Molecular Threading: Mechanical Extraction, Stretching and Placement of DNA Molecules from a Liquid-Air Interface

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Abstract

We present “molecular threading”, a surface independent tip-based method for stretching and depositing single and double-stranded DNA molecules. DNA is stretched into air at a liquid-air interface, and can be subsequently deposited onto a dry substrate isolated from solution. The design of an apparatus used for molecular threading is presented, and fluorescence and electron microscopies are used to characterize the angular distribution, straightness, and reproducibility of stretched DNA deposited in arrays onto elastomeric surfaces and thin membranes. Molecular threading demonstrates high straightness and uniformity over length scales from nanometers to micrometers, and represents an alternative to existing DNA deposition and linearization methods. These results point towards scalable and high-throughput precision manipulation of single-molecule polymers.


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Introduction

In solution, a double stranded DNA (dsDNA) double-helix adopts an entropically favourable compact random-coil conformation, and a single stranded DNA (ssDNA) molecule, having a smaller persistence length, is more compact still [1]. By applying tension to the strand, a DNA molecule can be stretched into an entropically unfavorable, elongated state.

The first method developed for mechanical elongation of DNA involved stretching fibres of its precipitated sodium salt [2] in air, and a number of techniques for stretching DNA from or in solution have been developed since. Existing bulk methods for stretching DNA include molecular combing [3,4,5,6], transfer printing [7], and shear-induced stretching in micro- and nanochannels [8]. These techniques have been used to prepare DNA for sequence motif mapping and to characterize chromosomal abnormalities [9,10]. Molecular combing - where DNA molecules are elongated on compatible surfaces by the action of a receding meniscus - combined with transfer printing and electron microscopy, shows promise as a tool to enable direct reading of genetic and epigenetic information from stretched DNA molecules [11].

Existing single-molecule manipulation techniques, such as optical and magnetic tweezers [12,13], atomic force microscopy [14], and micro-needle manipulation [15], have had success characterizing the behaviour of DNA and other bio-polymers under tension and torsion, as well as their interactions with other biomolecules [16,17].

Existing methods for stretching DNA have inherent limitations. Bulk methods such as molecular combing or transfer printing require either liquid (buffer) or solid (stamp) contact with the substrate, both of which are incompatible with sensitive or fragile surfaces. Additionally, stretching in molecular combing is sensitive to both the surface and the buffer [18,19,20]. A method to elongate DNA molecules while avoiding bulk contact of the carrier with the substrate is desirable because it circumvents these constraints. While single-molecule manipulation techniques can stretch DNA molecules independent of a surface, existing single-molecule methods do not enable the transfer of stretched molecules to a target substrate and are generally slow.

To overcome these limitations, we have developed an efficient tip-based method to mechanically stretch DNA molecules and deposit them onto a surface [21]. In this novel method, stretching is accomplished by mechanically pulling DNA segments from a droplet using a pulled-glass micro-needle and suspending them in...
air between a liquid-air interface and the micro-needle tip. The segments can be deposited onto a dry substrate in a separate step, avoiding bulk contact of the carrier medium with the deposition surface. We call this method “molecular threading.”

Results

I. Construction and operation of a molecular threading device for DNA

Molecular threading works by extracting one or more DNA molecules from a liquid buffer into air using the tip of a chemically-passivated pulled-glass micro-needle. The passivation layer, created by immersing the needle-tip in a suspension of Poly(methyl methacrylate) (PMMA) in acetone, provides simultaneous DNA affinity and fluid hydrophobicity. Threading proceeds by inserting the passivated needle-tip into a droplet of DNA-containing buffer solution. The needle is then extracted from the droplet, and one or more DNA molecules bound to the needle are pulled through the liquid-air interface of the droplet. Surface tension acting on the DNA at the liquid-air interface provides a restoring force that stretches the DNA segment suspended in air. The stretched DNA can be placed on a substrate positioned underneath it by lowering the needle-tip to the substrate surface, anchoring the molecule by weak forces to the substrate in a stretched configuration.

We designed an apparatus to independently actuate the three main components in threading (glass micro-needle, DNA suspension droplet, and target substrate) and enable real-time monitoring of the process. A schematic representation is shown in Figure 1A, and photographs of the apparatus are shown in Figures S1 and S2. Each component is rigidly attached to an independent long-range three-axis micropositioner for coarse alignment. The motion of the needle must be controlled with higher precision by a secondary computer-controlled piezoelectric nanopositioner mounted on a microchip of the same wafer. The components are aligned inside the nanopositioner working volume (120 μm × 120 μm × 120 μm) using cameras aimed in the -Z (substrate normal) and +Y (perpendicular to needle) axes. Various geometries are possible; the DNA-containing droplet and substrate may be placed in contact with each other or separated by a gap, and the needle may be configured to either contact the substrate during its stroke or avoid contact. In the latter no-contact case, the needle may be extended past the substrate edge along the needle (X) axis and then brought below the substrate edge in the Z-axis to simultaneously shear the strand while depositing it.

Substrate and droplet separation is advantageous in some applications; however we have typically found straighter deposited strands when the substrate is partially contacted by the droplet. This mode is shown in Figure 1B, which depicts the working volume of the instrument in the partially-contacted configuration. The droplet is immobilized between two low-wetting surfaces, and both sides of the liquid-air interface as well as the substrate surface are within the range of fine motion of the micro-needle tip. We call this configuration “spider threading” since the threaded strands extend from a partially-combed region like the legs of a spider. Spider threading has certain advantages: (1) reduced local evaporation rate since only part of the droplet is exposed, (2) straighter strands due to more consistent meniscus shape and reduced evaporation rate, and (3) additional deposition of a partially-combed region with high-density fibrous bundles which can serve as fiducials during imaging for identifying the maximum extent of the meniscus. Figure 1C illustrates the steps of the optimized “spider” molecular threading process:

i) The nanopositioner moves according to a pre-programmed path, causing the tip of the needle to penetrate the droplet meniscus and into the liquid.

ii) The positioner then reverses direction, extracting the needle from the droplet by a pre-programmed distance (extraction distance). Any DNA molecules attached to the needle, collectively called a “thread”, are mechanically separated from the droplet and its contents, and suspended in tension above the substrate between the withdrawing tip and the liquid-air interface by the restoring force of the meniscus. The number of attached and/or intertwined DNA molecules attached to the tip is not known until subsequent interrogation, e.g., via high-resolution imaging such as TEM or AFM. The thread remains normal to the droplet surface during manipulation; this effect is shown in Video S1 for a multi-strand thread large enough to be visible via light microscopy. As the thread exceeds its extraction limit and is pulled out of the meniscus, it un-stretches but remains bound to the needle. By reintroducing the needle to the droplet, the meniscus force on the DNA is restored and it can be re-stretched by the needle action. This indicates that if the DNA remains bound to the needle tip the threading process is reversible.

iii) When the tip is brought into contact with the substrate, the thread is transferred and immobilized onto the substrate surface. This contact with the surface also cleans the tip, freeing it to repeat the threading process.

iv) By translating the tip across the substrate transverse to the previously-deposited threads (Y-axis), an array of adjacent, approximately parallel threads can be created with an operator-defined lateral pitch.

The process may be repeated indefinitely over the same substrate area, producing arbitrarily dense arrays of straight parallel threads.

II. Properties of threaded DNA

Molecular threading is a high-throughput technique: the motion of the piezoelectric nanopositioner sets the speed of threading, which is able to exceed 10 Hz. To assess macroscale properties, we prepared arrays (typically 50–100 threads) from single and double stranded DNA by threading on low-wetting elastomer shelves using a nominal thread-to-thread pitch of 1 μm (Figures 2–3). The threaded DNA was stained with YOYO-1 intercalating dye and imaged by fluorescence microscopy. For investigations at higher resolution, we conducted electron microscopy (EM) imaging of threaded modified ssDNA strands stained with a thymidine-selective osmium tetroxide 2,2’-bipyridine (osbipy) contrast-enhancing label (Figures 4–5). The synthesis and labelling kinetics of this agent have been discussed in previous work [22].

III. Applicability to multiple DNA and substrate types

Molecular threading reliably produces arrays of parallel threads on PDMS substrates using both single-stranded and double-stranded DNA, and is compatible with a variety of DNA and buffer combinations. After staining the substrate surface with an intercalating dye these arrays can be detected using fluorescence microscopy (Figure 2). Figure 2A shows long (~380 kbp) m13mp18 ssDNA concatamers prepared using rolling circle amplification [23] that were threaded with an extraction distance of 40 μm. The 167 kbp double-stranded viral genome of bacteriophage T4 GT7 was also found to have similarly high
deposition efficiency at the same extraction distance (Figure 2B). However, the shorter, double-stranded viral genome of λ bacteriophage (48.5 kbp, approximately 16 μm crystallographic length) did not thread successfully at this extraction distance: only at a shorter extraction distances of less than 20 μm could DNA be detected (Figure 2C). This result indicates that the extraction distance for a given DNA length cannot exceed a critical value, and any attempt to extract a thread beyond this distance causes threading yields to drop precipitously. This is expected, since after extraction a sufficient length of excess DNA must remain in the droplet so as to allow controlled stretching and placement of the thread by the micro-needle.

Arrays of parallel threads may also be placed onto many types of unsupported thin films, which remain intact even after repeated contact with the tip of the needle. Commercially-available 5 nm thick Si and SiN TEM grids from TEMWindows and Dune

Figure 1. Molecular threading apparatus. (A) Device for stretching and positioning DNA molecules extracted at a liquid-air interface. A set of three 3-axis micro-controllers are used for coarse alignment of a DNA-binding tip, a target substrate, and the droplet containing a DNA suspension. Horizontal and vertical cameras are used to monitor the process (vertical camera not shown). (B) Illustration of the work area for “spider” threading. The droplet is immobilized between two low-wetting surfaces and a piezo-controlled tip is then moved between the surface of the suspension and the adjacent substrate. (C) Molecular threading at a liquid-air interface. (i) Needle tip penetrates into DNA suspension, binding to one or more strands (collectively called a “thread”). (ii) Needle tip extracts from the suspension parallel to substrate surface. The restoring force of the meniscus places the bound thread in tension. (iii) Needle tip is brought into direct contact with the target surface, transferring the strand to the substrate and the needle. (iv) A lateral translation of the needle after each extraction permits the creation of an array of linearized DNA. (D) Images of the needle, droplet, and substrate, taken under magnification, and corresponding to the illustrations in (C).

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Figure 2. Threading of single and double-stranded DNA. Various DNA types are compatible with molecular threading. (A) Concatamers of m13mp18 ssDNA prepared by rolling-circle amplification, extracted to 40 μm. (B) Double-stranded genomic T4 GT7 bacteriophage DNA, extracted to 40 μm. (C) Double-stranded λ bacteriophage DNA, extracted to 10 μm. All samples were threaded onto different elastomer substrates and stained with intercalating dye YOYO-1 prior to deposition.

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Sciences are compatible with threading, and floated carbon supports as thin as 2.5 nm can be threaded on if the droplet and needle do not contact the film. For tougher semiconductor film types, the DNA-containing droplet can be placed directly in contact with the bulk support adjacent to the film window that will be used to support threaded DNA (Figure S3). Strands are deposited by touching the needle at its full retraction to the window membrane (i.e., spider threading mode). Si tend to stick to the needle or tear after a few hundred depositions, whereas SiN films are more robust. In the alternate geometry where droplet and needle do not contact the substrate, threads can be pulled across the width of a microfabricated cantilever substrate and deposited by lowering the tip below the far side of the cantilever (Figure S4, S5). A more detailed description of cantilever threading mode is described in Text S1.

IV. Uniformity and straightness

We used fluorescence imaging to assay the angular distribution of m13mp18 threads approximately 20 μm long within two different arrays prepared on the same surface. A least-squares fit measuring thread direction was applied to each of 120 threads taken from four images (Figure 3A), which were found to fit a normal distribution with standard deviation of 1.16° for the 20 μm long threads (Figure 3B). The angle variability was found to be random across individual threads in the array. This knowledge allows us to estimate the probability that a thread of length L will overlap its neighboring thread at a given thread-to-thread pitch P. Using the fitted angular distribution, L can be bound by a function of P such that crossovers between threads fall within an acceptable error tolerance. To do so, we assume that (i) the width of a thread is small compared to both thread length and thread pitch and (ii) the distribution of thread-to-surface angles is independent of the thread-to-thread pitch. The location of the first plausible collision between two adjacent threads, one with angle θi and the other with angle θj, is Lc(θi, θj) = P/2 tan(θi) μm from the edge of the surface, or approximately 8 μm for the 1 μm pitch being examined here. A more typical collision would, for example, involve two adjacent strands at angles ±σ, yielding a distance of approximately 25 μm from the surface edge. Intertread collisions can typically be avoided for a given pitch by only extracting threads to a conservative Lc distance, while applications tolerant of overlap may use a more liberal Lc or longer extraction distance.

Besides the angular variation between threads, other types of variations are apparent. These are illustrated in the 12 μm wide area containing ten threads in Figure 3C (area sampled from...
Figure 3A), where the intensity profile from the fluorescent image is superimposed onto interval markers representing the programmed thread spacing of ~1 μm:

1) **Missing threads**: Two of eleven expected threads are missing in the 12 μm window.

2) **Intertwined multiples**: Some threads have higher fluorescence intensity than others, indicating multiple DNA strands.

3) **Forks**: The left-most profile is a pair of threads spaced too closely and which fork toward the bottom of the image, indicating that threads may not always bind the needle at the tip and may interact with each other during extraction.

4) **Lateral deviations from expected thread position**: Threads deviate laterally by up to ~200 nm in position.

**Discussion**

I. Single vs. multi-strand threads

One consideration for this technique is its ability to deliver single strands versus hairpins or fibrous, multi-strand threads. Based on scanning TEM images from spider threading experiments on thin semiconducting thin films (see Figure 4A-B for a representative example), we found strands are often thicker near the needle touch point, and thinner near the droplet meniscus, suggesting that these threads assume a hairpin shape. This observation is consistent with the limited threading success of short DNA in Figure 2C. In Figure 4C, four threads modified with osbipy label are observed with differing integrated intensities. The background-subtracted relative intensities indicate that the left-most thread comprises about four times as many strands as the other threads.

**Figure 4. SEM characterization of threaded DNA.** All images were taken on a Hitachi S-5500 high-resolution transmission SEM (~1 nm resolution), operated at 30 kV, with 50 μA maximum post-flash current. (A) Synthetic ssDNA modified with EM-dense labels to enhance contrast against 5 nm thick semiconductor support films. The deposition pattern indicates the behavior of DNA under the action of a receding meniscus compared to the molecular threading process. (B) High resolution image of a section of four deposited threads ((A) – red box). (C) Intensity profile for a cross section of (B) - yellow box). The non-homogeneous contrast between threads is likely explained by variability in the number of ssDNA molecules making up each thread, while non-homogeneous contrast between different regions of the same thread is likely explained by secondary structure. (D & E) Comparison between (D) molecular threading and (E) molecular combing of synthetic ssDNA, modified with a thymidine-specific label osmium 2-2'-bipyridine (osbipy), onto 5 nm thick amorphous Si substrates, with contrast around strand backbone enhanced for publication clarity.

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two right-most ones, and the second left-most thread comprises about three times as many. Correspondingly, we surmise that, as long as the DNA has a sufficient extraction length, a distribution of single-stranded and multi-stranded segments extending from the meniscus will be present. For applications where single-strandedness is crucial, as in EM-based DNA sequencing, the strands must be very long and extraction lengths scaled commensurately to ensure that the desired quantity of single-strand segments are deposited. At higher EM resolution, quantification of single-strandedness, base-to-base stretching, and observation of individual nucleotide labeling is possible, though this is beyond the scope of this paper and is discussed elsewhere [24, 25]. However, several qualitative features of threading that can be discerned from high-resolution images are examined in Figure S6.

II. Comparison to other linearization techniques

Threaded DNA is straighter than DNA obtained using molecular combing. Threaded osbipy-labeled DNA (Figure 4D) deviates laterally by less than 10 nm over longitudinal strand distances exceeding 1 μm, whereas for combed DNA (Figure 4E), deviations of the same type are often an order of magnitude larger over the same distances, since variations in boundary dynamics of the carrier droplet and local interactions between strand and substrate can cause substantial variations in combing quality.

We investigated local roughness and waviness of threads by atomic-resolution TEM, and compared our findings with combing based on several results from the literature [6, 19, 20, 22, 33, 34, 35, 36, 37, 38, 39, 40] (see Figure 5). Threads are extremely straight both locally (<100 nm length) and over

Figure 5. Advantages of molecular threading over combing. (A) Comparison of maximum angular deviation between threading and combing examples from the literature. (B) Comparison of maximum lateral deviation in μm for similar strand length. (C) Comparison between molecular threading (i) and combing (ii–iii) at high resolution, with synthetic DNA base-specifically labelled with targeted EM-dense contrast agents placed onto 5 nm thick amorphous Si substrates. In general, threaded strands are much straighter than combed strands. Combed strands can be quite straight in sections on the order of 100 nm (ii), but over larger distances large lateral deviations can occur in a different region along the same strand (iii). Composite images are 230 nm sections of imaged strands several μm long obtained on a Nion UltraSTEM 100 atomic-resolution STEM (~1 Å resolution) retrofitted to do automatic stage movement. The UltraSTEM is operated at 60 kV, with 100 pA current measured on the Ronchigram." Contrast around strand backbone is enhanced for publication clarity.

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larger subunit lengths (>100 nm length). In contrast, combed strands are wavy and less uniform even for short subunit lengths (<100 nm length). Combed strands can be very straight in some segments (>100 nm long—Figure 5Gi) exhibits small lateral deviation amplitude of just a few nm—but very large deviations exceeding 10’s of nm invariably manifest along the same combed backbone (see Figure 5Gli). A survey of molecular combing examples from the literature show that lateral position deviation for favorable combing (straightest and most uniform) is about 120 nm RMS along >10 μm long strand lengths. AFM images at higher magnification show 30 nm RMS fluctuations for shorter (~1 μm) strand lengths.

Threaded array density is also orders of magnitude higher than combing density (see Figure 4E). Threading is typically able to place threads spaced <100 nm apart with >50% yield (a yield of ~65% is showcased in Figure 3A), whereas combing typically places strands 1 μm to 10 μm apart. Moreover, as long as the position of the droplet is continuously adjusted to compensate for evaporation, the same needle can be used to re-thread in a previously-threaded region and thus permits arbitrarily high densities of linearized DNA. Thread yield is related to strandedness through droplet DNA concentration, so lower strand count per thread and high array density are competing parameters; if strand count is not a concern, thread yields of >90% can be obtained by using a sufficiently high (>25 ng/μL) DNA concentration. In contrast, the starting positions of combed strands are defined in stochastic pinning events during the combing process, so that strands are rarely found in high densities. The combing example in Figure 4E is a rare case where two combed strands were found very close together and is used for illustrative purposes; adjacent combed strands are rarely separated by less than 10 μm. In surveying the literature, we found combed samples of favorable quality had a density less than 0.01 strands per μm² of sampled image. Angular variability for threaded versus combed arrays was found to be <1° and ~7°, respectively (<100 μm image field).

Threaded samples are also much cleaner than combed ones since the droplet medium does not contact the array surface. This difference is apparent when comparing threading and combing from the same solution. In combing, uncombed supercoiled DNA balls deposited on the surface during the combing process are typically visible whereas the strand-to-ratio for threading is much higher. However, threading produces hairpin shapes as well as a distribution of strands comprising each thread, so combed DNA is more likely to be single-stranded and fully elongated.

The figures shown, as well as those surveyed in the literature, are selections of high quality threading and combing, although poorly threaded strands may also exhibit imperfections such as kinks, breaks, fibrousness, and gentle curves over micrometer lengths. However, as a general rule, we have found that threading gives more consistent results, orders-of-magnitude improvement in array density, and superior straightness compared to combing at all length scales down to the nanoscale. For DNA linearization, threading also has broader applicability than current AFM techniques. While AFM may currently possess scaling efficiencies for DNA extraction, and (ii) missing threads. For the first failure mode, the presence of multiple threads can be explained by the tip binding to multiple molecules in the droplet. Alternatively, multiple DNA molecules may interlink in solution (via hybridization, cross-linking, or physical tangling) such that the stretching of one thread bound to the needle induces the stretching and transport of one or more unbound strands. For the second failure mode, missing threads can result from a lack of binding between the needle and DNA, or from a bond molecule that is shorter than the extraction distance being prematurely pulled out of the meniscus. Strategies to reduce the number of failed threads include: (1) monofunctionalized needles to mediate specific interactions with chemically modified DNA and improve binding probability, (2) needles that impose steric constraints on the number of simultaneously bound strands, and (3) tuning the composition of the liquid droplet such that distinct DNA molecules strongly repel one another. Real-time monitoring using fluorescence rather than bright field imaging would also permit direct observation of DNA-nucleotide binding events, from which the time constant for attachment can be inferred.

IV. Implications of this work and conclusions

The field of DNA nanotechnology has historically been driven by bottom-up, self-assembly techniques [26] but complementary results can often be achieved through high-throughput, top-down nanomanipulation approaches using mesoscale or micro-scale tools. Molecular threading is one such approach which produces reliable results. Because threading stretches DNA in air rather than in liquid, the extended thread can be placed onto water-soluble, dry, or fragile surfaces. Since the positions of threads and the spacings between adjacent threads are determined by the programmed mechanical motion of the piezoelectric positioning rather than stochastic binding events, the user can precisely control the spatial arrangement of threads on the target surface. These advantages are significant for many DNA positioning applications, such as in DNA sequencing by transmission electron microscopy (TEM), where a highly reproducible density of single strands on ultra-thin electron-transparent supports is required [11,21,27,28,29]. The speed of molecular threading is limited by the motion of the nanopositioner, which can operate with <100 ms cycles. Additional throughput gains may be achieved by threading with arrays of parallel needles. One such scheme is detailed in Figure 5Si.

Applications beyond sequencing include nanofabrication, such as aperiodic templates for organic or inorganic materials using DNA as an organizing scaffold, or precision-patterned DNA nanowire arrays. Future work will explore the use of molecular threading as a tool for generating new types of DNA nanostructures such as anisotropic DNA sheets or webs, or asymmetric patterns, which have favourable electrical, optical or catalytic properties once modified. In certain applications, directed overlap between threads—as in Figure 3B for threads stretched more than $L_{str}$—can be advantageous, and this has been demonstrated in studies involving DNA [30].
Molecular threading shows great potential as a high-throughput DNA manipulation technology. Much work is needed before it can be reliably applied in scaled DNA sequencing or material templating technologies, but in its current form it demonstrates clear advantages over other manipulation techniques such as molecular combing. The straightness, reproducibility, and array density are at least an order of magnitude better than competing techniques. Important next steps include (1) improving the characterization of deposited threads given a range of environmental conditions and (2) improving our understanding of the physics for the DNA unwinding process at the liquid-air interface, which might be achieved through molecular modeling. Answers to these questions would point the way toward a new apparatus design with an eye toward industrial scalability.

**Materials and Methods**

**Nomenclature**

We term a “thread” to be one or more DNA strands elongated by a PMMA-passivated needle withdrawing from a liquid-air interface and deposited on a surface. A thread must subsequently be interrogated by high resolution imaging to determine whether it is a single strand or a collection of strands.

**Preparation of a DNA-binding Glass Micro-Needle**

Micro-needles were pulled from a 1.0 mm diameter borosilicate glass rod using a Sutter P-2000 laser-based micropipette puller with the following parameters: HEAT = 825, FIL = 2, VEL = 20, DEL = 255, PUL = 150. The PMMA suspension was prepared by dissolving 100,000 M.W. PMMA beads (Polysciences) in acetone, and then serially diluting in acetone to a final concentration of 0.1 mg/mL. The tip of the needle was immersed in the suspension for 1 s, then removed from the liquid and suspended in acetone vapor 1 cm above the surface of the PMMA-in-acetone suspension for 1 minute while sealing the container with a gloved hand.

**Instrumentation**

Three Sutter MP-225 micro-manipulators were connected to two daisy-chained Sutter MP-200 control boxes and controlled from a single Sutter ROE-200 unit. A PI P-611 NanoCube piezo stage nanopositioner was mechanically coupled to one manipulator using a machined aluminum interface and was connected to a corresponding PI E-664 NanoCube piezo controller. The controller, using a 3 X BNC/37 pin sub-D adapter, was connected to an Acces I/O USB-AO16-16A 16-bit analog/digital for control via USB from a desktop computer using software written in Java. Two Qioptiq Optem 70XL micro-inspection lenses were illuminated with two Dolan Jenner MI 150 fibre optic illuminators and were used for real time monitoring.

**Design of component interfaces**

A machined aluminum interface for the micro-needle (“needle holder”) connects the micro-needle to the piezo stage. It consists of a fixed aluminum base, attached to the piezo stage, and a removable aluminum needle holder coupled to the base using a spring-clip. A series of grooves, 1 mm in diameter, are cut radially into the needle holder, spanning 0° to −25° at 5° intervals (relative to the plane of the instrument in Figure 1A). A small aluminum block attached to an adjustable screw holds a micro-needle in one of the grooves. In this report, the micro-needle is always oriented at −15°. An image of the interface is shown in Figure S1. A glass rod using a Sutter P-2000 laser-based micropipette puller with a fixed aluminum base, attached to the piezo stage, and a removable aluminum needle holder coupled to the base using a clamp supplied with the device. A machined aluminum interface for the substrate (“substrate holder”) was also fabricated. It consists of a base, mechanically coupled to a second micro-manipulator, and a removable platform coupled to the base using a spring-clip. A substrate can be placed directly on the platform. A sample droplet holder formed by a 3/8” steel rod wrapped in paraffilm, was coupled to the third micro-manipulator using a clamp supplied with the device.

**Preparation of substrates**

An elastomer gel substrate was removed from a Gel-Pak Gel-Box chip-holder with tweezers, and cut into 0.5 cm² pieces with a razor-blade. A 45° incline was cut into one side of each piece, and each piece was washed in TE buffer for 3 min. A piece was then placed incline-forward on the aluminium substrate holder.

**Preparation of long ssDNA using rolling circle amplification**

10 μL of 10× reaction buffer (10× phi29 DNA Polymerase Buffer B7020, Enzymatics, 500 mMTris-HCl, 100 mM (NH4)2SO4, 40 mM DTT, 100 mM MgCl2, pH 7.5), 1 μL of 1 nM M13mp18 template (NEB), 2.5 μL of 100 mM primer (TCCAAAGTCAGGGGCGAAAAACG, ID 1) and 1.6 μL of dNTP mix (Enzymatics N2050L) was brought to a volume of 48 μL in water. The mixture was incubated at 95°C for 1 min, then 60°C for 1 min, then brought to 4°C. The mixture was put on ice, and 2 μL of phi29 DNA polymerase (10 μL/μL, Enzymatics P7020-LC-L) was added. The whole mixture was then incubated at 30°C for 4 hr, then brought to 4°C diluted in 450 μL of 1× PBS (pH 7.4). The recovered solution was then diluted 100× in PBS and stored at 4°C until needed. The dilution was brought to room temperature, mixed by pipetting half its volume up and down several times, and then directly used as described in the molecular threading procedure.

**Preparation of long dsDNA molecules**

T4GT7 phage stock solution (Wako, 166 kb, 300 ng/μL) was diluted 100× in TE buffer to a final concentration of 3 ng/μL and was stored at 4°C. φX phage stock solution (NEB, 48.5 kb, 500 ng/μL) was similarly diluted in TE buffer to a final concentration of 5 ng/μL and was stored at 4°C.

**Molecular threading procedure**

A PMMA-treated micro-needle was set at −15° using the needle holder. A cut elastomer substrate was placed with tweezers on the substrate holder. A suspension of DNA molecules was brought to room temperature from 4°C, and mixed by pipetting half its volume up and down several times. A 5 μL droplet was then pipetted on to the end of the paraffilm coated rod. To create an array of threads, the piezoelectric nanopositioner executed the following pre-programmed motions:

1. Translate x1 μm toward droplet (needle penetrates droplet).
2. Translate x2 μm away from droplet (needle and attached thread is withdrawn from droplet).
3. Simultaneously translate x3 μm away from droplet and z1 μm toward surface (needle contacts surface)
4. Translate z1 μm away from surface (needle detaches from thread)
5. Translate y1 μm parallel to droplet (needle prepares to thread next element in array).
6. Execute the above instructions N times.

Unless otherwise noted, RCA M13mp18 and T4GT7 phage were prepared with x1 = 40, x2 = 30, x3 = 10, z1 = 20, y1 = 1 and N = 90, while φX phage was prepared with x1 = 20, x2 = 10, x3 = 10, z1 = 10, y1 = 1, and N = 90.
Preparation of Surfaces for Fluorescence Microscopy

1 mM YOYO-1 intercalator (Invitrogen) was diluted 10000× in TE buffer (10 mM of Tris-HCl/1 mM of EDTA, pH 8) and a 5 µL droplet was pipetted onto a surface prepared by molecular threading. After a 3 min incubation the surface was washed twice with TE buffer and placed surface-down onto a glass cover slip.

Fluorescence Microscopy

All fluorescent images were obtained using oil immersion at 160× magnification using a Zeiss Axio Observer Z2 microscope with a Hamamatsu C9100 camera under a GFP filter set (excitation 450–490 nm, emission 500–550 nm). The images presented in Figure 2 were adjusted for brightness and contrast for publication clarity.

Electron Microscopy

Samples were imaged by electron microscopy in two steps, starting with a screening step at spatial resolutions of approximately 1 nm and followed by high resolution imaging at 1 Å. The screening was conducted in the Hitachi S-5500 - a high-performance scanning electron microscope (SEM) operating in the enhanced-resolution dark field transmission mode (Figure 4). Aberration-corrected scanning transmission electron microscopy (STEM) was carried out in a specialized Nion UltraSTEM 100 [43] modified for automated image acquisition over large sample areas at atomic resolution (Figure 5). Images were automatically captured and post-processed to create composite maps of complete strands. This customized system and its capabilities are described in detail elsewhere [25].

Image analysis

A line perpendicular to the edge of the substrate surface was superimposed on the left side of each fluorescence image of an array of threads, and a curve was traced on top of each thread in the image. ImageJ [44] was used to obtain a cross-section of each modified image at 1 μm intervals along the measurement axis, and the collection of distances between the measurement axis and the traced curves was used to fit a regression line to each thread. The slope of each regression line was then used to compute the angle made with the measurement axis, and a normal distribution and the collection of distances between the measurement axis and a normal distribution was fit to the distribution of angles. Three outliers were discarded.

Post-processed electron microscopy images were loaded into Gatan DigitalMicrograph, part of the Gatan Microscopy Suite v1.8.0. Straightness and orientation measurements of threaded and combed strands were conducted by placing a reference line for each thread using a least-squares estimate. Maximum deviation measurements were taken along this trendline for rolling 1 μm segments of 100 nm along the reference line. Images of combed samples from the literature used for the comparison in Figure 5 were first calibrated for scale after loading into DigitalMicrograph and then similarly processed to extract strand angles and lateral deviation figures. Poorly combed strands, such as hooks and loops, were considered outliers and excluded from the averages.

Supporting Information

Figure S1 The molecular threading apparatus and the primary threading components. (A) The complete apparatus is desktop-sized and relatively portable. (B) A small rod covered in parafilm is a sufficient support for the DNA containing droplet which provides the liquid-air interface. (C) The angle the pulled-glass micro-needle makes with the droplet surface can be adjusted in 5° increments using a machined holder. (D) The PDMS shelf sits on a machined holder; a screw can be turned to align the shelf with the needle and droplet. If a TEM support is placed on the shelf, threads can be directly deposited onto an unsupported thin film.

(TIF)

Figure S2 Preparation of the working volume using the molecular threading apparatus. (A) Apparatus with primary components detached. The interfaces for the micro-needle, droplet, and substrate holders are each attached to a three-axis long-range micro-positioner. The two cameras used for real time monitoring are positioned over the 120 μm³ working volume. (B) With the holders attached, the micro-needle, PDMS substrate, and DNA containing droplet are positioned in the working volume. (C) Alternate angle of image in (A). (D) Alternate angle of image in (B).

(TIF)

Figure S3 Molecular threading on Si and SiN thin films. (A) Top view. The DNA-containing droplet is brought into contact with the edge of a 5 nm Si unsupported thin film on a TEM grid (TEM Windows). The grid sits on top of the PDMS shelf shown in Figure 1. (B) Side view. The micro-needle moves according to Figure 1D and the description in Materials and Methods. DNA is extracted at the liquid-air interface and deposited when the needle comes into contact with the thin film.

(TIF)

Figure S4 Design of a cantilever-based substrate for molecular threading. (A): Mock-up of a cantilever substrate. The manipulation handle (i) is used as an attachment point for vacuum tweezers for transportation and alignment (with the droplet and micro-needle). The cantilever (ii) has several lithographically etched windows across which an unsupported thin carbon film is floated. Any threads deposited across the film can be subsequently examined via transmission electron microscopy. (B): An electron micrograph of the cantilever. (C): Alignment of the droplet, cantilever, and micro-needle in preparation for threading. Image taken under magnification.

(TIF)

Figure S5 DRIE/vapor HF processing of cantilever grids. (A) Optical micrograph showing two sizes of cantilever grids fabricated using DRIE/vapor-HF process. Smaller and larger grids are approximately 1 mm×1 mm and 2 mm×2 mm, respectively. Most of the grid is provided for handling and mounting. Sample is threaded across cantilever, perpendicular to long axis. (B) SEM image of cantilever from larger grid fabricated by DRIE/vapor-HF. Sample molecules are threaded across cantilever and over circular windows. Windows are approximately 3–4 nm thick and are fabricated with wrinkle-free, amorphous carbon. Inspection at higher magnification showed all windows to be intact.

(TIF)

Figure S6 High resolution imaging of threads. (A) A representative sample of threaded and labelled DNA used for high resolution imaging. (B) Regions of higher intensity along the strand are due to higher label density, likely caused by label aggregation, strand coiling, or cross-linking with small labelled oligonucleotides. (C) The needle-tip deposits a region of high label density when transferring its DNA handle to the surface. (D) Un-threaded DNA coils, similar to those found in molecular combing,
are also deposited during either the alignment or threading process.

**Figure S7** Future work with needle arrays to improve throughput. (A) One scheme to improve throughput via a needle array. (i) A “pool” containing long DNA molecules is aligned with an array of glass micro-needles treated with PMMA. A small lateral offset is applied to each row to avoid strand overlap. (ii) The entire array is inserted into the solution. (iii) Multiple DNA molecules are suspended at the liquid-air interface. A motorized stage supporting a PDMS substrate, or TEM support similar to the cantilever discussed in Text S1, is moved across the surface of the pool in order to collect the extracted strands. (iv) The collected strands can now be imaged via fluorescence microscopy or transmission electron microscopy, depending on the substrate. (B) Prototype ultra-high-aspect ratio microfabricated needle arrays to be studied in future work. Needle tips have ~10–20 nm radius of curvature.

**File S1** Microfabrication of “cantilever” EM grids designed for use with threading.

**References**


