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A protective role for BRCA2 at stalled replication forks

Gurushankar Chandramouly, Nicholas A Willis and Ralph Scully*

Abstract

The hereditary breast and ovarian cancer predisposition genes BRCA1 and BRCA2 account for the lion's share of heritable breast cancer risk in the human population. Loss of function of either gene results in defective homologous recombination (HR) and triggers genomic instability, accelerating breast tumorigenesis. A long-standing hypothesis proposes that BRCA1 and BRCA2 mediate HR following attempted replication across damaged DNA, ensuring error-free processing of the stalled replication fork. A recent paper describes a new replication fork protective function of BRCA2, which appears to collaborate with its HR function to suppress genomic instability.

The hereditary breast and ovarian cancer predisposition genes BRCA1 and BRCA2 have crucial roles in the control of double strand break (DSB) repair by homologous recombination (HR). BRCA1 functions early in HR, interacting with nucleases, including CtIP and the Mre11/Rad50/Nbs1 (MRN) complex to coordinate DNA end resection to form single-stranded DNA [1]. BRCA1 also recruits BRCA2 to the break site. BRCA2 is a key repair component that loads the Rad51 recombinase onto single-stranded DNA of the processed DSB [2-4]. DSBs in cycling somatic cells are thought to arise predominantly during the DNA synthesis phase of the cell cycle, when a replicative DNA polymerase stalls on abnormal DNA structure. Some models invoke the use of HR to repair the broken fork using the undamaged neighboring sister chromatid as a template [5-7]. However, not all HR requires a DSB intermediate; a fork-stalling DNA adduct may be bypassed without complete breakage (‘collapse’) of the fork. Because the stalled fork is structurally different from an isolated DSB, the mechanisms governing HR in these two contexts may differ. Circumstantial evidence suggests that critical events underlying breast cancer risk, as well as the therapeutic action of poly(ADP) ribose polymerase inhibitors on BRCA-linked cancers, are played out at sites of DNA polymerase stalling [8-10]. Paradoxically, however, the experimental tools available for studying HR at sites of mammalian DNA polymerase stalling are quite limited.

In a paper published recently in Cell, Schlacher and colleagues [11] have taken an interesting new approach to studying how BRCA2 affects stalled fork metabolism. The authors used single-molecule DNA fiber analysis to determine the fate of newly synthesized (‘nascent’) DNA strands just proximal to replication forks in cells treated with hydroxyurea (HU), an agent that depletes the nucleotide pool, causing genome-wide DNA polymerase stalling. In wild-type cells, the nascent strands at HU-arrested forks were protected from degradation. In contrast, cells lacking wild-type BRCA2 revealed progressive erosion of nascent DNA strands. The authors identified the DNA end resection complex MRN as a key mediator of this process. Precisely what DNA structures MRN acts on to erode the nascent strands at HU-arrested forks in BRCA2 mutants is not yet clear, although it likely involves MRN-mediated resection of a DNA end. In wild-type cells, the short period of exposure to HU described by Schlacher and colleagues should produce few if any DSBs as direct products of fork collapse [12]. In this regard, one previous report suggested that BRCA2 mutant cells are particularly susceptible to replication fork collapse following prolonged HU treatment [13]. This raises the possibility that MRN acts on DNA ends produced by fork collapse in BRCA2 mutants. If this were the case, however, only a fraction of all nascent strands should show the observed resection. The fact that Schlacher and colleagues observed coordinated resection of all nascent strands, a process that had evidently begun within minutes of HU exposure, suggests that a mechanism other than fork collapse generated the DNA ends that MRN attacks. Experiments in yeast and bacteria have revealed that stalled forks can undergo ‘fork reversal’ - a process whereby the fork backtracks on its path, generating a cruciform DNA structure called a
‘chicken foot’, with a single exposed DNA end composed of paired nascent strands [14,15]. Schlacher and colleagues speculate that an equivalent structure can form in mammalian cells and that BRCA2 either prevents its formation or helps to protect the free DNA end of this cruciform structure from attack by MRN.

BRCA2 protection of the Hu-stalled fork from MRN-mediated resection could reflect a non-canonical role for BRCA2/Rad51 at stalled forks. Indeed, Schlacher and colleagues found that a BRCA2 mutant lacking the carboxy-terminal Rad51 binding domain is defective for protection against Hu-triggered nascent strand resection, but retains intact HR repair of a chromosomal DSB. These observations, together with other experiments reported in the paper, suggest that BRCA2 has a ‘stalled fork-protective’ function mediated by its stabilizing effect on the Rad51 nucleoprotein filament but distinct from its role in traditional DSB repair by HR [11]. A caveat of this interpretation is the likelihood that HR at stalled forks involves mechanisms additional to those required for HR at an isolated DSB. Perhaps the ‘fork-protective’ function of BRCA2 is indeed required for HR at stalled forks, but not for HR at isolated DSBs. Data obtained with use of Hu also pose some problems of interpretation, since Hu produces genome-wide arrest of otherwise structurally normal replication forks. It is not clear to what extent this static form of ‘replication stress’ reflects the dynamic interactions that occur between the replication fork and a DNA polymerase-stalling carcinogenic DNA adduct.

Our current view of BRCA1 and BRCA2 suggests that regulation of HR at stalled replication forks influences the probability of a woman developing breast cancer as well as her response to therapy. The work by Schlacher and colleagues reveals new mechanisms underlying this relationship. The rewards of continuing to dig deeper into basic mechanisms of action of the BRCA genes will be a better understanding of breast cancer risk and breast cancer therapy. Perhaps this will include the discovery of new therapies that take advantage of the replication-recombination ‘Achilles heel’ of BRCA mutant breast cancers and of other cancers in which HR is defective.

Competing interests
The authors declare that they have no competing interests.

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Abbreviations
DSB, double strand break; HR, homologous recombination; HU, hydroxyurea; MRN, Mre11/Rad50/Nbs1.