Molecular Docking Simulation Based Virtual Screening for The Design of Potential Inhibitors of Heme Oxygenase of Corney Bacterium Diphtheria.

Somdutt Mujwar, Raghav Mishra, Isha Tomer, Alekh Gour

Abstract: Diphtheria is an infectious human disorder affecting upper respiratory tract which is characterized by fever, sore throat, and malaise. It is caused by Corynebacterium diphtheriae and other pathogenic strains of Corynebacterium. The pathogen invades in the nasopharynx and infects the host by releasing an exotoxin leading to the severe concerns in the kidneys, nervous system and cardiac muscles. Sometimes diphtheria infections may be fatal because of the circulatory failure caused by myocarditis. Hemeoxygenase is the rate limiting enzyme in heme degradation and catalyzes the NADPH-cytochrome P450 reductase-dependent cleavage of heme to biliverdin with the release of iron and carbon monoxide. In the present paper we have performed molecular docking simulation based in-silico virtual screening of an NCI diversity set-II containing 1593 diverse ligands to identify potential inhibitor of the Heme oxidase enzyme of Corneybacterium diphtheria. The lead molecules are shortlisted on the basis of their binding energy and these molecules are supposed to be further evaluated experimentally for development of a newer therapy for the treatment of diphtheria.

Keywords: Corney bacterium diphtheriae; Diphtheria; Heme oxygenase: Molecular docking: virtual screening.

1. INTRODUCTION

During the recent past, there is a rise in the human population which are getting affected by life-treating infectious diseases. Microbial drug resistance hasn’t simply supplemented in the expansion in the rate of mortality identified with infections and the frequency of diseases, still there isbudgetary increase in the health care expenses for the treatment for drug-resistant infections[1]. On account of the dramatic escalation in microbial infections, newer antimicrobial agents havingunique mechanisms are earnest need to fight against the multidrug-resistant infections. The medicinal chemists are primarily exercising nowadays on the discovery and development of new biologically active agents for the treatment of these alleviating ailments. The design and development of drug molecules clearly bidsundoubtedly with the utmost hopes for endeavors in the present and future era[2].

Diphtheria is an uncommon disease in most developed nations inerable from the routine childhood vaccination. However, the disease is seen at times among non-immunized children in developing countries. [3] Diphtheria is brought about by a toxigenic bacteria Corynebacteriumdiphtheriae which stays as one of the imperative reasons of illness and death among children. There are four principle biotypes of C. diphtheriae: Gravis, Intermedius, Mitis, and Belfanti. Globally, the Intermedius is most often associated with exotoxin production, despite all three strains are able of producing exotoxin. [4] It is an overwhelming life threatening situation, with the issues requiring timely recognition, fast treatment against the diphtheria antitoxin and in the most extreme circumstances critical intensive care interventions to comfort the obstructions of the upper respiratory system. Diphtheria toxin causes quick obstruction of the upper airway by the massive soft tissue oedema, acute and subacute cardiac, the formation of the pseudo-membrane in the pharynx, larynx and nares, acute and subacute cardiac, renal and neurological complications. In spite of the availability of the diphtheria vaccine since 1921, its appropriation was limited in the initial days and a million diphtheria cases were reported and over 60,000 deaths were happened every year in the 1970s [5]. The incorporation of diphtheria toxoid vaccine in the worldwide Expanded Programme on Immunization in 1974 saw a decrease of cases by over 90% from 1980 to 2000. With a fundamental proliferation number of 6 to 7 (average number of secondary cases in a fully susceptible population), only 85% of each birth associate need to be immunized to achieve the critical immunity threshold that gives diphtheria herd immunity. [6] Three dosage of the vaccine containing diphtheria toxoid has been continued universally since 2010, despite the fact that with the increasing country to country variations as well as the rise of re-centre proves the homogenous coverage of population by the diphtheria. The systemic indications of diphtheria are because of serious tissue damage throughout the body caused by the excretion of the diphtheria exotoxin [7]. The diphtheria toxin was encoded by the tox gene, which has been appeared to be under transcriptional control of the iron-dependent diphtheria toxin repressor protein [8]. It is a global iron-dependent repressor likened to the ferric uptake repressor protein of Gram-negative bacteria. Iron is a basic supplement required for the survival of most bacteria, and the capacity of pathogenic bacteria to get adequate iron during infection is essential for such pathogens to cause disease [9].

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To dodge the low availability of iron, pathogens have developed refined mechanisms to utilize the host’s iron- and heme-containing proteins. Hemeoxygenase is the rate limiting enzyme in heme degradation and catalyzes the NADPH-cytochrome P450 reductase-dependent cleavage of heme to biliverdin with the release of iron and carbon monoxide. [10] In the present paper we have performed molecular docking simulation based *in-silico* virtual screening of an NCI diversity set-II containing 1593 diverse ligands to identify potential inhibitor of the Heme oxidase enzyme of *Corneybacterium diphtheria*. The Heme oxidase inhibitors are supposed to interfere with the life cycle of the pathogen and should act as a potential therapy for the treatment of diphtheria.

II. EXPERIMENTAL

A. Selection of Macromolecule and its Preparation

The Hemeoxygenase (HeO) enzyme of *Corneybacterium diphtheria* bound with ligand protoporphyrin-IX (HEM) (pdb id-1v8x) was downloaded from the protein data bank (Figure 1). The Hemeoxygenase enzyme was prepared for molecular docking simulation by removing ligand and from its active binding site, removal of water molecules which are not interacting with the ligand, and addition of polar hydrogen atoms [11-12].

![Figure 1. Crystal structure of the Hemeoxygenase enzyme of C. diphtheria acquired from the protein data bank database. (PDB ID-1v8x)](image)

**B. Preparation of Ligand for Molecular Docking**

The bound ligand HEM is prepared for molecular docking simulation by providing the rotatable, non-rotatable as well as un-rotatable bonds present in the ligand to the AutoDock software [3].

**C. Identification of Binding Site**

The ligand binding site of the HemeOxygenase enzyme is identified by exploring the binding interaction of the bound ligand HEM by using PyMol software. The complexed ligand HEM bound in the receptor’s active binding site is separated from the complex molecule by using software chimera [14-15].

**D. Molecular Docking**

The binding site of the Hemeoxygenase is identified by using protein visualization PyMol software to enumerate the grid parameter points of the grid box required to perform the molecular docking simulation of ligand molecules with pathogenic Hemeoxygenase. These grid parameters were utilized for all docking runs. The grid-box was placed by centering the ligand molecule and covering all the residues involved in the binding of the ligand to ensure that all the extended conformations of ligand fits within the grid box (Figure 2).

![Figure 2. Three-dimensional grid-box covering the ligand binding site of the pathogenic Heme oxygenase enzyme.](image)

The separate map files for each of the atom types present in the receptor as well as ligand viz. A Cl OA SA F H P Br Fe HD N C I NA S etc. are prepared by running Autogrid utility of the AutoDock suite. These map file prepared by Autogrid is used for carrying out molecular docking simulations by AutoDock [16-17]. Lamarckian genetic algorithm (LGA) is one of the primary conformational search approach employed in AutoDock for molecular docking simulation. A trail population is created for various possible conformations, followed by the mutation, conformational parameters exchange, and compete in a manner kindred to biological evolution in successive generations for eventually selecting individuals with lowest binding energy. The individual conformational search for its local conformational space, discovering local minima, and then proceed this information to later generations is performed by “Lamarckian” aspect, which is its additional feature. The binding energy of the small molecules with macromolecular targets is predicted by using semi-empirical force field. The force field allows the assimilation of the intramolecular energies into the predicted binding energy by the evaluation of the energetics for both bound as well as unbound states based on a comprehensive thermodynamic model.
Docking parameter file required for the docking of each ligand molecule was prepared by using the 150 Genetic Algorithm (GA) runs, 250000 maximum numbers of evaluations, 27000 maximum numbers of generations and 0.02% rate of gene mutation [16-17].

E. Docking Method Validation

The position and orientations of the ligand obtained after the molecular docking study represents probable binding patterns of the inhibitors. The various docking parameters considered in the docking methods were validated by re-docking individually crystalized ligand HEM over Hemeoxygenase enzyme. The molecular docking simulation technique is validated by using following parameters:

(a) Overlay Methods

The validation of molecular docking method is performed by overlay method. The docked conformation of the bound ligand should be impeccably overlaid with reference to the bioactive conformation of the ligand present in the crystal structure of the downloaded protein.

(b) Chemical Resemblance

The molecular docking method is validated when the docked ligand should have same interactions with the residues of macromolecule as that present in the downloaded crystallized macromolecule.

F. In-silico Virtual Screening

In the present study the NCI diversity set-II containing 1593 diverse ligand molecules were virtually screened against validated pathogenic Hemeoxygenase enzyme. The molecular docking simulation based in silico virtual screening is performed to identify potential inhibitor molecules for Hemeoxygenase enzyme of *C. diphtheria*.

G. Analysis of Molecular Docking Simulation Results

After performing molecular docking simulation of the selected ligand molecules against the pathogenic Hemeoxygenase enzyme, the best ligand molecule were evaluated on the basis of their lowest binding energy. The Lamarckian Genetic Algorithm used for scoring. All the results obtained by molecular docking simulation were evaluated on the basis of hydrophilic and lipophilic interactions obtained between the binding residues present in the active ligand binding site of the macromolecule and ligand. The empirical range of the free binding energy should be in the range of -5 to -15 kcal/mol. The mathematical equation to calculate binding affinity of the specific ligand for a particular target is as follows:

$$K_i = e^{\frac{\Delta G}{RT}}$$

Where, $\Delta G$=change in free energy upon binding, $R$=gas constant and $T$=temperature.

H. Pharmacokinetic and Toxicity Evaluation

The physicochemical parameters as well as toxicity profile of all the selected lead molecules were evaluated by using DataWarrior software. DataWarrior software checks for the presence of major toxic effects such as mutagenicity, tumorigenecity, irritant effect and reproductive effects in the lead molecules. This software searches for the presence of those functional group which are responsible for the toxic effects of the ligand molecule. The DataWarrior program also calculates drug-likeness and drug score of the lead molecule on the basis of their physicochemical properties [17].

III. RESULTS AND DISCUSSION

A. Selection and Preparation of Macromolecule

Hemeoxygenase bound with ligand HEM (pdb id-1v8x) was downloaded from protein data bank database. The three dimensional structure model of protein is procured by using X-ray diffraction technique at a resolution of 1.85 Å by using *E. coli* as an expression system. The 1v8x protein complex consists of three identical polypeptide chain of 215 amino acids. The chain B and C were removed with the help of chimera software and chain A was selected for the experiment. The receptor molecule is prepared for molecular docking simulation process by adding polar hydrogen bonds, removing redundant water molecules and addition and distribution of charge. After processing the receptor molecule it was saved in *.pdbqt format by using AutoDock software.

B. Preparation of Ligand for Molecular Docking

Eight rotatable bonds were present in the bound ligand molecule. All the eight bonds were kept rotatable in the ligand molecule for the current experimental study. The prepared ligand was saved in the *.pdbqt format.

C. Identification of Binding Site and Grid-Box Preparation

The amino acid residues Tyr130 and Arg177 were involved in the active binding of the HEM ligand with the pathogenic Hemeoxygenase enzyme. The binding interactions of the bound ligand HEM ligand with the pathogenic Hemeoxygenase enzyme is shown in Figure 3.
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An appropriate grid box was prepared by covering all the macromolecular residues which are involved in the active binding of the bound ligand HEM ligand with the pathogenic Hemeoxygenase enzyme. The coordinates used for the preparation of the grid box were tabulated in table 1.

### Table 1. The grid coordinates for pathogenic Heme oxygenase enzyme.

| Proteins | x-D | y-D | z-D | Spacing (Å) | x-center | y-center | z-center |
|----------|-----|-----|-----|-------------|----------|----------|----------|
| 1v8x     | 40  | 40  | 40  | 0.425       | 25.134   | -9.27    | 1.154    |

### D. Molecular Docking Simulations and Its Validation

The results obtained after molecular docking of the bound ligand HEM ligand with the pathogenic Hemeoxygenase enzyme were presented in table 2.

### Table 2. Molecular Docking results of ligand HEM ligand with the pathogenic Heme oxygenase enzyme (1v8x).

| Protein | Interacting residue | RMSD | Binding energy (kcal/mol) |
|---------|---------------------|------|---------------------------|
| 1v8x    | Tyr130 and Arg177   | 0.41 | -11.92                    |

The molecular docking process for docking of particular ligand with a specific macromolecule was performed by considering following parameters:

(a) Overlay Method

The molecular docking method was validated as the docked conformation of the ligand should be perfectly overlaid with the crystal structure of the ligand present in the downloaded protein. The overlaid conformation of the docked ligand with reference to the crystal structure of the downloaded ligand was shown in figure 4.

(b) Chemical Resemblance

The molecular docking method was validated when the docked ligand had the similar interactions with the residues of macromolecule as that were present in the downloaded crystallized macromolecule. The interactions present in crystal structure and the interactions present in the docked structure were shown in figure 5.

### E. Virtual Screening

The binding affinity of all the ligand molecules were identified by analyzing the binding energy obtained for top ranking pose of each ligand and interactions of docked compound were visualized. The molecular docking results of all the nine flavonoid molecules were obtained after performing AutoDock based molecular docking simulation against the pathogenic Hemeoxygenase enzyme were shown in table 3.
Table 3. Binding energy of selected best ligand molecules against the pathogenic Heme oxygenase enzyme.

| S. No. | Name           | Structure                  | Binding Energy |
|-------|----------------|----------------------------|----------------|
| 1     | ZINC126709 09  | ![](image1.png)             | -11.02         |
| 2     | ZINC054927 94 | ![](image2.png)             | -10.58         |
| 3     | ZINC047058 02 | ![](image3.png)             | -9.79          |
| 4     | ZINC174659 58 | ![](image4.png)             | -9.77          |
| 5     | ZINC011632 59 | ![](image5.png)             | -9.70          |
| 6     | ZINC130990 27 | ![](image6.png)             | -9.65          |
| 7     | ZINC173756 49 | ![](image7.png)             | -9.54          |
| 8     | ZINC126709 03 | ![](image8.png)             | -9.43          |
| 9     | ZINC020349 99 | ![](image9.png)             | -9.40          |
| 10    | ZINC043768 56 | ![](image10.png)            | -9.36          |

F. Pharmacokinetic and Toxicity Evaluation

The toxicity evaluation of the lead molecules by observing the presence of functional groups responsible for the major toxic effects like mutagenicity, tumorogenicity, irritant effect as well as reproductive effect were identified by using DataWarrior software. The results obtained in the toxicity profiling of all the selected lead molecules are shown in the table 4. The results of toxicity profiling of all lead molecules suggests that the five out of ten compound, i.e. ZINC12670909, ZINC01163259, ZINC12670903, ZINC02034999 and ZINC04376856 are not having any functional group responsible for major toxic effects like mutagenicity, tumorogenicity, irritant effect as well as reproductive effect with very good pharmacokinetic profile.

Table 4: Toxicity profiling for the pathogenic Heme oxidase inhibitors.

| Lead Compound   | Mutagenic | Tumorigenic | Irritant | Reproductive Effect |
|-----------------|-----------|-------------|----------|---------------------|
| ZINC12670909    | None      | None        | None     | None                |
| ZINC05492794    | High      | None        | None     | High                |
| ZINC04705802    | None      | None        | High     | High                |
| ZINC17465958    | None      | None        | High     | High                |
| ZINC01163259    | None      | None        | None     | None                |
| ZINC13099027    | None      | None        | High     | High                |
| ZINC17375649    | None      | None        | None     | None                |
| ZINC12670903    | None      | None        | None     | None                |
| ZINC02034999    | None      | None        | None     | None                |
| ZINC04376856    | None      | None        | None     | None                |

IV. CONCLUSION

AutoDock based molecular docking simulation technique is very useful in identification of mechanism of action for some of the potential inhibitors for Heme oxidase enzyme of C. diphtheriae. The NCI diversity set-II containing 1593 diverse ligands are evaluated for their binding affinity against the pathogenic Heme oxidase enzyme. The ligand molecules having best binding affinity are identified as a lead molecules having potential inhibitory activity of pathogenic Heme oxidase enzyme. These selected lead molecules are further evaluated for their pharmacokinetic profiling and presence of toxic effects. The results of toxicity profiling of all lead molecules suggests that the five out of ten compound, i.e. ZINC12670909, ZINC01163259, ZINC12670903, ZINC02034999 and ZINC04376856 are not having any functional group responsible for major toxic effects like mutagenicity, tumorogenicity, irritant effect as well as reproductive effect with very good pharmacokinetic profile. Thus, these five molecules are supposed to be further evaluated experimentally for development of a newer therapy for the treatment of diphtheria.
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AUTHOR PROFILE

I'm a researcher working in the field of bioinformatics and computational biology having sound knowledge of Pharmaceutical Chemistry. My research interests revolve around understanding the pathophysiology of various diseases and using knowledge acquired from my pharmaceutical background and its application in my current research studies for designing novel lead molecules with high potency and free from side effects as well as development of drug resistance by the pathogenic microorganisms. Presently I’m working on the In-silico design of riboswitch based novel inhibitor molecules for various pathogenic microorganisms which can be free from associated side effects as well as development drug resistance by the pathogen. I am using AutoDock based virtual screening technique to screen potent lead molecules, which are further used for the development of pharmacophore. This designed pharmacophore structure is used for the development of novel inhibitor molecules by using biosisotropic substitution technique, which further evaluated for their pharmacokinetic properties on the basis of their physicochemical properties.

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