Synthesis, characterization, DFT analysis and docking studies of a novel Schiff base using 5-bromo salicylaldehyde and β-alanine

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ABSTRACT
Poly(ADP-ribose) polymerase-1(PARP-1) is a DNA-dependent enzyme, forming part of ADP-ribosyltransferase family. Although some PARP inhibitors find therapeutic applications in cancer therapy and exhibits crucial role in DNA damage response. Here a novel Schiff base, (E)-3-((5-bromo-2-hydroxybenzylidene) amino) propanoic acid was synthesized using 5-bromo salicylaldehyde and β-alanine. Characterization was carried out using IR, UV-Vis,1H and 13C NMR and mass spectrum. Present study involves the evaluation of a novel Schiff base as an inhibitor against human breast cancer cell lines (pdb:3GEY) using 2-(dimethylamino)-N-(6-oxo-5,6-dihydrophenanthridin-2-yl) acetamide (DDA) as a native ligand. In silico study of 3GEY inhibitor is a variant of PARP-15, docking with two different ligands (E)-3-((5-bromo-2-hydroxybenzylidene) amino) propanoic acid (SBL) and the native ligand. Synthesized ligand docked in to the B chain of PARP enzyme binding site to visualize the best docked pose and favorable ligand-protein binding interactions. Swiss ADME tool determines the drug likeness and strongly suggests that SBL can be a promising candidate to fight against breast cancer. DFT studies were done to support the experimental results using B3LYP/6-311++G(d,p) and geometry optimization was performed. Various thermodynamic parameters and NLO properties were found out. ECD and VCD spectrum were explained using DFT studies. Vibrational and Raman frequencies were also reported. HOMO-LUMO band gaps, Mulliken charges were calculated and the electrostatic potential surface was mapped with various properties. Experimental findings obtained are in good agreement with that of theoretical DFT analysis.

1. Introduction
PARP inhibitor protein (Poly(ADP-ribose)/polymerase) has DNA repairing properties so popularly called as guardian angel of DNA. It inhibits mitosis, inflammation, non prescription and cytotoxic properties [1]. Thus PARP-1 act as an anti-cancer agent while PARP inhibitors find applications as chemo-sensitizing agents in various kind of cancer growths [2, 3]. The parent PARP protein has six domains. Two of these six domains are Zn binding domains having the power to repair damaged DNA, and the third one is Zn binding domain that directs DNA-dependent enzyme activation [4, 5]. One of the domains of PARP-15 zinc finger is antiviral protein and it gives important antiviral response in mammals [6]. The isolated transferase domain of PARP-15 has considerable auto modification activity in invitro studies [7]. So pharmaceutical industries used PARP inhibitor protein as a new class of drug. Here amino acids in the active site of 3GEY is a variant of PARP target protein which makes interaction with a ligand containing functional groups. This protein has four ligands out of which one ligand is the active one. Higher score steric interaction between protein and ligand indicates its high binding affinity [1]. So, this ligand has a crucial part in DNA damage repair and enables to treat ovarian, fallopian tube, primary peritoneal and breast cancer [4]. Some recent studies were already reported with PARP protein and a novel Schiff base, 1,3,4-oxadiazoles, which was subjected to insilico analysis, against breast cancer cells. Molecular docking study by using Schrodinger software shows that enzyme shows a very high value of binding affinity against 1,3,4-oxadiazoles Schiff base [8].
5-bromosalicylaldehyde containing Schiff base have antitumor activity [9] is applied for manufacturing anti-cancer drug and Schiff bases has unique biological activity, can be used as low toxic in design and synthesis, and has got several applications like antibacterial and fungal, anti-inflammatory [10], anti-tubercular [11], anti-oxidant [12], and kills parasitic worms [13, 14]. A naturally occurring non-essential amino acid, β-alanine form part of carnosine, which in turn helps to buffer acid in muscles [15] may exhibit tumor progression [16], which has been most vastly investigated [17]. β-alanine treatment will bring out anti-tumor activity.

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Figure 1. Optimized geometry of SBL.

Figure 2. FT-IR spectra of SBL (a) Experimental (b) Simulated.
Figure 3. UV-Visible spectra of SBL (a) Experimental (b) Simulated.

Figure 4. Mass spectra of SBL.

Figure 5. Raman spectra of SBL.
results in selective tissues that express carnosine synthase [18]. When breast epithelial cells [19, 20] were treated with β-alanine, mechanism of action and specificity of the reagent need further investigation as a co-therapeutic agent in tumor treatment [21].

Schiff bases are much useful owing to their synthetic accessibility, structural features, and varieties. There have been many reported applications of antiviral activity and agriculture Schiff base has numerous applications in modern technology. It is used as photodetector in biological systems and used as a photo stabilizer and optical sound recording technology [12]. Schiff base ligands having oxygen and nitrogen donor atoms shows wide spectrum of biological properties. They can be used in homogeneous and heterogeneous catalysis [10]. Schiff bases possess an imine group –N=CH-, which is participating in transamination and racemization reactions [22]. Properties of Schiff base include chelation [23], thermal stability [24], non-linear optical property [25], and proton transfer [26]. Thermally stable ones are used as stationary phases in gas chromatography [27]. The nonlinear optical property of Schiff bases provides applications in electronic, opto-electronic and photonic systems [28]. Schiff bases are used as catalysts in photo-electrochemical processes. They can also be used as corrosion inhibitors of mild steel, copper, aluminum, zinc etc. [27].

The present study focuses on the synthesis of novel Schiff base (E)-3-((5-bromo-2-hydroxybenzylidene) amino) propanoic acid (SBL) by the condensation reaction of 5-bromosalicylaldehyde & β alanine in a 1:1 ratio using ethanol as the solvent. The synthesized Schiff base then characterized was performed using IR, UV, 1H and 13C NMR and mass spectrum. Present work focus in silico study of 3GEY inhibitor is a variant of PARP-15, docking with two different ligands (E)-3-((5-bromo-2-hydroxybenzylidene) amino) propanoic acid (SBL) and 2-(dimethylamino)-N-(6-oxo-5,6-dihydrophenanthridin-2-yl) acetamide (DDA). Here the SBL was optimized using DFT computational method [27]. Molecular docking into the protein 3GEY responsible for breast cancer, was carried out by evaluating binding affinities with SBL and finding drug-likeness using the SWISS ADME tool. Density functional theory calculations were done to support the experimental results using Gaussian16 W software. Visualization of results were done with GaussView 6.1 software. B3LYP/6-311+G(d,p) level of theory was chosen for optimizing geometry of SBL. Various thermo-chemical parameters were reported. Vibrational and Raman frequencies was also reported. HOMO-LUMO band gaps, NLO Properties and Mulliken charge were calculated and the electrostatic potential surface was mapped with various properties [29].
2. Experimental

2.1. Chemicals and instruments

All the chemicals used were of analytical grade and used without any further purification. 5-bromosalicylaldehyde, β-alanine and ethanol were purchased from Avra Chemicals. UV-Visible spectrum was recorded in the range of 200–800 nm in PerkinElmer Lambda 365. IR spectrum of synthesized ligand in the frequency of 400-4000 cm\(^{-1}\) was recorded on Perkin Elmer spectrometer IR version 10.6.1. \(^1\)H and \(^13\)C NMR spectrum was measured with 400 MHz Varian Joel spectrophotometer. HR-MS of the title compound was recorded using Accella 600 HPLC system with autosampler and PDA detector. DFT calculations were done using Gaussian 16W software and visualized using GaussView 6.1 software. Molecular docking was done using Auto Dock Vina v.1.2.0 tools.

2.2. Preparation

2.2.1. Synthesis of Schiff base

The Schiff base (E)-3-((5-bromo-2-hydroxybenzylidene) amino) propanoic acid [SBL] was synthesized by the condensation of 5-bromosalicylaldehyde and β-alanine [30] by dissolving each of the organic compounds separately in ethanolic solution in a beaker in 1:1 ratio. The solution in both the beakers are then poured onto RB flask which was then refluxed for about 4 h. A white colored compound formed was collected, filtered, recrystallized using ethanol, dried and characterized [29].

3. Results and discussions

3.1. Geometry optimization

The optimized geometry of synthesized ligand SBL (Figure 1) was done with B3LYP/6-311+G(d,p) level of accuracy. C10-N8 bond length is found to be 1.452\(\text{Å}\) and C7-N8 bond length is about 1.271\(\text{Å}\). C10-N8 bond length is slightly greater than C7-N8, because double bonds have shorter bond distances than the single bond. C6-O15 bond length is slightly less than C12-O13. The bond lengths of C6-O15, C12-O13, C3-Br9, C12 = O14 and O15-H25 are 1.368\(\text{Å}\), 1.357 \(\text{Å}\), 1.919\(\text{Å}\), 1.205\(\text{Å}\) and 0.9628\(\text{Å}\) respectively. The bond length of benzene ring is given by C1-C2 is 1.39\(\text{Å}\), C2-C3 is 1.393\(\text{Å}\), C3-C4 is 1.384\(\text{Å}\), C4-C5 is 1.402\(\text{Å}\), C5-C6 is 1.405\(\text{Å}\), C4-H18 and C2-H19 bond lengths are 1.081\(\text{Å}\) and 1.822\(\text{Å}\) respectively. The basic geometry of trigonal planar is 120\(°\). The bond angle of C2-C1-C6 is 120.495\(°\), C2-C6-C1 = 120.30\(°\), C1-C2-C3 = 119.18\(°\), C2-C7-C8 = 120.623\(°\), C2-C12-C13 = 120.454\(°\), C12-C6-C13 = 118.658\(°\), C6-O15-H25 bond angle is 109.94\(°\) whereas the C12-O13-H24 bond angle is 107.43\(°\). C4-O12-H25 angle is slightly greater than C12-O13-H24, because the region with one double bond and single bond has got high electron density so that the double bond causes slightly larger angle
The dihedral angles are H20–C10–N8–H19 and H21–C10–N8–C11 which are 3.5539° and 0.5856° respectively.

### 3.2. FT-IR spectra

The IR spectra of the synthesized Schiff base SBL (Figure 2) exhibits strong absorption. The experimental value of νC=O appears at 1669 cm⁻¹. The intense band of νC=N were observed at 1609 cm⁻¹ [29]. νC–H stretching is observed at 2916 cm⁻¹. A sharp peak observed at 749.52 cm⁻¹ corresponds to π(C–H) interactions. A strong absorption observed at 1506 cm⁻¹ indicates benzene ring C=C backbone stretching vibration peak. Several peaks observed in the range of 2916-2429.3 cm⁻¹ is due to the presence of aromatic C–H bond. The C–O stretching was observed at 1250-1360 cm⁻¹ with strong bands. The CH=NH groups stretching vibrations appear at 2978.90 cm⁻¹. The peak at 3082 cm⁻¹ that not including hydrogen bonding can be assigned to νOH.

The computed IR intensities of the novel compound SBL is calculated using B3LYP/6-311+G(d,p). A strong and broad peak of OH was obtained at 3208.76 cm⁻¹. The simulated result of SBL shows peak of νC = O at 1650 cm⁻¹. The intense band of νC = N were observed in the stretch at 1606 cm⁻¹ νC–H stretching is observed at 2978 cm⁻¹. The sharp peak obtained at 713 cm⁻¹ is C–H bending. The strong absorption at 1461-1442 cm⁻¹ due to the presence of benzene ring C=C stretching vibrational absorption peak. C–O stretching appears at 1211-1300 cm⁻¹. The CH = N group stretching vibrations appear at 2978.90 cm⁻¹. The peak at 3082 cm⁻¹ that not including hydrogen bonding can be assigned to νOH.

### Table 1. Thermochemical data from DFT analysis.

| Property                  | Value                  |
|---------------------------|------------------------|
| Thermal energy            | 126.533 kcal/mol       |
| Heat Capacity (CV)        | 53.567 Cal/mol-Kelvin  |
| Entropy (S)               | 129.711 Cal/mol-Kelvin |
| Electronic Energy (EE)    | 3241.852 Hartree        |
| Zero-point Energy Correction | 0.186738 Hartree       |
| Thermal Correction to Energy | 0.201643 Hartree       |
| Thermal Correction to Enthalpy | 0.202567 Hartree       |
| Thermal Correction to Free Energy | 0.140957 Hartree       |

**Figure 11.** HOMO and LUMO of SBL.

**Figure 12.** Ligand docked into the active site of B chain of 3GEY protein.
C–Br stretching is present at 634.39 cm\(^{-1}\). A strong & broad peak of OH is obtained at 3210.81 cm\(^{-1}\) \cite{32}. Experimentally observed FT-IR spectra of the title compound is in good agreement with that of the theoretical value.

### 3.3. UV-VIS spectra

The experimental and theoretical absorbance wavelength of the title compound SBL (Figure 3) were noted. The maximum absorbance wavelength of SBL was measured to be 307 nm experimentally. UV-Vis spectra were computed with TD-DFT. The theoretical maximum absorption wavelength is 305 nm which indicates the excitation of electrons of azomethine group \cite{33}. This is attributed to the azomethine group’s π-∗π and n-∗π transitions \cite{34}. Thus, experimentally determined absorption wavelength of SBL agrees well with that of the theoretical value.

### 3.4. High resolution mass spectra

The mass spectral fragmentation of the synthesized Schiff base SBL (Figure 4) can be studied using mass spectrometry. SBL was characterized using HR-MS spectra. Since one Bromine atom present in the synthesized compound causes two peaks in the molecular ion region i.e.; M+ and
M+2. M+ = 272.008 and M+2 = 274.013. The molecular weight of the synthesized compound C10H10O3NBr is 272.008 g/mol and the base peak is at m/z = 199.007(C7H5OBrN⁺).

3.5. Raman spectra

The synthesized Schiff base SBL (Figure 5) was checked for Raman vibrations calculated using B3LYP/6-311+G (dip) functional. Bands at 1634 cm⁻¹ and 1707 cm⁻¹ correspond to C=O stretching frequencies respectively. Bands at 3082, 3054, 3045 cm⁻¹ correspond to C–H stretching modes of the phenyl ring and the sharp peak obtained at 713 cm⁻¹ corresponds to out of plane π (C–H) interactions. Then a strong absorption at 1493-1403 cm⁻¹ is due to the presence of benzene ring C=C stretching band. C–O stretching frequency were shown by bands in the region 1211-1038 cm⁻¹. The signals near δ160.6 (C6)-174(C13) ppm due to azomethine carbon atoms.

3.6. ¹³C & ¹H NMR spectra

For the title compound, the experimental ¹H and ¹³C NMR was measured using a 400 MHz Varian Joel instrument using CDCl₃ as the solvent. Experimentally measured ¹H NMR spectra of SBL showed two sharp singlets which were interpreted for OH δ at 10.918 (s, 1H, OH) due to the carboxylic part of OH and δ 11.61 (s, 1H, OH) due to the aromatic OH proton [23]. ¹H NMR (δ, ppm, CDCl₃, 400 MHz) 7.66 (s, 1H, CH), 7.605 (s, 1H, CH), 7.43 (s, 1H, CH), 3.6 (t, 2H, CH₂), 2.73 (t, 2H, CH₂). The sharp singlet in 7.66 ppm attributed to azomethine(-HC=N-) proton.

Theoretical, ¹H NMR values were δ 11.09 (s, 1H, OH), 9.79 (s, 1H, OH), 7.58 (s, 1H, CH), 7.600 (s, 1H, CH), 8.41 (s, 1H, CH), 3.30 (t, 2H, CH₂), 2.66 (t, 2H, CH₂); ¹³C NMR values were δ 118 (C₁), 129(C₂), 132.9 (C₃), 139.8 (C₄), 160.6 (C₆), 155.5 (C₉), 66 (C₁₁), 38 (C₁₂), 174 (C₁₃). The differential absorption of left and right circularly polarized light linked with the electronic transitions of chromophores present in a
molecule is referred to as Electronic Circular Dichroism (ECD). The ECD spectrum of the title compound is found near UV region 309.20 nm (Figure 6), indicating the existence of an aromatic chromophore in the compound, according to DFT analysis the absorbance reported by their UV-Vis spectra at 307 nm is similar to this peak of absorption at 309.20 in the ECD spectrum.

ECD is extended from the UV to the IR area of the electromagnetic spectrum by vibrational circular dichroism (VCD) (Figure 7). The C=N stretching vibration is represented by a peak at 1616 cm\(^{-1}\) in the estimated VCD spectra. The absorption peak of vC = O in VCD bands is 1669 cm\(^{-1}\). The presence of a benzene ring C=C backbone stretching vibration absorption peak could explain the significant absorption at 1678 cm\(^{-1}\). At 3046 cm\(^{-1}\), the VCD spectrum displays a substantial C-H stretching. All of these estimated VCD spectra results are in good agreement with the IR spectral results.

### 3.8. Mulliken charge

Mulliken charges are obtained from Mulliken population analysis and estimates partial atomic charges, based on the linear combination of atomic orbitals and molecular orbital techniques. The Mulliken charge is derived from the total chemical charge of the molecule and the atomic charges. The Mulliken charges are calculated using the equation:

\[
q_i = \frac{1}{n} \sum_{j=1}^{n} n_{ij} \mu_{ij}
\]

where \(q_i\) is the Mulliken charge of atom \(i\), \(n_{ij}\) is the number of electrons in orbital \(j\) of atom \(i\), and \(\mu_{ij}\) is the coefficient of orbital \(j\) on atom \(i\).

### 3.9. Non-linear optical properties (NLO)

NLO properties are derived from the interaction of electromagnetic fields and molecules in the molecule which results in altered frequency, phase, amplitude and propagation features, resulting in non-linear visual phenomena [29]. In the DFT level, the electronic dipole moment, molecule polarizability, first and second-order hyperpolarizability values can be calculated to evaluate NLO property. The polarizability, hyperpolarizability, and dipole moment of the title chemical were determined using Gaussian program, with polarizability being 166.131 au. The dipole moment was found to be 3.580 Debye, and these findings clearly show that the examined molecule has nonlinear optical behavior [36].

### 3.10. Molecular electrostatic potential (MEP)

The MEP surface picturizes the reactivity against positive or negative reagents and attributes structure-activity relation of the system [37]. There exists a correlation between electrostatic potential and the dipole moment, electronegativity, and partial charges [38]. Near the phenyl ring, the greatest positive region is located, and zero potential spans over the remaining moieties. The strongest attraction is indicated by blue region, and the strongest repulsion is in red region. The blue colored positive regions are susceptible to nucleophilic reactivity and red negative regions were attributed to electrophilic reactivity. Negative potential is attributed to the presence of lone pairs in the electronegative atoms. In SBL, the negative region is localized over C=N and positive region near the phenyl group. The remaining surface is encapsulated by zero potential as shown in Figure 9. The difference in electron density between the first excited state from the total CI density matrix from TD-DFT calculation and ground state electron density from total SCF density is mapped onto the SCF density shown in Figure 10. The Blue region indicates positive values of difference in electron density where the ground state density is smaller than the excited state density. The red region indicates the reverse. Here in the present system electron density moves from phenyl moiety to C=N moiety as transition happens from the ground state to first excited state.

### 3.11. Frontier molecular orbital analysis (FMO)

The energy values of the HOMO and LUMO orbitals are very important in quantum chemistry with the HOMO being the valence orbital that gives out electrons, whilst the LUMO accepts electrons [39].

The electron in HOMO is more concentrated in the aromatic side and C=N bond, whereas in LUMO (Figure 11), the electron moves towards the aromatic side with Br atom and OH group the HOMO electron move away from the aromatic to aliphatic side. The energy gap is the difference in energy between the HOMO and LUMO orbitals, and it is an essential structure stability reflector [40]. In terms of charge transfer in compounds, a smaller HOMO-LUMO gap suggests that the molecule is less stable [41]. The conjugated molecules have a small HOMO-LUMO gap, which is due to a high degree of intramolecular charge transfer from the electron donor to the acceptor groups [30, 42]. The HOMO and LUMO energy values, in this case, are -0.24019 and -0.07065 eV, respectively. If \( E_{\text{HOMO}} = -0.24019 \text{ eV} \) can be used to express the ionization energy and electron affinity. The hardness and chemical potential are calculated using the equations (I-A)/2 and (I + A)/2, where I and A are the chemical species’ initial ionization potential and electron affinity, respectively [43]. The HOMO-LUMO energy gap is 4.613454664 eV for the title compound, with Ionization potential \( I = E_{\text{HOMO}} = -0.24019 \text{ eV} \), Electron affinity \( A = E_{\text{LUMO}} = -0.07065 \text{ eV} \), Global hardness = -0.5 eV, and Chemical potential = -0.08477 eV [44].

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### Table 2: Swiss ADME tool data of two ligands.

| Properties          | C_{3}H_{10}BrNO_{3}(SBL) | C_{7}H_{14}N_{2}O_{2}(native ligand) | A drug like property exhibiting a range |
|---------------------|---------------------------|--------------------------------------|----------------------------------------|
| **Physicochemical** |                           |                                      |                                        |
| 1. Molar refractivity| 61.01                     | 88.90                                | 40–130                                 |
| 2. TPSA             | 69.89 A\(^{0}\)           | 65.20 A\(^{0}\)                     | 20–130                                 |
| **Lipophilicity**   |                           |                                      |                                        |
| 1. LOGP             | 1.68                      | 2.01                                 |                                       |
| 2. xLOGP            | 1.33                      | 1.71                                 |                                       |
| 3. vlLOGP           | 2.05                      | 1.99                                 | Lipophilic compound                    |
| **Pharmacokinetics**|                           |                                      |                                        |
| 1. Log \(k_{p}\)    | -7.02                     | -6.89                                | More value high skin permeability      |
| **Drug likeness**   |                           |                                      |                                        |
| 1. Bioavailabilityscore | 0.85                     | 0.55                                 | Amount of dose get human body          |
| **Medicinal chemistry** |                         |                                      |                                        |
| 1. synthetic accessibility | 2.30                     | 2.12                                 | <5 Easy synthesis                     |
|                     |                           |                                      | >5 hard to synthesis                   |
|                     |                           |                                      | Range 1-10                             |
3.12. Thermochemistry

Thermodynamic property of the synthesized Schiff base (E)-3-((5-bromo hydroxy benzylidene) amino) propanoic acid obtained through DFT calculations are shown in Table 1.

4. Molecular docking

Synthesized Schiff base SBL were carried out for the molecular docking to measure the inhibition tendency of the target cancer protein 3GEY (Figure 12) which is the variant of PARP-15 [45]. Auto Dock vina v1.2.0 was employed for molecular docking studies. For in silico screening, molecular docking is a functional tool in the arsenal of drug design experiments [46, 47]. Docking is a computer approach for finding a molecule that fits the protein's biologically active binding site energetically and geometrically [48, 49]. Molecular docking has proven to be an effective approach for discovering new drugs that target proteins. Protein-ligand docking refers to search for the most error-free and stable ligand conformation or pose with a protein on a specific binding site [50, 51]. For molecular docking, the target protein PDB file 3GEY is obtained from RCSB-Protein data bank and SBL ligand was saved in PDB format from Gaussian 16W. The simulation of protein-ligand binding interactions is done through Autodock tools (v. 1.5.6) and Auto Dock Vina. For preparing protein, water molecules were removed followed by adding polar hydrogens and Kollman charges. Auto grid determines the position of native ligand on binding site on B chain of target protein by arranging grid coordinates. In docking process, the coordinates of the grid box shall be validated for ensuring for the ligands to bind on binding site in the accurate conformation. Using Auto Dock Vina biologically active conformations are simulated and lower energy scores demonstrate the best protein-ligand binding affinity. Visualization of docked poses of SBL with the protein target was done using BIOVIA Discovery Studio visualizer 2021 (64 bit) client.

Figures 13 and 14 visualizes 2D interaction diagram of the best pose for the native ligand of protein and SBL respectively. Figure 14 provides protein-ligand (SBL) interaction in LigPlot. LigPlot V.4.5.3 is a tool which creates graphical representation of protein-ligand interaction obtained from the three-dimensional coordinates incorporated in a pdb file of docked system (Figure 15). The interaction involves H-bonds designated as dashed lines and hydrophobic interactions indicated by Arc with Spokes directing towards contracting ligand atoms [52]. A total of ten amino acids are involved in ligand binding pocket of receptor for the native ligand with prominent interactions while 13 amino acids contribute to SBL ligand binding process.

The affinity of SBL ligands with receptor PARP-15 (pdb:3GEY) is given by prominent interactions to TYR 582, TYR 569, LEU 637, SER 577, GLY 538, THR 581, ALA 571 and these are the active residues in 3GEY target protein. Native ligand C17H17N3O3 illustrate HIS 537, LEU 637, TYR 569, GLY 538, THR 576, TYR 582, ALA 571 and which indicate two different ligands have mostly similar active residues. There are two key residues that have hydrogen bonds SER577, GLY 538 which restricted ligand in the binding domain. C-N bond interacts with an amino acid residue (GLY 538) in both cases. Compare to native ligand, SBL has extra three residual interactions with bromine atom they are TYR 569, LEU 637, TYR 582. Here the native ligand is replaced with SBL is the easy process they docked in to the active site of 3GEY protein by key and lock method. So, at this time we can conclude that biological features of native ligand are similar to SBL, SBL being lower molecular weight system compare to native ligand and therefore, easy to trap in to the cavity. The comparison chart reveals that some properties are favorable to SBL or comparable to that of native ligand which means SBL have potential drug likeness (Table 2). Here lipophilicity values are very much less than 5 in case of SBL which shows high Lyophilic character. Based on pharmacokinetics, more negative is the value has high skin permeability. Here SBL has got higher negative value of -7.02 where native ligand has got a value of -6.89 which mean that synthesized Schiff base shows high permeability through skin. The molecular weight of SBL is lesser than the native ligand and method of synthesis lot more easily than native ligand. From all these comparisons we can conclude that the SBL is way much better than the native ligand.

5. Conclusion

Bioinorganic chemistry is a prosperous field of drug research for cancer treatment so this Schiff base can be studied invitro study against PARP. Breast cancer is the most frequently detected cancer which accounts for 23% of the cancer cases and 14% of the cancer deaths. Over the past few years, Schiff bases find applications in medicine having multi-dimensional properties. Present study involves the synthesis of a Schiff base (E)- 3-((5-bromo-2-hydroxybenzylidene) amino) propanoic acid using 5-bromosalicylaldehyde and β-alanine. FT-IR, UV-Visible, 1H & 13C NMR and mass spectrum studies were used to characterize the synthesized compound. FT-IR spectrum indicates a strong band in 1609 cm⁻¹ which indicates C=O bond. The maximum absorption in the UV-Vis spectrum is found at 307 nm. A molecular ion at m/z = 274 was observed in the mass spectrum, which matches to the molecular weight of the synthesized Schiff base with the molecular formula C10H10O2NBr. The formation of the proposed compound was supported by 1H and 13C NMR. The experimental values are found to be in good agreement with theoretical values derived from DFT analysis. DFT analysis were carried with B3LYP/6-31+G(d, p), and the optimized geometry of the system was reported. Mulliken atomic charges and HOMO-LUMO band gaps were also determined. In this study, we aim to investigate how 3GEY, the variant of PARP-1 protein has high potent binding affinities between native ligand and SBL and comparison of interaction between ligand functional group with amino acid residues in protein employing Auto Dock tools. Visualization of docked pose of new Schiff base ligand shows promising interactions with active site residues of target protein. The drug likeness properties reported by using Swiss ADME tool also point towards a promising inhibitor candidate. The synthesized Schiff base, SBL can be further extended for its complex formation with different transition metal ions for enhanced biological activity.

Declarations

Author contribution statement

Meenukutty M. S, Arsha P Mohan: Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Vidya V. G, Viju Kumar V. G: Conceived and designed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

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Data availability statement

Data will be made available on request.

Declaration of interests statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.
