Design of Tail-Clamp Peptide Nucleic Acid Tethered with Azobenzene Linker for Sequence-Specific Detection of Homopurine DNA

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Abstract: DNA carries genetic information in its sequence of bases. Synthetic oligonucleotides that can sequence-specifically recognize a target gene sequence are a useful tool for regulating gene expression or detecting target genes. Among the many synthetic oligonucleotides, tail-clamp peptide nucleic acid (TC-PNA) offers advantages since it has two homopyrimidine PNA strands connected via a flexible ethylene glycol-type linker that can recognize complementary homopurine sequences via Watson-Crick and Hoogsteen base pairings and form thermally-stable PNA/PNA/DNA triplex structures. Here, we synthesized a series of TC-PNAs that can possess different lengths of azobenzene-containing linkers and studied their binding behaviours to homopurine single-stranded DNA. Introduction of azobenzene at the N-terminus amine of PNA increased the thermal stability of PNA-DNA duplexes. Further extension of the homopyrimidine PNA strand at the N-terminus of PNA-AZO further increased the binding stability of the PNA/DNA/PNA triplex to the target homopurine sequence; however, it induced TC-PNA/DNA/TC-PNA complex formation. Among these TC-PNAs, 9W5H-C4-AZO consisting of nine Watson-Crick bases and five Hoogsteen bases tethered with a beta-alanine conjugated azobenzene linker gave a stable 1:1 TC-PNA/ssDNA complex and exhibited good mismatch recognition. Our design for TC-PNA-AZO can be utilized for detecting homopurine sequences in various genes.

Keywords: peptide nucleic acid (PNA); tail-clamp PNA; detection of homopurine DNA; azobenzene-containing linkers

1. Introduction

Sequence-specific detection of target genes using oligonucleotides is a key technology for detecting pathogens and disease-related genes. The accuracy and sensitivity of the diagnosis relies on the chemical properties of the oligonucleotides. Thus, various types of chemically modified nucleic acids have been developed to improve binding affinity and sequence specificity.

Peptide nucleic acid (PNA) is a DNA mimic in which the phosphate backbone has been replaced by a neutral amide backbone composed of N-(2-aminoethyl) glycine linkages (Figure 1A) [1]. The advantages of PNA are its high binding affinity [2–4], good mismatch discrimination [5], nuclease and protease resistance [6], and low affinity for proteins [7]. Tail-clamp PNA (TC-PNA), which is composed of two homopyrimidine PNA strands connected via a linker molecule, 2-aminoethoxy-2-ethoxy acetic acid (AEEA, Figure 1B), can form a stable complex with its target homopurine DNA sequence consisting partly of a duplex via Watson-Crick base pairing and partly of a triplex via both...
We recently developed a novel type of azobenzene amino acid [17] and used it as a linker for TC-PNA to Watson-Crick and Hoogsteen base pairings [1,2,8,9]. Therefore, TC-PNA can form a thermally-stable PNA/DNA/PNA triplex [5,10,11]. This efficient DNA binding of TC-PNA has been used for down-regulating DNA transcription [3,12,13], DNA repair [14,15] and capturing bacterial genes [16]. We recently developed a novel type of azobenzene amino acid [17] and used it as a linker for TC-PNA to enhance the binding characteristics of PNA to a complementary RNA sequence on the influenza A virus gene [18].

![Chemical structure images](image1.png)

**Figure 1.** Chemical structure of (A) PNA monomer; (B) ethylene glycol-based aminoethoxyethoxy acetate linker (AEEA); (C) azobenzene amino acid linker (C0-AZO); and (D) spacer conjugated azobenzene amino acid linker (C3-, C4-, C5- and C6-AZO (n = 1, 2, 3 and 4)); (E) Schematic image of triplex formation between TC-PNA and ssDNA. The C-terminal strand (PNA (WC)) of TC-PNA was designed to form Watson-Crick base pairs with the N-terminal strand (PNA (H)) of TC-PNA was designed to form Hoogsteen base pairs with its target ssDNA; and (F) one possible image of a TC-PNA/TC-PNA/ssDNA trimer.

In this study, we synthesized homopyrimidine TC-PNAs that consist of different lengths of azobenzene linker (Figure 1C,D, TC-PNA AZO) and different numbers of homopyrimidine bases strands. Their binding mode of action to complementary homopurine single stranded DNAs (ssDNAs) was assessed by gel mobility shift assay, melting temperature analysis, and electrospray ionization mass spectrometry. Optimization of both the lengths of the azobenzene linker and the homopyrimidine strands of TC-PNA efficiently provided a 1:1 complex with complementary homopurine ssDNA with high binding affinity and good sequence specificity. This result allows us to design homopyrimidine TC-PNA that recognizes homopurine DNA.

## 2. Results and Discussions

### 2.1. Design of PNAs and TC-PNAs

We designed TC-PNA molecules composed of two PNA strands tethered with different types of spacer molecules (Table 1). TC-PNA has been demonstrated to form complex structures consisting of partially triplex and partially duplex regions with its target ssDNA by Watson-Crick and Hoogsteen base pairings [10] (Figure 1E). Since tethering two PNA strands reduces the entropic penalty paid during triplex formation by two distinct PNA molecules, TC-PNA exhibits improved binding affinity
to ssDNA [19]. On the other hand, TC-PNA potentially forms a trimer complex (Figure 1F) when excess TC-PNA is added to the target ssDNA.

Recently, we reported that TC-PNA tethered with an azobenzene linker showed a nearly 10-fold improved inhibitory effect on the reverse transcription of the influenza A virus gene by MuLV reverse transcriptase [18]. These reports indicate that the linker structure of TC-PNA affects its binding characteristics with target sequences. However, homopyrimidine TC-PNAs targeting homopurine sequences may potentially form a TC-PNA/TC-PNA/DNA trimer [20]. To investigate the effect of the linker on the binding characteristics of homopyrimidine TC-PNA to homopurine ssDNA, we synthesized a series of linker molecules (Figure 1B–D) and individually introduced them as linkers for TC-PNA by Fmoc-solid phase synthesis (Table 1). The homopyrimidine PNA strand contains a complementary sequence to a target ssDNA (DNA1), while the linker unit contains different number of a flexible ethylene glycol based linkers (AEEA) or different lengths of rigid azobenzene linkers. PNA1 and 2 were synthesized to confirm the effect of azobenzene on its binding characteristics to the complementary ssDNA (Table 1). PNA3–5 that possess different numbers of flexible AEEA linkers were synthesized to study the effect of linker length on the binding properties of TC-PNAs (Table 1). On the other hand, PNA6–10 that possess different lengths of azobenzene linkers were synthesized to study how azobenzene linkers affect the binding properties of TC-PNAs (Table 1). PNA11–17 were synthesized to study the effect of the Hoogsteen strand of TC-PNA on their binding characteristics to ssDNA (Table 1). All PNAs have a lysine molecule at the C-terminus to increase water solubility. Three consecutive lysine residues at the N-terminal of the TC-PNAs were used to enhance the binding affinity of the Hoogsteen strand to the target DNA [21].

| Entry | Name | PNA Sequence (N to C)/DNA Sequence (5′ to 3′) | Mass (Found) |
|-------|------|--------------------------------------------|-------------|
| PNA1  | 12W  | TCTCCCTTCTTT-Lys                          | 3266.44     |
| PNA2  | 12W-C0-AZO | AZO-TCTCCCTTCTTT-Lys                  | 3549.80     |
| PNA3  | 12W5H-(AEEA) | H-(Lys)3-CCTCT-(AEEA)-TCTCCCTTCTTT-Lys     | 5082.51     |
| PNA4  | 12W5H-(AEEA)2 | H-(Lys)3-CCTCT-(AEEA)2-TCTCCCTTCTTT-Lys   | 5227.70     |
| PNA5  | 12W5H-(AEEA)3 | H-(Lys)3-CCTCT-(AEEA)3-TCTCCCTTCTTT-Lys   | 5372.80     |
| PNA6  | 12W5H C0-AZO | H-(Lys)3-CCTCT-AZO-TCTCCCTTCTTT-Lys        | 5220.74     |
| PNA7  | 12W5H C3-AZO | H-(Lys)3-CCTCT-C3-AZO-TCTCCCTTCTTT-Lys    | 5277.81     |
| PNA8  | 12W5H C4-AZO | H-(Lys)3-CCTCT-C4-AZO-TCTCCCTTCTTT-Lys     | 5319.89     |
| PNA9  | 12W5H C5-AZO | H-(Lys)3-CCTCT-C5-AZO-TCTCCCTTCTTT-Lys     | 5305.86     |
| PNA10 | 12W5H C6-AZO | H-(Lys)3-CCTCT-C6-AZO-TCTCCCTTCTTT-Lys     | 5306.59     |
| PNA11 | 9W9H C4-AZO | H-(Lys)3-CCCTCT-C4-AZO-TCTCCCTTCTTT-Lys    | 5528.17     |
| PNA12 | 9W8H C4-AZO | H-(Lys)3-CCCTCT-C4-AZO-TCTCCCTTCTTT-Lys    | 5527.81     |
| PNA13 | 9W7H C4-AZO | H-(Lys)3-CCCTCT-C4-AZO-TCTCCCTTCTTT-Lys    | 5510.62     |
| PNA14 | 9W6H C4-AZO | H-(Lys)3-CCCTCT-C4-AZO-TCTCCCTTCTTT-Lys    | 4744.37     |
| PNA15 | 9W5H C4-AZO | H-(Lys)3-CCCTCT-C4-AZO-TCTCCCTTCTTT-Lys    | 4493.07     |
| PNA16 | 9W4H C4-AZO | H-(Lys)3-CCCTCT-C4-AZO-TCTCCCTTCTTT-Lys    | 4241.77     |
| PNA17 | 9W3H C4-AZO | H-(Lys)3-CCCTCT-C4-AZO-TCTCCCTTCTTT-Lys    | 3990.47     |

| DNA1 | Cy3-CATCATCAAAAGGAGGAGATGGTG | 7982.50 |
| DNA2 | Cy3-CATCATCAAAAGGAGGAGATGGTG | 6627.40 |
| DNA3 | Cy3-CATCATCAAAAGGAGGAGATGGTG | 7932.40 |

AEEA: 2-aminoethoxy-2-ethoxy acetic acid, AZO: Thioazobenzene, Lys: Lysine. Underline:Mismatch bases.

2.2. Recognition of Single Strand DNA by PNAs and TC-PNAs

We studied the effect of the linker structures of TC-PNA on binding efficiency to Cy-3 labelled ssDNA (DNA1, Cy3 and linker structure are shown in Figure S1) using gel mobility shift assays. Since Hoogsteen base pairs can be formed under acidic conditions [22], we incubated TC-PNA and Cy3-ssDNA in 10 mM phosphate buffer (pH 6.0) containing 1 mM EDTA. The relative intensities of
shifted bands upon non-denaturing poly-acrylamide gel electrophoresis is a measure of the binding efficiency of TC-PNA to the target ssDNA, while the mobility of each shifted band indicates the composition of the TC-PNA and ssDNA complex.

First, we incubated PNA1–6 and DNA1 in a 1:1 molar ratio according to manufacture provided molar extinction coefficients and analysed their binding characteristics using the gel mobility shift assay. The reaction mixture was added to 30% glycerol and carefully injected in each lane of polyacrylamide gel to avoid the pH change by gel running buffer and immediately subjected to gel mobility shift assay. As a result, single strand PNA (PNA1) binding to DNA1 was observed to be very limited (Figure 2A). However, PNA2, which has C0-AZO at the N-terminus, produced a single distinctly shifted band. This improved binding of PNA2 can be explained by the stacking interaction of AZO with its neighbouring base pairs that increases the duplex stability (Table 2).

![Figure 2](image_url)

*Figure 2.* Analysis of DNA/PNA complexes by gel mobility shift assay. (A) All lanes contain Cy3-labelled ssDNA (DNA1) at 100 nM. Each lane from the left side contains no PNA, 100 nM of PNA1, 2, 6, 3, 4, and 5, respectively. Each PNA and DNA1 were incubated in 10 mM sodium phosphate buffer and 1 mM EDTA (pH 6.0) at 25 °C for 10 min. Gel mobility shift assays of PNA/DNA complexes were performed using 15% non-denaturing polyacrylamide gels; and (B) quantitative band-intensity analysis of DNA and DNA/PNA complexes in Figure 2.

PNA3, 4, and 5 are TC-PNAs that contain (AEEA)1, (AEEA)2, and (AEEA)3 as the linker molecule, respectively. Based on the amount of unbound DNA1 shown in Figure 2A,B, PNA5 exhibited the highest binding affinity to DNA1 among them (shifted band; 66.0%). Corresponding to our results, Kuhn et al. reported that TC-PNA containing three consecutive AEEA linkers gives the most preferable PNA/DNA/PNA triplex with complementary ssDNA [10]. Additionally, PNA3–5 afforded multiple bands with DNA1. A similar phenomenon has also been reported in the case of bisPNA by Hansen et al. and those multiple bands were identified as a mixture of a bisPNA/DNA complex and bisPNA/bisPNA/DNA complexes [19]. PNA6 tethered with AZO linker showed an apparent single shifted band. The rigidity and hydrophobicity of AZO linker only allowed a preferable complex formation. On the other hand, PNA3–5 showed smear shifted bands (Figure 2A). This is probably due to the flexible and water-soluble feature of AEEA linker and, thus, allowed different types of complex formation. PNA3 and 6 that possess shorter linkers gave weaker shifted band (Figure 2B, PNA3; 37.3%, PNA6; 36.5%) than PNA4 and 5 that possess longer AEEA linkers (Figure 2B, PNA4; 53.4%, PNA5; 66.0%). From these results, the linker length for TC-PNA could be important for complex formation with the target ssDNA.

Second, we prepared azobenzene-tethered TC-PNA (PNA6–10) that contained different lengths of azobenzene linker molecules (C0-, C3-, C4- C5-, and C6-AZO, Figure 1C,D) in between two PNA strands (Table 1). PNA6–10 and DNA1 were incubated in a 1:1 molar-ratio for 10 min at room temperature and the mixtures were assessed using a gel mobility shift assay. PNA6, 7, 8, 9, and 10 provided a single shifted band and their relative band intensities were 44.4%, 44.9%, 81.7%, 41.2%,
and 68.4%, respectively (Figure 3A,C). It is noteworthy that the mobility of PNA6 and DNA1 was apparently less than the others (Figure 3A). To study the stoichiometry of TC-PNA and DNA1 in each complex, we added two equivalents of PNA6–10 to DNA1 and analysed them in the same way. Interestingly, PNA6 gave the same single shifted band (85.7%) even with excess amounts of PNA (Figure 3B), while PNA7–10 gave secondary shifted bands at lower mobility positions (Figure 3B, PD complex 1) in addition to higher mobility positions (Figure 3B, PD complex 2). According to our quantitative band-intensity analysis in Figure 3B,D, azobenzene linker molecules for TC-PNA affected their DNA binding properties. Further, PNA8 that possesses C4-AZO linker showed the most efficient binding property to give PD complex 2 among PNA6–10 (Figure 3C; 81.7% at one equivalent of PNA8, Figure 3D; 58.8% at two equivalents of PNA8).

![Shifted bands](image_url)

**Figure 3.** Effect of azobenzene linker length on complex formation between tail-clamp PNA (TC-PNA) and ssDNA. Each lane contains 100 nM of ssDNA. (A) Each lane from the left side contains no PNA, 100 nM of PNA6, 7, 8, 9, and 10; (B) Each lane from the left side contains no PNA, 200 nM of PNA6, 7, 8, 9, and 10, which contain C0-AZO, C3-AZO, C4-AZO, C5-AZO, and C6-AZO as linkers, respectively. Each TC-PNA and ssDNA were incubated in 10 mM sodium phosphate buffer and 1 mM EDTA (pH 6.0) at 25 °C for 10 min. Gel mobility shift assays of TC-PNA/DNA complexes were performed using 15% non-denaturing polyacrylamide gels; (C) quantitative band-intensity analysis of DNA and DNA/PNA complexes in Figure 3A; and (D) quantitative band-intensity analysis of DNA and DNA/PNA complexes in Figure 3B.

### 2.3. Effect of Homopyrimidine Bases of Azobenzene-Tethered PNAs on Complex Formation with Single Stranded DNA

Azobenzene-tethered homopyrimidine TC-PNA may potentially form TC-PNA/DNA/TC-PNA trimers. Therefore, we altered the number of homopyrimidine bases of TC-PNA-C4-AZO and examined their binding properties with homopurine ssDNA by gel mobility shift assay. When two equivalents of PNA11–14 were incubated with DNA1, there were two distinct shifted bands (Figure 4).
As a result, the band intensity of PD complex respectively (Figure 5A,B).

Indeed, the addition of two equivalents of PNA formed not only by Hoogsteen base pairing, but also by Watson-Crick base pairing.

To analyse the composition of TC-PNA and ssDNA complexes, we employed PNA, 8 and 15, since they exhibited a different gel mobility shift pattern as seen in Figure 3A,B and Figure 4. Indeed, the addition of two equivalents of PNA, 8, or 15 to DNA1 produced single bands with a lesser mobility band (61.3%), two shifted bands (81.8% in total), or a greater mobility band (88.4%), respectively (Figure 5A,B).
we observed a set of multiply charged ion peaks at 1845.8 (+10; calcd. 1845.7), 2049.4 (+9; calcd. (Figure S5B, Table S2; Supplemental Data). We also observed another set of multiply-charged ion peaks

Thus, we measured the melting temperature of our representative PNAs and DNA

2.5. Thermal Stability of TC-PNA and ssDNA Complex

By introducing a modest length of azobenzene linker into TC-PNA and reducing the number of homopurine ssDNA. This was also confirmed in the case of ssPNA (Figure S6A: PNA Supplemental data). Side peaks at 1546.8 (+8; calcd. 1544.4), 1767.6 (+7; calcd. 1764.9), and 2062.3 ion peaks at 1565.5 (+8; calcd. 1565.0), 1789.1 (+7; calcd. 1788.2), and 2086.9 (+6; calcd. 2086.4 that agree with the theoretical molecular mass (18,590.1 Da) of a PNA with the theoretical molecular mass (18,590.1 Da) of a PNA

The thermal stability of PNA and DNA gives us some insights into their binding modes of action. From these results, homopyrimidine TC-PNA-AZO potentially forms 2:1 complexes with DNA/PNA complexes were performed using 15% non-denaturing polyacrylamide gels. (B) Quantitative band-intensity analysis of DNA and DNA/PNA complexes in Figure 5.

The composition of the PNA6/DNA1 complex was analysed using nano ESI-MS. As a result, we observed a set of multiply charged ion peaks at 1845.8 (+10; calcd. 1845.7), 2049.4 (+9; calcd. 2050.6) and 2308.8 (+8; calcd. 2308.7 that agree with the theoretical molecular mass (18,447.9 Da) of a PNA6/DNA/PNA15 complex (Figure S5A, Table S1; Supplemental Data). This indicated that PNA6 initially formed Watson-Crick base pairing with DNA1; however, the length of the C0-AZO linker was insufficient to facilitate Hoogsteen base pairing. Thus, a second PNA6 was bound to the 1:1 PNA6/DNA1 complex and afforded a PNA6/DNA1/PNA6 complex.

In the case of PNA8, we observed a set of multiply charged ion peaks at 1899.7 (+7; calcd. 1900.6) that agree with the theoretical molecular mass (13,298.3 Da) of a 1:1 PNA8/DNA1/Na+ complex (Figure S5B, Table S2; Supplemental Data). We also observed another set of multiply-charged ion peaks at 1859.9 (+10; calcd. 1859.9), 2065.5 (+9; calcd. 2066.4) and 2323.5 (+8; calcd. 2324.6) that agree with the theoretical molecular mass (18,590.1 Da) of a PNA8/PNA8/DNA1/Na+ complex (Figure S5B, Table S2; Supplemental data). Taken together with our previous gel mobility shift assay data, the linker substitution of TC-PNA from C0-AZO to C4-AZO facilitates a 1:1 TC-PNA/ssDNA complex formation.

PNA15, which possesses a shorter homopyrimidine strand, produced a set of multiply-charged ion peaks at 1565.5 (+8; calcd. 1565.0), 1789.1 (+7; calcd. 1788.2), and 2086.9 (+6; calcd. 2086.4 that agree with the theoretical molecular mass (12,513.6 Da) of a PNA15/DNA1/Na+ (Figure S5C, Table S3; Supplemental data). Side peaks at 1546.8 (+8; calcd. 1544.4), 1767.6 (+7; calcd. 1764.9), and 2062.3 (+6; calcd. 2058.9), which agree with the theoretical molecular mass (12,348 Da) of the 1:1 complex of PNA15/DNA1/Na+ lacking a lysine residue, were also observed.

From these results, homopyrimidine TC-PNA-AZO potentially forms 2:1 complexes with homopurine ssDNA. This was also confirmed in the case of ssPNA (Figure S6A: PNA1, Figure S6B: PNA2, Tables S4 and S5; Supplemental Data). However, the 2:1 complex formation can be prevented by introducing a modest length of azobenzene linker into TC-PNA and reducing the number of homopyrimidine strands that can form Hoogsteen base pairs.

2.5. Thermal Stability of TC-PNA and ssDNA Complex

The thermal stability of PNA and DNA gives us some insights into their binding modes of action. Thus, we measured the melting temperature of our representative PNAs and DNA1. As a result, PNA2

| PNA | None | 1 | 2 | 6 | 8 | 15 |
|-----|------|---|---|---|---|----|
| DNA | +    | + | + | + | + | + |
| % ratio of shifted band | 0.0 16.9 76.4 61.3 81.1 88.4 |
| % ratio of remaining DNA band | 100.0 83.1 23.6 38.7 18.9 11.6 |

Figure 5. Effect of azobenzene linker length and the number of homopyrimidine bases of PNA on complex formation with complementary ssDNA. (A) Each lane contains 100 nM of DNA1. Each lane from the left side contains no PNA, 200 nM of PNA1, PNA2, PNA6, PNA8, and PNA15. Each DNA and PNA were incubated in 10 mM sodium phosphate buffer (pH 6.0) at 25 °C for 10 min. Gel mobility shift assays of PNA/DNA complexes were performed using 15% non-denaturing polyacrylamide gels.
showed approximately 5 °C higher Tm compared to PNA1/DNA1, probably due to the stacking interaction of azobenzene to the neighbouring bases (Table 2). PNA5, which has three consecutive AEEA linkers and five extended homopyrimidine bases, exhibited 18.3 °C higher Tm compared to PNA1 (Table 2). The three extended AEEA linkers and five homopyrimidine PNA bases contributed to stabilize the TC-PNA/DNA duplex. PNA6 and PNA9, which possess different lengths of azobenzene linkers, both exhibited approximately 11–13 °C higher TC-PNA/DNA duplex stability than PNA1, which was approximately 5–7 °C lower than that of PNA5 (Table 2). Although the azobenzene linker by itself tends to increase the PNA/DNA duplex stability by stacking interactions, the extended homopyrimidine PNA bases might have limited the stacking interaction by forming PNA/DNA base pairing.

Table 2. Melting temperatures between PNAs or TC-PNAs and DNA1.

| PNA Name | Tm  | ∆Tm * | ∆Tm ** |
|----------|-----|-------|--------|
| 12W      | 68.0|       |        |
| 12W-C0-AZO | 72.9| 4.9   |        |
| 12W5H-C0-AZO | 81.5| 13.5  |        |
| 12W5H-C4-AZO | 79.3| 11.3  |        |
| 12W5H-(AEEA)3 | 86.3| 18.3  |        |
| 9W9H-C4-AZO | 79.6| 19.9  |        |
| 9W7H-C4-AZO | 74.3| 14.6  |        |
| 9W5H-C4-AZO | 68.0| 8.3   |        |
| 9W3H-C4-AZO | 59.7| -     |        |

* melting temperature difference relative to PNA1(12W)/DNA1; ** melting temperature difference relative to PNA17(9W3H-C4-AZO)/DNA1.

To study the effect of homopyrimidine bases of TC-PNA-C4-AZO on its interaction with ssDNA, we analysed the Tm of PNA11(9W9H-AZO), 13(9W7H-AZO), 15(9W5H-AZO) and 17(9W3H-AZO) with DNA1. As a result, the Tm of PNA11, 13, 15, and 17 with DNA1 were 19.9 °C, 14.5 °C, and 8.3 °C higher than that of PNA17/DNA1, respectively. This result indicated that the extended homopyrimidine bases contribute to thermally stabilize the duplex in a sequence and length dependent manner. However, the extended homopyrimidine bases may induce a second PNA binding as shown in Figure 4. Optimizing the design of PNA is important not only for increasing its binding affinity, but also eliminating non-specific interactions with the target ssDNA.

2.6. Strand Displacement of dsDNA by TC-PNA-AZO 9W5H-C4

To determine if TC-PNA-C4-AZO can recognize a target sequence within duplex DNA, we prepared dsDNA by annealing 5'-Cy3-labelled target ssDNA (DNA1) and a complementary ssDNA (DNA2, Table 1). DNA1, DNA2, and the duplex DNA were individually incubated with PNA15 for 10 min at room temperature and subjected to gel mobility shift assays. PNA15 bound DNA1 and produced a single shifted band (Figure 6). When PNA15 was incubated with dsDNA, the duplex band intensity was reduced and a new, shifted band appeared on the gel (Figure 6). The mobility of the shifted band was identical to the band observed in the lane containing PNA15 and DNA1. This indicated that PNA15 can recognize the target sequence in dsDNA by forming a complex with DNA1 and eliminating DNA2 from the duplex. Interestingly, PNA15 did not induce multi-complex formation, even when five equivalents of PNA15 were added to DNA1 (Figure S3). Further, PNA15 also recognized mismatched bases in DNA3 (Figure S4).
were removed from the resin by a 10-min incubation in 100 µL of DMF 10 times. The concentration of each monomer (Fmoc-PNA OH Loaded Resin

The Novasyn TGR resin (200 mg, 0.24 mmol/g) was swollen in 5 mL DMF for 30 min prior to the synthesis. To activate the carboxyl groups of coupling monomers, 200 µL of base solution (0.3 M 2,6-lutidine, 0.2 M diisopropyl ethylamine and 0.33 M thiourea solution), and 0.5 M HATU were added to a mixture of Fmoc/Boc-protected lysine (Fmoc-Lys(Boc)-OH, 22.5 mg, 48 µmol) and Boc/Cbz-protected lysine (Boc-Lys(Cbz)-OH, 72.3 mg, 190 µmol) in 200 µL of DMF. For the coupling reaction, the mixture was added to the resin and incubated at ambient temperature for 60 min. The resin was then removed from the reaction solution and washed with DMF (5 × 5 mL). For capping of non-reacted amine groups, the resin was treated with 1.5 mL of capping solution (2,6-lutidine:Ac2O:pyridine = 6:5:89, v/v) for 5 min and washed with DMF (10 × 5 mL). For deprotection of Fmoc-groups, the resin was treated with 1 mL of deblock solution (40% (v/v) piperidine in DMF) for 5 min and washed with DMF (10 × 5 mL).

3.3. PNA Synthesis

Automated linear solid phase synthesis of PNA was performed using an Intavis ResPep parallel synthesizer equipped with micro scale columns (Köln, Germany). The lysine-loaded resin (200 mg, 16 µmol) was swollen in 5 mL DMF for 30 min and 20 mg of the resin was transferred to each column in the synthesizer. After removal of DMF, Fmoc-protecting groups of the lysine-loaded resin were removed from the resin by a 10-min incubation in 100 µL of deblock solution; the resin was subsequently washed with 100 µL of DMF 10 times. The concentration of each monomer (Fmoc-PNA monomers, Fmoc-AEEA-OH, Fmoc-AZO-OH [17] and Fmoc-Lys(Boc)-OH) was adjusted to 0.3 M in DMF solution. The activator solution contained 0.5 M HATU in DMF. The base solution contained

| DNA1 | + | + | + | + |
| DNA2 | - | - | + | + |
| PNA15 | - | + | - | + |

Figure 6. Strand displacement of dsDNA by TC-PNA-C4-AZO. Each lane contains oligonucleotides as indicated above. The concentrations of DNA1 and DNA2 were 100 nM, while PNA15 was 200 nM. Each DNA and PNA were incubated in 10 mM sodium phosphate buffer (pH 6.0) at 25 °C for 10 min. Gel mobility shift assays of PNA/DNA complexes were performed using 15% non-denaturing polyacrylamide gels.

3. Materials and Methods

3.1. Chemicals

Fmoc/Bhoc-protected PNA monomers were purchased from Panagene (Daejeon, Korea). Fmoc-Lys(Boc)-OH, poly-ethylene linker and TGR-resin were purchased from Merck Millipore (Tokyo, Japan). The coupling activators and HATU were purchased from Watanabe Chemicals (Hiroshima, Japan). DNA (salt free) was purchased from Sigma-Genosys (Ishikari, Japan). Other chemicals were purchased from Wako Pure Chemical (Osaka, Japan), Sigma-Aldrich (Tokyo, Japan) and Tokyo Chemical Industry (Tokyo, Japan). Reagents and solvents were used without further purification unless otherwise noted.

3.2. Preparation of Fmoc-Lys-(Boc)-OH Loaded Resin

The Novasyn TGR resin (200 mg, 0.24 mmol/g) was swollen in 5 mL DMF for 30 min prior to the synthesis. To activate the carboxyl groups of coupling monomers, 200 µL of base solution (0.3 M 2,6-lutidine, 0.2 M diisopropyl ethylamine and 0.33 M thiourea solution), and 0.5 M HATU were added to a mixture of Fmoc/Boc-protected lysine (Fmoc-Lys(Boc)-OH, 22.5 mg, 48 µmol) and Boc/Cbz-protected lysine (Boc-Lys(Cbz)-OH, 72.3 mg, 190 µmol) in 200 µL of DMF. For the coupling reaction, the mixture was added to the resin and incubated at ambient temperature for 60 min. The resin was then removed from the reaction solution and washed with DMF (5 × 5 mL). For capping of non-reacted amine groups, the resin was treated with 1.5 mL of capping solution (2,6-lutidine:Ac2O:pyridine = 6:5:89, v/v) for 5 min and washed with DMF (10 × 5 mL). For deprotection of Fmoc-groups, the resin was treated with 1 mL of deblock solution (40% (v/v) piperidine in DMF) for 5 min and washed with DMF (10 × 5 mL).
0.3 M 2,6-lutidine, 0.2 M diisopropyl ethylamine and 0.33 M thiourea in 5% NMM in pyridine solution (v/v). The deblock solution contained 40% pyperidine in DMF solution.

For the coupling reaction, 17.5 µL of monomer solution, 17.0 µL of activation solution and 8.50 µL of base solution were combined in a vessel and incubated for 2 min at ambient temperature. The coupling solution was then transferred to 20 mg of the resin and then incubated for 100 min at ambient temperature. After eluting the coupling solution from the resin by filtration, the resin was washed with 100 µL of DMF 10 times. This coupling procedure was repeated twice for each monomer elongation reaction. The resin was then incubated with 100 µL of capping solution for 10 min to protect the non-elongated amino groups with acetyl groups and subsequently washed with 100 µL of DMF 10 times. The coupling step of the next monomer and capping steps were repeated as described above until the desired PNA molecule was synthesized. Before the cleavage of PNA molecules from the resin, the resin was washed with 100 µL of DMF five times followed by washing with 100 µL of dichloromethane five times. After drying the resin, 1 mL of TFA/m-cresol (9:1, v/v) was added and incubated for 12 h. The resin was then filtered and the flow-through containing the cleaved PNA was transferred to a new tube. The PNA solution was added to 15 mL of ice-cold diethyl ether and the precipitate was collected by centrifugation at 4400 rpm for 4 min. The supernatant was transferred to another tube and discarded. The residue was dried under ambient atmosphere and then dissolved in 100 µL of distilled water.

3.4. PNA Purification and Analysis

All PNAs were purified by reverse-phase HPLC using a JASCO PU-2086 pump system (Tokyo, Japan) with a JASCO UV-2075 detector and a GL Science (Tokyo, Japan) Inertsil (150 mm × 4.6 mm, 5 µm) C-18 column for analytical runs or a GL Science Inertsil (20 mm × 250 mm, 3 µm) C-18 column for semi-preparative runs. Eluting solvents (analytical: A (0.1% TFA in water) and B (0.1% TFA in acetonitrile); semi preparative: A (0.1% TFA in water) and B (0.1% TFA in acetonitrile)) were used in a linear gradient at a flow rate of 1 mL/min for analytical and 5 mL/min for semi-preparative HPLC. The gradient for analytical runs was 0 → 50% B in 30 min and the gradient for semi-preparative runs was 0% B for 10 min, 0 → 5% B in 10 min, 5% B for 10 min, 5 → 10% B in 10 min, 10% B for 10 min, 10 → 20% B in 120 min, and 20 → 50% B in 10 min. Detection was performed using a UV-VIS detector at 260 nm. PNA molecular weights were analysed using an Ultraflextreme MALDI TOF Mass Spectrometer (Bruker Daltonics, Yokohama, Japan) (Figures S7–S23). The optical densities of PNA and DNA were measured at 260 nm with a UV1700 spectrometer (Shimadzu, Kyoto, Japan) using quartz cuvettes (4 × 10 mm). The extinction coefficient of PNA was calculated from the molar extinction coefficient obtained from https://secure.eurogentec.com/EGT/files/FileBrowse/Brochures/Oligonucleotides/PNA-guide.pdf. Measurements of absorption at 260 nm were carried out in a buffer solution (10 mM NaH₂PO₄, pH 7.0) at ambient temperature.

3.5. Gel Mobility Shift Analysis of PNA/DNA Complexes

PNAs were preheated at 95 °C for 5 min to prevent aggregation, then gradually cooled to 25 °C before being added to the DNA solution. DNA concentrations were quantified by absorbance at 260 nm using a molar extinction coefficient provided by manufacturer. PNA/DNA hybridization assays were conducted using 100 nM Cy3-labeled ssDNA with 1–2 equivalents of ssPNA, bisPNA, or TC-PNA in 10 mM sodium phosphate and 1 mM EDTA at pH 6.0 for 10 min at 25 °C. The reaction mixture for each condition was mixed with 0.2 volumes of a solution containing 30% glycerol, 0.025% bromophenol blue, and 0.025% xylencyanol (Sigma) and subjected to electrophoresis at 20 mA for 30 min on a 15% non-denaturing polyacrylamide gel using 1× TBE as a running buffer (89 mM Tris base, 89 mM borate, 2 mM EDTA, pH 8.1) at 4 °C in the dark. The gel images were created by use of a CCD digital image stock system, FAS-III (Toyobo, Osaka, Japan).
3.6. Analysis of TC-PNA/DNA Complexes Using Nano ESI-MS

The TC-PNA/DNA complexes were incubated under the conditions described above. The mixture was diluted and analysed by nano electrospray ionization mass spectrometry (nano ESI-MS) in 100 mM AcONH$_4$ aqueous solution. The experiments were performed on a Q-TOF II mass spectrometer (Micromass, Manchester, UK) in positive ion mode. To minimize complex dissociation, the capillary and cone voltages were optimized to 1.2 kV and 30 V, respectively. The nitrogen counter gas temperature was adjusted to 30 °C or 80 °C. Ar gas was introduced into a collision cell potentiated at 10 V, while the sample solution was introduced into a glass capillary (Proxeon, Thermo Fisher Scientific Inc., Waltham, MA, USA). Mass calibration was performed using cluster ions derived from NaI.

3.7. Thermal Melting Analaysis of PNA/DNA and TC-PNA/DNA

PNAs were preheated at 95 °C for 5 min to prevent aggregation, then gradually cooled to 25 °C before being added to the DNA solution. Melting profiles of PNA complexed with ssDNA were analysed on a UV1700 spectrophotometer (Shimadzu) using a microcell (eight cell, 1 mm) at 260 nm. PNAs and DNA1 were suspended in Na$_2$HPO$_4$ buffer (10 mM, pH 6.0) at 1 µM each. The temperature was ramped down from 95 °C to 10 °C at a rate of −1 °C/min.

4. Conclusions

In conclusion, homopyrimidine TC-PNA consisting of a C4-AZO linker, nine homopyrimidine bases in the N-terminal strand, and five homopyrimidine bases in the C-terminal strand can be used for detecting homopurine DNA target sequences with good binding affinity and sequence specificity. Our study into the design of TC-PNA should expand potential target sites and eliminate non-specific binding to other homopurine target sequences within a given gene.

Supplementary Materials: The following are available online.

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Abbreviations

The following abbreviations are used in this manuscript:

- AEEA: Aminoethoxy-2-ethoxy acetic acid
- BCIP: 5-Bromo-4-chloro-3'-indolyolphatase p-toluidine salt
- Bhoc: Benzhydryloxycarbonyl
- Boc: Tert-butyloxycarbonyl
- CDNA: Complementary DNA
- DMF: N,N-Dimethylformamide
- Fmoc: 9-Fluorenylmethoxycarbonyl
- HBTU: 2-(1H-Benzotriazole-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate
- HOBT: N-Hydroxybenzotriazole
- NMM: N-Methylmorpholine
- TFA: Trifluoroacetic acid
- PNA: peptide nucleic acid
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**Sample Availability**: Samples of the compounds are not available.