conservation of the human homolog of the *Schizosaccharomyces pombe rad2* gene, which is required for chromosome segregation and recovery from DNA damage. Mol. Cell. Biol. 14:4878–4888.
- 45. Nasmyth, K. A. 1977. Temperature-sensitive lethal mutants in the structural

gene for DNA ligase in the yeast Schizosaccharomyces pombe. Cell 12:1109– 1120.

- Nasmyth, K. A., and P. Nurse. 1981. Cell division cycle mutants altered in DNA replication and mitosis in the fission yeast *Schizosaccharomyces pombe*. Mol. Gen. Genet. 182:119–124.
- Phipps, J., A. Nasim, and D. R. Miller. 1985. Recovery, repair and mutagenesis in *Schizosaccharomyces pombe*. Adv. Genet. 23:1–72.
- Puranam, K. L., and P. J. Blackshear. 1994. Cloning and characterization of RECQL, a potential human homologue of the *Escherichia coli* DNA helicase RecQ. J. Biol. Chem. 269:29838–29845.
- Rowley, R., S. Subramani, and P. G. Young. 1992. Checkpoint controls in Schizosaccharomyces pombe: rad1. EMBO J. 11:1335–1342.
- Russell, P., and P. Nurse. 1986. cdc25 + functions as an inducer in the mitotic control of fission yeast. Cell 45:145–153.
- 51. Savitsky, K., A. Bar-Shira, S. Gilad, G. Rotman, Y. Ziv, L. Vanagaite, D. A. Tagle, S. Smith, T. Uziel, S. Sfez, M. Ashkenazi, I. Pecker, M. Frydman, R. Harnik, S. R. Patanjali, A. Simmons, G. A. Clines, A. Sartiel, R. A. Gatti, L. Chessa, O. Sanal, M. F. Lavin, N. G. J. Jaspers, M. R. Taylor, C. F. Arlett, T. Miki, S. M. Weissman, M. Lovett, F. S. Collins, and Y. Shiloh. 1995. A single ataxia telangiectasia gene with a product similar to PI 3-kinase. Science 268:1749–1753.
- 52. Seki, M., H. Miyazawa, S. Tada, J. Yanagisawa, T. Yamaoka, S. Hoshino, K. Ozawa, T. Eki, M. Nogami, and K. Okumura, H. Taguchi, F. Hanaoka, and T. Enomoto. 1994. Molecular cloning of cDNA encoding human DNA helicase Q1 which has homologue to *Escherichia coli* RecQ helicase and localization of the gene at chromosome 12p12. Nucleic Acids Res. 22:4566–4573.
- Shinohara, A., H. Ogawa, Y. Matsuda, N. Ushio, K. Ikeo, and T. Ogawa. 1993. Cloning of human, mouse and fission yeast recombination genes homologous to *RAD51* and *recA*. Nat. Genet. 4:239–243.
- Sidik, K., H. B. Lieberman, and G. A. Freyer. 1992. Repair of DNA damaged by UV light and ionizing radiation by cell-free extracts prepared from *Schizo-saccharomyces pombe*. Proc. Natl. Acad. Sci. USA 89:12112–12116.
- 55. Stewart, E., C. Chapman, F. Al-Khodairy, A. M. Carr, and T. Enoch. 1997. rqh1⁺, a fission yeast gene related to the Bloom's and Werner's syndrome genes, is required for reversible S-phase arrest. EMBO J. 16:2682–2692.
- Takao, M., R. Yonemasu, K. Yamamoto, and A. Yasui. 1996. Characterization of a UV endonuclease gene from the fission yeast *Schizosaccharomyces pombe* and its bacterial homolog. Nucleic Acids Res. 24:1267–1271.
- Taylor, E. M., R. J. McFarlane, and C. Price. 1996. 5-Azacytidine treatment of the fission yeast leads to cytotoxicity and cell cycle arrest. Mol. Gen. Genet. 253:128–137.
- Walworth, N., and R. Bernards. 1996. rad-dependent responses of the chk1encoded protein kinase at the DNA damage checkpoint. Science 271:353– 356.
- Walworth, N., S. Davey, and D. Beach. 1993. Fission yeast chk1 protein kinase links the *rad* checkpoint pathway to cdc2. Nature 363:368–371.
- Watt, P. M., I. D. Hickson, R. H. Borts, and E. J. Louis. 1996. SGS1, a homologue of the Bloom's and Werner's syndrome genes, is required for maintenance of genomic stability in Saccharomyces cerevisiae. Genetics 144: 935–945.
- Watt, P. M., E. J. Louis, R. H. Borts, and I. D. Hickson. 1995. Sgs1: a eukaryotic homolog of *E. coli* RecQ that interacts with topoisomerase II *in vivo* and is required for faithful chromosome segregation. Cell 81:253–260.
- Willis, A. E., and T. Lindahl. 1987. DNA ligase I deficiency in Bloom's syndrome. Nature 325:355–357.
- Yonemasu, R., S. McCready, J. M. Murray, F. Osman, M. Takao, K. Yamamoto, A. R. Lehmann, and A. Yasui. 1997. Characterization of the alternative excision repair pathway of UV-damaged DNA in *Schizosaccharomyces pombe*. Nucleic Acids Res. 25:1553–1558.
- 64. Yu, C.-E., J. Oshima, Y.-H. Fu, E. M. Wijsman, F. Hisama, R. Alisch, S. Matthews, J. Nakura, T. Miki, S. Ouasi, G. M. Martin, J. Mulligan, and G. D. Schellenberg. 1996. Positional cloning of the Werner's syndrome gene. Science 272:258–262.