d(CGGTGGT) forms an octameric parallel G-quadruplex via stacking of unusual G(:C):G(:C):G(:C):G(:C) octads

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ABSTRACT

Among non-canonical DNA secondary structures, G-quadruplexes are currently widely studied because of their probable involvement in many pivotal biological roles, and for their potential use in nanotechnology. The overall quadruplex scaffold can exhibit several morphologies through intramolecular or intermolecular organization of G-rich oligodeoxyribonucleic acid strands. In particular, several G-rich strands can form higher order assemblies by multimerization between several G-quadruplex units. Here, we report on the identification of a novel dimerization pathway. Our Nuclear magnetic resonance, circular dichroism, UV, gel electrophoresis and mass spectrometry studies on the DNA sequence dCGGTGGT demonstrate that this sequence forms an octamer when annealed in presence of K+ or NH4+ ions, through the 5'-5' stacking of two tetramolecular G-quadruplex subunits via unusual G(:C):G(:C):G(:C):G(:C) octads.

INTRODUCTION

G-quadruplexes are unique structures formed by Hoogsteen-type base pairing between four guanines and involving chelation of a metal ion. G-quadruplexes are typically formed by intramolecular folding of guanine-rich oligonucleotide (GRO) sequences or by intermolecular association of two or four sequences leading to the formation of dimeric or tetrameric complexes (1). The biological importance of these structures is 3-fold: (i) the occurrence of short G-rich sequences able to fold into G-quadruplex structures at the ends of telomeric DNA in eukaryotic chromosomes (2,3); (ii) the high prevalence of G-rich sequences in a large number of eukaryotic and prokaryotic genomes, and the increasing number of G-quadruplexes arising from these sequences (2,4,5); and (iii) their presence in the scaffold of several aptamers that have the ability to selectively bind to biologically relevant proteins and small molecules (6–10).

G-quadruplexes have the ability to form an array of conformations differing in structural features such as the molecularity, the relative orientation of the strands, the size of the loops connecting the strands and the glycosidic conformation of guanosine residues (syn or anti) (11). Further structural diversity arises from the quadruplex capacity to accommodate A-, T- and C-tetrads (12–14), as well as tetrads formed by modified residues (15,16). Moreover, in the last few years, several papers describing the higher order packing of intramolecular (17–19) or intermolecular (20,21) DNA quadruplexes have also been published. Particular attention has been devoted to the understanding of mechanisms responsible for the self-organization of human telomeric DNA into higher ordered G-quadruplex structures (17,19,22). Knowledge of such mechanisms is precious for shedding light on the in vivo role of quadruplex DNA as well as for the design of supramolecular DNA structures provided with novel functionalities. Understanding and controlling higher order structure formation is also of prime importance for in vitro studies of oligonucleotides derived from biologically relevant sequences, and for controlling the aggregation state of oligonucleotide aptamers aimed at therapeutic applications.

One of the identified structural moieties required for quadruplex dimerization is the presence of a stretch of guanines at the 5'- or 3'-termini of parallel G-quadruplex-forming GROs. These types of GROs have been reported to form higher ordered ‘dimer G-quadruplexes’ by either the association of interlocked slipped strands (23) or by end-to-end stacking (20,23). The presence of even a single...
base other than G at 5’- or 3’-termini has been reported to prevent the dimerization. In this context, we report here the results of our structural investigations on the sequence CGGTGGT under different conditions of oligodeoxy-nucleotide (ODN) concentration, pH and salts. We observe that the sequence folds into a ‘dimeric G-quadruplex’ (from now on 2Q), which is in fact an octamer of the sequence, coexisting in solution with minor amounts of other DNA secondary structures and with the single-stranded molecule. This higher ordered structure is obtained by head-to-head stacking between two G(3C):G(3C):G(3C):G(3C) octads, each belonging to one of the two ‘monomeric G-quadruplexes’ (1Q), as ascertained from the results described below.

MATERIALS AND METHODS

DNA sample preparation

DNA oligonucleotides TGGGGT, CTGTGTT, TGGTG GC, CCGTGGT and CCGGGT were chemically synthesized on an 8909 DNA/RNA synthesizer (Applied Biosystem), or purchased from Eurogentec (Belgium) for mass spectrometry experiments. DNA concentration expressed in strand molarity was calculated using a nearest-neighbor approximation for the absorption coefficients of the unfolded species. Ammonium acetate was obtained from a 5 M solution from Fluka (Molecular Biosystem), or purchased from Eurogentec (Belgium) for DNA sample preparation.

ODN samples were annealed by heating at 90°C for 15 min and then cooling and storing at 5°C. The pH was adjusted to 7.0 or 4.5 using HCl/KOH or AcOH/NH4OH aqueous solutions before the annealing procedure.

Polyacrylamide gel electrophoresis

The molecular size of structures were probed using non-denaturing polyacrylamide gel electrophoresis (PAGE) as previously described (24). ODN samples of 10 μM, obtained by diluting the annealed stock solutions just before the experiment, were loaded on a 15% polyacrylamide gel supplemented with 20 mM of KCl or NH4OAc. Glycerol of 10% was added just before loading. The gels were run at 26°C at constant voltage (100 V) for 2.5 h. The bands were visualized by UV shadowing and after ‘Syber green’ coloration.

Electrospray mass spectrometry

A Q-TOF Ultima Global mass spectrometer (Waters, Manchester, UK) was used to characterize the samples in ammonium acetate, and to perform a kinetics experiment. ODN samples (80 μM single-strand concentration), obtained as described in the PAGE paragraph with the addition of 10% methanol, were infused at 4 μl/min in the electrospray (ESI) source that was operated in negative ion mode (capillary voltage = −2.2 kV). Temperatures of ESI source and nitrogen desolvation gas were 80°C and 100°C, respectively. The source voltages were optimized for the best compromise to observe the single strand, tetramer and octamer: cone = 80 V, RF Lens 1 = 80 V, collision energy = 10 V and source pressure = 3.06 mbar. For the kinetics experiments, the reference strand dT6 was added to the dCGGTGGT strand, so as to follow the intensities of the reactant (dCGGTGGT single strand, noted G1), the intermediates (dimer = G2, trimer = G3, tetramer = G4 = 1Q) and the product (octamer = G8 = 2Q) relative to that of the reference. The method for obtaining relative response factors of the monomer, the tetramer 1Q and the octamer 2Q compared to the reference is described elsewhere (25), and the details of its application in the present case are given in Supplementary Data. In addition, a Bruker Apex-Qe9.4 T ESI-FTICR mass spectrometer was used to confirm the charge states from the isotopic distributions by recording high-resolution data, and to confirm the number of ammonium cations retained in each complex. The instrument voltages were chosen so as to minimize collision-induced dissociation and therefore keep the non-covalent complexes intact, including the inner ammonium cations: cap exit = −22 V, skimmer = −15 V, octapole offset = −4 V and cell cell trap = +4 V.

Circular dichroism

Circular dichroism (CD) spectra were measured at 25°C on a Jasco J-715 spectropolarimeter equipped with a Peltier Jasco JPT423S, using a 0.1-cm path length quartz cuvette with a reaction volume of 500 μl. CD spectra were recorded at different time points after the annealing on 40 μM ODN samples annealed at strand concentration of 200 μM or 6.0 mM. Spectra were averaged over three scans. A buffer baseline was subtracted from each spectrum and the spectra were normalized to have zero at 360 nm.

Nuclear magnetic resonance

Nuclear magnetic resonance (NMR) data were collected on a Varian Unity INOVA 500 MHz spectrometer equipped with a broadband inverse probe with z-field gradient, and on a Varian UNITY INOVA 700 MHz spectrometer equipped with a triple resonance cryoprobe. The data were processed using the Varian VNMRS software package. One-dimensional NMR spectra were acquired as 16 384 data points with a recycle delay of 1.0 s at 5, 25 and 65°C. Data sets were zero filled to 32 768 points prior to Fourier transformation and apodized with a shifted sine bell squared window function. Two-dimensional NMR spectra were all acquired at 25°C using a recycle delay of 1.2 s. NOESY spectra were acquired with mixing times of 100, 200 and 300 ms. TOCSY spectra were recorded with the standard MLEV-17 spin-lock sequence and a mixing time of 80 ms. For the experiments in H2O, water suppression was achieved by including a double pulsed-field gradient spin-echo (DPFGSE) module (26,27) in the pulse sequence prior to acquisition. In all 2D experiments, the time domain data consisted of 2048 complex points in t2 and 400 fids in t1 dimension. NMR samples were prepared at
6 mM strand concentration in 0.6 ml of 100 mM or 1.0 M K\(^+\) buffer obtained as above described. Sequence-specific resonance assignments for CGGTGGT were obtained by using NOESY and TOCSY spectra following the standard procedure (28,29). The self as well as sequential NOE connectivities from H\(^8\) to H\(^1\)' and H\(^2\)/H\(^2\)' were traced for all the nucleotides. The connectivities followed the standard patterns as for B-DNA, apart from noticeable exceptions between C\(_1\) and G\(_2\), and complete assignments of base and sugar protons was obtained.

Molecular modeling

The starting structure was generated using the builder module of Insight II (2005) program (Accelrys, Inc) on an Intel PC workstation running Red Hat Enterprise Linux. The model was parameterized according to the AMBER force-field, and calculations were performed using a distance-dependent macroscopic dielectric constant of 4.0 and an infinite cut-off for non-bonded interactions to partially compensate for the lack of solvent. The details of model building and energy minimization are given in Supplementary Data.

RESULTS

dCGGTGGT forms an octamer

The formation of a higher order assembly than a tetramer by the DNA sequence dCGGTGGT was first noticed in native PAGE experiments. Figure 1a shows PAGE experiments on the following DNA sequences: TGGGGT annealed in 0.1 M potassium buffer (lane 1) used as a tetramolecular quadruplex ([dTGGGGT]\(_4\)) size-marker, C TGTGTT (lane 2) used as a single-stranded 7-mer marker, TGGTGGC (lanes 3 and 4), CGGTGGT (lanes 5 and 6) and CGGGGT (lanes 7 and 8). The sequence TGGTGGC migrates similarly to [dTGGGGT]\(_4\); whereas, the sequence CGGTGGT shows a major band migrating much slower. We conclude that TGGTGGC forms prevalently a tetrameric assembly, in agreement with the reported tetramer G-quadruplex formation for this sequence (30), whereas C GGTGGT forms a higher order assembly. The polarity of the sequence therefore appears crucial in the higher order structure formation. The migration profile of dCGGGGT has also been investigated to determine whether the additional central thymine played any role in the higher order structure formation. Two bands are observed for dCGGG GT, the faster one migrating as the tetrameric [dTGGGG T]\(_4\), and the slower one migrating like the structure formed by CGGTGGT.

As the PAGE results are similar in NH\(_4\)\(^+\) and in K\(^+\) cations, we therefore used ESI-MS, which can only be carried out in NH\(_4\)\(^+\), to determine unambiguously the strand stoichiometry of the higher order assemblies formed by CGGTGGT and CGGGGT. The ESI-MS spectra of TGGTGGC, CGGTGGT and CGGGGT, each separately annealed at 6 mM in 150 mM NH\(_4\)OAc (pH = 7.0) and diluted to 80 \(\mu\)M final strand concentration, are shown in Figure 1b–d, respectively. All three sequences show peaks corresponding to a tetramolecular
assembly (noted G₄), but the spectra of CGGTGGT and CGGGGT also display peaks corresponding to an octamer assembly (noted G₈). We therefore assign the slow migrating PAGE band to an octameric self-assembly. In terms of relative intensities, although native PAGE and ESI-MS experiments were performed on samples obtained by dilution of the same annealed stock solution, the octamer is the darkest band in the gels of CGGTGGT (lanes 5 and 6), whereas the ESI-MS peaks of the octamer have lower intensities than the tetramer peaks. The analysis of the electrospray response factors in the kinetics ESI-MS experiments (see below and Supplementary Figure S2) indeed reveals the octamer has a lower ESI-MS response than the tetramer in the spectrometer used, and that the octamer is indeed the major species in solution.

The octamer contains stacked G-tetrads

All G-rich sequences studied here are expected to form G-quadruplex structures. Note that ‘G-quadruplex’ means that the structure contains stacked G-quartets, not necessarily that the assembly is a tetramer. CD was used to probe the relative orientation of the bases in the tetramer [TGGTGCG]₄ and in the octamer [CGGTGGT]₈. The overall CD profiles recorded in K⁺ (Figure 2a) and NH₄⁺ (Figure 2b) are in agreement with the formation of G-quadruplex structures (31,32) both in K⁺ and NH₄⁺ media, showing a positive maximum at about 260 nm and a negative minimum close to 240 nm, which are characteristics of head-to-tail arrangement of guanines, as typically found in parallel oligodeoxynucleotides G-quadruplexes (33–35). Interestingly, the CD spectra of CGGTGGT also show a weak, but clearly visible negative band centered at 290 nm. This minimum suggests that additional base stacking interactions could be involved in the formation of the higher ordered quadruplex species by allowing the quadruplex multimerization via end-to-end stacking. A typical heteropolar stacking in G-quadruplexes formed by strands with no inverted polarity usually results in a positive band at 290 nm (34), but negative bands at 290 nm have been observed in octameric self-assemblies of lipophilic guanosine derivatives (36). The type of stacking present in our octamers could therefore resemble that present in those lipophilic octamers.

![Figure 2. G-quadruplex formation probed by (a and b) CD in 100 mM KCl (a) and 150 mM NH₄OAc (b) of the sequences CGGTGGT (solid line) and TGGTGCC (dotted line), and by (c) counting the number of ammonium cations incorporated in the tetramer (noted G₄) and the octamer (noted G₈) of CGGTGGT using high resolution ESI-FTICR mass spectrometry.](image-url)
Because the CD signature is peculiar and does not allow to conclude unambiguously to a G-quadruplex structure, we decided to probe cation incorporation into the octamer. Indeed, successive G-quartets incorporate monovalent cations (here K+ or NH4+*) by coordination between eight guanines. High-resolution ESI-FTICRMS was used to count the number of ammonium cations per tetramer and per octamer (Figure 2c). Cation incorporation was found in all cases, confining G-quadruplex formation both in the tetramer and the octamer. The major peaks correspond to three cations per tetramer [dCGGTGGT]4 and eight cations per octamer [dCGGTGGT]8. The number of cations is indicative of the number of stacked quartets. Interestingly, the tetramer [dCGGTGGT]4 contains three ammonium cations, whereas the tetramer [dTGGTGGC]4 contains four (data not shown), suggesting a structural difference depending on the strand polarity. However, the incorporation of eight ammonium cations in the octamer suggests that within the octamer, each tetrameric subunit is now capable of incorporating four ammonium cations.

Influence of sample preparation procedure

In all experiments described above, the octamer was obtained by annealing the strand CGGTGGT at 6 mM single-strand ODN concentration at pH = 7.0 in 100 mM K+ or 150 mM NH4+. CD experiments carried out using a higher K+ concentration (1 M) show that higher cation concentration further favors the formation of the structure characterized by the negative band at 290 nm (Supplementary Figure S1). We also tested the influence of the solution pH, to test whether cytosine protonation might favor the formation of the new structure, but no clear influence of pH was found (see Supplementary Figure S1 and NMR results described below and in Supplementary Figure S4).

However, because of the octamer stoichiometry, we anticipated that the strand concentration upon annealing would have a great influence on the formation kinetics. Figure 3a shows the CD spectra recorded on CGGTGGT annealed at 6.0 mM or 0.2 mM ODN concentration in 1 M K+. The presence of the negative band at 290 nm was strictly dependent from ODN concentration: it is present when the sample is annealed starting from the 6 mM stock solution, whereas it is not observed when the sample is obtained by annealing the 0.2 mM ODN (Figure 3a). However, the CD spectrum of the species formed from the 0.2 mM ODN solution resembles that of a regular tetramolecular G-quadruplex. The influence of the strand concentration, therefore, indicates slow formation kinetics for the octamer.

To study the kinetics of octamer formation in the 6 mM stock solution, and to probe the nature of the reaction intermediates, we recorded ESI-MS spectra as a function of the time after addition of cation in the ODN solution, at room temperature. This experiment could be carried out only in ammonium acetate (150 mM). The single-stranded dT6 strand (0.75 mM) is added to the 6 mM single-stranded dCGGTGGT in water. Quadruplex formation is initiated by the addition of ammonium acetate (final concentration: 150 mM) at room temperature. At each time point, an aliquot of the sample is diluted to 80 μM final dCGGTGGT single-strand concentration, and injected in the ESI-MS. We therefore assume that, once formed, the octamer resists the dilution step. The signal intensity of the reference is used to normalize the intensity variations of the dCGGTG GT monomer, tetramer and octamer signal, and deduce their respective response factors (see Supplementary Figure S2). Figure 3b shows the time evolution of the relative proportions of single strand, tetramer and octamer, after correction for the relative response of each species, expressed as the fraction of single strand in each form. The single strand is quickly converted into a tetramer, but the tetramer-to-octamer conversion is slower. The dimerization is therefore the rate-limiting step in the octamer formation. The results also show that, although the octamer is not the most abundant peak in the mass spectra (see Figures 1b–d and 2c), it is nevertheless the predominant species. The predominance of the octamer at long time scales also confirms that, once formed, the octamer resists the dilution step.

Figure 3. Influence of the sample preparation conditions on the octamer formation. (a) Influence of strand concentration: CD spectra of CGGTGGT in 1.0 M K+ buffer annealed, respectively, at 6 mM (solid line) and 200 μM (dotted line) single-strand concentration. (b) Kinetics of tetramer and octamer formation of 6 mM CGGTGGT in 150 mM ammonium acetate at room temperature, as determined by time-resolved ESI-MS.
NMR investigation of the octamer structure

The $^1$H NMR spectrum of CGGTGGT in 100 mM K$^+$ buffer at 25°C recorded 1h after the annealing (Supplementary Figure S3, bottom left) is characterized by the presence of four well-resolved signals in the 11.0–11.6 ppm region attributable to the exchange-protected imino protons involved in the formation of Hoogsteen hydrogen bonds of four G-tetrads (37–39), in close analogy with what was previously observed for the parallel quadruplex structure formed by TGGTGGC (30).

Furthermore, the count of quadruplex NMR signals is the same as what must be expected for the NMR spectrum of the corresponding single strand, thus suggesting the formation of a highly symmetric parallel G-quadruplex possessing a 4-fold symmetry. However, the aromatic region of $^1$H NMR spectrum is populated by more than the expected seven signals belonging to the H8/H6 aromatic protons of nucleobases. In order to discriminate whether the minor peaks were due to sample impurities or to the coexistence in solution of alternative minor conformations of CGGTGGT, we recorded the $^1$H NMR spectrum also at 65°C, and the high-temperature spectrum was clearly in agreement with the second hypothesis as evidenced by the presence of just seven well-defined signals in the aromatic region of the spectrum. We therefore studied the influence of time elapsed after the annealing (Supplementary Figure S3), of K$^+$ concentration (Supplementary Figure S3), and of pH (Supplementary Figure S4) on the 1D NMR spectra. In agreement with the CD data, the NMR results confirmed that the best medium for the structural study of the new octameric structure formed by CGGTGGT requires 1.0 M K$^+$ and long folding times (Figure 4a), and that the change of pH from 7.0 to 4.5 does not affect the folding of the structure.

Figure 4. Study of the octamer structure by NMR. (a) $^1$H-NMR spectra recorded at 25°C of CGGTGGT in H$_2$O/D$_2$O 9:1, annealed at 6 mM strand concentration in 1 M K$^+$ at pH = 7.0. (b–g) Expansions of 2D NOESY spectrum of CGGTGGT (25°C, 200 ms mixing time, 1 M K$^+$, pH = 7.0). The sequential H8-H6 and self-peaks involving H1$^+$ protons have been labeled in (b) and (e), respectively. (d) and (g) show NOE cross-peaks involving T4/T7 methyl protons and H8/H6 or imino and amino protons, respectively. The T-tetrad diagnostic peaks between T4Me and T4NH and sequential T4Me-G3NH, T4Me-G3NH$_2$, and T4Me-G5NH are observable in panel (g). Imino–imino NOE correlations are shown in (e). The sequential imino–imino NOE correlations have been drawn in (e). NOE correlations between aromatic H8/H6 and imino and amino protons are shown in (f).
The combined analysis of 2D NOESY and TOCSY spectra (700 MHz, 25°C, 1 M K⁺, 15 days after the annealing) allowed us to get the complete assignment (Table 1) of both exchangeable and non-exchangeable protons of CG GTGGT. The sequential connectivities between the aromatic protons (Figure 4b) were traced, as well as the self and sequential NOE connectivities between H8/H6 base protons and H1', H2', H2'' and H3' sugar protons. The intensities of intranucleotide H8/H1' NOE cross-peaks (Figure 4c) indicate that all Gs adopt an anti glycosidic conformation. This finding together with the observation for each G-tetrad of a GiH8/GjN1H NOE cross-peak (Figure 4f), with i = j (where i and j represent the nucleotide labels), and with the observation of sequential G2N1H/G3N1H and G5N1H/G6N1H cross-peaks (Figure 4e), definitely allowed us to identify the presence of 4-fold symmetric parallel quadruplex motieties in the higher ordered structure formed by CGGTGGT when annealed in K⁺-containing buffer. In close analogy with what observed by Patel, P.K. and Hosur, R.V. for the quadruplex structure formed by TGTTGGC, we observed the formation of a central T-tetrad as ascertained by the appearance of the diagnostic T4NH/T4Me NOE cross-peak (Figure 4g) and supported by the observation of the strong shielding ring current experienced by T4 methyl protons that are significantly upfield-shifted when compared to the corresponding signals of the ‘free’ T (0.72 versus 1.64 ppm).

Although the overall observed NOEs account for a ‘classical’ tetramolecular parallel quadruplex, two peculiar NOE evidences drove us toward the disclosure of the higher ordered octamer structure. In Supplementary Figure S5, we show the 1D cross-section of the NOESY spectrum (200 ms mixing time) of CGGTGGT taken at 5.66 ppm (C1H1' protons). This figure shows that the NOE between C1H1' and C1H5 has almost the same intensity than that between C1H1' and C1H6. These NOEs are in turn stronger than the sequential NOE C1H1'-G2H8, but weaker, as expected, than those between C1H1' and self H2'/H2''. The relative intensities of the aforementioned NOEs involving C1H1' require that the C1H1'-C1H5 distance is in the same range of the intranucleotide C1H1'-C1H6 distance (2.31–3.74 Å depending on the glycosyl torsion angle). This experimental distance is compatible only with the occurrence of an intermolecular dipolar coupling between C1 bases, because the corresponding intranucleotide C1H1'-C1H5 distance would be in the range 4.64–5.49 Å. This finding, together with the observation of the very weak sequential NOE between C1H6 and G2H8 (marked with the arrow in Figure 4b), disclosed that, in the octamer, each C1 base does not stack over the flanking G2 tetrad, as one may expect, but projects outside the quadruplex scaffold in such a way to allow dipolar couplings with one of the four C1 bases belonging to the other tetramer. This unusual arrangement might be favored by the formation of additional C1NH2–G2N3 and G2NH2–C1N3 hydrogen bonds, thus allowing the formation of an unusual G2(C1):G2(C1):G2(C1):G2(C1) octad (Figure 5a) in each of the two tetrameric quadruplexes (1Q).

**DISCUSSION**

The structure adopted by the sequence dCGGTGGT must account for the following observations. It is a highly symmetrical octamer, formed from two parallel-stranded tetramers. In the octamer, each tetramer contains a central T-tetrad and the G-tetrads are stacked in head-to-tail arrangement. However, at the octamer’s interface between the two tetramers, the stacking of the G-tetrads is likely to be different, as determined from CD.

Interlocked structures like that formed by the (dGCGGTG)₄ tetramer (40) were initially taken in consideration to build an octamer model, but were discarded due to NMR inconsistencies and symmetry-related considerations. Unresponsiveness to pH changes also led us to discard structures involving protonated cytosines. The NOEs involving C1H1' point to the formation of a G2(C1):G2(C1):G2(C1):G2(C1) octad (Figure 5a) in each of the two tetrameric quadruplexes. The formation of the planar octad system would allow for further π–π stacking between C1 bases, thus adding more ‘hydrophobic glue’ between the two G-quadruplex tetramers that form the octamer.

Figure 5b shows the energy minimized model of the octamer formed by stacking of two tetramers, each containing a 5'-G(C):G(C):G(C):G(C) octad. The details of molecular modeling are given in Supplementary Figure S6, and additional ribbon and space-filled views of the model are shown in Supplementary Figure S6. The overall topology of each tetramer is largely similar to that of the tetramolecular quadruplex formed by TG GTGGC (see Figure 8 in reference 30), with the central T-tetrad causing a small decrease in the intertwining of the helices. Space filling views (Supplementary Figure S6) also evidence that the octamer structure is very compact, regardless the presence of two stacked octads. These are seen to determine only a slight increase in the thickness of the octamer at the dimerization interface.

Figure 5c–d also shows the head-to-head stacking between the two octads (c) and the stacking between tetrads G2-G3 (d). The stacking between tetrads G3-T4 and T4-G5 steps is shown in Supplementary Figure S7. Although C1 and G2 bases involved in the stacked octads appear slightly rotated with respect to the tetrad plane, Figure 5c shows good stacking between the six-membered rings of G2 bases and between the stacked C1 bases. The electric

**Table 1.** Proton chemical shift (ppm) for 2Q at 25°C, pH 7.0, 1 M K⁺

|            | Hα/Hβ | Hα/CH3 | NH | NH2b | H1' | H2' | H2'' | H3' | H4' | H5'/H5'' |
|------------|-------|--------|----|------|-----|-----|------|-----|-----|----------|
| C1         | 7.54  | 5.48   |    | 5.63 | 2.28 | 2.10 | 4.66 | 4.13 | 3.87 | 3.95     |
| G2         | 8.14  | 11.56  | 9.82 | 6.09 | 2.72 | 2.96 | 5.10 | 4.49 | 4.18 |
| G3         | 7.56  | 11.13  | 9.80 | 6.12 | 2.55 | 2.94 | 5.07 | 4.50 | 4.30 |
| T4         | 7.63  | 0.72   | 9.40 | 5.84 | 2.24 | 2.46 | 4.92 | 4.34 | 4.25 |
| G5         | 8.00  | 11.26  | 9.82 | 5.98 | 2.72 | 2.84 | 5.10 | 4.49 | 4.20 | 4.30 |
| G6         | 7.73  | 11.05  | 9.81 | 6.27 | 2.58 | 2.71 | 4.98 | 4.52 | 4.11 | 4.25 |
| T7         | 7.34  | 1.64   |    | 6.08 | 2.73 | 2.21 | 4.51 | 4.08 | 4.08 | 4.25 |

aRelative to H2O at 4.78 ppm.
bNH2i refers to the internal (i) H-bonded amino protons of guanosines.
transition moments corresponding to the guanine absorption band at approximately 250 nm are indicated in Figure 5c and d. The stacking interactions in the model allow to explain the CD data: G2-G3 stacking resembles the standard head-to-tail stacking encountered in parallel-stranded G-quadruplexes. However, the octad stacking gives rise to a heteropolar stacking that is distinct from those typically encountered in antiparallel G-quadruplexes. We, therefore, interpret the negative CD band at 290 nm as being due to the octad stacking shown in Figure 5c (36). Interestingly, a similar weak negative CD band at 290 nm was also observed, but not discussed, by Mastugami et al. (41) in the dimer formed by d(GGAGGAGGAGGA), which contains stacked G(:A):G(:A):G(:A):G heptads at the dimerization interface.

Our results therefore suggest that, like adenine, cytosine is also capable of inducing the formation of higher order structures by the formation of octads. One must note, however, that the formation of the octamer by the sequence dCGGTGGT is slow. ESI-MS kinetics experiments in ammonium acetate provide some insight into the mechanism of octamer formation. The first step is the formation of a tetramer that incorporates only three ammonium ions. This means either that the tetramer of [dCGGTGGT]₄ has either a central or a terminal tetrad disrupted, or that the tetramer contain slipped strands and hence incomplete tetrad formation (42). The rate-limiting step is the conversion of tetramer to octamer, and in contrast with the tetramer, the octamer incorporates four cations per tetrameric subunit. We therefore conclude
that the rate-limiting step is the rearrangement of the tetramer into a fully aligned parallel structure that is amenable to octad formation and stacking thereof. Further experiments with various sequences are underway to probe the effect of the central T-tetrad on stability and rate of formation of octamers.

CONCLUSIONS

In summary, the present study demonstrates that the formation of a quadruplex dimer (2Q) can occur when the 5′-cytosine-ending CGGTTGGT DNA sequence, but not the inverted TGTTGGC sequence, is annealed in the presence of K+ or NH4+ ions. The NMR and molecular modeling data disclosed that the formation of 2Q is allowed by the formation of an unusual octad arrangement, in each of the two 1Q units, involving C1 and G2 bases of each CGGTTGGT strand. Up to now, dimerization of G-quadruplexes had only been reported via guanine–adenine heptads (41), and recently via a shared uracil tetrad (43). Our finding of G-quadruplex dimerization with 5′-cytosine-containing strands adds further possibilities for the design of novel quadruplex-based DNA functional nanostructures, and further complexity to the mechanisms of folding and assembly of G-rich sequences. It is particularly important for in vitro studies of artificially designed (e.g. aptamers) or biologically relevant sequences.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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