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Anticancer potential of β-sitosterol and oleanolic acid as through inhibition of human estrogenic 17beta-hydroxysteroid dehydrogenase type-1 based on an *in silico* approach[†]

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The human estrogenic enzyme 17beta-hydroxysteroid dehydrogenase type-1 (HSD17B1) provides biosynthesis regulation of active estrogen in stimulating the development of breast cancer through cell proliferation. The β -sitosterol is classified as a steroid compound and is actually a type of triterpenoid compound that has a similar structure to a steroid. This similarity provides a great opportunity for the inhibitor candidate to bind to the HDS17B1 enzyme because of the template similarity on the active site. Several in silico approaches have been applied in this study to examine the potential of these two inhibitor candidates. Pharmacokinetic studies showed positive results by meeting several drug candidate criteria, such as drug-likeness, bioavailability, and ADMET properties. A combination of molecular docking and MD simulation showed good conformational interaction of the inhibitors and HSD17B1. Prediction of binding free energy (ΔG_{bind}) using the Molecular Mechanics-Generalized Born Surface Area (MM-GBSA) approach shows $\Delta G_{\rm bind}$ (kcal mol⁻¹) of C1-HSD17B1: -49.31 \pm 0.23 and C2-HSD17B1: -33.54 ± 0.34 . Meanwhile, decomposition energy analysis ($\Delta G_{bind}^{residue}$) suggested several key residues that were also responsible for the interaction with inhibitors, such as C1-HSD17B1 (six residues: Leu96, Leu149, Pro187, Met193, Val225, and Phe226) and C2-HSD17B1 (four residues: Ile14, Gly94, Pro187, and Val188). Hopefully, the obtained results from this research could be considered for the mechanistic inhibition of the HSDS17B1 enzyme at molecular and atomistic levels.

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Introduction

Steroids are involved in various biological processes, such as reproduction, aging, metabolism, and cancer.¹⁻⁴ The estrogen-

mediated effects of steroids are responsible for the genetic formation of breast tissue in women. One of the enzymes that plays a role in steroid regulation is human estrogenic 17betadehydrogenase type-1 (HSD17B1). The hydroxysteroid HSD17B1 enzyme is the first member of the HSD17B enzyme family that mediates the conversion of 17beta-hydroxysteroids, such as androgens and estrogens.5 This enzyme provides regulation in the final stages of active estrogen (estradiol) biosynthesis.6 It has been reported that the activity of this enzyme increases the level of estrogen. It stimulates the development of breast cancer cells through cell proliferation.7,8 Additionally, it has been reported that inhibition of HSD17B1 enzyme activity prevents proliferation in breast cancer cells in vitro and reduces tumor volume in human breast cancer lines grown (mice and rat models) in vivo.5,9 Therefore, this enzyme is a potential target in inhibiting the regulation of cancer cell development.6-9

Several research projects are looking for candidate inhibitors that can block the process of estrogen biosynthesis through enzyme HSD17B1 inhibition.^{4,9,10} There are several inhibitors with known activity (K_m) as HSD17B1 inhibitors, such as

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dihydrotestosterone (DHT): $32 \pm 9 \,\mu$ M, and androstandione: 26 \pm 9 $\mu M.^4$ It should be noted that both inhibitors have a similar structure, which is often known as a "mimic structure". The structural similarity is expected to have a better chance of binding to the targeted protein because of the template similarity on the active site.11 Thus, it is necessary to select a candidate inhibitor based on these considerations. Several natural product compounds can offer this feature, including triterpeand steroid groups as secondary metabolic noid compounds.12,13

Research on natural products as inhibitor candidates shows promising prospects in finding a potential inhibitor of the HSD17B1 enzyme.¹⁴ Additionally, natural products have provided good activity in inhibiting breast cancer cells.^{14,15} Some of them are β -sitosterol and oleanolic acid compounds, which were isolated and characterized from the stem bark of *Syzygium aqueum* in a previous study.¹⁵ The structural similarity of these compounds to dihydrotestosterone (a known HSD17B1 inhibitor) is expected to mean they will interact on the active site of the HSD17B1 enzyme. Structurally, it has –OH and –COOH groups at positions C3 (β -sitosterol) and C3, C28 (oleanolic acid) (Fig. 1). These groups increase the possibility of interacting with water molecules on the enzyme active site. Additionally, the presence of these groups can allow the formation of hydrogen bond interactions (donor or acceptor) with key residues on the enzyme active site at the molecular level.

Theoretical studies using an in silico approach provide an alternative way of finding candidate compounds that have potential as HSD17B1 inhibitors at the molecular level.^{16,17} A combination of molecular docking and molecular dynamics (MD) simulation provides a comprehensive structure-based approach to studying the interaction of the inhibitor with the HSD17B1 enzyme.¹⁶⁻¹⁸ Structure-based studies can provide a more detailed description of the inhibition mechanism through interactions with amino acid residues on the active site of the targeted protein. Molecular docking provides good calculation efficiency for inhibitor coordination on the enzyme active site.^{19,20} Meanwhile, the MD simulation offers a comprehensive analysis of the interaction between an inhibitor and the targeted protein during the simulation time.^{17,21} The evaluation proceeds by several considerations, such as grid-score²⁰ and binding free energy,²² aiming to see the binding affinity of inhibitor to HSD17B1. In addition, an evaluation of toxicity and drug-likeness provides an initial description of the pharmacokinetic properties of each inhibitor.23-25 Several considerations of the in silico approach offered in this research are expected to



Fig. 1 A suitable physicochemical space for the oral bioavailability prediction of candidates.

Methodology

Pharmacokinetic prediction

The drug-likeness and bioavailability calculations were performed by the SwissADME web service.²⁴ The criteria for druglikeness and bioavailability were calculated based on several rules, namely those of Lipinski and Veber. Meanwhile, prediction of ADMET (absorption, distribution, metabolism, excretion, and toxicity) of inhibitors as drug candidates used the web service pkCSM.²⁵ This prediction aims to study the activity of inhibitors as drug candidates when they enter the body.

Data set: ligand and receptor preparation

The selection of a target protein used the HSD17B1 co-crystal from the protein databank (PDB ID: 1JTV). The crystal complex contains a native ligand on the active site of the HSD17B1 enzyme as an inhibitor (testosterone, PDB ID: TES). The TES ligand was used as a reference or control in this study by considering it as an alternative binding mode to the HSD17B1 enzyme, which had been described in previous studies.⁴ The TES coordinates extracted from the co-crystals were used as initial coordinates for docking purposes. Meanwhile, the standard residues (amino acid residues) from co-crystals were used as receptors. Moreover, the key residues from the HSD17B1 active site, such as (Val143, Met147, Leu149, Pro187, Tyr218, His221, Val225, Phe226, Phe259, Leu262, Met279, and Val283) were used as the main targets for analysis purposes. β -Sitosterol (C1) and oleanolic acid (C2) were the candidate inhibitors used in this research. Those compounds were isolated (from Syzygium aqueum stem bark) and characterized in previous studies.¹⁵ The electrostatic potential (ESP) for candidate charge inhibitors was calculated using the Semiempirical Quantum Parametric Method-7 (SQM-PM7) contained in the Gaussian 16 package. Meanwhile, the AMBER FF14SB force field and the Austin Model 1-Bond Charge Correction (AM1-BCC) were applied to parameterize the receptors.

Molecular docking

Molecular docking was performed using the Dock6 package.²⁰ The inhibitor–HSD17B1 energy calculation was based on functional grid scoring (grid spacing: 0.30 Å, center *XYZ*: 11.08, 8.46, –11.24, and dimensions *XYZ*: 27.21, 28.28, 23.61). A flexible conformation with a functional grid scoring approach was mainly used to evaluate the redocking process. The successful redocking process gave the best superposition with an RMSD \leq 2.0 Å. The functional grid-based scoring process was analyzed in the gas state (eqn (1)) by considering parameters such as the van der Waals energy (E_{vdW}) and electrostatic energy (E_{ele}). The coordinates of each ligand obtained from the docking process were calculated in-depth using molecular dynamics simulations.

$$Grid-score = E_{vdW} + E_{ele}$$
(1)

Topology preparation

The coordinates of each ligand that had been obtained from the docking process were integrated into the MD simulation using the General AMBER Force Field (GAFF).²⁶ The topology preparation on each system used the *tleap* tools in the AMBER18 package. The topology preparation of each system took the form of ligand, receptor, complex, and complex-solvation. The solvent model used was TIP3P water solvent with a counter ion (Na⁺) as a neutralizing system. The system minimization process was carried out in three stages: water molecule minimization, complex minimization, and entire system minimization. All the minimization processes used 500 steps of steepest descent and 1500 steps of the conjugated gradient. All minimized systems were ready for use in the simulation process.

Molecular dynamics simulation

The MD simulation was performed using the AMBER18 package in several stages, such as heating, equilibrium, and trajectory production. The heating process was carried out gradually (10 K to 310 K) for 200 ps with a harmonic restraint of 30 kcal mol⁻¹ $Å^{-2}$. The periodic system equilibrium stages continued with harmonic restraints of 30, 20, 10, and 5 kcal mol⁻¹ $Å^{-2}$ for 1300 ps. The entire system was simulated to 100 ns (ensemble NPT: 310 K and 1 atm). The production stage during the simulation time (0–100 ns) aimed to produce trajectories for further analysis and evaluation purposes, such as conformational dynamics, binding affinity, hydrogen bonding, and solvent accessibility.^{22,27}

Binding free energy analysis

Calculation of binding free energy (ΔG_{bind}) and decomposition energy ($\Delta G_{\text{bind}}^{\text{residue}}$) was performed in the last 20 ns (80–100 ns) of the trajectories. This consideration was made based on calculation efficiency with the result that system stability was achieved in its trajectory range. The $\Delta G_{\rm bind}$ and $\Delta G_{\text{bind}}^{\text{residue}}$ calculations were made using the *MMPBSA.py* tools available in the AMBER18 package.²² MMPBSA.py is an efficient program with the flexibility to accommodate the needs of users performing end-state free energy calculations. In detail, the analysis of ΔG_{bind} and $\Delta G_{\text{bind}}^{\text{residue}}$ was undertaken using the Molecular Mechanics-Generalized Born Surface Area (MM-GBSA) approach. In summary, ΔG_{bind} can be described through eqn (2) by considering the energy complex ($\Delta G_{\text{ligand-}}$ HSD17B1), receptor ($\Delta G_{HSD17B1}$), and ligand (ΔG_{ligand}). Mathematically, ΔG_{bind} is described through eqn (3) by considering several parameters: the gas term (ΔG_{gas}), solvation term (ΔG_{sol}), and entropy change $(-T\Delta S)$. However, $-T\Delta S$ is neglected because of the high computational cost. Additionally, each ligand shows a similar structure, so the possibility of the resulting entropy change is not significant.28 Based on some of these considerations $-T\Delta S$ can be ignored in the ΔG_{bind} calculation. In detail, the energy component in ΔG_{bind} consists of bonded energy (ΔE_{bonded}), van der Waals energy (ΔE_{vdW}), and electrostatic energy (ΔE_{ele}). In particular, ΔE_{bonded} consists of bond, angle, and torsion energies with a conformational energy

equal to zero. Meanwhile, the energy component in $\Delta G_{\rm sol}$ is the total of the generalized Born models ($\Delta G_{\rm sol}^{\rm ele}$) and solvent-accessible surface area energy ($\Delta G_{\rm sol}^{\rm nonpolar}$). In detail, free energy binding can be calculated by considering some of these energy components (eqn (4)).

$$\Delta G_{\text{bind}} = \Delta G_{\text{ligand-HSD17B1}} - (\Delta G_{\text{HSD17B1}} + \Delta G_{\text{ligand}}) \qquad (2)$$

$$\Delta G_{\rm bind} = \Delta G_{\rm gas} + \Delta G_{\rm sol} - T\Delta S \tag{3}$$

$$\Delta G_{\text{bind}} = \Delta E_{\text{bonded}} + \Delta E_{\text{vdW}} + \Delta E_{\text{ele}} + \Delta G_{\text{sol}}^{\text{nonpolar}} + \Delta G_{\text{sol}}^{\text{ele}} \quad (4)$$

Results and discussion

Pharmacokinetic study: drug-likeness, bioavailability, and ADMET

Prediction of pharmacokinetic properties, such as druglikeness, bioavailability, and ADMET, aims to see the potential of a compound as a drug candidate. In this section, we try to describe some of the parameters responsible for the druglikeness and bioavailability of candidate compounds. The results showed that each compound has quite promising druglikeness and bioavailability criteria. Several drug-likeness criteria showed that there was only one violation of the Lipinski²⁹ and Veber³⁰ rules on M Log P (Table S1[†]). Notably, a good drug candidate shows fewer than a total of two violations of Lipinski's and Veber's rules. Compounds that meet the criteria of MW \leq 500 (g mol⁻¹), $M \log P \leq 4.15$, $\sum \text{HBA} \leq 10$, and Σ HBD \leq 5 (ref. 29) are highly appreciated as drug candidates. Moreover, compounds that have criteria of \sum rotatable bonds \leq 10 and TPSA \leq 140 Å² (ref. 30) are expected to increase the ability of drug candidates to penetrate cell membranes. Additionally, bioavailability shows that there are two violations in the range of oral bioavailability criteria for each candidate compound (Fig. 1). In general, the criteria for oral bioavailability meet the following requirements: lipophilicity (-0.7 <X Log P3 < 5.0), size (150 D < MW < 500 D), polarity (20 Å² < TPSA < 130 Å²), insolubility (0 < ESOL < 6), unsaturation (0.25 < Csp³ <1), and flexibility (0 < number of rotatable bonds <9).²⁴ Meanwhile, drug-likeness and bioavailability evaluations were also carried out on the TES ligand as a comparison (Table S1[†]). In summary, there was no violation of Lipinski's and Veber's rules. This data is supported by oral bioavailability that meets the overall criteria in a promising way (Fig. 1).

The ADMET study provides a clear image of a drug candidate's effects on the body. This parameter plays an important role in drug discovery.^{23–25} Prediction of ADMET properties shows that each compound shows promising potential (Table S2†). This is indicated by the value of Caco-2 permeability > 0.90 and intestinal absorption-human > 30% (+HIA).²³ These results showed that the compounds were absorbed into the body very well. The distribution of C1 compound showed that it was able to penetrate the blood–brain barrier (logBB > 0.3) and CNS permeability > -2 logPS. It identified that this compound could affect the work of the central nervous system (brain). On the other hand, compounds TES and C2 do not have the potential to penetrate the blood-brain barrier. However, they have the opportunity to penetrate the central nervous system (CNS). On the other hand, the metabolic parameters showed that each compound did not have any effect on the activity of metabolic enzymes (cytochrome isoenzymes), such as CYP1A2, CYP2C19, CYP2C9, CYP2D6, or CYP3A4. Hopefully, the compounds will not interfere with the activity of metabolic enzymes when consumed by the body. Crucial parameters such as toxicity are key parameters in the study of ADMET properties. Interestingly, all compounds belong to the non-toxic category (non-AMES toxicity and non-skin sensitization). However, compound C2 shows a category of hepatotoxicity that can cause damage to liver cells. Therefore, it is necessary to develop and modify C2 compounds for medicinal purposes. The ADMET results indicated that initial data on pharmacokinetic predictions show the compounds have potential worth considering.

Molecular docking: superposition and inhibitor-HSD17B1 conformation

Molecular docking can provide a clear initial picture of the coordinates of the active site on the targeted protein.³¹ However, it is necessary to pay attention to several parameters for the required process of coordinate selection and native ligand superposition. The redocking stage is one of the validation stages of the docking process for determining the active site of the targeted protein based on the coordinates of the native ligand. The redocking step has been widely implemented for efficient molecular docking calculations.19,20 The analysis was carried out on the TES coordinates as a reference for the cocrystals. The coordinates showed that the position of TES is in the pocket of the HSD17B1 enzyme (Fig. 2A). Additionally, the TES structural conformation indicated that the position of the -OH (O17) group is deep in the pocket. On the other hand, the position of the C=O (O3) group is in the outermost pocket (Fig. 2B and C). This is due to steric clashes between the 19methyl group and Leu149. The conformational stability of TES in the pocket is due to hydrophobic interactions and hydrogen bonding.4 The initial coordinate selection used a cluster sphere (radius 10.0 Å) selected from the TES coordinates (Fig. 2D). The results showed that the redocking process met the criteria with an RMSD value of 0.36 Å and a grid-score of -51.56 kcal mol⁻¹ (Fig. 2E). These results identify that the TES superposition has coordinates close to the co-crystal coordinates. It is intended that the docking inhibitor candidate process takes place at the coordinates of the HSD17B1 active site. Furthermore, the TES superposition shows hydrogen bonding (H-bond) interactions with His221 and Glu282 residues (Fig. 2F). The types of H-bond interactions are HBA (O17...H-N(His221)) and HBD ((Glu282) O···H-O17). Several parameters that have been described in the redocking stage were used for the candidate inhibitor docking process.

The optimized geometry of the candidate inhibitor was docked into the HSD17B1 active site. The results show that each candidate has a grid-score < grid-score reference, *i.e.*, C1: -69.90 kcal mol⁻¹ and C2: -61.46 kcal mol⁻¹ (Fig. 3). This indicates that the inhibitor candidate has good interaction in



Fig. 2 Redocking process: (A) targeting of the HSD17B1 active site. (B) Conformation of native ligand in the pocket area seen from the top. (C) Conformation of native ligand in the pocket area (rotated 90°). (D) Cluster sphere selected on the pocket area of the HDS17B1 enzyme. (E) Pose of native ligand for each conformation: crystal (dark gray), minimization (magenta), and flexible (green). (F) Interaction types between TES and HDS17B1 enzyme.

CI: -69.90 kcal/mal

Fig. 3 The molecular docking result of candidate-HSD17B1: the conformation of each candidate on the enzyme pocket area.

the gas term.³² In detail, the results of the energy contribution $(E_{vdW} + E_{ele})$ in kcal mol⁻¹ for each inhibitor were C1: $E_{vdW} =$ -70.25 and $E_{ele} = 0.35$ and C2: $E_{vdW} = -55.99$ and $E_{ele} = -5.47$. A similar energy contribution was also shown by TES-HSD17B1 (Fig. 2E). The interaction energy based on the functional grid scoring (gas term) for each inhibitor showed good binding stability with the targeted protein. This can be identified through a grid score that has a negative value. The grid score (higher negative value) indicated more thermodynamically stable interaction between the inhibitor and the targeted protein. Overall, the contribution of van der Waals energy (E_{vdW}) shows the largest contribution to the interaction between the inhibitor and HSD17B1 compared to electrostatic energy (E_{ele}) . Meanwhile, several residues are responsible for the inhibitor-HSD17B1 interaction (Fig. 2F and S1[†]), such as TES (7 residues: Leu149, Pro187, Tyr218, His221, Val225, Phe259, and Glu282), C1 (11 residues: Val143, Met147, Leu149, Tyr155, Cys185, Pro187, Val188, Tyr218, His221, Phe259, and Val283), and C2 (4 residues: Ser142, Leu149, Tyr155, and Gly186). Details of the overall interactions for each complex are provided in Table S3.† It should be noted that some of the parameters measured in the molecular docking are the initial conformations at the gas term. Therefore, the process of evaluating these results needs to be

studied using MD simulation to get more comprehensive results. However, molecular docking is quite helpful in efficiently determining the initial coordinates for each complex.

Conformational dynamics analysis

The topology of each system was prepared based on the coordinates of inhibitor–HSD17B1 obtained from molecular docking.^{16,17} The system that formed was evaluated more comprehensively to see its conformational dynamics. Several parameters were calculated to see the conformational dynamics quality of each system for 100 ns, including the total energy (E_{tot}) , potential energy (E_{pot}) , kinetic energy (E_{kin}) , the rootmean-square displacement (RMSD), radius of gyration (RoG), and the root-mean-square fluctuation (RMSF).

Energy analysis in each system in the form of E_{tot} , E_{pot} , and E_{kin} aimed to see the convergence achieved during the simulation time. Notably, the contribution of the E_{tot} parameter shows the contribution of $E_{pot} + E_{kin}$.³³ The results showed that each system had achieved good convergence. This can be identified by the absence of significant fluctuating changes in each system (Fig. S2†). The average value of the contribution of each energy is shown in Table S4.† A system with convergence can be used to continue the process of analyzing the compactness and stability.

System stability analysis is indicated by the complex RMSD value.^{21,34} The fact the simulation process reached 100 ns identified that each system has good stability. It is characterized by fairly good fluctuating stability from 15 ns to 100 ns (Fig. 4). Significant fluctuations occurred at the beginning of the simulation (0–200 ps) due to the heating process and increased gradually from 200 ps to 15 ns to reach an equilibrium process. Overall, each system has a stability difference that is not too significant. This is indicated by the average complex RMSD (nm) of TES-HSD17B1: 0.26 \pm 0.03, C1-HSD17B1: 0.26 \pm 0.03, and C2-HSD17B1: 0.27 \pm 0.03. Based on our assumptions, the stability that occurs in the candidate inhibitor system is caused by having a similar structure to the reference ligand. Thus, the positions of the C1 and C2 structures were able to occupy the HSD17B1 enzyme pocket well. However, this assumption will be



Fig. 4 Trajectory analysis: the root-mean-square displacement of each complex plotted during the 100 ns of MD simulation.

explained further in the next section by considering the binding affinity for each system.

The structural compactness is based on the RoG parameter.³⁵ The RoG analysis aims to see whether the structure is stably folded or not.

The calculation was carried out for 100 ns to see the RoG fluctuations shown in Fig. S3.† Changes in RoG fluctuations in each system did not show a significant difference. This is supported by the average values of RoG (nm), TES-HSD17B1: 19.73 \pm 0.14, C1-HSD17B1: 19.79 \pm 0.09, and C2-HSD17B1: 19.89 \pm 0.11. Changes in the values of these fluctuations indicate that the structure of each system is well folded. This is corroborated by the average structure taken per unit time of 20 ns for each system during the simulation time of 100 ns (Fig. S4†). Overall, there was no significant change in structural conformation during the simulation time (0–100 ns) in each system.

The RMSF was calculated to identify the flexibility of the protein over the simulation time (Fig. S5†). The TES-HSD17B1 system showed higher fluctuation compared to the other systems. In summary, the flexibility trend for each system was TES-HSD17B1 > C2-HSD17B1 > C1-HSD17B1 (Table S4†). Higher fluctuation (~2.00 nm) was shown by residues 130–133, 174–177, 199–201, 204–205, 208–210, 281, and 284–285 (alphahelix and loop regions). Moreover, twelve key residues (Val143, Met147, Leu149, Pro187, Tyr218, His221, Val225, Phe226, Phe259, Leu262, Met279, and Val283) that affect the main thermodynamic force for binding were also analyzed. The fluctuation changes from the key residues on the active site showed that His221, Met279, and Val283 residues had higher fluctuations compared to other residues (Fig. S6†).

The description of several parameters shows that the dynamic conformation formed during the simulation time had reached a fairly good convergence and stability. Therefore, further analysis was considered in the form of binding affinity, hydrogen bonding, and water accessibility using the last 20 ns (80–100 ns) of the trajectories. This consideration was made based on the efficiency of calculations that are often used for simulation analysis.^{21,34} It should be noted that this consideration is useful if the complex RMSD has achieved good enough stabilization.

Binding affinity prediction: binding free energy and key binding residues

The binding affinity was calculated for each complex using the MM-GBSA method available in the AMBER18 package.²² Determination of binding affinity was calculated based on 100 snapshots extracted from the last 20 ns of the trajectories. Additionally, the calculation was carried out by considering the energy contribution in the gas and solution terms described in Table 1. The calculation process for each energy component was the main focus for looking at the energy contribution to binding free energy ($\Delta G_{\rm bind}$).

The energy contributions at C1 and C2 show that ΔG_{bind} is promising as an inhibitor of the HSD17B1 enzyme. It is taken into consideration that the candidate ΔG_{bind} (C1 and C2) is stronger than that of the native ligand ΔG_{bind} as a reference (a

Table 1Binding free energy prediction calculated with the MolecularMechanics-Generalized Born Surface Area (MM-GBSA) method. Dataare shown as mean \pm standard error of mean (SEM)

Energy (kcal mol ⁻¹)	TES-HSD17B1	C1-HSD17B1	C2-HSD17B1
$\Delta E_{\rm vdW}$	-40.99 ± 0.27	-57.84 ± 0.24	-45.93 ± 0.34
ΔE_{ele}	-9.36 ± 0.29	-4.66 ± 0.33	-15.03 ± 0.41
$\Delta G_{\rm gas}$	-50.35 ± 0.33	-62.51 ± 0.36	-60.97 ± 0.43
$\Delta G_{\rm sol}^{\rm ele}$	23.12 ± 0.24	20.52 ± 0.30	32.84 ± 0.28
$\Delta G_{\rm sol}^{\rm nonpolar}$	-5.05 ± 0.01	-7.33 ± 0.02	-5.41 ± 0.02
ΔG_{sol}	18.07 ± 0.24	13.19 ± 0.29	27.42 ± 0.28
$\Delta G_{ m bind}$	-32.28 ± 0.25	-49.31 ± 0.23	-33.54 ± 0.34

higher negative value). Thus, the candidate inhibitors show promising potential to inhibit the HSD17B1 enzyme. It is hoped that candidates with good binding affinity will be able to bind strongly to the active site of the targeted protein.³⁶ The goal is to block the active site of the HSD17B1 enzyme, which is responsible for regulating the supply of active estrogen as fuel for breast cancer cell development.^{7,8}

Analysis of binding free energy (ΔG_{bind}) showed the energy trend (kcal mol⁻¹) of C1-HSD17B1 (-49.31 ± 0.23) < C2-HSD17B1 (-33.54 ± 0.34) < TES-HSD17B1 (-32.28 ± 0.25)

based on the MM-GBSA approach. The binding free energy cannot be separated from the effect of the energy contribution in gas (ΔG_{gas}) and solution (ΔG_{sol}) terms. In general, the energy contribution in the gas term ($\Delta E_{\text{vdW}} + \Delta E_{\text{elec}}$) has advantages compared to the solution term ($\Delta G_{\text{sol}}^{\text{nonpolar}} + \Delta G_{\text{sol}}^{\text{ele}}$). This is due to the unfavorable contribution of the polar solvation free energy ($\Delta G_{\text{sol}}^{\text{ele}}$) of the Generalized Born (GB) solvent model of each complex. In previous reports, the dielectric constant strongly influenced the calculation of $\Delta G_{\text{sol}}^{\text{ele}}$ and the variable dielectric model showed potential power.^{28,37} However, the energy contribution of $\Delta G_{\text{gas}} + \Delta G_{\text{sol}}$ showed quite promising results for each inhibitor.

Specifically, free energy key residues play a crucial role in observing the inhibitory mechanism of the HSD17B1 enzyme in calculating energy decomposition ($\Delta G_{\text{bind}}^{\text{residue}}$). The key residues with a promising free energy contribution were evaluated based on $\Delta G_{\text{bind}}^{\text{residue}} \leq -1.00 \text{ kcal mol}^{-1}$ criteria.³⁸ Calculation of $\Delta G_{\text{bind}}^{\text{residue}}$ was performed using 100 snapshots on the last 20 ns of the trajectories *via dcomp* tools. The results (Fig. 5) showed that there are key residues that have stronger free energy in each complex, including TES-HSD17B1 (five residues: Leu149, Pro187, His221, Ser222, and Val225), C1-HSD17B1 (six residues: Leu96, Leu149, Pro187, Met193, Val225, and Phe226), and



Fig. 5 The residual energy decomposition was plotted along with the simulation over the last 20 ns of each complex.



Fig. 6 Energy contribution from each residue of HDS17B1 to the binding of each ligand.

C2-HSD17B1 (four residues: Ile14, Gly94, Pro187, and Val188). These results identify that these key residues interact with inhibitors on the HSD17B1 pocket. The key residues Leu149 and Pro187 were highlighted in the inhibition mechanism of the HSD17B1 enzyme. Previous research reported that the Leu149 residue plays a role in steroid discrimination and modulates DHT levels in breast tissue.4 Additionally, the residue of Pro187 is a hydrophobic and aromatic residue that contributes to the main thermodynamic force for the binding mode. Therefore, the interaction evaluation of the inhibitors with key residues plays a crucial role in understanding the inhibition mechanism. Specifically, the energy contribution $\Delta G_{\rm bind}^{\rm residue}$ is affected by the energy contribution in the gas and solution terms, ΔE_{vdW} + $\Delta G_{sol(GB)}^{nonpolar}$ and ΔE_{ele} $\Delta G_{\rm sol(GB)}^{\rm ele}$ provided by Fig. 6. We can see that the energy contribution $\Delta E_{vdW} + \Delta G_{sol(GB)}^{nonpolar}$ is able to stabilize the free energy of each residue. It is inseparable from the contribution of ΔE_{vdW} energy in binding mode key residues³⁹

Hydrogen bonding analysis

The previous section described the interaction of amino acids with inhibitors at the molecular level. In this section, we try to present the interaction of amino acids with inhibitors at the atomistic level. Hydrogen bonding (H-bond) analysis is the main ligand–protein interaction.^{40,41} Analyses were performed using 2000 snapshots extracted from the last 20 ns of the trajectories. The evaluation process was carried out on number, type (HBD and HBA), distance, and % occupation of H-bond (Fig. 7).

The H-bond number parameter shows the number of H-bond interactions between ligand and protein. The results of the analysis during the last 20 ns of simulation time showed TES-HSD17B1 had three H-bonds, C1-HSD17B1 had three H-bonds, and C2-HSD17B1 had four H-bonds. The obtained number of H-bonds was used to look at occupation levels. This procedure aimed to see how long the bond was recorded during the simulation time. Surprisingly, the C1-HSD17B1 complex has only two H-bonds with a percentage occupation <10.0%. Meanwhile, the C2-HSD17B1 complex has four H-bonds, with two H-bonds having occupancy percentages >90.0%. Several reports state that the criterion for a strong category of H-bonding is H-bond \geq 70%.^{42,43} This explains that the H-bond interaction which has the highest percentage occupation deserves to be considered in the ligand-protein interaction. In detail, the TES-HSD17B1 interaction involves two H-bonds: (i) O17...HN(His221) with 20.8%, 2.97 Å, HBA and (ii) O17-H…O(Glu282) with 15.9%, 2.76 Å, HBD. The interaction of the H-bond with these two residues (His221 and Glu282) showed good correlation with the molecular docking results (Fig. 2F) and the results of previous studies.⁴ Meanwhile, the interactions of the H-bond in C1-HSD17B1 were (i) O3…H-O(Tyr155) with 4.45%, 3.08 Å, HBA, and (ii) O3-H… O(Glu194) with 2.45%, 2.78 Å, HBD. Additionally, the interactions of the H-bonds in C2-HSD17B1 were (i) O28a…HO(Tyr155) with 94.30%, 2.85 Å, HBA, (ii) O28b-H…O(Gly186) with 94.05%, 2.93 Å, HBD, (iii) O28b...HO(Ser142) with 9.10%, 3.17 Å, HBA, and (iv) O28a…HO(Ser142) with 4.50%, 3.12 Å, HBA. In particular, the C2-HSD17B1 complex has two strong H-bond categories with the % occupation of H-bonds \geq 70%. As explained, the presence of the -OH and -COOH groups contributes to the H-bond interaction. This further strengthens the existence of this group playing an important role in ligand-protein interactions at the atomistic level.

Solvent accessibility in the HSD17B1 active site

The solvent accessibility surface area (SASA) aims to see the ability of water molecules as solvents to access the surface of the active side area (pocket) during the simulation time. The role of water molecules on the active site surface plays an important role in mediating the interaction between the ligand and the targeted protein.44 It shows the solvent could access the surface of the active site up to ~9.30 nm². The C1-HSD17B1 system showed greater solvent access than the other systems. In detail, the SASA parameters (nm²) showed TES-HSD17B1: 7.91 \pm 1.01, C1-HSD17B1: 9.30 \pm 0.87, and C2-HSD17B1: 8.83 \pm 0.85. However, no system showed a significant difference in this parameter (Fig. S7[†]). As previously mentioned, the presence of water molecules on the surface of the active site of the targeted protein affects the stability of the protein structure. Moreover, its stability can be achieved through the interaction of hydrophobic contacts, hydrogen bonds, π -stacking, salt bridges, amide stacking, or cation- π interaction.⁴⁵

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Fig. 8 Trajectory analysis during the simulation over the last 20 ns: the radial distribution functions (g(r)) of the water oxygen atom and integration numbers (n(r)), up to the first minimum around the heteroatoms (black arrow).

The water molecule contained in the HSD17B1 active site needs to be studied for its distribution process during the simulation time. This aims to see the opportunities for water molecules to approach the ligand interface. In the previous section, the presence of the -OH and -COOH functional groups played an important role in the interaction with water molecules, in particular, the analysis of the oxygen (O) atoms in each ligand. Therefore, the possibility of water molecules approaching the oxygen atom can be analyzed through the radial distribution function (RDF).38 Analysis of this parameter was undertaken through the integration number on the first minimum value of the oxygen atom in each ligand.43 The TES-HSD17B1 complex showed that O3 atoms were more exposed to water molecules than O17 atoms. This can be seen in the atomic distance of O3 (n(r): 2.15 and g(r): 0.22) at the first peak, which is 0.35 nm. On the other hand, the O17 atom shows almost no hydration peak at all. Meanwhile, complexes C1-HSD17B1 and C2-HSD17B1 showed that water molecules were well distributed around the oxygen atom interface of each inhibitor. Except that O28b on the C2 inhibitor shows rather low access of water molecules at the interface. Meanwhile, the remaining atoms in each inhibitor C1 (O3) and C2 (O3 and O28a) showed promising access to water molecules (Fig. 8). In detail, the parameters measured for each atom are C1 (O3): distance = 0.33 nm, n(r) = 2.54, and g(r) = 0.37; C2 (O3): distance = 0.34 nm, n(r) = 2.51and g(r) = 0.32; and C2 (O28a): distance = 0.36 nm, n(r) = 1.03and g(r) = 0.04. The RDF analysis provides information about how the water molecules approach the oxygen atom in each ligand. The nonzero value at the first minimum peak indicates a high level of water transfer in the first solvation shell. These values are observed at TES (O3), C1 (O3), and C2 (O3 and O28a). This statement is supported by the value of n(r) being within \sim 2.1–2.5 at TES (O3), C1 (O3), and C2 (O3). In particular, C2 (O28a) has a consistent value, being stably solvated by one molecule of water. This can be seen through the first minimum distance at \sim 0.4 nm. As mentioned earlier, the oxygen atoms at TES (O17) and C2 (28b) have low hydration peaks. It can be seen that the first minimum of the two oxygen atoms is zero. This explains that there is no water transfer in the first solvation shell. Additionally, this condition also states that no water molecules are exposed at the interface of both oxygen atoms. Taking together all the findings discussed above, this explains that the presence of -OH and -COOH groups in each ligand can provide greater opportunities for interaction with water molecules. Based on the consideration of several parameters (SASA and RDF) above, the water accessibility of each system shows that the distribution of solvent molecules is quite well exposed.

Conclusions

In this work, we have presented an *in silico* approach including a pharmacokinetic study, molecular docking, and MD simulation to study β -sitosterol (C1) and oleanolic acid (C2) compounds as potential anticancer agents through the inhibition of human estrogenic 17beta-hydroxysteroid dehydrogenase type-1 (HSD17B1). Pharmacokinetic studies suggest that each candidate had non-toxic criteria worthy of consideration. A

study of the interaction between the inhibitor and the HSD17B1 enzyme at the molecular level showed good correlation through a combination of molecular docking and MD simulation. The correlation is shown through the grid-score and binding free energy (MM-GBSA) with the energy trend C1-HSD17B1 < C2-HSD17B1 < TES-HSD17B1. Meanwhile, key residues Leu149 and Pro187 were highlighted in the HSD17B1 enzyme inhibition mechanism showing $\Delta G_{\text{bind}}^{\text{residue}} \leq -1.00 \text{ kcal mol}^{-1}$. More specifically, the atomistic level was studied to see the interaction of ligands and amino acids through the hydrogen bond in the pocket area of the targeted protein. The results showed that the presence of -OH and -COOH groups in each inhibitor had a crucial influence in stabilizing the inhibitor-HSD17B1 interaction. In conclusion, all the calculation results suggest that candidate inhibitors C1 and C2 have worthy potential to inhibit the HSD17B1 enzyme.

Conflicts of interest

The authors declare no conflicts of interest.

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