In Silico Molecular Docking on Bioactive Compounds from Indian Medicinal Plants against Type 2 Diabetic Target Proteins: A Computational Approach

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Natural chemical compounds from medicinal plants used for the healing of diseases and disorders with fewer side effects, easy accessibility and economically cheap. The current study was aimed at finding novel drug like molecules as anti-diabetic compounds using in silico approach. Intermolecular interactions between target proteins and different antidiabetic compounds were observed. Five phytocompounds were selected from Plumbago zeylanica, Neolitsea cassia and Wrightia tinctoria and taken for molecular docking against human pancreatic alpha-amylase and human dipeptidyl peptidase IV using Autodock 4.2. Among the five phyto compounds, 6 urs-12-en-24-oic acid Plumbago zeylanica is the best compound for both the human pancreatic alpha-amylase and human dipeptidyl peptidase IV inhibition, as it possessed higher value in molecular dockings.

Key words: Molecular docking, in silico study, Autodock, alpha-amylase, dipeptidyl peptidase IV

Type 2 Diabetes Mellitus (T2DM) is a critical metabolic failure characterized by less insulin action and high blood glucose level[1]. T2DM is signified as the quickest worldwide threat to people’s health[2] and it causes blindness, lower limb amputation and kidney disease[3]. As stated by WHO, 350 million people may be affected in 2030 globally, if the action has not been taken against T2DM[4]. Alpha (α)-Amylase and Dipeptidyl Peptidase IV (DPP IV) inhibitors used to protect the digestion of starch and other carbohydrates with T2DM patients[5].

α-Amylase universally distributed throughout the animal, plant and microbial kingdoms. Many mammals secrete two types of α-amylase, one is salivary α-amylase by the parotid gland and another one is pancreatic α-amylase secreted by the pancreas[6].

α-Amylase inhibitors extensively used in the therapy of T2DM. DPP IV inhibitor has the potential to be a novel, efficient and considerable agent to treat T2DM[7]. The usage of DPP IV inhibitors has fewer side effects like hypoglycemia, increasing body weight and gastrointestinal tract disorders[8]. The research of oral glucose tolerance tests on animals revealed that genetic deletion of DPP IV has increased the glucose tolerance and the secretion of insulin[9]. It encourages the study of α-amylase and DPP IV inhibitor bioactive compounds from medicinal plants which have fewer side effects, low cost and are easy to obtain.

Numerous drugs such as meglitinides, biguanides, sulfonylureas, thiazolidinediones, α-amylase inhibitors, DPP IV inhibitors used to treat the T2DM[10]. The side effects of these enzymatic inhibitors are flatulence, low blood sugar and hepatitis[11,12]. In recent years, a variety of research has done on plants[13-19]. Besides, several herbs and fruits have the property of inhibitors[16,17].

The known natural inhibitors of digestive enzymes include phytocompounds which operate through various mechanisms[17]. Today, researchers emphasise primarily on the finding of practical and low side effect therapeutic drugs to treat T2DM[20]. Medicinal plants produce secondary metabolites that have effective antidiabetic potentials to regulate blood glucose level. In conventional treatment, several medicinal plants...
were used such as *Plumbago zeylanica* (*P. zeylanica*), Neolitsea cassia (*N. cassia*) and Wrightia tinctoria (*W. tinctoria*) that avoid difficulties in the control of T2DM.

Molecular docking is an essential method in the improvement of new medications. Docking strategy permits describing the conduct of a test molecule in the coupling site of the receptor target of interest. A docking method must have the ability to predict the useful binding strength between the ligand and the receptor complex[21]. In computational chemistry, *in silico* analysis has led to the discovery of new active drugs with fewer side effects[22]. Molecular docking analysis has directed to predict the binding attractions of different bioactive compounds and to explain specific sites of interaction between the bioactive compounds and the target proteins[23]. The aim of this *in silico* study was to explore the antidiabetic properties of phytocompounds derivatives namely 6 urs-12-en-24-oic acid, Beta (β)-Amyrin, 10,12-Docasadiyndioic acid, 1-(Ethynyltrimethylsilyl-2)-1-Chloro-2-Ethyl-2-methyl cyclopropane and androstane via molecular docking using Autodock software.

**MATERIALS AND METHODS**

**Selection of receptor:**

The Three Dimensional (3D) structure of the receptor binding sites of human Pancreatic α amylase (3OLD) (fig. 1a) and human DPP IV (5Y7H) (fig. 1b) retrieved from the Protein Data Bank (PDB) (http://www.rcsb.org).

**Selection of ligand:**

The 3D structures of bioactive compounds namely 6 urs-12-en-24-oic acid (fig. 2a) and β-amyrin (fig. 2b) from *P. zeylanica*, 10,12-Docasadiyndioic acid (fig. 2c) and 1-(Ethynyltrimethylsilyl-2)-1-Chloro-2-Ethyl-2-methyl cyclopropane (fig. 2d) from *N. cassia* and androstane (fig. 2e) from *W. tinctoria* collected from Pubchem, a compound database. The collected Structure Data File (SDF) files of identified bioactive compounds from the PubChem database were converted into PDB format using the EduPymol version 1.7.4.4[24].

**Protein ligand docking:**

Molecular docking analysis performed to evaluate the most preferred geometry of the protein ligand complex. Docking phase is meaningless without its two components such as target protein and ligand. Human pancreatic α amylase (3OLD) and human DPP IV (5Y7H) used for performing docking study as target proteins (fig. 1). Docking results identify native or native-like configurations of the protein-ligand complex. The selected proteins complex used after removal of already bonded ligand and water molecules. Docking was performed by AutoDock 4.2 software. The complete docking steps could be stated as follows: first of all, the water molecules were eliminated from the protein. After the removal of water molecules, the PDB
file of 3OLD and 5Y7H which provided as an input to the software. Kollman charges were computed for the macromolecule by AutoDock. Then the macromolecule was checked for the missing atoms and repaired. After repairing missing atoms, the hydrogens were added by keeping all the parameters at default settings. The macromolecule after all these modifications was saved as .PDB in the same directory. Then the ligand preparation was carried out. Like macromolecule, Kollman and Gasteiger charges were computed for the ligand.[25] Some of the torsions of the ligands were defined. The root was detected; the rotatable bonds were converted into non-rotatable bonds and vice versa. Also the number of active torsions was to most atoms rather than fewest. A Protein Data Bank, Partial Charge (Q) and Atom Type (T) (PDBQT) file was then created for the modified ligand with extension .PDBQT. After the preparation of a macromolecule and ligand, the rigid residue was prepared using the grid module provided in the AutoDock 4.0 Grid module employed .PDB file. This program was run using a searching grid extended over ligand molecules with box spacing 70×70×60; spacing was 0.608, x, y, z coordinates 8.485, -5.766, 15.737 respectively for 3OLD; 80×80×80, spacing was 0.508, x, y, z coordinates 52.271, 58.538, 45.819 respectively for 5Y7H while other parameters were default. The flexible macromolecule was saved with the .PDBQT extension. For molecular docking AutoDock 4.2 software was used.

It employed a configuration file referring to PDBQT files of macromolecules and compounds prepared using AutoDock 4.2 and grid properties. As an output AutoDock 4.2 generated log files and PDBQT files of energy models for the selected data set. The output file contained different energy models. Among these models, the lowest energy model against each ligand was selected and appended at the end of the original protein file. As a result of this step, docked files for the selected set were generated. For the interpretation of docking results; target protein and protein docked with the data set of compounds, the interactions between the active pocket of protein and compounds were found. The best docking poses were predicted to be the most stable conformation of each compound for binding to the protein active site. Consequently, the output of the docking process was analyzed utilizing EduPyrmol version 1.7.4.4.

RESULTS AND DISCUSSION

All the identified compounds (6 urs-12-en-24-oic acid (fig. 2a) and β-amyrin (fig. 2b) from *P. zeylanica*, 10,12-Docosadiyndioic acid (fig. 2c) and 1-(Ethynyltrimethylsilyl-2)-1-Chloro-2-Ethyl-2-methyl cyclopropane (fig. 2d) from *N. cassia* and androstane (fig. 2e) from *W. tinctoria*) were docked into that active site of human pancreatic α amylase (fig. 1a) and DPP IV (fig. 1b) by using AutoDock 4.2. The binding affinities of these compounds to target proteins are shown in Table 1 and Table 2 and fig. 3 and fig. 4. The identified five compounds have higher binding interaction compared to the co-crystallized ligand of 3OLD and 5Y7H were found lying deep into the binding cavity of target proteins showing all the significant interaction and favorable energy of interaction ranging from -3.55 to -10.58 kcal/mol and -4.33 to -9.45 kcal/mol respectively (Table 1 and Table 2). The bonded and non-bonded interactions of these compounds are shown in Table 1 and Table 2. Among

![Fig. 2: Chemical structure of identified phytocompounds investigated, (a) 6 urs-12-en-24-oic acid from *P. zeylanica*; (b) β-Amyrin from *P. zeylanica*; (c) 10,12-Docosadiyndioic acid from *N. cassia*; (d) 1-(Ethynyltrimethylsilyl-2)-1-Chloro-2-Ethyl-2-methyl cyclopropane from *N. cassia* and (e) Androstane from *W. tinctoria*](image-url)
| Parameters                          | 6 urs-12-en-24-oic acid from P. zeylanica | β-Amyrin from P. zeylanica | 10,12-Docosadiynidoic acid from N. cassia | 1-(Ethynyltrimethylsilyl)-2-Chloro-2-Ethyl-2-methyl cyclopropane from N. cassia | Androstane from W. tinctoria |
|-----------------------------------|------------------------------------------|-----------------------------|-------------------------------------------|-----------------------------------------------------------------------------|-----------------------------|
| Binding energy (Kcal/mol)         | -10.58                                   | -8.9                        | -3.55                                     | -4.3                                                                       | -7.58                       |
| Ligand Efficiency                 | -0.31                                    | -0.29                       | -0.14                                     | -0.33                                                                      | -0.4                        |
| Inhibition Constant               | 17.49nM                                  | 297.25nM                    | 2.52mM                                    | 705.84µM                                                                   | 2.77 µM                     |
| Intermolecular Energy             | -11.48                                   | -9.2                        | -9.81                                     | -5.19                                                                      | -7.58                       |
| vDW+Hbond+desolv Energy (Kcal/mol)| -11.28                                   | -9.05                       | -8.7                                      | -5.16                                                                      | -7.58                       |
| Electrostatic Energy              | -0.19                                    | -0.15                       | -1.11                                     | -0.03                                                                      | 0.0                         |
| Total Internal Energy             | 0.01                                     | 0.07                        | -0.52                                     | -0.37                                                                      | 0.0                         |
| Torsional Free Energy             | 0.89                                     | 0.3                         | 6.26                                      | 0.89                                                                       | 0.0                         |
| RMSD                              | 30.14                                    | 31.87                       | 27.89                                     | 31.01                                                                      | 30.29                       |
| Number of Hydrogen bonds          | 1                                        | 2                           | 3                                        | 0                                                                          | 0                           |
| Hydrogen bond Interaction         | Lys200-NZ: O31-Ligand                     | Asp197-OD1: H32-Ligand and Arg195-NH2: O31-Ligand | Ala106-N: O27-Ligand and Lys200-NZ:O15-Ligand | NA                                                                         | NA                          |
| Hydrogen Bond Length (Å)          | 3.2                                      | 1.9 and 3.2                 | 3.0, 3.0 and 2.6                          | NA                                                                         | NA                          |

**TABLE 2: MOLECULAR DOCKING STATISTICS OF THE DPP IV AND IDENTIFIED PHYTOCOMPOUNDS COMPLEX**

| Parameters                          | 6 urs-12-en-24-oic acid from P. zeylanica | β-Amyrin from P. zeylanica | 10,12-Docosadiynidoic acid from N. cassia | 1-(Ethynyltrimethylsilyl)-2-Chloro-2-Ethyl-2-methyl cyclopropane from N. cassia | Androstane from W. tinctoria |
|-----------------------------------|------------------------------------------|-----------------------------|-------------------------------------------|-----------------------------------------------------------------------------|-----------------------------|
| Binding energy (Kcal/mol)         | -9.45                                    | -8.81                       | -4.41                                     | -4.30                                                                       | -8.06                       |
| Ligand efficiency                 | -0.28                                    | -0.28                       | -0.17                                     | -0.33                                                                      | -0.42                       |
| Inhibition constant               | 117.59 nm                                | 350.78 nm                   | 582.5 µm                                  | 672.36 µm                                                                  | 1.24 µm                     |
| Intermolecular energy             | -10.35                                   | -9.1                        | -10.68                                    | -5.22                                                                      | -8.06                       |
| vDW+Hbond+desolv energy (Kcal/mol)| -10.2                                    | -9.01                       | -9.32                                     | -5.29                                                                      | -8.06                       |
| Electrostatic energy              | -0.15                                    | -0.09                       | -1.35                                     | 0.05                                                                       | 0.0                         |
| Total internal energy             | 0.08                                     | 0.07                        | -1.05                                     | -0.31                                                                      | 0.0                         |
| Torsional free energy             | 0.89                                     | 0.3                         | 6.26                                      | 0.89                                                                       | 0.0                         |
| RMSD                              | 86.09                                    | 89.38                       | 83.73                                     | 84.55                                                                      | 86.88                       |
| Number of hydrogen bonds          | 2                                        | 2                           | 6                                        | 0                                                                          | 0                           |
| Hydrogen bond interaction         | TRP627-NE1: O33Ligand and TRP563-CA: O31Ligand |                      | SER212-OG: O31-Ligand and Leu214-O: H32-Ligand | Arg471NH1: O14Ligand, Arg471NE: O15 Ligand, Ser458OG: O14 Ligand, Arg358O: H28Ligand, Arg356N: O27Ligand and Phe357N: O26Ligand | NA                           |
| Hydrogen bond length (Å)          | 2.9 and 2.7                              | 2.5 and 1.8                 | 2.9, 3.3, 2.7, 2.1, 3.1 and 2.8           | NA                                                                         | NA                          |

Molecular docking is now widely used to discover new ligands for the target of known structure. The free energy binding score denotes the interactions of ligand to the target protein; the lower free energy binding, the
higher binding affinity\textsuperscript{[26]. Also, inhibition constant can be calculated using computational methods. The lowest inhibition constant denotes the most active compound. It signifies the molecular interaction between ligands and the target protein. The high value of surface interaction indicates the possibilities of compounds binding with the target protein\textsuperscript{[27].}

Based on the\textit{ in silico} analysis, 6 urs-12-en-24-oic acid of \textit{P. zeylanica} (fig. 3a and fig. 4a) has the lowest value (-10.58 Kcal/mol for \textalpha-amylase and -9.45 Kcal/mol for DPP IV) in binding free energy while \textbeta-amyrin of \textit{P. zeylanica} (fig. 3b and fig. 4b) and androstane of \textit{W. tinctoria} (fig. 3c and fig. 4c) were in the second (-8.9 Kcal/mol for \textalpha-amylase and -8.81 Kcal/mol for DPP IV) and third position (-7.58 Kcal/mol for \textalpha-amylase and -8.06 Kcal/mol for DPP IV) with two target proteins (Table 1 and Table 2). 1-(Ethynyltrimethylsilyl-2)-1-Chloro-2-Ethyl-2-methyl cyclopropane from \textit{N. cassia} have shown the fourth (-4.3 Kcal/mol) and fifth (-3.55 Kcal/mol) positions for \textalpha-amylase respectively and fifth (-4.33 Kcal/mol) and fourth places (-4.41 Kcal/mol) for DPP IV respectively when compared with binding energies with other ligands (Table 1 and Table 2). Hydrogen bonds are a significant factor in defining the specificity of ligand binding\textsuperscript{[28]. 6 urs-12-en-24-oic acid of \textit{P. zeylanica} ranked highest in binding affinity with 3OLD and 5Y7H using a hydrogen bonds.
bond (Table 1 and Table 2). Lys200 of 3OLD formed one hydrogen bonds with 6 urs-12-en-24-oic acid of \( P. zeylanica \) between NZ of the residue and hydroxyl (O31) of the ligand played in binding affinity with a bond length of 3.2 Å. The TRP627 and TRP563 of 5Y7H active site residues formed two hydrogen bonds with O31and H32 of 6 urs-12-en-24-oic acid with the length of 2.9 and 2.7Å. β-amyrin from \( P. zeylanica \) formed two hydrogen bonds with both the target proteins.

β-amyrin and 3OLD complex formed hydrogen bonds by Asp197-OD1 and Arg195-NH2 of 3OLD bonded with H32 and O31 of Ligand respectively. Their hydrogen bond length was 1.9 and 3.2Å. β-amyrin and 5Y7H complex also formed through two hydrogen bond: SER212-OG and Leu214-O of 5Y7H with O31and H32 of ligand and the length were 2.5 and 1.8Å respectively. Similarly, 10,12-Docasadieniodio acid from \( N. cassia \) formed three hydrogen bonds with 3OLD and 6 hydrogen bonds with 5Y7H (Table 1 and Table 2). Though, 1-(Ethynyltrimethylsilyl-2)-1-Chloro-2-Ethyl-2-methyl cyclopropane from \( N. cassia \) (fig. 3d and fig. 4d) also showed good binding score but not interacted by hydrogen bonds with two target proteins (Table 1 and Table 2). It interacted through hydrophobic, electrostatic and Van der Waals interactions with both the target proteins.

Likewise, androstane from \( W. tinctoria \) (fig. 3e and fig. 4e) also is not interacted through hydrogen bond and electrostatic force; it is related to hydrophobic and Van der Waals interactions (Table 1 and Table 2). The lowermost value of free binding energy generates a useful binding molecule and then produces a potent biological activity. Free energy binding and surface interaction between ligands and target protein influence the inhibitory activity of selected bioactive compounds on α-amylase and DPP IV. The \textit{in silico} analysis exposed the prominence and presence of oxygen, nitrogen and carbonyl groups in the compound structure was essential and amplified the tendency of hydrogen bond interactions\textsuperscript{29}. Molecular docking studies are broadly exercising methods to analyze the promising drugs in the pharmaceutical industry. The binding pose of ligands or bioactive compounds to their target proteins divulges their possible affinity and activity of drugs.

The current study was aimed at finding novel drug like molecules as antidiabetic compounds using \textit{in silico} approach. Intermolecular interactions between target proteins and different antidiabetic compounds were observed. The protein-ligand interaction shows that 6 urs-12-en-24-oic acid from \( P. zeylanica \) having the best interactions compared to the other identified compounds from different medicinal plants toward the enzymes human pancreatic α-Amylase and human DPP IV. From the above observations, we can assume that \( P. zeylanica \) can be a prominent resource for antidiabetic compounds.

**Conflict of interests:**

The authors declared no conflicts of interest.

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