Real-Time Single-Molecule Observation of Rolling-Circle DNA Replication

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ABSTRACT

We present a simple technique for visualizing replication of individual DNA molecules in real time. By attaching a rolling-circle substrate to a TIRF microscope-mounted flow chamber, we are able to monitor the progression of single-DNA synthesis events and accurately measure rates and processivities of single T7 and *Escherichia coli* replisomes as they replicate DNA. This method allows for rapid and precise characterization of the kinetics of DNA synthesis and the effects of replication inhibitors.

INTRODUCTION

DNA replication is a fundamental biological process that requires the coordinated activities of a large number of enzymes organized in a multiprotein assembly termed the replisome. Using a variety of systems and approaches, researchers aim to understand how the various proteins work together and how their interactions affect replication activities (1–4). Traditionally, biochemical DNA replication assays rely on agarose gel electrophoresis and/or the incorporation of radioactive nucleotides to measure rates of DNA synthesis and processivity of replication events. From a practical standpoint, these techniques involve multiple time-consuming steps as well as the use of hazardous materials. Furthermore, these techniques average over a large ensemble of molecules, making it inherently difficult to obtain distributions of enzymatic rates and processivities, essential data for complete characterization of replisomal activities. Accurate determination of processivity typically involves either trapping or dilution techniques, both of which can inadvertently affect reaction concentrations and dissociation kinetics. Also, rare events and short-lived intermediates are not easily observed in bulk-phase techniques, obscuring some of the dynamic interactions and states that occur during DNA replication.

Information on rate, processivity and short-lived intermediate states can be obtained by observing individual replisomes at the single-molecule level. In recent years, numerous single-molecule techniques have been developed to characterize the activity of nucleic-acid enzymes such as DNA polymerases and helicases (5–9). Most studies have relied on the mechanical manipulation of individual proteins or on the imaging of fluorescent tags to report on the catalytic activity of replication proteins. Although quite powerful, these techniques are generally limited to observing a single-DNA substrate at a time, making it difficult to accumulate statistics for larger multiprotein complexes. Furthermore, the experimental setups can be prohibitively complex and expensive for the general user. Recently, we have demonstrated a multiplexed single-molecule replication assay based on flow-stretching of tethered bacteriophage λ DNA and used the technique to characterize helicase–primase and helicase–polymerase interactions in leading-strand synthesis (10–12). In these experiments, the difference in extension of single-stranded (ss) and double-stranded (ds) DNA at low pN forces is exploited to observe an enzyme acting on DNA. The requirement for a net conversion between ss and dsDNA makes it challenging to apply this method to coordinated replication where both DNA strands at the fork are copied. Using replication loops as an observable, we have recently used this flow-stretching technique to study the processes controlling the dynamics of loop formation at the T7 replication fork (13). Nevertheless, these experiments present only an indirect readout of replication-fork dynamics.

Here, we present a simple single-molecule assay for measuring coordinated DNA replication by individual replisomes in real time. We employ the rolling-circle DNA amplification scheme, as it facilitates highly processive DNA synthesis. Rolling-circle replication has been proven a useful method for monitoring real-time synthesis in bulk-phase assays (14–16) and at the single-molecule level to characterize telomere mimic sequence ssDNA (17). Briefly, we couple the 5’-end of the lagging strand...
of a rolling-circle substrate to the surface of a flow chamber and introduce replication components into the flow cell to initiate DNA synthesis. By applying a constant, laminar flow through the chamber we hydrodynamically stretch the growing DNA. Application of a low concentration of intercalating stain during the reaction allows us to directly image the time-dependent length of dozens of growing DNA molecules in real time. As a proof of principle, we characterize fully reconstituted replisomes from two systems, bacteriophage T7 and Escherichia coli, some of the largest protein complexes studied to date at the single-molecule level. Bacterial and viral DNA replication are obvious targets for therapeutic inhibition, and this assay readily lends itself to studying replication inhibitors, a principle demonstrated by examining the effects of a dideoxynucleotide triphosphate on the rate and processivity of T7 DNA replication.

MATERIALS AND METHODS

Rolling-circle substrate preparation

Substrate was prepared as described previously (18). A 66-mer tail oligonucleotide 5’-Biotin-TGATC-3’ (Integrated DNA Technologies) was annealed to M13mp18 ssDNA (New England Biolabs) and extended with 64 nM T7 polymerase in reaction buffer (see below) at 37°C for 12 min. The reaction was quenched with 100 mM EDTA, cooled, and the filled M13 extracted via phenol–chloroform extraction.

Coverslip surface preparation

To minimize non-specific interactions between the glass surface and proteins, we covalently couple high-molecular-weight polyethylene glycol (PEG) to the surface, including a small fraction of biotin moieties for tethering of our DNA substrates. First, the glass is coupled to the alkoxy group of an aminosilane, creating a surface with reactive amine groups that can subsequently be coated with a polymer of choice (19). By coupling a mixed population of biotinylated and non-biotinylated succinimidyl propionate-PEG to the amine-functionalized glass, we coat the coverslip in a layer of PEG displaying a mixture of biotin and non-reactive methyl groups. The biotinylated coverslip is incubated with a 1 mg/ml solution of streptavidin (Sigma) in PBS (pH 7.5) immediately before use in the experiment.

Flow-cell construction

Chamber construction was done as described previously (12). Briefly, a 2-mm channel was cut out of double-sided adhesive tape (Grace BioLabs) and attached to a quartz slide with pre-drilled holes and a functionalized coverslip (12,19). Polyethylene tubes (0.76 mm inner diameter, 1.22 mm outer diameter; Becton Dickinson) were inserted into the holes, and the entire chamber was sealed with quick-dry epoxy. The flow cell was incubated with 20 mM Tris pH 7.5, 2 mM EDTA, 50 mM NaCl, 200 µg/ml bovine serum albumin (BSA) and 0.005% Tween to block the surface and further prevent non-specific interactions.

TIRF microscope setup

The flow cell was placed on an Olympus IX-71 inverted microscope and connected to a syringe pump (Harvard Apparatus) used to control the flow of buffer (20). An airspring was inserted between the flow cell and pump to dampen flow fluctuations. SYTOX Orange (Invitrogen), present in the replication buffer was excited through a 60× oil-immersion TIRF objective (Olympus, NA = 1.45) with a 532 nm solid-state laser (Coherent Compass 215M-75). The resulting fluorescence was collected back through the objective, passed through an emission filter (Chroma HQ600/75 m) to eliminate residual laser light and recorded on an EM–CCD (Hamamatsu).

Replication reactions

Substrate DNA was flowed into the chamber at 25 pM for 30 min at 0.025 ml/min and excess DNA was removed via washing with buffer. For reactions performed at 37°C, a home-built aluminum block was affixed to the top quartz surface with high thermal conductivity paste and heated with a cartridge heater (Omega) controlled by variable power supply (Elenco). By measuring the buffer temperature in the flow chamber we were able to control the cartridge heater to achieve 37°C at the observed region.

T7 reaction

Replication reactions were performed and proteins purified as described previously (13,21,22). Reaction buffer contained 40 mM Tris pH 7.5, 50 mM potassium glutamate, 10 mM magnesium chloride (MgCl2), 100 µg/ml BSA with 5 mM dithiorthiol (DTT), 600 µM deoxyribonucleotides (dNTPs, New England Biolabs), 300 µM ATP (Sigma), 300 µM CTP and 15 nM SYTOX Orange added immediately before beginning the reaction. Proteins were added as: 5 nM gp4 (hexameric), 40 nM polymerase (1:1 gp5: thioredoxin) and 360 nM gp2.5. DideoxyGTP (ddGTP, Roche) was added at indicated concentrations immediately before starting the reactions.

E. coli reaction

DnaB helicase, DnaC helicase loader, DnaG primase, αβ polymerase, β processivity clamp and the τψϕϕrψϕψ and τψϕϕrψϕψ clamp loader assemblies were purified as described (12). PriA, PriB, DnaT and SSB were prepared from overproducing strains by methods similar to those described by Marians (23). Replication reactions were performed as described previously (12,24). Reaction buffer contained 50 mM HEPES pH 7.9, 12 mM magnesium acetate (MgOAc2), 80 mM potassium chloride (KCl) and 100 µg/ml BSA with 10 mM DTT, 40 µM dNTPs, 200 µM UTP, GTP and CTP, 1 mM ATP and 15 nM SYTOX Orange added before the reaction. Proteins were added as: 30 nM DnaB (hexameric), 180 nM DnaC (monomeric), 30 nM αβ, 15 nM τψϕϕrψϕψ or τψϕϕrψϕψ, 30 nM β (dimeric), 300 nM DnaG, 250 nM SSB (tetrameric), 20 nM PriA, 40 nM PriB and 480 nM DnaT.
Data analysis

To obtain length trajectories and minimize the contribution from transverse Brownian fluctuations of the DNA, we calculated intensity projections by summing over a narrow rectangular box of pixels (~2 μm wide) along the length of the DNA. The end of the DNA was defined by an intensity threshold set to 65% of the maximum intensity. Linear interpolation around the threshold determined the DNA end position to sub-pixel resolution. Rate was determined by linear regression of DNA length plotted versus time. Processivity was determined by deconvolving the length of the M13 DNA substrate from the final length of the extended DNA and converted to basepairs by determining the known length of tethered λ-phage DNA (48.5 kb) with our camera and using the calibrated pixel size in basepairs (here, 1 pixel = 785 bp).

RESULTS AND DISCUSSION

Experimental outline

A single-stranded, 7.2-kb circular M13mp18 DNA is primed and its complementary strand synthesized to form dsDNA. Upon completion of synthesis of the entire circular substrate, the polymerase acting at the 3′-end will encounter the 5′-end of the original primer and synthesis will continue by displacing the previously synthesized DNA as ssDNA (Figure 1a). In the presence of protein activities required for priming and other lagging-strand processes, the displaced ssDNA tail will be effectively converted into dsDNA by repetitive priming and synthesis of lagging-strand Okazaki fragments. To allow surface immobilization and single-molecule observation of the replication substrates, we used a 5′-biotinylated ‘tail’ primer in constructing the M13 rolling-circle substrate. This substrate is purified and introduced into a flow chamber constructed with a biotin–streptavidin-functionalized coverslip (12). The filled, surface-attached M13 can serve as a replication template upon addition of nucleotides and proteins. Flowing low picomolar concentrations of M13 DNA into the chamber results in hundreds of DNA molecules in a single field of view (125 μm × 125 μm), each of which can serve as a substrate for the introduced proteins (Figure 1b).

As the replication reaction proceeds, the DNA attaching the M13 circle to the surface is extended and stretched fully by hydrodynamic flow of buffer. We are able to visualize individual dsDNA molecules through total-internal reflection fluorescence (TIRF) microscopy by adding SYTOX Orange dsDNA intercalating stain to the reaction buffer, allowing for observation of coupled leading- and lagging-strand synthesis as a processively lengthening, stained dsDNA molecule. Addition of the necessary replication proteins resulted in the lengthening of the M13 template indicating processive DNA synthesis (see Supplementary Data). Trajectories of DNA length as a function of time reported on rates of DNA synthesis (Figure 2), and total product length at the end of synthesis was used to measure processivity.

T7 DNA replication

Our single-molecule rolling-circle assay is broadly applicable to general DNA replication. We examined two common model systems of DNA replication, the bacterial phage T7 and E. coli replisomes. After binding our rolling-circle substrate to the surface of the chamber, we introduced the replisomal proteins in replication buffer containing the intercalating dye SYTOX Orange and observed replication as the solution entered the field of view. The T7 replisome can be reconstituted with four proteins: gp4, helicase/primase; gp5 + thioredoxin, polymerase; gp2.5, ssDNA-binding protein (1,13,21). At room temperature (22 °C), we observed T7 replication at 75.9 ± 4.8 bp s⁻¹ with an average processivity of 25.3 ± 1.7 kbp (Figure 3). Rates of synthesis were consistent with the previously reported value of 80 bp s⁻¹ at room temperature (13).

E. coli DNA replication

The E. coli replisome is significantly more complex (1,2) with 13 core proteins (DnaB, helicase; DnaG, primase; βθ, polymerase; β, processivity clamp; τδ♀δ’γψ or τδδ’γψ, clamp loader; SSB, ssDNA-binding protein) and four additional proteins, DnaC, PriA, PriB and DnaT to assist helicase loading on the 5′-blocked DNA.
substrate (24). Replication by the *E. coli* replisome at 37°C was observed at 535.5 ± 39 bp s⁻¹, consistent with the previously reported 600 bp s⁻¹ (25–27), with an average processivity of 85.3 ± 6.1 kbp (Figure 3). Replication products with lengths of 350 kbp or greater were observed, demonstrating the high processivity of the *E. coli* replisome. The dramatic increase in rate and processivity seen with the *E. coli* as compared to T7 proteins was not merely due to temperature: increasing the temperature of the T7 reaction to 37°C increased the rate to 141.5 ± 8.9 bp s⁻¹ and the processivity to 51.6 ± 9.4 kbp.

Several lines of evidence lead us to conclude that we are observing true coordinated replication. As our assay measures coordinated DNA synthesis by observing replication of the lagging strand, omitting components required for coordination (DnaG primase or χ/ψ using different clamp loader complexes) results in no observable replication. Leading-strand synthesis alone produces no net gain of dsDNA as the circle ‘rolls’, and neither T7 gp5 + thioredoxin or *E. coli* αxθ is capable of strand invasion synthesis by polymerase activity alone, thus any aberrant or uncoupled replication is not observed in our assay, which visualizes exclusively dsDNA. These facts combined with rate values very similar to those published support our conclusion that each observed replication product is from a single, complete replisome.

**Effects of SYTOX and flow stretching DNA**

SYTOX Orange is an intercalating agent and can photodamage DNA (28), making it necessary to determine if its inclusion in our assay affects replication. Based on several control experiments, we conclude that for the systems we examined, SYTOX has no effect on replication. As noted above, rates of DNA synthesis for both T7 and *E. coli* are consistent with previous measures. Bulk-phase rolling-circle assays we performed showed no dependence on the presence or absence of SYTOX in the replication buffer. In other single-molecule experiments, SYTOX has been shown to have no effect on the size of T7 DNA replication loops or rates of synthesis (13). Performing the single-molecule rolling-circle experiment without SYTOX and staining after the reaction was finished yielded identical processivity distributions (*E. coli*, 85.6 ± 9.5 kbp). Laser excitation had no effect on
DNA synthesis, as products from reactions not illuminated until the end of the experiment displayed identical length distributions as those observed in real time (E. coli, 90.1 ± 13.6 kbp). Also, the force acting on the replication proteins themselves is vanishingly small, as they are located at the circle near the distal end of the DNA and experience effectively no force when stretched (29). Confirming this assumption, we performed the E. coli replication reaction in a tube, stopped the reaction by addition of 100 mM EDTA after 20 min and flowed the mixture into the flow cell. Introduction of SYTOX allowed visualization of the force-free reaction products with consistent length distribution (78.2 ± 11.8 kbp).

Analyzing replication inhibitors

The search for novel antibacterial and antiviral agents remains an important field of research, and DNA replication is an obvious target for pharmacological investigation (30,31). The ease of performing our experiment and the availability of automated fluorescence imaging systems make the assay reported here an ideal method for screening and characterizing replication inhibitors. We demonstrated this concept by measuring rates and processivities of the T7 replication reaction with varying amounts of deoxyGTP (ddGTP), a chain-terminating nucleotide that can be removed by the exonuclease activity of gp5 (32). Obviously, the incorporation of a single ddGTP depends upon concentration, and each inclusion requires the polymerase to switch between polymerase and exonuclease activities. The relative portion of time needed to excise the deoxynucleotides increases with the ddGTP concentration and results in a decrease in aggregate rate of DNA synthesis. We observed a rate decrease from 75.9 ± 4.8 bp s⁻¹ in the absence of ddGTP to 49.2 ± 3.1 bp s⁻¹ at 1 μM ddGTP (Figure 4a). In addition, product length decreased from 25.3 ± 1.7 kbp without ddGTP to 4.8 ± 0.5 kbp at 1 μM ddGTP (Figure 4b). It is interesting to note that the decrease in rate and product length scale differently with ddGTP concentration. Product length decreases by ~5-fold over the concentration range compared to ~1.5-fold for rate. These results suggest that each ddGTP incorporation increases the probability of replisome dissociation (‘off’ rate), leading to a decrease in total replication processivity. Translating our assay to larger-scale screening or testing would present a simple method for easily analyzed, multiplexed studies of the effects of small molecules on replisomal DNA replication.

CONCLUSIONS

Our assay allows for real-time readout of replication rate and processivity from a single active replisome while increasing dramatically the ease and speed of performing replication experiments with no need for radioactive or hazardous materials. By nature, the rolling-circle method promotes extensive synthesis limited only by the processivity of the replisome rather than the length of the template. This allows us to observe DNA replication up to several hundred kilobasepairs, measuring real values for replisome processivity, numbers difficult to measure precisely with bulk-phase methods due to the exclusion limitations of 50 kbp or fewer in agarose gel electrophoresis. Wide-field imaging allows us to observe hundreds of tethered M13 substrates and study the behaviors of their associated replisomes in a single field of view. Thus, even with modest yields of correct replisomal assembly and activity we are able to construct statistically meaningful (N > 50) rate and processivity histograms from a single 20 min experiment. Components of the assay are made using published methods and can be produced readily in large or reusable quantities. Extension of the assay using multichannel microfluidic devices will allow us to perform a high number of reactions in a short time, a necessary criterion for screening and characterizing the effect of replication inhibitors or other biochemical perturbations. The simplicity of the experiment along with the ability to observe single active replisomes in real time makes our technique a powerful and tractable method for studying the many elements of DNA replication.
SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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Conflict of interest statement. None declared.

REFERENCES