Structural Modeling, Evolution and Ligand Interaction of KMP11 Protein of Different Leishmania Strains

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Abstract

The kinetoplastid-specific KMP11 protein was first described for Leishmania donovani associated with the lypophosphoglycan molecule and is localized mainly around the flagellum and flagellar pocket. This protein is well conserved among kinetoplastids and plays an analogous role in all the flagellates, irrespective of their pathogenicity in humans. The structural elucidation of this important protein may bring about information required to target KMP11 to find valid drug candidates. The atomic-resolution model of KMP11 protein of six different Leishmania strains has been determined from its amino acid sequence by using homology modeling. The stereochemical validation of modeled protein has been done by PROCHECK and Profiles-3D scores. The ligand protein interaction of the KMP11 protein models were carried out with several anti-leishmanial drugs i.e. miltefosine, sitamaquine, pentamidine, amphotericin B, SAG (sodium antimony gluconate), leishmanial peptide, paromomycin and vinblastine and an anticancer compound, sulforaphane. Glutamic acid (E) and lysine (K) of KMP11 are the key amino acids during ligand-receptor interaction. From structural and docking analyses, it is hypothesized that KMP11 of a specific Leishmania strain interacts with a specific anti-leishmanial drug candidate i.e. miltefosine interacts only with KMP11 of L. braziliensis but not with KMP11 of any other Leishmania strain. Highest docking score was found in case of pentamidine. Anticarcinogenic compound, sulphoraphane has shown comparable docking scores and H-bonds with KMP11 protein of six Leishmania strains.

Keywords: KMP; KMP11; Kinetoplastid membrane protein-11; Anti-leishmanial drug; Pentamidine; Leishmania; Sulforaphane; VL; Kal-Azar; Leishmaniasis; Paromomycin

Introduction

Tropical disease like Visceral Leishmaniasis (Kala-azar) caused by Leishmania species has become a significant cause of morbidity and mortality in 88 countries (Ashford et al., 1992; Rosypal et al., 2003). It has been reported that VL in adult patients are co-infected with HIV from 33 countries (Rosenthal et al., 2000). Resistance to pentavalent antimonials [Sb (V)] has been reported earlier even from India which has been the first line of drug of choice for treatment of leishmaniasis (Faraut-Gambbarelli et al., 1997; Lira et al., 1999; Sunder et al., 2000). Second line drugs e.g. pentamidine and amphotericin B has severe side effects and high cost which limit their use (Mishra et al., 1992). Miltefosine (hexadecylphosphocholine) has been approved as the first oral drug for leishmaniasis. It can be used for both antimony-sensitive and antimony-resistant patients (Sunder et al., 1999). In vitro studies have indicated that a single point mutation may lead to miltefosine resistant in the parasite and miltefosine is contra-indicated in pregnancy.
The kinetoplastid membrane protein-11 (KMP11) was first described in *Leishmania donovani* associated to the lipophosphoglycan (LPG) molecule and is located throughout the parasite surface. The 11 kDa molecule was first isolated from *Leishmania donovani* and its primary structure was determined by protein and DNA sequencing (Jardim et al., 1995). KMP11 protein has a defined cellular localization mainly around the flagellum and flagellar pocket. This protein is well conserved among kinetoplastids and presents numerous universal characteristics such as amino acid composition, cellular localization, and a high expression level in the insect stages. It plays an analogous role in all these flagellates, irrespective of their pathogenicity in humans.

Serological tests have shown that KMP11 may be used to discriminate *L. chagasi* infection from active VL and may serve as a marker of response to therapy (Passos et al., 2005). KMP11 is a potent stimulator of human T cells (Tolson et al., 1994). The complete *L. donovani* KMP11 protein and peptide fractions of the protein have been shown to act as B and T cell immunogens during visceral leishmaniasis (Jensen et al., 1998).

Evaluation of immune modulatory properties of recombinant antigens kinetoplastid membrane protein-11 (KMP11) and *Leishmania* homologue of receptors for activated C kinase (LACK) in cutaneous leishmaniasis (CL) and mucosal leishmaniasis (ML) patients have shown that the modulatory properties of LACK and KMP11 are due to induction of IL-10 production and may be helpful for attenuating chronic inflammatory diseases. However, in some clinical conditions, as demonstrated for ML, these molecules are not able to suppress the IFN-gamma response, even inducing IL-10 production (Carvalho et al., 2005). Another experiment on KMP11 have also shown that upon stimulation with KMP11, mononuclear cells from leishmaniasis patients produces high levels of IL-10, while a predominant IFN-gamma production could be observed in cultures stimulated with H2A and soluble *Leishmania* antigen. KMP11 is recognized by cells and sera of patients with different clinical forms of leishmaniasis, and KMP11, through IL-10 production, proved to be a potent antigen in modulating type 1 immune response (de Carvalho et al., 2003).

Severe neurodegenerative disorders are likely to occur if *Leishmania* invades into the visceral organs of human. These disorders may not suitably be treated with anti-leishmanial drugs. It is still unknown, how the anti-leishmanial drugs are acting and whether there is any interaction between any of the surface molecules of the parasite and anti-leishmanial drugs. Hence there is requirement of study on structural and functional characteristics of different proteins of *Leishmania* strains to target the proteins to find novel anti-leishmanial drug. This research work was carried out with a view to know whether there is any interaction between the anti-leishmanial drugs and the flagellar protein i.e. KMP11 of different *Leishmania* strains. But x-ray crystallographic structure is not available for this important protein of *Leishmania* species. The main features of this research work was to generate the three-dimensional (3D) structure of the protein based on the available template structural homologues from protein databank and SCOP database, to validate the models by standard parameters, to know the evolution of this protein in different *Leishmania* strains and to identify the key amino acids involved in ligand-protein interactions.

### Methods

In this study various three dimensional structural models of the KMP11 protein of different *Leishmania* strains were generated. The models were validated by Ramachandran plots of PROCHECK and profiles-3D scores of discovery studio software v 2.0. The models of KMP11 were further tested for in silico docking study to know the presence of any interaction between the ligand and KMP11 protein. Various methods applied in this study are given below.

#### Homology Modeling

The homology modeling of KMP11 protein of different strains of *Leishmania* was performed using DS Modeling 2.0. The homologue search and sequence alignment were done by two modules, sequence analysis and protein families. Sequences of eight different strains of *Leishmania* were identified by searching over NCBI (The National Center for Biotechnology Information) website. Protein families calculate multiple sequence alignment using sequence and structure information, aligns sequences of six (because sequences of *L. major*, *L. infantum*, *L. tropica* are identical) different strains of *Leishmania* and its templates. The final 3-D model was generated by MODELER program of Discovery Studio2.0 which includes automated homology modeling and loop modeling.

#### Protein Simulation

KMP11 protein of six different strains of *Leishmania* models could be further refined by CHARMM (Brooks et
al., 1983) in DS Modeling 2.0, which provides powerful mechanics and dynamics protocols for studying the energetics and motion of molecules, from small ligands to multi-component physiological complexes. CHARMM force field (Accelrys) was used throughout the simulation. Constraint was applied to allow only binding site and ligand to be flexible during the simulation.

**Protein–ligand Interaction Study:**

This study was done by LigandFit / LigandScore (Venkatachalam et al., 2003) in DS Modeling 2.0, which is an automated tool for protein – small molecule docking/scoring, including:

- Define binding site (ligand-based or cavity-based).
- Generate ligand conformations (Monte Carlo trials).
- Dock each conformation (align shapes of ligand to binding site; 24 orientation of ligand; Rigid Body Energy minimization (RBM) with grid-based energy function).
- Save the top docked structures (diverse poses).
- Apply scoring function(s) to each docked structure for the best binding mode (binding affinity prediction).

**Mathematical Formula for Ligandfit Score (DS)**

\[
\text{Dock Score (force field)} = - (\text{ligand/receptor interaction energy} + \text{ligand internal energy}).
\]

There are two energy terms in the forcefield version of DockScore, internal energy of the ligand and the interaction energy of the ligand with the receptor. The interaction energy is taken as the sum of the van der Waals energy and electrostatic energy. The computation of the interaction energy can be quite time consuming. To reduce the time needed for this calculation, a grid-based estimation of the ligand/receptor interaction energy is employed. The van der Waals component of the force field interaction energy typically exhibits a steep rise at short interatomic distances, which can have undesirable consequences in the context of ligand-receptor docking. In particular, the combination of approximating the receptor structure as rigid and limited sampling of ligand conformational space tends to overly penalize poses with “mild” short contacts between the ligand and receptor, due to the “hard” nature of the van der Waals potential as defined in most standard force fields. To overcome this tendency, a softened form of the van der Waals potential is employed with the DockScore function. This softened potential rises to a large but finite value at zero interatomic separation. To maintain a proper balance between electrostatics and van der Waals, the electrostatic energy is also softened to prevent it from dominating the van der Waals energy at short separations. The internal energy of the ligand is computed when using the force field version of Dock Score. The purpose of including the internal energy is to avoid ligand conformations with bad internal nonbond clashes. By default, only the standard (not softened) van der Waals energy is used for the ligand internal energy. Including electrostatic energy as part of the ligand internal energy is optionally available.

Different structural models of KMP11 protein of various *Leishmania* strains were docked with different anti-leishmanial drugs; both presently used drugs and previously used drugs for the treatment of leishmaniasis. All the ligand structures were downloaded from PubChem database as *.sdf file.

Docking experiments were also performed in GOLD software using the default GOLD fitness function (VDW = 4.0, H-bonding = 2.5) and evolutionary parameters: population size = 100; selection pressure = 1.1; # operations = 100,000; # islands = 5; niche size = 2; migration = 10; mutation = 95; crossover = 95 (Jones et al., 1997). Scoring function “Goldscore” was used for evaluation of different docking.

**Mathematical Formula for GOLD Docking**

\[
\text{Fitness} = S(\text{hb}_{\text{ext}}) + 1.3750*S(\text{vdw}_{\text{ext}}) + S(\text{hb}_{\text{int}}) + 1.0000*S(\text{vdw}_{\text{int}}).
\]

Five docking runs were performed per structure. If at any time 3 of the 10 poses were within 1.5 Å RMSD of each other, the docking run for that structure was terminated and docking calculations began for the next structure. Best three poses and docking scores were outputted into a *.mol file and text file respectively.

**Result and Discussion**

**Structure Prediction and Evolution**

From BLAST and clustal analysis, KMP11 proteins of different *Leishmania* strains are have ninety percent identity to each other (figure1a). The phylogenetic analysis showed that KMP11 of *Leishmania* make a separate cluster distinct from *Trypanosoma* (figure1b). Evolution of KMP11 protein of different *Leishmania* strains occurred during the same time period.

KMP11 protein has been found to be an essential surface protein of *Leishmania*. Distant homologues were selected for modeling KMP11 protein using MODELER program. Dali program was run to identify template (Holm et al.,
The PDB ids of selected templates were 2odm, 2c5k and 1owa. PDB ‘2odm’ is the crystal structure of *S. aureus* YlaN, an essential leucine rich protein involved in the control of cell shape. PDB ‘2c5k’ is crystal structure of N-terminal domain of tlg1 complexed with N-terminus of Vps51. PDB ‘1owa’ is the solution structural studies on human erythrocyte alpha spectrin N terminal tetramerization domain. PDB templates ‘1nkp’ and ‘1am9’ were also detected to be distant homology of KMP11 from Dompred program (Marsden et al., 2002). Only one domain was detected in KMP11 protein of six different *Leishmania* strains from Dompred program. PDB templates ‘1nkp’ and ‘1am9’ were also detected to be distant homology of KMP11 from Dompred program (Marsden et al., 2002). Only one domain was detected in KMP11 protein of six different *Leishmania* strains from Dompred program.

**Figure 1a:** Clustal W shows very close identity among the KMP11 sequence of *Leishmania* and trypanosome sp.

**Figure 1b:** Phylogram shown here depicts the close analogy among different *Leishmania* sp & *T. cruzi* and *T. rangeli*.
antimony gluconate), leishmanial peptide, paromomycin, vinblastine, sulforaphane, ketoconazole and allopurinol.

It is interesting to note that the antileishmanial drug, pentamidine was able to find five different binding sites in KMP11 protein of various *Leishmania* strains e.g. *L. major*, *L. donovani*, *L. braziliensis*, *L. amazonensis*, *L. panamensis* and *L. guyanensis*. In some binding sites of KMP11 protein, ten different binding conformations of pentamidine have been observed.

Vinblastine is reported to be down regulating mRNA level of KMP11 in *Trypanosoma cruzi*. Vinblastine have no interaction with the KMP11 protein of different *Leishmania* strains as no ligand binding site has yet been found in this study. Hence it is known from this in silico study that vinblastine don’t have any effect on KMP11 protein phase but may have effect at RNA phase (Thomas and Garcia, 2000). Similarly amphotericin B, SAG (sodium antimony gluconate) and leishmanial peptide did not find any ligand binding site in this flagellar protein, KMP11.

In case of *L. braziliensis*, four antileishmanial drugs e.g. miltefosine, sitamaquine, pentamidine and paromomycin showed high dock scores, in case of both GOLD score and DS dock score, which signifies that these four compounds have high affinity for KMP11. KMP11 protein of *L. braziliensis* shows highest ligand protein interaction with antileishmanial drug, pentamidine (figure 3 and figure 4). No other ligand has shown any binding affinity for the modeled KMP11 protein. Ten different binding conformations have been observed during docking study in DS (Accelrys) in case of the above mentioned four drugs. Paromomycin is likely to form two H-bonds with lysine (74th)
Figure 3: A screenshot from docking of KMP11 protein of *L. braziliensis* with an antileishmanial drug, pentamidine. Different binding modes of pentamidine are shown. Green mesh part is the ligand binding site found using ligand receptor interaction tool of Discovery Studio tool (Structure-Flat Ribbons, Hide Sphere, Color, decolor Residue Atoms within Sphere-activate, and Hide residue around sphere).

Figure 4: Screenshot from docking of *L. braziliensis* with pentamidine (having highest dock score 80.939) with ligand-receptor interaction tool of Discovery Studio (Accelrys). It shows three H-bonds formed between different atoms of ligand and amino acids of KMP11 protein of *L. braziliensis* i.e. involved in Glu-52, Glu-71 and Glu (E)-82.
and asparagine (63rd) amino acids. Three amino acids e.g. Glu-52, Glu-71 and Glu-82 of one helix of the modeled KMP11) were found to be involved in formation of H-bonds with this ligand (figure 4). Hence glutamic acid (E) of this flagellar protein (KMP11) is the key amino acid responsible for ligand protein interaction in *L. braziliensis*. Two different ligand binding sites in KMP11 protein are known to be present for pentamidine only. It is interesting to be noted that one pose of pentamidine has highest ligandfit score (80.939) whereas second conformation of pentamidine didn’t show any affinity for KMP11 protein of *L. braziliensis* as it showed very low ligandfit score (7.06).

It’s well known that hydrogen bond plays an important role for the structure and function of biological molecules, especially for inhibition in a complex. Hence pentamidine is likely to be the best antileishmanial drug in case of *L. braziliensis*. It has also been reported in other research works that pentamidine is an aromatic diamidine that displays multiple effects and is active in vitro against a number of different bacteria, protozoa, and fungi.

In case of *L. donovani*, high dock scores have been observed in case of three antileishmanial drugs e.g. sitamaquine, pentamidine and paromomycin. In one of the conformations of ligand paromomycin, four H-bonds are formed & amino acid residues involved are H67, E52 (3H-bond, is bifractional). So glutamic acid (52nd) and histidine (67th) are vital for interaction of paromomycin with KMP11 of *L. braziliensis*. It has been earlier reported that an ointment containing 15% paromomycin and 0.5% gentamicin shows 100% effective in BALB/c mice (Grogl et al., 1999). Paromomycin is a broad-spectrum aminoglycosidic antibiotic. Hence it is predicted that paromomycin may be specifically acting against *L. donovani* as an antileishmanial drug i.e. acting as an antagonist to KMP11 rather than agonist. It has been earlier reported that paromomycin along with gentamycin is 100% effective in BALBc mice.

KMP11 of *L. amazonensis* and *L. guyanensis* showed interaction only with few conformations of pentamidine but it doesn’t show any interaction with other compounds or antileishmanial drugs. In case of *L. panamensis* only one ligand binding site was found, which showed docking with four antileishmanial drugs e.g. pentamidine, paromomycin, miltefosine and sitamaquine. These drug candidates have also proved high GOLD scores and ligandfit scores (DS).

From GOLD program, different number of H-bonds involved in docking and docking scores of antileishmanial drugs with KMP11 protein of different *Leishmania* strains has been shown in figure 5. Pentamidine have shown highest (51.42) GOLD score with KMP11 of *L. guyanensis*.

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![Figure 5: Bar diagram showing GOLD-scores and involved H-bonds between flagellar protein, KMP11 of different *Leishmania* strains and different ligands (antileishmanial drugs and sulphoraphane)](image_url)
Pentamidine have also shown second highest score (49.04) with KMP11 of *L. braziliensis*. Highest number (7) of H-bond involved during docking has been observed in case of paromomycin with KMP11 of *L. braziliensis* and *L. donovani*. In few cases there is no involvement of H-bonds e.g. sitamaquine with KMP11 of *L. braziliensis* and *L. panamensis*, miltefosine with KMP11 of *L. donovani* and *L. amazonensis*. Various amino acids at different positions of KMP11 protein have been found to be essential for ligand protein interaction with implicated antileishmanial drugs. Among them four amino acids namely glutamic acid (E), threonine (T), lysine (K) and methionine (M) have been identified to be key participants of KMP11 protein of different *Leishmania* strains.

Sulforaphane, an anticancer compound, first isolated from broccoli (Zhang et al., 1992) was taken for ligand protein interaction study to know whether it has any comparable number of H-bonds and docking scores to the implicated drugs used for the treatment of leishmaniasis. This compound has shown comparable results, which is learned from the bar diagram considering the number of involved H-bonds in docking of KMP11 protein of each *Leishmania* strain with different antileishmanial drugs and docking scores (GOLD) depicted in figure 5. Highest GOLD score (44.24) has been observed in case of *L. panamensis*, which is more than or comparable to the docking scores of currently implicated drugs for the management of leishmaniasis. Highest number (5) of H-bonds between KMP11 and sulforaphane has been found in case of *L. amazonensis*, which is the second lowest.

Ligand-protein interaction study of sulforaphane in discovery studio have shown highest dock score (51.31) with KMP11 of *L. braziliensis* (figure 6). The superimposition of ten different conformations of sulforaphane with ligand binding amino acids of KMP11 protein of *L. braziliensis* is shown in (figure 7). The labeled amino acids also form H-bond with the concerned ligand. From the figure 6 it is known that sulforaphane is the second ligand having affinity to KMP11 of all *Leishmania* strains considered in this study. Sulforaphane is the second ligand having highest ligandfit score. Four ligand binding sites in KMP11 and ten different conformations of sulforaphane have been detected from discovery studio. In case of *L. major*, sulforaphane binds to KMP11 in three different poses. The interesting thing is that the involved amino acid responsible for formation of H-bond is lysine (LYS29, LYS85 and LYS92) with...
Figure 7: The superimposition of ten different conformations of sulforaphane showing ligand binding amino acids of KMP11 protein of *L. braziliensis* involved in ligand protein interaction. It was found that the output is in agreement with the surface model generated by Ligandfit.

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<th>LigScore2</th>
<th>PLP1</th>
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<th>Jain</th>
<th>PMF</th>
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</table>

Table 1: Different ligands protein interaction carried out in LigandFit tool of Discovery Studio. Scoring of the docking has been done in different methods i.e. LigScore1, LigScore2, PLP1, PLP2, Jain, PMF and Dock scores. The table represents the highest scores of different ligands interacted with modeled KMP11 protein of six different *Leishmania* strains.
different poses of sulforaphane. LYS74 in case of L. amazonensis, LYS45 in L. panamensis, LYS74 and LYS76 in L. guyanensis are responsible for H-bond formation with sulforaphane. Hence it is hypothesized that sulforaphane, which is available in cruciferous vegetables, such as cauliflower, cabbage and kale, may be implicated for the treatment of leishmaniasis in human beings. This novel compound should be tested in vitro and in vivo for appropriate application in antileishmanial therapy.

A table (table 1) of different docking scores based on different scoring functions has been prepared considering top scoring (highest binding affinity) ligand protein interactions of KMP11 protein of six different Leishmania strains and different compounds including antileishmanial drugs from discovery studio (Accelrys). Different docking scoring functions described in the table are LigScore1 and LigScore2 (Krammer et al., 2005), Piecewise Linear Potential—PLP1 (Gehlhaar et al., 1995), PLP2 (Gehlhaar et al., 1999), Jain (Jain, 1996), PMF (Muegge and Martin, 1999). The scoring functions tend to fall into two major classes emphasizing either: H-bonding interactions or van der Waals, hydrophobic as well as polar attractive/repulsive interactions. GOLD score, PLP (1&2), as well as the Monte Carlo scoring functions all have highly weighted H-bonding terms. On the other hand, LigScore (1 & 2), Jain, PMF, and dock score contain highly weighted terms for van der Waals interactions, lipophilic interactions, and polar attractive/repulsive interactions as well as terms for buried and total polar surface areas. High ligandfit scores have also been observed without H-bonding. As H-bonding appears to play a critical role in binding of different ligands, it is not amazing that the scoring functions with H-bonding terms seemed to do better. It is predicted that the ligands, whose dock scores (DS and GOLD) are more than thirty, are likely to have better ligand-protein interaction with KMP11.

For evaluation of dock scores (GOLD score and dock score in table 1) given by these two tools, we have performed one tailed t-test. After the t-test, it was found that there is no significant difference between the two scores given by two software tools as the calculated value (0.1391) is less than the tabulated value ($t_{95} = 1.7247$) in one tailed two sample unequal variance t-test at ninety-five percent confidence interval and degree of freedom = 20.

**Conclusions and Perspectives**

Compounds selected for this study have earlier been reported in other works on leishmaniasis, one compound which hasn’t been reported about its antileishmanial activity is sulforaphane. There is no report that these compounds interact with KMP11 protein of *Leishmania*. This is the first report of ligand protein interaction about the currently used drugs for the treatment of leishmaniasis and KMP11 protein of *Leishmania*. The study of inhibitory capacity of antileishmanial drugs is a prerequisite for design of novel drug candidates against *Leishmania* species. Homology modeling of *Leishmania* KMP11 shed new light on the ligand binding features of this protein. Highest docking score along with more number of H-bonds were taken to be the best inhibitory compound for KMP11 protein of *Leishmania*.

From ligand protein interaction study, it is learned that few drugs, which have been implicated for leishmaniasis treatment, are specific to specific strain (pentamidine to *L. braziliensis*). Few ligands (e.g. vinblastine) don’t have any effect on KMP11. Glutamic acid (E) and lysine (K) of KMP11 are the key amino acids during ligand-receptor interaction. Pentamidine has been found to be interacting with KMP11 protein of different *Leishmania* strains. Hence it is hypothesized that pentamidine may be the best antileishmanial drug from this KMP11 protein and antileishmanial drug interaction study. Paromomycin have also been found to have best interaction with KMP11 of *L. donovani* (figure 8). Except miltefosine and vincristine,
another anticancerous drug, sulforaphane available from different cruciferous plants, have shown comparable results and hence may be implicated for the treatment of leishmaniasis. The treatment cost will be very less if sulforaphane proves to be a drug candidate for the treatment of leishmaniasis in comparison to current applied drugs for leishmaniasis as this is available from cruciferous plants and available throughout the world. These in silico study must be proved in laboratory before these antileishmanial drugs or likely to be antileishmanial drugs can be brought into clinical trials. As KMP11 is localized in the flagellum and flagellar pocket of Leishmania (which can be ascertained by fluorescent microscopic studies) the inhibition efficacy of the above mentioned compounds can be studied by incubating the insectal stage of Leishmania culture with various ligands taken in this study and measuring the IC50 values.

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