Human ribonuclease H1 resolves R-loops and thereby enables progression of the DNA replication fork

Faithful DNA replication is essential for genome stability. To ensure accurate replication, numerous complex and redundant replication and repair mechanisms function in tandem with the core replication proteins to ensure DNA replication continues even when replication challenges are present that could impede progression of the replication fork. A unique topological challenge to the replication machinery is posed by RNA–DNA hybrids, commonly referred to as R-loops. Although R-loops play important roles in gene expression and recombination at immunoglobulin sites, their persistence is thought to interfere with DNA replication by slowing or impeding replication fork progression. Therefore, it is of interest to identify DNA-associated enzymes that help resolve replication-impeding R-loops. Here, using DNA fiber analysis, we demonstrate that human ribonuclease H1 (RNH1) plays an important role in replication fork movement in the mammalian nucleus by resolving R-loops. We found that RNH1 depletion results in accumulation of RNA–DNA hybrids, slowing of replication forks, and increased DNA damage. Our data uncovered a role for RNH1 in global DNA replication in the mammalian nucleus. Because accumulation of RNA–DNA hybrids is linked to various human cancers and neurodegenerative disorders, our study raises the possibility that replication fork progression might be impeded, adding to increased genomic instability and contributing to disease.

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3 The abbreviations used are: SETX, Senataxin; AQR, Aquarius; RNAH, ribonuclease H; CO, chromosomal orientation; siCtrl, control siRNA; siRNH1, RNH1-directed siRNA; DIP, DNA–RNA immunoprecipitation; maRTA, microfluidic-assisted replication track analysis; CldU, 5-chloro-2'-deoxyuridine; IdU, 5-ido-2'-deoxyuridine; BrdC, 5-bromo-2'-deoxyuridine; qRT, quantitative reverse transcription; γH2AX, histone H2AX phosphorylation at Ser-139; FAM, 6-fluorescein amidite.
ing (13). Ectopic expression of RNH1 in yeast is sufficient to minimize transcription-dependent hyper-recombination, pausing of the replication fork, and hydroxyurea sensitivity (14–16). In mammalian cells, RNH1 has an established role in mitochondrial DNA replication, and its deletion is embryonically lethal, demonstrating that RNH2 cannot compensate in this setting (17–20). RNH1 localizes to the mammalian nucleus (21), and ectopic RNH1 expression is routinely exploited in mammalian cells to resolve RNA–DNA hybrids. Recently, RNH1 was shown to prevent unwanted recombination events at the telomere by resolving telomeric RNA–DNA hybrids in cells that utilize the alternative lengthening mechanism of telomere maintenance (22). Another recent study identified a link between DNA damage and the accumulation of RNA–DNA hybrids at the telomere (23). However, whether RNH1 plays a role in nuclear DNA replication outside of telomeres remains to be explored.

Given the role of RNA–DNA hybrids in replication impairment and the ability of RNH1 to resolve such hybrids, we sought to determine whether RNH1 impacts genomic integrity in the mammalian nucleus and if so how. We depleted RNH1 from human cell lines and found that RNH1 depletion resulted in increased RNA–DNA hybrids, DNA damage response, and slowing of DNA replication forks. Importantly, these phenotypes were dependent upon RNH1 nuclease activity, suggesting that the hybrids were responsible for these phenotypes. Our studies uncover a novel role of RNH1 in the mammalian nucleus and extend its important function in nuclear DNA replication.

Results

**RNH1 contributes to genome stability and preserves telomere integrity**

Although RNH2 has well-ascribed functions in the mammalian nucleus, the role of RNH1 has remained more obscure. Because R-loops form throughout the genome and RNH1 can resolve R-loops that would pose barriers to the replication machinery, we hypothesized that RNH1 might play an important role in the nucleus and that its loss might perturb replication fork progression and thus elicit a DNA damage response (4). To test this hypothesis, we depleted RNH1 from normal, checkpoint-competent RPE1 cells and measured the levels of histone H2AX phosphorylation at Ser-139 (checkpoint-competent RPE1 cells, and measured the levels in this setting (17–20). RNH1 localizes to the mammalian nucleus (21), and ectopic RNH1 expression is routinely exploited in mammalian cells to resolve RNA–DNA hybrids. Recently, RNH1 was shown to prevent unwanted recombination events at the telomere by resolving telomeric RNA–DNA hybrids in cells that utilize the alternative lengthening mechanism of telomere maintenance (22). Another recent study identified a link between DNA damage and the accumulation of RNA–DNA hybrids at the telomere (23). However, whether RNH1 plays a role in nuclear DNA replication outside of telomeres remains to be explored.

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**Nuclear RNA–DNA hybrid levels increase upon RNH1 depletion**

To demonstrate that RNA–DNA hybrids were responsible for the DNA damage and telomere loss phenotype upon RNH1 loss, we next measured the hybrid levels in the nucleus. We treated 293T cells with control (siCtrl) or RNH1-directed (siRNH1) siRNAs and collected cells 48 h later. Transfection with siRNH1 led to a significant reduction in RNH1 mRNA levels (2.5-fold) (Fig. 2A) and protein levels (3.5-fold) (Fig. 2B) compared with levels present in siCtrl cells. To measure the amount of RNA–DNA hybrids in control versus RNH1-depleted cells, we next extracted nuclear DNA lysate and subjected it to DNA–RNA immunoprecipitation (DIP) using the well-characterized RNA–DNA hybrid antibody S9.6 (27). We conducted a genomic quantitative PCR on a well-characterized hybrid-forming 5′ pause site of β-actin gene as a readout of hybrids. As a control for specificity, we also pretreated lysates with recombinant RNaseH enzyme in vitro to degrade existing RNA–DNA hybrids in both control and depleted cells. As expected, pretreatment with an in vitro RNaseH enzyme led to a 1.8-fold reduction of RNA–DNA hybrids in control and a 3.5-fold reduction in RNH1-depleted cells, confirming the specificity of the S9.6 antibody. Additionally, immunoprecipitation with an IgG control antibody failed to precipitate RNA–DNA hybrids, indicating that the signals we measured were bona fide RNA–DNA hybrids. Analysis of immunoprecipitations from RNH1-depleted cells revealed a significant 2-fold increase in the nuclear RNA–DNA hybrids compared with those in control cells (Fig. 2C). To further corroborate these findings, we also utilized the S9.6 antibody and carried out immunofluorescence on RPE1 cells. As expected, RNH1-depleted cells showed increased levels of RNA–DNA hybrids as represented by elevated S9.6 signal in the nucleus compared with that in the control cells (Fig. 2D and E). Together, these data demonstrate that RNH1 depletion can lead to a significant increase in RNA–DNA hybrids and that this increase correlates with increased DNA damage and telomere loss.
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RNH1 depletion results in replication fork slowing and increased termination and stalling

Given the increased RNA–DNA hybrids, DNA damage, and loss of both telomeric ends, indicative of a replication defect following RNH1 depletion, we hypothesized that RNA–DNA hybrids pose barriers to DNA replication forks. This hypothesis was supported by previous studies showing that the removal of RNA–DNA hybrids by ectopically expressed RNH1 can directly affect replication fork movement in yeast (28). To test this hypothesis, we used microfluidic-assisted replication track analysis (maRTA) to directly measure replication fork progression in RPE1 cells depleted of RNH1. To restrict our analysis to progressing replication forks, we measured IdU tracks that were directly preceded by a CldU track (Fig. 3B). Measuring the lengths of these IdU tracks, we found that RNH1-depleted cells had an average IdU track length of 9.3 ± 0.3 μm, whereas in the control cells was 14.5 ± 0.4 μm, indicating that the replication forks moved significantly slower in RNH1-depleted cells compared with control cells (Fig. 3C). Given the increase in RNA–DNA hybrids associated with RNH1 loss, these data suggest that RNH1 facilitates efficient DNA replication by clearing RNA–DNA hybrids that would otherwise impede replication fork progression during S phase.

To understand how the loss of RNH1 perturbed replication dynamics, we next asked whether other replication parameters including termination, stalling, and origin firing were affected upon RNH1 depletion. Premature termination and stalling events correspond to CldU (red)-only tracks (Fig. 3B). RNH1-depleted cells showed a significant 1.4-fold increase in termination and/or stalling events compared with control cells (Fig. 3C). Given the increase in RNA–DNA hybrids associated with RNH1 loss, these data suggest that RNH1 facilitates efficient DNA replication by clearing RNA–DNA hybrids that would otherwise impede replication fork progression during S phase.

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Figure 1. RNH1 contributes to genome stability and preserves telomere integrity.
A. Western analysis of RNH1 expression and γH2AX in control (shCtrl) and RNH1-depleted RPE1 cells (shRNH1). Bleomycin-treated cells (Bleo) served as a positive control for γH2AX. α-Tubulin is shown as a loading control. Molecular mass in kilodaltons is marked to the right for reference. B. Quantification of γH2AX intensity in shCtrl and shRNH1 cells from the Western blot in A. C. Representative metaphase chromosomes processed with CO-FISH from shCtrl or shRNH1 RPE1 cells. Leading strand-replicated telomeres are green, and lagging strand-replicated telomeres are red. Regions marked by white asterisks are magnified; white arrows indicate telomere free ends (TFE) in magnified images. D. Representative quantification of telomere loss in shCtrl and shRNH1 RPE1 cells. A minimum of 700 metaphase chromosomes were analyzed. p values were computed using a two-tailed Student’s t test (*, p < 0.05). Error bars represent S.E.
or red flanked by green on both sides was analyzed (Fig. 3B). However, no significant differences in origin firing were observed between RNH1-depleted and control cells (Fig. 3E). Collectively, these results indicate that RNH1 plays an important role in assisting fork movement during DNA replication. We suggest that RNH1 does this by resolving RNA–DNA hybrids that pose barriers to a progressing replication fork. Furthermore, our data provide the first evidence that RNH1 plays a role in global DNA replication in the mammalian nucleus.

Nuclease activity of RNH1 is required for efficient replication fork movement

To further characterize the role of RNH1 in DNA replication, we tested whether the nuclease function of RNH1 was required for this activity. To do this, we created a series of RPE1 cell lines ectopically expressing either a GFP-tagged wild-type (WT) or a well-characterized GFP-tagged nuclease-dead (D145N) form of RNH1 (32, 33). To establish a direct role in the nucleus, these RNH1 constructs lacked the mitochondrial targeting sequence present in the endogenous gene, thereby restricting their expression to the nucleus. We confirmed the nuclear localization of ectopically expressed proteins by visualizing GFP expression only in the nucleus (Fig. 4A). These stable RPE1 cells were transfected with an siRNH1 directed toward the 3′-UTR that did not target the ectopically expressed protein. We observed that RNH1 depletion using a 3′-UTR siRNA was comparable with that of a previously used coding sequence targeting siRNA (Fig. 4B). We also observed robust expression of our ectopically tagged RNH1 proteins and significant depletion of endogenous RNH1 levels (Fig. 4B).

Next, using maRTA, we again measured replication fork movement and found that ectopic expression of WT RNH1 restored fork movement to the levels observed in siCtrl cells (Fig. 4C). Indeed, although RNH1-depleted forks moved an IdU length of 8 μm, ectopic expression of WT RNH1 increased this to 12 μm, levels we observed in siCtrl cells. In contrast, ectopic expression of the catalytically dead D145N allele of RNH1 failed to rescue the replication fork movement defect (fork movement was 7.9 μm, nearly identical to that observed in RNH1-depleted cells). Together, these findings demonstrated that the nuclease activity of RNH1 is required for unperturbed movement of replication forks in mammalian cells. Similarly, we also measured fork termination and stalling events upon ectopic expression of WT and D145N alleles in RNH1-depleted cells. As expected, both events were reversed by ectopic expression of WT RNH1 but not the nuclease-dead allele, thereby reiterating the importance of the nuclease function of RNH1 in the fidelity of replication fork progression (Fig. 4D). Neither WT nor the D145N allele of RNH1 affected the levels of origin firing in RNH1-depleted cells (Fig. 4E). These data suggest that the nuclease activity of RNH1 is required for resolution of RNA–DNA hybrids during DNA replication.
hybrids and therefore efficient movement of replication forks during nuclear DNA replication.

Discussion

Our study establishes a role for RNH1 in genomic DNA replication. Indeed, we illustrate for the first time that RNH1 nuclease activity is required for efficient fork movement during nuclear DNA replication. Furthermore, we have established a correlation between the accumulation of RNA–DNA hybrids and replication defects observed upon RNH1 depletion. Taken together, we propose a model wherein RNH1 resolves RNA–DNA hybrids to assist the replication machinery in its uninterrupted movement during DNA replication.

The unscheduled formation and stabilization of RNA–DNA hybrids have been postulated to be detrimental to the replication machinery. The importance of resolving these structures is probably best underscored by the multitude of proteins that act on RNA–DNA structures. Indeed, helicases such as SETX, AQR, and DHX9 in mammalian cells and Sen1 and PIF1 in yeast have all been shown to resolve RNA–DNA hybrids (9, 34–36). Here, we add RNH1 to a growing list of proteins by showing that endogenous RNH1 is required to similarly remove RNA–DNA hybrids and that if these hybrids are not removed replication is significantly impacted. However, how RNH1 is regulated in the nucleus and how its activity is coordinated with the replication machinery remain unclear. A recent study from Nguyen et al. (37) elegantly demonstrated that replication protein A can interact with RNH1 and stimulate its activity, raising the possibility that RNH1 is tightly regulated by the DNA replication and repair machinery.

Given that RNH1 loss elicits replication defects such as fork slowing, termination, and fork stalling, it will be critical to determine how this impacts checkpoint activation and cell cycle progression. Furthermore, understanding the fate of accumulated RNA–DNA hybrids upon RNH1 depletion is another interesting avenue worth pursuing. As previously
noted, RNA–DNA hybrids arising from different sources can be processed via separate mechanisms (9). For example, those involved in class switch recombination are not processed via nucleotide excision repair, whereas those arising from loss of some RNA processing factors or camptothecin treatment are processed by nucleotide excision repair. It is also worth evaluating whether redundant nucleases and helicases including RNH2, SETX, and AQR could rescue the effects of RNH1 loss.

The study of R-loops and their resolution have sparked more attention in recent years due to the fact that R-loops are associated with a number of diseases including cancer and several neurodegenerative disorders (38). This underscores a need for understanding these structures and their origins, stabilization, and resolution along with their impact on cellular processes. By revealing the function of RNH1 in R-loop resolution in the nucleus, our study adds to the diversity of mechanisms targeting such structures. Furthermore,
our study identifies a previously unknown function of RNH1 in nuclear DNA replication. Together, our work broadens the understanding of RNA–DNA structures and places RNH1 as a novel mechanism to resolve those structures and assist in nuclear DNA replication.

**Experimental procedures**

**Cell culture**

Cells were cultured at 37 °C in 5% carbon dioxide and atmospheric oxygen as reported previously (39). 293T cells were obtained from Dr. Robert Weinberg (Massachusetts Institute of Technology) and cultured in high-glucose Dulbecco’s modified Eagle’s medium (DMEM) containing 10% heat-inactivated fetal bovine serum (FBS) and 1% penicillin/streptomycin (Sigma-Aldrich). RPE1 cells were obtained from ATCC and cultured in DMEM/Nutrient Mixture F-12 containing 7.5% heat-inactivated FBS and 1% penicillin/streptomycin.

**siRNA transfection**

siRNA transfection was performed using Invitrogen’s Lipofectamine RNAiMAX reagent (Thermo Fisher Scientific) according to the manufacturer’s protocol. siRNAs used were siCtrl (catalog number 4390843) and siRNH1 (catalog number 4390824, ID s48356) from Life Technologies or siRNA directed to the 3’UTR of RNH1 (hs.Ri.RNASEH1.13.1) from Integrated DNA Technologies.

**Virus production, infections, and stable cell lines**

Lentiviral production and transductions were carried out as reported previously (40). Briefly, 293T cells were transduced with pLKO.1-puro plasmid carrying an shRNA using TransIT-LT1 reagent (Mirus Bio, Madison, WI) and a mixture containing 8:1 ratio of pHR-CMV-8.2 R packaging plasmid and pCMV-VSV-G. Supernatant containing virus was collected 48 h post-transfection and filtered through a 0.45-µm PVDF membrane. RPE1 cells were infected for 4 h each on 2 consecutive days in the presence of 8 µg/ml protamine sulfate (Sigma). Following infection, transduced cells were selected with 15 µg/ml puromycin sulfate. Stable RPE1 cell lines were prepared by using either a GFP-tagged D145N RNH1 construct (a generous gift from Dr. Marteijn) (32) or its wild-type version (modified from D145N construct using site-directed mutagenesis; Agilent Technologies).

**Western blot analysis**

Western blot analysis was carried out as described previously with modifications (41). Briefly, cells were washed with PBS and lysed in radioimmunoprecipitation assay buffer (150 mM NaCl, 50 mM Tris-HCl, pH 8.0, 1.0% Nonidet P-40, 0.5% sodium deoxycholate, and 0.1% SDS) supplemented with 1 mM sodium orthovanadate, 1 mM sodium fluoride, 1 mM microcystin LR, 2 mM phenylmethylsulfonyl fluoride, protease inhibitor mixture (Sigma), and phosphatase inhibitor mixture set 1 (EMD Millipore, Billerica, MA). Following sonication and centrifugation, supernatant lystate was quantified using a protein assay (Bio-Rad). Lysates were subjected to SDS-PAGE and transferred to PVDF membranes for blotting. The following antibodies were used: mouse monoclonal anti-RNase H1 (H00246243-M01, Novus Biologicals, Littleton, CO), rat monoclonal anti-tubulin (NB600-506, Novus Biologicals), and mouse monoclonal anti-phospho(Ser-139) H2AX (05-636, Millipore).

**Metaphase chromosome preparation**

Metaphase chromosomes were prepared as described previously (42). Briefly, cultured RPE1 cells were treated with 0.5 µg/ml Colcemid (Sigma) for 6 h. Arrested metaphase cells were collected by mitotic shake-off, treated with 75 mM potassium chloride, and fixed in 3:1 solution of methanol and acetic acid. Chromosomes were spread by dropping onto glass slides. For analysis via CO-FISH, 0.3 µg/ml BrdU (Sigma) and 0.1 µg/ml BrdC (MP Biomedicals, Santa Ana, CA) were added to the cultured medium 18 h prior to collection of the cells.

**CO-FISH**

CO-FISH was performed as described previously with modifications (43). Briefly, spread metaphase chromosomes were aged at 65 °C for 18 h. Aged chromosomes were rehydrated in PBS, treated with 100 µg/ml RNase at 37 °C for 10 min, and refixed in 4% paraformaldehyde at room temperature for 10 min. Fixed chromosomes were UV-sensitized in 0.5 µg/ml Hoechst 33258 (Sigma) in 2× SSC at room temperature for 15 min and exposed to 365 nm UV light for 1 h using a UV cross-linker (Vilber-Lourmat, Marne-la-Vallée, France). Exposed chromosomes were digested with 3 units/µl exonuclease III (Promega, Madison, WI) at room temperature for 15 min, denatured in 70% formamide in 2× SSC, and dehydrated in cold ethanol before hybridization. Chromosomes were hybridized first using a 0.03 µg/ml concentration of a leading strand telomere PNA probe (FAM-(TTAGGG)6) followed by a 0.03 µg/ml concentration of a lagging strand PNA probe (Cy3-CCCTAA)6 (both probes from PNA Bio, Thousand Oaks, CA). Hybridized chromosomes were mounted using ProLong Gold (Life Technologies) with 125 ng/ml DAPI.

**maRTA**

maRTA was conducted as described previously (29, 44). Briefly, asynchronous RPE1 cells were labeled for 30 min each with 50 µM CldU and 50 µM IdU with two PBS washes in between. Labeled cells were collected and embedded in agarose plugs for lysis and DNA extraction. DNA was subsequently stretched, denatured, and subjected to immunostaining. Antibodies used were rat anti-CldU/BrdU (Abcam, ab6326), mouse anti-IdU/BrdU (BD Biosciences, 347580), goat anti-rat Alexa Fluor 594 (Invitrogen, A11007), and goat anti-mouse Alexa Fluor 488 (Invitrogen, A11001).

**Fluorescence imaging**

Metaphase chromosomes from CO-FISH and labeled DNA tracks from maRTA were imaged on a Nikon 90i epifluorescence microscope using a 100× 1.40 numerical aperture Plan Apo VC objective (Nikon Instruments, Melville, NY) with Cargille Type LDF immersion oil (Cargille Sacher Laboratories, Cedar Grove, NJ). Images were captured using a CoolSnap HQ2 charge-coupled device camera (Photometrics, Tucson, AZ) and deconvoluted with a blind algorithm using NISElements AR
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(Chromosomes) prior to quantification. RPE1 cells stably expressing GFP-tagged WT and D145N RNH1 were visualized and captured without any staining.

DIP

DIP was performed as described previously with modifications (6). Briefly, 293T cells were pelleted and resuspended in DIP lysis buffer (0.5% Nonidet P-40, 85 mM potassium chloride, and 5 mM PIPES). Following centrifugation, pelleted nuclei were lysed in DIP nuclear lysis buffer (1% sodium dodecyl sulfate, 25 mM Tris-HCl, pH 8, and 5 mM EDTA), sheared, and digested with two sequential rounds of 100 μg of proteinase K for 1.5 h each at 55 °C. DNA was phenol/chloroform-extracted and ethanol-precipitated (45) at which point one-half was subjected to an overnight digestion with recombinant ribonuclease H (Roche Applied Science). Samples were then diluted in DIP dilution buffer (1.1% Triton X-100, 0.01% sodium dodecyl sulfate, 1.2 mM EDTA, 16.7 mM Tris-HCl, pH 8, and 166.5 mM sodium chloride) and sonicated to generate ~200-bp-long DNA fragments. Resulting DNA was quantified using the PicoGreen assay following the manufacturer’s protocol (Life Technologies). Resulting DNA was quantified using the PicoGreen assay following the manufacturer’s protocol (Life Technologies). Resulting DNA was quantified using the PicoGreen assay following the manufacturer’s protocol (Life Technologies).

S9.6 immunofluorescence

S9.6 immunofluorescence was performed as described previously (9). Briefly, RPE1 cells transfected with siCtrl or siRNH1 were fixed with ice-cold methanol for 5 min at −20 °C. Fixed cells were blocked in 2% BSA and PBS for an hour at room temperature followed by incubation with the S9.6 primary antibody (1:200 dilution; 1 μg/ml) and goat anti-mouse Alexa Fluor 594-conjugated secondary antibody (1:1000) for 1 h each at room temperature. Finally, cells were washed in 0.5 μg/ml Hoechst 33258 in PBS to label the nuclei and mounted using ProLong Gold. Images were taken at 40× using a Nikon 90i epifluorescence microscope as described above. Only the nuclear staining of S9.6 signal was considered and analyzed using ImageJ. 1.50i.

Genomic quantitative PCR

Genomic quantitative PCR was performed using Power SYBR Green Master Mix (Life Technologies) following the manufacturer’s protocol. The 5′ region of the β-actin pause element (5′ pause site), known to form RNA–DNA hybrids, was amplified to assess hybrid formation. Reaction conditions were as described in the manufacturer’s instructions with 58.7 °C as the annealing temperature. Primers used were: 5′-TTACCC AGA GTG CAG GTG TG-3′ (forward) and 5′-CCC CAA TAA GCA GGA ACA GA-3′ (reverse).

Quantitative reverse transcription-PCR (qRT-PCR)

qRT-PCR was performed as described previously (39). Target genes used were RNH1 (Hs.PT.39a.22214836, Integrated DNA Technologies) and GAPDH (Hs.PT.39a.22214836, Integrated DNA Technologies).

Author contributions—Experiments were designed by S. A. S., A. V., S. P., and D. C. T. Experiments were performed by S. P., D. C. T., and J. J. Data were analyzed by S. A. S., A. V., B. M., and S. P. The manuscript was written by S. A. S., A. V., and S. P.

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