Riboswitches as Potential Targets for Aminoglycosides Compared with rRNA Molecules: In Silico Study

Elnaz Mehdizadeh Aghdam1,2, Mohammad Esmaeil Hejazi3, Mohammad Saeid Hejazi4* and Abolfazl Barzegar4*

1Department of Pharmaceutical Biotechnology, Faculty of Pharmacy, Tabriz University of Medical Sciences, Tabriz, Iran
2Student Research Committee, Tabriz University of Medical Sciences, Tabriz, Iran
3Faculty of Medicine, Tabriz University of Medical Sciences, Tabriz, Iran.
4Research Institute for Fundamental Sciences (RIFS), University of Tabriz, Tabriz, Iran

Abstract

Riboswitches are cis acting riboregulators in non-coding region of the mRNAs. Their possible contribution in antibiotic targeting especially for FMN, TPP and lysine riboswitches in bacteria has been revealed since a decade ago. Regarding some studies on the possibility of the interaction between aminoglycosides and the artificial riboswitches, in this study we attempted to evaluate the binding potential of different types of aminoglycosides including gentamicin, amikacin, kanamycin, neomycin, tobramycin, sisomicin and paromomycin with various classes of riboswitches using computational methods. Applying Auto Dock vina, it was shown that the binding energy of each kind of riboswitches with different types of aminoglycosides (riboswitches/aminoglycosides) is almost similar or sometimes more than the binding energy of the aminoglycoside with the corresponding binding cage of “16S rRNA A site” (16S rRNA A site/aminoglycosides) as aminoglycosides’ target site. The affinity between riboswitches and aminoglycosides is almost the same or higher than the affinity of riboswitches/natural ligands. In this study ampicillin was used as the negative control antibiotic and 5S rRNA was employed as the negative control RNA. Results showed that the binding energies of riboswitches/ampicillin and SS rRNA/aminoglycosides are usually lower than the energy of riboswitches/aminoglycosides. Accordingly, lysine, glycine and SAM-I riboswitches were recognized as the best RNA targets for all of the aminoglycosides because of their higher binding energy. In the next step, docking results were further validated by rDock program. Furthermore, it was shown that hydrogen binding makes a key role in the binding energy between aminoglycosides and riboswitches. Moreover, MD simulation studies on lysine riboswitch/paromomycin complex confirmed the stability of the docked structure in the solvent containing magnesium and chloride ions.

Keywords: Riboswitches; Aminoglycosides; Docking; Binding energy; rRNA

Introduction

Riboswitches as non-coding sequences usually located in 5’UTR of mRNAs, are involved in gene regulation through binding to various small molecules without protein interpretation. Ligand binding to the conserved structure of riboswitches causes the folding shift of RNA molecule and halting of transcription and/or translation of downstream related genes [1]. Their unique characteristics on specific and selective binding to various molecules make them interesting biological devices. Since early 2000s, when the riboswitches were primarily introduced [2], the possibility of their application as antibiotic targets has been raised dramatically. At the beginning, the riboswitch-targeting mechanism of four well-known antibiotics including rosofavin [3,4], pyrithiamine [5], L-Aminoethylcysteine (AEC) and DL-4-oxalysine [6] were approved as ligands for FMN, TPP and lysine riboswitches. Afterwards, some efforts have been taken to find drug-like compounds through drug discovery methods in order to achieve some reliable effective antibacterials.

The aminoglycoside antibiotics are important therapeutic compounds in the treatment of severe bacterial infections. They exert their effects via binding to A site of 16S rRNA in 30S ribosomal subunit and cause interference in mRNA translation [7-9]. The presence of the caticonic amine groups of the aminoglycosides causes binding to the negatively charged pockets in folded RNA [10]. However, in addition to “16S rRNA A site”, it has been shown that aminoglycoside neomycin could bind to several other RNAs including the Trans-Activating Response (TAR) [11], Rev response element (RRE) RNAs of HIV-1[12] and catalytic RNAs, such as group I introns, RNase P, and the hammerhead and hepatitis delta virus ribozymes [13-16]. It is established that conformational changes in the RNA molecules can be occurred by drug binding at such sites [17,18]. The binding capability of the aminoglycosides to folded RNA structures has been applied to produce RNA aptamers [19]. In addition, designing artificial riboswitches for aminoglycosides have been conducted in the past decade [20-23]. For instance, a neomycin B binding artificial riboswitch was designed [20] which have later been studied structurally for the binding of ribostamycin and tobramycin [22]. Moreover, an in vitro interaction evaluation has been shown that some aminoglycosides inhibit the glmS riboswitch [24].

According to the literature, there have been some reports about the structural similarity between riboswitches and rRNAs which raises the...
possibility of functional connection between these two types of RNA molecules. In 2007 some structurally similar motifs to tRNA structures have been detected in riboswitches [25]. In 2008, an artificial riboswitch for neomycin B (an aminoglycoside antibiotic), was engineered [20] which partially resembles the ribosomal A site, the natural target for aminoglycoside antibiotics [26]. A comprehensive study on this similarity was carried out by our group [27]. Regarding the structural similarity between RNAs and riboswitches, the possibility of binding of paromomycin, as a representative of aminoglycosides, to different types of riboswitches was verified [27]. In this study we are aimed to evaluate and validate the binding potential of 7 aminoglycosides including paromomycin, gentamicin, amikacin, kanamycin, neomycin, tobramycin, sisomicin against 9 types of riboswitches through computational methods.

Methods

Molecular docking

Molecular docking is an important tool in structural biology and computational drug design. It is commonly used to calculate the binding modes of ligands and drug candidates to their protein/nucleic acid targets to predict the affinity and activity of the small molecule drugs [28]. Consequently, there is a wide range of uses and applications for molecular docking, including drug discovery and affinity prediction [29]. The most cited docking tool, namely Auto Dock [30] was used to predict the binding orientation of aminoglycosides to 9 types of riboswitches. The last version of Auto Dock namely Auto Dock Vina significantly enhances the average accuracy and speed of the binding mode predictions compared to Auto Dock 4 for molecular docking [31].

Preparation of Macromolecules for AutoDock Vina

Nine riboswitch classes which have not only the most representatives in microorganisms [32], but also have available PDB structures, were selected. Their PDB codes which represent preferably unbound state of riboswitches were extracted first from Rfam (http://rfam.sanger.ac.uk/) and then PDB (http://www.rcsb.org/pdb/home/home.do) databases. These riboswitch classes included TPP (PDB code: 2gdi), FMN (PDB code: 2yi), SAM-I (PDB code: 3iqn), lysine (PDB code: 3dxr), glycine (PDB code: 3ox0), purine (PDB code: 4eqs), c-di-GMP-II (PDB code: 3q3g), preQ1 (PDB code: 3fu2), THF (PDB code: 3sy5) riboswitches. In addition, the PDB structures of "16S rRNA A site" were obtained from 1j7t, 2et3, 2g5q, 2esi, 2ets, 1lc4, 4f8u codes for paromomycin, gentamicin, amikacin, kanamycin, neomycin, tobramycin and sisomicin respectively. 5S Ribosomal RNA structure was extracted from 1j7t, 2et3, 2g5q, 2esi, 2ets, 1lc4, 4f8u codes for paromomycin, as a representative of aminoglycosides, to different types of riboswitches was verified [27]. In this study we are aimed to evaluate and validate the binding potential of 7 aminoglycosides including paromomycin, gentamicin, amikacin, kanamycin, neomycin, tobramycin, sisomicin against 9 types of riboswitches through computational methods.

Preparation of the ligand for Auto Dock Vina

The structures of seven aminoglycosides including paromomycin, gentamicin, amikacin, kanamycin, neomycin, tobramycin and sisomicin were obtained through splitting of PDB codes of "16S rRNA A site" including 1j7t, 2et3, 2g5q, 2esi, 2ets, 1lc4, 4f8u respectively, using Viewer Pro Version 5. In addition, the specific ligand of each riboswitch was split from the complex structure of the riboswitch from PDB codes of 3do0, 3owi, 2y2i, 3iqr, 3suh, 3fu2, 4eqs, 3q3g, 2gdi for lysine, glycine, FMN, SAM, THF, preQ, purine, c-di-GMP-II and TPP molecules respectively. Besides, the structure of ampicillin as a negative control antibiotic was acquired from 4kr4 PDB code. All rotatable bonds within the ligands were allowed to rotate freely and Gasteiger charges were added to the obtained structure of the ligand using the Auto Dock Tools package 1.5.6.

Docking procedure

All dockings were performed using Auto Dock Vina 1.1.2 [31] which is a new generation of docking software from the Molecular Graphics Lab (http://vina.scripps.edu). The grid box of each riboswitch was set according to the similar part of the riboswitches with "16S rRNA A site" based on our previous study [27]. In addition, the grid boxes for various A sites were obtained based on the binding site of the aminoglycosides. The number of 20 modes was set for each docking run. Other parameters were kept to their default values. First, docking was carried out for every aminoglycoside and riboswitches. The pbdqt file of each docking procedure was created and the best binding energy was acquired. The binding energies of the aminoglycosides with related "16S rRNA A sites" and also the binding energies of the riboswitches with their own specific ligands were considered as positive controls. Besides, the interaction between riboswitches and ampicillin as well as the interaction of 5S rRNA with aminoglycosides were considered as negative controls. The interaction features of selected conformations were analyzed using AutoDockTools package 1.5.6.

Docking validation

rDock program is a fast and versatile docking tool for docking small molecules against for nucleic acids [33]. Docked conformations of paromomycin and gentamicin (as sample aminoglycosides) with highest binding energy in Auto Dock Vina were selected to be validated and rescored via rDock.

Docking preparation for rDock

At first the system definition parameter file was developed to define the receptor file, ligand and scoring functions. Cavity mapping was carried out based on "Reference ligand method" and the radius of cavity mapping region, radius of small probe, minimum cavity volume to accept, maximum number of cavities to accept, receptor atom radius increment for excluded volume and grid resolution for mapping, were considered 4.0 Å, 1.0 Å, 100 Å³, 10, 0.0 Å and 0.5 Å, respectively. Receptor and ligand files were prepared in Mol2 and SD formats, respectively, using Open Babel 2.3.2 program. Finally cavity mapping was performed using rbcavity (one of the executable programs of rDock). Besides, where needed rbrmsd was used to calculate the RMSD between pose predictions and the PDB structure.

Docking processing for rDock

Having done the cavity mapping, docking process was performed using rbdock with definition of input files including ligand, system definition parameter file, docking protocol file and output SD format file. The file dock_solv.prm was considered as the docking protocol file based on SF5 scoring function which is compatible with nucleic acids according to the reference guide [33].

Post-processing and analysis of results

In order to sort out the docking conformations based on the total score, sdsort program was utilized. The output SD file of rbdock was considered as input file for sdsort. Afterwards, the summary of scores (total, inter, intra and vdw) was obtained through sdsort program. According to the program guide, total score is a weighed sum of intermolecular, ligand intramolecular, site intramolecular and external
restraint terms. Inter score is the most important term as it represents the receptor-ligand interaction score. Intra score shows the energy difference between the ligand and the input ligand conformation.

**Molecular dynamics simulation**

In order to confirm the data from docking results we performed additional computational molecular dynamics (MD) simulation approach via GROMACS program [34]. For this aim, the complex (lysine riboswitch/paromomycin) was placed in a cubic box center of 12.164 nm, 12.164 nm, 12.164 nm with period boundary condition and solvated by TIP3P water molecules [35]. Mg$^{2+}$ and Cl$^{-}$ counterions were added to maintain overall system electroneutrality. The simulation was performed with GROMACS 5.0 suite program using CHARMM27 force field [36]. The Berendsen temperature coupling was used to keep the system at 300 K, and the constant of coupling was 0.1 ps. The Particle Mesh Ewald (PME) algorithm was applied to calculate long-range electrostatics interactions with a cutoff of 0.9 nm and a cutoff of 1.4 nm was set for Van der Waals interactions. The Verlet leapfrog integrator with an integration time step of 2 fs was used and LINCS algorithm was employed to keep all bonds involving hydrogen atoms rigid.

At first, the system was subjected to 500000 steps of steep descent energy minimization. Then, position restrained molecular dynamics was subsequently carried out for 200 ps. Finally, MD simulation was run for 2000 ps to the whole system.

**Results**

**Docking of riboswitches with various aminoglycosides**

Auto Dock Vina [37] is an open-source program for molecular docking. In comparison to Auto Dock 4, Auto Dock Vina significantly enhances the average accuracy of the binding predictions. Docking was performed on 9 types of riboswitches (lysine, glycine, purine, FMN, SAM, TPP, PreQ, c-di-GMP-II, THF) as receptors interacting with 7 types of aminoglycosides (paromomycin, gentamicin, amikacin, kanamycin, neomycin, tobramycin, sisomicin) as ligands using Auto Dock Vina. The rRNA molecule was used as a control for all of 7 types of aminoglycosides because of having "16S rRNA A site" as a known binding site for aminoglycosides. Therefore, the docked results for riboswitches/aminoglycosides were compared with the binding energy of docked antibiotics with "16S rRNA A site". Moreover, the corresponding natural metabolites of riboswitches were docked and used as a positive control (named own ligand) to assay antibiotic interactions with riboswitches (Figures 1-7). The RMSD calculated by rDock between predicted poses of natural ligands and their PDB structure were almost 1~2 Å. The interactions of some complexes including the SS rRNA/aminoglycosides and riboswitches/ampicillin were considered as negative controls. The binding energy of such complexes was almost 3-4 kcal/mol (Figure 1-7).

As seen in Figure 1, the binding energy of all of the riboswitches/paromomycin is approximately close to the binding energy of the interaction between rRNA A site/paromomycin. It was revealed that the binding energy of every kind of riboswitches/paromomycin is more (negatively) than the binding energy of riboswitches/natural ligands except for c-di-GMP-II riboswitch. In addition, the affinity of aminoglycosides/riboswitches is almost 2-fold of the affinity of riboswitches/ampicillin and SS rRNA/paromomycin (Figure 1-8). Besides, the binding energies of the riboswitches/aminoglycosides are comparable to those of riboswitches/natural ligands. Even the binding energies of the glycine, lysine and purine riboswitches/aminoglycosides are higher than those of riboswitches/natural ligands. In addition,
Figure 2: (A) Chemical structure of gentamicin. (B) Dark black columns demonstrate the binding energies of gentamicin with different riboswitches, dark grey columns show the binding energy of each riboswitch with its own natural ligands and light grey columns show the binding energy of each riboswitch with ampicillin based on Auto Dock Vina results. (C) The interaction between the riboswitches and gentamicin. Hydrogen bindings are shown as green dot lines.

Figure 3: (A) Chemical structure of neomycin. (B) Dark black columns show the binding energies of neomycin with different riboswitches, dark grey columns show the binding energy of each riboswitch with its own natural ligands and light grey columns show the binding energy of each riboswitch with ampicillin based on Auto Dock Vina results. (C) The interaction between the riboswitches and neomycin. Hydrogen bindings are shown as green dot lines.
Figure 4: (A) Chemical structure of kanamycin. (B) Dark black columns represent the binding energies of kanamycin with different riboswitches, dark grey columns show the binding energy of each riboswitch with its own natural ligands and light grey columns show the binding energy of each riboswitch with ampicillin based on AutoDock Vina results. (C) The interaction between the riboswitches and kanamycin. Hydrogen bindings are demonstrated with green dot lines.

Figure 5: (A) Chemical structure of amikacin. (B) Dark black columns demonstrate the binding energies of amikacin with different riboswitches, dark grey columns show the binding energy of each riboswitch with its own natural ligands and light grey columns show the binding energy of each riboswitch with ampicillin based on AutoDock Vina results. (C) The interaction between the riboswitches and amikacin. Hydrogen bindings are shown as green dot lines.
Figure 6: (A) Chemical structure of sisomicin. (B) Dark black columns show the binding energies of sisomicin with different riboswitches, dark grey columns show the binding energy of each riboswitch with its own natural ligands and light grey columns show the binding energy of each riboswitch with ampicillin based on AutoDock Vina results. (C) The interaction between the riboswitches and sisomicin. Hydrogen bindings are shown as green dot lines.

Figure 7: (A) Chemical structure of tobramycin. (B) Dark black columns demonstrate the binding energies of tobramycin with different riboswitches, dark grey columns show the binding energy of each riboswitch with its own natural ligands and light grey columns show the binding energy of each riboswitch with ampicillin based on AutoDock Vina results. (C) The interaction between the riboswitches and tobramycin. Hydrogen bindings are shown as green dot lines.
according to Figure 1B, lysine, glycine and SAM-I riboswitches have the best binding energies to paromomycin in comparison to “16S rRNA A site”. Figure 1C illustrates the 3D structure of the interactions between paromomycin and lysine, glycine and SAM-I riboswitches and also “16S rRNA A site”. The green dot lines illustrate the hydrogen binding between the ligand and the RNA molecule. It was shown that hydrogen bindings exist mostly between paromomycin and guanine and adenine nucleotides.

Approximately the same pattern was observed for other types of aminoglycosides (Figures 2-7). As seen, of 9 classes of riboswitches, lysine, glycine and SAM-I riboswitches showed higher binding energy to interact with different aminoglycosides. For some riboswitches, the natural metabolites binding energies are less than aminoglycosides’ binding energies and vice versa for others. For instance, c-di-GMP-II riboswitch showed higher affinity to its own metabolite ligand in comparison to the binding energies of paromomycin (Figure 1B), gentamicin (Figure 2B) and neomycin (Figure 3B). However, for kanamycin (Figure 4B), amikacin (Figure 5B), sisomicin (Figure 6B) and tobramycin (Figure 7B), in addition to c-di-GMP-II riboswitch, SAM-I riboswitch also showed higher binding energy with their own ligands. Regarding interaction between antibiotics with lysine, glycine, SAM-I riboswitches and “16s rRNA A site”, most hydrogen bindings make connection between the ligands and guanine/adenine nucleotides.

Docking validation

rDock program which can be used for docking against nucleic acids very efficiently, was applied to rescore docked conformations. The conformations with higher affinity were conducted as ligands to be docked once again with rDock program. The main jobs were performed by rbcavity (cavity generation) and rbdock (docking programs) [33].

As shown in Figure 8A, according to total scores, all of the studied riboswitches showed considerable affinity to paromomycin in the range of -16.83--55.16 for c-d-GMP-II and lysine riboswitches, respectively. Then, the best total score belongs to lysine and THF riboswitches with the total score of -55.16 and -43.93. Regarding total score of -16.49 for “A site rRNA”, it can be inferred that all studied riboswitches showed more affinity to paromomycin than its known target. Also, according to Figure 8B, gentamicin showed even higher affinity to different riboswitches in the range of -48.39--78.32 for lysine and SAM-I riboswitches, respectively. Then, lysine riboswitch is also the best target for gentamicin in comparison to other riboswitches. Moreover, the difference between the total scores of riboswitches and “A site rRNA” (-25.33) is more than the case of paromomycin. In addition, the intermolecular scores are higher than intramolecular scores in paromomycin. However, in the case of gentamicin intermolecular scores are lower which reflects some differences in the involved forces of binding. On the other hand, van der waals forces are approximately equal between two aminoglycosides/riboswitches bindings.

Molecular dynamics simulation

GROMACS 5.0 suite program was used to confirm the docking results and evaluate the interaction in an environment filled with water, Mg2+ and Cl ions. The PDB structure of docked lysine riboswitch and paromomycin was applied as the starting structure. In addition, separate ligand and riboswitch structure (with approximate distance of 6 Å) was simulated as negative control.

Figure 9A demonstrates the Root-Mean-Square Deviation (RMSD) of the whole system relative to the initial system during the simulation time. As shown, the system reaches equilibrium after 1500 ps and RMSD value of 8 nm.

Figure 9B and Figure 9C shows the distances fluctuation between C4 atom of paromomycin and P atom of A138 in lysine riboswitch. Results showed that, the distances fluctuation is very high in negative control (Figure 9B), whereas the distances variation is very low in docked form of complex (Figure 9C). In other words, the interaction between the aminoglycoside and the riboswitch is strong enough to keep the ligand in the complex form during the simulation.
Discussion

Non-coding RNAs have been considered as important elements for gene regulations in the cell more than a decade ago. A sort of cis acting riboregulators, so-called riboswitches, has attracted so many attentions in this regard since their discovery in 2002 [2]. Their structures consist of two essential parts including aptamer domain and expression platform. Structurally conserved aptamer domain binds to specific ligand and makes alteration in the dimensional structures of expression platform allosterically, followed by down-regulation or up-regulation of the corresponding genes [38]. Several types of riboswitches with specificity to particular ligands have been identified up to now [1]. However, lots are looking forwards to be discovered. The possibility of antibiotic targeting by riboswitches has been studied considerably. Lysine [6] and guanine [39] riboswitches are most studied targets for antibiotic designing. Although, the success to design completely effective antibiotics is rare [40], full efforts have been taken to improve drug discovery in this regard.

Aminoglycosides are important tools for battling severe infections. Their known mechanism is related to halting the translation through binding to 16S rRNA A site structure [7]. In addition, the binding of some aminoglycosides to artificial riboswitches has been investigated in the number of studies [20-23]. On the other hand, a comprehensive study on the similarity of rRNA and different riboswitches structures has been conducted and possible targeting of riboswitches by paromomycin has been suggested based on docking results [27]. Accordingly in this study, the affinity of other aminoglycosides was investigated and verified via AutoDock Vina and rDock programs, respectively. Moreover, a sample MD simulation was conducted to evaluate the interaction in water and ions environment. In this study, first the PDB structures of the representatives of nine kinds of riboswitches were extracted. It should be noted that based on different studies on bound and free-state of the riboswitches, the RMSD of the atoms are not high (approximately lower than 2 Å) [41]. It means that the cell environment itself could fold the riboswitches properly [42]. Therefore, in spite of the rare PDB codes for free-state riboswitches, docking on their bound structure could be acceptable as well. However, selecting the unbound state was preferable. As shown in Figures 1-7, the binding energies of different riboswitches with related aminoglycosides are approximately similar or more than the affinity of the 16S rRNA A site with the aminoglycoside. However the binding energy of the lysine, glycine and SAM-I riboswitches are significantly higher than those of other riboswitches. In the case of paromomycin, lysine, THF and SAM-I riboswitches were best receptors according to our previous study [27]. This could be due to differences in the scoring function of AutoDock 4 (mostly based on electrostatic forces) and AutoDock Vina (mostly based on hydrophobic and hydrogen binding). However, lysine and SAM-I riboswitches have been considered as better targets according to both methods. In addition, the affinity of riboswitches/ aminoglycosides is more than the binding energy of negative controls including riboswitches/ampicillin and 5S rRNA/aminoglycosides complexes (Figure 1-7). The exception of c-di-GMP-II riboswitch for all aminoglycosides and SAM-I riboswitch for kanamycin (Figure 4B), amikacin (Figure 5B), sisomicin (Figure 6B) and tobramycin (Figure 7B) demonstrates that these riboswitches may fail to bind aminoglycosides in competition with their known ligands. Binding characteristics of different aminoglycosides showed that strong hydrogen bindings exist between the aminoglycoside molecules and guanine/adénine nucleotides. In this regard, there is no difference between the binding of aminoglycoside/A site and aminoglycoside/riboswitch interactions.

rDock program as a valid and strong newly developed docking method for ribonucleic acids [33] have been utilized to recore the results of AutoDock Vina. Based on the observations (Figure 8) almost all riboswitches/aminoglycoside demonstrate higher total scores in comparison to rRNA/aminoglycosides in the case of paromomycin and gentamicin. Showing high total score, lysine riboswitch can be considered as the most important target for aminoglycosides which is in accordance with the results of Auto Dock Vina and AutoDock 4 [27]. However, all riboswitches illustrated better affinity to aminoglycosides in comparison to rRNAs. According to the results, although the binding modes of paromomycin and gentamicin may be different with each other, vdw forces are almost equal among riboswitches and aminoglycosides.

Taken the whole, it seems that though different riboswitches show affinity to aminoglycosides, some of them are particularly better than others in terms of higher binding energy such as lysine riboswitch verified by rDock scoring function.

It should be noted that molecular docking has a number of limitations beside its strength. For example, its dependence on the search algorithm and scoring function makes the performance of this approach quite different. Besides, the rigidity of binding site is not quite similar with what happens in reality. In addition, calculation of the binding energy in the vacuum condition without considering the effect of environmental molecules is an important limitation of docking method. Particularly, the presence of cationic ions such as Mg$^{2+}$ is so important for the behavior of the riboswitches in the cell. As a result, in the next step, the complex of lysine riboswitch/paromomycin was studied through MD simulation. MD simulation enabled us to evaluate the interaction in the mixture of water and ions, especially Mg$^{2+}$. Regarding the RMSD value the system is equilibrated. The result of the simulation showed that the distance of the ligand and the receptor was not varied considerably during the simulation, while the distance variation was considerably high in the negative control simulation (Figure 9). These findings approved the possibility of interaction between lysine riboswitch and paromomycin even in the ionic environment.

However, computational methods cannot replace experimental validation of the results and further in vitro and in vivo studies are needed to confirm the results.

Conclusion

In this study the binding affinity of 7 aminoglycosides including paromomycin, gentamicin, amikacin, kanamycin, neomycin, tobramycin and sisomicin to different types of riboswitches was evaluated and validated though docking methods and MD simulation. In conclusion, according to Auto Dock Vina results, almost all kinds of aminoglycosides have considerable affinities to the riboswitches. Among them lysine, glycine and SAM-I riboswitches showed stronger binding energies. In addition, the binding energies of riboswitches/ aminoglycosides are more than the binding energies of negative control complexes including riboswitches/ampicillin and 5S rRNA/ aminoglycosides. Additionally, most hydrogen bindings are created with guanine and adenine nucleotides. Docking validation through rDock program confirmed data as it was shown that the total score is higher in all riboswitches/aminoglycosides interactions in comparison with “16S rRNA A site”/aminoglycosides interactions. MD simulation study on lysine riboswitch/paromomycin complex approved the docking results even within the solvent in the presence of magnesium and chloride ions. Further computational and experimental studies are
suggested regarding interaction of riboswitches with aminoglycosides as well as consequences, effectiveness and efficiency of the interaction.

Acknowledgment

This article was written based on a data set of Ph.D thesis of Elnaz Meh dizadeh Aghdam registered in Tabriz University of Medical Sciences. This study has been supported financially under Ph.D thesis proposal submitted at No. 80 in the Faculty of Pharmacy, Tabriz University of Medical Sciences, Tabriz, Iran.

References