Highly Parallel Magnetic Tweezers by Targeted DNA Tethering

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ABSTRACT: Single-molecule force-spectroscopy methods such as magnetic and optical tweezers have emerged as powerful tools for the detailed study of biomechanical aspects of DNA–enzyme interactions. As typically only a single molecule of DNA is addressed in an individual experiment, these methods suffer from a low data throughput. Here, we report a novel method for targeted, nonrandom immobilization of DNA-tethered magnetic beads in regular arrays through microcontact printing of DNA end-binding labels. We show that the increase in density due to the arrangement of DNA-bead tethers in regular arrays can give rise to a one-order-of-magnitude improvement in data-throughput in magnetic tweezers experiments. We demonstrate the applicability of this technique in tweezers experiments where up to 450 beads are simultaneously tracked in parallel, yielding statistical data on the mechanics of DNA for 357 molecules from a single experimental run. Our technique paves the way for kilomolecule force spectroscopy experiments, enabling the study of rare events in DNA–protein interactions and the acquisition of large statistical data sets from individual experimental runs.

KEYWORDS: Magnetic tweezers, single-molecule force spectroscopy, microcontact printing, increased packing density

Magnetic tweezers (MT) have become a widely used single-molecule technique for the study of the mechanics of macromolecules and the dynamics of enzymes that act on DNA or RNA.¹ The popularity of the MT technique is due to its low cost, simplicity of implementation, and the possibility to monitor and control the effects of torsional and tensional stress on enzyme–DNA interactions.²³⁴ In magnetic tweezers, a DNA molecule is tethered between a paramagnetic bead and the surface of a flow cell. An external magnet exerts a force and torque on the DNA molecule (see Figure 1a for a schematic).³ The length of the DNA molecule is measured by tracking the XYZ position of the paramagnetic bead using video microscopy, providing a means for monitoring, for example, enzyme-induced changes in DNA topology. The force and torque exerted on the molecule can be extracted from measurements of the position and thermomechanical noise spectrum of the motion of the bead.¹⁴

In standard MT experiments, only a single molecule of DNA is addressed in an individual experiment. Since several paramagnetic beads can be captured in the imaging field of view, MT inherently bear potential for multiplexing,⁵–⁸ thereby increasing the data throughput in single-molecule force—spectroscopy experiments. The achievable data throughput in multiplexed magnetic tweezers is limited by the density of isolated and trackable beads in the field of view. Here, we show that it is possible to fabricate regular arrays of closely spaced DNA-bead tethers, thus increasing the density of trackable beads in the field of view by up to 1 order of magnitude. We create such regular arrays of surface-immobilized tethers through microcontact printing of a pattern of DNA-end-binding labels on the surface of a flow cell (see scheme in Figure 1). The use of microcontact printing in this context is particularly advantageous as it offers a facile, cheap, and robust method for the fabrication of sub-100 nm protein structures.⁹ A lipid bilayer serves as a highly inert surface passivation layer that prevents nonspecific binding of the magnetic beads.

In the standard MT approach, DNA molecules are immobilized on a surface that is randomly covered with DNA-end-binding labels, leading to a sparse, random distribution of DNA-tethered beads. Following this procedure, DNA-tethered beads can only be used for further experimenting and analysis if they are spaced farther than an experiment-specific distance d₀ from their nearest neighbor. This requirement for a minimum bead-to-bead distance results from crosstalk in position tracking when the defraction pattern of the beads overlaps and from attractive forces that arise between neighboring beads as a result of magnetic dipole–dipole coupling that can lead to irreversible pull-in of DNA-tethered beads. The minimum separation thus depends on the size of the defocused bead image as well as the length of the DNA tethers.

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The maximum achievable number of useful DNA-tethered beads can be calculated from the probability density of finding the nearest neighbor at some given distance $r$. For point-like particles that are randomly distributed on a 2D plane, the probability density reads\(^{9}\)

$$\omega(r) = 2\rho\pi c e^{-\rho\pi r^2}$$  \hspace{1cm} \hspace{1cm} (1)

with $\rho$ as the surface density of DNA-tethered beads. The number of productive tethers (i.e., those with $r > d_0$) can therefore be calculated as

$$N_{\text{bead}} = A_{\text{FOV}}\rho \int_{d_0}^{\infty} \omega(r)dr = A_{\text{FOV}}\rho e^{-\rho\pi d_0^2}$$  \hspace{1cm} \hspace{1cm} (2)

with $A_{\text{FOV}}$ the area of the field of view. The optimal density that can be achieved is obtained for $dN/d\rho = 0$, yielding a maximum number of beads of

$$N_{\text{max}} = \frac{A_{\text{FOV}}}{d_0^2} \frac{1}{\pi\epsilon}$$  \hspace{1cm} \hspace{1cm} (3)

A regular distribution of beads results in a much higher stacking density. Full occupation of a square array pattern with pitch $d = d_0$ yields a surface density $N_{\text{max}} = A_{\text{FOV}}/d_0^2$. For a hexagonal pattern with the same pitch, $N_{\text{max}} = (2/\sqrt{3}) (A_{\text{FOV}}/d_0^2)$. Regular arrays can thus lead to an improvement in surface density compared to the case of a random distribution with a factor of $\pi\epsilon$ and $(2/\sqrt{3})\pi\epsilon (\sim 10)$ for a square and hexagonal pattern, respectively.

Here, we create regular arrays of DNA-tethered magnetic beads via surface patterning of antidigoxigenin protein, which specifically binds to a digoxigenin end-labeled DNA. The protein pattern is prepared via microcontact printing. The protocol for microcontact printing is described in Figure 2a. A negative image of the desired protein pattern is created in a silicon template using a combination of electron-beam lithography and dry etching. Figure 2b shows a scanning electron micrograph of a resulting ~100 nm deep square etch pit in silicon. Before reuse of the silicon template, organic residues are removed from the template (Piranha H$_2$SO$_4$/H$_2$O$_2$ 3:1, 10 min), and the silicon template is subsequently O$_2$-plasma treated to make the surface hydrophilic. A disposable, elastomeric stamp with a flat print surface is cut from a precast PDMS layer (Sylgard 184, Dow Corning) and inked with an antidigoxigenin solution (0.5 mg/mL in PBS buffer, 10 mM). After incubation (30 min), the solution is washed off and the surface-adsorbed layer of antidigoxigenin is transferred to the silicon template. A positive image of the desired protein pattern remains on the PDMS stamp. The pattern on the PDMS stamp is subsequently transferred to a glass slide through microcontact printing. For further details on the latter, see Coyer et al. Prior to printing, the glass slide is cleaned and treated with a TL1 solution (NH$_4$OH/H$_2$O$_2$ 3:1) to make the surface hydrophilic.

Figure 2c shows an atomic force microscopy (AFM) image of a square antidigoxigenin patch printed on a glass slide. The AFM scan shows that the conformation of the square defined in the
silicon master is excellently reproduced. The protein patch has a rough surface with median peak height of $\sim 4$ nm, consistent with a high coverage of a monolayer of printed protein, thereby confirming observations in ref 8. Figure 2d shows images of a hexagonal pattern of fluorescently labeled antidigoxigenin square patches (each 200 nm) printed on a glass slide. The image reveals a regularly spaced and uniformly occupied array of fluorescent protein patches ($d_0 = 9 \mu m$).

After microcontact printing of the glass slide, a flow cell for MT experiments is prepared with the protein-printed slide as the bottom layer. In our experiments, protein patterns with a patch size in the range of 200 to 600 nm and pitch in the range of 9 to 20 $\mu m$ are used. The size of the active, DNA-binding area in the resulting pattern is thus only a $10^{10}/C_0$ to $10^{14}/C_0$ fraction of the total area. This places a strong requirement on the ratio between DNA-specific binding and random nonspecific binding of DNA or beads to the surface ($> 10^3$ to $10^5$).

To meet this requirement, a highly inert lipid bilayer is formed in the area between the protein patches in order to prevent nonspecific DNA and bead binding. To this end, 50 nm lipid vesicles were prepared as described previously and suspended in a buffer (10 mM Tris, 100 mM NaCl (pH 8)). A lipid bilayer spontaneously assembles on the protein-free part of the surface upon adsorption and spontaneous rupture of the lipid vesicles (30 min). After surface passivation, double-stranded (ds)DNA molecules (7.3 kb, 500 pM) were introduced into the flow cell (buffer: Tris-HCl (pH 7.8), 100 mM NaCl, 4 mM MgCl$_2$). DNA molecules were prepared as described previously. The DNA is digoxigenin labeled at one end to allow specific binding to the antidigoxigenin patches on the surface, and a biotin label on the other end allows subsequent attachment of streptavidin-coated paramagnetic beads (for more details on the preparation of this 7.3 kb dsDNA construct, see ref 12.). After incubation (10 min), free DNA was removed from the flow cell and 1 $\mu m$ streptavidin-coated paramagnetic beads (MyOne, Invitrogen) are introduced and allowed to settle on the surface (10 min). After removal of nonbound magnetic beads, a regular pattern of surface-tethered magnetic beads remains.

Figure 2e shows an example of a square pattern of DNA-tethered magnetic beads of 1 $\mu m$ diameter. The image shows a zoom (40%) of a 300 $\times$ 400 $\mu m$ field of view in which 996 magnetic beads and bead clusters are organized in a square array. In this field of view, 787 spatially isolated beads were found (using an automated bead-finding procedure, see Supporting Information), that is, a factor of 4.8 more than can be achieved using a random distribution of tethers (see eq 3). An alternative protocol where DNA is first incubated with the magnetic beads and where DNA-bound beads are subsequently introduced in the flow cell yielded similar results (data not shown). Furthermore, backfilling of nonfully occupied arrays is possible through the addition of DNA-bound beads. In following this procedure, binding of DNA-tethered beads to an already occupied protein patch is prevented through steric hindrance (for attachment pads smaller than the bead diameter). As a result, the Poisson limit on the fraction of patches occupied with a single molecule can be overcome.

Figure 3. Highly parallel measurement and analysis of DNA mechanics. (a) Histogram of the measured bead height measured for a 7.3 kb dsDNA at a nominal force of 1.8 pN. $N_{\text{mol}} = 450$ beads were found in a hexagonal pattern with a pitch of 15 $\mu m$ (a fragment of such pattern is shown in the inset). Eighty percent ($N_{\text{mol}} = 357$) of these DNA-bead tethers were found to have an end-to-end distance longer than 0.5 $\mu m$. Red line indicates the expected length. (b) Example of a force response curve measured for a single DNA-bead tether in the array. (c) Scatter plot of the persistence length, $L_p$, and contour length, $L_c$, extracted by fitting a wormlike chain model to the force response of an ensemble of DNA-bead tethers ($N_{\text{mol}} = 250$). (d) Histogram of extracted $L_p$ for $N_{\text{mol}} = 105$. Two peaks are found, centered around $L_p = 48$ nm and $L_p = 25$ nm, corresponding to single and double tethered molecules, respectively.
The above-described microcontact printing based method allows the facile and robust preparation of dense regular arrays of DNA-bead tethers for multiplexed magnetic tweezers with minimal extra processing steps and time required for sample preparation. Particularly advantageous in this context is that the silicon master template, with e-beam defined nanostructures, can be reused to prepare many flow cells, thereby significantly reducing the sample cost. We found that >10 flow cells can reproducibly be prepared by reusing the same master template without noticeable degradation in sample quality.

To demonstrate the potential of extracting large statistically meaningful data sets out of a single experimental run by means of these highly parallel magnetic tweezers, we have measured and analyzed the mechanical properties of an ensemble of 7.3 kb dsDNA. The magnetic tweezers setup used in this work consists of a microscope imaging system with 25 × optical magnification and a 1.4 megapixel camera (Falcon1.4M, Dalsa) resulting in a 300 × 400 μm field of view. Modeling of the force fields generated by the magnet configuration used in this work indicates that the force-variation in the field of view is smaller than 1% in the force range used in the experiments (data not shown). To address the computational challenge of accurate positional tracking of all the magnetic beads in the field of view, we store images acquired by the camera on a hard drive and perform an image analysis offline after the experiment.

A flow cell was prepared with a hexagonal pattern of antidigoxigenin patches (d0 = 15 μm). A total of 450 beads were found in the field of view using an automated procedure (Supporting Information). Figure 3a shows a histogram of the maximum bead height measured during 25 s at a nominal force of Fmag = 1.8 pN in a 10 mM Tris buffer (pH 7.5). Out of 450 beads in the field of view, 80% (357) of the DNA-bead tethers displayed an end-to-end distance longer than 0.5 μm. The histogram of the measured lengths shows a most probable length of 2.31 ± 0.08 μm, close to the expected length = 2.26 ± 0.005 μm (at Fmag = 1.8 pN, 25 s measurement time). Several experimental factors contribute to the observed distribution in end-to-end distances: (1) the random distribution of the positions of attachment of DNA on the bead, (2) errors made in determining the position of the surface of the flow cell, (3) the variation in the conformation of attachment of the DNA-end-binding labels, and last (4) a variation in the number of DNA molecules bound to a single bead (see below).

In the next experiment, we performed a high-throughput analysis of the elastic response of a large set of DNA tethers. Figure 3b shows a measurement of the characteristic elastic response of dsDNA. The force required to extend a dsDNA molecule, thereby reducing the conformational entropy of the molecule, is well described by a wormlike chain (WLC) model.13,14 The model has two parameters, the contour length of the molecule, Lc, and the persistence length, Lp, a measure for the length scale over which orientational fluctuations decay. dsDNA has a well-defined Lc of 0.34 nm per base pair (2.48 μm for 7.3 kb).15 Lp is modestly dependent on salt and was measured to be Lp ≈ 50 nm in the range of 30 mM to 150 mM NaCl.13 In single-molecule pulling experiments, the extracted Lp values furthermore depend on the length of the molecule for short DNA substrates and are affected by degrees of freedom of bead fluctuations.15 We used the dynamic force spectroscopy method proposed by Kruthof et al. for the fast and accurate analysis of the DNA force response.16 The total time required for the experimental run was only 8 min (including measurements of the force response and calibration measurements of a bead-specific height-offset and a force calibration factor). Figure 3c shows a scatter plot of the extracted Lp and Lc values for 105 molecules. Figure 3d shows the distribution of the extracted Lp. Two peaks are easily distinguished and are centered around Lp = 48 nm and Lp = 25 nm, corresponding to singly tethered and doubly tethered beads respectively.17

The highly parallel magnetic tweezers technique used here will be highly useful in the systematic study of the mechanical properties of ssDNA, RNAs and protein-bound DNAs.18 DNA-binding proteins can significantly alter the mechanical parameters of the DNA, and high-throughput measurements of these parameters can yield crucial insight into protein function.3,12

In conclusion, we have described a new method for the targeted surface-attachment of DNA-bead tethers via microcontact printing of DNA-end-binding labels. DNA tethering in a regular array can yield a one-order-of-magnitude increase in the density of DNA-tethered beads on the surface, thereby greatly increasing the data throughput in magnetic tweezers experiments. Similar improvements in throughput can be realized for tethered-particle-motion experiments, holographic optical tweezers experiments and, experiments with torque-sensitive magnetic tweezers.4,19,20 To illustrate the potential of this technique in providing a high-throughput analysis in magnetic tweezers, we performed measurements of the mechanical parameters of up to 357 dsDNA molecules in a single experimental run. Further advances can be made by increasing the image field-of-view size, for example, by making use of a larger camera that leads to an increase of the field-of-view size without compromising the tracking accuracy.5

The method presented here paves the way for kilo-molecule level magnetic tweezers experiments. Kilo-molecule magnetic tweezers will greatly facilitate the force and torque spectroscopy studies of DNA—protein and RNA—protein interactions through the acquisition of large statistical data sets from individual experimental runs.

**ASSOCIATED CONTENT**

Supporting Information. Additional information and figures. This material is available free of charge via the Internet at http://pubs.acs.org.

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