Efflux-Mediated bile Resistance in Gram-Positive Pathogens

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Abstract

Gram positive pathogens are causing many serious infections that affect humans and result in mild to severe diseases worldwide. In order to survive and initiate infection, enteric pathogens must resist the physiochemical defence factors in the human intestinal tract. One of these defence factors is bile, a potent antibacterial like compound in the intestine. Efflux pumps are the important mechanism by which bacteria resist antibacterial agents such as bile. Efflux of antimicrobial substances outside the bacterial cell is considered as a key factor for intestinal colonization and virulence of enteric pathogens. This paper will review the research conducted on efflux – mediated bile resistance in *Staphylococcus aureus*, *Listeria monocytogenes*, *Enterococcus faecalis* and *Clostridium perfringens*. These bacteria colonize in the human & animal gastrointestinal tract and they have a multiple mechanism to resist the innate defences in the gut and antibacterial activity of bile. However, bile resistance in these bacteria is not fully understood. The evidence from this review suggests that Gram-positive pathogens have the ability to active transport of bile. Further research is needed to know how these pathogens sense bile and how bile regulates its virulence factor. In general, therefore, it seems that understanding the specific mechanism of bile resistance in enteric bacteria including gram-positive pathogens may involve in the development of novel strategies to control and treatment of gastrointestinal infections.

Keywords: Efflux pump, Bile salt, antimicrobial agent resistance, enteric gram positive pathogen

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(Received: August 2, 2021; accepted: November 29, 2021)
INTRODUCTION

Bile is a digestive secretion made by hepatocytes of the liver and enters the small intestine via the bile duct. The main functions of bile are emulsification and solubilization of lipids and as a hormone regulating food metabolism.1-2 Bile salts are the major component of bile and constitute about 50% of its organic component. It is well documented that bile salts are one of the detergents that have antibacterial action against a wide range of enteric bacteria. Enteric Gram-positive pathogens must resist the toxic action of bile salts to survive inside the intestinal tract. These toxic actions can disrupt the bacterial membrane, induce oxidative stress and DNA damage and dissociate membrane protein.2,3 Thus, these pathogens have developed different mechanisms to resist bile salts. The possible mechanisms by which bacteria resist bile include synthesis of various types of fatty acid and surface protein, reducing oxidative thiol modifications, restoring redox homeostasis, bile transport using efflux pump system & hydrolyze of bile acids using bile salt hydrolases (BSHs). However, the specific mechanisms are not fully understood yet.3-6

Bile efflux is one of the important mechanisms, whereby bacteria resist the detergent action of bile by pumping out the bile salt from their cytoplasm to the external environment. Pumping out of antibacterial agents including bile salt needs particular proteins known as efflux pumps.7 The antimicrobial and other harmful agents that affect the bacteria induce expression of the efflux pumps.8 Also, efflux pumps are more than only antibiotic resistant factors. They even help the microorganisms to pump out the heavy metals, solvents and biocides or any harmful agents.9

After discovering the efflux pumps in the 1990s, molecular microbiology improvement led to recognise different types of efflux pumps in Gram-positive bacteria such as, methicillin-resistant Staphylococcus aureus (MRSA), Streptococcus pneumoniae, Clostridium difficile, Enterococcus spp. and Listeria monocytogenes and Gram-negative bacteria like, Acinetobacter baumannii, Escherichia coli, Klebsiella pneumoniae, Stenotrophomonas maltophilia, Campylobacter jejuni, Pseudomonas aeruginosa, Vibrio cholerae and Salmonella spp.10,11

Basically, efflux pumps in bacteria are found as five groups depending on composition, source of energy and number of transmembrane spanning regions as; the resistance-nodulation-division (RND) family, the major facilitator superfamily (MFS), the ATP binding cassette (ABC) superfamily, the small multidrug resistance (SMR) family and the multidrug and toxic compound.12,8 Biologically, the resistance to the antibacterial agent means that the agent is unable to affect the target place with enough concentration that inhibits the activity of the pathogen. The resistance to tetracycline by E. coli was the first efflux pump that identified in bacteria.13,14 Also, several studies suggested that the inaction of some efflux pumps in Gram-positive pathogens results increase the sensitivity of these bacteria to bile.15 So, in this review we focused on the role of efflux pump systems on the ability of some Gram-positive pathogens such as staphylococcus aureus, Listeria monocytogenes, Enterococcus faecalis and Clostridium perfringens to resist bile and survive in the gastrointestinal tract.

Efflux pump systems in Gram-positive pathogens

The microorganisms have improved their defence mechanisms inside the host through various strategies as; inactivation of the enzymes,16,17 changing the drug targets or permeability18,19 or by efflux pump overexpression.17,11 Bacteria have either selective efflux pumps for one type of substance or nonselective to transport different classes of antibiotics.11,8 Essentially, bacterial antibiotic resistance has developed due to the exposure to the antibiotics or it could be naturally resistant to some antibiotics.20 For example, E. coli has marRAB locus that intrinsically carried by the chromosomal DNA which is responsible for E. coli resistance to some antibiotics as penicillin, cephalosporins, fluoroquinolones, tetracyclines, chloramphenicol, etc.21,22

Efflux pumps are transporter proteins that are located in the plasma membrane of the bacterial cell.23 There are five efflux transporter families in prokaryotes that are responsible for antibiotic resistance. Some efflux pump transporters resist to a narrow spectrum of...
antibiotics as TetM in *E. coli* against tetracyclines or MexCD-OprJ of *P. aeruginosa* against fourth generation cephalosporins. While others resist a wide range of antimicrobials such as AdeABC of *A. baumanii*, NorA of *S. aureus* and AcrAB-ToIC of *E. coli*. Lately, the sixth bacterial efflux pump family has been discovered in bacteria which is called the Proteobacterial Antimicrobial Compound Efflux (PACE) superfamily. Since, many researches focused on multidrug resistance MDR in Gram-negative pathogens due to the complex mechanisms of MDR in these bacteria because of the double-membrane around the bacterial cell that enables the expression of a triple efflux pump system as MexA/MexB/OprM of *Pseudomonas aeruginosa* or AcrA/AcrB/ToIC of *Enterobacteriaceae*. Here is focusing on efflux pump transporters in Gram-positive pathogens and in detail pumping of bile in *S. aureus* bacteria which belong to four main groups of efflux pump: major facilitator superfamily MFS, ATP-binding cassette ABC, Multidrug and Toxic Compound Extrusion MATE and small multidrug resistance SMR. **MFS (major facilitator superfamily)**

In Gram-positive bacteria, the MDR efflux transporters include SMR, MATE, ABC and MFS. MFS transporter family largely exists in prokaryotes and even in eukaryotes. Type of the transport is actively using the proton motive force (PMF) as a source of energy. MFS proteins are mostly 400 to 600 A.A. In length organized as 12 or 14 membrane-spanning helices and there is a large loop in location between helices six and seven. The MFS efflux proteins in Gram-positive bacteria that suitable examples for this family of transporters are, Nor A, NorB and MdeA in *Staphylococcus aureus*, LmrB, Bmr and Bmr3 in *Bacillus subtilis* and MeFe in *Streptococcus pneumoniae*. **SMR (small multidrug resistance)**

SMR transporter family is the smallest among other efflux pump groups. It is secondary multidrug transporters and belongs to the drug/metabolite transporter (DMT) family. The proteins in this family basically consist of 100-120 A.A., have 4 transmembrane helices and short loops between the helices act as homodimers. These transporters utilize proton motive force to pump toxic material from the bacteria to the outside. SMR transporters are found in Gram-negative and Gram-positive pathogens. The EmrE isolated of *E. coli* is the protein that studied well and has about 110 residues. Also, EbrAB of *Bacillus subtilis* and Smr, QacG, QacH of *Staphylococcus aureus* are famous examples of the SMR efflux pumps in Gram-positive bacteria. **MATE (Multidrug and Toxic Compound Extrusion)**

This group of transporters was discovered recently and exists in Gram-negative and Gram-positive bacteria. MATE proteins are mostly consisting of 400 to 700 A. A. with 12 transmembrane helices and all these transporters are similar in about 40% of their sequences. MepA protein of *Staphylococcus aureus* is a common example to MATE transporters in Gram-positive pathogens. In this group of efflux pumps, different antibiotics can be pumped out of the bacterial cell as norfloxacin, chloramphenicol, ciprofloxacin, kanamycin, ampicillin and others. Also, two sources of energy used by MATE proteins; the PMF and the sodium ion (Na+) gradient. NorM protein of *Vibrio parahaemolyticus* is the first famous example of this group. This transporter can also extrude aminoglycosides and dyes. In addition, MepA in *S. aureus* is considered as one of the well-studied MATE proteins in Gram-positive bacteria. Also, there are other transporters of MATE group were studied such as YdhE in *E. coli*, HmrM in *Haemophilus influenzae*, PmpM in *P. aeruginosa* and CdeA in *C. difficile*. **ABC (ATP-binding cassette)**

This group of efflux pumps is large and widely spread in prokaryotes and eukaryotes. ABC proteins are active transporters that use ATP hydrolysis in process of removing the toxic substances to the outside of the cell. These proteins are consisting of two nucleotide-binding domains (NBDs) and two transmembrane domains (TMDs). The TMDs include six transmembrane α-helices and they are homo or heterodimers and highly hydrophobic that able to recognise the harmful substances and extrusion. While, the NBDs are hydrophilic and act as the ATPase subunit in the cytoplasm which bind and hydrolyse ATP.
efflux families which is the source of energy through ATP hydrolysis. ATP binding and hydrolysis causes changes in ABC proteins structure that allows transport of materials to the outside. LmrA in *Lactococcus lacti* and Rv1217c–Rv1218c in *Mycobacterium tuberculosis* are ABC efflux transporters example in Gram-positive bacteria.

Moreover, ABC transporters act as a triple efflux pump in Gram-negative bacteria that work with the membrane fusion protein and an outer membrane protein such as MacB of *E. coli* which work with the outer membrane channel (ToIC) and the periplasmic adaptor (MacA). LmrA protein of *L. lactis* was the first ABC transporter that recognised in Gram-positive bacteria. Also, Sav1866 in *S. aureus* is the model for the ABC transporter-mediated multidrug efflux. In addition to LmrA and Sav1866 are homologous of the P-glycoprotein which are multidrug transporters in humans.

**Efflux pump inhibitors as new antibacterial agents**

Efflux pump inhibitors EPIs are molecules that are able to block efflux pumps activity and inhibit their function to extrude the antibacterial agents out of the cell. Also, they can restore the antibiotic role against the pathogens. Depending on their origin, the EPIs are either inhibitors from plant-source or synthetic inhibitors. In Gram-positive bacteria, for example, efflux pump inhibitors against the NorA system of *S. aureus* were fully studied. Both natural and synthetic EPIs are effective against NorA in *S. aureus* as Genistein derived from Lupinus argenteus, Spinosan A from Dalea spinosa and Tiliroside from Herissantiubae as a plant-origin efflux pump inhibitor in particular.

It was noticed that the synthesised EPIs have toxic side effects after use in most cases. While, many natural EPIs are very effective against different groups of efflux pumps and can be used safely. EPIs use different mechanisms to block the efflux pumps function which results in the stop delivery of drugs or any antimicrobial factor to the outside. Thus, EPIs act as adjuvants with the antibiotics can support their role and inhibit efflux pumps transporter expression.

Using efflux pump inhibitors EPIs appears to be the resolution to this problem or as a discovery of a new antibiotic. Also, it can assist to decrease the probability of producing a mutant generation in addition to pumping out the antibiotics. Moreover, EPIs mixing with antibiotics acts as a successful strategy to prevent the diseases caused by MDR pathogens via stopping the expression of efflux genes. In addition to the effect of efflux pumping the antimicrobial agent, they also decrease the concentration of the agent and may cause mutants.

![Fig. 1](https://example.com/fig1.png)

**Fig. 1.** Representation of different types of efflux pumps in Gram-positive bacteria and the main types of efflux pumps associated with bile resistance in some gram-positive pathogens.
Bile efflux system in gram-positive pathogen

Enteric G-positive pathogens have multiple mechanisms to resist the detergent effect of antibacterial agents such as bile. One of the important mechanisms in the bile resistance is the efflux pump system. Several studies documented the relationship between bile resistance in gram-positive pathogen and drug efflux pumps. Sannasiddappa et al. demonstrated that *S. aureus* MnhF a secondary efflux pump involved in the resistance of this bacteria to unconjugated bile acid. He found that inactivation of MnhF results in a significant decrease in the ability of *S. aureus* to resist the detergent effect of bile and the ability of the bacterium to survive in the in vitro human colonic model. Furthermore, expression of *S. aureus* mnhf in *E. coli* enhanced the resistance of the host cell to unconjugated bile acids.

EF0420 and EF1814 gene which encodes EmrB/QacA family drug resistance transporters protein in *Enterococcus faecalis* showed increased transcriptional activity upon exposure to 1% of bovine bile. *Enterococcus faecalis* EmrB/QacA efflux pump plays a role in PMF dependent extrusion of bile which is required for active extrusion of bile. In *Listeria monocytogenes*, several studies reported the role of efflux bases transporters in bile resistance. Exposure of *Listeria monocytogenes* to 1% porcine bile results in the induction of multidrug resistance efflux pump of the major facilitator superfamily MdrM and MdrT. *L. monocytogenes* MdrT mutant were found more susceptible to cholic acid and bile than wild type. Sleator et al. suggest that active efflux of bile from the bacterial cell is a more effective mechanism than other mechanisms such as bile detoxification. They identify a novel bile efflux system (BilE) in *Listeria monocytogenes*, which is a new virulence factor in this bacterium and play a critical role in intestinal colonization. BilE belongs to ABC type MDR efflux transporter in *Listeria monocytogenes* and have sequence similarity to members of the betaine carnitine choline transporter (BCCT) family. The mutation in BilE gene was associated with defects in removing bile and increasing the sensitivity of the bacterium to bile. Sleator and colleagues found that OpuC, an ABC transporter has a significant role in colonizing and survival of *L. monocytogenes* in the small intestine. Enteric bacteria including *L. monocytogenes* able to resist hyperosmotic stress in the gut lumen by accumulating of active osmotic compounds such as carnitine and betaine. A recent study demonstrates that OpuC is a high-affinity carnitine transporter. Mutation in the carnitine transport system, OpuC lead to an increase of the bacterium sensitivity to bile in general, therefore it seems that *L. monocytogenes* OpuC has a vital role in resistant the acute toxicity of bile. *Bacillus cereus* a gram-positive opportunistic pathogen, which is responsible for a wide range of infections in humans such as food poisoning. It is well documented that *Bacillus cereus* has the ability to germinate and sporulate in the mice gut, however, little data is available about the ability of the bacterium to survive and colonize the human gut. Transcriptional response to bile shows that several gen in *Bacillus cereus* was unregulated. Induction of EmrB/QacA transporter family in the presence of bile refers to the role of this type of efflux pump in bile response. A few experimental data is available about the role of Clostridium efflux pump in bile resistance.

**CONCLUSION**

This assay has reviewed the role of efflux pump system in bile resistance in gram-positive enteric pathogens. The ability of enteric bacteria to resist bile is required for colonization and initiating infection. The specific mechanism by which enteric pathogen resists bile is not fully understood, however, accumulating evidence suggests that efflux pump plays a key role in bile resistance in a wide variety of enteric pathogens. These pumps transport bile outside of the bacterial cell and then decrease its harm. As previously mentioned, bacterial efflux pumps can be classified into six main families, including ABC, MFS, SMR, RND, MATE and PACE family. This is the first study reporting efflux-mediated bile resistance in gram-positive pathogens. The investigation of efflux mediated bile resistance in *Staphylococcus aureus*, *Listeria monocytogenes*, *Enterococcus faecalis*, *Bacillus cereus* and *Clostridium perfringens* has shown that most transporters related to bile resistance belong to ABC and MFS family. ABC family transporters need energy (ATP) to drive transport. While MFS family drives transport using the proton motive force (PMF) as a source of energy. Upregulation of gene coding efflux
pump proteins upon exposure gram pathogen to a sublethal dose of bile, and increasing sensitivity to bile in pathogen lack to efflux pump (mutant) are the main methods were used in studies of bile tolerance. These studies suggest that gram-positive pathogens have the ability to active transport of bile, however Further research is needed to know how pathogenic bacteria sense bile and how bile involves in virulence factor regulation.

ACKNOWLEDGMENTS

None.

CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

AUTHORS’ CONTRIBUTION

All authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

FUNDING

None.

DATA AVAILABILITY

All datasets generated or analyzed during this study are included in the manuscript.

ETHICS STATEMENT

Not applicable.

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