Molecular Docking Studies of Phytochemicals of *Vitex Negundo* (L.) Against Adenosine A1 Receptor as Therapeutic Target in Cardiovascular Diseases

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Abstract

Recent days, Ayurveda medicine is becoming one of the best alternatives for the modern medicines for effective control of cardiovascular diseases (CVD) due to their limited side effects and ease of availability to a common man. *Vitex negundo* (L.) is an aromatic shrub known to possess the active phytochemicals such as 5,3′-dihydroxy-7, 8, 4′-trimethoxyflavanone; 5,3′- dihydroxy-6, 7, 4′-trimethoxyflavanone; 5-hydroxy-7,4′-dimethoxy flavone; betulinic acid; ursolic acid; n-hentriacontanol and β-sitosterol in its leaves which could be useful to treat CVDs. Adenosine A1 receptor (AAR) is a well known drug target of CVDs where its inhibition by AAR inhibitors is an ideal approach to control CVDs. Homology model of AAR was constructed and its stereo chemical quality was validated and its potential binding sites were explored using CASTP server. The compounds were successfully docked into the predicted binding site of AAR using LibDock module of Discovery studio software. Their affinities and binding mode orientations of the seven compounds in AAR binding site revealed that they can be used as best AAR inhibitors to control and manage CVDs.

Keywords: 5,3′-Dihydroxy-7,8,4′-trimethoxyflavanone; Adenosine A1 receptor; CVD; Molecular Docking

Introduction

Cardiovascular diseases (CVD) remain the principal cause of death in both developed and developing countries accounting roughly 20% of all deaths worldwide. It is estimated that 17.5 million people were died due to CVDs in 2012, representing 31% of global deaths. Among all these deaths, 7.4 million were because of coronary heart disease and 6.7 million were because of stroke[1,2]. Recent epidemiological survey indicated that urbanization and changes in life style are main causes for increase of CVDs. It is expected that by the year 2030, more than 23.3 million people will die annually due to CVDs. Asian countries are the home to 20% of the world’s population with the highest burden of CVDs in the world[3]. Myocardial Infarction occurs when the supply of blood to a part of the heart is interrupted. This is most commonly due to atherosclerotic plaque, which is an unstable collection of cholesterol and WBC (macrophages) in the wall of an artery[4]. The risk factors are family history, ethnicity and age, which cannot be controlled. Other risk factors that can be controlled include smoking, hypertension, obesity, high cholesterol, physical inactivity, type 2 diabetes, unhealthy diets, oxidative stress, lipoproteins, marker enzymes and harmful use of alcohol etc[5,6].

It is well known that, Adenosine is a key endogenous molecule and potentially important signaling molecule in the heart. The adenosine production by cardiac myocytes reflects the metabolic state of the myocardium. Adenosine acts as regulator of vasodilation in the coronary resistance vessels, thereby coupling blood flow to the energetic needs of the heart[7]. It is well known that adenosine receptors are important therapeutic targets for cardiovascular diseases[8]. Adenosine regulates tissue function by activating four G-protein-coupled adenosine receptors: A₁, A₂A, A₂B and A₃; which control signaling pathways involved in regulating many body functions, especially in the cardiovascular system[9]. Activation of Adenosine A1 receptor (AAR) protects the heart from oxy-

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gen deficiency and slows down the heart rate, while activation of the A2 receptors can improve the flow of blood to the heart and lower blood pressure. These pathways are involved in inducing the CVDs leading to complications and push the life of an individual into the risk\[10,11\]. Hence, ceasing the activity of AAR in this metabolic pathway is suggested to be an ideal approach to prevent CVDs.

Natural compounds have been widely used for the discovery of many important modern drugs with novel mode of action\[12\]. \textit{V. negundo} Linn. (Verbenaceae, Synonyms-Indian Privet; Nirgundi; Bana) is a woody, large aromatic shrub growing to a small tree. Different parts of \textit{V. negundo} have cardio-tonic, hepato-protective, antioxidant activity, anti-inflammatory, analgesic, cytotoxic activity, antihistamine property, antiulcerogenic, antiparasitic, antimicrobial, anti-fungal, antiviral activity and anti-arthritic potentials\[13-23\]. Leaves of \textit{V. negundo} contains phyto-compounds such as 5,3’-dihydroxy-7,8,4’-trimethoxyflavanone, 5,3’-dihydroxy-6,7,4’-trimethoxyflavanone, 5-hydroxy-7,4’ dimethoxy flavones, 5, 3’- dihydroxy-7, 8, 4’-trimethoxy flavanone, betulinic acid, ursolic acid, n-hentriacontanol and β-sitosterol\[24\]. There exist no reports so far on the bioactivity of \textit{V. negundo} with cardioprotective property and hence, in the present study we aimed to identify the cardio protective role of the phyto compounds of \textit{V. negundo} through molecular modeling studies.

Methodology

Ligand Study

The leaves of \textit{V. negundo} was reported to possess the active principles such as 5,3’-dihydroxy-7,8,4’-trimethoxyflavanone; 5,3’-dihydroxy-7,8,4’-trimethoxyflavanone; 5-hydroxy-7,4’ dimethoxy flavones ; 5, 3’- dihydroxy-7, 8, 4’-trimethoxy flavanone; betulinic acid; ursolic acid; n-hentriacontanol and β-sitosterol were 24 (Figure 1). We aimed to identify the inhibitory action of these compounds against AAR through molecular modeling studies. Initially the chemical structures of the compounds were retrieved from PubChem database and their three dimensional models were constructed in Discovery studio graphical environment and subjected to energy minimization in CHARMm force field to an RMS gradient of 0.01Å. The low energy and stabilized conformations of the compounds were saved and used for further studies.

Sequence Retrieval and Template Identification

The protein sequence of AAR from Homo sapiens (ID: NP000665) was retrieved from NCBI and similarity search was carried out using BLAST-P server against PDB database. The hit with highest identity and coverage was taken as template to construct the homology model of AAR.

Homology Modeling of AAR

The homology model of the AAR was constructed using Modeller 9v11 software tool\[25\]. The Blast-P similarity search AAR against PDB revealed that a maximum identity of 51% was found with Human Adenosine A2 a Receptor that can be used as a template to construct the homology model of AAR. A sequence alignment file was generated between the template and AAR sequences and mentioned in python modeling script. The script file along with template structure was submitted to Modeller to run the script. A total of 20 models were generated and among them the model with lowest DOPE (Discrete Optimized Potential Energy) was chosen as best model.

Protein Processing and Preparation through Molecular Dynamics Simulation

The structure constructed by using Modeller was optimized in Discovery studio software using CHARMm force field. The structure was energy minimized in 10,000 steps at a spherical cutoff of 12 Å for Vander Waals interactions and no periodic boundary conditions were added providing the implicit solvent environment. The energy minimized structure of AAR was then prepared using protein preparation wizard to clean up the common internal errors of the structure. Loops were defined with a maximum length of 20 residues with subsequent CHARMm minimization. The structure is then protonated with a protein dielectric constant of 10, pH 7, ionic strength of 0.145 M and with an energy cutoff of 0.9 kcal/mol. The protonated structure
was subjected to molecular dynamics simulations using standard dynamics cascade. The simulations were started with a primary minimization step using Steepest Descent algorithm to improve the poor conformational regions and followed by secondary energy minimization step using Conjugate Gradient algorithm to an RMS gradient of 0.1Å in a total of 2000 steps. The system was heated from 30 K to 300 K in a time steps of 100 picoseconds (ps) with a velocity frequency of 50 steps. Further the system was equilibrated in 100 ps and finally followed by production phase of 10000 ps. The generated conformations were studied for their total energies and RMSD to look into the stability of the system. The low energy conformation obtained at the end of the simulation was trapped from the total trajectories and used for the further studies.

Structure Validation
The stereo chemical quality of the final structure was analyzed by Ramachandran plot using PROCHECK validation server[20] and Z-Score plot of ProsaWeb server[21].

Binding Site Identification
CASTp online server was employed to predict the possible binding sites on the AAR structure. CASTp identifies and measures pockets and pocket mouth openings, as well as cavities. This program specifies the atoms lining pockets, volume and area of pockets and cavities; pocket openings, and buried cavities; and the area and circumference of mouth openings[22]. Further the functional patterns were also predicted by submitting the AAR protein sequence to PROSITE secondary database[23]. The idea behind the usage of these two servers is the correlation of results from both sources will help to predict more reliable binding site that can be targeted with the compounds.

Molecular Docking Studies
Docking studies were carried out using a high-throughput algorithm, LibDock of Discovery Studio. The energy minimized compounds were docked into the predicted binding site of AAR homology model with an XYZ coordinates of -51.24, -1.47 and -33.11 respectively. Ligand conformations were aligned with polar and non-polar receptor sites which are called as hotspots. A maximum of 100 hot spots were defined with a docking tolerance of 0.25 Å to accept or reject any specific conformation. BEST conformation method was applied to ensure the maximum conformational space of the ligand in the binding site. Multiple docking conformations were generated for each ligand and evaluated by LibDock scores. The docked conformations were subsequently energy minimized to an RMSD cutoff of 1Å in 1000 maximum steps. Energetically, the most favorable conformation was selected and their intermolecular interactions in the active site of AAR were analyzed in the best energy docked conformations[24].

Results and Discussion
Homology Modeling of AAR
A high percentage of sequence similarity should be more accurate alignment between the target sequence and template structure. In our study, we have chosen crystal structure of human adenosine A2a receptor as a reference structure for constructing AAR model (PDB ID: 3VG9). A total of 20 mod-
Figure 3: (A) Ramachandran plot showing the stereo chemical quality of the AAR model. Glycine residues are shown as triangle, Proline residues as square and remaining residues as circles. (B) ProsaWeb plot of AAR homology model showing a Z-score of -3.96 falling in the range of native X-ray crystallographic region.

Prediction of Binding Site

Multiple binding sites were predicted in the stabilized model of AAR using CASTp server. PROSITE results revealed that the AAR protein contains G-protein coupled receptors family 1 signature ranging from 93-109 residues. These residues were in correlation with the CASTp results and hence the same residues such as Ser93, Ser94, Ile95, Leu96, Ala97, Leu98, Leu99, Ala100, Ile101, Ala102, Val103, Leu107, Arg108 and Val109 were chosen as binding site. These residues were also observed to contain a perfect binding cavity conformation in the AAR structure and hence chosen as the more favorable sites to dock the ligands (Figure 4).

Figure 4: Homology model of AAR showing the predicted binding site. The AAR structure is represented in cartoon model and the binding site in sphere surface. Left: lateral view; Right: Top view. The binding site is confined to the pore region of the model.

Molecular Docking

The seven compounds were successfully docked into the predicted binding cavity of AAR model using LibDock module. Multiple conformations were generated for each compound and the conformations were ranked using LibDock score. The observation of LibDock score revealed that Lig1 showed a highest docking score of 111.36 kcal/mol and the lowest docking score of 70.47 kcal/mol was observed for Lig5 (Table 1). According to LibDock algorithm, higher the docking score higher the strength and lower the docking score lower the strength of the docking complex. So, it can be inferred from the docking scores that, Lig1 is having most binding affinity among all the compounds and could best inhibit the AAR activity.

Table 1: Molecular docking of Phytochemicals of *Vitex negundo* (L.) against human AAR protein structure.

| Ligand        | Name                                      | LibDock Score (kcal/mol) |
|---------------|-------------------------------------------|--------------------------|
| Lig1          | 5,3′-dihydroxy-7,8,4′-trimethoxyflavone    | 111.36                   |
| Lig2          | 5,3′-dihydroxy -6,7,4′-trimethoxyflavone   | 90.84                    |
| Lig3          | 5-hydroxy-7,4′-dimethoxyflavone            | 110.20                   |
| Lig4          | Betulinic acid                            | 80.15                    |
| Lig5          | Ursolic acid                              | 70.47                    |
| Lig6          | n-hentriacontanol                         | 70.83                    |
| Lig7          | β-sitosterol                               | 90.48                    |

These compounds were observed to interact with the AAR binding site by means of hydrogen bonding and hydrophobic interactions. Lig1 was observed to form hydrogen bonding with Ala43, Thr44 and Arg105 residues using its oxygen reactive centres. Such kinds of hydrogen bond interactions were also observed in Lig2, Lig3 and Lig7. The residue, Arg123 of AAR was observed to play a vital role to interact with the ligands by means of hydrogen bonding. Hydrophobic interactions were observed with Lig4, Lig5 and Lig6, which were favored by their aromatic rings (Figure 5). The hydrophobic residues such as Ala100, Val103, Tyr106, Val118, Val119 and Ala127 were observed to form hydrophobic spheres with the ligands. Hydrophobic interactions represent more stable interactions than hydrogen bonds. The compounds, which did not formed hydrogen bonds, were stabilized by hydrophobic interactions by orienting towards the hydrophobic amino acid residues in the AAR binding site. Although there exists variable docking scores among seven phytochemicals, the orientation of the compounds was observed to fit snugly in the binding site favoring their strong binding with AAR. Such a molecular interactions and affinity levels of these seven phytochemicals explains their inhibitory action against AAR protein. These results will finally suggest their use as AAR inhibitors and further their in vitro evaluation is needed to be ascertained.

Figure 5: Molecular interactions of phytochemicals of *Vitex negundo* (L.) in the binding site of AAR model. The ligands are shown in CPK colors and the AAR amino acid residues are shown in orange color stick models.
In the present study we aimed to identify the phytochemicals of *V. negundo* (L.) as AAR protein inhibitors so as to use for the control and management of CVDs. As a preliminary study, we have modeled the AAR protein structure by homology modeling method and stereo chemical quality was validated. A stabilized trajectory of the model was obtained by molecular dynamics simulations and a potential binding site was predicted using CASTp and PROSITE servers. Molecular docking studies of the seven phytochemicals using LibDock module revealed their potential (5,3′-dihydroxy-7,8,4′-trimethoxyflavanone) to inhibit AAR protein and in turn suggesting their use for the treatment and management of CVDs.

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