IN SILICO MOLECULAR DOCKING OF XANTHONE DERIVATIVES AS CYCLOOXYGENASE-2 INHIBITOR AGENTS

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ABSTRACT

Objective: To demonstrate the potential of different xanthone derivatives as cyclooxygenase-2 (COX-2) inhibitor agents and their selectivity against cyclooxygenase-1 (COX-1) and COX-2 using molecular simulation.

Methods: Nine novel xanthone derivatives (compounds A-I) were employed to dock against protein COX-2 (Protein Data Bank/PDB ID: 1CX2) and COX-1 (PDB ID: 3N8Z). Celecoxib, a selective COX-2 inhibitor, was chosen as a control compound. The free binding energy produced by the docking was scored using Protein-Ligand Ant System (PLANTS) and the hydrogen bonds (H-bonds) between ligands and enzymes were visualised using PyMol.

Results: Molecular docking studies revealed that celecoxib docked to the active site of COX-2 enzyme, but not to COX-1; whereas xanthone derivatives docked to the active site of both COX-2 and COX-1. Free binding energy of xanthone derivatives ranged between -73.06 to -79.25 against COX-2 and -78.13 against celecoxib. H-bonds in the molecule of xanthone derivatives and derivatives docked to the active site of both COX-2 and COX-1. The free binding energy of xanthone derivatives ranged between -73.57 to -79.18 and -78.13 against celecoxib. These results suggested that the inhibition was non-selective.

Conclusion: In conclusion, in silico studies showed that xanthone derivatives could be effective as potential inhibitors against COX-2, although they are not selective.

Keywords: Xanthones, Molecular docking, Anticancer, COX-2, Selectivity

INTRODUCTION

The cyclooxygenase (COX) enzyme plays an important role in the production of prostaglandin from arachidonic acid, which is involved in various processes in the body, including inflammation, pain, and hyperpyrexia [1]. It is widely known that COX has two isoforms, namely COX-1 and COX-2. COX-1 is a constitutive part of the body that maintains the normal function of the gastrointestinal organs, the kidneys, and platelets, while COX-2 is an inducible enzyme that is primarily expressed by various pro-inflammatory cytokines, lipopolysaccharides, mitogens, and oncogenes [2]. When COX-1 is inhibited, the inflammation process will decrease. Impairments in gastrointestinal mucosal protection as well as the normal functions of the kidneys and platelets, along with other side effects, may occur [1].

Clinical data shows that, although COX-2 plays a role in the repairment of injury, excessive expression of this enzyme will promote various pathological processes, including carcinogenesis and cancer growth [3]. COX-2 enzymes are found to be highly expressed in a number of inflammatory processes and tumours, such as inflammatory bowel disease (IBD) and colon cancer, whereas it is minimal or undetected at the normal colon cell [4]. This is why research on new anti-inflammatory drugs has been focusing on the identification of compounds with selective activity against COX-2 in preventing the inflammation process.

A number of studies show that the inflammation process is involved in carcinogenesis. The immune system, cytokines, chemokines, and transcription factors are directly associated with malignancy, which together form an “orchestra” in the pathogenesis of cancer. Immune cells in the microenvironment with inflammation stimulate the production of cytokines and chemokines. These will activate transcription factors responsible for spreading the tumor (mainly nuclear factor kappa B/NFkB and signal transducer and activator of transcription 3/STAT3) and stimulate other pro-inflammatory cytokines [5]. Therefore, inflammation can be considered as an initial form of tumor progression that may develop into a true cancer [6].

The use of aspirin and other non-steroidal anti-inflammatory drugs (NSAIDs) in chemoprevention of cancer has proven capable in reducing incidences of cancer and preventing mortality in some cancers, such as cancer of the colorectal [7], stomach [8], lung [9], breast [10], lymphatic tissue (Hodgkin lymphoma) [11], pancreas, oesophagus, kidney and bladder [12]. Aspirin and other NSAIDs act as anti-tumors by shifting the balance of Bax/Bcl-2 and activating a number of caspases [13], as antiplatelet drugs [14], by inhibiting the signalling process of NF-kB [16], as anti-angiogenesis drugs [17], and by inducing Rac1 gene involved in the apoptosis [18].

Selective COX-2 inhibitors were formerly developed as an anti-inflammatory drug with fewer gastrointestinal side effects when compared to COX-1 [19]. In long-term use, some COX-2 inhibitors (rofecoxib and valdecoxib) increase the risk of heart attacks and strokes; thus, they were withdrawn from the market. Currently, celecoxib is the only COX-2 selective inhibitor available in the market [19]. There is a rising urge to develop a new selective COX-2 inhibitor with fewer side effects through a more effective and efficient drug discovery process.

A method to investigate a more effective and efficient new drug is using molecular docking. This is a part of molecular modelling that predicts interaction orientations (conformations) between two molecules precisely, as shown by the formation of a stable complex. The most preferred orientation (the best conformer) is indicated by the lowest binding energy and is associated with the strongest interaction. This method allows us to explore and investigate many drugs for the same receptor at the same time. The drug with a better interaction between a ligand and a receptor will be chosen for use in laboratory experiments, and it saves resources and is less time-consuming [20].
This drug virtual-screening method is widely applied in structure-based drug design and can be performed for interaction modelling of ligands and proteins at an atomic level [21]. The chemical structure of certain drug receptor is downloaded from the Protein Data Bank (PDB) for investigation of their molecular docking capability at the active site, using computer algorithms and scoring function [22]. This consists of basic processes, namely ligand conformation predictions at the receptor active site, placement and orientation of the conformations, and their binding affinity [21]. Interactions between ligands and receptors, as indicated by their scoring function, are ranked; ligands with the lowest score will be investigated further. This step will make molecular docking an important part of more effective and efficient drug discovery, because it saves time and money [22].

Xanthone derivative is one compound known for analgesic and anti-inflammation activity. Natural xanthones can be isolated from a variety of plants, including Garcinia mangostana [23], Hypericum perforatum [24], Calophyllum inophyllum [25], Gamboge hanburyi [26], and Artocarpus optusus [27]. Xanthones have an anti-inflammatory effect by inhibiting COX-2 and prostaglandin synthesis at glia cell in rats [28], without affecting the constitutive COX-1 [29].

The docking of natural xanthones from Garcinia mangostana [30] and propoxy-and pentyloxy-substituted synthetic xanthones [31] are studied against COX-2 enzymes. The result showed that xanthone ligands were docked to the receptor and formed stable complexes through several interactions. The bond between drugs and receptors are categorised as weak and are formed through non-covalent interactions, such as van der Waals, hydrophobic, and hydrogen. This means the effects are reversible. A hydrogen bond (H-bond) has the strongest affinity with the distance below 3.2 Å [32] and is the main interaction contributing to binding energy.

A number of xanthone derivatives were newly synthesised by Yuanita (Organic Chemistry of Mathematics and Natural Sciences Faculty, Gadjah Mada University). These xanthones have not yet been studied for their COX-2 inhibitory activity. This study aims to investigate the molecular docking of hydroxy-and polyhydroxy-substituted xanthones against COX-1 and COX-2 receptors. This is to predict their inhibitory activities and the possibility of their selectivity.

MATERIALS AND METHODS

Material preparation

Three-dimension (3D) structures of enzymes were downloaded from the Protein Data Bank database (www.rosb.org) with PDB ID: 3NZF for COX-1 and 1CX2 for COX-2. 3NZF structure has a native ligand FLP (flurbiprofen), and 1CX2 is a complex of COX-2 enzymes with native ligand SC-558—a selective COX-2 inhibitor. A series of hydroxy-and polyhydroxy-substituted xanthones derivatives (compound A-I) were used as experimental ligands (fig 1).

The 3D structures of COX-1 and COX-2 that were downloaded were prepared with YASARA (http://www.yasara.org) in the standard setting. Only A-chain of the protein extracted from the PDB file and hydrogen atoms were included in the preparation process. The results were saved in the mol2 format and would be a virtual target for docking simulation. The downloaded native ligands (FLP and SC-558) were prepared with Marvin sketch by configuring them into two dimension (2D) format. They were protonated at pH 7.4, and ligand conformations were performed. The ten conformers form FLP and SC-558 ligands were saved in the mol2 format for the docking process.

The 2D structure of nine experimental ligands were constructed with Marvin sketch 5.2.5.1 (http://www.chemaxon.com) [33]. The preparation for these ligands was performed in the same way as the native ligands and the conformers saved in the mol 2 format for the docking process. Celecoxib (IUPAC name 4-[5-(4-methylphenyl)-3-(trifluoromethyl) pyrazol-1-yl] benzen and PubChem CID 2662) were used as a control for COX-2 selective inhibitors, and the ligand preparation was done in the same way as the others.

Docking molecular process

Docking program simulations used PLANTS/Protein-Ligand Ant System [34], and the docking parameter (root median square deviation/RMSD and free binding energy) were measured with YASARA (http://www.yasara.org). Root median square deviation for SC-558 was overlapping conformation of the copy ligand SC-558 after redocking against COX-2 active site, when compared to the ligand structure from measurements from the x-ray crystal. Also, RMSD for FLP was overlapping ligand FLP against the active site of COX-1. The smaller the RMSD value, the smaller the coordinate difference between two ligands, and the ligand structures that overlapped were more similar [35]. A docking program is valid and suitable for reproducing if the RMSD<2 Å [36]. In the PLANTS program, the binding site definition file is set to 5 Å from the coordinate location where SC-558 was bound to COX-2 and IBP was bound to COX-1. The position with the best

Fig. 1: Chemical structures of xanthone derivatives involved in the study
score (the most electronegative) was selected as the best predictive binding position [37].

The Pymol program (www.pymol.org) was used to see the H-bonds between ligand and the receptor [38]. Amino acid residues that interacted with the redocking compound were compared to those that interacted with the crystal molecule.

RESULTS AND DISCUSSION

COX-1 and COX-2 active sites have a similar structure, with the major differences being isoleucine substitution at amino acid residues 434 and 523 COX-1 (Ile434 and Ile523) to valine at COX2 (Val434 and Val523) at the side pocket[2]. The smaller valine residue at COX-2 forms a hydrophobic area and secondary pocket, due to pharmacophore SO₂Me and SO₂NH₂[39]. This secondary pocket in the COX-2 receptor is absent in COX-1, and become a target for the COX-2 selective inhibitor [40]. Another difference is that, in the side pocket, amino acid histidine (His513) for COX-1 is replaced by arginine (Arg513) in COX-2 and it allows interaction between the COX-2 receptor with polar compounds [41]. These basic differences at the COX-2 active sites are underlying selective mechanisms of COX-2 over the COX-1 receptor.

In this study, the inhibition of xanthone compounds against COX-2 and COX-1 enzymes was conducted in silico using a docking program named PLANTS. The PLANTS program has the ability to make a good prediction performance, and it can be performed in a short time [34]. The molecular docking process aims to predict the receptor ligand-binding mode, indicated by docking accuracy. Docking accuracy was considered successful if the RMSD value is less than 2 Å, and the process of docking the experimental compounds can be done.

Docking method validation

The redocking process of native ligands SC-558 and FLP was performed in a way as mentioned in the method. YASARA showed that the ligand copy almost coincides with COX-2 binding site at 1.3855 Å RMSD value, while COX-1 was at 0.33874 Å (both less than 2 Å); these met the validity criteria of docking. Comparison between the ligand conformation SC-558 and FLP from the x-ray crystal structure with redocking calculation results can be seen in fig. 2. The free binding energy between the copy ligand with COX-1 and COX-2 active site were-96.48 and-79.22, respectively. H-bonds or polar contacts of copy ligand with COX-1 enzyme were at amino acid residue Leu352, whereas those at COX-2 were in the positions of Ser353 and Arg513 (fig 3).

Docking molecular of celecoxib against COX-1 and COX-2 enzymes

The best conformation celecoxib docking with COX-1 and COX-2 enzymes is illustrated in fig. 4. It appears that the native ligand (yellow) and celecoxib (red) coincide; for the COX-2 enzyme, this overlap is very similar to the results of redocking native ligand (fig. 1b), while crossing each other for the COX-1.

Fig. 2: Comparison of conformation between the native ligand of the X-ray crystal structure (yellow) to the docking result (blue) and its RMSD values (<2 Å) (a) COX-1, (b) COX-2

(a) 3N8Z (RMSD = 0.3874 Å) (b) 1CX2 (RMSD = 1.3855 Å)

Fig. 3: H-bond positions among native ligands with amino acid residues of the COX-1 receptor (Leu352)(a) and with the COX-2 (Ser353 and Arg513) (b). Yellow dashed lines indicate the H-bonds between the ligands and protein

(a) COX-1 (b) COX-2

Fig. 4: Results of the celecoxib docking with native ligand of COX-1 (a) and COX-2 (b), performed with YASARA. Native ligand is yellow, whereas celecoxib is red

(a) 3N8Z (b) ICX2
The docking results of celecoxib and xanthone compounds against the enzyme COX-1 are shown in table 1, whereas that against COX-2 is listed in table 2. It appears that celecoxib, a COX-2 selective inhibitor, has free binding energy-78.13 (close to free binding energy of native ligand), while the enzyme COX-1 is much higher, at-34.01. Results of the celecoxib docking with native ligand 1CX2 and 3N8Z (fig. 4) may rationalise the extreme difference of free binding energy between the celecoxib with COX-1 and COX-2 enzymes. In this study, the more similar the docking overlap between the drugs with its receptor, the lower the energy that is generated; it also forms a more stable complex.

Table 1: Result of xanthones docking against the enzyme COX-1 (3N8Z), amino acid residues involved, and the number of hydrogen bonds

<table>
<thead>
<tr>
<th>Compound</th>
<th>Free energy</th>
<th>H-bonds</th>
<th>Atom component</th>
<th>Amino acid residues</th>
<th>n</th>
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<td>O atom (C=O) of Leu502</td>
<td>1</td>
<td></td>
</tr>
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<td>A</td>
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<td>O atom (C=O) of Met3122</td>
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<tr>
<td>B</td>
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<td>0 of C=O</td>
<td>H atom (OH) of Ser530</td>
<td>1</td>
<td></td>
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<tr>
<td>C</td>
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<td>H of OH (C-3)</td>
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<td></td>
</tr>
<tr>
<td>D</td>
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<td>0 of OH (C-6)</td>
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<tr>
<td>E</td>
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<td>0 of OH (C-6)</td>
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<td>F</td>
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<td>H of OH (C-3)</td>
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<tr>
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Table 2: Results of xanthones docking towards enzyme COX-2 (1CX2), amino acid residues involved, and the number of hydrogen bonds

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<th>Compound</th>
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<th>Atom component</th>
<th>Amino acid residues</th>
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<tr>
<td>I</td>
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<td>H atom (NH) of Arg120</td>
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The free binding energy between celecoxib with COX-1 is very high (-34.01) while the COX-2 is quite low (-78.13) and is close to the energy of the native ligand. This indicates the COX-2 selectivity of celecoxib. Celecoxib docked to COX-2 enzyme with minimal free binding energy and H-bond is similar to that ative ligand, at Ser153 and Arg113. H-bonds between ligand and enzyme can predict the strength and the catalytic activity of that complex [42].

It appears that free binding energy throughout xanthones derivatives of COX-2 enzyme is higher than celecoxib (except for compound A), and it is assumed that interactions of xanthone derivatives with the active site of COX-2 are weaker than celecoxib. The interaction energy against COX-1 is lower than celecoxib, which indicates that xanthones are also able to bind strongly to receptor enzyme COX-1; thus they are not selective for COX-2. The number of H-bonds between xanthones and COX enzymes is comparable for both COX-1 and COX-2 (table 1),
while the residue of primary amino acids are Arg$^{12}$, Ser$^{53}$ and Met$^{522}$ in COX-1, whereas in COX-2 they are Arg$^{12}$, Tyr$^{355}$, Tyr$^{385}$, and Ser$^{53}$ (table 2). These polar contacts of xanthones with both COX-1 and COX-2 enzymes differ greatly with celecoxib.

Fig. 5 shows the orientation of the ligand (celecoxib and compound F) against COX-1 enzyme, while against COX-2 is shown in fig. 6. It appears that celecoxib and compound F are capable of occupying the active site of COX-2 and COX-1 enzymes, though the H-bonds positions were at different amino acid residues. Celecoxib and compound F are located in an area bounded by several amino acid residues; among them are Trp$^{385}$, Trp$^{387}$, Phe$^{518}$, Val$^{349}$, Leu$^{352}$, Leu$^{359}$, Tyr$^{385}$, Trp$^{355}$, Arg$^{12}$, Ala$^{116}$, Phe$^{318}$, Val$^{322}$, Ala$^{127}$, and Leu$^{552}$. All the residues involved in protein-ligand interaction determine the inhibition activity of COX enzymes [2]. These amino acid similarities indicate that xanthones are able to occupy the active site of the COX-2 enzyme, and are believed to engage in inhibitory activity against the COX-2 enzyme.

(a) Celecoxib

(b) Compound F

Fig. 5: Orientation of celecoxib (a) and compound F (b) at the active site of the COX-1 enzyme

The active site of the COX-2 enzyme has three important regions. First there is the hydrophobic pocket, which is lined with amino acid residues Tyr$^{355}$, Trp$^{355}$, Phe$^{518}$, Ala$^{351}$, Tyr$^{349}$, and Leu$^{352}$. The second region is located at the gates of the active site and contains hydrophilic amino acid residues Arg$^{12}$, Glu$^{524}$, and Tyr$^{355}$; while the third is a side pocket with amino acid residues His$^{90}$, Arg$^{113}$, and Val$^{523}$ [19]. Some amino acid residues [Arg$^{12}$, Tyr$^{355}$, His$^{90}$, Arg$^{113}$, Val$^{523}$, Ser$^{53}$, and Glu$^{124}$] are believed to play a major role as a gateway entry of the ligand into the active site of the COX-enzyme, as shown in the analysis of the crystal structure of several inhibitors of COX-2 selective [43]. Several previous studies have shown that residue Arg$^{113}$ is involved in the interaction of H-bond formation in several compounds COX-2 selective inhibitors [44-45]. The low energy interactions and the similarity of amino acid residues with the native ligand in COX-2 docking process showed its activity as a selective inhibitor of COX-2.

(a) Celecoxib

(b) Compound F

Fig. 6: Orientation of celecoxib (a) and compound F (b) at the active site of the COX-2 enzyme

The interaction energy of xanthone compounds against the COX-2 enzyme varies between -73.57 to -79.18, with H-bonds found at amino acid residues Arg$^{12}$, Tyr$^{355}$, Trp$^{355}$, and Ser$^{53}$. These energies are slightly different to celecoxib (-78.13). H-bonds between the ligand and protein at amino acid residues Arg$^{113}$ and Tyr$^{355}$ are associated with formation of a stable complex in the COX-2 pockets in the celecoxib, indomethacin, and diclofenac; they are selective inhibitors of COX-2 [43]. The presence of H-bonds to the residue Arg$^{113}$, Tyr$^{355}$, Tyr$^{385}$, and Ser$^{53}$ are also associated with better COX-2 inhibitory activity [46]. In general, it appears that the overall xanthone derivatives show good docking capability of the protein COX-2; however they have
planar-shaped structures. This is slightly different from previous studies, in which it is mentioned that, if a compound is planar, it will have a little activity against receptor interactions COX-2 [47]. The free binding energy of xanthones against the enzyme COX-1 and COX-2 show small differences with-73.06 and-79.25 for COX-1, and-73.57 and-79.18 for COX-2. The slight difference of interaction energy and the number of H-bonds between xanthones and both COX suggest that the compounds have an equal COX-1 and COX-2 inhibitory activity or equipotency. For compounds with equipotency, the greater the molecule size, the more selective the inhibitory activity against COX-2. This is due to the increasing volume of the active site [48]. But of course this needs further investigation involving laboratory experimental studies.

CONCLUSION

Xanthone derivative compounds exhibit good inhibitory activity against the COX-2 enzyme. This is because they are able to occupy the COX-2 receptor in the active site comparable with celecoxib, but not at the same active sites. The free binding energy between xanthones with COX-2 and COX-1 enzymes is slightly different, so inhibition of COX-2 by xanthones are non-selective.

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CONFLICT OF INTERESTS

All the authors declared that there is no conflict of interest.

REFERENCES


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