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# Stability and free energy calculation of LNA modified quadruplex: a molecular dynamics study

Amit Kumar Chaubey · Kshatresh Dutta Dubey ·  
Rajendra Prasad Ojha

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**Abstract** Telomeric ends of chromosomes, which comprise noncoding repeat sequences of guanine-rich DNA, which are the fundamental in protecting the cell from recombination and degradation. Telomeric DNA sequences can form four stranded quadruplex structures, which are involved in the structure of telomere ends. The formation and stabilization of telomeric quadruplexes has been shown to inhibit the activity of telomerase, thus establishing telomeric DNA quadruplex as an attractive target for cancer therapeutic intervention. Molecular dynamic simulation offers the prospects of detailed description of the dynamical structure with ion and water at molecular level. In this work we have taken a oligomeric part of human telomeric DNA, d(TAGGGT) to form different monomeric quadruplex structures d(TAGGGT)<sub>4</sub>. Here we report the relative stabilities of these structures under K<sup>+</sup> ion conditions and binding interaction between the strands, as determined by molecular dynamic simulations followed by energy calculation. We have taken locked nucleic acid (LNA) in this study. The free energy molecular mechanics Poisson Boltzman surface area calculations are performed for the determination of most stable complex structure between all modified structures. We calculated binding free energy for the combination of different strands as the ligand and receptor for all structures. The energetic study shows that, a mixed hybrid type quadruplex conformation in which two parallel strands are bind with other two antiparallel strands,

are more stable than other conformations. The possible mechanism for the inhibition of the cancerous growth has been discussed. Such studies may be helpful for the rational drug designing.

**Keywords** G-quadruplex · LNA · MD simulation · Free energy calculation

## Introduction

Telomeric DNA occurs at the ends of eukaryotic chromosomes. It protects genome from degradation and participates in chromosomal fusion and recombination [1]. Human telomeric DNA consists of guanine-rich tandem repeats d(TTAGGG)<sub>n</sub>, and in human somatic cells is ~5–8 kb in length [2]. It is in duplex form for most of this length, with the exception of the 3' terminal 150–200 nucleotides, that comprise a single stranded overhang [3]. Due to the inability of DNA polymerase to fully copy the 3' ends, telomeric DNA progressive shortens during replication [4]. The end replication problem is overcome by the expression of the enzyme telomerase, which adds hexanucleotide repeats to the 3' end of the single stranded DNA, thereby preventing shortening of telomeres. Telomeres have been shown to be activated in 80–85% of human cancer cells [5], have suggested playing a key role in maintaining the malignant phenotype by stabilizing telomer length and integrity [6].

The structure and stability of telomeres are of great research interest as they are closely related with cancer [7–10], aging [11, 12] and genetic stability [13–15]. Single stranded G-rich telomeric sequences at the end of chromosomes can fold into intramolecular quadruplex structures, a DNA secondary structure consisting of stacked

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A. K. Chaubey · K. D. Dubey · R. P. Ojha (✉)  
Biophysics Unit, Department of Physics, DDU Gorakhpur  
University, Gorakhpur 273009, Uttar Pradesh, India  
e-mail: rp\_ojha@yahoo.com

G-tetrad planes connected by a network of Hoogsteen hydrogen bonds and stabilized by monovalent cations, such as Na<sup>+</sup> and K<sup>+</sup>. Human telomeric DNA repeats are highly conserved, which has been suggested to be related to their ability to form DNA G-quadruplexes [10–16]. The formation and stabilization of the DNA G-quadruplex in the human telomeric sequence have been shown to inhibit the activity of telomerase. Thus the telomeric DNA quadruplex has been considered to be an attractive target for cancer therapeutic intervention [7, 9, 10, 17–19].

The folding and formation of short telomeric G-quadruplex structures have been studied extensively by a variety of biophysical, structural and chemical probe methods. Structural information on the intact human telomeric DNA G-quadruplex formed under physiologically relevant conditions is necessary for structure based rational drug design. The human single stranded telomer end which can potentially fold into a number of four repeat quadruplexes, is a longer and more stable than in many other vertebrates. The potential role of quadruplexes in vivo has been highlighted with the recent development of therapeutic strategies designed to stabilize telomeric ends as G-quadruplex structures using specific small molecules, which can destabilize telomer maintenance in tumor cells [20–23].

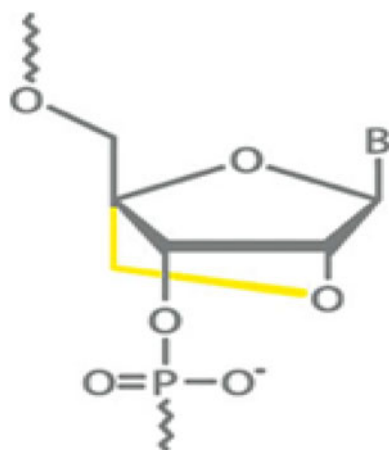
The crystal structure of a four-repeat human telomeric sequence d[AG<sub>3</sub>(T<sub>2</sub>AG<sub>3</sub>)<sub>3</sub>] crystallized from a potassium ion environment, shows that it forms an intramolecular G-quadruplex structure in which all four phosphate backbone strands are parallel to each other and the tetrads stabilized by the presence of K<sup>+</sup> ions in the central cavity [24]. Several NMR solution studies of the folded structure of the intramolecular human telomeric G-quadruplex formed in K<sup>+</sup> solution, demonstrate a novel hybrid type folding topology with mixed parallel-antiparallel G-strands [25]. This hybrid type G-quadruplex structure appears to be the physiologically relevant conformation of the human telomeric DNA. This folding of structure has also been independently reported by two other groups [26, 27].

The intramolecular human telomeric quadruplex can attain antiparallel basket-type, antiparallel chair-type, parallel propeller-type and mixed hybrid-type conformations, depending on the nucleotide length and ionic conditions [24, 25, 28, 29]. Literature based on NMR data shows that the Tel-22 sequence forms a well defined basket type intramolecular G-quadruplex in the presence of Na<sup>+</sup> ions [28]. From the platinum cross-linking method, it showed that the antiparallel basket-type conformation forms in the presence of both Na<sup>+</sup> and K<sup>+</sup> ions [29]. Another report based on the I-radioprobe technique supported the formation of the antiparallel chair-type conformation in the presence of K<sup>+</sup> ions and the antiparallel basket-type conformation forms in the presence of Na<sup>+</sup> ions [30]. Shafer et al. also found the antiparallel chair-type conformation to

be present under K<sup>+</sup> ions conditions with the chemical ligation method [31]. An X-ray crystal study of Tel-22 found the parallel propeller-type conformation in the presence of K<sup>+</sup> ions [24]. Another report based on simulations showed that Tel-22 adopts the chair-type conformation in the presence of K<sup>+</sup> ions [32]. Ambrus et al. [25] using 1D NMR analysis showed that, in the presence of K<sup>+</sup> ions, the Tel-22 sequence adopts two conformations, which were found to be the antiparallel and mixed hybrid-type G-quadruplexes according to CD data.

Nucleic acid is central to transmission, expression and conservation of genetic information. Consequently, high affinity binding of complementary nucleic acids has applications in biotechnology and medicine. In this context there are the development of nucleic acid with chemical modification rendering them high affinity and stability, since unmodified DNA or RNA oligonucleotides have moderate affinities for complementary targets and low stability in biological fluids [33]. Result of such modification is Locked Nucleic Acid (LNA) molecule, where the furanose conformation is chemically locked in an RNA like (C3'-endo) conformation by introduction of a 2'-O, 4'-C methylene linkage (Fig. 1). LNA nucleotides substantially increase the thermal stability when incorporated into both DNA duplexes and triplexes and LNA has the potential for applications in drug design. With such immense potential, it is of fundamental importance to understand the structural nature of complexes formed by LNA with DNA or RNA. There have been some studies in this direction, mainly with NMR spectroscopy and X-ray crystallography, where in most cases only selected nucleotides has been replaced by LNA nucleotide in a regular RNA or DNA based duplex [34–36]. These studies have provided useful information on the nature of duplex structure with the introduction of LNA. It was demonstrated that LNA modifications stabilize the tetrameric quadruplexes of TG<sub>3</sub>T when it is fully modified [37]. Also fully modified RNA and 2'-O-methyl-RNA quadruplexes are more stable than their unmodified d(TG<sub>4</sub>T) counterpart [38]. However, there are no structures showing the effect of LNA on 24-mer monomeric human telomeric quadruplex structures. Thus, we decided to investigate the stability and existence of LNA modified human telomeric quadruplex structures with K<sup>+</sup> ions.

We have used molecular modeling methods to build models of G-quadruplex arrays of monomeric structures formed from extended human telomeric DNA. Since different reports have suggested variable conformation of human telomeric quadruplex in the presence of K<sup>+</sup> ions, we used seven different types of human telomeric d(TAGGGT)<sub>4</sub> modified quadruplex conformations (Fig. 2), viz., parallel propeller type (all strands are parallel), basket-type, chair-type and mixed hybrid types, for simulation studies to understand their relative stabilities and



**Fig. 1** Chemical structure of locked nucleic acid, in which a methylene bridge connecting the O2' atom with C4' atom locks the ribose ring

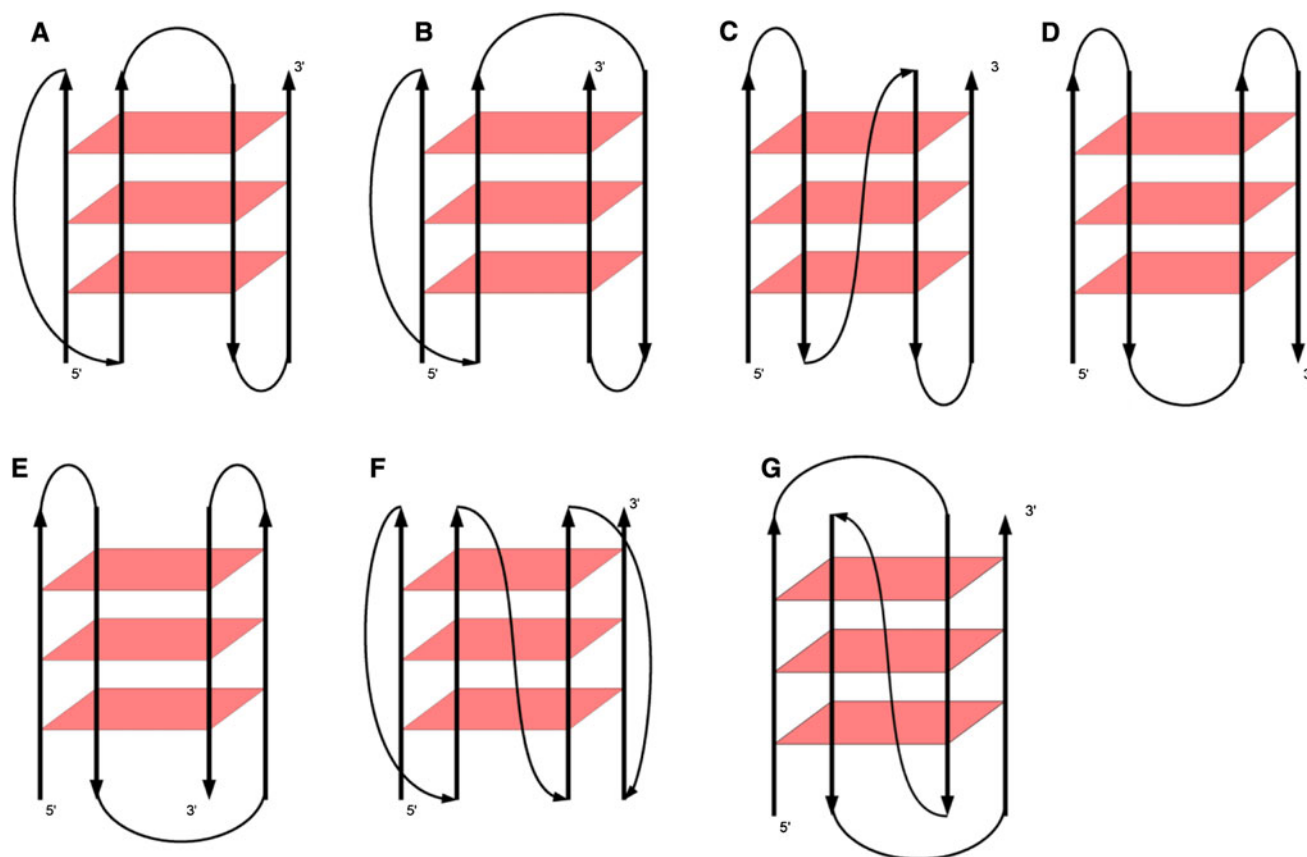
interaction between strands. Furthermore, to characterize the most energetically favorable quadruplex structures, we estimated the binding free energy and the entropy. As only a few earlier reports based on MD simulation of human telomeric quadruplex and the role of LNA with DNA in duplex [39, 40], triplex and quadruplex, this is the first

exhaustive study providing detailed insight into the fully LNA modified different type of monomeric quadruplex structures.

## Methodology

### Model generation

The crystal structure [24] of the 22-mer human telomeric DNA d [AG3(T2AG3)3] (PDB code 1KF1) was used as the primary unit for the construction of the higher-order models. It contains three stacks of guanine tetrads. The extreme 5' and 3' thymine base were first added at start and end of the sequence respectively, to generate 24-mer d (TAGGGT)<sub>4</sub> structures. We have generated seven different types of modified structures (Fig. 2), where (LNA) was considered for modification. For modification, all the nucleotides were changed into LNA. In seven generated modified structures, four are mixed hybrid-type (MOD1, MOD2, MOD3 and MOD4), one basket-type (MOD5), one chair-type (MOD6) and one parallel propeller-type (MOD7). In MOD1 and MOD2, strand 1-2-4 are parallel and strand 3 are antiparallel (Fig. 2), in MOD3 and MOD4



**Fig. 2** Schematic representations of human telomeric quadruplex conformations: **a–d** mixed hybrid type, **e** basket type, **f** chair type, and **g** parallel propeller type

**Table 1** Summary of the model names, nature of structures and production time of LNA modified 24-mer human telomeric quadruplex structures

Model	Nature and structure of modified quadruplex	Production time (ns)
MOD1	Parallel-1-2-4 antiparallel-3a (mixed hybrid-type)	20
MOD2	Parallel-1-2-4 antiparallel-3b (mixed hybrid-type)	20
MOD3	Parallel-1-4 antiparallel-2-3a (mixed hybrid-type)	20
MOD4	Parallel-1-4 antiparallel-2-3b (mixed hybrid-type)	20
MOD5	Parallel-1-3 antiparallel-2-4a (basket-type)	20
MOD6	Parallel-1-3 antiparallel-2-4b (chair-type)	20
MOD7	Parallel propeller-type	20

which is also a mixed hybrid-type structure, strand 1–4 are in parallel and strand 2–3 are in antiparallel (Fig. 2), in MOD5 and MOD6, strand 1-3 are in parallel and strand 2–4 are in antiparallel (Fig. 2) and in seventh modified structure (MOD7), all strands are in parallel (Fig. 2). Summary of the model names, their types and production time are listed in Table 1.

#### Force field parameter generation

Since the force field parameters for the modified nucleotides (LNA) are not available in literature, therefore, their parameters are generated using the GAUSSIAN 03 and RESP [41] program of AMBER10. All the ab-initio calculations are done at the DFT (B3LYP) level of theory with G-31G (d) basis set, using GAUSSIAN suite. Three letter codes for all the fitted nucleosides were developed to standardize the naming of the modified nucleosides in pdb files. The parameters for the modified nucleotides are available as supplementary material (Table S1).

The nomenclature of the modified nucleotides used in the text is as follows.

Modified adenine: LCA

Modified guanine: LCG

Modified thymine: TL5 and TL3

Where thymine is the ending nucleotide in all quadruplexes.

#### Simulation and equilibration

The X-ray structure shows a vertical alignment of consecutive K<sup>+</sup> ions along the axis within the central core of the structure, in the middle between the G-tetrad. Thus, the

ions were retained in the positions as observed in the crystal structure [24]. We placed 5 K<sup>+</sup> ions to the central channel of all the models manually. The models were solvated in a periodic TIP3P water box [42] at least 10 Å from any solute atom. Additional positively charged K<sup>+</sup> counter ions were included in the system to neutralize the charge on the DNA backbone. We have used recent amber force field ff03 [43–45] for parameter generation.

The simulation protocols were consistent for all of the systems. The systems were annealed from 0 to 300 K for 600 ps with continuous decrease of restrain (force constant was decreased from 100 to 5 kcal/mol Å<sup>2</sup> in six steps). The final stage of equilibration involved upto 1,200 ps runs using a force constant of 2 kcal/mol Å<sup>2</sup> on the solute and inner ions to fix them during equilibration. The final production run was carried out without any restrain on the system for 20 ns and coordinates were saved after every 10 ps for analysis of their trajectories.

All calculations were carried out with the SANDER module of AMBER10 [46]. Periodic boundary condition has been applied using the particle mesh Ewald (PME) method to treat long-range electrostatics. Hydrogen bonds were constrained using SHAKE [47]. A time step of 2 fs and a direct space non-bonded cut-off of 10 Å were used. The Langevin coupling with a collision frequency of 2.0 was used for temperature regulation. A constant pressure of 1 atm. has been attained with isotropic molecule based scaling with a relaxation time of 2 ps. The trajectories were analyzed using the PTRAJ module available in the AMBER10 and visualized by means of the CHIMERA program [48].

#### MM-PBSA calculation

We performed standard MM-PBSA (Molecular Mechanics Poisson-Boltzman, surface area) [49] method for free energy calculation. Here, the total free energy of binding is expressed as the sum of the contribution from the gas phase and solvation energy and an additional term of solute entropy. This can be expressed by following equation

$$\Delta G_{TOT} = \Delta E_{GAS} + \Delta E_{SOLV} - T\Delta S$$

where  $\Delta E_{GAS}$  is the total gas phase energy given by

$$\Delta E_{GAS} = \Delta E_{INT} + \Delta E_{VDW} + \Delta E_{ELEC}.$$

Here  $\Delta E_{INT}$  corresponds to bond, angle and torsion terms in the molecular mechanical force field.  $\Delta E_{SOLV}$  is the total solvation energy (polar and non polar), and  $T\Delta S$  corresponds to solute entropy effect. The detail of these terms can be found in our recent publication [50]. Analysis is done for the last 6 ns (14–20 ns) trajectory of all the complexes. The snapshots for these quadruplex are extracted at intervals of 20 ps. Prior to the analysis, all



water molecule and K<sup>+</sup> ions were stripped from the trajectory. Solvation free energy is computed as the sum of polar and nonpolar contributions using a continuum solvent representation.

The polar contribution is calculated by Molsurf, implemented in AMBER10. The non-polar solvent contribution is estimated from a SASA dependent term

$$\Delta E_{\text{SNP}} = \gamma \cdot \text{SASA} + \beta.$$

Here  $\gamma$  is set to 0.0072 kcal/Å<sup>2</sup> and  $\beta$  to 0. The calculation for solute entropy contribution is performed with the NMODE module in AMBER10. The snapshots were minimized in the gas phase using the conjugate gradient method for 1,000 steps, using a distance dependent dielectric of 4r (r is inter atomic distance) and with a convergence criterion of 0.1 kcal/mol Å for the energy gradient.

## Results

### Dynamic structure

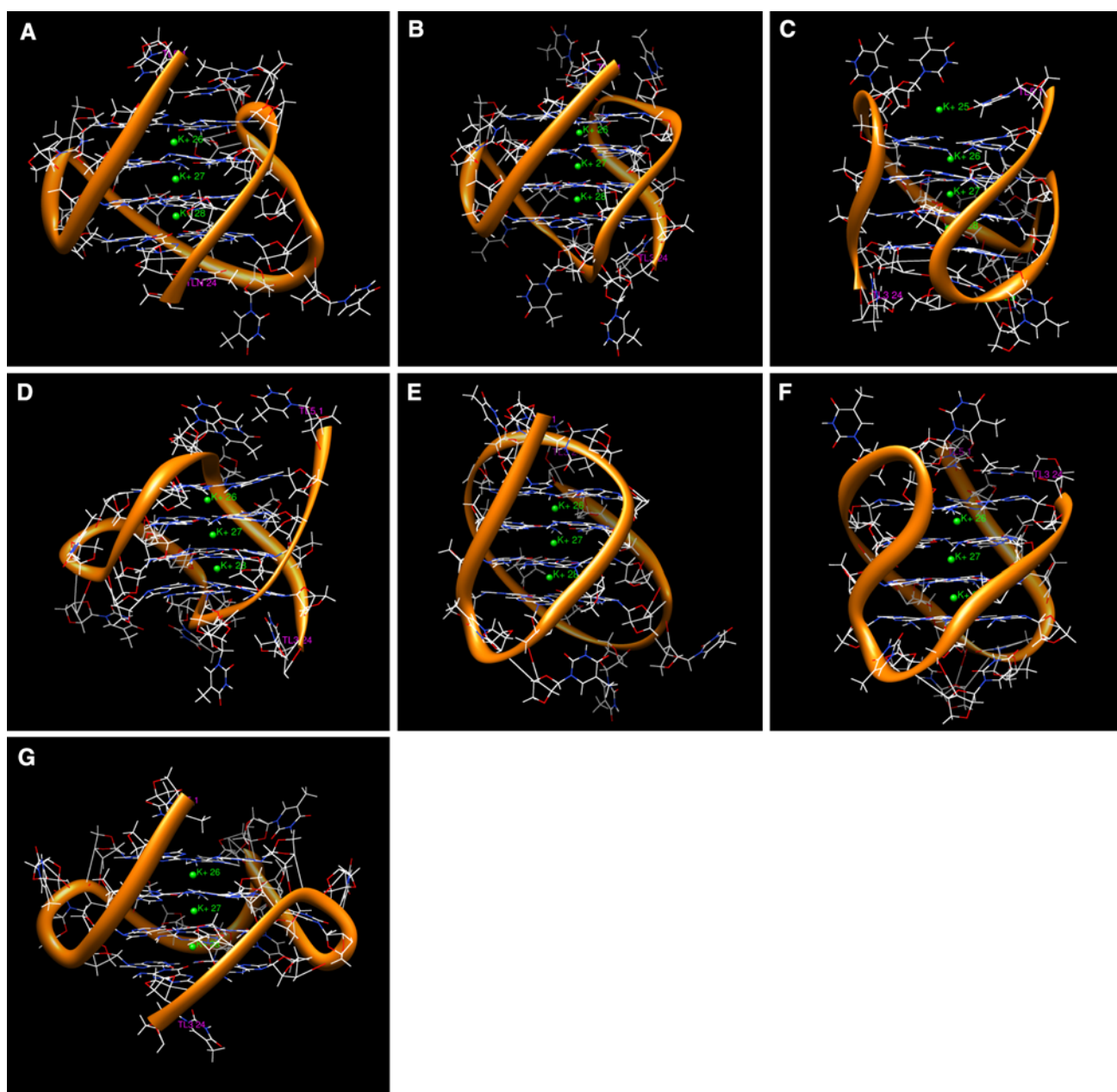
From MD simulation of all seven modified quadruplex structures, there are series of snapshots extorted at periodic intervals, out of which only the initial structures are shown in Fig. 3. During dynamics, different snapshots at interval of 10, 15 and 20 ns of MOD2 are represented in Fig. 4, while other snapshots of MOD1 and MOD3-MOD7 for interval 10, 15 and 20 ns are shown in Figure S1. During MD simulation, hydrogen bonding pattern is convincingly well maintained. The central core of stacked tetraplex has a line of potassium ions that is coincident with the helical axis. Here the metal is bipyramidally coordinated by eight equidistant carbonyl oxygen atoms. However, potassium ions which were inside quadruplex, deviate from the symmetrical geometry. From the Fig. 4, we see that deviated ions form bipyramidal binding with O6 atom of guanine bases.

The parallel three-tetrad quadruplex remains stable with two cations in the channel. This confirms capability to achieve smooth equilibrium by exchange of cations with the bulk solvent. This phenomenon was experimentally observed on the hundreds of microsecond to millisecond time scale for the central ions [51]. The stability of the quadruplex is immediately lost when ions are absent in the channel [52–54]. In the initial stage of simulation, the ions are slightly displaced towards the outside in all structures. However, after a long simulation, two ions are bound between the plane of quartet in MOD2 (Fig. 4), which shows the stability and occurrence of this model. In all other structures, except MOD2, the ions are moved outwards during a long simulation (Figure S1).

### Structural stability of the models

The root mean square deviation (rmsd) during simulation can be used as a standard of the conformational stability of a structure or a model. In this work, we intend to investigate the conformational stability of different modified G-quadruplex structures from telomeric DNA and in studying the dynamic effects of the structures. Figure 5 for MOD2 and Figure S2 for rest models compare the rmsd over the course of simulation. Each plot shows the rmsd of all-atom model (black), backbone atom only (red), base atom only (green) and middle base atom (blue). A jump in rmsd is observed during the first nanosecond which is a consequence of relaxation of the starting model. All trajectories are stabilized during simulation with relatively small fluctuation in the models. As evident from the plots of MOD1,2,4,5,6, the rmsd of all atom models are higher than that of the backbone atoms, where as in the plots of MOD3 and MOD7 the rmsd value of base atoms are higher than that of all atoms and backbone atoms. The backbone atoms are stabilized earlier during the simulation and the higher rmsd of all atoms model is a result of the wobbling effect of the nucleotide bases. All G-tetrad are held together in a coplanar array with eight hydrogen bonds which play a major role in ranking of these tetrads as most stable moieties in the models, which is shown by lowest rmsd for middle base atoms in all plots.

All models are more or less stable during the simulation except MOD7. The rmsd of MOD7 are extremely high which suggesting distortion of the structure from its starting conformation. The rmsd of backbone atoms are ~11.0 Å which also gives the rmsd value of all atoms, base atoms and middle base atoms. During the 4.7–13 ns simulation, its fluctuation remains constant with ~5.8–7.4 Å. The other more deviated structures are MOD1 and MOD3. The rmsd of MOD1 is ~8.5 Å for backbone atoms and ~9.1 Å for all atoms, which gives constant fluctuations during 14–20 ns simulation. In MOD3 the rmsd is ~8.0 Å for base atoms and ~8.3 Å for all atoms, which gives constant fluctuation during 11–19 ns simulation. In MOD4, the rmsd of all the atoms and base atoms coincide during last 6 ns run and give the rmsd of ~7.2 Å. For MOD5 and MOD6, the rmsd value is ~5.0 and ~5.4 Å for backbone atoms and for all atoms it gives ~5.5 and ~6.0 Å, which is constant during simulation. The rms deviation over the course of simulation for MOD1 and MOD3–7 are shown in Figure S2. Figure 5 shows the rmsd of different subsegments of MOD2. The rmsd value for all atoms is in the range of ~2.9 Å–3.3 Å, for backbone atoms it is ~2.6 Å and for base atoms it gives the deviation of ~1.8 Å. The model gives constant rmsd value of middle base atoms (~0.75 Å), during the simulation. This model gives lower rmsd value for all subsegments in



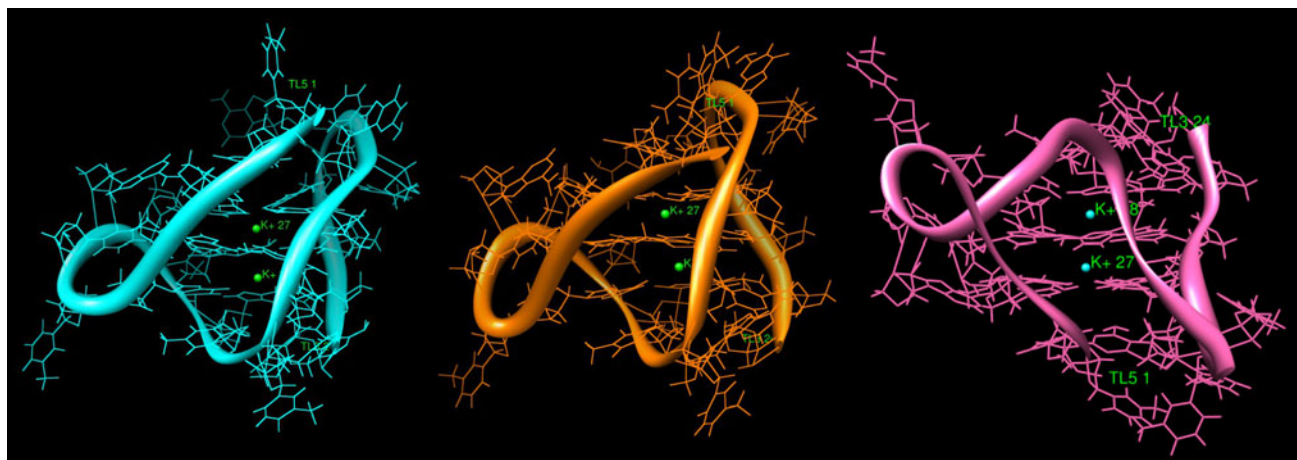
**Fig. 3** Models of modified human telomeric (a–d) mixed hybrid type quadruplex, e basket type quadruplex, f chair type quadruplex, and g parallel propeller type quadruplex structure. Three K<sup>+</sup> ion present in the central cavity are shown in green

comparison with all other models, which shows that MOD2 gives the highest stability, during simulation.

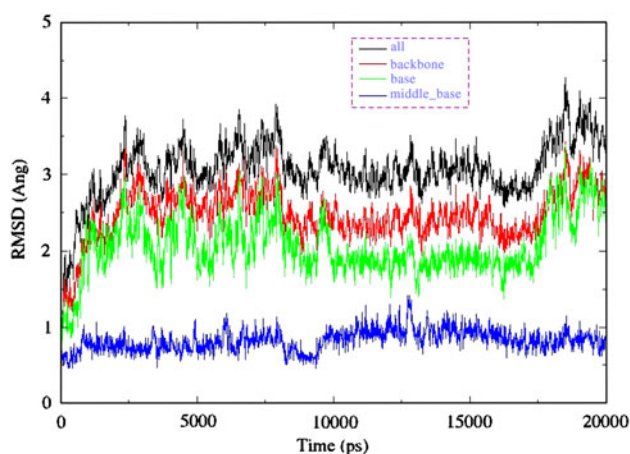
To identify the regions of high flexibility, per atom averaged mean square fluctuation (MSF) is reported for MOD2. Since, MOD2 exhibits a stable conformation; hence we performed  $\beta$  factor calculation of this model. The  $\beta$  factor regarding to MOD2 is represented in Fig. 6. Figure 6 reveals that atoms 575–685 show higher thermal fluctuation which corresponds to residue 17–20 of complex.

#### MM-PBSA calculations

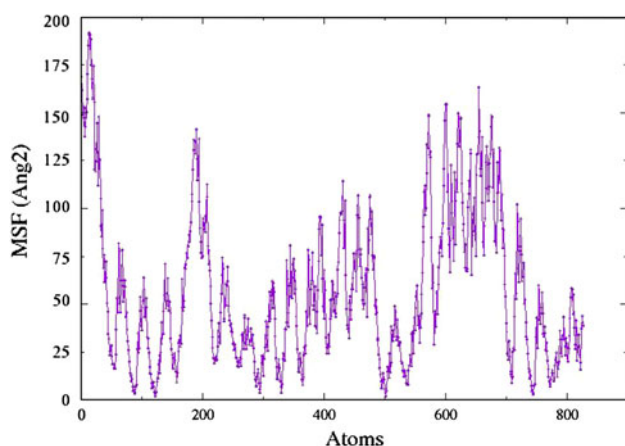
The free energy calculation using MM-PBSA method can be used to study thermo chemical properties of quadruplex models and to provide a semi quantitative estimate of their stability [55, 56]. The binding energy components for all the models of modified human telomeric quadruplex structures are summarized in Table 2. Models with lower binding energies are expected to be more stable than those with higher value. The total energy of the solute ( $E_{\text{GAS}}$ )



**Fig. 4** Snapshots of MOD2 structure during production dynamics at 10, 15 and 20 ns (snapshot for 10 ns is in cyan, 15 ns is in orange and 20 ns is in magenta colour)



**Fig. 5** RMSD plot showing the stability of the model during the MD run. RMSD values calculated for all atoms (black), backbone-only atoms (red), base only atoms (green) and middle base atoms (blue) are plotted for MOD2 structure



**Fig. 6** All-atom mean square fluctuation versus atom number for MOD2

includes the electrostatic energy, van der Waals energy derived from a Lennard-Jones potential and the internal energy. Finite difference Poisson–Boltzman equations were used to calculate the electrostatic potential field. The binding energy calculation for the quadruplex suggested a favorable contribution for van der Waals energy ( $E_{VDW}$ ) for all quadruplex models. The total binding energy ( $E_{TOT}$ ) ranged between  $-1982.94$  to  $-3482.13$  kcal/mol for all models. These values show that MOD6, chair type structure, where Ist and IIInd strand are parallel and second and fourth are antiparallel, has the least favorable binding energy ( $-1,982.94$  kcal/mol). Furthermore, MOD7, a parallel propeller type structure, gives the most favorable binding energy ( $-3,482.13$  kcal/mol) relative to other models. Since all the models have sufficient negative binding energy, they show stability in their structures.

The free energy components, for all the seven quadruplex structures (where first two strands are bound with last two strands) are summarized in Table 3. Here, electrostatic energy ( $\Delta E_{ELE}$ ) makes an unfavorable contribution while van der Waals energy ( $\Delta E_{VDW}$ ) provides favorable contributions. Moreover, the nonpolar solvation energy ( $\Delta E_{SNP}$ ) also has favorable contributions to the total binding free energy. This implies that van der Waals and nonpolar solvation energy play an important role in binding of quadruplexes. The solvent electrostatic energy ( $\Delta E_{SP}$ ) also makes favorable contribution. Moreover, we observed unfavorable entropy contributions ( $T\Delta S$ ) for all models of quadruplex. The solute entropy term of binding for all seven models ranged between  $-33.93$  to  $-48.83$  kcal/mol. The binding free energy ( $\Delta G_{TOT}$ ) of all seven structures varies from  $68.09$  to  $-4.87$  kcal/mol. These values show that, MOD3, which is a mixed hybrid type structure is stable among other proposed model because it has lowest binding energy ( $-46.74$  kcal/mol) and it has lowest



**Table 2** Total Energies of all the quadruplex structures

	MOD1	MOD2	MOD3	MOD4	MOD5	MOD6	MOD7
E <sub>ELE</sub>	1153.13 ± 65.15	1822.76 ± 66.66	1320.46 ± 76.38	1421.87 ± 118.48	1760.89 ± 78.17	1640.02 ± 70.83	1258.92 ± 132.32
E <sub>VDW</sub>	-149.77 ± 9.13	-207.52 ± 12.33	-159.64 ± 12.31	-170.77 ± 15.76	-214.64 ± 11.01	273.14 ± 19.42	-160.32 ± 16.83
E <sub>INT</sub>	2170.03 ± 22.10	2150.02 ± 27.30	2142.05 ± 23.83	2138.01 ± 15.54	2168.94 ± 19.87	3157.41 ± 27.65	2136.52 ± 19.17
E <sub>GAS</sub>	3173.39 ± 63.66	3765.26 ± 60.06	3302.88 ± 73.19	3389.11 ± 115.33	3715.19 ± 75.83	5070.57 ± 70.19	3235.11 ± 111.59
E <sub>SNP</sub>	42.41 ± 1.13	34.85 ± 0.82	40.30 ± 0.94	39.25 ± 1.04	34.13 ± 0.64	36.69 ± 0.80	40.24 ± 1.59
E <sub>SP</sub>	-6686.28 ± 62.41	-7204.11 ± 55.19	-6818.80 ± 73.79	-6897.86 ± 110.85	-7173.57 ± 52.69	-7090.21 ± 65.71	-6757.50 ± 99.08
E <sub>SOLV</sub>	-6643.87 ± 62.64	-7169.27 ± 55.66	-6778.50 ± 74.52	-6858.61 ± 117.70	-7139.44 ± 53.12	-7053.52 ± 66.06	-6717.28 ± 99.31
E <sub>TOT_ELE</sub>	-5533.15 ± 14.15	-5381.36 ± 28.18	-5498.34 ± 11.4	-5475.99 ± 13.22	-5412.68 ± 32.71	-5450.19 ± 18.13	-5498.58 ± 27.57
E <sub>TOT</sub>	-3470.48 ± 19.67	-3404.00 ± 28.18	-3475.63 ± 27.03	-3469.50 ± 15.32	-3424.24 ± 32.48	-1982.94 ± 19.70	-3482.15 ± 24.49

**Table 3** Comparison of binding free energies of quadruplex structures

	MOD1	MOD2	MOD3	MOD4	MOD5	MOD6	MOD7
ΔE <sub>ELE</sub>	2093.33 ± 37.69	2474.14 ± 48.67	2214.20 ± 45.68	2331.91 ± 78.71	2430.89 ± 35.11	2368.14 ± 49.25	2142.26 ± 73.57
ΔE <sub>VDW</sub>	-67.41 ± 5.64	-65.06 ± 7.24	-68.63 ± 6.55	-75.86 ± 5.90	-92.58 ± 6.68	-56.27 ± 4.67	-43.36 ± 9.87
ΔE <sub>INT</sub>	6.59 ± 1.71	4.73 ± 0.81	3.62 ± 1.28	6.28 ± 1.40	5.53 ± 1.38	20.52 ± 2.21	5.21 ± 1.55
ΔE <sub>GAS</sub>	2032.50 ± 34.87	2413.80 ± 43.75	2149.18 ± 46.30	2262.33 ± 74.29	2343.84 ± 35.33	2332.39 ± 47.92	2104.11 ± 67.79
ΔE <sub>SNP</sub>	-8.85 ± 0.86	-7.79 ± 0.79	-8.55 ± 0.64	-9.28 ± 0.74	-10.78 ± 0.48	-8.51 ± 0.45	-5.91 ± 1.08
ΔE <sub>SP</sub>	-2067.99 ± 35.99	-2371.85 ± 43.55	-2187.37 ± 44.35	-2297.71 ± 73.91	-2365.22 ± 26.16	-2332.25 ± 48.26	-2125.45 ± 68.21
ΔE <sub>SOLV</sub>	-2076.85 ± 36.10	-2379.65 ± 44.02	-2195.92 ± 44.40	-2306.99 ± 74.46	-2376.00 ± 26.25	-2340.77 ± 48.26	-2131.36 ± 68.77
ΔE <sub>TOT_ELE</sub>	25.34 ± 6.36	102.28 ± 21.42	26.82 ± 8.38	34.21 ± 6.83	65.66 ± 22.60	35.89 ± 7.27	16.81 ± 14.82
ΔE <sub>GAS+SOLV</sub>	-44.33 ± 6.62	34.16 ± 18.75	-46.74 ± 9.21	-44.66 ± 5.56	-32.16 ± 25.27	-8.37 ± 5.96	-27.25 ± 14.05
TΔS	-48.83 ± 11.31	-33.93 ± 11.43	-41.87 ± 8.02	-47.41 ± 8.99	-46.54 ± 11.50	-45.35 ± 9.30	-38.06 ± 10.68
ΔG <sub>TOT</sub>	4.50	68.09	-4.87	2.75	14.38	36.98	10.81

ΔE<sub>ELE</sub> = coulombic energy, ΔE<sub>VDW</sub> = vander walls energy, ΔE<sub>INT</sub> = internal energy, ΔE<sub>GAS</sub> = ΔE<sub>ELE</sub> + ΔE<sub>VDW</sub> + ΔE<sub>INT</sub>, ΔE<sub>SNP</sub> = nonpolar solvation energy, ΔE<sub>SP</sub> = polar solvation energy, ΔE<sub>SOLV</sub> = ΔE<sub>SNP</sub> + ΔE<sub>SP</sub> = total solvation energy, ΔE<sub>TOT\_ELE</sub> = ΔE<sub>ELE</sub> + ΔE<sub>SP</sub> = total electrostatic energy, ΔE<sub>GAS+SOLV</sub> = ΔE<sub>GAS</sub> + ΔE<sub>SOLV</sub> = enthalpy, TΔS = solute entropy, ΔG<sub>TOT</sub> = ΔE<sub>GAS+SOLV</sub> - TΔS = absolute free energy

All energy values are in kcal/mol

**Table 4** Hydrogen bonding distances between the atoms of G-tetrads of MOD2

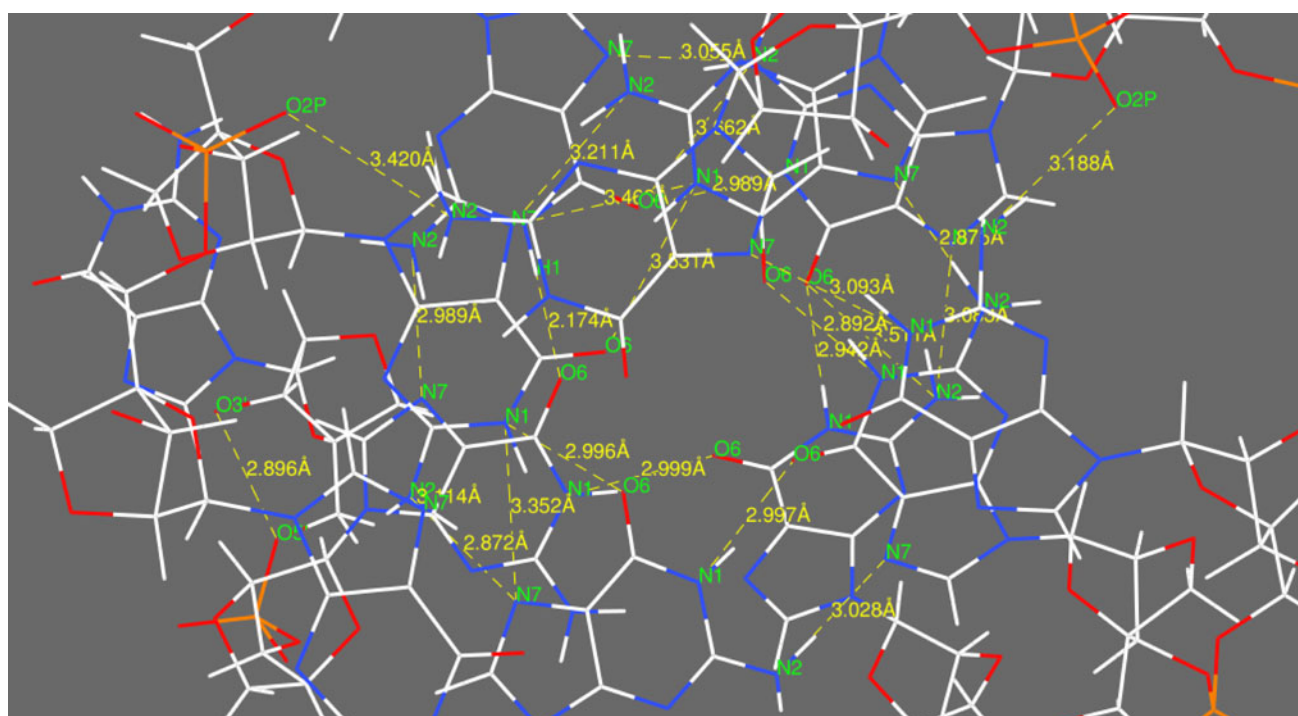
S. no.	Res. no.	Atom1	Res. no.	Atom2	Distance(Å)
1	LCG-9	O6	LCG-3	N1	2.942
2	LCG-3	N2	LCG-9	O6	3.511
3	LCG-3	N2	LCG-9	N7	3.083
4	LCG-9	N1	LCG-21	O6	2.989
5	LCG-21	O6	LCG-9	N2	3.662
6	LCG-9	N2	LCG-21	N7	3.055
7	LCG-16	O6	LCG-21	H1	2.174
8	LCG-16	N7	LCG-21	N2	2.989
9	LCG-16	N1	LCG-3	O6	2.999
10	LCG-17	O3'	LCG-17	O6	2.896
11	LCG-22	N1	LCG-15	N7	3.352
12	LCG-15	N7	LCG-22	N2	2.872
13	LCG-22	N2	LCG-23	N7	3.114
14	LCG-22	N7	LCG-10	N7	3.211
15	LCG-22	N7	LCG-10	N1	3.463
16	LCG-10	N1	LCG-22	O6	3.531
17	LCG-22	N1	LCG-15	O6	2.996
18	LCG-15	N1	LCG-4	O6	2.997
19	LCG-10	O6	LCG-4	N1	2.892
20	LCG-10	N7	LCG-4	N2	2.875
21	LCG-5	N1	LCG-11	N7	3.093
22	LCG-11	O2P	LCG-5	N2	3.188
23	LCG-11	N2	LCG-23	O2P	3.420
24	LCG-4	N7	LCG-15	N2	3.028

binding free energy ( $-4.87$  kcal/mol). MOD4 is the second most stable structure with respect to other models, which is also a mixed hybrid type structure, where, first and fourth strands are parallel and second and third strand are anti-parallel. The binding energy for this model is  $-44.66$  kcal/mol and binding free energy is  $2.75$  kcal/mol, which gives second lowest value. Among all structures, MOD2, which is also a mixed hybrid type structure where first, second and fourth strand are in parallel while third strand is in antiparallel, has lowest stability because its binding energy is  $34.16$  kcal/mol and also binding free energy for this model is most unfavorable (positive) relative to other models.

Consequently, the binding free energy calculation suggests that MOD3 exhibits higher stability than the other modified structures.

### Hydrogen bonding patterns

The starting fiber model [57], as well as high resolution crystal structure [58] of parallel quadruplex with coordinated cations, contains N1–H1–O6 and N2–H2–N7 (Hoo-gsteen type) hydrogen bonds between adjacent guanines, leading to eight hydrogen bonds per G-tetrad. Ab initio study on G-tetrad [56] also suggest that, in the absence of a coordinated cation, G-tetrad is stabilized by bifurcated hydrogen bonds between N1–H1–N7 and N2–H2–O6 atoms, whereas MD simulation of the 4-mer parallel quadruplex structure without any coordinated cations [52]

**Fig. 7** Hydrogen bonds and corresponding distances between the atoms of G-tetrads in MOD2

reported disruption of the G-tetrad geometry due to strand slippage. It is observed that during the molecular dynamics simulation, hydrogen bonding scheme within G-tetrad depends on the electrostatic interaction between the polar atoms of guanine bases and the coordinating molecules. The inter atomic distances between potential hydrogen bond forming groups within a G-tetrad and distances between different O6 atoms in the MD structures obtained during 20 ns dynamic run, which are listed in Table 4 and Table S2. During the simulation, hydrogen bonds in all G-tetrads (Fig. 7 and S3) are retained for all seven structures. However, due to the strong attractive force between coordinated K<sup>+</sup> ion and O6 atoms, guanine bases in some G-tetrads undergo in plane rotational motion. Because of this rotational motion, N<sub>1</sub>–H<sub>1</sub>–O<sub>6</sub> and in some cases N<sub>2</sub>–H<sub>2</sub>–N<sub>7</sub> hydrogen bonds are elongated. Whereas N<sub>1</sub> atom and N<sub>2</sub> atom of guanine base comes closer to N<sub>7</sub> and O<sub>6</sub> atom of neighboring guanine base and forms a N<sub>1</sub>–H<sub>1</sub>–N<sub>7</sub> and N<sub>2</sub>–H<sub>2</sub>–O<sub>6</sub> hydrogen bond. Thus, there centered hydrogen bond play additional role to stabilized these tetrads. As shown in Table 4, G-tetrad of MOD2 forms 24 hydrogen bonds, which is the maximum number of bond with association to all other models. The bonding distance and number of hydrogen bonds for MOD1 and MOD3-7 are listed in Table S2. Due to maximum number of H-bonding between tetrads, MOD2 structure shows more stability.

## Conclusion

It is well known that LNA modifications stabilize duplex and triplex DNA. In previous report a fully LNA modified quadruplex d(TG3T) was stabilized relative to the DNA quadruplex (37). This study explains the relative stability of modified d(TAGGGT)<sub>4</sub> G-quadruplex configurations. MD simulations are performed for seven different modified quadruplex structures. Among all modified structures, one of the mixed hybrid type structures, with two parallel and two antiparallel strands, shows lowest free energy. The energy calculations suggest that this mixed hybrid type conformation is the most stable among other conformations. RMS deviation of MOD2 complex, which is also a mixed hybrid type structure, is smaller. RMS deviation values of other complexes are very high and vary during the simulation. This study suggest that the modified monomeric DNA sequence d(TAGGGT)<sub>4</sub> can form only mixed hybrid type of structure. These modified sequences may help to inhibit the growth of the tumor and cancerous cell. The tendency of human telomeric G-quadruplex structure to adopt different conformations under varying environmental conditions which poses significant challenges in the design of specific drugs that could efficiently

bind and stabilize the structure. Our study provides in-depth insight into the conformational aspects of the quadruplex for subsequent evaluation of its respective interactions with drugs.

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