Characterisation of the MacA/MacB/TolC tripartite pump that confers resistance to macrolides in E. coli

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Characterisation of the MacA/MacB/TolC tripartite pump that confers resistance to macrolides in *E. coli*

By
Hong-Ting Lin M.Sc.

A thesis submitted for the degree of Doctor of Philosophy
In
The School of Biological and Biomedical Sciences
Centre for Infectious Diseases
University of Durham

November 2008

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Hong Ting Lin
November 2008
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Finally, I am indebted to my parents, family, friends and my beloved, Chen-Ju for their perpetual patience, encouragement and support. My little boys, Jing-Huan and Sing-Rung always bring me priceless joy.
Abstract

Gram-negative bacteria possess tripartite pumps, composed of a membrane fusion protein (MFP), an inner membrane protein (IMP) and an outer membrane protein (OMP), to transport drugs across the inner and outer membranes. The plasmid encoding MacA, MacB and TolC confers resistance to the macrolide erythromycin in the host *E. coli* cell Kam3, indicating the three proteins are assembled and actively functional. MFPs are believed to have an important role in the stabilizing the pump complex; intriguingly, we found that the MFP MacA not only interacts directly with the IMP MacB and the OMP TolC, but regulates the function of MacB, apparently increasing its affinity for both ATP and erythromycin. As MacB hydrolyzes ATP there is a burst in phosphate production that is symptomatic of the reaction being rate-limited by product release; but the burst disappeared in the presence of MacA. Since MacA caused only a marginal increase in the $k_{\text{cat}}$, but a significant decrease in the $K_m$, for the steady-state ATPase activity, this suggests that the disappearance of the phosphate burst is due to a decrease in the rate of hydrolysis, rather than an increase in the rate of product release. This kinetic behaviour indicates that MacA promotes and stabilizes the ATP-binding form of the transporter. MacA regulates the activity of MacB via its β-strand domain since *S. aureus* MacA, which lacks the coiled coil structure that is present in *E. coli* MacA and believed to be involved in the interaction with TolC, was able to abolish the Pi burst catalysed by MacB, in direct analogy with the effect of *E. coli* MacA on MacB. Analytical ultracentrifugation, mass spectrometry and atomic force microscopy indicated that MacB forms dimers, in analogy to ABC-transporters that require a pair of NBDs to bind ATP. Our data suggests a direct role for MacA in facilitating the delivery of drugs by MacB to TolC: by enhancing the binding of drugs to MacB and stabilizing the reorientation of MacB to the outward-facing conformation.
Abstract

Gram-negative bacteria possess tripartite pumps, composed of a membrane fusion protein (MFP), an inner membrane protein (IMP) and an outer membrane protein (OMP), to transport drugs across the inner and outer membranes. The plasmid encoding MacA, MacB and TolC confers resistance to the macrolide erythromycin in the host *E. coli* cell Kam3, indicating the three proteins are assembled and actively functional. MFPs are believed to have an important role in the stabilizing the pump complex; intriguingly, we found that the MFP MacA not only interacts directly with the IMP MacB and the OMP TolC, but regulates the function of MacB, apparently increasing its affinity for both ATP and erythromycin. As MacB hydrolyzes ATP there is a burst in phosphate production that is symptomatic of the reaction being rate-limited by product release; but the burst disappeared in the presence of MacA. Since MacA caused only a marginal increase in the $k_{cat}$, but a significant decrease in the $K_{m}$, for the steady-state ATPase activity, this suggests that the disappearance of the phosphate burst is due to a decrease in the rate of hydrolysis, rather than an increase in the rate of product release. This kinetic behaviour indicates that MacA promotes and stabilizes the ATP-binding form of the transporter. MacA regulates the activity of MacB via its $\beta$-strand domain since *S. aureus* MacA, which lacks the coiled coil structure that is present in *E. coli* MacA and believed to be involved in the interaction with TolC, was able to abolish the Pi burst catalysed by MacB, in direct analogy with the effect of *E. coli* MacA on MacB. Analytical ultracentrifugation, mass spectrometry and atomic force microscopy indicated that MacB forms dimers, in analogy to ABC-transporters that require a pair of NBDs to bind ATP. Our data suggests a direct role for MacA in facilitating the delivery of drugs by MacB to TolC: by enhancing the binding of drugs to MacB and stabilizing the reorientation of MacB to the outward-facing conformation.
### Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>ABC</td>
<td>ATP-binding cassette</td>
</tr>
<tr>
<td>ADP</td>
<td>Adenosine diphosphate</td>
</tr>
<tr>
<td>AFM</td>
<td>Atomic force microscope</td>
</tr>
<tr>
<td>AMP-PNP</td>
<td>(\beta, \gamma)-imidoadenosine 5'-triphosphate tetralithium salt</td>
</tr>
<tr>
<td>AMS</td>
<td>4-acetoamide-4'-maleimidylstilbene-2, 2'-disulfonic acid</td>
</tr>
<tr>
<td>AUC</td>
<td>Analytical ultracentrifugation</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>CCCP</td>
<td>Cyanide-M-chlorophenyl hydrazone</td>
</tr>
<tr>
<td>CD</td>
<td>Circular dichroism</td>
</tr>
<tr>
<td>DDM</td>
<td>n-dodecyl-(\beta)-D-maltopyranoside</td>
</tr>
<tr>
<td>DLS</td>
<td>Dynamic light scattering</td>
</tr>
<tr>
<td>DME</td>
<td>Drug-metabolite efflux</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
</tr>
<tr>
<td>EM</td>
<td>Electron microscope</td>
</tr>
<tr>
<td>ES</td>
<td>Electro-spray</td>
</tr>
<tr>
<td>FTIR</td>
<td>Fourier transform infrared spectroscopy</td>
</tr>
<tr>
<td>(k_{\text{cat}})</td>
<td>Catalytic constant</td>
</tr>
<tr>
<td>(K_m)</td>
<td>Michaelis constant</td>
</tr>
<tr>
<td>IPTG</td>
<td>Isopropyl-(\beta)-D-thiogalactopyranoside</td>
</tr>
<tr>
<td>IMP</td>
<td>Inner membrane protein</td>
</tr>
<tr>
<td>MALDI</td>
<td>Matrix-assisted laser desorption/ionization</td>
</tr>
<tr>
<td>MATE</td>
<td>Multi and toxic compound extrusion</td>
</tr>
<tr>
<td>MDR</td>
<td>Multidrug resistance</td>
</tr>
<tr>
<td>MESG</td>
<td>2-amino-6-mercapto-7-methylpurine riboside</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------</td>
</tr>
<tr>
<td>MFP</td>
<td>Membrane fusion protein</td>
</tr>
<tr>
<td>MFS</td>
<td>Major facilitator superfamily</td>
</tr>
<tr>
<td>MIANS</td>
<td>2-(4’-maleimidoanilino) naphthalene-6-sulfonic acid</td>
</tr>
<tr>
<td>MOPS</td>
<td>4-Morpholinepropanesulfonic acid</td>
</tr>
<tr>
<td>MS</td>
<td>Mass spectrometry</td>
</tr>
<tr>
<td>NBD</td>
<td>Nucleotide binding domain</td>
</tr>
<tr>
<td>NEM</td>
<td>N-ethyl maleimide</td>
</tr>
<tr>
<td>NRMSD</td>
<td>Normalized root mean squared deviation</td>
</tr>
<tr>
<td>OMP</td>
<td>Outer membrane protein</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>P-gp</td>
<td>P-glycoprotein</td>
</tr>
<tr>
<td>Pi</td>
<td>Inorganic phosphate</td>
</tr>
<tr>
<td>RE</td>
<td>Restriction enzyme</td>
</tr>
<tr>
<td>RND</td>
<td>Resistance-nodulation-cell division</td>
</tr>
<tr>
<td>SBP</td>
<td>Substrate binding protein</td>
</tr>
<tr>
<td>SD</td>
<td>Shine Dalgarno</td>
</tr>
<tr>
<td>SEC</td>
<td>Size exclusion chromatography</td>
</tr>
<tr>
<td>SMR</td>
<td>Small multidrug resistance family</td>
</tr>
<tr>
<td>TM</td>
<td>Transmembrane</td>
</tr>
<tr>
<td>TMD</td>
<td>Transmembrane domain</td>
</tr>
<tr>
<td>TPP⁺</td>
<td>Tetraphenylphosphonium</td>
</tr>
<tr>
<td>$V_{max}$</td>
<td>Maximum Velocity</td>
</tr>
</tbody>
</table>
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Membrane transporters are involved in the translocation of drugs and antibiotics, uptake of nutrients, protein secretion, toxin production, photosynthesis and many other vital functions in all organisms (Henderson, 1998). Such membrane proteins are notoriously difficult to study due to their hydrophobicity and low expression levels. These difficulties can explain that whilst membranes proteins account for 30% of all cells proteins, only around 160 membrane proteins structures have been solved, whereas the structures of over 8,000 soluble proteins have been solved (Michel, 2006; White, 2008).

Multidrug transporters, as implied by the name, confer resistance to multiple drugs by pumping them out of cells, decreasing the intracellular concentration of the drug to subtoxic levels (Borges-Walmsley, 2003). It therefore not only leads to prolonged and more complicated treatments but also an increase in the cost of diagnoses and cures. Recent progress in sequencing of microbial genomes and protein crystallography provide opportunities to identify novel antibacterial targets for the new development of drugs. To prevent and control infectious diseases caused by the multidrug-resistant bacteria, more research into multidrug transporters, which are recognized as a major contributor to multidrug resistance, are urgently required.

The MacB protein, an ATP binding casstte (ABC) macrolide antibiotic resistance transporter from *E. coli* (Kobayashi *et al.*, 2001), was studied in order to determine its biochemical and structural properties. Our interest was in determining the mechanism of regulation of the ATPase activity of MacB by MacA and the function of the tripartite complex pump: MacA/MacB/TolC. The ABC transporter MacB was easier to investigate than other pumps, such as those belonging to the RND family in terms of its interaction with MacA due its ATPase characteristic. Importantly, an understanding of the MacA/MacB/TolC system may give us more understanding of type I secretion systems in bacteria, such as the HlyB/HlyD/TolC system, where HlyB is an ABC transporter (Thanabalu *et al.*, 1998).

In the Type I secretion system, a continuous channel is formed by the assembly of multiple proteins into a complex that spans the inner, outer membranes and periplasmic region. The mechanisms of the type II and type V secretion systems are
characterized by a two-step process that involves periplasmic intermediates. The type III secretion system was mediated by approximately 20 proteins and it is sec-independent. The type IV secretion system is involved in genetic exchange, transfer of DNA (Gerlach and Hensel, 2007).
1.1 Antibiotics

Antibiotics, which literally means "against life", were originally produced by bacteria or fungi. They are used to attack the specific agent causing the infection, hence helping the infected host to eliminate the infection. However, extensive use of antibiotics has raised a serious public health problem due to multidrug resistance, in which bacteria become resistant to multi-antibiotics. A common clinical problem in patients with infectious diseases as well as in patients with cancer, multidrug resistance, a phenomenon which enable a disease-causing prokaryotic or eukaryotic microorganisms to resist distinct drugs of a wide variety of structure and function targeted at eliminating the organism. The efflux of drugs by the multidrug transporter p-glycoprotein is one of the principal ways for multidrug resistance.

To combat diseases caused by multidrug resistant bacteria, we need to learn more about the physiology of pathogens to aid in designing new drugs and extend the life of antibiotics (Alanis et al., 2005).
1.1.1 Target of antibiotics

Antibiotics are usually classified on the basis of their chemical structures and modes of action. The main classes of antibiotics inhibit four classical targets: (a) cell wall biosynthesis, (b) protein biosynthesis, (c) DNA and RNA biosynthesis, and (d) other targets, such as cytoplasmic membrane biosynthesis.

**Inhibition of bacterial cell wall biosynthesis**

β-lactam antibiotics, such as penicillin, penicillin G and amoxicillin, can inhibit the formation of peptidoglycan crosslinkings in the bacterial cell wall. The β-lactam (Figure 1.1) moiety of penicillin binds to the transpeptidase that links the peptidoglycan molecules in bacteria, weakening the cell wall of the bacteria, that can lead to cell cytolysis (Spratt and Cromie, 1988).

![The β-lactam ring](image)

**Figure 1.1 Penicillin core structure; with the β-lactam ring indicated.**

**Inhibition of protein biosynthesis**

There are a large number of steps involved in protein synthesis, and many kinds of antibiotics are used to bind to bacteria ribosome and block protein synthesis. This action is mainly bacteriostatic, but can also be bactericidal at higher concentrations. Bacteriostatic antibiotics interfere with bacterial growth, while bactericidal antibiotics kill bacteria. These two actions are accomplished by different mechanisms. Bactericidal actions can disrupt bacterial cell wall or plasma membrane, and bacteriostatic actions can interfere with protein synthesis, nucleic acid synthesis, or other vital metabolic processes, which may become bactericidal when the bacteria
deplete some essential materials, such as ATP, a main energy source for the majority of cellular functions.

For example, aminoglycosides and tetracyclines can interact with the 30S subunit of the bacterial ribosome; macrolides (Figure 1.2) and chloramphenicol can bind to the 50S subunit of the bacterial ribosome, therefore inhibiting translocation of tRNA (Brodersen et al., 2000; Katz and Ashley, 2005).

![Figure 1.2 Core structure of erythromycin. A first generation macrolide, with the macrolide ring indicated upper left.](image)

**Inhibition of DNA and RNA biosynthesis**

DNA replication is an essential process for all organisms. On the premise that eukaryotic cells do not contain DNA gyrase or topoisomerase IV, quinolone and fluoroquinolones antibiotics such as norfloxacin are used to inhibit the DNA gyrase or the topoisomerase IV enzyme to stop DNA replication and transcription, therefore actively killing bacteria (Li, 2005). Rifampicin inhibits RNA polymerase by binding to the β subunit of RNA polymerase, thus preventing transcription of mRNA and its subsequent translation into proteins (Campbell et al., 2001).

**Other targets**

The current generation sulfa drug sulfamethoxazole is used in combination with trimethoprim and each drug inhibits distinct steps in folic acid metabolism (Huovinen, 2001).

Antimicrobial peptides, secreted from both gram-positive and gram-negative bacterial, have been classified within the bacteriocins. Most antimicrobial peptides
such as nisin can attack cytoplasmic membranes, permeabilize target membranes, eventually leading to cell death (Hancock and Chapple, 1999).

1.1.2 Antibiotic resistance

Antimicrobial resistance is a growing problem in the world. For instance, the first case of antimicrobial resistance occurred in the late 1930s and 1940s, soon after the introduction of the first antibiotic classes, sulfonamides and penicillins (Alekshun and Levy, 2007). Common bacteria, such as strains of *S. aureus*, became resistant to these classes of antibiotics. In 2002, a report indicated that strains of Group A *Streptococcus* are becoming resistant to macrolides and spreading outside the hospital, causing community-acquired infections (Martin *et al.*, 2002).

The major mechanisms of antibiotic resistance are as follows:

**Avoiding accumulation of antibiotics**

The prevention of antibiotic action is by either increasing the efflux or decreasing the influx of antibiotics across biological membranes. Cells having mutant TolC, R367H or R390C, show hypersensitivity to drugs, which is presumably caused by having TolC tunnel open, which allow more antibiotics entered the cell than were expelled (Augustus *et al.*, 2004).

Drugs are pumped out by multidrug transporters faster than they can diffuse across membranes, so the drug level inside can be kept low, at levels that are much less effective. Multidrug transporters present in both gram-positive and gram-negative bacteria confer resistance to a wide variety of drugs and this is now recognized as a major mechanism for multidrug resistance in bacteria (Alekshun and Levy, 2007).

**Destruction of antibiotics by either modification or hydrolysis**

Enzymatic inactivation of antibiotics is a major mechanism of resistance in pathogenic bacteria to natural antibiotics such as β-lactams, aminoglycosides and chloramphenicol (Walsh, 2000). The classic case is the hydrolysis of the four-
membered β-lactam ring in penicillins by β-lactamases from resistant bacteria, resulting in the inactivation of the antibiotics.

**Modification of the drug target**

The antibiotic mechanism of target alternation is by modification of the normal target so that it does not bind the antibiotics. As previously stated, erythromycin can bind to the 50S subunit of the bacterial ribosome to inhibit the translocation of tRNA. The methyl transferase enzyme Erm can methylate a specific adenine residue of the 23S RNA component of the ribosome without the interference of protein biosynthesis and lower the affinity of the macrolides for the RNA (Walsh, 2000).
1.2 Multidrug transporters

The efflux of cytotoxic compounds out of cells by transmembrane proteins is one of the most important mechanisms for drug resistance in prokaryotes and eukaryotes (Borges-Walmsley et al., 2003). Most multidrug transporters have been divided into two groups by their energy requirements: those membrane transporters belonging to ATP binding Cassette (ABC) transporters which can couple transport to the hydrolysis of ATP; and secondary active transporter, which can use the potential energy of the electrochemical gradients of H⁺ or Na⁺ across the membrane (Figure 1.3).

1.2.1 Powerful multidrug transporters: What are they?

Secondary active transporters have been categorized into several families by their sequence similarities: these include the Resistance-Nodulation-cell Division (RND) family, the Major Facilitator (MF) superfamily, the Small Multidrug Resistance (SMR) family, the Multi and Toxic Compound Extrusion (MATE) family and the Drug-Metabolite Efflux (DME) family; while the DME family has only recently been described and are not well characterized, they are believed to have some similarity to the SMR family (Jack et al., 2001).
1.2.1 Secondary active transporters

MF family

The MF transporters form the largest family of the secondary active transporters (Figure 1.4) (Saidijam et al., 2006), and these MF family proteins transport drugs, ions, sugars sugar-phosphates, nucleosides, amino acids, and peptides.

In 1998, transport proteins from eukarya, archaea and bacteria were categorized on the basis of their molecular phylogeny (Saijijam et al., 2006) and transport proteins for each organism are classified on the basis of their energy coupling mechanism, protein phylogenic family and substrate specificity. Drug efflux proteins fall into five classes, where MF and ABC superfamily are the largest families among them.

MF transporters are typically 400 to 600 amino acids long and arranged into 12 trans-membrane α-helices (Figure 1.5), with a large cytoplasmic loop between helices six and seven (Saier et al., 1999). A smaller number of these transporters have 14 trans-membrane α-helices, such as the *S. aureus* tetracycline and divalent cation transporters TetA(K) (Ginn et al., 1997) and QacA, respectively (Paulsen et al., 1996).

![Distribution of Drug Efflux Systems](image)

**Figure 1.4** Distribution of putative drug efflux systems in different microorganisms. The colors signify as follows from the bottom: light blue, ABC; red, MFS; green, RND; dark blue, MATE; and orange, SMR. See ref. Saidijam et al., 2006 for more formation.
MdfA

MdfA, a 410 residue protein arranged into 12 transmembrane helices (Figure 1.5) (Adler et al. 2004; Adler and Bibi, 2002), confers resistance to a remarkably diverse group of cytotoxic compounds, including cationic, zwitterionic and uncharged substrates (Edgar and Bibi, 1997; Nishino and Yamaguchi, 2001; Lewinson et al., 2006).

Figure 1.5 Secondary structure model of MdfA (Adler et al., 2004)

Early studies revealed that the charged residue located in the transmembrane domain, Glu$^{26}$ in helices 1, is important in substrate recognition (Edgar and Bibi, 1999). Replacement of Glu$^{26}$ with the positively charged residue lysine eliminated the resistance against the positively charged drug ethidium bromide, but retained the resistance to chloramphenicol. Interestingly, when Glu$^{26}$ was replaced by the negatively charged residue aspartate, chloramphenicol recognition and transport greatly decreased; however, resistance against lipophilic cations remained. In addition, mutagenesis studies revealed many second site mutations that restored the function of inactive E26 mutants in chloramphenicol. Several of them were differentially affected by different substrates, suggesting different drugs may interact with different regions of the pocket (Adler and Bibi, 2004). A similar observation was made using direct binding studies, which showed positively charged drugs competed with the cationic compound tetraphenylphosphonium (TPP$^+$) in binding to MdfA, whereas some zwitterionic drugs did not. Besides, the neutral drug chloramphenicol stimulated the binding of TPP$^+$ by increasing its affinity for MdfA, implying that the two drugs could simultaneously bind to two distinct sites (Lewinson and Bibi, 2001).
*EmrD*

The MF family transporter EmrD structure was reported and the transport mechanism was proposed in 2006 (Yin et al., 2006). EmrD was first identified as an efflux pump transporting uncouplers of oxidative phosphorylation such as cyanide-M-chlorophenyl hydrazone (CCCP) (Naroditskaya et al., 1993), but subsequently reported to be able to export a broad spectrum of hydrophobic compounds (Nishino and Yamaguchi, 2001). The topology of EmrD is shown in figure 1.6a; EmrD has 12 transmembrane α helices and its N- and C- termini are located in the cytoplasm. As illustrated in figure 1.6b H3, H6, H9 and H12 face away from the interior and are not involved in the pore formation; on other hand, the four central helices H1, H4, H7, H10 and the four lateral helices H2, H5, H8, H11 surround a central cavity, which consists mostly of hydrophobic residues, allowing transportation of lipophilic compounds. Among them are the two pairs of stacked aromatic groups, Tyr^{52}/Tyr^{56} and Trp^{300}/Phe^{249}, which could be involved in substrate binding. In Bmr, a close homolog of EmrD, two phenylalanines have been implicated in substrate recognition. Yin et al in 2006 proposed that two long helical regions, (H4/L4-5/H5 and H10/L10-11/H11), and H1, located on the cytoplasmic side, could serve as a selectivity filter. A potential mechanism for drug transport is that CCCP can enter the central cavity, either through the membrane leaflet or cytoplasm, and is recognized by the long helical filter region, and is subsequently bound by hydrophobic residues in the cavity (Figure 1.6c).
Figure 1.6  The structure of EmrD by ribbon presentation. (a) Side view of the EmrD. (b) Top view of the EmrD. (c) The proposed mechanism of the EmrD transporter (Yin *et al*., 2006).
SMR family

The smallest multidrug transporters belong to the SMR family, which are around 100 amino acids long. The SMR family transporters extrude various drugs in exchange for protons, conferring bacteria resistance to these compounds (Yerushalmi et al., 1995; Schuldiner et al., 2001).

EmrE

![Secondary structure model of EmrE-His](image)

Figure 1.7  **Secondary structure model of EmrE-His.** Putative transmembrane domains are shown in boxes connected by hydrophilic segments. Charged residues are highlighted and the ones conserved in EmrE homologs are shown with black circles (Yerushalmi and Schuldiner, 2000).

The best studied SMR transporter EmrE from *E. coli*, which confers resistance to ethidium bromide and methyl viologen, is putatively arranged into four transmembrane α helices (Figure 1.7) (Yerushalmi et al., 1995; Yerushalmi and Schuldiner, 2000; Ninio et al., 2004). Cysteine scanning mutagenesis and FTIR experiments indicated that EmrE is composed of a tightly packed bundle of α-helices without any continuous aqueous domains, implying substrates are transported through a hydrophobic pathway in the protein (Mordoch et al., 1999; Arkin et al., 1996). Further study on the identity of amino acid residues in the substrate binding and transportation pathway have been provided by site-directed mutagenesis and biochemical experiments. Seven of the eight charged amino acids located in putative loops of EmrE can be replaced with either cysteine or other amino acids bearing the same charge without significant change on transport activity. The other charged amino
acid Glu^{14}, a carboxylic residue conserved in all EmrE homologs, located in the first putative transmembrane domain is essential for substrate binding and efflux of the drug TPP^+ (Yerushalmi and Schuldiner, 2000). In addition, further data exploring the drug binding domain indicated that some aromatic residues from transmembrane domain 2 and 3 form part of the binding site (Sharoni et al., 2005).

Although the oligomeric state in vivo is still under dispute, the minimal functional unit for substrate binding is a dimer (Butler et al., 2004; Ubarretxena-Belandia et al., 2003). The EmrE-detergent complex was extracted from membranes in a form that retains the ability to bind the drug TPP^+ in a 3 to 1 molar ratio (Muth et al., 2000), which may be due to some inactive monomer. A projection structure of EmrE at 7.0 Å resolution 2D crystals showed the repetitive unit in the crystal comprised of eight α-helices arranged in an asymmetric manner, indicating the minimal functional unit for EmrE was a dimer (Figure 1.8a) (Tate et al., 2001). When TPP^+ binds to EmrE in the 2D crystals, a small conformational change occurs that is shown to be mainly due to the movement of a single α-helix (Tate et al., 2003). The recent X-ray EmrE-TPP^+ structure revealed another characteristic in the EmrE structure, dual topology (Chen et al., 2007). In this structure, the SeMet sites are related by a pseudo-twofold rotational axis running along the dimer interface parallel with the membrane plane. Conserved Glu^{14} point toward the binding chamber and appear well placed to form ionic contacts with the positively charged TPP molecular (Figure 1.8b). As illustrated in 1.8c, it shows the Se sites in the first transmembrane helices (residues 21 and 21*) are located pseudo-symmetrically on opposite sides of the bilayer. This feature indicates that the X-ray structure of the EmrE-TPP^+ is an antiparallel dimer.

This EmrE structure is consistent with some biochemical data; for example, the membrane-embedded glutamate (E^{14}) is involved in substrate binding and translocation; the conserved Trp^{63} in the third transmembrane helix has been identified as an essential residue that is involved with binding of the substrate and protein activity (Elbaz et al., 2005); in the same helix, Tyr^{60} is also essential for activity; Tyr^{40} is conserved in the EmrE branch of the SMR family, and its replacement has an effect on the specificity of the transporter to substrates (Soskine et al., 2004; Sharoni et al., 2005). Tryptophan residues may play several roles in integral membrane proteins. For example, tryptophan residues can maintain interactions between a protein and its substrate using hydrophobic stacking interactions (Vazquez-Ibar et al., 2003); they
may also participate in cation/π interactions between protein and their ligands (Zacharias and Dougherty, 2002); moreover, the locations of tryptophan residues of membrane proteins tend to be at the ends of transmembrane α-helices (Arkin and Brunger, 1998), reflecting the amphipathic nature of a tryptophan residue that has the ability to form hydrogen bonds, as well as to exhibit hydrophobic character.

The proposed mechanism of transport involves the binding of TPP$^+$ by the two membrane-embedded Glu$^{14}$ residues located in transmembrane domain 1 (Figure 1.8d). The substrate enters EmrE either from the inner leaflet of the cytoplasmic membrane or directly from the cytoplasm. The binding site, occupied by the substrate, triggers a conformational change so that it becomes accessible to the other face of the membrane. The direct binding of protons to both Glu$^{14}$ residues induces the release of substrate at the periplasmic surface. The chamber where TPP$^+$ binds is sealed at the top by helix H; the creation of an opening on the opposite side of the membrane from the chamber entrance would require the move of this helix, whilst the proximity of helices G and F could mean they move as a group.
Figure 1.8 Structure of EmrE. (a) Projection map of EmrE at 7.0 Å resolution. The interpretation of each mark are as follows: P, an α-helix nearly perpendicular to the membrane plane; T, probably a single α-helix tilted with respect to the membrane normal; A, an arc of probably for tilted α helices (Tate et al., 2001). (b) Top view of the EmrE-TPP+ structure, with transmembrane helices labeled. Red spheres represent the residues that have been implicated in substrate binding and transport by biochemical and mutagenesis studies (Chen et al., 2007). (c) Front view of the EmrE-TPP+ structure, with two SeMet markers. (d) The proposed mechanism of transport by EmrE (Ubarretxena-Belandia et al., 2003).
RND family

RND transporters can be found in all three kingdoms of life, Eukarya, Acrhaea and Eubacteria. RND transporters are composed typically of 1000 amino acid residues and are much larger than other secondary active transporters (Borges-Walmsley et al., 2003). A characteristic feature of the topology of RND transporters is the presence of 2 large periplasmic loops between transmembrane helices 1 and 2 and 7 and 8. Since the N-terminal halves of RND transporters are homologous to the C-terminal halves, this suggests that these transporters have arisen by gene duplication.

AcrB

AcrB, the best studied member of the RND family, is a multidrug transporter that confers resistance to a wide range of antibiotics; including tetracycline, chloramphenicol, β-lactams, novobiocin, fusidic acid, nalidixic acid, and fluoroquinolones; the detergent SDS and Triton X-100; various cationic dyes, disinfectants, and even solvents (Poole, 2001).

The first crystal structure of AcrB was obtained at 3.5 Å resolution. The three protomers are fixed symmetrically by the crystal packing force (Murakami et al., 2002). The appearance of the AcrB trimer is like a jellyfish with a three-fold symmetry axis perpendicular to the membrane plane. It comprises a periplasmic headpiece and a transmembrane region with approximate dimensions of 70 Å by 50 Å, respectively. The AcrB structure contains three domains: the transmembrane (TM) domain, the porter domain and the TolC docking domain (Figure 1.9a). The porter domain is composed of four subdomains, PN1, PN2, PC1 and PC2 for each protomer. PN1 and PN2 locate between TM1 and TM2 and PC1 and PC2 locate between TM7 and TM8. Each subdomain contains two characteristic β-strand-α-helix-β-strand motifs. These four subdomains are packed with their β-sheets surrounding a substrate binding pocket (Figure 1.9b).

More recently, the structure of new crystal forms of AcrB have been determined in which each protomer adopts a different conformation (Murakami et al., 2006; Seeger M. et al., 2006), leading to a rotating pump mechanism.
Figure 1.9  **AcrB structure.** (a) The side view of the AcrB structure in ribbon representation. (b) The topology diagram of the protomer (Murakami et al., 2002).
Bound minocycline was observed in one of the three protomers at the distal end of the pocket, between the β-sheets of PN2 and PC1 (Figure 1.10a). The substrate binding pocket is rich in aromatic amino acid residues which may interact with the drug molecule by hydrophobic or aromatic-aromatic interactions. It has been shown that two different sets of amino acid can interact with different drugs, indicating different sets of residues are used for binding of different kinds of substrates.

If we compare the protomer involved in extrusion to the one involved in binding drugs, an asymmetric difference can be recognized (Sennhauser et al., 2007; Das et al., 2007; Seeger et al., 2008). PN2 and PC1 are moved back towards PN1 and PC2 in the extrusion protomers, making the pocket too small to bind substrates. The inclined central helix (Figure 1.10a) from the extrusion protomer fills the possible “exit gap” between PN1 and PN2 in the binding protomer and creates a space between PN1 and PN2 in the extrusion protomer opening to the top funnel. Based on the different conformations of the three protomers of AcrB, a three step, (access, binding and extrusion), mechanism was proposed (Figure 1.11). In the access state, the vestibule is open to the periplasm, but the binding pocket is shrunken, making it impossible for substrates to have access to the vestibule. In the binding state, the vestibule is kept open and the binding pocket is expanded for substrate binding. At this stage, exit from the binding site is blocked by the inclined central helix from the extrusion protomer. Final, in the extrusion state, the squeezing of the binding pocket pushes the bound drugs out via the top funnel. Theses protomer conformation can be expected to be coupled with proton translocation across the membrane. Asp\textsuperscript{407}, Asp\textsuperscript{408} and Lys\textsuperscript{940} in transmembrane helices have been identified as essential amino acid residues for AcrB function and are probably involved in proton translocation. In the access and binding protomers, Lys\textsuperscript{940} of TM10 is coordinated by salt bridges with Asp\textsuperscript{407} and Asp\textsuperscript{408} of TM4. In contrast, Lys\textsuperscript{940} in the extrusion protomer is turned towards Thr\textsuperscript{978} of TM11 and this causes movement of TM4 and TM10 (Figure 1.10b). Site mutagenesis and drug susceptibility studies have also shown that Thr\textsuperscript{978} is essential for drug transport (Takatsuka and Nikaido, 2006). Unfortunately, how the twisting of these transmembrane helices causes the movement of the subdomains and the central α-helix inclination is still unclear.
Figure 1.10  The asymmetric crystal structure of AcrB-minocycline complex. (a) Cut view of the porter domain from the distal side of the cell. The bound minocycline molecule is colored in orange. (b) The bottom view of the AcrB transmembrane domain structure from the periplasmic side (Murakami et al., 2006).

Figure 1.11  Schematic illustration of the proposed transporter mechanism. A, B and E represents the Access, Binding and Extrusion conformation respectively. (a) Side view of the proposed tripartite efflux pump AcrA-AcrB-TolC. (b) The proposed conformational change mechanism of three protomers of AcrB in a transport cycle (Schuldiner, 2006).
MATE family

MATE transporters can be found in all three kingdoms of life, Eukarya, Archaea and Eubacteria. In bacteria, these transporters are around 450 amino acids long and putatively arranged into 12 transmembrane α-helices (Figure 1.12) (Singh et al., 2006). Based on the similar size and topology, MATE transporters were once suggested to be members of the MF family; however, they have no similarity in protein sequence with MF transporters (Brown et al., 1999), such as the five-residue motif (RXGRR), located between TM2 and TM3. (Pao et al., 1998)

![Figure 1.12 Putative topology of NorM, with the 12 transmembrane helices boxed (Singh et al., 2006).](image)

**NorM**

Of all multidrug transporters, our knowledge of the MATE transporters is most rudimentary because they have had little study. NorM, a Na⁺/multidrug anti-porter, first found in *V. parahaemolyticus*, confers resistance to hydrophilic fluoroquinolone such as norfloxacin, ciprofloxacin and ethidium bromide, is the best characterized MATE transporter (Morita et al., 1998). The topology of NorM from *V. cholerae* originally predicted by the algorithm HMMTOP was validated by fusing C-terminal truncations of NorM to either the TEM β-lactamase or chloramphenicol acetyltransferase (CAT) reporter (Figure 1.12). These studies indicated that NorM has 12 transmembrane domains and both the N- and C-terminal ends are located in the cytoplasm. In addition, some residues located on the hydrophilic loops are suggested to be involved in norfloxacin efflux (Singh et al., 2006). Mutagenesis experiments indicated that the acidic amino acid residues Asp32, Glu251, and Asp367, highly conserved in cluster 1 MATE transporters, located on the putative transmembrane domains 1, 7 and 10 of NorM, are involved in Na⁺-dependent drug transport process (Otsuka et al., 2005).
1.2.2 ABC transporters

1.2.2.1 What are they?

ATP binding Cassette (ABC) transporters can couple ATP (Adenosine 5'-triphosphate) hydrolysis to transport of a tremendous variety of substrates including drugs, vitamins, peptides, polysaccharides, ions, sugars, lipids and even proteins across cellular membranes in all organisms (Higgins, 1992; Hollenstein et al., 2007).

ABC transporters have been conserved across the three kingdoms of Archaea, Eubacteria and Eukarya. Their ubiquitous distribution and primordial origin reflect the fundamental requirement of cellular homeostasis to import and concentrate essential nutrients, expel toxins acquired from the environment, and produce metabolic by-products. The report of the complete genome sequence of *E. coli* K-12 identified 80 ABC proteins, equals to 5% of the genome. In humans, ABC transporters can be grouped into seven different subfamilies (A, B, C, D, E, F, G) (Tatusov et al., 1997).

In humans, P-glycoprotein (p-gp) in the plasma membrane is probably the best studied ABC drug efflux transporter. It can transport a wide of substrates; including a number of drugs, such as the anti-cancer drugs doxorubicin and daunorubicin that are applied in therapeutic applications. MRP1, which localizes to the basolateral membrane of epithelial cells, was first identified in a cell line and shown to confer resistance to a range of anti-cancer drugs, such as anthracyclines (Cole et al., 1994; Evers et al., 1996).

ABC importers mediate the uptake of essential nutrients, whereas ABC exporters expel diverse substrates (Figure 1.13). The ABC exporter is formed by the transmembrane domains (TMD) and the nucleotide binding domains (NBD). In most cases, P-gp for example (Figure 1.13a), a single polypeptide includes the four domains, forming a TMD-NBD-TMD-NBD structure. On the other hand, the LmrA, a multidrug efflux pump, needs two polypeptides to form the functional unit.

Unlike ABC exporters, the ABC importers require the additional soluble protein substrate-binding-protein (SBP) to bind the substrate and deliver it to the translocator in the membrane. In many cases, each domain of ABC importers is present as a separate polypeptide, giving a total of five unique proteins (Figure 1.13b), such as the maltose importer MalEFGK2 and vitamin B12 importer BtuC2D2F.
The membrane-bound complex importer MalFGK2 from *E. coli*, with the help from MalE, a soluble periplasmic receptor, can mediate the uptake of the nutrient maltose from the periplasm across the membrane into the cytoplasm (Diederichs et al., 2000). Some ABC transporters in bacteria are around 35% identical to the P-gp and the region of homology extends from the nucleotide binding domains to the transmembrane domains, thus suggesting these ABC transporters not only have similar sequences but may transport similar substrates. LmrA, a multidrug efflux pump, from *L. lactis*, confers resistance to drugs and antibiotics and has been found to functionally substitute for human P-gp when expressed in lung fibroblast cells (van Veen et al., 1998). In addition, the ABC pump Sav1866 from *S. aureus* also shows significant sequence similarity to P-gp. The structure of the Sav1866-ADP complex has been determined and can be helpful in understanding the substrate recognition and transportation process (Dawson and Locher, 2006).

![Diagram of ABC transporters](image)

**Figure 1.13** The topology of different ABC transporters. TMDs are represented by squares, NBDs are represented by circles, and substrate binding proteins with ABC importers are represented by pac-men.
1.2.2.2 Human diseases associated with ABC transporter

Drugs are pumped out by ABC transporters faster than they can diffuse across membranes, so the drug level inside is kept low and is much less effective, causing problems with chemotherapeutic treatments. Besides, there are approximately 50 known ABC transporters in humans and there are currently 13 genetic diseases associated with defects in 14 of these transporters. A few human diseases related to ABC transporters are described as follows (Gottesman and Ambudkar, 2001).

Cancer

Multidrug resistance (MDR) means cells have a cross-resistance phenotype against several unrelated drugs which differ in structure and target specificities. Resistance to chemotherapy is a common clinical problem in patients with cancer, and cellular resistance to drugs is a major reason for treatment failure. The MDR phenotype was first described as P-gp mediated for cancer cell, but cancer cells can be MDR without the involvement of P-gp (Lage, 2003). The Alterations in the signal-transduction pathways can abolish the function of drugs that accelerate programmed cell death. For example, a mutated, inactive p53 that is no longer able to trigger apoptosis after the detection of DNA damage caused by drugs (Hannun, 1997; Lowe et al., 1993).

AIDS

Human immunodeficiency virus (HIV), which causes acquired immune deficiency syndrome (AIDS), has been studied for more than a decade. However, it can not be eradicated from infected patients partially due to multidrug resistance (Huisman et al., 2000). The ABC transporter P-gp is known to efflux a number of drugs that includes the human immunodeficiency virus protease inhibitor. A report indicated that overexpression of p-gp can protect cell lines from the cytotoxic effects of high concentrations of phosphatidylinositol (PI) (Choo et al., 2000). In vivo, pharmacological inhibition of the accumulation of PIs in brain and testes of mice (Choo et al., 2000) and p-gp knockout mice also demonstrated an increased antiretroviral drug saquinavir accumulation in the brain (Washington et al., 2000).
Surprisingly, growing evidence suggests that p-gp may also affect replication of enveloped viruses (Owen et al., 2005). A report showed that p-gp overexpression can block insertion of the influenza virus fusion protein into the plasma membrane, inhibiting the membrane fusion and infectivity of the influenza virus (Raviv et al., 2000).

**Cystic fibrosis**

Malfunctional ABC transporters can also cause diseases. The cystic fibrosis transmembrane conductance regulator (CFTR) is a cAMP-activated chloride channel, which creates sweat, digestive juices, and mucus, is expressed in the epithelia of the lung, intestine, pancreas and other tissues (Verkman et al., 2006). Mutations, such as the most common mutation ΔF508, in CFTR can cause cystic fibrosis (CF), a hereditary disease, where defective epithelial cell fluid transport produces chromic lung infection and deterioration of lung function, pancreatic insufficiency and male infertility.
1.3 The structure and mechanism of ATP-binding cassette transporters

1.3.1 Conserved domains and motifs

So far, each ABC transporter has been discovered to consist of four domains: two nucleotide binding domains (NBD) that serve as engines, where ATP can be hydrolyzed, coupling hydrolysis to active transport; and two transmembrane domains (TMD) that are believed to constitute the transport pathway.

**NBD in ABC transporters**

Many crystals of isolated NBDs are available, yielding detailed information about the overall structures of the nucleotide-binding sites (Davidson and Chen, 2004). Generally, the NBDs can be divided into two sub-domains: a RecA-like sub-domain and a helical sub-domain, the latter being specific for ABC transporters. The RecA-like fold is characterized by a central β-sheet flanked by α-helices and can be found in a broad subclass of topologically homologues including RecA, DNA and RNA helicases, F1-ATP synthase, DNA repair enzymes and ABC transporters (Ye et al., 2004).

Many conserved motifs, involved in ATP binding and hydrolysis or facilitating crucial interfaces in the assembled transporter (Higgins et al., 2004), can be recognized in the NBDs of ABC transporters (Figure 1.14) and will be described as follows:

*Walker A and Walker B*

The Walker A (Walker et al., 1982), also known as the P-loop, and Walker B motifs, present in many ATP-binding proteins and are not specific to ABC transporters. The highly conserved Walker A motif (G-X-X-G-X-G-K-S/T, where X represents any amino acid) is located after β-strand 3 and before the α-helix, and is involved in the coordination of the phosphates of ATP during the hydrolysis reaction. The less conserved Walker B motif (O-O-O-O-D-E, where O represents a hydrophobic amino acid) located on β-strand 7 provides a conserved aspartate that coordinates the Mg\(^{2+}\) ion in the nucleotide-binding site through H\(_2\)O. The last residue, a glutamate, in the Walker B motif arranges the nucleophilic attack on ATP via a water molecule and mutation of this residue leads to loss of ATPase activity (Moody et al., 2002; Smith et al., 2002). Zaitseva et al. suggested that residue H662 may interact with the γ-
phosphate moiety of ATP and water molecule 90, which forms hydrogen bonds to H662, E631, and via water molecule 109 to Q550, could serve as the catalytic water since a slight movement would be sufficient to put it in a proper orientation for attack of the scissile bond.

**Signature sequence**

The signature sequence, also known as the LSGGQ motif and C-loop, located at the joint point between the helical domain and the Walker B motif, binds to ATP, and is the motif that distinguishes ABC proteins from other nucleotide-binding proteins (Davidson, 2002). Mutations in the signature sequence inhibit ATPase activity (Schmees *et al.*, 1999; Bakos *et al.*, 1997) and structural studies indicated the motif contacts the nucleotide in the ATP-bound state (Smith *et al.*, 2002; Chen *et al.*, 2003; Dawson and Locher, 2006; Dawson and Locher, 2007).

**Q-loop**

The Q-loop, also known as the γ-phosphate linker, following β-strand 6, contains a glutamine that can form hydrogen bonds with the Mg$^{2+}$ and the attacking H$_2$O. Verdon and co-workers (Verdon *et al.*, 2003), on the basis of the structure of the ABC-ATPase GlcV, proposed that the role of the conserved Q-loop glutamine residue, and its interaction with Mg$^{2+}$, is to keep the two ABC subdomains in close contact to each other. For example, mutations in the conserved glutamine in the Q-loop of MalK had only marginally reduced the ATPase activity but led to defective transporters when they interated in the membrane protein complex.

**H motif**

A conserved histidine in the H motif, located before α-helix 7, is believed to serve as a linchpin in ATP hydrolysis (Zaitseva *et al.*, 2005). Chen and co-workers found that the H motif residue (His$^{192}$) not only forms a hydrogen bond with the γ-phosphate but is also involved in direct protein-protein interactions across the dimer interface in the ATP-bound MalK dimer (Chen *et al.*, 2003). Mutations of the conserved histidine in HisP (His$^{211}$) (Shyamala *et al.*, 1991), MalK (His$^{192}$) (Davidson and Sharma, 1997) and HlyB (H$^{662}$) (Zaitseva *et al.*, 2004) resulted in a substantive decrease in the steady-state ATPase activities of these proteins.
D-loop

The D-loop is located between the Walker B motif and α-helix 5 and contributes in the contact interface between the two NBDs. In the closed form of Malk, the conserved Asp$^{165}$ in the D-loop was found to form a hydrogen bond with Ser$^{38}$ in the Walker A motif of the other subunit (Chen et al., 2003).

A-loop

The A-loop, a highly conserved aromatic residue approximately 25 amino acids upstream of the Walker A motif, is essential for ATP-binding (Ambudkar et al., 2006). The structure of GlcV indicated that a phenylalanine residue is involved in the recognition of the adenine base (Verdon et al., 2003). Similarly, it was shown that a tyrosine residue interacts with the adenine ring of ATP in the HlyB-NBD structure (Zaitseva et al., 2005). Mutagenesis studies also indicated that ATP hydrolysis was abolished when Tyr$^{16}$ was mutated to cysteine in the histidine permease (Shyamala et al., 1991).

Figure 1.14  The conserved motifs in NBDs of ABC transporters. Crystal structure of the Hlyb-NBD dimer with bound ATP/Mg$^{2+}$ (Zaitseva et al., 2005). ATP (stick representation) and Mg$^{2+}$ (green spheres) are sandwiched at the interface of the two monomers.
TMD in ABC transporters

The TMDs create the channel that substrates pass through during translocation. Unlike the NBDs, the TMDs are poorly conserved. The TMDs of ABC transporters are diverse in primary sequence, length, and the number of transmembrane helices, reflecting the large diversity of substrates transported. Characteristically, ABC importers have between 10 and 20 transmembrane helices; while ABC exporters tend to form the two-times-six α-helix structure, making a total of 12 transmembrane helices per functional unit (Figure 1.15). However, this is not always the case. For example, P-gp and CFTR are functional as a single polypeptide that includes all four domains (Sheppard and Welsh, 1999; Loo and Clarke, 1995); whilst the ABC transporter MacB from *E. coli* is predicted to have a topology consisting of four transmembrane helices; which will be described further below (Kabayashi *et al.*, 2003).

![Figure 1.15 Illustration of the TMDs topology of ABC transporters. Sav1866 is shown in green, ModB is shown in yellow and BtuC, HI1471 are in red (Hollenstein *et al.*, 2007).](image)
1.3.2 Structures of ABC transporters

So far there are five complete ABC transporter structures that have been determined: the importers BtuCD, from *E. coli*, HI1470/1, from *H. influenzae* and ModB₂C₂A from *A. fulgidus*, and the exporter Sav1866, from *S. aureus* and MsbA. The four domains of the importers are on separate polypeptides and form a complex of homodimeric NBDs and TMDs (the structure of the periplasmic binding protein ModA is also shown in the ModB₂C₂A structure) whereas Sav1866 is a half transporter with one NBD and one TMD on each polypeptide (Locher *et al.*, 2002; Pinkett *et al.*, 2007; Hollenstein *et al.*, 2007; Dawson and Locher, 2006; Ward *et al.*, 2007).

*Sav1866*

*Sav1866* is a multidrug ABC transporter from *S. aureus*. When reconstituted into liposomes, Sav1866 ATPase activity can be stimulated by the anti-cancer drugs doxorubicin, vinblastine and the fluorescent dye Hoechst 33342 (Dawson and Locher, 2006), indicative of communication between the substrate binding site, in the TMD, and the cytoplasmic NBDs.

The Sav1866 structure is the first determined for an ABC exporter (Dawson and Locher, 2006). The Sav1866 transporter (Figure 1.16) is a 120 Å long, 65 Å wide and 55 Å deep homodimer and each polypeptide is composed of two domains: an N-terminal TMD (amino acid residues 1–320) and a C-terminal NBD (residues 337–578). Conserved ATP-binding and hydrolysis motifs, which are exposed at the interface, can be observed in the structure, revealing a “head-to-tail” arrangement. The electron density map indicated that the two bound ADP molecules were sandwiched at the NBD-NBD interface, in contact with the Walker A motif of one NBD and the ABC signature motif of the other.

Each Sav1866 subunit has 6 transmembrane helices, making a full transporter with 12 transmembrane helices. Bundles of transmembrane helices are divided into two wings that point away from one another towards the cell exterior, providing an outward-facing conformation in the crystal (Figure 1.16). Interestingly, each wing is composed of helices 1 and 2 from one subunit and helices 3 to 6 from the other. A large cavity was observed in the interface of the two TMDs, which is exposed to the
extracellular space. The cavity features a hydrophilic surface, lined primarily with polar and charged amino-acid side chains from helices 2 to 5, in the inner leaflet.

Figure 1.16  Sav1866 structure by ribbon representation. (a) Side view of the Sav1866 structure with subunits colored in yellow and green. (b) Stereo view rotated with respect to “(a)” by 90° around the vertical axis. (c) Close-up view of the interfaces between NBDs and TMDs of Sav1866. One subunit is colored in green ribbon representation and other is in yellow coil representation (Dawson and Locher, 2006).
The transmembrane helices are extended beyond the lipid bilayer by short extracellular loops (ECLs) and long cytoplasmic loops (ICLs), which protrude 25 Å into the cytoplasm. ICLs 1 and 2, which connected helix 2 and 3, and 4 and 5, respectively, both contain short helices, known as "coupling helices", and are orientated approximately parallel to the membrane of the shared interface formed by the TMDs (figure 1.16c). Surprisingly, coupling helix 1 contacts both subunits whereas coupling helix 2 contacts only the other, indicating that the ICLs of Sav1866 primarily contact the NBD of the opposite unit. In contrast, the TMDs of the ABC importer BtuCD contact only on NBD, resulting in a large gap at the center of the four protein domains (Locher et al., 2002). Residues around the conserved Q loops from the NBDs participate in the NBD-TMD contact interface and those residues can also be observed in the ABC importer BtuCD. In the BtuCD structure, residues around the Q loop dominate the interface between BtuC and BtuD. Side chains making specific contacts to the L loop of BtuC are located in helices H2, H3 of BtuD. Elements involved in ATP-binding and hydrolysis may be coupled to the conformation of the L loop from the membrane-spanning subunit and vice versa.

In the Sav1866, two prominent exceptions which were not observed in the BtuCD include the conserved residues Tyr391 and Glu473. The residue Glu473 contacts both ICLs and is found to be part of a previously unrecognized sequence motif (EVGERG), only conserved in ABC exporters. This newly-discovered specific sequence motif, called the X-loop, may indicate that ABC importers and exporters have diverse coupling mechanisms (Dawson and Locher, 2006).

Nucleotide-bound structures of MsbA provide another example of the outward-facing conformation of an ABC transporter (Figure 1.17). The electron-density maps of the MsbA-AMPPNP complex showed a conformation similar to the homologous Sav1866-ADP complex (Ward et al., 2007). Each monomer of MsbA has six transmembrane helices, forming a full transporter of 12 transmembrane helices. Similar to the Sav1866 structure, each wing of the MsbA full transporter is composed of helices 1 and 2 from one subunit and helices 3 to 6 from the other. In this study, the structural evidence arising from three MsbA structures, the nucleotide bound, open apo and closed apo forms, indicated that MsbA can undergo a large range of motions between the apo and nucleotide-bound states. This proposal will be described in the following sections.
Figure 1.17 The stereoview of the nucleotide-bound MsbA structure. One monomer is colored with a rainbow gradient (N to C terminus, blue to red), and the other is white. The AMP-PNP molecule is displayed as blue sticks.
The ModB$_2$C$_2$A is an archaeal molybdate ABC importer from *A. fulgidus*. This full-size transporter is composed of two NBDs (ModC) and two TMDs (ModB) that interact with a periplasmic binding protein (ModA). The structure of the complex has been determined to a resolution of 3.1 Å using X-ray crystallography (Hollenstein *et al.*, 2007) (Figure 1.18a). In the crystal, a single binding protein ModA, with bound tungstate, is attached to the external side of ModB$_2$. Each ModB subunit has 6 transmembrane helices, making a full transporter of 12 transmembrane helices. The twelve helices provide an inward-facing conformation, which is wide open to the cytoplasm. The nucleotide free ModC subunits reveal an open conformation, which contrasts with the “closed conformation” for the NBDs of Sav1866-ADP.

The ModB subunits form a large cavity, possibly representing the translocation pathway, shielded from the lipid bilayer and only accessible from the cytoplasm. The cavity opens to the cytoplasmic space, narrows down towards the external membrane boundary and is separated from the extracytoplasmic space by a closed gate below the interface with the binding protein ModA (Figure 1.18b). The gate is formed by the joining of four protein stretches (gate regions), two from each ModB subunit. Gate region 1 and 2 are located in transmembrane helices 3 and 5 respectively. In addition, the phenyl rings of the conserved Phe$^{200}$, in transmembrane helix 5, from the two ModB subunits are within van der Waals distance, implying their involvement in gating (Figure 1.18c).

An “head-to-tail” arrangement between the P loops and the ABC signature motifs was observed in the two ModC subunits. Not surprisingly, a gap was observed between the ATP-binding motifs of the nucleotide-free NBDs, indicating a so called “open conformation” of NBDs. The observation of the spacing between the Walker A motif and the ABC signature sequence is consistent with the biochemical studies of Mal ABC importers in which the solvent accessibility to the fluorescently labelled ATP binding sites is increased in the absence of ATP (Mannering *et al.*, 2001). In this study, Cys$^{40}$, in the Walker A motif of MalK, was labelled with 2-(4'-maleimidoanilino) naphthalene-6-sulfonic acid (MIANS) in order to observe the conformational changes in the Malk subunit.

The ModB-ModC interface is believed to transmit conformational changes, coupling ATP binding and hydrolysis to substrate transport. Similar to the Sav1866
The structure of the ABC transporter ModB$_2$C$_2$A. (a) The ModB subunits are in yellow and blue, the ModC subunits are in green and pink, the substrate binding protein ModA is in red, and with bound tungstate in van der Waals representation (yellow and blue spheres). The grey box indicates the possible location of the lipid bilayer on the basis of the hydrophobicity of the protein surface. (b) Topology of the ModB subunit. The conserved gated regions are colored in green and red. (c) Top view of transmembrane ModB subunits, with one subunit in yellow ribbon representation and the other in blue backbone trace (Hollenstein et al., 2007).
1.3.3 Conformational changes in NBDs

The Sav1866 and ModB2C2A structures are indicative of the “closed conformation” and “open conformation” of the NBDs respectively. The idea is drawn from comparison of the nucleotide-free-MalK and ATP-bound-MalK structures, in which it was possible to observe major conformational changes in the NBDs as ATP was bound, which has been likened to the action of a pair of tweezers (Figure 1.19). The distance between the two His\(^{89}\) on the Q-loop moved from the closed form to the semi-open or open conformations by 9 or 25 Å respectively (Chen et al., 2003; Oldham et al., 2007).

In addition to the movement of the whole NBDs, a hinge rotation of the helical subdomain relative to the RecA-like subdomain can be observed in the MalK structures. The difference can be found not only in different NBD structures but in the same protein crystallized in different crystal lattices and even between molecules in the same lattice (Chen et al., 2003). Interesting, the orientation of the helical subdomains are the same among the three ATP-bound structures: HisP, MalK and MJ0796, suggesting the helical subdomain has unique orientation in the presence of γ-phosphate. The flexibility of the helical subdomain was only observed in the NBD structures in the absence of the γ-phosphate of ATP, suggesting an important mechanistic role for this movement. When the helical subdomain is tilted away from the RecA-like subdomain, as in the nucleotide-free conformation, the two NBDs are unable to come into close enough contact to form the ATP-bound dimer. This rotation of the helical subdomain could also be coupled with ATP binding and hydrolysis and the characteristic flexibility may be important in adjusting the size of the opening of the translocation pathway for various sizes of substrates. This can also be found in full size ABC transporters by comparing the NBD structures from Sav1866 (closed form) and ModB2C2A (open form). A difference can be observed in the ModC\(_2\) dimer in the withdrawal of the helical subdomain from the interface with the opposite NBD, combined with a slight outward tilt. Also, several lines of biochemical evidence indicated that the association and dissociation of NBDs plays an important role in transport. The E171Q mutant of MJ0796 eliminated hydrolysis and stabilized the ATP-bound dimer (Smith et al., 2002). Addition of ATP enhanced disulfide-crosslinking between cysteines introduced at position 85 in the Q-loops of the MalK subunits (Hunke et al., 2000).
Figure 1.19  **Comparison of two MalK structures**, where closed form MalK with bound ATP is in yellow and semi-open form MalK without bound ATP is in blue (Chen et al., 2003).
1.3.4 Mechanisms for coupling of hydrolysis to transport

That the binding of ATP triggers the NBDs to dimerize has been described in the previous section. Two models for the transport mechanism, based on the Sav1866 and ModB$_2$C$_2$A structures, and MsbA structures, will be described below.

*A transport mechanism based on the Sav1866 and ModB$_2$C$_2$A structures*

![Figure 1.20 Conformational changes triggered by ATP. The structure of ModB$_2$C$_2$A indicates an inward-facing conformation and the Sav1866 structure indicates an outward-facing conformation. The black helices are the coupling helices from the TMDs (Hollenstein et al., 2007).](image-url)
A mechanism for ABC transporters has been proposed on the basis of the structures of the two full size ABC transporters, Sav1866 and ModB2C2A (Hollenstein et al., 2007) (Figure 1.20), where Sav1866 represents the "closed form" of the NBDs and the "outward-facing" conformation of the TMDs whilst ModB2C2A represents the "open form" of the NBDs and the "inward-facing" conformation of the TMDs. In the Sav1866 structure, the distance between the coupling helices is decreased by 10-15 Å when compared with the nucleotide-free structure for ModB2C2A. ABC importers may now accept substrates from their cognate binding proteins and ABC exporters may extrude bound drugs to the environment by presenting a cavity of low affinity to the extracellular medium (Loo and Clarke, 2005). The hydrolysis of ATP, and the subsequent release of ADP and inorganic phosphate, may return the transporter to an inward-facing conformation. ABC importers may now release their substrates into the cytoplasm, whereas ABC exporters may recruit new substrates into their high-affinity binding sites.

A transport mechanism based on the structures of three conformations of MsbA

Ward and co-workers (Ward et al., 2007) provided structural evidence, arising from different conformational states for MsbA, showing that MsbA can undergo a large range of motions between the apo and nucleotide-bound states (Figure 1.22a).

The conformational changes between difference MsbA structures are most noticeable when monomers from each state are aligned using transmembrane helices 1, 2, 3, and 6 (Figure 1.21). Comparison of the apo structures reveals that TM4/TM5/IH2 can pivot around 30° about a hinge formed by EL2/EL3 in a rigid-body motion with nearly conserved NBD alignment. In apo structures, this hinge creates an inward-facing conformation between TM3/TM6 and TM4/TM5 (Figure 1.21). Interestingly, although the NBDs were closed in the apo state, an ATP sandwich structure was not observed (Figure 1.22). In addition, the P-loops of opposing monomers are positioned next to one another (Figure 1.22c).

The transition from the closed apo conformation to the nucleotide-bound conformation can be accounted for by two motions. Firstly, there is a 10° pivot about the hinge, EL2/EL3 (Figure 1.21), bringing the NBDs closer together. Secondly, the NBDs form an ATP sandwich by tilting the TM4/TM5 pair around 20° out of plane
(relative to the first motion) about that pivot point centered along its length. Since TM3/TM6 are connected to TM4/TM5 via EL2 and EL3, the formation of the NBD sandwich pulls TM3/TM away from TM1/TM2 (Figure 1.22ab). An outward opening is created between TM3/TM6 and TM1/TM2, which is different from the inward opening created between TM3/TM6 and TM4/TM5. The conformational changes (Figure 1.21 and 1.22) in the NBDs cause a rearrangement of the packing of TMDs, switching access to the internal chamber from the inner to the outer leaflet of the bilayer. The large inward-facing opening may serve an important role in transport because the open and closed apo structures can accommodate substrates from the inner leaflet or the cytoplasm. Substrate binding to the inward-facing conformation may lead to the closure the TMDs and then rearrangement of the NBD, allowing the formation of an ATP sandwich conformation.

![Figure 1.21 Comparison of the MsbA monomers.](Image) The stereoview of the conformational changes within the MsbA monomer illustrates the mobility of the TM4/TM5/IH2. Monomers are aligned on transmembrane helices 1, 2, 3 and 6. Two apo open MsbA monomers that differ in the relative positioning of TM4/TM5/IH2 are shown in blue and cyan. The apo closed MsbA is colored in green and the nucleotide-bound MsbA is in yellow (Ward et al., 2007).
Figure 1.22  Conformational changes in MsbA. (a) Conformational changes in the open apo, closed apo and nucleotide-bound MsbA. (b) Model of the conformations changes. The arrows illustrate the motions required to go to the next state. (c) Top view of the NBDs. One monomer is colored in white and the other in gray (Ward et al., 2007).
1.4 Tripartite multidrug efflux pumps in gram-negative bacteria

Antibiotic resistance due to multidrug transporters is a growing problem in the world; to combat them, more research into the transport mechanism and structures of efflux pumps is necessary. Gram-negative bacteria possess tripartite complex pumps which are composed of an inner membrane protein, a periplasmic protein and an outer membrane protein, which work together to confer multidrug resistance. Recent studies have brought a better understanding of tripartite drug efflux pumps, particularly, AcrA/AcrB/TolC in E. coli and MexA/MexB/OprM in P. aeruginosa. The determination of the structures of the three components, AcrA, AcrB and TolC, provide us with clues to the function and mechanism of these tripartite efflux complexes (Koronakis et al., 2000; Mikolosko et al., 2006; Murakami et al., 2002).

TolC structure

TolC, a multifunctional outer membrane protein, is involved in type I secretion of toxins and proteases and plays an important role in antibiotic efflux. It can export haemolysin by a substrate-induced transient interaction with the ABC transporter HlyB and the membrane fusion protein HlyD (Thanabalu et al., 1998). On the other hand, the in vivo disulfide cross-linking between AcrB and TolC may suggest that the AcrA/AcrB/TolC efflux complex appears to be independent for the drug substrates and is assembled constitutively (Touze et al., 2004).

The TolC structure has been solved by X-ray crystallography (Koronakis et al., 2000), indicating that three TolC protomers assemble to form a 140 Å long channel spanning the outer membrane and periplasmic space (Figure 1.23a). The TolC molecule is composed of three domains: a β-barrel domain, an α-helical domain, and a mixed α/β-domain which forms a strap around the middle section of the helical barrel (Figure 1.23b). The α-helical barrel domain of trimeric TolC forms a cylinder of 100 Å in length and 35 Å internal diameter, which makes a mostly solvent-filled inner volume of roughly 43,000 Å³, one of the largest known in a protein structure. The upper end of the TolC structure is open, providing wide solvent access, while the lower end of the structure is closed by dense packing of 12 tunnel-forming α helices (Figure 1.23a).
Figure 1.23  The structure of outer membrane protein TolC. (a) The Side view of the TolC by a ribbon representation where three protomers are colored in red, yellow and blue. (b) TolC monomer by a ribbon representation where the β-barrel domain is in yellow, the α-helix domain is in green and the equatorial domain is in red (Koronakis et al., 2000). (c) A scheme of the mutations to stabilize channel opening of TolC. (d) The bottom view of the open and closed TolC structures (Bavro et al., 2008).
TolC must undergo a conformational change to open the entrance for translocation since it is too small for the substrates to enter the channel (Figure 1.23c). The disruption of the hydrogen bond between amino acid residue D\textsuperscript{153} and Y\textsuperscript{362} and the salt bridge between the D\textsuperscript{153} and R\textsuperscript{367} of TolC increase the conductance in lipid bilayers, enlarging the low end entrance (Figure 1.23c) (Andersen et al., 2002). The translocation of polypeptide substrate is prohibited by introducing disulphide bonds to constrain the entrance coiled coils in the close state (Eswaran et al., 2003), further supporting the proposal that the transition to the open state TolC is accomplished by the re-arrangements of the entrance helices.

Bavro and co-workers determined two TolC structures in its partially open state (Bavro et al., 2008). Previous report indicated that the disruption of the hydrogen bond between amino acid residue D\textsuperscript{153} and Y\textsuperscript{362} and the salt bridge between the D\textsuperscript{153} and R\textsuperscript{367} of TolC can enlarge the TolC periplasmic entrance (Andersen et al., 2002). The structure of the double mutant (Y362F and R367E) of TolC was solved in two crystallin forms at 3.2 and 3.3 Å resolution (Figure 1.23d). The crystal structures of the double mutant showed that three of the six pairs of coiled coils move radially outward from the central in comparison of the closed state form, repacking the helices and partially opening the channel (Figure 1.23d). It was found that the greatest conformational changes occur in the H7/H8 helices; the H3/H4 helices are comparatively static and are like a stator around which the H7/H8 swings. Although the movements of the helices in the double mutant TolC structure enable the opening for passage of molecules past the distal tip of TolC, the second aperture is still closed, suggesting that a further conformational change is required for greater channel opening.
**AcrA structure**

The membrane fusion protein AcrA is a component of the tripartite multidrug efflux pump AcrA/AcrB/TolC in *E. coli*. AcrA has been shown to interact with both AcrB and TolC, which suggests that AcrA serves as a scaffold in the tripartite efflux pump (Touze et al., 2004; Tikhonova et al., 2004). In addition, AcrA may also function in the opening of the periplasmic end of TolC channel (Bavro et al., 2008).

The crystal structure of a 28 kDa core of AcrA (residues 45-312) was shown to be comprised of an α-helical hairpin domain, a lipoyl domain and a β-barrel domain (Figure 1.24a). Indeed, it is closely related to the structure of the membrane fusion protein MexA from *P. aeruginosa* (Higgins et al., 2004; Mikolosko et al., 2006). Interestingly, in the lattice in the AcrA crystal, each α-helical hairpin of the four monomers has a different orientation (Figure 1.24b), which is not observed in the MexA structure. The difference in orientation of the α-helical hairpin of AcrA stems from an effective hinge, located between the α-helical hairpin and lipoyl domain, composed of residues 99-106 in α-helix 1 and residues 169-173 in α-helix 2.

In contrast to AcrB and TolC, which form trimers, the oligomeric state of AcrA is uncertain. In the crystals, AcrA are seen to form a dimer through extensive contacts involving the α-helical hairpins as well as adjoining lipoyl and β-barrel domains. On the other hand, AcrA was monomeric in solution and was found to be trimeric when crosslinked *in vivo* by DSG, which has a short spacer arm (7.7 Å) (Eswaran et al., 2004; Zgurskaya and Nikaido, 2000).
Figure 1.24  **Structure of AcrA.** (a) Ribbon representation of the crystal structure of AcrA (residues 45-312) monomer, with the α-helical hairpin domain in red, lipoyl domain in green and β-barrel domain in cyan. (b) Four conformations of AcrA in the crystal show the conformational flexibility of the α-helical hairpin (Mikolosko et al., 2006).
The AcrB/AcrA/TolC complex pumps

TolC is a homotrimeric outer membrane protein that comprises an outer-membrane β-barrel, prolonged by a 100 Å α-helical transperiplasmic tunnel. The TolC extracellular exit channel is constitutively open, with a pore diameter of 35 Å, but the periplasmic domain channel entrance was restrained by coiled-coil α-helices. Andersen and co-workers proposed an iris-like transition to the open state based on the realignment of the inner coils on the outer ones (Anderson et al., 2002). The inner membrane protein AcrB, known as being trimeric, consists of a transmembrane domain at its bottom and a huge periplasmic domain on top. The top docking domain forms a funnel, with about a 30 Å internal diameter that is consistent with that of the TolC conduit. The sum of the periplasmic length of AcrB and TolC is about 170 Å, which is enough to cross the periplasmic space (Koronakis et al., 2000; Murakami et al., 2002). Moreover, the length and topology of the periplasmic domains of both TolC and AcrB suggests that the two interact directly (Tamura et al., 2005). Substrates might access the AcrB central cavity from the periplasm through the AcrB lateral vestibules for transport to the cell exterior.

AcrA and TolC in E. coli have been found to form a stable intermembrane multidrug efflux complex with or without the substrate (Eswaran et al., 2004; Tikhonova and Zgurskaya, 2004). The AcrA-AcrB complex was identified upon purification from cell membranes after stabilization by in vivo crosslinking in the absence of TolC (Zgurskaya and Nikaido, 2000). Moreover, AcrA was found to be crosslinked with AcrB and TolC individually in vivo, and microcalorimetry studies demonstrated energetically favourable interactions of AcrA with both purified AcrB and TolC (Touze et al., 2004). AcrB can also be isolated as a complex with TolC by in vivo crosslinking; however, isothermal titration calorimetry did not detect an interaction between the purified AcrB and TolC proteins, suggesting AcrA may serve a scaffold to stabilize the complex in vivo (Eswaran et al., 2004).

The AcrB and TolC crystals structures indicated that the 70-Å- and 100-Å-long periplasmic domains of AcrB and TolC would come into close proximity (Murakami et al., 2002; Koronakis et al., 2000). The interaction between the apex of AcrB and the bottom of TolC was then revealed by in vivo disulfide crosslinking between AcrB and TolC without a crosslinking agent (Tamura et al., 2005). Domain swapping and crosslinking experiments suggest that the C-terminal region of AcrA is required for the
interaction with AcrB (Elkins and Nikaido, 2003; Touzé et al., 2004). The crystal structures of AcrA and TolC (Mikolosko et al., 2006; Koronakis et al., 2000; Federici et al., 2004) suggested that the AcrA hairpin and the TolC lower α-helical domain could be involved in complex assembly; a proposal that has been proven by hairpin domain swapping (Stegmeier et al., 2006; Eda et al., 2006).

Although AcrA, AcrB and TolC are proven to form a stable tripartite efflux pump in E. coli, some critical points about how the complex works are still uncertain. Touzé (Touzé et al., 2004) suggested the complex is assembled constantly since the assembly occurred independently of the presence of the substrate. Besides, it is estimated that around 500 toxic ethidium molecules are exported every second by each MexA/MexB/OprM pump in P. aeruginosa (Narita et al., 2003). On the other hand, Tamura suggested that there is not enough TolC, a multifunctional protein, to form permanent complexes with AcrB and AcrA, although the assembly is substrate independent (Tamura et al., 2005). Instead, he proposed two possibilities: (1) transient complex formation is simply substrate independent; or (2) an unknown intrinsic substrate which facilitates complex formation has not been discovered. As discussed in the previous section, we still do not know how TolC undergoes the conformational change from the close to open state during substrate transportation. Since there is no obvious conformational change in the TolC docking domain of AcrB during the translocation cycle, Murakami (Murakami et al., 2006) proposed the conformational change caused by proton translocation and drug binding may be transmitted, via AcrA, to TolC, thus causing opening of the TolC periplasmic channel. In this case, the AcrA α-helical hairpin may be involved in the opening TolC, via its conformational flexibility (Mikolosko et al., 2006).

Unlike trimeric AcrB and TolC, our knowledge of how many AcrA fit into the tripartite pump remains uncertain. A complex model, based on homology modelling and molecular docking, was proposed (Fernandez-Recio et al., 2004). In this model the lipoyl domain of AcrA docks into the AcrB clefts, whilst three AcrA fit simultaneously into the interprotomer grooves of AcrB and the open state of TolC, so that the predicted protomer stoichiometry is 1:1:1 for the MFP: IMP: OMP components of the tripartite pump (Figure 1.25). In that case, the conformational change caused by proton translocation and drug binding may be transmitted through AcrA via a mutual pushing and pulling movement at the cleft (Murakami et al., 2006), consistent with a recent
report showing the cleft flexibility of AcrB is essential for the pump function (Takasuka and Nikaido, 2007). A recent study also suggested that there is only a single binding site on the TolC protomer (Lobedanz et al., 2007); however, by using in vivo crosslinking to map the extent of intermolecular contacts, it showed that AcrA contacted the intramolecular groove between the inner and outer entrance coils of TolC H7/H8/H3 which is different from the contacts with the groove by helices H3/H4/H7 in the previous model.

Figure 1.25  Two sides of view of the assembled tripartite efflux pump model consisted of AcrA (red), AcrB (green) and TolC (blue). (a) Side view of the complex. (b) Top view of the complex (Fernandez-Recio et al., 2004).
Another model of the tripartite complex predicted a stoichiometry of 3:1:1 for the MFP: IMP: OMP components of the tripartite pump (Figure 1.26) (Higgins et al., 2004). The modelled ring of nine MexA (AcrA homologue) is sufficiently large to form a sheath around the open state TolC and AcrB, stabilizing the complex and making a conduit that spans the entire periplasm. This sheath model may be crucial for MF transporters, without a periplasmic domain, as a scaffold to interact with outer membrane proteins. Isothermal titration calorimetry data indicated the AcrA-AcrB and AcrA-TolC bindings are concentration dependent (Touze et al., 2004); moreover, it suggested that there would be enough space on each protomer TolC to accommodate up to three membrane fusion proteins (Mikolosko et al., 2006; Higgins et al., 2004).

Figure 1.26  The model of the tripartite efflux pump. (a) Two perpendicular views of the nine MexA molecules forming like a ring structure where the α-helices hairpin are in green and the β-strands of the lipoyl domains are in blue. (b) The side view of the complex consisted of AcrA (blue), AcrB (white) and TolC (green) (Higgins et al., 2004).
1.5 What is known about the MacA/MacB/TolC tripartite complex pump

The ABC transporter MacB was first found to confer MacA and TolC dependent resistance to macrolides in *E. coli* cells (Kobayashi *et al.*, 2001). By drug susceptibility test, Kobayashi and co-workers established that *E. coli* strain Kam3 cells (*ΔacrAB*), harboring a plasmid encoding MacAB had an obvious enhancement in their resistance to macrolides, such as erythromycin, clarithromycin, oleandomycin and azithromycin. Furthermore, Kam3 cells transformed with a plasmid encoding MacAB were found to take up less $[^{14}C]$ erythromycin than ones without MacAB, suggesting active efflux of erythromycin out of the cells. A homologue of the MacA-MacB ABC transporter of *E. coli* was identified in *N. gonorrhoeae* (Rouquette-Loughlin *et al.*, 2005); furthermore, Rouquette and co-workers indicated that *E. coli* cells JMZ120 (*ΔacrAB*) harboring a plasmid encoding MacAB have higher resistance to erythromycin and azithromycin.

It is predicated, by secondary structure analysis, that MacB putatively has four transmembrane helices, and there is a large periplasmic loop between the transmembrane helix 1 and 2; the NBD domain is located on the N terminal side of the polypeptide and the TMD domain is on the C terminal side. The MacB topology was determined by systematically replacing residues with cysteines and testing for the $[^{14}C]$ NEM binding (Figure 1.27) (Kobayashi *et al.*, 2003). MacA share a sequence identity of 21.3%, and similarity of 38.7% to its homologous protein *E. coli* AcrA, which comprised of an α-helical hairpin domain, a lipoyl domain and a β-barrel domain in its truncated form.

During the course of our study, another revealed that co-reconstitution of MacA and MacB into proteoliposomes increased the steady-state ATPase activity of MacB, suggesting that the MFP as not only serves a scaffold, for assembly of the tripartite complex, but also has a functional role (Tikhonova *et al.*, 2007). We have established that the periplasmic β-strand domain of MacA interacts with the periplasmic domain of MacB to modulate the ATPase activity of its cytoplasmic NBD. Our data suggests that this interaction promotes and stabilizes a conformation of MacB in which the substrate-binding site is outward-facing. This is concomitant with an increase in the resistance of cells to erythromycin that indicates that antibiotic
extrusion is stimulated. This suggests that MacA has a direct role in controlling the pump action to facilitate the transfer of substrates across the inner-membrane.

**Figure 1.27** Topology of ABC transporter MacB from E. coli. Nucleotide binding motifs are boxed in pink squares and the putative transmembrane helices are boxed in yellow squares. Cys$^{56}$, replaced by Ala, and the other two intrinsic cysteins are shown in bold letters. NEM labelings which was completely or largely blocked, and not affected by AMS were shown in blue ovals and red ovals, respectively. [$^{14}$C] NEM which was not reacted regardless of pretreatment or not with AMS was shown in open circles (Kobayashi *et al.*, 2003).
Chapter two: Material and methods

2.1 Web and computer resources

The nucleotide sequences of *E. coli* macB, macA and tolC were obtained from the National Center for Biotechnology Information (NCBI, www.ncbi.nlm.nih.gov). The Vector NTI Advance 10 (Invitrogen) was used for the design and analysis of oligonucleotides, and for the alignments of DNA and protein sequences for comparison. Protein secondary structures were predicted by Jpred from the University of Dundee (www.compbio.dundee.ac.uk/~www-ipred/). The Dichroweb at Birbeck College (www.cryst.bbk.ac.uk/cdweb/html/home.html) was used as an online Circular Dichroism (CD) data analysis program. The Research Collaboratory for Structural Bioinformatics protein data bank (RCSB, www.rcsb.org/pdb/home/home.do) was used as a source for protein structures.

2.2 Media and antibiotics

Luria-Bertani (LB) medium was used in molecular cloning. *E. coli* cells harboring plasmid(s) were grown in 2xYT medium at 37°C. The antibiotics carbenicillin (100 µg/ml), kanamycin (30 µg/ml) and inducing agent isopropyl-β-D-thiogalactopyranoside (IPTG) were added as required. Reagents were of analytical molecular biology grade and sourced from Difco, Melford and Sigma unless otherwise stated.

**Luria-Bertani (LB) Medium**

NaCl 10g
Trypton 10g
Yeast extract 5g
dH2O up to 1L
Mueller-Hinton Medium

Mueller-Hinton powder 23g
dH₂O up to 1L

SOC Medium

NaCl 0.585g
KCl 0.1865g
Trypton 20g
Yeast extract 5g
dH₂O up to 970ml

The above were autoclaved and the following filter sterilised components were added.

1M MgCl₂ 10ml
1M MgSO₄ 10ml
2M Glucose 10ml

2x Yeast Tryptone (2x YT) Medium

NaCl 5g
Tryptone 16g
Yeast extract 10g
dH₂O up to 1L

2.3 Bacteria strains, media and vectors

The bacterial strains and plasmids used in this study are listed in Table 2.1 and 2.2. *E. coli* strain Novablue(DE3) was used for blue and white screening of constructs. *E. coli* strain C43(DE3) was used for protein expression, and Kam3(DE3) was used for drug resistance tests.

Plasmid pGEM-T Easy vector (Promega) was used for blue and white screening of transformants in molecular cloning, and plasmids pET28a(+), pET21d(+) and pET21a (Novagen) were used as cloning and expression vectors for *macA*, *macB* and *tolC* individually. pETDuet™-1 was used as cloning vector for *macA*, *macB* and *tolC* for growth curve measurements.
Table 2.1  *E. coli* Strains.

<table>
<thead>
<tr>
<th>Strains</th>
<th>Genotype</th>
<th>Reference</th>
</tr>
</thead>
</table>
| NovaBlue(DE3) | * endA1  hsdR17  (rk12·mk12·)  supE44  thi-1  
|            | * recA1  gyrA96  relA1  lac  [F proA*B*  
|            | * lacI9ZΔM15::Tn10 (TetR*)                                                 | Novagen                        |
| C43(DE3)   | * E. coli B F  omp  had  S  (rB~ mB~)  dcm*  Tet'  
|            | * gal EndA The{argU ileY leuW Cam'}                                         | Miroux and Walker, 1996        |
| Kam3(DE3)  | * ΔacrAB                                                                  | Morita *et al.*, 1998          |

Table 2.2  Plasmids.

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Description</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>pGEM-T Easy</td>
<td>Cloning vector compatible with complementation, Ap&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Promega</td>
</tr>
<tr>
<td>pET21a(+)</td>
<td>T7 promoter expression vector, Ap&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Novagen</td>
</tr>
<tr>
<td>pET21d(+)</td>
<td>T7 promoter expression vector, Ap&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Novagen</td>
</tr>
<tr>
<td>pET28a(+)</td>
<td>T7 promoter expression vector, Km&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Novagen</td>
</tr>
<tr>
<td>pET45b(+)</td>
<td>T7 promoter expression vector, Ap&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Novagen</td>
</tr>
<tr>
<td>pETDuet™-1</td>
<td>T7 promoter expression vector with two cloning site, Ap&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Novagen</td>
</tr>
<tr>
<td>pBAD/gIII</td>
<td>pBR322-derived expression vectors for secreted recombinant protein expression</td>
<td>Invitrogen</td>
</tr>
</tbody>
</table>
2.4 Genetic manipulation

2.4.1 Polymerase chain reaction

Polymerase chain reaction (PCR) was achieved by a Qiagen hotstar taq DNA polymerase kit. All components were mixed gently in a PCR tube (Sarstedt) and then the mixture was divided equally into two tubes (each tube contains 50 µl), which were then transferred to a PCR machine. The reagent recipe used for PCR reactions is as follows.

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dH₂O</td>
<td>84.5</td>
</tr>
<tr>
<td>10X Buffer</td>
<td>10</td>
</tr>
<tr>
<td>dNTPs</td>
<td>2</td>
</tr>
<tr>
<td>Primer Forward</td>
<td>1</td>
</tr>
<tr>
<td>Primer Reverse</td>
<td>1</td>
</tr>
<tr>
<td>Hotstar taq</td>
<td>0.5</td>
</tr>
<tr>
<td>DNA template</td>
<td>1</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>100</strong></td>
</tr>
</tbody>
</table>

Primers were synthesised by Invitrogen Custom Primers, on a 50 nmol scale, and desalted. Stock solutions were prepared to a final concentration of 50 pmol/µl and stored at -20°C. Oligonucleotide primers used in this study are listed in Table 2.3.
Table 2.3 Oligonucleotide primers. Oligonucleotide names that end with ‘F’ and ‘R’ represent primer pairs. **Bold** sequences indicate restriction endonuclease sites. *Italicised* sequences indicate additional nucleotides inserted to keep target sequence in-frame with the start codon. **Underlined** sequences indicate stop codons incorporated to prevent addition of vector sequences into the transcription. Dotted underlined indicate Shine-Dalgarno (SD) sequences (Shine and Dalgarno, 1974). Shaded sequences indicate part of the leading sequence on pBAD/glll which permits secretion of recombinant protein into the periplasmic space (Rapoza and Webster, 1993; Chen et al., 1994)

<table>
<thead>
<tr>
<th>Primers</th>
<th>Nucleotide sequence (5' to 3')</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ndel-macB Forward</td>
<td><strong>CAT ATG</strong> ACG CCT TTG CTC GAA TTA AAG GAT</td>
<td>Chapter 3</td>
</tr>
<tr>
<td>Sacl-stop-macB Reverse</td>
<td><strong>GAG CTC</strong> TCA CTC TCG TGC CAG AGC ATC TAC</td>
<td>Chapter 3</td>
</tr>
<tr>
<td>Ncol-macA Forward</td>
<td><strong>CCA TGG</strong> GA GCA TAT GGA GTA TTC AGA AAA TTT</td>
<td>Chapter 4</td>
</tr>
<tr>
<td>Ncol-FD20-macA F</td>
<td><strong>CCA TGG</strong> GA GGA CTG ATT ACG TTA TGG AGA ATT</td>
<td>Chapter 4</td>
</tr>
<tr>
<td>Xhol-macA Reverse</td>
<td><strong>CTC GAG</strong> TTG TGC AGC TCC TGG TTT GGC CTC</td>
<td>Chapter 4</td>
</tr>
<tr>
<td>Ndel-tolC Forward</td>
<td><strong>CAT ATG</strong> AAG AAA TTG CTC CCC ATT CTT ATC</td>
<td>Chapter 4</td>
</tr>
<tr>
<td>Xhol-RD43tolC Reverse</td>
<td><strong>CTC GAG</strong> TTC CGG ATT AGT GGA AAC CGG TTT</td>
<td>Chapter 4</td>
</tr>
<tr>
<td>Ncol-TolC Forward</td>
<td><strong>CCA TGG</strong> GA AAG AAA TTG CTC CCC ATT CTT ATC</td>
<td>Chapter 5</td>
</tr>
<tr>
<td>EcoRI-MacB Forward</td>
<td><strong>GAA TTC</strong> G ACG CCT TTG CTC GAA TTA AAG GAT</td>
<td>Chapter 5</td>
</tr>
<tr>
<td>Sal1-stop-MacB Reverse</td>
<td><strong>GTC GAC</strong> TCA CTC TCG TGC CAG AGC ATC TAC</td>
<td>Chapter 5</td>
</tr>
<tr>
<td>Sal1-SD-macA Forward</td>
<td><strong>GTC GAC</strong> GAG GAA TAA TAA ATG GCA TAT GGA GTA TTC AGA AAA</td>
<td>Chapter 5</td>
</tr>
<tr>
<td>Sall-SD-glll Seqence F</td>
<td><strong>GTC GAC</strong> GAG GAA TAA TAA ATG AAA AAA CTG CTG TTC GCG ATT</td>
<td>Chapter 5</td>
</tr>
<tr>
<td>Not1-stop-T7tag-macA Reverse</td>
<td><strong>GCGG CCGC</strong> TCA ACC CAT TTG CTG TCC ACC AGT CAT GCT AGC CAT TTG TGC AGC TCC TGG TTT GGC CTC ACC AAT CAC CAC TTC ATC GC</td>
<td>Chapter 5</td>
</tr>
</tbody>
</table>
2.4.2 DNA preparation and electrophoresis

DNA purification was accomplished by a QIAGEN QIAquick gel extraction kit and a Promega Wizard<sup>®</sup> SV 96 Plasmid DNA Purification kit. DNA was then visualized on a 1% agarose gel containing adequate ethidium bromide (Sigma). A 1kb DNA ladder (Invitrogen) allowed estimation of DNA fragment size. The Components of 6X DNA loading buffer and 10X TAE (Tris-acetate-EDTA) agarose gel running buffer are listed as below.

6X DNA loading buffer

<table>
<thead>
<tr>
<th>Component</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bromophenol Blue</td>
<td>0.25%</td>
</tr>
<tr>
<td>Glycerol</td>
<td>30%</td>
</tr>
</tbody>
</table>

10X TAE buffer

<table>
<thead>
<tr>
<th>Component</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris Base</td>
<td>48.4g</td>
</tr>
<tr>
<td>Glacial acetic acid</td>
<td>10.9g</td>
</tr>
<tr>
<td>EDTA</td>
<td>2.92g</td>
</tr>
<tr>
<td>dH&lt;sub&gt;2&lt;/sub&gt;O</td>
<td>up to 1L</td>
</tr>
</tbody>
</table>

2.4.3 Cloning

2.4.3.1 Cloning into pGEM-T easy vectors

The pGEM-T easy vector system (Promega) was adopted for the cloning of PCR products. The pGEM-T Easy vector utilises single 3' thymidine overhangs on either side of the multiple cloning site to clone PCR products possessing complimentary 3' adenosine overhang generated by Taq DNA polymerase. The pGEM-T easy vector also contains T7 and SP6 RNA polymerase promoters flanking a multiple cloning region within the coding region for the α-peptide of β-galactosidase. The successful insertion of a target gene inactivated β-galactosidase, which could then be identified as white colonies on LB plates containing 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-Gal) and IPTG.
After cleanup, the amplified DNA was mixed with pGEM-T easy vector and T4 DNA ligase, and then left overnight at 4°C for ligation. A detailed recipe for the ligation reaction is as follows.

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (μl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2x Ligase buffer</td>
<td>5</td>
</tr>
<tr>
<td>PCR product</td>
<td>3</td>
</tr>
<tr>
<td>pGEM-T easy vector</td>
<td>1</td>
</tr>
<tr>
<td>T4 DNA ligase</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>10</td>
</tr>
</tbody>
</table>

The plasmid was then transformed (see 2.4.4 for details) into *E. coli* Novablue(DE3) for the screening of transformants. Novablue cells harboring plasmids were stored with 20% glycerol at -80°C until used.

2.4.3.2 Cohesive end DNA cloning

Restriction digestion was performed using the relevant restriction enzymes (Promega) in order to generate insert sequences from pGEM-T easy plasmids and linear overexpression vectors. All reagents were added in order as below and incubated for 3 to 5 hours at 37°C as needed.

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (μl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dH₂O</td>
<td>35.2</td>
</tr>
<tr>
<td>10X Buffer</td>
<td>8</td>
</tr>
<tr>
<td>100X BSA</td>
<td>0.8</td>
</tr>
<tr>
<td>Enzyme I</td>
<td>3</td>
</tr>
<tr>
<td>Enzyme II</td>
<td>3</td>
</tr>
<tr>
<td>Plasmid</td>
<td>30</td>
</tr>
<tr>
<td>Total</td>
<td>80</td>
</tr>
</tbody>
</table>
Ligations were carried out using the desired linear insert and vector. T4 DNA ligase HC (Invitrogen) catalyses the ligation of the complimentary ends of DNA fragment and linear vector, generated from digestion with the same restriction enzymes. The insert to vector molar ratio can have a significant effect on the outcome of a ligation and subsequent transformation step. Therefore, insert: vector molar ratios between 2 and 6 were utilized in this study. All components were added in the order listed as follows.

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (μl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5x Ligase buffer</td>
<td>4</td>
</tr>
<tr>
<td>Insert DNA</td>
<td>variable</td>
</tr>
<tr>
<td>Vector DNA</td>
<td>variable</td>
</tr>
<tr>
<td>T4 DNA ligase</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>10</td>
</tr>
</tbody>
</table>

The reaction mixture was incubated at 16°C for 90 min and then at 4°C overnight.

2.4.4 Transformation into *E. coli*

For transformation with DNA vectors, *E. coli* competent cells were prepared according to the procedure below. A chemical method for the induction of competence in *E. coli* strains is used for cloning.

1 ml of an overnight culture of the desired bacterial strain was inoculated into 100 ml of media and when the cells had reached an OD\textsubscript{600} of 0.5, they were cooled on ice for 5 min. The cells were then harvested by centrifuge at 5,000 xg for 5 min at 4°C and the supernatant was discarded. The cell pellet was resuspended in ice-cold TFB1 buffer containing 100 mM rubidium chloride, 50 mM manganese chloride, 30 mM potassium acetate, 10 mM calcium chloride, 15% (v/v) glycerol, pH 5.8, and kept on ice for 90 min. The cells were harvested by centrifuge at 4°C and the supernatant was discarded. The cell pellet was resuspended in 4 ml of ice-cold TFB2 buffer containing 10 mM MOPS, 10 mM rubidium chloride, 75 mM calcium chloride, 15% (v/v) glycerol, pH 6.8, and kept on ice for another 45 min. As described above, cells were
harvested, resuspended one more time in 4 ml ice-cold TFB2. Aliquots of 50 μL were flash frozen in a dry ice/ethanol bath and stored at -80°C.

Another method was applied to make *E. coli* strains, for use in protein expression, competent by using MgCl₂ and CaCl₂. The desired bacterial strain was grown in 5 ml LB media at 37°C for 5 hours and then placed on ice for 5 min before harvesting by centrifuge at 4°C. The cell pellet was resuspended in 1 ml ice-cold 0.1 M MgCl₂ and harvested by centrifugation at 5,000 xg for 6 min. As described above, the cell pellet was resuspended in 1 ml ice-cold 0.1 M CaCl₂ and harvested by centrifugation. Cells were then carefully resuspended in 200 μl ice-cold 0.1 M CaCl₂, and kept on ice for transformation.

Transformation was performed as follows. Competent cells were kept on ice until thawed. After adding 5 μl of plasmid, the competent cells were then put back on ice for a further 30 min. The cells were treated with a 42°C heat shock for 30~90 sec and placed on ice for another 1~5 min. LB plates, with relevant antibiotics, were used for plating out of cells and incubated at 37°C overnight. Cells harboring the desired plasmids were stored at -80°C until used.

### 2.4.5 λDE3 Lysogenisation of bacterial strains

A λDE3 Lysogenization Kit (Novagen) was applied for the integration of λDE3 prophage into an *E. coli* host cell chromosome, so that the lysogenized host could be used to express heterologous DNA from T7 expression vectors. Host cells were grown in LB media supplemented with 0.2% maltose and 10 mM MgSO₄, at 37°C to an OD₆₀₀ of 0.5, and 1~10 μl host cells were taken to mix with 10⁸ of pfu of λDE3, 10⁸ pfu of helper phage and 10⁸ pfu of selection phage. The mixture was incubated at 37°C for 20 min to allow the phage to adsorb to the host and spread on a LB plate. The plate was then incubated at 37°C overnight for cells to grow.

The presence of the λDE3 prophage in host cells was verified with a T7 phage deletion mutant, T7 tester phage. The T7 tester phage can make large halo-like plaques on authentic λDE3 lysogens in the presence of IPTG, while smaller plaques appear in the absence of inducer. In duplicate tubes, 100 μl of infected host cells, at an OD₆₀₀ of 0.5, were added with 100 μl of T7 tester phage (1 × 10³ pfu/ml). The mixtures were incubated at room temperature for 10 min to allow phage to adsorb to host and added
with 3 ml of molten agarose (no warmer than 47°C). The contents of one duplicate were poured out onto a LB plate and other one onto a LB plate with 0.4 mM IPTG. After 4 hours of incubation at 37°C the plates were ready to check.

2.5 Protein overexpression and purification

2L flasks of 2xYT media containing relevant antibiotics were inoculated with 8 ml of pre-incubated cells harboring plasmids and incubated at 37°C. 0.2~0.5 mM IPTG was added when the cells density reached an OD_{600} of 0.5. The temperature was adjusted to 25°C and protein overexpression was allowed to proceed overnight.

Cells were harvested by centrifugation at 4°C and resuspended in buffer containing 20 mM Tris-HCl pH 7.5, 500 mM, 10% (v/v) glycerol. Cell paste was added with a tablet of complete protease inhibitor (Roche) and 1000 units of DNAse I (Sigma) before going through a French press twice for cell lysis. Cell debris was removed by centrifugation at 24,000 x g (JA-20 rotor, Beckman), and cell membranes were isolated by ultracentrifugation at 168,000 x g (Type 50.2 Ti rotor, Beckman). Membranes pellets were resuspended in buffer containing 20 mM Tris-HCl pH 7.5, 20% (v/v) glycerol and stored at -80°C until needed.

2.5.1 Immobilized metal affinity chromatography

Protein-His\textsubscript{6} was solublized from cell membranes with Triton X-100 (5% [v/v]) (Promega) or n-dodecyl-β-maltopyranoside (DDM) (1% [w/v]) (Melford) and loaded into a 1 ml Ni\textsuperscript{2+} or Cu\textsuperscript{2+} charged HiTrap chelating column (GE Healthcare) by a peristaltic pump (Williamson) at 4°C. The column was then transferred onto an AKTA 10 purifier (GE Healthcare) for further purification steps. The column was washed with, 15-30 times column volume, Tris-HCl buffer containing a suitable concentration of NaCl and imidazole to strip off contaminants and then eluted in buffer containing Tris-HCl pH 7.5, 500 mM NaCl, 500 mM imidazole or 50 mM L-histidine and 0.2% Triton X-100 or 0.05% DDM to isolate the target protein. PD-10 desalting columns (GE Healthcare) were used for desalting and protein was stored at -80°C for further use.

Used columns were washed with distilled water to dispel salts, and 0.1 M EDTA was applied to strip off metal ions. After being washed thoroughly with
distilled water, the columns were recharged with 0.1 M NiSO$_4$ or CuSO$_4$ and kept at 4°C until needed.

2.5.2 Ion exchange chromatography

Protein extraction was followed by ion-exchange purification using a Hitrap Q column (GE Healthcare). The sample was loaded on an AKTA 10 purifier and a salt gradient (10–100% 500 mM NaCl) was used to generate a separate isolated protein peak. Used columns were washed with Tris buffer containing 1 M NaCl and distilled water in order. Columns were stored with 20% ethanol at 4°C.

2.5.3 Detergent exchange

Purified MacB in Triton X-100 (0.2%) was mounted onto a HiTrap Q column and washed with 10 column volume of Tris Buffer 1 (20 mM Tris-HCl buffer, 10 mM NaCl, 10% glycerol, 0.2% DDM, pH 7.5) to remove the Triton X-100 detergent. The column was then washed with an additional 2 column volume of the Tris-HCl buffer 1 containing 0.05% DDM. MacB-DDM was then eluted out with the Tris-HCl buffer 1 containing 100 mM NaCl, and 0.05% DDM.
2.6 Protein Analysis

2.6.1 Electrophoresis

Recombinant proteins were resolved by SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) to determine the efficiency of overexpression and purification. Proteins were visualized on NuPAGE pre-cast Bis-Tris polyacrylamide gels and gradient 4-12% or 12% gels which were adopted as appropriate. A NuPAGE LDS Sample Buffer (4X) (Invitrogen) was used in protein sample preparation and a SeeBlue pre-stained protein marker (Invitrogen) allowed estimation of the protein molecular weight. The recipes for the MOPS running buffer, stain solution and destain solution were as follows:

10X MOPS running buffer

<table>
<thead>
<tr>
<th>Component</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>MOPS</td>
<td>104g</td>
</tr>
<tr>
<td>Tris base</td>
<td>61g</td>
</tr>
<tr>
<td>SDS</td>
<td>10g</td>
</tr>
<tr>
<td>EDTA</td>
<td>3g</td>
</tr>
<tr>
<td>dH2O</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bring up to 1 Liter</td>
</tr>
</tbody>
</table>

Stain solution

<table>
<thead>
<tr>
<th>Component</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanol</td>
<td>40% (v/v)</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>10% (v/v)</td>
</tr>
<tr>
<td>Coomassie Brilliant Blue R250</td>
<td>0.1% (w/v)</td>
</tr>
<tr>
<td>dH2O</td>
<td>50% (v/v)</td>
</tr>
</tbody>
</table>

Destain solution

<table>
<thead>
<tr>
<th>Component</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanol</td>
<td>40% (v/v)</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>10% (v/v)</td>
</tr>
<tr>
<td>dH2O</td>
<td>50% (v/v)</td>
</tr>
</tbody>
</table>
2.6.2 Protein Assay

Protein concentrations were determined using a Coomassie protein assay kit (Bio-Rad). The assay, based on the method of Bradford, is a Coomassie-binding assay where a differential color change of a dye occurs in response to various concentrations of protein. The absorbance maximum for Coomassie Brilliant Blue G-250 dye shifts from 465 nm to 595 nm when it binds to protein and primarily to basic and aromatic amino acid residues, especially arginine (Compton and Jones, 1985).

Protein was diluted 200, 400 and 800 fold into 800μL dH2O to ensure the protein concentration measured was within the working range of the assay (1.2-10μg/ml). The samples were then added with 200 μl of dye reagent concentrate, mixed well, and left at room temperature for 5 min. The absorbance was measured at 595 nm against a dH2O blank. A calibration curve was generated using bovine serum albumin as a standard, and made by plotting the concentration (μg/μl) against the Abs595 and fitting the data by linear regression. Both standard and experimental reactions were performed in triplicate and the concentration of protein was then calculated.

2.6.3 Western blot

Protein samples were run using two SDS-PAGE gels by gel electrophoresis in duplicate, applied as experimental gel and control gel. The proteins on the experimental gel were transferred to a PVDF (Polyvinylidene difluoride) membrane (Millipore) where they were probed with the primary antibody. Mouse monoclonal anti-polyhistidine antibody (Sigma), S•Tag™ monoclonal antibody (Novagen) and T7•Tag® monoclonal antibody (Novagen) are used as primary antibodies as appropriate. The PVDF membrane was then incubated with a second antibody, goat anti-mouse immunoglobulin G AP (alkaline phosphatase) conjugate (Bio-Rad), and protein identified a colorimetric AP conjugate substrate kit (Bio-Rad) so that signal was viewable on a film. All buffers involved in the procedure are listed below:
Transfer buffer

Glycine 29g
Tris base 5.8g
SDS 0.37g
Methanol 20% (v/v)
dH₂O Bring up to 1 Liter

ELISA Wash buffer

*Phosphate stock

Phosphate stock* 30ml
NaCl 8.76g
Tween 20 0.05% (v/v)
dH₂O Bring up to 1 Liter

*Phosphate stock

KH₂PO₄ 6.465g
K₂HPO₄ 35.3g
Adjusted to pH 7.4

dH₂O Bring up to 1 Liter

Blocking buffer

ELISA Wash buffer 25ml
BSA (Bovine serum albumin) 0.75g
2.7 Biochemical assays

2.7.1 ATPase activity

The ATPase activity was determined using stopped flow spectrometer (PiStar-180 spectrometer, Applied Photophysics). EnzChek phosphate assay, a Pi quantification assay (Molecular Probes) was adopted in the measurements of protein ATPase activity. When Pi is present, PNP (purine nucleoside phosphorylase) can enzymatically convert MESG substrate (2-amino-6-mercapto-7-methylpurine riboside) into ribose 1-phosphate and 2-amino-6-mercapto-7-methylpurine, which results in a spectrophotometric shift in the maximum absorbance from 330 nm to 360 nm (Webb, 1992). A standard curve was generated, using KH$_2$PO$_4$ as a standard, by plotting the phosphate concentration (nmoles) against the $A_{360}$ and fitting the data by linear regression. The reaction components are as follows:

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dH$_2$O</td>
<td>740µl - x</td>
</tr>
<tr>
<td>20X Reaction buffer</td>
<td>50</td>
</tr>
<tr>
<td>MESG substrate</td>
<td>200</td>
</tr>
<tr>
<td>KH$_2$PO$_4$</td>
<td>x</td>
</tr>
<tr>
<td>PNP</td>
<td>10</td>
</tr>
<tr>
<td>Total</td>
<td>1000</td>
</tr>
</tbody>
</table>

Measurements at $A_{360}$ were recorded continuously when protein and ATP were mixed from two drive syringes. The reaction temperature was controlled at 25°C by a water bath (Jasco). The recipes for the two reservoirs are as follows:

**Protein sample syringe reservoir**

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dH$_2$O</td>
<td>740µl - x</td>
</tr>
<tr>
<td>20X Reaction buffer</td>
<td>50</td>
</tr>
<tr>
<td>MESG substrate</td>
<td>200</td>
</tr>
<tr>
<td>PNP</td>
<td>10</td>
</tr>
<tr>
<td><strong>Protein (s)</strong></td>
<td>x</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>1000</td>
</tr>
</tbody>
</table>
Substrate sample syringe reservoir

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Volume (μl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dH₂O</td>
<td>740μl - y</td>
</tr>
<tr>
<td>20X Reaction buffer</td>
<td>50</td>
</tr>
<tr>
<td>MESG substrate</td>
<td>200</td>
</tr>
<tr>
<td>PNP</td>
<td>10</td>
</tr>
<tr>
<td>ATP</td>
<td>y</td>
</tr>
<tr>
<td>Total</td>
<td>1000</td>
</tr>
</tbody>
</table>

The data were analyzed by fitting a single exponential plus slope equation:

\[ f(x) = a \cdot \exp(-kx) + bx + c \]

Once the a, b, c, k values were determined, the Abs\textsubscript{360} values were then collected and calculated into Pi concentration. The traces were plotted the phosphate concentration against time using Sigma Plot.

2.7.2 Equilibrium binding of \([^{14}C]\) erythromycin

50 μg of purified protein was incubated with 1, 5 and 10 μM of hot \([N\text{-methyl-}^{14}C]\) erythromycin (48.8 mCi/mmol, Perkin Elmer Life sciences, USA) and cold erythromycin for 30 min. The incubation was carried out in glass tubes in a final volume of 500 μl of buffer containing 10 mM Tris-HCl pH 7.4, 0.05% DDM. The samples were incubated at room temperature in the dark for 45 min, and rapidly filtered through 0.22 μM nitrocellulose filters (Whatmann), which were pre-equilibrated overnight in ice-cold wash buffer containing 10 mM Tris-HCl pH 7.4, 150 mM KCl, 5 mM MgCl\textsubscript{2}. The filters were back-to-back washed once more with 3 ml of ice-cold wash buffer and soaked in scintillation fluid for at least 3 hours until totally dissolved. Subsequently, filter-retained radioactivity was measured by liquid scintillation counting using the scintillant Ultima Gold XR (Perkin Elmer Life Sciences, USA). Non-specific binding to MacB was determined as the amount of \([^{14}C]\) erythromycin bound to half molar quantities of 12 transmembrane helices-containing sugar transporter GalP (Velamakanni \textit{et al.}, 2007), and was less than 30% of total binding.
2.7.3 Pull-down Assay

Pull-down assays were used to determine if there was a physical interaction between two proteins \textit{in vitro}. In a pull-down assay, a tagged bait protein is captured on an immobilized affinity ligand (Ni$^{2+}$ in this study) specific for the tag. The immobilized bait protein can be incubated with putative prey proteins to capture the one that interacts with the bait protein. Membrane pellets containing preyed protein with an S-tag, a 15 residues tag sequence, were dissolved in detergent and mixed with purified bait protein with a polyHis tag. The mixture was loaded into a Ni$^{2+}$ charged Hitrap chelating column (GE Healthcare), so that the bait protein could be immobilised on the column, along with the prey protein if there was an interaction between them. The column was then washed with 15-20 column volumes of 20 mM Tris-HCl buffer containing reasonably high imidazole (75-100 mM) to eliminate any false positive results due to non-specific interactions. The bait-prey complex was eluted with 20 mM Tris-HCl buffer containing 500 mM imidazole; followed by the gel electrophoresis to visualize the bait and prey proteins. A Western blot was carried out using S-Tag monoclonal antibody (Novagen) to probe the prey protein, confirming its identity. To eliminate false positive non-specific interactions between the prey protein and the column, a control experiment was done in a similar manner without the bait protein.
2.7.4 Protein cross-linking

Crosslinker Dithiobis succinimidypropionate (DSP) (Figure 2.1) was adopted for crosslinking of MacB. It contains an amine-reactive N-hydroxysuccinimide (NHS) ester at each end of an 8-carbon spacer arm. NHS esters react with primary amines at pH 7-9 to form stable amide bonds, along with release of the N-hydroxysuccinimide leaving group. DMA and DMS (Figure 2.1) belong to homobifunctional imidoester cross-linkers. The imidoester functional group is available for the modification of primary amines and has minimal cross reactivity toward other nucleophilic groups in proteins.

Figure 2.1 Structures of crosslinkers.
2 mg/ml MacB in buffer containing 50 mM Na₂HPO₄ pH 7.5, 50 mM NaCl, 10% glycerol was incubated with 1, 2, 10, 20, 50, 100-fold excess of DSP concentration for 30 min at 30°C. To stop the reaction, 1 M Tris-HCl was added to the samples and boiled for 3 min before being analyzed by gel electrophoresis. Pre-stained marker Seeblue (Invitrogen) and bench marker (Invitrogen) were used for estimation of protein molecular weight.

2 mg/ml MacA, 1 mg/ml MacB or MacA-MacB was also crosslinked by 5 mM DMA or DMS individually for 60 min at room temperature, and analyzed in an analogous manner to that for DSP.

### 2.7.5 Phenol-sulfuric acid colorimetric assay

Detergents are essential for membrane protein isolation and purification. DDM, a glycosidic nonionic detergent, has been used for the membrane proteins manipulated in this study. A rapid, simple and sensitive procedure for the quantification of glycosidic and bile salt-based detergents has been presented (Urbani and Warne, 2005). This method can quantify sugar or cholate moieties by colorimetric reactions with phenol and sulfuric acid, which can be used to quantify DDM. Furfural derivatives are generated when sugars are dehydrated in concentrated sulfuric acid; following a condensation reaction in the presence of phenol, an aromatic molecule, the p-semiquinonoid chromogen is formed, which has as visible absorbance at 490 nm. The colorimetric assay was undertaken to quantify the DDM contribution in the MacB-DDM complex via a similar approach to a previous report (Butler et al, 2004).

The MacB-DDM complex and processing buffer, 20 mM Tris HCl, 50 mM NaCl, 10% glycerol, with or without 0.2% DDM, pH 7.5, were collected after a detergent exchange and the buffer with DDM served as blank for correction for the background in absorbance measurements. The MacB concentration was determined as described in section 2.6.2 before assaying, which enable to estimate the total amount of DDM, from MacB-DDM complex and DDM micelles in the buffer, and keep it in the effective range the standard curve (0-60μg in 60μL). The MacB-DDM complex was diluted as necessary into the original purification buffer (Tris-HCl buffer) without DDM to make a final volume of 60μL. The sample was first mixed with 300 μL phenol (5%, w/v), then 720 μL concentrated sulphuric acid was carefully added and mixed
well by vortexing with the tube cap closed. One thing needs to be paid attention to is
the strong exothermic reaction that occurs on the addition of concentrated sulfuric acid,
during which the temperature can go up to 110-120°C. The mixture was allowed to
cool down at room temperature for 20 min and transferred to a cuvette for the
absorbance measurements at 490 nm using buffer as a background blank. A standard
curve was generated using DDM as a standard, and made by plotting the DDM
concentration (µg/60µL) against the Abs490, fitting the data by linear regression. The
molar ratio of protein to detergent was then calculated.

2.7.6 Growth curve measurements

In order to test the ability of the proteins to assemble into functional
transporters that confer resistance in *E. coli*, drug susceptibility assays were
undertaken by measuring cell growth curves in the presence of erythromycin. Cells of
*E. coli* strain Kam3(DE3), harboring plasmids or blank vectors, were grown in 2xYT
media at 37°C, with shaking at 200 rpm. 0.05-0.1 mM IPTG was added when the cells
density reached an OD600 of 0.5-0.6. Cells were grown for overnight and diluted into
media, incubated with the antibiotic erythromycin at the concentrations of 100 µg/ml.
Cells growth curves were recorded over the next 12 hours. For experiments to measure
the loss in growth due to erythromycin, cells were grown for 3 hours in the absence
and presence of 50 µg/ml erythromycin and the relative growth is the ratio of the
OD600 values.
2.8 Structural approaches

2.8.1 Dynamic light scattering (DLS)

A Zetasizer Nano instrument (Malvern) was used for DLS analyses in order to determine particle distribution and molecule weight for the MacB protein. The protein was dialyzed against buffer containing 20 mM Tris-HCl pH 7.5, 100 mM NaCl, 1% glycerol, and 0.005% βDMM overnight at 4°C and diluted to a final concentration of 1 mg/ml. The sample was centrifuged at 14,000 xg for 10 min in order to check protein’s stability before analysis. The data for the protein sample and buffer itself were recorded at 4°C and 16°C.

2.8.2 Analytical size exclusion chromatography (SEC)

A superdex 200 PC 3.2/30 column (GE Healthcare) was used in this experiment. The column was equilibrated with degassed and filtered buffer containing 20 mM Tris-HCl pH 7.5, 300 mM NaCl, 10% glycerol and 0.05% DDM for 2 column volume. The sample was filtered through a 0.22 μm filter (Millipore) and centrifuged at 14,000 xg for 5 min before loading onto a column. The sample injection was accomplished by using a 100 μl loop and a 1 ml syringe. A high molecular weight gel filtration calibration kit containing aldolase (Mr 158 000), catalase (Mr 232 000), ferritin (Mr 440 000), thyroglobulin (Mr 669 000) and blue dextran 2000 served as standards for estimation of molecular weight of the protein. The used column was washed with distilled water and stored with 20% ethanol at 4°C.

The Mr of protein was calculated from a calibration curve generated by plotting the log values of the Mr of the standards against their gel-phase distribution coefficient values ($K_{av}$). The $K_{av}$ values were calculated using their elution volumes ($V_e$), total bed volume of 2.4 ml ($V_i$) and the void volume ($V_o$) according to the equation $K_{av} = (V_e-V_o)/(V_c-V_o)$, where $V_c$ represents total column volume.

2.8.3 Analytical ultracentrifugation (AUC)

AUC is one of the most powerful and versatile techniques for the quantitative characterisation of macromolecular associations in solution (Howlett et al., 2006). Sedimentation equilibrium measurements were performed using a Beckman Optima XL-A analytical ultracentrifuge equipped with both absorbance and interference optics,
allowing optical signals to be measured as a function of radial distance. 100 μL of 0.2 mg/ml MacB in buffer 1 (20 mM Tris 8.0 150 mM NaCl, 1% glycerol and 0.006% βDDM) supplemented with 10%, 25% or 50% D₂O was placed in the sample compartment of a epon double-sector centerpiece and 110 μl buffer 1 was placed in the reference compartment. The samples were centrifuged at 10°C and 10,000, 15,000 and 25,000 rpm using an An60-Ti rotor. Scans were acquired using the absorbance optical system 15 hours after the start of the experiment and in 1 hour intervals until equilibrium was attained. The buffer densities and partial specific volume of MacB were calculated using the program SEDNTERP (Laue et al., 1992) and the partial specific volume of βDDM was taken as 0.809 cm³/g (Peterson et al., 1988). The equilibrium data was analyzed using SEDPHAT (Vistica et al., 2004). Sedimentation velocity measurements have been performed using the same hardware at 55,000 rpm at 10°C. The measurements were performed in buffer 1.

2.8.4 Circular dichroism (CD) spectroscopy

Circular Dichroism (CD) is a valuable technique for examining the structure of proteins in solution (Kelly et al., 2005). CD analysis was used to determine the secondary structure compositions of proteins from the peptide bond region in this study. CD spectroscopy and data collection was achieved using a CD spectrometer, with a Xe arc light source (Jasco J-810). It is difficult to make measurements below 180 nm by a conventional CD instrument since the intensity of the radiation falls off in this region; besides, both the N₂ used for purging the sample compartment and H₂O solvent absorb significantly. Protein secondary structure observations by CD spectroscopy occur in the region of 190-260 nm and the absorption is due to the peptide bond. There is a broad but weak signal centred around 220 nm based on n to π transition and a more intense π to π* transition centred around 190 nm. By determining the CD spectrum in this range, the secondary structure of a protein can be determined.

The priority for CD spectroscopy analysis is to keep the total absorbance of the sample within reasonable bounds in order to avoid excessive noise. The signal/noise ratio for CD is theoretically a maximum when the absorbance is 0.869 (Fasman, 1996). The absorbance of the sample is conveniently monitored by the trace of the High Tension voltage (the voltage applied to the photomultiplier). For reliable data, this
should be less than 700 V, but this value will depend on the particular instrument being used.

The NaCl and imidazole in the protein samples were reduced as much as possible by dialysis to minimize the strong absorbance from the components of the buffer. The spectrum of proteins in 20 mM Tris buffer, 20 mM NaF, pH 7.5 was recorded from 260 nm to 190 nm, six times and averaged; from this average spectrum the baseline spectrum for buffer itself was subtracted to get the final spectrum. Data was then input into the online server Dichroweb at Birbeck College (Whitmore and Wallace, 2004) and analyzed by a number of algorithms, including SELCON3, CDSSTR and CONTIN-LL.

2.8.5 Electrospray (ES) mass spectrometry

MacB sample purification and preparation were accomplished as described in section 3.1. Analyses were performed using a nanoflow ES mass spectrometers Q-ToF2 (Micromass) modified for transmission and isolation of high mass ions (Sobott et al., 2002). The following experimental parameters were used to record the mass spectra of 2 mg/ml MacB on the Q-ToF2 instrument: a needle voltage of 1.5 kV and an MCP of 2350 V.

2.8.6 Atomic force microscope (AFM)

MacB sample purification and preparation were accomplished as described in section 3.1. 1 mg/ml MacB in buffer containing 20 mM Tris pH 7.5, 50 mM NaCl, 10% glycerol and 0.05% βDDM was diluted 1000 times to a final concentration of 1 μg/ml, and 45 μl of the sample was allowed to adsorb to freshly cleaved mica. After 10 min incubation, the sample was washed with Milli-Q water and dried under nitrogen. Imaging was performed with a Multimode atomic force microscope (Digital Instruments, Santa Barbara, CA). Samples were imaged in air and experiments were carried out in tapping mode. The silicon cantilevers containing a diamond-like extra tip had a drive frequency of ≈300 kHz and a specified spring constant of 40 N/m (MikroMasch, Portland, OR). The applied imaging force was kept as low as possible (target amplitude ≈1.6-1.8 V and amplitude set-point ≈1.3-1.5 V).
The molecular volumes of the protein particles were determined from particle dimensions based on AFM images. After adsorption of the receptors onto the mica support, the particles adopt the shape of a spherical cap. The heights and half-height radii were measured from multiple cross-sections of the same particle, and the molecular volume was calculated by using the following equation:

\[ V_m = \frac{\pi h}{6} \left( 3r^2 + h^2 \right) \]  

where \( h \) is the particle height and \( r \) is the radius (Barrera et al., 2005; Schneider et al., 1998).

Molecular volume based on molecular mass was calculated by using the equation

\[ V_c = \left( \frac{M_0}{N_0} \right) \left( V_1 + dV_2 \right) \]  

where \( M_0 \) is the molecular mass, \( N_0 \) is Avogadro's number, \( V_1 \) and \( V_2 \) are the partial specific volumes of particle and water, respectively, and \( d \) is the extent of protein hydration (Barrera et al., 2005; Schneider et al., 1998). We used the value of 0.74 cm\(^3\)/g for the partial specific volumes of protein, and 0.4 g of water per g of protein for the extent of protein hydration (Grant, 1957). Using automated recognition of proteins (Barrera et al., 2007; Barrera et al., 2008), it was calculated the molecular volume of isolated particles as described above.

2.8.7 Transmission electron microscope (TEM)

MacA and MacB sample purifications and preparations were accomplished as described in section 4.1 and 3.1 respectively. Protein samples in buffer containing 20 mM Tris pH 7.5, 500 mM NaCl, 10% glycerol and 0.2% TX-100 were pipetted onto the carbon-coated side of a grid. After 1 min, filter paper was used to take off excessive liquid without touching the grid. The grid was then negatively stained using 1% uranyl acetate for 30 seconds. Filter paper was used to take off excess liquid without touching the grid. The grid was then examined using a transmission electron microscope instrument (Hitachi).
2.8.8 Crystallization

Crystallization is an important tool for the determination of the structure of biological macromolecules through the use of X-ray diffraction for analyzing the atomic arrangement of crystals. Proteins were initially screened with commercial sparse matrix screens that provided a broad range of parameters to identify good crystallization conditions. Once an initial condition, good for protein crystals to grow, was found, modification of the crystallization conditions was then carried on to obtain crystals that are of X-ray diffraction quality.

2.8.8.1 Sitting drops technique

Crystallization trials were undertaken using the vapour diffusion sitting drop method on Greiner crystallization plates, which have 96 round wells; crystallization conditions were screened automatically using a Cartesian robot. The volume of each screen drop was around 50 nl. After setting up the drops the whole tray was sealed and spun at 1000 xg for 1 min in order to pull the drops into good shape at the bottom of the well. Crystals were grown at 18 °C and their development was monitored intermittent periods by light microscopy.

2.8.8.2 Hanging drops technique

Crystallization trials were carried out using 24-well crystallization plates (Nextal). A droplet of protein solution was mixed with crystallization reagent on a platform inverted over the reservoir in vapor equilibration with the reagent. The initial reagent concentration in the droplet is less than that in the reservoir but will eventually reach equilibrium and settle between the drop and the reservoir when the reservoir pulls water from the droplet in the vapor phase. During this equilibration process, the sample is also concentrated, increasing the relative supersaturation of the sample in the drop. The plates were incubated at 18°C and the droplets were monitored by light microscopy to identify different stages of crystallization.
Chapter three: Determination of the MacB oligomeric state

The ABC transporter MacB has been shown to confer macrolides resistance in the presence of TolC via active drug efflux (Kobayashi et al., 2001). MacB has a topology that consists of one N-terminal cytoplasmic nucleotide binding domain, four transmembrane helices and one large periplasmic loop between the helix 1 and 2. The MacB topology was determined by systematically replacing residues with cysteines and testing for the \[^{14}C\] NEM binding (Figure 1.27) (Kobayashi et al., 2003)

Most ABC transporter, have been found to consist of four domains: two nucleotide binding domains (NBD) that serve as engines for ATP hydrolysis, which is coupled to active transport, and two transmembrane domains (TMD) that are believed to constitute the transport pathway. The 3D structures of such as Sav1866, MsbA and ModB\(_2\)C\(_2\)A are in an agreement with a four domain organization in full-size ABC transporters (Dawson and Locher, 2006; Hollenstein et al., 2007). Well studied ABC exporters such as LmrA, Sav1866 and MsbA are believed to form a full transporter with twelve transmembrane helices; six helices being contributed by each subunit (Borges-Walmsley et al., 2003). After detergent solubilization the ABC transporter BmrA, from Bacillus subtilis, was found to be a dimer using analytical ultracentrifugation (Ravaud et al., 2006). Exceptionally, MacB has only "four" transmembrane helices rather than six, leading to the question of MacB's functional oligomeric state. In the tripartite efflux pump AcrA-AcrB-TolC, AcrB and TolC are proven to form trimers, whereas the AcrA oligomeric state is uncertain. MacB confers TolC-dependent macrolide resistance, which might indicate that MacB also forms trimers in order to coordinate TolC.

In this chapter, I report on the development of procedures that provide a good yield and purity of MacB, which has enabled detailed characterisation of its functional and structural states. Size exclusion chromatography (SEC), analytical ultracentrifugation (AUC), electro-spray mass spectrometry and atomic force microscope (AFM) were applied to discover the oligomeric state of MacB when in a detergent-solubilized state.
3.1 MacB-His<sub>6</sub> construction and production

Cells of the *E. coli* strain Novablué, derived from *E. coli* K12, served as the source of chromosomal DNA for the *macA* and *macB* genes. The completed genome of *E. coli* is available from the NCBI database, as a source of sequence information for the genetic manipulation of these genes (section 2.1). A list of primers, vector plasmids and *E. coli* strains used in the study can be found in section 2.3 and 2.4. The *macB* gene of *E. coli* was amplified by PCR, incorporating artificial restriction enzyme sites, suitable for cloning into a chemically inducible expression vector. The detailed methods for the construction can be found in section 2.4, genetic manipulation.

The forward primer, Ndel-macB, and reverse primer, SacI-stop-macB, were designed to incorporate the restriction enzyme (RE) sites, Ndel and SacI, at the 5' and 3' ends of the amplified *macB*, respectively. Successful PCR amplification of the *macB* sequence was shown to be consistent with the predicted size on an agarose gel (Figure 3.1a). The amplified *macB* gene was extracted and purified from the agarose gel and ligated into the A-T propagation pGEM-T Easy vector. The ligation mixture was then transformed into chemically competent *E. coli* NovaBlue. The pET28a(+) expression vectors and the recombinant pGEM-T Easy vectors were digested with Ndel and SacI simultaneously, and the digests were then separated by gel electrophoresis. The *macB* gene and linear pET28a were cut out and purified; followed by cohesive end ligation using T4 DNA ligase. The ligation mixture was transformed into chemically competent *E. coli* NovaBlue. Plasmid DNA purified from recombinant colonies was digested with Ndel and SacI to confirm the presence of inserted DNA (Figure 3.1b).

The pET28a-*macB* expression construct was transformed into the *E. coli* strain C43(DE3) (Miroux and Walker, 1996) for protein production. The protocols for protein expression and for harvesting cell membranes were as described in section 2.5. The cell membranes were dissolved in 20 mM Tris buffer pH 7.5, 50 mM NaCl, 10 mM imidazole, 1 mM THP, 10% glycerol and 5% Triton X-100 on a rotary shaker overnight.

MacB-His<sub>6</sub> was purified by metal affinity chromatography using HiTrap Cu<sup>2+</sup> column (section 2.5.1). MacB-His<sub>6</sub> was eluted in buffer containing 20 mM Tris-HCl pH 7.5, 500 mM NaCl, 10% glycerol, 50 mM L-histidine and 0.2% Triton X-100. The buffer was exchanged into 20 mM Tris-HCl pH 7.5, 10 mM NaCl, 10% glycerol, and
0.2% Triton X-100 by using PD-10 columns prior to the second purification by ion exchange chromatography using HiTrap Q column (section 2.5.2).

The efficiency of expression and purification of MacB-His<sub>6</sub> was determined by gel electrophoresis (section 2.6.1) (Figure 3.2). The migration of the major protein band (lanes 2 to 8) is between the 51 kDa and 64 kDa of the SeeBlue protein marker (lane 1), which is marginally different from the MacB-His<sub>6</sub> calculated molecular weight of 72.97 kDa. The identification of MacB-His<sub>6</sub> protein was confirmed by Western blot, as shown in figure 3.2.

**Figure 3.1** pET28a/macB construction (a) PCR amplification of *macB* gene from *E.coli* K-12. Lane 1= DNA ladder. Lane 2= the amplified *macB* gene. (b) Restriction enzyme digestion analysis of the pET28a/macB plasmid. Lane 1= ladder. Lanes 2-3 indicate the presence of the *macB* gene.
Figure 3.2  SDS-PAGE of MacB-His<sub>6</sub>. Lane 1 = protein molecular weight (M<sub>r</sub>) marker. Lanes 2-8 indicate the MacB proteins. The signals for MacB-His<sub>6</sub> were shown on the X-ray film by using Western blot.

3.2 Size exclusion chromatography of MacB

Analytical SEC was used to determine the molecular weight of MacB in order to investigate its oligomeric state. The analysis was performed on an AKTA using a Superdex200PC3.2/30 column as described previously (section 2.8.2). The elution profiles of MacB were monitored using a 280 nm UV detector.

Blue dextran was applied to the column to determine the void volume (V<sub>o</sub>) of the column. The protein molecular mass standards included aldolase, catalase, ferritin and thyroglobin, which were run individually in order to determinate the K<sub>a</sub> for each standard (Table 3.1), which was then used to generate a calibration curve (r<sup>2</sup> > 0.99) (Figure 3.3).
<table>
<thead>
<tr>
<th>Standard</th>
<th>Mr (kDa)</th>
<th>Concentration (mg/ml)</th>
<th>$V_e$ (ml)</th>
<th>$K_{av}=(V_e-V_0)/(V_c-V_0)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blue dextran</td>
<td>--</td>
<td>2</td>
<td>1.01</td>
<td>--</td>
</tr>
<tr>
<td>Aldolase</td>
<td>158</td>
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<td>1.59</td>
<td>0.42</td>
</tr>
<tr>
<td>Catalase</td>
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<td>5</td>
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<tr>
<td>Ferritin</td>
<td>440</td>
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<td>0.22</td>
</tr>
<tr>
<td>Thyroglobin</td>
<td>669</td>
<td>3</td>
<td>1.18</td>
<td>0.12</td>
</tr>
</tbody>
</table>

Table 3.1 Generation of the gel-phase distribution coefficient values ($K_{av}$). 

$K_{av} = (V_e-V_0)/(V_c-V_0)$, $V_e$ = elution volume for the protein, $V_0$ = column void volume and $V_c$ = geometric column volume (2.4 ml).

Figure 3.3 A molecular weight calibration curve generated for protein standards on an analytical Superdex 200 PC 3.2/30 gel filtration column.
MacB was first extracted and purified in Triton X-100; however, Triton X-100 absorbed strongly at 280 nm, making it impossible to monitor the MacB elution profile. Thus, MacB-His$_6$ was exchanged into buffer containing Tris-HCl pH 7.5, 300 mM NaCl, 10% glycerol and 0.2% DDM before being analyzed by SEC (section 2.5.3). Using an Enzchek Assay, MacB was confirmed to retain ATPase activity and retained activity after detergent exchange (data not shown). The MacB-His$_6$ was applied to the column and it showed a sharp peak at an elution volume of approximate 1.32 ml ($V_e$), giving a $K_{av}$ value of 0.223, corresponding to a Mr for the MacB-His$_6$/DDM complex of 415 kDa (Figure 3.4). The average Mr for MacB-His$_6$/DDM complex calculated from 2 runs in the solution is 410 kDa ± 10 kDa.
3.3 Quantification of DDM contribution in MacB/DDM complex

Considering that MacB was in a detergent solubilized state, the detergent could make a significant contribution to the Mr of the protein-detergent complex (Ravaud et al., 2006). In order to determine the contribution of DDM in the MacB/DDM complex, a phenol-sulfuric acid colorimetric assay was applied to quantify the DDM contribution (section 2.7.5). A standard curve containing six points between 0 µg to 20 µg of DDM was produced with a line of best fit ($r^2 > 0.99$) (Figure 3.5).

![Standard curve of DDM quantity by using the colorimetric assay.](image)

A MacB protein sample and its elution buffer were obtained from the same batch used for the SEC analysis in the previous section, and used to determine the quantity of bound DDM using a colorimetric assay. This MacB sample was determined to have concentration of 0.85 mg/ml using coomassie protein assay kit (section 2.6.2). The colorimetric assays were performed in triplicate and two dilutions (1:10 and 1:20) to ensure that the amount of DDM detected was within the range of the standard curve. Using the equation generated from the standard curve, the amount of DDM bound by MacB was determined spectrometrically to be 1.2 g DDM/g MacB (Table 3.2), which is equivalent to a MacB: DDM molar ratio of 1: 171. Ravaud et al in 2006 concluded that the ABC transporter BmrA is a dimer in the detergent solubilized state and that the amount of DDM bound by BmrA was about 1.5 g DDM/g BmrA. BmrA consists of six transmembrane α-helices whereas MacB has only four transmembrane α-helices,
suggesting that the DDM: MacB ratio is reasonable. These results indicate that for every MacB molecule, bound DDM adds a further 88 kDa to the Mr of the complex, resulting in the Mr for a MacB monomer, with bound DDM, to be about 161 kDa. However, we should also consider that there would be a quantity of bound lipid, which would make a further contribution to the total Mr of the complex but this is probably less significant (Ravaud et al., 2006). The estimated Mr of 410 kDa for the complex from the SEC analysis suggested that there are at least two MacB protomers in the MacB/DDM/lipid complex, but the Mr of 410 kDa is slightly higher than expected for the dimer. The size estimate from the SEC analysis is on the basis of the radius of gyration. If the particle is elongated, it tends to overestimate the size of the particle. Since MacB has a membrane domain that is sandwiched by cytoplasmic and periplasmic domain it is likely to have an elongated structure.

<table>
<thead>
<tr>
<th>1 to 10 Dilution (MacB 5.1 μg/60 μL sample volume)</th>
<th>1 to 20 Dilution (MacB 2.55 μg/60 μL sample volume)</th>
</tr>
</thead>
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<tr>
<td><strong>Abs</strong>&lt;sub&gt;490&lt;/sub&gt; (blank off)</td>
<td><strong>Abs</strong>&lt;sub&gt;490&lt;/sub&gt; (blank off)</td>
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<tr>
<td>1</td>
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<td>0.2511</td>
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<tr>
<td>3</td>
<td>0.2489</td>
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</table>

Table 3.2 The MacB/DDM ratio in the protein-detergent complex. Both the g/g and Molar/Molar ratio have been represented.
3.4 Crosslinking of MacB in vitro

In order to stabilize the oligomeric state of the detergent solubilized complex, crosslinking experiments were undertaken in collaboration with Dr. Vassiliy N. Bavro at the University of Cambridge.

Since Tris can quench the crosslinking reaction, purified MacB was exchanged into buffer containing 50 mM Na$_2$HPO$_4$ pH 7.5, 50 mM NaCl, 10% glycerol and 0.05% DDM prior to crosslinking. The MacB crosslinking in vitro was performed using DSP, a crosslinker with 12 Å spacer arms (section 2.7.4), and analyzed by gel electrophoresis. The Mr of the MacB-His$_6$ monomer was predicated to be 72.97 kDa. On the gel, the MacB monomer was located between the 60 and 70 kDa protein makers and the most predominant band, predicated to be the MacB dimer, which was located between the 120 and 160 kDa markers (Figure 3.6a). However, the presence of some higher Mr bands may suggest MacB can also form higher order oligomers or erroneous crosslinking by the long spacer arm crosslinker DSP. In order to confirm the MacB oligomeric state and eliminate the possibility of erroneous crosslinking, the homobifunctional imidoester cross-linkers DMA (8.6 Å spacer arm) and DMS (11 Å spacer arm) were adopted in the MacB crosslinking. MacB dimers were clearly observed, whilst the higher order oligomers had vanished (Figure 3.6b), consistent with MacB forming dimers.

![Figure 3.6 SDS-PAGE of the crosslinked MacB-His$_6$.](image)

(a) MacB was crosslinked with 1, 2, 10, 20, 50, 100 folds (Lane 2-8) of concentration of DSP. (b) MacB was incubated with 5 mM DMA or DMS (Lane 2 and 3) individually.
3.5 AUC analysis of MacB

AUC analysis and data representation were carried out in collaboration with Dr. Martin Montcrieffe and Dr. Vassiliy N. Bavro at the University of Cambridge.

In order to investigate more evidence that MacB is in a dimeric state, a combination of sedimentation equilibrium and sedimentation velocity analyses using analytical ultracentrifugation was performed. The MacB was diluted into buffer containing low DDM (just below the CMC [critical micelle concentration]) (Section 2.8.3) in order to minimize the intervention of DDM micelles. The protein was stable in this buffer condition for over a week, allowing several datasets to be collected from the same sample. To determine the detergent contribution in the buoyant mass of the protein-detergent complex, a series of different density buffers were used, which were prepared using different D$_2$O concentrations (section 2.8.3).

The apparent Mr for MacB was determined from sedimentation equilibrium measurements to be 162.6 kDa (Figure 3.7a), which corresponded to a MacB dimer (The Mr of a MacB monomer is estimated to be 72.97 kDa). Since the equilibrium is only dependent upon the mass as it is calculated by specific buoyancy, buffers with different D$_2$O concentrations so as to provide different densities were prepared for the experiments in order to negate the detergent contribution in the complex. The idea is basically to create a buffer with the same background buoyancy as the detergent compound so that it is solely the mass of protein that is detected. For technical reasons we could not exactly match the buffer density to that of the DDM; fortunately, we used several different buffer compositions and for each of these we determined the equilibrium mass. Since the protein contribution is constant, it was possible to extrapolate the equilibrium mass to the theoretical point at which the buffer matches the specific density of DDM (Peterson et al., 1988). Sedimentation velocity is shape dependent and a sedimentation coefficient of 6.8 S was generated, which corresponded to a Mr of 160 kDa (Figure 3.7b). The sedimentation velocity, where detergent was taken into account, gave a very similar result to the sedimentation equilibrium, suggesting that few detergent molecules were bound. Our colorimetric data showed that there were 171 molecules of DDM bound to one MacB molecule in the buffer containing 0.2% DDM. However, the DDM in the sample for AUC analysis was much lower at 0.006%, just below the CMC, which may lead to the difference in the data.
Figure 3.7  **AUC analysis of MacB.** (a) Sedimentation equilibrium profiles of the MacB-DDM. It shows a representative sedimentation equilibrium profile from one of the runs (two different velocities of the same sample). Experimental data (dots) and fitted model (solid line) is shown for each. The bottom panel is the residuals (differences between data and model) after fitting. (b) Sedimentation velocity measurements of the MacB-DDM. Sedimentation velocity profiles of the MacB are consistent with a stable dimer formation.
3.6 Electrospray mass-spectrometry

Electrospray mass-spectrometry experiments and data representation were carried out in collaboration with Dr. Nelson P. Barrera and Dr. Vassiliy N. Bavro at the University of Cambridge.

Electrospray mass-spectrometry was used to determine accurately the Mr of the protein, under conditions that would dissociate the DDM, revealing a peak with a Mr of 145.96 ± 0.02 kDa, which is consistent with a MacB dimer (Figure 3.8). With known mass-to-charge ratio (m/z) and numbers of positive ion, the molecular weight of MacB can be calculated.

The use of mass spectrometry to determine the Mr of membrane proteins is a novel and challenging field and our results are the first, to the best of our knowledge, to report on the determination of the oligomeric state of a native membrane protein from the ABC transporter family.

![Mass spectrum of MacB](image)

**Figure 3.8** Mass spectrum of MacB. The charge states corresponding to the peaks are graphed.
3.7 AFM analysis of MacB

AFM analysis and data representation were carried out in collaboration with Dr. Nelson P. Barrera and Vassiliy N. Bavro at the University of Cambridge.

Using AFM larger particles could clearly be seen at high-resolution to consist of two protein domains that are highly suggestive of a dimer (Figure 3.9). Not surprisingly, in the presence of AMP-PNP, a non-hydrolysable analogue of ATP, the ratio of dimers to monomers on the AFM grids increased, which would be consistent with the nucleotide stabilizing the interaction between monomers (Figure 3.10).

Our findings are novel because detergent molecules tend to impair the resolution of AFM studies of detergent solubilized membrane proteins; so that in the case of MacB, we may be able to get information on the topology and stoichiometry of its assemblies with MacA and TolC. Indeed, when MacA, which was resolved as a single population of particles with a volume of 69 nm$^3$, was mixed with MacB, a number of particles with volumes far in excess of 238 nm$^3$ (e.g. MacB dimer volume) and 307 nm$^3$ (e.g. MacB dimer volume + MacA volume) were resolved, indicative not only of the interaction of MacA and MacB but also the formation of multi-subunit complexes (data not shown).
Figure 3.9  **AFM imaging of MacB.** (a) 3D picture of a low magnification image of MacB acquired in air in Tapping Mode with a diamond-like extratip of resonant frequency~300 kHz and spring constant of 40 N/m. Letters 'm' and 'd' represent MacB monomer and dimer, respectively. (b) High resolution images of structures where two small particles ('m'+m') are attached to one another, clearly indicative of dimer formation.
Figure 3.10  Frequency distribution of molecular volumes of MacB. (a) The curve indicates a fitted Gaussian function. The 'm' and 'd' peaks correspond to volumes of 118 ± 1 nm$^3$ (n= 1642) and 238 ± 5 nm$^3$ (n= 665). The data is consistent with the 'm' and 'd' peaks corresponding to the MacB monomer and dimer, respectively. (b) Frequency distribution of molecular volumes of MacB that had been incubated with the non-hydrolysable ATP analogue AMP-PNP. The peaks correspond to volumes of 112 ± 3 nm$^3$ (n= 94) and 218 ± 14 nm$^3$ (n= 116). This data indicates an increase in the dimer: monomer ratio in the presence of AMP-PNP.
3.8 Discussion

Membrane proteins are notoriously difficult to overexpress and manipulate due to their high hydrophobicity; and the integral membrane protein MacB, a ABC transporter from *E. coli*, is no exception. The *macB* gene was first cloned into the plasmid pQE80 (Qiagen) for overexpression in the *E. coli* strain M15/PREP4 (Qiagen), under the regulation of a T5 promoter. However, MacB tended to stop the cells from growing once induced and it degraded extensively using this expression system, resulting in an extremely low yield and stability (data not shown). The yield and stability of MacB protein was significantly improved using the T7 promoter regulated plasmid pET28a(+), along with the *E. coli* strain C43(DE3). Even so, using the latter expression system of the yield of pure MacB protein, after two purification steps, was only about 1 mg of protein from 2 L cells. MacB lost its ATPase activity when being extracted with 2% DDM, but retained activity when extracted with 5% Triton X-100 and subsequently switched into a buffer containing 0.2% DDM. This observation indicates that the activity of MacB is detergent-dependant, but the reason is unknown.

Our colorimetric analysis indicated that there were 171 molecules of DDM bound to one MacB molecule, which is consistent with the values found for other membrane proteins (Butler *et al.*, 2004; Ravaud *et al.*, 2006). EmrE, the best studied SMR protein, was found to be a dimer in the detergent-solubilized state, with a molar ratio of DDM/EmrE of 104.5 ± 6 (Butler *et al.*, 2004). BmrA was found to be dimeric in the detergent solubilized state and the molar ratio of DDM/BmrA was determined to be 194 (Ravaud *et al.*, 2006). However, our sedimentation velocity AUC analyses, where the detergent was taken into account, gave a DDM/MacB of molar ratio of 24. This difference might be due to using a DDM concentration of 0.006%, slightly below the CMC, in the buffer for AUC analysis.

In this study, ES-MS has been used to successfully identify MacB oligomeric state. ES-MS is now widely accepted as a powerful method to determine accurately the stoichiometry of intact protein complexes (Hernandez and Robinson, 2001; Sharon and Robinson, 2007). However, it has remained extremely difficult to analyze membrane proteins, solubilized in detergent due to their hydrophobicity, under similar conditions. So far, only a few studies have reported the observation of membrane proteins or their...
complexes in MS (Hanson et al., 2003; Ilag et al., 2004; Lengqvist et al., 2004; Meier et al., 2007). In this study, using nanoflow ES (a miniaturized form of ES with reduced flow rates) and a high collision energy, which facilitates the desolvation process and induces dissociation of detergent-proteins clusters, we demonstrated for the first time the native dimeric state of MacB. Under our experimental conditions, it was possible to successfully maintain the non-covalent subunit interactions, determining the oligomeric state of the protein complex without ambiguity. The development of strategies to tackle this field is challenging, primarily since the large quantities of detergent suppress the protein signal while the highly hydrophobic membrane proteins in aqueous buffers often blocks electrospray needles.

Also of interest is the association between structural results provided by MS and AFM. AFM imaging has been applied previously to determine the stoichiometry of purified membrane proteins (Barrera et al., 2005; Barrera et al., 2008). Interestingly, using diamond-like extratips to scan the topography of isolated MacB, it was possible to identify both subunits forming the complex. Although AFM imaging cannot give very accurate measurements of molecular masses of particles, it provides a reasonable range for the MacB protein (approximately 130-140 kDa). The structure of each component of the AcrA/AcrB/TolC complex pump has been determined, giving some insight into how the complex is assembled; however, the stoichiometry of the whole tripartite complex pump is still controversial (Fernandez-Recio et al., 2004; Higgins et al., 2004). In this study, using AFM, each subunit of the MacB dimer can be identified, suggesting the possibility to extend such analyses to determine how the tripartite complex is assembled.
Chapter four: Defining the role of the membrane fusion protein MacA in MacB’s ATP hydrolysis

Gram-negative bacteria such as *E. coli* employ membrane efflux systems to transport antibiotics, noxious chemicals and large protein toxins outside the cell (Koronakis *et al.*, 2004). Multidrug efflux pumps consist of three components: an energy-providing integral inner membrane protein, either an ABC transporter or a proton antiporter of the RND family or MFS (Borges-Walmsley *et al.*, 2003), cooperates with a protein of the TolC exit duct family. The third component of tripartite complex pumps is a membrane fusion protein, largely periplasmic and anchored to the inner membrane by a single transmembrane α-helix or an N-terminal lipid moiety. One of the best studied efflux pumps of *E. coli* is the AcrA/ArcB/TolC system, where the membrane fusion protein AcrA was proposed to provide a physical link between AcrB and TolC.

Kobayashi *et al* in 2001 showed that the MacAB complex confers TolC-dependent macrolide resistance via active drug efflux. In this chapter, the potential roles and domains of membrane fusion protein MacA to interact with MacB will be investigated.

4.1 MacA-His$_6$ construction and production

A construct was made to overexpress the MacA protein in this study. The details of the construction are described in section 3.1. In short, the forward primer, Ncol-macA, and reverse primer, XhoI-macA, were designed to incorporate the RE sites, Ncol and XhoI, at the 5’ and 3’ ends of the amplified macA, respectively. Successful PCR amplification of the macA sequence was shown to be consistent with the predicted size on an agarose gel (Figure 4.1a). Plasmid DNA purified from recombinant colonies was digested with Ncol and XhoI to confirm the presence of the inserted DNA (Figure 4.1b).

The pET21d-macA expression construct was transformed into the *E. coli* strain C43(DE3) for protein production. The cell membranes were dissolved in 20 mM Tris
buffer pH 7.5, 300 mM NaCl, 15 mM imidazole, 1 mM THP, 10% glycerol and 5% Triton X-100 on a rotary shaker overnight.

MacA-His$_6$ was purified by metal affinity chromatography using a HiTrap Ni$^{2+}$ column (section 2.5.1). MacA-His$_6$ was eluted in buffer containing 20 mM Tris-HCl pH 7.5, 500 mM NaCl, 10% glycerol, 500 mM imidazole and 0.2% Triton X-100. The efficiency of expression and purification of MacA-His$_6$ was determined by gel electrophoresis (section 2.6.1) (Figure 4.2). The migration of the major protein band (lanes 2 to 10) between the 39 kDa and 51 kDa bands of the SeeBlue protein marker (lane 1) is consistent with the calculated molecular weight of 41.8 kDa MacA-His$_6$. Further identification of the MacA-His$_6$ protein was achieved by Western Blot, as shown in figure 4.2.

![Figure 4.1 pET21d/macA construction](image)

(a) 1 2 3

1.6 Kb
1 Kb

← Amplified macA gene (1113bp)

(b) 1 2 3 4

← pET21d

← Inserted macA gene (1113bp)

**Figure 4.1** pET21d/macA construction. (a) PCR amplification of macA gene from *E.coli* K-12. Lane 2 = the DNA ladder. Lane 1 and 3 indicate the amplified macA gene. (b) Restriction enzyme digestion analysis of the pET21d/macA plasmid. Lane 1= DNA ladder. Lanes 2-3 indicate the presence of the macA gene.
4.2 MacA regulates the ATPase activity of MacB

Although MacB belongs to the ABC superfamily it has an exceptional architecture. The presence of an NBD in MacB affords the opportunity to test for an interaction with, and modulation of its ATPase activity by, MacA, which can be carried out in solution. The MacB protein was purified as described in section 3.1. and shown to have its detergent-dependent ATPase activity, being active in Triton X-100 but not in a high concentration of DDM. In order to investigate the potential role of the membrane fusion protein MacA in the ATP hydrolysis by MacB, the dye 2-amino-6-mercapto-7-methylpurine riboside (MESG) was used to monitor the time course of ATP hydrolysis by detecting the production of inorganic phosphate (Pi) (section 2.7.1). A standard curve was generated, which illustrated the amount of Pi released in nanomoles (nmoles) plotted against the absorbance at 360 nm (Abs\textsubscript{360}) and analyzed by linear regression. Points plotted were from an average of three independent experiments (Figure 4.3).
Figure 4.3  Standard curve of inorganic phosphate (Pi). Measurements were undertaken at absorbance 360 nm using the EnzChek phosphate assay.

The MacB ATPase activity was determined in the presence of MgCl₂ and various concentrations of ATP. The amount of Pi released in each sample was measured by the change in $A_{360}$ and was calculated by using the equation generated from the standard curve. The time course was characterized by a burst in Pi production, during the first 20 seconds, followed by a slower steady-state rate (Figure 4.4). This kinetic behavior indicated that the ATP was hydrolyzed rapidly in the burst phase of the trace, and further turnovers were rate limited by the slow release of product.

The amplitude of the burst phase, measured at eight different ATP concentrations, from 0.01 mM to 4 mM, ranged from 2.3-2.4 μM (Figure 4.4), which is equivalent to the MacB concentration (2.3 μM). When the MacB concentration was increased to 3.5 μM, the amplitude of the burst phase increased to 3.3 μM (Figure 4.5), indicating that the burst is approximately equivalent to the MacB concentration and that both NBDs within the MacB dimer are functional. The rate constant for the hydrolysis step was determined by fitting the burst phase to an exponential function, yielding a value of about 0.22 s⁻¹, which was independent of the ATP concentration.
Figure 4.4  A plot for Pi generation by MacB on concentration of ATP. MacB's ATPase activity was measured by the time course for the change in Pi concentration. 2.3 µM MacB was used and the ATPase activity was measured in the presence of various concentration of Mg-ATP, from 0.1 mM to 4 mM.

Figure 4.5  A plot for Pi generation by MacB. MacB's ATPase activity was measured by the time course for the change in Pi concentration. 3.5 µM MacB was used, and the ATPase activity was measured at Mg-ATP concentration of 1 mM.
Figure 4.6  A plot for Pi generation by MacAB on concentration of ATP. 2.3 μM MacA and MacB were used, and the ATPase activity was measured in the presence of Mg-ATP.

Figure 4.7  The steady-state rate of Pi production by MacB as a function of the ATP concentration in the absence (lower curve) and presence (upper curve) of an equivalent concentration of MacA. The data are characterized by $V_{\text{max}}$ and $K_{\text{m}}$ values of 8.9 nmoles ATP/mg MacB/min and 374 μM, respectively, for MacB alone; and of 12.3 nmoles ATP/mg MacB/min and 72 μM, respectively, for MacB in the presence of an equivalent concentration of MacA.
The steady-state phase was characterized by a rate of Pi production that increased in a hyperbolic manner with the ATP concentration (Figure 4.7). A maximal specific activity of 8.9 nmoles ATP/mg MacB/min and a $K_m$ of 374 $\mu$M were obtained by fitting the data to a hyperbolic function. The macrolides erythromycin and oleandomycin, substrates of MacB, were also added respectively in the reaction to investigate the possibility of a substrate-induced change in the ATPase activity; however, the presence of the macrolides did not affect ATP hydrolysis by MacB, which is consistent with a previous report on MacB (Tikhonova et al, 2007) but contrasts with the findings for Sav1866 (Dawson and Locher, 2006). The ATPase activity of Sav1866, reconstituted into liposomes, was stimulated by the anti-cancer drugs doxorubicin, vinblastine and the fluorescent dye Hoechst 33342.

An equivalent concentration of MacA was added to 2.3 $\mu$M MacB, the ATPase activity was determined in the presence of MgCl$_2$ and various concentration of ATP. Surprisingly, the Pi burst phase disappeared but the steady-state rate was enhanced (Figure 4.6). This behavior might suggest that MacA increased the rate of ADP dissociation from MacB, making this step no longer rate-limiting. However, $k_{cat}$ did not increase to a rate above that determined for the burst phase and so we must assume that the hydrolysis rate also decreases.

Figure 4.7 illustrated the steady state ATPase activity of MacB on various ATP concentrations. The non-linear regression line was obtained by fitting in a hyperbolic trace. The equation of the regression line is as follows:

$$y = \frac{a \times x}{b + x}$$

In the equation, "a" represents $V_{max}$ and "b" represents $K_m$. The $k_{cat}/K_m$ ratio is often thought of as a measure of enzyme efficiency. Both a big value of $k_{cat}$ (rapid turnover) or a small value of $K_m$ (high affinity to substrate) can makes $k_{cat}/K_m$ large.

Although MacA increased the specific activity of MacB from 8.9 to 12.3 nmoles ATP/mg MacB/min, which would correspond to an increase in $k_{cat}$ from 0.011 to 0.015 $s^{-1}$, the steady-state rate did not exceed the rate for hydrolysis in the absence of MacA (e.g. 0.22 $s^{-1}$). Accordingly, the increase in the steady-state rate, which was attributed
to product release, would not be rapid enough to prevent the phosphate burst. Consequently our results suggest that MacA hinders the rate of ATP hydrolysis, so that this process is no longer rate limited by the product release. This finding contrasts with a previous report on MacB, where the presence of MacA had no affect on the ATPase activity of MacB in solution. However, they did find that MacA, when co-reconstituted with MacB, had the effect of increasing both $k_{cat}$ (e.g. $k_{cat}$ from 0.1 to 0.78 s$^{-1}$) and the affinity of MacB for ATP (e.g. $K_m$ from 2.3 to 0.38 mM), which is similar to the effect of MacA on the steady-state kinetics of ATPase activity of MacB in a detergent solubilized state in our study.
4.3 The MacA periplasmic domain

The previous section indicated that MacA regulates the ATPase activity of MacB. MacA is a membrane fusion protein with an anchored transmembrane domain. To investigate the possibility that the transmembrane domain is involved in the interaction with MacB, soluble truncated MacA, in which the first N-terminal 20 residues were deleted, was generated based on the predication of its secondary structure, using the program Jpred at University of Dundee (Section 2.1). Figure 4.8 illustrates the short transmembrane α-helix (pink) and large periplasmic domain (blue).

![Diagram of secondary structure prediction of E. coli MacA. The predicted secondary structure elements of MacA are drawn above the amino acid sequence with the α-helices and β-sheets depicted as H and E respectively. Horizontal arrows drawn above secondary structure describe predicted protein topology (Green arrow indicates cytoplasmic domain, pink arrow indicates membrane spanning domain and blue arrow indicates periplasmic domain).]
4.3.1 Truncated MacA construction and production

A construct was made for the expression of the N-terminal truncated MacA. The details of the construction are described in section 3.1, indicating the changes in DNA template and primers. In short, the forward primer, Ncol-FD20macA, and reverse primer, XhoI-macA, were designed to incorporate the RE sites, Ncol and XhoI, at the 5' and 3' ends of the amplified macA sequence, respectively. Plasmid DNA purified from recombinant colonies was digested with Ncol and XhoI to confirm the presence of the inserted DNA (Figure 4.9a).

The pET21d-Δ20macA expression construct was transformed into the E. coli strain C43(DE3) for protein production. Δ20MacA-His6 was purified by metal affinity chromatography using a HiTrap Ni\(^{2+}\) column (section 2.5.1). MacA-His6 was eluted in buffer containing 20 mM Tris-HCl pH 7.5, 500 mM NaCl, 10% glycerol, and 500 mM imidazole. The efficiency of expression and purification of MacA-His6 was determined by gel electrophoresis (section 2.6.1) (Figure 4.9b). The migration of the major protein band (lanes 2 to 10) is just above the 39 kDa SeeBlue protein marker (lane 1), which is consistent with a calculated molecular weight of 39.3 kDa for MacA-His6.
Figure 4.9  pET21d/Δ20macA construction. (a) Restriction enzyme digestion analysis of the pET21d/ΔmacA plasmid. Lanes 1-2 indicate the presence of the truncated macA gene. (b) SDS-PAGE of truncated MacA-His$_6$. Lanes 1-9 indicate the truncated Δ20MacA protein. Lane 10= the protein M, marker.
4.3.2 N-terminal truncated MacA regulates the ATPase activity of MacB

When N-terminal truncated MacA was added to MacB, at an equivalent concentration, a similar behavior was observed as in the presence of full length MacA, which is different as observed in the absence of MacB (Figure 4.4), indicating that truncated MacA was equally capable of regulating the ATPase activity of MacB (Figure 4.10). Thus, we concluded that the N-terminal transmembrane domain is not required in the regulation of the ATPase of MacB. This finding is consistent with a report on the membrane fusion protein AcrA, in which, a lipid-deficient derivative of AcrA was shown to retain its ability to cooperate with AcrB and TolC to confer resistance to drugs (Zgurskahya and Nikaido, 1999).

![Figure 4.10 Pi generation by MacAB and Δ20MacA plus MacB. 2 μM MacB and Δ20 MacA were used, and the ATPase activity was measured at Mg-ATP concentration of 1 mM.](image-url)
4.4 The α-helical hairpin of MacA is not required in the regulation of the ATPase activity of MacB

The measurements of the ATPase activity of *E. coli* MacB in the presence of *S. aureus* MacA was undertaken in collaboration with Mrs. Helen Frankish from our group.

Both gram-negative and gram-positive bacteria, such as *E. coli* and *S. aureus*, possess *macAB* genes encoding macrolide transporters; however, gram-positive bacteria would not be expected to require MacA to interact with an OMP due to the lack of an outer-membrane. We have noted that in some strains of *S. aureus*, MacA has a large deletion that corresponds to the coiled-coil hairpin which is the domain believed to interact with the outer membrane protein (Mikolosko *et al.*, 2006; Bavro *et al.*, 2008).

The genome sequence of *S. epidermidis* reveals three genes that putatively encode a macrolide efflux pump: SE2405, encoding a membrane translocase; SE2406, encoding an ATPase; and SE2407, encoding a membrane fusion protein. SE2405 and SE2406 have 46.5% and 55.4% sequence similarity, respectively, to the corresponding domains of *E. coli* MacB and SE2407 has 26% sequence similarity to *E. coli* MacA (Figure 4.11). Furthermore, secondary structure predications suggest that SE2407 and *E. coli* MacA are arranged into similar domains, consisting of an N-terminal transmembrane α-helix and a pair of long helices between two β-strand containing domains. *S. aureus* possesses three homologous genes, with SAS0173 and SAS0174 having 76.8% and 72.1% amino acid sequence similarity to SE2405 and SE2406, respectively. Interestingly, the alignment of SAS0172 with SE2407 indicates that the *S. aureus* MFP MacA encoded by SAS0172 has lost a large fragment, spanning residues 88-206 in the MFP encoded by SE2407 (Figure 4.11). This is the region that incorporates the hairpin that interacts with TolC, suggesting that *S. aureus* MacA has lost this domain because of absence of an outer membrane in gram-positive bacteria. However, *S. aureus* still retains the other domains of the MFP, strongly suggesting that this plays an important functional role, such as activating its cognate ABC transporter composed of SAS0173 and SAS0174.
Figure 4.11 **MacA sequence alignment.** A sequence alignment of the MacA membrane fusion proteins from the gram-negative bacterium *E. coli* and gram-positive bacteria *S. epidermitis* and *S. aureus* was represented. This analysis indicates that while the proteins from *E. coli* and *S. epidermitis* have similar secondary structures, the large α-helical domain predicted to harbor the TolC interaction site is missing in the *S. aureus* protein. The illustration of MacA sequence alignment was undertaken in collaboration with Dr. Vassiliy N. Bavro at the University of Cambridge.
In order to test the hypothesis, the ATPase activity of MacB was determined in the presence *S. aureus* MacA (SAS0172). The addition of an excess of *S. aureus* MacA to *E. coli* MacB was able to abolish the Pi burst and this kinetic behavior was consistent with the effect of *E. coli* MacA on MacB (Figure 4.12), suggesting that the α-helical hairpin of MacA is not required for activation of MacB. This finding is consistent with a report on AcrA (Elkins and Nikaido, 2003). In this report, several chimeric mutants of AcrA were created by replacing with the corresponding sequences from YhiU, a homolog does not function with the AcrB. Drug susceptibility tests of these chimeras revealed that N-terminal region of AcrA could be replaced with the corresponding domain of YhiU up to residue 290 and these mutants still retained functional as native AcrA. However, AcrA lost its ability to function with AcrB when the region between residues 290 and 357 was replaced with corresponding sequence of YhiU, indicating a small region of AcrA close to its C terminus is involved in the interaction with its cognate pump protein, AcrB.

![Figure 4.12](image)

**Figure 4.12** Pi generation by *E. coli* MacB in the presence of *S. aureus* MacA. 2 μM MacB was used, and the ATPase activity was measured at Mg-ATP concentration of 1 mM.
4.5 CD analysis

In the previous section, we predicted that *S. aureus* MacA encoded by *SAS0172* has lost the coiled coil hairpin structure found in *E. coli* MacA (Figure 4.11). In order to determine the secondary structure of the N-terminal truncated MacA from *E. coli*, CD spectrum analysis was applied. Protein samples were dialyzed against buffer containing NaF rather than NaCl, minimizing the strong absorbance from Cl\(^-\) in the far UV region. In order to estimate the best protein concentration for CD analysis, the absorbance of the sample was measured at in range of the wavelength (190 nm-260 nm), on the priority of keeping the strongest absorbance value within 0.8-1, generating a concentration for CD at around 0.09 mg/ml (2.25 μM). The spectrum of proteins in Tris buffer containing 20 mM NaF, pH 7.5 was recorded from 260 nm to 190 nm, for an average of six and subtracted by baseline spectrum of buffer itself to get the final spectrum. A quench and a peak can be seen at around 210 nm and 195 nm respectively (Figure 4.13). The High Tension voltage, the voltage applied to the photomultiplier, reached 700 when the wavelength went down to 195 nm during the measurement of the protein sample; however, this situation can also be observed in the measurements of buffer and H\(_2\)O respectively. Generally, for reliable data, the High Tension voltage should remain within 700 V, but this value will depend on the particular instrument being used (Kelly *et al.*, 2005). The reason could be due to the combination of using a weak Xe arc light source and the strong absorbance of H\(_2\)O in the far UV region.

![Figure 4.13 The CD Spectrum of Δ20MacA. Data was collected from 260 nm to 190 nm for a 2 mm pathlength, with a 2 sec rate constant, 10 nm/min scanning speed and 1 nm bandwidth.](image)
Figure 4.14  The secondary structure prediction of Δ20MacA. The data were analyzed by three algorithms (a) Contin-LL, (b) CDSSTR and (c) SELCON3.
Data was then input into the online server Dichroweb at Birbeck College, and analyzed by the algorithms SELCON3, CDSSTR and CONTIN-LL. The NRMSD (normalized root mean square deviation) values in the analyses generated from these three algorithms CONTIN-LL, CDSSTR, and SELCON3 are 0.105, 0.027 and 0.086 respectively (Figure 4.14). Generally speaking, a low value for the NRMSD is a necessary but not sufficient condition for concluding that an analysis has produced a good result (Brahms S and Brahms J, 1980). That is, if the NRMSD is high (> 0.1), the correspondence of the calculated secondary structure with the actual one is probably ‘incorrect’, but a low NRMSD does not necessarily indicate an accurate analysis. Among the three algorithms, CDSSTR gave the lowest NRMSD value; however, this algorithm tends, but not always, to give the lowest NRMSD values for a given reference database, but this does not mean that those results are the most significant. Apart from the NRMSD, the total sum of the secondary structure compositions must be ≥ 1. Based on these rules, the selected predicted compositions of α-helix were 18.7%, 9%, and 10.4% from CONTIN-LL, CDSSTR, and SELCON3 respectively (Figure 4.14). The membrane fusion protein AcrA in *E. coli* in its truncated form (Mikolosko *et al.*, 2006) has approximate 15% α-helix composition. Thus, our data is consistent with the prediction that the truncated MacA has the coiled coil hairpin structure.
4.6 Discussion

Membrane fusion proteins have been found to be essential components of tripartite pumps utilizing RND, MFS, and ABC transporters (Putman et al., 2000). The ATPase domain of MacB supplies a good approach to study the interaction of MacA with MacB. At first, MacA was originally extracted from membranes using DDM, however, our EM analysis and gel filtration data (data now shown) indicated it aggregated immediately after purification. Oddly, both MacB and MacA lost stability when extracted with DDM, but they both retained activity when extracted with Triton X-100.

Our kinetic data indicated that MacB had a maximal specific activity of 8.9 nmoles ATP/min/mg MacB and a $K_m$ of 374 μM Mg-ATP. During the course of our studies another investigation reported on the ATPase activity of MacB (Tikhonova et al., 2007), in which it was found that MacB retained ATPase activity in Triton X-100 but was inhibited by DDM. In this report, MacB, solubilized in 0.2% Triton X-100, appeared to have a specific activity that was an order of magnitude higher than we had found; however, a discontinuous assay was used to determine the ATPase activity and this could have been exaggerated by the initial Pi burst in the reaction. Indeed, $k_{cat}$ was marginally slower than the hydrolysis rate determined from the Pi burst phase in our experiments (e.g. 0.17 s$^{-1}$ versus 0.22 s$^{-1}$). In this report, they found that MacA could stimulate the ATPase activity of MacB when both of them were co-reconstituted into proteoliposomes; however, MacA had no substantial effect on the modulation of the ATPase activity of MacB in the detergent-solubilized state, which contrasts with our findings in that MacA increases the rate of product release ($k_{cat}$ from 0.011 to 0.015 s$^{-1}$), whilst decreasing the rate of hydrolysis to a similar rate to product release. The reason for this discrepancy could be because a discontinuous assay was used to determine the ATPase activity in the study undertaken by Tikhonova et al (Tikhonova et al., 2007), and the initial Pi burst in the reaction could have gone unnoticed. However, they did find that co-reconstitution of MacB with MacA had the effect of increasing both $k_{cat}$ ($k_{cat}$ from 0.1 to 0.78 s$^{-1}$) and the affinity of MacB for ATP ($K_m$ from 2.3 to 0.38 mM). This behaviour is similar to the effect of MacA on the steady-state kinetics of the ATPase activity of solubilized MacB in our study. To know if the reconstituted protein behaves in an identical manner to the solubilized protein would require the
reconstitution of sufficient amounts of MacB to define any Pi burst, which is technically demanding.

We found that the addition of a slightly sub-stoichiometric concentration of *E. coli* MacA to MacB was sufficient block the Pi burst catalysed by MacB, suggesting that the stoichiometry of binding of MacA:MacB is 1:2. This makes sense in the light of our finding that MacB forms dimers, which presumably bind a single MacA. Considering that TolC forms a trimer this would suggest that it binds at least three MacA molecules; implying that the minimal stoichiometry for the tripartite complex is a trimer of MacB dimers, each of which is connected to TolC via a single MacA (e.g. a MacA/MacB/TolC assembly of 3:6:3). Consistent with this proposal our AFM studies have identified particles larger than expected for a complex of a MacA monomer and MacB dimer (e.g. 307 nm$^3$) and with volumes sufficient to accommodate a trimer of MacB dimers and MacA trimer (e.g. > 900 nm$^3$).

The CD data is consistent with our prediction that MacA possess a coiled-coil hairpin structure. We predicted that *S. aureus* MacA has lost a large fragment, spanning residues 88-206 in the MFP encoded by *SAS0172*, and that this is the region that would incorporate the α-helical coiled-coil domain that interacts with the OMP (Lobedanz *et al.*, 2007). The addition of either *E. coli* MacA or *S. aureus* MacA can regulate the ATPase activity of MacB, suggesting that it is the β-strand domain of MacA, which is involved in the MacA-MacB interaction.
Chapter five: Tripartite pump: MacA, MacB and open state TolC

The membrane fusion protein AcrA, the RND transporter AcrB and the multifunctional outer membrane protein TolC can form a functional complex, and the latter can also function with MF and ABC transporters (Saier and Paulsen, 2001). Although the structure of an assembled tripartite complex structure has not been determined, the structure of each component has revealed how those components may assemble. The MacA/MacB/TolC complex was shown to confer resistance to macrolides in *E. coli* (Kobayashi *et al.*, 2001); in this study, our aim was to investigate how the complex is assembled and functions.

5.1 A functional tripartite complex

To evaluate the function of the tripartite complex, we adopted measurements of the growth of cells expressing various combinations of the *macA*, *macB* and *tolC* genes. The expression level of wild type TolC was not stable so the plasmid encoding the open-state TolC, which has a deletion of the last 43 amino acids and two mutation points, Y362F and R367E, was a courtesy from the lab of Dr. Ben Luisi at the University of Cambridge (Bavro *et al.*, 2008), has been used in this experiment. This served as the template for PCR amplifications in all TolC constructs.
5.1.1 Constructions

Constructs were made in a similar manner as described in section 3.1. In short, the chromosomal macB region of *E. coli* Novablue was amplified using PCR and cloned into EcoRI and SalI restriction sites in the first cloning site of pETDuet™-1 to yield the pETDuet™-1/macB plasmid. The amplified tolC gene was then cloned into NdeI and XhoI restriction sites in the second cloning site of the pETDuet™-1/macB plasmid to yield the pETDuet™-1/macB-tolC plasmid (Figure 5.1a). A Shine-Dalgarno (SD) sequence was added upstream of the full length macA gene during PCR amplification. The amplified SD-macA gene was cloned into the SalI and NotI sites in the first cloning site of the pETDuet™-1/macB and the pETDuet™-1/macB-tolC plasmids to yield pETDuet™-1/macB-macA and pETDuet™-1/macB-macA-tolC plasmids, respectively (Figure 5.1b). In order to test whether the N-terminus of MacA is required for the functional assembly of the complex, the plasmid pETDuet™-1/macB-SS-Δ20macA-tolC plasmid was constructed in a similar manner but with a slight modification. The signal sequence gIII from the vector pBAD/gIII was fused to the 5' end of the SD-Δ20macA sequence in order to lead truncated MacA to the periplasm (Figure 5.1c).
Figure 5.1 Constructions for drug susceptibility tests. (a) Restriction enzyme digestion analysis of pETDuet™-1/macB-tolC plasmid. Lane 1 = the DNA ladder. Lanes 2-3 indicate the presence of the tolC gene. Lanes 4-5 indicate the presence of the macB gene. (b) Restriction enzyme digestion analysis of pETDuet™-1/macB-macA-tolC and pETDuet™-1/macB-D20macA-tolC. Lane 1 = the DNA ladder. Lanes 2-3 indicate the presence of the macA gene. Lanes 4-5 indicate the presence of the Δ20macA gene. (c) Restriction enzyme digestion analysis of pETDuet™-1/macB-gIII-Δ20macA-tolC. Lane 1 = the DNA ladder. Lane 2 indicates the presence of the gIII-Δ20macA gene.
5.1.2 Drug susceptibility test

Plasmids pETDuet$^{TM}$-1/macB, pETDuet$^{TM}$-1/macB-macA, pETDuet$^{TM}$-1/macB-tolC, pETDuet$^{TM}$-1/macB-macA-tolC and pETDuet$^{TM}$-1/macB-D20macA-tolC were transformed into the *E. coli* ΔacrAB strain Kam3(DE3), which is highly susceptible to a range of antibiotics (Morita *et al.*, 1998). The λDE3 lysogenisation of Kam3 was described in section 2.4.5.

The *macA*, *macB* and *tolC* genes were simultaneously expressed from pETDuet$^{TM}$-1 in the Kam3(DE3) strain, conferring elevated resistance to the macrolide erythromycin compared to the control Kam(DE3), harboring only the pETDuet$^{TM}$-1 vector, indicative of the formation of a functional complex (Figure 5.2).

Our previous data in section 4.3 suggested that the N-terminal transmembrane domain of MacA is not required in the regulation of the ATPase activity of MacB. In order to test whether the N-terminus of MacA is required for the functional assembly of the complex. A construct in which the gIII signal sequence was fused to truncated MacA, targetting it to the periplasm, was capable of conferring resistance to erythromycin (Figure 5.2), indicating that the N-terminal transmembrane domain is not essential for the assembly of the functional complex. This finding is consistent with two reports on membrane fusion proteins. Yoneyama *et al* (Yoneyama *et al.*, 2000) substituted cysteine 24 with phenylalanine or tyrosine and showed this mutant soluble MexA, without the lipid moiety, retained its function in the MexA/MexB/OprM complex pumps, retaining the antibiotic susceptibility. Membrane fusion protein AcrA, when being truncated the first 25 residues, still functional active in the AcrA/AcrB/TolC complex pumps, conferring erythromycin and novobiocin resistance in the ΔacrA deletion *E. coli* strain N43 (Zgurskaya and Nikaido, 1999).

Cells expressing *macB* with *tolC* conferred modest resistance to erythromycin in comparison to cells expressing *macB*, *tolC* and *macA*, indicating that MacA is required to couple MacB to TolC (Figure 5.2).
MacB alone conferred elevated resistance to erythromycin, probably due to its ability to pump the antibiotic into the periplasm, but we consistently found that expressing macB with either tolC or macA conferred greater resistance. Consequently, we sought to test if MacA could enhance this ability. However, it is difficult to distinguish the slight difference in resistance since cells harboring different plasmids tend to grow at different rates. Consequently, we adopted an alternative procedure of monitoring the growth of cells in the presence and absence of erythromycin and determining the loss in growth due to the antibiotic. This procedure enable us to determine the susceptibility of the MacB and MacAB expressing cells to erythromycin, revealing a more significant loss in growth of the MacB expressing cells compared to the MacAB expressing cells and indicating that the simultaneous expression of MacA and MacB increases the resistance of the cells to erythromycin (Figure 5.3). This data suggests that MacA enhances the ability of MacB to confer antibiotic resistance. Similarly, a previous study reported that MacAB, but not MacB alone, conferred resistance to macrolides (Kobayashi et al., 2001).
Figure 5.3 **Drug susceptibility histogram.** The cell growth of Kam3 cells harboring plasmids were measured at absorbance 600 nm up to 3 hours. Relative growth (%) was obtained by dividing the growth of cells in the presence of 50 μg/ml erythromycin by the growth of cells in the absence of erythromycin. MacAB represents the cells expressing MacAB proteins; MacB represents the cells expressing MacB protein; Control represents the cells harboring empty vector; and Wild represents the wild Kam3(DE3) *E. coli* strain.

### 5.2 MacA enhanced the erythromycin binding to MacB

The equilibrium binding of erythromycin experiments were undertaken in collaboration with Dr. Saroj K. Velamakanni at the University of Cambridge.

From the previous section, we concluded that MacA, MacB and TolC can form a functional tripartite pump, conferring *E. coli* resistance to erythromycin. Furthermore, the cells harboring the plasmid encoding MacAB showed greater resistance than the plasmid encoding MacB itself. Our kinetics data has indicated that MacA regulates MacB ATPase activity; in this section, we detected the binding of [N-methyl-\(^{14}\)C]-erythromycin to purified MacB in the presence and absence of MacA (section 2.7.2). The experiments were divided into three groups containing 50 μg MacA, 50 μg MacB, and 25 μg MacB with 25 μg MacA respectively, where equal amount of protein was in each group. Interestingly, our data indicated that MacB bound more erythromycin in the presence of MacA, in a concentration dependent manner (Figure 5.4). However, it was not possible to increase the erythromycin to a concentration that saturated MacB to determine affinity due to erythromycin high hydrophobicity, leading erythromycin
to precipitate with proteins at higher concentration. The data indicate that MacA not only modulates the ATPase activity of MacB but also enhances its capacity to bind erythromycin.

![Histogram of Erythromycin Equilibrium Binding](image)

**Figure 5.4** Histogram of erythromycin equilibrium binding to MacA, MacB and MacAB proteins. The bars represent the erythromycin bound by MacA (left, black), MacB (middle, light grey) and MacAB (right, dark grey).

### 5.3 *in vitro* Crosslinking of MacA-MacB

The protein crosslinking experiments and data representation were undertaken in collaboration with Dr. Vassiliy N. Bavro at the University of Cambridge.

Our drug susceptibility tests indicated that MacA, MacB and TolC can form a functional complex pump *in vivo*; whilst our kinetics data indicate that purified MacA can regulate the ATPase activity of purified MacB in a detergent solubilized state. In this section, protein-protein crosslinking was adopted to gain more evidence that MacB can interact with MacA *in vitro*. Purified MacB and MacA were incubated with the crosslinker DMA and DMS for 1 hr at room temperature. Interestingly, additional bands (Lane 5, 6, 9 and 10) appeared with an Mr greater than the MacB monomer, and the one in the lane 5 was cut out along with the control bands in lane 3 for MALDI (Matrix-Assisted Laser Desorption/Ionization) analysis. MALDI in gel-digestion indicated that both MacA and MacB proteins were present in the cross-linked band (Figure 5.6); a miniscule amount of MacA could only be detected in the control band. This data suggested that MacB can interact directly with MacA.
Figure 5.5  SDS-PAGE of the crosslinked MacA-MacB complexes by using DMA and DMS. MacB-MacA complexes are framed in red boxes. Lane 1= MacA, Lane 2= MacB, Lane 3= MacA + 5 mM DMA, Lane 4= MacB + 5 mM DMA, Lane 5= MacA + MacB + 5 mM DMA incubated for 1 h, Lane 6= MacB + 5 mM DMA to be incubated for 30 min + MacA for another 30 min, Lane 7= MacA + 5 mM DMS, Lane 8= MacB + 5 mM DMS, Lane 9= MacA + MacB + 5 mM DMS incubated for 1 h, Lane 10= MacB + 5 mM DMS incubated for 30 min + MacA for another 30 min.

Figure 5.6  MALDI analyses of the MacA-MacB complex fragment. Fragments positively identified by MALDI are colored in red. (a) MacB protein has 42% sequence coverage of the peptide map. (b) MacA protein has 37% sequence coverage of the peptide map.
5.4 *in vitro* interaction between components in the complex

Interactions between *E. coli* MacA, MacB and TolC were tested for using detergent solubilized proteins for pull down assays. The strategy for the pull-down assay involved using a bait protein and a prey protein (section 2.7.3). MacB-His<sub>6</sub> served as the bait protein in the detection of MacA-MacB and MacB-TolC interaction, and TolC-His<sub>6</sub> served as the bait protein in the detection of TolC-MacA interaction.

5.4.1 Construction and overexpression of the bait proteins

Purified MacB-His<sub>6</sub> protein was overexpressed and purified as previously described (section 3.1).

pET21a/tolC plasmid was constructed in a similar manner, to that described in section 3.1; and relevant primers, vector plasmids and *E. coli* strains can be found in section 2.3 and 2.4. In short, the amplified *tolC* gene was cloned into the *NdeI* and *XhoI* restriction sites of the pET21a(+) plasmid to yield the pET21a/tolC plasmid (Figure 5.7a). The pET21a/tolC expression construct was then transformed into the *E. coli* strain C43(DE3) for protein production. The protein expression and cell membranes harvest were manipulated as described in section 2.5. The cell membranes were dissolved in 20 mM Tris buffer pH 7.5, 300 mM NaCl, 10 mM imidazole, 1 mM THP, 10% glycerol and 5% Triton X-100 on a rotary shaker overnight.

TolC-His<sub>6</sub> was purified by metal affinity chromatography using a HiTrap Ni<sup>2+</sup> column (section 2.5.1), and eluted in buffer containing 20 mM Tris-HCl pH 7.5, 500 mM NaCl, 10% glycerol, 500 mM imidazole and 0.2% Triton X-100. The yield and purify of the TolC-His<sub>6</sub> was determined by gel electrophoresis (section 2.6.1) (Figure 5.7b). The migration of the major protein band (lanes 2 to 6) is just above the 51 kDa protein marker band (lane 1), which is consistent with the Mr of the TolC-His<sub>6</sub>, with a calculated molecular weight of 49.2 kDa.
Figure 5.7  **pET21a/tolC construction and overexpression** (a) Restriction enzyme digestion analysis of pET21a/tolC plasmids. Lane 1= the DNA ladder. Lanes 2-3 indicate the presence of the *tolC* gene. (b) SDS-PAGE of TolC-His₆. Lane 1= the protein Mᵣ marker. Lanes 2-6 indicate the TolC-His₆ proteins.
5.4.2 Construction of the prey proteins

Constructs were made in a similar manner to those described in the previous section. In short, the amplified Δ20macA, macA and tolC genes was cloned individually into the NcoI and XhoI restriction sites of the pET45b(+) plasmid to yield the pET45b/Δ20macA, pET45b/macA and pET45b/tolC plasmids (Figure 5.8a). Each of those expression constructs was transformed into the E. coli strain C43(DE3) for membrane slurry production.

Figure 5.8 S-tagged protein construction for pull-down assay. (a) Restriction enzyme digestion analysis of pET45b/macA and pET45b/Δ20macA plasmid. Lane 1= the DNA ladder. Lanes 2-3 indicate the presence of the Δ20macA gene. Lanes 4-5 indicate the presence of the macA gene. (b) Restriction enzyme digestion analysis of the pET45b/tolC plasmid. Lane 3= the DNA ladder. Lanes 1-2 indicate the presence of the tolC gene.
5.4.3 Pull down Assay

In the AcrA/AcrB/TolC tripartite complex pump, calorimetry demonstrated energetically favourable interactions of AcrA with both purified AcrB and TolC proteins (Touze et al., 2004). We previously concluded that MacA, MacB and TolC form a functional complex on the basis of our drug susceptibility test data and our findings indicate that MacB directly interacts with MacA in crosslinking studies.

Using pull-down assay (section 2.7.3), in which MacB-His₆, served as the bait protein, we were able to capture the prey protein MacA-Stag (Figure 5.9). Membrane pellets containing MacA-Stag was dissolved in Triton X-100 and added to purified MacB-His₆. The mixture was loaded into a Ni²⁺ charged chelating column to immobilize the MacB-His₆, along with MacA-Stag. Unbound MacA-Stag can be seen in the flow-through (Figure 5.9 lane 6); the column was then given highly stringent wash to eliminate unbound MacA-Stag. A fraction was collected before the elution of the MacB-His₆-MacA-Stag complex and only a very tiny amount of MacA-Stag could be detected, indicating little or no unbound MacA-Stag suspending in the column (Figure 5.9 lane 5). The MacB-His₆ and MacA-Stag can be clearly observed in the elution (Figure 5.9 lane 4), revealing MacA could interact with MacB. No MacA-Stag was seen in the elution of the control group (Figure 5.9 lane 1), suggesting that there was no non-specific interaction between MacA and the column.

Using MacB-His₆ and N-terminal truncated MacA-Stag for the pull-down assay, Δ20MacA can be observed in the elution (Figure 5.10), indicating that there is interaction between the periplasmic domain of MacA and MacB. This behaviour is consistent with our previous data that indicated that truncated MacA can regulate the ATPase activity of MacB. Similarly, using TolC-His₆ and MacA-Stag, a strong interaction can be detected (Figure 5.11). Using MacB-His₆ and TolC-Stag, an interaction can be observed (Figure 5.12), but this interaction is apparently a lot weaker than those between MacA and MacB/TolC.
Figure 5.9  **MacA-Stag pulled down by MacB-His$_6$.** Lane 1 = elution, Lane 2 = wash, Lane 3 = flow-through (Lanes 1-3 were collected in the absence of the MacB-His$_6$ protein). Lane 4 = elution, Lane 5 = wash, Lane 6 = FT (Lanes 4-6 were collected in the presence of the MacB-His$_6$ protein. Lane 7 = the MacB-His$_6$ protein and Lane 8 = the protein M, marker. The identity of the MacA-Stag protein was confirmed by Western blot using S-Tag monoclonal antibody.
Figure 5.10  **Δ20MacA-Stag pulled down by MacB-His$_6$.** FT represents flow-through, W1 represents early stage of wash, W2 represents later stage of wash, and E represents elution. “+” indicates the presence of the protein, whereas “-” indicates the absence of the protein. In the film, the prey protein was detected by Western blot using S-Tag monoclonal antibody.

Figure 5.11  **MacA-Stag pulled down by TolC-His$_6$.** FT represents flow-through, W1 represents early stage of wash, and E represents elution. “+” indicates the presence of the protein, whereas “-” indicates the absence of the protein. In the film, the prey protein was detected by Western blot using S-Tag monoclonal antibody.

Figure 5.12  **TolC-Stag pulled down by MacB-His$_6$.** FT represents flow-through, W2 represents later stage of wash, and E represents elution. “+” indicates the presence of the protein, whereas “-” indicates the absence of the protein. In the film, the prey protein was detected by Western blot using S-Tag monoclonal antibody.
5.5 Discussion

Gram-negative bacteria possess tripartite pumps composed of an inner membrane protein, a periplasmic protein and an outer membrane protein, which work together to confer multidrug resistance. In order to determine the function of the MacA/MacB/TolC tripartite-complex pump, we undertook growth curve measurement to determine the drug susceptibility of the strains expressing different combinations of the MacA, MacB and TolC proteins.

Originally, minimum inhibitory concentration tests (MIC) were adopted to determine whether the MacA/MacB/TolC complex pump confers resistance to the macrolide erythromycin in bacteria. However, *E. coli* strain Kam3(DE3) harboring plasmids encoding MacA, MacB and TolC didn't have a substantial effect on the resistance of the cells to erythromycin (data not shown). It could be that although the proteins assembled together to translocate drugs successfully, the high toxicity of the three proteins had a detrimental effect on the host cells, leading to the insignificance of drug resistance. As an alternative, growth curve measurements were undertaken to determine the function of the tripartite complex pump. During the grow curve measurements, the tubes were put in an incubator with shaking to enable more O₂ to be dissolved into the media, allowing the host cells to grow more vigorously. The effect was that in the growth curve measurements, the minimum concentration that inhibited growth of erythromycin is a lot greater than the one in the MIC tests (e.g. 300 μM vs 16 μM erythromycin).

Our drug susceptibility tests revealed that MacB can confer elevated resistance in bacteria to erythromycin in the presence of MacA (Figure 5.3) and our equilibrium erythromycin binding experiments indicated that MacA increases the binding ability of MacB to erythromycin (Figure 5.4); besides, our kinetics data suggested that MacA increased the rate of product release from MacB, speeding up the turn over rate. We speculated that MacA increases the binding ability of MacB for erythromycin to catch up the faster conformation switch between the outward-facing and inward-facing conformation, speeding up the transportation of drugs.
In the AcrA/AcrB/TolC complex pump, AcrA can be crosslinked with AcrB and TolC individually in vivo, and microcalorimetry studies demonstrated energetically favourable interactions of AcrA with both purified AcrB and TolC in vitro (Touze et al., 2004). Our pull-down data indicated that any two components of the MacA/MacB/TolC complex pump can interact directly in the absence of the third component. However, the intensity of the band in Figures 5.10, 5.11 and 5.12 suggests that the interaction between MacB and TolC is weak compared to those between MacA and MacB/TolC. This may suggest that the dimerization of MacB can recruit TolC, via MacA, and simultaneous open TolC for translocation.
Chapter six: Crystallization trials

Integral membrane proteins such as transporters and receptors are critical components of many fundamental biological processes. Furthermore, integral membrane proteins are important in biomedical and biotechnological applications; whilst the majority of drug targets are integral membrane proteins. The macrolide ABC transporter MacB putatively has an exceptional structure comprising of four transmembrane helices, which is drastically different to the topology of the structure determined for ABC exporters Sav1866 and MsbA, which has six transmembrane helices, in bacteria. In this chapter, I report on how the MacB protein was to crystallization screens in order to acquire crystal for structure determination.

6.1 Dynamic light scattering of MacB

Purified MacB in buffer containing 20 mM Tris-HCl pH 7.5, 50 mM NaCl, 0.05% DDM and 10% glycerol was analyzed by dynamic light scattering to determine the size distribution profile of MacB. The DLS profiles of 1 mg/ml MacB and the buffer itself were measured at 16°C and 4°C. A good correlation between the traces suggested that measurements and data are reliable, and MacB showed its high homogeneity (> 97%) at both 16°C and 4°C (Figure 6.1ab), indicating that it is monodispersed in solution. The analysis of the buffer itself served as the control, compared to MacB, eliminating the chance of a false signal from the buffer (Figure 6.1c).
Figure 6.1  **Size distribution analysis of MacB.** The DLS profile of MacB was measured at 16°C (a) and at 4°C (b), and the DLS profile of the buffer was measured at 4°C (c).
6.2 Preliminary sparse matrix screen of MacB

The protein crystallization and data collection were undertaken in collaboration with Dr. Vassiliy N. Bavro and Mr. Zbigniew Pietras at the University of Cambridge.

The \textit{macB} gene sequence was cloned on to the pET28a(+) plasmid vector, which introduces a thrombin digestion site between the N-terminal His tag and the cloning site, making it possible to make tag-free MacB proteins for crystallization trials. However, after being treated with thrombin enzyme (Novagen), a small proportion of MacB still retained the tags, detected by Western blot (section 2.6.3), causing heterogeneity in the protein prep (data not shown). To avoid heterogeneity, the tag and the cleavage sequence were left on the N-terminal on MacB when it was crystallized.

Purified MacB in Triton X-100 (0.2%) (section 3.1) was exchanged and dialyzed against buffer containing 20 mM Tris-HCl pH 7.5, 50 mM NaCl, 0.05% DDM and with or without 10% glycerol before being used in crystallization trials. Preliminary crystallization trials of MacB were accomplished using sparse matrix screens to discover a suitable starting condition.

The commercial sparse matrix screens used were SMI, MbClass, MbClassII, Classics, Classics light, pHClear II (Qiagen), MembFac (Hampton Research), and MemSys (Molecular Dimensions). Crystallization trials were undertaken using 10 mg/ml MacB in the vapour diffusion sitting drop method on 96 well plates with round bottoms and automated screening by a Cartesian robot. The development was monitored intermittently by light microscopy.

As shown in Figure 6.2, a few putative micro-crystals could be observed in five conditions, but only one of them was from non-glycerol MacB samples (#57 Classics light), suggesting that MacB is more stable in the presence of glycerol. Among the five conditions, 28-30% PEG400 is found in three conditions (Classics 74 and 75, and MbClassII 39), suggesting that precipitant PEG 400 is suitable for MacB crystalization; however, no specific salt or pH value is preferable for the formation of MacB crystals (Table 6.1).

Having identified a promising starting condition, the condition can be further optimised to improve crystal quality through adjustments in precipitant and salt concentration, and pH ranges (Delucas \textit{et al.}, 2005).
Figure 6.2 Photographs of putative MacB micro-crystals taken under light microscopy. Micro-crystals indicative of initial microcrystal development identified from (a) #74 Classics (b) #75 Classic (c) #57 Classics light (d) #39 MbClassII (e) #62 SM1 screen kits.
Table 6.1 Compositions of conditions yielding putative micro-crystals.

<table>
<thead>
<tr>
<th>Kit</th>
<th>Salt</th>
<th>Buffer</th>
<th>Precipitant</th>
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<tbody>
<tr>
<td>Classics 74</td>
<td>0.2 M Calcium chloride</td>
<td>0.1 M HEPES sodium salt pH 7.5</td>
<td>28 % (v/v) PEG 400</td>
</tr>
<tr>
<td>Classics 75</td>
<td>0.1 M Cadmium chloride</td>
<td>0.1 M Sodium acetate pH 4.6</td>
<td>30 % (v/v) PEG 400</td>
</tr>
<tr>
<td>Classics Light 57</td>
<td>0.5 M Ammonium sulfate</td>
<td>0.1 M tri-Sodium citrate pH 5.6</td>
<td>0.5 M Lithium sulfate</td>
</tr>
<tr>
<td>MbClassII 39</td>
<td>0.1 M Magnesium chloride</td>
<td>0.1 M Sodium acetate pH 4.6</td>
<td>30 % (v/v) PEG 400</td>
</tr>
<tr>
<td>SM1 62</td>
<td>--</td>
<td>0.1 M sodium acetate anhydrous pH 4.5</td>
<td>20% (w/v) PEG 3000</td>
</tr>
</tbody>
</table>
6.3 Preliminary sparse matrix screen of MacA/MacB/TolC complex

The protein crystallization and data collection were undertaken in collaboration with Dr. Vassiliy N. Bavro and Mr. Zbigniew Pietras at the University of Cambridge.

Purified MacA in Triton X-100 (0.2%) (section 4.1) and MacB (section 3.1) in Triton X-100 (0.2%) and open state TolC (supplied by Dr. Bavro) were exchanged and dialyzed against buffer containing 20mM Tris-HCl pH 7.5, 150 mM NaCl, 0.05% DDM and with 10% glycerol. 40 μL of MacA (3 mg/ml), 60 μL of MacB (8 mg/ml) and 20 μL of TolC (10 mg/ml) were mixed together to make a final concentration of around 6.5 mg/ml, allowing the Molar ratio of MacA: MacB: TolC = 1: 2: 2. Preliminary crystallization trials of the MaA/MacB/TolC complex were accomplished using sparse matrix screens to discover a suitable starting condition.

The commercial sparse matrix screens used were SMI, MbClass, and MbClassII, Classics (Qiagen). Crystallization trials were undertaken using the mixture of the proteins in the vapour diffusion sitting drop method on 96 well plates with round bottoms and automated screening by a Cartesian robot. The development was monitored intermittent by light microscopy.

As shown in Figure 6.3a, a micro-crystal could be observed in the SMI 66 condition, 0.2 M MgCl₂, 0.1 M Tris pH 7.0, and 10% (w/v) PEG 8000. The crystal was frozen using cryo-agent 30% PEG 400 and taken to the European Synchrotron Radiation Facility (ESRF) to test for X-ray diffraction. The crystal generated a low diffraction of 20 Å in the lattice (Figure 6.3). Unfortunately, there was a technical problem to get a definitive result indicating either single protein or the protein complex within the crystal since the crystal was too small to tell.

The optimizations of the crystal condition of the complex proteins were undertaken by a gradient of PEG 8000 (5.5-13%, with an interval of 1.5%) versus pH (6.5-8, with an interval of 0.5) and of PEG 8000 (5-15%, with an interval of 2%) versus MgSO₄ (50-200 mM, with an interval of 50 mM). However, no crystal was observed in this optimization. The possible reason could be that we used proteins for crystallization from batch to batch, causing the crystal not reproducible.
Figure 6.3  Images of the crystal of the MacA/MacB/TolC complex. (a) The image of the MacA/MacB/TolC crystal in the well of SM1 66. (b) The image of the loop and the crystal during scanning. (c) The complex diffraction image, with a diffraction of 20 Å limit.
6.4 Discussion

Membrane protein crystallography is a challenge. It is reasonable to state that from expression, purification to crystallization and structure determination of integral membrane proteins still remain non-trivial endeavors that require substantial time and resources (Caffrey, 2003; Wiener et al., 2004). Even more, diffraction-quality crystals are particularly difficult to get when a membrane protein source is used. The reason could be current limited ability to manipulate proteins bearing hydrophobic/amphiphilic surfaces that are usually enveloped with membrane lipid. For this reason there are only around 160 structures for unique membrane proteins (White, 2008).

MacB retained ATPase activity when detergent solubilized but this was detergent dependent, being active in Triton X-100 but not in high DDM. MacB was first extracted using 2% DDM but didn’t retain its ATPase activity under these conditions. However, interestingly, MacB remained functional when extracted with 5% Triton X-100 and then exchanged into low DDM concentrations (e.g. 0.05%). The reason why MacB does not remain functional when extracted by DDM is unknown. MacB went through two steps of purification before the crystallization trials; after which a DLS analysis showed that the MacB samples had reached 97% homogeneity. Figure 6.2 shows some putative micro-crystals of MacB but the majority of them do not have angles, indicating 10 mg/ml of MacB may be too high, resulting in shortage of time for crystal formation. Besides, MacB was extracted by 5% Triton X-100 to obtain the best yield, but excess detergent may result in excess delipidation, causing protein instability. (Boulter and Wang, 2001).

Another hindrance in MacB crystallization is that the tag and the cleavage sequence were left on the N-terminal on MacB when it was crystallized. This could be surmounted by taking a further step purification for affinity chromatography to isolate the tagged and tag-free MacB proteins. However, too many purification steps may lead to protein delipidation. Another suggestion is to put the His tag onto the C-terminal of
MacB; since the NBD is on the N-terminal of MacB, we speculated that a C-terminal tag on MacB may avoid a disordered hydrophilic contact.

Structure determination of ABC transporters has been challenging due to their dynamic nature in a detergent solubilized state, which often hinders formation of well-ordered, three-dimensional crystals. The addition of AMP-PNP may be helpful to fix the protein flexibility. Since MacB ATPase activity is detergent dependent, another approach to think about is to exchange into polyoxyethylene glycol detergents such as C_{12}E_8, which has a similar structure to Triton X-100 but without a benzene ring. Sav1866 and ModB_{2C_{2A}} were both crystallized successfully in the presence of the detergent C_{12}E_8.
Chapter seven: Final discussion

Tripartite pumps in gram-negative bacteria can facilitate the extrusion of a wide variety of substrates from the cell. There three components assembled in the tripartite pumps are as follows: an inner membrane protein (IMP), an outer membrane protein (OMP) and a membrane fusion protein (MFP), making it possible to translocate substrates across both the inner and outer membranes. Interestingly, gram-positive bacteria that lack an outer membrane are also found to possess MFPs, suggesting an additional function on top of serving as a scaffold to stabilize the pump assembly. Murakami proposed that conformational changes in AcrB can be transmitted via AcrA to TolC, suggesting AcrA may play a role in addition to stabilize the interaction between AcrB and TolC (Mikolosko et al., 2006). In order to verify this assumption, we chose the MacA/MacB/TolC tripartite complex pump in E. coli (Kobayashi et al., 2001) since there are clearly homologous systems in gram-positive bacteria. The MFP MacA not only stabilizes the tripartite-assembly by interacting with both the IMP MacB and the OMP TolC but also has a role in regulating the function of MacB, apparently increasing its affinity for both erythromycin and ATP. Moreover, a better understanding of this complex will give us a better insight into type I secretion systems in bacteria, such as HlyB/HlyD/TolC, where HlyB is an ABC transporter (Thanabalu et al., 1998).

Generally, the MFP is anchored to the inner membrane by either an α-helix or a lipid moiety, attached to a cysteine residue, located towards the N-terminus of the protein (Tikhonova et al., 2004). This might suggest that there is only a requirement to anchor the MFP to the inner membrane, rather than a need for specific interactions between the IMP and MFP. Using pull-down assay, we showed that the anchor, N-terminal α-helix, of E. coli MacA was not required for complex formation with either MacB or TolC. This is consistent with a previous report that N-terminal truncated MacA could interact with MacB (Tikhonova et al., 2007). However, using drug susceptibility tests, we showed that the transmembrane helix of MacA was required to target MacA to the periplasm, where the periplasmic domain of MacA could interact...
with the periplasmic domains of MacB and TolC to form a functional tripartite complex. This is consistent with a report about the MFP AcrA in *E. coli*, which concluded that the truncated lipid-deficient AcrA derivative was as functional as full length AcrA, as judged by the resistance of the cells to erythromycin and novobiocin (Zgurskaya et al., 1999). However, this behaviour is different from another report that the deletion of the N-terminal α-helix of MacA, whilst not blocking its interaction with MacB, yielded a derivative that when expressed with MacB didn’t confer elevated resistance to erythromycin (Tikhonova et al., 2007). The exact reason for the failure of this study to detect the elevated resistance of cells expressing N-terminal truncated MacA is unknown. One possibility is that we used different truncations; in our study we used Δ20MacA whilst Tikhonova *et al* used a Δ31MacA; alternatively maybe the Δ31MacA fused to the OmpA signal sequence failed to reach the periplasm?

The MFP component of the tripartite efflux systems of gram-negative bacteria is believed to act as a bridge between the outer and the inner membranes proteins. The MFP AcrA incorporates a pair of α-helices that form a coiled-coil, sandwiched between two β-strand domains, which have been shown to interact with the α-helical barrel of the OMP TolC (Lobedanz *et al*., 2007; Bavro *et al*., 2008). Furthermore by domain swapping experiments showed the β-strand domains are crucial for the interaction with AcrB, but the exact role of these domains remains obscure (Elkins and Nikaido, 2003). Here, we provided evidence that the β-strand domain activates the MacB ATPase. When excess *S. aureus* MacA was added to *E. coli* MacB, no phosphate burst was observed but the steady-state rate was enhanced (Figure 4.12), in direct analogy with the effect of *E. coli* MacA on *E. coli* MacB, indicating that this interaction is via the β-strand domains of the MFP MacA. We established that cells expressing only MacB were less resilient, in terms of their growth, than those expressing both MacA and B, when exposed to erythromycin, suggesting that the latter are more efficient in extruding the antibiotic (Figure 5.3). This may have acted as a selective pressure for gram-positive bacteria to retain the MFP. In the case of *S. aureus*, we speculate that there has been a selective pressure to retain MacA, but because there is no OMP with which it would interact, there has been no pressure to retain the coiled-coil domain that has been deleted from MacA. Recently, the role of the MFP
BesA in the function of the BesABC complex pump has been reported (Bunikis et al., 2008). BesC, a TolC homologue, interacts with the putative inner membrane transporter BesB, an AcrB homologue, and the AcrA homologue BesA, a periplasmic adaptor protein. Interestingly, in this system BesA lacks the α-helical hairpin. It could be that the interaction between IMP BesB and OMP BesC is sufficient to trigger and stabilize the opening of BesC. The reason for the retention of BesA in the BesABC system is most likely associated with its function in the activation of the inner-membrane component, BesB, which could explain the preservation of the hairpin-lacking MFPs in gram-positive bacteria, such as S. aureus.

MacB has a nucleotide-binding domain that incorporates Walker A and B motifs and an ABC signature sequence, which are characteristic features of members of the ABC superfamily. Biochemical and structural data suggested the conserved motifs are involved in the binding and hydrolysis of ATP, which are believed to be the key steps in the transport mechanism of ABC transporters. The binding of ATP to both soluble ATPase subunits (Chen et al., 2003) and to the NBDs within a complete ABC transporter (Dawson and Locher, 2007) have been shown to promote dimerisation as the ATP is bound at the interface of these nucleotide binding-sites, sandwiched between the Walker A motif of one NBD and the ABC signature motif of the other NBD. The ATP occlusion model in human p-gp, different from the ATP switch model as described previously, indicated that ATP-driven NBD dimerization was followed by a rapid, temperature-dependent occlusion of ATP at one of the ATP sites, which showed an asymmetry. The latter step is actually energetically driven, involves relatively large conformation changes, and may drive the conformational change in the TMDs (Sauna and Ambudkar, 2007).

In this chapter, using SEC, crosslinking, AUC, ES-Mass spectrometry, and AFM, MacB was proven to be dimeric in a DDM solubilized state. Although MacB has an exceptional four transmembrane helices, different from the six helices characteristics of ABC exporters such as Sav1866 and MsbA, its ability to form dimer suggests that MacB operates by a similar mechanism to ABC transporters.

Our AFM studies indicated that there are a higher proportion of MacB dimers in the presence of the non-hydrolysable nucleotide AMP-PNP. A proposed mechanism of
complete ABC transporters suggested that the binding of ATP triggers the transporter to adopt a conformation in which the substrate binding site is outward-facing, because ATP bridges the two NBDs, closing off the inward-facing substrate binding site (Hollenstein et al., 2007). Conversely, the release of the hydrolysis products ADP and phosphate promotes an inward-facing conformation, as the structural constraint imposed by binding of the nucleotide is released. Consistent with this proposal, several studies indicate that NBD dimerisation cannot be induced by ADP (Zaitseva et al., 2006), because interactions between the γ-phosphate of ATP and the signature sequence catalyse these events (Lu et al., 2005; Oloo et al., 2006). Our kinetic data indicated that MacA can increase the affinity of MacB for ATP and decrease the rate of ATP hydrolysis in the initial stage, so that MacA actually helps to stimulate and stabilize the ATP binding conformation of MacB, presumably the outward-facing conformation. In this manner, MacA could play a direct role in driving antibiotic translocation between MacB and TolC. Our equilibrium erythromycin binding data pointed out that MacB bound more erythromycin in the presence of MacA and that the amount bound increased in a concentration dependent manner, suggesting that MacA not only modulate the ATPase activity of MacB but also its capacity to bind drugs.

Our studies established that the conformational changes of MacB can be propagated from the periplasmic domain of MacA to the cytoplasmic NBD of MacB, so it is possible that TolC, by interacting with MacA, can detect the nucleotide state of MacB. The binding of ATP to MacB could stimulate MacA to interact with TolC, possibly via the coiled coil hairpin of MacA, trigger the formation of the tripartite complex pump. Murakami (Murakami et al., 2006) proposed the conformational change caused by proton translocation and drug binding may be transmitted, via AcrA, to TolC, thus causing opening of the TolC periplasmic channel. In this case, the AcrA α-helical hairpin may be involved in the opening TolC, via its conformational flexibility (Mikolosko et al., 2006).

There is an analogy in the mechanism of operation of ABC importer that works in conjunction with a periplasmic binding-protein (Daus et al., 2007). In the E. coli maltose transporter, the binding of ATP to the ATPase subunit MalK induces conformational changes in the periplasmic loops of the membrane subunits MalF
and MalG (Daus et al., 2006); whilst EPR studies revealed that ATP-binding, but not ADP-binding, caused an increase in affinity between the transporter and the maltose binding-protein (MBP), and opening of the MBP (Chen et al., 2001). The binding of ATP to MacB could induce reorientation of the substrate binding-site of MacB and recruit TolC, via MacA, and simultaneous open the TolC to receive the translocated antibiotic. The interaction between TolC and MacA would uncouple them from MacB, promoting reorientation of the substrate binding-site that would be assisted by the rapid hydrolysis of ATP to ADP.

**Figure 7.1** Proposed mechanism of action for the MacA/MacB/TolC pump. MacAs are represented by orange squares; MacBs are colored in red, in which NBDs are represented by circles and TMDs are represented by ovals; TolCs are represented by blue cylinders; substrates are represented by black stars; ATP molecules are represented by green spheres.
Based on our studies of MacB and knowledge of the structure and function of other ABC transporters we proposed the following mechanism of action for the MacA/MacB/TolC pump. MacB recruits the substrate from the inner leaflet of the membrane or the cytoplasm, triggering a conformational change of MacB, from the inward-facing conformation to the outward-facing conformation, and relocates the NBDs of MacB, allowing the formation of the ATP sandwich in the presence of nucleotide (Figure 7.1). Meanwhile, the MacA can interact with the MacB dimer, increasing the affinity of MacB for ATP, helping the formation of ATP sandwich and decreasing the initial ATPase activity of MacB to stabilize dimer formation. The recruitment of the TolC to the MacAB complex allows translocation of the substrate across the periplasm and leads to the ATP hydrolysis; at this stage, MacA helps to seal the channel across the periplasm and aids ATP hydrolysis by increasing the steady state ATPase activity of MacB due to an increase in the release rate of the products of ATP hydrolysis, allowing a faster conformational change of MacB, from the outward-facing to the inward-facing conformation, resetting the transporter in its original conformation.
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MacB ABC Transporter Is a Dimer Whose ATPase Activity and Macrolide-binding Capacity Are Regulated by the Membrane Fusion Protein MacA*

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Gram-negative bacteria utilize specialized machinery to translocate drugs and protein toxins across the inner and outer membranes, consisting of a tripartite complex composed of an inner membrane secondary or primary active transporter (IMP), a periplasmic membrane fusion protein, and an outer membrane channel. We have investigated the assembly and function of the MacAB/ToIC/ol complex that confers resistance to macrolides in Escherichia coli. The membrane fusion protein MacA not only stabilizes the tripartite assembly by interacting with both the inner membrane protein MacB and the outer membrane protein ToIC, but also has a role in regulating the function of MacB, apparently increasing its affinity for both erythromycin and ATP. Analysis of the kinetic behavior of ATP hydrolysis indicated that MacA promotes and stabilizes the ATP-binding form of the MacB transporter. For the first time, we have established unambiguously the structure of the MacB ABC transport domain. This domain is an oligomeric protein that is arranged into two major domains, each of which is further subdivided into an N-terminal nucleotide binding domain and a C-terminal substrate-binding domain. These domains interact with each other by forming a continuous path across the membrane. The structure of the MacB ABC transport domain is a dimer with a single nucleotide binding site and a single substrate binding site. The dimeric structure is stabilized by a single intersubunit salt bridge.

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The abbreviations used are: IMP, inner membrane protein; βDDM, β-dodecyl maltoside; MFP, membrane fusion protein; OMP, outer membrane protein; AUC, analytical ultracentrifugation; ES-MS, electrospray-mass spectrometry; AMP-PNP, adenosine 5'-β,γ-imino-triphosphate; S-tag, for proteins fused to the 15-amino acid polypeptide (e.g. Lys-Glu-Thr-Ala-Ala-Ala-Lys-Phe-Glu-Arg-Gln-His-Met-Asp-Ser) derived from RNase A; NBD, nucleotide binding domain; MBP, maltose binding protein; AFM, atomic force microscopy.
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the periplasmic space. The structure of AcrA revealed an elongated monomer that is composed of three subdomains as follows: a β-barrel, a lipoyl domain, and a 58-Å-long α-helical hairpin (8). AcrA has been reported to interact with both TolC, via the α-helical hairpin (12), and AcrB, via the β-domains (13), to stabilize the tripartite complex (4, 5). Several lines of evidence suggest that assembly of the tripartite complex is a dynamic process that induces conformational changes in the protein components. The structures of a number of OMPs indicate that the channel is closed at one or both sides, requiring conformational changes to allow passage of the substrate (11–14). Several studies support the hypothesis that a small number of key interactions between adjacent coiled-coils of the OMP are broken to allow the inner coils to untwist and realign with the outer coils, thereby opening the OMP entrance aperture (11, 15–17). The conformational flexibility of AcrA may enable realignment of the coiled-coil to stabilize the open state of TolC (6, 11, 15). Recently, the structures of a number of different conformations of AcrB have been determined that support a mechanistic model in which there is a functional rotation in the periplasmic domains driving transfer of drugs into the OMP (18–21). The conformational asymmetry in the IMP is also reflected in the asymmetry in the open state of the OMP TolC (11). Such conformational changes in AcrB could be transmitted to TolC via AcrA. Furthermore, studies of the RND transporter AcrD from Enterobacter cloacae, whose typical substrates, aminoglycosides, are not expected to diffuse spontaneously across the lipid bilayer, revealed that only when AcrD was reconstituted with AcrA was it able to take up aminoglycosides, implying that AcrA is needed to "activate" AcrD (22). Collectively, these findings suggest that the MFP supports conformational changes in both the IMP and OMP.

We have sought to further investigate the assembly of the tripartite complex using the MacABToLC system responsible for the extrusion of macrolides from E. coli (3). To date this is the only tripartite pump for antibiotics that has been shown to utilize an ABC transporter, MacB, which has a novel architecture, consisting of a four-helix transmembrane domain, an N-terminal cytoplasmic nucleotide binding domain (NBD), and a large periplasmic domain formed by the loops connecting helices 1 and 2 (3, 23). The fact that the ATPase activity of the transporter can be monitored, using purified proteins, provides an additional tool for elucidating how its activity is modulated by assembly and disassembly of the tripartite complex. Indeed, recent studies, reported while this work was in progress, established that the MFP MacA modulates the steady-state ATPase activity of the IMP MacB (24). Our studies have confirmed these findings and, by undertaking pre-steady-state analysis of the kinetics, extended them by identifying how MacA modulates the ATPase mechanism of MacB. In addition, we have established that MacA also increases the capacity of MacB for binding erythromycin. This behavior provides a rationale for the retention of MacA by related systems from Gram-positive bacteria, such as Staphylococcus aureus, which lack an outer membrane and OMP (supplemental Fig. 1). Such analyses will not only have a bearing on understanding the function of tripartite drug-pumps but also of related toxin transporters.

EXPERIMENTAL PROCEDURES

Strains and Plasmids—The E. coli strains and plasmids used are described in supplemental Table 1, and the primers used for construction of plasmid vectors are described in supplemental Table 2.

Protein Overexpression and Purification—All the proteins used in this study were purified as fusion proteins with a six-histidine tag from E. coli overexpressing strains according to the protocols given in the supplemental material.

Pulldown Assays—For pulldown assays, proteins were overexpressed with a C-terminal S-tag for use with the cognate protein that was His-tagged. Membrane pellets, dissolved in detergent, or cell supernatant containing the S-tagged prey protein were mixed with 2 mg of purified His-tagged bait-protein. The mixture was loaded onto a Ni²⁺-charged Hitrap chelating column (GE Healthcare), so that the bait-protein could be immobilized on the column along with the prey-protein if they interact. The column was then washed with 15–20 column volumes of Tris buffer containing 75–100 mM imidazole and Triton X-100 (0.2% (v/v)) to eliminate any false-positive results because of nonspecific interactions. The bait-prey protein complex was eluted with 500 mM imidazole Tris buffer, containing Triton X-100 (0.2% (v/v)), and SDS-PAGE was used to visualize the bait- and prey-proteins. A Western blot was performed with anti-S-tag antibodies to confirm the presence of the S-tagged prey-protein.

Growth Curve Analyses—E. coli cells, of strain Kam3(DE3), harboring plasmids were grown at 37°C, with shaking at 200 rpm, until the cell density gave an A₅₅₀ of 0.5, when 0.05–0.1 mM isopropyl l-thio-β-D-galactopyranoside was added to the cells. The cells were grown for a further 3 h, when the cells were diluted with 2X YT media containing 50 or 100 µg/ml erythromycin, and the growth curve was recorded over the next 3 h. For experiments to measure the loss in growth because of erythromycin, cells were grown for 3 h in the absence and presence of 50 or 100 µg/ml erythromycin, and the growth loss is the ratio of the A₅₅₀ values.

Analytical Ultracentrifugation—Sedimentation equilibrium measurements were performed using a Beckman Optima XL-A analytical ultracentrifuge equipped with both absorbance and interference optics. 100 µl of MacB in buffer 1 (20 mM Tris, pH 8.0, 150 mM NaCl, 1% w/v glycerol, and 0.006% (w/v) βDDM) supplemented with 10, 25, or 50% D₂O was placed in the sample compartment of a Epon double-sector centerpiece, and 110 µl of buffer 1 was placed in the reference compartment. The final protein concentrations used in the runs were between 0.5 and 1.0 mg/ml. The D₂O was used to match the density of the solvent to the density of the detergent as described previously (25). The samples were centrifuged at 283 K (10°C) and 10,000, 15,000, and 25,000 rpm using an An60-Ti rotor. Scans were acquired using the absorbance optical system 15 h after the start of the experiment and in 1-h intervals until equilibrium was attained. Sedimentation velocity measurements were performed using the same hardware at 55,000 rpm at 10°C in buffer 1. The details of the data analysis are given as supplemental material.

Mass Spectrometry—Analyses were performed in a nanoflow ES mass spectrometer Q-ToF2 (Micromass). The following
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experimental parameters were used to record mass spectra of 2 mg/ml MacB in the Q-ToF2 instrument; needle voltage of 1.5 kV and MCP 2350 V.

**Atomic Force Microscopy**—MacB was diluted to a final concentration of 1 µg/ml, and 45 µl of the sample was allowed to adsorb to freshly cleaved mica. Imaging in air was performed with a Multimode atomic force microscope (Digital Instruments, Santa Barbara, CA) in tapping mode. The silicon cantilevers containing a diamond-like extratip had a drive frequency of ~300 kHz and a specified spring constant of 40 newtons/m (MikroMasch, Portland, OR), and the applied imaging force was kept as low as possible (target amplitude ~1.6–1.8 V and amplitude set-point ~1.3–1.5 V). The molecular volumes of the protein particles were determined from particle dimensions based on AFM images (see supplemental material).

**ATPase Assays**—An EnzChek phosphate assay kit (Invitrogen) was used to determine the ATPase activity of MacB hydrolyzing MgATP to release phosphate, when the reactants were mixed in a stopped-flow device (see supplemental material).

Generally, 2.3 µm protein was mixed with varying concentrations of ATP, up to 4 mM, in the presence of 6 mM MgCl₂ to ensure that all the ATP was complexed with Mg²⁺. In control experiments, no ATPase activity was apparent in the absence of Mg²⁺. Generally, for MacB alone, the hydrolysis of ATP was characterized by a P₁ burst, which was of near equivalence to the MacB concentration, consistent with the ATPase activity being attributable to MacB, rather than any contaminant proteins.

**Quantification of Erythromycin Binding to Affinity-purified Mac Proteins**—The equilibrium binding of [N-methyl-³¹C]erythromycin to purified Mac proteins was determined by rapid filtration and quantification of the radioactivity remaining on 0.2-µm filters as outlined in the supplemental material.

**RESULTS**

*MacA Interacts with Both MacB and TolC via Its Periplasmic Domain*—Interactions between *E. coli* MacA, MacB, and TolC were tested using detergent-solubilized proteins for pulldown assays (Fig. 1). MacA interacted with MacB (Fig. 1B), which we confirmed by cross-linking the proteins (supplemental Fig. 2A), of TolC, which was only found in the flow-through (lane 6), establishing that S-tagged TolC does not bind to the column. A protein M, marker was run in lane 2. A pulldown of N-terminal truncated MacA by MacB. An SDS-polyacrylamide gel shows the His-tagged MacB (lane 1), and the corresponding Western blot using antibodies to the corresponding protein fractions (indicated with arrows). The pulldown assay was performed with His-tagged MacB immobilized on a Ni²⁺-agarose column, over which a slurry of detergent-solubilized membranes from cells overexpressing S-tagged MacA was passed (lane 2). An SDS-polyacrylamide gel (lanes 1–8) probed with antibodies to the corresponding protein fractions (indicated with arrows). A weak band, which was not present in the absence of MacB, was apparent in lane 5, indicating a weak interaction between MacA and TolC. A control experiment was performed in the absence of immobilized MacA in which TolC was passed through a Ni²⁺-agarose column and the flow-through (lane 6), the 100 µM imidazole wash (lane 8), and the 500 µM imidazole elution (lane 7) were tested for the presence of MacA, which was only found in the flow-through (lane 6), establishing that S-tagged TolC does not bind to the column. A protein M, marker was run in lane 2. A pulldown of N-terminal truncated MacA by MacB. An SDS-polyacrylamide gel shows the His-tagged MacB (lane 1), and the corresponding Western blot using antibodies to the corresponding protein fractions (indicated with arrows). The pulldown assay was performed with His-tagged MacB immobilized on a Ni²⁺-agarose column, over which a slurry of detergent-solubilized membranes from cells overexpressing S-tagged MacA was passed (lane 2). An SDS-polyacrylamide gel (lanes 1–8) probed with antibodies to the corresponding protein fractions (indicated with arrows). A weak band, which was not present in the absence of MacB, was apparent in lane 5, indicating a weak interaction between MacA and TolC. A control experiment was performed in the absence of immobilized MacA in which TolC was passed through a Ni²⁺-agarose column and the flow-through (lane 6), the 100 µM imidazole wash (lane 8), and the 500 µM imidazole elution (lane 7) were tested for the presence of MacA, which was only found in the flow-through (lane 6), establishing that S-tagged TolC does not bind to the column. A protein M, marker was run in lane 6.
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Although we detected an interaction between TolC and MacB (Fig. 1D), the intensity of the band suggested a weak interaction. N-terminal truncated MacA (Δ20-MacA) interacted with MacB (Fig. 1E), indicating that it is the periplasmic domains of these proteins that interact. The fact that in each case the cognate pump protein could be pulled out of a complex mixture of detergent-solubilized proteins from membranes or cells indicated that the interactions are specific.

E. coli MacA and MacB Form a Functional Complex with TolC—The simultaneous expression of macA, macB, and tolC in the E. coli ΔacrB strain Kam3 (26) conferred resistance to erythromycin, indicative of the formation of a functional complex (Fig. 2A). Cells expressing macB with tolC conferred modest resistance to erythromycin in comparison with cells expressing macB alone, indicating that MacA is required to couple MacB to TolC (Fig. 2A). We sought to test whether the N terminus of MacA, which incorporates an α-helix that could interact with MacB, is required for the functional assembly of the complex. A construct in which the gIII-signal sequence was fused to truncated MacA, targeting it to the periplasm, was capable of conferring resistance to erythromycin (Fig. 2A), indicating that the N-terminal domain is not essential for the assembly of the functional complex. This is consistent with a report that a truncated lipid-deficient AcrA derivative was functional as judged by resistance of the cells to erythromycin (27). MacB alone conferred elevated resistance to erythromycin, probably because of its ability to pump the antibiotic into the periplasm, but we consistently found that expressing macB with either tolC or macA conferred greater resistance. Consequently, we sought to test if MacA could enhance this ability. To overcome the difficulty in comparing the growth of cells overexpressing multiple proteins that tend to grow at different rates, we monitored the growth of cells in the presence and absence of erythromycin and determined the growth loss (for cells growing in the presence of erythromycin in comparison with cells growing in the absence of erythromycin) (Fig. 2B). This revealed a significant loss in growth of the cells expressing MacB compared with those expressing MacAB, indicating that the simultaneous expression of MacA and MacB increases the resistance of the cells to erythromycin (Fig. 2B), suggesting that MacA enhances the ability of MacB to confer antibiotic resistance. Similarly, a previous study reported that MacAB, but not MacB alone, conferred resistance to macrolides (3).

MacB Forms Dimers—MacB has an atypical structure for an ABC transporter as it is predicted to have an N-terminal cytoplasmic NBD, which is connected to a four-helix transmembrane domain, with a large periplasmic domain formed by the loops connecting helices 1 and 2 (3, 23). If MacB resembles other ABC transporters that use a pair of NBDs to bind ATP, then it should function as a dimer. However, many transporters, including ABC transporters, have 12 membrane-spanning helices; MacB could adopt a similar topology by forming trimers. Furthermore, AcrB (4, 5) and TolC (9), which assemble into a tripartite complex with AcrA, clearly form dimers. If the trimeric arrangement of the periplasmic domains in AcrB forms a necessary scaffold for binding of AcrA, so that it can effectively interact with TolC, then by analogy the periplasmic domain of MacB might also be forced into forming trimers when interacting with MacA and TolC.

Therefore, we sought to determine the oligomeric state of MacB. Size-exclusion chromatography indicated that it forms higher order oligomers consistent with a dimer (data not shown), but such measurements are not only dependent upon the molecular weight but also the shape of the protein. Furthermore, there is a need to determine the number of detergent molecules complexed by the protein. Consequently, to determine whether the detergent-solubilized MacB was dimeric, we added a cross-linker to trap the oligomers; when we ran the cross-linked protein on an SDS-polyacrylamide gel, the most prominent band ran between the 120- and 160-kDa markers, indicative of a dimer, which has a calculated molecular mass of 145.8 kDa (supplemental Fig. 2B).

To further confirm the basic oligomeric unit as a dimer, we used two other techniques, analytical ultracentrifugation (AUC) and electrospray mass spectrometry (ES-MS). For the AUC experiments, we reduced the βDDM concentration to
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A representative sedimentation equilibrium profile from one of the runs (two different velocities of the same sample) is shown. Experimental data (dots) and fitted model for a 162.6-kDa particle (solid line) is shown for each. The bottom panel represents the residuals after fitting. A, B, MacA sedimentation velocity profiles of MacB are consistent with the formation of a stable dimer. The upper panel shows sedimentation profile curves at different time points, and the lower panel presents a c(s) size distribution analysis with solutions of the Lamm equation. The sedimentation coefficient is 6.8 S corresponding to an apparent molecular mass of 160 kDa, which is consistent with a dimer.

just below the critical micelle concentration to avoid the formation of micelles. To determine the detergent contribution in the buoyant mass of the protein-detergent complex, we used a series of different density buffers prepared using a range of DDM concentrations. The apparent molecular mass for MacB was determined from sedimentation equilibrium measurements to be 162.6 kDa (Fig. 3A) and from sedimentation velocity measurements to have a sedimentation coefficient of 6.8 S, corresponding to a molecular mass of 160.0 kDa (Fig. 3B). This molecular mass is greater than expected for monomeric and less than expected for trimeric MacB, complexed with bound detergent, but it is highly consistent with a MacB dimer to which about 16 βDDM molecules are bound. Although the amount of detergent bound to the MacB dimer appears to be lower than reported for RND (28) and MF (29) transporters, this reflects the fact that in our experimental set-up the detergent concentration was close to the critical micelle concentration, the amount of bound detergent was similar to that of other membrane proteins (e.g. MacB solubilized in 0.05% w/v βDDM bound 1.2 g of βDDM/g of MacB, which is equivalent to a βDDM:MacB molar ratio of 164:1). Electrospray-mass spectrometry (ES-MS) was used in nontandem con-
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**FIGURE 4. AFM analyses, AFM Imaging of MacB.**

A three-dimensional picture of a low magnification image of MacB acquired in air in Tapping Mode with a diamond-like extra tip of resonant frequency ~300 kHz and spring constant of 40 newtons/m. m and d show particles that belong to the first and second peak in B, respectively. B, frequency distribution of molecular volumes of MacB. The curve indicates a fitted Gaussian function. The m and d peaks correspond to volumes of 118 ± 1 nm$^3$ (n = 1642) and 238 ± 5 nm$^3$ (n = 665), consistent with the monomer and dimer, respectively. C, frequency distribution of molecular volumes of MacB that had been incubated with the nonhydrolysable ATP analogue AMP-PNP. The peaks correspond to volumes of 112 ± 3 nm$^3$ (n = 94) and 218 ± 14 nm$^3$ (n = 116). These data indicate an increase in the monomer-dimer ratio in the presence of AMP-PNP. D, high resolution images of structures where two small particles (m + m) are attached to one another, clearly indicative of dimer formation.

The mal specific activity of 8.9 nmol of ATP/min/mg MacB and a $K_m$ of 374 μM. A progressive reduction in the amplitude of the burst phase for ATP concentrations below the steady-state $K_m$ value hindered analyses because the pre-steady-state phase tended to merge with the steady-state phase; consequently, we only used the steady-state rates determined at ATP concentrations of 0.1 μM and above. For comparison, the lipid A transporter MabA, an half-ABC transporter, was characterized by a $V_{max}$ of 37 nmol of ATP/min/mg and a $K_m$ of 878 μM (31).

When MacA was added to MacB, at an equivalent or higher concentration, with both proteins in detergent, no phosphate burst was observed (Fig. 5A). Considering that the detergent was present for both the ATPase assays with MacB and MacAB, we only used the steady-state rates determined at ATP concentrations of 0.1 μM and above. For comparison, the lipid A transporter MabA, an half-ABC transporter, was characterized by a $V_{max}$ of 37 nmol of ATP/min/mg and a $K_m$ of 878 μM (31).

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In contrast to a previous study, which indicated that the ATPase activity of reconstituted MacB was not activated by N-terminal truncated MacA (24), we found that when MacB was mixed with Δ20-MacA no P$_i$ burst was apparent (data not shown). In the same study, and consistent with our findings, N-terminal truncated MacA was shown to interact with MacB; however, in contrast to our findings, when MacB was expressed with N-terminal truncated MacA in erythromycin-susceptible
FIGURE 5. MacA regulates the ATPase activity of MacB. A, time course for the change in P, concentration, corresponding to the absorbance change of the 2-amino-6-mercapto-7-methylpurine riboside in A, where 2.3 μM MacB was mixed with 1 μM ATP in the absence (upper trace) and presence (lower trace) of an equivalent concentration of MacA. In the absence of MacA, MacB produced a phosphate (P,) burst, with a rate and amplitude of 0.235 (±0.001) s⁻¹ and 2.20 (±0.01) μM, respectively. MacB did not produce a P, burst in the presence of MacA, B, steady-state rate of P, production by MacB as a function of the ATP concentration in the absence (lower curve) and presence (upper curve) of an equivalent concentration of MacA. The data are characterized by Vmax and K₅0 values of 6.9 (±0.7) nmol of ATP/mg MacB/min and 374 (±126) μM, respectively, for MacB alone; and of 12.3 (±0.3) nmol of ATP/mg MacB/min and 72 (±22) μM, respectively, for MacB in the presence of an equivalent concentration of MacA.

cells, there was no increase in erythromycin resistance (24). The reason for the difference with our own findings is unclear; however, in the previous study the signal sequence of OmpA was used to target MacA, truncated at position 32, to the periplasm (24), raising the possibility that shorter MacA sequence cannot exclude the possibility of the formation of additional conformations, so as to maintain the accessibility of the cytosolic substrate-binding site (32). In the case of MacB, its ATPase activity might instead be stimulated by TolC, to reset its conformation following drug transfer to TolC.

FIGURE 6. MacA increases the capacity of MacB to bind erythromycin—MacA Increases the Capacity of MacB to Bind Erythromycin—Although we could not detect any effect of erythromycin on the ATPase activity of MacB, we could detect the binding of [3H]erythromycin to detergent-solubilized MacB (Fig. 6). Importantly, we found that the MacAB complex bound more erythromycin than MacB alone (which binds considerably more than MacA alone) and that the amount bound increased in a concentration-dependent manner, as would be expected if MacA increased the affinity of MacB for erythromycin. However, because of the insolubility of the antibiotic in aqueous solutions, it was not possible to test a full range of erythromycin concentrations that might saturate MacAB. Consequently, we cannot exclude the possibility of the formation of additional sites within the MacAB complex that are not apparent in either MacB or MacA alone. Our data indicate that MacA not only modulates the ATPase activity of MacB but also enhances its capacity to bind erythromycin (be this due to an increase in affinity of MacB for drugs or the formation of additional drug-binding sites within the MacAB complex).

DISCUSSION

Gram-negative bacteria possess tripartite pumps that facilitate the extrusion of protein toxins and cytotoxic compounds, such as antibiotics, from the cell. In these tripartite assemblies the IMP is coupled to an OMP by a periplasmic MFP, forming a pump that can translocate molecules across both the inner and outer membranes. Intriguingly, Gram-positive bacteria that lack an outer membrane also possess MFPs, suggesting that they play a role in addition to stabilizing the interaction of the IMP with the OMP. Indeed, we have noted in some of these MFPs a large deletion corresponding to the coiled-coil hairpin that would interact with the OMP (supplemental Fig. 1). To investigate this possibility, we sought to determine the role of MacA, the MFP that couples the IMP MacB, an ATP-driven transporter, with the OMP TolC in E. coli to extrude macrolide
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We established that MacA interacts with both MacB and ToIC (Fig. 1) to form a functional tripartite complex (Fig. 2). However, we also found that MacA enhanced the resistance to erythromycin conferred by MacB alone (Fig. 2B), suggesting that it modulated the transport activity of MacB, which is consistent with our data demonstrating that MacA regulates the drug binding and ATPase activity of MacB (Figs. 5 and 6). The role of the MFP BesA in the activity of the BesABC pump in Borrelia burgdorferi, the causative agent of Lyme disease, has recently been highlighted (33). It is quite remarkable that in this system BesA also lacks the α-helical hairpin. As such this periplasmic protein is unable to unlock the periplasmic entry site of the OMP, a function that is attributed to the hairpin (11). To compensate, the OMP has evolved to be constitutively leaky. The reason for the retention of BesA in the Borrelia system is most likely associated with its function in the activation of the inner membrane component, in that case an RND transporter. It is conceivable then, that a similar role is also present in the ABC transporters and their associated MFPs, and it could explain the preservation of the hairpin-lacking MFPs in Gram-positive bacteria, such as S. aureus (supplemental Fig. 1).

Although E. coli MacB has an NBD, which incorporates Walker A and B motifs and an ABC signature sequence that are characteristic features of members of the ABC superfamily, it is not a classic ABC transporter. It has a large periplasmic domain, reminiscent of that found in RND transporters, which form trimers in which these domains form substantive sites of contact between the protomers. Consequently, we sought to determine the oligomeric state of MacB and in doing so have developed a novel ES-MS approach to unambiguously establish that MacB forms dimers (Fig. 3C). ES-MS is now widely accepted as a powerful method to determine accurately the stoichiometry of intact protein complexes (34). However membrane proteins, solubilized by detergent or adsorbed in micelles, have remained difficult to analyze under similar MS conditions. The development of strategies to tackle this field is challenging, primarily because the large quantities of detergent suppress the protein signal, whereas the poor solubility of membrane proteins in aqueous buffers often causes the electrospray needle to block. To date only a few MS studies have reported the observation of membrane proteins or their complexes by MS (35-39). Here we report on the use of a miniaturized form of ES with reduced flow rates (nano-ES), and a high collision energy that facilitates the desolvation process and induces dissociation of detergent-proteins clusters, to determine the oligomeric state of an integral homomeric membrane protein. In our protocol we used lower quantities of βDDM, than have been reported previously, without apparent detrimental effects on the stability of the protein. Under our experimental conditions, we have successfully maintained the noncovalent subunit interactions, such that the oligomeric state of the protein complex could be determined without ambiguity. As would be expected for an ABC transporter in which the NBDs interact, our biophysical studies of MacB indicate that it forms dimers (Figs. 3 and 4 and supplemental Fig. 3). This is an interesting observation because it raises the following question. How does dimeric MacB interact with trimeric ToIC and the number of MacA molecules needed to stabilize the tripartite complex? Extension of our AFM studies will be vital for determining the stoichiometry of the interactions to provide an understanding of the assembly of the tripartite complex, which may be difficult to address by other methods, such as crystallization of the complex.

Our findings and previous studies, which have shown that disruption of the Walker A and B motifs not only inhibits the ATPase activity but also blocks the capacity of MacB to confer macrolide resistance (24), suggest that MacB operates by a similar mechanism to typical ABC transporters. Our understanding of how ABC transporters couple ATP hydrolysis to transport is still rudimentary, but the determination of the crystal structures of several ATPase subunits and complete ABC transporters has suggested conservation of key steps in the molecular mechanism. The binding of ATP to both soluble ATPase subunits (40), solubilized NBDs from ABC transporters (41, 42), and to the NBDs within a complete ABC transporter (43, 44) has been shown to promote dimerization as the ATP is bound at the interface of these nucleotide-binding sites, sandwiched between the Walker A motif of one NBD and the ABC signature motif of the other NBD. Our AFM studies indicated that there is a higher proportion of MacB dimers in the presence of the nonhydrolysable nucleotide AMP-PNP (Fig. 4). In complete ABC transporters, the binding of ATP causes the transporter to adopt a conformation in which the substrate-binding site is outward-facing, because ATP bridges the two NBDs, closing off the inward-facing substrate-binding site (45). Conversely, the release of the hydrolysates ADP and phosphate is thought to promote an inward-facing conformation, as the structural constraint imposed by binding of the nucleotide is released. Consistent with this proposal, several studies indicate that NBD dimerization cannot be induced by ADP (46), because interactions between the γ-phosphate of ATP and the signature sequence catalyze these events (47).

We found that MacA modulates these processes in MacB (Fig. 5), and so our kinetic data have clear implications in terms of such a mechanism. Most significantly, MacA increases the apparent affinity of MacB for ATP, while decreasing the rate of ATP hydrolysis, so as to promote and stabilize the ATP binding conformation, which we presume to be the conformation in which the antibiotic-binding site is outward-facing. In this manner, MacA would play a direct role in driving antibiotic translocation between MacB and ToIC.

During the course of our studies another investigation reported on the effect of MacA on the steady-state ATPase activity of MacB (24). In this study, MacB, solubilized in Triton X-100, appeared to have a specific activity that was an order of magnitude higher than we had found; however, a discontinuous assay was used to determine the ATPase activity, and this could well have been influenced by pooling the time points from both the burst and steady-state phases. Indeed, $k_{cat}$ was marginally slower than the hydrolysis rate determined from the $P_x$ burst phase in our experiments (e.g. 0.17 versus 0.24 s$^{-1}$). Possibly for similar reasons, this assay did not detect the effect of MacA on MacB solubilized in Triton X-100. However, they did find that MacB reconstituted into liposomes was characterized by a reduced $k_{cat}$ (e.g. $k_{cat} = 0.10$ s$^{-1}$) and decreased affinity of MacB for ATP (e.g. $K_m = 2.30$ mM). Co-reconstitution of MacB with MacA had the effect of increasing both $k_{cat}$ (e.g. $k_{cat} = 0.78$ s$^{-1}$)
and the affinity of MacB for ATP (e.g.  ) This behavior is similar to the effect of MacA on the steady-state kinetics of the ATPase activity of solubilized MacB in our study. To know if the reconstituted protein behaves in an identical manner to the solubilized protein would require the reconstitution of sufficient amounts of MacB to define any P, burst, and the kinetics of the ATPase activity of solubilized MacB in our study. Consequently, although these earlier studies support our conclusion that MacA affects the ATPase activity of MacB, they do not give the detailed insight into the ATPase mechanism provided by our pre-steady-state analyses.

Our studies established that conformational changes can be propagated from the periplasmic domain of MacA to the cytoplasmic NBD of MacB (Figs. 1 and 2), and consequently, it is plausible that TolC, by interacting with MacA, can detect the nucleotide state of MacB. The binding of ATP to MacB would stimulate MacA to interact with TolC, inducing the latter to adopt the open state. Because ATP and MacA stabilize in the outward-facing conformation of MacB, drug transfer from MacB to TolC would be facilitated. TolC would then communicate with the NBD of MacB (again, such communication is probably conveyed via MacA) to stimulate ATP hydrolysis, which would be required to reset MacB in the inward-facing conformation. If ATP hydrolysis is controlled by TolC, so that the MacB conformation is only reset after the interaction with TolC and productive transfer of drugs, this would provide an explanation of the apparent insensitivity of the ATPase activity of MacB to drugs. Indeed, without such a feedback control mechanism of drug export, it would appear that drug stimulation of the ATPase activity of MacB would be counter-productive because the MacB conformation could be reset before drug transfer to TolC. There is an analogy in the mechanism of operation of ABC transporters that work in conjunction with a periplasmic binding protein (47). In the E. coli maltose transporter, the binding of ATP to the ATPase subunit MalK induces conformational changes, detected by tryptic digestion, in the periplasmic loops of the membrane subunits MalF and MalG (48), although EPR studies revealed that ATP binding, but not ADP binding, caused an increase in affinity between the transporter and the maltose-binding protein (MBP), and “forcing open” the bound MBP to release its substrate (49). The transporter is reset in its original inward-facing conformation by ATP hydrolysis, which is stimulated by the MBP (50). In our model TolC replaces the MBP and, because the open state requires disruption of the second selectivity filter and subsequent twisting of the helices of the TolC channel to keep it fully open, there is a need for the MacA hairpins to stabilize this conformation, which could not be achieved by interaction with MacB alone. Although we have interpreted our findings in terms of an alternating conformation model for MacB, in which the drug-binding sites are inwardly and outwardly exposed, such a model could easily be refined to account for drug binding to a fixed periplasmic site, in analogy to AcrB, with ATP binding and hydrolysis coupled to conformational changes that induce TolC association and dissociation. Clearly the work presented here provides an important framework for further studies that will enable the elucidation of the mechanism underlying the dynamics of assembly of the MacAB/TolC tripartite pump in the future.

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MacB Dimer Is Regulated by MacA