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RecG controls DNA amplification at double-strand breaks and arrested replication forks

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Over the years since its discovery, different hypotheses have been put forward to explain the function of RecG in bacteria. These have ranged from branch migration and resolution of Holliday junctions [1–6] via the promotion and inhibition of RecA-mediated strand exchange [7,8] to replication fork reversal [9–15]. However, evidence has recently emerged that RecG is implicated in stabilising joint molecules [16] and in controlling DNA amplification by a mechanism that involves over-replication associated with DNA double-strand break repair (DSBR) [17–24]. These observations place RecG at the interface of DNA replication and DNA repair. But what is the function of RecG? Four hypotheses have been proposed to account for the role of RecG in preventing over-replication. In two of these, RecG prevents the formation of DNA double-strand ends that are associated with the generation of new origin-independent replication forks by two different mechanisms [17,21]. In the third hypothesis, RecG catalyses the formation of double-strand ends that are associated with the elimination of new origin-independent replication forks [23]. And in the fourth hypothesis, RecG prevents a form of origin-independent DNA replication known as constitutive stable DNA replication (cSDR), which is initiated at R-loops [25].

For many years, no eukaryotic homologue or orthologue of the bacterial RecG protein had been identified. However, recently several candidates have been proposed. These include the mitochondrial helicase Irc3 of Saccharomyces cerevisiae [26], the plastid and mitochondrial helicase RECG of Physcomitrella patens [27], the mitochondrial helicase RECG1 of Arabidopsis thaliana [28] and the human nuclear helicase SMARCAL1 [29]. All of these genes are implicated in the maintenance of DNA stability and all the plastid and mitochondrial genes show partial cross-complementation with recG. Irc3 and SMARCAL1 catalyse similar reactions to purified RecG on replication fork and Holliday junction substrates in vitro. SMARCAL1 is a particularly attractive orthologue of RecG as it is a nuclear DNA damage response protein that is a.

Abbreviations
cSDR, constitutive stable DNA replication; DSBR, double-strand break repair; DSBs, DNA double-strand breaks; iSDR, inducible stable DNA replication; SIOD, Schimke immunoosseous dysplasia.
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substrate for phosphorylation by ATR [30,31] and travels with the replication fork [32]. Cells lacking SMARCAL1 are prone to accumulate DSBs [32] and patients with a biallelic deficiency in SMARCAL1 have the Schimke immunoosseous dysplasia (SIOD) disease that includes cancer predisposition [33,34]. It is interesting to note that SMARCAL1 is required to accurately and effectively replicate telomeric DNA [35–37]. This is the DNA of eukaryotic chromosomes that is predicted to be most sensitive to replication restart because a stalled replication fork at this location cannot be rescued by a convergent fork from another replication origin.

In this review, we firstly discuss the importance of DNA double-strand breaks (DSBs) in DNA amplification. We then describe the evidence that RecG and RuvABC catalyse alternative steps in DNA repair by homologous recombination. This is followed by an overview of the biochemical activity of RecG and a discussion of whether the replication fork reversal reaction, which has been well documented to be catalysed by RecG in vitro, is implicated in DNA repair in vivo. We then discuss the recent evidence that RecG and RuvABC collaborate to stabilise joint molecules. Finally, we discuss the evidence that RecG prevents DNA amplification at DSBs and arrested DNA replication forks and assess the strengths of the four models that have been proposed to account for the function of RecG. Readers are encouraged to consult two recent reviews that take different perspectives. In the first of these, Piero Bianco concentrates on the biochemical activities of the protein with a particular emphasis on recent single-molecule approaches to studying replication fork reversal catalysed by RecG [38]. In the second, Christian Rudolph and colleagues discuss chromosome replication in the absence of RecG concentrating on the hypothesis that replication fork collisions are responsible for ‘pathological’ patterns of DNA replication and on the role of replication fork traps (where the Tus protein binds ter sites) in this context [39].

In eukaryotic cells, DSBs associated with DNA replication stimulate DNA amplification highlighting the importance of understanding the sources of replication-dependent DSBs and their association with over-replication.

DNA amplification, the formation of an abnormally high copy number of one or more genomic regions, is a characteristic of cancer and of the evolution of tumours that resist treatment with anticancer drugs [40–43]. It is also a mechanism that bacteria use to evolve resistance to antibiotics [44]. There is evidence that in eukaryotes DNA amplification is stimulated by impaired S-phase checkpoint activities and by chromosomal sites and treatments that elevate the frequency of DNA double-strand ends associated with DNA replication [45–50]. These amplification events are frequently associated with altered deoxyribonucleoside triphosphate pools and DNA replication stress leading to the early stages of cancer development [50–54]. For these reasons, it is critical to understand the pathways by which DNA double-strand ends are formed as a consequence of DNA replication and how these events may be associated with DNA amplification. Many of these pathways of DNA double-strand end formation have been initially investigated in prokaryotic systems but are not exclusive to prokaryotes. As depicted in Fig. 1, the pathways of replication-dependent DSB formation include: (A) replication fork reversal [55–57], (B) replication fork collapse [58], (C) replication fork rear-ending [59], (D) secondary structure cleavage [60,61], (E) replication fork restart at a 3’ flap [21], (F) template-switching with replication fork reversal [23], and (G) reverse-restart of an arrested replication fork [17]. Depending of the pathway, RecG has been proposed to promote the formation of double-strand ends (in pathways A and F) or to prevent the formation of double-strand ends (in pathways E and G). Pathways E and G postulate over-replication associated with the formation of DSBs and DNA amplification in Escherichia coli. We will evaluate below the arguments for and against the proposed in vivo roles of RecG.

RecG and RuvABC catalyse alternative steps in DNA repair and recombination

The recG gene was first identified by Storm and collaborators as a recombination-deficient mutant of E. coli K12 [62]. Cells with the recG162 or recG258 mutation were more sensitive to UV, ionising radiation and mitomycin C, and displayed reduced conjugal and P1 transductional efficiency [1,62,63]. More recent in vivo studies have confirmed the involvement of RecG in DSBR. Cells lacking RecG are sensitive to breaks induced by the I-SceI homing endonuclease [5], the EcoKI endonuclease [6] and cleavage of a 246 bp palindrome by the SbcCD DNA hairpin endonuclease [60]. The observation that (like RecA) RecG plays a role in several different homologous recombination pathways in E. coli suggests that it plays a fundamental role in DNA repair [63]. But, what does RecG do? Further understanding of the role of RecG came from genetic studies combining the recG mutation with other mutations in genes encoding proteins involved in
DNA repair and recombination [1,63]. recG mutants showed a modest additional sensitivity to UV when combined with either the recB (RecB subunit of the RecBCD enzyme, exonuclease IV, implicated in DNA double-strand end unwinding, resection and RecA loading during DSBR) or recJ (RecJ 5'-3' exonuclease, implicated in gap extension during single-strand gap repair) but not the recF mutation (RecF component of RecFOR, implicated in RecA loading during single-strand gap repair). However, more striking observations were obtained when recG was combined with ruv mutations (RuvABC implicated in the branch migration and cleavage of Holliday junctions). Double ruvA recG, ruvB recG and ruvC recG mutants exhibited a more dramatic increase in sensitivity to UV and ionising radiation, and a greater defect in recombination after conjugation or transduction when compared to either of the single mutants. These results suggest that RecG and RuvABC catalyse two alternative steps in the repair of DSBs by homologous recombination, potentially during the resolution of Holliday junctions [1]. This idea was supported by the study of rusA mutants that suppress the recombination deficiency phenotype of ruvA mutants. These suppressor strains have activated the expression of a Holliday junction resolvase gene encoded within a cryptic prophage [4]. The suppression observed in these rusA ruvA double mutants requires the presence of RecG, further suggesting that the alternative pathways catalysed by RuvABC or RecG might be for the resolution of Holliday junctions [4]. However, we describe below an alternative hypothesis to explain the redundancy of RecG and RuvABC.

**RecG protein unwinds and remodells branched DNA molecules in vitro**

Purified RecG protein has 3′–5′ helicase and nucleic acid translocase activities. In vitro, it can bind and unwind synthetic model Holliday junctions and various other types of branched DNA substrates including replication forks, D-loops and R-loops [2,3,8–10,64–68].
Unlike most other helicases, this enzyme unwinds DNA by translocating on dsDNA rather than on ssDNA. *In vitro*, RecG works as a monomer [69,70] and efficiently catalyses the re-pairing of template strands in substrates mimicking replication forks. Interestingly, RecG promoted unwinding reactions occur preferentially on substrates mimicking replication forks with a nascent strand annealed to the lagging-strand template [9,15].

RecG catalyses replication fork reversal (also known as replication fork regression) *in vitro* on a substrate containing both nascent strands (Fig. 1A) [9–14]. This RecG-catalysed replication fork reversal reaction has been observed using an oligonucleotide substrate with nascent strands annealed to both the leading- and lagging-strand templates [14], a replication fork in supercoiled plasmid DNA [71] and a replication fork blocked at a DNA lesion in an *in vitro* replication system where the DNA polymerase and the replicative helicase remain associated with the DNA [11]. These studies have led to the opinion that replication fork reversal is an important biochemical activity of RecG [9,11,12,14,15,38,66,67,70–77]. RecG can catalyse this reaction thanks to its unusual structure [70]. This 76-kDa enzyme possesses a unique translocation by RecG motif, which is located between the wedge and the helicase domains of the protein and contributes to the unwinding of branched molecules by forming a helical hairpin motif [78]. For a more detailed discussion of the structure of the RecG protein, readers are referred to the recent review [38].

**RecG does not catalyse replication fork reversal *in vivo***

In 1976, two papers proposed a mechanism for non-mutagenic replication bypass of a DNA lesion that involved reannealing of replicated template DNA strands and extrusion and pairing of newly synthesised DNA strands [55,57]. Over two decades later, a study of *E. coli rep* mutants provided evidence for the occurrence of this replication fork reversal reaction in cells with undamaged DNA but with compromised DNA replication [56]. It was proposed that the RecG-catalysed replication fork reversal reaction observed *in vitro* might also happen following UV irradiation *in vivo* [14]. The absence of this pathway in *recG* mutants would permit re-pairing of template strands to help repair DNA lesions [14]. However, none of the studies of replication fork reversal to date, using different ways of compromising DNA replication, has revealed any situation where RecG is required for the reaction *in vivo* [56,79–84]. Furthermore, a subsequent investigation showed little evidence that RecG promotes replication fork reversal following UV irradiation [85]. This generated a conundrum. Why would RecG be so good at catalysing replication fork reversal *in vitro* but unable to catalyse the reaction *in vivo*? A clue to this might be the observation that when PriA is present, RecG initiates the re-pairing of parental strands but only proceeds as far as bringing the 3′ end of the nascent leading-strand to the fork junction point, whereupon the DNA is bound by PriA in a fork-stabilising configuration (Fig. 2) [86]. We shall return to this observation later.

![Fig. 2.](image-url)
RecG and RuvABC collaborate to stabilise joint molecules during DSBR

As described above, there is good evidence that RecG and RuvABC catalyse alternative steps in the pathway of recombination, which would explain the high DNA damage sensitivity and recombination deficiency of a ruv recG double mutant. Since RuvABC is known to act as a branch migration and Holliday junction resolution complex [87], it was attractive to hypothesise that this redundancy arose from two alternative pathways of resolution of Holliday junctions. One possibility was that RecG with the help of a topoisomerase might catalyse the dissolution of structures containing two Holliday junctions as had originally been proposed for bacteriophage lambda recombination [88] and has been shown in eukaryotic chromosomes by a combination of BLM, TopoIIIa and Rmi1 (see [89]). However, a substantial proportion of chromosome dimers is generated among recombinants formed in the absence of RuvABC, indicating that crossing over has taken place in conditions where the hypothetical RecG-mediated resolution pathway would be operating [5,6]. This observation is not compatible with a dissolution pathway catalysed by RecG as topoisomerases do not catalyse crossing over and has prompted two alternative hypotheses. First, an unknown nuclease could participate in the RecG pathway of resolution [5] and second, resolution could be mediated by the next round of chromosomal DNA replication passing through the Holliday junction [6].

On the assumption that RuvABC and RecG catalyse alternative pathways of Holliday junction resolution, it was logical to look for evidence of accumulation of Holliday junction intermediates in a ruvAB recG double mutant. However, very surprisingly this double mutant failed to accumulate Holliday junction intermediates while a ruvAB mutant readily did (Fig. 3A) [16]. This result clearly showed that RuvABC is responsible for the resolution of Holliday junctions in cells containing RecG. However, few joint molecules of any kind were detected in a strain lacking both RuvAB and RecG. Clearly, the presence of either RuvAB or RecG is required to generate stable joint molecules (including molecules with Holliday junctions) in the first place [16]. This led Mawer and Leach to suggest that the branch migration activities of RuvAB and/or RecG might provide alternative ways of stabilising an initially formed and otherwise unstable form of joint molecule, thus explaining the genetic redundancy observed previously. Since joint molecules could not be stably recovered in the absence of RuvAB and RecG, it was hypothesised that initially formed intermediates generated in the absence of these proteins might consist of D-loops that could be destabilised by a helicase. Further work revealed that this helicase is PriA [17].

The stabilisation of initially formed joint molecules, consisting of D-loops generated by the RecA protein, through the branch migration activities of RuvAB and RecG is readily understandable. Given that RuvAB branch migrates Holliday junctions prior to their resolution by RuvC, it is highly probable that the stabilising activity of RuvABC operates at the Holliday junction end of a D-loop by extending the region of base pairing between the recombining duplexes, leading to their covalent exchange following cleavage and ligation (Fig. 3Bi). However, the site of action of RecG is less clearly defined by the biochemistry of the enzyme, since this protein can catalyse both the migration of Holliday junctions and the remodelling of replication forks. During DSBR both of these structures are present, one at each end of a D-loop. A clue as to the nature of the RecG substrate in vivo comes from the observation that a class of suppressors of the recG recombination-deficient phenotype carries mutations in PriA, either reducing or eliminating the helicase activity of the protein [90]. PriA plays a critical role in the reloading of DnaB, the replicative helicase, onto various DNA structures [91–94]. It does so by binding to a replication fork substrate with a 3’ end at the fork junction in a configuration whereupon the fork is stabilised and the helicase activity of PriA is switched off [95]. The helicase-defective mutants of priA that suppress the recombination-deficient phenotype of recG mutants are indeed competent for catalysing replication restart [96]. This suppression, coupled with the observation that RecG delivers PriA to a replication fork substrate in its 3’ end-binding mode [86], argue strongly for a joint molecule stabilising role of RecG associated with the replication fork end of a D-loop (Fig. 3Bii). Accordingly, we propose that D-loops are stabilised in the presence of RuvABC and RecG by activities at both DNA junctions (Fig. 3Biii). Furthermore, we conclude that this overlap in function could be responsible for the genetic redundancy of recG and ruvABC mutants.

RecG controls DNA amplification during DSBR and at arrested replication forks

It has long been known that there is a link between RecG and DNA replication. cSDR is induced in the absence of RecG [97]. cSDR is a form of DNA synthesis [98,99] that requires RecA [100,101], transcription
and is stimulated in rnhA mutants [102]. It is therefore proposed to originate from persistent R-loops that may be generated through the action of RecA. recG rnhA double mutants are not viable and it has been proposed that RecG either unwinds persistent R-loops or prevents their formation through opposing the action of RecA [97]. Inducible stable DNA replication (iSDR) is also elevated in the absence of RecG [104,105]. iSDR requires the induction of the SOS response [106], the action of RecBCD [105,107] and is insensitive to inhibition of transcription [108], consistent with resulting from DSBs. The reader is directed

Fig. 3. Stabilisation of joint molecules by RuvABC and RecG. (A) DSBR intermediates visualised by 2D gel electrophoresis. RuvAB and RecG do not simply provide alternative pathways for the resolution of Holliday junctions, as previously suggested. Four-way Holliday junction intermediates accumulate in the absence of RuvAB but not in the absence of RecG. The accumulation of Holliday junctions in the absence of RuvAB requires the presence of RecG [16]. Data reproduced with permission from PLoS Genetics. (B) Role of RuvABC and RecG in the stability of joint molecules (i) Joint molecule stabilisation by RuvABC. In the absence of RecG, RuvAB migrates the Holliday junction away from the site of initiation of DSBR and leads to its cleavage by RuvC. Both branch migration and cleavage stabilise the joint molecule. (ii) Joint molecule stabilisation by RecG. In the absence of RuvABC, RecG manipulates the replication fork end of the D-loop to allow PriA to bind in its 3’ end-binding fork-stabilising mode. This allows the initiation of DNA replication that stabilises the joint molecule. (iii) Joint molecule stabilisation by RuvABC and RecG. In the presence of both RuvABC and RecG, both the Holliday junction and replication fork ends of the D-loop are stabilised.
to the review [108] for a more detailed description of cSDR and iSDR.

During DSBR in *E. coli*, the RecBCD enzyme resects broken ends for distances of up to several kilobases [109]. It is therefore essential that the degraded DNA is restored. This is normally carried out by establishing DNA replication initiated through the action of PriA [60], arguing for the loading of the replicative helicase DnaB and the replicative DNA polymerase PolIII. However, in the absence of RecG, DNA over-replication is observed following DNA damage [19–21]. At a site-specific DNA break, this over-replication flanks the site of DSBR [17]. Furthermore, even in the absence of DNA damage, recG mutants over-replicate the terminus region of their chromosome between termination sites *terA* and *terB* [17,18,20,22,23]. This over-replication is mediated by PriA and PriB and is suppressed by combining the recG mutation with PriA-helicase mutations [22]. These results suggest that the replicative helicase DnaB loads onto DNA substrates generated in this region.

Four alternative hypotheses have been proposed, none of which is free from limitations, to explain the observation that DNA amplification is prevented by RecG.

First (Fig. 4A), DNA amplification is a consequence of DNA flaps that are hypothesised to arise when replication forks collide [18,19,21,22]. It is hypothesised that replication fork collisions frequently give rise to 3′ flaps that can be converted into 5′ flaps by RecG, and then these 5′ flaps are degraded by 5′–3′ exonucleases [18,21]. In the absence of RecG, the 3′ flaps persist and are converted into new replication forks through the action of PriA [18,21]. The existence of 3′ flaps is supported by the preference of RecG for processing 3′ flaps over 5′ flaps [15,86] and the observation of DNA over-replication in the terminus region of the chromosome of a triple 3′–5′ exonuclease mutant, *xseA xonA sbcDC* [22]. The products of the *xseA*, *xonA* and *sbcDC* genes (exonuclease VII, exonuclease I and SbcCD exo/endonuclease respectively) are the major 3′–5′ exonucleases in *E. coli*. They participate in several DNA repair and genome stability pathways and the reader is directed to review [110] for a more detailed discussion of their functions. The over-replication in the *xseA xonA sbcDC* mutant is very interesting and does indeed suggest the existence of a pathway of DNA amplification involving 3′ overhangs. However, contrary to the prediction of the model that RecG can remove 3′ flaps by converting them to 5′

![Fig. 4. Four different models proposed to explain how RecG controls DNA amplification. (A) Fork collision and restart at a 3′ flap. When two replication forks (moving in the directions of the green arrows) collide, it is hypothesised that in the absence of RecG a 3′ flap is generated that leads to the assembly of a replication fork. In the presence of RecG, the 3′ flap is converted into a 5′ flap that can be degraded by 5′–3′ exonucleases [18,19,21,22]. (B) Fork collision and template-switching followed by replication fork reversal. When two replication forks (moving in the directions of the green arrows) collide, it is hypothesised that template switching occurs leading to over-replication. This is corrected by RecG-dependent replication fork reversal and DNA degradation at one (or both) of the replication forks [23]. (C) cSDR and termination at Tus/ter blocks. It is proposed that, in the absence of RecG, cSDR initiates at sites of transcription around the genome leading to replication forks that are blocked by Tus/ter. This results principally in over-replication of the region between termination sites (at the positions of blocked red arrows) as cSDR forks are removed by colliding with origin-initiated replication forks [25]. (D) Reverse-restart of an arrested replication fork. At an arrested replication fork (at the position of the blocked red arrow) RecG prevents the assembly of the replicative helicase on the newly synthesised lagging-strand. In the absence of RecG, this loading is permitted and backwards-directed DNA replication occurs [17].
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flaps, RecG is unable to prevent this pathway as the amplification observed in the xseA xonA sbcDC mutant occurs in the presence of RecG. Conversely in the absence of RecG, the three 3′–5′ exonucleases are not able to prevent over-replication. Therefore, either a single 3′ flap processing pathway is delicately balanced between the activities of RecG on one hand and the three 3′–5′ exonucleases on the other, or there are two separate pathways operating on different substrates. The synthetic lethality of a recG xseA xonA sbcDC quadruple mutant provides some indirect evidence for the existence of a single substrate but it is not conclusive since the phenotype of DNA over-replication in the terminus region is not lethal and the cause of lethality of the quadruple mutant is unknown [18]. Furthermore, although a priA300 helicase defective mutation suppresses the DNA damage sensitivity of a recG mutant, it does not suppress the DNA damage sensitivity of an xseA xonA sbcDC mutant [18], presenting a counter-argument in favour of the existence of two distinct substrates. In this first model, PriA is hypothesised to recruit DnaB without acting in PriA’s 3′ end-binding and fork-stabilising mode, which does not fit easily with the biochemical observation that RecG remodels a replication fork substrate to favour PriA binding in its 3′ end-binding mode [86]. The DNA ends generated during this process should be at multiple positions where collisions happen between replication forks and should be pointing in both directions but in fact they are primarily generated at ter sites where they are unidirectional [17]. Finally, a complete inversion of chromosome replication is observed in a dnaA recG tus rpo* mutant [22] where replication forks cannot form at the origin of DNA replication so there is no prediction of fork collisions in the chromosome terminus region, from where replication is nevertheless observed to originate.

Second (Fig. 4B), DNA amplification is caused by replication forks sliding past each other in the terminus region of the chromosome [23]. This reaction is corrected by RecG that catalyses replication fork reversal on one (or both) of the replication forks, generating one or more DNA double-strand ends that can be degraded by RecBCD. This hypothesis differs from the first hypothesis in two principal respects. First, RecG is predicted to generate DNA double-strand ends rather than to remove a precursor of DNA double-strand ends and second the sliding of replication forks past each other requires a rather complex double DNA template switch. We now know that there is an increase in the frequency of DNA double-strand ends that bind RecA protein in the terminus region of the chromosome of a recG mutant [17], which is not predicted by this model. As with the first hypothesis, this model does not explain the inversion of chromosome replication observed in a dnaA recG tus rpo* mutant, since this model also predicts that over-replication of the terminus region requires the meeting of replication forks coming from the origin, which are absent in this mutant [22].

Third (Fig. 4C), DNA amplification in the terminus region is simply a consequence of cSDR that is allowed to occur in a recG mutant and proceeds through the terminus region until it reaches a Tus/ter block [25]. cSDR may indeed contribute in some ways to the pattern of DNA replication observed in a recG mutant. However, this hypothesis does not explain the origin of the DNA double-strand ends that bind RecA at ter sites in a recG mutant [17]. Furthermore, the unusual replication observed in a recG mutant is different from that observed in an rnhA mutant as only the former can be suppressed by a priA300 helicase-defective mutant [111]. These observations argue against the involvement of cSDR in the terminus over-replication formed in the absence of recG. In contrast, the stimulation of iSDR in a recG mutant could be related to the over-replication observed in the absence of RecG as proposed by the first and fourth hypotheses. iSDR occurs as a consequence of DSBR by homologous recombination and the recombination deficiency of recG mutants is known to be suppressed by priA300 [112].

Fourth (Fig. 4D), DNA amplification is caused by the incorrect loading of PriA at a site of replication fork arrest or at a newly formed replication fork [17], leading to the formation of a backwards-directed replication fork. This reverse-restart hypothesis is based on two observations. (a) RecG loads PriA onto a model replication fork in the 3′ end-binding and fork-stabilising mode [86], predicted to facilitate the loading of DnaB to restart the fork correctly. (b) DNA double-strand ends bound to RecA protein are detected at the sites of initiation of DNA amplification at an induced DSB and in the terminus region of the chromosome between terA and terB [17]. As attractive as this model is, it does not explain all the previous observations either. For example, it does not explain the observation of DNA amplification in the terminus region of a RecG+ cell in the absence of the 3′–5′ exonucleases. It also does not directly explain the inversion of chromosome replication observed in a dnaA recG tus rpo* mutant [22]. However, DSBs have been observed surrounding the dif site [17,113]. These breaks could provide the DNA replication initiation sites that would allow this inversion of chromosome replication to occur according to this model.
Only the first and fourth hypotheses propose that over-replication occurs as a consequence of DNA double-strand ends that are generated in the absence of RecG. The detection of RecA bound to DNA double-strand ends in the terminus region, which is specifically enhanced in a recG mutant, provides support for these two models. This stimulation of DSBR is also consistent with iSDR being induced in a recG mutant.

Conclusions and perspectives

It is clear that RecG prevents DNA amplification at a site of induced DSBR in the lacZ gene [17]. This is also the case in the terminus region of the E. coli chromosome where DNA amplification in the absence of RecG is similarly associated with DSBR [17]. These observations are only in accordance with hypotheses one and four (Fig. 4A,D). We favour the simple explanation, prevention of reverse-restart, that is described in Fig. 4D. RecG directs the correct loading of PriA, at replication forks that have lost (or not yet acquired) the DNA replication machinery. Appropriate binding of PriA allows DNA replication to proceed correctly via loading of the replicative helicase DnaB. In the presence of RecG, the formation of normal replication forks is predicted to occur at sites of DSBR where they are required to replace the DNA lost during resection. In the absence of RecG, PriA and DnaB can be loaded incorrectly to replication forks that have been created by DSBR or replication forks that have arrested and lost their replisomes. Incorrect loading of DnaB leads to DNA amplification (Fig. 4D) [17].

However, if this explanation is not correct and DNA double-strand ends arise as a consequence of replication fork collisions in the absence of RecG (Fig. 4A), then these collisions must occur primarily at terA and terB sites in a recG mutant as this is where RecA binding to DNA double-strand ends is detected by ChiP [17]. The ChiP data reveal that RecA binding is at one-ended DNA breaks all pointing in one of the two possible directions at each of the ter sites [17]. This implies that any fork collision occurring at a ter site would have to lead to a specific orientation of break. This may be possible if the direction of replication fork movement upon collision with a ter site can determine the strand on which the hypothetical 3’ single-strand is generated.

Why a xseA xonA sbcDC triple 3’–5’ exonuclease mutant stimulates DNA amplification in the terminus region of the chromosome remains to be determined. Does this amplification arise from the same pathway as the over-replication in a recG mutant, or is it mediated by a separate pathway controlled by 3’ overhangs? How DNA replication is initiated in the terminus region of a dnaA recG tus rpoB mutant also remains to be determined. Is this replication initiated by the DSBs detected on the two sides of the dif site [17,113]? Further investigations are required to answer these questions.

DNA replication restart is stringently restricted in eukaryotic cells. However, one might predict that such a pathway could exist to ensure completion of replication between the most telomere proximal origin of replication and the end of the chromosome. One might also predict that, even in the absence of a pathway for restart, incorrect loading of a replicative helicase at the site of a stalled replication fork, to allow reverse-restart, should be prevented to avoid DNA amplification. Perhaps this is where SMARCAL1 plays a role in maintaining genome stability.

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Author contributions

Both authors have contributed to writing the manuscript.

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