Monocytic leukemia-associated antigen-42 (MLAA-42) is a novel-identified monocytic leukemia-associated antigen which is overexpressed in acute monocytic leukemia. In an attempt to uncover the mechanism of MLAA-42 overexpression, it significantly works as an elongation factor which plays an important role in polypeptide elongation in leukemia as shown in the biochemical pathway (Figure 1) [5]. Furthermore, MLAA-42 plays a critical role in the transformation of normal cells to malignant phenotype through binding with GTP bound protein leading to unrestrained proliferation of the abnormal white blood cells which lead to AMoL. In the meantime, inhibition of proliferation of bone marrow cells, which takes place if there is no binding between MLAA-42 and GTP, is caused directly to prevent formation of AMoL [5].

Heterocyclic compounds have been found to gain extensive interest due to their considerable wide range applications in medicinal, pharmaceutical, and pharmacological activities especially as anticancer agents [6, 7]. In the present study, computer aided virtual screening studies were carried out using ChemBank™ database to report new molecular entities which can be used as potent inhibitors against MLAA-42 protein. This study warrants further investigations to indicate that MLAA-42 is a novel target for
designing newer drug-like molecules for leukemia treatment, using in silico approaches.

2. Results and Discussion

2.1. Generation of the Three-Dimensional Structure of MLAA-42. For the case where the target MLAA-42 is not experimentally determined by means of X-ray crystallography and nuclear magnetic resonance (NMR), the computational-based techniques are used for 3D structure prediction [8]. The FASTA sequence of the target protein (ID: Q6W6M8) with 151 amino acid residues was identified from Universal Protein Resource "UniProtKB" database (https://www.uniprot.org/uniprot/Q6W6M8) [9]. The template was recognized by subjecting the obtained sequence to BLAST, Jpred3, and Domain Fishing servers, and Table 1 represents the resulted maximum E-value. The 1N0V "Crystal structure of elongation factor 2" was selected as a template protein, on the basis of parameters such as query cover (77%), maximum identity (63%), and statistical E-value (8 × 10^{-47}) with the target [10]. ClustalW server was used to perform the alignment of both elongation factor proteins MLAA-42 and 1N0V (541-720 aa), as shown in Figure 2. The conserved amino acid residues are elucidated as (*) (39.3%), highly similar residues as (:), and weakly similar residues as (.)(51.7%). The individual percentages, resulted from the server, refer to the identity and similarity of the target residue sequence with its template sequence, respectively.

Twenty-five models of MLAA-42 were generated by using Modeller 9.11 software, depending on satisfying spatial restraints in terms of probability density function [11]. The 3D model with fewest restraint violations, lowest probability density function (855.33), and acceptable geometry was selected for further validation [12]. Figure 3 represents the 3D model of the target, which consists of five alpha-helices and six beta-strands. The secondary structures of both 1N0V and MLAA-42, which represent the amino acid sequences of helices and strands in both proteins, are represented in Figures 4 and 5 and (Figure S1 and Tables S1 and S2) in the supplementary data section, using PDBsum server [13]. To obtain a stable confirmation, energy minimization and protein loop modeling were performed by GROMOS96 Force-Field set, using Swiss-Pdb Viewer [14]. To get insight of the superimposition between the proteins MLAA-42 and 1N0V, the root-mean-square deviation RMSD for the backbone of MLAA-42 and 1N0V is 0.51 Å (within the permissible range i.e. ≤2 Å), which indicates close homology and ensures reliability of the model, as declared in Figure 6.

2.2. Physicochemical Properties and the Abundance of Amino Acids in MLAA-42. The physicochemical properties of MLAA-42 were predicted using ProtParam tool [15]. The protein sequence has 151 amino acid residues with molecular weight of 16.871 kDa. Furthermore, the most abundant amino acid residues are LEU, SER, GLU, ALA, VAL, LYS, GLY, and ASP, respectively, in high percentages in the target protein, as shown in Figure 7. Lucine has the highest abundance (9.9%), and methionine has the lowest abundance (0.7%). The physicochemical parameters predicted a

Table 1: The template protein selection.

<table>
<thead>
<tr>
<th>Server name</th>
<th>Parameters for template selection</th>
<th>E-values</th>
<th>PDB code</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCBI-Blast</td>
<td>Sequence similarity</td>
<td>9e^{-42}</td>
<td>1N0V</td>
</tr>
<tr>
<td>Jpred3</td>
<td>Secondary structure prediction</td>
<td>6e^{-40}</td>
<td>1N0V</td>
</tr>
<tr>
<td>Domain fishing</td>
<td>Domain similarity</td>
<td>3e^{-40}</td>
<td>1N0V</td>
</tr>
</tbody>
</table>

1N0V was selected as a template protein using different servers, based on query cover, maximum identity, and statistical E-value.

Figure 1: Biochemical pathway of MLAA-42 protein involves transformation of normal cells to malignant phenotype through binding with GTP bound protein leading to unrestrained proliferation of the abnormal white blood cells which lead to AMoL.
negatively charged protein as the result of the high number of negatively charged residues (aspartic acid 6.0% and glutamic acid 8.6%) in contrast with positively charged ones (arginine 5.3% and lysine 7.3%). The atomic composition of MLAA-42 is 2364, with 747 carbon (C), 1176 hydrogen (H), 204 nitrogen (N), 233 oxygen (O), and 4 sulfur (S). In addition, the protein is acidic, with an isoelectric point (pI) of 5.47. The estimated half-life of the target showed that it can remain intact without being degraded for 3.5 h in humans, 10 min in yeast, and less than 10 h in *E. coli*, and its extinction coefficient is 24075 M$^{-1}$ cm$^{-1}$. Finally, the generated aliphatic index was 82.65, with grand average of hydrophobicity (GRAVY) of $-0.426$ and an instability index of 34.55.

Figure 2: Pairwise sequence alignment of MLAA-42 with 1N0V. The conserved amino acid residues are elucidated as (*), highly similar residues as (:), and weakly similar residues as (.).

Figure 3: 3D structure of MLAA-42 protein. It consists of five $\alpha$-helices and six $\beta$-strands.

Figure 4: The secondary structure of the target MLAA-42 obtained by the PDBsum server.
2.3. Validation of the Model and Active Site Analysis. The 3D model was validated by different tools such as PROCHECK (Ramachandran plot) and ProSA to check its structural integrity. Ramachandran plot (Figure 8) of the modeled protein represents 87.3% (117 aa) of the total residues in the most favored regions and 11.9% (16 aa) in additionally allowed regions, indicating a good quality model. Moreover, ProSA Z-score (dark spot) is $-2.85$, which falls within the values range of the known proteins determined by X-ray (light blue) and NMR (dark blue) (see Figure S2, in the supplementary data section), predicting a reasonable quality model. Figure S3, in the supplementary data section, shows the 3D model quality by plotting energies ($X$-axis) as a function against residue sequence position ($Y$-axis). The figure shows that most of residues of MLAA-42 have negative energies, which indicates an acceptable model. In view of the facts mentioned above about the unavailability of 3D structure of MLAA-42 in protein data bank PDB, thus, there are no previous data obtained for the binding regions of the target. Therefore, to overcome the lack of experimental data, herein, CASTp and SiteMap cheminformatics tools were used to identify the binding site cavities of the target (Figures S4 and S5 and Tables S3 and S4), in the supplementary data section, respectively. CASTp server gives information about hydrophobic pocket regions; in addition, it measures cavity volume and area. The results declared four sites, site 1 has volume of $40.9 \, \text{Å}^3$, site 2 with volume of $28.2 \, \text{Å}^3$, site 3 with volume of $50.7 \, \text{Å}^3$, and site 4 has volume of $105.4 \, \text{Å}^3$. Among the four different cavities obtained from CASTp, the amino acids from PHE44 to ILE96 are consistently present in all sites. SiteMap gives information about the hydrophilic and hydrophobic regions present on the protein surface. It identified two binding cavities on MLAA-42 protein. The site 1 with volume $1222.79 \, \text{Å}^3$ and size of $4.38 \, \text{Å}$ and site 2 has volume of $46.30 \, \text{Å}^3$ and size of $32 \, \text{Å}$. The active sites predicted by SiteMap are from TYR38 to ILE100. In conclusion, these computational prediction techniques give information about the binding site regions that are composed of TYR38 to ILE100 residues are responsible for MLAA-42 docking with ligand molecules. The docking interactions show that the residues TYR38, LYS40, THR89, ILE96, ASP99, and ILE100 are observed to be crucial to identify the antagonists against MLAA-42 protein. Figure S6, in the supplementary data section, represents the grid with dimensions $80 \times 80 \times 80 \, \text{Å}$ around the binding site, to perform further studies.

2.4. Molecular Docking Analysis and ADMET Profile. The compounds with heterocyclic core structures are pharmaceutically important class of compounds because of their diverse range of biological importance such as anticancer...
activity against cancer cell lines [6, 7]. In view of the facts mentioned above and as a part of our efforts to identify new anticancer drug candidates [16], a ChemBank™ library of 2344 small molecules with an anticancer activity was downloaded [17] for specific docking. Computer-aided screening protocol at the binding regions of the target was performed using virtual screening workflow docking program of Glide, Schrodinger suite. The glide docking program identifies specific structural motifs and provides exceptionally large contribution to enhance binding affinity [18]. The docking approach was run in a flexible docking mode which automatically generates different confirmations for each ligand molecule. The total isomeric structures (13,354) obtained after ligand preparation was subjected to high throughput virtual screening (HTVS) mode and 10% hit structures (1335) obtained as output, followed by 10% of them, 133, screened in standard precision (SP) mode, and finally the 10% of the resultant were screened in the Extra precision (XP) mode to obtain 13 hits. The docked compounds were prioritized according to their binding energies (in range of $-12.19$ to $-9.22$ Kcal/mol) after docking to the active site pocket of the target protein, as tabulated in Table 2.

A sample of six best docking interactions between the ligands and protein, arranged rankwise based on their binding energies, is depicted in Figure 9 and Table 2. The docked compounds were prioritized according to their binding energies (in range of $-12.19$ to $-9.22$ Kcal/mol) after docking to the active site pocket of the target protein, as tabulated in Table 2.

![Figure 7: The abundance of amino acids in MLAA-42. Graphical representation of the abundance of 20 amino acids present in the target. Lucine has the highest abundance (9.9%), and methionine has the lowest abundance (0.7%).](image)

![Figure 8: Stereochemical analysis of MLAAA-42. The red region declares the most favorable area of residues; the yellow region is additionally allowed; and generously allowed residues in the light-yellow region. The RC plot declares 99.2% of residues falling in the allowed region.](image)
Table 2: The chemical structures of top six scoring ligand molecules against MLAA-42 protein.

<table>
<thead>
<tr>
<th>S. no.</th>
<th>Structure and IUPAC name of molecule</th>
<th>Glide energy (KJ/Mol)</th>
<th>Binding energy (Kcal/mol)</th>
<th>The interacting amino acid residues</th>
<th>Bond distance (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6-([‘N’-[2-(4-Chloro-phenoxy)-acyl]-hydrazino]-6-oxo-hexanoic acid N’-[2-(4-chloro-phenoxy)-acyl]-hydrazide</td>
<td>-63.59</td>
<td>-12.19</td>
<td>H-bonds: LYS40: N – L1</td>
<td>2.91</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>HYD94: O – L1</td>
<td>2.32</td>
</tr>
<tr>
<td>2</td>
<td>2-[2-(1-Methyl-buta-1,3-dienyloxy)-acetylaminio]-5,6-dihydro-4H-cyclopenta[b]thiophene-3-carboxylic acid (2-ethoxy-phenyl)-amide</td>
<td>-50.27</td>
<td>-11.83</td>
<td>H-bonds: TYR38: N – L2</td>
<td>2.89</td>
</tr>
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<td></td>
<td></td>
<td>ILE100: N – L2</td>
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</tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>TYR38: O – L2</td>
<td>1.94</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>π–π interactions: TRP85-L2</td>
<td>4.95</td>
</tr>
<tr>
<td>3</td>
<td>N-[3-(Naphthalen-1-ylcarbamoyl)-4,5,6,7-tetrahydro-benzo[b]thiophene-3-carboxylic acid (2-ethoxy-phenyl)-amide</td>
<td>-42.22</td>
<td>-11.61</td>
<td>H-bonds: ILE100: H – L3</td>
<td>1.68</td>
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<td>THR98: O – L3</td>
<td>2.36</td>
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<tr>
<td>4</td>
<td>1-(Naphthalene-2-sulfonyl)-pyrrolidine-2-carboxylic acid (2-(2-chloro-phenoxy)-ethyl) amid</td>
<td>-47.52</td>
<td>-11.30</td>
<td>H-bonds: LYS40: N – L4</td>
<td>2.95</td>
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<td>THR98: N – L4</td>
<td>2.65</td>
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<td>THR98: O – L5</td>
<td>2.13</td>
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<td>ASP99: OD1-L5</td>
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<td></td>
<td></td>
<td>π–π interactions: TRP85-L5</td>
<td>4.87</td>
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<tr>
<td>6</td>
<td>N-(2,4-Dichloro-phenyl)-3-(N’-thioacetylhydrazine-carbonyl)-propanamide</td>
<td>-40.65</td>
<td>-11.10</td>
<td>H-bonds: ILE100: N – L6</td>
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<td>TYR38: O – L6</td>
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<td></td>
<td></td>
<td>THR98: OG1-L6</td>
<td>2.29</td>
</tr>
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</table>

Table represents top six ligand molecules with binding energies in range of -12.19 to -9.22 Kcal/mol, hydrogen bonds, and pi-interactions to MLAA-42’s active site pocket.
Figure 9: Continued.
Figure 9: Interactions between MLAA-42 with ligand molecules (a)–(f). (A) Three-dimensional docked poses: the binding residues of MAA-42 are represented in yellow stick, the ligand molecules are shown in the purple stick model, the hydrogen bonds are shown in black dotted lines, and pi-pi interactions in orange lines. (B) Two-dimensional docked poses: the residues are shown in a 3-letter code, hydrogen bonds in pink lines, and pi-pi interactions in green lines.
(Figure 9). While in 2D representation, the residues are shown in a 3-letter code, the hydrogen bonds are represented in pink lines, and π-interactions are shown in green lines. The ligand molecules with heterocyclic moieties such as thiophene, pyrazole, and pyrrole and amide groups such as carbamoyl (-CONH-) and sulphonamide (-SO₃NH-) in ligands L1-L6 are observed to be common pharmacophore groups which interact with the active site region (TYR38, LYS40, TRP 85, ILE96, THR98, ASP99, and ILE100) of MLAA-42 through a network of hydrogen bonds and π-interactions, as represented in Table 2. The carbamoyl group, which is present in all ligand molecules, formed intermolecular hydrogen bonds with the residues TYR38, LYS40, TRP 85, ILE96, THR98, ASP99, and ILE100. In addition, the sulphonamide group, which is present in ligand L4, formed a hydrogen bond interaction (N---H---O) with the residue THR98 at the distance of 2.65 Å. Furthermore, the tryptophan (TRP85) formed π-π interactions (face to face) with the benzene ring of ligand L2 on its side chain plus the pyrazole ring of ligand L5, at the distance of 4.95 and 4.87 Å, respectively. The intermolecular hydrogen bonds and π-interactions add more stability to the docked complexes [21]. The other docked compounds to the target are included in supplementary data section as Figure S7.

SASA values of MLAA-42, before and after docking, were analysed using Discovery Studio Visualizer (see Figure S8, in the supplementary data section). The decrease in SASA values confirms that the amino acid residues (TYR38 to ILE100) are involved in bonds formation with ligand molecules (a)–(f).

The ADMET properties for the resulted ligand molecules were determined in silico by using the QikProp module of the Schrödinger suite. The molecules with agreeable ADMET properties are considered as new novel drug candidates, as shown in Table 3. Interestingly, all the compounds have a molecular weight in the acceptable range of 349.7 to 511.3 Daltons (less than 725 Dalton). BBB+ value describes the ability of the compound to cross the blood-brain barrier, which is in the permissible ranges for all compounds. They had <10 hydrogen bond acceptors and <5 hydrogen bond donors, and log P values of <5. These properties are in the reasonable range of Lipinski’s rule of five (LORF) [22]. The human oral absorption, partition coefficient (QLogPo/w), and water solubility (QLogS) values are in the agreeable range defined for human use, which indicates their possible ability to be drug candidates, as shown in Table 3. Also, the values show that the compounds can be absorbed by the human intestines. Furthermore, all molecules (a)–(f) displayed negative AMES toxicity test which means that the ligands are non-mutagenic. Also, they displayed negative carcinogenicity test. These ligands can be considered as highly potent inhibitors against leukemia.

3. Materials and Methods

3.1. 3D Homology Model of MLAA-42. The protein’s three-dimensional structure is required to understand its function [23]. Herein, an in silico homology modeling generates a three-dimensional (3D) for an unknown structure of protein (target) depending on one or more proteins of known structures (templates) as reported earlier by Aboubakr et al. in 2016 [24, 25]. The generation of (3D) structure of MLAA-42 was carried out using comparative modeling techniques. The amino acid sequence of MLAA-42 is retrieved from the UniProtKB database in FASTA format. The resulted sequence is submitted to BLASTp, Jpred3, and Domain fishing servers, to identify suitable templates [26–28]. E-value is identified the similarity between the target and template proteins [29]. The template with maximum identity is selected for generating the 3D structure of the target. The pairwise sequence alignment of MLAA-42 with its template is performed using the ClustalW tool to recognize the structurally conserved regions [30]. ClustalW server uses the Gonnet matrix algorithm to make certain the presence of conserved motifs and to minimize the atomic gaps. The % identity of alignment is measured as a ratio number of the identical residues in the alignment and the total number of residues of the target. The % similarity is measured as a ratio of the total number of identical and similar residues to the total number of residues of the target. The 3D structure of the target is generated using Modeler 9.11. Then, the model is energy minimized using SPDBV and Impref in Schrödinger to obtain a stable conformation [31]. The RMSD value, which is calculated using the Swiss-Pdb viewer, is used to find the best superimposition for protein structures. The lower value (≤2 Å) means the best alignment between the structures.

3.2. 3D Structure Validation. Subsequently, quality of the generated model is evaluated by computational protocols such as PROCHECK and Protein Structural Analysis (ProSA) [32, 33]. PROCHECK, a program that relies on Ramachandran plots for structure verification, figures out the stereochemical quality of the model. Furthermore, ProSA is applied to check for energy criteria in comparison with a large set of known protein structures with similar size [34].

3.3. Binding Site Prediction and Ligand Preparation. It is the hydrophobic cavities on the surface of a protein, which is responsible for its specificity [35]. It is theoretically determined by means of tools based on recent theoretical and algorithmic results of computational geometry such as CAST-p and SiteMap of Schrödinger suite [36, 37]. These computational prediction tools analytically furnished the area and volume of cavities. The grid is created around the binding cavity to perform further structure-based virtual screening studies [38]. LigPrep module of Schrödinger suite is used to generate various conformers of small molecule depending upon its structural features [39].

3.4. In silico Molecular Docking, SASA, and ADMET Analysis. Molecular docking protocol is used to predict the preferred orientation of a ligand molecule to the target protein to form a stable complex [40, 41]. The docking accuracy is determined by finding how closely the binding confirmation with
the lowest energy of the cocrystallized ligand molecule predicted by the object scoring function; G-score (Glide score) resembles an experimental binding modes determined by X-ray crystallographic technique [42]. Once the 3D model of MLAA-42 is validated and the target binding pockets are defined, the docking-based virtual screening study is performed using the Glide docking tool incorporated in the Schrödinger package by Maestro [43]. The virtual screening approach is performed through hierarchical flexible docking methods: High Throughput Virtual Screening (HTVS), Standard Precision (SP), and Extra Precision (XP). The ligand molecules are prioritized on the basis of the docking score, docking energy in each step, and by default 10% of the molecules selected and considered for the next hierarchical step [18]. Solvent Accessible Surface Area (SASA) of the target, before and after docking, is computed using Accelrys Discovery Studio Visualizer 3.5 software. Prediction of drug-like profiles, such as physicochemical, pharmacokinetic, and safety of the compounds, is performed using the QikProp module [44].

4. Conclusion

In this work, the computer-aided drug design protocols were used to identify novel leads against MLAA-42 protein. The homology model of the target was evaluated by homology modeling techniques. In silico molecular docking were also adopted to identify the lead compounds. The resulted molecules with heteroscaffolds and amide groups (-CONH- and –SO2NH-) exhibited better estimated binding energy values and agreeable pharmacokinetic properties and were ranked as potent drug-like candidates against the target protein. Hence, MLAA-42 has emerged as a therapeutic target for treatment of leukemia carcinoma.

Abbreviations

MLAA-42: Monocytic leukemia-associated antigen-42
UniProt: Universal protein resource crystal structure of
1N0V: elongation factor 2
3D: Three-dimensional
GEF: Guanine nucleotide exchange factors
GAP: GTPase-activating proteins
BLASTp: Basic alignment tool program
Jpred3: Java prediction V 3.0

Table 3: ADME properties of the top docked ligand molecules.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>CNS</th>
<th>M. (Wt)</th>
<th>SASA</th>
<th>Volume</th>
<th>Donor HB</th>
<th>Accept HB</th>
<th>QPlogPo (w)</th>
<th>QPPCaco</th>
<th>QPlogBB</th>
<th>Percent human oral absorption</th>
<th>*NandO</th>
<th>Rule of three</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1</td>
<td>−2</td>
<td>511.3</td>
<td>905.7</td>
<td>1554.4</td>
<td>1</td>
<td>8</td>
<td>4.7</td>
<td>64.5</td>
<td>−2.8</td>
<td>74.0</td>
<td>10</td>
<td>1</td>
</tr>
<tr>
<td>L2</td>
<td>0</td>
<td>436.5</td>
<td>751.0</td>
<td>1348.2</td>
<td>1</td>
<td>5</td>
<td>5.5</td>
<td>2264.9</td>
<td>−0.5</td>
<td>100</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>L3</td>
<td>−2</td>
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<td>740.5</td>
<td>1298.5</td>
<td>2</td>
<td>6</td>
<td>4.4</td>
<td>48.1</td>
<td>−1.7</td>
<td>83.1</td>
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<tr>
<td>L4</td>
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<td>458.9</td>
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<td>8</td>
<td>3.9</td>
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<td>−0.6</td>
<td>100</td>
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<tr>
<td>L5</td>
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<td>349.7</td>
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<td>4</td>
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<td>−0.2</td>
<td>100</td>
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<td>0</td>
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<tr>
<td>L6</td>
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<td>411.3</td>
<td>708.3</td>
<td>1210.2</td>
<td>3</td>
<td>7</td>
<td>3.9</td>
<td>538.7</td>
<td>−0.8</td>
<td>100</td>
<td>6</td>
<td>0</td>
</tr>
</tbody>
</table>


Impref: Impact refinement module
ProSA: Protein structure analysis
Castp: Computed atlas of surface topography of protein
RMSD: Root-mean-square deviation
SPDV: Swiss PDB viewer
OPLS: Optimized potentials for liquid simulations
PPIs: Protein-protein interactions
SASA: Solvent accessible surface area
ADME: Adsorption, distribution, metabolic, and excretion.

Data Availability

The data used to support the findings of this study are included within the article.

Conflicts of Interest

The authors declare no conflicts of interest.

Authors’ Contributions

I. S, A.H.A., and H. R. M. R designed the study. I. S. carried out the homology modeling of the target and evaluated the 3D model. A. H. A performed the virtual screening approach and wrote the preliminary draft of the manuscript. H. R. M. R was responsible for identification of ADME profiles of the molecules and wrote the corresponding section. All authors contributed to the revision of the drafts and agreed on the final version to be submitted.

Acknowledgments

The authors thank Prof. V. Uma, Molecular Modeling Research Lab., UCS, Osmania University, India, for the help and support.

Supplementary Materials

Figure S1: the secondary structure of MLAA-42. The 3D model of the target protein consists of five helices as red cylinders and six strands as pink arrows. Figure S2: The 3D model quality of MLAA-42. The score (−2.85) is falling in the range of PDB proteins identified by NMR (dark-blue region) and X-ray crystallography (light-blue region). Figure S3: ProSA of MLAA-42 protein. The plot shows local model
quality by plotting energies as a function against amino acid sequence position. Figure S4: binding site of MLAA-42 from the CAST-p server. Figure S5: binding site of MLAA-42 by SiteMap module. Figure S6: receptor grid generation by Schrodinger Suite. The grid generated with 80 Å × 80 Å × 80 Å dimensions using receptor grid generation of Glide module. Figure S7: two-dimensional interactions between MLAA-42 with other ligand molecules. Figure S8: the surface accessibility of MLAA-42 and docked complex (L1). Red-colored peaks represent SASA value of MLAA-42 (after docking), blue-colored peaks represent before docking. X-axis represents the amino acid residues and Y-axis represents SASA value. Tables S1 and S2: the secondary structure details of MLAA-42 protein (S1) the α-helices in MLAA-42 protein. (S2) the β-strands in MLAA-42 protein. Table S3: binding site of the MLAA-42 by CAST-p server. Table S4: binding site of MLAA-42 by SiteMap module. (Supplementary Materials)

References


