Modeling and Docking of Cysteine Protease-A (CPA) of Leishmania donovani

Sindhuprava Rana¹*, Jay Prakash Mahato¹, Maneesh Kumar², Surendra Sarsaiya¹
¹Department of Biotechnology, Sri Satya Sai University of Technology & Medical Sciences, Bhopal, Madhya Pradesh, India. ²Department of Biotechnology, College of Commerce, Arts & Science, (Magadh University, Bodh Gaya), Patna-800020, Bihar, India.

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ABSTRACT

Leishmaniasis is a tropical disease and causes severe public health issues. The disease caused by parasitic protozoan Leishmania species. Many people around the globe are affected in dramatic way, which would be fatal if untreated. Drugs have been made that become resistant to present strains. Cysteine proteases (CPs) are common protein degrading cysteine protease enzymes present in Leishmania donovani. In present scenario, to achieve effective enzyme inhibition mechanism, it should be taken into consideration to develop some effective and novel inhibitors to block the binding sites of that protein. Homology modeling of cysteine protease has been done and docked with suitable inhibitors by using various servers and computational tools. The model was designed, compared and validated by DOPE and Verify 3D scores by using DSv3.5. Licochalcone-a alone showed 37 LibDock conformations with 6 different poses were suitably docked at the site 1. The study would help to design the protein with new drugs in respect of resistant one for the treatment of harmful visceral leishmaniasis.

Abbreviations: CHARMM: Chemistry at HARvard Macromolecular Mechanics; CPs: Cysteine proteases; DOPE: Discrete Optimized Protein Energy; DSv3.5: Discovery Studio 3.5; I-TASSER: Iterative Threading ASSEMBly Refinement; NCBI: National Center for Biotechnology Information; pBLAST: protein Basic Local Alignment Search Tool; PDB: Protein Data Bank; RMSD: Root Mean Square Deviation; VL: visceral leishmania.

INTRODUCTION

Visceral Leishmaniasis (VL) is a chronic tropical disease, responsible for severe public health concern around the world. The parasitic species mainly involve in this infection are belong to protozoan Leishmania donovani (Das et al., 2013). This serious infection is basically mediated by sand flies (Phlebotomus argentipes) that transmit the disease to about 12 million people as only reservoir and is endemic in the tropical regions of Africa, America, the Indian subcontinental and subtropical regions of Southeast Asia (Selzer et al., 1999).

The survival of the sand fly vector depends on many biotic (human/animal blood meals) and abiotic (precipitation and temperature) factors. Such conditional habitat found near Gangetic plains (Ostyn et al., 2015). L. donovani is an active causative agent that causes human tegumentary and VL that also known as kalazar, in local regions. It would be fatal for human survival if not properly diagnosed and treated well on time (Alvar et al., 2012, Silva-Almeida et al., 2012). In past decades, many approached have been to create effective vaccine but not much success would achieve in its cure so far (Ait-Oudhia et al., 2011, Bhattacharya et al., 2016). Cysteine proteases (CPs) are common protein degrading enzymes that cleave various peptides. They are the essential virulence factors in leishmania parasitic. They play key role in many infectious protozoa and viral agents (Mahmoudzadeh-Niknam and McKerrow, 2004; Vicik et al., 2006).
Due to its biological importance it would become the potential target to develop new drug to cure the leishmaniasis (Das and Ali, 2014). CPs can also participate in various other cellular metabolisms like excystment, exsheathing and, some parasite immunoevasion and tissue invasion. Many parasitic CPs have some potential immunogenic properties, which have been exploited for diagnostic purposes and vaccine targets (Sajid and Mc Kerrow, 2002). In the Leishmania, the mechanism of nucleophilic cysteine thiol is involved catalytic mechanism in catalytic dyad. So, the creation of novel inhibitors is the use of electrophilic moieties that covalently bind to the cysteine residue of the active site. The present aim to achieve effective enzyme inhibition, this should be taken into consideration that this electrophile would incorporate into peptidic or peptidomimetic sequences present in the substrate binding sites. In L. donovani, CPs is the virulence factor in human and animals (McKerrow, 1999; Mahmoudzadeh-Niknam and Mc Kerrow, 2004; Rosenthal, 2004).

Many inhibitors are used for docking purpose that potentially inhibits the cysteine protease of L. donovani. Among them licochalcone-A is a novel estrogenic, natural phenolic compound that has antimalarial, anticancer, antitumor and antiviral activities. It also induces apoptosis at some instant that prevents any biological harm to the host cell (Fu et al., 2004). Primin is another naturally organic compound has clear antimicrobial activities (Brondani et al., 2007). Sitamaquine and Miltefosine are the best known drugs used in the treatment of VL through oral route (Sundar and Olliaro, 2007; Loiseau et al., 2011). Amarogentin is an smart organic compound from Gentiana lutea, a herbal bitter is generally used in digestive disorders (Keil et al., 2000). Emetine is an effective anti-protozoal alkaloid and induces vomiting. It is also used in the treatment of severe invasive amoebiasis (Kattwinkel, 1949). Fisetin is the active flavonoid, found in various fruits and green vegetables, generally used against numerous harmful diseases. It is an good anti-inflammatory and antioxidant compound (Khan et al., 2013). In this present work, various tools and softwares have been used to understand in silico biological role of cysteine protease of L. donovani within the affected host. The structured cysteine protease model would give the proper idea about the protein-ligand interaction. The work clears the selection of some novel drugs which might paved the way to design them for the betterment of human society against leishmaniasis.

METHODS AND MATERIALS

3D Structure Prediction and Model Prediction

The amino acid sequence of cysteine protease (accession: AAL09448) of Leishmania donovani has taken from the protein database of NCBI (Sayers et al., 2011). A BLASTP search was done (Altschul et al., 1997) against Protein Data Bank (PDB) (Berman et al., 2000). This protein sequence has used for our homology modeling, computational assessment and to predict the suitable protein-ligand interaction. The 3D structure of the cysteine protease of L. donovani was performed by using various servers online servers like HMM-HMM comparison (HMpred) (Söding, 2005; Remmert et al., 2012), the hierarchical method of protein structure and function (1-TASSER) (Zhang, 2008), the protein structure prediction (Raptor X) (Källberg et al., 2012), the knowledge based approach (Swiss Model) (Arnold et al., 2006).

Protein Stimulation and Active Sites Prediction

The best structure of cysteine protease that was modeled by Swiss model web server, based on the DOPE Score (Shen and Sali, 2006) was selected with lowest value for further study. The actual positions and nature of amino acids within the modeled structure was evaluated by Ramachandran Plot via PROCHECK (Laskowski et al., 1996), ERRAT (Colovos and Yeates, 1993) and Verified 3D (Bowie et al., 1991; Luthy et al., 1992). The protein was also analyzed by verify Protein (Profiles 3D) from DSv3.5. Along with these, other validation like refinement of flexible loop and side chain were done by Discovery Studio version 3.5 (DSv3.5) (Inc, 2012). The refinement was followed by protein stimulation via CHARMM (Brooks et al., 1983) with the help of DSv3.5.

The CHARMM is a dynamic in silico molecular computational program that minimize the energy value during docking while protein has been stimulated under the effect of dielectrics of the atoms of amino acids (Sahoo et al., 2009, Sahoo et al., 2014). The model structure of the cysteine protease with all its computational parameters were further taken for the docking purpose using the same DSv3.5.

Docking

The accurate identification and characterization of binding sites of the modeled cysteine protease of the L. donovani have been identified by using the DSv3.5. The modeled protein was selected and allowed for Protein Preparation under the effect of CHARMM force, followed by cavities identification as active sites where the selected ligands have allowed docking. The three active sites were shown by DSv3.5, in which only the site1 has actively participated for the docking work. Site2 and site3 haven’t shown any docked score. After the protein preparation has been performed, the ligand library was made by using different ligands like licochalcone-A, Primin, Sitamaquine, Miltefosine, Amarogentin, Emetine and Fisetin with their known IC50 value (Ansari et al., 2012) were taken from NCBI PubChem database. The docking were performed by LibDock algorithm inbuild in DSv3.5 (Rao et al., 2007).

RESULTS AND DISCUSSION

The computational analysis using both software and servers of the modeled protein structure enable us the biological properties. The cysteine protease of L. donovani, with the hypothetical protein modeled that was prepared and retrieved from Swiss model tool.
Modeled Structure prediction using homology modeling approaches

The proper template identification after pBLAST, followed by modeled protein quality assessment enables to understand the suitable functions, and localization of the subcellular protein and their interaction with ligands. The most common and approximate prediction method is homology modeling that provides a proper idea about the protein. The homology model of cysteine protease is shown in Figure 1. The figure labeled as sequence alpha (\(\alpha\)), beta (\(\beta\)) and flexible loops (FL). The cysteine protease consists of seven alpha helixes, ten beta sheet, three \(\eta\) (small helixes) and few flexible loops. The secondary sequence was clearly mentioned the actual amino acids residues involved in making alpha helixes, beta sheet, flexible loops and small helixes (with three amino acids residues) [Figure 2]. All the models which were designed from the different servers were compared and validated by DOPE scores by using DSv3.5. The suitable template model that retrieved from the Swiss model has been taken for our study which has the lowest DOPE value of -430.174938 as the best-modeled structure. The model structure was proved by Verify 3D that showed 83.06 % value. The protein verify by using DSv3.5, shows the 111.74 verify score which lies in between verify expected high score (139.723) and verify expected low score (62.8754). The model validation PROCHECK tool was used to determine Ramachandran plot [Figure 3] that assures the quality model. The result of Ramachandran plot showed the amount of amino acid residues i.e. 85% favored regions, 13.1% additional allowed regions, 1.5% generously allowed regions and 0.4% disallowed regions favored represented as a reliable and good quality model. The reliability of the modeled protein was also checked by using ERRAT that showed 83.162 overall model quality [Figure 4].

**Fig 1:** 3D Model Cysteine Protease was produced by DSv3.5 having \(\alpha\) helices are in red, \(\beta\) sheets are in blue and flexible loops are in grey.

**Fig 2:** Secondary Structure of the modeled cysteine protease by DSv3.5.
Protein-Ligand Interaction

After assessment of suitable cysteine protease protein model detecting by different computational tools and servers, some active binding sites of the model cysteine protease were created, then we tried to analyze the specific binding affinity of some suitable substrate ligands that were effectively docked with the 3D modeled structure. There are three different binding sites having charged amino acids, were detected by using receptor cavities protocol of DSv3.5. The Licochalcone-A has a highest LibDock score of 71.728 also showed the absolute energy value of 86.5400 for the binding site 1 (the position value of the site 1: -28.787, 22.2787, 20.1339, 19.6) of the model protein. Licochalcone-A has showed 37 poses of LibDock conformations and 5 hits, in which only 6 different poses were suitably docked at the site 1. Licochalcone-A along with some others ligands which are also perfectly bound at this site 1 are providing the LibDock scores [Table 1], along with their LibDock Score. They are Primin, Sitamaquine, Miltefosine, Amarogentin, Emetine and Fisetin also exhibited high binding activities with the receptor protein. The binding between model cysteine protease protein and ligand Licochalcone-A binding was shown in Figure 5. The subcellular localization of the cysteine protease protein is present within the extracellular regions and some part of periplasmic and cytoplasmic also. The proper Protein-Ligand interaction is shown in Figure 6. Here, the frequently involved amino acid of model protein that forms hydrogen bonds with the ligand is His 81. The group i.e.: ND1 of His 81 interacts with :O4 of Licochalcone-A. Our docking result and the structure analysis suggest that the model protein having active binding site to ligand properly. The study would help to design the new drugs in respect of resistant one for the cure of leishmaniasis.

Table 1: Comparative LibDock Scores of different effective ligands

<table>
<thead>
<tr>
<th>Ligands</th>
<th>LibDock Score</th>
<th>Absolute Value</th>
<th>Relative Value</th>
</tr>
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<tbody>
<tr>
<td>Licochalcone-A</td>
<td>71.728</td>
<td>86.5400</td>
<td>13.3502</td>
</tr>
<tr>
<td>Primin</td>
<td>71.5692</td>
<td>27.9008</td>
<td>09.4004</td>
</tr>
<tr>
<td>Sitamaquine</td>
<td>70.4248</td>
<td>63.1020</td>
<td>17.5402</td>
</tr>
<tr>
<td>Miltefosine</td>
<td>69.6103</td>
<td>27.3418</td>
<td>12.7868</td>
</tr>
<tr>
<td>Amarogentin</td>
<td>68.3701</td>
<td>90.1020</td>
<td>18.6526</td>
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<tr>
<td>Emetine</td>
<td>67.4750</td>
<td>81.6668</td>
<td>16.2672</td>
</tr>
<tr>
<td>Fisetin</td>
<td>66.8878</td>
<td>40.5343</td>
<td>00.0039</td>
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Conflict of Interests: There are no conflicts of interest.

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