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# Computational analysis suggests a highly bendable, fragile structure for nucleosomal DNA

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#### ABSTRACT

Eukaryotic chromosomal DNA coils around histones to form nucleosomes. Although histone affinity for DNA depends on DNA sequence patterns, how nucleosome positioning is determined by them remains unknown. Here, we show relationships between nucleosome positioning and two structural characteristics of DNA conferred by DNA sequence. Analysis of bendability and hydroxyl radical cleavage intensity of nucleosomal DNA sequences indicated that nucleosomal DNA is bendable and fragile and that nucleosome positional stability was correlated with characteristics of DNA. This result explains how histone positioning is partially determined by nucleosomal DNA structure, illuminating the optimization of chromosomal DNA packaging that controls cellular dynamics.

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# 1. Introduction

Eukaryotic chromosomal DNA is packaged into nucleosomes and then into chromatin, which has a repetitive, compact structure. A nucleosome comprises about 147 base pairs (bps) coiled around a histone octamer, which consists of two copies of four core histones (H2A, H2B, H3, and H4) (Luger et al., 1997; Davey and Richmond, 2002). Neighboring nucleosomes are separated by linker regions of DNA (approximately 50 bps) without histones (Van Holde, 1989).

Previous studies have shown that nucleosomes have various functions besides the packaging of chromosomal DNA. For example, many cellular processes that involve DNA respond dramatically to changes in the positions of and modifications of nucleosomes, and in the nucleus, the dynamics of the positioning of nucleosomes and their flanking regions play a key role in cellular processes (Strahl and Allis, 2000; Yuan et al., 2005; Segal et al., 2006; Barski et al., 2007; Schones et al., 2008; Lister et al., 2009; Tolstorukov et al., 2009). For instance, nucleosome positioning regulates eukaryotic transcription by affecting the accessibility of transcription factors to DNA or by affecting

Abbreviations: ChIP-chip, ChIP combined with microarray technology; ChIP-Seq, ChIP combined with deep sequencing; GC, guanine-cytosine; PRS, positive reference set; RRS, random reference set.

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transcriptional elongation by polymerase (Yuan et al., 2005; Schones et al., 2008; Cairns, 2009).

The efficiency of the coiling of DNA around histones to form nucleosomes depends strongly on DNA sequence (Yuan et al., 2005; Ogawa et al., 2010). *In vitro* studies have shown that the range of affinities between histones and DNA is at least 1000-fold and that DNA sequence pattern is an important factor for accurate nucleosome positioning (Thastrom et al., 1999). Furthermore, the results of several other studies suggest that the DNA coiled around the histones is bendable, and these studies have shed light on the importance of DNA structure for nucleosome positioning (Drew and Travers, 1985; Balasubramanian et al., 1998; Johnson et al., 2006; Miele and Vaillant, 2008; Rohs et al., 2009). However, no detailed, systematic analysis of the bendability of coiled DNA has been performed, and whether DNA bendability is chromatin wide is not clear.

Along with the progress in computational methods for analyzing nucleosome data (Zhang et al., 2008; Tolstorukov et al., 2009), recent remarkable progress in chromatin immunoprecipitation (ChIP) techniques, including ChIP-chip (ChIP combined with microarray technology) and ChIP-Seq (ChIP combined with deep sequencing), has permitted detection and prediction of nucleosome positioning at the whole-genome level (Yuan et al., 2005; Barski et al., 2007; Park, 2009). The increase in the number of large-scale nucleosome data sets has permitted systematic analysis of nucleosome data in conjunction with other "omic" data (e.g., genome-wide relationships between nucleosome positioning and transcription regulation (Schones et al., 2008; Lister et al., 2009)).

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In this study, we used a computational analysis of a large-scale nucleosomal data set to evaluate two overall characteristics of chromosomal DNA structure: DNA bendability and DNA cleavage intensity. DNA bendability is calculated from parameters for a set of trinucleotide patterns (Brukner et al., 1995) or tetranucleotide patterns (Packer et al., 2000) in a given sequence. The parameters for each trinucleotide pattern were obtained by Brukner et al. (1995) using DNA binding proteins which generally bind to DNA that bends toward the major groove. The parameters were derived from 709 DNA binding regions. Bendability is frequently used to analyze DNA structure. For example, the location of transcription initiation promoters is associated with areas of bendable DNA (Abeel et al., 2008; Akan and Deloukas, 2008; Choi and Kim, 2009).

Cleavage intensity indicates the likelihood of DNA cleavage by hydroxyl radicals and provides a map of local variation in the shape of the DNA backbone. Cleavage intensity is calculated from parameters for a set of tetranucleotide patterns in a given sequence. The parameters were derived by Greenbaum et al. (2007) from experiments in which DNA sequences were exposed to hydroxyl radicals; they analyzed how cleavage of DNA sequences depends on their sequence patterns. Cleavage intensity also reflects the solvent-accessible surface area of DNA. This index of DNA fragility has also been used to characterize structural features of DNA (Tullius and Greenbaum, 2005; Greenbaum et al., 2007).

These two structural parameters cover all combinations of trinucleotide or tetranucleotide patterns and are sufficiently reliable to analyze DNA structure (Brukner et al., 1995; Greenbaum et al., 2007). We conducted genome-wide computational analysis of the DNA bendability and cleavage intensity of nucleosome and linker DNA regions and found that nucleosomal DNA was overall highly bendable and fragile. We also determined whether the positional stability of nucleosomes under various cellular conditions depended on these two structural properties of DNA, and we found that stable nucleosomal DNA was more bendable and cleavable than unstable nucleosomal DNA. We discuss which types of DNA structure confer the optimal DNA packaging in chromosomes and which types are optional for dynamic cellular processes.

# 2. Materials and methods

# 2.1. Genomic DNA and nucleosome positioning data

The reference genome sequence of Homo sapiens (hg18) was obtained from the University of California at Santa Cruz genome database (http://genome.ucsc.edu/). To determine genomic histone positions, we used the previously reported ChIP-Seq data set for histone H3 of human CD4+ T-cells obtained under two cellular conditions, with and without the T-cell receptor signal, which are designated as the activated and resting conditions, respectively (Schones et al., 2008). We modified a previously developed method (Tolstorukov et al., 2009) to define nucleosomal positions from the ChIP-Seq of both resting and activated cells (for details of the method, see Supplementary methods). We assessed the positional stability of nucleosomes under the two cellular conditions by computing the distance of the genomic position of a given nucleosome in the resting cell from its nearest nucleosomal position in the activated cell. For all the nucleosomes identified under the resting conditions, we defined stable nucleosomes as those for which the distance between the nucleosome under resting conditions and the nearest nucleosome under activated conditions was within 12 bps, and unstable nucleosomes as those for which that distance ranged from 13 to 73 bps (half the size of one nucleosomal DNA unit). Nucleosomes that did not fit either definition were not used for the positional stability analysis. The boundary between the two types of nucleosomes (i.e., the 12-13-bp boundary) was defined from the inflection point in the distance distribution (Supplementary methods).

#### 2.2. DNA bendability calculation

To assess nucleosomal DNA bendability, we used the trinucleotide bendability parameter profile obtained by Brukner et al. (1995). For each of the nucleosomes identified under the resting conditions (including both stable and unstable nucleosomes), DNA bendability was calculated for every trinucleotide in the region from -400 to +400 bps from the center position of the nucleosomal DNA region. For each position relative to the central position of the nucleosome, the average DNA bendability was calculated.

#### 2.3. DNA cleavage intensity calculation

We used the OH radical cleavage intensity database (ORChID; http://dna.bu.edu/orchid/) program, which is used to estimate the hydroxyl radical cleavage strength and solvent-accessible surface area using a tetranucleotide DNA cleavage intensity profile (Greenbaum et al., 2007), to calculate the DNA cleavage intensity. (The source code of ORChID was kindly provided by its authors; ORChID is a web-based application.) For each of all the nucleosomes identified under the resting conditions, we computed tetranucleotide DNA cleavage intensities for all the DNA positions from -400 to +400 bps from the center position by querying to ORChID. For each position relative to the central position of the nucleosome, the average DNA cleavage intensity was calculated.

## 2.4. Guanine-cytosine content and statistical analysis

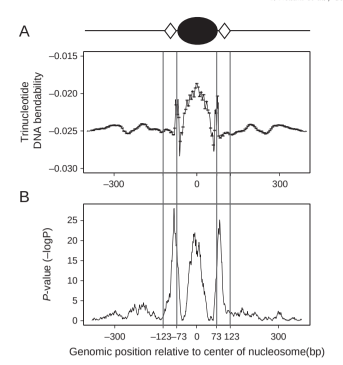
For a control analysis with respect to guanine-cytosine (GC) content, we randomly sampled 10000 out of the ~5000000 nucleosomes identified under the resting conditions as a positive reference set (PRS) to reduce the calculation time. For each nucleosome in the PRS, sliding window averages of DNA bendability, cleavage intensity, and GC content for the region surrounding the nucleosome (from -400 to +400 bps) were calculated using a 30-bp window with 1-bp displacement. To prepare a random reference set (RRS), we randomly selected 1000000 genomic positions and used their surrounding sequences (from -400 to +400 bps) to calculate DNA bendability, cleavage intensity, and GC content using a 30-bp sliding window average. Additionally, the DNA bendability and cleavage intensity expected from each GC content value were calculated from the above data, and we set them as RRS with GC content. Then, we randomly correlated the sliding window for each of the 10000 nucleosomes in the PRS with the sliding windows of the RRS having the same GC content, so that the DNA bendabilities and cleavage intensities of the same relative sliding window positions could be compared with the DNA bendabilities and cleavage intensities expected from the GC content and the genomic context. For each sliding window position relative to the center of the nucleosomal region, differences in DNA bendability and cleavage intensity between the PRS and the RRS were evaluated by t-test.

#### 3. Results

### 3.1. Bendability of nucleosomal DNA

To investigate the structural properties of nucleosomal DNA, we first analyzed the bendability of the DNA sequences surrounding the nucleosomal region. Using all the nucleosomal positions of human CD4+ T-cells identified under the resting conditions, we calculated average DNA bendabilities of the genomic positions relative to the central position of the nucleosome (from -400 to +400 bps) (Fig. 1A). Bendability scores within the nucleosomal region were significantly higher than those within the linker regions ( $P<10^{-15}$ ; t-test). Moreover, the bendabilities in the nucleosomal region gradually increased toward the central position of the nucleosome,

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**Fig. 1.** Bendability of nucleosomal DNA and the surrounding regions. (A) Trinucleotide DNA bendability versus genomic position relative to the central position of the nucleosome. The black oval denotes the nucleosome region (from -73 to +73 bps), and the white diamonds denote linker regions (from -123 to -74 bps and from +74 to +123 bps). The polygonal line and the error bars represent average trinucleotide bendabilities and standard deviations (s.d.) (Note that s.d. values are shown only for every 5th bp. to improve viewability.) (B) Statistical significance (P-values) of differences in DNA bendability between PRS and RRS as determined by t-test (see Materials and methods). The polygonal line represents P-values for each genomic position.

which is consistent with the symmetric structure of the histone protein (Luger et al., 1997). The bendability distribution showed a ~10-bp periodic fluctuation within the nucleosomal DNA region. Even more striking, DNA bendabilities at the boundaries between the nucleosomal regions and linker regions on both sides of the nucleosome were quite high, which implies that the unbending boundary regions also played an important structural role in nucleosome positioning.

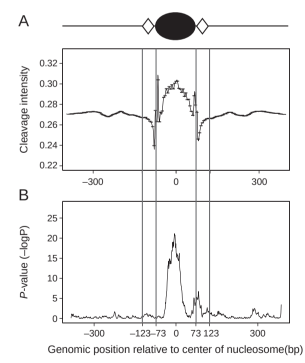
Previous work has suggested that DNA bendability is affected by GC content (Peckham et al., 2007), and several machine learning approaches have shown that nucleosome positioning is associated with DNA sequence pattern (Peckham et al., 2007; Ogawa et al., 2010). Therefore, we performed a control analysis of GC content to determine whether the high bendability of the nucleosomal DNA was a consequence of high GC content that is possibly correlated with the nucleosomal DNA sequence context. For each position relative to the central position of the nucleosome, DNA bendabilities were compared with those expected from the RRS (see Materials and methods). P-value peaks ( $P < 10^{-20}$ ; t-test) were observed at three positions: the central position of the nucleosome and at both boundaries between the nucleosome and the linker regions (Fig. 1B). In contrast, the other regions did not show statistical significance for DNA bendability even within nucleosomal region, meaning that the DNA bendabilities of these regions could be expected from GC content alone. The central position of nucleosomal DNA and the nucleosome-linker junction were suggested to have high bendabilities that could not be explained by GC content alone.

## 3.2. Fragility of nucleosomal DNA

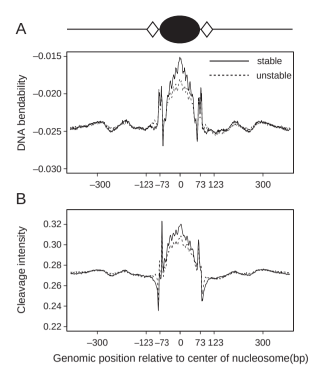
We next computed DNA cleavage intensity, which is an estimate of hydroxyl radical cleavage strength and the solvent-accessible area, for the DNA sequence surrounding the nucleosome using ORChID. Like the DNA bendability pattern, the DNA cleavage intensity pattern for the nucleosomal DNA region was symmetrical. The cleavage intensity in the nucleosomal region was higher than that of the linker regions on both sides ( $P < 10^{-15}$ ; t-test), and a  $\sim 10$ -bp periodic fluctuation of DNA cleavage intensity was also observed, especially in the nucleosomal DNA region (Fig. 2A). The cleavage intensity of the nucleosomal region was statistically higher than the intensities of the RRS expected from the GC content, especially at the central position of the nucleosome (Fig. 2B). Moreover, interestingly, the cleavage intensities in the regions immediately inside the nucleosome–linker junctions were markedly higher than average, whereas the intensities in the regions immediately outside the junctions were markedly lower than average (that is, the distribution dropped sharply from the inside to the outside of the nucleosome region). However, the difference from the RRS was statistically significant only for the downstream junction ( $P < 10^{-5}$ ).

# 3.3. Structural features of nucleosomal DNA and positional stability of the nucleosome

If nucleosomal DNA structure determines nucleosome positioning, the positional stability of the nucleosome should correlate with structural characteristics of nucleosomal DNA. To determine whether this correlation was in operation, we compared nucleosome positions obtained from the resting state of CD4+ T-cells with the positions obtained from the activated state, and we then defined stable and unstable nucleosomes (see Materials and methods). For each group of stable and unstable nucleosomes, we analyzed DNA bendability and cleavage intensities of the region surrounding the nucleosome (from  $-400\ {\rm to}\ +400\ {\rm bps})$  and found that the overall nucleosomal DNA bendability and the overall cleavage intensity of stable nucleosomes were markedly higher than those of unstable nucleosomes (Fig. 3).



**Fig. 2.** Cleavage intensity of nucleosomal DNA and the surrounding regions. (A) DNA cleavage intensity. The black oval denotes the nucleosome region (from -73 to +73 bps), and the white diamonds denote linker regions (from -123 to -74 bps and from +74 to +123 bps). The polygonal line and the error bars represent average DNA cleavage intensity and s.d. (Note that s.d. values are shown only for every 5th bp, to improve viewability.) (B) Statistical significance (P-values) of differences in cleavage intensity between PRS and RRS as determined by t-test (see Materials and methods). The polygonal line represents P-values for each genomic position.



**Fig. 3.** Structural characteristics of DNA around stable and unstable nucleosomes. (A) DNA bendability and (B) DNA cleavage intensity. The black oval denotes the nucleosome region (from -73 to +73 bps), and the white diamonds denote linker regions (from -123 to -74 bps and from +74 to +123 bps).

#### 4. Discussion

In this study, we computationally analyzed two structural features of the DNA sequences around nucleosomes and found several significant differences between nucleosomal and linker DNA regions. The sequence around the nucleosome overall had a higher DNA bendability score and a higher DNA cleavage intensity score than did the linker regions. These observations were true overall, but not every local region of DNA showed exactly the same tendencies. However, we found many local regions that did show such tendencies (an example is presented in Supplementary Figure 2). We note that our preliminary results indicated that these structural features of DNA sequences associated with nucleosomal and linker regions were also found in *Arabidopsis thaliana* (Supplementary Figure 3), which implies that these structural features may be conserved in other species.

Previous studies have reported the existence of specific DNA sequence patterns in the nucleosomal DNA region (Segal et al., 2006; Peckham et al., 2007; Ogawa et al., 2010). For instance, a previous support vector machine-based learning approach was used to derive short DNA fragments that frequently appear in the nucleosomal region (Ogawa et al., 2010). However, the correlation between these short DNA fragments and nucleosome formation and DNA bendability scores is far from perfect. Although the trinucleotides CCA/TGG and CAG/CTG have been reported to have the highest nucleosomeformation potentials (Peckham et al., 2007; Ogawa et al., 2010), the bendability scores of CCA/TGG and CAG/CTG, -0.246 and 0.175, respectively, are lower and higher, respectively, than the average bendability score of -0.019. ATA/TAT, TAA/TTA, and AAA/TTT have been suggested as having the potential to inhibit nucleosome formation (neither of the papers offered any reasons for this low frequency) (Peckham et al., 2007; Ogawa et al., 2010), but their bendability scores (0.182, 0.068, and -0.274, respectively) are higher, higher, and lower, respectively, than the average. Furthermore, although a previous study suggested that DNA bendability is strongly correlated with GC content (Peckham et al., 2007), we showed that bendable DNA structures around the center positions of nucleosomes and in the regions around the nucleosome-linker junctions cannot be explained by GC content alone. Accordingly, the high bendability score of the nucleosomal DNA region is not merely a consequence of coincidence between DNA bendability parameters and previously known sequence features of nucleosomal DNA, and this fact supports the plausibility that nucleosomal DNA is bendable.

We also found that high DNA cleavage intensity scores, especially at the central positions of the nucleosomes, were independent of GC content. We also calculated average DNA cleavage intensities of tetranucleotides containing CCA/TGG and CAG/CTG trinucleotides. Although CCA/TGG and CAG/CTG were the two sequence fragments with the highest potentials to form nucleosomes (Ogawa et al., 2010) (as mentioned previously), the average DNA cleavage intensities of tetranucleotides containing CCA/TGG and CAG/CTG were 0.37 and 0.55, respectively, that is, the same as and higher than the average tetranucleotide cleavage intensity (0.37). This result implies that, like DNA bendability, the high DNA cleavage intensity score of the nucleosomal DNA region is not due completely to the known sequence features enriched within the nucleosomal DNA region.

A recent study has suggested a relationship between the bendabilities and cleavage intensities of DNA sequences: these features commonly exist within positions of the minor groove of the DNA structure (Heddi et al., 2009). Our results indicate a similarity between the distribution patterns of DNA bendability and cleavage intensity with respect to genomic position relative to the central position of the nucleosome, and both distributions have ~10-bp periodic fluctuation, which is consistent with length of one pitch of the double-stranded DNA structure (i.e., the periodicity of the minor groove positions that face the histone). To determine whether the similar distribution patterns were independent from or tightly associated with each other, we compared trinucleotide bendability scores and DNA cleavage intensity scores of tetranucleotides. For each of the trinucleotides having DNA bendability scores, we calculated the average DNA cleavage intensity of all tetranucleotides containing the corresponding trinucleotide in the nucleosomal DNA region. We found that the Pearson's correlation coefficient between the trinucleotide bendability scores and their corresponding average tetranucleotide cleavage intensity scores was 0.26, which suggests no direct relationship between bendability and cleavage intensity.

Taken together, our results suggest that the high bendability scores of the nucleosomal DNA region computed in this study and the symmetric bendability pattern around the nucleosomes reflect the fact that, regularly repeated, bendable DNA regions have evolved so that chromosomal DNA can sinuously coil around histones and form the compact chromatin structure. Moreover, the high cleavage intensity of the nucleosomal region suggests that histones confer durability to nucleosomal DNA against hydroxyl radicals and that this DNA region has not needed to gain the durability evolutionarily. Abrupt changes in DNA bendability and cleavage intensity were found at the nucleosome-linker junctions. In particular, the DNA bendability of the junction region significantly exceeded that expected from the GC content, which suggests that the DNA at nucleosome-linker junctions plays an important structural role in the stability of nucleosome positioning. The overall trends in DNA bendability and cleavage intensity for the stable nucleosomes were more remarkable than those for the unstable nucleosomes; this result supports our speculations about the biological functions of bendability and cleavage intensity. Gene expression is thought to be controlled by nucleosomal positioning and stability (Cairns, 2009). The relationship between the positional stability of nucleosomes and DNA structural preferences also suggests that nucleosomal DNA sequences are encoded to control gene expression. Although known DNA sequence features that are suggested to facilitate high-efficiency formation of nucleosomes and DNA sequences conferring DNA bendability and cleavage intensity must not be independent from each other, we believe that our results shed light on how nucleosome positioning's control of cellular dynamics (e.g., gene expression pattern (Schones et al., 2008; Cairns, 2009)) is regulated by DNA structural preferences.

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# Appendix A. Supplementary data

Supplementary data to this article can be found online at doi:10.1016/j.gene.2011.02.004.

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