DOCKING STUDY OF METHYL HESPERIDIN AS NUCLEOSIDE REVERSE TRANSCRIPTASE INHIBITOR

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INTRODUCTION

Acquired immunodeficiency syndrome (AIDS) is a global and deadly disease in the world. Based on the global report of the World Health Organization (WHO), the amount of death in the world caused by AIDS reached 1.8 million people [9]. Based on the data from the Health Ministry of Republic Indonesia, the incident number of AIDS in September 2014 reached 55,799 people in Indonesia [8]. Drug therapy for AIDS known as Retroviral Drug now has resistance and side effect problems affecting medication [10, 16, 21]. Therefore, there is a necessity to discover a new agent as anti-HIV because the drug discovery is begun from a plant [11].

Orange plant, one of a typical Kalimantan Barat plant, has a potential to be developed into anti-HIV because this plant produces a few secondary metabolites such as hesperidin, tangeretin, hesperitin, naringin and naringenin. Hesperidin has been proved as antidiabetic, anticholesterol, antiinflammation, antioxidant and antifertility [6]. There are no toxic effects found in the trial of male and female mice. Moreover, hesperidin could be easily digested and is not accumulated in the body and hesperidin also does not cause an allergic reaction [5]. In another trial, 70% ethanol extract of Justicia gendarussa Burm. f leaves had been proved to inhibit the activity of reverse transcriptase HIV-1 and apigenin is the main secondary metabolite detected in the extract [4]. Apigenin had also been proved to inhibit the activity of reverse transcriptase HIV-1 as a competitive substrate, in vitro [3]. Hesperidin has a similar chemical structure with apigenin. They are flavonoid that has three rings with same chemical structures. Based on these data, hesperidin could possibly inhibit the activity of reverse transcriptase HIV-1.

Reverse transcriptase has an important role in the HIV life cycle because of the enzyme reversely transcript ribonucleic acid (RNA) virus into deoxyribonucleic acid (DNA) virus. DNA of virus nest is incorporated into the DNA of the host cell and when host cell transcribes the code area containing this DNA, proteins needed to create a new virus are produced. All of those proteins and enzymes needed to create a new virus are originated from the host cell. Only reverse transcriptase, integrase and virus RNA are originally brought by the virus [7]. Therefore, the inhibition of the activity of reverse transcriptase HIV-1 could stop virus replication.

Although hesperidin has a potency as an anti-HIV activity, hesperidin could not be absorbed well in the body. Solubility, one of the physicochemical properties of hesperidin is low in the body causing low bioavailability. The solubility in water, stomach and intestine are respectively 57 mg/l, 49,1 mg/l and 68,0 mg/l [2]. Log P of hesperidin are 0,30 [1]. Therefore, there is a necessary to improve the solubility of hesperidin. Chemical structure modification is a way to improve hesperidin solubility by methyl addition, a nonpolar functional group, into hesperidin chemical structure and also eliminates the glicon functional group from the structure.

It is needed to analyze that the addition of methyl into the chemical structure does not change the anti-HIV activity of hesperidin. The one of research methods could be chosen is a computerized method. The method could predict the solubility and the affinity of interaction between ligand and receptor. This study aims to analyze the physicochemical properties related to hesperidin solubility and permeability properties in the body and the docking study of methylated hesperidin as a competitive substrate of reverse transcriptase HIV-1.

MATERIALS AND METHODS

Materials

Hardware used for physicochemical property analysis and docking study of ligand is Notebook PC ASUS X455L Series with the specifications of Intel Core i3-4030U CPU 1,90 GHz, system operation Windows 8.1 Enterprise with 64-bit windows and 2GB of RAM. Software used for physicochemical property analysis of ligand is ChemDraw 15.0. Software used for docking study of ligand is Autodock Vina with Autodock Tools 1.1.2 version and software used for docking visualization of ligand-receptor interaction is Discovery Studio.

Docking methodology

2 dimensions (2D) and 3 (3D) dimensions, the chemical structure of ligand was obtained by drawing ligand structure and setting the
minimal sterical energy of ligand structure. Receptor, reverse transcriptase HIV-1 substrate, was downloaded from RCSB Protein Data Bank (http://www.rcsb.org/pdb/home/home.do). Grid parameter was set using Autodock Tools to coordinate 18.177 x 25.99 x 37.741 Å (Amstrong) and dimension 30 x 30 x 30 Å. Grid dimension was set same between the ligand and positive control. Docking study was calculated using Autodock Vina with the grid parameter and type of interaction was visualized using Discovery Studio.

The analysis of physicochemical properties

Physicochemical properties of ligand were analyzed using Chem Draw 15.0 [12].

Data analysis

The analysis result of ligand physicochemical properties (logP, BM, H H acceptor and donor) are analyzed using the rule of five Lipinski to predict the properties of ligand absorption and permeability. The affinity of ligand-receptor interaction was analyzed using Autodock Vina and Discovery Studio to analyze ligand-receptor binding energy and amino acid residues involved from ligand-receptor interaction (kcal/Mol) [18-20].

RESULTS AND DISCUSSION

Chemical structure of modification is a part of drug development. Addition of functional group of atoms in the chemical structure of a compound would change the polarity and affinity of a compound to a receptor [12]. In order to certain, the effect of methyl addition into hesperidin against the solubility and the affinity of methyl hesperidin to the receptor, physicochemical property analysis and docking study was conducted in this research based on a very large mathematical operation in computational chemistry. The method integrates chemical theory and modelling with experimental observations based on mathematical algorithms, statistics and databases. It calculates the fundamental laws of physic of an atom such as charged particles, positive nuclei, negative electrons and geometrical arrangements of the nuclei correspond to stable molecules, and how atoms interact each other. Therefore, this method could contribute as preliminary studies for supporting in vitro and in vivo experimental method [18].

3D chemical structures of hesperidin, methyl hesperidin and (positive control) zidovudine obtained using ChemDraw and after minimizing energy are shown in fig. 1.

Minimizing the energy of ligand structure obtained using Chem Office program is conducted to obtain the most stable conformation of the ligand that are close to the real ligand binding to the receptor in the human body. In order to obtain the confirmation of a compound that is as similar as the one in the nature, the minimizing energy would give the best arrangement of atoms position in a space for a compound so net inter-atomic force and sterical energy of each atom are close to zero and the position on the potential energy surface is a stationary point. The confirmation of structure would affect the real physicochemical properties of a compound. It also affects the docking score showing the affinity of ligand-receptor [13].

The analysis result of ligand physicochemical properties showed that log P of methyl hesperidin is more than hesperidin. Same as hesperidin, methyl hesperidin have a molecular weight less than 500 and C log P is less than 5. The amount of NH functional group for Amida are 0, and O-H functional group for carboxylate and phenol are 2, the total are 2, means less than 5. The amount of O atom is 5 and N atom are 0, so the total are 5, means less than 10. According to Lipinski Rule, these could be predicted that hesperidin and methyl hesperidin could be absorbed and has good permeability.

Lapinski Rule reveals that a drug would be pharmacology active if it has physicochemical properties contributing to drug pharmacokinetics in the human body including absorption, distribution, metabolism and excretion. This rule proved based on empirically screening program for the most oral drug that a range of physical properties is consistent with oral activity such as Acyclovir, Alprazolam, Proxiam, Vinblastin and 2000 others. In general, a drug has low absorption and permeability because the molecular weight is more than 500 Dalton, an octanol-water partition of coefficient log P is greater than 5, hydrogen bond acceptors of a drug are more than 10 for nitrogen or oxygen atoms, and hydrogen bond donors for a drug are more than 5 for nitrogen or oxygen atoms with one or more hydrogen atoms [12].

Minimizing the energy of ligand structure obtained using Chem Draw 15.0, A): Hesperidin, B): methyl hesperidin, C): zidovudine

Both ligand, methyl hesperidin, hesperidin and positive control were optimized using Autodock Tools to determine various factors contributing to ligand affinity to receptors such as partial charges, the addition of both polar and nonpolar hydrogen atoms, rotatable bonds, the number of torsional degree, identifying an atom as a central root and the torsion count.

Crystal structure of the receptor was downloaded from RCSB Protein Data Bank (http://www.rcsb.org/pdb/home/home.do), in the form of reverse transcriptase HIV-1 complex with DNA and zidovudine, with the code of receptor 3V4I. The receptor was optimized using Autodock Tools such as eliminating water, adding hydrogen atoms and partial charges.

Grid parameters for methyl hesperidin and hesperidin were set close to positive control grid box by choosing the similar dimension area. It was conducted in order to keep the docking result is in the active site of the receptor. The active site of receptor chosen are amino acid residues affecting the function or activity of the enzyme.

The binding energy score of hesperidin and methyl hesperidin against the solubility and the affinity of methyl hesperidin to the receptor cause releasing energy called exotherm. Binding energy shows the inter-atomic force, the sterical energy and potential energy resulted from ligand-receptor binding. The lower energy binding score from ligand-receptor binding, the more stable the ligand-receptor to bind. This because the better affinity of ligand-receptor is determined by the stability of ligand-receptor binding[13]. The RMSD score shows the iteration of conformation model of ligand against the receptor with the binding energy score shown in table 1 is significantly not different [15]. It could be predicted that methyl hesperidin, hesperidin could be bound to the receptor as positive control.

Fig. 1: Ligand 3D structure obtained using ChemDraw 15.0, A): Hesperidin, B): methyl hesperidin, C): zidovudine

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Table 1: Binding energy and binding site obtained using autodock vina 1.1.2.

<table>
<thead>
<tr>
<th>Ligand</th>
<th>Binding energy score (kcal/Mol)</th>
<th>RMSD</th>
</tr>
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<tbody>
<tr>
<td>Hesperidin</td>
<td>-8.0</td>
<td>0.000</td>
</tr>
<tr>
<td>Methyl Hesperid</td>
<td>-8.8</td>
<td>0.000</td>
</tr>
<tr>
<td>Zidovudine</td>
<td>-9.3</td>
<td>0.000</td>
</tr>
</tbody>
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Hesperidin, methyl hesperidin and Zidovudin could inhibit the activity of the enzyme, reverse transcriptase. This could be proved first by comparing the hesperidin and methyl hesperidin binding energy score with a proven antiretroviral drug, Zidovudin. Second, the relationship between binding energy score with the bioactivity of the drug is shown in a docking study of Glimepirid and Glibenclamid, antidiabetic drugs with the binding energy score that is lower than zero [19]. Moreover, another study shows the relationship between the binding energy score with bioactivity study. A compound in Buddleja salviifolia (L). Lam leaf showed the binding energy score is lower than zero and has bioactivity as anti-inflammatory and antioxidant [20].

Type of interaction between ligand, hesperidin and receptor, DNA, using Discovery Studio is shown in fig. 2.

Type of interaction between ligand, methyl hesperidin and receptor, DNA, using Discovery Studio is shown in fig. 3.

Fig. 2: Visualization of hesperidin-receptor interaction using discovery studio, A): Type of interaction between hesperidin and receptor, B): Hesperidin is in the position of the receptor substrate

Fig. 3: Visualization of methyl hesperidin-receptor interaction using discovery studio, A): Type of interaction between methyl hesperidin and receptor, B): Methyl hesperidin is in the position of the receptor substrate

Type of interaction between methyl hesperidin and receptor is more than hesperidin and the receptor as shown in fig. 2 and 3. This contributes to the different binding energy score between both ligands. Both ligands could be bound in the active site of the substrate position of receptor shown in fig. 2 and 3. Because both ligands have no atoms that could facilitate phosphate bond to the next incoming substrate, both ligands could act as stop codon for reverse transcription of virus RNA into DNA [14].

Type of interaction between positive control, zidovudine, and DNA using Discovery Studio are shown in fig. 4. Zidovudine has the most interactions contributing to the greatest zidovudine binding energy score. Zidovudine could be bound in the substrate position of the receptor. Because zidovudine has no atoms that could facilitate phosphate bond to the next incoming substrate, zidovudine could act as stop codon for reverse transcription of virus RNA into DNA [14].
Hesperidin and methyl hesperidin could inhibit the activity of reverse transcriptase because of both ligand bind to an amino acid residue in the active site of the substrate. There is a relationship between the binding sites of docking studies with the bioactivity of the drug. In a study of proven antidiabetic drugs showed that Glimipride bind to the different kind of amino acid residue with other antidiabetic drug but both ligand has good bioactivity [19]. In other studies showed that there is a relationship between docking studies and the bioactivity of the compound in Buddleja salviifolia (L.). Lam leaf [20], and the consistent result between docking result and in vitro antibacterial of N-[2-chloroquinolin-3-yl] methylidenel-2-aniline [17]. This relationship could be obtained because the different kind of amino acid residue is located in the active site of receptor whose receptor activity.

In order to obtain the information about the methyl hesperidin activity as anti-HIV, especially related to the quantity of compound that could inhibit the activity of reverse transcriptase, we need to conduct a research experiment. Computational chemistry is a qualitative method used to determine whether or not atoms of a ligand interaction with atoms of a receptor. The method was conducted only based on the calculation of theoretical approach [18].

CONCLUSION

The results first showed that according to Lapinski Rule, methyl hesperidin has physicochemical properties contributing to drug pharmacokinetics in the human body. It could be predicted that methyl hesperidin would be pharmacology active. The second showed that the binding energy of methyl hesperidin is lower than zero, as zidovudine, a proven antiretroviral drug. It means that methyl hesperidin would be pharmacology active. The second pharmacokinetics in the human body. It could be predicted that methyl hesperidin would be pharmacology active. The second pharmacokinetics in the human body. It could be predicted that methyl hesperidin would be pharmacology active.

REFERENCES


