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Kyoto University
Development of a visible nanothermometer with a highly emissive 2′-O-methylated guanosine analogue†

Seiji Yamamoto,* Soyoung Park* and Hiroshi Sugiyama*ab

We have synthesized a fluorescent base analogue, 2-aminothieno[3,4-d]pyrimidine based G-mimic deoxyribonucleoside, 2′-OME-thG, and investigated its photophysical properties and DNA incorporation. The 2′-methoxy group of 2′-OME-thG effectively induces the Z-form DNA. Finally, we have constructed a visible nanothermometer based on the B–Z transition of DNA using 2′-OME-thG.

The development of nanodevices and molecular machines such as nanorobots and molecular switches has been a very active research area in the field of nanotechnology. A variety of nanomaterials such as gold nanoparticles, graphene oxide, and mesoporous silica have been investigated, and are now widely utilized to develop nanodevices. DNA is an outstanding natural building block with which to construct two- and three-dimensional nanostructures because of its unique complementarity, chemical stability, and dynamic conformational changes by external stimuli.1–5 Besides natural nucleobases (A, T, C, G), with the rapid advancements in chemical biology, multifarious functionalized nucleobase analogues have been developed, and these artificial genetic alphabet characters widen the sphere of DNA technology applications.6–10 Previously, Tor and coworkers have developed isomorphic fluorescent RNA nucleosides derived from thieno[3,4-d]pyrimidine and demonstrated their attractive photophysical properties including visible light emission and a high quantum yield.11 Recently, we have synthesized a highly emissive deoxyguanosine analogue, thDG, as a fluorescent probe and found that thDG fluorescence could be applied to visualize B to Z transitions of DNA based on different π-stacking of B- and Z-DNA (Fig. 1).12 In this context, we have focused the nucleotide modifications to enhance the utility of thDG and synthesized a highly emissive 2′-O-methylated guanosine analogue, 2′-OMe-thG, and investigated its photophysical properties as a fluorescent base analogue.

2′-O-Methyl-modification of oligonucleotides is a well-known strategy used to increase the binding affinity of oligonucleotides for their target and to enhance the thermal stability of the resulting duplex structure. This methylation also has the advantage that the 2′-O-methyl substituent inhibits the hydrolysis of oligonucleotides in vivo.13–18 Herein, we report the development of a visible nanothermometer by a combination of highly emissive 2′-O-methylated guanosine analogue, 2′-OMe-thG, and distinctive B–Z transition of DNA.

The synthesis of 2′-OMe-thG (3) was achieved based on reported procedures for thDG nucleosides and 2′-O-methyl-modified oligonucleotides (Scheme 1).19a,19e,19f The protected 2′-O-methylated ribonucleoside (2) was obtained through Friedel–Crafts C-glycosylation between thienoguanine (1) and an acylated sugar derivative. This coupling mainly afforded a β-anomer in 83% yield. The benzoyl-protecting groups were removed in a methanolic base and the N,N-dimethylformamide group was introduced for protection of the purine amino group (4). Subsequently, the 5′- hydroxyl group was protected...
with the dimethoxytrityl ether (DMTr) and the desired β-anomer (5) could be isolated in 75% yield. The configuration at the C-1 carbons of the β-anomer was confirmed by 1D and 2D (NOESY) 
\( ^1\)H NMR experiments (see ESI†).

To evaluate oligonucleotides containing the 2′-O-methylated guanosine analogue, the phosphoramidite 2′-O-Me-\( ^{th}\)G (6) was synthesized and incorporated into the center of 18-mer DNA oligonucleotides of 5′-(CGTCCGTCXACGCACGC)-3′, where X = 2′-O-Me-\( ^{th}\)G, by automated solid-phase synthesis. The complementary strands of ODN1 containing matched or mismatched bases and the corresponding native DNA duplexes with G were also prepared. The 2′-O-Me-\( ^{th}\)G-C base pair showed almost identical thermal stability (\( T_m = 72.1\) °C) compared with native duplex DNA with a G-C base pair (\( T_m = 72.1\) °C), as shown in Fig. 2. The complementary strands containing mismatched bases (ODN4–7) decreased the melting temperature compared with one obtained using the complementary strands containing dG. The thermodynamic stability and base pairing selectivity indicate that 2′-O-Me-\( ^{th}\)G could replace a G base in the strand without structural disruption. The photophysical properties of 2′-O-Me-\( ^{th}\)G monomer were also investigated (see ESI†).

The fluorescence of 2′-O-Me-\( ^{th}\)G ribonucleoside (1) shows absorption at 320 nm and visible emission at 457 nm with a high quantum yield of 0.652 under neutral conditions in water.

We prepared a self-complementary decamer 5′-d(CGXCGGCGG)-3′ (ODN8), where X = 2′-O-Me-\( ^{th}\)G, and examined its conformation using circular dichroism (CD) spectra. Fig. 3a shows the CD spectra of ODN8 in various concentrations of NaClO4 at 5 °C. At 1–3 M NaClO4, a negative Cotton effect at around 250 nm and a positive Cotton effect at around 285 nm were observed; ODN8 maintained the B-form. When we increased the NaClO4 concentrations from 5 to 9 M, we observed a positive Cotton effect at around 260 nm and a negative Cotton effect at around 290 nm; 2′-O-Me-\( ^{th}\)G-containing decamer duplex converted to Z conformation. In a recent study, we demonstrated that the B–Z transition could be visualized by the fluorescence intensity of \( ^{th}\)dG.\(^{12}\) Although \( ^{th}\)dG indicated the comparable resemblance to the native dG nucleosides regarding thermodynamic stability and base pairing selectivity, the B–Z transition became more difficult when \( ^{th}\)dG was incorporated as a replacement for a dG nucleotide. Because \( ^{th}\)dG favors the anti-conformation to stabilize B-form DNA, 8-methylguanine (m8G) was additionally introduced into DNA sequences as a Z-stabilizing unit.\(^{28}\) Fortunately, the obtained results indicate that the oligonucleotide possessing 2′-O-Me-\( ^{th}\)G could convert Z conformation without the aid of a Z-DNA inducer. Subsequently, we observed the change in fluorescent intensity by B–Z transition of ODN8. As shown in Fig. 3b, the fluorescence of ODN8 increased significantly with increasing NaClO4 concentration. This result indicated that strong fluorescence enhancement was observed in Z-DNA compared with B-DNA.

Previously, we have found that B–Z transition can be controlled by temperature and high salt conditions, and demonstrated a DNA-based switching device that responds to
Based on previous studies and the photo-physical properties of 2'-OMe-thG, we devised a visible thermometer. To test this concept, the 2'-OMe-thG-containing oligonucleotide 5'-d(CGCGCXXCGCGC)-3' (ODN9), where X = 2'-OMe-thG, was prepared and conformational changes by temperature were investigated in 3.5 M NaClO4. At 5 °C, ODN9 predominantly converted to Z-DNA because of its lower entropy. As the temperature increased from 5 °C to 40 °C, the proportion of B-DNA gradually increased (Fig. 4a and c). We ascertained that the equilibrium between B-DNA and Z-DNA of ODN9 could be controlled by temperature. Therefore, the fluorescence of ODN9 was observed at different temperatures in 3.5 M NaClO4. At 5 °C, ODN9 predominantly converted to Z-DNA because of its lower entropy. As the temperature increased from 5 °C to 40 °C, the proportion of B-DNA gradually increased (Fig. 4a and c). We ascertained that the equilibrium between B-DNA and Z-DNA of ODN9 could be controlled by temperature. Therefore, the fluorescence of ODN9 was observed at different temperatures in 3.5 M NaClO4. To our delight, the fluorescence intensity of ODN9 changed depending on the temperature in conjunction with its B–Z transition. As shown in Fig. 3b, very strong fluorescence enhancement was observed at 5 °C, whereas the fluorescence intensity of ODN9 decreased significantly at 40 °C. The proportions of Z-DNA, B-DNA, and single-strand DNA are shown in the ESI (Fig. S6†). To examine the utility of 2'-OMe-thG-containing oligonucleotide as a nanothermometer, we monitored the fluorescence of ODN9 under a repetitive temperature cycle between low temperature (5 °C) and high temperature (40 °C). Consequently, a reproducible fluorescence of ODN9 was observed according to the change in the proportion of Z- and B-DNA (Fig. 4c). As shown in Fig. 4d, the distinguishable blue
emission of the nanothermometer at 5 °C was visible to the naked eye.

For further investigation of the devised nano-thermometer we prepared two 2′-OMe-thG-containing oligonucleotide 5′-d(CGCCXCCGCG)-3′ (ODN10), where X = 2′-OMe-thG, and evaluated it by CD spectra and fluorescent spectra measurements. Although ODN10 did not indicate dramatic B→Z transition in the range of 0 M to 5 M NaClO4, fluorescent intensity strongly increased with increasing NaClO4 concentration from 7 M to 9 M (Fig. S7†). Therefore, CD and emission spectra of ODN10 were measured in 7 M NaClO4 at various temperatures. Interestingly, ODN10 indicated that the population of Z-form strongly increased with increasing NaClO4 concentration from 5 °C to 35 °C and the emission increased in line with the proportion of Z conformation (Fig. 5); the thermal response of ODN10 was in inverse to that of ODN9. It is known that Z-RNA is the more stable at higher temperature.21,22 This result suggests that four 2′-OMe group in duplex ODN10 may induce RNA-like thermal conformational change and we can reverse the response to the same stimuli by control the incorporation of 2′-OMe-thG.

In conclusion, we synthesized a useful fluorescent guanine analogue, 2′-OMe-thG, and developed a visible nanothermometer using a combination of distinctive B→Z transition of DNA and robust brightness of 2′-OMe-thG in the visible region. Temperature is a critical factor influencing various biochemical transformations in living systems.23 A visible nanothermometer based on a nucleic acid may be a useful tool for the development of biocompatible nanodevices to monitor the temperature in an intracellular environment.24–29 Further investigations into the application of 2′-OMe-thG are ongoing in our laboratory.

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Notes and references


