SURVEY AND SUMMARY

Replication fork reversal and the maintenance of genome stability

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ABSTRACT

The progress of replication forks is often threatened in vivo, both by DNA damage and by proteins bound to the template. Blocked forks must somehow be restarted, and the original blockage cleared, in order to complete genome duplication, implying that blocked fork processing may be critical for genome stability. One possible pathway that might allow processing and restart of blocked forks, replication fork reversal, involves the unwinding of blocked forks to form four-stranded structures resembling Holliday junctions. This concept has gained increasing popularity recently based on the ability of such processing to explain many genetic observations, the detection of unwound fork structures in vivo and the identification of enzymes that have the capacity to catalyse fork regression in vitro. Here, we discuss the contexts in which fork regression might occur, the factors that may promote such a reaction and the possible roles of replication fork unwinding in normal DNA metabolism.

INTRODUCTION

The idea that replication forks break down and act as a significant source of genome instability has gained widespread acceptance over the last decade. This popularity has not arisen because of a realization that lesions within the template DNA can block replication—such inhibition was demonstrated many decades ago (1,2). Rather, the potential links between genome instability and replication fork breakdown have gained prominence because of in vivo and in vitro studies that suggest movement of replication forks is inhibited far more frequently than previously suspected (3). Central to many models of how impeded replication forks are processed is the idea that a blocked fork can be unwound to generate a four-stranded DNA structure resembling a Holliday junction formed during recombination, first proposed in 1976 (Figure 1) (4,5). This so-called ‘fork regression’ has been proposed by many groups to aid the repair of damaged forks although the details of how such regressed forks (also referred to as ‘chicken foot’ structures (6)) might be processed differ between models. Indeed, at first glance, it is difficult to see how unwinding of a blocked replication fork could enhance the efficiency of repair of such a fork. The original block would be unaffected by this unwinding (Figure 1) and so would presumably be able to block the progression of a reassembled replisome. However, the strand switching involved in regression might facilitate excision repair of single-stranded DNA lesions by relocation of such damage into regions of duplex DNA, a process that might also facilitate lesion bypass rather than repair (see below, Figure 5). More generally, access of repair enzymes to DNA lesions could be hampered by the presence of a blocked replication fork, and fork unwinding might relieve this inhibition (see below, Figures 6–8).

This review aims to summarize the evidence that replication fork unwinding does occur, the situations in which it might occur and with what frequency. The possible consequences of fork regression and subsequent processing of these regressed forks on genome duplication and stability will also be explored. As will be seen, there are many unanswered questions within this field, in part because much of the data concerning fork regression is indirect in nature. We have aimed therefore for a balanced overview of this area, which might give the impression of equivocation. However, we believe that the many uncertainties regarding fork regression demand such an approach until more direct evidence becomes available.

WHEN MIGHT FORK REGRESSION OCCUR?

The assumption behind most models invoking replication fork reversal is that fork unwinding occurs only upon disengagement of the replication machinery from the forked DNA. Consideration of the disposition of

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components of the replication machinery at the replication fork supports this assumption. Firstly, disruption of the primer-template junction on the leading strand upon fork regression would inhibit interaction of the leading strand polymerase with DNA. Secondly, fork regression would also displace the replicative helicase away from the ends of the newly synthesized DNA strands.

If fork unwinding occurs after, or concomitant with, disengagement of the replication machinery from the fork, when might such disengagement occur? The high processivities of replication suggest that the probabilities of active replisomes disengaging from the chromosome spontaneously are very low. Disengagement of the replication machinery might therefore occur only upon blockage of replication. Many types of DNA lesion within single strands of template DNA are known to inhibit replicative DNA polymerases (7,8) whilst interstrand cross-links inhibit processes involving DNA strand separation (9). Such DNA damage can arise due to both endogenous and exogenous agents (10). Moreover, proteins bound to the template DNA, in particular transcribing RNA polymerases, present barriers to fork progression both in vivo and in vitro (11,12) as do covalent protein–DNA complexes (13). Protein–DNA barriers to replication also include pre-programmed blocks such as the Escherichia coli Tus-ter system and the eukaryotic replication fork barrier (RFB) located within rDNA repeats (14–16).

If replication fork regression occurs only upon blockage of a fork and disengagement of the replication enzymes from the fork, then the frequency of fork regression will be determined, at least in part, by the frequency of blockage and the stability of replisome association with a blocked fork. These two critical parameters will be dealt with below.

FREQUENCY OF REPLICATION BLOCKAGE

Bacteria

Analysis of replication blockage often involves the use of DNA-damaging agents such as UV light or depletion of deoxyribonucleotide pools by treatment with hydroxyurea, all of which increase the probability of replication blockage. However, estimating the frequency with which forks come to a halt in vivo in the absence of exogenous agents has proven difficult, largely because of the multiple overlapping mechanisms that exist in cells in order to repair blocked forks. For instance, in E. coli replication might be promoted past a lesion in the leading strand template by a specialized translesion DNA polymerase (17), by processing of forks by the strand exchange protein RecA (18) or by PriC-directed replication fork reloading downstream of the lesion (19). Thus, the phenotypic consequences of inactivation of a single fork repair pathway are often uninformative as regards the probability of replication blockage.

One potential method of avoiding the complications of overlapping pathways is to use mutants deficient specifically in the ability to reload the replication apparatus after blockage. A study of E. coli cells with a temperature-sensitive mutation in DnaC, thought to be required only for the loading of the replicative helicase onto the chromosome, revealed that ~18% of cells failed to complete chromosome duplication during a single round of replication under non-permissive conditions (20). These data suggest that replication fork blockage is a relatively rare event, at least in E. coli. However, whether the mutant DnaC used in this study is completely defective at the non-permissive temperature is unknown and so cells might have retained at least some ability to rebuild blocked replication forks. Indeed, other studies have suggested a higher frequency of fork blockage in E. coli. Reloading of the replication apparatus back onto the chromosome away from the normal origin of replication, oriC, requires either PriA or PriC in addition to DnaC (21). Strains with deletions in either priA or priC are viable, whereas strains deleted for both genes are inviable, implying that most cells do require the ability to reload replisomes away from oriC even in the absence of elevated levels of DNA damage (21). The lethality of combinations of mutations in recombination and repair genes [see, for example, (22,23)], also hints at a high frequency of replication fork problems.

Eukaryotes

Estimates of the frequency of replication fork blockage in eukaryotes are even less precise. Little is known concerning replication fork reloading mechanisms in eukaryotes (see below) and so specific disabling of any putative replisome reloading mechanism(s), as described above for E. coli, is currently not possible. Mutations in recombination enzymes that may promote fork restart, such as the strand exchange protein Rad51, do lead to loss of viability (24). However, recombination enzymes are involved in many aspects of DNA metabolism, not just the processing of blocked replication forks. Thus the pleiotropic effects of recombination defects make it difficult to correlate viability defects with specific defects in resumption of replication. However, as in E. coli, growth defects exhibited by combinations of mutations in genes encoding...
recombination and/or repair enzymes do indicate that fork progression is often interrupted [see, for example, (25,26)].

**BLOCKED FORK STABILITIES**

**Bacteria**

If fork unwinding is likely to occur only upon fork blockage and replisome disengagement, how long do replisomes remain associated with DNA after blockage? The functional half-life of blocked *E. coli* replisomes reconstituted in vitro is 4–6 min regardless of whether forks are halted by accumulation of positive torsional strain (27) or by collision with protein–DNA complexes (28). Whether this loss of functionality in vitro is associated with complete dissociation of the replisome remains unknown but these data do imply that at least some components of a blocked replisome have limited stability. However, blocked fork stabilities of minutes in vitro contrast with the observed association of *E. coli* replication enzymes at the site of a site-specific replication block for several hours in vivo, as measured by live-cell imaging (29,30). Such forks, even after being blocked for 2 h, retained the ability to resume replication rapidly upon removal of the block (29).

Comparison of blocked fork stabilities in vitro and in vivo might suggest the presence of stabilizing factors present within *E. coli* cells but absent in reconstituted replication systems in vitro. However, the apparent prolonged association of replication enzymes at a blockage site in vivo might also reflect continual dissociation and then reloading of the replication machinery via PriA or PriC (21). It remains possible therefore that disengagement of the replisome, and thus the potential for unwinding of the fork, occurs within a few minutes of fork blockage in *E. coli*.

**Eukaryotes**

*Saccharomyces cerevisiae* replisomes blocked by nucleo-protein complexes or by depletion of deoxynucleotid pools remain associated with blocked forks for up to 60 min in vivo, as measured by chromatin immunoprecipitation and PCR (31,32). These data support extended fork stabilities in vivo. However, whilst fork-reloading systems similar to those found in *E. coli* remain to be identified in eukaryotes, the ability to initiate replication at recombination intermediates (33) implies such systems do exist. Indeed, restart of at least partially disassembled replication forks has been documented in *Xenopus* (34). Thus monitoring of blocked fork stabilities in eukaryotic cells by chromatin immunoprecipitation might be complicated by continual replisome dissociation and reloading, as in *E. coli*. However, although there is little evidence that stabilization factors exist in bacteria, there is evidence of active stabilization of blocked forks in eukaryotes. In yeast, Mrc1 and Tof1 travel with replication forks (35,36) and may play a structural role within the replisome to ensure concerted unwinding and DNA polymerization (37,38) even in the absence of replication blocking agents (39). The replication checkpoint also promotes retention of function of eukaryotic forks blocked by hydroxyurea treatment or DNA alkylation, facilitating continued replication upon removal of the block (40,41). Although the molecular basis of this stabilization by Mec1 and Rad53 kinases in *S. cerevisiae* is unclear, there is evidence that Mec1 promotes stabilization of DNA polymerases at blocked forks whilst Rad53 promotes retention of the replicative helicase at such forks (32,42). Such fork stabilization mechanisms imply that any regression of forks might be delayed in comparison with blocked forks in bacteria.

This possible reduced opportunity for fork regression in eukaryotes might reflect the multiple origins of replication present in eukaryotic chromosomes. Thus, the problem of a blocked fork in a eukaryotic chromosome might be solved by the arrival of a converging fork from an adjacent origin (43). Rapid initiation of blocked fork processing in eukaryotes might therefore be unnecessary, and possibly even dangerous, given the potential for genome rearrangements upon engagement of recombination mechanisms (44). However, the apparently greater stability of blocked forks in eukaryocytes as opposed to bacteria does not exclude the possibility that active processing of blocked forks, including fork regression, is a requirement for the maintenance of genome stability and/or cell viability given that a single unreplicated region within a genome could prove lethal. Frequency of use might not therefore reflect importance.

**THE STRUCTURE OF BLOCKED REPLICATION FORKS**

Different blocks to DNA replication are likely to result in the formation of different blocked fork structures. The semi-discontinuous nature of DNA synthesis at replication forks suggests that single-strand-specific blocks on the lagging strand template can be bypassed by re-priming lagging strand synthesis on the 3’ side of the lesion, leaving a ssDNA gap in the lagging strand duplex to be repaired later by recombination (Figure 2A) (45–48). Single-strand-specific lesions on the leading strand template may result in a different outcome. The leading strand polymerase might be inhibited but continued movement of the replicative helicase would continue to generate ssDNA, allowing lagging strand synthesis to continue some way beyond the 3’ end of the blocked leading strand resulting in a blocked fork with a leading strand gap, possibly of many hundreds of bases (Figure 2Bii) (49–52). However, recent work using *E. coli* proteins has suggested that re-priming of leading strand synthesis can occur at a fork in vitro (Figure 2Bii) (19), providing a mechanistic explanation for the repeated detection of what appears to be discontinuous leading strand synthesis in the presence of UV light-induced pyrimidine dimers or defective DNA ligase (53–56). Leading strand lesions might therefore be bypassed by re-priming in the same manner as lagging strand lesions. Such models imply that exposure of cells to DNA-damaging agents that affect primarily single strands of DNA should not have a major impact on replication fork progression, at least in bacteria which do not possess the replication checkpoints.
found in eukaryotes. However, *E. coli* cells exposed to UV light experience substantial delays in replication, recovery depends on lesion removal and also on the ability to reload replisomes back onto the chromosome, all of which suggests that re-priming of leading strand synthesis downstream of lesions might not be a frequent event (57–60). The extent to which leading strand synthesis is re-primed downstream of a lesion is therefore unclear.

In contrast to single-strand-specific blocks, double-strand-specific blocks, such as interstrand cross-links or nucleoprotein complexes, are likely to inhibit both the polymerases and the helicase. Analysis of *S. cerevisiae* replication forks halted by the preprogrammed nucleoprotein block in the rDNA cluster revealed the presence of only three bases of ssDNA on the leading strand template (61) whilst reconstituted *E. coli* replisomes blocked by the Tus-ter nucleoprotein complex possessed a ssDNA region on the lagging strand template of 50–70 bases (62). Regardless of the precise disposition of the leading and lagging strands at such blocks, both these studies imply little if any gap in the leading strand and a gap of at most a few tens of bases on the lagging strand.

**EVIDENCE THAT FORK REGRESSION DOES OCCUR**

**Initial observations**

The idea of fork regression was first suggested in 1974 (63) and received experimental support in 1976 (4,5). The formation of heavy/heavy DNA using bromodeoxyuridine pulse-labelling of replicating human cells treated with an alkylation agent or with UV light implied that pairing of the nascent DNA strands could occur during DNA replication (4,5). Strauss and co-workers (4) also detected four-armed structures in partially replicated DNA using electron microscopy. These data could be explained by extrusion of a duplex from a replication fork, resulting in the formation of double-stranded DNA in which both strands were newly synthesized (Figure 1). Formation of duplexes, containing only nascent DNA, was detected in several subsequent studies (64–67) but some of this work suggested that branch migration of replication forks was an artefact of DNA extraction or of labelling (64,65). Regardless of whether fork regression could occur in vivo, these studies did highlight the possibility that replication forks had the potential to unwind and form four-stranded DNA structures that resembled Holliday junctions.

**Genetic evidence**

The concept of fork regression received little attention during the 1980s, possibly due to the lack of evidence that such a reaction had a physiological role. However, fork regression began to be employed to explain various genetic observations in the 1990s. Hyperrecombination was detected in the replication termination region of the *E. coli* chromosome (68), a region in which protein–DNA complexes (Tus-ter) act as pre-programmed polar blocks to replication (15). Regression of a blocked fork was proposed to initiate these recombination events, with the extruded fourth arm suggested to be a substrate for recombination (68). Other work demonstrated that defects

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**Figure 2.** Possible structures of forks halted by single-strand-specific blocks such as UV-light-induced pyrimidine dimers located on either the lagging strand template (A) or the leading strand template (B).
in the *E. coli* replicative helicase DnaB or the putative accessory replicative helicase Rep resulted in the accumulation of double-stranded DNA breaks (69), which could be explained by regression and subsequent processing of blocked replication forks (70). Fork regression has also been invoked to explain genome instability associated with fork blockage in the *S. cerevisiae* rDNA region (71,72) and processing of forks blocked by transcribing RNA polymerases (73) and UV light-induced DNA damage (18) in *E. coli*.

This rebirth of the concept of fork regression led to the application of fork regression models to explain, at least in part, many aspects of cell viability, DNA repair and genome stability in relation to the interplay between replication and recombination in *E. coli* (3,43,74). Such models were also adopted to explain observations made in eukaryotic systems (44,75).

Thus, the ability of models invoking fork regression to explain at least some mutant phenotypes supports the concept of fork regression, albeit indirectly. However, such models are often contradictory which may reflect the difficulty in extrapolating what might happen in wild-type cells from observations made using mutants. For instance, cells deficient in an enzyme that processes a specific intermediate of recombination might accumulate this intermediate. An increased concentration of this intermediate might then allow processing by a second enzyme which results in a series of reactions that normally does not occur in wild-type cells. Contradictory models might also reflect the use of different methods to block replication. The nature of the initial block will have a profound effect on the structure of the blocked fork (see above) and thus how the blocked fork might be processed, as highlighted by a recent comparison of processing of forks in *E. coli* blocked either by inactivation of the replicative helicase or by UV light-induced DNA damage (76).

These interpretation problems aside, genetic analyses are a powerful tool in studying fork blockage, regression and processing. Unlike more direct detection methods, genetic assays are exquisitely sensitive to the presence of a blocked and unrepaired replication fork. Just one persistent blocked fork can have major impacts on cell viability, providing a sensitive assay for analysis of blocked fork processing (29,77–80).

**Electrophoretic evidence**

Electrophoretic techniques provide more direct monitoring of the structures of blocked replication forks. Neutral-neutral 2-D gel electrophoresis has been particularly informative in this regard since separation of DNA is based on both size and shape (81). Thus, branched DNA structures corresponding to replication bubbles, replication forks and four-armed Holliday junction structures can all be resolved (82). This approach has been used to detect DNA structures that might have arisen by fork regression since such Holliday junction-like structures (hereafter referred to as Holliday junctions) appear to migrate as a cone/spike above the ‘Y’ arc within such gels that is sensitive to treatment with a Holliday junction-specific endonuclease (71).

Although recent work has highlighted apparent discrepancies between predicted migration patterns of regressed forks and the cone-shaped signal in such gels (83), a subsequent study suggested that exonucleolytic degradation of the regressed fourth arm might explain such discrepancies (84) (Figure 6).

These cone-shaped signals have been detected in a variety of systems including the highly repetitive rDNA array in *S. cerevisiae* which contains a pre-programmed block to replication (71), hydroxyurea-treated *S. cerevisiae* cells (41), UV-treated *E. coli* (18) and a bacteriophage T4 origin of replication (84). However, whether 2-D gels provide an accurate measure of the frequency with which fork regression occurs in vivo is unknown, given that any regressed fork signals in 2-D gels reflect both the rates of formation and dissolution of such structures. Holliday junction structures can spontaneously branch migrate (85) and so any regressed forks could migrate to reform a fork structure or to form two linear duplexes, potentially reducing the intensity of any signal from such structures. Rapid processing of regressed forks in vivo could also mask the actual frequency with which fork regression occurs. The extent to which any fork unwinds to form a fourth duplex arm will also constrain the ability of electrophoretic techniques to detect regressed forks. How many base pairs at a fork might be extruded to form a fourth duplex during regression is unknown and may also differ depending on the nature of the blockage and the mechanism driving fork regression. It is conceivable that extrusion of only a few tens of base pairs might still lead to enzymatic processing of the resultant Holliday junction, but render this regressed fork structure unresolvable from a normal fork in 2-D gels.

**Electron microscopy**

A second direct detection method is electron microscopy, the technique first used to detect regressed fork structures (4). EM does not suffer from the problem of assigning electrophoretic signals to specific DNA structures since DNA structures are visualized directly. ssDNA and dsDNA can also be distinguished by EM, whereas little work has been done using 2-D gels to analyse structures containing extensive ssDNA regions (82). EM studies have shown that in *S. cerevisiae* wild-type cells treated with hydroxyurea, blocked forks accumulated only small regions of ssDNA and regressed fork structures were rarely seen (86). In contrast, in cells deficient in the DNA replication checkpoint, extensive regions of ssDNA were seen at blocked forks and many structures resembling regressed forks were observed (86). These data suggest that fork regression might be a rare occurrence upon replication blockage in a wild-type eukaryotic cell, and that fork regression might be a pathological rather than a normal physiological event in eukaryotes (41). However, whilst EM has several advantages over 2-D gels, both techniques are limited by their ‘snapshot’ nature—levels of detectable regressed forks might be influenced significantly by rapid processing and also by limited rather than extensive extrusion of a fourth arm. Thus, direct techniques such as EM and 2-D gels are more
appropriate for comparing relative levels of fork regression rather than absolute frequencies of regression.

In summary, fork regression has been detected using direct techniques but the frequency of regression appears low as judged by these methods. This low apparent frequency might be explained by regression being a pathological event (41,86) but it might also reflect the short-lived nature of regressed forks rather than their frequency of occurrence. Indeed, genetic analyses suggest that unwinding of blocked forks, and subsequent processing of these Holliday junctions, could play important roles in the completion of genome duplication and the maintenance of genome stability. Thus, whilst the frequency of fork regression is in question, a low frequency and/or rapid processing does not necessarily preclude an important role for regression in genome duplication.

Discussion up to this point has assumed that fork regression is a favourable reaction, at least under some conditions. Studies demonstrating that DNA topology and specific enzymes can promote regression support this view, providing support for the concept of fork regression.

WHAT DRIVES FORK REGRESSION?

DNA supercoiling

Separation of the parental strands at a replication fork causes overwinding of the parental duplex resulting in an increase in positive superhelicity ahead of the fork, some of which may equilibrate across the fork to form precatenanes (87,88). This torsional strain must be relieved by topoisomerases to allow continued strand separation but DNA ahead of a moving replisome is likely to be overwound to some extent (87). However, analysis of partially replicated plasmid DNA in vitro, in which DNA-intercalating agents were used to generate overwinding of the parental DNA and so mimic the overwinding found near forks in vivo, did not generate the expected numbers of supercoils or precatenanes as judged by gel electrophoretic methods (6,89). Scanning force microscopy and EM of such DNA structures revealed that fork regression had occurred, resulting in rewinding of the parental DNA strands in front of the fork and relief of the mechanical strain caused by binding of intercalating agents (6,89). This ability of positive torsional strain within template DNA to drive fork regression was supported by the use of Holliday junction-specific endonucleases to cleave replication forks reconstituted in vitro and blocked by the accumulation of positive supercoiling (90). In contrast, fork regression was inhibited in negatively supercoiled DNA (90).

These data imply that positive supercoiling may promote fork regression in vivo. Relief of torsional strain by fork regression would come with little thermodynamic cost associated with disruption of base pairs since any base pair disruption during fork regression would be matched by the formation of base pairs between the newly-formed nascent-nascent duplex and the reformed parental duplex (6). One caveat to these studies is that the forked DNA structures were embedded within plasmids and it is unknown whether the small topological domain size of plasmids provide good models for the larger, more fluid domains found in vivo (91,92).

Furthermore, the probability of regression occurring via relief of positive torsional strain would depend on the rate of dissociation of the replisome from the blocked fork (assuming replisome dissociation must precede regression) versus the rate at which topoisomerases act in the vicinity of blocked forks to provide alternative relief. Blocked fork stabilities of a few minutes (27,28) might therefore provide sufficient time for topoisomerases to remove positive supercoils ahead of the fork, reducing the likelihood of any regression being driven by positive torsional strain. Even when removal of such excess positive windings was inhibited by use of a partial loss-of-function DNA gyrase mutation in E. coli (the main topoisomerase involved in the removal of positive superhelicity ahead of forks) genetic analysis suggested that although replication fork breakdown was frequent, regression of these forks was infrequent (93). Whether positive supercoiling ahead of blocked forks does result in fork regression in vivo is therefore uncertain.

Helicases/translocases

(i) RecG: This is a branched DNA-specific helicase found in E. coli, the absence of which confers moderate defects in DNA repair and recombination (94). RecG is also the first enzyme shown to unwind DNA forks to form Holliday junctions in vitro (73). This monomeric enzyme promotes regression of forked DNA in vitro by simultaneous translocation along the leading and lagging strand templates resulting in coupled unwinding of both daughter strands (Figure 3) (73,90,95). Moreover, RecG can catalyse regression of negatively supercoiled in vitro replication intermediates even though such supercoiling inhibits spontaneous fork regression (96,97). Analysis of small DNA substrates with heterologous arms indicated that RecG preferentially bound to and unwound forks with a lagging strand but no leading strand positioned at the branch point (90) implying that RecG might target forks halted by a leading strand template lesion (Figure 2Bi). However, using forked DNA structures with homologous daughter duplex arms, reflecting the homology found in such structures in vivo, RecG promoted efficient regression of forks with varied dispositions of leading and lagging strands at the branch point (73,98,99). RecG might therefore be able to catalyse fork regression regardless of the disposition of leading and lagging strands at the branch point.

Structural analysis of Thermatoga maritima RecG bound to a forked DNA substrate revealed that the two helicase domains interact with the parental duplex of the fork indicating that the helicase motor of RecG translocates along dsDNA (100). The third domain makes critical contacts with the branch point of the fork (Figure 3A) (100–102). Within this third domain each of the
parental strands are located within grooves separated by a so-called 'wedge' motif, with each groove able to accommodate ssDNA but not dsDNA (100). These data are consistent with translocation of the helicase domains of RecG along the parental duplex catalysing disruption of hydrogen bonding within the two daughter duplexes, resulting in re-annealing of the parental DNA strands and consequent annealing of the leading and lagging strands (100,101,103).

RecG-catalysed fork regression has been used to explain the DNA repair defects found in cells lacking RecG (73,104) whilst the co-localization of Bacillus subtilis RecG with replication forks in vivo (105) and the interaction of E. coli RecG with the C-terminus of SSB in vitro (106) support a role for RecG in processing replication forks. However, RecG can unwind a range of branched nucleic acid substrates including Holliday junctions, three-strand junctions, D-loops and R-loops (107–110) reactions which are all consistent with the known contacts between RecG and forked DNA (90,100). Whilst any helicase that catalyses fork regression might also be expected to catalyse branch migration of the resultant Holliday junction, unwinding of other branched DNA substrates makes it difficult to establish unequivocally that RecG promotes fork regression in vivo. Indeed, a broad DNA substrate specificity in vitro is a recurring theme amongst enzymes with the ability to catalyse fork regression. However, whilst it is tempting to search for a single in vivo function for any particular enzyme, multiple functions are also possible and might reflect the multiplicity of DNA structures that arise during DNA replication, repair and recombination. Ascribing in vivo function by correlating phenotypes of mutants with in vitro activities of the corresponding enzyme is perhaps the most challenging aspect of analysing helicases involved in genome stability.

(ii) RuvAB: This is a helicase complex found in E. coli which, in conjunction with the endonuclease RuvC, branch migrates and cleaves Holliday junctions during the late stages of homologous recombination (111,112). This specificity for Holliday junctions is reflected in the structure of the RuvAB complex in which a RuvA tetramer binds to one face of a Holliday junction, whilst two hexamers of RuvB each encircle duplex DNA arms emerging from opposing sides of the RuvA tetramer (Figure 4A) (112). Translocation of the RuvB hexamers along the duplex DNA results in branch migration of the DNA substrate, with the DNA 'spooling' across the face of the RuvA tetramer (113–115). Thus, although RuvAB can catalyse unwinding of a range of branched DNA substrates (113,116), the structure of the RuvA tetramer, the interaction of RuvAB with the RuvC Holliday-junction-specific endonuclease and the roles of RuvA, B and C in DNA repair and recombination indicate RuvAB(C) acts primarily on Holliday junctions (112,114).

However, RuvAB has been proposed to catalyse fork regression in vivo in strains bearing defects in replisome components or in the putative accessory replicative helicase Rep (117–119). In vitro, RuvAB can unwind model DNA forks in a direction consistent with regression (Figure 4B) but this directionality is only displayed on DNA substrates in which loading of RuvB is artificially restricted to the parental DNA duplex (116). Moreover, such restriction severely inhibits RuvAB helicase activity (116). In the absence of such a restriction, RuvAB does not appear to promote efficient fork regression in vitro (73), but instead catalyses the opposite reaction resulting in unwinding rather than re-annealing of the parental duplex (Figure 4C) (116,118). These apparent discrepancies might be reconciled if RuvAB, in certain mutant strains, is needed to promote extensive extrusion and/or stabilization of the fourth duplex arm from a fork that has already undergone limited regression, catalysed either by RecG or by positive supercoiling. The very limited processivity of RecG during regression of model fork structures in vitro (120) is compatible with RecG-catalysed initial fork regression in vivo (100) which, in conjunction with the endonuclease RuvC, branch migrates and cleaves Holliday junctions during the late stages of homologous recombination (111,112).
regression followed by RuvAB-catalysed branch migration of the resultant Holliday junction.

(iii) RecQ family helicases: These are a highly conserved family of enzymes with key roles in the maintenance of genome stability (121). Unicellular organisms tend to contain only a single RecQ family member whereas multi-cellular organisms possess multiple RecQ-type enzymes (75). The single RecQ helicase found in E. coli, the founding member of this helicase family, does not promote regression of model forks in vitro (98,99). However, RecQ-type helicases are thought to perform a variety of functions in vivo which may be reflected in the variety of different domains found within members of this family in addition to the highly conserved helicase domains (122). Humans lacking the Bloom’s syndrome helicase (BLM), a member of the RecQ family, display increased genome instability (123) and, whilst the molecular basis of these phenotypes has yet to be fully established, BLM has been shown to catalyse the regression of model fork structures in vitro (98,99). Regression of blocked forks by BLM might provide one mechanism of maintaining genome stability, possibly via template switching to allow replication past ssDNA lesions without running the risk of interchromosomal recombination (4,98,99). A second human RecQ helicase, WRN, has also been shown to promote fork regression in vitro (99,124). Werner syndrome patients lacking WRN also display elevated levels of genome instability and suffer many phenotypes associated with ageing (125). WRN also possesses a 3' to 5' exonuclease activity, not seen in other human RecQ helicases, which has been implicated in degradation of the leading strand in model fork substrates (124). Notably, forks containing a gap in the leading strand at the branch point are preferred substrates for WRN-catalysed regression, suggesting WRN exonuclease and helicase activities might co-operate during any fork regression (124).

As with RecG, BLM and WRN have the ability to unwind a range of branched DNA structures (126), thus complicating any correlation between phenotypes of cells lacking each helicase and in vitro helicase activities. Indeed, BLM can resolve a recombination intermediate containing a double Holliday junction in concert with human topoisomerase IIIα (127) and can also disrupt Rad51-ssDNA filaments (128), activities, which could also account for the maintenance of genome stability by BLM. The ability of BLM and WRN to unwind a fork in the direction required for regression can also be inhibited by RPA, a human ssDNA-binding protein, if there is a significant stretch of ssDNA present at the branch point (129). Fork regression activity of BLM and WRN might therefore be restricted to specific blocked fork structures.

A third human RecQ family helicase, RecQ5β, can also unwind model forks with a leading strand gap to promote re-annealing of the parental strands and annealing of the daughter strands (129). Human RecQ5β co-localizes with sites of DNA replication in vivo and interacts physically with PCNA whilst mice and chicken cells lacking RecQ5 have high levels of genome instability, all of which are consistent with a possible role of fork regression in processing blocked forks (129–131). However, in contrast to RecG, BLM and WRN, RecQ5β does not unwind the daughter strands within model forks in a concerted fashion (129). Thus, formation of a Holliday junction from a fork by RecQ5β would presumably occur via two independent unwinding events. Whether such a reaction would result in efficient formation of a Holliday junction from a fork remains unknown but could be tested by the use of large homologous fork structures such as those employed in studies of RecG and BLM (73,98).

(iv) UvsW: This helicase plays multiple roles in replication and repair of bacteriophage T4 DNA and can also complement many of the phenotypes of E. coli recG strains (132). The DNA substrate

Figure 4. Action of RuvAB on Holliday junctions and forks. (A) Action of RuvAB on Holliday junction structures. Translocation of the two RuvB hexamers along opposing duplex arms results in movement of duplexes as indicated by dashed arrows. Strand separation results from spooling of the DNA strands across the ‘acidic pins’ found on the surface of the RuvA tetramer (176). (B) Catalysis of fork regression by RuvAB would necessitate loading of a single RuvB hexamer onto the parental duplex. (C) Binding of two RuvB hexamers onto opposing duplex arms results in unwinding of the junction in the direction opposite to that required for regression.
specification of UvsW is also similar to RecG, with forks and Holliday junctions being preferred substrates suggesting that, like RecG, UvsW may catalyse regression of forks (133,134). Indeed, UvsW is required for accumulation of regressed forks at a T4 origin of replication in vitro, as detected by gel electrophoresis, and catalyses fork regression in vitro, providing compelling evidence that UvsW-driven fork regression is a physiologically important reaction in T4 (135). However, although UvsW might be functionally similar to RecG, it has no structural homology to RecG outside of the two helicase domains but instead resembles eukaryotic Rad54 (136). Although Rad54, an Snf2 family translocase, plays a central role in eukaryotic homologous recombination and can unwind various branched DNA substrates (137), there is currently no evidence that this enzyme is involved in fork regression.

(v) Rad5 and HARP: Saccharomyces cerevisiae Rad5 is required for the bypass of UV light-induced DNA damage via a mechanism that does not involve recombination, properties consistent with a template switching mechanism of DNA damage tolerance (138). In vitro, Rad5 can promote regression of model fork substrates and branch migration of the resultant Holliday junction, consistent with damage tolerance via template switching (139). Rad5-catalysed regression occurs via concerted unwinding of the daughter strands at the fork (139) but the enzyme has no detectable helicase activity on non-branched DNA substrates (140) in contrast to RecG, BLM and WRN. This feature of Rad5 may be related to it being a member of the Snf2 family of helicases/translocases, many of which translocate along both do not unwind DNA (141). Snf2 enzymes and RecG both define families within the Superfamily 2 of helicases/translocases (142) and so it is tempting to speculate that the helicase/translocase domains of Rad5, like RecG (100), move along the parental duplex of a fork in an ATP-dependent manner resulting in disruption of daughter duplexes and fork regression.

The recent identification of a human Snf2 family member, HARP, with the ability to re-anneal ssDNA bubbles bound by RPA via ATP-driven translocation (143) is also reminiscent of the dsDNA-specific translocation by RecG that can be coupled to fork regression (90,100). It has been argued that this HARP annealing activity is distinct from fork regression since HARP does not exhibit helicase activity on partial duplex substrates (143). However, Rad5 is also an ATP-driven translocase that does not possess the ability to unwind partial duplexes but can still regress forks in vitro (139). The possibility remains therefore that HARP might catalyse fork regression.

(vi) FANCM: This is a component of the Fanconi anaemia (FA) core complex in humans (144), disruption of which results in many chromosome instability phenotypes, both spontaneous and damage-induced (145). FANCM is a Superfamily 2 helicase that, although it cannot unwind partial duplex substrates (144), can unwind both forks and Holliday junctions and can also catalyse regression of model forks in vitro (146,147). Given that FA cells are especially sensitive to DNA cross-linking agents, FANCM-catalysed regression has been suggested to counter the movement of a replisome towards an interstrand cross-link, thus maintaining access of repair enzymes to the lesion (147). The Schizosaccharomyces pombe FANCM homologue Fml1 can also promote regression of a large model fork, and cells lacking Fml1 are also sensitive to DNA cross-linking agents, demonstrating conservation of in vitro and in vivo function of this motor (148). However, whether such a model might appertain to human FANCM is unclear.

(vii) Hjm/Hel308: This is a Superfamily 2 helicase from the archaeon Sulfolobus tokodaii that might also promote regression of forks bearing both leading and lagging strands at the branch point (149). However, this reaction appears to be complex since Hjm/Hel308A can promote annealing of complementary DNA strands, and also apparently translocate in both the 5’→3’ direction and the 5’→5’ direction along ssDNA (149). More details concerning this regression reaction are needed.

Strand exchange proteins

Strand exchange proteins play central roles in DNA repair and recombination in all organisms. The strand exchange protein in E. coli, RecA, functions by binding to ssDNA in a cooperative manner to form a dynamic nucleoprotein filament (150). This filament then interacts with dsDNA and, upon association with homologous duplex DNA, can catalyse strand exchange. Such exchange is thought to play key roles in the post-replication repair of ssDNA gaps thought to be generated by re-priming of DNA synthesis downstream of single-strand-specific DNA lesions (45,47). In addition to catalysing exchange of strands between ssDNA and homologous duplex DNA, polymerization of RecA in the 5’→3’ direction along ssDNA can also extend into adjacent duplex DNA and drive strand exchange between two homologous duplexes, so-called four strand exchange (151–153).

RecA has been implicated in catalysing fork regression in mutants bearing a temperature-sensitive allele of the replicative helicase gene at the restrictive temperature (154). In vitro labelling studies coupled with 2-D electrophoretic analyses also demonstrated that, in cells exposed to UV light, regressed forks form in a RecA-dependent manner (15,59,155). Moreover, RecA can catalyse
regression in vitro of model forks bearing a 2000-nt gap in the leading strand (120,156), reflecting the possible structure of a fork blocked by a lesion in the leading strand template (see Figure 2Bi).

This ability to catalyse regression in vitro appears to be a general property of strand exchange proteins, with T4 bacteriophage UvsX and human Rad51 both being able to promote regression of model forks with a leading strand gap (157,158). The requirement for ssDNA to allow initial binding and nucleation of strand exchange proteins (150,159) implies that blocked forks possessing limited ssDNA would not be a target for these enzymes. Such fork structure specificity is supported by genetic analyses in E. coli strains with mutations in a range of helicases, with the ability of RecA to catalyse fork regression being proposed to depend on the nature of the helicase mutation (154,160).

**POSSIBLE FUNCTIONS OF FORK REGRESSION IN NORMAL DNA METABOLISM**

Many models of replication fork repair have incorporated fork regression as a critical step on the road to resuscitation of a blocked fork. However, these models are often complex and also contradictory, presenting a confusing picture of how fork regression might aid replication. This confusion is due to: (i) the problem of assigning function to an enzyme in a wild-type organism based on the phenotype of a mutant lacking this enzyme; (ii) difficulties in assigning in vivo function to an enzyme when the enzyme can act on multiple DNA substrates in vitro; (iii) problems associated with detection of regressed forks in vivo; and (iv) the use of different types of block in different studies to analyse fork processing.

If the discrepancies between different models are laid aside, how might fork regression aid genome duplication in the face of blocks to replication? In considering this question, it is critical to consider the nature of replicative blocks. Replication past DNA lesions requires either removal or bypass of the block (161). Such removal or bypass could involve reorientation of the original DNA template (see below) whilst bypass could also be effected by recruitment of a specialized translesion polymerase (7,8). In contrast, the majority of protein–DNA complexes present lower barriers to fork movement such that reloading of the replisome might allow replication to proceed through the original protein–DNA block. Indeed, stochastic blockage by most nucleoprotein complexes is likely the key difference between such ‘accidental’ blocks and pre-programmed protein–DNA blocks such as E. coli Tus-ter that present efficient replicative blocks.

Generalized models of how fork regression might aid genome duplication will now be discussed.

**Fork regression and reversal**

The most basic model invoking regression involves unwinding of the fork to form a Holliday junction and then reversal of this unwinding process to reform the original fork (4). Why might such a reaction occur? In the case of a DNA lesion within the leading strand template, lagging strand synthesis may be able to continue some way beyond this lesion resulting in the damage being within a region of ssDNA (Figure 2Bi). Thus, no intact complementary strand would be available for excision repair. Fork reversal might promote replication past DNA lesions by provision of an undamaged complementary strand via template switching, the original proposed function of fork regression in which the blocked leading strand is extended using the lagging strand as a template (Figure 5A–D) (4). Reversal of the original regression reaction followed by reloading of the replisome onto the restored fork would result in bypass of the original lesion (Figure 5D). Repair rather than bypass of the original blocking lesion could also be promoted since regression would reposition the lesion opposite the uncorrupted lagging strand template (Figure 5B, E and F). Regression and reversal might therefore facilitate bypass and/or repair of leading strand template lesions. Evidence is accumulating that post-replicative bypass mechanisms are important in eukaryotes (162) and the involvement of Rad5 (see above) in such mechanisms provides indirect evidence for lesion bypass by regression and template switching (139).

Regression followed by reversal may also promote genome duplication in other ways. Movement of the
Given that most ‘accidental’ nucleoprotein blocks to replication are likely to have a low probability of blocking an individual replisome, reinitiation of replication upstream of the block (Figure 6C) might also facilitate movement of the replisome through such a block on the second attempt (80).

Given the multiplicity of exonucleases present in all organisms it might be expected that fork restoration would more likely occur via degradation of the extruded duplex rather than reversal of fork unwinding. *E. coli* RecBCD is a rapid, processive helicase complex with a potent nuclease activity that can initiate degradation of both strands at a blunt duplex DNA end (163) and has been implicated in degradation of such dsDNA ends at regressed forks in specific mutant backgrounds (70,164,165) (Figure 6B). Whilst no enzyme complex equivalent to RecBCD has been identified in eukaryotes, multiple helicases and nucleases have recently been implicated in processing of DNA ends in eukaryotes in a manner functionally equivalent to RecBCD (166–168). Moreover, *S. cerevisiae* Exo1 endonuclease may act at reversed forks, although whether degradation occurs via a duplex extruded by fork regression is unclear (169). A second *E. coli* exonuclease, RecJ, may also degrade strands at blocked forks. RecA-catalysed reversal of forks blocked by UV lesions (see above) might reposition the lesion back into the parental duplex, facilitating excision repair (18,76). Targeting of the nascent lagging strand by RecQ helicase and RecJ 5'-3' exonuclease might then restore a fork structure onto which the replication apparatus could be reloaded, allowing resumption of replication (76). However, this model does not invoke re-annealing of the nascent leading and lagging strands. Instead, the leading strand is retained in the RecA nucleoprotein filament whilst the lagging strand is degraded.

**Regression followed by cleavage of the Holliday junction**

All organisms appear to possess endonucleases that specifically bind to and cleave Holliday junctions symmetrically, resulting in the generation of DNA duplexes (170,171). Such cleavage reactions are critical in resolving Holliday junctions formed during recombination. In the case of a regressed fork, cleavage of the four-stranded structure would generate one intact duplex (after sealing of the nick by DNA ligase) and a second duplex with a dsDNA end (70) (Figure 7A-C). Such cleavage would therefore destroy rather than remodel the blocked fork structure, on the face of it a reckless act. However, dsDNA ends can act as substrates for recombination, with strand exchange between the dsDNA end and an intact sister duplex resulting in D-loop formation and assembly of the replication apparatus onto this D-loop (21,33). At a regressed fork cleaved by a Holliday junction resolvase the end result of such recombination would be re-creation of a replication fork upstream of the original block (Figure 7D-F), the same outcome as expected of regression followed by degradation of the extruded duplex (Figure 6C).

Such a reaction might, as described above, promote access of repair enzymes to the block (Figure 7C). However, with regressed fork cleavage followed by
recombination, there would be a risk of inaccurate recombination occurring. Indeed, in *E. coli* the Holliday junction resolvase RuvC might cleave regressed replication forks only in the absence of RecBCD (70,164,165). The implication is that degradation of the extruded duplex arm (by RecBCD) is a more efficient reaction than cleavage of regressed forks. However, work analysing the interplay between replication and transcription in *E. coli* implies that direct resetting of a blocked fork by RecBCD is unlikely to provide an efficient means of restarting replication and, by implication, that cleavage of regressed forks by RuvC does (73). These apparent contradictions might simply reflect the different mutant
strains analysed in different studies and may provide a good example of how context is critical in understanding blocked fork processing.

Regression followed by recombination of the extruded duplex

Implicit in the reversal followed by degradation model (Figure 6) is the assumption that degradation would halt once the extruded arm was completely degraded. However, this processing could also result in generation of a 3’ ssDNA tail and subsequent homologous recombination (68). For example, in *E. coli* the highly processive RecBCD helicase/exonuclease (172) might continue to unwind and degrade the nascent leading and lagging strands via the blunt duplex DNA end formed by regression, in effect continuing the regression process. In such a situation, RecBCD might continue until it encounters *χ*, an octameric DNA sequence that modulates RecBCD activity such that degradation of the strand with a 3’ end is inhibited (173). The resultant 3’ ssDNA tail would then become a target for recombination with strand exchange most likely occurring with the homologous sequence found within the reformed parental duplex. The end result would be reassembly of the replication machinery at a D-loop, generating a replication fork with two Holliday junctions upstream of the fork (70) (Figure 8). Resolution of this double Holliday junction structure would therefore recreate a replication fork.

As with regression followed by Holliday junction cleavage, recombination of the extruded duplex would be accompanied by the risk of inaccurate homologous recombination between non-identical sequences. However, this risk may be reduced by the physical association of the recombining dsDNA end with the target donor sequence (the reformed parental duplex downstream of the regressed fork). In contrast, there would be no direct linkage between donor and target after cleavage of a regressed fork (Figure 7D).

Most enzymatic steps involved in this putative processing have been characterized in detail within other contexts, usually double-strand break repair (33,150,163). Recent work has also suggested a possible alternative mechanism of dealing with a double Holliday junction structure that does not involve symmetrical cleavage of both junctions by endonucleases. The human RecQ-type helicase BLM in combination with human topoisomerase IIIα can catalyse dissolution of such a structure in vitro (127) which may avoid problems associated with directionality of cleavage of Holliday junctions.

The only direct evidence that regression followed by recombination might occur *in vivo* relates to a bacteriophage T4 origin of replication. During the early stages of infection replication occurs at *ori(34)* via the formation of an R-loop that initiates DNA synthesis in one direction (174). Initiation of DNA synthesis from the fork structure at the opposite end of the origin bubble requires regression of this fork by UvsW (135)(see above) and subsequent processing by a Holliday junction resolvase (EndoVII) and also an ATPase/exonuclease complex (gp46/47) that generates ssDNA from dsDNA ends to facilitate loading of a strand exchange protein (84). Whilst the Holliday junction resolvase could conceivably operate before the exonuclease, as shown in Figure 7, the T4 origin is not a recombination hotspot (175) which supports an intramolecular (Figure 8) rather than an intermolecular recombination (Figure 7) step (84).

**SUMMARY**

A large body of evidence is accumulating that suggests replication fork regression does occur and may help to promote genome duplication in wild-type organisms even in the absence of exogenous DNA damage. In particular, the identification of an increasing number of enzymes that can catalyse regression *in vitro* (Table 1) implies that many organisms possess the potential to unwind blocked replication forks. However, most of the evidence in support of a physiological role for fork regression is indirect. The obvious technical difficulties in direct

| Factor                  | Organism                        | Properties                                                                 |
|-------------------------|---------------------------------|---------------------------------------------------------------------------|
| DNA supercoiling        | All                             | Non-enzymatic                                                             |
| RecG                    | *E. coli*                       | Superfamily 2 helicase, highly conserved in bacteria but no obvious (nuclear-encoded) eukaryotic homologues |
| RuvAB                   | *E. coli*                       | Efficient initiation of fork regression but could promote branch migration of Holliday junctions formed by regression |
| RecQ homologues         | *H. sapiens*                    | Highly conserved group of Superfamily 2 helicases. Not all RecQ-type helicases possess regression activity |
| UvsW                    | T4 bacteriophage                | Superfamily 2 helicase, bears functional (but not structural) similarity to RecG |
| Rad5                    | *S. cerevisiae*                 | Superfamily 2 DNA translocase but no detectable helicase activity |
| FANCM/Fml1              | *H. sapiens/S. pombe*           | Superfamily 2 helicases. FANCM also contains an endonuclease domain |
| Hjm/Hel308              | *S. tokodai*                    | Superfamily 2 helicase                                                   |
| Strand exchange proteins| *E. coli*/T4 bacteriophage/*H. sapiens* | Ubiquitous. Initial binding requires ssDNA, implying regression by these enzymes may be fork structure-specific |

With the exception of RuvAB all helicases/translocases currently suspected of catalysing regression are Superfamily 2 motors. Whether this reflects specific properties of Superfamily 2 motors that are needed for efficient fork regression is unknown.
detection of fork regression in vivo might explain the indirect nature of most of this evidence but it remains possible that regression occurs only infrequently under normal physiological conditions. Ultimately, determining whether fork regression does play a role in genome duplication will require two problems to be addressed. Firstly, many models imply that fork regression can promote repair or require two problems to be addressed. Secondly, an unequivocal link between the ability of an enzyme to catalyse fork regression in vitro with its function in vivo is also required. Until both of these issues are addressed, models of blocked fork processing that invoke regression will remain speculative.

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