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Corresponding author: Neni Frimayanti, E-mail: nenifrimayanti@gmail.com Research article

Insight on the In Silico Study and Biological Activity Assay of Chalcone-Based 1, 5-Benzothiazepines as Potential Inhibitor for Breast Cancer MCF7

Neni Frimayanti^{1,*}, Marzieh Yaeghoobi², Ihsan Ikhtiarudin¹, Dhea Rizki Wannisyah Putri¹, Hamid Namavar² and Fatemeh S.Bitaraf³

Sekolah Tinggi Ilmu Farmasi (STIFAR) Riau, Jalan Kamboja, Pekanbaru, 28293, Indonesia
 School of Medicine, Shahroud University of Medical Sciences, Shahroud, Iran
 Department of Medical Biotechnology, School of Medicine, Shahroud University of Medical Sciences, Shahroud, Iran

Abstract In silico study was performed to twelve 1,5-benzothiazepine chalcone derivatives with the protein target from the crystallographic structure modeling of the enzyme tyrosine kinase. The objective of this study is to execute and to estimate the biological activity of chalcone-based 1,5-benzothiazepine derivatives as potential inhibitors for breast cancer MCF7. To get insight into potential anticancer activities, molecular docking, molecular dynamic and ADME prediction were performed. Docking results reported that compound MA9 with binding free energy of -11.2 kcal / mol can interact through hydrogen bonds with amino acids Cys788 on 1T46 protein active site. In addition, the lowest binding free energy conformation indicated its stability during molecular dynamic simulation. MA9 is also shown to have drug likeness properties based on ADME prediction. In order to evaluate the modeling outcomes, MTT assay were performed for some of the most and least promising benzologs (i.e., MA1, MA6, MA8 and MA9). As expected, compound MA9 with the best calculated anticancer properties revealed the best inhibition against MCF7cell line in vitro. Thus, this compound was chosen as the reference for the next stage in the drug design.

Keywords: ADME, Benzothiazepine, Docking, MCF7, Molecular Dynamic

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INTRODUCTION

Cancer is one of the leading causes of death worldwide. It is an abnormal growth of body tissue cells that turn into malignant cells that can grow further and spread to other parts of the body and may result death. Breast cancer is one type of cancer; which is very frightening for women throughout the world. According to The American Cancer Society, in the incidence of invasive breast cancer in 2018 occurred as many as 266.120 cases in women, 2.550 cases in men, 63.960 cases of carsinoma in situ (CIS) and 40.610 cases (14%) deaths due to breast cancer (Rebbeca et al., 2018).

Benzothiazepines are seven membered heterocyclic compounds, containing nitrogen and sulfur (Vyawahare et al., 2010) and 1, 5-benzotiazepine derivatives are important pharmacophores from this family (Kurokawa et al., 1997). According to IUPAC (International Union of Pure and Applied Chemistry), pharmacophore is a steric and electronic factor needed to ensure the optimal molecular interactions with specific biological target structures (Mannhold et al., 2006; Narendran et. al., 2014). 1, 5-Benzothiazepine is a pharmacophore that can provide a broad spectrum of biological activity such as Ca²⁺ channel antagonists (Kurokawa et al., 1997), antimicrobials (Gaikwad et al., 2013), lung cancer (Ameta et al., 2013) and anticonvulsants (Garg et al., 2010). Rationally, in silico studies are playing important role in drug design (Neni et al., 2020). In the process of drug design, in silico studies (i.e. molecular docking and molecular dynamic) are used to find the best conformation with low binding free energy. In addition, they can provide useful information for predicting the orientation of a candidate drugs bond to its target (i.e. protein). To date, although some in silico investigations have been carried out on potential anticancer properties of chalcone derivatives like pyrazolines (Jasril et al., 2017) and guercetins (Lelita et al., 2017) but unfortunately, thus far, chalcone-derived heterocycles like 1,5-benzothiazepines have not been extensively studied for their potential anticancer activities. Moreover, there are not many reports on discovering an anticancer drugs using in silico tools such as molecular docking, molecular dynamic (MD) and ADME calculation. This study aimed to explore new potential agents from 1, 5-benzothiazepine against breast cancer MCF7 using tyrosine kinase as the target.

MATERIALS AND METHODS

Molecular docking

All 1, 5-benzothizepines were synthesized, purified and characterized following reported procedure (Yaeghoobi, 2012). The molecular structure of these benzologs and Doxorubicin as positive control were drawn using Chemdraw Ultra 13.0 and stored in "cdx" format and converted to PDB format using Discovery studio visualizer (DSV) software package. The energies of all the ligands were minimized for then the minimized structures were subsequently prepared with detected root of torsion and number of torsion. Molecular structure of twelve 1, 5-benzothiazepine and doxorubicin are shown in Table 1.



 Table 1. Molecular structures of 1, 5-benzothiazepine

The structure of protein was downloaded from the protein databank (www.pdb.org PDB ID: 1T46), then the water molecules and initial ligands in the protein were removed. Further preparation carried out by adding hydrogen atoms, polar only and merged non polar. A grid box of the protein structure was then achieved with a grid spacing of 1 Å, dimensions of $31 \times 25 \times 35$ points along the x, y and z-axes and centered on the protein active site. Docking was performed using Autodock vina software packages (Trott et al., 2010).

Molecular dynamic simulation

NAMD (NAnoscale Molecular Dynamics program v 2.9) was excreted to execute the molecular dynamic simulation (i.e. preliminary study). The selected force field of CHARMM27 (Chemistry at HARvard Macromolecular Mechanics) was utilized as the best force field. TIP3P water box with 2.5 Å water layer for each direction of coordinated structure was used to achieve modeled protein.

NVT ensemble from 0 to 300 K over 100 ps was applied to heat the system gradually. MD simulations was conducted on 50 ns time scale for each system in an isothermal isobaric ensemble (NPT) with periodic boundary conditions. The coupling of temperature and pressure parameters were set on 1.0 ps. The coordinates were saved at every 0.1 ps during the sampling process. The simulations were generated the conformations for then they were used for further binding free energy calculations and decomposition process.

Adsorption, distribution, metabolism and excretion (ADME) prediction

ADME profiles of potential compounds were calculated to get better insight into the physicochemical and pharmacokinetic properties as well as prediction of drug likeness of candidates. In this study, the ADME profiles were calculated using SwissADME server (http://www.swissadme.ch/index.php).

Biological assay using MTT assay

To evaluate the modeling results, MTT assay were performed for compounds with the most and least MCF7 cells were seeded into 96-well plates at an initial density of 104 cells/well. MA1, MA6, MA8, MA9 and Doxorubicin were then added as standard drug to achieve a final concentration of 100, 50, 25, 10, 5, 2.5, or 1.25 µgr/ml. MTT(Methyl Thiazole Tetrazolium) was dissolved in PBS to obtain the concentration of 5 mg/mL, then it filter sterilized and added in a volume of 10 µL to each well to attain the final concentration of 0.5 mg/mL and incubated for 4 h. The MTT-containing medium was replaced by 100 µL of dimethylsulfoxide (DMSO, SantaCruz, Dallas, TX, USA), and the plates were gently agitated for 10 min. Absorbance was obtained at 570 nm with background subtraction at 690 nm using Biotek Cytation 5imaging reader (Biotek Instruments, Winooski, VT, USA). All of the experiments were performed in triplicate.

RESULTS

Molecular docking

Molecular docking is a good tool to predict and match the desired binding sites, understanding possible conformation of the compounds and further to clarify the binding interactions between ligand and receptor.

In this study, twelve 1,5-benzothiazepines were docked as ligands into protein (PDB ID: 1t46) to elucidate the ligand interactions with protein binding site. The docking results are listed in Table 2 which highlighted the tendency of these chalcone-based benzologs to bind tightly to the active site as well as allosteric sites of tyrosine kinase as receptor. The best poses of complex ligand-protein were selected according to the lowest value of binding free energy and also Root Mean Squared Deviation (RMSD) value. Based on the docking results, **MA9** estimated to be active as breast cancer MCF7 inhibitor. Although **MA5** has also displayed a hydrogen bond with Cys788 residue, yet,

its binding free energy value is higher than the positive control and it cannot be considered as a potentially active ligand. In addition, compounds **MA1** and **MA6** have shown the binding free energy value of -9.3kcal/mol and -9.1kcal/mol, respectively. However, **MA1** has not exhibited any hydrogen bonding or other interaction with amino acid residue. Although **MA6** has displayed hydrogen bonds with Asp810 and Glu 640, nevertheless, these bindings are different from the amino acid sequence of the positive control. Hence, these compounds can not be considered as potential inhibitors for MCF7.

	Parameters				
Compound	Binding free energy (kcal/mol)	RMSD value	Hydrogen bond		
MA1	-9.3	0.000	-		
MA2	-9.4	0.000	-		
MA3	-10.2	0.000	-		
MA4	-9.9	0.000	-		
MA5	-9.3	0.000	Cys788		
MA6	-9.1	0.000	Ásp810, Glu640		
MA7	-9.7	0.000	-		
MA8	-10.1	0.000	Cys788		
MA9	-11.2	0.000	Cys788		
MA10	-10.0	0.000	-		
MA11	-9.8	0.000	-		
MA12	-10.0	0.000	-		
Doxorubicin	-9.9	0.000	Asp677, Tyr672		

 Table 2. The docking results.

Compound **MA8** with binding free energy value of -10.3kcal/mol, was observed to have one hydrogen bonding with amino acid residue Cys788, van der Waals interaction with Ser639 Glu640, Leu644, Tyr570, Ile571 and Ile789. The spatial arrangement of MA8 is presented in Figure 1.



Figure 1. Spatial arrangement of compound MA8

The estimated active compound, **MA9**, with binding free energy value of -11.4kcal/mol, interacted via hydrogen bonding with Cys788. Furthermore, **MA9** is exhibited to interact through van der Waals interaction with amino acid Glu640, Ser639, Leu647, Leu783, Ile571 dan Tyr570. Thus, **MA8** and **MA9** predicted to be active compounds. The existence of methoxy groups in ring A may enhance the hydrophobicity of these molecules, which may presumably cause compounds **MA8** and **MA9** to have better binding affinity compare to other compounds (Noval et al., 2020). Figure 2 Superimpose was performed to check the orientation of **MA8** and **MA9** ligands simultaneously. It seems that both of them have the same orientation to bind with the protein where in both cases, the seven-membered ring flipped slightly. However, other parts of chalcone backbones (i.e. rings A and B) maintained their orientations. Superimpose for **MA8** and **MA9** shown in Figure 3.



Figure 2. Spatial arrangement of compound MA9.



Figure 3. Superimpose for MA8 (pink) and MA9 (purple).

Based on the docking results, it was found that doxorubicin has binding free energy value of -9.9kcal/mol and an RMSD value of 0,000. Docking visualization results shown that Doxorubicin (i.e. positive control) can bind to 13 amino acid residues on the active site of the receptor, namely Glu671, Ala621, Leu799, Val603, Leu595, Phe811, Asp677, Asn680, Cys674, Tyr657, Gly676, Cys673 and Cys788. It interacts with receptors through hydrogen bonds with amino acid residues Asp677 and Tyr672, van der Waals interaction with residues Cys673, Gly676, Tyr675, Asn689, Cys788 and Phe811, pi-alkyl bonds with residues Ala621, Leu799 Leu595 and Val603, pi-sigma

bonds with Leu 799 and Leu595 residues, and pi-donor hydrogen bonds with Asp677 residues. Figure 4 is depicted the spatial arrangement for Doxorubicin.

According to the result, the binding free energies for **MA8** and **MA9** are lower than Doxorubicin and moreover, they both exhibited crucial binding affinities (i.e. hydrogen bond) with residue Cys788 at the protein active site which seems to be significant for activity prediction. Hence, **MA8** and **MA9** considered potentially active as MCF7 inhibitors and can be further utilized in drug designing process.



Figure 4. Spatial arrangement of doxorubicin.

Molecular dynamic

MD simulation was inspected on these two compounds to explore the interactions between ligand and receptor (Adel, et. al 2019). The stability of MD simulation was examined to confirm the binding profile of ligands and to give an overall impression about the estimated active 1,5-benzothiazepine, MD simulation was subjected to run for 50 ns. In the present study, MD simulation was performed to ensure that interaction between protein and active compounds are still maintained (Neni et al., 2020). It was initiated using the high stability with energy minimum at temperature of 300K to see affinity of the ligand to binding site.

The results were inspected by examining the efficiency of hydrogen binding in both potentially active compounds i.e. **MA8** and **MA9** and also the most inactive compounds i.e. **MA1** and **MA6** before and after 50 ns and 300K MD simulation. Generally, in case of compound **MA9**, the conformation was maintained to bind well with the same residues before and after operation and the hydrogen bond distance is less than 2.9 Å. However, compound **MA8** has hydrogen bond distance higher than 2.9 Å and the rest of tested benzologs seemed to lose their activity due to the presence of some interactions between ligands and receptor that were not maintained. According to the docking results, **MA8** has potency as breast cancer inhibitor, unfortunaely according to MD simulation this compound was not able to maintain the existence of hydrogen bond and the distance of hydrogen bonding is higher than 2.9 Å (Duan et. al., 2019). Table 3 is listed the interactions with amino acid after the MD simulation. Based on MD simulation, **MA9** seemed to retain its interactions with the same amino acids after simulation which indicates that it can be used as potential inhibitors for breast cancer. Visualization of MD simulation is depicted in Figure 5.

Compound	After docking	MD simulation	Distance of Hbond
MA1	-	-	-
MA6	Glu 640, Asp810	-	-
MA8	Cys788, Ser639, Glu640, Leu644, Tyr570, Ile571, Ile789, Asp810, Val643, Pro573, Ala636.	Ala636, Pro573, Tyr570, Cys788, Asp810, Ile571, Glu640, Leu644,	3.2 Å
MA9	Cys788, Glu640, Ser639, Leu647, Leu783, Ile571 , Tyr570, Asp810 , Val643, Cys788, Ala636, Pro573 Leu644	Ala636, Pro573, Tyr570, Cys788, Asp810, Ile571, Glu640, Leu644, Ser639	2.9 Å

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ADME profiles calculation

The in silico absorption, distribution, metabolism and excretion (ADME) profile is also called as pharmacokinetic profile. Hence, ADME was used to expect reduce the risk of late-stage attrition of drug development and to optimize the screening process. However, it only can be used for the promising compounds (Yamashita et al., 2004).

A compound can be determined to have bioavailability if it follows the Lipinski's rule of five. This rule consisted when the maximum MW is 500, Log P is not greater than 5, hydrogen bond donor is less than 5 and hydrogen bond acceptor is less than 10 (Lipinski et al., 2012).Lipinski's rule of five calculation was performed to achieve the absorption levels of potentially active compounds to across the lipid bilayers in the human body. ADME profile was generated the drug likeness properties based on Lipinski's rule of five for compounds **MA1, MA6, MA8** and **MA9** (Table 4). Based on calculations, the potentially active compounds shown the reasonable drug properties. It is indicated that **MA9** is safe to be used as potential anti cancers for MCF7.



(a)

(b)

Figure 5. MD Visualization for (a) MA8 and (b) MA9

Profiles	Compound MA1	Compound MA6	Compound MA8	Compound MA9
MW (g/mol)	330.47	380.91	421.51	395.52
Consensus Log Po/w	3.79	3.91	3.44	3.95
Hydrogen bond donor	1	2	1	0
Hydrogen bond acceptor	0	1	5	2
Rotatable bonds	2	2	5	3
Drug likeness (Lipinski)	Yes	Yes	Yes	Yes

 Table 4. Drug likeness properties of MA8 and MA9.

Bological assay

The effect of **MA1**, **MA6**, **MA8**, **MA9** and Doxorubicin at various concentrations (100, 50, 25, 10, 5, 2.5, or 1.25 µgr/ml) on the MCF7 cell viability were determined using MTT assay. MCF7 cells incubated for 72 h. The IC₅₀ value for **MA9**, **MA8**, **MA1** and **MA6** were obtained at the concentrations about 550 and 100µgr/ml respectively, whereas the IC₅₀ value for Doxorubicin was observed at about 10µgr/ml. Furthermore, at 5µgr/ml, the cell survival assessed value for **MA9** was significantly lower than the rest of tested compounds. The result revealed that amongst all tested compounds, **MA1**, **MA6** and **MA8** were confirmed to be inactive while **MA9** with the lowest concentration for IC₅₀, exhibited the higher chance of cell survival. Hence, **MA9** with the best experimental result can be considered as a potential lead for breast cancer treatment. The MTT assay results summarized in Figure 6 (a and b)



Figure 6. Percentage of cell survival for tested compounds and Doxorubicin in various concentrations (a) and (b)



(b)

Figure 6. Percentage of cell survival for tested compounds and Doxorubicin in various concentrations (a) and (b)

CONCLUSION

In silico study has been successfully estimated the activity of twelve chalcone-based 1,5-benzotiazepines as potential inhibitors for breast cancer MCF7. Based on the in silico results, **MA9** with binding free energy of -11.2 kcal, can interact through hydrogen bonds with amino acids Cys788 on the active site of 1T46 protein. This potentially active candidate exhibited lowest binding free energy conformation which indicated its stability during the molecular dynamic simulation. ADME profiles also proofed that its drug likeness properties. The computational results were accordance with the biological activity assay. Hence, compound MA9 was then chosen as the reference for the next stage in the drug design.

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