Gold rotor bead tracking for high-speed measurements of DNA twist, torque and extension

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Single-molecule measurements of DNA twist and extension have been used to reveal physical properties of the double helix and to characterize structural dynamics and mechanochemistry in nucleoprotein complexes. However, the spatiotemporal resolution of twist measurements has been limited by the use of angular probes with high rotational drag, which prevents detection of short-lived intermediates or small angular steps. We introduce gold rotor bead tracking (AuRBT), which yields >100× improvement in time resolution over previous techniques. AuRBT employs gold nanoparticles as bright low-drag rotational and extensional probes, which are monitored by instrumentation that combines magnetic tweezers with objective-side evanescent darkfield microscopy. Our analysis of high-speed structural dynamics of DNA gyrase using AuRBT revealed an unanticipated transient intermediate. AuRBT also enables direct measurements of DNA torque with >50× shorter integration times than previous techniques; we demonstrated high-resolution torque spectroscopy by mapping the conformational landscape of a Z-forming DNA sequence.

Single-molecule tracking experiments can yield rich information about the structural dynamics of a molecular system but are fundamentally limited by Brownian noise1,2, which must be overcome to resolve discrete transitions on biologically relevant timescales. To analyze structural transitions in DNA-protein complexes, several groups have introduced real-time methods for simultaneously measuring two broadly relevant structural properties3: changes in DNA contour length and changes in linking number (∆Lk) resulting from trapped writhe4,5 or topoisomerization6. Direct measurements of ∆Lk rely on observing the rotation of a probe attached to the DNA molecule6,7; the spatiotemporal resolution of these measurements depends on the rotational hydrodynamic drag of the probe3. To date, the highest-resolution angular measurements have been achieved using rotor-bead tracking (RBT)3,6, which takes advantage of probe specialization: a micrometer-scale magnetic bead is used to apply piconewton-scale forces to stretch the DNA and a lower-drag secondary probe bound to the side of the molecule (the ‘rotor bead’, typically a ~300 nm polystyrene sphere) reads out molecular twist and extension. RBT achieves subsecond relaxation times. However, angular steps smaller than half a rotation (∆Lk = 0.5) have required several seconds to distinguish from noise6. Currently, RBT and alternative methods such as freely orbiting magnetic tweezers (FOMT)12 are thus inadequate for resolving unitary steps and substeps in the unperturbed millisecond-to-second timescales typical of many biological processes6,15,16.

Torque resolution suffers from a closely related limitation17. Direct torque measurements18 have been used to quantify the torsional rigidity of DNA19–21, study the plectonemic buckling transition20–24 and analyze sequence-dependent structural transitions22,25. However, current methods require seconds of integration at a minimum, and typically much longer times, to achieve biologically relevant torque precisions on the order of 1 pN nm (refs. 3,17,22,25). In the Brownian-limited regime, drag is the only parameter that influences torque-noise density3. Torque-measurement methods that use low-drag probes are thus needed for high-resolution torque spectroscopy and for studies of torque generation by molecular motors that unwind26, supercoil27 or helically track14 DNA.

Here we exploited probe specialization to achieve large gains in spatiotemporal resolution over RBT. Using 80–140 nm gold particles as rotors, AuRBT reduces rotational drag by orders of magnitude over existing methods3,6,12,20–22,25,27, enabling torque and twist measurements on the millisecond timescale and reduced Brownian noise over longer timescales. Our method includes a simultaneous readout of molecular extension using evanescent nanometry28,29, which provides a second degree of freedom with which to monitor nucleoprotein dynamics. We illustrate AuRBT’s enhanced spatiotemporal resolution with three experiments. First, we demonstrate the detection of subtle helicity changes induced by changes in DNA tension owing to twist-stretch coupling. Next, we present a high-resolution study of DNA gyrase, which revealed a previously unknown structural substate. Finally, we use AuRBT torque spectroscopy to map the B-Z transition in a short DNA sequence and show that the increased torque resolution provides...
the sampling power to measure torque probability distributions for meaningful comparisons to theoretical predictions.

RESULTS

Combined evanescent scattering and magnetic tweezers

AuRBT uses darkfield microscopy to track the three-dimensional (3D) position of a gold nanoparticle that is bound to the side of a stretched DNA molecule (Fig. 1 and Supplementary Video 1). Gold nanoparticle tracking has previously been implemented using a variety of darkfield configurations, including designs that rely on laser illumination through a condenser and objective-type evanescent excitation. Here we chose evanescent excitation so that a nanoparticle can be imaged adjacent to the surface without contamination from scattering by the larger magnetic bead attached to the opposite end of the DNA molecule. We chose objective-side illumination to accommodate magnetic tweezers on the condenser side of the sample. In a departure from previous work, our implementation uses two small mirrors positioned below the objective to couple laser excitation both in and out of the sample at high numerical aperture (Supplementary Figs. 1 and 2), which is similar to a design previously used for multiwavelength fluorescence measurements. The scattering image is collected on an electron-multiplying charge-coupled device (EMCCD) camera at high frame rates (up to 6.3 kHz); the instantaneous angle can then be determined from in-plane coordinates of the rotor (Fig. 1b), and vertical displacements of the rotor can be calculated using evanescent nanometry (Supplementary Fig. 3).

Small probes reduce integration time by orders of magnitude

AuRBT detects small twist signals by rapidly averaging over Brownian noise. Using rotor beads with radii several times smaller than those used with previous methods reduced drag by nearly two orders of magnitude, which enabled us to achieve angular diffusion constants in excess of 1,000 rad² s⁻¹ orders of magnitude, which enabled us to achieve angular diffusion constants in excess of 1,000 rad² s⁻¹. The highest-resolution measurements we show here used 80-nm rotor beads and 420-base-pair (bp) DNA tethers (referred to as AuRBT (diameter (d) = 80 nm, length (L) = 420 bp)), which exhibited a ~1 ms relaxation time and enabled precise twist determination >200× faster than can be achieved with currently published techniques.

For Brownian noise–limited DNA twist measurements, the signal-to-noise ratio (SNR) depends on the extent to which fluctuations can be reduced from their full-bandwidth amplitude of \((k_B T/\kappa)\)¹/² given by equipartition, where \(k_B\) is the Boltzmann constant, \(T\) is temperature and \(\kappa\) is the torsional stiffness of the tether. Analogous considerations have been extensively reviewed for the case of extension measurements. Angle measurements are correlated over the relaxation time \(t_r = \gamma/\kappa\), where \(\gamma\) is the drag of the probe. Fast relaxation times permit rapid sampling of fluctuations; therefore, in any given time interval, higher SNR can be achieved by either reducing drag or stiffening the tether. The expected root mean square (RMS) noise \(\sigma\) for a signal integrated over time \(t\) (Fig. 2a) is given by:

\[
\sigma = \sqrt{\frac{2k_B T}{\kappa} \left( t - \left( \frac{t}{t_r} \right)^2 \left( 1 - \exp \left( -\frac{t}{t_r} \right) \right) \right)}
\]  

Figure 1 | AuRBT design and data collection. (a) For AuRBT, an infrared laser is introduced using a small mirror to achieve total internal reflection. The return beam is extracted using a second mirror, and stray light is rejected with an iris. The darkfield image of a gold rotor bead is imaged on a high-speed camera. (b) A rotor bead was attached above a 420-bp torsionally constrained DNA segment and held under 10 pN of tension. The x-y position was recorded at 6.3 kHz and accumulated over 150 s to generate a 2D histogram, which shows restriction to an annulus and a preferred equilibrium angle. The instantaneous in-plane angle \(\phi\) can be calculated from the \(x\) and \(y\) measurements. The measured diameter \(d\) of the orbit for this bead (black circle) was 69.4 nm, which is a measure of the size of the particle. (c) Simultaneous angle and extension measurements of a rotor \((d = 68.1\text{ nm})\) attached above a 420 bp torsionally constrained segment and held under 26 pN of tension by a magnetic bead dimer. Extension was measured using evanescent nanometry (Online Methods). Full-bandwidth traces are shown along with 50 Hz and 1 Hz lowpass filtered data.

Measurements of RMS noise as a function of integration time confirm the expected performance of AuRBT (Fig. 2). The relaxation time \(t_r\) for AuRBT ranged from ~1 ms to 10 ms depending on the bead and tether chosen (Supplementary Table 1). The large \(t\) limit \((t >> t_r)\) illustrates how spatiotemporal resolution is affected by drag:

\[
\sigma = \sqrt{\frac{2k_B T \gamma}{\kappa^2 t}} = \frac{2k_B T \gamma}{\kappa^2 \sigma^2}
\]  

To detect a given step size with a given SNR, the required integration time thus scales linearly with drag, which in turn depends on the curve of the probe diameter. AuRBT provides biologically relevant gains in resolution. For example, previously published techniques would require >20 s of integration to detect the angular signal corresponding to unwinding of a single base pair (\(|\Delta L| = 0.095\)), such as could result from unitary steps of RNA polymerase. Using AuRBT, this measurement can be performed in ~80 ms with SNR of 3 (Fig. 2a). As an alternate representation of resolution comparisons, we show noise power spectral density plots (Supplementary Fig. 4).
We confirmed cubic scaling of drag with particle size by plotting all measured angular diffusion constants as a function of measured bead diameters (Fig. 2c). A global fit to the approximate expected relationship $D = \eta_{\text{eff}} T/(14\pi r^3)$ gives an effective viscosity $\eta_{\text{eff}} = 1.2$ mPa s, which is slightly higher than the viscosity of the bulk solution, as may be expected due to the proximity of the surface (Supplementary Fig. 5).

Rapid measurement of twist-stretch coupling in DNA

We challenged the high twist resolution of AuRBT by measuring a known physical property of the double helix: DNA overwinds when stretched. This small effect has been previously observed using RBT, relying on long integration times\(^{40}\). No subsequent studies have measured twist-stretch coupling in the freely fluctuating twist ensemble, although several groups have determined the sign and magnitude of the elastic coupling term by measuring changes in extension upon overwinding in the fixed twist ensemble\(^{40–42}\). We used AuRBT with 80-nm beads to perform rapid observations of twist-stretch coupling in a 4,130-bp DNA segment (Fig. 3). We torsionally constrained the DNA tether at the magnetic bead, but it was free to swivel at the coverslip, so that the rotor bead reported the twist of the upper segment (Fig. 3a). We increased force in ~1.5-pN steps from ~4.5 pN to ~13.5 pN, which induced stepwise helicity changes of ~0.01%, as expected based on previous results\(^{40–42}\). Helicity changes were well separated from Brownian noise using only 60 s of integration (Fig. 3b,c). This experiment may be compared with that in ref. 40 figure 1c, in which ~10x longer integration times had been used to resolve changes in twist induced by ~3x larger steps in force.

AuRBT reveals a transient DNA gyrase intermediate

As a representative system for AuRBT measurements of nucleoprotein complexes, we investigated the structural dynamics of DNA gyrase, an essential bacterial molecular motor that harnesses ATP hydrolysis to introduce supercoils into DNA\(^{43}\) (Fig. 4). Gyrase has been previously studied using RBT\(^{6,7}\), and substeps in its mechanochemical cycle have been characterized by analyzing changes in angle and extension (z) at limiting [ATP]\(^6\). Single-molecule gyrase traces are characterized by proessive bursts of activity\(^{6,7}\), in which each enzymatic cycle introduces two rotations owing to duplex-strand passage\(^6\). The dominant kinetic dwell in the cycle occurs in the Ω state, in which >100 bp of DNA contour length are sequestered, but no supercoils are trapped in the complex\(^6\). An ATP-accelerated remodeling transition converts Ω into a chirally wrapped intermediate, dubbed the α state, which can be detected as a rotational substep at low [ATP]\(^6\). The chiral wrap is critical for guaranteeing that subsequent strand passage will directionally introduce supercoils, differentiating gyrase from other members.
of the type II topoisomerase family. Post–strand-passage states are presumed to be short-lived and had not been detected in prior RBT experiments under any conditions. At saturating [ATP], the α state was also undetectable, and even the dominant Ω dwell could not be visualized in every cycle. Spatiotemporal resolution has thus been a limiting factor in achieving a complete understanding of DNA gyrase mechanochemistry.

We used AuRBT with 140-nm beads and 420-bp tethers to study DNA gyrase at high resolution, which yielded well-resolved Ω dwells even at saturating [ATP]. Rapid transient excursions to supercoil-trapping states (previously seen only at limiting [ATP]) were also visible in the angle traces (Fig. 4b), which suggested a more dynamic picture of gyrase mechanochemistry than has been previously appreciated. Finally, simultaneous angle and z tracking revealed a previously undetected intermediate: the otherwise contracted z signal often exhibited transient (±10–150 ms) excursions toward zero; these excursions typically occurred in between Ω dwells in successive cycles. To display the angle-z dynamics, we used a color scale to represent z.

**Figure 3** High-resolution measurement of twist-stretch coupling. (a) ‘Top-constrained’ experimental geometry for measuring changes of twist in a long DNA segment. (b) Full-bandwidth (color) and filtered (black) angle traces for a DNA molecule, acquired during stepwise increments of tension exerted every 60 s. Dashed line indicates extrapolated mean twist at zero force. (c) Mean twist as a function of force, taken from the data shown in b, showing the linear increase expected for a negative twist-stretch coupling coefficient. Error bars show the estimated RMS twist noise (see Fig. 2b) for the integration time \( t = 120 \) s (totaled over two 60-s dwells at each force).

**Figure 4** High-resolution analysis of gyrase dynamics at 1 mM ATP. (a) Schematic of ‘bottom-constrained’ experimental geometry. During gyrase activity, the angle and height of the bead reflect changes in Λk and extension of the lower DNA segment. Measurements were performed using 140-nm rotors under 1.1 pN of tension, and angle and z were both low-pass–filtered to 50 Hz. (b) Excised traces of single-enzyme bursts showing angle as a function of time, with color indicating the instantaneous value of relative DNA extension (z). Dominant dwells are visible every two rotations as expected. Short regions flanking each burst show the extension of free DNA. *, events in which the DNA extension briefly increases, primarily coinciding with entrance into a new rotational dwell. (c) A single gyrase trace, projected along angle and z axes; brief excursions (*) are visible between dominant dwells. (d) 2D histogram of angle and z coordinates accumulated over a total of 87 gyrase cycles belonging to 19 enzymatic bursts on five separate DNA tethers; 1.5 s of data flanking each burst were included to show the position of free DNA. (e) Proposed model explaining excursions in extension (*). DNA contour length is released from one or both C-terminal domains (CTDs) after strand passage, as the enzyme enters a newly defined Ω state. The DNA is then recaptured to reset the enzyme for the next cycle.
Figure 5 | AuRBT torque spectroscopy. (a) Schematic of ‘doubly constrained’ experimental geometry for static torque measurements. Total twist \( \theta \) can be varied by rotating the magnets, and torque is measured from the angular deflection of a calibrated transducer segment, \( \tau = k \Delta \theta \). A sequence of interest can be placed in the lower segment, such as the 50 bp GpC repeat used here to examine Z-DNA formation. (b) Torque resolution of AuRBT and of published alternate methods for measuring DNA torque: RMS torque noise was measured as a function of integration time, using 4,130-bp transducer segments assayed in a ‘top-constrained’ geometry. (80-nm rotor results are plotted from the same data set as AuRBT–80-nm rotor, 4,130-bp tether in Fig. 2a.) Corresponding power spectral density plots are shown in Supplementary Figure 6. Models (equation (3)) were parameterized from spectral analysis for AuRBT and RBT and based on reported parameters for eMTT20 and OTW4. The dashed horizontal line indicates the RMS torque noise required for measuring a 1 pN nm change with SNR of 3, and its intersection point with each curve indicates the necessary duration of signal averaging. (c) Torque as a function of imposed twist for a DNA molecule containing a 50-bp GpC repeat. Similar traces were obtained for a total of four molecules under two buffer conditions. AuRBT was used with a 110-nm rotor under 5 pN of tension, and twist was ramped linearly at 0.05 rotations/s. Low-pass–filtered data for four individual rewinding curves on the same molecule and full-bandwidth data (6.3 kHz) are shown. A statistical mechanical model for the B-Z transition22,25 was used to fit each data set. Model prediction (Fig. 4b) for measuring the torque on a single DNA molecule 17–22,24,27. (d, e) Distributions of measured torque as a function of imposed twist. The expectation torque curve from \( \kappa \) was replotted in black. Combined full-bandwidth data from all traces in \( \kappa \) were binned in order to plot \( P(\kappa) \) (heat-mapped) for each value of twist (d). Predicted \( P(\kappa) \) was calculated (equations 9–12 in Supplementary Note 1) based on the parameters determined from fitting the expectation torque curve, and plotted for comparison with experiment (e).

Probing ensembles with high-resolution torque spectroscopy

Over the past decade, a number of methods have been introduced for measuring the torque on a single DNA molecule. Among them, the optical torque wrench (OTW) uses a polarized optical trap to measure and apply torques. Magnetic tweezer (MTT) and electromagnetic torque tweezers (eMTT) uses a vertical field for force application, with a small fixed (MTT) or controllable (eMTT) horizontal component to measure and apply biologically relevant torques. Soft magnetic tweezers differ in implementation but also measure displacement in a weak magnetic potential as do nanorod magnetic tweezers. Static RBT uses the deflection of a calibrated ‘transducer’ DNA segment to measure torque in the molecule. For any torque measurement technique that measures the displacement of a rotational probe with drag \( \gamma \) in a harmonic potential with stiffness \( \kappa \), the RMS torque noise is given by an expression closely related to equation (1):

\[
\sigma_T = \kappa \sqrt{\frac{2k_BT}{\kappa} \frac{t_r}{t} \left( \frac{t_r}{t} \right)^2 \left( 1 - \exp \left( -\frac{t}{t_r} \right) \right)}
\]

(3)

where \( t \) is the integration time and \( t_r \) is the relaxation time, as before. In the limit of long integration times, this expression depends only on the rotational drag:

\[
\sigma_T \approx \frac{2k_BT\gamma}{t}
\]

(4)

We reasoned that we could make precise DNA torque measurements on much shorter timescales than using previously published methods, by adapting the ‘static RBT’ method to use gold nanosphere probes (Fig. 5). We characterized the fluctuations of gold nanosphere rotors attached to DNA transducers to confirm the expected gains in torque resolution (Fig. 5b, Supplementary Fig. 6 and Supplementary Table 2). We saw up to 55x reductions in required integration times relative to previous static RBT measurements and even larger improvements relative to other techniques.

To test the performance of static AuRBT on a well-characterized molecular system, we examined the B-Z transition in DNA molecules containing a 50 bp GpC repeat sequence of interest.
This sequence (here referred to as Z50) displays a localized transition from right-handed B-DNA to left-handed Z-DNA under negative supercoiling. As expected, we observed a characteristic torque signature that was well-described by a statistical mechanical model similar to the Ising model at fixed magnetization. Upon unwinding, the B-DNA molecule initially builds up torque linearly. Additional unwinding causes an abrupt torque jump that corresponds to the formation of an initial Z-DNA domain. The domain can then be extended at approximately constant torque until the entire sequence of interest is converted to Z-DNA, which restores linear elastic behavior.

AuRBT revealed the mean torque as a function of imposed twist at higher resolution than reported earlier and validated the previously proposed model with high fidelity (Fig. 5c). Additionally, the improved sampling allowed for a new kind of comparison between theory and experiment for this transition. Based on the best-fit parameters obtained from the averaged torque alone (Fig. 5c), we used the statistical mechanical model (Supplementary Note 1) to predict the entire distribution of measured torques at each value of total twist. This distribution can be compared with the experimental distribution obtained for each sampled twist value, and this comparison showed excellent agreement between theory and experiment. We show the comparison in 2D histograms (Fig. 5d,e) and in an animated plot (Supplementary Video 2).

**DISCUSSION**

Two major goals that drive the development of single-molecule approaches are to directly observe dynamic transitions and fleeting intermediates in actively functioning biomolecular complexes and to characterize distributions of molecular properties, as opposed to ensemble averages. In practical single-molecule manipulation experiments, slow Brownian fluctuations can interfere with both of these goals. For measurements of DNA twist and torque, AuRBT provides the spatiotemporal resolution to detect intermediates, and the sampling power to define distributions. AuRBT allows biologically relevant angular displacements to be resolved on previously inaccessible timescales that are relevant to enzymology. For example, the twist signal expected from unwinding of a single DNA base pair can now be detected with less than 100 ms of integration, and larger changes can be resolved on ~1-ms timescales. When different measurement coordinates are available for detecting the same biomolecular event, direct angular measurements can now sometimes be the most sensitive alternative: for example, indirect measurements of ΔLk via plectonemic amplification require 1 s of integration or longer to discriminate single-base-pair differences in unwinding. Published high-resolution optical tweezers assays of DNA translocating motors require ~1 s of integration to discriminate changes in extension arising from single-base-pair steps. For the subset of DNA translocating motors that track the helical groove, AuRBT may be used to detect single-base-pair steps on shorter timescales, by relying on the angular coordinate instead.

AuRBT offers distinct advantages and some tradeoffs compared to other methods for direct measurements of DNA twist and torque. AuRBT should be considered when high resolution is paramount: for example, AuRBT enables precise twist measurements using >50,000× shorter integration times than FOMT and precise torque measurements using >500× shorter integration times than OTW.

For twist measurements, FOMT has the compensatory advantage that it can be performed with minor modifications to many existing magnetic tweezers setups and that a single molecule can easily be assayed successively in fixed-angle and freely fluctuating modes. Additionally, twist assays with larger probes, including FOMT and RBT, can generally be performed at lower forces than AuRBT, which relies on tension to suppress lateral fluctuations (Supplementary Fig. 7). The highest-resolution AuRBT measurements reported here required a minimum of ~5 pN of tension, although we performed high-resolution assays of the tension-sensitive DNA gyrase enzyme (using 140-nm gold rotors) at ~1 pN. Similar tradeoffs between tension and resolution are seen in high-resolution optical trapping experiments, which require high tensions (typically >10 pN) to reduce tether compliance.

For torque measurements, OTW can be advantageous for some experiments because it also includes a simultaneous dynamic high-resolution force measurement. In addition, OTW (and related techniques) can be performed over a wide range of torques, whereas static AuRBT is restricted to the torque range over which the B-DNA transducer displays linear twist elasticity. This important limitation might be overcome in the future by using stable transducers constructed from DNA origami, expanding the accessible torque range.

AuRBT contributes to new single-molecule approaches in which high-resolution measurements of multiple degrees of freedom provide detailed dynamic pictures of molecular processes. AuRBT combines direct measurements of angle with high-bandwidth extension measurements (Supplementary Fig. 8), which we exploited to discover a previously unknown structural intermediate in DNA gyrase, opening the door for further dissections of the gyrase mechanism and illustrating the potential for investigating other complex DNA-protein machines. Other recent high-resolution multiparameter methods have included an elegant combination of Ångstrom-resolution optical trapping with single-molecule fluorescence. In principle, there are no barriers to augmenting AuRBT with single-molecule fluorescence as well: AuRBT measurements are based entirely on near-infrared scattering, which leaves the visible spectrum free for future experiments in which fluorescent probes report synchronously on internal degrees of freedom. High-resolution and multidimensional single-molecule methods will help bridge the gap between atomically detailed crystal structures and dynamic functional assays, illuminating the complete conformational landscapes of molecular machines at the heart of biology.

**METHODS**

Methods and any associated references are available in the online version of the paper.

**Note:** Any Supplementary Information and Source Data files are available in the online version of the paper.

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ONLINE METHODS

Instrumentation. Experiments were performed on a modified Nikon Eclipse Ti-S inverted microscope (Supplementary Fig. 1). Samples were mounted on a three-axis nanopositioning stage (Mad City Labs PDQ-series). Excitation was provided by an intensity-stabilized 845-nm laser diode (Lumics LU0845M200) directly coupled to a polarization-maintaining fiber; laser power fluctuations were ±0.1%. Laser power used (measured after the fiber) ranged from 2.5 mW to 80 mW; the highest powers are required for the smallest beads. Under these conditions, nanoparticle heating and optical forces are generally small (Supplementary Note 2), and there was no evidence of optical damage, with many molecules lasting for hours of observation. The laser was collimated into a ~2 mm beam and then focused onto the objective back focal plane (Fig. 1 and Supplementary Figs. 1 and 2) using a 500 nm achromatic lens. A half-wave plate was used to achieve s-polarization at the sample interface. The return beam was collected on a position-sensitive detector (Supplementary Fig. 1) to provide a signal for focus stabilization, which was controlled by a Matlab callback function implementing proportional-integral gain at 1 Hz (Supplementary Fig. 9). Scattered light was collected with a Nikon Apo TIRF (60×/1.49 numerical aperture, oil) objective and imaged through an optical path splitter (Cairn, Optosplit II) and onto an EMCCD camera (Andor Ixon+) equipped with liquid cooling (Koolance Exos-2).

Magnetic tweezers were used to apply calibrated stretching forces. A pair of 0.25 inch × 0.25 inch × 0.5 inch rectangular neodymium magnets (K&J Magnetics B448) was held above the sample in a machined aluminum mount with dipoles in opposing directions (Fig. 1a), as described previously. A hollowed shaft connected the magnets to a rotary servomotor (Physik Instrumente C-150.PD), which was mounted on a vertical servomotor (Physik Instrumente C-150.PD1) to enable rotation about and translation along the optical axis. Collimated light from a superbright LED (Thorlabs M660F1) entered the sample from above through a 0.6 mm gap in the magnets, providing illumination for 3D ring tracking of magnetic beads during force calibrations.

A complete list of parts for AuRBT is provided in Supplementary Note 3.

Preparation of DNA, beads and protein. DNA tethers for rotor bead tracking were prepared by ligation of restriction enzyme digested PCR products (or hybridized oligonucleotides in the case of the Z50 sequence of interest), as in previous work. The details of tether construction are shown in Supplementary Table 3. Magnetic beads were prepared by cross-linking 1 μm carboxyl-modified superparamagnetic beads (MyOne, Invitrogen) with rabbit anti-fluorescein (A-889, Invitrogen). Rotor beads were neutravidin-coated gold nanospheres (Nanopartz catalog numbers C11-70-TN-50, C11-90-TN-50 and C11-125-TN-50 for ~80 nm, ~110 nm and ~140 nm beads, respectively) or Power-Bind streptavidin-coated polystyrene spheres (Thermo Scientific; 300 nm beads). E. coli GyrA and GyrB subunits were individually expressed and purified as described, mixed to reconstitute tetrarmers, and stored at ~80 °C in 50 mM Tris-HCl, pH 7.5, 100 mM potassium glutamate, 2 mM DTT, 1 mM EDTA and 10% glycerol.

Chamber preparation. Custom flow cells were made of laser-cut paraffin-based (Nescofilm) channels that were melted at 73 °C between a 24 mm × 50 mm No. 1.5 glass coverslip (VWR) and a 24 mm × 50 mm hole-punched vinyl coverslip (Rinzle). Imaging was performed through the glass side, and watertight connections to polyethylene tubing (SAI, PE-10) were made at the ports on the vinyl side by using small rubber o-rings. Tubings were mated with o-rings after being pulled through undersized holes in custom machined aluminum bars, which also facilitated clamping of the o-rings to the surface. The bars compressed the o-rings against the vinyl coverslip ports, sealing the free end of the tubing in a small space near the channel entry port. The active surface of the glass coverslip was spin-coated with 0.1% w/v nitrocellulose (E.F. Fullam) in amyl acetate >30 min before chamber assembly.

To prepare bead-DNA complexes, 25 μl of gold nanospheres were washed by centrifugation in 150 μl of binding buffer (500 mM NaCl, 40 mM Tris-HCl pH 8.0, 0.2% Tween-20, 200 μg/ml BSA, 5 mM EDTA and 0.01% sodium azide), resuspended in 25 μl of binding buffer, mixed with 3 μl of DNA tethers (~50 pM in 10 mM Tris-HCl pH 8.0 and 5 mM EDTA) and incubated overnight.

Channel preparation began by flowing 40 μl of 2 μg/ml antidigoxigenin (Roche) in PBS and incubating for 60 min and subsequently washing with 400 μl of passivation buffer (a one-to-one mixture of 10 mg/ml NEB BSA and a solution containing 1 M NaCl, 80 mM Tris-HCl pH 8.0, 0.4% Tween-20, 10 mM EDTA and 0.02% sodium azide) and incubating for 1–2 h. After channel passivation, the bead-DNA mixture was introduced for 1 h, and then the channel was washed with 400 μl of binding buffer. Finally, 10 μl of magnetic beads in PBS were mixed with 15 μl of binding buffer and introduced for 1 h, followed by an additional wash with 40 μl binding buffer. Twist-stretch coupling and torque spectroscopy measurements were performed in 40 mM Tris pH 8.0, 100 mM NaCl, 5 mM EDTA, 200 μg/ml BSA (NEB), 0.01% sodium azide and 0.2% Tween-20. Gyrase experiments were performed in GB (35 mM Tris-HCl, pH 7.6, 24 mM potassium glutamate, 4 mM MgCl₂, 2 mM DTT, 250 μg/ml BSA (NEB), 0.2 mM spermidine, 0.2% Tween-20 and 1 mM ATP), essentially as described previously.

Selecting DNA tethers for AuRBT. To select a rotor bead assembly for analysis, the coverslip surface was first scanned to visually identify a scattering gold particle colocalized underneath a magnetic bead. A short bead tracking data set was then acquired to check for expected rotor bead behavior. Beads were rejected from further analysis if the x-y positions failed to form an annulus (Fig. 1b), if the beads showed periods of heavily restricted mobility (reflecting surface sticking) or if the angular signal was unconstrained (reflecting DNA nicking or unconstrained attachment). Magnets were then rotated to verify the attachment geometry. For bottom-constrained tethers (as in Fig. 4a), magnet rotation at low force did not cause the magnetic bead to approach the surface; for top-constrained tethers (as in Fig. 3a), the rotor bead angle followed the rotation of the magnets. For doubly constrained tethers (as in Fig. 5a), we rejected molecules if rotation of the magnetic bead did not produce a corresponding rotation of the rotor bead attenuated by the expected partition ratio (the ratio of the lower segment length to that of the entire molecule).
Data acquisition and PSF fitting. Rotor-bead images were recorded at frame rates of 1.9 kHz, 3.4 kHz and 6.3 kHz for 140-nm, 110-nm and 80-nm rotor beads, respectively. The EMCCD’s ‘isolated crop’ mode at 10 × 10 pixels was used in all three cases to achieve the necessary frame rates. A quasi–real-time acquisition and analysis loop was run in Matlab (Supplementary Software) at a loop rate of 1 Hz. During each loop, all available images were downloaded from the camera and immediately fit using a 2D Gaussian fitting function, which was written in C and compiled as a Matlab executable. Fit parameters included an additive offset \( c \), peak height \( A \), \( x \) position \( x_0 \), \( y \) position \( y_0 \), \( x \) s.d. \( \sigma_x \) and \( y \) s.d. \( \sigma_y \).

After fitting, all raw images and fit parameters were written directly to a six-disk redundant array of independent disks (RAID) array, and proportional feedback was applied to the sample stage to stabilize the \( x \)-\( y \) position of the molecule in the center of the crop.

Rotor bead tracking data analysis, calibration and corrections. To calibrate \( x \)-\( y \) trajectories, lateral magnification was determined by scanning a surface-bound gold nanoparticle across a rectangular lattice of \( x \)-\( y \) positions, and tracking its position. Mean row and column separations gave an average magnification of 153×. Further corrections were applied to \( x \)-\( y \) trajectories before calculation of angle. First, residual drift was corrected by subtracting the center of an ellipse fitted to each 1-s interval of data. Next, overall ellipticity was removed using a fit to the entire trajectory (Supplementary Fig. 10). Rotor bead angle was determined by computing the four-quadrant inverse tangent of each corrected pair of \( (x,y) \) coordinates and then unwrapping the result to determine cumulative angle (Supplementary Video 1). Bead sizes were determined by fitting histograms of measured radii to Rician distributions\(^{52} \), in order to extract each orbital radius independent of lateral fluctuations.

Rotor bead heights were determined using evanescent nanometry\(^{28,29} \), computed as \( \Delta z = -\log (I/I_{<1>)} \), where \( I \) is proportional to the instantaneous scattering intensity, and determined by 2D Gaussian fitting \( (I = A \sigma_x \sigma_y) \) and \( <I_0> \) represents the average scattering intensity at a given in-plane angle \( \phi \) (Supplementary Fig. 11). The field’s decay length \( \lambda \) was determined by cross-calibration with dual focus imaging (Supplementary Fig. 3). Evanescent field decay lengths were typically 130–200 nm.

To allow unambiguous angle tracking, lateral fluctuations of the rotor bead were suppressed by tension. Rare zero crossings (in which a bead trajectory passes through the \( x \)-\( y \) origin, causing erroneous jumps of integer rotations) were identified and corrected for using semi-automated analysis in Matlab. Tensions were chosen to give a maximum tolerated zero crossing rate of ~10–20 per million frames; these conditions also ensure that positional noise contributes negligible errors to angle determination (Supplementary Fig. 7).

Force calibration. The average force as a function of magnet height was computed as the product of lateral stiffness along the field direction with the extension of the molecule\(^{53} \). Lateral stiffness was estimated from Lorentzian fits to position noise power spectra\(^{54} \) generated from 70,000 image frames taken at a sampling rate of 1.6 kHz (the maximum observed Lorentzian cutoff frequency of ~50 Hz). At each magnet position, the focal depth of the magnetic bead was determined relative to the feedback-stabilized focal plane by using a lookup table for the radial profile of its diffraction ring pattern as a function of defocus\(^{55} \). Absolute DNA extension was determined by comparing the focal depth of the tethered bead to that of a surface-bound reference bead and correcting for off-center attachments\(^{56} \).

Torque spectroscopy. Static AuRBT torque spectroscopy was performed under 5 pN of tension. Torque-twist curves were acquired by rotating the magnetic tweezers at 18°/s from +5 rotations to −15 rotations and back. The magnet angle corresponding to \( \theta = 0 \) was determined by finding the magnet position which maximized extension of the molecule, as in refs. 22,25. The torsional rigidity of the transducer segment was determined by analyzing the noise power spectra of free twist fluctuations in top constrained molecules, which contained the identical DNA transducer.

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