



A Comprehensive Investigation of the Interactions between Proteins and Ligands in the Crystal Structures of Mycobacterium Tuberculosis

Rajesh Nanaware ^a, Anuruddha Chabukswar ^{a*},
Vishal Tekade ^b, Kunal Kashid ^b, Tejas Vidhate ^b
and Rushikesh Karmure ^b

^a Department of Pharmaceutical Sciences, School of Health Sciences and Technology, Dr. Vishwanath Karad MIT-World Peace University, Pune-411038, Maharashtra, India.

^b Department of Pharmacy, TPCT'S Terna College of Engineering Osmanabad-413501, Maharashtra, India.

Authors' contributions

This work was carried out in collaboration among all authors. Author RN designed the study, wrote the protocol and first draft of the manuscript. Author AC managed the analyses of the study. Authors VT, KK, TV and RK managed the literature searches. All authors read and approved the final manuscript.

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ABSTRACT

Mycobacterium tuberculosis (MTB), an acid-fast aerobic bacterium that may grow on gram stain as either a gram-positive or gram-negative bacterium, is the disease-causing agent of tuberculosis (TB). Rifampin, isoniazid, pyrazinamide and ethambutol, the first-line anti-tubercular drugs, can all have hepatotoxic side effects. The new medicine needs to work through a novel mode of action or

*Corresponding author: E-mail: anuruddha.chabukswar@mitwpu.edu.in;

to a novel target, be more active than presently available treatments, and shorten the course of treatment for the MDR-TB and XDR-TB, also active against both active and latent bacteria, and does not interact with antiretroviral medications because many TB patients also have HIV. Additionally, it must work well with other anti-TB medications to form at least an effective three-drug regimen. This article discusses the analysis of a few FDA-approved anti-tubercular medications and their binding locations with respective targeted proteins. This mainly focuses on the amino acids of the proteins that are responsible for the formation of interactions with a drug molecule. So, researchers can modify the existing drugs or their derivatives or can construct a new molecule according to the binding sites of enzymes corresponding to mycobacterium tuberculosis.

Keywords: Anti-tubercular drugs; Mycobacterium tuberculosis; MDR-TB; XDR-TB.

1. INTRODUCTION

TB is still among the deadliest infectious diseases in the world. Nearly 30,000 people become ill with this preventable and treatable disease every day, and close to 4400 people lose their lives to TB. Access to TB diagnosis and treatment as well as the burden of TB disease continue to be negatively impacted by the COVID-19 epidemic. Global TB objectives are off track and advancement made in the years leading up to 2019 has slowed, stopped, or reversed. The number of people who were infected with TB was expected to be 10.6 million (95% UI: 9.9-11 million) in 2021, an increase of 4.5% from 10.1 million (95% UI: 9.5-10.7 million) in 2020. The number of new cases of tuberculosis (TB) per 100,000 people increased by 3.6% between 2020 and 2021, reversing annual decreases of roughly 2% for most of the preceding two decades. The net decrease from 2015 to 2021 was 10%, which is barely halfway to the first End TB Strategy milestone [1]. Reductions in the number of new TB cases in 2020 and 2021 indicate more people with the disease are undiagnosed and untreated, which first leads to more TB deaths and community transmission of the infection before, after a lag, increasing the number of new cases of TB. We anticipate that the impact of COVID-19 will vary depending on the context. For instance, nations with high rates of tuberculosis (TB), like India and Vietnam, have seen substantially disparate COVID-19 occurrences [2, 3]. The Mycobacterium tuberculosis infection medications approved by the FDA include Rifampin, Isoniazid, Pyrazinamide, and Ethambutol [4]. Approximately 13 new compounds have been recently discovered, including Bedaquiline, Delamanid, Pretomanid, Contezolid, Delpazolid, Sutezolid, GSK-3036656, Macozinone, OPC-167832, Q203, SQ109, TBA-7371, and TBI-166. This occurs after almost 40 years of active research and development work

without any novel approved TB drugs. Based on their phase II clinical trials, two of these drugs bedaquiline and delamanid have been given accelerated or conditional regulatory approval [5-7]. Linezolid, Levofloxacin, Moxifloxacin, and Nitrazoxanide are the additional substances. For better TB illness therapy, more research is required. Although TB typically assaults the lungs, it can also affect the kidneys, spine, and brain [8]. Droplets from speaking, coughing, and sneezing propagate the illness. Micro-bacterial cultures, chest X-rays, and tuberculin skin tests are used to make the diagnosis [9]. The appearance of tuberculosis varies, and based on clinical manifestation, it is categorized into pulmonary TB (PTB) and extrapulmonary TB (EPTB) [10]. Because extrapulmonary TB (EPTB) accounts for more than 50% of all TB diagnoses in HIV-positive individuals, the situation is made more difficult by the HIV pandemic [11]. EPTB frequently presents a significant challenge for early diagnosis due to its heterogeneity of presentation. Constitutional symptoms including fever, anorexia, weight loss, malaise, and exhaustion may be present [12]. Drug resistance is a recurring issue in the treatment of TB. For instance, in 2016, it was reported that 490000 instances of multidrug-resistant tuberculosis (MDR-TB) and an additional 110000 cases of TB that were susceptible to isoniazid but resistant to rifampicin (RR-TB), the best first-line anti-TB medication [13]. MDR-TB is resistant to two most potent anti-TB medications, rifampicin and isoniazid. Extensively drug-resistant tuberculosis (XDR-TB), also known as MDR-TB and is resistant to at least one fluoroquinolone plus a second-line injectable medication (kanamycin, capreomycin, or amikacin), is a different form of resistant TB. In the intensive phase, which lasts for 6 to 12 months, all XDR-TB cases are treated with injections of Capreomycin, Moxifloxacin (MIFX), PAS high-dose Isoniazid, clofazimine, Linezolid, and Co-amoxiclav. The rest of the drugs, except

for the injections, are continued for 18 months in the continuation phase. The following obstacles must be overcome by researchers working in the field of TB treatment, among others: the newer drug must be more effective than currently available to shorten the length of treatment; it must also be active against both active and latent bacteria; it must act by a novel mechanism of action or to a novel target, particularly for the treatment of MDR-TB and XDR-TB; and it must not interact with antiretroviral drugs because many TB patients also take these drugs [14,15]. In this review, 14-FDA-approved Anti-tubercular drugs obtained from different literature and their protein-ligand interactions are discussed.

2. FDA-APPROVED DRUGS USED IN TB

2.1 Bedaquiline

For the treatment of tuberculosis, bedaquiline, a diarylquinoline, is offered under the trade name Sirturo. Both the FDA and the EMA authorized bedaquiline for the treatment of MDR-TB in December 2012 and March 2014, respectively [16-20].

Mechanism of Action: Bedaquiline is extremely effective and works through a novel mechanism that involves inhibition of the proton pump of mycobacterial ATP synthase [21].

Binding Pockets: In their Nucleic Acid Research publication, Kailu Yang and Junjie Zhang

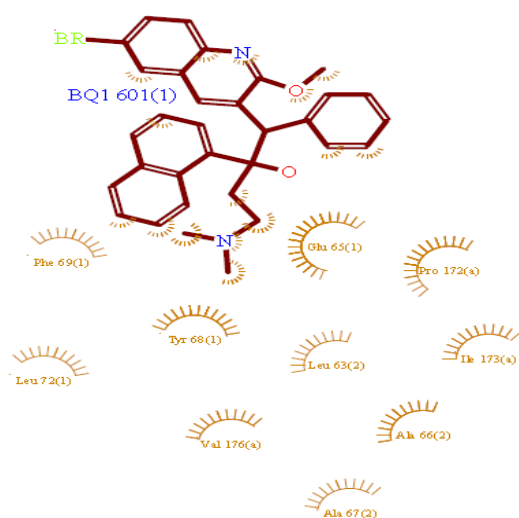
reported that Bedaquiline has hydrophobic connections with PHE-691, LEU-721, PRO-172, VAL-176, GLU-654, ILE-705, ALA-669, and TYR-681 [22].

2.2 Delamanid

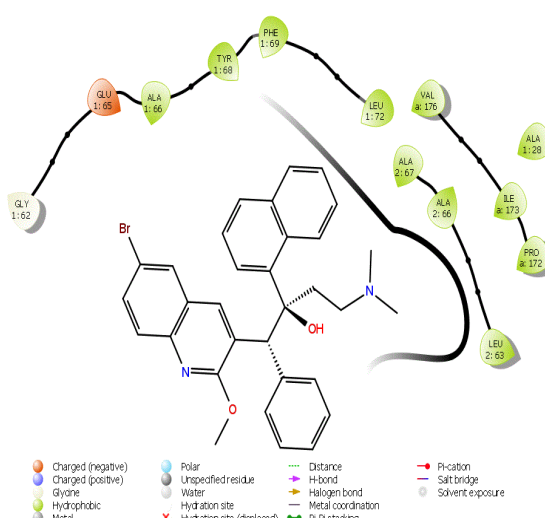
Delamanid, also marketed as Deltyba, is a derivative of nitroimidazooxazole (OPC-67683). On April 28, 2014, the EMA granted Delamanid marketing authorization for use.

Mechanism of Action: Delamanid works by preventing mycolic acid synthesis. A Keto-mycolic and methoxy-mycolic acid, two crucial elements of the *M. tuberculosis* cell wall, are inhibited, which results in bacterial cell death [23].

Binding pockets: Cellitti S.E. came to the conclusion in their study that the Delamanid exhibits hydrophobic and other interactions with LYS-55A, ARG-60A, PRO-63A, LEU-64A, and MET-87A, VAL-46A, LYS-55A, THR-56A, ASN-62A, PRO-63A, TYR-65A, ALA-76A, LYS-79A, SER-78A, MET-87A, TRP-88A, and A5N-91A are other molecules with which the delamanid formed H-bonds. Delamanid also creates the TYR-65A that stacks. Delamanid joins forces with GLY-53A, TYR-133A, LYS-104A, ARG-54A, and PRO-86A to form hydrophobic and other connections [24]. Liglot and 2D complexes of bedaquiline and delamanid are shown in Fig. 1.



A



B

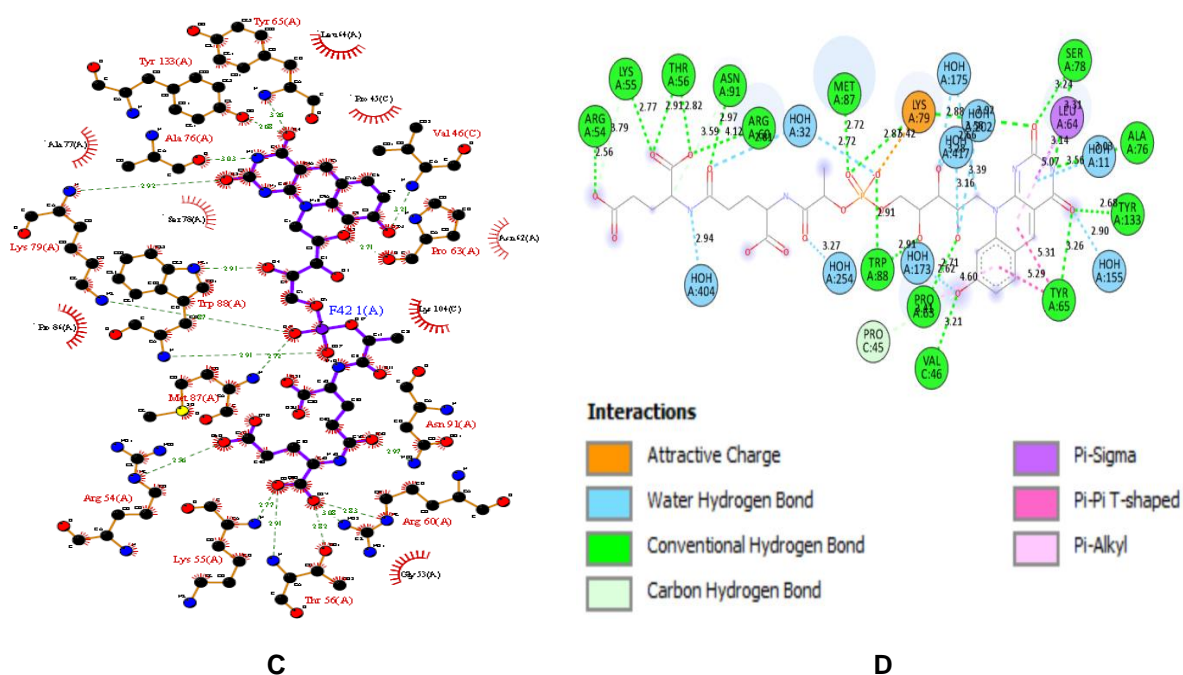


Fig. 1. A) Ligplot and B) 2D Complex of Bedaquiline with 7JGC. C) Ligplot and D) 2D Complex of Delamanid with 3R5R.

2.3 Pretomanid

Pretomanid, also known as PA-824, is a nitroimidazole medication that is presently undergoing clinical studies to see whether it has any effect on both drug-susceptible and drug-resistant TB. Pretomanid, which shortens the duration of TB treatment, surpassed rifampicin, and pyrazinamide in terms of its ability to kill non-replicating Mycobacterium tuberculosis [25,26].

Mechanism of Action: The suppression of lipid and protein synthesis is its mode of action. It prevents the formation of the mycolates found in mycobacterial cell walls [27]. A recent investigation on *M. smegmatis*, a species that is genetically related to MTB, however, revealed that pretomanid may operate instead by interfering with bacterial sugar phosphate metabolism and causing the bacteria to produce the deadly substance methylglyoxal [28].

Binding pockets: Barry C. Finzel and Courtney C. Aldrich in the Journal of Medicinal concluded that pretomanid exhibits hydrophobic interactions with PRO-24A and TYR-25A. Pretomanid stack with TRP-64A at various distances and TYR-157A. Additionally, GLY-156A forms halogen bonds with it. Together with ILE-256A, ALA-226A, GLY-227A, PRO-24A, MET-91A, THR-

318B, and PH-92B, it also forms other bonds [29].

2.4 Contezolid

The first oxazolidinone antibiotic to show potential for use in the management of infections driven on by Gram-positive pathogens was contezolid or MRX-I [30].

Mechanism of Action: In general, oxyazolidinones work by preventing the production of new proteins. It was believed that they accomplished this by obstructing translational integrity [31,32]. Fig. 2 depicts the Liglot and 2D complexes of pretomanid and contezolid.

2.5 Sutezolid

An oxazolidinone derivative of linezolid called sutezolid (PNU-100480 or U-100480) is now being tested in clinical studies to treat tuberculosis. In a murine model, sutezolid was found to be more effective than linezolid, and its combination with I and II-line anti-TB medications demonstrated excellent bactericidal action with the ability to accelerate therapy.

Mechanism of Action: Sutezolid inhibits protein synthesis [33-35].

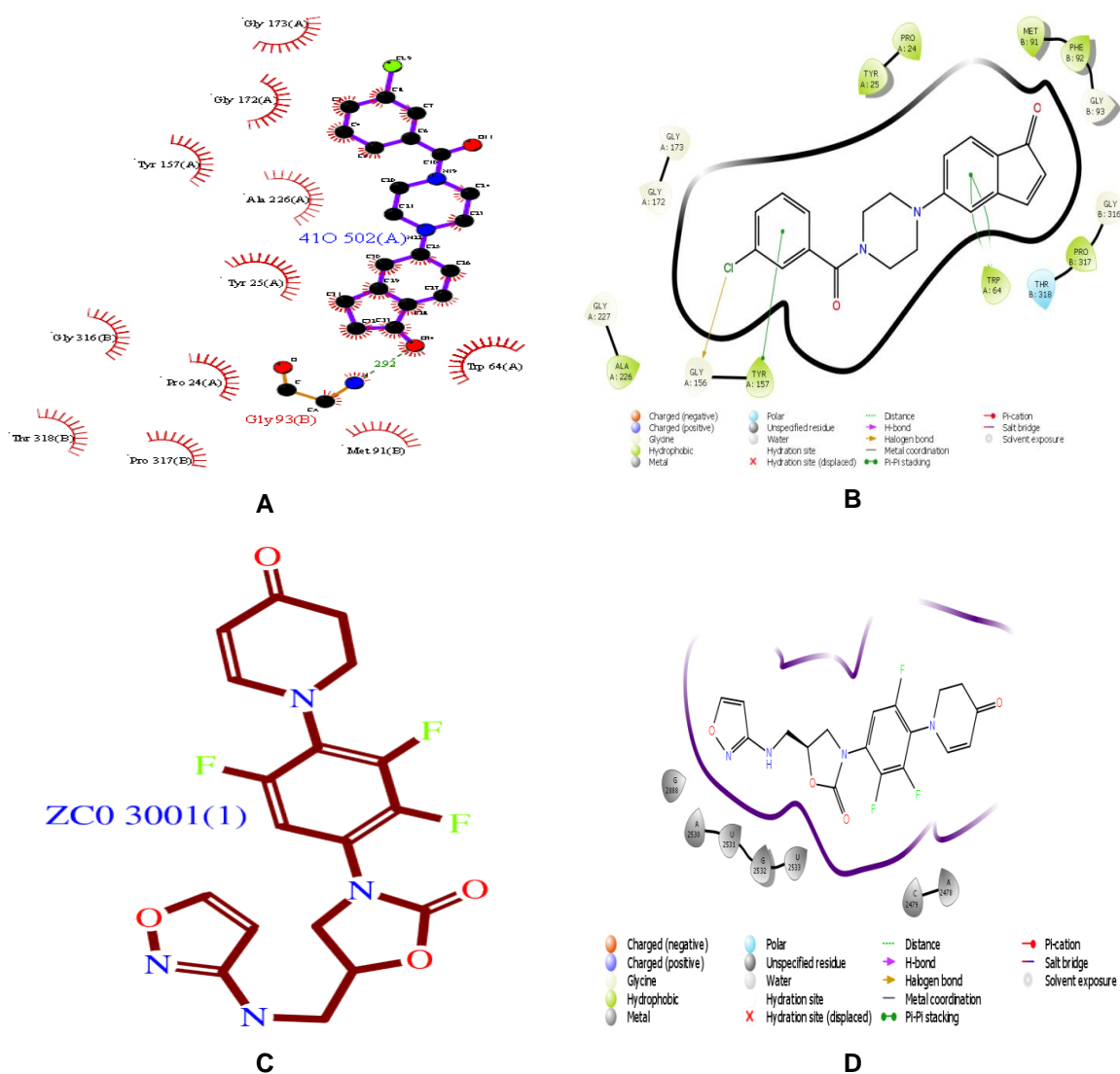


Fig. 2. A) Ligplot and B) 2D Complex of Pretomanid with 4XJO. C) Ligplot and D) 2D Complex of Contezolid with 6WQN.

Binding Pockets: According to the research paper by Mendes, V. Sutezolid formed hydrophobic and other interactions with ASN-179A, TRP-103, PHE-110-GLU-180, LEU-183, PHE-114, and MET-142, TRP-145, TRY-148, THR-149, TRP-138, ILE107, GLY-106, and LEU-87 [36].

2.6 Macozinone

Macozinone did not interact negatively or favorably with I-line or II-line drugs in a trial conducted by Lupine and colleagues. Macozinone had improved bactericidal action in the lungs and spleen that was comparable to isoniazid when tested in a chronic TB murine model.

Mechanism of Action: By binding covalently to a cysteine residue in the active site of decaprenylphosphoryl-b-D-ribose 2'-epimerase (DprE1) inhibits the formation of cell walls [37].

Binding pockets: According to the research paper by Vadim Makaro and Stewart T. Cole, Macozinone exhibits hydrophobic interactions with VAL-365A. Macozinone joins LYS-134A and LYS-418A to produce H-bonds. It also displays ASN-385A halogen bonding. Additionally, it displays interactions with SER-228A, GLY-133A, HIS-132A, PHE-369A, GLY-117A, and CYS-387A as well as additional interactions [38]. Fig. 3 depicts the Liglot and 2D complexes of sutezolid and macozinone.

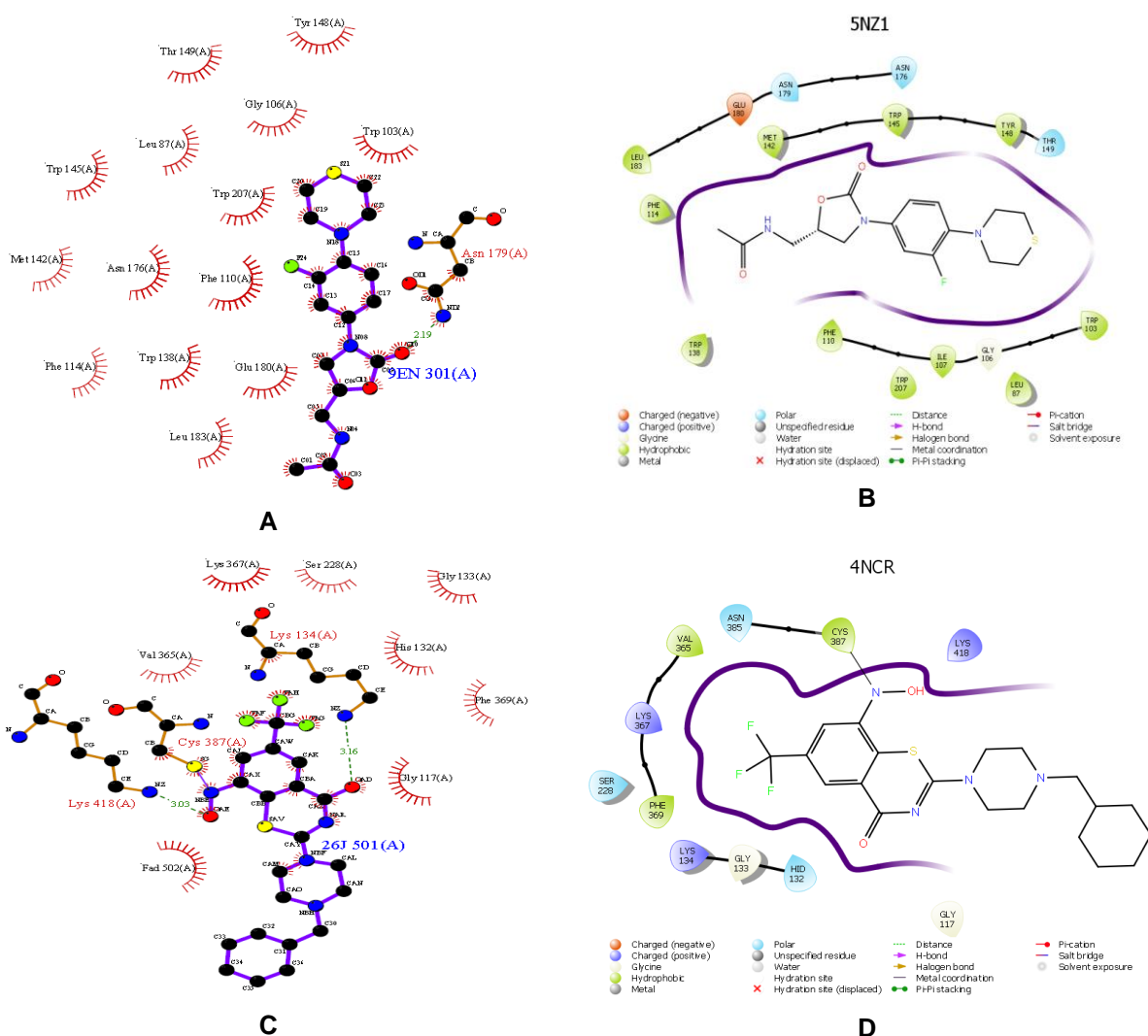


Fig. 3. A) Ligplot and B) 2D Complex of Sutezolid with 5ZN1. C) Ligplot and D) 2D Complex of Macozinone with 4NCR.

2.7 Linezolid

Linezolid exhibits significant action against *M. tuberculosis* that is rapidly developing, slowly growing, and incapable of reproducing itself, demonstrating its potential to be used in TB therapy regimens that are shorter in duration [39].

Mechanism of Action: An oxazolidinone antibiotic called linezolid works to suppress translation by attaching to the ribosome's 50S subunit and competing with the cell's natural substrates [40-41].

Binding pockets: According to the research paper by Mendis V., Linezolid exhibits hydrophobic and other interactions with ASN-179, ASN-176, VAL-152, THR-149, TYR-148,

TRP-145, MET-142, LEU-183, MET-102, TRP-103, TRP-207, ILE-107 and PHE-110 [38].

2.8 Levofloxacin

The L-isomer of the D, L-racemate ofloxacin, levofloxacin is an oral fluoroquinolone antibiotic that was created in Japan. Levofloxacin has a tolerability profile that is comparable to other oral fluoroquinolones, with gastrointestinal and central nervous system side effects being the most frequent. The pharmacokinetics of levofloxacin is not influenced by food, age, or gender.

Mechanism of Action: Modifications to the cell membrane porin channels or the DNA gyrase A or B subunits are two examples of chromosomal abnormalities that might cause bacterial resistance to levofloxacin [42-46].

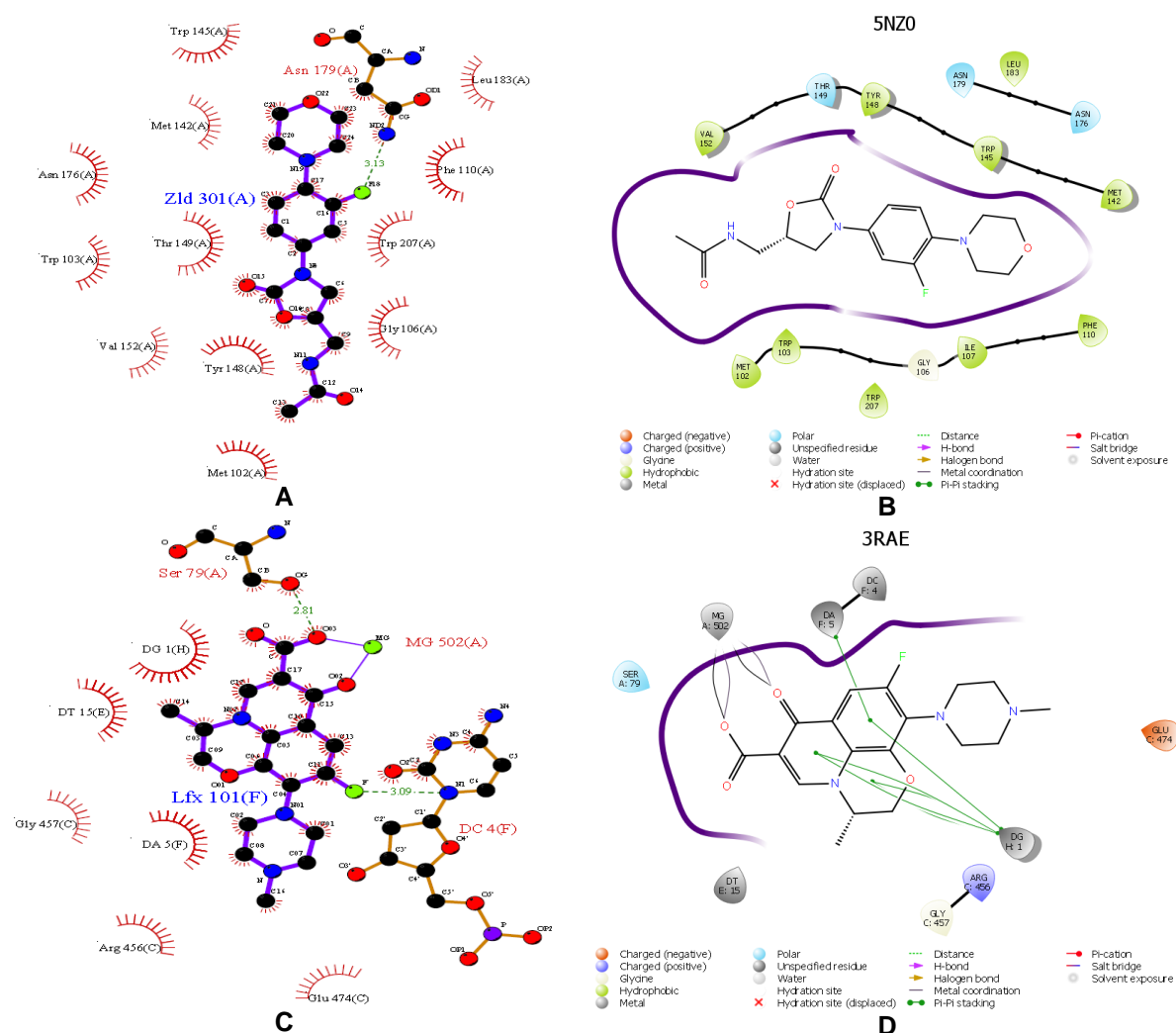


Fig. 4. A) Ligplot and B) 2D Complex of Linezolid with 5NZ0. C) Ligplot and D) 2D Complex of Levofloxacin with 3RAE.

Binding pockets: Levofloxacin is used in the research work by L. Mark Fisher and Mark R. Sanderson, and they discovered that it exhibits hydrophobic interactions with ILE-170A. Along with ARG-117A, GLY-173A, ALA-175A, ASN-326A, SER-436D, LYS-458C, ASN-461C, and ASN-473, it also forms H-bonds with them [47]. Fig. 4 displays the linezolid and levofloxacin liglot and 2D complexes.

2.9 Moxifloxacin

A fluoroquinolone antibiotic that exhibits a broad spectrum of activity, moxifloxacin is used to treat endocarditis, sinusitis, pneumonia, and TB [48,49].

Mechanism of Action: Moxifloxacin inhibits the synthesis of bacterial DNA [50-52].

Binding pockets: In their article, Pan F. Chan, Velupillai Srikanthasan, and Jianzhong Huang concluded that moxifloxacin exhibits hydrophobic interactions as well as others with ARG-122A, GLY-459B, SER-84A, and GLY-473B [53].

2.10 Rifampicin

Rifampicin, also known as rifampin, was developed in 1968 and is still one of the main medications used to treat tuberculosis (TB) because of its sterilizing properties and capacity to accelerate recovery times when used at high doses [54-58]. To treat patients with pulmonary TB who were chronically resistant to treatment, rifampicin was initially used in clinical settings in the late 1960s [59-61].

Mechanism of Action: The enzyme responsible for DNA transcription in bacteria, RNA polymerase, is inhibited by rifampicin [62].

Binding pockets: Gerard D. Wright and his research team demonstrated in their study, "Rifamycin Resistance Enzyme with an Unprecedented Mechanism of Action," that rifampicin interacts hydrophobically with LEU-176A, PHE-257A, and PRO-284A. The H-bonds with GLY-9A, THR-285A, ARG-213A, MET-205, ALA-204, VAL-215, ARG-201, MET-342, GLY-286, GLY-287, PHE-74, VAL-93, GLY-44, GLN-43, VAL-69 and ARG-196 are also shown [63].

2.11 Isoniazid

This synthetic drug was first designated by Meller and Melley in the year 1912. The antibiotic isoniazid, also referred to as isonicotinic acid hydrazide, is used to treat tuberculosis. It is often combined with rifampicin, pyrazinamide, and either streptomycin or ethambutol to treat active

tuberculosis. It is frequently used alone to treat latent tuberculosis.

Mechanism of Action: Isoniazid inhibits mycolic acid biosynthesis [64].

Binding pockets: According to Gegia M.'s study, the isoniazid interacts hydrophobically with PHE-78, PRO-79, ALA-80, LEU-88, and TYR-85A. Instead, it displays an H-bond with the amino acid THR-307A. It displays other interactions with THR307, HIS-269, and HIS-84 [65]. Fig. 5 displays the moxifloxacin, rifampicin and isoniazid liglot and 2D complexes.

2.12 Pyrazinamide

Because of its unique method of action (interference with ATP synthesis), pyrazinamide will probably continue to play a significant role in the treatment of drug-susceptible and multidrug-resistant TB (MDR-TB).

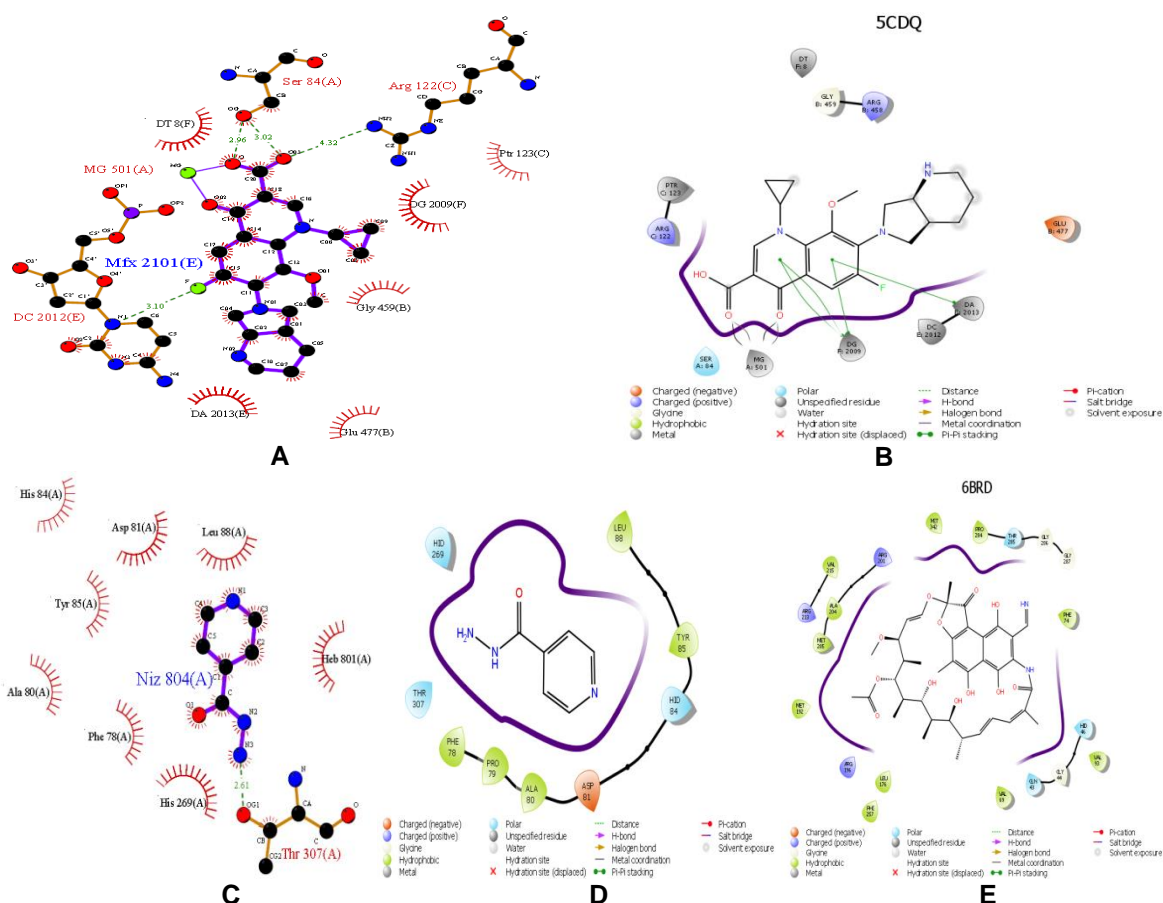


Fig. 5. A) Ligplot and B) 2D Complex of Moxifloxacin with 5CDQ. C) Ligplot of Isoniazid complexed with 4PAE. D & E) 2D Complex of Isoniazid with 4PAE & Rifampicin with 6BRD.

Mechanism of Action: Pyrazinamide inhibits the growth of bacteria [66-67].

Binding pockets: According to Petrella, S. study published in the PLoS One journal, pyrazinamide exhibits interactions with the following amino acids: ARG-255A, HIS-109, GLN-105A, ASP-108A, GLU-258, and PRA-598 [68].

2.13 Ethambutol

The main purpose of the drug ethambutol is to cure tuberculosis. It is typically administered along with other anti-tuberculosis drugs such as isoniazid, rifampicin, and pyrazinamide. Mycobacterium avium complex and Mycobacterium kansasii may also be treated with it.

Mechanism of Action: The bacteriostatic medication ethambutol prevents bacilli from reproducing by interfering with the formation of arabinogalactan in the cell wall [69].

Binding pockets: In their article, Zhang, L. demonstrated that ethambutol has hydrophobic and other TRP interactions with VAL-1004A, ASP-279A, TRP-965, TRP-572, TYR-282, ILE-283, TYR-314 and ARG-383 [70].

2.14 Para-amino Salicylic Acid

In the 1940s, it was discovered that para-amino salicylic acid (PAS), a second-line anti-TB medication (SLD), was useful for treating tuberculosis.

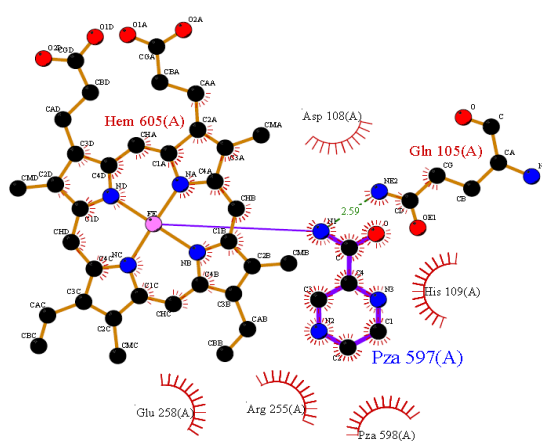
Mechanism of Action: Para-amino salicylic acid inhibits folic acid synthesis [71].

Binding pockets: In their article, Schreuder, H. A. demonstrated that the para-amino salicylic acid exhibits interactions with TRP-185, TYR-222, PRO293, ALA-296, LEU-210, TYR-201, LEU-199, VAL-47, ARG-214, SER-212, THR-296 and ARG-44 that are hydrophobic and others [72]. Fig. 6 displays the pyrazinamide, ethambutol and para-amino salicylic acid liglot and 2D complexes.

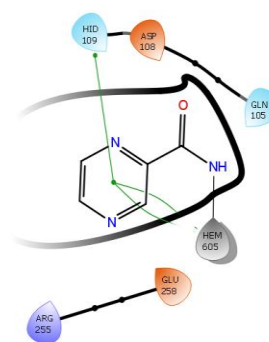
3. RESULTS AND DISCUSSION

The 3D crystal structures of all the enzymes were retrieved from the protein data bank-RCSB (<http://www.rcsb.org>). For interpretation and visualization of interactions of protein-ligand complex, Mastero and Biovia discovery studio visualizer and freely accessible server PDBsum (<http://www.ebi.ac.uk/pdbsum>) for Ligplot were used [73].

Bedaquiline forms non-bonded contacts with amino acids of 7JGC protein. The oxygen and carbons of amino acid proline-172A form non-bonded contacts with carbon and oxygen atoms of bedaquiline with varying distances (3.05-3.65Å). Isoleucine-173A forms two non-bonded contacts with O and C atoms of bedaquiline with a distance of 3.05-3.24Å. Valine-176A formed two interactions with carbons having distances 3.51 and 3.35Å. Five different number carbons of Glutamic acid-65 interacts with C and N atoms of bedaquiline and oxygen atoms of seven different positions of Glutamic acid-65 form interactions with C and O atoms of the molecule BQ with varying distances. The Tyrosine-68 residue forms non-bonded contacts with different C and N atoms of bedaquiline. Also, C and O atoms of Alanine-66 and 67, Leucine-63 and 72, and Phenylalanine-69 residue create interactions with C atoms of BQ (distances ranging from 3.10 to 3.90 Å).



A



B



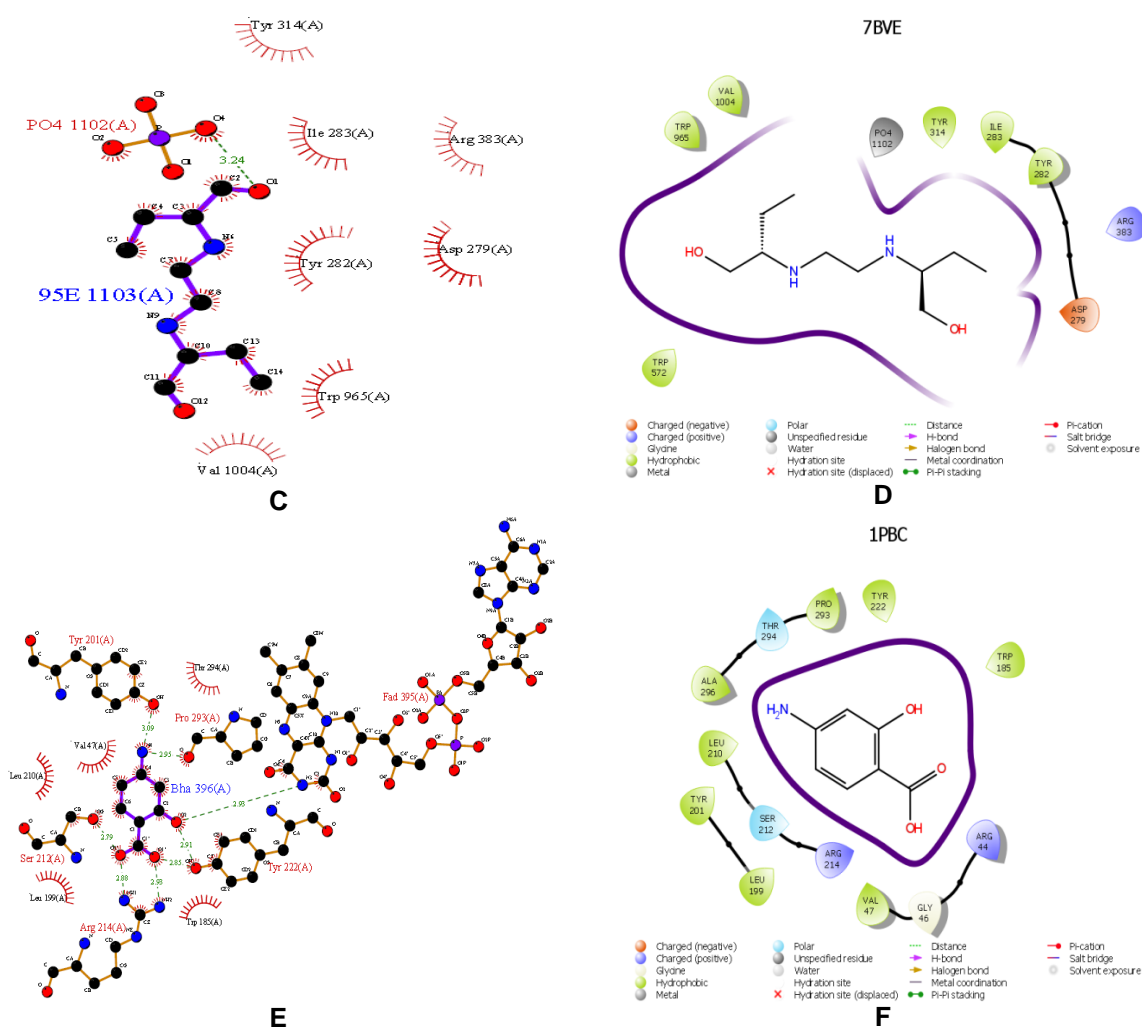


Fig. 6. A) Ligplot and B) 2D Complex of Pyrazinamide with 3PL1. C) Ligplot and D) 2D Complex of Ethambutol with 7BVE. E) Ligplot and F) 2D Complex of PAS with 1PBC.

The drug delamanid formed 18 (hydrogen bonds) bonded and 140 non-bonded interactions with 3R5R. O and N atoms of amino acids from A chain Arginine-54 and 60, Lysine-55 and 79, Threonine-56, Proline-63, Tyrosine-65 and 133, Alanine-76, Methionine-87, Tryptophan-88, Asparagine-91 and Valine-46 formed hydrogen bonds (having distance ranges from 2.56 to 3.26Å) with different number of O and N atoms of delamanid molecule. Pretomanid forms one hydrogen bond and 62 non-bonded interactions with protein 4XJO. Nitrogen atom of Glycine-93B formed hydrogen bond with oxygen-34 atom of pretomanid molecule with a distance of 2.92Å. In non-bonded interactions of Proline-24A and 317B, Tyrosine-25A, Tryptophan-64A, Tyrosine-157A, Glycine-172A, 316B and 93, Methionine-91, Threonine-318B amino acids were involved. Sutezolid forms one hydrogen bond and 67 non-bonded interactions with protein 5NZ1. Nitrogen

atom of Asparagine-179A formed hydrogen bond with oxygen-10 of sutezolid molecule with a distance of 2.19Å. Macozinone molecule forms 3-(hydrogen bonds) bonded and 33 non-bonded contacts with protein 4NCR. Nitrogen atom of Lysine-134A and 418A interacts with the oxygen of the molecule with a distance of 3.16Å and 3.03 respectively. The S atom of Cystine-187A formed a hydrogen bond with N atom with a distance of 1.66 Å.

Linezolid forms a single hydrogen bond and 53 non-bonded interactions with protein 5NZ0. Nitrogen atom of Asparagine-179A formed hydrogen bond with the C-18 atom of a linezolid molecule with a distance of 3.13Å. MET-102A, TRP-103A, GLY-106A, ILE-107A, PHE-110A, TRP-145A, TYR-148A, THR-149A, ASN-176A, ASN179A and TRP-207A residues contributes to non-bonded contacts. Levofloxacin

Table 1. FDA Approved Antitubercular drugs MOA and their PDB details

S.N.	Drug	PDB Code	Classification	Organism	Mutation	Resolution	Mechanism of Action
1	Bedaquiline	7JGC	Hydrolase	M. Smegmatis	No	3.40Å°	Inhibits the Proton pump ATP Synthase
2	Delamanid	3R5R	Oxidoreductase	M. tuberculosis	No	2-10Å°	Inhibits mycolic acid biosynthesis.
3	Pretomanid	4XJO	Transferase inhibitor	M. tuberculosis	No	1.50 Å°	Inhibition of mycolic acid synthesis
4	Contezolid	6WQN	Ribosome	S. aureus	Yes	2.90Å°	Inhibits protein synthesis.
5	Sutezolid	5NZ1	Transcription	M. Tuberculosis	No	2.33 Å	Inhibits protein synthesis.
6	Macozinone	4NCR	Oxidoreductase	M. Tuberculosis	No	1.88 Å°	By forming a covalent bond with a cysteine residue, it prevents the synthesis of the cell wall.
7	Linezolid	5NZ0	Transcription	M. tuberculosis	No	1.82 Å	Inhibits translocation by binding to the 50s subunits of ribosomes.
8	Levofloxacin	3RAE	Isomerase/ DNA/Antibiotic	S. pneumoniae	No	2.90Å°	Inhibits DNA gyrase subunit (Gyrase A & Gyrase B) topoisomerase iv subunit (Par C & Part E).
9	Moxifloxacin	5CDQ	Hydrolase	S.aureussubaureus	No	2.95Å°	Inhibits DNA gyrase subunit (Gyrase A & Gyrase B) topoisomerase iv subunit (Par C &Part r E).
10	Rifampicin	6BRD	Oxidoreductase	S.venezuelige	No	3.32 Å°	Inhibits RNA polymerase.
11	Isoniazid	4PAE	Oxidoreductase	Synechococcus elongatus	Yes	3.21 Å	Inhibit synthesis of mycolic acid.
12	Pyrazinamide	3PL1	Oxidoreductase	A. flavus	Yes	1.68 Å°	Inhibits the growth of bacteria.
13	Ethambutol	7BVE	Transferase	M. Smegmatis	No	2.81 Å	Inhibition of arabinosyl transferase.
14	PAS	1PBC	Oxidoreductase	Pseudomonas fluorescens	No	2.80 Å	Inhibition of folic acid synthesis.

forms a hydrogen bond and 87 non-bonded interactions with protein 3RAE. The oxygen atom of Serine-79A formed a hydrogen bond with O-18 atom of the levofloxacin molecule with a distance of 2.81Å. Moxifloxacin molecule forms 3-(H-bond) bonded and 87 non-bonded contacts with protein 5CDQ. Nitrogen atom of Lysine-134A and 418A interacts with the oxygen of the molecule with a distance of 3.16 and 3.03Å respectively. Rifampicin formed 2 hydrogen bonds and 52 non-bonded interactions with protein 6BRD. Nitrogen from Arginine-196 and 216A binds with oxygen atoms no.8 and 2 of the rifampicin molecule with distances 2.67 and 3.18Å respectively. Isoniazid formed one hydrogen bond and 53 non-bonded contacts with protein 4PAE. Amino acid Thyronine-307A binds with N-3 of the Isoniazid molecule with a distance of 2.61Å. Pyrazinamide formed 1 hydrogen bond and 5 non-bonded contacts with protein 3PL1. Amino acid Glanine-105A formed a hydrogen bond with N of pyrazinamide molecule with a distance of 2.59Å. Ethambutol formed 1 hydrogen bond and 20 non-bonded interactions with the protein 7BVE. Aspartic acid-279A formed a hydrogen bond with the nitrogen of ethambutol with a distance of 5.14Å. Para-amino salicylic acid formed 10 hydrogens and 42 non-bonded contacts with the protein 1PBC. The residues involved in the hydrogen bonds are Tyrosine-201 and 222A, Serine-212A, Arginine-214 and Proline-293A. The information about crystal structures of the entire protein-ligand complexes was summarized in Table 1.

4. CONCLUSION

Collectively, our research offers a structural basis for understanding the significance of binding sites of amino acids of a particular protein. It serves as a useful tool for understanding the relationship between structure and function. Based on significant binding pockets, including donors and acceptors of amino acid residues, the researchers can investigate the newer molecular architectures for treating M. tuberculosis. The main finding is that future research using combined ligands and receptor-based design will generate a new class of anti-tubercular drugs that will benefit millions of patients.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

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