Yra1-bound RNA–DNA hybrids cause orientation-independent transcription–replication collisions and telomere instability

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R loops are an important source of genome instability, largely due to their negative impact on replication progression. Yra1/ALY is an abundant RNA-binding factor conserved from yeast to humans and required for mRNA export, but its excess causes lethality and genome instability. Here, we show that, in addition to ssDNA and ssRNA, Yra1 binds RNA–DNA hybrids in vitro and, when artificially overexpressed, can be recruited to chromatin in an RNA–DNA hybrid-dependent manner, stabilizing R loops and converting them into replication obstacles in vivo. Importantly, an excess of Yra1 increases R-loop-mediated genome instability caused by transcription–replication collisions regardless of whether they are codirectional or head-on. It also induces telomere shortening in telomerase-negative cells and accelerates senescence, consistent with a defect in telomere replication. Our results indicate that RNA–DNA hybrids form transiently in cells regardless of replication and, after stabilization by excess Yra1, compromise genome integrity, in agreement with a two-step model of R-loop-mediated genome instability. This work opens new perspectives to understand transcription-associated genome instability in repair-deficient cells, including tumoral cells.

[Keywords: Yra1; R loop; transcription–replication collision; telomeres]

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RNA–DNA hybrids are produced cotranscriptionally when the nascent transcript threads back hybridizing with the template DNA, leading together with the displaced nontemplate ssDNA to a structure termed the R loop. Hybridss have a tendency to accumulate preferentially at highly transcribed protein-coding genes, peaking at promoters and terminators, rDNA, tRNA-coding genes, Ty elements, centromeres, and telomeres (Ginno et al. 2012, Chan et al. 2014, El Hage et al. 2014, Wahba et al. 2016). RNA–DNA hybrids may benefit cell physiology, as shown in some cases of transcription initiation and termination, mitochondrial DNA replication, or immunoglobulin class switching (Aguilera and García-Muse 2012). However, RNA–DNA hybrids may also have a strong impact on genome instability, as shown in cells defective in specific mRNP assembly factors such as the THO complex or the SRSF1 RNA-binding protein, topoisomerase I, RNA–DNA helicases, or RNase H, a ribonuclease that specifically degrades the RNA moiety of RNA–DNA hybrids [Santos-Pereira and Aguilera 2015; Sollier and Cimprich 2015]. Accumulating evidence indicates that most of this genetic instability is due to the ability of R loops to stall the progression of the replication fork, leading to its collapse (García-Muse and Aguilera 2016). Thus, RNA–DNA helicases are required for the replication of highly transcribed regions [Boubakri et al. 2010], and R-loop accumulation impairs replication fork progression from bacteria to human cell lines [Welling et al. 2006; Tuduri et al. 2009; Gan et al. 2011; Castellano-Pozo et al. 2012, Hamperl et al. 2017]. Alternatively, R loops may generate genomic instability if the displaced
ssDNA is recognized and processed by flap endonucleases (Sollier et al. 2014). Despite reports showing that R loops alter replication, causing genome instability, the mechanism is still unclear. However, recent evidence indicates that R loops are not deleterious per se but require a second step, such as a local chromatin compaction, to compromise genome integrity (Castellano-Pozo et al. 2012; García-Pichardo et al. 2017).

RNA–DNA hybrids also have a physiological role at telomeres. Telomeres are transcribed into a long non-coding RNA [lncRNA] called TERRA that recruits telomeric proteins, contributes to heterochromatin formation (Maicher et al. 2014), and prevents activation of the DNA damage response (Flynn et al. 2011). A small proportion of TERRA RNA hybridizes with the DNA, forming telomeric RNA–DNA hybrids. This is restrained by the actions of RNase H and the THO complex (Balk et al. 2013; Pleifler et al. 2013; Arora et al. 2014; Yu et al. 2014) and favored by telomere shortening. TERRA RNA–DNA hybrids promote homologous recombination (Graf et al. 2017) between telomeric repeats, enabling alternative lengthening of telomeres (ALT), a mechanism used by telomerase-deficient tumor cells to prevent telomere shortening (Balk et al. 2013; Arora et al. 2014; Yu et al. 2014).

A number of transcription and RNA processing factors, such as THO or SRSF1, control cotranscriptional R-loop formation (Santos-Pereira and Aguilera 2015). Yra1 is an RNA-binding protein conserved in metazoans [ALY/REF] that acts as an adaptor for mRNA export factors (Kohler and Hurt 2007). It is cotranscriptionally loaded onto RNA by directly interacting with the C-terminal domain [CTD] of RNA polymerase II [RNAPII] (MacKellar and Greenleaf 2011) or with other mRNP assembly factors (Johnson et al. 2011; Ma et al. 2013; Ren et al. 2017). Interestingly, Yra1 stoichiometry is tightly regulated in the cell via a mechanism relying on Yra1 inhibition of YRA1 pre-mRNA splicing (Rodríguez-Navarro et al. 2002; Preker and Guthrie 2006; Dong et al. 2007). Thus, removal of the YRA1 intron from the gene bypasses this autoregulatory circuit, causing Yra1 overexpression with a strong negative impact on mRNA export and cell viability (Rodríguez-Navarro et al. 2002; Preker and Guthrie 2006). Interestingly, we reported recently that high Yra1 intron levels alter genome dynamics by accumulating DNA damage and causing transcription-associated spontaneous recombination that is suppressed by RNase H overexpression (Gavaldá et al. 2016).

Aiming to understand how Yra1 overexpression and transcription compromise genome integrity, we demonstrate here that excess Yra1 is recruited to chromatin in an RNA–DNA hybrid-dependent manner, thereby stabilizing R loops and converting them into genome integrity threats. Importantly, an excess of Yra1 increases R-loop-mediated genome instability regardless of transcription–replication orientation. Consistent with the fact that Yra1 also binds to telomeres, where R loops accumulate, we show that Yra1 overexpression causes an increase of RNA–DNA hybrids at telomeres, telomere alterations, and accelerated senescence in telomerase-deficient cells.

Our results demonstrate not only that excess of Yra1/ALY binds and stabilizes R loops but that R loops are transiently formed in cells regardless of replication, although they need to be stabilized to compromise genome integrity.

Results

Excess Yra1 binds to chromatin in an RNA–DNA hybrid-dependent manner

Since YRA1 overexpression increases genomic instability in an RNase H-sensitive manner (Gavaldá et al. 2016), we wondered whether overexpressed Yra1 stabilized naturally formed RNA–DNA hybrids. For this, we analyzed whether localization of overexpressed Yra1 to transcribed genes was dependent on RNA–DNA hybrids. We performed an Yra1 chromatin immunoprecipitation (ChIP) in wild-type cells overexpressing or not overexpressing YRA1 using an HA-tagged Yra1 protein expressed from either an intron-deficient [HA-YRA1α] or an intron-containing [HA-YRA1] version of YRA1, respectively (Gavaldá et al. 2016). To determine whether putative Yra1 binding to chromatin was dependent on the presence of RNA–DNA hybrids, we overexpressed RNase H, an enzyme that specifically degrades the RNA moiety of RNA–DNA hybrids. Under YRA1-overexpressing conditions, we found a clear increase of Yra1 recruitment to the endogenous GCN4 and PDR5 genes previously reported to accumulate RNA–DNA hybrids (García-Benítez et al. 2017) and to the YRA1 intron region. Overexpressing RNase H in the cell, we restored Yra1 basal levels in all of the regions analyzed (Fig. 1A). These observations argue that Yra1 is recruited to RNA–DNA hybrids when artifically overexpressed in addition to their natural putative RNA.

Yra1 increases RNA–DNA hybrid accumulation in vivo

Next, we determined whether RNA–DNA hybrids were increased, presumably due to stabilization, when YRA1 was overexpressed. We performed a DNA–RNA immunoprecipitation [DRIP] with the S9.6 antibody in wild-type cells overexpressing [HA-YRA1α] or not overexpressing [HA-YRA1] YRA1 to detect the hybrids. We focused on the genomic regions where we showed a hybrid-dependent Yra1 localization: GCN4, PDR5, and YRA1 introns. We observed a S9.6 signal in the three regions analyzed that was significantly reduced by RNase H treatment [Fig. 1B]. The RNA–DNA hybrids detected in GCN4, PDR5, and YRA1 introns were significantly increased when YRA1 was overexpressed [Fig. 1B].

Next, we examined whether Yra1 recruitment was enriched at regions naturally forming RNA–DNA hybrids genome-wide when overexpressed. We performed S9.6 DRIP-seq [DRIP combined with high-throughput sequencing] analysis in wild-type cells and compared the data with our previously published ChIP–chip data on Yra1 recruitment [HA-Yra1 and HA-Yra1α] (Gavaldá et al. 2016). Our DRIP-seq data are consistent with previously published S9.6 ChIP-seq [ChIP combined with high-
Yra1 in mutants that accumulate R loops, such as *hpr1Δ*, *mtf1Δ*, or *tho2Δ* mutants lacking the THO complex involved in mRNP biogenesis or the double-mutant *top1Δ top2-1* that accumulates negative supercoiling favoring R loops [Supplemental Fig. S2]. As can be seen, the results showing a decrease in cell viability suggest that the increase in persistent RNA–DNA hybrids in *YRA1*-overexpressing cells could contribute to cell death.

**An mRNA export defect is not sufficient to increase R loops**

Since overexpression of Yra1 impairs mRNA export [Rodríguez-Navarro et al. 2002], we wanted to confirm that the nuclear mRNA accumulation resulting from an mRNA export defect was not sufficient to increase R loops. We analyzed three different nucleoporin mutants (*nup42Δ* and *nup60Δ*, known to be affected in mRNA export, and *nup100Δ*, not affected) [Bonnet and Palancade 2014] and the positive control (*mlp1Δ*, a nuclear pore mutant known to accumulate R loops) [García-Benítez et al. 2017]. R loops were inferred using a genetic method based on the hyperrecombination ability of the human activation-induced cytidine deaminase (hAID), an enzyme that modifies cytidines in the ssDNA moiety of the R loop, as shown previously [García-Benítez et al. 2017; García-Pichardo et al. 2017]. Recombination in *nup42Δ* and *nup100Δ* was similar to the wild type, whereas in *nup60Δ* it was increased (4.3×), but this increase was not suppressed by RNase H overexpression, consistent with its known sensitivity to HU and MMS [Niño et al. 2016] and in contrast to the positive control *mlp1Δ* [Supplemental Fig. S3]. Therefore, we conclude that accumulation of mRNA in the nucleus due to an RNA export defect does not induce R loops per se.

**Yra1 binds RNA–DNA hybrids in vitro**

Since Yra1 localizes to RNA–DNA hybrid-enriched regions and is a well-characterized RNA-binding protein, we reasoned that Yra1 might directly bind to RNA–DNA hybrids. To test this idea, we isolated a recombinant His6-tagged version of Yra1 from bacteria, confirming it as shown previously (García-Benítez et al. 2017; García-Pichardo et al. 2017). Recombination in *nup42Δ* and *nup100Δ* was similar to the wild type, whereas in *nup60Δ* it was increased (4.3×), but this increase was not suppressed by RNase H overexpression, consistent with its known sensitivity to HU and MMS [Niño et al. 2016] and in contrast to the positive control *mlp1Δ* [Supplemental Fig. S3]. Therefore, we conclude that accumulation of mRNA in the nucleus due to an RNA export defect does not induce R loops per se.

**Figure 1. YRA1 overexpression causes RNA–DNA hybrid accumulation. (A) ChIP analysis of Yra1 using anti-HA antibody in a wild-type strain overexpressing [green, HA-YRAΔi] or not overexpressing [blue, HA-YRA1] YRA1 and RNaseH [RNH+ or RNH−] at the GCN4 and PDR5 genes or the YRA1 intron. n > 3. (B) DNA–RNA immunoprecipitation (DRI) with the S9.6 antibody in wild-type asynchronous cultures overexpressing [HA-YRAΔi; green] or not overexpressing [HA-YRA1; blue] YRA1 at the GCN4 and PDR5 genes or the YRA1 intron. n > 3. Samples were treated (+) or not (−) in vitro with RNase H [RNH] prior to the immunoprecipitation. Means and SEM are plotted in all panels. (*) P < 0.05, two-tailed Student’s t-test.**
Therefore, Yra1 is able to bind in vitro not only ssRNA but also ssDNA, dsRNA, and a RNA–DNA duplex. To confirm these interactions, we competed Yra1 binding with different cold nucleic acid species. First, we incubated purified Yra1 with labeled dsRNA and increasing amounts of cold ssRNA or RNA–DNA hybrid. Both nucleic acids reverted Yra1 interaction with dsRNA at similar concentrations (5 µM) (Fig. 3B). Next, we challenged the Yra1–hybrid interaction with increasing amounts of cold hybrid, ssRNA, or ssDNA. In agreement with its binding to RNA–DNA hybrids, cold ssRNA, ssDNA, or RNA–DNA hybrids competed for Yra1 binding, since RNA–DNA hybrids are slightly better competitors than ssRNA and ssDNA [Fig. 3C]. Altogether, these results indicate that Yra1 is able to bind RNA–DNA hybrids, supporting the conclusion that when overexpressed in vivo, Yra1 can bind R loops in chromatin, contributing to their stabilization.

R-loop-mediated genome instability is linked to head-on transcription–replication

We showed previously that transcription causes hyperrecombination when occurring in a head-on orientation with respect to replication but not when occurring codirectionally [Prado and Aguilera 2005]. Since RNA–DNA hybrids are an obstacle for replication fork progression [Wellinger et al. 2006; Gan et al. 2011], we examined whether they could explain the orientation-dependent transcription–replication conflicts responsible for genome instability. We used the previously reported plasmids pGAL-OUT and pGAL-IN [Prado and Aguilera
that contain leu2 truncated repeats transcribed from an inducible GAL1 promoter in either a codirectional (OUT) or head-on (IN) orientation with respect to replication driven from the early replication origin ARSH4 [Fig. 4A], respectively. DNA damage driven by the transcription–replication collision would be repaired by recombination between the leu2 direct repeats, generating a wild-type LEU2 gene. Consequently, we could measure genome instability as the frequency of Leu2+ recombinant colonies. We observed that the frequency of recombination in the absence of transcription was low and similar in both systems [Fig. 4B], consistent with previous results [Prado and Aguilera 2005]. However, upon transcription, induction recombination was highly increased [7.7-fold] in the IN system with head-on transcription–replication, whereas only a twofold increase in recombination was observed in the codirectional OUT system [Fig. 4B]. When similar experiments were performed in both systems after RNase H overexpression, the transcription-dependent hyperrecombination observed in the head-on IN system was suppressed, whereas the recombination frequencies in the codirectional OUT system did not change [Fig. 4B]. These results argue that RNA–DNA hybrids are an important source of genome instability in systems undergoing head-on transcription–replication conflicts but not in those undergoing codirectional conflicts. This interpretation would be consistent with the recent observation in human cells using similarly designed plasmid-based constructs in which transcription is driven from a bacterial T7 promoter and replication is driven from an Epstein-Barr virus replication origin [Hamperl et al. 2017] as well with R-loop-dependent replication impairment observed in the Bacillus subtilis genome [Lang et al. 2017]. However, none of these results provide any answer to whether R loops are formed only in head-on transcription–replication or after replication.

Yra1-stabilized R loops induce instability regardless of transcription–replication orientation

If Yra1 binds and stabilizes transient RNA–DNA hybrids present in the genome, we might expect that Yra1 will increase transcription–replication conflicts mediated by these structures. To test this hypothesis, we measured recombination in the codirectional and head-on systems either overexpressing [YRA1Δ] or not overexpressing [YRA1] YRA1. Yra1 overexpression enhanced recombination 7.2 times in the head-on IN system, an increase that was majorly suppressed by RNase H overexpression [Fig. 4C]. Importantly, an 8.7-fold increase was also observed in the codirectional OUT system, which was also suppressed by RNase H [Fig. 4C]. This result indicates that RNA–DNA hybrids also form under codirectional transcription–replication, but such hybrids are not stable.
Figure 4. R-loop-mediated transcription–replication conflicts are orientation-independent in YRA1-overexpressing cells. (A) Schemes of the centromeric plasmids harboring the recombination systems in head-on (IN) and codirectional (OUT) orientation. The arrows indicate the RNAPII-driven transcription orientation of the leu2 gene repeats from the GAL1 promoter and the direction of replication forks initiated at ARSH4. (B) The effect of RNaseH1 overexpression on the recombination of the head-on (IN) and codirectional (OUT) systems in wild-type cells. Cells grown in either glucose (TRX) or galactose (TRX) and overexpressing RNase H (RNH+ or RNH−) or not overexpressing RNase H. The fold increase over no transcription and no RNase H expression is indicated at the right. (C) The effect of RNaseH1 overexpression on the recombination of transcribed IN or OUT plasmids in wild-type cells overexpressing YRA1Δ1 or not overexpressing [YRA1 YRA1] and RNase H (RNH− or RNH+). The fold increase over no overexpression of YRA1 and Rnase H is indicated at the right. Means and SEM are plotted in all panels. Asterisks indicate statistically significant differences between the strains indicated, according to Student’s t-tests. [*] P < 0.05; [**] P < 0.005; [***] P < 0.0005.

enough to drive high genome instability. However, under Yra1 overexpression, hybrids are stabilized, resulting in a significant increase in recombination, presumably by constituting a stable block to replication fork progression regardless of transcription–replication orientation.

Therefore, our hypothesis predicts that RNA–DNA hybrids should be present in both plasmids. Consequently, we performed a DRIP analysis using the S9.6 antibody in samples with or without RNase H treatment. In the analysis, we included the previously described hybrid-accumulating PDR5 gene as an internal positive control and PRE1 as a negative control. The results confirmed the presence of RNA–DNA hybrids in the plasmid-borne LEU2 gene of the recombinant head-on IN system, in agreement with the in vivo data [Fig. 5A]. Notably, we also detected hybrids in the recombined OUT system where LEU2 transcription is codirectional to replication fork progress [Fig. 5A]. Therefore, we conclude that RNA–DNA hybrids are formed during transcription in both codirectional and head-on constructs but significantly induce genomic instability, measured as recombination, when replication and transcription are head-on and not when they are codirectional. However, when the hybrid is stabilized via binding to Yra1, replication would stall regardless of the transcription orientation, leading to a similar increase in recombination [sevenfold to eightfold above the wild-type levels]. These results clearly support that R loops are formed transiently regardless of replication and not as a consequence of transcription–replication conflicts.

Although unlikely, it might be possible that circular plasmids impose a specific topological constraint different from chromosomes that could enhance R-loop accumulation and Yra1 binding. To confirm that this was not the case and that R loops occurred in linear chromosomes, we constructed a transcription–replication collision system in chromosome III and analyzed RNA–DNA hybrids under the same conditions tested in the plasmid construct. To do so, we integrated the LEU2 gene under the inducible GAL1 promoter in a head-on orientation with respect to replication driven from the early-firing ARS315 origin [Fig. 5B], since this orientation was the one with the potential to cause DNA opening that could favor R loops. DRIP analyses of cells cultured in galactose-containing medium [transcription ON] versus glucose-containing medium [transcription off] revealed that RNA–DNA hybrids appeared in the LEU2 gene only when transcribed [Fig. 5B]. In the PDR5 gene used as a positive control, hybrids were observed in both conditions, consistent with the fact that PDR5 is constitutively transcribed in both media [Fig. 5B]. Importantly, the RNA–DNA hybrids formed in the chromosomal transcription–replication collision system increased when YRA1 was overexpressed [Fig. 5B], in agreement with the increase in recombination observed in the plasmid [Fig. 4C].

Cumulative evidence with concomitant studies on RNA–DNA hybrids, γH2AX, and 53BP1 foci and recombination, comet, and DNA-combing assays suggests that R-loop-dependent γH2AX is the result of DNA damage (Garcia-Rubio et al. 2015; Schwab et al. 2015; Salas-Armenteros et al. 2017). To assess whether the RNA–DNA hybrids detected in the LEU2 gene cause DNA damage, we measured H2A-P [the equivalent in yeast to γH2AX]) levels by ChIP under conditions of both active and inactive transcription of the GAL1::LEU2 fusion, using telemeric repeats as a positive control (Kitada et al. 2011). An increase in H2A-P was observed when transcription of LEU2 was active, but not when it was inactive, as well as in the telemeric controls [Fig. 5C]. Therefore, head-on transcription of the chromosomal LEU2 gene also generates DNA breaks. Notably, such breaks were suppressed by RNase H overexpression [Fig. 5C], confirming that they were R-loop-dependent. Most important, such H2A-P foci were significantly increased when Yra1 was overexpressed, and this increase was suppressed by...
R-loop–replication conflicts enhanced by Yra1

We showed recently that an excess of Yra1 causes a cell senescence-like phenotype and a slight telomere shortening in telomerase-positive cells and is enriched at Y′ telomeric regions (Gavalda et al. 2016). Since Yra1 is a highly efficient RNA-binding protein [Fig. 3, Strasser and Hurt 2000] and since telomerase activity in wild-type cells relies on an RNA molecule that is used as a template for telomere synthesis, it was possible that an excess of Yra1 sequestered the telomerase RNA molecule TLC1, leading to a telomerase-deficient phenotype instead of acting directly on the telomere. To test this possibility, we determined the effects of Yra1 overexpression in telomerase-minus cells (est2Δ). Heterozygous EST2/EST2 diploids harboring the pGAL::Yra1Δi plasmid were sporulated in glucose medium. Next, haploid est2Δ pGAL::Yra1Δi spore clones were selected in glucose medium and further propagated in galactose liquid medium via serial dilutions [Hardy et al. 2014]. The senescence profiles, kinetics of telomere shortening, and the type of survivors formed in multiple est2Δ and est2Δ pGAL::Yra1Δi clones grown in galactose were analyzed (Fig. 6). We showed that est2Δ pGAL::Yra1Δi clones (overexpressing Yra1) exhibited a rapid and deep premature senescence compared with est2Δ clones [Fig. 6A]. Strikingly, telomere length analysis at the first time point of the senescence assay indicated that telomeres were shorter in the est2Δ pGAL::Yra1Δi clones compared with est2Δ clones [Fig. 6B–D]. This result suggests an abrupt telomere shortening upon overexpression of Yra1 different from the progressive telomere erosion normally observed in the absence of telomerase. Consistently, survivors appeared earlier in est2Δ cells overexpressing YRA1 [Fig. 6C]. As predicted, RNA–DNA hybrids were increased in est2Δ cells [Fig. 6E] as shown in the single TelVI-R telomere and consistent with previous results indicating that shortening of telomeres in est2Δ cells triggers TERRA transcription and RNA–DNA hybrid accumulation [Graf et al. 2017]. Importantly, such hybrids were significantly increased under Yra1 overexpression in RNA–DNA hybrid stabilization [Fig. 6E].

Figure 5. Effect of Yra1 overexpression on transcription–replication collision-mediated genome instability. (A) DRIP with the S9.6 antibody in a wild-type strain carrying the recombined head-on [IN] and codirectional [OUT] plasmid. Samples from asynchronous cultures grown in galactose were treated (+) or not (−) in vitro with RNase H (RNH). The regions assayed were the LEU2 gene of the plasmid and the chromosomal PDR5 and PRE1 (negative control) genes. n > 3. (B) DRIP assay in wild-type strain with the GAL::LEU2 [URA3] system integrated in chromosome III (Chr III) close to ARS315 in head-on orientation. Nucleic acids from cells grown in either glucose (Trx−) or galactose (Trx+) were treated with RNase H (RNH) prior to the immunoprecipitation with S9.6 antibody. Samples from cells grown in galactose (Trx+)–overexpressing [GAL::YRA1Δi] YRA1 are shown in green. The regions assayed were the LEU2 gene at the head-on OUT integrated system and the chromosomal PDR5 gene. n > 3. (C) ChiP analysis of H2A-P in the GAL::LEU2 [URA3] construct integrated in chromosome III (Chr III) in head-on orientation with respect to replication. Cells transformed with GAL::RNH1 [RNH−] or with the empty vector [RNH−] were grown in glucose (Trx−) or galactose (Trx+). n > 3. Cells overexpressing [GAL::YRA1Δi] YRA1 are shown in green. The regions assayed were the LEU2 gene at the head-on IN integrated system and the telomere (TELVI-R). n > 3. Means and SEM are plotted in all panels. Asterisks indicate statistically significant differences between the strains indicated, according to Student’s t-tests. [*] P < 0.05; [**] P < 0.005; ['**'] P < 0.0005.
expression on senescence and telomere dynamics in significance (Supplemental Fig. S5).

We noticed an increase of Y amplification upon Yra1 overexpression, although this increase did not reach a statistical significance (Supplemental Fig. S5). These results unequivocally indicate that it is the high accumulation of overexpressed Yra1 at telomeres (Gavaldá et al. 2016) rather than an inhibition of telomerase activity due to the sequestering of the RNA template that causes a quick and profound shortening of telomeres and premature senescence. We also noticed an increase of Y amplification upon Yra1 overexpression, although this increase did not reach a statistical significance (Supplemental Fig. S5).

Finally, we tried to determine the impact of RNH1 overexpression on senescence and telomere dynamics in est2Δ cells overexpressing Yra1. However, RNase H overexpression leads to a very sick phenotype of telomerase-deficient est2Δ cells overexpressing Yra1. These cells entered crisis quickly with a very low viability compared with est2Δ cells overexpressing only Yra1. Unfortunately, it is not possible to know whether this behavior is the result of two independent types of stresses (RNase H and Yra1 overexpression) or whether the two processes are interdependent, and any interpretation about the type of survivors was hampered by the very low cell density at crisis in cells co-overexpressing Rhn1 and Yra1. Indeed, since survivors result from clonal events, the large heterogeneity might simply reflect differences among their very low cell densities. Therefore, we cannot conclude whether RNase H overexpression counteracts Yra1 overexpression or potentiates its effect on telomere dynamics, both possibilities being consistent in any case with the increased presence of RNA–DNA hybrids at telomeres (Fig. 6E).

**Figure 6.** Premature senescence and accelerated shortening of telomeres promoted by Yra1 overexpression. (A) Mean growth curves over time of wild type (n = 2), GAL::Yra1Δi (n = 2), est2Δ (n = 10), and est2Δ GAL::Yra1Δi (n = 10). Each clone corresponds to a spore isolated from the heterozygous diploid strain, propagated in liquid culture through daily serial dilutions every 2 d. OD600 was measured every 2 d to estimate the cell density. Population doublings (PDs) were estimated from the initial spores. (B, top) Schematic representation of wild-type, type I, and type II telomeres. All telomeres contain one X element in the subtelomeric region and from zero to four long [L] or short [S] Y subtelomeric sequences. Type I survivors show an amplification of Y sequence and interstitial TG1 repeats. Type II survivors display an elongation of TG1 terminal repeats. Positions of the Xhol sites are shown. (C) TG1Δ-probed Southern blot analysis of the telomere length of representative clones of est2Δ and est2Δ GAL::Yra1Δi during senescence. (D) Southern blot analysis and mean telomere length at the first time point. n = 10. (E) DRIP with the S9.6 antibody in wild-type and est2Δ strains. Nucleic acids were treated with RNase H (RNH+) prior to the immunoprecipitation with S9.6 antibody. Samples from cells grown in galactose (Trx+) overexpressing (GAL::YRA1Δi) mean growth curves over time of wild type (n = 2), GAL::Yra1Δi (n = 2), est2Δ (n = 10), and est2Δ GAL::Yra1Δi (n = 10). Each clone corresponds to a spore isolated from the heterozygous diploid strain, propagated in liquid culture through daily serial dilutions every 2 d. OD600 was measured every 2 d to estimate the cell density. Population doublings (PDs) were estimated from the initial spores. (B, top) Schematic representation of wild-type, type I, and type II telomeres. All telomeres contain one X element in the subtelomeric region and from zero to four long [L] or short [S] Y subtelomeric sequences. Type I survivors show an amplification of Y sequence and interstitial TG1 repeats. Type II survivors display an elongation of TG1 terminal repeats. Positions of the Xhol sites are shown. (C) TG1Δ-probed Southern blot analysis of the telomere length of representative clones of est2Δ and est2Δ GAL::Yra1Δi during senescence. (D) Southern blot analysis and mean telomere length at the first time point. n = 10. (E) DRIP with the S9.6 antibody in wild-type and est2Δ strains. Nucleic acids were treated with RNase H (RNH+) prior to the immunoprecipitation with S9.6 antibody. Samples from cells grown in galactose (Trx+) overexpressing (GAL::YRA1Δi) YRA1 are shown in green. The region assayed was the telomere (TELVI-R). n = 3. Means and SEM are plotted in all panels. Asterisks indicate statistically significant differences between the samples indicated, according to Student’s t-tests. [*] P < 0.05; [**] P < 0.005; [***] P < 0.0005.

**Discussion**

After demonstrating that the RNA-binding factor Yra1 can bind RNA–DNA hybrids in vitro and in vivo, we showed that dynamic DNA–RNA hybrids are bound by overexpressed Yra1 and threaten genome integrity by promoting transcription–replication collisions. This is a general effect occurring all over the genome. Hyperrecombination caused by transcription–replication collisions is R-loop-dependent and specific to head-on orientation, not being observed for codirectional transcription. Importantly, overexpression of Yra1 causes a similar and significant sevenfold to eightfold increase in R-loop-dependent recombination in systems undergoing either head-on or codirectional transcription–replication collisions. This suggests that RNA–DNA hybrids are cotranscriptionally formed independently of replication, but only when the replication fork approaches an R loop from downstream from the RNAp does it compromise genome integrity. Thus, the replication fork progressing in the same direction as transcription may easily resolve an R loop, unless Yra1 is bound to it. The ability of Yra1 to bind R loops and block replication dynamics is supported by the fact that Yra1 overexpression shortens telomeres by a telomerase-
helicase with in vitro RNA R loops. Indeed, Yra1 inhibits loading of Dbp2, an RNA leading to the accumulation of stable or more persistent likely impeding the action of helicases and RNase H and (Ma et al. 2013, 2016). However, the in vitro ssDNA-bind-

interestingly, be its role when present at wild-type levels in the cell. In- and in vivo when overexpressed, although this would not have consequence on genome integrity. However, here we showed that R loops by themselves may not be strong inducers of genome instability; instead, they rely on a subsequent step to compromise genome integrity (García-Pichardo et al. 2017). Thus, naturally formed R loops are presumably transient, with no consequence on genome integrity. However, we demonstrate that by artificially stabilizing transient R loops by binding to overexpressed Yra1, R loops accumulate at high levels and compromise genome integrity.

We showed previously that transcription induces hyperrecombination only when it occurs concomitantly with replication and in head-on orientation [Prado and Aguilera 2005]. Now we show that the hyperrecombination induced in the head-on orientation is largely dependent on R loops [Fig. 4] but not codirectional collisions. These results are consistent with two recent reports in bacteria and human cells showing that codirectional transcription–replication conflicts have little impact on the stability of the construct-containing plasmid or on bacteria viability, while head-on orientation is unstable and causes

![Figure 7. Analysis of type II recombination events at TELVI-R.](image)

**A** The clones used in Figure 6 were analyzed by single-telomere Southern blot at the first time point of senescence and after the appearance of survivors. Each band corresponds to a single type II recombination event in the cell population independently of its intensity. **B** The frequency of type II recombination events at TELVI-R. Details are as in Figure 6D.

independent mechanism presumably linked to R loops generated by TERRA RNA, which is yet to be understood. These data indicate that R loops form transiently in the cell regardless of replication and generate genome instability mainly in head-on orientation or when they are stabilized by proteins such as Yra1 by impeding the progression of replication.

We showed here that the Yra1 RNA-binding factor can bind and presumably stabilize RNA–DNA hybrids in vitro and in vivo when overexpressed, although this would not be its role when present at wild-type levels in the cell. Interestingly, *Arabidopsis* AtNDX, a homeodomain-containing protein, stabilizes R loops by binding to the displaced ssDNA [Sun et al. 2013]. In contrast to AtNDX, Yra1 is an RNA-binding protein that presumably affects R-loop dynamics by binding to the RNA–DNA hybrids, likely impeding the action of helicases and RNase H and leading to the accumulation of stable or more persistent R loops. Indeed, Yra1 inhibits loading of Dbp2, an RNA helicase with in vitro RNA–DNA-unwinding activity (Ma et al. 2013, 2016). However, the in vitro ssDNA-bind-

How transcription induces genome instability is not fully understood yet, but research conducted in the last two decades points to conflicts with replication as a major source of transcription-associated instability [García-Rubio et al. 2003; Prado and Aguilera 2005; Mirkin and Mirkin 2007; Boubakri et al. 2010]. The transcription-dependent recombination results obtained in the codirec-

Figure 7. Analysis of type II recombination events at TELVI-R. **A** The clones used in Figure 6 were analyzed by single-telomere Southern blot at the first time point of senescence and after the appearance of survivors. Each band corresponds to a single type II recombination event in the cell population independently of its intensity. **B** The frequency of type II recombination events at TELVI-R. Details are as in Figure 6D.
lethality in RNase H-deficient bacteria (Hamperl et al. 2017; Lang et al. 2017). Nevertheless, the S9.6 DRIP analysis reveals that RNA–DNA hybrids are formed in LEU12 in not only the head-on but also the codirectional system, although they do not largely compromise the genome integrity. We detected R loops regardless of transcription–replication orientation, but when Yra1 is overexpressed, an R-loop-dependent increase in recombination occurred in both systems, whether head-on or codirectional, demonstrating that they are present in both situations. Therefore, our data indicate that R loops are formed cotranscriptionally regardless of replication and may be present before the replication fork arrives; that is, it is not the transcription–replication collisions the cause of the R loops (Bermejo et al. 2011; Hamperl et al. 2017; Lang et al. 2017). Consistently, R loops are also observed in the codirectional orientation in a bacterial chromosome too (Lang et al. 2017) and are detected in G1 cells in yeast and human cells (Castellano-Pozo et al. 2012; Bhatia et al. 2014). Therefore, we conclude that R loops are the cause and not the consequence of transcription–replication collisions, consistent with the genome-wide accumulation of RNA–DNA hybrids detected in our study (Fig. 2; Supplemental Fig. S1) or other previously published reports (Chan et al. 2014; El Hage et al. 2014; Wahba et al. 2016).

We propose that the different outcome of a replisome encountering an R loop codirectionally or head-on may depend on the structure encountered. In codirectional orientation, the replisome would meet the RNA–DNA hybrid directly in the leading strand template, presumably being able to remove it easily with the help of some helicase function (Fig. 8A). However, in the head-on orientation, the fork would encounter first the RNA polymerase that thus would be trapped ahead of the nascent RNA forming the RNA–DNA hybrid in the lagging strand template. In this orientation, the RNA polymerase would be a major contributor to the stalling of the replisome, presumably incapable of backtracking due to the hybrid behind (Fig. 8A). Indeed, it has been shown in bacteria that backtracked RNA polymerase unable to resume transcription would cause DNA breaks when encountered by a replication fork (Nudler 2012). It is worth noting that a physical contact between the replisome and the RNA polymerase might not be necessary, as the positive topological constraint generated in between the two advancing polymerases may be sufficient to block both processes. However, if the R loop is blocked artificially by binding with Yra1 when it is in excess in the cell, then it becomes an obstacle to replication no matter from which direction it is encountered (Fig. 8B).

The importance of R-loop stabilization becoming a general obstacle to replication is supported by the analysis of the impact of Yra1 on telomere dynamics and homeostasis, where R loops have been shown to play a role. We show here that Yra1 overexpression causes a telomere shortening phenotype that is not explained by a potential ability of overexpressed Yra1 to sequester the telomerase TLC RNA. Telomeres are shortened in the absence of telomerase when Yra1 is overexpressed (Fig. 6), favoring the idea that Yra1 excess would be shortening the telomere through R-loop stabilization. Telomere repeats and subtelomeric regions are transcribed in IncRNAs known as TERRA, which form unstable R loops (Maicher et al. 2014; Azzalin and Lingner 2015) that need to be removed before chromosomal ends are replicated (Graf et al. 2017). Therefore, it is likely that, by stabilizing TERRA R loops, Yra1 overexpression impairs telomere replication progression. Consistent with this idea, overexpressed Yra1 is recruited to telomeress, and Rrm3, the replicative helicase that resolves replication fork stalls, accumulates at telomeres in this situation (Gavalda et al. 2016). Importantly, the presence of RNA–DNA hybrids at telomeres is higher in est2Δ cells, as expected, and significantly enhanced under Yra1 overexpression (Fig. 6E). Fork collapse mediated by stable R loops might be responsible of the telomere shortening, thus causing premature senescence (Simon et al. 2016). Indeed, deletion of RNase H or the THO complex in cells unable to undergo recombination (rad52Δ) increases TERRA R loops and leads to rapid telomere loss and senescence (Balk et al. 2013). We observed a reduced appearance of type II telomeres and slightly higher Y subtelomeric amplification in the survivors after senescence. This suggests that Yra1 overexpression may favor type I recombination between subtelomeric Y elements at the expense of type II recombination. This is consistent with the observation that RNA–DNA hybrids formed by

**Figure 8.** Model to explain Yra1-mediated transcription–replication collisions. [A] In the codirectional orientation, the replication fork [RF] encounters the R loop, removing it. In the head-on orientation, either the positive supercoiling accumulated ahead or the RNA polymerase itself would block the replication fork. [B] Overexpressed Yra1 (+Yra1 OE) binds to the R loop, stabilizing it and blocking replication whether it proceeds codirectionally or in head-on orientation.
In conclusion, we uncovered the role of RNA–DNA hybrids in transcription–replication conflicts regardless of the orientation, indicating that R-loop formation is independent of replication. Artificially stabilized RNA–DNA hybrids cause genome instability when encountered by the replication fork in both head-on and codirectional orientation; otherwise, only RNA–DNA hybrids involved in head-on transcription–replication collisions have a major effect on genome instability. These findings may help understand the role in preventing R-loop accumulation and R-loop-mediated genome instability of specific functions such as histone deacetylase Sin3A, the THO complex, or the Fanconi anemia pathway, which have been shown to impair replication fork progression through R loops when mutated (García-Rubio et al. 2015; Schwab et al. 2015; Madireddy et al. 2016; Salas-Armenteros et al. 2017). Similarly, these results raise the question of whether overabundance of the Yra1 ortholog ALY protein in a significant proportion of tumoral cells may be related to high levels of stabilized RNA–DNA hybrids that contribute to genome instability (Dominguez-Sanchez et al. 2011). This work thus should open new perspectives to understand the mechanisms of R-loop-mediated genome instability and its implications in cancer development.

Materials and methods

Strains, primers, and plasmids

Yeast strains, primers, and plasmids used are listed in Supplemental Tables S1–S3, respectively. The BGL2 strain was generated by integrating in BY4741 the GAL1::LEU2 (URA3) construct at position 225,807 of chromosome III in a head-on orientation relative to replication starting at ARS315. This strain was crossed with Ybp250 to generate BYGL2-10B. The recombinant plasmid pARSGLBIN-Leu2 was used as a template to amplify the GAL1::LEU2 (URA3) construct, adding the BglII restriction site and 40-bp of homology to the region upstream of or downstream from position 225,807 of chromosome III at both sides. The linear PCR fragment was purified and transformed in the BY4741 strain. Ura+ transformants were checked for GAL1::LEU2 (URA3) integration by Southern blot.

Genome-wide experiments

DRIP was performed as described (García-Pichardo et al. 2017). For sequencing, immunoprecipitation and input samples were used as templates with a GenomePlex Complete whole-genome amplification kit [Merck] to amplify the DNA. The library was prepared to obtain a fragment size of 200 bp according to the manufacturer’s protocol [Ion Torrent PGM]. The quality of the samples was certified with a Bionalyzer and a high-sensitivity DNA analysis kit, and runs were performed in an Ion 316 CHiP version 2. The data generated are available under Gene Expression Omnibus accession number GSE113580. Bioinformatic analysis was performed as described [see the Supplemental Material]; DRIP-seq signal values were multiplied by three for visualization.

Yra1 expression and purification

His6-tagged Yra1 was expressed from pET-Yra1 [MacKellar and Greenleaf 2011] in BL21 Rossetta Escherichia coli DE3 cells [Novagen]. Bacteria were grown in 1 L of LB medium with ampicillin and chloramphenicol, and protein expression was induced with 0.2 mM IPTG overnight at 16°C. Cells were lysed and His6–Yra1 purified through Ni-Sepharose Fast Flow resin [GE Healthcare] as described [MacKellar and Greenleaf 2011] followed by SP-Sepharose [GE Healthcare] [Ma et al. 2013].

Electrophoretic mobility shift assay

RNA and DNA oligonucleotides purchased from Integrated DNA Technologies [IDT] were end-labeled with[32]P-ATP using T4 poly-nucleotide kinase according to the manufacturer’s instructions. The same nucleic acids were used as cold competitors and added at the concentration indicated. The binding reactions were performed with the indicated nucleic acids at 30–50 nM final concentration and increasing amounts of recombinant Yra1 (170 nM to 1.4 mM) in binding buffer (10 mM Tris- HCl at pH 7.5, 25 mM NaCl, 1 mM DTT, 10% glycerol) supplemented with 0.05 mg/mL BSA, 1× cocktail protease inhibitors, and RNase OUT for 10 min at 30°C. Samples were resolved in 6% PAGE and 0.5× TBE run for 1 h at 15 mA. Gels were dried, and images were taken with a Fujifilm Life Science FLA-5100 imaging system.

DRIP assays

DRIP in est2Δ strains was performed as described [Garcia-Rubio et al. 2018] in haploid spore products of diploid PAY316 obtained by crossing PAY76 with PAY321 that were heterozygous for EST2 [EST2/est2Δ] and carried the pRS415GAL or pRS413GAL::YRA1Δ plasmid. After 3 d of growth at 30°C, all of the spore colonies were transferred to 10 mL of liquid SRaf.

ChiP assays

Yeast mid-log cultures growing at 30°C were processed as described [Hecht and Grunstein 1999] with minor modifications. Cells were broken in a multibead shaker at 4°C in lysis buffer (50 mM HEPES-KOH at pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate). Chromatin was sonicated to an average fragment size of 400–500 bp and immunoprecipitated with 5 µL of anti-histone H2A [phospho-129] or anti-HA antibody coated to Protein A Dynabeads [Invitrogen].

Genome instability analysis in Saccharomyces cerevisiae

Spontaneous recombination frequencies were obtained as the average value of median frequencies obtained by six to 10 fluctuation tests performed with two to three independent transformants. For each fluctuation test, six independent colonies were analyzed as described previously [Prado and Aguilera 2005].
Senescence assays and telomere analysis

Senescence assays in liquid medium were performed as described previously (Churikov et al. 2014) from haploid spore products of diploid, PAY247 or PAY249 obtained by crossing PAY267 or PAY269 with PAY76 that were heterozygous for EST2 [EST2Δ] and carried the pEST2-URA3 plasmid and the pRS413GAL or pRS413GAL::YRA1Δi plasmid. To ensure homogeneous telomere length before sporulation, the diploids were propagated for at least 50 generations on YPD plates. After 3 d of growth at 30°C, all of the spore colonies were transferred to 2 mL of liquid SGal-his to estimate the number of population doublings, and the suspensions were immediately diluted to 10^5 cells per milliliter. Cells were serially passaged in 15 mL of liquid SGal-his at 10^5 cells per milliliter at 48-h intervals. Replicative senescence was calculated as the average of two to 10 independent spores with identical genotype. Telomere analysis of the samples was performed as described (Churikov et al. 2014, Hardy et al. 2014).

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Author contributions: M.G.-R., P.A. and A.G.R. performed all experiments, with the exception of the DRIP-seq and its bioinformatic analysis, which were performed by J.L.-B., and the construction of the head-on transcription–replication system, which was performed by I.F.R. The study was designed by M.G.-R., P.A., A.G.R., M.-N.S., V.G., and A.A. who also analyzed the data. A.G.-R. and A.A. wrote the first draft of the manuscript. All authors contributed to the final version.

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