Identification of G-quadruplexes in functional RNAs using FOLDeR

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RNA G-quadruplex (G4) structures are thought to affect biological processes, including translation and pre-mRNA splicing, but it is not possible at present to demonstrate that they form naturally at specific sequences within long functional RNA molecules. We have developed a novel strategy, footprinting of long 7-deaza-guanine substituted RNAs (FOLDeR), that allows the formation of G4s to be confirmed in long RNAs and in functional conditions.
G-rich sequences in DNA or RNA have the potential to form four-stranded structures known as G-quadruplexes (G4). DNA G4 formation has been demonstrated in vivo and implicated in a very wide range of gene expression processes, notably telomere maintenance, transcription and genome instability. RNA G4s exist in vivo and have been implicated in various post-transcriptional processes. They would be expected to form more readily in RNA than in DNA, but it is still difficult to predict whether they do form in any given RNA sequence.

G4s have been suggested previously to be of functional significance in pre-mRNA splicing. Most of these studies used similar approaches, using bioinformatic tools to identify well-characterized G4-forming sequences and biophysical methods to show that the predicted G4 can form in isolated short sections of RNA. However, such sequences are naturally part of longer pre-mRNAs in which the propensity of a sequence to form a G4 is determined by competition with secondary structures and protein binding. Other methods for identifying G4s include the use of ligands or antibodies that bind selectively to G4s, but a general hazard is that the binding reagent affects the equilibrium between different structures and thus might induce the formation of otherwise unstable non-functional G4s. Mutagenesis is often used for confirmation of G4 formation in vivo, but mutants need to be designed so as not to affect the binding of regulatory proteins or perturb likely secondary structures.

We describe here a method in which we probed the secondary structure of an entire functional unit of pre-mRNA, comparing native RNA with 7-deazaguanine (7-deaza-G) substituted RNA, which can form the same secondary structures but not G4s. Differences, highlighted regions in which G4 might form. These differences were
confirmed in functional splicing conditions by ribonuclease H digestion in nuclear extracts. This method allowed the identification of functionally relevant G4s.

The splicing reaction we have studied is that of the human B-cell lymphoma-extra large (Bcl-x) pre-mRNA, which has two alternative 5’ splice sites (5’SS) in its exon 2. The major isoform, Bcl-xL (XL), is an anti-apoptotic factor, while the alternative isoform, Bcl-xS (XS), is pro-apoptotic17. There are a number of guanine-rich (G-rich) sequences in the regions around the Bcl-x alternative splice sites that could form G4s.

We designed a functional splicing transcript, Bcl-x-681, by preserving the sequence between the alternative XS and XL 5’SS, shortening the intron and the exon 3 and adding an additional 5’SS at the 3’end of the construct (Fig. 1a). Splicing of this transcript substantially favored the Bcl-xL isoform, recapitulating the preference for Bcl-xL observed in HeLa cells18 (Fig. 1b, Supplementary Fig. 1).

The presence of G4s in this functional RNA was assessed by an electrophoretic mobility shift assay (EMSA) using a previously described G4-specific antibody, BG42,4, that induced a shift in the migration of the Bcl-x-681 RNA (Fig. 1c, left). This transcript contains 6 G-rich sequences predicted to potentially form a G419, one upstream of the XS splice site (Q1), three between the XS and the XL splice sites (Q2, Q3 and Q4), and one downstream of the 3’ splice site (3’SS) (Q6). All of these short sequences displayed a circular dichroism spectrum that is typical for parallel G4s, with a positive signal at 265 nm and a negative signal at 240 nm (Supplementary Fig. 2). The stability of these G4s ranged from 37°C (Q4) to >70°C (Q2). However, these
experiments did not provide information on whether these sequences do form in the face of competing secondary structures in functional Bcl-x pre-mRNA.

Therefore, our strategy for identifying the location of G4 elements in this RNA was to (i) map structured regions using ribonucleases (RNases), and then (ii) repeat the assay with 7-deaza-G-RNA, in which G4s would not form\textsuperscript{16}. For (i), we performed RNA footprinting using RNases T1, T2 and V1, providing structural restraints at nucleotide resolution (Supplementary Figs. 3 and 4). In total, we defined 163 and 91 nucleotides as single-stranded and double-stranded, respectively (Supplementary Fig. 4, Supplementary Dataset 1). We used these restraints with a secondary structure prediction software (Mfold\textsuperscript{20}) to derive the best fit model (Supplementary Fig. 6). This model predicted that the Bcl-x-681 RNA is highly structured and suggested the presence of four structurally independent domains, denoted as X\textsubscript{s}, X\textsubscript{l}, intron and 3’ss domains according to the splicing elements they contained (Supplementary Fig. 6).

We transcribed each structural domain separately and RNA footprints of each domain showed a very similar footprinting pattern to that of the full-length RNA (Supplementary Fig. 7), supporting the structural model.

To identify potential G4s within this structure, we replaced all guanines by 7-deaza-G during \textit{in vitro} transcription. The absence of the N7 in the 7-deaza-G was shown previously to abolish the possibility of Hoogsteen base pairing and thus of G4 formation, whilst still permitting Watson-Crick base-pairing and the formation of stem-loops\textsuperscript{16}. We confirmed the absence of G4s in the 7-deaza-G Bcl-x-681 RNA by EMSA. As expected, BG4\textsuperscript{2-4} did not induce a shift in the migration of the 7-deaza-G RNA (Fig. 1c, right).
We performed RNA footprinting on the 7-deaza-G RNA (Supplementary Fig. 4) and a comparison between the RNA footprinting of the 7-deaza-G-substituted transcripts and the native transcript was assessed by calculating an index factor corresponding to 

\[(\text{V1 7-deaza-G} \div \text{V1 native}) / (\text{T2 7-deaza-G} \div \text{T2 native})\] (Supplementary Dataset 2, left). The majority of the 40 nucleotides having a index value below 0.3 or above 3, indicating a three-fold difference in structural probing between the native and the 7-deaza-G RNA) were located in the XS and the XL domains, and many of these nucleotides coincided with two of the possible G4 regions (Q2, and Q5) (Fig. 1d).

We therefore concluded that the Bcl-x-681 transcript contains G4s located near the XS and XL splice sites. Previous reports identified G4s by comparing structural probing of RNAs in the presence of either potassium (K⁺) or Lithium (Li⁺) ions⁸, although on much shorter sequences than Bcl-x-681. For comparison, we performed an RNA footprinting experiment of Bcl-x-681 in the presence of Li⁺ or K⁺ ions (Supplementary Dataset 3) and calculated an index as 

\[(\text{V1 K⁺} \div \text{V1 Li⁺}) / (\text{T2 K⁺} \div \text{T2 Li⁺})\]. Only 5 nucleotides displayed an index below 0.3 or above 3. In order to compare this strategy with our strategy using deaza-G-substituted RNA, we therefore considered the 40 nucleotides that displayed an index below 0.5 or above 2 (Supplementary Fig. 89). The majorities of these nucleotides were located in the XS and XL domains and coincided mainly with Q2. These results validated the use of our 7-deaza-GTP substitution strategy for mapping G4s in long RNAs and, importantly, showed that the use of 7-deaza-G substitution is more sensitive than the comparison of structural probing in K⁺ and Li⁺ containing buffers.
A major benefit of using 7-deaza-G RNA in structural probing is that, in contrast to the comparison between Li\(^+\) and K\(^+\) buffers, it can be done in functional conditions, such as in nuclear extracts. Therefore, we tested the accessibility of native and 7-deaza-G modified Bel-x-681 RNAs using RNase H cleavage directed by 10-mer DNA oligonucleotides in functional splicing conditions in nuclear extracts (Supplementary Fig. 109). We focused our analysis on the X\(_S\) and X\(_L\) domains. We assigned protected or accessible regions on the basis of either \(< 40\%\) or \(> 60\%\) cleavage, respectively (Supplementary Fig. 109). The RNase H cleavage patterns of native RNA in nuclear extract were in good agreement with the structure predicted in solution (Fig. 2a). Putative stems were protected in both the X\(_S\) and X\(_L\) domains, while several of the predicted loops were accessible. Comparison of the RNase H patterns of the native and the 7-deaza-G RNA (Fig. 2b) showed major differences mainly near the putative G4s, especially Q2 and Q5, consistent with the footprinting results. These data demonstrated that the use of 7-deaza-G RNA can identify G4s in functional splicing conditions.

In conclusion, we have described FOLDeR, a new method that allowed us to demonstrate the presence of G4s within long RNAs by using 7-deaza-G RNA modifications in combination with secondary structure probing (Fig. 1). By coupling footprinting with ribonuclease H cleavage assays on the two different types of transcript, it was also possible to validate the findings in functional conditions, such as nuclear extracts (Fig. 2). Although we demonstrated the power of this method using nuclease digestions, its validity relies on the modification of the RNA and not on the structural probing method. Therefore, the use of 7-deaza-G RNAs to probe
G4s could also be done in combination with other structural probing methods such as selective 2′-hydroxyl acylation analyzed by primer extension (SHAPE) chemistry or dimethyl sulphate (DMS) footprinting. It is now becoming clear that RNA G4s have a higher propensity to form than their DNA counterparts. Recently, structural data revealed the presence of RNA G4s that could not be predicted by bioinformatics tools. The development of an experimental method for identifying and locating G4s is therefore a major step forward. The existence of G4s in a functional RNA can be tested using the BG4 antibody and their location established using 7-deaza-G-RNA as described here.

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Author Contributions
C.W. performed all experiments under the guidance of I.C.E. and C.D. Footprinting experiments and structural model calculation were performed under the guidance of I.B.A. and C.B. G.A.B. and L.H.H. contributed to the development and the validation of the strategy. C.W., I.C.E., and C.D. interpreted the results and wrote the manuscript.
The authors declare no competing financial interests
References


Figure Legends:

**Figure 1: The functional Bcl-x-681 RNA contains G4s in vitro**

(a) Schematic representation of native Bcl-x (top) and Bcl-x-681 transcript (bottom) used in this study. The size of introns and exons are indicated in nucleotides below the diagrams. X_L and X_S 5’ splice sites are indicated above the diagrams. (b) In vitro splicing assay of Bcl-x-681. Bands corresponding to the unspliced transcript, the X_L and the X_S products are labelled. The full-length gel is displayed in Supplementary Fig. 1 (c) Electrophoretic-mobility shift assay of native (left) and 7-deaza-G substituted (right) Bcl-x-681 in the absence or presence of increasing amount of BG4 antibody (0, 150 pM, 1.5 nM, 15 nM, 150 nM and 1.5 μM). The full-length gel is displayed in Supplementary Fig. 10. (d) Mapping of RNA footprinting of Bcl-x-681. Nucleotides having a significantly different footprinting pattern upon 7-deaza-GTP substitution are circled and are located mainly to the X_S and X_L domains. Putative G4s are labelled Q1 to Q6.

**Figure 2: The Bcl-x-681 RNA contains G4s in functional conditions**

(a) Schematic drawing of the RNAse H cleavage of native Bcl-x-681 on the secondary structure model of Bcl-x-681. Accessible regions (average >60% cleavage) are in black, and protected regions (average <40% cleavage) are in grey. RNAse cleavage experiments were done in triplicate and are presented in supplementary Figure 9A. (b) Percentage of change of RNAse H cleavages in 7-deaza-G versus native RNA. Positive changes indicate increase in accessibility while negative changes indicate increase in protection. Data represent the difference between the average of three RNAse H cleavage experiments on native RNA (supplementary
Figure 9A) and the average of three RNase H cleavage experiments on 7-deaza-G RNA (supplementary Figure 9B).
Online Methods:

**In Vitro Transcription of RNA**

A transcription reaction containing 40 mM Tris (pH 7.5), 20 mM MgCl₂, 10 mM NaCl, 2 mM spermidine HCl, 10 mM DTT, 4 mM NTPs, 1 μg DNA template (PCR product), 5% RNaseOut (Invitrogen), and 5% T7 polymerase (1:20 dilution) was incubated at 37 °C for 4 h. DNA fragments were removed using an S-300 column (GE Healthcare). The product was extracted by phenol-chloroform and precipitated by ethanol. The pellet was dissolved in the desired amount of TE buffer and stored at −20 °C. For radiolabelled RNA, a transcription reaction containing 40 mM Tris (pH 7.5), 6 mM MgCl₂, 10 mM NaCl, 2 mM spermidine HCl, 0.5 mM ATP, 0.5 mM CTP, 0.5 mM UTP, 0.05 mM GTP, 50 ng DNA template (PCR product), 4% RNaseOut, 10% T7 (1:20 dilution), 1 mM Ribo m7G Cap analogue (Promega) and 0.33 μM [α-32P] GTP (10 mCi/ml, 3000 Ci/mmol) (Pelkin Elmer) was incubated at 37 °C for 1.5 h. Samples were then purified via a denaturing gel.

**In vitro transcription of 7-deaza RNA**

A transcription reaction containing 40 mM Tris (pH 7.5), 20 mM MgCl₂, 10 mM NaCl, 2 mM spermidine HCl, 10 mM DTT, 4 mM NTPs (c7GTP instead of GTP), 1 μg DNA template (PCR product), 5% RNaseOut, 2 mM Ribo m7G Cap analogue (Promega) and 5% T7 (1:20 dilution) was incubated at 37 °C for 4 h. DNA fragments were removed using an S-300 column (GE Healthcare). The product was extracted by phenol-chloroform and precipitated by ethanol. The pellet was dissolved in the desired amount of TE buffer and stored at −20 °C. For radiolabelled deaza RNA, a transcription reaction containing 40 mM Tris (pH 7.5), 6 mM MgCl₂, 10 mM NaCl,
2 mM spermidine HCl, 0.5 mM ATP, 0.5 mM CTP, 0.5 mM UTP, 0.05 mM GTP, 50 ng DNA template (PCR product), 4% RNaseOut, 10% T7 (1:20 dilution), 1 mM Ribo m7G Cap analogue (Promega) and 0.33 μM [α-32P] GTP (10 mCi/ml, 3000 Ci/mmol) (Pelkin Elmer) was incubated at 37 °C for 1.5 h. Samples were then purified via a denaturing gel.

**In Vitro Splicing of RNA**

Samples containing 1.53 mM ATP, 20 mM CrPi, 3.2 mM MgCl₂, 20 mM Hepes (pH 7.5), 50 mM KGlu, 0.125% NP-40, 50% nuclear extract (Cilbiotech) and 10–20 cps of radiolabelled RNA transcript were set up in 10 μl reactions on ice then incubated at 30 °C for 2 h in a pre-heated waterbath. Aliquots of 2 μl were taken into a microtitre plate on dry ice at timepoints 0, 30, 60 and 120 min. Samples were then treated by addition of 50 μl of 0.4 mg/ml proteinase K and incubation for 10 mins at 37 °C. Samples were precipitated twice with 100% ethanol, and dissolved in 10 μl formamide dyes. To prepare for loading on an 8% denaturing polyacrylamide gel, samples were placed in boiling water for 30 sec and run until bromophenol blue has just run out of the gel. Gels were then fixed, dried and exposed to a phosphorimaging screen overnight. Quantification of pre-mRNA and both mRNA products was done using OptiQuant software, and intensities were normalised to account for the number of guanines in each RNA species.

**RNase probing**

Samples containing 400 ng RNA (either native or 7-deaza-substituted), 0.5 mg/ml yeast tRNA, and either 150 mM KCl, 1.5 mM MgCl₂ and 20 mM Hepes pH 7.6 (K⁺ containing buffer) or 150 mM KCl, 1.5 mM MgCl₂ and 20 mM Hepes pH 7.6 (Li⁺ containing buffer).
containing buffer) in a total volume of 9 µl, were heated at 65 °C in a water bath for 10 min then slow cooled to 37 °C. 1 µl of either Tris buffer (0.5 mM Tris pH 7.6, 1 M KCl or 1M LiCl, and 25 mM MgCl₂), RNase T1 (Roche Diagnostics), RNase T2 (Ambion) or RNase V1 (Mobitech) was added. Reactions were incubated at ambient temperature for 6 min then stopped with 101.8 µl of stop buffer (3.9 mM EDTA and 0.2 µg yeast tRNA) and 101.8 µl phenol-chloroform then spun at 13,500 rpm for 10 min. The top layer was removed and added to 700 µl ethanol (96%), 12 ng glycogen and 22 mM NaAc (pH 3.0). Samples were then ethanol-precipitated and dissolved in 4 µl potassium borate or sterile water.

**Reverse transcription of probed RNA**

A hybridization mix was made containing 0.25 µl hybridization buffer (reverse transcription buffer without MgCl₂), 1 µl of 100 cps/µL radiolabelled primer and 0.25 µl water. 1 µl of probed RNA was incubated with 1.5 µl of hybridization mix at 65 °C for 10 min then cooled on ice. 2.5 µl of extension mix (0.1 µl 5 mM dNTPs, 0.25 µl reverse transcriptase buffer, 0.25 µl 1/10 diluted AMV reverse transcriptase (MB Biochemical) and 1.9 µl water) was added to each probed RNA mix and incubated at 42 °C for 30 min. 3 µl formamide dye was added and the sample was heated to 96 °C for 2 min, and then put on ice for 2 min. 2 µl was loaded onto 7% DNA sequencing gel and run for 2 h at 100 W. Gels were dried and exposed to a phosphorimaging screen overnight and quantified using SAFA.
**RNase H assays**

RNase H cleavage was done in triplicate in splicing conditions by adding DNA oligonucleotide at 10 µM after incubation for 30 min and continuing incubation for 5 min. Reactions were processed as for splicing reactions.

**Electromobility Shift assays (EMSA)**

The BG4 antibody was purchased from Absolute Antibody (Ab00174-1.1). Radiolabelled RNA at 10 count per seconds was incubated with various concentrations (0, 150 pM, 1.5 nM, 15 nM, 150 nM and 1.5 μM) of BG4 antibody in a microtitre plate. 3 µl of native gel dye was added, and 4 µl was run on a small 5% native polyacrylamide gel at 4°C. The gel was run at 5 W for 1 h. Gels were dried and exposed to a phosphorimager screen overnight.

**Circular Dichroism**

RNA sequences corresponding to Q1, Q2, Q3, Q4, Q5, and Q6 were purchased from Dharmacon, GE Healthcare, deprotected according to the manufacturer instructions, lyophilised, and resuspended in 20mM KH₂PO₄, pH7, 100mM KCl at a final concentration of 20μM. Circular dichroism spectra were acquired on a Chirascan spectrometer at 20°C and data were collected between 350 and 220nm in triplicate and averaged. Buffer baseline was subtracted for each spectrum. CD Melting curves were obtained by measuring the CD signal at 265nm from 20 to 80°C at 1°C heating rate.