A human transcription factor in search mode

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SUPPLEMENTARY INFORMATION
Section 1. DNA helicoidal analysis of the MTERF1 recognition complex

Figure S1. Structural analysis of DNA in the MTERF1-DNA specific complex (1). Using Curves (2), the base pair step parameters rise distance (top left), bending angle (top right), opening angle (bottom left), and twist angle (bottom right) were calculated. For reference, the parameters for B-DNA are shown in each panel.

Section 2. Helix fitting

Defining an irregular helix that makes only one revolution (or less) such as MTERF1 is a challenging problem. Many curve fitting methods exist, but few can deal with the one-turn helix-fitting problem (3). Methods that can solve such a helix suffer from requiring that atomic positions are regularly spaced and helically symmetric (4-6). Our approach is related to the Riemann helix-fitting method of Frühwirth et al. (7), though simpler since we numerically search helical orientations, enabling simpler transformations. A helix-fitting procedure that geometrically simplifies the problem was developed. The normal vector to a plane coincides with the helical axis of a cylindrical helix if the coordinates of the helix project a perfect circle onto the plane.
Our approach to calculate the helical parameters – pitch and radius – of the MTERF1 superhelix was to find the plane onto which the superhelical residues projected the best circle, the normal plane to the assumed helical axis. The plane is defined in terms of the direction cosines \((a,b,c) = (\sin\phi\cos\theta, \sin\phi\sin\theta, \cos\phi)\) of the helical axis (with \(\phi\) and \(\theta\) the spherical coordinates). A 2D-grid search - \(\phi\) and \(\theta\) - of helical axis orientations was performed by rotating the Ca of the \(N\) superhelical residues \((X,Y,Z)\) to \((x,y,z)\), in spherical coordinates,

\[
\begin{pmatrix}
\sqrt{1-a^2} & 0 & a \\
-\frac{ab}{r} & c/r & b \\
-\frac{ac}{r} & -\frac{b}{r} & c
\end{pmatrix}
\begin{pmatrix}
X \\
Y \\
Z
\end{pmatrix}
\]

Eqn. S1

The projection depends only on the normal plane’s orientation, and not on its position, so it is assumed to pass through the (arbitrary) origin. On the projection plane, the centre and radius of the circle are determined from the linear least squares problem,

\[
\begin{pmatrix}
1 & \sum_{i=1}^{N} x_i & \sum_{i=1}^{N} y_i \\
1 & \sum_{i=1}^{N} 2x_i & \sum_{i=1}^{N} 2y_i \\
1 & \sum_{i=1}^{N} 2x_i & \sum_{i=1}^{N} 2y_i
\end{pmatrix}
\begin{pmatrix}
\rho \\
x_0 \\
y_0
\end{pmatrix}
= \begin{pmatrix}
\sum_{i=1}^{N} x_i^2 + \sum_{i=1}^{N} y_i^2 \\
\sum_{i=1}^{N} x_i^2 + \sum_{i=1}^{N} y_i^2 \\
\sum_{i=1}^{N} x_i^2 + \sum_{i=1}^{N} y_i^2
\end{pmatrix}
\]

Eqn. S2

where \(\rho\) is the radius of the helix and \((x_0, y_0)\) is its centre. The \(\phi\) and \(\theta\) of the orientation with the lowest residual,

\[
Res^2 = \sum_{i=1}^{N} (x_i - x_0)^2 + (y_i - y_0)^2 + \rho^2 - 2\rho\sqrt{(x_i - x_0)^2 + (y_i - y_0)^2}
\]

Eqn. S3

is the helical frame. With the coordinates already rotated in Eqn. S1, pitch is simply \(|z_N - z_1|\Phi/2\pi\), where \(\Phi\) is the parametric angle of the circle and thus the helix – the helical sweep.

Finding the helical parameters of apo MTERF1

An unconstrained fit of the superhelical residues was performed for the full ensemble of apo MTERF1 structures, after first RMS aligning the structures to the superhelical residues (using the recognition mode structure as the reference) to remove rotational and translational motions. Then the dependence of the helical parameters was analysed, including fit residual, on the helical axis orientations as apo MTERF1 samples a breadth of conformations (Figure S2). A scatter plot of helical axis orientations shows the full range of \(\theta\) is being sampled while a 90° range in \(\phi\) is being sampled, but clusters of distinct orientations are apparent (Figure S2A). To determine why different structures of apo MTERF1 lie along rather different helical axes, heat maps were plotted of the residual (Figure S2B), radius (Figure S2C), and pitch (Figure S2D). Configurations with low residual - \(\phi\) in [50°,70°] and \(\theta\) in [240°,300°] – lie in the same region of the grid where structures exhibit low radius. That same region of the grid also presents a range of superhelical pitch in which 30 Å is the minimum.

apo MTERF1 structures with representative values of superhelical pitch were selected from the MD ensemble. Figure S2E shows a surface representation of an apo MTERF1 structure with low superhelical pitch – 35 Å. Two other conformations of MTERF1 with high pitch – 52 Å – and very high pitch – 64 Å – are presented in Figure S2F and S2G, respectively. The very high pitch structure (Figure S2G) is representative of conformations
whose orientations can lead to artefactual helical parameters. The helical axis may not be the long axis of the protein, instead it is possible to be in a half-rotated orientation (the helix would be wider and more extended, with large pitch and radius but << 360° sweep) or full rotated orientations (the helix would be disk-like, with << 30 Å pitch, >> 20 Å radius). Our confined grid search of the same structures prevents these spurious helices by restricting helical orientations to [50°,70°] and θ in [240°,300°].

Figure S2. Calculating superhelical parameters of apo MTERF1. (A) Scatter plot of superhelical axes. Heat map of (B) residual, (C) radius, and (D) pitch. (E) Low pitch apo MTERF1. (F) High pitch apo MTERF1, the conformation corresponding to the structure found in the specific complex. (G) Very high pitch apo MTERF1. In (E) and (F) the unconstrained helical axis orientations were in ϕ in [50°,70°] and θ in [240°,300°] whereas the
superhelical residues of the structure in (G) adopted an orientation in an alternate region of the map, the pitch and radius of which are shown in red. Data represents one of the 8apo MTERF1 simulations.

Section 3. Anisotropic Network Model details

A Hessian matrix was constructed and diagonalized to calculate the normal modes. A distance cutoff that provided high correlation between calculated and experimental B-factors (8) was chosen. Cutoffs of 8, 10, 12, 15, 18, 21, and 24 Å were tested (8) resulting in correlation coefficients of 0.5778, 0.6209, 0.6217, 0.6229, 0.6311, 0.6472, and 0.6619, respectively.

Section 4. Finding DNA parameters with our general helix frame

An ideal DNA geometry with a base pair step rise of 3.38 Å and a step twist of 36.0 degrees was analysed (Table S1). As expected, our method reproduces the helical rise parameters of B-DNA built using NAB (14) (36° twist, 10 base pair per revolution multiplied by 3.38 Å = 33.8 Å). Our method also reproduces twist. The radius of the major groove was defined as the radius of the helix traced by centre of the C1’ atoms.

<table>
<thead>
<tr>
<th>Atoms</th>
<th>Count</th>
<th>Residual (Å)</th>
<th>Radius (Å)</th>
<th>Pitch (Å)</th>
<th>Rise (Å)</th>
<th>Twist (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1’ (W)</td>
<td>22</td>
<td>0.023</td>
<td>5.85</td>
<td>33.80</td>
<td>3.38</td>
<td>36.00</td>
</tr>
<tr>
<td>C1’ (C)</td>
<td>22</td>
<td>0.015</td>
<td>5.85</td>
<td>33.81</td>
<td>3.38</td>
<td>36.00</td>
</tr>
</tbody>
</table>

(W) indicates the Watson strand, (C) the Crick strand in the ABC definition (15). The residual in our fitting procedure measures the deviation of the projected atomic coordinates from a perfect circle in units of Å (Section 2). Our sweep parameter is analogous to twist.

What is the pitch of B-DNA?

An upper limit of 42 Å B-DNA pitch was identified to be that which MTERF1 could bind in search mode. This value was arrived at by using two approaches. First, the literature in which the base pair step parameters rise was reported was reviewed (rise was multiplied by 10 since 10 bp/360°). Second, the helical pitch of B-DNA of our own MD simulations was measured.

A maximum value of rise in the central dinucleotide of CGCA/TGCG is 4.5 Å in nucleosome core particle crystal structures (16), which is almost identical to the average rise for all DNA sequences plus 3 standard deviations (4.4 Å) found from MD simulations of the 136 tetranucleotide sequences (17). Pitch for B-DNA, which contains 10 nucleotides per helical turn, is then 44 Å (horizontal lines in Figure 5). In a third example from the literature, Olson et al. reported average rise values of 3.36 Å and standard deviations of 0.25 Å of protein-DNA complexes in which complexes containing broken base pairs were omitted (41.1 Å pitch for B-DNA) (18). Together, the literature supports the assumptions that a B-DNA molecule a protein would randomly encounter in solution would likely posses a ~42 Å pitch, or less.
4 independent MD simulations of B-DNA were then performed. The systems were built, equilibrated, and production performed exactly as the search mode complexes. The helical parameters of the B-DNA in the MD trajectories were then analysed (Figure S3A and S3B). Overall, the distributions of DNA pitch agree with the above conclusions. The average pitch was observed to be 35.8 Å for the HS and 33.3 Å for the LS, with 2.31 Å standard deviation in the HS and 3.35 Å standard deviation in the LS. Thus the HS accesses pitch with 42.7 Å pitch (average plus 3σ) and the LS, 43.4 Å.

Section 5. Generating a nonspecific complex

Thus 15 apo MTERF1 structures were docked to B-DNA using DOT2.0 (9), which has been previously been shown to be suitable for protein-DNA docking (10). Our procedure followed previously published protocols (9). Briefly, REDUCE (11) parameters for heavy and polar hydrogen protein and DNA atoms were used, electrostatic potentials were calculated with APBS (12) and 0.200 M ionic strength, and electrostatic clamping was used to flatten pathological energies (9). van der Waals energies were estimated by counting DNA atoms that were within an interaction region coating the protein; the inner surface of the region was defined by the MSMS (13) molecular surface with a 1.4 Å probe radius, and the outer surface was defined by expanding the protein van der Waals atomic radii by 3.0 Å (9). Desolvation energies were not included in the calculations. 54,000 orientations of apo MTERF1 and B-DNA were evaluated for each of the low pitch protein structures. The protocol was validated by docking MTERF1 and DNA from crystallography, reproducing the experimental complex (RMSD < 3 Å for the 7 highest ranked structures).

Section 6. Calculating how well MTERF1 tracks a major groove

To test whether the highest ranked (lowest-energy) docked poses were reasonable models of a nonspecific complex, the 30 best poses from each of the 15 docking calculations were filtered by how well MTERF1 tracked the major groove, with the correct polarity. First, major groove sites were defined (Figure S4A). For B-DNA, a line connecting P atoms on opposite strands separated by 5 nucleotides (the P of nucleotide 1 on one strand and the P of nucleotide 6 on the other strand) is nearly parallel to the helical axis with a length of 21 Å. The midpoint of the line, ~10 Å from the P atoms and ~8 Å from nucleobase functional groups, was defined as a major groove site.
(Figure S4B). The distance between superhelical Cα atoms and major groove sites was measured, and each were expected to be a distance of 6-10 Å from each major groove site, based on inspection of the N- and C-site in the crystal structure and analysis of simulations with the specific MTERF1-DNA complex (~7 Å, see Figure S5). Poses were classified as nonspecific complexes when MTERF1 tracked the major groove and consecutive superhelical Cα atoms resided in consecutive major groove sites. An average distance between these protein residue and major groove site pairs was < 11 Å when the docked pose visually appeared to be tracking the major groove (Figure S6). The pairing sequence of Cα superhelical atoms and major groove sites was set by the pair with the smallest separation for a particular pose. For example, if the smallest separation between a superhelical Cα and a major groove site was the fourth Cα and the fifth major groove site (a pair), then the first Cα was automatically paired with the second major groove site, the second Cα paired with the third major groove site, and so on. The pairing rule causes very large Cα-major groove site distances (>25 Å average) when MTERF1 is docked in the reverse polarity, or criss-crosses the major and minor grooves (the screws are cross-threaded).
**Figure S4.** Method to geometrically measure how well MTERF1 tracks a major groove. **(A)** Major groove sites are the midpoints (pink spheres) between successive P atoms (dark red spheres) on opposite strands offset in sequence by 5, shown as pink dotted lines. **(B)** Major groove site-superhelical residue pairing scheme.
Figure S5. The expected values of our new major groove-binding distance was established by analysing the probability density of the average major groove distance between each superhelical residue and its nearest major groove site for control simulations of the specific MTERF1-DNA complex. 40 bins were used (the integral of the bins is one).
**Figure S6.** Quantifying the position of MTERF1 in the major groove of B-DNA. The upper pose has a groove tracking distance of 9.55 Å and the lower pose is 14.03 Å. In the upper pose, the first superhelical residue is paired to the tenth major groove site, the second superhelical residue the ninth site and so on, the individual
distances of which are 13.5, 11.1, 6.0, 8.3, 5.4, 9.2, 12.1, 14.5, and 5.9 Å. For the lower pose, the first superhelical residue is paired with the fourteenth major groove site, the second superhelical residue with the thirteenth major groove site and so on, the individual distances of which are 11.2, 11.6, 10.6, 13.5, 13.6, 17.0, 16.8, 15.9, and 16.3 Å.

Section 7. Equilibration details

Table S2. Equilibration procedure for explicit solvent MD simulations.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Ref</th>
<th>EOM</th>
<th>Steps ($x 10^3$)</th>
<th>Temp (K)</th>
<th>Ensemble</th>
<th>Group</th>
<th>Force constant (kcal/molÅ²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>xtal</td>
<td>min</td>
<td>10</td>
<td>-</td>
<td>-</td>
<td>A</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>MD</td>
<td>100</td>
<td>100/300</td>
<td>NVT</td>
<td>A</td>
<td>100</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>MD</td>
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<td>300</td>
<td>NPT</td>
<td>A</td>
<td>10</td>
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<td>MD</td>
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<td>300</td>
<td>NPT</td>
<td>A</td>
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<tr>
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<td>min</td>
<td>10</td>
<td>-</td>
<td>-</td>
<td>B</td>
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</tr>
<tr>
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<td>100</td>
<td>300</td>
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<tr>
<td>7</td>
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<tr>
<td>8</td>
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<td>300</td>
<td>NPT</td>
<td>B</td>
<td>0.1</td>
</tr>
<tr>
<td>9</td>
<td></td>
<td>MD</td>
<td>2500</td>
<td>300</td>
<td>NPT</td>
<td>-</td>
<td>0</td>
</tr>
</tbody>
</table>

Ref: reference coordinates. EOM: equation of motion: min, minimization; MD, molecular dynamics. Ensemble: NPT and NVT used a weak temperature coupling thermostat with isotropic position scaling. Group: atoms restrained to reference structure (Ref): A for apo MTERF1, all protein heavy atoms except the sidechains of residues 1 and 2, B for apo MTERF1, all protein backbone atoms – Ca, N, and C; A for holo MTERF1, all protein and DNA heavy atoms except, in apo MTERF1, the side chains of residues 1 and 2, B for holo MTERF1, all protein and DNA backbone atoms – C1’, C2’, C3’, C4’, O4’, C5’, O3’, O5’, OP1, OP2, P; A for search mode MTERF1, superhelical Cα atoms and all DNA heavy atoms, B for search mode MTERF1, only DNA backbone atoms (MTERF1 fully unrestrained). Force constant: harmonic force constant for Cartesian restraints. For search mode, force constants were 1/10th of those illustrated above, except for stage 8, which had no force applied. The Berendsen thermostat (19) was used for all stages of MD including production. Stage 2 and 3 used bath coupling constants of 0.1 ps; all subsequent stages of MD including production used 0.5 ps coupling constants.

Section 8. Docking B-DNA to MTERF1 from the recognition structure

To determine whether the conformation of MTERF1 in recognition mode (the crystal structure) could bind B-DNA, a docking procedure followed by MD was used to evaluate the energetics of the complex. All poses in which B-DNA was docked into the binding cleft of MTERF1 were tested for stability using our fully atomistic MD procedure (see Section 7) and found to be energetically unstable (>10⁸ kcal/mol). Some poses appear reasonable (Figure S7A and B) and indeed MD energies were stable, though not structurally stable (RMSD > 7 Å after only 50 ns). Closer inspection of these poses reveals that the C-site of the DNA would clash if extended. To show these poses were simply artefacts of using short DNA, the DNA in all the poses were lengthened and the systems subjected to the same equilibration procedure. As expected, every single docked poses was not energetically viable (>10⁸ kcal/mol) due to van der Waals clashes.
Section 9. RMSD analysis of control and apoMTERF1 simulations

To establish a baseline for the apo MTERF1 conformational change, 4 independent 1.5 µs control simulations of the MTERF1-DNA specific complex were performed. Small structural fluctuations were expected and the conformation of MTERF1 would on average be similar to the crystal structure. To quantify how similar our MD structures were to the reference crystal structure, the root mean square positional deviations (RMSD) between our MD snapshots and the equilibrated crystal structure were measured using cpptraj (20). The terminal nucleotides (three per strand, for both 5’ and 3’ ends) were excluded to avoid overweighting deviations arising from end-fraying (21). The evolution of RMSD in the control simulations displayed in Figure S8A shows that the conformation of MTERF1 throughout the simulations remains similar to that of the reference. In recognition mode, MTERF1 and the DNA remain tightly bound with relatively little conformational fluctuation compared to apo MTERF1.

Figure S7. Docking B-DNA to MTERF1 from the recognition mode structure fails to produce poses in which the protein tracks the major groove. (A) One of two DOT2.0 docked poses in which MD equilibration energy did not result in high energy (>10^9 kcal/mol) using the exact procedure used to dock low-pitch apo MTERF1 to B-DNA and to dock MTERF1 from recognition mode to the corresponding unwound DNA from the crystal structure. (B) The second pose. (C) After 75 ns of unrestrained MD of the pose from (A), MTERF1 dissociates from the DNA (all atom RMSD > 7 Å) (D) After 50 ns of unrestrained MD of the pose from (B), MTERF1 dissociates from the DNA (all atom RMSD > 8 Å).
Figure S8. RMSD analysis of apo MTERF1 and holo MTERF1 (specific complex) unrestrained MD simulations. (A) RMSD of the specific MTERF1-DNA complex (crystal structure) protein backbone, excluding the first mterf motif and the C-tail, and the DNA C1' and P atoms, excluding the 3 terminal base pairs at each end. (B) RMSD of only the protein in the specific complex using the same atoms as in (A). (C) RMSD of MTERF1 in the unbound protein simulations using the same atoms as in (A) and (B).

The RMSD was also used as a more routine metric of conformational heterogeneity to help gauge the degree to which our helical analyses resolve the hypothesized apo MTERF1 conformational change. Thus the time-resolved RMSD of apo MTERF1 is reported in Figure S8C. The analysis highlights the potential problems of a crude metric of conformational such as RMSD. Simulations in which the RMSD was lower than other simulations never sampled structures with superhelical pitch < 42 Å while simulations with high RMSD never sampled low superhelical pitch.

Section 10. The similarity of ANM and MD lowest frequency motions

Table S3. ANM and MD eigenvector RMSIP similarity analysis

Root mean square inner product (RMSIP)(22) of all top ten eigenvectors (1 to 10), the top four eigenvectors (1 to 4), the top three (1 to 3), and the top two (1 to 2). RMSIP provides a global similarity comparison of eigenvector
overlaps, accounting for the possibility that corresponding MD and ANM eigenvectors are not in the same order. The eigenvectors defining each of the ANM modes and MD PCs would be parallel if they were identical and orthogonal if completely unrelated; the dot product of parallel vectors is zero if they are orthogonal and one if they are parallel.

<table>
<thead>
<tr>
<th>ANM (mode)</th>
<th>MD (PC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 to 2</td>
<td>1 to 3</td>
</tr>
<tr>
<td>1 to 2</td>
<td>0.776</td>
</tr>
<tr>
<td>1 to 3</td>
<td></td>
</tr>
<tr>
<td>1 to 4</td>
<td></td>
</tr>
<tr>
<td>1 to 10</td>
<td></td>
</tr>
</tbody>
</table>
Section 11. Histograms of holo MTERF1 and apo MTERF1 helical parameters

Figure S9. Histograms of helical parameters of holo and apo MTERF1 MD ensembles. Probabilities of holo MTERF1 pitch (top left), holo MTERF1 radius (top right), apo MTERF1 pitch (bottom left), and apo MTERF1 radius (bottom right).
Section 12. Productive, energetically stable nonspecific docked complexes

Figure S10. The 12 models of the nonspecific complex that best track the major groove obtained by docking low-pitch apo MTERF1 structures to B-DNA. The protein conformations that were used to generate the pose has the following helical parameters (pitch, radius, sweep): for (A) and (B): 41.1 Å, 9.8 Å, 372°; for (C): 41.6 Å, 10.1 Å, 368°; for (D), (E), (F) and (G): 40.3 Å, 11.4 Å, 340°; for (H) and (I): 39.8 Å, 16.2 Å, 276°; for (J): 37.8 Å, 16.0 Å, 287°; for (K) and (L): 34.5 Å, 9.8 Å, 389°.
Section 13. Quantifying how tightly MTERF1 binds DNA

To complement the analysis in Section 12, the area shared by MTERF1 and DNA was evaluated using the Linear Combination of Pair Overlaps (LCPO) method (23). Since we were not calculating forces, the non-recursive LCPO algorithm (less accurate than the recursive algorithm used in GB simulations) was sufficient. The cpptraj module in AmberTools (20) was used to perform the calculation and a bash script to post-process the three sets of surface areas – one half of the protein surface area plus one half of the DNA surface area minus one half of the complex surface area.

Figure S11. The shared surface area of MTERF1 and DNA in the 12 distinct search mode complexes. The abscissa is time, in units of µs.
Figure S12. The shared surface area of MTERF1 and DNA in the recognition complex for each of the four independent simulations. The abscissa is time, in units of µs.
Section 14. Measuring MTERF1 translocation along DNA

To quantify MTERF1 sliding a similar approach to that used by the Levy lab (24) was used. The metric was used to distinguish geometries that were sliding, hopping, or freely diffusing. Here, the measurement has a specific meaning here for two reasons. First, MTERF1 wraps around the DNA, so the protein centre-of-mass (COM) falls near the DNA axis. Second, we use the sliding distance in combination with our major groove-tracking distance to ensure the sliding distance is reporting on translocation and not unbinding.

![A](image1.png) ![B](image2.png)

Figure S13. Visualization of our sliding distance metric. The protein (grey) and DNA (green) are shown as translucent cartoons to highlight the centres-of-mass. The centre-of-mass (COM) of the superhelical residues is shown as a grey sphere. The COM of the C1’ atoms in the DNA, excluding 4 bases at each end of the duplex to prevent end-fraying artefacts, is shown as a green sphere. (A) The equilibrated snapshot (0.5 µs) of the search mode. (B) A snapshot of the search mode (1.4 µs) that shows increased distance between the COM of the protein and the DNA.

REFERENCES


