Replicative DNA polymerases promote active displacement of SSB proteins during lagging strand synthesis

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ABSTRACT

Genome replication induces the generation of large stretches of single-stranded DNA (ssDNA) intermediates that are rapidly protected by single-stranded DNA-binding (SSB) proteins. To date, the mechanism by which tightly bound SSBs are removed from ssDNA by the lagging strand DNA polymerase without compromising the advance of the replication fork remains unresolved. Here, we aimed to address this question by measuring, with optical tweezers, the real-time replication kinetics of the human mitochondrial and bacteriophage T7 DNA polymerases on free-ssDNA, in comparison with ssDNA covered with homologous and non-homologous SSBs under mechanical tension. We find important differences between the force dependencies of the instantaneous replication rates of each polymerase on different substrates. Modeling of the data supports a mechanism in which strong, specific polymerase-SSB interactions, up to ~12 k_BT, are required for the polymerase to dislodge SSB from the template without compromising its instantaneous replication rate, even under stress conditions that may affect SSB–DNA organization and/or polymerase–SSB communication. Upon interaction, the elimination of template secondary structure by SSB binding facilitates the maximum replication rate of the lagging strand polymerase. In contrast, in the absence of polymerase–SSB interactions, SSB poses an effective barrier for the advance of the polymerase, slowing down DNA synthesis.

INTRODUCTION

Genome duplication induces the formation of transient single-stranded DNA (ssDNA) intermediates, usually thousands of nucleotides long, which are rapidly covered by single-stranded DNA-binding (SSB) proteins. SSBs bind ssDNA in a sequence-independent manner with high affinity, generally in the picomolar range (1–4). SSB binding protects ssDNA from degradation and greatly stimulates DNA replication in vitro. This stimulation is in part due to the many roles attributed to SSBs (reviewed in (1,2,4)), such as prevention of ssDNA degradation, removal of ssDNA secondary structure, increased primer recognition and initiation, decreased non-specific binding of DNA polymerase to the template, and stimulation of the average primer extension and strand displacement activities of the DNA polymerase. Due to their many functions during DNA replication, SSBs are found in all organisms, and are absolutely required for the synthesis of genomic and mitochondrial DNA.

In humans, the replication mechanism of mitochondrial DNA remains actively debated, though increasing experimental evidence indicates that during replication of the L-strand (equivalent to the lagging strand of nuclear DNA), the mitochondrial replicase, PolY, works on long stretches of ssDNA intermediates covered by mitochondrial SSB, mtSSB (5–9). Human mtSSB has significant sequence and structural homology (10,11) with the well-characterized...
SSB from *Escherichia coli* (EcoSSB). Sequence comparison showed that the major differences between these two proteins are that residues 1–9 and 55–59 of mtSSB are missing in EcoSSB and that EcoSSB contains an additional C-terminal tail missing in mtSSB (11). Apart from these differences, these two SSBs share fast binding ($10^9$–$10^{11}$ M$^{-1}$ s$^{-1}$) and slow dissociation ($10^{-1}$–$10^{-2}$ s$^{-1}$) kinetics to and from ssDNA (12,13) and bind preformed ssDNA as tetramers in at least two different modes. In each mode, ~30 (mtSSB$\beta_0$) or ~60 (mtSSB$\beta_{1}0$) nucleotides are organized per tetramer depending on the salt and protein concentrations (3,12–14). In fact, both proteins stimulate *in vitro* the primer extension activity of Pol (15,16). Interestingly, biochemical and physiological studies showed that a mtSSB variant (mtSSB$\beta_2$,3) carrying a deletion of amino acids 51–59 (those missing in EcoSSB) stimulates the activity of Pol $\beta$ less than wild-type mtSSB (17). These nine amino-acids pertain to loop 2.3, which is a flexible domain characteristic of vertebrate mtSSBs. The loop 2.3 is located on the surface of mtSSB close to the ssDNA-binding channel but is not involved in ssDNA binding. These findings suggest that specific structural elements of mtSSB may play a role in facilitating the primer extension activity of Pol (i.e. via species-specific interactions).

To date, several questions remain unanswered about how apparently competing functions of polymerase and SSB are coordinated during synthesis of the lagging strand. First, the mechanism by which replicative DNA polymerases dislodge the long-lived SSB–DNA complexes in a manner that does not compromise the advance of the replication fork is unknown. This is a remarkable task considering that during lagging strand synthesis the high density of SSB bound to the template would restrict the capacity of a single SSB to diffuse (18,19), and the absence of free ssDNA ends would prevent SSB from being pushed outward by the advance of the polymerase (20). Second, the nature of the stimulatory effect of SSB on the average replication velocity described in bulk studies is not known. Is stimulation due to direct polymerase–SSB connections, which enhance the instantaneous primer extension rate of the polymerase? Or instead, is it due to an indirect role of SSB in the organization of the template?

Here, we aimed to answer these questions using optical tweezers to determine the coordinated action of human mtSSB and Pol $\beta$, in a single-molecule assay mimicking lagging strand synthesis under different mechanical tensions. Mechanical tension applied to the ends of ssDNA modulates the replication rate of DNA polymerases (21–26), and when applied to the ends of ssDNA–SSB complexes, it regulates the binding modes of SSB proteins (14,27,28). Consequently, mechanical tension can be employed as a useful variable to determine the mechanistic processes involved in the replication of SSB–DNA complexes, and to quantify the energies involved in the mechanism of SSB release by DNA polymerases.

In our studies, we measured and compared the real-time kinetics of individual Pol holoenzymes replicating under mechanical tension either free-ssDNA or ssDNA covered by either human wild-type mtSSB (mtSSB$\beta_{WT}$), the mtSSB variant mtSSB$\beta_{2,3}$ or EcoSSB. In order to explore whether our findings can be extended to other DNA replication systems, we also measured the effect of bacteriophage T7 SSB (gp2.5), mtSSB$\beta_{WT}$ and EcoSSB on the instantaneous replication rate of the T7 DNA polymerase (T7DNAp) under mechanical tension. Interestingly, the two DNA polymerases under study responded similarly to the absence or presence of homologous and non-homologous SSBs in the template. For both DNA polymerases, the instantaneous replication rate increased rapidly with tension (~6 pN), when DNA synthesis occurred on free-ssDNA. In contrast, in the presence of their homologous SSBs, both polymerases replicated at their maximum rates within the entire range of tension at which SSB remained stably bound to the template (~8 pN). Interestingly, non-homologous or mutant SSBs covering the DNA template prevented both DNA polymerases from achieving their maximum replication rates. Overall, these results are compatible with a model in which the secondary structure of free DNA presents a significant barrier (~2 k$_{B}$T) to the advance of these replicative DNA polymerases. Destabilization of the secondary structure by SSB binding favored maximum instantaneous replication rates but only when functional interactions between polymerase and SSB promoting SSB release are established. In particular, for the mitochondrial replication proteins we found that specific polymerase–SSB interactions, as high as ~12 k$_{B}$T (22°C) between residues of loop2.3 of mtSSB and Pol $\beta$, allow this polymerase to displace the SSB tetramers from the template without hindering its replication rate, even under mechanical stress conditions that affect the SSB–DNA structure. The similarities found in the mitochondrial and bacteriophage T7 DNA replication systems suggest that their respective polymerases may use similar mechanisms to dislodge the tightly bound SSBs during lagging strand DNA synthesis.

**MATERIALS AND METHODS**

**Proteins and DNA constructs**

Recombinant mtSSB$\beta_{WT}$ and mtSSB$\beta_{2,3}$ proteins were prepared from bacterial cells as described previously (15). Recombinant catalytic and accessory subunits of human Pol $\beta$ were prepared from S/E9 and bacterial cells, respectively, as described previously and combined in a 1:1.5 molar ratio to reconstitute the holoenzyme (29). Recombinant EcoSSB was purchased from Thermofisher and recombinant T7 DNAp and gp2.5 were purchased from New England Biolabs and LSBio, respectively. The gapped DNA template, consisting of ~900 nt of ssDNA flanked by ~3550 bp dsDNA handles, was prepared from pBacgus11 vector (Novagen) and labeled with digoxigenin (Dig) at one end and biotin at the other, as described in (21). The approximate length of the ssDNA fragment was determined by denaturing gel electrophoresis (Supplementary Figure S1).

**Optical tweezers experiments**

We used a counter propagating dual-beam optical tweezers instrument to manipulate individual DNA molecules (30). Data were monitored at 500 Hz at 22 ± 1°C.

**SSB-free and SSB-bound ssDNA force–extension curves (FECs).** ssDNA was obtained by unzipping mechanically
a dsDNA hairpin of 510 bp (31) using the oligonucleotide hybridization method described previously, Supplementary Figure S2 (14). SSB proteins were introduced inside the flow cell after dilution to 50 nM (or 100 nM for gp2.5) in the replication buffer without dNTPs: 50 mM Tris pH 8.5, 30 mM KCl, 10 mM Dithiothreitol (DTT), 4 mM MgCl₂, 0.2 mg/ml bovine serum albumin. SSB binding to preformed ssDNA was carried out at constant tensions <8 pN. Once equilibrium was reached, tension was varied with a constant pulling rate of 150 nm s⁻¹, in order to obtain the final FEC of each SSB–DNA complex. A minimum of 5 FECs were obtained for ssDNA and for each SSB–DNA complex.

**Primer extension activities.** A gapped DNA molecule was tethered between an anti-Dig and streptavidin covered beads. Polγ and T7DNAp were introduced inside the flow cell diluted to 2 nM (unless otherwise indicated) in the replication buffer containing the four dNTPs (50 μM). Individual replication traces were obtained in free or SSB-covered gapped-DNA substrates. SSB concentration was 50 nM in all cases except for gp2.5, which was 100 nM. Data were taken in the ‘constant force feedback’ mode. In this mode, the distance between the beads was adjusted to maintain a constant tension in the DNA as the single-stranded template is replicated to dsDNA.

**Data analysis**

**Secondary structure.** The amount of secondary structure in ssDNA as a function of mechanical tension was quantified from experimental FECs of ssDNA (Figure 1C) according to Bosco et al. (32), Supplementary Figure S3.

**SSB-binding mode and protein coverage determination.** The binding mode (number of nucleotides wrapped per SSB tetramer) and protein coverage (the fraction of the ssDNA covered by SSB) of mtSSB and EcoSSB were determined from experimental ssDNA–SSB FECs according to a phenomenological model described previously (33), Supplementary Figure S2.

**ΔGSSB calculation.** The average Gibbs energy to unwrap one nucleotide from an SSB tetramer at a specific force value was calculated by integrating the area between the FECs of the SSB-bound and SSB-free ssDNA above that specific force (Supplementary Figure S2). The area was divided by the total number of nucleotides and corrected by the protein coverage. The change of ΔGSSB with force for mtSSBWT, mtSSB₁,₃ and EcoSSB are shown in Supplementary Figure S2D.

**Average velocity and processivity.** The number of nucleotides incorporated by Polγ and T7DNAp was obtained by dividing the distance change between the beads by the change in extension at a given force accompanying the conversion of one SSB-free, or SSB-bound, single-stranded nucleotide into its double-stranded counterpart. The average extensions of SSB-free and SSB-bound ssDNA nucleotides as a function of force are shown in Figure 1C. The dsDNA was approximated with the worm-like chain model for polymer elasticity with a persistent length of $P = 53$ nm and stretch modulus $S = 1200$ pN/nm (34). The average replication rates at each tension were determined by a line fit to the traces showing the number of replicated nucleotides versus time. The final rate at each tension was obtained by averaging over all of the traces taken at similar tension values (±1 pN).

**RESULTS**

**Detection of Pol activity and determination of SSB–ssDNA force–extension curves**

We used a highly stable, miniaturized optical tweezers device (30) to investigate the real-time kinetics of individual Pol molecules replicating a gapped DNA molecule containing a ~900 nt ssDNA segment flanked by ~3550 bp dsDNA handles attached between two micro-sized beads (‘Materials and Methods’ section). One bead is held in the optical trap and the other on top of a micropipette (Figure 1A). We monitored at constant force, from ~1 to 17 pN, the end-to-end distance change of the DNA construct ($Δx$ in Figure 1A) as Pol converts to dsDNA the ssDNA segment, either not covered or covered in situ by mtSSBWT, mtSSB₁,₃ or EcoSSB proteins (Figure 1B and ‘Materials and Methods’ section). Experiments were performed at salt conditions appropriate for DNA synthesis (30 mM KCl and 4 mM MgCl₂) (14,17) and at 50 nM SSB concentration (for all three SSB proteins). These conditions are known to favor the low binding site-size mode of mtSSBWT, mtSSB₁,₃ and EcoSSB (1,3,13,14), which has been recently shown to represent the binding mode selected preferentially during the process of DNA replication (14). Indeed, analysis of the force–extension curves (FECs) of the three SSB–DNA complexes (Figure 1C) indicated that under these conditions all three SSBs bind to ssDNA in the low site-size mode (~30 nt/tetramer) and cover 80–90% of the template at tensions below 8 pN, Supplementary Figure S2 (33). Above this tension, we and others have shown that force promotes significant ssDNA unwrapping from SSB, leading to release of the protein from the template (14,28).

**Template secondary structure hinders the instantaneous replication rate by Pol**

We first aimed to characterize the real-time kinetics of Pol on SSB-free ssDNA under mechanical tension. Analysis of independent replication traces of Pol on ssDNA ($N = 100$) showed that at the lowest tensions (~1 pN) the average number of nucleotides (nt) incorporated was $450 ± 25$ nt (processivity), and the average rate of DNA synthesis was $6 ± 1$ nt s⁻¹ (Supplementary Figure S5). These values are in good agreement with the average processivity, ~560 nt and average primer extension rate, ~6 nt s⁻¹ described in vitro.
for the reconstituted human holoenzyme under similar salt conditions (6,15,35,36). Polymerase concentration was kept low (2 nM) to ensure that each DNA synthesis event corresponds to the activity of a single holoenzyme. The time interval between synthesis events was typically about 10 times larger than the duration of a single event. In addition, varying the polymerase concentration 50-fold (from 0.2 to 10 nM) did not change significantly either the average replication velocity or the processivity (Supplementary Figure S6). These observations suggest that each replication run corresponds to the activity of a single polymerase.

We developed a new method (velocity prominence method) to calculate the average replication rate without pauses, $V_{\text{max}}$, from molecular motor trajectories presenting slow averaging rates, such as PolV (Supplementary Data and Supplementary Figure S4). Validation of the method on simulated traces (as described in 37) indicated that $V_{\text{max}}$ values reported for our experimental data may present an average underestimation of ∼13% (Supplementary Data). Analysis of $V_{\text{max}}$ showed that in SSB-free ssDNA $V_{\text{max}}$ depends strongly on the tension applied to the template: $V_{\text{max}}$ increased with tension <6 pN, from ∼15 to 25 nt s$^{-1}$, to gradually decrease toward stalling at tensions >6 pN (Figure 2A). Similar dependencies of the replication rate on tension have been described for other replicative DNA polymerases (23–26).

The activity of PolV, as with majority of replicative DNA polymerases, is hampered by stable secondary structure of the template (38–40). Accordingly, we noticed that PolV reaches its maximum replication rate at ∼6 pN, which is the tension value that promotes the destabilization of ∼100% of the template secondary structure under our experimental conditions (Materials and Methods’ section and Supplementary Figure S3).

Based on these observations, we considered that $V_{\text{max}}$ is modulated by the work to convert a single-stranded nucleotide to double-stranded form, plus the work required to disrupt the secondary structure characteristic of free ssDNA at each tension. We thus fit the measured force–velocity data to the Arrhenius law (41):

$$V_{\text{max}}(f) = k(0) \exp\left(-\frac{\phi f}{k_B T}\right) + \Delta G_{\text{eff}} \cdot (1 - \psi(f)) / k_B T$$

(1)

The free-variables in this model are: $k(0)$, which is the replication rate at zero tension ($f = 0$ pN) in the absence of template secondary structure and $\Delta G_{\text{eff}}$, or the average energy barrier imposed by the template secondary structure on the advance of the polymerase. The parameter $\delta$ defines the kinetic step size of the DNA polymerase and was fixed to 1 nt/step, according to bulk, structural and single-molecule data (42–45). The fraction of ssDNA template bases forming secondary structure as a function of force, $(1 - \psi(f))$, together with the lengths of single- and double-stranded nucleotides at each tension ($x_{ss}(f)$ and $x_{ds}(f)$, respectively), were quantified from experimental FECs (Materials and Methods’ section and Figure 1C; Supplementary Figure S3), $k_B$ is the Boltzmann constant and $T$ the absolute temperature. Equation (1) was fitted to data (Figure 2A) minimizing the mean square difference given for the two free parameters: $k(0) = 23.0 ± 0.7$ nt s$^{-1}$ and $\Delta G_{\text{eff}} = 1.4 ± 0.2$ k$_B$T (Table 1). The value of $k(0)$ matches the intrinsic nucleotide incorporation rates determined previously in vitro for Drosophila melanogaster, 15 nt s$^{-1}$ and human, 37 nt s$^{-1}$ mitochondria DNA polymerases (46–48).

The model points out the important role of the salt-stabilized template secondary structure on modulating the replication rate at low tensions (i.e. it represents an effective barrier of ∼1.4 k$_B$T for PolV under the current experimental conditions). Below 6 pN, the initial increase in the replication rate is...
mainly due to disruption of the template secondary structure by tension, while above 6 pN, in the absence of secondary structure ($\phi(f) = 1$), the replication rate decreases as a result of the work to convert a single- to a double-stranded nucleotide at each tension. Alternative models to interpret the force–velocity behavior of DNA polymerases are discussed in Supplementary Data and Supplementary Figure S7. We note that, although Pol$\gamma$ exhibits both polymerization and proofreading (exonucleolytic) activities in vitro and in vivo, with our current resolution, we did not detect processive force-induced exonucleolytic activity (FIEA) at any tension below 60 pN, as was previously observed for other DNA polymerases (25,26) (see ‘Discussion’ section).

Removal of template secondary structure by mtSSB$\text{WT}$ increases the instantaneous replication rate by Pol$\gamma$

Next, we studied the real-time kinetics of individual Pol$\gamma$ holoenzymes in replicating ssDNA covered with mtSSB$\text{WT}$ at different template tensions. Analysis of independent replication traces ($N = 109$) showed that the presence of mtSSB$\text{WT}$ increased the average replication rate and processivity of Pol$\gamma$, which is consistent with previous bulk studies (15,48,49). Under these conditions, Pol$\gamma$ replicated the full length of the template, $\sim$900 nt, with an average rate at 1 pN of $\sim$9 nt s$^{-1}$ (Supplementary Figure S5). The presence of mtSSB$\text{WT}$ also had a strong effect on $V_{\text{max}}$ (Figure 2A): at $\sim$1 pN $V_{\text{max}} = 24 \pm 2$ nt s$^{-1}$, which is the maximum rate measured on free-ssDNA at a tension destabilizing most of the template secondary structure ($\sim$6 pN). In addition, below 8 pN, $V_{\text{max}}$ presents much weaker force dependence than that measured on free-ssDNA (Figure 2A). These observations suggest that at tensions in which mtSSB$\text{WT}$ remains stably bound to ssDNA (<8 pN), its activity in removing the secondary structure of the template (14) plays a key role in stimulating the instantaneous replication rate by Pol$\gamma$. In fact, the combined effect of tension and mtSSB$\text{WT}$ on $V_{\text{max}}$ is well explained by the model described above Equation (1), when there is no contribution of the secondary structure of the template, $\phi(f)$~1 ($k(0)$ was fixed to $\sim$23 nt s$^{-1}$, by the fit described in the previous section). In other words, in this case, the dependence of $V_{\text{max}}$ on tension simply depends on the work to convert in each nucleotide incorporation step ($\delta = 1$) the extension of a SSB-bound nucleotide ($x_{\text{SSB}}(f)$) to its double-stranded form ($x_{\text{ds}}(f)$). The values of $x_{\text{SSB}}(f)$ and $x_{\text{ds}}(f)$ were determined experimentally from FECs (Figure 1C); therefore, all parameters of Equation (2) (below) were fixed:

$$V_{\text{max}}(f) = k(0) \exp(-\delta \cdot f \cdot x_{\text{SSB}}(f)) / k_B T$$

Regardless of the model used to explain the data, it is striking that mtSSB$\text{WT}$ promotes the maximum rate of replication despite its tight association with ssDNA (below 10 pN). This observation suggests strongly the existence of a specific mechanism for ‘active’ release of mtSSB$\text{WT}$ from the template by Pol$\gamma$. 

![Figure 2](https://academic.oup.com/nar/article-abstract/47/11/5723/5436771)

**Figure 2.** Effect of homologous and non-homologous SS Bs on Pol$\gamma$ replication kinetics. (A) Change of $V_{\text{max}}$ (nt s$^{-1}$) with tension in ssDNA-free (blue symbols, $N = 100$) and mtSSB$\text{WT}$-covered (red symbols, $N = 109$) ssDNA. Lines represent best fit to data with Equation (1) (solid blue line), Equation (2) (solid red line) and Equation (4) (dotted red line). (B) Tension dependence of $V_{\text{max}}$ (nt s$^{-1}$) in ssDNA covered either with mtSSB$\text{WT}$ (red symbols), mtSSB$_{\text{B3}}$ (green symbols, $N = 57$) or EcoSSB (black symbols, $N = 30$). Solid lines represent best fit of the data with Equation (4). For reference, dotted blue line shows fit to $V_{\text{max}}$ in ssDNA-free ssDNA (as shown in A). (C) Moving probability (average velocity/$V_{\text{max}}$) of Pol$\gamma$ on ssDNA (blue) and on ssDNA covered with mtSSB$\text{WT}$ (red), mtSSB$_{\text{B3}}$ (green) and EcoSSB (black). For all figures, error bars show standard errors.
Polymerase–SSB interactions are required for active removal of SSB

In order to characterize the mechanism used by Poly to displace SSB from the template during DNA synthesis and to determine the putative role of species-specific interactions, we next measured the effect of the human mitochondrial SSB variant mtSSB2,3 on the tension-dependent kinetics of Poly. mtSSB2,3 lacks nine residues corresponding to the loop 2.3 (S51–L59), which contribute to the stimulation of ssDNA binding (17). In fact, under our current experimental conditions, both mtSSBWT- and mtSSB2,3- ssDNA complexes respond similarly to tension, which is reflected by their nearly identical FEs (Figure 1C).

These experiments were performed at tensions below 8 pN, under which no significant protein detachment is expected, which might complicate data interpretation (14,28). We observed that the replication rate of Poly on mtSSB2,3-DNA presented an entirely different force-dependence from that measured on mtSSBWT-DNA complexes (N = 57, Figure 2B): at the lowest tension (~1 pN), $V_{\text{max}}$ was similar to that measured with mtSSBWT ($V_{\text{max}} = 23 \pm 2$ nt s$^{-1}$), but it dropped rapidly (by ~30%) as tension increased to 8 pN, Figure 2B (average replication rates and processivity are shown in Supplementary Figure S5). These results would argue that loop 2.3 plays a key role in the mechanism of SSB displacement from the template by Poly.

Next, we aimed to check if the measured inhibition of Poly rate is due to a particular characteristic of the mtSSB2,3-DNA complex acquired with application of tension or, instead, is due specifically to the elimination of communication between the two proteins through the loop 2.3. In order to test this possibility, we measured the effect of EcoSSB on the force dependent kinetics of Poly. EcoSSB represents an ideal model system to test this possibility because of its high degree of sequence, structural and functional homology with mtSSBWT (3,10–14,27,28): both proteins bind to ssDNA as tetramers, with nearly identical binding and apparent dissociation constants, salt dependent binding site-sizes and, they respond similarly to tension on the ssDNA (Figure 1C and Supplementary Figure S2). However, EcoSSB does not contain the equivalent residues corresponding to the loop 2.3 characteristic of animal mitochondrial SSBs. Interestingly, these experiments showed that EcoSSB modulates the force-dependent kinetics of Poly in a manner very similar to that of mtSSB2,3 (N = 30, Figure 2B); at the lowest tension (~1 pN), $V_{\text{max}}$ is equal within error to that measured on mtSSB2,3-DNA, $V_{\text{max}} = 21 \pm 2$ nt s$^{-1}$, and decreases rapidly as force increases to ~8 pN. The characteristic velocity drop with tension, measured for both mtSSB2,3 and EcoSSB indicates that a significant conformational change induced by tension hinders the polymerase advance on DNA templates covered with SSBs lacking residues of the loop 2.3.

Overall, the data suggest that specific interactions between Poly and mtSSBWT mediated by residues of the loop 2.3 prevent the deleterious effect of tension on the replication rate, arguing that these interactions play a crucial role on the mechanism of active release of mtSSBWT from the template by Poly.

Effect of SSBs on the moving probability of Poly

In order to estimate the probability to find the polymerase in a pause or moving state during each replication condition, we calculated the moving probability, which was obtained directly from the ratio between the average velocity and $V_{\text{max}}$. Figure 2C shows that the probability of finding Poly in a moving state while replicating ssDNA was ~37% and mostly independent of tension below 8 pN. Interestingly, at the lowest tension (~1 pN) the presence of any of the three tetrameric SSBs under study did not change the moving probability, suggesting that template secondary structure is not the main source of the low moving probability (or high pause occupancy) characteristic of Poly. Intriguingly, the moving probability decreased by a ~20% further when replication was measured on ssDNA covered by any of the SSBs at tensions above 2 pN, Figure 2C. This effect suggests that the conformational change induced by tension on the SSB-DNA complexes (see below) also increased the probability of Poly being in a paused state and/or promoted additional pause state(s). However, proper identification of the type, frequency and length of pauses would be required to interrogate the nature of the high pause occupancy of the mitochondrial polymerase. We note that the method for $V_{\text{max}}$ identification presented here (‘velocity prominence’ method, Supplementary Data) did not allow us to identify pause events directly from individual replication traces.

Determination of interaction energies and conformational changes relevant for SSB displacement

We next sought to determine the magnitude of the conformational change induced by tension on the polymerase–SSB-DNA complexes and to quantify the energetics of
polymerase–SSB interactions required for ‘active’ SSB release. To this end, we sought the simplest possible model to interpret the different effects of mechanical tension on the replication kinetics of Poly on the three types of SSB–DNA templates (Supplementary Figure S7). We found that the model described above (Equation 2) explains well the force-dependent replication rate of Poly on the three SSB covered templates when two additional and contrasting factors are considered:

(i) The probability of the polymerase–SSB complex to establish functional interactions that favor SSB release from ssDNA, $P_{int}(f)$. This probability can be written as:

$$P_{int}(f) = \frac{\exp \left( -\Delta G_{int}^v/f \right)}{1 + \exp \left( -\Delta G_{int}^v/f \right)}$$  

(3)

where $\Delta G_{int}$ corresponds to the magnitude of interaction energy between the polymerase–SSB pair that facilitates the release of SSB from the template. $\Delta G_{int}$ is compromised by the magnitude of the conformational change induced by tension, $d$, which in turn, would decrease the probability of interaction between Poly and SSB.

(ii) The average energy barrier to unwrap $n$ nucleotides from SSB at each tension, $n\Delta G_{SSB}(f)$, which would hinder polymerase movement. $\Delta G_{SSB}(f)$ was calculated experimentally from the FECs of each SSB–DNA complex (‘Materials and Methods’ section and Supplementary Figure S2).

Then, $V_{max}$ can be expressed as:

$$V_{max}(f) = k(0) \exp \left[ -\delta(f) - \chi_{SSB}(f) - \chi_{ds}(f) \right]$$

$$\times \left[ P_{int}(f) + (1 - P_{int}(f)) \exp \left( -n\Delta G_{SSB}(f)/k_BT \right) \right]$$  

(4)

The free parameters, $\Delta G_{int}$, $d$ and $n$, were fixed by least squares fit of Equation (4) to experimental data, Figure 2A and B. The values obtained in each case and their corresponding errors are summarized in Table 1. We note that the other parameters of Equation (4) were determined either experimentally or were set by previous fits: $\delta = 1$ nt/step (42–45), $k(0) = 23$ nt s$^{-1}$ (fits from Equation (1)), $\chi_{SSB}(f)$, $\chi_{ds}(f)$ and $\Delta G_{SSB}(f)$ were determined from experimental FECs (Figure 1C and Supplementary Figure S2D). The results from the fits indicated that an identical tension-induced conformational change, $d \sim 2.7$ nm (Table 1), is compatible with all three situations. The main difference between them lies in the value of the interaction energy between each polymerase–SSB pair, $\Delta G_{int}$. When replication occurs along ssDNA covered with SSBs lacking residues of loop 2.3, the values of $\Delta G_{int}$ obtained from the fits were $-2.3 \pm 0.8$ k$_B$T and $-1.0 \pm 0.4$ k$_B$T for mtSSB$_{WT}$ and EcoSSB, respectively. The combination of relatively low $\Delta G_{int}$ and a relatively large conformational change has a strong effect on $P_{int}(f)$, which becomes $<<1$ rapidly with tension (Supplementary Figure S8). Consequently, as the probability of functional interactions between the two proteins decreases, the replication rate decreases concomitantly with the energy barrier imposed by the SSBs, $n\Delta G_{SSB}(f)$. In contrast, when Poly replicates ssDNA covered with mtSSB$_{WT}$, the least squares fit to data yielded $\Delta G_{int} = -12$ k$_B$T (Table 1 and Supplementary Figure S9). In this case, the magnitude of the interaction is high enough to counterbalance the effect of the conformational change induced by tension, thus ensuring, a high probability of interaction between the two proteins over a wide range of forces, $P_{int}(f) \sim 1$ from 1 to 10 pN (Figure 4 and Supplementary Figure S8). Please note that when $P_{int}(f) \sim 1$, Equation (4) equals Equation (2), and therefore, the replication rate is not hindered by the average energy barrier to unwrap n nucleotides from the SSB tetramer, $n\Delta G_{SSB}(f)$. The experimental values of $\Delta G_{SSB}$ in the absence of tension, $\Delta G_{SSB}(0)$, were $\sim 0.3$ k$_B$T for both mtSSBs and $\sim 0.18$ k$_B$T for EcoSSB (Supplementary Figure S2D), whereas the values of $n$ obtained from the fits were $\sim 2$ and $\sim 5$ nt for the mitochondrial and E. coli SSBs, respectively (see Table 1 and ‘Discussion’ section).

Effect of homologous and non-homologous SSBs on T7DNAp activity

We sought to understand further the role of catalytically relevant interactions in the mechanism of SSB release by DNA polymerases, and to explore whether the behavior observed for the mitochondrial polymerase can be extended to other DNA replication systems. To this end, we next measured the effect on the instantaneous replication rate of bacteriophage T7 DNA polymerase (T7DNAp) of homologous T7 SSB (gp2.5) and non-homologous (mtSSB$_{WT}$ and EcoSSB) SSB proteins under mechanical tension. Of note is that gp2.5 presents entirely different structural and ssDNA-binding properties than the tetrameric mitochondrial and E. coli SSBs used in this study (2). These differences are in part reflected by the FECs of gp2.5–DNA complexes (Figure 1C). First, we characterized the force dependence of the replication kinetics of T7DNAp on ssDNA under the same experimental conditions used for Poly (‘Materials and Methods section’). As reported previously (23,26,50), the replication rate without pauses ($V_{max}$) of T7DNAp was strongly dependent on tension; it increased rapidly from 1 to 6 pN, decreased toward stalling at tension >6 pN and a strong FIEA prevailed above $\sim 30$ pN (Figure 3A and Supplementary Figure S10). As in the case of Poly, the entire force–velocity behavior of T7DNAp on ssDNA was explained by the model described in Equation (1) (with $\delta = 1$), which considers the combined effect of tension and template secondary structure on the replication rate (Figure 3A). The values and corresponding errors obtained for the free variables of the model are shown in Table 1.

Next, we measured the activity of T7DNAp on ssDNA templates covered in situ by gp2.5, mtSSB$_{WT}$ or EcoSSB at increasing tensions (1–8 pN), and under the same experimental conditions used for Poly (‘Materials and Methods’ section). Interestingly, in the presence of the homologous T7 SSB (gp2.5), T7DNAp replicated at $V_{max} \sim 90$ nt s$^{-1}$ at all forces below 6–8 pN (Figure 3B), which is the maximum rate we measured for this polymerase on free-ssDNA at tensions destabilizing most of the template secondary structure (Table 1 and Figure 3A). This behavior was well explained with our model (Equation 4) when the probability of interaction between the two proteins was $P_{int}(f) \sim 1$ (Figure 3B).
Effect of homologous and non-homologous SSBs on T7DNA replication kinetics. (A) $V_{\text{max}}$ (nt s$^{-1}$) change with tension in SSB-free ssDNA ($N = 42$). Solid line represents best fit to data with Equation (1). (B) $V_{\text{max}}$ (nt s$^{-1}$) change with tension on ssDNA covered with gp2.5 ($N = 30$, pink symbols), mtSSBWT ($N = 32$, red symbols) or EcoSSB ($N = 31$, black symbols). Solid lines represent best fits of data with Equation (4): $P_{\text{int}}(f) \sim 1$ for homologous gp2.5 (magenta) and $P_{\text{int}}(f) \sim 0$ for non-homologous mtSSB (red line) and EcoSSB (black line). For comparison, dotted line shows fit to $V_{\text{max}}$ of T7DNA in SSB-free ssDNA (as shown in A). (C) Moving probability (average velocity/$V_{\text{max}}$) of T7DNA on ssDNA (blue) and on ssDNA covered with gp2.5 (magenta), mtSSBWT (red) and EcoSSB (black). For all figures, error bars show standard errors.

Figure 3. Effect of homologous and non-homologous SSBs on T7DNA replication kinetics.
unravel the secondary structure characteristic of ssDNA at low tensions (Equation (1)). As for the majority of replicative DNA polymerases, the in vitro activities of Poly and T7DNAp are hampered by the secondary structure of the template. Accordingly, $V_{\text{max}}$ increases rapidly as tension disrupts the secondary structure of the template and peaks at those tensions, $\sim 6 \, \text{pN}$, promoting the full destabilization of various possible configurations that characterize the condensed phase of the ssDNA at low forces under our experimental conditions (Supplementary Figure S3). Above $6 \, \text{pN}$, in the absence of secondary structure, the replication rate decreases with the work to convert a single- to a double-stranded nucleotide at increasing tensions. Interestingly, the results from the fits indicated that the average effective energy barrier imposed by the template secondary structure to the advance of the polymerase, $\Delta G_{\text{eff}}$, is significantly greater for T7DNAp, $\Delta G_{\text{eff}} \sim 2.6 \, k_B T$, than for Poly $\Delta G_{\text{eff}} \sim 1.4 \, k_B T$ (Table 1). This difference suggests an enhanced ability of the mitochondrial DNA polymerase to cope with the high configurational entropy characteristic ssDNA at low forces. Other alternative models have been used to explain similar force-velocity dependencies of other DNA polymerases replicating on ssDNA (23,26,53–55 and discussed in Supplementary Data). In contrast with these models, the major strengths of our model are: (i) it is based on measurable experimental observations (does not require assumptions regarding the nature of rate-limiting steps or polymerase–DNA organizations) and (ii) it fits well the entire data set with a kinetic step size of $1 \, \text{nt}$, in full agreement with previous biochemical, structural and single-molecule observations (42–45). Future single-molecule manipulation experiments, which should quantify the amount of secondary structure of the template, will help to test the efficacy of our model and of alternative models to explain the force-dependent rate of different replicative DNA polymerases.

Notably, we did not detect processive FIEA at any tension below $60 \, \text{pN}$ for Poly, which contrasts with the fast ($\sim 200 \, \text{nt} \, s^{-1}$) FIEA observed here and in previous studies for T7DNAp at forces $>30 \, \text{pN}$, Supplementary Figure S10 (26,50). We suspect that the slow exonucleolytic rate of Poly on ssDNA, which is $>10^3$ times slower than that of T7DNAp (46), either prevents FIEA to occur or deters its detection with our current resolution. In spite of its slow exonucleolytic rate, Poly presents a high intrinsic fidelity of replication, $\sim 2 \times 10^{-6}$, and very low probability of excision of a correctly base paired nucleotide ($\sim 0.14\%$) (46,47). This suggests strongly that the occurrence of single exonucleolytic events, which would be detected as pauses under our experimental conditions, are highly improbable and thus, other factors are likely responsible for the low moving probability, $3\%$, (or high pause occupancy) measured for this polymerase (Figure 2C). However, as mentioned before, a detailed identification of the type and characteristics of pause events would be required to reveal their nature.

For both DNA polymerases, the presence of SSB had a marked effect on $V_{\text{max}}$ and its response to tension (Figures 2 and 3). In the presence of their homologous SSBs, Poly and T7DNAp replicated at their maximum rates at all tensions $<8 \, \text{pN}$. In contrast, non-homologous SSBs presented a barrier for the advance of both polymerases, slowing down their $V_{\text{max}}(f)$, although in slightly different ways. Regarding the magnitude of this barrier ($n \Delta G_{\text{SSB}}(f)$), our data indicated that $n \sim 2–3$ and 5 nucleotides are unwrapped from mtSSB and EcoSSB, respectively, in each nucleotide incorporation step by Poly and T7DNAp (Table 1). These differences may suggest the existence of significant variations in the way ssDNA is unwrapped from SSBs of different origin. However, the energy to unwrap a single nucleotide from mtSSB is higher than that for EcoSSB, $\Delta G_{\text{SSB}}(0) \sim 0.3 \, k_B T$ versus $0.18 \, k_B T$, respectively (Supplementary Figure S2D; 14,27,28). Therefore, the energy barrier imposed by the mitochondrial (wild-type and variant) and E. coli SSB proteins to the advance of Poly and T7DNAp at zero tension is $n \Delta G_{\text{SSB}}(0) \sim 1 \, k_B T$ per nucleotide incorporation step.

According to our model (Equation 4), functional interactions between polymerase and SSB must be established so that the energy barrier of ssDNA unwrapping from the SSB does not slow down the replication rate. As shown in Equation (3), the probability to establish these functional interactions, $P_{\text{int}}(f)$, depends on the magnitude of the interaction energy between the polymerase and the SSB, $\Delta G_{\text{int}}$, and on the magnitude of the tension-induced conformational change in the SSB–DNA complex, $d$. The evidence for a conformational change come from the gradual decrease of $V_{\text{max}}$ with tension observed for Poly when replicating on ssDNA covered by mtSSB or EcoSSB. Importantly, a conformational change of $\sim 2.7 \, \text{nm}$ explained well all $V_{\text{max}}$ dependencies on tension measured when replication occurred along ssDNA covered by the mitochondrial and E. coli SSB proteins. This result would reflect the similarity of the ssDNA-binding properties (and their response to tension) of the tetrameric SSBs, and suggests strongly that the tension-induced conformational change mainly affects the SSB–DNA complex. According to recent single-molecule and molecular dynamics simulations studies, stretching a single ssDNA–EcoSSB complex below $8 \, \text{pN}$ aligns the SSB tetramer along the pulling coordinate and favors the unwrapping of several nucleotides from the SSB ends (28,56). For example, considering an average distance per ssDNA nucleotide within the $1–8 \, \text{pN}$ range of $\sim 0.3 \, \text{nm}/\text{nt}$, the unwrapping of $\sim 3 \, \text{nt}$ from each side of the SSB tetramer ($3 \, \text{nt} \times 0.3 \, \text{nm}/\text{nt} \times 2 = 1.8 \, \text{nm}$) plus tilting of the mtSSB tetramer along its longer axis (mtSSBWT and EcoSSB dimensions $\sim 8 \times 4 \, \text{nm}$ (10, 11)) could easily account for the $\sim 2.7 \, \text{nm}$ conformational change obtained from fits to the data. Note that this conformational change implies a reorganization of the SSB–DNA complex and does not necessarily imply a physical separation between the polymerase and the SSB by 3 nm (Figure 4). Importantly, this conformational change does not inhibit the replication rate of Poly on mtSSBWT–DNA and does not further decrease the replication rate of T7DNAp in EcoSSB- or mtSSBWT-covered ssDNA. These results argue that the conformational change induced by tension on the SSB–DNA complex does not generate an artificial barrier for polymerase progression; instead, it affects specifically the likelihood to establish functional interactions between the polymerase and the SSB relevant for SSB displacement, $P_{\text{int}}(f)$ (see below).
The main differences accounting for the particular force–velocity dependence of each polymerase–SSB complex rely on the interaction energies between the two proteins, \( \Delta G_{\text{int}} \). For Pol\(\gamma \), when replication occurs in the presence of its homologous mtSSB\(_{\text{WT}}\), \( \Delta G_{\text{int}} \sim -12 \, k_B T \). This high interaction energy ensures a high probability of functional interactions between the two proteins (Figure 4 and Supplementary Figure S8), implying a null probability to establish functional interactions, \( P_{\text{int}}(f) \sim 0 \) (Supplementary Figure S8), which could facilitate the release of SSB.

In contrast, the lower interaction energies between non-homologous polymerase–SSB couples modulated differently the observed response of \( V_{\text{max}} \) with tension. On one hand, for Pol\(\gamma \), the interaction energies with mtSSB\(_{3,3}\) and EcoSSB are about 6 to 12 times lower, \( \Delta G_{\text{int}} \sim -2 \, k_B T \) and \( -1 \, k_B T \) respectively, than that of Pol\(\gamma \) with its homologous mtSSB\(_{\text{WT}}\) (Table 1). Under these conditions, the probabilities to establish functional interactions are still significant at the lowest tensions (Supplementary Figure S8). However, the lower \( \Delta G_{\text{int}} \) values make the probabilities of interaction more sensitive to external factors, such as external tension and therefore, to decrease rapidly due to the conformational change induced by tension on the SSB–DNA complex (Figure 4 and Supplementary Figure S8). As the probabilities to establish functional interactions decrease, the replication rates are slowed down concomitantly by the effective energy barrier imposed by each SSB tetramer, \( n \cdot \Delta G_{\text{SSB}}(f) \).

Overall, our results support a mechanism in which SSB promotes the maximum replication rate of the lagging strand polymerase indirectly, by elimination of the secondary structure of the template. However, for this to occur, an interaction energy between the polymerase/SSB pair is required to overcome the energy of binding of SSB to ssDNA and for the polymerase to dislodge actively the SSB from ssDNA. At present, our data do not reveal the nature of polymerase–SSB interactions required for active SSB release. Physical interactions between mtSSB and proteins involved in DNA repair have been shown (57, 58), but there is no evidence to date of direct physical interactions between Pol\(\gamma \) and mtSSB. In contrast, T7DNAp is known to interact physically with the C-terminal tails of the homologous gp2.5 SSB and non-homologous EcoSSB (2,59). However, our data show that this interaction is not sufficient for the active displacement of EcoSSB, as this tetrameric SSB prevents T7DNAp from reaching its maximum replication rate (Figure 3B). These observations suggest that instead of (or in addition to) physical interactions, other types of interactions would modulate the communication between homologous polymerase–SSB couples, which is required for proper synthesis of the lagging strand. In consideration of
this and previous studies (59), we support a mechanism in which the nature of the interaction energy between the polymerase and the SSB required for active SSB release during lagging DNA synthesis involves repulsive electrostatic interactions. In the case of the mitochondrial DNA replication system, these may result from interactions of negatively-charged residues (S51, S54, Y57, D53, 55E and Q58) within each of the four loops 2.3 (one per monomer) that are located at the surface of mtSSBs, and the significantly net negative charge of the catalytic subunit of Poly (isoelectric point ~6.15 under our experimental conditions, pH 8.5). Interestingly, the ability of Poly to displace actively mtSSB₃₃ and EcoSSB at the lowest tension (~1 pN) suggests that additional residues, other than those of loop 2.3, play lesser, although significant, roles in the interaction between the two proteins. However, these are apparently not sufficiently robust to overcome the energy barrier to release SSB either under stress or changing physiological conditions. Based on the similarities found between the human mitochondrial and bacteriophage T7 DNA replication systems (comprising SSBs from different origin and with different structural and biochemical properties), it is tempting to speculate that other eukaryotic and prokaryotic DNA replication systems may use similar mechanisms of SSB displacement by replicative DNA polymerases during lagging strand DNA synthesis.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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