INSIGHTS INTO THE ACTION MECHANISM OF AZACYANINES: THEIR TOPOISOMERASE IIα INHIBITION POTENTIAL AND NUCLEIC ACID SELECTIVITY

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INSIGHTS INTO THE ACTION MECHANISM OF AZACYANINES:
THEIR TOPOISOMERASE IIα INHIBITION POTENTIAL AND NUCLEIC ACID SELECTIVITY

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INSIGHTS INTO THE ACTION MECHANISM OF AZACYANINES: THEIR TOPOISOMERASE IIα INHIBITION POTENTIAL AND NUCLEIC ACID SELECTIVITY

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Topoisomerase II alpha (Topo IIα) is one of the essential enzymes in cell viability. It is required for cell cycle progression due to its role in regulating the topological constraints during DNA replication and transcription. Due to its role, Topo IIα became one of the extensively exploited targets in chemotherapy, and there is an on-going research in design of small molecules targeting topoisomerase’s catalytic activity at different stages using different mechanisms.

Within the scope of this thesis first, the effects of five azacyanine derivatives on Topoisomerase IIα enzyme (Topo IIα) at molecular level have been assessed by using an in-vitro Topoisomerase Drug Screening kit. Three of the azacyanines (Azamethyl, Azaethyl and Azaisobutyl) used were benzimidazole derivatives differing from each
other in terms of the alkyl chain length on the benzimidazole ring and two of them were benzothiazoles (Aza4 and Aza5). Second, the affinity and selectivity of these compounds towards different nucleic acid sequences and structures have been investigated by using competition dialysis method. Our primary goal was to accentuate azacyanines as probable Topo IIα inhibitors, and provide some mechanistic explanations for their effects.

Our results revealed that three of the five azacyanine molecules investigated in here decreases the catalytic activity of Topo IIα. Detailed investigation of the effect of Azamethyl on Topo IIα revealed that the effect on Topo IIα was concentration dependent, such that the higher Azamethyl concentrations led to stronger effect as determined by the intensity of the DNA bands in ethidium bromide (EtBr) stained agarose gels. Moreover, when used at the same concentrations (50.0 µM), Azamethyl showed stronger effect than Etoposide which is a well-known Topo IIα inhibitor being used in chemotherapy. Even better, the efficiency of 50.0 µM Azamethyl was higher than 500.0 µM Etoposide.

To shed light onto the mechanism of Topo IIα catalytic activity change, we further investigated the affinity and selectivity of azacyanines towards different nucleic acid sequences and structures via competition dialysis. We included three more azacyanines (Azapropyl, Azabutyl and Azaisopropyl) in our competition dialysis setup to understand the effect of the chain length and branching on the benzimidazole ring in nucleic acid binding affinity and selectivity. Our results revealed that the azacyanines were highly selective towards triple helical nucleic acid structure poly(dA).[poly(dT)]₂. More importantly, their affinity and selectivity towards poly(dA).[poly(dT)]₃ was decreasing with the increasing linear alkyl chain length, and increasing branching. The order of affinity was Azamethyl > Azaethyl > Azapropyl > Azabutyl > Azaisobutyl > Azaisopropyl.

Altogether, the findings presented in here demonstrate the potential of azacyanines to inhibit Topo IIα’s catalytic activity.
**Keywords:** Azacyanine, Catalytic Activity Change, Topoisomerase II Alpha, Selectivity, Nucleic Acid Structure, Competition Dialysis.
ÖZ

AZASIYANINLERIN ETKİ MEKANİZMASI:
TOPOİZOMERAZ IIα ENGELLEME POTANSİYELİ
VE
NÜKLEİK ASİT SEÇİCİLİĞİ

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Topoizomeraz II alfa (Topo IIα) hücre yaşamında görev alan temel enzimlerden birisidir. DNA replikasyonu ve transkripsiyonu sırasında DNA’nın topolojik kısıtlamalarının çözülmesini sağladığı için hücre döngüsü ilerleyişinde gerekli bir enzimdir. Bu rolü sebebi ile Topo IIα kemoterapide sıkça kullanılan bir hedef ve topoizomeraz enziminin katalitik aktivitesini değişik mekanizmalar ile farklı adımlarda etkileyebilecek küçük moleküllerin dizaynı konusunda devam eden birçok bilimsel çalışma bulunmaktadır.

Bu tez kapsamında, ilk olarak beş farklı azasiyanın türevinin Topo IIα üzerindeki etkisi moleküler düzeyde in-vitro Topoizomeraz tarama kiti ile incelenmiştir. Kullanılan üç azasiyanın molekülü (Azametil, Azaetil ve Azaisobütil)

Elde ettiği sonuçlar, inceленen bu beş azasiyanın molekülünden üçünün Topo IIα enziminin katalitik aktivitesini azalttığını göstermiştir. Ayrıntılı incelemelerde, etidyum bromür (EtBr) boyalı agaroz jellerindeki bantların yoğunluğu bakılarak Azametil molekülünün kullanılan konsantrasyona bağlı olarak TopoIIα enziminin katalitik aktivitesine negatif olarak etkisi görülmüştür. Özellikle aynı konsantrasyonlarda (50.0 µM) kullanıldığında Azametil molekülünün Etoposid molekülüne göre çok daha etkili olduğu saptanmıştır. Etoposid molekülü kemoterapide kullanılan ve etkisi kanıtlanmış bir Topo IIα inhibitörüdür. Daha da önemlisi, 50.0 µM Azametil molekülünün 500.0 µM Etoposid molekülünden bile daha etkili bir şekilde Topo IIα’nın katalitik aktivitesini düşürdüğü gözlemlenmiştir.

Topo IIα’nın katalitik aktivitesinin değişmesine ışık tutmak amacı ile azasiyanın molekülerinin farklı nükleik asit sekanslarına ve yapılarına karşı olan ilgişi ve seçiciliği rekabetçi diyaliz yöntemi kullanılarak araştırılmıştır. Daha önce çalışılmamış üç azasiyanın molekülü (Azapropil, Azabütil ve Azaizopropil) de zincir uzunluğunun ve halka yapısının nükleik asitlere bağlanmadaki etkisini incelemek üzerine rekabetçi diyaliz dengesine eklenmiştir. Sonuçlarımız azasiyanın molekülerinin üçlü yapı sarmalı yapısı oluşturulan poly(dA).[poly(dT)]₂ nükleik asit yapısına yüksek afinitite ve seçicilik ile bağlandığıını göstermiştir. Daha da önemlisi bu ilgi ve seçicilik benzimidazol halka yapısındaki artan zincir uzunluğu ve zincirin dallanma özelliğine göre azalma göstermiştir. Molekülerin poly(dA).[poly(dT)]₂ üçlü DNA yapısına karşı olan ilgileri aşağıdaki sıraya göre gerçekleşmiştir. Azametil > Azaetil > Azapropil > Azabütil > Azaisobütil > Azaisopropil.
Burada sunulan bulgular azasiyanın moleküllerinin muhtemel Topo IIα katalitik inhibitörü olabilme potansiyellerine ışık tutmaktadır.

**Anahtar kelimeler:** Azasiyanın, Katalitik Aktivite Değişikliği, Topoizomeraz II Alfa, Sıkı, Nükleik Asit Yapı, Rekabet Diyaliz.
TO

SCIENCE

&

HUMANITY
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<thead>
<tr>
<th>Abbreviation</th>
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<tbody>
<tr>
<td>ANOVA</td>
<td>Analysis of variance</td>
</tr>
<tr>
<td>ATM</td>
<td>Ataxia telangiectasia mutated</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>ATR</td>
<td>ATM Rad3 related</td>
</tr>
<tr>
<td>Aza4</td>
<td>Azacyanine 4</td>
</tr>
<tr>
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</tr>
<tr>
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<td>Azacyanine butyl</td>
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<tr>
<td>Azaethyl</td>
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<tr>
<td>Azaisobutyl</td>
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</tr>
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</tr>
<tr>
<td>Azapropyl</td>
<td>Azacyanine propyl</td>
</tr>
<tr>
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<td>Buffered phosphate EDTA with sodium chloride</td>
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<tr>
<td>bp</td>
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<tr>
<td>c-Myc</td>
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</tr>
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<tr>
<td>EDTA</td>
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</tr>
<tr>
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</tr>
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<td>g</td>
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</tr>
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CHAPTER 1

INTRODUCTION

1.1. DNA Structures:

Deoxyribonucleic acid (DNA), which is the carrier of genetic information, plays a central role in life. It is the main player in replication and transcription, and therefore responsible for the development and maintenance of life. DNA that is mainly condensed inside the nucleus is mostly in B form, Watson-Crick DNA. It is made up of two anti-parallel polynucleotide strands that wind around each other with a right-handed twist, forming the double helical structure. In double helical structure, the bases occupy the inside of the helix and the sugar-phosphate chains face to the outside. Its conformation provides a direct and easy access to replication and transcription related proteins like polymerases. Furthermore, B-form DNA is the most compatible form for packaging the long polymer inside the nucleus due to its tight bending capability. Its stability, flexibility and dynamic conformation are thought to be the main reasons behind the selection of DNA as the genetic material instead of ribonucleic acid (RNA). (Travers & Muskhelishvili, 2015).

After the discovery of right-handed double helical structure of B-form DNA by Watson and Crick in 1953 (Watson & Crick, 1953) there have been many revolutions in our understanding of the DNA structure and function.

Besides B-form DNA, other kinds of DNA structures such as A-DNA, Z-DNA, i-motifs, hairpins, bulges, stem-loops, pseudoknots, cruciforms, triplexes, and G-
quadruplexes have been discovered (Bachurin, Kletskii, Burov, & Kurbatov, 2018) (Figure 1). All of these structures that mostly have non-canonical (non-Watson-Crick) base pairing arrangements are found to play distinctive crucial roles in cell cycle regulation. Especially triple helical (triplex) and quadruplex structures received great attention in the last two decades due to their likely roles in gene expression and uncontrolled cell division associated with formation of cancerous tumors (H. Han & Hurley, 2000; Van Dyke & Nelson, 2013). It has been shown that triplex and quadruplex structures also have essential roles such as protecting DNA from degradation and keeping promoters silent (Rigo, Palumbo, & Sissi, 2017; Toscano-Garibay & Aquino-Jarquin, 2014).

![Figure 1](image_url)

**Figure 1.** DNA structures observed in the cell. a. B-form DNA and Z-form DNA. b. Cruciform. c. Triplex. (Bochman, Paeschke, & Zakian, 2012). Reprinted from (*Nat. Rev. Genet.*), Vol.13, Copyright (2012) with permission from (Springer Nature).
Triplex structures are generally formed by the placement of a third strand in the major groove of the double helical structure. Four stranded G-quadruplex structure are generally formed by the association of guanine rich strands. (Bissler, 2007; Gacy et al., 1998; Hanahan & Weinberg, 2011).

1.1.1. Triplex DNA and DNA/RNA Hybrids:

Although triple helical structures are known for a long time, it took quite some time to realize their importance. In triple helix formations basically a third strand, so called triplex-forming oligonucleotide (TFO), binds to a duplex DNA structure via non-canonical Hoogsteen base pairing (Frank-Kamenetskii & Mirkin, 1995). Triple helical structures are mostly classified as purine motifs or pyrimidine motifs. In a purine motif, purine rich TFO binds to the purine strand of the duplex via reverse-Hoogsteen base pairing in an antiparallel orientation. On the other hand, in the pyrimidine motif, pyrimidine rich TFO binds to the purine strand of the duplex via Hoogsteen base pairing in a parallel orientation (Bing et al., 2017). DNA itself and RNAs such as long non-coding RNAs, promoter-associated RNAs, promoter inhibiting RNAs, promoter RNAs, small interfering RNAs, and micro RNAs are all considered as potential TFOs (Toscano-Garibay & Aquino-Jarquin, 2014) (Figure 2).
Figure 2. Possible triplex structures in the cell. (Toscano-Garibay & Aquino-Jarquin, 2014). Reprinted from (Biochim. Biophys. Acta), Vol.1839, Copyright (2014) with permission from (Elsevier).

Triplex forming oligonucleotides gained great attention in the last two decades with respect to controlling gene expression. Oligonucleotides can be used for controlling gene expression via antisense approach, which is composed of different strategies such as antisense and antigene strategies (Helene, Thuong, & Harel, 1992; Rakoczy, 2001). In the antisense strategy, oligonucleotides can be used to target messenger RNAs therefore downregulating protein production. On the other hand, in the anti gene strategy, oligonucleotides can be used as triplex forming oligos (TFOs) by binding specific sequences on DNA thereby inhibiting the transcription of the gene (Helene et al., 1992; Rakoczy, 2001). As for controlling the gene expression, it has been shown that triple helices are related to gene silencing (Toscano-Garibay & Aquino-Jarquin, 2014), DNA damage, and DNA repair mechanisms (Ni et al., 2018). Therefore, triplex formation can be used in silencing the oncogenes to prevent tumor growth.
Furthermore, there are also several in-vitro studies, published recently, showing the emerging roles of triplex nucleic acid structures in various areas such as biosensors, sequence specific labeling, DNA based nanostructures, and RNA biology (Chen et al., 2015; Conrad, 2014; M. S. Han, Lytton-Jean, & Mirkin, 2006; Hu, Cecconello, Idili, Ricci, & Willner, 2017).

Nevertheless, triplex structures are generally unstable compared to their corresponding duplex structures because of the weaker Hoogsteen base pairing. Since their stability is essential in impeding gene expression and/or regulation, stabilizing such structures has become a major focus in rational drug design research. One way to increase the stability of these structures is to design small molecules that tightly and selectively bind to them. Despite many different types of stabilizing small molecules present there is still a need for better and more effective triplex binding molecules (Chaires et al., 2003; Escudé et al., 1998; M. S. Han et al., 2006).

On the other hand, the formation of stable triplex sequences in-vivo could be harmful for normal cells. It has been shown that polypurine-polypyrimidine mirror repeats (triplex forming sequences) could give rise to various diseases by causing mutations or altering gene expression (Bissler, 2007). There are a number of inherited as well as acquired diseases associated with the potential formation of stable triple helical structures due to the presence of trinucleotide repeats such as Friedreich’s ataxia, Tuberosclerosis complex, Autosomal dominant polycystic kidney disease, and Follicular lymphoma (Bissler, 2007).

Among these diseases Friedreich’s ataxia, is an autosomal recessive disease which exhibits progressive degeneration of nerve tissues located in the spinal cord and nerves in the legs and arms. The gene, responsible for this disease, is located on chromosome 9 and normally contains 7-22 GAA repeats. However, in Friedreich’s ataxia patients number of repeats increases abruptly, to hundreds or thousands. These increased numbers of repeats are in return cause the formation of hinged DNA (H-DNA) that interferes with the gene expression. This process gives rise to reduced
number of frataxin protein which is a mitochondrial protein that controls cellular iron homeostasis. Reduced number of frataxin, then, causes reduced muscle activity (Bissler, 2007; Frank-Kamenetskii & Mirkin, 1995).

1.1.2. Quadruplex DNA Structures:

It is known that the formation of quadruplex structures is dependent on guanine nucleotides. Planar arrangement of four guanine bases forms G-quartets, where the stacking of these G-quartets results in the formation of the G-quadruplex structures (Bing et al., 2017). There are structurally different quadruplexes based on the number of guanine bases and the number of strands involved in the formation of the structure (Figure 3).

![Figure 3. Different forms of G-quadruplexes.](Harkness & Mittermaier, 2017). Reprinted from (Biochim. Biophys. Acta), Vol.1865, Copyright (2017) with permission from (Elsevier).
The formation of G-quadruplex structures in-vitro is well established and the propensity of guanine rich sequences to form G-quadruplex structures in-vivo is well accepted. Genome-wide bioinformatic analyses revealed that the guanine rich sequences are particularly present at the end of the chromosomes, telomeres, and in promoter regions such as the ones seen in c-myc gene (Rigo et al., 2017) (Figure 4).

Figure 4. Possible locations of G-quadruplex structures in the cell. (Rigo et al., 2017). Reprinted from (Biochim. Biophys. Acta), Vol.1861, Copyright (2017) with permission from (Elsevier).

Guanine rich sequences are believed to play very important roles in regulation of many cellular functions such as telomere maintenance, transcription regulation, and translational & post-translational regulation of RNAs (Bing et al., 2017). For instance, roles of guanine rich sequences at the end of the telomeres include prevention of
chromosome fusions which would otherwise give rise to genomic instability and prevention of degradation of chromosome ends. Furthermore, guanine rich sequences at transcription start sites are believed to control the gene expression (Rigo et al., 2017). Another function of guanine rich sequences appears in RNA biology. G-quadruplex structures on RNA molecules could have many roles such as translation regulation, 3’end formation, transcription termination, RNA localization, splicing regulation, and ribonucleoprotein formation (Fay, Lyons, & Ivanov, 2017).

Due their possible roles in gene expression and gene regulation, guanine rich sequences are one of the most sought after structures in cancer treatment. For instance, it is proposed that driving the formation of quadruplex structures and stabilizing them in guanine rich oncogenes such as c-Myc, could be enough to halt the over-expression of these genes in cancer cells (Siddiqui-Jain, Grand, Bearss, & Hurley, 2002).

1.1.3. Targeting DNA by Using Small Molecules:

Nucleic acids play essential roles in various biological processes like gene storage, transcription, replication etc. Therefore, one can regulate many biological processes by targeting the nucleic acid structures in-vivo. This approach forms the basis of the most conventional drug design studies (M. Wang, Yu, Liang, Lu, & Zhang, 2016). Nucleic acids are the most preferred targets compared to proteins and protein-like factors mainly due to their upstream roles. Additionally, the existence of alternative nucleic acid structures, namely duplex, triplex, and quadruplex makes them desirable targets for various drugs in the treatment of several diseases (M. Wang et al., 2016).

Wang et al. summarized the importance and the recent advances in developing small molecules that target nucleic acid structures. Small molecules could interfere with biological processes by binding to the nucleic acid structures, and altering either their structure or their interactions with other proteins and/or nucleic acid structures. For instance, the binding of a small molecule to a DNA sequence could block the enzyme binding, such as topoisomerase or DNA polymerase, to that very same sequence by blocking the recognition of the sequence by these proteins due to a
change triggered in DNA’s conformation upon small molecule binding (M. Wang et al., 2016).

There are numerous examples of small molecules targeting duplex DNA using different modes of binding. For instance, psoralen, aflatoxin B, N^4C-ethyl-N^4C, mitomycin C, and N^2G-ethyl-N^2G target DNA duplex by forming covalent bonds with the DNA structure. On the other hand, while netropsin, distamycin, DAPI, berenil, pentamidine, and Hoechst 33258 are targeting DNA duplex by binding to the minor groove, neomycin, nogalamycin, and neocarzinostatins are targeting DNA duplex by binding to the major groove via non-covalent interactions. Additionally, daunomycin, adriamycin, ditercalinium, and cryptolepine which are known as intercalators are also targeting DNA duplex via non-covalent interactions, mainly by stacking between base pairs. Furthermore, there are molecules targeting DNA duplex via multiple binding modes such as PBD-BIMZ and PBD-naphthalimide (M. Wang et al., 2016).

On the other hand, there are also numerous studies targeting triplex nucleic acid structures. For example, benzopyridoindole derivatives such as benzo[e]pyridoindole (BePI) and benzo[g]pyridoindole (BgPI) have been demonstrated as successful triplex stabilizing small molecules. Moreover, coralyne, a berberine alkaloid, is a well-known natural product that stabilizes triplex structures. Other studies also showed the binding and selectivity of naphtylquinoline, bis-amidoantraquinones, and amidoglycosides towards triplex DNA structures (Fox & Darby, 2004). Additionally, neomycin-pyrene computational modelling also demonstrated that these small molecules target triplex DNA structures through both groove binding and intercalation (M. Wang et al., 2016).

Quadruplex DNA structures are also targets for small molecules as mentioned before. For instance, distamycin A, daunomycin, RHPS4, Tmpyp4, Telomestatin, and BRACO 19 targets quadruplex structures (M. Wang et al., 2016). Moreover, it had been recently shown that newly synthesized small molecule- CX-5461, which is a
polymerase I derived rRNA inhibitor, has the ability of stabilizing G-quadruplex structures on DNA (Xu et al., 2017).

1.2. Topoisomerases:

Throughout DNA transcription, replication, and recombination several proteins and enzymes play crucial roles. During these processes unwinding DNA is a prerequisite for proteins to enter to the site. However, unwinding DNA subsequently creates some topological problems since unwinding one site would give rise to overwinding elsewhere on the DNA. This topological problem is solved by topoisomerase enzymes. They are responsible from linearizing and/or releasing the torsion on the DNA. By doing so other proteins can enter to the site and required process may continue. Linearizing DNA by topoisomerases, however, is a very complex mechanism involving DNA cleavage, DNA relaxation, and finally religation of the phosphate backbone in order to restore the integrity of the DNA continuity (Winkler, 2011). Detailed molecular mechanisms are given in the following sections.

1.2.1. Classification of Topoisomerases:

Topoisomerases are classified mainly into two groups as Type I (Topo I) and Type II (Topo II) based on their ability to cleave only one or both of the DNA strands respectively. In other words, while Type I topoisomerases introduce single strand breaks, Type II topoisomerases introduce double strand breaks (Champoux, 2001; J. C. Wang, 1998). Odd numbered enzymes (e.g. “I” or “III”) generally fall into type I topoisomerases, whereas evenly numbered enzymes (e.g. “II” or “IV”) generally fall into type II topoisomerases. There are also many sub classifications of topoisomerases where letters “A”, “B”, or “C” indicate the subtype. Subtypes usually differ from each other in amino acid sequence, structure, and/or function (Vos, Tretter, Schmidt, & Berger, 2011).

There are six functionally different topoisomerases encoded by the human genome which are organized into three main groups; type I (type Ia and type Ib), type II (type
IIa and type IIb), and type III (type IIIa and type IIIb) (Pommier, 2013). All of these enzymes have distinctive roles in the cell. Moreover, while topoisomerase type I and type II enzymes are abundant in the nucleus, topoisomerase type III enzyme is abundant in mitochondria. The distinctive roles of Topo I and Topo II enzymes in the cell are explained in detail below.

1.2.2. **Topoisomerase I:**

Type I topoisomerase enzyme helps fork movement during relaxation of DNA and all higher eukaryotes contain at least one type I topoisomerase enzyme. Type I topoisomerases are further divided into two sub-families; type Ia which relaxes only the negatively supercoiled DNA and type Ib which relaxes both the negatively and positively supercoiled DNA (Goto, Laipis, & Wang, 1984; Liu & Miller, 1981). Mammalian cells have type Ib sub-family. Type Ib topoisomerases share no homology with the other type of topoisomerases and are functionally distinct from type Ia topoisomerases (Uday Bhanu & Kondapi, 2010).

Topoisomerase I (Topo I) has essential roles for the normal development of the cell (Morham, Kluckman, Voulomanos, & Smithies, 1996). It simply relaxes supercoiled DNA thus ease the entrance of other enzymes and/or proteins into the required site. Unless Topo I works properly, helical constraints of the supercoiled DNA may hinder DNA replication or/and transcription processes (Gilmour, Pflugfelder, Wang, & Lis, 1986; J. Wu, Phatnani, Hsieh, & Greenleaf, 2010). It has been shown that inactivation of Topo I reduced the transcription rate of *Saccharomyces cerevisiae* and disturbed the development of *Drosophila melanogaster* (Uday Bhanu & Kondapi, 2010).

The working principle of Topo I is illustrated in Figure 5. It is based on trans-esterification reaction that occurs between the tyrosine residue located in the topoisomerase’s active site (tyrosine 723 in humans) and 3’ phosphodiester backbone of the DNA. Then rotation of free 5’ hydroxyl end over the intact DNA strand takes place. After the relaxation of the duplex, 5’ hydroxyl end acts as a nucleophile and reverse trans-esterification occurs towards the phosphotyrosil bond. Topo I is liberated
after DNA re-ligation and catalyzes another nicking, closing, relaxation cycle elsewhere in the genome (Pommier & Cherfils, 2005). After relaxation, DNA or RNA polymerases will continue with replication or transcription (Barros et al., 2013; Uday Bhanu & Kondapi, 2010).

Figure 5. Working principle of Topoisomerase I (Lodisch et al., 2000).

However, despite the importance in relieving the torsional stress, Topo I also possesses serious dangers to the cells. It can trigger genomic instability when encounters with a non-B DNA structure or might interfere with continuing gene expression. As a result of interference, negatively supercoiled strands might accumulate at one site and eventually promote the formation of R-loops and/or stable non-B DNA structures like quadruplexes. This action, at the end, has the capability of interfering with DNA replication and provide highly toxic double strand breaks (N. Kim & Jinks-
Robertson, 2017). Thus, targeting Topo I and/or interfering with its activity can be used in the treatment regime of cancer to kill the malignant cells.

1.2.3. Topoisomerase II:

Topoisomerase II enzymes (Topo II) are believed to play a role in DNA replication, chromosome condensation, chromosome segregation, transcription, DNA repair, and neuronal activities although some of these roles are controversial (J. Nitiss, 2009; Vos et al., 2011). In mammalian cells, there are two Topo II isozymes (type IIα and type IIb) while in other eukaryotic cells there is only one Topo II (J. L. Nitiss, 2009).

As mentioned before, in order to solve the topological problems in DNA, introduction of DNA strand breaks is obligatory. Topoisomerases provide a safe way for DNA breaks by protecting the cut edges of the breaks. Topo II’s function seems more important in this aspect because it creates double strand breaks while Topo I creates single strand breaks, which can be compensated by other strand of the DNA. If the cut site is not protected by the enzyme after Topo II cleavage, mutations might arise due to the unrepaired double strand break. Topoisomerase II enzymes function in two different ways; decatenation of catenated DNA and/or relaxation of supercoiled DNA (Figure 6).
1.2.4. Working Principle of Topoisomerase II:

Topoisomerase II enzymes are homodimeric where each subunit breaks one DNA strand to create a double strand break. As illustrated in Figure 7, Topo II interacts with two strands of DNA to complete the passage. It introduces a double stranded break in one DNA strand which is called as G segment (gate segment), and then passes the other DNA strand, which is called the T segment, through the break. T segment is passed through the G segment by the enzyme via ATP dependent reaction. Then the break in G segment is resealed (J. Nitiss, 2009).

Enzyme cleavage of G strand is achieved in the presence of Mg$^{2+}$ by forming a phosphotyrosine linkage between each of the DNA single strands. On the other hand, it produces a free tyrosine in each subunit of the enzyme. Then a closed clamp is formed by ATP binding to the enzyme. This closed clamp captures T segment and passes it through the break in the G segment. After the passage, T segment leaves the enzyme through the carboxy terminus. ATP hydrolysis occurs twice during the whole process. First ATP is used to assist in strand passage. The second ATP is used to re-open the
clamp. G segment may be released after the process or, if it is necessary, enzyme may go through another catalytic cycle without dissociating the G segment (Figure 7). This mechanism of Topo II provides some advantages like protection of DNA ends of the break and the ability of quick and efficient religation of DNA breaks (J. Nitiss, 2009).

**Figure 7.** Working principle of Topoisomerase II (J. Nitiss, 2009). Reprinted from *Nat. Rev. Cancer*, Vol.9, Copyright (2009) with permission from (Springer Nature).

**1.2.5. Biological Functions of Topoisomerase II:**

As mentioned above, Topo II enzyme has a vital role in several processes during cell cycle and metabolism. The role of Topo II in some of these specific processes are discussed in detail below (Ferguson & Baguley, 1994; J. Nitiss, 2009; Vos et al., 2011).
1.2.5.1. Topoisomerase II During DNA Replication:

DNA replication is a semi-conservative process which occurs via unwinding of two DNA strands followed by the replication of each strand. In the absence of a functional topoisomerase in the replication fork, positively supercoiled DNA accumulates. The accumulation of DNA results in several topological problems and ultimately blocks the replication. Early stages of replication only require Topo I activity to relax the DNA. But in the later stages of replication when two replication forks impinge on each other, Topo I cannot fit into the replication site to relax the DNA. At this stage where two interlinked catenanes present, Topo II activity is required for decatenation of the dimer (J. Nitiss, 2009; Vos et al., 2011). Topo II action during the replication elongation includes catenation of pre-catenane structures (Lucas, Germe, Chevrier-Miller, & Hyrien, 2001).

Type II topoisomerases are composed of two subtypes; Topo IIα and Topo IIβ. It is believed that human Topo IIα has different functions than Topo IIβ and other type II topoisomerases at lower eukaryotic cells. It is observed that Topo II relaxes positively supercoiled DNA more actively than negatively supercoiled DNA (McClendon, Rodriguez, & Osheroff, 2005). Since it is the positively supercoiled DNA that is accumulating during the replication fork movement, Topo IIα is hypothesized to play a more crucial role than Topo IIβ. Yet, several studies have proven this hypothesis. Li et al. had shown that the phosphorylation of Topo IIα during S-phase is a prerequisite for the replication process (H Li, Wang, & Liu, 2008). Moreover, it has been shown that while Topo IIβ knockout cells can be recovered and continue with the replication, Topo IIα knocked down cells failed to continue with the replication (J. Nitiss, 2009)

1.2.5.2. Topoisomerase II During Chromosome Segregation:

During mitosis, duplicated chromosomes must be segregated from each other once the replication is completed. Topoisomerases aid during this process by compacting DNA and holding sister chromatids together during chromosome line up at
the center of the cell. Afterwards sister chromatids are separated from each other by topoisomerase action. Although catenated DNA is a problem for early replication process, at chromosome segregation step, catenated sister chromatid generation helps for proper chromosome lining at the center of the cell. DNA compaction via catenation, performed by topoisomerases, facilitates the chromosome segregation (J. Nitiss, 2009; Vos et al., 2011).

It was once believed that catenated DNAs were the sole reasons behind the staying of sister chromatids together. But it had been clearly shown that the specialized proteins called cohesins are the main players in keeping sister chromatids together (Diaz-Martinez, Gimenez-Abian, & Clarke, 2008; Losada & Hirano, 2005). It had been observed that even though the cohesins are essential for the sister chromatid cohesion, their absence is partly tolerated (Huang, Milutinovich, & Koshland, 2005; Michaelis, Ciosk, & Nasmyth, 1997). One possible explanation for this situation is the maintenance of cohesion with catenated DNA formation.

Another relation of topo II enzyme with the chromosome segregation is its association with the factors like aurora B kinase and the polo-like kinase (Plk-1) which are responsible for centromere formation and segregation (Coelho et al., 2008; H Li et al., 2008). Although these interactions are not well understood, co-localization of topo II with Plk-1 and aurora kinase B might be important for centromere resolution (Vos et al., 2011).

1.2.5.3. Topoisomerase II During Transcription:

During transcription, RNA polymerase action gives rise to localized positively supercoiled DNAs ahead of transcription bubble (Liu & Wang, 1987; H. Y. Wu, Shyy, Wang, & Liu, 1988) (Figure 8). If these positively supercoiled strands are left unchecked, stable R loop (RNA-DNA hybrids) formation might be observed (Marc Drolet, Bi, & Liu, 1994). Since the formation of such structures cause genomic instability, the cell growth will be impaired due to the formation of DNA breaks (M Drolet et al., 1995; Tuduri et al., 2009).
Figure 8. Topo II action during transcription (Vos et al., 2011). Reprinted from (Nat. Rev. Mol. Cell Biol.), Vol.12, Copyright (2011) with permission from (Springer Nature).

Although it has been shown that topo I enzymes work more actively at transcription site to remove the negative supercoils, a sub-type of topo II enzyme, topo IIβ has been observed at many promoter sites of highly transcribed genes (J. Nitiss, 2009; Vos et al., 2011). Ju et al. showed topo IIβ localization on the promoter of genes which were activated by nuclear hormone receptors by means of chromatin immunoprecipitation (ChIP) experiments (Ju et al., 2006; Ju & Rosenfeld, 2006). The finding in their studies was the co-localization of topo IIβ with DNA repair related proteins (PARP, Ku70/Ku86, and DNA dependent protein kinase) at the transcription initiation site. This type of localization and function is different than the function of topo IIβ in double stranded DNA breakage. It is believed that topo IIβ is recruited to promoters for its enzymatic function. Nevertheless, the absolute function and the
underlying mechanism of topo IIβ during transcription still needs to be elucidated (J. L. Nitiss, 2009)

1.2.5.4. Topoisomerase II in Neuronal Cells:

Topo IIβ had been also found to have a unique role in certain neuronal cells. No other type of topoisomerase has been found to have such a role (J. L. Nitiss, 2009). Topo IIβ had been established as one of the crucial enzymes for the neural development (Lyu & Wang, 2003; Yang, Li, Prescott, Burden, & Wang, 2000). Experiments in these studies were based on the loss of function of topo IIβ. Mice with homozygous deleted topo IIβ genes were inviable because of several neuronal deficits related to motor neuron failures. Microarray analysis of the same mice showed, approximately 1-4% of expressed genes were different. Furthermore, scientists showed that the localization of topo IIβ on various genes that are mostly related to developmentally regulated genes. Together these findings showed the importance of topo IIβ on neural cell development both directly and indirectly (J. L. Nitiss, 2009).

1.2.5.5. Topoisomerase II in Recombination, Checkpoints, and DNA Repair:

Various types of topoisomerases in the cell give rise to double stranded DNA (dsDNA) breaks. These dsDNA breaks in a way, impacts different mechanisms in the cell such as cell cycle checkpoint response activation, meiotic recombination, and DNA repair (J. Nitiss, 2009; Vos et al., 2011).

Several other topoisomerase like proteins accompany topoisomerase enzymes throughout these processes. For example: topo IIβ-A subunit homolog, Spo 11, creates dsDNA breaks during homologous recombination (Bergerat et al., 1997; Keeney, Giroux, & Kleckner, 1997; Pecia et al., 2002). After formation of dsDNA break by Spo11, protein must be detached from the DNA to prevent mutations (Neale, Pan, & Keeney, 2005). At this point, double strand break repair proteins like Rad50-Mre11-Nbs1 (MRN) complex (Keeney, 2008; Neale et al., 2005), Rad51 and Dmc1-like factors (Baudat, Manova, Yuen, Jasin, & Keeney, 2000; W. Li & Ma, 2006), and
nuclease Sae2/CtIP (Hartsuiker et al., 2009) remove Spo11 from the DNA in an independent mechanism from ATM (ataxia telangiectasia mutated) and ATR (ATM Rad3-related), which are the most known repair pathways.

As a consequence of their role on introducing strand breaks, topoisomerasases can also promote DNA repair mechanism. Cliby et al. showed ATM and ATR pathway activation upon Topo I caused breakage (Cliby, Lewis, Lilly, & Kaufmann, 2002). Moreover Treszezamsky et al. showed BRCA1 and BRCA2 pathway activation after topoisomerase II caused dsDNA breaks (Treszezamsky et al., 2007). All of these pathways are known to have a role both in cell cycle arrest and DNA repair mechanism. Several other studies showed that SUMOylation and ubiquitylation mediated destruction of topoisomerase enzymes, in which they inadvertently trapped with DNA in a covalent complex. Once the topoisomerase is destroyed, repairing enzymes such as Tdp1 and Tdp2 help to repair the covalent tyrosine-DNA adducts. In other words, the activation of the checkpoint response and DNA repair is dependent on both the type of the breakage and the enzymes involved in the process. However, the exact mechanism is not elucidated in detail yet (Vos et al., 2011).

There are also several studies showing the role of Topo II in cell cycle arrest. In one study, it had been shown in yeast cells that whereas a complete depletion of Topo II did not lead to cell cycle delay during mitosis, expression of the inactive Topo II led to a mitotic delay (Baxter & Diffley, 2008). This observation suggests that Topo II presence is needed to trigger the cell cycle arrest (Andrews et al., 2006). On the contrary, in another study, it had been observed that removal of Topo II via Tet-off system caused a mitotic delay but the depletion of Topo IIα via siRNA system did not induce mitotic delay (Hongchang Li, Wang, & Liu, 2008). The latter experiment suggests that knockdown of Topo IIα was insufficient to induce DNA repair which supports the conclusion of Andrews et al. (Andrews et al., 2006).
1.2.6. Topoisomerases as Targets in Cancer Treatment:

Although topoisomerase enzymes are very important for cell viability, any corruption on their DNA cleavage activity would give rise to mutagenic DNA breaks which may eventually lead to cell death. Scientists exploited this situation to deliberately kill malignant cells via topoisomerase inhibition in cancer therapy since 1980s. Several proteins, small molecules, and protein-like factors that can inhibit topoisomerase activity had been discovered. The mode of action of these molecules varies greatly. One kind attenuates the DNA binding activity of these enzymes, another attenuates the enzyme’s ability to cleave DNA (Vos et al., 2011).

1.2.7. Targeting Topoisomerases Using Small Molecules:
1.2.7.1. Examples of Small molecules Targeting Topoisomerases:

There are two broad classes of topoisomerase targeting small molecules based on their mode of action: “inhibitors” affect the enzyme activity and “poisons” stabilize DNA-enzyme complex. Whereas the effects of inhibitors are reversible, the effects of poisons are irreversible. And therefore poisons are considered to be more toxic for cells than inhibitors (Nitiss, 2009).

There are several molecules used for decades in significant therapeutics to inhibit the topoisomerase activity but their mechanisms of action have only been understood rather recently. For example; Camptothecin and its derivatives like Topotecan and irinotecan (Figure 9) affect Topo Iβ by intercalating between the active site of the enzyme and the cleaved DNA ends, thus blocking DNA religation and relaxation of supercoils. Therefore, they are classified as Topo Iβ poisons (Barros et al., 2013; Staker et al., 2002). Another example is Fluoroquinolones, whose targets are gyrase and Topo IV in bacteria, are used as antibiotics. Additionally, semisynthetic podophyllotoxin derivatives such as Etoposide and Teniposide (Figure 10) are being used as chemotherapeutic agents. Etoposide is a Topo II poison which binds between 5' and 3' end of a broken DNA and prevents the resealing and the releasing of the strand (Laponogov et al., 2010). Other examples are novo- and chlorobiocins (Lewis et al.,
1996; Tsai et al., 1997), radicicol (Corbett & Berger, 2006), and bisdioxopiperazines (Classen, Olland, & Berger, 2003). They all block the ATPase activity of the enzyme. Moreover, simocyclinone D8 affects DNA binding (Edwards et al., 2009) and NTBI GSK299423 affects DNA cleavage (Bax et al., 2010).

In addition to aforementioned examples, m-AMSA (Amsacrine) (Figure 11) needs particular attention. Because it is the first synthetic compound which targets topoisomerase II (Hornedo & Van Echo, 1985). Its working principle is based on intercalation and blocking the movement of DNA-enzyme complex (Ketron, Denny, Graves, & Osheroff, 2012). Its discovery led scientists to explore new, less toxic synthetic compounds for topoisomerase inhibition.

Together these findings not only highlight the variety of topoisomerase targeting molecules but also show the abundance of molecular scaffolds capable of topoisomerase inhibition (Vos et al., 2011).
Figure 9. Some Topo I poisons. a. camptothecin. b. topotecan. c. irinotecan.
1.2.7.2. Cell Response to Topoisomerase Targeting Small Molecules:

In most of the cells, accumulation of mutations, ubiquitinylation, SUMOylation, and the formation of reactive oxygen species (ROS) is observed upon the intake of a topoisomerase inhibitor or poison. All of these changes occurring inside the cell lead to
mitotic delay, induce DNA repair mechanisms and ultimately induce cell death due to unrepai red DNA breaks (Vos et al., 2011).

For instance, in bacteria, it had been observed that some fluoroquinolones (Topo IV poison) promote cell death by inducing ROS production (Dwyer, Kohanski, Hayete, & Collins, 2007; Kohanski, Dwyer, Hayete, Lawrence, & Collins, 2007). Interestingly, some other fluoroquinolones like moxifloxacin and PD161144 were shown to kill cells even though the ROS cascade is blocked (oxygen absence). This result suggests that there might be other pathways for anti-topoisomerase action (X. Wang, Zhao, Malik, & Drlica, 2010).

In eukaryotes, both Topo I and Topo II poisons induce ubiquitinylation and SUMOylation (Vos et al., 2011). These processes are needed to initiate repair response as described earlier. Repair response mechanism acts to decrease the effect of the poison on topoisomerase. One way to mitigate the effect of the repair response mechanism is to administer the repair response mechanism inhibitor concurrently with the poison. For instance, it had been shown that fibroblasts of mice lacking PARP-1, one of the crucial enzymes in repair mechanism, are found to be more sensitive to topo II inhibitor (C-1305) (Węsierska-Gądek, Schloffer, Gueorguieva, Uhl, & Składanowski, 2004). In another study, it had been shown that doxorubicin’s (topo II inhibitor) activity in liver cells was enhanced by ANI which is a PARP inhibitor (Muñoz-Gámez et al., 2011).

1.3. Effects of Azacyanines in Yeast Cells:

Azacyanines (Figure 12 a, b, and c) were first synthesized by Kurth et al. as ion channel inhibitors (Haddadin, Kurth, & Olmstead, 2000). The interactions of these azacyanines named as Azamethyl (named as Aza3 in previous studies), Aza4 and Aza5 with different DNA structures had also been established by lateral studies. By using NMR, UV-Vis and CD spectroscopy, it had been shown that these molecules bind selectively to G-quadruplex forming human telomeric sequence (tel24) over duplex DNA sequences (Çetinkol, Engelhart, Nanjunda, Wilson, & Hud, 2008). They had also
been able to induce the formation of the poly(A) self-structure in-vitro (Çetinkol & Hud, 2009).

Moreover, it has been shown that these azacyanines were able to induce genomic instability in yeast cells. In his thesis, Kim showed that azacyanine derivatives (Azamethyl, Aza4, and Aza5) induce chromosomal fragility in yeast cells. He hypothesized that this might be due to the stabilization of triple helical structures formed by \((GAA)_n\) triplex repeats. Moreover, it has been shown that increased chromosome fragility, caused by azacyanines, induced DNA repair surveillance system and caused G2/M phase arrest. It is also emphasized that azacyanines inhibit cell growth in a dose dependent manner (Kim, 2009).

However, there is no study, up to date, showing neither azacyanines’ high affinity towards triple helical structures nor their selectivity towards different nucleic acid structures. Though their chemical structure resembles especially to the intercalators mentioned above, meaning that they might have a potential to target several structures in a cell such as DNA, RNA, and/or DNA-RNA hybrids in addition to their complexes such as DNA-topoisomerase complexes. Therefore, inspired by these findings mentioned above we hypothesized that the observed effects of azacyanines in yeast cells might be due to their binding ability to different nucleic acid structures, especially to triplex and quadruplex structures and/or topoisomerase II alpha.
1.4. Thesis Focus:

This thesis focuses on the effect of several azacyanines on topoisomerase IIα and their selectivity towards different nucleic acid structures. Our main goal is to have a grasp on the mechanism of action of azacyanines observed in the yeast cells (Kim, 2009). Chemical structures of azacyanines used in our studies are given in Figure 12.

Here first, we focused on the effect of azacyanines on Topoisomerase II alpha enzyme (Topo IIα) which is the main expressed type II topoisomerase in the mammalian cells. Recently, a number of studies about synthetic topoisomerase inhibitors showed successful inhibition of Topo IIα. Since azacyanines (Figure 12) share similar aromatic ring structures with these molecules namely camptothecin, topotecan, irinotecan, etoposide, teniposide, and amsacrine (Figures 9-11), we hypothesized that the cell cycle arrest observed in yeast cells might be due to their ability to inhibit Topo IIα (Kim, 2009).

Secondly, we focused on revealing the binding affinity and selectivity of these azacyanines towards different nucleic acid structures via using competition dialysis to test Kim’s hypothesis on their binding ability to triple helical structures. Our goal was to shed light to the effect of azacyanines on genome instability observed in yeast cells via understanding their interactions with different DNA structures (Kim, 2009).

As a whole, at the end of this thesis, we aim to propose some alternative mechanisms on toxic effects of these azacyanine molecules in cells and reveal their potential as possible topoisomerase inhibitors and chemotherapeutic agents.
CHAPTER 2

POSSIBLE EFFECTS OF AZACYANINE DERIVATIVES ON TOPOISOMERASE II ALPHA (TOPO IIα)

2.1. Introduction:

Topoisomerase II alpha (Topo IIα) has long been utilized as a target in chemotherapeutic applications in clinical oncology. Usage of small molecules as therapeutic agents such as etoposide and doxorubicin (classified as Topo IIα poisons) to target Topo IIα due to their toxicity is one of the most commonly applied practices. Topo IIα poisons’ main function is to stabilize the Topo IIα-DNA complex, and eventually lead to apoptosis due to the accumulation of DNA damage. Though, it should also be noted that the utilization of such Topo IIα poisons bears also the risk of secondary malignancies and cardiotoxicity (Lipshultz et al., 2010; Vrooman et al., 2011).

In addition to Topo IIα poisons, there are several other small molecules identified as catalytic inhibitors. Catalytic inhibitors inhibit Topo IIα enzyme at different stages of its catalytic cycle without stabilizing the DNA-Topo enzyme complex (Figure 13) (Bau, Kang, Austin, & Kurz, 2013; Larsen, Escargueil, & Skladanowski, 2003; Pommier, 2013).
Based on the structural similarity, such as the extended aromatic ring system with the positive charge, of many topoisomerase II inhibitors to our azacyanines (Figures 9-12), we hypothesized that our molecules might also have the capability of inhibiting Topo IIα.

We utilized a kit from TopoGEN Inc. (Buena Vista, USA) which is optimized for revealing the catalytic inhibition efficiency of small molecules. The proposed reaction mechanisms with the catenated DNA (kDNA) substrate provided by the supplier are given in Figure 14. We have investigated the effect of Azamethyl, Aza4 and Aza5 on Topo IIα, which were shown to trigger cell cycle arrest in yeast cells (Kim, 2009). In addition to these three molecules we have included Azaethyl and Azaisobutyl compounds in our assay in order to understand the effect of small
molecule structure, mainly the effect of chain length and branching on the benzimidazole ring, on inhibition efficiency. After the incubation of the compounds with the enzyme (Topo IIα) and substrate (catenated DNA), the inhibition efficiency of azacyanines were assessed via agarose gel electrophoresis.

![Diagram of reaction mechanisms](image)

**Figure 14.** Proposed reaction mechanisms for Topoisomerase IIα kDNA based drug screening kit. kDNA: catenated DNA (substrate), VP-16 (Etoposide): proved topo II poison, CPT (Camptothecin): proved topo I inhibitor. Image taken from manufacturer’s user manual (TopoGEN Inc., Catalog no: TG1019-2).

### 2.2. Materials and Methods:

All the small molecules were synthesized and characterized in our lab according to previously published protocols (Haddadin *et al.*, 2000). Stock solutions of azacyanines were prepared by dissolving in DMSO (Sigma Aldrich, M81802) and further diluted with Tris-HCl therefore the effect of DMSO on the enzyme was minimized. Preparation of 10.0 mM Tris-HCl is described in detail in Appendix A. The
concentrations of the stock solutions were determined using calculated extinction coefficient values given in Table 1 (the Beer-Lambert’s Law: absorbance = extinction coefficient x path length x concentration) via UV-Vis spectrometer. Millipore de-ionized water was used throughout all of the experiments.

Table 1. Extinction coefficients of azacyanines.

<table>
<thead>
<tr>
<th>Name of the compounds</th>
<th>Extinction Coefficient (M⁻¹.cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azamethyl</td>
<td>343 nm – 45,700</td>
</tr>
<tr>
<td>Aza4</td>
<td>387 nm – 24,240</td>
</tr>
<tr>
<td>Aza5</td>
<td>407 nm – 25,000</td>
</tr>
<tr>
<td>Azaethyl</td>
<td>343 nm – 44,700</td>
</tr>
<tr>
<td>Azaisobutyl</td>
<td>343 nm – 44,700</td>
</tr>
<tr>
<td>Azapropyl</td>
<td>343 nm – 43,300</td>
</tr>
<tr>
<td>Azabutyl</td>
<td>343 nm – 44,000</td>
</tr>
<tr>
<td>Azaisopropyl</td>
<td>343 nm – 45,100</td>
</tr>
</tbody>
</table>

Topoisomerase II Drug Screening Kit (kDNA based) (TG1019-2) was purchased from TopoGEN Inc. (Buena Vista, USA). Experiments were conducted as instructed by the supplier. Briefly, required amount of ingredients (water, buffer, DNA, test compound, and enzyme) were pooled together in an eppendorf tube on ice followed by 37°C, 40 minutes incubation. The reactions were stopped by addition of 10% SDS at 37°C and followed by the addition of Proteinase K and 30 minutes incubation at 37°C to digest the enzyme. Afterwards loading dye was added to each sample and the samples were loaded into 1% agarose gel with EtBr. Agarose gels were prepared in 1xTAE buffer as described before (Lee, Costumbrado, Hsu, & Kim, 2012). Samples were poured to Thermo Fischer Owl Easycast B2 mini gel electrophoresis system and then run for 180 minutes at 100 Volt. At the end of the electrophoresis, gels were
stained with EtBr and de-stained in water. Gels were visualized using BioRad XR Molecular Gel Imager. Details of preparation of 50X TAE buffer, 1% agarose gel, and EtBr solution are given in Appendix A.

2.3. Results and Discussion:

2.3.1. Determination of the Optimal Concentration of Azamethyl for Topoisomerase II Alpha Inhibition:

We started our investigations on the effect of azacyanines on Topo IIα by determining the effect of different concentrations of Azamethyl on Topo IIα along with the proven Topo II poison Etoposide (VP-16) (Liu, 1989). Here all the reaction mixtures, unless said otherwise, include the substrate, 200.0 ng kDNA (catenated DNA) and 5.0 units of Topo IIα.

The result of our initial screening is given in Figure 15. In lane 1, kDNA, which was our substrate in our assay, was used as a control. Linearized and decatenated DNA products were also used as control markers in lanes 2 and 5 respectively. Lane 3 and 4 were our positive control 50.0 μM and 500.0 μM VP-16 respectively. VP-16 is expected to decrease the enzyme activity in a concentration dependent manner. In lane 6, a “blank” well that only includes the solvent DMSO and the buffer (Tris-HCl) was also used as a control in order to see the effect of our solutions on Topo IIα activity. Here the ratio of DMSO:Tris-HCl was 1:9 which was the ratio used in preparation of positive control VP-16 samples in lanes 3 and 4. Lane 7, a “no drug” well, was another control which only includes the reaction mixture without any inhibitor. Here, we expect to see the full enzyme activity on our substrate. Lanes 8-11 had different concentrations of Azamethyl added to the reaction mixture.

As shown in Figure 15, our enzyme was active under the reaction conditions and our solution used in the preparation of VP-16 and azacyanines had little effect on our enzyme’s activity (Lanes 6 and 7). Our substrate, kDNA by itself barely enters the gel (Lane 1). But in the presence of Topo IIα (Lanes 6 and 7), several bands were observed due to the formation of decatenated kDNA products by Topo IIα. Decatenated
bands in lanes 3, 4, and 6-11 are not at the same alignment with the decatenated DNA marker because of the long running time (3 hours) and should not be considered as a linear band. VP-16 is clearly inhibiting Topo IIα in a dose dependent manner (Lanes 3 and 4). The intensity of the decatenated DNA bands was decreased in the presence of 50.0 μM VP-16. The measured intensities of the decatenated DNA bands were given in Figure 16. Here, the intensities of decatenated DNA bands in all samples were normalized to the intensity of the decatenated DNA band belonging to the sample that has full Topo IIα activity in the absence of any inhibitor (lane 7-no drug). The intensity decreased to 0.63 in the presence of 50.0 μM VP-16 (lane 3) compared to the lane 7 which was taken as 1.00. The intensity of the band decreased further to 0.20 in the presence of 500.0 μM VP-16. Our compound, Azamethyl, showed incredible catalytic activity decrease compared to VP-16. The intensities of the bands were 0.75, 0.26, 0.12 and 0.01 in the presence of 1.0 μM, 10.0 μM, 50.0 μM and 100.0 μM Azamethyl respectively. It is clear that Azamethyl decreases the catalytic activity of Topo IIα in a concentration dependent manner. However, at this point, we cannot classify its inhibition efficiency as a Topo IIα poison, since we did not observe the band belonging to the linear DNA in the presence of Azamethyl. The linear band formation, even though it was weak, was observed in the presence low concentrations of VP-16. More importantly, our results revealed that the effect of Azamethyl on Topo IIα was stronger compared to VP-16. Even at 50.0 μM concentration Azamethyl’s efficiency was greater than 500.0 μM of VP-16.
Figure 15. Agarose gel electrophoresis results for the effect of Azamethyl on Topo IIα. Lane 1: “kDNA”, catenated DNA substrate, Lane 2: “Linear DNA marker” Topo IIα poison indicator, Lane 3 and 4: Topo IIα and kDNA in the presence of “VP-16” (Etoposide) which is a proven Topo IIα poison, Lane 5: “Decaten. DNA marker”, decatenated DNA products as an indicator of the enzyme activity, Lane 6: “Blank” only drug solvents (DMSO:Tris-HCl - 1:9), Lane 7: “no drug” includes only the enzyme Topo IIα. Lane 8-11: Topo IIα and kDNA in the presence of “Azamethyl” which is the tested drug. Gel conditions: 1%agarose in 1xTAE buffer; 100 V; 180 minutes run. Black arrow indicates the band that is the main indicator of the enzyme activity. Red circle: all intensities are normalized according to this band.
Figure 16. Agarose gel electrophoresis results obtained by analyzing the gel image given in Figure 15. Left table is the normalized intensity values. Normalization was performed relative to our negative control (no drug sample) that only includes the substrate (kDNA) and the enzyme (Topo IIα). The contents of the samples were given in Figure 15. Error bars represent 5% of the normalized intensities. Intensities measured by Image Lab, Version 5.2.1 (BioRad Laboratories).
2.3.2. Determination of the Effects of Different Azacyanines on Topoisomerase II Alpha:

Once we have seen that Azamethyl was decreasing the catalytic activity of Topo IIα, we decided to investigate the effects of other azacyanines also on Topo IIα. We included Aza4 and Aza5 in our assay since these two molecules were studied by Kim in his studies along with Azamethyl (Kim, 2009). In addition, we included two Azamethyl analogs named as Azaethyl and Azaisobutyl to understand the effect of alkyl chain length and branching on Topo IIα catalytic activity. Once again, we have included VP-16 as our positive control. Since the efficiency of Azamethyl was strong at 50.0 μM concentration, here we tested all the drugs efficiencies at that concentration on 5 units of Topo IIα. Again we have included kDNA as our substrate along with linearized and decatenated DNA markers. No drug lane and the blank lane (DMSO:Tris-HCl, 1:9) were also used as our negative controls.

Our results given in Figure 17 clearly demonstrated that Azamethyl, Aza4 and Aza5 can decrease the catalytic activity of Topo IIα. The intensity of the decatenated DNA decreased in the presence of Azamethyl, Aza4 and Aza5. The measured intensities given in Figure 18 were 0.16, 0.26 and 0.37 respectively. We did not observe the formation of the linear band in the presence of Azamethyl, Aza4 and Aza5 thus cannot classify them as poisons. Decatenated bands in lanes 3, 5-11 are not at the same alignment with the decatanated DNA marker because of the long running time (3 hours) and should not be considered as a linear band. To our surprise, Azaethyl and Azaisobutyl did not show any change on Topo IIα catalytic activity. (Figures 17 and 18). On the contrary, the activity of Topo IIα seemed to increase slightly in the presence of Azaethyl and Azaisobutyl. The intensities of the decatenated DNA band were measured as 1.20 and 1.21 in the presence of Azaethyl and Azaisobutyl, respectively.
Figure 17. Agarose gel electrophoresis results for different azacyanines on Topo IIα. Lane 1: “kDNA”, catenated DNA substrate, Lane 2: “Linear DNA marker”, Lane 3: kDNA and Topo IIα in the presence of “VP-16” (Etoposide) which is a proven Topo IIα poison, Lane 4: “Decaten. DNA marker”, decatenated DNA products as an indicator of the enzyme activity, Lane 5: “Blank” only drug solvents (DMSO:Tris-HCl - 1:9), Lane 6: “no drug” includes only the enzyme Topo IIα. Lane 7-11: kDNA and Topo IIα in the presence of 50.0 µM azacyanine derivatives. Lane 7: “Azamethyl”, Lane 8: “Aza4”, Lane 9: “Aza5”, Lane 10: “Azaethyl”, Lane 11: “Azaisobutyl”. Gel conditions: 1%agarose in 1xTAE buffer; 100 V; 180 minutes run. Black arrow indicates the band that is the main indicator of the enzyme activity. Red circle: all intensities are normalized according to this band.
Figure 18. Agarose gel electrophoresis results obtained by analyzing the gel image given in Figure 17. Left table is the normalized intensity values. Normalization was performed relative to our negative control (no drug sample) that only includes the substrate (kDNA) and the enzyme (Topo IIα). The contents of the samples were given in Figure 17. Error bars represent 5% of the normalized intensities. Intensities measured by Image Lab, Version 5.2.1 (BioRad Laboratories).

2.4. Conclusions:

Our results revealed that Azamethyl, Aza4 and Aza5 cause to decrease in Topo IIα’s catalytic activity, while Azaethyl and Azaisobutyl do not.
Among the molecules investigated in here, Azamethyl showed the highest decrease in TopoIIα’s catalytic activity. The effect of Azamethyl on Topo IIα was concentration dependent, such that there was a slight decrease on catalytic activity of Topo IIα even in the presence of 1.0 μM Azamethyl. More importantly, our results revealed that Azamethyl is more powerful than the proven Topo IIα poison, Etoposide (VP-16). While the topoisomerase was 63% active in the presence of 50.0 μM VP-16, its activity was only 12% in the presence of 50.0 μM Azamethyl. To our surprise, 50.0 μM Azamethyl showed stronger effect than 500.0 μM VP-16. We have also tested the effect of Azaethyl, Azaisobutyl, Aza4 and Aza5 at 50.0 μM small molecule concentration. Our results revealed that while Aza4 and Aza5 also decrease the catalytic activity of Topo IIα, Azaethyl and Azaisobutyl do not show such ability. On the contrary, to our surprise the activity of Topo IIα seemed to increase slightly in the presence of Azaethyl and Azaisobutyl. The interactions of these molecules with Topo IIα need to be investigated in detail in order to be able to explain such an increase. However, we think that the underlying cause might be related to the interactions of these molecules with the DNA rather than TopoIIα. As will be explained in chapter 3, azacyanines have the capability of binding to different nucleic acids. Therefore if Azaethyl and Azaisobutyl do not interfere with the Topo IIα, they might be binding to decatenated DNA products and this could cause the increased intensities.

Results obtained here provide us an alternative explanation on how azacyanines might be affecting the cells in addition to the observations by Kim (Kim, 2009). It is now clear that Azamethyl, Aza4, and Aza5 decreases Topo IIα enzyme’s catalytic activity. This might be one of the reasons behind the cell cycle arrest observed by Kim (Kim, 2009). They might be interfering with the enzyme at different steps of its catalytic cycle. For example, they might be interfering with the binding of the enzyme to DNA, stabilizing the non-covalent DNA-enzyme complex, or inhibiting ATP binding to the enzyme (Larsen et al., 2003). Yet, at which stage of its catalytic cycle that our small molecules interfere with Topo IIα needs to be elucidated further.
CHAPTER 3

SELECTIVITY OF AZACYANINES TOWARDS NUCLEIC ACID STRUCTURES

3.1. Introduction:

For more than 40 years, mainly because of the participation of small molecules as therapeutic drugs, designing of small molecules that have the ability to bind nucleic acids has been an active area (Waring & Wakelin, 2003). Targeting nucleic acid structures (DNA and/or RNA) with small molecules gives an opportunity for the genetically originated diseases’ treatment (Martinez & Chacon-Garcia, 2005; Thurston, 1999). Particularly, small molecule binding to non-canonical nucleic acid structures such as triplex and quadruplexes has been a major focus for rational drug design. However, designing molecules that can bind to nucleic acid structures tightly and selectively is the first step to understand such interactions (Hurley, 1989; Mergny & Helene, 1998; Thurston, 1999).

Small molecules, upon binding to certain nucleic acid structures, change the conformation and therefore could either stabilize or destabilize these structures and cause problems such as impaired gene expression patterns and/or mutations (Bissler, 2007). This subsequently induces DNA repair systems and drives cells to cell cycle arrest and could even lead to cell death. Their stabilization ability, have been widely used in drug design for several treatments such as cancer to get rid of undesired cells (M. Wang et al., 2016). Moreover, selectivity of various other small molecules towards
specific nucleic acid sequences had been shown with different studies (Chaires et al., 2003; Ren & Chaires, 1999; Chaires, 1998).

Selectivity of Azamethyl, Aza4, and Aza5 to quadruplex structures along with duplex DNA structures and poly(A) had been reported previously (Çetinkol et al., 2008; Çetinkol & Hud, 2009). Additionally, Kim investigated the effects of these molecules in-vivo using yeast cells, expressing GAA•TTC repeat sequences. It had been revealed that all the molecules were blocking the DNA synthesis, inducing chromosome fragility and inhibiting cell growth in these yeast cells in a dose dependent manner. Based on his results, Kim hypothesized that the observed results might be due to the binding of these molecules to triple helical structures. He even suggested that these molecules can form the basis for the development of novel antitumor drugs that bind to triple helical structures tightly and act via the inhibition of cellular proliferation (Kim, 2009).

In Chapter 2, we have observed that Azamethyl, Aza4, and Aza5 catalytically inhibit Topo IIα. Yet at which catalytic step do they inhibit the enzyme’s activity is not known. Therefore, we hypothesized that if azacyanines have the capability of binding to different nucleic acid structures; they might be affecting the enzyme at the first step of its catalytic activity, which is the binding of the enzyme to DNA as shown in Figure 7. Azacyanines might be interfering with the DNA and thus changing the conformation of the DNA. This might be preventing the enzyme binding to DNA and thus inhibiting the enzyme activity.

Here, using competition dialysis method, we have investigated the selectivity of azacyanines towards various nucleic acid structures. We have also included Azaethyl, Azapropyl, Azaisopropyl, Azabutyl and Azaisobutyl in addition to Azamethyl, Aza4, and Aza5, in our competition dialysis experiments to see the effect of alkyl chain length and branching of benzimidazole ring on nucleic acid binding affinity and selectivity.
3.2. Materials and Methods:

Competition dialysis experiments were conducted as previously described (Chaires et al., 2003; Ragazzon, Garbett, & Chaires, 2007; Ren & Chaires, 1999). Components and preparation of the buffers used throughout the experiments are given in Appendix A and the oligonucleotides used in the experiments are given in Appendix B. Construction of standard curves are given in Appendix C.

The oligonucleotide samples were prepared as described in Ren & Chaires, 1999. Briefly, oligonucleotides including poly(dA), poly(dT), poly(dA).poly(dT), poly(dAdT).poly(dAdT), poly(rA).poly(rU), poly(dGdC).poly(dGdC) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Natural DNA of Micrococcus lysodeikticus purchased from Sigma-Aldrich (St. Louis, MO, USA) was sonicated for 30 minutes (5 minutes sonication following 5 minutes rest) followed by phenol-chloroform extraction (Chaires, Dattagupta, & Crothers, 1982). 32 base long oligonucleotides including (dA)$_{32}$ and (dT)$_{32}$ and quadruplex sequences including Tel24 (5′-TTGGG[TTAGGG]$_3$A- 3’) and 5′- T$_2$G$_{20}$T$_2$ -3’ were purchased from Integrated DNA Technologies (IDT) (Coralville, IA, USA). Remaining oligonucleotides were produced by mixing the required nucleic acids as described below:

- **Triplex DNA - poly(dA).[(poly(dT)]$_2$** was prepared by mixing poly(dA).poly(dT) and poly(dT) in a 1:1 equimolar ratio,
- **Triplex DNA - (dA)$_{32}$.[(dT)$_{32}$.** was prepared by mixing (dA)$_{32}$ and (dT)$_{32}$ in a 1:2 equimolar ratio,
- **Triplex RNA - poly(rA).[(poly(rU)]$_2$** was prepared by mixing either poly(rA).poly(rU) and poly(rU) in a 1:1 equimolar ratio, or poly(rA) and poly(rU) in a 1:2 equimolar ratio
- **Duplex DNA - (dA)$_{32}$.[dT]$_{32}$** was prepared by mixing (dA)$_{32}$ and (dT)$_{32}$ in a 1:1 equimolar ratio.
All the prepared oligonucleotides were annealed by heating at 95°C for 5 minutes in a water bath and then left overnight at room temperature for cooling down slowly to assure the formation of the proper secondary structure before used in competition dialysis experiments.

For each competition dialysis experiment, 0.5 mL of 75.0 µM of each oligonucleotide, in a monomeric unit (nucleotide, base pair, triplet, quartet), were put into Pierce (Thermofischer Scientific, USA) 7000 Da molecular weight cutoff dialysis cassettes. Then, cassettes were dialyzed against 500.0 mL of 1.0 µM of the selected ligand (Azamethyl, Azaethyl, Azapropyl, Azabutyl, Azaisobutyl, Azaisopropyl, Aza4, or Aza5) solution for 24 hours. 1xBPES buffer (1.0 mM Na$_2$EDTA, 6.0 mM Na$_2$HPO$_4$, 2.0 mM NaH$_2$PO$_4$, and 185.0 mM NaCl); pH 7.0, was prepared (see Appendix A) as previously described (Ren & Chaires, 1999) and used in all experiments.

At the end of the dialysis period, aliquots were taken from each dialysis cassettes, and 1% (w/v) SDS was added to release the bound ligands from nucleic acids and to ensure that the ligands were free in the cassettes. Afterwards, the ligand concentrations in each cassette were determined via fluorescence spectroscopy (Varian, Cary Eclipse). Device setup was as following: Emission spectra collected: 325 to 675 nm, excitation wavelength: 324 nm, excitation and emission slits: 2.5 and 5.0, respectively. For each azacyanine derivative, standard curves were constructed using the same fluorescence spectrophotometer prior to competition dialysis (see Appendix C). This allowed us to determine the concentration of ligands by using fluorescence intensities. Appropriate corrections were made in calculating the ligand concentration in each cassette considering the dilution due to the added SDS and 1X BPES buffer. Two independent experiments were performed for each molecule.

The ratio of the bound ligand to free ligand concentration ($C_b/C_f$) was determined by: $C_b/C_f=(C_t/C_f)-1$ where $C_t$ (total ligand concentration) is the concentration of the ligand in each dialysis cassette and $C_f$ (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (that only
includes 1X BPES buffer) which typically did not significantly vary from initial small molecule concentration that was 1.0 µM.

Descriptive statistics was used to analyze our competition dialysis results. SPSSv23.0.0.0 (IBM corp., Armonk, NY) was utilized for statistical analysis. Kruskal Wallis non-parametric test was performed for non-normal distributions. One-way ANOVA test was performed, with Bonferroni and Sidak post-hoc tests, for normal distributions (see Appendix D).

3.3. Results and Discussions:

3.3.1. Azamethyl:

At the end of the 24 hours long competition period, the concentration of Azamethyl was different in each dialysis cassette as indicated by the varying fluorescence intensities among the cassettes containing different nucleic acids as shown in Figure 19. And more importantly, the fluorescence intensities in certain cassettes were also highly different than the fluorescence intensity obtained for the control cassette that is only containing buffer solution. This means that Azamethyl was binding to certain nucleic acid structures and therefore diffusing into the cassettes that are containing these nucleic acid structures. Using the fluorescence intensities, the exact concentration of Azamethyl in each dialysis cassette was calculated as explained in Appendix C. In each trial, $C_b/C_f$ for each cassette was calculated, and then $C_b/C_f$ ratios for both of the replicate experiments were averaged out and recorded as the final result (Table 2). The final average $C_b/C_f$ values for each nucleic acid structure were plotted as shown in Figure 20. Our statistical results were also given in Figure 20.
Figure 19. Average fluorescence spectra of Azamethyl. Measurements were obtained at the end of the 24 hours long competition dialysis against Azamethyl from dialysis cassettes containing different nucleic acid structures.
Table 2. Competition dialysis results for Azamethyl.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>Folds over buffer (C_b/C_f)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1st replicate</td>
</tr>
<tr>
<td>poly(dA),[poly(dT)]_2 - triplex DNA</td>
<td>10.11</td>
</tr>
<tr>
<td>(dA)_32,[(dT)_32]_2 - triplex DNA</td>
<td>10.78</td>
</tr>
<tr>
<td>5'-T3G20T2 - 3' - quadruplex DNA</td>
<td>2.51</td>
</tr>
<tr>
<td>poly(rA),[poly(rU)]_2 - triplex RNA</td>
<td>1.60</td>
</tr>
<tr>
<td>M.lysodeiktus - dsDNA</td>
<td>1.32</td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>1.26</td>
</tr>
<tr>
<td>poly(dGdC),poly(dGdC) - dsDNA</td>
<td>1.14</td>
</tr>
<tr>
<td>poly(dA),poly(dT) - dsDNA</td>
<td>0.77</td>
</tr>
<tr>
<td>poly(rA),poly(dT) - DNA-RNA hybrid</td>
<td>0.48</td>
</tr>
<tr>
<td>poly(rA),poly(rU) - dsRNA</td>
<td>0.64</td>
</tr>
<tr>
<td>poly(dAdT),poly(dAdT) - dsDNA</td>
<td>0.27</td>
</tr>
<tr>
<td>(dA)_32,(dT)_32 - dsDNA</td>
<td>0.09</td>
</tr>
<tr>
<td>poly(rA) - ssRNA</td>
<td>0.13</td>
</tr>
<tr>
<td>poly(dT) - ssDNA</td>
<td>0.02</td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
</tr>
<tr>
<td>tank</td>
<td>0.03</td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>-0.01</td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>0.06</td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration (C_b/C_f) was determined by: C_b/C_f=(C_t/C_f)-1 where C_t (total ligand concentration) is the concentration of the ligand in each dialysis cassette and C_f (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Figure 20. Competition Dialysis results for Azamethyl. *; p<0.05 from control by Kruskal Wallis non-parametric test. Error bars are ±1 standard deviations of the mean with a confidence level of 95%.

Our competition dialysis results revealed undoubtedly that Azamethyl is binding to triple helical structures of poly(dA).poly([dT])2 and (dA)32.([dT]32)2 with high affinity and selectivity. The amount of Azamethyl collected in the dialysis cassettes containing poly(dA).poly([dT])2 and (dA)32.([dT]32)2 was substantial compared to other samples. The average calculated Cb/Cf values were 11.69 and 10.65 with significance levels of p=0.012 and p=0.014 respectively according to non-parametric Kruskal–Wallis test. Azamethyl also showed significant binding to 5’-T2G20T2-3’ quadruplex and poly(rA).[poly(rU)]2 RNA triplex with Cb/Cf values of 2.53 and 1.68, respectively. Our results also revealed that, Azamethyl’s affinity towards
double stranded structures was low with even lower affinity towards RNA structures compared to DNA structures. Furthermore, Azamethyl had no affinity towards single stranded nucleic acid structures under these conditions, where the obtained \( C_b/C_f \) values were within the experimental errors.

### 3.3.2. Azaethyl:

The fluorescence spectra of the samples against Azaethyl in each dialysis cassette were given in Figure 21. The concentration of Azaethyl was also different in each dialysis cassettes after 24 hours long competition dialysis. Then \( C_b/C_f \) was calculated, as mentioned in materials and methods, and \( C_b/C_f \) ratio for both of the replicate experiments were averaged out and recorded as the final result (Table 3). The average \( C_b/C_f \) values with the statistical analysis were plotted in Figure 22.

![Azaethyl](image)

**Figure 21.** Average fluorescence spectra of Azaethyl. Measurements were obtained at the end of the 24 hours long competition dialysis against Azaethyl from dialysis cassettes containing different nucleic acid structures.
Table 3. Competition dialysis results for Azaethyl.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>Folds over buffer (C_b/C_t)</th>
<th>1st replicate</th>
<th>2nd replicate</th>
<th>Average</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>poly(dA)_2[poly(dT)]_2 - triplex DNA</td>
<td>6.95</td>
<td>6.88</td>
<td>6.92</td>
<td>0.05</td>
<td></td>
</tr>
<tr>
<td>(dA)_2[(dT)_2]_2 - triplex DNA</td>
<td>1.75</td>
<td>1.00</td>
<td>1.37</td>
<td>0.53</td>
<td></td>
</tr>
<tr>
<td>5'-T2G20T2 – 3’- quadruplex DNA</td>
<td>0.88</td>
<td>0.72</td>
<td>0.80</td>
<td>0.11</td>
<td></td>
</tr>
<tr>
<td>poly(rA)_2[poly(rU)]_2 - triplex RNA</td>
<td>-0.02</td>
<td>-0.01</td>
<td>-0.02</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>M.lysodeiktus - dsDNA</td>
<td>0.49</td>
<td>0.32</td>
<td>0.40</td>
<td>0.12</td>
<td></td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>0.24</td>
<td>0.15</td>
<td>0.19</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td>poly(dGdC)_2.poly(dGdC) - dsDNA</td>
<td>0.20</td>
<td>0.13</td>
<td>0.17</td>
<td>0.05</td>
<td></td>
</tr>
<tr>
<td>poly(dA)_2.poly(dT) - dsDNA</td>
<td>0.36</td>
<td>0.42</td>
<td>0.39</td>
<td>0.04</td>
<td></td>
</tr>
<tr>
<td>poly(rA)_2.poly(dT) - DNA-RNA hybrid</td>
<td>0.08</td>
<td>-0.02</td>
<td>0.03</td>
<td>0.08</td>
<td></td>
</tr>
<tr>
<td>poly(rA)_2.poly(rU) - dsRNA</td>
<td>0.00</td>
<td>-0.08</td>
<td>-0.04</td>
<td>0.05</td>
<td></td>
</tr>
<tr>
<td>poly(dAdT)_2.poly(dAdT) - dsDNA</td>
<td>0.04</td>
<td>0.03</td>
<td>0.03</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>(dA)_2[(dT)_2]_2 - dsDNA</td>
<td>0.00</td>
<td>0.02</td>
<td>0.01</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>poly(rA) - ssDNA</td>
<td>-0.03</td>
<td>-0.03</td>
<td>-0.03</td>
<td>0.00</td>
<td></td>
</tr>
<tr>
<td>poly(dT) - ssDNA</td>
<td>0.00</td>
<td>-0.04</td>
<td>-0.02</td>
<td>0.03</td>
<td></td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td></td>
</tr>
<tr>
<td>tank</td>
<td>0.04</td>
<td>0.03</td>
<td>0.04</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>-0.03</td>
<td>-0.02</td>
<td>-0.03</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>-0.07</td>
<td>-0.04</td>
<td>-0.05</td>
<td>0.02</td>
<td></td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration (C_b/C_t) was determined by: C_b/C_t=(C_t/C_f)-1 where C_t (total ligand concentration) is the concentration of the ligand in each dialysis cassette and C_f (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Azaethyl also showed significant selectivity, though not as dramatic as Azamethyl’s, towards poly(dA)[poly(dT)]$_2$ and (dA)$_{32}$[(dT)$_{32}$]$_2$ triplex DNA structures with average $C_b/C_f$ values of 6.92 and 1.37 and significance levels of $p=0.026$ and $p=0.033$, respectively, according to non-parametric Kruskal–Wallis test. Other than its binding to poly(dA)[poly(dT)]$_2$ and (dA)$_{32}$[(dT)$_{32}$]$_2$ triplex DNA structures, its binding to 5’-T$_2$G$_{20}$T$_2$-3’ quadruplex was also significant according to our statistical analysis ($p=0.044$). Its binding was found to be nonsignificant to any other sequences.
3.3.3. Azapropyl:

The fluorescence spectra of the samples in each dialysis cassette were depicted in Figure 23. The concentration of Azapropyl was also different in each dialysis cassettes after 24 hours long competition. The calculated $C_b/C_f$ values for both of the replicate experiments were given in Table 4 along with the average $C_b/C_f$ values. The average $C_b/C_f$ values with the statistical analysis were also plotted in Figure 24.

Figure 23. Average fluorescence spectra of Azapropyl. Measurements were obtained at the end of the 24 hours long competition dialysis against Azapropyl from dialysis cassettes containing different nucleic acid structures.
Table 4. Competition dialysis results for Azapropyl.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>Folds over buffer (C_b/C_f)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1st replicate</td>
</tr>
<tr>
<td>poly(dA).[poly(dT)]_2 - triplex DNA</td>
<td></td>
</tr>
<tr>
<td></td>
<td>4.62</td>
</tr>
<tr>
<td>(dA)_32.[(dT)_32] - triplex DNA</td>
<td>0.65</td>
</tr>
<tr>
<td>5'-T_G20T_2 - 3' - quadruplex DNA</td>
<td>1.27</td>
</tr>
<tr>
<td>poly(rA).[poly(rU)]_2 - triplex RNA</td>
<td>0.13</td>
</tr>
<tr>
<td>M.lysodeiktus - dsDNA</td>
<td>0.39</td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>0.24</td>
</tr>
<tr>
<td>poly(dGdC).poly(dGdC) - dsDNA</td>
<td>0.21</td>
</tr>
<tr>
<td>poly(dA).poly(dT) - dsDNA</td>
<td>0.32</td>
</tr>
<tr>
<td>poly(rA).poly(dT) - DNA-RNA hybrid</td>
<td>0.05</td>
</tr>
<tr>
<td>poly(rA).poly(rU) - dsRNA</td>
<td>-0.05</td>
</tr>
<tr>
<td>poly(dAdT).poly(dAdT) - dsDNA</td>
<td>-0.02</td>
</tr>
<tr>
<td>(dA)_32.(dT)_32 - dsDNA</td>
<td>0.10</td>
</tr>
<tr>
<td>poly(rA) - ssRNA</td>
<td>-0.02</td>
</tr>
<tr>
<td>poly(dT) - ssDNA</td>
<td>0.05</td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
</tr>
<tr>
<td>tank</td>
<td>0.16</td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>0.08</td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>0.03</td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration ($C_b/C_f$) was determined by: $C_b/C_f = (C_t/C_f) - 1$ where $C_t$ (total ligand concentration) is the concentration of the ligand in each dialysis cassette and $C_f$ (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Figure 24. Competition Dialysis results for Azapropyl. *; p<0.05 from control by Kruskal Wallis non-parametric test. Error bars are ±1 standard deviations of the mean with a confidence level of 95%.

Our results have revealed that Azapropyl also shows the highest affinity towards poly(dA)[poly(dT)]$_2$ triplex DNA structure. However, its affinity to 5’-T$_2$G$_{20}$T$_2$-3’ quadruplex sequence was higher than its affinity to (dA)$_{32}$[(dT)$_{32}$]$_2$ triplex DNA unlike Azamethyl and Azaethyl molecules. The obtained C$_b$/C$_f$ values for poly(dA)[poly(dT)]$_2$, 5’-T$_2$G$_{20}$T$_2$-3’, and (dA)$_{32}$[(dT)$_{32}$]$_2$ were 5.18 (p=0.024) and 1.28 (0.031), and 0.75 (p=0.041), respectively.
3.3.4. Azabutyl:

The fluorescence spectra of the Azabutyl samples obtained from dialysis cassettes after 24 hours of competition dialysis were displayed in Figure 25. Again, the concentration of Azabutyl was varied between different dialysis cassettes. The calculated $C_b/C_f$ ratios for both of the replicate experiments were given in Table 5 and the average $C_b/C_f$ values with the statistical analysis were plotted in Figure 26.

![Figure 25. Average fluorescence spectra of Azabutyl. Measurements were obtained at the end of the 24 hours long competition dialysis against Azabutyl from dialysis cassettes containing different nucleic acid structures](image-url)
Table 5. Competition dialysis results for Azabutyl.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>1st replicate</th>
<th>2nd replicate</th>
<th>Average</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>poly(dA).[poly(dT)]₂ - triplex DNA</td>
<td>4.36</td>
<td>5.53</td>
<td>4.94</td>
<td>0.82</td>
</tr>
<tr>
<td>(dA)₃₂.[(dT)₃₂]₂ - triplex DNA</td>
<td>0.31</td>
<td>0.31</td>
<td>0.31</td>
<td>0.00</td>
</tr>
<tr>
<td>5′-T₃₂G₂₀T₂ – 3′- quadruplex DNA</td>
<td>4.30</td>
<td>4.94</td>
<td>4.62</td>
<td>0.45</td>
</tr>
<tr>
<td>poly(rA).[poly(rU)]₂ - triplex RNA</td>
<td>0.04</td>
<td>0.03</td>
<td>0.03</td>
<td>0.00</td>
</tr>
<tr>
<td>M.lysoseiktus - dsDNA</td>
<td>0.71</td>
<td>0.57</td>
<td>0.64</td>
<td>0.10</td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>0.64</td>
<td>0.61</td>
<td>0.62</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(dGdC).poly(dGdC) - dsDNA</td>
<td>0.18</td>
<td>0.29</td>
<td>0.23</td>
<td>0.08</td>
</tr>
<tr>
<td>poly(dA).poly(dT) - dsDNA</td>
<td>0.26</td>
<td>0.37</td>
<td>0.32</td>
<td>0.08</td>
</tr>
<tr>
<td>poly(rA).poly(dT) - DNA-RNA hybrid</td>
<td>0.18</td>
<td>0.16</td>
<td>0.17</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(rA).poly(rU) - dsRNA</td>
<td>0.04</td>
<td>0.11</td>
<td>0.07</td>
<td>0.05</td>
</tr>
<tr>
<td>poly(dAdT).poly(dAdT) - dsDNA</td>
<td>0.10</td>
<td>0.07</td>
<td>0.09</td>
<td>0.02</td>
</tr>
<tr>
<td>(dA)₃₂.(dT)₃₂ - dsDNA</td>
<td>0.13</td>
<td>0.07</td>
<td>0.10</td>
<td>0.05</td>
</tr>
<tr>
<td>poly(rA) - ssRNA</td>
<td>0.07</td>
<td>-0.02</td>
<td>0.02</td>
<td>0.07</td>
</tr>
<tr>
<td>poly(dT) - ssRNA</td>
<td>-0.01</td>
<td>0.16</td>
<td>0.07</td>
<td>0.12</td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>tank</td>
<td>0.19</td>
<td>0.34</td>
<td>0.26</td>
<td>0.11</td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>0.00</td>
<td>0.10</td>
<td>0.05</td>
<td>0.07</td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>0.04</td>
<td>0.01</td>
<td>0.03</td>
<td>0.02</td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration (Cₘ/Cₘ) was determined by: Cₘ/Cₘ=(Cₜ/Cₘ)-1 where Cₜ (total ligand concentration) is the concentration of the ligand in each dialysis cassette and Cₘ (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Figure 26. Competition Dialysis results for Azabutyl. *; p<0.05 from control by Kruskal Wallis non-parametric test. Error bars are ±1 standard deviations of the mean with a confidence level of 95%.

Azabutyl showed similar affinity towards poly(dA)[poly(dT)]₂ triplex and 5’T₂G₂₀T₂-3’ quadruplex with Cₜ/Cₛ values of 4.94 and 4.62, respectively. Interestingly, it did not show any significant affinity to (dA)₃₂[(dT)₃₂]₂ triplex. Such a result indicates that the affinity of the small molecules depends not only to the structure of the nucleic acid but also its length. Conversely, Azabutyl also showed statistically significant binding to Micrococcus lysodeikticus (Cₜ/Cₛ = 0.64 folds) and tel24 (Cₜ/Cₛ =0.62 folds) according to our analysis.
3.3.5. Azaisopropyl:

The fluorescence spectra of the Azaisopropyl samples recovered from dialysis cassettes were provided in Figure 27. The concentration of Azaisopropyl was only slightly different in each dialysis cassette as given in Table 6. Average $C_b/C_f$ values given in Figure 28 were also not that different from each other.

**Figure 27.** Average fluorescence spectra of Azaisopropyl. Measurements were obtained at the end of the 24 hours long competition dialysis against Azaisopropyl from dialysis cassettes containing different nucleic acid structures
Table 6. Competition dialysis results for Azaisopropyl.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>1st replicate</th>
<th>2nd replicate</th>
<th>Average</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>poly(dA).[poly(dT)]_2 - triplex DNA</td>
<td>0.22</td>
<td>0.13</td>
<td>0.17</td>
<td>0.06</td>
</tr>
<tr>
<td>(dA)_32.[(dT)_32] - triplex DNA</td>
<td>0.13</td>
<td>-0.02</td>
<td>0.05</td>
<td>0.10</td>
</tr>
<tr>
<td>S’-T_2G_20T_2 – 3’- quadruplex DNA</td>
<td>0.26</td>
<td>0.14</td>
<td>0.20</td>
<td>0.08</td>
</tr>
<tr>
<td>poly(rA).[poly(rU)]_2 - triplex RNA</td>
<td>0.01</td>
<td>-0.02</td>
<td>-0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>M.lysodeiktus - dsDNA</td>
<td>0.15</td>
<td>0.09</td>
<td>0.12</td>
<td>0.05</td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>0.12</td>
<td>-0.03</td>
<td>0.05</td>
<td>0.11</td>
</tr>
<tr>
<td>poly(dGdC).poly(dGdC) - dsDNA</td>
<td>0.08</td>
<td>0.08</td>
<td>0.08</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(dA).poly(dT) - dsDNA</td>
<td>0.06</td>
<td>0.03</td>
<td>0.04</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(rA).poly(dT) - DNA-RNA hybrid</td>
<td>0.07</td>
<td>-0.06</td>
<td>0.01</td>
<td>0.09</td>
</tr>
<tr>
<td>poly(rA).poly(rU) - dsRNA</td>
<td>0.11</td>
<td>-0.04</td>
<td>0.03</td>
<td>0.11</td>
</tr>
<tr>
<td>poly(dAdT).poly(dAdT) - dsDNA</td>
<td>-0.03</td>
<td>-0.02</td>
<td>-0.03</td>
<td>0.00</td>
</tr>
<tr>
<td>(dA)_32.(dT)_32 - dsDNA</td>
<td>0.02</td>
<td>-0.02</td>
<td>0.00</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(rA) - ssRNA</td>
<td>0.01</td>
<td>-0.06</td>
<td>-0.03</td>
<td>0.05</td>
</tr>
<tr>
<td>poly(dT) - ssDNA</td>
<td>0.02</td>
<td>-0.08</td>
<td>-0.03</td>
<td>0.07</td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>tank</td>
<td>0.09</td>
<td>0.06</td>
<td>0.07</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>0.05</td>
<td>-0.02</td>
<td>0.01</td>
<td>0.05</td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>-0.02</td>
<td>0.00</td>
<td>-0.01</td>
<td>0.01</td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration (C_b/C_f) was determined by: C_b/C_f=(C_t/C_f)-1 where C_t (total ligand concentration) is the concentration of the ligand in each dialysis cassette and C_f (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Azaisopropyl’s behavior was significantly different than the azacyanines which bear a linear alkyl chain in their benzimidazole ring. It did not show any selectivity to any kind of nucleic acid structure. The amount of the Azaisopropyl concentration in dialysis cassettes containing different nucleic acid structures was no different than the control dialysis cassette containing only the buffer solution. Since the data showed normal distribution (Kolmogorov-Smirnov; p=0.200 and Shapiro-Wilk; p=0.620), we performed one-way ANOVA following Bonferroni and Sidak post-hoc tests in here.
and confirmed that the binding of Azaisopropyl to any of the nucleic acids structures was not statistically significant.

3.3.6. Azaisobutyl:

Average fluorescence spectra of Azaisobutyl samples obtained at the end of the 24 hours long competition dialysis from dialysis cassettes containing different nucleic acid structures were given in Figure 29. $C_b/C_f$ values are given in Table 7 and statistical analysis of the samples was plotted in Figure 30.

![Figure 29](image_url)

**Figure 29.** Average fluorescence spectra of Azaisobutyl. Measurements were obtained at the end of the 24 hours long competition dialysis against Azabutyl from dialysis cassettes containing different nucleic acid structures
Table 7. Competition dialysis results for Azaisobutyl.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>Folds over buffer (C_b/C_f)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1st replicate</td>
</tr>
<tr>
<td>poly(dA).[poly(dT)]_2 - triplex DNA</td>
<td>1.16</td>
</tr>
<tr>
<td>(dA)_32.([d(T)]_32]_2 - triplex DNA</td>
<td>0.58</td>
</tr>
<tr>
<td>5'-T_g20T_2 - 3'- quadruplex DNA</td>
<td>2.49</td>
</tr>
<tr>
<td>poly(rA).[poly(rU)]_2 - triplex RNA</td>
<td>0.05</td>
</tr>
<tr>
<td>M.lysodeiktus - dsDNA</td>
<td>0.31</td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>0.24</td>
</tr>
<tr>
<td>poly(dGdC).poly(dGdC) - dsDNA</td>
<td>0.22</td>
</tr>
<tr>
<td>poly(dA).poly(dT) - dsDNA</td>
<td>0.51</td>
</tr>
<tr>
<td>poly(rA).poly(dT) - DNA-RNA hybrid</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(rA).poly(rU) - dsRNA</td>
<td>0.06</td>
</tr>
<tr>
<td>poly(dAdT).poly(dAdT) - dsDNA</td>
<td>0.02</td>
</tr>
<tr>
<td>(dA)_32.(dT)_32 - dsDNA</td>
<td>0.33</td>
</tr>
<tr>
<td>poly(rA) - ssRNA</td>
<td>0.00</td>
</tr>
<tr>
<td>poly(dT) - ssDNA</td>
<td>0.10</td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
</tr>
<tr>
<td>tank</td>
<td>0.32</td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>0.04</td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>-0.12</td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration (C_b/C_f) was determined by: C_b/C_f=(C_t/C_f)-1 where C_t (total ligand concentration) is the concentration of the ligand in each dialysis cassette and C_f (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Figure 30. Competition Dialysis results for Azaisobutyl. *, p<0.05 from control by Kruskal Wallis non-parametric test. Error bars are ±1 standard deviations of the mean with a confidence level of 95%.

Our competition dialysis results for Azaisobutyl revealed that Azaisobutyl was not that different from Azaisopropyl in term of its binding affinity and selectivity towards different nucleic acid structures. It did not show any significant binding affinity to any of the structures we investigated as affirmed by the lack of a significant variation among C_b/C_f values (Table 7 and Figure 30). Azabutyl only showed slight binding to 5’-T_2G_20T_2-3’ quadruplex sequence with C_b/C_f value 2.50, but our statistical analysis revealed the binding was not significant (p-value=0.198).
3.3.7. Aza 4:

The nucleic acid affinity and selectivity of Aza4 and Aza5, which are benzothiazole derivatives, was also investigated using competition dialysis. The fluorescence spectra of the Aza4 samples obtained from each dialysis cassette were shown in Figure 31. The $C_b/C_f$ ratios for both of the trials along with the average $C_b/C_f$ values were given in Table 8 and the average $C_b/C_f$ values along with the statistical analysis were plotted in Figure 32.

Figure 31. Average fluorescence spectra of Aza4. Measurements were obtained at the end of the 24 hours long competition dialysis against Aza4 from dialysis cassettes containing different nucleic acid structures
Table 8. Competition dialysis results for Aza4.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>1&lt;sup&gt;st&lt;/sup&gt; replicate</th>
<th>2&lt;sup&gt;nd&lt;/sup&gt; replicate</th>
<th>Average</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>poly(dA),[poly(dT)]&lt;sub&gt;2&lt;/sub&gt; - triplex DNA</td>
<td>10.23</td>
<td>10.94</td>
<td>10.59</td>
<td>0.50</td>
</tr>
<tr>
<td>(dA)&lt;sub&gt;32&lt;/sub&gt;,(dT)&lt;sub&gt;32&lt;/sub&gt; - triplex DNA</td>
<td>2.19</td>
<td>1.94</td>
<td>2.06</td>
<td>0.18</td>
</tr>
<tr>
<td>5’-T&lt;sub&gt;2&lt;/sub&gt;G&lt;sub&gt;20&lt;/sub&gt;T&lt;sub&gt;2&lt;/sub&gt; - 3’- quadruplex DNA</td>
<td>0.97</td>
<td>0.88</td>
<td>0.92</td>
<td>0.06</td>
</tr>
<tr>
<td>poly(rA),[poly(rU)]&lt;sub&gt;2&lt;/sub&gt; - triplex RNA</td>
<td>0.25</td>
<td>0.20</td>
<td>0.23</td>
<td>0.03</td>
</tr>
<tr>
<td>M.lysodeiktus - dsDNA</td>
<td>0.45</td>
<td>0.39</td>
<td>0.42</td>
<td>0.04</td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>0.33</td>
<td>0.34</td>
<td>0.33</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(dGdC),poly(dGdC) - dsDNA</td>
<td>0.18</td>
<td>0.19</td>
<td>0.19</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(dA),poly(dT) - dsDNA</td>
<td>0.67</td>
<td>0.66</td>
<td>0.66</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(rA),poly(dT) - DNA-RNA hybrid</td>
<td>0.00</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(rA),poly(rU) - dsRNA</td>
<td>0.06</td>
<td>0.04</td>
<td>0.05</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(dAdT),poly(dAdT) - dsDNA</td>
<td>0.05</td>
<td>0.02</td>
<td>0.04</td>
<td>0.02</td>
</tr>
<tr>
<td>(dA)&lt;sub&gt;32&lt;/sub&gt;,(dT)&lt;sub&gt;32&lt;/sub&gt; - dsDNA</td>
<td>0.10</td>
<td>0.02</td>
<td>0.06</td>
<td>0.06</td>
</tr>
<tr>
<td>poly(rA) - ssRNA</td>
<td>0.01</td>
<td>-0.01</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(dT) - ssDNA</td>
<td>0.05</td>
<td>0.02</td>
<td>0.04</td>
<td>0.02</td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>tank</td>
<td>0.06</td>
<td>0.01</td>
<td>0.04</td>
<td>0.04</td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>0.06</td>
<td>-0.01</td>
<td>0.02</td>
<td>0.05</td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>0.03</td>
<td>-0.05</td>
<td>-0.01</td>
<td>0.06</td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration ($C_b/C_f$) was determined by: $C_b/C_f = (C_t/C_f) - 1$ where $C_t$ (total ligand concentration) is the concentration of the ligand in each dialysis cassette and $C_f$ (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Aza4 also showed significant affinity and selectivity towards poly(dA)[poly(dT)]_2 and (dA)_32[(dT)_32]_2 triplex DNA structures with the average \( C_b/C_f \) values of 10.59 and 2.06 with the significance levels of \( p=0.015 \) and \( p=0.020 \), respectively, according to non-parametric Kruskal–Wallis test. It’s binding affinity to poly(dA)[poly(dT)]_2 triplex DNA was very similar to Azamethyl’s. On the other hand, its binding affinity to (dA)_32[(dT)_32]_2 triplex DNA was significantly lower than Azamethyl’s. Additionally, its binding to 5’-T_2G_20T_2-3’ quadruplex and poly(dA).poly(dT)-dsDNA was also significant according to our statistical analysis.
with $p=0.027$ and $p=0.038$, respectively. Its binding to any other sequence was not statistically significant.

### 3.3.8. Aza5:

The fluorescence spectra of the Aza5 samples in each dialysis cassette were depicted in Figure 33. The concentration of Aza5 was also found to be different in each dialysis cassette Table 9. The average $C_b/C_f$ values with the statistical analysis were plotted in Figure 34.

![Figure 33. Average fluorescence spectra of Aza5. Measurements were obtained at the end of the 24 hours long competition dialysis against Aza5 from dialysis cassettes containing different nucleic acid structures](image-url)
Table 9. Competition dialysis results for Aza5.

<table>
<thead>
<tr>
<th>Oligonucleotides</th>
<th>1st replicate</th>
<th>2nd replicate</th>
<th>Average</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>poly(dA)].[poly(dT)]₂ - triplex DNA</td>
<td>11.99</td>
<td>12.58</td>
<td>12.29</td>
<td>0.42</td>
</tr>
<tr>
<td>(dA)$<em>{32}$.[(dT)$</em>{32}$]₂ - triplex DNA</td>
<td>5.48</td>
<td>5.38</td>
<td>5.43</td>
<td>0.07</td>
</tr>
<tr>
<td>5'-T₃G₂₀T₂ - 3'- quadruplex DNA</td>
<td>4.07</td>
<td>4.13</td>
<td>4.10</td>
<td>0.04</td>
</tr>
<tr>
<td>poly(rA).[poly(rU)]₂ - triplex RNA</td>
<td>1.90</td>
<td>2.00</td>
<td>1.95</td>
<td>0.07</td>
</tr>
<tr>
<td>M.lysodeiktus - dsDNA</td>
<td>1.06</td>
<td>1.17</td>
<td>1.11</td>
<td>0.08</td>
</tr>
<tr>
<td>tel24 - quadruplex DNA</td>
<td>2.32</td>
<td>2.54</td>
<td>2.43</td>
<td>0.16</td>
</tr>
<tr>
<td>poly(dGdC).poly(dGdC) - dsDNA</td>
<td>0.49</td>
<td>0.64</td>
<td>0.56</td>
<td>0.11</td>
</tr>
<tr>
<td>poly(dA).poly(dT) - dsDNA</td>
<td>0.80</td>
<td>0.83</td>
<td>0.82</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(rA).poly(dT) - DNA-RNA hybrid</td>
<td>0.29</td>
<td>0.51</td>
<td>0.40</td>
<td>0.16</td>
</tr>
<tr>
<td>poly(rA).poly(rU) - dsRNA</td>
<td>0.06</td>
<td>0.12</td>
<td>0.09</td>
<td>0.05</td>
</tr>
<tr>
<td>poly(dAdT).poly(dAdT) - dsDNA</td>
<td>0.03</td>
<td>0.14</td>
<td>0.08</td>
<td>0.08</td>
</tr>
<tr>
<td>(dA)$<em>{32}$.[(dT)$</em>{32}$] - dsDNA</td>
<td>0.07</td>
<td>0.07</td>
<td>0.07</td>
<td>0.00</td>
</tr>
<tr>
<td>poly(rA) - ssRNA</td>
<td>0.01</td>
<td>-0.04</td>
<td>-0.01</td>
<td>0.04</td>
</tr>
<tr>
<td>poly(dT) - ssDNA</td>
<td>-0.04</td>
<td>-0.01</td>
<td>-0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>buffer</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>tank</td>
<td>0.02</td>
<td>0.05</td>
<td>0.04</td>
<td>0.02</td>
</tr>
<tr>
<td>poly(dA) - ssDNA</td>
<td>-0.04</td>
<td>-0.06</td>
<td>-0.05</td>
<td>0.01</td>
</tr>
<tr>
<td>poly(rU) - ssRNA</td>
<td>-0.03</td>
<td>-0.02</td>
<td>-0.03</td>
<td>0.00</td>
</tr>
</tbody>
</table>

The ratio of the bound ligand to free ligand concentration ($C_b/C_f$) was determined by: $C_b/C_f = (C_t/C_f) - 1$ where $C_t$ (total ligand concentration) is the concentration of the ligand in each dialysis cassette and $C_f$ (free ligand concentration) is the concentration of the ligand in a positive control dialysis cassette (buffer) that only includes 1X BPES buffer which typically did not significantly vary from initial small molecule concentration that was 1.0 µM in the solution (tank).
Our competition dialysis results revealed undoubtedly that Aza5 is also binding to triple helical structures of poly(dA).[poly(dT)]_2 and (dA)_32.[(dT)_32]_2 with very high affinity and selectivity (Figure 33-34, Table 9). The amount of Aza5 collected in the dialysis cassettes containing poly(dA).[poly(dT)]_2 and (dA)_32.[(dT)_32]_2 was substantial. The average $C_b/C_f$ values of 12.29 and 5.43 with significance levels of $p=0.007$ and $p=0.009$, were obtained respectively, according to non-parametric Kruskal–Wallis test.

Compared to Azamethyl, its binding affinity to poly(dA).[poly(dT)]_2 triplex DNA structure was slightly higher, whereas its binding affinity to (dA)_32.[(dT)_32]_2 was lower. Aza5 also showed statistically significant binding to 5’-T2G20T2-3’ quadruplex, tel24,
poly(rA).[poly(rU)]₂ RNA triplex, *Micrococcus Lysodeiktus* duplex DNA, poly(dA).poly(dT) duplex DNA, poly(dGdC).poly(dGdC) duplex DNA, and poly(rA).poly(dT) DNA-RNA duplex hybrid with $C_b/C_f$ values of 4.10, 2.43, 1.95, 1.11, 0.82, 0.56, and 0.40 respectively. Furthermore, Aza5 had no affinity towards single stranded nucleic acid structures under these conditions, where the obtained $C_b/C_f$ values were within the experimental errors. Aza5 showed the highest selectivity towards triplex and quadruplex structures among all azacyanines studied here.

### 3.4. Conclusions:

Here we have examined the affinity and selectivity of eight azacyanines towards different nucleic acid sequences and structures. Six of them were benzimidazole derivatives differing in alky chain length and branching on the benzimidazole ring from each other. Two of them were benzothiazole derivatives differing in substitution on the sixth carbon of the benzimidazole ring from each other.

Our results revealed that Azamethyl, Azaethyl and Azapropyl possess high selectivity and binding affinity towards poly(dA).[poly(dT)]₂ – triplex DNA structure. Especially Azamethyl’s affinity towards poly(dA).[poly(dT)]₂ was remarkable. $C_b/C_f$ value was 11.7 ($p = 0.012$) meaning that 11.7 μM Azamethyl was accumulated to the dialysis cassette that contains poly(dA).[poly(dT)]₂ due to high binding affinity where the free Azamethyl concentration was only about 1.0 μM outside. Azabutyl was also binding poly(dA).[poly(dT)]₂ with relatively high affinity. However, it was not really selective towards poly(dA).[poly(dT)]₂. Its affinity towards poly(dA).[poly(dT)]₂ triplex DNA structure and 5’-T₂G₂₀T₂- 3’ quadruplex DNA structure were similar. Overall, the affinity of benzimidazole derivatives towards poly(dA).[poly(dT)]₂ triplex DNA structure was decreasing with the increasing alky chain length. The order of affinity was Azamethyl > Azaethyl > Azapropyl > Azabutyl. In other words, the effect of molecular structure on binding ability of the small molecules to nucleic acid structures was crucial.
Also for these azacyanines with the linear alkyl chain length in the benzimidazole ring, we observed that their affinity towards DNA structures was higher than their affinity to RNA structures. Moreover, their affinity towards DNA-RNA hybrid structures was in between. We also observed that none of the azacyanines showed significant binding affinity towards any of the single stranded nucleic acid structures under these conditions.

Azaisopropyl and Azaisobutyl, which bear a branched alkyl chain in their benzimidazole ring, showed no significant binding affinity statistically to any of the nucleic acid structures examined. Especially, in the case of Azaisopropyl there was no binding at all. It was evident that the branching on the benzimidazole ring was detrimental, preventing these molecules from binding to any of the nucleic acid structures included in our assay.

Based on benzothiazole derivatives, our results revealed that Aza4 and Aza5 also possess high selectivity and binding affinity towards poly(dA).[poly(dT)]2 – triplex DNA structure. Especially Aza5’s affinity towards poly(dA).[poly(dT)]2 was remarkable. \( C_b/C_f \) value was 12.3 (\( p = 0.007 \)). Aza5 showed the most dramatic results in terms of its affinity and selectivity towards triplex poly(dA).[poly(dT)]2 triplex DNA structure among all of the azacyanine derivatives.

Aza4 also showed high affinity and selectivity towards poly(dA).[poly(dT)]2 triplex DNA structure. \( C_b/C_f \) value was 10.6 with the \( p \) value: 0.015. Furthermore, its binding to (dA)\(_{32}\).[(dT)\(_{32}\)]\(_2\) triplex DNA was also noteworthy with the \( C_b/C_f \) value 2.06 (\( p = 0.020 \)).

Previously, using surface plasma resonance (SPR) spectroscopy it has been shown that the binding affinity of Azamethyl, Aza4 and Aza5 towards G-quadruplex- tel 24 structure were quite similar (Çetinkol \& al., 2008). Furthermore, in term of their binding affinity to poly(A) at high drug concentrations, Azamethyl and Aza5 had similar binding affinities but the binding affinity of Aza4 was quite lower (Çetinkol & Hud, 2009). Moreover, in his studies Kim revealed that Aza5 showed the most
remarkable effect on yeast cells with (GAA)$_n$ repeats. In addition, Kim hypothesized that the effect they observed was due to the binding and the stabilizing effect of azacyanines to triple helical nucleic acids (Kim, 2009). Here, our results confirmed the studies of Kim that azacyanines were binding to poly(dA).[poly(dT)]$_2$ triplex DNA structure with high affinity and selectivity. Their affinity order was Aza5 > Azamethyl > Aza4 in an agreement with Kim’s data (Kim, 2009). In terms of their binding abilities to tel24, even though their binding affinity is way lower compared to their affinity to poly(dA).[poly(dT)]$_2$ triplex DNA, we observed a very similar trend. Aza5 had slightly higher affinity than Azamethyl, and Azamethyl had slightly higher affinity than Aza4.

In conclusion, our competition dialysis results revealed that certain azacyanine molecules possess high affinity and selectivity towards triplex and quadruplex DNA structures. More specifically Aza5, Azamethyl, and Aza4 showed the most dramatic binding selectivity towards poly(dA).poly(dT)$_2$ triplex DNA structure. Their ability of binding to other triplex and quadruplex structures was also statistically significant. Moreover, our results showed that, among the benzimidazole derivatives, the binding affinity of the small molecules towards poly(dA).poly(dT)$_2$ triplex DNA was decreasing with the increasing alkyl chain length and branching. Especially the effect of branching on binding affinity of these molecules was significant. Neither Azaisopropyl nor Azaisobutyl showed significant binding affinity towards any of the nucleic acid structures included in our competition dialysis assay.
CHAPTER 4

THESIS CONCLUSION

4.1. Azamethyl, Aza4 and Aza5 Catalytically Inhibit Topo IIα:

Here, we first investigated the effect of different azacyanines on Topo IIα. Our results revealed that Azamethyl, Aza4, and Aza5 can decrease the catalytic activity of Topo IIα with high efficiency. More importantly, their efficiency was higher than the proven Topo IIα poison (Etoposide). Etoposide (also known as Toposar and VePesid) is one of the medications used in chemotherapy and included in the World Health Organization’s “List of Essential Medicines” in 2015 as an anticancer drug (World Health Organization, 2015).

In our assay, we have observed that 50.0 µM Azamethyl decrease Topo IIα catalytic activity even more strongly than 500.0 µM of Etoposide. However, we have observed that Azamethyl affects Topo IIα catalytically. In other words, azacyanines cannot be considered as poisons, like Etoposide. This means that they are probably less toxic than the Etoposide. The lower toxicity of azacyanines might have a significant impact on their potential use in chemotherapy. Even though Etoposide is a commonly used medicine, it cannot be used in high doses due to its highly toxic side effects to normal cells which limit its effectiveness in chemotherapy (Marigny, Aubin, Burgot, Le Gall, & Gandemer, 2005). Therefore, azacyanines with low toxicity, meaning lower side effects, might also be good choices to inhibit Topo IIα in chemotherapeutic applications.
Unlike Azamethyl, Aza5, and Aza4; Azaethyl and Azaisobutyl did not change Topo IIα’s activity. Surprisingly, they both strengthen the intensity of the bands belonging to decatenated DNA products in our gel. This indicates that they do not decrease the catalytic activity of Topo IIα enzyme but rather increase its activity. However, this outcome requires further detailed investigation.

The results obtained here provide an alternative insight about the mechanistic explanation of azacyanine derivatives’ effects in yeast cells observed also by Kim (Kim, 2009) and demonstrate the potentials of Azamethyl, Aza4 and Aza5. However, their effect on different types of cancer cells should also be examined by in-vivo studies. Only after such examinations, their real potential as Topo IIα interfering molecules will be truly realized.

4.2. Azamethyl, Aza4, and Aza5 Bind to Triplex and Quadruplex Nucleic Acid Structures with High Affinity and Selectivity:

Our competition dialysis results showed that Azamethyl, Aza5, and Aza4 bind with high affinity and selectivity especially to poly(dA).[poly(dT)]_2 - triplex DNA and 5’-T_2G_20T_2- 3’ - quadruplex DNA structures. In addition to these molecules, Azaethyl, Azapropyl, and Azabutyl also showed differential binding affinity towards the same triplex and quadruplex structures. On the other hand, Azaisopropyl and Azaisobutyl did not show any binding affinity towards any of the nucleic acid structures examined here.

In terms of their triplex binding affinity, Aza5 showed the highest binding affinity. Its C_v/C_f values were 12.3 (p=0.007) and 5.4 (p=0.009) towards poly(dA).poly[(dT)]_2 and (dA)_32.[(dT)_32]_2 triplex DNA structures respectively. This means that Aza5 binds to poly(dA).poly(dT)_2 triplex DNA sequence 12.3 folds more than our control (includes only buffer). Its affinity to quadruplex structures was also high with C_v/C_f values of 4.1 (p=0.011) and 2.43 (p=0.013) towards 5’- T_2G_20T_2 – 3’ and tel24 quadruplex DNAs respectively. These findings are important because as explained in chapter 1.1, stabilizing triplex and quadruplex structures could be a promising strategy in cancer treatment.
Moreover, Azamethyl also showed high affinity and selectivity towards triplex and quadruplex DNA structures. Its $C_b/C_f$ values were 11.7 ($p=0.012$) and 10.7 ($p=0.014$) towards poly(dA).poly[(dT)$_2$] and (dA)$_{32}$.[(dT)$_{32}$]$_2$ triplex DNA structures, respectively. Furthermore, Aza4 also showed relatively high affinity towards triplex DNA structures. Azaethyl, Azapropyl, and Azabutyl also showed high affinity and selectivity towards poly(dA).poly[(dT)$_2$] triplex DNA structure but their binding affinity and selectivity were not as high as Azamethyl’s, Aza4’s, or Aza5’s. On the other hand, Azaisopropyl and Azaisobutyl showed no notable affinity towards any nucleic acid structures investigated here.

Among the benzimidazole derivatives (Azamethyl, Azaethyl, Azapropyl, Azabutyl, Azaisopropyl, and Azaisobutyl) investigated, it is obvious that the affinity and selectivity of these molecules towards poly(dA).poly[(dT)$_2$] triplex DNA structure is highly dependent on the linear alkyl chain length and branching. With the increasing length of linear alkyl chain and/or branching, the binding ability of azacyanines to poly(dA).poly[(dT)$_2$] triplex DNA structure was decreasing. To our knowledge, this is the first work displaying how the systematic change in molecular structure of the small molecule can alter the triplex binding affinity and selectivity.

Our competition dialysis results also provide an alternative explanation about the mechanism of azacyanines’ effects on cells observed previously by Kim (Kim, 2009). He hypothesized that the reason behind the observed double stranded breaks in yeast cells might be due to the high binding affinity and selectivity of Azamethyl, Aza4 and Aza5 toward triplexes. Here, we have clearly demonstrated that all three molecules have high affinity and selectivity towards certain triplex structures.

4.3. Concluding Remarks:

In conclusion, we have found two probable explanations about the observed effects of Azamethyl, Aza4 and Aza5 in yeast cells demonstrated by Kim (Kim, 2009).
The first explanation is the ability of these molecules to change the catalytic activity of Topo IIα. Our results explicitly revealed that catalytic activity of Topo IIα decreased with Azamethyl, Aza4, and Aza5. Moreover, the effect was significantly higher than the inhibition ability of Etoposide which is a proven Topo IIα poison. Therefore, these molecules might have a high potential to be effective chemotherapeutic agents. However, further investigation on their effect in different cancer cell lines are needed to realize their full potential.

Our second explanation, which supports the findings and hypothesizes of Kim (Kim, 2009), is the fact that these molecules have high binding affinity and selectivity especially towards certain triplex DNA structures. They were also shown to have relatively high binding affinity towards certain quadruplex structures.

At this point we do not know exactly at which catalytic step (Figure 7) Topo IIα is affected, but we suspect that the binding of these molecules (Azamethyl, Aza4, and Aza5) to higher order nucleic acid structures might be the reason behind their effects on Topo IIα’s catalytic activity. These molecules might be binding to enzyme binding sites on the DNA before the enzyme and therefore preventing the formation of an intact DNA-enzyme complex by changing the DNA conformation or they might be directly targeting the enzyme and blocking its binding to the DNA.

Overall, these molecules might have potential to be used in chemotherapy due to their negative effects on Topo IIα’s catalytic activity and can also be used in gene targeting due to their triplex binding affinity. The findings presented in this thesis, inspire us to further investigate azacyanines’ potentials.
REFERENCES


Overexpression of RNase H partially complements the growth defect of an Escherichia coli delta topA mutant: R-loop formation is a major problem in the absence of DNA topoisomerase I. *Proceedings of the National Academy of Sciences of the United States of America*, 92(8), 3526–3530.


51(8), 1730–1739.


1315–1324.


APPENDIX A

PREPARATION OF BUFFERS AND STOCK SOLUTIONS

1X BPES buffer (pH:7.0) solution for competition dialysis experiments:

6.0 mM Na$_2$HPO$_4$, 2.0 mM NaH$_2$PO$_4$, 1.0 mM Na$_2$EDTA, 185.0 mM NaCl.

Concentrations were arranged by using following equation;

$$M_1 \times V_1 = M_2 \times V_2$$

Thus for 1.0 L of 1X BPES buffer, following amounts of ingredients were mixed. Preparation of each ingredient will be given next.

- Na$_2$HPO$_4$ : 300 mM (stock) x $V_1$= 6 mM x 1000 mL; $V_1 = 20.0$ mL,
- NaH$_2$PO$_4$ : 400 mM (stock) x $V_1$= 2 mM x 1000 mL; $V_1 = 5.0$ mL,
- Na$_2$EDTA : 100 mM (stock) x $V_1$= 1 mM x 1000 mL; $V_1 = 10.0$ mL,
- NaCl : 500 mM (stock) x $V_1$= 185 mM x 1000 mL; $V_1 = 370.0$ mL,
- 500.0 mL deionized water added to 405.0 mL solution,
- pH was adjusted to 7.0 by NaOH and/or HCl,
- Total volume was arranged to 1.0 L with deionized (Millipore) water.

300 mM Na$_2$HPO$_4$ stock solution for 1X BPES buffer:

For 300.0 mM; 100.0 mL stock solution 4.26 gram was needed. Calculations are below;

$$n \text{ (mol)} = \text{molarity (M) } \times \text{volume (L)}$$

$$0.3 \text{ M} \times 0.1 \text{ L} = 0.03 \text{ mol.}$$
m = mol x \( M_w \) (141.96 g/mol for Na\(_2\)HPO\(_4\))

\[ 0.03 \times 141.96 = 4.26 \text{ g.} \]

**400 mM Na\(_2\)HPO\(_4\) stock solution for 1X BPES buffer:**

For 400.0 mM; 100.0 mL stock solution 4.80 gram was needed. Calculations are below;

\[ n \text{ (mol)} = \text{molarity (M)} \times \text{volume (L)} \]

\[ 0.4 \text{ M} \times 0.1 \text{ L} = 0.04 \text{ mol.} \]

\[ m = \text{mol} \times \text{M}_w \text{ (119.98 g/mol for NaH}_2\text{PO}_4\) \]

\[ 0.04 \times 119.98 = 4.80 \text{ g.} \]

**100 mM Na\(_2\)EDTA stock solution for 1X BPES buffer:**

For 100.0 mM; 100.0 mL stock solution 3.72 gram was needed. Calculations are below;

\[ n \text{ (mol)} = \text{molarity (M)} \times \text{volume (L)} \]

\[ 0.1 \text{ M} \times 0.1 \text{ L} = 0.01 \text{ mol.} \]

\[ m = \text{mol} \times \text{M}_w \text{ (372.24 g/mol for Na}_2\text{EDTA.2H}_2\text{O\) \]

\[ 0.01 \times 372.24 = 3.72 \text{ g.} \]

pH of the solution must be 8.0 and arranged with concentrated NaOH and/or HCl.

**500 mM NaCl stock solution for 1X BPES buffer:**

For 500.0 mM; 1.0 L stock solution 29.22 gram was needed. Calculations are below;

\[ n \text{ (mol)} = \text{molarity (M)} \times \text{volume (L)} \]

\[ 0.5 \text{ M} \times 1.0 \text{ L} = 0.5 \text{ mol.} \]

\[ m = \text{mol} \times \text{M}_w \text{ (58.44 g/mol for NaCl\) \]

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10% (w/v) SDS stock solution for competition dialysis experiments:
10.0 g SDS (Sigma Aldrich, 862010) dissolved in 80.0 mL deionized water. Then volume was completed to 100.0 mL with deionized (Millipore) water.

1% Agarose Gel for Topoisomerase kit visualization:
1.00 g agarose (VWR-Amresco RA, N605) was dissolved in 100.0 mL of 1X TAE buffer in a microwave oven. After cooling down, 0.5 µg/mL of EtBr was added. Next, the gel was poured into “B2” Thermo Fischer Owl Easycast Mini Gel Electrophoresis system and left for the complete polymerization of agarose for about 30 to 40 minutes. 1X TAE buffer was poured on the gel and into the tray as a running buffer. Later, the samples were loaded into the wells. The gel was run at 100 Volt for 3 hours. Afterwards, the gel was stained with 1.0 µg/mL EtBr-water mixture for about 40 minutes followed by 30 minutes of de-staining with distilled water. Finally, the image of the gel was taken using BioRad XR+ molecular gel imager system.

1X TAE (Tris Acetate EDTA) buffer (pH:8.5) solution for agarose gel electrophoresis experiments:
40.0 mM Tris (pH: 7.6), 20.0 mM acetic acid, and 1.0 mM EDTA.
50X TAE buffer was prepared and diluted to 1X TAE buffer,
1X TAE buffer (1.0 L): 20 mL of 50X TAE buffer and 980 mL Millipore deionized water.

Preparation of 50X TAE buffer:
For 100.0 mL 50X TAE buffer, following amounts of ingredients were mixed. Preparation of ingredients will be given next.

- 24.20 g Tris base (M_w: 121.14 g/mol),
- 50.0 mL deionized water to dissolve Tris base,
• 5.71 mL glacial acetic acid,
• 10.0 mL; 0.5 M (pH:8.0) EDTA,
• Arrange pH around 8.5 using concentrated NaOH and/or HCl,
• Complete total volume to 100.0 mL with deionized water.

For 1X TAE buffer, dilute 20.0 mL of 50X TAE buffer with 980.0 mL deionized water.

**0.5 M, pH: 8.0 EDTA solution for 50X TAE buffer:**

For 500.0 mM; 100.0 mL stock solution 18.61 gram was needed. Calculations are below;

\[
\text{n (mol) = molarity (M) x volume (L)}
\]

\[
0.5 \text{ M} \times 0.1 \text{ L} = 0.05 \text{ mol.}
\]

\[
m = \text{mol} \times M_w (372.24 \text{ g/mol for Na}_2\text{EDTA}.2\text{H}_2\text{O})
\]

\[
0.05 \times 372.24 = 18.61 \text{ g.}
\]

**10.0 mM, pH: 7.5 Tris-HCl solution for drug dilutions:**

For 10.0 mM; 100.0 mL stock solution 0.12 gram Tris and 20.0 μL 11.0 M HCl were needed. Calculations are below;

\[
\text{n (mol) = molarity (M) x volume (L)}
\]

\[
0.01 \text{ M} \times 0.1 \text{ L} = 0.001 \text{ mol.}
\]

\[
m = \text{mol} \times M_w (121.1 \text{ g/mol for Tris})
\]

\[
0.01 \times 121.1 = 0.12 \text{ g. pH was arranged to 7.52 with HCl.}
\]

**EtBr solution for staining of nucleic acids:**

2.80 mg EtBr (Sigma Aldrich, E1510) was dissolved in 1.40 mL deionized water as a stock solution (2.0 μg/μL). To have 0.5 μg/mL EtBr in a gel and 1.0 μg/mL in staining water, following amounts of EtBr was added to the gel and staining water;
For 100.0 mL gel, we put 25.0 µL of EtBr stock solution into 100.0 mL 1X TAE buffer thus 50.0 µg EtBr will be in the gel. At the end, it makes 0.5 µg/mL.

For 400.0 mL staining water, we put 200.0 µL of EtBr into 400.0 mL distilled water therefore 400.0 µg EtBr will be in the water. At the end, it makes 1.0 µg/mL for staining.
OLIGONUCLEOTIDES USED IN COMPETITION DIALYSIS

Oligonucleotides given in Table B.1 were purchased from different companies and used throughout the competition dialysis experiments. Detailed instruction for preparation of double stranded oligonucleotides, DNA-RNA hybrid, RNA double stranded oligonucleotides, triplex DNA structures, and quadruplex DNA structures are given in chapter 3.2. Extinction coefficient values (ε) were either taken from the literature or purchased company. Monomeric units (nucleotide, base pair, triplet, and quartet) were used to prepare 75.0 μM of stock solutions.
Table B.1. Oligonucleotides used throughout the competition dialysis experiments.

<table>
<thead>
<tr>
<th>Conformation</th>
<th>Index</th>
<th>Nucleic Acid</th>
<th>λ (nm)</th>
<th>ε (M⁻¹·cm⁻¹)</th>
<th>Monomeric Unit</th>
<th>Company</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single-stranded DNA</td>
<td>1</td>
<td>poly(dA)</td>
<td>257</td>
<td>8600ᵃ</td>
<td>Nucleotide</td>
<td>Sigma-Aldrich P0887</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>poly(dT)</td>
<td>264</td>
<td>8520ᵃ</td>
<td>Nucleotide</td>
<td>Sigma-Aldrich P6905</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>poly(rA)</td>
<td>258</td>
<td>9800ᵃ</td>
<td>Nucleotide</td>
<td>Merck</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>poly(rU)</td>
<td>260</td>
<td>9350ᵃ</td>
<td>Nucleotide</td>
<td>Merck</td>
</tr>
<tr>
<td>Single-stranded RNA</td>
<td>5</td>
<td>(dA)₃₂.(dT)₃₂⁺</td>
<td>260</td>
<td>(dA)₃₂: 387400ᵇ (dT)₃₂: 259800ᵇ</td>
<td>Base pair</td>
<td>IDT</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>poly(dA).poly(dT)</td>
<td>260</td>
<td>12000ᵃ</td>
<td>Base pair</td>
<td>Sigma-Aldrich P9764</td>
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<td>Double-stranded DNA</td>
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<td>poly(dAdT).poly(dAdT)</td>
<td>262</td>
<td>13200ᵃ</td>
<td>Base pair</td>
<td>Sigma-Aldrich P0883</td>
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<tr>
<td></td>
<td>8</td>
<td>poly(dGdC).poly(dGdC)</td>
<td>254</td>
<td>16800ᵃ</td>
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<tr>
<td>DNA-RNA Hybrid</td>
<td>9</td>
<td>Microccoccus Lysodeikticus (72% GC)</td>
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<td>10</td>
<td>poly(rA).poly(dT)*</td>
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<td>Base pair</td>
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<td></td>
<td></td>
<td></td>
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<td>Double-stranded RNA</td>
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<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*a from (Ragazzon et al., 2007) ;b from Integrated DNA Technologies (IDT); *prepared from more than one nucleic acid – for detailed information see the section 3.2. Materials & Methods.
C.1. Standard Curve Preparation:

The standard curves used in determination of the small molecule concentrations were prepared for each small molecule (azacyanine) using Fluorescence Spectroscopy.

For each small molecule, samples with varying concentrations of the small molecule were prepared from a stock solution of 50.0 µM. The concentration of the 50.0 µM stock solution was determined via UV-Vis spectroscopy using the calculated extinction coefficients for each molecule (Table 1). Samples were prepared by taking appropriate volumes of 50.0 µM small molecule stock solution, 1X BPES buffer, and 10%SDS. As shown in Table C1, all components (1X BPES buffer and 10%SDS) were kept at the same volume except the added small molecule stock solution. A representative calculation used in the preparation of the first and the second sample were given below. The concentrations of all the other samples were calculated the same way and the samples were prepared accordingly:

\[ M_1 \times V_1 = M_2 \times V_2 \]

- 50.0 µM x \( V_1 = 0.1 \) µM x 2000.0 µL; \( V_1 = 4.0 \) µL.
- 50.0 µM x \( V_2 = 0.2 \) µM x \([2000+V_2 \) µL]; \( V_2 = 8.0 \) µL – 4.0 µL (already in the mixture) therefore 4.0 µL from the stock is added to get a 0.2 µM sample.
Table C1. Preparation of samples for the standard curve construction:

<table>
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<tr>
<th>Drug Stock Solution -50µM</th>
<th>Concentration (µM)</th>
<th>Added drug volume (µL)</th>
<th>Total drug volume (µL)</th>
<th>10% SDS (µL)</th>
<th>1X BPES buffer (µL)</th>
<th>Total volume (µL)</th>
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<td>1964.0</td>
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<td>245.8</td>
<td>32.0</td>
<td>1964.0</td>
<td>2241.8</td>
<td></td>
</tr>
</tbody>
</table>

Afterwards, the emission spectra for all samples were collected using Varian, Cary Eclipse fluorescence spectrometer using the parameters given below:

**Azamethyl, Azaethyl, Azapropyl, Azabutyl, Azaisopropyl, and Azaisobutyl;**


**Aza4;**


**Aza5;**

Later, two standard curves were constructed for each small molecule using the fluorescence intensities at two different wavelengths (reference wavelengths). Two equations from “fluorescence intensity vs. the concentration” graphs were derived for each sample by performing linear regression. These equations were used to calculate the concentrations of the small molecules in the samples obtained from each of the competition dialysis cassettes (section 3.3). The concentrations obtained for a sample, by using both of the standard curves, are averaged out, and recorded as the final concentration.

The reason behind choosing two different wavelengths was to minimize our experimental errors. The first wavelength we choose was the wavelength where the maximum absorbance was occurring. However, at that wavelength the fluorescence intensity was out of range for our concentrated samples hence we could not observe any absorbance for the first chosen wavelength. Therefore, we chose another wavelength whose intensity was not out of range for the small molecule concentration we have examined.

Since all azacyanine molecules have different chemical structures (Figure 12), obtained spectra from fluorescence spectrophotometer were different for some of the molecules. Therefore, different excitation wavelengths were chosen as a reference for each molecule’s standard curve constructions. Reference excitation wavelengths were 379 nm & 440 nm for Azamethyl, Azaethyl, Azapropyl, Azabutyl, Azaisopropyl, and Azaisobutyl, 445 nm & 465 nm for Aza4, and 475 nm & 520 nm for Aza5.

Small molecule concentrations of the samples obtained from 24 hours long competition dialysis (section 3.3) were calculated based on the equations obtained through the linear regression. Here, the measured fluorescence intensity belonging to a sample obtained from the dialysis cassette containing a certain nucleic acid structure was put in place of Y value on the equation after dilution corrections. Then, the concentrations corresponding to the X value on the equation were calculated.
C.2. Standard Curve for Azamethyl:

a. 

Figure C.1. Standard curve construction for Azamethyl. a. Fluorescence intensity vs wavelength measurements for different concentration of Azamethyl. b. The standard curve for Azamethyl at 379 nm. c. The standard curve for Azamethyl at 440 nm.
curve using the fluorescence intensities at 379 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 440 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.

C.3. Standard Curve for Azaethyl:

a.

b.
**Figure C.2.** Standard curve construction for Azaethyl. a. Fluorescence intensity vs wavelength measurements for different concentration of Azaethyl. b. The standard curve using the fluorescence intensities at 379 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 440 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.

**C.4. Standard Curve for Azapropyl:**

a.
Figure C.3. Standard curve construction for Azapropyl. a. Fluorescence intensity vs wavelength measurements for different concentration of Azapropyl. b. The standard curve using the fluorescence intensities at 379 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 440 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.
C.5. Standard Curve for Azabutyl:

a.

[Graph showing fluorescence intensity vs wavelength for different concentrations of Azabutyl]

b.

[Graph showing standard curve for Azabutyl at 379 nm, with the equation $y = 301.96x + 4.90$ and $R^2 = 0.999464$]

c.

[Graph showing standard curve for Azabutyl at 440 nm, with the equation $y = 144.31x + 11.45$ and $R^2 = 0.999241$]

**Figure C.4.** Standard curve construction for Azabutyl. a. Fluorescence intensity vs wavelength measurements for different concentration of Azabutyl. b. The standard
curve using the fluorescence intensities at 379 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 440 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.

C.6. Standard Curve for Azaisopropyl:

a. 

b. 

\[
y = 322.10 \times x + 13.90 \\
R^2 = 0.99902
\]
Figure C.5. Standard curve construction for Azaisopropyl. a. Fluorescence intensity vs wavelength measurements for different concentration of Azaisopropyl. b. The standard curve using the fluorescence intensities at 379 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 440 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.

C.7. Standard Curve for Azaisobutyl:

a.
Figure C.6. Standard curve construction for Azaisobutyl. a. Fluorescence intensity vs wavelength measurements for different concentration of Azaisobutyl. b. The standard curve using the fluorescence intensities at 379 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 440 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.
C.8. Standard Curve for Aza4:

a. 

![Graph a: Fluorescence intensity vs wavelength measurements for different concentration of Aza4.]

b. 

![Graph b: The standard curve for Aza4 at 445 nm.]

\[ y = 242.91x - 4.91 \]
\[ R^2 = 0.999965 \]

c. 

![Graph c: The standard curve for Aza4 at 465 nm.]

\[ y = 181.60x - 2.39 \]
\[ R^2 = 0.999904 \]

**Figure C.7.** Standard curve construction for Aza4. a. Fluorescence intensity vs wavelength measurements for different concentration of Aza4. b. The standard curve
using the fluorescence intensities at 445 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 465 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.

C.9. Standard Curve for Aza5:

a.

![Graph showing Aza5 fluorescence intensities at 5.5 μM and 0.1 μM](image)

b.

![Graph showing Aza5 standard curve at 475 nm with equation and R^2 value](image)
Figure C.8. Standard curve construction for Aza5. a. Fluorescence intensity vs wavelength measurements for different concentration of Aza5. b. The standard curve using the fluorescence intensities at 475 nm vs concentration (circles). c. The standard curve using the fluorescence intensities at 520 nm vs concentration (circles). The linear regression lines and the obtained equations were also given on the graphs in b and c.

Table C2. Equations derived from standard curves for each azacyanine molecule:

<table>
<thead>
<tr>
<th></th>
<th>379 nm</th>
<th>440 nm</th>
<th>445 nm</th>
<th>465 nm</th>
<th>475 nm</th>
<th>520 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azamethyl</td>
<td>y = 304.09.x + 5.43</td>
<td>y = 144.25.x + 13.65</td>
<td>y = 242.91.x – 4.92</td>
<td>y = 181.60.x – 2.39</td>
<td>y = 163.83.x + 5.29</td>
<td>y = 96.30.x + 2.77</td>
</tr>
<tr>
<td>Azaethyl</td>
<td>y = 231.74.x + 2.87</td>
<td>y = 116.92.x + 2.78</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azapropyl</td>
<td>y = 276.05.x + 10.08</td>
<td>y = 133.36.x + 8.20</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azabutyl</td>
<td>y = 301.96.x + 4.90</td>
<td>y = 144.31.x + 11.45</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azaisopropyl</td>
<td>y = 322.10.x + 13.90</td>
<td>y = 159.05.x + 16.98</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azaisobutyl</td>
<td>y = 387.90.x + 13.25</td>
<td>y = 168.15.x + 19.35</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azaisobutyl</td>
<td>y = 322.10.x + 13.90</td>
<td>y = 159.05.x + 16.98</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azaisobutyl</td>
<td>y = 387.90.x + 13.25</td>
<td>y = 168.15.x + 19.35</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
APPENDIX D

STATISTICAL ANALYSIS

The statistical analysis of our competition dialysis data is given in Table D1. The competition dialysis data (C_b/C_f values) represented here is the mean of two independent experiments (see section 3.3). In our statistical analysis confidence levels were taken as 99% and 95%. p<0.01 and p<0.05 were considered as significantly different from the positive control that is small molecule concentration in the cassette containing only the buffer solution. The Kruskal Wallis non-parametric test was performed for all molecules except Azaisopropyl. Since the Azaisopropyl data showed a normal distribution (Kolmorogov-Smirnow: p=0.200 and Shapiro-Wilk: p=0.620) one-way ANOVA followed by Bonferronni and Sidak Post-Hoc tests were performed for it.

In the Table D1 statistical analysis results for the azacyanines that are found to be binding significantly to a certain nucleic acid structure are presented. If the statistical analysis results were not given in the table below, it signifies that the binding of the small molecule to that particular nucleic acid structure was not found statistically significant. “Dashes” on the Table D1 means that the p values exceeded 0.05 and therefore were found as non-significant.
### Table D.1. Statistical analysis results of competition dialysis experiments:

<table>
<thead>
<tr>
<th>DNA Structure</th>
<th>Azamethyl C&lt;sub&gt;i&lt;/sub&gt;/C&lt;sub&gt;f&lt;/sub&gt;</th>
<th>p-value</th>
<th>Aza4 C&lt;sub&gt;i&lt;/sub&gt;/C&lt;sub&gt;f&lt;/sub&gt;</th>
<th>p-value</th>
<th>Aza5 C&lt;sub&gt;i&lt;/sub&gt;/C&lt;sub&gt;f&lt;/sub&gt;</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>poly(dA).poly(dT) triplex DNA</td>
<td>11.7</td>
<td>0.012</td>
<td>10.6</td>
<td>0.015</td>
<td>12.3</td>
<td>0.007</td>
</tr>
<tr>
<td>(dA)&lt;sub&gt;32&lt;/sub&gt;-(dT)&lt;sub&gt;32&lt;/sub&gt;2 triplex DNA</td>
<td>10.7</td>
<td>0.014</td>
<td>2.1</td>
<td>0.020</td>
<td>5.4</td>
<td>0.009</td>
</tr>
<tr>
<td>5'-T&lt;sub&gt;2&lt;/sub&gt;G&lt;sub&gt;20&lt;/sub&gt;T&lt;sub&gt;2&lt;/sub&gt;3' - quadruplex DNA</td>
<td>2.5</td>
<td>0.017</td>
<td>0.9</td>
<td>0.027</td>
<td>4.1</td>
<td>0.011</td>
</tr>
<tr>
<td>poly(rA).poly(rU) triplexRNA</td>
<td>1.7</td>
<td>0.021</td>
<td>-</td>
<td>-</td>
<td>1.9</td>
<td>0.016</td>
</tr>
<tr>
<td>Micrococcus Lysodeicticus- dsDNA</td>
<td>1.3</td>
<td>0.026</td>
<td>-</td>
<td>-</td>
<td>1.1</td>
<td>0.021</td>
</tr>
<tr>
<td>5'-TTGGG [TTAGGG]&lt;sub&gt;3&lt;/sub&gt;A - 3' quadruplexDNA</td>
<td>1.1</td>
<td>0.032</td>
<td>-</td>
<td>-</td>
<td>2.4</td>
<td>0.013</td>
</tr>
<tr>
<td>poly(dGdC).poly(dGdC) dsDNA</td>
<td>1.1</td>
<td>0.038</td>
<td>-</td>
<td>-</td>
<td>0.6</td>
<td>0.033</td>
</tr>
<tr>
<td>poly(dA).poly(dT) dsDNA</td>
<td>0.9</td>
<td>0.047</td>
<td>0.7</td>
<td>0.038</td>
<td>0.8</td>
<td>0.026</td>
</tr>
</tbody>
</table>
Table D.1. Statistical analysis results of competition dialysis experiments:

<table>
<thead>
<tr>
<th></th>
<th>Azaethyl</th>
<th>Azapropyl</th>
<th>Azabutyl</th>
<th>Azaisopropyl</th>
<th>Azaisobutyl</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C_b/C_f</td>
<td>p-value</td>
<td>C_b/C_f</td>
<td>p-value</td>
<td>C_b/C_f</td>
</tr>
<tr>
<td>poly(dA).[poly(dT)]_2 triplex DNA</td>
<td>6.9</td>
<td>0.026</td>
<td>5.2</td>
<td>0.024</td>
<td>4.9</td>
</tr>
<tr>
<td>(dA)_32.[(dT)_32]_2 triplex DNA</td>
<td>1.4</td>
<td>0.033</td>
<td>0.8</td>
<td>0.041</td>
<td>-</td>
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<tr>
<td>5'-T_G20T_3' - quadruplex DNA</td>
<td>0.8</td>
<td>0.044</td>
<td>1.3</td>
<td>0.031</td>
<td>4.6</td>
</tr>
<tr>
<td>poly(rA).[poly(rU)]_2 triplexRNA</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Micrococcus Lysodeicticus-dsDNA</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.6</td>
</tr>
<tr>
<td>5'-TTGGG [TTAGGG]_3A - 3' quadruplexDNA</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.6</td>
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<tr>
<td>poly(dGdC).poly(dGdC)dsDNA</td>
<td>-</td>
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<td>-</td>
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<tr>
<td>poly(dA).poly(dT) dsDNA</td>
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</tr>
</tbody>
</table>