Structure of a two-G-tetrad intramolecular G-quadruplex formed by a variant human telomeric sequence in K⁺ solution: insights into the interconversion of human telomeric G-quadruplex structures

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ABSTRACT

Human telomeric DNA G-quadruplex has been considered as an attractive target for cancer therapeutical intervention. The telomeric sequence shows intrinsic structure polymorphism. Here we report a novel intramolecular G-quadruplex structure formed by a variant human telomeric sequence in K⁺ solution. This sequence forms a basket-type intramolecular G-quadruplex with only two G-tetrads but multiple-layer capping structures formed by loop residues. While it is shown that this structure can only be detected in the specifically truncated telomeric sequences without any 5'-flanking residues, our results suggest that this two-G-tetrad conformation is likely to be an intermediate form of the interconversion of different telomeric G-quadruplex conformations.

INTRODUCTION

Telomeres are specialized DNA sequences that cap the ends of linear chromosomes (1–3). The structure and stability of telomeres are closely related with cancer (4–7), aging (8,9) and genetic stability (3,10,11). Human and vertebrate telomeres consist of highly conserved tandem repeats of the hexanucleotide d(TTAGGG)₅₋₁₀ kb in length 5’ to 3’ toward the chromosome end, terminating in a single-stranded 3’-overhang of 100–200 bases (12–15). In normal cells, each cell replication results in a 50- to 200-base loss of the telomere. After a critical shortening of the telomeric DNA is reached, the cell undergoes apoptosis (16). However, telomeres of cancer cells do not shorten on replication, due to the activation of a reverse transcriptase, telomerase, that extends the telomeric sequence at the chromosome ends (17). Telomerase is activated in 80–85% of human cancer cells and has been suggested to play a key role in maintaining the malignant phenotype by stabilizing telomere length and integrity (18,19). The G-rich telomeric sequence can form G-quadruplex DNA secondary structures that consist of stacked G-tetrad planes connected by a network of Hoogsteen hydrogen bonds and stabilized by monovalent cations, such as Na⁺ and K⁺. The G-quadruplex formation by the single-stranded human telomeric DNA can inhibit the activity of telomerase (20). Compounds that stabilize the intramolecular DNA G-quadruplexes formed in the human telomeric sequence have been shown to inhibit the activity of telomerase and disrupt telomere capping and maintenance, making the intramolecular human telomeric DNA G-quadruplex an attractive target for cancer therapeutic intervention (4,6,7,21–23).

The human telomeric sequence was first shown more than a decade ago to form a basket-type intramolecular structure in Na⁺ solution by NMR, using the 22-nt telomeric sequence d[AGGG(TTAGGG)]₂ (24). The same 22-nt telomeric sequence was reported to form a parallel-stranded intramolecular G-quadruplex in crystallographic analysis in the presence of K⁺ (25). Recent results show that the hybrid-type intramolecular G-quadruplex

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structures appear to be the major conformations formed in human telomeric sequences in K⁺ solution, with a dynamic equilibrium between hybrid-I and hybrid-2 conformations (26–33). The hybrid forms appear to be the major form for human telomeric sequences in K⁺ solution even in the co-presence of high concentration of Na⁺ (26). The two hybrid-type telomeric G-quadruplexes are distinct and yet closely related, and appear to be largely determined by the 3'-flanking sequence and thus the different capping structures (27,28). The highly conserved human telomeric sequence shows intrinsic structure polymorphism [see recent review (34)]; however, little is known about the interconversion of different telomeric G-quadruplex conformations. Interestingly, while the energy difference between the two hybrid-type forms appears to be low as both conformations coexist in K⁺ solution, the kinetics of the interconversion between the two hybrid forms is rather slow and very few exchange peaks were observed on the NMR time scale (27), indicating a high energy barrier of the intermediate state(s). In addition, recent reports have shown a multi-step melting transition of telomeric G-quadruplexes in K⁺ solution and was reported to be more stable than the K⁺ form and the basket-type Na⁺ form telomeric G-quadruplexes (35,36).

In this study, we have determined an interesting intramolecular G-quadruplex structure formed by a variant human telomeric sequence with a G-to-I substitution in K⁺ solution. This sequence forms a basket-type intramolecular G-quadruplex with only two G-tetrads but multiple-layer capping structures formed by loop residues. The stable formation of this two-G-tetrad basket-type structure is driven by the extensive capping structures and loop interactions. We have also shown that this structure can be clearly detected in a specifically truncated telomeric sequence, GGG(TTAGGG)₃T; however, the addition of any 5'-flanking residue(s) abolishes this conformation. A very similar NMR structure has very recently been reported using an 8-Br-incorporated telomeric sequence (37). The structure was suggested to be a new form of the natural human telomeric sequence in K⁺ solution and was reported to be more stable than the hybrid-type structures. Our results differ from the previous report in several aspects. First, our results show substantial differences in the capping structures and loop interactions. Most importantly, our results suggest that this two-G-tetrad conformation is likely to be an intermediate form of the interconversion of different telomeric G-quadruplex conformations, rather than a stable new form of the extended human telomeric sequence, as this structure can hardly be detected in any telomeric sequence with a 5'-flanking segment. We have previously proposed a model for the interconversion between the hybrid-type K⁺ form and the basket-type Na⁺ form telomeric G-quadruplex structures through a strand-reorientation mechanism (26). This two-G-tetrad basket-type structure appears to fit the intermediate state in our proposed transition model, and may also be a transition state of the interconversion of the two hybrid-type conformations. This will be the first report of a NMR-determined structure that represents an intermediate state of G-quadruplex structure interconversion.

### MATERIALS AND METHODS

#### Sample preparation

The DNA oligonucleotides were synthesized using β-cyanoethylphosphoramidite solid-phase chemistry on an Expedite™ 8909 Nucleic Acid Synthesis System (Applied Biosystem, Inc) in DMT-on mode, and were purified using C18 reverse-phase HPLC chromatography, as described previously (26–28,38–41). Samples in D₂O were prepared by repeated lyophilization and final dissolution in 99.96% D₂O. Samples in water were prepared in 10%/90% D₂O/H₂O solution. The final NMR samples contained 0.1–2.5 mM DNA in 25 mM K-phosphate buffer (pH 7.0) and 70 mM KCl.

#### NMR experiments

NMR experiments were performed on a Bruker DRX-600 spectrometer as described previously (26–28,38–41). Standard 2D NMR experiments, including NOESY, TOCSY and DQF-COSY, were collected at 1, 7, 15, 20, 25 and 30°C to obtain the complete proton resonance assignment. Non-exchangeable protons were estimated based on the NOE cross-peak volumes at 50–300 ms mixing times, with the upper and lower boundaries assigned to ±30% of the estimated distances. The methyl base proton Me pseudoatom–H6 distance (2.99 Å) was used as a reference. The distances involving the unresolved protons, e.g. methyl protons, were assigned using pseudo-atom notation in X-PLOR.

#### Distance geometry and simulated annealing (DGSA) calculations

Metric matrix distance geometry (MMDG) and simulated annealing calculations were carried out in X-PLOR (42) to embed and optimize 100 initial structures based on an arbitrary extended conformation for the single-stranded I14Tel23 sequence, as described previously (27,28,38,40). The experimentally obtained distance restraints and G-tetrad hydrogen-bonding distance restraints were included during the calculation.

#### NOE-distance restrained molecular dynamics calculations

All of the 100 molecules obtained from the DGSA calculations were subjected to NOE-restrained molecular dynamics refinement in XPLOR (42) with a distance-dependent dielectric constant, as described previously (27,28,38,40). The force constants were scaled at 10–30 and 80–100 kcal/mol Å² for NOE and hydrogen bond distance restraints, respectively. A total of 508 NOE distance restraints, of which 276 are from inter-residue NOEs, were incorporated into the NOE-restrained structure calculation. Dihedral angle restraints were used to restrict the glycosidic torsion angle (ϕ) for the experimentally assigned syn and anti conformations. A dihedral angle restraint of 60(±40)° was applied to the syn G-tetrad guanines, and a dihedral angle restraint of 240(±40)° was applied to the anti G-tetrad guanines. The force constants of dihedral angle restraints were 10 kcal/mol rad². The eight best molecules were selected based both on their minimal energy terms and number of
NOE violations and have been deposited in the Protein Data Bank (PDB ID 2kka).

**CD spectroscopy**

CD spectra of the oligonucleotides were recorded on a Jasco J-810 spectropolarimeter (Jasco, Easton, MD). A quartz cell of 1 mm optical path length and an instrument scanning speed of 100 nm/min with 1 s response time was used for measurements. The measurements are the averages of three repetitions between 200 and 320 nm at room temperature. Spectra were baseline-corrected and the signal contributions of the buffer were subtracted. DNA solutions for CD measurements were made in 25 mM K(+/Na)-PO4, 70 mM K(+/Na)Cl, pH 7.0 buffer with 10 μM strand concentrations.

**RESULTS**

A variant human telomeric sequence forms a stable intramolecular G-quadruplex structure in K⁺ solution

The 22-nt human telomeric sequence A–Tel21, d[AGGG(TTAGGG)₃] (Figure 1A), has been shown to form a single stable unimolecular G-quadruplex in Na⁺ solution (24) and was used for the crystallography study in the presence of K⁺ (25). However, we found this sequence forms a mixture of multiple G-quadruplex conformations in K⁺ solution as indicated by 1D ¹H NMR (Figure 1B) (26). We have carried out single G-to-I substitution at various guanine positions of the A–Tel21 sequence. Inosine differs from guanine by lacking the NH₂ group at the C2 position. We found that the G-to-I substitution at position 14 results in a sequence (I₁₄–Tel22, Figure 1A) that gives rise to a very different but well-resolved NMR spectrum (Supplementary Figure S1), and the presence of well-resolved peaks between 11 and 12 ppm indicates the formation of a G-quadruplex structure. Minor conformations are present in I₁₄–Tel22 as indicated by the presence of weak resonances, which are much more evident in 2D NOESY spectra (data not shown). The addition of a thymine residue at the 3'-end (I₁₄–Tel23, Figure 1A) stabilizes the major structure and improves the NMR spectral quality (Figure 1B). The ¹H NMR spectrum of I₁₄–Tel23 is much cleaner as the population of minor conformations is clearly reduced as compared to I₁₄–Tel22. The 1D ¹H NMR spectrum of I₁₄–Tel23 in K⁺ solution shows eight well-resolved imino proton resonances at 11–12 ppm with sharp line widths (~6 Hz at 20°C) (Figure 2B), indicating the formation of a stable G-quadruplex structure. The melting temperature of this G-quadruplex is concentration-independent as shown by both NMR and CD, indicating that the I₁₄–Tel23 G-quadruplex structure is unimolecular.

Assignment of proton resonances for the I₁₄–Tel23 G-quadruplex in K⁺ solution

Interestingly, I₁₄–Tel23 shows only eight strong imino peaks in the 1D proton spectrum between 10.6 and 12 ppm in K⁺ solution (Figure 2B bottom). This result indicates that, instead of three G-tetrads, only two G-tetrads appear to form in the I₁₄–Tel23 G-quadruplex. We have site-specifically labeled each guanine of I₁₄–Tel23, one at a time, with low-enrichment (6%) incorporation of ¹⁵N-labeled guanine nucleoside. Using ¹⁵N-filtered experiments as previously reported (26–28,39,40), the imino (one-bond connection to N1) and base aromatic
H8 (two-bond connection to N7) protons of guanines were unambiguously assigned. The assignment of imino protons of guanines of I14–Tel23 is shown in Figure 2. In addition, I14–Tel23 also shows four weaker imino peaks, with two more upfield (10.1–10.5 ppm) and two more downfield (12.1–12.6 ppm) relative to the eight strong imino protons of the two G-tetrads (Figure 1B bottom). We were also able to assign the three weaker imino guanine protons (G10, G22 and G4) using 15N-filtered experiments. The base H6 and methyl proton resonances of thymines have been unambiguously assigned by substituting deoxyuridine (dU) for each thymine one at a time. Complete assignment of the proton resonances of I14–Tel23 was accomplished by using 2D-NOESY, TOCSY and COSY data at various temperatures, and the standard DNA sequential assignment procedure (Figure 3 and Table 1). We were also able to assign the weak imino proton at 12.3 ppm at 20°C to be I14NH1 (Figure 1B bottom) based on the strong NOE with its H2 proton (Figure 3A). The VT studies of I14–Tel23 by NMR shows most imino protons of the two G-tetrads can be detected at 60°C (Supplementary Figure S2). Notably, the imino proton of G10 can still be detected at 50°C. The melting temperature of this G-quadruplex is 54°C as measured by CD (Supplementary Figure S3).

**I14–Tel23 adopts a basket-type G-quadruplex structure with only two G-tetrads and extensive loop interactions in K⁺ solution**

Many inter-residue NOE interactions are observed in 2D-NOESY of I14–Tel23 in K⁺. Critical inter-residue NOE interactions of I14–Tel23 are schematically summarized in Figure 5, which clearly define the overall structure of the unique G-quadruplex structure formed in I14–Tel23. I14–Tel23 forms a basket-type intramolecular G-quadruplex with two G-tetrads and multiple layers of capping structures in K⁺ solution (Figure 4A). The two G-tetrads are G2–G15–G21–G9 and G3–G16–G20–G8, with reversed arrangements of sugar glycosidic conformations, i.e. syn-syn-anti-anti and anti-anti-syn-syn, respectively. The assignment of the imino and base H8 protons of guanines leads to the direct determination of the folding topology of the G-quadruplex structure formed in I14–Tel23 in K⁺ solution (Figure 4A). In a G-tetrad plane with a Hoogsteen-type H-bond network, the imino proton NH1 of a guanine is in close spatial vicinity to the NH1s of the two adjacent guanines, and to the base H8 of one of the adjacent guanines (Figure 4A, left). Two G-tetrad planes were determined based on the NOE connectivities of exchangeable protons (Figure 3A). For example, the GNH1/GNH1 NOE interactions, including G2NH1/G15NH1, G15NH1/G21NH1, G21NH1/G9NH1, and G9NH1/G2NH1 (Figure 3A, bottom); and the GH1/GH8 NOE interactions, including G2NH1/G15H8, G15NH1/G21H8 and G21NH1/G9H8 (Figure 3A, top), define the G-tetrad plane of G2–G15–G21–G9 (Figure 4A). Similarly, G3NH1/G8NH1, G8NH1/G20NH1, G20NH1/G16NH1, G16NH1/G3NH1 and G3NH1/G8NH1, G8NH1/G20H8, G20NH1/G16H8, G16NH1/G3H8 (Figure 3A) define the G-tetrad plane of G3–G16–G20–G8 (Figure 4A). The strong NOE interactions of G2NH1/G16NH1, G15NH1/G20NH1, and G21NH1/G8NH1 (Figure 3A) indicate that the two G-tetrads have reversed G-arrangements (Figure 4A). An expanded region for base and sugar H1 protons of non-exchangeable proton NOESY with sequential assignment is shown in Figure 3B. Four guanine residues are in syn-conformation, including G2, G8, G15 and G20, as indicated by the very strong H8–H1 NOE intensities (Figure 3B). The regular sequential NOE connectivities are either missing or very weak at the N(i)- synG(i + 1) steps, i.e., A1–G2, A7–G8, I14–G15 and A19–G20. The characteristic synG(i)H8/G(i + 1)H1 NOEs are observed, including those of G2–G3, G8–G9, G15–G16 and G20–G21 (Figure 3B). A characteristic downfield shift is observed for the H2/H2′ sugar protons of the syn-guanines (Figure 3B and Table 1). For the N(i)- antiG(i + 1) steps, the sequential NOE connectivities of the base H8 protons to the 5′-flanking
Figure 3. (A) The expanded H1–H8/H2/H6 (top) and H1–H1 (bottom) regions, with labeling, of the exchangeable proton 2D JR-NOESY spectrum of I14–Tel23 in K⁺ solution at 1°C. Only the aromatic H2 protons are specified in the labels. NOEs are labeled as following: intra-tetrad NOEs are labeled in red, inter-tetrad NOEs are labeled in purple, intra-triple NOEs are labeled in green, NOEs between the G-tetrads and the stacking triple structures are labeled in blue. (B) The expanded H8/H6–H1₀ region of the non-exchangeable 2D-NOESY spectrum of I14–Tel23 in K⁺ solution at 20°C. The sequential assignment pathway is shown. The H8–H1₀ NOEs are labeled with residue names. The residue names of the guanines with syn conformation are in red. A3 and A15 show much broader peaks, likely related with the dynamic motion. Missing connectivities are labeled with arrows. The characteristic G(i)H8/G(i + 1)H1₀ NOEs for the syn G(i)s are labeled in green. Conditions: 25 mM pH 7.0 K-phosphate, 70 mM KCl, 2.0 mM DNA.
residue sugar H1'/H2'/H2" protons, typical for right-handed DNA twist, are clearly observed (Figure 3B).

Remarkably, the two G-tetrads in the I14–Tel23 G-quadruplex show extensive interactions with the loop residues. Multi-layer capping structures appear to form involving the loop residues, covering the centrally located two G-tetrads. Specifically, the G2–G15–G21–G9 G-tetrad is covered by a G10–I14–G22 triplet which is further capped by a T12–T23 base pair, while the G3–G16–G20–G8 G-tetrad is covered by a G4–A7–A19 triplet which is further capped by a T6–T18 base pair (Figure 4A). Each loop residue exhibits extensive inter-residue NOE interactions as summarized in Figure 5 and Supplementary Table S1. The four layers of the capping structures are defined by extensive NOE interactions.

We have also site-specifically labeled each guanine of I14–Tel22 and have collected a full set of 2D NMR data on I14–Tel22 to have a complete assignment of I14–Tel22 (data not shown). Our NMR data indicate that the major G-quadruplex formed in I14–Tel22 is the same as that formed in I14–Tel23. However, minor conformations are much more evident in I14–Tel22, as the G10–I14–G22 triple structure was not stably formed in I14–Tel22 due to the absence of T23.

**NOE-derived molecular structure of I14–Tel23 in K+ shows well-defined, multi-layer capping structures formed by loop residues**

Solution structures of the I14–Tel23 G-quadruplex were calculated using a NOE-restrained distance geometry (DGSA) and molecular dynamics (RMD) approach (Figure 6, PDB ID 2kka), starting from an arbitrary extended single-stranded DNA. A total of 508 NOE distance restraints, of which 276 are from inter-residue NOE interactions, were incorporated into the NOE-restrained structure calculation (Table 2). Dihedral angle restraints were used for the glycosidic torsion angle (χ) based on the experimentally determined syn and anti conformations. The structure statistics are listed in Table 2.

The I14–Tel23 in K+ forms a well-defined G-quadruplex with multi-layer capping structures formed by loop residues (Figure 6A). The two central G-tetrad layers are each covered by a middle triple structure, namely the G10–I14–G22 triple and the G4–A7–A19 triple, which are each further capped by an outer T–T base pair, i.e., T12–T23 and T6–T18. A representative model of the I14–Tel23 G-quadruplex structure in K+ is shown in two different views in Figure 6B. For the 23–nt I14–Tel23 sequence, except for the 8 guanines involved in the G-tetrad formation, there are 15 loop and flanking residues. Remarkably, 10 of the 15 loop residues are involved in the well-defined capping structures. For the remaining 5 loop residues, three thymines, T5, T11, and T17, are located in the grooves; while A1 and A13 have more flexible conformations.

A well-defined G10–I14–G22 triple structure is found to cover the G2–G15–G21–G9 tetrad, with G10 stacking over G9 and G2, I14 stacking over G2 and G15, and G22 stacking over G21 and G9 (Figure 7A). Potential H-bonds are formed within the triple structure. G10NH1 is H-bonded with I14O6, I14NH1 is H-bonded with G22O6, and G22NH1 is H-bonded with G10O6, while G10NH2b is H-bonded with I14N7, and...
G22NH2b is H-bonded with G10N7 (Figure 4B, left, and Figure 7A). The imino protons of all three residues, i.e. G10NH1, I14NH1, and G22NH1, can be clearly detected in 1H NMR up to 30°C between 10–12.5 ppm (Figure 1B and Supplementary Figure S2). The chemical shifts of the imino protons of G10 and G22 are consistent with the NH-O hydrogen bonding, while the downfield shifting of I14 is typical for an inosine base. In addition, the NOEs between the imino protons and amino protons NH2b of G10 and G22 are both observed by NMR, indicating that the amino proton of the two guanines are also involved in H-bonding (data not shown). Moreover, NOE interactions between G10, I14 and G22 are observed, such as G22NH1/G10NH8, G10NH1/I14H8, and G10NH1/I14NH1, and G10NH1/G22NH1 (Figures 3A and 5). The three residues of the G10–I14–G22 triple are well-stacked with the G2–G15–G21–G9 G-tetrad, as defined by extensive NOEs (Figure 5 and Supplementary Table S1). For example, G10 is above G9 along the G9–G2 side of the G2–G15–G21–G9 G-tetrad, as G10NH1 shows a strong NOE with G9NH1 and a weaker NOE with G22NH1 (Figures 3A and 5). I14 appears to stack above G15 along the G15–G2 side, with a strong NOE observed between I14H2/G15NH1 (Figure 3A). G22 appears to stack above G21–G9, with a strong NOE observed between G22NH1/G21NH1 and a clear NOE between G22NH1/G9NH1 (Figure 3A).

A second well-defined G4–A7–A19 triple structure is found to cover the G3–G16–G20–G8 tetrad, with G4 stacking over G3 and G16, A7 stacking over G8, and A19 stacking over G20 and G16 (Figure 7B). Potential H-bonds are formed within the G4–A7–A19 triple structure (Figure 4B, right). The imino proton of G4 is observed in 1H NMR (Figure 1B). The G4–A7–A19 triple structure is also supported by experimental NMR data (Figure 5 and Supplementary Table S1). The downfield-shifted chemical shift of G4NH1 indicates a N–H–N hydrogen bonding. In addition, the amino protons of G4 show clear NOEs with G4NH1 and appear to be H-bonded. Within the G4–A7–A19 triple, a very strong NOE is detected between G4NH1 and A19H2, and a clear NOE is detected between A7H2 and A19H2. Three residues of this triple structure, A19, A7, and G4, are all shown to stack well with the G3–G16–G20–G8 tetrad by NOE interactions (Figure 5 and Supplementary Table S1). A19H2 shows strong NOEs with G20NH1 and G16NH1, medium NOE with G3NH1 and weak NOE with G8NH1 (Figure 3A, top), indicating A19 is stacked right above the G20–G16 edge of the G3–G16–G20–G8 tetrad. A7H2 and H2 show clear NOEs with G8 and G3 (weaker) (Figure 3A, top), indicating A7 is stacked also above the G3–G16–G20–G8 tetrad but towards the G8–G3 edge. G4 shows nice stacking with G3, as indicated by inter-residue NOEs. In addition, G4NH1 shows a clear NOE with G16NH1.

In addition to the two triple structures immediately covering the two G-tetrads, each triple structure is further covered and stabilized by a T–T capping structure at each end of the I14–Tel23 G-quadruplex. Specifically, the G10–I14–G22 triple is capped by the T12–T23 base pair (Figure 7C), while the G4–A7–A19 triple is covered by the T6–T18 base pair (Figure 7D). The two T–T base pairs are also well supported by experimental NMR data. For the T12–T23 base pair, while T23 is well stacked with G22, T12 shows very good stacking with G10 (Figure 5 and Supplementary Table S1). Indeed, the presence of T23, and thus the formation of T23–T12 base pair, appears to be important in stabilizing the unique two-G-tetrad G-quadruplex. While the stabilization effect of the
T23–T12 base pair to the G10–I14–G22 triple appears to improve the stability of the overall I14–Tel23 G-quadruplex, the T23–T12 base pair is critical in stabilizing the two-G-tetrad G-quadruplex in the Tel21–T sequence (37). Similarly, T18 and T6 also appear to stack well with A19 and G4, respectively. A notable upfield-shifting is observed for the sugar and base protons of T12 and T6 (Table 1), indicating that the two residues are positioned right above the neighboring triple structure and experiencing strong ring-current effect.

The I14–Tel23 G-quadruplex can only be clearly detected in the specifically truncated telomeric sequences with no 5′-flanking. We have previously shown that Hybrid-1 and Hybrid-2 types of intramolecular G-quadruplexes co-exist in extended four-G-tract human telomeric sequences with various lengths of 5′- and 3′-flanking segments (27). It is important to find out whether this new two-G-tetrad G-quadruplex can also form in the human telomeric sequences. We have tested all four-G-tract human telomeric sequences with various lengths of 5′- and 3′-flanking TTA segments. The formation of the new two-G-tetrad G-quadruplex can be monitored by the characteristic imino peak of G10 (Figure 1B). We found that the new two-G-tetrad G-quadruplex can only be clearly detected in the truncated telomeric sequences with no 5′-flanking. A 3′-flanking thymine (T23) is critical in stabilizing the two-G-tetrads G-quadruplex. The two-G-tetrad G-quadruplex is mostly clearly detected in Tel21–T and accounts for ~60–65% of the total population (Figure 1B). This can be explained by

Figure 5. Schematic diagram of inter-residue NOE connectivities of I14–Tel23 G-quadruplex formed in K⁺ solution. The guanines in syn conformation are represented using gray circles. The NOE connectivities clearly define the G-quadruplex conformation and provide distance restraints for structure calculation.
the structural data, in that T23–T12 base pair stabilizes the G10–I14–G22 triple conformation and thus the overall two-G-tetrad G-quadruplex. However, as soon as a 5′-flanking segment is added, even a single adenine residue, the population of this two-G-tetrad form is prohibited. A–Tel21–T is the only telomeric sequence with a 5′-flanking residue (Figure 1A) in which the two-G-tetrad form can be detected; however, the population of the two-G-tetrad form is reduced to less than 10% of the total population in A–Tel21–T (Figure 1B), and is undetectable in any other four-G-tract human telomeric sequences with a 5′-flanking segment (Figure 1B and Supplementary Figure S1).

CD spectroscopic study

We have measured CD spectra for I14–Tel23, I14–Tel22, and wild-type four-G-tract telomeric sequences with various flanking segments in 100 mM K+ solution, pH 7. CD spectra of several sequences are shown in Figure 4C. I14–Tel23 and I14–Tel22 show very similar CD profiles in K+ solution. It is noteworthy that the CD profile of A–Tel21 (Figure 1A) in Na+ solution, which has also been shown to form a basket-type G-quadruplex (24),

Table 2. Structural statistics for the I14–Tel23 G-quadruplex

| NMR distance and dihedral constraints |  |
|--------------------------------------|--|
| Distance restraints                  |  |
| Total NOE                            | 508 |
| Intraresidue                         | 232 |
| Interresidue                         | 276 |
| Sequential (|i–j| = 1)                   | 184 |
| Non-sequential (|i–j| > 1)                   | 92  |
| Hydrogen bonds                       | 21  |
| Total dihedral angle restraints      | 16  |
| Structure statistics                 |  |
| NOE violations                       |  |
| numbers (>0.2Å)                      | 2.6 ± 0.9 |
| r.m.s.d. of violation (Å)            | 0.040 ± 0.014 |
| Deviations from idealized geometry   |  |
| Bond length (Å)                      | 0.007 ± 0.000 |
| Bond angle (°)                       | 1.345 ± 0.016 |
| Impropers (°)                        | 0.983 ± 0.005 |
| pairwise r.m.s.d. of heavy atoms (Å) |  |
| Two G-tetrads                        | 1.10 ± 0.16 |
| With two triads                      | 1.21 ± 0.10 |
| With two T–T base pairs              | 1.26 ± 0.10 |
| All residues                         | 1.59 ± 0.17 |

The ensemble of eight structures is selected based on the minimal energy terms and number of NOE violations.

Figure 6. (A) Stereo view of the superimposed 8 NMR-refined structures of the two-G-tetrad G-quadruplex formed in I14–Tel23 in K+ solution. The G2–G15–G21–G9 tetrad is in light pink, and the G3–G16–G20–G8 is in yellow. The guanines in the G10–I14–G22 and G4–A7–A19 triples are in magenta, I14 in orange, adenines are in red, and thymines in blue. (B) A representative model of the NMR-refined I14–Tel23 G-quadruplex structure from two different views, prepared using GRASP (43) (guanine = yellow, adenine = red, thymine = blue).
DISCUSSION

We report here a unique G-quadruplex structure formed in a variant human telomeric sequence, I14-Tel23 (Figures 6 and 7). This G-quadruplex is a basket-type structure with only two G-tetrads but multiple layers of capping structures. The extensive capping structures and loop interactions appear to stabilize the formation of this two-G-tetrad G-quadruplex. The two G-tetrads are sandwiched between the two triple capping structures, which are each further capped and stabilized by a T–T base pair (Figure 4). Specifically, the first G-tetrad G2–G15–G21–G9 (from the 5′-end of the sequence) is covered by the G10–I14–G22 triple, which is further capped by the T12–T23 base pair. The substitution of inosine for G14 of the human telomeric sequence (Figure 1A) appears to drive the formation of this two-G-tetrad G-quadruplex form, and the 3′-flanking T (T23) further stabilizes this structure, indicating that the formation of the G10–I14–G22 triple capping structure is critical for the stable formation of this two-G-tetrad quadruplex structure. The second G-tetrad G3–G16–G20–G8 is covered by the G4–A7–A19 triple, which is further capped by the T6–T18 base pair.

The four layers of the capping structures are very well defined by extensive NOE interactions (Figure 5 and Supplementary Table S1). For example, the G10–I14–G22 triple structure (Figure 7) is very well defined by the NMR experimental data. The imino protons of G10, I14 and G22 are clearly observed in 1D 1H NMR at 20°C (Figure 1B), indicating that all three imino protons are involved in H-bonding. This observation is consistent with the G10–I14–G22 triple conformation, in which G10NH1 is H-bonded with I14O6, I14NH1 is H-bonded with G22O6, and G22NH1 is H-bonded with G10O6 (Figure 4B). The G10–I14–G22 triple structure appears to be quite stable with the imino proton of G10 detectable even at 50°C (Supplementary Figure S2). A similar structure was very recently reported using a 8-Br-G-substituted Tel21–T sequence (BrG7–Tel22) in K+ (37). The structure formed by BrG7–Tel22 is also a basket-type two-G-tetrad G-quadruplex with two triple capping structures; however, our structure differs from that structure in the capping structures and loop interactions in several aspects. While both structures showed two capping triples, the conformations of both triples in our structure differ from that reported for BrG7–Tel22. Specifically, for the G10–I14–G22 triple, the conformation of I14 in our structure is different from G13 of the equivalent G9–G13–G21 triple in the structure reported for BrG7–Tel22 (37). Note that our numbering system is based on Tel 23 (A–Tel21–T, see Figure 1A) and thus is one number larger. G13NH1 is not shown to form any H-bond in the BrG7–Tel22 structure. However, this imino proton is clearly detected at ~11.2 ppm at 20°C for the native Tel22 sequence (Supplementary Figure S1, G14 of Tel21–T). In addition, the remarkable stability of G10NH1 [also observed for the equivalent G9NH1 of BrG7–Tel22 (37)] may be better explained by the G10–I14–G22 triple conformation in our structure (Figure 4B), in which the additional hydrogen bonds formed by G10NH2:G14N7 and I14NH1:G22O6 can stabilize the G10NH1:I14O6 hydrogen bond. For the G4–A7–A19 triple, the conformation of A7 in our structure is different from the...
G16 in the basket-form (stabilized by specific loop interactions of a specific sequence. For example, the presence of I14 loosens the H-bonding connection between G14 and D2 in the basket-form (stabilized by specific loop interactions of a specific sequence. For example, the presence of I14 loosens the H-bonding connection between G14 and D2 in the basket-form (Figure 8). Schematic diagram of a proposed model for interconversions between the Na⁺ basket form (A) and the K⁺ hybrid form of telomeric G-quadruplexes (D and D2). Guanines with known anti conformation are in red, and guanines with syn conformation are in purple. The A–Tel21 forms a single stable basket-type intramolecular G-quadruplex in Na⁺ solution. In the presence of K⁺ the basket form (A) is not favored and A–Tel21 can convert to the hybrid forms (D and D2) through a strand-reorientation mechanism (C and C2). The two-G-tetrad conformation is likely to be the intermediate state of this strand reorientation and may be stabilized by specific loop interactions of a specific sequence. For example, the presence of I14 loosens the H-bonding connection between G14 and G2 in the basket-form (A) and thus frees the 5'-end for its sliding-up dissociation from the core G-tetrads. However, this strand sliding-up reorientation is captured at the two-G-tetrad stage (B) due to the stable formation of the G10–I14–G22 capping structure. A similar sliding (B2) and dissociations (C2) could also potentially happen at the 3'-end of the basket-form structure to convert to the hybrid-2 form G-quadruplex by the 3'-end swing-back (D2). The two-G-tetrad form may also be an intermediate state of the interconversion between the two hybrid-type forms (D+C+B+B2+C2+D2 circle).

![Figure 8](image-url)

The G-strand of the two-G-tetrad form is abolished (Supplementary Figure S1). A–Tel21–T is the only telomeric sequence with a 5'-flanking segment (Figure 4B) in which the two-G-tetrad form can be weakly detected; however, the population of the two-G-tetrad form is less than 10% in A–Tel21–T. The two-G-tetrad form cannot be detected in any other four-G-tract human telomeric sequences with a 5'-flanking segment (Figure 1B and Supplementary Figure S1). Therefore, this two-G-tetrad form is unlikely to be a general form of the G-quadruplex formed in the extended human telomeric sequences. Instead, our results suggest that this two-G-tetrad conformation is likely to be an intermediate transition form of the interconversion between different telomeric G-quadruplex conformations (Figure 8). We have previously proposed a model for the interconversion between the hybrid-type K⁺ form and the basket-type Na⁺ form telomeric G-quadruplex structures through a strand-reorientation mechanism (26). The 5'-G-strand of

Most importantly, our results show that this structure can only be detected in the specifically truncated telomeric sequences without any 5'-flanking residues, and thus do not support the conclusion by Lim et al. that the two-G-tetrad structure is a stable new form for the extended human telomeric sequences (37). As soon as a 5'-flanking segment is present, even with a single adenine residue, the formation of this two-G-tetrad form is
the basket-type G-quadruplex (Figure 8A) may dissociate from the core G-tetrads (Figure 8C) and swing back to the other side of the second G-stand to form a parallel-stranded structural motif with a double-chain-reversal loop (Figure 8D). Intriguingly, the two-G-tetrad form appears to fit the intermediate state between the transition from Figure 8A to C: when the 5'-strand is sliding up to dissociate from the core G-tetrads, a two-G-tetrad conformation can be formed transiently, which may then be stabilized by specific loop interactions of a specific sequence (Figure 8B). For example, in the case of I14–Tel23, the substitution of G14 by inosine appears to first destabilize the 5'-G-tetrad of the three G-tetrad form and subsequently capture the formation of the two-G-tetrad form. Specifically, as inosine lacks the amino group at C2, the G-to-I substitution at G14 loosens the H-bonding connection between G14 and G2 in the 5'-G-tetrad of the three G-tetrad basket form (Figure 8A). The 5'-end of the basket-type form becomes more flexible and can start to slide up, which is subsequently captured at the two-G-tetrad stage (Figure 8B) due to the stable formation of the G10–I14–G22 capping structure (which is further stabilized by the presence of T23 due to the T23–T12 capping structure). Interestingly, the glycosidic conformations of the tetrad guanines in the two-G-tetrad form are the same as those in the hybrid-1 form (Figure 8D); thus the presence of I14 appears to loosen up the 5'-end and favor the 5'-end dissociation (Figure 8B) for the conversion to the hybrid-1 form by the 5'-end swing-back rearrangement. A similar sliding (Figure 8B2) and dissociation (Figure 8C2) process could also happen at the 3'-end of the basket-form structure for the conversion to the hybrid-2 form G-quadruplex by the 3'-end swing-back rearrangement mechanism (Figure 8D2). In addition, the two-G-tetrad form may also be an intermediate state of the interconversion between the two hybrid-type forms (Figure 8). As previously shown, while the energy difference of the two hybrid-forms is quite small so that the two forms can co-exist in the extended telomeric sequences, the kinetics of the interconversion between the two forms appears to be rather slow, which cannot be detected on the NMR time scale (ms) (27), indicating the energy barrier of the intermediate state(s) is likely to be high. In addition, it has been shown that the unfolding processes of the telomeric G-quadruplexes have multiple intermediate states (35,36). Taken together, a possible model may be proposed for the interconversion of the two hybrid forms with multiple intermediate states, including the two-G-tetrad forms (Figure 8, D ↔ C ↔ B ↔ B2 ↔ C2 ↔ D2).

It is noteworthy that the CD profile of this basket-type two-G-tetrad G-quadruplex is very different from the CD profile of the basket-type telomeric G-quadruplex formed by A–Tel21 in Na+ solution (Figure 4C). In addition, the CD melting profile shows that the melting temperature of the I14–Tel23 quadruplex is about 54°C (Supplementary Figure S3). The two-G-tetrad form does not appear to be more stable than the hybrid forms, as suggested in the previous report (37). Although the Tel21–T sequence shows a higher melting temperature, the increased Tm is likely caused by the presence of multiple structures, as evident by its 1D 1H NMR spectrum (Figure 1B). Furthermore, the A–Tel21 sequence, which does not appear to form the two-G-tetrad structure (Figure 1B), shows a similarly increased melting temperature (Supplementary Figure S3).

ACCESSION NUMBER
PDB ID 2kka.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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