DOCKING STUDIES FOR VARIOUS ANTIBACTERIAL BENZILATE DERIVATIVES

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ABSTRACT

Objectives: In this study, we have focused on discovering the leads for the enzyme targets of infectious disease tuberculosis. We employed computer-aided drug design docking tool, to discover new leads for Mycobacterium tuberculosis (MTB).

Methods: Five compounds were synthesized and they are made to dock into the active site of the enzyme; retrieved from protein data bank.

Results: The docking studies and structure-activity relationship reveals that the compound 2'-chloro-4-methoxy-3-nitro benzilic acid after three different docking strategies reveals that the score was found to be higher compared with others (−5.568 kcal/mol).

Conclusion: On the closer analysis of this molecule, the molecule showed stacking interaction and the compound has also found to be surrounded by non-polar amino acids, which makes this molecule potent toward antibacterial drug discovery.

Keywords: Antibacterials, Docking, Absorption, Distribution, Metabolism and excretion study, Resistance.

INTRODUCTION

As there is an increased number of drug-resistant bacterial cases worldwide, there is an urgent need for novel therapeutic interventions including innovative antibacterial and antymycobacterial drugs [1,2] with no cross-resistance to available drugs in the market. The most pathogenic bacteria Mycobacterium tuberculosis (MTB), the causative agent for tuberculosis in humans leads to a bacterial killer worldwide [3]. The bacteria have led to the emergence of multi-drug resistant and extensively drug resistant strains of bacteria. The treatment is based on the combination of two or more antibiotics, and the side effects are many. To limit the medications and side effects the preliminary studies have been done for the small molecule inhibitors which have been shown good antibacterial activity toward many pathogenic bacteria. Recently, benzilic acid derivatives were synthesized as potent antibacterial agents with good activity range. They inhibit the pathogenic bacteria’s which includes Staphylococcus aureus, Klebsiella pneumoniae, Escherichia coli, and Staphylococcus epidermidis. Virtual screening, a computational method where the compounds could be assessed for their potential to bind specific sites on target molecules such as proteins, was employed in the study [4]. Furthermore, the pharmacokinetic properties were also predicted for future perspective of the small molecule compounds.

METHODS

Computational details

The computational details for this study were conducted in an Intel Core i5 capacity processor with memory of 4GB random access memory running with the windows 7 operating system. The virtual screening options for high throughput virtual screening (HTVS), standard precision (SP) and glide extra precision (XP) docking were all checked to be executed. The module glide XP of Schrodinger 9.3 (glide, version 5.7, Schrodinger, LLC, and New York, 2015) was utilized to perform docking studies. Suitable bonding and the charges were added to the hetero atoms and the corresponding hydrogen atoms were added to all the atoms [5].

Preparation of ligands

The synthesized molecules were processed through the Lipinski filters to enable the drug property. Ligand preparation was performed for the synthesized molecules using LigPrep module available in the software (LigPrep v2.2, Schrodinger LLC, NY) and Epik (Epik v1.6, Schrodinger, LLC, NY) to expand protonation and tautomeric states at 7.0±2.0 pH units. Conformational sampling was also performed for all database molecules using the Confgen search algorithm. Confgen with OPLS 2005 force field was applied for the generation of conformers with duplicate poses eliminate if the RMSD was <2.0 Å. A distance-dependent dielectric constant of four and maximum relative energy difference of 10 kcal/mol were applied [6].

Molecular docking

Docking studies for the synthesized compounds were performed using glide module of Schrodinger, LLC. 2015. Primarily, using glide module [7,8] (grid based ligand docking with energetics), we examined for important interactions based on the reference ligand and the protein of interest in the flexible mode docking. The glide module with three modes of docking, HTVS, SP, and XP mode was employed sequentially. The XP mode was used for exhaustive sampling and advanced scoring resulting in even higher enrichment.
Finally, the shortlisted hit molecules were selected based on the visual inspection of amino acid interaction, docking score, and the active site cavity [9].

**ADME prediction**

All the synthesized compounds for our study were selected, and the molecules were subjected to ADME predicted analysis using QikProp module of Schrodinger. The important properties such as octanol-water coefficient (logP), human oral absorption, Lipinski’s rule of five, blood-brain barrier (BBB) coefficient, HERG property, and CaO2-2 permeability property were predicted for the synthesized compounds, and also the predicted results were checked for any violations to determine the nature of the compounds.

**RESULTS AND DISCUSSIONS**

Synthesized compounds have taken for docking studies to establish the structure-activity relationship using crystal structure of MTB co-crystallized with inhibitor thiazole benzamide (protein data bank ID:4WYC). Analysis of crystal structure of 4WYC revealed with hydrogen bonding interactions with nonpolar interaction like Trp398. The inhibitor is well associated with hydrophobic amino acids met61, Trp398, Tryr407, and phe402 [10]. To validate the active site pocket the reference ligand was redocked and the docking score was found to be −6.032 kcal/mol. Redocking results showed that the compound exhibited similar interactions as that of crystal structure and showed an RMSD of 1.02 Å. Further, the compounds synthesized were screened based on three different docking strategies [11]. The ligand interaction with protein was depicted in Fig. 1. The docking score and the ligand interactions for the compounds were tabulated in Table 1.

The compound 2'-chloro-4-methoxy-3-nitro benzilic acid was found to inhibit the pathogenic bacteria’s S. aureus, K. pneumoniae and E. coli at a distance of 10 mm using disc diffusion method, when compared to other compounds. The compound after three different docking strategies reveals that the score was found to be −5.568 kcal/mol. On the closer analysis of this molecule, the molecule showed similar stacking interaction like the reference molecule; the compound has also found to be surrounded by nonpolar amino acids, which makes this molecule potent toward antibacterial drug discovery. The binding analysis and ligand interaction diagram for the compound 2'-chloro-4-methoxy-3-nitro benzilic acid was depicted in Fig. 2.

Based on our docking studies, it has confirmed that the structure changes in the compounds series were found to be well correlated with in vitro antibacterial results. The compound 4, 4'-dibromo benzilic acid possessed high docking score of −5.228 kcal/mol with the stacking interaction with amino acid Trp64. The activity of this compound also found to be well correlated with the reference ligand. The close analysis of this compound revealed that the compound is well packed with nonpolar interactions which make this compound more active against the pathogenic bacteria. The binding analysis and ligand interaction diagram for the most active compound 4, 4'-dibromo benzilic acid are shown in Fig. 3. The docking score and its ligand interaction for the synthesized compounds are tabulated in Table 1.

The compound 2, 2'-dichloro benzilic acid was found to inhibit the bacteria at a distance of 8 mm. This is quite lesser than the other molecules. This makes this molecule more effective binding, and the docking score was found to be −5.121 kcal/mol. The binding analysis of this compound reveals that the compound well fitted into the active site pocket and the group phenyl chloride was found to interact with nonpolar amino acids Tyr25 and Trp64 which reveals that there are two stacking interactions making this compound more stable for further processing as better drug compound [12]. The binding analysis and ligand interaction for the compound 2, 2'-dichloro benzilic acid was depicted in Fig. 4.

The compound benzilic acid was found to inhibit the bacteria K. pneumoniae at a distance of 12 mm. This compound after in silico screening analysis was found to possess good docking score −5.069 kcal/mol. On the closer analysis of this compound reveals that the molecule has well fitted into the active site pocket of the protein; also their ligand interaction shows that the molecule was surrounded by nonpolar amino acid and it is found to be interact with an amino acid Trp64 which is an important interaction of original ligand. The binding

**Table 1: Docking score and ligand interaction results for the synthesized compounds**

| S. No. | Compound name                         | Docking score kcal/mol | Ligand interaction |
|-------|---------------------------------------|------------------------|--------------------|
| 1     | 2'-chloro-4-methoxy-3-nitro benzilic acid | −5.568                 | Trp64              |
| 2     | 4, 4'-dibromo benzilic acid            | −5.225                 | Tyr157             |
| 3     | 2, 2'-dichlorobenzilic acid            | −5.121                 | Trp64, Trp25       |
| 4     | Benzilic acid                         | −5.069                 | Trp64              |
| 5     | Methyl benzilate                       | −3.140                 | Trp64              |
using 3.078 −3.195 100 BB 1558.995 1558.995 QPP −0.229 100 QPlog 324.014 4.169 −0.619 −4.996 L-alanine QPlog 86.397 antibacterial analysis. Further, these compounds 2.98 −0.403 −2.726 antimicrobial, anthelmintic and docking studies, the compound possesses less docking score −3.140 kcal/mol. The compound methyl benzilate does not show any antibacterial activity with any of the pathogenic bacteria’s. After molecular docking studies, the compound possesses less docking score −3.140 kcal/mol. The closer analysis of this reveals that the compound does not well fit into the active site of the enzyme. Furthermore, the benzyl moiety has facing outward which might have makes this compound less active when compared with other compounds virtually and biologically. The binding and ligand interaction for the compound methyl benzilate was shown in the Fig 6. 

ADME prediction
To further account for the potential of the compounds to act as efficient drug candidates, their absorption, distribution, metabolism, and excretion (ADME) properties were also calculated in silico using Qikprop. The obtained values for molecular logP, HERG property, CaCO accessibility, BBB, and human oral absorption; it is also used to assess violation of Lipinski’s rule of five if any. All the compounds were shown to correlate well with the human oral absorption. BBB separates the human brain from the direct contact of circulatory system, thus protecting the brain for unwanted solute particles. Both the predicted compounds were shown to be BBB negative ensuring their administration safe for the brain. The ADME predictions for the synthesized compounds were tabulated in Table 2.

CONCLUSIONS
We utilized the medicinal chemistry tools of structure-based drug design strategy. Docking studies were performed to identify new scaffold molecules. This strategy revealed hitherto unknown binding pockets and inhibitor binding modes distinct from the earlier reported inhibitors and will be exploited successfully in further antitubercular drug development process. The most active compound 2’-chloro-4’-methoxy-3-nitro benzilic acid was found to be most active in both in silico and in vitro antibacterial analysis. Further, these compounds will be carried out for their antitubercular property as these are small molecule leads could easily cross the cell barrier systems in mycobacteria.

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Table 2: ADME prediction for the synthesized compounds

| Compound name                  | QPlog | QPlog Po/w | QPP | QPlog HERG | Percent human oral absorption |
|-------------------------------|-------|------------|-----|------------|-----------------------------|
| Methyl benzilate              | 3.078 | −4.996     | 1558.995 | −0.403     | 100                         |
| 2,2’-dichlorobenzilic acid    | 4.169 | −2.726     | 324.014   | −0.229     | 96.291                      |
| 2’-chloro-4-methoxy-3-nitro benzilic acid | 3.078 | −4.996 | 1558.995 | −0.403 | 100 |
| 4,4’-dibromo benzilic acid    | 4.199 | −3.195     | 222.95    | −0.303     | 93.562                      |
| Benzilic acid                 | 2.98  | −3.193     | 222.314   | −0.619     | 86.397                      |

*Predicted octanol/water partition coefficient logP (acceptable range: −2.0 to 6.5); *Predicted IC50 value for blockage of HERG K+ channels (below −5); *Predicted apparent CaCO-2 cell permeability in nm/seconds (<25 poor; >500 great); *Predicted brain/blood partition coefficient (<3.0-1.2); *Percent human oral absorption (<25% is poor and >80% is high); *Rule of 5 violation (mol_MW <500, QPlogPo/w <5, donorHB ≤5, accptHB ≤10)
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