2'-O,4'-C-Methylene Bridged Nucleic Acid Modification Promotes Pyrimidine Motif Triplex DNA Formation at Physiological pH

THERMODYNAMIC AND KINETIC STUDIES

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Extreme instability of pyrimidine motif triplex DNA at physiological pH severely limits its use in an artificial control of gene expression in vivo. Stabilization of the pyrimidine motif triplex at physiological pH is, therefore, crucial in improving its therapeutic potential. To this end, we have investigated the thermodynamic and kinetic effects of our previously reported chemical modification, 2'-O,4'-C-methylene bridged nucleic acid (2',4'-BNA), modification of triplex-forming oligonucleotide (TFO), on pyrimidine motif triplex formation at physiological pH. The thermodynamic analyses indicated that the 2',4'-BNA modification of TFO increased the binding constant of the pyrimidine motif triplex formation at neutral pH by ~20 times. The number and position of the 2',4'-BNA modification introduced into the TFO did not significantly affect the magnitude of the increase in the binding constant. The consideration of the observed thermodynamic parameters suggested that the increased rigidity itself of the 2',4'-BNA-modified TFO in the free state relative to the unmodified TFO may enable the significant increase in the binding constant at neutral pH. Kinetic data demonstrated that the observed increase in the binding constant at neutral pH by the 2',4'-BNA modification of TFO resulted from the considerable decrease in the dissociation rate constant. Our results certainly support the idea that the 2',4'-BNA modification of TFO could be a key chemical modification and may eventually lead to progress in therapeutic applications of the antigenic strategy in vivo.

In recent years, triplex DNA has attracted considerable interest because of its possible biological functions in vivo and its wide variety of potential applications, such as regulation of gene expression, site-specific cleavage of duplex DNA, mapping of genomic DNA, and gene-targeted mutagenesis (1–3). A triplex is usually formed through the sequence-specific interaction of a single-stranded homopurine or homopyrimidine triplex-forming oligonucleotide (TFO) with the major groove of homopurine-homopyrimidine stretch in duplex DNA (1–5). In the pyrimidine motif triplex, a homopyrimidine TFO binds parallel to the homopurine strand of the target duplex by Hoogsteen hydrogen bonding to form A:T and C-G:C triplets (1–5). On the other hand, in the purine motif triplex, a homopurine TFO binds antiparallel to the homopyrimidine strand of the target duplex by reverse Hoogsteen hydrogen bonding to form A:A:T (or T:A:T) and G-G:C triplets (1–5).

Because the cytosine bases in a homopyrimidine TFO must be protonated to bind with the guanine bases of the G-C duplex, the formation of the pyrimidine motif triplex needs an acidic pH condition and is thus extremely unstable at physiological pH (6–8). Instead, the pH-independent formation of the purine motif triplex is available at physiological pH. However, the purine motif triplex formation is severely inhibited by physiological concentrations of certain monovalent cations, especially K⁺. Undefined association between K⁺ and the guanine-rich homopurine TFO has been applied to explain the inhibitory effect (9, 10). Therefore, stabilization of the pyrimidine motif triplex at physiological pH is of great importance in improving its therapeutic potential to artificially control gene expression in vivo. Numerous efforts such as the replacement of cytosine bases in a homopyrimidine TFO with 5-methylcytosine (7, 11–13) or other chemically modified bases (14–18), the conjugation of different DNA intercalators to TFO (19, 20), and the use of polyamines such as spermine or spermidine as triplex stabilizers (21) have been made to improve the stability of the pyrimidine motif triplex at physiological pH.

We first synthesized and developed a new class of chemical modifications of nucleic acids, bridged nucleic acid (BNA), such as 2'-O,4'-C-methylene BNA (2',4'-BNA; Fig. 1a; Refs. 22–27) and 3'-O,4'-C-methylene BNA (3',4'-BNA; Refs. 30–32). The 2',4'-BNA modification of TFO increased the thermal stability of the pyrimidine motif triplex DNA at neutral pH using a homopyrimidine-homopurine target duplex and its specific cytosine-rich TFO (25). However, the mechanistic explanation for the 2',4'-BNA modification-mediated triplex stabilization was not clearly understood. Here, therefore, we have further ex-

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1 The abbreviations used are: TFO, triplex-forming oligonucleotide; BNA, bridged nucleic acid; 2',4'-BNA, 2'-O,4'-C-methylene bridged nucleic acid; 3',4'-BNA, 3'-O,4'-C-methylene bridged nucleic acid; EMSA, electrophoretic mobility shift assay; CD, circular dichroism; ITC, isothermal titration calorimetry; IAsys, interaction analysis system; Pur, purine; Pyr, pyrimidine; Bt, biotinylated; NS, nonspecific; Tm, melting temperature; koff, off-rate constant; kassoc, association rate constant; koff, dissociation rate constant.

2 After our report on the first synthesis of 2',4'-BNA monomers, Wengel’s group demonstrated some properties of 2',4'-BNA (28, 29).
tended our previous study to explore thermodynamic and kinetic effects of the 2',4'-BNA modification on the pyrimidine motif triplex formation at neutral pH. The thermodynamic and kinetic effects of the 2',4'-BNA modification on the pyrimidine motif triplex formation between a 23-base pair homopurine-homopyrimidine target duplex (Pur23A-Pyr23T; Fig. 1b) and its specific 15-mer unmodified or 2',4'- BNA-modified homopyrimidine TFO (Pyr15T, Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, and Pyr15BNA5-2; Fig. 1b) have been analyzed by electrophoretic mobility shift assay (EMSA; Refs. 33, 34), UV melting, isothermal titration calorimetry (ITC; Refs. 34–40), and interaction analysis system (IAsys; Refs. 34, 39–43). Results from these independent lines of experiments have clearly indicated the significant effect of the 2',4'-BNA modification to promote the pyrimidine motif triplex formation at neutral pH. The binding constant at neutral pH for the pyrimidine motif triplex formation with the 2',4'-BNA-modified TFO was ~10–20 times larger than that observed with the corresponding unmodified TFO. Kinetic data have also demonstrated that the contribution for the increase in the binding constant by the 2',4'-BNA modification of TFO resulted from the considerable decrease in the dissociation rate constant. The ability of the 2',4'-BNA modification of TFO to promote the pyrimidine motif triplex formation at physiological pH would support further progress in therapeutic applications of the antigenic strategy in vivo.

MATERIALS AND METHODS

Preparation of Oligonucleotides—We synthesized 23-mer complementary oligonucleotides for the target duplex Pur23A and Pyr23T (Fig. 1b), a 15-mer unmodified homopyrimidine TFO specific for the target duplex Pyr15T (Fig. 1b), and a nonspecific homopyrimidine oligonucleotide, Pyr15NS (Fig. 1b), on an Applied Biosystems DNA synthesizer using the solid-phase cyanophosphoramidite method and purified them with a reverse-phase high performance chromatography on a Wakosil DNA column. The 15-mer 2',4'-BNA-modified homopyrimidine TFOs Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, and Pyr15BNA5-2 (Fig. 1b) were synthesized and purified as described previously (22, 28). 5’ biotinylated Pyr23T (denoted Bt-Pyr23T) was prepared using biotin phosphoramidite. The concentration of all oligonucleotides was determined by UV absorbance. Complementary strands Pur23A and Pyr23T were annealed by heating to up to 90 °C, followed by a gradual cooling to room temperature. The annealed sample was applied on a hydroxyapatite column (Koken Inc.) to remove unpaired single strands. The concentration of the duplex DNA (Pur23A-Pyr23T) was determined by UV absorption considering the DNA concentration ratio of 1 optical density unit = 50 μg/ml, with an M, of 15180.

EMSA—EMSA experiments were performed essentially as described previously (34). In 9 μl of reaction mixture, 700-μl labeled duplex (~1 ng) was mixed with increasing concentrations of the specific TFO (Pyr15T, Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2) and the nonspecific oligonucleotide (Pyr15NS) in a buffer containing 50 mM Tris acetate, pH 7.0, 100 mM sodium chloride, and 10 mM magnesium chloride. Pur15NS was added to achieve equimolar concentrations of TFO in each lane as well as to minimize adhesion of the DNA (target duplex and TFO) to plastic surfaces during incubation and subsequent loss during processing. After 6 h of incubation at 37 °C, 2 μl of 50% glycerol solution containing bromphenol blue was added without changing the pH and salt concentrations of the reaction mixtures. Samples were then directly loaded onto a 15% native polyacrylamide gel prepared in buffer containing bromphenol blue was added without changing the pH and salt concentrations of the reaction mixtures. Samples were then loaded onto a 15% native polyacrylamide gel prepared in buffer solution containing bromphenol blue. The positions of the DNA bands were determined by UV absorbance. Complementary strands Pur23A and Pyr23T were annealed by heating to up to 90 °C, followed by a gradual cooling to room temperature. The annealed sample was applied on a hydroxyapatite column (Koken Inc.) to remove unpaired single strands. The concentration of the duplex DNA (Pur23A-Pyr23T) was determined by UV absorption considering the DNA concentration ratio of 1 optical density unit = 50 μg/ml, with an M, of 15180.

UV Melting—UV melting experiments were carried out using a Jasco J-720 spectrophotometer equipped with an HEC-363 Peltier-type cell holder controlled by a TPU-436 temperature programmer. UV melting profiles were measured in buffer A (10 mM sodium cacodylate-cacodylic acid, pH 6.8, containing 200 mM NaCl and 20 mM MgCl2) at a scan rate of 0.5 °C/min to 260 nm. The first derivative was calculated from the UV melting profile. The peak temperatures in the derivative curve were designated the melting temperatures (Tm). Cell path length was 1 cm. The triplex DNA concentration used was 1 μM.

Circular Dichroism (CD) Spectroscopy—CD spectra at room temperature were recorded in buffer A on a Jasco J-720 spectropolarimeter interfaced with a microcomputer. Cell path length was 1 cm. The triplex DNA concentration used was 4 μM.

RESULTS

Electrophoretic Mobility Shift Assay of Pyrimidine Motif Triplex Formation at Neutral pH—The pyrimidine motif triplex formation of the target duplex (Pur23A-Pyr23T; Fig. 1b) with unmodified (Pyr15T; Fig. 1b) or 2',4'-BNA-modified (Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2; Fig. 1b) TFO was examined at pH 7.0 by EMSA (Fig. 2). The total oligonucleotide concentration (specific TFO (Pyr15T, Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2; Fig. 1b) + [non-
specific oligonucleotide (Pyr15NS; Fig. 1b)) was kept constant at 1 μM to minimize loss of DNA during processing. Although incubation with 1 μM Pyr15NS alone did not cause a shift in electrophoretic migration of the target duplex (see Fig. 2, lane 1 for Pyr15T), those with Pyr15T, Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2 at particular concentration caused retardation of the duplex migration owing to triplex formation (33). The $K_0$ of triplex formation was determined from the concentration of the TFO, which caused half of the target duplex to shift to the triplex (33). The $K_0$ of the triplex with Pyr15T was estimated to be 1.0 μM. In contrast, the $K_0$ of the triplex with Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2 was 0.06 μM, indicating that the 2′,4′-BNA modification of TFO increased the binding affinity of the pyrimidine motif triplex formation at neutral pH by ~20 times. The increase in the $K_0$ by the 2′,4′-BNA modification was similar in magnitude among the four modified TFOs.

Spectroscopic Characterization of Pyrimidine Motif Triplex at Neutral pH—The thermal stability of the pyrimidine motif triplex with unmodified (Pyr15T) or 2′,4′-BNA-modified (Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2) TFO was investigated at pH 6.8 by UV melting (Fig. 3 and Table I). All the triplexes showed two-step melting. The first transition at a lower temperature, $T_{m1}$, was the melting of the triplex to a duplex and a TFO, and the second transition at a higher temperature, $T_{m2}$, was the melting of the duplex (Fig. 3). Although the $T_{m2}$ was almost identical among all the triplexes, the $T_{m1}$ for Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2 was significantly higher than that for Pyr15T (Table I), indicating that the 2′,4′-BNA modification of TFO increased the thermal stability of the pyrimidine motif triplex at neutral pH.

To further characterize the triplexes involving unmodified (Pyr15T) or 2′,4′-BNA-modified (Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2) TFO, CD spectra of the triplexes were measured at 25 °C and pH 6.8 (Fig. 4). The overall shape of the CD spectra was similar among all the profiles. A negative band in the short-wavelength (210–220-nm) region was observed for all the profiles, confirming the triplex formation involving each TFO (44, 45). The intensity of the negative short-wavelength (210–220-nm) band for the triplexes involving Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2 was larger than that observed for the triplex involving Pyr15T, indicating that all the pyrimidine motif triplexes involving the 2′,4′-BNA-modified TFO had more aspects in the A conformation than that involving the unmodified TFO (45).

Thermodynamic Analyses of Pyrimidine Motif Triplex Formation at Neutral pH by ITC—We examined the thermodynamic parameters of the pyrimidine motif triplex formation between a 23-base pair target duplex (Pur23A–Pyr23T) and its specific 15-mer unmodified (Pyr15T) or 2′,4′-BNA-modified (Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2) TFO at 25 °C and pH 6.8 by ITC. To investigate the pH dependence of the pyrimidine motif triplex formation, the thermodynamic parameters of the triplex formation between Pur23A–Pyr23T and Pyr15T were also analyzed at 25 °C and pH 5.8 by ITC. Fig. 5a compares the ITC profiles of the initial
The thermodynamic analyses of the pyrimidine motif triplex formation with Pyr15T or Pyr15BNA7-1 at pH 6.8 and with Pyr15T at pH 5.8 by ITC. a, ITC profiles of the initial three injections for the binding of Pur23A-Pyr23T to Pyr15T or Pyr15BNA7-1 at 25 °C and pH 6.8 and to Pyr15T at 25 °C and pH 5.8. The TFO solution (120 µl in buffer A or buffer B; see "Materials and Methods") was injected in 5-µl increments into 5-µl Pur23A-Pyr23T solution in the same buffer. Injections occurred over 10 s at 10-min intervals. b, total ITC profile for the triplex formation between Pyr15BNA7-1 and Pur23A-Pyr23T. The Pyr15BNA7-1 solution was injected 20 times into the Pur23A-Pyr23T solution. Other experimental conditions were the same as in a. c, titration plots against the molar ratio of [TFO]/[Pur23A-Pyr23T]. The data were fitted by a nonlinear least-squares method.

three injections for the binding of Pur23A-Pyr23T with Pyr15T or Pyr15BNA7-1 at pH 6.8 and with Pyr15T at pH 5.8. The magnitudes of the exothermic peaks for Pyr15BNA7-1 at pH 6.8 and for Pyr15T at pH 5.8 were larger than those observed for Pyr15T at pH 6.8. Fig. 5b shows an ITC profile over 200 min for triplex formation with Pyr15BNA7-1 at pH 6.8. An exothermic heat pulse was observed after each injection of Pyr15BNA7-1 into Pur23A-Pyr23T. The magnitude of each peak decreased gradually with each new injection, and a small peak was still observed at a molar ratio of [Pyr15BNA7-1]/[Pur23A-Pyr23T] = 2. The area of the small peak was equal to the heat of dilution measured in a separate experiment by injecting Pyr15BNA7-1 into the same buffer (data not shown). The area under each peak was integrated, and the heat of dilution of Pyr15BNA7-1 was subtracted from the integrated values. The corrected heat was divided by the moles of injected solution, and the resulting values were plotted as a function of a molar ratio of [Pyr15BNA7-1]/[Pur23A-Pyr23T], as shown in Fig. 5c. The resultant titration plot was fitted to a sigmoidal curve by a nonlinear least-squares method. The binding constant, $K_a$, and the enthalpy change, $\Delta H$, were obtained from the fitted curve (37). The Gibbs free energy change, $\Delta G$, and the entropy change, $\Delta S$, were calculated from the equation, $\Delta G = -RT \ln K_a = \Delta H - T \Delta S$ (37) where $R$ is gas constant and $T$ is temperature. The titration plots for Pyr15T at pH 5.8 and pH 6.8 are also shown in Fig. 5c. The thermodynamic parameters for Pyr15T at pH 5.8 and 6.8 were obtained from the titration plots in the same way. The ITC profiles and the titration plots for Pyr15BNA7-2, Pyr15BNA5-1, and Pyr15BNA5-2 at pH 6.8 were almost the same as those observed for Pyr15BNA7-1 at pH 6.8 (data not shown). The thermodynamic parameters for Pyr15BNA7-2, Pyr15BNA5-1, and Pyr15BNA5-2 at pH 6.8 were obtained in the same way.

Table II summarizes the thermodynamic parameters for the pyrimidine motif triplex formation with all the TFOs at 25 °C and pH 6.8 and those with Pyr15T at 25 °C and pH 5.8, obtained from ITC. The signs of both $\Delta H$ and $\Delta S$ were negative under all the conditions. Because an observed negative $\Delta S$ was unfavorable for the triplex formation, the triplex formation was driven by a large negative $\Delta H$ under each condition. The magnitudes of the negative $\Delta H$ of the triplex formation for Pyr15T at pH 5.8 and for Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2 at pH 6.8 were 2.5 or 1.6–1.7 times larger than that observed for Pyr15T at pH 6.8, consistent with the ITC profiles in Fig. 5a. The $K_a$ for Pyr15T at pH 5.8 was ~20 times larger than that observed for Pyr15T at pH 6.8, confirming, like others (6–8), that neutral pH is unfavorable for the pyrimidine motif triplex formation involving C-G-C triads. In addition, the $K_a$ for Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2 at pH 6.8 was 10–20 times larger than that observed for Pyr15T at pH 6.8, indicating that the 2'-4'-BNA modification of TFO increased the $K_a$ of the pyrimidine motif triplex formation at neutral pH by 10–20 times. The increase in the $K_a$ by the 2’,4’-BNA modification of TFO was similar in magnitude among the four modified TFOs.

Kinetic Analyses of Pyrimidine Motif Triplex Formation at Neutral pH by IAsys—To examine the putative mechanism involved in the increase in $K_a$ of the pyrimidine motif triplex formation by the 2’,4’-BNA modification (Fig. 2 and Table II), we assessed the kinetic parameters for the association and dissociation of TFO (Pyr15T, Pyr15BNA7-1, Pyr15BNA7-2, Pyr15BNA5-1, or Pyr15BNA5-2) with Pur23A-Pyr23T at 25 °C and pH 6.8 by IAsys. Fig. 6a compares the sensorgrams representing the triplex formation and dissociation involving 90 µM of the specific TFO (Pyr15T or Pyr15BNA7-1). The injection of Pyr15T over the immobilized Bi-Pyr23T-Pur23A caused an increase in response. The response was substantially delayed on the injection of Pyr15BNA7-1, indicating that the 2’,4’-BNA modification decreased the association rate constant of the triplex equilibrium. In contrast, the change in the dissociation curve with time for Pyr15BNA7-1 was much smaller than that for Pyr15T. The result clearly indicated that the 2’,4’-BNA modification of TFO remarkably decreased the dissociation rate constant of the triplex equilibrium. The similar profiles were obtained for Pyr15BNA7-2, Pyr15BNA5-1, and Pyr15BNA5-2.

To understand the kinetic parameters more quantitatively, we analyzed a series of association and dissociation curves at the various concentrations of TFO. As shown in Fig. 6b, an increase in the concentration of the 2’,4’-BNA-modified TFO (Pyr15BNA7-1) led to a gradual change in the response of the association curves. The $k_{on}$ was obtained from the analysis of each association curve. Fig. 6c shows a plot of $k_{on}$ against the Pyr15BNA7-1 concentrations. The resultant plot was fitted to a straight line by a linear least-squares method. The association rate constant ($k_{on}$) and the dissociation rate constant ($k_{dissoc}$) were determined from the slope and the intercept of the regression line, respectively (41–43). The kinetic parameters for the triplex formation with Pyr15BNA7-2, Pyr15BNA5-1, and

Pyrimidine Motif Triplex DNA Formation at Physiological pH
Pyrimidine Motif Triplex DNA Formation at Physiological pH

**Table II**

<table>
<thead>
<tr>
<th>TFO</th>
<th>pH</th>
<th>$K_a$ (relative)</th>
<th>$\Delta G$ kcal/mol</th>
<th>$\Delta H$ kcal/mol</th>
<th>$\Delta S$ cal/mol/K</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyr15T</td>
<td>5.8a</td>
<td>(3.83 ± 0.74) $\times 10^6$</td>
<td>19.4</td>
<td>−8.98 ± 0.13</td>
<td>−85.6 ± 2.6</td>
</tr>
<tr>
<td>Pyr15T</td>
<td>6.8b</td>
<td>(1.97 ± 0.43) $\times 10^7$</td>
<td>10.0</td>
<td>−7.22 ± 0.15</td>
<td>−34.9 ± 2.2</td>
</tr>
<tr>
<td>Pyr15BNA7–1</td>
<td>6.8b</td>
<td>(2.28 ± 0.32) $\times 10^7$</td>
<td>11.6</td>
<td>−8.67 ± 0.09</td>
<td>−55.5 ± 1.5</td>
</tr>
<tr>
<td>Pyr15BNA7–2</td>
<td>6.8b</td>
<td>(2.16 ± 0.08) $\times 10^7$</td>
<td>11.0</td>
<td>−8.64 ± 0.11</td>
<td>−57.3 ± 2.0</td>
</tr>
<tr>
<td>Pyr15BNA5–1</td>
<td>6.8b</td>
<td>(3.65 ± 0.54) $\times 10^7$</td>
<td>18.5</td>
<td>−8.95 ± 0.10</td>
<td>−60.7 ± 1.4</td>
</tr>
<tr>
<td>Pyr15BNA5–2</td>
<td>6.8b</td>
<td>(2.06 ± 0.42) $\times 10^7$</td>
<td>10.5</td>
<td>−8.61 ± 0.14</td>
<td>−60.5 ± 2.6</td>
</tr>
</tbody>
</table>

*10 mM sodium cacodylate-cacodylic acid, pH 5.8, 200 mM sodium chloride, and 20 mM magnesium chloride.

not shown), that is, the dissociation was very slow, we could not directly determine the $k_{\text{dissoc}}$ from the dissociation curves. Thus, the $k_{\text{dissoc}}$ obtained from the association curves was presented for Pyr15BNA7–1, Pyr15BNA7–2, Pyr15BNA5–1, and Pyr15BNA5–2, although its S.D. was relatively large (Table III). On the other hand, although the $k_{\text{assoc}}$ for Pyr15T was obtained in the same way as described above, the $k_{\text{dissoc}}$ was determined in the following way owing to the much faster dissociation. The off-rate constant ($k_{\text{off}}$) was obtained from the analysis of each dissociation curve (Fig. 6a; data not shown). Because $k_{\text{off}}$ is usually independent of the concentration of the injected solution, the $k_{\text{dissoc}}$ was determined by averaging $k_{\text{off}}$ for several concentrations (41–43). $K_a$ was calculated from the equation $K_a = k_{\text{assoc}}/k_{\text{dissoc}}$.

Table III summarizes the kinetic parameters for the pyrimidine motif triplex formation with all the TFOs at 25 °C and pH 6.8, obtained from IAsys. The magnitudes of $K_a$ calculated from the ratio of $k_{\text{assoc}}$ to $k_{\text{dissoc}}$ (Table III) were consistent with those obtained from ITC (Table II). The 2′-4′-BNA modification of TFO increased the $K_a$ for the pyrimidine motif triplex formation at neutral pH, which supported the results of EMSA (Fig. 2) and ITC (Table II). The $k_{\text{dissoc}}$ of the triplex formation decreased 40–70 times by the 2′,4′-BNA modification of TFO. In contrast, when the $k_{\text{dissoc}}$ of the triplex formation was compared, 1–1.3 times smaller $k_{\text{dissoc}}$ was obtained by the 2′,4′-BNA modification of TFO, which opposes the increase in $K_a$. Thus, the much larger $K_a$ by the 2′,4′-BNA modification resulted from the decrease in $k_{\text{dissoc}}$. The kinetic effect to increase the $K_a$ by the 2′,4′-BNA modification was similar among the four modified TFOs.

**DISCUSSION**

The $K_a$ of the pyrimidine motif triplex formation with Pyr15T at pH 5.8 was 20 times larger than that observed with Pyr15T at pH 6.8 (Table II), which is consistent with the previously reported results that the neutral pH is unfavorable for the pyrimidine motif triplex formation involving C–GC triads (6–8). The $K_a$ of the triplex formation with Pyr15BNA7–1, Pyr15BNA7–2, Pyr15BNA5–1, or Pyr15BNA5–2 at pH 6.8 was 10–20 times larger than that observed with Pyr15T at pH 6.8 (Table II). Thus, the much larger $K_a$ by the 2′,4′-BNA modification of TFO was supported by the results of EMSA (Fig. 2) and IAsys (Table III). In addition, the 2′,4′-BNA modification of TFO increased the thermal stability of the pyrimidine motif triplex at pH 6.8 (Table I). These results indicate that the 2′,4′-BNA modification of TFO considerably promotes the pyrimidine motif triplex formation at neutral pH.

The $\Delta S$ on the triplex formation measured by ITC reflects a major contribution from the hydrogen bonding and the base stacking involved in the triplex formation (38, 46–48). On the other hand, the $\Delta S$ on the triplex formation measured by ITC includes a positive entropy change from release of structured water on the triplex formation and a major contribution of a
negative conformational entropy change from the conformational restraint of TFO involved in the triplex formation (38, 46–48). Because the formed triplex structure involving Pyr15T at pH 5.8 and that involving Pyr15T at pH 6.8 is the same, the magnitude of $\Delta H$ and $\Delta S$ on the triplex formation measured by ITC could be the same between the two conditions. However, unexpectedly, the magnitudes of $\Delta H$ and $\Delta S$ for Pyr15T at pH 6.8 were significantly smaller than those observed for Pyr15T at pH 5.8 (Table II). When the $\Delta H$ and $\Delta S$ are calculated from the fitting procedure of ITC, the heat observed by ITC is divided not by the effective concentration really involved in the triplex formation but by the apparent concentration added to the triplex formation (37). The calculation does not take it into consideration what percentage of the added concentration is really effectively involved in the triplex formation. Thus, if the triplex formation is less stoichiometric under a certain condition, the magnitudes of $\Delta H$ and $\Delta S$ for the less stoichiometric triplex formation estimated by ITC are smaller than those observed for the more stoichiometric triplex formation under another condition. Therefore, the significantly smaller magnitudes of $\Delta H$ and $\Delta S$ for Pyr15T at pH 6.8 relative to those for Pyr15T at pH 5.8 (Table II) suggest that the triplex formation with Pyr15T at pH 5.8 was significantly less stoichiometric than that with Pyr15T at pH 5.8, which was also supported by the significantly smaller magnitudes of $K_a$ and $\Delta G$ for Pyr15T at pH 5.8 (Table II). In contrast, the $K_a$ and $\Delta G$ for Pyr15T at pH 5.8 and those for the 2',4'-BNA modified TFOs at pH 6.8 were quite similar (Table II), suggesting that the triplex formation under the two conditions was similarly quite stoichiometric. We conclude that the triplex formation with Pyr15T at pH 5.8 was significantly less stoichiometric than that with Pyr15T at pH 5.8 and that with the 2',4'-BNA-modified TFOs at pH 6.8. Thus, to discuss the promotion mechanism of the triplex formation by the 2',4'-BNA modification, the comparison of the $\Delta H$ and $\Delta S$ between Pyr15T at pH 6.8 and 2',4'-BNA-modified TFOs at pH 6.8 is not valid because of the significantly reduced stoichiometry for Pyr15T at pH 6.8. The comparison of the $\Delta H$ and $\Delta S$ between Pyr15T at pH 5.8 and 2',4'-BNA-modified TFOs at pH 6.8 with similar stoichiometry will provide a reasonable promotion mechanism for the triplex formation by the 2',4'-BNA modification, as discussed below.

Although the $K_a$ and $\Delta G$ for Pyr15T at pH 5.8 and those for the 2',4'-BNA-modified TFOs at pH 6.8 were quite similar (Table II), the ingredients of $\Delta G$, that is, $\Delta H$ and $\Delta S$, were obviously different from each other. The magnitudes of the negative $\Delta H$ and $\Delta S$ for the 2',4'-BNA-modified TFOs at pH 6.8 were smaller than those observed for Pyr15T at pH 5.8 (Table II). The hydrogen bonding and the base stacking involved in the triplex formation are usually considered the major sources of the negative $\Delta H$ on the triplex formation (38, 46–48). Thus, the difference in $\Delta H$ for the stoichiometric triplex formations between Pyr15T at pH 5.8 and the 2',4'-BNA-modified TFOs at pH 6.8 (Table II) suggests that the hydrogen bonding and the base stacking of the triplex with the 2',4'-BNA-modified TFOs are significantly different from those with the corresponding unmodified TFO. In fact, the CD spectra show that the triplexes with the 2',4'-BNA-modified TFO had the A-like conformation (Ref. 45 and Fig. 4). The A-like conformation by the 2',4'-BNA modification of TFO may result in the difference in the negative $\Delta H$ between the unmodified and 2',4'-BNA-modified TFOs. On the other hand, the negative $\Delta S$ on the triplex formation is mainly contributed by a negative conformational entropy change attributable to the conformational restraint of TFO involved in the triplex formation (38, 46–48). Therefore, the smaller magnitude of the negative $\Delta S$ for the 2',4'-BNA modified TFOs at pH 6.8 relative to that for Pyr15T at pH 5.8 (Table II) suggests that the 2',4'-BNA-modified TFO in the free state is more rigid than the corresponding unmodified TFO.

The increased rigidity of the 2',4'-BNA modified TFO in the free state relative to the corresponding unmodified TFO causes the smaller entropic loss on the triplex formation with the 2',4'-BNA-modified TFO at neutral pH, which provides a favorable component to the $\Delta G$ and leads to the increase in the $K_a$ of the triplex formation at neutral pH. We conclude that the increased rigidity of the 2',4'-BNA-modified TFO in the free state may be one of the factors that increases the $K_a$ of the pyrimidine motif triplex formation at neutral pH.

The increase in the $K_a$ by the 2',4'-BNA modification was similar in magnitude among the four modified TFOs (Table II and III), indicating that the number and position of the 2',4'-BNA modification did not significantly affect the magnitude of the increase in the $K_a$ at neutral pH. The rigidity itself of the 2',4'-BNA-modified TFO may be more important to achieve the increase in the $K_a$ at neutral pH than the variation of the number and position of the 2',4'-BNA modification. Thus, other modification strategies to gain the increased rigidity of TFO may also be useful to increase the $K_a$ at neutral pH.

Kinetic data have demonstrated that the 2',4'-BNA modification of TFO considerably decreased the $k_{\text{dissoc}}$ of the pyrimidine motif triplex formation (Table III). The decrease in the $k_{\text{dissoc}}$ is a plausible kinetic reason to explain the remarkable gain in the $K_a$ at neutral pH by the 2',4'-BNA modification (Fig. 2 and Tables II and III). Both our group (38) and others (49) have previously proposed a model that triplexes form along nucleation-elongation processes: in a nucleation step only a few base contacts of the Hoogsteen base pairings provide a reasonable promotion mechanism for the triplex formation by the 2',4'-BNA modification, as discussed below.

### Table III

<table>
<thead>
<tr>
<th>TFO</th>
<th>$k_{\text{assoc}}$</th>
<th>$k_{\text{dissoc}}$</th>
<th>$K_a$ (relative)</th>
<th>$K_{\text{dissoc}}$ (relative)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyr15T</td>
<td>$(6.31 \pm 0.18) \times 10^2$</td>
<td>$1.0$</td>
<td>$1.0$</td>
<td>$(5.41 \pm 0.91) \times 10^4$</td>
</tr>
<tr>
<td>Pyr15BNA7–1</td>
<td>$(4.86 \pm 0.69) \times 10^2$</td>
<td>$0.77$</td>
<td>$0.013$</td>
<td>$(3.09 \pm 0.39) \times 10^6$</td>
</tr>
<tr>
<td>Pyr15BNA7–2</td>
<td>$(5.05 \pm 0.77) \times 10^2$</td>
<td>$0.80$</td>
<td>$0.200$</td>
<td>$(2.15 \pm 1.36) \times 10^6$</td>
</tr>
<tr>
<td>Pyr15BNA5–1</td>
<td>$(5.93 \pm 0.93) \times 10^2$</td>
<td>$0.94$</td>
<td>$0.025$</td>
<td>$(2.00 \pm 1.02) \times 10^6$</td>
</tr>
<tr>
<td>Pyr15BNA5–2</td>
<td>$(6.32 \pm 0.34) \times 10^2$</td>
<td>$1.0$</td>
<td>$(2.38 \pm 1.00) \times 10^6$</td>
<td></td>
</tr>
</tbody>
</table>

Parameters were at 25°C and pH 6.8 in 10 mM sodium cacodylate-cacodylic acid, 200 mM sodium chloride, and 20 mM magnesium chloride, obtained from IAsys.
The present study has clearly demonstrated that the 2',4'-BNA modification of TFO promotes pyrimidine motif triplex formation at neutral pH. We conclude that the design of TFO to bridge different positions of sugar moiety with the alkyl chain to gain the increased rigidity of TFO is certainly a promising strategy for the promotion of triplex formation under physiological condition and may eventually lead to progress in therapeutic applications of the antigene strategy in vivo.

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REFERENCES