Quantitative Formation of Monomeric G-Quadruplex DNA from Multimeric Structures of c-Myc Promoter Sequence

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G-Quadruplex (G4)-forming DNA sequences have a tendency to form stable multimeric structures. This can be problematic for studies with synthetic oligodeoxynucleotides. Herein, we describe a method that quantitatively converts multimeric intermolecular structures of the Pu27 sequence from the c-myc promoter into the desired monomeric G4 by alkaline treatment and refolding.

Guanine-rich DNA sequences are able to adopt three-dimensional structures called G-quadruplex (G4) DNA. Four guanines can form a G-quartet through Hoogsteen base pairing while the O6 oxygens are coordinated by cations like K⁺, NH₄⁺ or Na⁺. At least two of these square planar G-quartets are involved in the formation of a G4 (Scheme 1).[1]

G4-forming sequences are widespread in the human genome.[2] They are enriched in promoter regions of many genes including oncogenes like c-myc,[3] c-kit,[4] VEGF[5] and bcl-2[6] which is of great interest for therapeutic applications like anticancer drug design. The protooncogene c-myc encodes a protein that is a central regulator of cell growth, differentiation, proliferation and apoptosis.[7] Overexpression of c-myc is associated with various cancers like colon, breast, cervix and small cell lung carcinomas.[8] Expression of c-myc is mainly controlled by the nuclear hypersensitivity element III1 (NHE III1), that is located −142 to −115 base pairs upstream the P1 promoter.[9] The template strand of NHE III1 contains the G-rich sequence Pu27 that is able to form a G4. There is an equilibrium between transcriptionally active forms and G4 structures which function as transcriptional repressor element.[10,11] When G4s are formed in vitro a variety of different intra- and intermolecular structures can be obtained.[12] G-rich oligodeoxynucleotides (ODN) can not only form monomeric G4 but also bi-, tri- or tetramolecular G4 and other higher order structures like G-wire.[13] This can be a major problem when studies with defined monomeric G4 are intended. A common method to obtain defined structures is to modify the sequence by substitution or deletion of guanine nucleotides.[14] However, this approach can be problematic because the modified G4 might lose its biological relevance. For the native Pu27 sequence for example, a defined structure can be obtained by deleting five G residues and replacing two G residues with T.[15] The resulting MYC22-G14T/G23T sequence is quite different compared to the native one.

Trent and co-workers investigated the influence of strand concentration, annealing process and buffer composition on the distribution of monomer and higher-order structures of Pu27 after thermal denaturation at 100 °C.[16] They found that the highest monomer yield is obtained at low K⁺ and low strand concentrations. However, even under optimized conditions a significant fraction of the G4 exists in a multimeric form and there is no method described that exclusively forms the biologically relevant monomeric G4 of Pu27.

Because it is also possible to denature duplex DNA[17] as well as G-rich ODN[18] under basic conditions, we were interested to investigate alkaline denaturation for transforming Pu27 ODN into monomeric G4. G-rich ODN were treated with NaOH followed by neutralization with buffer containing K⁺ ions. For comparison, we performed thermal treatment which is commonly used for unfolding and refolding G4. The efficiency of monomer formation of chemically and thermally treated as well as untreated ODN was analyzed by size-exclusion chromatography (SEC), which is well suited to investigate the oligomerization state of G4.[19a,17] The chromatogram of Pu27 after DNA synthesis with RP-HPLC purification shows that the structure of Pu27 is very polymorphic without treatment (Figure 1). A small fraction elutes as monomer with a retention time of around 8 min but most of the ODN elutes earlier, which can be attributed to multimeric structures with higher molecular weight. After thermal treatment at 95 °C for 10 min and cooling to room temperature, the amount of monomer increases but a significant fraction of the ODN is still in its multimeric forms.

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Scheme 1. Structure of a G-quartet with hydrogen bonds between N1 and O6 highlighted in red and transformation of various multimeric structures into monomeric G4 DNA.

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This demonstrates that thermal denaturation and renaturation by cooling to room temperature is not sufficient to quantitatively produce Pu27 monomers. Only with the chemical treatment all multimeric structures are disrupted and the monomer forms exclusively.

For the chemical treatment ODN (250 μM) was incubated in an NaOH solution (150 mM) for 5 min at room temperature and then diluted with 50 times the volume of K⁺-containing buffer (25 mM KH₂PO₄, pH 7.0) to neutralize the solution and facilitate monomer formation. Lower NaOH concentrations, even at elevated temperature, did not yield pure monomer (Figures S1 and S2). The disruption of multimeric structures with 150 mM NaOH can be explained by removal of protons from N1 of guanine residues, which leads to loss of essential hydrogen bonds for G-quartet formation (Scheme 1). Loss of N1 protons in Pu27 is supported by a change of the UV spectrum upon base treatment which resembles the UV change observed for 2'-deoxyguanosine (Figure S3).

We also tested the different treatment methods for derivatives of Pu27 (Table 1). In the sequence of Pu22 the first G-run, which is not involved in monomer G4 formation of Pu27, is deleted in order to reduce the polymorphism. The second and fourth G-run contain four G residues as in the native Pu27. In the Myc22 sequence the first G-run related to Pu27 is deleted, just as in Pu22, and in addition the second and fourth G-run have three instead of four G in order to reduce polymorphism even more. In the SEC analysis of Pu22 (Figure 2A) and Myc22 (Figure 2B) it becomes obvious that even these sequences form multimeric structures if not treated (black curves). The more the sequence is modified, the more monomers can be observed. Thermal treatment results in less multimers for Pu22 and only monomer is observed for the more modified Myc22 sequence (green curves). After chemical treatment all higher-order structures also disappear for Pu22 (red curve). The results show that sequence modifications of native Pu27 have a positive influence in reducing the formation of multimeric structures, but if only monomer of Pu27 or slightly modified Pu22 is desired, alkaline treatment is necessary.

We also examined the treatment methods for the three sequences using polyacrylamide gel electrophoresis (PAGE; Figure 3). On the NuPAGE gel the untreated strands of Pu27 and Pu22 display a fast migrating band and many slower migrating bands that correspond to multimeric structures. After thermal treatment the multimeric structures are still present but the relative amounts of monomer increase. Only denaturation under basic conditions and neutralization with K⁺-containing buffer converts all multimeric structures into monomers. For Myc22 very low amounts of multimers can be detected without

![Figure 1](image1.png)

**Figure 1.** Size-exclusion chromatography (SEC) of native c-myc ODN Pu27 without treatment (black), after thermal treatment at 95 °C for 10 min and cooling to room temperature (green) as well as after chemical denaturation with 150 mM NaOH and neutralization with buffer containing K⁺ (red). Very large macromolecules, e.g., bovine thyroglobulin (670 kDa), elute around 5 min.

![Figure 2](image2.png)

**Figure 2.** SEC of c-myc-derived sequences A) Pu22 and B) Myc22 without treatment (black), after thermal treatment (green) and after chemical treatment (red).

![Figure 3](image3.png)

**Figure 3.** NuPAGE of Pu27, Pu22 and Myc22 without treatment, after thermal treatment and after chemical treatment.
or with thermal treatment and chemical treatment results in pure monomer.

The circular dichroism (CD) spectra of Pu27 before and after the different treatment methods (Figure 4) have a maximum around 260 nm and a minimum around 240 nm what is expected for Pu27 which forms a parallel propeller-type G4.\textsuperscript{[3,4]} These CD spectra show no significant difference although the SEC and NuPAGE analysis demonstrate that there are many different structures. This indicates that CD spectroscopy of Pu27 is not suitable to distinguish between the monomeric and multimeric structures. The CD spectrum of the ODN that was chemically treated also displays the signature of a parallel G4 which demonstrates that the monomer species is not a random coil but a folded G4. It can also be concluded that the multimeric structures have a parallel orientation and even without treatment Pu27 forms G4 structures.

G4 DNA has another interesting spectroscopic property namely that the absorbance at 295 nm decreases upon melting. We, therefore, investigated the melting behavior of Pu27 and compared the untreated, thermally and chemically treated ODN. The melting temperatures indicated by the inflection points of the melting curves are very similar (Figure 5).

However, a major difference is observed in the amplitude of absorbance change that is highest for the chemically treated and smallest for the untreated ODN. This indicates that the change in absorbance is caused by the melting of the monomeric G4 and that most of the multimeric structures melt above 90 °C. The untreated and thermal treated samples contain a much smaller amount of monomeric G4 and for that reason the change in absorbance is smaller upon heating. The observation that a large fraction of multimeric structures of Pu27 do not melt at 90 °C gives a plausible explanation why the thermal treatment fails to form G4 monomers quantitatively. For the c-myc-derived sequences Pu22 and Myc22, which contain less or almost no multimeric structures to begin with, the amplitudes of absorbance change are very similar or almost identical for the samples without, with thermal or with chemical treatment (Figures S4 and S5).

In this work we demonstrated that multimeric G4 DNA structures can be completely converted into monomeric G4 under convenient conditions. By treatment of Pu27 with 150 mM NaOH and neutralization with potassium ions monomeric G4 is formed exclusively. We have shown by SEC and PAGE that thermal treatment is not sufficient to disrupt all multimeric structures of Pu27 and its derivative Pu22. In addition, our results demonstrate that the presence of multimeric structures can easily be overlooked when only CD spectra and UV melting points are analyzed. The possibility of having multimeric structures in G4 preparations is rarely discussed in the literature. The presence of higher-order structures can be problematic for studies with G4 DNA where correct concentrations are needed to obtain binding constants for ligand or protein interactions. In addition, the multimeric structures might behave different than the monomers which could lead to false structure-function correlations. These problems can be easily overcome by alkaline denaturation followed by refolding in the presence of potassium ions.

**Experimental Section**

Oligodeoxynucleotides (ODN) were purchased RP-HPLC-purified and lyophilized from Integrated DNA Technologies (IDT; for sequences see Table 1) and dissolved in water. Water was purified using a Milli-Q purification system and passed through a 0.22 μm filter. Reagents were purchased from Acros Organics, Gerbu, Merck, Serva and Sigma-Aldrich with analytical grade and used without further purification. The following buffers were prepared: KPP buffer: 25 mM KH2PO4 (pH 7.0); SEC buffer: 25 mM KH2PO4, 1 mM NaNO3 (pH 7.0); NuPAGE gel buffer: 357 mM Bis–Tris (pH 6.5–6.8, NuPAGE running buffer: 50 mM MOPS, 50 mM Tris–X, 5 mM EDTA, 0.1 % SDS (pH 7.6); SDS loading buffer (5x): 250 mM Tris–HCl, 12.5 % SDS, 50 % glycerol, 0.05 % Bromphenol Blue.

**Chemical denaturation and refolding of G4 ODN:** An aqueous ODN solution (5 nmol, 5 μL) was supplemented with a solution of NaOH (15 μL, 200 mM) to obtain a final concentration of 150 mM.

**Table 1. Sequences of G4-forming oligodeoxynucleotides (ODN).**

| ODN   | Sequence (from 5' to 3' direction)            |
|-------|-----------------------------------------------|
| Pu27  | TGGGGAGGGTTGGGAGGGGTTGGGGAAGG                |
| Pu22  | TGGGAGGGTTGGGAGGGGTTGGGAA                   |
| Myc22 | TGGGGAGGGTTGGGAGGGGTTGGGTGTGAGTG            |
Afterwards KPP buffer (980 μL) was added to the reaction mixture, and the solution was incubated at room temperature for 5 min. The sample was then heated to 90°C, and the melting curves were measured at a temperature range from 35°C to 90°C. The data interval was 0.1°C with a heating rate of 0.3°C/min. The first derivation of the obtained curves was calculated and the melting temperature obtained by averaging the three lowest values.

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**Conflict of Interest**

The authors declare no conflict of interest.