

Stabilization of telomeric i-motif structures by (2' S)-2'-deoxy-2'-C-methyl-cytidine residues

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Abstract: G-quadruplex and i-motif are tetraplex structures present in telomeres and promoter regions of oncogenes. The possibility of producing nanodevices with pH-sensitive functions has triggered the interest for modified oligonucleotides with improved structural properties. We synthesized C-rich oligonucleotides carrying conformationally restricted (2' S)-2'-deoxy-2'-C-methyl-cytidine units. The effect of this modified nucleoside on the stability of intramolecular i-motifs related to vertebrate telomere was investigated by means of spectroscopic methods (UV, CD and NMR). The replacement of selected positions of the C-core by the appropriate C-modified residues induces the formation of stable intercalated tetraplexes at pHs near neutrality. The study demonstrates the possibility of enhancing the stability of i-motif by chemical modifications.

Introduction

Nucleic acids can adopt secondary structures that play important roles in several cellular functions. Among them, i-motifs are present in oncogenic regions as well as in human telomeric DNA regions indicating their possible role during the regulation of oncogene expression at the transcription level.^[1-3] Recently, i-motif structures were also found in the CCG triplet repeats^[4], in the G₂C₄ hexanucleotide repeat expansion mutation sequence,^[5] and in centromeric sequences.^[6] In addition, several proteins

have been studied that specifically bind C-rich telomeric sequences.^[7-10]

These i-motifs are intercalated tetraplex structures formed by the association of a hemiprotonated cytosine base pair (C-CH⁺), and close sugar-sugar contacts stabilized by CH⁺•••O interactions^[11] in C-rich sequences. The C-CH⁺ bond has been compared to the strong hydrogen bonds produced by equally hydrogen sharing observed in basic organic molecules with low hopping energy barriers.^[12] Moreover, there are several analytical and biomedical applications of cytosine-rich oligonucleotides^[13] exploiting the advantages of i-motif pH-dependent formation especially in the nanotechnology field.^[14,15]

Chemically modified nucleic acids are widely explored as therapeutic drugs, and diagnostic tools, in target validation for drug development and in other applications in the field of biomedicine and nanotechnology. In addition, chemically modified nucleic acids are good molecular models for the elucidation of nucleic acids structures, as well as excellent tools for physicochemical studies and for probing protein-nucleic acids interactions. To date, a large number of modifications have been developed and analyzed, providing important information for the design of new derivatives with tailor-made properties, including pairing specificity, nuclease resistance and cellular uptake.^[16]

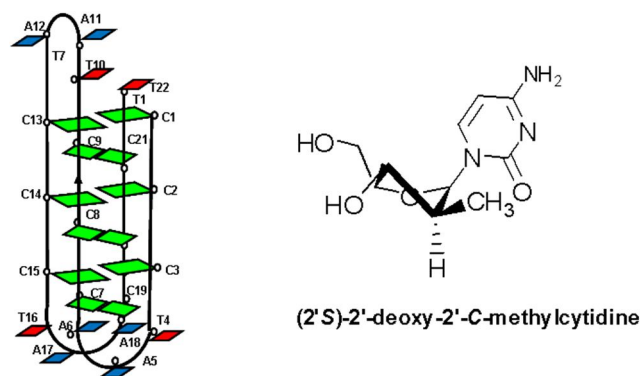
Several C-rich modified oligonucleotides have been developed, with the aim of increasing nuclease resistance and i-motif stability.^[17] In most of cases, modifications are not well-tolerated in an i-motif because they might either destabilize the structure or prevent its formation.^[17,18] Base modifications such as 5-methylcytosine and 5-hydroxymethylcytosine are permissive bases that can be adapted to the i-motif structures. They show a position-dependent stability and different behavior under experimental crowding conditions.^[19-21] These bases are important epigenetic markers in the human genome.

The i-motif structure consists of two wide and two extremely narrow grooves. The stability of this structure depends on the repulsion between adjacent negatively charged phosphates backbones across these minor grooves. Various backbone substitutions have been studied in order to form intermolecular and intramolecular i-motifs.^[17,18] Modifications such as Locked Nucleic Acid (LNA)^[22,23], Unlocked Nucleic Acids (UNA),^[24] acyclic threoninol^[25] and peptide nucleic acids (PNA)^[26] backbones have been introduced in biologically relevant i-motif structures.

The most common conformation of the sugars found in the i-motif structures is the 3'-endo. The ribose in RNA adopts a 3'-endo pucker but the i-motif is destabilized.^[27] Steric interactions such as 2'-O-Me^[27] or 2'-F^[28] also produce a decrease in the stability. However, modifications with arabinose derivatives in which the 2'-OH group points toward the major groove are well tolerated within the i-motif structure.^[29] Recently, 2'-F-araC have been studied showing stabilized i-motifs over a wide pH range without modifying the overall structure.^[30]

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To further explore the tolerance of 2'-substituents in the arabino sugar configuration, we incorporated the (2'-S)-2'-deoxy-2'-C-methyl-cytidine unit ($C_{Me}Up$)^[31-33] (Scheme 1) in a well described C-rich fragment of the vertebrate telomere (C_3TA_2)₃C₃T. This i-motif structure folds intramolecularly with four cytidine stretches connected by three TAA loops and six C-CH⁺ base pairs (Scheme 1).^[34] A similar sequence has also been reported to show a polymorphism with two possible conformations.^[35] The $C_{Me}Up$ derivative is characterized by its nuclease stability^[36] and for its influence as conformationally restricted analogue on the activity of 10-23 DNAses.^[36,37]



Scheme 1. Representation of C-rich fragment of the vertebrate telomere (C_3TA_2)₃C₃T studied in this work and chemical structure of (2'-S)-2'-deoxy-2'-C-methylcytidine unit ($C_{Me}Up$)

The introduction of carbon substituents at the 2'-position of deoxynucleotides deeply affects the sugar conformation state. In this sense, the 2'-deoxy-2'-C-methyl-nucleosides provided interesting properties since different preferred sugar conformations were observed depending on the absolute conformation at the 2'-carbon.^[38-40] The (2'-S)-2'-deoxy-2'-C-methyl-nucleosides mainly adopt the C3'-endo sugar puckering while the (2'-R)-2'-deoxy-2'-C-methyl-nucleosides prefer the C2'-endo conformation. We rationalized that (2'-S)-2'-deoxy-2'-C-methylcytidine ($C_{Me}Up$) with a methyl group in a pseudo-equatorial orientation that stabilizes a North (RNA-like) pucker was an interesting unit to introduce in i-motif structures. Therefore, in this work we replaced dC by $C_{Me}Up$ in different positions of the C-core stretches of the vertebrate telomere (C_3TA_2)₃C₃T. We selected single substitutions at 5'-terminal stretch (VT1 $C_{Me}Up$), internal positions (VT2 $C_{Me}Up$) or double substitution involved in the same internal C-CH⁺ base pairs (VT2,14 $C_{Me}Up$) (Table 1 and Figure S1).

Table 1. Sequences used in this study. C : (2'-S)-2'-deoxy-2'-C-methylcytidine unit ($C_{Me}Up$)

Name	Sequence
VTWT	CCCTAACCCCTAACCCCTAACCCCT
VT1 $C_{Me}Up$	CCCTAACCCCTAACCCCTAACCCCT
VT2 $C_{Me}Up$	CCCTAACCCCTAACCCCTAACCCCT
VT2,14 $C_{Me}Up$	CCCTAACCCCTAACCCCTAACCCCT

Results

CD spectra

First, the $C_{Me}Up$ i-motif formation was investigated by circular dichroism (CD). Figure 1 and Figure S2 shows the CD spectra recorded for all sequences at pH 5.5 and 15 °C. We obtained the characteristic i-motif CD signature with positive and negative peaks at 287 and 262 nm, respectively, for all the modified oligonucleotides prepared in this study and at different pHs. In all cases, similar CD signature as unmodified VTWT was observed, indicating that the modified nucleoside is well incorporated in the overall structure and at all pH studied.

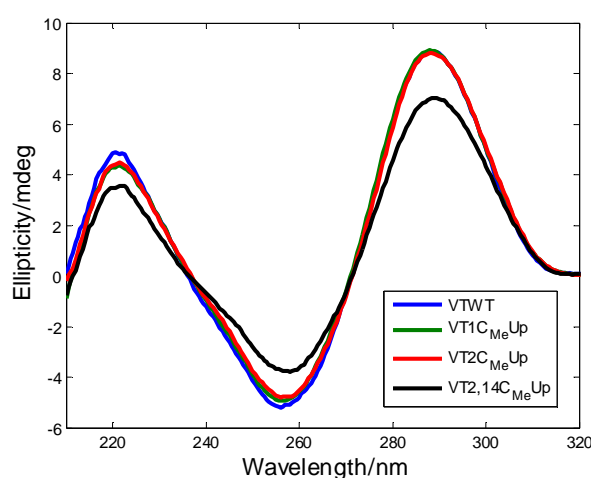


Figure 1. CD spectra of the sequences measured at pH 5.5 and 15 °C.

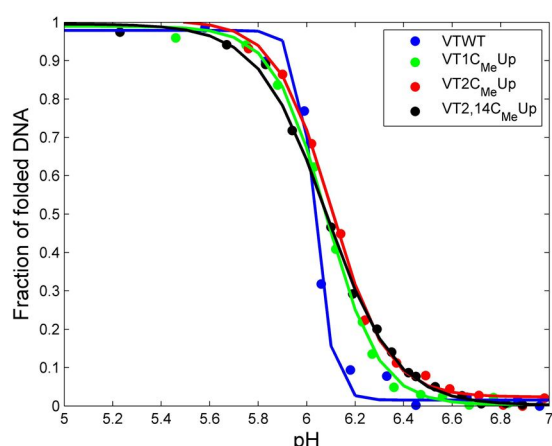
Acid-base titrations

CD spectra of $C_{Me}Up$ modified i-motif against pH variation were recorded. Acid-base titrations were carried out in the pH range of 3-7 and at 25 °C (Figure 2 and Figure S3) show the DNA folded fraction vs. pH adjusted to a sigmoidal curve. The pH transition midpoint ($pH_{1/2}$) at which 50% of the oligonucleotide is folded into an i-motif structure was determined according to the procedure previously described. In all cases, the titration curves were analyzed assuming a single protonation event in order to extract populations of folded (protonated) or unfolded (deprotonated) states. This assumption was previously checked by means of multivariate analysis (data not shown). The calculated parameters from each curve are summarized in Table 2. In general, the modifications increase slightly the pH transition midpoint. This increase is almost within the range of experimental uncertainty. The $C_{Me}Up$ derivative has a methyl group at the 2' position of the ribose. For this reason the introduction of the methyl group is not expected to have a strong impact in the pH transition midpoint. The proposed mutations

1 have a more significant effect on the pH transitional range (R_T),
 2 which varies from 0.22 (for the wild-type) to 0.5 for the
 3 VT1C_{Me}Up and VT2C_{Me}Up, and even 0.64 for the double
 4 modification. In another words, the acid-base titration curve of
 5 the unmodified sequence is very sharp indicating a strong
 6 cooperativity in the deprotonation of the i-motif. Once the first
 7 protons are removed, it is easier to remove the rest of them
 8 leading to a quick destabilization of the i-motif. On the contrary
 9 the shape of the curves obtained for the C_{Me}Up modified
 10 sequences are wider (less cooperative). These differences may
 11 explain the stabilization of the i-motif at pHs higher than the
 12 midpoint (see below).

13 **Table 2.** pH transition midpoints and transitional range for the studied DNA
 14 sequences.

Name	pH _{1/2}	pH _{10%}	pH _{90%}	R _T
VTWT	6.03±0.02	5.92	6.14	0.22
VT1C _{Me} Up	6.08±0.02	5.83	6.33	0.50
VT2C _{Me} Up	6.10±0.02	5.85	6.39	0.54
VT2,14C _{Me} Up	6.08±0.02	5.76	6.40	0.64



39 **Figure 2.** Dependence of the fraction of folded DNA with pH. The
 40 acid-base titrations were carried out at 25 °C.

44 Thermal stability of i-motif

45 Next, we studied the thermal stability of the oligonucleotides
 46 using CD and molecular absorption spectroscopies. The thermal
 47 denaturation curves were obtained by measuring the UV
 48 absorbance of the oligonucleotide solutions at 295 nm at pHs
 49 5.0-7.0 (Fig. S4). At this wavelength the denaturation of the i-
 50 motif affords a decrease in the absorbance from where the
 51 midpoint (T_m) can be determined as the minimum of the first
 52 derivative (Table 3 and Figure S4). The denaturation may also
 53 be followed at 260 nm obtaining similar results but in this case
 54 the denaturation process increases the absorbance as shown in
 55 Figures S5 and S6.

15 **Table 3.** Melting temperatures determined for the studied sequences in 20 mM
 16 phosphate buffer and 150 mM KCl at 295 nm. A two-state process has been
 17 considered. The uncertainty in the determination of T_m at pH 6.0, 5.5 and 5.0
 18 is ±0.7 °C.

Sequence	pH	T_m (°C)	ΔT_m (°C)
VTWT	6.5	≤12	-
VT1C _{Me} Up	6.5	~15	~3
VT2C _{Me} Up	6.5	~15	~3
VT2,14C _{Me} Up	6.5	18	~6
VTWT	6.0	23.9	-
VT1C _{Me} Up	6.0	24.5	0.6
VT2C _{Me} Up	6.0	27.0	3.1
VT2,14C _{Me} Up	6.0	28.5	4.6
VTWT	5.5	43.1	-
VT1C _{Me} Up	5.5	44.3	0.8
VT2C _{Me} Up	5.5	44.8	1.7
VT2,14C _{Me} Up	5.5	45.4	2.3
VTWT	5.0	54.8	-
VT1C _{Me} Up	5.0	55.5	0.7
VT2C _{Me} Up	5.0	55.4	0.6
VT2,14C _{Me} Up	5.0	54.6	-0.2

19 The determined T_m values showed that the C_{Me}Up-modified
 20 oligonucleotides slightly stabilize the i-motif structure, depending
 21 on the pH. shifting their T_m to higher values. At pH 5.0, the small
 22 changes on T_m values are not significant, indicating that the
 23 modification do not change the stability of the i-motif structure at
 24 this pH. The differences are more pronounced at pH 5.5, with an
 25 increase in thermal stability up to 2.3 °C in the double modified
 26 variant and around 1.7 °C for the single modification in
 27 VT2C_{Me}Up. At pH 6.0 there is a clear stabilization of the double
 28 modified variant ($\Delta T_m = 4.6$ °C) followed by VT2C_{Me}Up ($\Delta T_m = 3.1$
 29 °C). The shifts in T_m values in relation to that of the wild-type
 30 sequence agree with the behavior of the pH transitional range
 31 R_T mentioned above. Therefore, as this range becomes wider,
 32 the i-motif structure can be observed at higher pH values, even
 33 though as a minor species. Despite the analysis of the
 34 transitions at pH 6.5 is difficult, it is still observable for the double
 35 modified variant. Surprisingly, at pH 7, in which almost no i-motif
 36 is observed in the natural sequence, the denaturation curve
 37 measured at 295 nm of VT2,14C_{Me}Up variant (Figure S4) still
 38 shows a small hypochromic change that indicates the presence
 39 of the i-motif structure. These results suggest C_{Me}Up
 40 modification stabilizes human vertebrate i-motif structures even
 41 near neutral pH values.

42 Similar results were obtained by following the denaturation of the
 43 i-motif sequences by circular dichroism (see supporting
 44 information). At pH 5.5 and 6.0 the higher stability of C_{Me}Up-
 45 modified oligonucleotides was confirmed being the double
 46 modified variant the most stable followed by VT2C_{Me}Up and
 47 VT1C_{Me}Up. Although there are some differences in T_m values
 48 obtained by CD- and UV-absorbance methods, the relative order
 49 of stability was confirmed. At pH 6.5 the transitions are still
 50 observable but the first derivatives are too noisy and the
 51 estimated T_m 's are not reliable. At pH 7.0 only the double
 52 modified variant shows changes in the CD spectra that may
 53 confirm the presence of i-motif structure at low temperatures.

54 NMR spectra

We further analyzed the structure of the modified i-motifs by NMR spectroscopy. 1D NMR spectra show the characteristic cytosine imino protons in hemi-protonated C:CH⁺ base pairs at around 15-16 ppm (Figure 3a,c). At neutral pH i-motif formation is significantly favored for VT2,14C_{Me}Up in comparison to the non-modified sequence, which does not exhibit the characteristic i-motif signals under these conditions. 2D NOESY spectra exhibit the characteristic cross-peak pattern of i-motif (Figure S7). Although the complete assignment of the NMR spectra is beyond the scope of this study, the 2'-CH₃ signals of the modified nucleotides could be identified (0.46 and 0.81 ppm). Their NOEs with the amino protons of hemiprotonated cytosines, shown in Figure S7 indicate that these methyl groups are oriented towards the i-motif major grooves, as expected (Fig. 3e). No steric clash with neighboring residues is expected in this orientation. To confirm this point, a molecular dynamics calculation was carried out with the AMBER program. After a 1-ns molecular dynamics run, the model structure of VT2,14C_{Me}Up remains stable (see Figure S8), with the two 2'-methyl groups well accommodated in the major grooves, and the sugar of the C_{Me}Up residues in North conformations (Figure 4).

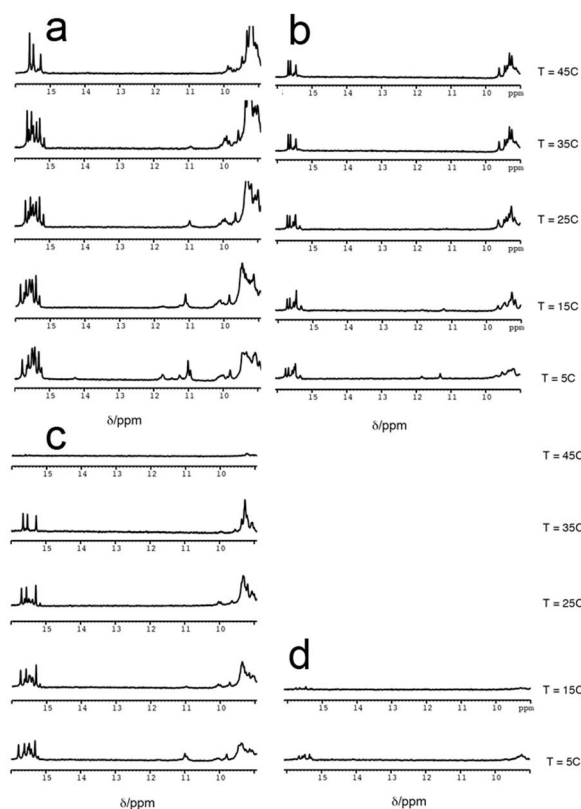


Figure 3. Exchangeable proton region of the NMR spectra at different temperatures of VT2,14C_{Me}Up at pH 5 (a) and pH 7 (c), and the unmodified human telomeric sequence at pH 5 (b) and pH 7 (d).

These results confirm that the arabinose sugar configuration is well tolerated within the i-motif structure^[29,30], and permits different substitutions at the 2' positions which are not allowed in the ribose configurations due to bad steric contacts throughout the i-motif minor groove. 2'-Fluorine substitutions provoke the most dramatic effect, since the fluorine electronegativity causes favourable changes in the charge distribution within the sugar. These changes in charge distribution are not expected in the case of 2'-CH₃ substitutions and, consequently, the stabilization effect is not so pronounced. However, (2'*S*)-2'-deoxy-2'-C-methyl-cytidine has a strong tendency to adopt a C3'-*endo* conformation, which is usually preferred in i-motifs. Preorganization of sugar in the proper conformation is most probably the reason of the observed stabilization of i-motif.

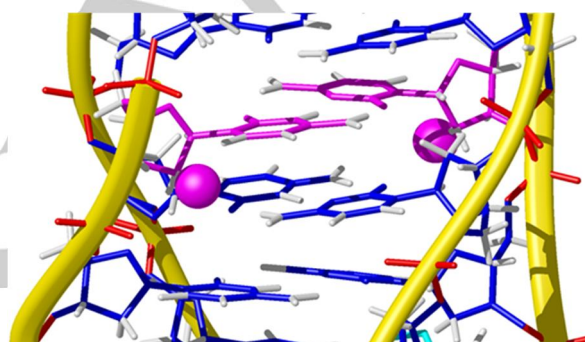


Figure 4. Model of VT2,14C_{Me}Up based on the structure of the telomeric sequence. (2'*S*)-2'-deoxy-2'-C-methyl-cytidines are shown in magenta with 2'-CH₃ groups displayed as spheres.

Discussion

Due to the potential applications of i-motif structures, there is an increasing interest in designing of modified cytosine derivatives capable to form stable i-motif particularly at near neutral pHs.^[13-15] One of the first studied modifications was the backbone-modified derivatives known as peptide nucleic acids.^[26] Some stabilization of the i-motif at neutral pH was observed, but these results were obtained in short tetrameric structures and are difficult to extrapolate to biologically relevant intramolecular i-motifs. As cytosine may be subjected to epigenetic modifications, the presence of 5-methyl-C and 5-hydroxymethyl-C in the C-rich telomere sequence has been studied, thus showing small changes in the stability of the modified i-motifs.^[19-21] Interestingly, it has been observed that the inclusion of ribocytidines destabilizes i-motifs^[27] but arabinocytidines provoke the opposite effect.^[29] It has also demonstrated that the presence of 2'-fluoro-araC produces very stable i-motifs even at neutral pH,^[30] mainly due to favorable electrostatic interactions provoked by the electrowithdrawing properties of the fluorine atom. In this work we have studied the effect of the (2'*S*)-2'-deoxy-2'-C-methylcytidine derivatives on the stability of the i-motif, demonstrating that the substitution of a -H for a CH₃ group gives

rise to more stable i-motifs and that these modified structures can also be observed at pH near neutrality. We replaced dC by C_{Me}Up in different positions of the C-core stretches of the C-rich fragment of the vertebrate telomere (C₃TA₂)₃C₃T selecting single substitutions at 5'-terminal external position (VT1C_{Me}Up), and single (VT2C_{Me}Up) and double replacements (VT2,14C_{Me}Up) in the next internal position. Thermal denaturation studies followed by UV- and CD-analyses showed that the most stable variant at mild acidic pH was the double modified sequence (VT2,14C_{Me}Up) followed by the single modified sequence at internal position (VT2C_{Me}Up). The single modified sequence at external position (VT1C_{Me}Up) is the less stable but still it is more stable than the unmodified sequence at mild acidic pH. The most evident demonstration of the existence of i-motif at neutral pH for the double modified variant comes from the presence of the exchangeable protons at pH 7.0 in the NMR spectra of VT2,14C_{Me}Up up to 35 °C. Moreover, UV- and CD-spectroscopies confirm a distinct behavior of the double modified variant at pH 7.0 in accordance with the presence of an i-motif structure for VT2,14C_{Me}Up at low temperatures.

A remarkable observation found is the low influence of the C_{Me}Up-modification on the pH midpoint transition values which oscillates between 6.03 and 6.10 (Table 2). On the contrary, other cytidine derivatives such as 5-methyl-C^[20] and 2'-Fluoro-araC^[30] induce larger variations in the i-motif p_{H1/2} values that explain the changes in the stability. In our case the C_{Me}Up modification induces an alteration in the acid-base profile of i-motifs indicating that the protonation and deprotonation of cytidines are less cooperative. Therefore, the observed trend is that at pHs higher than the midpoint the stability of the i-motifs increases while at lower pHs it decreases.

It is interesting to compare these results with the stabilization provoked by the 2'-F-araC derivatives. Both studies confirm that i-motifs tolerate well 2'-substitutions in the arabino sugar configuration. However, 2'-F-araC derivatives stabilize i-motif at all pHs while (2'S)-2'-deoxy-2'-C-methyl-cytidine do not stabilize i-motif at pH 5.0. In contrast to the 2'-F substitution, the methyl group at 2' should not alter the charge distribution in the sugar and, therefore, the favorable electrostatic interactions involving 2'-F-arabinoses are absent. Consequently, the stabilization observed for (2'S)-2'-deoxy-2'-C-methyl-cytidine substitution is less pronounced. In this case, the preference of C_{Me}Up derivatives for the north-sugar conformation is probably the main factor in the stabilization. These results open the door to the design of other stabilizing C- derivatives.

Conclusions

In conclusion, we studied the effect of the replacement of the dC by C_{Me}Up in the ability to form intramolecular i-motifs. The relative stability of the modified structures have been analyzed by UV, CD and NMR spectroscopies, showing that C_{Me}Up residues induce a stabilization of the i-motif being the C_{Me}Up: C_{Me}Up pair more stable than the C_{Me}Up: dC pair. This

stabilization could be used to modulate the stability of i-motif structures at mild acidic to neutral pH values.

Experimental Section

Oligonucleotide synthesis

The oligonucleotides listed in Table 1 were prepared on a DNA/RNA Applied Biosystems 394 synthesizer by solid-phase 2-cyanoethylphosphoramidite chemistry at a 0.2 μmol scale. Sequences containing C_{Me}Up were prepared using (2'S)-5'-O-(4,4'-dimethoxytrityl)-2'-deoxy-2'-C-methyl-N⁶-benzoylcytidine-3'-O-(2-cyanoethyl-N,N-disopropyl)-phosphoramidite that was prepared as reported previously^[32,33].

The oligonucleotides were synthesized in the DMT-ON mode. After the standard ammonia treatment (conc. aq. NH₃, 55 °C, 6h), the oligonucleotides were purified using Glen-Pack™ DNA purification cartridges (Glen Research), analyzed by HPLC on a Semipreparative column: X-Bridge™ OST C₁₈ (10x50 mm, 2.5 μm); 10 min linear gradient from 0 % to 30%, flow rate 2 mL·min⁻¹; solution A was 5% acetonitrile (ACN) in 0.1 M aqueous triethylammonium acetate (TEAA) and B 70% ACN in 0.1 M aqueous TEAA and confirmed by Matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF) spectra (Table 1, Figure S1).

UV Melting studies

UV thermal denaturation data^[41] were obtained on a JASCO V-650 spectrophotometer equipped with a Peltier temperature control. Oligonucleotides were resuspended in 20 mM citrate/phosphate buffer pH 5.0 and 5.5, phosphate buffer pH 6.0, 6.5 and 7.0 containing 50 mM KCl to form solutions with a 5 μM concentration. These solutions were heated at 85 °C and slowly cooled down to room temperature for annealing. The resulting solutions were stored at 4 °C overnight. The thermal denaturing curves were obtained following the change of the absorption at 295 nm from 10 °C to 80 °C with a linear temperature ramp of 0.5 °C·min⁻¹. The dissociation melting temperatures were determined as the midpoint of the transition (T_m) using the first derivative of the experimental data. T_m values were determined as an average of three experiments.

CD spectroscopy

Oligonucleotides were resuspended in 20 mM citrate/phosphate buffer and 50 mM KCl at pH 5.0, 5.5 and in phosphate buffer at pH 6.0, 6.5 and pH 7.0 to form solutions with a concentration of 1.3 μM. These solutions were heated at 85 °C and slowly cooled down to room temperature for annealing and stored at 4 °C overnight. The thermal denaturing curves were obtained in a single experiment following the change of the CD absorption at 280 nm from 5 °C to 80 °C with a linear temperature ramp of 0.5 °C·min⁻¹ on a JASCO spectropolarimeter J-815V equipped with a Julabo F-25/HD temperature control unit. Spectra were registered at 15 °C over a range of 210-350 nm with a scanning speed of 100 nm·min⁻¹, a response time of 4 s, a 0.5 nm data pitch and a 1nm bandwidth. CD absorption at 280 nm data as a function of temperature were analysed as described elsewhere.^[42]

Acid-base titrations

Acid-base titrations^[25,43] of the sequences were monitored using the Jasco spectropolarimeter above mentioned. Experimental conditions were as follows: 25 °C and 20 mM phosphate buffer with 50 mM KCl. Titrations were carried out by adjusting the pH value of solutions containing 2 μM of oligonucleotides in a Hellma quartz cell (10 mm path length, 3 mL). CD spectra were recorded in a pH-stepwise fashion. The ellipticity values at 280 nm were converted into "fraction of folded DNA" using the equation:

$$\text{fraction of folded DNA} = 1 - \frac{BS_{\text{fold}} - \text{ellipticity}_{\text{pH}}}{BS_{\text{fold}} - BS_{\text{unfold}}}$$

Where ellipticity_{pH} is the ellipticity at 295 nm at a given pH, and "BS_{fold}" and "BS_{unfold}" correspond to the baseline values of the folded and unfolded species, respectively. These chosen baseline values correspond to pH values around 3.5 and 7.5, respectively. The transition

midpoint of each transition ($pH_{1/2}$) was obtained by plotting the "fraction of folded DNA" vs. pH, and fitting to a sigmoidal using the Curve Fitting tool included in the Optimization Toolbox for Matlab (MathWorks, Natick, MA, USA). From the calculated values, the pH transitional range (R_T , range between 10% and 90% of the unfolded structure) was also determined. The validity of this two-state transition process was validated prior to the calculations using multivariate analysis.

NMR spectroscopy and molecular modeling of structures

Samples for NMR experiments were dissolved in either D_2O or 9:1 H_2O/D_2O in 10 mM sodium phosphate buffer. The sample concentration was 0.4 mM. The samples were annealed before the NMR experiments. Experiments were carried out at different pH values, ranging from 5.0 to 7.0. The pH was adjusted by adding aliquots of concentrated solution of either DCl or NaOD. All NMR spectra were acquired on Bruker Advance spectrometers operating at 600 and 800 MHz, equipped with cryoprobes and processed with the TOPSPIN software. NOESY spectra were acquired with mixing times of 150 and 250 ms. TOCSY spectra were recorded with the standard MLEV-17 spin-lock sequence and a mixing time of 80 ms. In most of the experiments in H_2O , water suppression was achieved by including a WATERGATE module^[44] in the pulse sequence prior to acquisition. Molecular structures were calculated with the SANDER module of the molecular dynamics package AMBER.

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Keywords: i-motif, (2'-S)-2'-deoxy-2'-C-methyl-cytidine, telomere, Nuclear Magnetic Resonance, Circular Dichroism, thermal denaturation studies

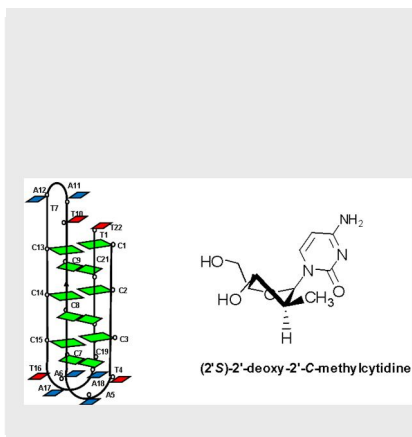
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Stabilization of telomeric i-motif structures by (2'S)-2'-deoxy-2'-C-methyl-Cytidine residues