Replication protein A binds RNA and promotes R-loop formation

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Replication protein A (RPA), a major eukaryotic ssDNA-binding protein, is essential for all metabolic processes that involve ssDNA, including DNA replication, repair, and damage signaling. To perform its functions, RPA binds ssDNA tightly. In contrast, it was presumed that RPA binds RNA weakly. However, recent data suggest that RPA may play a role in RNA metabolism. RPA stimulates RNA-templated DNA repair in vitro and associates in vivo with R-loops, the three-stranded structures consisting of an RNA-DNA hybrid and the displaced ssDNA strand. R-loops are common in the genomes of pro- and eukaryotes, including humans, and may play an important role in transcription-coupled homologous recombination and DNA replication restart. However, the mechanism of R-loop formation remains unknown. Here, we investigated the RNA-binding properties of human RPA and its possible role in R-loop formation. Using gel-retardation and RNA/DNA competition assays, we found that RPA binds RNA with an unexpectedly high affinity (KD ≈ 100 pm). Furthermore, RPA, by forming a complex with RNA, can promote R-loop formation with homologous dsDNA. In reconstitution experiments, we showed that human DNA polymerases can utilize RPA-generated R-loops for initiation of DNA synthesis, mimicking the process of replication restart in vivo. These results demonstrate that RPA binds RNA with high affinity, supporting the role of this protein in RNA metabolism and suggesting a mechanism of genome maintenance that depends on RPA-mediated DNA replication restart.

Replication protein A (RPA) is a major ssDNA-binding protein in eukaryotes (1). It is a highly conserved trimeric protein composed of three subunits, RPA70, RPA32, and RPA14, which all are essential for cell viability (2). RPA plays a critical role in most, if not all, metabolic processes that involve ssDNA, including DNA replication, repair, transcription, and DNA damage signaling (2–5). RPA binding protects ssDNA from degradation and unfolds DNA secondary structures. RPA interacts with various proteins helping to coordinate different cellular processes.

Recently, it was found that RPA is closely associated with R-loops in vivo (6–8). R-loops are currently known to exist in the genomes of bacteria, yeast, and higher eukaryotes (9–11). In humans, R-loops occur over tens of thousands of genomic loci covering up to 5% of the genome (12, 13).

It was suggested that R-loops may play an important role during DNA repair by initiating transcription-coupled homologous recombination in actively transcribed genome regions (6, 14, 15). It was also proposed that R-loops may promote restart of replication forks stalled at damaged DNA (16, 17). The role of R-loops in priming replication was actually the first biological function proposed for this structure in bacteria (18). More recently, it was found that in eukaryotes, persistent RNA-DNA hybrids initiate DNA synthesis in ribosomal DNA in a replication origin-independent manner (19). Being an important regulator of cellular processes such as transcription, gene expression, DNA replication, and DNA repair, R-loops also represent a source of genome instability, if not timely processed or repaired (20, 21). The mechanism of R-loop formation in vivo remains to be understood.

RPA has a strong binding affinity to ssDNA (2, 22); therefore, it was thought that RPA association with R-loops is due to its binding to the displaced ssDNA strand generated during R-loop formation. Surprisingly, until recently, RPA binding to RNA had not been explored. It was presumed that RPA binds to RNA weakly, because in early studies the affinity of RPA for RNA had not been explored. It was previously anticipated. We found that RPA binds RNA with high affinity (KD ≈ 100 pm). Furthermore, we demonstrate that RPA has a unique ability to form an active complex with RNA, which promotes formation of bona fide R-loops through invasion of RNA into homologous covalently closed duplex DNA. Using RPA-generated R-loops, we reconstituted DNA synthesis in vitro using human DNA polymerases, supporting the role of R-loops in the mechanism of DNA replication restart.

Results

RPA binds to RNA with high affinity

First, using electrophoretic mobility shift assay (EMSA), we examined the RPA affinity for RNA. Previously, it was reported that the RPA-binding affinity for RNA is approximately the same as for dsDNA and ~1000-fold weaker than for ssDNA.
Surprisingly, we found that RPA binding to a 48-nt RNA (no. 501; Table S1) ($K_D = 101.4 \pm 17.0 \text{ pM}$) is 300–400-fold stronger than for homologous 48-bp dsDNA (nos. 211/212) ($K_D = 35.5 \pm 7.0 \text{ nM}$) and only 30–60-fold weaker than for a 48-mer ssDNA of the identical sequence (no. 211) ($K_D = 3.1 \pm 0.6 \text{ pM}$; Fig. 1 and Fig. S1). The presence of 100 mM NaCl, the condition that was used in previous studies, had no significant effect on the RPA affinity ($K_D = 72.0 \pm 10.2 \text{ pm}$) for RNA (no. 501) (Fig. S2). We also tested the RPA-binding affinity for RNA-DNA hybrid (nos. 501/212), which appeared to be 2 times weaker ($K_D = 85.9 \pm 4.5 \text{ nM}$) than for dsDNA of identical sequence (nos. 211/212) (Fig. 1E and Fig. S1C).

Then we examined RPA binding to RNA using competitors. When RNA was used as a competitor against ssDNA, we found that the RPA affinity for RNA (no. 501) was ~60-fold weaker than for ssDNA of identical sequence (no. 211) (Fig. S3, A and B). When nonhomologous supercoiled pHSG299 plasmid dsDNA was used as a competitor, the affinities of RPA for RNA and ssDNA were ~500- and ~33,000-fold, respectively, stronger than for plasmid dsDNA (Fig. S3, C and D). Thus, these results were consistent with the RPA $K_D$ values for RNA and DNA indicated above.

Then we tested the RPA binding to four other 48-nt RNAs of different sequences (Fig. S4). For three of them (nos. 3R, 7R, and 8R), the RPA-binding affinity was strong ($K_D$ in the range of 62.9–248.1 pm), and for one of them (no. 540), it was significantly weaker ($K_D > 4 \text{ nm}$). Inspection of the RNA structures showed that no. 540 has a much stronger propensity to form secondary structures than other tested RNAs (Table S2). We also tested the RPA binding to homopolymers: poly(rA) (no. 10R, 48 nt) and poly(rU) (no. 11R, 48 nt) (Fig. S5, A and B). For poly(rU), the RPA-binding affinity ($K_D = 184.4 \pm 26.4 \text{ pM}$) was in the range with other tested RNA molecules of the same size, except for no. 540, whereas for poly(rA), it was significantly weaker ($K_D = 1.4 \pm 0.2 \text{ nM}$). Moreover, poly(rA) appeared to be an extremely weak competitor against ssDNA (no. 211, 48 nt), even weaker than could be expected based on the $K_D$ measurements. Consistent with the reports from Wold’s group (23), a 1000-fold excess of poly(rA) was not sufficient for a 2-fold decrease in RPA binding to ssDNA (Fig. S5C). An increase in incubation time up to 3 h had no apparent effect on the outcome of poly(rA) competition with ssDNA for RPA binding. We suggest that this weak competitiveness of poly(rA) may be related to the kinetics of RPA binding to this substrate. For
instance, RPA-poly(rA) complexes may have a significantly shorter lifetime than the RPA-ssDNA complexes.

Overall, these data show that RPA binds RNA with high affinity. They also indicate that RPA binding to RNA is lowered by RNA secondary structures and by poly(rA) sequences.

**RPA promotes R-loop formation in vitro**

The finding that RPA binds RNA strongly, taken together with the known association of RPA with R-loops in vivo, prompted us to test whether RPA has the R-loop formation activity (Fig. 2A). Indeed, we found that RPA can promote R-loop formation between a $^{32}$P-labeled 48-mer RNA (no. 501) and homologous supercoiled pUC19 plasmid DNA (Fig. 2, B and C). In these experiments, the plasmid DNA was prepared by a non-denaturing method to avoid formation of irreversibly denatured plasmid DNA, a source of a potential artifact due to RNA/DNA annealing. We then tested the authenticity of the RPA-promoted R-loops. In contrast to RNA-DNA hybrids

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**Figure 2. RPA promotes R-loop formation.**

A, reaction scheme. $^*$, $^{32}$P label. B, kinetics of the R-loop formation by RPA (200 nM) analyzed by electrophoresis in a 1% agarose gel. 180$^0$, reaction (180 min) in the absence of RPA. The stoichiometric ratio of RNA/dsDNA was 5:1 (in molecules). C, graphical representation of R- and D-loop formation by RPA. D, sensitivity of R-loops to EcoRI cleavage. $^{32}$P-labeled RNA (no. 501; 3 $\mu$M, nt) was incubated with supercoiled pUC19 dsDNA (67.2 $\mu$M, nt) for 3 h in the presence of RPA (200 nM, lanes 2 and 3) or in its absence (lanes 4 and 5). The R-loops were then incubated with EcoRI (lane 3). The products were analyzed by electrophoresis in 1% agarose gel. Controls include [$^{32}$P]RNA (no. 501; 3 $\mu$M, nt) (lane 1) and a mixture of $^{32}$P-labeled RNA (no. 501; 3 $\mu$M, nt) and pUC19 incubated with EcoRI storage buffer (lane 4) or with EcoRI (lane 5). The gel was autoradiographed (top) to visualize $^{32}$P-labeled RNA and R-loops and then stained with ethidium bromide (bottom) to monitor intactness of supercoiled pUC19 dsDNA and its cleavage by EcoRI. Note that R-loops co-migrate in the gel with supercoiled pUC19 DNA. E, sensitivity of R-loops to RNase H. The R-loops produced as in D (lane 2) were incubated with RNase H (5 units) (lane 2) or with the storage buffer (lane 2). The products were analyzed as in D. F, RNA length dependence of R-loop formation by RPA. R-loops were formed by RPA (200 nM) in pUC19 DNA (67.2 $\mu$M, nt) using $^{32}$P-labeled RNAs: 12-mer (no. 4R), 24-mer (no. 5R), 36-mer (no. 6R), and 48-mer (no. 501) (each 3 $\mu$M, nt). R-loops were analyzed by electrophoresis in a 1% agarose gel. G, data from F presented as a graph. Error bars, S.E.
produced by annealing or RNA-protein complexes that can resist deproteinization, R-loops, similar to D-loops, are sensitive to plasmid dsDNA cleavage (outside of the R-loop region) with a restriction endonuclease (24). The cleavage causes loss of plasmid dsDNA superhelicity and R-loop dissociation due to DNA branch migration. We found that dsDNA linearization with EcoRI indeed causes R-loop dissociation, confirming their bona fide nature (Fig. 2D). As expected, the R-loops were also sensitive to RNase H, which digests the RNA moiety in the RNA-DNA hybrid (Fig. 2E).

When RNA was replaced with a 48-mer ssDNA of the identical sequence (no. 211), the efficiency of the reaction (D-loop formation) was reduced significantly, indicating that the RPA activity was specific for R-loop formation (Fig. 2C and Fig. S6A). R-loop formation strictly requires homology between RNA and dsDNA; no R-loops were formed with a nonhomologous RNA (no. 534, 48 nt) (Fig. S6B). Not all tested RNAs were equally proficient in RPA-promoted R-loop formation (Fig. S7). This proficiency inversely correlates with the RNA propensity to form secondary structures (Table S2). It does not generally correlate with the RPA-binding affinity for the tested RNAs, as RPA has similar $K_D$ for nos. 501 and 3R, which differ dramatically in their ability to support R-loop formation (Table S2). However, by titrating the RPA-RNA complexes with NaCl, we found that the RPA complex with RNA no. 501 is more stable than with RNA no. 3R (Fig. S8). Thus, the stability of RPA-RNA complexes may contribute to R-loop formation efficiency. The yield of R-loop formation rises with the increase of RNA length from 24 to 48 nt. No R-loops formed with a 12-nt RNA (Fig. 2, F and G), consistent with poor RPA binding to short RNAs (Fig. S9).

The R-loop–forming activity shows evolutionary conservation among RPA orthologs. *Saccharomyces cerevisiae* RPA (ScRPA) promotes R-loop formation, albeit with an ~6-fold reduced efficiency. We also tested an RPA functional homolog from *Escherichia coli* (EcSSB) for R-loop formation activity (Fig. 3). No activity was observed in a broad range of EcSSB protein concentrations under standard R-loop formation conditions. We also tested two other conditions, in which EcSSB protein showed the strongest annealing activity, such as a buffer with pH 5.5 or the presence of 2 mM spermidine (pH 7.0) (25). However, at any of the tested conditions, EcSSB did not show R-loop formation activity. The optimal RPA concentration for R-loop formation was one RPA heterotrimer per 15 nt and 30 nt of RNA for human and yeast orthologs, respectively (Fig. S10). Human RPA produces R-loops over a broad range of Mg$^{2+}$ dependence with a maximal R-loop yield at 2 mM (Fig. S11). We also found that RAD52 or RAD51 recombinase, which efficiently promoted D-loop formation, did not promote R-loop formation (Fig. 4). RAD52 at different concentrations was mixed with supercoiled pUC19 dsDNA (9.3 nM (molecules) or 50 nM (nt)) first, followed by the addition of RNA (93 nM (molecules) or 4.5 nM (nt)) (Fig. S12). We found that under these conditions, R-loops can be formed, but their yield even under optimal RAD52 concentration (200 nm) was very low (0.6%). These results indicate that RAD52 promotes inverse RNA strand exchange in proximity to dsDNA.

Figure 3. Human and yeast RPA promote R-loop formation. A, human RPA (HsRPA) (200 nM) and yeast RPA (ScRPA) (100 nM), but not *E. coli* SSB (EcSSB) (270 nM), promote R-loop formation between a 48-mer RNA (no. 501; 3 μM, nt) and pUC19 dsDNA (67.2 μM, nt). The R-loops were analyzed by electrophoresis in a 1% agarose gel. B, data from A presented as a graph. C, kinetics of R-loop formation by ScRPA (100 nM) between 48-mer RNA (no. 501; 3 μM, nt) and pUC19 dsDNA (67.2 μM, nt) analyzed by electrophoresis in a 1% agarose gel. D, data from C presented as a graph. Error bars, S.E.
Figure 4. Human RAD52 and RAD51 promote formation of D-loops, but not R-loops. A, scheme of D/R-loop formation. *, $^{32}$P label. B, kinetics of RAD52-promoted D- and R-loop formation. RAD52 (450 nM) was preincubated with a 48-mer ssDNA (no. 211; 3 $\mu$M, nt) or RNA (no. 501; 3 $\mu$M, nt) of the same sequence. The reactions were initiated by the addition of supercoiled pUC19 dsDNA (50 $\mu$M, nt), and the products were analyzed by electrophoresis in a 1% agarose gel. 60sp, RAD52-independent (spontaneous) D/R-loop formation after 60 min of reaction. C, data from B shown as a graph. D, kinetics of RAD51-promoted D- and R-loop formation. RAD51 (1 $\mu$M) was preincubated with a 48-mer ssDNA (no. 211; 3 $\mu$M, nt) or RNA (no. 501; 3 $\mu$M, nt). The reactions were initiated by the addition of supercoiled pUC19 dsDNA (50 $\mu$M, nt). The products were analyzed by electrophoresis in a 1% agarose gel. 60sp, RAD51-independent (spontaneous) D/R-loop formation after 60 min of incubation. E, data from D shown as a graph.
break ends. Thus, R-loop formation appeared to be a unique activity of RPA among tested proteins.

Reconstitution of DNA replication restart using R-loops

Previously, it was suggested by Kogoma (16) that R-loops may be used to initiate the restart of DNA replication stalled at DNA damage sites. The ability of RPA to form R-loop may be especially relevant to this hypothesis because of a strong and well-documented RPA association with DNA replication. Thus, RPA was initially discovered in human cell extracts as a component essential for SV40 DNA replication in vitro (26–28). Here, we wanted to test whether human DNA polymerases pol δ, pol α, and pol ε and the translesion polymerase pol η can use R-loops for initiation of DNA replication. Pol η was shown to promote DNA synthesis from homologous recombination intermediates (D-loops) (29). In our experiments, DNA polymerases were directly added to the R-loops generated by RPA with 32P-labeled RNA (no. 501) and pUC19 DNA in the presence of four dNTPs (Fig. 5A). RNA extension by DNA polymerases was visualized by electrophoresis in denaturing gels. In control (lane 1), DNA polymerases were omitted. 32P-labeled markers are shown in lanes 6–8. C, effect of RNA, ssDNA, RPA, and RPA-RNA on DNA synthesis by pol ε. RNA extension by pol ε (50 nM) was carried out using deproteinized and purified R-loops (1 nM) (lane 2). The R-loops were premixed with ssDNA (no. 2; 3 μM, nt) (lane 3), RNA (no. 517; 3 μM, nt) (lane 4), RPA (5 nM) (lane 5), or a mixture of RPA (200 nM) and RNA (no. 517; 3 μM, nt) (lane 7) prior to pol ε addition. In control (lanes 1 and 6), pol ε was substituted with storage buffer.

Figure 5. In vitro reconstitution of DNA synthesis restart from R-loops. A, experimental scheme. *32P label at 5′-end of RNA (48 nt of no. 501). Blue arrow, extension of RNA by DNA polymerases. B, R-loops (3 nM) were generated in pUC19 using RPA. RNA extension in R-loops was carried out using DNA pol α (50 nM), η (38 nM), ε (50 nM), or pol δ (0.5 nM). The products of RNA extension were analyzed by electrophoresis in 8% polyacrylamide denaturing gels. In control (lane 1), DNA polymerases were omitted. 32P-labeled markers are shown in lanes 6–8. C, effect of RNA, ssDNA, RPA, and RPA-RNA on DNA synthesis by pol ε. RNA extension by pol ε (50 nM) was carried out using deproteinized and purified R-loops (1 nM) (lane 2). The R-loops were premixed with ssDNA (no. 2; 3 μM, nt) (lane 3), RNA (no. 517; 3 μM, nt) (lane 4), RPA (5 nM) (lane 5), or a mixture of RPA (200 nM) and RNA (no. 517; 3 μM, nt) (lane 7) prior to pol ε addition. In control (lanes 1 and 6), pol ε was substituted with storage buffer.

RPA promotes R-loop formation

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Figure 6. Proposed RPA-dependent DNA replication restart initiated at R-loop. RPA promotes formation of the R-loop that serves as a primer for a DNA polymerase during replication restart.

4 O. M. Mazina and A. V. Mazin, unpublished results.
To investigate whether the poor ability of pol ε to use native R-loops for DNA extension is intrinsic or is caused by an inhibitory effect of RPA present in the reaction, we deproteinized and purified R-loops. In this case, pol ε and other tested DNA polymerases efficiently extended RNA (Fig. 5C and Fig. S13). Moreover, the reactions were not affected significantly when RPA was added back to purified R-loops at a concentration that was sufficient to cover the displaced ssDNA strand in R-loops at a stoichiometry of 1 trimer per 15 nt. However, when free RNA (no. 517; 3 μM, nt) or RPA-RNA complexes were added to R-loops, we found a strong inhibition of pol ε (Fig. 5C, lanes 4 and 7). Thus, the presence of RPA-RNA complexes inhibited activity of pol ε in the reconstitution experiments with nondeproteinized R-loops. Pol ε was also sensitive to free ssDNA (no. 2; 3 μM, nt), albeit to a lesser degree (Fig. 5C, lane 3). Strong inhibition of pol ε with RNA (Fig. 5C, lane 4) was unexpected and requires further investigation. Among other tested polymerases, only pol η showed some mild sensitivity to free ssDNA and RNA (Fig. S13, compare the RNA primer uptake in lane 7 with those in lanes 8 and 9), and none of them showed significant sensitivity to RPA under tested conditions (Fig. S13, lanes 5, 10, and 18). Thus, RPA-generated R-loops can be used for initiation of DNA synthesis by human DNA polymerases: pol α, pol η, pol δ, and pol ε.

**Discussion**

In this study, we identified novel unanticipated activities of RPA: high-affinity binding to RNA and formation of R-loops between RNA and homologous supercoiled dsDNA. We show that human DNA polymerases α, δ, ε, and η can utilize RPA-generated R-loops for initiation of DNA synthesis in vitro, supporting a previously proposed role of R-loops in DNA replication restart (16,17).

Whereas binding of RPA to RNA was demonstrated in early studies, the RPA-binding affinity for RNA was underestimated. The affinity of RPA for RNA was measured indirectly using RNA as a nonlabeled poly(rA) and poly(rIC) competitor against labeled ssDNA in the filter-binding assay (23). As we show here, poly(rA) is an exceptionally poor competitor against ssDNA, not adequately representing RNA with mixed-base composition. Poly(rIC) readily forms double-stranded structures to which RPA is known to bind poorly. Recently, using fluorescence anisotropy, the $K_D$ values for RPA binding to ssDNA and RNA were determined as 4 and 15.2 nm, respectively (32). Based on these data, one could conclude that RPA binds to RNA with only 3–4-fold lower affinity than to ssDNA. However, the RNA and ssDNA concentrations (5 nM) in this study were too close to the reported $K_D$ values, especially for ssDNA, making the accuracy of these measurements problematic. In addition, the length and the structure of this RNA substrate were not reported in the paper. Moreover, in other experiments, the poly(dU) sequences were referred to as “RNA,” making the need for the substrate description even stronger. In contrast, our data show that RPA binds to RNA with high affinity ($K_D \sim 100$ pm for a 48-mer), about 500-fold stronger than to dsDNA but still 30–60-fold weaker than to ssDNA.

The high affinity of RPA to RNA in vitro may suggest that RPA binds RNA also in vivo. Recent proteomics studies support this proposal. Thus, RPA has been identified among RNA-interacting proteins in mammalian (33,34) and yeast cells (35). Bonasio’s group (33), by protein-RNA photocross-linking and quantitative MS, identified RPA among the proteins that interact with RNA regardless of its polyadenylation status in the nuclei of mouse embryonic stem cells. The RNA-cross-linked peptide $^{263}$VYFSK$^{268}$ was mapped in the DNA-binding domain A of RPA70. Mendell’s group (34) identified RPA among the proteins that interact with long noncoding RNA NORAD. In that study, biotinylated RNA fragments of NORAD were incubated with human HTT-116 whole-cell lysates, and the proteins that bind to these fragments were eluted and identified by MS. RPA70, RPA32, and RPA14 subunits were among the proteins that specifically bound NORAD RNA. Parker’s group (35), by UV cross-linking proteins to mRNAs, identified RPA among the proteins that directly interact with mRNA in vivo. mRNA-protein complexes were then purified under denaturing conditions using oligo(dT) columns, and the RNA–bound proteins were analyzed by LC–MS/MS (35). ScRFA1 subunit (ortholog of HsRPA70) was identified among the mRNA-bound proteins. The biological role of RPA-RNA interactions remains to be understood. RPA may protect RNA from RNases or recruit proteins that are involved in RNA metabolism. A putative role of RPA in mRNA nuclear export was reported (36). Additional studies are needed to further characterize RPA-RNA interactions in vivo.

Even though RNA is abundant in the cell, RPA binding to RNA may not necessarily interfere with its well-established functions in DNA metabolism that require RPA binding to ssDNA. It is likely that RPA will transfer from RNA to ssDNA generated during DNA replication stress or damage due to its 60-fold higher affinity for ssDNA. The dynamic nature of RPA binding was demonstrated for ssDNA; RPA can migrate along the ssDNA axis and transfer from one polynucleotide to another (22,37–39).

R-loop formation promoted by RPA is a unique type of strand exchange, as it is initiated by a complex that RPA forms with RNA. In contrast, all other known types of strand exchange, both forward and inverse, are initiated by a recombinase-DNA complex. For instance, *E. coli* RecA promotes formation of R-loops or RNA-DNA heteroduplexes in inverse RNA strand exchange by assembling an active complex on dsDNA, which then engages free RNA (17,40). Similarly, RAD52 promotes formation of RNA-DNA heteroduplex in inverse RNA strand exchange by forming an active complex with dsDNA (41). R-loop formation was reported for ICP8, the herpes simplex virus type-1 ssDNA-binding protein (42). However, this reaction occurred only with the alkali-denatured form of plasmid dsDNA through the annealing mechanism. Thus, RPA appeared to be the first known protein that promotes formation of *bona fide* R-loops by forming an active complex with RNA and by promoting invasion of RNA into covalently closed duplex DNA.

Although the mechanism of R-loop formation by RPA remains to be investigated, several assumptions can be made. During the initial step of R-loop formation, RPA acts in a
complex with RNA due to its ~500-fold higher affinity for RNA compared with the plasmid dsDNA. Moreover, the optimal amount of RPA required for R-loop formation corresponds to its stoichiometric coverage of RNA, but not dsDNA (Fig. S10). Next, RPA-RNA complex needs to engage dsDNA in the homology search process. The RPA trimer has at least four DNA-binding domains (3, 22), which could potentially provide binding space to both RNA and dsDNA, juxtaposing them for RNA-DNA pairing. Binding of dsDNA by the RPA-RNA complex should be by necessity weak to allow multiple association-dissociation steps during the homology probing. After homology is found and initial R-loops are formed, RPA may not remain bound to the newly formed RNA-DNA heteroduplex but be transferred to the displaced ssDNA strand, to which it has much higher affinity (Fig. 1). This RPA binding to the displaced ssDNA strand may help to stabilize and further expand the R-loop. A relatively weaker RPA binding to RNA compared with ssDNA may favor its R-loop formation activity as opposed to D-loop formation. Because of a strong binding to RPA, ssDNA may occupy all available binding space, preventing dsDNA binding that is needed for formation of D-loops.

It is highly plausible that R-loop formation by RPA is not the only mechanism that exists in the cell. Because we did not find such activity in EcSSB, we assume that E. coli may use other mechanisms that remain to be identified. RecA-mediated inverse RNA strand exchange for R-loop formation was previously proposed to be one of these mechanisms (17). We also cannot exclude the possibility that some auxiliary proteins are required for stimulation of EcSSB R-loop formation activity. Recent data indicate that R-loops are a common structure in genomes of humans and other species (12, 13). The biological role of R-loops is currently under intense investigation. It was found that R-loops are essential for repair of DNA double-strand breaks in actively transcribed genome regions through transcription-coupled homologous recombination (6, 14, 15) or non-homologous end joining (43). It was proposed by Kogoma (16) that R-loops may serve as a primer to restart DNA replication stalled at DNA lesions (Fig. 6). The R-loop formation activity of RPA may be especially relevant to replication restart because of a strong RPA association with DNA replication (26–28). The RPA32 subunit was directly UV cross-linked with the RNA strand of the nascent RNA-DNA primer during SV40 replication in nuclei of monkey CV-1 cells (44). It was demonstrated that RPA interacts with pol α, RFC, and pol δ (45, 46). RPA stabilizes a complex between short RNA primer and pol α and then coordinates loading of RFC, PCNA, and pol δ to initiate DNA synthesis. We found that all tested human DNA polymerases, pol α, pol δ, pol ε, and pol η can utilize RPA-generated R-loops for initiation of DNA synthesis in vitro. These in vitro reconstitution experiments further support Kogoma’s hypothesis and suggest the mechanisms of genome maintenance that depend on RPA and RNA.

**Experimental procedures**

**Proteins, DNA, and RNA**

Human DNA polymerases pol η, the catalytic core of pol α p180(335–1257), the FLAG-tagged four-subunit pol δ, RFC, and PCNA were purified as described (50–53). The catalytic core of human pol ε p261(1–1172)exo + was purified according to Ref. 54 with the following modifications: His tag was placed on the N terminus before a SUMO tag and removed by SUMO protease after the first purification step, which included the nickel ion affinity chromatography. The oligodeoxyribonucleotides (Table S1) were purchased from IDT Inc. and further purified by electrophoresis in polyacrylamide gels containing 50% urea (55). HPLC-purified oligoribonucleotides were purchased from IDT Inc. All experiments with RNA were carried out in the presence of 1 × Ambion RNasecure RNase inactivation reagent. Double-stranded oligonucleotides were prepared by annealing of equimolar (molecules) amounts of complementary oligonucleotides (55). When indicated, oligonucleotides were 5-end–labeled with [γ-32P]ATP using T4 polynucleotide kinase (New England Biolabs). Supercoiled pUC19 plasmid dsDNA was prepared by a method that did not involve DNA denaturation (56) with modifications. Briefly, E. coli host cells were treated with lysozyme and lysed with Triton X-100. The lysate was cleared by centrifuged at 4 °C for 30 min at 40,000 × g. The cleared lysate was mixed with ethidium bromide to 700 μg/ml and CsCl (0.59 g/1 ml of cleared lysate) and loaded at the top of CsCl (1.58 g/ml in water) solution that filled the bottom half of the centrifuge tube. The samples were centrifuged in an angle rotor for 24 h at 200,000 × g at the ambient temperature. Isolated supercoiled plasmid DNA was further purified by 3× butanol extractions followed by gel filtration on a Sephacryl S-500 column. Supercooled pHSG299 plasmid dsDNA purified by CsCl-ethidium bromide gradient centrifugation was purchased from Takara Bio Inc. pHSG299 is a derivative of pUC19 plasmid in which an ampicillin-resistant gene was replaced with a kanamycin-resistant gene. DNA and RNA concentrations are expressed in moles of molecules or, when indicated, in moles of nucleotides.

**RPA binding to RNA, ssDNA, dsDNA, and RNA-DNA hybrid using EMSA**

20-μl mixtures contained human RPA at the indicated concentrations, 25 mM Tris-acetate (pH 7.5), 10 mM KCl (added with the protein stock), 2 mM DTT, 1 mM magnesium acetate, 100 μg/ml BSA, and 32P-labeled 48-mer nucleic acid substrates: RNA (no. 501; 5 PM, molecules), ssDNA (no. 211; 0.5 PM, molecules), dsDNA (nos. 211/212; 3 NM, molecules), or RNA-DNA hybrid (nos. 501/212; 3 NM, molecules). The mixtures were incubated for 15 min at 37 °C and then placed on ice. Each sample was supplemented with 3 μl of 50% glycerol and loaded onto a 6% polyacrylamide gel (29:1) in 0.25× TBE buffer (22.5 mM Tris, 22.5 mM borate, and 0.25 mM EDTA, pH 8.3). Bromphenol blue was added only in the sample containing 32P-labeled probe without RPA. Electrophoresis was carried out at 13 V/cm for 1 h at room temperature. The gels were dried on Amersham Biosciences Hybond-N+ membrane (GE Healthcare) and analyzed using a Typhoon FLA 7000 biomolecular imager. The KD and Bmax values were obtained by fitting the data to a one-site binding hyperbola in GraphPad Prism 5.0.
RPA promotes R-loop formation

RPA-promoted R-loop formation was carried out for 3 h at 37 °C, as described above. Then 1 μl of 10× RNase H reaction buffer and 1 μl (5 units) of RNase H (New England Biolabs) were added to 8 μl of the reaction mixture, and incubation was continued for another 30 min. The samples were deproteinated and analyzed by electrophoresis in 1% agarose gels. The gels were dried and autoradiographed and analyzed using a Typhoon FLA 7000 biomolecular imager as described above for EMSA. The dried gels were then rehydrated by soaking in water, detached from the Amersham Biosciences Hybond-N+ membrane, stained with ethidium bromide, and analyzed as described above for EcoRI cleavage of R-loops.

Reconstitution of DNA synthesis restart using R-loops

RPA-promoted R-loop formation between pUC19 and 32P-labeled 48-mer RNA (no. 501) was carried out in buffer containing 25 mM Tris-acetate, pH 7.5, 1 mM magnesium acetate, 100 μM each of four dNTPs, 250 μg/ml BSA, and 10 mM DTT for 3 h at 37°C. Then KCl was added to a final concentration of 40 mM. To initiate DNA synthesis from R-loops (3 nM, molecules), 9-μl aliquots were mixed with DNA pol α (50 nM), pol η (38 nM), or pol ε (50 nM) and incubated for 30 min at 37°C. The addition of the DNA polymerases increased final KCl concentration to 60 mM.

For RNA extension by pol δ, RPA-promoted R-loop formation was performed in standard buffer A for 3 h at 37°C. Reconstitution reactions (10 μl) contained R-loops (3 nM, molecules), RFC (8 nM), PCNA (48 nM), pol δ (0.5 nM), 30 mM Tris-acetate, pH 7.5, 5 mM magnesium acetate, 100 μM each of four dNTPs, 1 mM ATP, 250 μg/ml BSA, 10 mM DTT, 60 mM KCl. R-loops were preincubated with RCF and PCNA for 5 min at 37°C, and then pol δ was added, and incubation was continued for another 30 min.

All DNA polymerization reactions were terminated by adding 15 μl of 99.9% formamide, containing 0.1% bromphenol blue. The samples were heated for 4 min at 80°C, and the products of RNA extension were analyzed by electrophoresis in an 8% denaturing PAGE (19:1), containing 50% urea. The migration markers M1–M3 were 63-nt RNA, 70–150-nt ssDNA oligonucleotides, and 166–910-nt denatured Ddel restriction fragments of pUC19, respectively. After electrophoresis, the gels were fixed in 10% glacial acetic acid and 10% ethanol for 20 min at room temperature, dried, and analyzed using a Typhoon FLA 7000 biomolecular imager.

RNA extension by DNA polymerases using deproteinized R-loops

RPA-promoted R-loop formation was performed in buffer A for 3 h at 37°C. The R-loops were deproteinized by treatment with proteinase K (1 mg/ml) and 0.8% SDS for 30 min at 37°C, and then 1 mM EDTA, pH 8.0, was added to chelate magnesium ions. The deproteinized R-loops were purified by passing twice
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through S-400 spin columns (GE Healthcare) equilibrated with 25 mM Tris-HCl (pH 7.5) and 25 mM KCl. The purified R-loops were supplemented with 1 mM of magnesium acetate and kept at −20 °C.

Reactions (10 μl) were initiated by adding DNA pol α (50 nM), pol η (4 nM), or pol ε (50 nM) to deproteinized R-loops (1 nM) in 30 mM Tris-HCl (pH 7.5), 1 mM magnesium acetate, 100 μM each of four dNTPs, 250 μg/ml BSA, 10 mM DTT, 60 mM KCl and carried out for 30 min at 37 °C. For DNA pol δ, the reactions (10 μl) were carried out in 30 mM Tris-acetate, pH 7.5, 5 mM magnesium acetate, 100 μM each of four dNTPs, 1 mM ATP, 250 μg/ml BSA, 10 mM DTT, and 60 mM KCl. Deproteinized R-loops (1 nM, molecules) were preincubated with RCF (2 nM) and PCNA (10 nM) for 5 min at 37 °C, and then pol δ (0.5 nM) was added, followed by incubation for another 30 min.

Quantification and statistical analysis

For statistical analysis, GraphPad Prism 5 software was used. In vitro experiments were repeated at least three times; S.E. values are presented on the graphs.

Data availability

All data are contained within the article and supporting information.

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Abbreviations—The abbreviations used are: RPA, replication protein A; EMSA, electrophoretic mobility shift assay; nt, nucleotide (s); pol, polymerase; RFC, replication factor C; PCNA, proliferating cell nuclear antigen; SUMO, small ubiquitin-like modifier.

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