NMR solution structure of the major G-quadruplex structure formed in the human BCL2 promoter region

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ABSTRACT

BCL2 protein functions as an inhibitor of cell apoptosis and has been found to be aberrantly expressed in a wide range of human diseases. A highly GC-rich region upstream of the P1 promoter plays an important role in the transcriptional regulation of BCL2. Here we report the NMR solution structure of the major intramolecular G-quadruplex formed on the G-rich strand of this region in K⁺ solution. This well-defined mixed parallel/antiparallel-stranded G-quadruplex structure contains three G-tetrads of mixed G-arrangements, which are connected with two lateral loops and one side loop, and four grooves of different widths. The three loops interact with the core G-tetrads in a specific way that defines and stabilizes the overall G-quadruplex structure. The loop conformations are in accord with the experimental mutation and footprinting data. The first 3-nt loop adopts a lateral loop conformation and appears to determine the overall folding of the BCL2 G-quadruplex. The third 1-nt double-chain-reversal loop defines another example of a stable parallel-stranded structural motif using the G₃NG₃ sequence. Significantly, the distinct major BCL2 promoter G-quadruplex structure suggests that it can be specifically involved in gene modulation and can be an attractive target for pathway-specific drug design.

INTRODUCTION

The BCL2 (B-cell CLL/lymphoma 2) gene product is a mitochondrial membrane protein that plays an essential role in cell survival. The BCL2 protein exists in delicate balance with other apoptosis-related proteins and functions as an inhibitor of cell apoptosis (1,2). Deregulation of the BCL2 gene is associated with aberrant cell growth in many human diseases. For example, BCL2 has been found to be aberrantly overexpressed in a wide range of human tumors, including B-cell and T-cell lymphomas (3,4) and breast (5), prostate (6), cervical (7), colorectal (8), and non-small cell lung carcinomas (9). Moreover, BCL2 overexpression has also been associated with poor prognosis and has been found to interfere with traditional cancer therapeutics (10,11). Inhibition of BCL2 expression by small molecules (12,13), peptidomimetics (14), or antisense oligonucleotides (15,16) has been shown to reduce cellular proliferation and to enhance chemotherapeutic efficacy. In contrast, aberrantly reduced BCL2 expression is associated with cardiovascular diseases and neurological disorders, such as ischemia/reperfusion injury of cardiac and renal tissues (17,18), multiple sclerosis (19), Alzheimer’s and Parkinson’s diseases (20), and tissue damage related with stroke (21) and spinal cord injuries (22). Thus, BCL2 has also emerged as an attractive target for neuroprotective and tissue-protective therapies.

There is accumulating in vitro evidence of G-quadruplex structures formed in promoter regions of several genes as transcriptional regulators, such as the transcriptional repressors reported in the human MYC (c-MYC) gene (23,24), the chicken β-globulin gene (25) and several muscle-specific genes (26,27), as well as a transcriptional enhancer reported in the human insulin gene (28,29). The human BCL2 gene has two promoters, P1 and P2. The major promoter, P1, located 1386–1423 bp upstream of the translation start site, is a TATA-less, GC-rich promoter containing multiple transcriptional start sites and positioned within a nuclease hypersensitive site (30,31). A highly GC-rich 39-bp region located 58–19 bp upstream of the P1 promoter has been implicated in playing a major role in the regulation of BCL2 transcription (Figure 1A) (32). Multiple transcription factors have been reported to bind to or regulate BCL2 gene expression through this region, including CREB (33), WT1 (34), Sp1 (30), E2F (35), NF-xB (36), and NGF (37). For example, CREB functions to activate BCL2 expression (33), while WT1 functions to repress BCL2 expression (34).

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concentrations. Conditions: 25°C.

Figure 1. (A) The promoter sequence of the BCL2 gene and its modifications. Bcl2Pu39 is the wild-type BCL2 39mer sequence; bcl2MidG4 is the BCL2 23mer sequence containing the middle four consecutive G-runs, which forms the most stable G-quadruplex structure; bcl2Mid is the mutant 23mer with G-to-T mutations at positions 15 and 16, bcl2Midm2 is the mutant 23mer with G-to-T mutations at positions 15 and 19, and bcl2Midm3 is the mutant 23mer with G-to-T mutations at positions 18 and 19. The six G-runs are underlined and numbered using Roman numerals; the BCL2 23mer is the mutant 23mer with G-to-T mutations at positions 15 and 16, bcl2Midm3 is the most stable G-quadruplex structure; bcl2Mid is the mutant BCL2 23mer sequence containing the middle four consecutive G-runs, which forms the most stable G-quadruplex structure. (B) The imino regions of 1D 1H NMR spectra of bcl2Mid samples at 0.1 mM (upper) and 1.5 mM (lower) strand concentrations. Conditions: 25°C, 20 mM K-phosphate, 40 mM KCl, pH 7.0.

Our previous results have shown that the 39mer guanine-rich strand (bcl2Pu39, Figure 1A) in the BCL2 promoter can form a mixture of three distinct intramolecular G-quadruplexes in K+ containing solution, and that the G-quadruplex formed on the middle four consecutive runs of guanines is the most stable one (38,39), which adopts a novel folding of mixed parallel/antiparallel-stranded structure (39). In this paper we report the NMR solution structure for the predominant G-quadruplex structure formed in the BCL2 promoter region. The solution structure provides not only the molecular details of this G-quadruplex but also important insights for its loop conformations and interactions with the core tetrad structures.

MATERIALS AND METHODS

Sample preparation

The DNA oligonucleotides were synthesized using β-cyanoethylphosphoramidite solid-phase chemistry on an Expedite™ 8909 Nucleic Acid Synthesis System (Applied Biosystem, Inc.) in DMT-on mode, and were purified using C18 reverse-phase HPLC chromatography, as described previously (39–41). Deprotection was carried out using 80% AcOH for 1 h, followed by ether extraction. DMT-off DNA was further purified by HPLC followed by successive dialysis against 150 mM NaCl and H2O, 6% 1,2,7-15N, 2,13C-labeled guanine phosphoramidite (42) was used for site-specific labeled DNA synthesis. Samples in D2O were prepared by repeated lyophilization and final dissolution in 99.96% D2O. Samples in water were prepared in 10%/90% D2O/H2O solution. The final NMR samples contained 0.2–2 mM DNA oligonucleotides in 20 mM K-phosphate buffer (pH 7.0) and 40 mM KCl.

NMR experiments

NMR experiments were performed on a Bruker DRX-600 spectrometer as described before (39–41,43). At the determined melting point (61°C), stoichiometric titration of the melted and the folded strands as a function of total strand concentration from 0.01 to 0.1 mM was performed (43). Assignment of guanine imino and H8 protons were obtained by 1D 15N-filtered experiments using site-specific labeling. Standard 2D NMR experiments, including NOESY, TOCSY, and DQF-COSY, were used at 1, 7, 15, 20, 25, 30 and 35°C to obtain complete proton resonance assignment. The NMR experiments for samples in water solution were performed with Watergate or Jump-and-Return water suppression techniques. Relaxation delays were set to 3 s. The acquisition data points were set to 2048 × (350–512) (complex points). The 45° or 60° shifted sine-squared functions were applied to NOESY and TOCSY spectra. The fifth-order polynomial functions were employed for the baseline corrections. The final spectral sizes are 2048 × 1024. Peak assignments and integrations were achieved using the software Sparky (UCSF). Only relatively isolated peaks were used for NOE-restrained structure calculation. Severely overlapping peaks were discarded. The NOE peaks were integrated using the peak fitting function and volume integration of Sparky. We have manually checked every peak to make sure the fitted line shape agrees with the experimental data. The average linewidth of NMR peaks is 10–15 Hz, and the digitized resolution of the NOESY spectra is sufficient for the line-fitting and accurate volume integration in Sparky. Distances between non-exchangeable protons were estimated based on the NOE cross-peak volumes at 50–300 ms mixing times, with the upper and lower boundaries assigned to ±20% of the estimated distances. Distances between exchangeable protons were assigned with looser boundaries (±1–1.5 Å). The cytosine base proton H5–H6 distance (2.45 Å) was used as a reference. The distances involving the unresolved protons, e.g. methyl protons, were assigned using pseudo-atom notation to make use of the pseudo-atom correction automatically computed by X-PLOR.

The 31P NMR spectra were collected on a DNA sample at 1.5 mM in D2O (20 mM potassium-phosphate buffer, 40 mM KCl, pH 7.0) at 25°C and were referenced to an external standard of 85% H3PO4, including the 1D proton-decoupled phosphorus spectrum, and 2D heteronuclear 31P–1H Correlation Spectroscopy (COSY) and Heteronuclear Single Quantum
Correlation Spectroscopy (HSQC). A series of $^{31}$P–$^1$H HSQC spectra were collected at spectral widths of 10 p.p.m. ($^1$H) × 5 p.p.m. ($^{31}$P) with 2048 × 128 complex points, using various heteronuclear INEPT transfer delays corresponding to J-couplings of 5, 10, 15, 20 and 25 Hz. The non-selective and H3'-selective $^{31}$P–$^1$H COSY experiments (44) were carried out in States-TPPI mode using the same spectral width with 2048 × 128 complex points and 256 scans. Assignments of the individual $^{31}$P resonance were accomplished by a combination of 2D $^1$H/$^1$H NOESY, COSY, TOCSY and heteronuclear $^{31}$P–$^1$H COSY.

**Distance geometry and simulated annealing (DGSA) calculations**

Metric matrix distance geometry (MMDG) calculations were carried out using X-PLOR (45) to embed and optimize 100 initial structures. An arbitrary extended conformation was first generated for the single-stranded bc12Mid sequence. Substructure embedding was performed to produce a family of 100 DG structures. The embedded DG structures were then subjected to simulated annealing regularization. The experimentally obtained distance restraints and G-tetrad hydrogen-bonding distance restraints were included during the calculation. All distance restraints were specified with the SUM averaging option in X-PLOR (45). After simulated annealing, 97% of the molecules were folded in the correct topology, whereas 3% of the molecules were misfolded, e.g. in left-handed folding topology.

**NOE-distance restrained molecular dynamics calculations**

All of the 100 molecules obtained from the DGSA calculations were subjected to NOE-restrained Simulated Annealing refinement in XPLOR (45) with a distance-dependent dielectric constant. Atoms participating in hydrogen bonds in the G-tetrad planes were restrained with distances corresponding to ideal hydrogen bond geometry. Each individual hydrogen bond was restrained using two distance restraints (heavy atom–heavy atom–proton). Hydrogen-bonding distance restraints were also applied to A10:T15 bp, with larger distance bounds (±0.4 Å). All experimentally obtained distance restraints were specified ambiguously with the sum-averaging option. The force constants were scaled at 30 and 100 kcal mol$^{-1}$ Å$^{-2}$ for NOE and hydrogen bond distance restraints, respectively. A total of 476 NOE-distance restraints, which of 168 are from inter-residue NOE interactions, were incorporated into the NOE-restrained structure calculation.

Dihedral angle restraints were used to restrict the glycosidic torsion angle ($\chi$) for the experimentally assigned syn configuration, i.e. G1, G7, G8, G17 and G21, tetrad-guanines [60(±30)°], and A13 and A14 in the 3'-loop [60(±60)°], as well as for some of the experimentally assigned anti configuration bases, i.e. C4, G5 and C6 in the 5'-loop [220(±40)°]. Dihedral angle restraints were also used to restrain the sugar backbone torsion angles $\beta$, $\gamma$ and $\epsilon$ (46). Based on the J-coupling constants of $^{31}$P($n$)–H5'/H5''($n$) and H3'–H4'–H5'/H5''($n$) obtained from $^{31}$P–$^1$H COSY and HSQC experiments with various J-couplings, the $\beta$ angles were restrained to the $\beta$–t conformation at 180(±90)° for all the residues, except for G21 whose $\beta$ angle was restrained to −60(±20)°. The $\epsilon$ angles were restrained to 80(±20)° for G19, and to 90(±40)° for T15 and C6. Based on the relative intensities of H3'–H5'/H5'' and H4'–H5'/H5'', the $\gamma$ angles of the majority of residues with resolved H5'/H5'' are in the regular $\gamma$- conformation (−60°) or sometimes in the $\gamma$– conformation (−90°), because for each residue, the H3'–H5' (or H3'–H5'') NOE is clearly stronger than the H3'–H5'' (or H3'–H5') NOE, except for A14 and T15 which show similar intensities for H3'–H5' and H3'–H5'', and thus fall in the $\gamma$–t region (46). Thus only the $\gamma$ angles of A14 and T15 were restrained to 170(±90)°. The force constants of dihedral angle restraints were 10 kcal mol$^{-1}$ rad$^{-2}$ for $\chi$ and 5 kcal mol$^{-1}$ rad$^{-2}$ for $\beta$, $\gamma$ and $\epsilon$.

NOE-restrained simulated annealing refinement calculations were initiated at 300 K. The temperature was gradually increased to 1000 K in 4 ps. The system was equilibrated at 1000 K for 20 ps, and then slowly cooled to 300 K in 10 ps. The 20 best molecules were selected based both on their minimal energy terms and number of NOE violations, and were further subjected to NOE-restrained molecular dynamics calculations at 300 K for 25 ps. The coordinates saved every 0.1 ps during the last 2.0 ps of NOE-restrained molecular dynamics calculations were averaged, and the resulting averaged-structure was subjected to minimization until the energy gradient of 0.1 kcal mol$^{-1}$ was achieved. A soft planarity restraint of 1 kcal mol$^{-1}$ Å$^{-2}$ was imposed on the tetrads before the heating process and was removed at the beginning of the equilibration stage. The time steps for all processes of heating, cooling, and equilibration were equal to 0.5 fs. The 10 best molecules were selected based both on their minimal energy terms and number of NOE violations and have been deposited in the Protein Data Bank (accession no. 2F8U).

**RESULTS AND DISCUSSION**

**Basis for selection of the bc12Mid sequence with its dual G-to-T mutations as the predominant G-quadruplex in the BCL2 promoter**

In a recently published study (38), we have demonstrated that the six runs of G-tracts containing three or more contiguous guanines in the wild-type sequence (bc12Pu39, Figure 1A) can form three overlapping G-quadruplex structures, i.e., 5'G4, midG4, and 3'G4 (see Supplementary Table S1). Of these three overlapping sequences, midG4, or bc12MidG4 (Figure 1A), is by far the most stable. Since this sequence contains a run of five continuous guanines (Supplementary Table S1), the embedded DG structures were then subjected to simulated annealing regularization. The resulting averaged-structure was subjected to minimization until the energy gradient of 0.1 kcal mol$^{-1}$ was achieved. A soft planarity restraint of 1 kcal mol$^{-1}$ Å$^{-2}$ was imposed on the tetrads before the heating process and was removed at the beginning of the equilibration stage. The time steps for all processes of heating, cooling, and equilibration were equal to 0.5 fs. The 10 best molecules were selected based both on their minimal energy terms and number of NOE violations and have been deposited in the Protein Data Bank (accession no. 2F8U).
In addition, this same mutated sequence was previously shown to be the most stable using polymerase stop assay (see Figure 6 in Ref. 38). Therefore, the bcl2Mid sequence with the dual (G-to-T) mutation at the 5'-end of bcl2MidG4 (positions 15 and 16, Figure 1A) was chosen as the sequence for NMR structure determination. The reason why the other possible contiguous runs of three guanines in Figure 1A are unstable is that these mutational sequences eliminate the G3NG3 single-nt double-chain-reversal loop which provides stability to the quadruplex. Thus, it is clear that the dual 5'- (G-to-T) mutation selected can give rise to a specific and stable fold containing this G3NG3 single-nt double-chain-reversal motif. Importantly, the two mutated guanines (G15 and G16, Figure 1A) give rise to the predominant DMS cleavage pattern found in the wild-type sequence (38). Furthermore, the bcl2Mid sequence gives rise to a CD spectrum very similar to that of the wild-type sequence (39). We are therefore confident that the G-quadruplex from this sequence is not only the predominant one found in a single-stranded BCL2 promoter DNA template, but its dual G-to-T mutations force the sequence into the predominant loop isomer. This is an important point because mutants that change the folding pattern or result in isolation of the least stable species are less likely to be biologically relevant.

**bcl2Mid forms a monomeric G-quadruplex structure**

The bcl2Mid molecule exhibits a stronger propensity to aggregate than other G-quadruplex forming sequences we have worked with. Minor conformations are also present as indicated by the presence of weak resonances (Figure 1B), whose intensities are <5% of the major species and thus do not interfere with the unambiguous structural analysis of the predominant BCL2 G-quadruplex structure. A sample of bcl2Mid in potassium solution with a concentration over 3 mM, as has been routinely used for other G-quadruplex structures (41,43,47,48), shows a markedly increased background after a week and readily aggregates to a gel form after annealing from 95°C. We therefore used a lower concentration of 1.5 mM of bcl2Mid in potassium for our NMR structure determination (Figure 1B, lower panel). The sample of 1.5 mM bcl2Mid is stable in potassium solution for over 1 month which is sufficient for multiple 2D experiments. The two 1D1H spectra of the 1.5 mM bcl2Mid NMR sample, freshly made or in NMR solution for 5 weeks, are shown in Supplementary Figure S2A&B. No noticeable differences in 1D NMR spectra were observed for this sample after 5 weeks in solution. However, after over 1 year in NMR solution, a significantly increased level of broad resonance background in 1D NMR spectra is observed for this sample, even though the sharp peaks from the major species are still clearly observable (Supplementary Figure S2D).
The expanded H8/H6–H1′ region with assignments of the non-
exchangeable 2D-NOESY spectrum of bcl2Mid. The sequential assignment
pathway is shown. Missing connectivities are labeled with asterisks. The H8–
H1′ NOEs of the nucleotides with syn configuration are labeled by residue
names, while the H5–H6 NOEs of cytosines are also labeled for reference.

Figure 4. The expanded H8/H6–H1′ region with assignments of the non-
exchangeable 2D-NOESY spectrum of bcl2Mid. The sequential assignment
pathway is shown. Missing connectivities are labeled with asterisks. The H8–
H1′ NOEs of the nucleotides with syn configuration are labeled by residue
names, while the H5–H6 NOEs of cytosines are also labeled for reference.

The major G-quadruplex formed on the central BCL2 pro-
moter sequence has been shown to adopt a mixed parallel/
antiparallel-stranded folding (Figure 3A) (39). The imino
NH1 and base aromatic H8 protons of guanine residues of
this bcl2Mid were unambiguously assigned by the site-
specific low-concentration (6%) incorporation of 1, 2, 7-15N,
2,13C-labeled guanine nucleoside at each guanine position of
the sequence, one base at a time (Figure 3B and Supple-
mentary Figure S4) (39). The base H6 proton resonances of
thymines and cytosines were unambiguously assigned by sub-
stituting deoxyuridine (dU) for dT/dC one at a time at each
thymine/cytosine position of the sequence. Multiple 2D
experiments, including 2D-NOESY, TOCSY and COSY,
were carried out for bcl2Mid at a strand concentration of
1.5 mM in 20 mM pH 7.0 K-phosphate and 40 mM KCl at
various temperatures. Standard DNA sequential assignment
procedure was utilized for the assignment of the proton reso-
nances of bcl2Mid (Figure 4). The assignment of the aromatic
protons allowed the direct assignment of H1′ and H2′/H2′
resonances, which was then extended to other regions, includ-
ing those of H3′, H4′ and H5′/H5′. The proton chemical
shifts at 25°C are listed in Table 1. All proton resonances
have been unambiguously assigned, except some H5′/H5′
protons which cannot be differentiated. The ambiguity of
H5′/H5′ protons should not affect the NMR structure calcu-
lation, since the NOE intensities associated with H5′ or H5′
do not vary much regardless whether a resonance is assigned
as H5′ or H5′. In addition, very few NOE intensities associ-
it with H5′ or H5′ were used (see below). For all the resi-
dues with resolved H2′ and H2′ resonance, the H1′–H3′ NOE
is weaker than both the H1′–H2′ and H1′–H2′ NOEs, while
the H1′–H2′ NOE is stronger than the H1′–H2′ NOE, indicat-
ing a C2′-endo sugar pucker conformation.

Phosphorus resonance assignments of bcl2Mid

We have carried out 31P experiments on bcl2Mid, including
proton-decoupled 1D phosphorus spectroscopy, as well as
2D heteronuclear 31P–1H COSY and 31P–31P HSQC. The 1D
31P spectrum at 25°C in D2O is shown in Figure 5A, refer-
cenced to the phosphoric acid standard. While the majority of
the 31P resonances of bcl2Mid are clustered around
−1 ± 0.3 p.p.m., which has been shown to be the chemical
shifts for 31P resonances of regular right-handed B-DNAs,
there are a number of 31P resonances that are shifted outside
this region. Using 2D 31P–31P HSQC (Figure 5B) and the
proton assignments, we were able to assign all of the 31P reso-
nances of bcl2Mid (Table 1). Every phosphorus resonance
was assigned by using the assignments of sugar protons H3′, H4′,
and H5′/H5′, and correlating it to its 3′-coupled H3′ proton
and 3′-coupled H4′ and H5′/H5′protons. Moreover, the phos-
phorus assignments confirmed the proton assignments of
H3′(n − 1), H4′(n), and H5′/H5′(n) for each step. Very
interestingly, the phosphorus resonances that are observed out of
the −1 ± 0.3 p.p.m. region are all from the residues in the loop
regions of unique conformations, including the
downfield-shifted (from −1 p.p.m.) G21P (at C20–G21 step,
ΔP 1.5 p.p.m.), T15 (ΔP 0.88 p.p.m.), C4 (ΔP 0.54
p.p.m.), and C6 (ΔP 0.43 p.p.m.), and the upfield-shifted
G19 (ΔP −0.93 p.p.m.), C20 (ΔP −1.12 p.p.m.), and T16
(ΔP −1.61 p.p.m.). In particular, those phosphorus reso-
nances associated with the unique double-chain-reversal
single-nt loop (G19, C20, G21, see Figure 3A) and with the
unique conformation at T15–T16 (Figure 3A) are the most shifted.

Proton resonance assignments of bcl2Mid

The major G-quadruplex formed on the central BCL2 pro-
moter sequence has been shown to adopt a mixed parallel/
antiparallel-stranded folding (Figure 3A) (39). The imino
NH1 and base aromatic H8 protons of guanine residues of

1D proton spectra of bcl2Mid at various temperatures
(VTs) indicate that the melting temperature for the quadru-
plex structure in 60 mM K+ is around 65°C (Supplementary
Figure S3). The melting process of the G-quadruplex struc-
ture of bcl2Mid appears to be rather homogeneous throughout
the G-tetrad core structure. As described in our previous
report (39), bcl2Mid forms a monomeric intramolecular
G-quadruplex structure, as demonstrated by the independence
of the melting temperature from the concentration, the sharp
NMR spectral line widths (Figure 1B), and the EMSA
(electrophoretic mobility shift assay) results. The formation
of a unimolecular structure in bcl2Mid is unambiguously con-

figured by the NMR stoichiometry titration experiment at
the melting temperature using DNA concentrations from
0.01 to 0.1 mM (43,49) (Figure 2). The NMR spectra of
bcl2Mid at a strand concentration of 1.5 mM (for 2D NMR
experiments and structure determination) and 0.1 mM (for
1D experiments and the titration experiment) are the same
(Figure 1B, upper and lower panels), indicating that the same
unimolecular G-quadruplex structure is formed in all
the conditions used for our NMR analysis. In addition,
DOSY (Diffusion Ordered Spectroscopy) experiments
on bcl2Mid also confirmed that the sharp NMR signals are
from a molecular weight equivalent to a monomeric structure.
(The DOSY results will be published elsewhere.)
The J-coupling constants of $^{31}$P($\pi$)–H5/H5$^\prime$(n) and H3($\pi$)–$^{31}$P(n + 1) were obtained from $^{31}$P–$^1$H COSY and confirmed by HSQC experiments with various J-couplings. The $^{31}$P–H5$^\prime$ and $^{31}$P–H5$^\prime$ coupling constants can define the $\beta$ torsion angle quite well (46). Except for G21, all other observed $^{31}$P($\pi$)–H5/H5$^\prime$(n) coupling constants are below 10 Hz, indicating the $\beta$ torsion angles fall in the common trans-region, 180(±90)$^\circ$. G21 gives rise to a $^{31}$P–H5$^\prime$ coupling constant of $\sim$12 Hz and no observable $^{31}$P–H5$^\prime$ coupling, indicating a $\beta$ torsion angle in the $\beta$-region, $\sim$60(±20)$^\circ$. Most H3$^\pi$–$^{31}$P couplings are <10 Hz, indicating that the $\varepsilon$ angles fall in the common trans/gauche- conformation (170$^\circ$–300$^\circ$), at which range the $\varepsilon$ angles can not be accurately determined by the H3($\pi$)–$^{31}$P(n + 1) coupling constants due to several possible $\varepsilon$ angles for one H3$^\pi$–$^{31}$P coupling in this region (46). However, when the H3($\pi$)–$^{31}$P(n + 1) coupling constants are larger than 10 Hz, it is possible to more accurately determine the $\varepsilon$ angles. The H3($\pi$)–$^{31}$P(n + 1) coupling constants for G19(–C20P), T15(–T16P) and C6(–G7P) are 15, 12 and 12 Hz, respectively, indicative of $\varepsilon$ angles in the gauche+/trans- region (centered $\sim$80–90$^\circ$). It is again interesting to note that the residues exhibiting unusual large coupling constants are all associated with the loop regions, especially with the single-nt double-chain-reversal loop. It may be noted that for non-B-DNA structures, the $^{31}$P chemical shifts show a very poor correlation with the $\varepsilon$ angle values (50).

**NOE interactions within G-tetrad regions in the bcl2Mid G-quadruplex**

Many inter-residue NOE crosspeaks are observed in 2D-NOESY and are summarized in Figure 6. Only one NOE intensity is associated with H5$^\prime$ or H5$^{\prime\prime}$. An expanded NOESY spectrum of base and sugar H1$^\prime$ protons is shown in Figure 4. Five guanine residues of the core G-tetrads are in the syn-configuration, including G1, G7, G8, G17 and G21 (Figure 3A), as indicated by the very strong H8–H1$^\prime$ NOE intensities (Figure 4). A characteristic downfield shift is observed for the H2/H2$^\prime$ sugar protons of the syn-guanines (Table 1). The sequential NOE crosspeak connectivities of the base H8 protons to the 5$^\prime$-flanking residue sugar H1$^\prime$/H2$^\prime$/H2$^{\prime\prime}$ protons, typical for right-handed DNA twist, are observed for bcl2Mid (Figure 4). Bcl2Mid adopts a mixed parallel/antiparallel-stranded G-quadruplex structure, with the first, third and fourth G-strands being parallel with each other, and the second G-strand being antiparallel with the rest of the strands (Figure 3A) (39). The top two G-tetrads have the same anti/syn distribution and are connected by guanines with the same sugar configurations, while the third G-tetrad has a reversed anti/syn distribution and is connected with the middle G-tetrad by guanines with different sugar configurations (Figure 3A). In accord with the topology, the sequential NOE connectivities are indeed either missing or very weak at the N(i)–synG(i + 1) steps, i.e. C6–G7, G7–G8, T16–G17 and C20–G21. The characteristic G(i)H8/G(i + 1)H1$^\prime$ NOEs are observed when G(i) is in the syn configuration, such as G1–G2, G7–G8, G8–G9 and G17–G18 (Figure 4). The same configurations of guanines (syn–syn or anti–anti steps) connecting the top two G-tetrads are reflected by clear NOE interactions of adjacent guanine base protons, e.g. G2H1/G3H1, G7H1/G8H1, G18H1/G19H1 and G22H1/G23H1 (Supplementary Figure S5). The reversed configurations of guanines connecting the bottom two G-tetrads are clearly reflected by strong inter-tetrad NOE interactions of guanine imino protons, such as G2H1/G9H1.
G8H1/G17H1, G18H1/G21H1 and G22H1/G1H1 (Supplementary Figure S5). Furthermore, the right-handedness of the DNA backbone of the G-quadruplex is clearly indicated by inter-tetrad NOE interactions, including G3H1/G8H8, G7H1/G18H8, G19H1/G22H8 and G23H1/G2H8 (Figure 7).

NOE interactions within loop regions in the bcl2Mid G-quadruplex

The sequential NOE connectivities are interrupted at the loop regions, including C4–G5–C6, A10–G11–G12–A13–A14–T15–T16, and G19–C20–G21 (Figure 4), indicating a poorly stacked loop conformation. All residues in the loop regions are in the anti conformation except A13 and A14, as indicated by the strong H8–H1' NOE intensities (Figure 4). A number of interesting NOE interactions are observed for both the lateral loops, whereas the sequential NOE connectivities are almost all missing for the double-chain-reversal single-nucleotide C loop, except a couple of NOEs between the sugar protons of G19 and C20 (Figure 6). For the 5'-CGC lateral loop, sequential connectivities are clearly observable within G5–C6–G7. Interestingly, clear NOEs are observed between G5/C6 and the imino protons of the top tetrad, such as G7H1/C6H6, H5&H1', G19H1/C6H5&H6, G3H1/G5H8&H1', and G23H1/G5H1' (Figure 4, complete summary in Figure 6), indicating that the G5 and C6 residues are stacking over the top G-tetrad. On the other hand, unusual NOEs are observed within the G3–C4–G5 region. Strong NOEs are observed for C4 base protons H5&H6 with G3 sugar protons H1'&H4', e.g. C4H5/G3H4' and C4H6/G3H1' (strong), C4H5/G3H1' and C4H6/G3H4' (medium strong), but not with G3H3' or H2'. No NOEs are observed between C4H6/H5 and G3H8 for base stacking. For the C4–G5 step, clear NOEs are observed for G5H8 with C4H3' (strong), H2' (strong) and H2'' (medium strong), but not with C4H1' or H4'. These NOE data suggest an unusual structure adopted within this region, where the C4 base is not stacked over the top G-tetrad but is positioned into the groove that is close to the H1'/H4' side of the G3 sugar. This will be discussed in more detail later.
For the middle 7-nt lateral loop, A10–G11–G12–A13–A14–T15–T16, A10 is very well stacked with G9 and the bottom G-tetrad, as is evident by the clear NOE interactions, such as A10H2/G9H1&G17H1 (Figures 6 and 7). Very interestingly, clear NOEs are observed between T15, instead of T16, and the bottom G-tetrad, such as T15H6/G1H1&G21H1, and T15H1/G21H1 (Figures 6 and 7), indicating that T15, but not T16, is stacked right on the bottom G-tetrad. Furthermore, a clear imino proton is observed for T15H3, which exhibits a very strong NOE interaction with A10H62 (Figure 7), indicating that T15 is likely to be involved in a stable H-bonded conformation with A10.

Mutational analysis for loop segments

We have carried out systematic analysis of BCL2 promoter sequences with mutated residues in the loop regions to determine their functional role in BCL2 G-quadruplex formation and stability. We incorporated one mutation at a time for each loop residue, using various substitutions as listed in Table 2, and then collected NMR spectra on each sample. Mutations that do not induce changes in the NMR spectra are marked with a ‘/C0’ symbol, whereas those that do induce clear spectral changes are marked with a ‘+/C+’ symbol.

Interestingly, the most sensitive positions are C4 and A10, which cannot tolerate any substitutions.

NOE-restrained structure calculation

NOE data (Figure 6) define the overall topology of this mixed antiparallel/parallel G-quadruplex and were used for NOE-restrained structure calculation of the major BCL2 G-quadruplex formed with bcl2Mid. Solution structures of this G-quadruplex were calculated using a NOE-restrained distance geometry (DGSA) and molecular dynamics (RMD) approach, starting from an arbitrary extended single-stranded DNA model. A total of 476 NOE distance restraints, of which 168 are from inter-residue NOE interactions, were

Table 2. Effect of single base substitutions on the G-quadruplex formation of the Bcl2Mid sequence

| Residue | C | T | U | A | I |
|---------|---|---|---|---|---|
| C4      | + |  |   |   |   |
| C5      |   |   |   |   |   |
| C6      |   | + |   |   |   |
| A10     |   | + |   |   | + |
| G11     |   |   |   |   |   |
| G12     |   |   |   |   |   |
| A13     |   |   |   |   |   |
| A14     |   |   |   |   |   |
| T15     |   |   |   |   |   |
| T16     |   |   |   |   |   |
| C20     |   |   |   |   |   |

*: The G-quadruplex formation is not affected; +: the G-quadruplex formation is destabilized.

Table 3. Structural statistics for the Bcl2Mid

| Distance restraints | 476 |
|---------------------|-----|
| Intraresidue        | 309 |
| Interresidue        | 167 |
| Sequential (|i–β| = 1) | 107 |
| Non-sequential (|i–|j| > 1) | 60 |
| Hydrogen bonds      | 26  |
| Total dihedral angle restraints | 38 |

| Violations (mean and s.d.) | 0.03 ± 0.003 |
| Bond length (Å)           | 0.007 ± 0.0001 |
| Bond angle (°)            | 1.40 ± 0.01  |
| Improper (°)              | 1.05 ± 0.01  |
| Average pairwise r.m.s.d. of heavy atoms (Å) | 1.01 ± 0.12 |
| G-tetrads                | 1.01 ± 0.12 |
| With C20                 | 1.14 ± 0.14 |
| With C4,G5,C6            | 1.04 ± 0.12 |
| With A10,T15             | 1.13 ± 0.14 |
| All residues             | 2.55 ± 0.65 |

*: The ensemble of 10 structures is selected based both on the minimal energy terms and number of NOE violations.

Interestingly, the most sensitive positions are C4 and A10, which cannot tolerate any substitutions.

For the middle 7-nt lateral loop, A10–G11–G12–A13–A14–T15–T16, A10 is very well stacked with G9 and the bottom G-tetrad, as is evident by the clear NOE interactions, such as A10H2/G9H1&G17H1 (Figures 6 and 7). Very interestingly, clear NOEs are observed between T15, instead of T16, and the bottom G-tetrad, such as T15H6/G1H1&G21H1, and T15H1/G21H1 (Figures 6 and 7), indicating that T15, but not T16, is stacked right on the bottom G-tetrad. Furthermore, a clear imino proton is observed for T15H3, which exhibits a very strong NOE interaction with A10H62 (Figure 7), indicating that T15 is likely to be involved in a stable H-bonded conformation with A10.
incorporated into the NOE-restrained structure calculation (Table 3). Dihedral angle restraints were used for the glycosidic torsion angle ($\chi$) for all the syn residues, and for the three anti residues, i.e. C4, G5 and C6, in the 5'-loop. Dihedral angle restraints were also used to restrain the sugar backbone torsion angles $\beta$, $\gamma$, and $\epsilon$ (Materials and Methods).

The superimposition of the 10 lowest energy structures produced by the refinement is shown in Figure 8 (PDB ID 2F8U). The structure statistics are listed in Table 3. An average of ~23 restraints per residue was used for the solution structure calculation, including experimentally observed H-bond interactions for the G-tetrads, whereas no planarity constraints were used for RMD calculations.Remarkably, out of the 10 lowest energy structures, the RMS deviation for distance violations is only 0.03 Å (Table 3). The bcl2Mid G-quadruplex structure is very well defined, with a RMSD of 1.01 Å for the three G-tetrads. The 5'-CGC lateral loop is also well defined, as the RMSD of the three G-tetrads and the 5'-CGC loop is 1.04 Å. The RMSDs are 1.14 Å and 1.13 Å, respectively, when including the C20 single-nucleotide double-chain-reversal loop or the A10/T15 of the middle 7-nt lateral loop. The RMSD for all residues is 2.55 Å, indicating a much less-defined middle 7-nt loop.

Molecular structure of the major G-quadruplex structure in the BCL2 promoter

A representative model of the bcl2Mid G-quadruplex structure is shown in different views in Figure 9 and Supplementary Figure S6. The G-quadruplex consists of three G-tetrads linked with mixed parallel/antiparallel right-handed G-strands that are connected by the first two lateral loops (CGC and AGGAATT) and a third single-nucleotide (nt) double-chain-reversal side loop (C), as also shown in our previous report (39). The first, third and fourth G-strands of this G-quadruplex are parallel with each other, while the second G-strand is antiparallel with the rest of the G-strands. The bcl2Mid G-quadruplex contains one wide groove (groove I, between the first and second antiparallel G-strands), one narrow groove (groove II, between the second and third antiparallel G-strands), and two intermediate grooves (groove III and IV, between the third, fourth, and first parallel G-strands) (Supplementary Figure S6). All the $\delta$ torsion angles fall in...
the range of 110–150°C, consistent with the C2′-sugar pucker conformations indicated by the NMR data.

The three G-tetrads are well defined, with the top two G-tetrads having the same arrangement of guanine configuration (anti:syncanti) and the bottom G-tetrad having the reversed arrangement (syn:anti:sync:sync). Extensive stacking between the guanine five-membered rings is observed for adjacent guanines of the bottom two G-tetrads with the alternate guanine glycosidic configurations (Figure 10A), while only partial stacking is observed for adjacent guanines of the top two G-tetrads with the same guanine glycosidic configurations (Figure 10B).

Loop conformation and functional role of loop residues

The NMR solution structures indicate that the three loop regions interact with the core G-tetrads in a specific way that defines and stabilizes the unique BCL2 G-quadruplex structure. The first C4–G5–C6 loop linker that connects the first and second antiparallel G-strands forms a unique lateral loop conformation which is very well defined (Figures 9A and 10C). G5 and C6 stack right on the top G-tetrad, where G5 stacks very well with G3 and C6 is positioned above the center of G7 and G19 (Figure 10C). Interestingly, the C4 residue adopts a unique conformation in which its base is positioned in the wide groove I and is perpendicular to the G-tetrad planes (Figure 9A), covering the outside of the G3 residue of the top tetrad. This unique conformation of C4 appears to be rather favored, as over 90% of the 100 structures from our DGSA calculation are in this conformation.

The conformation of the C4–G5–C6 loop is well defined as indicated by the small root mean square deviation of this loop region (Figure 8 and Table 3) (see above). The unique conformation of C4 explains the unusual NOE interactions observed between C4 and G3/G5 as discussed before, e.g. strong NOEs between the C4 H5/H6 and G3 H1′/H4′, and the lack of NOE between the G5H8 and the C4 H1′ (Figure 6).

The second 7-nt A10–G11–G12–A13–A14–T15–T16 linker that connects the second and third antiparallel G-strands also forms a lateral loop conformation in the NMR structure (Figure 9). A10 stacks very well with G9 and the bottom tetrad, while T15, which is the second residue from the other end of the loop, stacks well with the bottom G-tetrad. Potential reversed Watson–Crick hydrogen bonds could be formed between A10 and T15 (Figure 10D), as the imino proton H3 of T15 can be detected at 1°C (Figure 7). T16, which is sequentially adjacent to G17, however, is looped out from the bottom G-tetrad (Figure 9). The remaining four residues of this long lateral loop are not very well defined (Figure 8) and are mostly exposed to the solvent (Figure 9).

Significantly, bcl2Mid contains a G3NG3 sequence motif, which adopts a single-nucleotide (C20) double-chain-reversal loop conformation connecting two parallel G-strands (Figure 3A), representing another example of a very stable parallel-stranded structural motif, in addition to the one first observed in the MYC promoter sequence (41,51,52). The conformation of this single-nucleotide double-chain-reversal loop (C20) is very similar to those of the MYC G-quadruplex (41). The right-handed DNA backbone twist brings the ends of the two adjacent parallel G-strands very close to each other spatially so that the single-nucleotide double-chain-reversal loop conformation is rather favored (Figure 9B and Supplementary Figure S6).

The loop conformations are defined by the experimental NOEs and agree well with the experimental data (Figure 6 and Table 2). For the first 3-nt CGC lateral loop, C4, which is positioned in groove I, cannot even tolerate a substitution of uridine, which is the closest nucleotide to cytosine, indicating that the amino group of cytosine is clearly more favored for the groove positioning, as reported previously for DNA groove-binding ligands (53,54). On the other hand, G5 and C6, which appear to be able to tolerate various substitutions, can presumably still retain the stacking interactions with different residues. For the second 7-nt AGGAATT lateral loop, in accord with the NMR structure in which A10 forms a well-defined H-bonded conformation that stacks extensively with G9 of the bottom tetrad, the A10 residue cannot tolerate any substitution in our mutational analysis. Interestingly, T15 and T16, which are the two mutated residues, display much higher tolerance and can be substituted with uridines and adenosines, indicating a greater conformational flexibility of these two positions. The rest of the loop residues display much higher flexibility and can tolerate most substitutions. Indeed, G11 and G12, which adopt an extended loop conformation, show enhanced cleavage in the DMS footprinting study (38). Finally, the C20 single-nucleotide double-chain-reversal loop appears to be able to tolerate various substitutions.

Figure 10. Stacking interactions between (A) the middle (magenta) and bottom (green) G-tetrads, which have reversed anti:syncanti:anti and sync:anti:sync:sync arrangements, and (B) the top (cyan) and middle (magenta) G-tetrads, which have the same anti:syncanti:anti arrangements; and stacking interactions between (C) the first lateral loop C4–G5–C6 (cyan) and the top G-tetrad (pink), and (D) the bottom G-tetrad (orange) and the A10:T15 bp (green) from the second lateral loop, A10–G11–G12–A13–A14–T15–T16. Figures are prepared using PyMOL.
The BCL2 G-quadruplexes and their relationship to transcriptional control and other G-quadruplex-containing promoter elements

The G-quadruplexes in the BCL2 promoter element represent a more complex architecture than those found in other promoters, such as the c-Myc NHE III. In this contribution we have characterized the major loop isomer from the bcl2MidG4 sequence that overlaps with two other G-quadruplexes (5′G4 and 3′G4). In principle there is a total of 15 different possible loop and structural isomers in the BCL2 promoter element. Since this element (bcl2Pu39, Figure 1A) also overlaps with a binding site for WT1 transcriptional factor, which is a suppressor protein, it may be that formation of one or more of the G-quadruplexes results in transcriptional activation. In addition to the possible transcriptional activation role of the G-quadruplexes in this region, it is also possible that the emergence or elimination of individual G-quadruplexes in this region may differentially regulate gene expression by selectively interacting with transcriptional factors that either activate or suppress BCL2 gene expression.

A common feature of the most stable BCL-2 loop isomer studied and the c-Myc promoter G-quadruplex (41,51,52), and likely the VEGF and HIF-1 studied and the c-Myc promoter G-quadruplex (41,51,52), is the single-nitrogen-containing double-chain-reversal loop. Only this isomer of the bcl2MidG4Pu23 retains this feature that is essential for stability of these G-quadruplexes. The single less stable G4-TTA-G4 double-chain-reversal loop found in the human telomeric G-quadruplex (43) may be important for the inherent more facile reversibility of the G-quadruplex to alternative folding structures.

Last, the diversity in sequences and resulting folding patterns found within the BCL2 promoter gives rise to differential drug binding (38), which may mimic protein recognition differences. From a drug design perspective, this diversity in drug binding provides opportunities for selectivity. Furthermore, the major G-quadruplexes formed in BCL2 and MYC promoter sequences are different (41), with distinct folding patterns, G-tetrad conformations, loop conformations, and groove conformations, suggesting that such regions can be differentially targeted by G-quadruplex-interactive agents. The structural diversity of the G-quadruplexes formed in oncogene promoter regions makes such regions attractive targets for pathway-specific drug design.

SUPPLEMENTARY DATA
Supplementary Data is available at NAR Online.

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