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Sources, resolution and physiological relevance of R-loops and RNA–DNA hybrids

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Abstract

RNA–DNA hybrids are generated during transcription, DNA replication and DNA repair, and are crucial intermediates in these processes. When RNA–DNA hybrids are stably formed in double-stranded DNA, they displace one of the DNA strands and give rise to a three-stranded structure called an R-loop. R-loops are widespread in the genome and enriched at active genes. R-loops have important roles in regulating gene expression and chromatin structure, but they also pose a threat to genomic stability, especially during DNA replication. To keep the genome stable, cells have evolved a slew of mechanisms to prevent aberrant R-loop accumulation. Although R-loops can cause DNA damage, they are also induced by DNA damage and act as key intermediates in DNA repair, such as transcription-coupled and RNA-templated DNA break repair. When the regulation of R-loops goes awry, pathological R-loops accumulate, which contribute to diseases such as neurodegeneration and cancer. In this Review, we discuss the current understanding of the sources of R-loops and other RNA–DNA hybrids, mechanisms that suppress and resolve these structures, the impact of these structures on DNA repair and genomic stability, and opportunities to therapeutically target pathological R-loops.

Introduction

R-loops are a three-stranded nucleic acid structure consisting of RNA–DNA hybrid and a displaced strand of DNA. R-loops and other RNA–DNA hybrids, such as those formed at DNA replication forks and at processed double-stranded DNA (dsDNA) ends, are important intermediates in transcription, DNA replication and DNA repair. Studies in multiple organisms have revealed that R-loops have crucial roles in cellular processes such as chromatin organization and chromosome segregation¹⁻⁴. The RNA–DNA hybrids formed at DNA replication forks and processed DNA ends do not displace single-stranded DNA (ssDNA) and thus do not constitute R-loops, but are also important for DNA replication and repair^{4,5}. Paradoxically, emerging evidence suggests that R-loops are a major source of genomic instability, highlighting the importance of proper regulation of R-loops.

Recent genetic and biochemical studies have uncovered numerous regulators of R-loops, providing insights into how R-loops are generated and resolved in various cellular and chromosomal

contexts. The functions of R-loops in different cellular processes, especially in DNA repair are also increasingly appreciated. New technical advances in mapping R-loops, such as DNA–RNA immunoprecipitation sequencing (DRIP-seq) and ribonucleases H1 (RNaseH1) chromatin immunoprecipitation (R-ChIP)^{6,7}, have allowed us to assess the distribution and dynamics of R-loops throughout the genome and detect changes in pathological contexts. Notably, alterations of R-loop levels are detected in cancer models and are associated with transcription–replication conflicts [G] and genomic instability⁸⁻¹¹, offering insights into the source of genomic instability in cancer cells and a new opportunity for cancer therapy.

In this Review, we first discuss new findings of the sources of R-loops and RNA–DNA hybrids in both physiological and pathological conditions. We then discuss the factors and pathways that generate or resolve R-loops and other RNA–DNA hybrids in various cellular contexts, and effects of R-loops and RNA–DNA hybrids on the genome, including positive and negative effects at active genes, DNA replication forks and DNA damage sites, are also discussed. Finally, we discuss the causes and consequences of aberrant R-loops in cancer cells, and the opportunity to target aberrant R-loops in cancer therapy. Through this Review, we aim to provide an updated and integrated view of the fast-evolving research of R-loops and RNA–DNA hybrids.

[H1] Sources and distribution of hybrids

The sources of R-loops and RNA–DNA hybrids, as well as their functions, are distinct in various cellular, chromosomal and sequence contexts. In this section, we discuss how R-loops and RNA–DNA hybrids form naturally during transcription, DNA replication and DNA repair throughout the genome. Furthermore, R-loops accumulate in specific regions of the genome, such as at telomeres and centromeres, and in mitochondrial DNA.

[H2] Co-transcriptional R-loops

During replication, the nascent RNA very transiently anneals to the DNA template within the active site of the the RNA polymerase, giving rise to a short, transient RNA–DNA hybrid, which is resolved by release of the nascent RNA through a dedicated channel¹². R-loops form when nascent RNA transiently re-anneals back onto the template behind the RNA polymerase. At genes transcribed by RNA polymerase II (Pol II), which include the protein-coding genes, co-

transcriptional R-loops mainly accumulate at promoters and transcription start sites (TSSs)¹³ and at transcription termination sites (TTSs)^{14,15}. R-loops at promoters are involved in transcription activation¹⁶, and R-loops at Pol II pause sites near G-rich 'terminator' elements aid transcription termination and formation of gene-repressive chromatin^{14,15} (Fig. 1a). In immunoglobulin genes, R-loops also accumulate in the highly repetitive, GC-rich switch regions, where they aid class switch recombination [G] ¹⁷.

Both increased Pol II transcription^{9,18,19} and conversely Pol II pausing²⁰, have been linked to Rloop accumulation. Increased Pol II transcription generates more RNA that can form R-loops, whereas Pol II pausing increases the chance of RNA hybridization with the DNA template. Active ribosomal DNA repeats are highly transcribed by Pol I, and accumulate R-loops that can interfere with Pol I transcription even in normal conditions²¹. High levels of R-loops, particularly in the 5' regions of the 18S genes, correlate with Pol I pile-up²¹. Pol II transcription of the intergenic spacers between ribosomal DNA genes also generates R-loops that limit Pol I transcription of intergenic non-coding RNAs, which could otherwise disrupt nucleolar structure²². The 5S rRNA genes, which are transcribed by Pol III, are also prone to R-loop formation in normal growth conditions, and other Pol III genes such as tRNA genes accumulate R-loops only in absence of RNase H activity (see below)²³.

[H2] R-loops at telomeres and centromeres

Telomeric repeat-containing RNA (TERRA) is a long non-coding RNA (lncRNA) transcribed by Pol II from telomeres and sub-telomeric regions^{24,25}. TERRA contains the UUAGGG repeats, which can hybridize with the C-rich strand of telomere DNA [G]. In human cells, TERRA associates with telomeres²⁴, interacts with a number of telomere-binding proteins and chromatin modulators such as telomere repeat binding factor 2 (TRF2) and ATRX, and is important for telomere maintenance²⁶. TERRA-generated R-loops are detected at telomeres in human cells²⁷ (Fig. 1b). A recent study showed that RNA consisting of the UUAGGG repeats is sufficient to form telomeric R-loops in *trans* through a RAD51 recombinase-mediated mechanism, and that the formation of telomeric R-loops increases telomere fragility²⁸. Thus, telomeric R-loops allow TERRA to associate with and maintain telomeres, but they also generate genomic instability. The proper regulation of TERRA and telomeric R-loops is likely important for the proper function of telomeres.

Similar to telomeres, centromeres are also transcribed by Pol II²⁹. RNA transcripts of centromeric and pericentric satellite DNA are detected in human cells. The RNA transcripts of a satellite repeats associate with centromeres^{30,31}, suggesting that they form R-loops in *cis* (Fig. 1b). Centromeric RNAs are required for the association of histone H3-like centromeric protein A (CENPA) and centromere protein C (CENP-C) with centromeres^{30,31}, rendering these RNAs important structural components of centromeres. Centromeric R-loops are readily detected in mitotic chromosomes³². The kinase ataxia telangiectasia mutated and Rad3-related (ATR) is recruited to centromeres during mitosis in an R-loop-dependent manner, allowing ATR to promote Aurora B activation through CHK1³² (Fig. 1b). This mitosis-specific and centromere-specific mechanism of ATR activation ensures accurate microtubule-kinetochore attachment and faithful chromosome segregation. However, although centromeric R-loops are important for centromere assembly and function, they are also a source of replication stress. In S phase, CENPA is required for suppressing transcription and R-loops at centromeres, thereby protecting centromeres from DNA damage³³. Thus, centromeric R-loops have both positive and negative effects on centromeres, and the levels of centromeric R-loops and their associated proteins may be intricately orchestrated during the cell cycle.

[H2] Hybrids generated during replication

The most prominant source of replication-associated RNA–DNA hybrids is lagging-strand replication, during which the DNA polymerase α (Pol α)-primase complex synthesizes an 8–10-nucleotide-long RNA primer every ~200 nucleotides³⁴ (Fig. 2a). However, these hybrids are routinely removed during Okazaki fragment maturation, when the RNA primers and the majority of their Pol α -synthesized DNA extensions are displaced by DNA polymerase δ and the resulting flaps are removed by flap endonuclease 1 (FEN1), acting alone or with DNA2 (Ref.³⁵). RNA-primer-based RNA–DNA hybrids might also be produced during bypass of DNA lesions and repriming by Pol α -primase or DNA-directed primase/polymerase protein (PrimPol), although in this case the mechanism of RNA–DNA hybrid removal is much less certain^{36,37} (Fig. 2a). Another form of RNA–DNA hybrid that is frequently generated during DNA replication is a

misincorporated ribonucleotide. The high concentration of ribonucleotides in the cell leads to their misincorporation by replicative DNA polymerases at an estimated rate of once every 7.6 kb, which would equate to 1 million sites per cell division³⁸. Misincorporated ribonucleotides are proposed to be the most frequent genomic lesion³⁸ and require ribonucleotide excision repair (RER) for their removal³⁹ (Fig. 2a).

[H2] DNA damage induced hybrids

In human cells, RNA–DNA hybrids are detected at sites of DNA double-stranded breaks (DSBs). These hybrids are formed by RNA-DNA hybridization on the ssDNA overhangs of DSBs or by invasion of RNA into dsDNA. Several types of RNA contribute to the hybrids at DSBs. Dicergenerated and Drosha-generated small RNAs are reported to form hybrids around DSBs^{40,41}. DNA damage-induced long non-coding RNAs (dilncRNAs) also form hybrids at DSBs⁴². The synthesis of dilncRNAs is linked to Pol II recruited to DSBs by the MRE11-RAD50-NBS1 (MRN) complex^{42,43} (Fig. 2b). Pol III also contributes to RNA synthesis at DSBs⁴⁴. These findings suggest that de novo RNA synthesis at DSBs promotes the formation of RNA-DNA hybrids. However, genome-wide mapping of RNA-DNA hybrids following induction of DSBs at specific sites reveals that the increase in hybrid levels primarily occurs in transcriptionally active regions⁴⁵, suggesting that preexisting RNA transcripts have a predominant role in the formation of damageinduced RNA-DNA hybrids (Fig. 2b). Consistently, when DSBs and single-stranded breaks (SSBs) are induced by reactive oxygen species (ROS) at a specific chromosomal locus, hybrids are only detected in the presence of local transcription⁴⁶. It was proposed that the pausing of Pol II by DSBs and SSBs leads to formation of R-loops^{4,46}. How Pol II is paused by DSBs and SSBs is not understood. The exposure of ssDNA in R-loops at active genes may also allow de novo RNA synthesis⁴, thereby contributing to the formation of RNA–DNA hybrids at DNA damage sites.

[H2] Mitochondrial R-loops

R-loops also accumulate in mitochondrial DNA (mtDNA), which consists of a heavy (H) strand and a light (L) strand with distinct nucleotide compositions⁴⁷. R-loops are detected across the control region of mtDNA [G], and the 3' ends of the RNAs in R-loops coincide with ori-H, the origin of heavy-strand synthesis. The formation of R-loops at ori-H displaces ssDNA and initiates RNA priming for heavy-strand synthesis. After the nascent heavy strand is extended beyond oriL, the synthesis of the light strand is initiated from ori-L⁴⁸, allowing mtDNA to be fully duplicated. Ribonuclease H1 (RNaseH1) is crucial for R-loop processing and DNA replication in mitochondria⁴⁹. Although R-loops are important for mtDNA replication, high levels of R-loops cause mtDNA instability. The mitochondrial degradosome complex, which consists of the helicase SUV3 and the ribonuclease polyribonucleotide nucleotidyltransferase 1, prevents the accumulation of pathological R-loops in mitochondria⁵⁰.

[H2] R-loops generated during genome editing

In addition to the naturally occurring R-loops and RNA–DNA hybrids, R-loops can be artificially generated during genome editing by CRISPR–Cas^{51,52}. In CRISPR–Cas systems, Cas proteins in complex with small guide RNAs (gRNAs) recognize and cleave target DNA sequences. During the recognition of target DNA, the gRNA invades dsDNA and forms RNA–DNA hybrids. The RNA–DNA hybrids formed by Cas9 and gRNAs are about 16–20-nucleotide long. The crystal structure of the Cas9–gRNA–dsDNA complex shows that the RNA–DNA hybrid is largely buried inside the Cas9 protein⁵³. The formation of these RNA–DNA hybrids displaces the non-target strand of DNA and positions the RNA–DNA hybrid and the ssDNA near the active sites of the HNH and RuvC nuclease domains of Cas9, respectively⁵³. The ability of CRISPR–Cas systems to recognize target DNA by forming R-loops also enables them to mediate base editing and prime editing^{54,55}. Other Cas proteins also use R-loops to recognize and cleave targets^{56,57}.

[H1] Impact of hybrids on genomic stability

Although R-loops and RNA–DNA hybrids have contributing roles in many cellular processes, they are also a source of genomic instability when they accumulate at abnormally high levels, or in the wrong context. For example, aberrant R-loops impose a threat to replication forks and increase the nucleolytic cleavage of DNA.

[H2] R-loop-replication fork collisions

Collisions between R-loops and replication forks are a source of DNA damage in bacteria, yeast and human cells⁵⁸⁻⁶⁰. Using R-loop-forming episomes carrying a unidirectional Epstein-Barr virus (EBV) replication origin, head-on collisions of R-loops and replication forks were shown to generate more DNA damage than co-directional collisions⁵⁹ (Fig. 3a). Head-on collisions also

increase R-loop levels in the episomes and in the genome, whereas co-directional collisions decrease R-loop levels⁵⁹. Furthermore, the intermediates generated by head-on collisions or codirectional collisions are processed differently⁵⁹. Head-on collisions were suggested to lead to stalling of replication forks, whereas co-directional collisions allow the CDC45–MCM–GINS (CMG) replicative helicase to unwind RNA–DNA hybrids. Notably, only head-on collisions activate the ATR pathway, which orchestrates the cellular response to DNA replication stress⁵⁹. Subsequent studies revealed that the activation of ATR by R-loops requires replication forks are forced to undergo reversal and MUS81-mediated processing when they encounter R-loops⁶¹. ATR inhibition in cells with high levels of R-loops leads to increased MUS81-dependent DNA damage⁶¹, suggesting that R-loop-induced DNA damage arises from MUS81 cleavage of stalled forks, R-loops, or both.

R-loops could interfere with replication fork progression through a number of mechanisms. The RNA–DNA hybrids and paused Pol II in R-loops or the positive supercoiling induced by R-loops may interfere with the progression of replication forks and cause replication fork collapse. Topoisomerase 1 (TOP1), which relaxes DNA supercoiling, suppresses DNA damage at TTSs where head-on collisions occur⁶² (Fig. 3b). The displaced ssDNA in R-loops can potentially form various types of secondary structures. In sequences with high GC skew, the displaced G-rich ssDNA could form G-quadruplexes [G] (G4s), which are barriers to replication forks⁶³. At telomeres, G4s are a source of replication stress; defects in helicases that resolve G4, such as regulator of telomere elongation helicase 1 (RTEL1), Fanconi anemia group J protein (FANCJ) and Bloom syndrome protein (BLM) increase genomic instability⁶⁴⁻⁶⁶. R-loops also induce genomic instability at DNA trinucleotide repeats, possibly by promoting the formation of DNA secondary structures^{67,68}.

R-loops are associated with gene-repressive histone modifications and chromatin compaction, which could impose replication stress and genomic instability. At TTSs, which are prone to headon collisions⁶², R-loops induce gene-repressive di-methylation of histone H3 Lys9 and recruitment of heterochromatin protein 1 (HP1)¹⁵ (Fig. 3c). R-loops are also associated with high levels of phosphorylated histone H3 Ser10 and with chromatin compaction at centromeres and pericentromeric regions⁶⁹ (Fig. 3c), which are prone to replication stress³³. Notably, depletion of the linker histone H1, which promotes chromatin compaction, suppresses R-loops and replication-associated genomic instability in *Drosophila melanogaster*, supporting the idea that R-loop-associated chromatin compaction contributes to collisions with replication forks and to genomic instability⁷⁰.

[H2] Nucleolytic processing of R-loops

Xeroderma pigmentosum group F (XPF) and XPG, two structure-specific endonucleases involved in transcription-coupled nucleotide excision repair (TC-NER), generate DNA damage in an Rloop-dependent manner⁷¹. Other TC-NER factors, such as XPB and XPD, are also required for the formation of R-loop-dependent DNA damage⁷¹. These results suggest that TC-NER proteins have a role in processing R-loops. Camptothecin induces transcription-dependent DSBs through XPF and XPG, suggesting that they cleave co-transcriptional R-loops⁷². XPG is not required for the activation of ATR by R-loops⁶¹, indicating that it acts independently of replication.

The displaced ssDNA in R-loops is a potential substrate of DNA modifying enzymes. During immunoglobulin class-switch recombination in B cells, activation induced cytidine deaminase (AID) converts cytosines to uracil in the ssDNA exposed by R-loops in the switch regions¹⁷. The uracil generated by AID is processed by the uracil-DNA glycosylase into abasic sites, which directly stall DNA polymerases and also give rise to SSBs through base excision repair, which are converted into DSBs during replication (Fig. 3d). In addition to AID, other cytidine deaminases such as APOBEC3A and APOBEC3B may also attack the ssDNA in R-loops and generate abasic sites and DNA breaks⁷³.

[H1] R-loop suppression and resolution

To ensure that R-loops are properly regulated and avoid genomic instability, multiple pathways act in concert to prevent aberrant R-loop formation and remove excessive R-loops.

[H2] RNaseH1 and RNaseH2

RNase H proteins are endoribonucleases that specifically cleave the RNA strand in RNA–DNA hybrids. In eukaryotes, this activity is shared between RNaseH1 and RNaseH2 (Fig. 4a, b; Table

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1). RNaseH1 removes R-loops in the nucleus and in the mitochondria, where it is essential for mitochondrial DNA replication⁷⁴. RNaseH2 is purely nuclear, consists of three subunits, and interacts with proliferating cell nuclear antigen (PCNA), which may help to recruit it to sites of DNA replication and repair⁷⁵. The RNaseH2A subunit provides the enzymatic activity, and the RNaseH2B subunit is required for PCNA interaction and nuclear localization^{75,76}. RNaseH2 can remove R-loops and misincorporated ribonucleotides from DNA, the latter through its action in the RER pathway³⁹. RER is initiated by ribonucleotide incision by RNaseH2, followed by DNA repair by the replication machinery, which involves strand displacement DNA synthesis by DNA polymerases δ (Pol δ) and Pol ε , flap cleavage by FEN1 or exonuclease 1 (EXO1), and DNA ligation by DNA ligase I (LIG1)³⁹. Although FEN1 provides the dominant RNA–primer removing activity, RNaseH2 is also implicated in RNA primer removal⁷⁵. Work in budding yeast suggests that RNaseH2 is cell cycle regulated and more highly expressed and active in late S and G2 phase, when it is required for RER and R-loop removal⁷⁷. RNaseH1 is active independently of cell cycle stage, but may require R-loop-induced stress for its recruitment to chromatin⁷⁷.

[H2] R-loop suppression in transcription

RNA processing suppresses R-loop accumulation at all stages of transcription (Fig. 4a; Table 1). During RNA synthesis, the intrinsic transcript-cleavage activity of Pol II, which requires transcription elongation factor S-II (TFIIS; also known as TCEA1), suppresses R-loop formation by reducing Pol II pausing and backtracking [G]⁷⁸. During co-transcriptional RNA processing, splicing factors such as serine/arginine-rich splicing factor 1 (SRSF1)⁷⁹, splicing factor proline and glutamine rich (SFPQ)⁸⁰ and splicing factor 3B subunit 1 (SF3B1)⁸¹ bind the nascent RNA as it leaves the RNA polymerase active site and prevent it from re-annealing to the template. The transcription export (TREX) complex, which contains the core components of the THO complex and additional RNA binding proteins⁸², suppresses R-loop formation by promoting the assembly and nuclear export of the messenger ribonucleoprotein⁸³. The TREX-2 complex also suppresses R-loop formation. Cleavage of lncRNAs mediated by SPT6 and the Integrator complex [G] prevents R-loop formation by extended lncRNA transcripts⁸⁵. Degradation of some enhancer RNAs by the RNA exosome [G] prevents R-loop accumulation at the corresponding enhancers⁸⁶.

TOP1 resolves both positive and negative supercoiling during transcription and DNA replication (Table 1). Loss of TOP1 results in R-loop accumulation in active genes, possibly because the increase of negative supercoiling behind elongating Pol II favors RNA–DNA hybrid formation⁸⁷ (Fig. 4a). The TOP1 inhibitor camptothecin induces cotranscriptional R-loops⁷². TOP1 suppresses R-loop-associated DNA damage at TTSs where head-on collisions of R-loops and replication forks occur⁶².

The helicase senataxin is also involved in R-loop suppression at TTSs (Fig. 4a; Table 1)^{14,88}. The stability of senataxin is regulated by the deubiquitinase by the ubiquitin carboxyl-terminal hydrolase USP11, which also controls R-loop levels⁸⁹. Depletion of the RNA helicase aquarius increases R-loops and genomic instability, suggesting that Aquarius unwinds RNA-DNA hybrids⁷¹. Several helicases of the DEAD-box (DDX) and DEAH box (DHX) families are involved in R-loop suppression. DDX39B unwinds RNA-DNA hybrids in vitro and suppresses cotranscriptional R-loops genome-wide90. DDX21 also unwinds RNA-DNA hybrids and it cooperates with sirtuin-7 to suppress R-loops at specific genes⁹¹. DHX9 unwinds R-loops *in vitro*, associates with R-loops in cells, and suppresses camptothecin-induced cotranscriptional Rloops^{92,93}. Paradoxically, DHX9 was shown to also promote R-loop formation in cells with impaired splicing⁸⁰. In fission and budding yeasts, the senataxin homolog Sen1⁹⁴ and the helicases Pif1 and Rrm3 (Ref.^{95,96}) are required for proper transcription termination at tRNA genes and for maintaining genomic stability, although the functions of these proteins are not linked with Rloops^{94,95}. Together, these findings suggest that cells use multiple RNA–DNA helicases to suppress cotranscriptional R-loops and their associated genomic instability (Table 1). Whether and how these helicases function in a context-specific manner and how they are regulated remains largely unknown.

A number of DNA repair proteins are also involved in the suppression of cotranscriptional R-loops (Table 1). Breast cancer type 1 susceptibility protein (BRCA1) associates with TTSs and recruits senataxin to suppress R-loops⁸⁸. In human breast luminal epithelial cells with *BRCA1* mutations, R-loops accumulate at promoter-proximal Pol II pause sites; and suppression of Pol II pausing alleviates R-loop accumulation in mouse BRCA1 deficient cells⁹⁷. Loss of BRCA2 also leads to increased Pol II pausing and R-loop accumulation at promoter-proximal sites⁹⁸. BRCA2 binds the

TREX-2 complex to suppress R-loops⁸⁴. In addition, BRCA2 recruits DDX5 to suppress R-loops in transcribed regions⁹⁹. In BRCA1-deficient or BRCA2-deficient cancer cells, DHX9 is recruited to R-loops by RNF168 to keep R-loops at tolerable levels¹⁰. Several proteins of the Fanconi anemia repair pathway, such as FANCD2, FANCI and FANCA, are important for R-loop suppression^{100,101}. The FANCD2–FANCI complex interacts with RNA–DNA hybrids and recruits RNA-processing factors, such as heterogeneous nuclear ribonucleoprotein U and DDX47, to remove R-loops^{102 103}. The ssDNA-binding replication protein A (RPA) is also present at R-loops, possibly on the displaced ssDNA¹⁰⁴. RPA recruits and stimulates RNaseH1 to prevent excessive R-loop accumulation. Structure-specific nucleases such as XPF, XPG, FEN1 and CtIP (also known as RBBP8) are also involved in the processing and removal of cotranscriptional R-loops^{72,105}.

[H2] R-loop responses during DNA replication

The suppression of cotranscriptional R-loops is important for preventing collisions of R-loops and replication forks. For example, TOP1 resolves R-loop associated DNA supercoiling at TTSs to suppress the induction of DNA damage by head-on collisions of replication forks and R-loops⁶². A number of DNA repair proteins are involved in the response to collisions of replication forks with R-loops (Table 1). The helicase FANCM unwinds telomeric RNA-DNA hybrids in vitro and suppresses replication stress during alternative lengthening of telomeres (ALT)^{106,107}. BLM also unwinds RNA-DNA hybrids in vitro and suppresses R-loops in cells¹⁰⁸. BRCA1, FANCM and BLM act in concert to resolve R-loop-induced replication stress at ALT telomeres¹⁰⁹. The helicase senataxin associates with replication forks and protects forks in transcribed genes¹¹⁰. In response to DSB formation in such genes, senataxin is recruited to DSBs to remove RNA-DNA hybrids⁴⁵. DDX1, DDX5, DDX21 and DHX9 interact with ATPase family AAA domain-containing protein 5 (ATAD5), which offloads PCNA replication forks¹¹¹. Depletion of this group of helicases increases R-loops in S phase and reduces DNA synthesis, suggesting that they remove R-loops during replication. DDX19 enters the nucleus following DNA damage in an ATR-dependent manner to remove R-loops¹¹² (Fig. 4b). Structure-specific nucleases such as MUS81 and MRE11 are also involved in the suppression of R-loops, possibly by cleaving DNA structures induced by collisions of R-loops and replication forks^{61,113}.

Several proteins involved in DNA damage responses, such as ataxia telangiectasia mutated (ATM), ATR, CHK1 and CHK2, suppress R-loops¹¹⁴. These checkpoint proteins were proposed to be important for preventing collisions of replication forks with R-loops or for repairing collapsed forks¹¹⁴. Head-on collisions between replication forks and R-loops activate ATR signaling, whereas codirectional collisions activate ATM, suggesting that there could be different replication-restart pathways for head-on versus codirectional collisions⁵⁹. Replication forks undergo reversal at R-loops, which generates substrates for nucleolytic processing by MUS81–EME2 (the MUS81 complex most active in S phase)^{61,115} (Fig. 4b). MUS81 processing of reversed forks at R-loops activates ATR⁶¹, allows Pol II to passage through stalled forks, and eventually promotes fork restart using strand annealing by RAD52 and nick ligation by LIG4¹¹⁵ (Fig. 4c). Such a process is reminiscent of, but distinct from break-induced replication [G] (BIR)¹¹⁵.

There is evidence that replication forks can bypass R-loops, either through RNA-DNA hybrid unwinding by the replicative helicase or by repriming replication downstream of the R-loop (Fig. 4b). As discussed above, codirectional collisions of replication forks and R-loops reduce R-loop levels⁵⁹, possibly because the helicase MCM (part of the CMG complex) can unwind RNA-DNA hybrids on the leading strand. PrimPol promotes replication fork progression through R-loops at GAA repeats; R-loops could promote the formation of a secondary structures such as hybrid triplex structure formed between a GAA repeat and the opposite CTT repeat. If the GAA repeat was on the leading strand template such a triplex structure could stall the leading-strand DNA polymerase¹¹⁶ (Fig. 4b). The repriming by PrimPol on the leading strand allows replication forks to avoid stalling. PrimPol also limits the formation of R-loops in GAA repeats, suggesting that in absence of PrimPol, excessive ssDNA could become accessible for RNA polymerase to produce more RNA–DNA hybrids¹¹⁶. As ssDNA generally forms at stalled forks, RNA polymerases could also promote R-loop formation in response to other replication-blocking lesions. Behind the progressing fork, the PCNA offloader ATAD5 can restrict R-loop formation, possibly by preventing PCNA from interfering with transcription, as well as by recruiting RNA-DNA helicases as described above¹¹¹.

[H2] Chromatin modulators

In addition to factors that function in transcription, replication and DNA repair machineries, several chromatin reorganizing activities are implicated in preventing R-loop accumulation (Table 1). The SWI/SNF^{100,117} and INO80¹¹⁸ chromatin remodeling complexes and the histone chaperone complex facilitates chromatin transcription (FACT)¹¹⁹ aid replisome progression in presence of Rloops (Fig. 4b). FACT is proposed to replace histones at sites of transcription-replication conflicts, possibly in response to R-loop-induced formation of repressive histone modifications^{69,119}. INO80 is similarly suggested to resolve R-loops by facilitating chromatin decompaction but could also help resolve R-loops through its interaction with RNA-DNA helicases such as DDX5 or by facilitating removal of stalled Pol II¹¹⁸. SWI/SNF might suppress R-loops by controlling the accessibility of chromatin to the DNA repair machinery¹⁰⁰. The mechanism is still unclear, as SWI/SNF increases chromatin accessibility genome-wide, but it limits chromatin accessibility at sites of transcription-replication conflicts¹⁰⁰. SWI/SNF acts in the Fanconi anemia pathway together with senataxin and FANCD2, and in absence of SWI/SNF R-loops accumulate specifically in S phase. This result suggests that the function of SWI/SNF is specific to replicating cells, such as removing R-loops at stalled replication forks¹⁰⁰. SWI/SNF is also implicated in recruiting TOP2A to R-loop sites¹¹⁷.

Several chromatin modifiers including Polycomb repressive complex 1 (PRC1)^{120,121}, the histone Lys acetyltransferase KAT8¹²² and the histone deacetylase (HDAC) SIN3A complex⁸³ have been implicated in suppressing R-loops (Fig. 4; Table 1). As PRC1 and SIN3A function as transcription repressors, R-loop accumulation in their absence could directly result from increased transcription. However, SIN3A also suppresses R-loop accumulation through a physical interaction with the THO complex subunit THOC1 (Ref.⁸³). The acetyllysine readers bromodomain-containing protein 2 (BRD2) and BRD4 suppress R-loop formation, either by preventing Pol II pausing (BRD4) or through recruitment of TOP1 (BRD2)¹²³⁻¹²⁵. The majority of these factors have known roles in DNA replication and the DNA damage response (reviewed in¹²⁶) (Fig. 4a,b), which promotes R-loop processing during collisions with replication forks. As the cited studies were performed in cycling cells, it is uncertain to what extent the above factors also directly and specifically suppress R-loops during transcription. This question could be further investigated using cells in G1 phase or non-cycling cells.

[H1] Functions of hybrids in DNA repair

Although R-loops can induce genomic instability, R-loops and RNA–DNA hybrids are also induced by DNA damage. The R-loops and RNA–DNA hybrids induced by DNA damage may have several distinct roles in promoting the removal of DNA damage in different contexts.

[H2] Repair of double-strand DNA breaks

Both the RNA generated in transcribed regions before DNA damage and the RNAs induced by DNA damage contribute to the formation of RNA–DNA hybrids at DSBs. The RNA–DNA hybrids at DSBs may promote homologous recombination (HR) through several distinct mechanisms.

[H3] Recruitment of repair proteins. Several types of RNA–DNA hybrids have been proposed to recruit DNA repair proteins to DSBs. DSB-induced small RNAs (diRNAs) or DNA damage response small RNAs (DDRNAs) are small non-coding RNAs generated by Dicer at DSBs^{40,41}. diRNAs form a complex with Argonaute-2 and RAD51, which promotes the localization of RAD51 to DSBs, possibly through diRNA hybridization with DNA¹²⁷ (Fig. 5a). The DDRNAs are derived from damage-induced long non-coding RNAs (dilncRNAs), which are transcribed at DSBs¹²⁸. dilncRNAs form RNA–DNA hybrids at DSBs, thereby promoting the recruitment of the HR proteins BRCA1, BRCA2 and RAD51129 (Fig. 5a). BRCA1 was shown to bind RNA-DNA hybrids, and BRCA1 foci in cells in S and G2 phases were reduced by RNase H treatment, suggesting that BRCA1 may be a sensor of R-loops¹²⁹. When ROS are locally induced at a specific chromosomal locus, R-loops are generated in a transcription-dependent manner⁴⁶, suggesting that RNA transcripts hybridize with DNA at sites of DNA damage (Fig. 5a). The repair of ROSinduced DSBs requires a non-canonical HR pathway mediated by Cockayne syndrome group B (CSB), a protein also involved in TC-NER, and RAD52 and RAD51⁴⁶. Both CSB and RAD52 directly bind RNA-DNA hybrids in vitro and localize to sites of DNA damage in an R-loopdependent manner in cells^{46,130}, suggesting that they are also sensors of DNA damage-induced Rloops. In G2 cells exposed to ionizing radiation or laser microirradiation, RAD52 is recruited to DNA damage sites in a transcription and RNA-DNA hybrid-dependent manner¹³¹. Furthermore, RPA may act as a sensor of R-loops by recognizing the displaced ssDNA¹⁰⁴. Collectively, these findings suggest that R-loops or RNA-DNA hybrids at DSBs may directly or indirectly promote

the recruitment of a number of DNA repair proteins, which may enhance DSB repair by increasing the local concentrations of these repair factors.

DNA end resection **[G]** is important for the choice of DNA repair pathways. Whereas resection is required for HR, it antagonizes canonical non-homologous end joining. Whether and how the R-loops and RNA–DNA hybrids at DSBs affect the resection of DNA ends is still controversial. In yeast, overexpression of RNaseH1 destabilizes RNA–DNA hybrids at DSBs and induces excessive resection¹³², suggesting that hybrids restrict resection. In human cells, however, RAD52, which localizes at DSBs in a hybrid-dependent manner, recruits the XPG to cleave R-loops and promotes ssDNA formation (Fig. 5b)¹³¹. Loss of Pol III, which generates RNA–DNA hybrids at DSBs, also reduces resection ^{(Fig. 5b)⁴⁴}. Furthermore, DHX9 recruits BRCA1 to Pol II-generated RNA at DSBs to enhance resection¹³³. These findings suggest that hybrids promote resection in human cells. Nonetheless, depletion of senataxin did not significantly affect resection⁴⁵. Notably, reducing hybrids at DSBs by RNaseH1 overexpression in yeast or depletion of Pol III in human cells results in sequence loss around DSBs^{44,132}, suggesting that hybrids protect ssDNA overhangs.

[H3] RNA-templated or bridged repair. In budding yeast, RNA transcripts can serve as template for DSB repair when RNaseH and reverse transcriptase activities are absent¹³⁰. The RNAtemplated repair of DSBs requires Rad52, which can promote the formation of RNA–DNA hybrids and strand exchange between homologous ssRNA and ssDNA^{130,134}. RNA transcripts were hypothesized to hybridize with 3' overhangs at DSBs, thereby allowing RNA-templated DNA synthesis (Fig. 5c). The resulting extended ssDNA overhang then captures the ssDNA from the opposite DSB end, joining the two ends together. In human cells, the repair of ROS-induced DSBs, which is dependent on RNA–DNA hybrids, can occur in G0 and in G1 (before DNA replication), suggesting that it may also involve RNA templates¹³⁵. Using purified proteins and RNA, RAD52 was shown to promote RNA-templated DNA recombination *in vitro*¹³⁶. A recent study shows that DNA polymerase θ , which is required for a DSB repair pathway termed theta-mediated DNA end joining can efficiently synthesize DNA using RNA as a template and promote RNA-templated DSB repair in a cell-based reporter assay¹³⁷. Although RNA–DNA hybrids can enable RNAtemplated repair of DSBs, whether this type of repair counts for a significant portion of repair activity in cells remains unknown. In addition, RNA transcripts could potentially function as a molecular bridge to hold two DNA ends together. In the presence of RAD52, an RNA oligo that can anneal with ssDNA on both sides of a DSB enables the joining of DNA ends *in vitro*¹³⁶. Whether RNA can function as a bridge during DSB repair in cells and how important this mechanism is *in vivo* remains unclear.

[H3] DR-loops. Using HR reporter assays, in which local transcription at DSB sites can be switched on and off, local transcription was shown to substantially stimulate HR¹³⁸. Importantly, even in the absence of local transcription, tethering homologous RNA transcripts to the vicinity of DSBs by nuclease-inactive Cas9 enhances HR, recapitulating the effects of transcription¹³⁸. These findings suggest that preexisting RNA transcripts at transcriptionally active sites have an important role in HR once DSBs emerge. Furthermore, the stimulation of HR by transcription or tethered RNA transcripts is dependent on RAD51 associated protein 1 (RAD51AP1) and USP1 associated factor 1 (UAF1), which form a complex that promotes R-loop formation^{138,139}. Surprisingly, DSBinduced and RAD51AP1-generated RNA–DNA hybrids are detected not only at DSBs but also in the DNA serving as the donor DNA [G] for repair, suggesting that DNA–DNA and RNA–DNA hybrids co-exist in donor DNA. Indeed, purified RAD51AP1 and RAD51 can promote the invasion of both ssRNA and ssDNA into donor dsDNA in vitro, generating DR-loops, which contain both DNA-DNA and RNA-DNA hybrids (Fig. 5d). The formation of DR-loops is associated with enhanced RAD51-mediated ssDNA invasion, suggesting that DR-loops are HR intermediates favoring RAD51 function. These findings on DR-loops suggest that RNA is not only a regulator of DNA repair proteins, but also a direct participant in recombination. This role of RNA in HR may be crucial for protecting transcribed regions of the genome.

[H3] RNA modifications. The *N*⁶-methyladenosine (m⁶A) modification of mRNA is associated with R-loops and promotes R-loop suppression and resolution^{140,141}, although it may also facilitate transcription termination through R-loops¹⁴². In response to ultraviolet irradiation damage, m⁶A is rapidly and transiently generated by methyltransferase 3, *N*⁶-adenosine-methyltransferase complex catalytic subunit (METTL3) at DNA damage sites¹⁴³. In the absence of m⁶A, the translesion synthesis **[G]** DNA polymerase Polκ cannot be recruited efficiently to DNA, and ultraviolet damage repair is impaired¹⁴³. METTL3 catalyzes the m⁶A modification of RNA at DSBs, to which

the m⁶A 'reader' YTHDC1 is recruited to stabilize RNA–DNA hybrids¹⁴⁴. Loss of METTL3 leads to defective HR¹⁴⁴.

In response to ROS-induced DNA damage, the 5-methylcytosine (m⁵C) modification of RNA is rapidly induced at DNA damage sites¹⁴⁵ (Fig. 5a). This induction requires local transcription, suggesting that local mRNA is modified by m⁵C¹⁴⁵. RNA–DNA hybrids are directly bound by tRNA aspartic acid methyltransferase 1 (TRDMT1; also known as tRNA (cytosine(38)-C(5))-methyltransferase), which is required for efficient m⁵C formation at DNA damage sites and repair of ROS-induced DSBs. Both RAD52 and RAD51 are recruited to sites of ROS-induced DNA damage in a TRDMT1-dependent manner¹⁴⁵ (Fig. 5a). RAD52, which is required for recruiting RAD51 to ROS-induced DSBs, has higher affinity for RNA–DNA hybrids containing m⁵C-modified RNA. These results suggest that a TRDMT1–m⁵C–RAD52 axis operates on R-loops to promote the repair of ROS-induced DSBs. In addition to RNA methylation, adenosine-to-inosine RNA editing [G] is increased near DSBs¹⁴⁶. This response is mediated by adenosine deaminases acting on RNA 2 (ADAR2; also known as double-stranded RNA-specific editase 1), which is required for efficient HR and cellular resistance to DNA damage.

[H2] Repair of collapsed replication forks

As discussed above, BIR or related pathways are involved in restarting replication forks stalled by R-loops. At sites of DNA damage or genomic instability, DNA damage-induced R-loops may stall replication forks and induce BIR, allowing BIR to alleviate the replication stress. For example, in yeast, TERRA-generated R-loops accumulate at critically short telomeres, promoting the DDR to extend telomeres¹⁴⁷. In human cancer cells, telomeric R-loops promote the BIR-related ALT pathway to extend telomeres^{148,149}. Local induction of ROS at telomeres triggers an R-loop-dependent CSB–RAD52–DNA polymerase delta subunit 3 (POLD3) pathway to remove DNA damage at telomeres, thereby linking R-loops with POLD3-mediated BIR¹⁵⁰. A number of proteins involved in these BIR (and related) pathways, such as RAD52, RAD51AP1 and CSB, can directly bind RNA–DNA hybrids. Although damage-induced R-loops may induce BIR by stalling replication forks, whether and how R-loops directly participate in the process of BIR is still unknown.

[H2] Repair of single-strand DNA breaks

A complex of BRCA1 with RNAi machinery was recently shown to generate single-stranded DNA-damage-associated small RNAs (sdRNAs) 24–40 nucleotides-long from RNA transcripts in response to DNA damage¹⁵¹ (Fig. 5e). These sdRNAs interact with partner and localizer of BRCA2 (PALB2) and RAD52 to promote the repair of SSBs at the TTSs of active genes. R-loops were speculated to be involved in the formation and function of sdRNAs. Consistent with this hypothesis, sdRNAs function at the loci from which they are generated, and R-loops are detected at these loci. Further studies are needed to determine whether R-loops directly participate in sdRNA generation and function.

[H2] Resolution of hybrids during DNA repair

Although RNA–DNA hybrids have positive roles in DNA repair, they must be removed before the completion of repair. In yeast, RNaseH is required for suppressing hybrids at DSBs and for efficient HR^{132,152}. However, other studies argue that RNaseH is not required for DSB repair¹⁵³. In human cells, BRCA2 recruits DDX5 and RNaseH2 to DSBs to remove RNA–DNA hybrids^{129,154}. Furthermore, senataxin promotes RAD51 recruitment to DSBs and reduces illegitimate rejoining of distant DSBs, supporting the idea that hybrid removal is important for accurate DSB repair⁴⁵.

[H1] Pathological R-loops

Aberrant accumulation of R-loops can cause increased replications stress and/or DNA damage, which can underlie cancer susceptibility or neurodegeneration. R-loop levels could be elevated in both types of disorders owing to loss of transcription regulation or R-loop suppressing activities. The genomic location of R-loop accumulation might vary depending on the nature of the genetic alterations²⁰, and on whether the affected tissue is proliferating or post-mitotic.

[H2] Cancer-associated R-loops

A number of oncogenic events can increase the levels of R-loops and associated genomic instability. Activated oncogenes and bacterial carcinogens can increase R-loop formation, which elevates replication stress and genomic instability (Fig. 6a). Oncogenic estrogen signaling¹⁸, the HRAS^{V12} oncoprotein¹⁹ and the EWS-FLI1 fusion oncoprotein⁹, all increase R-loop levels genome-wide, promoting replication stress and DNA damage. HRAS-induced and estrogen-

induced R-loop formation is linked with increased transcription, especially at the target genes of estrogen and RAS signaling^{18,19}. EWS-FLI1, a fusion of EWS RNA binding protein 1 (EWSR1) and Friend leukemia integration 1 transcription factor (FLI1), is also proposed to increase transcription by stimulating Pol II activity; depletion of wild-type EWSR1 to mimic the effects of EWS-FLI1 fusion induced R-loops especially in highly expressed regions⁹ (Fig. 6a). Expression of the SS18-SSX1 oncoprotein, a fusion of the SS18 (also known as SSXT) subunit of BAF chromatin remodeling complex (mammalian SWI/SNF) with SSX1, which drives synovial sarcoma, also increased R-loop levels¹¹. Although the underlying mechanism is not known, it is of note that SS18-SSX alters SWI/SNF targeting to chromatin¹⁵⁵. *Heliobacter pylori* infections, which act as a carcinogen promoting gastric cancer, also increase RNA synthesis by activating NF-κB signaling and thereby inducing R-loop-dependent replication stress¹⁵⁶.

However, elevated RNA synthesis is not the only way in which oncogenes can increase R-loop levels. Suspected oncogenic mutations in the genes encoding the splicing factors SRSF2 and U2 small RNA auxiliary factor 1 (U2AF1; also known as U2AF35), which are prevalent in myelodysplastic syndrome (MDS)^{20,104,157}, were found to induce R-loops and replication stress, possibly through increasing Pol II pausing or other mechanisms²⁰ (Fig. 6b). Finally, the oncogenes MYCN¹⁵⁸ and MDM2 (Ref.¹²¹) have been reported to suppress R-loop formation, acting in concert with BRCA1¹⁵⁸ or the chromatin modifier PRC1 (Ref.¹²¹), respectively. The R-loop suppressing effects of MYCN and MDM2 may allow cancer cells to tolerate high levels of transcription and replication stress. The discrepancy between MYCN and other oncogenes such as HRAS^{V12} might be explained by timing, as MYCN suppresses R-loops in the first few hours following oncogene induction¹⁵⁸, whereas HRAS^{V12}-induced R-loops were detected after three days of induction¹⁹. It will be interesting to investigate the effects of prolonged MYCN overexpression on R-loop levels.

Factors involved in the repair of replication-associated DNA damage such as FANCM, the MRN complex^{113,159}, BRCA1⁸⁸ and BRCA2⁸⁴, and factors required for activation of cell cycle checkpoints such as ATM¹⁶⁰, function as tumor suppressors. The R-loop suppressing functions of these proteins^{84,88,113,159,160} could be important in preventing conflicts between R-loops and replication, thereby limiting cancer-promoting genomic instability in proliferating cells (Fig. 6b).

The regulation of TERRA and telomeric R-loops is altered in a subset of human cancers. TERRA is generally upregulated in cancer cells that use the ALT pathway to extend telomeres¹⁶¹. Loss of ATRX, which is a chromatin remodeller acting at subtelomeres and frequently mutated in ALT-positive cancers, increases TERRA levels¹⁶² (Fig. 6c). Whether ATRX loss is the primary cause of TERRA upregulation in ALT-positive cancers remains unclear¹⁶¹. Consistent with the upregulation of TERRA, telomeric R-loops are more abundant in ALT-positive cancer cells²⁷. Loss of ATRX also increases formation of G-quadruplexes at telomeres^{163,164}, which may indirectly contribute to R-loop formation. A recent study showed that inhibition of TERRA transcription diminishes ALT activity¹⁴⁸, supporting the idea that telomeric R-loops drive ALT by inducing replication stress^{27,148,165,166} (Fig. 1b). The helicase FANCM has the ability to unwind telomeric R-loops, and it restricts ALT activity to a tolerable level in ALT-positive cancer cells^{106,107} (Fig. 6d). BRCA1 directly binds TERRA and suppresses telomeric R-loops and DNA damage¹⁶⁷ (Fig. 6d).

Cancer cells can contain variants in the telomeric repeat sequence, and A–C mismatches between DNA and TERRA RNA arising from this variation inhibit RNaseH2 activity. RNA editing by the adenosine-to-inosine editing enzyme ADAR1 (also known as double-stranded RNA-specific adenosine deaminase) converts A–C mismatches to I–C matched base pairs, thereby allowing RNA processing by RNaseH2 specifically at telomeres, which is required for the proliferation of non-ALT or telomerase-activated cancer cells¹⁶⁸. Interestingly, depletion of the helicase RTEL1, which binds the G-quadruplexes formed by TERRA, reduces telomeric R-loops, but increases telomere instability¹⁶⁹. These results suggest that telomeric R-loops need to be maintained at an appropriate level to keep telomeres stable or to not activate ALT above a tolerable level.

[H2] R-loops in the central nervous system

Excessive R-loop formation in post-mitotic neurons could promote transcription-associated DNA damage and genomic instability independently of conflicts with DNA replication, thereby contributing to neurodegenerative diseases and neurodevelopmental disorders. Mutations in DNA repair genes such as *ATM* and *SETX* underlie cerebellar ataxias [G] and are linked to increased DNA damage in neurons^{170,171}. At least one of the affected genes, *SETX*, encodes senataxin, a protein with direct roles in R-loop resolution¹⁴. Increased R-loop formation could plausibly

contribute to DNA damage in these disorders¹⁷⁰. Mouse models of cerebellar ataxias, including ataxia telangiectasia (caused by mutations in *ATM*), ataxia oculomotor apraxia type 1 (AOA1; mutations in aprataxin), AOA2 (mutations in senataxin), and spinocerebellar ataxia with axonal neuropathy (mutations in tyrosyl-DNA phosphodiesterase) all display increased levels of R-loops in proliferating tissues¹⁶⁰ (Fig. 6d). However, in line with the observation that mice tend to be poor models for neurodegeneration that frequently fail to develop the neurological symptoms of the human disorders, R-loops are not detected in the brains of these mice¹⁶⁰. By contrast, human non-proliferating AOA2-derived neural progenitor cells do display increased R-loop signals in nucleoli and the cytoplasm, as well as increased oxidative DNA damage¹⁷⁰. Other *SETX* mutations underlie amyotrophic lateral sclerosis 4 (ALS4), in which R-loops are reduced in gene promoters, which reduces the expression of affected genes and may contribute to disease etiology¹⁷².

Several neurodegenerative diseases such as amyotrophic lateral sclerosis and frontotemporal dementia (ALS/FTD) and Friedreich ataxia are linked with expansions of short guanine-containing repeat sequences, which are prone to forming stable R-loops^{173,174}. Several mechanisms have been proposed to explain how increased R-loop formation in one gene might promote neurodegeneration. R-loops at expanded repeats in *C9orf72*, which underlies ALS, and in the frataxin gene, which underlies Friedreich ataxia, have been linked to impaired Pol II transcription and gene silencing, again suggesting that R-loops could promote pathology by posing blocks to gene expression^{173,174} (Fig. 6d). In addition, RNA binding proteins such as TDP-43 form dysfunctional aggregates in ALS and FTD, which can further promote R-loop accumulation¹⁷⁵. Repeat expansions in *C9orf72* are linked also with P62 accumulation, causing defective ATM signaling, which exacerbates R-loop-mediated DNA damage¹⁷⁶. Mutations in splicing factors can also underlie neurodegenerative disorders. R-loops accumulate specifically in the embryonic neurons of zebrafish lacking the splicing factor SF3B1, where they promote DNA damage and apoptosis⁸¹.

R-loop formation in expanded repeats is implicated also in the neurodevelopmental disorder Fragile X syndrome¹⁷³. Mutations in RNaseH2 or in the deoxynucleoside triphosphate triphosphohydrolase SAMHD1, which are factors involved in R-loop prevention¹⁷⁷, cause Aicardi-Goutieres syndrome, a neuro-inflammatory disorder that has been linked to the presence of DNA

in the cytosol and thus to activation of the cGAS–STING pathway [G] rather than to R-loop accumulation^{178,179}.

[H1] Targeting aberrant R-loops in cancer

Although aberrant R-loop accumulation in cancer cells may promote tumorigenesis by altering gene expression and fueling genomic instability, it also confers a vulnerability to cancer cells that can be exploited therapeutically. Recent studies suggest that cancer cells harboring high levels of R-loops can be preferentially killed by drugs that exacerbate R-loop-associated DNA damage and block the R-loop response.

[H2] Inhibition of the ATR checkpoint

An R-loop dependent ATR response is detected in cells depleted of SRSF1, suggesting that ATR is activated in response to aberrant R-loop accumulation⁶¹. In contrast to the activation of ATR by the replication inhibitor hydroxyurea, ATR activation by R-loops required replication-fork reversal and the endonuclease MUS81–EME2⁶¹. Inhibition of ATR in cells with high levels of R-loops leads to an increase of MUS81-EME1–generated DNA damage, suggesting that ATR functions in a feedback loop to suppress R-loop-associated DNA damage by limiting MUS81 activity⁶¹. ATR also suppresses transcription–replication collisions and promotes fork recovery in cells with high levels of R-loops⁶¹. Together, these results suggest that ATR is a key regulator of the R-loop response.

Several splicing-factor mutations prevalent in MDS and acute myeloid leukemia (AML), such as U2AF1^{S34F} and SRSF2^{P95H}, were shown to increase R-loops in cells^{20,104,157}. An ATR response was induced by these spliceosome mutants in an R-loop-dependent manner. Furthermore, inhibition of ATR in cells expressing the spliceosome mutants increases R-loop-dependent DNA damage, suggesting that ATR protects the cells against aberrant R-loops. Thus, inhibition of ATR is an attractive strategy to exploit the R-loop-associated vulnerability of cancer cells (Fig. 7a). How R-loops are induced by spliceosome mutants and how R-loops contribute to the ATR inhibitor (ATRi) effects is still not fully understood.

[H2] Inhibition of RNA splicing

The splicing-factor mutations found in MDS affect alternative splicing of many genes. RNA splicing inhibitors were developed to kill MDS cells by exacerbating splicing defects. In particular, inhibitors of SF3B1, such as pladienolide B, E7107, and H3B-8800, have been shown to preferentially kill cells expressing splicing-factor mutants¹⁸⁰. Splicing inhibitors were also found to increase R-loops and stimulate the ATR response in cells expressing U2AF1^{S34F} (Ref. ¹⁵⁷) (Fig. 7b). Consequently, splicing inhibitors increase the sensitivity of U2AF1^{S34F}-expressing cells to ATRi¹⁵⁷. Thus, combinations of ATRi and drugs that increase R-loop levels may be effective in treating MDS.

[H2] Nonsense-mediated mRNA decay inhibition

Cells expressing MDS-associated splicing-factor mutants, including U2AF1^{S34F} and SF3B1^{K700E}, were recently shown to be sensitive to an inhibitor of serine/threonine-protein kinase SMG1 (SMG1), which is required for nonsense-mediated mRNA decay **[G]** (NMD)¹⁸¹. SMG1 inhibition or depletion of the NMD factor UPF1 increase R-loop levels and induce DNA damage in an R-loop-dependent manner, suggesting that inhibition of NMD preferentially kills MDS cells by exacerbating R-loop-associated genomic instability (Fig. 7b).

[H2] R-loops-associated DNA repair defects

As described above, EWS-FLI expression increases Pol II elongation and R-loop formation, and activates the ATR pathway⁹. Furthermore, EWS-FLI reduces HR efficiency⁹. In Ewing's sarcoma cells, BRCA1 associates with elongating Pol II and thus fails to localize at DSBs, conferring HR deficiency⁹. Because ATR is crucial for both HR and the R-loop response and as poly(ADP-ribose) polymerase (PARP) inhibitors selectively kill HR-deficient cells, the HR deficiency of Ewing's sarcoma cells renders them sensitive to inhibitors of ATR and PARPs (Fig. 7c).

[H2] Stabilization of G-quadruplexes

When R-loops are formed in sequences with high GC skew, the displaced G-rich ssDNA can give rise to G4s. Stabilization of G4s by G4 ligands renders R-loops more persistent, thereby enhancing their impacts on genomic stability. Furthermore, G4s themselves can interfere with DNA replication and cause DNA damage. In BRCA2-deficient cells, G4 ligands induce DNA damage in an R-loop-dependent manner¹⁸² (Fig. 7d). G4-stabilizing ligands also interfere with the

resolution of R-loops at sites of ROS-induced damage in transcribed regions, thereby compromising DSB repair¹⁰⁸. Telomeric R-loops are associated with G4s, and in ALT⁺ cancer cells, G4-stabilizing ligands enhance ALT activity and its associated telomere instability¹⁸³. These findings suggest that G4-stabilizing drugs can be used to exploit R-loop associated genomic instability in cancer cells.

[H2] Other R-loop-augmenting drugs

Though not originally designed to induce R-loops, several targeted cancer treatments such as TOP1 inhibitors (camptothecin), HDAC inhibitors (romidepsin) and BET protein inhibitors (JQ1) increase R-loop formation, presumably by inhibiting the R-loop suppressing activities of their targets^{123-125,173,184}. There are also reports that tumor-treating electric fields induce R-loop accumulation, possibly as a by-product of inhibiting DNA repair or inducing replication stress¹⁸⁵, and the natural compound anti-cancer drug trabectedin and its derivative lurbinectedin promote R-loop-partially dependent DNA damage¹⁸⁶. Such findings may have implications for combination therapy with other R-loop-targeting treatments.

[H1] Conclusions and future perspective

Our understanding of the sources, resolution and impact of R-loops and other RNA–DNA hybrids has extended considerably in the past few years owing to the development of new analytic tools and technologies, which allowed us to map R-loops genome-wide, to identify R-loop associated proteins, and to follow the effects of R-loops in various biological contexts. It has become clear that R-loops are not a homogenous structure, but a population of distinct and dynamic RNA–DNA–protein assemblies. The effects of R-loops on the genome are also diverse, depending on where, when and how R-loops are generated, how abundant and stable the R-loops are, and which proteins the R-loops interact with. Although numerous regulators of R-loops have been identified, we still know little about how these regulators function in concert in cells. Why do cells need so many R-loop regulators? Do these regulators function in a context-specific manner? If so, what controls the context specificities of R-loop regulators?

Going forward, it will be crucial to investigate how different types of R-loops and RNA–DNA hybrids are generated, processed and resolved, and how they function in specific chromosomal,

cellular and tissue contexts. This effort should include more detailed investigations of R-loopmediated genomic instability in cycling versus non-cycling cells and of the relative contribution of DNA replication to R-loop-mediated genomic instability. Finally, our understanding of the roles of R-loops in pathological and therapeutic contexts is still far from complete. It will be important to establish clinically useful assays to measure the accumulation of R-loops in patients, to understand how R-loops and associated genomic instability contribute to pathogenesis and to develop new strategies to alleviate or exploit R-loop-associated defects in therapy.

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Competing interests

The authors declare no competing interests.

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	R-loop suppressors	Description	References
	an an 1		70
	SRSFI	Splicing factor; binds the Pol II CID	19
	CEDO NONO	and hascent RIVA transcripts	187
	SFPQ, NONO	Involved in RNA splicing; suppress	
		Sulicing fortage mutated in	104 157
	SF3B1, UZAF1,	Splicing lactors; mutated in	20
	SKSF2	and cancer	
RNA	SDT6	Pol II associated alongation factor	85
nrocessing	5110	rearries the Integrator complex to	
factors		suppress y generated from long non-	
		coding RNA	
	TEIIS	Recognizes backtracked Pol II and	78
	11 115	stimulates transcript cleavage	
	THO TREX and	Promote transcription termination	188
	TREX-2 complexes	and assembly and nuclear export of	189
		messenger ribonucleoproteins	
	RNA exosome	3'-5'exoribonuclease: degrades non-	86
		coding RNAs and prevents R-loop	
		accumulation	
	TOP1	Removes supercoiling during	21
		replication and transcription;	62
Topoisomerase		suppresses R-loop-associated	
		replication stress at transcription	
		termination sites.	
	RNaseH1	Cleaves RNA in RNA–DNA hybrids	190
RNase H	RNaseH2	Cleaves RNA in RNA-DNA hybrids;	
		removes ribonucleotides mis-	59
		incorporated in DNA	14
	SETX	Putative RNA–DNA helicase;	88
		suppresses R-loops at transcription	45
		termination sites and DSBs.	71
	AQK	Putative RNA–DNA helicase;	
		Leavinda DNA DNA hybrida in	192
		vitro: suppress P loops in human	
		cells	
	FANCM	Unwinds telomeric RNA_DNA	106
		hybrids: suppresses telomeric R-	193
		loops in human cells.	

 Table 1: Proteins involved in suppression and resolution of R-loops and RNA–DNA hybrids

	DDX1	Associates with ATAD5 and	111
		suppresses R-loops during	
RNA-DNA		replication.	
helicases	DDX5	Associates with BRCA2 and	194
	22110	suppresses R-loops at DSBs.	111
		Associates with ATAD5 and	154
		suppresses R-loops during	
		replication	
	DDX19	Enters the nucleus following DNA	112
	DDAI)	damage in an ATR-dependent	
		manner: suppresses R-loops	
	DDV21	Inswinds PNA DNA hybrids in	91
	DDA21	vitro: functions with SIRT7 to	111
		suppress P loops at specific gapes	
		A appletes K-100ps at specific genes.	
		Associates with ATADS, suppresses	
	DDV20D	R-loops during replication.	90
	DDX39B	Unwinds RNA–DNA hybrids in	
		<i>vitro</i> ; suppresses co-transcriptional	
	DUNIO	R-loops genome-wide.	02
	DHX9	Binds RNA–DNA hybrids and	95 80.02
		unwinds R-loops.	111
		Associates with ATAD5 and	111
		suppresses R-loops during	
		replication.	
		Also shown to promote R-loop	
	D.D.G.L.I	formation.	00
	BRCA1	Suppresses R-loops at promoter-	88
		proximal Pol II pause sites and	97
		transcription termination sites.	
		Recruits SETX to transcription	
		termination sites.	
	BRCA2	Suppresses R-loops at promoter-	84
		proximal Pol II pause sites. Interacts	98
DNA repair		with DDX5 and TREX-2.	
proteins	Fanconi anemia	FANCD2–FANCI bind RNA–DNA	195
(non-helicases)	factors	hybrids and RNA processing factors	159
	(FANCD2, FANCI,	suppress R-loops to coordinate	103
	FANCA)	replication and transcription	
	RPA	Recruits and stimulates RNaseH1.	104
	XPG, XPF	Structure-specific nucleases involved	71
		in transcription-coupled nucleotide	
		excision repair (TC-NER), may	
		cleave and remove R-loops during	
		transcription.	

	MUS81	A structure-specific nuclease that	61
		cleaves Holliday junctions and stalled	115
		or reversed replication forks, may	
		process R-loops upon collisions of R-	
		loops and replication forks.	
	CtIP	May process R-loops at active genes.	105
	MRE11	Suppresses R-loops upon collisions	113
		of R-loops and replication forks,	
		promotes functions of FA proteins.	
	APTX	Suppresses R-loops through its	160
		function in single-strand break repair.	
	TDP1	Suppresses R-loops through its	160
		function in single-strand break repair.	
	SAMHD1	Deoxynucleoside	177
		triphosphohydrolase (dNTPase) and	
		3'-5' -exoribonuclease. Does not	
		resolve R-loops directly but can	
		recruit other factors such as MRE11.	
	ATR, Chk1	May suppress R-loops by preventing	61
DNA-damage		collisions between R-loops and	114
checkpoint		replication forks.	
proteins	ATM, Chk2	May suppress R-loops by promoting	114
		the repair of DSBs at replication	
		forks collapsed at R-loops.	
	SWI/SNF	ATP-dependent chromatin-	100
		remodeling complexes, may suppress	
		R-loops during transcription–	
		replication conflicts.	
	INO80	Part of the ATP-dependent	118
		chromatin-remodeling complex, may	
Chromatin		reorganize chromatin to resolve R-	
modulators		loops.	
	FACT	Histone chaperone, may reorganize	119
		chromatin at sites of R-loop-	
		replication collisions.	101
	PCR1	Promotes repressive chromatin,	
		suppresses R-loops by decreasing	
		transcription.	02
	SIN3A	A histone deacetylase that interacts	85
		with the THO complex.	100
	KAT8	A histone acetyltransferase that	122
		functions with BRD2 and BRD4 to	
		suppress R-loops.	

BRD2	A reader of histone Lys acetylation;	123
	recruits TOP1.	
BRD4	A reader of histone Lys acetylation;	196
	suppresses R-loops at specific genes	125
	by preventing Pol II pausing.	

APTX, aprataxin; AOR, RNA helicase aquarius; ATAD5, ATPase family AAA domaincontaining protein 5; ATM, ataxia-telangiectasia mutated; ATR, ATM and Rad3-related; BLM, Bloom syndrome protein; BRCA1, breast cancer type 1 susceptibility protein; BRD2, bromodomain-containing protein 2; Chk1, checkpoint kinase 1; CtIP, DNA endonuclease RBBP8; DDX1, ATP-dependent RNA helicase DDX1; DHX9, ATP-dependent RNA helicase A; DSBs, DNA double-stranded breaks; FA, Fanconi anemia; FANCD2, Fanconi anemia group D2 protein; KAT8, Histone acetyltransferase KAT8; Lys, lysine; MRE11, double-strand break repair protein MRE11; MUS81, crossover junction endonuclease MUS81; NONO, Non-POU domaincontaining octamer-binding protein; Pol II CTD, RNA polymerase II C-terminal domain; PCR1, Polycomb repressor complex 1; RNaseH1, ribonuclease H1; RPA, replication protein A; SAMHD1, deoxynucleoside triphosphate triphosphohydrolase SAMHD1; SETX, senataxin; SIRT7, NAD-dependent protein deacetylase sirtuin-7; SFPQ, splicing factor, proline- and glutamine-rich; SF3B1, splicing factor 3B subunit 1; SPT6, transcription elongation factor SPT6; SRSF1, serine/arginine-rich splicing factor 1; TFIIS, transcription elongation factor A protein 1; TDP1, tyrosyl-DNA phosphodiesterase 1; TOP1, topoisomerase 1; TREX-2; three prime repair exonuclease 2; U2AF1, splicing factor U2AF 35 kDa subunit; XPG, Xeroderma Pigmentosum complementation group G.

Figure legends

Fig. 1. Sources of R-loops and other RNA–DNA hybrids (I).

- **a.** *Co-transcriptional R-loops.* Left: Formation of R-loops at RNA polymerase II (Pol II) pause sites near transcription start sites (TSSs), at sequences with high GC skew in gene bodies, and at transcription termination sites (TTSs). Right: The pausing of Pol II at TTSs induces antisense transcription, which recruits Argonaute-2 (AGO2) and the histone Lys methyltransferase G9a (also known as EHMT2)¹⁵. G9a di-methylates histone H3 Lys9 (H3K9me2) and promotes the recruitment of heterochromatin protein 1 (HP1) and the formation of repressive chromatin.
- **b.** *Telomeric and centromeric R-loops.* Top: The long non-coding RNA telomeric repeatcontaining RNA (TERRA) forms R-loops at telomeres. TERRA interacts with telomere repeat binding factor 2 (TRF2) and ATRX, thereby contributing to telomere maintenance. Telomeric R-loops also induce replication stress, which promotes alternative lengthening of telomeres (ALT). Bottom: RNA transcripts from centromeric repeats form R-loops in *cis.* Centromeric RNAs bind histone H3-like centromeric protein A (CENPA), thereby promoting the localization of CENPA and centromere protein C (CENPC) to centromeres. In mitosis, centromeric R-loops recruit and activate the kinase ATR, which activates CHK1 and aurora kinase B (AurB) at centromeres and enables accurate chromosome segregation.

Fig. 2. Sources of R-loops and other RNA–DNA hybrids (II).

- **a.** *R-loops generated during replication.* Top: The DNA polymerase α (Pol α)-primase complex synthesizes RNA primers in Okazaki fragments on the lagging strand, thereby forming RNA–DNA hybrids. The RNA primers are subsequently removed by flap endonuclease 1 (FEN1) and the DNA is ligated by DNA ligase 1 (LIG1). Ribonucleotides (NTP) misincorporation during DNA replication can also generate RNA–DNA hybrids. Ribonucleases H2 (RNaseH2)-mediated ribonucleotide excision repair (RER) removes misincorporated ribonucleotides. Bottom: When a leading-strand DNA polymerase is stalled, DNA-directed primase/polymerase protein (PrimPol) re-primes replication ahead of the stalled polymerase.
- b. DNA damage-induced hybrids. Left: RNA polymerase II (Pol II) is recruited to DNA double-stranded breaks (DSBs) by the MRE11–RAD50–NBS1 (MRN) complex, enabling Pol II to carry out *de novo* RNA synthesis. Pol II-displaced DNA flaps are bound by replication protein A (RPA). Right: DSBs in transcriptionally-active regions induce Pol II pausing, allowing RNA transcripts to form RNA–DNA hybrids with help from RAD51 associated protein 1 (RAD51AP1) and USP1 associated factor 1 (UAF1)¹³⁸. Pol II may pause before or after DSBs, and the RNA–DNA hybrids at DSBs may exist in R-loops or on ssDNA ssDNA overhangs.

Fig. 3. Effects of R-loops and RNA–DNA hybrids on genomic stability.

- **a.** *Co-directional and head-on collisions of R-loops and replication forks.* Top: Codirectional collisions of replication forks with R-loops allow the replicative CDC45– MCM–GINS (CMG) helicase to unwind RNA–DNA hybrids, leading to dissociation of RNA and polymerase II (Pol II) and reduction of R-loop levels. Bottom: Head-on collisions of replication forks and R-loops force fork reversal. The processing of reversed forks by the structure-specific endonuclease MUS81 promotes activation of the kinase ATR, which in turn limits the activity of MUS81 and suppresses DSB formation.
- **b.** *DNA supercoiling-induced R-loops and collisions.* DNA supercoiling ahead of Pol II promote head-on collisions of R-loops and replication forks at transcription termination sites (TTSs). Topoisomerase 1 (TOP1) resolves the supercoils and alleviates replication fork collapse and consequently the formation of DNA double-stranded breaks (DSBs) at TTSs. In the absence of TOP1, replication protein A (RPA), which binds single-strand DNA (ssDNA) at TTSs becomes increasingly phosphorylated by ATR, which may facilitate the repair of DSBs.
- **c.** *R-loop-associated chromatin compaction.* R-loops are associated with gene-repressive histone modifications such as histone H3 Lys9 di-methylation (H3K9me2) and histone H3 Ser10 phosphorylation (H3S10p) and with compacted chromatin. H3K9me2 recruits heterochromatin protein 1 (HP1) to form heterochromatin. The compact chromatin induced by R-loops interferes with DNA replication and increases genomic instability.
- **d.** *Single-strand DNA damage in R-loops.* The displaced ssDNA in R-loops is a substrate of activation induced cytidine deaminase (AID) and APOBEC deaminases. The uracil resulting from cytosine deamination is recognized by uracil-DNA glycosylase (UNG) who generates abasic sites, which are subsequently converted to SSBs through base excision repair. Both abasic sites and SSBs interfere with replication forks, leading to stalling of DNA polymerases and fork collapse.

Fig. 4. Mechanisms of R-loop suppression and resolution.

- **a.** *R-loop suppression during transcription.* Ribonuclease H1 (RNaseH1 or RNH1) and RNaseH2 (RNH2) cleave the RNA in RNA–DNA hybrids, and RNA–DNA helicases such as senataxin (SETX) unwind R-loops. R-loop suppressors include chromatin modifiers that repress transcription, such as the histone H2A ubiquitylating (Ub) Polycomb repressive complex 1 (PRC1) and the histone deacetylase complex SIN3A. Lys acetyltransferase 8 (KAT8) can also suppress R-loops, by acetylating (Ac) histone H4 and thus recruiting bromodomain-containing protein 2 (BRD2) and BRD4; in turn, BRD2 recruits topoisomerase 1 (TOP1), which resolves DNA supercoiling and helps suppress R-loops. Prevention of Pol II pausing and backtracking by transcription regulators such as BRD4 counteracts R-loop formation. RNA splicing factors, such as serine and arginine rich splicing factor 1 (SRSF1), and factors involved in the assembly of messenger ribonucleoproteins (mRNPs) for mRNA export, such as the THO complex (THO), also prevent RNA–DNA hybrid formation.
- **b.** *R-loop bypass or resolution during DNA replication.* Bypass R-loops by replication forks is aided by chromatin remodelers such as the complex facilitates chromatin transactions (FACT), the INO80 complex and the SWI/SNF complex, which act in concert with Fanconi anemia group D2 protein (FANCD2) and the helicase senataxin (SETX). The activation of the kinase ATR by reversed replication forks following collision with R-loops prevents excessive fork cleavage by the structure-specific endonuclease MUS81 and promotes nuclear import of the helicase DEAD-box19 (DDX19) to remove R-loops. Replisomes might also resolve R-loops through unwinding by the replicative helicase minichromosome maintenance (MCM), or bypass R-loops through repriming by Polα-primase (not shown) and DNA-directed primase/polymerase protein (PrimPol), for example at triplex structures formed by GAA repeats. RNaseH2 (RNH2) interacts with PCNA and can remove R-loops at replication forks.
- **c.** *Repair and restart of DNA replication forks.* Replication fork reversal at R-loops can lead to fork processing by MUS81, followed by fork repair and restart involving the recombination protein RAD52 and DNA ligase IV (LIG4), and resumption of transcription activity. Fork repair might be aided by chromatin modifiers such as PRC1 and KAT8. H2A, histone H2A; H4, histone H4

Fig. 5. Roles of R-loops and RNA–DNA hybrids in DNA repair.

a. *Recruitment of DNA repair proteins.* Left: At sites of double-stranded DNA breaks (DSBs), damage-induced long non-coding RNAs (dilncRNAs) are transcribed and converted into double-strand RNAs, which are processed into small RNAs (DDRNAs or diRNAs) by the ribonuclease III Dicer pathway. diRNAs interact with Argonaute-2 (AGO2) and the homologous recombination (HR) factor RAD51 to promote the localization of RAD51 at DSBs. The RNA–DNA hybrids formed by dilncRNAs are also recognized by breast cancer type 1 susceptibility protein (BRCA1), thereby promoting the recruitment of BRCA2 and RAD51. Right: Reactive oxygen species (ROS) generate single-stranded DNA breaks (SSBs) and DSBs in transcribed regions, thereby inducing R-loop formation. The R-loops are recognized by Cockayne syndrome B (CSB) and RAD52, enabling RAD51 recruitment and DSB repair. The enzyme tRNA aspartic acid methyltransferase 1 (TRDMT1) modifies

the RNA in R-loops with 5-methylcytosine (m⁵C), thereby promoting RAD52 and RAD51 recruitment.

- **b.** *Single-stranded DNA and DNA end resection.* Left: In cells in G2 phase, DSB-induced R-loops are recognized by RAD52. RAD52 recruits xeroderma pigmentosum group G (XPG) to cleave R-loops, generating ssDNA that is bound by replication protein A (RPA), thereby enhancing HR activity. Right: RNA polymerase III (Pol III) is recruited to DSBs by the MRN complex and CTBP-interacting protein (CtIP). Pol III-mediated RNA synthesis promotes DNA end resection by unwinding double-stranded DNA (ssDNA). The ssDNA flap displaced by Pol III is bound by RPA and removed by DNA replication helicase/nuclease 2 (DNA2) and exonuclease 1 (EXO1).
- c. *RNA-templated repair*. When DSBs form in transcribed regions, the preexisting RNA transcripts can hybridize with ssDNA overhangs by RAD52 thereby enabling DNA polymerase θ (POL θ) to extend the ssDNA using RNA as template. The extended ssDNA from one DSB end can captures the ssDNA from the other DSB end, allowing DSB repair by gap filling.
- **d.** *DR-loops.* Preexisting RNA transcripts at DSBs can invade homologous DNA with the help of RAD51 associated protein 1 (RAD51AP1) and USP1 associated factor 1 (UAF1). The formation of R-loops in homologous DNA promotes ssDNA invasion and extension, giving rise to DR-loops. The RNA–DNA hybrids in DR-loops must be removed before the completion of HR.
- e. *Repair of single-stranded DNA breaks.* When SSBs form at the transcription termination sites, a complex of BRCA1 and RNAi factors processes local RNA transcripts into single-stranded DNA-damage-associated small RNAs (sdRNAs), which interact with RAD52 and partner and localizer of BRCA2 (PALB2) to localize them at SSB sites to repair the lesions.

Fig. 6. R-loops in pathology.

- **a.** *Gain of R-loops through increased RNA synthesis.* Left: Oncogenes such as EWS-FLI1 (fusion of EWS RNA binding protein 1 (EWSR1) and Friend leukemia integration 1 transcription factor (FLI1)), HRAS^{V12} and the estrogen receptor (ER) can promote R-loop formation by increasing RNA synthesis. Right:
- **b.** *Gain of R-loops through loss of RNA processing or DNA repair.* Suspected oncogenic mutations in splicing factors such as U2 small nuclear RNA auxiliary factor 1 (U2AF1) and serine/arginine-rich splicing factor 2 (SRSF2) may promote R-loop formation by increasing RNA polymerase II (Pol II) pausing owing to defects in co-transcriptional splicing. Loss of BRCA1 or BRCA2 also leads to increased Pol II pausing, and loss of BRCA2 additionally results in defective mRNP export by the TREX-2 complex, leading to R-loop accumulation. Loss of BRCA1 or Fanconi anemia group M (FANCM) may increase R-loop levels through loss of DNA repair.
- **c.** *Telomeric R-loops in cancer cells.* Loss of ATRX is associated with increased telomeric R-loops in cancer cells with alternative lengthening of telomeres (ALT) activity. FANCM and BRCA1 suppress the levels of telomeric R-loops, thereby keeping telomere replication stress and ALT activity at tolerable levels.
- **d.** *Excessive R-loop formation in neurodegenerative disorders.* In cerebellar ataxias caused by loss of DNA repair proteins such as ataxia telangiectasia mutated (ATM), tyrosyl-DNA phosphodiesterase 1 (TDP1) or senataxin (SETX), R-loop accumulation could exacerbate

cytotoxic DNA damage and inhibit Pol II, leading to cell death. In amyotrophic lateral sclerosis, frontotemporal dementia and Friedreich ataxia, R-loops accumulating owing to senataxin mutations or guanine (G)-rich repeat expansions could inhibit Pol II, leading to reduced expression of essential genes and cell death.

Fig. 7. Targeting R-loop-associated vulnerabilities in cancer therapy.

- **a.** *Inhibition of ATR.* In myelodysplastic syndrome and acute myeloid leukemia cells, mutant splicing factors such as U2AF1^{S34F} increase R-loop accumulation and activate ATR. ATR provides a protective feedback to suppress R-loop-induced double-strand DNA breaks (DSBs). ATR inhibitior (ATRi) disrupts the ATR-mediated feedback loop, thereby increasing DSBs and promoting cell death.
- **b.** *Inhibition of nonsense-mediated mRNA decay.* Splicing factor 3B subunit 1 (SF3B1) and nonsense mediated decay (NMD) associated PI3K related kinase (SMG1) prevent aberrant R-loop accumulation during transcription. In MDS and AML cells carrying spliceosome mutations, SF3B1 inhibitor (SF3B1i) and SMG1 inhibitor (SMG1i) further increase R-loop levels, exacerbating R-loop-associated DNA damage and driving cell death.
- **c.** *Inhibition of DNA repair.* The Ewing's sarcoma oncogene EWS-FLI increases R-loop levels and the interaction between elongating Pol II and BRCA1. The binding of BRCA1 to Pol II compromises the localization of BRCA1 to DSBs, thereby reducing homologous recombination (HR) activity and rendering Ewing's sarcoma cells sensitive to poly(ADP-ribose) polymerase inhibitors (PARPi).
- **d.** *Stabilization of G-quadruplexes.* Cancer cells with defects in R-loop suppressors such as BRCA2, are prone to form G-quadruplexes (G4s) in sequences with high GC skew owing to the displacement of G-rich single-strand DNA in R-loops. G4-stabilizing drugs can increase the levels of G4s and R-loops in cancer cells. Collisions of replication forks with G4-containing R-loops (left) or with G4s alone (right) can lead to DNA damage and cell death.

Glossary

BACKTRACKING

Backward movement of transcribing RNA polymerase, which enables proofreading and regulation of transcript elongation.

BREAK-INDUCED REPLICATION

A process in which one-ended DNA breaks generated at replication forks are extended using homologous DNA as template.

CEREBELLAR ATAXIA

Progressive neurological disorder caused by damage to the cerebellum, characterised by inability to control balance, gait and muscle coordination.

CGAS-STING PATHWAY

Cellular signaling pathway that senses DNA in the cytoplasm as a sign of viral or bacterial infection and activates innate immunity responses.

CLASS SWITCH RECOMBINATION

The process of switching the antibody type produced by mature B cells, during which the immunoglubulin gene is subject to transcription-dependent DNA damage followed by repair-mediated rearrangements.

DONOR DNA

The DNA sequence used as template for DNA repair during homologous recombination.

CONTROL REGION OF MTDNA

A non-coding region of the mitochondrial genome that controls RNA and DNA synthesis.

C-RICH STRAND OF TELOMERE DNA

Telomere DNA consists of TTAGGG repeats on the G-rich strand and CCCTAA repeats on the C-rich strand.

G-QUADRUPLEX

A DNA secondary secture formed by guanine (G)-rich sequences through G–G base pairing.

INTEGRATOR COMPLEX

A multisubunit protein complex with RNA endonuclease activity that controls the expression and processing of Pol II transcripts.

RNA EXOSOME

A multisubunit protein complex with 3'-5' exoribonuclease activity that degrades non-coding Pol II transcripts.

Nonsense-mediated mRNA decay

A cellular surveillance mechanism that detects and degrades mRNAs harboring premature termination codons.

REPLICATION FORK REVERSAL

Backward movement of the replication fork during which the nascent newly-synthesized strands dissociate from the template strands and anneal together to form a four-way junction.

REPLICATION STRESS

A plethora of DNA replication impedements that compromise the efficiency or fidelity of DNA synthesis and increase genomic instability.

DNA end resection

A process in which exonucleases cleave one of the DNA strands at DNA double-stranded breaks to generate overhangs.

RNA EDITING

Post-transcriptional enzymatic process that changes RNA nucleotides, for example by deaminating adenosine to inosine.

TRANSCRIPTION-REPLICATION CONFLICTS

The conllisions between transcription and DNA replication complexes and consequences in both transcription and DNA replication.

TRANSLESION SYNTHESIS

A process in which a group of specialized DNA polymerases at or behind replication forks bypass DNA lesions to ensue replication.

TUMOR-TREATING ELECTRIC FIELDS

A non-invasive treatment where alternating electric fields are applied to tumour sites to inhibit cancer cell proliferation

Co-transcriptional R-loops





DNA damage-induced R-loops and RNA-DNA hybrids

b



Collisions between R-loops and replication forks





а



d



Fork collapse





Transcription restart

а

Recruitment of DDR proteins





b



DSB Pol III CtIP Pol III 3' End processing 5' RPA 5' RPA 5' RPA 3' Flap removal \downarrow DNA2 EX01



Removal of RNA-DNA hybrids

е



С

Loss of DNA repair а b 000 FANCM **Replication stress** Increased RNA synthesis (BRCA1) RNA Pol II pausing BRCA1 **Replication stress** Pol II Pol II BRCA2 HRASMUT EWS-FLI1 SRSF2^{MUT} TREX-2 U2AF1^{MUT} Loss of mRNP Loss of DNA repair ALT telomeres d С Tolerable replication TDP1 stress and ALT activity Loss of RNA-DNA helicase ATM SETX DNA damage ∞ Pol II Pol II FANCM G-rich repeat BRCA1 Suppress telomeric R-loops

Transcription inhibition

Fig. 6

Inhibition of the ATR-Chk1 pathway



b

Inhibition of RNA splicing or NMD



С

Inhibition of DNA repair



Stabilization of G4s



Fork collisions with R-loops

Fork collisions with G4s (after R-loops removal)