Homology Modeling of Leishmanolysin (gp63) from *Leishmania panamensis* and Molecular Docking of Flavonoids

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1. INTRODUCTION

Leishmaniases, a neglected spectrum of diseases considered to have negative implications and many consequences, are caused by protozoa parasites of the genus *Leishmania,* transmitted by female sandflies of *Phlebotomus* (old world) and *Lutzomyia* (new world). Among the molecular factors that contribute to the virulence and pathogenesis of *Leishmania* are metalloproteases, e.g., glycoprotein 63 (gp63), also known as leishmanolysin or major surface protease (MSP). This protease is a zinc-dependent metalloprotease that is found on the surface of the parasite, abundant in *Leishmania* promastigote and amastigote. This study describes the prediction of three-dimensional (3D) structures of leishmanolysin (UniProt ID A0A088RJX7) of *Leishmania panamensis* employing a homology modeling approach. The 3D structure prediction was performed using the SWISS-MODEL web server. The tools PROCHECK, Molprobyty, and Verify3D were used to check the quality of the model, indicating that they are reliable. Best docking configurations were identified applying AutoDock Vina in PyRx 0.8 to obtain a potential anti-leishmanial activity. Flavonoids such as lanaroflavone, podocarpusflavone A, amentoflavone, and podocarpusflavone B showed good scores among these molecules. Lanaroflavone appears to be the most suitable compound from binding affinity calculations.

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and is eliminated during the maturation and activation. gp63 is the most abundant surface protein of promastigotes and contains 1% of the total parasite proteome but is upregulated in amastigotes. gp63 has been reported to interact with the fibronectin receptor and therefore could further aid the adherence of the parasite to macrophages. From these diverse findings, it is clear that by cleavage and/or degradation of various proteins, gp63 can profoundly affect the macrophage functions, favoring the survival of Leishmania.

Even though glycoproteins have been one of the most important targets of the Leishmania parasite, few relevant studies with glycoprotein have been performed. In the search for new drugs for the treatment of leishmaniasis, Shaukat et al. conducted a study of benzimidazole derivatives, whose results both in vitro and in silico through molecular docking showed that these compounds could serve as the basis for the future treatment of leishmaniasis.

Computational methods have been successfully applied to predict protein structures and ligand–protein interactions. Molecular docking is a method that is used to predict the preferred orientation of predominantly small organic molecules (ligands) within biological macromolecules (proteins). In this sense, natural products can be a valuable alternative to provide huge diversity of chemical structures for biological screening tests of Leishmania species; flavonoids are promising drug candidates for the treatment of all forms of leishmaniasis. This study reports a three-dimensional (3D) structure of the leishmanolysin protein, gp63, from Leishmania panamensis that was built through homologous modeling and optimized using molecular dynamics (MD) simulation. The leishmanolysin protein from L. major (ID PDB: 1LML) was used as a template. The structure was validated to check the quality of the models, indicating that they are reliable. Based on the 3D structure, the ligand-binding modes of flavonoid compounds were elucidated using molecular docking in proteins both crystallized and built by homology modeling. The docking analysis revealed that gp63 from L. major and L. panamensis can structurally accommodate various flavonoid-like ligands and helped us in the molecular recognition of the design of the possible inhibitors and provided new knowledge that can be used to treat the disease caused by the Leishmania species.

2. RESULTS AND DISCUSSION

2.1. Homology Model. The structural models of leishmanolysin from L. panamensis were built by comparative modeling using the crystal structure of gp63 of L. major (PDB ID: 1LML) as the template (Figure 1).

The alignment (Figure S1) between leishmanolysin proteins of L. major and L. panamensis indicated the presence of conserved regions throughout the protein under study. The structures were selected based on their similarity, identity, and number of gaps. The alignment is consistent with experimental results. The comparison with the L. major leishmanolysin sequence showed the identity displayed 74.8% with Lpgp63.

These results are within the range of 59–71% reported previously by Sutter et al. for a variety of L. major leishmanolysin proteins coded by chromosome 10. The HEXXHXXGXXH motif of the active site, typical of the zinc-dependent metalloproteinases, is highly conserved. There is an insertion of 62 residues between glycine and the third histidine in each of the proteins modeled for the Leishmania species, as reported by Yao et al. The position of the active site is highlighted with two of the histidines being part of the α-helix (H8) and the other in a loop in the central domain. The zinc atom of the protein lies between the N-terminal and central domains. The alignment is consistent with experimental outcomes reporting that nine disulphide bridges in the template structure are completely aligned with the leishmanolysin sequence. The high conservation of the sequence is also reflected in the similarity of some properties.

2.2. Molecular Dynamics Simulation. Molecular dynamics simulations were employed to optimize and establish the stability of the protein constructed by homology modeling. The MD trajectory was analyzed by computing the root-mean-square deviation (RMSD) of the structures generated during the MD simulation were calculated to characterize the mobility of particular residues (Figure 2A). As expected, the RMSF plot shows a variation in amino acid residues between the different leishmanolysin proteins of L. major and L. panamensis. In the model, the greatest fluctuations were observed in the N- and C-terminal...
domains, while the central domain where the active site is located remains rigid, as reported by Bianchini et al.18 On the other hand, are evidenced mainly in the change of the length of the amino acids in the active site concerning the zinc atom. It is observed in this study that the bond distance of zinc to the Nε2 atoms of His residues undergoes variations.17–19 It is additionally mentioned that the zinc atom shortens its distance to the OE2 of the GLU residue (Figure S2).

A Ramachandran plot was built, and it showed that 92.1 and 91.1% of residues were in the most favored regions and 7.1 and 8% were in the additional allowed regions. After the MD refinement, a checking process was also undertaken. The Ramachandran plot of the 1LML protein and model built by homology showed that 96.0 and 96.3% of residues were found in the most favored allowed regions and 4 and 3.7% were in the additional allowed region (Figure S3). Rodriguez et al. report that the homology model of L. braziliensis shows percentage of residues in favorable regions very similar to those reported in this article for the study of leishmanolysin.20,21

The analysis of the models with ProSA-web service shows a Z-score between −8.27 and −9.81 (acceptable values are below 0.5). The global score was between −0.98 and −1.49. Otherwise, the quality of the models was confirmed by the Verify3D analysis in an average of 91% of the residues with a score >2 in the 3D/one-dimensional (1D) profile.

2.3. Molecular Docking. Molecular docking has been implemented to reveal the interaction between flavonoid compounds and leishmanolysin proteins of L. major and L. panamensis. Table 1 shows 24 flavonoid molecules to have the lowest binding energy docked at the active site for the two leishmanolysin proteins of L. major and L. panamensis, respectively. The chemical structures are indicated in the Supporting Information, Table S1. Inspection of the docked structures shows the most favorable interactions within the active site, particularly between hydrogen bonds and hydrophobic interactions.

Docking of selected ligand molecules and reference compounds onto the binding site (pocket) of the leishmanolysin (gp63) protein and the homologous protein constructed was performed by AutoDock Vina in the PyRx 0.8 module. After that, for each molecule, the best pose based on its conformation and docking binding energy was selected. Since leishmanolysin structures do not contain a co-crystallized ligand, the study was conducted employing the active site reported by Schlagenhouf et al.10 The binding site comprised amino acids HIS226, GLU263, HIS268, HIS334, and MET345 and a Zn atom, and the grid box was centered on these residues. As reported by Shaukat et al.,14 amphotericin B was also used as a reference molecule. The antiparasitic activity of amphotericin B is related to the ability of this antibiotic to form permeable channels in the cellular membrane of the pathogen. These channels cause a leakage of monovalent ions and small organic molecules from the cellular membrane.1,4,22 In both amastigotes and promastigotes, leishmanolysin is present on the surface, and it is possible that amphotericin will interact with this protein. The docking performed with the antibiotic using the gp63 protein showed that it interacts with the amino acids GLU220 and GLU265 and the Zn atom for the case of gp63 of L. major. The proteins constructed by the homology of L. panamensis follow the same scheme described above (GLU218, GLU263, and Zn) (Figure S4).

The docking pose of lanarolavone exhibited the lowest binding affinity, followed by podocarpusflavone A, amento-

Table 1. Binding Affinity (kcal/mol) of the Most Interesting Compounds Obtained by Docking with Different Leishmanolysin of the Leishmania Species

| molecule                        | 1LML | Lpgp63 |
|---------------------------------|------|--------|
| amphotericin B                  | −11.5| −10.7  |
| lanarolavone                    | −10.5| −9.9   |
| podocarpusflavone A             | −10.1| −9.7   |
| amentoflavone                   | −9.9 | −9.7   |
| podocarpusflavone B             | −9.6 | −9.1   |
| pseudotussogal                   | −9.5 | −9.2   |
| tetrahydrorobustaflavone        | −9.4 | −9.7   |
| 3,3′-dehydrodibinib              | −8.7 | −8.3   |
| rhuscalone VI                   | −8.6 | −8.4   |
| epigallocatechin                | −8.6 | −7.3   |
| agathidavone                    | −8.4 | −8.6   |
| SN00000577                     | −8.4 | −7.2   |
| methyltetrahydrodamenteoflavone | −8.4 | −8.5   |
| SN00000355                     | −8.4 | −6.4   |
| SN00000558                     | −8.3 | −7.2   |
| abyssinone IV                  | −8.2 | −7.6   |
| quercitrin                     | −8.2 | −7.5   |
| SN00000365                     | −8.2 | −7.2   |
| 4-hidroxylonchocarpine         | −8.1 | −7.2   |
| (α-naphthoflavone) SN00000328 | −8.1 | −7.3   |
| bipinnatisone A                | −8.0 | −6.5   |
| medicagenina                   | −8.0 | −7.5   |
| SN00000367                     | −8.0 | −7.3   |
| SN00000357                     | −8.0 | −6.8   |
| SN00157618                     | −8.0 | −8.1   |

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flavone, amentoflavone, and podocarpusflavone A when docking was performed with 1LML and Lpgp63 (Figure 3). The most significant interactions between the ligands and proteins were hydrophobic, H-bonding, metal-contact, and π-stacking (π−π T-shaped) interactions. Table 2 summarizes the principal interactions between the first four ligands with the lowest binding affinity and each of the proteins studied. Lanaroflavone, podocarpusflavone, and amentoflavone formed an H-bond with residues LEU224, ALA225, SER418, and GLN341 within the active site, while podocarpusflavone B interacted with GLY222, SER418, and SER465 through H-bonds. All ligands presented hydrophobic links, especially π-stacking, with VAL261, HIS264, LEU257, PRO460, and ALA350 with the rings of biflavonoid compounds. Podocarpusflavone B

Table 2. H-Bond Interaction Information from Docking Calculations between Ligands and Modeled Proteins

| molecule        | 1LML H-bond distance (Å) | Lpgp63 H-bond distance (Å) |
|-----------------|--------------------------|-----------------------------|
| amentoflavone   |                          |                             |
| LEU224 NH−O=C-4 | 1.98                     | LEU222 NH−C==O-4            | 3.00 |
| ALA225 C==O−OH-5 | 2.80                    | ALA223 C==O−OH-5            | 2.24 |
| ALA349 NH−O-1   | 2.19                     | ASP416 O−OH-5*              | 2.80 |
|                 |                          | SER448 OH−OH-4*"           | 2.31 |
|                 |                          | THR459 OH−C==O-4"          | 2.77 |
|                 |                          | THR459 OH−OH-5*            | 2.80 |
| lanaroflavone   |                          |                             |
| HIS264 NH−OH-7  | 2.81                     | LYS339 NH−C==O-4"          | 2.98 |
| ALA346 C==O−OH-7| 2.92                     |                             |
| GLN341 C==O−OH-4* | 2.98             |                             |
| ALA349 NH−O-1   | 2.61                     |                             |
| LEU224 NH−O=C-4 | 1.96                     | LEU222 NH−C==O-4            | 2.02 |
| ALA225 C==O−OH-5 | 2.64                    | ALA223 C==O−OH-5            | 2.16 |
| podocarpusflavone A |                  |                             |
| ALA349 NH−O-1   | 2.40                     |                             |
| SER418 OH−OH-4* | 2.09                     |                             |
| podocarpusflavone B |                  |                             |
| GLY222 C==O−OH-5 | 2.91                     | LEU222 NH−C==O-4            | 2.01 |
| SER418 OH−OH-4* | 2.44                     |                             |
| SER465 OH−OH-4* | 2.77                     |                             |

Figure 3. Docking poses of the binding interaction of molecules in the active site of leishmanolysin from L. major: (A) lanaroflavone, (B) amentoflavone, (C) podocarpusflavone A, and (D) podocarpusflavone B.
interacted with amino acids GLY222, SER418, and SER465 through H-bonds.

An interaction with the zinc atom was observed in lanoraflavone where the oxygen of the hydroxyl group in position 7 favorably interacts with this metal. In podocarpusflavone A and amentoflavone, the O–metal bond of the same hydroxyl group in position 7 was favorable, perhaps because of the distance between the two atoms, while in podocarpusflavone B, there was no interaction with the zinc as was observed.

In the case of leishmanolysin, lpgp63, of *L. panamensis* constructed by homology, the four molecules mentioned above presented the same pattern of interactions as in gp63 of *L. major* (Table 2). Lanaroflavone, amentoflavone, and podocarpusflavone A and B interact through H-bonds with residues LEU222, ALA223, and ALA334. Regarding hydrophobic interactions, the amino acids involved are ILE255, HIS262, and VAL347. The zinc atom has a metallic interaction with the hydroxyl group of the flavonoids (Figure 4).

### 2.4. Solvent-Accessible Surface Area (ASA)

Figure 5A,B shows the solvent-accessible surface area for the two proteins studied, 1LML and lpgp63, respectively. The graph shows that the docked molecules produce an alteration in the access to solvent, being more noticeable in amentoflavone. The HIS264 and GLU265 residues decrease their ASA values with respect to the native protein. The HIS334 residue does not show significant changes for lanaroflavone, podocarpusflavone A, and podocarpusflavone B, while in amentoflavone, the value of the ASA is drastically decreased. On the other hand, the zinc atom is also significantly affected when the docked compounds are compared to the native protein. Table S2 shows the ASA values for all protein residues that interact with docked molecules.

### 2.5. Validation

The molecular docking performed with each of the models and the crystallographic structure was validated by the receiver operating characteristic (ROC) curve. The area under the curve (AUC) for 1LML and LpGp63 was 0.926 ± 0.026 and 0.928 ± 0.017, respectively (Figure S5). The ROC curve discriminates among active ligands and decoy structures, indicating the good overall predictive performance (AUC > 0.5) of docking for the models of lpgp63 and 1LML.
Since the calculation of the enrichment factor (EF1%) represents another method to evaluate the performance of virtual screening in discriminating active from inactive compounds, we performed a docking study with a decoy set including 86 active and 90 inactive molecules. Particularly, analyzing the top 1% of the ranking obtained by sorting the scores in ascending order, it was possible to find good EF values. For 1LML and LpGp63, the EF values were 45 and 44, respectively, which show that the model demonstrated sufficient discriminating capability for exploring the top 1% of docked compounds.

The correlation analysis performed for 32 flavonoids presented a linear relationship between theoretical free binding energy (kcal/mole) vs pIC$_{50}$ for 1LML and the model developed by homology (Figure S5). Both correlations indicate a value of R $>$ 0.5. Therefore, we can conclude that the outcome achieved in the virtual screening with AutoDock Vina for the two proteins tested was favorable (Table S3).

Flavonoids represent a broad family of polyphenolic compounds found in vegetables and fruits. It is generally accepted that these compounds are safe and nontoxic. The most common flavonoids are flavones and isoflavones. Naturally occurring flavones have been reported to have leishmanicidal activity. Various flavonoids have demonstrated activity against amastigotes in macrophage infection, such as quer cetin that reduces intracellular load by 70% at 45 μM. Weniger et al. have reported that lanaroflavone is a prodrug, which needs to be metabolized by the cell to exert its effect.

The molecules pseudotusangol and rhuschalcone VI have a binding affinity of $-9.6$ and $-8.7$ kcal/mol, respectively. These two compounds presented activity against L. major. Further, it was shown that these two molecules exhibit notable attraction for the nucleoside hydrolase enzyme of Leishmania. Methyltetrahydroamanto flavone ($-8.4$ kcal/mol) has a positive interaction energy with L. major methionyl-tRNA synthetase, according to the same authors.

The analyzed compounds, in general, demonstrate affinity for the proteins considered. Although they are molecules of natural origin, Lipinski’s rules are not so relevant in predicting their drug-likeness, as reported for other biologically interesting natural products. We believe that this is because nature has learned to maintain low hydrophobicity and intermolecular H-bond donating potential when it needs to make biologically active compounds with significant molecular weight and considerable numbers of rotatable bonds. In most cases, natural products do not necessarily abide by Lipinski’s rule because they are thought to enter the human body not by passive diffusion but by more complex mechanisms like active transportation and hence are not expected to comply with the rules for bioavailability. Besides, natural products are more likely to resemble biosynthetic intermediates or endogenous metabolites than purely synthetic compounds and hence take advantage of active transport mechanisms.

3. CONCLUSIONS

In this study, the 3D structures of different Leishmania spp. gp63 proteins were constructed by homology modeling using the crystal structure of leishmanolysin (gp63) of L. major as a template. The reliability of three models was assessed by Ramachandran plots, ProSA-web, PROCHECK, Verify3D, and molprobity. As expected, the resulting structure was similar to that of the template. The data reported here are in agreement with what has been suggested by some authors, with the N-terminal and C-terminal regions being the regions of most appreciable variation. The findings indicate a more rigid central domain, although some regions (especially the loop region) were extremely flexible, despite certain points where protein mobility was observed.

Docking studies made it possible to predict affinity, activity, binding, and the orientation of the binding of flavonoids with leishmanolysin proteins. The analysis was based on the evaluation of their binding energy. Compounds like flavonoids, chalcones, and bioflavonoids showed great affinity to the leishmanolysin protein from L. major. Lanaroflavone, the molecule with the lowest binding energy, also showed good antileishmanial activity, despite violating the rule of five (Lipinski’s rules). Flavonoid derivatives are natural products that exhibit promising antileishmanial activity and deserve further research.
4. EXPERIMENTAL SECTION AND COMPUTATIONAL METHODS

4.1. Homology Model. The protein sequence of gp63 from L. panamensis was retrieved from the UniProt database. The Uniprot accession number is A0A088RJX7 with 594 amino acid residues. The crystal structure of gp63 from L. major, employed as a template, was retrieved from the protein database PDB (PDB ID: 1LML).10 The model was generated using the SWISS-MODEL server.38 The best model was selected based on QMEAN and Z-score of ProSA-web. The quality of 3D prediction was assessed using Molprobity.39 ProSA-web,40 and Verify3D.41 The final representation was visualized using Discovery Studio Visualizer.42 The gp63 protein of L. major was obtained from the PDB database with PDB ID 1LML; all water molecules were removed and hydrogen atoms were added. On the other hand, the protonation states of the surrounding histidine residues for both proteins were assigned using PROPKA;43 they were selected as neutral compared to those reported by Sutter et al. (Figure 5).

4.2. Molecular Dynamics Simulation. Molecular dynamics (MD) simulations of 200 ns were performed for the modeled protein and the template from L. panamensis and L. major, respectively, using the GROMACS 2016.5 package;44 all simulations were carried out using Amber99sb as a force field.45 Both macromolecules were solvated by a cubic periodic box in which each protein was solvated with TIP3P water under periodic boundary conditions (PBCs).46 The system was neutralized, and the ionic strength (0.1 mol L$^{-1}$) of the medium was adjusted by adding Na$^+$ and Cl$^-$ ions, keeping the number of particles constant. After these steps, energy minimization of the systems was performed until convergence, which was followed by equilibration with pressure and temperature (NVT and NPT ensembles) kept constant at 300 K and 1.0 bar, respectively; equilibration periods were 1.0 ns, production runs were of 10 ns duration, and the V-rescale thermostat and Parrinello–Rahman thermostat were used. The LINCS57 and SETTLE58 algorithms were employed to determine bond lengths of hydrogen atom distance constraints, respectively, whereas long-range interactions were calculated employing the particle-mesh Ewald (PME) method,49,50 used to constrain the geometry of the water molecules. The equilibrated system was subjected to the final MD production run of 10 ns, applying the periodic boundary conditions (PBCs) and integrating the equation of motion every 2 fs. GROMACS and VMD51 software packages were used to analyze the MD trajectories. Root-mean-square deviation (RMSD) of Ca residues and root-mean-square fluctuation (RMSF) were also calculated. In total, the production time was 200 ns for each system.

4.3. Ligand Database and Preparation. In total, 5470 drug-like compounds (chalcones, flavonoids, and bilavonoids) were retrieved from the Super Natural II database,52 as a single file in spatial data file (SDF) format. This file was imported in Open Babel53 and all ligand structures were converted into PDB format. Then, the compounds were optimized with density functional theory (DFT) at the B3LYP/6-31G level using the Gaussian 09 package.54 Amphoterin B was obtained from the PubChem database55 and employed as reference molecules in molecular docking studies according to the suggestions by Saukot et al.

4.4. Molecular Docking. Molecular docking was performed using AutoDock Vina56 in PyRx 30.8 (virtual screening tools).57 Then, the resulting collection of potential ligands was docked into the leishmanolysin proteins. Hydrogen atoms were added to all proteins, and partial atomic charges were calculated. Initially, flexible-ligand docking was done. The grid box size was set to $35 \times 35 \times 35$ points with a spacing of 0.375 Å. For the calculation, 150 runs of the Lamarckian genetic algorithm (LGA)58 with 25 000 000 evaluations and 270 000 generations were performed.

4.5. Validation. To validate docking, scoring functions were assessed using different parameters like the area under the curve (AUC), receiver operating characteristic (ROC) curve, and enrichment factor (EF1%) according to Meekum et al.59 A dataset of 176 molecules with strong, moderate, and weak antileishmanial activity was employed to validate and to assess the selectivity and sensibility of our methodology. A linear correlation analysis was performed between the biological activity (pIC$_{50}$) and the affinity of flavonoids reported and the calculated affinity values were, in addition, used for validation of the virtual screening done by AutoDock Vina (Figure S6).

The statistical significance of the AUC value of different models was evaluated with a $p$ test with a 95% confidence limit.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.0c01584.

Homology modeling, molecular dynamics simulation, and docking were performed on leishmanolysin of Leishmania panamensis using gp63 of L. major as a template. A series of flavonoids were docked into the active site of gp63 from L. major and L. panamensis, and their scores were in the order laranorolavone > amentoflavone > podocarpusflavone A > podocarpusflavone B (PDF)

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Author Contributions
J.M.-C., H.G.-E., A.P., and L.C.-C. conceived and designed the experiments; J.M.-C., A.P.H.G.-E., and L.C.-C. performed the experiments; J.M.-C., H.G.-E., R.V.-R., and M.L.S.G. analyzed the data; R.V.-R. was also responsible for the correspondence of the manuscript. All authors discussed, edited, and approved the final version of the manuscript.

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Notes
The authors declare no competing financial interest.

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