

Intrinsic flexibility of B-DNA: the experimental TRX scale

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ABSTRACT

B-DNA flexibility, crucial for DNA–protein recognition, is sequence dependent. Free DNA in solution would in principle be the best reference state to uncover the relation between base sequences and their intrinsic flexibility; however, this has long been hampered by a lack of suitable experimental data. We investigated this relationship by compiling and analyzing a large dataset of NMR ³¹P chemical shifts in solution. These measurements reflect the BI ↔ BII equilibrium in DNA, intimately correlated to helical descriptors of the curvature, winding and groove dimensions. Comparing the ten complementary DNA dinucleotide steps indicates that some steps are much more flexible than others. This malleability is primarily controlled at the dinucleotide level, modulated by the tetranucleotide environment. Our analyses provide an experimental scale called TRX that quantifies the intrinsic flexibility of the ten dinucleotide steps in terms of Twist, Roll, and X-disp (base pair displacement). Applying the TRX scale to DNA sequences optimized for nucleosome formation reveals a 10 base-pair periodic alternation of stiff and flexible regions. Thus, DNA flexibility captured by the TRX scale is relevant to nucleosome formation, suggesting that this scale may be of general interest to better understand protein–DNA recognition.

INTRODUCTION

The DNA genomic sequence of numerous organisms is now available. Nevertheless, this considerable amount of

data does not by itself illuminate the genetic messages contained in these genomes. It is largely because these messages must be deciphered by numerous DNA-binding proteins such as transcription factors. The DNA also influences chromatin folding via its recruitment of histone octamers to form the nucleosomes. So, the cellular DNA is continuously ‘read’ by proteins. Specific and non-specific DNA–protein binding are known to depend on the conformational preferences of the nucleotidic sequences (1–10). Thus, understanding DNA–protein interactions requires knowledge of the intrinsic flexibility of DNA sequences.

Quantifying the sequence-dependence of DNA flexibility remains a major goal of molecular biology. The flexibility of DNA can be studied at various levels of details, from base-pair step properties to the long range behavior of the double-helix considered as a flexible rod. Most DNA–protein contacts engage 15–20 base pair (bp) DNA fragments (11), apart from cases such as the nucleosome where the interface covers 146–147 bp. Thus, dinucleotide or tetranucleotide sequences provide an appropriate (and ultimately required) resolution to address intrinsic DNA flexibility as a determinant of DNA/protein interaction. One approach to quantify the dinucleotide/tetranucleotide flexibility is to observe their conformational spread on populations large enough for meaningful statistics. Assembling such a dataset is not trivial.

A range of experimental methods has been used to capture DNA deformability (12). Analyses of the variability in X-ray structures of B-DNA devoid of proteins contributed to characterize the conformational properties of the ten dinucleotides in such context. This approach has well-known limitations, notoriously the crystal packing forces which impact the details of DNA structures (13–16) and the effective temperature of the X-ray

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statistical ensemble (12). A large set of crystallized DNA-protein complexes was also analyzed (5), assuming that deformations in bound DNA are likely to reflect its intrinsic structural properties, and that protein-bound DNA may be less subjected to crystal packing than free DNA. The dataset was deemed large enough to average the DNA distortions imparted by the proteins. Extracting the fluctuations and correlations of bound DNA, structural parameters yielded a complete set of sequence-dependent empirical energy functions, providing a quantification of the dinucleotide step malleabilities, aimed at predicting the energetic cost of DNA deformations upon protein binding. The ranking of step malleability depended on the helicoidal parameter considered, but YpR appeared the most easily deformed steps. This approach was applied to the question of DNA folding in chromatin, accounting quite well for nucleosome locations in six nucleosome-positioning sequences (17,18).

However, a comparison of free and bound X-ray DNA (19) showed that, in contrast with free DNA, all DNA/protein oligomers exhibited numerous north sugars and a large range of $\alpha/\beta/\gamma$ backbone conformations. In addition, the range of helical parameter values are larger in bound than in free DNA (5). Yet, the couplings between various helical parameters (5) or between helical parameters and BI/BII backbone states (19) were qualitatively retrieved in free and bound DNA ensembles. Thus, free and bound X-ray DNA structures obey the same overall intrinsic mechanics. This reinforces the idea that proteins exploit the free DNA dynamical properties upon binding. At the same time, proteins are able to extend the DNA energy landscape, probably altering some conformational energetic barriers, in particular in the DNA backbone (20). It is thus unclear to which extent the sequence-dependent flexibility of free B-DNA in solution can be extrapolated in practice from the distortions observed in X-ray bound DNA structures.

Two recent reports analyzed datasets mixing X-ray and NMR DNA structures (21,22), but the accuracy of DNA NMR structures is still an issue. NMR provides experimental restraints which are combined with molecular simulations. With unlabeled DNA, the low density of protons, together with relatively frequent overlaps in the NOE peaks, limit the number of measured distances. Hence, the derived structures may depend on the refinement protocols and the force-fields (23). Developments requiring labeled DNA (24–27) increase the number of observables, and improve the quality of the refined structures, although the interpretation of these additional data is not straightforward and may introduce specific biases (28). Approaches using numerous restraints combined with potentials of mean-force (29) or extensive MDs in explicit solvent (23) seem to provide another avenue to produce reliable structures of DNA free in solution, with a representation of their dynamics. Unfortunately, such accurate structures are too few to provide a general guide to DNA intrinsic flexibility.

Computer modeling methods have frequently been used to study DNA deformability, based on simple potential energy calculations or molecular dynamics (MD) in

explicit solvent (1,30), reviewed in (12). In particular, the elastic properties of the ten complementary dinucleotides were evaluated from MD trajectories (30–35). Studies in the same vein were performed with a variety of MD protocols: (i) with different force-fields [AMBER parm94: (31,32,35,36); parm99: (1,33,34); parmbsc0 and CHARMM27: (30)], (ii) unconstrained (1,30–33,35,36) or under stress (34), and (iii) considering the six helicoidal parameters [twist, roll, tilt, rise, shift and slide (31–33,36)] or focusing on specific parameters [twist: (34); roll: (35)]. These studies reached convergent conclusions in identifying three distinct groups of steps in terms of purine (R)/pyrimidine (Y) classification: the flexible YpR, intermediate RpR•YpY, and stiff RpY steps. However, force-field limitations still exist (23,37–40). The comparison of NMR observables to their MD counterparts has unveiled shortcomings in the representation of salient details of B-DNA structure and dynamics by the commonly used DNA force-fields (23). It showed that the best current force-fields do not reproduce the modulation of DNA dynamics by its sequence, although they implement much of the intrinsic mechanical couplings within B-DNA. Therefore, the characterization of DNA sequence-dependent intrinsic flexibility still requires experimental input.

Here, we approach the intrinsic flexibility of DNA by probing its backbone properties in solution, in absence of proteins. The phosphate groups in B-DNA can adopt two conformations, the BI and BII states (Figure 1). They differ in the torsion angles ϵ and ζ which are *trans/g-* in BI ($\epsilon-\zeta$ centered at -90°) and *g-/trans* in BII ($\epsilon-\zeta$ centered at $+90^\circ$). These two states, initially identified from crystallographic studies (41), were then detected in NMR by measurements of $^3J_{H3'-P}$ spin-spin coupling constants and/or ^{31}P chemical shifts (δP) (42,43). Backbone and deoxyribose conformations are correlated (19,44,45). Most importantly, BI and BII conformations are also mechanically coupled to the DNA helicoidal parameters of roll, twist and X-disp, related to curvature, winding and groove dimensions, respectively (19,23,46–52). Furthermore, the propensity to undergo the BI/BII transition is sequence dependent (19,48,53). A method was recently devised to quantify for the first time the BI and BII populations in solution, from the very sensitive and accurate measurements of δP (48). Indeed, NMR studies highlighted strong linear correlations between δP and the three measured internucleotide distances $H2'_i-H6/8_{i+1}$, $H2''_i-H6/8_{i+1}$ and $H6/8_i-H6/8_{i+1}$ (48,54). These correlations, also present in very high resolution X-ray structures where δP can be substituted by ($\epsilon-\zeta$) values, allowed to establish a quantitative relation between δP and BI/BII ratios. They also confirm that the BI/BII ratio is associated to inter base pair parameters. The couplings between the BI \leftrightarrow BII equilibrium, deoxyribose conformational exchange (44) and helicoidal parameters (48) establish that the BI \leftrightarrow BII motions reflect the overall B-DNA structure and dynamics in solution.

In the present work, 328 δP values previously reported for various B-DNA sequences in solution were collected. The influence of the dinucleotide and tetranucleotide sequence context on δP was analyzed. It provides

evidence that DNA flexibility of a given step is primarily dominated by the corresponding dinucleotidic sequence, although secondary sequence effects beyond this immediate environment are observed on several steps. This forms the basis of a new and general experimental scale that characterizes the intrinsic properties of all 10 DNA complementary dinucleotides in terms of Twist, Roll and X-disp parameters, and is thus called the TRX scale.

Applying this new concept to sequences identified to preferentially bind to histone octamers and form nucleosomes (55–57) illustrates how the TRX scale provides new insights into the structural organization of DNA and its biological importance.

MATERIALS AND METHODS

BI/BII propensities in solution, from NMR

We previously (48) explained how ^{31}P NMR chemical shifts could be converted to BII percentages by the equation $\text{BII}(\%) = 143 \delta\text{P} + 85$ for δP referenced to phosphoric acid, or $\text{BII}(\%) = 143 \delta\text{P} + 621$ for δP referenced to trimethylphosphate. This conversion is based on the linear cross-correlations observed between the three $\text{H}2'_i - \text{H}6/8_{i+1}$, $\text{H}2''_i - \text{H}6/8_{i+1}$, and $\text{H}6/8_i - \text{H}6/8_{i+1}$ sequential distances and δP in NMR or $\varepsilon - \zeta$ values in X-ray structures. Both δP and $\varepsilon - \zeta$ reflect the interconversion between the backbone BI ($\varepsilon - \zeta < 0^\circ$) and BII ($\varepsilon - \zeta > 0^\circ$) states. The same method was used to compile the BII percentages per dinucleotide presented here. These statistics were gathered from a total of 328 ^{31}P chemical shifts reported in the literature for B-DNA without mismatches, chemical modifications or ligands (see Supplementary Table S1). The BII propensities per complementary dinucleotides ($\text{N}_i\text{pN}_{i+1} \bullet \text{N}_j\text{pN}_{j+1}$) are the half-sum of the BII percentages for the two corresponding phosphates. BI propensities are equal to 100-(BII propensities).

Crystallographic datasets

The X-ray structures of free B-DNA include the 37 DNA oligomers (2 octamers, 23 decamers, 11 dodecamers, 1 oligomer with 15 bp) of 2.0 Å resolution or better, the same dataset used in a previous analysis (19). These structures are all double-stranded DNAs, and do not include mismatches or modified bases, sugars or backbones.

Numerous nucleosome core particle (NCP) X-ray structures are available, but, to benefit from reliable structural details, we examined only the structures with resolution ≤ 2.5 Å, crystallized without histone mutants or groove ligands [PDB codes: 1EQZ (58), 1KX5 and 1KX3 (59), 1M19 (60), 2CV5 (61)]. 1EQZ and 1KX3 structures are quasi identical (RMSD ~ 0.8 Å); the significance of the 2CV5 DNA fine structure appeared problematic, due to tight DNA–DNA contacts within the crystal that notably altered the DNA conformation (61). Therefore, the NCP DNA structures analyzed here are those from entries 1KX3, 1M19 and 1KX5.

Nucleosomal sequences

The sequences used in the ‘Implications for DNA recognition by proteins’ section came from Dr. Segal’s web site <http://genie.weizmann.ac.il/pubs/nucleosomes06/index.html>. The sequences are given in Supplementary Table S4.

Structure analysis

Analyses of DNA structures were carried out using CURVES (62), which calculates the optimal helical axis and a complete set of conformational and helicoidal parameters. Analyses were made in terms of both local and global parameters; however, since both analyses give quasi-identical values, only the global values are reported here. X-disp values are given for base pairs, i.e. for strand 1 with strand 2.

The phosphate conformations were analyzed in terms of BI and BII states, defined by the values of the two torsion angles ε ($\text{C}_4 - \text{C}_3 - \text{O}_3 - \text{P}$) and ζ ($\text{C}_3 - \text{O}_3 - \text{P} - \text{O}_5$), with BI corresponding to $\varepsilon/\zeta = t/g$ ($\varepsilon - \zeta < 0^\circ$ and centered around -90°), and BII corresponding to $\varepsilon/\zeta = g/t$ ($\varepsilon - \zeta > 0^\circ$ and centered on $+90^\circ$).

Fourier analysis

Values from the TRX scale for all the dinucleotidic steps along the DNA sequences were taken as input signal. Subtracting the average value $\langle \text{trx} \rangle = (\text{trx}(1) + \dots + \text{trx}(N))/N$ gives a centered signal $b(n) = \text{trx}(n) - \langle \text{trx} \rangle$. The Fourier series is the sum, over the N steps of the segment, of plane waves

$$f(k) = N^{-1/2} \sum_{n=1}^N b(n) \exp(-i2\pi kn).$$

The function f is periodic of period 1; as such, it is completely specified by its values in the unit interval $[0,1]$, best seen as a unit circle. All our Fourier analyses (Figure 6 and Table 2) represent the Fourier intensity or power spectrum $|f(k)|^2$.

Correlation length

The Fourier intensity at period ~ 10 is not enough to match the relative affinities for nucleosome formation. Some coherent flexibility/stiffness is also required. The dinucleotides can be classified into two categories delimited by the average TRX score of 21 (Table 1): those, flexible, with a significant propensity to adopt BII conformation (TRX score > 21 , coded as +1) and those, stiffer, predominantly confined to BI (TRX score < 21 , code -1). Then the DNA sequence is naturally subdivided into blocks of consecutive equal values, the flexible blocks of consecutive +1 and the stiff blocks of consecutive -1 . The mean length of these blocks, readily computed from histograms of the type depicted in Figure 7, defines a characteristic length, one for each category: the flex ξ_f and the stiff ξ_s characteristic lengths. The overall average is the correlation length $\xi = (\xi_f + \xi_s)/2$.

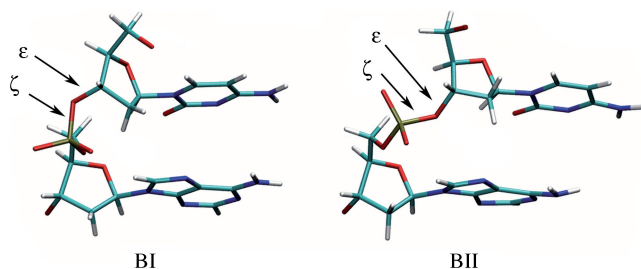


Figure 1. Illustration of the BI (left) and BII (right) phosphate linkage conformations with a CpA dinucleotide. BI and BII backbone conformations differ in the torsion angles ϵ and ζ which are respectively *trans/g-* in BI ($\epsilon - \zeta = -90^\circ$) and *g-/trans* in BII ($\epsilon - \zeta = +90^\circ$).

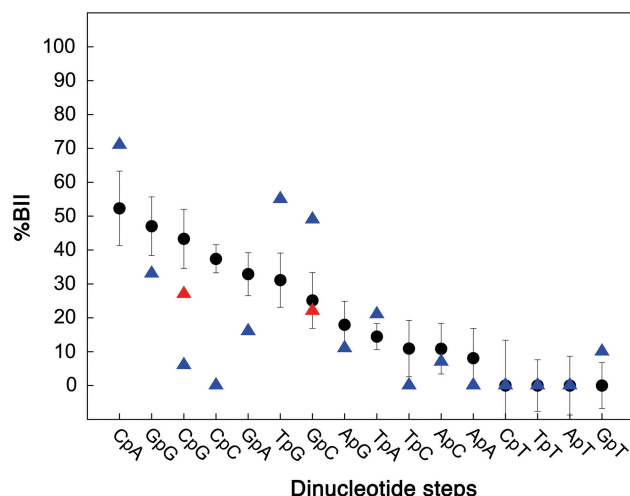


Figure 2. Influence of the dinucleotide sequences on BII percentages in B-DNA. BII percentages (%BII) of free B-DNA inferred from δP in solution (black circles, with vertical bars for standard deviations) compiled from data published in the literature (see Supplementary Table S1). The conversion of δP in terms of BII percentage, using a published procedure (48), is detailed in ‘Materials and Methods’ section. BII percentages in solution are compared with those extracted from X-ray structures (triangles; blue: all structures, red: decamers only).

RESULTS

BI/BII dinucleotide sequence effect in B-DNA

The phosphate linkages in B-DNA adopt both the BI ($\epsilon - \zeta < 0$) and BII ($\epsilon - \zeta > 0$) states (Figure 1). ^{31}P chemical shifts (δP), very accurately measured in solution by NMR, reflect the BI \leftrightarrow BII equilibrium (48,63). We collected from the literature a large set of 328 δP values previously reported for various B-DNA sequences in solution, excluding the phosphate linkages submitted to end effects at the sequence termini (Supplementary Table S1). Although monovalent counterions can modulate the BI/BII equilibrium (54), these subtle effects are expected to be averaged in our data-set, the δP being measured in variations of the physiological conditions.

Each of the 16 B-DNA dinucleotides is characterized by a specific δP average value (Supplementary Figure S1), with small standard deviations (0.06 ppm on average). This influence of dinucleotide sequences on the BI \leftrightarrow BII equilibrium is presented in Figure 2, translating the δP

Table 1. Influence of DNA base sequence on the BII percentages in free DNA in solution

	<i>N</i>	%BII	TRX score
CpG•CpG	25	43•43	43
CpA•TpG	28	52•31	42
GpG•CpC	11	47•37	42
GpC•GpC	22	25•25	25
GpA•TpC	25	33•11	22
TpA•TpA	12	14•14	14
ApG•CpT	19	18•0	9
ApA•TpT	17	11•0	5
ApC•GpT	23	8•0	4
ApT•ApT	22	0•0	0

The DNA sequence is expressed in terms of the 10 complementary dinucleotides, of frequency *N* in the NMR data collected from the literature. The BII percentages (%BII) are given for each partner in a complementary dinucleotide. The average standard deviation of %BII is ± 8 . The TRX scores are the half-sums of the %BII observed for two facing phosphates in a complementary dinucleotide. The %BII and TRX scores higher (in bold) and lower than average (which is 21) correspond to enhanced and restricted flexibilities, respectively. A maximal flexibility on this scale corresponds to a TRX score of 50.

from our data set to BII percentages with an established method (48) (see ‘Materials and Methods’ section). This analysis shows that (i) the averaged BII percentage in solution is 21% and (ii) 7 out of the 16 dinucleotides exhibit BII populations higher than average. These steps can thus be regarded as BII-rich steps. The maximal flexibility is observed for the phosphates of CpG, CpA and GpG, with a BI/BII ratio close to 1 (50% BI, 50% BII).

Considering the BII propensities of the 10 possible complementary dinucleotide sequences $N_i p N_{i+1} \bullet N_j p N_{j+1}$ (Table 1; *N*: any base) highlights that the facing phosphates tend to exhibit similar behaviors overall. In a first group containing ApN•NpT and TpA•TpA, the phosphates are very rarely in BII. This group, mainly confined in BI•BI, is thus characterized by a restricted backbone flexibility. In another group of dinucleotide steps (GpG•CpC, CpG•CpG, GpC•GpC and CpA•TpG) the facing phosphates can adopt BI and BII conformations, with a higher-than-average BII percentage. This family exhibits an enhanced backbone flexibility, potentially able to explore the three possible combinations, BI•BI, BI•BII (and its counterpart form BII•BI) or BII•BII. GpA•TpC is the only case where the two facing phosphates exhibit behaviors in marked contrast, with GpA able to significantly populate BII, but not TpC.

The BII percentages in solution can be compared with those extracted from crystal structures of free B-DNA (Figure 2). Considering the 16 dinucleotides, the backbone of ApA, (A/T)pC and NpT is overwhelmingly in BI in both environments. In contrast, the solution and crystal states often differ for steps with a significant propensity to populate BII. These propensities are either lower (CpG, CpC, GpG, GpA) or higher (CpA, TpG, GpC) in crystals than in solution. In X-ray structures, the behavior of the phosphates in GpG•CpC remains ambiguous since these steps are underrepresented (6 cases). Dinucleotides GpC and CpG most likely

illustrate crystalline packing biases. Indeed, a large majority of GpC and CpG (34/44 and 51/69, respectively) are located at dodecamer ends, where adjacent molecules are in contact *via* their grooves (64). There are no such contacts in the decamers that are stacked end-to-end, where the GpC and CpG BII percentages are very close to those inferred from NMR data (Figure 2). The remaining apparent discrepancy concerns CpA•TpG, due to a tetranucleotide sequence bias in the crystal structures, explained in the next section.

In addition, the NMR BII propensities do not fit with those from the protein-bound DNA X-ray structures. Actually, a previous study (19) comparing high resolution X-ray structures of free and protein-bound DNA showed that, although the global DNA intrinsic mechanics is preserved (for instance the coupling between several helical parameters and the backbone states), the BII populations are globally reduced in protein-bound DNA.

For free B-DNA, taking the solid state biases into account allows to sufficiently reconcile sequence effects in solution with their crystal counterpart. The fact that the BII propensities follow the same trends overall in both environments is a very strong indication that these trends represent intrinsic properties of the DNA molecule.

BI/BII tetranucleotide sequence effect in B-DNA

Experimental (65–68) and theoretical (21,33,53,69) results indicated that the behavior of some dinucleotides in B-DNA is influenced by their two bracketing nearest neighbors (flanking bases). The size of our NMR data set does not allow the analysis of all the 136 tetrameric combinations. A reasonable simplification is to consider instead the 3'- and 5'-contexts of the dinucleotides in terms of purine R or pyrimidine Y (Y–Y, R–R, Y–R, R–Y). In our NMR data set, these four categories are significantly populated for most dinucleotides (Supplementary Table S2).

The flanking bases do influence the BI ↔ BII equilibrium of the central phosphate in most BII rich dinucleotides (Figure 3). Indeed, a Y–R context reinforces the BII populations in CpA, GpG, CpG, TpG. The BI-rich steps, together with GpA, appear mainly indifferent to their flanking bases (Figure 3). Yet, in this category, it seems possible to obtain substantial BII populations in YApAR, RTpCY and RApCY (Figure 3), but YApAY and RTpCY are represented by only two cases, and the four δP collected for RApCY are especially scattered (from -0.39 to -0.19 ppm).

Overall, the four possible sequence contexts are so unequally populated in crystals that it prevents a systematic analysis of the tetranucleotide effect in this environment. However, the tetrameric sequence effect can account for the differences observed in Figure 2 between the NMR and X-ray BII percentages of CpA•TpG and CpG. Indeed, 14 out of 21 CpA•TpG steps in the X-ray data belong to the Y(CpA/TpG)R pattern that enhances BII and thus leads to dinucleotide average BII percentages higher than those found in solution. Conversely, 13 out of 18 CpG•CpG in crystals are surrounded by R–Y•R–Y

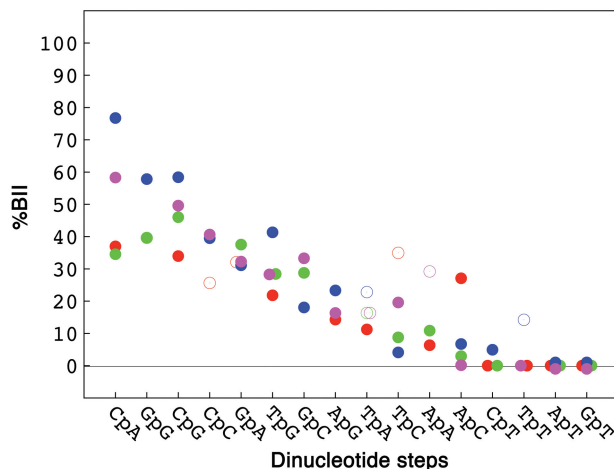


Figure 3. Influence of the tetranucleotide sequence on the BII percentages of the central phosphate group. BII percentages (%BII), compiled and translated from δP values published in the literature (see ‘Materials and Methods’ section and Supplementary Table S1), are given for the dinucleotide steps in four different tetranucleotide contexts: R–R in green, R–Y in red, Y–R in blue and Y–Y in magenta. Categories that contain more or less than three cases are represented by filled and open circles, respectively.

or R–R•Y–Y, reducing their apparent global BII propensity.

More work is needed to understand the tetranucleotide effects, probably under-estimated, although potentially crucial for DNA–protein recognition (23,68). For the present purpose, this secondary influence does not disrupt the flexibility rankings detected at the dinucleotide level. In sum, the BI/BII propensities of phosphate groups are primarily controlled at the dinucleotide level, finely modulated at the tetrad level.

Relationship between helical parameters and backbone states

X-ray B-DNA structures are useful to demonstrate the general intrinsic mechanics of DNA. The principles underlying this mechanics do not depend on the sequence, and sound statistical analyses can be undertaken on a large set of high resolution oligomers. Such approaches have established that roll and twist values are correlated in both free and protein-bound DNA (5,46). Crucially, these inter-base descriptors depend on the conformations of the two phosphates facing one another across the strands (19,23,46) (Figure 4). BI•BI is associated with positive rolls ($2.7 \pm 4.1^\circ$), whereas BI•BII ($-4.2 \pm 5.1^\circ$) and especially BII•BII ($-10.87 \pm 2.1^\circ$) correspond to negative rolls. The twist values associated to BI•BI ($33.9 \pm 5.0^\circ$) are lower than those with BI•BII ($39.4 \pm 2.8^\circ$) or BII•BII ($48.1 \pm 4.1^\circ$). The only exception is for the semi-flexible GpA•TpC steps, their twist and roll appearing rather uncorrelated, as previously noticed (5). Phosphate conformational combinations are also correlated to the slide (Supplementary Figure S2) that becomes positive in BII•BII steps. This parameter is very strongly correlated to the roll (5,18). For instance, roll and slide variations work in concert to generate

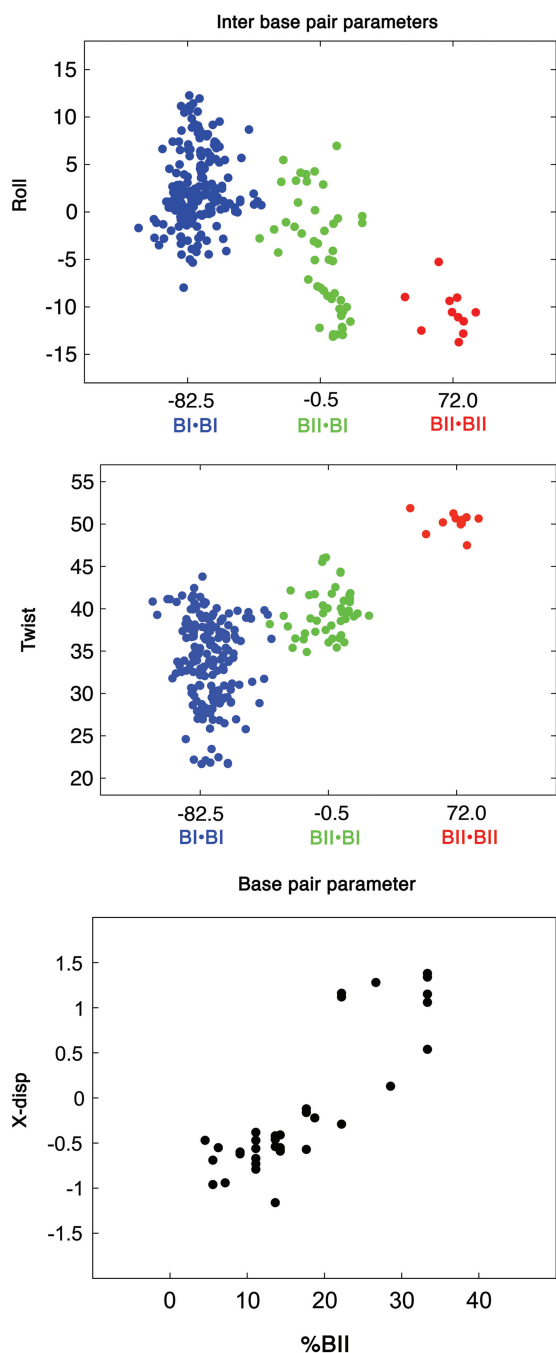


Figure 4. Influence of BII phosphates on helical descriptors. Inter base pair parameters: Twist ($^{\circ}$) and roll ($^{\circ}$) plotted versus $(\epsilon - \zeta)_{av}$, averaged on the two facing phosphates. BI•BI (blue, $(\epsilon - \zeta)_{av} = -82.5^{\circ}$), BI•BII (green, $(\epsilon - \zeta)_{av} = -0.5^{\circ}$) and BII•BII (red, $(\epsilon - \zeta)_{av} = +72^{\circ}$) are the three possible conformational combinations of facing phosphates. Base pair parameter: X-disp (\AA), averaged along whole oligomer sequences plotted versus the BII percentage (%BII) per oligomer (number of BII phosphates/total number of phosphates). The data were extracted from high resolution X-ray structures of free DNA (19).

sharp bending in DNA/protein complexes (18). Due to this equivalence, we do not elaborate further on the slide in the following.

BII phosphates also act on groove dimensions, as previously shown by solid state NMR studies on fibrous

DNA (51) and by analyzing X-ray data (19,46). Indeed, they displace base pairs towards the major groove, influencing the X-disp parameter. X-disp is defined for one base pair, positive and negative values reflecting displacement towards the major and minor grooves, respectively. Analysis of X-ray structures detects no more than two BII among the four phosphate linkages pIp•pJp surrounding one base pair I•J. This is consistent with previous X-ray analysis (19,46) and with a NMR study of S^2 order parameters that highlighted that a BII rich phosphate precludes high BII populations in its 5' and 3' neighbors (44). Examining the X-disp values in the present X-ray data set yields $X\text{-disp} = -0.6 \pm 0.4$, -0.2 ± 0.5 and $+0.85 \pm 0.7 \text{\AA}$ when zero, one and two BII linkages surround the central base pair I•J. Several proximal (but not successive) BII phosphates enhance this effect, preserving an optimal base stacking (47,53). So, the average X-disp is $\geq 1.0 \text{\AA}$ for deca- and dodecamers in which more than 22% of phosphates are simultaneously in BII (for instance, 22% correspond to 4 BII on the 18 phosphates encountered in a decamer) (19,23) (Figure 4). Such cumulative shifts in X-disp values mediate the influence of BII rich regions on the major groove overall shape, making it convex (depth of $-1.0 \pm 0.5 \text{\AA}$) instead of concave (usual depth being $+5 \pm 2 \text{\AA}$).

The helical parameters are not directly accessible by NMR, but their mechanical couplings are revealed through linear correlations between the δP values and the three sequential internucleotide distances $H2'_i - H6/8_{i+1}$, $H2''_i - H6/8_{i+1}$ and $H6/8_i - H6/8_{i+1}$ (48,54), related to the twist and roll parameters (49). Also, the relationship between the BI/BII conformations and the base pair displacements was confirmed by NMR on fibrous DNA (51). In addition, these couplings were retrieved by modeling of DNA in explicit solvent (46,52), and in simulations under NMR constraints (23).

So, different facing backbone BI/BII conformations define sub-states in B-DNA, influencing both the local (e.g. roll and twist) and neighboring structure (e.g. X-disp) of the double helix. In particular, strong negative roll and very high twist appear distinctive of synchronous facing BII phosphates.

The TRX scale

The strong correlations between BI \leftrightarrow BII phosphate populations and helical parameters mean that the BII propensities characterizing each of the ten complementary dinucleotides reflect the intrinsic flexibility of the corresponding twist, roll and X-disp. Consequently, the scale of BII propensities at the dinucleotide level (Table 1) is a realistic and general representation of the sequence dependency of DNA intrinsic plasticity. Considering the above-mentioned similar average behavior of facing phosphates, this flexibility can be quantified by the half-sum of the BII propensities of the two facing phosphates in a complementary dinucleotide (right-most column in Table 1). This constitutes an experimental scale of local intrinsic flexibility, called 'TRX' in reference to the Twist, Roll and X-disp parameters that it represents.

The steps with a low TRX score (low BII propensity) have a restricted flexibility: low or moderate twists associated with null or positive rolls, respectively; those bases tend to be minimally displaced away from the axis of the double helix. The steps associated with higher than average TRX score (>21) are characterized by plurimodal conformational populations. They can explore a larger conformational space, from low twists/positive rolls ('BI state') to high twists/negative rolls ('BII state'); the movement of these bases towards the major groove is facilitated. High plasticities are observed for steps CpA•TpG, CpG•CpG, GpG•CpC, with TRX scores of 42–43. These values are close to 50, the maximal flexibility in the context of the TRX scale (50% of the time in BI, 50% of the time in BII). Thus, the value of the TRX score has a physical meaning, directly related to the BII populations in free B-DNA in aqueous solution. These populations are a signature of the intrinsic mechanical flexibility of the associated base steps.

The examination of the variability of inter-base helical parameters in the X-ray datasets of free DNA [our dataset and another reported on <http://rutchem.rutgers.edu/~olson/pdna.html> (5)] globally agree with the TRX scale. The only serious discrepancy is the rank of step TpA. According to the X-ray data, this step is more flexible than GpG•CpC, CpG•CpG and GpG•CpC. Conversely, the δP based observations classify TpA in the 'stiff' category while GpG•CpC, CpG•CpG and GpG•CpC are ranked flexible (Table 1). But in the X-ray dataset GpG•CpC are poorly represented (less than 2% of the dinucleotide steps) and GpC•GpC and CpG•CpG are subjected to intermolecular contacts, as explained above. Another source of discrepancy could come from the sugar north \leftrightarrow south exchange, coupled to the BI \leftrightarrow BII equilibrium (44), but possibly underestimated in solid phase, in particular on cytidine residues (27). So, the TpA flexibility rank deduced from X-ray structures could be biased by an incorrect estimation of the properties of GpG•CpC, CpG•CpG and GpG•CpC steps.

The TRX scale promises to be helpful for deciphering the intrinsic flexibility of free DNA underpinning DNA–protein interactions. Indeed, intrinsic structural preferences of free DNA, and its possible pre-organization, are expected to limit the energetic and entropic penalties upon protein binding.

Implications for DNA recognition by proteins

To demonstrate the concrete use of the TRX scale, we illustrate how it can be applied to DNA sequences bound preferentially by the histone core. The nucleosome core particle (NCP), consisting of 146–147 bp of DNA wound twice around an octameric core of four histone proteins, is the fundamental building block of packaged DNA in eukaryotic cells. Earlier studies on chicken nucleosomal DNA established that some DNA sequences favored the NCP formation (70,71). Sequences having an exceptionally high affinity for histone octamers were then identified using the SELEX technique (55–57), leading to identify a consensus for the central 73 bp (56). Using large

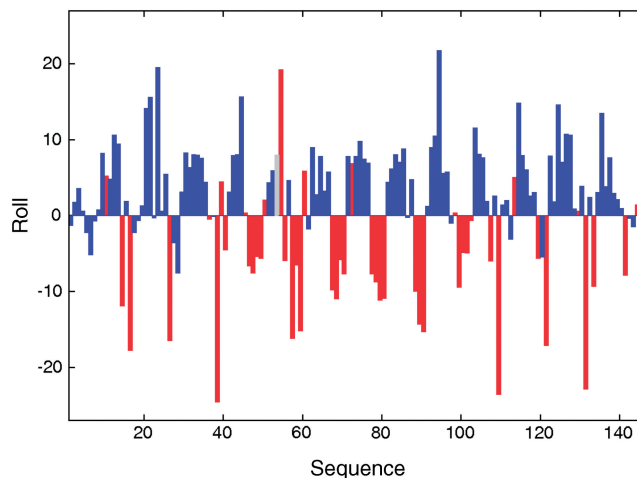


Figure 5. Relationship between roll and backbone conformations in NCP X-ray structures. Roll ($^{\circ}$) profile along the DNA sequence in nucleosome X-ray structure 1KX5. The bars giving the roll values are colored according to the state of the facing phosphates in the complementary dinucleotide steps: blue for BI•BI and red for BI•BII or BII•BII. The grey bar represents an unclassifiable step (BI on strand 1 and $(\epsilon - \zeta) \sim 170^{\circ}$, i.e. neither BI nor BII, on strand 2). Structures 1KX3 and 1M19 give similar patterns.

pools of natural sequences, the nucleosome intrinsic sequence preferences were recently characterized primarily by ~ 10 bp periodicities of specific dinucleotides along the nucleosome length, thought to facilitate the sharp bending of DNA around the nucleosome (56,57,71–78). However, in absence of reliable descriptors of dinucleotide properties, the physical basis of this organization remains obscure.

The NCP X-ray structures revealed a periodic alternation of positive and negative roll tracts along the double helix resulting in the DNA superhelical path (79). Our analysis of several high resolution NCP X-ray structures [PDB codes 1KX3 (59), 1M19 (60) and 1KX5 (59)] shows that positive and negative rolls are mainly associated to BI and BII phosphates, respectively (Figure 5 for 1KX5). Some steps escape this rule, as illustrated by steps 53 and 54 in 1KX5 (Figure 5). Step 53 is associated with an atypical $(\epsilon - \zeta)$ value of $\sim 170^{\circ}$, and in step 54 BI•BII is associated to a strong positive roll. Such peculiar behaviors, together with unusual $\alpha/\beta/\gamma$ angles, help to remember that the backbone conformational landscape of bound DNA is not exactly as in free DNA (19). Nevertheless, rolls and $(\epsilon - \zeta)$ averaged on facing phosphates are as well correlated in 1KX5, 1KX3 and 1M19 (correlation coefficients from -0.72 to -0.65) as in free X-ray DNA (correlation coefficient of -0.67). Thus, the relationship between BI/BII and roll appears robust, as well as those involving twist and X-disp (not shown). So, BI and BII rich regions are the centers of 10 bp fragments curved towards the major and the minor grooves, respectively. Furthermore, the NCP BII steps are in majority encountered on CpA•TpG, GpG•CpC and GpC•GpC (Supplementary Table S3), consistent with the TRX scale. That both BII structural and sequence effects in free DNA are transferable to the

Table 2. Nucleosomal sequences characterized according to the TRX scale

sequences	% _s	140 dinucleotide steps			72 dinucleotide steps			$\Delta\Delta G$	RA	RA
		POW ₁₀	ξ_s	ξ_r	POW ₁₀	ξ_s	ξ_r			
601	46	0.0065	2.2	2.6	0.0064	2.4	2.8	-2.9	1	
618	51	0.0054	1.9	2.0	0.0086	1.7	1.9	-2.3		
f2	58	0.0094	2.9	2.1	0.0102	2.5	2.3		2.47	
f3	55	0.0065	2.4	2.3	0.0102	2.5	2.3		1.60	
h2	47	0.0035	2.0	2.3	0.0029	2.4	2.1		0.11	
h3	58	0.0021	2.7	2.0	0.0001	4.1	1.5		0.15	
g1	45	0.0076	1.8	2.5	0.0082	2.0	2.6			1
g2	39	0.0025	1.8	3.2	0.0007	2.1	4.4			0.16
g3	40	0.0031	1.6	2.7	0.0027	1.5	2.8			0.15
g4	36	0.0054	1.5	2.7	0.0047	1.4	2.8			0.14
g5	37	0.0020	1.4	3.1	0.0005	1.1	3.7			0.12

The 147 bp sequences (Supplementary Table S4), their nomenclature and the relative affinities (RA) are from Dr Segal's web site <http://genie.weizmann.ac.il/pubs/nucleosomes06/index.html>. The relative affinities are referenced to either 601 (values in columns 9 and 10) or g1 (values in last column) sequences. The central regions of f2 and f3 are identical. The $\Delta\Delta G$ (kcal/mol) values are referenced to that measured for the 5S sequence (55). The percentage of stiff complementary dinucleotides (%_s) is given for each sequence. FT analyses were performed considering the TRX scores as a signal along either the 140 innermost or the 72 central dinucleotide steps of the DNA sequences. POW₁₀ is the integrated power of the FT peak at 0.1 ± 0.01 frequency (periodicity of 10 ± 1 bp). The characteristic lengths were calculated for the stiff (ξ_s) and flexible (ξ_r) blocks defined by their TRX scores (Table 1).

NCP structures makes this system particularly relevant to analyze how DNA intrinsic flexibility, quantified with the TRX scale, influences its assembly with proteins.

We analyze here the structural properties of synthetic sequences related to the SELEX approach for which affinity measurements are available (Table 2 and Supplementary Table S4). Artificial and natural sequences exhibit the same lexicographic periodicities: ApA•TpT, TpA and ApT oscillate out of phase with GpG•CpC, CpG•CpG and GpG•CpC, with a period of ~ 10 bp (56). The related SELEX pool contains very high affinity sequences that do not have their equivalent in genomic positioning DNA. We wondered if the characteristics of these very high affinity sequences would emerge particularly clearly from a TRX analysis and contrast with those of lower affinity sequences. So, we first considered the SELEX sequence best at forming NCP (sequence 601), and one of the worst (sequence 618) (57). These sequences 601 and 618 present a weak similarity, with $\sim 30\%$ of identical bases. A set of sequences deriving from sequence 601 were also analyzed (<http://genie.weizmann.ac.il/pubs/nucleosomes06/index.html>). f2 and f3 correspond to sequence 601 enriched in 10 bp periodic TpA steps either along the whole sequence (f2) or only in its central part (f3). Thus the central regions of f2 and f3 are identical; their affinities for the histone core are higher than that of sequence 601. h2–3 and g1–5 differ from sequence 601 by ~ 50 mutated bases between positions 31 and 122, i.e. in the central region of the 147 bp segment. Apart from g1, these sequences display low propensities to form nucleosomes. Overall, a representative range of affinities is covered by these 11 sequences.

Considering the TRX scores as signals along the DNA sequence, Fourier transform (FT) analyses were performed on i) the 140 dinucleotides (141 bp) corresponding to the DNA length in an NCP less its outermost base pairs and ii) the central 72 dinucleotides (73 bp), given that most of the considered sequences differ from the 601 reference

by mutations in this region. Regarding the NCP, the most interesting spectrum section covers the frequencies from 0.09 to 0.11, corresponding to periodicities of ~ 11 and 9 bp, respectively. Indeed, the NCP sequence periodicities inferred from various biochemical experiments vary within this range [(80) and references herein]. The direct estimates of periodicity offered by high-resolution crystal structures of NCP reconstituted on a unique DNA sequence give periodicity values from 10.15 to 10.3 bp (59,79).

Sequences 601, f2, f3 and g1 are characterized by a strong FT signal corresponding to a ~ 10 bp periodicity (see Figure 6 for 601 and f2). The corresponding powers, i.e. integrated the FT signal between frequencies 0.09 and 0.11, show maximal scores (Table 2) that are further strengthened in the central regions for f2, f3 and g1 (Table 2). This means that these preferential DNA sites for assembling into nucleosome are characterized by ~ 10 bp alternations of blocks of enhanced and restricted flexibility. Considering that enhanced and restricted flexibilities correspond to TRX scores higher and lower than the average, respectively, we find that the average lengths of each type of block are globally equivalent (characteristic lengths in Table 2), especially in the central region (Table 2 and Figure 7). This organization, illustrated in Schema 1 for the sequences f2, 601 and g1, correlates the X-ray NCP structures in which the BI or BII tracts include at least three consecutive dinucleotides (Figure 5).

Incidentally, considering TpA as flexible as CpA•TpG, as indicated by the X-ray data, results in a net decrease of all the ~ 10 bp FT signals. For instance, the power of the 10 bp FT peak of f2 drops from 0.0094 (TpA stiff) to 0.003 (TpA flexible). When assuming a flexible TpA, the FT analysis detects a strong 15 bp periodicity on these high affinity sequences, not compatible with the NCP. This reinforces the notion developed above that the flexibility of TpA estimated from X-ray datasets could be inappropriate to DNA in solution.

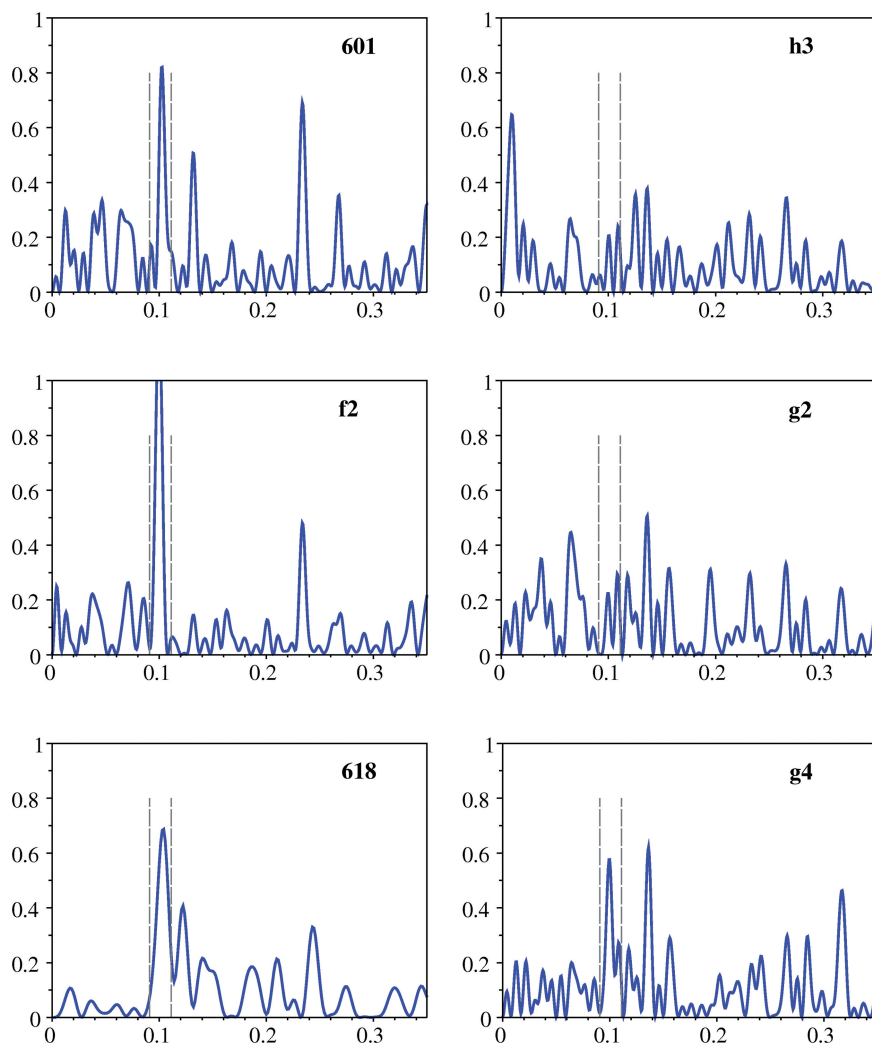


Figure 6. Fourier analysis of the TRX signal along DNA nucleosomal sequences. Fourier transform (intensity in arbitrary units) as a function of wave vector k (bp^{-1}) of the TRX signal along sequences representative of high (601 and f2), moderate (618) and low (h3, g2 and g4) propensity for forming nucleosome (Table 2). The Fourier analysis was performed considering the TRX scores (Table 1) as a signal along the innermost 140 dinucleotides of the nucleosomal sequences, apart from sequence 618, analyzed on the 72 central dinucleotides. The dotted vertical lines indicate the window corresponding to the period 10 ± 1 bp.

Sequence 618, less attractive than 601 for the histone core, gives a less intense TRX-FT signal at frequency of 0.1 ± 0.01 when the entire sequence is taken into account, but the 10 bp periodicity is noticeably enhanced in its central part (Figure 6 and Table 2). Nevertheless, the block lengths are globally shorter than those of very high affinity sequences (Table 2, Figure 7 and Scheme 1), implying that the number of steps pushed at the inverse of their intrinsic structural trends in NCP is greater with 618 than with 601.

The ~ 10 bp periodicity is even more subdued for the low affinity sequences h2, h3 and g2–5 (Figure 6 for h3, g2 and g4). FT peaks at 0.1 ± 0.01 are still observable but their intensities are very low (Table 2). In addition, the average lengths of stretches of restricted and enhanced flexibility are not balanced (Table 2). Either stiff (h3) or flexible (g2–5) dinucleotides are dominant in the central region of these low affinity sequences (Table 2, Figure 7

and Scheme 1 for h3, g2 and g4). Such configuration is likely to increase the energetic penalty in NCP formation, with a large deformation cost (excess of stiff dinucleotides) or a loss of entropy (excess of flexible dinucleotides). This idea is supported by the very poor affinity of sequence g4 that presents a significant 10 bp periodicity (Figure 6, Table 2) but with almost all its central blocks of restricted flexibility corresponding to only one dinucleotide (Figure 7, Table 2, Scheme 1).

In sum, the TRX scale reveals that the preferential DNA sites for assembling into nucleosome are primarily characterized by a 10 bp periodicity that corresponds to an alternation of blocks characterized by restricted and enhanced flexibility. Our analysis also suggests that blocks of minimal length are required to define a high affinity sequence. The central 72 dinucleotides may be more crucial for the positioning than their flanking external parts since both the periodicity and the block

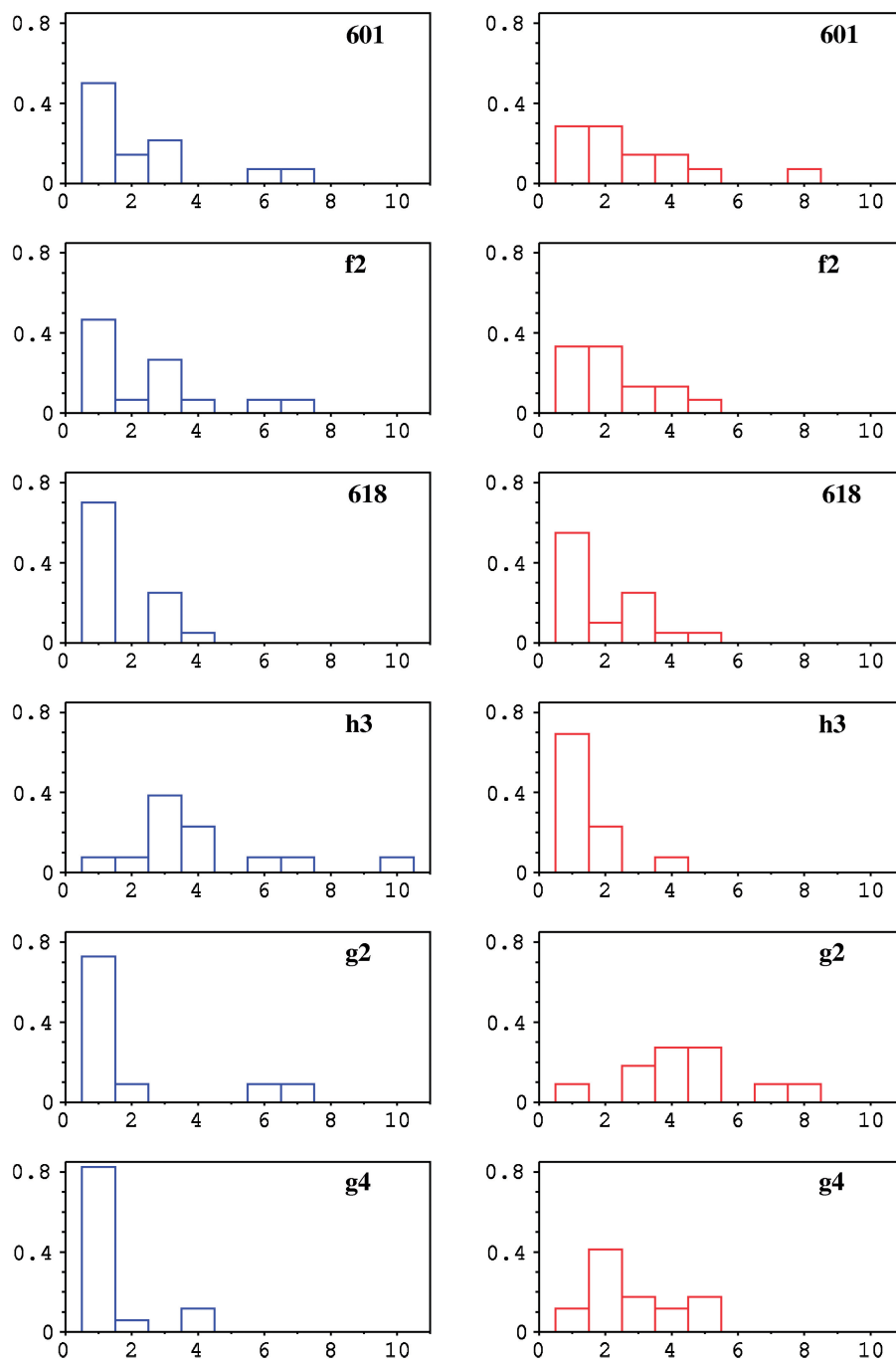


Figure 7. Histograms of the length (expressed in number of dinucleotides) of stiff (left panels, in blue) and flexible (right panels, in red) blocks in the central 72 dinucleotides of the nucleosomal DNA sequences. The sequences presented here are the same as in Figure 6, with nucleosomal affinity ranging from very high (f2) to low (h3, g2, g4).

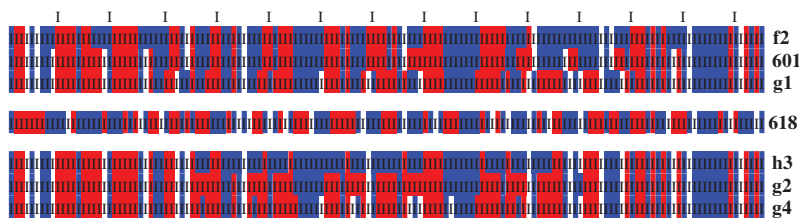
length balance found for the best assembling sequences are especially marked in the central parts of the 147 bp fragments analyzed here.

DISCUSSION AND CONCLUSION

In B-DNA, any phosphate group is potentially able to oscillate between the BI and BII states, the BI/BII ratio reflecting the helicoidal twist, roll and X-disp values of the

corresponding step (19,23,46–52). The BI/BII ratio is accessible by the solution NMR measurements of ^{31}P chemical shifts (δP) (48).

The present work establishes firmly the dinucleotide sequence effect on the BI \leftrightarrow BII equilibrium and its populations, based on a new and extensive compilation of the available δP measured on free B-DNA in solution. The influence reflecting the category (purine or pyrimidine) of the nearest neighbors bracketing the dinucleotide is detected but this secondary effect does



The sequences represented here are representative of high (f2, 601 and g1), moderate (618) and low (h3, g2 and g4) propensity for nucleosome formation. Each sequence contains 146 phosphodiester linkages scanned in terms of stiff (blue, TRX scores higher than average, see Table 1) or flexible (red, TRX scores lower than average) dinucleotides (vertical bar). The semi-flexible GpA•TpC step is not colored. The first line of black bars marks the dimeric steps at position 10, 20, 30 ... bp from the left end.

Scheme 1. Nucleosomal sequences coded by the TRX flexibility scale.

not disrupt the flexibility rankings detected at the dinucleotide level. Due to the predominant impact of the dinucleotide sequence, each complementary dinucleotide step can be characterized by a number corresponding to the experimental average BII population of its facing phosphates (Table 1) and thus the phosphate group flexibility. Crucially, this number also represents the malleability in terms of Twist, Roll and X-disp (base pair displacement), due to the tight coupling between helical parameters and backbone states. Thus, the numbers associated to the ten complementary dinucleotide steps result in a scale of intrinsic flexibility that we call 'TRX'. The most inherently flexible and rigid steps correspond to TRX scores of 50 (50% BI/50% BII) and 0 (100% BI), respectively.

Broadly speaking, two dinucleotide categories emerge from the TRX scale. The first has restricted flexibility, i.e. low or moderate twists associated with positive and null rolls. Their corresponding bases are preferentially located close to the axis of the double helix. The second group exhibits an enhanced flexibility where the dinucleotides can adopt either low twists/positive rolls or high twists/negative rolls. Such bases are more easily displaced towards the major groove, away from the helical axis. According to the TRX scale, ApA•TpT, ApT•ApT, TpA•TpA have restricted flexibility. In contrast, GpG•CpC, GpC•GpC, CpG•CpG have enhanced flexibility. The steps which are mixed in terms of base composition have flexibilities either restricted (ApC•GpT and ApG•CpT) or enhanced (CpA•TpG), GpA•TpC being intermediate. Thus, the DNA dynamical properties are not easily assigned solely by reading the lexicographic nucleotide chaining.

Our approach of flexibility may go beyond the intrinsic plasticity of DNA in its B form, since the contrasted behavior of GpG•CpC, GpC•GpC, CpG•CpG on one hand, and ApA•TpT, ApT•ApT, TpA•TpA on the other hand, according to δP echoes the influence of sequences on double helix polymorphism. Indeed, contrary to A•T rich sequences, G•C and CpA•TpG rich DNAs can adopt the Z form (81–83) and facilitate the B to A transition (84–86). The A and Z forms correspond to large molecular reorganizations occurring generally in non-physiological environments, but their formation may take advantage of the dinucleotide properties highlighted here.

Most importantly, the TRX scale emerges as useful to understand how the DNA/protein interactions depend on DNA sequence-dependent intrinsic malleability. Here, the TRX scale was used to investigate the properties of nucleosomal positioning sequences. Indeed, it is established that the nucleosome reconstitutes better on certain DNA sequences than others (56,77,87). Our analysis of the high resolution NCP crystal structures reveals that the tight coupling between BI and BII backbone states and helical parameters, together with the marked sequence dependency of the BI/BII propensities, echoes our findings on free DNA. This supports the idea that intrinsic DNA properties reduce the cost of marked distortions required by wrapping in the NCP complex (56,77,87).

Our TRX analysis was performed on artificial sequences that offered a range from very high to low affinity for nucleosome reconstitution. We show that the high affinity sequences are characterized by a ~ 10 bp periodicity, corresponding to a regular alternation of stiff (limited to positive and null rolls) and flexible stretches (able to adopt negative rolls). Thus, high affinity sequences consist of a succession of blocks phased to distort as required in the NCP structure. This periodicity is maximal when considering the central 73 bp. It appears also that, in these 73 bp central parts, a minimal length of both stiff and flexible blocks is required to obtain optimal nucleosomal sequences.

These findings are consistent with several observations. First, our analysis explains why sequences considerably enriched in A•T, unable to adopt negative rolls without conformational strain, function as nucleosome exclusion signals (70,71,88,89), likely due to a large deformation energy penalty. Also, it suggests that too much DNA flexibility (e.g. with a flexible dinucleotide at every step) may lower the DNA/protein affinity because of a loss of entropy when going from the flexible free form to the nucleosomal structure (90,91). Second, the importance of the sequence central part is consistent with (i) the consensus found in the central part of artificial nucleosomal sequences (56); (ii) the nucleosome reconstitution *in vivo* and *in vitro* that begins with the central (H3–H4)₂ tetrasomal part, followed by incorporation of the two terminal (H2A–H2B) dimers (92,93) and (iii) a recent study showing that 73 bp fragments are efficient to analyze the nucleosome occupancy along the genomes of

Saccharomyces cerevisiae and *Drosophila melanogaster* (94). Third, the TRX scale allows to rationalize the dinucleotide position frequencies found on both artificial (56,57,87) or natural nucleosomal DNA sequences *in vitro* (72,75–78) or *in vivo* (95). These studies converge to characterize the nucleosome sequence preferences primarily by a ~10 bp periodicity of ApT, TpA and ApA•TpT that oscillate out of phase with GpC, CpG, GpG•CpC and CpA•TpG along the nucleosomal DNA constituent. Both groups are each composed by a mixture of YpR, RpR•YpY and RpY. Neither the modeling nor the crystallographic approaches do properly account for DNA plasticity since they all conclude to flexible YpR, intermediate RpR•YpY, and stiff RpY steps. From the TRX point of view, ApT, TpA and ApA•TpT are characterized by a restricted plasticity, contrary to the flexible GpC, CpG, GpG•CpC and CpA•TpG. Thus, the TRX approach clarifies the lexicographic periodicities encountered in nucleosomal sequences, artificial and natural.

In sum, examination of free DNA backbone motions in solution, monitored by δP , correlated to several helical parameters, allows to uncover the intrinsic B-DNA flexibility. This flexibility appears dominated by the dinucleotide sequence, although finely modulated at the tetranucleotide level. The quantification of the DNA intrinsic malleability of the ten complementary dinucleotides yields an experimental scale, called TRX (Table 1). We started to decipher the affinity of artificial DNA sequences for nucleosome formation in terms of their TRX signals. This shows that the TRX scale offers an understanding of the preferred recognition of DNA sequences by proteins, based on flexibility criteria.

A key strength of the TRX scale is its experimental foundation, directly inferred from very precise δP NMR measurements in solution, free from dependence on any structural models or limitations (e.g. force-field influences or crystal packing). To our knowledge, the TRX scale is currently the only experimental quantitative scale characterizing the ten B-DNA complementary dinucleotide steps in terms of average structure and flexibility, thus offering a conceptual and practical framework to make progress regarding the role of DNA dynamics in DNA–protein recognition mechanisms.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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