THE IMPACT OF IONS ON RNA APTAMER-AMINOGLYCOSIDE INTERACTIONS

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Signature:
ABSTRACT

THE IMPACT OF IONS ON RNA APTAMER-AMINOGLYCOSIDE INTERACTIONS

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Aptamers are small nucleic acid or peptide molecules that can bind to specific targets with high affinity. They have been applied for diagnostic, therapeutic and imaging purposes. In order to have the best performance of aptamers, it is worth understanding their structural dynamics. Previous studies have demonstrated that aptamer structures and thus their performances are affected by the surrounding sodium, potassium and magnesium ion concentrations. Based on these findings, we hypothesized that these ions may differently affect the binding affinity of nucleic acid aptamers. In our study, we investigated the effects of varying concentrations of sodium, potassium and magnesium ions on RNA aptamers that bind aminoglycosides. Using Isothermal Titration Calorimetry (ITC) and UV spectrophotometry, four neomycin-B binding RNA aptamers were tested in buffer solutions with different ionic contents. Our findings showed that the presence of magnesium leads to a significant change in both binding mode and affinity; whereas, potassium and sodium only affect the binding affinity of the aptamers. Depending on these results, we performed fluorescence assays with fluorophore-modified aptamers and the same buffers as used before. The change in the fluorescence signal between bound and unbound aptamers verified specific ligand binding and indicated that the concentrations and composition of these
ions alter the three-dimensional structure and consequently affect the binding characteristics of aptamers. The main reason for the observed variation in aptamer performance seems to be the result of aptamers adapting to a predominant structure under a specific buffer condition. Therefore, components of the buffer systems that the aptamers are selected and optimized need to be carefully considered by researchers, because in the light of our results, the binding may get affected once the buffer system is modified by end-users.

Keywords: RNA Aptamers, Metal Ions, Aminoglycosides, Isothermal Titration Calorimetry, UV Spectroscopy
RNA YAPILI APTAMERLERİN AMİNOGLİKOZİTLERLE ETKİLEŞİMİNDE METAL İYONLARININ ETKİSİ

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Aralık 2019, 59 sayfa

değişiklik, bağlanmanın hedefe spesifik gerçekleştiğini doğrulamış ve bu iyonların
konsantrasyon ve kompozisyonlarının aptamerlerin üç-boyutlu yapılarını ve buna
bağlı olarak bağlanma karakteristiklerini değiştirdiğini göstermiştir. Aptamer
performanslarında görülen bu farklılığın temel sebebi, aptamerlerin belirli bir tampon
cözelti içinde baskı bir yapıya sahip olmalarından kaynaklanır. Bu nedenle,
aptamerlerin seçildiği ve optimize edildiği tampon çözeltisi sisteminin içeriği
araştırmacılar tarafından dikkate alınmalıdır, çünkü bu sonuçların ışığında, son
kullanıcının tampon çözeltide yapacağı herhangi bir değişiklik bir bağlanmayı
etkileyebilir.

Anahtar Kelimeler: RNA Aptamerleri, Metal İyonları, Aminoglikozitler, İzotermal
Titrasyon Mikrokalorimetresi, UV Spektroskopisi
To my amazing mother…
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LIST OF ABBREVIATIONS

ABBREVIATIONS

DNA  deoxyribonucleic acid
RNA  ribonucleic acid
SELEX  Systematic Evolution of Ligands by Exponential Enrichment
PCR  Polymerase Chain Reaction
RT-PCR  Reverse Transcription Polymerase Chain Reaction
SPR  Surface Plasmon Resonance
ITC  Isothermal Titration Calorimetry
AFM  Atomic Force Microscope
TPP  Thiamine pyrophosphate
NMR  Nuclear Magnetic Resonance
nM  nanomolar
µM  micromolar
mM  millimolar
µL  microliter
mL  milliliter
K\textsubscript{d}  dissociation constant
ddH\textsubscript{2}O  distilled water
CHAPTER 1

INTRODUCTION

1.1. Nucleic Acids

In the year 1868, a Swiss scientist named Friedrich Miescher was studying the chemical composition of leukocytes and his studies eventually led to the discovery of DNA. He discovered a novel molecule which is resistant to proteases, contains large amounts of phosphorus and lacks sulphur. He figured that this molecule is not a protein or lipid and named it “nuclein” (Dahm, 2008). After this landmark discovery, in the early twentieth century, subsequent studies rapidly expanded the knowledge about this molecule, which is now called deoxyribonucleic acid, DNA. In 1944, Oswald Avery, Colin MacLeod and Maclyn McCarty claimed that DNA was the active carrier of genetic information and not proteins (Avery, Macleod, & McCarty, 1944). This theory was later confirmed in 1952 by Alfred Hershey and Martha Chase as a result of their discoveries concerning the mechanism of replication and the genetics of viruses. They observed that bacteria are infected with T2 bacteriophage viral genome but not proteins, and later on, the new viruses produced from the infected bacteria carried only viral DNA (Hershey & Chase, 1952). A year after this discovery, by extending the work of Rosalind Franklin and Maurice Wilkins, the structure of the DNA was confirmed to be a right-handed double helix using X-Ray crystallography (Watson & Crick, 1953). Thanks to all these studies and many more, today we know that DNA is the storage of all the genetic information, and the expression of the genomic material is needed to maintain the biological functions of all living organisms. The flow of genetic information from DNA into RNA and the synthesis of proteins are collectively termed the “central dogma”. Francis Crick’s version of the central dogma explains the flow of information from the DNA is transcribed into RNA, and then it’s translated into proteins (Dahm, 2010).
In the second half of the twentieth century, extensive studies on DNA were carried on, and the understanding of how the genetic information is processed in living organisms became more clear. In 1990, the prominent Human Genome Project was initiated. The project aimed to decipher the human genome by drawing a map of the sequences of all the genes in the human genome. Researchers successfully sequenced all 3.2 billion base pairs in less time than initially expected (Chial, 2008). The project was, in fact, a huge success in identifying the genes encoded in the human genome, but it also revealed that the human genome is much more complex, and still a lot of mysteries to be solved. Upon completion of the Human Genome Project in 2003, several follow-up studies were conducted. One of these studies was the Encyclopedia of DNA Elements (ENCODE) project, that aimed to identify all the functional components of the genome (Rosenbloom et al., 2012). The motivation was to determine the role of the remaining components of the genome, once improperly referred to as “junk DNA”.

Recent studies have shown that non-coding DNAs have roles in the regulation of transcription and translation; some of them are even transcribed into functional non-coding RNAs such as tRNA, rRNA, regulatory RNAs and long non-coding RNAs (lncRNAs) (Goff & Rinn, 2015). LncRNA used to be considered “transcriptional noise” because they were believed not to code for any proteins (J. Zhang et al., 2019). Surprisingly, extensive studies with lncRNAs identified that they have many vital roles in cellular functions including regulation of gene expression by sequestering miRNAs and transcriptional factors (Azlan, Obeidat, Yunus, & Azzam, 2019) and thus they were associated with embryonic development, some types of cancers, Hepatitis C and other diseases (Nam & Bartel, 2012).

RNA used to be recognized only as a passive carrier of genetic information through central dogma. However, the number of biological functions attributed to folded RNA molecules is still increasing with new studies. RNA has crucial roles in various pathways of cellular metabolism, such as protein synthesis and transportation, alternative RNA splicing, and regulation of translation. Many RNA types have been described over the past years, some called ribozymes having catalytic activities in
biochemical reactions just like enzymes. Some RNAs have important roles in the regulation of gene expression. Other types of functional RNAs include; small nuclear RNAs (snRNAs), needed for intron splicing in the nucleus; the signal recognition particle (SRP), helping protein transportation through membranes; microRNAs (miRNA) small regulatory RNAs involved in gene expression and the ribozymes catalyzing reactions on both RNA and DNA (such as group I introns) (Clancy, 2008). Another important role of RNA in gene regulation is mediated by riboswitches in both prokaryotes and eukaryotes. Riboswitches are RNA sensors that directly affect gene expression upon metabolite binding due to their ability to switch between conformations (Edwards & Batey, 2010). In all of these cases, metal ions play an essential role in providing structural stability, enabling catalytic functions and regulation. By the multifunctional nature of RNA biopolymers, RNA can serve as conformational switches, catalysts for chemical reactions and rigid structural scaffolds (Stampfl, Lempradl, Koehler, & Schroeder, 2007).

1.1.1. Chemical Structure of Nucleic Acids

‘Nucleic acid’ term refers to polynucleotide chains consisting of nucleotide bases as their monomeric unit. A nucleotide is formed of a heterocyclic nucleobase, either a purine base (A, adenine or G, guanine) or pyrimidine base (C, cytosine; U, uracil or T, thymine), covalently bound with a glycosidic bond to a pentose sugar (a nucleoside), which is attached to a phosphate with a phosphodiester bond at the 5’ or 3’ end. Nucleotides (NTP, where N may be A: adenosine; G: guanosine; C: cytidine; T: thymidine; or U: uridine) are linked together through phosphate groups by the 5’ and 3’ hydroxyl groups to form phosphodiester bonds and make up oligonucleotide chains, or attached to additional phosphate groups to form either NDPs with diphosphates, or NTPs with triphosphates. Structures of nucleotides are illustrated in Figure 1.1.
The two classes of nucleic acids; ribonucleic acids (RNA) carry a hydroxyl group (-OH) at the 2’ position on the pentose sugar; whereas, deoxyribonucleic acids (DNA) have a hydrogen atom (-H) instead. The heterocyclic bases A, G, C, and T are commonly seen in the DNA, but the T is replaced by U in RNA. These heterocyclic bases are typically planar, and their exocyclic groups lie in the center of the rings, and lonepair electrons are delocalized into the π-systems of the rings. The tautomeric forms of the heterocyclic bases are found in trace concentrations in solution.
Compared to DNA, the 2’-OH group of RNA provides more structural complexity and more options for metal ion binding. Although this 2’-OH group gives RNA a better structural stability under acidic pH conditions, it also makes RNA susceptible to depolymerization due to transesterification caused by polyvalent metal ions at neutral pH.

Water plays an essential role in nucleic acid structure as nucleic acids usually remain highly hydrated in solution. Two H₂O molecules per phosphodiester bond and up to six H₂O per base pair are found in both DNA and RNA. Water interaction with nucleic acids can be classified in two different ways. Firstly, they form hydrogen bonds and create a highly ordered hydration shell around the nucleic acid. This is crucial for intra- and intermolecular interactions between nucleic acid residues and in metal ion-nucleic acid binding, because these sites need to be partially dehydrated to form direct bonds. The second means of water interaction is through a partially ordered water layer weakly bound to the nucleic acid, usually through secondary H-bonds or induced dipole-dipole interactions. These water molecules can be easily replaced with metal ions on the nucleic acid surface. The polarization of water molecules around the nucleic acids can weaken the electrostatic interactions with nucleic acids (Kazakov & Hecht, 2016).

The chemical composition of nucleic acids provides a wide variety of metal-binding sites to form ionic, coordinative (donor-acceptor), hydrogen (-H), and covalent bonds. Specific and nonspecific interactions with metal ions are known to be essential for both RNA folding and stabilization (Lipfert, Doniach, Das, & Herschlag, 2014). To perform all the biological functions, RNA molecules are folded into unique tertiary structures that are highly complex, compact and stable. Even small RNAs are relatively structured chains, which target terminal regions of mRNA for regulating gene expression. A number of RNA structural motifs require certain ions to stabilize the RNA chain. Many ribozymes and riboswitches depend on metal ions as cofactors to form functional configurations (Serganov & Patel, 2012).
1.1.2. RNA Folding

RNA molecules gain their tertiary structures through a two-step process. The first step of RNA folding is the formation of RNA secondary structure, which contains double-stranded regions through complementary base pairing and structural motifs including stem-loops, pseudoknots, internal bulges, hairpins, kissing loops, G-quadruplexes, three-way junction etc. Because of the fact that nucleic acids are highly negatively charged, the RNA molecules depend on cations and molecules to shield the negative charges, so to reduce the repulsion of polyanionic chains. In turn, this shielding promotes folding and helps the RNA adopt a stable secondary structure. Therefore, this process is mediated by monovalent and divalent cations, cationic polyamines, and some basic proteins. Figure 1.2 shows a simplified version of RNA folding pathway and the predominant metal ions involved at each step.

*Figure 1.2 Metal Ion Driven RNA Folding*
The second stage of RNA folding is the formation of a unique tertiary structure built to perform a specific biological activity. It is formed by intramolecular interactions between bases (base stacking), between sugars (sugar packing) and phosphate backbone. The formation of tertiary structure of RNA is particularly dependent on its sequence, type of metal ions (divalent cations Mg\(^{2+}\) and Ca\(^{2+}\); monovalent cations Na\(^+\) and K\(^+\)) and RNA’s binding partners (RNA binding proteins) (Pyle, 2002).

1.2. Nucleic Acid-Metal Ion Interactions

Metal ions are proven to be essential to maintain structural stability and ensure functionality of the nucleic acids in general (Pyle, 2002). Because nucleic acids are often in contact with metal ions, they are under the influence of the electrostatic interactions. As a result, the thermodynamic stability of their secondary and tertiary structures as well as the kinetic mechanism of RNA folding are affected. (Woodson, 2005).

Close to a hundred metal ions can be found in nature but most of them are not functionally required in biological organisms and some of them are even toxic. Cellular functions like catalysis depend on specific metal ions as cofactors for such reactions (P. Y. Watson & Fedor, 2011). Magnesium (Mg\(^{2+}\)) and potassium (K\(^+\)) are the most frequent ions found in intracellular fluids, whereas the others can be found only in trace amounts and unlikely to interact with nucleic acids. Sodium (Na\(^+\)) is the most abundant monovalent cation in extracellular fluids, so it’s rarely in contact with intracellular nucleic acids (Xi, Wang, Xiong, Zhang, & Tan, 2018). However, it is usually used in the buffer systems to perform binding assays, nuclear magnetic resonance (NMR) spectroscopy (Barnwal, Yang, & Varani, 2017) or X-Ray crystallographic studies (Westhof, 2015) with nucleic acids.

Na\(^+\) and K\(^+\) are the two monovalent cations used during structural analyses of nucleic acids. Numerous studies demonstrated that K\(^+\) is essential for RNA functions. For example, studies report that ribosomal activity is strongly dependent on the presence
of K⁺ or ammonium (NH⁴⁺) cations but is inhibited by Na⁺ (Rozov et al., 2019). Mammalian ribosomes go through a significant unfolding and lose their functions in the absence of K⁺ (Näslund & Hultin, 1971). However, several studies indicated that some RNA structural motifs specifically require Na⁺ ions for structural stabilization. The stability of hairpin motifs and some ribozymes have been shown to decrease when Na⁺ ions were substituted by K⁺ ions in the buffer system (Johnson-Buck, McDowell, & Walter, 2011).

Mechanistically, monovalent cations contribute to charge neutralization and help bring the polynucleotide chains together. However, they are usually required in a lot higher concentrations than divalent cations to stabilize the RNA structures. In the absence of divalent cations, they are unable to lead to the active folding of large RNA chains. For example, Na⁺ and K⁺ cations are needed for the activation of ribozymes, but they don’t seem to be directly involved in catalysis (Johnson-Buck et al., 2011). In terms of divalent cations, Mg²⁺ is intracellularly the most abundant and mainly involved in RNA structural stability and functionality. Mg²⁺ has a smaller ionic radius than K⁺ and Na⁺, and it is the preferred divalent cation for charge neutralization over monovalent ions because of its higher charge density and lower entropic cost for localization (Auffinger, Grover, & Westhof, 2011).

1.2.1. Metal Ion Interactions with RNA

Metal ion binding to RNA usually occurs in two different means; either through diffuse or site-specific binding (ion chelation). “Diffuse ions” are mostly hydrated in a solution and they are attracted to the electrostatic field of RNA. Diffuse binding provides charge shielding and reduces the electrostatic repulsion between RNA backbone. Such shielding can be implemented by a number of mono- or divalent cations that is necessary for RNA to acquire both secondary and tertiary structures (Pyle, 2002). The concentration of diffuse ions in proximity of an RNA molecule is determined by the magnitude of electrostatic potential there. (Draper, 2004). Divalent
Mg$^{2+}$ ions are also shown to diffuse around nucleic acids. So, both monovalent and divalent cations contribute to diffuse ion atmosphere and help stabilization RNA tertiary structure. Although few RNAs can fold into stable tertiary structures with only monovalent ions present, diffuse Mg$^{2+}$ ions are proven to have a significant impact on stabilizing nucleic acids by directly interacting with phosphate backbone (Pyle, 2002).

The water ligands of metal ions can interact with RNA bases and backbone substituents on the folded RNA’s inner or outer spheres. Outer sphere interactions usually don’t have a direct effect on RNA functions, but they are needed to maintain specific motifs. Interactions of metal ions with RNA inner sphere are usually needed for catalytic function (Batey, Rambo, Lucast, Rha, & Doudna, 2000).

The second mechanism of ion interaction is ion chelation. “Chelated ions” can interact with specific sites on the RNA structure directly and they are held in that location by electrostatic forces. K$^+$ and Mg$^{2+}$ are the cations that mostly interact with other atoms and molecules through electrostatic forces because they both have a closed shell electronic structure. Ion chelation can lead to the displacement of some of the water molecules around diffuse ions on the RNA surface and this has a great energetic cost due to repulsion of these diffuse ions. In order to form direct bonds with RNA, metal ions have to be partially dehydrated. Although, dehydration of Mg$^{2+}$ ions is energetically very expensive, it is required to bring neighboring phosphate groups in close proximity (Draper, 2004). For instance, Mg$^{2+}$ binding to internal loop motifs results in bringing the neighboring chains into an unusual proximity (Correll, Freeborn, Moore, & Steitz, 1997) or dehydrated K$^+$ ion binding to the A-platform motif. The latter interaction inhibits RNA function until the potassium is replaced by thallium (Tl$^+$) (Basu et al., 1998). On folded RNA molecules, metal ions usually bind to deep and narrow groove as demonstrated by NMR and X-Ray crystallography (Ferré-D’Amaré & Doudna, 2000). Studies on electrostatic surface potential on RNAs showed that the major groove has a high negative potential, thus, it attracts many ions (Chin, Sharp, Honig, & Pyle, 1999). One of the most common binding sites on the major groove is the tandem G-U wobble base pairs which have a
high negative charge, so it is a preferred binding site for metal ions and coordinated water molecules (Cate & Doudna, 1996).

1.3. Aptamers

Aptamers are single-stranded deoxyribonucleic acid (DNA) or ribonucleic acid (RNA) oligomers or small peptides that bind to their specific targets with high affinity and selectivity. The term “aptamer” is coined from the Latin word *aptus*, which means fit, and the Greek *meros*, meaning part or region. Their potential ligands include proteins and peptides, macromolecules like carbohydrates and lipids, complex molecular suprastructures like viruses and other microorganisms or even whole cells (Dunn, Jimenez, & Chaput, 2017). Aptamers bind to their ligands in a structure-specific manner, by folding into their unique and complex three-dimensional conformations naturally. They are obtained by a selection method called Systematic Evolution of Ligands by Exponential Enrichment (SELEX) (Y. Zhang, Lai, & Juhas, 2019). After selection, oligomers fold and obtain tertiary structural motifs such as stems, loops, bulges, hairpins, pseudoknots, triplexes, or quadruplexes. Although aptamers can be functional in their native 3D forms, they are often further modified with various functional groups to enhance stability, for immobilization and cellular intake. As a result of their high selectivity and affinity to a certain ligand, they have been the preferred tool for many biotechnological applications like diagnostic, therapeutic and imaging purposes. They have also been utilized in drug targeting and novel immunotherapy approaches to treat various diseases (Wengerter et al., 2014).

Due to their similar applications, aptamers and antibodies are often compared. Table shows the main comparison points between aptamers and antibodies. The prominent advantage of aptamers over antibodies is their stability. As they are nucleic acid-based molecules, aptamers are more thermally stable than protein-based antibodies, which go under irreversible degradation at high temperatures; whereas, aptamers can be refolded into their tertiary structures and gain functionality, so they can be used
repeatedly. Nucleic acids are usually not recognized by the human immune system, so aptamers have very low immunogenicity and toxicity in the body. Another important advantage is the ease of aptamer production. While the production of monoclonal antibodies is a very laborious and expensive process which sometimes can take months, upon selection, aptamers can easily be obtained in high amounts with great accuracy using solid state synthesis, during which aptamers can be further modified with other functional groups. Antibody production requires an immune response, so they cannot recognize certain nonimmunogenic ligands such as ions and small molecules for which aptamers show a high affinity and selectivity (Song, Lee, & Ban, 2012).

Table 1.1 Comparison of Aptamers and Antibodies

<table>
<thead>
<tr>
<th>Feature</th>
<th>Aptamer</th>
<th>Antibody</th>
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<tbody>
<tr>
<td>Molecular Weight</td>
<td>Small (&lt;30 kDa)</td>
<td>Relatively Large (&gt;75 kDa)</td>
</tr>
<tr>
<td>Stability</td>
<td>Refolds to functional tertiary structure after denaturation</td>
<td>Irreversibly denatured at high temperatures</td>
</tr>
<tr>
<td>Affinity &amp; Specificity</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Targets</td>
<td>Any molecule or cell</td>
<td>Immunogenic molecules</td>
</tr>
<tr>
<td>Minimum Target Size</td>
<td>Small, ~60 Da</td>
<td>~600 Da</td>
</tr>
<tr>
<td>Immunogenicity</td>
<td>Very low</td>
<td>High</td>
</tr>
<tr>
<td>Production time</td>
<td>Several days to weeks</td>
<td>Several months</td>
</tr>
<tr>
<td>Reproducibility</td>
<td>Very low batch variation</td>
<td>Very high batch variation</td>
</tr>
<tr>
<td>Cost and Manufacture</td>
<td>Cheap, in vitro</td>
<td>Expensive, in vivo</td>
</tr>
<tr>
<td>Chemical Modifications</td>
<td>Easily modified with functional groups</td>
<td>Difficult to chemically modify</td>
</tr>
<tr>
<td>Storage</td>
<td>Stable at room temperature</td>
<td>Easily denatured, needs freezing</td>
</tr>
<tr>
<td>Shelf Life</td>
<td>Several years (if dried)</td>
<td>~6 months</td>
</tr>
</tbody>
</table>

1.3.1. Systematic Evolution of Ligands by Exponential Enrichment (SELEX)

Aptamers are selected against a specific target from a randomized oligonucleotide library by stimulating evolution in vitro under a defined set of conditions. This
selection method is called SELEX and it was first discovered in 1990 by three separate
groups; namely Tuerk and Gold (Tuerk & Gold, 1990), Ellington and Szostak
(Ellington & Szostak, 1990) and Robertson and Joyce (Robertson & Joyce, 1990).
SELEX includes a number of selection rounds (5 to 20 rounds); at each round, a
random oligonucleotide pool is incubated with the target molecule and the best
binding oligonucleotides are isolated for enrichment by PCR amplification (PCR for
dNA libraries, RT-PCR for RNA libraries). At the end of these selection rounds, the
sequences of the oligonucleotides with the highest affinity toward the target are
identified.

Before a SELEX procedure, a randomized oligonucleotide library is obtained. The
initial oligonucleotide libraries usually consist up to $10^{15}$ unique sequences, which all
carry a random region (20-50 nucleotides long) flanked by two fixed primer binding
sites for PCR amplification. A conventional SELEX experiment is usually carried in
four steps: 1. Incubation of the oligonucleotide pool with the target 2. Partitioning of
the binding sequences 3. Amplification of these sequences with PCR 4. Sequencing

![Figure 1.3 Schematic Illustration of a SELEX Procedure](image-url)
the best binding sequences to identify aptamers. Before sequencing the most promising oligonucleotides, first three steps of this procedure are followed for several rounds (5-20 rounds) to amplify the bound sequences. At each round unbound oligos are separated and bound oligos are amplified using PCR for DNA libraries or RT-PCR for RNA libraries. For DNA SELEX, biotin-labeled primers can be used in PCR, and the resulting double-stranded forms are separated using methods such as streptavidin bead capture. For RNA, T7 RNA polymerase promoter containing primers are used in RT-PCR after which RNA libraries are amplified by in vitro transcription. The target molecules are usually immobilized on a matrix to achieve partitioning of the bound oligos. To eliminate the sequences that bind to the matrix in a target-independent manner, a negative selection round is usually included, which is done by passing the oligonucleotide library through the supporting matrix. A similar approach, called counter-selection, utilizes analogs or structurally similar targets to remove the non-specific binding sequences from the mixture. These procedures are applied in most of the modified SELEX methods to ensure that the obtained sequences are selective and specific to the targets. After the final round, the selected oligonucleotides are sequenced to identify the aptamers. Generally, next-generation sequencing is used to obtain sequences of binding oligos from different rounds of selection. Comparative analysis of these sequences shows the recurring motifs involved in recognition of the target. The most promising sequences are amplified and further characterized.

A number of modified SELEX techniques that aim to shorten the time needed for high-affinity aptamer selection or to obtain aptamers specifically designed for certain applications have been developed. For example, Golden et al. suggested photochemical SELEX (photoSELEX) which enables selection of modified aptamers and required fewer number of selection rounds than traditional SELEX methods. The selected aptamers can form photoinduced covalent bonds with their ligands, to make them more specific (Tombelli, Minunni, & Mascini, 2005). A recently developed method that aims to cut down the amount of time and labor required for SELEX is robotic assisted selection procedure. It is a fully automated method that can perform
12 consecutive selection rounds in less than two days and select aptamers against 8 different targets simultaneously, without any manual interruption needed (Breuers, Bryant, Legen, & Mayer, 2019).

### 1.3.2. **Aptamer Modifications**

Despite having the advantages of low immunogenicity, easy production, low cost and high thermal stability, aptamers may undergo degradation in cells, and get rapidly extracted from the body through renal filtration. Nucleic acid aptamers are recognized by nucleases, and peptide aptamers are susceptible to protease degradation (Ni et al., 2017). To circumvent these problems, numerous chemical modifications can be implemented for better aptamer stability *in vivo*. Among these chemical modifications, terminal end modifications such as 5’ PEGylation and 3’ capping with inverted thymidine, and the modifications on the nucleobases, sugar rings or the phosphodiester bonds are the most commonly used ones (Zhuo et al., 2017). Attachment of certain functional groups is also needed for immobilization of aptamers onto solid supports for biosensor development, such as amine (-NH₂), thiol (-SH) or carboxyl (-COOH) groups (E. Wang, Wu, Niu, & Cai, 2011).

Although most of the modifications on aptamer structures are usually applied after selection to enhance stability, this can cause a decrease in affinity towards the target because it may prevent the aptamer ligand interaction or alter the structure of the aptamer. Alternative to the traditional selection methods, modified SELEX techniques that utilizes pre-modified nucleotides have been developed in order to obtain inherently stable aptamers with high affinity. Mod-SELEX methods use chemically modified oligonucleotide libraries such as 2’-amino pyrimidines, 2’fluoro pyrimidines or 2’-O-methyl nucleotides (Keefe & Cload, 2008).

To date, hundreds of aptamers against numerous targets have been selected with different SELEX methodologies, and further modified for various applications. Selected aptamers are often used as recognition molecules for biosensor platforms,
which are called “aptasensors”. Aptamers offer many advantages over antibodies, such as their small size, affinity and selectivity, cost effectiveness and most importantly their flexibility and compatibility for novel biosensor designs. Incorporation of aptamers with a diverse range of nanostructures and sensing methodologies has led to the development innovative biosensors, which are highly sensitive and selective; including electrochemical, optical, microcantilever, and acoustic detectors (Mehlhorn, Rahimi, & Joseph, 2018).

Analyzing the binding properties of aptamers is important before using them for research and development purposes. Binding characteristics of aptamers include affinity, kinetics, specificity and buffer sensitivity, which are crucial information for aptamer design and use. In biosensor development, one of the most important aspects of sensing molecules is the affinity towards the target. The high binding affinity of the aptamers is the result of hydrogen bonds, structural compatibility, aromatic ring stacking, hydrophobic and electrostatic interactions and van der Waals forces (Zhuo et al., 2017). Measurement of the binding affinities of aptamers can be done using several biophysical methods; such as Surface Plasmon Resonance (SPR), Isothermal Titration Calorimetry (ITC), Flow Cytometry, Microscale Thermophoresis (MST), Atomic Force Microscope (AFM) etc. ITC is one of the most widely used methods to determine binding affinity of molecular interactions in solution. It is a label-free, quantitative technique to measure the heat release upon aptamer-ligand binding and determine the binding affinity (dissociation constant, $K_d$), enthalpy changes, stoichiometry and thermodynamic parameters of molecular interactions (Zhuo et al., 2017).

1.3.3. Structure-Function Relation

In order to form stable tertiary structures, RNA requires positive charges to shield the high negative charges on its phosphate backbone and reduce the repulsion between closely packed chains. Thus, metal cations have an essential role in facilitating RNA
folding in order to become biologically functional molecules. An extensive literature is available regarding how the metal ions interact and effect nucleic acid structures and their function. Aptamers are nucleic acids in nature, so they are also affected by the metal ions. Numerous studies show that aptamers may need certain cations to function or get inhibited by the ionic environment. Characterization of aptamers is needed to understand their molecular recognition, because it is important to know their key properties, such as affinity, kinetics, specificity, ion dependence and buffer sensitivity, in order to efficiently use them in diverse applications (Chang, McKeague, Liang, & Smolke, 2014).

Aptamers are nucleic acid molecules with negatively charged structures, so they are highly affected by the positive ions (cations) surrounding them. Their free form in a solution is highly dependent on the electrostatic interactions with specific ions, ionic strength and pH because these factors affect the charge distribution of the aptamers and consequently their ligand interactions. In different ionic conditions, aptamers will follow different folding patterns and thus their binding affinity and selectivity towards their target will be altered. For this reason, the properties of aptamers and their performances in future applications depend on the buffer system used during the SELEX procedure (Ilgu, Fazlioglu, Ozturk, Ozurekci, & Nilsen-Hamilton, 2019).

Aptamers bind their ligands in a structure-specific manner through their unique tertiary structures, which they naturally fold into with their surrounding ions. Their binding sites are mainly determined by the secondary structural motifs such as stem-loops, hairpin, pseudoknots, internal bulges, kissing loops, three-way junctions and G-quadruplexes. Based on the primary sequences, these motifs can be predicted (Parisien & Major, 2008) and help to determine the binding site on the aptamer’s tertiary structure, which accounts for only a short section of the aptamer. To enhance aptamer selectivity and affinity, truncation is an effective way to minimize the sequence and increase the specific contact points and make the aptamer more accessible for ligand binding (Hasegawa, Savory, Abe, & Ikebukuro, 2016).
1.3.4. Riboswitches and Ion Dependency

Riboswitches are non-coding RNA sequences that carry an aptamer domain as sensing element and a downstream expression domain that regulates the transcription and translation. Regulation by riboswitches is possible through conformational changes thanks to their flexible structures upon specific metabolite binding to the aptamer domain. Riboswitches can regulate gene expression through transcription termination, translation inhibition or inducing alternative splicing of mRNA (McCluskey et al., 2019).

A study with thiamine pyrophosphate (TPP) sensing riboswitches showed that interaction with Mg$^{2+}$ is essential for TPP binding and it cannot be substituted by other monovalent or divalent metal ions. The X-ray crystallography studies of the thiC TPP riboswitch from Arabidopsis thaliana revealed that oxygen atoms are chelated in the pyrophosphate moiety of TPP by only one Mg$^{2+}$ ion when TPP is bound to the RNA. On the other hand, when the thiM riboswitch from Escherichia coli is bound to TPP, two bridging Mg$^{2+}$ ions chelate oxygen atoms in the pyrophosphate region. In case of this system, the presence of Mg$^{2+}$ ions is strictly needed for binding of TPP to its riboswitches. However, some riboswitches do not depend on Mg$^{2+}$ ions specifically to function. For example, it was shown that the GlmS riboswitch, which is a ribozyme that undergoes self-cleavage when it interacts with glucosamine-6-phosphate (GlcN6P), does not rely on Mg$^{2+}$ ions for this reaction. Here, the system only requires the positive charges for the electrostatic stabilization of the structure, so high concentrations of monovalent cations, other divalent cations or Co(NH$_3$)$_6^{3+}$ can substitute Mg$^{2+}$ ions. (Noeske, Schwalbe, & Wöhnert, 2007).

In another work, which investigates the effect of Mg$^{2+}$ ions on conformational dynamics and affinity of B. subtilis lysC lysine-binding riboswitch, it was shown that sub-millimolar concentrations of Mg$^{2+}$ have a huge impact on aptamer function and affinity by altering the conformation. This study confirmed that cellular
concentrations of Mg\(^{2+}\) cations are high enough to affect RNA functions, stabilize their structures, and also play a role in modulation of gene expression (McCluskey et al., 2019).

1.4. Aminoglycosides as Aptamer Targets

Aminoglycosides are a large family of potent antibiotics that target bacterial ribosome and disturb its functionality. They can bind to the ribosomal decoding site (A-site) or to several ribozymes and inhibit their activity. Aminoglycosides have hydrophilic structures, so they need an electron transport system from the host’s respiratory system to penetrate the hydrophobic cell membranes of the bacteria. As a result, they are only effective against aerobic bacteria (Germovsek, Barker, & Sharland, 2017; Krause, Serio, Kane, & Connolly, 2016). In the cytosol, they bind to the 16S rRNA decoding region (A-site) on the 30S small subunit of the bacterial ribosome. Upon aminoglycoside binding, the position of the two adenines, A1492 and A1493, which are needed for the codon–anticodon interaction, get shifted; so, the protein synthesis is inhibited (Fourmy, Recht, Blanchard, & Puglisi, 1996; Ramakrishnan, 2002).

Aminoglycosides are highly flexible molecules with multiple positive charges, which facilitate their binding to the negatively charged major grooves of a variety of RNA molecules (Vakulenko & Mobashery, 2003). It was shown that aminoglycoside binding leads to displacement of divalent metal ions required for both RNA and ribozyme functionality. A study revealed structural data for a small RNA representing 16S rRNA decoding region in complex with several aminoglycosides using NMR spectroscopy. They showed that divalent Pb\(^{2+}\) ions induce cleavage on various sites on rRNA A-site which are needed for functionality, and upon binding with aminoglycoside neomycin B, Pb\(^{2+}\) induced cleavages are inhibited. This data, along with other similar studies, suggest that binding of aminoglycosides to rRNA A-site leads to divalent ion displacement, hence causing dysfunctionality (codon misreading).
Aminoglycosides contain a highly conserved aminocyclitol ring to which amino sugars are attached by glycosidic linkages. The aminocyclitol ring consists of a 2-deoxystreptamine (2-DOS) and has 1,3-diamino functionality and three or four hydroxyl groups to bind amino sugars. Aminoglycosides are generally divided into three structural subclasses depending on the position of these glycosidic linkages: 4-monosubstituted, 4,5- or 4,6-disubstituted (Kulik, Goral, Jasiński, Dominiak, & Trylska, 2015). Despite the differences in their substitutions, aminoglycosides in each subclass have high structural resemblance. Figure 1.4 illustrates the chemical structures of aminoglycosides.

![Figure 1.4 Structures of Aminoglycosides](image_url)
4-monosubstituted aminoglycosides include neamine, paromamine and apramycin, among which apramycin is the only one used pharmaceutically and it carries a bicyclic ring 1. Paromamine and neamine only differ in their R substituent groups and they are not used as drugs alone.

4,5-disubstituted aminoglycosides include ribostamycin, butirosin B, neomycin B, paromomycin and lividomycin A. The antibiotics in this group differ in their substitutions on the third ring. Lividomycin A is the only aminoglycoside in this group that contains five rings.

Kanamycin A, kanamycin B, kanamycin C, tobramycin, sisomicin, netilmicin and amikacin are in a subcategory of 4,6-disubstituted aminoglycosides, and the other subcategory includes gentamicin C1, gentamicin C2, gentamicin C1A and geneticin. The major reason of dividing this group into such subcategories is the presence of an extra R group attached to the 6’ position of the first ring as well as the third ring connected to the 2-DOS (Chittapragada, Roberts, & Ham, 2009).

1.4.1. Aminoglycoside Detection with Aptamers

A number of high affinity aptamers have been selected against aminoglycosides such as neomycin B (Wallis, von Ahsen, Schroeder, & Famulok, 1995), tobramycin (Wang & Rando, 1995), lividomycin, kanamycin A (Lato, Boles, & Ellington, 1995) and kanamycin B (Kwon, Chun, Jeong, & Yu, 2001). These selections were done using libraries that have 30 to 74 nucleotide long random regions (Klug & Famulok, 1994). Selections against aminoglycosides resulted in isolation of aptamers that could promiscuously bind to other types of aminoglycosides more or less specifically, due to not applying counterselection procedures. For example, Kwon et al. selected the K8 aptamer originally to bind kanamycin B, but it was shown to have 10-times higher affinity against tobramycin (Kwon et al., 2001). Another aptamer (W13 RNA) was isolated to bind tobramycin, which could recognize neomycin too. Similarly, it is known that most of the lividomycin aptamers can recognize neomycin B and
paromomycin as well (Lato et al., 1995). Based on the outcome of these studies, it is clear that most aminoglycoside-binding aptamers are structurally similar, although they might have different primary sequence and binding patterns. In general, the aminoglycoside-binding RNA aptamers have stem-loop structures and the stems often carry internal bulges or small hairpin loops, resulting in a widened major groove. The loops, essential for binding are quite variable in sequence but seem to show a recurring motif (Lorenz & Schroeder, 2006).

Structural studies of tobramycin and neomycin aptamers revealed similarities in binding patterns; the loops are closed with non-canonical base pairing and the antibiotic is trapped in the major groove (L. Jiang et al., 1999; Licong Jiang, Suri, Fiala, & Patel, 1997). Also, neomycin riboswitches selected in yeast shared this structural feature (Weigand et al., 2011).

Since their discovery in the 1940s, aminoglycosides have been the most commonly used antibiotics around the world due to their high efficiency as bactericidal drugs and their low cost. Antibiotics are often misused for humans or in veterinary medicine and in the food industry, for example as growth enhancers in livestock. The unnecessary intake of antibiotics leads to the evolution of antibiotic-resistant bacteria which has recently become a serious problem. This results in current antibiotics losing their effectiveness in treating microbial diseases (Furusawa, Horinouchi, & Maeda, 2018). According to the reports, infections with antibiotic-resistant bacteria account for approximately 33000 deaths each year (Cassini et al., 2018). There have been several successful initiatives to prevent misuse of antibiotics.

In parallel, researchers have successfully developed aptasensors to detect antibiotics in body fluids, remnants in dairy products and food samples in several laboratories today. Selected examples of developed aptasensors for aminoglycoside detection are summarized in Table 1.2.
### Table 1.2 Aptsensors for Aminoglycoside Detection

<table>
<thead>
<tr>
<th>Target</th>
<th>Nucleotide Backbone</th>
<th>Sequence</th>
<th>Dissociation Constant (Kd, nM)</th>
<th>Aptsensor Type/Detection Method</th>
<th>Limit of Detection (LOD, nM)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Neomycin B</strong></td>
<td>RNA</td>
<td>GGA CUG GGC GAG AAG</td>
<td>115 ± 25</td>
<td>Fluorometric</td>
<td>10</td>
<td>(Ling et al., 2016; Wallis et al., 1995)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>UUC AGU CC</td>
<td></td>
<td></td>
<td></td>
<td>(De-los-Santos-Álvarez et al., 2007)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GCC CUG GGC GAG AAG UUU AGG CC</td>
<td>2500 ± 900</td>
<td>Impedimetric Electrochemical</td>
<td>5</td>
<td>(De-los-Santos-Álvarez et al., 2009)</td>
</tr>
<tr>
<td></td>
<td>DNA</td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>78.8</td>
<td>Cantilever Aptsensor</td>
<td>5000</td>
<td>(Bai et al., 2014)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>78.8</td>
<td>Colorimetric/UV-Vis</td>
<td>25</td>
<td>(Song et al., 2011)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>78.8</td>
<td>Colorimetric/UV-Vis</td>
<td>2.5</td>
<td>(Niu et al., 2014)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>8.38</td>
<td>Colorimetric Aptsensor</td>
<td>1.49</td>
<td>(Kumar Sharma et al., 2014)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>78.8</td>
<td>Colorimetric/UV-Vis</td>
<td>0.014</td>
<td>(Zhou et al., 2014)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>78.8</td>
<td>Colorimetric Aptsensor</td>
<td>4.5</td>
<td>(Y. Xu et al., 2015)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CGG AAG GCC GCC ACC CCA TCG GCG GGG GCG AAG CTT GCG</td>
<td>92.3 ± 29.1</td>
<td>Colorimetric Aptsensor</td>
<td>3.35</td>
<td>(Ha et al., 2017; Ha et al., 2017)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>0.0778</td>
<td>Colorimetric Aptsensor</td>
<td>0.0778</td>
<td>(J. Liu et al., 2018)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>Liquid Crystal Assay</td>
<td>&lt;1</td>
<td>(Ying Wang et al., 2017)</td>
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<tr>
<td></td>
<td></td>
<td>AAG TGG GGG TTT AGG CTA AGC CGA</td>
<td>0.612</td>
<td>Fluorometric</td>
<td>0.612</td>
<td>(Khabeaux et al., 2015)</td>
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<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>1.58</td>
<td>Fluorometric</td>
<td>1.58</td>
<td>(C. Liu et al., 2015)</td>
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<tr>
<td></td>
<td></td>
<td>TGG GGG TTT AGG CTA AGC CGA</td>
<td>Electrochemiluminescence</td>
<td>0.35</td>
<td>(Feng et al., 2019)</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>GGG ACT TGG TTT AGG TAA TGA GTC CC</td>
<td>650,000</td>
<td>Amperometric Electrochemical/Square Wave Voltammetry</td>
<td>(Rowe et al., 2011; Wang and Rando, 1995)</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>GGG ACT TGG TTT AGG TAA TGA GTC CC</td>
<td>281,000</td>
<td>Amperometric Electrochemical/Square Wave Voltammetry</td>
<td></td>
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<td></td>
<td>GGG ACT TGG TTT AGG TAA TGA GTC CC</td>
<td>450,000</td>
<td>Amperometric Electrochemical/Square Wave Voltammetry</td>
<td></td>
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<tr>
<td><strong>Kanamycin</strong></td>
<td>RNA</td>
<td>GGG ACT TGG TTT AGG TAA TGA GTC CC</td>
<td>72,000</td>
<td>Amperometric Electrochemical/Square Wave Voltammetry</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>GGG ACT TGG TTT AGG TAA TGA GTC CC</td>
<td>80,000</td>
<td>Amperometric Electrochemical/Square Wave Voltammetry</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>DNA</td>
<td>DNA</td>
<td></td>
<td></td>
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</tr>
<tr>
<td><strong>Gentamicin</strong></td>
<td>RNA</td>
<td>GGG ACT TGG TTT AGG TAA TGA GTC CC</td>
<td>800,000</td>
<td>Amperometric Electrochemical/Square Wave Voltammetry</td>
<td></td>
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</table>
Table 1.3 Aptsensors for Aminoglycoside Detection, Continued

<table>
<thead>
<tr>
<th>Target</th>
<th>Nucleotide Backbone</th>
<th>Sequence</th>
<th>Dissociation Constant (Kd, nM)</th>
<th>Aptsensor Type/Detection Method</th>
<th>Limit of Detection (LOD, nM)</th>
<th>Reference</th>
</tr>
</thead>
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<td>Tetracyclins</td>
<td>DNA</td>
<td>TCC GTG TAG AGG TCG GGT CTC TGG CCA ACT GAT TCG TAG AAA AGT ATA GGC CCG CAG GGC</td>
<td>260</td>
<td>Surface Plasmon Resonance</td>
<td>500</td>
<td>(Cappi et al., 2015; Spiga et al., 2015)</td>
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<tr>
<td></td>
<td></td>
<td>TAG GGA ATT CGT CCG CCG AAT CAT GGC AGG TTA GTG GCG CGA TCG ATG ATA GGC CCG CAG GCA</td>
<td>56.9</td>
<td>Colorimetric Aptsensor</td>
<td>37.9</td>
<td>(Han et al., 2018)</td>
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<tr>
<td></td>
<td></td>
<td>CCG CCG CCG ACG TTA TGG GTC GAT AGA GCC CAC GTC GAC GCA</td>
<td>46.8</td>
<td>Colorimetric Aptsensor</td>
<td>37.10</td>
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<td>GCC ACT TGG TTG AGG TAA TGA GTC GAC</td>
<td>48.4</td>
<td>Colorimetric Aptsensor</td>
<td>37.11</td>
<td></td>
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<tr>
<td></td>
<td>RNA</td>
<td>GGC AGG AGG UUG AGC UAC ACU CGU GCC</td>
<td>400</td>
<td>Impedimetric Electrochemical/ Faradic Impedance Spectroscopy</td>
<td>400</td>
<td>(González-Fernández et al., 2011; Wang and Rand, 1995)</td>
</tr>
<tr>
<td>Streptomycins</td>
<td>DNA</td>
<td>GGG GTT TGG TGT TCT GGC TGG TGT TCT CTA GCA AGA GCC AGG TTT GTC CCA ACG</td>
<td>199.1</td>
<td>Colorimetric/UV-Vis</td>
<td>200</td>
<td>(Zhou et al., 2013)</td>
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<td>TAG AGG GTG TGG TCT AGA GTA TGG GTC TAC GTA AGG TAC GGA TCT TGG GTC GTC</td>
<td>221.3</td>
<td>Colorimetric/UV-Vis</td>
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<td>AGC TGG GTG GGG GCC ACC AGG AGG TAT ACG TCT TTT TGG</td>
<td>272.0</td>
<td>Colorimetric/UV-Vis</td>
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<td>TGG TGG TCT GGG GTG CTC GGC TGG TTA CGG GCC TGG GGG GAA</td>
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<td>Colorimetric/UV-Vis</td>
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<td>GGG GTT TGG TGT TCT GGC TGG TGT TCT CTA GCA AGA GCC AGG TTT GTC CCA ACG</td>
<td>199.1</td>
<td>Colorimetric Aptsensor</td>
<td>86</td>
<td>(Zhou et al., 2017; Zhou et al., 2013)</td>
</tr>
<tr>
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<td></td>
<td>TAG GGA ATT CGT CCG CCG AAT CGG GGT GCT CTC CTC CGT GGC TGG GCC GCC GCA</td>
<td></td>
<td>Photoelectrochemical Aptsensor</td>
<td>0.033</td>
<td>(X. Xu et al., 2017)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>TAG GGA ATT CGT CCG CCG AAT CGG GGT GCT CTC CTC CGT GGC TGG GCC GCC GCA</td>
<td></td>
<td>Amperometric Electrochemical/Differential Pulse Voltammetry</td>
<td>11.4</td>
<td>(Dansok et al., 2016)</td>
</tr>
</tbody>
</table>
CHAPTER 2

MATERIALS AND METHODS

2.1. Materials

2.1.1. Aptamers

All aptamers used in this project were purchased as lyophilized from Integrated DNA Technologies (IDT, Coralville, IA). They were then resuspended in sterile MiliQ water (RNase-free) to prepare 1mM stock solutions. Stocks were stored in -20 °C and kept on ice while sample preparations. All of the aptamers used in this research are single-stranded RNA aptamers. Their sequences are given in Table 2.1 and Table 2.2, and secondary structures with the lowest free energies are illustrated in Figure 2.1 and Figure 2.2 (RNAstructure Web Servers for RNA Secondary Structure Prediction, Matthews Lab, https://rna.urmc.rochester.edu/RNAstructureWeb/).

Table 2.1 Aptamers used in ITC Experiments

<table>
<thead>
<tr>
<th>Aptamer</th>
<th>Backbone</th>
<th>Size</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>31NEO3A</td>
<td>RNA</td>
<td>31nt</td>
<td>GGC AUA GCU UGU CCU UUA AUG GUC CUA UGU C</td>
</tr>
<tr>
<td>24m5NEO2A</td>
<td>RNA</td>
<td>24nt</td>
<td>CAC UGC AGU CCG AAA GGG CCA GUG</td>
</tr>
<tr>
<td>30NEO4A</td>
<td>RNA</td>
<td>30nt</td>
<td>CUU UGC GAU GUC CUU UAA UGG UCC GCG AGG</td>
</tr>
<tr>
<td>27m2NEO5A</td>
<td>RNA</td>
<td>27nt</td>
<td>GGC UGC UUG UCC UUU A/2AP/U GGU CCA GUC</td>
</tr>
</tbody>
</table>
Table 2.2 Aptamers used in fluorescence assays

<table>
<thead>
<tr>
<th>Aptamer</th>
<th>Backbone</th>
<th>Size</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>23m17NEO1A</td>
<td>RNA</td>
<td>23nt</td>
<td>GGA CUG GGC GAG /6-FAM/AG UUU AGU CC</td>
</tr>
<tr>
<td>25m10NEO2A</td>
<td>RNA</td>
<td>25nt</td>
<td>CAC UGC /6-FAM/GU CCG AAA AGG GCC AGU G</td>
</tr>
</tbody>
</table>

Figure 2.1 Secondary structures of aptamers used in ITC experiments

Figure 2.2 Secondary structures of 6-FAM modified aptamers
2.1.2. Antibiotic Targets

Neomycin-B (Sigma; Catalog number: N6910-25G, Lot number: 081K08832), Kanamycin-B (Sigma; Catalog number: B5264-100MG; Lot number: 099K1597) were purchased from Sigma Aldrich and Ampicillin (AppliChem, Catalog number: A6352,0025; Lot number: 6L008514) was purchased from AppliChem BioChemica. Stock solutions of 20 mM were prepared using sterile MiliQ water (RNase-free) and they were stored in -20 °C until they were used for experiments.

Table 2.3 Buffer systems used in experiments

<table>
<thead>
<tr>
<th>Buffers</th>
<th>Ionic Compositions</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>20 mM Na₂HPO₄, pH 7.4</td>
</tr>
<tr>
<td>B</td>
<td>20 mM Na₂HPO₄ + 5 mM MgCl₂, pH 7.4</td>
</tr>
<tr>
<td>C</td>
<td>20 mM Na₂HPO₄ + 125 mM KCl, pH 7.4</td>
</tr>
<tr>
<td>D</td>
<td>20 mM Na₂HPO₄ + 125 mM KCl + 5 mM MgCl₂, pH 7.4</td>
</tr>
</tbody>
</table>

2.2. Methods

2.2.1. ITC Experiments

Isothermal Titration Calorimetry (ITC) is a thermoelectric device that measures the temperature difference between the two cells (sample and reference cells) and a second sensor measures the temperature difference between the cells and the jacket. The result is given as the integral of the power required to keep the two cells at a constant temperature over time, which is equal to the change in heat resulting from the process.
The change in measured heat release upon ligand binding is directly proportional to binding strength. Ligand is titrated from the syringe in 30 injections into the sample cell containing aptamer solution. Once the aptamer is saturated with ligand, heat signal decreases. Calculated heat release from the injections gives either a hyperbolic or sigmoidal curve, which can be fit to one or two sets of binding site models, from which we can calculate the binding affinity (dissociation constant, $K_d$), stoichiometry, $\Delta H$ (enthalpy), $\Delta S$ (entropy) and $\Delta G$ (Gibbs free energy) of aptamer-ligand interaction (Duff, Grubbs, & Howell, 2011).
Before an ITC measurement, the equipment is thoroughly washed to avoid contamination or dilution. The sample cell is washed with 30 mL of ddH2O and 10 mL of binding buffer. The syringe is washed with 30mL of ddH2O and 10 mL of binding buffer as well.

The reference cell is filled with degassed ddH2O and sealed tightly.

Table 2.4 Sample preparation of aptamer solutions for ITC experiments

<table>
<thead>
<tr>
<th></th>
<th>Volume</th>
<th>Final Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aptamer</td>
<td>2.5 µL</td>
<td>2.5 µM</td>
</tr>
<tr>
<td>Binding Buffer</td>
<td>200 µL</td>
<td>20 mM (for NaP), 125 mM (for KCl), 5 mM (for MgCl2)</td>
</tr>
<tr>
<td>ddH2O</td>
<td>797.5 µL</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1 mL</td>
<td></td>
</tr>
</tbody>
</table>

Table 2.5 Sample preparation of titrant (ligand) solutions for ITC experiments

<table>
<thead>
<tr>
<th></th>
<th>Volume</th>
<th>Final Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neomycin-B</td>
<td>3 µL</td>
<td>60 µM</td>
</tr>
<tr>
<td>Binding Buffer</td>
<td>200 µL</td>
<td>20 mM (for NaP), 125 mM (for KCl), 5 mM (for MgCl2)</td>
</tr>
<tr>
<td>ddH2O</td>
<td>797 µL</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1 mL</td>
<td></td>
</tr>
</tbody>
</table>

Aptamer solutions were prepared to have a final concentration of 2.5 µM with each buffer system shown in Table 2.4. All the aptamers are single-stranded RNA aptamers,
so, to ensure that they are folded in their correct three-dimensional conformations after unfreezing, a refolding process is required. After aptamer solution is prepared in 1.5 mL Eppendorf tubes, they are placed in 95 °C heating block for 2 minutes to unfold all the nucleic acid chains, then it is cooled slowly on the bench to room temperature for 90 minutes to allow refolding.

Titrant solutions with neomycin-B ligand were prepared to have a final concentration of 60 µM in the same buffers as the aptamers as shown in Table 2.5. For a typical ITC experiment, the titrant is prepared in about 20-fold higher concentration than the macromolecule. This way proper saturation can be achieved, and the resulting curve would be fit well.

For each ITC experiment, the aptamer and ligand solutions were prepared according to the volumes given in Table 2.4 and Table 2.5. The tubes were carefully vortexed and spun down then the solutions are transferred to 5 mL glass vials. Both samples were degassed using a MicroCal ThermoVac sample degassing thermostat for approximately 5 minutes at 20 °C. This step is crucial for any ITC run, to avoid any air bubbles in the samples.

The samples were loaded in the MicroCal VP-ITC machine. The aptamer solution was transferred to the sample cell using a long needle Hamilton syringe. The injection syringe was filled with the titrant solution with the help of a plastic disposable syringe. The sample in the syringe was purged and refilled three times automatically using the VPViewer 2000 software.

The parameters for each ITC run are given in the Appendix A.
All the ITC assays were done in at least two repeats. The blank run for each experiment was performed by titrating the ligand samples into the buffer systems with same concentrations of ions.

The analysis of the obtained data was carried out using Origin 7 software. The data from the blank run was subtracted from the aptamer-ligand titration and the released heat values were fit to one or two sets of binding models to obtain the thermodynamic parameters, such as $K_d$, stoichiometry, $\Delta H$ (enthalpy), $\Delta S$ (entropy) and $\Delta G$ (Gibbs free energy).

2.2.2. Fluorescence Spectroscopy

Aptamers might obtain different conformations in their free and bound states in solution. Fluorophore-labelled aptamers work in signal-on or signal-off modes, depending on the structural change upon ligand binding either exposing or hiding the fluorophore, respectively. The change in fluorescence signal relatively shows the specificity of the aptamer towards the ligand. In these fluorescence assays, the aim is to test the ion-dependency of these aptamer applications. In addition, we may get some information about the effects of different ionic compositions on the conformational change of the same aptamers.

Two aptamers were labelled with 6-FAM (6-Carboxyfluorescein) fluorophore which has an excitation wavelength of 495 nm and an emission wavelength of 517 nm for fluorescence assays. 23m17NEO1A (Ilgu et al., 2014) was tagged at the 13th residue on its pentaloop, and 25m10NEO2A (from the same family as 24m5NEO2A) was tagged at the 7th residue on its internal bulge, as shown in Figure 2.2. Aptamers were prepared in final concentrations of 2.5 µM in the same buffers, as given in Table 2.6. The aptamer samples were placed in the heating block at 95 °C for 2 minutes to unfold
and slowly cooled on the bench at room temperature for 90 minutes to assure proper refolding.

*Table 2.6 Sample preparation for fluorescence assays*

<table>
<thead>
<tr>
<th></th>
<th>Volume</th>
<th>Final Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aptamer (10 µM in stock)</td>
<td>2.5 µL</td>
<td>2.5 µM</td>
</tr>
<tr>
<td>Antibiotic (10x in stock)</td>
<td>10 µL</td>
<td>Varying concentrations, NeoB (0.1 µM-10 µM), KanB (0.5 µM-50 µM), Amp (0.5 µM-50 µM)</td>
</tr>
<tr>
<td>Binding Buffer (2x in stock)</td>
<td>48.75 µL</td>
<td>20 mM (for NaP), 125 mM (for KCl), 5 mM (for MgCl₂)</td>
</tr>
<tr>
<td>ddH₂O</td>
<td>48.75 µL</td>
<td></td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>100 µL</strong></td>
<td></td>
</tr>
</tbody>
</table>

An aminoglycoside kanamycin and a penicillin type antibiotic ampicillin were used as control groups. Neomycin-B, kanamycin-B and ampicillin antibiotics were prepared in increasing concentrations as given in the Figure 2.4, in the same buffers as aptamers.

96 well Greiner black plates were used for fluorescence assays. Plate design is illustrated in Figure 2.4.
Fluorescence measurements are done using the SpectraMax Paradigm plate reader in two repeats. The average values of the obtained data are analyzed using R software.
3.1. ITC Results

As seen in the Figure 3.1, the comparison between 31NEO3A binding to neomycin-B in only NaP (left) buffer and in buffer containing Mg$^{2+}$ ions (right) showed significant difference in binding mode. Addition of 5mM Mg$^{2+}$ into the buffer system has led to a shift from two sets of binding to one set of binding model for this aptamer (indicated in red boxes). This might be due to Mg$^{2+}$ ions chelating around nucleic acids and...
blocking the second binding site or competitive binding of Mg\textsuperscript{2+} to the same site as neomycin-B.

![Figure 3.2 Binding curves for 31NEO3A-Neomycin B binding in buffers C and D](image)

Figure 3.2 shows the comparison of 31NEO3A-Neomycin-B binding in Na\textsuperscript{+} and K\textsuperscript{+} buffer (left), and in Na\textsuperscript{+}, K\textsuperscript{+} and Mg\textsuperscript{2+} containing buffer (right). The binding modes in these buffers are not significantly different, due to high concentrations of metal ions in both environments.

All the aptamers reacted differently to the change in surrounding ion concentrations.
24m5NEO2A binds to neomycin-B in two sets of binding sites model and with a high affinity in 20mM NaP buffer. Mg$^{2+}$ addition to the buffer leads to a decrease in affinity and a faster saturation rate.

Figure 3.3 Binding curves for 24m5NEO2A-Neomycin B binding in buffers A and B
In high monovalent ion concentration, the binding mode of 24m5NEO5A to neomycin-B changed to one binding site mode. The difference in the binding curves suggests that in high K$^+$ and Mg$^{2+}$ buffer, the saturation rate of the aptamer increased and as a result the affinity decreased.

Figure 3.4 Binding curves for 24m5NEO2A-Neomycin B binding in buffers C and D
30NEO4A-neomycin-B binding mode changed from two binding sites to one binding site mode upon addition of Mg\(^{2+}\). Fast saturation seen in the graph implies that Mg\(^{2+}\) addition may block one of the binding sites. In comparison, no significant change in ΔH was observed.
30NEO4A-neomycin B binding in high monovalent ion concentration was one binding sites mode and showed a very fast saturation rate. Between the binding graphs from high K⁺ buffer and Mg²⁺ buffer (Figure 3.5), no significant change was observed. But upon addition of Mg²⁺ into K⁺ buffer, ΔH decreased drastically due to Mg²⁺ binding. Heat release is caused by conformational change and ligand binding. Mg²⁺ compensates for the conformational change by stabilizing the nucleic acid chain. As a result, ligand binding becomes less ΔH dependent.

Figure 3.6 Binding curves for 30NEO4A-Neomycin B binding in buffers C and D
Figure 3.7 Binding curves for 27m2NEO5A–Neomycin B binding in buffers A and B

Binding mode of 27m2NEO5A-neomycin B interaction changed from two binding sites to one binding site models upon Mg\(^{2+}\) addition. But, no change in ∆H was observed.
No significant change was observed for 27m2NEO5A-neomycin-B binding in buffers C and D. Both graphs show one binding site modes and a fast saturation rate.

Table 3.1 $K_D$ values of aptamer-ligand binding in each buffer system

<table>
<thead>
<tr>
<th></th>
<th>NaP</th>
<th>NaP+KCl</th>
<th>NaP+MgCl₂</th>
<th>NaP+KCl+MgCl₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>31NEO3A</td>
<td>$K_D = 1.25 \pm 0.55 \mu M$</td>
<td>$K_D = 0.84 \pm 0.58 nM$</td>
<td>0.91 ± 0.19 µM</td>
<td>3.31 ± 0.07 µM</td>
</tr>
<tr>
<td>24m5NEO2A</td>
<td>N/A</td>
<td></td>
<td>1.28 ± 0.43 µM</td>
<td>1.27 ± 0.46 µM</td>
</tr>
<tr>
<td>30NEO4A</td>
<td>$K_D = 2.18 \pm 1.48 nM$</td>
<td>$K_D = 0.69 \pm 0.26 \mu M$</td>
<td>0.14 ± 0.03 µM</td>
<td>0.66 ± 0.05 µM</td>
</tr>
<tr>
<td>27m2NEO5A</td>
<td>$K_D = 5.12 \pm 2.84 nM$</td>
<td>$K_D = 2.19 \pm 2.03 \mu M$</td>
<td>0.63 ± 0.14 µM</td>
<td>1.24 ± 0.23 µM</td>
</tr>
</tbody>
</table>
Table 3.2 ΔS values of aptamer-ligand binding in each buffer system

<table>
<thead>
<tr>
<th>Buffer System</th>
<th>NaP</th>
<th>NaP+KCl</th>
<th>NaP+MgCl2</th>
<th>NaP+KCl+MgCl2</th>
</tr>
</thead>
<tbody>
<tr>
<td>31NEO3A</td>
<td>ΔS1 = -13.0 cal/K, ΔS2 = -41.6 cal/K</td>
<td>ΔS = -206.92 cal/K</td>
<td>ΔS = -207.46 cal/K</td>
<td>ΔS = -123.2 cal/K</td>
</tr>
<tr>
<td>24m5NEO2A</td>
<td>N/A</td>
<td>ΔS1 = -35.72 cal/K</td>
<td>ΔS = -24.8 cal/K</td>
<td>ΔS = -28.1 cal/K</td>
</tr>
<tr>
<td>30NEO4A</td>
<td>ΔS1 = -52.9 cal/K, ΔS2 = -49.5 cal/K</td>
<td>ΔS = -86.33 cal/K</td>
<td>ΔS = -95.69 cal/K</td>
<td>ΔS = -97.65 cal/K</td>
</tr>
<tr>
<td>27m2NEO5A</td>
<td>ΔS1 = -94.58 cal/K, ΔS2 = -37.53 cal/K</td>
<td>ΔS = -235.11 cal/K</td>
<td>ΔS = -144.64 cal/K</td>
<td>ΔS = -158.60 cal/K</td>
</tr>
</tbody>
</table>

Table 3.3 ΔG values of aptamer-ligand binding in each buffer system

<table>
<thead>
<tr>
<th>Buffer System</th>
<th>NaP</th>
<th>NaP+KCl</th>
<th>NaP+MgCl2</th>
<th>NaP+KCl+MgCl2</th>
</tr>
</thead>
<tbody>
<tr>
<td>31NEO3A</td>
<td>ΔG1 = 3.8 kcal, ΔG2 = 12.26 kcal</td>
<td>ΔG = 59.44 kcal</td>
<td>ΔG = 60.53 kcal</td>
<td>ΔG = 36.44 kcal</td>
</tr>
<tr>
<td>24m5NEO2A</td>
<td>N/A</td>
<td>ΔG1 = 10.54 kcal</td>
<td>ΔG = 7.30 kcal</td>
<td>ΔG = 25.95 kcal</td>
</tr>
<tr>
<td>30NEO4A</td>
<td>ΔG1 = 15.61 kcal, ΔG2 = 14.62 kcal</td>
<td>ΔG = 24.74 kcal</td>
<td>ΔG = 27.88 kcal</td>
<td>ΔG = 28.89 kcal</td>
</tr>
<tr>
<td>27m2NEO5A</td>
<td>N/A</td>
<td>ΔG = 66.30 kcal</td>
<td>ΔG = 43.26 kcal</td>
<td>ΔG = 159.01 kcal</td>
</tr>
</tbody>
</table>

Binding assays with ITC for these aptamers to neomycin B in each buffer were done in two repeats. The results are given as the average $K_d$ values in the Table 3.1. Increasing ion concentrations has led to significant decrease in binding affinities.

Calculated average entropy (ΔS) and Gibbs free energy (ΔG) values are given in tables 3.2 and 3.3, respectively.
3.1.1. Fluorescence Spectroscopy Results

The average values of the fluorescence signals from both repeated assays are given as a graph, shown in the Figure 3.9. Both aptamers showed limited binding to kanamycin-B and no binding to ampicillin, so their specificity to neomycin-B was confirmed. Also, for both aptamers, ion dependency was observed, although they differ in response to varying ions.

Kanamycin-B is the same type of antibiotic as neomycin-B and it is known that aminoglycoside aptamers in general may exhibit binding to other aminoglycosides more or less specifically. Fluorescence signals from the aptamers indicate binding to kanamycin-B, as expected, but the signals are decreased compared to neomycin-B binding. 25m10NEO1A in high ionic strength showed binding to Neomycin-B but not to Kanamycin-B.

Both aptamers showed no binding to ampicillin in any buffer system. In the presence of ampicillin, the increased fluorescence signal of 25m10NEO2A is due to diffuse binding of Mg$^{2+}$ ions.
Neomycin-B binding of 23m17NEO1A tended to decrease in high K⁺ concentration, but no significant difference was observed as shown in Figure 3.9 (a). 25m10NEO1A displayed best binding to neomycin-B in low metal ion concentration, but in contrast to 23m10NEO1A, significant decrease was observed upon addition of K⁺ and Mg²⁺. These results indicate that both of these aptamers are suitable for neomycin-B detection in solution under low ionic strength and fluorescence measurement can be implemented as the detection system.

Aptamers are affected by the surrounding ions differently. This might be due to their folding patterns in secondary and tertiary structures. Another criterion could be the selection of the site where fluorescent modification is applied. This may also change the aptamer responses to the ligands how they get affected by monovalent and divalent ions that surround residues differently.
Aptamers are functional oligonucleotides that can be used in diverse applications. As a result of being nucleic acids in nature, they are expected to be affected by the metal ions in their surrounding environment. The aim of this study was to investigate the behavior of aminoglycoside RNA aptamers under different sets of ionic conditions.

Binding affinity and specificity of aptamers are the two main characteristics that determine the suitability of the aptamers to detect a certain ligand. In this study, we utilized Isothermal Titration Calorimetry (ITC) and Fluorescence Spectroscopy methods to determine such characteristics of selected RNA aptamers.

We observed a significant decrease in affinity upon addition of divalent cations (Mg$^{2+}$) or high concentrations of monovalent cations (K$^+$) to the buffer system. ITC results showed that, even in low concentrations, addition of Mg$^{2+}$ causes a decline in binding affinity from nM to µM range. Fluorescence measurements also confirmed the ion dependency of aminoglycoside aptamers. They performed the best in low ionic conditions.

As a general statement, we can conclude that even though each aptamer reacted differently to the change ions in their surroundings, their binding affinities drop drastically when exposed to a buffer with high ionic strength. Similarly, their specificities are altered due to the potential conformational change caused by the surrounding ions. In the light of our results, it has become clear that understanding how the aptamers are affected by certain metal ions in solution is critical to get the best performance out of the aptamers. To sum up, the impact of the buffer components should be considered when selecting an aptamer and during the final application.
REFERENCES


Cate, J. H., & Doudna, J. A. (1996). Metal-binding sites in the major groove of a large


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APPENDICES

A. ITC Parameters

- Number of injections: 30
- Temperature: 25 °C
- Reference power: 18 μcal/sec
- Initial delay: 60 sec
- Syringe concentration: 0.060 mM
- Cell concentration: 0.0025 mM
- Stirring speed: 300 rpm
- Injection volume: 10 μL (first injection 1μL)
- Duration: 20 sec (first injection 2 sec)
- Spacing: 180 sec
- Filler period: 2 sec