Review

Structure-based in silico approaches for drug discovery against *Mycobacterium tuberculosis*

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**Abstract**

*Mycobacterium tuberculosis* is the causative agent of TB and was estimated to cause 1.4 million death in 2019, alongside 10 million new infections. Drug resistance is a growing issue, with multi-drug resistant infections representing 3.3% of all new infections; hence novel antimycobacterial drugs are urgently required to combat this growing health emergency. Alongside this, increased knowledge of gene essentiality in the pathogenic organism and larger compound databases can aid in the discovery of new drug compounds. The number of protein structures, X-ray based and modelled, is increasing and now accounts for greater than > 80% of all predicted *M. tuberculosis* proteins; allowing novel targets to be investigated.

This review will focus on structure-based in silico approaches for drug discovery, covering a range of complexities and computational demands, with associated antimycobacterial examples. This includes molecular docking, molecular dynamic simulations, ensemble docking and free energy calculations. Applications of machine learning onto each of these approaches will be discussed. The need for experimental validation of computational hits is an essential component, which is unfortunately missing from many current studies. The future outlooks of these approaches will also be discussed.

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**Abbreviations:** cMD, Classical Molecular Dynamic; cryo-EM, cryogenic electron microscopy; CV, collective variable; LIE, Linear Interaction Energy; MD, Molecular Dynamic; MDR, multi-drug resistant; MMPB(GB)SA, Molecular Mechanics with Poisson Boltzmann (or generalised Born) and Surface Area solvation; Mt, Mycobacterium tuberculosis; ns, nanosecond; PTC, peptidyl transferase centre; RMSD, root-mean square-deviation; Tuberculosis, TB.

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1. Introduction

*Mycobacterium tuberculosis*, the causative agent of tuberculosis (TB), contributed to an estimated 1.4 million deaths in 2019, with approximately 10 million new infections in the same year [1]. Also, it is predicted that *M. tuberculosis* latently infects approximately one-third of the world’s population [1,2] and resistance to the current drug-treatment regime is also on the rise, with 3.3% of new cases being multi-drug resistant (MDR); this number increases drastically to 17.7% for previously treated infections [1]. If this global epidemic is to be stopped, it requires the identification and exploitation of novel drug targets, alongside other preventative approaches and treatment options [1,3].

The development of new antimycobacterial drugs is particularly challenging, in part due to the unique adaptations that *M. tuberculosis* employs which are not present in other bacterial species. The unique mycobacterial cell envelope structure, composed of modified peptidoglycan, mycolic acids and arabinogalactan, provides a waxy hydrophobic barrier which prevents penetration of several antibiotics [4,5]. In addition, *M. tuberculosis* can enter a hypoxia-induced latent growth-state, characterised by reduced metabolic activity [2,3]. This has been coupled to lower efficacy of several antibiotics, including isoniazid and beta-lactams, as their killing activity relies on active growth or metabolism [6].

The four front-line antimycobacterial drugs in current use (ethambutol, isoniazid, pyrazinamide and rifampicin), were all discovered and developed through traditional compound screening experimental methodologies [7–9]. These studies resulted in the development of ethambutol from polyamines, isoniazid and pyrazinamide from nicotinamide and rifampicin from rifamycin [7–9]. In addition, drug repurposing studies have led to the identification of many second-line antimycobacterial drugs, including fluoroquinolones, linezolid and clofazimine [10]. Repurposed drugs also represent one-third of all the new TB drugs currently in clinical trials [11]. These phenotypic drug-to-target approaches have continued to be used to successfully identify new drugs, such as delamanid and pretomanid from nitroimidazooxazoles [12,13].

However, the screening of large compound libraries is financially expensive and high re-discovery rates coupled with fewer novel hits per high-throughput screen, demonstrates that alternative approaches are required for the discovery and development of new anti-TB therapies. In this regard, the use of computational approaches for initial virtual screening, followed by concurrent experimental and computational analysis has the potential to reduce costs and increase the quality of compounds taken forward towards the developmental pipeline.

To date, two conventional computational approaches are utilised for drug discovery/repurposing projects which are either, ligand-based [14] or structure-based [15–17]. The former primarily focuses on data mining of chemical structures and associated biological activity, while the latter is concerned with the interactions of potential drugs with targets of biological interest. Both approaches aim to find chemical structures which are the most active against a particular target/organism, however, structure-based approaches have greater potential to find novel chemical structures [18].

This review focuses upon structure-based methods related to anti-TB drug discovery efforts. Several different in silico approaches will be covered, across a range of complexities and computational demands, and recent examples of their application to target *M. tuberculosis* highlighted. The application of machine-learning on several of these approaches will also be covered, alongside the increased need to perform experimental validation on computational predictions. However, before structure-based approaches can be undertaken, the selection of a target of interest and a chemical compound library to screen is essential [16,17], hence, these will be briefly covered.

2. Protein target selection and structures

Drug target selection is a major challenge in the field of drug discovery, as it usually requires a detailed understanding of the biological role and molecular genetics associated with genes that are required for bacterial survival or establishment of infection. Therefore, a common approach of target-based drug discovery research is to focus on only essential *M. tuberculosis* genes. In this regard, several highly useful studies detailing *M. tuberculosis* gene essentiality have provided guidance to the field [19,20].

Once a protein drug-target has been identified, protein structures required for downstream screening can be obtained in several ways, including crystallographic methods, cryo-electron microscopy (cryo-EM) and homology modelling. Crystallographic methods are labour intensive and produce an average protein structure, normally utilising X-rays to solve experimentally obtained protein crystals. Cryo-EM is a more recent development, which rapidly freezes proteins in aqueous environments, trapping them in ice crystals, and then uses transmission electron microscopy to solve the structures. This allows structural determination of proteins which do not readily crystallise, including membrane proteins. Homology modelling is a computational approach which uses the primary protein sequence and known crystal structures of homologous proteins, to generate the most likely protein structure. In addition, newer ab initio methods are rapidly increasing in accuracy, such as AlphaFold [21], and these deep learning approaches may dominate computational methods in the future. However, crystallographic methods are currently still the preferred approach due to their accuracy and experimental validation.

Crystal structures are currently available for a large number of *M. tuberculosis* proteins within Protein Data Bank [22], with 2,630 structures based on X-ray diffraction and a further 41 derived from electron microscopy. The great quantity of *M. tuberculosis* protein structures is in part due to the effort of the TB-structural genomics consortium [23]. If the crystal structure is not available, then the SWISS-MODEL repository [24] provides an alternative, containing a collection of modelled structures, using ProMod3 to perform homology modelling [25]. *M. tuberculosis* is one of the core species for which new models are generated and updated on a weekly basis, to account for new crystal structures, and currently 3,366 protein-encoding sequences out of a predicted 3,993 have modelled structures [24]. Alternative services for structural modelling...
include I-TASSER [26] and Phyre2 [27]; they utilise different methods to ProMod3, and hence may provide alternative structures. The use of both crystal structures and models can allow the majority of M. tuberculosis proteins to be utilised for drug screening and further structural studies.

3. Chemical libraries – virtual and tangible

Chemical libraries used for in silico screening can either be taken from databases containing modelled chemical structures or derived from tangible libraries normally utilised for high-throughput screening. The latter provides a pre-selected set of compounds, whereas the former approach generally requires selection of a sub-set of compounds. Hence, the compound libraries chosen can focus on either a broad-range of physiochemical properties, structural diversity, TB-specific compounds, or compounds for drug-repurposing. The type of compounds selected may also be influenced by the target of interest or the aim of the in silico screen, such as novel compound identification or drug-repurposing.

If the main goal of a virtual screening campaign is to identify a novel chemical compound with a strong predicted binding affinity, then large compound databases will contain the greatest structural diversity. These include the ZINC database, approximately 230 million compounds [28], the ChEMBL database, approximately 2 million compounds [29] and the Enamine REAL database, approximately 1.4 billion compounds [30], among many others [31]. These large general compound databases may be ideal for searching for unique compound structures; however, they have two main limitations; 1. their size might make them too computationally demanding for screening, 2. antimycobacterial drugs have been shown to possess different physiochemical properties compared to ‘typical’ drugs [32]. As these compound databases are normally generated around ‘typical’ drug physiochemical properties, they may not contain many compounds which are suitable for targeting M. tuberculosis. Hence, TB-specific compound libraries may provide a more focussed effort. These include the CDD-TB library of compounds, approximately 7,000 compounds [33] and the WuXi antituberculosis library, approximately 10,000 compounds [34]. The former is a virtual set of chemical compounds, whereas the latter is a physical compound library typically used in high-throughput screens which could be adapted for in silico use [34].

Drug repurposing provides an alternative strategy for drug-screening studies, utilising specific compound libraries containing only clinically approved drugs, which are also commercially available. One obvious inherent limitation of these focussed drug libraries is reduced compound structural diversity. However, since all drugs in clinical use come with an abundance of in vivo clinical data (absorption, distribution, metabolism, excretion, and toxicity etc), one clear potential benefit of screening repurposed libraries is the rapid expedition of hits towards the clinic. Examples of such libraries include the Prestwick library, 1,520 compounds [35] and the Broad Institute drug repurposing library, 6,798 compounds [36]. For the former, the molecules’ 3D structures are available from the ZINC database [28], whereas for the latter, the SMILES strings can be obtained from the Drug Repurposing Hub [36].

Selecting a suitable compound library for virtual screening largely depends on which chemical properties are the focus of the study and the parameters imposed to select for hits. Once a target protein and compound library have been selected, a variety of in silico approaches can be employed to identify compounds that bind the protein target.

4. Molecular docking

4.1. Molecular docking approaches

Molecular docking is a computational process which aims to study the interactions occurring between a protein and compound of interest. Various docking programmes exist based on this premise, but they all use different algorithms to try and fit a compound into the binding site of a protein [15]. They generate several potential conformations, or poses, which are then scored, and the top scoring poses, in a ranked order, are the main output [15]. These programmes focus on the compounds’ flexibility while treating the protein as completely rigid [37,38]; this approach saves computational power, but also decreases the accuracy [39].

The range of docking programmes available includes; GOLD [40,41]; AutoDOCK Vina [42]; Glide [43,44] and PharmScreen [45], amongst several others [15,46,47]. These programmes display diversity in both their search and score methods. They also differ in their availability due to licensing [48], with the majority being restricted to commercial use only.

As each of these programmes use different approaches for generating docking poses, their efficacy can be difficult to compare. Most programmes have been shown to generate binding site poses similar to crystallographic structures, but with varying abilities to reproduce binding affinity data [47]. A previous study found GOLD and Glide to be superior to other docking programmes [46], however many docking programmes, including AutoDOCK Vina and PharmScreen were developed more recently than this study was performed. The success of molecular docking can also be dependent on the protein–ligand pair being simulated and CNN_DockBench, a deep-learning based programme, has been developed to predict which molecular docking programme would be most successful for correct pose predictions [49].

4.2. Molecular docking applied to M. tuberculosis

Molecular docking is the most widely used computational approach for virtual screening against M. tuberculosis proteins and has resulted in numerous published studies that are summarised in Table 1. The majority of proteins targeted by this approach represent proteins encoded by essential genes, based on the Himar1 transposon mutagenesis study of DeJesus (2017) [19]; with the exceptions being antigen 85c, BioA, EthR, NarL and LipU. However, non-essentiality assigned by this approach is based on in vitro growth and does not guarantee these genes are non-essential in vivo [50]. For example, the NarL protein is required for anaerobic survival during infection and BioA is essential for biofilm synthesis during M. tuberculosis latency [51,52]. In addition, the EthR protein is involved in ethionamide resistance and hence survival during drug treatment [53].

Focussing on the types of proteins being targeted, the majority are involved in either intermediary metabolism or lipid metabolism within M. tuberculosis. Then regulatory proteins and cell wall regulator proteins make up the remainder of the targets. InhA and DprE1 have been the targets of the most virtual screening campaigns, with at least three each so far. InhA is the eventual target of both isoniazid and ethionamide, following activation of these produgs [54] and hence InhA represents a validated target, whose inhibition has an in vivo impact on M. tuberculosis survival. While DprE1 is targeted by several antimycobacterial drugs in the current anti-TB development pipeline, and hence it represents another validated target [1,10].

The majority of the studies outlined in Table 1 have taken compounds from general chemical databases containing millions of compounds. While the remainder are equally split between TB-
| Docking Programme | TB Protein | Protein function | Compounds Screened | Computational follow-up | Experimental Validation | Reference |
|-------------------|------------|------------------|--------------------|-------------------------|-------------------------|-----------|
| AutoDOCK Vina     | MurB       | Peptidoglycan biosynthesis | FDA-approved compounds: Drug Bank (1932) | MD simulations and MMPBSA energy calculations | No | [68] |
| AutoDOCK 3.05     | Isocitrate lyase | Glyoxylate bypass | Malaysian Local Natural Compound Database – 3,000 compounds | MD simulations – then ensemble docking | In vivo testing against M. smegmatis. | [73] |
| AutoDOCK 4.0      | RmlD       | Carbohydrate biosynthesis | Super Natural-II database – 570 compounds | MD simulations – then ensemble docking | No | [74] |
| CDOCKERIER        | BioA       | Biotin biosynthesis | Enamine REAL database – 4.5 million compounds | ADMET predictions | In vivo confirmation against M. tuberculosis H37Rv | [51] |
| FRIGATE           | Antigen 85c | Lipid metabolism | ZINC database – 2 million compounds | No | NMR binding against Antigen 85c and MIC against M. smegmatis | [76] |
| Glide             | LipU       | Lipid hydrolysis | 6,282 FDA-approved drugs | MD simulations and Prime MMGBSA calculations | No | [77] |
| AutoDOCK          | AroB       | Shikimate pathway | 1,082 compounds preselected from DrugBank database | MD simulations | No | [78] |
| AutoDOCK          | GinA1      | Glutamine biosynthesis | ChEMBL antimycobacterials – 56,400; FDA-approved drugs – 1596; natural products – 419 & phytochemicals – 918 compounds | MD simulations and MMPBSA calculations | No | [79] |
| AutoDOCK          | DprE1      | Arabinogalactan biosynthesis | 30,789 ChEMBL antimycobacterial compounds | ADMET predictions; MD simulations and MMPBSA & MMGBSA calculations | No | [80] |
| GOLD              | EthR       | Transcriptional regulator | Drugs Now subset of ZINC database – 405,201 compounds | Follow-up performed MD simulations & binding energy calculations | In vivo confirmation against M. tuberculosis H37Rv & crystal structures of compound-EthR complexes | [53] |
| LibDock           | KasA       | Mycolic acid biosynthesis | Top 50 diverse compounds selected by machine learning(85) | Toxicity predictions | In vitro binding to purified KasA protein | [86] |
| AutoDOCK, GOLD, GOLD & Plants | InhA | Mycolic acid biosynthesis | 2,050 compounds pre-selected from Enamine database | MD simulations | In vitro inhibition assay against Mtb & in vivo MIC against M. tuberculosis H37Rv | [88] |
| GOLD Score        | MbtA       | Mycobactin synthesis | 4379 diverse compounds, selected from 9 million 67 trisubstituted benzimidazoles analogues | MD simulations | In vitro inhibition assay against AroQ | [89] |
| AutoDOCK 4.2 & Surflex Dock | AroQ | Shikimate pathway | 2,050 compounds pre-selected from Enamine database | MD simulations | No | [90] |
| AutoDOCK 4.2 & Surflex Dock | PtsZ | Cell division | Top 50 diverse compounds selected by machine learning(85) | Toxicity predictions | In vitro inhibition assay against Mtb & in vivo MIC against M. tuberculosis H37Rv | [88] |
specific, natural product and drug repurposing focussed libraries. Hence, completely novel chemical discovery remains the focus of these preliminary drug discovery campaigns. For the majority of these studies (Table 1) the apparent absence of follow-on experimental data (either in vitro or in vivo) confirming compound bioactivity is an obvious limitation that will prevent these predicted compounds from being taken forward.

4.3. Machine learning applied to molecular docking

A wider issue for the use of molecular docking is the propensity for false positives to be generated when screening compound libraries. This may be linked to many programmes’ scoring functions containing inherent biases for large molecules [55]. False negatives can also occur if drug binding requires conformational changes to the protein [56], as molecular docking is focussed on compound flexibility. In contrast, machine learning approaches have been successfully used to predict known protein–ligand affinities with higher accuracy than conventional molecular docking [55,57]. Many comprehensive reviews have been undertaken on the application of machine learning on the scoring functions within molecular docking [58–60], and hence this information won’t be repeated herein.

Several freely available machine-learning programmes exist as open-access resources, these target either the scoring-functions or pose generation of molecular docking programmes [57–59,61]. NNScore2.0 [62], SIEVE-Score [63] and RF-Score-VS [64] are some examples of machine-learning programmes that focus on scoring-functions, all of which are trained on ligand–receptor binding characteristics and associated KD values. In turn, these programmes re-score pre-generated docked poses, thus providing an alternative ranking of all compounds being investigated. These programmes are particularly interesting as they can be applied retroactively to previous docking campaigns, to assess whether promising compounds have been missed. Currently, there is little wide-spread adoption of these new machine learning methods in the field of drug discovery. The lack of uptake is likely due to a lack of immediate accessibility of these programmes, compared to more established molecular docking programmes, and required computational knowledge represents a barrier to entry for programme implementation. These programmes have also been trained on specific sets of drug-protein binding data, SIEVE-Score and RF-Score-VS were trained on the Directory of Useful Decoys, Enhanced dataset [65], while NNScore was trained on 4,141 protein–ligand complexes selected from the protein data bank [66]. Training on specific drug-protein binding data can allow high quality predictions, if the downstream application’s proteins and potential drugs align with the training set [59]. However, if the properties of the training and application proteins/drugs are different, then user-supplied input data to train the models could be utilised to increase the accuracy of predictions [58,59].

Machine learning has been applied to a docking-based approach targeting the ribosomal peptidyl transferase centre (PCT) of M. tuberculosis [67]. This study provided validation for machine learning to be applied for drug discovery, as the generated model predicted binding efficiency to the PCT which matched experimental results [67]. The model was trained using the pose outputs from molecular docking, focussing on the eleven-atom core structure of all the phenylthiazole compounds screened. One limitation of this study was the lack of testing for the model’s ability to predict activity of novel untested structures.

To date, no scoring-function focussed machine learning study, for M. tuberculosis protein molecular docking, has been published in the literature, thereby highlighting a largely unexplored area of research. Providing suitable training sets become available, these machine learning approaches have the potential to greatly improve the efficiency of M. tuberculosis virtual screening studies.

5. Molecular dynamic simulations

5.1. Classical molecular dynamic simulations

Molecular dynamic (MD) simulations are a computational approach to model the motions of atoms over short nano or microsecond timeframes [91–93]. These simulations require an initial input structure such as a crystal structure or homology model, which specifies where all the atoms are and at what velocities they are currently travelling. Then the simulation uses pre-determined force fields, CHARMM [94] and AMBER [95] force fields are commonly used [92,96], to calculate the forces acting on each atom. This information is used to solve the Newtonian equations of motion generating a trajectory and the atoms are moved to these new positions. This movement normally represents a time-step of two femtoseconds. This process is repeated for each new set of atom positions until a pre-set number of timesteps has elapsed [93]. For a short ten nanosecond simulation, five million timesteps need to be calculated. At specific timesteps, for example every 5,000 timesteps, the atom positions and trajectories can be saved as trajectory frames. This process allows a subset of the total number of timesteps, rather than the whole simulation output, to be processed during downstream analysis. Several programmes exist to undertake these MD simulations, including NAMD [92], GROMACS [97], AMBER [98] and CHARMM [99], with the first two being utilised the most. The necessary input files for these programmes can be generated using the CHARMM-GUI webserver [100] or manually using VMD [101], among several other resources [93].

Despite there being several programmes in currently use to perform MD simulations, no one programme is favoured in the published literature [96]. One main distinguishing feature of these MD simulation programmes is whether the license is free or commercial [48]. Generally, MD programmes are found to output similar results, despite them utilising different force fields and algorithms [96,102]. The efficacy of each programme will depend on the system being simulated, the parameters required, the type of MD simulation being undertaken and the hardware available. For example, GROMACS has been developed to allow its execution on any hardware from laptops to supercomputers [97], whereas, NAMD has been developed for optimum scalability across high performance clusters [92].

5.2. Enhanced sampling MD simulation methods

The above description of MD simulations has focussed upon classical MD simulations (cMD), using only the Newtonian equation of motion and force fields to solve them [92]. However, these approaches are limited in the amount of conformational space they can sample in the limited timescale of the simulation. This lack of sampling viable conformations and hence, the inability to simulate larger-scale structural changes, is one of the major challenges in MD simulations [103–105]. As a result, several modification to MD simulations have been introduced to include additional parameters which speed up this sampling process [48,92,103,104,106].

These alterations to the MD simulations can be split into two main groups: collective variable (CV) alterations and tempering [48]. The main difference is the former group of approaches requires existing knowledge of the system being modelled, whereas the latter set of approaches does not. However, these two groups of approaches have also been combined [103,107]. The CV MD approach, most commonly metadynamic MD simula-
tions, is based on simulation variables which can be measured, including: bond distances, dihedral angles, root-mean square-deviation (RMSD) from a known structure, radius of gyration or ligand–protein distances [105]. These variables are then used to determine when additional biasing parameters will be applied to the simulation [48, 103, 105, 108] and two to three CVs are normally specified per simulation [48, 103]. The two most common tempering approaches are accelerated MD [109, 110] and replica-exchange MD [111]. Tempering MD simulations generally enhance the conformational sampling by uniformly boosting all degrees of freedom of the system. This is not an exhaustive list of enhanced sampling methods and several comprehensive review of these methods related to MD simulations have been undertaken elsewhere [103–105].

MD simulation outputs generally are not used alone for analysing drug-protein binding, rather data is extracted from the simulations and then used for further analysis within drug discovery. Two main pieces of information can be gained: 1, the conformational changes of the protein and 2, the energy of the protein and its associated interactions.

5.3. MD simulations applications to M. Tuberculosis

MD simulations have been applied to several M. tuberculosis proteins, as summarised in Table 1. The majority of simulations have been utilised either in ensemble docking or predictions of protein–ligand binding and these will be discussed later. Two studies are exceptions, using MD simulations to model drug-binding interactions in more detail. In the case of trisubstituted benzimidazoles binding to FtsZ, the MD simulations were used to validate the docking-predicted binding interactions and assess the protein–ligand complexes’ stability [90]. The other paper, utilising MD simulations, focussed on simulations of the RND efflux pumps involved in antimycobacterial drug resistance, specifically MmpL5 [112]. This work focussed on the potential interactions of two antimycobacterial drugs with this potential efflux pump and explored the effects of binding on both the MmpL5 trimer and drugs. Both these papers highlight the greater level of information which can obtained on drug-protein interaction through MD simulations compared to molecular docking approaches. To date, there are no reported studies that have applied enhanced sampling MD simulation approaches to M. tuberculosis proteins for drug discovery purposes.

5.4. Machine learning applied to MD simulations

The main drawbacks of MD simulations are the computational resource demands, the large timescales required to sample many protein conformations and errors due to the underlying force fields. To combat these issues, machine learning has been applied to several different aspects of MD simulations, including improvements to the underlying force fields [113–115]; increasing the protein conformations which are sampled [116] and improving the analysis of MD simulations [116, 117]. The application to the improvement of underlying forcefield, while able to achieve large timescale reductions [115], have so far only been applied to simple organic molecules [113, 115, 118] or large single-component systems [119], rather than biomolecules. Selected examples for machine learning-based enhanced sampling of protein conformations will be highlighted, as they have been applied most extensively to biomolecular MD simulations.

Several methods have been applied to enhance the sampling of protein conformations during MD simulations, either through biasing of ongoing MD simulations [120] or through selection of starting-point protein conformations for short MD simulations [121, 122]. Anncolvar uses machine learning to approximate CVs for metadynamic MD simulations, which would otherwise be too computationally expensive to apply during a simulation, such as molecular surface area calculations [120]. Hence, allowing novel biases to be applied, to generate more diverse protein conformations. An alternative approach has been to use an MD trajectory as the machine learning input, and then generate predicted protein conformations, expanding the structures which are sampled without increased simulation times [123].

These machine learning approaches are still in the early stages of development, with many different methodologies being employed concurrently, and as yet, no one approach is favoured [113, 116]. No applications of machine learning onto MD simulations for antimycobacterial drug discovery have been published. Once these approaches become more accessible, they may allow faster screening of compounds and may make MD simulations accessible to more resource-limited settings.

6. Ensemble docking

One of the major disadvantages of molecular docking of compounds against protein targets is the inability to account for the protein flexibility during these simulated drug interactions. Addition of protein flexibility to current docking programmes would impede their use for screening large compound libraries due to the computational demands. One method which has been adopted to help reduce this problem is use of ensemble docking [37, 48, 124]. This approach incorporates the flexibility of the protein by performing molecular docking on an ensemble of protein structures and then a weighted average score for each compound is calculated.

The structural diversity of a protein can be found in two ways. First, if several crystal structures of the protein exist, either the apo- or compound-bound forms, they can all be used as known protein conformations [37, 125]. Alternatively, MD simulations can be performed on a protein structure, either a crystal structure or model, then the trajectory frames can be clustered. The most common or representative protein conformations found by clustering are used for ensemble docking [48, 126]. Several methods are available for the clustering of MD trajectory frames, including: g-cluster within GROMACS, which clusters by a user-specified RMSD cut-off [97]; clustering within Chimera, which clusters by an algorithm-determined RMSD cut-off [127]; POMVE which clusters by binding pocket shape [128] and principal component analysis which groups structures along trajectories representing the largest structural differences. A major issue of these clustering approaches is how to select for the best clustered structures to use for ensemble docking, as the most abundant conformations may not be the drug-binding conformations [106]. Hence, a balanced approach is required between selecting diverse conformations and selecting conformations that can discriminate between active and inactive ligands, to allow ensemble docking to be most effective.

Several different types of MD simulations can be used to generate the structural diversity. Replica exchange, accelerated and metadynamics MD simulations, described above, are some of the methods which have been used to generate increased structural diversity across the trajectory frames [48, 104, 106]. These enhanced sampling methods increase the likelihood that the ensemble of protein structures used for docking represent the physiological structures which occur within the cell and ideally the drug-binding conformations.

Ensemble docking has also been utilised against M. tuberculosis proteins specifically, using cMD trajectories for ensemble generation, for Glf2 [129], RmID [74] and Isocitrate lyase [73]. The trajectory lengths of the MD simulations used were 100, 50 or 18 ns, respectively; with 10 ns MD simulations previously being suffi-
cient to sample druggable binding pockets in multiple targets [106]. Based on these trajectories, 100, 13 and 22 different structures were obtained by clustering for downstream ensemble docking, using cpptraj ([129,130], no further information provided), g_cluster using a 2.0 Å cut-off [74] or active site clustering by backbone RMSD [73], respectively. Validation of these conditions/methods for drug-binding predictions, through in vitro protein-binding analysis, was not undertaken during these studies [73,74,129]. However, in vitro antimycobacterial testing against *M. smegmatis*, undertaken in one study, indicated potential antimycobacterial inhibitors from the crude plant extracts under investigation [73].

An example of the application of machine learning onto ensemble docking is ENRI, a programme that has been developed to enrich protein conformations which can discriminate between active and inactive compounds [131]. The ENRI programme initially focussed on nuclear receptors, which are not known to be present within *M. tuberculosis*; hence, if suitable training sets for *M. tuberculosis* protein families were developed, this could then be used to adapt ENRI’s use. Ideally, this approach will allow fewer protein conformers to be utilised for ensemble docking, representing more accurate drug-binding conformations, leading to more successful hits from these virtual screens. In addition, as ensemble docking utilises both MD simulations and molecular docking, any machine learning programmes which have previously been mentioned, could be combined and applied to further increase ensemble docking success.

7. Protein-Ligand binding energies

Ensemble docking allows the protein flexibility to be accounted for during molecular docking compound screening, improving upon one limitation. However, the other main limitation of molecular docking programmes are the scoring functions, which are used to predict the binding energies of compounds. The scoring functions generally only consider bonding interactions, such as hydrogen bonding, electrostatic and hydrophobic interactions [15]. This excludes the contributions of solvation, both the displacement of solvent from the active site and the stability of the molecule in solvent, and entropic contributions [39,132,133]. Several more advanced approaches exist to take account of these discrepancies, including Linear Interaction Energy (LIE), Molecular Mechanics with Poisson Boltzmann (or generalised Born) and Surface Area solvation (MM-PB(GB)SA) and alchemical methods [132–135]. The first two methods generally utilise the end-point outputs from MD simulations for their calculations, whereas the last set of approaches require new MD simulations. These methods can generate accurate predictions of the ligand–protein binding energies, but the current computational demands make this inaccessible for more than a few compounds.

MM-PB(GB)SA is the most widely adopted approach for estimating ligand binding energy, after molecular docking [132,136,137]. This approach performs calculations on the free ligand, free protein and ligand–protein complex, then the differences are used to estimate the free energy of ligand binding [132,138]. These calculations are performed in three sections, Molecular Mechanics (MM), Poisson Boltzmann (PB) (or generalised Born (GB)) and Surface Area solvation (SA), before the summation is used to estimate the binding energy [137]. Using one MD simulation of the ligand–protein complex, rather than three separate MD simulations for each component, appears to be the favoured approach, giving similar accuracy with lower computational requirements [132,139,140].

The calculation of drug–protein binding energies using MMPB(GB)SA has been applied to several drug discovery attempts against *M. tuberculosis* proteins. These include: LipU [77], GlnA1 [79], DprE1 [80], PknA [81], NarL [52], PanC [141], MurB [68] and MurE [68]. In the majority of cases, Glide was used to perform the initial virtual screen. For the MMPB(GB)SA calculations, either the Prime MMGBSA method was used, or MD simulations were performed, and the trajectory outputs used for the calculations. The former approach performs energy minimisation of the ligand, protein and docked ligand–protein complex in place of an MD simulation, then performs MMGBSA calculations [142], this was used for LipU and MraY drug-binding calculations [77,83]. The latter approach used MD simulations which varied between 15 and 100 ns per ligand–protein complex. Depending on the MD programme used, either GROMACS or AMBER, the associated script to calculate MMPPBSA energies, or MMPB(GB)SA energies for DprE1 & PanC [80,141], was used. Unfortunately, as no in vitro binding studies were performed following these computational studies, a comparison of the relative accuracies of Prime MMGBSA vs MD simulation-MMPBSA is not possible.

The other methods to calculate drug-protein energies have not been applied to *M. tuberculosis* proteins as often. However, LIE has been used for drug binding energies against EthR [84] and an alchemical method to estimate the absolute ligand binding energy has been applied to RmIC [143].

Machine learning has rarely been applied to these current binding energy calculation methods, likely due to large computational costs of performing them. The only application found was the use of machine learning to filter the MD trajectory frames used to calculate MMPPBSA binding energies [144]. This led to an increased correlation between experimental and predicted binding energies for MMPPBSA calculations, but no difference for MMGBSA calculations [144]. More generally, any machine learning approaches which improve the efficiency of MD simulations, will also increase the efficiency of these follow-up binding energies calculations, as they also required MD simulations.

In contrast to the lower number of applications of machine learning onto current methods, many machine learning based programmes have been developed to predict protein–ligand binding affinity [57]. The focus on novel approaches is likely due to the high computational costs of the current methods. However, these approaches have not been widely tested, with only one machine-learning based on approach used in the latest SAMPL host–guest challenge [145].

8. Hit analysis

Computational approaches for drug discovery have the obvious drawback of generating many potential hits, which often include many false positives due to inherent software biases. This is especially true of molecular docking and hence further down-stream analysis of the hits should be undertaken in confirmatory studies. Potential hits can also be mis-classified as false negatives, especially during molecular docking due to the forced protein rigidity [37,38,56]. The true scale of the false negatives problem may be underestimated, as they are rarely evaluated following virtual screening campaigns, despite one study finding two of five inhibitor classes were missed by molecular docking compared to high-throughput screening [56]. However, ensemble docking may reduce this issue by accounting for protein flexibility [56,126,146] and allowing multiple drug-binding modes to be assessed.

When selection of follow-up hits is being undertaken on the basis of physiochemical properties, it should be noted that the current TB-drug space has significantly different physiochemical properties in comparison to those specified by the Lipinski’s rule of five [32,147]. The difference in compound properties is in part due to the hydrophobic *M. tuberculosis* cell envelope, which pre-
vents penetration of many drug compounds. To aid in the evaluation of hits from an in silico drug discovery screen, several computational approaches exist for predicting penetration through the M. tuberculosis membrane \[148,149\] and predicting in vivo M. tuberculosis activity \[150\]. This can allow downstream development of hits which are predicted to both strongly bind a protein of interest and penetrate the M. tuberculosis cell envelope.

An initial study modelled the M. tuberculosis cell envelope, using MD simulations to evaluate compound penetration, however, this could only test a small number of compounds at once \[151\]. More recently machine learning has been used to predict compound permeability and the MycPermCheck programme has been developed to allow for routine screening of compounds \[149\]. Alongside predictions of M. tuberculosis membrane penetration, further machine learning models have been developed to predict compound activity against M. tuberculosis in vivo \[85,150\]. This has been undertaken using either whole cell M. tuberculosis activity data \[85\] or in vivo data from mouse infection models \[150\]. The former machine learning model has been successfully applied to InhA docked hits, to prioritise molecules with potential whole-cell M. tuberculosis activity \[69\]. While the activity data used to train these in vivo activity models is publicly available, the trained models themselves are not, and hence are less accessible for use within hit analysis, compared to MycPermCheck.

The predictive power of machine learning based models is heavily tied to the training dataset used, both the breath of compound properties and the quality of the data \[152\]. Hence, these programmes will effectively predict membrane penetration/compound activity for some novel compounds but will generate inaccurate predictions for many others. In the short-term, experimental validation of any predictions would need to be undertaken rather than complete reliance on these computational analysis methods.

9. Experimental validation

Whilst many of the published studies summarised in this review (Table 1) have provided many novel "potential" hits, and are useful for testing computational drug discovery approaches, the lack of follow-up in vitro or in vivo confirmatory studies means that many of these projects have stalled. To allow the hit compounds of these studies to be turned into antimycobacterial lead compounds, at least some experimental validation is required. As a minimum, either the hits should be tested for binding against the protein of interest or tested for bacteriostatic and/or bactericidal activity against Mycobacteria, ideally M. tuberculosis. Both of these approaches provide evidence that development of these new hits may be successful. The latter validation method is more informative for further development, as it confirms a drug can penetrate the M. tuberculosis cell envelope and is not removed by efflux or deactivated by metabolism before it exerts its bacteriostatic/bactericidal activity. In addition, due to the increasing levels of M. tuberculosis drug resistance, testing of novel drug compounds against clinical MDR-TB strains would provide strong evidence for their future development \[10\], due to limited number of treatment options \[1\].

An exemplary approach for in silico structure-based drug discovery against M. tuberculosis was undertaken by Tatum et al. \[53,84\] (Table 1). An initial virtual screening approach, using GOLD, docked 409,201 compounds, selected from the DrugsNow subset of the ZINC database, against EthR. Then, hits were shortlisted for chemical diversity and by visual inspection of binding poses. This led to 85 compounds which were tested for in vitro binding against EthR, and 20 compounds showed binding. Fifteen of these hits were then tested against M. tuberculosis H37Rv and co-crystallisation trials of EthR-drug were undertaken, leading to four crystal structures. This study provides compelling evidence for the development of the four potential EthR inhibitors \[53\] and for adopting similar approaches for future drug discovery efforts. A follow-up study then focussed on MD simulations and LIE calculations for the four potential inhibitors, providing computational comparisons of the experimental binding modes \[53\]. Further studies similar to this work are required to move from promising drug hits into lead compounds that can be tested within preclinical trials.

10. Summary and outlook

The current antimycobacterial drug discovery approach has focussed on in vitro high-throughput screening and drug repurposing. These are likely to remain important areas of research and the in-silico approaches discussed herein can be complementary methods and aid in the discovery of new compounds. These approaches should allow a larger number of compounds to be virtually screened, before testing in vitro and allow the diversity and ideally quality of hits to increase. This will hopefully increasing the likelihood of this ligand discovery being translated into novel drug discovery \[152\].

Molecular docking represents the most widely applied method for discovery of drug compounds targeting M. tuberculosis proteins, likely due to its wider accessibility. There is a higher availability of user-friendly programmes for this approach, whereas MD simulations and its associated analysis have a greater barrier to entry, hence explaining their lower frequency of utilisation. However, the other in silico methods outlined here, especially MMPBSA calculations and machine learning-based hit analysis, provide other approaches for filtering molecular docking hits before in vitro testing needs to be undertaken. Ideally, they will allow fewer, but higher quality, hit compounds to be tested, reducing the overall costs of early drug screening, and providing a stronger basis for lead development and testing.

Out of all the applications of machine learning described herein, its application to molecular docking is likely to have the greatest impact in the short-term, as this initial step is becoming more universal in the early stages of target-based drug discovery. Hence, as machine-learning applications have been shown to generate more accurate protein binding predictions, this will likely increase the number of true positive hits that are tested in vitro. In the longer-term, the use of machine-learning methods to accurately calculate ligand–protein binding energies may revolutionise how protein specific inhibitors could be found. However, this would be dependent on the calculations being at a computational efficiency accessible for large-scale screening of compounds and accurately to fully model in vitro binding.

Alongside the application of these current and novel methods to M. tuberculosis drug discovery, they need to be tied to experimental validation of any hit compounds. This helps to validate both the methodology and generates more activity data that can be used to train machine learning models. Only through validation of computational hits can novel drug compounds be found to combat the growing drug resistance present within M. tuberculosis worldwide.

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