Sequence-dependent Kinetic Behavior of Protein-induced B- to Z-DNA Transition

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Z-DNA is a left-handed duplex DNA that contains a zigzag backbone. However, Z-DNA is merely not a mirror-image of the right-handed B-DNA. Since the first crystal structure of Z-DNA was revealed,\(^1\) there have been many studies about chemical and physical characteristics of Z-DNA. It is well known that Z-DNA formation has a preference for sequences with alternating pyrimidine/purine (APP), especially alternating dC and dG.\(^2\) Poly d(GC) can be stabilized in Z conformation in the presence of salts such as 4 M NaCl.\(^3\) In addition, positively charged molecules such as spermine and cobalt hexaamine can facilitate B-Z transition of double stranded DNA (dsDNA) into Z conformation.\(^2\) Chemical modifications on bases also stabilize dsDNA in Z conformation; for instance, poly d(G\(^{32}\)C) requires much lower salt concentrations to form Z-DNA than those used for poly d(GC).\(^2\) In vivo, it is thought that negative supercoiling occurring from various biological processes such as RNA transcription is a major driving force to stabilize Z-DNA.\(^4\) Recently, a novel Z-DNA binding motif Z.\(\alpha\) was discovered by Rich group.\(^5\) The Z.\(\alpha\) was first identified from human ADAR1 (double-stranded RNA adenosine deaminase 1);\(^6\) an RNA editing enzyme that is responsible for A-to-I RNA editing by deaminating adenosine. Thus, human Z.\(\alpha\) from ADAR1 (hZ.\(\alpha\)ADAR1) has been a subject of works about both its Z-DNA binding activity and its use for Z-DNA study.\(^6\)

It is suggested that Z-DNA might have a functional role in transcription from the work of Ho and his colleagues,\(^7\) in which Z-forming sequences are widely found near transcription start sites. In addition, the recent study by Liu et al. also demonstrated that Z-DNA formation is necessary for the transcription initiation of the \(CSF1\) (colony stimulating factor 1) gene.\(^8\) Thus, understanding B-Z transition mechanism of dsDNA would help to elucidate the role of Z-DNA as a potential modulator of eukaryotic transcription in vivo.

B-Z transition can be achieved in the presence of high-salt or Z-DNA binding protein, Z.\(\alpha\). Salt-induced B-Z transition is well-established in terms of chemical equilibrium. In kinetic aspect, there are two models to explain how B-Z transition undergoes in the presence of Z-DNA inducers. The all-or-none model proposed by Pohl and Jovin\(^9\) assumes that Z-forming nucleation events take place only at the ends of polymer. On the other hand, the model of Walker and Aboul-ela\(^10\) suggested that nucleation events occur at partially open or distorted base pair regions within the B-form DNA. The well-designed study by Perez et al.\(^11\) demonstrated the existence of a salt threshold in B-Z transition. A pathway of B-Z transition mechanism depends on whether the salt concentration is higher or lower than the threshold. On the other hand, so far there is almost no study available about kinetic behaviors of B-Z transition induced by hZ.\(\alpha\)ADAR1.

Our approach to study B-Z transition mechanism is to utilize hZ.\(\alpha\)ADAR1 as a Z-DNA inducer. In the co-crystal structure of hZ.\(\alpha\)ADAR1-Z-DNA,\(^12\) hZ.\(\alpha\)ADAR1 specifically interacts with phosphate backbones of Z-DNA, which explains why hZ.\(\alpha\)ADAR1 binds to Z-DNA specifically and tightly. hZ.\(\alpha\)ADAR1 is a powerful agent to induce/stabilize Z conformation in dsDNA so that it is able to flip many different sequences of DNA other than d(CG) repeats in low salt condition.\(^12\) Z-DNA has been thought to be formed in sequences composed of d(CA/TG) repeats. However, it is energetically much less favorable to form Z-DNA. Only under extreme salt conditions, poly d(AC/GT) is also able to adopt Z conformation \textit{in vitro}.\(^12\) In this study, taking advantage of hZ.\(\alpha\)ADAR1 as a powerful Z-DNA inducer, B-Z transitions of DNAs containing d(CA/TG) repeats as well as d(CG) repeats were investigated.

B-Z transitions of DNAs with different sequence compositions of d(CG) repeats and d(CA/TG) repeats were studied by using circular dichroism (CD). Table 1 summarized dsDNA substrates used for this study. First, fourteen base pair dsDNA substrates in different d(CG) and d(CA/TG) contents were designed to compare the rates of B-Z transition induced by hZ.\(\alpha\)ADAR1. CG7 and CA7 have seven repeats of d(CG) and d(CA/TG), respectively, while CACG7 has a mixed sequence containing three d(CG) repeats and four d(CA/TG) repeats, \textit{i.e.} d(CACGGCACCGCA/ TGGCGTCCGCGT). Based on thermodynamic stability for dinucleotide base pairs in Z-DNA calculated by Kagawa et al.,\(^14\) the d(CA/TG) base pair is energetically less favorable to be stabilized in Z conformation than d(CG). Thus, the order of DNA substrates energetically favorable for Z-DNA formation would be CG7 > CACG7 > CA7.
Table 1. Sequences of DNA substrates used for B-Z transition in this study

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>CA7</td>
<td>5’-CACACACACACACAC-3’</td>
</tr>
<tr>
<td>CACG7</td>
<td>5’-CACGCACGCAACAC-3’</td>
</tr>
<tr>
<td>CG7</td>
<td>5’-GCCGGCCGCACGCGC-3’</td>
</tr>
</tbody>
</table>

In the physiological condition with a low salt concentration (10 mM HEPES, pH 7.4, 10 mM NaCl), CD spectra between 240 nm and 310 nm were taken to detect B-Z transitions of dsDNA substrates by hZ$_{ADAR1}$. For kinetic study, changes of the CD signals at 255 nm were recorded for 40 min after adding hZ$_{ADAR1}$. Various temperatures ranging from 5 °C to 25 °C were used for each reaction. Figure 1 showed that hZ$_{ADAR1}$ stabilized all dsDNA oligomers (CG7, CA7, CACG7) into Z conformation. The rates of B-Z transition monitored at 255 nm showed a dependency on the (CG) dinucleotide content in the substrate DNAs (Fig. 2). Surprisingly CA7 had the fastest flipping rate among three substrates regardless of temperature (Fig. 2D). Next we carried out B-Z transition of poly d(CG) and poly d(AC/GT) induced by hZ$_{ADAR1}$ to check the effect of polymer length on flipping rate. As shown in Figure 3, the B-Z transition of poly d(AC/GT) was still faster than that of poly d(GC) in the presence of hZ$_{ADAR1}$. Thus, hZ$_{ADAR1}$ induced B-Z transition has a dependency on sequence context, not on polymer length.

Our result clearly demonstrated that hZ$_{ADAR1}$ can induce Z-DNA formation in the dsDNAs. However, at the present stage, we do not know how B-Z transition mechanism by hZ$_{ADAR1}$ occurs in detail. Nevertheless, it is possible that there are two pathways to shift B-Z equilibrium of dsDNA by hZ$_{ADAR1}$. Firstly, hZ$_{ADAR1}$ only interacts with Z conformation of DNA within DNA substrates. Small amounts of Z-DNA always exist as DNA stays in the B-Z equilibrium. When hZ$_{ADAR1}$ binds to Z-DNA, hZ$_{ADAR1}$-bound Z-DNAs are sequestered from the rest of the B-Z equilibrium. Consequently, thus, apparent B-Z equilibrium moves toward Z-DNA by accumulating hZ$_{ADAR1}$ bound Z-DNA complexes in the reaction. Secondly, hZ$_{ADAR1}$ may actively engage with DNAs to convert B-DNA into Z-DNA. When hZ$_{ADAR1}$ collides with DNA, this collision may stimulate formation of Z-DNA and it results in formation of hZ$_{ADAR1}$ bound Z-DNA complex. Inverse relationship between stabilities of DNA sequences in Z-DNA and their flipping rates by hZ$_{ADAR1}$ indicates that hZ$_{ADAR1}$ may actively shift B-Z equilibrium toward Z-DNA. Overall, our result implies that the rate of flipping by hZ$_{ADAR1}$ is dependent on strength of base-pairings on DNA track, which in turn hZ$_{ADAR1}$ could readily switch the backbone of DNA into Z conformation.

Since our kinetic study demonstrates that the flipping rates of dsDNAs by hZ$_{ADAR1}$ were inversely related to strength of base pairings within dsDNAs. This observation implies that a crucial step for B-Z transition induced by hZ$_{ADAR1}$ may need strand separation or breakage of base pair events. In this pathway, encounters between DNA and hZ$_{ADAR1}$ in the reaction may be important for the rates of B-Z transition. When hZ$_{ADAR1}$ collides with DNA, hZ$_{ADAR1}$ may rearrange DNA backbones into Z conformation. This could occur more readily if strength of base-pairing is weaker to break, which explains why B-Z transition of DNA containing high content of AT base pair is faster.

In summary, our study demonstrated that B-Z transition induced by hZ$_{ADAR1}$ behaves quite unexpectedly. When hZ$_{ADAR1}$ works as a powerful Z-DNA inducer, it stimulates Z-DNA formation by the mechanism that depends on the strength of base-pairings in DNA substrates. Consequently, the weaker base-pairing within DNA track results in the faster B-Z transition induced by hZ$_{ADAR1}$. Finally, our work extends the scope of Z-DNA study by using hZ$_{ADAR1}$ as a Z-DNA inducer. Many different sequences of dsDNAs that

Figure 1. CD spectra of the reaction mixtures after 1 hr. CD spectra of reactions with three different DNA substrates at 25 °C were obtained before adding hZ$_{ADAR1}$ (dotted line) and in the presence of hZ$_{ADAR1}$ (solid line) after 1 hr incubation, respectively. All DNA samples showed typical B-DNA spectra, while spectral changes to Z-DNA were observed as hZ$_{ADAR1}$ was added. Inversion of the CD spectrum arounds 255 nm and 292 nm is a typical characteristic of B-Z transition.
do not easily form Z conformation could be studied by their Z-DNA formation using hZαADAR1.

**Experimental Section**

**Protein and DNA preparation.** hZαADAR1 peptide was expressed and purified as described previously. Briefly, the Zα domain gene from human ADAR1 was cloned into pET28a expression vector. hZαADAR1 with the N-terminal His-tag was expressed in *E. coli* and purified by affinity purification using a His-affinity column (Amersham BioSciences). After thrombin digestion to remove the N-terminal His-tag, hZαADAR1 peptides were then further purified to homogeneity with a Hi-Trap SP column (Amersham BioSciences). The hZαADAR1 peptide was then dialyzed against buffer A (5 mM HEPES, pH 7.5, 10 mM NaCl), and concentrated to > 1 mM.

DNA oligonucleotides were synthesized and purified by HPLC. DNAs were dissolved in buffer B (10 mM Tris-Cl, pH 7.4, 50 mM NaCl, 0.1 mM EDTA) prior to annealing to form duplex DNAs. Annealing of DNA oligomers
was achieved by heating DNA samples at 90 °C for 3 min and slowly cooled down to 4 °C. Polymer DNAs were purchased from Amersham and dissolved in Buffer B prior to use.

**CD measurement.** Measurements were taken on 30 μg/μL (45 μM in base pair) of DNA in 2 mL of CD buffer (10 mM HEPES, pH 7.4, 10 mM NaCl, 0.1 mM EDTA) in a 1 cm quartz cell. CD spectra were taken at various temperatures using a J-810 CD spectrometer (Jasco, Japan). hZαADAR1 peptide was added to the sample in the final concentration of 20 μM from a concentrated stock solution, not exceeding 5% of the total volume. After 1 hr incubation, CD spectra of the reaction mixtures were then recorded between 240 nm to 310 nm at 1 nm intervals averaged over 3 sec. For kinetic profiles of the B-Z transition, time-dependent change of CD spectrum at 255 nm was monitored to measure the rate of B-Z transition induced by hZαADAR1. For this measurement, CD signal changes at 255 nm were collected at 1-sec intervals after a 5-sec dead time.

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**References**