Stability of telomeric G-quadruplexes

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ABSTRACT

In most eukaryotes, telomeric DNA consists of repeats of a short motif that includes consecutive guanines and may hence fold into G-quadruplexes. Budding yeasts have telomeres composed of longer repeats and show variation in the degree of repeat homogeneity. Although telomeric sequences from several organisms have been shown to fold into G-quadruplexes in vitro, surprisingly, no study has been dedicated to the comparison of G-quadruplex folding and stability of known telomeric sequences. Furthermore, to our knowledge, folding of yeast telomeric sequences into intramolecular G-quadruplexes has never been investigated. Using biophysical and biochemical methods, we studied sequences mimicking about four repetitions of telomeric motifs from a variety of organisms, including yeasts, with the aim of comparing the G-quadruplex folding potential of telomeric sequences among eukaryotes. G-quadruplex folding did not appear to be a conserved feature among yeast telomeric sequences. By contrast, all known telomeric sequences from eukaryotes other than yeasts folded into G-quadruplexes. Nevertheless, while G3T1-4A repeats (found in a variety of organisms) and G4T2A repeats (found in ciliates) folded into stable G-quadruplexes, G-quadruplexes formed by repetitions of G2T2A and G2CT2A motifs (found in many insects and in nematodes, respectively) appeared to be in equilibrium with non-G-quadruplex structures (likely hairpin-duplexes).

INTRODUCTION

Telomeres are specialized nucleoprotein complexes that cap and protect the extremities of linear eukaryotic chromosomes. In most eukaryotes telomeric DNA consists of a tandem array of a short motif of 5–8 nt, that includes two, three or four consecutive guanines, with an impressive prevalence of the G3T2A motif (1). The G3T2A motif is found in many phylogenetically distant organisms, including vertebrates (2,3), Trypanosoma brucei (4,5), several fungi (6), slime moulds (7,8) and several aquatic invertebrates (9). Variants of this motif are found in many other organisms: G3TA in the parasite Giardia intestinalis (10); G3T3A in a variety of plants (11), such as Arabidopsis thaliana (12); G3T4A in the unicellular green alga Chlamydomonas reinhardtii (13); G3T2A in the silkworm Bombyx mori and in many other insects (14); G2CT2A in the nematodes Ascaris lumbricoides (15), Caenorhabditis elegans and in other nematodes (16,17). Telomeric motifs bearing four consecutive guanines are found in ciliates: G4T2 in holotrichous ciliates such as Tetrahymena (18) and G4T4 in hypotrichous ciliates such as Oxytricha. Euplotes and Stylonychia (19). Degenerated telomeric motifs have been reported for several organisms: GGGTTTA and GGGTTCA in Plasmodium species (20,21); GGGTTTA and GGGTTA in Lycopersicon esculentum (tomato plant) (22); GGTTA and GGGTTT in Paramecium species (23,24), where a single telomerase RNA, encoding G1T2 repeats, has been identified (25).

Unlike in most eukaryotes, telomeres have rapidly evolved among budding yeasts; these organisms possess the most variable set of telomeric repeats (26,27). A number of budding yeast species have telomeres composed of long repeats (8–26 bp) and show variations in the degree of repeat homogeneity (28,29). The budding yeast Saccharomyces cerevisiae has short heterogeneous telomeric repeats of the general form TG2-3(TG)1-6 (30,31), encoded by a single telomerase RNA (32); the most abundant type of repeat is TG3TG (29). G3T2A repeats are found in the immediate subtelomeric region of S. cerevisiae, suggesting that G3T2A-like telomeres were present prior to yeast divergence (26). Despite
G-quadruplex folding potential of telomeric motifs has been emphasized since early studies (46,47). Several G-quadruplex structures formed by telomeric repeats from different organisms have been resolved, notably the intramolecular G-quadruplexes formed by Oxytricha and Tetrahymena telomeric sequences in sodium (48–50), by human (vertebrate) telomeric sequences in sodium and in potassium (51–58) and by Bombyx mori and Giardia modified telomeric sequences in potassium (59,60) (Figure 1).

Although several telomeric sequences have been shown to fold into G-quadruplexes, data have been acquired with different techniques and under different conditions, hence do not allow a straightforward comparison of G-quadruplex folding and stability of telomeric sequences among eukaryotes. Furthermore, to our knowledge, folding of yeast telomeric sequences into G-quadruplexes has never been proved, with the exception of sequences mimicking two or three repetitions of S. cerevisiae heterogeneous telomeric motifs (46,61). Using biophysical and biochemical methods, we undertook an investigation of sequences consisting of about four repetitions of telomeric motifs from a variety of organisms (the minimal length required to allow formation of intramolecular G-quadruplexes); the aim was to (i) probe their propensity to fold into G-quadruplexes, (ii) provide some information about their folding and (iii) study/compare their stability. The studied sequences are listed in Table 1. Among yeasts, we chose to limit our study to S. cerevisiae and S. pombe (which are model organisms in the study of telomeres), to Candida glabrata and Candida guillermondii. Among the diverse telomeric sequences of budding yeasts (other than species of the genus Saccharomyces), C. glabrata and C. guillermondii telomeric sequences present the shortest loops (6 nt) connecting consecutive guanines (28,29); they should hence be the most prone to fold into G-quadruplexes. Actually, several studies on model sequences indicate that increasing loop length leads to a decrease in G-quadruplex stability (62–64). S. cerevisiae and S. pombe have heterogeneous telomeric sequences; we chose to study repetitions of the most frequently found motifs. In a previous study, we provided evidence of folding into stable G-quadruplexes by repetitions of Plasmodium degenerated telomeric motif (GGGT TYA, where Y is T or C) under the same conditions used here (65); melting temperatures and thermodynamic data from this work were included in Table 1.

MATERIAL AND METHODS

Oligonucleotides

Reverse-Phase Cartridge Gold™ (RPC) purified oligonucleotides were purchased from Eurogentec (Seraing, Belgium), dissolved in bi-distilled water at a concentration of 200 μM and stored at −20°C. Concentrations were determined by ultraviolet (UV) absorption using the extinction coefficients provided by the manufacturer. For spectroscopic measurements, RPC-purified oligonucleotides were used without further purification; for non-denaturing PAGE experiments, oligonucleotides
Table 1. Sequences, melting temperatures and thermodynamic parameters for G-quadruplex folding of the studied oligonucleotides

| Sequence name (species) | Sequence (from S' to 3') | NaCl | KCl |
|-------------------------|--------------------------|------|-----|
|                         |                          | $T_m$ (°C) | $\Delta H^\circ$ (kcal mol$^{-1}$) | $\Delta S^\circ$ (cal K$^{-1}$ mol$^{-1}$) | $\Delta G^\circ$ (310 K) (kcal mol$^{-1}$) |
| $G_2$ motifs            |                          |      |     |     |       |
| Bom17 (Bombyx)          | (GGTTA)$_2$GG            | a    | a   |     |       |
| Asc20 (Ascaris)         | (GGCTTT)$_2$GG          | a    | a   |     |       |
| Spom20 (S. pombe)       | (GGTTACA)$_2$GG         | no $G_4^b$ |     |       |
| Spom23 (S. pombe)       | (GGTTACA)$_2$GG         | no $G_4^b$ |     |       |
| Cgi26 (C. guillermondii)| (GGTTACT)$_2$GG         | no G4 |     |       |
| $G_3$ motifs            |                          |      |     |     |       |
| Gia18 (Giardia)         | (GGGTTA)$_2$GG          | 60   | $-53 \pm 7$ | $-157 \pm 19$ | $-3.7 \pm 0.6$ | 68 | $-63 \pm 4^f$ | $-185 \pm 12^f$ | $-5.7 \pm 0.4^f$ |
| Hum21 (H. sapiens)      | (GGGTTA)$_2$GG          | 59   | $-55 \pm 4$ | $-167 \pm 12$ | $-3.7 \pm 0.4$ | 65 | $-60 \pm 3$ | $-177 \pm 9$ | $-5.0 \pm 0.3$ |
| Par21 (Paramecium$^e$) | (GGGTTT)$_2$GG          | 50   | $-53 \pm 8$ | $-162 \pm 23$ | $-2.2 \pm 0.5$ | 64 | $-64 \pm 10^f$ | $-189 \pm 29^f$ | $-5.2 \pm 0.9^f$ |
| Scer21 (S. cerevisiae)  | (GGGTTT)$_2$GG          | 50   | $-48 \pm 5$ | $-148 \pm 15$ | $-1.9 \pm 0.2$ | 68 | $-68 \pm 8^f$ | $-200 \pm 25^f$ | $-6.2 \pm 0.9^f$ |
| Ara24 (Arabidopsis)    | (GGGTTT)$_2$GG          | 57   | $-60 \pm 5$ | $-179 \pm 14$ | $-3.6 \pm 0.4$ | 64 | $-73 \pm 10$ | $-216 \pm 28$ | $-5.7 \pm 0.9$ |
| Plasmodium$^e$          | (GGGTTA)$_2$GG, Y=T or C | 55   | $-53 \pm 4$ | $-158 \pm 11$ | $-2.8 \pm 0.4$ | 64 | $-63 \pm 5$ | $-186 \pm 14$ | $-4.9 \pm 0.6$ |
| Tom24 (L. esculentum$^e$) | (GGGTTA)$_2$GG         | 51   | $-54 \pm 4$ | $-168 \pm 13$ | $-2.3 \pm 0.2$ | 64 | $-77 \pm 5$ | $-228 \pm 13$ | $-6.1 \pm 0.4$ |
| Chla27 (Chlamydomonas)  | (GGGTTT)$_2$GG          | 46   | $-56 \pm 4$ | $-176 \pm 13$ | $-1.6 \pm 0.3$ | 57.5 | $-68 \pm 8$ | $-207 \pm 24$ | $-4.2 \pm 0.6$ |
| $G_4$ motifs            |                          |      |     |     |       |
| Tet22 (Tetrahymena)     | (GGGTTT)$_2$GG          | 64   | $-54 \pm 8$ | $-161 \pm 24$ | $-4.4 \pm 0.7$ | >80 |       |       |       |
| Oxy28 (Oxytricha)       | (GGGTTT)$_2$GG          | 66   | $-82 \pm 13$ | $-242 \pm 38$ | $-6.9 \pm 1.3$ | >80 |       |       |       |
| $G_3/G_4$ motif         |                          |      |     |     |       |
| Gla26 (C. glabrata)     | G$_3$T(GTG)$_2$G$_3$ | 48   | $-53 \pm 12$ | $-164 \pm 36$ | $-1.8 \pm 0.5$ | 63 | $-70 \pm 12$ | $-206 \pm 34$ | $-5.4 \pm 0.8$ |

The number in the sequence name denotes the sequence length (in nucleotides); telomeric motifs are in italic font.

Melting temperatures ($T_m$) reported in this table depended neither on wavelength nor on oligonucleotide strand concentration (3, 10 and 30 μM).

Melting curves at 295 nm were analysed according a two-state equilibrium model and assuming linear low- and high-temperature absorbance baselines. Standard enthalpy and entropy changes ($\Delta H^\circ$ and $\Delta S^\circ$) for folding were determined by linear fitting ln$K$ versus $1/T$, where $K$ is the equilibrium constant between the folded and the unfolded state; standard Gibbs free energy changes ($\Delta G^\circ$) were extrapolated at 310 K, from the relation $\Delta G^\circ (T) = \Delta S^\circ - T \Delta S^\circ$. $\Delta H^\circ$, $\Delta S^\circ$ and $\Delta G^\circ$ reported in this table are the mean values obtained from analysis of melting curves at 3 and 30 μM strand concentration upon varying linear low- and high-temperature absorbance baselines ± maximum deviation.

The presence of a G-quadruplex and of a non-G-quadruplex competing structure made $T_m$ determination not straightforward.

$^a$Not determined: $T_m$ of Spom20 in KCl could not be determined accurately because of the incertitude in low-temperature absorbance baseline.

$^b$Paramaecium and L. esculentum (tomato plant) have degenerated telomeric motifs: GGGKTT in Paramaecium (K=T or G) and GGGTTWA (W=A or T) in tomato plant; for W=T, the tomato telomeric motif comes back to the Arabidopsis one; for K=G, the Paramaecium telomeric motif comes back to the Tetrahymena one.

$^c$Data from a previous study (65) (mean values from analysis of the eight possible variant sequences); for Y=T, the Plasmodium telomeric sequence comes back to the Arabidopsis one.

$^d$For Gia18, Par21 and Scer21 in KCl at 30 μM strand concentration a two-state model is not appropriate, as revealed by non-denaturing PAGE; nevertheless linear van’t Hoff graphs were obtained.

$^e$Data from a previous study (65) (mean values from analysis of the eight possible variant sequences); for Y=T, the Plasmodium telomeric sequence comes back to the Arabidopsis one.
were purified by PAGE according to standard protocols. For Asc20, Gia18 and Tet22, spectroscopic measurements were carried out on two different batches of RPC purified oligonucleotides. For Gia18, spectroscopic measurements were also carried out on a third batch, PAGE purified by the manufacturer; data reproducibility was excellent. The quality of Gia18 oligonucleotides (RPC and PAGE purified) was checked by denaturing PAGE.

**Thermal difference spectra, UV-melting curves and circular dichroism**

UV-absorbance measurements were acquired on an Uvikon 940 (Kontron) spectrophotometer or, for tomato plant and *Paramecium* variant sequences, on an Uvikon XL (Secomam). For melting experiments, samples were heated at 92°C for a few minutes, cooled from 92 to 2°C at a rate of 0.2°C min⁻¹, kept at 2°C for 30 min and heated from 2 to 92°C at a rate of 0.2°C min⁻¹. The temperature was varied with a circulating water bath; evaporation at high temperatures and condensation at low temperatures were prevented by a layer of mineral oil and by a dry air flow in the sample compartment, respectively. Temperature was measured with an inert glass sensor immersed into a water-filled quartz cell or, for tomato plant and *Paramecium* variant sequences, with a sensor embedded in a solid cuvet. The absorbance was monitored at 245, 260, 273, 295 and 405 nm; the absorbance at 405 nm was subtracted from each wavelength. For each melting curve, the melting temperature (T_m) was graphically determined as the intercept between the melting curve and the median line between low-temperature and high-temperature absorbance linear baselines (66). For each sample, the thermal difference spectrum (TDS) was obtained by subtracting the absorbance spectrum at 2°C from the one at 92°C (67). The spectrum at 92°C was recorded after heating the sample at 92°C for a few minutes, whereas the spectrum at 2°C was recorded after annealing from 92 to 2°C at a rate of 0.2°C min⁻¹. Circular dichroism (CD) spectra were recorded on a Jasco J-815 spectropolarimeter or, for tomato plant and *Paramecium* variant sequences, on a Jasco J-815, at 4°C, after annealing from 92°C at a rate of 0.2°C min⁻¹. Each spectrum was obtained by averaging three scans at a speed of 500 nm min⁻¹ and by subtracting the contributions of the solvent and quartz cell. All experiments (melting, TDS and CD) were carried out at 3, 10 and 30 µM oligonucleotide strand concentration (±5%), in a cacodylic acid buffer (10 mM) at pH 7.2 (adjusted with LiOH), containing KCl (100 mM) or NaCl (100 mM), using 1 cm (for 3 and 10 µM strand concentration) or 0.2 cm (for 30 µM strand concentration) path-length quartz cells. Stock solutions (200 µM) were heated at 90°C for a few minutes immediately before sample preparation. TDS, UV-melting and CD experiments were carried out at least twice.

**Non-denaturing PAGE**

Stock solutions (200 µM) of PAGE-purified oligonucleotides were heated at 90°C for a few minutes immediately before sample preparation. Oligonucleotides (30 µM strand concentration) were annealed from 92 to 2°C at a rate of 1°C min⁻¹ in the same solvents used for spectroscopic measurements (10 mM cacodylic acid buffer, pH 7.2 adjusted with LiOH, 100 mM NaCl or 100 mM KCl). Samples, supplemented with sucrose (11.5%), were then loaded (10 µl) into a 16.5 cm × 14.5 cm × 0.1 cm 12% polyacrylamide gel (acrylamide: bisacrylamide 19:1, prepared in a TBE buffer supplemented with 20 mM NaCl or KCl). Electrophoresis was run in a cold room at 4°C, for 2 h and 30 min, at 3 W per gel, in a TBE buffer supplemented with NaCl (20 mM) or KCl (20 mM); the temperature attained in the gels was about 15°C. DNA was detected by UV-shadow at 254 nm with a Gel:BOX (Syngene). An oligothymidylate marker (dT21) and two double-stranded markers of 9 and 12 bp (ds9: 5’GGGTATCGGG3’ + 5’CCGATACGC3’; ds12: 5’GGGTTTGTTTGGT3’ + 5’CCGAGGTACGC3’) were loaded into the gels to provide internal migration markers. We point out that dTn oligonucleotides are not appropriate markers for single-strand DNA migration in PAGE (68).

**Fluorescence resonance energy transfer melting curves**

The melting of the two double-dye labelled oligonucleotides FBOM17T and FASC20T was followed by fluorescence resonance energy transfer (FRET) on a FluoroMax-3 spectrophluorimeter (FBOM17T: FAM-5’GGTTAGGTTAGG3’-TAMRA, FASC20T: FAM-5’GGGTTAGGTTAGG3’-TAMRA, FAM: 6-carboxyfluorescein, TAMRA: carboxytetramethylrhodamine, purchased from Eurogentec, Seraing, Belgium). The temperature was raised from 5 to 90°C with a circulating water bath; FAM emission (excitation at 470 nm, emission at 520 nm) was recorded at intervals of 1°C, after an equilibration time of 30 s. FAM emission versus temperature plots were normalized between 0 and 1 and the temperature of half-dissociation (T_1/2), corresponding to an emission value of 0.5, was determined. FRET-melting experiments were carried out at 0.2 µM oligonucleotide strand concentration, in a cacodylic acid buffer (10 mM) at pH 7.2 (adjusted with LiOH), containing NaCl (100 mM) or KCl (100 mM).

**RESULTS**

**Thermal difference spectra**

A simple method to have an insight into the structure adopted by a nucleic acid sequence is to record its absorbance spectra at high temperature (unfolded form) and at low temperature (folded form). The difference between these spectra, called thermal difference spectrum (TDS), provides a spectroscopic signature for nucleic acid structures (67,69).

TDS spectra are shown in Figure 2. With the exception of the C. guilliermondii (Cgi26) and of the S. pombe (Spom20 and Spom23) sequences, all the studied telomeric sequences displayed TDS signatures characteristic of G-quadruplex structures, both in NaCl and in KCl: two major positive peaks around 240 and 270 nm and a negative peak around 295 nm.
The C. guillermondii sequence (Cgi26) neither in NaCl nor in KCl displayed a G-quadruplex-type TDS signature. Concerning the S. pombe sequences (Spom20 and Spom23), an unambiguous TDS G-quadruplex signature was obtained only for Spom20 in KCl. TDS of Spom20 in NaCl and Spom23 in NaCl and KCl exhibited a very small negative peak at 295 nm (autoscaled TDS are shown in Figure S1).

**CD spectra**

CD spectra were recorded at 4°C, a temperature at which the equilibrium was strongly shifted towards the folded forms, as shown by thermal melting profiles (Figures S2 and S3). CD spectra at 3 μM oligonucleotide strand concentration are shown in Figure 2. The Bombyx (Bom17), Ascaris (Asc20) and Oxytricha (Oxy28) sequences exhibited a negative band around 260 nm and a positive band around 290 nm (a ‘Type II’ signature), both in NaCl and in KCl. All the other G-quadruplex folding sequences (i.e. sequences displaying G-quadruplex TDS) exhibited distinct CD signatures in NaCl and in KCl: a Type II signature in NaCl (with the exception of Tet22), and a positive peak around 290 nm and a positive peak or shoulder in the 250–270-nm region in KCl; this supports structural differences in NaCl and in KCl.

With the exception of the Giardia (Gia18), Tetrahymena (Tet22) and S. cerevisiae (Scer21) sequences in KCl, the CD spectra of all the other sequences were independent from oligonucleotide strand concentration (3, 10 or 30 μM), both in NaCl and in KCl. Conversely, the CD spectra of Giardia (Gia18), Tetrahymena (Tet22) and S. cerevisiae (Scer21) sequences in KCl strongly depended on oligonucleotide strand concentration (Figure 3). For Gia18 and Tet22 increasing the oligonucleotide concentration from 3 to 30 μM led to the appearance of a CD positive peak at about 260 nm (Figure 3A and B). For Scer21, increasing the concentration led to a change in the ratio r between the CD peak at about 260 nm and the one at about 290 nm: (r < 1 for 3 μM, r ≈ 1 for 10 μM and r > 1 for 30 μM) (Figure 3C). The dependence of Gia18, Tet22 and Scer21 CD spectra on strand concentration in KCl supports the presence of multi-stranded structures.

Concerning C. guillermondii (Cgi26) and S. pombe (Spom20 and Spom23) sequences, only the S. pombe sequence Spom20 in KCl displayed an unambiguous Type II G-quadruplex CD signature, consistently with the results obtained by TDS.

Hence, TDS provided evidence of G-quadruplex folding for most of the sequences listed in Table 1; CD allowed (i) highlighting structural differences in NaCl and in KCl for most of the sequences (at least for Gia18, Hum21, Par21, Scer21, Ara24, Tom24, Chla27, Tet22 and Gla26) and (ii) providing evidence of the presence of multi-stranded structures for a few sequences (Gia18, Tet22 and Scer21 in KCl). Individual CD spectra will not be discussed in detail for the following reason. Empirically, two types of CD spectra are considered indicative of DNA G-quadruplexes: spectra with a negative band around 240 nm and a dominant positive band around 260 nm (‘Type I’ spectra), and spectra with a negative band around 260 nm and a positive band around 290 nm (‘Type II’ spectra). Theoretical calculations give an insight into the origin of Type I and Type II signals:
stacking of two G-quartets with the same hydrogen-bond orientation originates a Type I signal, stacking of two G-quartets with opposite hydrogen-bond orientation originates a Type II signal (70,71) (G-quartet hydrogen-bond orientation is illustrated in Figure 1). Nevertheless, G-quadruplexes are highly polymorphic and these empirical and theoretical evidences do not allow an unambiguous interpretation of CD spectra of potential G-quadruplex folding sequences.

UV melting experiments

TDS allowed verifying G-quadruplex folding at low temperature. To study stability, we recorded melting profiles at wavelengths of absorbance maximum variations upon heating, determined by TDS, i.e. 245, 273 and 295 nm. All melting profiles were reversible (melting curves at 295 nm are reported in Supplementary Figures S2 and S3). Melting transitions of all G-quadruplex folding sequences presented a single transition (a single first derivative peak) and occurred (with the exception of Spom20 in KCl) in a relatively narrow range of temperature (about 30°C for Bom17 and Asc20 and about 20°C for the other sequences), thereby the graphically determined melting temperatures \( T_{m(s)} \) did not significantly depend on the choice of the high- and low-temperature absorbance linear baselines; \( T_{m(s)} \) could then be determined with a precision of \( \pm 1°C \) or better.

\( T_{m(s)} \) of sequences bearing G3 and G4 motifs (included Gia26) were higher in potassium than in sodium: \( T_{m(s)} \) ranged from 46°C (Chia27) to 64–66°C (Tet22 and Oxy28) in NaCl and from 57.5°C (Chal27) to >80°C (Tet22 and Oxy28) in KCl; they did not depend on wavelength (245, 273 and 295 nm) and are reported in Table 1. The flatness of the S. pombe Spom20 melting profile in KCl at 295 nm (Supplementary Figures S2 and S3) prevented us from determining an accurate \( T_{m} \) (incertitude in the determination of a low-temperature absorbance baseline); its \( T_{m} \) was roughly estimated to be around 27°C. Bombyx (Bom17) and Ascaris (Asc20) differed from all the other G-quadruplex folding sequences for several features. (i) Surprisingly, the graphically determined \( T_{m(s)} \) at 295 nm were higher in sodium than in potassium: 46°C in NaCl and 44°C in KCl for Bom17, 41°C in NaCl and 39°C in KCl for Asc20. The maxima of the first derivatives of their melting curves (\( T_{max} \)) showed the same trend than melting temperatures determined by graphical analysis:

\[ T_{max}(\text{NaCl}) > T_{max}(\text{KCl}). \]

(ii) Asc20 \( T_{m(s)} \)s depended on wavelength (\( T_{m(s)} \)s at 245 nm were around 45°C both in sodium and potassium); this feature suggests the presence of a competing structure. (iii) The amplitude of Asc20 melting transition at 295 nm in NaCl was about half of the amplitude in KCl (Figure 4A); Bom17 melting transitions at 295 nm in NaCl and in KCl had nearly equal amplitudes (Figure 4B). Conversely, for all G-quadruplexes with \( T_{m(\text{NaCl})} < T_{m(\text{KCl})} \), the amplitudes of melting transitions at 295 nm in NaCl were larger than in KCl (Figures S2 and S3) (this feature appears to be characteristic of many other G-quadruplexes).

Finally, for all the sequences, \( T_{m} \) did not depend on oligonucleotide strand concentration (3, 10 and 30 μM), not even for those sequences displaying concentration-dependent CD spectra, i.e. Gia18 and Scer21 in KCl (Tet22 \( T_{m} \) in KCl was too high to be determined). In particular, we verified that \( T_{m} \) of Gia18 in KCl did not vary upon increasing oligonucleotide strand concentration up to 100 μM (data not shown). The independence of \( T_{m} \) from oligonucleotide strand concentration suggests formation of intramolecular structures.

FRET melting experiments

The ‘non-classical’ behaviour of Bombyx (Bom17) and Ascaris (Asc20) prompted us to a deeper investigation. We studied the double-dye labelled sequences FAsc20T (FAM-5(GGCTTA)3GG3-TAMRA) and FBom17T (FAM-5(GGTTA)3GG3-TMRA), FAsc20T neither in NaCl nor in KCl folded into G-quadruplexes, as revealed by CD spectroscopy (Figure 4C); it folded into a structure where the two dyes were close to each other, the stability of which \( (T_{1/2} = 54°C) \) did not depend on the cation (very likely a hairpin-duplex), as revealed by thermal melting followed by FRET (Figure 4E). Conversely, FBom17T folded into G-quadruplexes, as revealed by CD spectroscopy (Figure 4D), more stable in potassium \( (T_{1/2} = 55°C) \) than in sodium \( (T_{1/2} = 45°C) \) (Figure 4F).

Gel electrophoresis

Non-denaturing PAGE was carried out at 30 μM oligonucleotide strand concentration, i.e. the highest concentration used for spectroscopic measurements. DNA was detected by UV-shadow (Figure 5). The temperature of

![Figure 3. Circular dichroism spectra at 4°C of (A) Gia18, (B) Tet22 and (C) Scer21 in KCl at 3, 10 and 30 μM oligonucleotide strand concentration (circles, crosses and triangles, respectively).](image-url)
Figure 4. (A and B) Thermal melting (cooling and heating) followed by absorbance at 295 nm of (A) Asc20 and (B) Bom17 in NaCl (circles) and KCl (triangles), at 3 μM oligonucleotide strand concentration. (C and D) CD spectra at 4°C of (C) FAsc20T and (D) FBom17T in NaCl (circles) and KCl (triangles), at 3 μM oligonucleotide strand concentration. (E and F) Thermal melting (heating) followed by FRET (excitation at 470 nm, emission at 520 nm) of (E) FAsc20T and (F) FBom17T in NaCl (circles) and KCl (triangles), at 0.2 μM oligonucleotide strand concentration.

Figure 5. Non-denaturing PAGE in (A) NaCl and (B) KCl at 30 μM oligonucleotide strand concentration. dT21 is an oligothymidylate marker; dx9 and dx12 are two double-stranded markers of 9 and 12 bp, respectively. Oligonucleotides were detected by UV-shadow.
the gels was about 15°C; at this temperature, all the G-quadruplex folding sequences (i.e. all the sequences except Spom20, Spom23 and Cgi26 in NaCl and Spom23 and Cgi26 in KCl) were folded both in KCl and in NaCl, as could be inferred by the inspection of melting profiles (Supplementary Figures S2 and S3). We used an oligothymidylate marker (dT21) and 9- and 12-bp duplexes (dx9 and dx12, respectively) as internal migration standards.

In NaCl, a single band was detected for all the telomeric sequences (Figure 5A). All the oligonucleotides displayed a higher mobility than dT21; most of them had a mobility intermediate between the ones of dx9 and dx12. In NaCl, the mobility of the sequences displaying a TDS G-quadruplex signature (i.e. all the sequences except Spom20, Spom23 and Cgi26) slightly decreased with the length of the sequence, with the exception of Asc20 and Tet22. The yeast Spom20, Spom23 and Cgi26 sequences, which did not fold into G-quadruplexes in NaCl, as revealed by TDS, were the slowest migrating species.

In KCl, the migration patterns were more complex (Figure 5B). Most of the sequences (Hum21, Bom17, Asc20, Tom24, Ara24, Glab26, Chla27, Spom20, Spom23 and Cgi26) migrated as single bands and faster than dT21. Among these single-band migrating sequences, Spom23 and Cgi26 had the lowest electrophoretic mobility: according to spectroscopic measurements in KCl, these two sequences are not folded into G-quadruplexes. The *Giardia* (Gia18) and *Tetrahymena* (Tet22) sequences displayed a fast- and a slow-migrating band, the latter migrating slower than the internal migration standard dT21. For the *Paramecium* (Par21) and the *Oxytricha* (Oxy28) sequences, two bands were detected, both fast migrating as compared to dT21. In particular, the two bands of Oxy28 (a principal and a secondary one) were very close to each other. Finally, three bands were detected for the *S. cerevisiae* sequence (Scer21): two fast migrating bands (between dx9 and dx12) and a slow migrating band (between dx12 and dT21).

The migration pattern obtained in KCl for Gia18, Tet22, Scer21, Par21 and Oxy28 provides evidence of the presence of at least two (or three, for Scer21) different structures at the tested oligonucleotide strand concentration (30 μM). For Gia18, Tet22 and Scer21, the slow migrating band detected in KCl may be ascribed to the presence of multi-stranded species; this hypothesis is supported by changes in their CD spectra upon increasing oligonucleotide strand concentration from 3 to 30 μM (Figure 3).

**DISCUSSION**

In the present work we studied telomeric sequences from different organisms, with the aim of probing their propensity to fold into G-quadruplex structures, providing some information about their folding and studying their stability.

**G-quadruplex folding**

Thermal difference spectra provided evidence of the formation of G-quadruplex structures, both in sodium and in potassium, for all the studied telomeric sequences, with the exception of the *C. guilliermondii* (Cgi26) and *S. pombe* (Spom20 and Spom23) sequences. By contrast, TDS and CD in sodium and in potassium did not support formation of G-quadruplexes by Cgi26. Spom20 in potassium displayed unambiguous TDS and CD G-quadruplex signatures. Finally, the very small negative peak at 295 nm in TDS of Spom20 in sodium and Spom23 in both sodium and potassium suggests that a very minor fraction of these sequences folds into G-quadruplexes; this fraction may contribute to the observed CD spectra.

**Molecularity**

For the telomeric sequences displaying a TDS G-quadruplex signature, three lines of evidence supported folding into intramolecular structures in sodium: the independence of (i) melting temperatures and (ii) CD spectra from oligonucleotide strand concentration; (iii) the presence of a single and fast migrating band (as compared to dT21) in a non-denaturing PAGE experiment. The same conclusion could be drawn for most of the sequences in potassium (Bom17, Asc20, Hum21, Ara24, Tom24, Glab26, Chla27 and Spom20).

For the *Paramecium* (Par21) and *Oxytricha* (Oxy28) sequences in potassium, two bands were detected by PAGE; the mobility of these bands (as compared to the internal markers) and the independence of Par21 *T*<sub>m</sub> and Par21–Oxy28 CD spectra from oligonucleotide strand concentration suggest folding into intramolecular structures of different electrophoretic mobilities (Oxy28 *T*<sub>m</sub> in potassium were too high to be determined).

The PAGE migration pattern of *Giardia* (Gia18), *Tetrahymena* (Tet22) and *S. cerevisiae* (Scer21) sequences in potassium provided evidence of the presence of dramatically different structures (at least two for Gia18 and Tet22 and at least three for Scer21). For these three sequences, the slow migrating band may be ascribed to multi-stranded structures. This hypothesis is supported by the change in Gia18, Tet22 and Scer21 CD spectra upon increasing oligonucleotide strand concentration. Furthermore, the PAGE migration pattern of Gia18 in KCl was shown to be concentration-dependent: the slow migrating band was not detected at 100-fold lower oligonucleotide strand concentration (300 nM) (gel detected by Syber Gold staining, data not shown). This demonstrates that the slow migrating band of Gia18 corresponds to multi-stranded species.

On the other hand, the independence of Gia18 and Scer21 *T*<sub>m</sub> from oligonucleotide strand concentration (up to 100 μM for Gia18) does not support folding into intermolecular structures, such as bimolecular or tetramolecular G-quadruplexes (Tet22 *T*<sub>m</sub> in potassium were too high to be determined). This puzzling behaviour might be explained by folding into intramolecular G-quadruplexes (monomers) and formation of stacked monomers upon increasing oligonucleotide strand concentration from 3 to 30 μM. The appearance of a positive CD band around 260 nm might arise from stacking of terminal G-quartets with the same hydrogen-bond orientation (70,71). The hypothesis of stacked intramolecular
G-quadruplexes is not appropriate (or is only partially appropriate) for Scer21, which present a more complex non-denaturing PAGE migration pattern.

**Structural differences in sodium and in potassium**

Besides providing evidence of a concentration-dependent behaviour of Gia18, Tet22 and Scer21 in potassium, CD allowed highlighting structural differences in sodium and potassium.

The CD spectra of most of the telomeric sequences (Gia18, Hum21, Ara24, Chla27, Par21, Scer21, Tom24 and Tet22) strongly depended on the nature of the cation present in solution (sodium or potassium), suggesting different G-quadruplex conformations or a different equilibrium among multiple possible G-quadruplex conformations according to the nature of the cation. Only the Bombyx (Bom17), Ascaris (Asc20) and Oxytricha (Oxy28) sequences displayed a similar (although not strictly identical) CD signature in sodium and in potassium.

It would be unwarranted to speculate about CD signatures and G-quadruplex folding topologies, for several reasons. (i) The simple empirical rule often used to interpret CD spectra (Type I spectrum = parallel folding, Type II spectrum = antiparallel folding) is based on a few resolved structures and is not of general validity (71). (ii) Interpretation of CD spectra in terms of relative strand orientation is not formally correct (70,71). (iii) Interpretation of CD spectra in terms of folding topologies relies on the implicit assumption that a sequence folds into a single G-quadruplex structure. A single sequence may fold into multiple G-quadruplex structures in the same solution; this is the case, for example, of human and Giardia telomeric sequences in potassium (Figure 1). Structure determination requires high-resolution structural studies, such as nuclear magnetic resonance (NMR) or X-ray crystallography, and is beyond the scope of this work.

**Stability of G3 and G4 motifs**

All sequences bearing three or more consecutive guanines folded into G-quadruplexes more stable in potassium than in sodium (higher Tm), as expected for G-quadruplexes. Sequences bearing three consecutive guanines (including Gla26, for a total of eight motifs) had Tm ranging from 46 to 60°C in sodium and from 57.5 to 68°C in potassium. Notably, for the G3T4A motif sequences, Tm slightly decreased with n for n = 1, 2 and 3 (Gia18, Hum21 and Ara24, respectively); increasing n to 4 (Chla27) led to a dramatic decrease in Tm (about 10°C). The Tetrahymena (Tet22) and Oxytricha (Oxy28) sequences, bearing four consecutive guanines, were the most stable; they had roughly the same Tm in sodium (64 and 66°C, respectively) and Tm higher than 80°C in potassium.

We carried out a thermodynamic analysis of melting transitions (Table 1). A two-state equilibrium model, assuming linear low- and high-temperature absorbance baselines, resulted in linear van’t Hoff graphs. Non-denaturing PAGE experiments revealed that such a model is not appropriate for Gia18, Par21 and Scer21 in KCl, nevertheless linear van’t Hoff graphs could be obtained even for these sequences in KCl. Thermodynamic values did not depend on oligonucleotide strand concentration (3 and 30 μM). For G3 and G3/G4 motifs, ΔHm of folding varied between −60 and −48 kcal mol−1 in NaCl and between −77 and −60 kcal mol−1 in KCl; ΔGo of folding, extrapolated at 37°C, varied between −3.7 and −1.6 kcal mol−1 in NaCl and between −6.2 and −4.2 kcal mol−1 in KCl. The analysis of G4 motifs was possible only in NaCl: standard enthalpy change of Oxy28 folding (ΔH = −82 ± 13 kcal mol−1) was more negative than standard enthalpy changes of Tet22 folding (ΔH = −54 ± 8 kcal mol−1) and of G3 and G3/G4 motifs. For Tet22, G3 and G3/G4 motifs in NaCl, Tm and ΔGo were linearly correlated (linear regression coefficient R = 0.9877). In KCl, Tm and ΔGo correlation was poorer; this might be ascribed to a greater error in ΔGo determination in KCl than in NaCl (due to a greater uncertainty in high-temperature baseline determination and to extrapolation of ΔGo at a temperature far below the temperature ranges where melting transitions occurred).

The influence of G-tract length, loop length and loop sequence on G-quadruplex structure and stability has been explored on model sequences (62–64,72–77); nevertheless, no general rules has emerged (78). In particular, the number of consecutive guanines does not necessarily correspond to the number of G-quartets (50,57,60). NMR studies of the Oxytricha 3(G4T4)4-G4 and of the Tetrahymena 3(T3G4)4 telomeric sequences in sodium revealed four stacked G-quartets in the Oxytricha G-quadruplex (48,49) (Figure 1H) and only three stacked G-quartets in the Tetrahymena G-quadruplex, connected by a GGG lateral loop, a central TTG lateral loop and a TT double-chain-reversal loop (50) (Figure 1F). Our data do not allow drawing conclusions about the number of stacked G-quartets in the Tetrahymena telomeric sequence Tet22, which differed from the one used in the NMR study by two nucleotides (TT). Nevertheless, in sodium, the amplitude in absorbance of the 295 nm melting transition of Tet22 (0.047 variation in absorbance recorded at 3 μM oligonucleotide strand concentration and 1 cm optical path length) was very similar to the amplitudes of the sequences bearing three consecutive guanines (0.049 ± 0.008 on average) and significantly smaller than the amplitude of Oxy28 (0.072) (Figure S2). This feature, along with ΔHm values discussed above, might argue stacking of three G-quartets in Tet22 and of four G-quartets in Oxy28.

For a comprehensive analysis of tomato plant and Paramecium degenerated telomeric sequences, we studied the sixteen following variant sequences: 5(GGG TTWA)3GGG3 (where W is A or T) (tomato) and 5(GGGKTT3)GGG3 (where K is T or G) (Paramecium) (Table 2). Not surprisingly, all these sequences folded into stable G-quadruplexes (TDS are shown in Supplementary Figures S4 and S5). Tm did not strongly vary among tomato variant sequences: it ranged from 55 to 58°C in NaCl and from 64 to 69°C in KCl (Table 2). Tm of Paramecium variant sequences spanned a broader range: from 52 to 60°C in NaCl and from 68 to 76°C in KCl (Table 2). Two correlations between sequence and stability
and between sequence and CD signature emerged. (i) Among *Paramecium* variant sequences, stability in NaCl correlated with the sequence of the third repeated unit (more precisely of nucleotides 16–18): G-quadruplexes formed by \( (G_3KTT)_2G_3TTT \) sequences were less stable (\( T_{m,NaCl} \) of 52–54°C) than the ones formed by \( (G_3KTT)_2G_3GTGTG_3 \) sequences (\( T_{m,NaCl} \) of 58–60°C). (ii) Among tomato plant variant sequences, the sequence of the first loop connecting runs of guanines (nucleotides 4–7) correlated with CD spectra in KCl: \( G_3TTA(A_3TWA)_2G_3 \) sequences had Tom24-like CD spectra, while \( G_3TTT(A_3TWA)_2G_3 \) sequences had Ara24-like CD spectra (Figure S4).

### Stability of G2 motifs

Among the telomeric sequences bearing two consecutive guanines, only the *Bombbyx* (Bom17) and the *Ascaris* (Asc20) sequences folded into G-quadruplexes both in sodium and in potassium. Permuting a cytosine from the first position (Asc20) to the last position (Spom20) of the G-tract connecting sequences abolished G-quadruplex folding in sodium. The analysis of Bom17 and Asc20 melting transitions at 295 nm revealed higher \( T_{m,\text{NaCl}} \) in potassium than in sodium (about 2°C). This behaviour is unique to Bom17 and Asc20 sequences, since the model sequences \( (G_3TWA)_2G_3 \) with \( n = 3 \) and \( n = 4 \) folded into G-quadruplexes with higher \( T_{m,\text{NaCl}} \) in potassium than in sodium (data not shown). An apparent higher \( T_{m,\text{NaCl}} \) at 295 nm in sodium than in potassium can result from the equilibrium between a G-quadruplex and a non-G-quadruplex structure. Asc20 and Bom17 can in principle fold into (hairpin-) duplexes with, respectively, 2 out of 3 and 1 out of 3 nucleotides engaged in Watson–Crick base pairs. The presence of a competing structure is suggested by results obtained in LiCl and results obtained with fluorescent conjugates. Folding of Asc20 and Bom17 into non-G-quadruplex structures, melting at about 38 and 25°C, respectively, was indeed observed in LiCl (Figure S6); folding into these structures did not result into absorbance variation at 295 nm. Results obtained with the double-dye-labelled sequences suggest than both Asc20 and Bom17 fold into two structures (a G-quadruplex and a non-G-quadruplex structure) and that the presence of the dyes completely shift the equilibrium of the *Ascaris* sequence toward the non-G-quadruplex structure and the equilibrium of the *Bombbyx* sequence toward the G-quadruplex (a ‘classical’ G-quadruplex, i.e. more stable in potassium than in sodium). If a competing structure is present, then the \( T_{m,\text{NaCl}} \) at 295 nm is not a ‘true’ G-quadruplex \( T_{m,\text{NaCl}} \), even if the competing structure does not contribute to melting profiles at 295 nm. More precisely, if \( f \) is the G-quadruplex folded fraction at low temperatures, then the graphically determined \( T_{m,\text{NaCl}} \) at 295 nm is the temperature at which a fraction \( f/2 \) is folded into a G-quadruplex; this temperature, which depend on \( f \) (i.e. on the equilibrium between the G-quadruplex and the competing structure) cannot be used as a parameter for G-quadruplex stability. The presence of a non-G-quadruplex competing structure may also explain the small amplitude of Asc20 melting transition at 295 nm in sodium compared to the one in potassium (Figure 4A). This feature may result from a smaller G-quadruplex folded fraction in sodium than in potassium at low temperatures. This would mean that G-quadruplexes formed by Asc20 are less stable in sodium than in potassium (under the reasonable hypothesis that the stability of the competitive structure does not depend on the nature of the cation). This hypothesis is not in contradiction with higher \( T_{m,\text{NaCl}} \) at 295 nm in sodium than in potassium, as discussed above. Finally, the independence of Asc20 and Bom17 melting temperatures from oligonucleotide strand concentration (3, 10 and 30 μM) supports the intramolecular nature of both the G-quadruplex and the non-G-quadruplex structures. The PAGE migration patterns of Asc20 and Bom17 are not inconsistent with the presence of G-quadruplexes and hairpin-duplexes, when compared to the internal migration standard dx9 and dx12.

The inability of the Cgi26 and Spom23 sequences to fold into G-quadruplexes or stable G-quadruplexes does not result just from the length of the loops separating the runs of guanines (6 and 5 nt, respectively). Actually the model sequence \( (G_3TWA)_3G_3 \) folded into G-quadruplexes both in sodium and in potassium (with a \( T_{m} \) of 16 and 25°C, respectively), whereas the model sequence \( (G_3TWA)_3G_3 \) folded into stable G-quadruplexes in potassium only (with a \( T_{m} \) of 18°C) (data not shown). Hence, the behaviour of the telomeric sequences bearing just two consecutive guanines (from Bom17 to Cgi26) arises from their peculiar loop sequences.

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**Table 2.** Tomato plant and *Paramecium* variant telomeric sequences and \( T_{m,\text{NaCl}} \) determined by analysis of melting curves recorded at 295 nm, at 3 μM oligonucleotide strand concentration

| Sequence name | Sequence (from 5’ to 3’) | \( T_{m,\text{NaCl}} \) (°C) in NaCl | \( T_{m,\text{KCl}} \) (°C) in NaCl |
|---------------|---------------------------|-----------------------------|-----------------------------|
| Tomato plant  |                           |                             |                             |
| Tom24         | G_{3TTAAG},TTAAG,TTAAG    | 55                          | 67                          |
| TAA           | G_{3TTAAG},TTAAG,TTAAG    | 58                          | 65                          |
| ATA           | G_{3TTAAG},TTAAG,TTAAG    | 57                          | 69                          |
| AAT           | G_{3TTAAG},TTAAG,TTAAG    | 58                          | 67                          |
| TTA           | G_{3TTAAG},TTAAG,TTAAG    | 58                          | 66                          |
| TAT           | G_{3TTAAG},TTAAG,TTAAG    | 57                          | 64                          |
| ATT           | G_{3TTAAG},TTAAG,TTAAG    | 56                          | 67                          |
| Ara24         | G_{3TTAAG},TTAAG,TTAAG    | 58                          | 66                          |
| *Paramecium*  |                           |                             |                             |
| Par21         | G_{3TTG},TTTG,TTTG        | 52                          | 68                          |
| GTT           | G_{3TTG},TTTG,TTTG        | 54                          | 72                          |
| TGT           | G_{3TTG},TTTG,TTTG        | 52                          | 68                          |
| TTG           | G_{3TTG},TTTG,TTTG        | 59                          | 72                          |
| GGT           | G_{3TTG},TTTG,TTTG        | 54                          | 71                          |
| GTG           | G_{3TTG},TTTG,TTTG        | 58                          | 74                          |
| TGG           | G_{3TTG},TTTG,TTTG        | 58                          | 73                          |
| GGG           | G_{3TTG},TTTG,TTTG        | 60                          | 76                          |

\( T_{m,\text{NaCl}} \) reported in this table were obtained on a different spectrophotometer from those reported in Table 1. Temperature was measured with a sensor in a sealed cuvet provided by the manufacturer; this may explain the slightly higher \( T_{m,\text{NaCl}} \) reported in this table as compared to \( T_{m,\text{NaCl}} \) reported in Table 1 for Tom24, Ara24 and Par21. TDS and CD spectra are reported in Supplementary Figures S4 and S5.
In summary, with regard to the ability of folding into intramolecular G-quadruplexes, our study highlights at least two different sets of telomeric motifs: (i) yeast telomeric motifs and (ii) telomeric motifs from other eukaryotes. On the one hand, the ability to fold into G-quadruplex structures does not appear to be a conserved feature among the exceptionally diverse yeast telomeric sequences: unlike C. glabrata, Giuillermondii telomeric sequence did not fold into G-quadruplexes; unlike repetitions of the TG3TG motif (frequently occurring in S. cerevisiae telomeres), repetitions of G2TTAC(A) motifs (frequently occurring in S. pombe telomeres) were not prone to fold into stable G-quadruplexes. On the other hand, evolution of telomeric sequences in eukaryotes other than yeasts (repetitions of G1T1A4A in a variety of organisms, G2T2A4 in ciliates, G2T2A in many insects and G2CT2A in nematodes) appears to have preserved the G-quadruplex folding potential, even if these structures pose a potential problem to telomeric G-strand replication (79,80). Notably, telomeric sequences with three or more consecutive guanines folded into G-quadruplexes with melting temperatures above temperatures where organisms known to have these telomeric sequences thrive. Curiously, no telomeric motif with an adenine following the run of guanines has been found, with the exception of the putative G3AAT telomeric motif of the hot-spring unicellular red alga Cyanidioschyzon merolae (81). Intriguingly, an adenine as the first base in the loops connecting runs of two or three consecutive guanines is detrimental to G-quadruplex folding (47,77). Cyanidioschyzon merolae is an extremophilic organism, thriving in acid hot springs (pH 1.5, 45°C). Its telomeric sequence \(5'G3AATG3'\) folded into very stable G-quadruplexes, melting at 70°C in NaCl and stable at temperatures as high as 90°C in KCl (Supplementary Figure S7). Unlikely, telomeric motifs of other thermophilic eukaryotes (such as fungi able to grow at 55–60°C, hot-spring algae other than C. merolae, the worms Paralvinella suffincola and Alvinella Pompejana, tardigrades) are not known, to our knowledge.

In conclusion, the present study provides an insight into intramolecular G-quadruplex folding potential of telomeric sequences from a variety of organisms. (i) All the telomeric sequences bearing three or more consecutive guanines (in particular G1T1A4A motifs and G2T2A4 ciliate motifs) folded into stable G-quadruplexes both in potassium and in sodium. (ii) The G2TTA telomeric motif sequence (found in many insects, such as Bombyx mori) and the G2CTTA telomeric motif sequence (found in the nematodes Ascaris lumbricoides and Caenorhabditis elegans) folded into G-quadruplexes both in potassium and in sodium. Our results strongly suggest that, for these two sequences, the G-quadruplex structures are in equilibrium with non-G-quadruplex structures (likely hairpin-duplexes). (iii) G-quadruplex folding does not appear to be a characteristic feature of budding yeast telomeres, despite a widespread conserved TG-rich motif of about six base pairs (28). Actually, unlike repetitions of the S. cerevisiae frequent occurring motif TG3TG and of the Candida glabrata CTGTG4TCTG3TG motif, the Candida guillermondii telomeric sequence (TACTG2TG motif) was unable to fold into G-quadruplexes. (iv) The most frequent telomeric motifs in the fission yeast Schizosaccharomyces pombe are G2TTAC and G2TTAC. Only repetitions of the GGTTAC motif folded into G-quadruplexes and just in potassium. In order to verify if the S. pombe expanded consensus telomeric motif gggggGTTTACe (where lower case letters denote rare insertions) (34,35) is prone to fold into G-quadruplex structures a deeper investigation of combinations of possible S. pombe telomeric units is required. (v) For most of the telomeric sequences folding into G-quadruplexes, structural differences between the sodium and the potassium forms were revealed by CD.

(vi) Finally, circular dichroism and PAGE supported folding of the Giardia intestinalis GGGTA, of the Tetrahymena GGGGTT and of the S.cerevisiae TGGGTG motif sequences into intramolecular and multi-stranded structures in potassium; the independence of their \(T_m\) from oligonucleotide strand concentration suggests that these multi-stranded structures may be stacked intramolecular G-quadruplexes.

Besides allowing comparison of G-quadruplex folding and stability of eukaryotic telomeric sequences, the present study enriches the dataset necessary to build and refine algorithms predicting G-quadruplex stability (82).

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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**REFERENCES**

1. Wellinger, R.J. and Sen, D. (1997) The DNA structures at the ends of eukaryotic chromosomes. *Eur. J. Cancer.*, **33**, 735–749.
2. Moyzis, R.K., Buckingham, J.M., Cram, L.S., Dani, M., Deaven, L.L., Jones, M.D., Meyne, J., Ratliff, R.L. and Wu, J.R. (1988) A highly conserved repetitive DNA sequence, \(5'TTAGGG3'\), present at the telomeres of human chromosomes. *Proc. Natl Acad. Sci. USA*, **85**, 6622–6626.
1. Meyne, J., Ratliff, R.L., and Moyzis, R.K. (1989) Conservation of the human telomere sequence (TTAGGG)n among vertebrates. Proc. Natl Acad. Sci. USA, 86, 7049–7053.
2. Blackburn, E.H. and Challoner, P.B. (1984) Identification of a telomeric DNA sequence in Trypanosoma brucei. Cell, 36, 447–457.
3. Van der Ploeg, L.H., Liu, A.Y. and Borst, P. (1984) Structure of the growing telomeres of Trypanosomes. Cell, 36, 459–468.
4. Schechtman, M.G. (1990) Characterization of telomere DNA from Neurospora crassa. Gene, 88, 159–165.
5. Emery, H.S. and Weiner, A.M. (1981) An irregular satellite sequence is found at the termini of the linear extrachromosomal rDNA in Dictyostelium discoideum. Cell, 26, 411–419.
6. Forney, J., Henderson, E.R. and Blackburn, E.H. (1987) Identification of the telomeric sequence of the acellular slime molds Daidymium trids and Physarum polycephalum. Nucleic Acids Res., 15, 9143–9152.
7. Sinclair, C.S., Richmond, R.H. and Ostrand, G.K. (2007) Characterization of the telomere regions of selarctinian coral, Acropora succulenta. Genetica, 129, 227–233.
8. Le Blanc, S.M., Kasc, R.S. and Van der Ploeg, L.H. (1991) Analysis of the Didymium lamblia rRNA encoding telomere with [TAGGG]n as the telomere repeat. Nucleic Acids Res., 19, 5790.
9. Fuchs, J., Brandes, A. and Schubert, I. (1995) Telomere sequence localisation and karyotype evolution in higher plants. Plant Syst. Evol., 196, 227–241.
10. Richards, E.J. and Ausubel, F.M. (1988) Isolation of a higher eukaryotic telomere from Arabidopsis thaliana. Cell, 53, 127–136.
11. Petracek, M.E., Lelebvre, P.A., Silbow, C.D. and Berman, J. (1990) Chlamydomonas telomere sequences are A + T-rich but contain three consecutive G-C base pairs. Proc. Natl Acad. Sci. USA, 87, 8222–8226.
12. Okazaki, S., Tsuchida, K., Ishikawa, H. and Fujiwara, H. (1993) Identification of a pentanucleotide telomeric sequence at the terminus of the extrachromosomal rDNA sequence of Didymium iridis molds. J. Mol. Biol., 235, 1133–1139.
13. Wicky, C., Villeneuve, A.M., Lauper, N., Codourey, L., Tobler, H. and Muller, F. (1996) Telomeric repeats (TTAGGG)n are sufficient for chromosome capping function in Caenorhabditis elegans. Proc. Natl Acad. Sci. USA, 93, 8983–8988.
14. Oka, Y. and Thomas, C.A. Jr (1987) The cohering telomeres of Neocallimastix tetragona. Nucleic Acids Res., 15, 649–654.
15. Boretz, C., Berger, L., Postberg, J., Hanes, I., Lips, H.J. and Pluckthun, A. (2001) In vitro generated antibodies specific for telomere-guanine-quadruplex DNA react with Styloynchia lemnae macronucleus. Proc. Natl Acad. Sci. USA, 98, 8572–8577.
16. Paeschke, K., Simonsson, T., Postberg, J., Rhodes, D. and Lips, H.J. (2005) Telomere end-binding proteins control the formation of G-quadruplex DNA structures in vivo. Nat. Struct. Mol. Biol., 12, 847–854.
17. Baroin, A., Prat, A. and Caron, F. (1987) Telomere site position heterogeneity in macronuclear DNA of Paramecium primaurelia. Nucleic Acids Res., 15, 1717–1728.
18. Forney, J.D. and Blackburn, E.H. (1988) Developmentally controlled telomere addition in wild-type and mutant paramecia. Mol. Cell. Biol., 8, 251–258.
19. McCormick-Graham, M. and Romero, D.P. (1996) A single telomerase RNA is sufficient for the synthesis of variable telomeric DNA repeats in ciliates of the genus Paramecium. Mol. Cell. Biol., 16, 1871–1879.
20. Teixeira, M.T. and Gilson, E. (2005) Telomere maintenance, function and evolution: the yeast paradigm. Chromosome Res., 13, 535–548.
21. Lue, N.F. Plasticity of telomere maintenance mechanisms in yeast. Trends Biochem. Sci., 35, 8–17.
22. McEachern, M.J. and Blackburn, E.H. (1994) A conserved sequence motif within the exceptionally diverse telomeric sequences of budding yeasts. Proc. Natl Acad. Sci. USA, 91, 3453–3457.
23. Cohn, M., McEachern, M.J. and Blackburn, E.H. (1998) Telomeric sequence diversity within the genus Saccharomyces. Curr. Genet., 33, 83–91.
24. Wang, S.S. and Zakian, V.A. (1990) Sequencing of Saccharomyces telomeres cloned using T4 DNA polymerase reveals two domains. Mol. Cell. Biol., 10, 4415–4419.
25. Singer, M.S. and Gottschling, D.E. (1994) TLC1: template RNA component of Saccharomyces cerevisiae telomerase. Science, 266, 404–409.
26. Leonard, J., Box, J.A., Bunch, J.T. and Baumann, P. (2008) TER1, the RNA subunit of fission yeast telomerase. Nat. Struct. Mol. Biol., 15, 26–33.
27. Webb, C.J. and Zakian, V.A. (2008) Identification and characterization of the Schizosaccharomyes pombe TER1 telomerase RNA. Nat. Struct. Mol. Biol., 15, 34–42.
28. Hiraoka, Y., Henderson, E. and Blackburn, E.H. (1998) Not so peculiar: fission yeast telomere repeats. Trends Biochem. Sci., 23, 126–131.
29. Trujillo, K.M., Bunch, J.T. and Baumann, P. (2005) Extended DNA binding site in Pot1 broadens sequence specificity to allow recognition of heterogeneous fission yeast telomeres. J. Biol. Chem., 280, 9119–9128.
30. Neidle, S. and Balasubramanian, S. (2006) Quadruplex Nucleic Acids. RSC Biomolecular Sciences, Cambridge.
31. Williamson, J.R., Rehurman, M.K. and Cech, T.R. (1989) Monovalent cation-induced structure of telomeric DNA: the G-quartet model. Cell, 59, 871–880.
32. Neidle, S. and Balasubramanian, S. (2006) Quadruplex Nucleic Acids. RSC Biomolecular Sciences, Cambridge.
33. Wang, S.S. and Zakian, V.A. (1990) Sequencing of Saccharomyces telomeres cloned using T4 DNA polymerase reveals two domains. Mol. Cell. Biol., 10, 4415–4419.
34. Singer, M.S. and Gottschling, D.E. (1994) TLC1: template RNA component of Saccharomyces cerevisiae telomerase. Science, 266, 404–409.
35. Leonard, J., Box, J.A., Bunch, J.T. and Baumann, P. (2008) TER1, the RNA subunit of fission yeast telomerase. Nat. Struct. Mol. Biol., 15, 26–33.
36. Webb, C.J. and Zakian, V.A. (2008) Identification and characterization of the Schizosaccharomyces pombe TER1 telomerase RNA. Nat. Struct. Mol. Biol., 15, 34–42.
37. Hiraoka, Y., Henderson, E. and Blackburn, E.H. (1998) Not so peculiar: fission yeast telomere repeats. Trends Biochem. Sci., 23, 126–131.
38. Trujillo, K.M., Bunch, J.T. and Baumann, P. (2005) Extended DNA binding site in Pot1 broadens sequence specificity to allow recognition of heterogeneous fission yeast telomeres. J. Biol. Chem., 280, 9119–9128.
39. Neidle, S. and Balasubramanian, S. (2006) Quadruplex Nucleic Acids. RSC Biomolecular Sciences, Cambridge.
40. Williamson, J.R., Rehurman, M.K. and Cech, T.R. (1989) Monovalent cation-induced structure of telomeric DNA: the G-quartet model. Cell, 59, 871–880.
41. Wang, S.S. and Zakian, V.A. (1990) Sequencing of Saccharomyces telomeres cloned using T4 DNA polymerase reveals two domains. Mol. Cell. Biol., 10, 4415–4419.
42. Singer, M.S. and Gottschling, D.E. (1994) TLC1: template RNA component of Saccharomyces cerevisiae telomerase. Science, 266, 404–409.
60. Hu, L., Lim, K.W., Bouaziz, S. and Phan, A.T. (2009) Giardia telomere repeat d(TAGGG)4 G-quartet. *Structure*, 17, 1141–1156.

61. Wang, Y. and Patel, D.J. (1993) Solution structure of the human telomere repeat d(TAGGG)4 G-quartet. *Structure*, 1, 263–282.

53. Dai, J., Carver, M., Punchihewa, C., Jones, R.A. and Yang, D. (2007) Intramolecular G-quadruplexes in K+ solution: insights into structure polymorphism of the human telomeric sequence. *Nucleic Acids Res.*, 35, 927–940.

54. Lim, K.W., Amrane, S., Bouaziz, S., Xu, W., Mu, Y., Patel, D.J., Lui, K.N. and Phan, A.T. (2009) Structure of the human telomere in K+ solution: a stable basket-type G-quadruplex with only two G-tetrad layers. *J. Am. Chem. Soc.*, 131, 4301–4309.

55. Amrane, S., Ang, R.W., Tan, Z.M., Li, C., Lim, J.K., Lim, J.M., Lim, K.W. and Phan, A.T. (2009) A novel chair-type G-quadruplex formed by a Bombyx mori telomeric sequence. *Nucleic Acids Res.*, 37, 931–938.

56. Hu, L., Lim, K.W., Bouaziz, S. and Phan, A.T. (2009) Giardia telomeric sequence d(TAGGGH4 forms two intramolecular G-quadruplexes in K+ solution: effect of loop length and sequence on the folding topology. *J. Am. Chem. Soc.*, 131, 16824–16831.

57. Venczel, E.A. and Sen, D. (1993) Parallel and antiparallel G-DNA structures from a complex telomeric sequence. *Biochemistry*, 32, 6220–6228.

58. Hazel, P., Huppert, J., Balasubramanian, S. and Neidle, S. (2004) Loop-length-dependent folding of G-quadruplexes. *J. Am. Chem. Soc.*, 126, 16405–16415.

59. Guедин, A., De Cian, A., Gros, J., Lacroix, L. and Mergny, J.L. (2008) Sequence effects in single-base loops for quadruplexes. *Biochimie*, 90, 686–696.

60. Guедин, A., Gros, J., Alberti, P. and Mergny, J.L. (2010) How long is too long? Effects of loop size on G-quadruplex stability. *Nucleic Acids Res.*, 38, 7858–7868.

61. De Cian, A., Grellier, P., Mouray, E., Depoix, D., Bertrand, H., Monchaud, D., Teulade-Fichou, M.P., Mergny, J.L. and Alberti, P. (2008) Plasmodium telomeric sequences: structure, stability and quadruplex targeting by small compounds. *ChemBioChem*, 9, 2730–2739.

62. Mergny, J.L. and Lacroix, L. (2009) UV melting of G-quadruplexes. *Curr. Protoc. Nucleic Acids Chem.*, 17, 1.1–17.1.15.

63. Mergny, J.L., Li, J., Lacroix, L., Amrane, S. and Chaires, J.B. (2005) Thermal difference spectra: a specific signature for nucleic acid structures. *Nucleic Acids Res.*, 33, e138.

64. Kajiyama, K., Kyo, M. and Vorlickova, M. (2007) Oligo(dT) is not a correct native PAGE marker for single-stranded DNA. *Biochem. Biophys. Res. Commun.*, 353, 776–779.

65. Mergny, J.L., Phan, A.T. and Lacroix, L. (1998) Following G-quartet formation by UV-spectroscopy. *FEBS Lett.*, 435, 74–78.

66. Gray, D.M., Sen, D., Yamada, K., Repges, R., Repges, C., Raabe, G. and Fleischhauer, J. (2008) Measured and calculated CD spectra of G-quartets stacked with the same or opposite polarities. *Chirality*, 20, 431–440.

67. Masierno, S., Trotta, R., Pieraccini, S., De Tito, S., Perone, R., Randazzo, A. and Spada, G.P. (2010) A non-empirical chromophoric interpretation of CD spectra of DNA G-quadruplex structures. *Org. Biomol. Chem.*, 8, 2683–2692.

68. Risitano, A. and Fox, K.R. (2003) Stability of intramolecular DNA quadruplexes: comparison with DNA duplexes. *Biochemistry*, 42, 6507–6513.

69. Rachwal, P.A., Findlow, I.S., Werner, J.M., Brown, T. and Fox, K.R. (2007) Intramolecular DNA quadruplexes with different arrangements of short and long loops. *Nucleic Acids Res.*, 35, 4214–4222.

70. Rachwal, P.A., Brown, T. and Fox, K.R. (2007) Effect of G-tract length on the topology and stability of intramolecular DNA quadruplexes. *Biochemistry*, 46, 3036–3044.

71. Bugaut, A. and Balasubramanian, S. (2008) A sequence-independent study of the influence of short loop lengths on the stability and topology of intramolecular DNA G-quadruplexes. *Biochemistry*, 47, 689–697.

72. Smargiasso, N., Rosu, F., Hsia, C., Colson, P., Baker, E.S., Bowers, M.T., De Pauw, E. and Gabelica, V. (2008) G-quadruplex DNA assemblies: loop length, cation identity, and multimer formation. *J. Am. Chem. Soc.*, 130, 10208–10216.

73. Guедин, A., Alberti, P. and Mergny, J.L. (2009) Stability of intramolecular quadruplexes: sequence effects in the central loop. *Nucleic Acids Res.*, 37, 5559–5567.

74. Kumar, N. and Maiti, S. (2008) A thermodynamic overview of naturally occurring intramolecular DNA quadruplexes. *Nucleic Acids Res.*, 36, 5610–5622.

75. Crabbe, L., Verdun, R.E., Haggblom, C.I. and Karlseder, J. (2004) Defective telomere lagging strand synthesis in cells lacking WRN helicase activity. *Science*, 306, 1951–1953.

76. Sfeir, A., Kosiyatrakul, S.T., Hockemeyer, D., MacRae, S.L., Bowers, M.T., De Pauw, E. and Gabelica, V. (2008) G-quadruplex DNA assemblies: loop length, cation identity, and multimer formation. *J. Am. Chem. Soc.*, 130, 10208–10216.

77. Crumble, B., Verduin, R.E., Haggblom, C.I. and Karlseder, J. (2004) Defective telomere lagging strand synthesis in cells lacking WRN helicase activity. *Science*, 306, 1951–1953.

78. Fein, A., Kosiyatrakul, S.T., Hockemeyer, D., MacRae, S.L., Barshes, J., Schildkraut, C.L. and de Lange, T. (2009) Mammalian telomeres resemble fragile sites and require TRF1 for efficient replication. *Cell*, 138, 99–103.

79. Nozaki, H., Takano, H., Misumi, O., Terasawa, K., Matsuzaki, M., Maruyama, S., Nishida, K., Yasigawa, F., Yoshida, Y., Fujitawa, T. et al. (2007) A 100%–complete sequence reveals unusually simple genomic features in the hot-spring red alga *Cyanidioschyzon merolae*. *BMC Biol.*, 5, 28.

80. Siegel, O., Payet, L., Mergny, J.L., MacKay, D.J. and Huppert, J.L. (2009) Predicting and understanding the stability of G-quadruplexes. *Bioinformatics*, 25, 1374–1382.

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