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Histone-DNA assemblies in archaea : shaping the genome on the edge of life

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CHAPTER 3

Quantitation of DNA binding affinity using Tethered Particle Motion

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Abstract

The binding constant is an important characteristic of a DNA-binding protein. A large number of methods exist to measure the binding constant, but many of those methods have intrinsic flaws that influence the outcome of the characterization. Tethered Particle Motion (TPM) is a simple, cheap and high-throughput single molecule method that can be used to reliably measure binding constants of proteins binding to DNA, provided that they distort DNA. In TPM, the motion of a bead tethered to a surface by DNA is tracked using light microscopy. A protein binding to the DNA will alter bead motion. This makes it possible to measure binding properties. We use the bacterial protein Integration Host Factor (IHF) as an example to show how specific binding to DNA can be measured. Moreover, we show a new intuitive quantitative approach to displaying data obtained via TPM.

Introduction

Determination of binding affinity is an important part of the characterization of DNA-binding proteins. Typically, the binding affinity is expressed in terms of the binding constant (K_D). The K_D is the ratio of the off-rate constant, k_{off} , and the on-rate constant, k_{on} . Operationally, the K_D is defined as the concentration at which half of the substrate is bound by ligand (here: DNA and protein, respectively). For protein-DNA binding, the K_D is conventionally determined using Electrophoretic Mobility Shift Assays (EMSA), Filter Binding Assays, and Fluorescence Anisotropy (256-258). Additionally, Isothermal Titration Calorimetry (ITC), Surface Plasmon Resonance (SPR) and Microscale Thermophoresis (MST) were used (259-261). These techniques measure protein-DNA affinity in bulk, and therefore cannot distinguish different populations of bound and unbound molecules. Some of these techniques separate the protein-DNA complexes from solution or use small sample volumes, thus introducing large surface areas that may influence effective compound concentrations. Single molecule techniques such as Optical Tweezers, Magnetic Tweezers, Acoustic Force Spectroscopy (AFS) and Atomic Force Spectrometry (AFM) can be used to estimate the K_D (262, 263), but require a high level of expertise and often expensive experimental setups. Single molecule Förster Resonance Energy Transfer (smFRET) allows for measuring of DNA that is not bound to a surface and/or a bead (264), but requires fluorescently labeled DNA and protein and/or a sufficiently large and predictable conformational change of the DNA.

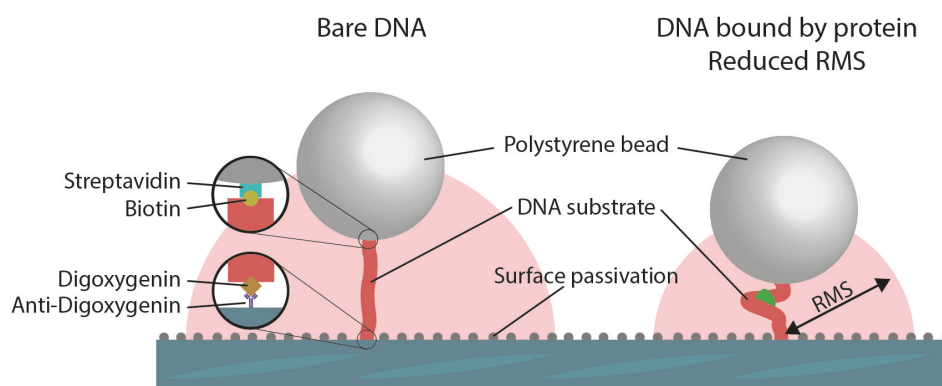


Figure 3.1: Principle of Tethered Particle Motion (TPM). A bead is tethered to a glass surface by DNA. The position of the bead is tracked over time. Proteins that bind to the DNA and thereby deform it, change its apparent persistence length, which results in a change in the Root Mean Square displacement (RMS) of the bead.

Tethered particle motion (TPM) is a simple, label-free single molecule technique that can be used to characterize DNA-binding proteins in a high-throughput manner. The technique relies on the tracking of a bead in solution that has been tethered to a glass surface by a DNA molecule or other macromolecule (**FIGURE 3.1**) (265). The bead exhibits Brownian motion, which is restricted by the tether. In contrast to some of the above-mentioned techniques, no additional external force is applied to the bead. The xy-positions of the bead are recorded over time and are used to calculate the root mean square displacement (RMS) of the bead (**EQUATION 3.1**). TPM has a high throughput (*see NOTE 1*), which makes it a powerful tool to study the characteristics of DNA-binding proteins. DNA-binding proteins that deform the DNA upon binding, change the apparent persistence length (L_p) of the DNA, a measure for rigidity, when bound, thereby altering the RMS (266). Although the L_p cannot be directly calculated with TPM, the change in RMS is a reliable measure for protein binding. TPM has been used to measure DNA binding by architectural proteins, proteins involved in DNA replication, transcription, DNA repair and recombination (123, 127, 267-274).

$$RMS = \sqrt{\frac{1}{n} \sum_{i=1}^n [(x_i - \bar{x})^2 + (y_i - \bar{y})^2]} \quad (3.1)$$

Equation 3.1. Calculation of the Root Mean Square displacement. x and y are the coordinates of the two-dimensional position of the bead. The summation runs over all measurements taken in the 45 second time window.

In this chapter we describe the use of TPM to measure the K_D of sequence-specific DNA-binding proteins. As an example we characterized the bacterial Integration Host Factor (IHF) (275). IHF is a nucleoid-associated protein (NAP) that is involved in shaping genome architecture in bacteria. IHF also plays a role in up- and downregulation of genes (276). This function is attributed to the ability of IHF to bend DNA and bring regulatory elements together, as well as due to direct interactions with RNA polymerase (277, 278). IHF is known to bend DNA when bound at its high affinity recognition sequence (279) and also bends DNA when bound to DNA non-specifically (280). Bound at its specific site, binding angles of 120–160° have been reported (281-283).

Materials & Methods

All materials are listed in the **SUPPLEMENTARY INFORMATION** section.

Parafilm semi-circles

1. Cut a strip of parafilm at a length fitting the grid of the laser cutter.
2. Remove the protection layer from the parafilm and place the parafilm onto the grid of the laser cutter.
3. Load the template of the semi-circles in the laser cutter software of choice. Design has to be such that two parafilm semicircles on a 28 mm coverglass create a 6 mm wide channel. Position the parafilm so that the template virtually overlays the parafilm.
4. Fix the parafilm on the grid with office tape.
5. Cut the parafilm (*see NOTE 9*).
6. Remove parafilm strip. The semi-circles will stick to the grid of the laser cutter. Take the semi-circles off the grid individually with tweezers.
7. Store the parafilm semi-circles at 4°C to prevent sticking to each other.

Sample chambers

The following steps should be carried out with gloves, both to protect the hands from exposure to chemicals and to keep the sample chambers free of contaminants.

1. Place the cover glasses in the slide holder using the tweezers with pointed tips (*see NOTE 10*).
2. Place the holder in a beaker and submerge in acetone. Place the beaker in a sonication bath and sonicate for one cleaning cycle (*see NOTE 11*).
3. Transfer the slide holder into a new beaker and submerge the slide holder in ethanol. Place the beaker in a sonication bath and sonicate for one cleaning cycle.
4. Remove the holder from the beaker. Place the cover glasses on a precision wipe and dry the cover glasses by gently wiping them with precision wipes. Turn the cover glasses over and leave exposed to air for 10 minutes to dry.
5. Using tweezers place two parafilm semi-circles in parallel on opposing sides of a 28 mm cover glass, so that a straight channel is created (**FIGURE 3.2**). The parafilm should align with the glass on the round edges. Push the parafilm firmly but carefully onto the cover glass with the tweezers so that the parafilm adheres to the cover glass (*see NOTE*

- 12).
6. Place the 35 mm cover glass on top of the parafilm-covered side of the 28 mm cover glass using tweezers, carefully centered onto the 28 mm cover glass. Push firmly but carefully with the tweezers so that the slides stick together.
7. Place the sample chamber on the hot plate (pre-heated to 100°C) using tweezers. Cover the sample chamber with the lid and press firmly and evenly on the lid for 10 seconds. Remove the lid from the sample chamber (see NOTE 13).
8. Lift the sample chamber off the hot plate using tweezers and let it cool down on a precision wipe. Store the sample chamber in a small plastic storage box at room temperature.

Bead suspension

1. Dilute the stock of streptavidin-coated beads to 0.01% w/v in 5 ml BB (see NOTE 16).
2. Vortex thoroughly for 5 minutes.
3. Sonicate the bead suspension on ice for 3 minutes (active), 2.5 s on and 9.9 s off, at 25% of the maximum power of the immersion sonicator. The total run time will be around 15 minutes. The tip of the immersion sonicator should be submerged by roughly 3 cm. Never sonicate the bead

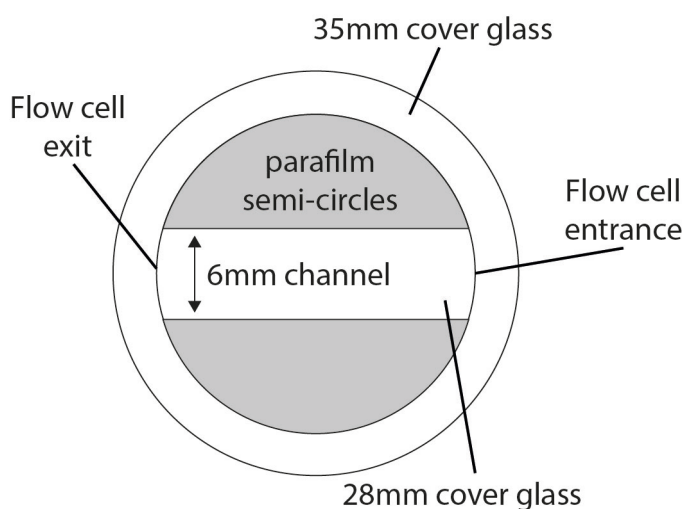


Figure 3.2: Sample chamber for TPM. The sample chamber consists of two parafilm semi-circles sandwiched between two cover glasses. The sample chamber is agglutinated by heating to 100°C.

suspension for more than 5 minutes of active time, as this may damage the beads and its coating.

4. Flow 30 μl of bead suspension into an empty sample chamber. If less than 1% of the beads is clustered, it is considered a good bead suspension.
5. Store the bead suspension at 4°C.

DNA substrate

1. Add the following components to a PCR reaction tube on ice:

Component	Quantity
DNA template in 10 mM Tris pH 7.5	10 ng
Biotinylated forward primer	25 pmol
Digoxigenin-labeled reverse primer	25 pmol
Phusion master mix	25 μl
DMSO	1.5 μl
Milli-Q	To total volume of 50 μl

2. Place the PCR reaction tube in the thermal cycler and use the following touchdown protocol (*see NOTE 14*):

Temperature	Duration	Cycles
98°C	30 s	
98°C	10 s	15x
72°C (-1°C per cycle)	20 s	
72°C	60 s	
98°C	10 s	25x
57°C	20 s	
72°C	60 s	
72°C	600 s	
12°C	∞	

3. Purify the PCR product using a PCR clean-up kit (*see NOTE 15*).
4. Run 2 μl of PCR product on a 1% agarose gel, pre-stained with GelRed, alongside a suitable DNA marker. Verify the length of the DNA substrate and estimate the concentration using a spectrophotometer. Store the DNA substrate at -20°C at a concentration of 100-500 pM.

Anti-digoxigenin solution

1. Dissolve 200 µg of anti-digoxigenin in 1 ml of BB.
2. Aliquot per 20 µl and store the anti-digoxigenin at -20°C.
3. When using the anti-digoxigenin solution, thaw an aliquot and dilute 10 times with BB.

DNA-tethered bead in sample chamber

1. Take a clean sample chamber and flow in 20 µg/ml anti-dig solution by holding the tip of a pipette against the opening on the right side of the sample chamber at a 45° angle and carefully pressing the plunger button (*see NOTE 17*). Incubate 10 minutes at room temperature.
2. Wash the sample chamber with 100 µl passivation solution and incubate for 10 minutes at room temperature. The excess liquid flowing out of the sample chamber is collected using Whatman paper. This promotes an undisturbed and uninterrupted flow in the channel.
3. Wash the sample chamber with 100 µl BB (*see NOTE 18*).
4. Flow in 100 µl 100 pM DNA substrate (*see NOTE 19*). Incubate for 10 minutes at room temperature.
5. Wash the sample chamber with 100 µl BB.
6. Pipette the bead suspension stock up and down and flow in 100 µl bead suspension. Incubate for 10 minutes at room temperature.
7. Wash the sample chamber with 100 µl EB.
8. Flow in 100 µl protein solution, diluted to the desired concentration in EB. Incubate for 10 minutes at room temperature.
9. Wash the sample chamber with 100 µl protein solution of the same concentration as in step 8.
10. Seal the sample chamber on both sides by applying nail polish to the opening of the sample chamber. When the nail polish is dry, the sample chamber is ready to be used.

Dilution series

1. Calculate a dilution series with a range from 0 to 1000 nM using twofold dilution steps (*see NOTE 20*). Determine the concentration range in which DNA binding is to be expected. Literature values can be used if available; if not, preliminary experiments may be required.
2. Dilute a protein stock aliquot to a concentration that can be used to make further protein dilutions with EB (*see NOTE 21*). Store this stock

on ice (*see NOTE 22*).

3. Using the working stock, make a dilution series by diluting part of the working stock with EB to the desired concentration. Dilute part of this sample with EB in order to create the next sample (*see NOTE 21*). Store samples on ice (*see NOTE 22*).
4. Measure at least two, but preferably three dilution series (*see NOTE 23*).

Imaging

1. An hour before imaging, turn on the lamp and the heating stage.
2. Place immersion oil on the objective and start imaging software.
3. Place a sample chamber in the sample chamber holder.
4. Raise the objective until the beads are in focus (*see NOTE 24*).
5. Close the isolation chamber. The temperature of the chamber and the objective will now rise to a set temperature (*see NOTE 25*).
6. Re-focus on the beads and ensure that no focal drift is observed (*see NOTE 26*).
7. Select a field of view by moving the stage of the microscope. Select the beads and track them for at least 1500 frames. Images are taken at 25 Hz, with an exposure time of 20 ms. At least 100 'good' bead-tether combinations are needed for quantification after quality check (explained in 'Data analysis'; *see NOTE 27*).
8. Use the tracking software to determine x- and y-positions of the beads (*see NOTE 8*).

Data analysis

1. Calculate the anisotropic ratio (**EQUATION 3.2**), the root mean square displacement (RMS) and the standard deviation of the RMS for all measurements at a given concentration combined (*see NOTE 28*). Discard tethers with an anisotropic ratio of above 1.3. Discard measurements with a standard deviation of the RMS larger than 6% of the RMS. We thereby exclude beads sticking to the surface during the measurement (*see NOTE 29*).
2. Plot all RMS values that meet the requirements as explained in step 1 in a histogram.
3. Fit a Gaussian curve to the histogram to determine the average RMS and standard deviation for each condition.
4. Calculate the standard error of the mean for each condition.

$$\alpha = \frac{l_{major}}{l_{minor}} \quad (3.2)$$

Equation 3.2: Calculation of the anisotropic ratio (α). l_{major} and l_{minor} represent the length of the major and minor axis of the xy-scatter plot, respectively.

Analysis of specific protein-DNA binding

1. Determine the RMS of the DNA-tethered bead in the absence of protein. This is the position of the peak in the Gaussian fit at 0 nM (**FIGURE 3.3A**).
2. Determine the RMS at saturation levels of specific binding. This is the concentration at which the peak seen at 0 nM has completely disappeared and a peak at a different RMS has appeared (**FIGURE 3.3F**).
3. Concentrations at which two peaks are observed, represent a situation in which a subpopulation of the DNA is bound by protein (**FIGURE 3.3B-E**).
4. For the concentrations at which subpopulations of bound and unbound states can be observed, calculate the occupancy (**EQUATION 3.3**). Occupancy is the ratio between bound and unbound DNA substrate.
5. Find the K_D and Hill coefficient (n) by solving the Hill binding model (**EQUATION 3.4**) for K_D and n (see **NOTE 30**).
6. Fit the occupancies using the Hill binding model (**FIGURE 3.4**; see **NOTE 31**). Use graphing and data analysis software to display the occupancies per concentration and the Hill fit in a graph.

$$\theta = \frac{h_{peak, bound}}{h_{peak, bound} + h_{peak, bound}} \quad (3.3)$$

Equation 3.3: Calculation of occupancy. The occupancy is the fraction of substrate bound by protein and can be calculated from the area of the peaks of the fitted Gaussian distributions.

$$\theta = \frac{1}{\left(\frac{K_D}{[L]}\right)^n + 1} \quad (3.4)$$

Equation 3.4: Hill binding model. Occupancy θ , binding constant K_D , ligand concentration $[L]$, Hill coefficient n .

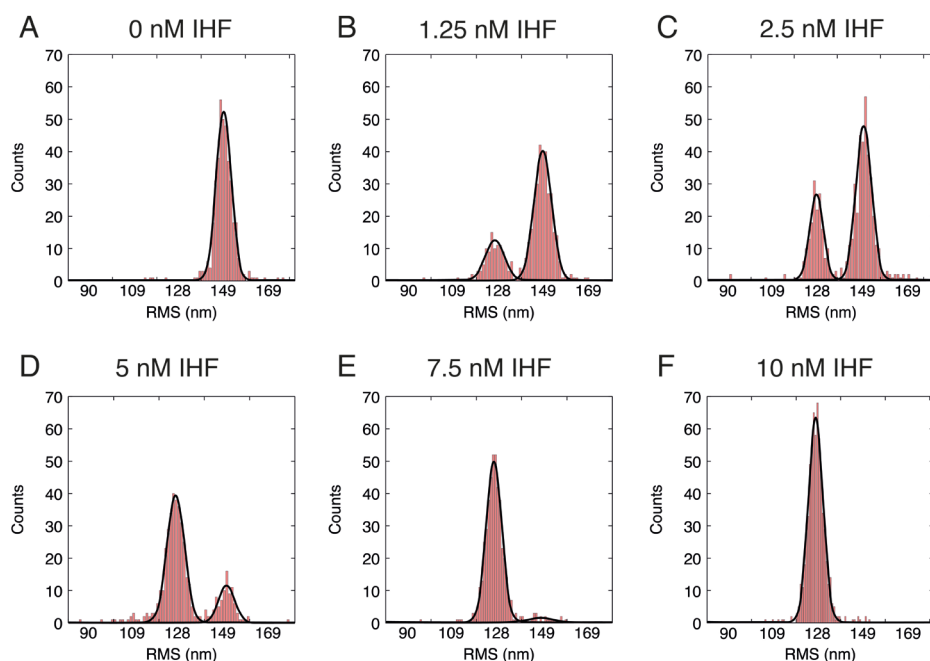


Figure 3.3: Specific binding of IHF to DNA at 0-10 nM. A-F: histograms of RMS values obtained for DNA-tethered beads at IHF concentrations of 0, 1.25, 2.5, 5, 7.5 and 10 nM. The histograms have been fitted with a Gaussian function in KaleidaGraph (black, semi-opaque line).

Example of K_D estimation for DNA-binding protein IHF

We used IHF as an example to illustrate how to characterize specific DNA-binding proteins. In our example, we observe a reduction in RMS at low (1.25 nM) IHF concentrations from 149 nm to 127 nm (FIGURE 3.3). This reduction in RMS is caused by DNA bending as a consequence of IHF binding at its specific binding site (*see NOTE 32*). Saturation of specific binding is achieved at 10 nM, the concentration at which only the population with reduced RMS is observed. Fitting the data using the Hill binding model (χ^2 : 4.6) yields a binding affinity (K_D) of 3.0 ± 0.3 nM, which is in agreement with values reported using EMSA on different variants of the binding consensus sequence (284). A Hill coefficient (n) of 2.5 ± 0.5 was found. A Hill coefficient >1 indicates positively cooperative binding; $n < 1$ indicates negatively cooperative binding (*see NOTE 33*).

Data representation

TPM is a single molecule method with a high throughput. As a consequence data are often reported in terms of an average RMS value. However, displaying the full population of RMS values can be insightful to illustrate distribution shape and

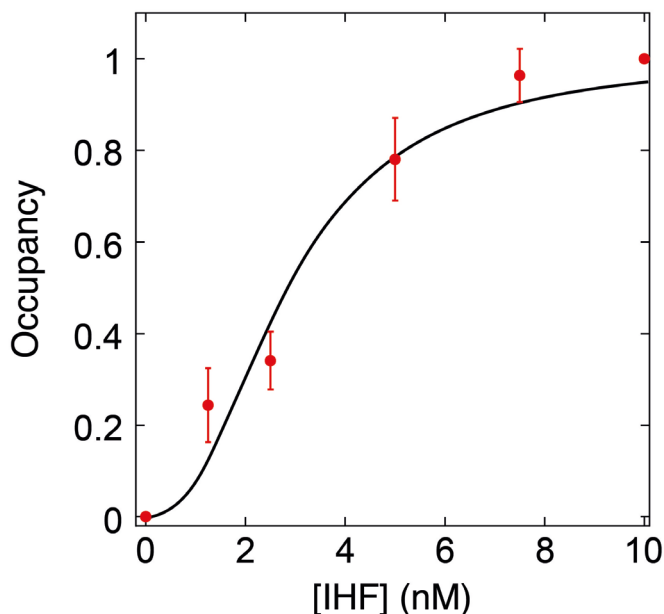


Figure 3.4: Binding curve for specific binding of IHF to DNA. The data points were fit using the Hill binding model. The Hill coefficient (n) is 2.5 ± 0.5 , and the binding affinity (K_D) is 3.0 ± 0.3 nM. Error bars represent the standard error of the mean.

population heterogeneity. The new representation approach presented here accommodates both average values, as well as unprocessed population information.

1. Plot all data points as black squares using the graphing and analysis software.
2. In the same plot, include the values of the average RMS at all concentrations and display in a color different from that of the data points.
3. Export the plot as a vector image.
4. Open the plot in Adobe Illustrator and select all black squares. Change the opacity from 100% to 5% (FIGURE 3.5; see NOTE 34).

In our example, specific and non-specific binding by IHF can be observed. Specific binding of IHF is observed between 0 and 10 nM, where the RMS decreases from 149 ± 5 nm to 127 ± 5 nm and is characterized by the two populations. Intensity profiles of the populations are equivalent to the histograms of FIGURE 3.3. In the non-specific binding regime, at IHF concentrations >30 nM, we find a single population of which the RMS continues to reduce as a function of IHF concentration up to 400 nM. This further reduction in RMS is reduced to sequence unspecific binding of the protein. At concentrations >400 nM the RMS increases,

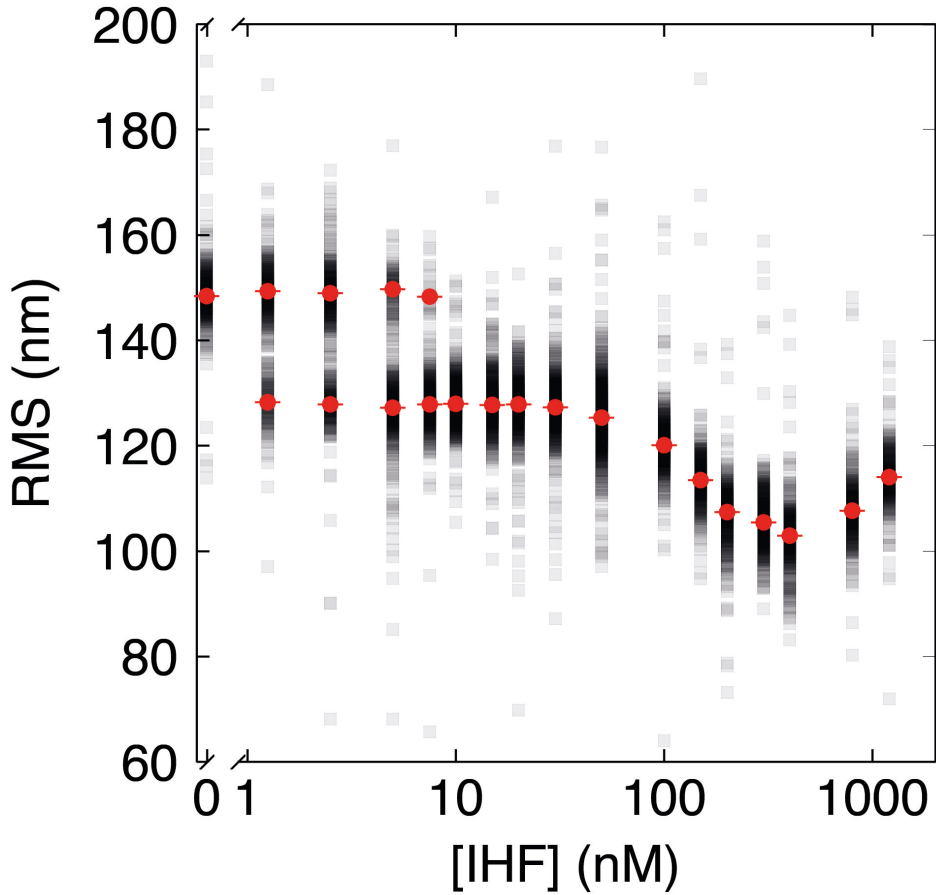


Figure 3.5: Compaction of DNA by IHF. Black, partially transparent squares represent the individual RMS values of DNA-bead complexes after application of anisotropic ratio and standard deviation of the RMS as selection criteria. Red dots and bars represent the average RMS that resulted from fitting with a Gaussian function.

which is attributed to stiffening of DNA by IHF bound at high density along the DNA, similar to the effects of observed for the HU protein (285).

Notes

- 1) Up to 100 beads were measured for 1 minute.
- 2) A custom made hot plate was used to heat the sample chambers. The hot plate is made from a circular massive metal block in which a shallow cavity is carved (diameter 35 mm). A sample chamber fits into the cavity. A metal lid (diameter 35 mm) with a plastic handle covers the sample chamber. The hot plate is placed on a heating block and heats the sample chamber to 100°C.
- 3) Nitrile gloves were used, as the powder that is often found in rubber gloves may contaminate the sample chamber.
- 4) As a template the plasmid pGp1 was used (286), which contains part of the bacteriophage mu genome. A 685bp fragment that contains 1 IHF binding site exactly at the center was amplified, which is expected to result in the highest effect on end-to-end distance (283). Our experience is that a DNA substrate in the order of 700 bp is ideal, as protein binding results in a clearly detectable change in RMS.
- 5) The forward and reverse primers that we used are 5' BIO-TGATT AAGGTTGTGGTTAATTTGTTTATCAGTTCCG 3' and 5' DIG-ATTACTTCCGGTTTAGTTTCTAAGGCG 3'.
- 6) The experimental buffer contained 10 mM Tris-HCl pH 8.0 and 100 mM KCl, which is suitable for specific binding of IHF (287). The osmolarity of the buffer should be sufficiently high to prevent non-specific binding of specific binding proteins. A different protein of interest may require buffer optimization.
- 7) Recombinant IHF at a concentration of 120 μ M was used here. This stock was diluted 100-fold at minimum. The number of freeze-thaw cycles was minimized by aliquoting the protein stock, to guarantee protein activity.
- 8) Software written for Acoustic Force Spectrometry was used (288), which is freely available online*. Other particle tracing software has been described and compared by Chenouard *et al.* (289). Our lab also

* http://figshare.com/articles/AFS_software/1195874

has positive experience with PolyParticleTracker (290).

- 9) Here, the computer-aided design software AutoCAD was used to design the cutting template in combination with the VersaLaser tabletop Laser cutter. We cut the parafilm at 55% laser power, 30% of the maximum speed, 1000 pulse and 1.00 mm Z-axis. Different laser cutters may require different settings, allowing the parafilm to be cut straight and without melting or burning.
- 10) Ensure that each slot is occupied by only one cover glass. The cover glasses tend to stick together, which prevents them from being cleaned properly.
- 11) For sonication of the glass slides, the Emag EMMI 4 sonicator was used, which is pre-programmed in cycles of 9 minutes at 42.5 kHz.
- 12) Note that the parafilm leaves grease marks on the cover glass, so avoid sliding after placing them.
- 13) Heating the sample chamber for too long may result in excessive melting of the parafilm, which causes a non-straight sample chamber. This prevents a good flow in the sample chamber. As a result, coating of the surface with anti-digoxygenin and/or passivation agent may be heterogeneous, which reduces throughput and results in lower effective protein concentrations. Especially experiments at low concentration may then suffer from reduced reproducibility.
- 14) Conventional protocols with fixed annealing temperatures may be used as well, but need to be individually optimized for different primer-template combinations and synthesized fragment lengths.
- 15) Purification using the GenElute PCR clean-up kit separates the PCR product from other components that were added to the reaction tube to facilitate the PCR reaction, such as excess primers, nucleotides and salts. The result of this purification is the labeled DNA-substrate in 10 mM Tris-HCl pH 8.5 (provided with the kit).
- 16) The bead stock is vortexed and sonicated to homogenize the suspension. Beads tend to stick together, which will distort measurements.

- 17) Check if the sample chamber has no cracks in either of the cover glasses, as defects may create a flow inside the sample chamber while measuring. This will affect the RMS by giving it a directional bias. Also, check if the channel is straight; see **FIGURE 3.3**. Left-handed people may prefer using the opening on the left side of the sample chamber for sample loading.
- 18) The volumes used for washing depend on the volume of the sample chamber. As a rule of thumb, we wash with approximately three times the volume of the sample chamber.
- 19) The DNA substrate concentration is 100 pM, but when we find that this concentration is not sufficient for a particular experiment, we raise the concentration up to 500 pM.
- 20) A dilution series with concentrations from 0 to 1000 nM is used, with twofold dilution steps. Based on results and literature values, an additional range and/or different dilution steps may be used, which may result in a more reliable calculation of the K_D .
- 21) When measuring many samples, consider breaking up the dilution series in two or more overlapping ranges. Here, we diluted a 120 μ M IHF stock to a working stock of 1.2 μ M. From that, we made a serial dilution which included IHF concentrations of 800, 400, 300, 200, 150, 100 and 50 nM: the high range concentrations. For the low range concentrations, we diluted the 1.2 μ M working stock directly to 50 nM, from which we subsequently made a serial dilution which included IHF concentrations of 30, 20, 15, 10, 7.5, 5, 2.5 and 1.25 nM. We verified that the RMS values at 50 nM from the high range dilution series were identical to the RMS values at 50 nM from the low range dilution series. We also measured the RMS in the absence of IHF, indicated with 0 nM.
- 22) Do not freeze the protein solutions, since the activity of the protein may be affected.
- 23) For reliable results and to minimize the effect of pipetting errors, experiments should be carried out at least in duplicate, but preferably in triplicate, and independently from each other. Especially when the concentration range over which binding takes place is narrow, small changes in sample concentrations can be of significant influence on the

outcome of the binding curve.

- 24) Due to a sedimentation effect, there may be a slight difference in RMS between beads tethered to the bottom surface and beads tethered to the top surface. Therefore, always choose the same surface for measurements. Also, the focus for top and bottom surface may differ.
- 25) It may take 5 to 20 minutes, depending on the temperature, before the isolation chamber has completely reached the desired temperature.
- 26) While the temperature of the immersion oil on the objective rises, the refractive index of the oil changes slightly. As a result, focal drift is observed. Wait until the immersion oil has reached the desired temperature and then refocus. When no focal drift is observed, the immersion oil has the desired temperature.
- 27) At least 100 good bead-tether combinations are needed, but the actual amount of measured bead-tether combinations to reach this limit may differ. For IHF, 300 measured bead-tether combinations always resulted in more than 100 good combinations. The amount of bead-tether combinations that need to be measured, depends on the protein that is used.
- 28) Verify that individual experiments done at the same concentration give similar outcomes. If so, RMS values can be combined and can be fitted together. If one measurement is a clear outlier due to a known cause, discard the measurement. In other cases, re-measure the sample and check if the results align with any previous measurements.
- 29) Discard any bead with an anisotropic ratio >1.3 , which is considered non-symmetrical. Non-symmetrical beads indicate for example that a bead is attached to two DNA tethers. An extensive study that links motion patterns to setup defects has been done by Visser *et al.* (291) Also discard any bead with a standard deviation of the RMS of $>6\%$, since this indicates rapid protein dissociation and sticking and/or releasing of the bead to or from the sample chamber surface.
- 30) The Hill equation can be solved using IGOR Pro, which was done here, but MatLab and other data analysis programs are also suitable for this.
- 31) As error bars, the standard error of the mean of the individual measure-

ment series per concentration has been used.

- 32) If desired, the effective persistence length of the DNA can be determined for a single protein bound to DNA, which could be used to calculate the DNA bending angle as described by Kulić and coworkers (266). However, this requires an extensive simulation series, which reveals the relation between RMS and L_p . The simulations are highly dependent on the length and sequence of the DNA substrate, buffer conditions and temperature of the experiment. This means that for every experiment in which one or more of these parameters is changed, a new simulation series is required. The simulations have been described by Driessen *et al.* (292).
- 33) Here, the Hill coefficient is 2.5, which indicates positively cooperative binding. However, in this example, only one IHF dimer binds to DNA. The apparent positively cooperative binding is possibly a result of the dimeric nature of the protein. Both components of the dimer have to bind to the DNA in order to bend the DNA.
- 34) For representation of all data points, squares are most suitable, because squares are easy distinguishable when data points overlap. Circles may make it harder to distinguish individual data points in the plot. The opacity that suits your data best depends on the number of data points. When using more data points than shown in the example, it may be useful to use a lower opacity. The Gaussian distribution of the data points should be visible from the intensity of the combination of data points.

Supplementary information

Parafilm semi-circles

1. Parafilm
2. Office tape
3. Laser cutter
4. Computer-aided design (CAD) software (for example AutoCAD or CorelDRAW)
5. Tweezers

Sample chambers

1. Round cover glass, diameter 35 mm (VWR)
2. Round cover glass, diameter 28 mm (Thermo Scientific)
3. Parafilm semi-circles (*see above*)
4. Cover slip rack (Diversified Biotech)
5. Ethanol
6. Acetone
7. Beaker
8. Sonication bath
9. Heating block, capable of heating up to 100°C, with hot plate for 35 mm cover glasses (*see NOTE 2*)
10. Tweezers with pointed tips
11. KIMTECH SCIENCE* precision wipes tissue wipers (Kimberly-Clark)
12. Nitrile gloves (*see NOTE 3*)
13. Whatman paper

Bead suspension

1. Streptavidin-coated polystyrene beads, diameter 0.44 μm (Kisker Biotech)
2. Immersion sonicator with fine tip
3. Bead buffer (BB) (10 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM EDTA pH 8.0, 1 mM DTT, 3% glycerol, 100 $\mu\text{g/mL}$ BSA-acetylated [Ambion])
4. Vortex

Anti-digoxigenin solution

1. Anti-digoxigenin (anti-Dig) polyclonal antibodies 200 µg (Roche)
2. Bead buffer (BB) (*see above*)

DNA substrate

1. Template for DNA substrate containing the binding site of interest. PCR amplification with compatible primers should yield a substrate of desired length and sequence (*see NOTE 4*).
2. Biotinylated forward primer, 10 µM in 10 mM Tris, pH 7.5, designed to be compatible with the template (*see NOTE 5*)
3. Digoxigenin-labeled reverse primer, 10 µM in 10 mM Tris, pH 7.5, compatible with the template (*see NOTE 5*)
4. Phusion master mix (New England Biolabs)
5. DMSO
6. Milli-Q
7. GenElute PCR clean-up kit (Sigma Aldrich)
8. PCR reaction tubes
9. Biorad C1000 Thermal Cycler

Substrate quality control

1. Agarose gel, 1% in Tris/Borate/EDTA (TBE) buffer
2. GelRed nucleic acid gel stain (Biotium)
3. GeneRuler DNA marker
4. Spectrophotometer to measure DNA concentrations (Nanodrop, Qubit, SimpliNano or comparable device)

Sample chamber reagents

1. Bead buffer (BB) (*see above*)
2. Anti-digoxigenin solution
3. Passivation solution (4 mg/ml Blotting Grade Blocker [Bio-Rad] dissolved in SB)
4. Experimental buffer (EB) suitable for protein binding (*see NOTE 6*)
5. Protein of interest, preferably at high concentrations to avoid as much as possible components of the protein storage buffer on the EB (*see NOTE 7*)

Imaging

1. Isolation table
2. Inverted microscope (Nikon Diaphot 300) (**FIGURE S3.1**)
3. 100x oil-immersion objective (NA 1.25)
4. Immersion oil for microscopy (Merck)
5. Heat chamber (Bioscience tools TC-HLS-025 heating element for heating of the objective and a custom sample chamber heating element consisting of 40 parallel resistors); Pt100 sensors attached to the sample chamber holder and the objective, and two SA200 PID digital temperature controllers)
6. Climatized room
7. Isolation box
8. Thorlabs CMOS camera DCC1545M
9. Lens paper
10. Nacre-free nail polish
11. Particle-tracking software (software available online (288); *see NOTE 8*)

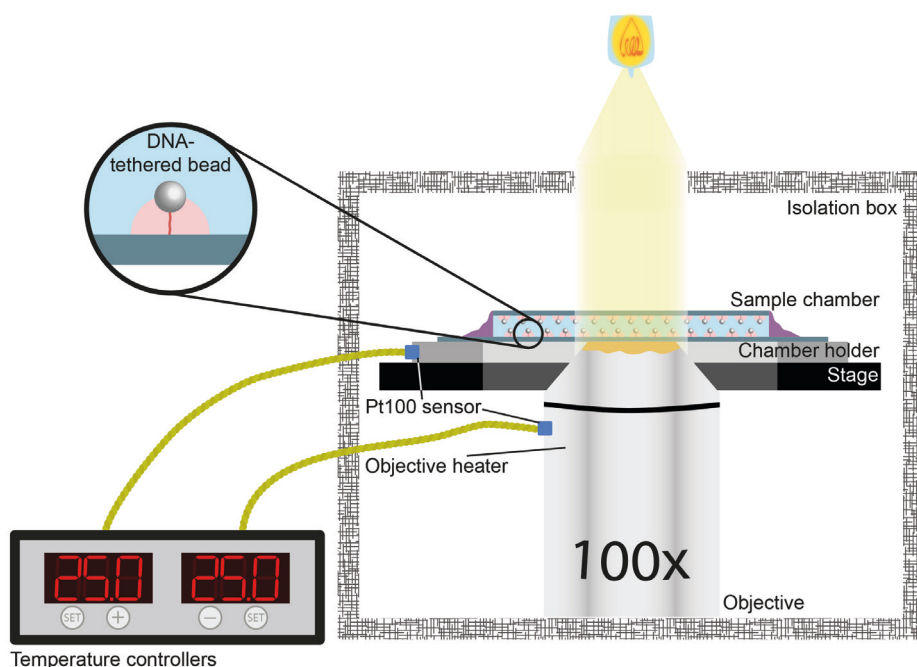


Figure S3.1: Schematic of the Tethered Particle Motion setup. The sample chamber is heated by two heating elements, one placed in the sample chamber holder and one on the objective. An isolation box covers the stage, sample chamber and heating elements. Schematic is not to scale.

Data analysis and representation

1. Computer
2. Custom MatLab routine for calculating RMS, standard deviation of the RMS and anisotropic ratio. The routine also selects the beads that meet the requirements for reliable beads (*see* 'Data analysis' in the **METHODS** section)
3. Graphing and data analysis software, such as KaleidaGraph, Origin or IGOR Pro
4. Adobe Illustrator

