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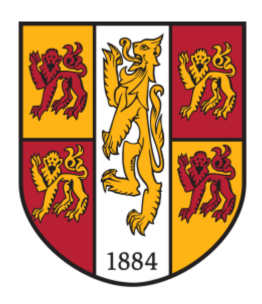
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EPIDEMIOLOGICAL GENE CARRIAGE STUDY OF GRAM-NEGATIVE EXTENDED SPECTRUM β-LACTAMASE-PRODUCING BACTERIA IN URINARY TRACT INFECTION PATIENTS IN NORTH WALES; ISOLATION OF A NOVEL CTXM-TYPE ENZYME



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A thesis submitted for the degree of Doctor of Philosophy January 2020

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Dedication

To the memory of my beloved parents, who would have been happy to see me Follow in this step.

To my loving wife, a constant source of inspiration and motivation to me.

To my brothers and sisters.

I dedicate my work with wholehearted love and gratitude to you all.

Yasir

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Abstract

Antibiotic resistance is considered a major public health concerns in the 21st century. Microbes that were once susceptible to antibiotics are becoming more and more difficult to treat as they acquire rapidly a wide range of resistance mechanisms including enzymes of the β-lactamase family. These proteins are typically plasmid-borne and inactivate beta-lactam ring containing antibiotics that are worldwide used to treat infections. This enzyme family with more than 2800 proteins is rapidly evolving through the spontaneous acquisition of missense mutations that enhance protein stability, increase catalytic turn-over and/or improve substrate specificity.

This study is an epidemiological snapshot of the antibiotic resistance conferred by CTX-M-type extended-spectrum β -lactamases (CTX-M type ESBL) in patients based in North Wales who were diagnosed with urinary tract infection (UTI).

Multiplex PCR amplification of β -lactamase genes of the *bla*CTX-M groups 1, 2, 9, and 8/25 followed by DNA sequencing of ESBL-producing isolates was performed with samples from three referral hospital in North Wales; Ysbyty Gwynedd, Glan Clwyd Hospital, and Wrexham Maelor Hospital

In line with previous studies, the β-lactamase CTX-M-15 was found to be the dominant CTX-M variant in the study region. The analysis of samples from Ysbyty Gwynedd hospital at Bangor revealed however an as yet unknown CTX-M variant that differed from CTX-M-14 by only three missense mutations. The amino acid substitutions A55T, A273P and R277C reside in the beta-strand domain opposite of the active site of the enzyme. CTX-M-15, CTX-M-14 and the novel variant were expressed in the periplasm of recombinant *E.coli* using the inducible vector pASK-IBA2C. Determination of the Minimum Inhibitory Concentration (MIC) of a series of clinical relevant antibiotics using the induced recombinant strains showed a significant increase in Nitrofurantoin resistance and a partial increase in Cefoxitin resistance that were both specifically linked with the three mutations. In vitro Kinetic studies using protein from periplasmic extracts of induced strains showed however that neither the affinity for Nitrofurantoin nor for Cefoxitin was increased by these substitutions. Taken together, the three missense mutations in CTX-M-14 may render UTI patients isolates more resistant to Nitrofurantoin because the three amino acid changes increase enzyme stability, its expression level in the resistant coliform bacteria or result in a higher turn-over rate of the mutated β -lactamase by facilitating conformational changes.

Abbreviations

APS Ammonium persulfate

AMR Amicrobial resistance

AMP Ampicillin

AMC Co-amoxiclav

AK Amikacin

Amo Amoxicillin

Aug Augmentin

Ami Amikacin

bp Base pair

BLAST Basic Local Alignment Search Tool

CTX Cefotaxime

CAZ `Ceftazidime

CA Clavulanic acid

CL Cephalexin

CIP Ciprofloxacin

CPD Cefpodoxime

CT Cefotaxime

CFU Colony Forming Unit CFU

C-terminal Carboxy-terminal domain

CDC Centres for Disease Control

CLSI Clinical & Laboratory Standards Institute

DNA Deoxyribonucleic acid

DDST Double-disk synergy test

DMF Dimethylformamide

ddH2O distilled water

EUCAST European Committee on Antimicrobial Susceptibility Testing

E.coli Escherichia coli

EDTA Ethylenediamine tetra-acetic acid

ESBLs Extended-spectrum b –lactamases

ETP Ertapenem

FX Cefoxitin

GN Gentamicin

GP General Practice

HGT Horizontal gene transfer

HCl Hydrochloric Acid

IMP ImipenemIMI ImidazoleIP InpatientskD Kilodalton

KPC Klebsiella pneumonia carbapenemase

L Litter

Mg Milligram

MgCl2 Magnesium chloride

ml Millilitre mM Millimolar

MHT Modified Hodge test

MALDI Matrix Assisted Laser Desorption Ionization-Time of Flight

MIC Minimum inhibitory concentration

MRSA Methicillin-resistant S. aureus

MEM Meropenem

MHB Mueller Hinton Agar

μg Microgram μL Microliter

NARMS The national antimicrobial resistance monitoring system

NHS National Health Service

NaCl Sodium chloride

N-terminal Amino-terminal domain

NaOH Sodium hydroxide

Ni Nitrofurantoin

oligo-dT Oligodeoxythymidylic acid

OMPs Outer membrane proteins

OD Optical density

OP Outpatients

PAGE Polyacrylamide gel electrophoresis

PBPs Penicillin-binding proteins

PBS Phosphate-buffered saline
PCR Polymerase chain reaction
PCR Polymerase chain reaction
PBP Penicillin binding protein
Ptz Piperacillin/tazobactam

Pi Isoelectric point

SD Standard deviation

SDS Sodium dodecyl sulphate

SDS-PAGE Sodium dodecyl sulfate-Polyacrylamide gel

SHV Sulfhydryl variable

TBE Tris-borate-EDTA

TEMED Tetramethylethylenediamine

Tet Tetracycline

TZP Piperacillin/tazobactam

TZ Cefotazidime

UV Ultraviolet

UTI Urinary Tract Infection

Vmax Maximum Velocity

WB Western blot

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Chapter 1: INTRODUCTION

1.1. Antibiotics: The 'Miracle' Drugs

Infectious diseases were the main cause of human deaths worldwide in first half of the 20th century and the second leading cause of mortality worldwide in recent decades (Nathan & Cars, 2014). For millennia, medicine was in the dark ages, relying on plants, fungi, and lichen to treat infections (Bernardini et al, 2018), until research into microbial secondary metabolites led to the antibiotic era (Landecker, Hannah & Blackman, 2016). Antibiotics are one of the most important inventions in the pharmacological field and have been considered 'miracle' drugs since they were first introduced to the health care system in the 1940s (Brandt et al, 2014). The discovery of antibiotics revolutionized the medical sector, forever, changing the way medicine is practiced (Laxminarayan et al, 2013).

Many antibiotic agents have been developed to kill bacteria and inhibit their growth or reproduction (Yount, 2013). Antibiotics are categorised into three main groups according to the mechanism through which this action occurs. The first group are inhibitors of bacterial cell wall biosynthesis. These compounds cause the rupturing of the bacterial cell wall by inhibiting the action of transglycosidases, the enzymes responsible for the cross-linking of peptidoglycan (sugar polymers) molecules that give the cell wall strength (Kapoor et al, 2017). The second group includes the inhibitors of bacterial protein synthesis. These compounds target the 30S or 50S subunit of the bacterial ribosome (70S ribosome), which is responsible for the translation of mRNA into protein (Lin et al, 2018). The third group are the inhibitors of nucleic acid production. These are classified into DNA inhibitors and RNA inhibitors. DNA inhibitors, such as quinolones, act upon DNA gyrase as a topoisomerase inhibitor (Koo et al, 2017). RNA inhibitors, such as rifampin, acts upon DNA-dependent RNA polymerase (Harms et al, 2016). Researchers may also classify antibiotics into two groups according to their specific targets. Broad spectrum antibiotics affect a wide range of bacteria and narrow spectrum antibiotics target only specific types (Wright, 2007). Table 1.1 shows the most commonly used antibiotics and their target.

Table 1.1 The most commonly used antibiotics and their targets.

Antibiotic class	Example(s)	Target	Reference
β-Lactams	Penicillins (ampicillin),	Peptidoglycan	Sauvage et al,
	cephalosporins	biosynthesis	2016
	(cephamycin), penems		
	(meropenem),		
	monobactams (aztreonam)		
Aminoglycosides	Gentamicin, streptomycin,	Translation	Hong et al, 2014
	spectinomycin		
Glycopeptides	Vancomycin, teicoplanin	Peptidoglycan	Binda et al, 2014
Tetracyclines	Minocycline, tigecycline	Translation	Goodman et al,
			2016
Macrolides	Erythromycin, azithromycin	Translation	Jelić et al, 2016
Lincosamides	Clindamycin	Translation	Matzov et al,
			2017
Streptogramins	Synercid	Translation	Noeske et al,
			2014
Oxazolidinones	Linezolid	Translation	Belousoff et al,
			2017
Phenicols	Chloramphenicol	Translation	Stavrianidi et al,
			2018
Fluoroquinolone	Ciprofloxacin	DNA replication	Hangas et al,
			2018
Pyrimidines	Trimethoprim	C1 metabolism	Holmes et al,
			2016
Lipopeptides	Daptomycin	Cell membrane	Müller et al, 2016
Cationic peptides	Colistin	Cell membrane	O'Driscoll et al,
			2018

The targets of the most commonly used antibiotic are peptidoglycan biosynthesis and bacterial ribosomes (70S ribosome). Peptidoglycan biosynthesis is affected through the inhibition of transglycosidases, the enzymes responsible for giving the cell wall strength through the cross-linking of peptidoglycan molecules; this results in the rupture of the bacterial cell wall. Antibiotics that target the bacterial 70S ribosome, the complex molecule responsible for the translation of mRNA into protein, act as the inhibitors of nucleic acids, which affects bacterial DNA replication.

For decades, the aforementioned antibiotics and other antimicrobial drugs have proven effective in reducing rates of illness and death from infectious diseases. However, the misuse and overuse of antibiotics has resulted in the emergence of pathogens possessing multi-antibiotic resistant genes (Van Boeckel et al, 2014), creating a significant health concern for modern medicine – antibiotic resistance.

1.2. Antibiotic Resistance: A Fast-Growing Global Health Problem

The successful use of any drug to treat infectious disease is determined by the potential for the bacteria to develop a tolerance or resistance to the drug from the time of the drug's introduction (Banin et al, 2017). This can be seen with antibiotics, which have shown continually decreasing efficacy since they were first discovered (Sabtu, 2015). Microbes that were once susceptible to antibiotics are becoming increasingly difficult to treat due to the development of a wide range of chemical and physiological mechanisms of resistance inside the bacterial cells (Davies & Davies, 2010).

Global efforts are necessary to prevent and control the development of further antibiotic resistance. In response to this urgent need, many research studies and reports have highlighted the severe threat of antibiotic resistance (Aslam et al, 2018). The Review on Antimicrobial Resistance (2014), commissioned by the UK Government and chaired by Jim O'Neill¹ reported that 50,000 lives are claimed each year by antimicrobial-resistant infection in Europe and the United States alone. In addition, it is expected that by 2050, antibiotic resistant pathogens will be the leading cause of mortality globally (Figure 1.1; Review on Antimicrobial Resistance, 2014).

¹ Jim O'Neill, a former chairman of Goldman Sachs Asset Management and a former UK Treasury Minister, is Chair of Chatham House.(https://www.project-syndicate.org)

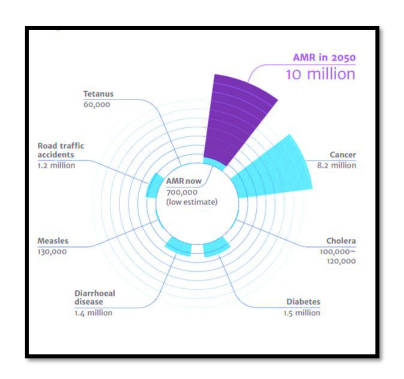


Figure 1.1 The expected global deaths in 2050.

The predicted causes of global deaths in by 2050. The number of people expected to die because of antibiotic resistance is more than the number expected to die due to cancer (Review on Antimicrobial resistance chaired by Jim O' Neill, 2014)

By 2050, 300 million people are expected to die prematurely because of antibiotic resistance, resulting in a 2-3.5% decline in the gross world product (GWP), which equates to the loss of between 60 and 100 trillion USD worth of economic output if the problem of antibiotic resistance continues (Review on Antimicrobial Resistance, 2014). The threat of antimicrobial resistance should concern every country regardless of its economic level. For example, in 15 European countries, methicillin–resistant strains have been found to be the main cause of 10% of all blood stream *Staphylococcus aureus* infections, and the resistance rates in some of these countries is as high as 50% (EARS-NE, 2013).

1.2.1. Overuse and Misuse as the Main Cause of the Antibiotic Resistance

Overuse or over prescribing of antibiotics is considered one of two main causes of antibiotic resistance in many pathogens, particularly in outpatient settings such as clinics and emergency departments (Ventola, 2015), where in parts of Europe 80% to 90% of antibiotic prescriptions are written by general practitioners (Llor et al, 2014). Globally, the consumption of antibiotics has increased in parallel with the increase in antibiotic resistance (WHO, 2009). There is also

high variation in the prevalence of antibiotic resistance between countries (WHO, 2018), which is likely caused by high variation in antibiotic consumption between countries. The WHO (2018), for example, reported antibiotic consumption rates across 65 countries varied between 1 and 2225 tonnes per year and the consumption of total antibiotics in Defined Daily Doses(DDD)² per 1000 inhabitants per day in these 65 countries ranged from 4.4 (in Burundi , Africa region) to 64.4 (in Mongolia ,Western Pacific Region) DDD per 1000 inhabitants per day.

Misuse of antibiotics, such as using inappropriate antibiotics to treat a particular type of infection, is considered the other main cause of antibiotics resistance (Viroj et al, 2018). The Centres for Disease Control and Prevention in the United States has reported that at least 30% of outpatient antibiotic prescriptions are unnecessary, meaning that they are prescribed for non-bacterial infections such as viral respiratory infections, or self-prescribed (CDC, 2016). Another type of antibiotic misuse is the prescribing of broad-spectrum antibiotics, which are classified as non-first line antibiotics, to treat infections that are susceptible to treatment by narrow-spectrum antibiotics. Typically, the prescripting of such broad-spectrum antibiotics occurs before confirming the specific type of bacterial infection (Llor et al, 2014).

1.2.2. The Evolution of Antibiotic Resistance

The resistance to certain antibiotics can exist naturally in some bacterial species. However, some susceptible bacteria can become resistant through either genetic mutations or through acquiring resistance genes from another bacterium (Lin et al, 2015).

Antibiotic resistance resulting from spontaneous genetic changes are rare and thought only to occur in about one in 1 million to one in 10 million bacterial cells (Long et al, 2016). Resistance can result from many different genetic mutations. Some mutations cause the target to which the antibiotic attaches to the bacterial cell to be revoked or cause the cell membrane pores through which the antibiotics enter the bacterial cell to close (Epand et al, 2016). Other mutations result in the production of enzymes that activate antibiotics compounds, such as β -lactamases (Palzkil et al, 2018). In addition, mutations can result in the production of a transport system that can export the antibiotic back outside of the bacterial cell, e.g. AmpC β -lactamase (Kohlmann et al, 2018).

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² Defined Daily Doses (DDD) is the assumed average maintenance dose per day for a drug used for its main indication in adults.

Antibiotic resistance can be acquired by the exchange of genetic material between resistant and susceptible bacteria in a process known as horizontal gene transfer (HGT) (Hong et al, 2018). Unlike vertical gene transfer, where genetic material is passed from parent to offspring, HGT involves genetic material being passed between bacterial cells of the same generation (Paquola et al, 2018). In this process, resistance genes are transferred in one of three vectors, either plasmids, transposons, or integrons, by one of three mechanisms, either transformation, conjugation, or transduction (Gyles & Boerlin, 2014).

Transformation is a process in which the genetic material carried by a bacteria (recipient) is altered by the incorporation of foreign (exogenous) DNA from another dead or degraded bacterium (Croucher et al, 2016; Figure 1.2). This process is considered a form of genetic recombination that occur predominantly between homologous DNA regions, which carry approximately the same nucleotide sequences (Johnston et al, 2014). The transformable or competent bacteria (recipient) is characterised by its ability to bind much more DNA than non-competent bacteria and to kill non-competent cells to release DNA for transformation (Blokesch & Melanie, 2016). *Streptococcus pneumoniae* and *Neisseria meningitides* are well-known of examples naturally competent bacteria (Bakkali, 2013).

Fragmented DNA from dead bacteria

Uptake fragmented DNA by competent bacteria

Incorporation by foreign (exogenous) DNA

Bacterial chromosome

Donor receipt

Transformed Bacteria

Figure 1.2 Bacterial transformation

Steps of Transformation; up taking of degraded DNA from dead bacterial cells by competent cell, followed by the incorporation of recipient DNA by foreign (exogenous) DNA.(modified from "conjugation by Adenosine [license; CC BY-SA 3.0])

The other mechanism of the Horizontal gene transfer (HGT) is conjugation. Here, the genetic material of chromosome from two sexually distinct types of cells are brought together through direct cell-to-cell contact (Grohmann et al, 2003). A partial gene from the donor bacteria moves indirectly through the pilus (Figure 1.3), which is produced by the donor bacteria and bridges the two cells, to complete gene transfer into the recipient bacterial cell (Shannon et al, 2015). This process is followed by the separation of the bacterial cells, after which the combined genetic material are altered further in the recipient cell (Vazquez et al, 2017). Some genetic material can be transferred in the reverse direction, from the recipient to donor bacteria, which is known as retro transfer (Sultan et al, 2018).

Plasmid Donor receipt

Figure 1.3 Bacteria conjugation.

Transferring of a partial gene from the donor bacteria to the recipient through pilus during the process of conjugation.(Modified from Zhang P et al,2015)

Bacterial genetic material, including resistance genes, can also be transferred horizontally by viral vector (bacteriophages), in a process known as transduction (De Sousa et al, 2018). In this process, bacteriophage bind to the bacterial cell and inject DNA into the cell through the cell membrane, whereupon small pieces of bacterial DNA can be packaged into the bacteriophage genome during the production of new phage particles (Casjens & Sherwood, 2003). The new phage particle is replicated instead of the bacterial genetic

material, which is degraded by the phage during this process (Harada et al, 2018). The bacteriophage then lyse the bacterial cell releasing phage carrying the packaged part of bacterial DNA. This will be injected into another bacterium (recipient) as part of the phage replication cycle (Rodríguez-Rubio et al, 2013; Figure 1.4).

Bacterial(host)
gene

Transduction

Phage replication

Lysis

Lysis

Phage carrying the packaged part of bacterial DNA

Figure 1.4 Bacterial Transduction.

The process of transduction which starts with the binding of the bacteriophage to the bacterial cell and the injection of its DNA into the bacterial cell. A small part of the bacterial DNA is packaged by the injected phage DNA, which then replicate inside bacteria instead the bacterial DNA, which is degraded by the phage. The host bacteria are then lysed and phage carrying the packaged part of bacterial DNA are released to bind the cell wall of other hosts (Modified from "conjugation by Adenosine.(modified from "conjugation by Adenosine [license; CC BY-SA 3.0]).

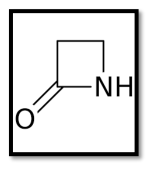
Transduction is classified into generalized transduction, in which any bacterial DNA can be transferred to another bacterium by a bacteriophage, and specialized transduction, in which a restricted set of bacterial genes is transferred to another bacterium (Chiang et al, 2019). The *mecA* gene³ carried by methicillin-resistant *S. aureus* (MRSA) is an example of a resistance gene that is transferred horizontally by transduction (Paterson et al, 2014).

Aforementioned evolutions of antibiotic resistance grows with the increasing number of bacterial species becoming resistant to the most commonly used antibiotics. A group of these antibiotics that represents approximately over 65% of global antibiotic use (Poole, K,2004) and are considered the largest class of bacteria cell wall synthesis inhibitors are beta-lactam antibiotics (Livermore, 1998).

1.3. Beta-Lactam Antibiotics

 β -lactam antibiotics are the most commonly used agents in the treatment of bacterial infections which are classified under the inhibitors of cell wall biosynthesis (Bush, 2018). They contain a four-atom ring (β -lactam) structure (Figure 1.5) and include penicillins, cephalosporins and carbapenems (Meletis, 2016).

Figure 1.5 β-Lactam ring.



 β -Lactam ring is a four-membered lactam. (Gilchrist T.1987).

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³ mecA is a gene found in bacterial cells that confers resistance to antibiotics such as methicillin, penicillin, and other penicillin-like antibiotics. mecA codes for PBP2a, which has a lower affinity for beta-lactams, which prevents cell death by maintaining the structural integrity of the cell wall.

1.3.1. The Main Types of βeta-Lactam Antibiotics

1.3.1.1 Penicillin (penams)

Penicillin is considered the most commonly used β -lactam antibiotic and was the first discovered antimicrobial drug. It was derived originally from a fungus "moulds" and used to treat the bacterial infection caused by *streptococci* and *staphylococci* (Thom, 1945). Penicillin V (taken orally), Penicillin G (taken intravenously) and procaine penicillin are different forms of penicillin (Bush et al, 2016). These forms share the three basic chemical compounds; a thiazolidine ring, β -lactam ring, and a side chain (Lobanovska et al, 2017) (Figure 1.6).

Figure 1.6 Penicillin Structure.

The chemical structure of Penicillin. Note the β -lactam ring; thiazolidine and side chain. (Gonzalez-Estrada A et al.,2015)

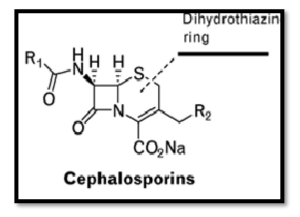
Based on chemical substitutions on the residue attached to the β -lactam ring, which confer different activities (Tahlan & Jensen, 2013), the penicillin family is categorised into five groups; natural penicillin, aminopenicillins, penicillinase-resistant penicillins, extended spectrum penicillins, and aminopenicillin/ β -lactamase inhibitor combinations (Miller & Lane, 2002).

Despite the wide range of pathogens that can be combated by penicillin the first sign of antibiotic resistance has appeared soon after the discovery of penicillin when penicillinase producing *E. coli* was detected by Abraham and Chain in 1940 (Sillankorva at el, 2019).

1.3.1.2. Cephalosporins

Cephalosporin is considered the largest group of related β -lactam antibiotics, which are characterised by their broad spectrum of activity (Bush et al, 2016). Cephalosporins are utilised to treat many infections including meningitis, bacteraemia, and skin and soft tissue infections (Thønnings et al, 2016). Structurally, it differs from penicillin by having a dihydrothiazin ring (Figure 1.7).

Figure 1.7 Cephalosporin Structure.



The chemical structure of cephalosporin. Note the β-lactam ring, dihydrothiazin ring, and side chain. (Feng H et al., 2017)

The large group of cephalosporins are categorised traditionally into five generations based on their antimicrobial spectrum and agents in the same generation are classified further according to their pharmacokinetics (Kim et al, 2014). First-generation cephalosporins exhibit higher activity with gram positive than that with the gram-negative bacteria, e.g. Cefazolin (Katzung et al, 2015). The second-generation cephalosporins, such as Cefoxitin, show the highest activity

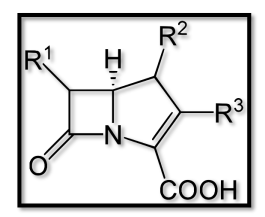
against gram-negative organisms and the lowest against gram-positive organisms (Bratzler et al, 2013).

The third-generation are the most prescribed cephalosporins and are the first generation to be considered as extended-spectrum cephalosporin. They are characterised by their greater resistance to the β -lactamase enzymes produced by gram-negative bacteria that are exhibited by first and second generations e.g. Cefotaxime and Ceftazidime which are considered the most commonly used antibiotics of this generation (Chaudhry et al, 2019). The fourth-generation retain the expanded activity of the third-generation against gram-negatives, in addition to an improved gram-positive spectrum, e.g. Cefpirome (Proticetal, 2016). The fifth-generation, which includes ceftaroline, is known as anti-methicillin-resistant *S. aureus* (MRSA) cephalosporins as it has high activity against gram-negative rods like MRSA and *Streptococcus* (Stevens et al, 2014)

1.3.1.3. Carbapenem

Carbapenems is the third main type of β -lactam antibiotics which is being used increasingly in recent years as a result of the rising the prevalence of cephalosporin resistant isolates (Hawkey et al, 2012). Among hundreds of β -lactam antibiotics, carbapenems are used as a last-line agent for the serious infections that cannot be treated by other β -lactams as it possess the broadest spectrum of activity and greatest potency against gram-positive and gram-negative bacteria (Papp-Wallace et al, 2011). The broad spectrum antibacterial activity of carbapenems is due to its a unique structure that is defined by a Carbapenem coupled to a β -lactam ring (Figure 1.8), which provides protection against most β -lactamases, such as metallo- β -lactamase (MBL), as well as extended spectrum β -lactamases (Codjoe et al, 2017).

Figure 1.8 Carbapenem Structure



The chemical structure of Carbapenem is defined as the 4:5 fused ring lactam of penicillin with double bond between C-2 and C3 but with the substitution of carbon for sulphur at C-1 (Papp-Wallace et al, 2011)

Based on their antibacterial activity, carbapenems are classified into three groups: Group 1, broad-spectrum carbapenems, such as Ertapenem, with limited activity against non-fermentative gram-negative bacilli; Group 2 broad-spectrum carbapenems, such as Imipenem, meropenem, and doripenem, with activity against non-fermentative gram-negative bacilli; and Group 3 which includes carbapenems that shows activity against methicillin-resistant *Staphylococcus aureus* (MRSA) and are currently under development (El-Gamal et al, 2017).

In addition to the three main β -Lactams, β eta-lactamase inhibitors confer approximately the same antimicrobial activity with an expanded spectrum of activity (Toussaint et al, 2015). These expanded spectra promote the potency of the antibiotic and provide resistance against the hydrolytic activity of the β -lactamase enzyme (Frère et al, 2005). Clavulanic acid is the most commonly used and effective β -lactamase inhibitor, which was produced as a result of natural product screening and was used initially to inhibit the activity of most plasmid penicillinases (Bush, Karen & Bradford, 2016)

The efficiency of β -lactam antibiotics is continuously challenged by resistant bacterial strains. This is most often due to the resistant cells, particularly non-fastidious gram-negative bacteria, producing an enzyme known as β -lactamase (Bush & Jacoby, 2010). β -lactamase—based

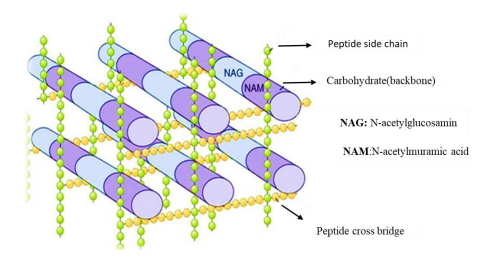
resistance cannot be understood without first knowing the mode of action of β -lactam antibiotics.

1.3.2. Beta-Lactam's Mode of Action

In all bacterial species a cell wall surrounds the bacterial cell, maintaining its shape and integrity, while also preventing cell lysis from high osmotic pressure (Yang et al, 2016). The cell wall is composed of a complex cross-linked peptidoglycan layer that contains repeated units of alternating disaccharides of N-acetyl glucosamine and N-acetylmuramic acids. A-5 amino-acid peptide is linked to N-acetylmuramic acid sugar and terminates in D-alanyl-D-alanine (Yadav et al, 2018).

In both gram-positive and gram-negative bacteria enzymes called penicillin-binding proteins (PBPs) act through a subgroup enzyme, transpeptidases (DD-transpeptidases), to remove the terminal alanine and catalyse the process of forming a cross-link with a nearby peptide (Rioseras, 2016; Figure 1.9). This cross-bridging mechanism is intimately intertwined with the discovery of the PBPs as it was concluded by some studies that the impact of penicillin on the synthesis of peptidoglycan in the transpeptidation reaction leads to the crosslinking of glycan chains is inhibited by B lactams (Penicillin) (Zapun et al, 2008).

Figure 1.9 Chemical structure of peptidoglycan.



Peptidoglycan structure; N-acetylglucosamine (NAG) and N-acetylmuramic acid (NAM) chains linked by peptide cross-bridges (Modified from clinicalgate.com).

PBPs are commonly classified into three main groups based on their molecular weight and domain structure; high molecular weight PBPs, which include two families called class A and class B, and low molecular weight PBPs, which consists of class C (Sauvage et al, 2016). All PBPs are characterized by the presence of three conserved motifs in the active site: SxN, SxxK, and KTG, where x is any residue (Nicola et al, 2010). The penicillin-binding domain that catalyses transpeptidation (transpeptidase) is constituted in the C-terminal region of class A PBPs (Welsh et al, 2017).

β-lactams inhibits the activity of transpeptidase by acting as a peptidoglycan precursor, which is targeted by the transpeptidases enzyme (Nikolaidis et al, 2014). The transpeptidation catalysed by the transpeptidase enzyme is completed in two steps. First, DD-trans peptidase interact with an acyl donor containing a pentapeptide stem ending in D-Ala-D-Ala, resulting in releasing of the second D-Ala, followed by an ester bond formation between the carbonyl of the first D-Ala and the catalytic serine residue of the enzymes. Second, the resulting acylenzyme is attacked nucleophilically by acyl acceptor generates the peptidoglycan cross-link (Edoo et al, 2017).

In the presence of β -lactam, the carbon carbonyl of β -lactams is attacked by the catalytic serine, similar to the first step of the transpeptidation reaction, which results in the breaking of the amide bond of the β -lactam ring and the inactivation of the D,D-transpeptidases (Vandavasi et

al, 2017) preventing the formation of the cross-link and leading to the lysis of the bacterial cell (Tidwell, 2008).

1.3.3 Resistance to βeta-Lactams

The emergence of resistant bacteria has reduced the effectiveness of β -lactams. There are three main resistance mechanisms for β -lactams (Figure 1.10): (1) alteration of the target to which the β -lactam antibiotics binds, which in turn changes the penicillin-binding protein and makes the β -lactam less effective in disrupting cell wall synthesis (Kocaoglu et al, 2015); (2) altering the porins and efflux pump system to reduce the amount of β -lactam that reaches the target; specifically, efflux transport increases the amount of the drug pumped out and porin mutations reduce cell membrane permeability to decrease the amount of drugs entering the cell (Masi, 2017); and, the subject of this thesis, (3) enzyme degradation of β -lactams, totally or partially, by the β -lactamase family of enzymes (Bonomo et al, 2017).

Antibiotic

Figure 1.10 Three mechanisms of antibiotic resistance against β -lactam.

Three main resistance mechanisms for β -lactam; alteration of the target, altering the porins and efflux pump system to reduce the amount of β -lactam and the enzymatic degradation (Todar, Ph.D., textbookofbacteriology.net).

1.3.3.1 β-Lactamases

β-lactamases are enzymes that hydrolyse the amide bond in β-lactam compounds and they are most frequently found in gram-negative bacterial isolates (Palzkill & Timothy, 2013). Abraham & Chain published the first paper identifying β-lactamase nearly 75 years ago. They documented the β-lactamase producing E. coli, but they were unable to extract the enzyme from these bacteria (Abraham & Chain, 1940). Four years later, Kirby successfully isolated the enzyme, which he called 'penicillin inactivator', and warned that its emergence could pose a significant clinical danger (Drawz & Bonomo, 2010).

The health concerns related to β -lactamase production has grown over the years as the number of unique, naturally occurring β -lactamases discovered has grown to more than 2,800 proteins (Bush, 2018). Treatments for many serious infections are still reliant on cephalosporins and carbapenems, which have continuously shown a declining efficiency over the years due to β -lactamase-based resistance (Shaikh et al, 2015).

1.3.3.1.1. β-Lactamase Classifications

Since the emergence of β -lactamase–based resistance, many attempts have been made to classify these enzymes. Richmond & Sykes (1960) proposed the first classification scheme. They based their classifications on whether the enzyme showed more hydrolytic activity against penicillin or cephaloridine, and whether it could be inhibited by P-chloromercuribenzoate and/or cloxacillin (Richmond & Sykes, 1973).

Jacoby & Bush (2010) have proposed another functional classification scheme in which the β -lactamases are categorised into four groups according to substrate and inhibitor profile: group 1 consists of cephalosporinase that are not inhibited well by clavulanic acid; group 2 is generally inhibited by active-directed β -lactamase inhibitors; group 3 is poorly inhibited by all classical β -lactamases inhibitors except EDTA and p-chloromercuribenzoate; and group 4 includes penicillinase that cannot be inhibited by clavulanic acid (Jacoby & Bush ,2010).

The most widely used classification is the sequence-based scheme, which was proposed by Ambler in 1980 and has been reviewed many times in the subsequent years. This classification divides β -lactamases into four classes (A, B, C, and D) based upon their amino acid sequences (Silveira et al, 2018). Ambler originally specified two classes: class A, whose activity depends on the presence of serine in the active site; and class B, which contain the metallo- β -lactamases that require zinc (Zn+ 2) for their activity (Palzkill &Timothy, 2013).

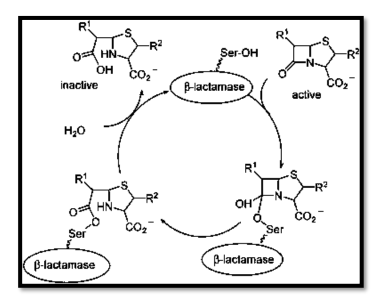
Class C, which is known as AmpC, and class D were found later and described as active-site serine β -lactamase but bore little sequence similarity to the serine class A enzyme (Öztürk et al, 2015).

A classification scheme based on functionality resulted in three major groups: group 1, cephalosporinase (class C); group 2, serine β -lactamase (class A and D); and group 3, metallo β -lactamase (class B) (White et al, 2017).

Group 1 Cephalosporinase: belonging to molecular class C, which are encoded predominantly on chromosome of *Enterobacteriaceae* (Ruppé et al, 2015). In terms of resistance, this group is more active on cephalosporins than benzylpenicillin and resistant to inhibition by clavulanic acid. Furthermore, this group shows a high affinity for aztreonam and a lack of activity on Cefoxitin (Bush & Jacoby, 2009).

Group 2 Serine β-lactamase: represents the largest group of β-lactamase and includes molecular classes A and D (Chandra & Seema, 2013). These enzymes are predominantly chromosomal, although some *staphylococcal* penicillinases are plasmid encoded (Livermore, 1995). The name of this group is derived from the role of serine in the active site of the enzyme's hydrolytic activity, which starts with the formation of a covalent Michael complex as a result of a non-covalent association between the enzyme and the β-lactam antibiotic (Brem et al, 2016). After this association, the active-site serine performs a nucleophilic attack on the carbonyl of the β-lactam, which results in a high-energy tetrahedral acylation intermediate (Lee et al, 2016). Next, as a result of the protonation of β-lactam nitrogen and the cleavage of the C-N bond, the tetrahedral acylation intermediate transitions into a lower-energy covalent acylenzyme, which is then attacked by activated water molecules and results in the formation of a high tetrahedral deacylation intermediate and hydrolysis (Meini et al, 2015). The bond between the oxygen of the nucleophilic serine and the carbonyl group of the β-lactam antibiotic regenerates the active enzyme and releases the inactive antibiotic. (Drawz & Bonomo, 2010) (Figure 1.11).

Figure 1.11 Serine β -lactamase.



General mechanism of hydrolysis of a β -lactam antibiotic by a serine β -lactamase (Touchet et al., 2011).

Group 3 MBLs: metallo- β -lactamases (MBLs) are chromosomal enzymes that are produced mainly from gram-positive bacteria and differ structurally from the other β -lactamases due to their requirement for zinc ions at the active site (Sun et al, 2018). Functionally, MBLs are characterised by their hydrolytic activity against carbapenems, which cannot be hydrolysed by serine β -lactamases (Lisa, 2017). Unlike serine β -lactamases, MBLs cannot be inhibited by clavulanic acid or tazobactam, but they are inhibited by metal ion chelators, e.g. EDTA (Patel & Bonomo, 2013). Furthermore, the class B β -lactamases utilises zinc to activate a water molecule and catalyse its direct addition to the β -lactam ring (Figure 1.12). This is unlike the class A, C, and D β -lactamases, which depend on opening the β -lactam ring via covalent acyl enzyme intermediates for their hydrolytic activity (Jeon et al, 2015).

Figure 1.12 Mechanism of hydrolysis by metallo-β-lactamases

The role of zinc ion in the mechanism of action of metallo-β-lactamases (Fernandes et al, 2013).

MBLs are classified structurally into three subclasses: B1, B2, and B3, which differ from each other in the consensus of amino acid at the zinc-active site (Table 1.2).

Table 1.2 MBL subclasses

	Number of	Consensus amino acids at active site		
β subclasses	active site Zn² atoms	Ligands to Zn ¹	Ligands to Zn ²	References
B1a	2	His116	Asp120	Garau & Getal, 2004
B1b	2	His118	Cys221	Young et al, 2009
		His196	His223	
		His116	Asp120	
B2	1	His118	Cys221	Ackerman et al, 2013
		His196	His263	
		Na	Asp120	
			Cys22	
			His263	
В3	2	His/Gln118	His121	Fre're et al, 2005
		His196	His263	

1.3.3.1.2. Main Types of β-Lactamase

There are three main types of β -lactamases that vary in their mechanism of action and in their antibiotic target (Madec et al, 2017), and each type is categorised under one of the aforementioned molecular classes. These types include:

(1) AmpC β -lactamase, which are classified as C class, are predominantly detected in Gramnegative organisms, e.g. E. coli and K. pneumoniae (Fenollar-Ferrer et al, 2008), and exhibit great hydrolytic activity against narrow and broad cephalosporinase and alpha-methoxy β -lactam, e.g. cefoxitin and aztreonam. In addition, they are poorly inhibited by clavulanic acid (Manoharan et al, 2012).

The *AmpC* gene is predominantly encoded on chromosomes, and here the production of the AmpC enzyme is known as inducible (as a result of exposure to inducible genes, e.g. penicillin, cefoxitin, and Carbapenem; Kaur et al, 2016), or it can be acquired by a transferable AmpC gene on a plasmid. This is termed plasmid-mediated AmpC β-lactamase (Coertze et al, 2017).

The resistance conferred by AmpC β -lactamase producing gram-negatives is enhanced by controlling the delivery of the β -lactam antibiotic to the periplasm. This is performed by the transport system in the outer membrane (Moreira et al, 2004), where the porin channels facilitate the penetration of β -lactam, and the porins and efflux pumps transport the β -lactam out of the bacterial cell (Pages et al, 2016; Figure 1.9).

The phenotypic detection of AmpC producing bacteria in clinical laboratories is always being improved as it can give a positive result with extended spectrum β -lactamase (ESBLs), the third main type of β -lactamases (Jacoby, 2009). The Three-Dimensional Test is still the most commonly used method to identify the AmpC producers. Here, extended-spectrum cephalosporin (Cefoxitin) is placed 3 mm from the inoculated slit with the test organism on a Mueller-Hinton agar. A positive result is indicated by the emergence of a distorted zone in the slit side (Gupta et al, 2014).

(2) Carbapenemases β -lactamases are characterised by their high hydrolytic activity against Carbapenem antibiotics, unlike the other β -lactamase types (Jeone et al, 2015). This ability is due to their distinct chemical structure: a 4:5 fused lactam ring with the presence of double bonds between C2 and C3 with the hydroxymethyl side (Nordmann & Poirel, 2002; Figure 8).

Based on the Ambler classification, carbapenemases can be categorised under three classes of β -lactamases; Class A and class D (Serine) carbapenemases β -lactamases and class B (zinc, metallo) carbapenemases β -lactamases (Codjoe et al, 2017). KPC, VIM, IMP, NDM and OXA-48 types are considered the most well know and effective carbapenemases (Poirel et al, 2012). Among the different carbapenemase-producing species, *Klebsiella pneumonia* (KPC) is among the most widespread carbapenemase producer and is one of the main causes of nosocomial infections (Van Duin et al, 2017).

The clinical identification of carbapenemase producers is based on the Modified Hodge Test (MHT) in which the tested isolates are streaked on lines away from Carbapenem disk (meropenem or Ertapenem) on Mueller Hinton agar, which has been inoculated with a Carbapenem-susceptible *E. coli* strain. A positive result in the MHT is indicated by a cloverleaf shaped growth of Carbapenem-susceptible *E. coli* around the streak line close to the disk of Carbapenem, which is permitted due to the decreasing concentration of the carbapenemase enzyme away from the disk (Tamma et al, 2018).

(3) The third main type of β -lactamases, which is also the most rapidly increasing type worldwide, is extended spectrum β -lactamases (ESBL). One of its subtypes is the subject of this study.

1.3.3.1.2.1. Extended Spectrum β-Lactamase (ESBL)

Extended spectrum β -lactamases are the third main type of β -lactamase and were first detected in Western Europe during the mid-1980s. Since then, their prevalence has been increasing steadily and has been associated with the widespread use of cephalosporins (Kim et al, 2017). The resulting mutation of the *TEM1*, *TEM2*, and *SHV* genes leads to the alteration of the amino acid configuration around the active site of the β -lactamase enzyme and, thus, ESBL production (Bajpai et al, 2017).

ESBL-producing bacteria exhibits resistance to most β -lactam antibiotics, including third and fourth generation cephalosporins (Rahmanetal, 2018), which severely limits the range of treatments available for infections caused by ESBLs (Blaak et al, 2014). However, these ESBL-producing strains can be inhibited by β -lactamase inhibitors, such as clavulanic acid, tazobactam, and sulbactam (Rawat & Nair, 2010).

The ESBL isolates were initially detected in hospital infections caused by *Klebsiella* pneumonia, but are now also associated with community acquired infections, particularly

urinary tract infections caused by ESBL-producing *E. coli* (Koksal et al, 2019). In many parts of the world *E. coli* and *Klebsiella pneumoniae* represents 10-40% of the detected ESBL producers (Rupp & Fey, 2003).

ESBL enzymes are predominantly plasmid encoded, and are characterised by carrying genes encoding resistance to other antibiotics, such as aminoglycoside, which may result in further problems in the treatment of infections caused by ESBL-producing strains (Bush & Jacoby, 2010). This problem therefore requires the use of appropriate antibacterial agents, which could be unachievable due to misleading of phenotypic detections used in clinical laboratories (Poulou et al, 2014).

1.3.3.1.2.1.1. ESBL Types

Many classifications of ESBLs have been introduced since the first emergence of this enzyme. The most common classification is based on the type of β -lactam that is hydrolysed by the ESBL type. The most important types are: SHV, TEM, OXA, and CTX-M.

SHV ESBL

The name of this group is derived from the sulfhydryl variable, as it was thought that the active site of the enzyme can be bound with P-chloromercuribenzoate by sulfhydryl⁴ to inhibit the activity of cephaloridine (Drawz et al, 2010). Since then, it has been reported that the active site of SHV type β -lactamase is serine hydroxyl sulfhydryl (Manageiro et al, 2012). Based on the study conducted by Chaves et al (2001), the most well-known SHV-producing bacteria are found in the *Enterobacteriaceae* family, which includes *Klebsiella pneumonia* where SHV-product is detected with a frequency of 80-90%.

Genetically, all SHV variants are derived from the *bla*SHV-1 gene, which is generally considered a plasmid-encoded enzyme with one to seven amino acid substitutions (Drieux et al, 2008). The premature protein consists of 286 amino acids (Thai et al, 2010), and the mature enzyme is yielded by removing the 21 amino acids at the N-terminus from the signal sequence (Kuzin et al, 1999). The *bla*SHV gene encoding has mutated rapidly and has transferred to other gram-negative bacteria in different geographical regions. Jacoby & Bush (2012) have described these as the 117 SHV variants.

⁴ The sulfhydryl group is a family of organic compounds that contains an R group bound to a sulphur atom and a hydrogen atom that play an important role in biochemical processes and participate in enzymic reactions for the formation and transfer of acyl residues.

In resistance terms, initially SHV ESBL has conferred great hydrolytic activity against penicillin and first generation cephalosporins (Matthew et al, 1973). This hydrolytic activity has expanded over the years to inactivate extended spectrum cephalosporins and monobactam by some structural mutations in the *blaSHV-1* gene (Bush & Jacoby, 2010).

Based on functional properties and molecular characteristics SHV ESBL can be categorised in three main subgroups: subgroup 2b, which has ability to hydrolyse penicillin and only early cephalosporins, e.g. cephalothin and cephaloridine, and can be inhibited by clavulanic acid; subgroup 2br, which confers a broad-spectrum of activity against β -lactamase inhibitors including clavulanic acid; and subgroup 2be, which include SHV β -lactamase and includes enzymes that show hydrolytic activity towards oxyimino β -lactams; cefotaxime, Ceftazidime, and aztreonam (Liakopoulos et al, 2016).

TEM ESBL

TEM gene was first detected in 1965 in *E. coli* isolates (Bonardi et al, 2018). Over 100 TEMs have been discovered and the majority of these are ESBLs (Paterson & Bonomo, 2005). The most effective subtypes of this enzyme include TEM1, TEM2, which structurally substitute lysine for glutamine at position 39 (Grigorenko et al, 2018), and TEM3. TEM1 is capable of hydrolysis with penicillin and first generation cephalosporins, but it is unable to inactivate the oxyimino cephalosporin (Palzkill, 2018). Ninety percent of resistance against penicillin in *E. coli* is due to TEM1 production (Livermore, 1995).

TEM2, which is not categorised as an ESBL, has the same hydrolytic profile as TEM1, but it has a more active native promoter and differences in its isoelectric point (5.6 compared to 5.4; Jain et al, 2008). TEM3, which was originally named CTX-M-1 due to its activity against cefotaxime, is considered the first TEM variant with increased activity against extended spectrum cephalosporins (Rupp & Fey, 2003).

Some TEM derivatives confer negligible hydrolytic activity against extended spectrum cephalosporins and show low affinity for some β -lactamase inhibitors. These TEM derivatives are not classified as ESBL enzymes (Paterson & Bonomo, 2005). However, some recovered mutants in TEM β -lactamase, which are described as complex mutants, develop the hydrolytic activity of this enzyme against third generation cephalosporins and inhibitors, such as clavulanate (Ruppé et al, 2015).

OXA ESBL

The name of the OXA enzyme is derived from its oxacillin-hydrolysing abilities (Scriver et al, 1994), as it confers a hydrolysing rate for oxacillin and cloxacillin that is 50% greater than that of benzyl penicillin (Bush et al, 2010 .The most common OXA subtypes have been detected predominantly in *Pseudomonas aeruginosa* and *E. coli* (Antunes et al, 2014). Unlike other ESBL types, most OXA subtypes do not confer hydrolytic activity against extended spectrum cephalosporins to a significant degree (Santillana et al, 2007).

The different subtypes of the OXA enzyme are classified into two groups according to the hydrolytic activity against cefotaxime, ceftriaxone, and aztreonam. Group (1) includes OXA-11, OXA-14, OXA-16, OXA-17, OXA-19, OXA-15, OXA-18, OXA-28, OXA-31, OXA-32, OXA-35, and OXA-45, which all confer frank resistance against the three aforementioned antibiotics. Group (2) includes one member, OXA-10, which shows weak hydrolytic activity against the same antibiotics. However, it is described as a Carbapenem-hydrolysing enzyme (Toleman et al, 2003).

Most members of Group 1 OXA do not originate as ESBLs, but become ESBLs through mutations derived mainly from the OXA-10 ESBL (Bradford, 2001). The sequencing of any new OXA variant enzyme shows two amino acid changes from the OXA-10 enzyme, altering an arginine at amino acid 143 to serine and a glycine at position 157 to aspartate (Evans & Amyes, 2014). OXA-11 is considered the first example of an OXA enzyme that shows alteration from non-ESBL to ESBL characteristics as a result of OXA-10 mutations (Paterson & Bonomo, 2005).

The fourth type of ESBL β -lactamase enzymes, which has become the most common and rapidly growing health concern, is CTX-M β -lactamase. The epidemiology of different variants of this enzyme in North Wales and studying its activity characteristics are the main subjects of this study.

1.3.3.1.2.1.2. CTX-M Enzyme: The Most Rapidly Increasing Type of ESBLs

The name of the CTX-M enzyme is derived from cefotaxime antibiotics, which can be hydrolysed by CTX-M β -lactamase with a minimum inhibitory concentration (MIC) of less than 64 μ g/mL (Baraniak, 2002). The CTX-M has become the predominant type of ESBL detected in many regions of the world (Zhao & Hu, 2013). Potz et al (2006) studied 16 British hospitals to determine the different ESBL types contributing to antibiotic resistance and found that out of 19,252 clinical isolates detected over a 12 week period, CTX-M-producing strains

accounted for 1.7%, higher than the other ESBL-producing strains that were predominantly detected in *E. coli* and *Klebsiella* spp. The concern is that these enzymes have contributed to the increased detection of resistant *E. coli* in outpatients and patients admitted in hospitals for short periods of time, which suggests the community acquisition of these strains (Canton, 2006).

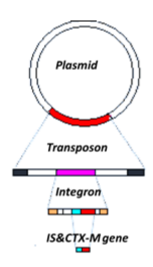
Functionally, based on Kinetic studies the CTX-M-type β-lactamases show higher hydrolytic activity against cephaloridine or cephalothin more than against benzyl penicillin, and they preferentially hydrolyse cefotaxime over Ceftazidime (Shaikh et al, 2015). The CTX-M enzyme's ability to inactivate cefotaxime is approximately 35 times greater than against Ceftazidime, which is inactivated with more efficiency by TEM and SHV cefotaxime (Bonnet, 2004). Furthermore, compared to TEM and SHV, CTX-M has the highest hydrolytic activity against the narrow-spectrum cephalosporins (Drieux et al, 2008). However, since the emergence of CTX-M, the most effective inhibitor for this enzyme has been the combination of tazobactam and penicillin (Bush & Bradford, 2016).

The bla CTX-M Gene and Hydrolytic Activity

Unlike other ESBL types, SHV, TEM and OXA, which are generated by substitutions of amino acids in their parent enzymes the different subtypes of CTX-M ESBL are acquired by a horizontal gene transfer between bacteria in which the CTXM gene is carried on either plasmids or transposons (Knudsen et al, 2018). In comparing the genes inside the cefotaxime-susceptible *Kluyvera spp* with the same genes inside the resistant one, it has been observed that the resistant species contain genes showing weak expression for β -lactamase, and they can be inhibited by cefotaxime. These genes, known as *bla*KLU, are found to acquire the CTX-M phenotypes via an insertion sequence (IS; Rodríguez et al, 2004).

ISs are the smallest elements that can transpose independently in the organism, and they can cause genome rearrangements and insertion mutations (Mahillon & Chandler, 1998). One of the main roles of the IS is to move IS mobilised genes among transposons, plasmids, integrons and chromosomes, and this increases the possibility of transferring the resistant determinant (Zhao & Hu, 2013). ISEcp1, which was detected in the plasmid pST01 carried by *E. coli*, is considered the most common insertion sequences and is characterised by having a high-level expression promoter for CTX-M (Bevan et al, 2017). These insertion sequences lie adjacent to the *blaCTX-M* gene and are present in transposon, which is integrated with a plasmid (Figure 1.13; Lartigue et al, 2004).

Figure 1.13 Plasmid-encoding extended-spectrum β -lactamase CTX-M.



blaKLU acquire the CTX-M phenotypes via an insertion sequence (ISEcp1), which is characterised by its high-level expression promoter for CTX-M gene (Modified from Cantón,et al.,2012)

The *blaCTX-M* gene is predominantly encoded on large plasmids that tend to carry additional resistance genes, such as those exhibiting resistance to fluoroquinolones and aminoglycosides. These plasmids range in size from 7 to 430 Kb (Cao et al, 2002). The *blaCTX-M* gene is mainly encoded on the incompatibility group FII plasmids, which are characterised by low copy numbers and are known as epidemic plasmids due to their potency to readily acquire and disseminate the resistance genes in *Enterobacteriaceae* members (Zhuo et al, 2013). The most well-known examples of these plasmids are the pandemic dissemination of CTX-M-15, favoured by IncFII plasmids (Irrgang et al, 2017). Unlike most other resistance-carrying plasmids, FII plasmids are thought to be widely distributed and adapted to their hosts even before exposure to antibiotics (Partridge et al, 2011).

In addition to FII plasmids, there are many other plasmids involved in the dissemination of different CTX-M types, including: the IncN plasmids that has been shown to contribute to the

dissemination of CTX-3 in Poland (Baraniak et al, 2002); the F33:A-:B-type plasmids encoding CTX-M-65 were first detected in China (He et al, 2013); and the IncK plasmids were responsible for the dissemination of CTX-M-14 in the United Kingdom (Stokes et al, 2012).

Although *the blaCTX-M* gene is encoded predominantly in plasmids, the chromosomal *blaCTX-M* has been reported in some studies worldwide. In Korea, Song et al (2011) detected chromosomal *CTX-M* genes in 21 out of 25 *P. mirabilis* isolates. Furthermore, the chromosomal *blaCTX-M-*25 and *blaCTX-M-*41 have been detected in *P. mirabilis* isolates in Israel (Navon-Venezia et al, 2008).

The *blaCTX-M* gene encodes 291 amino acids and a single change of an amino acid constitutes a new CTX-M type (Giedraitienė et al, 2017). Unlike other ESBLs, this enzyme does not enlarge the active site to accommodate the antibiotics, but instead its hydrolytic activity depends on point substitutions (Taylor & Francis, 2015). The Ceftazidime antibiotic protects itself from the hydrolytic activity of the CTX-M enzyme by having a C7 β-amino thiazoloxyimino-amide side chain compound, which can only be hydrolysed by Asp 240Gly CTX-M (Gwon et al, 2019). The resistance to Ceftazidime is frequently associated with mutation at amino acid 167 and 240 (Delmas et al, 2008).

CTX-M-Producing Isolates Causing UTI: Limitation of Phenotypic Detection

Urinary tract infections (UTIs) are considered one of the most infectious of diseases in both the healthcare setting and in the general community (Beahm et al, 2017). Every year 150 million people around the world are diagnosed with a UTI (Flores-Mireles et al, 2015) and approximately 40% of women worldwide develop an infection of urinary tract at least once in their lifetime (Micali et al, 2014). The estimated cost resulting from the eleven million people are diagnosed with UTI annually in US is five billion dollars (Foxman, 2014).

UTI are classified generally into two types; Complicated UTIs predominantly occur in elderly people, particularly among post-menopausal women and men with prostatic hypertrophy, and are associated with serious complications, including antibiotic resistance (Pallett & Hand, 2010). Uncomplicated UTIs typically occur across age groups of both sexes (Magliano et al, 2012) and are mostly associated with functional or structural abnormalities of the urinary tract such as obstructions, renal failure, and pregnancy (Durojaiye & Healy, 2015).

In addition to the dysfunction of urinary tract a high residual volume (RU) is considered another risk factor that can lead to the development of a UTI, even without any symptoms in the lower

urinary tract (Cho & Kim, 2010). Adult male patients with residual urinary volume more than 180 ml are more likely to have bacteriuria (Jhang & Kuo, 2017). However, the role of RU in the development of UTIs in females is controversial (Truzzi et al, 2008). In addition, behavioural risk factors can contribute in recurrent UTIs among females, of which sexual intercourse is considered the main risk factors, particularly for young women (Mishra et al, 2016).

Uropathogenic E. coli (UPEC) as a Main Cause of UTIs

There are a broad range of pathogens that cause UTIs, but uropathogenic *E. coli* (UPEC) remains the most common, representing 80-90% of pathogens associated with community acquired UTIs (Noormandi & Dabaghzadeh, 2015). UPEC is continuously developing resistance against the most commonly used antibiotics (Flores et al, 2015). UPEC is classified into four main phylogroups (A, B1, B2 and D) based on the occurrences of genomic pathogenicity islands (PAI) and the expression of virulence factor acquired by *E coli* to cause infections of urinary tract, such as toxins, adhesins, iron-acquisition system and surface polysaccharides (Hannan et al, 2012).

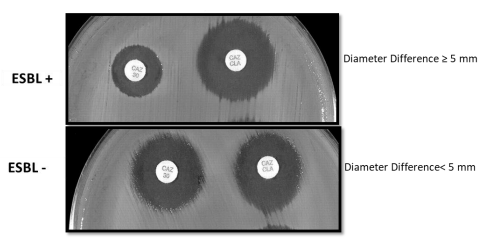
UTIs can be caused by other pathogens, although less frequently than UPEC, including *K. pneumonia*, *P. mirabilis*, *Enterococcus faecalis*, *Streptococcus bovis*, and *Pseudomonas aeruginosa*, in addition to the fungus Candida (Palou et al, 2013).

Role of CTX-M Type ESBL-Producing E. coli in UTI

The detection of resistant *E. coli* in UTI patients has increased mainly due to the spread of ESBL-producing bacteria (Laupland et al, 2008). Before 2003 these were TEM and SHV penicillinase-producing bacteria, however they are now being replaced by CTX-M-producing *E. coli*, which is the strain most often acquired in hospitals (Livermore, 2005).

In clinical diagnostic laboratories, the detection of ESBL is primarily based on phenotypic testing, e.g. a double-disc synergy test (Figure 15), which is considered the crucial step in the appropriate management of the patient. This test is highly cost-effective and easy to use (Woodford & Sundsfjord, 2005), but phenotypic methods are unable to distinguish between the different types of ESBLs or to determine the gene responsible for the production of each enzyme. This results in treatment failures in many cases due to the use of inappropriate antibiotics, and to the outbreak of multi-resistant gram-negative pathogens that require expensive control efforts (Tenover, 2007).

Figure 1.14 Combination disc method.



Combined disc diffusion test used to identify ESBL producing bacteria. CAZ = Ceftazidime and CLA = clavulanic acid. Note the difference in the diameter between the two discs, which is the determinant if the test is positive (≥ 5 mm) or negative (Modified from Kumar et al, 2012).

This study attempts to respond to this need by using molecular-based techniques for ESBL detection with the additional ability to detect low-level genetic resistance. With these techniques, it would be possible to determine the contribution of the most rapidly increasing type of ESBL (CTX-M) to the resistance conferred by the causative pathogens of urinary tract infections. In addition, this study will create an epidemiological snapshot of the antibiotic resistance pattern based on CTX-M enzyme production in UTI patients in North Wales.

The samples of the study are ESBL-producing bacteria isolated from 300 urine samples of UTI patients in three referral hospitals in North Wales: Ysbyty Gwynedd, Glan Clwyd Hospital and Wrexham Maelor Hospital.

The research question of this study is to determine to what extent CTX-M-type ESBLs contribute to the resistance phenomenon detected in isolates from UTI patients in North Wales, and whether these CTX-M-producing bacterial isolates carry the same or different CTX-M genes.

1.4. Objectives

This research aims to study the antibiotic resistance of bacterial species isolated from urine of urinary tract infection (UTI) patients at three referral hospitals in North Wales in order to:

- (1) Determine the role of CTX-M-type ESBL β -lactamases in the antibiotic resistance of UTI patients in North Wales.
- (2) Search for novel CTX-M-type gene variants.
- (3) In vitro characterisation of recombinant CTX-M-type β -lactamases with novel mutations, including determination of posttranslational modifications (e.g. phosphorylation status) and enzyme kinetics.

Chapter 2 : MATERIALS AND METHODS

Overview

The aims of the study were reached through five main steps. First, samples and data were collected, including patient ages, gender as well as phenotypic resistance to various antibiotics of clinical samples. Second, genes of interest were screened using multiplex PCR amplification of β-lactamase genes *blaCTX-M 1, 2, 9* and *8/25* followed by gel electrophoresis and sequencing. Third, the mutant *CTXM* gene was cloned into an *E.coli* expression plasmid allowing for the purification of a C-terminally strep-tagged recombinant protein. Fourth, protein expression and protein purity were verified using western blotting technique and fifth the minimum inhibitory concentration (MIC) and kinetic constants of induced strep-tagged recombinant proteins were determined against selected antibiotics.

2.1 Microbiology—Cultivation

2.1.1 Sample collection

The samples of the study were Extended Spectrum Beta Lactamase (ESBL)-producing bacteria which were isolated from patients with urinary tract infections and whose specimens were submitted to three referral hospitals in North Wales: Ysbyty Gwynedd, Glan Clwyd Hospital and Wrexham Maelor Hospital (Figure 2.1). One hundred clinical samples of ESBLs were obtained from each hospital when the study started in 2014. The collected data of the study included age, gender and phenotypic resistance to various antibiotics.

Figure 2.1 Location of the three referral hospitals of North Wales.



Map created by geography department, Bangor University, shows the location of the three-referral hospital in North Wales, UK. Ysbyty Gwynedd, Glan Clwyd Hospital and Wrexham Maelor Hospital.

2.1.2 Sub culturing of bacterial isolates

The isolates were all sub cultured from their agar slopes on the primary UTI Agar. In order to identify *Enterococcus spp.*, *Escherichia coli*, and *coliform* bacteria, samples were cultured on the primary UTI Agar (#2421517).

The UTI Agar (formerly Chromogenic UTI Agar) is a chromogenic medium for the presumptive identification and differentiation of all the main micro-organisms that cause urinary tract infections (UTIs) and contain a chromogenic substrate that is hydrolysed by enzymes produced by these bacteria. The β -D-glucosidase produced by *E. coli* hydrolyses the chromogenic substrate to produce pink-red colonies, and the β -glucosidase from coliform bacteria hydrolyses the chromogenic substrate to produce blue colonies (Oxoid Ltd, 2008).

The isolates were sub cultured using a plastic loop for the streak dilution method to achieve single colonies, which ensures the purity of the isolates to carry out the multiplex PCR assay. The plates were then incubated at 37°C for 18–24 h, and the growth of the colonies was

recorded. Two types of colonies were observed in the cultured samples: pink colonies (indicating the presence of *E. coli*) and blue colonies (*coliform bacteria*).

2.1.3 Identification of isolates using the Bruker Daltonik MALDI Biotyper technique

Principle: Bacterial identification by matrix-assisted laser desorption/ionisation (MALDI) is based on the use of mass spectrometry to measure the protein content of bacteria, using the mass-to-charge ratio of the proteins. The bacteria are exposed to an ion source to ionise and separate their proteins based on their mass-to-charge ratio, and the mass spectrum generated is compared against a database of mass spectra. Of note, the proteins extracted by the lysis solvantin acidic conditions are mostly basic cytoplasmic proteins, specifically mitochondrial, ribosomal and RNA binding proteins, which all have a range of 4000 to 15000 DA (Ryzhov V. & Fenselau C, 2001).

Sample preparation: First, the bacteria are extracted to generate sufficient spectra. The isolate is dissolved in 50 μ L of 70% ethanol, placed in a micro centrifuge and centrifuged at 20,800 rpm for 10 min. The obtained supernatant is discarded, and the cell pellet is dried and mixed with acetonitrile and 50 μ L of 70% formic acid before being centrifuged for 2 min at 20,800 rpm. Then, 1 μ l of the obtained supernatant is placed in the target plate, dried and loaded with 2 μ l of the matrix. Pre-treatment is beneficial because it inactivates the organisms and enhances detection of biomarkers above 15 KD.

Once the samples are loaded, they are introduced into a high-vacuum environment in which they are ionised by a precise laser release of a cloud of protein that is accelerated by an electric charge. The speed at which the proteins unravel is based on their weight: the heavier the protein, the slower it travels. The time of flight (TOF) is recorded, and when the proteins reach the end of the path, a mass spectrum is created for each sample by comparing its obtained spectrum against a library of mass spectra for different bacteria that can be identified at three levels: species, genus and family.

2.1.4 Microbanking the isolates

All isolates were maintained in Microbank cryovials for long-term storage and retrieval (Figure 2.2). Each MicrobankTM vial contains approximately 25 sterile coloured beads (single colour) and specially formulated cryopreservative. The specially treated beads are of a porous nature which allows microorganisms to readily adhere onto the bead surface. After inoculation, the MicrobankTM vials are kept at -80°C for extended storage. When a fresh culture is required, a

single bead is easily removed from the MicrobankTM vial and is used to directly inoculate a suitable culture medium (White & Sand., 1985).

2.2. Molecular Analysis

Genotype detection was used to determine the dominant *bla*CTX-M genes responsible for CTX-M-type-ESBL production among *bla*CTX-M groups 1, 2, 9 and 8/25, which were detected between UTI patients. CTX-M-type-ESBL genotypic detection was conducted using the following three procedures: preparing cell-free lysate, amplifying the PCR of *bla*CTX-M genes with oligonucleotide primers and screening these genes using gel electrophoresis and visualization of the gel strips using Quantity One software followed by sequencing.

2.2.1 Preparing cell-free lysates

The isolates of interest were lysed to extract their genomic DNA, which was used as the DNA template. The lysis solution was prepared by making a solution of 0.125 g of 25% sodium dodecyl sulphate (SDS) and 1 g of 0.05% sodium hydroxide (NaOH) dissolved in distilled water to make a volume of 50 ml. A small amount of sub-cultured bacteria from each plate was scraped with a sterile pipette tip, which was dipped into a PCR tube containing 20 μ L of lysis buffer and labelled before loading into the PCR machine.

The following protocol was used: the samples were heated at 95° C for 15 min, after which they were maintained at 10° C until the machine was turned on. Once the samples were unloaded from the PCR machine, $180 \,\mu\text{L}$ of sterile water was added to each sample, and each was stored at 20° C until it was subject to PCR amplification.

2.2.2. The primer designs

The primers used in this study for blaCTX-M groups 1, 2, 9 and 8/25 were designed using the following specifications to produce accurate results: length of (19–22) nucleotide, base composition of 50–55% (G-C) and average melting temperature of 52–60°C. Computer software was used for the primer design, such as OLIGO 6, which guarantees minimizing the probability of stable primer-dimer formation (Table 2-1).

2.2.3. Primer preparation

The primers were purchased from the Eurofins MWG Operon Company. All primers were diluted to a concentration of 100 pmol/ μ L, as indicated in the manufacturer's instructions. The working solution was prepared by the further dilution of 100 pmol/ μ L solution to 10 pmol/ μ L (1:10 dilution).

2.2.4. Multiplex PCR amplification of β-lactamase genes blaCTX-M 1, 2, 9 and 8/25

The β -lactamase genes blaCTX-M 1, 2, 9 and 8/25 were amplified using a multiplex PCR. A total volume of 20 μ L of the PCR reaction mixture was prepared for each cell lysate sample. The master mix for the multiplex PCR included 10 μ L of Biomex red PCR, 2 μ L of forward primers (0.5 μ L of the specific primer for each of the groups 1, 2, 9 and 8/25); 2 μ L of reverse primers (0.5 μ L of the specific primer for each of the groups 1, 2, 9 and 8/25); and 5 μ L of water to create a final volume of 19 μ L, which was added to 1 μ L of bacterial cell lysate. The 3 published primer was used in the multiplex PCR of blaCTX-M genes, as shown in Table 2.

Table 2.1 Primer sets used in the PCR amplification of blaCTX-M genes

β-lactamase gene target	Primer sets	Sequences (5'-3')	Amplicon (bp)	Reference
	MultiCTXMGp1- for	TTAGGAARTGTGCCGCTGYAb/		Dallenn
CTX-M	MultiCTXMGp1-2 Orev	CGATATCGTTGGTGGTRCCAT ^b	688	et al, 2010
group 1				
	MultiCTXMGp2-for	CGTTAACGGCACGATGAC ^b /		Dallenn
CTX-M	MultiCTXMGp1-2 Orev	CGATATCGTTGGTGGTRCCAT ^b	404	et al, 2010
group 2				
	MultiCTXMGp2-for	TCAAGCCTGCCGATCTGGT/		Dallenn
CTX-M	MultiCTXMGp1-2 Orev	TGATTCTCGCCGCTGAAG	561	et al, 2010
group 9				
	CTX-Mg8/25 for	AACRCRCAGACGCTCTACb/		Dallenn
CTX-M	CTX-M8/25 rev	TCGAGCCGGAASGTGTYAT ^b	326	et al, 2010
group 8/25				

The positive control (*E. coli* ESBL ATCC51446) was included in the assay. All the labelled PCR tubes were mixed briefly and centrifuged at 13,000 rpm for 30 s before they were arranged in the PCR thermo cycler, which was set in the BLAM program. The conditions of the multiplex PCR were set according to the method published by Dallennet al. (2010) as shown in Table 2.2.

Table 2.2 The multiplex PCR program

Heated lid	105 °C	1 min
Denaturation	94 °C	40 sec
Annealing	60 °C	40 sec
Extension	72 °C	1 min
Final extension	72 °C	7 min
Number of cycles	30	-

The amplified PCR products were run on agarose gels.

2.2.5 Agarose gel preparation

A 2 % agarose gel was prepared as follows: 100 mL of TBE buffer (1X) was added to 900 mL of dH2O to achieve a final volume of 1000 mL of diluted buffer. Then, 3 g of agarose powder (#9012-36-6) was weighed in a clean flask and 150 mL of diluted buffer was added. The dissolved mixture was then boiled in a microwave for 3–5 min and cooled to 45°C before 15 μ L of nucleic acid dye (ethidium bromide) was added; the solution was poured into a gel mould once the gel was solidified. The set gel was placed in the BIO-RAD tank and filled with 500 mL of 1x TBE (Tris-borate-EDTA) buffer.

2.2.6 Samples loading on agarose gel

The first lane of the gel was loaded with 5 μ L aliquot of 100 bp ladder (Promega) 5 μ L of PCR products. And positive were loaded into the appropriate wells of the gel. After 45 min of electrophoresis, the PCR products were visualised using the Doc200 machine (Bio-Rad) and Quantity One 1-D analysis software (Bio-Rad).

2.2.7 DNA purification

Separating PCR products on agarose gel, the bands of interest were cut under UV light and transferred into an Eppendorf tube in order to purify its DNA using ISOLATE II PCR and Gel Kit (Bioline) (#IS494-BO37210) based on the manufacturer's instructions.

2.2.8 Sequencing the gene-of-interest using the pJET-1.2 forward primer

Samples containing successful recombinant plasmids were sequenced using the pJet-1.2 forward primer, one of Thermo Scientific's sequencing primers that has single-stranded oligonucleotides with 5'-hydroxyl and 3'-hydroxyl ends. The pJET1.2 sequencing primers

flanked the Eco32I site in the eco47IR gene of positive selection cloning vector pJET1.2. All primers were supplied as 10 µM aqueous solutions (Thermo Scientific, n.d.).

Samples containing successful recombinant plasmids were sequenced using a pJET plasmid primed with Thermo Scientific sequencing primers that have single-stranded oligonucleotide with 5'-hydroxyl group and 3'-hydroxyl ends. All primers were supplied as 10 µM genes of positively cloned victors. Flank the *E.coli* site.

2.2.9 Cloning the gene of interest into an *E.coli* expression pASK-IBA2C plasmid

Genes of interest were further analysed upon cloning of the mutant gene into an *E.coli* expression plasmid, pASK-IBA2C, which allows for the purification of strip-tagged recombinant proteins.

To clone genes of interest, full-length genes were amplified from the corresponding samples with the forward and reverse primers (Table 2.3). Both primers contain the BsaI restriction site, which is located in the multiple cloning site of the plasmid (Figure 2.2). The trick is that the enzyme generates an asymmetric cut with long sticky ends (GGTCTCN1 on the forward strand and CCAGAGN5 on the reverse strand). Since the enzymes cut five nucleotides away from the recognition site and because these sequences are different at either site of the two BsaI cuts in the plasmid, the plasmid cannot re-ligate, and only the cloned fragments should give colonies.

Table 2.3 Primer sets

Primer sets	Sequences (5'-3')
CTX-M-14 like-F	atggcataat ggtctc a ggcc ATGGTGACAAAGAGAGTGCAACGG
CTX-M-15-F	atggcataat ggtctc a ggcc ATGGTTAAAAAATCACTGCGCCAG
CTX-M-14-like-R	atggcataat ggtctc a gcgct CAGCCCTTCGGCGATGATTC
CTX-M-15-R	atggcataat ggtctc a gcgct CAAACCGTCGGTGACGATTTTAG

Nool BsaI pASK-IBA2C EcoRI 3061 bp Sat Konl Multiple **OmpA** BamHI cloning site Strep-tag Xhol Soll Cam R Pst BsaI Nool Eco47III HindIII

Figure 2.2 Features of pASK -IBA2C plasmid.

pASK-IBA2C plasmid features: Promoter (37 to 72 bp) ,tet-repressor (1893 to 2516 bp) ,OmpA signal sequence (139 to 201 bp),multiple cloning site (232 to 318 bp),Strep-tag® from (202 to 231bp), forward primer binding site (57 to 76 bp), reverse primer binding site (407 to 423bp), CamR resistance gene from (1023 to 1883 bp) (Modified from IBA Data Sheet ,2010)

2.2.10. Ligation and transformation of competent *E.coli* with pASK-IBA2C plasmid carrying the gene of interest

The ligation of the expression plasmid with the gene of interest was performed by the T4 DNA ligase enzyme (Promega). This enzyme catalyses the joining of DNA strands between the 5′-phosphate and the 3′-hydroxyl groups of adjacent nucleotides in either the blunt-ended or the cohesive-ended configuration. The pASK-IBA2C plasmid was used as victor to be inserted within the gene of interest as well as the T4 DNA Ligase Buffer (10X), as seen in Table 2.4.

Table 2.4 Ligation mixture

T4 DNA Ligase Buffer (10X)	2 μL
Pask-IBA2C plasmid	2 μL
Insert DNA	5 μL
T4 DNA Ligase	1 μL
Nuclease-free water	Up to 20 μL

The ligation mixture was then incubated at 16°C overnight to be transformed into competent cells (competent *E.coli*).

2.2.11 Transformation of plasmid DNA into E. coli using the heat shock method

In order to replicate the plasmid carrying the gene of interest, the plasmid was transformed into chemically competent E.coli (BIOLINE) (#Bio-85025) using the heat shock method. Then, 5 μ L of ligation mixture was added to 50 μ L of competent E coli and placed on ice before it was incubated at 42°C for 30 min (heat shock) and then placed back in the ice; 1mL of LB broth media (Table 2.5) was then added to the transformed cells before they were incubated at 37°C for 30 min with agitation.

Table 2.5 500 mL LB broth preparation

Trypton	5 g
NaCl	5 g
Yeast Extract	2.5 g
ddH2O	Up to 500 mL & Autoclave (121 C, 20 minutes)

2.2.12 Plating of transformed cells on LB Chloramphenicol agar

As pASK-IBA2C plasmid carries Chloramphenicol-resistant genes, the transformed cells with the plasmid were plated on 25 μ g/mL Chloramphenicol LB agar (Table 2.6) in order to isolate the cells containing recombinant plasmid from those that did not. The plated transformed cells were then incubated at 37°C overnight.

After overnight incubation, the growth colonies were incubated overnight in LB broth at 37°C with agitation.

Table 2.6 LB Chloramphenicol agar preparation

Trypton (#D23821)	5 g
NaCl (#38011-38036)	5 g
Yeast Extract (#12/MFM/1102)	2.5 g
Agar (#9002-18-0)	6 g
ddH2O	Up to 500 mL & Autoclave (121 C, 20 minutes)
chloramphenicol	50 μL from (50mg/ml Chloramphenicol Stock)

2.2.13 Purification of recombinant plasmids from the transformed *E. coli*.

In order to confirm the presence of the gene of interest in the cloning site of pASK-IBA2C plasmid, the recombinant plasmid from the transformed *E.coli* was purified using ISOLATE II PLASMID KIT (Bioline) (#IS493-BO27120) according to the manufacturer's instructions.

The purified recombinant plasmid was then digested by the restriction enzymes HindIII (Promega) and Xbal (Promega) (Table 2.7). Agarose gel was used to analyse the restricted recombinant plasmid.

Table 2.7 Restriction reaction mix

Recombinant plasmid	5 μL
HindIII(Promega)	1 μL
Xbal(Promega)	1 μL
10x NEBuffer 2 (BioLabs)	2 μL
Nuclease-free water	Up to 20 μL

The restriction mixture was then incubated at 37°C for 2h before it was analysed by 1% agarose gel.

2.2.14 Cell growth, gene expression and preparation of bacterial protein extract

2.2.14.1 Host strains

Chemically competent *E. coli* (Bioline) was used as a host strain for periplasmic secretion from recombinant gene expression with the promoter system. The expression of a Strep-Tag fusion protein was induced by the addition of an anhydrotetracycline to logarithmically growing cells. The protein was then exported to the periplasmic space of *E. coli* strain by the ompA leader sequence. The leader sequence was removed when the protein reached the periplasmic space (Plfickthun & Skerra, 1989).

2.2.14.2 Culture, cell harvest and periplasmic extract

To produce the protein, 5 mL of LB/chloramphenicol was incubated with fresh colonies of *E.coli* strains harbouring expression plasmids and was left overnight at 37°C (200 rpm). The preculture was then transferred to 20 mL LB/chloramphenicol in 50-mL Falcon tubes; it was

incubated again with agitation (200 rpm) at 25°C for periplasmic extract. For this extract, the optical density (OD) of the culture was monitored at 550 nm, and when an OD550 reached 0.5 gene expression, it was induced by adding 10µL of an anhydrotetracycline solution (aTe; 2 mg/mL in dimethylformamide, DMF), and shaking continued for 3 hr. The induced *E. coli* samples were then harvested by centrifugation (4200g, 12 min, 4°C).

The supernatant was discarded, and the cell pellets were suspended in 200–1000 µl of cold buffer P (Table 2.8) before they were incubated on ice for 30 min and centrifuged for 5 min at 14k rpm. The supernatant, which consisted of recombinant proteins released from periplasmic space, was carefully pipetted as the periplasmic extract and transferred to an Eppendorf tube. This protein solution was directly readied for Strep-Tactin affinity chromatography, and the cell pellet, which consisted of recombinant proteins left in the cytoplasm, was resuspended in the same volume that was used in the buffer P. in 1× SDS sample buffer to be tested with the periplasmic extract on SDS-PAGE.

Table 2.8 Preparation of buffer P

Tris (pH 8.0)	100 mM
Sucrose	500 mM
Na2EDTA]	1 Mm

2.2.15 Protein verification by the Western blotting technique (SDS-PAGE analysis)

The aim of the western blotting technique is to identify a desired protein by using a specific antibody which can only bind to the antigen of this protein in complex mixture of proteins. These proteins have been fractionated by a polyacrylamide gel and transferred into blotting membrane to be identified based on their molecular weight in the polypeptide chain.

The immunoblotting can be done in six main steps: denaturing of the extracted protein; separating the proteins of the sample using sodium dodecyl sulfatepolyacrylamide (SDS-PAGE) gel; transferring the separated polypeptides to a blotting membrane; blocking nonspecific binding sites on the membrane; incubating the blocking membrane with the specific antibodies; and detecting and visualising the desired protein based on molecular weight.

2.2.15.1 Protein denaturation and sample preparation

Before loading the protein samples for SDS-PAGE gel electrophoresis, $10~\mu L$ from the soluble extract was added to $10~\mu L$ of $4\times SDS$ sample buffer proteins and heated for 3 min at 95°C in order to denature the proteins. This enables the separation of the proteins based on their sizes and prevents proteases enzymes from degrading the samples.

2.2.15.2 Normal SDS-PAGE preparation

Two gels were prepared to analyse the protein extraction samples: lower separation gel and upper stacking gel. The samples were loaded onto different concentrations of lower SDS-PAGE gels according to their size, as they were analysed using 10% gel and then 12%. The four SDS-PAGE concentrations used in this study are shown in Table 2.9.

Table 2.9 Preparation of lower separation gel

Concentration	8%	10%	12%	15%
Number of gels	4	4	4	4
1 M Tris-HCl pH = 8.8	7.5 mL	7.5 mL	7.5 mL	7.5 Ml
40% Acrylamide/Bis 37.5:1	4mL	5 mL	6 mL	7.5 mL
10% *APS	100 μL	100 μL	100 μL	100 μL
20% *SDS	150 μL	150 μL	150 μL	150 μL
ddH2O	8.5 mL	7.5 mL	6.5 mL	5 mL
TEMED	30 μL	30 μL	30 μL	30 μL
Total	20 mL	20 mL	20 mL	20 mL

^{*}SDS: Sodium dodecyl persulfate.

The gel mixture was poured out between Bio-Rad Blotting System glass plates, and the bubbles at the top of the gels were removed by adding drops of isopropanol. The gels were left to set at room temperature for 30 min, and then the isopropanol was washed by dH2O.

Once the lower separation gel was set, the upper stacking gels were prepared, as shown in Table 2.10. Then, the mixture of stacking gels was placed on top of the prepared separation gel layers, and combs were inserted immediately to form the loading. The gels were then left to set at room temperature for 15 min.

^{*}APS: Ammonium Persulfate. *TEMED: Tetramethylethylenediamine

Table 2.10 Preparation of upper stacking gel

Concentration of gel	4%
Number of gels	4
40% Acrylamide/Bis 37.5:1	1 mL
1 M Tris-HCl pH = 6.8	1.5 mL
10% APS	100 μL
20% SDS	50 μL
dH2O	7.5 mL
TEMED	10 μL
Total	20 mL

2.2.15.3 Running samples on SDS-PAGE

The glass plates carrying the gels were placed in Bio-Rad protein electrophoresis tanks filled with running buffer 1x SDS (Table 2.11). The combes were removed, and the first well of the gel was loaded with 5 μ L of the protein standard ladder (Thermo Fisher Scientific), which is a commercially available mixture of proteins of known molecular weights, and the other wells were loaded with the protein samples.

Table 2.11 Preparation of 1x SDS buffer

Glycine(#4037241194)	28.8 g
Tris base(#26841)	6.04 g
SDS(#3634737/88)	2 g
dH2O	up to 2 L

Once the samples were loaded, the electrophoresis gels were run at 100 V for 1.45 hr. When the electrophoresis was finished, the gels were removed from the tanks and transferred onto blotting membranes.

2.2.15.4 Membrane transfer

The aim of this step was to make the proteins accessible to antibody detection by transferring the protein from the gel to the blotting membrane, PVDF, or to the nitrocellulose membrane (#115197535710) (GE Health Life Sciences). This was done using an electric current that can

pull the negatively charged proteins on the gel towards the membrane on the positively charged anode. The PVDF and nitrocellulose membranes were chosen for their non-specific protein binding properties (i.e., it binds all proteins equally well).

The transfer sandwich was set up in the following order: the black cassette was on the bottom, whereas the pre-wet sponge, filter paper (GE Health Life Sciences), equilibrated gel, PVDF or nitrocellulose membrane, filter paper, pre-wet sponge and red cassette were on top. The transfer sandwiches were then placed in the transfer tank filled with 1x transfer buffer (Table 2.12), and the transfer was run at 65 V for 2.5 hr or at 10 V overnight.

Table 2.12 Preparation of 1x transfer buffer.

Glycine	28.8 g
Tris base	6.04 g
Methanol	200 mL
dH2O	2 L

2.2.15.5 Membrane blocking, antibody staining

In order to prevent the interaction between the membrane and the antibodies used to detect the target protein, the membrane was placed in a 3% milk blocking buffer (3% milk powder in 1x PBS, 0.05% Tween 20) for 1 hr on the rocking platform. This enabled the milk buffer to attach to the membrane in all the places where the target proteins had not attached so that the added antibodies could bind to only the target protein.

Once the membrane was blocked, the diluted primary antibody (Table 2.13) was added to the membranes in sealed plastic bags and incubated at -20°C overnight. The primary antibodies were then removed by using the rocking platform and washing them with 1x washing buffer (1x PBS with 0.05% Tween 20) for three times (10 min each) at room temperature.

The washed membranes were then incubated with the diluted secondary antibody in the blocking buffer (Table 2.13) in sealed plastic bags for 1 hr at room temperature using the rocking platform. The membranes were then washed three times (10 min each) by 1x washing buffer to remove any non-binding secondary antibodies.

Table 2.13 Antibodies used in this study

Name of Antibody	Туре	Company	Dilution	Product number
Anti-step-TagII	Primary antibody	Bioscience	1:1000	193988
Anti-Rabbit	Secondary antibodies	Millipore	1:10000	401315

2.2.15.6 Detection and visualization

Once the unbounded secondary antibodies were washed, the bounded proteins were detected using X-ray film. The washed membranes were covered completely with white (0.5 mL) and brown (0.5 mL) Western Lightning Plus-ECL solution (PerkinElmer), which is a non-radioactive light-emitting system designed to detect proteins immobilized on a membrane (perkinelmer website).

The membranes were then placed in plastic wallets. These were then put in a dark developing room, under an X-ray film in the cassette of exposure. After few min of exposure (10–20 min), the film was then developed using the OptiMax X-ray Processor machine.

The protein of interest then was identified based on its size by comparing the obtained bands on the X-ray film with the marker or ladder loaded during electrophoresis.

2.2.16 Purification of Strep-Tag® fusion proteins using Strep-Tactin® Spin Columns

The produced recombinant protein was tagged with a short peptide Strep-Tag® II (8 amino acids, WSHPQFEK), which can be genetically fused upstream or downstream to the reading frame of any gene and can be expressed as fusion peptide and has a high selectivity to Strep-Tactin®.

The tagged protein can be purified by the binding affinity between Strep-Tag II and Strep-Tactin using prepacked chromatography columns (iba #18000069), which allows for a gravity flow purification of the Strep-Tag fusion proteins. Table 2.13 shows the buffers needed for the purification.

Table 2.14 Buffers used in the protein purification

Buffers	concentrations	
Buffer W	100 mM Tris/HCl, pH 8.0	
	150 mM NaCl	
	1 mM EDTA	
Buffer BE (elution buffer)	100 mM Tris/HCl, pH 8.0	
	150 mM NaCl	
	1 mM EDTA	
	2 mM D-biotin	

Before the extracts were loaded to the columns, they were equilibrated three times. They were filled with 500 μ L Buffer W with centrifuging at each step for 30 seconds at 700 x g (approx. 2000 rpm), which rehydrates the dried Strep-Tactin® resin for subsequent use. The lysate was also centrifuged at 13.000 rpm, for 5 min, and at 4°C in order to remove any aggregates that could clog the column. The Tactin® Spin Column.

The preequilibrated Strep-Tactin® spin column was then loaded with up to $500 \,\mu\text{L}$ supernatant of cleared lysate and centrifuged for 30 seconds at $700 \, x$ g (approx. $2000 \, \text{rpm}$), and $2 \, \mu\text{L}$ of collected flow through were analysed with SDS-PAGE. The column was then washed 4 times by $100 \, \mu\text{L}$ Buffer W with centrifuging for 30 seconds at $13.000 \, \text{rpm}$; $2 \, \mu\text{L}$ of the first washing fraction and $20 \, \mu\text{L}$ of each subsequent fraction were taken to be analysed with SDS-PAGE.

The recombinant protein in the washed Strep-Tactin® spin column was eluted by placing the column into a fresh 1.5 mL reaction tube and by adding 150µL Buffer BE (Biotin-Elution-Buffer) three times to the column with centrifuging for 30 seconds at 700 x g (approx. 2000 rpm) in the first time and 15 seconds at maximum speed in the subsequent centrifuging. The purified protein was then analysed with SDS-PAGE.

2.2.17 Protein concentrating using Vivaspin 500 Centrifugal Concentrators

In order to maximise the concentration the protein yield, the elute was placed in Vivaspin® 500 centrifugal concentrators (#1703013), which is an ultrafiltration device that offers a simple, one step procedure for sample concentration with starting volumes of up to $500 \, \mu L$.

2.2.18 Using a Q5 $^{\circ}$ Site-Directed Mutagenesis Kit for the mutagenesis of the gene of interest

The gene of interest was mutated to create substitutions using a Q5® Site-Directed Mutagenesis Kit (#0071605). This was selected because, unlike other Mutagenesis Kits used to mutate genomic DNA (e.g. CRISPR-CAS9), it is capable of mutating plasmid DNA. Mutagenesis was carried out in three steps: (1) exponential PCR amplification using mutagenic primers (Table 2.15, 2.16) and a Q5 Hot Start High-Fidelity DNA Polymerase; (2) the circularisation of amplified material; and (3) the transformation at room temperature into competent *E. coli* (BioLabs website).

Table 2.15 Mutagenic primers sets used in the exponential PCR amplification

Primer sets	Sequences (5'-3')
CTXM-14-P273A C277R F	CCGCAACAGAACGCAGAGAGCCGCAGAGATGTGCTGGC
CTXM-14-P273A C277R R	CTGGGTAAAATAGGTCACCAG
CTXM-14-A55TR	ACGCCCAGCCGCCTCCGTGC
CTXM-14-A55TR	AGAGATGTGCTGGCCTGGGT

Table 2.16 Mutagenesis PCR reaction mix

Q5 Hot Start High-Fidelity 2X Master Mix	12.5 μL
10 μM Forward Primer	1.25 μL
10 μM Reverse Primer	1.25 μL
Template DNA	1 μL
Nuclease-free water	Up to 25 μL

The PCR reaction mixture was run using the PCR program, as shown in Table 2.16.

Table 2.17 PCR program of the PCR Amplification

STEP	TEMP	TIME
Initial Denaturation	98°C	30 seconds
25 Cycles	98°C	10 seconds
	50–72°C	10–30 seconds
	72°C	20–30 seconds
Final Extension	72°C	2 minutes
Hold	4–10°C	

The amplified material was circularized by adding Kinase-Ligase-DpnI (KLD) enzyme mix, as shown in Table 2.18.

Table 2.18 KLD Reaction

PCR Product	1 μL
2X KLD Reaction	5 μL
Buffer	
10X KLD Enzyme	1 μL
Mix	
Nuclease-free Water	Up to 10 μL

After 5 μL of KLD mix was added to 50 μL of competent *E coli* and placed on ice, it was incubated at 42°C for 30 min (heat shock) and then placed back in the ice; 1mL of LB broth media (Table 2.6) was then added to the transformed cells before they were incubated at 37°C for 30 min with agitation. The transformed cells with the plasmid were plated on 25 μg/mL Chloramphenicol LB agar (Table 2.7). The plated transformed cells were then incubated at 37°C overnight. After overnight incubation, the growth colonies were incubated in LB broth at 37°C overnight with agitation.

The recombinant plasmid carrying the gene with mutation was purified from the transformed *E. coli* using the ISOLATE II PLASMID KIT and prepared for sequencing. After sequencing,

the mutated gene was induced, purified and concentrated (see sections 2.2.15.5, 2.2.15.6 and 2.2.15.7).

2.2.19 Biochemical characterization

Biochemical characterization involved studying the phosphorylation status of B-lactamase proteins, which was completed by two methods:

2.2.19.1 Isoelectric-focused analysis of the CTX-M proteins

The isoelectric (2 D) gel electrophoresis technique is a molecular technique in which proteins are distributed across the 2 D gel profile; they are first separated by their isoelectric point (pI) in isoelectric focusing and then are further separated by molecular weight using sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE). This technique expands the number of proteins that can be identified, and it efficiently provides detailed data for proteomics analysis (Florian Weiland et al., 2014).

In the 2D analysis, 15 μ L of protein extract was further diluted in 110 μ L of IEF buffer, and the total 125 μ L was loaded onto the 2D tray. The immobilized pH gradient (IPG) strips (#163-2000) nonlinear (pH 3–10, Bio-Rad) were then put over the protein solution, and the strips were then overlaid with 2 mL of mineral oil, and the 2D program was set, as shown in Table 2.19.

Table 2.19 Programme of 2D analysis

Running	Active dehydration at 50 V for 12 hr
Focusing	Rapid method, 10,000 V, hold to 500 V

After the run was completed, the 2D strips were washed in 2D tray using the rocking platform with 2D solution: solution I (6 M urea, 0.375 M Tris-HCl [pH = 8.8], 2% SDS, 20% glycerol, 2% [w/v] Dithiothreitol [DTT]) for 10 min; and solution II (Tris-HCl pH = 8.8, 2% SDS, 20% glycerol, 2.5% [w/v] iodoacetamide) for another 10 min.

The 2D strips were run on a 10% normal SDS-PAGE gel with only the lower separation layer (Table 2.9). The separation gels were cast right to the top of the glass plates, leaving approximately 5 mm space to apply the strip. The electrophoresis gels were run at 100 V for

1.5 hr and the transfer, blocking and visualizing were performed, as mentioned in sections 2.2.15.5 and 2.2.15.6.

2.2.19.2 Phos-tag gels

In order to analyse the phosphorylation status of the proteins, the manganese (II) Phos-tag SDS-PAGE was used, which specifically delays the mobility of the phosphorylated proteins at a neutral pH. The lower Phos-tag SDS was prepared in the way shown in Table 2.20.

Table 2.20 Preparation of phos-tag gel

Acrylamide/Bis 29:1	1.4 mL
1 M Tris-HCl pH = 8	1.8 mL
5 mM Phos-tag	70 μL
10 mM MnCl2	70 μL
20% SDS	27 μL
TEMED	15 μL
10% APS	70 μL

The gel mixture was poured out between the Bio-Rad Blotting System glass plates, and the bubbles at the top of the gels were removed by adding drops of isopropanol. The gels were left to set at room temperature for 30 min, and then the isopropanol was washed by dH2O.

Once the lower separation gel was set, the upper stacking gels were prepared, as shown in (Table 2.10) then, the mixture of stacking gels was placed on top of the prepared separation gel layers, and combs were inserted immediately to form the loading. The gels were left to set at room temperature for 15 min.

The gels were then run with electrophoresis as described in section 2.2.15.3. Once the run was finished, the gels were washed three times in 1x transfer buffer with 20 mM EDTA to remove the Mn++ ions, as they interfere with the transfer. The transfer, blocking and visualizing were performed as mentioned in sections 2.2.15.5 and 2.2.15.6.

2.2.20 Determining the MIC of induced strep-tagged recombinant genes using the E-Test

The determination of minimum inhibitory concentrations (MICs) of induced strep-tagged recombinant genes using the E-Test method reveals how active the β -lactamases are against a range of different antibiotics. The E-Test method is a well-established assay for testing antimicrobial resistance by applying a predefined gradient of antibiotic concentrations on a plastic strip onto a layer of bacterial cells (Biomerieux website).

Materials needed for E-Test are, Agar plate with selected media for susceptibility test, inculcation suspension media, swabs used for inclusion of suspended strains on agar plate, 0.5 McFarland turbidity standards, and antibiotic strips.

To determine the MIC of the recombinant genes, the fresh colonies of the *E.coli* strains that harboured expression plasmids with the gene of interest were induced before they were plated with E-Test strips of selected antibiotic (Table 2.21). Plates were incubated at 35 °C. MIC detection was repeated in biological triplicates for each antibiotic.

Table 2.21 E-test strip of selected antibiotics

Name of Antibiotic strip	Company	Product number
Nitrofurantoin	BIOMÉRIEUX	10047676160
Ceftazidime	BIOMÉRIEUX	1004719296
Cefoxitin	BIOMÉRIEUX	1004857420
Cefotaxime	BIOMÉRIEUX	100495820
Imipenem	BIOMÉRIEUX	100412373

2.2.21 Determination of kinetic constants

The kinetic constants of the protein of interest was measured by mixing a limited amount of purified β -lactamase proteins with different concentrations of selected lactam antibiotics at 25°C in the assay buffer (50 mM phosphate buffer, pH 7.0) in a 1-cm quartz cuvette using a spectrometer for 12 min.

The absorbance was measured in triplicate with each concentration and then a nonlinear regression of the Michaelis-Menten equation was applied to the data in order to determine the Km (He, DD et al., 2015).

The protein concentration in the tested samples at absorbance (at λ 280nm) was determined using Protein Concentration Calculator web (www.aatbio.com/tools/calculate-protein-concentration). The determination of the protein in solution was based on substituting of the molecular weight of the protein, extinction coefficient, and λ max (280nm), which was measured by spectrophotometer, into a derived form of the Beer-Lambert law.

Chapter 3: URINARY TRACT INFECTIONS DATA ANALYSIS

3.1 Introduction

Urinary tract infections (UTIs) are considered one of the most infectious diseases occurring both in the healthcare setting and in the general community (Rowe & Juthani-Mehta, 2013). UTIs can often be diagnosed quickly and treated effectively with antibiotics (Schmiemann et al., 2010). However, some UTI patients cannot be treated with commonly-used antibiotics, a situation thought to be caused by resistant species, of which producers of Extended-spectrum beta-lactamases (ESBLs) are the most common and detectable pathogens in UTI patients (Al Yousef et al., 2016).

ESBLs are enzymes produced mainly by gram-negative bacteria (*E. coli*, coliforms, *Acinetobacter spp.* and others), and they can exhibit a resistance to most beta-lactam antibiotics, including third- and fourth-generation cephalosporins, which severely limits the applicable treatments for infections caused by ESBLs (Blaak H et al., 2013). However, these ESBL-producing strains may be inhibited by beta-lactamase inhibitors such as clavulanic acid, tazobactam and sulbactam (Engler et al., 2017). The spread of ESBL-producing bacteria has become a major global health concern over the past generation (Kuralayanapalya et al., 2019).

Despite the high number of patients referred for urinalysis due to infection of the urinary tract in North Wales health care settings, as it is shown in table 3.1, between 2011 and 2013 from the total number of 193142 cases 68% were referred from their local GP to the hospital, few epidemiological studies have looked for a pattern of antibiotic resistance resulting from ESBL producers.

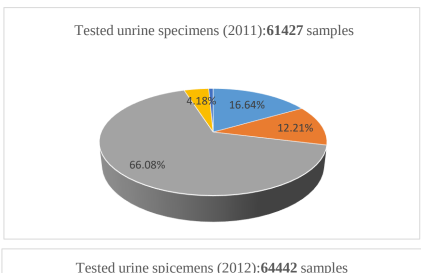
This part of the study attempts to respond to this research need by analysing the data of UTI patients in a North Wales tertiary care hospital, Ysbyty Gwynedd Hospital, to identify the risk factors associated with isolations of *E. coli* carrying ESBL in urine samples from hospitalised patients and outpatients.

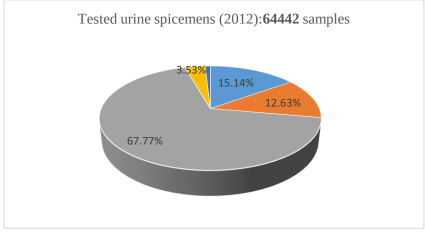
Table 3.1 Urine specimens tested from 2011 to 2013 in Ysbyty Gwynedd Hospital

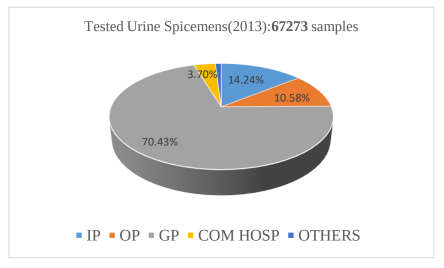
Source/Years	2011	2012	2013
IP	10227	9761	9584
OP	7503	8145	7120
GP	40658	43677	47386
COM HOSP	2570	2278	2493
OTHERS	469	581	691
Total	61427	64442	67273

Data taken from Ysbyty Gwynedd Hospital for patients referred for urine tests over three years (2011-2013); the groups are IP (Inpatients), OP (Outpatients), GP (General practice), com hospital (Community hospital: Dolgellau, Alltwen Hospital in Porthmadog and Stanley Hospital in Holyhead), and other sources.

Figure 3.1 The number of urine specimens submitted to Ysbyty Gwynedd Hospital over the three years (2011-2013)







Number of patients referred for urinalysis due to symptoms of the urinary tract infection to one of the three main referral hospital in in North Wales. It is noticeable that the number of tested urine samples has increased from 61427 in 2011 to 67273 in 2013. It is noticeable also that GP samples represents more than two thirds (68%) from the total number of tested samples over the three years which indicates to increasing in community acquired infection.

Research ethics

The patient specimens were routine diagnostic requests sent to the Microbiology laboratory for testing from primary and secondary care settings. Once in the laboratory the data collected included source, age and gender of patients, phenotypic antibiotic screening and causative organisms. Each data set was anonymised and no patient identifiable information was recorded.

3.2 Results

3.2.1 Data collection

Isolates from Ysbyty Gwynedd Hospital were screened and tested phenotypically and genetically to detect the cause of CTXM-type ESBL-producing bacteria in the antibiotic-resistance pattern among UTI patients in North Wales. Despite there being 300 such isolates at the three main referral hospitals in North Wales, access to only 100 samples and medical data was available from Ysbyty Gwynedd Hospital. These 100 isolates were statistically analysed in terms of frequency of UTI among ages and genders, detected ESBLs and the distribution of phenotypic resistance against various antibiotics. Table 3.2 shows the medical data of 100 samples (No 201–300) collected from Ysbyty Gwynedd Hospital in 2014.

Table 3.2 Characterisation of isolates from Ysbyty Gwynedd Hospital (201-300)

									Pher	noty	pic a	ntib	iotic						
Lab#	Isolate #	Age	Sex	AMO	CPD	AUG	LΙΝ	TRI	CTX	CAZ	GEN	AMI	₹	MER	PTZ	CIP	CPD/CV	ID	GP/Hosp
27697	201	77	F	R	R	S	S	R	R	R	S	S	S	S	S	S	+	Colif	GP
G28282	202	55	М	R	R	S	S	R	R	S	S	S	S	S	S	R	+	Colif	GP
27855	203	56	M	S	S	S	S	R	S	S	S	S	S	S	S	S	+	E. coli	GP
28469	204	88	M	R	R	S	R	R	R	R	S	S	S	S	S	S	+	E. coli	Hosp
25696	205	72	M	R	R	S	S	S	R	R	R	S	S	S	S	S	+	E. coli	Hosp
23484	206	71	F	R	R	R	R	R	R	R	S	S	S	S	S	R	+	E. coli	GP
23623	207	69	F	R	R	S	S	S	R	R	S	S	S	S	S	S	+	E. coli	GP
23414	208	100	F	R	R	R	S	R	R	R	S	S	S	S	S	R	+	E. coli	Hosp
23842	209	77	M	R	R	R	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
12030	210	75	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
12294	211	92	F	R	R	R	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
12675	212	93	F	R	R	S	R	R	R	R	S	S	S	S	S	R	+	E. coli	GP
12640	213	90	F	R	R	S	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
13095	214	68	М	R	R	R	S	R	R	R	R	S	S	S	R	R	+	E. coli	GP
15309	215	57	F	R	R	S	S	R	R	R	S	S	S	S	S	R	+	E. coli	GP
15227	216	59	М	R	R	R	R	R	R	R	S	S	S	S	S	R	+	E. coli	GP
20057	217	83	F	R	R	R	S	R	R	S	R	S	S	S	R	R	+	E. coli	GP
27700	218	93	М	R	R	S	R	R	R	R	R	S	S	S	S	S	+	Prote	GP
28287	219	90	М	R	R	S	S	R	R	R	R	S	S	S	S	R	+	Colif	GP
11773	220	80	М	R	R	S	R	R	R	R	R	S	S	S	S	S	+	Colif	Hosp
28316	221	92	F	R	R	S	R	R	R	R	S	S	S	S	S	S	+	Colif	GP
23635	222	68	М	R	R	R	S	R	R	R	S	S	S	S	S	R	+	E. coli	GP
23456	223	92	F	R	R	S	S	R	R	R	S	S	S	S	S	R	+	E. coli	GP
23670	224	85	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
23362	225	81	М	R	R	S	R	R	R	R	S	S	S	S	S	R	+	E. coli	Hosp
12472	226	81	F	R	R	S	S	R	R	R	S	S	S	S	S	R	+	E. coli	Hosp
12564	227	62	F	R	R	S	S	R	R	R	R	S	S	S	S	R	+	Colif	Hosp
12672	228	52	М	R	R	S	R	R	R	R	S	S	S	S	S	S	+	Colif	GP

					Phenotypic antibiotic														
Lab #	Isolate#	Age	Sex	АМО	CPD	AUG	ΗN	TRI	CTX	CAZ	GEN	AMI	₫	MER	PTZ	CIP	CPD/CV	ID	GP/Hosp
12447	229	70	М	R	R	R	R	R	R	R	S	S	S	S	S	S	+	Colif	GP
13428	230	62	F	R	R	R	S	R	R	R	R	S	S	S	S	R	+	Colif	GP
15581	231	97	F	R	R	R	R	R	R	R	R	S	S	S	S	R	+	E. coli	Hosp
15217	232	40	F	R	R	R	S	R	R	R	R	S	S	S	R	R	+	E. coli	GP
18069	233	84	М	R	R	S	R	R	R	R	R	S	S	S	S	R	+	Colif	Hosp
27717	234	96	F	R	R	S	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
15306	235	79	F	R	R	S	S	S	R	R	S	S	S	S	S	R	+	E. coli	GP
15468	236	73	F	R	R	R	R	R	R	R	R	S	S	S	S	R	+	E. coli	Hosp
13373	237	82	М	R	R	S	S	R	R	S	S	S	S	S	S	R	+	E. coli	GP
13388	238	86	F	R	R	R	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
13232	239	74	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
13291	240	88	F	R	R	R	R	R	R	R	S	S	S	S	S	R	+	Colif	GP
13369	241	77	F	R	R	R	R	R	R	R	S	S	S	S	S	S	+	Colif	Hosp
12405	242	66	F	R	R	S	S	R	R	R	S	S	S	S	S	S	+	E. coli	GP
23471	243	93	F	R	R	R	S	R	R	R	S	S	S	S	S	S	+	E. coli	Hosp
23604	244	25	F	R	R	R	R	R	R	R	R	S	S	S	S	R	+	Colif	Hosp
23631	245	74	М	R	R	R	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
28223	246	87	F	R	R	S	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
28096	247	96	F	R	R	R	R	R	R	R	R	S	S	S	S	R	+	E. coli	GP
28194	248	25	F	R	R	R	R	R	R	R	R	S	S	S	S	R	+	Colif	GP
27651*	249	91	F	R	R	S	R	R	R	R	R	S	S	S	S	R	+	Colif	GP
27551*	250	92	F	R	R	S	R	R	R	R	R	S	S	S	S	R	+	Protu	GP
27717	251	96	F	R	R	S	R	R	R	R	S	S	S	S	R	R	+	E. coli	GP
23946	252	74	М	R	S	S	R	R	R	S	S	S	S	S	S	R	+	Steno	Hosp
27301	253	94	F	R	R	R	S	S	R	S	R	S	S	S	R	S	+	E. coli	GP
27297	254	86	F	R	R	S	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
27181*	255	73	М	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
27181*	256	73	М	R	R	R	R	R	R	R	R	S	S	S	R	R	+	Prote	Hosp
26083	257	91	F	R	R	S	S	R	R	R	R	S	S	S	R	R	+	Colif	GP
26988	258	71	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
26548	259	86	F	R	R	S	S	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp

					Phenotypic antibiotic														
Lab #	Isolate #	Age	Sex	АМО	CPD	AUG	FIN	TRI	CTX	CAZ	GEN	AMI	Σ	MER	PTZ	CIP	CPD/CV	ID	GP/Hosp
25942	260	64	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
25673	261	73	М	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
23781	262	91	F	R	R	S	R	R	R	R	R	S	S	S	R	R	+	Colif	GP
24954	263	75	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
24543	264	88	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
24623	265	73	F	R	R	R	S	R	R	R	R	S	S	S	R	R	+	E. coli	GP
23096	266	80	F	R	R	R	S	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
25706	267	84	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
23759	268	89	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
24026	269	90	F	R	R	S	S	R	R	R	R	S	S	S	R	R	+	E. coli	GP
24526	270	73	М	R	R	S	S	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
23760	271	22	F	R	R	R	R	S	R	R	R	S	S	S	R	R	+	E. coli	GP
23787	272	3	F	R	R	S	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
23789	273	94	F	R	R	S	S	R	R	R	R	S	S	S	R	R	+	E. coli	GP
23975	274	20	F	R	R	R	S	R	R	R	R	S	S	S	S	S	+	E. coli	GP
24542	275	86	М	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
24654	276	86	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
25006	277	88	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	Colif	GP
25065	278	95	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
25864	279	89	М	R	R	R	R	R	R	R	R	S	S	S	S	S	+	Colif	GP
24785	280	80	М	R	R	S	S	R	R	R	R	S	S	S	R	R	+	E. coli	GP
25792	281	90	М	R	R	S	S	R	R	R	R	S	S	S	R	R	+	Colif	Hosp
26477	282	80	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	Colif	GP
25691	283	90	М	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
25898	284	49	F	R	R	S	S	R	R	R	S	S	S	S	R	R	+	E. coli	GP
26523	285	100	F	R	R	R	R	R	R	R	S	S	S	S	R	R	+	E. coli	GP
26592	286	80	F	R	R	S	S	R	R	R	S	S	S	S	R	R	+	E. coli	Hosp
27083	287	80	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
26602	288	20	F	R	R	R	S	R	R	R	S	S	S	S	S	S	+	E. coli	GP
26852	289	86	F	R	R	S	S	R	R	S	S	S	S	S	R	R	+	E. coli	Hosp
26822	290	93	F	R	R	R	S	R	R	R	S	S	S	S	R	R	+	E. coli	Hosp

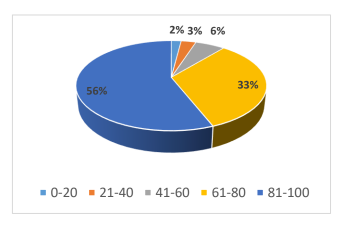
					Phenotypic antibiotic														
Lab #	Isolate #	Age	Sex	AMO	CPD	AUG	ΙN	TRI	CTX	CAZ	GEN	AMI	₫	MER	PTZ	CIP	CPD/CV	ID	GP/Hosp
26675	291	50	F	R	R	S	S	R	R	R	S	S	S	S	R	R	+	E. coli	GP
27449	292	79	М	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
27554	293	90	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
27442	294	79	М	R	R	S	S	R	R	S	S	S	S	S	R	R	+	E. coli	GP
27611	295	76	М	R	R	R	R	R	R	R	S	S	S	S	R	R	+	E. coli	GP
27184	296	59	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	Hosp
27303	297	93	F	R	R	S	S	R	R	S	S	S	S	S	R	S	+	E. coli	Hosp
27231	298	69	М	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
27383	299	80	F	R	R	R	R	R	R	R	R	S	S	S	R	R	+	E. coli	GP
27112	300	79	F	R	R	R	R	R	R	R	S	S	S	S	R	R	+	E. coli	Hosp

F: female, M: male, R: resistant, S, Sensitive, Amo: Amoxicillin, CPD: Cefpodoxime, CIP: ciprofloxacin, Aug: Augmentin, Ni: Nitrofurantoin, CTX: cefotaxime, CAZ: Ceftazidime, Gen: Gentamicin, Ami: amikacin, IMI: Imidazole, Mer: Meropenem, Ptz: piperacillin/tazobactam, GP: general practice

3.2.2 Frequency of UTI according to patient age and gender

The data collected in the present study (Table 3.2) show that the prevalence of UTI may be affected by age. People aged 0–20 represents the lowest frequency of UTI problems (3%) while those aged 61–100 are most susceptible to UTIs (89%), followed by those aged 41–60 (8%) and 21–40 (4%).

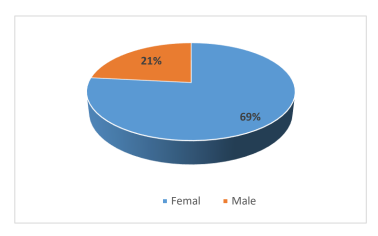
Figure 3.2 UTI prevalence based on patient age



The cut-off value used to group the prevalence (i.e. percentage) of each age group from 100 UTI patients (>20, 21–40, 41–60, and 81–100) based on the patient ages provided in Table 3.2 (Medical data collected from Ysbyty Gwynedd Hospital in 2014.

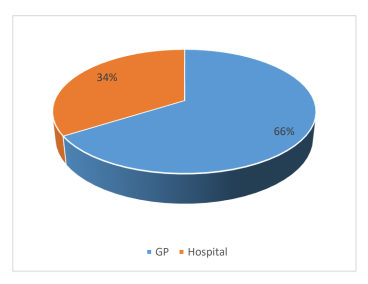
The data also shows that 69% of diagnosed UTIs occur in females, which is thought to be due to anatomical differences between the sexes (Figure 3.3). Moreover, most of the infections were community-acquired, with 66% of the infection cases detected by general practitioners (GPs; Figure 3.4).

Figure 3.3 UTI prevalence based on patient gender



Susceptibility for urinary tract infections according to gender. % of females to males from 100 UTI patients (Medical data collected from Ysbyty Gwynedd Hospital in 2014.

Figure 3.4 Hospital and community acquired UTI



Percentage of community acquired to hospital acquired urinary tract infections in 100 UTI patients (Medical data collected from Ysbyty Gwynedd Hospital in 2014). * GP (General practice).

3.2.3 ESBL-producing isolates as a causative organism for UTI

The screening of extended-spectrum beta-lactamase was carried out in the microbiology department of Ysbety Gwynedd by the Combination Disc (diffusion method) in accordance with CLSI (Clinical and Laboratory Standards institution) guidelines which propose diffusion methods for 21 antibiotics susceptibility testing used in secreting of ESBL-producing isolates (CLSI,2016).

In the combination Disc method two discs containing antibiotics are positioned on the agar plates: one Cefpodoxime disc (CPD) and one Cefpodoxime with clavulanic acid (CPD/CV) disc. Following incubation with the same bacterial isolate, the diameter of the inhibition zone around the two discs is measured, and the test is considered positive (ESBL-producer) if the difference in diameter between the two discs is ≥ 5 mm (Carter et al., 2000).

Medical data from Ysbyty Gwynedd Hospital on UTI patients shows that 99% of the detected isolates among UTI patients display a resistance to CPD which are susceptible to the combination of Cefpodoxime and clavulanic acid. The resistant to Cefpodoxime indicates the expression of an Extended-spectrum beta-lactamases (ESBL) (Figure 3.5). Only one isolate in the study exhibited sensitivity to CPD disc (isolate #203).

The high prevalence of ESBLs in the study samples is emphasised by the ESBL data from Ysbyty Gwynedd Hospital (Tables 3.3, 3.4 and 3.5), which show that the total number of detected CPD-resistant isolates among UTI patients from different areas in North Wales has increased over three years (2011–2013) from 831 to 1345, while the detection of ESBL-producing isolates-which represent two thirds of detected CPD-resistant isolates (Figure 3.6)-has grown accordingly from 569 to 963 over the same period.

Table 3.3 Total Number of detected *E.coli/coliforms* amongst UTI patients from 2011-2013 in YG hospital.

Source/Years	2011	2012	2013
IP	1424	1570	1964
OP	872	1195	1252
GP	7793	9970	11654
COM HOSP	414	462	624
OTHERS	52	50	56
Total	10555	13247	15549

Table 3.4 Detected CPD-resistant isolates in *E.coli/coliforms* between 2011 and 2013 at Ysbyty Gwynedd Hospital

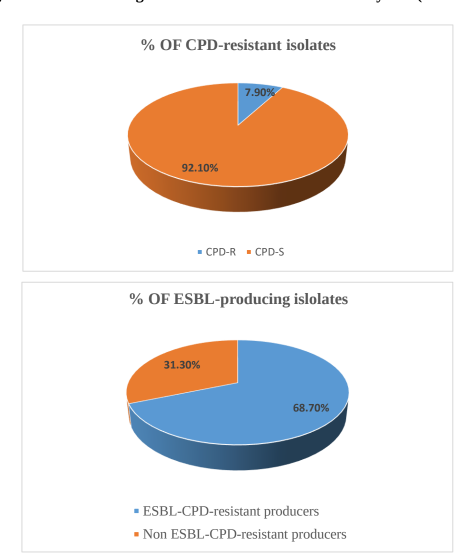
Source/Years	2011	2012	2013
IP	155	163	274
OP	67	59	94
GP	552	654	864
COM HOSP	51	57	109
OTHERS	6	5	4
Total	831	938	1345

Table 3.5 Number of detected extended-spectrum beta-lactamase (ESBL)—positive isolates from CPD-resistant isolates between 2011 and 2013 at Ysbyty Gwynedd Hospital

Source/Years	2011	2012	2013
IP	116	113	211
OP	40	35	65
GP	371	410	590
COM HOSP	38	46	94
OTHERS	4	4	3
Total	569	608	963

Data taken from Ysbyty Gwynedd Hospital on patients referred for urine tests over three years (2011–2013): (**Table 3.3.**) shows the total number of *E. coli* and coliform, which increased from 10,555 to 15,549 over the three years; (**Table.3.4**) shows the total number of CPD-resistant isolates (Cefpodoxime is one of the antimicrobial agents recommended by the National Committee for Clinical Laboratory Standards for screening isolates of *Klebsiella spp.* and *Escherichia coli* for extended-spectrum β-lactamase (ESBL) production), which rose from 831 to 1,345 over the same three years); (**Table.3.5**) shows the number of detected ESBL-producing isolates, which approximately doubled between 2011 and 2013.* IP (Inpatients) *OP (Outpatients),* GP (General practice),* com hospital (Community hospital: Dolgellau , alltwen hospital in porthmadog and stanley hospital in Holyhead)

Figure 3.5 ESBL among CPD-resistant isolates over three years (2011-2013)



The percentage of CPD-resistant isolates among 39351 *Ecoli/Coliforms* detected in urine samples tested over three years 2011-2013:**7.9%** (3114). Among 3114 CPD-resistant *Ecoli/Coliforms* more than two thirds (68.70%) of isolates show ESBL production-based resistance.

3.2.4 Antibiotic resistance profile

Antibiotic profile includes agreed first line antibiotic usage for treatment of UTIs in primary and secondary care with diagnostic protocol following HPA (Health Protection Agency) and UCAST guideline.

It can be observed from Table 3.2 that the isolates displayed the highest resistance to amoxicillin (99%), Cefpodoxime (98%), cefotaxime (96%), Ceftazidime (91%) and trimethoprim (91%), while the isolates were almost completely susceptible to amikacin, imidazole and meropenem (Figure 3.6).

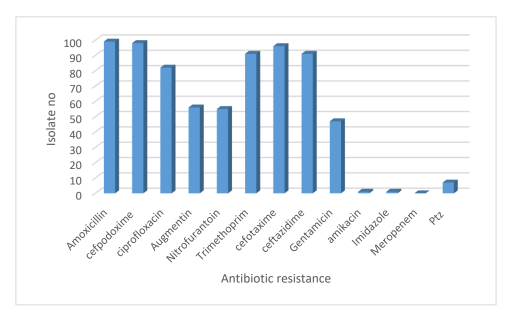


Figure 3.6 Phenotypic resistance to various antibiotics.

Charts shows the resistance of ESBL producing isolates to agreed first line antibiotic usage for treatment of UTIs in primary and secondary care with diagnostic protocol following HPA (Health Protection Agency) and UCAST guideline.

3.2. Discussion

According to the data from Ysbyty Gwynedd Hospital (Table 3.1), 67,273 patients were referred to the hospital between January and December 2013 for urinalysis after reporting symptoms of the urinary tract infections. This is a high incidence rate, representing approximately 34.5% of population of the Gwynedd and Anglesey study area (population of 194,780; Welsh Government, statistics and research department, 2019).

One possible explanation is that North Wales is a rural area, with approximately 80% land managed for farming (National Assembly for Wales, 2017). The consumption of antibiotics by livestock, which is considered a major risk factor for the emergence of antibiotic resistance, has increased significantly in recent years according to the review on antimicrobial in agriculture and the environment published in 2015 (Review on Antimicrobial Resistance, 2015). However, it should be remembered that AB usage is much reduced on the beef and sheep farms typical of North Wales (UK Veterinary Antibiotic Resistance and Sales Surveillance Report, 2019), making this less likely to be the main explanatory factor for our results.

The present study confirms previous reports of a significantly higher prevalence of UTIs among females. Based on data by Ysbyty Gwynedd Hospital, women represent around two thirds of infected patients. Proposed reasons for this cite anatomic differences including shorter urethral length and the moist periurethral environment in women (Hickling et al., 2015). However, unlike other studies including that by Sobel (2014) which conclude that young, sexually-active women 18–24 years of age have the highest incidence of UTIs, the present study shows that the majority of women with UTIs in North Wales are over 40; only 7.2% are aged under 25.

As can be seen in Table 3.2, a large majority of patients (88%) were over 60 years of age, which supports the conclusion of many studies that increasing age is itself a risk factor for UTIs. There are many possible reasons for the susceptibility of elderly people to UTIs, including: urinary incontinence and urinary retention, hospitalisations and accompanying urinary catheterisations, long-term medical institutionalisation, immunity senescence as well as other factors such as anatomic abnormalities of the urinary tract, particularly those which produce incontinence or urinary retention (e.g., prostatic hyperplasia) (Wang et al.,2017). The high UTI incidence in North Wales can be explained by the high percentage of old people in the population of North Wales Which, according to Welsh Government services and

information, is expected to rise from 18% (as determined in 2008) to almost 26 % of the Welsh population by 2033.

Regarding community and nosocomial (hospital) acquisition of UTIs, the data support previous research finding that community-acquired (uncomplicated) UTI infections are of a higher prevalence than those acquired in hospitals. According to many reviews (e.g. Kucheria et al. 2005), *E. coli* causes less than 50% of nosocomial UTIs but causes more than 80% of community-acquired infections. However, the present study shows that 27 out of 34 (79%) hospital-acquired UTIs are caused by *E. coli*. This may indicate an increase in the spread of *E. coli* in hospitals as an uropathogen.

Analysis of ESBL data in this study can be added to other studies that show an alarming increase in the threat posed by antibiotic resistance conferred by ESBLs due to the two-fold increase in the number of ESBLs detected over the three years of this study, 2011-2013 (Table 3.5). The antibiotic resistance profiles of the samples analysed in this study showed that 68.70% of the CPD (Cefpodoxime) resistant isolates tested over the three years (2011-2013) belong to ESBL positive class. The resistance in the remaining isolates may be caused by other β -lactamase enzymes that show hydrolytic activity against cephalosporins, including CPD. AmpC β -lactamase could be responsible for CPD resistance in these isolates.

Lewis et al. (2015) studied the prevalence of AmpC promoter mutations in groups of ESBL-negative uropathogenic *E.coli* strains that confer CPD resistance. They detected AmpC-carrying plasmids in only 10 isolates from 50 clinical samples, while the analysis of the chromosomal AmpC promoter regions in the remaining strains revealed mutations at 16 different positions and they found that AmpC mediated resistance in uropathogenic *E.coli* to Cefpodoxime can result from three point mutations in chromosome.

One of the possible reasons behind the rapid growth of ESBL-based resistance is the inaccurate phenotypic methods used for ESBL screening in clinical laboratories (e.g. the double-disc synergy test; Bajpai et al., 2017). These methods are unable to distinguish between the approximately 200 types of ESBLs identified to date (Ghafourian et al., 2015) and this is likely to lead to treatment failures due to the use of inappropriate antibiotic and outbreaks of ESBL-based antibiotic resistance.

To highlight the inadequacies of the routine screening methods for ESBLs and to detect the responsibility of the most rapidly growing ESBL types, the CTXM-type-ESBL gene, which

confers resistance in the isolates of the study to most antibiotics, were screened further genetically, using PCR followed by sequencing (Chapter 4).

Chapter 4: DETECTION OF GENES ENCODING CTX-M-TYPE-ESBL IN ISOLATES FROM NORTH WALES UTI PATIENTS

4.1 Introduction

The detection of resistant isolates in diagnostic laboratories is primarily based on phenotypic testing, which is characterised by cost-effectiveness and ease of use (Ayefoumi et al., 2019). However, the limitations of the phenotypic methods result in their inability to determine the gene responsible for antibiotic resistance, which often results in treatment failures due to the use of inappropriate antibiotics, in addition to the outbreak of multi-resistant, gram-negative pathogens (Srivastava et al., 2018). Such outbreaks could be prevented by the correct molecular analysis, which enables the identification of the genetic determinants of pathogenicity and antibiotic resistance (Hiltunen et al., 2017).

Genes that code for multiple types of β -lactamases (*bla* genes), predominately located on plasmids, are considered the most common antibiotic resistance genes (Chen et al., 2014). In a significant part, it refers to extended-spectrum β -lactamases (ESBLs), which inactivate most beta-lactam antibiotics, such as penicillins, cephalosporins (except for cephamycins) and monobactams, in addition to third- and fourth-generation cephalosporins, by hydrolysis of the β -lactam ring (Bachiri et al., 2017). This severely limits the possibilities of treatment for infections caused by ESBLs. In urinary tract infections (UTIs), *E. coli* remains the most common producer of ESBLs (Damus-Białek et al., 2018).

Many classifications of ESBLs have been introduced since the first emergence of this group of broad substrate enzymes. The most common classification is based on the type of beta-lactam that is hydrolysed by this ESBL type more than the other type (Rahman et al., 2018). The most important types are SHV-type-ESBL, which efficiently hydrolyses penicillin and first-generation cephalosporins, TEM-type-ESBL, which hydrolyses ampicillin at a greater rate than carbenicillin, oxacillin or cephalothin, OXA-type-ESBLs which are so named because of their oxacillin-hydrolysing abilities, and CTX-M-type-ESBLs' name reflects their potent hydrolytic activity against cefotaxime in addition to rare ESBL types e.g. VEB-1 and BES-1 (Taylor & Francis, 2015).

CTX-M has become the predominant type of ESBL enzymes detected in many regions of the world (Chaudhary et al., 2016). It has been reported that among ESBLs producing *E. coli* detected in clinical laboratories, the most commonly identified are enzymes from the family

CTX-M (mostly CTX-M-1 lineage), followed by SHVs and recently the less frequent TEMs (Maria et al., 2018).

The *blaCTX-M* gene is predominantly encoded on large plasmids that tend to carry additional resistance genes, which explains the increasing detection of resistant *E. coli* in outpatients and patients admitted to hospitals for short periods. This suggests that this plasmids spread in bacterial communities outside of hospitals for example in farming environments (Riccobono et al., 2015).

The *blaCTX-M* gene encodes 291 amino acids, and a single change of an amino acid constitutes a new CTX-M type (Giedraitienė et al., 2017). Unlike other ESBLs, this enzyme does not enlarge the active site to accommodate the antibiotics, instead its hydrolytic activity depends on point substitutions (Taylor & Francis, 2015).

Since the emergence of the CTX-M enzyme there have been at least 128 CTX-M types that have been described (Po et al., 2017). The phylogenetic tree of CTX-M-type ESBLs has been constructed by using amino acid sequences in which the CTX-M enzyme is divided into 5 groups: CTX-M-1, CTX-M-2, CTX-M-8, CTX-M-9 and CTX-M-25 (Figure 4.1). They are each named after the archetypal enzymes of their individual groups:

CTX-M-1 cluster; CTX-M-1, 3, 10, 12, 15, 22, 23,

CTX-M-2 cluster; CTX-M-2, 4, 5, 6, 7, 20, 76, 77,

CTX-M-8 cluster; CTX-M-8, 40, 63,

CTX-M-9 cluster; CTX-M-9, 14, 15, 16, 17, 18, 19,

CTX-M-25 cluster; and CTX-M-25, 26, 39, 41, 91.

The 5 groups differ from each other by ≥ 10 amino acid residues, while each group includes a number of minor allelic variants that differ from each other by ≤ 5 amino acid residues (Bevan et al., 2017). Furthermore, there are at least 4 *CTX-M* variants that show a hybrid structure: CTX-M-45, which is a hybrid of CTX-M-14 with a protein of unknown origin; CTX-M-64; CTX-M-123; and CTX-M-132 variants, which are hybrids of CTX-M-15 with different segments of CTX-M-14. Most of these variants are categorised under the CTX-M groups 1 and 9 (Canton et al., 2012).

Group 2 CTX-M-2/4-7/20/31/43/44/56/59/ 74/75/92/97/124/131 Group 1 CTX-M-5/76/77/95/124 CTX-M-1/3/10-12/15/22/23/28-30/ KLUA-1-6-7/8-12 32-34/36/42/52-55/57/58/60-62/ (K. ascorbata) Group 25 66/68/69/71/72/79/80/82/88/96/ CTX-M-25/26/39/ 101/107-109/114/116/117/133 41/89/91/94/100 CTX-M-45 CTX-M-78 (K. georgiana) CTX-M-3 (K. ascorbata) CTX-M-37 (K. cryocrescens) Group 8 CTX-M-132 CTX-M-8/40/63 KLUG-1 (K. georgiana CTX-M-123 Group 9 **KLUC Group** CTX-M-9/13/14/16-19/21/24/ CTX-M-64 KLUC-2-4 27/38/46-51/65/67/81/83-87/90/ 93/98-99/102/104-106/110-113/ 121/122/126/134 KLUC-1 (K. cryocrescens)

Figure 4.1 The phylogenetic tree of CTX-M-type ESBLs

Tree diagram showing the similarity among the enzymes of the CTX-M lineage and the clustering of members of different CTX-M groups. The tree was constructed with the TREEVIEW program on the basis of the amino acid sequence alignment of available sequences of CTX-M and cognate proteins from the Kluyvera spp. Available at the Lahey Clinic website (http://www.lahey.org/Studies/).

KLUY-1-4 (K. georgiana)

Clinically, each CTX-M type can be inhibited by a specific antibiotic and the inability of phenotypic methods to distinguish between the different types of CTX-M or to determine the gene responsible for the production of each enzyme, which results in treatment failures due to the inappropriate use of antibiotics, is a major problem. Using molecular-based techniques would help to guide the use of appropriate antibiotics, while have the additional advantage of being able to detect low-level genetic resistance. Genotyping method could provide a fast surveillance tool for detecting resistant strains.

In order to create an epidemiological snapshot of the antibiotic resistance pattern in UTI patients in North Wales, and to determine to what extent CTX-M-type ESBLs contribute to the resistance phenomenon detected in isolates from UTI patients in North Wales and whether these CTX-M-producing bacterial isolates carry the same or different *CTX-M* genes, 300 ESBL-producing isolates from 3 referral hospitals in North Wales (Ysbyty Gwynedd, Glan Clwyd Hospital and Wrexham Maelor Hospital) were sub cultured on UTI agar, from which I was able to identify *Enterococcus spp.*, *Escherichia coli* and coliform bacteria before they were tested by multiplex PCR using primers for genes' phylogenetic groups (Gp) (Table 2.1):

CTX-MGp1 (688bp), Gp2 (404bp), Gp9 (561bp) and Gp8/25 (326bp) based on percentages of similarity followed by gel electrophoresis (Dallenne et al., 2012).

Sequencing analysis of the amplicons was performed on approximately 50% of the samples that had bands with lengths corresponding with 1 of the CTX-M groups from each hospital in order to identify *blaCTX-M* genes. Sequences were then compared with known blactamase gene sequences via multiple-sequence alignments using the BLAST programme.

4.2 Results

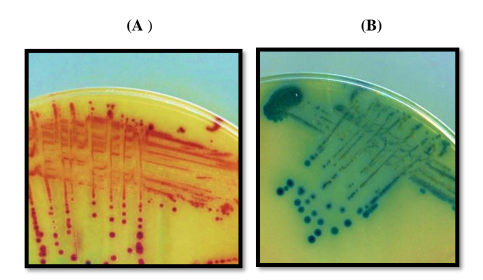
4.2.1 ESBL-producing isolates from Wrexham Maelor Hospital (1-100)

The screening of ESBL-producing isolates for the CTX-M groups was done in 3 main steps: sub culturing of preserved isolates from labelled bijoux tubes of nutrient agar slop, testing of extracted DNA for *blaCTX-M* 1, 2, 9 and 8/25 using multiplex PCR followed by electrophoresis and DNA sequencing analysis of amplicon PCR products.

4.2.1.1 Bacterial culturing on primary UTI agar

In order to achieve single colonies that would ensure the purity of the isolates to carry out the multiplex PCR identification of *Enterococcus spp.*, *Escherichia coli* (*E. coli*) and coliform bacteria, the preserved samples were cultured on primary UTI agar. Two types of colonies were observed in the 100 cultured samples (Figure 4.2): pink colonies (indicative of the presence of *E. coli*) and blue colonies (coliform bacteria). The growth of the colonies was recorded (Table 4.1) with their hospital lab numbers.

Figure 4.2 Sub culture isolates on UTI agar



Sub cultured isolates from UTI patients on primary UTI agar; (A) $E.\ coli$ with pink colonies (sample 44); (B) coliform bacteria with blue colonies (sample 55). The agar contains a chromogenic substrate that is hydrolysed by enzymes produced by these bacteria; β -D-glucosidase produced by $E.\ coli$ hydrolyses the chromogenic substrate to produce pink-red colonies, and β -glucosidase from coliform bacteria hydrolyses the chromogenic substrate to produce blue colonies (Oxoid Ltd., 2008).

Table 4.1 Bacterial identification based on colony characteristics for samples 1-100

Isolate number Lab number Date of subculture on colony colour 1 19152 7/5/2015 E. coli 2 18620 7/5/2015 E. coli 3 18257 7/5/2015 E. coli 4 18570 7/5/2015 E. coli 5 17893 7/5/2015 E. coli 6 19166 7/5/2015 E. coli 7 18322 7/5/2015 E. coli 8 18144 7/5/2015 E. coli 9 16548 7/5/2015 E. coli 10 17046 7/5/2015 E. coli 11 17464 7/5/2015 E. coli 12 17578 7/5/2015 E. coli 13 16920 7/5/2015 E. coli 14 17571 7/5/2015 E. coli 15 17084 7/5/2015 E. coli 16 17392 7/5/2015 E. coli 17 17212 7/5/2015 E. col				Indicative bacteria based
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15 17084 7/5/2015 E. coli 16 17392 7/5/2015 E. coli 17 17212 7/5/2015 E. coli 18 16785 7/5/2015 E. coli 19 16497 7/5/2015 E. coli 20 16632 7/5/2015 E. coli 21 17580 18/5/2015 Coliform 22 14379 18/5/2015 Coliform 23 13843 18/5/2015 E. coli 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 E. coli 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli </th <th>13</th> <th>16920</th> <th>7/5/2015</th> <th>E. coli</th>	13	16920	7/5/2015	E. coli
16 17392 7/5/2015 E. coli 17 17212 7/5/2015 E. coli 18 16785 7/5/2015 E. coli 19 16497 7/5/2015 E. coli 20 16632 7/5/2015 E. coli 21 17580 18/5/2015 Coliform 22 14379 18/5/2015 Coliform 23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 Coliform 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	14	17571	7/5/2015	Coliform
17 17212 7/5/2015 E. coli 18 16785 7/5/2015 E. coli 19 16497 7/5/2015 E. coli 20 16632 7/5/2015 E. coli 21 17580 18/5/2015 Coliform 22 14379 18/5/2015 Coliform 23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 E. coli 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	15	17084	7/5/2015	E. coli
18 16785 7/5/2015 E. coli 19 16497 7/5/2015 E. coli 20 16632 7/5/2015 E. coli 21 17580 18/5/2015 Coliform 22 14379 18/5/2015 Coliform 23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 E. coli 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	16	17392	7/5/2015	E. coli
19 16497 7/5/2015 E. coli 20 16632 7/5/2015 E. coli 21 17580 18/5/2015 Coliform 22 14379 18/5/2015 Coliform 23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 Coliform 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	17	17212	7/5/2015	E. coli
20 16632 7/5/2015 E. coli 21 17580 18/5/2015 Coliform 22 14379 18/5/2015 Coliform 23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 E. coli 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	18	16785	7/5/2015	E. coli
21 17580 18/5/2015 Coliform 22 14379 18/5/2015 Coliform 23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 E. coli 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	19	16497	7/5/2015	E. coli
22 14379 18/5/2015 Coliform 23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 Coliform 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	20	16632	7/5/2015	E. coli
23 13843 18/5/2015 Coliform 24 12456 18/5/2015 E. coli 25 13808 18/5/2015 Coliform 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	21	17580	18/5/2015	Coliform
24 12456 18/5/2015 E. coli 25 13808 18/5/2015 Coliform 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	22	14379	18/5/2015	Coliform
25 13808 18/5/2015 Coliform 26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	23	13843	18/5/2015	Coliform
26 14098 18/5/2015 E. coli 27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	24	12456	18/5/2015	E. coli
27 13651 18/5/2015 E. coli 28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	25	13808	18/5/2015	Coliform
28 13830 18/5/2015 E. coli 29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	26	14098	18/5/2015	E. coli
29 14122 18/5/2015 E. coli 30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	27	13651	18/5/2015	E. coli
30 14386 18/5/2015 E. coli 31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	28	13830	18/5/2015	E. coli
31 14152 18/5/2015 E. coli 32 13756 18/5/2015 E. coli	29	14122	18/5/2015	E. coli
32 13756 18/5/2015 E. coli	30	14386	18/5/2015	E. coli
	31	14152	18/5/2015	E. coli
33 13738 18/5/2015 E. coli	32	13756	18/5/2015	E. coli
	33	13738	18/5/2015	E. coli

34	13700	18/5/2015	E. coli
35	14384	18/5/2015	E. coli
36	13856	18/5/2015	E. coli
37	13745	18/5/2015	E. coli
38	13656	18/5/2015	E. coli
40	14758	18/5/2015	E. coli
41	15314	2/6/2015	E. coli
42	15378	2/6/2015	E. coli
43	15029	2/6/2015	E. coli
44	15340	2/6/2015	E. coli
45	14822	2/6/2015	E. coli
46	15194	2/6/2015	E. coli
47	15211	2/6/2015	E. coli
48	15060	2/6/2015	Coliform
49	24341	2/6/2015	E. coli
50	19677	2/6/2015	E. coli
51	19630	2/6/2015	E. coli
52	19658	2/6/2015	E. coli
53	19349	2/6/2015	E. coli
54	19274	2/6/2015	E. coli
55	20650	2/6/2015	Coliform
56	21176	16/6/2015	E. coli
57	21184	16/6/2015	E. coli
58	21087	16/6/2015	Coliform
59	20050	16/6/2015	E. coli
60	14773	16/6/2015	E. coli
61	15638	16/6/2015	E. coli
62	15643	16/6/2015	E. coli
63	15571	16/6/2015	E. coli
64	15393	16/6/2015	E. coli
65	15395	16/6/2015	E. coli
66	16013	16/6/2015	E. coli
67	16092	16/6/2015	E. coli
68	16368	16/6/2015	E. coli
69	15419	16/6/2015	E. coli
70	15457	16/6/2015	E. coli
71	15979	30/6/2015	E. coli
72	15881	30/6/2015	E. coli
73	15787	30/6/2015	E. coli
	L	l .	1

74	15920	30/6/2015	E. coli
75	14819	30/6/2015	E. coli
76	14838	30/6/2015	E. coli
77	15278	30/6/2015	E. coli
78	15190	30/6/2015	E. coli
79	14620	30/6/2015	E. coli
80	14437	30/6/2015	Coliform
81	14465	30/6/2015	E. coli
82	14479	30/6/2015	E. coli
83	14611	30/6/2015	Coliform
84	15051	30/6/2015	E. coli
85	14575	30/6/2015	E. coli
86	14506	30/6/2015	E. coli
87	15039	20/7/2015	E. coli
88	14648	20/7/2015	E. coli
89	15248	20/7/2015	E. coli
90	23810	20/7/2015	E. coli
91	23631	20/7/2015	E. coli
92	23955	20/7/2015	E. coli
93	22860	20/7/2015	E. coli
94	24147	20/7/2015	E. coli
95	23999	20/7/2015	Coliform
96	22802	20/7/2015	E. coli
97	23733	20/7/2015	E. coli
98	24236	20/7/2015	E. coli
99	24337	20/7/2015	E. coli
100	12986	20/7/2015	E. coli

4.2.1.2 Multiplex PCR amplification of β -lactamase genes *blaCTX-M* 1, 2, 9 and 8/25 followed by gel electrophoresis of samples 1-100

The isolates of interest were lysed to extract their DNA, which was used as the DNA template for PCR amplification. The total volume of 20 µL of the PCR reaction mixture was prepared for each cell lysate sample. After 2 h of amplification, the amplified PCR products were run on agarose gels. Around 91% of the samples had bands located at the length of 1 of *CTX-M* groups.

4.2.1.2.1 PCR gel analysis of the *blaCTX-M* genes (Isolates 1-14)

As shown in Figure 4.3, which depicts the gel analysis of isolates 1-14, 12 samples had bands with lengths that corresponded to 2 groups of CTX-M genes: *CTX-MGp1* (688bp) samples 1 (*E. coli*), 4 (*E. coli*), 5 (*E. coli*), 6 (*E. coli*), 7 (*E. coli*), 8 (*E. coli*), 9 (*E. coli*), 10 (*E. coli*), 11 (*E. coli*) and 12 (*E. coli*). Two samples had bands with lengths of *CTX-MGp2* (404bp), samples 2 (*E. coli*) and 3 (*E. coli*).

4.2.1.2.2 PCR gel analysis of the *blaCTX-M* genes (Isolates 15-28)

Gel analysis of samples 15-28 shows that 13 samples had bands with lengths that corresponded to *CTX-MGp1* (688bp): samples 15, 16, 17, 18, 19, 20, 24 and 26 (all *E. coli*) and 21, 22, 23 and 25 (all coliforms). Sample 17 had 2 bands, 1 with a length corresponding to *CTX-MGp1* (688bp) and another corresponding to *CTX-MGp9* (561bp) (Figure 4.4).

4.2.1.2.3 PCR gel analysis of the *blaCTX-M* genes (Isolates 29-42)

Fourteen samples (29-42) displayed bands located at the lengths of 3 groups of *CTX-M* genes on gel analysis: *CTX-MGp1* (688bp) samples 29, 30, 31, 32, 34, 35, 38 and 42 (all *E. coli*), CTX-MGp2 (404bp) samples 36, 40 and 41 (all *E. coli*) and *CTX-MGp9* samples 37 and 39 (all *E. coli*) (Figure 4.5).

4.2.1.2.4 PCR gel analysis of the *blaCTX-M* genes (Isolates 43-56)

This group of samples showed bands with lengths that corresponded to 2 groups of *CTX-M* genes: *CTX-MGp1* (688bp) samples 43, 44, 45, 46, 49, 50, 51, 52, 53 and 56 (all *E. coli*) and 48 and 55 (all coliforms) and *CTX-MGp9* (561bp) sample 47. Sample 54 had 2 bands; the first corresponded to *CTX-MGp1* and the other corresponded to CTX-MGp9 (Figure 4.6).

4.2.1.2.5 PCR gel analysis of the *blaCTX-M* genes (Isolates 57-70)

All samples of this group had bands with lengths that corresponded to *CTX-MGp1* (688bp): samples 57,59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69 and 70 (all *E. coli*) and 58 (coliform) (Figure 4.7).

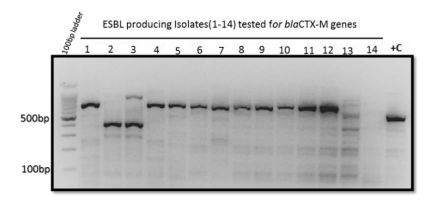
4.2.1.2.6 PCR gel analysis of the *blaCTX-M* genes (Isolates 71-84)

As shown in Figure 4.8, 11 samples displayed bands located at lengths of 2 groups of CTX-M genes: *CTX-MGp1* (688bp), which includes samples 72, 75, 76, 77, 79, 81, 82 and 84 (all *E. coli*), and *CTX-MGp9* (561bp), which includes samples 71 (*E. coli*) and 80 (coliform). Sample 73 had 2 bands; the first corresponded to *CTX-MGp1* and the other corresponded *to CTX-MGp9*.

4.2.1.2.7 PCR gel analysis of the *blaCTX-M* genes (Isolates 85-100)

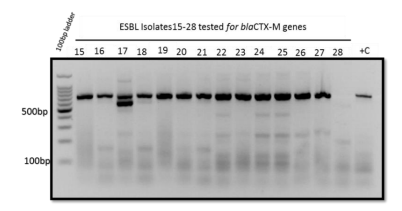
Based on gel analysis, 2 types of *CTX-M* genes seem to be carried by 15 isolates from this group: *CTX-MGp1* (688bp), which includes samples 86, 87, 88, 89, 90, 92, 93, 94, 96, 98 and 99 (all *E. coli*) and sample 95 (coliform); and *CTX-MGp9* (561bp), which includes samples 91, 97 and 100 (all *E.coli*) (Figure 4.9).

Figure 4.3 Multiplex PCR assay for the blaCTX-M genes, samples 1-14



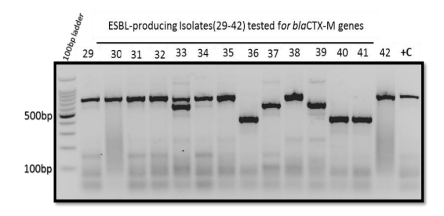
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1: the 100-bp ladder; lanes 2–15: the 14 ESBL samples. Twelve samples had bands with lengths that corresponded to two groups of *CTXM* genes; *CTXMGp1* (688 bp) samples (1, 4, 5, 6, 7,8,9,10,11 and 12) and *CTXMGp2* (404 bp) samples 2&3.Lane 16: positive control.

Figure 4.4 Multiplex PCR assay for the blaCTX-M genes, samples 15-28



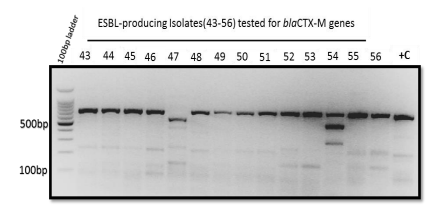
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1: 100-bp ladder. Lanes 2–15: 14 ESBL samples. Thirteen samples had bands with lengths that corresponded to *CTXMGp1* (688 bp); samples (15-27) and sample (3) had two bands one with length corresponded to CTXMGp1 (688 bp) and another with *CTXMGp9* (561bp). Lane 16: positive control.

Figure 4.5 Multiplex PCR assay for the blaCTX-M genes, samples 29-42



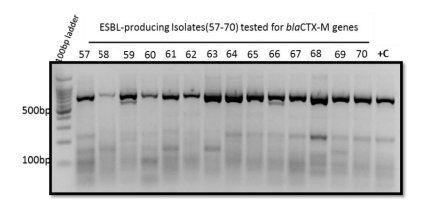
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1: 100-bp ladder. Lanes 2–15:14 ESBL samples. Fourteen samples had bands with lengths that corresponded to three groups of *CTXM* genes; *CTXMGp1* (688 bp) samples (29,30,31,32,34,35,38,42), *CTXMGp2* (404bp)samples(36,40 and 41) *CTXMGp9*(37&39).

Figure 4.6 Multiplex PCR assay for the blaCTX-M genes, samples 43-56



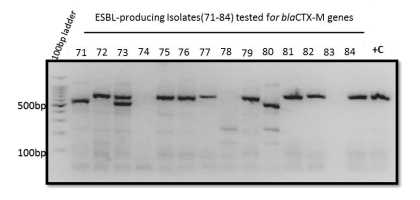
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1: the 100-bp ladder. Lanes 2–15: 14 ESBL samples. samples had bands with lengths that corresponded to two groups of *CTX-M* genes; CTXMGp1 (688 bp) samples (43,44,45,46,48,49,50,51,52,53,55,56) , *CTXMGp9* (561 bp) sample(47),sample 54 had two bands one corresponded to *CTXMGp1* and another to *CTXMGp9*. Lane 16:positive control.

Figure 4.7 Multiplex PCR assay for the blaCTX-M genes, samples 57-70



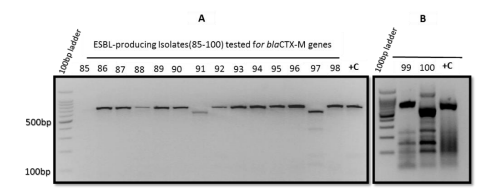
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1:100-bp ladder. Lanes 2–15: 14 ESBL samples. Fourteen samples had bands with lengths that corresponded to *CTXMGp1* (688 bp) samples (57-70). Lane 16: positive control.

Figure 4.8 Multiplex PCR assay for the blaCTX-M genes, samples 71-84



Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1:100-bp ladder. Lanes 2–15: 14 ESBL samples. Eleven samples had bands with lengths that corresponded to two groups of *CTXM* genes; *CTXMGp1* (688bp) samples (72,75,76,77,79,81,82 and 84) ,*CTXMGp9* (561bp) sample (71&80),sample (73) had two bands one corresponded to *CTXMGp1* and another to *CTXMGp9*. Lane 16:positive control.

Figure 4.9 Multiplex PCR assay for the blaCTX-M genes, samples 85-100



A) Gel analysis of *blaCTX-M* amplicon on 2% agarose gel. Lane 1: the 100bp ladder. Lanes 2–15: 14 ESBL samples. Thirteen samples had bands with lengths that corresponded to 2 groups of *CTX-M* genes: *CTX-MGp1* (688bp) samples (86, 87, 88, 89, 90, 92, 93, 94, 95, 96 and 98) and CTX-MGp9 (561bp) samples 91 and 97. Lane 16: positive control. (**B**) Gel analysis of *blaCTX-M* amplicon on 2% agarose gel. Lane 1: the 100bp ladder. Lanes 2–3: 2 ESBL samples. Two samples had bands with lengths that corresponded to 2 groups of *CTX-M* genes: *CTX-MGp1* (688bp) sample 99 and *CTX-MGp9* (561bp) sample 100. Lane 16: positive control.

The screening of ESBL-producing isolates from Wrexham Maelor Hospital (1–100) for the CTX-M groups using multiplex PCR followed by gel analysis shows, based on the amplicon size, that the predominant CTX-M group is group 1 (Table 4.2).

Table 4.2 PCR amplicon sizes of the blaCTX-M genes, samples 1-100

CTX-M group	Amplicon size (bp)	Sample No
CTX-M group 1	688	1, 4, 5, 6, 7, 8, 9, 10, 11, 12, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 29, 30, 31, 32, 34, 35, 38, 42, 43, 44, 45, 46, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 72, 73, 75, 76, 77, 79, 81, 82, 84, 86, 87, 88, 89, 90, 92, 93, 94, 95, 96, 98 and 99.
CTX-M group 2	404	2, 3, 36, 40, 41, 91, 97 and 100.
CTX-M group 9	561	17, 37, 39, 47, 71, 80.
CTX-M group 8/25	326	

4.2.1.3 Sequencing analysis of multiplex PCR products (1-100)

The DNA of amplified PCR was purified using the ISOLATE II PCR kit for sequencing in order to identify the *CTX-M* genes detected in the multiplex PCR assays. The sequencing analysis was performed on 47 (50%) samples that displayed bands with sizes corresponding to *CTX-M* groups. The sequenced samples were compared with DNA sequences of known b-lactamase genes using the BLAST programme. Table 4.3 shows the BLAST output of the sequenced samples.

Table 4.3 Sequencing analysis of positive samples from 1-100

		Accession	CTX-M group		ıp	DNA	
Sample No	Blast output	number	Gp1	Gp2	Gp9	—— Gp8/25	sequence
1	CTX-M-15 gene	KP325146.1	√				Appendix I
3	CTX-M-59 gene	MH661248.1		√			Appendix I
5	CTX-M-3 gene	CP034325.1	√				Appendix I
7	CTX-M-15 gene	KR338941.1	✓				Appendix I
9	CTX-M-15 gene	KY640551.1	√				Appendix I
11	CTX-M-66 gene	NG_049017.1	√				Appendix I
15	Klebsiella pneumoniae complete genome	CP011985.1					Appendix I
17 (lower band)	CTX-M-108 gene	JF274245.1			✓		Appendix I
19	CTX-M-15 gene	MK113957.1	√				Appendix I
21	CTX-M-15 gene	KY640536.1	√				Appendix I
23	CTX-M-32 gene	MH900527.1	√				Appendix I
25	CTX-M-15 gene	MH900522.1	√				Appendix I
27	CTX-M-15 gene	KY640528.1	√				Appendix I
29	CTX-M-55 gene	MH900523.1	√				Appendix I
31	E coli strain MNCRE44, complete genome	CP010876.1					Appendix I
33(lower band)	CTX-M-9 gene	MF797877.1			√		Appendix I
35	CTX-M-15 gene	MK113960.1	√				Appendix I
36	CTX-M-27 gene	MH900525.1			√		Appendix I
41	CTX-M-2 gene	MH661245.1			✓		Appendix I
43	CTX-M-15 gene	MH900522.1	√				Appendix I
45	CTX-M-15 gene	MK113960.1	√				Appendix I
47	CTX-M-9 gene	KT459753.1			√		Appendix I
49	CTX-M-15 gene	MK405591	√				Appendix I
51	CTX-M-15 gene	LT628520	√				Appendix I
54(lower band)	<i>CTX-M-27</i> gene	MH900525			√		Appendix I
57	CTX-M-15 gene	CP040398	√				Appendix I
59	CTX-M-1 gene	MH037035	√				Appendix I

61	CTX-M-15 gene	KY640536	✓		Appendix I
63	E.coli4928STDY7071340	LR607331			Appendix I
	genome assembly				
65	CTX-M-55 gene	MF958462	✓		Appendix I
67	CTX-M-15 gene	KP325147	✓		Appendix I
69	<i>CTX-M-163</i> gene	KP681698	✓		Appendix I
71	CTX-M-15 gene	MK113957	✓		Appendix I
73	CTX-M-16 gene	AY029068		✓	Appendix I
75	CTX-M-15 gene	KP325146	✓		Appendix I
77	CTX-M-15 gene	KY640536	✓		Appendix I
80	CTX-M-9 gene	MF797877		✓	Appendix I
82	CTX-M-160 gene	NG_048945	✓		Appendix I
84	CTX-M-15 gene	LT628516	✓		Appendix I
86	CTX-M-15 gene	KY640534	✓		Appendix I
88	CTX-M-225 gene	NG_064720.	✓		Appendix I
90	CTX-M-1 gene	MG255315	✓		Appendix I
91	Ecoli strain VRES	LR607054			Appendix I
	hospital6495320 genome				
	assembly				
93	CTX-M-172 gene	NG_048957	✓		Appendix I
95	CTX-M-15 gene	LT628516.	√		Appendix I
97	CTX-M-9 gene	MF797877		√	Appendix I
98	CTX-M-15 gene	MK113957	√		Appendix I
100	CTX-M-15 gene	MH900525		√	Appendix I

The BLAST output shows that the most common (dominant) *CTX-M* in Wrexham Maelor Hospital isolates is *CTX-M-15*, which is classified under group 1 as 23 (48.9%) shows high identity with *CTX-M-15*, followed by *CTX-M-9*; 4 (8.5%), *CTX-M-27*, *CTX-M-2 gene* and *CTX-M-55*; 2 (4.2%), *CTX-M-59*; 1 (2.1%), *CTX-M-3*; 1 (2.1%), *CTX-M-172* gene; 1 (2.1) *CTX-M-66*; 1 (2.1%), *CTX-M-108*; 1 (2.1%), *CTX-M-32*; 1 (2.1%), *CTX-M-1*; 1 (2.1%), *CTX-M-160*; 1 (2.1%), *CTX-M-163*; 1 (2.1%) and *CTX-M-225*; 1 (2.1%).

4.2.2 ESBL-producing isolates from Glan Clwyd Hospital (101-200)

The 3 steps described in section 4.2.1 were carried out in order to screen the Glan Clwyd Hospital isolates for CTX-M genes.

4.2.2.1 Bacterial culturing on primary UTI agar

Isolates from Glan Clwyd Hospital were cultured on primary UTI agar as prescribed in section 4.2.1.1. The growth of the colonies was recorded (Table 4.4) according to the hospital lab numbers.

 $Table \ 4.4 \ Bacterial \ identification \ based \ on \ colony \ characteristics \ for \ samples \ 101-200$

Isolate number	Lab number	Date of subculture	Indicative bacteria
			based on colony colour
101	12457	17/9/2015	E. coli
102	107166	17/9/2015	E. coli
103	12249	17/9/2015	E. coli
104	12261	17/9/2015	E. coli
105	12823	17/9/2015	E. coli
106	12906	17/9/2015	E. coli
107	112575	17/9/2015	Coliform
108	12808	17/9/2015	E. coli
109	12847	17/9/2015	E. coli
110	12167	17/9/2015	E. coli
111	12609	17/9/2015	E. coli
112	11512	17/9/2015	E. coli
113	13108	17/9/2015	E. coli
114	10604	17/9/2015	Coliform
115	12734	17/9/2015	E. coli
116	11471	17/9/2015	E. coli
117	12472	17/9/2015	E. coli
118	12722	17/5/2015	E. coli
119	13138	17/5/2015	E. coli
120	107156	17/9/2015	E. coli
121	19800	17/9/2015	E. coli
122	22522	17/9/2015	E. coli
123	225512	17/9/2015	E. coli
124	22903	17/9/2015	E. coli
125	22867	17/9/2015	E. coli
126	22664	18/9/2015	E. coli
127	22941	18/9/2015	E. coli
128	22877	18/9/2015	E. coli
129	22803	18/9/2015	E. coli
30	22847	18/9/2015	E. coli
131	22429	18/9/2015	E. coli
132	22426	18/9/2015	E. coli
133	22658	5/10/2015	E. coli
134	22753	5/10/2015	Coliform
135	22662	5/10/2015	E. coli

136	22655	5/10/2015	E. coli
		5/10/2015	
137	22804		E. coli
138	23621	5/10/2015	E. coli
139	24056	5/10/2015	E. coli
140	23789	5/10/2015	E. coli
141	23870	5/10/2015	E. coli
142	23641	5/10/2015	E. coli
143	23799	5/10/2015	E. coli
144	28433	5/10/2015	E. coli
145	20012	5/10/2015	E. coli
146	18481	5/10/2015	E. coli
147	19702	5/10/2015	E. coli
148	18431	5/10/2015	Coliform
149	18720	5/10/2015	E. coli
150	18694	5/10/2015	E. coli
151	19037	5/10/2015	E. coli
152	18514	5/10/2015	E. coli
153	18251	2/11/2015	E. coli
154	19107	2/11/2015	E. coli
155	19927	2/11/2015	E. coli
156	18601	2/11/2015	E. coli
157	18543	2/11/2015	E. coli
158	18477	2/11/2015	E. coli
159	21810	2/11/2015	E. coli
160	21677	2/11/2015	E. coli
161	21640	2/11/2015	E. coli
162	20782	2/11/2015	Coliform
163	21649	2/11/2015	E. coli
164	21793	2/11/2015	E. coli
165	22041	2/11/2015	E. coli
166	21761	2/11/2015	E. coli
167	21988	2/11/2015	E. coli
168	22192	2/11/2015	E. coli
169	20541	2/11/2015	E. coli
170	20510	2/11/2015	E. coli
171	20665	2/11/2015	E. coli
172	21862	2/11/2015	E. coli
173	20853	2/11/2015	Coliform
174	22238	2/11/2015	E. coli
L			

175	21361	2/11/2015	E. coli
176	25798	2/11/2015	E. coli
177	26377	2/11/2015	E. coli
178	25705	2/11/2015	E. coli
179	28162	2/11/2015	E. coli
180	25927	2/11/2015	E. coli
181	28381	6/1/2016	E. coli
182	25784	6/1/2016	E. coli
183	25886	6/1/2016	E. coli
184	26672	6/1/2016	E. coli
185	28022	6/1/2016	E. coli
186	28211	6/1/2016	E. coli
187	26280	6/1/2016	E. coli
188	28374	6/1/2016	E. coli
189	28006	6/1/2016	E. coli
190	25956	6/1/2016	E. coli
191	28333	6/1/2016	E. coli
192	28502	6/1/2016	E. coli
193	28511	6/1/2016	E. coli
194	28457	6/1/2016	E. coli
195	26668	6/1/2016	E. coli
196	28648	6/1/2016	E. coli
197	26576	6/1/2016	E. coli
198	26894	6/1/2016	E. coli
199	26864	6/1/2016	Coliform
200	28084	6/1/2016	E. coli

4.2.2.2 Multiplex PCR amplification of β -lactamase genes blaCTX-M 1, 2, 9 and 8/25, followed by gel electrophoresis of samples 101-200

The extracted genomic DNA of each isolate was amplified using multiple PCR as explained in section 4.2.1.2. Around 73% of the samples had bands located at the length of 1 of the *CTX-M* groups.

4.2.2.2.1 PCR gel analysis of the *blaCTX-M* genes (Isolates 101-114)

Thirteen samples from this group displayed bands located that corresponded to 2 groups of *CTX-M* genes: *CTX-MGp1* (688bp), which includes samples 101, 103, 104, 105, 109, 110, 112

and 113 (all *E. coli*) in addition to samples 107 and 114 (all coliforms), and *CTX-MGp2* (404bp), which includes samples 106, 108 and 111 (all *E. coli*) (Figure 4.10).

4.2.2.2.2 PCR gel analysis of the *blaCTX-M* genes (Isolates 115-128)

As shown in Figure 4.11, 7 samples had bands with lengths that corresponded to *CTX-MGp1* (688 bp): samples 115, 118, 120, 122, 124 and 125 (all *E. coli*).

4.2.2.2.3 PCR gel analysis of the *blaCTX-M* genes (Isolates 129-142)

The gel analysis of these samples shows that 2 groups of CTX-M seem to be carried by the isolates: *CTX-MGp1* (688bp), which includes samples 132, 133, 134, 135, 136, 137, 138, 140, 141 and 142 (all *E. coli*), and *CTX-MGp2* (404bp), which includes sample 129 (Figure 4.12).

4.2.2.2.4 PCR gel analysis of the *blaCTX-M* genes (Isolates 143-156)

Bands with lengths corresponding to *CTX-MGp1* (688bp) were displayed in samples 143, 144, 145, 146, 147, 150 and 151 (all *E. coli*) in addition to sample 48 (coliform) (Figure 4.13).

4.2.2.2.5 PCR gel analysis of the *blaCTX-M* genes (Isolates 157-170)

Samples from this group displayed bands located at the lengths of 3 *CTX-M* groups: *CTX-MGp1* (688bp), which includes samples 157, 160, 161, 164, 165, 166, 167 and 168 (all *E. coli*); CTX-MGp2 (404bp), which includes sample 158 (*E. coli*); and *CTX-MGp9*, which includes sample 159 (*E. coli*) (Figure 4.14).

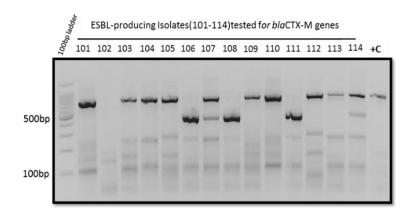
4.2.2.2.6 PCR gel analysis of the *blaCTX-M* genes (Isolates 171-184)

As shown in Figure 4.15, 12 samples—172, 174, 175, 176, 177, 178, 179, 180, 181, 182 and 184 (all *E. coli*) in addition to sample 173 (coliform)—had bands with sizes corresponding to *CTX-MGp1* (688bp).

4.2.2.2.7 PCR gel analysis of the *blaCTX-M* genes (Isolates 185-200)

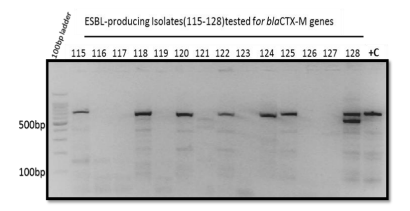
Twelve samples from this group displayed bands located at the lengths of *CTX-MGp1* (688bp): 185, 186, 187, 189, 190, 192, 193, 194, 195, 196, 198 and 200 (*all E.coli*) (Figure 4.16).

Figure 4.10 Multiplex PCR assay for the blaCTX-M genes, samples 101-114



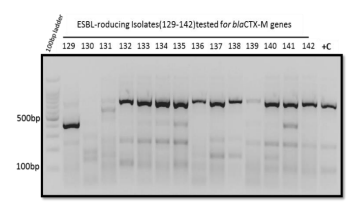
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Thirteen samples had bands with lengths that corresponded to two groups of *CTX-M* genes; *CTX-MGp1* (688bp) samples (101,103,104,105,107,109,110,112,113,114), CTXMGp2 (404bp) sample (106,108 and 111). Lane 16: positive control.

Figure 4.11 Multiplex PCR assay for the blaCTX-M genes, samples 115-128



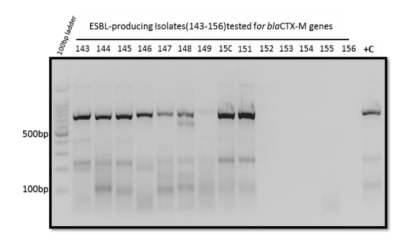
Analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1: the 100bp ladder; lanes 2–15: 14 ESBL samples. Seven samples had bands with lengths that corresponded to *CTXMGp*1 (688 bp) samples (115,118,120,122,124,125), sample (128) had two bands one corresponded to CTXMGp1 and another to *CTXMGp9*. Lane 16: positive control.

Figure 4.12 Multiplex PCR assay for the blaCTX-M genes, samples 129-242



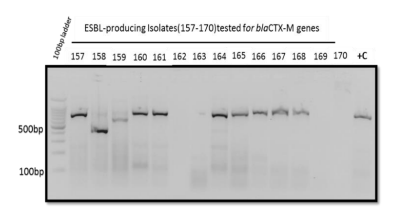
Gel analysis of *blaCTX-M* amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Eleven samples had bands with lengths that corresponded to two groups of CTXM genes; CTXMGp1 (688bp) samples (132,133,134,135,136,137,138,140,141,142), CTXMGp2 (404bp) sample (129). Lane 16: positive control.

Figure 4.13 Multiplex PCR assay for the blaCTX-M genes, samples 143-156



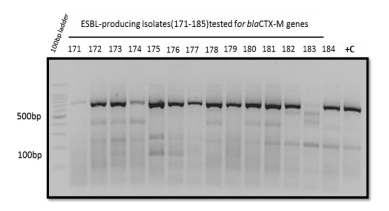
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Eleven samples had bands with lengths that corresponded to CTXMGp1 (688bp) samples (143,144,145,146,147,148,150,151). Lane 16: positive control.

Figure 4.14 Multiplex PCR assay for the blaCTX-M genes, samples 157-170



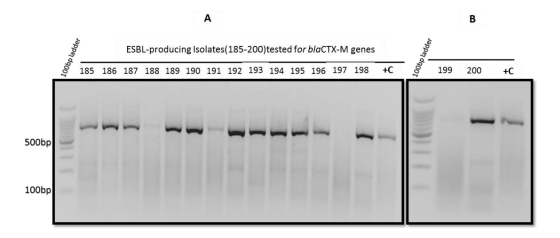
Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Ten samples had bands with lengths that corresponded to two groups of CTXM genes; CTXMGp1 (688bp) samples (157,160,161,164,165,166,167,168), CTXMGp2 (404bp) sample (158), CTXMGp9 (sample159).Lane 16: positive control.

Figure 4.15 Multiplex PCR assay for the blaCTX-M genes, samples 171-184



Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Twelve samples had bands with lengths that corresponded to *CTXMGp1* (688bp) samples (172,173,174,175,176,177,178,179,180,181,182 and 184). Lane 16: positive control.

Figure 4.16 Multiplex PCR assay for the blaCTX-M genes, samples 185-200



(A) Gel analysis of *blaCTX-M* amplicon on 2% agarose gel. Lane 1: the 100bp ladder. Lanes 2–15: 14 ESBL samples. Eleven samples had bands with lengths that corresponded to *CTX-MGp1* (688bp) samples 185, 186, 187, 189, 190, 192, 193, 194, 195, 196 and 198. Lane16: positive control. (B) Gel analysis of *blaCTX-M* amplicon on 2% agarose gel. Lane 1: the 100bp ladder. Lanes 2–3: 2 ESBL samples. One sample had bands with lengths that corresponded to *CTX-MGp1* (688bp) sample 200. Lane 16: positive control.

The screening of ESBL-producing isolates from Wrexham Maelor Hospital (101–200) for the CTX-M groups using multiplex PCR followed by gel analysis shows, based on the amplicon size, that the predominant CTX-M group is group 1 (Table 4.5).

Table 4.5 PCR amplicon sizes of the blaCTX-M genes, samples 101-200

CTX-M group	Amplicon size (bp)	Sample No
CTX-M group 1	688	101, 103, 104, 105, 107, 109, 110, 112, 113, 114, 115, 118, 120, 122, 124,125, 132, 133, 134, 135, 136, 137, 138, 140, 141, 142, 143, 144, 145, 146, 147, 148, 150, 151, 157, 160, 161, 164, 165, 166, 167 168, 172, 173, 174, 175, 176, 177, 178, 179, 180, 181, 182, 184, 185, 186, 187, 189, 190, 192, 193, 194, 195, 196, 198 and 200.
CTX-M group 2	404	106, 108, 111, 129, 158
CTX-M group 9	561	159
CTX-M group 8/25	326	

4.2.2.3 Sequencing analysis of multiplex PCR products (201-200)

The sequencing analysis was performed on 46 (60%) samples that displayed bands with sizes that corresponded with 1 of the CTX-M groups. The BLAST programme was then used to compare each sequence with DNA sequences of known b-lactamase genes. Table 4.6 shows the BLAST output of the sequenced samples.

Table 4.6 Sequencing analysis of positive samples from 101-200

		Accession	CTX-M group		up	DNA	
Sample No	Blast output	number	Gp1	Gp2	Gp9	—— Gp8/25	sequence
101	CTX-M-15 gene	KP325147.1	√				Appendix II
103	CTX-M-15 gene	MH900522.1	√				Appendix II
106	CTX-M-59 gene	MH661247.1		✓			Appendix II
108	CTX-M family	MK896928.1					Appendix II
109	CTX-M-15 gene	MK113960.1	√				Appendix II
111	Escherichia coli O1:H42 strain CLSC36 chromosome, complete genome.Sequence	CP041300.1					Appendix II
113	CTX-M-15 gene	MH900522.1	✓				Appendix II
115	CTX-M-15 gene	MK405591.1	√				Appendix II
118	CTX-M-15 gene	MH900522.1	√				Appendix II
120	CTX-M-15 gene	KY640536.1	√				Appendix II
122	<i>CTX-M-203</i> gene	NG_055269.1					Appendix II
124	CTX-M-66 gene	NG_049017.1	✓				Appendix II
128(lower band)	Escherichia coli strain 4928STDY7071340 genome assembly	LR607331.1					Appendix II
129	CTX-M-9 gene	MF797877.1			✓		Appendix II
132	CTX-M-15 gene	MH523447.1	✓				Appendix II
135	CTX-M-32 gene	MH900527.1	✓				Appendix II
136	CTX-M-15 gene	MH900522.1	√				Appendix II
138	CTX-M-15 gene	KY640528.1	√				Appendix II
141	CTX-M-15 gene	CP040398.1	√				Appendix II
142	CTX-M-1 gene	MH037035.1	✓				Appendix II
143	CTX-M-108 gene	JF274245.1	√				Appendix II
145	CTX-M-15 gene	MK113958.1	√				Appendix II
147	CTX-M-15 gene	MH900522.1	√				Appendix II
150	CTX-M-15 gene	MH900522.1	✓				Appendix II
151	CTX-M-15 gene	MH900522.1	√				Appendix II
157	CTX-M-65 gene	KX495605.1	√				Appendix II

158	CTX-M-51 gene	NG_049002.1			√	Appendix II
159	Escherichia coli strain MNCRE44, complete genome	CP010876.1				Appendix II
161	CTX-M-15 gene	KP325147.1	√			Appendix II
164	CTX-M-15 gene	MH900522.1	√			Appendix II
166	CTX-M-15 gene	KY640528.1	√			Appendix II
168	CTX-M-59 gene	MH661247.1		√		Appendix II
172	CTX-M-15 gene	CP040398.1	√			Appendix II
174	CTX-M-172 gene	NG_048957.1				Appendix II
175	CTX-M-15 gene	MH900522	✓			Appendix II
177	CTX-M-15 gene	MK113956.1	√			Appendix II
179	CTX-M-3 gene	CP034325.1				Appendix II
181	CTX-M-15 gene	KY640536.1	✓			Appendix II
184	CTX-M-15 gene	NG_048947.1	√			Appendix II
185	CTX-M-15 gene	KP325147.1	√			Appendix II
187	CTX-M-15 gene	MH523447.1				Appendix II
190	CTX-M-15 gene	LT628518.1	√			Appendix II
193	CTX-M-55 gene	MH900523.1	√			Appendix II
195	CTX-M-2 gene	MH661246.1		✓		Appendix II
198	CTX-M-27 gene	MH900525.1			√	Appendix II
200	CTX-M-15 gene	MH891569.1	✓			Appendix II

The sequence analysis using the BLAST tool shows that 27 (58%) of Glan Clwyd Hospital isolates had a high identity with *CTX-M-15*, followed by *CTX-M-59*: 2 (4%), *CTX-M-66*: 1 (2%), *CTX-M-203*: 1 (2%), *CTX-M-9*: 1 (2%), *CTX-M-32*: 1 (2%), *CTX-M-1*: 1 (2%), *CTX-M-108*: 1 (2%), *CTX-M-65*: 1 (2%), *CTX-M-51*: 1 (2%), *CTX-M-55*: 1 (2%), *CTX-M-172*: 1 (2%), *CTX-M-3*: 1 (2%), *CTX-M-2*: 1 (2%), *CTX-M-27*: 1 (2%).

4.2.3 ESBL-producing isolates from Ysbyty Gwynedd (YG) Hospital (201-300)

The 3 steps described in section 4.2.1 were carried out in order to screen the YG Hospital isolates for CTX-M genes.

4.2.3.1 Bacterial culturing on primary UTI agar

Isolates from YG Hospital were cultured on primary UTI agar as prescribed in section 4.2.1.1. The growth of the colonies was recorded in Table 4.5 according to the hospital lab number.

 $Table \ 4.7 \ Bacterial \ identification \ based \ on \ colony \ characteristics \ for \ samples \ 201-300$

Isolate number	Lab number	Date of subculture	Indicative bacteria
			based on colony colour
201	27697	12/05/2016	Coliform
202	28282	12/05/2016	Coliform
203	27855	12/05/2016	Coliform
204	28469	12/05/2016	E. coli
205	25696	12/05/2016	E. coli
206	23484	12/05/2016	E. coli
207	23623	12/05/2016	E. coli
208	23414	12/05/2016	E. coli
209	23842	12/05/2016	E. coli
210	12030	12/05/2016	E. coli
211	12294	12/05/2016	E. coli
212	12675	12/05/2016	E. coli
213	12640	12/05/2016	E. coli
214	13096	12/05/2016	E. coli
215	15309	12/05/2016	E. coli
216	15227	12/05/2016	E. coli
217	20057	12/05/2016	E. coli
218	27700	10/08/2016	Coliform
219	28287	10/08/2016	Coliform
220	11773	10/08/2016	Coliform
221	28316	10/08/2016	Coliform
222	23635	10/08/2016	E. coli
223	23456	10/08/2016	E. coli
224	23670	10/08/2016	E. coli
225	23362	10/08/2016	E. coli
226	12472	10/08/2016	E. coli
227	12564	10/08/2016	Coliform
228	12672	10/08/2016	Coliform
229	12447	10/08/2016	Coliform
230	13428	10/08/2016	Coliform
231	15581	10/08/2016	E. coli
232	15217	10/08/2016	E. coli
233	18069	10/08/2016	Coliform
234	27717	10/08/2016	E. coli
235	15306	10/08/2016	E. coli

236	15468	10/08/2016	E. coli
237	13373	10/08/2016	E. coli
238	13388	10/08/2016	E. coli
239	13232	10/08/2016	E. coli
240	13291	10/08/2016	E. coli
241	13396	10/08/2016	Coliform
242	12405	10/08/2016	E. coli
243	23471	10/08/2016	E. coli
244	23640	10/08/2016	Coliform
245	23631	10/08/2016	E. coli
246	28223	10/08/2016	E. coli
247	28096	10/08/2016	E. coli
248	28194	08/02/2017	Coliform
249	27651*	08/02/2017	Coliform
250	27651*	08/02/2017	E. coli
251	27717	08/02/2017	E. coli
252	23946	08/02/2017	E. coli
253	27301	08/02/2017	E. coli
254	27297	08/02/2017	E. coli
255	27181*	08/02/2017	E. coli
256	27181*	08/02/2017	Coliform
257	26083	08/02/2017	Coliform
258	26988	08/02/2017	E. coli
259	26548	08/02/2017	E. coli
260	25942	08/02/2017	E. coli
261	25673	08/02/2017	E. coli
262	23781	08/02/2017	Coliform
263	24954	08/02/2017	E. coli
264	24543	08/02/2017	E. coli
265	24623	08/02/2017	E. coli
266	23096	08/02/2017	E. coli
267	25706	08/02/2017	E. coli
278	23759	08/02/2017	E. coli
279	24026	08/02/2017	E. coli
270	24526	08/02/2017	E. coli
271	23760	08/02/2017	E. coli
272	23787	11/04/2017	E. coli
273	23789	11/04/2017	E. coli
274	23975	11/04/2017	Coliform
		1	

275	24542	11/04/2017	E. coli
276	24654	11/04/2017	E. coli
277	25006	11/04/2017	E. coli
278	25065	11/04/2017	E. coli
279	25864	11/04/2017	Coliform
280	24785	11/04/2017	E. coli
281	25792	11/04/2017	Coliform
282	26477	11/04/2017	Coliform
283	25691	11/04/2017	E. coli
284	25898	11/04/2017	E. coli
285	26523	11/04/2017	E. coli
286	26592	11/04/2017	E. coli
287	27083	11/04/2017	E. coli
288	26602	11/04/2017	E. coli
289	26852	11/04/2017	E. coli
290	26822	11/04/2017	E. coli
291	26675	11/04/2017	E. coli
292	27449	11/04/2017	E. coli
293	27554	11/04/2017	E. coli
294	27442	11/04/2017	E. coli
295	27611	11/04/2017	E. coli
296	27184	11/04/2017	E. coli
297	27303	11/04/2017	E. coli
398	27231	11/04/2017	E. coli
399	27383	11/04/2017	E. coli
300	27121	11/04/2017	E. coli

4.2.3.2 Multiplex PCR amplification of β -lactamase genes *blaCTX-M* 1, 2, 9 and 8/25 followed by gel electrophoresis of samples 201-300

The extracted genomic DNA of each isolate was amplified using multiple PCR as explained in section 4.2.1.2. Around 89% of the samples had bands located at the length of 1 of the *CTX-M* groups.

4.2.3.2.1 PCR gel analysis of the *blaCTX-M* genes (Isolates 201-214)

Based on the sizes of the displayed bands, these isolates seem to carry *CTX-M* genes: *CTX-MGp1* (688bp) samples 201 (coliforms) and 204, 205, 206, 207, 208, 209, 210, 211 and 212 (all *E. coli*), in addition to *CTX-MGp2* (404bp) samples 202 (*E. coli*) and 203 (coliform) (Figure 4.17).

4.2.3.2.2 PCR gel analysis of the *blaCTX-M* genes (Isolates 215-228)

As shown in Figure 4.18, 13 samples from these isolates corresponded to 2 groups of *CTX-M* genes: *CTX-MGp1* (688bp), which includes samples 215, 216, 217, 218, 221, 222, 223 and 225 (all *E. coli*) and 219, 220 and 227 (all coliforms), in addition to CTX-MGp9 (561bp) samples 226 (*E. coli*) and 228 (coliform).

4.2.3.2.3 PCR gel analysis of the *blaCTX-M* genes (Isolates 229-242)

Nine isolates from this group—229, 231, 236, 237, 238, 239 and 242 (all *E. coli*) in addition to 240 and 241 (all coliforms)—displayed with lengths that corresponded to *CTX-MGp1* (688bp) (Figure 4.19).

4.2.3.2.4 PCR gel analysis of the *blaCTX-M* genes (Isolates 243-256)

Three *CTX-M* groups seem to be carried by these isolates based on the displayed bands: *CTX-MGp1* (688bp) in samples 244 (coliform), 245 (*E. coli*), 247 (*E. coli*), 248 (coliform), 249 (coliform), 250 (*E. coli*), 251 (*E. coli*), 253 (*E. coli*), 255 (*E. coli*) and 256 (coliform); *CTX-MGp2* (404bp) in samples 252 and 254 (all *E. coli*); and *CTX-MGp9* (561bp) in sample 246 (*E.coli*) (Figure 4.20).

4.2.3.2.5 PCR gel analysis of the *blaCTX-M* genes (Isolates 257-270)

Only 1 type, *CTX-MGp1* (688bp), of the CTX-M genes seemed to be detected in11 isolates of this group, which includes samples 258, 259, 260, 263, 264, 265, 266 and 267 (all *E.coli*), in addition to samples 268, 257 and 262 (all coliforms) (Figure 4.21).

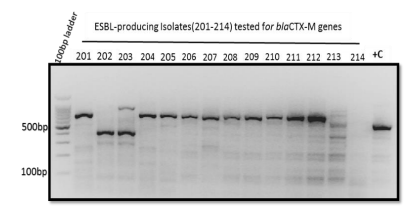
4.2.3.2.6 PCR gel analysis of the *blaCTX-M* genes (Isolates 271-284)

Gel analysis of this group shows that the 14 samples had bands with lengths that corresponded to *CTX-MGp1* (688 bp): 271, 272, 273, 274, 275, 276, 276, 277 and 278 (all *E. coli*) in addition to 279, 280 and 281 (all coliforms) (Figure 4.22).

4.2.3.2.7 PCR gel analysis of the *blaCTX-M* genes (Isolates 285-300)

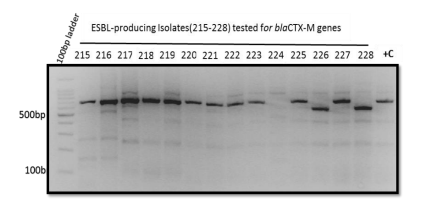
The displayed bands of the gel analysis show that 2 types of CTX-M genes seem to be carried by this group of isolates: *CTX-MGp1* (688bp) samples, which include samples 285, 286, 287, 288, 289, 290, 291, 292, 295, 296, 297, 298, 299 and 300 (all *E.coli*) and *CTX-MGp9* (561bp) samples 293 and 294 (both *E.coli*) (Figure 4.23).

Figure 4.17 Multiplex PCR assay for the blaCTX-M genes, samples 201-214



Gel analysis of *bla*CTXM amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Twelve samples had bands with lengths that corresponded to two groups of CTXM genes; CTXMGp1 (688bp) samples (201,204,205,206,207,208,209,210,211,212), CTXMGp2 (404bp) sample (202&203). Lane 16: positive control.

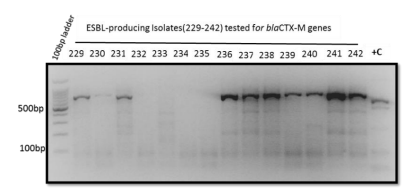
Figure 4.18 Multiplex PCR assay for the blaCTX-M genes, samples 215-228



Gel analysis of blaCTX-M amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Fourteen samples had bands with lengths that corresponded to two groups of CTX-M genes; CTXMGp1 (688bp) samples (244,245,247,248,249,250,251,253,255,256), CTXMGp2 (404bp) sample (252&2254), CTXMGp9 (561bp) sample (246). Lane 16: positive control.

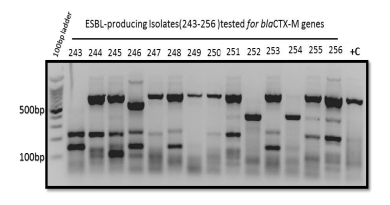
107

Figure 4.19 Multiplex PCR assay for the blaCTX-M genes, samples 229-242



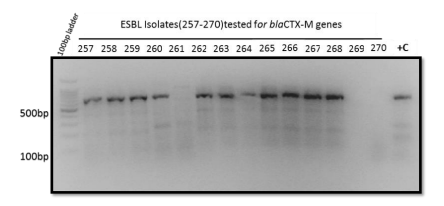
Gel analysis of blaCTX-M amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Nine samples had bands with lengths that corresponded to CTXMGp1 (688bp) samples (229,231,236,237,238,239,240,241,242). Lane 16: positive control.

Figure 4.20 Multiplex PCR assay for the blaCTX-M genes, samples 243-256



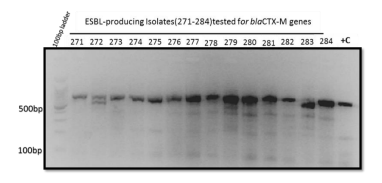
Gel analysis of blaCTX-M amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Fourteen samples had bands with lengths that corresponded to two groups of CTX-M genes; CTXMGp1 (688bp) samples (244,245,247,248,249,250,251,253,255,256), CTXMGp2 (404bp) sample (252&2254), CTXMGp9 (561bp) sample (246). Lane 16: positive control.

Figure 4.21 Multiplex PCR assay for the blaCTX-M genes, samples 257-270



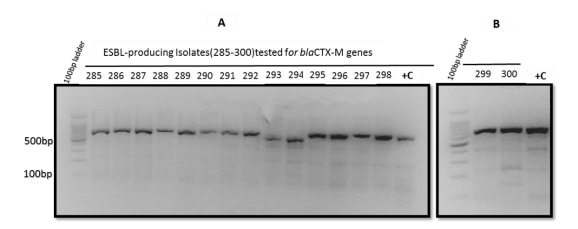
Gel analysis of blaCTX-M amplicon on 2% agarose gel. Lane 1:100bp ladder. Lanes 2–15: 14 ESBL samples. Eleven samples had bands with lengths that corresponded to CTXMGp1 (688bp) samples (257,258,259,260,262,263,264,265,266,267,268). Lane 16: positive control.

Figure 4.22 Multiplex PCR assay for the blaCTX-M genes, samples 271-284



Gel analysis of *blaCTX-M* amplicon on 2% agarose gel. Lane 1: 100bp ladder; lanes 2–15: 14 ESBL samples. Fourteen samples had bands with lengths that corresponded to *CTX-MGp1* (688 bp) samples (271-284). Lane 16: positive control.

Figure 4.23 Multiplex PCR assay for the blaCTX-M genes, samples 285-300



(A) Gel analysis of *bla*CTX-M amplicon on 2% agarose gel. Lane 1: the 100bp ladder. Lanes 2–15: 14 ESBL samples. Fourteen samples had bands with lengths that corresponded to 2 groups of *CTX-M* genes: *CTX-MGp1* (688bp) (samples 285, 286, 287, 288, 289, 290, 291, 292, 295, 296, 297 and 298) and *CTX-MGp9* (561bp) (samples 293 and 294). Lane 16: positive control. (B) Gel analysis of *blaCTX-M* amplicon on 2% agarose gel. Lane 1: the 100bp ladder. Lanes 2–3: 2 ESBL samples. Two samples had bands with lengths that corresponded to *CTX-MGp1* (688bp) samples (299 and 300). Lane 16: positive control.

The screening of ESBL-producing isolates from Wrexham Maelor Hospital (201–300) for the CTX-M groups using multiplex PCR followed by gel analysis shows, based on the amplicon size, that the predominant CTX-M group is group 1 (Table 4.8).

Table 4.8 PCR amplicon sizes of the blaCTX-M genes, samples 201-300

CTX-M group	Amplicon size (bp)	Sample No
CTX-M group 1	688	201, 204, 205, 206, 207, 208, 209, 210, 211, 212, 215, 216, 217, 218, 219, 220, 221, 222, 223, 225, 219, 227, 229, 231, 236, 237, 238, 239, 240, 241, 242, 244,
		245, 247, 248, 249, 250, 251, 253, 255, 256, 258, 259, 260, 263, 264, 265, 266, 267, 268, 257, 262, 271, 272, 273, 274, 275, 276, 276, 277, 278, 279, 280, 281, 285, 286, 287, 288, 289, 290, 291, 292, 295, 296, 297, 298, 299, 300.
CTX-M group 2	404	202, 203, 246.
CTX-M group 9	561	293 and 294.
CTX-M group 8/25	326	

4.2.3.3 Sequencing analysis of multiplex PCR products (201-300)

Forty-five (50%) of samples that displayed bands with sizes that corresponded with 1 of the CTX-M groups were sequenced and then compared with DNA sequences of known b-lactamase genes using the BLAST programme Table 4.9 shows sequencing analysis of positive samples from 201-300.

Table 4.9 Sequencing analysis of positive samples from 201-300

Sample No		Accession	CTX-M group			DNA	
	Blast output	number	Gp1	Gp2	Gp9	Gp8/25	sequence
201	CTX-M-15 gene	MH900522	✓				Appendix III
202	CTX-M-2 gene	LC229068		✓			Appendix III
205	CTX-M-15 gene	MF346615	√				Appendix III
207	CTX-M-108 gene	JF274245	√				Appendix III
209	Escherichia coli strain 3385 chromosome, complete genome	CP029420					Appendix III
211	CTX-M-15 gene	MF346615	✓				Appendix III
215	CTX-M-172 gene	NG_048957.1	√				Appendix III
217	CTX-M-15 gene	CP040398	√				Appendix III
219	CTX-M-15 gene	MK405591	√				Appendix III
223	CTX-M-15 gene	LT628518	√				Appendix III
226	CTX-M-14(like)	CP032888			✓		Appendix III
228	CTX-M-27 gene	MfH900525			√		Appendix III
229	CTX-M-15 gene	LT628518.1	√				Appendix III
231	CTX-M-15 gene	MK113959	√				Appendix III
238	CTX-M-15 gene	LT628520	√				Appendix III
241	CTX-M-32 gene	MH900527	√				Appendix III
242	CTX-M-15 gene	LT628518	√				Appendix III
244	CTX-M-15 gene	MK234851.1	√				Appendix III
246	CTX-M-14(like)	CP032888			✓		Appendix III
248	CTX-M-15 gene	MH900522	√				Appendix III
250	CTX-M-15 gene	MK405591	√				Appendix III
252	CTX-M-2 gene	MH900526.1		✓			Appendix III
254	Escherichia coli strain 4928STDY7071340 genome assembly.	LR607331.1					Appendix III
257	CTX-M-14(like)	CP032888			✓		Appendix III
259	CTX-M-15 gene	LT628516	✓				Appendix III
262	CTX-M-15 gene	MK113960	✓				Appendix III

264	CTX-M-15 gene	MK405591	√		Appendix III
266	CTX-M-15 gene	MF977517.1	√		Appendix III
268	CTX-M-15 gene	MK405591.1	√		Appendix III
271	CTX-M-15 gene	KY640534	√		Appendix III
272(lower band)	Escherichia coli strain 4928STDY7071340 genome assembly	LR607331.1			Appendix III
275	CTX-M-90 gene	MF418175.1		✓	Appendix III
277	<i>CTX-M-163</i> gene	NG_048948.1	√		Appendix III
279	CTX-M-15 gene	KR338941	√		Appendix III
281	CTX-M-15 gene	MK113960	√		Appendix III
283	CTX-M-14(like)	CP032888		✓	Appendix III
286	CTX-M-15 gene	MH891569	√		Appendix III
288	CTX-M-15 gene	KY640534	√		Appendix III
290	CTX-M-15 gene	MK113957	√		Appendix III
292	CTX-M-15 gene	KR338941	√		Appendix III
293	CTX-M-9 gene	CP031724.1		✓	Appendix III
294	CTX-M-14(like)	CP032888		✓	Appendix III
296	CTX-M-15 gene	KY640536.1	✓		Appendix III
298	CTX-M-1 gene	MH037035.1	√		Appendix III
300	CTX-M-15 gene	MH900522.1	✓		Appendix III

The BLAST output of the sequenced YG Hospital sample shows that the *CTX-M-15* gene is the most prevalent type of *CTX-M*; 27(60%) followed by *CTX-M-14* (like); 5 (11%), *CTX-M-2*: 2 (4%), *CTX-M-108*: 1 (2.2%), *CTX-M-172*: 1 (2.2%), *CTX-M-27*: 1 (2.2%), *CTX-M-32*: 1 (2.2%), *CTX-M-90*: 1 (2.2%), *CTX-M-163*: 1 (2.2%), *CTX-M-9*: 1 (2.2%).

Sequencing of $\approx 50\%$ of ESBL-producing isolates, from all three hospitals, that had bands on gel analysis correspond to one of the CTX-M genes showed that 55.73% were CTX-M-15 producers. Table 4.10 summarises the percentage of each detected CTX-M type among the sequenced samples.

Table 4.10 Table 4.10 CTX-M genes identified at the three hospital sites

Blast out	CTX-M group	Number (%)
CTX-M-15	Gp1	77 (55.7%)
CTX-M-14 (like)	Gp9	5 (3.62%)
CTX-M-9	Gp9	5 (3.62%)
CTX-M-27	Gp9	4 (2.8%)
CTX-M-59	Gp2	4 (2.8%)
CTX-M-2	Gp2	4 (2.8%)
CTX-M-108	Gp9	3 (2.1%)
CTX-M-32	Gp1	3 (2.1%)
CTX-M-1	Gp1	3 (2.1%)
CTX-M-172	Gp1	3 (2.1%)
CTX-M-55	Gp1	3 (2.1%)
CTX-M-3	Gp1	2 (1.4%)
CTX-M-66	Gp1	2 (1.4%)
CTX-M-163	Gp1	2 (1.4%)
CTX-M-160	Gp1	1 (0.72%)
CTX-M-225	Gp1	1 (0.72%)
CTX-M-203	Gp1	1 (0.72%)
CTX-M-65	Gp1	1 (0.72%)
CTX-M-51	Gp9	1 (0.72%)
CTX-M-90	Gp9	1 (0.72%)
CTX-M family		1 (0.72%)
Others e.g,Escherichia coli strain MNCRE44, complete genome		11 (7.9%)

4.3 Discussion

4.3.1 Primers competition in Multiplex PCR

In multiplex PCR, more than one target sequence can be amplified by including multiple pairs of primers in the reaction, which increases the detection of resistance genes in a single test, and results in a considerable saving of time and effort (Elnifro et al., 200). Limitations of multiplex PCR include a lack of optimization and false negatives. The latter is caused by primer competition for reagents, which could favour stronger over weaker binding primers, and increases the chance of obtaining spurious amplification products, due primarily to the formation of primer dimers (Brownie, 1997). However, this competition for reagents is not so crucial because it mainly reduces the yield, and in this study the problem could be overcome by using a published primer in which the design parameters had been selected to account for the homology of primers with their target nucleic acid sequences, their length, the GC content, and their concentration.

4.3.2 CTX-M-15 is the dominant CTX-M-type ESBL in North Wales

CTX-Ms have become the most rapidly growing type of ESBL since they were first identified in Germany, France and South America in 2000. Many studies have been established to draw the global picture of CTX-M variants, and it has been concluded that blaCTX-M-15 has increased over time in most countries and is considered the dominant E.coli resistant gene in most regions (Edward et al., 2017). Interestingly, the exception from this rule are regions like South America, where blaCTX-M-2 is the most detectable CTX-M gene, and Spain, South-East Asia, China, Japan and South Korea, where the group 9 variants are the most prevalent CTX-Ms (Woerther et al., 2013).

The United Kingdom is classified as a region with a high prevalence of *blaCTX-M-15*. However, the previous studies by Mushtaq et al. (2003), who conducted a study at 26 hospitals in the UK and Ireland where the researchers collected up to 200 consecutive, clinically significant isolates from separate inpatients during a multi-centre survey of inpatient isolates concluded that among 122 cephalosporin-resistant isolates of *Enterobacteriaceae*, there were only 4 *CTX-M-15*-producing isolates that were collected at hospitals in London (2 isolates), Newcastle-upon-Tyne and Belfast. The region of this study, North Wales, was part of Mushtaqs et al' study.

The present study shows that the prevalence of *CTX-M-15* has reached North Wales as among the 138 (50% of samples had bands on gel analysis corresponded with of CTX-M genes)

sequenced ESBL-producing isolates collected from 3 main referral hospitals in North Wales, 77 (55.73%) have been detected as *CTX-M-15* producers. This is considered a remarkable increase in the UK epidemiology of *CTX-M-15* and presents huge challenges to healthcare as it restricts options to treat infections caused by CTX-M-producing bacteria.

In terms of antibiotic resistance, it has been reported that *CTX-M-15* possesses a high activity against Ceftazidime (Malik et al., 2018). The YG Hospital data (Table 3.2) indicates that the isolates that were identified genotypically as *CTX-M-15* producers conferred resistance to other antibiotic profiles, such as amoxicillin and ciprofloxacin, suggesting the presence of other mechanisms in addition to *CTX-M-15* or novel mutations in the *CTX-M-15* gene that may develop as a result of misuse of antibiotics due to the limitations of ESBL phenotypic dictions in clinical laboratories (Rahman et al., 2018). An alternative explanation may be provided by the rural environment of the North Wales hospitals which serve a large farming community. Given the presence of antibiotic resistant *Enterobacteriaceae* in food producing animals like lambs (Gozi kS et al., 2019), the resistance conferring plasmids may be transmitted to humans.

It has been suggested (e.g. Carattoli, 2013; Orhue et al., 2017) that the high global spread of *CTX-M 15* is promoted by its location on the conjugative plasmid *IncF*, which facilitates the horizontal transfer of the antibiotic resistance plasmid by the conjugation in *Enterobacteriaceae*. This singles *CTX-M-15* out as other *blaCTX-M* genes like *blaCTX-M-14*, the second dominant CTX-M type is carried on a variety of plasmid types, including IncF, like the IncK plasmid that is prevalent in the Far East, (Ang et al., 2015). Moreover, the main host of the IncF plasmid is *Enterobacteriaceae* (Fortini et al., 2015), which explains the high percentage of *E.coli* that carries the *CTX-M-15* gene in compression with coliforms among the isolates of this study.

4.3.3 The novel beta-Lactamase, CTX-M-14-LIKE

Sequencing of the YG Hospital samples revealed that, in addition to *CTX-M-15* as the dominant CTX-M type, some isolates ;226 (*E. coli*), 246 (*E. coli*), 257 (coliform), 283 (*E. coli*) and 296 (*E. coli*), which represent 11% of the sequenced samples, revealed a new β-lactamase gene that is closely related to *CTX-M-14* at the protein level as it substitutes only 3 amino acids when compared to the known CTX-M-14 protein. The three amino acids are T55A, A273P and R277C (Figures 4.24). Because of its close similarity to CTX-M-14, the novel gene was named *CTX-M-14-Like*.

Figure 4.24 Sequence alignment of CTX-M14&CTX-M14 LIKE proteins

Query	1	mvtkrvqrmmfaaaaciplllgsaplyaqtsavqqklaalekssggrlgvalid <mark>A</mark> adntq	60
		MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALID ADNTQ	
Sbjct	1	${\tt MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALIDTADNTQ}$	60
Query	61	$\verb VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM $	120
		VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	
Sbjct	61	VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	120
Query	121	TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	180
		TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	
Sbjct	121	TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	180
Query	181	RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	240
		RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	
Sbjct	181	RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	240
Query	241	gdygttndiaviwpqgraplvlvtyftqpqqn <mark>P</mark> esr <mark>C</mark> dvlasaariiaegl 291	
		GDYGTTNDIAVIWPOGRAPLVLVTYFTOPOON ESR DVLASAARIIAEGL	
Sbjct	241	GDYGTTNDIAVIWPOGRAPLVLVTYFTOPOONAESRRDVLASAARIIAEGL 291	
عات ر بدد	Z 4 I	ADIAIIMDIYAIMI ÄQVYEDADAIILI TÄLÄÄÄNYESVVDADYOVYVIIVERI 531	

The sequence alignment of CTX-M-14 (query sequence 1) and CTX-M-14 LIKE (query sequence 2) proteins using the Clustal Omega tool, as indicated by the 3 highlighted differences between CTX-M-14 and CTX-M-14 LIKE, is T55A, A273P and R277C.

BLAST output CTX-M-14 [Enterobacteriaceae], Sequence ID: ref|WP_001617865.1|.

The two CTX-M genes, *CTX-M-15* as the dominant CTX-M in North Wales and the novel *CTX-M-14-LIKE* β-lactamase gene were then further analysed upon cloning into an *E.coli* expression plasmid (pASK-IBA2), which allowed for the purification of a C-terminally streptagged recombinant protein. The activity of the purified, recombinant enzymes was then tested with the most commonly used antibiotic for UTI patients in the clinical laboratories of North Wales hospitals. This work is detailed in Chapter 5 and 6 respectively.

Chapter 5 : OVEREXPRESSION AND PURIFICATION OF RECOMBINANT CTX-M ENZYMES

5.1. Introduction

As the study of a protein's properties requires a large number of proteins to study (and these proteins usually cannot be produced by native hosts), protein overexpression was developed to increase protein production by an appropriate host and to facilitate the purification of proteins (Tripathi & Nagesh, 2016). Three main elements are required for protein overexpression: a gene, a vector (an expression plasmid) carrying the gene of interest and an expression host, which maximises the amount and quality of the protein produced by the vector—gene combination (Goh et al., 2017).

An expression plasmid is generally engineered to carry regulatory sequences, that is, the enhancer and promoter regions responsible for the transcription of the inserted gene in the expression plasmid (Singha et al., 2017). An appropriate vector should be selected precisely to ensure the efficient production of proteins. Thus, the expression plasmid chosen in this study to obtain an abundant amount of proteins from *CTX-M* genes was the *pASK-IBA2C* plasmid. *pASK-IBA2C* is an expression plasmid that is under the transcriptional control of the tetracycline promoter/operator and that is characterised by its ability to export the expressed recombinant protein into the periplasm of *Escherichia coli* to facilitate protein harvesting and purification (Schmidt & Skerra, 2015). Chemically competent *E. coli* cells, which is considered one of the most popular hosts for protein expression, was used as an organism for the overexpression of recombinant CTX-M proteins.

The purification of the produced recombinant protein can be achieved using a tag protein affinity system, in which the protein is genetically tagged with a small amino acid peptide sequence. This is a predominantly viral protein sequence that can be attached to the N-terminus or C-terminus of the recombinant proteins based on the needs of the experiment and the properties of the protein of interest (Kosobokova et al., 2016). In the *pASK-IBA2C* plasmid, the included tag sequence is the Strep-tag, which constitutes the nine amino acid peptide sequence Ala-Trp-Arg-His-Pro-Gln-Phe-Gly-Gly. This small amino acid sequence was engineered to reversibly bind to the protein Streptavidin and is in this sense distinct from other commonly used affinity peptides like HA or Myc that are peptides from natural proteins (Chromatogr, 1994).

The reversible binding of the Strep-tag sequence to a genetically modified Streptavidin protein known as Strep-Tactin®, allows for a one-step protein purification (Yeliseev et al., 2017). Transfer of the recombinant protein is facilitated by the OmpA leader peptide that is later removed during the transport of the recombinant protein by a membrane-bound protease inside the periplasmic space (Inouye& Halegoua, 1980). The OmpA leader peptide is encoded by the *pASK-IBA2C* plasmid and sits at the N-terminus of the recombinant protein. Figure 5.1 shows the Strep-Tactin®XT purification system used to purify the recombinant Strep-tagged CTX-M proteins.

Recombinant CTX-M protein Strep-tag

Figure 5.1 Protein purification using Strep-Tactin®

Purification of Strep-tagged CTX-M proteins using the Strep-Tactin® purification system, which relies on binding the Strep-Tag II® and Strep-Tactin® included in prepacked chromatography columns (Modified from IBA Data Sheet, 2017).

The objectives of this experiment were to obtain an abundant amount of pure CTX-M-14, CTX-M-14-LIKE and CTX-M-15 proteins to measure the enzymatic activity of these beta-lactamases against commonly used antibiotics for UTI patients and to compare the catalytic

activity of the novel B-lactamase, CTX-M-14-LIKE, with the known CTX-M-14 enzyme. CTX-M protein expression was completed in four steps: the full-length gene encoding *CTX-M* was inserted into the plasmid *pASK-IBA2C*, the transformation of *pASK-IBA2C* into carrying *CTX-M* in *E. coli*, production of the protein of interest by the expression host followed by lysing and purification from the periplasmic space and, finally, the verification of the protein expression level using the Western blot technique.

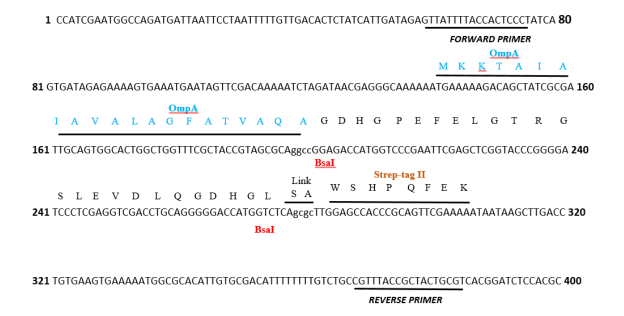
5.2. Results

5.2.1. Cloning of a mutant *CTX-M* gene into a *pASK-IBA2C* plasmid expressing *E. coli*

The two *CTX-M* genes, the novel *CTX-M-14-LIKE* and *CTX-M-15*, were cloned into a *pASK-IBA2C* plasmid expressing *E. coli* to allow for the purification of a Strep-tagged recombinant protein. This vector carries a *Tet* promoter, which was developed to produce C-terminal fusion proteins and the Strep-tag II®. The produced recombinant protein is transported into the periplasm of *E. coli* via the vector-encoded OmpA signal sequence.

The full-length *CTX-M* genes were amplified from the corresponding samples using forward and reverse primers (Section 2.2.9, Table 2.3). Both primers contained BsaI restriction sites at multiple cloning sites in the plasmid (Figure 5.2).

Figure 5.2 Multiple Cloning Site of pASK-IBA2C

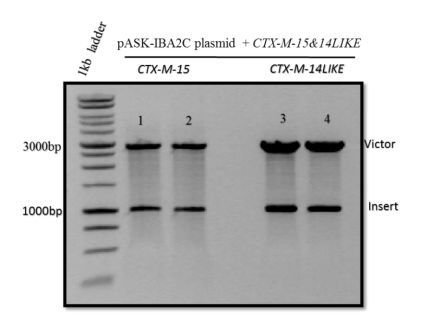


Features of multiple cloning sites of *pASK-IBA2C*, which are responsible for transferring the protein produced in the *E. coli* cytoplasm to the periplasmic space, Strep-tag II® and the restriction sites of BsaI (Modified from IBA Data Sheet, 2010).

The ligation of the *pASK-IBA2C* plasmid vector carrying the *CTX-M-14-LIKE* and *CTX-M-15* genes was performed using the T4 DNA ligase enzyme. To replicate the plasmid carrying the gene of interest, the plasmid was transformed into chemically competent *E. coli* using the heat shock method (Section 2.2.11).

The cells transformed with the plasmid were plated on 25 μ g/mL of LB agar with chloramphenicol (Section 2.2.12). After overnight incubation, the growth colonies were incubated overnight in LB/chloramphenicol at 37 °C, and the recombinant plasmid was agitated. The transformed *E. coli* was purified using an ISOLATE II Plasmid Kit. The purified recombinant plasmid was then digested by BsaI enzyme (Section 2.2.13). In order to verify the insertion, agarose gel was used to analyse the restricted recombinant plasmid, where two bands can be seen in the lane of the digestion reaction of the *pASK-IBA2C* plasmid vector and *CTX-M* genes; the vector had a length of 3,000 bp and an insert of 1,000 bp (Figure 5.3).

Figure 5.3 Gel analysis of pASK-IBA2C, CTX-M-14-LIKE and CTX-M-15



Plasmid mapping analysis on 0.8% agarose gel. Lane 1: 6 μ L of the DNA ladder (1 kp of the ladder). Lanes 2 and 3: 10 μ L of the digestion reaction of the *pASK-IBA2C* plasmid vector and *CTX-M-15* (plasmids 1 and 2). Lanes 6 and 7: 10 μ L of the digestion reaction of the *pASK-IBA2C* plasmid vector and *CTX-M-14-LIKE* (plasmids 3 and 4). The vector had a length of 3,000 bp and an insert of 1,000 bp.

5.2.2 Sequencing analysis of a cloned pASK-IBA2C plasmid with CTX-M-14-LIKE and CTX-M-15

The purified pASK-IBA2C plasmid carrying CTX-M-14-LIKE and CTX-M-15 was sequenced using pASK-IBA2C forward primer in order to verify the insertion of the CTX-M genes into the expression plasmid. Sequencing analysis was performed on one plasmid from each CTX-M gene (plasmids 1 and 3). The sequenced samples were compared with the DNA sequences of known β -lactamase genes using the BLAST programme. Table 5.1 shows the BLAST output of the sequenced samples.

Table 5.1 Sequencing analysis of the pASK-IBA2C plasmid carrying CTX-M-14-LIKE and CTX-M-15

Plasmid Number	BLAST Output	Accession
		Number
	Escherichia coli plasmid pryc110.1	
	insertion sequence ISCR1 putative	
1	recombinase gene, partial cds and beta-	GQ892051.1
	lactamase CTX-M-14 (blaCTX-M-14b)	
	gene, complete cds	
	Klebsiella pneumoniae strain KPTR7239-	
	17 insertion sequence ISEcp1, partial	
3	sequence and beta-lactamase CTX-M-15	MH544748.1
	(blaCTX-M) gene (blaCTX-M-15) allele,	
	complete cds	

5.2.3 Translation and sequence alignments of the recombinant CTX-Ms

The plasmids that shared a high sequence identity with the *CTX-M-14-LIKE* and *CTX-M-15* genes were translated to protein sequences using ExPASy (Figures; 5.4(A) &5.5(B)), which allows for the translation of a nucleotide sequence to a protein sequence. After, the sequences were aligned using Clustal Omega, which arranges the protein sequences to identify regions of similarity that may be a consequence of functional, structural or evolutionary relationships between the sequences. Figure 5.5(B) shows the three substitutions between *CTX-M-14-LIKE* and *CTX-M-14*: T55A, A273P and R277

Figure 5.4 Protein Sequence of plasmid (1); pASK-IBA2C plasmid carrying CTX-M-15

(A)

MKKTAIAIAVALAGFATVAQAMVKKSLRQFTLMATATVTLLLGSVPLYAQTADVQQKLAELER QSGGRLGVALINTADNSQILYRADERFAMCSTSKVMAAAAVLKKSESEPNLLNQRVEIKKSDLV NYNPIAEKHVNGTMSLAELSAAALQYSDNVAMNKLIAHVGGPASVTAFARQLGDETFRLDRTEP TLNTAIPGDPRDTTSPRAMAQTLRNLTLGKALGDSQRAQLVTWMKGNTTGAASIQAGLPASWV VGDKTGSGGYGTTNDIAVIWPKDRAPLILVTYFTQPQPKAESRRDVLASAAKS

OmpA Leader (incomplete sequence)CTX-M-15Strep Tag

(B)

Query	1	MVKKSLRQFTLMATATVTLLLGSVPLYAQTADVQQKLAELERQSGGRLGVALINTADNSQ	60
		MVKKSLRQFTLMATATVTLLLGSVPLYAQTADVQQKLAELERQSGGRLGVALINTADNSQ	
Sbjct	1	MVKKSLRQFTLMATATVTLLLGSVPLYAQTADVQQKLAELERQSGGRLGVALINTADNSQ	60
Query	61	ILYRADERFAMCSTSKVMAAAAVLKKSESEPNLLNQRVEIKKSDLVNYNPIAEKHVNGTM	120
		ILYRADERFAMCSTSKVMAAAAVLKKSESEPNLLNQRVEIKKSDLVNYNPIAEKHVNGTM	
Sbjct	61	ILYRADERFAMCSTSKVMAAAAVLKKSESEPNLLNQRVEIKKSDLVNYNPIAEKHVNGTM	120
Query	121	SLAELSAAALQYSDNVAMNKLIAHVGGPASVTAFARQLGDETFRLDRTEPTLNTAIPGDP	180
		SLAELSAAALQYSDNVAMNKLIAHVGGPASVTAFARQLGDETFRLDRTEPTLNTAIPGDP	
Sbjct	121	SLAELSAAALQYSDNVAMNKLIAHVGGPASVTAFARQLGDETFRLDRTEPTLNTAIPGDP	180
Query	181	RDTTSPRAMAQTLRNLTLGKALGDSQRAQLVTWMKGNTTGAASIQAGLPASWVVGDKTGS	240
		RDTTSPRAMAQTLRNLTLGKALGDSQRAQLVTWMKGNTTGAASIQAGLPASWVVGDKTGS	
Sbjct	181	RDTTSPRAMAQTLRNLTLGKALGDSQRAQLVTWMKGNTTGAASIQAGLPASWVVGDKTGS	240
Query	241	GGYGTTNDIAVIWPKDRAPLILVTYFTQPQPKAESRRDVLASAAK 285	
		GGYGTTNDIAVIWPKDRAPLILVTYFTQPQPKAESRRDVLASAAK	
Sbjct	241	GGYGTTNDIAVIWPKDRAPLILVTYFTQPQPKAESRRDVLASAAK 285	

(A)Forward protein Translation of *CTX-M-15* gene using Expasy tool .The three elements needed for protein overexpression (OmpA lead, inserted gene and strep tag are highlighted (B) Protein alignments created Using Clustal Omega tool. Protein Blast output: class A extended-spectrum beta-lactamase *CTX-M-15*.

Sequence ID: WP_000239590.1

Figure 5.5 Protein Sequence of plasmid (3); pASK-IBA2C plasmid carrying CTX-M-14LIKE

(A)

ASALAGFATVAQAMVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALIDAA DNTQVLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTMTLAELS AAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDPRDTTTPRAMAQTLRQLT LGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGSGDYGTTNDIAVIWPQGRAPLVLVTYFT QPQQNPESRCDVLASAARIIAEGLSAWSHPQFEK

OmpA Leader (incomplete sequence) CTX-M-14 like Strep Tag

(B)

Query	1	${\tt MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALID} {\color{red} {\bf A}} {\tt ADNTQ}$	60
		MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALID ADNTQ	
Sbjct	1	MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALIDTADNTQ	60
Query	61	VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	120
2		VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	
Sbjct	61	VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	120
•	101		100
Query	121	TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP TLAELSAAALOYSDNTAMNKLIAOLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	180
Sbjct	121	TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	180
Query	181	RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	240
- 1		RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	
Sbjct	181	RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	240
Query	241	GDYGTTNDIAVIWPQGRAPLVLVTYFTQPQQN P ESR C DVLASAARIIAEGL 291	
		GDYGTTNDIAVIWPQGRAPLVLVTYFTQPQQN ESR DVLASAARIIAEGL	
Sbjct	241	GDYGTTNDIAVIWPQGRAPLVLVTYFTQPQQNAESRRDVLASAARIIAEGL 291	

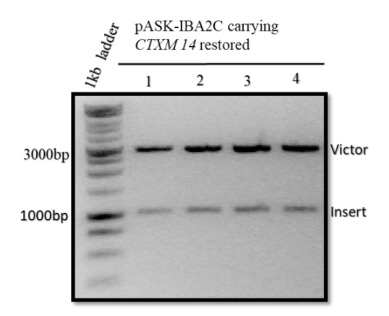
(A) Forward protein Translation of *CTX-M-14LIKE* gene using Expasy tool. The three elements needed for protein overexpression (OmpA lead, inserted gene and strep tag are highlighted. (B) Protein alignments created Using Clustal Omega tool. Protein Blast output: class A extended-spectrum beta-lactamase *CTX-M-14*. Sequence ID: WP_001617865.1.As indicated the three highlighted differences between CTXM-14 & CTXM-14 – Like are: T55A, A273P and R277C.

5.2.4 Using a Q5® Site-Directed Mutagenesis Kit for the reversion of the three *CTX-M-14-LIKE* changes

In order to compare CTX-M-14 and CTX-M-14-LIKE enzymes in terms of their enzymatic activity against commonly used antibiotics for UTI patients, T55A, A273P and R277C were mutated back to the CTX-M-14 wild-type sequence using the Q5® Site-Directed Mutagenesis Kit (Section 2.2.18). After the triple mutagenesis of the CTX-M-14-LIKE gene, the pASK-IBA2C plasmid carrying the restored CTX-M-14 gene was transformed into E. coli, plated on LB agar with chloramphenicol, incubated overnight in LB broth, purified using an ISOLATE II Plasmid Kit and restricted using BsaI before sequencing.

Figure 5.6 shows the gel analysis of the restricted and purified *pASK-IBA2C* plasmid carrying the restored *CTX-M-14* in which two bands can be seen in the lane of the digestion reaction: the vector at length of 3,000 bp and an insert of 1,000 bp.

Figure 5.6 Gel analysis of pASK-IBA2C and the triple reverted CTX-M-14



Plasmid mapping analysis on 0.8% agarose gel. Lane 1: 6 μ L of the DNA ladder (1 kp of the ladder). Lanes 2, 3 and 4: 10 μ L of the digestion reaction of the pASK-IBA2C plasmid vector and *CTX-M-14* restored (T55A, A273P and R277C). The vector had a length of 3,000 bp and an insert of 1,000 bp.

The *pASK-IBA2C* plasmid carrying triple reversion of *CTX-M-14-LIKE* (A55T, P273A and C277R) was sequenced using the *pASK-IBA2C* forward primer in order to verify that *CTX-M-14-LIKE* was mutated back to *CTX-M14*. The sequencing analysis was performed on plasmids 1 and 2. The sequenced samples were compared with the DNA sequences of known β -lactamase genes using the BLAST programme and then translated and aligned using ExPASy and Clustal Omega, respectively (Figure 5.7).

Figure 5.7 Protein Sequence of plasmid (1); pASK-IBA2C plasmid carrying the triple reversion of CTXM 14-LIKE

(A)

MKKTAIAIAVALAGFATVAQAMVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEK SSGGRLGVALIDTADNTQVLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLV NYNPIAEKHVNGTMTLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEP TLNTAIPGDPRDTTTPRAMAQLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVG DKTGSGDYGTTNDIAVIWPQGRAPLVLVTYFTQPQQNAESRRDVLASAARIIAEGLSAWSHPQFE K

OmpA Leader (incomplete sequence) CTX-M-14 likeStrep Tag

(B)

Query	1	${\tt MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALID} {\bf T} {\tt ADNTQ}$	60
		${\tt MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALIDTADNTQ}$	
Sbjct	1	${\tt MVTKRVQRMMFAAAACIPLLLGSAPLYAQTSAVQQKLAALEKSSGGRLGVALIDTADNTQ}$	60
Query	61	VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	120
		VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	
Sbjct	61	VLYRGDERFPMCSTSKVMAAAAVLKQSETQKQLLNQPVEIKPADLVNYNPIAEKHVNGTM	120
Query	121	TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	180
		TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	
Sbjct	121	TLAELSAAALQYSDNTAMNKLIAQLGGPGGVTAFARAIGDETFRLDRTEPTLNTAIPGDP	180
Query	181	RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	240
		RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	
Sbjct	181	RDTTTPRAMAQTLRQLTLGHALGETQRAQLVTWLKGNTTGAASIRAGLPTSWTVGDKTGS	240
Query	241	GDYGTTNDIAVIWPQGRAPLVLVTYFTQPQQN A ESR R DVLASAARIIAEGL 291	
		GDYGTTNDIAVIWPQGRAPLVLVTYFTQPQQNAESRRDVLASAARIIAEGL	
Sbjct	241	GDYGTTNDIAVIWPQGRAPLVLVTYFTQPQQNAESRRDVLASAARIIAEGL 291	

(A)Forward protein Translation of *triple reversion of CTXM 14-LIKe* gene using Expasy tool. The three elements needed for protein overexpression (OmpA lead, inserted gene and strep tag are highlighted (B) Protein alignments created Using Clustal Omega tool. Protein Blast output: class A extended-spectrum beta-lactamase *CTX-M-14* [*Enterobacterales*] Sequence ID: WP_001617865.1

Length: 291 .As indicated the three highlighted changes of *CTX-M-14LIKE*: T55A, A273P and R277C were mutated back to *CTX-M-14*.

5.2.5 Producing Strep-tag fusion proteins for CTX-M-14, CTX-M-14-LIKE and CTX-M-15

The production of the Strep-Tag fusion proteins for CTX-M-14, CTX-M-14-LIKE and CTX-M-15 was performed in three main steps: growth of host strains harbouring the expression plasmid using the three CTX-Ms, gene expression of the Strep-Tag fusion proteins induced by the addition of 10 μL of anhydrotetracycline solution (aTe;2 mg/mL in dimethylformamide, DMF) logarithmically growing cells (Section 2.2.14.2) and protein extraction (Sections 2.2.14.1 and 2.2.14.2). Upon induction, the protein produced in the cytoplasm of the host cell was exported to the periplasmic space by the OmpA leader sequence (Figure 5.8), which was removed while the protein was transported through the inner membrane (Plfickthun & Skerra, 1989).

Periplasmic space

Cytosol

ompA seq

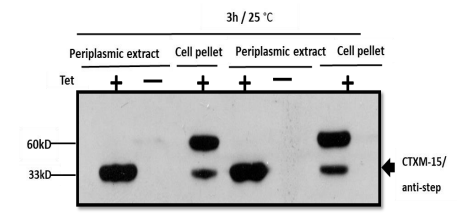
Unfolded preprotein(CTX-M)

Figure 5.8 Periplasmic Expression of recombinant CTX-M enzymes

Periplasmic secretion strategy for production of recombinant CTX-M proteins in *E.coli*. The exportation of the produced unfolded recombinant protein to the periplasmic space is led by OmpA leader sequence.

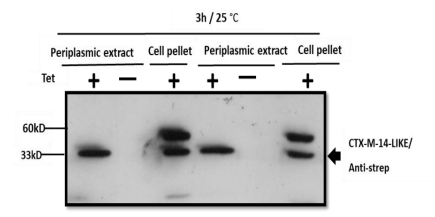
The expression level of the extracted protein was verified using the Western blot technique (Section 2.2.15). Figures 5.9, 5.10, and 5.11 show the Western blot analysis of the extracted periplasmic CTX-M proteins; CTX-M-15, CTX-M-14-LIKE and CTX-M-14, the supernatant of centrifuged, which had bands with sizes corresponding with the expected sizes of the CTX-M proteins (33 KD), induced and uninduced strains, as well as the remaining cell pellet. The cell pellet consisted of the recombinant proteins left in the cytoplasm to check the efficiency of the export from the cytoplasm to the periplasmic space. Retention in the cytoplasm is frequently caused by the formation of inclusion bodies made up of mis-folded, recombinant protein.

Figure 5.9 Western blot analysis of Induced strep-tagged recombinant CTX-M-15



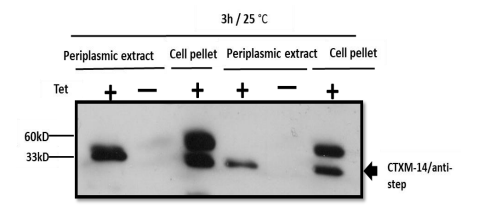
Western blot analysis after 3h induction at 25°C with anhydrotetracycline of strep-tagged recombinant CTX-M-15.(+) = induced strains with anhydrotetracycline; (-) = Uninduced strains (negative control). Tested induced strains cell pellet (i.e., Cytoplasmic produced recombinant protein), show the bands' sizes corresponding with the expected sizes of the CTX-M proteins (33 KD), which indicates that some of protein hasn't been exported despite the high yielded of recombinant CTX-M extracted from the periplasmic space, in addition to a band at approximately 60 KD, could be dimeric 60KD Which is insoluble and therefore in the pellet.

Figure 5.10 Western blot analysis of Induced strep-tagged recombinant CTX-M-14LIKE



Western blot analysis after 3h induction at 25°C with anhydrotetracycline of strep-tagged recombinan.t *CTX-M-14LIKE*. (+) = induced strains with anhydrotetracycline; (-) = Uninduced strains (negative control). Tested induced strains cell pellet (i.e., Cytoplasmic produced recombinant protein), show the bands' sizes corresponding with the expected sizes of the CTX-M proteins (33 KD), which indicates that some of protein hasn't been exported despite the high yielded of recombinant CTX-M extracted from the periplasmic space, in addition to a band at approximately 60 KD, could be dimeric 60KD Which is insoluble and therefore in the pellet.

Figure 5.11 Western blot analysis of Induced strep-tagged recombinant CTX-M-14



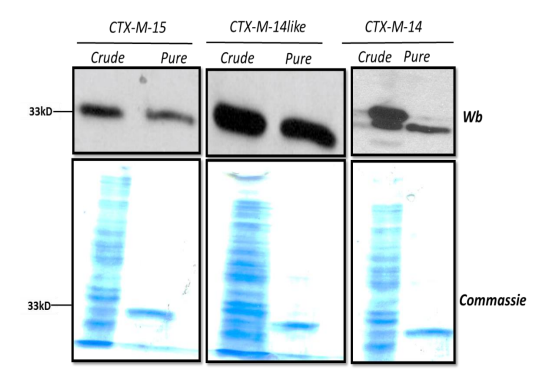
Western blot analysis after 3h induction at 25°C with anhydrotetracycline of streptagged recombinan.t *CTX-M-14*. (+) = induced strains with anhydrotetracycline; (-) = Uninduced strains (negative control). Tested induced strains cell pellet (i.e., Cytoplasmic produced recombinant protein), show the bands' sizes corresponding with the expected sizes of the CTX-M proteins (33 KD), which indicates that some of protein hasn't been exported despite the high yielded of recombinant CTX-M extracted from the periplasmic space, in addition to a band at approximately 60 KD, could be dimeric 60KD which is insoluble and therefore in the pellet.

5.2.6 Protein purification using Strep-Tactin®XT followed by protein concentration using VIVASPIN® 500

The recombinant protein was C-terminally tagged with the short peptide Strep-Tag II®, and has a high selectivity to Strep-Tactin®. The tagged recombinant CTX-M proteins with a Streptag can be purified from the crude extract using the binding affinity between the Strep-Tag II® and Strep-Tactin® included in prepacked chromatography columns, which allows for the gravity-flow purification of the Strep-tagged fusion proteins (Section 2.2.16). The purified CTX-M proteins were then concentrated using VIVASPIN® 500 to increase the concentration of the protein yield.

To check the purity of the CTX-M-14, CTX-M-14-like and CTX-M-15 proteins, the three enzymes were analysed using Coomassie stains, which are a family of dyes commonly used to stain proteins in SDS-PAGE gels. The gels were soaked in dye, and the excess stain was eluted with a solvent. This treatment allows the proteins to be visualised as blue bands on a clear background. Figure 5.14 shows the verification of the three purified CTX-M proteins using Coomassie stains by comparing the crude extract with the eluate of the affinity columns (purified proteins).

Figure 5.12 Checking the purity of the three CTX-Ms using Coomassie stains



Coomassie stain analysis of the purified CTX-M proteins. The expression level of the three produced proteins were verified with Western blot analysis followed by purity verification Using Coomassie stains by comparing the crude extract with the eluate of the affinity columns (purified proteins).

5.2.7 Biochemical characterisation of the CTX-M proteins

One important biochemical characterisation that should be taken into account when studying protein properties is the protein phosphorylation status, which is responsible for a variety of important protein functions, including subcellular localisation, protein degradation and stabilisation and biochemical activities (Manning Getal, 2002). This study sought to determine how dephosphorylating affects the activity of CTX-M enzymes against commonly used antibiotics.

NetPhos predicts the serine, threonine and tyrosine phosphorylation sites of proteins. It showed that CTX-M-14 and CTX-M-15 possess both some potential phosphorylation sites. Figure 5.15 shows the three-dimensional structures of the two CTX-Ms as well as the predicted phosphorylated amino acids.

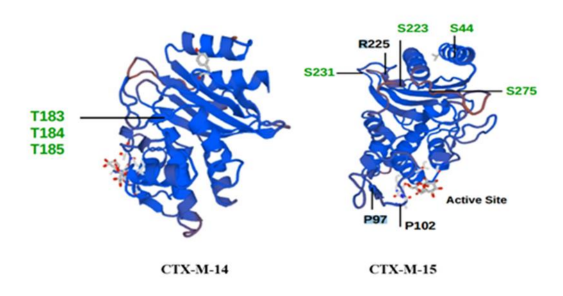


Figure 5.13 Three-dimensional CTX-M-14 and CTX-M-15 structures

Three-dimensional structures of CTX-M-15 and CTX-M-14. The green highlighted amino acids indicate the expected phosphorylation sites using NetPhos. The active site is indicated with a beta-lactam antibiotic. The structure was generated using the SWISS-MODEL (99.62% identity).

The phosphorylation sites of the CTX-M proteins were analysed using two techniques: isoelectric gel electrophoresis and Phos-tag gel analysis.

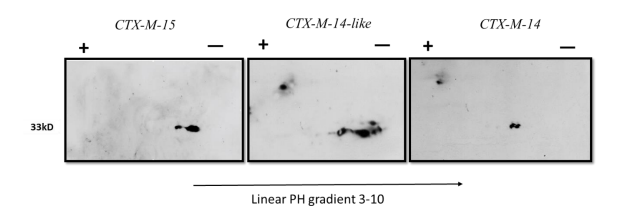
5.2.7.1 Isoelectric-focused analysis of the CTX-M-14, CTX-M-14-LIKE and CTX-M-15 proteins

The three CTX-M proteins were analysed in two-dimensional gels in order to check for posttranslational modifications that either add or remove protein charges (e.g. phosphorylation, acetylation) and to find out how homogenous the purified CTX-M protein samples are. This was achieved by separating the protein first based on its isoelectric point (pI) and then by molecular weight using normal SDS-PAGE (Section 2.2.19.1). Figure 5.16 shows the Western blot analysis of the distributed proteins in the two-dimensional gel. At least two isoforms with the expected MW of 33kD were detected in the three protein extracts. This indicates phosphorylation or another charge changing post-translational modification.

5.2.7.2 Phos-tag gel analysis of the CTX-M-14, CTX-M-14-LIKE and CTX-M-15 proteins

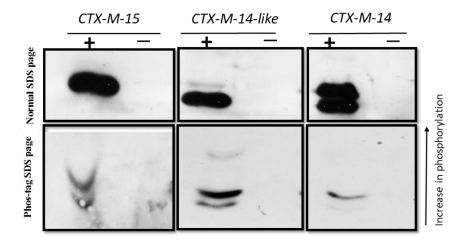
Manganese(II)-Phos-tag SDS-PAGE (Table 2.20) was used to verify whether the observed changes in the isoelectric values are due to phosphorylation as the Phos-tag reagent reacts specifically with phosphorylared protein and slows down their mobility on SDS page (Horinouchi et al,2016) (Section 2.2.19.2). Phos-tag can bind the phosphate groups of the phosphorylated proteins, which delays their electrophoretic mobility so that only phosphorylated proteins can be seen in the Phos-tag gel. As shown in Figure 5-17, at least two bands were observed for all three CTX-M proteins in the presence of Phos-tag. While the band intensity was comparable for CTX-M-15, the slower migration, phosphorylated band was more intense in the case of CTX-M-14 and CTX-M-14-LIKE. Especially, the intensity of the faster migrating band of CTX-M-14-like was very low. CTX-M-14-like displayed a third, strongly retarded band that might indicate a higher phosphorylation level. It was however interesting to observe that even in the absence of the Phostag reagent CTX-M-14-like showed a slower migrating band. The latter finding is not inconsistent with phosphorylation as some phosphorylated proteins migrate slower in normal SDS page (Caspari T, et al, 200).

Figure 5.14 Two-dimensional gel analysis of the CTX-M-14, CTX-M-14-LIKE and CTX-M-15 proteins



Two-dimensional gel analysis showed that the 33 kDa protein exists in two forms (CTX-M-14 and CTX-M-15) and four forms (CTX-M-14-like) with different pI, indicating phosphorylation. CTX-M-14 and CTX-M-14-like exist in heavier molecular weight forms at a higher acidity (90 kDa).

Figure 5.15 Phos-tag gel analysis of the Strep-tagged recombinant CTX-M-15, CTX-M-14-like and CTX-M-14 proteins



Phosphorylation analysis of the CTX-M-14, CTX-M-14-like and CTX-M-15 proteins using manganese (II)-Phos-tag SDS-PAGE. At least one shift band can be seen in the Phos-tag gel analysis of the three CTX-M proteins, which indicates phosphorylation.

5.3 Discussion

5.3.1 Possibility of the pASK-IBA2C plasmid's self-ligation with the BsaI restriction enzyme

The BsaI restriction enzyme has restriction sites located at multiple cloning sites of the plasmid and should be selected based on its ability to generate asymmetric cuts with long sticky ends (GGTCTCN1 on the forward strand and CCAGAGN5 on the reverse strand). It should therefore be unable to re-ligate as BsaI cuts five nucleotides away from the recognition site. In addition, these sequences are different at either site of the two BsaI cuts in the plasmid, meaning that only the cloned fragments should provide colonies after the transformation of the pASK-IBA2C plasmid carrying the *CTX-M* genes in *E. coli* strains (Schmidt & Skerra, 2015).

5.3.2 Periplasmic expression as a suitable strategy for producing CTX-M proteins

There are two strategies for the secretion of proteins from cloned genes in the *pASK-IBA2C* plasmid: cytosolic expression and periplasmic expression. Periplasmic expression was selected in the production of CTX-M proteins due to its advantages. First, it is able to separate the produced recombinant proteins from host cell proteases, which are mostly resident in the bacterial cytosol. Second, it is easy to extract molecules from the periplasmic space by osmotic shock which results in an already highly enriched protein extract (Van Den Berg et al., 2015). Third, unlike cytosolic secretion, periplasmic expression allows for efficient disulphide bond formation, which prevents the aggregation or degradation of unfolded polypeptides (Gąciarz et al., 2017). However, the aggregate formation in the periplasmic space and the production of non-secretory proteins are considered disadvantages of periplasmic expression (Baumgarten., 2018).

5.3.3 Efficiency of the OmpA sequence in exporting proteins to the periplasmic space

Despite the high yield of recombinant proteins produced in the *E. coli* expression system, they are mainly expressed as insoluble and inactive inclusion bodies, which must be refolded (Costa et al., 2014). This problem has been overcome by two main strategies. The first is the use of fusion tags, which enables the purification of the tagged protein based on an affinity system. The second involves exporting the recombinant proteins to the periplasmic space to enhance their solubility and function by fusing the N-terminus of the protein to a signal peptide, which acts as the leader in exploring the heterologous protein from the cytoplasm into the periplasmic space (Rosano et al., 2014).

However, It was reported by some studies (e.g., Zhang et al., 2018) that the presence of a signal peptide does not guarantee the efficient exporting of the protein to the periplasmic space. This conclusion can be emphasised by the Western blot analysis conducted in this study on the induced strains cell pellet (i.e., Cytoplasmic produced recombinant protein), figures 5.10, 5.11 and 5.12, which show the bands' sizes corresponding with the expected sizes of the CTX-M proteins (33 KD), which indicates that some of protein hasn't been exported despite the high yielded of recombinant CTX-M extracted from the periplasmic space, in addition to a band at approximately 60 KD, could be dimeric 60KD which is insoluble and therefore in the pellet.

5.3.4 Analysis of phosphorylation statues

Phosphorylation status analysis of the purified CTX-M proteins using Phos-tag gel indicate phosphorylation or another post-translational modification of the three CTX-M proteins (CTX-M-15, CTX-M-14 and CTX-M-14-LIKE). This is supported by the detection of at least two isoforms with the expected size of CTX-M proteins (33KD) using western blot analysis of the distributed proteins in 2D gel. The 2D and Phos-tag data support the conclusion that all the CTX-M proteins are phosphorylated when recombinant expressed in *E.coli*. CTX-M-14-LIKE might even be hyper-phosphorylated. However, phosphorylation status of these protein could be a consequence of the overexpression in *E. coli*, which could be further investigated by testing the same proteins from the parental strains under endogenous conditions followed by protein sequencing.

Determination of any marked differences in MIC (minimum inhibitory concentration) and the kinetics of hydrolysis between the purified, recombinant CTX-M proteins with antibiotics commonly used to treat UTI patients is detailed in Chapter 6.

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Chapter 6: ANTIMICROBIAL SUSCEPTIBILITY OF DOMINANT CTX-M PRODUCING BACTERIA IN NORTH WALES

6.1 Introduction

Antimicrobial susceptibility testing (AST) is a critical procedure that aims to identify the antimicrobial agents that are effective against a particular infectious disease (Lagier et al, 2015). National programs for control and prevention of infectious diseases, as well as hospitals and clinics, rely on the information provided through AST to maintain effective treatment regimens (Bayot & Bragg, 2019). AST aims to determine the lowest concentration of antibiotic required to inhibit visible growth of bacteria over a defined time period, which is known as the minimum inhibitory concentration (MIC)(Lowman, 2018).

The antibiotic concentrations associated with the classification of bacteria as sensitive (S), intermediate (I), and resistant (R) are defined based on these the AST results (Ersoy et al, 2017). The minimum inhibitory concentration (MIC) breakpoints (S, I, R) have been defined by the European Committee on Antimicrobial Susceptibility Testing (EUCAST). A classification of sensitive (S) indicates that the infection should be treated using the recommended antimicrobial dosage for the species and site of infection. A classification of intermediate (I) indicates that the normal dosing of the antimicrobial agent should be increased to achieve a clinical response. A classification of resistant (R) indicates that the microorganism cannot be inhibited with normal dosing of the antimicrobial agent (EUCAST, 2018). These definitions are adopted in this study.

There are many techniques for determining the MIC values of antimicrobial drugs, but the most commonly used method is the broth dilution test. In this test a standardised bacterial suspension $(1.5 \times 10^5 \, \text{CFU})^5$ is inoculated in tubes with two-fold dilutions of antibiotic (e.g., 1, 2, 4, 8) µg/mL) followed by incubation at 35°C for 24 hours (Schumacher et al, 2018). The lowest concentration that prevents bacterial growth (MIC) is determined by examining the turbidity of the antibiotic suspension (Jorgensen & Ferraro, 2009). Despite the method's popularity there are considerable short-comings to this procedure, including the excessive amount of reagents and space required, and the high potential for errors to occur during the preparation of antibiotic solutions (Rennie et al, 2012). The latter challenge has been overcome by replacing the broth

⁵ Colony Forming Unit (CFU) unit used to estimate the number of viable bacteria (bacteria which is able to multiply via fission under the controlled conditions) in a sample.

dilution procedure with the antimicrobial gradient diffusion method in which an antimicrobial concentration gradient is established to test susceptibility of cultured isolates on an agar medium known as Epsilometer test or E-test (Sader & Pignatari, 1994).

The E-test procedure consists of a plastic strip that carries a concentration gradient of dried and stabilised antimicrobial drugs (Leigue et al, 2016). The MIC is determination by placing the graduated antibiotic strips on inoculated agar medium with adjusted bacterial suspension followed by incubation. Due to an immediate release of antibiotics from the E-test strip to the agar surface during the incubation a symmetrical inhibition ellipse is formed along the strip and the MIC value in μ g/mL is read from where the ellipse edge intersects the strip (Khan et al, 2019). This procedure is used for antibiotic susceptibly testing in this study. The results of antibiotic susceptibility testing in this study was further confirmed by the determination of the kinetic parameters of enzyme produced by the tested isolates (β -lactamase enzymes), which can give addition information about the interaction between the antimicrobial agents and enzyme-based resistances conferred by some bacterial species (Zygmunt et al, 1992).

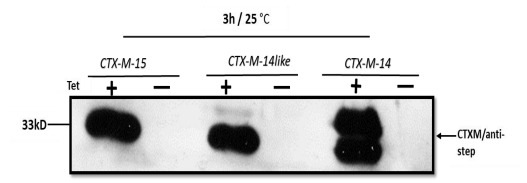
The main objective of this part of the study is to determine any marked differences in MIC and the kinetics of hydrolysis between CTX-M-14 and CTX-M-14-LIKE with antibiotics commonly used to treat UTI patients, i.e. to determine whether the three detected mutations (T55A, A273P, and R277C) change the substrate specificity of CTX-M-14 β-lactamase in addition to the characterisation of CTX-M 15. The latter enzyme has been identified as a dominant CTX-M β-lactamase in North Wales that may in part explain why some UTIs fail to respond to β-lactam antibiotics. To achieve this aim, three recombinant *CTX-Ms*, (*CTXM-15*, *CTXM-14* and *CTX-M-14* LIKE) were recombinantly expressed in *E.coli* and purified (sections 5.2.5 & 5.2.6) before for their antibiotic susceptibility and enzymatic activity were tested using the E-test method and in vitro assays, respectively.

6.2 Results

6.2.1 Overexpression of step-tagged recombinant CTX-M genes.

Freshly cultured chemically competent *E. coli* carrying the plasmid *pASK-IBA2-CTX-M* were induced with a 10µL of anhydrotetracycline solution (2mg/mL dimethylformamide, DMF) at a cell density of 0.5 measured at 550nm (OD550), and incubated with shaking for 3 hr at 25°C. The induced bacteria were then plated on MHB medium with chloramphenicol and E-test strips of five commonly used antibiotics were applied to the plates. The expression level of the three proteins, CTXM-15, CTXM-14 and CTX-M14 LIKE was verified using the Western blot technique (Figure 6.1).

Figure 6.1 Expression level verification of CTX-M proteins using western blot.



Western blot analysis after 3h induction at 25°C with 10μl of anhydrotetracycline of strep-tagged recombinant *CTX-M-15*, *CTX-M-14like and CTX-M-14*. Periplasmic extraction of the three proteins can be seen in the expected size of CTX-M protein (33KD).

6.2.2 Determination of the minimum inhibitory concentration (MIC) of induced strep-tagged recombinant CTX-M genes using E-test

The minimum inhibitory concentrations (MICs) of *E.coli* carrying *pASK-IBA2-CTX-M* were measured for five β-lactams commonly used as a first line treatment for UTIs (Imipenem [IP], Cefoxitin [FX], Nitrofurantoin [NI], Cefotazidime [TZ], and Cefotaxime [CT]) using the E-test protocol (Section 2.2.20) following the MIC break points guidelines of European Committee on Antimicrobial Susceptibility Testing (EUCAST). MIC detection was repeated in biological triplicates for each antibiotic.

6.2.2.1 MIC determination of Imipenem with CTXM-15, CTXM-14 and CTX-M-14-LIKE

An E-test strip with the Imipenem antibiotic was positioned on Mueller Hinton Agar (MHB) with chloramphenicol agar inoculated with suspended (in MHB medium with chloramphenicol) induced or uninduced *E.coli* carrying *pASK-IBA2-CTX-M-15*, *CTXM-14* and *CTXM-14-LIK* with 10 μ L anhydrotetracycline solution followed by incubation at 37°C for 18-24 hrs. The approximate number of the suspended bacteria was standardized before the inoculation by McFarland Standards in which the inoculation suspension was diluted 1:1000 and then compared and adjusted with the turbidity of 0.5 McFarland standard, which gives approximately 1.5×10^8 bacterial cells per mL.

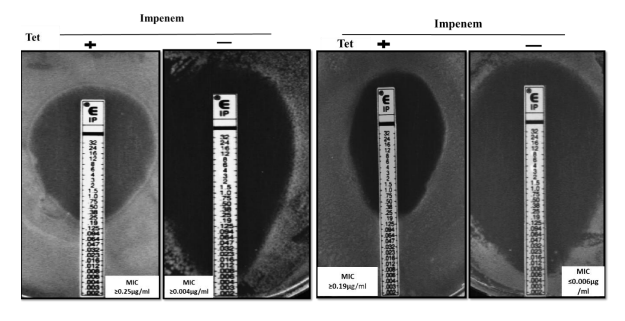
As it is shown in Figure 6.2, the MIC of Imipenem with the induced strains carrying *CTXM-15*, *CTX-M-14*, and *CTXM-14-like* are \geq 0.25 µg/mL (intermediate), \geq 0.19 µg/mL (susceptible), and 0.064 µg/mL (susceptible), respectively. Test for the uninduced strains show MIC < 0.003 µg/mL with Imipenem.

6.2.2.2 MIC determination of Cefoxitin with CTX-M-15, CTX-M-14 and CTX-M-14LIKE

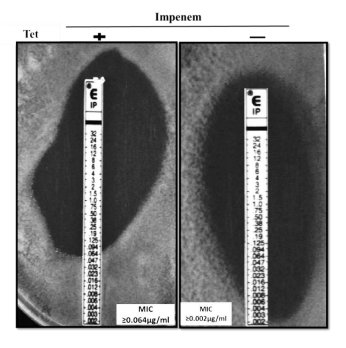
Induced *E. coli* strains carrying *pASK-IBA2-CTX-M-15*, *CTXM-14*, and *CTXM-14-LIK* with $10\mu l$ an anhydrotetracycline solution (Section 6.2) was inoculated on Mueller Hinton Agar (MHB) with an E-strep strip containing the Cefoxitin antibiotic after standardising using the 0.5 McFarland standard (1.5×10^8 approximate bacterial cells per mL) followed by incubation at 37°C for 18-24 hrs.

The highest MIC of Cefoxitin was observed with strains carrying pASK-IBA2- CTXM-14-lik (3 μ g/ml) (intermediate resistance), while the MIC value of the strains carrying *pASK-IBA2-CTX-M-15 or pASK-IBA2-CTXM-14* with Cefoxitin 2 μ g/mL and 1 μ g/mL respectively indicated susceptibility (Figure 6.3).

Figure 6.2 MIC of Imipenem against *E. coli* carrying pASK-IBA2-CTX-M.



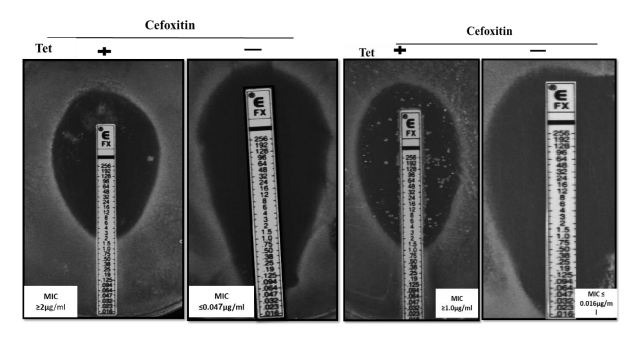
CTXM-15 CTXM-14



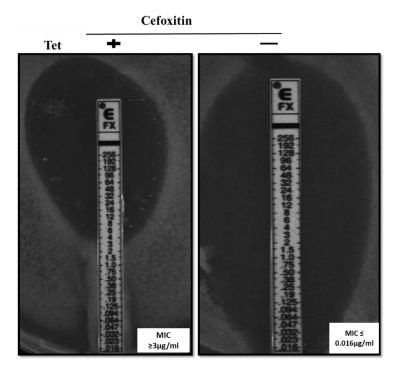
CTXM-14-LIKE

MIC of Imipenem against *E. coli* strains carrying CTX-M genes (CTXM-15, CTX-M-14 and CTX-M-14-LIKE), measured using E-test strips. The highest MIC of Imipenem can be seen with CTX-M-15 (\geq 0.25 µg/mL) which is considered a value of resistance based on EUCAST guideline followed by CTX-M-14(\geq 0.19 µg/mL) and CTX-M-14LIKE (\geq 0.064 µg/mL) which are corresponding with susceptible range. (+) = Induced strain with anhydrotetracycline, (-) = uninduced strain.

Figure 6.3 MIC of Cefoxitin against E. coli carrying pASK-IBA2-CTX-M.



CTX-M-15 CTX-M-14



CTX-M-14-LIE

MIC of Cefoxitin against *E. coli* strains carrying CTX-M genes (CTX-M-15, CTX-M-14 and CTX-M-14-LIKE) measured using E-test strips. Cefoxitin has the highest MIC with CTX-M-14-like ($\geq 3~\mu g/mL$) followed by CTX-M-15 ($\geq 2~\mu g/mL$) and CTX-M-14($\geq 1~\mu g/mL$). (+) = Induced strain with anhydrotetracycline, (-) = uninduced strain (negative control).

6.2.2.3 MIC determination of Nitrofurantoin with CTX-M-15, CTX-M-14, and CTX-M-14-LIKE

The MIC of Nitrofurantoin for induced strains carrying *pASK-IBA2-CTX-M15*, *CTX-M-14* or *CTX-M-14LIKE* was determined by placing a Nitrofurantoin E-test strip on Mueller Hinton Agar (MHB) cultured with the induced or uninduced strains (Section 6.2) and incubated at 37° C for 24 hrs. The suspended strains were adjusted by dilution to 0.5 McFarland standard $(1.5 \times 10^{8} \text{ approximate bacterial cells per mL)}$ before culturing. MIC of an uninduced strain was determined using the same conditions.

The MIC values of Nitrofurantoin with the three tested strains are illustrated in Figure 6.4. This shows that the highest MIC of Nitrofurantoin is with the strain carrying *pASK-IBA2-CTX-M-14LIKE* (\geq 512 µg/mL), which is categorised as a value of Nitrofurantoin resistance (> 64 µg/ml), while the strains expressing CTX-M-15 and CTX-M-14 show a susceptibility against Nitrofurantoin (\leq 64 µl/mL); \geq 4 µg/mL and \geq 6 µg/mL, respectively.

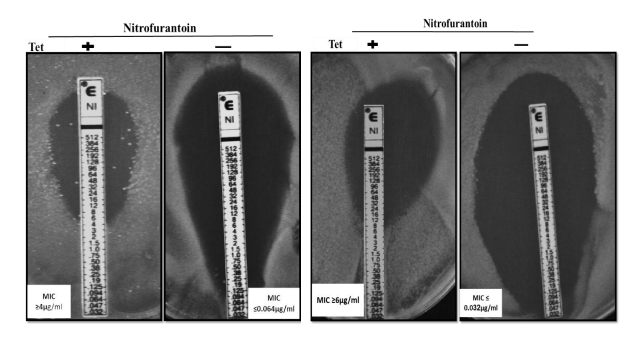
6.2.2.4 Determination of Ceftazidime MIC with CTX-M-15, CTX-M-14, and CTX-M-14 like

In order to determine MIC of Ceftazidime with strains expressing CTX-M14, CTX-M-14LIKE or CTX-M-15 genes, the strains were induced using 10µl anhydrotetracycline (section 6.2) before they were adjusted with 0.5 McFarland standard (1.5 × 10^8 approximate bacterial suspension per mL) and cultured on Mueller Hinton with chloramphenicol Agar with Ceftazidime E-test strips. These were then incubated 37°C for 24 hrs. Figure 6.4 shows that the three strains are susceptible to Ceftazidime ($\leq 1 \mu g/mL$); $\leq 1 \mu g/mL$, $\leq 1 \mu g/mL$, and $\leq 0.75 \mu g/mL$ respectively.

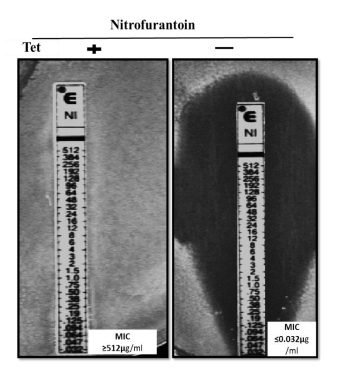
6.2.2.5 Cefotaxime MIC Determination with CTX-M-15, CTX-M-14, and CTX-M-14 like

Cefotaxime E-test strips were used to determine the MIC of Cefotaxime with strains carrying *pASK-IBA2-CTX-M-15*, *CTX-M-14* or *CTX-M-14LIKE*. This was conducted in three steps: (1) induction of strains by anhydrotetracycline (section 6.3) adjusting the suspended strains with 0.5 McFarland standard $(1.5 \times 10^8 \text{ approximate bacterial cell suspension per mL})$; (2) culturing the induced strains on Mueller Hinton with chloramphenicol Agar with E-test strips of Ceftazidime; and (3) incubation at 37°C for 24 hrs. The strains carrying pASK-IBA2-CTX-M-15 are the only strains to have a value of MIC corresponded to a Cefotaxime resistance (32 μ g/mL), while the MIC of cells harbouring CTX-M-14 (1.5 μ g/mL) or CTXM-14-LIKE (0.5 μ g/mL) indicate a susceptibility of these strains to Cefotaxime ($\leq 1 \mu$ g/mL).

Figure 6.4 MIC of Nitrofurantoin against E. coli carrying pASK-IBA2-CTX-M.



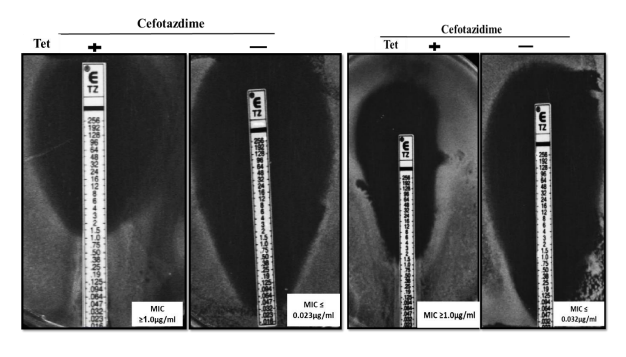
CTX-M-15 CTX-M-14



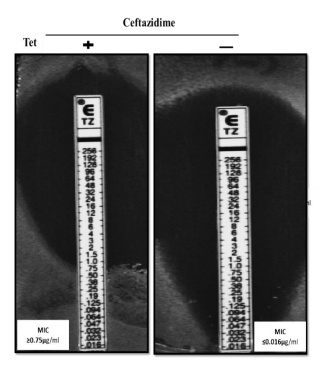
CTX-M-14-LIKE

MIC of Nitrofurantoin against *E. coli* strains carrying CTX-M genes (CTXM-15, CTX-M-14 and CTX-M-14-LIKE), measured using E-test strips. Noticeably, strains carrying *CTX-M-14-LIKE* show high MIC for Nitrofurantoin (\geq 512 μ g/mL) in comparison with the MIC for Nitrofurantoin shown in strains carrying CTX-M-15 (\geq 4 μ g/mL) and CTX-M-14 (\geq 6 μ g/mL). (+) = induced strains with anhydrotetracycline; (-) = Uninduced strains (negative control).

Figure 6.5 MIC of Ceftazidime against E. coli carrying pASK-IBA2-CTX-M.



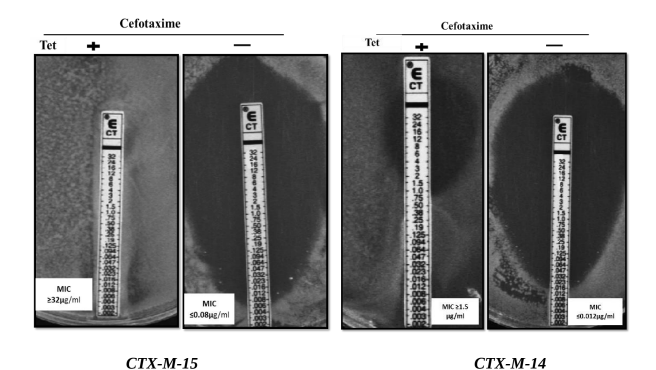
CTX-M-15 CTX-M-14



CTX-M-14-LIKE

MIC of Ceftazidime against *E.coli* strains carrying CTX-M genes (CTX-M-15, CTX-M-14 and CTX-M-14-LIKE), measured using E-test strips. The MIC value for the three proteins is approximately the same (0.75-1.0 μ g/mL). (+) = induced strain with anhydrotetracycline; (-) = negative control (uninduced strain)

Figure 6.6 MIC of Cefotaxime against *E.coli* carrying pASK-IBA2-CTX-M



CTX-M-14-LIKE

MIC of Cefotaxime against *E. coli* strains carrying CTX-M genes (CTX-M-15, CTX-M-14 and CTX-M-14-LIKE), measured using E-test strips. Strains carrying *CTX-M-15* show high MIC for Cefotaxime (\geq 32 µg/mL) in comparison with strains carrying CTX-M-14 (\geq 1.5 µg/mL) and CTX-M-14-LIKE (\geq 0.5 µg/mL). (+) = induced strains with anhydrotetracycline; (-) = uninduced strains (negative control).

Comparison of the MIC breakpoints of all E-Test experiments (Table 6.1) shows that two selected antibiotics, Nitrofurantoin and Cefoxitin, have a higher MIC indicating an enhanced resistance with strains carrying *CTX-M-14-LIKE* than those carrying *CTX-M-14*. This indicates that the three amino acid changes (T55A, A273P, and R277C) in CTX-M-14-Like could enhance the recognition and/or turn-over of Nitrofurantoin and Cefoxitin.

Table 6.1 MICs of *E. coli* isolates harbouring *blaCTX-M*, *15*, *14 or 14-LIKE* towards selected antibiotics.

	MIC(μg/mL)									
	IP		FX		NI		TZ		CT	
blaCTX-M	+	-	+	-	+	-	+	-	+	-
M-15	≥0.25	≤0.004	≥2	≤0.047	≥4	≤0.064	≥1.5	≤0.023	≥32	≤0.08
	≥0.19	≤0.004	≥1.5	≤0.047	≥4	≤0.064	≥1	≤0.023	≥32	≤0.06
	≥0.25	≤0.004	≥2	≤0.047	≥3	≤0.064	≥1	≤0.023	≥32	≤0.08
SD	0.03	0	0.2	0	0.5	0	0.2	0	0	0.01
M-14	≥0.19	≤0.006	≥1	≤0.016	≥6	≤0.032	≥1	≤0.047	≥1.5	≤0.012
	≥0.19	≤0.006	≥1	≤0.016	≥8	≤0.032	≥1	≤0.032	≥1.5	≤0.012
	≥0.19	≤0.006	≥1	≤0.016	≥6	≤0.032	≥1	≤0.032	≥1	≤0.012
SD	0	0	0	0	1.15	0	0	0.008	0.2	0
M-14-LIKE	≥0.064	≤0.002	≥3	≤0.016	≥512	≤0.032	≥0.75	≤0.016	≥0.75	≤0.06
	≥0.064	≤0.002	≥3	≤0.016	≥512	≤0.032	≥0.75	≤0.016	≥0.75	≤0.06
	≥0.064	≤0.002	≥3	≤0.016	≥512	≤0.047	≥1	≤0.016	≥0.75	≤0.06
SD	0	0	0	0	0	0.008	0.14	0	0	0

MIC breakpoints of selected antibiotics with *E. coli* strains carrying CTX-M-15, CTX-M-14 and CTX-M-14-LIKE; (+) = induced strains; (-) = uninduced strains; IP = Imipenem; FX = Cefoxitin; NI = Nitrofurantoin; TZ = Ceftazidime; CT = Cefotaxime; SD = standard deviation. It is noticeable that two antibiotics out of the selected antibiotics show higher MIC with CTX-M-14-LIKE carrying strains than those with strains harbouring CTX-M-14; Nitrofurantoin (\geq 512 µg/mL with CTX-M-14-LIKE and 6 µg/mL with CTX-M-14; significantly higher) and Cefoxitin (3 µg/mL and 1 µg/mL towards CTX-M-14-LIKE and CTX-M-14; not significantly higher). MIC detection was repeated in biological triplicates for each antibiotic.

The MIC values of Nitrofurantoin and Cefoxitin indicated by the E-test assay were subsequently confirmed by determining the affinity (Km) constants of these antibiotics with CTX-M-14 and CTX-M-14-LIKE in order to determine whether the three revealed changes (T55A, A273P, and R277C) could affect the CTX-M-14 enzyme specificity and render cells resistant to Nitrofurantoin and less susceptible to Cefoxitin.

6.2.3 Determination of kinetic constants

The kinetic parameters of the purified CTX-M-14 and CTX-M-14-LIKE proteins, in addition to CTX-M-15, were determined with antibiotics that shows higher MIC with CTX-M-14-LIKE than with CTX-M-14 (see section 2.2.21). These were Nitrofurantoin and Cefoxitin, plus Ceftazidime, which shows low MIC with the three CTX-Ms.

6.2.3.1 Km of CTX-M-14, CTX-M-14-LIKE, and CTX-M-15 towards Nitrofurantoin

The velocity of hydrolysis of Nitrofurantoin by CTX-M-15, CTX-M-14 or CTX-M-14-LIKE were measured by monitoring the changes of absorbance at $\lambda = 266$ nm in a Quartz cuvette in a spectrophotometer over 12 min (Tables: 6.2, 6.3, 6.4, 6.5, 6.6, and 6.7, respectively). The assay was set up such that the amount of the purified protein was not rate limiting and the linear velocities of antibiotic turnover could be monitored by the decline in absorbance, a measure for the hydrolysis of the tested antibiotic. Figures 6.7, 6.8 and 6.9 show the Km value of CTX-M-15, CTX-M-14, and CTX-M-14-LIKE respectively.

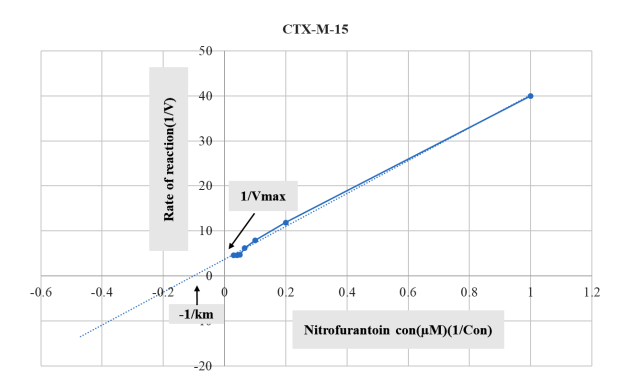
Table 6.2 OD of Nitrofurantoin + CTX-M-15 protein mixture

Time/min	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.02	0.031	0.182	0.309	0.436	0.604	0.784	0.879
1	0.02	0.023	0.17	0.285	0.418	0.585	0.747	0.843
2	0.02	0.012	0.157	0.274	0.397	0.562	0.711	0.807
3	0.02	0	0.143	0.256	0.378	0.538	0.675	0.77
4	0.02	0	0.129	0.238	0.352	0.514	0.639	0.732
5	0.02	0	0.113	0.211	0.296	0.484	0.604	0.698
6	0.02	0	0.098	0.189	0.26	0.45	0.568	0.658
7	0.02	0	0.077	0.167	0.22	0.401	0.531	0.628
8	0.02	0	0.054	0.144	0.169	0.343	0.498	0.592
9	0.02	0	0.03	0.114	0.117	0.272	0.464	0.554
10	0.02	0	0.006	0.088	0.057	0.19	0.424	0.515
11	0.02	0	0	0.06	0.003	0.105	0.389	0.481
12	0	0	0	0.037	0	0.004	0.353	0.447

Table 6.3 Velocity Values (decline OD/min) of Nitrofurantoin + CTX-M-15

Δ /min									
T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ	
0	0	0	0	0	0	0	0	0	
1	0	0.008	0.012	0.024	0.018	0.019	0.037	0.036	
2	0	0.019	0.025	0.035	0.039	0.042	0.073	0.072	
3	0	0.031	0.039	0.053	0.058	0.066	0.109	0.109	
4	0	0.031	0.053	0.071	0.084	0.09	0.145	0.147	
5	0	0.031	0.069	0.098	0.14	0.12	0.18	0.181	
6	0	0.031	0.084	0.12	0176	0.154	0.216	0.221	
7	0	0.031	0.105	0.139	0.0216	0.203	0.253	0.252	
8	0	0.031	0.128	0.165	0.267	0.261	0.286	0.287	
9	0	0.031	0.152	0.195	0.319	0.332	0.32	0.325	
10	0	0.031	0.176	0.221	0.379	0.412	0.36	0.364	
11	0	0.031	0.182	0.249	0.433	0.499	0.395	0.398	
12	0	0.031	0.182	0.269	0.436	0.6	0.431	0.432	
Average	0	0.025	0.092	0.126	0.197	0.215	0.216	0.217	

Figure 6.7 Km determination of CTX-M-15 towards Nitrofurantoin.



Determination of hydrolytic activity of CTX-M-15 towards the Nitrofurantoin antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda = 266$ nm over 12 mins. The concentration of Nitrofurantoin that corresponded to $\frac{1}{2}$ Vamx ($\frac{1}{0.217} \approx 4.60$), solid line, indicates a Km value ($\frac{1}{0.11} \approx 9 \mu M$).

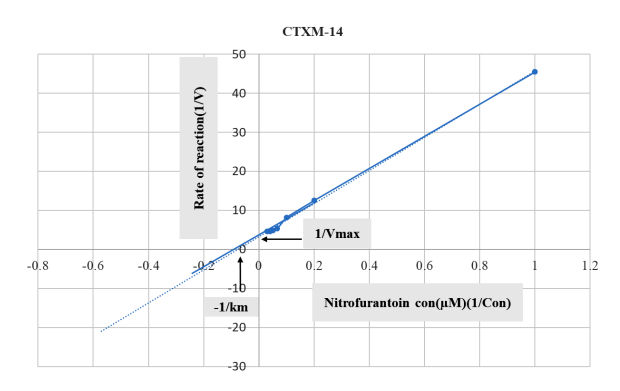
Table 6.4 OD of Nitrofurantoin+ CTX-M-14 protein mixture

Time/min	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.02	0.027	0.141	0.24	0.38	0.522	0.693	0.798
1	0.02	0.02	0.13	0.217	0.362	0.504	0.657	0.762
2	0.02	0.011	0.117	0.206	0.342	0.481	0.621	0.727
3	0.02	0.005	0.105	0.189	0.323	0.458	0.584	0.691
4	0.02	0	0.091	0.171	0.298	0.433	0.549	0.652
5	0.02	0	0.076	0.147	0.243	0.405	0.514	0.618
6	0.02	0	0.061	0.123	0208	0.37	0.478	0.582
7	0.02	0	0.044	0.102	0.168	0.322	0.443	0.553
8	0.02	0	0.024	0.079	0.118	0.263	0.407	0.516
9	0.02	0	0.001	0.05	0.058	0.193	0.373	0.478
10	0.02	0	0	0.024	0	0.111	0.34	0.441
11	0.02	0	0	0	0	0.013	0.301	0.408
12	0	0	0	0	0	0	0.266	0.374

Table 6.5 Velocity Values (decline OD/min) of Nitrofurantoin+CTX-M-14 mixture

				∆/min				
T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0	0	0	0	0	0	0	0
1	0	009	0.011	0.019	0.018	0.018	0.036	0.037
2	0	0.019	0.024	0.031	0.038	0.041	0.072	0.071
3	0	0.025	0.036	0.047	0.057	0.064	0.109	0.108
4	0	0.027	0.05	0.066	0.082	0.089	0.144	0.146
5	0	0.027	0.065	0.089	0.137	0.117	0.179	0.18
6	0	0.027	0.08	0.108	0.172	0.152	0.215	0.216
7	0	0.027	0.097	0.132	0.212	0.2	0.25	0.245
8	0	0.027	0.117	0.158	0.262	0.259	0.286	0.282
9	0	0.027	0.14	0.188	0.322	0.329	0.32	0.32
10	0	0.027	0.141	0219	0.38	0.411	0.353	0.357
11	0	0.027	0.141	0.24	0.38	0.509	0.392	0.39
12	0	0.027	0.141	0.24	0.38	0.522	0.451	0.424
Average	0	0.022	0.08	0.122	0.187	0.208	0.215	0.214

Figure 6.8 Km determination of CTX-M-14 towards Nitrofurantoin



Determination of hydrolytic activity of CTX-M-14 towards the Nitrofurantoin antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda = 266$ nm over 12 mins. The concentration of Nitrofurantoin that corresponded to ½ Vmax (1/0.214 \approx 4.67), solid line, indicates a Km value (1/0.1 \approx 10 μ M).

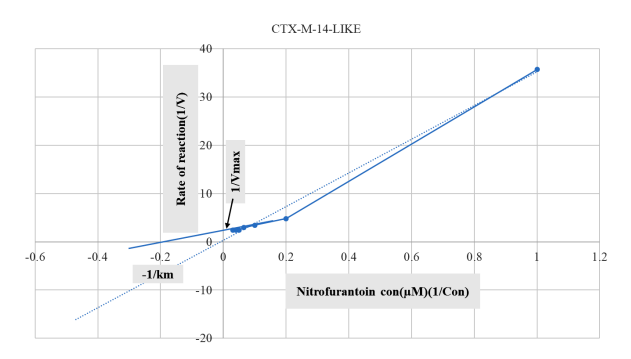
Table 6.6 OD of Nitrofurantoin+ CTXM-14-LIKE protein mixture

T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.03	0.032	0.24	0.358	0.511	0.693	0.797	0.931
1	0.03	0.016	0.138	0.299	0.463	0.662	0.767	0.898
2	0.03	0.004	0.031	0.218	0.405	0.616	0.722	0.852
3	0	0	0	0.117	0.335	0.565	0.672	0.802
4	0.02	0	0	0	0.259	0.428	0.606	0.736
5	0.02	0	0	0	0.167	0.34	0.533	0.662
6	0.02	0	0	0	0.057	0.237	0.445	0.574
7	0.02	0	0	0	0	0.117	0.326	0.471
8	0.02	0	0	0	0	0	0.189	0.351
9	0.02	0	0	0	0	0	0.038	0.213
10	0.02	0	0	0	0	0	0	0.074
11	0.02	0	0	0	0	0	0	0
12	0.02	0	0	0	0	0	0	0

Table 6.7 Velocity Values (Decline OD/min) of Nitrofurantoin+CTX-M-14-LIKE protein mixture

Δ/min										
T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ		
0	0	0	0	0	0	0	0	0		
1	0	0.016	0.102	0.059	0.048	0.031	0.033	0.033		
2	0	0.028	0.209	0.14	0.106	0.077	0.078	0.078		
3	0	0.032	0.24	0.241	0.176	0.128	0.132	0.129		
4	0	0.032	0.24	0.358	0.252	0.265	0.206	0.195		
5	0	0.032	0.24	0.358	0.344	0.353	0.294	0.269		
6	0	0.032	0.24	0.358	0.454	0.456	0.415	0.357		
7	0	0.032	0.24	0.358	0.511	0.576	0.552	0.46		
8	0	0.032	0.24	0.358	0.511	0.693	0.687	0.58		
9	0	0.032	0.24	0.358	0.511	0.693	0.789	0.718		
10	0	0.032	0.24	0.358	0.511	0.693	0.798	0.857		
11	0	0.032	0.24	0.358	0.511	0.693	0.798	0.931		
12	0	0.032	0.24	0.358	0.511	0.693	0.798	0.931		
Average	0	0.028	0.208	0.29	0.34	0.41	0.42	0.46		

Figure 6.9 Km determination of Nitrofurantoin towards CTX-M-14-LIKE.



Determination of hydrolytic activity of CTX-M-14-LIKE towards the Nitrofurantoin antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda=266$ nm over 12 mins. The concentration of Nitrofurantoin that corresponded to ½ Vmax (1/0.46 $\approx\!2.17$), solid line, indicates a Km value (1/0.2 $\approx\!5$ $\mu M)$

6.2.3.2 Km of CTX-M-15, CTX-M-14 and CTX-M-14-like towards Cefoxitin

Absorbance changes (at $\lambda = 235$ nm) were monitored (section 6.2.3.1) to measure the hydrolysis velocity of Cefoxitin by the three CTX-M proteins (Tables 6.8, 6.9, 6.10, 6.11, 6.12 and 6.13 respectively). Km values of CTX-M-15, CTX-M-14, CTX-M-14-LIKE, CTX-M-14-are shown in Figures 6.10, 6.11 and 6.13 respectively.

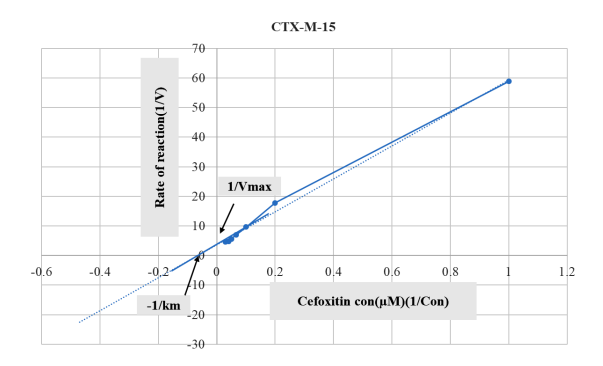
Table 6.8 Cefoxitin+ CTX-M-15 protein mixture

Time(min)/con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.008	0.019	0.078	0.171	0.227	0.356	0.403	0.492
1	0.008	0.006	0.066	0.148	0.209	0.338	0.367	0.456
2	0.008	0	0.055	0.136	0.189	0.315	0.33	0.421
3	0.008	0	0.042	0.124	0.17	0.292	0.292	0.385
4	0.008	0	0.028	0.105	0.146	0.267	0.258	0.346
5	0.008	0	0.013	0.08	0.091	0.239	0.223	0.312
6	0.008	0	0	0.056	0.036	0.204	0.187	0.276
7	0.008	0	0	0.034	0.001	0.156	0.152	0.24
8	0.008	0	0	0.011	0	0.097	0.116	0.211
9	0.008	0	0	0	0	0.027	0.082	0.174
10	0.008	0	0	0	0	0	0.049	0.136
11	0.008	0	0	0	0	0	0.01	0.097
12	0.008	0	0	0	0	0	0	0.066

Table 6.9 Velocity Values (decline OD/min) of Cefoxitin+CTX-M-15

			∆/min					
T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0	0	0	0	0	0	0	0
1	0	0.013	0.012	0.023	0.018	0.018	0.036	0.036
2	0	0.019	0.023	0.035	0.038	0.041	0.073	0.071
3	0	0.019	0.036	0.047	0.057	0.064	0.111	0.107
4	0	0.019	0.05	0.066	0.081	0.089	0.145	0.146
5	0	0.019	0.065	0.091	0.136	0.117	0.18	0.18
6	0	0.019	0.078	0.115	0.191	0.152	0.216	0.216
7	0	0.019	0.078	0.137	0.226	0.2	0.251	0.252
8	0	0.019	0.078	0.16	0.227	0.259	0.287	0.281
9	0	0.019	0.078	0.171	0.227	0.329	0.321	0.318
10	0	0.019	0.078	0.171	0.227	0.356	0.354	0.356
11	0	0.019	0.078	0.171	0.227	0.356	0.393	0.395
12	0	0.019	0.078	0.171	0.227	0.356	0.403	0.426
Average	0	0.017	0.056	0.104	0.144	0.179	0.213	0.214

Figure 6.10 Km determination of Cefoxitin towards CTX-M-15.



Determination of hydrolytic activity of CTX-M-15 enzyme towards the Cefoxitin antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda=235$ nm over 12 mins. The concentration of Cefoxitin that corresponded to ½ Vmax (1/0.214≈4.67), solid line, indicates a Km value (1/0.066≈ 15.1 μ M).

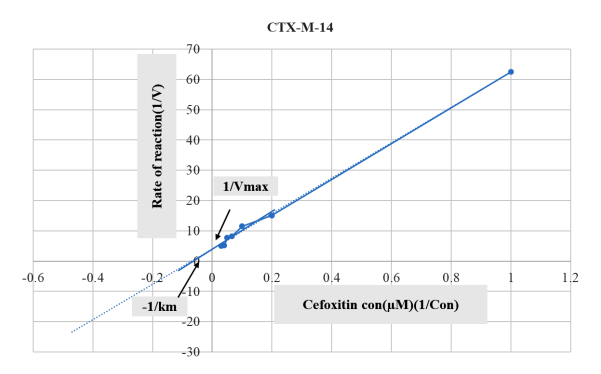
Table 6.10 OD of Cefoxitin+CTX-M-14 protein mixture

T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.011	0.018	0.098	0.13	0.18	0.209	0.297	0.34
1	0.011	0.01	0.087	0.108	0.162	0.189	0.261	0.304
2	0.011	0.002	0.075	0.098	0.143	0.171	0.223	0.27
3	0.011	0	0.061	0.082	0.125	0.149	0.189	0.235
4	0.011	0	0.046	0.066	0.101	0.126	0.157	0.198
5	0.011	0	0.031	0.044	0.048	0.098	0.124	0.165
6	0.011	0	0.014	0.021	0.013	0.065	0.091	0.132
7	0.011	0	0	0.002	0	0.018	0.056	0.102
8	0.011	0	0	0	0	0	0.021	0.066
9	0.011	0	0	0	0	0	0	0.029
10	0.011	0	0	0	0	0	0	0.004
11	0.011	0	0	0	0	0	0	0
12	0.011	0	0	0	0	0	0	0

Table 6.11 Velocity Values (decline OD/min) of Cefoxitin+CTX-M-14 protein mixture

				∆/min				
T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0	0	0	0	0	0	0	0
1	0	0.008	0.011	0.022	0.018	0.02	0.036	0.036
2	0	0.016	0.023	0.032	0.037	0.038	0.074	0.07
3	0	0.018	0.037	0.048	0.055	0.06	0.187	0.105
4	0	0.018	0.052	0.064	0.079	0.083	0.14	0.142
5	0	0.018	0.067	0.086	0.132	0.111	0.173	0.175
6	0	0.018	0.084	0.109	0.167	0.144	0.206	0.208
7	0	0.018	0.098	0.128	0.18	0.191	0.241	0.238
8	0	0.018	0.098	0.13	0.18	0.209	0.276	0.274
9	0	0.018	0.098	0.13	0.18	0.209	0.297	0.311
10	0	0.018	0.098	0.13	0.18	0.209	0.297	0.336
11	0	0.018	0.098	0.13	0.18	0.209	0.297	0.34
12	0	0.018	0.098	0.13	0.18	0.209	0.297	0.34
Average	0	0.015	0.066	0.087	0.12	0.13	0.193	0.198

Figure 6.11 Km determination of Cefoxitin towards CTX-M-14.



Determination of hydrolytic activity of CTX-M-14 towards the Cefoxitin antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda=235$ nm over 12 mins. The concentration of Cefoxitin that corresponded to ½ Vmax (1/0.198 \approx 5.05), solid line, indicates a Km value (1/0.055 \approx 18 μ M).

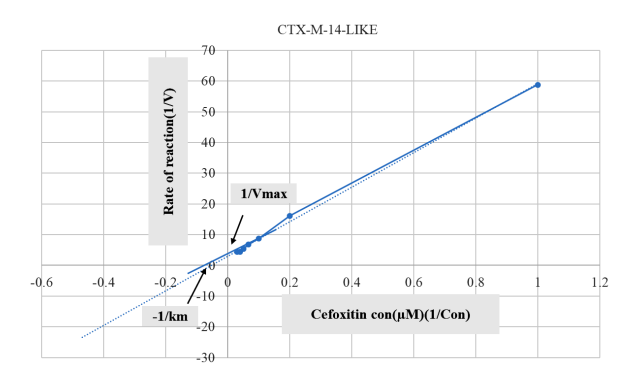
Table 6.12 OD of cefoxitin+CTX-M-14-LIKE mixture

T(min)/con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.013	0.019	0.085	0.177	0.233	0.363	0.411	0.498
1	0.013	0.004	0.072	0.152	0.212	0.345	0.372	0.459
2	0.013	0	0.057	0.138	0.191	0.319	0.332	0.421
3	0.013	0	0.042	0.118	0.173	0.293	0.292	0.383
4	0.013	0	0.025	0.097	0.145	0.265	0.254	0.341
5	0.013	0	0.007	0.069	0.087	0.234	0.216	0.304
6	0.013	0	0	0.042	0.049	0.197	0.177	0.265
7	0.013	0	0	0.018	0.005	0.148	0.139	0.232
8	0.013	0	0	0	0	0.085	0.101	0.192
9	0.013	0	0	0	0	0.014	0.066	0.151
10	0.013	0	0	0	0	0	0.028	0.11
11	0.013	0	0	0	0	0	0	0.075
12	0.013	0	0	0	0	0	0	0.038

Table 6.13 Velocity Values (decline of OD/min) of Cefoxitin+CTX-M-14-LIKE

	Δ/ min											
T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ				
0	0	0	0	0	0	0	0	0				
1	0	0.015	0.013	0.025	0.021	0.018	0.039	0.039				
2	0	0.019	0.028	0.039	0.042	0.044	0.079	0.077				
3	0	0.019	0.043	0.059	0.06	0.07	0.119	0.115				
4	0	0.019	0.06	0.079	0.088	0.098	0.157	0.157				
5	0	0.019	0.078	0.108	0.146	0.129	0.195	0.194				
6	0	0.019	0.085	0.135	0.184	0.166	0.234	0.233				
7	0	0.019	0.085	0.159	0.228	0.215	0.272	0.266				
8	0	0.019	0.085	0.177	0.233	0.278	0.31	0.306				
9	0	0.019	0.085	0.177	0.233	0.349	0.345	0.347				
10	0	0.019	0.085	0.177	0.233	0.363	0.383	0.388				
11	0	0.019	0.085	0.177	0.233	0.363	0.411	0.423				
12	0	0.019	0.085	0.177	0.233	0.363	0.411	0.46				
Average	0	0.017	0.062	0.114	0.148	0.188	0.227	0.229				

Figure 6.12 Km determination of Cefoxitin towards CTX-M-14-LIKE.



Determination of hydrolytic activity of CTX-M-14LIKE enzyme towards Cefoxitin antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda = 235$ nm over 12 min. The concentration of Cefoxitin that corresponded to ½ Vmax (1/0.229 \approx 4.36), solid line, indicates a Km value (1/0.07 \approx 13 μ M).

6.2.3.3 Km of CTXM-14, CTX-M-14-LIKE and CTX-M-15 towards Ceftazidime

Km of Ceftazidime (which had low MIC of the three CTX-Ms) were determined by monitoring Absorbance changes (at $\lambda 274$ nm) (section 6.2.3.1). (Tables 6.14, 6.15, 6.16, 6.17, 6.18 and 6.19). Figures 6.13, 6.14 and 6.15 show Km values of CTX-M-15, CTX-M-14, and CTX-M14-LIKE respectively.

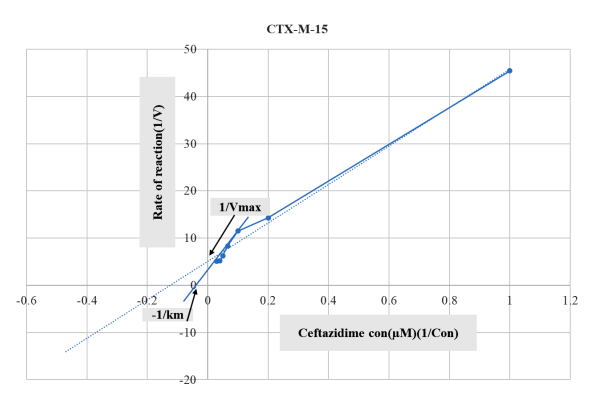
Table 6.14 OD of Ceftazidime+CTX-M-15

T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.011	0.026	0.106	0.12	0.188	0.218	0.308	0.349
1	0.011	0.009	0.095	0.117	0.169	0.197	0.268	0.313
2	0.011	0.002	0.083	0.107	0.149	0.179	0.23	0.278
3	0.011	0	0.069	0.099	0.133	0.157	0.197	0.243
4	0.011	0	0.054	0.073	0.108	0.134	0.175	0.208
5	0.011	0	0.039	0.053	0.057	0.106	0.132	0.172
6	0.011	0	0.022	0.029	0.019	0.043	0.098	0.139
7	0.011	0	0	0.009	0	0.013	0.063	0.109
8	0.011	0	0	0	0	0	0.028	0.074
9	0.011	0	0	0	0	0	0	0.036
10	0.011	0	0	0	0	0	0	0
11	0.011	0	0	0	0	0	0	0
12	0.011	0	0	0	0	0	0	0

Table 6.15 Velocity values (decline of OD/min) + CTX-M-15

T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0	0	0	0	0	0	0	0
1	0	0.017	0.011	0.003	0.021	0.021	0.04	0.036
2	0	0.018	0.023	0.013	0.039	0.039	0.078	0.071
3	0	0.026	0.037	0.021	0.055	0.061	0.111	0.106
4	0	0.026	0.052	0.047	0.08	0.084	0.133	0.141
5	0	0.026	0.067	0.067	0.131	0.112	0.176	0.177
6	0	0.026	0.084	0.091	0.169	0.175	0.21	0.21
7	0	0.026	0.106	0.111	0.188	0.205	0.245	0.24
8	0	0.026	0.106	0.12	0.188	0.218	0.28	0.275
9	0	0.026	0.106	0.12	0.188	0.218	0.308	0.313
10	0	0.026	0.106	0.12	0.188	0.218	0.308	0.349
11	0	0.026	0.106	0.12	0.188	0.218	0.308	0.349
12	0	0.026	0.106	0.12	0.188	0.218	0.308	0.349
Average	0	0.022	0.07	0.087	0.12	0.16	0.192	0.2

Figure 6.13 Km determination of Ceftazidime towards CTX-M-15.



Determination of hydrolytic activity of CTX-M-15 enzyme towards the Ceftazidime antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda=274$ nm over 12 mins. The concentration of Ceftazidime that corresponded to ½ Vmax $(1/0.2\approx5)$, solid line, indicates a Km value $(1/0.036\approx27.5~\mu\text{M})$.

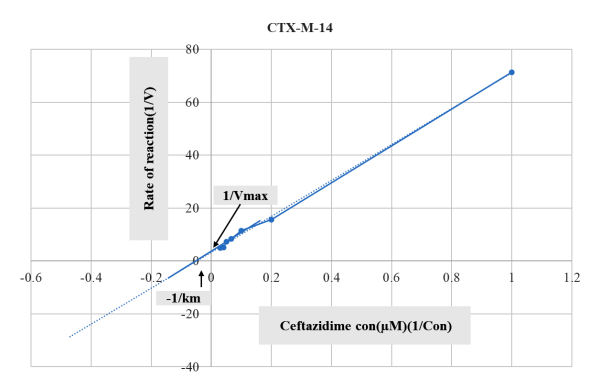
Table 6.16 OD of Ceftazidime+CTX-M-14

T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0.009	0.015	0.095	0.129	0.179	0.205	0.295	0.339
1	0.009	0.08	0.084	0.105	0.159	0.186	0.267	0.301
2	0.009	0.002	0.072	0.095	0.14	0.168	0.22	0.24
3	0.009	0	0.059	0.079	0.122	0.146	0.185	0.232
4	0.009	0	0.043	0.063	0.098	0.122	0.154	0.194
5	0.009	0	0.029	0.04	0.046	0.095	0.12	0.161
6	0.009	0	0.012	0.022	0.09	0.062	0.088	0.128
7	0.009	0	0	0.003	0	0.015	0.053	0.097
8	0.009	0	0	0	0	0	0.018	0.062
9	0.009	0	0	0	0	0	0	0.025
10	0.009	0	0	0	0	0	0	0.002
11	0.009	0	0	0	0	0	0	0
12	0.009	0	0	0	0	0	0	0

Table 6.17 Velocity values (decline of OD/min) +CTX-M-14

T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0	0	0	0	0	0	0	0
1	0	0.007	0.011	0.024	0.02	0.019	0.028	0.038
2	0	0.013	0.023	0.034	0.039	0.037	0.075	0.099
3	0	0.015	0.037	0.05	0.057	0.059	0.11	0.107
4	0	0.015	0.052	0.066	0.081	0.083	0.141	0.145
5	0	0.015	0.067	0.089	0.133	0.111	0.175	0.178
6	0	0.015	0.084	0.107	0.167	0.144	0.207	0.211
7	0	0.015	0.095	0.126	0.179	0.191	0.242	0.242
8	0	0.015	0.095	0.129	0.179	0.205	0.277	0.277
9	0	0.015	0.095	0.129	0.179	0.205	0.295	0.314
10	0	0.015	0.095	0.129	0.179	0.205	0.295	0.337
11	0	0.015	0.095	0.129	0.179	0.205	0.295	0.339
12	0	0.015	0.095	0.129	0.179	0.205	0.295	0.339
Average	0	0.013	0.064	0.087	0.12	0.138	0.197	0.205

Figure 6.14 Km determination of Ceftazidime towards CTX-M-14.



Determination of hydrolytic activity of CTX-M-14 enzyme towards the Ceftazidime antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda=274$ nm over 12 mins. The concentration of Ceftazidime that corresponded to ½ Vmax $(1/0.205\approx4.87)$, solid line, indicates a Km value $(1/0.036\approx27.5\mu\text{M})$.

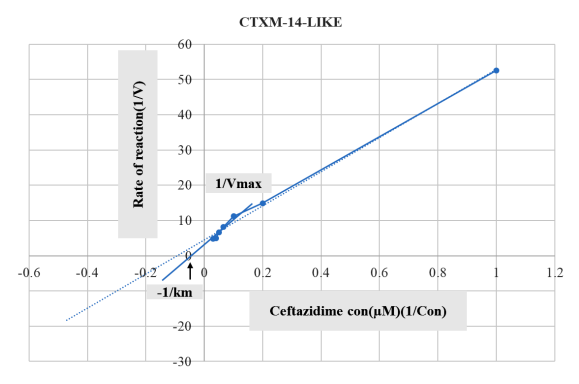
Table 6.18 OD of Ceftazidime+CTX-M-14-LIKE

1 0.013 0.004 0.09 0.112 0.165 0.192 0.264 0.304 2 0.013 0 0.08 0.101 0.146 0.174 0.227 0.27 3 0.013 0 0.061 0.085 0.128 0.152 0.192 0.237 4 0.013 0 0.049 0.069 0.105 0.129 0.16 0.198 5 0.013 0 0.034 0.047 0.051 0.101 0.126 0.167 6 0.013 0 0.017 0.024 0.017 0.069 0.094 0.134 7 0.013 0 0 0.006 0 0.022 0.059 0.105 8 0.013 0 0 0 0 0 0.024 0.069	T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
2 0.013 0 0.08 0.101 0.146 0.174 0.227 0.27 3 0.013 0 0.061 0.085 0.128 0.152 0.192 0.237 4 0.013 0 0.049 0.069 0.105 0.129 0.16 0.198 5 0.013 0 0.034 0.047 0.051 0.101 0.126 0.167 6 0.013 0 0.017 0.024 0.017 0.069 0.094 0.134 7 0.013 0 0 0.006 0 0.022 0.059 0.105 8 0.013 0 0 0 0 0 0 0.024 0.069 9 0.013 0 0 0 0 0 0 0 0 10 0.013 0 0 0 0 0 0 0 0 11 0.013 0 0 0 0 0 0 0	0	0.013	0.021	0.101	0.134	0.183	0.212	0.3	0.346
3 0.013 0 0.061 0.085 0.128 0.152 0.192 0.237 4 0.013 0 0.049 0.069 0.105 0.129 0.16 0.198 5 0.013 0 0.034 0.047 0.051 0.101 0.126 0.167 6 0.013 0 0.017 0.024 0.017 0.069 0.094 0.134 7 0.013 0 0 0.006 0 0.022 0.059 0.105 8 0.013 0 0 0 0 0 0.024 0.069 9 0.013 0 0 0 0 0 0 0 10 0.013 0 0 0 0 0 0 0 11 0.013 0 0 0 0 0 0 0	1	0.013	0.004	0.09	0.112	0.165	0.192	0.264	0.304
4 0.013 0 0.049 0.069 0.105 0.129 0.16 0.198 5 0.013 0 0.034 0.047 0.051 0.101 0.126 0.167 6 0.013 0 0.017 0.024 0.017 0.069 0.094 0.134 7 0.013 0 0 0.006 0 0.022 0.059 0.105 8 0.013 0 0 0 0 0 0.024 0.069 9 0.013 0 0 0 0 0 0 0 0.031 10 0.013 0 0 0 0 0 0 0 0 11 0.013 0 0 0 0 0 0 0	2	0.013	0	0.08	0.101	0.146	0.174	0.227	0.27
5 0.013 0 0.034 0.047 0.051 0.101 0.126 0.167 6 0.013 0 0.017 0.024 0.017 0.069 0.094 0.134 7 0.013 0 0 0.006 0 0.022 0.059 0.105 8 0.013 0 0 0 0 0 0.024 0.069 9 0.013 0 0 0 0 0 0 0 0.031 10 0.013 0 0 0 0 0 0 0 0 11 0.013 0 0 0 0 0 0 0	3	0.013	0	0.061	0.085	0.128	0.152	0.192	0.237
6 0.013 0 0.017 0.024 0.017 0.069 0.094 0.134 7 0.013 0 0 0.006 0 0.022 0.059 0.105 8 0.013 0 0 0 0 0 0.024 0.069 9 0.013 0 0 0 0 0 0 0.031 10 0.013 0 0 0 0 0 0 0 11 0.013 0 0 0 0 0 0 0	4	0.013	0	0.049	0.069	0.105	0.129	0.16	0.198
7 0.013 0 0 0.006 0 0.022 0.059 0.105 8 0.013 0 0 0 0 0 0.024 0.069 9 0.013 0 0 0 0 0 0 0 0.031 10 0.013 0 0 0 0 0 0 0 0 11 0.013 0 0 0 0 0 0 0	5	0.013	0	0.034	0.047	0.051	0.101	0.126	0.167
8 0.013 0 0 0 0 0 0.024 0.069 9 0.013 0 0 0 0 0 0 0 0.031 10 0.013 0 0 0 0 0 0 0 0 11 0.013 0 0 0 0 0 0 0	6	0.013	0	0.017	0.024	0.017	0.069	0.094	0.134
9 0.013 0 0 0 0 0 0 0.031 10 0.013 0 0 0 0 0 0 0 0.007 11 0.013 0 0 0 0 0 0 0	7	0.013	0	0	0.006	0	0.022	0.059	0.105
10 0.013 0 0 0 0 0 0 0.007 11 0.013 0 0 0 0 0 0 0	8	0.013	0	0	0	0	0	0.024	0.069
11 0.013 0 0 0 0 0 0 0	9	0.013	0	0	0	0	0	0	0.031
	10	0.013	0	0	0	0	0	0	0.007
12 0.013 0 0 0 0 0 0 0	11	0.013	0	0	0	0	0	0	0
	12	0.013	0	0	0	0	0	0	0

Table 6.19 Velocity values (decline of OD/min) +CTX-M-14-LIKE

T(min)/Con	0 μΜ	1 μΜ	5 μΜ	10 μΜ	15 μΜ	20 μΜ	25 μΜ	30 μΜ
0	0	0	0	0	0	0	0	0
1	0	0.017	0.011	0.022	0.018	0.02	0.036	0.036
2	0	0.021	0.023	0.033	0.037	0.038	0.074	0.09
3	0	0.021	0.037	0.048	0.055	0.06	0.187	0.105
4	0	0.021	0.052	0.064	0.079	0.083	0.14	0.142
5	0	0.021	0.067	0.086	0.132	0.111	0.175	0.175
6	0	0.021	0.084	0.109	0.167	0.144	0.208	0.208
7	0	0.021	0.101	0.128	0.183	0.191	0.3	0.242
8	0	0.021	0.101	0.134	0.183	0.212	0.3	0.274
9	0	0.021	0.101	0.134	0.183	0.212	0.3	0.313
10	0	0.021	0.101	0.134	0.183	0.212	0.3	0.336
11	0	0.021	0.101	0.134	0.183	0.212	0.3	0.346
12	0	0.021	0.101	0.134	0.183	0.212	0.3	0.346
Average	0	0.019	0.067	0.089	0.122	0.151	0.201	0.207

Figure 6.15 Km determination of Ceftazidime towards CTX-M-14-LIKE.



Determination of hydrolytic activity of CTX-M-14-LIKE towards Ceftazidime antibiotic. The velocity of the reaction is measured by monitoring of the change in absorbance of protein-antibiotic mixture by spectrophotometer at $\lambda=274$ nm over 12 mins. The concentration of Ceftazidime that corresponded to ½ Vmax (1/0.207≈4.83) indicates, solid line, a Km value (1/0.036 ≈ 27.5 μ M).

Comparison of km values (Table 6.20) shows that CTX-M-14-LIKE, with the three substitutions (A55T, A273P, R227C), shows lower Km value (higher hydrolytic activity) than CTX-M-14 for Nitrofurantoin which had a high MIC with CTX-M-14-LIKE by E-test.

Table 6.20 Km values of five purified CTX-M enzymes

	Nitrofu	rantoin	Ce	foxitin	Ceftazidime	
	Vmax	Km(µM)	Vmax	Km(µM)	Vmax	Km(µM)
blaCTX-M						
CTX-M-15	0.217	9	0.214	15.1	0.2	27.5
CTX-M-14	0.214	10	0.198	20	0.205	28.5
CTX-M14-LIKE	0.46	5	0.229	15	0.207	28.5

Km values of three CTX-M enzymes; CTX-M-15, CTX-M-14, CTX-M-14-LKE with three selected antibiotics; Nitrofurantoin, Cefoxitin and Ceftazidime which showed high, medium and low MIC respectively with CTX-M-14-LIKE in comparison to CTX-M-14. As it can be observed CTX-M-14-LIKE with the three changes (A55T, A273P, R227C) shows lower Km value (higher affinity) than CTX-M-14 for Nitrofurantoin.

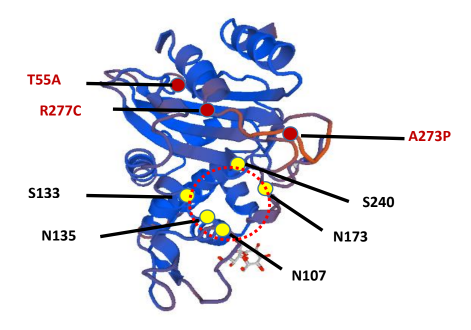
6.3. Discussion

This part of the study aims to determine whether the mutations in CTX-M-14 (T55A, A273P, and R277C) can affect the antibiotic specificity of CTX-M-14 β-lactamase. This was achieved by comparing the minimal inhibitory concentrations (MICs) and the enzyme kinetics in *E. coli* strains expressing either CTX-M-14 or CTX-M-14-LIKE against selected commonly used antibiotics used in the treatment of UTI patients (Nitrofurantoin, Cefotazidime, Cefotaxime, Cefoxitin and Imipenem). The results of the E-tests and in vitro assays show that the three changes in the novel CTX-M (CTX-M-14-LIKE) significantly affect the susceptibility of the strains carrying the recombinant CTX-M-14-LIKE genes makes them more resistant to the Nitrofurantoin antibiotic. In addition, there is a slight effect on Cefoxitin susceptibility in the same strains (i.e. Show bit more resistance to Cefoxitin than those expressing CTXM-14).

As it is shown in Table 6.20, insignificant difference were found in the Km values between CTX-M-14 and CTX-M-14-LIKE with Nitrofurantoin and Cefoxitin. CTX-M-14-LIKE was two folds faster and two fold more affine than CTX-M-14 with Nitrofurantoin and approximately more than one fold faster with Cefoxitin. However, the interesting observation is that the affinities are not significantly different between CTX-M-14 and CTX-M-14-like, although the latter cells are more resistant to Nitrofurantoin (MIC is 64-times higher with CTX-M-14-like than with CTXM-14). This strongly indicates that the three mutations, which are far away from the active site, do not affect the affinity but the turnover for these two antibiotics (Nitrofurantoin and Cefoxitin).

The cells become more resistant because the three mutations seem to promote the conformational changes for Nitrofurantoin and Cefoxitin increasing the rate at which CTX-M-14-like is able to hydrolyse the two antibiotics. This is consistent with the structure model of CTX-M-14 that shows the three amino acid substitutions situated in two loop regions in the beta-strand domain which sits on top of the active site (Figure 6.16). It is therefore not predicted that the three mutated amino acids make direct contact with the β -lactam substrate. Interestingly, two of the substitutions (A273P and R277C) affect the large loop next to the active site. Taken together these findings imply an indirect effect of the substitutions on the conformational changes or the structure of the enzyme.

Figure 6.16 3D structure of the CTX-M-14-LIKE protein.



The 3D structure of CTX-M-14 protein using the SWISS-MODEL tool. The positions of the three changes (T55A, A273P, and R277C) are highlighted. The dotted line highlights the active site with the five amino acids (N107, S133, N135, N173, and S240) that make direct contact with the β -lactam antibiotic substrate. Note that two of the mutated amino acids are predicted to be located in a large loop above the substrate binding site (A273P and R277C).

The catalytic cycle of β -lactamase enzymes is controlled by many structural elements, including as the Ω -loop. The Ω -loop is located at the bottom of the active site entrance and contains highly conserved residues that are important in the catalytic activity of the enzyme. The mutation of these residues results in the loss of enzymatic activity (Pimenta et al., 2014). The effects of structural elements and residues that are located close or in the active site has been demonstrated by many studies. However, few studies have looked into the effect of substitutions in residues distal from the active site on CTX-Ms catalytic activity.

It was reported that the evolution of genetic elements encoding CTX-M enzymes can result from random mutations and recombinant different resistance genes particularly between CTX-Ms classified under groups 1 and group 9, from which CTX-M-15 and CTX-M-14 originate, respectively (Canton et al., 2012). CTX-M-64 is considered an example of this as it contains the middle part of CTX-M-14, with the N- and C-terminals of CTX-M-15, making it more active than CTX-M-14 (Tian et al., 2014). This example show how can the residues distal to the active site are involved in the catalytic cycle of CTX-M enzymes.

The study conducted by He D et al. (2016) has investigated the molecular basis of the differential mechanisms exhibited by CTX-M-14 and of its hybrid (CTX-M-64), which has a higher catalytic activity when compared to CTX-14, in order to check the contribution of residues located distal from active site, in the N and C terminals, to the enzymatic activity. The study concluded that the greater stability of CTX-M-64, which was analysed by thermal stability assay, was behind the enhanced extended-spectrum activity of CTX-M-64.

In this study the location of the three mutations in CTX-M-14-LIKE, which are far away from the active site, emphasise this conclusion as they do not affect the affinity significantly, but do affect the turnover of the two antibiotics (Nitrofurantoin and Cefoxitin). The greater integrity and stability of CTX-M-14-LIKE, which requires further investigation using thermal stability assays, could be associated with the revealed mutation in CTXM-14-LIKE and may drive the high catalytic activity of this enzyme.

The importance of this observation is that Nitrofurantoin has been considered as a standard treatment for UTIs since 2010 (Gardiner et al., 2019). Since then, Nitrofurantoin has shown high activity against uropathogen, including *E. coli* and *Enterococci*, including most extended spectrum β-lactamase producing bacteria addition to all causative bacterial of nosocomial lower UTIs (McKinnnel & Miller, 2011). Generally, many multidrug resistant organisms are still susceptible to Nitrofurantoin (Sanchez et al., 2016). Hence, the spread of CTX-M-14-LIKE β-lactamase in the local communities is expected to worsen the antibiotic resistance problem among UTI patients as the 3 mutations may affect protein stability and/or turnover rates.

Chapter 7: GENERAL DISCUSSION AND CONCLUSIONS

One of the most significant public health concerns in the 21^{st} century is antibiotic resistance. Many bacteria species have developed a range of physiological mechanisms that confer resistance to many of the most commonly used antibiotics (Munita et al., 2016) Resistance to antibiotics results from developing one of three general mechanisms: (1) revoking the target to which antibiotic use to attach to the bacterial cell; (2) decreasing the permeability barrier of the bacterial cell envelope, which reduces the accumulation of antibiotics in the cell; and (3) producing enzymes that inactivate or export the antibiotic. The family of β -lactamases which degrade β -lactam ring antibiotics belongs to the third group (Babic et al., 2006). It is worth noting that within the family of more than 2800 β -lactamases a rapid evolution continues as individual enzymes undergo random mutation thus generating novel substrate specificities or protein features posing the greatest health concerns (Bush and Karen, 2018).

This research focused on a family of β -lactamase enzymes considered to be the most rapidly evolving type of proteins, the CTX-M-type extended spectrum β -lactamases (CTX-M type ESBL). Specifically, this research investigated the extent to which CTX-M-type ESBLs contribute to the resistance phenomenon detected in isolates from urinary tract infection (UTI) patients in North Wales. It was assessed whether these CTX-M producing bacteria carry the same or different CTX-M genes and the extent to which different CTX-M enzymes vary with regard to the antibiotic resistance spectrum.

The aim of the study was achieved in four main steps: (1) the analysis of UTI patient data from one of the main referral hospitals in North Wales in order to identify the risk factors associated with isolations of ESBL-producing *E. coli* in urine samples from hospitalised patients and outpatients; (2) by determining the contribution of CTX-M-type ESBL, and its dominant type, to antibiotic resistance in 300 ESBL-producing isolates obtained from UTI patients in three referral hospitals in North Wales; Ysbyty Gwynedd, Glan Clwyd Hospital, and Wrexham Maelor Hospital by testing the isolates with multiplex PCR using primers for phylogenic groups of CTX-M family followed by sequencing; (3) by cloning of dominant CTX-Ms in expression plasmids that produce C-terminally affinity tagged protein to facilitate its purification to obtain abundant amount of recombinant CTX-M protein to measure its catalytic activity against commonly used antibiotics for UTI patients; and (4) by determining minimal

inhibitory concentrations (MIC) and catalytic activities of the dominant CTX-Ms against antibiotic used to treat UTIs compared to antibiotics used for routine ESBL screening.

7.1. The significantly higher incidence of UTIs in North Wales

Data from Ysbyty Gwynedd (YG) hospital in Bangor indicates a significantly higher number of patients referred for urinalysis due to symptoms of urinary tract infections over the three years of the study in comparison to other regions around the world with approximately the same population size and health care level. The suggested explanation for this high UTI incidence in North Wales is the high percentage of old people in the population of North Wales Which, according to Welsh Government services and information, is expected to rise from 18% (as determined in 2008) to almost 26 % of the Welsh population by 2033. This age-group is more likely to experience UTIs than younger individuals. Another explanation for the increased UTI prevalence in North Wales is the high percentage of rural communities with approximately 80% of Welsh land being managed as farmland, and the close proximity of these community to antibiotic treated livestock. However, it should be remembered that AB usage is much reduced on the beef and sheep farms typical of North Wales (UK Veterinary Antibiotic Resistance and Sales Surveillance Report, 2019), making this less likely to be the main explanatory factor for our results.

7.2. Uropathogenic E. coli (UPEC) as causative organism of nosocomial UTIs

Conclusions of many studies (e.g., Kollef et al., 2016) indicate that uropathogenic *E. coli* represents less than 50% of pathogens causing nosocomial (hospital) acquired UTIs, while community acquired UTIs are caused predominantly by *E. coli*. The outcomes of this study indicates an increased spread of uropathogenic *E. coli* in hospitals as it represents 79% of the detected pathogens of hospital acquired UTIs. The possible reason behind this rise of uropathogenic *E. coli* in hospitals is the divergence of some *E. coli* strains from a commensal to pathogenic form inside the body through the acquisition of specific virulence factors via horizontal DNA transfer of plasmids, transposons, and bacteriophage. The transfer of genetic material also moves antibiotic resistance genes between bacteria thus increasing both the pathogenicity and resistance of *E. coli* (Gyles et al., 2014).

Unlike other uropathogenic bacteria such as *Klebsiella pneumonia*, *Pseudomonas aeruginosa*, *Proteus mirabilis* and *Enterococcus* (Hof, 2017), *E. coli* is characterised by the presence of mobile genetic elements, pathogenicity islands that carry specialised virulence factors which provide *E.coli* with a high degree of genetic diversity (Wiles, 2008). The virulence factors of uropathogenic *E. coli* is associated with the development of UTIs making infections more

complicated to treat. These factors can be divided into two groups: (1) surface virulence factors, which are characterised by various types of adhesive fimbriae that enable $E.\ coli$ to attach within the urinary tract and (2) secreted virulence factors, which include the toxins produced by colonizing $E.\ coli$ e.g. α -haemolysin which causes the inflammatory response that leads to the symptoms of UTIs (Bien, 2012).

7.3. Remarkable rise of antibiotic resistance conferred by ESBL-producing isolates

The precise definition of extended spectrum β -lactamase has not been agreed, but the commonly used definition is that ESBLs are a group of β -lactamases that are characterised by their broad hydrolytic activity against first, second, and third generations of cephalosporins, in addition to aztreonam, and can be inhibited only by β -lactamase inhibitors (Rahman et al., 2018). This makes ESBL-producing isolates a major problem for clinical therapeutics as many clinically used antibiotics contain a β -lactam ring as the functional element (Shakya et al., 2018).

The data analysed in this study can be added to other studies that show an alarming increase in the threat posed by antibiotic resistance conferred by ESBLs due to the two-fold increase in the number of ESBLs detected over the three years of this study, 2011-2013, (Chapter 3; Table 3.5). One of the possible reasons behind the rapid growth of ESBL-based resistance is the inaccurate phenotypic methods used for ESBL screening in clinical laboratories, e.g. the double-disc synergy test (Bajpai et al., 2017) which is unable to distinguish between approximately 200 types of ESBLs identified to date (Ghafourian et al., 2015). This means that routine screening methods are unable to determine the strain and gene responsible for the production of each ESBL enzyme, which are not all inhibited by the same antibiotic. In addition, ESBL genes are often found on low copy number IncF plasmids with multiple other resistance genes, such as the aminoglycoside/fluoroquinolones resistance conferring gene (Robberts et al., 2009). Together, these factors could lead to treatment failure in many cases due to the use of inappropriate antibiotic resulting in outbreaks of ESBL-based antibiotic resistance. Developing molecular-based techniques with the additional ability to detect low level genetic resistance for detection of ESBL in clinical diagnostic laboratories could decrease the false diagnosis of ESBLs and limit the rapid growing of ESBL-based resistance.

7.4 Dominant CTX-M-type ESBLs in North Wales

Screening of isolates diagnosed in the clinical laboratories of the main three referral hospital in North Wales as ESBL producers for CTX-M genes, using multiplex PCR followed by sequencing, revealed that the most detectable CTX-M among these isolates is CTX-M-15. This isolate represented more than half of all sequenced samples (Chapter 4, Tables: 4.2,4.4 and 4.6). This is a very large increase in CTX-M-15 prevalence in North Wales when compared to that found in previous studies e.g. Mushtaq et al., (2003) who only detected CTX-M-15 in the South and North of England, in addition to some regions of Northern Ireland. North Wales, was part of Mushtaqs et al' study.

Unlike other CTX-M variants, the *CTX-M-15* gene on the conjugative plasmid *IncF*, which facilitates its horizontal transfer by the conjugation process, provides an explanation for the high global prevalence of CTX-M-15 (Orhue et al., 2017).

Some studies (e.g. Laurent Poirel et al., 2013) have investigated the spread of ESBL-producing isolates among companion animals and found through the molecular screening of these isolates high rates of CTX-M-15 producing *Klebsiella pneumonia*. Acquiring resistance confirmed by CTX-M-15 from animal reservoirs is very likely in North Wales hospitals with its large farming community that lives in close proximity to food producing animals. Where resistance in animals are found the resistance plasmid may be transferred horizontally to human. Acquiring resistance genes from animal reservoirs has become an increasingly common practice over recent years and it is used for highly effective antibiotics like colistin. A discussion of this can be found in Wang R et al.'s study conducted in China, which detected the emergence and spreading of colistin resistance gene mcr-1 in pig isolates (Wang R et al., 2018)

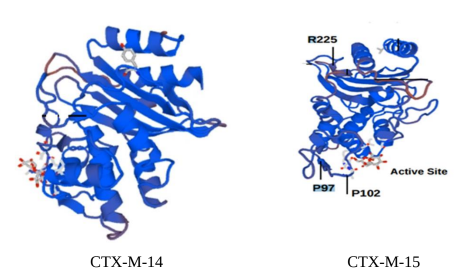
CTX-M-15 is characterised by its high activity against Ceftazidime (Po et al., 2017). The antibiotic profiles of the isolates carrying CTX-M-15 in one the three referral hospitals of the study (Ysbyty Gwynedd hospital at Bangor) shows resistance of these isolates also to other antibiotics like amoxicillin and ciprofloxacin that are known to inhibit the activity of CTX-M-15 (Chapter 3;Table 3.2). The existence of a novel mutation in the *CTX-M-15* gene which was selected for by the misuse of antibiotics for treatment of patients due to inaccurate phenotypic detection, could be the reason behind the development of resistance to these antibiotics.

7.5 The novel CTX-M-14 type ESBL revealed in North Wales

In addition to the spread of CTX-M-15 in the screened samples of this study, sequencing has revealed also a new β -lactamase gene that is closely related to *CTX-M-14* at the protein level, with the substitution of only three amino acids (T55A, A273P, and R227C) in the *CTX-M-14* gene. This novel β -lactamase was detected in approximately 11% of sequenced samples from one of the three hospitals of the study (Ysbyty Gwynedd, Bangor). The presence of this novel CTX-M at only one of three hospital sites included in this study supports the idea that the novel β -lactamase has been transferred horizontally on a plasmid vector from the farming community. Because the close similarity of the novel CTX-M to CTX-14, the novel gene was named *CTX-M-14-LIKE*.

CTX-M14 is the second most frequently reported globally after CTX-M-15. These two variations are classified under two different CTX-M groups: CTX-M group 1 and CTX-M-group 9, respectively (D'andrea et al., 2013). Figure 7.1 shows the 3D structure of the two CTXMs.

Figure 7.1 3D structure of CTX-M-14 and CTX-M-15



3D structure of CTX-M-14 and CTX-M-15. The active site is indicated. Proline 97 and Proline 102 and arginine 225 are Key differences between CTXM-15 and CTXM-14 in otherwise highly conserved regions. The images were Generated using Swiss Model (1ylp.1.A 99.62% identity).

The second part of this study focused on the comparison between the two CTX-Ms (CTX-M-14 and CTX-M-14-like) in terms of antibiotic resistance and how the substitution between the two enzymes impact on their hydrolytic activity against antibiotics commonly used to treat patients with UTIs.

7.6 Production and purification of recombinant CTX-M proteins

In order to study the enzymatic activity of the dominant CTX-Ms (CTX-M-15 and CTX-M-14-LIKE) and to compare the catalytic activity of the novel β-lactamase (CTX-M-14-LIKE) with the activity of CTX-M-14, recombinant proteins of these CTX-Ms were required. These were obtained by the cloning of the full length CTX-M genes in the *E. coli* expression plasmid *pASK-IBA2C* which allows for the purification of C-terminally strep-tagged recombinant proteins after their induction from the periplasmic space. Among two strategies of protein secretion by pASK-IBA2C; cytosolic and periplasmic expression, the later strategy was used to produce proteins of the three CTX-Ms (CTX-M-15, CTX-M-14, and CTX-M-14-like) because it has several advantages. Among these are, the ability to obtain highly enriched protein extract due to the ease of extraction of proteins from the periplasmic space by osmotic shock that removes the outer membrane and to separate cytoplasmic host cell protease from the recombinant protein product. In addition, the periplasmic expression of the genes reduces aggregation or degradation of unfolded polypeptides as the protein that forms protein multimers is not exported (Van Den Berg et al., 2015).

The export of the recombinant CTX-M proteins from cytoplasm into the periplasmic space is led by a signal peptide included in the pASK-IBA2C plasmid, the OmpA (Outer Membrane Protein A) leader sequence which is fused in the N-terminus of the protein. The export of the CTX-M protein into the periplasmic space enhances their solubility and function as recombinant protein produced in the cytoplasm of *E. coli* predominantly forms insoluble and inactive inclusion bodies (Rosano et al., 2014).

The efficiency of the OmpA leader sequence in exporting the CTX-M protein to the periplasmic space has been checked by western blot analysis of induced cell pellet (i.e., Cytoplasmic produced recombinant protein) in ordered to verify the full exporting of recombinant CTX-M proteins to periplasmic space (Chapter 5 ,Figures;5.9,5.10 and 5.11). This analysis by western blot has shown bands with sizes corresponding to the expected size of the CTX-M protein of 33KD, which indicates that some of protein hasn't been exported despite the high yielded of recombinant CTX-M extracted from the periplasmic space. In addition, a

band at approximately 60KD was detected in cell pellet analysed by western blot, this band could be dimeric 60KD which is insoluble and therefore in the pellet.

The produced recombinant CTX-M was C-terminally tagged with the short peptide Strep-Tag (Ala-Trp-Arg-His-Pro-Gln-Phe-Gly-Gly) to purify the protein from the periplasmic extract by its affinity to strep-tactin that is included in prepacked columns and allows for gravity-flow purification of the strep-tagged CTX-M proteins. The purity of the proteins was checked using Coomassie stain to detect additional protein in the preparations that is not Strep-tagged (Chapter 5; Figure 5.14).

7.7 Phosphorylation status analysis of the CTX-M proteins

Protein phosphorylation is responsible for a variety of protein properties, including protein stabilization and degradation, and some biochemical activities (Lin J et al., 2010). Phosphorylation of beta-lactamase produced in *E.coli* has been reported in many studies (e.g. phosphorylated TEM-b-lactamase was determined to be phosphorylated by threonine residue, which effects its secretion to periplasmic space) (Udo H et al. 1995). Analysing of CTX-M proteins sequences bioinformatically using the NetPhos tools revealed that CTX-M-14 and CTX-M-15 both possess some potential phosphorylation sites at serine, threonine, and tyrosine residues (Chapter 5; Figure 5.13). This provided the motivation to analyse the phosphorylation status of the purified CTX-M proteins using two techniques, Phos-tag gel and two dimensional (2D) gel analysis, in order to determine how de-phosphorylation could affect the enzymatic activity of CTX-M proteins against commonly used antibiotics.

Bands detected in the Phos-tag gels (as the only phosphorylated protein can be visualised in this gel as the modified protein moves slower) indicate phosphorylation of the three CTX-M proteins (CTX-M-15, CTX-M-14 and CTX-M-14-LIKE) (Chapter 5; Figure 5.15). This is supported by the detection of at least two isoforms with the expected size of CTX-M proteins (33KD) using western blot analysis of the distributed proteins after isoelectric focusing (2D gel analysis) (Chapter 5; Figure 5.14). It is noteworthy that the 2D technique detects all posttranslational modifications that affect the overall charge of the protein of which phosphorylation is only one form. But combined with the finding that the fraction of the recombinant protein runs more slowly in the presence of the phos-tag that is specific for phosphorylation. It is feasible to conclude that a sub fraction of the recombinant CTX-M protein is phosphorylated. It is however possible that the phosphorylation status of these

proteins is a consequence of the overexpression in *E. coli*, which could be further investigated by testing the same proteins from the parental strains under endogenous conditions.

7.8 Comparative analysis of catalytic activity between CTX-M-14 and CTX-M-14-LIKE

Analysis of the catalytic activity of recombinant CTX-M-14 and CTX-M-14-LIKE was conducted in order to determine any marked difference in the minimal inhibitory concentration (MIC) and the kinetics of hydrolysis between these two enzymes to determine whether the three substitutions (T55A, A273P and R277C) change the substrate specificity of CTX-M-14 β -lactamase.

Comparison of the MIC values of strains expressing either CTX-M-14 or CTX-M-14-LIKE against selected antibiotics commonly used in the treatment of UTI patients shows that the three changes in the novel CTX-M (CTX-M-14-LIKE) render cells more resistant to Nitrofurantoin (MIC; \geq 512 µg/mL with CTX-M-14-LIKE while \geq 8 µg/mL with CTX-M-14) (Chapter 6;Table 6.1). An only two-fold increase in the affinity (Km) of CTX-M-14-LIKE compared to CTX-M-14 was found for Nitrofurantoin and one fold was found for Cefoxitin (chapter 6; Table 6.20). Although the affinity (Km) is independent on the amount of active protein in the in vitro test the maximal velocity (Vmax) is however dependent on the percentage of active protein in the purified preparation, it is difficult to conclude that a factor of two is a significant increase. The reason is that the graphical Laneweaver-Burk method has not such a high resolution.

Given that the affinities are not significantly different between CTX-M-14 and CTX-M-14-like for Nitrofurantoin; instead, MIC is 64-times higher with CTX-M-14-like than with CTXM-14, the three mutations that reside in the beta-strand domain opposite to the active site (Chapter 6; Figure 6.16) affect the conformational changes required for the effective hydrolysis of Nitrofurantoin thus increasing the turnover rate for this antibiotic. Another possibility is that the mutations increase the protein stability or amount in the endogenous coliform as shown for other mutations in β -lactamases (Jacquier et al., 2013).

Modelling the three amino acid substitutions onto the protein structure of CTX-M-14-like revealed that all three substitutions map to the domain opposite the active site. Interestingly, two of the substitutions A273P and R277C affect the large loop next to the active site which may have an effect on the catalytic cycle of the mutated enzyme Taken together these findings

imply an indirect effect of the substitutions on the conformational changes or the structure of the enzyme.

The effects of amino acid substitutions on CTX-M β -lactamase hydrolytic activity has been demonstrated by many studies, with the focus of these studies mostly on amino acids located in active site (e.g. Patel et al., 2015). In contrast, few studies have looked into the effect of substitutions in residues distal from the active site on CTX-Ms catalytic activity. He D et al., (2016) detected some differences in residues between CTX-M-14 and one of its variants CTX-M-64 that are located mainly at both the C and N termini (e.g., V29A, L48A, V286A) . These substitutions are reported to increase the hydrolytic activity of CTX-M-14 variant CTX-M-64 toward some B-lactams due to its greater stability which was analysed by thermal stability assay.

The result presented here support the conclusions of the studies regarding the impact of distal residues on CTX-Ms hydrolytic cycle, but expand these findings to other antibiotic agent, most notably Nitrofurantoin. The enhanced activity of CTX-M-14-LIKE associated with the three revealed mutations (T55A, A273P and R277C) could be resulted from the greater integrity and stability of CTX-M-14-LIKE, which requires further investigation using thermal stability assays.

7.9 Conclusion

This study aimed to create an epidemiological snapshot of the antibiotic resistance conferred by CTX-M-type ESBL producing bacteria in UTI patients in North Wales and to determine to what extent do the different CTX-M enzymes vary with regard to the antibiotic resistance spectrum. The data obtained from the three study hospitals; Ysbyty Gwynedd, Glan Clwyd Hospital, and Wrexham Maelor Hospital indicate a significantly high incidence of UTIs in North Wales which may correlate with high percentage of rural communities in the region that live in close proximity to the antibiotic treated livestock.

Multiplex PCR amplification of β -lactamase genes of the *bla*CTX-M groups 1, 2, 9, and 8/25, followed by DNA sequencing of ESBL-producing isolates from three referral hospital in North Wales revealed that CTX-M-15 is the dominant variant of CTX-M found in UTI patients in the study region. In addition, this study revealed a new β -lactamase gene that is closely related to CTX-M-14 at the protein levels and was unique to Ysbyty Gwynedd at Bangor.

Antimicrobial susceptibility testing in addition to determinations on Km constants of CTX-M-14 and CTX-M-14-LIKE with antibiotics commonly used to treat UTI patients indicates enhanced activity of CTX-M-14-LIKE against Nitrofurantoin (significantly) and Cefoxitin (slightly). This is due to the effects of three substitutions A55T, A273P and R277C that reside distal to the active site. This strongly supports the conclusion that, in addition to active-site mutations, the hydrolytic cycle of CTX-M enzymes can be affected by distal residues which are most likely to be associated with the substrate turn-over rates and/or stability of the enzyme.

The impact of the study and the implications for public health policy

Since the most common clinical phenotypic detection methods currently in use are unable to provide detailed information about ESBL gene types, the main impact of this study is to motivate clinical laboratories to invest in the development of quick and accurate molecular techniques. These methods might include PCR-based assays for detecting the genes responsible for the resistance and an efficient molecular-based identification method for the detection of resistant species. This approach could also enable non-expressing ESBL genes to be detected, which would improve the ability of clinicians to diagnose and treat antibiotic resistance UTIs, in addition to creating an epidemiological snapshot of the antibiotic resistance pattern in UTI patients in North Wales and potentially globally.

7.10 Future work

One of the limitations of this study is the difficulty in obtaining clinical samples and data from the study hospitals, which limits the number of samples used in the study. Further, test of Nitrofurantoin resistant isolates will be a priority of future work in order to check whether the novel CTX-M protein exists in the general population of Wales and to investigate the origin of this gene, which to date has only been detected in one of the three study hospitals (Ysbyty Gwynedd). In addition, more information is required on the background of the patients whose samples were provided. This could be difficult to obtain due to issues of data protection and patient anonymity, but at a minimum it would be useful to distinguish "rural" from "urban" samples as this would help to test hypotheses regarding why there was a disproportionately high prevalence of patients with AB-resistant UTIs. Determination of CTX-M-14-LIKE hydrolytic activity against more β -lactam antibiotics should also be a priority for future work.

The phosphorylation analysis of the recombinant CTX-M was promising and further investigation of the phosphorylation status by testing the same proteins from the parental strains under physiological conditions (i.e. not under recombinant conditions) should be a main focus of subsequent studies. This could be achieved by a proteomics approach sequencing endogenous material from Nitrofurantoin resistant isolates. If post-translational modifications also exist, phosphor-specific antibodies could be generated to continue the study under physiological conditions. Further investigation should be conducted to determine how the three revealed residues that are located distal to the active site of CTX-M-14-LIKE result in higher catalytic activity. This work should use thermal stability assays to determine whether these three distal amino acid substitutions can affect the integrity and stability of the protein.

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Appendix I

Wrexham hospital samples DNA sequences;

3:

BLAST output: Klebsiella pneumoniae strain KP58 extended-spectrum betalactamase **CTX-M-59** (blaCTX-M) gene, blaCTX-M-59 allele, partial cds Sequence ID: MH661248.1Length: 817Number of Matches: 1

ACGCAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTA
ACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCA
CCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGT
TAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAA
AGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCG
ATAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGT
TCGCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACA
CCGCCATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGC
GGAATCTGACGCTGGGTAAAGCATTGGGCGACCAACGGGCGCAGCTGGTGACATGGA
TGAAAGGCAATACCACCGGTGCAGCGAGCAACTTCAGCTGGACTTCCTTGGGGTT
GTGGGGGGGATAAAACCGGCAGCGAGCGACCTATG

BLAST output: Klebsiella pneumoniae isolate KSH203 plasmid pKSH203-**CTX-M-3**, complete sequence

Sequence ID: CP034325.1Length: 156910Number of Matches: 1

7

BLAST output: CTX-M-15 (blaCTX-M-15) gene, partial cds Sequence

ID: gb|KR338941.1|Length: 867Number of Matches: 1

9:

 ${\tt CGCAAAAACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTGATGG}\\ {\tt CCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTTG}\\$

AGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGA
CGATGTCACTGGCTGAGCTTAGCGCGGCCGCCGCTACAGTACAGCGATAACGTGGCGATGA
ATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGG
GAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCG
ATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGG
GTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCA
CCGGTGCAGCGAGCATTCAGGCTGGACTGCCTTCCTGGGTTGTGGGGGATAAAACCG
GCAGCGGTGGCTATGGCACCACAACCGAATATCGAAAATTTC

BLAST output: Klebsiella pneumoniae strain E161KPMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds

Sequence ID: KY640551.1Length: 632Number of Matches: 1

11:

BLAST output: Proteus mirabilis PM341/03 pPM341/03 blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-66**, complete CDS

Sequence ID: NG 049017.1Length: 1076Number of Matches: 1

15:

TTAGGAAGTGTGCCGCTGTATATCATCGATACCGCA

GGCCTGCGCGATGCTAATGACGAAGTGGAACGTATCGGCATCGAGCGCGCCTGGCAGGAG

ATTGCCCAGGCCGATCGGGTGCTGTTTATGGTCGATGGCACCACCAACGATATCG

BLAST output: Klebsiella pneumoniae UHKPC07, complete genome Sequence ID: gb| CP011985.1|Length: 5322270Number of Matches: 1

17:

 $CGCAAGATTTAGGAAGTGTGCCGCTGTATGCGCAAACGGCGGACGTACAGCAAAAACTTGCCG\\ AATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAGCAGATAATTCGC\\$

BLAST output : *Shigella sp.* **SH223 blaCTX-M-108 (CTX-M-108) g**ene, partial cds Sequence ID: gb| JF274245.1|Length: 864Number of Matches: 1

19:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAAC
ACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACC
AGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAAGTGAAAGCGAACCGAATCTGTTA
AATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAG
CACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGAT
AACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTC
GCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACC
GCCATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAAACTCTGCGG
AATCTGACGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATG
AAAGGCAATACCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTTCCTGGGTTGTG
GGGGGATAAAACCGGCAGCGAGCGAGCTATGGCACAGC

BLAST output : Klebsiella pneumoniae subsp. pneumoniae strain KPTR13-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence

ID: MK113957.1Length: 1027Number of Matches: 1

21:

BLAST output: Escherichia coli strain E78ECMO CTX-M-15 beta-lactamase

(blaCTX-M-15) gene, partial cds

Sequence ID: KY640536.1 Length: 641 Number of Matches: 1

23:

ACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACCACTTCACCT

BLAST output: Escherichia coli strain IPK192 beta-lactamase CTX-M-32

(blaCTX-M) gene, blaCTX-M-32 allele, complete cds

Sequence ID: MH900527.1Length: 876Number of Matches: 1

25:

AACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCA
GTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATC
AGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAA
TGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCCTACAGTAC

BLAST output : Escherichia coli strain IPK182 beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1Length: 876Number of Matches: 1

27:

TACGCAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAAC
ACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACC
AGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTA
AATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAG
CACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCCTACAGTACAGCGAT
AACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTC
GCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGT

BLAST output :Klebsiella pneumoniae strain E12KPMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds Sequence ID: <u>KY640528.1</u>Length: 640Number of

29

Matches: 1

GATGCCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAA

BLAST output: Escherichia coli strain IPK174 beta-lactamase **CTX-M-55** (blaCTX-M) gene, blaCTX-M-55 allele, complete cds

Sequence ID: MH900523.1Length: 876Number of Matches: 1

31:

TCGAGCCGGAACGTGTCATGCGGGCGTCAGGCTG

CCGTAATGGCGATTTGCGCCCGGACCAGGCCGCACGGGGAAACTCTGCGGCCTTTTTCGT
TCTTACTGCGGGTAAGGCACCCAGTCGCCGCCGTTCAGGCGAACGTACGGTTTATCCTGG
TATTGAATAACTACTGCATTTGAGTTCTCGGAGACCGGTGCTGTTTGTGGTAACCCACTG
GTGAGTTTTTCCAGTCAACATTGTCTTCGGTGAAAATCTTGCCATCAAGAACGCGAACC
ACCAGATCGGCAGGCTTGA

BLAST output: Escherichia coli strain MNCRE44, complete genome

Sequence ID: gb| CP010876.1|Length: 5010884Number of Matches: 3

33:

BLAST output: Salmonella enterica strain Sal1426 class A extended-spectrum beta-lactamase **CTX-M-9** (blaCTX-M) gene, blaCTX-M-9 allele, partial cds Sequence ID: MF797877.1Length: 867Number of Matches: 1

35:

ACGCAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTA
ACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCA
CCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGT
TAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAA
AGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCG
ATAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGT
TCG

BLAST output: Klebsiella pneumoniae strain KPTR1-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-115 allele, complete cds Sequence ID: MK113960.1 Length: 1008Number of Matches: 1

GCTGCTGGGAGCGCCGCTTTATGCGCAGACGAGTGCGGTGCAGCAAATGCTGGCGGCG CTGGAGATAAGC

BLAST output: Escherichia coli strain IPK37 beta-lactamase CTX-M-27 (blaCTX-

M) gene, blaCTX-M-27 allele, complete cds

Sequence ID: MH900525.1 Length: 876Number of Matches: 1

37:

GCTGGGCAGCGCCGCTTTATGCGCAGACGAGTGCGGTGCAGCAAAAGCTGGCGGCGCT GGAGAAAAGCAGCGGAGGGCGGCTGGGCGTCGCGCTCATCGATACCGCAGATAATACGCA GGTGCTTTATCGCGGTGATGAACGCTTTCCAATGTGCAGTACCAGTAAAGTTATGG

BLAST output : Escherichia coli strain YSP8-1 plasmid pYSP8-1-**CTX-M-14**, complete sequence Sequence ID: <u>CP037912.1</u>Length: 149304Number of Matches: 1

41:

ACCCCTGCTATTTAGCAGCGCAACGCTGCATGCGCAGGCGAACAGCGTGCAACAGCAGCTGGA
AGCCCTGGAGAAAAGTTCGGGAGGTCGGCTTGGCGTTGCGCTGATTAACACCGCCGATAATTCG
CAGATTCTCTACCGTGCCGATGAACGTTTTGCGATGTGCAGTACCAGTAAGGTGATGGCGGCCG
CGGCGGTGCTTAAACAGAGCGAGAGCGATAAGCACCTGCTAAATCAGCGCGTTGAAATCAAGA
AGAGCGACCTGGTTAACTACAATCCCATTGCGGAGAAACACGTTAACGGCACGATGACGCTGG
CTGAGCTTGGCGCAGCGGCGCTGCAGTATAGCGACAATACTGCCATGAATAAGCTGATTGCCCA
TCTGGGTGGCCCGATAAAGTGACGGCGTTTGCTCGCTCGTTGGGTGATGAGACCTTCCGTCTGG
ACAGAACCGAGCCCACGCTCAATACCGCCATTCCAGGCGACCCGCGTGATACCACCACGCCGCT
CGCGATGGCGCAGACCCTGAAAAATCTGACGCTGGGTAAAGCGCTGGCGGAAACTCAGCGGGC
ACAGTTGGTGACGTGGCTTAAGGGCAATACTACCGGTAGCGCGAGCATTCCGGCGGGTCTGCCG
AAATCATGGGTAGTGGGCGATAAAACCGGCAGCGGAGCATTATGGCACCACCAACGATATCGCG
GTTATCTGGC

BLAST output : Klebsiella pneumoniae strain KP72 extended-spectrum betalactamase **CTX-M-2** (blaCTX-M) gene, blaCTX-M-2 allele, partial cds Sequence ID: MH661245.1Length: 826Number of Matches: 1

43:

AACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGC ACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTG TTAAATCAGCGAGTTGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAA AAGCACGTCAATGGGA **BLAST output**: *Escherichia coli* strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence

ID: MH900522.1 Length: 876Number of Matches: 1

45:

BLAST output :Klebsiella pneumoniae strain KPTR1-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase CTX-M-15 (blaCTX-M) gene,

blaCTX-M-115 allele, complete cds Sequence ID: MK113960.1 Length: 1008Number of Matches: 1

47:

ATGGTGACAAAGAGAGTGCAACGGATGATGTTCGCGGCGGCGGCGTGCATTCCGCTGCTG
CTGGGCAGCGCGCGCTTTATGCGCAGACGAGTGCGGTGCAGCAAAAAGCTGCGGCGCT
GAGAAAAGCAGCGGAGGGCGACGCTTTCCAATGTGCAGTACCAGTAAAGTTATGGCGGCC
GCGGCGGTGCTTAAGCAGAGTGAAACGCAAAAGCAGCTGCTTAATCAGCCTGTCGAGATC
AAGCCTGCCGATCTGGTTAACTACAATCCGATTGCCGAAAAACACGTCAACGGCACAATG
ACGCTGGCAGAGCTGAGCGCGGCCGCGTTGCAGTACAGCGACAATACCGCCATGAACAA
TTGATTGCCCAGCTCGGTGGCCCGGGAGGCGTGACGGCTTTTGCCCGCGCGATCGGCGAT
GAGACGTTTCGTCTGGATCGCACTGAACCTACGCTGAATACCGCCATTCCCGGCGACCCG
AGAGACACCACCACGCCGCGGGCGATGGCACAGACGTTGCGTCAGCTTACGCTGGGTCA
GCGCTGGGCGAAACCCAGCGGGCGCAGTTGGTGACGTGGCTCAAAGGCAATACGACCGC
GCAGCCAGCATTCGGGCCGGCTTACCGACGTCGACTGAATAA

BLAST output: Enterobacter cloacae strain CERIC152-401 class A extended spectrum beta-lactamase **CTX-M-9** (CTX-M-9) gene, partial cds Sequence

ID: KT459753.1Length: 867Number of Matches: 2

TGAGGGCCGCGGCGGTGCTGAAAAAAAGTGAAAGCAAACCGAATCTGTTAAATCAGC GAGTTGAGATCAAAAAATGTG

BLAST output: *Escherichia coli* strain CR4 plasmid class A extended-spectrum beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence ID: MK405591.1Length: 876Number of Matches: 1

51:

BLAST output :Klebsiella pneumoniae partial **CTX-M-15** gene for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 327

Sequence ID: LT628520.1Length: 626Number of Matches: 1

54:

ATGGTGACAAAGAGAGTGCAACGGATGATGTTCGCGGCGGCGGCGTGCATTCCGCTGCTG
TGGGCAGCGCGCCGCTTTATGCGCAGACGAGTGCGGTGCAGCAAAAGCTGGCGGCGCTG
GAGAAAAGCAGCGGAGGGCGGCTGGGCGTCCGCTCATCGATACCGCAGATAATACGCAG
GTGCTTTATCGCGGTGATGAACGCTTTCCAATGTGCAGTACCAGTAAAGTTATGGCGGCC
GCGGCGGTGCTTAAGCAGAGTGAAACGCAAAAAGCAGCTGCTTAATCAGCCTGTCGAGATC
AAGCCTGCCGATCTGGTTAACTACAATCCGATTGCCGAAAAACACGTCAACGGCACAAT
AGAGACACCACCACGCCGGGGCGATGGCACAGACGTTGCGTCAACGCTTACGCTGGGTCAT
GCGCTGGGCGAAACCCAGCGGGCGCAGTTGGTGACGTGGCTCAAAGGCAATACGACCGGC

BLAST output: *Escherichia coli* strain IPK37 beta-lactamase **CTX-M-27** (blaCTX-M) gene, blaCTX-M-27 allele, complete cds

BLAST output: Escherichia coli strain BA22372 **plasmid pCTX-M-15**_22372, complete sequence Sequence ID: <u>CP040398.1</u>Length: 98470Number of Matches: 1

59:

BLAST output: Escherichia coli strain A19 plasmid beta-lactamase **CTX-M-1** (blaCTX-M) gene, blaCTX-M-CTX-M-1 allele, partial cds Sequence ID: MH037035.1Length: 810Number of Matches: 3

61:

ACGCAAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAAC
ACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACC

AGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTA
AATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAG
CACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGAT
AACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTC
GCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACC
GCCATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGG
AATCTGACGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATG
AAAGGCAATACCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTTCCTGGGTTGTG
GGGGATAAAACCGGCAGCGGTGGCTATGGACA

BLAST output: *Escherichia coli* strain E78ECMO **CTX-M-15** beta-lactamase (blaCTX-M-15) gene, partial cds Sequence ID: <u>KY640536.1</u>Length: 641Number of Matches: 1

63

TTGCCATCAAGAACGCGAACCACCAGATCGGCAGGCTTGA

BLAST output: *Escherichia coli* strain 4928STDY7071340 genome assembly, chromosome: 1 Sequence ID: LR607331.1Length: 5120867Number of Matches: 1

65:

BLAST output: Escherichia coli strain CH-15-5 extended spectrum beta-lactamase **CTX-M-55** (blaCTX-M-55) gene, complete cds; and hypothetical protein gene, partial cds

Sequence ID: MF958462.1Length: 980Number of Matches: 1

BLAST output : Escherichia coli strain F170 extended-spectrum beta-lactamase CTX-M-15 gene (CTX-M 1 group) , partial cds Sequence

ID: gb|KP325147.1|Length: 843Number of Matches: 1

69:

BLAST output: Escherichia coli strain 1125509 class A beta-lactamase (blaCTX-M-163) gene, complete cds Sequence ID: gb|KP681698.1|Length: 876Number of Matches: 1

71:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAAC

BLAST output : Klebsiella pneumoniae subsp. pneumoniae strain KPTR13-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence ID: MK113957.1Length: 1027Number of Matches: 1

73:

CTGCTTAATCAGCCTGTCGAGATCAAGCCTGCCGATCTGGTTAACTACAATCCGATTGCC
GAAAAACACGTCAACGGCACAATGACGCTGGCAGAACTGAGCGCGGCCGCGTTGCAGTAC
AGCGACAATACCGCCATGAACAAATTGATTGCCCAGCTCGGTGGCCCGGGAGGCGTGACG
GCTTTTGCCCGCGCGATCGGCGATGAGACGTTTCGTCTGGATCGCACTGAACCTACGCTG
AATACCGCCATTCCCGGCGACCCGAGAGACACCACCACGCCGCGGGCGATGGCGCAGACG
TTGCGTCAGCTTACGCTGGGTCATGCGCTGGGCGAAACCCAGCGGGCGCAGTTGGTGACG
TGGCTCAAAGGCAATACGACCGGCGCAGCCAGCATTCGGGCCGGCTTACCGACGTCGTGG
ACTGTGGG

BLAST output :Escherichia coli beta-lactamase CTX-M-16 gene, complete cds AY029068.1

75:

CGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTCCGAG CCGACGTTAAACACCGCCATTT

BLAST output: Escherichia coli strain F147 **extended-spectrum beta-lactamase CTX-M-15 gene**, partial cds . Sequence ID: gb|KP325146.1|Length: 822Number of Matches: 1

77:

BLAST output: Escherichia coli strain E78ECMO **CTX-M-15** beta-lactamase (blaCTX-M-15) gene, partial cds,Sequence ID: <u>KY640536.1</u>Length: 641Number of Matches **80**:

BLAST output: Salmonella enterica strain Sal1426 class A extended-spectrum beta-lactamase **CTX-M-9** (blaCTX-M) gene, blaCTX-M-9 allele, partial cds Sequence ID: MF797877.1Length: 867Number of Matches: 1

00	
XX	-
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ATGATGAGAAAAAGCGTAAGGCGGGCGATGTTAATGACGACAGCCTGTGTTTCGCTGCTG
TTGGCCAGTGTGCCGCTGTGTGCCCAGGCGAACGATGTTCAACAAAAGCTCGCGGCGCTG
GAGAAAAGCAGCGGGGGACGACTGGGTGTGCGTTGATTACACCGCCGATAACACGCAG
ACGCTCTACCGCGCCGACGAGCGTTTTGCCATGTGCAGCACCAGTAAAGTGATGGCGGTA
GCGGCGGTGCTTAAGCAAAGTGAAACGCAAAGGGCTTGTTGAGTCAGCGGGTTGAAATT
AAGCCCTCAGACTTGGTTAACTACAACCCCATTGCGGAAAAACACGTCAATGGCACGATG
GACACGTTCCGGCTCGATCGTACCGAGCCGACCCCG

BLAST output: Proteus mirabilis 1111641 blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-160**, complete CDS .Sequence

ID: NG 048945.1 Length: 876Number of Matches: 2

84:

BLAST output: Klebsiella pneumoniae partial CTX-M-15 gene for Class A Extended Spectrum beta-lactamase **CTX-M-15**, isolate 122. Sequence ID: LT628516.1Length: 622Number of Matches: 1

86:

GGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAACCGGCAGCGGTGGCTATGGCACCACCAACG ATATCGACGAT

BLAST output: Escherichia coli strain E74ECMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds.Sequence ID: KY640534.1 Length: 640Number of Matches: 1

88:

ACGCAAAACTTGCCGAATTAGAGCGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACA CAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAA AGTGATGGCCGCGGCGGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCG AGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAA

BLAST output :Escherichia coli 275 unnamed blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-225**, complete CDS.Sequence ID: NG_064720.1Length: 1048Number of Matches: 1

90:

AACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCA
GTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATC
AGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAC
CGCGTTCGCCCGACAGCTGGGGACGAAACGTTCCGTCTCGACCGTACACGTTGGCGGCCCGGCT
AGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAAACGTTCCTCCGCGTGATACCACTTCACC
TCGGGCAATGGCGCAAACTCTGCGTAATCTGACGCTGGGTAAAGCATTGGGTGACAGCCAACG
GGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCAGGCTGGACT
GCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGACTATGGCACCACCAACA

BLAST output : Escherichia coli strain RN 1 plasmid cefotaximase class A extended-spectrum beta-lactamase **CTX-M-1** (**blaCTX-M**) **gene**, blaCTX-M-1 allele, partial cds

Sequence ID: MG255315.1Length: 487Number of Matches: 3

91:

AACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCA
GTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATC
AGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAA
TGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCCTACAGTAC

BLAST output : Escherichia coli strain VRES-hospital6495320 genome assembly, plasmid: 1 .Sequence ID: <u>LR607054.1</u>Length: 111743Number of Matches: 1

93

BLAST output : Escherichia coli blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-172**, complete CDS

Sequence ID: NG_048957.1Length: 876Number of Matches: 1

95:

BLAST output : Klebsiella pneumoniae partial **CTX-M-15 gene** for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 122.Sequence ID: LT628516.1Length: 622Number of Matches: 1

97:

ATGGTGACAAAGAGAGTGCAACGGATGATGTTCGCGGCGGCGGCGTGCATTCCGCTGCTG

CTGGGCAGCGCGCCGCTTTATGCGCAGACGAGTGCGGTGCAGCAAAAGCTGGCGGCGCTG
GAGAAAAGCAGCGGAGGGCGTCACCGCGTCGCGCTCATCGATACCGCAGATAATACGCAG
GTGCTTTATCGCGGTGATGAACGCTTTCCAATGTGCAGTACCAGTAAAGTTATGGCGGCC
GCGGCGGTGCTTAAGCAGAGTGAAACGCAAAAGCAGCTGCTTAATCAGCCTGTCGAGATC
AAGCCTGCCGATCTGGTTAACTACAATCCGATTGCCGAAAAACACGTCAACGGCACAATG
ACGCTGGCAGAGCTGAGCGCGGCCGCGTTGCAGTACAGCGACAATACCGCCATGAACAA
TTGATTGCCCAGCTCGGTGGCCCGGGAGGCGTGACGGCTTTTGCCCGCGCGATCGGCGAT
GAGACGTTTCGTCTGGATCGCACTGAACCTACGCTGAATACCGCCATTCCCGGCGACCCG
AGAGACACCA

BLAST output : Salmonella enterica strain Sal1426 class A extended-spectrum beta-lactamase **CTX-M-9** (blaCTX-M) gene, blaCTX-M-9 allele, partial cds .Sequence ID: MF797877.1Length: 867Number of Matches: 1

98:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAAC
ACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACC
AGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTT
AAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAA
GCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGAT
AACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTC
GCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACC
GCCATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGG
AATCTGACGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATG
AAAGGCAATACCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTCCTGGGTTGTG
GGGGGATAAAACCGGCAGCGAGCGAGCTATGGCACAGC

Klebsiella pneumoniae subsp. pneumoniae strain KPTR13-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence ID: MK113957.1 Length: 1027Number of Matches: 1

100:

ACGGATGATGTTCGCGGCGGCGGCGTGCATTCCGCTGCTGCTGGGCAGCGCGCCGCTTTATGCG CAGACGAGTGCGGTGCAGCAAAAGCTGGCGGCGCTGGAGAAAAGCAGCGGAGGGCGGCTGGG CGTCGCGCTCATCGATACCGCAGATAATACGCAGGTGCTTTATCGCGGTGATGAACGCTTTCCA ATGTGCAGTACCAGTAAAGTTATGGCGGCCGCGGCGGTGCTTAAGCAGAGTGAAACGCAAAAG CAGCTGCTTAATCAGCCTGTCGAGATCAAGCCTGCCGATCTGGTTAACTACAATCCGATTGCCG
AAAAACACGTCAACGGCACAATGACGCTGGCAGAACTGAGCGCGGCGCGCGTTGCAGTACAGCG
ACAATACCGCCATGAACAATTGATTGCCCAGCTCGGTGGCCCGGGAGGCGTGACGGCTTTTGCC
CGCGCGATCGGCGATGAGACACTTCGTCTGGATCGCACTGAACCTACGCTGAATACCGCCATTC
CCGGCGACCCGAGAGACACCACCACCGCGGGCGATGGCGCAGACGTTGCGTCAGCTTACGC
TGGGTCATGCGCTGGGCGAAACCCAGCGGGCGCAGTTGGTGACG

Escherichia coli strain IPK37 beta-lactamase **CTX-M-27** (blaCTX-M) gene, blaCTX-M-27 allele, complete cds.Sequence ID: MH900525.1Length: 876Number of Matches: 1

Appendix II

Glan Clwyd Hospital samples DNA sequences

101:

TGCCGAATTAGAGCGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAGCAGA
TAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGT
GATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCG
AGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAA
TGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGGCTACAGTACAGCGATAACGTGGC
GATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACA
GCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCC
GGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGAC
GCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAA
TACCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTTCCTGGGTTGTGGGGGGATAA
AACCGGCAGCGAGCTATGGCACCACCACCACCACGATATCG

Blast result : Escherichia coli strain F170 extended-spectrum beta-lactamase CTX-M-15 gene (CTX-M 1 group) , partial cds. Sequence ID: gb|KP325147.1|

Length: 843Number of Matches: 1

Blast result: Escherichia coli strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cdsequence ID: MH900522.1 Length: 876Number of Matches: 1

106

CGACAATACTGCCATGAATAAGCTGATTGCCCATCTGGGTGGTCCCGATAAAGTGACAGC
GTTTGCTCGCTCGTTGGGTGATGAGACCTTCCGTCTGGACAGAACCGAGCCCACGCTCAATACC
GCCATTCCAGGCGACCCGCGTGATACCACCACGCCGCTCGCGATGGCGCAGACCCTGAAAAATC
TGACGCTGGGTAAAGCGATGGCGGAAACTCAGCGGGCACAGTTGGTGACGTGGCTTAAGGGCA
ATACTACCGGTAGCGCGAGCATTCGGGCGGGTCTGCCGAAATCATGGGTAGTGGGCGATAAAA
CCGGCAGCGGAGATTATGGCACCACCAACGA

BLAST output: Klebsiella pneumoniae strain KP10 extended-spectrum betalactamase **CTX-M-59** (blaCTX-M) gene, blaCTX-M-59 allele, partial cds Sequence ID: MH661247.1 Length: 843Number of Matches: 1

108:

GCGCTGGCGAAACTCAGCGGCACAGTTGGTGACATGGCTTAAGGGCAATATACCGGTAGCG CGAGCATTTGGGCGGGTCTGCCGAAATCATGGGTAGTGGGCGATAAAACCGGCAGCGGAGATT ATGGCACCACCAACGATATCGCGGTTATCTGGCCGGAAAACCACGC

ACCGCTGGTTCTGGTGACCTACTTTACCCA

BLAST output: Escherichia coli strain 42.120E1 CTX-M family extended-spectrum beta-lactamase (blaCTX-M) gene, partial cds

Sequence ID: MK896928.1 Length: 618Number of Matches: 1

109:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACA
GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG
TGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCAATACCACCGGTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGGCTATGGCACCACC
AACCGA

BLAST output: Klebsiella pneumoniae strain KPTR1-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase CTX-M-15 (blaCTX-M) gene, **blaCTX-M-115** allele, complete cds. Sequence ID: MK113960.1 Length: 1008Number of Matches: 1

111:

CGGTGCCGATGACCCCGTTACAGGAGTTCTGGCACTATTTTAAACGCAACAAAGGCGCGG TTGTCGGACTGGTTTACGTCGTCATCGTGCCGTTAACGACTGCC

BLAST output:Escherichia coli O1:H42 strain CLSC36 chromosome, complete genome.Sequence ID: <u>CP041300.1</u> Length: 5083072Number of Matches: 1

113:

CAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGC AGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTG CTTCCTGGGTTGTGGGGGATAAGACC

BLAST output:Escherichia coli strain IPK182 beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1Length: 876Number of Matches: 1

115:

CGATGTGCAGCACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAAGTGAAAGCGAAC
CGAATCTGTTAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGC
GGAAAAGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAG
CGATAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTC
GCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCC
ATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGA
CGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATA
CCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTTCCTGGGTTGTGGGGGGATAAAACCG
GCAGCGGTGGCTATGGCACCACAA

BLAST output: Escherichia coli strain CR4 plasmid class A extended-spectrum beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence ID: MK405591.1 Length: 876Number of Matches: 1

118:

BLAST output: Escherichia coli strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1 Length: 876Number of Matches: 1

120:

GCAAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAGC
AGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTG
ATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTT
GAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGGACGA
TGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAAGCT
GATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGAACCG
TTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACCA
CTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACAG
CCAACGGGCGCAGCTGGTGACATGGATGAAAAGCAATACCACCGGTGCAGCGAGCATTCAGGC
TGGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAACCGGCAGCGTGGCTATGGCACCACCACC
GATATC

BLAST output:Escherichia coli strain E78ECMO **CTX-M-15** beta-lactamase (blaCTX-M-15) gene, partial cds

Sequence ID: KY640536.1 Length: 641Number of Matches: 1

122:

BLAST output: Escherichia coli ESBL20160070 blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-203**, complete CDS

Sequence ID: NG_055269.1 Length: 876Number of Matches: 1

124:

GATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGGGTTCGCCCGACAGCTGGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATCCAGGCGACTGCTTCCTGGGTTGTGGGGGATAAAA

BLAST output :Proteus mirabilis PM341/03 pPM341/03 blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-66**, complete CDS

Sequence ID: NG_049017.1 Length: 1076Number of Matches: 1

128:

AATCAATCGTGCCGTTAACGAGATTAGCGCACCGGTGCCGATGACCCCGTTACAGGAGTT CTGGCACTATTCTAAACGCAACAAAGGCGCGGTTGTCGGGCTGGTTTACGTCGTCATCGT GCCGTTAACAAAAC

BLAST output :Escherichia coli strain 4928STDY7071340 genome assembly, chromosome: 1.Sequence ID: LR607331.1 Length: 5120867Number of Matches: 1

129:

BLAST output :Salmonella enterica strain Sal1426 class A extended-spectrum beta-lactamase CTX-M-9 (blaCTX-M) gene, **blaCTX-M-9** allele, partial cdsSequence ID: MF797877.1 Length: 867Number of Matches: 1

132:

TGGTGACATGGATGAAAGCCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTTC
CTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGGCTATGGCACCGCCC

BLAST output : Klebsiella pneumoniae **KL8-ctx-m-15 gene** cluster, complete sequence

Sequence ID: MH523447.1 Length: 20281Number of Matches:

235:

BLAST output : Escherichia coli strain IPK192 beta-lactamase CTX-M-32 (blaCTX-M) gene, **blaCTX-M-32** allele, complete cds

Sequence ID: MH900527.1 Length: 876Number of Matches: 3

136:

Escherichia coli strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1 Length: 876Number of Matches: 1

138:

TACGCAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACA GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG TGATGGCCGCGGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGT

BLAST output: Klebsiella pneumoniae strain E12KPMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds.Sequence ID: <u>KY640528.1</u> Length: 640Number of Matches: 1

141:

CAAAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAGC
AGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTG
ATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTT
GAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGGACGA
TGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAAGCT
GATTGCTCACGTTGGCGGGCCCGGCTACACGTCACCGCGTTCGCCCGACAGCTGGGAGACGAAACG
TTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACCA
CTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACAG
CCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCAGGC
TGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGGCTATGGCACCACCAAC
CGATATCGAGA

BLAST output: Escherichia coli strain BA22372 plasmid **pCTX-M-15**_22372, complete sequence. Sequence ID: <u>CP040398.1</u> Length: 98470Number of Matches: 1

142:

AGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAA
GTGATGGCCGCGCGCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGA
GTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCACCGCGT
TCGCCCGACAGCTGGGGACGAAACGTTCCGTCTCGACCGTACACGTTGGCGGCCCGGCTAGCGT
CACCGCGTTCGCCCGACAGCTGGGAGACGAAACGTTCCTCCGCGTGATACCACTTCACCTCGGG
CAATGGCGCAAACTCTGCGTAATCTGACGCTGGGTAAAGCATTGGGTGACAGCCAACGGGCGC
AGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTG
CTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGACTATGGCACCACCACA

BLAST output : Escherichia coli strain A19 plasmid beta-lactamase **CTX-M-1** (blaCTX-M) gene, blaCTX-M-CTX-M-1 allele, partial cds.Sequence ID: MH037035.1 Length: 810Number of Matches: 3

143:

GTCAGTTCACGCTGATGGCGACGGCAACCGTCACGCTGTTGTTAGGAAGTGTGCCGCTGTATGC
GCAAACGGCGGACGTACAGCAAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGG
TGTGGCATTGATTAACACACGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCG
ATGTGCAGCACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCG
AATCTGTTAAATCAGCGAGCTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGG
AAAAGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCG
ATAACGTAGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGC
CCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATT
CCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGC
TGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCAATACCA
CCGGTGCAGCGAGCATTCAGGCTGGACTCCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAG
CGGTGGCTATGGCACCACCAACGATATCGCGGTGATCTGGCCAAAAGATCGTGCGCCGCTGATT
CTGGTCACTTACTTCACCCAGCCTCAACCTAAGGCAGAAAGCCGTCGCGATGTATTAGCGTCGG
CGGCTAAAATCGTCA

BLAST output : Shigella sp. SH223 blaCTX-M-108 (CTX-M-108) gene, partial cds Sequence ID: <u>JF274245.1</u> Length: 864Number of Matches: 1

145:

ACGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACAA
GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG
TGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAACCGGCAGCGGTGCAGCGACCAAA

BLAST output : Klebsiella pneumoniae subsp. pneumoniae strain KPTR12-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence ID: MK113958.1 Length: 1027Number of Matches: 1

147:

BLAST output : Escherichia coli strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, **blaCTX-M-15** allele, complete cds.Sequence ID: MH900522.1 Length: 876Number of Matches: 1

150:

AGTGATGGCCGCGGCGGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCG
AGTTGAGATCACAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGG
ACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATA
AGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGA
AACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGAT
ACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCG
ACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCAATACCACCGGTGCAGCGAGCATTC
AGGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAGACCG

BLAST output: Escherichia coli strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence ID: MH900522.1 Length: 876Number of Matches: 1

151:

BLAST output : Escherichia coli strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence ID: MH900522.1 Length: 876Number of Matches: 1

157:

BLAST output: Escherichia coli strain Hb-9 plasmid pHb-9 insertion sequence ISEcp1, partial sequence; beta-lactamase CTX-M-65 (blaCTX-M-65) and truncated transposase of IS903D (tnpA) genes, complete cds; insertion sequence IS26, complete sequence; fosfomycin resistance protein (fosA3), hypothetical protein, CadC-like protein, and putative transcriptional regulator, TetR family genes, complete cds; and insertion sequence IS26, partial sequence Sequence ID: KX495605.1 Length: 5927Number of Matches: 1

158:

BLAST output :Escherichia coli pMSP520 blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-51**, complete CDS.Sequence ID: NG 049002.1 Length: 876Number of Matches: 1

159:

TCGAGCCGGAACGTGTCATGCGGGCGTCAGGCTGCCGTAATGGCGATTTGCGCCCGGACCAGGCCGCCGTCGCCGGGGAAACTCTGCGGCCTTTTTCGTTCTTACTGCGGGTAAGGCACCCAGTCGCCGCCGT

TCAGGCGAACGTACGGTTTATCCTGGTATTGAATAACTACTGCATTTGAGTTCTCGGAGACCGGT GCTGTTTGTGGTAACCCACTGGTGAGTTTTTTCCAGTCAACATTGTCTTCGGTGAAAATCTTGCC ATCAAGAACGCGAACCACCAGATCGGCAGGCTTGA

BLAST output: Escherichia coli strain MNCRE44, complete genome. Sequence

ID: gb| CP010876.1| Length: 5010884Number of Matches: 3

161:

BLAST output: Escherichia coli strain F170 extended-spectrum beta-lactamase CTX-M-15 gene (CTX-M 1 group), partial cds.Sequence ID: gb[KP325147.1]

Length: 843Number of Matches: 1

164:

BLAST output: Escherichia coli strain IPK182 beta-lactamase CTX-M-15

(blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1 Length: 876Number of Matches: 1

166:

TACGCAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACA
GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG
TGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGT

BLAST output: Klebsiella pneumoniae strain E12KPMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds

Sequence ID: KY640528.1 Length: 640Number of Matches: 1

168:

BLAST output: Klebsiella pneumoniae strain KP10 extended-spectrum betalactamase CTX-M-59 (blaCTX-M) gene, blaCTX-M-59 allele, partial cds.Sequence ID: MH661247.1 Length: 843Number of Matches: 1

172:

BLAST output: Escherichia coli strain BA22372 plasmid **pCTX-M-15**_22372, complete sequence.Sequence ID: <u>CP040398.1</u> Length: 98470Number of Matches: 1

174:

BLAST output: Escherichia coli blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-172**, complete CDS.Sequence

ID: NG_048957.1Length: 876Number of Matches: 1

175:

BLAST output: Escherichia coli strain IPK182 beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence

ID: MH900522.1 Length: 876Number of Matches: 1

177:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACA GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG TGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG TTGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCCTACAGTACAGCGATAACGTGGCGATGAATAGC
TGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAAAC
GTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACC
ACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACA
GCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCA

BLAST output: Klebsiella pneumoniae subsp. pneumoniae strain KPTR7239-17 insertion sequence ISEcp1, partial sequence; and beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence ID: MK113956.1 Length: 1026Number of Matches:

179:

BLAST output: Klebsiella pneumoniae isolate KSH203 plasmid pKSH203-**CTX-M-3**, complete sequence.Sequence ID: <u>CP034325.1</u>Length: 156910Number of Matches: 1

181:

BLAST output: Klebsiella pneumoniae strain E161KPMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds.Sequence ID: KY640551.1 Length: 632Number of Matches: 1

184:

TCACTGAGCCAGTTCACGCTGATGGCGACGGCAACCGTCACGCTGTTGTTAGGAAGTGTGCCGC
TGTATGCGCAAACGGCGGACGTACAGCAAAAACTTGCCGAATTAAGCGGCAGTGTCGGGAGGC
AGACTGGGTGTGGCATTGATTAACACAGCAGATAATTCGCAATACTTTATCGTGCTGATGAGCG
CTTTGCGATGTGCAGCACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAAGTGAAAG
CGAACCGAATCTGTTAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCG
ATTGCGGAAAAGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGT
ACAGCGATAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGC
GTTCGCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGAACTCTGCGGAATC
TGACGCTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCA
ATACCACCGGTGCAGCGAGCATTCAGGCTGG

BLAST output : Klebsiella oxytoca 1125476 blaCTX-M gene for class A extended-spectrum beta-lactamase **CTX-M-162**, complete CDS.Sequence ID: NG_048947.1 Length: 876Number of Matches: 1

185:

BLAST output: Escherichia coli strain F170 extended-spectrum beta-lactamase CTX-M-15 gene (CTX-M 1 group), partial cds.Sequence ID: gb| KP325147.1|

4.0=

Length: 843Number of Matches: 1

187:

CAAAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAGC
AGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTG
ATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTT
GAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGACGA
TGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAAGCT
GATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACCGAAACG
TTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCACCATTCCGGGCGATCCGCGTGATACCA
CTTCACCTCGGGCAATGGCGCAAACTGTGCGGAATCTGACGCTGGGTAAAGCATTCGGCCACAG
CCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCAGGC
TGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGTGGCTATGGCACCACCA

BLAST output : Klebsiella pneumoniae KL8-ctx-m-15 gene cluster, complete sequence

Sequence ID: MH523447.1 Length: 20281Number of Matches: 1

190:

BLAST output : Klebsiella pneumoniae partial CTX-M-15 gene for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 243.Sequence ID: <u>LT628518.1</u> Length: 626Number of Matches: 1

193:

CGCCAGTTCACGCTGATGGCGACGGCAACCGTCACGCTGTTGTTAGGAAGTGTGCCGCTGTATG
CGCAAACGGCGGACGTACAGCAAAAACTTGCCGAATTAGAGCGCAGTCGGGAGGCAGACTGG
GTGTGGCATTGATTAACACAGCAGATAATTCGCAAATACTTTATCGCGCTGATGAGCGCTTTGC
GATGTGCAGCACCAGTAAAGTGATGGCTGTGGCCGCGGTGCTGAAGAAAAAGTGAAAGCGAACC
GAATCTGTTAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCG
GAAAAGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGC
GATAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCG
CCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCAT
TCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAAACTCTGCGGAATCTGACG

CTGGGTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACC
ACCGGTGCAGCGAGCATTCAGGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAACCGGCA
GCGGTGGCTATGGCACCACCAACGATATCGCGGTGATCTGGCCAAAAGATCGTGCGCCGCTGAT
TCTGGTCACTTACTTCACCCAGCCTCAACC

BLAST output : Escherichia coli strain IPK174 beta-lactamase CTX-M-55 (blaCTX-M) gene, blaCTX-M-55 allele, complete cds

Sequence ID: MH900523.1 Length: 876Number of Matches: 1

195:

CTCAATGTTAACGGTGATGGCGACGCTACCCCTGCTATTTAGCAGCGCAACGCTGCATGCGCAG
GCGAACAGCGTGCAACAGCAGCTGGAAGCCCTGGAGAAAAGTTCGGGAGGTCGGCTTGGCGTT
GCGCTGATTAACACCGCCGATAATTCGCAGATTCTCTACCGTGCCGATGAACGTTTTGCGATGTG
CAGTACCAGTAAGGTGATGGCGGCCGCGGGGGGTGCTTAAACAGAGCGAGAGCGATAAGCACCT
GCTAAATCAGCGCGTTGAAATCAAGAAGAGCGACCTGGTTAACTACAATCCCATTGCGGAGAA
ACACGTTAACGGCACGATGACGCTGGCTGAGCTTGGCGCAGCGGCGCTTCAGTATAGCGACAAT
ACTGCCATGAATAAGCTGATTGCCCATCTGGGTGGTC

BLAST output : Klebsiella pneumoniae strain KP75 extended-spectrum betalactamase **CTX-M-2** (blaCTX-M) gene, blaCTX-M-2 allele, partial cds

Sequence ID: MH661246.1 Length: 829Number of Matches: 1

198:

GTGCAACGGATGATGTTCGCGGCGGCGGCGTGCATTCCGCTGCTGCTGCAGCAGCGCGCCGCTTT
ATGCGCAGACGAGTGCGGTGCAGCAAAAGCTGGCGGCGCGCTGGAGAAAAGCAGCGGAGGGCGG
CTGGGCGTCGCGCTCATCGATACCGCAGATAATACGCAGGTGCTTTATCACGGTGATGAACGCT
TTCCAATGTGCAGTACCAGTAAAGTTATGGCGGCCGCGGCGGTGCTTAAGCAGAGTGAAACGCA
AAAGCAGCTGCTTAATCAGCCTGTCGAGATCAAGCCTGCCGATCTGGTTAACTACAATCCGATT
GCCGAAAAACACGTCAACGGCACAATGACGCTGGCAGAACTGAGCGCGGCCGCGTTGCAGTAC
AGCGACAATACCGCCATGAACAAATTGATTGCCCAGCTCGGTGGCCCGGGAGGCGTGACGGCT
TTTGCCCGCGCGATCGGCGATGAGACACCACCACGCCGCGGGCGATGGCGCAGACCTTGCGTCAGC
CCATTCCCGGCGACCCGAGAGACACCACCACCACGCCGCGGGCGAACCTTGCGTCAACC
TTACGCTGGGTCATGCGCTGGGCGAAACCCACCACGCCGGGCGCAGTTGGTGACCTCAAAGGCA
ATACGACCGGCGCAGCCAGCATTCGGGCCGGCTTACCGACGTCGTGGACTGTGGTGATAAGA
CCGGCAGCGGCGCACCCACCACCACCACCACCACGACGTCGTGGACTGTGGGTGATAAGA

BLAST output :: Escherichia coli strain IPK37 beta-lactamase CTX-M-27 (blaCTX-M) gene, blaCTX-M-27 allele, complete cds

Sequence ID: MH900525.1 Length: 876Number of Matches: 1

200:

CGCAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACAC
AGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAA
GTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGA
GTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGGA
CGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAAGCATCCACCGGTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAACCGGCAGCGGTGCAGCGACCCA

BLAST output : Klebsiella pneumoniae strain KPCSQ15a extended spectrum beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, partial cds Sequence ID: MH891569.1 Length: 731Number of Matches: 1

Appendix III

YG Hospital samples DNA sequences.

201:

CGATGTGCAGCACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAAGTGAAAGCGAAC
CGAATCTGTTAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGC
GGAAAAGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAG
CGATAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTC
GCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCC
ATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGA
CGCTGGAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACC
ACCGGTGCAGCGAGCATCAGGCTGGACTGCCTG

CTTCCTGGGTTGTGGGGGATAAAACCGGCAGCGGTGGCTATGGCACCACCAACGATATCG

BLAST output: Escherichia coli strain IPK182 beta-lactamase CTX-M-15

(blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1Length: 876Number of Matches: 1

202

BLAST output : Salmonella enterica subsp. enterica serovar Manhattan plasmid **blaCTX-M-2** gene for beta-lactamase CTX-M-2, complete cds, strain: 3377

Sequence ID: LC229068.1Length: 876Number of Matches: 1

205:

BLAST output : Escherichia coli strain EP174a beta-lactamase (blaCTX-M15) gene, partial cds

Sequence ID: MF346615.1Length: 871Number of Matches: 1

207:

AGTTCACGCTGATGGCGACGGCAACCGTCAGCTGTTGTTAGGAAGTGTGCCGCTGTATGCGCAA ACGGCGGACGTACAGCAAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTG GCATTGATTAACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGT GCAGCACCAGTAAAGTGATGGCCGCGGCCGGGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATC
TGTTAAATCAGCGAGCTGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAG
CACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAAC
GTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGAC
AGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGG
CGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGT
AAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGAAAGGCAATACCACCGGT
GCAGCGAGCATTCAGGCTGGACTGCTTCCTGGGTTGTGGGGGATAAAA

BLAST output: Shigella sp. SH223 blaCTX-M-108 (CTX-M-108) gene, partial cds Sequence ID: <u>JF274245.1</u>Length: 864Number of Matches: 1

209:

AACGTGTCATGCGGGCGTCAGGCTGCCGTAATGGCAATTTGCGCCCGGACCAGGCCGCAGGGG GAAACTCTGCGGCCTTTTTCGTTCTTACTGCGGGTAAGGCACCCAGTCGCCGCCGTTCAGGCGA ACGTACGGTTTATCCTGGTATTGAATAACTACTGCATTTGAGTTCTCGGAGACCGGTGCTGTTTG TGGTAACCCACTGGTGAGTTTTTTCCAGTCAACATTGTCTTCGGTGAAAATCTTGCCATCAAGAA CGCGAACCACCAGATCGG

BLAST output : Escherichia coli strain 3385 chromosome, complete genome Sequence ID: CP029420.1Length: 4910422Number of Matches: 3

211:

AGTTCACGCTGATGGCGACGGCAACCGTCACGCTGTTGTTAGGAAGTGTGCCGCTGTATGCGCA
AACGGCGGACGTACAGCAGAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGT
GGCATTGATTAACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATG
TGCAGCACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAAT
CTGTTAAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAA
AGCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATA
ACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCG
ACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCG
GGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGG
GTAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGAAAGGCAATACCACCG
GTGCAGCGAGCATCCAGGCTGGACTGCCTGCTTCCTGGGTTGT

BLAST output :Escherichia coli strain EP174a beta-lactamase (blaCTX-M15) gene, partial cds Sequence ID: MF346615.1Length: 871Number of Matches: 1

215:

AGCAAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACA
CAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAA
AGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCG
AGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGG
ACGATGTCACTGGCTGAGCTTAGCGCGGCCGCCTACAGTACAGCGATAACGTGGCGATGAATA
ACCGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCAATACCACCGGTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAACCGGC

Escherichia coli blaCTX-M gene for class A extended-spectrum beta-lactamase CTX-M-172, complete CDS Sequence ID: NG_048957.1Length: 876Number of Matches: 1

217:

ACGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAA
CACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCAC
CAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTT
AAATCAGCGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAA
GCACGTCAATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGA
TAACGTGGCGATGAATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTT
CGCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACAC
CGCCATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCG
GAATCTGACGCTGGGTAAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGAT
GACAGGGT

BLAST output : Escherichia coli strain BA22372 plasmid pCTX-M-15_22372, complete sequence

Sequence ID: CP040398.1Length: 98470Number of Matches: 1

219:

GGATGGCTCGAGTTTTCAGCAAGATTTAGGAAATGTGCCGCTGTATGCGCAAACGGCGGACGT
ACAGCAAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAA
CACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGT
AAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAG
CGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATG
GGACGATGTCACTGGCTGAGCTTAGCGCGGCATCTTTCTAGAAGATCTCCTACAATATTCTCAGC

BLAST output :Escherichia coli strain CR4 plasmid class A extended-spectrum beta-lactamase CTX-M-**15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence ID: MK405591.1Length: 876Number of Matches: 1

223:

BLAST output :Klebsiella pneumoniae partial CTX-M-15 gene for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 243 Sequence ID: <u>LT628518.1</u>Length: 626Number of Matches: 1

226:

BLAST output :Escherichia coli strain SCEC020022 plasmid pCTXM14_020022, complete sequence.Sequence ID: CP032888.1Length: 109553Number of Matches: 1

228:

AACACGTCAACGGCACAATGACGCTGGCAGAACTGAGCGCGGCCGCGTTGCAGTACAGCGACA
ATACCGCCATGAACAAATTGATTGCCCAGCTCGGTGGCCCGGGAGGCGTGACGGCTTTTGCCCG
CGCGATCGGCGATGAGACGTTTCGTCTGGATCGCACTGAACCTACGCTGAATCCGCCATTCCCG
GCGACCCGAGAGACACCACCACCGCGGGGCGATGGCGCAGACGTTGCGTCAGCTTACGCTGG
GTCATGCGCTGGGCGAAACCCAGCGGGCGCAGTTGGTGACGTGGCCAAAGGCAATACGACCGG
CGCAGCCAGCATTCGGGCCGGCTTACCGACGTCGTGGACTG

BLAST output :Escherichia coli strain IPK37 beta-lactamase CTX-M-27 (blaCTX-M) gene, blaCTX-M-27 allele, complete cds.Sequence

ID: MH900525.1 Length: 876Number of Matches: 1

229:

GCATATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTGATGGC
CGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTTGA
GATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAA
TAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGG
AGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGA
TCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGG
TAAAGCATTGGGCGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCCAATACCAC
CGGTGCAGCGAGCATTCAGGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGG
CAGCGGTGGCTATGGCACCCAA

BLAST output :Klebsiella pneumoniae partial CTX-M-15 gene for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 243

Sequence ID: LT628518.1Length: 626Number of Matches: 1

231:

BLAST output :Klebsiella pneumoniae subsp. pneumoniae strain KPTR2-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MK113959.1Length: 1027Number of Matches: 1

238:

CGCAAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAG
CAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGT
GATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGT
TGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGACG
ATGTCACTGGCTGAGCTTAGCGCGGCCGCCTACAGTACAGCGATAACGTGGCGATGAATAAGC
TGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACCGAAAC
GTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACC
ACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTCAGG
CCCAACGGGCGCAGCTGGTGACATGGATGAAAGCCACCGCCA
CTGGACTGCTTCCTGGGTTGTGGGGGGATAAAACCCGCCA

BLAST output :Klebsiella pneumoniae partial **CTX-M-15** gene for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 327 .Sequence ID: <u>LT628520.1</u>Length: 626Number of Matches: 1

241:

TCACCGCGTTCGCCCGACAGCTGGGAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTT

AAACACCGCCATTCCGGGCGATCCGCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTG

CGTAATCTGACGCTGGGTAAAGCATTGGGTGACAGCCAACGGGCGCAGCTGGTGACATGGATG

AAAGGCA

BLAST output :Escherichia coli strain IPK192 beta-lactamase CTX-M-32 (blaCTX-M) gene, blaCTX-M-32 allele, complete cds.Sequence

ID: MH900527.1 Length: 876Number of Matches: 1

242:

TAAAGTGATGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCA
GCGAGTTGAGATCACAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAAT
GGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCCGCTACAGTACAGCGATAACGTGGCGATG
AATAAGCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAG
ACGAAACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCG
TGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTG
GGCGACAGCCAACGGGCGCAGCTGGTGACATGGAAAGCAATACCACCGGTGCAGCGAGC
ATTCAGGCTGGACTGCCTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGGCTATGGCA
CCCCACCA

BLAST output :Klebsiella pneumoniae partial **CTX-M-15** gene for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 243

Sequence ID: LT628518.1Length: 626Number of Matches: 1

244:

BLAST output : Escherichia coli class A extended-spectrum beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence ID: MK234851.1 Length: 877Number of Matches: 1

246:

BLAST output :Escherichia coli strain SCEC020022 plasmid pCTXM14_020022, complete sequence.Sequence ID: CP032888.1Length: 109553Number of Matches: 1

248:

AACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGC
ACCAGTAAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTG
TTAAATCAGCGAGTTGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAA
AAGCACGTCAATGGGA

Blast result : Escherichia coli strain IPK182 beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1 Length: 876Number of Matches: 1

250:

GGATGGCTCGAGTTTTCAGCAAGATTTAGGAAATGTGCCGCTGTATGCGCAAACGGCGGACGT
ACAGCAAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAA
CACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGT
AAAGTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAG
CGAGTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATG

Blast result : Escherichia coli strain CR4 plasmid class A extended-spectrum beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence ID: MK405591.1Length: 876Number of Matches: 1

252:

Blast result : Escherichia coli strain IPK118 beta-lactamase CTX-M-2 (blaCTX-M) gene, blaCTX-M-2 allele, partial cds.Sequence ID: MH900526.1Length: 793Number of Matches: 1

254:

AGATGCCGTGATTAGCGCTCCGGTGCCGATGACCCCGTTACAGGAGTTCTGGCACTATTT
TAAACGCAACACAGGCGCGGTTGTCGGGCTGGTTTACGTCGTCATCGTGCCGTTTACAAG
GCACCACCACCACGAC

Blast result : Escherichia coli strain 4928STDY7071340 genome assembly, chromosome: 1.Sequence ID: LR607331.1Length: 5120867Number of Matches: 1

257:

TCAGCGCTCAGCCCTTCGGCGATGATTCTCGCCGCTGAAGCCAGCACATCGCGGCGGCTCTCTG
CGTTCTGTTGCGGCTGGGTAAAATAGGTCACCAGAACCAGCGGCGCACGACCCTGCGGCCAGAT
CACCGCAATATCATTGGTGGTGCCGTAGTCGCCGCTGCCGGTCTTATCACCTACAGTCCACGAC

GTCGGTAAGCCGGCCCGAATGCTCGCTGCGCCGGTCGTATTGCCTTTGAGCCACGTCACCAACT
GCGCCCGCTGGGTTTCGCCCAGCGCATAACCCAGCGTAAGCTGACGCAACGTCTGTGCCATCGC
CCGCGGCGTGGTGGTCTCTCGGGTCGCCGGGAATGGCGGTATTCAGCGTAGGTTCAGTGCGA
TCCAGACGAAACGTCTCATCGCCGATCGCGCGGGCAAAAGCCGTCACGCCTCCCGGGCCACCG
AGCTGGGCAATCAATTTGTTCATGGCGGTATTGTCGCTGTACTGCAACGCGGCCGCGCTCAGCT
CTGCCAGCGTCATTGTGCCGTTGACGTGTTTTTCGGCAATCGGATTGTAGTTAACCAGATCGGCA
GGCTTGATCTCGACAGGCTGATTAAGCAGCTGCTTTTGCGTTTCACTCTGCTTAAGCACCGCCGC
GGCCGCCATAACTTTACTGGTACTGCACATTGGAAAGCGTTCATCACCGCGATAAAGCACCTGC
GTATTATCTGCGGTATCGATGAGCGCGACGCCCAGCCGCCCTCCGCTGCTTTTCTCCAGCGCCGC
CAGCTTTTGCTGCACCGCACTCGTCTGCGCATAAAGCGGCGCGCTGCACAGCAGCAGCAGCAGAATG
CACGCCGCCGCCGCGCAACATCATCCGTTGCACTCTCTTTTGTCACCATGGCCTGAGACCATTATGC
CAT

BLAST output :Escherichia coli strain SCEC020022 plasmid pCTXM14_020022, complete sequence.Sequence ID: CP032888.1Length: 109553Number of Matches: 1

259:

Blast result : Klebsiella pneumoniae partial CTX-M-15 gene for Class A Extended Spectrum beta-lactamase CTX-M-15, isolate 122Sequence ID: LT628516.1Length: 622Number of Matches: 1

262:

GCGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACAC
AGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAA
GTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGA
GTTGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGA
CGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA

CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCA GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAAACGGGCAGCGGTGGCTATGGCACCAC AAC

Blast result: Klebsiella pneumoniae strain KPTR1-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-115 allele, complete cds Sequence ID: MK113960.1Length: 1008Number of Matches: 1

264:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACA
GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG
TGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGTGGC

Blast result :Escherichia coli strain CR4 plasmid class A extended-spectrum betalactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence ID: MK405591.1Length: 876Number of Matches: 1

266:

Blast result :Escherichia coli strain MIAE02105 extended-spectrum class A beta lactamase CTX-M-15 (bla CTX-M15) gene, partial cds.Sequence

ID: MF977517.1 Length: 867Number of Matches: 1

268:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACA
GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG
TGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGGC

Blast result: Escherichia coli strain CR4 plasmid class A extended-spectrum betalactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-15 allele, complete cds Sequence ID: MK405591.1 Length: 876Number of Matches: 1

271:

GCAAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAGC
AGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTG
ATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTT
GAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGACGA
TGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAAGCT
GATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACCGAAACG
TTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACCA
CTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACAG
CCAACGGGCGCAGCTGGTGACATGGATGAAAAGCAATACCACCGGTGCAGCGAGCATTCAGGC
TGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGGCTATGGCACCACCAAC
GATATCGACGAT

Blast result : Escherichia coli strain E74ECMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds.Sequence ID: <u>KY640534.1</u>Length: 640Number of Matches: 1

272 (lower band):

TAGTTAGGCTCTTCAACCCAGTCGCCCTCTGTTGCCCACCAGTGATTAAATTCGTCGTGA
TTTGCTGAAACTGCCATACCCTATCGCCTGTCGTTTTTTTATTAACGACAATGACTATAGA
TGTTTAGCTGAGGAAAATCTTAATAATACGTGTTGTATTGACGAGTATCTTATGCCGGGA
ACAAGTCATCTCG

Blast result: Escherichia coli strain 4928STDY7071340 genome assembly,

chromosome: 1

Sequence ID: LR607331.1Length: 5120867Number of Matches: 1

273:

Blast result :Escherichia coli strain IPK182 beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds

Sequence ID: MH900522.1Length: 876Number of Matches: 1

275:

AGGGCGGCTGGGCGTCATCGATACCGCAGATAATACGCAGGTGCTTTATCGCGGTGAT
GAACGCTTTCCAATGTGCAGTACCAGTAAAGTTATGGCGGTCGCGGCGGTGCTTAAGCAGAGTG
AAACGCAAAAGCAGCTGCTTAATCAGCCTGTCGAGATCAAGCCTGCCGATCTGGTTAACTACAA
TCCGATTGCAGAAAAACACGTCAACGGCACAATGACGCTGGCAGAACTGAGCGCGGCCGCGTT
GCAGTACAGCGACAATACCGCCATGAACAAATTGATTGCCCAGCTCGGTGGCCCGGGAGGCGT
GACGGCTTTTGCCCGCGCGATCGGCGATGAGACATTCGTCTGGATCGCACTGAACCTACGCTG
AATACCTCCATTCCCGGCGACCCGAGAGACACCACCACGCCGCGGGCGATGGCGCAGACGTTG
CGTCAGCTTACGCTGGGTCATGCGCTGGGCGAAACCCAGCGGGCGCAGTTGGTGACGTCGAAGCTCA
AAGGCAATACGACCGCGCGCGCGCGCGCGCGCGCCGCTTACCGACGTCGTGG

Salmonella enterica subsp. enterica serovar California strain GZ680 extendedspectrum beta-lactamase **CTX-M-90** (blaCTX-M-90) gene, partial cds

Sequence ID: MF418175.1Length: 823Number of Matches: 1

277:

BLAST output : Escherichia coli 1125509 blaCTX-M gene for class A extendedspectrum beta-lactamase CTX-M-163, complete CDS.Sequence

ID: NG_048948.1Length: 876Number of Matches: 1

279:

Blast result : Escherichia coli strain HG16 extended spectrum beta-lactamase CTX-M-15 (blaCTX-M-15) gene, partial cds.Sequence

ID: gb|KR338941.1|Length: 867Number of Matches: 1

281:

GCGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACAC
AGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAA
GTGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGA
GTTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAAGCACGTCAATGGGA
CGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAAGCAATACCACCGGTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAAAACCGGCCAGCGTGGCTATGGCACCAC
AAC

Blast result: Klebsiella pneumoniae strain KPTR1-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase CTX-M-15 (blaCTX-M) gene, blaCTX-M-115 allele, complete cds.Sequence ID: MK113960.1Length: 1008Number of Matches: 1

283:

BLAST output :Escherichia coli strain SCEC020022 plasmid pCTXM14_020022, complete sequence.Sequence ID: CP032888.1Length: 109553Number of Matches: 1

286:

CGCAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACAA GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG TGATGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCCACCGCTGCTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGGCTATGGCACCA

Blast result : Klebsiella pneumoniae strain KPCSQ15a extended spectrum betalactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, partial cds Sequence ID: MH891569.1Length: 731Number of Matches: 1

288:

GCAAACTTGCCGATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACAGC
AGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAGTG
ATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAGTT
GAGATCAAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGACGA
TGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAAGCT
GATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACCGAAACG
TTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATACCA
CTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGACAG
CCAACGGGCGCAGCTGGTGACATGGATGAAAAGCAATACCACCGGTGCAGCGAGCATTCAGGC
TGGACTGCCTGCTTCCTGGGTTGTGGGGGGATAAAAACCGGCAGCGGTGGCTATGGCACCACCAAC
GATATCGACGAT

Blast result : Escherichia coli strain E74ECMO beta-lactamase **CTX-M-15** (blaCTX-M-15) gene, partial cds.Sequence ID: KY640534.1 Length: 640Number of Matches: 1

290:

CGCAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGCAGACTGGGTGTGGCATTGATTAACACA
GCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCACCAGTAAAG
TGATGGCCGCGGCCGCGGTGCTGAAGAAAAGTGAAAGCGAACCGAATCTGTTAAATCAGCGAG
TTGAGATCAAAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGAC
GATGTCACTGGCTGAGCTTAGCGCGGCCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA

CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGGCAATACCACCGGTGCAGCGAGCATTCA GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGGGATAAAACCGGCAGCGGTGGCTATGGCACAGC

Blast result: Klebsiella pneumoniae subsp. pneumoniae strain KPTR13-18 insertion sequence ISEcp1, partial sequence; and beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence

ID: MK113957.1Length: 1027Number of Matches: 1

292:

Blast result : Escherichia coli strain HG16 extended spectrum beta-lactamase CTX-M-15 (blaCTX-M-15) gene, partial cds.Sequence

ID: gb|KR338941.1|Length: 867Number of Matches: 1

293:

ACAAAGAGAGTGCAACGGATGATGTTCGCGGCGGCGGCGTGCATTCCGCTGCTGCTGGGCAGC
GCGCCGCTTTATGCGCAGACGAGTGCGGTGCAGCAAAAGCTGGCGGCGCTGGAGAAAAGCAGC
GGAGGGCGGCTGGGCGTCGGCTCATCGATACCGCAGATAATACGCAGTGCTTTATCGCGGTGAT
GAACGCTTTCCAATGTGCAGTACCAGTAAAGTTATGGCGGCCGCGGGGGGTGCTTAGCAGAGGAA
ACGCAAAAGCAGCTGCTTAATCAGCCTGTCGAGATCAAGCTGCCGATCTGGTTAACTACAATCC
GATTGCCGAAAAACACGTCAACGGCACAATGACGCTGGCAGAGCTGAGCGCGGCCGCGTTGCA
GTACAGCGACAATACCGCCATGAACAAATTGATTGCCCAGCTCGGTGGCCCGGGAGGCGTGAC
GGCTTTTGCCCGCGCGATCGGCGATGAGACATTCGTCTTGGATCGCACTGAACCTACGCTGAAT
ACCGCCATTCCCGGCGACCCGAGAGACACCACCACCACGCGGGGCGATGGCACAGACGTTGCGT
CAGCTTACGCTGGGTCATGCGCTGGGCGAAACCCACCACGCGGGCGCAGTTGGTGACGTGGCTCAAA
GGCAATACGACCGGCGCAGCCAGCATTCGGGCCGGCTTACCGACGTCGTGGACTGCAGGTGAT
AAGGCGACTACGGCACCACCACCACCACCACGCCGCGCGCTCGTGGACTGCAGGTGAT

BLAST output: Enterobacter hormaechei strain WCHEH020038 plasmid pCTXM9_020038, complete sequence >Sequence ID: CP031724.1 Length: 296580Number of Matches: 1

294:

BLAST output :Escherichia coli strain SCEC020022 plasmid pCTXM14_020022, complete sequence.Sequence ID: CP032888.1Length: 109553Number of Matches: 1

296:

BLAST output: Escherichia coli strain E78ECMO **CTX-M-15** beta-lactamase (blaCTX-M-15) gene, partial cds.Sequence ID: KY640536.1 Length: 641Number of Matches: 1

298:

ACGTACAGCAAAAACTTGCCGAATTAGAGCGGCAGTCGGGAGGAAGACTGGGTGTGGCATTGA
TTAACACAGCAGATAATTCGCAAATACTTTATCGTGCTGATGAGCGCTTTGCGATGTGCAGCAC
CAGTAAAGTGATGGCCGTGGCCGCGGTGCTGAAGAAAAAGTGAAAGCGAACCGAATCTGTTAAA
TCAGCGAGTTGAGATCAAAAAAATCTGACTTGGTTAACTATAATCCGATTGCGGAAAAGCACGTC
GATGGGACGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCG
ATGAATAAGCTGATTTCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGG
GAGACGAAACGTTCCGTCTCGACCGTACCGAGCCGATGTTAAACACCGCCATTCCGGGCGATCC
GCGTGATACCACTTCACCTCGGGCAATGGCGCAAACTCTGCGTAATCTGACGCTGGGTAAAGCA
TTGGGTGACAGCCAACGGGCGCAGCTGGTGACATGGATGAAAAGCCAATACCACCGGTGCAGCG
AGCATTCAGGCTGGACTGCCTTCCTGGGTTGTGGGGGGATAAAACCGGCAGCGGTGACTATG
GCACCACCAACGATATCGCGGTGATCTGGCCAAAAGATCGTGCTCCGCTGATTCTGGTCACTTA
CTTCACCCAGCCTCAACCTAAGGCAGAAAGCCGTCGC

BLAST output: Escherichia coli strain A19 plasmid beta-lactamase **CTX-M-1** (blaCTX-M) gene, blaCTX-M-CTX-M-1 allele, partial cds

Sequence ID: MH037035.1Length: 810Number of Matches: 1

300:

TGATGCCGCGGCCGCGGTGCTGAAGAAAAGATGAAAGCGAACCGAATCTGTTAAATCAGCGA
GTTGAGATCACAAAATCTGACCTTGTTAACTATAATCCGATTGCGGAAAAGCACGTCAATGGGA
CGATGTCACTGGCTGAGCTTAGCGCGGCCGCGCTACAGTACAGCGATAACGTGGCGATGAATAA
GCTGATTGCTCACGTTGGCGGCCCGGCTAGCGTCACCGCGTTCGCCCGACAGCTGGGAGACGAA
ACGTTCCGTCTCGACCGTACCGAGCCGACGTTAAACACCGCCATTCCGGGCGATCCGCGTGATA
CCACTTCACCTCGGGCAATGGCGCAAACTCTGCGGAATCTGACGCTGGGTAAAGCATTGGGCGA
CAGCCAACGGGCGCAGCTGGTGACATGGATGAAAGCAATACCACCGGTGCAGCGAGCATTCA
GGCTGGACTGCCTGCTTCCTGGGTTGTGGGGGATAAAACCGGCAGCGG

BLAST output: Escherichia coli strain IPK182 beta-lactamase **CTX-M-15** (blaCTX-M) gene, blaCTX-M-15 allele, complete cds.Sequence

ID: MH900522.1Length: 876Number of Matches: 1