

# Using Comparative Genomics and Virtual Screening for Antibiotic Drug Discovery

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## Abstract

Methods from comparative genomics, sequence analysis, and virtual screening were combined to predict new drug targets and the chemical compounds that bind most strongly to those targets.

**Motivation:** The evolution and spread of antibiotic resistant pathogenic bacteria has been rapid and often lethal, while the pipeline for new antibiotics has remained virtually bone dry. We face an urgent need for new antibiotics and more cost effective methods to support drug discovery; greater efficiency may be achieved by prioritizing *in vitro* testing.

**Results** We found that this combination of methods provided a cost effective way to predict suitable drug targets, their active sites, and prioritize chemical compounds for the *in vitro* testing of their antimicrobial properties, especially in the light of domain expertise.

**Availability** Software and data are available upon request.

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**Supplementary Information** Supplementary tables and figures are available with the online version.

## 1 Introduction

In this thesis we report on our application of methods from comparative genomics and virtual screening to discover new bacterial drug targets and chemical compounds that bind to those targets. We hypothesized that given the growing abundance of genomic sequence data and computing power, the combination of sequence analysis and virtual screening provide an efficient way to identify probable active sites on essential proteins and the chemical compounds that bind to them with the greatest affinity. Our hope is that this research will contribute the antibiotic drug discovery process and that new antibiotics will be developed and, eventually, be used to promote health and save lives.

The specific objectives of this research include the following:

- Acquire the complete proteomes of *Bacillus subtilis* and 10 to 100 diverse bacterial proteomes.
- Identify the cell wall associated proteins in *Bacillus subtilis* that are the most broadly conserved across bacterial species, both Gram-positive and Gram-negative, and use sequence analysis to determine probable active sites.
- Determine target proteins: broadly conserved proteins have a known 3D structure.
- Virtually screen chemical compounds against probable active sites on target proteins.
- Identify the “best” compounds for *in vitro* testing against pathogenic bacteria.
- Develop software and techniques, especially those that use high throughput distributed computing, that scale to very large data sets.

The treatment of infectious disease with antibiotics has been one of the major successes of the 20th century and has led to large increases in life expectancy. However, microorganisms are becoming increasingly resistant to multiple classes of antibiotics and there is an urgent need for new compounds [43]. For example, *Staphylococcus aureus* (*S. aureus*) is the leading cause of bacterial infections of the skin, soft tissue, bloodstream, and lower respiratory tract in developed countries [11]. Although penicillin was effective initially, resistant strains emerged in the mid-1940s, methicillin resistant *S. aureus* (MRSA) was discovered in the 1960s [11], and strains with some degree of vancomycin resistance have recently emerged worldwide [24]. The implication is clear. If one of the greatest successes of the 20th century is lost, we face the possibility of a commensurate decrease in life expectancy.

In an effort to meet this challenge, the first major part of our research applies sequence analysis

to recently determined genomic data. The availability of bacterial genome sequences was expected have a large impact on antibiotic discovery by enabling new genomics-based approaches for identifying new molecular targets [32] but, it should be emphasized, genomics does not significantly alter the time frame for the drug discovery process, which can take up to 12 years [41]. In practice, genomics has delivered novel essential drug targets for target based discovery efforts [41; 42].

The second major part of our research uses the power of distributed computing for virtual screening of compounds. This application of virtual screening was motivated by several considerations. First, computational methods have advanced to become a crucial component of many drug discovery programs, and virtual screening techniques are now widely used [31]. Second, the size of compound libraries has been growing exponentially from the output of robotic “combinatorial chemistry” [23]. Third, the number of known 3D structures continues to grow, with thousands being added each year [70]. Fourth, small organic ligands are integral parts of many protein structures [37, p. 2] and, more importantly, they usually participate directly in the function of a protein and, consequently, may be one of the most interesting components of a structure to study [37, p. 103]. And finally, we noted that virtual screening to discover new inhibitors is becoming a common practice in modern drug discovery [61].

Although pathogenic bacteria have proven to be remarkably adaptable, the cell wall remains a good target for antibiotics [62; 63; 60] and more than 100 genes involved in bacterial cell wall biosynthesis and division have been identified. Starting with these genes, we used a comparative genomics approach [3] to identify a set of homologous genes/proteins in multiple bacterial species.

We used BLAST [1] for pairwise sequence alignment and MUSCLE [13] for multiple sequence alignment (MSA), Jalview [71] for visualization of the MSA and calculation of conservation scores for each amino acid in the *B. subtilis* query protein, and PyMOL [10] for 3D molecular visualization.

We downloaded ligand files representing  $\approx$  100,000 compounds from the ZINC database [26]; almost 90,000 were from the Natural Products Database (NPD) meta data set. Given a promising set of candidate proteins, probable active sites, and ligands we used AutoDockTools (ADT) [59] to prepare receptor and ligand input files and AutoDock Vina [69] molecular docking software

for virtual screening to identify small molecule inhibitors [43].

We found that amino acids with high conservation scores corresponded well with known active sites, i.e., organic ligand binding sites according to the Protein Data Bank (PDB). We also found that by the application of molecular evolutionary principles, sequence analysis and virtual screening, we were able to prioritize bacterial species, their proteins and matching compounds for wet lab experiments. In summary, we considered 123 *B. subtilis* proteins and  $\approx$  100,000 chemical compounds. Four proteins, FtsZ, CoaD, YwtF, and RacE were identified as candidates for virtual screening; FtsZ and YwtF and a small number of specific compounds were selected for testing which, based on empirical evidence, had the highest probability of exhibiting antimicrobial properties.

This thesis is organized as follows. Section 2 provides background information on bacteria, antibiotics, and the motivations underlying our research. Section 3 provides a conceptual introduction to the methods used in our research and why they were chosen. Section 4 describes how our methods were applied. Section 5 presents our results and interpretations. Section 6 provides a brief discussion of our findings. In addition, the Supplementary Material in the Appendix provides many detailed tables and figures.

## 2 Bacteria and Antibiotics

### 2.1 Background

The ‘golden era’ of antibiotic research occurred from the late 1940s to the late 1960s [41]. In fact, nearly all classes of antibiotics (acting on the same target) currently in use were developed prior to 1970 [32]. Since the 1980s, there has been a decline in the number of new antibiotics brought into clinical practice, and even these new antibiotics bind to the same target molecules as their predecessors. And now, as pan-resistant strains of pathogenic bacteria have become a clinical reality, the pipeline of new drugs is “virtually bone dry” [55].

This has been due to a number of factors including the high expense of developing new drugs, the relatively low profit potential of antibiotics, industry concentration through mergers and acquisition, and higher regulatory hurdles [54; 55; 51]. Consequently, new classes of antibiotics are desperately needed to combat the global emergence of bacterial pathogens – this has been described

as one of the most important challenges facing the pharmaceutical industry [7].

Rational drug design has been based on detailed structural information of the drug target. However, the success of this paradigm, which emerged in the late 1960s, has been limited by the small number of well defined molecular targets and progress has been limited to a few chemical classes, e.g.,  $\beta$ -lactamase inhibitors, carbapenems and fluoroquinolones [32]. This limitation and the lack of new antibiotic classes since 1970 is shown in Figure 1 where the graph flattens out in the late 1960s.

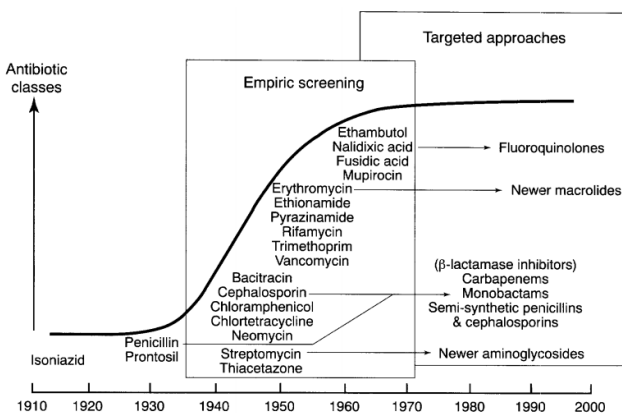


Figure 1: Discovery of new classes of antibiotics: Empiric screening has generated antibiotic classes, while targeted approaches (i.e., rational drug design) have generated antibiotic agents that act by related mechanisms. Some representative antibiotics are shown here. Image: [32].

## 2.2 Antibiotic Resistance

Pathogens such as *Staphylococcus aureus* (*S. aureus*), *Streptococcus pneumoniae* (*S. pneumoniae*) and *Enterococcus faecalis* (*E. faecalis*) have become increasingly resistant to multiple antibiotics [41]. Methicillin-resistant *S. aureus* (MRSA) and vancomycin-resistant *Enterococci* (VRE) infections tend to occur in hospitals whereas multi-drug-resistant *pneumococci* are more likely to occur in the community [2; 52; 28]. If current trends continue, even vancomycin, a last line of defense for hospitalized patients, will no longer be effective [41].

For example, *S. aureus USA300* is resistant to methicillin, ampicillin, penicillin, mupirocin, macrolides, lincosamides, tetracycline, ciprofloxacin and streptogramin. Antibiotic resistance genes are present on strain USA300's chro-

mosome and plasmid, they include *mecA* on the chromosome, *ermC* and *ileS* on both the chromosome and plasmid, and gene *blaZ* on the plasmid [12]. *S. aureus USA300* sensitive to vancomycin [12]. Vancomycin resistant *S. aureus* (VRSA) is a rare strain of *S. aureus* that was identified in 2002 [72]; it has become resistant to the glycopeptide antibiotic vancomycin [11].

The main mechanisms by which microorganisms exhibit resistance include drug inactivation or modification through the production of  $\beta$ -lactamases, alteration of a target site, alteration of a metabolic pathway, and reduced drug accumulation by decreasing permeability and/or increasing active efflux of the drugs across the cell surface [19]. However, the acquisition of drug resistance is not necessarily the product of a slow evolutionary process.

Nonresistant bacteria may rapidly acquire resistance from resistant bacteria by horizontal gene transfer using one of three mechanisms: transformation, transduction and conjugation [49]. “Unlike eukaryotes, which evolve principally through the modification of existing genetic information, bacteria have obtained a significant proportion of their genetic diversity through the acquisition of sequences from distantly related organisms. Horizontal gene transfer produces extremely dynamic genomes in which substantial amounts of DNA are introduced into and deleted from the chromosome” [49]. The significance of horizontal gene transfer for bacterial evolution was recognized when multidrug resistance patterns emerged on a worldwide scale [9].

## 2.3 Broad Spectrum Antimicrobial Agents

The range of different microbes against which an antimicrobial agent acts is called its *spectrum of activity*, and agents that are effective against bacteria from a wide range of taxonomic groups, including Gram-positive and Gram-negative, are called *broad spectrum*. Broad spectrum drugs are appropriate where a patient is seriously ill with an infection but the organism is not known [6], i.e., such a drug will have a higher probability of success. Consequently, in this thesis, broadly conserved cell wall related proteins, especially those conserved in both Gram positive and Gram negative bacteria, are analyzed for their drug target potential.

## 2.4 The Cell Wall and Membrane as Targets

The cell wall is an essential structure for virtually all bacteria as it protects the cell from damage and osmotic lysis; it is the target of our best antibiotics [36]. In general, antimicrobial drugs act on important structures or functions that exist in bacteria (prokaryotic cells) but not in humans (eukaryotic cells). For example, Gram-positive bacteria have high osmotic pressure and would burst without a sturdy cell wall. Penicillin and cephalosporin contain a  $\beta$  lactam ring which attaches to enzymes that cross-link peptidoglycans. Interference with the cross-linking of tetrapeptides prevents cell wall synthesis [6].

Although all cell membranes are similar, those of bacteria are significantly different from animal cells. For example, some polypeptide antibiotics such as polymyxins act as detergents and distort bacterial cell membranes – they are especially effective against Gram-negative bacteria which have an outer membrane rich in phospholipids [6].

## 3 Comparative Genomics and Virtual Screening

In this section we provide general background information on comparative genomics and virtual screening.

The application of comparative genomics for target identification in this paper is similar to the application in [3] where conserved proteins were identified, determined to be essential and, therefore, potential drug targets. We chose comparative genomics to identify antibacterial targets because, while not a flood [54], an unprecedented number of novel targets have been discovered via this approach [50].

Given broadly conserved proteins and their sequences, amino acids (and their locations) in those sequences that are important functionally and for binding tend to be conserved across species during evolution [75, p586]. Given a 3D structure, an *active site* is a localized combination of amino acids (which may be far apart in the amino acid sequence) that can interact with a chemically specific substrate and provide the protein with biological activity; as a result, proteins with very different amino acid sequences may fold into structures that produce the same active site [46, p415]. Consequently, *in silico* screening of chemical compound libraries forms the second major part of our

approach. Once an active site or probable binding site has been identified, the binding of small molecule ligands can be modeled. Molecular modeling is a high throughput way of rapidly investigating the binding potential of large numbers of small molecule drugs [75, p587].

While the design, analysis and enhancement of ligands are also important steps in the drug discovery process [38], they are outside the scope of this thesis.

### 3.1 Comparative Genomics

The first insights into genomic drug discovery approaches were made possible by the comparative analysis of the complete genome sequences of 10 bacterial pathogens [17]. Broadly conserved genes conserved tend to be essential, which makes them attractive targets for new broad-spectrum antibiotics [17]. In [3], researchers used *Escherichia coli* (*E. coli*) as a reference organism. They identified four novel genes (*yggD*, *ycfB*, *yihA*, *yjeQ*, and their respective orthologs). These genes were attractive targets, i.e., they were novel, essential, broadly conserved (homologs in a wide range of bacteria including *H. influenza*, *S. pneumoniae*, *H. pylori*, and *B. burgdorferi*), and low toxicity for higher organisms.

In other research, it was reported that many cell division proteins are conserved across Gram-positive and Gram-negative bacteria [40]. Since essential bacterial genes tend to be more conserved than nonessential genes over short and long evolutionary time scales [29], the most highly conserved genes associated with cell wall biosynthesis should be the best targets for new broad spectrum antibiotics.

*B. subtilis* has approximately 4,100 genes; 192 were shown to be indispensable, 79 were predicted to be essential – with about 44 of these involved in the synthesis of cell envelope and the determination of cell shape and division [33]. As of 2000, about 150 essential genes were identified in *S. aureus* [27].

### 3.2 Sequence Analysis

#### 3.2.1 Foundation

The fundamental basis for our approach is the well established notion that similar sequences tend to have similar structures and, consequently, similar functions. Burkhard Rost analyzed more than a million alignments of pairs of protein sequences to determine a threshold for sequence identity [57],

i.e., the minimum level of sequence identity required for an alignment to provide a reliable measure of homology. He found that 90% of sequence pairs with at least 30% identity over their whole length were structurally similar. Below 25% sequence identity, only 10% of the aligned pairs were structurally similar.

As a result, 30% sequence identity is often used to justify an initial presumption of homology [75]. The region between 30% and 20% has been called the *twilight zone* – where homology may exist but cannot be assumed without additional evidence – and below 20% lies the *midnight zone* [75].

### 3.2.2 Pairwise Sequence Alignment

The Basic Local Alignment Search Tool (BLAST) [1] may be applied as a first step in sequence analysis. BLAST uses heuristics to find high scoring pairs and the computationally expensive Smith-Waterman algorithm [64] to generate the final alignment [1]. In general, each query is “blasted” against a database of proteomes to find similar proteins in other organisms. This generates one set of similar sequences for each query protein.

### 3.2.3 Complexity

Alignment algorithms such as Needleman-Wunsch [47] and Smith-Waterman [64] are based on dynamic programming: pairwise sequence alignments have  $O(n^2)$  time and space complexity; multiple sequence alignments of  $k$  sequences have  $O(n^k)$  time and space complexity and are not feasible for alignments of more than three sequences [34]. Hence, all available methods are approximations, i.e., hierarchical clustering of pairwise alignments roughly approximates the phylogenetic tree and guides the multiple alignment [34]. This hierarchical approach to progressive sequence alignment, an idea presented in [16], reduces the  $O(n^k)$  multiple alignment problem to a series of  $O(n^2)$  problems [34].

### 3.2.4 Multiple Sequence Alignment (MSA)

MSA may be viewed as a extension of pairwise sequence alignment that may reveal small active sites in otherwise dissimilar sequences. CLUSTAL is a multiple sequence alignment algorithm introduced in [68]. It uses heuristics, as do all available MSA methods [34], to generate an alignment that is not guaranteed to be optimal for a given scoring scheme. CLUSTALW [22] is an improved version

CLUSTAL [21] – the W stands for weighted. Although CLUSTALW is still the most widely used multiple sequence alignment tool, no significant improvements have been made to the algorithm since its introduction and several more recently developed algorithms perform better in terms of accuracy, speed or both [14].

In a recent review, MUSCLE [13] and T-COFFEE [48] were listed as the most accurate programs [14]. On nearly all benchmarks, these newer programs outperformed CLUSTALW in terms of average accuracy. Given a typical desktop PC, the authors recommended MUSCLE for aligning over 100 sequences that are approximately globally alignable [14]. Since we expected to use a typical desktop PC for MSA of large numbers of sequences that are at least 30% identical, we chose MUSCLE generating for generating all multiple sequence alignment and Jalview [71] for MSA visualization – an example is shown in Figure 2.

## 3.3 Virtual Screening

Virtual screening is an *in silico* structure-based design strategy where small molecules are “docked” into the structures of target molecules and scored for their complementarity to binding sites. This process models similar events that occur in nature. For example, Figures 3a and 3b show two proteins, FtsZ and Coad. An organic ligand (in green) bound to an active site on the surface of each protein; the figures also show polar contacts as defined in the Protein Data Bank (pdb) file for each protein.

Fortunately, accurate energy calculations and scores are not necessarily required for meaningful compound selection and, in a typical docking study, a large compound database may be reduced to a short list of  $\sim 100$  preferred candidates [31]. Docking and scoring are important steps in the filter between a total potential library and testing at the bench [38] and, by some measures, its major achievement has been to eliminate inactive compounds from further consideration [5]. Indeed, such computational approaches have been widely used for hit identification and lead optimization [31].

A central problem in drug discovery is the identification of a compound that will bind tightly and specifically to a target protein. Tight binding promotes efficacy at low concentrations, and specificity helps to minimize side effects [38]. Although it is difficult to estimate absolute affinities, comparative docking can indicate relative affini-

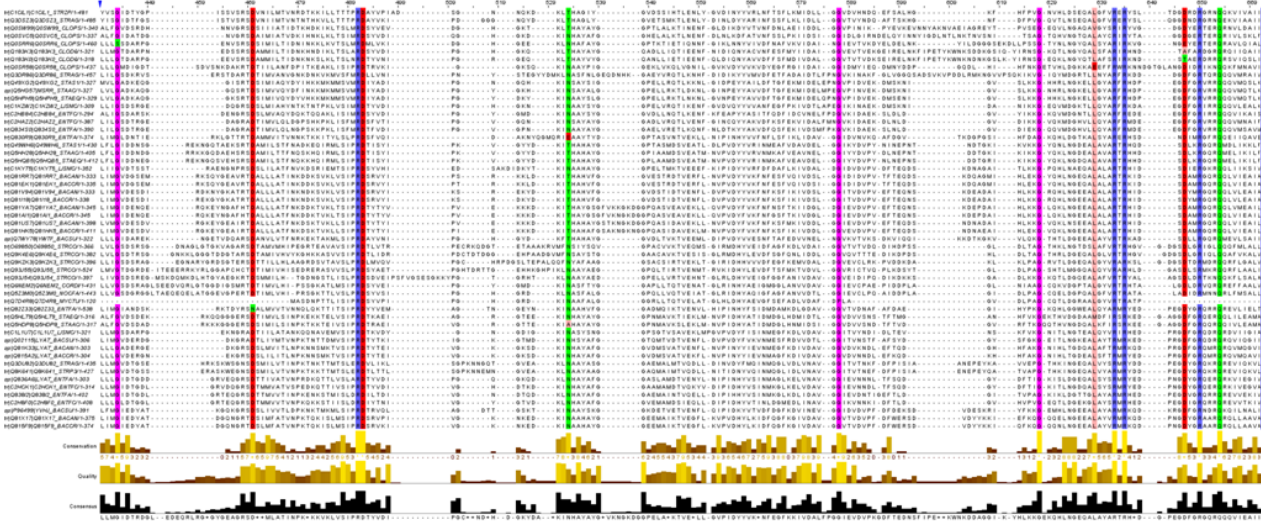


Figure 2: Multiple sequence alignment for sequences similar to *B. subtilis* (columns with 100% conservation are colored).

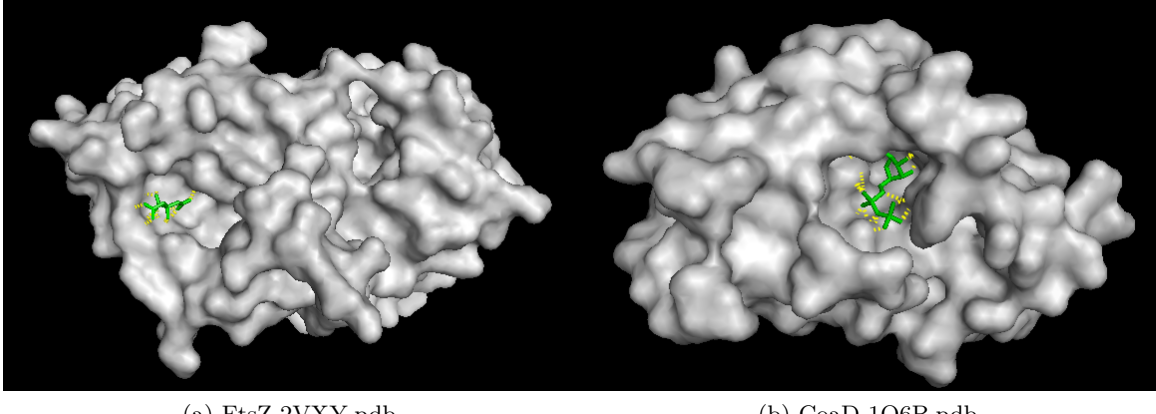


Figure 3: Pymol images of FtsZ and CoaD with organic ligand using Protein Data Bank files.

ties. This means that given a scoring function that can rank ligands in approximate order of affinity, we can select compounds and set priorities for experimental testing [38].

However, the probability of finding a novel antibacterial compound with broad spectrum activity depends on the diversity of biologically relevant compounds available for screening. Chemical diversity is an extremely important factor when searching for new drugs [51], and ZINC provides a large database of millions of commercially available compounds for virtual screening [26].

Docking *in silico* of large sets of ligands is computationally expensive and, consequently, high-throughput computing clusters and algorithms are widely used for virtual screening [53]. We found that virtual screening is well suited to distributed computing. Screening one compound does not de-

pend on the output from screening other compounds. As a result, many compounds can be screened simultaneously on many CPUs located anywhere on a network.

## 4 Methods

In this section we describe the application of our methods and algorithm parameters.

A list of 123 *B. subtilis* proteins was provided by Dr. Nick Allenby of Demuris Ltd., Newcastle Upon Tyne, United Kingdom. We downloaded these *B. subtilis* proteins from the UniProt website. Complete protein lists may be found in the Supplementary Material, Tables 14 and 15. We used the Database of Essential Genes [74; 73] to confirm whether or not broadly conserved genes are essential.

Next we downloaded 60 complete bacterial proteomes in fasta format from the Integr8 sequence database [30]. The 60 proteomes were selected for pathogenicity and diversity. They included 25 Gram positive, 25 Gram negative, and 10 proteomes, such as Mycoplasmas, that do not have a cell wall [18]. Complete proteome lists may be found in the Supplementary Material, Table 16.

As our first computational task, we compared each of the 123 *B. subtilis* proteins with every protein in each of the 60 proteomes in order to identify potential drug targets, i.e., we used BLAST [1] for pairwise sequence alignment to identify broadly conserved cell wall proteins. Then we used MUSCLE [13] for multiple sequence alignment to identify highly conserved residues in each query sequence and mapped conservation scores for those residues to the query protein’s 3D surface to indicate probable active sites. The workflow for the these tasks is shown in Figure 4.

We also used BLAST to identify proteins conserved in the 25 Gram positive and the 25 Gram negative proteomes; and to gauge the redundancy of broadly conserved proteins in the *B. subtilis* proteome, i.e., we assumed that relatively unique proteins are more likely to be essential.

After identifying potential targets, we docked NPD ligands from the online ZINC database with each potential target protein *in silico*, i.e., we used Vina [69] to perform virtual screening to identify chemical compounds for *in vitro* testing. The search space that included the most promising, highly conserved region of the receptor molecule – this region either included a known active site or a predicted active site. And finally, top scoring ligands were selected for *in vitro* testing. The workflow for these tasks is shown in Figure 5.

#### 4.1 Pairwise Sequence Alignment

We used the Basic Local Alignment Search Tool (BLAST) [1] to identify the set of proteins that have homologs in many species of pathogenic bacteria. BLAST uses heuristics in an attempt to align pairs of sequences such that the resulting alignment’s similarity score  $S$  is maximized and, consequently, its  $E$ -value, which is a measure of the reliability of the  $S$  score, is minimized [35].

$$E = kmne^{-\lambda S} \quad (1)$$

Equation 1 states that  $E$ , the number of alignments expected by chance, is a function of several parameters: (1) the size of the search space  $m * n$ ;

(2) the normalized score  $\lambda S$ , and (3) a minor constant  $k$ . In a database search, the size of the search space is the product of the number of letters in the query  $m$  and the number of letters in the database  $n$ . As equation 1 indicates, a given increase in  $S$  results in an exponential decrease in the expected number of alignments  $E$  with a similarity score at least as good as  $S$ ; therefore, the smaller the  $E$  value, the more significant the alignment [35].

From Equation 1: A lower  $E$  value reduces false positives at the expense of an increase in false negatives. In fact, very short sequence alignments (i.e., small  $m$ ) with very high  $S$  scores will be filtered out because, all else being equal, a shorter sequence will result in a higher e-value and, therefore, deemed less statistically significant. Consequently, alignments with very short *B. subtilis* query proteins were filtered out and not analyzed.

The Expect value ( $E$ ) is a parameter that describes the number of hits expected by chance when searching a database of a particular size. The lower the  $E$  value, or the closer it is to zero, the more "significant" the match is. However, virtually identical short alignments have relatively high  $E$  values because the calculation of the  $E$  value takes into account the length of the query sequence. These  $E$  values are high because shorter sequences have a higher probability of occurring in the database by chance alone [<http://blast.ncbi.nlm.nih.gov> ].

The pseudo code in Table 1 represents our use of BLAST [1] to identify one or more homologous proteins for each *B. subtilis* query protein. **Input:** a query fasta file of 123 protein sequences, 60 subject proteomes, and a set of thresholds T: a BLAST E value of 0.00001 (a smaller E value will filter out more alignments than a larger one) and a minimum percent identity of 30%.

```
let file = the B_subtilis fasta file

foreach protein P in file
    blast P against the proteomes
        for each hit > T1
            record hit

foreach protein P in file
    if P had >= 1 hit in >= T2 proteomes
        save set of hit sequences to file
```

Table 1: Pseudo code for using BLAST to identify homologous proteins for each *B. subtilis* query protein.

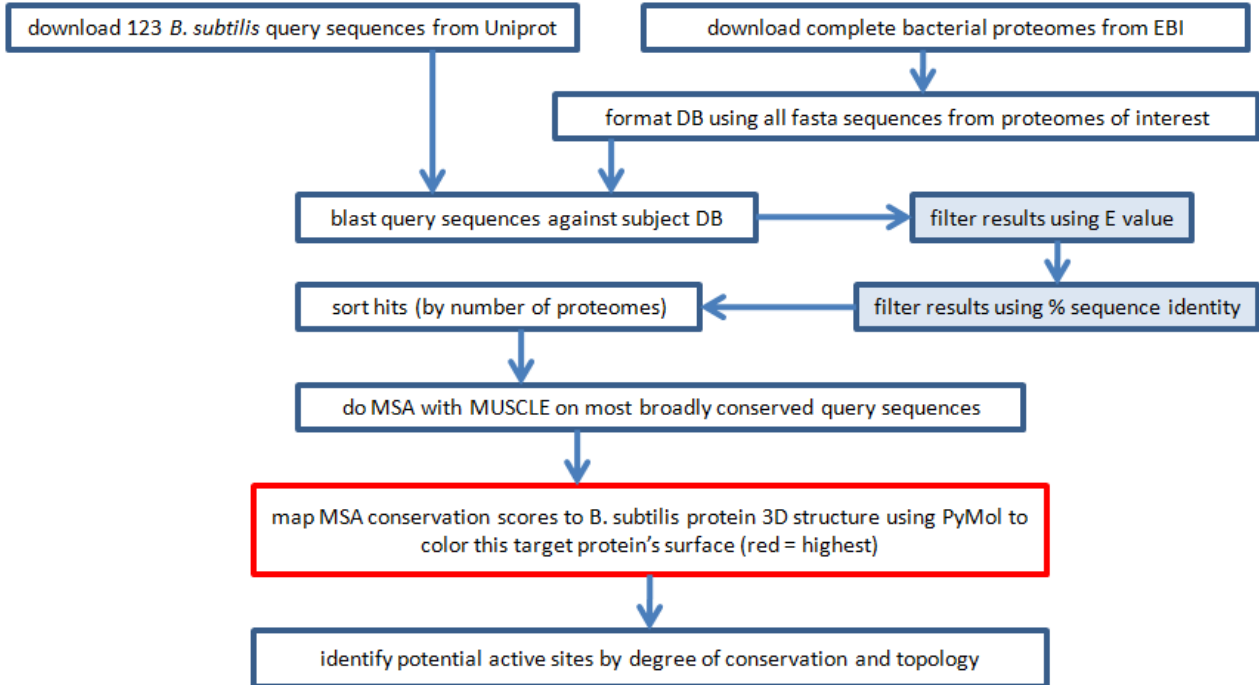


Figure 4: Workflow 1: pairwise and multiple sequence alignment; mapping conservation scores by color to surface of *B. subtilis* 3D structure.

**Output:** a list of *B. subtilis* proteins that are probably homologous to proteins in subject species ranked in order of the number of subject species with one or more probable homologs (hits).

Time complexity: blasting 123 *B. subtilis* proteins against 60 proteomes is relatively fast:  $\approx 162$  seconds, or 2.7 seconds per bacterial proteome. This step of our research was greatly facilitated by software that we implemented in the Java programming language, shown in Figure 14 of the Supplementary Material.

Although outside the scope of this thesis, we note that additional work may be needed for aligning subsequences that constitute active sites, i.e., multiple protein sequences may lack global similarity but possess critical local similarity.

## 4.2 Multiple Sequence Alignment

After we determined which sequences were suitable, based on E value, sequence identity, and whether or not the top scoring *B. subtilis* proteins had a known 3D structure, we generated multiple sequence alignments using MUSCLE [13]. Each MSA included the *B. subtilis* sequence and, typically, about 50 very similar sequences from at least 40 of the 60 proteomes in our local database.

The MSAs produced by MUSCLE were visualized with Jalview [71] and conservation scores were extracted with a Perl script, mapped to each amino acid in the *B. subtilis* protein of interest,

saved to a pml file, and visualized with PyMOL [10].

## 4.3 Virtual Screening

The virtual screening method applied in this research is a receptor based screen [31] where we attempted to “dock” members of a chemical library against a given protein structure *in silico* and software predicted the conformation and binding affinity of NPD ligands.

### 4.3.1 Chemical Compounds used as Ligands

Natural Products Database (NPD) of commercially available natural products and natural product derivatives was downloaded from the ZINC database [26]. The 2008/5 version of the NPD contains 89,425 entries for commercially available compounds from seven vendors that advertise their compounds as being of natural origin, either pure natural products, or chemical derivatives of natural products. We also downloaded the “usual” data sets of 11 random micro vendors, approx 11,000 from the ZINC database. The vendors are listed in Tables 12 and 13 of the Supplementary Material.

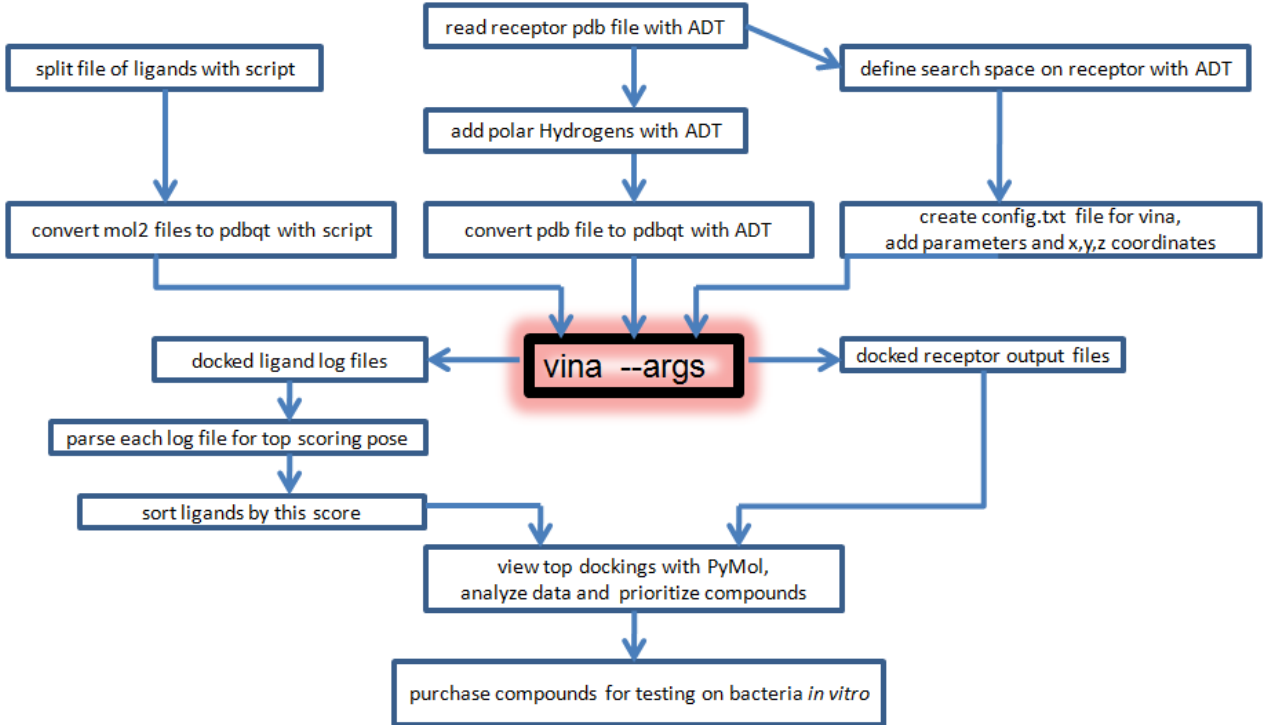


Figure 5: Workflow 2: virtual screening and ligand selection.

#### 4.3.2 Docking Software

For virtual screening, we considered AutoDock 4 [44; 25] and AutoDock Vina (Vina) [69]. Both have been found to be equally capable in ranking smaller molecules with few rotatable bonds, and both exhibit a size-related bias in scoring [8]. However, AutoDock Vina has an improved local search algorithm and the ability to detect and utilize multiple CPUs. As a result, Vina executes more quickly than AutoDock 4 and ranks larger molecules more accurately, “researchers should look to it first when undertaking a virtual screen” [8].

Hence, given a set of target proteins and the Natural Products Database (NPD) ligands from the ZINC database [26], we used Vina [69] to screen these ligands versus our target proteins, i.e., we used Vina to predict how well our drug candidates will bind *in vitro* to probable active sites (identified above by sequence analysis) of known 3D structure. Note: to achieve high throughput, depending on availability, up to 88 receptor-ligand docking operations were performed in parallel on a cluster of computers with a total of 88 CPUs (cisbclust at Newcastle University). This reduced expected time complexity from months to days, i.e., from  $\approx 30$  to 60 seconds to less than 1 second per docking.

#### 4.3.3 Ligand and Receptor Preparation for Vina

The “usual” set NPD ligands were downloaded from ZINC in the mol2 file format (p0.0, p0.1, p0.2, p0.3, p0.4, p1.0). Each file, which contained thousands of ligands, was split into single ligand files in mol2 format. Each single ligand file was prepared for docking and converted to pdbqt format using Perl and the prepare\_ligand4.py script from AutoDockTools [59]. Each protein receptor was prepared and converted to pdbqt format using the AutoDockTools graphical user interface [59]: polar hydrogens were added before conversion to pdbqt format.

#### 4.3.4 Search Space for Vina

AutoDockTools [59] was also used calculate the  $x, y, z$  coordinates of the search space, Table 2, to be explored by Vina [69]; spacing was set to 1.000 Angstrom. The probable active sites identified from our sequence analysis helped to limit this search space and reduce computation time. We noted that the quality of the starting coordinates for each receptor, in effect, put a limit on the accuracy of our docked results [45].

receptor	center_x	center_y	center_z	size_x	size_y	size_z
FtsZ	-17.047	31.454	24.275	22	22	22
CoaD	126.100	69.632	117.280	20	20	20
YwtF	24.077	58.595	85.242	22	22	22
CoaD	-.0972	12.775	18.182	34	24	28

Table 2: Vina Search Space Parameters (Angstroms).

#### 4.3.5 Vina Parameters

We used Vina’s default parameters except for exhaustiveness, which we set to 6 instead of 8. This allowed us to uniformly screen the ligands against more receptor proteins in the time available and, time permitting, subsequently screen a small number of high scoring ligands with exhaustiveness set to at a higher level to make a final selection for *in vitro* testing.

## 5 Results

Our BLAST results indicated that four *B. subtilis* proteins with known 3D structure were highly conserved; FtsZ and CoaD in both Gram positive and Gram negative bacteria, Table 3; and YwtF and RacE in Gram positive bacteria only, Table 4. Complete tables, including information indicating the existence of a known 3D structure for matching proteins on other proteomes, may be found in the Supplementary Material, Tables 17, 18 and 19. We confirmed that two are listed as essential genes for multiple species of bacteria, at the Database of Essential Genes (DEG) web site [74; 73]. This database listed 12 species for FtsZ, 6 species for CoaD, 0 species for YwtF (function is unknown), and only *B. subtilis* was listed for RacE. However, our results suggest that all four may serve as viable drug targets.

Our multiple sequence alignment results showed that highly conserved regions of each protein’s 3D structure was consistent with known or plausible active sites. We also found that virtual screening by Vina predicted binding sites for the NPD ligands that were consistent with sites characterized by high conservation, pocket topologies, and experimentally verified organic ligand binding sites as recorded in the FtsZ (2VXY) and CoaD (1O6B) pdb files from the Protein Data Bank; and the positions of high scoring ligands docked with YwtF (3MEJ) and RacE (1ZUW) pdb files were consistent with regions found by our sequence analysis to be highly conserved.

#### 5.1 BLAST

The following four *B. subtilis* proteins, FtsZ, CoaD, YwtF, and RacE, have a known 3D structure and were found to be broadly conserved in our BLAST results, Tables 3 and Table 4. While this alone suggests that they might serve as suitable drug targets, FtsZ and CoaD are unique in the *B. subtilis* proteome – as Table 6 shows, FtsZ and CoaD had no matches with other proteins in their own proteome and YwtF and RacE had only one match other than with themselves. The complete table may be found in the Supplementary Material, Table 20.

##### 5.1.1 Four proteins: FtsZ, CoaD, YwtF, RacE

In this section we provide a concise summary of our four proteins of interest, but provide somewhat more detail for YwtF and RacE because their drug target potential may be less obvious. **FtsZ** may be the most critical component of bacterial cell division machinery [15]. It is an essential bacterial protein [33], guanosine triphosphatase, and a homolog of mammalian beta-tubulin that polymerizes and assembles into a ring to initiate cell division [20]. **CoaD** was found to be essential for several species of bacteria, including *Streptococcus pneumoniae* [67; 66; 65]. The structure of CoaD was reported in [4]. **YwtF:** Although the 3D structure for this protein has been determined its function is unknown [<http://dx.doi.org/10.2210/pdb3mej/pdb>]. Specifically, this entry describes a domain of unknown function that is found in the predicted extracellular domain of a number of putative membrane-bound proteins, which includes protein psr: described as a penicillin binding protein 5 (PDP-5) synthesis repressor. Another putative membrane-bound protein is Bacillus subtilis LytR. And a third protein, CpsA, is described as a putative regulatory protein involved in exocellular polysaccharide biosynthesis [ InterPro entry IPR004474 ]. **RacE:** D-glutamate is an essential building block of the peptidoglycan layer in bacterial cell walls and can be synthesized

from L-glutamate by glutamate racemase (RacE). The structure of a complex of *B. subtilis* RacE [33] with D-glutamate reveals that the glutamate is buried in a deep pocket. This structure provides new insights into the RacE mechanism and an explanation for the potency of a family of RacE inhibitors [58].

## 5.2 MSA and active sites

Based on our BLAST results for *B. subtilis* versus 60 proteomes, we used MUSCLE to generate MSAs using multiple sequences: 54 for FtsZ, 50 for CoaD, 57 for YwtF, and 38 for RacE. The four proteins vary in length: FtsZ 382, CoaD 161, YwtF 322, and RacE 272 amino acids; and various numbers of gaps were inserted into their MSAs. Figure 6 shows an overview of the MSA for FtsZ, CoaD, YwtF and RacE respectively. These visualizations were generated by Jalview such that colored bars appear only at positions where residues are 100% conserved.

The MSA for FtsZ in Figure 6a shows the greatest degree of conservation, and this is even more apparent in Figure 7a where the conservation scores calculated by Jalview, ranging from 0 (lowest) to 11 (highest), have been mapped to the surface of FtsZ as colors: gray {0..5}, blue {6,7}, yellow {8}, orange {9}, and red {10,11}. Perhaps the most striking feature of this FtsZ 3D image is the large highly conserved area, colored red, and a deep pocket on the left side of Figure 7a. In contrast, the opposite side, Figure 7b, shows less conservation and lacks a similar pocket.

The MSA for CoaD in Figure 6b shows less conservation than FtsZ and its highly conserved residues are grouped closer together. Figure 7c shows a more compact highly conserved region (mostly red) and a substantial pocket; in stark contrast, the opposite side, shown in Figure 7d, shows very little conservation or concave areas.

The MSA for YwtF in Figure 6c shows much less conservation than FtsZ. However, as Figure 8a shows, this protein's highly conserved residues map to one highly conserved surface region (mostly red) which lacks a substantial pocket but nonetheless presents a region that probably plays an important role. In contrast, the opposite side, shown in Figure 8b, shows very little conservation.

The MSA for RacE in Figure 6d appears to show more conservation than the MSA for YwtF, but this greater conservation is not evident from the surface colors for RacE in Figures 8c and 8d. The

implication is that many of RacE's highly conserved residues are located in the interior. This was confirmed by rendering RacE as lines in PyMOL, shown in Figure 16 of the Supplementary Material. Consequently, unlike the other three proteins, RacE does not present an obvious target for docking ligands – where highly conserved surface regions, especially those characterized by a substantial pocket, are probably good drug targets – but its highly conserved residues are internal, the topology of RacE is complex and, given the potency of a family of RacE inhibitors developed as novel antibiotics [58], merits further investigation.

## 5.3 Virtual Screening

A concise illustration of our results from screening  $\approx$  90,000 ligands against the four proteins identified from our BLAST results is shown in Figure 9. Scores were relatively consistent between proteins, i.e., NPD ligands docked with FtsZ tended to score the best (lowest binding energy as measure by kcal/mol), followed by CoaD, YwtF, and RacE; Table 7. In addition, NPD ligands scored consistently higher than the micro vendor ligands, Table 23 of the Supplementary Material.

Docking scores indicate that FtsZ contains the most promising drug target, while CoaD which has a similar appearance (Figure 7c), also scores relatively well. Figures 11a, 11b, 11c, and 11d show the top scoring ligand for each protein docked with FtsZ, CoaD, YwtF, and RacE respectively. We noted that the top scoring ligands for FtsZ tend to have a bend similar to the one shown in Figure 11a such that they conform closely to the contours of the pocket, which is also a characteristic of high scoring ligands for the other proteins. This is consistent with other research, which has found that in most cases the largest pocket/cavity is the active site [39].

## 5.4 Validation

We validated our *in silico* results using against the data contained in the Protein Data Bank pdb files and ZINC mol2 ligand files.

(1) Highly conserved regions in FtsZ and CoaD were compared with their known active sites. We found that the locations of highly conserved residues corresponded very closely to known active sites.

(2) Although Vina's accuracy was confirmed in [8], we compared the predicted binding site of lig-

<b>Proteomes</b>	<b>Hits</b>	<b>mean id.</b>	<b>UniProt #</b>	<b>GN</b>	<b>3D</b>
54	54	55%	P17865	<i>ftsZ</i>	yes
51	70	50%	P70965	<i>murAA</i>	no
51	70	47%	P19670	<i>murAB</i>	no
51	61	46%	O31751	<i>uppS</i>	no
50	54	39%	Q03523	<i>murE</i>	no
49	49	46%	O34797	<i>coaD</i>	yes
48	49	46%	P14192	<i>glmU</i>	no
48	60	38%	P40778	<i>murC</i>	no
47	77	34%	P31114	<i>hepT</i>	no
46	80	39%	P54383	<i>ispA</i>	no
46	66	47%	O31822	<i>yngB</i>	no
46	64	51%	Q05852	<i>gtaB</i>	no
45	45	36%	P96613	<i>murF</i>	no
44	57	39%	P96612	<i>ddl</i>	no
43	46	37%	Q06755	<i>ispD</i>	no
42	43	37%	Q03522	<i>murD</i>	no
41	55	43%	P54523	<i>dxs</i>	no

Table 3: *B. subtilis* query proteins versus 60 Proteomes

<b>Proteomes</b>	<b>Hits</b>	<b>mean id.</b>	<b>UniProt #</b>	<b>GN</b>	<b>3D</b>
25	25	63%	P17865	<i>ftsZ</i>	yes
25	25	51%	P14192	<i>glmU</i>	no
25	25	50%	O34797	<i>coaD</i>	yes
25	34	48%	O31751	<i>uppS</i>	no
24	27	41%	Q03523	<i>murE</i>	no
23	29	43%	P96612	<i>ddl</i>	no
21	24	47%	P94556	<i>race</i>	yes
21	58	34%	Q7WY78	<i>ywtF</i>	yes
19	20	34%	O05412	<i>yrpC</i>	no
16	18	51%	O31753	<i>dxr</i>	no
16	33	37%	P39581	<i>dltA</i>	yes
15	16	60%	P54482	<i>ispG</i>	no

Table 4: *B. subtilis* query proteins versus 25 Gram Positive Proteomes

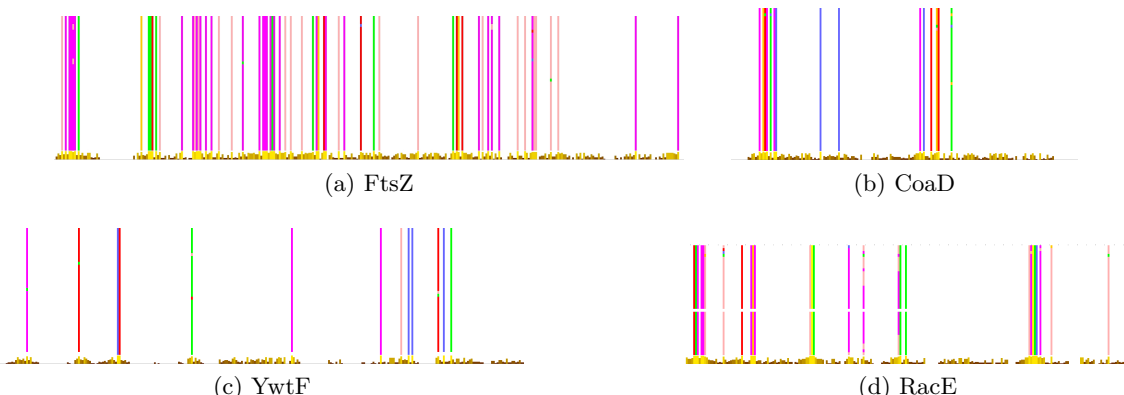


Figure 6: Multiple sequence alignments: columns are in color where residues are 100% conserved. The FtsZ sequence shows the greatest degree of conservation.

ands similar to the organic FtsZ ligand, CIT, with the images in Figure 10 indicate, the organic ligand's experimentally determined binding site. As and's known position closely matches the position

<b>Proteomes</b>	<b>Hits</b>	<b>mean id.</b>	<b>UniProt #</b>	<b>GN</b>	<b>3D</b>
25	25	47%	P17865	<i>ftsZ</i>	yes
25	26	46%	P70965	<i>murAA</i>	no
25	25	45%	O31751	<i>uppS</i>	no
25	26	42%	P19670	<i>murAB</i>	no
25	27	36%	Q03523	<i>murE</i>	no
24	24	42%	O34797	<i>coaD</i>	yes
23	25	40%	P14192	<i>glmU</i>	no
23	43	37%	P54383	<i>ispA</i>	no
23	39	32%	P31114	<i>hepT</i>	no
23	34	32%	P40778	<i>murC</i>	no
22	22	56%	Q06756	<i>ispF</i>	no
22	28	42%	P54523	<i>dxs</i>	no
22	22	33%	P96613	<i>murF</i>	no
21	27	46%	Q05852	<i>gtaB</i>	no
21	31	43%	O31822	<i>yngB</i>	no
21	29	36%	P96612	<i>ddl</i>	no
21	43	34%	P38422	<i>dacF</i>	no
21	22	32%	Q03522	<i>murD</i>	no
20	20	36%	Q06755	<i>ispD</i>	no
19	20	37%	P54473	<i>ispH</i>	no
19	19	34%	O05412	<i>yrpC</i>	no
19	40	33%	P35150	<i>dacB</i>	no
18	18	41%	O31753	<i>dxr</i>	no
17	95	37%	P96740	<i>pgdS</i>	no

Table 5: *B. subtilis* query proteins versus 25 Gram Negative Proteomes

<b>Proteomes</b>	<b>Hits</b>	<b>mean identity</b>	<b>UniProt #</b>	<b>GN</b>	<b>3D</b>
1	1	100%	P17865	<i>ftsZ</i>	yes
1	1	100%	O34797	<i>coaD</i>	yes
1	1	100%	P39844	<i>dacC</i>	yes
1	1	100%	P50740	<i>fni</i>	yes
1	1	100%	P39131	<i>mnaA</i>	yes
1	1	100%	P27623	<i>tagD</i>	yes
1	2	66%	O05412	<i>yrpC</i>	no
1	2	66%	P94556	<i>racE</i>	yes
1	3	59%	P96499	<i>yvhJ</i>	no
1	3	59%	Q02115	<i>lytR</i>	no
1	3	58%	Q7WY78	<i>ywtF</i>	yes

Table 6: Redundancy of 123 *B. subtilis* proteins in the *B. subtilis* Proteome. Using BLAST with an E value of 0.1 and filtered with a sequence identity threshold of  $\geq 30\%$ .

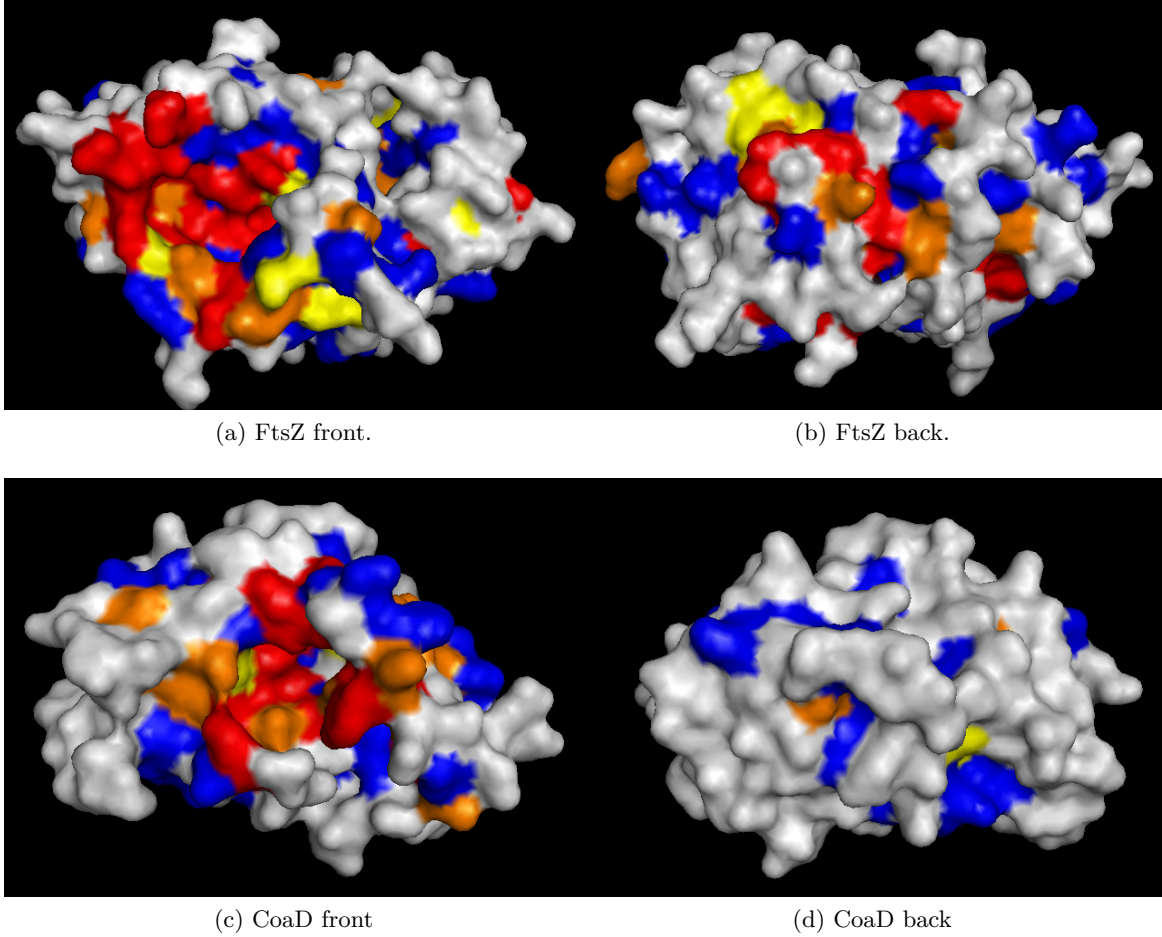


Figure 7: FtsZ and CoaD with surface colored by conservation score: gray {0..5}, blue {6,7}, yellow {8}, orange {9}, red {10,11}.

Protein Name	UniProt #	pdb id	Top Scoring Ligand	Score kcal/mol
FtsZ	P17865	2VXY	ZINC04236084	-12.10
CoaD	O34797	1O6B	ZINC03846570	-10.20
YwtF	Q7WY78	3MEJ	ZINC08791231	-9.10
RacE	P94556	1ZUW	ZINC03844349	-7.70

Table 7: Top scoring NPD ligands using predicted site

predicted *in silico* for a virtually identical ligand.

(3) We compared the docking scores of a set of 10777 NPD ligands (p0.4) when docked within a search area centered on our predicted active sites versus docking scores for the same set of ligands in a search space of the same size but centered on a poorly conserved region on the opposite side of each protein. Although results for FtsZ and CoaD were as expected – the ligands docked on the “back” side did not score as well – ligands docked on the “back” side of YwtF and CoaD scored better, Table 8. This was quite interesting. However, Figure 12c shows that the top scoring ligand for YwtF in this case is docked deep in an elaborate pocket characterized by some conservation (blue

={6,7} on the scale from 0 to 11), whereas the “front” side of YwtF presents a large highly conserved area, Figure 11c, lacks a substantial pocket. In the case of RacE, Figure 12d shows the top scoring ligand docked in a small pocket in a relatively conserved area which, in hindsight, is not obviously a less likely binding site than the one shown in Figure 11d.

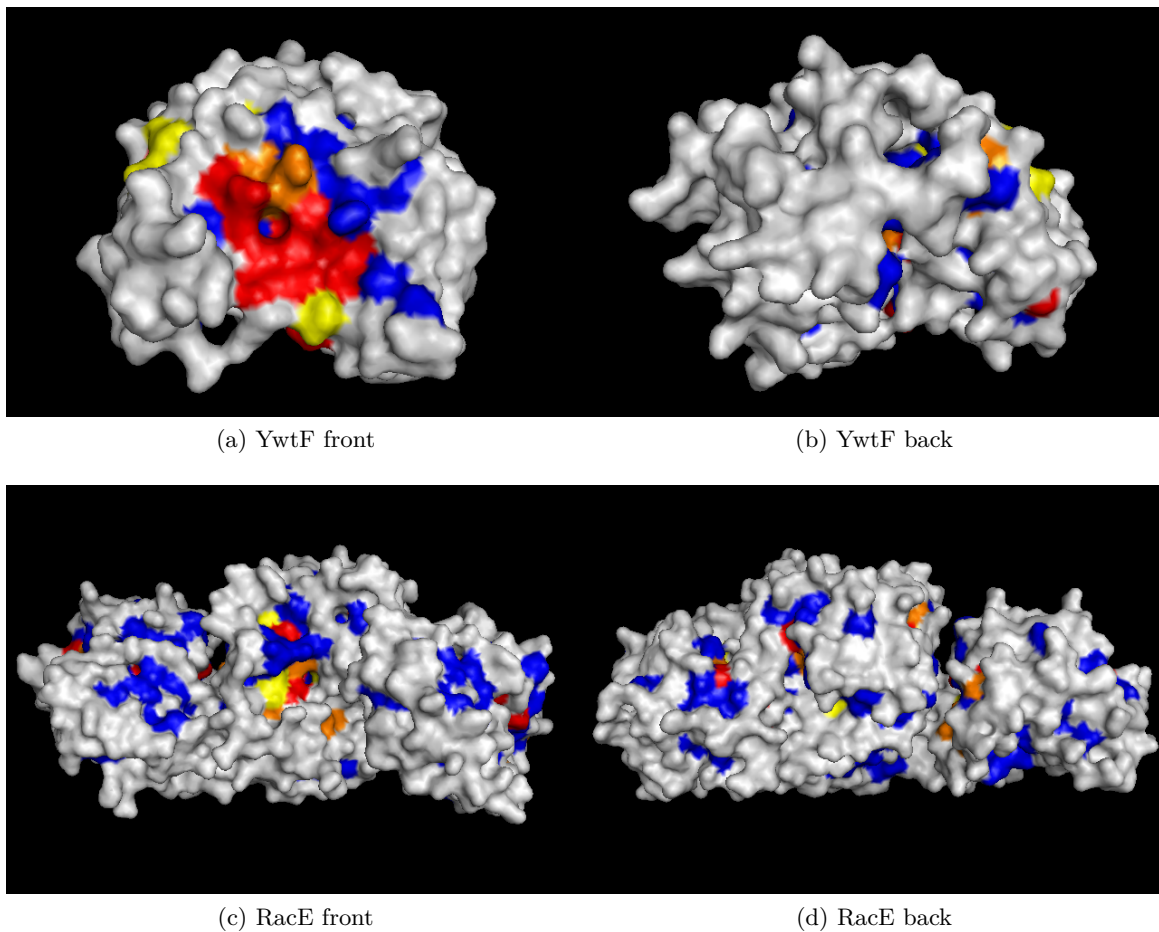


Figure 8: FtsZ and CoaD with surface colored by conservation score: gray {0..5}, blue {6,7}, yellow {8}, orange {9}, red {10,11}.

receptor	$mean_a$	$mean_b$	$min_a$	$min_b$	$max_a$	$max_b$
FtsZ	-8.24	-5.89	-11.90	-8.40	-3.40	-2.50
CoaD	-6.97	-4.61	-9.60	-6.60	-2.80	-1.90
YwtF	-6.41	-6.75	-9.00	<b>-11.40</b>	-2.60	-2.70
RacE	-5.21	-5.64	-7.60	-8.30	-2.10	-2.60

Table 8: Docking Scores for NPD p0.4 ligands: search areas include predicted active site (a) versus predicted inactive site (b).

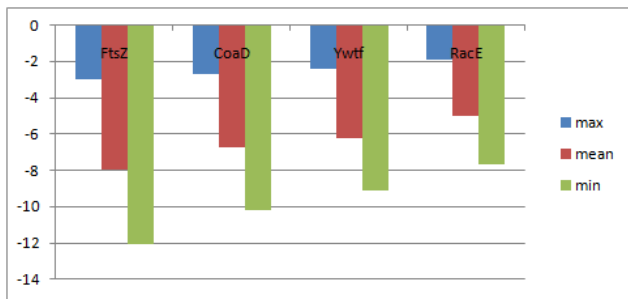


Figure 9: Docking scores in kcal/mol for NPD ligands. A more negative value indicates a lower binding energy.

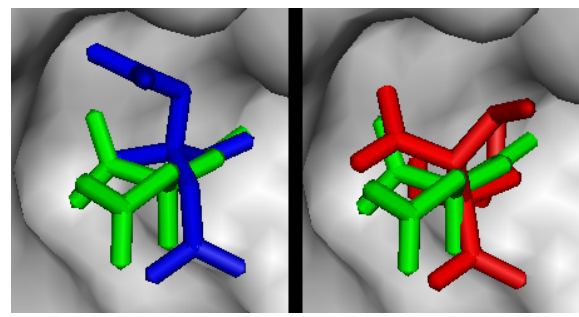


Figure 10: FtsZ with organic ligand (green) in known position versus docked ligands (a) and (b).

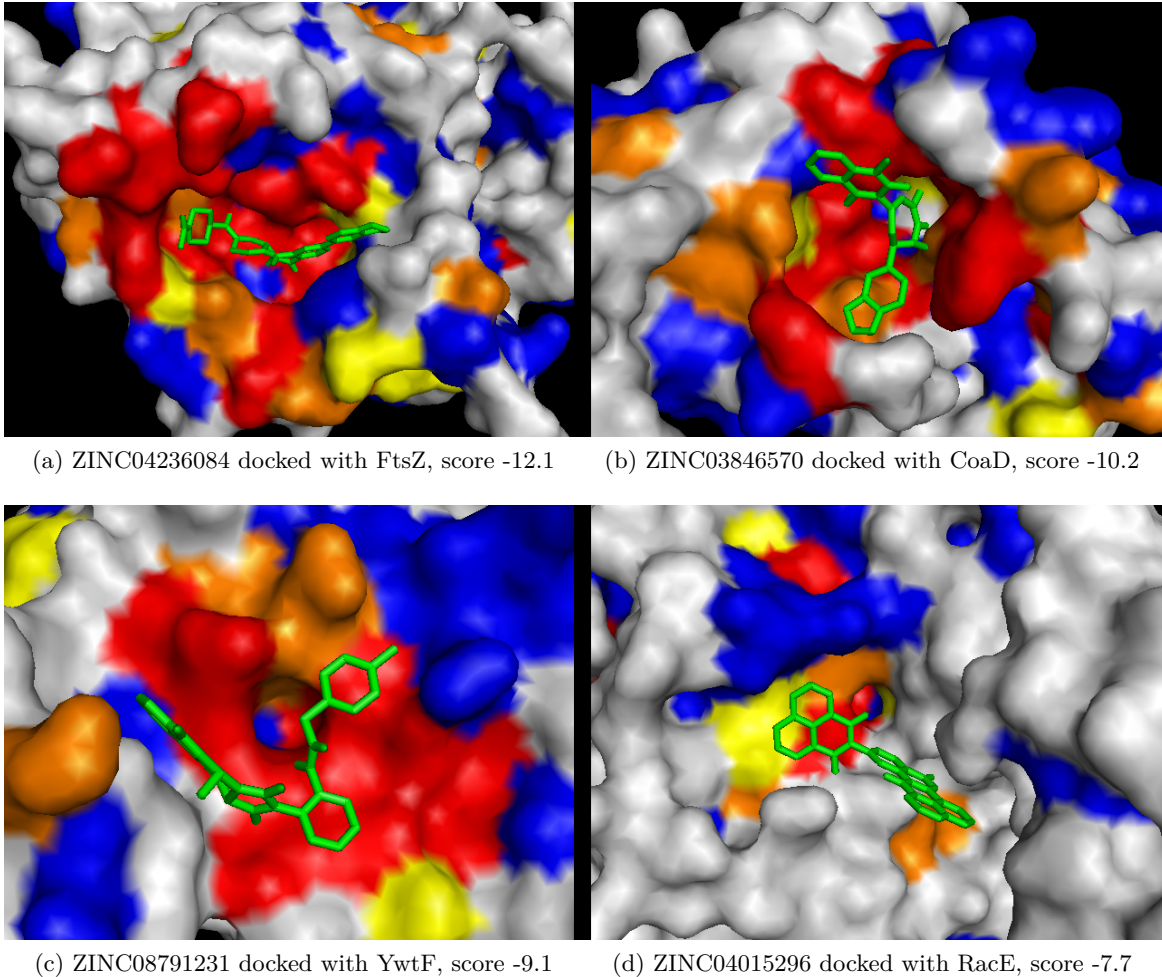


Figure 11: Top scoring NPD ligands (complete set) docked in search areas that include **predicted active sites**.

## 6 Discussion

In this thesis we considered 123 genes associated with bacterial cell wall biosynthesis and cell division from the *B. subtilis* genome. From these genes we used a comparative genomics approach to identify essential proteins in multiple bacterial species because the proteins most conserved across species are most likely to be essential for the survival of more species of bacteria [3] and, therefore, good potential drug targets [17; 56]. We found that the conserved residues in each protein coincided with either known active sites, or sites where the best scoring ligands docked with the lowest binding energy. However, while our results indicate that the combination of very highly conserved residues and a substantial pocket clearly indicate a potential drug target, less favorable conservation scores and surface topology indicate the need for further investigation and probably a larger search space for the docking software.

While the images of proteins presented in this

paper may appear to be precisely colored according to conservation and, therefore, functional importance, we can only say that based on the data and algorithms used, some surface regions are probably more important than others. First, the sequences used for MSA depended not only on the set of input sequences but on BLAST (which uses heuristics) and its parameters. Second, the MSAs produced by MUSCLE may differ from those produced by other widely used algorithms and, of course, all are probabilistic. Different MSAs may yield different conservation scores and different 3D images. Third, blasting against a different set of proteomes may also lead to different conservation scores. Fourth, the results from virtual screening depended on multiple factors including the set of ligands and the search parameters used by AutoDock Vina. Another cause for concern is that the 3D structure of the vast majority of proteins is not known, which means our investigation has in fact been limited to a small minority of *B. subtilis*

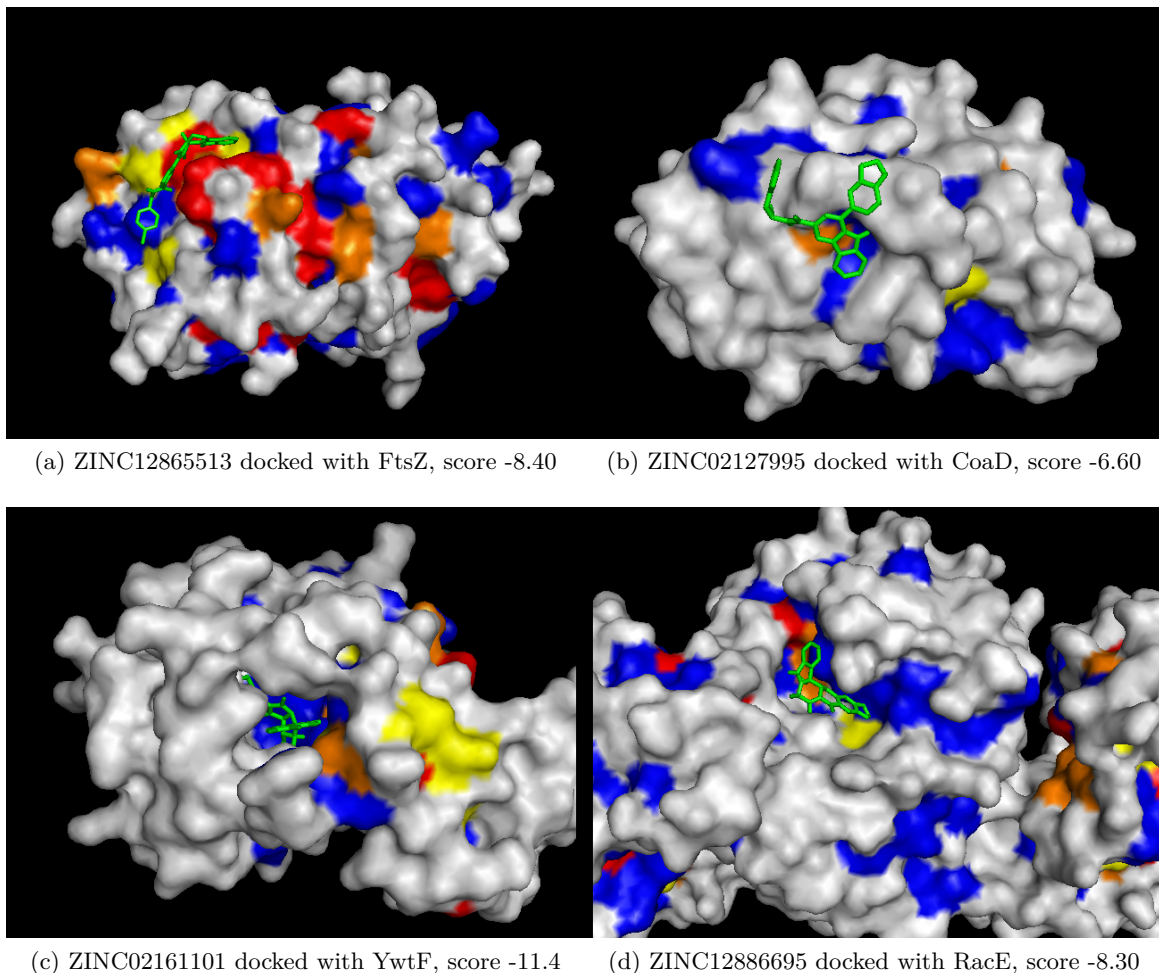


Figure 12: Top scoring NPD p0.4 ligands docked in search areas that include **predicted inactive sites** (using 10777 ligands instead of  $\approx 90000$ ).

proteins.

## 7 Future Work and Conclusion

The scope for future work includes many areas. First, this work could be extended to consider the 3D structure of proteins similar to essential *B. subtilis* proteins. Second, additional methods and resources could be combined with ours to form an ensemble with greater accuracy. Third, the data analyzed in this paper should be amenable to machine learning techniques. For example, which features of a chemical compound are the best predictors binding energy may depend on multiple factors and the receptor protein, but this could be learned from the data produced by our methods and data from online databases such as ZINC, UniProt, PDB, and many others.

In conclusion, we found that the methods presented provide a principled and rational way to reduce costs and manage complexity. Given an

extremely large number of possible protein ligand combinations, our results were used to decide which proteins to target, which ligands to purchase, and thereby prioritize *in vitro* experiments designed to test compounds for antimicrobial properties.

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