Introduction

Tuberculosis is a common as well as one of the deadly infectious diseases caused by *Mycobacterium tuberculosis*. It affects most of the world’s population, mainly in developing countries (Harries and Dye, 2006; Lopez and Mathers, 2006). Antibiotics were prescribed but effective treatment is challenging due to the complicated structure and chemical composition of the mycobacterium cell wall. The unusual structure of the bacterial cell wall makes many antibiotics ineffective and check the entry of drugs (Jain and Mondal, 2008).

Isoniazid and ethambutol have been used for the decades as frontline drugs to inhibit *M. tuberculosis*, but the rise of multi-drug resistant and extensively drug resistant strains poses a serious threat to present treatment of tuberculosis (Burris, 2004; Mcllerton, et al., 2009; Zhang, et al., 2014). Ethambutol inhibits the synthesis of essential components of the mycobacterial cell wall. Ethambutol targets the biosynthesis of the cell wall, inhibiting the synthesis of both arabinogalactan and lipoarabinomannan. It is assumed to act via inhibition of arabinosyl transferases (Amin, et al., 2008). An arabinosyltransferase is a transferase enzyme acting upon arabinose belongs to the family of glycosyl transferases. Ethambutol has been also reported for several toxic effects such as optic neuritis, color blindness etc (Kumar, et al., 1993). Therefore, the need of new and alternative drug candidate for tuberculosis is obvious and current approaches is aim to screened lead molecule from chemical database using Computer aided screening methodology based of known chemical...
structure of ethambutol.

Materials and Methods

Receptor and ligand retrieval and analog design for ethambutol: The 3-D structure of arabinosyl transferase (3PTY) was retrieved from Protein Data Bank. The structure of ethambutol was also retrieved from Drug Bank. Structural similarity, sub-structure, identity (70%) search were performed and carried out for ethambutol like compounds using Molsoft ICM Browser 3.5-1p and ChemBioDraw Ultra 12.0 software (Gogoi, et al., 2012; Lagunin, et al., 2000). Compounds library were collected from ZINC Database, PubChem, Chemspider, ChemBank cheminformatics site in sdf format. Ethambutol structure based analogs were also designed manually using Chem Sketch software. Around 100,000 compounds were considered and screened for ethambutol like candidate lead compound. Open BabelGUI tool was used for chemical file conversion purposes.

Ligand structure optimization and physicochemical properties calculation: Screened ligands were optimized before docking using MM2 force field of ChemBio 3D ultra. Physicochemical properties (Hydrogen bond acceptor, hydrogen bond donor, number of rotatable bond, calculated log P, molecular weight, etc) were predicted and checked for non-violation of drug like and Lipinski’s rules using PreADMET server.

Potential protein binding sites prediction and molecular docking study: The potential ligand binding site of arabinosyl transferase receptor was computed at MVD workspace. Volume and Surface of the binding site were computed and optimum binding site was selected to perform docking. The screened compounds were imported in the Molegro Virtual Docker workspace. The bonds flexibility of the ligands was set and the side chain flexibility of the amino acids in the binding cavity was set with a tolerance of 1.10 and strength of 0.90 for docking simulations. RMSD threshold for multiple cluster poses was set at <2.00 Å. The docking algorithm was set at a maximum iteration of 1500 with a simplex evolution size of 50 and a minimum of 20 runs. Molecular docking was carried out using Molegro Virtual Docker (MVD) (Molegro APS: MVD 5.0) (Thomsen and Christensen, 2006). MVD is molecular visualization and molecular docking software which is based on a differential evolution algorithm; the solution of the algorithm takes into account the sum of the intermolecular interaction energy between the ligand and the protein and the intramolecular interaction energy of the ligand. The docking energy scoring function is based on the modified piecewise linear potential (PLP) with new hydrogen bonding and electrostatic terms included. Interaction of Ligands with receptor was studied to know the best binding orientation of receptor-ligand complex in terms of minimum energy score.

ADME and toxicity prediction: Absorption, Distribution, Metabolism, Excretion and Toxicity were studied for top ranking compounds were computed using PASS (Prediction of Activity Spectra for Substances) Inet and Pre ADMET server (Gogoi, et al., 2012; Lagunin, et al., 2000). PASS Inet predicts 3678 pharmacological effects, mechanisms of action, mutagenicity, carcinogenicity, teratogenicity and embryotoxicity. MDCK cell permeability, human intestinal absorption, blood-brain barrier penetration and plasma protein binding scores were studied and compared (Nordin and Bergstrom, 2006).

Result and Discussion

Herein, we have screened out 3148 compounds structurally similar with ethambutol from 11,74,583 compounds based on chemical similarity (structural) using ZINC database. We have also retrieved ethambutol like 5 compounds from Chemspider on the basic of calculated property, 10 from ChemBank on the basic of substructure and 3 from Pubchem on the basic of property. We calculated physicochemical property for 222 compounds in Molsoft ICM-Browser software and observe that most of the compounds follows Lipinski’s rule of Five as presented in the Table I including few analogues of ethambutol.

The compounds with the predicted drug likeness of more than 80% with Lipinski’s qualification were used to study their ADME properties. 222 compounds were checked for absorption and distribution in human body using PreADMET as given in Table II. Each compound was checked for carcinogenic, embryo toxin and teratogenic and 31 non-toxic compounds were chosen for molecular docking analysis.

Receptor model was exported and potential bindings sites were predicted in the Molegro Virtual Docker workspace as presented in the Table III with their coordinate position in the workspace. Missing coordinates of receptor was checked before loading. Amino acid residues around the binding cavity were given in the Table IV.

Molecular docking is a novel approach to study small compound inhibition to receptor protein. We docked 31 non toxic compounds with receptor model of M. tuberculosis arabinosyl transferase using Molegro Virtual Docker (MVD) software. MVD is molecular visualization and molecular docking software which is based on a differential evolution algorithm; the solution of the algorithm takes into account the sum of the intermolecular interaction energy between the ligand and the protein and the intramolecular interaction...
Table I: Physicochemical property of top ranking database compounds

| Compound ID | Formula         | HBA | HBD | Rot B | MW     | ClogP |
|-------------|-----------------|-----|-----|-------|--------|-------|
| ZINC0388344 | C₆H₆NO          | 1   | 1   | 1     | 130.123| 0.796 |
| ZINC0441875 | C₆H₅NO₂         | 1   | 3   | 4     | 199.181| 0.57  |
| ZINC0690002 | C₆H₅NO          | 1   | 1   | 4     | 142.123| 0.60  |
| ZINC1731684 | C₆H₅NO          | 1   | 1   | 4     | 142.123| 0.60  |
| ZINC1988907 | C₆H₅NO₂         | 2   | 1   | 4     | 311.293| 1.04  |
| ZINC1988907 | C₆H₅NO₂         | 2   | 1   | 5     | 271.262| 0.03  |
| ZINC2044196 | C₆H₅NO₂         | 2   | 3   | 3     | 199.181| 0.44  |
| ZINC2044196 | C₆H₅NO₂         | 2   | 2   | 5     | 269.283| 2.99  |
| ZINC2044196 | C₆H₅NO₂         | 2   | 3   | 6     | 231.251| -0.134|
| ZINC2044196 | C₆H₅NO₂         | 2   | 3   | 6     | 231.251| -0.134|
| Pubchem1793772 | C₆H₅NO₂       | 3   | 3   | 10    | 204.147| 1.00  |
| Pubchem18542010 | C₆H₅NO₂      | 5   | 2   | 9     | 204.099| 0.20  |
| Pubchem21811791 | C₆H₅NO₂     | 4   | 2   | 10    | 204.136| -0.028|
| Chemspider8464931 | C₆H₅NO₂S   | 4   | 3   | 7     | 268.088| 0.55  |
| Chemspider8464933 | C₆H₅NO₂S   | 4   | 3   | 7     | 268.088| 0.55  |
| Chemspider16740754 | C₆H₅NO     | 1   | 4   | 6     | 193.157| 0.13  |
| ChemBank10036 | C₆H₅NO₂      | 6   | 2   | 11    | 376.199| 0.66  |
| ChemBank1176 | C₆H₅NO₂      | 4   | 4   | 9     | 204.183| 0.118 |
| ChemBank1608 | C₆H₅NO₂      | 7   | 5   | 13    | 405.226| -1.822|
| ChemBank11000260 | C₆H₅NO₂     | 4   | 4   | 9     | 204.183| 0.118 |
| ChemBank1049255 | C₆H₅NO₂     | 4   | 4   | 5     | 256.205| 1.246 |

Table II: ADME of compounds

| Compounds   | HIA (%) | IVCEL (nm/sec) | INVMCM (nm/sec) | IVSP (log(kp, cm/hour)) | IVPPB (%) | IVBBBP (%) |
|-------------|---------|----------------|-----------------|--------------------------|-----------|------------|
| Chem2057082 | 92.492  | 53.79          | 77.47           | -2.45                    | 62.111    | 0.172      |
| Chemspider763024 | 91.515 | 1.430          | 73.606          | -3.216                   | 60.233    | 0.019      |
| ZINC0566632 | 99.302  | 49.650         | 177.164         | -1.771                   | 60.233    | 0.019      |
| ZINC0088846 | 99.027  | 25.796         | 264.592         | -3.555                   | 0.00      | 0.9054     |
| ZINC05105206 | 99.039 | 50.917         | 173.184         | -2.635                   | 0.00      | 1.115      |
| Pubchem1793772 | 78.811 | 21.325         | 8.355           | -3.54                    | 30.154    | 0.060      |
| ZINC17353697 | 87.424 | 37.884         | 227.952         | -4.869                   | 7.397     | 0.406      |
| Etha11      | 87.280  | 21.959         | 30.751          | 2.266                    | 85.488    | 5.110      |
| Etha17      | 70.69   | 0.410          | 1.71            | -5.002                   | 0.00      | 0.482      |
| ZINC0088344 | 99.039  | 25.799         | 264.59          | -3.02                    | 0.00      | 0.9050     |
| Etha10      | 82.468  | 19.289         | 0.478           | -4.203                   | 38.100    | 0.365      |

Table III: Predicted binding sites of the receptor

| Cavity | Position | X   | Y   | Z   | Volume (Å³) | Surface (Å²) |
|--------|----------|-----|-----|-----|-------------|--------------|
| 1      | 95.116   | -6.436 | 6.514 | 97.28 | 368.64      | 84.48        |
| 2      | 90.257   | -18.415 | 1.179 | 74.24 | 240.64      |
| 3      | 70.878   | -0.667 | 16.219 | 25.6  | 111.36      |
| 4      | 70.878   | -0.667 | 16.219 | 19.968 | 84.48       |

Table IV: Amino acid residues around the potential binding site

| Site       | Ser739 | Asn740 | Leu743 | Ala743 |
|------------|--------|--------|--------|--------|
|            |        |        |        |        |
|            |        |        |        |        |
|            |        |        |        |        |

The energy of the ligand. The docking energy scoring function is based on the modified piecewise linear potential (PLP) with new hydrogen bonding and electrostatic terms included.

The ligands were optimized before docking for proper structural stabilization. We calculated stretch, bend, steth, torsion, non-1,4 VDW, 1,4 VDW, total energy (Kcal/mol) using MM2 module of Chembio office tool. Docking computation was done based on the parameters mentioned in the methodology.

While docking the receptor was set rigid and docked with the receptor binding site inside the constraint...
where, bond flexibility of lignds was set as “on”. The docking result has predicted two database compounds and two analogues of ethambutol based on least energy score of rerank, moldock and H bond values as presented in the Table V. Dock poses were further inspected for hydrogen bonding interaction with the receptor. The compound Chemspider20572082 and Zinc00388344 showed highest rerank score of -117.902 and -107.278 with optimum hydrogen bonding with the receptor including two analogue of ethambutol as shown in the Table VI.

The compound Chemspider20572082 interacts with the amino acid residue Gly1058, Asp1052 and Asp1056 and forming three hydrogen bond interaction at the distance of 2.43, 2.35 and 1.99 Å respectively. ZINCOO388344 is forming three hydrogen bonds with Val1045 and Leu1060 and clearly reflecting its novelty as a inhibitor of M. tuberculosis arabinosyl transferase in compared with ethambutol having poor reranking score of -39.980.

Screening for alternative and effective drug is urgently needed to combat the drug resistance strains of M. tuberculosis (Burris, 2004; McIlennon, et al., 2009; Zhang, et al., 2014). The failure of ethambutol is another challenge. Therefore to meet the present challenges for inhibition of M. tuberculosis arabinosyl transferase by these compounds would be a useful starting point to design better therapeutics of M. tuberculosis and in vitro experiment on compounds, viz. Chem2057082 and ZINCOO388344 is recommended.

In this investigation, virtual screening has been performed using various filters. The screened compounds are subjected to molecular docking and result are analysis

| Ligand            | MolDock score | Rerank score | HBond  |
|-------------------|---------------|--------------|--------|
| Chemspider20572082| -117.902      | -93.096      | -22.399|
| Zinc00388344      | -107.278      | -80.301      | -08.963|
| Etha9             | -104.055      | -76.741      | -05.465|
| Etha17            | -94.996       | -45.846      | -05.651|
| Etha10            | -61.488       | -44.818      | -04.162|
| Ethambutol        | -55.620       | -39.980      | -12.403|

Table VI: Hydrogen bonding between ligands and receptor

| Ligand name | Ligands | Distance (Å) | Protein |
|-------------|---------|--------------|---------|
| Etha 9      | H(1) 17 | 3.25         | O(8) 1934 Val1054 |
|             | H(1) 17 | 2.54         | N(7) 1943 Scr1047 |
|             | H(1) 17 | 2.24         | O(8) 1948 Scr1047 |
|             | H(1) 23 | 2.24         | O(8) 2036 Asp1056 |
| Chem2057082 | H(1) 25 | 2.43         | O(8) 2055 Gly1058 |
|             | H(1) 30 | 2.35         | O(8) 1989 Asp1052 |
|             | H(1) 26 | 1.99         | O(8) 2036 Asp1056 |
| ZINCOO388344 | H(1) 22 | 2.12         | O(8) 1934 Val1045 |
|             | H(1) 22 | 2.22         | O(8) 2061 Leu1060 |
|             | H(1) 22 | 2.26         | O(8) 2064 Leu1060 |
| Ethambutol  | H(1) 20 | 2.03         | O(8) 1948 Ser1047 |
|             | H(1) 29 | 2.32         | O(8) 2055 Gly1058 |
|             | H(1) 29 | 3.01         | O(8) 2036 Asp1056 |
on the basic of rerank score and hydrogen bond interaction and it was found that 3 compounds showed better result out of 31 docked compound than the control drug. Further ADME and pharmacological effects of these compounds observed comparatively better bioavailability, distribution, absorption, drug likeness, and pharmacological effects than ethambutol. Hence, it could be concluded that these three compounds could be considered as potent drug candidate of *M. tuberculosis*.

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