Computational View of β-Asarone–Human Serum Albumin Interaction

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Abstract: β-Asarone (BAS), a bioactive phytochemical from the medicinal herb, Acorus calamus Linn., has shown many pharmacological activities. Computational docking studies unveiled the interaction site of BAS on the human plasma carrier, albumin. The primary binding arrangement of BAS was placed at Sudlow’s Site I of HSA, which is pinpointed in subdomain IIA of albumin. Hydrophobic and van der Waals forces together with hydrogen bonds appear to secure the BAS-albumin complex. The BAS at Site I was surrounded by more hydrophobic and polar residues than those seen at Site II, as evidenced by LigPlot+. Therefore, the interaction between BAS and albumin at Site I seems to be comparatively more stable owing to more vital interactions.

Keywords: β-asarone; human serum albumin; drug-albumin interaction; computational docking; cluster analysis.

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1. Introduction

The β-asarone (1-propenyl-2,4,5-methoxybenzol) (BAS) with known chemical structure (Figure 1) is obtained from the medicinal herb, Acorus calamus Linn.[1]. Different pharmacological activities associated with BAS are anti-inflammatory, antiepileptic, antifungal, and antidepressant activities [2–5]. Several studies have reported that BAS functions as a promising anticancer agent in treating various carcinomas, viz., lung, gastric, hepatocellular, and colorectal cancers [6–9]. Due to its ability to cross the blood-brain barrier, BAS is frequently distributed in the cerebral tissue [10]. BAS has also shown positive results for treating neurodegenerative diseases, viz., Alzheimer’s and Parkinson’s diseases, depressive disorder, and ischemic stroke [5, 11–12].
Drug-plasma protein binding influences a drug's transportation and distribution to its specific target locations [13,14]. Such interaction significantly impacts the drug's pharmacokinetics, such as enhanced \textit{in vivo} half-life, prolonged efficacy, improved solubility, fast elimination, and reduced toxicity [15–17]. Therefore, the molecular interaction study between a drug and the protein is vital. Both experimental and \textit{in silico} techniques are being used to characterize ligand-protein interactions. \textit{In silico} analysis allows the first-hand interpretation of the preferred binding fashion of the drug to the protein's binding site at the atomic level [18–25]. However, these results need to be validated by experimental approaches. Albumin, present in human serum (HSA), is primarily responsible for carrying many drug molecules to their target organs throughout the bloodstream [15]. The drug's binding to albumin is expedited owing to the existence of two drug-binding sites, \textit{i.e.}, Sudlow's Sites I and II, placed in subdomains IIA and IIIA, respectively [26,27]. Several recent reports have shown the binding preference of various ligands to HSA at either of these two ligand-binding sites [28–33]. This paper describes the anticipated binding fashion and interaction of BAS with HSA at the atomic level using molecular docking assessments.

2. Methods

2.1. Molecular docking.

The computational docking simulation required the three-dimensional structures of BAS and HSA. BAS structure was generated using Avogadro software and optimized with the MMF94 force field [34,35]. Meanwhile, the PDB website was used to obtain the HSA structure (PDB ID: 1BM0). Optimization of HSA configuration involved the discharge of crystallized water, placement of polar hydrogens, as well as Kollman charges via AutoDockTools [36]. The docking simulations of BAS at the HSA Sites I and II were conducted independently with 100 search runs utilizing the Lamarckian genetic algorithm. The grid box was centralized at Site I (x: 35.36, y: 32.41, z: 36.46) and Site II (x: 14.42, y: 23.55, z: 23.31) separately with a dimension of 70 × 70 × 70 and 0.375 Å spacing. Other docking parameters were fixed as mutation (0.02), crossover operator weights (0.8), and elitism (1). The analysis of molecular docking results involved clustering the runs based on the 2.0 Å root-mean-squared deviation (RMSD). Further interaction and visualisation were done utilizing UCSF Chimera and LigPlot+ [37,38].

3. Results and Discussion

3.1. Molecular docking results.

The atomic interactions generated between BAS and HSA can be predicted and visualized through molecular docking simulation. The independent site-specific docking
simulation also enabled the identification of BAS binding priority on HSA Sudlow's binding sites. Based on the docking cluster analyses (Figure 2), Sites I and II were associated with overall 5 and 8 multi-member conformational clusters, respectively. The maximum populated cluster on Site I comprised 68 members ($\bar{X} = -25.7 \text{ kJ mol}^{-1}$), whereas Site II included 49 members ($\bar{X} = -17.2 \text{ kJ mol}^{-1}$). Meanwhile, the least calculated binding energy of BAS at Site I was $-26.9 \text{ kJ mol}^{-1}$ compared to $-18.2 \text{ kJ mol}^{-1}$ at Site II. All these initial analyses stated that the preference of BAS was leaning towards Site I of the protein.

**Figure 2.** The cluster analyses of 100 AutoDock docking runs (RMSD = 2.0 Å) of BAS to the drug-binding sites, *i.e.*, (A) Site I and (B) Site II of HSA (PDB: 1BM0).

Further analyses were accomplished by utilizing the lowest binding energy complex structures of BAS and albumin at Sites I and II (Figure 3A–C). The investigation revealed that two H-bonds were formed when BAS was bound at Site I, while no hydrogen bond involvement was seen at Site II (Table 1). The surrounding pocket contributing to the polar and hydrophobic interactions (Figure 4A and 4B) showed engagement of many residues in this interaction when BAS was bound at Site I compared to Site II. The presence of hydrogen bonds plus other non-covalent interactions formed between BAS and Site I of HSA would have increased the complex stability. Overall, these analyses revealed the capability of BAS forming a stable complex while it is bound at Site I of HSA compared to Site II, hence favored its binding choice towards Site I of the protein.
Figure 3. (A) Binding alignment of BAS (rendered in the sticks) in albumin binding sites I and II on the basis of least binding energy. Different domains I, II, and III of HSA are represented in orange, sky blue, and green colors, respectively. The enlarged view of the binding sites showed hydrogen bonds (green lines) within the protein's amino acid residues (rendered in the yellow stick) and BAS at Site I (B) and Site II (C) of HSA.

Table 1. Anticipated hydrogen bonds between atoms of amino acid residues of HSA and BAS at Sites I and II.

| HSA binding sites | HSA residue : atom   | BAS atom | Distance (Å) |
|------------------|----------------------|----------|--------------|
| Site I           | Tyr-150 : hydrogen-hydrogen | O        | 2.04         |
|                  | Arg-257 : hydrogen-electron | O        | 1.89         |
| Site II          | --                   | --       | --           |

Figure 4. LigPlot+ maps present the hydrophobic and polar interactions between atoms of BAS and the amino acid residues of HSA at Site I (A) and Site II (B).

4. Conclusions

The findings obtained from our study indicated that the locus of BAS binding site on HSA was predicted at Site I (subdomain IIA). The stabilizing forces for the formed complex...
were found to be hydrophobic and polar interactions along with H bonds. The orientation of BAS bounded at Sudlow's Site I of HSA permitted substantial binding strength energy owing to increased non-covalent interactions and subsequently a stable complexation.

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**Conflicts of Interest**

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