Research Article

Novel Indolocarbazole Derivative 12-(α-L-arabinopyranosyl)-indolo[2,3-a]pyrrolo[3,4-c]carbazole-5,7-dione Is a Preferred c-Myc Guanine Quadruplex Ligand

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Received 26 November 2010; Accepted 4 March 2011

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The indolocarbazole derivative 12-(α-L-arabinopyranosyl)indolo[2,3-a]pyrrolo[3,4-c]carbazole-5,7-dione (AIC) has demonstrated a high potency (at nanomolar to submicromolar concentrations) towards the NCI panel of human tumor cell lines and transplanted tumors. Intercalation into the DNA double helix has been identified as an important prerequisite for AIC cytotoxicity. In this study, we provide evidence for preferential binding to the G-quadruplex derived from the c-Myc oncogene promoter (Pu18 d(AG3TG4)2; G-c-Myc). The association constant for AIC : G-c-Myc complex was ∼100 times and 10 times greater than the respective values for the complexes AIC : c-Myc duplex and AIC : telomeric d(TTAGGG)4 G-quadruplex. The concentrations at which AIC formed complexes with G-c-Myc were close to those that attenuated the steady-state level of the c-Myc mRNA in the human HCT116 colon carcinoma cell line. We suggest that preferential binding of AIC to G-c-Myc rather than to the c-Myc duplex might favor the quadruplex formation in the cells, thereby contributing to downregulation of the c-Myc expression by AIC.

1. Introduction

The c-Myc oncoprotein is an important transcription factor that plays a pivotal role in cell proliferation and survival [1]. Deregulation of c-Myc gene expression has been detected in many tumor types and is believed to be an important step in tumorigenesis [2]. A major control element of the human c-Myc gene is a purine/pyrimidine rich region (Pu27; the nuclease-hypersensitive element 1) located 115 bases upstream from the P1 promoter. This element controls up to 95% of the total c-Myc transcription [3, 4]. The Pu27 segment has been shown to adopt an intramolecular DNA tetraplex (G-quadruplex) conformation under physiological conditions [5]. The oligomer containing four consecutive 3′ runs of guanines d(AGGGTGAGGGTGAGGG) (Pu18) is the minimal prerequisite for the c-Myc G-quadruplex (G-c-Myc) formation [6, 7].

Involvement of the G-quadruplex structures in regulation of gene transcription [8, 9], RNA metabolism [10] and telomere function [11] opens an area for design of small molecular weight compounds that can preferentially interact with G-quadruplex DNA. The distinctive nature of quadruplex topologies suggests that quadruplex ligands may selectively silence a given gene [8] with little effect on other DNA regions containing noncanonical higher-order structures. A number of G-quadruplex ligands have been designed that demonstrated either similar affinity to various G-quadruplexes or preferential binding to a particular tetraplex. For example, the porphyrin derivative TMPyP4, the cyclic polycation telomestatin, and 2,6-pyridin-dicarboxamide recognized both G-c-Myc and telomeric G-quadruplex (G-tel) [12, 13]. In contrast, fluoroquinolone derivative quinofloxin, the first therapeutic G-c-Myc ligand that entered clinical trials, did not show the cellular behavior characteristic of
the telomere-targeting agent [11, 13]. The trisubstituted isoalloxazines specifically interacted with the c-Kit sequence and downregulated c-Kit oncogene expression [14].

Indolocarbazoles and their congeners, a class of extensively investigated antitumor drugs, interact with multiple intracellular targets [15]. The mechanisms of antitumor effects of carbohydrate derivatives of indolocarbazoles include DNA damage and inhibition of the enzymes critical for cell viability, in particular, topoisomerase I, cyclin-dependent kinases, and the checkpoint kinase 1 [15–17]. The carbohydrate moiety of indolocarbazoles can enhance the binding of the conjugate to a target [16]. However, the binding constants to the duplex and G-quadruplex DNAs of indolocarbazoles containing one carbohydrate substituent are relatively low (Ka ≤ 2 × 10^5 M^−1) [18–20]. In contrast, we found the association constant to native double stranded DNA (dsDNA) of the novel indolocarbazole derivative, 12-(α-L-arabinopyranosyl)indolo[2,3-a]pyrrolo[3,4-c]carbazole-5,7-dione (AIC, Figure 1) to be substantially higher (1.6 × 10^6 M^−1 ≤ Ka ≤ 3.3 × 10^6 M^−1) [21, 22]. AIC potently inhibited growth of 60 cultured human tumor cell lines (National Cancer Institute drug screening panel). This compound demonstrated promising in vivo therapeutic efficacy against transplanted tumors. The concentrations of AIC required for the formation of intercalative complexes with dsDNA and the concentrations that induced tumor cell death were similar, suggesting that intercalation into DNA is an important factor of cytotoxicity of this indolocarbazole derivative [22].

In this study we analyzed the interactions of AIC with G-quadruplex DNA structures. We explored the affinity and selectivity of AIC interaction with the intramolecular G-c-Myc formed by Pu18 [6]. The association constants of AIC:G-c-Myc complexes were exceptionally high, ~100 times greater than the respective value for the complexes of AIC with 18-bp c-Myc duplex DNA. The concentrations at which AIC formed complexes with G-c-Myc were close to the concentrations that attenuated the steady-state level of the c-Myc mRNA in the human HCT116 colon carcinoma cell line.

2. Material and Methods

2.1. Reagents and Sample Preparation. All reagents were from Sigma-Aldrich, St. Louis, Mo unless otherwise specified. The oligonucleotides d(AG3TG4)2 (G-c-Myc), d(C4AC3T)2 (comp-c-Myc) and d(TTAGGG)4 (G-tel) were synthesized by Syntol (Moscow). The DNA preparations were dissolved in Hanks’ buffer (0.137 M NaCl, 5.4 mM KCl, 0.25 mM Na2HPO4, 0.44 mM KH2PO4, 1.3 mM CaCl2, 1 mM MgSO4, 4.2 mM NaHCO3, pH 7.2) immediately before the experiments. The molar extinction coefficient of AIC was ε(220) = 22 000 M^−1. All experiments were performed in Hanks’ buffer at 20°C.

2.2. Instruments. Absorption spectra were acquired with a Jasco V-550 spectrophotometer (Japan). Fluorescence was registered with a Cary Eclipse spectrophotofluorimeter (Varian Inc.). CD spectra were recorded using a Jasco 715 spectropolarimeter (Japan). Molar dichroism (Δε) was calculated per mole of nucleotides. All instruments were equipped with thermostated cell holders.

2.3. Binding of AIC to DNA. The binding of AIC to various DNAs was monitored by UV light absorption and spectrofluorimetry [23]. The AIC:DNA complex formation was represented by Scatchard plots, that is, by the ratio r/Ci as the function of r, where Ci is concentration of free AIC, and r is the number of bound AIC molecules per G-quadruplex or DNA duplex. The experimental data were fitted by (1) for several independent types of bound molecules:

\[
r = \sum_{i=1}^{N} \frac{n_i K_i C_i}{1 + K_i C_i},
\]

where N is the number of independent types of binding, n_i is the maximal number of molecules that participate in one type of binding, K_i is the association constant of i type of complex, and C_i is the concentration of unbound AIC.

Studies of energy transfer and circular dichroism (CD) spectra were performed as described by Kaluzhny et al. [22].

2.4. Measurement of c-Myc mRNA by Quantitative Real-Time PCR. The HCT116 human colon carcinoma cell line (American Type Culture Collection, Manassas, Va) was cultured in RPMI-1640 supplemented with 5% fetal calf serum (HyClone, Logan, Ut), 2 mM L-glutamine, 100 U/mL penicillin, and 100 μg/mL streptomycin at 37°C, 5% CO2 in a humidified atmosphere. Cells (2 × 10^5 in 6 mL of culture medium) were plated onto 60 mm Petri dishes overnight. Then AIC was added, and the cells were incubated for 24–48 h. An Agilent Total RNA mini-isolation kit (Agilent, Tex) was used for RNA isolation. Reverse transcription was performed as previously described [24]. For quantitative real-time PCR, cDNA samples were amplified with primers

![Figure 1: Structure of 12-(α-L-arabinopyranosyl)indolo[2,3-a]pyrrolo[3,4-c]carbazole-5,7-dione (AIC).](image-url)
corresponding to c-Myc (forward 5′-AAAGACACGGCG-AGCCCGAA-3′; reverse 5′-CCAGTCCTGCATGCCTCGC-AAG-3′) and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (forward, 5′-GAAGGTGAGATCAACGAGATT-3′; reverse 5′-GCCAAAAGGGTCATCATCTCT-3′). The reactions were performed using Stratagene Brilliant II SYBR Green QPCR Master Mix on a Stratagene MX 3005P thermocycler. The MxPro (QPCR Software) was used for data analysis. The amounts of c-Myc mRNA in AIC-treated cells were expressed as the percentage of the signal intensity to that in control (untreated) samples (regarded as 100%).

3. Results

3.1. UV Absorption of AIC:DNA Complexes. The UV absorption spectra of free AIC showed two prominent bands at 285 nm and 318 nm and a weaker band at 425 nm [22]. The absorption near λ = 318 nm decreased upon binding of AIC to G-c-Myc, G-tel, and c-Myc duplex d(AGGGTTGGG)2(d(CCCACACCT)); in Hanks’ buffer at 20°C (Figure 2). The absorption maximum shifted from 318 nm to ∼325 nm. The spectral changes were similar to those observed for AIC intercalation into dsDNA [22]. Using the absorption at λ = 318 nm we plotted the AIC-binding curves to G-c-Myc, G-tel, and c-Myc duplex in Scatchard coordinates, r/C1 on r (Figure 3, triangles). Based on these curves we calculated maximal numbers of AIC molecules bound to the oligonucleotides: three molecules to G-c-Myc, two to G-tel, and five to the c-Myc duplex.

3.2. Fluorescence of AIC:G-Quadruplex Complexes. Fluorescence emission at 500 < λ < 650 nm centered at ∼550 nm was detectable upon binding of AIC to G-c-Myc (Figure 4(a), solid curve) and to G-tel (Figure 4(a), triangles). Similar fluorescence emission was typical for AIC intercalation into dsDNA [22]. In aqueous solution the fluorescence of the drug was quenched [20]. Therefore, the fluorescence spectrum of free AIC in ethanol was taken for comparison with the respective spectra of complexes AIC:G-c-Myc and AIC:G-tel (Figure 4(a), dashed curve).

The dependencies of fluorescence intensities (I550) of AIC:G-c-Myc and AIC:G-tel on r value (calculated using the binding curves in Figures 3(a) and 3(b); triangles) were plotted in Figure 4(a), inset (triangles and circles, resp.). The fluorescence intensity of AIC:G-c-Myc complexes increased linearly with the concentration of bound AIC (0 < r ≤ 1). The fluorescence intensity increased much more slowly at r > 1 (note ~10-fold smaller slope of the curve). Thus, the fluorescence of the second and the third bound AIC molecules (1 < r ≤ 3) quenched significantly. In contrast, the fluorescence intensity of AIC:G-tel grew linearly at 0 < r ≤ 2. Apparently, two AIC molecules bound to G-tel fluoresced with the same quantum yield.

These results strongly suggest that only the first AIC molecule tightly bound to G-c-Myc is fluorescent being placed in a hydrophobic environment. Such a highly fluorescent AIC molecule may stack between the terminal G-quartet and 5′-adenine [7]. Two other AIC molecules whose fluorescence is markedly quenched are supposed to bind to the two propeller loops in a less hydrophobic environment. One may hypothesize that the fluorescence of AIC bound to the loops might be quenched by water molecules. Two hydrophobic binding sites are present in the antiparallel G-tel. The AIC molecules may be stacked between each of the terminal G-quartets and the bases of lateral and diagonal loops across the quartets [25]. The structures of the two sites are not identical; therefore, the binding curve is nonlinear (Figure 3(b), triangles).

3.3. Parameters of AIC Binding to Various DNA Structures. We generated the Scatchard plot (r/fC1 on r) for the binding of fluorescent AIC molecules to G-c-Myc (Figure 3(a), squares). The C1 values were derived from the binding curve drawn on the basis of UV absorption data (Figure 3(a), triangles). The binding curve revealed a single “strong” binding site, Kf = (7.0 ± 1) · 106 M−1. We subtracted the contribution of the fluorescent molecules (Figure 3(a), squares) from the binding curve constructed using UV absorption (Figure 3(a), triangles). The resulting Scatchard plot revealed the two quenched ligand molecules with the association constant K2 = (4 ± 1) · 105 M−1 (Figure 3(a), circles). The curve in Figure 2(a), triangles, reflects the formation of two types of complexes, that is, the fluorescent K1 = (7 ± 1) · 105 M−1, n1 = 1 and the nonfluorescent (quenched) K2 = (4 ± 1) · 105 M−1, n2 = 2, respectively (Table 1). This curve was analyzed using a model for two independent types of drug-binding sites (1). The theoretical plot corresponding to the K1, K2, n1, n2 values (Table 1) is shown in Figure 2(a), solid curve.

The curve of AIC binding to the G-tel generated using the fluorimetric method (Figure 3(b), squares) coincided with the binding curve obtained by UV absorption (Figure 3(b), triangles). The binding curve may be approximated with two independent binding sites. The calculated association constants are given in Table 1. The association constants for the strong and the weak AIC:G-tel complexes were ~10 times lower than those for AIC:G-c-Myc complexes (Table 1).

The Scatchard plot for AIC binding to the 18 bp c-Myc duplex (generated by UV absorption) was linear at 2 ≤ r ≤ 4 (Figure 3(c). The binding curve was approximated with the equation for a single type complex. The 18 bp c-Myc duplex formed complexes with five inner AIC molecules, K = (8 ± 1) · 105 M−1 (Table 1). The association constant is close to that for strong AIC binding to dsDNAs of various nucleotide content 1.6 · 106 < K1 ≤ 3.3 · 106 M−1 [22]. Importantly, the affinity of AIC to c-Myc duplex appeared to be ~100 times weaker than to G-c-Myc (Table 1). Hence, the G-quadruplex conformation(s) may be energetically favorable for binding of AIC to DNA within the c-Myc regulatory region. Altogether, G-c-Myc emerges as a preferred target for AIC.

3.4. Energy Transfer from G-Quadruplexes to AIC. To investigate whether the stacking contacts and intercalation might be the modes of AIC:G-quadruplex complex formation, we studied the energy transfer from the bases of G-c-Myc or
Figure 2: UV absorption spectra of AIC:DNA complexes: (a) AIC:G-c-Myc, (b) AIC:G-tel, and (c) AIC:c-Myc duplex. The concentrations of oligomers Pu18 and d(TTAGGG)4 were 0–2.5 μM (strands) and of c-Myc duplex were 0–3 μM (strands). Curves 1–3 indicate the spectra of free AIC, partially bound AIC and fully bound AIC, respectively: Hanks’ buffer, t = 20°C.

Figure 3: Scatchard plots of AIC:DNA binding. (a) The curve of AIC binding to G-c-Myc (Pu18) registered by UV absorption (triangles) and spectrofluorimetry (squares). The circles represent absorption of nonfluorescent complexes. (b) The curve of AIC binding to G-tel registered by UV absorption (triangles) and spectrofluorimetry (squares). (c) The curve of AIC binding to Pu18 c-Myc duplex registered by UV absorption. The parameters r and \( r_f \) were calculated per DNA strand: Hanks’ buffer, t = 20°C.
molecules per G-c-Myc in 95% ethanol solution (dashed line). Fluorescence intensity I normalized to 1. The parameter r in Figure 4(b). The fluorescence intensity of free AIC in ethanol were taken for comparison with the respective spectra of complexes AIC:G-c-Myc (solid line), AIC:G-tel (circles). (b) Excitation spectra of complexes AIC:G-c-Myc (solid line), AIC:G-tel (triangles), and free AIC in 95% ethanol solution (dashed line). Fluorescence intensity I_{550} registered at 320 nm was normalized to 1. The numbers of bound AIC molecules per G-c-Myc and G-tel were r = 1.

**Figure 4: Fluorescence of AIC:G-quadruplex complexes.** (a) The fluorescence spectra of free AIC in 95% ethanol solution (dashed line), AIC bound to Pu18 G-c-Myc (solid line) and to G-tel (triangles). The intensity of fluorescence was registered after excitation at 320 nm and normalized to 1. The parameter r was ~1 for all complexes. Inset: fluorescence of AIC:G-quadruplex complexes depending on r. Complexes AIC:G-c-Myc (triangles), AIC:G-tel (circles). (b) Excitation spectra of complexes AIC:G-c-Myc (solid line), AIC:G-tel (triangles), and free AIC in 95% ethanol solution (dashed line). Fluorescence intensity I_{550} registered at 320 nm was normalized to 1. The numbers of bound AIC molecules per G-c-Myc and G-tel were r = 1.

**Table 1: Binding parameters of AIC:DNA complexes.**

| DNA        | [a]K_1 (×10^{-6}), M^{-1} | [b]n_1 | [a]K_2 (×10^{-6}), M^{-1} | n_2 |
|------------|-----------------------------|--------|-----------------------------|-----|
| G-c-Myc    | 70 ± 10                     | 1      | 4 ± 1                       | 2   |
| G-tel      | 6.5 ± 1.0                   | 1      | 0.33 ± 0.05                 | 1   |
| c-Myc duplex| 0.8 ± 0.1                   | 5      | —                           | —   |

[a]: K_1, K_2 are the association constants of AIC:DNA complexes; [b]: n_1, n_2 are the maximal numbers of AIC molecules bound to a particular DNA.

G-tel to bound AIC molecules. Because the unbound AIC molecules fluoresce only negligibly in aqueous solution, the excitation spectra of free AIC in ethanol were taken for comparison with the respective spectra of complexes AIC:G-c-Myc and AIC:G-tel. The excitation spectra of AIC:G-c-Myc, AIC:G-tel, and the free drug in ethanol are shown in Figure 4(b). The fluorescence intensity I_{550} of AIC was registered upon excitation in the region 240 < λ < 340 nm. All spectra were normalized with respect to the fluorescence intensity I_{550} corresponding to excitation at λ = 320 nm. The occupancy value r for AIC:G-c-Myc and AIC:G-tel was ~1, that is, the AIC concentration was low enough for predominant formation of strong AIC:G-quadruplex complexes. Virtually no excitation signal of free AIC was detectable at ~260 nm (Figure 4(b), dashed line). The excitation spectra of AIC:G-quadruplexes demonstrated a markedly increased fluorescence at 250 < λ < 280 nm, the region of nucleotide absorption (Figure 4(b), bold curve and triangles). The signal in the region 250 < λ < 280 nm in the excitation spectra of AIC:G-quadruplexes indicated the energy transfer from the bases of G-quadruplexes to AIC. This effect is known to occur if a planar molecule is involved in stacking interactions with the adjacent bases. In contrast, no energy transfer is detectable when fluorescent molecules bind to the DNA grooves [20, 26]. Thus, the energy transfer data supported the stacking interaction of AIC with nucleotides, possibly involving an end stacking to terminal G-quartets in G-c-Myc and G-tel.

3.5. **Circular Dichroism Spectra of AIC:G-c-Myc Complexes.** We detected no induced CD signal in the wavelength region of AIC absorption upon binding of the drug to the oligonucleotides studied. To monitor the changes of the Pu18 G-c-Myc conformation upon AIC binding, we registered CD spectra of the complexes in the wavelength region of nucleotide absorption (220 < λ < 320 nm) (Figure 5). The parameters of CD spectra proved that free G-c-Myc adopted the structure of a parallel Pu18 G-quadruplex [6]. The CD spectra of G-c-Myc changed upon binding to AIC. The magnitude of the positive CD band at λ = 265 nm decreased after the addition of the drug. At maximal occupancy, r = 3, the magnitude of the band decreased by ~25%. These results may be interpreted as a reduction of stacking contacts between G-quartets. Nevertheless, the parallel conformation of the quadruplex remained intact upon drug binding.

We performed CD experiments to reveal whether AIC binding induces c-Myc duplex dissociation followed by G-quadruplex formation. The CD spectrum of the sample (Figure 5(b), open circles) is typical for a double-stranded oligonucleotide rich in G-runs. Only small changes around 280–290 nm occurred upon binding of 3 AIC molecules per duplex (Figure 5(b), filled circles). The excess of AIC (12 molecules per duplex) did not affect the CD spectrum of the oligonucleotide, however, a positive CD signal above 320 nm was detectable (Figure 5(b), filled squares). This induced CD band may reflect the interactions of AIC molecules with the quadruplex.
3.6. Downregulation of c-Myc Expression by AIC. To reveal whether AIC, a strong G-c-Myc ligand (see above), can modulate the steady-state level of c-Myc mRNA in living cells, we analyzed c-Myc transcripts in HCT116 human colon carcinoma cell line by real-time PCR after reverse transcription. As shown in Figure 6, the amounts of c-Myc mRNA decreased in cells treated with 1 μM of AIC for 24–48 h. By 24 h the amount of c-Myc mRNA was 4–6 times smaller than that in untreated cells. The decreased level of c-Myc transcripts remained sustained for up to 48 h of exposure (Figure 6). In contrast, no effect on in vitro telomerase activity was detectable even at concentrations of AIC up to 10 μM (data not shown). These results are in line with preferential affinity of AIC to G-c-Myc rather than to G-tel.

4. Discussion

Previously we have shown a high potency of AIC in a panel of human tumor cell lines and in vivo tumor models [22]. The duplex DNA, in particular, DNA breaks have been identified as intracellular AIC targets important for cytotoxicity of this indolocarbazole derivative. In the present study we identified G-c-Myc as a new selective target of AIC. The high affinity of AIC to G-c-Myc ($K_1 = 7 \pm 1 \cdot 10^7 \text{M}^{-1}$) was ~100 times greater than that to the c-Myc derived duplex DNA. AIC demonstrated a ~10-fold preference for the parallel G-c-Myc conformation over the antiparallel G-tel (Table 1). Accordingly, treatment of HCT116 cells with 1 μM of AIC attenuated the amount of c-Myc mRNA whereas the in vitro telomerase activity remained unaltered even by 10 μM of the drug.

Design of therapeutic G-quadruplex binding drugs may be based on differential structure of individual tetraplex DNAs and/or on simultaneous targeting of quadruplexes by a particular ligand. A structural preference for the parallel c-Myc or c-Kit quadruplexes over the antiparallel G-tel has been reported for several ligands [14, 27, 28]. Taking advantage of different degrees of structural plasticity of parallel and antiparallel quadruplexes in binding ligands, Garner et al. have identified an extended heteroaromatic 1,4-triazole (TRZ) as a compound with some selectivity for parallel folds whereas the polycyclic fluorinated acridinium cation RHPS4 showed selectivity for antiparallel conformations [29]. However, RHPS4 and TRZ were similarly
potent telomerase inhibitors [29]. Interestingly, the porphyrin derivative 5,10,15,20-tetrakis(1-methyl-4-pyridyl)-21H, 23H-porphine (TMPyP4) demonstrated the properties similar to AIC. The binding parameters of TMPyP4 to G-c-Myc at ionic strength $K^+ = 0.1 \text{M}$ ($K_1 = 7 \pm 1 \times 10^7 \text{M}^{-1}$, $n_1 = 1$ and $K_2 = 1 \times 10^6 \text{M}^{-1}$, $n_2 = 2$) [28] were close to AIC binding to G-c-Myc Pu18 in Hanks’ buffer (Table 1). Furthermore, TMPyP4 preferentially bound to G-c-Myc rather than to c-Myc duplex DNA [28]. Tight TMPyP4 binding sites were suggested to be at the terminal G-quartets while the weaker binding sites were located at the nucleotide loops [6, 27]. Our data suggest that binding of AIC to G-c-Myc may occur via similar sites. The signal in the region 250 < $\lambda$ < 280 nm of the excitation spectra of AIC:G-c-Myc complexes indicated energy transfer from the bases of G-quadruplexes to AIC. Thus, the energy transfer data supported the mode of AIC:nucleotide interactions, most likely involving an end stacking to terminal G-quartets in G-c-Myc and G-tel.

Although much of the structural work on G-quadruplexes has been done in vitro, evidence is growing to prove the roles of these structures in vivo [30, 31]. The formation of G-quadruplexes is facilitated by negative superhelical stress produced during transcription and can be stabilized by monovalent cations such as $K^+$ and $Na^+$ that intercalate between the G-tetrads and coordinate the bonds with guanine carbonyl groups. Transcription-associated supercoiling has been demonstrated to evoke dynamic effects on DNA conformation in the c-Myc regulatory region: the duplex DNA can be converted into nonduplex DNA, even at considerable distances from the transcriptional start site [32]. These nonduplex DNA structures that control both transcriptional switch on/off and the rate of transcription firing, are amenable to targeting by small molecular weight ligands. The fact that AIC is a strong ligand not only for c-Myc-derived G-quadruplex but also for c-Myc duplex (withstanding the preference of the former conformational structure) implies that this ligand might be relevant for treatment of c-Myc positive tumors due to the ability to interact with various regions within this oncogene. One can hypothesize that the formation of drug-DNA complexes at multiple sites is linked to a decrease of c-Myc mRNA in AIC-treated cells (see below).

The conformational transition from a duplex to a G-quadruplex within the c-Myc promoter has been proposed as a mechanism of c-Myc inhibition by TMPyP4 [27, 28]. Accordingly, the high selectivity of AIC for G-c-Myc may lead to prevalence of the quadruplex over the duplex DNA conformation in the cell. We believe that the formation of high affinity complexes AIC:G-c-Myc is likely to be a prerequisite for downregulation of c-Myc expression. Still, drug-quadruplex interaction may not be the sole mechanism of c-Myc silencing by AIC. Other events including the interference with the formation of the transcriptional complexes and/or the traverse of the transcriptional machinery along the DNA template, as well as deregulation of mRNA stability, cannot be ruled out at this stage of the study. Moreover, dual targeting by a quadruplex ligand might be therapeutically beneficial. Indeed, the effect of the G-c-Myc ligand quarfloxin involves disruption of the inter- action between ribosomal DNA G-quadruplexes and nucleolin [11, 13]. Gunaratnam et al. have identified a naphthalene diimide derivative as a compound that recognized both G-tel DNA and the quadruplex sites in the c-Kit promoter. This compound inhibited telomerase activity and suppressed c-Kit mRNA and protein expression, the effects associated with the induction of growth arrest in a patient-derived gastrointestinal stromal tumor cell line [33]. Whatever the mechanism(s), the identification of AIC as a preferred binder to c-Myc promoter-derived G-quadruplex may expand the therapeutic perspective of indolocarbazole derivatives. Together with the confirmed ability to target some protein kinases and dsDNA, these compounds emerge as high affinity ligands for noncanonical DNA structures and potential modulators of gene expression, tentatively via DNA conformation-dependent regulatory mechanisms.

Acknowledgments

The authors thank M.A. Livshits for helpful discussions. This work was supported by Grant 08-04-12104-ofi of Russian Foundation for Basic Research. A. A. Shtil and O. F. Borisova share equal senior authorship.

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