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## NMR analysis of <sup>15</sup>N-labeled naphthyridine carbamate dimer (NCD) to contiguous CGG/CGG units in DNA<sup>+</sup>

Takeshi Yamada, ();<sup>a</sup> Shuhei Sakurabayashi,<sup>a</sup> Noriaki Sugiura,<sup>a</sup> Hitoshi Haneoka<sup>b</sup> and Kazuhiko Nakatani ()\*<sup>a</sup>

The structure of the complex formed by naphthyridine carbamate dimer (NCD) binding to CGG repeat sequences in DNA, associated with fragile X syndrome, has been elucidated using <sup>15</sup>N-labeled NCD and <sup>1</sup>H-<sup>15</sup>N HSQC. In a fully saturated state, two NCD molecules consistently bind to each CGG/CGG unit, maintaining a 1:2 binding stoichiometry.

The aberrant expansion of trinucleotide repeat (TNR) sequences in the human genome leads to over 40 serious neurodegenerative diseases, exemplified by fragile X syndrome (FXS, caused by CGG repeat), Huntington's disease (HD, CAG), and myotonic dystrophy type 1 (DM1, CTG).<sup>1-3</sup> Unfortunately, there is currently no effective therapy for TNR diseases. This genomic instability results in both contraction and expansion of the repeats.<sup>4–6</sup> Aberrantly expanded repeats tend to form metastable slip-outs, comprising hairpin structures with repetitive CXG/CXG units, where X-X mismatches are flanked by two C-G base pairs during replication and transcription reactions.<sup>7</sup> The chemical stability of the slip-outs is a contributing factor to the genomic instability of TNR. Recent studies from our group have demonstrated that perturbing slipout structures with small external molecules can modulate the chemical stability of the hairpin structure and, consequently, genomic instability,<sup>8,9</sup> highlighting the importance of developing such small molecules and elucidating their binding modes.<sup>10–13</sup>

We have reported a variety of small molecules binding to TNR sequences, represented by naphthyridine carbamate dimer (NCD, Fig. 1A),<sup>14</sup> which showed a remarkable binding to the CGG repeat DNA. Recently, we reported the NMR structural

<sup>a</sup> Department of Regulatory Bioorganic Chemistry, SANKEN, Osaka University, Mihogaoka 8-1, Ibaraki, Osaka 567-0047, Japan. determination of the complex comprised of NCD and a model DNA containing a single CGG/CGG triad<sup>15</sup> (Fig. 1B, PDB: 7YVW). The NMR structure elucidated the distinctive binding pattern of NCD. Two NCD molecules bind to the CGG/CGG triad, and the



**Fig. 1** (A) Chemical structures of NCD and <sup>15</sup>N-NCD. The partial numberings of guanine and naphthyridine rings are shown in red and green, respectively. (B) Illustration of complexes composed of two NCD molecules and a CGG/CGG unit, as elucidated by NMR structural determination (ref. 15). The proposed structures of complexes involving NCD molecules and consecutive (C) two or (D) three repetitive CGG/CGG units. 'NC' refers to the naphthyridine carbamate moiety of NCD.

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E-mail: nakatani@sanken.osaka-u.ac.jp

<sup>&</sup>lt;sup>b</sup> Comprehensive Analysis Center, SANKEN, Osaka University, Mihogaoka 8-1, Ibaraki, Osaka 567-0047, Japan

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<sup>‡</sup> Current address: Nucleotide and Peptide Drug Discovery Center, Tokyo Medical and Dental University, Yushima 1-5-45, Bunkyo-ku, Tokyo 113-8519, Japan.

four naphthyridine carbamate (NC) moieties form hydrogen bonds with four guanines in the CGG/CGG. This interaction leads to the flip-out of two cytosines. To gain insight into the mode of NCD binding to the CGG repeat, we measured the mass spectrometry of a CGG repeat DNA  $(5'-d(CGG)_{10}-3')$  in the presence of NCD, and confirmed that even numbers of NCD molecules bound to the CGG repeat. Based on these experimental results, we have postulated that the complex of NCD molecules and CGG repeat DNA would adopt repetitive structures consisting of one CGG/CGG unit with two NCD molecules, as depicted in Fig. 1C and D.

In the proposed binding model of NCD to the CGG repeat DNA, one question arises as to whether NCD binding to the CGG repeats followed the nearest-neighbor exclusion principle as observed for the binding of DNA intercalators.<sup>16</sup> When an intercalator binds to a duplex DNA, the two adjacent intercalation sites would remain unoccupied due to the perturbation of the base pair structure by intercalation. When CGG repeat DNA formed the hairpin secondary structure, there are contiguous CGG/CGG units, and these CGG/CGG units are in different environments depending on the position, e.g., the end or the middle of the hairpin structure. Furthermore, NCD-binding to one CGG/CGG unit should perturb the structure of the neighboring CGG/CGG unit and, hence, the binding of NCD. Although MS data revealed an even number of NCD molecules bound to the CGG repeat DNAs, our understanding of the NCD binding mode to CGG repeat DNA remains limited. To address the question raised above, we synthesized <sup>15</sup>N-labeled NCD (<sup>15</sup>N-NCD, Fig. 1A) and measured the <sup>15</sup>N NMR spectrum to see how the <sup>15</sup>N signal changed as the number of CGG/CGG units increased. In the proposed model of (CGG/CGG)<sub>2</sub> with four NCD molecules (Fig. 1C), the NC moieties (NC1, NC2, NC3, and NC4) shown in blue has likely a similar chemical environment to those in the complex of (CGG/CGG)<sub>3</sub> with six NCD molecules (Fig. 1D). Likewise, NC moieties shown in green should be in similar environments but different from those in blue. In addition, the NC moieties colored in cyan in Fig. 1D would be in different environments from those colored in blue and green, although the difference could be small. The different chemical environments of NC moieties could be observed by the difference in the chemical shift in NMR analysis. Here, we report that <sup>1</sup>H-<sup>15</sup>N HSQC spectra of <sup>15</sup>N-NCD with DNA involving one, two, and three units of CGG/CGG clearly indicated that NCD could bind to the contiguous CGG/CGG sites.

2-Amino-1,8-naphthyridine labeled with <sup>15</sup>N at the amino group was synthesized by reacting 2-chloro-1,8-naphthyridine with 22% aqueous <sup>15</sup>N ammonia in the presence of 1.0 equivalent copper in 40% yield. Boc-protected <sup>15</sup>N-NCD and <sup>15</sup>N-NCD were prepared following the reported procedure<sup>17</sup> (Scheme S1 in ESI<sup>†</sup>). The <sup>1</sup>H-NMR of Boc-protected <sup>15</sup>N-NCD in CDCl<sub>3</sub> is presented in Fig. S1 in the ESI.<sup>†</sup> The amide protons of <sup>15</sup>N-NCD appeared at 7.69 ppm as a superposition of two doublets due to the heteronuclear coupling between <sup>1</sup>H and <sup>15</sup>N.

Next, <sup>15</sup>N-NCD was used to assign amide protons in the complexes involving NCD and a CGG/CGG unit. The dsDNA GG1, composed of DNA1: 5'-d(CTAA CGG AATG)-3' and DNA2:



**Fig. 2** (A) dsDNAs used in this study, denoted as GGn (n = 1 to 3) based on the number of CGG/CGG units. The green numbers indicate residue numbering in GG1. (B)  ${}^{1}\text{H}{-}{}^{15}\text{N}$  HSQC of GG1 (200  $\mu$ M) in 5% D<sub>2</sub>O in sodium phosphate buffer (20 mM, pH = 6.8) containing 100 mM NaCl in the presence of  ${}^{15}\text{N}{-}\text{NCD}$  (400  $\mu$ M). The imino-proton region (10.5 to 12.5 ppm) of the  ${}^{1}\text{H}$  and  ${}^{15}\text{N}$  HSQC spectra is displayed. The  ${}^{1}\text{H}{-}\text{NMR}$  of the NCD-GG1 complex is presented above the HSQC. Amide protons of NCD are labeled as NCn (n = 1 to 4), and the imino-protons of guanines are denoted as Gn (n =numbering). (C) Schematic illustration of the binding of two  ${}^{15}\text{N}{-}\text{NCD}$  molecules to the CGG/CGG unit in GG1. Amide protons next to  ${}^{15}\text{N}$  are colored in blue and observed in the HSQC shown in Fig. 3B.

5'-d(CATT CGG TTAG)-3', was used for the NMR structural determination of the NCD-CGG/CGG complex in our previous study<sup>15</sup> (Fig. 2A). The <sup>1</sup>H-<sup>15</sup>N HSQC spectrum of GG1 (200  $\mu$ M) in the presence of <sup>15</sup>N-NCD (400  $\mu$ M) is depicted in Fig. 2B. For clarity, the <sup>1</sup>H-NMR spectrum of the NCD-GG1 complex is overlaid above the HSQC spectrum. In the <sup>1</sup>H-<sup>15</sup>N HSQC, four cross peaks were observed at (<sup>1</sup>H, <sup>15</sup>N) coordinates: (11.02, 118.4), (11.09, 119.6), (11.18, 119.8), and (11.63, 119.6), and identified as amide protons of NC1, NC2, NC3, and NC4, respectively (Fig. 2C). As two amide protons would be anticipated for one NCD binding, the HSQC spectra clearly indicated the binding of two NCD molecules to GG1.

Continuing, we employed another model DNA named GG2, comprising two consecutive CGG/CGG units (Fig. 2A, n = 2) for titration experiments using <sup>15</sup>N-NCD. The sequences of GG1 and GG2 are identical except for the number of CGG/CGG units. The titration results are presented in Fig. 3A. In the <sup>1</sup>H-NMR spectrum of GG2, the imino-protons of thymidine were observed between 13.5 and 14.0 ppm, while those of guanine were between 12.5 and 13.0 ppm. Upon titration with <sup>15</sup>N-NCD, the signal initially became complex (*e.g.*, GG2: <sup>15</sup>N-NCD = 1:3), but as the titration proceeded, the spectrum simplified. Eventually, the



Fig. 3 (A) One-dimensional <sup>1</sup>H spectra of GG2 (500  $\mu$ M) in sodium phosphate buffer (pH 6.8, 20 mM) containing 100 mM NaCl in the presence of <sup>15</sup>N-NCD with various GG2-<sup>15</sup>N-NCD ratios at 283K. The ratio displayed in the left side of each spectrum represents the proportion of GG2 and <sup>15</sup>N-NCD. (B) <sup>1</sup>H-<sup>15</sup>N HSQC of GG2 (500  $\mu$ M) in 5% D<sub>2</sub>O in sodium phosphate buffer (20 mM, pH = 6.8) containing 100 mM NaCl in the presence of <sup>15</sup>N-NCD (3 mM). The <sup>1</sup>H-NMR spectrum of the NCD-GG2 complex is presented above the HSQC. Dashed guidelines illustrate the correlation of <sup>1</sup>H-NMR and HSQC. A red star highlights a superposition of two peaks. (C) The 3D view of Fig. 2B. The peak highlighted with a red star has approximately twice the intensity of the other peaks.

shifts of signals saturated upon the addition of 6 equivalents of <sup>15</sup>N-NCD. In Fig. 3B, the <sup>1</sup>H-<sup>15</sup>N HSQC of the GG2 and <sup>15</sup>N-NCD complex under saturated conditions is displayed. The <sup>1</sup>H-NMR spectrum shown above the HSQC spectrum is the GG2 and unlabeled NCD complex under the saturated conditions. At a glance, seven cross peaks in the HSQC spectrum were identified at (<sup>1</sup>H, <sup>15</sup>N) coordinates: (10.89, 118.6), (10.94, 119.8), (11.03, 119.1), (11.03, 119.2), (11.03, 119.8), (11.11, 118.8), and (11.48, 119.6). Upon further analysis using a 3D view (Fig. 3C), the peak at (11.11, 118.8), marked with a red star, displayed approximately twice the intensity compared to other peaks, indicating a superposition of two peaks. Therefore, eight cross peaks corresponding to the <sup>1</sup>H-<sup>15</sup>N coupling were identified, suggesting that four NCD molecules bound to GG2.

In a similar manner, the model DNA GG3, comprising three consecutive CGG/CGG units, was subjected to titration with <sup>15</sup>N-NCD (Fig. 4A). Corresponding to the results of Fig. 3A, a complex spectrum was observed at the early stages of titration, but it simplified as the titration progressed. Ultimately, with the addition of 15 equivalents of <sup>15</sup>N-NCD, the changes in the peaks reached saturation. In Fig. 4B, the <sup>1</sup>H–<sup>15</sup>N HSQC spectrum, capturing the complex of GG3 and <sup>15</sup>N-NCD under saturated conditions, is presented. At a glance, nine peaks are visible at (<sup>1</sup>H, <sup>15</sup>N) coordinates: (10.90, 118.6), (10.92, 119.2), (10.94, 119.8), (11.03, 118.9), (11.04, 119.1), (11.04, 119.2), (11.04, 119.8), (11.09, 118.9), (11.48, 119.6). Through a 3D view analysis (Fig. 4C), it was determined that the peaks at (10.92, 119.2),



Fig. 4 (A) One-dimensional <sup>1</sup>H spectra of GG3 (500  $\mu$ M) in sodium phosphate buffer (pH 6.8, 20 mM) containing 100 mM NaCl in the presence of <sup>15</sup>N-NCD with various GG3–<sup>15</sup>N-NCD ratios at 283K. (B) <sup>1</sup>H–<sup>15</sup>N HSQC of GG3 (500  $\mu$ M) in 5% D<sub>2</sub>O in sodium phosphate buffer (20 mM, pH = 6.8) containing 100 mM NaCl in the presence of <sup>15</sup>N-NCD (3 mM). Dashed guidelines illustrate the correlation of <sup>1</sup>H-NMR and HSQC. Three red stars highlight the superpositions of two peaks. (C) The 3D HSQC view. Three peaks highlighted with red stars show twice the intensity compared to other peaks.

(11.03, 118.9), and (11.09, 118.9) were superpositions of the two peaks. Consequently, twelve cross peaks are observed as amideprotons, indicating the binding of six NCD molecules to GG3.

In these studies, three different DNAs containing one (GG1), two (GG2), and three units (GG3) of CGG/CGG were employed to measure <sup>1</sup>H-<sup>15</sup>N HSQC spectra of the complexes with <sup>15</sup>Nlabeled NCD. In GG1, the CGG/CGG unit was sandwiched between Watson-Crick base pairs. GG2 had two adjacent CGG/CGG units at one side with Watson-Crick base pairs, and GG3 additionally included an "inner" CGG/CGG unit, sandwiched by two "outer" CGG/CGG units.

Comparison of the cross peaks observed in the HSQC spectra for GG1, GG2, and GG3 with <sup>15</sup>N-NCD was conducted (Fig. 5). In the HSQC spectra of GG1, four cross peaks were identified, exhibiting different <sup>1</sup>H and <sup>15</sup>N chemical shifts (Fig. 5A). These cross peaks were assigned based on prior NMR studies.<sup>15</sup> A notable difference in the <sup>1</sup>H and <sup>15</sup>N chemical shifts between NC1 and NC4 amide N–H was attributed to the adjacent nucleotide bases, such as adenine for NC1 and thymine for NC4.

In the HSQC spectra of GG2 with <sup>15</sup>N-NCD, these four cross peaks shifted upfield in terms of the <sup>1</sup>H chemical shift, while no significant changes in the <sup>15</sup>N chemical shift were observed. Additionally, three new cross peaks, totaling four N–H, appeared (Fig. 5B, circled with a red line). These seven cross peaks in the GG2–NCD complex displayed almost the same <sup>1</sup>H and <sup>15</sup>N chemical shifts in the HSQC spectra of the GG3–NCD complex, where two new cross peaks, each containing two N–H, emerged (Fig. 5C, circled with a green line).

The remarkable similarity in the <sup>1</sup>H and <sup>15</sup>N chemical shifts of the eight NC moieties in both GG2 and GG3 complexes



Fig. 5 Comparison of  $^1H-^{15}N$  HSQC. (A) GG1 and  $^{15}N$ -NCD, (B) GG2 and  $^{15}N$ -NCD, and (C) GG3 and  $^{15}N$ -NCD complex.

indicated that the chemical environment of these NC moieties remained consistent. This suggests that NCD-binding to the two "outer" CGG/CGG units in GG3 is similar to that in GG2. The two new cross peaks in GG3 are evidently due to NCD binding to the "inner" CGG/CGG unit and have different <sup>1</sup>H and <sup>15</sup>N chemical shifts, indicating a distinct chemical environment compared to the NC moieties bound to the "outer" unit. These observations strongly indicate that NCD can bind to the contiguous CGG/CGG units in the CGG repeat hairpin, even though the "inner" unit may experience structural perturbations due to NCD binding to neighboring units.

Another crucial observation was made in these studies. In the titration experiments with both GG2 and GG3, the spectrum initially became complicated but eventually converged. Such complexity was not observed in the titration of NCD to GG1.<sup>15</sup> The binding pathway of NCD in the presence of contiguous CGG/CGG units likely involves intermediates but converges to a specific structure under equilibrium, suggesting that NCD can likely bind to CGG repeat DNA, even if the repeat is substantial in length.

In conclusion, our investigation focused on the binding mode of NCD to repetitive CGG/CGG units using <sup>15</sup>N-labeled

NCD and  ${}^{1}\text{H}{-}{}^{15}\text{N}$  HSQC NMR techniques. By comparing the  ${}^{1}\text{H}{-}{}^{15}\text{N}$  HSQC spectra of complexes with three DNA models containing one, two, and three CGG/CGG units, we confirmed that NCD can bind to all three CGG/CGG units in GG3. This implies the plausible scenario of NCD binding to long CGG repeat DNA.

#### Conflicts of interest

There are no conflicts to declare.

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