Photoreactivity of the linker region of two consecutive G-quadruplexes formed by human telomeric DNA†

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We report the application of a photoreaction method for probing two consecutive G-quadruplexes formed by human telomeric DNA. This method can discriminate the loop structure located between two consecutive G-quadruplexes formed by eight TTAGGG repeats in K⁺ and Na⁺ solutions.

The human telomeric DNA contains thousands of tandem TTAGGG repeats and an approximately 200 bp 3′ single-stranded overhang that is capable of forming G-quadruplex structures.1 Recent visualization studies provided evidence of G-quadruplex formation at the telomeric region in cells.2 Stable G-quadruplexes formed by telomere overhang can protect chromosomes from degradation, end fusion, and being recognized as double-strand breaks (DSBs).3,4 To date, various intramolecular G-quadruplex structures formed by four TTAGGG repeats in the presence of K⁺ or Na⁺ ions have been elucidated.5–9 However, this view of single G-quadruplex formation is incomplete, as higher-order G-quadruplex structures are more biologically relevant considering the existence of the long telomere repeats. The consecutive G-quadruplexes formed by longer telomeric DNA have been investigated in several experimental studies, including those that used CD spectra, NMR, atomic force microscopy (AFM), and optical tweezers.10–14 Moreover, the study of higher-order G-quadruplex structures offers new and selective binding sites for drugs that target telomeric sequences. Several ligands that can selectively bind to higher-order G-quadruplexes have been reported.15,16

Because the single G-quadruplex units of the telomeric sequence in K⁺ and Na⁺ solutions are highly polymorphic, various higher-order G-quadruplex structures may be formed by long telomeric DNA.17 The sequence of eight TTAGGG repeats and its variant sequences have been frequently used in the study of higher-order structures. It was suggested that these sequences could form two consecutive G-quadruplexes that were connected by a TTA loop.18,19 Stable consecutive G-quadruplex structures can be formed by two hybrid G-quadruplexes (hybrid-1 or hybrid-2) in K⁺ solution, or by two antiparallel G-quadruplexes in Na⁺ solution. Molecular modelling studies indicate that the stacking model of hybrid-1 and hybrid-2 was most stable in K⁺ solution.20 As single G-quadruplex formation is quite different in K⁺ and Na⁺ solutions, consecutive G-quadruplex structures formed by two G-quadruplex subunits have different stacking interactions and TTA loop structures in the linker region.

Conversely, the thymine (T) base of DNA can be replaced with 5-halouracil (BrU or I). We have demonstrated that the photoreactions of 5-halouracil-containing DNA can be used to probe various conformations of DNA, including the A, B, and Z forms and G-quadruplexes.21–25 Under 302 nm irradiation, the photochemical products formed by hydrogen abstraction are highly dependent on conformation. Regarding G-quadruplexes, the photoreaction efficiency and products are closely related to the loop structure. In our previous study, we found that the 2′-deoxyribonolactone residue was effectively and selectively produced in the diagonal loop of the antiparallel G-quadruplexes formed by 5-iodouracil-containing telomeric DNA. However, there was almost no photoreactivity in the loop structures of hybrid G-quadruplexes formed in K⁺ solution.26

In this study, we applied our photoreaction method to detect the loop structure located between two consecutive G-quadruplexes formed by eight TTAGGG repeats in K⁺ and Na⁺ solutions. The second thymine of the TTA loop was replaced with BrU, as it was most reactive in the diagonal loop of an antiparallel G-quadruplex in the case of I. We proved that, during the photoreaction process, the cleavage of C–Br occurs via a heterolysis pathway, whereas the cleavage of C–I occurs via both homolysis and heterolysis pathways.27 It was also reported that electron transfer can occur from a G-quadruplex to the T–T dimer, which is a chemical electron trap, similar to BrU.28 Therefore, we used BrU instead of I to focus on the electron transfer from G-tetrads to BrU during the photoreaction process.

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Table 1: Oligonucleotides used in this study

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>Sequence (5’ to 3’)</th>
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<tbody>
<tr>
<td>ODN 1</td>
<td>FAM-TTAGGGTAGGGBrUAGGGTTAGGG</td>
</tr>
<tr>
<td>ODN 2</td>
<td>FAM-TTAGGGTFAAGGTTAGGGTTAGGGTTAGGGBrBADAGGGTTAGGG</td>
</tr>
<tr>
<td>ODN 3</td>
<td>FAM-TTAGGGTAGGGTTAGGGTAGGGBrBADAGGGTTAGGG</td>
</tr>
<tr>
<td>ODN 4</td>
<td>FAM-TTAGGGTAGGG</td>
</tr>
<tr>
<td>ODN 5</td>
<td>FAM-TTAGGGTAGGGTTAGGGTTAGGG</td>
</tr>
<tr>
<td>ODN 6</td>
<td>FAM-TTAGGGTAGGGTTAGGGTTAGGGTTAGGG</td>
</tr>
</tbody>
</table>

All of the oligonucleotides used in this study are shown in Table 1. ODN 1, with four TTAGGG repeats, was designed to confirm the photoreaction in the diagonal loop of an antiparallel G-quadruplex reported previously. ODN 2, which may form two consecutive antiparallel G-quadruplexes, had two BrU replacements in the diagonal loops. It was designed to study the photoreaction of the diagonal loop in two consecutive G-quadruplexes formed in Na⁺ solution. ODN 3, with a BrU replacement in the TTA loop located between two G-quadruplex-forming sequences, was designed to investigate the photoreaction of the loop located between G-quadruplexes in Na⁺ and K⁺ solutions. ODN 4–6 were used as reference DNAs. All ODNs were 5’-end FAM labeled.

The CD spectra were analyzed to confirm the G-quadruplex formation. As shown in Fig. S1 (ESI†), both ODN 1 and ODN 2 formed antiparallel G-quadruplexes in Na⁺ solution and hybrid G-quadruplexes in K⁺ solution. The CD signals of ODN 2, which contained eight TTAGGG repeats, in Na⁺ and K⁺ solution had about two times the amplitude of the CD signals of ODN 1 in Na⁺ and K⁺ solutions, respectively. Therefore, the eight TTAGGG repeats labeled with FAM at the 5’ end used in the current study might mainly form two consecutive G-quadruplex structures.²⁰

Previously, we demonstrated a proposed mechanism in the hot-spot sequence (5’-(G/C)-[A]₅₋₇ BrU BrU-3’) of duplex DNA and its reverse sequence (Fig. S2, ESI†).²³,²⁴ Similar to duplex DNA, 2’-deoxyribonolactone products were obtained from the photoreaction of G-quadruplexes in our previous study.²⁶ The formation of guanine radical cations in G-quadruplexes was investigated using transient UV-vis spectra and time-dependent DFT study.²⁹,³⁰

Therefore, we assumed that a similar electron transfer mechanism occurs in the photoreaction of G-quadruplexes. Based on this mechanism, DNA samples in different ion solutions were photoirradiated separately at 0 °C for 15 min using a monochromator (302 nm) and were then subjected to heating and cleavage at 95 °C for 15 min. Subsequently, the resulting 5’-end FAM-labeled DNA fragments were analyzed by denaturing gel analysis. Finally, photoreactive sites were determined by comparison with the reference ODNs.

The results obtained for ODN 1 are shown in Fig. 1a and Fig. S3 (ESI†). In the presence of Na⁺, efficient cleavage was observed after heating, indicating the formation of deoxyribo- nolactone products in the diagonal loop of the antiparallel G-quadruplex. The product of the photoreaction also showed a similar migration in the gel vs. the reference DNA ODN 4. However, in K⁺ and Li⁺ solutions, we did not observe clear bands of deoxyribonolactone photoreaction products. These results were in accordance with the photoreaction of 5-iodouracil-containing telomeric DNA revealed in a previous HPLC analysis.²⁶ Therefore, the deoxyribonolactone products of photoreactions and photoreaction efficiency can be investigated via gel analysis using 5’-end FAM-labeled DNA. We also calculated the efficiency of photoreaction by measuring the percentage of the cleavage band. For ODN 1, the efficiency of the photoreaction using 15 min of irradiation was about 35%.

Next, we investigated the photoreaction of ODN 2 and the results were shown in Fig. 1b. ODN 4 and ODN 6 were used as reference DNAs for gel analysis. Two cleavage bands that showed similar migration to those of ODN 4 and ODN 6 were obtained for the products of ODN 2 in Na⁺ solution. Therefore, ODN 2 showed efficient photoreaction in two diagonal loops in the presence of Na⁺, suggesting the formation of two antiparallel G-quadruplexes by ODN 2. These two bands showed different intensity, suggesting the two diagonal loop structures were not exactly the same. In contrast, gel analysis showed that no efficient photoreaction occurred in the presence of K⁺. One band with similar gel migration to that of ODN 6 was observed in Li⁺ solution. The CD spectra revealed that ODN 2 formed a stable hairpin structure. Therefore, we assumed that this band might be a result of the photoreaction cleavage of the hairpin structure. The results obtained for ODN 1 and ODN 2 led us to conclude that efficient photoreaction occurred in the diagonal loop of...
antiparallel G-quadruplexes formed by human telomeric sequences. Furthermore, the two cleavage bands of ODN 2 showed that ODN 2 might form two consecutive antiparallel G-quadruplexes. In contrast, no efficient photoreaction was found in these loop structures of the hybrid G-quadruplexes formed in K⁺ solution.

The diagonal loop structure of an antiparallel G-quadruplex in Na⁺ solution is shown in Fig. 2. A1 was positioned between BrU(T12) and the G-tetrad, and was directly stacked with BrU(T12).

Fig. 2 NMR structure of an antiparallel G-quadruplex in Na⁺ solution (PDB code 143D, 5'-AGGGTTAGGGTTAGGGTTAGGG-3') and a stacking model of thymine (yellow), 5-bromouracil (red), and adenine (blue) in the diagonal loop region.

Considering the photoreaction mechanism of hot-spot sequences in duplex DNA, we proposed that A1 prevented rapid electron back transfer, and increased the lifetime of the anion radical formed by BrU. Therefore, an efficient photoreaction occurred in the diagonal loop of the antiparallel G-quadruplex formed by ODN 1 in Na⁺ solution. Moreover, ODN 2, which might form two consecutive antiparallel G-quadruplexes with two diagonal loops, exhibited two photoreaction cleavage sites. However, in the hybrid-1 and hybrid-2 G-quadruplex structures there were no such stacking models in the loop structures (Fig. S4, ESI†).

Subsequently, we investigated the photoreaction of ODN 3, in which one thymine in the loop located between two G-quadruplex-forming sequences was substituted with BrU. ODN 5, which contains four TTAGGG repeats, was the reference DNA. As shown in Fig. 3a, a clear cleavage band similar to that of ODN 5 was observed in the presence of K⁺. However, in Na⁺ solution, the photoreaction efficiency was lower compared with the photo-reaction efficiency observed in K⁺ solution. Moreover, in Na⁺ solution was not strong and the two thymines of the TTA linker were not well stacked with the G-tetrads of the two G-quadruplexes. Therefore, the photoreaction kinetics of ODN 3 in K⁺ solution were lower than that of ODN 2 in Na⁺ solution.

To compare the TTA loop structures of two G-quadruplex subunits formed by eight TTAGGG repeats in Na⁺ and K⁺ solutions, we constructed two consecutive structures according to a method reported previously, and compared their linker regions based on the results of molecular simulation.20 The dimeric model of two antiparallel units in Na⁺ solution and the dimeric model of hybrid-1 and hybrid-2 in K⁺ solution were generated as shown in Fig. 4. First, molecular simulation showed that the consecutive structure formed by hybrid-1 and hybrid-2 in K⁺ solution had stronger stacking interactions, in which the TTA linker was involved. In contrast, the interaction of two antiparallel G-quadruplexes in Na⁺ solution was not strong and the two thymines of the TTA linker were not well stacked with the G-tetrads of the two G-quadruplexes. Second, in the TTA loop located between the hybrid-1 and hybrid-2 structures, two adenines (A19 and A37) were positioned between the stacking of thymines (BrU) and G-tetrads. Similar to what was observed for the hot-spot sequence in duplex DNA and the diagonal loop of the antiparallel G-quadruplex, these two adenines prevented rapid electron back transfer, and increased the lifetime of the anion radical formed by BrU. However, in the case of a dimeric model...
in Na⁺ solution, there was no such stacking in the TTA loop between two antiparallel G-quadruplexes. A19stacked with T30, which was located in the diagonal loop of one antiparallel G-quadruplex. Finally, in the TTA loop located between the hybrid-1 and hybrid-2 structures, the second thymine, T18 (BrU), of the TTA linker was located quite close to the C1' of the first deoxythymidine, T17, which can facilitate hydrogen abstraction during photoreaction. Overall, the photoreaction efficiency was determined both by the stacking models of loop bases with G-tetrads and by the TTA loop structure itself. Although in Na⁺ solution the stacking of BrU, adenine and G-tetrad in the linker region was not as strong as the stacking in K⁺ solution, electron transfer from G-tetrad to the linker loop region may still occur. However, adenines which were not located between BrU and G-tetrad could not prevent the electron back transfer and increase the lifetime of the anion radical formed by BrU.

In conclusion, we have demonstrated that 5'-end FAM-labeled DNA can be used to easily analyze the photoreaction of 5-halouracil-containing DNA via denaturing gel analysis. Using this method, we discovered that the photoreaction of 5-bromouracil-containing DNA could be used to probe the loop structure of G-quadruplexes formed by longer human telomeric DNA. Different photoreaction efficiencies were observed for eight TTAGGG repeats in K⁺ and Na⁺ solutions, indicating different loop structures among two consecutive G-quadruplexes. Dimeric models of the consecutive structures in K⁺ and Na⁺ solutions were also generated by molecular simulation, and the loop regions located between two G-quadruplex units were compared. Therefore, our method can be used to detect higher-order G-quadruplex formation in long human telomeric DNA. Furthermore, as efficient photoreaction was observed in K⁺ solution, which is biologically relevant, this method may be used to probe the G-quadruplex formation of telomeric DNA in cells.

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Notes and references