Multiplexed single-molecule force spectroscopy using a centrifuge

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We present a miniature centrifuge force microscope (CFM) that repurposes a benchtop centrifuge for high-throughput single-molecule experiments with high-resolution particle tracking, a large force range, temperature control and simple push-button operation. Incorporating DNA nanoswitches to enable repeated interrogation by force of single molecular pairs, we demonstrate increased throughput, reliability and the ability to characterize population heterogeneity. We perform spatiotemporally multiplexed experiments to collect 1,863 bond rupture statistics from 538 traceable molecular pairs in a single experiment, and show that 2 populations of DNA zippers can be distinguished using per-molecule statistics to reduce noise.
The ability to mechanically manipulate single molecules is leading to insights throughout biomedical research, from the action of molecular motors in replication and transcription to the role of mechanical forces in development\(^1\)–\(^3\). While in principle these approaches enable the full characterization of individual molecular complexes and the study of population heterogeneity at the single-molecule level, in practice key challenges exist. The first challenge for force spectroscopy studies is the low throughput of most single-molecule approaches, which is just starting to be addressed with recently developed multiplexed methods\(^6\)–\(^13\). Furthermore, sufficient statistics must be collected not only for the population\(^14\), but also for each individual molecule, which can be a challenge for studying catastrophic transitions such as bond rupture\(^15\)–\(^17\). Another challenge is the positive identification of the single-molecule interactions of interest over non-specific and multiple interactions. Finally, there is the subtle challenge of noise, both thermal and experimental, that makes distinguishing different populations of molecules with similar force properties difficult.

We have met all of these challenges with spatiotemporally multiplexed force spectroscopy, a combination of massively parallel spatial multiplexing with repeated interrogation enabled by self-assembled nanoscale devices. First, we introduce a miniature centrifuge force microscope (CFM) for high-throughput single-molecule experimentation that utilizes a commercial benchtop centrifuge. Functionally similar to our first prototype device\(^18\), an entire microscope imaging system is rotated to observe microscopic objects subjected to uniform centrifugal force (unlike earlier ‘spinning disk’ centrifuge microscopes\(^19\),\(^20\)). This inexpensive design brings new features including temperature control and high-resolution particle tracking (~2 nm). Second, we introduce a high-throughput CFM assay that integrates mechanical nanoswitches\(^17\),\(^21\),\(^22\) to provide important new functionality. The nanoswitches serve two roles—one as a molecular signature to facilitate reliable and automated analysis of large data sets, and the second to enable the repeated interrogation of each single-molecule pair, increasing throughput and enabling new measurements of heterogeneity in single-molecule experiments. By making repeated force measurements on hundreds of single-molecule complexes, we can collect multiple statistics on each molecule that comprises the population. We additionally show that by averaging multiple rupture forces on a per-molecule basis, we can reduce noise to enable super-resolved force spectroscopy—the identification of different populations of molecules below the thermal force-resolution limit. Averaging allows us to reduce the spread in force distributions (averaging reduces noise by a factor of \(\sim \sqrt{N}\)) without losing information about differences between molecules. Furthermore, the rich and relatively large data sets provided by our technique could also complement other analysis techniques for statistical deconvolution\(^23\),\(^24\).

**Results**

**Benchtop CFM with nanoscale tracking.** The redesigned CFM is a miniaturized microscope that fits into a commercial centrifuge so that force can be applied to samples while their micro- to nano-scale motions are tracked (Fig. 1 and Supplementary Fig. 1). To accommodate the microscope components in the 400 ml bucket volume, we selected components with size and weight in mind, and induced two right-angle bends in the optical path (Supplementary Fig. 1b). To ensure accessibility of the instrument, most components are commercially available and can be easily assembled (Supplementary Fig. 1d,e and Supplementary Table 1). A three-dimensional (3D)-printed housing was made to encapsulate the device and to ensure a tight fit within the bucket (Solidworks drawings provided as Supplemental Files). Transmission of camera data out of the CFM during centrifugation is accomplished by converting the camera’s gigabit Ethernet signal to a fibre-optic signal, then passing this data out of the centrifuge through a fibre rotary joint (Supplementary Fig. 1a,f). Details of the design are presented in the Methods.

We implemented a unique solution to measure nanometer-level extensions of tethers in the CFM by projecting tether length changes onto the \(x\)-\(y\) plane of the coverslip, enabling these measurements to be made in a relatively simple and computationally efficient way. Taking advantage of the fact that we can precisely control the direction of force application by mechanically constraining the angle of the centrifuge bucket, we intentionally misaligned the force and imaging axes (Supplementary Fig. 2a and Supplementary Note 1). Using this technique, we can ‘tune’ our tether length resolution based on the bucket angle with a range of \(\sim 2.5\) nm (at 20°) to 12 nm (at 80°) based on our lateral tracking resolution of \(\sim 2\) nm (Supplementary Fig. 2b,c). To validate and demonstrate this approach, we measured DNA force extension for over 100 molecules simultaneously over a span of <1 min (Supplementary Fig. 3a), and fit each with the standard worm-like chain model. The most likely contour length and persistence length was 8.2 ± 0.2 \(\mu\)m and 46 ± 1 nm, respectively, in agreement with expected values\(^25\). We additionally performed multiplexed overstretching measurements of lambda DNA (Supplementary Fig. 3c), yielding an overstretching force of 63.5 ± 1.7 pN (mean ± s.d.), consistent with previous measurements at these conditions\(^25\).

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**Figure 1 | Overview of the benchtop CFM.** (a,i) Illustration of a spinning centrifuge rotor containing the CFM in the red swing bucket; the rotation induces forces on molecules tethered between beads and the coverslip surface. The direction of force application can be set by constraining the angle of the centrifuge bucket, as described in Supplementary Fig. S. (a,ii) Schematic of the CFM. (b) Tether extension is monitored as a function of time while force is applied. When the DNA nanoswitch is integrated, molecular transitions such as bond rupture between a receptor (R)–ligand (L) pair causes a well-defined change in tether extension, providing a distinct signature for detecting interactions between two molecules of interest.
DNA nanoswitches for authenticating single-molecule data. To enable robust and repeatable rupture experiments, we integrated DNA nanoswitches into the CFM. These molecular switches are designed to adopt a looped structure when the molecules of interest are interacting and a linear structure when they are not (Fig. 2a), providing a distinct ‘signature’ (that is, increase in tether length) for rupture events (Supplementary Fig. 4b). We performed DNA unzipping experiments on a 29 bp DNA interaction, and used the ‘signature’ unlooping of the nanoswitches to positively identify and discriminate valid single-molecule data from multiple tethers and non-specific interactions (Fig. 2a and Supplementary Fig. 5). Following this approach, unzipping force measurements were carried out in the presence and absence of magnesium ions, with hundreds of rupture statistics for each condition collected in under 30 s of centrifuge run time. We found that magnesium stabilizes the duplex as observed previously,

Temperature-dependent DNA unzipping. We additionally performed DNA unzipping experiments at four temperatures, 4, 13, 23 and 37°C (Fig. 2c). Seminal works introduced temperature control into optical tweezers over 10 years ago,

Repeated interrogation and super-resolved force spectroscopy. Finally, we demonstrate spatiotemporally multiplexed force spectroscopy using the DNA nanoswitch assay on the CFM to repeatedly interrogate a population of molecules at the single-molecule level (Fig. 3a). We measured 1,863 statistics of DNA unzipping from repeated pulls of 538 molecules in a single sample to demonstrate the large amounts of single-molecule force data that can be accumulated with this approach (Fig. 3b). Even more significantly, the nanoswitches enable the unique properties of each molecule in a sample to be characterized from repeated measurements, as we demonstrate by determining a rupture-force histogram for each molecule in a sample, illuminating population heterogeneity at the single-molecule level (Fig. 3c). Furthermore, we show that by averaging data from multiple pulls of the same molecular pair the spread in force is reduced without losing the unique characteristics of each molecule. When applied to data from a single population, this per-molecule force averaging generates a super-resolved histogram with the expected narrowing when compared with the raw histogram of all the data (Fig. 3d). When applied to combined statistics from two populations of DNA zippers (introducing another G–C-rich zipper with a higher unzipping force), we show that the super-resolved histogram generated from per-molecule averaging can separate out two populations that are unresolvable from the raw histograms due to the intrinsic broadening of force that results from thermal noise and instrumental noise (Fig. 3e).

Discussion

As we have shown, together the CFM and the DNA nanoswitch assay provide powerful single-molecule capabilities, and give non-specialists unprecedented access to force spectroscopy. And even separately these technologies provide unique benefits that can make single-molecule experiments better, cheaper and faster. Our miniature CFM presented here, which integrates into standard benchtop centrifuges, has important new features including an expanded biologically relevant dynamic force range (Supplementary Fig. 6), easily implemented temperature control between 4 and 37°C, improved safety, lower cost (Supplementary Table 1 and Supplementary Note 3), and greater ease of use compared with the first-generation CFM. These last two points are important strides towards increasing the accessibility of single-molecule experiments for non-specialists. The total reduction in cost is particularly significant when considering the additional infrastructure typically needed for single-molecule experiments for non-specialists.

Figure 2 | DNA unzipping force measured using DNA nanoswitches on the CFM. (a, i) Schematic of the unzipping construct. Two complementary oligos (red and purple strands) hybridize to form a looped DNA nanoswitch. Force can unzip the two complementary strands, resulting in a measurable increase in tether length, providing a signature of DNA unzipping. (a, ii) Images of a bead tethered to the surface via a DNA nanoswitch showing the looped and unlooped states. The scale bar, 1 μm long. (a, iii) In one example of rupture-force measurement, we identified 381 tethers with the DNA nanoswitch transitions signature to collect rupture forces while the remaining 673 transitions that corresponded to bead detachment and improper transitions were omitted (see also Supplementary Fig. 5 for detail). (b) Unzipping force histograms of 29 bp dsDNA measured with the DNA nanoswitch under two different buffer conditions. (c) Average unzipping force of 29 bp dsDNA under different temperatures with PBS buffer (Total n = 306), with histograms of rupture forces shown as an inset. The theoretical line is calculated using a previously described thermodynamic model.
measurements that are obviated by our approach, as the centrifuge itself provides an isolated, temperature-regulated enclosure. In addition, our instrument can be operated without significant training, even by undergraduate researchers. Similarly, the DNA nanoswitch-based single-molecule assay has its own benefits, significantly reducing the effort to obtain reliable and accurate measurements by standardizing sample preparation, and by supplying a distinct molecular signature that enables reliable data verification and automated analysis. Furthermore, it enables repeated interrogation of each molecular complex for building up statistics of both the population and the individual molecules, revealing population heterogeneity at the single-molecule level.

There is difficulty in generalized comparisons of single-molecule instruments due to the wide variety of designs, but the miniature CFM presented here (using the 2.8 µm Dynabeads) is comparable to our own optical tweezers set-up in force range (~0.1 to 100 pN), force resolution (~3–5%) and spatial resolution (~2 nm). It is worth noting, however, that the most advanced optical tweezers push spatial resolution into the sub-Angstrom range, have force resolution of <0.1 pN and have temporal resolutions >10 kHz (this CFM is 15 Hz)². Regarding multiplexing, magnetic tweezers have achieved up to 145 statistics in a single experiment⁶ and other multiplexing methods show great promise as well², twelve, thirteen. However, our demonstration of 1,863 validated single-molecule statistics in one experiment is to our knowledge more than in any previous work.

As the potential for single-molecule approaches to address critical problems in biology becomes increasingly clear, the availability of such simple and powerful tools that do not require months of training or hundreds of thousands of dollars becomes a key. We believe the technologies presented here are an important step in that direction, enabling both a more detailed view of single molecules as part of a population, and a more streamlined way to amass large numbers of statistics. It is not unreasonable to imagine that this approach, paired with sample scanning and/or fast force cycling could potentially enable the collection of millions of single-molecule statistics from a single sample. With these advances, previously unrealistic applications such as high-throughput single-molecule screening and diagnostics become more viable, and single-molecule analysis may start to become as ubiquitous as standard bulk assays.

**Methods**

**CFM instrumentation.** The CFM consists of an optical microscope and digital image acquisition system, which is integrated into a centrifuge (Fig. 1a and Supplementary Fig. 1). We used a refrigerated benchtop centrifuge (Heraeus X1R, Thermo Scientific) with two modifications. First, the TX-400 rotor was modified to mount a fibre-optic rotary joint along the central axis (Princetel MJX). This was accomplished by removing the rotor’s central push-release mechanism (by loosening the screw on the side of the button), and threading the four existing
holes to access 10–32 screws. An adapter was designed and installed on the rotor to hold the slip ring (Supplementary Fig. 1a,b). To enable the fibre optic to pass through the lid, the central plastic viewing window was removed. Second, the centrifuge’s control module was upgraded to enable computer control. An upgraded module was kindly provided by Thermo Fisher Scientific to support this project, and was a simple drop-in replacement.

The light microscope of the benchtop CFM was constructed using mainly Thorlabs components (Supplementary Fig. 1b,c,e and Supplementary Table 1). A red LED (LED630E, Thorlabs) threaded to a tube mount (SILEDM, Thorlabs) served as the illumination source. A glass diffuser positioned between the LED source and sample cell provided uniform illumination across the entire field of view. A 300 µm diameter single-cell sample was compatible with the SM1 lens tube (see below CFM sample cell). The sample was magnified and imaged onto a CCD camera (GC 2,450, Prosilica, AVT) with a Thorlabs compatible with the SM1 lens tube (see below CFM sample cell). The sample was a simple drop-in replacement.

For the overstretching experiment, the surface tethering was strengthened by replacing digoxigenin–anti-digoxigenin with biotin–streptavidin—in other words, biotin–streptavidin interactions were used to anchor both ends of each tether. The nitrocellulose surface was functionalized by incubating it with 1 mg ml⁻¹ streptavidin in PBS solution for 30 min, followed by incubation with 10 mg ml⁻¹ Roche Blocking Reagent for 12 h, followed by incubation with passivation solution (10 mg ml⁻¹ Roche Blocking Reagent in PBS buffer) for 1 h. The channel was then washed and incubated with 5 µl of dual-biotin X-DNA for 15 min before loading in the streptavidin-coated beads. Under such conditions, the density of streptavidin on the surface was sparse enough that only one end of the biotin-labelled X-DNA was bound to the surface, leaving the other biotinylated end free to bind to the streptavidin-coated bead.

**Swinging window angle measurement.** To measure the angle of the swinging window relative to the axis of rotation, we attached an oil-based marker to the bottom of the bucket, which marks the height of the bucket on the wall as the centrifuge spins (Supplementary Fig. 2d). The uncertainty of the angle measurement was based on the distance measurement error estimate of 1 mm. At the rotational speed of 300 r.p.m. the bucket swings out to an angle of (81.4±0.8)°. At a much higher speed of 1,800 r.p.m. the angle increases by 2.3%. Further increase of the rotational speed does not increase the angle beyond the error of the measurement.

**Data analysis.** The loop opening signature was identified by tracking the beads’ x and y positions. As the angle of the centrifuge swing bucket was not 90° with respect to the axis of rotation, a component of the centrifugal force was directed in the x-y plane (Supplementary Fig. 5). As the rotational speed of the CFM increased, the looped DNA tethers were continuously extended. The opening of the loop was identified as a discontinuous change in extension. Each movie contained ~1,000 beads. To track these beads, they were first identified using the Matlab function imfindcircles. A template image for each bead was stored. To identify the bead in the subsequent frame, the template image was scanned in the x–y plane to find the position of maximum correlation. First the image was scanned in the x direction over a 25 pixel search region centred on the bead position from the previous frame. A 2nd order parabola was then fit to the correlation coefficient as a function of position. The position of maximum correlation was identified as the new bead position. The template image was then centred on the new x position, and the same procedure was done in the y direction. During the course of each experiment, there was some drift in the x–y plane. This was corrected for by taking the average of the change in x and y from the previous frame. When the x and y movements are the same, we found this drift correction to be sufficient for identifying the looped to unlooped transition. Transitions were identified by filtering out all bead trajectories except those that contained a discontinuous change in extension of both the correct magnitude and direction (Supplementary Fig. 5b,c). This removed false transitions that may have resulted was published in 0.7 mm diameter ports were drilled that served as the solution inlet and outlet. The cover and support glasses were cleaned by immersing in 100 ml a 1% (v/v) Hellmanex III solution, microwaving for 1 min, then sonicating for 30 min. Subsequently, the slides were rinsed thor- oughly with Millipore water. The CFM sample cell was a stainless steel hollow cylinder with a circular hole in one side and biotin on the other.

**CFM sample cell.** The mini-CFM sample cell was constructed using double-sided Kapton tape sandwiched between a 25 mm diameter support glass and a 19 mm diameter cover glass (3346, General Motor). The 0.7 mm diameter ports were ordered from S.J. Howard Glass (D263) and two 1 mm diameter ports were drilled that served as the solution inlet and outlet. The cover and support glasses were cleaned by immersing in 100 ml a 1% (v/v) Hellmanex III solution, microwaving for 1 min, then sonicating for 30 min. Subsequently, the slides were rinsed thoroughly with Millipore water. The CFM sample cell was a stainless steel hollow cylinder with a circular hole in one side and biotin on the other.

For DNA unzipping experiments, the number of transitions as a function of time was converted to the number of transitions as a function of force as follows:

\[
\text{Number of transitions} = \frac{\text{Number of DNA unzipping experiments}}{\text{Time}} \times \text{Force} 
\]
The rotational speed of the centrifuge was recorded during each movie using WinMess software (provided by Thermo Fisher Scientific, R&D) which enables communication and control with the computer. The rotational speed was then converted to force (F) as a function of time using the following equation:

\[ F = m_{\text{sample}} \times \omega^2 \times \frac{R_{\text{sample}}}{r^2} \]

where \(R_{\text{sample}}\) and \(\omega\) are the rotational radius and speed, respectively. To determine the effective mass of the beads (\(m_{\text{sample}}\)) in solution, we measured the Invitrogen M-270 bead density through sink-float analysis using aqueous sodium polytungstate solution (71913, Sigma-Aldrich) and obtained a density of 1.61 ± 0.02 g cm\(^{-3}\) that confirmed the manufacturer’s (Life Technologies) reported value of 1.6 g cm\(^{-3}\). The manufacturer also provided the bead diameter of our specific lot giving a mean of 2.80 μm with <1.6% CV (lot #144315600). The density of the silica beads used in the DNA force extension and overstretching experiment was measured similarly using the sodium polytungstate solution, yielding a density of 1.50 ± 0.03 g cm\(^{-3}\). The diameter of the silica beads was measured using transmission electron microscopy, yielding an average and s.d. of 4.27 ± 0.16 μm.

Tethered particle motion. We carried out tethered particle motion analysis for our tethered beads prior to the rupture-force measurement. We analysed the lateral fluctuations of each bead, calculating the root mean square of the drift-subtracted displacement and the asymmetry ratio, following previously established methods. Here the asymmetry ratio was calculated as the square root of the ratio between the maximum and maximum eigenvalues of the covariance matrix for the in-plane displacement. The in-plane position of each bead was recorded for 10 s at an acquisition rate of 10 Hz.

Repeated cycles of rupture-force measurement. Multiple rupture events were collected for each molecule in a set by repeatedly spinning the same sample multiple times. For the data presented in Fig. 3, the sample was spun up with an effective force-loading rate of 1 pN s\(^{-1}\) to a maximum force of 22 pN. The speed was then ramped down to 0 rpm for ~1 h to allow rebinding between each pair of molecules. Beads were identified from cycle to cycle by their positions relative to a set of fiducial beads, which were common to each cycle. For the measurement of the two populations of molecules, we made two different DNA nanoswitch unzipping constructs, one with 48% GC content (5'-CACGAATTCTCTGACCTCTCCTTGTAAACCGTGTAG-3') and one with 31% GC content (5'-CCTCAATATCACAACCTCAGTAT-3').

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