Biophysical And Biochemical Studies of Penicillin Binding Proteins And Novel PBP3 Inhibitors

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University of Warwick

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To my friends and family who have listened, encouraged and supported me throughout: I could not have done this without you. Thank you so much.
Declaration

This thesis is submitted to the University of Warwick in support of my application for the degree of Doctor of Philosophy. It has been composed by myself and has not been submitted in any previous application for any degree.

The work presented (including data generated and data analysis) was carried out by the author except in the cases outlined below:

- Dom Bellini (LMB, Cambridge) prepared PBP3 proteins used in Chapter 2 and Chapter 5; Chris Graham (University of Warwick) provided MreC protein (Chapter 2); Julie Todd (University of Warwick) provided some proteins used in Chapter 6.
- Sarah Bennett (University of Warwick) operated the liquid handling system during antimicrobial assays (Chapter 3).
- Mukesh Gangar (University of Cape Town) synthesised Compound 16 (Chapter 5). Robert Lesniak (Stanford University) synthesised S2d (Chapter 5).
- Dom Bellini crystallised and collected data for all crystal structures in Chapter 5, but these were refined by the author. Dom Bellini crystallised, collected data and refined (with input from the author) crystal structures in Chapter 6. All other crystal structures are taken from the protein data bank (PDB code referenced in the text).
- Joe Eyermann (University of Cape Town) generated the docked model of a PBP:12 complex (Chapter 5).
- \(^{13}\)C NMR for compounds in Chapter 5 were collected and assigned by Alen Krajnc (University of Oxford). High resolution mass spectra were collected by Thomas Corner (University of Oxford).
- Mass Spectrometry of PBP:nitrocefin complexes (Chapter 6) was performed by Alen Krajnc and Anthony Tumber (University of Oxford).
• Proteomic Mass Spectrometry was performed by Warwick Proteomics Research Technology Platform (sample preparation was performed by the author).

Parts of this thesis have been published by the author:


This work formed the basis for Chapter 4 and is summarised in section 4.1.


The author contributed structural activity relationship input to this work, discussed in section 1.3.3 and published two crystal structures (PDB: 6Y6U and 6Y6Z)


Chapter 5 was modified and submitted for publication: it was accepted, reviewed, revised and is currently returned to the reviewers for their approval (as of 11th June 2021). As part of this, 10 crystal structures (5.S2) were published on the PDB.

All three papers are included in the appendix.
Abstract

Antimicrobial resistance presents a worrying and growing threat to modern medicine. Combatting this challenge requires the discovery of new antimicrobial compounds, which are insensitive to current resistance pathways. One of the most important classes of antibiotics, β-lactams, target penicillin binding proteins (PBP3), a family of enzymes that synthesise the bacterial cell wall. In most clinically important gram-negative bacteria, PBP3 is a ubiquitous and essential drug target, whose inhibition can prevent the growth of bacterial cells. Novel mode of action inhibitors of PBPs are needed to combat rising resistance to β-lactams, but finding such inhibitors requires improved understanding of these proteins and new discovery methods.

This thesis describes the development of two tools for inhibitor discovery: biophysical methods (surface plasmon resonance) in an attempt to characterise the interactions of PBP3 with its ligands; and a high throughput microbiology platform designed to provide low cost, low volume, rapid screening of early stage compounds.

PBP3 mutation-mediated resistance mechanisms have been investigated by analysing structural data to provide insight into how mutations distal to the active site may lower the susceptibility of PBPs to β-lactams. Novel compounds are used to show the indirect link between the active site and these distant regions.

A crystallographic fragment screen, and follow up medicinal chemistry, was used to identify benzoxygenoroles as inhibitors of PBP3 with a novel, di-covalent binding mode, engaging two highly conserved serines in the active site. Studies with a non-clinical probe β-lactam, nitrocefin, also appear to show an unprecedented binding mode involving engagement of two serines. These results introduce benzoxygenoroles and other compounds as warheads that can bind both serine residues, thereby opening up new avenues for research in the pursuit of non-β-lactam PBP inhibitors.
## Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>2,2-DTP</td>
<td>2,2'-Dithiodibispyridine</td>
</tr>
<tr>
<td>$A_{\text{xxx nm}}$</td>
<td>Absorbance at the specified (XXX) wavelength</td>
</tr>
<tr>
<td>ABC</td>
<td>Ammonium Bicarbonate</td>
</tr>
<tr>
<td>AbPBP3</td>
<td>Acinetobacter baumannii PBP3</td>
</tr>
<tr>
<td>AMR</td>
<td>Antimicrobial resistance</td>
</tr>
<tr>
<td>ATCC</td>
<td>American Type Culture Collection</td>
</tr>
<tr>
<td>αN/ βN</td>
<td>Notation for secondary structure of proteins; α: alpha helices; β: beta strands. N refers to its index (in order N-C terminal). Two codes separated by a dash e.g. α11-β5 indicates the loop between these structures</td>
</tr>
<tr>
<td>caMHB</td>
<td>Cation-adjusted Mueller-Hinton broth 2</td>
</tr>
<tr>
<td>CDI</td>
<td>1,1'-carbonyldiimidazole</td>
</tr>
<tr>
<td>CENTA</td>
<td>$\text{((6R,7R)-3-(((3\text{-carboxy-4-nitrophenyl})\text{-thio) methyl})-8\text{-oxo-7-(2-(thiophen-2-yl)acetamido)-5-thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid}}\text{)}}$</td>
</tr>
<tr>
<td>CLSI</td>
<td>Clinical &amp; Laboratory Standards Institute</td>
</tr>
<tr>
<td>CFU</td>
<td>Colony forming units</td>
</tr>
<tr>
<td>DHFR</td>
<td>Dihydrofolate reductase</td>
</tr>
<tr>
<td>DMF</td>
<td>N,N-dimethylformamide</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl Sulphoxide</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
</tr>
<tr>
<td>DOS</td>
<td>Diversity orientated synthesis</td>
</tr>
<tr>
<td>DTNB</td>
<td>5,5-dithio-bis-(2-nitrobenzoic acid)</td>
</tr>
<tr>
<td>DTT</td>
<td>Dithiothreitol</td>
</tr>
<tr>
<td>EC50/90</td>
<td>Effective concentration at which 50/90 % of the growth is inhibited</td>
</tr>
<tr>
<td>Term</td>
<td>Definition</td>
</tr>
<tr>
<td>--------</td>
<td>------------------------------------------------</td>
</tr>
<tr>
<td>EcPBP3</td>
<td>Escherichia coli PBP3</td>
</tr>
<tr>
<td>EDC</td>
<td>1-ethyl-3-(3-dimethylaminopropyl)carbodiimide</td>
</tr>
<tr>
<td>EDTA</td>
<td>Ethylene diamine tetraacetate</td>
</tr>
<tr>
<td>equiv.</td>
<td>Equivalents</td>
</tr>
<tr>
<td>ER</td>
<td>Ensemble refinement</td>
</tr>
<tr>
<td>ES(I)</td>
<td>Electrospray (ionisation)</td>
</tr>
<tr>
<td>FA</td>
<td>Fluorescence Anisotropy</td>
</tr>
<tr>
<td>FBDD</td>
<td>Fragment based drug discovery</td>
</tr>
<tr>
<td>FIC(I)</td>
<td>FIC(I): fractional inhibitory concentration</td>
</tr>
<tr>
<td></td>
<td>(index)</td>
</tr>
<tr>
<td>g</td>
<td>Gravitational Force</td>
</tr>
<tr>
<td>GC</td>
<td>Gonococcal</td>
</tr>
<tr>
<td>GSK</td>
<td>GlaxoSmithKline</td>
</tr>
<tr>
<td>GT</td>
<td>Glycosyltransferase</td>
</tr>
<tr>
<td>HiPBP3</td>
<td>Haemophilus influenzae PBP3</td>
</tr>
<tr>
<td>HMM</td>
<td>High molecular mass</td>
</tr>
<tr>
<td>HPLC</td>
<td>High Performance Liquid Chromatography</td>
</tr>
<tr>
<td>HRMS</td>
<td>High resolution mass spectrometry</td>
</tr>
<tr>
<td>IPTG</td>
<td>Isopropyl β- d-1-thiogalactopyranoside</td>
</tr>
<tr>
<td>K_i</td>
<td>Kinetic constant: defined in Scheme 5.1.</td>
</tr>
<tr>
<td>K_M</td>
<td>Kinetic constant: defined in equation 6.55</td>
</tr>
<tr>
<td>K_s0</td>
<td>Kinetic constant: defined in equation 6.56</td>
</tr>
<tr>
<td>k_cat</td>
<td>Enzyme turnover number: defined in equation 6.7</td>
</tr>
<tr>
<td>k_x</td>
<td>Rate constant for the indicated step</td>
</tr>
<tr>
<td>LB</td>
<td>Lysogeny broth</td>
</tr>
<tr>
<td>LB agar</td>
<td>Lysogeny broth agar</td>
</tr>
<tr>
<td>LC</td>
<td>Liquid chromatography</td>
</tr>
<tr>
<td>LCMS</td>
<td>Liquid chromatography mass spectrometry</td>
</tr>
<tr>
<td>LMM</td>
<td>Low molecular mass</td>
</tr>
<tr>
<td>M</td>
<td>Species mass (in the context of mass</td>
</tr>
<tr>
<td></td>
<td>spectrometry)</td>
</tr>
<tr>
<td>MCA</td>
<td>Multichannel arm</td>
</tr>
<tr>
<td><strong>Abbreviation</strong></td>
<td><strong>Full Form</strong></td>
</tr>
<tr>
<td>-----------------</td>
<td>--------------</td>
</tr>
<tr>
<td>MD</td>
<td>Molecular dynamics</td>
</tr>
<tr>
<td>MIC</td>
<td>Minimum inhibitory concentration</td>
</tr>
<tr>
<td>MS</td>
<td>Mass spectrometry</td>
</tr>
<tr>
<td>MR</td>
<td>Relative molecular mass,</td>
</tr>
<tr>
<td>MRSA</td>
<td>Methicillin-resistant Staphylococcus aureus</td>
</tr>
<tr>
<td>NCTC</td>
<td>National Collection of Type Cultures</td>
</tr>
<tr>
<td>NgPBP2</td>
<td>Neisseria gonorrhoeae PBP2</td>
</tr>
<tr>
<td>NHS</td>
<td>N-hydroxysuccinimide</td>
</tr>
<tr>
<td>NMR</td>
<td>Nuclear magnetic resonance</td>
</tr>
<tr>
<td>OD&lt;sub&gt;600&lt;/sub&gt;</td>
<td>Optical density at 600 nm</td>
</tr>
<tr>
<td>PaPBP3</td>
<td>Pseudomonas aeruginosa PBP3</td>
</tr>
<tr>
<td>PBP</td>
<td>Penicillin binding protein</td>
</tr>
<tr>
<td>PDB</td>
<td>Protein data bank</td>
</tr>
<tr>
<td>PDEA</td>
<td>2-(2-pyridinylthio)ethaneamine hydrochloride</td>
</tr>
<tr>
<td>pI</td>
<td>Protein isoelectric point</td>
</tr>
<tr>
<td>pIC&lt;sub&gt;50&lt;/sub&gt;</td>
<td>Negative logarithm of half maximal inhibitory concentration</td>
</tr>
<tr>
<td>pK&lt;sub&gt;a&lt;/sub&gt;</td>
<td>Negative logarithm of the acid dissociation constant</td>
</tr>
<tr>
<td>SA</td>
<td>Streptavidin (an SPR Chip)</td>
</tr>
<tr>
<td>SDS PAGE</td>
<td>Sodium dodecyl sulphate polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>SMILES</td>
<td>Simplified molecular-input line-entry system</td>
</tr>
<tr>
<td>SOC</td>
<td>Super optimal broth with catabolite repression</td>
</tr>
<tr>
<td>SPR</td>
<td>Surface plasmon resonance</td>
</tr>
<tr>
<td>TCEP</td>
<td>Tris(2-carboxyethyl)phosphine</td>
</tr>
<tr>
<td>TFA</td>
<td>Trifluoroacetic acid</td>
</tr>
<tr>
<td>TG</td>
<td>transglycosylation</td>
</tr>
<tr>
<td>TLS</td>
<td>Translation/libration/screw - a refinement parameter</td>
</tr>
</tbody>
</table>
TOF  Time of flight
TP   Transpeptidation
Tris Tris(hydroxymethyl)aminomethane
RA   Residual activity

$R_{\text{max}}$  Theoretical maximal amount of analyte that can be bound to the chip. Defined in equation 2.2

$R_{\text{XXX}}$  Response of an SPR detector. XXX refers to the binding species
UDP  Uridine diphosphate
UMP  Uridine monophosphate
UPLC  Ultra Performance Liquid Chromatography
rpm  Revolutions per minute
rt   Room temperature

$V_{\text{max}}$  Maximal velocity of an enzyme-substrate interaction
RMSD  Root-mean-square deviation
RoMa  Robotic manipulator arm
RU   Response units
v/v  Volume to volume ratio
wt   Wild type
w/v  Weight to volume ratio

Other notation

[XXX]  Concentration of XXX

Amino acid substitutions are referred to as e.g. S349A: meaning the substitution of a serine residue for an alanine residue at the indicated (numerical) position.

In chapter 6 subscripts are used to indicate the substitution mutations (or wt) of a protein: e.g.: PaPBP3$_{S349A}$ is the S349A substitution mutation of PaPBP3
List of Chemical Structures

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“[W]e shall therefore continue to labour under discouragement so long as we are faced with the bugbear of drug resistance.”

Sir Charles Harington, 1957
Chapter 1. Introduction

1.1 Antimicrobial Resistance

Antimicrobials have been described as the “Fire extinguishers of medicine” \(^1\), in an analogy that elegantly captures their role: mundane and overlooked yet essential. Their use allows for the safe undertaking of surgery \(^2\) and cancer treatment \(^3\) as well as treatment of all types of infections within the community \(^4\). Their ubiquity in modern health systems has led to rising cases of resistance, with bacteria evolving mechanisms which reduce the efficacy of antibiotics \(^5,6\).

Resistance to antibiotics is an ancient \(^7-9\) and inevitable \(^10\) phenomenon, resulting from the historical co-evolution of antibiotic-resistant organisms alongside antibiotic-producing ones. Before the widespread use of antibiotics, it was predicted (by Fleming himself) that their “underdosage” would lead to the proliferation of resistance. It is not possible to prevent the development of resistance in the long-term and at the moment the only strategy against it has been to innovate with new drugs at a faster rate than the development of resistance to the current therapies. During the “golden age” of antimicrobial discovery (1930-1962) this was successful, with 20 classes of antibiotic discovered. Since then only 2 new classes of antibiotics have been marketed\(^11\). As drugs discovered more than 60 years ago become increasingly ineffective, a serious and growing lack of treatment options are emerging. The lack of new drugs within the antimicrobial sector is not a problem encountered across the rest of the pharmaceutical industry, which continues to discover drugs of all classes at a high rate \(^12\).

The combined challenges of low discovery rate and the development of resistance are creating a “slow tsunami” \(^13\) of untreatable infections. Both of these topics shall be discussed briefly below, but they are interlinked. The continual development of resistance creates the imperative for novel drug discovery, and resistance slows discovery efforts by introducing barriers rarely encountered in other areas of pharmaceutical discovery science. Before investigating these twin issues, it is worth considering how our current antibiotics were discovered.
1.1.1 A Brief History of Antimicrobial Drug Discovery

Perhaps the first modern antimicrobial screen was that of Paul Ehrlich in 1910 in his efforts to discover an effective treatment for syphilis by \textit{in vivo} screening of syphilitic rabbits \textsuperscript{14,15}. In the 1930s, \textit{in vivo} screening of mice with streptococcal sepsis was similarly used to discover pronosil \textsuperscript{15,16}, an antibiotic prodrug and the first in class of the sulfonamide “sulfa” drugs, of which many derivatives were later made \textsuperscript{17} such as the essential medicine sulfamethoxazole \textsuperscript{18}.

The most famous of the early antibiotics was penicillin \textsuperscript{19,20}. Whilst the antimicrobial activity of moulds was known, from 1928 onwards Fleming studied them systematically and quantitatively, by streaking bacteria over penicillin-spiked agar \textsuperscript{19}. The successful extraction of penicillin in 1940 \textsuperscript{20} and subsequent “public-private” partnership established to mass produce it during the second world war stimulated the US pharmaceutical industry \textsuperscript{21,22} and led to the start of the “golden age” of antibiotic discovery. During this period, almost all current classes of antibiotics \textsuperscript{23}, as many as 250 individual compounds \textsuperscript{21} were marketed as systematically administratable products. Many of these compounds came from screening soil samples to identify active compounds, beginning with the work of Hotchkiss and Dubos \textsuperscript{24} who discovered gramicidin and tyrocidine and later works using \textit{actinomyces} species to extract streptomycin \textsuperscript{25} (notable for being the first effective treatment for tuberculosis \textsuperscript{26,27}). Waksman, the Nobel prize-winning inventor of streptomycin, in fact coined the term antibiotic in 1941. In this period (1949-1962), chloramphenicol, tetracyclines, aminoglycosides, macrolides then glycopeptides, streptogramins and quinolones were discovered in quick succession \textsuperscript{28}. Only quinolones were fully synthetic, discovered (apparently serendipitously) as a byproduct of chloroquine production \textsuperscript{29,30}. In 2000, following a chemical screening program at Du Pont \textsuperscript{31}, oxazolidinones were released as the first novel class of inhibitors in 40 years. These represent the third class of synthetic antibiotics, with the rest all coming from natural product origins \textsuperscript{32}.

In summary, the antibiotics we have today benefited in their discovery from government investment, a lack of competing compounds on the market, and readily available natural products produced by soil microorganisms, which could be developed into drugs relatively quickly by medicinal chemistry techniques. Often, as is the case with penicillin, the intracellular target was discovered many years later \textsuperscript{33}. 

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and the fundamental biochemistry was not considered important for the drug
discovery efforts.\textsuperscript{34,35}

### 1.2 Challenges of Antibiotic Resistance

In contrast, the past few decades have yielded few compounds with genuinely novel
mechanisms of action and the pipeline of upcoming drugs remains empty.\textsuperscript{36} The
challenges of antimicrobial resistance are three-fold: socio-economical, regulatory
and scientific (Figure 1.1).

![Figure 1.1. Summary of the various challenges related to antimicrobial resistance](image)

**1.2.1 A Tripod of Socio-economic Challenges**

Steven J. Hoffman and Kevin Outterson have described antimicrobial resistance as
a tripod of three intersecting and mutually supporting pillars, each representing an
individual challenge within the overarching AMR crisis.\textsuperscript{37} These pillars are: access,
innovation and conservation. Solutions to AMR will require that each of these challenges is addressed: access is necessary to ensure that these essential medicines can be used by all: currently more people are killed by antibiotic-treatable infections than from resistant organisms \(^{38,39}\). It would be unethical and counterproductive \(^{40}\) to address AMR in rich countries without simultaneous work to lower the burden of infectious disease as a whole. Conservation and appropriate use of antibiotics will ensure that resistance is kept at a minimum level to allow successful cycles of drug development to keep pace with drug resistance in the clinic. In this way, public-awareness and a “One Health” approach to antibiotics, which encompasses limitations on their use in veterinary practice \(^{38,41}\), and better control of their end of life in wastewater treatment plants \(^{38,42}\) become crucial to ensuring innovation efforts are not wasted.

In their use of the tripod analogy, Hoffman and Outterson \(^{37}\) recognised that a singular focus on any of the three issues in isolation would be insufficient to prevent the rising antibiotic crisis leading us to a post-antibiotic world. Whilst the majority of this thesis is focussed on the technical aspects of the “innovation” leg of the tripod, it is worth briefly covering the other two to put the need for (and limits of) innovation in context.

Stymying attempts to challenge each of these three pillars is the fact that each has characteristics of a “commons” problem, one in which individuals can benefit in the short term by acting selfishly but that over the long term has severe negative effects on all stakeholders \(^{43}\). For example, in antimicrobial conservation a single prescriber can inappropriately prescribe antibiotics, driving resistance, with the consequences spreading outside their community whilst they benefit by quickly getting the patient out of the consulting room and higher patient satisfaction \(^{44}\). In this example the “prescriber” could be a single hospital with insufficient guidelines, or entire nations failing to correctly regulate antibiotic use, thereby spreading resistance across the globe.

Within innovation there is a similar problem. Antibiotic development, as with typical drug development, is very expensive; the “all-in” cost of bringing a modern drug to market is estimated at around $1.7 billion \(^{45}\). Given that a new antibiotic should be carefully regulated in its use, a novel compound may struggle to recoup this cost before exclusivity rights expire \(^{38,46,47}\). In the majority of cases antibiotics are only prescribed for short periods (~ weeks) compared to drugs for chronic conditions
which may require treatment over years or for the rest of the patient’s life. This reduces the sales volume and may drive pharmaceutical companies to different therapeutic areas. Poor sales after release have led to the recent failures of companies launching new antibiotics: Melinta and Achaogen. As such, new models are needed to “delink” the relationship between drug sales and payments to the drug developers. Multiple models have been proposed to do this. Fundamentally, a model must recognise the value of antibiotics to society and finance the developers appropriately. The social value of an antibiotic for the treatment of hospital acquired/ventilator associated bacterial pneumonia antibiotics for example, is estimated to be worth $12 billion. In this context, the development cost of a new compound is justifiable.

Even in the event of public money being used for antimicrobial drug development, global access could still be a commons problem as only rich countries can afford to pay but all countries will need access to the new drugs as resistance spreads without recognition of borders. Policy makers and the public will need to understand that this apparent altruistic and ethical behaviour is actually essential to slow the spread of resistance globally.

Commons problems must be addressed through well designed and well managed international agreements with buy-in from all nations. Politicians and the public must be engaged with these challenges now. Lessons in this can be learned from the concurrently burgeoning commons problem of the climate crisis.

1.2.2 Regulatory Challenges

Companies now face an increasing burden of regulation by the approval agencies (greater patient size and increasingly stringent non-inferiority trials) and postmarketing obligations. The cost of these expensive trials cannot be recouped by drugs that are 3 times less profitable than cancer drugs and 11 times less profitable than musculoskeletal drugs, so corporate policy has shifted budgets towards these more profitable areas. In addition to the cost, recruiting patients to trials can be difficult as those with the most intractable and drug-resistant infections are rare and often critically ill, but trials must start as soon as the patient is admitted to hospital and before they receive other treatments.
1.2.3 Scientific Challenges

1.2.3.1 Big Pharma in a post-golden age

Following the “golden age” of antibiotic discovery (see A Brief History of Antimicrobial Drug Discovery above) new chemical matter to fill the pipeline began to become increasingly rare and the first wave of scaling back by the pharmaceutical industry began. Advances in high throughput biochemical screening from the 1980s onwards using largely synthetic “robotically mass produced” chemical libraries and the start of the “genomic era” (following the publication of the first complete genome sequence of Haemophilus influenzae in 1995), failed to bring in a new wave of antibiotic discoveries. After a decade of Big Pharma efforts, it appeared a new golden era would not be forthcoming, and a second wave of exits occurred.

Researchers at Merck noted in the 1990s that antibiotic drug discovery was unlike other therapeutic areas the company worked in. There was not a simple relationship between the inhibition of the isolated biochemical target and “the alleviation of the disease state.” They were able to find potent inhibitors of bacterial proteins which could not be used to inhibit activity at the cellular level. GlaxoSmithKline (GSK) have estimated that even in target based screens antibiotic drug discovery success rate “was four- to five-fold lower than for targets from other therapeutic areas”, a fact that made their pursuit of antibacterial drugs financially unviable. Similarly, detailed analysis by AstraZeneca of their screens of hundreds of thousands of compounds revealed hit rates far lower than required to sustain these expensive projects.

Despite a large increase in the size of chemical libraries available and the use of modern automated technologies, high throughput methods on very large compound libraries failed to deliver for antibiotic discovery as it has for other targets (compounds in many pharma chemical libraries are focussed upon entry into mammalian cells and do not possess the properties required for entry into bacteria, especially gram-negative pathogens for which there is a particular dearth of new inhibitors). Now just 6 large pharmaceutical companies remain in the anti-infectives field and only recently have increasing numbers of small and medium sized companies begun developing antimicrobials.
1.2.3.2 Novel Compounds and Novel Targets

Finding a new molecular target for antibiotics and then screening against this seemed like an attractive prospect in the genomic era. Genomics, it was hoped, would allow essential, but previously untargeted, genes to be identified which would lack any cross resistance to current antibiotics. GSK’s analysis of their antibiotic discovery efforts is illustrative of the challenges faced when trying to find new targets for antibiotic treatment:

- They identified targets that were effective in in vivo infection models, but which did not inhibit bacterial growth in vitro.
- High throughput screens for the new target would fail to yield any tractable hits.
- They would find hits but then could not develop these hits into leads as activity could not be increased.
- Of the 67 high throughput screens on different bacteria protein targets, only 5 led to a lead, the majority of which eventually proved fruitless.

Discovery efforts by other companies appeared to face similar issues, severely slowing the rate of antibiotic innovation. The failure of high throughput screens has since been the focus of further work, for example it has been noted that antibacterials seem to have different physicochemical properties to non-antibiotic chemicals, and this may be the source of the challenges in identifying hits. Interestingly, AstraZeneca had fewer issues identifying hits, but still failed to convert these into leads.

Novel targets remain elusive. A potential target must fulfil many criteria. In addition to being lethal when inhibited, any effective new target must be well conserved amongst bacteria to ensure a sufficient spectrum of activity. A few hundred such targets might exist. A target which is essential and non-cumbersome and has multiple individually essential homologues within the cell, will experience a lower rate of development of resistance. It has been suggested that selecting single enzymes as targets during the genomic era led to a more rapid onset of resistance. Penicillin binding proteins (PBPs), the focus of this thesis, make excellent targets of chemotherapy, partly due to the fact so many homologues exist in the cell that can be targeted by the same warhead. As a result, for some bacterial species (such as *Escherichia. coli*), target-mediated resistance is relatively uncommon. In contrast, attempts by GSK to inhibit a single aminoacyl tRNA
synthetase lead to the rapid development of resistance *in vivo* \textsuperscript{77}, which may not have occurred had the compound been able to inhibit multiple tRNA synthetases. Multi-targeting should be high on the list of important criteria.

The majority of current antibiotics for systemic therapy target either the cell wall (Figure 1.2); or DNA, RNA and protein production centres of the bacteria \textsuperscript{74,78}, and were all identified in the pre-genomic era. The challenge of finding new targets led GSK to instead focus on finding new chemical matter to target these well validated centres \textsuperscript{66}, and almost all antibacterial compounds in the current pipeline are focussed towards them too \textsuperscript{36}. Whilst known to be effective in killing bacteria when modulated, an over reliance on these few mechanisms of inhibition may increase cross-resistance and speed the rate of resistance to a new agent.

1.2.3.3 The membrane challenges

Any compound aiming to inhibit an intracellular (or periplasmic) protein must accumulate near the target in sufficient quantities for sufficient time. This requires a balance in favour of the flow of the drug in rather than out of the cell. Drugs can enter the cell via passive diffusion or through porins \textsuperscript{71} and are removed by efflux pumps \textsuperscript{79}. In addition to its on-target affinity, a compound must be optimised with properties that maximise influx whilst avoiding efflux. A particular challenge in the discovery of novel drugs to target gram-negative pathogens is their outer membrane, which makes chemical permeation much more challenging by providing orthogonal filters that exclude potential drugs \textsuperscript{71}. Methods to study the ability of compounds to enter cells are being developed \textsuperscript{80}, but the rules about what chemical features are needed to ensure access have not been elucidated \textsuperscript{71,73,81-84}. Without such rules, phenotypic screening has been used to probe the effect of knocking in porins \textsuperscript{82} and knocking out outer membrane efflux pumps \textsuperscript{83} and porins \textsuperscript{85} to understand their internal logic.

Factors distinguishing between an effective gram-negative agent from an exclusively gram-positive one appeared to be low molecular weight (<600 Da) and higher polarity \textsuperscript{70,73,86}. The need for polarity may be a reason for the lack of success of large library high throughput screens, as these libraries are biased towards lipophilic molecules which are easier to synthesise \textsuperscript{72}. Recently, assays for determining whether a compound was able to accumulate in *Escherichia coli* identified rigidity, globularity, amphiphilicity and the presence of an amine group as
contributing to increased accumulation. The simple addition of amine groups onto known compounds was demonstrated to increase their accumulative properties.

With these complex demands for new molecules, an optimal strategy may be to use high throughput assays to find compounds that can permeate the membrane and add antimicrobial properties to them afterwards. Gram-negative membranes have thus been described as presenting additional drug targets to optimise towards, on top of that of the inhibited protein, essentially doubling the challenge.

1.2.3.4 Clinical Considerations

The clinical demands on antibiotics are also significant. As with all drugs, the compound must be designed with tolerable pharmacokinetic and pharmacodynamic properties which allow it to reach sufficient concentrations at the site of infection for long enough to kill the bacteria, whilst being non-toxic to humans. For certain diseases, this is made even more challenging by particular pathologies, for example the inaccessibility of Pseudomonas aeruginosa in the lungs of cystic fibrosis patients. Unlike other areas of medicine, further consideration must be made of the consequences of the dosing regime selected, as suboptimal dosing can result in increased resistance.

Certain infections, such as those causing sepsis, require such rapid treatment that broad spectrum agents are necessary, whilst for others a narrow spectrum antibiotic coupled to a strain-identifying diagnostic is sufficient, and avoids deleterious effects on gut bacteria.

Any putative antibiotic must then fulfil many scientific requirements whilst being discovered on a budget by the few remaining scientists still dedicated to this area of research. This is a huge challenge.

1.2.3.5 Molecular Mechanisms of Antibiotic Resistance

Resistance can develop by horizontal transfer of existing determinants or de novo mutations that arise by chance. If conditions are correct, mutations of a gene in a single cell can increase its fitness sufficiently to allow the cell line to become the dominant strain. This mutant gene can then be passed onto a plasmid and be transmitted rapidly around the globe, as demonstrated recently in the spread of resistance to the last-resort antibiotic colistin. In this way, a single incidence of
resistance development in one locale can, within a few years, become a global challenge.

Horizontal transfer can occur between different species, and the natural competency of species such as *Neisseria gonorrhoeae* allows them to drive their resistance to "superbug" status. High-level resistance is unlikely to arise from a single point mutation, but instead requires complex changes in cellular biochemistry, which is unlikely to occur in a single patient. Combining resistance elements from many species is often necessary for high level resistance.

On a molecular level, bacteria can become resistant to an antimicrobial through 3 main mechanisms:

I. Bacteria can modify the active compound to reduce its activity. Examples of this mechanism include β-lactamases which hydrolyse the amide of β-lactam antibiotics or the enzymatic adenylation, acetylation or phosphorylation modifications of aminoglycosides.

II. The drug target can be altered to change compound affinity. This can occur as a resistance mechanism against penicillin binding proteins (PBP), the targets of β-lactam antibiotics (see below), vancomycin, trimethoprim as well as rifampicin.

III. Bacteria can use efflux pumps and changes to membrane permeation to prevent entry or prevent build up of inhibitory compounds. Penetration into gram-negative bacteria which have two lipidic membranes with orthogonal "sieving properties" is particularly challenging, and it is these organisms which top the World Health Organisation’s priority pathogens list.

The first two mechanisms are mostly specific to a single class of antibiotics: for example, β-lactamase induction may be effective against penicillin but will not affect the efficacy of vancomycin. In contrast, the third mechanism has been recognised to be a significant source of cross resistance as the induced efflux pumps can have broad specificity to many drugs. Such cross resistance is concerning as it leads to organisms resistant to many families of antibiotics.
1.2.3.6 Methods to Slow Resistance

Many mechanisms of resistance will be associated with a fitness cost \(^{114}\). If the fitness cost presents too high a burden for the organism under conditions of low selective pressure, reversion to the wildtype (non-resistant) may occur, alternatively, compensatory mechanism by further mutation may lower the burden \(^{115-117}\). If reversion is possible, it has been suggested to use cycling of antibiotics to periodically revert the population and keep resistance low, although more evidence for the effectiveness of this is required \(^{118,119}\).

Another method suggested to slow the development of resistance is to use more than one antibiotic concurrently \(^{120}\). This can reduce mortality rates compared to a monotherapy \(^{121}\) and reduces the probability of emergent resistance phenotypes. Certain combinations of drugs have synergistic interactions, making combinations more effective than the sum of their individual effects \(^{122}\). However, many possible combinations of drugs and their relative doses exist and high throughput methods are required to determine the optimal strategy\(^{123}\). Whilst methods like this will likely be of benefit to the slowing of resistance, new molecular entities are required.

1.3 Approaches to finding new antibiotics

1.3.1 Whole cell vs Target-based methods

As mentioned above (section 1.1.1), many of the classes of drugs we use today were found by screening soil samples for molecules with antimicrobial activity. Generally, modern attempts to revisit these methods often only end up finding the same compounds \(^{124}\). An exception to this trend was found by screening uncultured species leading to the discovery of a novel, gram-positive active compound: teixobactin \(^{125,126}\).

Whole cell screening may identify compounds with antimicrobial activity but without providing insight into the mechanism of action. Additionally, without having control over the target selected, either non-specific (and potentially toxic) hits or the same, known biological targets may end up being continually re-selected \(^{65}\). Finding the mechanism of action is a significant hurdle, especially if a screen identifies thousands of compounds with distinct activities \(^{66,127}\). New methods are
being developed that can accelerate this process \cite{127,128}, or the whole cell screen itself can be designed to detect inhibitors for a particular target of interest \cite{65,124,129,130}. An additional advantage of whole-cell screening is that (if the correct bacterial strains are chosen) any hits will \textit{a priori} have the necessary properties to permeate the cell, removing the need to add this property after the fact (which may be necessary for target-based screening), and which can be very challenging.

Alternatively, a purified enzyme (or an entire substrate pathway \cite{65}) can be the target of a screen (as was the case for many of the screens attempted by GSK for example\cite{66}). If the target has previously been well characterised and found to be amenable to inhibition, then this method has the advantage of not needing the challenging mechanisms of action studies of a whole cell approach. In addition to the challenges discussed above (section 1.2.3.2), ignoring more complex aspects of the protein’s role in the cell (e.g. the formation of protein complexes) or simplifications made to assay the target’s activity (e.g. artificial substrates) may lead to a mismatch between the compound’s on-target affinity and its cellular activity. On the other hand, subsequent cellular assays may demonstrate a compound has a broader inhibitory profile than anticipated, which would require mechanism of action studies to investigate.

\textbf{1.3.2 Library Selection}

Once the screening method has been devised, a library of chemicals needs to be found to screen against. As pharmaceutical companies and their large libraries were broadly unsuccessful \cite{65}, other methods to provide chemical diversity are needed. Natural product libraries represent one such pool of molecules and have been used to identify leads \cite{124}, as they were used in the 20th century. Others have noted that high throughput screening libraries are enriched in achiral and apolar molecules, properties unlike natural products. Their lack of these properties is a potential reason for the failure of high throughput screens \cite{131}.

Diversity Oriented Synthesis (DOS) can be used to generate molecules more like natural products \cite{132}, where properties of structural complexity and diversity of a library are the objectives as opposed to a specific chemical pharmacophore. However, even using diversity methods, very few of the possible arrangements of atoms (the “chemical space”) have been synthesised. The number of organic
molecules with up to 17 heavy atoms has been enumerated at 1.6 x10^{10}, but only 6 x10^6 compounds (including those with >17 heavy atoms) have reportedly been synthesised \(^{133}\). If, as some have suggested, there exists a small molecule that can inhibit each druggable protein target \(^{134}\), we need methods to more completely search chemical space and find that lead. Fragment-based drug discovery offers a solution to this challenge \(^{135}\).

1.3.3 Fragment Based Drug Discovery (FBDD)

The use of smaller chemical molecules: “fragments” allows a larger chemical space to be covered with fewer experiments because the chemical space is smaller at lower molecular weights \(^{133}\). Fragments are small, partially drug-like compounds, which have few functional groups but conform to the rule of three (similar to Lipinski's famous rule of 5 \(^{136}\): <300 Da, three or fewer hydrogen bond donors and acceptors, cLogP <3 \(^{137}\). Simpler molecules make better starting points for lead optimisation and may have a larger probability of binding to a target as they lack non-binding groups that reduce the overall affinity \(^{138}\).

Conversely however, if a hit is found its affinity is likely to be lower affinity as it has few opportunities to make interactions with the protein. This typically requires assays capable of detecting low affinity for protein-drug interactions. One method is to use X-Ray crystallography, which allows for very high concentrations of ligands to be used and non-specific interactions to be filtered out as the binding site can be seen in the model. In order to be used for screening many of the stages of typical X-ray crystallography required optimisation to increase their throughput. The XChem technology, developed at Diamond Light Source, has done this \(^{134,139}\). Proteins are crystallised then soaked with fragments from a series of libraries before high throughput (400 crystal per run) crystallography is used to screen the crystals for any binding fragments. In order to identify low occupancy fragments, “PanDDA” software is used to find a background for the unligated structure and then this is compared to the new structure to identify any differences in electron density \(^{140}\).

In a recent example, via collaboration with the University of Cambridge we screened a DOS library of ~1,300 compounds against PBP3 crystals and identified a single hit. We then used the crystal structure to identify vectors in which the fragment could be “grown” to access further regions of the active site, although we were not able to
determine detectable inhibition in assays \textsuperscript{141}. Where two or more fragments are found within the active site, they can be “linked” or “merged” to increase in their potency \textsuperscript{135}.

FBDD has become increasingly popular since its initial description \textsuperscript{142,143}, against antimicrobial targets \textsuperscript{144,145}. This thesis describes an attempt to use FBDD against PBPs, which resulted in the identification of benzoxaboroles as PBP inhibitors with a novel binding mode (Chapter 5).

In actuality, a balance between screening specific targets and whole cell screening to find previously undiscovered modes of action which can then be further elucidated by mode of action studies has been recognised as the best approach \textsuperscript{32}. This work also covers the design of a high throughput screen for whole cell screening at the University of Warwick.

\section*{1.4 PBPs and β-lactams}

PBPs, particularly PBP3, are the focus of the target-based methods described in this thesis. As the name suggests, PBPs are the target of penicillin, and all other members of the β-lactam class of antibiotics (section 1.4.3). Since they were found to be the targets of this important drug class \textsuperscript{34,35}, they have been relatively well characterised \textsuperscript{146} and well validated as a drug target. The β-lactams, through bacterial evolution and rational medicinal chemical design satisfy many of the requirements for antimicrobial drugs described above, and are the most frequently prescribed antibiotic in primary care \textsuperscript{147}. This section will describe the structure and function of PBPs and their interactions with β-lactams.

\subsection*{1.4.1 Functional Role}

The bacterial cell wall is a component of the bacterial envelope, which bacteria use to protect and isolate themselves against the external environment \textsuperscript{148}. The cell wall must be constantly maintained to withstand turgor pressure upon the cell, to determine cell shape and accommodate cell division. One component of the cell wall; peptidoglycan, is a strong, elastic and adjustable biopolymer which wraps around the cell \textsuperscript{149}. The chemistry of the peptidoglycan is broadly similar in both gram-positive and -negative bacteria although it is thicker in the former \textsuperscript{149}.
Gram-negative and gram-positive bacteria both possess a number of PBPs which are used to produce and maintain the bacterial cell wall \textsuperscript{146,150}. \textit{E. coli} has twelve PBPs, but only four are essential to the cell and lethal when knocked out \textsuperscript{151}. Their names follow a convention from their descending order of molecular weights when run on an SDS polyacrylamide gel \textsuperscript{152}, and are typically classified as either high molecular weight (HMM) or low molecular weight (LMM). HMM PBPs are responsible for catalysing glycosyltransferase (GT) and transpeptidation (TP) reactions needed for the polymerisation of peptidoglycan, whereas LMM PBPs have roles in peptidoglycan recycling and maturation \textsuperscript{146,153,154}. The PBPs can also be separated by their functions: class A PBPs have both GT and TP activity, whereas class B PBPs have only TP activity and lack a transglycosylase domain \textsuperscript{146,153}. Penicillin binding proteins have different roles within the cell. For example in \textit{E. coli}: cellular elongation is driven by the mutually redundant PBP1a and PBP1b as well as PBP2, while PBP3 is responsible for cell division at the septum \textsuperscript{146,152}. Inhibition of the functional activity carried out by these enzymes is lethal to the cell.

Peptidoglycan is composed of a long glycan strand, made of alternating N-acetylglucosamine (GlcNAc) and N-acetylmuramic acid (MurNAc) amino sugars, interlinked by a short stem peptide chains (Figure 1.2a). It is synthesised from its constituent sugars in the cytoplasm by the sequential reactions of the “Mur” enzymes \textsuperscript{155,156}. MurA-F append the 5-amino acid chain to a uridine diphosphate-MurNAc (UDP-MurNAc) sugar, the product of which MraY then appends to a lipid tail (undecaprenyl phosphate), with the concomitant loss of uridine monophosphate (UMP). MurG catalyses the glucoaminylation reaction of this molecule (“lipid I”) to UDP-GlcNAc to form “lipid II” \textsuperscript{157}. Lipid II is “flipped” into the periplasm where transglycosylases polymerise it \textsuperscript{146} (Figure 1.2a).

Certain bifunctional PBPs, such as PBP1b from \textit{E. coli} have the transglycosylase activity, but non-PBP transglycosylases also contribute to this activity \textsuperscript{155,158–160}. Transpeptidation is then used to cross-link the peptides of adjacent glycan stands. Whilst all PBPs have a \textit{DD}-peptidase domain they may not all perform transpeptidation reactions, with others performing endopeptidase and carboxypeptidase roles \textsuperscript{146}. In almost all bacteria, the 4th and 5th amino acids (distal to the glycan chain) of the stem peptide are both \textit{d}-alanine residues \textsuperscript{161}. The reaction occurring between the transpeptidation (TP) domain of PBPs and these terminal residues is shown in Figure 1.2b. The stem peptide (by convention, referred to as the “donor”) enters the active site, forming a non-covalent complex, which is
attacked by the active site nucleophilic serine to form an acyl-enzyme complex concomitant with the release of the terminal d-Ala. A subsequent attack by a second nucleophile (by convention this is the “acceptor”) de-acylates the enzyme. In transpeptidation reactions of gram-negative organisms, the acceptor nucleophile is the ε-amine of the meso-diaminopimelyl (DAP) residue (the 3rd amino acid in the stem peptide), but in a carboxypeptidation reaction, this role is performed by water. This chemistry is also exploited by β-lactams which, as Tipper and Strominger noted, share structural similarity with the d-Ala-d-Ala dipeptide (highlighted in orange, Figure 1.2b). β-lactams are recognised by this structural motif, allowing them to be acylated onto the PBP, but once acylated, their rate of de-acylation is significantly slower than for the natural substrate so the β-lactam remains acylated to the PBP, resulting in a suicidal, steric inhibition. It is likely that the β-lactam’s substrate-mimicking mechanism of action and structure has contributed to their success and conservation through evolution. Serine β-lactamases exploit their structural similarities with PBPs to bind β-lactams, but are optimised for much faster de-acylation, leading to the chemical destruction of the β-lactam and resistance to β-lactams for the bacteria.
Figure 1.2. Functional role of the PBPs within the gram-negative periplasm.
(a) Peptidoglycan synthesis machinery is complex and involves many enzymes. Cytoplasmically, the UDP-MurNAc is synthesised and the amino acids of the peptide pentamer are sequentially added. This molecule is then appended to a lipid tail (undecaprenyl phosphate) with the concomitant release of UMP to make lipid I and then glucominylated with UDP-GlcNAc to produce lipid II. Lipid II is "flipped" into the periplasm where it is polymerised by the PBPs which catalyse transglycosylation and transpeptidation reactions. Figure reproduced, with permission from K. Smart. (b) Outline chemical mechanism of the transpeptidation domain. The peptide bond is acylated by nucleophilic attack from the active site sertine, passing through a tetrahedral transition state to reach an acyl-enzyme complex. A second nucleophilic attack (by an amine or water molecule) de-acylates the acyl-enzyme complex, releasing the protein. The figure shows these reactions occurring with the terminal d-Ala-d-Ala of the stem peptide or with \( \beta \)-lactams (a generic penicillin in this example). The structural similarity between the two species is highlighted in orange.
1.4.2 PBP3

1.4.2.1 Functional Role of PBP3

Partial inhibition of PBP3 has long been known to cause an elongated *E. coli* phenotype \(^{152}\), rationalised as PBP3 having a role in promoting peptidoglycan transpeptidation at the dividing cell septum. PBP3 interacts with other proteins as part of the “divisome complex”. Work using filamenting temperature-sensitive (*fts*) mutants of *E. coli* \(^{165}\) showed a sequential assembly of proteins at the site of division, starting with FtsZ which forms a cytoskeletal Z-ring. The *Fts* genes are all found in the “division of the cell wall” (*dcw*) gene cluster, which is conserved, in order, amongst many bacterial species \(^{166}\). One of the gene products to complex later in the process is FtsW, a transmembrane, non-PBP transglycosylase that itself recruits PBP3 (gene name: *ftsI*) into the divisome \(^{159,167}\). The binding of PBP3 to *P. aeruginosa* FtsW was shown to lead to the activation of the glycan polymerase activity of FtsW, which presumably supplies the PBP3 with the feedstocks for transpeptidation \(^{159}\). *In vitro* natural substrate processing by PBP3 has yet to be demonstrated, although thioester substrate-mimics can be turned over by PBP3 \(^{168}\). PBP3 of *P. aeruginosa* is the only PBP of that species which is genetically essential \(^{169}\), making it an attractive target for inhibition. Confusingly, *P. aeruginosa* also expresses a class B PBP called PBP3a (or PBP3x), but this appears to be non-essential \(^{170}\).

1.4.2.2 Topology

As monofunctional transpeptidases (class B PBPs), the PBP3 of gram-negative organisms such as *E. coli*, *P. aeruginosa*, *Acinetobacter baumanii*, and *H. influenzae* as well as the PBP2 of *N. gonorrhoeae* share significant structural homology \(^{105}\), despite low sequence identity (e.g. *E. coli* and *P. aeruginosa* share just 45.1% identity \(^{170}\)) (Figure 1.3). These proteins have three domains, an inner membrane spanning domain, a non-penicillin binding N-terminal domain and the penicillin binding transpeptidase domain \(^{154,171,172}\) (Figure 1.4b). The transmembrane helix spans the membrane once \(^{173}\) and is itself capable of directing the protein (or even GFP) to the septum \(^{174}\). The role of the N terminal domain has not been fully described but appears to mediate PBP3 interactions with other divisome proteins \(^{175,176}\). Interestingly, certain interactions with divisome proteins have been shown to sensitise PBP3 to acylation by cephalixin (but not other β-lactams) \(^{177}\), raising the possibility of allosteric regulation via the N-terminal domain.
The PBP3 C-terminal penicillin binding domain shares a common fold with many PBPs and β-lactamases in the penicilloyl-serine transferase superfamily. Members of the family align structurally and have conserved active site sequence motifs of SXXK, SXN, K(S/T)G \(^{153}\) and a consistent protein fold \(^{178,179}\) (Figure 1.4). PBP3 retains β-lactam binding activity if truncated to the TP domain only \(^{105,180}\), indicating this domain alone is sufficient for catalytic activity. Some proteins such as \(P.\ aeruginosa\) PBP3 do not tolerate this truncation \(^{179}\) (D. Bellini unpublished observations). Crystal structures have been produced of only transpeptidation domains \(^{105,180}\) or more commonly with the transpeptidation and N-terminal domains but lacking the transmembrane domain which is removed to allow for crystallisation and aqueous solubility \(^{105,173,179,181–183}\).

1.4.2.3 Crystallography of the Transpeptidase Domain

Crystal structures of the class B PBP transpeptidase domain reveal a highly ordered core with a wide, shallow central cleft, which has the active site serine (e.g. in \(P.\ aeruginosa\) PBP3 Ser294) at its centre \(^{105,178,179,181,182,184}\) (Figure 1.4). The fold of the transpeptidase domain is shared by all PBPs and β-lactamases of the penicilloyl-serine transferase superfamily \(^{185}\). The core of the domain is formed by a 5 stranded β-sheet (β1a and 1b and β3-5), with β3 forming the floor of the active site cleft (Figure 1.4c and d). The "jaws" of the active site cleft are formed by α11 on the bottom and the short and consecutive β2a-d and α4 elements on the top (Figure 1.4b). The catalytic residues are stabilised by their location near the centre of the domain, on α6 (the active site serine and nearby lysine in the SXXK motif), α4 (the serine of SXN) or β3 (lysine of the K(S/T)G motif). Mobile loops extend between the secondary structure elements and are important for binding. Notable are the β5-α11 loop which moves up and towards the active site upon piperacillin binding \(^{179}\) and the β3-β4 loop which is poorly resolved in some structures \(^{105}\) (Chapter 4).
Figure 1.3. Sequence alignments of five clinically important class PBPs from gram-negative pathogens. Sequences of the transpeptidase domain were aligned with Clustal Omega\(^{166}\) then the figure was generated by ESPript \(^{187}\). Numbering refers to residues of \( P. \) aeruginosa PBP3. Red arrows below the sequence show locations of important motifs (SXXK, SXN and K(T/S)G).
1.4.3 β-lactams

The β-lactams are a family of drugs containing a 4-membered (β) 2-azetidinone (lactam) ring. They are some of the most important antibiotics being "well tolerated, efficacious and well prescribed" 103. The first example of each class were found from natural product screens: penicillin was identified from fungal extracts 19 as were cephalosporins 188–190, whilst carbapenems and monobactams 191,192, were extracted from bacterial isolates. Many derivatives of each class have been synthesised and are used in the clinic 193, although just one example of a monobactam, aztreonam, has been marketed. Since the release of aztreonam 191,194 in the 1980’s no further classes have been discovered although many iterations of drugs within these classes have been developed. The most recent to be released is Cefiderocol which features an iron-conjugating catechol group to drive its uptake into gram-negative bacteria 195,196. Other innovations have been the development of various combinations of a β-lactam and β-lactamase inhibitor (see below) 103. Given the wide arsenal of β-lactams available (Scheme 1.1), clinical decisions on the choice of β-lactam administered depend on the pathogenic agent, route of administration, and the risk of allergic reaction 193,197,198.
Scheme 1.1 β-lactams and β-lactamase inhibitors. (a) Example structures of four common classes of clinical β-lactam drugs. The characteristic core group for each of the classes is coloured. The R1 and R2 side chains are labelled on the structure of ceftazidime, and the numbering system of the penicillin core group is indicated. The C-3 carboxylate is the carboxylic acid at the 3-position on penicillin G. (b) Selected example structures of β-lactamase inhibitors. Note the presence of a β-lactam ring in clavulanic acid.
1.4.3.1 Kinetics

When β-lactams react with PBPs they form covalent acyl-enzyme complexes with the active site serine (Figure 1.2). The steps of this reaction are summarised in Scheme 1.2. An initial non-covalent Michaelis complex \((E \cdot I)\) is in rapid equilibrium with the free enzyme \((E)\). The PBP and β-lactam composing \(E \cdot I\) react to give the covalently acylated \(EI\). This acylation step is typically considered to be irreversible. The acyl-enzyme complex can be hydrolysed to give the product \((P\), a ring-opened penicilloic or cephalosporic acid) and free enzyme. This de-acylation is typically slow (< 0.02 min\(^{-1}\)), thus the inhibition of a PBP is primarily sensitive to the ratio of the rate of covalent bond formation and the non-covalent dissociation constant. Chromogenic nitrocefin appears to act differently to other β-lactams, with a much higher turnover rate (up to ~ 40 min\(^{-1}\)) in, a phenomenon which is discussed in more detail in Chapter 6.

The non-covalent components of the interaction are strong enough to allow several non-acylated β-lactams to be observed in crystals in the PBP3 active site. The cephalosporic acid of cefoperazone (but not the penicilloic acid of azlocillin) can stabilise the protein to thermal melting. However, thermodynamic analysis of \(E. coli\) PBP5 found that the noncovalent interactions of four different β-lactams did not correlate with the experimentally determined acylation rates (which includes both non-covalent interaction and the covalent bond formation). This indicates that the non-covalent component is the less important component.

![Scheme 1.2 Reaction pathway of β-lactams interacting with PBPs](image)

\(E + I \rightleftharpoons E \cdot I\) (Pre-acyl Complex)
\(E \cdot I \rightarrow EI\) (Acyl-enzyme Complex)
\(EI \rightarrow E + P\) (De-acylation)

\(E\) represents the enzyme, \(I\) the β-lactam, \(E \cdot I\) is a non-covalent complex, \(EI\) is an acyl-enzyme covalent complex, \(P\) is the penicilloic acid (or equivalent) product of ester hydrolysis.
1.4.3.2 Crystallographic Interactions with β-lactams

Several important regions in the cleft contribute to β-lactam binding. Crystal structures of *P. aeruginosa* PBP3 (PaPBP3) are studied throughout this work and Figure 1.4e shows the principal residues of this protein that interact with piperacillin. The active site serine (Ser294) reacts with the β-lactam bond to form an acyl-enzyme complex. Nearby, and forming an interacting hydrogen bonding network are Ser349, Lys484 and Lys297 (blue in Figure 1.4e), which are thought to activate the Ser294 for nucleophilic attack \(^{205-210}\). Positioned on the β3 adjacent to Lys484 are Ser495 and Thr497 (residues in red in Figure 1.4e). These residues form an “acid binding pocket” that hydrogen bonds with the C3 carboxylate group (or equivalent in monobactams or cephalosporins) found in all β-lactams. Backbone amines of Ser294 (active site serine) and Thr487 (on the β3 strand) hydrogen bond with the β-lactam carbonyl, creating an oxyanion hole which polarises the carbonyl, activating it during its initial acylation, and later de-acylation \(^{185}\). Gly486 (glycine of the K(S/T)G motif - directly preceding Thr487), is apparently the key determinant of positioning of the amine of Thr487 as this residue is conserved in all members of the penicilloyl-serine transferase superfamily\(^{211}\).

A “hydrophobic wall” \(^{179}\) of residues Tyr503, Tyr532 and Phe533 can form adjacent to the β-lactam phenyl group as shown in Figure 1.4e. The R1 side chain of the β-lactam is engaged by a channel formed of tyrosines (Tyr328, Tyr407, Tyr409 and Tyr498) and the β3 strand (Arg489 side chain and several β3 backbone amides). Asn351 is a highly conserved residue \(^{153}\) which interacts with the first amide carbonyl of the β-lactam (this is another frequent motif of β-lactams). Val333 forms hydrophobic interactions with the β-lactam and the hydrophobic wall, creating the column seen spanning the active site cleft in Figure 1.4d.

The above description of interactions is generally true for simple penicillins \(^{105,179,183,212}\), aztreonam \(^{179}\), cefoperazone \(^{183}\), ceftazidime \(^{181}\) and even β-lactams without a covalent link to the protein \(^{183,203}\). For meropenem \(^{179}\), β-lactams with large siderophore conjugates \(^{179,213-216}\) and ceftobiprole (a cephalosporin with a large R2 sidechain \(^ {217}\)) many of the same regions participate but accommodation is made to fit the different groups.
Figure 1.4. Structural views of class B PBPs. (a) Overall topology of a class B PBP (PaPBP) shown in cartoon representation. The first domain is a short, single spanning transmembrane region including a short cytoplasmic sequence. This domain has never been crystalised and is sketched only. The crystal structure of the non-penicillin binding N-terminal domain is shown in pink and the functional transpeptidase domain in cyan with the active site serine highlighted in red. (b) Secondary structure elements of the penicillin binding domain of *P. aeruginosa* PBP3 (PDB code 6HZR 103) following the convention from 178. Rendered in cartoon representation with colours in rainbow from dark blue at the N-terminal to red at the C-terminal. Sequences for these secondary structure features are shown in
Figure 1.3. Despite the high resolution of this structure, the β3-β4 loop is poorly resolved, likely due to its flexibility. A dotted arrow indicates the missing region. Secondary structure elements are labelled following convention 178. (c) Sequence conservation of the transpeptidase domain. The same colours (red for highly conserved, blue for partially conserved and white for non-conserved) are shown for each residue on the 3D crystal structure as used in Figure 1.3. The highly conserved (across the entire penicilloyl-serine transferase superfamily) active site motifs are labeled. (d) Surface representation of the *P. aeruginosa* PBP3 reacted with piperacillin (PDB: 6R3X 105), coloured according to degree of hydrophobicity 218,219 shows the “jaws” around the shallow cleft of the active site. A column formed by the hydrophobic interactions of Val333 and Phe533 that spans the cleft. (e) The active site of the piperacillin-acylated PBP3 shown as a transparent surface. The β3 strand which runs parallel to the β-lactam along the floor of the active site is shown in cartoon representation. Residues interacting with the piperacillin are shown in stick representation and coloured according to binding region: catalytic residues (blue); acid binding pocket (red); hydrophobic wall (orange); R1 binding pocket (green); residues on the roof of the cleft (purple). Hydrogen bonds between piperacillin and residue side chains are shown (dashed lines).

1.4.3.3 Physiological Effect of β-lactam Inhibition of PBPs

Early microscopic observations noted that *E. coli* when treated with penicillin transformed from their typical rod shapes into protoplasts, which lyse under unfavourable osmotic conditions 220,221. If osmosis was prevented by isotonic conditions and the penicillin was washed off, growth could recover 221. Similarly if the *E. coli* were treated with penicillin in media that did not support growth, the drug treatment did not lead to lysis 220. These results indicate that penicillin (and β-lactams more generally) only acts indirectly through its inhibition of the cell wall synthesis. Later work showed that hydrolases play an important role in breaking down the peptidoglycan, particularly during cell division, and the disruption of the balance between production and destruction of the macromolecule at the cell septum leads to cell lysis 222. Pneumococcal cells without autolytic hydrolases were insensitive to penicillin 223,224. More recent evidence suggests that β-lactams may induce futile cycles of glycan strand production, which hastens cell death 225,226.

β-lactams each have individual specificities for the various PBPs of a cell 227–230 and it has been shown, using β-lactams with single PBP affinity, that the inhibition of more than one PBP in *E. coli* causes rapid bactericidal effects where the inhibition of just one causes only bacteriostasis 231. Thus, the relationship between target engagement and physiological effect is inexact. Additionally, each β-lactam has different membrane permeation properties and susceptibility to β-lactamase attack, further complicating this relationship 232. Ultimately, the broad spectrum antimicrobial activity (both gram negative and gram positive) of the β-lactams as a group derives
from their diverse on-target profile, which in turn derives from their mechanistic similarity to the PBP substrate. Consequently, a β-lactam can be found with activity against most organisms.

1.4.4 Resistance to β-lactams

Resistance of bacteria to β-lactams has been observed to occur by all three of the resistance mechanisms described above (see “Mechanisms of Antibiotic Resistance”) 233.

1.4.1 Penetration-efficiency-mediated Resistance to β-lactams

Resistance to β-lactams in E. coli can occur through mutations to the OmpC and OmpF porin proteins, which are a major entry route of β-lactams into this species 113,234. In P. aeruginosa this mechanism is less common and instead changes in the composition of the lipopolysaccharide, a layer of polymers beyond the outer membrane has been shown to influence the ability of β-lactams to enter the cell 235,236. This layer contributes to the “intrinsic resistance” of P. aeruginosa, along with expression of efflux pumps 113,237. Efflux pumps alone can lead to high levels of resistance in certain species such as Klebsiella pneumoniae 238.

1.4.2 Target-mediated Resistance to β-lactams

Target-mediated Resistance to β-lactams is discussed in Chapter 4. Broadly, this route appears to vary in its use significantly between species. In certain gram-positive species (e.g. methicillin resistant Staphylococcus aureus) acquisition and expression of a gene (e.g. mecA) encoding new, low affinity PBPs occurs 239, but this has not been observed for gram-negative class B PBPs. N. gonorrhoeae and H. influenzae are frequently observed to reduce the β-lactam affinity of target PBPs by generating variants in the PBP sequence. However, the exact mechanisms that lead to reduced affinity are not yet deciphered 105.

1.4.3 β-lactamase-mediated Resistance to β-lactams

Perhaps the best studied resistance mechanism is β-lactamase expression. Resistance through β-lactamase expression was observed even before the widespread clinical use of β-lactams 240,241 and evidence suggests they have existed for millions of years 242,243, likely co-evolving alongside β-lactams 193, in a bacterial “arms race”. There exists a β-lactamase that can hydrolyse each class of β-lactam
(although not all β-lactamases can hydrolyse all β-lactams) and β-lactamases are continually evolving, increasing their activity and broadening their substrate specificity. New β-lactams experience β-lactamase-mediated resistance within a few years of their release. The impact of β-lactamases is not limited to a single cell and they can serve as communal defense mechanisms, protecting susceptible cells in liquid media and in biofilms.

The Ambler classification separates β-lactamases on the basis of their sequence into 4 groups A, B, C and D. A, C and D are all serine β-lactamases, with acylation and de-acylation mechanisms similar to PBP. Class B enzymes are metallo-β-lactamases, for which there are no approved inhibitors (although aztreonam is not effectively turned over by this class, providing a therapeutic option). An alternative classification method, the Bush-Jacoby-Medeiros classification scheme separates β-lactamases by their function, which is the most important consideration for clinicians encountering a β-lactamase in a medical context. In this scheme group 1 contains mostly chromosomally encoded, Ambler class C cephalosporinases; group 2 (the largest group, with 12 subgroups) contains penicillinases, cephalosporinases and carbapenemases from Ambler class A and D; and group 3 is composed of metallo-β-lactamases (Ambler class B), which cannot be inhibited by clavulanic acid and tazobactam and generally have poor affinity for monobactams. Serine β-lactamases appear to have evolved from a primordial PBP, obtaining mutations that maintain β-lactam recognition capabilities and their overall fold, but which give them improved catalytic properties and decreased affinity of the PBP substrate. Genetic analysis indicates that the different classes of β-lactamase diverged from distinct ancestors, and each β-lactamase is more closely related to a PBP than to another β-lactamase. The core fold of the transpeptidase domain is thus able to support a wide array of different enzymatic functions each tuned to the enzyme’s function.

Attempts have been made to counter β-lactamase resistance by using a number of different β-lactamase inhibitors to protect a co-administered β-lactam. Three examples of β-lactamase inhibitors are shown in Scheme 1.1b. The earliest of these inhibitors, clavulanic acid was discovered from natural product screens. The β-lactam bond undergoes the typical serine acylation reaction with class A β-lactamases (Figure 1.2), however unlike other β-lactams, this releases a reactive intermediate that forms an irreversible complex with the β-lactamase, leading to suicidal inhibition. Like most β-lactamase inhibitors, clavulanic acid lacks
antimicrobial activity and it is administered alongside a β-lactam (often amoxicillin). More recent β-lactamase inhibitors classes include the diazabicyclooctanes, typified by avibactam, a reversible inhibitor of all serine β-lactamases, and boron-based inhibitors such as vaborbactam (class A and C activity). Whilst β-lactamase inhibitor/β-lactam combinations have been able to restore the efficacy of β-lactams to β-lactamase expressing strains, their efficacy depends on the ability of both components to reach the site of infection at sufficient concentrations. A single agent, β-lactamase-resistant therapy would not have this limitation.

1.4.5 Non-β-lactam compounds

PBPs are an excellent drug target. Many attempts have been made to find non-β-lactam PBP small molecule inhibitors (reviewed in 266–268), with the hope of inducing the same physiological effect as β-lactams but without the liability to β-lactamase attack. These are summarised in Table 1.1. These can be broadly divided into 3 groups. (i) Those that act as suicidal covalent inhibitors of PBPs (similarly to β-lactams), (ii) those with non-covalent mechanisms of action and (iii) those that acylate the active site with a boron warhead (which forms a reversible covalent interaction). Additionally, cyclic peptides have previously been investigated for PBP and β-lactamase inhibition and are currently being investigated by Bicycle Therapeutics for PBP inhibition. Unfortunately, in many cases even if a warhead has been demonstrated to engage PBPs, this can be insufficient to cause bacterial growth inhibition: this may be due to low potency, inability to reach the target, chemical instability within the cell or a combination of these factors.

Whilst a large number of pharmacophores have been investigated as PBP inhibitors, none have reached clinical medicine. The diazabicyclooctane (DBO) avibactam and the boronate vaborbactam are used as β-lactamase inhibitors and the bicyclic boronate taniborbacatam (formerly VNRX-7145) is in Phase III clinical trials for combination with cefepime. New boron-based β-lactamase inhibitor QPX7728 has recently entered Phase I clinical trials. The DBO literature offers hope that successful β-lactamase inhibitors may one day find use against PBPs. Avibactam has been shown to bind PBPs and ongoing work may prove DBOs to be an effective PBP-targeting therapeutic. Crystal structures show that DBOs including zidebactam, avibactam, ETX0462 and WCK 4234 can bind PBP2 and PBP3 and that there is an "enhancer effect" on the activity of the
\(\beta\)-lactams activity PBPs \(^{275,277,278}\) when co-administered with DBOs such as nacubactam and zidebactam. Additionally, the DBOs have efficacy via direct inhibition of PBPs and \(\beta\)-lactamases.

The diversity of non-\(\beta\)-lactam covalent warheads that have been investigated indicate the unique properties of the \(\beta\)-lactam, benefiting from substrate analogy, active site affinity and correct chemistry of the warhead. The \(\beta\)-lactam bond is one of a number of privileged warheads which have reactivity towards serine, (like boronates) \(^{282}\). The amide carbonyl of the \(\beta\)-lactam is destabilised by the ring strain of the 4 membered ring, making nucleophilic attack by the alcohol more likely \(^{283}\). The activation energy for this reaction must be low enough for the attack to be possible by serine but avoid rapid spontaneous hydrolysis \(^{214}\). The energy barrier is tuned by the side chain chemistry and the type of core found in the molecule \(^{284}\). As has been noted with the novel lactivicin core \(^{214}\), any new serine-binding warhead must carefully compromise between reactivity to serine attack and hydrolysis.

Lactivicin and \(\gamma\)-lactam derivatives in particular were the subject of large pharmaceutical investigations, which may eventually yield new medicines \(^{214,266,285-289}\). Adding sidechains from \(\beta\)-lactams to these molecules has been used to improve their affinity and the crystal structures demonstrate that they are using many of the \(\beta\)-lactam-binding regions shown in Figure 1.4.

A number of different compounds apparently demonstrate non-covalent inhibition of PBPs (at least 2 of which are thought to be non-specific promiscuous binders, whose activity is ablated in the presence of detergent \(^{267,268}\)) (Table 1.1). The low \(\mu\)M potency (against \(P.\ aeruginosa\) PBP3) and antimicrobial activity of the pyrrolidine-2,3-diones offers promise \(^{291}\). However with all the non-covalent inhibitors, their mode of action is unclear. They have proved difficult to observe crystallographically \(^{291-293}\), with the only successful example being found in the allosteric site of PBP2a from \(S.\ aureus\), (a site not found in gram-negative class B PBPs \(^{294-297}\)). Similarly, our crystallographic fragment screen of 1,300 diversity-orientated fragments against PBP3 found only a single hit (a \(\gamma\)-lactone) and this was covalently bound \(^{141}\). This is a hit rate several orders of magnitudes lower than other projects using a similar technique. Why PBPs are so difficult to crystallise with non-covalent inhibitors is unclear, and is reminiscent of the paucity of structural information about the protein-natural substrate complex (at least for HMM PBPs). Until this is understood, crystallographic FBDD screens may not be very productive.
The boron-based inhibitors will be discussed in more detail in Chapter 5, but they are distinct in that they are typically considered transition state inhibitors, mimicking the transition state of the β-lactam binding cycle (Figure 1.2). Boron is uniquely able to fill this role as it can exist in both an \( sp^2 \) and \( sp^3 \) hybridisation states.

Table 1.1 demonstrates that there is still much to understand about the behaviours of non-β-lactam inhibitors of PBPs.

<table>
<thead>
<tr>
<th>Chemistry</th>
<th>Active in a live organism</th>
<th>PBP Class investigated</th>
<th>TargetedCOMSITE PBP</th>
<th>Spectral Antibiotic activity</th>
<th>Antibacterial activity</th>
<th>Comments</th>
<th>References</th>
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</thead>
<tbody>
<tr>
<td>Pyrazolidinone γ-lactam</td>
<td>C</td>
<td>H</td>
<td>Y</td>
<td>6VQ</td>
<td>+ve &amp; -ve</td>
<td>S</td>
<td>Many derivatives have been synthesised (see reviews).</td>
</tr>
<tr>
<td>Lactivicin</td>
<td>C</td>
<td>H</td>
<td>Y</td>
<td>4OOL</td>
<td>+ve &amp; -ve</td>
<td>BLI</td>
<td>Many derivatives have been synthesised (see reviews).</td>
</tr>
<tr>
<td>β-lactone</td>
<td>C</td>
<td>H</td>
<td>E</td>
<td>ND</td>
<td>+ve &amp; -ve</td>
<td>S</td>
<td>Found in a natural product screen. Poorly stable in solution. β-lactones have been used for labelling PBPs</td>
</tr>
<tr>
<td>γ-lactone</td>
<td>C</td>
<td>H</td>
<td>Y</td>
<td>6Y6Z</td>
<td>N</td>
<td>ND</td>
<td>Found in an X-ray crystallographic screen</td>
</tr>
<tr>
<td>Cyclobutanone</td>
<td>C</td>
<td>ND</td>
<td>ND</td>
<td>R</td>
<td>N</td>
<td>BLI</td>
<td>In equilibrium with the hemiacetal in solution and in the active site</td>
</tr>
<tr>
<td>β-sultam</td>
<td>C</td>
<td>L</td>
<td>ND</td>
<td>1YQS</td>
<td>ND</td>
<td>BLI</td>
<td>Tested only in R61 from S. Pneumoniae, slow cyclation of the active site is observed</td>
</tr>
<tr>
<td>Triazoline-3-one</td>
<td>C</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>N</td>
<td>ND</td>
<td>Triazoline-3-thiones were also tested, similarly inactive</td>
</tr>
<tr>
<td>Diazabicyclooctane</td>
<td>C</td>
<td>H</td>
<td>Y</td>
<td>6G9S</td>
<td>+ve &amp; -ve</td>
<td>BLI</td>
<td>First investigated as a β-lactamase inhibitor, with later derivatives optimised for PBP inhibition (focussed on E. coli PBP2)</td>
</tr>
<tr>
<td>Compound</td>
<td>Structure</td>
<td>C</td>
<td>ND</td>
<td>N</td>
<td>ND</td>
<td>BLI (w)</td>
<td>Little investigation into this group. Unstable in solution</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-----------</td>
<td>---</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>---------</td>
<td>------------------------------------------------------------</td>
</tr>
<tr>
<td>Activated Alanine</td>
<td>R-CH₂Cl</td>
<td>C</td>
<td>L</td>
<td>N</td>
<td>ND</td>
<td>ND</td>
<td>R is a Boc- and carboxybenzyl-protected lysine</td>
</tr>
<tr>
<td>Trifluoroketone</td>
<td>HOOC-CH₂-</td>
<td>C</td>
<td>L</td>
<td>ND</td>
<td>3ZCZ</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Cyclic Phosphate</td>
<td></td>
<td>C</td>
<td>L</td>
<td>ND</td>
<td>1SCW</td>
<td>ND</td>
<td>BLI</td>
</tr>
<tr>
<td>Phosphonate</td>
<td></td>
<td>C</td>
<td>L</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>BLI</td>
</tr>
<tr>
<td>Anthraquinine</td>
<td></td>
<td>PR</td>
<td>H</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>N</td>
</tr>
<tr>
<td>Arylalkylidene Rhodanine</td>
<td></td>
<td>NC</td>
<td>Y</td>
<td>ND</td>
<td>ND</td>
<td><em>ve</em> &amp; -ve</td>
<td>Shows slow, reversible inhibition. Inhibition of analogues are detergent sensitive - possible promiscuous binders</td>
</tr>
<tr>
<td>Arylalkylidene Iminothiazolidin-4-one</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ortho-phenoxyl diphenylurea</td>
<td></td>
<td>PR</td>
<td>H</td>
<td>A</td>
<td>ND</td>
<td>ND</td>
<td>P</td>
</tr>
<tr>
<td>Naphthalene Sulphonamide</td>
<td></td>
<td>PR</td>
<td>L+H</td>
<td>ND</td>
<td>ND</td>
<td>*ve &amp; -ve</td>
<td>This pharmacophore has been identified in two separate screens</td>
</tr>
<tr>
<td>Benzamidobenzoic acid</td>
<td></td>
<td>PR</td>
<td>L+H</td>
<td>ND</td>
<td>ND</td>
<td>*ve</td>
<td>R</td>
</tr>
<tr>
<td>Compound</td>
<td>PR</td>
<td>L+H</td>
<td>Y</td>
<td>ND</td>
<td>N</td>
<td>ND</td>
<td>S</td>
</tr>
<tr>
<td>---------------------</td>
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<td>---</td>
<td>----</td>
<td>---</td>
<td>----</td>
<td>---------</td>
</tr>
<tr>
<td>4-Quinolone</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>discovered in a computational docking screen against PBP5 (details not given)</td>
</tr>
<tr>
<td>Thiazolidine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>similarity to the core of the Aryalkylidene Imidoothiazolidin-4-ones is noted. Possibility of covalent acylation though amide, but not identified in mass spectrometry experiments, MIC&lt;IC50*</td>
</tr>
<tr>
<td>Linked Phenyls</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>presence of reactive hydroxyl nitrobenzaldehyde group could be a covalent warhead. MIC&lt;IC50*</td>
</tr>
<tr>
<td>Thiochromone</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>* these compounds are a selection of 28 hits identified in a screen of 50,000 compounds against PBP2 using a bocillin fluorescence anisotropy assay. Note that the MICs are significantly lower than the IC50 for PBP2, which may indicate other cellular targets.</td>
</tr>
<tr>
<td>Triazole</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OXADIAZOLE</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>discovered in a computational docking screen against Methicillin-resistant Staphylococcus aureus PBP2a</td>
</tr>
<tr>
<td>Quinazolinone</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>crystal structure shows binding to an allosteric site of Staphylococcus aureus PBP2a, as reported for a β-lactam. This causes structural rearrangement of the active site. The authors propose that the compound’s addition inhibition of PBP1 (which lacks an allosteric site) indicates active site binding is possible as well</td>
</tr>
</tbody>
</table>

* MIC<IC50: Minimum Inhibitory Concentration less than the Inhibition Concentration 50%.
<table>
<thead>
<tr>
<th>Compound</th>
<th>Structure</th>
<th>Binding Mode</th>
<th>Activity</th>
<th>BLI</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrrolidine-2,3-dione</td>
<td><img src="image1" alt="Structure" /></td>
<td>PR H Y</td>
<td>ND /ve</td>
<td>ND</td>
<td>Discovered by screening PaPBP3 with an S2d based assay. Compounds show low µM activity vs P. aeruginosa cells and no cytotoxicity.</td>
</tr>
<tr>
<td>Alkyl boronic acid</td>
<td><img src="image2" alt="Structure" /></td>
<td>C L+H ND 2Y2K</td>
<td>+ve</td>
<td>BLI</td>
<td>Investigated as a natural substrate mimic of tigata. Can form trivalent complexes with PBP3s.</td>
</tr>
<tr>
<td>Aryl boronic acid</td>
<td><img src="image3" alt="Structure" /></td>
<td>C L+H Y</td>
<td>3ZVT N</td>
<td>BLI</td>
<td>Can form trivalent complexes with PBP3s.</td>
</tr>
<tr>
<td>Vaborbactam</td>
<td><img src="image4" alt="Structure" /></td>
<td>C H Y</td>
<td>7AUH N</td>
<td>BLI</td>
<td>The first clinically used boron-based BLI. Crystal structures show remarkable conservation of binding mode with β-lactams.</td>
</tr>
<tr>
<td>Bicyclic Boronate</td>
<td><img src="image5" alt="Structure" /></td>
<td>C L+H</td>
<td>+ve &amp; -ve</td>
<td>BLI</td>
<td>Investigation by Venatorx for use as a PBP and BLI. Very encouraging results, including good MICs.</td>
</tr>
<tr>
<td>Benzoxaborole</td>
<td><img src="image6" alt="Structure" /></td>
<td>C H Y</td>
<td>7AU1 N</td>
<td>BLI</td>
<td>Subject of a large section of my work. Promising new warhead for PBP3s. Investigated but later dropped by AZ as a BLI.</td>
</tr>
</tbody>
</table>

**Table 1.1. Non-β-lactam PBP binding small molecules previously reported.**

References: A: 260,285–287, 214,266,288,289; B: 260,274,276,277, 268, 299,310, K: 299,311, L: 267,312, M: 267,311,313,314, N: 315, O: 267,290, P: 292, Q: 316–318, R: 316,317, S: 319, T: 318, U: 293, V: 294–298,320, W: 291, X: 299,311,312,325, Y: 322,326, Z: 329–331, (Chapter 5), AA: 262,327,328, BB: 252,332, (Chapter 5). Data listed is a summary of the properties of each class and not reflective of the individual molecule shown. Abbreviations: C: Covalent: Warhead designed to engage covalently; PR: Reduced binding of probes (S2d, BOCILLIN FL, nitrocefin, or radiolabelled penicillin to the active site but mechanism not confirmed to be competitive); NC: When probed with S2d, behaves non-competitively. L: Activity demonstrated in a LMM PBP; H: Activity demonstrated in a HMM PBP E: Elongation phenotype was observed microscopically following treatment of bacteria: indicative of PBP3 inhibition 152; R: Crystallography reported, not found in the PDB; A: Crystallography attempted, but could not determine crystal structures with the inhibitor bound; +ve: gram-positive activity; -ve: gram-negative activity; BLI: β-lactamase Inhibitor; S: β-lactamase susceptible (w) weak.
1.4.6 PBP Assays

In order to quantitatively assess the ability of a compound to bind to PBPs, assays are required. The assays reported for investigation of PBPs are listed in Table 1.2 and the structures of the probes in Figure 1.5. Early investigation of PBPs and the binding affinities of various β-lactams were typically carried out with radiolabelled analogues of important β-lactams. When run on a gel, these reagents could give a picture of the affinity of all the PBPs in a membrane preparation for a given β-lactam \(^{152,334}\). Alternatively, the binding of a radiolabeled probe β-lactam to a purified enzyme can be used to find the kinetics of an individual protein:β-lactam interaction, and by extension the kinetics of a competing ligand \(^{315}\).
<table>
<thead>
<tr>
<th>Assay</th>
<th>Protein-based probe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radio-labeled β-lactams</td>
<td>Y</td>
</tr>
<tr>
<td>Nitrocefin and CENTA</td>
<td>Y</td>
</tr>
<tr>
<td>Bocillin FL fluorescence anisotropy</td>
<td>Y</td>
</tr>
<tr>
<td>Bocillin-labelled β-lactams in SDS-PAGE gels</td>
<td>Y</td>
</tr>
<tr>
<td>Biotinylated β-lactams</td>
<td>Y</td>
</tr>
<tr>
<td>S2d thioester substrate</td>
<td>N</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Methodology</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiolabelled analogues of various β-lactams are used to bind to PBPs which are then separated either on a gel and imaged using photographic paper or by filter then measured by scintillation</td>
<td>- Chemically (but not physically) identical analogues of β-lactams can be used, minimising artefacts. - Membrane preparations allow the observation of many PBPs simultaneously. - Can be used in a microplate assay format</td>
<td>- Long exposures times required (days to weeks) to develop signal - Health and safety concerns when handling radioactive substances - End point assay - Generally low throughput</td>
</tr>
<tr>
<td>Reaction of nitrocefin/CENTA with a PBP leads to a colour change that is detected in absorbance spectrophotometry</td>
<td>- Quick and easy assay - Good sensitivity of signal - Continuous assay</td>
<td>- Non-clinical β-lactam, results may not be relevant to all β-lactams - Substrate turnover not observed for all enzymes - Coloured inhibitors can interfere</td>
</tr>
<tr>
<td>Reaction of bocillin FL with a PBP can be measured by a change in the rate of rotation of the fluorophore, as determined by polarisation of fluorescent light</td>
<td>- Quick and easy assay - Good sensitivity of signal - Continuous assay - Most PBPs acylated - Very little protein consumed - Can observe acylation and de-acylation separately - Little interference from coloured compounds</td>
<td>- Presence of large fluorophore may affect normal β-lactam kinetics - Assay interference if aggregation occurs - Indirect substrate-concentration relationship</td>
</tr>
<tr>
<td>Bocillin binds to PBPs which are then run on an SDS-PAGE gel and imaged using a scanning fluorescence imager. Quantification possible by densitometry</td>
<td>- Membrane preparations allow the observation of many PBPs simultaneously - Direct substrate-concentration relationship - Less prone to interference than bocillin anisotropy</td>
<td>- End point assay - Presence of large fluorophore may affect normal β-lactam kinetics - Quantification can be a challenge - Low throughput</td>
</tr>
<tr>
<td>Biotinylated β-lactams bind to PBPs, which are then quantified using a streptavidin-conjugated horseradish peroxidase (HRP) and a fluorescent, oxidised substrate</td>
<td>- Has been developed into a high throughput assay - Good sensitivity of signal</td>
<td>- End point assay - Presence of large fluorophore may affect normal β-lactam kinetics - Quantification can be a challenge</td>
</tr>
<tr>
<td>Thioester substrate can react with the PBP, releasing a free thiol which is detected by an absorbance or fluorescence coupling reagent</td>
<td>- Supposedly mimics the transpeptidase reaction (amino acids can be used as “acceptor substrates” to accelerate de-acylation) - Quick and easy assay - Good sensitivity of signal - Continuous assay - Non-β-lactam probe</td>
<td>- Not all PBPs can process the substrate - some prohibitively slow</td>
</tr>
<tr>
<td>Method</td>
<td>N</td>
<td>Y</td>
</tr>
<tr>
<td>--------------------------------------------</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Mass spectrometry (MS)</td>
<td>N</td>
<td>Y</td>
</tr>
<tr>
<td>Surface Plasmon Resonance</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Thermodynamic analysis by reversible denaturation</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Thermal shift assay</td>
<td>N</td>
<td>Y</td>
</tr>
<tr>
<td>Natural substrate assays</td>
<td>N</td>
<td>N</td>
</tr>
</tbody>
</table>

Table 1.2. Assays used for the identification and classification of Class B PBP binding agents References: A, B, C, D, E, F, G, H, I, J, K, L. Abbreviations: Y: Yes; N: No; Practicality: a qualitative scale from challenging (- -) to simple (++), which aims to encompass the cost of the equipment and reagents for the assay, technical skill required to use it, and time taken to run the assay. For example, mass spectrometry assays score “- -” due to the high cost of the instrument, whilst nitrocefin scores “++” as it requires only a common benchtop plate reader and cheap, readily available reagents.
Later, fluorescent probes which are easier to handle, were developed\textsuperscript{163,318,340}, which can be used to stain PBPs in membrane extracts in an analogous fashion and with similar results to the radiolabelled probes\textsuperscript{340}. This method does not allow for continuous observations of the reaction. Because the fluorescence intensity does not typically change following acylation, the fluorescent anisotropy method is used to differentiate between the bound and unbound states\textsuperscript{163,340}. The theory behind fluorescence anisotropy has been understood since the 1920s\textsuperscript{349}, but the first plate reader based instruments were only designed in the mid 1990’s, leading to a “burst” of research activity, resulting in many assays exploiting this method\textsuperscript{350}. Figure 1.5b and c give a simplified overview of the physical aspects of the process. At the simplest level, the binding of BOCILLIN FL to a larger molecule slows its rotation and this is measured by the intensities of the parallel and perpendicular emitted light.
a

Nitrocefin

CENTA

S2d

Detect with coupling reagent

Bocillin FL

Acylation of PBP

Biotinylated Ampicillin

b

c

Fluorescence Anisotropy

Fluorescence Anisotropy
Figure 1.5. Chemical probes for the behaviour of PBPs. (a) Chemical structures of some of the probes used in Table 1.2. The reaction that produces the observable change is shown. Release of the 2-mercaptoacetic acid leaving group of S2d can be monitored by absorbance, fluorescence or fluorescence of coupling agents, or by direct observation of the change in absorbance at 250 nm due to the cleavage of the thioester 168,200. (b and c) Simplified scheme describing fluorescence anisotropy (or fluorescence polarisation). (b) Fluorophores (ovals) are randomly orientated in solution, with those orientated parallel to the polarised incident light excited by it (cyan), those that are not parallel (black) are not excited. Excited fluorophores emit light (green) after their average fluorescence lifetime $r$. After time $r$ the fluorophores will have randomly rotated due to Brownian motion of the solution. If the speed of rotation is much greater than $r$ (as it is for small BOCILLIN FL molecules) emitted light from the solution has no overall polarisation, due to the randomness of the orientations of the excited molecules. The fluorescence anisotropy of the sample is correspondingly low. (c) In this case, molecules of fluorophore are bound to a much larger molecule (as would be the case for BOCILLIN FL binding to PBP3). Only molecules starting parallel to the polarisation of the incident light are shown. Now, in the time interval $r$, much less rotation of the fluorophore has occurred because larger molecules rotate more slowly due to Brownian motion. Emitted light will therefore have a larger component polarised parallel to the polarisation of the incident light and the fluorescence anisotropy signal will be greater.

Biochemically, CENTA, nitrocefin and S2d behave quite similarly. The ligand is turned over by the PBP and released to give a change in absorbance that is monitored by a spectrophotometer. The species responsible for the colour changes are shown in Figure 1.5a. PBP3 from P. aeruginosa turns over these substrates rapidly, faster than related class B PBPs (Chapter 6). The kinetics of nitrocefin and CENTA are discussed in more detail in Chapter 6.

There are no assays that accurately utilise the natural product of class B PBPs, leaving open questions as to its exact mechanism in nature. A recent paper investigating natural product turnover in class A PBPs showed there were few chemical recognition elements required by the donor substrate (left hand side of Figure 1.2b) and even fewer specifying the characteristics of the acceptor nucleophile (right hand side nucleophile in Figure 1.2b) 348. Functional assays are important because assays that rely on β-lactam probes may miss novel classes of compounds with functional inhibitory activity against PBPs (i.e. can disrupt the binding of the natural substrate), but which do not affect the acylation of β-lactams (or probes like S2d).

Mass spectrometry methods can provide label free, very short time scale (if combined with quenched flow) insight into the behaviour of covalent binders of PBPs 178,344. This has been used to find the acylation rates of PBPs by β-lactams,
without the need for a competing substrate. If a protein is covalently bound to an inhibitor and then digested by proteases such as trypsin, the masses of the resultant peptides can be analysed to find the location of the acylation. These techniques are investigated in Chapter 6.

Chapter 2 covers SPR in more detail.

Thermodynamic methods, investigating the effect of the ligand binding on protein stability are useful for understanding the EI complex, but provide little kinetic information. For β-lactams, it has been shown that the non-covalent components of the active site interactions contribute little to the overall acylation efficiency.

Each of the techniques shown in Table 1.2 has its own advantages and often a combination of multiple methods is the best approach, and gives the broadest insight into the problem (Chapters 5 and 6).

1.5 Conclusions

The successful use of antimicrobials allows for much of modern medicine, but antimicrobial resistance can render them useless. Like fire extinguishers, we do not appreciate the value of antimicrobials until we no longer have access to them in an emergency. Better policy and incentives are required to ensure future generations have access to these essential medicines. The threat of resistance necessitates constant discovery of new compounds whilst low sales volumes and the need to keep drugs as a "last line of defense" disincentivises companies from investing in research. At the same time, data from pharmaceutical companies appears to show that discovery of good antimicrobials is more scientifically challenging than discovery drugs for other therapeutic areas.

Inhibition of PBPs is a well-validated antibacterial approach but β-lactams, the only drugs to target them, are increasingly subject to resistance that threatens their efficacy. Other classes of compounds have been shown to bind and inhibit PBPs but none with the efficacy of the β-lactams. As such, the employment of new methods may yet lead to effective new inhibitor classes. This thesis describes the development of tools for drug discovery, insight into PBP3-mediated mechanisms of
resistance and two examples of previously undescribed PBP inhibitor binding modes.

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Chapter 2. Feasibility Studies of a Surface Plasmon Resonance system for PBPs

2.1 Introduction

When attempting to analyse small molecule interactions, it is often desirable to have multiple, independent measurements of the same interaction. Multiple assays to probe PBPs have been designed (see Table 1.2) but in many cases these cannot measure the on- or off-rates of the protein-small molecule interaction. Instead, for this, biophysical methods must be used. A number of these methods exist, each using a different biophysical property of the protein. Surface plasmon resonance (SPR) and biolayer interferometry both exploit optical phenomenon whilst other methods such as isothermal titration calorimetry measure thermodynamic properties of an interaction. SPR is sensitive, label free and can be used to find the $k_{on}$ and $k_{off}$ of a bimolecular reaction. Whilst these desirable attributes have seen its use increase over recent years, its sensitivity demands great care for successful use which can make the method development challenging.

This chapter describes attempts to develop a SPR system for measuring small molecule interactions with PBP3, examining various coupling methods and optimisations of the method required for detecting binding.

Many proteins, including PBPs and β-lactamases have been assayed using SPR. Various methods have been reported to couple a PBP to the chip: through azide “click chemistry”, binding of a β-lactam, moenomycin on to the chip surface to link the protein to the chip at the transpeptidation (TP) domain or transglycosylation (TG) domain respectively. Much of this work has been carried out by Vollmer and colleagues in order to investigate protein-protein interactions. However, due to the much higher molecular mass of proteins, SPR systems investigating protein analytes present very different challenges to those investigating small molecules. Since PBP3 lacks a TG domain, an accessible TP site for investigating inhibitor binding is required, so coupling via the TP domain is not
feasible. At the commencement of this work, SPR had not been used in the
detection of small molecule binding at the PBP transpeptidase site. However in April
2021 a report described SPR for use in the determination of the binding of
pyrrolidine-2,3-diones to *P. aeruginosa* PBP3 (PaPBP3)\(^{16}\), this is compared to this
work in section 2.3.6.

**Note on terminology**

*In contrast to other areas of life sciences, SPR literature the term “ligand” refers
to the protein bound on to the surface of the chip, whilst “analyte” is used for the
molecules flown over the top whose interactions are investigated. For consistency
with the literature, this notation will be used throughout this chapter.*

### 2.1.1 Biophysics of SPR

An SPR instrument is able to detect changes in the characteristics of a thin metallic
surface by detecting small changes in its refractive index. The detection method is
based on physical phenomena that occur when light is incident on the interface with
a material (e.g. gold) with a lower refractive index than the refractive index of the
material (e.g. glass) it is currently travelling through. At low angles, total internal
reflection (TIR) can occur, in which the incident light is reflected, but an additional
electronic wave, termed a "evanescent wave field" is generated through the material
with the lower refractive index. This wave decays exponentially and typically only
penetrates around a quarter of a wavelength of the incident light into the surface.
Away from certain resonant wavelengths, no energy from the incident beam is
absorbed and the intensity of reflected light equals that of the incident. However, if
the interfacing material surface is a metal, at specific combinations of the
wavelength of light, the angle of incidence and the refractive index of the interfacing
material, resonance with the electrons of the metallic surface (in practice this is a
gold surface), leads to absorption of energy by the metallic surface which can be
seen as a “dip” in the ratio of incident/ reflected light (Figure 2.1). In Biacore
systems, only monochromatic light is used, meaning the angle of incidence can be
varied to find and observe the dips \(^{17,18}\). The shape and position (as defined by the
angle) of the dips are affected by the refractive index of the material (Figure 2.1b).
Due to the thinness of the surface, its refractive index can be changed by binding proteins and even small molecules to the surface. This response is linear with the mass of bound material \(^{12,13}\). The changes in the position of the minima of the dip following changes in refractive index are converted into “Response Units” (RU) in Biacore machines, which can then be used to quantify binding events.

**Figure 2.1. The fundamental components of a Biacore SPR instrument.** Not to scale. (a) The microfluidics and gold surface are combined on a disposable chip cassette. Once inserted into the machine the gold is placed against the glass prism. Monochromatic light shines onto the gold surface, is reflected with total internal reflectance, and the intensity of the reflected beam is measured by the diode array. The angle of the incident beam can be varied to detect the “dips”. The underside of the gold chip is coated with carboxymethylated dextran to which the ligand is coupled. Analytes can be flown through the microfluidics once the ligand has been bound and the response to their binding measured. (b) Dips recorded by the SPR of the flow cell 1 (blue) and 2 (grey) of the ligand-coupled SA chip (see below). Reflectance, the ratio of the intensity of the reflected and incident beams are plotted against the angle of incidence. The peak minima is converted to the response in response units (RU). The effect of the ligand binding can be seen in the right shift of the curve (as shown by the red arrow), this change corresponds to a 5700 RU change in response. Reflectance is measured by ratio of reflected light to incident light.
2.1.2 Instrumentation

Continually monitoring the surface whilst flowing analytes across it allows measurement of the interactions of the ligand (bound to the metal surface) and the analyte (in solution). An important factor when designing an SPR system is to find an appropriate coupling system for attaching the ligand to the surface. An ideal coupling method is one that gives a stable baseline, with the protein exposed to the flowing buffer at the correct density. For Biacore systems (on which all work in this chapter was carried out) 21, a variety of coupling methods exist as pre-prepared kits 18 or custom surfaces can be created from blank chips 22.

The choice of coupling method depends on: the protein; the aim of the experiment; the ease of regeneration of the system (see below); and size of the analyte being investigated. During the course of this work, five surface coupling methods were investigated.

In any experiment the first step is to attach the protein to the chip surface. The practicalities of this are discussed below but whichever method is used to carry this out, a linear relationship between the mass bound and change in response remains. As such, following immobilisation, the concentration of the protein on the chip can be easily calculated using Equation 2.1 23,24. For example, for couplings in which a 10,000 RU of PaPB3 (MR: ~60,000 Da) is bound to the chip ($R_{ligand}$) a concentration of ~1.67 mM of protein is bound to the flow cell. The matrix of dextran used on the CM5 chips (used for amide coupling and streptavidin capture) is not a monolayer as depicted in Figure 2.1a but forms a 3D surface.

\[
[\text{ligand}] \ (\text{mM}) = \frac{R_{\text{ligand}} \times 10}{MR_{\text{ligand}}}
\]  

(2.1)

Additionally, the theoretical maximal amount of analyte ($R_{\text{max}}$) that can be bound to the chip can be calculated by equation 2.2 using ratio of the molecular masses of the analyte and ligand and the valency of the interaction 25. In all cases, a simple 1:1 valency is assumed.
\[ R_{max} = \frac{M_{\text{analyte}}}{M_{\text{ligand}}} \times R_{\text{ligand}} \times \text{valency} \] (2.2)

2.1.3 Coupling and Regeneration

During SPR, the surface can be simplistically described as being in one of the three states shown in Figure 2.2: the uncoupled chip surface (state 1), the ligand coupled to the chip (state 2) and the analyte bound to the ligand, which itself is coupled to the chip (state 3). After analyte binding, the surface must be regenerated in order for a new analyte or different concentration of the same analyte to be applied. As shown in Figure 2.2, this regeneration can go via two routes, either returning to state 1, a protein-free chip (route A in Figure 2.2) or to state 2, an unbound protein (route B in Figure 2.2). Route B is perhaps the more common method, but route A can be used for effectively irreversible interactions such as those of β-lactams with PBPs. Three methods for route A were initially investigated: a streptavidin “CAPture” system and two methods utilising a thiol linkage \(^{21}\). Route B regeneration was also attempted for an amide-coupled and a streptavidin-coupled chip.

![Diagram showing coupling and regeneration methods]

**Figure 2.2. Generalised coupling and regeneration methods.** A chip with an unmodified surface (1) is first coupled to the ligand (2) then the analyte is bound (3). The chip can be regenerated via route A, in which the surface is returned to a ligand-free state, or route B in which the analyte is removed from the ligand.
2.2 Methods

2.2.1 Instrument

A Biacore T200 instrument was used for all experiments. Coupling experiments were run using the in-built “Biacore T200 Control Software” (Version 2.0); either as a Manual Run (Thiol coupled experiments) or in-build “Wizard” programs (all others). Analyte binding experiments were performed using the single cycle kinetics or multi cycle kinetics programs of the software. The sample compartment and chip were maintained at 25 °C throughout. The “Desorb” cleaning program was implemented between runs and the “Desorb and Sanitise” program ran every month on the instrument.

Details of the procedures for the coupling regimes are given in section 2.3.

Prior to running the analyte samples 20 “conditioning” start up cycles were used to stabilise the baseline response of the instrument. Solvent correction was run before and after the analyte experiments and the average of the two used to generate calibration curves.

All chips were “S series” chips purchased from Cytiva, handled following manufacturers directions.

2.2.2 Buffers

Running buffers were typically either: 0.01 M N-(2-hydroxyethyl) piperazine N’-(2-ethanesulfonic acid) (Hepes) pH 7.4, 0.15 M NaCl, 3 mM ethylene diamine tetraacetate (EDTA), 5 % (v/v) dimethyl sulfoxide (DMSO), 0.005 % (v/v) Surfactant P20 (for CAPture and thiol coupled experiments); or 10mM sodium phosphate, 2.7 mM KCl, 137 mM NaCl pH 8, 5 % (v/v) DMSO, 0.05 % (v/v) Tween 20 (for amide and streptavidin coupled experiments). Buffers were made up from 10 x stock solutions to 1.05 x solutions then 5 % (v/v) DMSO was added. Buffers were filtered and degassed under vacuum prior to use. Buffers were used within days of their preparation. A separate aliquot of the buffer without 5 % DMSO was kept for preparing the solvent correction curves. Analytes were diluted from high (either > 100 mM in DMSO or dissolved powder) concentration into the running buffer.
immediately prior to the beginning of the experiment. Upon changing the buffer system in the instrument in preparation for a run, the “Prime” procedure was run twice.

2.2.3 Reagents

Biotinylated and cysteine mutant PaPBP3 proteins were produced by Dr. Dom Bellini. Briefly, plasmids used previously for the production of PaPBP3 (residues 50 - 579) \(^\text{26}\) were modified at the C-terminus (immediately before the STOP codon) by the addition of either a short ‘Avitag’ (GLNDIFEAQKIEWHE) or a single residue of cysteine (no other cysteines are present in the protein). For the biotinylated protein, the plasmid was transformed into \textit{E. coli} BL21 cells as previously \(^\text{26}\), with an additional plasmid \textit{pBirAcm} (Avidity) co-transformed into the cells. The manufacturer's protocol (Avidity) was followed for the simultaneous expression of PaPBP3 and BirA as well as the enzymatic reaction to add \(\text{d-biotin at the Avitag site. Both proteins were purified as previously described } \(^\text{26}\). MreC (truncated to residues 58-367) was a kind gift from C. Graham.

Reagents were purchased from Sigma or Cytiva.

Solutions of dithiothreitol (DTT), tris(2-carboxyethyl)phosphine (TCEP), N-hydroxysuccinimide (NHS), 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC) and 2-(2-pyridyldithio)ethaneamine hydrochloride (PDEA) were prepared directly before use from powder stocks.

Analytes were diluted to their desired concentration from high concentration stocks (either > 100 mM in DMSO or dissolved powder) by serial dilution into the running buffer, keeping DMSO (if present in the stock) constant across all the injections.
2.2.4 Procedure for preparation of activated thiol PaPBP3

**Figure 2.3. Preparation of activated thiol PaPBP3.** PaPBP3 with a C-terminal cysteine was partially dimerised (by disulphide bonds between molecules) after purification, presumably due to spontaneous oxidation in air. 2.5 mM TCEP was used to reduce the dimers to monomers, which were then treated with 2 mM 2,2’-Dithiodispyridine (2,2-DTP) to produce 2-thiopyridine conjugates of PaPBP3, across an asymmetric disulphide bond. Unreacted TCEP and 2,2-DTP were separated from the protein by elution through a gravity-driven PD-10 column.

To prepare activated PaPBP3 for surface thiol coupling, and remove any disulphide-bridged dimers of PaPBP3 (formed by spontaneous oxidation during purification) the following procedure was followed (Figure 2.3). PaPBP3 with a C-terminal cysteine insertion (Cys580, 20 µM, in 10mM 2-amino-2-(hydroxymethyl)propane-1,3-diol (Tris), 500 mM NaCl, pH 8 buffer) was reacted with 2.5 mM TCEP before 2 mM 2,2’-Dithiodispyridine (2,2-DTP) was added. The solution was then loaded onto a gravity-driven PD10 column (Cytiva, running buffer: 100 mM sodium phosphate, pH 6 buffer) and the protein-containing fractions pooled, to allow separation from the TCEP- and 2,2-DTP-containing fractions. The lack of dimerisation (through Cys580) of the protein was confirmed by gel electrophoresis (without reducing agents in the loading dye). Protein fractions did not react with 5,5-dithio-bis-(2-nitrobenzoic acid) (DTNB), indicating a lack of free thiols. Conjugates similar to this have been shown to have stability in the order of months \(^7\), although this conjugation was used within days of preparation.

2.2.5 Reference Surfaces

The microfluidics within the Biacore chips allow for configurations in which the analyte is first flown over a “reference surface” which lacks the protein of interest so
that the non-specific and bulk phase changes caused by the analyte injection can be accounted for. Reference surfaces are used in all analyte binding experiments but varied between the coupling systems used; for the CAPture system streptavidin without biotinylation was used. An injection of the oligo-streptavidin solution only was carried out for this surface. For the thiol-based systems a free cysteine was used, coupled to the chip by NHS/EDC; for the streptavidin system streptavidin without biotinylation was used (no modification); for the amide coupling system ethanolamine was used (or MreC for section 2.3.5.4), coupled to the chip by NHS/EDC. In each case the reference surface was prepared using the same procedures as the active surface, but with the protein injection omitted.

Typically, the reference surface was in flow cell 1 and the “active” (ligand-activated) surface was in flow cell 2. During an analyte binding experiment, the instrument is set to record the response of flow cell 1 and 2 and find the difference between them to give the response signals.

2.2.6 DMSO Calibration Curves

DMSO has a significant effect on SPR, with a 1 % difference in DMSO concentration between the injected sample and the running buffer leading to 1000 RU response. Making up samples to the same exact concentration of DMSO as the running buffer is challenging, and reference subtraction is generally insufficient because the DMSO displaces a volume of water on the chip surface but the extent of this varies depending on the surface modification. It is therefore important to know how the reference and active surfaces respond to a DMSO pulse. For this a DMSO calibration curve is needed. An example is shown in Figure 2.4. A series of known concentrations of DMSO are injected and the response on the active and reference surfaces are recorded. The reference response and the difference between the active and reference are plotted against each other and used to construct a calibration curve. After the sample containing DMSO is injected, a square pulse (correcting for the difference in the response of the active and reference cells to DMSO) is subtracted from the reference-subtracted signal, which corrects for the calculated difference in response of the active and reference cells to DMSO. This correction is carried out automatically by the control software.
Figure 2.4. A typical Solvent Correction Curve. Eight concentrations of DMSO were flown over both the active (blue) and reference cells (orange) to generate a solvent correction calibration set. Due to the different surfaces, the active and reference flow cells respond differently to DMSO, one such difference is highlighted. Concentrations of DMSO injected are shown on the graph. The buffer contained a nominal 5 % (v/v) DMSO.

2.2.7 Analysis

Curve analysis was performed in the “Biacore T200 Control Software” (Version 2.0). This program can perform reference and solvent corrections. Figures of the traces were produced in Matlab (Version 2020a) using custom scripts. Figures of the coupling models were generated in ChemDraw (Professional version 17.0, PerkinElmer Informatics).

2.3 Results and Discussion

2.3.1 CAPture System
**Figure 2.5. Route A regeneration: CAPture coupling system.** (a) Scheme for the coupling system. Biotin was added to the PaPBP3 during purification by the addition of the small Avitag (Avidity) sequence to the protein sequence, which is recognised by the enzyme BirA (Avidity) which ligated biotin at a specified point within the binding sequence \(^{28,29}\). The chip (which is supplied with a single stranded oligonucleotide bound to the surface) was prepared by addition of a complementary streptavidin-oligonucleotide conjugate. The oligonucleotides annealed, resulting in a streptavidin-activated surface. When the protein was flown over this surface, the streptavidin bound to the biotin found on the protein with high affinity, coupling the protein to the chip. Once the analyte has bound, the chip can be regenerated by the addition of a solution of guanidine in NaOH to denature the double stranded oligo and return to state 1. Numbers in bold correspond to the three states of the chip shown in Figure 2.2. (b) Response of flow cell 2 to reagents. The chip was activated by flowing over oligonucleotide-coupled streptavidin (proprietary solution, 50 \(\mu\)g/mL, flow rate 2 \(\mu\)L/mL, 5 minute contact time), before the biotin-coupled PBP3 (1.25 mg/ml, flow rate 2 \(\mu\)L/mL, 10 minute contact time) was injected. (c) an injection of guanidine HCl (6 M with 250mM NaOH, flow rate 30 \(\mu\)L/mL, 2 minute contact time) removed both the streptavidin-coupled PaPBP3 and the chip was ready to be used again.
With the CAPture system \(^{30}\), regeneration is possible due to the reversible denaturing and annealing of complementary DNA oligonucleotides, attached to a streptavidin molecule, to capture a biotin-bound protein. Denaturing of the DNA allows the protein to be removed and the surface regenerated (Figure 2.5a). This kit is intended for the use of protein-protein interactions, but there is no fundamental reason why it cannot be used to study protein: small molecule interactions. This method utilises a commercially available kit (Cytiva), in which a chip is supplied with oligos coupled to its surface at predetermined and unchangeable density. Unfortunately, this determines the maximum load of the protein onto the chip, since only one protein molecule can be coupled to each oligo. Consequently, several attempts were made at coupling the protein onto the chip and each time only \(~4000\) RU of protein could be bound, although regeneration was successful and reversible. Additionally, there appeared to be significant amounts of baseline drift which remained even after long equilibrations.

### 2.3.2 Thiol-based System

The basis of reversibility in thiol-based systems is the reversibility of disulphide reduction/oxidation cycles (Figure 2.6). There are two similar systems.

\[
\text{Disulphide reduction/oxidation reaction}
\]

\[
R_1-S-S-R_1 \xleftarrow{\text{Reduction}} \xrightarrow{\text{Oxidation}} R_1-SH \times 2
\]

\[
\text{Thiol/disulphide exchange}
\]

\[
R_1-S-S-R_2 + HS-R_3 \xleftarrow{\text{Thiol/Disulphide exchange}} R_1-S-S-R_3 + R_2-SH
\]

**Figure 2.6. Simplified reactions of thiols used for the preparation of the thiol-coupled chip.** Disulphide reduction can be driven by reducing agents such as TCEP or DTT. Oxidation can happen spontaneously in air over time and will inactivate any free thiols, preventing them from acting as nucleophiles in the thiol/disulphide exchange reaction. Thiol/disulphide is the reversible exchange of one thiol for another within a disulphide bond. The position of equilibrium between the two states is determined by the relative reactivity and stability of each thiol as well as relative concentrations \(^{35,36}\).
In the first system “ligand-thiol”, the reduced, nucleophilic, sulphur is found on the ligand \(^{31,32}\), whilst in second “surface-thiol”, the nucleophilic sulphur is on the chip (Figure 2.7 and 2.8) \(^{32-34}\). The resultant ligand-coupled complex is similar (note the additional carboxyl group in the surface-thiol system (Figure 2.7 and 2.8), but the different methods differ in their re-activation following regeneration.

Using both methods, immobilisation was successful, with 4,552 RU and 7,367 RU binding for the surface and ligand thiol systems respectively. However, in both cases it was not possible to regenerate the chip, with either DTT or TCEP reducing agents at concentrations up to 1 M. Injections of sodium hydroxide (100 mM) did not result in a large reduction in binding, indicating it was not a purely non-covalent attachment (which would not be removed by reducing agents). Long injections (1 hour) with DTT also failed to remove ligand, indicating this is not an issue simply with the rate of reaction.

The lack of ability to remove the PaPBP3 from the chip could therefore be explained several ways: i) incomplete reaction of PDEA or cysteine (depending on the method) and activated chip surface, which would leave free, unreacted NHS-esters to couple covalently to the protein’s free amines (same reaction as used in amide coupling). Long injections with excess of PDEA or cysteine were used to attempt to prevent this issue from occurring, but it is possible these were insufficient. ii) Disulphide bonds may have been inaccessible to the reducing agents, by the protein burying the newly-formed thiol away from the solvent. This is feasible, especially given the electrostatic attraction between the negatively charged surface and positively charged protein at neutral pH. It could potentially be addressed by introducing longer linking molecules, which would act to separate the protein and surface, allowing more access to the disulphide bond. The reversibility of the thiol bond in solution was confirmed by gel electrophoresis in the presence of DTT.

Without the expected regeneration it was decided to not pursue this surface further.
Figure 2.7. Route A regeneration: thiol coupling. (a) Scheme for ligand thiol coupling. A CM5 (Cytiva) chip was used as the starting surface, which is activated by NHS and EDC coupling reagents. PDEA was coupled to the surface by its amine group. Thiol exchange (Figure 2.6) with a free sulphur on the PaPBP3 (introduced by the engineering of a cysteine into the C-terminal of the protein) coupled the protein to the chip. Once the analyte was bound to the surface, Dithiothreitol can be used to reduce the disulphide, regenerating the surface, before another thiol exchange reaction with pyridine-2-thiol re-activates the surface. (b) Scheme for surface thiol coupling. The protein was prepared as described (section 2.2.4) to produce a 2-thiopyridine conjugate of PaPBP3. The chip was prepared using NHS/EDC coupling reagents which were used to couple a molecule of cysteine onto the chip surface. This was then reduced by TCEP to produce a free sulphur which reacted with the protein disulphide to couple the protein onto the chip. After analysis, regeneration can occur by reduction of the disulfide, once again releasing a free-sulphur to react on the next cycle.
2.3.3 Streptavidin coupling

A second capture system uses the high affinity interactions of biotin (on a biotinylated protein) and streptavidin (pre-bound to the chip surface)\textsuperscript{37}, but unlike the CAPture system, this is irreversible and requires route B regeneration (Figure 2.9). The method may be better than the amide coupling system because all proteins are orientated similarly due to the fixed position of the biotin tag and nocoupling of important lysines can occur.
Figure 2.9. Route B regeneration: streptavidin coupling system with biotinylated PaPBP3. (a) Biotinylated PaPBP3 was prepared as with the CAP system, using the BirA/Avitag system (Avidity). The protein was then linked to the chip by the high affinity biotin-streptavidin interaction. As with amide coupling this is an irreversible immobilisation and regeneration is only possible by route B. (b) Response in flow cell 2 to reagent injections. The streptavidin on the chip was washed three times (3 x contact time 1 minute, flow rate 5 µl/min) with a solution of NaOH and NaCl (50 mM and 1 M respectively in water) then the biotinylated PaPBP3 (14 µM, in sodium phosphate buffer 10 mM, 500 mM NaCl, pH 8, total contact time ~17 minutes, flow rate 5 µl/minute) is flown over the surface until saturation was achieved: 5,700 RU of protein was bound.
2.3.4 Amide Coupling

An amide coupling system can be used to bind the ligand to the chip. This method will couple any solvent exposed amines on the protein to the chip surface. PaPBP3 has 25 lysine side chains as well as the N-terminal amine which may be coupled. These amines form an amide with carboxylic acid groups of the CM5 carboxymethylated dextran (Figure 2.10). The disadvantage of this method is that catalytically important residues (in particular Lys297 and Lys484) may get acylated which will presumably block the active site and lower the effective \( R_{\text{max}} \). Additionally the random sampling of coupled lysine residues leads to a range of orientations and conformations of the protein on the chip, which can create an inhomogeneous surface that may interfere with the measurement of kinetics.

In order to optimise the coupling reaction, “pre-concentration experiments” were performed by the instrument “Wizard”. In these experiments protein was injected onto the unmodified CM5 chip surface and the response recorded. If sufficient protein was able to bind non-covalently to the chip surface, a large peak will be recorded during the injection (Figure 2.10b). The weakly bound protein quickly dissociates after the injection (with a NaOH wash to disrupt higher affinity charge-charge interactions) allowing the surface to be activated for covalent interaction. Preliminary experiments found that 1 \( \mu \)M of PaPBP3 in a low salt, pH 6 sodium phosphate buffer (50 mM NaCl, 10mM sodium phosphate) gave a large pre-concentration response.

2.3.5 Analyte Binding

2.3.5.1 Clinical \( \beta \)-lactams

Initially, the interactions of PaPBP3 with clinical \( \beta \)-lactams (ampicillin and cefotaxime) were investigated. In these cases a technique called single cycle kinetics was used. This method attempts to collect several concentrations of analyte binding data using sequential increases in analyte concentrations. This avoids
Figure 2.10. Route B regeneration: amide coupling system. (a) Amide coupling links the PaPBP3 to the CM5 chip with an amide bond between the carboxyl of the carboxymethylated dextran chip surface and any free amines (lysine side chains and terminal amine) on the protein. This is an irreversible reaction and regeneration can only be achieved via route B. (b) Response of flow cell 2 to coupling reagents. A preconcentration test was initially carried out (see text), achieving around 16,000 RU of protein onto the surface. The majority of this protein was washed off the chip at the end of the injection but any remaining non-specifically bound protein remaining was removed with an injection of NaOH (100 mM in water, contact time 1 minutes, flow rate 30 μl/min). A mixture of NHS/EDC (0.1 M and 0.4M respectively in water, contact time 7 minutes, flow rate 10 μl/min) was used to prepare the carboxymethylated dextran surface for covalent coupling to the protein. PaPBP3 was then flown over the surface in multiple injections (total contact time ~17 minutes, flow rate 5 μl/min) achieving a final protein load of 12,086 RU. Ethanolamine (1.0 M, pH 8.5, contact time 7 minutes, flow rate 10 μl/ml) was then injected to block any unreacted activated surface molecules.
the issue of a single high concentration binding to all the active sites and inactivating the protein, preventing any more data from being collected. For three chip surfaces (CAPture, streptavidin coupling and amide coupling) β-lactam interactions did not generate any specific signal (Figure 2.11). The cause of this is unclear.

**Figure 2.11. β-lactam binding to various chip surfaces** (a) Injections of 5 concentrations of cefotaxime (12, 80, 400, 2,000, 10,000 nM) were flown over a PaPBP3 CAPture coupled chip surface using single cycle kinetics. Expected $R_{\text{max}}$ of 30 RU. Additionally the baseline following injections was unstable which is undesirable. (b) Injections of 5 concentrations of Ampicillin (8, 16, 32, 64, 128 μM) were flown over a streptavidin coupled chip surface using single cycle kinetics. Expected $R_{\text{max}}$ of 27 RU (c) Injections of 5 concentrations of cefotaxime (33.3, 100, 333, 1000, 3330 nM) were flown over an amide coupled chip surface using single cycle kinetics. Expected $R_{\text{max}}$ of 46 RU. Non-specific binding is observed (steep, concentration non-specific increases during injection). In each case the black bars indicate the injection pulse (contact time 120 s, flow rate 30 μl/min).
2.3.5.2 Nitrocefin

In contrast to clinical β-lactams, nitrocefin could be used to generate positive response curves on an amide coupled surface (Figure 2.12). Nitrocefin has previously been used in an SPR system to investigate β-lactamase behaviour in a custom-made coupled SPR-absorbance system. A concentration dependent response was observed for nitrocefin, with agreement between repeats. Nitrocefin was anticipated to be a better probe due to its ability to react with PaPBP3 then de-acylate (Chapter 6), which is effectively route B regeneration. However, the response could not be fully regenerated with the baseline not returning to zero between injections, even after 1000 s of dissociation or being left overnight. If the value of $k_{\text{cat}}$ for PaPBP3 nitrocefin turnover (~ 40 min$^{-1}$: Chapter 6) is assumed to occur under these conditions then it would be expected that the protein would regenerate with a half-life of 1.5 seconds. The lack of regeneration meant that the baseline increased after every injection.

This increase in the baseline did not prohibit further binding of nitrocefin to the protein. For example, due to the sequencing of the injections, half of the injections in Figure 2.12a occur after the 8 μM injection (see legend of Figure 2.12), which by itself lead to ~80 RU of response, more than the expected $R_{\text{max}}$ (76 RU). Such agreement between repeats as is seen would not be expected if the protein was becoming progressively inhibited. The total change in response over the experiment was much greater than the expected $R_{\text{max}}$, with a total change in response in flow cell 2 of 300 RU. Such a response can be indicative of a large contribution to the signal from non-specific binding (but high affinity binding, hence its irreversibility) of the nitrocefin to the protein.

The irreversible binding was not due to interactions of nitrocefin with the chip. Figure 2.12b shows the individual responses of flow cell 1 and 2 to an injection of 8 μM nitrocefin. The trace shows that whilst there is an interaction of nitrocefin with the chip, this dissociates after the analyte pulse. This means that the gain in response seen in Figure 2.12a is due to binding of nitrocefin to the protein itself (which is only present in flow cell 2).
It is unclear whether the observation of apparent greater-than-stoichiometric binding of nitrocefin to PaPBP3 (discussed in Chapter 6) is relevant here. PaPBP3 does not appear to approach saturation even after a mass equivalent to ~4 x nitrocefin molecules bind (change in baseline/expected $R_{\text{max}}$: 300/76 Figure 2.12), in contrast to the two binding events anticipated from crystallographic studies. SPR would be a useful tool for the investigation of specific greater-than-stoichiometric binding but unless conditions are identified in which non-specific binding does not occur these studies are not feasible.

![Graph](image)

**Figure 2.12. Investigation of nitrocefin binding.** (a) Reference- and solvent-corrected responses to five concentrations of nitrocefin (0.5-8 μM, see legend) flown over a PaPBP3 amide-coupled CM5 chip. Injections (contact time 120 s, flow rate 30 μl/min, dissociation 1000 s) were run in sequence from the lowest concentration to the highest and then lowest to highest again (giving the two repeats), with a control injection of the running buffer (0 μM nitrocefin) between each experimental injection. Expected $R_{\text{max}}$ of 76 RU. Significant increases in the baseline are observed throughout the experiment ({(reference-subtracted response at the end) - (reference-subtracted response at the beginning)}) such that there was an increase in the baseline of almost 300 RU across the entire experiment. In (a) each of the traces is zeroed at the beginning of the experiment to allow easier interpretation. (b) Responses of flow cells 1 (reference) and 2 (active) to the first 8 μM injection.
2.3.5.3 Benzoxaborole

To examine whether the issues of lack of interpretable binding were limited to β-lactams, a benzoxaborole (13) analyte was investigated. Biochemical evidence suggests that this binds in a weak reversible manner ($K_i$ of ~80 μM) and may be more amenable to SPR. This ligand appears to show PBP-binding activity, however its behaviour is very difficult to interpret, and does not fit into the expected models of ligand:analyte interaction (Figure 2.13).

The rapid increase in the response after the analyte is injected is either indicative of a very rapid on-rate or non-specific interactions. The shape of the curves during injection cycles themselves is not a smooth curve and even at low concentrations does not appear to show any of the curvature expected for bimolecular interactions (see the 8 μM curve, Figure 2.13b). Without this curvature, it is difficult to distinguish specific and non-specific binding (e.g. see the square shape of the non-specific binding curve in Figure 2.12b).

Another issue is that $R_{max}$ (around 100 RU) is not approached, even for high concentrations of analyte (2 mM is 25 times above the biochemically determined equilibrium constant of 80 μM). This could perhaps be accounted for by inactive conformations of the ligand, which would reduce the effective $R_{max}$. However, this is another sign that the system is not behaving as expected, and should be interpreted with caution. Using such high concentrations of analytes typically leads to more non-specific binding.

SPR curves can be simulated with known constants. Figure 2.13b shows an example using values of $k_{on}$ and $k_{off}$ which could give a $K_i$ of 80 μM (although an infinite amount of other combination of these values would also give this dissociation constant). Some similarities can be drawn between Figure 2.13a and b, but it is difficult to be confident in this interaction.
Figure 2.13. Investigation of benzoxaborole binding. (a) Reference- and solvent-corrected responses to nine concentrations (8 µM - 2 mM) of benzoxaborole 13 (Chapter 5) flown over a PaPB3 amide-coupled CM5 chip. Injections (contact time 120 s, flow rate 30 µl/min, dissociation 1000 s) were run from lowest to highest concentration with repeat injections of a concentration run in pairs. Control injections of the running buffer (0 µM benzoxaborole) were run at the beginning of the experiment. Expected $R_{\text{max}}$ of 96 RU. In each of the traces here, the response is zeroed at the beginning of the experiment to allow easier interpretation. (b) Simulated SPR traces using biochemical data (Chapter 5). Data was generated using the program "SPR-Simulation" (Version 1.3.2.3 Arnoud Marquat, sprpages.nl) using the following settings; $k_{\text{on}}$: $1 \times 10^3$ M$^{-1}$ s$^{-1}$; $k_{\text{off}}$: $8 \times 10^{-2}$ s$^{-1}$; $R_{\text{max}}$: 100 RU; Noise: 0.4 RU; Drift: 0.02 RU s$^{-1}$; Contact time 120 s. These values of $k_{\text{on}}$ and $k_{\text{off}}$ give an equilibrium constant ($k_{\text{on}}/k_{\text{off}}$) of 80 µM, approximately the value determined by bocillin competition (Table 5.3).
2.3.5.4 Reasons for Unresponsiveness

For the clinical β-lactams tested, it appears as if no interaction is occurring with the protein at all. This would suggest that the protein is inactive under these conditions. In solution, the protein was demonstrated to be active in the buffers used, however the environment of the coupled chip may cause the protein to behave differently than it does in solution.

The effective concentration of PaPBP3 on the chip surface is in the mM range (see section 2.1). This high concentration may be leading to local aggregation of the protein (the flexible 3D dextran matrix may allow contact between adjacent protein molecules), and its inactivation. However aggregation is typically observable in the response of the chip (Claire Shepherd, Cytiva, personal communication). Alternatively, the protein may be induced into an inactive conformation by protein-protein interactions under these conditions. Steric crowding of the proteins by nearby proteins or the dextran matrix (for CM5 chips used in amide coupling reactions) may be blocking access to the protein active site. The coupling method and surface are known to have a significant effect on the activity of the protein (Claire Shepherd, Cytiva, personal communication).

2.3.5.5 Non-specific Binding and Alternate Reference Surface

For nitrocefin and the benzoxaborole, there appears to be a large amount of non-specific (irreversible and non-stoichiometric) binding to the protein. This obscures any specific interaction that may be occurring between ligand and analyte and makes the results difficult to interpret. The non-specific binding appears to be mainly with the protein and not the chip surface itself, and occurs despite the presence of the Tween 20 detergent (0.05 % (v/v)). Similar non-specific binding appears to occur for another PaPBP3 system (section 2.3.6).

An alternative, protein-coated reference surface can be used to account for this difference in the degree of analyte binding to the active surface and the simple, ethanolamine reference surface. A surface was loaded with MreC (a bacterial cell wall protein not known to bind β-lactams) using amide coupling as shown for PaPBP3 (Figure 2.10), to a response level of 3500 RU and then the analyte was
flown over and the sensorgrams for the PaPBP3-containing active cell was corrected with the response of the same analyte over the MreC reference cell.

![Graphs showing binding response](image)

**Figure 2.14. Binding of benzoxaborole 13 corrected with an MreC reference.** (a) Active cell - coupled to PaPBP3 corrected with the response on an MreC-coupled reference cell. (b) Response on the MreC cell to 13. Colours correspond to the concentration of 13. Injections (contact time 120 s, flow rate 30 µl/min, dissociation 1000 s) were run from lowest to highest concentration with repeat injections of a concentration run in pairs. Expected $R_{\text{max}}$ of 96 RU. In each of the traces here, the response is zeroed at the beginning of the experiment to allow easier interpretation.

Compared to Figure 2.13a, the addition of the MreC reference leads to an improvement in the corrected signal by reducing the degree of non-specific binding.
However, challenges remain: there are significant differences between the two repeats of the concentrations (although these are generally not as pronounced in the dissociation phase). Even at \( \sim 2 \) mM of 13 there is not saturated binding of analyte (it is still increasing towards the end of the pulse and far from the \( R_{\text{max}} \) of 96 RU, contrast to the simulated Figure 2.13b), which would be expected if the value of the \( K_i \) found by the fluorescence anisotropy assay is correct (\( \sim 74 \) \( \mu \text{M} \) - Chapter 5). Additionally, the signal from the reference cell (Figure 2.14b) is not the square wave expected for a simple fast associating and dissociating non-specific analyte-ligand interaction (For example that seen at flow cell 1 during the interaction of nitrocefin with ethanolamine in Figure 2.12b). Under these circumstances, subtracting this reference signal from the active may lead to the introduction of further errors. Using another protein to coat the reference surface may avoid this challenge.

2.3.5.6 Buffer Screen

One factor relevant to non-specific binding is the buffer used. This was briefly investigated using an inactivated protein. Five different buffers were investigated for their effect on the non-specific binding of ampicillin, with the trace of the active and reference and difference between the two shown. Omission of the salt from the buffer leads to a small amount of non-specific binding of the ampicillin to the protein, which is only slowly washed off. This is likely due to the high \( \text{pI} \) of PaPBP3 (\( \text{pI} \) 8.8 as estimated from the sequence by protPAM 40) which may indicate that the protein has a significant positive charge at neutral pH. Non-specific binding can be suppressed by raising the salt concentration ((Figure 2.15c) or raising the pH (Figure 2.15d,e) which makes the protein less charged and the interactions less strong. However, the protein may be less stable at these conditions and the interactions less physiological.
Figure 2.15. Buffer Screening. In each case ampicillin was injected over a streptavidin coupled chip equilibrated in the following buffers: (a) No NaCl, 10mM Tris pH7, 0.05 % (v/v) Tween 20, 5 % (v/v) DMSO (b) 10mM Tris pH7, 150mM NaCl, 0.05 % (v/v) Tween 20, 5 % (v/v) DMSO (c) 500 mM NaCl, 10mM Tris pH7, 0.05 % (v/v) Tween 20, 5 % (v/v) DMSO (d) 10mM Tris pH8, 0.15M NaCl, 0.05 % (v/v) Tween 20, 5% (v/v) DMSO (e) 10mM 2,2’-(propane-1,3-diyl)iminobis [2-(hydroxymethyl)propane-1,3-diol] (Bis-tris propane), pH 9, 0.15 M NaCl, 0.05 % Tween 20, 5 % (v/v) DMSO. Injection is ampicillin (20 μM, contact time 120 s, flow rate 30 s). The response of the active (blue) and reference (orange) as well as the active-reference subtraction (green) is shown. No specific binding is expected as the protein was previously treated by the addition of an excess of ampicillin, which should block the active site. Before each injection the “prime” procedure was performed and the chip exposed to 20 startup cycles in the new buffer.

2.3.6 Comparisons with López-Pérez et al. (2021)

During the write-up of this thesis, López-Pérez et al. (2021) reported an SPR system for the detection of the binding of pyrrolidine-2,3-diones (Table 1.1) to PaPBP3. The SPR system they report uses amide coupling with a similar buffer
system to ours (although the salt and buffer concentrations were much lower: (13 mM NaCl, 257 μM KCl, 171 μM KH₂PO₄, 950 μM Na₂HPO₄, 0.05% Tween-20 and 5% DMSO)) and the pH was lower: pH 7.4 vs pH 8. They also use a Biacore T200 and a CM5 chip, on which they achieved comparable coupling levels (12,086 RU vs 11,820 RU, for this work and López-Pérez et al. (2021) respectively), with an ethanolamine reference surface. Kᵋ values determined for 4 out of 5 of their compounds were on the same order of magnitude as the IC₅₀s determined by S2d and bocilllin FL fluorescent polarisation, but one showed significantly weaker binding as determined by SPR compared to the other assays. This validates their SPR method to some degree, but they did not include an external control (e.g. a β-lactam) to compare the results to, likely as a result of similar challenges to those described in section 2.3.5. Sensorgrams have a similar shape to those shown in e.g. Figure 2.13, which they ascribe to very fast association and dissociation rates. At least some of this response appears to be due to non-specific binding because at least for one compound low concentrations of inhibitor do not approach zero response (the authors didn’t comment on this). Additionally they use harsh conditions for regeneration (50 % DMSO for 30 s, whereas the chip CM5 tolerance for DMSO only 10 % for 1 minute ⁴¹) which may be required to remove irreversible binding of the compound (although this is not referenced). This regeneration method was not tested in our system. This report appears to partially corroborate our findings that non-specific binding is a problem for amide coupled PBP3 SPR systems, but demonstrates it can be used to determine Kᵋ values that correlate to competition assays.

2.4 Future Work

It is possible that other coupling methods or conditions could be found in which the protein is active. Both the thiol-coupled and biotin coupled proteins were modified at the C-terminus, so N-terminal modification could be investigated. After the conclusion of this work, a team at AstraZeneca published a paper describing three methods of creating a stable but reversible SPR system ⁴². These include the use of “switchavidin”: a mutant of streptavidin with biotin affinity that can be modified by changing the pH ⁴². Investigation of these methods for a PBP3-based SPR system may give better responses than the reversible systems tried here (sections 2.3.1 and 2.3.2).
The buffer screen provided initial evidence that the buffer conditions can affect the response of the chip (both non-specific and specific interactions). Further screening for activity under various conditions (including different detergents) may be fruitful. A PBP3 from a different species could also be tested as the issues here may be unique to PaPBP3.

2.5 Conclusions

An SPR system for PBP3 would be highly beneficial to the study of novel inhibitors or even the interactions of the natural substrate at the PBP transpeptidation site. This work was initiated to provide in-solution biophysical evidence of the interactions of boronates with PBPs (Chapter 5). A direct observation (i.e. no biochemical competition) technique would be very useful in the study of target mechanisms of resistance in PBPs (Chapter 4). The results of feasibility studies for the design of a PaPBP3 SPR system, were insufficient to justify continuation of this work. A reliable response could not be obtained using any of the analytes tested. An ideal system should corroborate biochemical data and demonstrate specific, readily dissociating (route B) binding.

This work was made more challenging by a lack of convenient, reversible, high affinity, positive controls for the ligand-analyte interaction. Initial attempts to validate the system using β-lactams in a route A regenerating system by two methods (CAPture oligonucleotide denaturation (2.3.1) and thiol reduction (2.3.2)) were unsuccessful. Using β-lactams (including nitrocefin) as controls in a non-regenerating or route B regenerating system did not provide necessary validation. Work with a benzoxaborole (13) was partially successful but demonstrated large amounts of non-specific interaction, which obscured the specific signal. The relatively weak affinities of the benzoxaboroles (~80 μM) makes them a poor ligand for validation of the system. Recent reports demonstrate that SPR can be a useful tool in the context of PBPs, and further optimisation of the system may find more applications.
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Chapter 3. Design and Optimisation of a High-Throughput Antimicrobial Screening Platform

3.1 Introduction

One of the potential avenues to fill the presently empty antibiotic pipeline is the use of whole cell screening (Chapter 1). The smaller companies now responsible for the majority of antibiotic development \(^1\) are typically less likely to have significant biological testing capacity in house and instead outsource this work to specialist "contract research organisations", such as Warwick Antimicrobial Screening Facility \(^2\).

Warwick Antimicrobial Screening Facility provides an array of microbiological services to academic and industrial clients for early stage drug development run by an experienced, UK NEQAS accredited team. Most work is done manually following Clinical & Laboratory Standards Institute (CLSI) protocols \(^3\), which is possible for fewer than 50 compounds, but becomes increasingly expensive and time consuming for larger libraries. In order for larger libraries to be screened, development of automated methods was required.

In developing higher throughput low volume antimicrobial screens, requiring significantly less compound than traditional MIC methodology, we hope to facilitate the early stage development of novel antibiotics. The CO-ADD platform at the University of Queensland, Australia is driven by similar principles and even offers free screening in the hope of “emptying the drawers” and finding antimicrobial hits in compound libraries that otherwise would never have been screened against bacteria \(^4\).

We therefore aimed to create a highly automated assay (to minimise costs) with quick results turnaround and low sample volume. By minimising sample consumption, antimicrobial activity data can be included in an early stage in the discovery cycle, placing whole cell activity data at the centre of discovery efforts. Additionally, by miniaturising the assay consumable costs are lowered and throughput is increased
because more wells can be run on each plate. To create this platform it was necessary to optimise the bacterial growth rates within the system, create methods for liquid handling as well as develop and validate data processing pipelines to determine the minimum inhibitory concentration (MIC) of a given compound with minimal human intervention.

3.2 Methods

3.2.1 Antimicrobial Assays

The assay was run in 80 µl volume in 384-well black-walled, clear bottom plates (Greiner Bio-One) at 37 °C and the plates were prepared and analysed in a Tecan Freedom EVO liquid handling robot with an 8 channel “LiHa” liquid handling arm, a 96 multichannel arm “MCA 96”, a robotic manipulator arm “RoMa”, a cooling plate, an incubator and a plate reader (Tecan Infinite M200Pro) 5.

Source plates, separately containing inoculating media and antibiotic stocks were placed on carriers on the deck of the robot at the start of the run.
When testing organisms other than E. coli ΔToIC, a different plate reader (Tecan Spark 10M) in a biosafety level 2 laboratory was used.

3.2.2 Inocula

Bacteria were recovered from glycerol stocks (kept at -80 °C), and spread onto a petri dish containing lysogeny (LB) agar (except for N. gonorrhoeae, which was grown on chocolate agar in a 5 % (v/v) CO₂-enriched environment) then allowed to grow overnight at 37 °C. This dish was kept at 4 °C for up to 1 week. The inoculum was prepared by diluting bacteria collected from the dish by a cotton wool swab into phosphate buffered saline (137 mM NaCl, 2.7 mM KCl, 10 mM phosphate buffer pH 7.4) to a concentration of 1.5 x 10⁸ CFU/ml, as determined by comparison with a 0.5 McFarland standard. This stock was then diluted into 2 x cation-adjusted Mueller-Hinton broth 2 (caMHB, Merck) to a concentration of 9.2 x 10⁶ CFU/ml and pipetted into 2 ml 96-well blocks (Brand). All but three wells of the block contained inoculum whilst the remaining wells contained 2 x caMHB for the negative control. This was prepared
immediately before the initiation of the experiment within the robot. Whilst in the liquid handling robot, the inoculum was placed on a cooled carrier to maintain its temperature at 4°C.

3.2.3 Antibiotics Stocks

Antibiotic stocks were prepared at 4 x the desired final concentration (e.g. 512 µg/ml for a highest final concentration of 128 µg/ml) and manually pipetted into 96 well plates (Falcon). The bottom row of the 96 well plate is reserved for controls: 2 containing only water for injection into the positive and negative controls and 2 wells for known antibiotic controls (also at 4 x the desired final concentration) (Figure 3.1). Protocols for running the liquid handling system were written and operated by Dr. Sarah Bennett. Briefly, all the wells of a 384 well ‘destination’ plate were filled with 40 ul of sterile milliQ water using the MCA and 40 µl of each compound was pipetted using the LiHa, from the source plate into separate wells in the “destination” plate. The wells were mixed using aspirate and dispense commands, followed by a 40ul transfer into a new well containing 40 µl of water, thereby initiating a serial dilution that continued across the plate. At the last dilution step, 40 µl of diluted compound was removed and discarded. Once the inhibitor was plated out, 40 µl of inoculum was added to each well using the MCA. The RoMa then fitted a clear lid (Greiner microplate lid sterile) to the plate and transferred the plate to a plate reader (Tecan Infinite M200Pro). The plate was incubated at 37 °C, and OD₆₀₀ measurements were taken at 600 nm every 10 minutes for 4-8 hours.

3.2.4 Gonococcal (GC) media

For the GC media brain-heart infusion broth (Merck) was prepared, then supplemented with 5 % (v/v) horse blood (E & O Laboratories) and autoclaved. After cooling, the supernatant was poured off and filtered through a 0.22 µm sterile filter (Merck) before use.
3.3 Results and Discussion

3.3.1 Optimisation of Growth Rates

In order to increase throughput of the assay and minimise the amount of incubator space required, the microbroth serial dilution method of measuring growth inhibition, which is typically undertaken in 96 well plates with 100 μl total volume was miniaturised into a 384-well, 80μl format. Additionally, by moving from a typical endpoint of 18-24 hours to an endpoint of less than 8 hours we are able to increase throughput of automated runs. However, we needed to ensure growth was sufficient in this period to differentiate between cells affected and unaffected by the treatment.

Investigation of the literature shows various methods for monitoring growth implemented in phenotypic screens have been used previously. : i) visual inspection for turbidity, as used in low throughput screens, ii) commercial luminescence detection kits, iii) fluorescent staining, iv) strains mutated to induce bioluminescence or fluorescence, v) resazurin to detect changes in the redox environment, vi) changes
in turbidity detected by measurements at 600 nm \(^{14-16}\).

Whilst staining, fluorescent and luminescent methods may increase sensitivity as these methods amplify the signal, they also risk possible chances for compound interference in the read out and false discovery. Additionally genetic modifications to cells cannot be carried out on clinical strains. Therefore, we opted to use turbidity as the read out for this assay.

A validated mutant of *Escherichia coli* with a knock out of the transmembrane efflux pump *TolC* gene (*E. coli* Δ*TolC*) was selected for use during the development phase of this work as it is a biosafety level 1 organism and it is a useful tool for screening since it reduces the permeability barrier (see thesis Introduction), making hit discovery more likely \(^{17}\).

Figure 3.2 shows the growth curves produced when inoculating wells with different initial concentrations of *E. coli* Δ*TolC* cells in 80 µl of media in a 384 well plate. These growth curves were used to select an appropriate time point for measuring the endpoint in subsequent runs as well as a dilution of bacterial cells for the inoculum. As expected, end point change in optical density correlates with concentration of inoculum due to the binary growth of bacteria (Figure 3.2b). Using this data, an inoculum concentration of 4.6 x10\(^{6}\) CFU/ml was chosen as this gives good balance between sufficient change in optical density to detect growth within a short time frame (Figure 3.2b), and with relatively low variance between repeats (Figure 3.2c). After 240 minutes bacteria are in the middle of the growth phase.
Figure 3.2. Optimisation of Growth Conditions. (a) Overlay of growth curves of *E. coli* ΔToIC over 8 hours. Colours indicate the concentration of cells in the inoculum (see key), the negative control (black) has no inoculation. The red lines show the threshold for a change in optical density (OD_{600}) of 0.05 (horizontal) and 240 minutes (vertical). The intersections of these lines and the growth curves are shown in (b) and (c) respectively. b) For each concentration of inoculum, time for the *E. coli* growth curve to have a change in OD_{600} of 0.05 is shown, darker markers show more data points at that position. (c) Similarly, the change in OD_{600} after 240 minutes is shown for each concentration of *E. coli* inoculum. In (b) and (c) the x axis is a log_{2} scale.

Comparison of the turbidity of a suspension of bacterial cells to a 0.5 McFarland standard is used to estimate the concentration of bacteria. Due to the potential sensitivity of the method to errors caused by variations in estimations of turbidity when making up the inoculum, replicates with the same starting concentration of inocula were used to confirm the reproducibility of the growth curves. Eight different inocula ("replicates") with bacteria at a concentration of 4.6 x 10^{6} CFU/ml, as determined by comparison with a McFarland standard 8 different times, were used to initiate grow in
48 wells each. The data show that starting concentration is responsible for a small amount of variation between “replicates”, but that variation within each “replicate” is consistent (Figure 3.3). When running screens, all wells were inoculated with bacteria made up as a single inoculum, which will eliminate variation between “replicates” but variation within “replicates” is likely to have similar distribution. Variation between runs can be standardised by comparison of the positive controls.

![Graph showing variation due to starting inoculant](image)

**Figure 3.3. Variation due to starting inoculant.** (a) Overlay of growth curves of *E. coli* ΔTolC over 8 hours, darker grey indicates more data at that position. The red line is set at 240 minutes. (b) The change in turbidity for each curve in (a) is plotted against the “replicate”, where each replicate was made up by diluting bacterial cells to a concentration estimated to be equal to 4.6 x 10⁶ CFU/ml, as determined by comparison with a McFarland standard.

### 3.3.2 Z-factor

The Z-factor is a parameter which can be used to assess the quality of an assay. It is a measure of the signal to noise ratio of an assay, taking into account the standard deviation and means of the positive and negative controls of the assay. It is defined as in equation 3.1:

\[
Z = 1 - \frac{3\sigma_p + 3\sigma_n}{|\mu_p - \mu_n|} \tag{3.1}
\]
where $\sigma$ is the standard deviation and $\mu$ is the mean of the positive ($\mu_p$) and negative ($\mu_n$) controls.

The Z-factor was found to be 0.75 for this assay (E. coli ΔTolC growth in 4 hours with automated liquid handling), which is defined as “an excellent assay” by Zhang et al. 18, in terms of its ability to distinguish between growth and no growth.

### 3.3.3 Robotic Liquid Handling

These initial investigations showed that bacteria tolerated the low volume well and grew reproducibly. We then developed automated methods to measure and determine MICs. We envisaged two example use cases for the platform: a prototypical 12 concentration MIC with two-fold concentration steps, and a minituirised MIC test in which only four concentrations of compound were investigated, which allows for increased throughput. Protocols for both these methods were developed.

![Diagram of liquid handling methods](image)

**Figure 3.4. Liquid handling methods to transfer solutions between 96 well and 384 well plates and generate concentration series.** (a) Linear two-fold serial dilution across the plate. (b) Two-fold serial dilution for four concentrations in square grids. In both (a) and (b) lightness of the colour indicates increasing dilution. Two-fold dilution is achieved by mixture of equal volumes (40 µl) of compound-containing solution and water (c) Inoculating the wells with a constant concentration of bacterial cells. The operation shown in (a) can only be achieved by the 8 channel “LiHa” arm, whilst the operation shown in (b) and (c) can be carried out by the MCA 96 arm.

The protocol takes two “source” plates: one loaded with bacterial inoculum and one with test compounds at a single concentration (Figure 3.1). Four controls are included on every “destination” plate: negative, which has no inocula, positive which has no
inhibitor, and two antibiotics of known MIC to ensure consistency of dilution.

The contents of these plates are then distributed to the “destination” plates by the robotic liquid handling system before the assay can begin. The Tecan Freedom EVO liquid handling robot has 2 liquid handling heads: a 8 channel “LiHa” and a 96 multichannel arm “MCA 96”, and a robotic manipulator arm “RoMa”.

The robot can mimic a hand-held 8-well multichannel pipette and quickly dilute a compound across the long axis of a plate, as shown in Figure 3.4a. Alternatively, when test only 4 compound concentrations, the MCA 96 arm can be used to dilute the compounds in a 2 x 2 grid (Figure 3.4b). Due to the high multiplicity of the arm, this is very rapid. Following dilution of the compounds, the wells are inoculated with bacteria using the MCA 96 (Figure 3.4c), initiating the experiment.

The RoMa arm transfers the completed plates to a 37 °C incubator, and then moves the plate between the incubators and the plate reader at the reading intervals. The data is then outputted ready for processing.

3.3.4 DMSO

High-throughput compound libraries are often solvated in DMSO, so determining the sensitivity of the assay to DMSO is important to ensure any effect observed is not due to DMSO. The sensitivity of \textit{E. coli} ΔToIC to DMSO is shown in Figure 3.5. The results show total inhibition above 10 % but significant inhibition is not seen for concentrations below 2 % DMSO. A 2 % limit on DMSO concentration when diluting compounds was used in all subsequent work.
3.3.5 Data Processing

3.3.5.1 Workflow

Figure 3.6 shows the workflow used to process the data. It requires three inputs (shown as triangular boxes): the growth data (taken from a microplate reader); the names of the compounds and the “Maps”. Maps are directory files used to describe the layout of the destination plate (equivalent to the matching of colours in Figure 3.1. They increase the flexibility of the system since these can be easily changed to meet the desired number of concentrations of compounds to be tested, different arrangements of compounds on the destination plate and the types of controls used.
Figure 3.6. Overview of the workflows used to prepare a report of a set of MICs. Each box represents a process (typically run by an individual script) required to process the inputted data into the final report. Operations in white are those carried out by the code, in blue are client-facing operations, in red are experimental procedures. FIC(I): fractional inhibitory concentration (index).
One of the complications of the data processing is that the source plates are 96-well plates, whilst the destination plates are 384-well. 96-well plates are used due to their higher capacity and easier handling (pipettes and the robotic arms have tips spaced at these dimensions). The solution is the use of Maps, which correlate the positions in the two plates and allow the growth data to be stored in a database alongside information on the compound concentration and name so that this can correctly be outputted together in the report.

A series of Matlab scripts then prepares the data into a format in which the MIC can be determined. Essential scripts are those that reformat the data; calculate the growth over 4 hours; fit models to the data; and generate reports.

3.3.5.2 Determining the MIC

When determining an MIC using manual methods, the MIC is defined as the last well in which no turbidity can be detected by eye following incubation of the test compound. This is possible because over 16-24 hours an antimicrobial compound will typically prevent visible growth at its MIC, creating a clearly defined “cliff” of efficacy. Preliminary experiments with different inhibitors using short (4 hours) growth times revealed a number of possible shapes of the dose-response curves (Figure 3.7). Some inhibitors, such as polymyxin B (Figure 3.7a), produce a clear “cliff” in response at a critical threshold of activity, but others, such as tetracycline (Figure 3.7b) have log-linear responses because weak growth is still possible. Models to fit the curves to find a value for the MIC were investigated.
Figure 3.7. Dose responses of *E. coli* ΔToIC to different compounds. (a) Polymyxin B (b) Tetracycline (c) a novel compound (undisclosed). Each example shows a different slope of the curve. (c) is fitted with 4 models, Models A-D in Table 3.1: **Model A** (orange dashed line) the threshold for change in absorbance of 50% of the positive control; **Model B** (purple dashed line) the threshold for change in absorbance of 10% of the positive control; **Model C** (green solid line) and **Model D** (blue solid line). The MICs determined using each model for this compound are shown in Table 3.1. Grey circles show the change in absorbance over 4 hours for each concentration of compound. In (a) and (b) each circle indicates a repeat, darker circles indicating multiple (2-4) data points overlapping. This compound was determined to have an MIC of 16 μg/ml by methods following CLSI standards.

Four methods were tested for their ability to accurately determine a compound’s MIC from the dose-response curves (Figure 3.7c and Table 3.1). Models A and B determine
the MIC using a fixed threshold (Threshold) in optical density relative to the average of the positive and negative controls for that screen (equation 3.2). For A the threshold is 50 % (k = 0.5), for B the threshold is 10 % (k = 0.1). The MIC is simply found as the first concentration of inhibitor that gives a change in optical density of less than Threshold.

\[ \text{Threshold} = k \times (\text{positive control} - \text{negative control}) + \text{nag} \quad (3.2) \]

Models C and D use a 4-term sigmoidal curves to fit the data (equation 3.3) \(^{20}\), using the "fittype" curve fitting function in Matlab \(^{24}\).

\[ y = \text{bottom} + \frac{\text{top} - \text{bottom}}{1 + \left( \frac{x}{\text{EC}_{50}} \right)^{\text{slope}}} \quad (3.3) \]

Where \( y \) is the response and \( x \) is the log\(_2\) of the concentration of the inhibitor. C constrains the top and bottom terms of equation 3.3 to the change in OD\(_{600\text{ nm}}\) of the positive and negative controls of the experiment, whilst D allows these terms to be dynamically fitted but constrains the slope term to >2, which prevents the data being fit to too shallow a curve, which was shown to lead to very poor fits during evaluative fitting.

The EC\(_{90}\), defined as the effective concentration at which 90 % of the growth is inhibited, is then found (equation 3.4). For C, the 10 % threshold (Thres10) is defined using equation 3.2 with \( k = 0.1 \). Thres10 is substituted into equation 3.4 to find a value for EC\(_{90}\). This can be graphically visualised as the intersect of the green line (C) and the purple dashed line (B) in Figure 3.7c. For D, after fitting the data with the sigmoid from equation 3.3, Thres10 is simply defined relative to the top and bottom terms found by the fit (equation 3.5) and then the EC\(_{90}\) found using equation 3.4.

In both cases, after finding the EC\(_{90}\), the MIC is reported as the lowest concentration tested which was greater than the EC\(_{90}\).
\[ EC_{90} = EC_{50} \times \sqrt[\text{slope}]{\frac{\text{top-bottom}}{\text{Thres10-bottom}}} - 1 \]  \hspace{1cm} (3.4)

\[ \text{Thres10} = k \times (\text{top} - \text{bottom}) + \text{bottom} \]  \hspace{1cm} (3.5)

When analysing the results of high-throughput screens, ensuring the correct fitting of the data for every compound is important. The \( R^2 \) (coefficient of determination) term is determined by the Matlab \textit{fit} function and describes the goodness of fit of the curve to the data, where 1 is a perfect fit. This value can be used to flag data sets in which the curve fits the data very poorly, so they can be manually inspected. Compounds in which all concentrations are far above or below the MIC do not show sigmoidal curves, but can be instead compared to the positive and negative controls and the MIC reported as greater than the highest concentration tested or less than the lowest concentration tested.

<table>
<thead>
<tr>
<th>Model</th>
<th>Constraints</th>
<th>( EC_{90} ) (µg/ml)</th>
<th>MIC Reported (µg/ml)</th>
<th>Results from the test set</th>
<th>Results within 4 fold of standardized result (%)(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A  Threshold</td>
<td>N/A</td>
<td>N/A</td>
<td>4</td>
<td></td>
<td>73.2</td>
</tr>
<tr>
<td>(50%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B  Threshold</td>
<td>N/A</td>
<td>N/A</td>
<td>128</td>
<td></td>
<td>79.8</td>
</tr>
<tr>
<td>(10%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C  Sigmoid</td>
<td>\textit{top} and \textit{bottom}; see text</td>
<td>19.5</td>
<td>32</td>
<td></td>
<td>85.7</td>
</tr>
<tr>
<td>equation 3.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D  Sigmoid</td>
<td>\textit{slope} &gt;2</td>
<td>8.5</td>
<td>16</td>
<td></td>
<td>76.8</td>
</tr>
<tr>
<td>equation 3.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3.1. Details of the models investigated for determining the MIC in the automated pipeline N/A: not applicable. \(^a\) The percentage of results indicated by each model to be within 4 fold of the result from a standardised test is listed. This is used as a measure of accuracy of the model on a novel data set. See Figure 3.8.

These models were used to determine the MIC for unseen inhibition data of novel compounds. 51 novel compounds (provided by a collaborator), along with 5 known
antibiotics (amikacin, ticarcillin, minocycline, rifampin, clarithromycin) were tested in triplicate and each of the 168 runs was analysed individually (Figure 3.8) and compared with the results determined by a standardised, low throughput method (following CLSI protocols \(^2\)). As expected, the model A overestimates compound potency relative to the standardised data set (Figure 3.8a). The reason for this can easily be understood from Figure 3.7c as concentrations which give weak inhibition are designated as the MIC. It is likely this method would be more accurate for compounds which generate very steep inhibition curves (large slope term), of the type observed with polymyxin B (Figure 3.7a). A comparison of the histograms of the B and C demonstrates the benefit of using a sigmoidal fit to find \(EC_{90}\) rather than using only a simple threshold. Whilst the distributions of Figure 3.8b and Figure 3.8c are generally similar, B produces a left handed tail which is absent from C. As illustrated in Figure 3.7c, this occurs when points lie close to, but do not cross the 10% threshold, leading to the model underestimating the compounds efficacy. Finally D, gives results broadly similar to C but has many outliers, these occur due to poor fitting of the data, leading to a major misinterpretation of the result.

One case where model D is particularly useful is to fit curves well for compounds such as aztreonam. Aztreonam is not biocidal, even at concentrations many fold above the MIC, due to its specific inhibition of PBP3, which prevents division but allows cellular elongation \(^{19-21}\). This means that even at high concentrations of inhibitors there is a non-zero change in absorbance over the course of the experiment. The novel compound in Figure 3.7c is displaying this kind of behaviour, as demonstrated by the deviation between the green (C) and blue (D) at concentrations greater than 16 \(\mu g/ml\). Allowing variation in bottom helps fit the data more accurately in this case.
Figure 3.8. Results generated by each model (A-D) when analysing MICs of novel compounds. Models: (a) A; (b) B; (c) C; (d) D. The automated method was used to generate inhibition curves for 56 compounds in triplicate. The MICs calculated by each model for each individual replicate were compared to the MIC for that compound determined by standardised methods, expressed as a ratio (“Ratio of MICs”). This ratio is logged (base 2) and plotted against the frequency that result was observed by that model. 0 on the x-axis indicates the model generated the same result as the standardised test, negative values indicate that result from the standardised method was lower (more effective inhibition) than the result generated by the model, with the opposite being true for positive results. Any results which were recorded as less than the lowest value tested were treated as being 8-fold lower than the lowest concentration tested, similarly for when no inhibition was observed (treated as 8-fold higher. This is likely to be overly conservative.

In the early stages of a screening program, a result needs to be rapidly returned to give an indication of the potential of a novel compound. The exact measure of the MIC may
not be important in the early stages. For this purpose scripts were also written into the pipeline which allow just four concentrations to be tested for each compound. This allows a more than three-fold increase in the throughput as dilution of the type shown in Figure 3.4b can be used which gives a further improvement in throughput. Concentrations can be chosen to give a categorical description; e.g. a compound with an MIC of 64 μg/ml or greater may be categorised as ineffective, whilst one with an MIC of 8 μg/ml may be worth follow-up.

3.3.6 Screening Boronates

The liquid handling platform was used to screen a small library (49 compounds) of boron-containing compounds (those in Chapter 5 as well as a small collection of phenylboronic acids) against E. coli ΔTolC (Figure 3.9). Few active compound hits were identified, with the exception of tavaborole, an antifungal agent \(^{22,23}\), which was found to have an MIC of 2 μg/ml.

![Figure 3.9. Histogram of MICs of boron-containing compounds screened. 49 boron-containing compounds were screened in duplicate and their MIC determined by model C.](image-url)
3.3.7 Synergy Studies

We have additionally developed methods to investigate compound synergy using the automated platform. Synergy occurs between two antimicrobial compounds when their activity together is greater than the sum of their parts \(^{26}\). Due to the number of possible combinations of compounds, even from a small library, these studies are very labour-intensive when done by hand, making them an excellent candidate for automation. Protocols were established for the liquid handling system which allows both compounds to be diluted separately, before they are combined to a 2D gradient of concentration. Each compound is diluted two-fold in perpendicular directions and then 20 \(\mu\)l of each is added to the plate before inoculation. Protocols were designed to allow either 4 x 4 well grids (i.e. 4 concentrations of each of the inhibitors) or 11 x 7 well grids. As shown in Figure 3.6, the data processing pipeline is similar, but also includes the optional addition of calculations of the fractional inhibitory concentration index (FICI) \(^{27}\) (equation 3.6), a measure of the size of the synergistic effect of a combination. This is found after the determination of MIC of each of the two compounds (\(i\) and \(j\)).

Examples of the 4 x 4 well “checkboards”: grids of two concentrations of inhibitors - are shown in Figure 3.10. Beneficial interaction of the \(\beta\)-lactamase inhibitor clavulanic acid and \(\beta\)-lactams is well characterised for \(\beta\)-lactamase-expressing strains and this combination is used in the clinic \(^{24-26}\). In this case it is not possible to calculate the FICI because clavulanic acid does not have antimicrobial effect alone (no MIC, term for equation 3.6).

\[
\text{FICI} = FIC_i + FIC_j = \frac{[i]}{MIC_i} + \frac{[j]}{MIC_j}
\]  
(3.6)
Figure 3.10. Synergy Studies. 4 x 4 “checkerboards” showing interactions between clavulanic acid and a β-lactam when inhibiting the growth of β-lactamase (CTX-M-15) expressing E.coli strain NCTC 13353. (a) Penicillin G (b) Aztreonam. Note that the MIC of aztreonam is much lower than penicillin G so the concentrations tested are also lowered. Shading in each cell is greyscale with black indicating lack of growth and white indicating the same amount of growth as the positive control.

3.3.8 Growth of Other Organisms

With the protocols established for a basic dose response curve against E. coli ΔToIC, we aimed to expand the capabilities of the platform. Screening compounds against E. coli ΔToIC mutant is informative for certain projects, but novel compounds need to be effective against clinically relevant, priority organisms. The growth of these needed to be validated in the low volume, high throughput format (Figure 3.11). Currently the liquid handling robot is not in a Biological Safety Level 2 environment, so inoculation must be carried out manually in the appropriate environment. Additionally, P. aeruginosa, K. pneumoniae, A. baumannii and N. gonorrhoeae required higher initial concentrations of inoculum to ensure growth in the timescale, and in some cases different media. N. gonorrhoeae grew poorly and can often fail to grow all together, and may require further optimisation of conditions for this to be sufficiently robust for high throughput screening (Figure 3.11). One interesting observation from the growth curves of Salmonella typhimurium (Figure 3.11b), Enterobacter cloacae (Figure 3.11c) and to a lesser extent E.coli ΔToIC (Figure 3.3a) is the ‘jagged’ appearance of the growth curves in the exponential phase, compared to the smooth exponential growth of e.g. E. coli NCTC 25922 (Figure 3.11a). This is termed “diauxic growth”, and can be caused by sequential substrate utilisation with the different substrates allowing for different growth rates.
Figure 3.11. Growth curves of clinically relevant species. Growth curves for 10 species of bacteria are shown: (a) *E. coli* NCTC 25922, (b) *S. typhimurium* ATCC 19585, (c) *E. cloacae* NCTC 13405, (d) *Enterococcus faecalis* ATCC 47077, (e) *Staphylococcus aureus* ATCC 29213, (f) *P. aeruginosa* PAO1, (g) *K. pneumoniae* ATCC 700603, (h) *A. baumannii* NCTC 19606, (i) *P. aeruginosa* NCTC 13437 (j) *N. gonorrhoeae* ATCC 49226. Different growth media is required for the different species: (a-e) calcium adjusted Mueller Hinton broth, (f & g) Brain heart infusion (h-j) specialised GC media (section 3.2.4). For *A. baumannii* NCTC 19606, *P. aeruginosa* NCTC 13437 and *N. gonorrhoeae* (h-j) a 5% CO₂-enhanced environment (generated using the plate reader) was used to further stimulate growth as well as the use of GC media (despite the name this media also stimulates the growth of other bacteria). The concentration of inoculating bacteria is indicated by colour, as shown in the key. Note that the time scale for (j) is in hours, due to the slow growth of *N. gonorrhoeae*. 
3.3.9 Case Studies

Several academic groups have also developed automated and semi-automated screens for antimicrobial discovery, as well as the screening of large chemical libraries. Their screens have had different objectives, including drug repurposing, “unconventional” screening and discovery of synergistic partnerships, and investigation of cellular permeability. Examples of recent large scale screening attempts by academic groups are given below, demonstrating the potential uses of this platform.

3.3.9.1 Repurposing Screens

One accessible source of chemical matter is the commercially available libraries of known and approved compounds \(^{29-32}\), which can be used to search for new uses for old drugs, termed “repurposing” \(^{33}\). The advantage of repurposing is its significantly lower cost since the expensive safety stages of the drug discovery process have already been carried out, allowing a quick and cheap route to the clinic for a repurposed compound \(^{34}\), compared with the estimated $1.7 billion cost of a novel compound \(^{35}\).

An extensive screen by the American National Institutes of Health screened 5,170 known compounds in an attempt to find inhibitors of \(K.\ pneumoniae\). This screen found only 25 hits (0.5%) and follow-up work showing that the maximum concentration possible in the blood for these compounds was too low for them to be effective as antimicrobials \(^{14}\). Disappointingly, other recent repurposing attempts \(^{6,7,9}\) have similarly often only identified compounds with known antimicrobial activity (whether that be antimalarial, antiviral or antifungal). Potential reasons for the low hit discovery rate are discussed in the thesis Introduction.

3.3.9.2 Synergy Screens

Automated screening can also be used to search for novel synergistic partners of known antibiotics, in order to bypass mechanisms of resistance \(^{36-38}\) or unlock cryptic activity. Eric Brown, Gerard Wright and colleagues at McMaster Canada have implemented a semi-automatic platform in a series of “unconventional” screens \(^{15,16,39,40}\), which aim to leverage observations such as the increased sensitivity of \(E. coli\) to vancomycin when cold stressed to find novel inhibitors of outer membrane proteins and
the ability of loperamide (Imodium) to potentiate the antimicrobial potency of tetracyclines against multi-drug resistant *P. aeruginosa* 39. These screens can also help identify targets which have yet to be challenged by antibiotics 16. For example, observations of the β-lactam-potentiating activity of tunicamycin in methicillin-resistant *Staphylococcus aureus* 41 has lead to several attempts to find more drug-like inhibitors of the target of the highly cytotoxic tunicamycin, the wall teichoic acid synthesis machinery 15,42-44. The identification of compounds such as tarocins 43 and the anti-platelet drug ticlopidine 15 which lack antimicrobial activity but bypass resistance to β-lactams demonstrates the importance of synergy studies in combating antibiotic resistance.

Synergistic studies can also be used to find drug combinations that lower the clinical dose required and avoid the effects of resistance to a single therapy. Sun et al. used their high throughput “HI/GA” platform to screen large numbers of combinations of 2 or 3 known antibiotics to find cocktails that could be given to patients with multiple drug resistance infections where single therapy is failing 14. Without automation screening the many potential combinations would be prohibitive.

### 3.4 Conclusion

We have developed and validated a series of methods that can allow antimicrobial screening data to be obtained quickly and cheaply, with little compound usage. The accuracy of results is within the tolerance needed for a first round screen, as successful candidates are likely to be repeated via standardised methodologies. This platform will be used for projects within our group as well as being offered as a commercial service to small and medium drug discovery companies via Warwick Antimicrobial Facility.

### 3.5 References

20 Spratt BG. Distinct penicillin binding proteins involved in the division, elongation,


Chapter 4. Target-Mediated Resistance in PBP3: Role of Loop Flexibility

Chapter 4.S (p170-173) contains additional crystal views and data referred to in this chapter

4.1 Introduction

In response to treatment with antibiotics, bacteria have developed a number of mechanisms of resistance (Chapter 1) \(^1\). Target mediated resistance occurs when mutations are introduced into a protein which affects drug-protein interactions, and protects the bacterium from the drug’s effects. Mutations could introduce resistance to β-lactams through a number of mechanisms: (i) reduction of the pre-acyl affinity of the β-lactam for the active site, (ii) reduction of the acylation rate (covalent bond formation or (iii) increase of the de-acylation rate, leading to faster clearance of the drug from the protein \(^2\). Change in the protein sequence at just a few specific points is sufficient to induce resistance via one of these mechanisms \(^2-7\).

4.1.1 Mapping the Locations of Mutations

We first undertook a literature search for clinically relevant mutations that occur within class B PBPs of gram-negative species and mapped them onto new and improved high-resolution crystal structures \(^6\) (Table 4.1 and Figure 4.1). There is extensive literature on the presence of point mutations causing β-lactam-resistance in \textit{H. influenzae} PBP3 (HiPBP3, mutations prefixed by \(^{16}\) \(^3,8-19\)) and in \textit{N. gonorrhoeae} PBP2 (NgPBP2, mutations prefixed by \(^{16}\) \(^2,4,5,20-25\)), whereas only a few mutations have been reported in \textit{P. aeruginosa} PBP3 (PaPBP, mutations prefixed by \(^{Pa}\) \(^26,27\)) or \textit{E. coli} PBP3 (EcPBP, mutations prefixed by \(^{Ec}\) \(^28-30\)) (Table 4.1). By plotting the mutations onto crystal structures (Figure 4.1), it was clear that they cluster primarily in loops adjacent to the active site cleft of class B PBPs (Figure 1.4 for an overview of the protein). Loops are defined as stretches of residues between secondary structure elements, but they are increasingly understood to have important roles within proteins beyond simple connectivity \(^31\). The four loops
identified by mutation clustering were the β2 region (comprising of a series of loops between the short β strands β2a-d), the α10–β3 loop, the β3–β4 loop and the β5–α11 loop (Table 4.1).

<table>
<thead>
<tr>
<th></th>
<th>PaPBP3</th>
<th>HiPBP3</th>
<th>NgPBP2</th>
<th>EcPBP3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>β2 region</strong></td>
<td>–</td>
<td>H^{i} Ser357Asn</td>
<td>N^{β}_{Asp346a} insertion</td>
<td>E^{βc}(T/R)IPY,</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ec^{βc}YRI(N/K) insertions (around Tyr334)</td>
</tr>
<tr>
<td><strong>α10–β3</strong></td>
<td>–</td>
<td>H^{i} Arg501Leu,</td>
<td></td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>H^{i} Ala502Val,</td>
<td></td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>H^{i} Val511Ala</td>
<td></td>
<td>–</td>
</tr>
<tr>
<td><strong>β3–β4</strong></td>
<td>Pa^{a} Arg504Cys/His</td>
<td>H^{i} Arg517His,</td>
<td>N^{β}_{Phe504Leu},</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>H^{i} Asn526Lys</td>
<td></td>
<td>N^{β}_{Ala510Val},</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>N^{β}_{Asn512Tyr},</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>N^{β}_{Ala516Gly}</td>
</tr>
<tr>
<td><strong>β5–α11</strong></td>
<td>Pa^{a} Pro527Ser/Thr,</td>
<td>H^{i} Gly555Glu,</td>
<td>N^{β}_{Gly542Ser},</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>H^{i} Tyr557His</td>
<td></td>
<td>N^{β}_{Gly545Ser},</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>N^{β}_{Pro551Ser/Leu}</td>
</tr>
</tbody>
</table>

**Table 4.1.** Mutants observed in the class B PBPs of three species of gram-negative pathogens cluster in various regions. Modified from Bellini et al. 6.

### 4.1.2 β2 region

Simple comparisons of the crystal structures of wild type and mutant proteins can give an initial indication of the role of a particular mutation. To see if the presence of the N^{β}_{Asp346a} on the β2 region had an effect on its conformation, we crystallised and solved a novel structure (NgTP2^{βIR-6140}, PDB: 6HZJ) 6 of an NgPBP2 mutant with 5 clinical mutations (Pro551Ser + Phe504Leu + Ala510Val+ Ala516Gly + the insertion Asp346a) from *N. gonorrhoeae* strain FA6140 21. The conformation of the β2 region changes significantly compared to a lower resolution structure of the same protein (NgTP2^{βIR-6140}, PDB: 4U3T) 20 or the wild type protein (PDB: 3EQU) 4. Insertions of 4 residues into this loop have been repeatedly reported in carbapenem-expressing *E. coli* strains 28–30. The exact sequence inserted varies between strains but the effect is likely to be a rearrangement of the β2 loop. In both EcPBP3 and NgPBP2 as this loop sits above the active site cleft, these changes may restrict active site accessibility.
Figure 4.1. Plotting locations of clinical mutants on the class B transpeptidase domain. (a) Locations of the mutations listed in Table 4.1, plotted onto a NgPBP2 crystal structure (PDB: 6HZJ). Mutations are coloured by the originating class B PBP: PaPBP (cyan), HiPBP3 (orange), NgPBP2 (pink) and EcPBP3 (green). Labelled loops are highlighted in black. (b) Locations of mutations appear to cluster within 4 loops: the β2 region (in NgPBP2: residues 324-353), the α10-β3 loop (in NgPBP2: residues 477-494), the β3-β4 loop (in NgPBP2: residues 501-513) and the β5-α11 (in NgPBP2: residues 537-548).
4.1.3 β3-β4 loop

The β3-β4 loop is a flexible region where clinical mutations can arise. In the four available PaPBP3 wild type apo structures (PDB: 3OC2, 3PBN, 6HR4 and 6HZR) this loop is unresolved, likely due to conformational flexibility. The binding of piperacillin (PDB: 6R3X) or cefoperazone (PDB: 5DF8) appears to constrain the loop and allow it to be fully resolved.

Comparison of the apo and ceftriaxone reacted forms of wildtype NgPBP2 found that the β3-β4 loop was closer to the active site serine in the reacted form. A twisting of the β3 strand accompanied this change. Recent crystallographic evidence in a mutant NgPBP2 suggests that the effect of mutations in the β3-β4 region may be to prevent the contraction of the β3-β4 loop upon ceftriaxone binding, as the apo and ceftriaxone reacted forms of a clinical mutant of this protein had little structural variation.

A structure of PaPBP3 with the PaArg504Cys mutation appeared to have no structural differences with the wild type, when apo or piperacillin-reacted. The β3-β4 loop is the site of two frequently observed mutations (93 % in one study) in HiPBP3: HiArg517His and HiAsn526Lys. These mutations provide little resistance by themselves but precede the development of further, high resistance, mutations in H. influenzae.

4.1.4 β5-α11 loop

The β5-α11 loop was observed in β-lactam reacted PaPBP3 structures to transition from an outward "open" conformation (PDB: 3PBN and 6HZR) to an inward "closed" conformation upon binding of compounds such as piperacillin (PDB: 6R3X) or aztreonam (PDB: 3PBS) (Figure 4.S2). In the closed conformation residues PaTyr503, PaTyr532 and PaPhe533 form a "hydrophobic wall" against a hydrophobic section of a reacted compound. Mutations that occur in this region (Table 4.1) are primarily to proline or glycine residues which are likely to change the flexibility and mobility of this loop and its ability to do this movement. In NgPBP2, the role of the NgGly545Ser mutation (located at the N-terminal end of the α11 helix) has been
suggested to play a more direct role, as Ser545 was observed to form a hydrogen bond to Thr500 on the β3 strand, resulting in a conformational shift in the position of the ceftriaxone carboxylate group ².

4.1.5 α10-β3 loop

There has been little crystallographic evidence of the flexibility of the α10-β3 loop, but recent boronate-bound PaPBP3 structures show conformational plasticity here (see below).

From the locations of the mutations and the crystallographic conformational flexibility already seen in some of these regions, we hypothesised that these loops have a more important role in PBP3 biology than previously understood ⁶. Mutations responsible for β-lactam resistance may cause their effect by changing the flexibility of the loops and this is what causes changes in the dynamic behaviour of the protein. This chapter investigates loop flexibility using published crystallographic data as well as insight from novel PBP3:boronate complexes and then discusses methods beyond crystallography that could be used to further understand the dynamics of loops in PBP3s.

4.2 Methods

4.2.1 Ensemble refinement

In order to generate the ensemble models of PBP3s, Ensemble refinement (ER) was run using Phenix (Version 1.18.2) ³⁷⁻³⁹. Published structures (PDB: 6HZR and 6HZJ) were used for the input. To optimise the ensemble refinement, different values of the TLS (translation/libration/screw) parameter can be used. This parameter allows for flexibility of subsections of the model ³⁹. Refinement was run with TLS input values of 0.8, 0.9 and 1.0 and for both proteins it was found that 0.8 gave the lowest R factors. The ER led to a smaller Rwork value for the PaPBP3 and NgPBP2 models and a decreased Rfree value for the PaPBP3 model, but essentially unchanged Rfree value for the NgPBP2 model, compared to the deposited structure (Table 4.2). Resultant structures containing all of the conformational models (85 and 50 for the PaPBP3 and NgPBP2 models respectively) were exported and analysed by
MDanalysis\textsuperscript{40,41} and PyMOL (Open-Source PyMOL Molecular Graphics System, Schrodinger, LLC. Version 3.8.2).

4.2.2 RMSDs
The difference in the positions of main chain atoms (C, CA, N and O, following PDB notation) of a given residue was found for each of the conformational models generated by ER, compared to the deposited structure. The value for each residue was found by taking the mean across all the conformational models. The difference was assessed by determination of the root-mean-square deviation (RMSD). RMSD was found using the \texttt{MDAnalysis.analysis.rms} module\textsuperscript{42,43} of MDAnalysis, using custom scripts.

When determining the per-residue RMSD between two structures (e.g. Figure 4.S1), a similar procedure was used, but with only two input structures. Regions in which sections of the model were missing in one of the structures were omitted from analysis.

4.2.3 B factors
Refined B factors were extracted from the published model files by custom scripts in MDAnalysis. The mean of the B-factors of each main chain atom (C, CA, N, O, following PDB notation) of each residue was determined for each model or subunit.

4.2.4 PDBFlex
The PDBFlex server\textsuperscript{44} was used to examine the conformations of PaPBP3 structures in the “4kqrA cluster” which consisted of 38 structures (in the 16\textsuperscript{th} May 2020 PDB release). These correspond to structures with PDB codes: 3OC2, 3OCL, 3OCN, 3PBN, 3PBO, 3PBQ, 3PBS, 3PBT, 4FSF, 4KQO\textsuperscript{*}, 4KQQ\textsuperscript{*}, 4KQR, 4L0L, 4OOL, 4OOM, 4WEJ, 4WEK, 4WEL, 5DF7\textsuperscript{*}, 5DF8\textsuperscript{*}, 5DF9, 6HR4, 6HR6, 6HR9, 6HRZ, 6H1E, 6R3X, 6R42, 6R42, 6UN1, 6UN3, 6VJE and 6VOT, all of which have > 99 \% sequence similarity. Structures marked with an asterix have both A and B subunits, which were considered separately. Only 5 of these are apo structures (PDB codes 3OC2, 6HR4, 3PBN, 6HRZ, 6R40), with the rest being bound to a ligand. The average local root mean squared deviation of the C\textalpha
carbon (RMSD) was exported and plotted against the residue number in Matlab.
Boron-containing crystal structures were crystallised as described in section 5.2 and reference 45.

4.3 Results and Discussion

4.3.1 Assessing Backbone Flexibility with Crystallography

Crystallographic data can be further exploited to gain insight into the role of loops in PBPs. We used three methods to assess loop flexibility: ensemble refinement (ER), refined structure B factors, and the PDBFlex server.

In high resolution data maps, it may be possible to fit a region of electron density with more than one model, which can be used to understand the flexibility of that region. The Phenix.Ensemble refinement program uses molecular dynamics simulations to generate alternative models of the protein conformation and then refines them into the available electron density 38,39. Structures of PaPBP3 (PDB code 6HZR 6) and NgTP2HR-6140 (PDB code 6HZJ 6) were refined with ER in Phenix.Ensemble and the resultant models were compared to the published structure (Figure 4.2a and d). ER led to unchanged or improved R values (Table 4.2), indicating that the ER models fit well to the crystallographic data.

<table>
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<tr>
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*Table 4.2. R values of different refinement methods.* Comparison of R values between published values and ER values.

The B-factor is a measure of the change in intensity of the X-ray scattering of an atom due to its dynamic and static disorder, which is determined for each atom during refinement cycles 46. B-factors can be used to describe the intrinsic flexibility of a crystallographic model 47.
Finally it is possible to exploit the large number of structural datasets available for PaPBP3 using the "PDBFlex" server 44, to see flexibility of the protein across different crystal preparations and with different ligands. This server clusters and aligns protein structures with high sequence similarity (> 95 % identity) and assesses regions of structural similarity. There are 38 PaPBP3 structures in their cluster (none of the structures described in the Chapter 5 of this thesis were included in the dataset) but there are only 5 in the cluster of NgPBP2, which is insufficient for representative work.

The amount of conformational flexibility observed using the Phenix.Ensemble method correlates well with the crystallographic B-factor of a residue (Figure 4.2a and d). The 3D structures of the ensemble models show the positions that are possible: the positional variations are not random but are constrained to specific movements, within specific sections of the loop (Figure 4.2b and d). The results from the PDBFlex server cluster generally agree with the results from the single structure in Figure 4.2a, but are typically less noisy. The flexibility of the β3-β4 loop appears to have been underestimated by the server because it is not present in many of the structures analysed.
Figure 4.2. Investigation of loop movements of PaPBP3 and NgPBP2. (a) Results of ensemble refinement (ER) by Phenix.Ensemble of PaPBP3 (PDB code 6HZR 6). Root-mean-square deviations (RMSD) of each residue of P. aeruginosa PBP3 following ER. The RMSD of each of the 64 models generated by Phenix.Ensemble are compared to their position in the published crystal structure and then averaged and plotted for each residue (blue, left axis). The refined B-factor of each atom in the published crystal structure is shown (orange, right axis). Loop regions are highlighted in grey and labelled, the N-terminal domain (up to residue 221) is highlighted with a red box. Residues 489-501 are not resolved in the crystal structure and are missing from the analysis. (b) Overlays of the models generated by ER of PaPBP3 (PDB code 6HZR 6). The protein is represented in cartoon and certain loops are coloured and labelled. The greatest loop positional variation is seen for the β5-α11 and α10-β3 loops. The β3-β4 loop is too flexible to be resolved in the crystal model. (c) The PDBFlex server generated a per-residue RMSD using a cluster of 38 models of PaPBP3. These are plotted similarly to (a). The dual peaks in the β3-β4 loop is likely due to the frequent absence of residues in this region from refined structures. (d) Results of ER of the transpeptidase domain of NgTP246-6140 (PDB code 6HZJ 8). As in (a), the RMSD of each of the 50 models generated by ER are compared to their position in the same subunit in the
published crystal structure and then averaged and plotted for each residue (blue, left axis). The refined B-factor of each atom in the published crystal structure is shown (orange, right axis). Two subunits (A and B) are found in the asymmetric unit, each with slightly different conformations. A is shown by a solid line and B with a dashed line. (e) Overlays of the models generated by ER of subunit A of NgTP2\textsuperscript{HRI6140} (PDB code 6HZJ \textsuperscript{6}). The protein is represented in cartoon and certain loops are coloured and labelled.

By all three analysis methods, flexible loops appear as sharp spikes compared to the overall relatively static protein (see protein median values in Table 4.S1). In the PaPBP3 transpeptidase domain, the flexibility of the β2 region, the β3-β4 loop and particularly the β5-α11 is clear. The α10-β3 region has a high RMSD in the single high resolution crystal structure (Figure 4.2a) but this is not consistent across the PaPBP3 crystals (Figure 4.2c). In NgPBP2, the subunits within the asymmetric unit each have differently flexible loops. In subunit A the β3-β4 loop has a high degree of flexibility, whilst the β5-α11 loops are static. The opposite is true in subunit B. The α10-β3 loop has no more flexibility than an average section of the mainchain (Figure 4.2d).

Three other flexible regions are highlighted by three flexibility analysis techniques (Figure 4.2a, c and d). These regions located within the following loops on the “back” face of the transpeptidase domain: β1b-α2 (residues 266-293), in the nominal α8 region (residues 374-398), α9-α10 (residues 427-450) (Figure 4.3). Mutations were not consistently identified in these regions.
Figure 4.3. Loops on the “back” face of PaPBP3. Models generated by ER of PaPBP3, as in Figure 4.2b. Analysis (Figure 4.2 in the main text), shows flexibility occurring in an additional 3 loops to those discussed in detail in the text: the β1b-α2 loop (green, residues 266-293), in the nominal 49 α8 region (yellow, residues 374-398), the α9-α10 loop (cyan, residues 427-450). The α8 helix is not fully resolved in the PaPBP3 crystal structure. These regions have little significant secondary structure, but still only show flexibility in limited sections of the loops. All three loops are on the “back” face of the protein relative to the active site. (b) Replot of the graph shown in Figure 4.2c (generated by the PDBFlex server), highlighting the peaks in the RMSD curve due to the flexibility of the 3 loops.

The core secondary structure provided by the α + β fold of the transpeptidase domain (shared across the penicilloyl-serine transferase superfamily, Chapter 1) gives overall rigidity to the protein. Its secondary structure inflexibility is shown by the very low RMSDs of the main chain atoms (Figure 4.2). For example, residues
294-305 of PaPBP3 (helix α2) for which all residues have an average RMSD of less
than 0.1 Å, as assessed by ER and PDBflex (Figure 4.2a and c). Consequently, the
catalytic residues: (e.g. in PaPBP3 Ser294, Ser349, Lys297 and Lys484) all exhibit
low flexibility (Table 4.S1) as these residues are all located with the core structural
elements. It has been suggested that rigidity of the active site residues allows for
the fast catalysis of β-lactamases 49. Rigidity of these residues may ensure they stay
optimally spaced to allow for rapid activity once the substrate binds50,51. The
possible movements of the loops are constrained by the rigidity of the adjacent
secondary structure elements.

These results build a picture in which certain privileged regions (with some
consistency between species) are flexible, but much of the protein is highly
constrained. It has previously been observed that the global motions (slowest
dynamics of a protein) are defined by the protein fold 52 and that the movement of
individual loops are determined by the global motions of the protein 53,54. Like the
secondary structure, loop dynamics may be a “structure-encoded” property of the
penicilloyl-serine transferase superfamily fold.

4.3.2 Novel Ligands Generate Novel Conformations

Chapter 5 describes the interactions of PBPs with boronates. While investigating the
crystal structures of boronates bound to PaPBP3, it was observed that the
β5-α11 and α10-β3 loops adopt novel conformations compared to previously
published β-lactam reacted or apo structures. Whilst the protein used was wildtype,
its response to novel ligands may indicate cryptic behaviours of the protein, which
may be contributing to the resistance mechanism.

4.3.2.1 Conformations of the β5-α11 Loop

In the majority of the PaPBP3:boronate structures investigated in Chapter 5, the
density of residues in the β5-α11 loop was too poor to allow a model to be fit, likely
due to the flexibility of the region. However, structures with 1, 2 and 14 all display
unique conformations of the loop (Figure 4.4). Whilst the density of the side chains
is not complete in all models, they provide some insight into the flexibility of the
region. Density for each of the structures is shown in Figure 4.4b-d. The
conformation adopted by 1 is particularly unique, with the loop folding closer to the
β3 strand than it is observed in any other structure (Figure 4.4a and b).
These structures provide further evidence of the flexibility of the β5-α11 loop and also indicate that its conformation is dependent on the nature of the ligand binding at the active site, despite the fact that these ligands make few direct interactions with residues of the loop (1 and 14 with Gly534 and Gly535 and 2 with only Gly534).

Figure 4.4. Boronate binding can affect the conformation of the β5-α11 loop in PaPBP3. (a) Conformations of the β5-α11 loop in PaPBP3 structures with 1 (lilac), 2 (yellow), 14 (brown), and reacted piperacillin (black, molecule of reacted piperacillin itself not shown, PDB code: 6R3X) bound, as well as the apo structure (white, PDB code: 6HZR). Electron density for the loop with: (b) 1, (c) 2 and (d) 14 bound. The PaPBP3-14 complex (d) and the piperacillin reacted PaPBP3 have similar main chain conformations, however residues Tyr503, Tyr532 and Phe533 have a different arrangement of their π-stack. Electron densities were contoured at 1 σ and produced using the comfit function in the CCP4 suite. 
4.3.2.2 Conformations of the $\alpha10$-$\beta3$ Loop

The $\alpha10$-$\beta3$ loop has not been observed to exist in many conformations in previously published structures of PaPBP3 (i.e. Figure 4.2c shows a low RMSD for this region). However, compounds 2 and 13 both change the conformation of the $\alpha10$-$\beta3$ loop.

The structure of PaPBP3:13 shows two alternate conformations of four residues (476-480), each with comparable occupancy (48 % and 52 %) (Figure 4.5). Movement of this portion of the loop is within the bounds set by the ER (Figure 4.2a and b), but other PaPBP3 structures (e.g. piperacillin reacted (6R3X), apo (6HZR 6) nor any of the other structures with boronate described in this thesis) do not have sufficient density to so clearly define the two alternate conformations.

![Image](image_url)

**Figure 4.5. The crystal structure of PaPBP3:13 shows two alternate conformations of the $\alpha10$-$\beta3$ loop.** Omit map density (blue mesh) for residues 474-480 is shown. Electron densities were contoured at 1 $\sigma$ and produced using the comit function in the CCP4 suite 55.
Figure 4.6. Boronate binding can affect the conformation of the α10-β3 loop in PaPBp3. (a) Comparison of the α10-β3 loop position in the crystal structure of PaPBp3:2 (cyan and orange) and in a piperacillin reacted structure (PDB: 6R3X, black and grey). The loop is significantly displaced between residues 466 and 473, compared to the piperacillin reacted structure, which is itself much like the apo structure (PDB: 6HZR, Figure 4.S2). (b) Omit map density (blue mesh) for residues 466-473 of the PaPBp3:2 structure. The density of all the residues is well defined. Electron densities were contoured at 1 σ and produced using the comit function in the CCP4 suite. (c) Proposed mechanism for the transmittance of changes at the active site to changes in the α10-β3 loop. (1) the binding of the imidazole group at the ‘acid binding pocket’ (hydrogen bonding to Ser485 and Gly534) leads to a rearrangement of the β5-α11 loop and the α11 helix. (2) The sidechain of Phe472 moves towards the α11 helix, with a resultant change in the position of the α10-β3 loop (3).

The change in conformation observed with the PaPBp3:2 is more significant (Figure 4.6) and to my knowledge entirely novel. The loop extends out from the protein and away from the preceding α10 helix which aligns well to the piperacillin reacted structure (PDB: 6R3X), as does the loop beyond residue Ala474. The cause of the change in conformation is unclear, as no new hydrogen bond interactions, (e.g. with the α11 helix) are formed. The effect is not caused by the
tri-covalent bonding of the protein (see Chapter 5), as the binding of 1 (also tri-covalent) does not elicit this change in PaPBP3. It could be linked to the novel conformation of the β5-α11 loop that is observed in the structure of PaPBP3:2 (Figure 4.4a and c). One explanation could be that the binding of the pyrazole group of 2 causes the change in the conformation of the β5-α11 loop, and tightens the first twist of the α11 helix, leading to a displacement of residues Gly535 and Leu536 (1), Figure 4.4c). The displacement allows Phe472 to shift towards the active site (2) with the accompanying shift in the α10-β3 loop (3) (Figure 4.4c). In this way changes at the active site are transmitted to a change at the α10-β3 loop.

Separately, in both TEM-1 56 and SHV-1 57 β-lactamases, the region between the α10-β3 loop and helix α11 has been shown to be a “cryptic” binding site for weak allosteric inhibitors (Figure 4.5). Demonstration of the conformational flexibility of this region raises the possibility of allosteric inhibition of PBP3, something which is previously unreported.

4.3.2.3 Networks of Coupled Promoting Motions

The above examples suggest the presence of an indirect link between the binding of a specific ligand at the active site and the conformation of peripheral loops. This same link could be used to transfer the effect of mutations on the peripheral loops to changes in the active site. In bacterial dihydrofolate reductase (DHFR, the target of trimethoprim), it has been suggested that resistance mutations can affect the rate of catalysis through “networks of coupled promoting motions” 58,59. These networks consist of interacting chains of amino acids which transmit information through the protein by their concerted motions 58,59. Studies show that point mutations up to 20 Å away can influence the rate of catalysis as much as 200 fold 58,60 by this proposed mechanism. These networks are suggested to be a common feature of protein families, which may explain the conservation of locations of mutations across class B PBPs from different gram-negative species (Table 4.1). As has been previously observed with PBP3 6, the clinical mutations of a protein target do not necessarily lead to a change in the protein crystal structure 60,61. Instead, mutations can elicit a change in the protein dynamics, which affects the ligand processing 60,61.
4.4 Future Work: Beyond Cryo-Crystallography

Methods that do not rely on crystallographic evidence are required to assess protein dynamics. An over-reliance on crystallographic data may lead to overinterpretation of conformations that are actually the consequence of crystallographic artifacts such as crystal packing and are not biologically relevant. Crystal structures of the NgTP2\textsuperscript{HR-6140} and NgTP2\textsuperscript{39-6140} are illustrative (Figure 4.S1): the crystal structures prepared by different researchers have significant deviations in conformations of local regions, which are difficult to account for \textsuperscript{6,20}.

The change in dynamic behaviour of the protein (assessed with some of the tools below) could be correlated with the β-lactam-binding activity of the protein (using some of the methods listed in Table 1.2), for each of the mutations of interest. In this way, the changes in protein dynamics could be used to explain the mechanism of resistance, and perhaps even the contributions of each of the three pathways (i-iii, described above).

4.4.1 Room Temperature Crystallography

Protein crystallography is typically conducted at around 100 K, which helps reduce the effect of radiation damage that samples experience when exposed to the high energy beams of a synchrotron \textsuperscript{62}. However, this rapid freezing is thought to further bias the crystals into conformations which may not exist under physiological conditions \textsuperscript{63,64}. When X-ray diffraction patterns are instead collected at room temperature, studies have shown that different side chain conformations can be adopted and that additional backbone conformations become possible \textsuperscript{63}, presumably making the protein more similar to the physiological solution state. Due to the complex energy networks within the protein, \textit{a priori} prediction of the effects of warming on protein structure is challenging \textsuperscript{63}. Methods to study proteins at room temperature (including techniques which circumvent the issue of increased radiation damage) are being constantly improved, such as the VMXt beamline (being commissioned at Diamond Light Source) \textsuperscript{65} and X-ray free electron lasers \textsuperscript{66}.
4.4.2 Tryptic Digestion

A simple method to observe flexibility is to use limited tryptic digestion of a protein, which was used to validate the effects of mutations in NgPBP2 ⁶ and PBP2x from S. pneummoniae ⁶⁷. Decreased flexibility of the β2 loop region (caused by the introduction of mutations that provide resistance to β-lactams) was used to explain the lack of tryptic digestion within this section of the protein, compared to the wild type. When more rigid, the protein does not allow the trypsin to access the site of proteolysis. This technique is quick and highly site specific, but provides a very qualitative insight into the changes occurring.

4.4.3 NMR

Nuclear magnetic resonance (NMR), can show the interactions and dynamics of amino acids within proteins in solution at physiological temperature with atomic resolution ⁶⁸ and is well suited for the task of analysing protein dynamics. NMR tools are available which can study protein dynamics on timescales of ps-ns or μs-s ⁶⁹. Whilst solving a full protein model by NMR is challenging, NMR can be used in a more limited context to understand individual movements. After tagging mutated cysteines with fluorine-based probes the Schofield group and others from the University of Oxford were able to study the loop dynamics of metallo-β-lactamases using ¹⁹F NMR ⁷⁰,⁷¹. The ¹⁹F signal is highly sensitive to local changes in the environment, with movements of the loops the probes are placed on and binding of ligands to the active site leading to a change in signal. Two loops can even be tagged simultaneously and their varying interactions with different ligands observed ⁷¹. One limitation of this is that it requires the addition of a cysteine residue to the loop which itself may act as a resistance determining mutation.

4.4.4 Molecular dynamics

Molecular dynamics, where the internal structure of a protein is mechanically modelled as a series of masses and springs can be used to simulate the movements of proteins ⁷². The method analyses the protein as a series of very small (~femtosecond), discrete timesteps, so long simulations can be computationally expensive (typically up to a few nanoseconds of simulation is possible). However
the global dynamics of the protein (large scale movements) such as loop movements, allostery and substrate binding occur on the timescale of microseconds to milliseconds \(^{73-75}\). This means that feasible simulations may be too short to appreciate loop dynamics. This type of analysis has been applied to penicilloyl-serine transferase superfamily proteins and revealed their dynamics \(^{49,76-79}\).

A recent work using MD to understand the loop dynamics of class A \(\beta\)-lactamases (TEM-1 and KPC-2) shows the loops of these \(\beta\)-lactamases have important roles in \(\beta\)-lactamases, as we propose for PBP3 \(^{79}\). Simulations at equilibrium on the enzymes showed greatest fluctuations for the equivalent regions to the (PBP3) \(\alpha10-\beta3\) (referred to as the ‘hinge’ and \(\alpha11\) in TEM-11), the \(\beta3-\beta4\) loop (\(\beta7-\beta8\) in TEM-1) and the \(\beta5-\alpha11\) loop (\(\beta9-\alpha12\) in TEM-1), similar to the conclusions I have drawn from the crystal structures of class B PBP3 (section 4.3.1).

The authors also investigated the effect of removing a TEM-1 allosteric inhibitor (Figure 4.53), which sits between the TEM-1 equivalent of the \(\alpha10-\beta3\) loop and the \(\alpha11\) helix. Simulations of non-equilibrium protein dynamics for the 5 ns following removal of the allosteric ligand showed ‘signal propagation’ that led to changes in the equivalent region to \(\beta2\) region, ~33 \(\AA\) from the allostERIC site. One of the signal propagation pathways proposed was via the equivalent to the \(\alpha11-\beta5\) loop. As in PBP3, clinical mutants map onto the regions proposed for the signal propagation route \(^{79}\). This example parallels our hypothesis that the active site loops of PBP3 have important roles for \(\beta\)-lactam binding and that clinical mutants modify this.

Combinations of molecular dynamics and NMR can be a powerful tool for studying protein dynamics. In TEM-1 \(\beta\)-lactamase, NMR analysis of rigidity (measured over a short \(<\text{ps} \text{ timescale}\)) and molecular dynamics simulations (which model movements occurring over a ps-low ns timescale) showed that loop regions are more flexible than NMR analysis alone predicts \(^{49}\). The authors develop a model in which there is high rigidity over short time scales to increase the rate of catalysis, coupled with conformational changes happening over a \(\mu\text{s}-\text{ms} \text{ timescale}\), which has been supported by further NMR in chimeric \(\beta\)-lactamases \(^{80}\).
4.5 Conclusions

Our study on the locations of point mutations that cause target-mediated resistance in PBP3 as well as observations from crystal structures lead us to hypothesise certain loops may play a more important role in PBP3 structural dynamics than previously understood. Analysis with ensemble refinement and comparisons across many structures of PaPBP3 indicates flexibility within these same regions, but tools beyond crystallography are needed to better characterise their movements. Work in DHFR on correlations between mutations and flexibility provide a useful precedent for how work on PBPs may be conducted.

Use of unligated proteins will only provide limited information, and a more complete picture of target mediated resistance requires a better understanding of the exact nature of protein:drug interactions. Currently, only crystal structures of the apo and covalently reacted PBP3s (as well as a few examples of hydrolysed β-lactam structures in the active site) are available. Work with cephalosporins bound to mutant and wild-type NgPBP2 is helping to explain the effects of some mutations. We have demonstrated that novel ligands such as boronates can influence the conformations of the loop regions. Further crystal structures with novel ligands may be beneficial in this effort. Perhaps the most important ligand to crystallise is the natural substrate, for which there remains no crystal structures in a class B PBP. Any successful mutation to a PBP must reduce β-lactam binding, but allow natural substrate processing to proceed at sufficient rates to allow growth. Studies are required to assess how this is possible.

It is interesting to consider the impact of introduction of new, non-β-lactam PBP inhibitors (Table 1.1). For example, if inhibitors are found which lack β-lactamase susceptibility (e.g. boronates: Chapter 5) will this lead to a bias in the evolved mechanism of resistance towards target-mediated resistance? Better knowledge of PBP3 target mediated resistance may allow for its emergence to be predicted and for ligands to be developed in which resistance by this mechanism is minimised.
4.6 References


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https://doi.org/10.25080/Majora-629e541a-00e.


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Chapter 4.S. Supplementary Information for Chapter 4

4.S1 Additional Crystallographic Views and Data

Figure 4.S1. Comparisons between two structures of the transpeptidase domain of NgPBP2 from strain FA6140. The two structures NgTP2^{6HZJ} and NgTP2^{4U3T} (PDB: 6HZJ \(^1\) (blue) and 4U3T \(^2\) (orange)) are of the same protein but have different resolutions (1.4 vs 2.2 Å respectively) and different crystallisation conditions. All data here are comparisons of subunit A in both structures. (b) Root-mean-square deviation (RMSD) between the backbone atoms of each residue of the two structures. Large deviations between the structures are observed in the later part of the β2 region (c) and at the β3-β4 loop (d), as well as in the α9-α10 loop (residues 451-461) on the “rear” of the protein (Figure 4.3). Part of the β4-α11 loop was not fully resolved in NgTP2^{6HZJ} subunit A so these four residues (residues 542-545) were omitted from the analysis. (c) The β2 region, with residues Asp346 and Asp346a (sometimes denoted as Asp345a and Asp346 respectively \(^2\)) shown in stick representation. The latter part of the region (residues 345-354) has the greatest conformational deviation. (d) The β3-β4 loops of the proteins adopt different conformations.
**Figure 4.52. Comparison of apo and piperacillin reacted PaPBP3 crystal structures.** Apo PaPBP3 (PDB: 6HZR 1, white) and piperacillin reacted PaPBP3 (PDB: 6R3X 1, black) show different conformations, particularly at the β5-α11 (residues 488-503) and β3-β4 (residues 525-537) loops. The β5-α11 loop transitions from the open conformation in the apo model to the closed formation in the piperacillin reacted model (indicated by a red arrow). Residues Tyr503, Tyr532 and Phe533 are shown in sticks and labelled. They form a hydrophobic wall against the phenyl group of reacted piperacillin in the closed conformation. The β3-β4 loop is unresolved in the apo structure, but resolved in the piperacillin reacted model, which may indicate a decrease in conformational flexibility upon β-lactam binding. (b) The RMSD between the two structures, plotted for each residue. The N terminal domain is highlighted in pink and the 4 loop regions are labelled. The β5-α11 loop is the region of greatest difference between the two structures.
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<td>0.0162</td>
<td>10.07</td>
<td>11.17</td>
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</tr>
<tr>
<td>K(S/T)G</td>
<td>8.26</td>
<td>0.0833</td>
<td>0.1442</td>
<td>10.53</td>
<td>11.43</td>
<td>0.1216</td>
</tr>
</tbody>
</table>

Median Value*  
10.86 0.0985 0.04 13.31 13.85 0.1717 0.1788

Table 4.S1. Local flexibility of active site residues. Values are means of the value for the residue given in bold and the adjacent 2 residues either side of the residue in bold. For PaPBP3 the residues are (top to bottom): Ser294 (SXXK), Lys297 (SXXK), Ser349 (SXN) and Lys 484 (KSG), for NgPBP2 they are: Ser310 (SXXK), Lys310 (SXXK), Ser362 (SXN), Lys 497 (KTG). See Figure 4.1a. See Figure 4.1c. See Figure 4.1d. *Median value for the structure/subunit.
Figure 4.S3. TEM-1 was found to have a cryptic allosteric site between the α11 helix and the α10-β3 loop. The crystal structure of TEM-1 bound to the tetrazole-containing compound shown (pink, PDB: 1PZP), reveals two binding sites for the compound. One of these is between the α10-β3 loop and the α11 helix. TEM-1 and PBP3 are both members of the penicilloyl-serine transferase superfamily and share the same structural fold, the structure of TEM-1 is compared to the crystal structure of PaPBP3:2 (cyan), with the secondary structure of PaPBP3 labelled. The cryptic allosteric binding site of the tetrazole-containing compound is shown by the red dashed line. If the α10-β3 loop is sufficiently flexible, it may be possible to find similar allosteric inhibitors of PBP3.

4.S2 References


Chapter 5. Boron-based Inhibitors of PBP3: Benzoxyaboroles with Novel Binding Modes

Chapter 5.S (p219-234) contains additional information referred to in this chapter

5.1 Introduction

Currently, the most effective way to combat AMR is to discover compounds with novel modes of action or with novel chemistry for old targets. Previous attempts to target PBPs with novel chemistry are summarised in Section 1.4.5. One of the groups of compounds described therein (Table 1.1) are the boronates.

In the early 2000’s, the first boron-containing compound, bortezomib, was approved for use against multiple myeloma \(^1\)\(^2\) (Table 5.1A), affirming observations since the 1970’s that boronates could be used to bind and inhibit serine (and threonine) proteases \(^3\)\(^4\). Vaborbactam (approved)\(^5\)-\(^7\) and bicyclic boronate taniboractam (in clinical trials) \(^8\)-\(^10\) demonstrate that boronates can be clinically useful against penicilloyl serine transferase family proteins (both of these examples are \(\beta\)-lactamase inhibitors). However there are no compounds in academic literature which can act with the same high affinity against high molecular mass HMM PBPs, although other investigations have been reported \(^11\)-\(^22\). Boronate PBP inhibitors would be insensitive to \(\beta\)-lactamase hydrolysis which is a highly desirable property.

Boronates inhibit serine- (and metallo-) \(\beta\)-lactamases by mimicking the transition states in penicillin catalysis \(^11\)\(^22\), which themselves are analogous to the transitions states passed through during the natural substrate turnover of PBPs (Figure 5.1). On the catalytic pathway of both \(\beta\)-lactam hydrolysis and transpeptidation, the central carbonyl carbon transitions (twice) from \(sp^3\) to \(sp^2\) then back to \(sp^2\) hybridisation (Figure 5.1). Boron can similarly transition between a neutral \(sp^2\) hybridisation state and an anionic \(sp^3\) hybridisation state, and it is this ‘morphing’ property in particular that makes it successful at inhibiting \(\beta\)-lactamases \(^23\).
Figure 5.1. Boron mimicry of transition states on the transpeptidation and penicillin catalysis pathways of PBPs. General mechanism of (a) transpeptidation by class B PBPs and (b) the reaction of a β-Lactam with a PBP, exemplified with penicillin. This reaction is described in Figure 1.2. (c) Boron transitions between \( sp^2 \) and \( sp^3 \) hybridisation states and is able to mimic the tetrahedral transition states and inhibit the PBP. Hybridisation states of the central carbonyl carbon (or boron) of each species is shown with a coloured circle; yellow: \( sp^2 \), pink: \( sp^3 \).

Crystal structures of boronates with various proteins (including PBPs and β-lactamases) have shown the boron atom can form high valency complexes: with the hydroxlys of the ribose sugars of the 3'-adenosine monophosphoryl moiety of tRNA (Table 5.1B), with the active site residues of R39 pd-peptidase (Table 5.1C) and with nucleophilic serine and histidine residues in trypsin (Table 5.1D and E).
Table 5.1. Crystallography shows boronates can form a number of interesting and high valency complexes. (A) The first approved boron-containing drug was bortezomib, a peptido-mimetic proteasome inhibitor. It reacts with a nucleophilic threonine to reversibly inhibit the proteasome of cancer cells. This example is from a yeast homolog of the human proteasome (PDB: 2F16). (B) Tavaborole (a benzoxaborole) is an approved topical antifungal agent. It engages the vicinal hydroxyls of the ribose moiety of the 3’ adenosine of tRNA within the leucyl-tRNA synthetase editing site (PDB: 2VOG). The first reported example of tri-covalent binding of a boronate. This alkyl boronate can engage 3 residues of the active site of R39 (a LMM PBP) of Actinomadura sp. The residues shown are from the SXXK, SXN and K(S/T)G conserved motifs, like the examples shown in this chapter (PDB: 3ZVT). (D) and (E) examples of boronates reacting with trypsin. Boronate crystal structures indicate fragments can bind at multiple sites within the protein (some sites have a lower occupancy) (PDB: 2A32 and 2A31, respectively). A di-covalent His/Ser complex was also observed with a boronate in a chymotrypsin crystal structure. In (D) only one of the two refined alternate conformations of benzene boronic acid is shown. In the other conformation, the positions of the phenyl and hydroxyl groups are switched. (F) The benzoxaborole warhead has previously been studied for its use as a β-lactamase inhibitor. In AmpC, the residue equivalent to Ser349 of PaPBP3 is replaced by a tyrosine (Tyr177). Perhaps as a result, di-covalency is not observed, and the catalytic serine (Ser62, equivalent to Ser294) attacks the benzoxaborole from the other face compared to the direction of attack in PaPBP3 (Figure 5.S1).
The chemistry of boronates makes them particularly able to engage nucleophilic serines, with selectivity over cysteine nucleophiles. An NMR study of the reactivity of cysteine and serine residues in solution found that the benzoaborole pharmacophore was the most reactive warhead tested towards serine, but had low reactivity against cysteine.

The high valency reversible covalent attachment, serine selectivity and "morphing" properties of boron are all highly desirable for a warhead designed to engage PBPs. As antimicrobials, boronates (against other targets) have precedent in gram-negative bacteria, gram-positive bacteria and Mycobacterium tuberculosis.

A previous attempt by our group to use the XChem fragment screening capabilities available at Diamond (section 1.3.3) to screen PaPBP3 with a library of 1,300 diverse fragments returned only one hit (Table 1.1), which is a far lower hit rate than anticipated for this technique. The fact that the sole hit covalently bonded to the protein seemed remarkable so we decided to re-screen the protein with a library of known serine-enzyme covalent warheads, further enriched with boron-containing compounds.

5.2 Methods

5.2.1 Proteins and Crystallography

Proteins were expressed, purified and crystallised by D. Bellini, who was also responsible for collecting the crystallographic data. Protocols as previously published were followed, further details are described in.

All collected crystallographic data were processed using the automated pipeline at Diamond Light Source with data reduction by XDS (Version Nov. 11, 2017) and data scaling by AIMLESS (Version 0.6.3). Processing was carried out by either autoPROC or STARANISO. STARANISO was used when there was significant anisotropy of the data, (all PaPBP3 complexes except PaPBP3:8 and PaPBP3:16). PaPBP3:8 and PaPBP3:16, were isotropic and processed with autoPROC. Structures were phased with molecular replacement by Phaser_MR using a high
resolution structure of PaPBP3 (PDB: 6HZR)\(^{35}\) as the search model. Manual
building and ligand fitting were performed with COOT.\(^{41}\) Refinement was carried
primarily using REFMAC5 \(^{42}\) within the CCP4 suite \(^{43}\) as well as in phenix.refine.\(^{44}\)
Ligand constraints were generated by the grade webserver.\(^{45}\) Structures were
validated with MolProbity.\(^{46}\) Figures of structures were prepared using PyMOL (The
PyMOL Molecular Graphics System, Schrödinger, LLC) or CCP4mg.\(^{47}\) Composite
Omit maps that are shown they were calculated by ‘comit’ in the ccp4 suite.\(^{43}\)

The structures are published in the PDB under the following IDs; PaPBP3 in
complex with: \(1\), 7ATM; \(2\), 7ATO; \(3\), 7ATW; \(4\), 7ATX; \(8\), 7AU0; \(13\), 7AU1; \(14\), 7AU8;
\(15\), 7AU9; \(16\), 7AUB; **Vaborbactam**, 7AUH. Note that there are fewer compounds in
the published paper for these entries \(^{37}\), so the compound numbers in the PDB
entries do not correlate to those in this thesis.

Docking protocols (performed by J. Eyermann) are given in 5.5.4.

### 5.2.2 BOCILLIN FL Assays

Unless otherwise stated, assays were run in triplicate, using 60 nM purified protein,
30 nM BOCILLIN FL in pH 7, 100 mM sodium phosphate buffer, with 0.01 % Triton
X-100 to reduce promiscuous ligand binding \(^{12}\). Assays were in a volume of 50 \(\mu\)L,
in black, flat bottom, 384 well microplates (Greiner Bio-One) at 30 °C. The change in
fluorescence anisotropy \((r)\) was measured using a ClarioStar plate reader (BMG
Labtech) with polarised filters at excitation: 482 ± 16 nm, emission: F: 530 ± 4 nm
and calculated using MARS software v3.32 (BMG Labtech) using the equation:

\[
\begin{align*}
   r & = \frac{I_{parallel} - I_{perpendicular}}{I_{parallel} + 2 \cdot I_{perpendicular}} \\
\end{align*}
\]

where \(I_{parallel}\) and \(I_{perpendicular}\) are the fluorescence intensity parallel and perpendicular
to the excitation plane respectively (Section 1.4.6 and Figure 1.5).

Residual activities were found by pre-incubating the test compound and protein with
1024 \(\mu\)M test compound for 1 hour at 30 °C, before the reaction was initiated by the
addition of BOCILLIN FL. The change in fluorescence anisotropy after 30 minutes
was compared to the uninhibited control to determine the residual activity.
In order to calculate $K_i$, the compound and BOCILLIN FL were mixed and the reaction was initiated by the addition of the PBP. Also included in the experiment were progress curves of nine concentrations of PBP (40–110 nM). Progress curves were analysed with Global Kintek Explorer 8.0 (KinTek, USA), as described $^{48,49}$. Reported errors represent the standard error of the fit of the data to the model, produced by the software. The reaction between the PBP and BOCILLIN FL (Scheme 5.1) is simplified to a one-step model, with rate constant $k_1$ $^{48}$. However, unlike the previous studies, we found it was necessary to include a term ($k_2$) to account for the de-acylation of BOCILLIN FL from the PBP, which releases a hydrolysed product ($P$) (Scheme 5.1). The fluorescence intensity was constant throughout the reaction. Inhibitor binding was modelled as a reversible reaction to form the enzyme-inhibitor complex ($EI$) (Scheme 5.1). $K_i$ was calculated as the ratio of the off-rate $k_{off}$ to the on-rate $k_{on}$ (Scheme 5.1).

![Scheme 5.1. The model used to determine $K_i$ in Kintek Global Explorer. Terms $E$, $B$, $EB$, $P$, $I$ and $EI$ represent the PBP, BOCILLIN FL, enzyme-BOCILLIN FL complex, hydrolysed BOCILLIN FL, the inhibitor and the enzyme-inhibitor complex, respectively. $k_1$ models the acylation rate of BOCILLIN FL and $k_2$ the de-acylation rate of BOCILLIN FL$^{48}$. The enzyme-inhibitor interactions are modelled by the inhibitor $k_{off}$ and $k_{on}$ rates, with the $K_i$ the ratio of these values.](image)

### 5.2.3 S2d Assay

To confirm compound activity in an analogous assay, residual activities of substrate analogue S2d turnover by PaPBP3 $^{13,51–53}$ were determined in the presence of the compounds. Assays were conducted in 50 μL in a 384 well, clear bottom, black walled microplate (Greiner Bio-One). PaPBP3 (400 nM) was incubated with 2 mM of
each compound for one hour at 30 °C in 100 mM sodium phosphate, pH 7, supplemented with 0.01 % (v/v) Triton X-100 (total volume 25 μL). A solution of 5,5'-dithiobis-(2-nitrobenzoic acid) (DTNB) and S2d (total volume 25 μL), diluted in the same buffer (to give a final concentration in the assay of 1 mM for both reagents) was added to each well to initiate the reaction (final volume 50 μl). The final concentration of protein was 200 nM and the final inhibitor concentration was 1 mM. The reaction was followed by a ClarioStar plate reader (BMG Labtech) measuring the change in absorbance at 412 nm at 30 °C. The assay was conducted in the absence of an inhibitor (positive control) as well as with an excess of aztreonam (1 mM) which completely inhibits PaPBP3. Treatment with aztreonam was therefore used to find the rate of spontaneous S2d hydrolysis in the buffer. The initial rate of S2d turnover was calculated using Prism 9 (Prism 9 for macOS, GraphPad Software LLC) and the standard error calculated from three replicates. The S2d turnover rate was corrected for non-enzymatic hydrolysis then expressed as a ratio of the positive control:

\[
Residual\ Activity\ (\%) = 100 \times \frac{[\text{Initial rate in the presence of inhibitor}]-[\text{non-enzymatic rate}]}{[\text{Initial rate of positive control}]-[\text{non-enzymatic rate}]}
\]

5.2.4 Nitrocefin Assay

Nitrocefin \(^{54,55}\) turnover by PaPBP3 \(^{11,56}\) was used to determine the effect of a two-fold dilution on the \(pIC_{50}\) of 13. Assays were performed in clear bottom 384 well plates (Greiner Bio-One). Assays at an initial volume of 40 ml, contained 50 mM bisTris propane, pH 8.5, containing 1 % (v/v) Triton X-100 and 20 mM MgCl₂ and 150 μM Nitrocefin (Abcam). Assays were initiated by the addition of 238 mM PaPBP3, and the absorbance at 482 nm was monitored for 120 s at 30 °C using a ClarioStar plate reader (BMG Labtech). At this point, an additional 40 μL of buffer was then added (final volume: 80 μL), and the plate was shaken (40 s at 500 rpm), following which additional readings were taken. Rates of these linear curves (corrected for background nitrocefin turnover) were calculated using the in-built data analysis package of the plate-reader (MARS; BMG Labtech) and compared to the untreated control to determine relative rates. Relative rates were then plotted against the inhibitor concentration. \(pIC_{50}\) was determined using Prism 9 (Prism 9 for macOS, GraphPad Software LLC).
5.2.5 Antimicrobial Assays

Minimum inhibitory concentration (MIC)s of compounds were determined against CLSI reference strains of the following Gram negative organisms: E. coli (NCTC 25922), P. aeruginosa (PAO1 and a permeabilised strain which was a kind gift from Zgurskaya and colleagues), H. influenzae (ATCC 49766), A. baumannii (ATCC 19606) and N. gonorrhoeae (ATCC 49226) by the broth microdilution method using a control antibiotic. CLSI procedures were strictly adhered to throughout, with the exception of total volume, which was reduced to minimise compound consumption. The E. coli and P. aeruginosa strains were tested in a 10 μL final volume in cation-adjusted Mueller Hinton broth on the lids of inverted 96 well Costar microplates (Corning, USA) and incubated at 37 °C for 18 hours in a humidor (~98% humidity) to reduce evaporation. H. influenzae, A. baumannii and N. gonorrhoeae were tested in 50 μL of cation-adjusted Mueller Hinton broth supplemented with 5% lysed blood in 96-well Costar microplates (Corning, USA) and incubated at 37 °C for 18 hours in ~5% CO₂. Growth was determined by visual inspection of the plate after the incubation period. Data with the E. coli ΔToIC strain was collected as part of the automated screen of boronates, carried out by the high throughput screening platform (Chapter 3).

Synergy studies were conducted using the checkerboard titration method with E. coli and P. aeruginosa in a total volume of 10 μL cation adjusted Mueller Hinton broth, piperacillin at concentrations in the range of 32 to 0.25 μg/mL and benzoxaborole concentrations varied between 64 and 1 μg/mL, diluted in perpendicular directions across the plate. Growth was determined by visual inspection of the plate after 18 hours at 37 °C at 95 % humidity.

5.2.6 Chemoinformatics

Data pipelining software KNIME v3.5.3 was used to triage fragments of the “Serine Focused Covalent Fragments” compound set from Enamine. The compound set was manually assessed and then grouped by warheads of interest (using SMILES of the functional groups shown in Table 5.2). The desired number of compounds from each group was chosen and then the most diverse subset of fragments from each of the warhead groups was purchased for screening. Diversity was selected for using the “Diversity Picker” module by RDKit, which implements the MaxMin algorithm. Further compounds were added manually after the
selection. The molecular properties shown in Figure 5.2 were calculated using the “Molecular Properties” core KNIME module and the XlogP module from CDK \(^{59,62}\). A complete list of the fragments screened is given in Table 5.S1.

### 5.2.7 Synthetic Chemistry

All commercial reagents were from Sigma-Aldrich, Fisher Scientific, Combi-Blocks, Enamine, or Fluorochem and were used without further purification. 1 and 3 were from Combi-Blocks, Inc., 2 and 4 were from Enamine. These were used without further purification in the X-ray fragment screen; 13 and 14 were purchased from Wuxi Appotec; 16 was a gift from Mukesh Gangar (H3D, University of Cape Town); vaborbactam was purchased from MedChemExpress; 2-((Benzoyl-D-alanyl)thio)acetic acid (S2d) was a gift from Robert Lesniak (University of Oxford).

Solvents were used as received. Flash column chromatography was performed using a Teledyne ISCO flash purification system using a Silicycle SiliaSep ™ C18 cartridge. Purity of all final derivatives (except 7: 88 %) for biological testing was confirmed to be > 95 % (section 5.S5) as determined using an Agilent UPLC–MS: Agilent Technologies 6150 quadrupole mass spectrometer with positive mode electrospray ionisation, coupled with an Agilent Technologies 1290 Infinity II series UPLC system Agilent 1290 series UPLC at two wavelengths 254 and 280 nm. The following conditions were employed: Chromatography was performed using a 50 mm x 2.1 mm Kinetex 1.7 μm particle size Evo C18 100A, LC column 50 mm x 2.1 mm, in solvent A of (0.1 % (v/v) formic acid in water), and solvent B of (0.1 % (v/v) formic acid in acetonitrile). The structures of the final products were confirmed by NMR and mass spectrometric analysis. \(^1\)H and \(^{13}\)C nuclear magnetic resonance (NMR) spectra were collected on a Varian Mercury 300 MHz spectrometer or a Bruker AVIII 600 MHz spectrometer. Deuterated solvents were used as supplied. Residual solvent peaks were used to reference chemical shifts (δ) which are reported in parts per million downfield from the residual solvent peak as an internal standard. Peak multiplicity is expressed as s (singlet), d (doublet), t (triplet), q (quartet), m (multiplet), br (broad), or a combination of these. Coupling constants (J) are reported in hertz (Hz) to the nearest 0.5 Hz. High-resolution mass spectra were recorded using a Bruker MicroTOF instrument with an electrospray ionisation source and Time of Flight (TOF) analyser. The parent ion is quoted with the indicated ion: [M - H]^+ or [M + Na]^+.
Scheme 5.2. Synthesis of benzoxaborole derivatives (A) 5-11 and 15, (B) 12.
Reagents and conditions: (a) 1,1'-carboxyldiimidazole (CDI), N,N-dimethylformamide (DMF), 40 °C, 4 – 16 hours; (b) LiOH·H₂O, 1,4-dioxane:H₂O (3:1), 40 °C, 60 min; (c) HCl (4 M solution in 1,4-dioxane), CH₂Cl₂, rt, 16 hours; Note that low isolation yields for substituted benzoxaboroles (e.g. 9-11) in part reflect significant losses during purification on silica gel and provide scope for further optimisation. For B1, R₁ = H and R₂ = CH(R₃)CO₂Me. The complete structures of 5 – 8, 15 and 9 - 11 are shown in Table 5.3. Dppf: 1,1'-bis(diphenylphosphino)ferrocene; Boc, tert-butoxycarbonyl. Thank you to Alen Krajnc for assistance with this figure.

5.2.7.1 General Protocol 1: amide coupling

To a solution of the appropriate carboxylic acid (1 equiv.) in N,N-dimethylformamide (2 mL) was added 1,1'-carboxyldiimidazole (2 equiv.). The reaction was stirred for 5 minutes at room temperature; the appropriate amine (1, 1.2 or 1.5 equiv.) was then added, and the reaction was stirred 4–16 hours at 40 °C. The solvent was removed in vacuo, and the crude product was purified using a Teledyne ISCO CombiFlash chromatography system eluting a C18 column with a reverse phase solvent gradient of MeOH in 0.1 % (v/v) CH₃CO₂H in water. The product-containing fractions were then combined, and the organic solvent was removed in vacuo. When amide coupling yielded a target intermediate compound (e.g. methyl esters of general structure B1, Scheme 5.2) it was used in the next step without further purification, else lyophilisation was used to afford the desired products as solids (i.e. for 5 - 11, and 15).

1-Hydroxy-N-[2-(methylamino)-2-oxo-ethyl]-3H-2,1-benzoxaborole-6-carboxamide (5)

General Protocol 1 was followed using the following quantities of reagents: 3 (100
mg, 0.56 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (182 mg, 1.12 mmol, 2 equiv.); 2-amino-\(N\)-methylacetamide. HCl (104 mg, 0.84 mmol, 1.5 equiv.). Product: crystalline solid (47 mg, 32 %). Purity: >96 % (by HPLC). 

\[^{1}H\text{NMR}\] (600 MHz, DMSO-\(d_{6}\)) \(\delta\) 9.55 (br s, 1H, OH), 8.71 (t, \(J = 6.0\) Hz, 1H, NH), 8.18 (s, 1H, Ar-H), 7.93 (dd, \(J = 8.0, 2.0\) Hz, 1H, Ar-H), 7.85 (q, \(J = 5.0\) Hz, 1H, NH\(CH_{3}\)), 7.49 (d, \(J = 8.0\) Hz, 1H, Ar-H), 5.02 (s, 2H, -\(CH_{2}\)OB), 3.85 (CH\(_{2}\), obscured by solvent peak), 2.59 (d, \(J = 5.0\) Hz, 3H, CH\(_{3}\)); \(^{13}C\text{NMR}\) (151 MHz, DMSO-\(d_{6}\)) \(\delta\) 170.3, 167.8, 157.6, 133.2, 130.3, 121.9, 70.4, 43.1, 26.1; \text{LCMS (ESI^+, m/z) calculated for C\(_{11}\)H\(_{13}\)N\(_{2}\)O\(_{4}\)^{10}\text{B}, [M+H]^+ : 249, found 249; HRMS (ESI-TOF, m/z) [M - H] : 247.0896, found 247.0895.}

\(N\)-\([1R]-2\text{-Amino-1-benzyl-2-oxo-ethyl]-1-hydroxy-3H-2,1-benzoxaborole-6-carboxamide (6)\)

General Protocol 1 was followed using the following quantities of reagents: 3 (100 mg, 0.56 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (182 mg, 1.12 mmol, 2 equiv.); (R)-2-amino-3-phenylpropanamide HCl (169mg, 0.84 mmol, 1.5 equiv.). Product: crystalline solid (80 mg, 43 %). Purity: >98 % (by HPLC).

\[^{1}H\text{NMR}\] (600 MHz, DMSO-\(d_{6}\)) \(\delta\) 9.31 (br s, 1H, OH), 8.45 (d, \(J = 8.5\) Hz, 1H, NH), 8.18 – 8.15 (m, 1H, Ar-H), 7.89 (dd, \(J = 8.0, 2.0\) Hz, 1H, Ar-H), 7.59 – 7.52 (m, 1H, Ar-H), 7.46 (d, \(J = 8.0\) Hz, 1H, Ar-H), 7.33 – 7.30 (m, 2H, NH\(_{2}\)), 7.24 (t, \(J = 7.5\) Hz, 2H, Ar-H), 7.20 – 7.13 (m, 1H, Ar-H), 7.12 – 7.08 (m, 1H, Ar-H), 5.02 (s, 2H, -\(CH_{2}\)OB), 4.66 (ddd, \(J = 10.5, 8.5, 4.0\) Hz, 1H, -\(CHNH\)), 3.12 (dd, \(J = 14.0, 4.0\) Hz, 1H, -\(CH_{2}\)Ph), 2.99 (dd, \(J = 14.0, 4.0\) Hz, 1H, -\(CH_{3}\)Ph); \(^{13}C\text{NMR}\) (151 MHz, DMSO) \(\delta\) 173.8, 166.9, 157.3, 139.0, 133.5, 130.3, 130.2, 129.6, 128.5, 126.7, 121.6, 70.4, 55.2, 37.7; \text{LCMS (ESI^+, m/z) calculated for C\(_{11}\)H\(_{13}\)N\(_{2}\)O\(_{4}\)^{10}\text{B}, [M+H]^+ : 325, found 325; HRMS (ESI-TOF, m/z) [M + Na]^+ : 347.1174, found 347.1176.}

\(N\)-\([1R]-2\text{-Amino-2-oxo-1-phenyl-ethyl]-1-hydroxy-3H-2,1-benzoxaborole-6-carboxamide (7)\)

General Protocol 1 was followed using the following quantities of reagents: 3 (100 mg, 0.56 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (182 mg, 1.12 mmol, 2 equiv.); (R)-2-amino-2-phenylacetamide (127 mg, 0.84 mmol, 1.5
equiv.). Product: crystalline solid (74 mg, 37 %). Purity: 88 % (by HPLC). ³¹H NMR (600 MHz, DMSO-d₆) δ 9.32 (br s, 1H, OH), 9.20 (d, J = 7.0 Hz, 1H, NH), 8.26 (t, J = 1.0 Hz, 1H, Ar-H), 7.99 (dd, J = 8.0, 2.0 Hz, 1H, Ar-H), 7.52 – 7.45 (m, 3H, Ar-H), 7.43 – 7.31 (m, 3H, Ar-H), 5.68 (d, J = 7.0 Hz, 1H, -CH=NH), 5.04 (s, 2H, -CH₂OB), 3.66 (s, 3H, -OCH₃); ¹³C NMR (151 MHz, DMSO-d₆) δ 172.2, 166.7, 157.5, 139.3, 133.3, 130.5, 130.4, 128.8, 128.0, 127.9, 121.8, 70.4, 57.3; LCMS (ESI⁺, m/z) calculated for C₁₁H₁₅N₂O₄⁹¹B, [M+H]+: 311, found 311.

Methyl (2R)-2-[(1-hydroxy-3H-2,1-benzoxaborole-6-carbonyl)amino]-2-phenyl-acetate (8)

General Protocol 1 was followed using the following quantities of reagents: 3 (100 mg, 0.56 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1’-carbonyldiimidazole (182 mg, 1.12 mmol, 2 equiv.); methyl (2R)-2-amino-2-phenyl-acetate (139 mg, 0.84 mmol, 1.5 equiv.). Product: crystalline solid (38 mg, 20 %). Purity: >96 % (by HPLC). ¹³H NMR (600 MHz, DMSO-d₆) δ 9.32 (br s, 1H, OH), 9.20 (d, J = 7.0 Hz, 1H, NH), 8.26 (t, J = 1.0 Hz, 1H, Ar-H), 7.99 (dd, J = 8.0, 2.0 Hz, 1H, Ar-H), 7.52 – 7.45 (m, 3H, Ar-H), 7.43 – 7.31 (m, 3H, Ar-H), 5.68 (d, J = 7.0 Hz, 1H, -NHCH), 5.04 (s, 2H, -CH₂OB), 3.66 (s, 3H, -OCH₃); ¹³C NMR (151 MHz, DMSO-d₆) δ 171.6, 167.4, 157.7, 137.6, 136.7, 133.0, 130.7, 129.3, 128.7, 127.6, 70.4, 57.4, 52.8; LCMS (ESI⁺, m/z) calculated for C₁₁H₁₅N₂O₄⁹¹B, [M+H]+: 326, found 326; HRMS (ESI-TOF, m/z) [M + Na]+: 348.1014, found 348.1016.

N, N-Dibenzyl-1-hydroxy-3H-2,1-benzoxaborole-6-carboxamide (15)

General Protocol 1 was followed using the following reagents: 3 (100 mg, 0.56 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1’-carbonyldiimidazole (182 mg, 1.12 mmol, 2 equiv.); dibenzylamine (133 mg, 0.67 mmol, 1.2 equiv.). Product: Crystalline solid (20 mg, 10 %). Purity: >99 % (by HPLC). ¹³H NMR (300 MHz, DMSO-d₆) δ 9.26 (br s, 1H, OH), 7.85 (s, 1H, Ar-H), 7.61 – 7.52 (m, 1H, Ar-H), 7.47 (d, J = 8.0 Hz, 1H, Ar-H), 7.44 – 7.08 (m, 10H, Ar-H), 5.02 (s, 2H, -CH₂OB), 4.75 – 4.24 (m, 4H, 2 x -CH₂Ph), ¹³C NMR (151 MHz, DMSO-d₆) δ 171.6, 167.4, 157.7, 136.7, 133.0, 130.7, 130.6, 129.3, 129.0, 128.7, 121.7, 70.4, 57.4, 52.7; LCMS (ESI⁺, m/z) calculated for C₂₃H₂₆N₂O₃⁹¹B, [M+H]+: 358 found 358; HRMS (ESI-TOF, m/z) [M + Na]+: 380.1429, found 380.1429.
5.2.7.2 General Protocol 2: Synthesis of 9, 10 and 11 (Scheme 5.2A)

**Step (i):** General Protocol 1 was followed to afford an appropriate methyl ester intermediates, B1, which were then directly subjected to saponification. **Step (ii):** To a solution of B1 (1 equiv.) in 1,4-dioxane/water (3:1; 10 mL) was added lithium hydroxide monohydrate (either 4 or 6 equiv.) in one-portion. The reaction mixture was then stirred for 1 hour at 40 °C, before the volatiles were removed *in vacuo*. The residue thus obtained was then lyophilised to afford corresponding free carboxylic acids (confirmed by LC-MS analysis) as solids. **Step (iii):** Crude carboxylic acids were immediately coupled with selected piperazin-2-one derivatives using conditions outlined in General Protocol 1, giving target benzoarboroles as solids.

1-Hydroxy-N-[2-oxo-2-(3-oxo-piperazin-1-yl)ethyl]-3H-2,1-benzoxaborole-6-carboxamide (9)

General Protocol 2 was followed with the following quantities of reagents: Step (i): 3 (250 mg, 1.40 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (455 mg, 2.8 mmol, 2 equiv.) methyl glycinate HCl (133 mg, 0.67 mmol, 1.2 equiv.); step (ii): methyl 2-[(1-hydroxy-3H-2,1-benzoxaborole-6-carbonyl)amino]acetate (B1, 349 mg, 1.40 mmol, 1 equiv.); 1,4-dioxane/water (2.5 ml water, 7.5 ml dioxane); lithium hydroxide (134 mg, 5.61 mmol, 4 equiv.); step (iii): 2-[(1-hydroxy-3H-2,1-benzoxaborole-6-carbonyl)amino]acetic acid (50 mg, 0.21 mmol, 1 equiv.); N,N-dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (69 mg, 0.69 mmol, 2 equiv.); piperazin-2-one (32 mg, 0.32 mmol, 1.5 equiv.). Product: crystalline solid (14 mg). Purity: >97 % (by HPLC). H NMR (600 MHz, DMSO-$_d_6$) δ 9.37 (br s, 1H, OH), 8.63 – 8.56 (m, 1H, NH), 8.33 – 8.20 (m, 1H, Ar-H), 7.96 (dd, J = 8.0, 2.0 Hz, 1H, Ar-H), 7.51 (d, J = 8.0 Hz, 1H, Ar-H), 5.05 (s, 2H, -CH$_2$OB), 4.21 – 4.10 (m, 3H, CH$_2$), 3.96 (s, 1H, CH$_3$), 3.72 – 3.61 (m, 2H, CH$_2$), 3.32 – 3.18 (m, 2H, CH$_2$); C NMR (151 MHz, DMSO-$_d_6$) δ 172.1, 155.5, 135.4, 129.2, 129.2, 129.2, 129.1, 129.1, 127.9, 127.6, 127.5, 122.1, 70.4, 52.1; LCMS (ESI$^+$, m/z) calculated for C$_{10}$H$_{13}$N$_2$O$_2$B, [M+H]$^+$: 318, found 318; (ESI-TOF, m/z) [M + Na]$^+$ : 340.1076, found 340.1076.

(R)-1-Hydroxy-N-(1-(4-methyl-3-oxo-piperazin-1-yl)-1-oxo-3-phenylpropan-2-yl) -1,3-dihydrobenzo[c][1,2]oxaborole-6-carboxamide (10)
General Protocol 2 was followed using the following quantities of reagents: step (i): 3 (100 mg, 0.56 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (182 mg, 1.12 mmol, 2 equiv.); methyl D-phenylalaninate hydrochloride (121 mg, 0.56 mmol, 1 equiv.); step (ii): methyl(1-hydroxy-1,3-dihydrobenzo[c][1,2]oxaborole-6-carbonyl)-D-phenylalanine (B1, 187 mg, 0.55 mmol, 1 equiv.); 1,4-dioxane/water (2.5 ml water, 7.5 ml dioxane); lithium hydroxide (79.23 mg, 3.31 mmol, 6 equiv.); step (iii): (1-hydroxy-1,3-dihydrobenzo[c][1,2]oxaborole-6-carbonyl)-D-phenylalanine (30 mg, 0.09 mmol, 1 equiv.); N,N-dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (30 mg, 0.18 mmol, 2 equiv.); 1-methylpiperazin-2-one (16 mg, 0.14 mmol, 1.5 equiv.).

Product: crystalline solid (16 mg). Purity: >97% (by HPLC). 1H NMR (300 MHz, DMSO-d6) δ 8.85 (br s, 1H, OH), 8.24-8.16 (m, 1H, NH), 7.98 - 7.86 (m, 1H, Ar-H), 7.48 (d, J = 8.0 Hz, 1H, Ar-H), 7.36 - 7.14 (m, 6H, Ar-H), 5.20-4.95 (m, 2H, -CH2OB), 4.20 - 3.93 (m, 2H), 3.34 - 2.96 (m, 5H), 2.80 (s, 3H); LCMS (ESI+, m/z) calculated for C22H24N3O510B, [M+H]+: 422, found 422.

1-Hydroxy-N-[2-(4-methyl-3-oxo-piperazin-1-yl)-2-oxo-ethyl]-3H-2,1-benzoxaborole-6-carboxamide (11)

General Protocol 2 was followed with the following quantities of reagents: step (i): 3 (250 mg, 1.40 mmol, 1 equiv.); dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (455 mg, 2.8 mmol, 2 equiv.) methyl glycinate HCl (133 mg, 0.67 mmol, 1.2 equiv.); step (ii): methyl 2-[(1-hydroxy-3H-2,1-benzoxaborole-6-carbonyl)amino]acetate (B1, 349 mg, 1.40 mmol, 1 equiv.); 1,4-dioxane/water (2.5 ml water, 7.5 ml dioxane); lithium hydroxide (134 mg, 5.61 mmol, 4 equiv.); step (iii): 2-[(1-hydroxy-3H-2,1-benzoxaborole-6-carbonyl)amino]acetic acid (47 mg, 0.2 mmol, 1 equiv.); N,N-dimethylformamide (2 mL); 1,1'-carbonyldiimidazole (65 mg, 0.4 mmol, 2 equiv.); 1-methylpiperazin-2-one (34 mg, 0.30 mmol, 1.5 equiv.). Product: crystalline solid (17 mg). Purity: >96% (by HPLC). 1H NMR (300 MHz, DMSO-d6) δ 8.58 (br s, 1H, OH), 8.26 (s, 1H, Ar-H), 7.97 (d, J = 6.0 Hz, 1H, Ar-H), 7.51 (d, J = 8.0 Hz, 1H, Ar-H), 5.05 (s, 2H, CH2), 4.24 - 4.07 (m, 3H, -CH2OB and CH2), 4.05-3.94 (m, 2H, CH2), 3.48-3.30 (m, 2H, CH2), 2.89 (s, 3H, -NCH3); 13C NMR (151 MHz, DMSO-d6) δ167.6, 167.3, 165.2, 157.5, 133.4, 130.2, 130.0, 121.8, 70.3, 48.0, 47.5, 46.2, 41.2, 33.9; LCMS (ESI+, m/z) calculated for C15H18N3O510B, [M+H]+: 332, found 332;
5.2.7.3 Synthesis of 12 (Scheme 5.2A)

**Step (i):** To a solution of (12a) ( tert-butoxycarbonyl-D-phenylalanine) (300 mg, 1.13 mmol) in N,N-dimethylformamide (3 mL) was added 1,1-carboxyldiimidazole (367 mg, 2.26 mmol); the reaction was stirred for 5 minutes at room temperature. To the reaction mixture was then added piperazin-2-one (170 mg, 1.70 mmol) and the resultant solution was stirred for 4 hours at 40 °C. Ethyl acetate (10 mL) and water (10 mL) were added. The layers were separated. The aqueous layer was extracted with ethyl acetate (3 × 20 mL). Combined organic layers were dried (MgSO₄), filtered, and concentrated in vacuo to afford 12b as a yellow oil (350 mg, 88 %), which was used in the next step without further purification.

**Step (ii):** To a stirred solution of (12b) (350 mg, 1.01 mmol, 1 equiv.) in CH₂Cl₂ (10 mL) was added HCl (4M in 1,4-dioxane, 0.76 mL, 3.02 mmol, 3 equiv.) at room temperature under nitrogen. The reaction mixture was stirred overnight to afford an insoluble precipitate. The precipitate, 12c, was washed three times with CH₂Cl₂, then filtered, dried in air and used in the next step without further purification (54 mg, 19 %).

**Step (iii): General Protocol 1** was followed using the following quantities of reagents: 3 (54 mg, 0.3 mmol, 1 equiv.); N,N-dimethylformamide (3 mL); 1,1-carboxyldiimidazole (98 mg, 0.600 mmol, 2 equiv.); 12c (103 mg, 0.360 mmol, 1.2 equiv.). Product (12): white powder (20 mg, 16 %). Purity: >99 % (by HPLC).

**¹H NMR** (300 MHz, methanol-d₄); δ 8.31 (br s, 1H, OH), 8.14 (dd, J = 8.0, 2.0 Hz, 2H, Ar-H), 7.93 (d, J = 8.0 Hz, 1H, Ar-H), 7.56 – 7.45 (m, 2H, Ar-H), 7.36 – 7.22 (m, 3H, Ar-H), 5.15 (m, 5H, 2 x CH₂ and CH₃), 4.33 – 3.96 (m, 2H, CH₂), 3.88 – 3.40 (m, 3H, CH₂), 3.28 – 3.06 (m, 3H, CH₂); **¹³C NMR** (151 MHz, DMSO-d₆) δ 170.5, 167.3, 167.3, 157.8, 132.1, 131.9, 130.3, 129.6, 128.7, 127.1, 122.5, 70.4, 51.5, 46.1, 42.3, 40.4, 37.5; **LCMS** (ESI⁺, m/z) calculated for C₂₁H₂₈N₂O₅⁺⁹B, [M+H]⁺: 408, found 408; **HRMS** (ESI-TOF, m/z), [M + Na]⁺: 430.1543, found 430.1546.
5.3 Results

5.3.1 X-ray Fragment Screen

A small compound library was assembled to be screened against PaPBP3 on the XChem platform. The library (261 compounds in total) was selected to include a small number of known electrophilic warheads (nitriles, epoxides, sulphonyl fluorides and esters), but was especially enriched with boron-containing compounds (Table 5.2). Most of the compounds were selected from Enamine’s “Serine focused Covalent Fragments” library 63. Within each of the chemotype groups, selections from the catalogue were made to make the most chemically diverse set possible. The library generally conformed to the rule of 3 of fragment libraries 64 (after Lipinski’s rule of 5 65) (Figure 5.2), although this may not be as relevant to covalent fragment libraries, which may be more dependent on the chemistry of the warhead.

![Graphs showing properties of the fragments screened](image)

**Figure 5.2. Properties of the fragments screened.** The fragments screened mostly conformed to the ‘rule of 3’ 64, such that the molecular mass was <300 Da (a), the calculated log of partition coefficient (XLogP) 62 (b) and hydrogen bond donors (d) are <3. The number of hydrogen bond acceptors (predicted that <3 is best) (c) is the least compliant, but has a median of 4 hydrogen bond acceptors.
Boron-containing compounds are known to exhibit pH-dependent behaviour, so the library was screened against PaPBP3 once with the crystals at pH 6 and once with the crystals at pH 8.

<table>
<thead>
<tr>
<th>Chemotype</th>
<th>Number of compounds incorporating chemotype</th>
<th>Hits found by XChem</th>
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<tr>
<td>R≡N</td>
<td>Nitrile</td>
<td>5</td>
</tr>
<tr>
<td>△O</td>
<td>Epoxide</td>
<td>4</td>
</tr>
<tr>
<td>R−S−F</td>
<td>Sulphonyl fluoride</td>
<td>4</td>
</tr>
<tr>
<td>A−B</td>
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<td></td>
</tr>
<tr>
<td>A = aromatic C</td>
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<tr>
<td>A = aliphatic C</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>≡N</td>
<td>Benzoxaborole</td>
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</tr>
<tr>
<td>≡N</td>
<td>6,6-bicyclic boronate</td>
<td>16</td>
</tr>
<tr>
<td>≡N</td>
<td>Ester</td>
<td>92</td>
</tr>
</tbody>
</table>

Table 5.2. Electrophilic fragments screened against PaPBP3. * Some compounds incorporate multiple chemotypes. b Boronic acids bonded to aliphatic or aromatic carbons are distinguished. The boronic acid chemotype is explicitly defined as two free hydroxyls, such that benzoxaboroles or 6,6-bicyclic boronates are not a subset of this group. A complete list of fragments screened is given in Table 5.S1.

In total, 34 ‘hits’ were identified by the screen, although not all had complete density for the ligand. All hits were from the boronic acid (26 hits, all tri-covalent) or benzoxaborole (7 hits, all di-covalent) chemotypes, and all showed covalent bonding to the protein through the boron atom. The correlation of the valency of the binding and the chemotype was striking. Tri-covalent boronate binding has been
described in LMM PBPs\textsuperscript{16}, and di-covalency has been previously reported for other boronate compounds\textsuperscript{27,28} (Table 5.1), but this is the first time di-covalency has been reported for benzoxaboroles and the first time multi-covalent interactions have been observed in HMM PBPs.

5.3.2 Fragment Hit Examples

![Diagram of fragment hits example](image)

\textbf{Figure 5.3. Tri-covalency is observed in the reaction between boronic acids 1 and 2 and PaPBP3 by crystallography.} Boron atoms in (a) 1 and (b) 2 are ligated by Ser294, Ser349, and Lys484. Residues neighbouring the ligand are as shown in sticks and are labelled, hydrogen bonds are represented by black dashed lines. A network of hydrogen bonded waters (w) is also shown. Composite omit maps (contoured at 1σ) are shown as light blue mesh around the ligand and covalently bonded residues. A DMSO molecule is also found in the active site of both structures, but omitted for clarity.

A selection of the fragment hits which were representative of the binding modes were investigated in more detail. Boronic acids 1 and 2 react with PaPBP3 such that the boron becomes \(sp^3\) hybridised and tri-covalently bonded to Ser294 (the ‘active site’ or ‘catalytic’ serine in the SXXK motif), Ser349 (in the SXN motif), and Lys484 (in the KSG motif), forming a tri-covalent complex Figure 5.3. The aromatic groups of 1 and 2 occupy similar, but not identical, regions of the active site. Ser294 is displaced to the right by rotation of its side chain bonds, compared to its position when reacting with \(\beta\)-lactams. This allows a water to occupy the
‘oxyanion hole’ which the β-lactam-derived carbonyl often occupies. The tetrazole
group of 1 makes four hydrogen bonds to PaPBP3: two to the backbone nitrogens
of Gly534 and Gly535; two to the flanking residues Thr487 and Ser485. Residues
Thr487 and Ser485 are typically engaged by the C-3 penicillin carboxylate (or
equivalent group) found on all β-lactams.

Hydrogen bonds are also formed with the backbone nitrogens of Gly535 and the
Ser485 by the imidazole group of 2. These structures therefore provide support for
future design of PBP3 inhibitors incorporating either weakly acidic non-carboxylates
or neutral groups 67 that interact with the ‘acid binding pocket’ of PBP3 (Figure
5.S2).

A similar binding mode is reported for a boronate bound to R39 0,0-peptidase
from Actinomadura sp. (PDB: 3ZVT 16). In particular, the boron atoms and bonded
side chains are similarly positioned and superimpose well, regardless of the enzyme
and the different boron-bonded functional group, i.e. alkyl (Zervosen et al.16) versus
phenyl (this work) (Figure 5.S2d).

In contrast to boronic acids, benzoaborole fragments bind to PaPBP3 with a
di-covalent binding mode (Figure 5.4). The boron engages both Ser349 and Ser294
and the benzoaborole also hydrogen bonds through the ‘endo cyclic’ 2-position
oxygen (atom numbering shown in Figure 5.5) to Lys484. As with the phenyl boronic
acids, a water is found in the oxyanion hole. Both 3 and 4 form hydrogen bonds to
the sidechain NH2 group of Asn351, and both PaPBP3:3 and PaPBP3:4 complexes
show hydrogen bonding between Tyr409 and the backbone carbonyl of Thr487
(Figure 5.4).
Figure 5.4. Benzoazaborole fragments form di-covalent complexes with PaPBP3. Crystal structures of (a) PaPBP3:3 and (b) PaPBP3:4 complexes. The boron atom of both molecules reacts with the side chains of Ser294 and Ser349. Hydrogen bonds (black dashed lines) are formed with Lys484 and Asn351. A water (w) occupies the oxyanion hole. Composite omit maps (light blue mesh, maps contoured at 1σ) are shown for the ligand and the bonded serine sidechains.

5.3.3 Hit Expansion: Design and Synthesis

The novel binding mode of benzoazaboroles was investigated by the generation of a small number of chemical derivatives in an attempt to improve their affinities. The derivatives were designed using computational docking starting from fragments 3 and 4 as well as the complexes of PaPBP3 with β-lactams. It was hoped compounds would interface with the protein at the regions engaged by reacted piperacillin (Figure 1.4e, Figure 5.5). From examination of the PaPBP3:3 complex it was hypothesised that substitution of the benzoazaborole at the 6-position would allow side chain conformations which aligned closely to the reacted piperacillin R1 (C-6) side chain (Figure 5.5).
Figure 5.5. Binding modes of reacted piperacillin and benzoxaboroles 12 (predicted) and 8 (observed) complexed with PaPBP3. (a) Reacted piperacillin (PDB: 6R3X) engages multiple regions of PaPBP3 (Figure 1.4e), as observed by crystallography. (b) Predicted binding mode of 12 determined by docking, showing the regions of PaPBP3 which it was hoped would be engaged by parts of the sidechain, in analogy with piperacillin. (c) Crystallographically observed binding modes of 8 in complex with PaPBP3. Hydrogen bonds (black dashed lines) are shown. A composite omit map (light blue mesh, map contoured at 1σ) is shown for 8 and the bonded serine sidechains. Functional groups of piperacillin (d) and 12 (e): Serine binding region (orange), C-3 carboxylate (pink), group 1 (blue) and group 2 (green). Atom numbering on the penicillin and benzoxaborole cores is given.

Initial derivatives (5-8) were produced by coupling C-terminally protected amino acids (glycine, p-phenylglycine and p-phenylalanine) to the C-6 carboxyl group of 3, with additional derivatives (9-12) incorporating a ketopiperazine group at the C-terminal end of the first amino acid. 15 was synthesised with a C-6 dibenzylamine group in order to investigate the binding of a non-amino acid benzoxaborole. All reactions were carried out by amide coupling with 1,1'-carbonyldiimidazole (CDI). Purification was challenging, and several other compounds which had been designed could not be purified after synthesis, a known challenge for benzoxaboroles.

Initial crystal structures (as exemplified by 8) of benzoxaboroles 5-12, showed that whilst all derivatives bound in the same di-covalent binding mode, the benzoxaborole side chains were angled incorrectly for engagement with the active site as proposed.
The C-3 carboxylate is ubiquitous in β-lactams, so derivatives incorporating this group were investigated (13, 14 and 16 were synthesised by WuxiAppTec and collaborators). The incorporation of a second acidic group requires a more complex synthetic route involving ring closure of the benzoxaborole 5 membered ring. The presence of the acid group did not affect the position of the benzoxaborole ring within the active site of the protein and 3, 4, 8, and 13 - 16 all showed di-covalent binding. 16, which is derivatised at the 5-position, offers new avenues for further synthesis. Side chains originating from this point may be better positioned to form piperacillin-like interactions with the protein (Figure 5.6d). Similarly the non-amino acid 15 also exploits a region of the protein that is not occupied by the amino acid derivatives e.g. 3, 4, 8, 13 or 14 (Figure 5.6c).
Figure 5.6. PaPB3 complex with C-3 acid group containing benzoxaboroles (13, 14 and 16) and 15. Crystal structures of (a) the PaPB3:13 complex; (b) the PaPB3:14 complex; (c) the PaPB3:15 complex; (d) the PaPB3:16 complex. There are two alternative conformations of Tyr409 in the PaPB3:13 complex but only one is shown in (a). Both conformations are shown in Figure 6.S1b. Relevant hydrogen bonds (dashed black lines) are shown. Composite omit maps (light blue mesh, maps contoured at 1σ) are shown for the ligand and the bonded serine sidechains.
5.3.4 Vaborbactam

Vaborbactam, a boron-based serine β-lactamase inhibitor \(^{5-7}\) lacking antimicrobial activity, was also investigated alongside the benzoxaboroles to allow the behaviour of these different scaffolds to be compared. The PaPBP3:vaborbactam complex shows that vaborbactam reacts with Ser294 mono-covalently (Figure 5.7).

Figure 5.7. The PaPBP3:vaborbactam complex as observed by crystallography. (a) Vaborbactam bonds covalently to Ser294 and engages the active site with a number of hydrogen bonds (black dashed lines). Composite omit maps (light blue mesh, maps contoured at 1σ) are shown for vaborbactam and the bonded serine sidechain. (b) Overlay of vaborbactam (purple) and amoxicillin (PDB code: 6f1E, grey)\(^{36}\) binding PaPBP3. Residues neighbouring the ligand are shown in the same colour as the ligand and are labelled. The overlay shows considerable alignment of both residue side chains and reacted ligand conformations.

Comparison of the PaPBP3:vaborbactam complex with the structure of amoxicillin reacted PaPBP3 shows they superimpose well, particularly at the the C-3 carboxylate, the nucleophilic serine and the R1 side chain of amoxicillin (Figure 5.7b). Hydrogen bonds are formed between vaborbactam and residues Asn351, Tyr409, Thr487, Ser485, and the hydrophobic wall (residues Tyr503, Tyr532 and Phe533, Figure 1.4e) forms around the thiophene group.
5.3.5 Inhibition Assays

The ability of 1-16, as well as vaborbactam, to engage PaPBP3 in solution was assessed by competition with BOCILLIN FL in a fluorescence anisotropy (FA) assay and an assay of PBP-mediated thioester S2d hydrolysis \(^{13, 51-53}\) (Table 5.3, Figure 5.8). The results obtained by both assays were consistent with each other, but most compounds exhibit little inhibition even at the high concentration tested (1 mM). Dissociation constants \((K_d)\) were determined for 13, 14, 16, and vaborbactam, which had the lowest residual activities in both assays (Figure 5.8a-d). The most potent was found to be 13 (which is a racemate) with a \(K_d\) of 73.9 ± 0.8 μM (Table 5.3). Vaborbactam showed only weak affinity, despite its crystallographic homology with amoxicillin and apparently successful engagement with the protein. Overall binding is weak, which is consistent with other reports of μM affinities for boronates binding to HMM PBPs \(^{12-14, 21, 22}\).
Figure 5.8. Progress curves of boronate inhibition of PaPBP3. (a-d) BOCILLIN FL fluorescence anisotropy competition curves are shown for (a) 13, (b) 14, (c) 16 and (d) vaborbactam reacting with PaPBP3. Kintek Global explorer was used to fit the fluorescence anisotropy progress curves in the presence of increasing concentrations of each inhibitor to the model in Scheme 5.1. (e) Progress curves for determining residual activities by PaPBP3-mediated S2d turnover, following the absorbance of DTNB at 412 nm. (f) Progress curves of the inhibition of PaPBP3-mediated nitrocefin turnover by compound 13 to investigate reversibility. The rates were determined before and after the assay volume was doubled (at the indicated point), then used to create the pIC₅₀ curve shown in Figure 5.9.
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<th>Residual Activity by S2d Turnover (%)</th>
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<tr>
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<tr>
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<td>16 ± 6</td>
<td>19.6 ± 0.4</td>
<td>73.9 ± 0.8</td>
</tr>
<tr>
<td>14</td>
<td>42 ± 6</td>
<td>47.5 ± 0.6</td>
<td>172.0 ± 3.0</td>
</tr>
<tr>
<td>15b</td>
<td>87 ± 3</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>16</td>
<td>20 ± 5</td>
<td>ND</td>
<td>78.1 ± 0.9</td>
</tr>
</tbody>
</table>

Vaborbactam 54 ± 5 75.8 ± 0.7 201.0 ± 2.3

Table 5.3. Activity profiles of 1-16 and vaborbactam against PaPBP3. Residual activities were determined in the presence of 1 mM inhibitor and expressed as a percentage of the activity of the untreated control. Errors are standard errors (n = 3) from independent measurements. $K_i$ determined using global fitting in Kintek Global Explorer, errors are standard errors generated by fitting. *Residual activity measured at 100 μM due to insolubility at 1 mM. ND, not determined.
13 was assessed for its ability to bind class B PBPs from *E. coli*, *A. baumannii*, *H. influenzae*, and *N. gonorrhoeae* (Table 5.4) by competition with BOCILLIN FL. The apparent differences in the ability of 13 to bind each PBP indicates a degree of selectivity of the compound, given the similarity between these proteins (Figure 1.3).

<table>
<thead>
<tr>
<th>Protein Tested</th>
<th>Residual Activity by BOCILLIN FL FA (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBP3 from <em>P. aeruginosa</em></td>
<td>16 ± 6</td>
</tr>
<tr>
<td>PBP3 from <em>H. influenzae</em></td>
<td>32 ± 3</td>
</tr>
<tr>
<td>PBP3 from <em>A. baumannii</em></td>
<td>73 ± 3</td>
</tr>
<tr>
<td>PBP3 from <em>E. coli</em></td>
<td>&gt;90</td>
</tr>
<tr>
<td>PBP2 from <em>N. gonorrhoeae</em></td>
<td>&gt;90</td>
</tr>
</tbody>
</table>

Table 5.4. Inhibition of various class B PBPs by 13. Residual activities were determined in the presence of 1 mM 13 and expressed as a percentage of the activity of the untreated control. Errors are standard errors (n = 3) from independent measurements. *N. gonorrhoeae* PBP2 was a transpeptidase domain only construct.

The interaction of 13 with PaPBP3 was further confirmed in assays with nitrocefin (Table 1.2), which allowed a $pIC_{50}$ to be found (Figure 5.9). The reversibility of the binding of 13 could also be assessed by this method: determining the half maximal inhibitory concentration $pIC_{50}$ before and after a two-fold dilution of the assay solution showed a doubling of the $pIC_{50}$. Which indicates the reaction was rapidly reversible (equilibrium established in <30 s). Irreversibly binding ceftazidime does not exhibit a change in $pIC_{50}$ following the dilution of the assay. Similarly, pre-incubating 13 with PaPBP3 for 0, 30 or 60 minutes prior to initiation of the BOCILLIN FL assay appeared to have no significant effect on level of inhibition and the BOCILLIN FL fluorescence anisotropy progress curves can be fit well using a simple reversible, one step binding model (Scheme 5.1).
Figure 5.9. The effect of 2-fold dilution on the PaPBP3 pIC$_{50}$ of 13 and ceftazidime. The rate of nitrocefin turnover (measured at 482 nm) by PaPBP3 in the presence and absence of inhibitors was determined before and after a 2-fold dilution of the assay and the rates compared to the uninhibited control. Errors shown for each point are standard deviations from three repeats, errors on pIC$_{50}$ values are standard errors of the mean as determined by Prism 9 (Prism 9 for macOS, Graphpad Software, LLC).
5.3.6 Microbiology

Several compounds were screened against wild type *E. coli*, *P. aeruginosa*, *H. influenzae*, *A. baumannii* and *N. gonorrhoeae* strains. All were ineffective (MICs ≥ 64 µg/ml) (Table 5.5). Screening against *E. coli ΔToIC* (Chapter 3.3.6) or a *P. aeruginosa* strain engineered to remove the outer membrane permeability barrier by the introduction of a large pore \(^7\) did not show any antimicrobial activity (Table 5.5), indicating the lack of activity was not due to permeability issues. Synergy with piperacillin (which can be used to identify weak PBP inhibition) was not observed in non-β-lactamase-expressing *E. coli* and *P. aeruginosa*.

A separate screen of a library of phenyl boronic acids was conducted by the robotic screening platform (Chapter 3), which included all of the compounds in this study (except vaborbactam). This screen (against *E. coli ΔToIC*) did not identify any hits amongst the compounds in this chapter (Table 5.5).

<table>
<thead>
<tr>
<th>P aeruginosa</th>
<th>N CTCC93022</th>
<th>N CTCC43228</th>
<th>P aeruginosa Permeabilised</th>
<th>N CTCC43228</th>
<th>A baumannii ATCC 19660</th>
<th>H influenzae ATCC 49760</th>
<th>P aeruginosa Synergy</th>
<th>P aeruginosa Synergy</th>
<th>E coli NCTC 27/96</th>
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</tr>
<tr>
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<td>(&gt;64 \mu g/\text{mL})</td>
<td>(&gt;64 \mu g/\text{mL})</td>
<td>(&gt;64 \mu g/\text{mL})</td>
<td>(&gt;64 \mu g/\text{mL})</td>
<td>(&gt;64 \mu g/\text{mL})</td>
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<td>No Effect</td>
<td>NT</td>
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Table 5.5. Minimum inhibitory concentrations of selected compounds for a panel of gram-negative bacteria. *Permeabilised strains with the introduction of a FhuA pore and knockout of export pumps* \(^7\).
5.4 Discussion

5.4.1 Boron Binding Modes

Complexes of boronates with PaPBP3 reveal three distinct binding modes: (i) mono-covalent (reaction with Ser294; Figure 5.10a), (ii) di-covalent (reaction with Ser294 and Ser349; Figure 5.10b), and (iii) tri-covalent (reaction with Ser294, Ser349 and Lys484; Figure 5.10c). The binding mode correlates with the nature of the boron compound: monocyclic vaborbactam bind mono-covalently, benzoxaboroles (3-16) bind di-covalently, and phenyl boronates (1 and 2) bind tri-covalently. Tri-covalent \(^{16}\) and mono-covalent \(^{11,14-16,18,19}\) interactions of boron compounds with PBPs are known, but to my knowledge there are no reports on the di-covalent bonding of benzoxaborole compounds to PBPs. Crystallographic data was collected at pH 6 and pH 8 but the binding mode doesn't appear to relate to the crystallisation conditions, only to the warhead (Table 5.S3).
Figure 5.10. Three distinct binding modes of complexes of PBPs with boron-containing compounds. (a-c) Outline mechanisms for the formation of the different complexes. (a) Vaborbactam reacts mono-covalently with Ser294 of PaPBP3. (b) Benzoaxaboroles (3-16) bind di-covalently to Ser294 and Ser349 of PaPBP3. (c) Phenyl boronic acids (1 and 2) bind tri-covalently to Ser294, Ser349 and Lys484 of PaPBP3, similarly to alkyl boronic acids reacting with R39 DD-peptidase.
from *Actinomadura* sp.\(^{16}\). Boron must transition between \(sp^2\) and \(sp^3\) hybridisation states (highlighted in yellow and pink, respectively, Figure 5.1) to exchange hydroxyls for the nucleophilic side chain. The order in which nucleophilic side chains covalently react with the boron is unknown; (d) Two views of the overlay of crystallographically observed states of different boronates binding to PaPBP3, the lysine N\(_e\) is labelled. (e-g) The position of the boron atom within the active site relates to rotations of the chi1 and chi2 angles of the Ser294 sidechain from PaPBP3 (e). Ser349 (which binds to the boron of 13 and 1) and Lys484 (which binds to the boron of 1) are not shown. (e) Atom and angle labels of serine. (f) Newman projection aligned along the Ca-C\(\beta\) bond of Ser294 of PaPBP3 and crystal structures of vaborbactam and 1. Like a \(\beta\)-lactam reacted structure, the mono-covalently bonded vaborbactam structure has a Ser294 chi1 angle gauche’ to the serine backbone amine, whilst the tri-covalent (1) complex with PaPBP3 has a chi1 angle trans to the serine backbone amine. The di-covalently reacted PaPBP3:13 also has a gauche’ chi1 angle (Figure 5.S3), but unlike the mono-covalent structures, it has a Ser294 chi2 angle of -171°. A complete set of structural views and Newman projections are shown in Figure 5.S3.

Structures of 3, 4, 8, 10, 13, 14, 15, and 16 with PaPBP3 show a consistent positioning of the benzoxaborole relative to the active site, irrespective of C-6, C-5 or C-3 functionalisation (Figure 5.11). This is likely caused by the constraint of having both serines reacted to the boron.

![Figure 5.11. Overlay of various benzoxaborole conformations observed in complex with PaPBP3. Crystal structures of PaPBP3 with benzoxaborole compounds 3 (brown), 4 (orange), 8 (pink), 13 (green), 14 (purple), 15 (cyan), 16 (blue), reveal that they bind in a conserved mode. The protein structure is hidden for clarity. All compounds bind di-covalently: the tetrahedral boron is reacted with the hydroxyl groups of residues Ser294 and Ser349 (not shown for clarity) of the active site.](image)

Analysis of the bond angles of the nucleophilic residues (Ser294, Ser349 and Lys484) shows how each of the different binding modes affects the protein (Figure 5.10 and Figure 5.S3). In each of the binding modes, the side chains of
Ser349 and Lys484 are not displaced significantly, with the Oy of Ser349 and the Nc of Lys484 moving <1 Ångström relative to their position in the piperacillin reacted structure PaPBP3 (PDB: 6R3X). In contrast, Ser294 has a number of rotations of the first and second bonds of the side chain (described by dihedral angles chi1 and chi2, Figure 5.10e). The PaPBP3 structures of mono-covalently reacted β-lactams (e.g. piperacillin) and vaborbactam, as well as di-covalently reacted benzoxaboroles (3, 4, 8, 10, 13, 14, 15, and 16) all have a Ser294 chi1 that is gauche’ (~ -60°) to the serine amine (as defined in reference 71), whilst the tri-covalently reacted structure PaPBP3:1 has a trans chi1 angle (Figure 5.10f and Figure 5.S3). The Ser294 chi2 angle of the di-covalently reacted structures however is almost orthogonal to the chi2 of the mono-covalently reacted structures (-171° vs 109°, relative to the Ca), which displaces the boron to the right (as depicted in Figure 5.10d, left hand side image).

The previous report of a boronate engaging a PBP in a tri-covalent binding mode suggested that binding to the catalytic Ser49 (equivalent to Ser294) occurs first, with the other nucleophilic substitutions happening subsequently 16. Our data cannot be used to determine an order of reactions. Tri-covalent substitution of benzoxaboroles with a reaction by Lys484 is prevented by the strength of the endocyclic B1-O2 bond of the benzoxaborole 5-membered ring 72,73.

As depicted in Figure 5.10, Lys484 must be neutrally charged to act as a nucleophile and form the observed tri-covalent complex. If a lysine is buried and in a hydrophobic region of the active site, it is possible for its pKₐ to become more acidic and low pKₐ lysines have been described in the literature 74–76. The equivalent neutral lysine has previously been proposed to act as a general base in the β-lactam de-acylation of PBP5 18, although generally the lysine equivalent to Lys297 is proposed as the general base in the PBP active site 79–81.

The initial data presented here indicates that binding equilibrium is achieved at least within minutes. Previous reports have suggested that boron binds with a fast, weak reversible mode, which agrees with this data 20. The time resolution of the dilution method used is not good enough to observe behaviour on the seconds timescale, but work on the tri-covalent PBP:boron complex has suggested that the interaction is a rapid equilibrium association followed by a slower phase in which the covalent reaction occurs and the mono-covalent complex forms 16. Following this, it was suggested the tri-covalent complex forms in rapid equilibrium with the
mono-covalent complex (Figure 5.10c) \(^{16}\). As shown in Figure 5.1, boronates are thought to act as analogues of the PBP tetrahedral transition state \(^{17-19}\). The relevance of the multi-covalent complexes to this mimicry is unclear, and the evidence here is not enough to conclusively determine whether the multi-covalency affects the potency; it may only be a crystallographic artefact.

### 5.4.2 Structural Views

The benzoxaboroles were designed to engage the hydrogen bond network which piperacillin exploits when reacting with PaPBP3 (Figure 5.5), but the constrained benzoxaborole ring (Figure 5.11) prevented the formation of important active site hydrogen bonds. In particular, the hydrogen bond to Thr487 (which is formed by the C-6 amide of β-lactams (Figure 5.S4)) was not made. The conformation of the β3 β-strand is thought to be associated with formation of the hydrophobic wall formation (Figure 1.4) \(^{68}\), perhaps by interactions of the inhibitor with Thr487 and Arg489. The β3 strand of the PaPBP3:13 has a similar conformation to the β3 strand in the meropenem reacted PaPBP3 structure (Figure 5.12a). Additionally, in these structures Tyr409 forms a hydrogen-bond to the backbone carbonyl of Thr487. In contrast, when amoxicillin, aztreonam, ceftazidime and piperacillin react with PaPBP3, the inhibitors form a hydrogen-bond to the backbone carbonyl of Thr487 (Figure 5.S4). The constrained benzoxaborole core positions the C-6 amide ‘above’ the C-6 amide of reacted piperacillin and the larger distance prevents the benzoxaborole amide from interacting with Thr487 ((Figure 5.12b and c). A C-5 amide may be better positioned to engage this residue. A hydrogen bond is made by the C-6 amide to Asn351 (e.g. Figure 5.4b), but a C-5 amide may be unable to form this interaction. Interaction with Asn351 may be important as it is found in many β-lactams (e.g. piperacillin, Figure 5.5a).
Figure 5.12. Comparisons between PaPBP3 reacted with meropenem and 13 reveal similar conformation for the β3 strand, unlike that of the piperacillin reacted crystal structure. (a) Active site of the PaPBP3:meropenem complex (beige, PDB code: 3PBR) compared to the PaPBP3:13 complex (cyan). (b and c) Two views of the β3 strand conformation of the PaPBP3:13 complex (cyan) compared to the meropenem (beige) and piperacillin (black, PDB code: 6R3X) reacted structures. Tyr409 and its hydrogen bond (dashed line) to the backbone amine of Thr487 is shown. A hydrogen bond between Thr487 and the reacted piperacillin amine is shown. The protein structure shown is the PaPBP3:13 complex, but the β3 strand of the piperacillin reacted structure is overlaid. The conformation of the β3 strand in the PaPBP3:13 or meropenem complexes is unlike the PaPBP3 β3 strand conformation seen when reacted with other β-lactams (piperacillin, ceftazidime, amoxicillin, aztreonam): Figure 5.S4.

The β3 strand and β5-α11 loops of the PaPBP3:14 complex have conformations analogous to those of the piperacillin reacted PaPBP3 structure (Figure 5.13). 14 has a ketopiperazine substituent (designed to mimic the diketopiperazine of piperacillin), but lacks a phenyl group analogous to that of the d-Phe of piperacillin. Like the diketopiperazine of piperacillin, the ketopiperazine of 14 forms hydrogen bonds with the On oxygens of Tyr328 and Tyr407 (Figure 5.13), it is unclear whether there is link between the hydrogen bonds and the β3 strand conformation.

One clear conclusion from comparing the compound structures and their affinity is that the presence of a C-3 acid group is important. The affinities of 14 ($K_i = 172.0 \pm 3.0 \mu M$) and 11 (residual activity by BOCILLIN FL FA > 90 %) or 13 ($K_i = 73.9 \pm 0.8 \mu M$) and 7 (RA > 90 %) nicely demonstrate the effect of a C-3 acid group (Table 5.3). This is consistent with the conservation of the C-3 (or equivalent) carboxylate group in β-lactams as well as studies which show that the addition of a C-3 acid group to benzoxaboroles leads to up to 100 fold increases in β-lactamase binding affinity 82.
Figure 5.13. Comparisons between crystal structures of PaPBP3 reacted with piperacillin (black, PDB code: 6R3X<sup>53</sup>) and 14 (orange). (a) The protein backbone conformation is similar in both structures, particularly the β3 strand and the β5-α11 loop. The β3-β4 loop is incomplete in the PaPBP3:14 complex, likely due to flexibility of this region (Chapter 4). (b) The hydrogen bonding network (dashed lines) around the ketopiperazine group of 14 and the diketopiperazine of piperacillin with Tyr328 and Tyr407. Waters from the PaPBP3:14 complex (W<sub>14</sub>) and the piperacillin reacted structure (W<sub>p</sub>), are shown.

The β5-α11 loop is a flexible region that undergoes induced fit upon β-lactam binding and forms a “hydrophobic wall” (Chapter 4). Several of the benzoaboroles (10, 12, 13, 15 and 16) were designed such that their phenyl groups might engage the hydrophobic wall. In all cases this was unsuccessful and the wall was not formed (Figure 5.14). The structures of the PaPBP3:15 and PaPBP3:16 show a phenyl is situated in the same region of the active site as the phenyl rings of reacted piperacillin, but with a different orientation of the ring. The phenyl of 13 sits at a distance from the expected position of the hydrophobic wall. It is possible that designing compounds that better exploit this part of the protein would improve their affinity, although it is not essential to inhibition of the protein as inhibitors such as meropenem do not lead to wall formation (PDB: 3PBR)<sup>68</sup>.
2 induces a novel conformation of both the PaPBP3 α10-β3 and β5-α11 loops and ligands 1 and 14 also modify the β5-α11. These changes can be used to understand the conformational flexibility of the protein; this is discussed further in Chapter 4.

5.5 Future Work and Conclusions

The benzoaborole compounds investigated were not very potent in activity but this study demonstrated their potential as a new class of PBP inhibitors. Future studies investigating benzoaboroles will be able to benefit from the structural data presented here, in particular the observation that the benzoaborole warhead binds to the protein in a highly constrained and consistent manner (Figure 5.11), a conformation which appears to be relatively insensitive to modification at three different positions of the benzoaborole (C-3, C-5 and C-6). This fact, and some of the insights gained from the crystal structures (section 5.3.7) could be used to generate further iterations of the compound series, with the aim of improving affinity. Derivatisation of the C-4, and C-7 positions, and further exploration of the C-5 positions may be interesting and lead to improved potency.

High affinity HMM PBP boron-based inhibitors have yet to be discovered 11-14, with only a single example in the academic literature of a PBP-targeting,
boron-containing compound having antimicrobial activity. The most potent (and with antimicrobial activity) are 6,6-bicyclic boronates (a taniboractam-like scaffold, Table 1.1) in the patent literature. The efficacy of boronates against the structurally related β-lactamases offers hope for the discovery of more effective boronic PBP inhibitors, but an explanation for the much weaker affinities of reported boronates against PBPs is not yet forthcoming.

The lack of antimicrobial activity is disappointing. This can be explained to some degree by the compounds' relatively low on-target affinity, perhaps coupled to poor cellular accumulation (although the 'permeabilised' E. coli and P. aeruginosa strains showed no increase in activity). An alternative hypothesis posits that the mode of binding is important for inhibition of PBP cellular functioning and the apparent rapid equilibrium binding of these boronates is not sufficient to prevent the binding of the polymeric PBP3 substrate. In this scenario, the natural substrate is fed continuously into the PBP3 activity site at a high effective local concentration, easily displacing any bound boronate. In contrast, the effective irreversible inhibition with a β-lactam entirely prevents natural substrate binding and transpeptidation. It may be possible to use novel natural substrate assays to test the ability of boronates to prevent transpeptidation and create new benzoaborole derivatives which do this more effectively, thereby improving their antimicrobial activity.

The three binding modes of boronates engaging with HMM PBPs have not previously been reported, but the binding modes have precedence in other proteins. More work is needed to understand the significance of these observations for inhibitor design and their importance for potency. It might be expected that the multivalent reaction increases the potency, but it is not possible to make this conclusion from the data. A better understanding of the time-dependence of the reaction may be beneficial. Mass spectrometry or NMR (particularly 11B NMR) studies would perhaps be the best approach for this as these techniques are able to probe the hybridisation state of the boron; mass spectrometry would be able to identify the mass change associated with the loss and gain of water in the transition to multivalency (e.g. Figure 5.10b). A working SPR system (Chapter 2) would provide an insight into the kinetics of the reaction but is unlikely to give the same level of detail about the state of the boron atom. Kinetic studies employing a S349A mutant of PaPBP3 (Chapter 6) could be used to study the binding of benzoaboroles under conditions in which only a single serine is available in the active site.
The pKa of a boronate is a measure of its ability to transition between the sp\(^2\) and sp\(^3\) states (Figure 5.15). It has been shown that substitutions on the boronaborole can modify the boronate pK\(_a\) by changing the stability of the anionic (sp\(^3\)) species \(^{84,85}\), which in turn can affect a compound’s covalent \(^{85}\) and non-covalent interactions \(^{86}\). Given the likely importance of the covalent interaction for the binding of boronaboroles to PBPs, optimisation of the pKa of the boron may be important as optimisation of the non-covalent interactions of the sidechains. It may be difficult to distinguish the contributions of the two components, as any additional group will also contribute to the non-covalent interactions of the compound.

In the next chapter the possibility that the di-covalently bound boronaborole is acting as a transition state analogue in the transfer from one active serine to another is discussed. It appears that boronate compounds may have many uses in the inhibition and study of PBPs.

5.6 References


41 Emsley P, Lohkamp B, Scott WG, Cowtan K. Features and development of
78 Isom DG, Castanedo CA, Cannon BR, Garcia-Moreno E. B. Large shifts in pKa values of lysine residues buried inside a protein. Proc Natl Acad Sci 2011; 108:


### 5.S2 Crystallography Statistics

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Table 5.52. Crystallography Statistics. Values for the highest resolution shell are given in parentheses. All data, except structures PaPBP3:8 and PaPBP3:16, were processed using STARANISOSO (Global Phasing) and the ellipsoidal completeness is given. PaPBP3:8 and PaPBP3:16 were processed with autoPROC (Global Phasing), and the value given is the spherical completeness.
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**Table 5.S3. PaPBP3 crystal pH.** Different pHs were used when growing and soaking each ligand into PaPBP3 crystals. The pH that gave the most complete density around the ligand was selected for analysis and deposition. Good quality data was collected at both pHs. Based on these data, pH does not appear to correlate with the binding mode observed.
5.5 Further Crystallography Views

Figure 5.51. Benzoxaboroles engage β-lactamases (AmpC) and PaPB3 differently. Selected residues from crystal structures of *P. aeruginosa* AmpC and PaPB3. The Ca carbons of residues of the SXXK, SXN and K(S/T)G motifs (residues 294/90, 297/93, 349/177 and 484/342 in PaPB3/PaAmpC respectively) of the PaPB3:13 complex (cyan) and the PaAmpC:benzoxaborole complex (PDB: 4WYY 4). In AmpC the benzoxaborole is mono-covalently reacted. The PaAmpC benzoxaborole is rotated by ~50° (through its long axis) relative to its position in the PaPB3 crystal structure which is associated with attack of the benzoxaborole on the other face. The C-3 acid group is positioned similarly in both complexes. Only one of the two alternate conformations of the side chain of PaPB3 Lys297 is shown for clarity.
Figure 5.S2. Comparison of the interactions of reacted piperacillin and 1 with PaPBP3. Crystal structures of PaPBP3 with (a) reacted piperacillin and (b) 1 are shown with serine-binding groups coloured in orange and groups engaging the acid binding pocket shown in pink. (c) Active site of PaPBP3:1 (orange and cyan), with the molecule of reacted piperacillin overaid (black, PDB code: 6R3X). A molecule of DMSO found in the active site of PaPBP3:1 is shown, as well as certain water molecules (w). The right hand water molecule occupies the oxyanion hole. (d) Comparison of the interactions of residues of PaPBP3 and residues of Actinomadura sp. R39 (AsR39) with boronate 1 and an R39 inhibitor from Zervosen et al. All three residues (294/49, 349/298 and 484/410 in PaPBP3/AsR39 respectively) have similar conformations as they engage the boron atom.
Figure 5.S3. Crystal structure views and Newman projections of Ser294 of PaPBP3 reacted with boron-containing inhibitors and piperacillin. (a) Enlarged view of Figure 5.10. Overlay of Ser294 of PaPBP3 reacted to boron-containing compounds (1, 13 and Vaborbactam) and piperacillin (PDB: 6R3X). Residues Ser349 and Lys484 are not shown in any of these figures. The position of boron within the active site correlates with changes in the Ser294 chi1 and chi2 angles. (b) Atoms and side chain angles of Ser294. (c-g) Structure views and Newman projections aligned along the Ser294 Ca-Cβ bond (c-e) and the Ser294 Cβ-Oy bond (f and g) for each of the boron-containing compounds compared to the piperacillin-reacted structure (black): vaborbactam (c and f), 13 (d and g), 1 (e). Chi1 dihedral angles are defined relative to the Ser294 Na, Chi2 dihedral angles are defined relative to the Ser294 Co. Mono-covalent structures (vaborbactam and β-lactams) and di-covalent structures (benzoxaboroles) are differentiated by their Ser294 chi2 angles but have similar (gauche) Ser294 chi1 angles. Tri-covalent structures (phenyl boronic acids 1 and 2 and previous observations 6) have trans serine chi1 angles.
Figure 5.S4. β-lactams piperacillin, ceftazidime, amoxicillin and aztreonam make a consistent hydrogen bond with Thr487. Crystal structures of PaPB3 bound to various β-lactams. The β-lactam derived nitrogen:Thr487 hydrogen bond is shown for PaPB3 reacted with piperacillin (black, PDB code: 6R3X)\(^6\), amoxicillin (white, PDB code: 6I1E)\(^5\), ceftazidime (green, PDB code: 3POB)\(^7\) and aztreonam (blue, PDB code: 3PBS)\(^8\). The hydrogen bond (dashed lines, coloured the same as the compound) is similarly positioned in each view. Chemical structures of ceftazidime, amoxicillin and aztreonam are shown.
5.4 Computational Chemistry

Computational docking of boronates was performed by Charles Eyermann (University of Cape Town). The protocol is reproduced here, with permission.

The benzoaborole di-covalent binding mode of compound 3 in complex with PaPBP3 was used as a template for modelling benzoaborole design ideas. Given the β5-α11 loop was not defined in the PaPBP3: 3 structure (PDB: 7ATX), presumably due to its inability to engage the residues on the β5-α11 loop, a model of 3 bound to the PaPBP3 structure as observed in the piperacillin-reacted PaPBP3 structure (PDB: 6R3X) was developed. All modelling studies were performed using the Schrodinger Suite of programs (Schrodinger LLC, New York). Benzoaborole design ideas which incorporate the key binding interactions between reacted piperacillin and PaPB3 were built by modifying compound 3 using Schrodinger’s 3D Builder tool. All designs, including compound 11, were docked into PaPBP3 using the Glide SP software. Glide SP docking calculations were performed on a noncovalently bound ligand where the oxaborole portion of the benzoaborole was deleted and the remaining phenyl group was constrained to the position observed in PaPBP3: 3. Hydrogen-bond constraints to Tyr328, Asn351 and Arg489 were applied in the Glide SP docking. The Glide docking poses of the inhibitor were then reconstituted to a complete covalently bound benzoaborole and the inhibitor and PaPBP3 residues within 6 Å of the inhibitor were then minimized using Prime. The only explicit water included in the minimization calculations was a crystallographically observed (in PaPBP3: 3) water that hydrogen bonds to the backbone NHs of Ser294 and Thr487 (w in Figure 5.4). The minimizations were performed using the OPLS2005 force field, the variable-dielectric generalized Born solvation model for water, a dielectric constant of 80, and 40 iterations of 200 steps each.
5.S5 LC-MS traces

Compound

![LC-MS trace](image)

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Do not use Multiplier & Dilution Factor with ISTDs.

**SI g/mL:** 1.845 A, SI g=454.4, Ref=450, 1.05

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### Area Percent Report

**Signal 1: DNQ, A, S=599, Ref=595, 10**

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Total area: 100.000

### Area Percent Report

**Signal 1: DNQ, A, S=599, Ref=595, 10**

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Total area: 100.000

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5.6 References


(2) Tickle, I. J.; Flensburg, C.; Keller, P.; Paciorek, W.; Sharff, A.; Vonrhein, C.; Bricogne, G. STARANISO; Global Phasing Ltd.: Cambridge, United Kingdom.


Chapter 6. Double Reaction of Nitrocefin with PBP3

Chapter 6.5 (p268-274) contains additional information referred to in this chapter

6.1 Introduction

Nitrocefin (Figure 6.1) was initially studied as an antibiotic compound, but whilst it was found to have antimicrobial effects against some species, (e.g. 0.05 μg/ml against non-β-lactamase producing S. aureus) 1,2, its value was recognised as an agent for detecting β-lactamase producing organisms 2; in the presence of β-lactamases, the colour changes from yellow to dark red. Its chromogenic properties have been used to study the kinetics of purified β-lactamases 3-9 and a few PBP 10-15, but to my knowledge it has only been used in gram-negative class B PBP to obtain simple IC₅₀ 16,17. Whilst undertaking structural characterisation of the reaction of nitrocefin with PaPBP3 mutants, a crystal structure in which two nitrocefin molecules were bound was identified; this work covers attempts to understand this complex.

Figure 6.1. Chemical structure of nitrocefin. Atoms of the cephalosporin core and side chains are labelled. Nitrocefin molecular mass is 516.5 Da. On the right hand side, the R = the enzyme (linked via serine) when the nitrocefin has formed an adduct or R = H if the molecule is deacylated and in solution.
6.1.1 Simple Saturation and Substrate Inhibition Kinetics

Scheme 6.1. Schemes of the simple reaction pathway (a) and the substrate inhibition pathway (b). Where: E: enzyme; S: substrate; ES: enzyme:substrate complex with one substrate bound; P: product; ESS: enzyme:substrate complex with two substrates bound. Rate constants ($k_+^e, k_-^e, k_+^a, k_+^b$, and $k_2$) are shown for their respective steps. (b) This is a simple substrate inhibition scheme in which the second substrate can only bind to the ES complex and the ESS complex does not yield a product directly (see section 6.4.4).

The kinetic constants of many bimolecular enzymatic reactions can be found by measuring the steady state initial rate of reaction at varying substrate concentrations to find a “substrate concentration graph”. The substrate concentration graphs of many enzymes can be fitted to a “Briggs-Haldane” kinetic model, where the initial reaction velocity ($v$) increases hyperbolically with substrate concentration ($[S]$), scaled by two constants: the maximum reaction velocity ($V_{\text{max}}$) and the constant $K_m$ (equation 6.1, Scheme 6.1a). This is termed simple saturation kinetics as the dependence of the rate of reaction on concentration of substrate follows a single phase, saturating at $V_{\text{max}}$. This model assumes: steady state kinetics (where the concentration of intermediate ES is constant throughout the reaction); that $[S]$ is in great excess of the enzyme concentration; and that the rate limiting step is $k_{\text{cat}}$ (Scheme 6.1, equation 6.1a) $^{18-22}$:

$$
  v = \frac{V_{\text{max}} [S]}{K_m + [S]}
$$

(6.1)

In some enzymes $^{23,24}$, a non-hyperbolic relationship is observed with increasing substrate concentration (Figure 6.2), where there is a maximum rate attained at a given substrate concentration, above which, it declines. In cases where such substrate inhibition is observed, an extra term $(1 + [S]/K_s)$ can be included to account for the decrease (equation 6.2) $^{21}$:
\[ V = \frac{V_{\text{max}} [S]}{K_m + [S] \left(1 + \frac{[S]}{K_{SI}}\right)} \] (6.2)

The new term is a consequence of the ability of the \( ES \) complex to combine with substrate to form \( ESS \) (Scheme 6.1b). \( K_{SI} \) is the dissociation constant for complex \( ESS \) (equation 6.6). Rearrangement of equation 6.2 makes it plain that the initial velocity at a given concentration of substrate is a weighted reduction of \( V_{\text{max}} \) (scaled by the values of \( K_m \) and \( K_{SI} \))

\[ V = \frac{V_{\text{max}}}{1 + \frac{K_m}{[S]} + \frac{[S]}{K_{SI}}} \] (6.3)

Equation 6.1 can be considered a case of equation 6.2 where \( K_{SI} \) is infinite, meaning \([S]/K_{SI}\) approaches zero and the term is ignored. Simulations with various arbitrary values of \( K_m \), \( K_{SI} \) and \( V_{\text{max}} \) are shown in Figure 6.2. Derivation of equation 6.3 from Scheme 6.1b is shown in Chapter 6.S. The Briggs-Haldane and substrate inhibition models (Scheme 6.1) were used to fit the kinetic curves of the reaction various class B PBPs with nitrocefin.

---

**Figure 6.2. Simulations of substrate inhibition kinetics by plotting equation 6.3 with different values of \( K_{SI} \) and \( V_{\text{max}} \).**

- Green: Simple saturation kinetics, \( K_{SI} = \infty \), \( V_{\text{max}} = 1 \); Weak substrate inhibition, \( K_{SI} = 100 \), \( V_{\text{max}} = 1 \); Blue: strong substrate inhibition, \( K_{SI} = 10 \), \( V_{\text{max}} = 1 \); Purple: Substrate inhibition with lower \( V_{\text{max}} \): \( K_{SI} = 100 \), \( V_{\text{max}} = 0.33 \). In all examples \( K_m = 5 \). Arbitrary units of concentration for substrate and arbitrary reciprocal time units for velocity. The values of \( K_m = 5 \) and \( V_{\text{max}} = 1 \) are shown with dotted lines. For substrate inhibition, the maximal velocity observed occurs at the geometric mean of \( K_m \) and \( K_{SI} \) (\( \sqrt[3]{K_m K_{SI}} \)); \( V_{\text{max}} \) can no longer be reached experimentally (red and blue lines) because one of either \( K_m[S] \) or \( [S]/K_{SI} \) will always be large in the denominator.

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6.2 Methods

6.2.1 Protein Expression and Purification

PBP3 from A. baumanii (AbPBP3) and E. coli (EcPBP3) were cloned by D. Bellini and purified by J. Todd. All the proteins have both the C- and N-terminal domains, but were solubilised by the removal of the single transmembrane helix (the first ~50 residues). Plasmids containing the genes for each of the proteins: wild type P. aeruginosa PBP (PaPBP3<sub>W</sub>), PaPBP3<sub>S294A</sub>, PaPBP3<sub>S349A</sub> and PaPBP3<sub>y409A</sub> were a kind gift from D. Bellini. All plasmids were pET47b vector-based with kanamycin resistance and N-terminal 6xHis-tag.

Chemically competent E. coli BL21-DE3 cells (50 µL) were inoculated with 2 µL of plasmid and incubated on ice for 30 minutes. Heat shock (42 °C for 45 seconds) was then performed before further incubation on ice (5 minutes). 400 µL of SOC media (2% (w/v) tryptone, 0.5% (w/v) yeast extract, 10 mM NaCl, 2.5 mM KCl, 10 mM MgCl₂, 10 mM MgSO₄, 20 mM glucose) was added and the bacteria were allowed to grow (37 °C, 1 hour, 180 rpm shaking). This inoculum (100 µL) was then spread onto agar plates containing kanamycin (50 µg/mL) and grown overnight (37 °C). A single colony from the plate was picked and used to inoculate lysogeny broth (LB) containing kanamycin at 50 µg/ml, which was then allowed to grow as a starter culture overnight (37 °C, 180 rpm shaking). The following day, 3 x 1 L of LB (containing kanamycin at 50 µg/mL) were inoculated each with 10 mL of the overnight culture and allowed to grow to an optical density (A₆₆₀ nm) of 0.6 in baffled flasks (37 °C, 180 rpm shaking). The cultures were then cooled to 20 °C and induced with 1 mM of Isopropyl β-D-1-thiogalactopyranoside (IPTG) and allowed to express protein overnight (20 °C, 180 rpm shaking).

The next day, cells were pelleted at 9,000 x g in a Beckman JLA 8.1000 rotor for 20 minutes at 4 °C. The cell pellet was resuspended in lysis buffer (20 mM Sodium Phosphate pH 8, 500 mM NaCl, 20 mM imidazole (binding buffer), supplemented with 5 mg/ml lysozyme and 5 mg/ml bovine DNase) and then lysed by ultrasonication (10 x (30 s bursts interspersed by a 30 s pause on ice), using a Bandelin Electronic sonicator. Centrifugation (30,000 x g for 30 minutes at 4 °C in a Beckman JA 25.50 rotor) was used to remove cell debris. All subsequent chromatographic steps were performed at room temperature, flow rate 3 ml/min:
the supernatant was loaded onto a 5 ml HisTrap FF Crude column (Cytiva). Subsequently the column was washed with binding buffer (25 mL) supplemented with 10% (w/v) glycerol. An ÄKTA FPLC system (Cytiva) was equilibrated in binding buffer + 10% (w/v) glycerol and then eluted with a gradient of 10% to 100% of elution buffer (20 mM Sodium Phosphate pH 8, 500 mM NaCl, 500 mM imidazole, 10% (w/v) glycerol) across the column, which eluted the protein (which have an N-terminal his-tag). Dialysis was used to exchange the protein into a storage buffer (20 mM Sodium Phosphate pH 8, 500 mM NaCl, 20% (w/v) glycerol) then the protein was aliquoted and frozen at -20 °C. Protein concentration was determined by absorbance at 280 nm on a NanoDrop (Thermo Scientific) using a calculated extinction coefficient from ProtParam \(^2\). Purity (> 95%) was confirmed by gel electrophoresis and mass spectrometry.

### 6.2.2 Kinetics Assays

Measurements were carried out at 30 °C in triplicate using a Hellma micro quartz cuvette (1 cm pathlength) in a Cary 100 UV/vis spectrophotometer (Agilent). Data was acquired every 0.1 s. Nitrocefin (Abcam) or CENTA ((6R,7R)-3-(((3-carboxy-4-nitrophenyl)thio)methyl)-8-oxo-7-(2-(thiophen-2-yl)acetamido)-5-thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, Calbiochem) in a buffer of 50 mM bis-Tris propane (Merck) pH 8.5, 75 mM NaCl, 1% (v/v) DMSO (Merck) was added to the cuvette before the protein (in the same buffer) was added to initiate the reaction. The final volume in the cuvette was 200 µL. The initial rate immediately after mixing was found using the spectrophotometer’s software. A dead time of ~ 10 s occurred whilst the substrates were mixed and the measurement initiated. The background turnover due to the spontaneous hydrolysis of nitrocefin or CENTA was measured alongside the experiment and subtracted from the rate. Reactions with nitrocefin were followed at 482 nm (ε\(_{1\,cm,\,482\,nm} = 17,400 \, M^{-1} \cdot cm^{-1}\))\(^5\) and reactions with CENTA were followed at 405 nm (ε\(_{1\,cm,\,405\,nm} = 6,400 \, M^{-1} \cdot cm^{-1}\))\(^3\). The specific rates (reaction velocity divided by the concentration of enzyme (equation 6.33)) were fitted to either Equation 6.3 or Equation 6.1 using non-linear regression in Prism 9 (GraphPad Software), and the best fit parameters are reported alongside the standard errors of the mean found by the software.

The protein concentration for each reaction used was chosen such that the turnover rate was at least 10-fold higher than the background nitrocefin turnover rate. For reactions with nitrocefin, the protein concentrations were: PaPBP3\(_\text{wt}\): 0.4 µM;
PaPBPr₃₃₄₉A: 2.4 μM; PaPBPr₃₄₅₉₈A: 2.4 μM; AbPBPr₃: 0.8 μM; EcPBPr₃: 1.8 μM. For the reaction with CENTA, 0.8 μM of PaPBPr₃₃₄₉ was used. For PaPBPr₃₃₂₈₄₄₉, conditions for steady state turnover at a sufficient rate were not identified (up to 10 μM was tested).

The upper limits for concentrations of substrates were determined by the solubility of the substrate. Partial solubility was observed >500 μM of nitrocefin and >1 mM of CENTA. ~2 fold less than this was used to ensure insolubility was not causing the observed effects. The lowest concentration of nitrocefin tested was always >10 fold higher than the protein concentration.

### 6.2.3 BOCILLIN FL Stained Gels

7.5 μg of each protein was reacted with 1 μM of BOCILLIN FL for 20 minutes in a buffer of 50 mM bis-Tris propane (Merck) pH 8.5, 75 mM NaCl, 1 % (v/v) DMSO, before the reaction was terminated by the addition of SDS loading dye (16.7 mM Tris pH 8.8, 0.67 % (w/v) SDS, 6.7 % (v/v) glycerol and 0.005% (w/v) bromophenol blue, 10 mM DTT). The samples (0.75 μg) were loaded into Mini-PROTEAN TGX Precast 4-20 % gels (Bio-Rad) along with a Color Prestained Protein Standard ladder (New England Biolabs). Electrophoresis was carried out at 140 V for 50 minutes in 25 mM Tris pH 8.3, 190 mM glycine and 0.1 % (w/v) SDS. The gels were rinsed in water then imaged by a Typhoon FLA 9500 (Cytiva) imager with a 473 nm excitation laser and a 510 nm emission filter (LPB (510LP)). The gels were then stained with coomassie dye (Instant Blue, Exedeon) and imaged by white light transillumination in a GeneSnap G:Box Gel Doc (Syngene).

### 6.2.4 Denaturing Mass Spectrometry

Denaturing mass spectrometry was carried out on a RapidFire RF365 high throughput sampling robot connected to a 6550 QTOF, by collaboration with the University of Oxford (C. Schofield group): with thanks to Dr. A. Krajnc and Dr. A. Tumber. 1 μM of protein was reacted with 10 μM of nitrocefin in 50mM sodium phosphate, pH 7.5 buffer and the mass spectra recorded in positive mode. Full methods will be published in a forthcoming publication (*in preparation*).

Methods investigated for tryptic digest mass spectrometry are given in section 6.5.3.
Methods for the crystallography (Carried out by D. Bellini) are given in section 6.3.4.

6.3 Results

6.3.1 Crystal Structures

During analysis of crystallographic data of β-lactams reacted with *P. aeruginosa* PBP3 (PaPBP3), the double binding of nitrocefin to a Y409A substitution mutant of PaPBP3 (PaPBP3\textsubscript{Y409A}) was unexpectedly observed (Figure 6.3). In contrast, only a single nitrocefin is observed in the crystal structure with the wild type protein (PaPBP3\textsubscript{wt}) (Figure 6.4). This is a novel configuration for a PBP:β-lactam complex. The Y409A mutant was originally of interest as Tyr409 is one of the only active site residues (outside of the flexible loop regions, see Chapter 4) with significant conformational flexibility (Figure 6.3).
Figure 6.3. Crystal structure of nitrocefin-reacted PaPBP_{Y409A}. Nitrocefin reacts separately with Ser294 and Ser349 in the PaPBP_{Y409A} structure. (a) Neighbouring residues are shown in sticks and are labelled. Hydrogen bonds (black dashed lines) are shown from the nitrocefin ligands to nearby residues. Two waters (W_{Y1A} and W_{Y2}) are found in the active site. Composite omit maps (contoured at 1 σ) are shown around the nitrocefin ligands. A comparison of the active sites of both structures is shown in Figure 6.4. (b) LigPlot 2D representation of the active site, showing polar (green dashed lines) and non-polar interactions (red arcs) of the two reacted nitrocefin molecules (orange). The shortest distance between the two molecules (3.75 Å) and the distance from W_{Y1A} and W_{Y2} to the β-lactam-derived carbonyl carbons are shown with purple dashed lines.

The PaPBP_{3Y409A} crystal structure does not have complete coverage of the ligand; in particular the flexible nitrophenyl-containing side chain, R2 (Figure 6.1), has only very weak density in both observations within the PaPBP_{Y409A} structure (Figure 6.3), as well as in the single nitrocefin-reacted PaPBP3_{wt} structure (6.S2). This is consistent with another report of nitrocefin bound to a PBP from the actinobacterium
*Actinomadura* in the PDB (PDB: 1W8Y\textsuperscript{28}), although the nitrophenyl was fully resolved in two other PBP-nitrocefin complexes (PDB: 2UWX\textsuperscript{14} and 1MWS\textsuperscript{29}). Double binding of nitrocefin was not observed in any of these structures.

The “second” (novel) molecule of nitrocefin reacts with the side chain γ-OH of Ser349, to form an analogous acyl-enzyme complex to the β-lactam adduct of the canonical Ser294 side chain. The second nitrocefin extends into the active site cleft to the right of the first (Figure 6.3) and significantly disrupts the conformation of the α10-β3 loop (Figure 6.4), preventing it from being resolved in the crystal structure. The second nitrocefin makes hydrogen bonds with Ser485 and Lys484 via its C-4 acid group, but no other polar interactions to active site residues. The two reacted nitrocefin molecules make no direct interactions to one another and at their closest they are 3.75 Å apart (Figure 6.3b).

In the two nitrocefin-reacted structures (PaPBP3\textsubscript{mr} and PaPBP3\textsubscript{Y409A}) Ser349 is observed in three different conformations, with two alternate conformations seen in the PaPBP3\textsubscript{mr}:nitrocefin complex (Figure 6.4c). Ser349 faces out of the active site (in a *gauche*\textsuperscript{*} conformation) when reacted with nitrocefin, in contrast to its inward conformation (trans) when reacting with boronate compounds (Chapter 5) (Figure 6.4d). Figure 6.4c-g shows a comparison of the position of Ser349 in both of the nitrocefin-reacted structures as well as in the PaPBP3:13 and PaPBP3:piperacillin complexes.
Figure 6.4. Electron density of nitrocefin-reacted PaPBP3\textsubscript{wt} (a) and comparison of the active sites of crystal structures of PaPBP3\textsubscript{wt} and PaPBP3\textsubscript{Y409A} reacted with nitrocefin (b), including the positions of serine residues (c-f). The PaPBP3\textsubscript{wt} protein structure is coloured in cyan and the reacted nitrocefin ligands coloured orange. The PaPBP3\textsubscript{Y409A} protein structure (b) is coloured in pink and the reacted nitrocefin ligands coloured purple. Nitrocefin reacts with Ser294 in the PaPBP3\textsubscript{wt} structure, and with Ser294 and Ser349 (separately) in the PaPBP3\textsubscript{Y409A} structure. Neighbouring residues are shown in sticks and are labelled. Hydrogen bonds (black dashed lines) are shown from the nitrocefin ligands to nearby residues. Residues Ser294, Ser349 and Tyr409/Ala409 are shown in darker blue in the PaPBP3\textsubscript{wt} structure and purple in the PaPBP3\textsubscript{Y409A} structure. In the PaPBP3\textsubscript{wt} structure, only one of the two alternate conformations of Ser349 is shown. A water (\(w_{\text{wt}}\) and \(w_{\text{Y409A}}\)) for the structures with PaPBP3\textsubscript{wt} and PaPBP3\textsubscript{Y409A} respectively) is shown positioned similarly in the active site of both structures. (c) The position of Ser294 is similar in the PaPBP3\textsubscript{wt} (cyan) and PaPBP3\textsubscript{Y409A} (pink) nitrocefin-reacted structures, but Ser349 can rotate. In the PaPBP3\textsubscript{wt} structure, two conformations of Ser349 were observed with occupancies of: 19\% (pale blue) and 81\% (dark blue). A water (\(w_{\text{wt}}\)) sits in a similar position. (d) Comparison of the position of both serine residues with other PaPBP3 structures: PaPBP3:13 (yellow, see Chapter 5) and PaPBP3:piperacillin (black, PDB: 6R3X\textsuperscript{26}). There is a difference of 1.3 Å between the position of the Ca of Ser349 of the PaPBP3:13 complex and that of the PaPBP3\textsubscript{wt}:nitrocefin structure. (e) Serine atom and dihedral bond angle labelling. (f) The \(\text{chi}1\) bond angles of Ser349 of the PaPBP3\textsubscript{wt} and PaPBP3\textsubscript{Y409A} nitrocefin reacted structures, compared to the piperacillin reacted structure. Ser349 is unreacted in PaPBP3\textsubscript{wt} structures, but reacted with nitrocefin in the PaPBP3\textsubscript{Y409A} structure. Dihedral bond angles are measured relative to the serine N\textalpha\ atom.

6.3.2 Substrate Inhibition Kinetics

6.3.2.2 Nitrocefin Turnover by gram-negative class B PBPs

![Figure 6.5. Change in absorbance at 482 nm due to nitrocefin turnover.](image)

Individual readings each of the 3 repeats are plotted as dots. The PaPBP3\textsubscript{wt}-catalysed turnover rate (dark colours) is compared to the background turnover rate (light colours): nitrocefin at 48 µM (red and pink) and nitrocefin at 160 µM (blue and cyan).
Nitrocefin turnover can be observed by measuring the change in absorbance at 482 nm ($A_{482\text{nm}}^{nm}$) (Figure 6.5). After acylation by a PBP, the red cephalosporic acid produced is strongly shifted from the yellow nitrocefin ($\lambda_{\text{max}} = 390 \text{ nm}$) (Figure 1.5). Measuring the rate of nitrocefin turnover at several concentrations (8-200 $\mu$M of nitrocefin) revealed that several of the enzymes investigated (PaPBP3$_{\text{wt}}$, AbPBP3, EcPBP3) appear to have substrate inhibition kinetics (Figure 6.6 and Table 6.1). The PaPBP3$_{\text{K409A}}$ protein had a highly attenuated turnover rate (~40-fold reduction in $k_{\text{cat}}$, compared to PaPBP3$_{\text{wt}}$) and appeared to show simple saturation kinetics.

To probe the role of Ser349 in the putative substrate inhibition mechanism, a S349A mutant (PaPBP3$_{\text{S349A}}$) was produced. The PaPBP3$_{\text{S349A}}$ has very different kinetics to the wild type protein: the $K_m$ was ~ 8-fold higher and the $k_{\text{cat}}$ > 10-fold lower. A mutant of the active site serine: Ser294 (PaPBP3$_{\text{Ser294}}$) showed insignificant turnover compared to the background turnover rate under the conditions tested.

To test the generality of the substrate inhibition, the nitrocefin turnover kinetics of class B PBPs from two other gram-negative species were investigated: A. baumannii PBP3 (AbPBP3) and E. coli PBP3 (EcPBP3). AbPBP3 and EcPBP3 have slower turnover (8-fold and 20-fold lower respectively) than PaPBP3, but appear to have substrate inhibition kinetics. For EcPBP3, the $K_m$ was too low to be determined (<15 fold above the protein concentration) and the shallow negative gradient of the right hand portion of the graph (Figure 6.6c; $-1.4 \times 10^{-3} \text{ min}^{-1} \cdot \mu\text{M}^{-1}$ compared to $-44.6 \times 10^{-3} \text{ min}^{-1} \cdot \mu\text{M}^{-1}$ for PaPBP3$_{\text{wt}}$) also makes it impossible to define the $K_{SI}$ of the interaction accurately.
Figure 6.6. The kinetics of nitrocefin and CENTA turnover by PBPs. (a-d) Substrate concentration nitrocefin turnover graphs in the presence of: (a) PaPBP3<sub>wt</sub> (protein concentration: 0.4 μM); (b) PaPBP3<sub>S349A</sub> (red, squares) and PaPBP3<sub>V409A</sub> (orange, circles) (both: 2.4 μM); (c) AbPBP3 (green, triangles, 0.8 μM), EcPBP3 (blue, diamonds, 1.8 μM). (d) CENTA turnover rates with PaPBP3<sub>wt</sub> (PBP concentration: 0.8 μM). Data points were fitted using non-linear regression (solid line) in Prism 9 for Mac OS (Graphpad) to equation 6.2 (a and c) or equation 6.1 (b and d). The best fit parameters are shown in Table 6.1. For all graphs, standard deviation of 3 points is shown for each point.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Protein</th>
<th>Model&lt;sup&gt;a&lt;/sup&gt;</th>
<th>(k_{cat} \text{ (min}^{-1})^{b})</th>
<th>(K_M \text{ (μM)}^{b})</th>
<th>(K_S \text{ (μM)}^{b})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrocefin</td>
<td>PaPBP3&lt;sub&gt;wt&lt;/sub&gt;</td>
<td>SI</td>
<td>39.0 ± 2.5</td>
<td>12.1 ± 1.7</td>
<td>259.1 ± 49.1</td>
</tr>
<tr>
<td></td>
<td>PaPBP3&lt;sub&gt;S329A&lt;/sub&gt;</td>
<td>ND</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Nitrocefin</td>
<td>PaPBP3&lt;sub&gt;S349A&lt;/sub&gt;</td>
<td>SS</td>
<td>3.6 ± 0.2</td>
<td>81.3 ± 10.2</td>
<td>-</td>
</tr>
<tr>
<td>Nitrocefin</td>
<td>PaPBP3&lt;sub&gt;V409A&lt;/sub&gt;</td>
<td>SS</td>
<td>1.2 ± 0.03</td>
<td>&lt;24&lt;sup&gt;d&lt;/sup&gt;</td>
<td>-</td>
</tr>
<tr>
<td>Nitrocefin</td>
<td>AbPBP3</td>
<td>SI</td>
<td>5.0 ± 0.2</td>
<td>3.8 ± 0.7</td>
<td>552.9 ± 107.5</td>
</tr>
<tr>
<td>Nitrocefin</td>
<td>EcPBP3</td>
<td>SI</td>
<td>1.7 ± 0.2</td>
<td>&lt;24&lt;sup&gt;d&lt;/sup&gt;</td>
<td>&gt;200&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>CENTA</td>
<td>PaPBP3&lt;sub&gt;wt&lt;/sub&gt;</td>
<td>SS</td>
<td>7.3 ± 0.2</td>
<td>20.0 ± 2.9</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 6.1. Kinetics of nitrocefin and CENTA turnover. <sup>a</sup>Models- SI: substrate inhibition, fitted with equation 6.2; ND: constants could not be determined under the conditions tested; SS: simple saturation, fitted with equation 6.1. Substrate concentration curves for SI and SS models are shown in Figure 6.6. <sup>b</sup>Errors are standard errors of the mean calculated by Prism 9. <sup>c</sup>Conditions for nitrocefin turnover were not identified. <sup>d</sup>\(K_M\) is lower than the lowest concentration of nitrocefin tested. <sup>e</sup>\(K_S\) is not determinable with confidence from the data collected.
6.3.2.3 CENTA turnover by PaPBP3

To determine whether substrate inhibition is observed with other β-lactams, turnover of another chromogenic cephalosporin, CENTA\(^{1,3}\) (12 - 400 nM) was investigated. Upon reaction, CENTA loses a leaving group, which leads to the change in absorbance\(^3\) (Figure 1.5). Unlike nitrocefin, this appears to have simple saturation kinetics when reacting with PaPBP3 (Figure 6.6d and Table 6.1).

6.3.3 Reactions with BOCILLIN FL

The ability of the PBPs to bind BOCILLIN FL was investigated with gel electrophoresis (Table 1.2). All proteins except PaPBP3\(_{G294A}\) and PaPBP3\(_{S349A}\) were able to react with BOCILLIN FL under the conditions tested (Figure 6.7).

![Figure 6.7. BOCILLIN FL (a) and coomassie (b) staining of an SDS-PAGE gel. (a) BOCILLIN FL (1 μM) was reacted with each PBP (7.5 μg) for 20 minutes before the reaction was terminated by the addition of SDS. Fluorescence was measured using a Typhoon FLA 9000. The ladder has a fluorescent marker protein at 72 kDa. (b) Coomassie staining was carried out on the same gel after the fluorescence was measured. Gel representative of 2 replicates.](image-url)
6.3.4 Denaturing Mass Spectrometry

Figure 6.8. Denaturing mass spectrometry of the reaction of PaPBP3<sub>wt</sub> (a-d) and PaPBP3<sub>V409A</sub> (e-h) with nitrocefin. Deconvoluted mass spectra show masses corresponding to the apo protein ion for (a) PaPBP3<sub>wt</sub>, predicted mass.
(from sequence): 59809.29 Da, observed 59809.34 Da and (e) PaPBP3_{Y409A}, predicted mass: 59717.19 Da, 5916.73 Da. A red line indicates the position of the apo protein with additional peaks corresponding to adducts of nitrocefin (expected incremental Δmass: 516.50 Da). Significant peaks are labelled and the differences between peaks are given in daltons. Reaction time courses were measured by sampling from the same reaction at multiple time points. The instrument dead time was ~10 s, preventing measurement of the reaction before this time. For both proteins, a PBP-nitrocefin complex was formed within 10 s (b and f). For PaPBP3_{wt} (a-d), sufficient nitrocefin (initial concentration: 10 μM) was turned over by 6.5 minutes to observe a mass corresponding to apo protein (c), which was the major ion after 11 minutes (d). For PaPBP3_{Y409A}, the reaction velocity is presumably too slow (Table 6.1) to fully turnover the nitrocefin so the apo protein mass was not observed after 11 minutes. A mass corresponding to the reaction of two nitrocefin molecules (n = 2) with the PaPBP3_{Y409A} (observed: 60751.24, expected: 60749.73 Da) is large relative to the major ion after 1.5 minutes of reaction (g) but has reduced abundance after 11 minutes (h).

Mass spectra of the wild type and three active site mutants of PaPBP3 (Figure 6.8 and Figure 6.9) reacting with nitrocefin were recorded under denaturing conditions. PaPBP3_{wt} and PaPBP3_{Y409A} react rapidly (<10 s) and completely (i.e. no apo protein mass is observed in Figure 6.8b or d) with nitrocefin to produce an ion corresponding to a singly reacted nitrocefin molecule (i.e. (PBP3 + nS)^+, where S is nitrocefin and n = 1). For PaPBP3_{wt} it appears little nitrocefin remains after 11 minutes as the apo protein ion is once again observed. The lower rate of nitrocefin turnover recorded in absorbance assays for PaPBP3_{Y409A} is reflected in the lack of reformation of the apo ion after 11 minutes (Figure 6.8h) with this mutant.

The mass spectral analysis of each protein reacting with nitrocefin shows a mass corresponding to the adduct of two or more molecules of nitrocefin (i.e. (PBP3 + nS)^+, where n ≥ 1). The presence of additional adducts is particularly clear in the PaPBP3_{S294A} spectra (Figure 6.9a), where they appear up to (PBP3 + 3S)^+, with each subsequent adduct (increasing values of n) having progressively lower abundance. It is possible these adducts correspond to nonspecific interactions of protein and ligand^{31}, as masses corresponding to (PBP3 + 2S)^+ are not anticipated in the Ser294 and Ser349 mutants and they appear at roughly constant abundance (< 0.3 x 10^4) in all mass spectra (Figure 6.8 and Figure 6.9).
Figure 6.9. Denaturing mass spectrometry of the reaction of PaPBP35294A (a and b) and PaPBP35394A (c and d) with nitrocefin. Both proteins were reacted with nitrocefin and their mass spectra were recorded after ~10 seconds (a and c) and 5 minutes (b and d). Significant peaks are labelled and the differences between peaks are given in daltons. The red line indicates the position of the apo protein, confirmed by a spectra of the unreacted protein. Both proteins have an expected mass of 59793.29 Da, but the observed apo protein mass is ~59849 for both proteins, a difference of ~56, which in the absence of detectable amino acid substitutions may correspond to an adduct of an $^{56}$Fe ion possibly at the His-tag: further work is needed to confirm this. Additional masses corresponding to adducts of up to 3 nitrocefin molecules ($n = 3$) are observed in the spectra with both proteins, after both 10 s and 5 minutes.

The PaPBP34094A mass spectrum (Figure 6.8g) has a relatively large (PBP3 + 2S)$^+$ peak after 1.5 minutes, which is not seen after 11 minutes (Figure 6.8g). This may correspond to a specific interaction of nitrocefin with Ser349 in the active site. However, the presence of apparently nonspecific (PBP3 + nS)$^+$ ions makes it difficult to assign the specific (PBP3 + 2S)$^+$ with confidence.
6.3.5 Tryptic Digest Mass Spectrometry

Analysis of a PBP-nitrocefin complex by tryptic digest mass spectrometry could be used to confirm the binding site(s) of nitrocefin within the protein. The addition of the mass of the nitrocefin-derived adduct should be detectable by liquid chromatography coupled to mass spectrometry analysis (LC-MS/MS)\textsuperscript{32}. This method has previously been used to detect the β-lactam binding site of β-lactamases \textsuperscript{33,34}. Disappointingly, it was not possible to identify the site of nitrocefin reaction by tryptic mass spectrometric digest, despite several attempts (Table 6.2, section 6.3.5). Peptides covering almost all of the protein were found but none were shown to have the nitrocefin-derived adduct (Table 6.2). This result is consistent with a previous unsuccessful attempt (using a similar method) by researchers at Eli Lilly to identify the catalytic residues of MRSA PBP2a which interact with nitrocefin \textsuperscript{13}.

<table>
<thead>
<tr>
<th>Trial</th>
<th>Protein</th>
<th>Amount of PBP digested</th>
<th>Compound</th>
<th>Reaction Conditions</th>
<th>Unfolding Conditions</th>
<th>Trypsin Digest\textsuperscript{a}</th>
<th>Protocol</th>
<th>Coverage\textsuperscript{b}</th>
<th>Adduct</th>
</tr>
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<tbody>
<tr>
<td>A</td>
<td>PaPBP3\textsubscript{K80A}, PaPBP3\textsubscript{Ac}, AbPBP3</td>
<td>100 µg</td>
<td>Cefazidime and Nitrocefin, 100 µM</td>
<td>5 minutes, rt</td>
<td>8 M Urea</td>
<td>Overnight, 1:50</td>
<td>I</td>
<td>70%: Ser294 but no Ser349 coverage</td>
<td>No</td>
</tr>
<tr>
<td>B</td>
<td>PaPBP3\textsubscript{K80A}</td>
<td>20 µg</td>
<td>Nitrocefin, 100 µM</td>
<td>5 minutes, rt</td>
<td>8 M Urea</td>
<td>4 hours, 1:20</td>
<td>I</td>
<td>83%</td>
<td>No</td>
</tr>
<tr>
<td>C</td>
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<td>20 µg</td>
<td>Nitrocefin, 100 µM</td>
<td>2 minutes, rt</td>
<td>99 °C 5 minutes heat shock.</td>
<td>4 hours, 1:20</td>
<td>II</td>
<td>96%</td>
<td>No</td>
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<td>D</td>
<td>PaPBP3\textsubscript{K80A}</td>
<td>5 µg</td>
<td>Nitrocefin, 100 µM</td>
<td>3 minutes, rt</td>
<td>99 °C 5 minutes heat shock.</td>
<td>4 hours, 1:20</td>
<td>II</td>
<td>58%: Ser294 but no Ser349 coverage</td>
<td>No</td>
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<tr>
<td>E</td>
<td>PaPBP3\textsubscript{Ac}</td>
<td>60 µg</td>
<td>Nitrocefin, 1 mM</td>
<td>5 minutes, rt</td>
<td>SDS</td>
<td>4 hours, 1:20</td>
<td>III</td>
<td>97%</td>
<td>No</td>
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Table 6.2. Screens to identify tryptic digestion mass spectrometry conditions for stable adduction. \textsuperscript{a}Time exposed to trypsin and protein:trypsin ratio given. \textsuperscript{b}Coverage is the percentage of the peptide sequence observed by mass spectrometry. rt: room temperature; SDS: sodium dodecyl sulphate.
6.4 Discussion

6.4.1 Structural Analysis

The crystallographic observation of two molecules of a β-lactam acylating the active site of a PBP is, to my knowledge, without precedent amongst the many reported PBP:β-lactam crystal structures. The density around the “second” nitrocefin reacting with Ser349 is not complete in the PaPBP3Y409A structure, but there is sufficient density around the core of the compound to be confident of its presence. The R1 and R2 side chains (Figure 6.1) of the “second” nitrocefin that reacts with PaPBP3Y409A as well as the nitrocefin in the PaPBP3wt structure are poorly defined by the electron density, likely due to their flexibility (they make few direct hydrogen bonds, Figure 6.3). The position of the R2 side chain of the “second” nitrocefin reacted with PaPBP3Y409A is particularly ambiguous. It is modelled extending into a region normally occupied by the α10-β3 loop of the protein. Like the side chain, the α10-β3 loop lacks clear density and its conformation is very poorly resolved. It appears that in the presence of the second nitrocefin the conformation of the loop is disrupted, increasing its flexibility, and preventing its resolution. This disruption has analogy with the dynamic conformations of the α10-β3 loop observed when various novel boronate compounds bind to PaPBP3 ( Chapters 4 and 5). The α10-β3 loop of the PaPBP3wt:nitrocefin structure is fully resolved.

In neither the PaPBP3wt nor the PaPBP3Y409A nitrocefin-reacted structure is the β5-α11 loop resolved (residues 530-536 and 531-534 respectively are missing). This is consistent with the structures of PaPBP3:boronate complexes, many of which lack electron density for this loop (Chapter 5). The nitrophenyl group of the R2 sidechain of the Ser294-reacting nitrocefin is located in a similar position to the hydrophobic wall residues Tyr532 and Phe533 of the PaPBP3:piperacillin complex (PDB: 6R3X) , which may prevent the β5-α11 loop from entering the “closed” conformation (Chapter 4).

The crystallography data was collected after soaking crystals with 10 mM of nitrocefin for 60 minutes. This concentration is many fold higher than the concentrations used in the assays or mass spectrometry. Soaking (where a high concentration of the compound is added to a pre-grown crystal) was used to add nitrocefin to PaPBP3. This means that the reaction did not occur in solution but in
the crystal phase of the protein (which also has a much higher concentration than in solution), which may affect the kinetics. It is possible that the observation of nitrocefin bound to Ser349 is a crystallographic artefact, but the lack of the second nitrocefin in the wild type structure or at the various other solvent-exposed serines within the protein indicates specificity of the reaction.

6.4.2 Nitrocefin Kinetics

The PaPBP3<sub>wt</sub> nitrocefin turnover rates are much faster than values reported for PaPBP3<sub>wt</sub> turnover of other β-lactams (e.g. meropenem turnover (the rate limiting deacylation step) is just 0.02 min<sup>-1</sup>)<sup>36</sup>, but still much slower than nitrocefin turnover rates by β-lactamases (e.g. TEM-1 can turnover nitrocefin at rates of > 4 x 10<sup>4</sup> min<sup>-1</sup>; more than one thousand fold faster than PaPBP3<sup>)</sup><sup>37</sup>. Comparison between PaPBP3<sub>wt</sub>, AbPBP3 and EcPBP3, show significant differences in their behaviour, despite their structural and sequence similarities (Figure 1.3). Catalytic fine tuning as a result of small differences in the spatial geometry of the active site and/or non-conserved residues may cause the variance in the behaviour of these proteins. Although weak, there does appear to be substrate inhibition in AbPBP3 and possibly in EcPBP3, which suggests conservation of the substrate inhibition mechanism between species.

The differences in the <em>K</em><sub>M</sub> and <em>k</em><sub>cat</sub> of CENTA and nitrocefin for PaPBP3<sub>wt</sub> are consistent with other reports which show that substrate specificity for CENTA and nitrocefin varies between enzymes.<sup>3</sup> The apparent lack of substrate inhibition (under the conditions tested) in CENTA is interesting and suggests that the mechanism may have some specificity to nitrocefin. CENTA has a 3-position leaving group<sup>3</sup> but nitrocefin does not, which may be important. Unfortunately, studies (using similar techniques) with other β-lactams are experimentally challenging in PBP<sub>s</sub> due to their low turnover rates and their generally small extinction coefficients (typically < 1000 M<sup>-1</sup> cm<sup>-1</sup>)<sup>38</sup>, which would necessitate very high protein concentrations to get measurable signals.

6.4.3 β-lactam Kinetics Beyond Gram-Negative Class B PBPs

Other HMM PBP<sub>s</sub> have been investigated for their reactions with nitrocefin: S. <i>aureus</i> PBP2a<sup>39</sup>, <i>Streptococcus pneumoniae</i> PBP1b<sup>14</sup> and PBP2b<sup>12</sup> and E. <i>coli</i>
PBP1b \(^\text{10}\). For each of these proteins, the nitrocefin turnover rate was either not measured or < 0.03 min\(^{-1}\). For \(S.\ pneumoniae\) PBP2b, mutations in a clinical strain were shown weakly to increase the rate of deacylation (Chapter 4)\(^\text{12}\); Interestingly, Page describes EcPBP1b as having biphasic kinetics and greater-than-stoichiometric acylation of the protein \(^\text{10}\). After several enzyme turnovers the formation of a “new inhibited species” was observed which slows the turnover. The new species was attributed to a conformational change within the protein \(^\text{10}\), but the formation of a doubly acylated EcPBP1b is also consistent with these observations.

Examples across all classes of serine \(\beta\)-lactamase (class A, C, and D) have shown non-stoichiometric burst kinetics for the reaction of \(\beta\)-lactamase with many different \(\beta\)-lactams tested \(^\text{4,7,8,40–42}\). This behaviour is typically described as “substrate-induced inactivation” \(^\text{42}\) and not substrate inhibition: that is to say the transition to the less active form is a consequence of interaction with \(\beta\)-lactam substrates, but does not appear to be substrate concentration dependent. Waley suggested that the inactivation is the result of branching at the level of the \(ES\) complex to a less active form \(^\text{42}\), a mechanism that is similar to the model shown in Scheme 6.1b, but without the molecule of substrate required for the transition from \(ES\) to \(ESS\).

### 6.4.4 Kinetics Models

Considering first only the reaction of a single nitrocefin with PBP3, the kinetic scheme (Scheme 6.1a) used to fit the data to Briggs-Haldane steady state models is a simplification of the scheme presented in Scheme 1.2. It is expected nitrocefin will pass through both of the complexes described (\(E\cdot I\) and \(EI\)) during a turnover cycle, however the kinetic model reduces the two catalytic reactions (covalent bond formation and bond hydrolysis) to a single term: \(k_{\text{cat}}\), effectively ignoring the \(EI\) complex \(^\text{43}\). The magnitude of \(k_{\text{cat}}\) will be limited by the slowest of the two steps. Additionally, \(K_M\) also includes the \(k_{\text{cat}}\) term (equation 6.5), which means that mutants affecting \(k_{\text{cat}}\) also affect \(K_M\). Therefore within the context of a system obeying steady-state kinetics, it is not correct to say that the increase in \(K_M\) measured with PaPBP3\(^{3349A}\) is solely due to the change in affinity of the protein for nitrocefin: there is also a change in catalytic rate \(^\text{44}\).

In addition to the simplifications present in kinetic model for Brigg-Haldane steady state nitrocefin turnover, the kinetic model of substrate inhibition (Scheme 6.1b)
makes a number of assumptions. It assumes that: (I) the “second” molecule of nitrocefin can only bind to the ES complex; (II) ESS is unproductive (or nitrocefin turnover from ES is so much faster than turnover from ESS that the ESS complex appears to be unproductive); (III) ES is in rapid equilibrium with ESS (necessary for the validity of equation 6.2).

More complex kinetic models can be designed in which the ESS complex is productive (Scheme 6.S1a) or in which the “second” nitrocefin can bind to both E and ES (Scheme 6.S1b). Fitting of the rate equation to the former model (Scheme 6.S1c) to the data for PaPBP3 wt (Figure 6.6a) was attempted. An equivalent fit of the data was achieved, but several of the parameters were indeterminate, indicating that the model is too complex for the data. Therefore the simplest possible model for substrate inhibition (Scheme 6.1 and equation 6.3) was chosen for analysis of the data. This model may be sufficient given that $K_{si}$ is so much larger than $K_m$ (almost 7 fold for PaPBP3 wt), so the fraction of enzyme in the doubly-bound form (i.e. ESS/(E + ES+ ESS)) is small at the concentrations tested and the reactions of ESS make little contribution to the overall rate.

Unlike the kinetics experiments, both denaturing mass spectrometry and the BOCILLIN FL stained gel only measure the EI complex as the non-covalent E: I complex is lost due to the denaturing process. These techniques compared to the kinetics are making orthogonal observations of the same processes but both techniques here are only used qualitatively.

6.4.5 S294A and S349A Mutants

Alanine substitution of two serines in the active site: Ser294 (SXXK) and Ser349 (SXN) have differing effects. S294A leads to a complete loss of kinetic activity, consistent with the importance of this residue for catalysis. The mass spectrometry for PaPBP3 S294A (Figure 6.9a and b) shows little nitrocefin reaction beyond the rapid formation of apparently nonspecific nitrocefin adducts (see below). It therefore appears that this mutant is unable to acylate nitrocefin.

In contrast, the S349A substitution leads to an increase in $K_m$ and decrease in $k_{cat}$. The $k_{cat}/K_m$ for PaPBP3 S349A is 0.044 μM⁻¹ min⁻¹: 73 fold lower than the value for PaPBP3 wt (3.22 μM⁻¹ min⁻¹), indicating the mutation has a significant effect on the
catalytic activity of the protein. Substrate inhibition is not observed in PaPBP3<sub>S349A</sub>, which is consistent with S349A being the binding site for the second (inhibition-causing) molecule of nitrocefin. However, nitrocefin insolubility prevents measurement of turnover at higher nitrocefin concentrations, so it is not possible to confirm a lack of substrate inhibition: ideally values closer to \( k_{cat} \) should be observed.

The mass spectrometry of PaPBP3<sub>S349A</sub> shows (unlike PaPBP3<sub>wt</sub>) that the majority of the protein remained unligated after 10 s of reaction (Figure 6.9c), which may indicate this mutation is reducing the rate of acylation. Consistent with this, is the lack of (or slow) reaction with BOCILLIN FL, which could be due to a slow acylation rate.

The equivalent serine in Class A β-lactamases (Ser130) is hypothesised to assist in the β-lactam bond cleavage during acylation by hydrogen bonding to nitrogen and donating a proton \(^{45-47}\), so it may have an analogous role in HMM PBP. The effect of S349A mutation on the rate of catalysis is within values reported for the analogous substitution in other proteins: S130A substitution in a class A β-lactamase (\textit{Streptomyces albus} G β-lactamase) led to a 10 fold decrease in the \( k_{cat}/K_M \) for nitrocefin \(^{46}\); S110A substitution in \textit{E. coli} PBP5 (a LMM carboxypeptidase) decreased both acylation (83-fold vs wild type) and deacylation (8 fold vs wild type) of benzylpenicillin \(^{49}\).

### 6.4.6 The Tyr409Ala Mutant

Steady state kinetic analysis shows that the Y409A substitution mutation leads to an almost 40-fold decrease in the nitrocefin \( k_{cat} \) compared to PaPBP3<sub>wt</sub> (39.0 ± 2.5 min<sup>-1</sup> and 1.2 ± 0.03 min<sup>-1</sup> for PaPBP3<sub>wt</sub> and PaPBP3<sub>Y409A</sub> respectively), whilst the \( K_M \) appears to remain low (<24 μM for both proteins). Furthermore, mass spectrometry shows the mutation does not prevent the rapid (< 10 s) acylation of the protein (Figure 6.8f). The kinetics of PaPBP3<sub>Y409A</sub> were modelled with Briggs-Haldane kinetics (Equation 6.1), as opposed to substrate inhibition kinetics (Equation 6.3). However, this does not necessarily mean that substrate inhibition is not occurring, as the large reduction in the \( k_{cat} \) makes the absolute decrease in velocity at higher nitrocefin concentrations smaller (Figure 6.2), and therefore harder to measure with
certainty. Alternatively, the mutation might increase the value of $K_{\text{mi}}$ to such an extent as to render it practically impossible to demonstrate substrate inhibition.

The role of Tyr409 in the active site may be that it helps to position a water near to Ser294 for catalysis of the $\beta$-lactam-derived acyl bond. In the piperacillin-reacted PaPBP3 structure (PDB: 6R3X\textsuperscript{25}), a water is hydrogen bonded to Tyr409 as well as the $\beta$-lactam-derived carbonyl (Figure 6.10). In the PaPBP3$_{y409A}$:nitrocefin structure the water is still present, but it is displaced slightly (0.3 Å) away from the $\beta$-lactam-derived carbonyl with concurrent loss of hydrogen bonds to Asn351. This could result in the decreased catalysis rate observed (Table 6.1). Curiously, in the PaPBP3$_{\text{wt}}$:nitrocefin this water is not observed, perhaps due to the structure’s lower resolution (2.53 Å compared to 1.99 Å in the PaPBP3$_{\text{wt}}$ and PaPBP3$_{y409A}$ nitrocefin-reacted structures respectively).

Figure 6.10. Waters near to the active site serine. Crystal structures of PaPBP3$_{y409A}$:nitrocefin (pink) and PaPBP3$_{\text{wt}}$:piperacillin (black, PDB: 6R3X\textsuperscript{25}) complexes show a water (labelled $w_{\text{Y409A}}$ and $w_{\text{pip}}$ respectively in the two structures). Hydrogen bonds are formed from $w_{\text{pip}}$ to Ser294, Asn351, Tyr407 and Thr487, whereas $w_{\text{Y409A}}$ only hydrogen bonds to Ser294 and Tyr407. The water is displaced away from Ser294 in the PaPBP3$_{y409A}$:nitrocefin complex: the distance from $w_{\text{Y409A}}$ to the $\beta$-lactam derived carbonyl carbon is 3.1 Å but the equivalent distance in the PaPBP3$_{\text{wt}}$:piperacillin is 2.7 Å. The distance from the water to the $\delta$O of Asn351 increases from 3.0 Å in the PaPBP3$_{\text{wt}}$:piperacillin structures to 3.7 Å in the PaPBP3$_{y409A}$:nitrocefin structure, with the associated loss of a hydrogen bond. The water $w_{\text{Y409A}}$ is also shown to the right of Ser294. A hydrogen bond between Tyr409 and Thr487 is shown.
6.4.7 Mass Spectrometry

Assuming uniform efficiency of ionisation of species, denaturing mass spectrometry of nitrocefin-reacted proteins creates a snapshot of the solution state of covalent complexes at a given time point. After denaturing the protein, only covalent adducts should remain, with non-covalent interactions dissociated. The results presented here appear to show that for PaPBP3_{49}, PaPBP3_{S349A}, PaPBP3_{Y409A} a single nitrocefin-derived acyl-enzyme complex is formed as a major ion, whilst additional adducts of \( n \geq 1 \) are formed non-specifically. The adducts are designated to be non-specific from their presence in the catalytically inactive PaPBP3_{S329A}, their consistent low abundance and apparent high affinity (they are formed in <10 s). Investigation of comparable adducts in other proteins concluded that they form as the results of a random chemical process during the gas phase of the electrospray technique \(^{31,50}\). Work to improve the quality of the mass spectra is ongoing. The use of a reference protein to measure only the non-specific portion of the interaction \(^{50}\) or increasing concentration of salt ions to suppress non-specific interactions \(^{31}\) may be effective.

Alternatively, the non-specific adducts could be the result of the reaction of relatively reactive nitrocefin with solvent-exposed serines outside the active site. Adducts to other serines (i.e. not Ser294 or Ser349) are not observed crystallographically, but could occur in solution or be unobservable due to high mobility. Non-specific adducts such as these could provide an explanation for the observations of high affinity, irreversible interactions seen between nitrocefin and PaPBP3 in SPR experiments (Figure 2.11).

The mass spectrometry did not provide strong evidence of specific double binding of nitrocefin to PaPBP3. For PaPBP3_{Y409A}, a (PBP3 + 2S)\(^+\) ion was observed to form and decay (Figure 6.8e-h), although its abundance is too low to be sure of its specificity. Increasing the ligand concentration may increase the abundance of the doubly bound ion. Ideally concentrations above the kinetically determined \( K_S \) (> 260 \( \mu \text{M} \)) should be used for the mass spectrometry experiments but this is experimentally limited (to ~ 100 fold above protein concentration) by ion suppression which increases with ligand concentration \(^{50}\).
6.4.8 Hypotheses on the Observation of Two Nitrocefin Molecules

One possibility for the presence of the two molecules of nitrocefin in the PaPBP3<sub>Y409A</sub> structure proposes that the ‘second’ nitrocefin is able to react with Ser349 due to the slow nitrocefin catalysis of PaPBP3<sub>Y409A</sub>. The structures of di-covalently reacted benzoazaboroles (e.g. Figure 5.6, Chapter 5) may offer a potential mechanism for this. Mono-covalently reacting boronates have previously been described as transition state mimics of β-lactam acylation<sup>16,51</sup>; speculatively, di-covalently reacting boronates could be transition state mimics of nucleophilic attack on the Ser294 acyl-enzyme complex by Ser349, the result of which is the observed Ser349:nitrocefin complex (Figure 6.11). The di-covalent benzoazaborole crystal structures demonstrate that the two serine residues are sufficiently close and that the necessary reorientation of Ser294 is possible by rotation of <i>chi1</i> and <i>chi2</i> angles (Figure 5.10). This pathway may become dominant in PaPBP3<sub>Y409A</sub> as the nucleophilic water w<sub>VA2</sub> is displaced and becomes less reactive (Figure 6.10), leaving Ser349 as the most reactive nucleophile. An analogous mechanism in which the equivalent ‘second’ serine attacks an acylated catalytic site serine was proposed to explain a structure in which two serines are bonded to two different species following reaction of SHV-1 with tazobactam<sup>52</sup>.

Alternatively, the double reaction with nitrocefin may be an indirect consequence of the Y409A substitution. The substitution may have a similar effect to clinical mutations described in Chapter 4 (to my knowledge, mutations at position 409, or equivalent in other species, have not been identified in any clinical mutations). For example, this could be mediated via the loss of interactions of Tyr409 with backbone carbonyl of Thr487 on the β3 strand (Figure 6.10, 6.S1). Changes to the β3 strand could lead to a rearrangement of the α10-β3 loop, which opens up to allow nitrocefin to reach and react with Ser349.
Figure 6.11. Di-covalently binding benzoxaboroles may mimic a transition state in a nitrocefin serine transfer mechanism. (a) A Ser294 nitrocefin adduct is first formed by the classical β-lactam reaction (e.g. Figure 1.2). After reorientation of the serine side chain and its β-lactam-derived adduct, nucleophilic attack by Ser349 of the β-lactam-derived carbonyl carbon can occur. Either Lys297 or Lys484 could act as the general base for the activation of Ser349 (shown with green or blue dashed arrows respectively). The result of the nucleophilic substitution is the transfer of the nitrocefin adduct from Ser294 to Ser349. (b) The transition state in this reaction is analogous to the di-covalent benzoxaborole complex (e.g. Figure 5.6).

Presently there is insufficient evidence to confidently determine the mechanism that leads to the formation of the doubly acylated complex. Nucleophilic attack by Ser349 (necessary in both mechanisms) will likely require activation by a general base, most likely either neutral Lys297 or Lys484 (green and blue arrows respectively in Figure 6.11). The general base mechanism is analogous to the acylation mechanism proposed in other penicillin reactive enzymes where Lys297 (or equivalent) activates Ser294 (or equivalent) for nucleophilic attack on the β-lactam ring \( ^{45,46,53–56} \).

However, it cannot be discounted that the complex may solely have arisen as the result of a crystallographic artefact in the PaPBP3\(_{Y409A}^-\).
6.4.9 Future Work

Further work is required to fully characterise the doubly reacted PBP3 structure. Evidence for such a complex in solution remains ambiguous. Further kinetic experiments are required to establish the link between the crystallographic species and the substrate inhibition observed. The use of pre-steady state kinetics may provide insight into the rate constants for individual steps, as has been applied to study substrate-induced inactivation in β-lactamases\textsuperscript{42,57}. Kinetic studies at a range of pHs could be used to determine the pK\textsubscript{a} of residues involved in catalysis\textsuperscript{6,54,58,59}. In particular it would be of interest to see if the degree of substrate inhibition is affected by pH, as it may be possible to distinguish reaction at the two serines by their pH profiles. In general, the catalytic mechanisms of residues in the HMM PBP active site\textsuperscript{59} have been understudied compared to LMM PBPs or β-lactamases so further work may have a wider benefit.

Substitutions of Lys297 and Lys484 may be interesting to investigate. Whichever residue serves as a general base to Ser349, its mutation should slow or prevent nucleophilic attack by Ser349. To probe the role of Tyr409, further substitution of the position may be valuable. For example, substitution with hydrophilic phenylalanine or acidic glutamic acid may affect the behaviour of water w\textsubscript{Y409} and perhaps affect double acylation.

Optimisations of the conditions for conducting denaturing mass spectrometry experiments may lead to clearer evidence of the doubly reacted species in solution. Mass spectrometry could be used to investigate whether other β-lactams are able to doubly react with PBPs, since this is challenging spectrophotometrically for non-chromogenic compounds. Once conditions have been established for the denaturing mass spectrometry, they may be applied to improve the quality of the tryptic digest mass spectrometry data to allow confirmation of double binding by this method.

Finally the penamaldate assay, once used to establish the stoichiometric binding of β-lactams to β-lactamases\textsuperscript{44,60,61}, could be used to independently establish the reaction stoichiometry, although HgCl\textsubscript{2} is toxic, presenting an experimental challenge.
6.5 Conclusion

This work focuses on initial efforts to understand the observation of double reaction of an active site mutant of PaPBP3 with the chromogenic β-lactam nitrocefin. Steady state turnover of nitrocefin by PaPBP3mt and several active site mutants was carried out which showed substrate inhibition, a behaviour consistent with the binding of a second molecule of substrate. Substrate inhibition was observed in other class B PBPs from related species. Mass spectrometry of nitrocefin-reacted proteins shows turnover of the substrate, but does not provide firm evidence of doubly-reacted proteins. Tryptic digest mass spectrometry was unsuccessful and nitrocefin adducts could not be recorded. For now, caution must be taken when interpreting this interesting observation as a biologically relevant phenomenon. This effect may be limited to nitrocefin, which appears to have an unusually high turnover rate compared to other β-lactams 36. Nitrocefin (but not imipenem) has been reported to bind reversibly to the active site of a L,D-transpeptidase from Enterococcus faecium: a peptidoglycan cross-linking enzyme structurally unrelated to PBPs 62. Nitrocefin itself then may be a unique β-lactam, capable of reacting in unique ways. If this behaviour can be better understood (perhaps with further insight from boronate binding), it may be possible to design a novel generation of inhibitors which exploit reactions at both Ser294 and Ser349, similarly to the reactions of e.g. clavulanic acid and tazobactam with β-lactamases 52,63–65.

6.6 References

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Chapter 6.S. Supplemental
Information for Chapter 6

6.S1 Structural Views

Figure 6.S1. Tyr409 conformational flexibility in PaPBP3. (a) Comparison of the PaPBP3:aztreonam (green, PDB: 3PBS\(^1\)) and PaPBP3:piperacillin (black, PDB: 6R3X\(^2\)) crystal structures. Tyr409 is pointed towards the active site serine in the piperacillin-reacted structure, but faces outwards in the aztreonam-reacted structure. (b) Structure of PaPBP3:13 (see Chapter 5) which has two alternate conformations for Tyr409. Occupancies were determined by occupancy refinement in Phenix\(^3\) as 58 % and 42 % for the left and right tyrosine respectively. Both conformations form a hydrogen bond with the Thr487 backbone carbonyl. See also Figure 5.6.

6.S2 Kinetics

6.S2.1 Derivation of Equation 6.3

The substrate inhibition equation (equation 6.3) is derived by treating the scheme shown in Scheme 6.1b with the steady state method (derivation modified from \(^4\)). \([ES]\) is assumed to be constant throughout the reaction. This produces three equations. The first is the equilibria of the production (right hand side) and destruction (left hand side) of \([ES]\):

\[
k_{+1}[S][E] - [ES] - [ESS] + k_{-2}[ESS] = (k_{-1} + k_{cat} + k_{+2}[S])[E][S]
\]  

(6.S1)
The second corresponds to the equilibrium between $ES$ and $ESS$:

$$k_{+2} [S] \cdot [ES] = k_{-2} [ESS] \quad (6.2)$$

The third is the uses the assumption that the rate ($v$) is limited by the $k_{\text{cat}}$ (i.e. $k_{\text{cat}} << k_{+1}$):

$$v = k_{\text{cat}} \cdot [ES] \quad (6.3)$$

These simultaneous equations can then be combined and simplified to express the rate in terms of the $[E], [S]$ and the rate constants.

$$v = \frac{k_{\text{cat}} [E]}{1 + \frac{k_{-1} + k_{\text{cat}}}{k_{+1}[S]} + \frac{[S] \cdot k_{+2}}{k_{-2}}} \quad (6.4)$$

Finally we substitute the rate constants for the constant $K_M$ (equation 6.5) and the substrate inhibition constant ($K_{SI}$ equation 6.6) and use the expression for $V_{\text{max}}$ (equation 6.7) to eliminate the $[E]$ term, yielding equation 6.3:

$$K_M = \frac{k_{-1} + k_{\text{cat}}}{k_{+1}} \quad (6.5)$$

$$K_{SI} = \frac{k_{-2}}{k_{+2}} \quad (6.6)$$

$$V_{\text{max}} = k_{\text{cat}} \cdot [E] \quad (6.7)$$

$$v = \frac{V_{\text{max}}}{1 + \frac{K_M [S]}{K_{SI} + [S]}} \quad (6.3)$$
6.S2.2 More Complex Models

Scheme 6.S1 Alternative schemes (a-b) and rate equations (c-d) for substrate inhibition. (a and c) A model that includes turnover of ESS but like Scheme 6.1 only permits the binding of the second nitrocefin to the ES complex; (b and d) a model in which two binding sites can each independently bind a molecule of substrate (in either order), with turnover permitted from the doubly- or singly-reacted species. Schemes and rate equations modified from Yoshino et al.⁵.

6.S3 Tryptic Digest Mass Spectrometry

6.S3.1 Sample Preparation Methods

6.S3.1.1 Protocol I

Filter-Aided Sample Preparation ⁶
The reaction with nitrocefin was conducted at room temperature (Trial A buffer: 500 mM NaCl, 10 mM sodium phosphate, pH 8, Trial B buffer: 300 mM ammonium bicarbonate (ABC)) in a volume of 100 μl (Trial A) or 50 μl (Trial B), and then terminated by the addition of 8 M urea (400 μl). The sample was added to a protein spin column (0.5 mL, 10 kDa molecular weight cutoff, Amicon Ultra) and centrifuged at 8000 x g for 20 minutes. The sample was then buffer exchanged into 50 mM ABC by the sequential addition of ABC (400 μl) then centrifugation to reduce the volume, three times in total. Trypsin (sequencing grade, Promega) was then added and digestion allowed to proceed at room temperature: Trial A: 2 μg trypsin added, digestion overnight; Trial B: 1 μg trypsin added, digestion for 4 hours. After the digestion, the peptides were eluted by centrifugation through a spin column (8000 x
g, 20 minutes), then the spin column was additionally washed with 400 µl of water and centrifuged (8000 x g, 20 minutes). Both the elution and the washes were concentrated by SpeedyVac vacuum concentrator (30 °C, Thermofisher Scientific) until dry, then resuspended in a solution of 2% (v/v) acetonitrile, 0.1% (v/v) trifluoroacetic acid (TFA) in water.

**C₁₈ StageTip**
To ensure sample purity before MS, the sample was purified by C₁₈ fractionation in a “StageTip” (a pipette tip fitted with a filter in an adapter above an eppendorf tube). The StageTip was conditioned by applying the following solutions in order (each 50 µl) and then centrifuging at 2000 rpm: methanol then acetonitrile, then a solution of 2 % (v/v) acetonitrile, 0.1 % (v/v) TFA in water. The sample (150 µl) was then loaded onto the StageTip and centrifuged (15 minutes, 2000 rpm), then washed with first 1 % (v/v) TFA in ethyl acetate and centrifuged (50 µl, 4 minutes, 2000 rpm) then 2 % (v/v) acetonitrile, 0.1% (v/v) TFA in water and centrifuged (50 µl, 4 minutes, 2000 rpm). The peptides were then eluted in 80 % (v/v) acetonitrile in water before being concentrated to a dry powder by vacuum concentration. Peptides were stored at -20 °C, until they were ready to be run on the mass spectrometer (6.S3.2). Before analysis they were resuspended in 2 % (v/v) acetonitrile, 0.1 % (v/v) TFA in water (with 10 minutes ultrasonication to ensure solvation).

**6.S3.1.2 Protocol II**
Due to concerns about the long time taken to conduct the washing steps in Protocol I and III, Protocol II was designed to have minimal washing but required the sample to be pure to avoid contaminating the MS chromatography columns or machine itself.

To remove sodium chloride from the storage buffer of the protein, the protein was first dialysed into 300mM ammonium acetate (pH 8) and the nitrocefin was added (see Table 6.2 for reaction conditions). After heat shock (5 minutes, 99 °C), the sample was quickly cooled on ice, then trypsin was added (Trial C: 1 µg; Trial D: 0.25 µg;) and peptide digestion proceeded at room temperature for 4 hours. The sample was then concentrated by vacuum concentration to a dry mass, then stored at -20 °C until the sample was ready to be analysed (6.S3.2), at which time it was resuspended in 2 % (v/v) acetonitrile, 0.1 % (v/v) TFA in water (Trial C) or water (Trial D) and sonicated for 10 minutes.
6.S3.1.3 Protocol III (in-gel digest)

The reaction with nitrocefin was carried out at room temperature (50 µl reaction volume, buffer: 500 mM NaCl, 10 mM sodium phosphate, pH 8), then the reaction was terminated by unfolding the protein in SDS (8 % (w/v)). The sample was loaded onto a precast TruPAGE polyacrylamide gel (Merck) and run at 150 V for 40 minutes in 25 mM Tris pH 8.3, 190 mM glycine and 0.1% (w/v) SDS. The gel was stained for 10 minutes with coomassie dye (Instant Blue, Exedeon) to allow the bands to be visualised. The protein-containing band was cut out of the gel and then cut into 4 mm slices and washed with a destaining buffer (50 % (v/v) ethanol, 50mM ABC, 3 x 40 µl) until the colour was lost. The gel fragments were dehydrated in ethanol for 5 minutes whilst shaking then trypsin (50 ng in 30 µl of 50 mM ABC) was added and the sample was digested for 4 hours at room temperature. After digestion, the buffer around the gel was collected and the gel fragments were washed with 3 x 20 µl of 25 % (v/v) acetonitrile, 5 % (v/v) formic acid in water with sonication (10 minutes) to extract all peptides. The peptide extractions were concentrated by vacuum concentration and stored at -20 °C until they were ready to be run on the mass spectrometer (6.S3.2). Before analysis, they were resuspended in 2 % (v/v) acetonitrile, 0.1 % (v/v) TFA in water and ultrasonicated for 10 minutes.

6.S3.2 Proteomic Mass Spectrometry Methods

An analysis was performed by LC-MS/MS on a Ultimate 3000-RSLCnano system ( Dionex) coupled to an Orbitrap-Fusion Mass Spectrometer (Thermo-Scientific) as previously described 7. Data was analysed by MaxQuant software 8 against the known sample sequence and the E. coli database. Peptide sequences were assigned to the MS/MS spectra. Acylation by the nitrocefin (516.04 ± 0.5 Da, serine residues), methionine oxidation (15.99 ± 0.5 Da, methionine residues) and acetylation (42.01 ± 0.5 Da, peptide N-terminus) were treated as a variable modifications. Data was viewed in Scaffold4 (Proteome Software).

6.S4 Protein Crystallography Methods

Crystallography was carried out by D. Bellini.
Crystallisation and structure solution of nitrocefin:PaPBP3 adducts. Diffraction quality crystals of PaPBP3\textsubscript{wt} and its PaPBP3\textsubscript{Y409A} were obtained at 294 K using the hanging drop vapour diffusion method from mixing equal volumes of protein with a precipitant solution made up of 25\%(w/v) polyethylene glycol 3350, 0.1M Bis-Tris propane, 1 \%(w/v) protamine sulphate, pH 8. Crystals were cryoprotected with 20 \%(v/v) glycerol prior to flash-cooling in liquid nitrogen. Nitrocefin-protein complexes were obtained by soaking crystals with 10 mM compound for 60 minutes. Diffraction data were collected on Diamond beamlines I03 and I04. Structures were phased using molecular replacement with MrBump \textsuperscript{10}. All data were processed using Dials \textsuperscript{11}. Manual building and ligand fitting was performed with COOT \textsuperscript{12}. Structures were refined with REFMAC5 \textsuperscript{13} and validated with MolProb \textsuperscript{14}. The structures are available online: PDB codes 6HR6 (nitrocefin-reacted PaPBP3\textsubscript{wt}) and 6HR9 (nitrocefin-reacted PaPBP3\textsubscript{Y409A}).

6.S5 References


Chapter 7. Concluding Remarks

Antimicrobial resistance will be one of the greatest healthcare challenges of the coming decades and loss of these clinical fire extinguishers could lead to the deaths of millions. But tackling this challenge is made more difficult by intertwining socio-economic and scientific hurdles as well as the need for broad engagement with the necessary solutions. The intractability of the problem and a lack of profitability of antimicrobial research has led to decades of neglect that only recently is being properly addressed. As progressive resistance makes current antimicrobials ineffective, new therapies must be discovered and approved to replace them. The challenges of antimicrobial resistance are perennial and will necessitate continued and ongoing research. Originality in the mode of action of new inhibitors is desirable, and new methods, including those discussed here, are needed in the search for this novelty.

The case of β-lactams is emblematic of the crisis in antimicrobial discovery: PBP is a well-established and effective target, but no novel inhibitors have been approved since aztreonam in the 1980s. This is not without attempts; over the last 20 years, numerous academic groups and some pharmaceutical companies have published research reporting non-β-lactam PBP inhibitors but none of these compounds even approach the efficacy of β-lactams. This perhaps indicates how challenging this task is and once again underlines how well β-lactams are adapted for PBP inhibition. Boron based inhibitors (e.g. vaborbactam) have proven their efficacy against β-lactamases, but PBP-targeting boronates have yet to be realised as antimicrobials, for reasons that remain elusive.

Our fragment screen of serine covalent inhibitors found for the first time that boronates could bind to a high molecular mass PBP in di- or tri-covalent binding modes (depending on the warhead). Whilst the present on-target affinity is modest, further optimisation of derivatives (facilitated by 10 new structures of PBP:boronate complexes) may allow for improved antimicrobial activity. Utilisation of novel assays which employ natural substrates may allow better assessment of the ability of the compounds to inhibit the physiological functions of PBPs. This could lead to compounds with better in vitro efficacy.
The proposed role of boronates as transition state inhibitors of the reactions of PBPs with their substrates may be extended to explain the novel observation of two molecules of nitrocefin in the active site. Which I hypothesise may occur as the result of the transfer of the nitrocefin adduct from one active site serine to another. Whatever the mechanism, observation of the two nitrocefin molecules in the active site of mutant PBP3 is interesting and development of the observation could yield new inhibitors which exploit this phenomenon.

It is possible that non-β-lactam inhibitors will catalyse the further propagation of target-mediated resistance as β-lactamases become ineffective. In PBPs from gram-pathogens, the mechanisms underlying target-mediated resistance are not well understood but the frequent occurrence of clinical mutations in flexible loops adjacent to the active site suggests the loops have a role to play. Boronate compounds complexed to PBP3 show that signals can be transmitted between the active site and loops not in direct contact with the compound, perhaps by a network of interacting amino acids.

Finally this thesis has explored two tools to aid in drug discovery. A robotic liquid handling system has been utilised for antimicrobial screens which will find use in routine screening, aiding drug discovery. Attempts to characterise the interactions of PBP3 using a surface plasmon resonance, which had previously never been reported, were ultimately unsuccessful. One of the problems preventing development of the system was a lack of high affinity, reversibly binding compounds to act as a positive control. The outlook for finding such inhibitors is positive but further understanding of the protein and of inhibitors is required.
Appendix: Publications by the Author

This work formed the basis for Chapter 4 and is summarised in section 4.1.

The author contributed structural activity relationship input to this work, discussed in section 1.3.3 and published two crystal structures (PDB: 6Y6U and 6Y6Z)

Chapter 5 was modified and submitted for publication: it was accepted, reviewed, revised and is currently returned to the reviewers for their approval (as of 11th June 2021). As part of this, 10 crystal structures (5.S2) were published on the PDB.
As this is not yet publicly available, the supporting information is also included.