Timeless couples G quadruplex detection with processing by DDX11 during DNA replication

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Regions of the genome with the potential to form secondary structure pose a frequent and significant impediment to DNA replication and must be actively managed in order to preserve genetic and epigenetic integrity. The fork protection complex (FPC), a conserved group of replisome-associated proteins including Timeless, Tipin, and Claspin, plays an important role in maintaining efficient replisome activation, ensuring optimum fork rates, sister chromatid cohesion and checkpoint function. It also helps maintain the stability of sequences prone to secondary structure formation through an incompletely understood mechanism. Here, we report a previously unappreciated DNA binding domain in the C-terminus of Timeless, which exhibits specific binding to G quadruplex (G4) structures. We show that, *in vivo*, both the C-terminus of Timeless and the DDX11 helicase act collaboratively to ensure processive replication of G4 structures to prevent genetic and epigenetic instability.

DNA can create significant impediments to its own replication through formation of secondary structures. When unwound, certain sequences, often repetitive or of low complexity, can adopt a variety of non-B form structures, including hairpins, cruciforms, triplexes and quadruplexes ¹. It is becoming clear that secondary structure formation is a frequent event during replication, even at genomically abundant sequences previously thought not to be a major source of difficulty ². To prevent such sequences causing havoc with the genetic and epigenetic stability of the genome, cells deploy an intricate network of activities to counteract secondary structure formation and limit its effects. These activities include proteins that bind and destabilise DNA structures and specialised helicases that unwind them ³. In addition, the repriming activity of PrimPol can be deployed to confine a structure into a minimal region of single stranded DNA, limiting the potential dangers of exposing extensive ssDNA in a stalled replisome ^{2,4}.

G quadruplexes (G4s) are one of the most intensively studied and potent structural replication impediments. G4s arise in consequence of the ability of guanine to form Hoogsteen base-paired quartets ⁵. In favourable sequence contexts, comprising runs of dG separated by variable numbers of non-G bases, stacks of G quartets form G4s secondary structures. Current estimates suggest that over 700,000 sites in the human genome have the potential to form G4s ⁶. While some of these G4s may have important roles in genome physiology, all pose a potential threat to DNA replication and sites with G4-forming potential have been linked to both genetic and epigenetic instability ^{7,8}.

Precisely how DNA structures are detected and resolved by the replication machinery remains unclear. Many of the factors involved in processing G4 secondary structures, for instance FANCJ and REV1 9-13, do not appear to be constitutive components of the replisome 14. It is thus likely that core components of the replisome will act as 'first responders' to DNA structures and play an important role coupling their detection with suppressing their deleterious effects on DNA synthesis. Particularly interesting in this context is a subset of replisome components known as the fork protection complex (FPC). The FPC comprises four main proteins, Timeless, Tipin, Claspin and And-1 and is conserved from yeast to mammals ¹⁵. FPC components associate with the replication fork via direct interactions with the CMG replicative helicase and replicative polymerases α , δ and ε^{16-19} . They also interact with DNA both directly ²⁰ and indirectly via replication protein A ²¹. These interactions allow the FPC to remain at the fork, which ensures a normal speed of DNA synthesis ²². Additionally, the FPC has a series of functions that ensure normal replisome function and fork integrity: it is essential to avoid uncoupling of pol ε from the replicative helicase and consequent formation of long stretches of ssDNA ^{23,24}. It also has a conserved role in S-phase checkpoint activation in response to DNA damage, including checkpoint kinase activation, cell cycle arrest and maintenance of the integrity of the replication fork ^{25–28}. It also plays an important, but incompletely understood, role in maintaining chromosome cohesion ^{29,30}.

The FPC is thus well placed to play a role in the detection and metabolism of DNA secondary structures that could impede DNA synthesis. Indeed, deficiency of both Tof1 in yeast and Timeless in human cells leads to replication fork stalling, repeat instability and fragility at secondary structure-forming sequences ^{31–35}, underscoring the potential importance of Timeless in maintaining processive replication through regions of the genome capable of forming secondary structures.

Although Timeless itself does not appear to possess a catalytic activity that would process DNA secondary structures, it interacts with the helicase DDX11 ³⁶. DDX11 (or CHLR1) is a 5'-3' Fe-S helicase and belongs to the same superfamily 2 as FANCJ, RTEL and XPD ³. In humans, mutations in DDX11 cause Warsaw Breakage Syndrome, an extremely rare autosomal recessive disease characterised by microcephaly, growth retardation, cochlear abnormalities and abnormal skin pigmentation ³⁷. *In vitro*, DDX11 has unwinding activity on several non-duplex DNA structures, such as G4s ^{38,39}, triplex DNA ⁴⁰ and D-loops ³⁹. Further, the helicase activity of DDX11 is enhanced by Timeless ³⁶. However, it remains unclear how Timeless and

DDX11 collaborate *in vivo* in detecting and processing G4s during replication. Here, we provide *in vivo* evidence that Timeless and DDX11 operate together to ensure processive replication of G4-forming DNA. We report a previously unappreciated DNA binding domain in the C-terminus of Timeless, which exhibits specificity towards G4 structures. We propose that the C-terminus of Timeless plays a role in the detection of G4 structures at the replication fork, recruiting DDX11 to unwind them to ensure processive replication is maintained, thereby avoiding G4-induced genetic and epigenetic stability.

Results

Timeless is required for processive replication of a genomic G4-motif

To address whether Timeless is involved in maintaining processive replication of G4-forming DNA in vivo, we disrupted the TIMELESS locus in chicken DT40 cells with CRISPR/Cas9induced deletions. We isolated several timeless mutants with biallelic disruptions in exon 1, around the guide site (Supplementary Fig. 1). The timeless mutant cells were sensitive to cisplatin (Supplementary Fig. 2), as previously observed in human cells depleted of Timeless ⁴¹. To assess the role of Timeless in the replication of a G4-forming sequence, we took advantage of the Bu-1 loss variant assay 42. The stable expression of the BU-1 locus in DT40 is dependent on the maintenance of processive replication through a G4 motif located approximately 3.5 kb downstream of the promoter (Fig. 1a). Prolonged pausing of leading strand replication at this motif leads to loss of epigenetic information around the promoter of the gene and a permanent and heritable change in its expression 4,42-44. This stochastic and replication-dependent generation of Bu-1 loss variants can be monitored by flow cytometry as BU-1 encodes a surface glycoprotein. Small pools of Bu-1^{high} cells of wild type and timeless mutants were expanded in parallel for ~ 20 divisions (15 – 21 days) and the proportion of cells in each pool that had lost their Bu-1^{high} status determined. We detected increased levels of Bu-1 expression instability in *timeless* DT40 cells compared to the wild-type cells, which retained their stable Bu-1^{high} expression (Fig. 1b & c). The instability of Bu-1 expression in *timeless* cells was fully complemented by expression of human Timeless (Fig. 1c) and is dependent on the +3.5 G4 motif (Fig. 1c). Cells deficient in Tipin ⁴⁵, a constitutive interactor of Timeless within the FPC, also exhibit instability of BU-1 expression (Fig. 1d). These results show that Timeless is necessary to maintain processive DNA replication of a genomic G4-motif.

Identification and characterisation of a Timeless DNA-binding domain

As a core component of the replisome, Timeless is intimately associated with DNA synthesis at the replication fork ²². *In vitro* data show that the Timeless-Tipin complex can bind to single-stranded DNA (ssDNA) through RPA ²¹ and the Swi1-Swi3 complex, the fission yeast orthologue of Timeless-Tipin, was also shown to bind DNA ²⁰. Inspection of the amino acid sequence of human Timeless revealed the presence of a conserved domain in its C-terminal half (residue 816 to 954) (Fig. 2a), with a predicted fold similarity to the myb-like proteins of the homeodomain-like superfamily, that bind double-stranded DNA (dsDNA) with a tandem repeat of 3-helix bundles (named here N-term and C-term).

We used X-ray diffraction and NMR spectroscopy to investigate experimentally the structure and dynamics of the newly discovered Timeless domain. The 1.15 Å crystal structure of amino acids 885 to 947 (C-term), corresponding to a single myb-like fold, confirmed the presence of a three-helix bundle characteristic of the homeodomain superfamily of DNA-binding proteins, extended by the presence of a fourth C-terminal alpha helix unique to the Timeless domain (Fig. 2b). The NMR structural ensemble of amino acids 816 to 954 revealed two well-converged domains (824-880 and 891-944, backbone r.m.s.d. 0.4 and 0.5 Å, respectively) connected by a linker (881-890) that was significantly less well converged, implying a high degree of flexibility between N- and C-term domains (Fig. 2c). This observation was confirmed by backbone dynamics measurements (Supplementary Fig. 3). The N- and C-terminal domains adopted the same three-dimensional fold; in particular, the N-terminal repeat shared the presence of an additional fourth helix, H4, as seen in the C-terminal repeat (Fig. 2c).

In keeping with the similarity of its structure to known DNA-binding domains, we examined the ability of the Timeless domain to interact with DNA. We found that it bound with low micromolar affinity to both ss- and ds-DNA probes (Fig. 2d; top panel). A distinguishing feature of the Timeless DNA-Binding Domain (DBD) is the presence of a fourth alpha helix in both N- and C-terminal 3-helix bundle repeats. Superposition of the DBD C-term onto the structurally homologous domains of the telomeric protein TRF1 ⁴⁶ and bacterial cell-cycle regulator GcrA ⁴⁷ in complex with their double-stranded DNA substrates (Fig. 2e) shows that, if the DBD were to adopt a similar model of dsDNA binding, the fourth helix of both its N-term and C-term repeats would likely lead to a steric clash with the phosphate backbone of the DNA.

Although it is conceivable that the DBD might rearrange its conformation upon DNA binding, the fourth helix of both repeats participates in core hydrophobic interactions, mediated by conserved residues L872 and F879 (N-terminal repeat), and L936, V937 and L943 (C-terminal repeat), making such rearrangements unlikely. Given the functional context in which Timeless operates as a replisome component and the observations, presented in Fig. 1, that it is required to maintain processive replication of the *BU-1* G4, we speculated that the C-terminal DNA-binding activity of Timeless might be directed towards recognition of DNA secondary structures that form transiently on the unwound template, such as G4s.

Indeed, when we tested a well-characterised G4 sequence present in the promoter of the MYC gene ⁴⁸, the DBD bound to it with nanomolar affinity (Fig. 2d, bottom panel), and about one order of magnitude tighter than ds- or ssDNA. This observation prompted us to ask whether the Timeless-Tipin complex, like the isolated DBD, is able to bind to the same G4 motif. To mimic the unwound template DNA, we embedded the G4 within a longer ssDNA sequence (ssG4) (Supplementary Table 2). We found that the Timeless-Tipin complex bound to ssG4 with low micromolar affinity and in a selective fashion, as it did not show measurable interactions with a hairpin DNA embedded within the same ssDNA (ssHP), or a mutated ssG4 sequence that had lost the ability to fold into a guanine quadruplex (ss) (Fig. 3a, Supplementary Fig. 4, Supplementary Table 2). We next tested a series of G4 sequences, found in different genomic contests and with different folding topologies: we found that the Timeless-Tipin complex bound to all of them, albeit with different affinities that varied several fold, whereas it did not show appreciable binding to ss- or dsDNA (Fig. 3b).

These findings show that Timeless contains, in its C-terminal half, a previously unrecognised DNA-binding domain, which closely resembles in structure the tandem repeat of three-helix bundles found in the homeodomain-like superfamily of transcription factors. While Timeless DBD binds to both ss- and dsDNA, it binds with ~10-fold greater affinity to a defined G4 DNA sequence. The preference for G4 DNA is retained by the Timeless-Tipin complex.

The Timeless C-terminus is crucial for processive G4 replication in vivo

To further explore the *in vivo* contribution of Timeless C-terminus to G4 replication, we generated a DT40 cell line expressing a version of Timeless truncating the gene before the DBD, using CRISPR-Cas9 gene targeting to exon 16. This truncation also removes a domain previously reported to bind PARP1, the PARP1-binding domain (PBD) ⁴⁹. This cell line

exhibits instability of BU-I expression comparable to the *timeless* mutant (Fig. 4) suggesting a role for the C terminus of the protein in G4 replication. To further dissect this observation, we complemented *timeless* cells with truncated versions of human Timeless, lacking only the DBD (Δ DBD) or the PBD (PARP*). Timeless lacking each of these domains individually largely, but not completely, restores the BU-I expression instability of the *timeless* mutant suggesting that they act redundantly. Additionally, co-immunoprecipitation experiments show that neither region is required for binding the DDX11 helicase (Supplementary Fig. 5), a known binding partner of Timeless and whose ability to unwind G4s is stimulated by Timeless 36 . These results indicate that the C terminus of the Timeless protein has an important role in G4 replication with the DBD and the PDB both contributing to that function independently of DDX11 recruitment.

DDX11 ensures processive G4 replication in vivo

Since Timeless itself lacks any catalytic activity, we next explored the extent to which the genetic interaction between Timeless and the DDX11 helicase accounted for G4 processing in this system. As noted above, Timeless interacts with DDX11 29,36,50 , and this interaction has been shown to be important for sister chromatid cohesion 50 and preservation of fork progression in perturbed conditions 36 . Further, *in vitro*, DDX11 can unwind several DNA structures including G4s $^{38-40}$ and the activity of DDX11 is stimulated by Timeless 36 . Recent work has shown that DDX11 interacts with Timeless through a domain in exon 4 50 . We therefore examined BU-1 stability both in a ddx11 mutant generated by conventional gene targeting, which disrupts exons 7-12 downstream of the Timeless interaction domain 45 and in CRISPR/Cas9-generated deletion mutants that targets exon 4 and introduces downstream frameshifts disrupting expression of the Timeless interaction domain (Supplementary Fig. 1). The newly generated CRISPR/Cas9-generated ddx11 mutants are sensitive to cisplatin (Supplementary Fig. 2), similar to the previously reported mutant generated by gene targeting

All *ddx11* mutants examined exhibited elevated rates of Bu-1 loss variant generation (Fig. 5a). This expression instability was supressed by complementation with full-length chicken DDX11, but not with the K87A helicase-dead form of DDX11 (the chicken equivalent of the human K50A mutation in the Walker A motif), demonstrating that the helicase activity of DDX11 is essential for G4 replication *in vivo* (Fig. 5b). Importantly, the *BU-1* expression instability of *ddx11* cells is also dependent on the presence of the +3.5 G4 in *BU-1* (Fig. 5b).

DDX11 works with Timeless in preventing G4-dependent instability of BU-1 expression. We next asked whether DDX11 and Timeless participate in the same pathway for G4 resolution at the replication fork. We generated ddx11/timeless double mutants by targeting Timeless in the CRISPR/Cas9-generated ddx11 clone 1 cells. Fluctuation analysis showed that the double-mutant ddx11/timeless does not exhibit higher rates of Bu-1 loss-variant formation, compared to the single Timeless or ddx11 mutants (Fig. 5c). This suggests that these proteins act in the same pathway for G4 replication. This finding is in contrast to the genetic interaction between DDX11 and the FANCJ helicase, which also plays a prominent role in replication of G4s $^{9-12}$. fancj DT40 exhibit instability of BU-1 expression 44 that is dependent on the helicase activity of FANCJ (Supplementary Fig. 6). A double fancj/ddx11 mutant 51 exhibits significantly more BU-1 expression instability than either single mutant (Fig. 5d), suggesting that these helicases operate on the BU-1 G4 motif independently of each other.

Timeless and DDX11 deficiency leads to decreased proliferation and increased activation of DDR in presence of pyridostatin (PDS)

We next asked whether deficiency in Timeless and DDX11 exacerbates the globally detrimental cellular consequences of G4-binding ligands. We treated *timeless*, ddx11, and ddx11/timeless cells with the G4 ligand pyridostatin (PDS) ⁵². All mutant cell lines, but not the wild type, exhibited an intrinsic reduction in doubling time with cellular proliferation further decreased in the presence the G4 ligand (Fig. 6a). We next looked at the induction of histone H2AX phosphorylation (γ -H2AX) using permeabilised cell flow cytometry. We have previously observed that PDS induces little or no γ -H2AX in wild type DT40 cells ⁴³ and this was also true in our cytometric assay. However, *timeless*, and ddx11 mutants both exhibit increased levels of γ -H2AX (detected by flow cytometry) following treatment with 4 μ M PDS for 3 days compared with wild type cells (Fig. 6b & c). γ -H2AX levels in the double ddx11/timeless mutant after PDS treatment were not significantly higher than the *timeless* single mutant, again supporting the observation that these the two proteins operate in the same pathway for avoiding G4 ligand-induced DNA damage signalling.

Deficiency of DDX11 and Timeless leads to dysregulation of a common set of genes harbouring G4s in close proximity to their TSS

We have previously reported that the instability of expression of the BU-1 locus in mutants defective in G4 replication is observed in other loci genes across the genome 44,53 . We have

shown that correlations between the identity of the genes dysregulated in different mutants, along with the direction and magnitude of those changes, can be used to infer genetic relationships between G4 processing enzymes 44,53. We applied this approach to the relationship between Timeless and DDX11, assessing gene expression changes with RNA-seq. Loss of DDX11 or Timeless induced marked changes in gene expression (timeless 1752; ddx11 821 genes with altered expression at p > 0.95), with an approximately equal number of genes being up- or down-regulated (Supplementary Fig. 7a) in both cases. Moreover, there was a significant (p $< 2.2 \times 10^{-16}$) overlap in the identity of the deregulated genes in *timeless* and ddx11 cells (Supplementary Fig. 7b) and a significant correlation (Spearman correlation 0.82, p value $< 10^{-16}$) both in the magnitude and direction of the change in expression at the level of individual genes (Fig. 7a). Consistent with our model ⁴², genes dysregulated in *timeless* and ddx11 mutants are more likely to harbour a G4 close to their TSS. Using a G4 regex of (G₃N₁₂)₃G₃, we found that the region 1.5 kb around the TSS in genes dysregulated in *timeless* and ddx11 is enriched for G4 motifs relative to all genes (Fig. 7b), an observation that is not simply explained by a higher GC content (Fig. 7c). Further, the enrichment of G4s near the TSS was similar to that seen in cells lacking the FANCJ helicase (Fig. 7b & 7d) and the changes in gene expression in *timeless* and *ddx11* also correlated strongly with those observed in *fancj* cells, which we have previously studied ^{44,53} (Supplementary Fig. 7c, Spearman correlation ~ 0.9, p value $< 10^{-16}$), providing further support for a common, G4-dependent mechanism for the gene dysregulation in these three mutants. Together, these results suggest that DDX11 and Timeless collaborate to ensure expression stability through replication of a subset of genes with G4s close to their transcription start site.

Discussion

Accumulating genetic evidence suggests that DNA secondary structure formation is a frequent challenge to vertebrate DNA replication ³. Numerous factors collaborate to ensure that these structures do not result in persistent stalling of DNA synthesis or that the resulting tracts of single strand DNA formation are limited, for instance by efficient repriming ⁴. By limiting uncoupling of DNA unwinding and DNA synthesis, these mechanisms prevent both genetic and epigenetic instability.

A key to ensuring that polymerase pausing at secondary structures does not lead to extensive single stranded DNA formation should be efficient sensing of structure formation by the replisome coupled to the recruitment of factors that will remove the structure and, if needed, reprime DNA synthesis. However, it remains to be established how secondary structure formation on the DNA template is detected by the replisome.

Timeless, along with other components of the FPC, appears to be a constitutive component of the core replisome ⁵⁴. Although not essential for DNA replication, the FPC increases the speed and efficiency of fork progression and DNA synthesis ²². The intimate association of the FPC with the replisome potentially puts it in an ideal place to coordinate the response of the replisome to secondary structure formation. Indeed, loss of components of the FPC leads to instability of structure-forming repeat sequences ^{31,32,34,35,55}.

Here we show that the C-terminus of Timeless contains a DNA binding domain (DBD) that is able to bind G4 DNA more avidly than unstructured single and double stranded DNA *in vitro*. Using a sensitive *in vivo* assay that monitors episodes of replication pausing at a defined secondary structure forming sequence, we show that Timeless is required to maintain processive replication of G4 DNA. This requires the C-terminus of the protein, which contains both the newly identified DBD and an adjacent domain previously shown to bind PARP1 ^{49,56}. Interestingly, cells expressing Timeless lacking either domain exhibit significant, although not complete, complementation suggesting that there is significant redundancy between these two domains. The precise role of PARP1 in G4 sensing during replication remains to be explored, but it is noteworthy that PARP1 has been previously reported to bind G4 DNA *in vitro* ⁵⁷ suggesting, potentially, that Timeless could recognise G4s both directly through its DNA binding domain and indirectly via recruitment of PARP1 to the G4.

Timeless has also been shown to interact with the G4 helicase DDX11 and this interaction stimulates the helicase activity of DDX11 ³⁶. We have shown here that loss of Timeless and DDX11 are epistatic for G4-induced *BU-1* expression instability consistent with them operating as a unit. Interestingly, loss of both DDX11 and the related G4-unwinding Fe-S helicase, FANCJ, result in more rapid generation of Bu-1 loss variants than either single mutant suggesting that these two helicases independently contribute to replication of a the +3.5 *BU-1* G4. Our analysis of genome wide gene expression dysregulation suggests that this division of

labour between DDX11/Timeless and FANCJ is likely to be extended to other G4s in the genome.

Much work remains to be done to dissect the interactions between protein factors that process G4s and other fork-stalling DNA structures, to ensure smooth progress of DNA synthesis. Mechanistically, advances in replisome structural biology should help inform where G4 sensing by Timeless takes place relative to the CMG helicase. It is reasonable to anticipate that it will detect G4s forming transiently in the negatively supercoiled DNA emerging from CMG (Fig. 7e). The large size and multi-domain structure of Timeless indicates that its DBD could also reach and interact with DNA structures within a wide radius around the fork. This could potentially include those structures present in the DNA before it reaches CMG (Fig. 7f). Such structures might then be 'traversed' by the helicase, analogous to its proposed behaviour at a DNA-protein crosslink {Sparks et al., 2019, #91435}, before stalling the leading strand polymerase {discussed further in Lerner and Sale, 2019, #92536}. In either case, detection and resolution of the structure will ensure the smooth execution of DNA synthesis, avoiding the generation of potentially deleterious ssDNA.

Methods

Molecular cloning

Oligonucleotide sequences can be found in Supplementary Table 1. The Timeless DNA binding domain deletion (Δ DBD) and PARP-binding domain truncation mutant plasmids were constructed by PCR. pcDNA4-Flag-hTimeless was linearized with PCR using outward primers flanking the DNA binding domain (fragment 816-965) or the PARP1-binding domain (V1000-XX). 10 fmol mutant plasmids were recircularised (through the overlapping sequences) via Gibson assembly (1 h, 50°C). Recombinant DNA was treated with 10U of *DpnI* to minimize carryover of the original plasmid, and transformed into DH5 α competent bacteria. Correct mutagenesis was confirmed by restriction digest and Sanger sequencing.

FANCJ point mutations were generated by site-directed mutagenesis using a Quick Change II XL Site-Directed Mutagenesis Kit (Agilent), according to the manufacturer's instructions, in the pEAK8-CFP-hFANCJ plasmid (generated in this study). This plasmid was generated by digesting a CFP-containing plasmid, a hWT FANCJ cDNA-plasmid and a pEAK8 backbone

plasmid with *Hin*DIII and *Sal*I, *Sal*I and *Not*I and *Hin*DIII and *Not*I, respectively. All fragments were gel-purified with a QIAEX II Gel Extraction Kit (QIAGEN) and ligated in a three-way ligation reaction using a Rapid Ligation Kit (Merck) for 30 minutes at room temperature.

Cell lines and transfections

All DT40 cell lines were cultured at 37°C in a humidified 10% CO₂ atmosphere in RPMI 1640 with Glutamax (Thermo Fisher Scientific) supplemented with 7% fetal bovine serum (Thermo Fisher Scientific), 3% chicken serum (Sigma-Aldrich), 50 μM 2-mercaptoethanol and 1% penicillin/streptomycin, as previously described ⁵⁹. HEK293T cells were cultured at 37°C in a humidified 5% CO₂ atmosphere in DMEM (Thermo Fisher Scientific) supplemented with 10% fetal bovine serum and penicillin/streptomycin.

DT40 ddx11 (CRISPR), timeless and ddx11/timeless mutants were generated in this study by CRISPR-Cas9-mediated gene disruption. DT40 WT BU- $1^{\Delta G4}$ cells were generated by deleting the +3.5 G4 motif from both alleles of the BU-1 locus ⁴². Other mutant lines have also been described previously: fancj ⁴⁴, ddx11, fancj/ddx11 and tipin ^{45,51}.

Complemented *ddx11* cells were obtained by transfecting the KO cell lines with pEGFP-C1 (Clontech) harbouring chicken *DDX11* cDNA or a helicase-dead variant (K87A) ⁴⁵ with selection of G418-resistant (2 mg/ml) clones followed by screening for GFP expression by flow cytometry. Complemention of *timeless* was achieved by transfection of plasmids encoding the human Timeless WT cDNA (Addgene plasmid 22887), or truncated versions thereof, selected with zeocin (1 mg/ml) followed by screening for Flag-positive clones by western blot. Complemented *fancj* cells were obtained as follows. First, *fancj*+/- cells harbouring a tamoxifen-regulatable Cre recombinase, Mer-Cre-Mer ⁶⁰ were transfected with pXSPN ⁶¹ with wild type human YFP-FANCJ flanked by loxP sites. The second allele of the *FANCJ* locus was then disrupted with a previously described targeting construct ⁴⁴. For testing the effect of FANCJ mutations, a second pXPSN plasmid was introduced harbouring CFP-FANCJ[mut] without loxP sites, where 'mut' is the desired mutation. Treatment with tamoxifen results in excision of the wild type FANCJ transgene leaving the cells either FANCJ-deficient or solely expressing FANCJ[mut].

Transfections for CRISPR-Cas9 targeting were performed using a Neon electroporator (Thermo Fisher Scientific), 20 x10⁶ cells and 20 μg plasmid DNA and 3 pulses of 1400V with

intervals of 10 msec. For cDNA expression either the Neon, with the same conditions, or a BioRad electroporator with 1 pulse of 250V 950 μ F 100 Ω in 4 mm cuvettes, was used.

Gene disruption using CRISPR-Cas9

Guide RNA sequences used for disrupting *DDX11* and *TIMELESS* in DT40 cells are listed in Supplementary Table 1. Guide RNAs were designed using the CRISPOR online tool of the Zhang lab (https://zlab.bio/guide-design-resources). Each guide was cloned into the pSpCas9(BB)-2A-GFP (PX458) plasmid ⁶² and transfected as described above. 24 h after transfection cells were collected by centrifugation at 400g for 5 minutes and sorted at single live (PI⁻), GFP-positive cell per well in 96-well plates using a MoFlo (Beckman Coulter) or a Synergy (Sony) cell sorter. Cells were grown for 2 weeks, clones were collected and genotyped by Sanger sequencing of the targeted region amplified by PCR, gel-purified with a QIAquick Gel Extraction Kit (QIAGEN) and blunt-cloned using Zero Blunt TOPO PCR Cloning Kit (Thermo Fisher Scientific). Sequences were aligned against the WT DT40 genome using MacVector software.

Bu-1 staining and fluctuation analysis for generation of Bu-1 loss variants

Bu-1 staining and fluctuation analysis were performed as described previously 42,43 . Briefly, cells (confluency between $0.4 - 2 \times 10^6$) were directly stained with anti-Bu-1 conjugated with phycoerythrin (Santa Cruz 5K98-PE 70447 or Invitrogen 21-1A4-PE MA5-28754) at 1:100 dilution for 10 minutes at room temperature. Cells were analysed by flow cytometry using an LSRII flow cytometer (Beckman Coulter) and the FloJo software. Bu-1 expression in single live cells was gated using side scatter (SSC) in the y-axis and PE fluorescence in the x-axis. To perform fluctuation analysis, Bu-1-positive single cells were sorted and grown for ~ 20 generations before staining and flow cytometry analysis as described above.

G-quadruplex ligand

The small molecule pyridostatin (PDS) ⁵² was purchased from Sigma-Aldrich.

Cell proliferation and viability assays

 5×10^5 cells were seeded in 10 cm dishes containing 10 ml of media with G4 ligand/DMSO. Living cells were counted at 24, 48, 72 and 96 h on a Vi-Cell cell counter (Beckman Coulter). Doubling time was calculated during the exponential growth phase using the formula Doubling Time = duration * log (2) / log (Final Concentration) – log (Initial Concentration).

Cell viability was determined 72 h after PDS or cisplatin treatment using a CellTiter 96 AQueous One Solution Cell Proliferation Assay (Promega), according to the manufacturer's instructions. 1 x 10⁵ cells/well were seeded in 96-well plates in 200 µl media containing PDS or cisplatin. 72 h later, 20 µl of MTS reagent was added to 100 µl cells and incubated for 2-4 h at 37°C. Absorbance was measured at 492 nm (formazan salt) in a PHERAstar plate reader (BMG Labtech). The percentage of cell viability was calculated as follows: Cell viability (%) = (Test absorbance/Control absorbance) * 100.

Flow cytometry to monitor γ-H2AX

Cells were collected, fixed with 1% paraformaldehyde for 15 minutes in ice, washed with PBS and fixed with 70% ethanol for at least 24 h at -20° C. Cells were blocked and permeabilized in BD wash/permeabilization Buffer (BD Biosciences) and then incubated with anti- γ -H2AX antibody (05-636, Merck, diluted at 1:500 in BD Buffer) for 2 h at room temperature. Samples were washed twice with BD Buffer, incubated with anti-mouse AF594 antibody (A11062, Thermo Fisher Scientific, diluted at 1:200 in BD Buffer) for 1 h at room temperature in the dark, and then washed twice with BD Buffer. Cells were resuspended in PBS containing 0.5% BSA and 1 μ g/ml DAPI. Cells were analysed by flow cytometry using an LSRII flow cytometer (Beckman Coulter) and the FloJo software. A gate for γ -H2AX-positive cells was defined based on untreated cells that were considered negative for γ -H2AX staining (see Fig. 6b).

Protein expression and purification

Full-length human Timeless (1-1208) was amplified from I.M.A.G.E cDNA clone IRATp970C0576D (Source Bioscience) and cloned into pRSF-Duet1 (Novagen) with an N-terminal His₁₄-SUMO tag. Our previous structural analysis had demonstrated that residues 239-330 comprise a large disordered loop within the folded N-terminal region of Timeless ⁶³ and their removal improved dramatically the biochemical behaviour of the protein; they were therefore replaced by a short linker consisting of residues (Gly-Ser-Thr)₂. All experiments reported here were performed with this optimised Timeless construct. The full-length human Tipin gene (1-301) was codon-optimised for *E. coli* expression (Life Technologies), and cloned into the pGAT3 vector for expression fused to an N-terminal His₆-GST tag ⁶⁴. Timeless and Tipin were co-expressed in *E. coli* Rosetta2 (DE3). The complex was purified using Ni-NTA agarose (Qiagen) followed by cleavage of the tags with TEV and SUMO proteases, ion exchange chromatography (HiTrap Q HP, GE Healthcare) and size-exclusion chromatography

(Superdex 200 16/60, GE Healthcare). The DNA-binding domain of human Timeless (DBD; residues 816-954) was cloned into pRSF-Duet1 (Novagen) with an N-terminal His₁₄-SUMO tag and expressed in *E. coli* Rosetta2 (DE3). Purification entailed Ni-NTA agarose (Qiagen), His₁₄-SUMO tag cleavage with SUMO protease, Ni-NTA re-capture of the cleaved tag and size-exclusion chromatography (Superdex 75 16/60, GE Healthcare). Timeless 883-947 (DBD C-term) was expressed and purified in the same way as DBD.

X-ray crystal structure determination

The DBD C-term was crystallised by vapour diffusion, mixing equal volumes of protein at 800 μM and a solution of 200 mM sodium formate and 18% PEG 3350 at 19 °C. For cryoprotection of the crystals the mother liquor was replaced with a 2:2:1 (volume ratio) mixture of reservoir solution, protein buffer and 100% glycerol. X-ray diffraction data for a native crystal was collected at the I24 beam line at Diamond Light Source, Oxford, UK. The dataset was processed with XDS ⁶⁵ in space group P65 to a resolution of 1.15 Å.

For structure determination by anomalous scattering, crystals were grown of Se-Met labelled DBD. Selenomethionine was incorporated in the DBD using metabolic inhibition in the presence of Se-Met during bacterial expression in minimal media. Diffraction data at the peak wavelength of 0.9793 Å was collected at the Proxima2A beam line of the Soleil Synchrotron, Gif-sur-Yvette, France. The data was processed to 2.4 Å with XDS ⁶⁵ in the same space group and cell dimensions as for the native crystal. The structure was solved exploiting the anomalous signal of the peak dataset using Autosol (Phenix) ⁶⁶. A largely complete Se-Met model was used to solve the native dataset by molecular replacement using PHASER ⁶⁷. The model of the resulting solution was subsequently extended by Autobuild (Phenix) ⁶⁶ and completed by iterative cycles of manual building and model refinement in Coot ⁶⁸ and Phenix ⁶⁶. The structural figures were generated with Chimera ⁶⁹.

NMR spectroscopy

For expression of ¹⁵N- or ¹³C, ¹⁵N-labeled Timeless 816-954, ¹³C₆-glucose and/or ¹⁵NH₄Cl was used as the sole carbon/nitrogen source in M9 minimal medium. NMR measurements were made on ~0.5 mM solutions in 10% ²H₂O, 10 mM phosphate (pH 6.4), 40 mM KCl, 0.5 mM EDTA. Experiments were recorded at 25 °C on Bruker AVANCE III 600 or 800 MHz spectrometers equipped with QCI or TXI cryoprobes. Data were processed using the AZARA

suite of programs (v. 2.8, © 1993-2019; Wayne Boucher and Department of Biochemistry, University of Cambridge, unpublished). Backbone assignments were derived from established versions ⁷⁰ of HNCA, HN(CO)CA, HN(CO)CACB, HNCACB and HNCO experiments acquired with non-uniform sampling, and side chains using 3D (H)CC(CO)NH-TOCSY, TOCSY-15N-HSQC, NOESY-15N-HSQC, HC(C)H-TOCSY and NOESY-13C-HSQC. Assignment was carried out using CcpNmr Analysis v. 2.4 71. For the heteronuclear NOE experiments, either 4 s of ¹H saturation using a 120° pulse train or a 4-s delay was employed prior to the first ¹⁵N pulse. Inter-proton distance restraints were derived from NOE peak heights in the 3D NOESY spectra using the relaxation matrix method (which accounts for spin diffusion) and r^{-6} summation for ambiguous NOEs. Backbone ϕ and ψ torsion-angle restraints were obtained using TALOS+ 72. The restraints provided the input for the iterative assignment protocol ARIA v.1.2 73. For the final iteration, 100 structures were calculated with the violation tolerance set to 0.1 Å. After the last iteration, the 20 lowest-energy structures were subjected to a final water refinement to give an ensemble of 20 structures. Torsion-angle molecular dynamics simulations were performed using CNS ⁷⁴. The simulated annealing protocol used to calculate each structure included 60,000 molecular dynamics steps, including those for refinement, and two cooling stages (to 1000 K and 50 K).

EMSA-based DNA binding experiments

G-quadruplex sequences (Supplementary Table 2) were folded by heating the DNA to 95°C followed by gradual cooling to 10°C, in TE buffer supplemented with 150 mM KCl. Reactions contained 5 μM Timeless-Tipin and 5 μM 6FAM-labelled DNA in 25 mM HEPES pH 7.1, 150 mM KCl, 5 mM MgCl₂ and 1 mM DTT. Reactions were incubated on ice for 30 minutes, after which native loading dye was added and the samples analysed on a 1 % agarose gel in EMSA buffer (0.5x TBE, 10 mM KCl). Gels were run for 1 hour at 50 V and 4°C, and bands were visualized under UV light.

Fluorescence anisotropy analysis of DBD domain and Timeless-Tipin complex binding to DNA.

DNA sequences were annealed by heating to 95°C followed by gradual cooling to 10°C, using a thermocycler, in TE buffer supplemented with 150 mM KCl. Fluorescence anisotropy measurements were recorded at 25°C in a plate reader (PHERAstar FS; BMG Labtech). Excitation for Cy3-3'labeled DNA was at 540 nm and emission at 590 nm (20 nm bandwidth

for both). Excitation for FAM6-5'labeled DNA was at 485 nm and emission at 520 nm (10 nm bandwidth for both). For DBD titrations, each well contained 10 nM 3'Cy3-labelled DNA and increasing concentrations of DBD protein in assay buffer (25 mM HEPES-NaOH pH 7.2, 150 mM KCl, 0.5 mM EDTA, 0.005% Tween-20). For Timeless-Tipin complex titrations, each well contained 10 nM 6FAM-labelled DNA, and increasing concentrations of the complex in assay buffer (25 mM HEPES pH 7.1, 150 mM KCl, 1 mM DTT, 5 mM MgCl₂). The voltage gains and focal heights were adjusted using the instrument software using free fluorescein as reference (35 mP) and data collection was set to 200 flashes. Each data point is the mean of at least 3 independent measurements. The averaged data was analysed using nonlinear fits of ligand-depletion binding isotherms adapted for fluorescence-anisotropy measurements, assuming one to one binding model and using the *ProFit* software package (Quantum Soft). No corrections were necessary for changes in the quantum yield as a function of protein concentration.

Co-immunoprecipitation (Co-IP) and Western Blot (WB)

For whole-cell extracts, the protein samples were lysed in Lysis Buffer (50 mM Tris-HCl pH 7.5, 20 mM NaCl, 1 mM MgCl₂, 0.1% SDS, protease inhibitors (Complete Protease Inhibitor Cocktail Tablets, Merck), phosphatase inhibitors (Halt Phosphatase Inhibitor Cocktail (Thermo Fisher Scientific) and 1 µl/ml benzonase (Merck)) for 15 min at room temperature under agitation, and cleared by centrifugation at 16000 g for 20 min at 4°C. Samples were quantified using the Bradford method and approximately 30 µg were mixed with 5X SDS Page Sample Buffer (Jena Bioscience), boiled for 3 min, fractionated in 4-12% or 10% Nu-PAGE Bis-Tris gels (Thermo Fisher Scientific) in MOPS (200 mM MOP, 50 mM Sodium Acetate, 10 mM Na₂EDTA), and transferred to nitrocellulose membranes using an iBlot2 apparatus (Thermo Fisher Scientific) at 25 V for 7 min. Membranes were blocked for 1 h at room temperature with TBS 0.1% Tween-20 5% skimmed milk. Membranes were incubated with primary antibodies diluted in TBS-T 5% milk overnight at 4°C under agitation, washed with three washes of 10 min each with TBS-T, and incubated with HRP-conjugated secondary antibodies at room temperature for 1 h, followed by three washes with TBS-T. Membranes were developed with Immobilon Crescendo Western HRP substrate (Merck) for 1 min and exposed to X-ray films.

For Co-IP experiments, pcDNA4-Flag-Timeless plasmids (WT and truncated versions) were co-transfected into 2x10⁶ HEK293T cells using Lipofectamine 2000 (Thermo Fisher

Scientific) seeded in 10 cm dishes 16 h prior to transfection (80% confluency at transfection) with pcDNA3-hDDX11 ⁴⁵ (30 μg total DNA). 24 h after transfection, cells were detached with trypsin and washed in cold PBS. Pellets were lysed in NETN-M buffer with glycerol (150 mM NaCl, 50 mM Tris-HCl pH8, 0.5% Igepal, 1 mM EDTA, 5 mM MgCl₂ and 10% glycerol, supplemented with protease and phosphatase inhibitors) and benzonase for 15 min at room temperature under agitation. Samples were sonicated with 8 cycles of 30s with 30s intervals at the low setting in a Bioruptor Plus water bath sonicator (Diagenode) and centrifuged for 15 min at 16000 g at 4°C. Supernatant was collected, quantified and 2 mg of total cell extract were mixed with 30 μl Flag-M2 magnetic beads (M8823, Sigma-Aldrich). Samples were incubated at 4°C for 2 h under agitation. Beads were then washed 5 times for 5 min each with lysis buffer, and proteins were eluted by boiling with Sample Buffer Laemli 2X (Sigma-Aldrich) for 5 min at 95°C. Samples were subjected to analysis by WB as described above, using the indicated antibodies.

Primary antibodies used: mouse anti-Flag (M2 F3165, Merck), mouse anti-DDX11 (D-2 271711, Santa Cruz) and mouse anti- α -tubulin (T6074, Sigma). Secondary antibody: goat anti-mouse HRP (P0447 Dako).

RNA extraction and RNA-seq library preparation

RNA was extracted from three independent cell populations using the RNeasy Mini Kit (QIAGEN), according to the manufacturer's instructions. RNA sample quality was checked in BioAnalyzer RNA Pico chips (Agilent) and only high quality samples (RIN > 7) were used to build the libraries. 750 ng RNA were used to build the next generation sequencing libraries with the NEBNext Ultra II RNA Library Prep Kit for Illumina (New England BioLabs), NEBNext Poly(A) mRNA Magnetic Isolation Module and NEBNext Multiplex Oligos for Illumina (Index Primers Set 1 and 2), according to the manufacturer's intructions. Library quality was checked in BioAnalyzer DNA High Sensitivity chips (Agilent), and quantified using the KAPA Library Quanti Kit (Illumina) Universal qPCR Mix (KAPA Biosystems) on an ABI Prism ViiA 7 Real-Time PCR System (Thermo Fisher Scientific). Libraries were sequenced on a HiSeq4000 (Illumina).

RNA-seq library alignment

RNA sequencing reads were aligned to the GRCg6a chicken genome using Bowtie 2 ⁷⁵. Expression estimates (transcripts per million; TPM) were calculated using the rsem-calculate-

expression command of RSEM ⁷⁶ and the version 96 of the chicken genome gene annotation from Ensembl. Only transcripts with an expression above or equal at 1 TPM were considered for further analysis. Gene expression matrixes were built using rsem-generate-data-matrix and three independent biological replicates. Differentially expressed genes were called using EBseq ⁷⁷. Change in gene expression was estimated using the EBseq posterior Fold Change (PostFC) value and genes were considered differently expressed when the EBseq posterior probability of being differentially expressed (PPDE) was above or equal at 0.95.

Promoter sequence analysis

Gene promoter sequences were recovered from the GRCg6a chicken genome assembly using the getfasta command of BEDTools 78 . Sequence analyses were performed using custom scripts in the R environment (http://www.R-project.org/). Assessment of the enrichment of G-quadruplexes within the promoter of DDX11- and TIMELESS-dependent genes was based on regular expression matching algorithms to monitor the cumulative count of the G3N12 motif within promoters. This was computed by identifying the number of sequences of the form $d(G_{3+}N_{1-12}G_{3+}N_{1-12}G_{3+})$, where N is any base, which represents the loose definition of G4 forming sequences 79 . Densities of G-quadruplex forming sequences around promoters were computed by assessing the number of promoter sequences containing sequences of the form $d(G_{3+}N_{1+}G_{3+}N_{1+}G_{3+}N_{1+}G_{3+})$ in windows of 50 nucleotides sliding by 10 nucleotides normalised to the total number of promoter sequences analysed.

Statistical analysis of RNA-seq data

Data were analysed and statistics performed in the R environment. Overlaps between gene lists were tested using Fisher's exact tests. Differences between distributions were tested using Kolmogorov–Smirnov tests.

Figure Legends

Figure 1. Timeless and Tipin are required to maintain processive replication past G4 structures *in vivo*.

a. The BU-1 locus as a model system to record G4-dependent replication stalling. The leading strand of a replication fork entering the locus from 3' end stochastically stalls at the +3.5 G4, leading to the formation of a region of ssDNA, with interruption of parental histone recycling and of histone modifications necessary to maintain normal expression of the locus ⁴². **b.** Instability of BU-1 expression in timeless cells. FACS plots of wild type and timeless (clone 1) DT40 cells stained with anti-Bu-1 conjugated with phycoerythrin. Each line represents the Bu-1 expression profile of an individual clonal population. Unstained controls are shown in blue. c. Fluctuation analysis for Bu-1 loss in wild-type DT40 cells and two independent timeless clones generated by CRISPR-Cas9 targeting (clones 1 and 2; Supplementary Fig. 1), timeless (clone 1) complemented by expression of human Timeless cDNA and a timeless mutant on a background in which the endogenous +3.5 G4 has been deleted (Δ G4) ⁴². **d.** Fluctuation analysis for Bu-1 loss in DT40 wild-type and *tipin* cells. In c. and d., each symbol represents the percentage of cells in an individual clone expanded for 2-3 weeks that have lost Bu-1high expression. At least two independent fluctuation analyses were performed, with 24-36 individual clones each cell line per repeat. Bars and whiskers represent median and interquartile range, respectively. **** p < 0.0001; one-way ANOVA.

Figure 2. Identification and characterisation of a DNA-binding activity in Timeless

a. Schematic drawing of human Timeless, showing its domain structure, interacting proteins Tipin and Parp1, and its DNA-binding domain (DBD). A multiple sequence alignment of vertebrate Timeless sequences is shown underneath, with amino acid conservation coloured according to the Clustal colour scheme. The alignment is annotated with the extent and secondary structure elements of the two helical domains (N-term and C-term) composing the DBD. **b.** Ribbon drawing of the 1.15 Å crystal structure at of the DBD C-term. Helices are in red and labelled H1 to H4. **c.** Ribbon drawings of the N-term and C-term domains of the DBD determined by NMR. The two domains are shown in the same orientation to highlight their high degree of three-dimensional similarity. The superposition of the 20 lowest energy structures is shown for each domain. **d.** The DNA-binding affinity of DBD was measured by fluorescence anisotropy, titrating DBD protein against Cy3 3'-labelled ssDNA, dsDNA and G-quadruplex (G4) DNA. Top panel shows binding curves for ss- and dsDNA, the bottom panel

shows the binding curve for the G4 DNA substrate. The data points represent the mean of at least 3 independent experiments and the error bars indicate one standard deviation (SD). e. Ribbon diagram of the superposition of DBD C-term with the highly similar DNA-binding domains of telomeric protein TRF1 (PDB ID 1W0T) ⁴⁶ and the bacterial cell-cycle regulator GcrA (PDB ID 5Z7I) ⁴⁷ in complex with their DNA substrates. A similar DNA-binding mode by DBD would cause a steric overlap of helix H4 with the phosphate backbone of dsDNA. DBD C-term is in light blue, TRF1 and GcrA proteins in brown and their DNA substrates in khaki.

Figure 3. The Timeless-Tipin complex shows a preference for binding G-quadruplex DNA.

Fluorescence anisotropy was used to measure the binding affinity of Timeless-Tipin for the indicated DNA sequences. **a.** ssG4: G4 flanked by single-stranded DNA; ssHP: hairpin flanked by single-stranded DNA; ss: single-stranded DNA (see Supplementary Table 2 for sequence details). **b.** Binding affinity of Timeless-Tipin for a range of G-quadruplex DNA sequences (see Supplementary Table 2 for sequence details and references). Single-stranded (ss20: 5'-6FAM-ATAAGAGTGGTTAGAGTGTA) and double-stranded (ds20: ss20 annealed to complementary sequence) DNA were also tested as controls. Each data point is the mean of at least 3 independent experiments and the error bars indicate one SD.

Figure 4. The C-terminus of Timeless is required for processive G4 replication.

Fluctuation analysis for the generation of Bu-1 loss variants in wild-type cells, *timeless* (clone 1) cells, *timeless* (clone 1) complemented with human Timeless ΔDBD cDNA (hTim $\Delta 81$ -965) and *timeless* (clone 1) complemented with human Timeless truncated at the PARP binding domain (hTim[1:1000]) cDNA and *timeless* cells generated by CRISPR-Cas9 targeting exon 16 which truncates the protein removing the CTD containing both the DBD and the PARP binding domains. At least two independent fluctuation analyses were performed with 24-36 individual clones each cell line per repeat. Bars and whiskers represent median and interquartile range, respectively. * p < 0.05 and **** p < 0.0001; one-way ANOVA for comparison with the wild-type cells.

Figure 5. DDX11 is required for processive replication in collaboration with Timeless.

a. Fluctuation analysis for Bu-1a loss in wild-type DT40 cells, two independent ddx11 clones generated by CRISPR-Cas9 targeting (clones 1 and 2) and one ddx11 clone generated by conventional homologous recombination gene targeting (clone 3). Each symbol represents the percentage of cells in an individual clone expanded for 2-3 weeks that have lost Bu-1a^{high} expression. **b.** Fluctuation analysis for Bu-1a loss variant generation in wild-type cells, ddx11 (clone 1) cells, ddx11 (clone 1) complemented by expression of chicken DDX11 WT cDNA, ddx11 (clone 1) complemented by expression of helicase-dead form of chicken DDX11 (K87A) cDNA, and a ddx11 clone generated in cells in which the endogenous +3.5 G4 has been deleted (Δ G4). **c.** Fluctuation analysis for Bu-1 loss in two independent ddx11/timeless double-mutant clones. Fluctuation analyses for wild type, timeless (clone 1) and ddx11 (clone 1) are shown for comparison, as are fluctuation analyses fancj and fancj/ddx11 double-mutants. In all cases, at least two independent fluctuation analyses were performed, with 24-36 individual clones each cell line per repeat. Bars and whiskers represent median and interquartile range, respectively. ** p < 0.01 and **** p < 0.0001; one-way ANOVA for comparison with wild-type cells.

Figure 6. Timeless and DDX11-deficient cells have impaired growth and increased H2AX phosphorylation in the presence of a G4 ligand.

a. Growth curves for DT40 wild type, ddx11, timeless and ddx11/timeless cells, with and without 4 μM pyridostatin (PDS). Cells were seeded at 5 x10⁴ cells/ml on day 0 and the viable cells were counted each 24 h for 4 days. Bars represent SD of two independent experiments performed in duplicate. Doubling times (DMSO): WT 13 hours, timeless 18 hours, ddx11 16 hours, ddx11/timeless 24 hours. Doubling times (PDS): WT 13.6 hours, timeless 27 hours, ddx11 25.7 hours, ddx11/timeless 47.5 hours. **b.** DDR signalling detected by phosphorylation of histone H2AX (γ-H2AX) by flow cytometry in untreated cells or cells exposed to 4 μM PDS for 3 days. Pale histogram, untreated; dark histogram, treated; black dotted line, positive control cells treated with 0.1 μM cisplatin, also for 3 days. **c.** Quantification of γ-H2AX in DT40 wild type, ddx11, timeless and ddx11/timeless cells treated with 4 μM PDS for 3 days. The whiskers represent the minimum to maximum variation of three independent experiments performed in duplicate. * p < 0.05, ** p < 0.01, *** p < 0.001 unpaired, 2-tailed t-test for each pairwise comparison +/- PDS.

Figure 7. Loss of DDX11 and Timeless leads to genome wide expression dysregulation of genes with G4s around the transcription start site (TSS). a. Correlation of magnitude and direction of change of genes dysregulated (relative to wild type) in timeless vs. ddx11 DT40 cells. r_s (Spearman rho) is shown for each correlation. **b.** Cumulative distribution of genes containing n (x-axis) G4 motifs (see Methods) within 1.5kb of the TSS in genes deregulated in timeless (red) and ddx11 (blue) compared with all genes (black). p values calculated with the Kolmogorov–Smirnov test. c. GC content around the TSS in in *timeless* (red) and *ddx11* (blue) compared with all genes (black). **d.** Metagene analysis showing G4 frequency (see Methods) separated for coding and template strands around the TSS of genes dysregulated in timeless (left panel), ddx11 (centre panel) and fancj (right panel) mutants compared with all genes (black line). e & f. Two models, which are not mutually exclusive, for how Timeless might detect G4s at the replication fork. e. Recognition of a G4 by the Timeless C-terminus formed in the negatively supercoiled ssDNA ejected behind the CMG helicase, possibly in conjunction with an interaction with PARP1 bound to the structure ⁵⁷ with recruitment of DDX11. Failure of this mechanism would lead to uncoupling of the replicative helicase and leading strand polymerase. For further discussion of this issue and how G4s might interact with the replisome, see Lerner & Sale, 2019 ³. **f.** G4 detection and processing ahead of the CMG helicase. A leading strand G4 formed ahead of the replisome must be traversed by the CMG helicase, in an analogous manner to that proposed for traverse of an DNA interstrand crosslink ⁵⁸. This would lead the structure to stall the leading strand DNA polymerase and uncouple DNA unwinding and synthesis. Detection of the structure by the C-terminus of Timeless, Recruitment of DDX11 by Timeless would unwind the structure, obviating the need for further remodelling of the replicative helicase and preventing the structure meeting the leading strand polymerase.

Competing interests

The authors declare no competing interests.

Data and Reagent Availability

Coordinates and structure factors have been deposited in the Protein Data Bank (https://www.rcsb.org) under accession codes 6T9Q for the crystal structure of the C-terminal repeat of DNA-binding domain and 6TAZ (BMRB ID 34443) for the NMR structure of the DNA-binding domain. The RNA sequencing data has been deposited in GEO

(<u>https://www.ncbi.nlm.nih.gov/geo/</u>) with accession number GSE139256. Requests for materials can be addressed to the corresponding authors.

Author Contributions

J.E.S., L.P., L.K.L. and S.H. conceived the overall project. L.K.L. generated the cell lines and performed the molecular biology experiments (flow cytometry, survival, IP, RNAseq). S.S. constructed the truncated versions of Timeless plasmids. D.S. generated the mer-cre-mer FANCJ cell line. P.M. performed the RNAseq data analysis. S.H. expressed and purified both DBD and DBD C-term, determined the x-ray crystal structure of the C-term, and performed the initial characterisation of the DNA-binding properties of the DBD. M.L.K., A.B. and J.D.M performed the DNA-binding experiments. K.S. and S.H. determined the NMR structure of the DBD. D.B. provided key KO cell lines. L.K.L., J.E.S. and L.P. wrote the initial manuscript with assistance and editing by all authors.

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