Research Article

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Molecular Docking Analysis on Epidermal Growth Factor Receptor Wild Type (EGFR^{wt}) with Quinazoline Derivative Compounds as Tyrosine Kinase Inhibitors

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Abstract

Molecular docking analysis for protein EGFR^{wt} with quinazoline derivatives had been carried out. Six quinazoline derivatives obtained from previous experiment and two compounds predicted by QSAR were docked into EGFR^{wt} using AutoDock program. Comparing between six compounds and erlotinib, these compounds have lower binding energies of -6.54, -6.48, -6.22, -6.24, -6.11 and -6.09, respectively than erlotinib (-4.84 kcal/mol). Docking result of two compounds resulted from QSAR exhibited lower binding energies of -4.85 and -5.96 kcal/mol than erlotinib. Binding pose from those compounds took place in amino acid residue Met769 in distance range of hydrogen bond 1.7 until 2.1 Å.

Keywords: Molecular docking, EGFR^{wt} protein, AutoDock, Quinazoline

1 Introduction

Since 1985, lung cancer had been the most leading cancer in the world. There was 17.6% people death in the world caused by lung cancer and approximately 49.9% of this cases happened in developing countries [1]. In 2008, lung cancer became the main cause of cancer death in men with 951.000 deaths [2]. Nowadays, inhibition of cancer growth focused on protein receptor which had an important role in inhibition mechanism, such as EGFR protein [3], [4]. EGFR protein is a member of erbB1 family. This protein could influence the tumor growth including proliferation, angiogenesis, metastasis, and inhibition in apoptosis process [5].

One of strategies to inhibit EGFR protein

was by used small molecules tyrosine kinase inhibitors. This type inhibitor bind an H-bond to ATP binding site, thereby blocking signal transduction from the EGFR [6]. Some quinazoline derivative compounds had been used as tyrosine kinase inhibitor, such erlotinib [7] as the first generation of EGFR inhibitiors, afanitib or BIBW2992 [8] the second generation, and Osimertinib as the third generation of EGFR inhibitiors [9].

The Structure of co-crystal of erlotinib [10] with EGFR protein contribute an important information in search of the bioactive compound. From this co-crystal, binding pose of inhibitor and protein could be studied and utilized to design a new bioactive compound which have the same key interaction. Afatinib as analogue compounds which

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H. Rasyid et al., "Molecular Docking Analysis on Epidermal Growth Factor Receptor wild type (EGFRwt) with Quinazoline Derivative Compounds as Tyrosine Kinase Inhibitors."



Figure 1: Design strategy of novel EGFR inhibitors [13].

have similar interaction in EGFR protein, due to the same binding mode on amino acid residue Met793 and Cys797 [11]. The compound, PD153035, didn't show any side effect until the doses up to 2.5 μ M [12]. Modification through cyclization in compound PD153035 at the position 5- and 6- of quinazoline ring (Figure 1) had been done and complete a novel series of EGFR inhibitors [13].

Due to the novel series of EGFR inhibitor, QSAR analysis had been carry out and successfully design two derivatives compound that had better activities which could be described in a lower IC_{50} value [14]. However, docking analysis of these two predictive compounds had not been analysis. This research will focus on looking for binding site, binding pose, and inhibition mechanism of some quinazoline derivatives compounds.

Nowadays, computational chemistry has been considered as an effective way to help in understanding the interaction of drug and protein [15], [16]. Autodock program could help to predict and rank the structures arising from the association between ligand and a target protein in 3D structure [17].

2 Computational Methods

Three dimensional structure of EGFR protein was taken from Protein Data Bank (PDB ID: 1M17). Erlotinib as a ligand standard was extracted from the PDB file. Some of quinazoline derivative compounds were taken from previous study for compound B1, B2, B3, B4, B5, and B6 [13] to know pose of binding site and also two compound (Q1 and Q2) resulted from QSAR analysis [14]. List of compounds can be seen in Table 1. Software packaging used in this study were AutoDock 4.0 software with the help of AutoDock Tools, Pymol, Chimera, Gaussian 09 [18].

Preparation of protein molecule. Complex protein-ligand in the PDB file was cleaned from all

 Table 1: Core structure and list of docked compounds



of residue such as water and ligand erlotinib. Then, all of the sequence protein was added with hydrogen and charge using dockprep analysis and saved in pdb format file.

Preparation of ligand standard. Erlotinib as standard ligand was taken from the PDB code 1M17. Ligand erlotinib was selected and choosed inverted

all molecule. Then, delete all of selected molecule. All of hydrogen and charge were added to the ligand and saved in pdb format file.

Redocking analysis. Protein EGFR and erlotinib docked in affinity grid maps size was $30 \times 30 \times 30 \text{ Å}$ with 1.000 Å of spacing. Lamarckian Genetic Algorithm (LGA) was used to search the lowest energy for each docking conformation [19]. Resulted 10 conformations for each automated docking was analysed to know free energy binding, inhibition constant, and pose of binding site. Successful redocking result could be seen in lower RMSD, usually lower than 2 Å [20].

Preparation of ligand and docking analysis. Eight of quinazoline derivative compounds was drew and optimized in Gaussian using DFT (B3LYP/6-31G). All of compounds was docked in binding site which known in redocking analysis. The affinity grid maps size and LGA was set up same with redocking.

3 Results and Discussions

3.1 Redocking analysis

Standard ligand (erlotinib) was docked into EGFR protein to know the binding site of protein. Coordinate of resulted binding site was 22.0732; 0.288; 52.7863. Table 2 served the result of dockingsimulation which produced 10 conformations. There are seven conformations which have RMSD value lower than 2 Å. Successful redocking analysis will be seen from RMSD value and the matching pattern in hydrogen bond interaction



Figure 2: The overlapping structure of erlonotib before docking (green) and conformation 7 after redocking (pink).

compared with experimental result. Conformation 7 exhibited the lowest RMSD value (1.31 Å) between all of the cluster rank and showed an agreement in hydrogen bond interaction, due to the matching structure orientation between both of them. Indicating this conformation was in stable condition when binding with protein receptor. Overlapping structure of erlotinib before docking and structure of conformation 7 served in Figure 2. In this step, we can know that binding pose of erlotinib to the EGFR protein was influenced with amino acid residue of Met769. This result met with previous result which said that inhibitor interaction of erlotinib due to erlotinib accepts an H-bond from Met769 amide nitrogen [10].

Table 2: Redocking result of EGFR protein with erlotinib

| - | - | | | |
|--------------|-----------------------|--------------|------------------|----------|
| Conformation | ΔG (kcal/mol) | $K_i(\mu M)$ | Hydrogen Bond | RMSD (Å) |
| 1 | -4.94 | 240.14 | Cys773 (1.915 Å) | 1.80 |
| 2 | -4.83 | 290.01 | Met769 (2.115 Å) | 1.58 |
| 3 | -4.75 | 309.68 | - | 1.55 |
| 4 | -4.99 | 220 | Cys773 (1.926 Å) | 1.74 |
| 5 | -3.87 | 146 | Met769 (2.08 Å) | 2.93 |
| 6 | -4.88 | 266 | - | 1.37 |
| 7 | -4.84 | 282.66 | Met769 (2.117 Å) | 1.31 |
| 8 | -4.93 | 243.01 | Cys773 (2.164 Å) | 1.63 |
| 9 | -3.96 | 125 | Met769 (2.12Å) | 3.03 |
| 10 | -3.63 | 217 | - | 4.01 |

H. Rasyid et al., "Molecular Docking Analysis on Epidermal Growth Factor Receptor wild type (EGFRwt) with Quinazoline Derivative Compounds as Tyrosine Kinase Inhibitors."



Figure 3: Conformation of binding pose for compound B1-B6 (a:B1, b:B2, c:B3, d:B4, e:B5, f:B6, g:Q1, h:Q2).

3.2 Docking analysis for all compounds

Molecular docking of B1–B6 and Q1–Q2 were carried out using the resulted binding site from redocking analysis. All of compounds were set up to have the same position with erlotinib. Figure 3 displayed the best conformation of binding pose for each compound. Binding pose of compound B1 showed the same binding pattern with erlotinib in amino acid residue of Met769 in distance of hydrogen bond was 1.867 Å. This result appropriate with previous study [21] which docked compound B1 using GOLD and obtained only one hydrogen bond in Met769 in distance of bond was 2.14 Å. Compound B1also execute using AutoDock package and found two hydrogen bond directly through compound between N atom of quinazoline with Met769 and fluoro atom with Thr766 [13].

The interaction of B2–B6 and Q1–Q2 compounds was located in amino acid residue of Met769. This result indicating that these derivative compounds have a correct binding site, due to the same amino acid interaction in redocking analysis. This result was in good agreement with SAR analysis which could be seen in N11 atom of quinazoline with Met769 [6].

Autodock4 could convert binding energy to the inhibition constant by using equation (1) below.

$$K_i = \exp\left(\frac{\Delta G}{(R \ge T)}\right) \tag{1}$$

Where ΔG is the docking energy, *R* in calorie is 1.98719, and *T* is 298.15 K [20].

Table 3 displayed the docking result for each analogues compound. There were 10 conformations cluster rank for each docked compound. Compounds with the best conformation will have the lowest energy binding. The best conformation rank was taken from the lowest energy and lowest inhibition constant for each compound. Comparing with erlotinib (–4.84 kcal/mol), all of these quinazoline derivative compounds had lower binding energy value and inhibition constant, indicating these analogues compound had better anticancer activities. Inhibition constant of all compounds are also lower than erlotinib (282.6 μ M). This result suggested that these all compounds have a better stability and stronger interaction with protein receptor.

| Conformation of Compound | Binding Energy | K _i (μM) | Hydrogen Bond |
|-----------------------------|-------------------|---------------------|------------------|
| r r r | (kcal/mol) | | |
| B1 | -6.54 | 16.06 | Met769 (1.867 Å) |
| B2 | -6.48 | 17.86 | Met769 (1.981 Å) |
| B3 | -6.22 | 27.74 | Met769 (2.196 Å) |
| B4 | -6.24 | 26.74 | Met769 (2.173 Å) |
| В5 | -6.11 | 33.4 | Met769 (1.957 Å) |
| B6 | -6.09 | 34.42 | Met769 (2.120 Å) |
| Q1 | -4.85 | 276.56 | Met769 (1.898 Å) |
| | | | Lys721 (1.768 Å) |
| Q2 | -5.96 | 42.52 | Met769 (1.807 Å) |

Table 3: Docking result of all compounds

4 Conclusions

Redocking analysis using compound erlotinib as standar ligand had been done and resulted conformation 7 as the best conformation, due to the lowest RMSD value and the matching binding pose. Docking analysis in protein EGFR^{wt} with compound B1–B6 and Q1–Q2 gave a lower binding energy than erlotinib. Inhibition constant of these compounds are also lower than erlotinib. This result indicated that all of compounds have a stronger interaction in protein EGFR^{wt}. The binding pose of all compound took place in amino acid residue Met769 with a distance of hydrogen bond in range 1.7 until 2.1 Å.

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