Study of Interaction Between Bovine Serum Albumin and Dolutegravir Intermediate: Fluorescence and Molecular Docking Analysis

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Abstract: Novel (4R,12aS)-7-methoxy-4-methyl-6,8-dioxo-3,4,6,8,12,12a-hexahydro-2H-pyrido-[1′,2′:4,5]-pyrazino[2,1-b][1,3]oxazine-9-carboxylic acid (L) was synthesized and characterised. The interaction between bovine serum albumin (BSA) with L was scrutinized by steady-state fluorescence spectroscopy, fluorescence anisotropy, fluorescence lifetime, and molecular docking methods. The fluorescence titration experiments of BSA resulted in fluorescence quenching with the incremental addition of L. The conformational binding of L to BSA has been investigated by molecular docking analysis. The molecular probe's best conformation showed the affinity as free binding energy release of -7.93 Kcal/mol. The docking analysis confirms that ligand binds in the near vicinity of TRP-213 in the binding pocket of subdomain IIA.

Keywords fluorescence; anisotropy; lifetime; docking analysis; receptor.

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1. Introduction

The study of protein-ligand interactions has played an incredible role in its industrial, biological, cosmetics, and pharmaceutical applications. Serum albumins are the most abundant protein in plasma [1-5]. Its remarkable acceptor capabilities with various active molecules are a dynamic tool in developing novel therapeutic agents, the prognosis of the pharmacokinetic behavior of drugs, and pharmacodynamics modulations [6-10]. BSA acts as a kind of serum albumins, which has been used as a model protein for various studies. BSA is a large globular protein. Its molecular weight is 66000 Da [11-14]. It is constituted by 582 amino acid residues. It also comprises 17 disulfide bridges and a free -SH groups. It is made up of three distinct homologous domains I, II, and III; each homologous domain is divided into two subdomains A and B. It embraces two tryptophan residues, namely, Trp-213 and Trp-134. It is well known that the Trp-134 is located in a hydrophillic environment close to the protein surface (subdomain-I B). However, Trp-213 is located in the largest hydrophobic cavity, domain II (subdomain-II A). The major binding sites of BSA are situated in subdomains IIA and IIIA [15-23]. For protein-ligand interactions, site maker fluorescence molecular probes are utilized. The interactions between BSA with ligands led to important information about the absorption and distribution of the drug. The fluorescent emission plays an important role in protein-ligand interactions due to their high selectivity, excellent sensitivity, and quick response abilities. The
fluorescence emission can provide essential information about the binding mechanism, binding mode, binding constants, and site [24-28].

2. Materials and Methods

2.1. Materials.

BSA and spectroscopic grade solvent were obtained from Sigma, USA, and Merch, India and used. The 1H-NMR spectra were recorded on Varian-AS 400 spectrometers. The fluorescence spectra were obtained by using a Perkin-Elmer LS-55 spectrofluorimeter. The derivative (L) [29] by reported literature is shown in Figure 1. A 10 mM phosphate buffer pH 7 was prepared to prepare protein solution and fluorescence titration experiments.

![Figure 1](https://biointerfaceresearch.com/)

2.2. Protein sequence and ligand preparation.

Bovine Serum Albumin (BSA) protein sequence was retrieved from the Protein Data Bank (PDB id: 4f5s). The file opened in Discovery Studio Visualiser where Heteratoms and chain B were removed and the protein file saved in pdb format. The L was built in the Chemdraw (Chemoffice-17). The L was opened in Chem3D and subjected to energy minimized using the force field MMFF94 with a maximum number of iteration 500 and a minimum RMS gradient of 0.1. The energy-minimized structures were saved in pdb format for further use for docking purposes.

2.3. Molecular docking study.

The docking study was performed using Autodock 4.2.5. Auto dock predicts the bound conformations of flexible ligands to bio macromolecular targets using the Lamarckian genetic algorithm. It was done according to the literature report [30-31]. In brief, the PDBQT files were generated for protein and ligands molecules by adding polar hydrogen and Kollman charges in the case of the macromolecule, whereas in molecular probes, non-polar bonds are merged, and Gasteiger charges were applied. The grid box dimension was fixed to 74,68,66 (x, y, z coordinates) with default 0.375 Å grid point spacing. For docking simulation and conformational search, Lamarckian Genetic Algorithm (LGA) was chosen Moreover, the number of the evaluation were set to be 20 and rest other parameters were kept default. The molecular docking result was evaluated in PyMOL (The PyMOL Molecular System, Version 2.0 Schrodinger, LLC, and Discovery Studio Visualiser).

2.4. Binding constant calculation equation.

Binding constants of BSA and HSA with substrates were determined using the Benesi-Hildebrand equation by the fluorescence method in Eq. 1.
\[
\frac{1}{\Delta F} = \frac{1}{\Delta F_{\text{max}}} + \left(\frac{1}{K_a[C]}\right) \times \left(\frac{1}{\Delta F_{\text{max}}} \right)
\]  \hspace{1cm} (1)

Here \(\Delta F = (F_x - F_0)\) and \(\Delta F_{\text{max}} = F_\infty - F_0\), where \(F_0, F_x,\) and \(F_\infty\) are the emission intensities of BSA in the absence of substrate, at an intermediate substrate concentration, and at a concentration of the complete interaction, respectively. \(K_a\) is the binding constant, \(C\) is the concentration of the substrate, and \(n\) is the number of substrate-bound each to BSA (here \(n = 1\)).

3. Results and Discussion

The derivative (L) was prepared based on the literature report [29]. Fluorescence titration experiments show the interaction between proteins and ligands. BSA comprises three fluorophores, tryptophan, tyrosine, and phenylalanine [30-33]. Due to the low quantum yield of phenylalanine and almost quenched tyrosine features, the intrinsic fluorescence occurred mainly due to tryptophan residues. The interaction between proteins with various molecular probes can be diminished the fluorescence intensity of protein, which is known as fluorescence quenching. The fluorescence quenching mechanism can be dynamic, static; the static and dynamic quenching is simultaneously involved. Dynamic quenching ascribes from interaction through the collision between the fluorophores and quencher [34-35]. However, static quenching results from the formation of a complex between the fluorophores and quencher [36]. Fluorescence quenching of the protein provides important evidence about the molecular microenvironment modifications in the chromophore molecules' vicinity and the protein-ligands interactions [37-40].

BSA shows a strong emission band at 344 nm upon excitation at 295 nm. The peak at 344 nm is occurred due to the presence of tryptophan residue in the protein. The sequential addition of ligand to BSA shows the decreased fluorescence emission with increasing the concentration of ligand. It indicates the binding of the ligand with BSA. Upon addition of ligand to BSA, fluorescence emission quenched along with a redshift (13 nm) of emission band from 344 to 357 nm indicates the ligand's binding to BSA shown in Figure 2a. The literature found that fluorescence emission shifted from lower to higher wavelengths upon protein unfolding [41-42]. The equilibrium association constant (\(K_a, 6.80 \times 10^4 \text{ M}^{-1}\)) determined from the plot shown in Figure 2b. is presented in Table 1.

![Figure 2](https://biointerfaceresearch.com/13104)

**Figure 2.** Fluorescence spectra of 6 \(\mu\text{M}\) (3 mL) BSA in the presence of (a) molecular probe L; with increasing concentrations in the range of 0 to 48 \(\mu\text{M}\) (phosphate buffer, pH 7); (b) Binding constant plots of L with BSA. Solid lines indicate the fit using the Hill equation.
To gain the more information regarding protein-ligand interaction, it has been performed steady-state fluorescence anisotropy of BSA in the absence or the presence of the compounds by using equation 2. The increase in anisotropy values of BSA in the presence of compounds support their binding properties. From these values, it is clear that BSA’s initial symmetric state with anisotropy value close to zero [43] changed upon interactions with L. The higher anisotropy values indicated their ability to change the shape of the BSA protein. This is also in agreement with the observed order of the binding constants.

\[ r = \left( \frac{1_{\text{VV}} - G_{\text{VH}}}{1_{\text{VV}} + 2G_{\text{VH}}} \right) \] (2)

### Table 1. Binding constant and Anisotropy of BSA with L.

| Sample | \( K_a \) [M⁻¹] \( \times 10^4 \) | Anisotropy of BSA |
|--------|-------------------------------|-------------------|
| BSA+L  | 6.80                          | 0.116             |

Note: a = 2 μM ( 3 mL) BSA in 20 mM in phosphate buffer, pH 7.  
b = Anisotropy of BSA = 0.082

### 3.1. Molecular docking result.

BSA contains three homologous domains I, II, and III. Each homologous domain is divided into two subdomains, A and B. The molecular docking analysis can be used to retrieve information about the protein-ligand interactions [44]. The interaction between protein-ligands occurred mainly in subdomains IIA and IIIA. The literature envisaged that the molecular probes or drugs bind at site IIA or IIIA [45-46]. To gather more information about protein-ligand interactions, we have performed molecular docking analysis. The docking calculations are depicted in Table 2. Figure 3a shows that the molecular probe bind in subdomain IIA near the TRP-213 residue. The hydrogen bond architecture between the molecular probes and amino acid residues plays a tremendous role in stabilizing protein-ligand interactions. Besides that, L is enclosed by several hydrophobic and polar moieties. Hydrophobic interactions and charged and polar residues also provided extra stability of protein-ligand interactions through various amino acid residues such as ASP-450, SER-453, LEU-454, ARG-194, LEU-197, ARG-198, SER-201, LEU-210, LEU-480, LEU-346, VAL-342, SER-0343, and ALA-341. From these observations, it was demonstrated that ligand was bound with BSA.

### Table 2. Key amino acids displayed polar interaction and hydrophobic interaction (underline) on docking.

| Ligand | Predicted free energy of binding by docking, \( AG \), Kcal/mol | Polar interactions | Residue within 4 Å region |
|--------|---------------------------------------------------------------|-------------------|--------------------------|
| L      | -7.93                                                         | ARG-217           | ASP-450, SER-453, LEU-454, ARG-194, LEU-197, ARG-198, SER-201, LEU210, LEU480, TRP213, LEU-346, VAL-342, SER343, ALA-341 |

Zhang and his coworkers reported that increased protein-ligand complex stability is due to the increased hydrophobicity nature in his pioneer work. The formation of hydrogen bond architecture between protein and ligand diminished the hydrophilicity characteristics and helped increases hydrophobicity, which stabilizes the protein-ligand interaction [47]. The best conformation of ligand molecules showed the affinity as free binding energy release of -7.93 Kcal/mol. It is due to the spontaneous complexation reaction. The docking analysis confirms that ligand binds in the near vicinity of TRP-213 in the binding pocket of the site I.
3.2. Fluorescence lifetime study.

The fluorescence lifetime decays were studied for BSA with ligand (Table 3). Depending on the conditions, native BSA showed three [48-51] or two lifetimes [52-54]. It has been observed that three lifetimes for BSA as the best-fit decay profile is obtained from three exponential rather than the two exponential functions. At the same time, the interactions of the ligand with BSA changes these lifetimes significantly.
The drastic change in the lifetime suggests the strong interactions by the tryptophan residue in the BSA, as shown in Figure 4b. This occurs as this lifetime is characteristic of a free tryptophan [55-60]. Upon addition of ligand to BSA, the lifetime changes drastically, and values are listed in Table-3.

| Samples   | \(\lambda_{em}\) (nm) | \(\tau^1\) (ns)/(f)% | \(\tau^2\) (ns)/(f)% | \(\tau^3\) (ns)/(f)% | \(\chi^2\) |
|-----------|-----------------------|-----------------------|-----------------------|-----------------------|-----------|
| BSA       | 344                   | 0.449 (2.63)          | 3.762 (35.24)         | 7.327 (62.13)         | 1.06      |
| BSA + L   | 4.214(28.26)          | 7.241(35.92)          | 1.118 (35.82)         | 1.14                  |           |

### 4. Conclusions

In this present study, the interaction between the synthesized derivatives with BSA was investigated using steady-state fluorescence, fluorescence anisotropy, fluorescence lifetime, and molecular docking analysis under physiological condition pH 7. Steady-state fluorescence showed quenching fluorescence emission. BSA's binding with L was mainly due to hydrogen bonding, polar interactions, and hydrophobic interaction. Fluorescence anisotropy and lifetime experiments studies have clearly confirmed the binding of the ligand with BSA. Further, the molecular docking analysis confirmed the interaction between BSA and ligand.

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### Conflicts of Interest

The authors have declared that no competing interests exist.

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