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A pyrene-imidazolium derivative that selectively Recognizes G-Quadruplex DNA

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ABSTRACT

G-quadruplexes, formed of four stranded guanine bases stabilized by monovalent cations, serve important role in cancer cell growth and control gene expression in telomere. Since there are various types of quadruplex structures, rapid and simple screening methods with high selectivity, sensitivity and nontoxicity are required for understanding about the biological roles of quadruplex DNA as well as in designing therapeutics. Herein, we report a pyrene-imidazolium derivative, **JY-1**, which can with high selectivity recognize G-quadruplex using fluorescence and NMR spectroscopy. This is the first example based on the imidazolium derivative, which can detect the G-quadruplex directly to utilize the excimer/monomer emission change in pyrene fluorophore. The selectivity of strong binding to a specific sequence can allow for quadruplex sensing and the detection method presented here is very simple, using fluorescence and NMR study. Also, the groove binding characteristic of **JY-1** to the G-quadruplex has a relatively low nonspecific toxicity and the structure-specific differences in fluorescent character between DNA duplex and G-quadruplex may offer more discovery and application in biological study. © 2011 Elsevier Ltd. All rights reserved.

1. Introduction

For many years, the development of DNA sensing and recognition technology has attracted a great deal of attention since various diseases and genetic disorders have been found to be directly related to changes in DNA [1–4]. Among the various types of secondary structures in human DNA, such as B-DNA, Z-DNA, hair pins, *etc* [5]., guanine rich sequences like, G-quadruplexes serve an important role in cancer cell growth and control gene expression in telomeres [6]. G-quadruplexes are composed of four stranded guanine bases stabilized by monovalent cations such as Na⁺ or K⁺ between tetrad guanine bases [7]. Recent studies show that this unique structure can be formed not only in the telomeric region of the genome, but also among other guanine rich sequences having the potential to form quadruplex elsewhere in the human genome [8,9]. Efficient detection of G-quadruplex in addition to stabilization of the G-quadruplex structure is very helpful in designing cancer drugs and in diagnosis of various diseases [10,11]. Detection method taking advantage of fluorescence is so much simple and sensitive [12–15] that there have been reports on fluorophore labelled G-quadruplex oligonucleotide [16–18] or label free probe systems for sensing G-quardruplex [19-23]. However, these studies mainly set out to probe potassium ion stabilizing in Gquartet structure, or detect the conformational change of the Gquartet. Some studies were performed using a labelled oligonucleotide system, which is expensive and laborious. In addition, the intercalative methods for detection of the quadruplex can be cytotoxic with relatively low selectivity and can induce the destabilization of the quadruplex within telomeres, which is critical for telomerase activity inhibition [6]. Since there are various types of quadruplex structures, rapid and simple screening methods with high selectivity, sensitivity and nontoxicity are required for understanding the biological roles of quadruplex DNA as well as for designing therapeutics [24,25].

Herein, we report a pyrene-imidazolium derivative which can with high selectivity recognize G-quadruplex. Imidazolium based receptors have been investigated for the reconition of various anionic species [26–35]. We have synthesized a quadruplex selective fluorescent probe, **JY-1** (Fig. 1A), designed on the structural combination of a highly effective binding with DNA by using





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Fig. 1. (a) Chemical structure of pyrene derivatives, JY-1. (b-c) Fluorescence emission spectra of JY-1 complexed with (b) DNA-1 or (c) DNA-2 duplexes in the aqueous solution containing 10 mm sodium phosphate (pH = 7.0) at various D/P ratios. (d) Fluorescence emission spectra of JY-1 complexed with DNA-3 (D/P = 1.0) in the aqueous solution containing 10 mM HEPES buffer (pH = 7.4) with various KCl concentrations. (Control : JY-1 only, D: DNA, P: JY-1).

imidazolium unit, and inducing distinctive fluorescent emission changes. In order to account for the selective recognition of the JY-1 to G-quartet DNA, we have performed NMR and fluorecence study on the **JY-1** complexed with various DNA molecules (see Table 1).

2. Materials and methods

2.1. Synthesis

Compound 2: The suspension of 1-pyrenemethanol 4 (2 g, 8.6 mmol) in toluene (100 mL) was cooled to 0 °C followed by addition of phosphorus tribromide (1 mL, 10.5 mmol) via syringe. The mixture was stirred at 0 °C for 1 h and then warmed to room temperature, during which the reaction became homogeneous. Saturated Na₂CO₃ solution 50 mL was added slowly and the reaction was stirred until it cooled to room temperature. The phases were separated, and the organic phase was washed with H_2O (50 mL \times 2), brine (50 mL \times 2) and dried over Mg₂SO₄. The yellow filtrate was concentrated to minium volume. The yellow needle-like solid was collected and dried. The mother liquid was concentrated again and repeated the crystallization process. The total product was 2.3 g in 91% yield. m.p. = 136 °C ¹H-NMR (CDCl₃, 250 MHz) δ 5.23 (s, 2H), 8.02 (m, 5H), 8.21 (m, 3H), 8.35 (d, J = 9.3 Hz, 1H). ¹³C-NMR (CDCl₃, 62.5 MHz) & 32.28, 122.80, 124.58, 124.84, 125.07, 125.61, 126.26, 127.32, 127.67, 128.01, 128.22, 129.03, 130.51, 130.73, 131.17, 131.92.

Compound JY-1:A solution of 1-bromomethylpyrene 2 (0.66 g, 2.24 mmol) and bisimidazole 3 (0.15 g, 1 mmol) in 160 mL acetonitrile was refluxed for 24 h under argon. After cooling to the room temperature, the precipitate was filtered and washed with ether. The bromide salt (694 mg, 93%) was dissolved in 25 mL DMF. (During the dropwise addition of saturated aqueous KPF₆ solution, precipitate was formed. After washing the precipitate several times with water, desired product was obtained as a white solid (663 mg, 81%). ¹H-NMR (DMSO, 250 MHz) (for bromide salt) & 6.31 (s, 4H), 6.58 (s, 2H), 8.02 (s, 2H), 8.03 (s, 2H), 8.08-8.52 (m, 18 H), 9.51 (s, 2H). ¹³C-NMR (DMSO, 62.5 MHz) & 50.39, 58.29, 122.36, 123.53, 123.56, 124.04, 125.17, 125.83, 125.93, 126.06, 126.54, 126.71, 127.19, 128.29, 128.72, 128.84, 130.06, 130.61, 131.61, 137.62.; HRMS (FAB) calcd for C41H30F6N4P [M-PF6]+ 723.2101, found 723.2111.

2.2. Fluorescence study

Oligonucleotides were purchased from Bioneer(South Korea) and dissolved in 10 mM phosphate buffer, containing 100 mM NaCl, 0.1 mM EDTA, pH 7.0. This buffer solution was used during the whole fluorescent experiments except for DNA-3 study with KCl. The compound (1 mM) stock in CH₃CN was prepared and the final test samples were made up in 1% CH₃CN and 99% buffer solution. For all measurements, excitation was at 343 nm. Both excitation and emission slit widths were 3 nm or 5 nm. Fluorescence emission spectra were obtained using RF-5301/PC Spectrofluorophotometer (Shimadzu).

2.3. Circular dichroism (CD) measurement

The CD spectra were recorded on a JASCO J-810 spectropolarimeter. Solutions containing the probe and oligonucleotides were placed in a quartz cell (1 cm path length), and the spectra were recorded in the 190-450 nm region. The parameters of measurement were 1 nm bandwidth, standard sensitivity and response time of 4 s. Each sample was scanned 3 times and the averages were obtained. The baseline was corrected from the buffer solution containing 1% CH₃CN.

L	al	bl	е	1

Sequence contexts of DNA	studied here([a]: Single-stranded DN	IA).

Name	Sequence	Туре
DNA-1	d(ACCCACCCAA)/d(TTGGGTGGGT)	duplex
AC-10	d(ACCCACCCAA)	ssDNA[a]
TG-10	d(TTGGGTGGGT)	ssDNA
DNA-2	d(GTCATGGTTA)/d(TAACCATAGC)	duplex
DNA-3	d(GGTTGGTGTGGTTGG)	G-quartet
GGG	d(GGG)	ssDNA trimer
TGG	d(TGG)	ssDNA trimer
GTG	d(GTG)	ssDNA trimer
GGT	d(GGT)	ssDNA trimer
TTG	d(TTG)	ssDNA trimer

2.4. NMR experiment

All DNA and **JY-1** samples were dissolved in 60% H₂O/10% D₂O/30% CD₃CN buffer containing 5 m_M sodium phosphate (pH 8.0) and 50 m_M NaCl. All NMR experiments were performed on a Varian Inova 600 MHz spectrometer (KAIST, Daejeon) using a HCN triple-resonance probe. One dimensional (1D) NMR data were processed with either the program VMRR J (Varian, Palo Alto) or FELIX2004 (Accelrys, San Diego), whereas 2D data were processed with the program NMRPIPE and analyzed with the program Sparky.

3. Results and discussion

3.1. Design, synthesis, and fluorescence study of compound JY-1

For the synthesis of **JY-1** (Fig. 1A), 1-bromomethylpyrene (2) was prepared by treating 1-pyrenemethanol (4) with phosphorous tribromide in toluene (see Scheme 1). 1-bromomethylpyrene was then reacted with bisimidazole (3) followed by anion-exchange with KPF₆, yielding 81% **JY-1** (see Scheme 1). Fig. 1 shows the fluorescence emission spectra of **JY-1** with various DNA in aqueous solution. In the absence of DNA, **JY-1** shows the distinctive spectra with the monomer emission and excimer emission at 375 and 482 nm, respectively. There were no significant changes in the emission spectra of JY-1 when 10 equivalents of various phosphate and nucleotide anions were added (Supporting information S-Figure 1). Surprisingly, **JY-1** displayed fluorescent enhancement of the excimer peak (482 nm) upon the addition of DNA-1 duplex even at the DNA/**JY-1** molar ratio (D/P ratio) = 0.1 in 10 mM sodium phosphate buffer (Fig. 1B). However, when DNA-2, random DNA decamer duplex, was added, no significant enhancement of the excimer fluorescence peak at 482 nm was observed even with the D/P ratio of 20 (Fig. 1C). On the other hand, the intensity of monomer peak increased upon interaction with DNA-2, as the D/P ratio increased (Fig. 1C). These data indicate that **JY-1** binds with DNA-1 through unusual molecular interaction unlike DNA-2.

3.2. NMR studies for prove the interaction between **JY-1** and DNA complexes

To understand this unusual interaction, we performed ¹H-NMR experiments on the **JY-1**/DNA-1 complex. The assignment for the proton resonances of the **JY-1** were made by the analysis of NOESY spectra at 25 °C (see Fig. 2). Fig. 2A shows the titration of the DNA-1 duplex with **JY-1**, where the imino proton resonances extensively broadened as **JY-1** was added and then completely disappeared at the D/P ratio = 1.0. ¹H-NMR spectra of **JY-1** were also dramatically changed by addition of the DNA-1 duplex, indicating specific interaction between DNA-1 and **JY-1** (Fig. 2B). These results clearly demonstrated that the double helix of DNA-1 was destabilized upon interaction with **JY-1** and subsequently melted out to single strands. To further clarify this interaction, the titrations of **JY-1** with each strand of DNA-1 (TG-10 and AC-10) were monitored by 1D NMR spectra (Fig. 2C and D). Interestingly, like DNA-1, the TG-10 strand caused a significant change in the NMR spectra of **JY-1**

(Fig. 2B). Even though the D/P ratio is only 0.1, the NMR spectra of these complexes were completely different from that of free **IY-1** (Fig. 2D). For example, at D/P = 0.1, the peaks a and e are disappeared upon binding to the TG-10 strand and new four resonances observed at that region (Fig. 2C). In the free JY-1, each methylene moiety which connect between two imidazolium rings (peak a) or between imidazolium and pyren rings (peak e) shows only one signal in the proton NMR spectrum. However, in the case of the JY-1/TG-10 complex, the methylene protons positioned at a and e exhibit more than two resonances because the pyren and imidazolium rings cannot rotate freely any more. In the case of AC-10, no clear change in the NMR spectra of **JY-1** was observed, as the D/P ratio increased up to 1.0 (Fig. 2D). The slight upfield shifts of some resonances of **JY-1**, such as a resonance at 6.3 ppm (peak e), imply a nonspecific interaction between the imidazolium cation of **IY-1** and the phosphate anion of AC-10 (Fig. 2C). From these results, the molecular interaction between JY-1 and DNA-1 duplex can be summarized as follows: i) **JY-1** destabilizes the double helix of DNA-1 and then separates it into two single strands, TG-10 and AC-10; *ii*) JY-1 selectively binds to the single-stranded TG-10 and exhibits a unique conformation in the complex; iii) JY-1 in this unique conformation can emit the excimer fluorescence.

3.3. NMR study of compound **JY-1** for selective GGG trimer detection

The single-stranded TG-10 has a G-rich sequence and can form G-quartet DNA helix under KCl buffer conditions. To address the correlation between the G-quartet DNA structure and the excimer fluorescence of **JY-1**, the fluorescence study on the TG-rich 15mer, DNA-3, which is known as a thrombin-binding aptamer and can form the G-quartet structure with K⁺ ion [12], was performed. Here, the HEPES buffer (pH 7.4) was used instead of 10 mM sodium phosphate buffer containing 100 mM NaCl to exclude the Na⁺ ion effect because some metal cations including K⁺, Na⁺, or Pb²⁺ can stabilize the G-quartet structure [36,37]. Surprisingly, **JY-1** showed significant excimer fluorescence as the KCl concentration increased (Fig. 1D). From these results, we can suggest the hypothesis that the excimer fluorescence is a unique characteristic of **JY-1** complexed with G-quartet DNA.

In order to verify this hypothesis, we performed the fluorescence and NMR experiments on the **JY-1** complexed with five DNA trimers (TTG, TGG, GGG, GGT, and GTG), which are designed from cutting the TG-10 into trimers and GG dimer (Fig. 3). This approach provides the information for the minimum sequence requirement for DNA binding of the **JY-1** molecule. Only GGG trimer induced the large enhancement of excimer emission from **JY-1**, whereas other trimers and GG dimer did not induce the excimer emission (Fig. 3A). These results indicate that there is a unique interaction between **JY-1** and GGG trimer sequence. To prove the interaction, the NMR study of **JY-1** with GGG and TTG trimers was also performed and then compared with each other (Fig. 3C and D). Fig. 3C shows the titration of the GGG trimer to **JY-1** dissolved in the NMR



Scheme 1. Synthesis of JY-1.



Fig. 2. (A) 1D ¹H-NMR spectra of free DNA-1 (top) and DNA-1–**JY-1** complexes in 60% H₂O/10% D₂O/30% CD₃CN buffer containing 5 mM sodium phosphate (pH 8.0) and 50 mM NaCl at 25 °C. The **JY-1**/DNA-1 ratios (inverse D/P ratios) are shown on the left of each spectrum. 1D ¹H-NMR spectra of **JY-1** in 70% D₂O/30% CD₃CN buffer containing 5 mM sodium phosphate (pH 8.0) and 50 mM NaCl at 25 °C upon titration with (B) DNA-1 duplex, (C) TG-10, or (D) TG-10. 1D ¹H-NMR spectra of free DNA-1, TG-10 and AC-10 are shown on the top of the spectra.

buffer (60% H₂O/10% D₂O/30% CD₃CN, 5 mM sodium phosphate (pH = 8.0) and 50 mM NaCl). Like TG-10, at D/P = 1.5, the peaks a and e are disappeared upon binding to the GGG trimer and new resonances are observed at that region (Fig. 3C). This data indicates that the **JY-1** binds to the GGG trimer with similar binding mode to TG-10 strand. Thus, NMR structural study of the **JY-1**/GGG complex can provide structural information about the general interaction between **JY-1** and G-rich DNA strand. Surprisingly, six sharp peaks at 10–11.5 ppm, corresponding to guanine imino protons of G-quartet structure, appeared as the amount of GGG increased (Fig. 3C). Under this condition, free GGG showed no imino proton resonances (Fig. 3B). These results indicate that, under our experimental condition, free GGG did not form G-quartet structure but **JY-1** interacts with unstructured GGG and induces the G-quartet

structure of the GGG trimer, which shows some imino resonances at 10–11.5 ppm. When TTG was added to **JY-1**, this type of imino proton resonance was not observed, indicating that **JY-1** could not induce the G-quartet structure in the case of the TTG trimer (Fig. 3D).

3.4. Structural binding model studies

In order to indentify the suitable geometry of the induced G-quatet structure, the NOESY experiments on the **JY-1**–GGG complex were performed. Six imino peaks of this complex indicate that four molecules of GGG form two-fold symmetric G-quartet structure. The non-exchangeable base (H8) and sugar (H1', H2', H2'') protons of GGG in the complex were assigned according to their



Fig. 3. (A) Fluorescence emission spectra of **JY-1** complexed with DNA trimers (TTG, TGG, GGG, GGT, or GTG) or GG dimer in the aqueous solution containing 10 mm sodium phosphate (pH = 7.0) at D/P = 1.0. (B) ¹H-NMR spectrum of free GGG in 60% H₂O/10% D₂O/30% CD₃CN buffer containing 5 mm sodium phosphate (pH = 8.0) and 50 mm NaCl at 25 °C 1D ¹H-NMR spectra of **JY-1** in 60% H₂O/10% D₂O/30% CD₃CN buffer containing 5 mm sodium phosphate (pH = 8.0) and 50 mm NaCl at 25 °C 1D ¹H-NMR spectra of **JY-1** in 60% H₂O/10% D₂O/30% CD₃CN buffer containing 5 mm sodium phosphate (pH = 8.0) and 50 mm NaCl at 25 °C upon titration with the (C) GGG or (D) TTG trimers. The D/P ratios are shown on the left of spectra. The part of spectrum in which imino proton region are expaned to emphasize the six imino proton peaks are added on top of Fig. 3C.



Fig. 4. (A) Schematic structure of tetrameric parallel G-quartet of GGG. (B) Expanded regions (imino-to-base: upper; imino-to-amino: lower) of 2D Watergate NOESY spectra of the **JY-1**–GGG complex (D/P = 1.0) in 60% H₂O/10% D₂O/30% CD₃CN buffer containing 5 mM sodium phosphate (pH 8.0) and 50 mM NaCl at 10 °C. Sequential imino-to-imnio connectivities are represented with red lines. (C) Specific imino-H8 connectivity pattern around a G-tetrad (G•3G•G3•G6) indicated with magenta lines. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



Fig. 5. Representative model for the JY-1 complexed with (A) parallel G-quartet of GGG and (B) B-form DNA. The JY-1 atoms are green. Positive charged nitrogen atoms of JY-1 are blue. The sugar and base atoms of DNA are yellow and white, respectively. Phosphate backbone anions are red. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

intra-residue and sequential NOE connectivities (S-Figure 2). The exchangeable protons were assigned according to their sequential imino-to-imino connectivities and confirmed by intra-residue strong imino-to-amino and weak imino-to-H8 NOE cross peaks in the NOESY spectra (Fig. 4). The chemical shifts of the proton resonances of the GGG in the **JY-1**/GGG complex are given in S-Table 1. The pattern of the imino proton resonances of this complex is very similar to that of the U6 variant of human telomeric sequence. d(TAGGGUTAGGGT), which exhibits dimeric parallel G-quartet structure [38]. The two-fold symmetric parrallel G-quartet structure of GGG satisfied well the sequential imino-to-imino connectivities and unique imino-to-H8 NOE data (Fig. 4). However, none of the models for anti-parrallel G-quartet structures could not explain the NMR NOESY data we observed. Thus, we suggested that **JY-1** is thought to be not just the G-quartet structure-recognizing seonsor but the "G-quartet inducer" which is interacts with the G-rich DNA region and then produces the G-quartet structure.

The main intermolecular interaction in the **JY-1**–DNA complex is electrostatic interaction between the imidazolium moiety of JY-1 and phosphate anion of DNA. When two imidazolium cations maintain the stacked structure of two pyrene moieties during interaction with DNA, JY-1 shows the excimer fluorescence. However, when the stacked structure of two pyrene moieties is disrupted to make two imidazolium-DNA interactions, the excimer fluorescence disappears. Fig. 5A shows the possible model structure of the parrallel G-quartet DNA complexed with the **JY-1** that has the stacked two pyrene moieties. Surprisingly, two imidazolium groups could interact with the G1-p-G2 phosphate anion of one strand and with the G5-p-G6 phosphate anion of next strand. respectively (Fig. 5A). However, in the case of normal B-DNA, the phosphate anions across the minor groove cannot maintain the stacked structure of two pyrene moieties through electrostatic interaction with imidazolium groups of JY-1 (Fig. 5B). This representive model could suggest that JY-1 binds to GGG sequences on the groove of G-quartet tightly and that this interaction induces excimer emission enhancement of JY-1. The binding constant was calculated as $5.5\times10^5\,M^{-1}$ (S-Figure 3) and the excimer emission was shown visually in S-Figure 4.

3.5. Circular dichroism and fluorescent change measurement

Circular dichroism (CD) was also used to consider the binding mechanism between JY-1 and DNAs [39,40]. DNA-3, known as thrombin-binding aptamer, forms the antiparallel G-quadruplex structure in the presence of K⁺ ion and exhibits a negative band near 265 nm and a positive band at 290 nm, respectively [41]. As the concentration of **JY-1** increased, there was no appreciable distortion of DNA-3 in the presence of 6 mM KCl, and no induced circular dichroism (ICD) was observed (Fig. 6C). DNA-1 and DNA-2 duplex complexed with IY-1 showed lowered CD at 280 nm which is related with DNA helix unwinding [42]. We noticed DNA-1 complexed with JY-1 did not show ICD at the UV absorption range of JY-1, but only DNA-2 duplex with JY-1 showed a weak negative ICD from the pyrene absorption region (Fig. 6A and B). In the presence of DNA duplex, negative ICD band indicates that JY-1 could destabilize the DNA duplexes. These CD features are consistent with the excimer and monomer emission differences between three DNA duplexes complexed with JY-1 in the fluorescence study, and no ICD of DNA-3 with **JY-1** supports the groove binding interaction.

To further study the binding mode of **JY-1** with DNA oligonucleotides, we examined the fluorescent changes of **JY-1** with the anionic quencher $Fe(CN)_6^{4-}$. If the binding of **JY-1** involves intercalation to the DNA complex, the fluorescent quenching by $Fe(CN)_6^{4-}$ should be minimally or not effective because the negative charge of the phosphate group creates an electrostatic barrier, so the



Fig. 6. CD spectra changes of (A) DNA-1 duplex and (B) DNA-2 duplex (6 μ M) with various concentration of **JY-1** in 10 mM sodium phosphate buffer (pH 7.0). (C) CD spectra of DNA-3 (6 μ M) complexed with various concentration of **JY-1** in HEPES buffer containing 6 mM KCl. DNA/compound **JY-1** ratios ranged from 0.1 to 2 for each titration. (Control : DNA only, D: DNA, P: **JY-1**).

quencher can't approach the **JY-1** [43]. As shown in S-Figure 5, the anionic quencher Fe(CN) $\frac{6}{6}^{-}$ caused a decrease of excimer emission by **JY-1** complexed with DNA-1 duplex (at D/P = 0.1) and this result also confirms that the excimer emission of **JY-1** was induced by the outside binding of DNA complexes.

4. Conclusions

We have described a quadruplex selective fluorescent probe, **JY-1**, designed on the structural combination of highly effective binding ability with DNA as well as inducing distinctive fluorescent emission changes. The selectivity of strong binding to a specific sequence can allow for quadruplex sensing and the detection method presented here is very simple, using fluorescence and NMR study. We found that **JY-1** is thought to be the "G-quartet inducer" which is interacts with the G-rich DNA region and then produces the G-quartet structure, contrast to other sensors recognizing the G-quartet DNA structure. Also, the groove binding characteristic of **JY-1** to the G-quadruplex has a relatively low nonspecific toxicity and the structure-specific differences in fluorescent character between DNA duplex and G-quadruplex may offer more discovery and application in biological study.

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Appendix. Supplementary material

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.biomaterials.2011.11.073

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