Multiplexed single-molecule flow-stretching bead assay for DNA enzymology

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Single-molecule techniques have been used successfully to visualize real-time enzymatic activities, revealing transient complex properties and heterogeneity of various biological events. Especially, conventional force spectroscopy including optical tweezers and magnetic tweezers has been widely used to monitor change in DNA length by enzymes with high spatiotemporal resolutions of ∼nanometers and ∼milliseconds. However, DNA metabolism results from coordination of a number of components during the processes, requiring efficient monitoring of a complex of proteins catalyzing DNA substrates. In this mini-review, we will introduce a simple and multiplexed single-molecule assay to detect DNA substrates catalyzed by enzymes with high-throughput data collection. We conclude with a perspective of possible directions that enhance capability of the assay to reveal complex biological events with higher resolution. [BMB Reports 2019; 52(10): 589-594]

MULTIPLEXED SINGLE-MOLECULE FORCE SPECTROSCOPY

Single-molecule studies have extensively explored a variety of biological problems by following individual molecules going through their biochemical or biophysical reactions (1-3). Single-molecule assays have provided real-time snap shots of molecular events, resolving molecular sub-populations, transient intermediate states, rare molecular events, and non-uniform kinetics in the enzymatic reaction (4). In contrast to direct observation of probe-labeled enzymes on DNA using fluorescence microscopy (5), the most commonly used force spectroscopy, including optical tweezers (OT), magnetic tweezers (MT), and atomic force microscopy (AFM), can manipulate individual biomolecules by external forces in the range of 0.01-10,000 pN to unravel structural and mechanical properties (6). The force spectroscopy has been also used to precisely measure displacement as a function of force with high spatiotemporal resolution of sub-nanometers and sub-milliseconds to detect enzyme activities on substrates in DNA replication, DNA repair, transcription, and translation (7-11).

However, traditional force spectroscopy is difficult to parallelize measurement at one time, limiting monitoring of enzymatic activity by multi-proteins due to inefficient in vitro reconstitution of the biological system of interest (12). To achieve high-throughput data collection, optical tweezers and magnetic tweezers have been modified to efficiently collect data in parallel. Although OT uses a focused laser beam to trap a polystyrene/silica bead linked to DNA, holographic light using a spatial light modulator can trap ~100 beads per field of view (13) (Fig. 1A). However, this multiplexed OT is complex and needs professional knowledge and high cost to build and operate. MT uses a magnetic force applied to a superparamagnetic bead linked to DNA molecules by a magnet to manipulate bead-DNA immobilized on the surface. To increase the number of DNA-tethered beads, an array of DNA molecules linked to beads is immobilized on the patterned surface of the imaging chamber (~300 DNA-tethered beads per field of view) (Fig. 1B) (14, 15). Recently, another high-throughput force spectroscopy, called acoustic force spectroscopy (AFS), was developed, which can detect tens-thousands of DNA molecules per field of view depending on the magnification of an objective lens (16). The AFS uses acoustic waves to trap polystyrene microspheres attached to surface tethered-DNA molecules, resulting in the extension of multiple DNA molecules immobilized on the surface in an upward direction to the surface (Fig. 1C). These multiplexed MT and AFS require complex instruments and a calibration profile of diffraction pattern images of beads for measurement of bead displacement due to the vertical motion of trapped beads to the imaging plane.

In this mini review, we describe a simple, robust, low cost, and multiplexed single-molecule force spectroscopy to study processive enzyme activities on stretched-DNA substrates.
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Fig. 1. Multiplexed single-molecule force spectroscopy. (A) Holographic optical tweezers. (B) Multiplexed magnetic tweezers. (C) Acoustic force spectroscopy.

Fig. 2. Schematic representation of single-molecule flow-stretching bead assay. (A) A setup based on a conventional optical microscope. A buffer solution including target proteins flows through the flow chamber with constant rate by a syringe pump. (B) ~300 beads (2.8 µm in diameter) attached to tethered-DNA molecules on the surface under 10x magnification objective. (C) Bead-DNA is immobilized on the streptavidin-coated surface that is passivated with biotin-PEG/PEG (1:100). A laminar flow stretched bead-DNA. (D) Intensity profile of a bead image shows a Gaussian distribution.

using a hydrodynamic force, called flow-stretching bead assay.

SINGLE-MOLECULE FLOW-STRETCHING BEAD ASSAY (smFS)

Overview of smFS
To monitor individual DNA in the smFS, DNA molecules are immobilized on the surface passivated with biotin-PEG (polyethylene glycol) via biotin-avidin interactions while the opposite end with digoxigenin is attached to a super-paramagnetic bead (2.8 µm in diameter) functionalized with anti-digoxigenin antibody. A steady buffer flow with a constant rate given by a syringe pump drags the bead linked to the tethered-DNA, resulting in extension of DNA (Fig. 2A). A
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Flow-stretching bead assay using different elastic properties between dsDNA and ssDNA. (A) Force-extension curve for dsDNA and ssDNA. (B) Transition of dsDNA (ssDNA) to ssDNA (dsDNA) is shortening (lengthening). (C) The scheme of the length measurement of dsDNA and ssDNA at a given force.

Fig. 3. Flow-stretching bead assay using different elastic properties between dsDNA and ssDNA. (A) Force-extension curve for dsDNA and ssDNA. (B) Transition of dsDNA (ssDNA) to ssDNA (dsDNA) is shortening (lengthening). (C) The scheme of the length measurement of dsDNA and ssDNA at a given force.

damper made by a Falcon centrifuge tube that is filled with water and an air layer is installed between the syringe pump and flow chamber to filter high-frequency noises coming from mechanical fluctuation of the syringe pump (Fig. 2A). The position of beads is monitored to measure enzymatic activities on DNA substrates. Beads linked to immobilized-DNA molecules are visualized under a standard optical microscope and recorded by a charge coupled device (CCD) camera (Fig. 2A). Dark spots in the resulting image correspond to the beads (Fig. 2B). Optimal number of beads linked to DNA is \( \sim 300 \) per field of view under 10x objective. In the ssFS, we can monitor enzyme activities on DNA substrates by measuring changes in the length of individual DNA molecules by imaging beads and tracking their position.

Flow chamber construction
A flow chamber was constructed with a biotin-PEG functionalized cover slip, PEG-coated glass slide, double-sided tape, and tubing (Fig. 2C). A pair of inlet/outlet holes is drilled on a glass slide before it is passivated with PEG. A 100 \( \mu \)m thick double-sided tape is cut for a channel with 3 mm in width and 25 mm in length. Width and height of the channel must be maintained to perform experiments under an identical force once the external force is calibrated to flow rate by the syringe pump. A solution of streptavidin or neutravidin is evenly spread on the biotin-PEG cover slip and then washed with deionized water after 10 minutes of incubation. A chamber is made by putting the prepared cover slip and glass slide together with adhesive double-sided tape and then sealing with epoxy. Tubing is inserted into each inlet/outlet hole and then epoxy is applied to the junction between the hole and tubing. The flow chamber filled with a blocking buffer is ready to do a single-molecule flow-stretching experiment. For more detailed information on construction of the flow chamber, we refer to a demonstration movie published by the journal of visualized experiments (JoVE) (17).

Localization of beads in a nanometer scale
The position of a bead is determined by two dimensional (2D) Gaussian fit to the intensity profile of the bead (Fig. 2D). The 2D Gaussian function is defined as

\[
I(x, y) = I_0 + A \exp\left[ -\frac{(x-x_0)^2 + (y-y_0)^2}{2\sigma^2} \right]
\]

Where \( I_0 \) is a constant due to background noises, \( A \) is the amplitude of the fitted function, \( \sigma \) is the standard deviation of the fitted function, and \( x_0 \) and \( y_0 \) are the bead’s fitted position. By taking all possible noise sources into account, the precision of \( (x_0, y_0) \) is theoretically given by (18)

\[
\sigma = \sqrt{\frac{s^2}{N} + \frac{\alpha^2}{12N} + \frac{88n \beta^2}{\alpha^2 N^2}}
\]

where \( N \) is the number of collected photons, \( \alpha \) is the pixel size of a CCD camera, and \( \beta \) is the standard deviation of the background. The first term is photon shot noise, the second term is due to the finite pixel size of the CCD camera, and the last term is the background effect. This formula explicitly indicates that the more photons collected, the higher position...
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Fig. 4. Time trajectories of enzyme activities with flow-stretching bead assay. (A) Helicase unwinds duplex DNA (dsDNA → ssDNA). Slope in time trajectory is the unwinding rate. Helicase dissociation drives reannealing of unwound strands (ssDNA → dsDNA). (B) The primer extension of DNA polymerases (ssDNA → dsDNA). (C) The strand excision by exonuclease (dsDNA → ssDNA).

accuracy is determined. A powerful tracking software is freely available at DiaTrack (www.diatrack.org).

THE LENGTH CHANGE BETWEEN dsDNA AND ssDNA

The smFS detects activities of DNA enzymes using the different elastic property of single-stranded (ssDNA) and double-stranded DNA (dsDNA). The ssDNA is more flexible (shorter persistence length) than dsDNA and easily forms secondary structure (hairpin-coil) by base pairing in the same strand. Thus, coiling of ssDNA results in shortening of dsDNA at low stretching forces (< 6 pN) (Fig. 3A). At forces higher than 6 pN that is sufficiently strong to break the secondary structure, the ssDNA is extended longer than dsDNA because distance between base pairs in ssDNA is longer than that in the double-helical dsDNA (Fig. 3A). Therefore, conversion from dsDNA to ssDNA or vice versa can be monitored through the motion of beads in the opposite direction (a decrease in total DNA length) or in the same direction as the flow (an increase in total DNA length) (Fig. 3B).

Bead position on the nanometer scale obtained from the pixel size of the CCD camera (effective pixel size = physical pixel size/magnification) is converted to the number of nucleotides (nt) transitioned from dsDNA (ssDNA) to ssDNA (dsDNA) by measuring the length of dsDNA and ssDNA. The ssDNA is prepared by denaturing the dsDNA like 48.5 kilo-base pair (kb) long λ-phage DNA is incubated in 2 mM NaOH solution at 99°C for five minutes and then immediately quenched in a cold blocking buffer (~4°C) to prevent re-annealing. Distance (d_{dsDNA(ssDNA)}) between one end of dsDNA (ssDNA) (anchor point on the surface) and the center of a bead is determined by measuring half the distance (l) between the centers of the bead at opposite directions of laminar flow (Fig. 3C). The length (l) of DNA is $l = d_{dsDNA(ssDNA)} - R_{bead}$, where $R_{bead}$ is a bead radius. Conversion factor for transition between dsDNA and ssDNA is given by $nt/(l_{dsDNA} - l_{ssDNA})$.

APPLICATIONS IN DNA ENZYMEOLOGY

Since the single-molecule kinetics study of λ-exonuclease (19), the smFS has been successfully used for studying enzyme activities by multi-protein complexes in DNA replication (20-22) and DNA repair (23, 24). During DNA replication, helicase transforms from dsDNA to ssDNA by unwinding duplex DNA at the replication fork, resulting in shortening of DNA (Fig. 4A). Single-strand binding protein (SSB) is associated with resulting ssDNA before DNA synthesis by DNA polymerase and is also involved in DNA repair and recombination. DNA polymerase in the replisome synthesizes DNA on resulting ssDNA as a template, resulting in processivity of enzymatic activity (Fig. 4A, B). The strand excision by exonuclease is a crucial step to repair DNA, resulting in transition of dsDNA to ssDNA (Fig. 4C). Using the smFS, Park et al. showed the proofreading activity of E. coli DNA polymerase III when it incorrectly synthesizes DNA (26).
Jeon et al. revealed the mechanism of strand elimination by human ExoI exonuclease through sequential coordination of components in DNA mismatch repair (24).

In addition to the detection of the transition of dsDNA (ssDNA) to ssDNA (dsDNA), since SSB binding to ssDNA extends ssDNA, but SSB-bound ssDNA is shorter than dsDNA, the smFS can visualize the SSB association and dissociation from ssDNA (21, 23). A transient intermediate state, appearing as a pause in the time trajectory, was also detected; the primer synthesis by primase (20) and the stoppage of the Tus-Ter complex in the termination of DNA replication (27). The smFS can visualize in real time the conformational change in dsDNA; the dynamic loop formation and release of the lagging strand during the Okazaki fragment synthesis (21).

**PROSPECTS**

The smFS is a powerful toolbox to study enzymatic activities in a multi-protein complex in various DNA-associated biological events. Especially, it is suitable for highly processive enzymes. However, for higher resolution (more information) of molecular activity in the protein complex, the smFS can be combined with fluorescence imaging. The correlative system such as OT-fluorescence microscopy (37-40) and MT-fluorescence microscopy (28-36) can visualize directly fluorescently-labeled enzymes on the substrate as well as the product as change in microscopy (37-40) can visualize directly fluorescently-labeled enzymes on the substrate as well as the product as change in microscopy. The correlative system such as OT-fluorescence microscopy (37-40) and MT-fluorescence microscopy (28-36) can visualize in real time the conformational change in DNA length. In fact, Loparo et al. developed the smFS combined with total internal reflection fluorescence (TIRF) microscopy to study the exchange of DNA polymerases in DNA replication (41). The micron-scale bead attached to DNA is not compatible with TIRF microscopy due to 100 μm penetration depth of the evanescent wave. Instead, quantum dot (QD) to image change in DNA length was attached to a location close to the replication fork. Remaining DNA after the QD contributes to drag the QD, reducing thermal fluctuations of QD due to the small Stokes drag force on the QD. However, they need much faster flow rate (much more volume of proteins) and only observed less than 100 DNA-tethered QDs per field of view under 60x objective lens with high NA (> 1.2) for single fluorophore imaging. If the TIRF microscopy is replaced with highly inclined thin illumination (HiLo) with longer illumination depth, the larger bead can be used to increase spatial resolution. Immobilization of an array of DNA molecules on the patterned surface can dramatically increase the number of DNA substrates per field of view. We expect that the multiplexed correlation smFS with fluorescence imaging would greatly contribute to reveal unknown mechanisms in DNA-related biological events.

**ACKNOWLEDGEMENTS**

This study was supported by Global Research Lab Program through the NRF of Korea funded by the Ministry of Science and ICT (NRF-2017K1A1A2013241) (J.-B.L.).

**CONFLICTS OF INTEREST**

The authors have no conflicting interests.

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