Molecular Dynamics Simulation of SIRT1 Inhibitor from Indonesian Herbal Database

Andika1, Linda Erlina1, Azminah1, Arry Yanuar1
1Faculty of Pharmacy, Universitas Indonesia, Depok, West Java, INDONESIA.

ABSTRACT
Objective: Sirtuins are protein deacetylases regulating cellular metabolism, lifespan, stress responses, and linked with diseases pathogenesis such as cancer and neurodegenerative diseases. SIRT1, one of seven sirtuins members, is characterized by a conserved catalytic core and unique additional N-terminal and/or C-terminal sequences of variable length.1

Conclusion: Hits compounds dregamine and 5-oxocoronaridine obtained free energy MMPBSA calculation about -23 kcal/mol while occupancy hydrogen bonding of residues Ile347 and Asp348 about 80%.

Key Words: Sirtuin, SIRT1, Pharmacophore, Herbaldb, Molecular Dynamics Simulations.

INTRODUCTION
Molecular dynamics (MD) simulations method is one of the first of pioneering applications for fluid dynamics by Alder and Wainwright and by Rahman in the late 1950s and early 1960s. Because of the revolutionary advances in computer technology and algorithmic improvements, MD then become a valuable tool in many fields of physics and chemistry. Since the 1970s the MD has been widely used to study the structure and dynamics of macromolecules, such as proteins or nucleic acids.1 Sirtuins are a nicotinamide adenine dinucleotide (NAD1)-dependent histone deacetylase with highly conserved from bacteria to mammals and have been shown to extend lifespan in yeast.2,3 In mammals, there are 7 members (SIRT1-SIRT7) in the sirtuin family, which belong to class III histone deacetylases (HDACs) and show different functions, structure, and localization.4 Sirtuin1 (SIRT1), one of seven sirtuins members, has been extensively studied and evidently would be implicated in metabolism, senescence, and cancer.5 The NAD+-dependent deacetylase, has been described in the literature as a major player in the regulation of cellular stress responses. Its expression has been shown to be altered in cancer cells, and it targets both histone and non-histone proteins for deacylization and thereby alters metabolic programs in response to diverse physiological stress. Interestingly, many of the metabolic pathways that are influenced by SIRT1 are also altered in tumor development. Not only does SIRT1 have the potential to regulate oncogenic factors, it also orchestrates many aspects of metabolism and lipid regulation and recent reports are beginning to connect these areas. SIRT1 influences pathways that provide an alternative means of deriving energy (such as fatty acid oxidation and gluconeogenesis).6 In addition, SIRT1 also involved in some neurodegenerative diseases, such as Huntington’s and Alzheimer diseases.7,8 SIRT1 characterized by a conserved

MATERIALS AND METHODS
Materials
SIRT1 is one of important drug target in drug discovery field. Three-dimensional (3D) crystal structures for SIRT1 protein was obtained from the Protein Data Bank, PDB ID: 4I5I9 and PDB ID: 4ZZI.10

Methods
Molecular dynamics simulations using the AMBER 1211 and ff99SB force fields. Ligand parameters obtained from the general AMBER force field GAFF.12 Preparation of protein-ligand complexes, in addition to the counter ion, solvation, and preparation parameter/topology and coordinate files made using tLEaP module in AMBER. Four hits compound SIRT1 selected to proceed to the molecular dynamics simulations. Positive control compound selected in MD simulation is a ligand co-crystal and salermide. Partial atomic charge is calculated by the AM1-BCC method that uses Antechamber software. The system was solvated using the water model TIP3BOX.13 Then all of the complex dissolved in water (using a water model TIP3BOX) and cut boundary conditions octahedral periodic been applied (a cutoff distance of 10 Å). The entire systems were neutralized by adding Na+ counter ions by replacing solvent molecules. The systems were subjected to 10,000 steps steepest descent energy minimization. The production run was 10 ns. All simulations were run under periodic boundary conditions at the temperature (300 K) and the pressure constant (1 bar). Analysis of molecular dynamics simulations covers the RMSD (Root mean square deviation), RMSF
RESULTS

On the RMSD simulations, which lasted for 950 frames, each system has increased RMSD backbone indicating that the macromolecules unfold. There is an increase of four inhibitor compounds varied after frame 375. Ligand co-crystal 415 and hits compound shows RMSD value until a stable frame 950. 5-oxocoronaridine and dregamine increased to 4 Å as well as salermide.

The value of RMSF outline illustrates the shift in the conformation of amino acid residues each that provide the flexibility of the protein. On RMSF against macromolecules PDB ID: 4I5I seen does not indicate high fluctuating during the simulation took except on the last residue number. This indicates the residue is stable and does not provide the flexibility. Important residues such as Ala262, Phe273, Phe297, Ile314, Gln345, Asn346, Ile347, Asp348, and Val412 showed a low value of RMSF. This indicates that the residue is stable to bind to macromolecules and lacks flexibility (Figure 1). On RMSF backbone atoms against macromolecules PDB ID: 4ZZI at residue numbers 183-230 indicate fluctuating and high flexibility as a domain allosteric non-bonded ligand inhibitor compound (Figure 2).

Based on the docking scores and MMGBSA/MMPBSA binding free energies in Table 1, of the four compounds derived inhibitor hits compound 5-oxocoronaridine has the lowest free energy while salermide has the highest free energy between all four of these compounds. Meanwhile on the macromolecule PDB ID: 4ZZI dregamin hits compound has the lowest free energy while salermide has the highest free energy of the four of these compounds (Table 2).

Ligand co-crystal 415 have hydrogen bonding and binding affinity was good at residue Ile347 (94.21%), Asp348 (66.53%), Asn346 (63.16%) and Gln345 (95.89%). 5-oxocoronaridine showed hydrogen binding interactions and binding affinity was good in Ile347 and Phe297 with occupancy respectively 87.37% and 50%. Dregamin showed hydrogen bonding interactions and binding affinity on Ile316 with occupancy 73.05%. The active compound in SIRT1, salermide have hydrogen bonding and good affinity at Glu315 with occupancy 69.89% (Table 3).

On the PDB ID: 4ZZI co-crystal 1NS form hydrogen bonds with residues Phe273, Asn346, Ile347, Asp348 and Val412 with value occupancy respectively 71.05%; 54.32%; 99.89%; 96.11%; 77.68%. 5-oxocoronaridine has hydrogen bonding and binding affinity was good at residue Ile347 (95.89%), Asp348 (66.53%) and Gln345 (95.89%). Dregamin showed hydrogen bonding interactions and binding affinity on Ile316 with occupancy 73.05%.

![Figure 1: RMSD backbone atoms on ligands simulation with SIRT1 Macromolecules PDB ID: 4I5I and 4ZZI.](image1.png)

![Figure 2: RMSF backbone atoms on ligands simulation with SIRT1 Macromolecules PDB ID: 4I5I and 4ZZI.](image2.png)

<table>
<thead>
<tr>
<th>Table 1: Value Free Energy MMGBSA, MMPBSA and docking scores ligand / hits against SIRT1 macromolecule PDB ID: 4I5I</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compounds</td>
</tr>
<tr>
<td>--------------------------------</td>
</tr>
<tr>
<td>Ligand co-crystal 415</td>
</tr>
<tr>
<td>Dregamine</td>
</tr>
<tr>
<td>5-oxocoronaridine</td>
</tr>
<tr>
<td>salermide</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 2: Value Free Energy MMGBSA, MMPBSA and docking scores ligand / hits against SIRT1 macromolecule PDB ID: 4ZZI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compounds</td>
</tr>
<tr>
<td>--------------------------------</td>
</tr>
<tr>
<td>Ligand co-crystal 1NS</td>
</tr>
<tr>
<td>Dregamine</td>
</tr>
<tr>
<td>5-oxocoronaridine</td>
</tr>
<tr>
<td>salermide</td>
</tr>
</tbody>
</table>
Analogs, sirtinol, tenovin, salermide, thiobarbiturate, cambinol, 2-anilinobenzamide and other compounds. Simulation of molecular dynamics carried out by Karaman and Sippl (2015) by calculating MMGBSA with thieno core molecules [3,2-d] pyrimidine-6-carboxamide to SIRT1 shows all hydrogen bond interactions between the thieno [3,2-d] pyrimidine-6-carboxamide and residue Ile347, Asp348 as well as water molecules required for inhibitor activity. The region amino acids residue Thr344 – Asp348 referred to as “floor” of the binding pocket possibility ligand binding for interaction site. Ile347 and Ile348 were extensively residue domain to binding pocket for the ligand. The results of the hydrogen bond analysis also show that residual Ile347 and Asp348 have high occupancy that plays a role in inhibitor activity. Based on research results the 5-oxocoronaridine and dregamine compounds are potential candidates for SIRT1 inhibitors.

**CONCLUSION**

Hits compounds dregamine and 5-oxocoronaridine against two of macromolecules SIRT1 inhibitor (PDB ID: 4ISI and 4ZZI) obtained free dere tense hydrogen bonds with residues Ala262, Gln345 and Asn346 with value occupancy respectively 73.58%; 95.16%; 68.84%. The residue Asn346, Ile347, Asp348 interacts with the hit compound dregamine with value occupancy respectively 82.42%; 99.89%; 93.16%. As for the active compound salermide form hydrogen bonds with residues Val412 with occupancy 63.68% (Table 4).

**DISCUSSION**

Extensive studies of sirtuins began in early 2000 when Guarente and coworkers found that Sir2 prolongs the life of yeast and lower organisms such as worms. Although well documented to prolong the lifespan of yeast, worms, flies, and mammals, SIRT1 has recently been involved in the initiation and progression of many dangerous diseases such as cancer. Sirtuin inhibitors may also be useful as therapeutic agents because of SIRT1 regulation of cancer. This raises the possibility of inhibition of SIRT1 can suppress the proliferation of cancer cells. Several small molecule acts as sirtuin inhibitor that has potentially useful as therapeutic agent and has been characterized, including splitomicin and its analogs, sirtinol, tenovin, salermide, thiobarbiturate, cambinol, 2-anilinobenzamide and other compounds. Simulation of molecular dynamics carried out by Karaman and Sippl (2015) by calculating MMGBSA with thieno core molecules [3,2-d] pyrimidine-6-carboxamide to SIRT1 shows all hydrogen bond interactions between the thieno [3,2-d] pyrimidine-6-carboxamide and residue Ile347, Asp348 as well as water molecules required for inhibitor activity. The region amino acids residue Thr344 – Asp348 referred to as “floor” of the binding pocket possibility ligand binding for interaction site. Ile347 and Ile348 were extensively residue domain to binding pocket for the ligand. The results of the hydrogen bond analysis also show that residual Ile347 and Asp348 have high occupancy that plays a role in inhibitor activity. Based on research results the 5-oxocoronaridine and dregamine compounds are potential candidates for SIRT1 inhibitors.
energy MMPBSA calculation about -23 kcal/mol meanwhile occupancy hydrogen bonding of residues Ile347 and Asp348 about 80%.

ACKNOWLEDGEMENT

Author (AY) thanks to the Publikasi Internasional Terindeks Untuk Tugas Akhir Mahasiswa (PITTA) 2016 grant from the Universitas Indonesia, for funding support.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

REFERENCES


Article History: Submission Date : 24-08-2017 ; Revised Date : 07-10-2017 ; Acceptance Date : 25-11-2017.
Cite this article: Andika, Erlina L, Azminah and Yanuar A. Molecular Dynamics Simulation of SIRT1 Inhibitor from Indonesian Herbal Database. J Young Pharm. 2018;10(1):3-6.