Uncoupling of Sister Replisomes during Eukaryotic DNA Replication

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SUMMARY

The duplication of eukaryotic genomes involves the replication of DNA from multiple origins of replication. In S phase, two sister replisomes assemble at each active origin, and they replicate DNA in opposite directions. Little is known about the functional relationship between sister replisomes. Some data imply that they travel away from one another and thus function independently. Alternatively, sister replisomes may form a stationary, functional unit that draws parental DNA toward itself. If this “double replisome” model is correct, a constrained DNA molecule should not undergo replication. To test this prediction, lambda DNA was stretched and immobilized at both ends within a microfluidic flow cell. Upon exposure to Xenopus egg extracts, this DNA underwent extensive replication by a single pair of diverging replisomes. The data show that there is no obligatory coupling between sister replisomes and, together with other studies, imply that genome duplication involves autonomously functioning replisomes.

INTRODUCTION

The spatial and functional relationship between the two sister replisomes (Figure 1Ai) that emanate from each origin of DNA replication is not understood (Bochman and Schwacha, 2009; Takahashi et al., 2005). In one scenario, sister replisomes move apart after initiation and function independently (Figure 1Aii). In other models, sister replisomes must remain physically coupled after origin firing to allow unwinding by the replicative helicase (Falaschi, 2000; Sciafani et al., 2004; Weisshart et al., 1999; Wessel et al., 1992) (Figure 1Aiii). If this model is correct, a constrained DNA template should not undergo replication. To differentiate between these two models, we used Xenopus egg extracts to replicate DNA that was constrained at one or both ends, and replication was visualized at the single molecule level. The data show that in this vertebrate model system, efficient replication is independent of physical coupling between sister replisomes.

RESULTS

Replication of Immobilized DNA in Xenopus Egg Extracts

Biotinylated λ phage DNA (48.5 kb) was coupled at one or both 3' ends to the streptavidin-coated, bottom surface of a microfluidic flow cell (see Figure S1A available online). To replicate these DNA molecules, we used a soluble cell-free system derived from Xenopus egg extracts (Walter et al., 1998). DNA is first exposed to a high-speed supernatant (HSS) of egg cytoplasm that supports ORC-dependent but sequence-independent recruitment of MCM2-7 complexes to DNA. A second, nucleoplasmic
extract (NPE) is then added, which supports Cdk2-dependent activation of the MCM2-7 helicase, origin unwinding, replisome assembly, and replication of the DNA.

We first examined replication kinetics of λ phage molecules immobilized at only one end, leaving the DNA template unconstrained. After coupling λ phage DNA to the surface, HSS was drawn into the flow cell and allowed to incubate for 10 min (Figure 1Bi; for details, see the Experimental Procedures). Subsequently, HSS was exchanged with NPE containing digoxigenin labeled dUTP (dig-dUTP). After a further 15 min, proteins were removed by washing the flow cell with SDS-containing buffer, fluorescein-conjugated anti-digoxigenin antibody (anti-dig) was added, and the DNA was stretched by buffer flow (Figure 1Bii). Using total internal reflection fluorescence (TIRF) microscopy, we observed colinear tracts of anti-dig, indicating that replication of the immobilized λ phage DNA had occurred (Figure 1Ci). We also imaged the DNA using SYTOX Orange ("SYTOX"), a fluorescent DNA intercalating dye that labels duplex DNA in a sequence-independent fashion. SYTOX staining revealed that the immobilized λ phage DNA contained alternating tracts of weak and strong fluorescence intensity (Figure 1Cii). The strong tracts were twice as intense as the weak tracts (Figure S1Bi, orange trace), and they colocalized with anti-dig staining (Figure 1C, compare i and ii; Figure S1Bi), suggesting the strong tracts were due to the presence of two daughter duplexes (schematically depicted in Figure 1B).

Both the anti-dig tracts and intense SYTOX tracts disappeared in the presence of Geminin, an inhibitor of MCM2-7 loading (Figures 1Ciii and 1Civ and Figure S1Bii), indicating that these two signals reflect chromosomal DNA replication. Our results demonstrate that Xenopus egg extracts can efficiently replicate DNA templates immobilized within a microfluidic flow cell, and that this process is readily detected by two independent means (see Figure S2A for additional examples).

**Replication Kinetics of Singly Tethered DNA Molecules**

To characterize replication of singly tethered molecules, we quantified several properties of the replication products (Figures 2A–2C, black bars). As shown in Figure 2A, the average number of bubbles per λ DNA was 4.54 ± 1.82, suggesting an average interorigin distance of 10.7 kb, which agrees well with previous measurements in Xenopus egg extracts (Blow et al., 2001). The lengths of the replication bubbles showed an exponential distribution (Figure 2B), implying that initiation events occurred randomly in time (Herrick et al., 2000). Most λ DNA molecules were more than 60% replicated after 15 min in NPE (Figure 2C). These observations suggest that replication of singly tethered DNA molecules is similar to what was previously observed in Xenopus egg extracts (see also below).
Stretched DNA Is Efficiently Replicated in Extracts from Multiple Origins

To test whether physical coupling between sister replisomes is required for their function, we repeated the experiment on DNA that was stretched and doubly tethered. To achieve this condition, DNA molecules biotinylated at both 3' ends were introduced into the flow cell at high flow rates. Under these conditions, DNA molecules attached to the surface of the flow cell at one end, whereupon they were instantaneously stretched by buffer flow before binding to the surface at the other end. Using this procedure, we achieved end-to-end distances corresponding to ~90% of the expected contour length of B-form DNA (Figure S2C, see the Experimental Procedures for details). Importantly, such stretched (doubly tethered) DNA molecules replicated efficiently from multiple origins (Figures 2A–2C, gray bars; Figure S2B). When doubly tethered DNA molecules were incubated in extracts supplemented with Geminin, there was no DNA replication, demonstrating that replication of constrained DNA molecules is also MCM2-7 dependent (data not shown). Importantly, there was no significant difference in the replication of singly tethered and doubly tethered molecules (Figures 2A–2C, compare black and gray bars), suggesting that no physical contact between sister replisomes is required for replication in our system.

A Single Pair of Diverging Forks Replicates Stretched DNA

Given that there was on average about 10% slack present in the doubly tethered DNA, the replication observed above could have involved many short, neighboring replicons synthesized by physically coupled replisomes. To address this caveat, we examined whether a single pair of diverging sister forks can replicate stretched DNA to an extent larger than the slack originally present. To ensure that only a single pair of replisomes was activated on each DNA molecule, we used p27Kip, a Cdk2 inhibitor that blocks new initiations but does not affect elongation (Walter and Newport, 2000). Thus, 2–5 min after replication was initiated with NPE, we flowed in fresh NPE containing p27Kip (Figure 3A). Under these conditions, the majority of DNA molecules exhibited one or no replication bubbles (Figure 3C). To verify that the observed bubbles were produced by two diverging replication forks, we supplied dig-dUTP 15 min after the initial NPE addition and allowed replication to proceed for a further 25 min (Figure 3A).

In the replicated molecules, two tracts of dig-dUTP were visible whose outer edges coincided with the boundaries of the intense SYTOX tract, as expected for bidirectional replication from a single origin (Figure 3B and Figure S3A). Figure 3D shows that the extent of replication on every DNA molecule examined was much greater than the slack present in the substrate. For a single pair of physically coupled replisomes to produce such large replication bubbles, the molecule would have to be stretched well beyond the contour length of B-form DNA. Because the force required to overstretch DNA to such an extent (van Mameren et al., 2009) is larger than any reported for individual DNA motors (Smith et al., 2001), and because such overstretching would almost entirely denature the DNA substrate (van Mameren et al., 2009), it is very unlikely that sister replisomes remained connected during replication in this system. We conclude that sister replisomes can function independently on immobilized DNA molecules.

Uncoupling of Sister Replisomes Does Not Affect Fork Rates

To test whether the uncoupling of sister replisomes affects their replication dynamics, we compared fork rates on stretched,
doubly tethered and relaxed, singly tethered DNAs that had undergone a single initiation event (Figure S3). Dividing the lengths of the anti-dig tracts by the duration of the dig-dUTP pulse yielded a mean fork rate of 267 ± 160 bp/min for stretched DNA (n = 91; Figure 4A, gray bars) and 268 ± 161 bp/min for relaxed DNA (n = 98; Figure 4A, black bars). The presence of dig-dUTP did not affect the rate of DNA synthesis (Figure S3C). The measured rates were close to the lower estimates of fork rates in conventional, nuclear assembly Xenopus egg extracts (Lu et al., 1998; Mahbubani et al., 1992). Thus, replisomes on constrained and unconstrained molecules move at the same rates, demonstrating that replisome uncoupling does not adversely affect replication fork progression.

Correlation between Rates of Sister Forks on Singly Tethered but Not Doubly Tethered DNA

Previous studies in different experimental systems showed varying but significant degrees of correlation between the rates at which sister replication forks progress (Conti et al., 2007; Dubey and Raman, 1987; Tapper and Depamphilis, 1980). We looked for a correlation between progression of sister forks in our system. Figure 4B plots the length of the left versus right anti-dig tracts within single replication bubbles on singly and doubly tethered DNA. On singly tethered DNA, there was a weak, positive correlation between the rates at which the two sister replisomes moved (Figure 4B, black squares, n = 48; R = 0.26, p = 0.07). Since uncoupling of sister replisomes did not affect fork progression in our system, the correlation between sister forks in relaxed DNA is unlikely to be related to a functional interaction between sisters. Consistent with this, termination of one replication fork by a double-strand DNA break in yeast does not affect progression of the sister fork (Doksani et al., 2009). Therefore, correlations that we and others observed likely represent chromatin microenvironments that result in similar activity of nearby replisomes (Conti et al., 2007), perhaps due to similar concentrations of key replication factors. Consistent with this idea, sister replisomes moving on stretched DNA, which are separated in space, showed no correlation (Figure 4B, gray squares, n = 45; R = −0.1, p = 0.47).

DISCUSSION

It has been proposed that sister replisomes function as an obligatory dimeric complex (Falaschi, 2000; Kitamura et al., 2006; Ligasová et al., 2009; Sclafani et al., 2004). However, our data demonstrate that no physical association is required between sister replisomes on λ DNA replicating in Xenopus egg extracts, suggesting that replisomes can function independently during vertebrate DNA replication. Together with previous results which failed to find evidence of MCM2-7 double hexamers in S phase using coIP approaches (Gambus et al., 2006), and recent experiments using purified MCM2-7 holocomplexes (Ilves et al., 2010; Moyer et al., 2006), our data suggest that sister MCM2-7 helicases (and replisomes) normally uncouple upon activation, as seen in bacteria (Reyes-Lamothe et al., 2008).
associate through their N termini (Valle et al., 2000). In addition, that T-ag loads onto the SV40 origin as two hexamers, which by the analysis of SV40 T-ag. Electron microscopy showed 2002; Weisshart et al., 1999). Taken together, these studies unwinding activity than single hexamers (Alexandrov et al., 1992). Moreover, muta-

during T-ag-mediated DNA unwinding, a fraction of DNAs adopt a ``rabbit ear'' conformation, in which two loops of single-stranded DNA stick to the surface, the flow cell was incubated with blocking buffer (20 mM Tris [pH 7.5], 50 mM NaCl, 2 mM EDTA, 0.2 mg/ml BSA) for at least 15 min.

to attach λ DNA (New England Biolabs) to the streptavidin coated surface at one end, the single-stranded 5' tails of λ DNA were annealed and ligated to complementary oligonucleotides 5'-AGGTGCGGCCC-Biotin-3' and 5'-GGGCGGCGGACCT-3' (Integrated DNA Technologies). For double tethering, both oligos contained biotin at the 3' end. Biotinylated λ DNA (15–50 pM) in blocking buffer was injected into the flow cell at a constant rate of 20–100 μl/min. At a flow rate of 50–100 μl/min, λ DNA biotinylated at both ends was stretched to 70–80% of its contour length (16.5 μm) (Figure S2O–S2U). To stretch DNA further, we used chloroquine, which intercalates into and extends the pitch of dsDNA (Cohen and Yielding, 1965). When injected at 100 μl/min in the presence of 100 μM chloroquine in blocking buffer, end-to-end distance of doubly tethered λ DNAs was 85%–95% of its contour length (Figure S2Oc). After DNA injection and before addition of extract, chloroquine was removed by extensive washing of the flow cell with blocking buffer (5 min at 100 μl/min).

to limit our analysis to DNA molecules that remained doubly tethered during the entire replication reaction, we used a reduced flow rate (25 μl/min) for all buffer exchanges following replication. In this way, even if a DNA molecule that detached from one end in extracts became doubly tethered during subsequent washes, it would be stretched to a much lesser extent than molecules that remained stretched throughout the experiment. Thus, λ DNA stretched to 85%–95% of its contour length at the end of the experiment must have stayed doubly tethered during replication. Therefore, we analyzed only those molecules that were stretched to 85% or more as doubly tethered.

**Replication of Immobilized DNA**

In a separate line of investigation, we recently discovered that DNA replication in Xenopus egg extracts requires a minimum threshold concentration of DNA (~1 ng/μl in HSS and NPE (Lebofsky et al., 2010)). Since the effective concentration of λ DNA immobilized in the flow cell was extremely low, we supplemented HSS (Walter et al., 1998) and NPE (Walter et al., 1998) with “carrier” plasmid to raise the overall DNA concentration to levels that are compatible with DNA replication. Thus, after immobilizing λ DNA on the functionalized surface, HSS containing carrier plasmid (5–10 ng/μl of pBlueScript II KS [−]) was injected at 10 μl/min for 2 min and further incubated for 8 min without flow. Next, a 2:1 mixture of NPE and HSS supplemented with 5–10 ng/μl of pBS (“replication extract”) was flowed in at 10 μl/min for 80 s, followed by incubation for different lengths of time without flow, as indicated. dIg-dUTP (Roche Inc.) (7 μM) was also included in the replication extract for labeling
of replicated regions. All reactions were carried out at room temperature (22°C).

To observe bidirectional replication involving single initiation events, immobi-

lized λ DNA was incubated with HSS/carrier plasmid and subsequently re-

placed with replication extract (lacking dig-dUTP). After the time specified, a second replication extract containing 66 μg/ml p27THF was injected and incubated further. Finally, a replication extract containing 66 μg/ml p27THF and 7 μM dig-dUTP was introduced. In each case, replication extract was injected at 10 μl/min for 80 s.

To stop the replication reaction, the flow cell was washed with SDS buffer (20 mM Tris [pH 7.5], 50 mM NaCl, 12 mM EDTA, 0.1% SDS) for 10 min at 25 μl/min. To label dig-dUTP, anti-digoxigenin-fluorescein antibody (Roche Inc.) was diluted to 0.4 μg/ml with buffer (10 mM HEPES [pH 7.7], 2.5 mM MgCl2, 50 mM KCl, 0.2 mg/ml BSA) was drawn into the flow cell for 20 min at 25 μl/min. Excess antibody was removed by washing the flow cell with blocking buffer. Finally, blocking buffer containing 15 nM SYTOX Orange (Invitrogen) was introduced to fluorescently label dsDNA.

Use of Oligonucleotides as Carrier DNA

As described above, DNA replication of tethered λ DNA in the flow cell was carried out using egg extracts (HSS and NPE) that were supplemented with high concentrations of a carrier plasmid. Like the immobilized λ DNA, the free carrier plasmid is also expected to undergo replication in the flow cell. To ensure that the activity of replicosomes on the immobilized λ DNA was not dependent on interactions with replicosomes on the carrier plasmid, we replaced the carrier plasmid with a short, double-stranded oligonucleotide (29 bp; Lebofsky et al., 2010) that does not support loading of MCM2-7 helicase due to its small size and therefore cannot undergo replication (Edwards et al., 2002). We have shown that such a nuclease is able to replace carrier plasmid to promote licensing in HSS (Lebofsky et al., 2010).

Doubly tethered λ DNA was licensed with HSS containing 10 ng/μl of the oligo duplex. Subsequently, standard replication extract (NPE/HSS/carrier plasmid) was injected. Importantly, the carrier plasmid supplied with the replication extracts did not undergo DNA replication because it was added to a mixture of HSS and NPE, and NPE contains very high concentrations of Geminin, which block DNA replication (Figure S4A) (Walter et al., 1998). Thus, in this sequence, none of the added carrier DNA (oligo or plasmid) underwent DNA replication, yet single initiations on many stretched λ DNA molecules still produced bubbles larger than the slack present on stretched λ DNA (Figure S4B). These observations confirm that replication of carrier plasmid does not mediate replication of immobilized λ DNA molecules.

TIRF Microscopy

Immobilized λ DNA molecules were imaged on an objective-type TIRF config-

uration using an inverted microscope (IX-71; Olympus) equipped with a 60× oil objective (PlanApo, N.A. = 1.45; Olympus) and a 1.3× magnification unit. A multiwavelength Ar-Kr ion laser (Innova 70C-Spectrum, Coherent Inc.) was used for illumination. SYTOX Orange and fluorescein were excited with 568 nm and 488 nm laser light, respectively, using varying intensities and 100 ms exposures per frame. Images were acquired using an Andor iXon back-

illuminated electron-multiplying CCD camera (Andor Technology) at 2 Hz.

Single tethered molecules were imaged at a flow rate of 100–125 μl/min (for stretching), while doubly tethered DNAs were imaged in the absence of flow since they were already stretched.

Image Processing

To improve the signal-to-noise ratio on fluorescence images of replicated λ DNA, multiple (5–30) consecutive images of SYTOX and fluorescein were averaged separately using ImageJ and merged using Adobe Photoshop. The end-to-end distance of each DNA molecule was measured via the SYTOX image, and the size of a replication bubble was determined using the SYTOX or the fluorescein signal.

SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures and Supplemental References and can be found with this article at doi:10.1016/j.molcel.2010.11.027.


