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Investigating the role of the cell envelope in *E. coli* relating to silver sensitivity and resistance

by

Connor Westersund

A THESIS

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Abstract

Silver (I) is an antimicrobial agent that has established antimicrobial activity, yet the mechanism of action is unclear. This thesis follows up on an observation that Ag⁺ ions cause the cell membrane to separate from the cell wall in *Escherichia coli*. Data from a chemical genomic screen was utilized, identifying 6 Ag-responsive cells with single gene deletions (*damX*, *rodZ*, *minC*, *sanA*, *ybhO*, *tolB*), which are genes involved in maintenance of the cell envelope. When these mutants were grown in the presence of Ag⁺, cells demonstrated extensive cell envelope damage as seen by transmission electron microscopy. Evaluation of kill curves, some mutants conferred resistance and others sensitivity to silver in comparison to wildtype. From the data collected, it was recognized that functions around cell wall and transport across the cell membranes have roles in both silver resistance and sensitivity.

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Abbreviation and terms	Technical definition
Ab ^R	Antibiotic resistance cassette
Amp ^R	Ampicillin resistance, generated from an ampicillin resistance cassette.
ATP	Adenosine triphosphate
bet	A lambda phage gene that produces a Recombination protein that functions in general recombination and in the late, rolling-circle mode of lambda DNA replication.
Biofilm(s)	Any group of microorganisms in which cells stick to each other and often also to a surface. These adherent cells become embedded within a slimy extracellular matrix that is composed of extracellular polymeric substances (EPS).
BSC	Biosafety cabinet
<i>Cm^R</i>	Chloramphenicol resistance, generated from a Chloramphenicol resistance cassette or a <i>cat</i> (Chloramphenicol acetyltransferase) gene.
CBD	Calgary Biofilm Device is a lid that fits a 96 well microtiter plate, which contains 96 polystyrene equivalent pegs.
CLSM	Confocal laser scanning microscope
DMSO	Dimethyl sulfoxide is an organosulfur compound with the formula (CH ₃) ₂ SO
dNTP	nucleoside triphosphate (NTP) is a molecule containing a nucleoside bound to three phosphate groups.
dsDNA	Double stranded DNA
E. coli	Escherichia coli
EDTA	Ethylenediaminetetraacetic acid

Section 1.0: Symbols, Abbreviation, and Nomenclature

EtOH	Ethanol
exo	A lambda phage gene that produces a protein exonuclease that facilitates phage DNA recombination through the double-strand break repair (DSBR) and single-strand annealing pathways
Flp	Flippase (Flp) derived from the 2 μ m plasmid of baker's yeast <i>Saccharomyces cerevisiae</i> .
FRT	flippase recognition target
gam	A lambda phage gene that produces protein that protects linear double- stranded DNA from host exonuclease degradation
GC	Growth control
gDNA	Genomic DNA
g	g-force
HGT	Horizontal gene transfer
IM	Inner membrane
Kn ^R	Kanamycin resistance, generated from a Kanamycin cassette.
keV	Kiloelectron volt
LB	Lysogeny broth
LPS	lipopolysaccharide
MCS	Multiple cloning site
MDR	Multidrug- resistant
NSLB	No Salt Lysogeny Broth medium
OAS	Outside antisense primer, confirmation primer that binds to the antisense (non-coding) strand of DNA 3'-5'

OD	Optical density
OD600	Optical density measurement at 600 nm
ОМ	Outer membrane
O/N	Refers to overnight growth of culture medium
ORF	Open reading frame
OS	Outside sense primer, confirmation primer that binds to the coding strand of DNA 5'-3'
PCR	Polymerase chain reaction
Planktonic	floating as single cells in liquid medium
PVD	Physical vapor deposition
ROS	Reactive oxygen species
rpm	Revolutions per minute
RT	Room temperature
(-)/SC	Sterility control
SDS	Sodium dodecyl sulfate
SEM	Scanning electron microscope
SOC	Super optimal broth with catabolite repression recovery media
TAE	Tris-acetate-EDTA

Taq	A thermostable DNA polymerase named after the thermophilic bacterium Thermus aquaticus from which it was originally isolated from	
TEM	Transmission electron microscope	
UV	Ultra violet radiation	
V	Volts	
WT	Wildtype strain	

Chapter One: Introduction

1.1 Metal antimicrobials

Metals are essential for all forms of life, fulfilling certain structural and catalytic roles in biological processes, including precursor biosynthesis, DNA replication, transcription, respiration, and response to oxidative stress (Palmer and Skaar, 2016; Lemire *et al.*, 2013). For instance, transition *d*-block metals, like: manganese, iron, cobalt, nickel, and copper, have unfilled *d*-orbitals and therefore are redox active (Palmer and Skaar, 2016). These redox-active essential metals are found ubiquitously in several types of living organisms, functioning nearly exclusively as constituents of proteins, such as enzymes, storage proteins, and transcription factors (Lemire *et al.*, 2013; Porcheron *et al.*, 2013). Furthermore, their finely tuned redox activity permits essential metals to transfer electrons as part of vital biological processes, such as: respiration, N fixation, and photosynthesis (Finney and O'Halloran, 2003; Lemire *et al.*, 2013). Although these metals are essential for many crucial biological processes, these metals are lethal at high intracellular concentrations (Lemire *et al.*, 2013). Moreover, particular non-essential metals such as: silver (Ag), mercury (Hg), lead (Pb), and tellurium (Te) have been demonstrated to be extremely toxic to most microorganisms and have bactericidal activity at exceptionally low concentrations (Harrison *et al.*, 2004; Lemire *et al.*, 2013; Nies, 1999).

Since metal ions demonstrate extremely potent toxicity towards bacteria and yeast, particular metals have been used for their antimicrobial properties since ancient times (Lemire *et al.*, 2013). For example, vessels impregnated or made of Cu and Ag metal have been used in water disinfection and food preservation practices since the time of the Persian kings (Lemire *et al.*, 2013). This implementation of metals to reduce contaminant loads of resources was then later adopted by other ancient civilizations such as the Phoenicians, Greeks, Romans, and Egyptians (Alexander, 2009; Lemire *et al.*, 2013). Moreover, the medicinal use of metals has also been prevalent throughout antiquity to treat an array of different ailments such as leprosy, tuberculosis, gonorrhea, and syphilis (Lemire *et al.*, 2013). However, with the discovery of penicillin by Nobel laureate Sir Alexander Fleming in the 1920s (Aminov, 2010; Clardy *et al.*, 2009; Tan and Tatsumura, 2015), and the subsequent commercial development of new antibiotic drugs, metal antimicrobials prevalence in medicine rapidly diminished (Lemire *et al.*, 2013; Tan and Tatsumura, 2015).

Today, antimicrobial metal compounds can be found in wound dressings, coatings of indwelling medical devices, and as metallic surfaces (Afessa *et al.*, 2010; Kollef *et al.*, 2008; Saint *et al.*, 1998). Nonetheless, with the emergence and rise of multi-drug resistant (MDR) bacteria accompanied by the severe decline in the rate of development of new antibiotics, a resurgence of research into alternatives antimicrobials, like metals, is generating new applications for industry, agriculture, and healthcare (Allen *et al.*, 2014; Lemire *et al.*, 2013). This research has led to the determination and discovery that particular metals disrupt antibiotic-resistant biofilms (Teitzel and Parsek, 2003), exert synergistic bactericidal activity with other antimicrobials (Harrison *et al.*, 2008), inhibit metabolic pathways (Figure 1.0) (Middaugh *et al.*, 2005; Macomber *et al.*, 2011; Macomber and Elsey, 2009), and kill multidrug-resistant bacteria (Kaneko *et al.*, 2007), resulting in metal based-antimicrobials having real potential for profiling sustainability solutions to infection care and health (Turner, 2017).

However, in the era of MDR bacteria, metals unfortunately are not the end-all cure-all solution that the medical field is looking for (WHO, 2014; Hobman and Crossman, 2014). Just like with the rampant exposure to antibiotics giving rise to antibiotic resistance bacteria (Džidić et al., 2008; Spratt, 1994; McDermott et al., 2003), the natural exposure of bacteria to bioavailable metals (both essential and toxic) over the past billions of years has likely been the driver for the evolution of the ability of microorganisms to control cellular levels of these bioavailable metal ions (Hobman and Crossman, 2014). Thus, bacteria have evolved mechanisms for the acquisition of essential metals, controlling intracellular levels of metals, and eliminating those that are in excess and are deleterious (Hobman and Crossman, 2014). Microorganisms can resist metal toxicity by extracellular or intracellular sequestration of the metal, reduction in permeability, alteration of target sites, enzyme detoxification or efflux of the metal ions, highlighted in Figure 1.1. (Hobman & Brown, 1997). For example, a well characterized genetic elements, which confers resistance to silver (Ag⁺) is the *sil* operon, which microorganisms obtain through horizontal gene transfer (HGT). This operon is comprised of a gene cassette, which includes *silP*, *silA*, *silB*, *silC*, *silR*, *silE*, ORF105, and *silABC* (Belly and Kydd, 1982). Since their initial identification, sil genes have been identified in Salmonella Enterobacter, Escherichia coli, Pseudomonas, Acinetobacter, Klebsiella, and methicillin-resistant coagulase-negative Staphylococcus aureus (Finley et al., 2015). The Sil system is believed to mediate silver resistance by limiting the accumulation of silver in the cell through a combination of silver

sequestration in the periplasm (via SilE and SilF binding) and active efflux transporter SilCBA and the putative P-type ATPase transporter SilP (Randall *et al.*, 2015). Although many different types of metal resistant mechanisms have been well characterized (Hobman and Crossman, 2014), there have been far fewer investigations of the biochemical and biophysical mechanisms through which metals exert toxicity in bacteria (Lemire *et al.*, 2013). This area of study has been neglected and overshadowed by the complexity and metabolic diversity of microorganisms, which complicates the study of metal toxicity (Lemire *et al.*, 2013). Uncovering the unique biochemical pathways and toxicity mechanisms of different metal-based antimicrobials is imperative if we are to understand why resistance to these elements evolves and to improve the design of metal-based biocide agents to kill bacterial pathogens (Allen *et al.*, 2014; Hobman and Crossman, 2014; Lemire *et al.*, 2013).



Figure 1.0: General schematic of metal ion toxicity mechanisms: Mechanisms include: ROS generation causing DNA, protein and lipid damage; inhibition and disruption of membrane transport; impaired membrane function; redox reactions with thiol groups in proteins and enzymes resulting in enzyme damage and dysfunction.



Figure 1.1: Illustration of general metal ion resistance in bacteria. Mechanisms include: extracellular or intracellular sequestration of the metal; reduction in permeability; alteration of target sites; enzyme detoxification or efflux of the metal ions; acquisition of foreign DNA coding from antibiotic resistance; Mutations in the chromosome leading to resistance.

<u>1.2 The metal antimicrobial, silver (Ag+)</u>

Since ancient times, the antimicrobial activity of silver has been known to be effective against a broad range of microorganisms (Akram et al., 2016; Jung et al., 2008; Lemire et al., 2013). Herodotus, the Father of History, accounts that no Persian king, including Cirrus, would drink water that was not boiled and transported in flagons of silver, for the silver reduced the contaminant load of the water, while keeping it fresh for years (Alexander, 2009; Lemire et al., 2013). Over the millennia, silver has been utilized for variety of purposes due to its low exhibited toxicity towards eukaryotic cells in comparison to its lethality at low concentrations to prokaryotic organisms (Alexander, 2009). Now, in the 21st century, silver ions (Ag⁺) are being utilized to control infection and contaminant loads in a variety of medical practices and devices, including dental restorations, coatings on indwelling medical devises (catheters and endotracheal tubes), eyedrops, and bandages. (Klasen, 2000; Lemire et al., 2013; Silver and Phung, 1996; Slawson et al., 1992). Silver has also started to become implemented into a number of nonmedical consumer products including: deodorants, toothbrushes, toilet seats, drinking glasses, and even washing machines (Jung et al., 2008; Turner, 2017). With silver's resurgence in applications of both the medical and consumer product fields of industry, it is surprising that the mechanisms by which silver ions (Ag^+) impose their toxicity onto microbial cells is still incomplete (Celment and Jarret, 1994). In the literature, there have been many observations of different Ag⁺ cellular targets, which has led to the establishment of some mechanisms of the antimicrobial action of silver ions (Jung et al., 2008). What has been observed is that the antimicrobial action of silver ions is closely correlated to their interactions with thiol (sulphhydryl, R-SH) groups (Belly and Kydd, 1982; Bragg and Rainnie, 1974; Furr et al., 1994; Lemire et al., 2013). This relationship is described by the hard-soft acid base theory, which has been determined empirically and provides an ordering of transition metals according to their preferences for specific organic ligands (Jung et al., 2008; Lemire et al., 2013.) Essentially, Ag(I) is classified as a soft acid and consequently has electron-sharing affinities that can result in the energetically favorable formation of covalent bonds between soft bases, such as the sulphhydryl groups that are found in proteins (Lemire et al., 2013). Thus, the interaction and binding of silver ions to thiol groups in enzymes and proteins, results in their deactivation (Dakal et al., 2016). As a result, cells exposed to Ag⁺ ions can have a depletion of antioxidant

reserves, particularly glutathione, which is responsible for preventing damage to important cellular components caused by reactive oxygen species (ROS).

Further *in vitro* experiments have revealed that micromolar concentrations of Ag^+ ions directly inactivate purified fumarase A, a member of the dehydratase family (Xu and Imlay, 2012). Additional examinations, utilizing electron paramagnetic resonance analysis and measurements of released iron confirmed that damage was directly correlated to the destruction of the [4Fe-4S] cluster, and indeed, the reconstruction of the cluster fully restored fumarase activity (Xu and Imlay, 2012). Furthermore, it is thought that the direct or indirect destruction of [4Fe-4S] clusters caused by Ag^+ ions could result in the release of additional Fenton-active Fe^{2+} into the cytoplasm, generating hydroxide (OH⁻) and the highly reactive hydroxyl radical (OH⁺) (Lemire *et al.*, 2013; Park *et al.*, 2009; Xu *et al.*, 2012). However, the precise mechanism of ROS-mediated antibacterial activity of Ag^+ is not fully clear (Pellieux *et al.*, 2000).

Another established Ag⁺ toxicity mechanism is silver's ability to disrupt the activity of the bacterial electron transport chain (ETC) (Bragg and Rainnie, 1974; Gorden *et al.*, 2010; Lemire *et al.*, 2013 Lok *et al.*, 2006). In the organism *Vibrio harveyi*, Ag(I) has been demonstrated to inhibit the activity of NADH:quinone oxidoreductase (NQR), which is a component of the respiratory chain of some bacteria and generates a redox-driven transmembrane Na⁺ potential by translocating Na⁺(Lemire *et al.*, 2013). It is proposed that Ag(I) dissipates the chemiosmotic potential of the membrane by destroying the proton motive force of the ETC, causing proton leakage through the membrane. (Dibrov *et al.*, 2002; Lemire *et al.*, 2013).

One observed toxicity phenotype with no established mechanism involves Ag^+ ability to jeopardize the integrity of the cellular envelope in *E. coli* and *Staphylococcus aureus* (Jung *et al.*, 2008). The associated phenotype suggests that Ag^+ ions severely compromise the cell envelope, resulting in the cell lysis and separation of the cell membrane from the cell wall (Jung *et al.*, 2008). What remains unclear from morphological studies examining the toxicity of Ag^+ towards the cell envelope is that it is difficult to determine whether membrane disruption or the observed detachment of the cell wall are a cause or a consequence of cell death (Jung *et al.*, 2008). Here from this thesis work, I look to add investigative insight into possible genetic determinants associated with the cell envelope of *E. coli* to be possibly involved in Ag(I) incomplete toxicology.

<u>1.3 Hypothesis and aims</u>

Following a similar methodology to Baryshnikova et al., 2010 "Quantitative analysis of fitness and genetic interactions in yeast on a genome scale", Dr. Lemire from Dr. Tuner's Microbial Biochemistry Laboratory at the University of Calgary, preformed a genetic chemical screen of the Keio collection, an ordered mutant library of E. coli. The Keio collection was systematically created by Baba et al., 2006, to contain all single-gene deletions of all nonessential genes in E. coli K-12. The chromosomal deletions of the Keio collection were facilitated by the Lambda Red Homologous Recombination-based technique (Baba et al., 2006). The Recombinase technique enabled Baba and colleagues to target 4288 open reading frames of coding genes and replace them with a kanamycin cassette flanked by FLP recombinase recognition target sites (FRT), resulting in their inactivation (Baba et al., 2006). The Keio collection offer researchers a new resource for systematic analyses of unknown gene functions and gene regulatory networks, but also for genome-wide testing of mutational effects and responses to antimicrobial agents in a common strain background, E. coli K-12 BW25113 (Baba et al., 2006). Dr. Lemire utilized the Keio collection to screen Keio collection E. coli colony size as a proxy for fitness, of 3985 confirmed non-essential E. coli mutants to 100µM of AgNO₃ infused in M9 minimal noble agar media against their unchallenged counterparts. From the genetic chemical screen, Dr. Lemire discovered over 225 mutants silver sensitive and 177 mutants to display a resistant phenotype. The Sensitive ('sicker') or resistant ('healthier') mutants were selected based on a difference of 2 standard deviations of colony size from the normal, which was determined through photographed images that were processed using software that measures colony areas in terms of pixels. Of these mutants, 20 sensitive and 4 resistant mutants were found to be involved in cell envelope homeostasis. From these 24 genes, 6 (damX, rodZ, minC, sanA, ybhO, and tolB) were chosen to be further characterized based of their possible collective phenotype having possible involvement in Ag(I) cell envelope toxicity mechanism.

The overarching goal of this thesis was to provide answers in regard to the cell envelope associated genetic determinants chosen from the genetic chemical screen, having potential involvement in Ag^+ mechanism of bacterial toxicity. Validation of the *E. coli* cells phenotypic responses to Ag^+ ions would provide understanding to the biochemical and genetic basis of Ag^+ toxicity. In this thesis we aimed to provide insight into specific questions:

- 1.) Are there genotypic differences between Keio collection BW25113 and K-12 MG1655 cells, that lead to different metal response phenotypes in the presence of metal antimicrobials?
- 2.) Is the growth of the wildtype and cell envelope associated Ag-responsive BW25113 and MG1655 *E. coli* cells effected while grown in the presence of Ag⁺ and other metal toxins?
- 3.) Do Ag⁺ ions effect planktonic and surface biofilms of both wildtype and cell envelope associated Ag-responsive mutant *E. coli* cells? Is the total biomass accumulation of *E. coli* Ag-responsive mutants and wildtype biofilms effected by the presence of Ag⁺ ions?

We aimed to utilize a variety of microbiological techniques to answer the postulated questions. First, using similar methodology to Datsenko *et al.*, 2000, we utilized the Red-recombinase recombination method to generate chromosomal disruptions in the MG1655 genomic background of *E. coli*, aimed to be used for a genomic comparative analysis against Keio collection mutants. Second, utilizing metal exposure-response curves, I aimed to evaluate Ag^+ and a comparative metal antimicrobial Cu^{2+} , ability to inhibit the proliferation of both wildtype and cell envelope associated mutants. For the third aim, I will generate insight into answering the proposed questions from three different types of microscopy techniques: CLSM, SEM and TEM, which aim to characterize the surface morphology of biofilms (CLSM & SEM) and planktonic cell morphology (TEM) in the presence and absence of Ag^+ .

It is known that the antimicrobial activity of Ag based compounds is intrinsically dependent on the formation of the Ag⁺ ions (Walker and Parsons, 2014). Thus, the metal challenges that were selected for the characterization of the collective phenotyping of the Ag-responsive mutants and wild type cells were: silver nitrate (AgNO₃), silver sulfadiazine (AgSD), and Copper (II) sulfate (CuSO₄). AgNO₃ was chosen based of its high solubility, while AgSD was chosen based of its clinical relevance for it is frequently used as prophylactic agent in burn patients (Bessey, 2007), and CuSO₄ was chosen as a comparator metal antimicrobial.

Chapter Two: Methods

2.0 Materials and media

2.1.1 Bacterial strains

K-12 MG1655 and BW25113 *Escherichia coli* strains (standard laboratory strains used commonly in the study of antimicrobial resistance) were provided by National BioResource Project *E. coli* K-12 single-gene-knockout Keio Collection. The *E. coli* DH5 α strain, which contained template and helper plasmids (see Table 2.0 for genotype) are available in Turner lab archive. All plasmid preparation and isolation procedures were conducted on DH5 α cells containing the particular helper plasmid needed for individual experiments within this thesis work. The Red-recombinase expressing plasmid was provided by Dr. Joe Harrison (University of Calgary) K-12 MG1655::pKD46 and utilized for λ Red-recombinase gene replacement. All strains were stored at -80°C in 8% (v/v) dimethyl sulfoxide (DMSO) in Lysogeny–Broth medium (pH 7.0, 5.0g NaCl, 5.0g yeast extract, and 10.0 g tryptone per litre of double distilled water (ddH₂O) in a glass vial. The individual strains and associated genotypes are depicted in the table 2.0 below.

Strain	Genotype	Antibiotic Resistance	Function of deleted gene of interest
DH5a	F- Φ80lacZΔM15 Δ(lacZYA- argF)U169 recA1 endA1 hsdR17 (rK– , mK+) phoA supE44 thi-1 gyrA96 relA1	None	N/A
K-12 MG1655	F [•] , lambda ⁻ , <i>rph-1</i>	None	N/A
BW25113	F-, Δ(araD-araB)567, ΔlacZ4787(::rrnB-3), λ-, rph-1, Δ(rhaD-rhaB)568, hsdR514	None	N/A
BW25113	BW25113; $\Delta dam X::Kn^R$	Kn ^R	<i>DamX</i> is a non-essential cell division protein; it contains a C-terminal SPOR domain which targets the protein to the septal ring.
	Ag ⁺ - Resistant		
BW25113	BW25113; $\Delta rodZ$:: Kn^R	Kn ^R	<i>rodZ</i> is a bitopic inner membrane protein that is involved in the maintenance of cell shape through interaction with the MreB cytoskeleton.
	Ag ⁺ - Sensitive		
BW25113	BW25113; $\Delta minC::Kn^R$ Ag ⁺ - Sensitive	Kn ^R	The <i>minC-minD-minE</i> system acts to direct septation to the proper (central) site in the dividing <i>E. coli</i> cell.
BW25113	BW25113; $\Delta sanA::Kn^R$	Kn ^R	Multi-copy expression of <i>sanA</i> complements the vancomycin sensitivity of an <i>E. coli</i> K-12 mutant which exhibits outer membrane permeability defects
	Ag ⁺ - Sensitive		
BW25113	BW25113; $\Delta ybhO::Kn^R$	Kn ^R	ybhO (clsB) is the second cardiolipin synthase in E. coli
	Ag ⁺ - Sensitive		
BW25113	BW25113; ∆ <i>tolB</i> ::Kn ^R Ag ⁺ - Sensitive	Kn ^R	<i>tolB</i> is a periplasmic component of the Tol-Pal system - a group of interacting proteins which span the cell envelope of <i>E. coli</i> K-12. The Tol-Pal system plays a role in outer membrane invagination during cell division and is important for maintaining outer membrane integrity.

Table 2.0: Associated genotypes of *E. coli* strains used within this study.

* Kn^R = Kanamycin resistance, Cm^R =Chloramphenicol resistance, Amp^R =Ampicillin Resistance

2.1.2 Chemicals and reagents

All chemicals used in the preparation of media were supplied by Sigma Aldrich (Missouri, USA), EMD (Darmstadt, Germany), Bio-Rad Laboratories (California, USA) and BD Biosciences (New Jersey, USA). Reagents and enzymes used in cloning procedures were purchased from Thermo Fisher Scientific (Massachusetts, USA).

2.1.3 Medium and culture conditions

Throughout our studies—present and past—we have observed that the growth media chosen to culture bacterial cells is a significant factor that dictates the metal toxicity. Hence, we selected a media that provides a rich environment to ensure robust bacterial growth in each strain. Minimal salts media (M9) (10.0 mL M9 5x salts, 38.5 mL ddH₂O, 1.0 ml dextrose, 0.10 mL MgSO₄, and 0.005 mL CaCl₂) was utilized for metal susceptibility testing. The decision was made not to use rich media (high NaCl concentration) based on ionic bonding chemistry, where reducing the amount of free Cl⁻ ions in solution would reduce precipitation AgCl (s) and allow for more available Ag⁺ ions in solution. For routine overnight starting cultures (16 or 24-hour growth), diluting cultures for screening assays, transformation, and plasmid isolation, strains were incubated in test tubes at 37°C under aerobic conditions in Lysogeny broth, M9 minimal salts media, or on LB plates (1.25% (w/v) agar). Antibiotics ampicillin (*Amp*), chloramphenicol (*Cm*), and Kanamycin (*Kn*) were added in LB medium or while pouring LB plates as required to a final concentration of 100 µg/mL, 10-34 µg/mL, and 10-25 µg/mL respectively.

2.1.4 Antimicrobial metal stock solutions

Silver nitrate (AgNO₃), copper (II) sulfate (CuSO₄), gallium (III) nitrate (Ga(NO₃)₃·H₂O) and nickel sulfate (NiSO₄ \cdot 6H₂O) were all obtained from Sigma-Aldrich (St. Louis, MO, USA). Aluminum sulfate (Al₂(SO₄)₃·H₂O) was obtained from Matheson Coleman and Bell (Norwood, OH, USA) and zinc sulfate (ZnSO₄ \cdot 7H₂O) was received from Thermo Fisher Scientific (Fair Lawn, NJ, USA). Stock solutions of CuSO₄, Al₂(SO₄)₃ \cdot H₂O were made up to 1M, ZnSO₄ \cdot 7H₂O was made up to 1.5M, NiSO₄ \cdot 6H₂O to 2.5M and AgNO₃ to 500mM in distilled and deionized (dd)H₂O. All stock metal solutions were stored in glass vials at 21°C for no longer than two weeks.

2.1.5 Stock antibiotics solutions

Ampicillin (Sigma-Aldrich) and Kanamycin (Sigma-Aldrich) were prepared as stock solutions in double-distilled water (ddH₂O) at 100 and 25 μ g/ mL respectively. Chloramphenicol (Sigma-Aldrich) was prepared in 50 % ethanol (99.8%) and treated identically to the other antibiotics. The antibiotics were syringe-filtered and stored at – 20°C.

2.1.6 Plasmids and vectors

Table 2.1: All Plasmids	and vectors used in t	he work of this thesis.

Vectors	Description	Source			
Template plasmid λ Red recombinase					
pKD13	Amp ^R and Kn ^R ; plasmid markers are oriR6K γ Tl3 _λ (Terminator), <i>bla</i> , <i>rgn</i> B (Terminator), <i>kan</i> ; Accession Number AY048744	CGSC			
pKD32	Amp ^R and Cm ^R ; plasmid markers are oriR6K, tL3 ₂ (Terminator), <i>bla</i> , <i>rgn</i> B (Terminator), <i>kan</i> ; Accession Number AY048744	CGSC			
Temperature sensitive plasmid λ Red recombinase					
pCP20	Amp ^R and Cm ^R ; (only grows at 30 °C), $[cU857](\lambda)ts$, bla , cat , $[Flp]$; carries the yeast Flp recombinase gene (flp)	CGSC			
pKD46	Amp ^R , <i>repA101</i> (ts), <i>araBp-gam-bet-exo</i> , <i>oriR101</i> , <i>bla</i> ; more efficient that pKD20 at making gene disruptions; Accession number AY048746	CGSC			
Complementation					
pTZ57R/T	Linearized pTZ57R vector with 3'-ddT overhangs for TA cloning of PCR products with blue/white screening.	Thermo Fisher Scientifc			
pMS199EH _A	Cloning Vector- Amp ^R	E. Lanka			
Metal exposure-response					
pBBRIMC-2	Cloning Vector- Kn ^R	M.E. Kovach			

*CGSC = Coli Genetic Stock Center; Amp^R = ampicilin resistance; Cm^R = chloramphenicol resistance; Kn^R = kanamycin resistance.

2.1.7 Optical density

McFarland standards were utilized to visually approximate the concentration of cells in a suspension. The McFarland scale represents specific concentrations of CFU/mL and is used for estimating concentrations of Gram negative bacteria such as *E. coli*. Note that this estimate becomes uncertain with organisms outside the normal usage as different species of bacteria differ in size and mass. Use of this method requires calibration and validation. McFarland Standards are generally labeled 0.5 through 10 and filled with suspensions of Barium salts. The standards were created by preparing a 1.0% solution of anhydrous BaCl₂ and a 1.0% solution of H₂SO₄. The standards were stored in the dark, in a tightly sealed container at 20-25°C and should be stable for approximately 6 months. The advantage of the use of these standards is that no incubation time or equipment is needed to estimate bacterial numbers. The disadvantage is that there is some subjectivity involved in interpreting the turbidity, and that the numbers are valid only for those microorganisms similar to *E. coli*.

2.1.8 Chemically competent E. coli cells preparation and transformation

Chemically competent K-12 MG1655 and BW25113 E. coli cells used in the study were prepared using calcium chloride method (Chan et al., 2013). Overnight cultures were diluted to 1 McFarland standard d (~1.0 x 10⁸ colony forming units(CFU) mL⁻¹) in 7.0 mL of LB and incubated aerobically at 37°C to mid-log phase until an optical density at 600 nm (OD₆₀₀) of 0.3-0.4 units was reached. The cultures were incubated on ice for 15 min and centrifuged at 5000 rpm for 5 min at 4°C to obtain cell pellet. The cell pellet was re-suspended in 10 mL of ice-cold 1M CaCl₂·2H₂O and incubated on ice for 1 hour. Another round of centrifugation followed by resuspension in 10 mL of ice-cold 1M CaCl₂·2H₂O and incubation on ice for 1 hour was done. After discarding the supernatant, the pellet was re-suspended in 2 mL of cold 1M CaCl₂·2H₂O, 16% (v/v) glycerol 200 μ L aliquots were made in sterile 1.5 mL microfuge tubes. All freshly prepared competent cells were flash frozen and stored as cryogenic stock at -80 °C. All transformed and co-transformed E. coli strains used within this thesis were generated by heat shock transformation method or by electrophoresis. For plasmid transformation, $1-2 \mu L$ (50-100 ng) of plasmid DNA was added to 50 µL of competent cells, which were aseptically aliquoted into 1.5 mL microfuge tubes. For cloning purposes, 15 µL of ligation reaction mix was added to 50 µL competent E. coli DH5a cells. Mixture of DNA and competent cells was incubated on ice

for 20 minutes and then heat shocked for 90 seconds at 42 °C in water bath. After the heat shock treatment, the centrifuge tubes were returned to ice for 10 minutes. 500 mL of fresh LB was added to each of the tubes and incubated at 37 °C for 1 hour in a shaking incubator at 200 rpm. Following incubation 100 μ L of the culture was aseptically transferred and spread onto LB plates (1.25% (w/v) agar) (section 2.1.3).

2.1.9 Biofilm production

Following a protocol similar to Harrison et al., 2010, biofilms where grown from cryogenic stock stored at -80°C, a 200 μ L pipette tip was used to collect the different bacterial strains (Table 2.0) and inoculate 7 sterile 22.0 mL glass test tubes filled with 7.0 mL of M9 or LB media (6 tubes for the mutants and one for wildtype cultures). The test tubes were grown overnight for 24 hours at 37°C and 150 rpm. Following the overnight growth of the preculture, colonies were resuspended into fresh M9 media or LB media and matched to 1.0 McFarland standard. Next, the cultures were then poured into reagent reservoirs and using a multichannel micropipette, $180 \,\mu\text{L}$ of the bacterial cultures were aliquoted into each well from rows B-G, columns 2-12 of a sterile 96 well microtiter plate. Wells in rows A and H acted as negative controls, which contained 180 µL sterile 0.9% saline, while column one's rows B-G contained 180 µL of bacterial culture without an antimicrobial challenge (Growth Control). Both wildtype and mutant *E. coli* biofilms cells had 3 technical replicates when subjugated to the metal exposure-response experiment and subsequent crystal violet assay. The 96 well microtiter plate layout is depicted in Figure 2.0 and 2.1. Using a multichannel micropipette, 180 μ L of the concentrated metal challenges (tables) were transferred into column 2 wells of rows B-G (The site of highest antimicrobial challenge during testing). The contents of the wells were thoroughly mixed by pipetting the solution up and down several times. After mixing, transfer 180 μ L from the wells in column 2 to the corresponding wells in column 3. Mix the contents of column 3 and transfer 180 μ L from each well into the corresponding wells of column 4. The Serially mix and transfer process was continued down the length of the microtiter plate until the last column (12) was reached. Once the final column is met, discard 180 µL of the mixed contents to create equal volumes of wells for the entire microtiter plate. After the wells received the proper inoculum volumes, a Calgary Biofilm Device (CBD) was aseptically transferred to cover the 96 well microtiter plate. The CBD lid contains 96 equivalent polystyrene pegs, which serve as a solid surface for planktonic bacterium to adhere to in order to undergo physiological changes

associated with biofilm production. The plates were then onto a gyrorotary shaker set to 150 rpm and incubated for 24 hours at 37°C.

2.1.10 Primers

Table 2.4 lists all the different primers generated and used throughout the experiments of this thesis.
Gene Knockout Primer	Sequence	GC%	Tm (°C)	Template homology	
	(Homology extensions)				
Knockout primers					
rodZ KO	cgg atg ttt cgc ggt gtg g	63.0	61.0	K-12 MG1655	
rodZ RC	gtt acc cgt ctg tta ctg cg	55.0	60.5	K-12 MG1655	
mreB KO	cgc tgc tgc gtg tgg ttg gta aag taa gcg	57.0	65.7	K-12 MG1655	
mreB RC	ctg cct gca tcc gat tac tct tcg ctg aac	53.0	64.4	K-12 MG1655	
minC KO	cat gcc tta tag tct tcg gaa cat cat c	43.0	58.0	K-12 MG1655	
minC RC	cgg ttg aac ggt caa agc g	58.0	59.0	K-12 MG1655	
damX KO	ccg ctt atc aag cgg tct att agc	50.0	59.0	K-12 MG1655	
damX RC	gat cgg cct gta cct gac g	63.0	59.0	K-12 MG1655	
ispG KO	ccc tca atg ccg aac aat c	53.0	57.5	K-12 MG1655	
ispG RC	ctt ccc tca atg ttt cgg g	53.0	57.5	K-12 MG1655	
P1* (added to forward primer)	gtg tag gct gga gct tc	59.0	62.5	pKD13 OR pKD32	
P4* (added to reverse complement primer)	att ccg ggg atc cgt cga cc	65.9	64.6	pKD13 OR pKD32	
Confirmation Primers for λ Red- recombinase	l				
rodZ F confirmation primer (OS)	gtg cgt ttt tgc tga ctt taa g	41.0	58.4	K-12 MG1655	
<i>rodZ</i> RC confirmation primer (OAS)	gat ttg att gac cgt tgc ttc	43.0	57.5	K-12 MG1655	
<i>mreB</i> F confirmation primer (OS)	gcg gtt gca aac agg cga g	63.0	61.5	K-12 MG1655	
<i>mreB</i> RC confirmation primer (OAS)	gta gaa agg act gac ggc gg	60.0	62.5	K-12 MG1655	

Table 2.2: Primers utilized throughout this thesis.

<i>minC</i> F confirmation primer (OS)	cac aac cca taa tca ggt cga gat tac g	46.0	59.9	K-12 MG1655
minC RC confirmation primer (OAS)	aga tgc tgt cac gat tag ggt gc	52.0	57.1	K-12 MG1655
<i>damX</i> F confirmation primer (OS)	gcc gtt taa tat cat caa gca ggg	46.0	63.6	K-12 MG1655
<i>damX</i> RC confirmation primer (OAS)	gcg ctt aat tct tcc gtt ggc	52.0	61.2	K-12 MG1655
<i>ispG</i> F confirmation primer (OS)	cag tac aga tcc agt atc aag g	45.0	60.1	K-12 MG1655
<i>ispG</i> RC confirmation primer (OAS)	tgc cgc gaa tgg ctt gaa tg	55.0	60.5	K-12 MG1655
sanA F confirmation primer (OS)	ctc tcg gct gtc aca gta tc	55	56.0	BW25113
sanA RC confirmation primer (OAS)	Ccg aag ccg ata tat gcc ac	55	59.0	BW25113
<i>ybhO</i> F confirmation primer (OS)	cgg act gga tga gat tit tac c	45	55.0	BW25113
ybhO RC confirmation primer (OAS)	gca gccc gta aat cag ata gct g	50	58.0	BW25113
<i>tolB</i> F confirmation primer (OS)	tcc cga aaa cca ccgc cag	60	60.0	BW25113
tolB RC confirmation primer (OAS)	ccg ccg ttc gca tcc ata c	63	61.0	BW25113
Complementation Primers		Antibiotic resistance	Gene of interest	Restriction Enzyme site
pMS199EHA::rodZ	Forward: includes start (ATG) codon	Amp	rodZ	XbaI:TCTAGA
	ATATTCTAGAAGGAGAAATAATATG AATACTGAAGCCACGCACGA			HindIII:AAGCTT
	Reverse: includes stop (ATT) codon			
	ATATAAGCTTTTACTGCGCCGGTGATTGTTCG			
pTZ57R/T::rodZ	Forward:	-	rodZ	XbaI: <mark>TCTAGA</mark>
	ata <mark>ttetagaa</mark> ggagaaataatatg aataetgaageeaegaa			HindIII:AAGCTT
	Reverse:			
	atataagcttttactgcgccggtgattgttcg			

*KO= Knockout forward primer, RC= Reverse complement Primer. *The combination of both primers KO+P1 and RC+P4 sequences from Table 2.0, generated the complete knockout primer that is utilized to facilitate the insertion of the template plasmids antibiotic cassette in place of gene of interest. Further, it is not necessary to include the 5' homology extensions in the calculation of melting TM. *F= Forward primer, RC= Reverse complement Primer, OS =outside sense, OAS= outside antisense. TCTAGA= XbaI restriction enzyme cut site, AAGCTT=HindIII restriction enzyme cut site. ATG=start codon, TTA= stop codon.

2.2 λ Red-recombinase mediated gene replacement

The process of λ Red-recombinase mediated gene replacement contains three key molecular tools. The first being a plasmid with a temperature sensitive replicon and an inducible operon that expresses the lambda (λ) phage genes *gam*, *bet*, and *exo*. Second, a plasmid that serves as a template for generating the gene replacements. This plasmid cannot replicate in desired *E. coli* strain (strain that is used for the targeted gene deletion) and must have an antibiotic cassette that is flanked by Flp-recombination targets (FRT) sites. Finally, a helper plasmid with a temperature sensitive replicon that transiently expresses the yeast (*Saccharomyces cerevisiae*) recombinase Flp (Datsenko and Wanner, 2000).

The process of generating a gene replacement allele utilizing the Red-recombinase molecular tool is accomplished through polymerase chain reaction (PCR). Primers are used to amplify the template plasmid (pKD13 or pKD32) antibiotic cassette, which are flanked by FRT sites. The primers used to amplify this cassette are created with a 5' flanking homology extension that are 18-26 nucleotides in length. These homology extensions precisely match upstream and downstream nucleotide sequences in the *E. coli* chromosome of the targeted loci for deletion. This PCR product is transformed into electrocompetent *E. coli* (K-12 MG1655) cells that contain the temperature sensitive helper plasmid (pKD46) that expresses the Redrecombinase genes (*gam, bet,* and *exo*) through induction with arabinose. The Gam protein prevents the endogenous RecBCD nucleases from digesting linear DNA introduced into the *E. coli*. Exo protein is a 5' \rightarrow 3' dsDNA-dependent exonuclease and will degrade linear dsDNA starting from the 5' end. The Beta protein protects the ssDNA created by Exo and promotes its annealing to a complementary ssDNA target in the cell. The recombinase then mediates the sitespecific insertions of the antibiotic cassette (FRT -Ab^R-FRT) through a double crossover event between the 5' homology extensions and their complementary gene sequences on the *E. coli*

chromosome. This helper plasmid can be further cured from the cells during antibiotic selection for insertions by growing at non-permissible temperatures of >30 °C. After selection, mutant *E. coli* cells can have the antibiotic cassette removed from the chromosome with the insertion of another temperature sensitive plasmid (pCP20) that transiently expresses the yeast recombinase, Flp. Following transformation and selection, the helper plasmid can be cured from the cell with growth at a non-permissive temperature (>30 °C). The then unmarked deletion mutants can be identified through their inability to grow on media supplemented with antibiotics (Datsenko and Wanner, 2000).

2.3 Complementation

Following the successful generation of the deletion mutant K-12 MG1655 $\Delta rodZ$, plasmid-based complementation of K-12 MG1655 \(\Delta rodZ\) was conducted through cloning methods utilizing restriction enzymes and T4 ligase. The Amp- resistant expression vector pMS119EH_A generated by Beketskaia et al., 2014, was selected as the expression system for complementation because the vector has previously demonstrated to be a low copy number plasmid with about 20-30 copies per cell and also having low levels of expression ("leaky expression") from the PtacI promoter in the absence of induction by IPTG (Bay and Turner, 2009). This expression type has proven to be advantageous for the current study to obtain low non-toxic levels of RodZ protein as previous literature has reported that over-expression of membrane proteins can have detrimental effects in the host cells (Jensen et al., 2017). Further, the selection of the pMS119EH_A vector was further chosen for its previous success complementing genes for membrane proteins by the Turner Research Group (Turner et al., 2017; Beketskaia et al., 2014). In this experiment, because we are cloning the ORF of rodZ and inserting (through ligation) the ORF into an expression vector, we include the start codon (ATG) and the stop codon (TGA). The forward primer was designed with a 18-21 bp homology sequence (refer to Table 2.2), which would bind and amplify the target ORF. Additionally, the forward primer was generated to have an XbaI restriction site (TCTAGA) added to the 5' end of the primer. Finally, the forward primer consisted of the addition of 3-6 bp upstream of the inserted cut site XbaI, in order to improve cutting efficiency. The reverse primer has similar design; however, the reverse complement was constructed utilizing OligoCalc: Oligonucleotide Properties Calculator software. Similarly, 18-21 bp with homology to the ORF were used to generate the primer with the addition of the HindIII restriction site (AAGCTT) on the 5' end.

Following the restriction site 4 bp were added to improve restriction enzyme digestion. The full protocol for generating a plasmid-based complementation can be found in the appendix section A of this thesis.

2.4 Culture density analysis

2.4.1 Bacterial growth curves

A bacterial growth curve is a graph illustrating the changes in size of a bacterial population over time. The bacteria are cultured in a sterile nutrient medium and incubated at the optimal temperature for growth. Aliquoted samples are removed at intervals and the concentration of bacteria is evaluated. A logarithmic growth curve is plotted, which shows various phases of growth. In the *lag* (or *latent*) phase there is only small increases in numbers as the bacteria imbibe water, and synthesize ribosomal RNA and subsequent enzymes, in adjusting to their new environmental conditions (Navarro et al., 2010; Rolfe et al., 2012) Different bacteria reproduce at different rates. The time it takes the population to double is called the generation time. As this generation time of the cells decreases, they enter the log (or exponential) phase, in which the cells reach a maximum rate of reproduction and the number of bacteria increases directly with time, giving a straight slope on a logarithmic scale. With time, as the population grows, it enters the stationary phase, where the nutrients and electron acceptors are depleted, and the pH drops as carbon dioxide and other waste poisons accumulate (Navarro et al., 2010). Eventually the cell's energy storage begins to deplete resulting in cultures having a drastic decrease in the rate of cell divisions. Finally, the death (or final) phase occurs when the rate at which the bacteria die exceeds the rate at which they are produced; the population declines as the levels of nutrients fall and toxin levels increase.

For this experiment, it was imperative to measure the different growth capabilities of the different cell envelope mutant *E. coli* strains to investigate whether they had similar growth patterns, which would allow for the determination of the appropriate exposure times for metal exposure-response experimentations. The different mutant and wildtype strains tested, are displayed in Table 2.0.

2.4.2 Inoculation and growth

Bacteria from cryogenic DMSO stocks at -80°C (section 2.1.1). Utilizing a 200 µL pipet tip frozen cryogenic stock cultures were inoculated into minimal salts media (M9) and incubated

overnight at 37°C and shaking at 250 rpm. Following the O/N growth, 100 μ L of culture were then aliquoted onto LB (1.25% (w/v) agar plates and incubated at 37°C overnight. A single colony was then selected from the O/N growth and inoculated into 12 sterile 22.0 mL glass test tubes containing 5.0 mL of M9 media and placed into a 37°C incubator and shaken at 125 rpm. The 12 test tubes represent the different time intervals at which 1.0 mL of sample was extracted and placed into 2.0 mL cuvette and their turbidity was measured through a SpectraMax Plus 384 Microplate Reader. The optical density (OD) for each culture was recorded for 1, 2, 4, 6, 8, 10, 12, 16, 18, 20, 24, and 48-hour time intervals for BW25113 wildtype and mutant cells. The resulting OD₆₀₀ values were recorded in biological triplicates with 3 technical trials for each wildtype and mutant cell to *E. coli* cell to ensure accuracy of the results.

2.5 Evaluation of different metal exposure-responses of BW25113 E. coli cells

2.5.1 Metal exposure-response curves

The metal exposure-response experiment was conducted to visualize the dose and exposure-response relationship between mutant and wildtype MG1655 and BW25113 *E. coli* cells against three different metal challenges: silver nitrate (AgNO₃), silver sulfadiazine (AgSD), and Copper (II) sulfate (CuSO₄)1This experiment aims to describe the phenotypic changes in *E. coli* caused by differing levels of exposure to an antimicrobial stressor after a certain exposure time. The end result will produce a dose-response curve, which is a simple X-Y graph relating to the magnitude of a stressor (concentration of the antimicrobial in μ M) to the response of the receptor (the respective OD or viable cell counts of an organism).

2.5.2 Generating the metal exposure-response toxicity curve

In order to generate a metal exposure-response curve a spectrophotometer was used to analysis the OD₆₀₀ antimicrobial challenge plate samples. The spectrophotometer method measures culture turbidity directly and allows for the analysis of culture concentration of bacteria relative to antimicrobial challenge in the suspension. The plates were removed from the incubator and taken to analyzed by a SpectraMax Plus 384 Microplate. The machine was switched on and the SoftMax[®] Pro optical density reading software was initialized. The program was constructed with the correct format that highlights the proper controls and bacterial culture samples that are illustrated in Figures 2.0 and 2.1.

2.5.3 Growing subcultures, inoculum preparation and preparation of antimicrobial challenge plate

Utilizing methods similar to Lemire et al., 2015, cryogenic stock stored at -80°C (Section 2.1.1), a 200 µL pipette tip was used to collect the different bacterial strains (Table 2.0) and inoculate 7 sterile 22.0 mL test tubes filled with 7.0 mL of M9 media. The test tubes were grown O/N for 16-hours at 37°C and 150 rpm. During this incubation time the antimicrobial metal challenges were prepared. The challenges were stored in a cold room at 4°C until needed for the metal exposure-response inoculum preparation. The 10 different bacterial suspensions were then prepared to equal turbidity of 1.0 McFarland standard ($\sim 1.0 \times 10^8$ colony forming units (CFU) mL⁻¹) inside a BioSafety Cabinet (BSC). This was achieved by aliquoting 1.0 mL of each bacterial culture into 7.0 mL of fresh M9 media. The cultures were then poured into reagent reservoirs and using a multichannel micropipette, 180 µL of the sterile bacterial cultures were aliquoted into each well from rows B-G, columns 2-12 of a sterile 96 well microtiter plate. Rows A and H acted as negative controls containing 180 µL sterile 0.9% saline (Sterility growth Control, SC), while column one's rows B-G contained 180 µL of bacterial culture without an antimicrobial challenge (Growth Control = GC). This determination of having these two rows act as negative control wells is because of published literature confirming increased evaporation rate of these exterior wells of a 96 well microtiter plate (Harrison et al., 2010). Both wildtype and mutant *E. coli* cells had 3 technical replicates when subjugated to the metal exposure-response experiment. The 96 well microtiter plate layout is depicted in Figures 2.0 & 2.1. Using a multichannel micropipette, 180µL of the concentrated metal challenge was transferred into column 2 wells of rows B-G (The site of highest antimicrobial challenge during testing, 50 or 256 µM). The contents of the wells were thoroughly mixed by pipetting the solution up and down several times. After mixing, 180 μ L was transferred from wells in column 2 to the corresponding wells in column 3. The contents were Mixed and 180 µL was again transferred from each well into the corresponding wells of column 4. The Serially mix and transfer process was continued down the length of the microtiter plate until the wells of the last column (12) was reached. Once the final column is met, 180 µL of mixed bacteria and metal solution was discarded to create equal volumes of wells for the entire microtiter plate. The 96 well microtiter plate was then aseptically covered, and the lid and base of the plate was sealed and wrapped with paraffin wax to aid in decreased evaporation rates. The plates were placed onto a gyrorotary shaker set to 150 rpm and incubated overnight at 37°C. Plates were removed from the incubator

and their turbidity was determined using a SpectraMax Plus 384 Microplate described in section 2.1.7.

	1	2	3	4	5	6	7	8	9	10	11	12
Α	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC
В	GC	50.0	25.0	12.5	6.25	3.125	1.56	0.781	0.390	0.195	0.097	0.048
С	GC	50.0	25.0	12.5	6.25	3.125	1.56	0.781	0.390	0.195	0.097	0.048
D	GC	50.0	25.0	12.5	6.25	3.125	1.56	0.781	0.390	0.195	0.097	0.048
Е	GC	50.0	25.0	12.5	6.25	3.125	1.56	0.781	0.390	0.195	0.097	0.048
F	GC	50.0	25.0	12.5	6.25	3.125	1.56	0.781	0.390	0.195	0.097	0.048
G	GC	50.0	25.0	12.5	6.25	3.125	1.56	0.781	0.390	0.195	0.097	0.048
Н	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC

Figure 2.0: preparation of the antimicrobial challenge plate with highest metal challenge concentration being 50 μ M. Using a sterile 96 well microtiter plate, the following steps described in section (2.5.4) illustrate how to aseptically prepare the challenge plate for metal toxin exposure-response analysis. The above microtiter plate represents a serial two-fold dilution gradient of a single antimicrobial challenge plate. This plate outline is kept the same for all three metal challenges. Rows B-D consisted of three technical replicates of one *E. coli* strain, while rows E-G consisted of another strain. In total there were 4 antimicrobial challenge plates for each of the metal challenges AgNO₃, AgSD, and CuSO4. SC= sterility growth control (0.9% Saline), **GC**= Growth control of one *E. coli* strain.

	1	2	3	4	5	6	7	8	9	10	11	12
Α	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC
В	GC	256	128	64.0	32.0	16.0	8.00	4.00	2.00	1.00	0.50	0.25
С	GC	256	128	64.0	32.0	16.0	8.00	4.00	2.00	1.00	0.50	0.25
D	GC	256	128	64.0	32.0	16.0	8.00	4.00	2.00	1.00	0.50	0.25
Ε	GC	256	128	64.0	32.0	16.0	8.00	4.00	2.00	1.00	0.50	0.25
F	GC	256	128	64.0	32.0	16.0	8.00	4.00	2.00	1.00	0.50	0.25
G	GC	256	128	64.0	32.0	16.0	8.00	4.00	2.00	1.00	0.50	0.25
Η	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC	SC

Figure 2.1: preparation of the antimicrobial challenge plate with highest metal challenge concentration being 256 μ M. Rows B-D consisted of three technical replicates of one *E. coli* strain, while rows E-G consisted of another strain. The above microtiter plate represents a serial two-fold dilution gradient of a single antimicrobial challenge plate. In total there were 4 antimicrobial challenge plates for each of the metal challenges AgNO₃, AgSD, and CuSO₄. SC= sterility growth control (0.9% Saline), GC= Growth control. GC= Growth control of one *E. coli* strain.

2.5.4 Viable cell counts

Viable cell counts are a method used in microbiology to evaluate the number of cells capable of growing/dividing within a culture. The method which viable cell counts can be represented is through the plate count method. Microliter volumes of bacterial culture are aseptically pipetted onto the solid surface of an agar plate. The plates are incubated, and the resulting cultures are counted and the corresponding colony forming units per mL (CFU/mL) are generated through the formula:

(CFU/mL) = (number of colonies) x (dilution factor)

Volume of culture plate

The resulting values are used to graph the cell counts as a function to the metal concentration they were challenged with as a logarithmic unit. From these graphs the respective minimal bactericidal concentration (MBC) and minimal biofilm eradication concentration (MBEC) can be determined.

2.5.5 Preparation of antimicrobial challenge plate and manual counts of (CFU/mL) for viable cell counts

The methodology for preparing viable cell counts for planktonic and biofilm cells from an antimicrobial challenge plate (Figure 2.1) is the exact same to that of the optical density procedure highlighted in section 2.5.3. However, following the O/N growth (24 hours) the antimicrobial challenge plates were removed from the incubator and placed on the lab bench till ready. New 96 well microtiter plates were prepared containing 180 μ L of sterile 0.9% saline. Column 1 of the 96 well microtiter plates contained an addition of 20.0 μ L of universal neutralizer (0.5 g 1⁻¹ histidine (Sigma, St Louis, MO, USA), 0.5 g 1⁻¹ cysteine (Sigma) and 0.1 g 1⁻¹ reduced glutathione (Sigma) in ddH₂O) (used to bind free metal ions). To determine the CFU/mL of the distributed planktonic and biofilm cells (after sonication) bacterial populations, 11 dilutions with a dilution factor of 10 were performed. This was accomplished using a multichannel micropipette to transfer 20.0 μ L of culture from wells of column 1, which were mixed by pipetting up and down and transferred to the corresponding wells in column 2. The Serially mix and transfer process was continued down the length of the microtiter plate until the last column (12) was reached. Using a p20 pipette, 20μ L of the samples were then plated onto LB agar to determine the colony forming units and subsequently incubated overnight at 37°C.

2.5.7 Crystal violet assay

This Assay can be used for the indirect quantification of total biomass to determine differences in biofilm adherence to polystyrene pegs upon challenge with a metal antimicrobial agent (O'Toole, 2011). This method relies on the staining of attached cells with crystal violet dye, which binds to proteins and DNA. The difference in optical densities between treated and untreated mutant and wildtype biofilms will be quantified utilizing a spectrophotometer giving qualitative data on the differences in biofilm production and adherence of the different mutant strains biofilms (Table 2.0), when subjected to the three different metal antimicrobial challenges.

2.5.8 Crystal violet dye exposure and biofilm cultivation

Following the biofilm formation, outlined in section 2.1.9, and utilizing a similar crystal violet staining procedure to O'Toole, 2011, the 96 well microtiter plates containing wildtype and the 6-cell envelope BW25113 mutant *E. coli* strains (Table 2.0), which were exposed to a two-fold 50 μ M concentration gradient of the three metal antimicrobial challenges had dye plates prepared. The dye plates consisted of new sterile 96 well microtiter plates, which contained 150 μ L of 0.5 % crystal violet staining solution administered to each well of the microtiter plate. 12 dye plates were created for each of the corresponding CBD lids of the challenged wildtype and mutant *E. coli* biofilms. The CBD lids were placed into the dye plates and incubated at RT for 20 minutes on a bench rocker with a frequency of 20 oscillations per minute. The CBD lids were then inverted and placed on filter paper and tapped gently to remove any remaining liquid from the pegs. The CBD lids were then allowed to air-dry for 30 minutes at RT. The CBD lids were then placed into new 96 well microtiter plates containing 200 μ L of 30% (v/v) acetic acid (Biofilm's biomass cultivation plates). The lids were incubated in solution for 20 minutes at RT on a bench rocker with a frequency of 20 oscillations per minute. These plates were than taken to have their OD₅₇₀ read by SpectraMax Plus 384 Microplate reader. Following the plate reading,

the average OD_{570} of the wells without cells was subtracted from the OD_{570} of each well on the plate to eliminate background fluorescence picked up by the plate reader.

2.6. Microscopy

Three different microscopy techniques (CLSM, TEM, and SEM) were utilized to visualize differences in planktonic and surface biofilm morphology when in the presence and absence of AgNO₃.

2.6.1 Confocal laser scanning microscopy

The confocal laser scanning microscope (CLSM) is an optical imaging technique that captures multiple two-dimensional images at different depths in a sample, which enables for the reconstruction of three-dimensional structures within an object (Pawley, 2006). Given that microbes frequently live in multi or single species solid surface biofilms, which consist of architectural features important for population heterogeneity and emergent cell function (Harrison *et al.*, 2006). This CLSM technique offers an important insight into the understanding of biofilm morphology and the changes associated when an antimicrobial challenge is introduced (Harrison *et al.*, 2006). Bacterial biofilms have been widely studied and have been shown to modulate their architecture to changes in external and internal process when exposed to antimicrobial agents. Here in this experiment, the main goal was to visualize the morphological changes associated with each *E. coli* strain's biofilm when grown in the presence of a Ag⁺.

2.6.2 Microscopy biofilm production

Utilizing similar protocol to Harrison *et al.*, 2006, cryogenic stock stored at -80°C (Section 2.1.1), of the different bacterial strains (Table 2.0) was collected by a 200 μ L pipette tip and was inoculated 7 sterile 22.0 mL test tubes filled with 7.0 mL of M9 or LB media. The test tubes were incubated overnight for 16 hours at 37°C and 150 rpm. Following the O/N growth the cultures were sub-cultured under the same conditions. Following sub-culturing, 1.0 mL of culture was resuspended into fresh 7.0 mL M9 media and matched to 1.0 McFarland standard. Next, the cultures were then poured into reagent reservoirs and using a multichannel

micropipette, 180 μ L of the bacterial cultures were aliquoted into the wells of column 2, rows B-G, column 3 row B-2, column 4 rows B-G, and column 5, row B-4 of a sterile 96 well microtiter plate. The wells in column 4 rows B-G, and column 5, row B-4 acted at growth controls (GC) and were not subjected to any metal challenge. Furthermore, the remaining unused wells of the plate were aliquoted with 180 μ L of 0.9% saline, which acted as the sterile control (-). Using a multichannel micropipette, 180 μ L of concentrated 50 μ M AgNO₃ metal challenge was transferred into the wells of column 2 rows B-G and column 3 row B-2. The contents of the wells were thoroughly mixed by pipetting the solution up and down several times. After the wells received the proper inoculum volumes, a CBD lid was aseptically transferred to cover the 96 well microtiter plate. The plates were then placed onto a gyrorotary shaker set to 150 rpm and incubated for 24 hours at 37°C.

2.6.3 Conducting confocal laser scanning microscopy

The overnight growth plates (section 2.6.2) were removed from the incubator and were placed onto the lab bench. The CBD lid was removed and the corresponding polystyrene pegs with the different *E. coli* strains biofilms were aseptically broken off with alcohol flamed pliers. The pegs were then transferred to a new sterile 96 well microtiter plate. The pegs were rinsed in 200 μ L 0.9% saline twice and then placed into 200 μ L of 5.0% of glutaraldehyde PBS buffer and incubated at 37°C for 30 min. During the time of incubation, 0.1% acridine orange dye was prepared by adding 0.05 g of acridine orange in 50 mL of PBS buffer. The pegs were removed from the incubator and placed in 150 μ L of 0.1% acridine orange dye. The pegs were incubated for 15 min and wrapped in tinfoil as the stain is light sensitive. Once the incubation was finished, the pegs were washed again one more time in 200 μ L 0.9% saline. Following the last wash, the pegs were transferred to microscope slides using tweezers and a drop of ddH₂O was placed onto the pegs. The biofilms of the challenged and unchallenged pegs were then examined through Bitplane: Imaris© x 64 Microscopy image analysis software.

2.6.4 Transmission electron microscopy

The transmission electron microscope (TEM) is a powerful tool utilized in microbiology to investigate microorganisms through which electrons can pass through specimens and generate a projection image. Traditionally, TEM has been the "gold standard" for imaging dividing

microbial cells (Hazelton and Gelferblom, 2003). This microscopy method offers detailed information about *E. coli* cells ultrastructure.

2.6.5 Whole cell transmission electron microscopy preparation

Utilizing the cryogenic stocks of the different E. coli strains (Table 2.0) stored at -80°C (section 2.1.1), a 200 µL pipette tip was used to collect the different bacterial strains and inoculate 7 sterile 22.0 mL test tubes filled with 5.0 mL of M9 media. These test tubes were placed in an incubator set to 37°C and shaking at 150 rpm and incubated for 24-hours. After the incubation, 1.0 mL of the 5.0 mL cultures was removed and added into fresh 4.0 mL of M9 media. These test tubes served as the challenged cells samples (treated with AgNO₃). The remaining 4.0 mL of the overnight culture is kept and utilized as the negative control sample. The 4.0 mL was centrifuged at >10,000 x g for 2 minutes to produce a cell pellet and was stored at -20°C. The challenged test tubes received 25.0 µL of 10mM AgNO₃ stock solution to generate a final antimicrobial challenge of 50 μ M. These tubes were then allowed to incubate overnight for 16-24 hours at 37°C and 150 rpm. Following the incubation period and 30 minutes prior to imaging, the *E. coli* cells were spun down >10,000 x g for 2 minutes to obtain a cell pellet. Both control and challenge cell pellets were then resuspended in 5.0 mL of sterile ddH_2O . A 1/10 dilution was then performed, which required 100 µL of the resuspended cultures being transferred to a 1.5 mL microcentrifuge tube containing 900 µL of sterile ddH₂O. subsequently, $5.0-10 \mu$ L of each sample was placed onto the carbon side of the carbon-copper coated TEM grids. 5.0 µL of 1.0% phosphotungstic acid contrasting agent (used to improve image visibility) was added onto each sample, which were then left to air-dry for 10 minutes. Each carbon-copper grid was loaded into the TEM and imaged utilizing Gatan Digital Micrograph[©] and FEI microscope interface[®] at 80-100 -keV.

2.6.6 Scanning electron microscopy

The scanning electron microscope (SEM) uses a focused beam of high-energy electrons to generate a reflected image/signal at the surface of solid specimens. The signals that derive from electron-sample interactions reveal information about the sample particularly external morphology. This technique was utilized to observe the changes in physical characteristics of the *E. coli* biofilms when challenged by Ag^+ .

2.6.7 Sputter deposition/coating and fixing of E. coli biofilms for SEM imaging

Biofilm formation for SEM imaging followed the same procedural steps as section 2.6.2. The overnight growth plates were removed from the incubator are placed onto the lab bench. The CBD lid was removed and the corresponding polystyrene pegs with the different E. coli strains biofilms were aseptically broken off with alcohol flamed pliers. The pegs were then transferred to a new sterile 1.5 mL microcentrifuge tubes and placed in a BSC to air-dry for 24 hrs. The pegs were then transferred to microscope slides, which were placed into the coater glass chamber of a Hummer Sputter Coater. The voltage and the main power to the machine was turned on allowing for the creation of a vacuum system within the glass gas chamber. Once the vacuum's gage reached 50 millitorr (takes roughly 30 minutes), the valve for argon gas was opened until the pressure inside the chamber was 400 millitorr. The argon valve was then closed, and the glass chamber was again pumped to create another vacuum with a pressure of 50 millitorr. At this time, the high voltage control knob was set to 9 and turned on. The argon valve was reopened, and the gas flow was adjusted until the amp meter read 10 milliamperes. When a pinkish purple glow of argon plasma begins to emit from the top of the glass chamber, a timer of four minutes was set. At this time physical vapor deposition (PVD) is taking place, whereby atoms are ejected from a target or source material (platinum) that is to be deposited on a substrate (polystyrene pegs) (Bouzakis and Michailidis, 2014). Once the four-minute timer is reached, the samples are well coated in the platinum source material. The glass gas chamber is re-pressurized, and the samples are removed. The pegs are then transferred onto double sided carbon tape (improves imaging), which is placed on the sample stand inside the SEM chamber and were visualized using Phillips and Scandium© software at 120 -keV.

2.7 High throughput metal resistance profiling:

2.7.1 Disk diffusion assay

The purpose of the disk diffusion susceptibility test is to determine the sensitivity or resistance of aerobic bacteria to various antimicrobial compounds. 6-mm filter paper disk are impregnated with a known concentration of an antimicrobial compound and placed onto spread agar plates containing the different bacterial inoculum. immediately water is absorbed into the disk from the agar and the antimicrobial begins to diffuse into the surrounding agar. The rate of diffusion through the agar is not as rapid as the rate of extraction of the antimicrobial out of the disk, therefore the concentration of antimicrobial is highest closest to the disk and a logarithmic reduction in concentration occurs as the distance from the disk increases (Jorgensen and Turnidge, 2008). The rate of diffusion of the antimicrobial through the agar is dependent on the diffusion and solubility properties of the antimicrobial compound in the agar and its molecular weight (Bauer *et al.*, 1966). Larger molecules will diffuse at a slower rate than lower molecular weight compounds. These factors, in combination, result in each antimicrobial having a unique breakpoint zone (zone of inhibition) size, which indicates the susceptibility of a bacterial culture to that antimicrobial compound.

2.7.2 Preparation of Inoculum and placement of antimicrobial disks

Utilizing cryogenic stock stored at -80°C, a 200 μ L pipette tip was used to collect the different bacterial strains (Table 2.0) and inoculate 7 sterile 22.0 mL test tubes filled with 7.0 mL of M9 media. The test tubes were incubated overnight for 16 hours at 37°C and 150 rpm. Following the O/N growth of the pre-culture, the colonies were suspended in 0.9% NaCl to match the density of a 1.0 McFarland standard. Subsequently, 250 μ L of each mutant bacterial strain and wildtype strain were aliquoted onto fresh LB agar plates and spread evenly with an ethanol flamed glass spreader. Sterile antimicrobial disks, presoaked in each metal stock solution found in section 2.1.4 for 2 hours, were placed on the LB agar plates and incubated for 16 hours at 37°C. The region of bacterial susceptibility (zone of inhibition) was measured with a ruler from edge to edge across the zone of inhibition and recorded as a diameter value in millimeters.

Chapter Three: Results

3.1 Choice of cell envelope associated genes

Gene choice was based off the chemical genetic screen conducted by Dr. Lemire (unpublished results), which related mutant colony size to fitness and enabled identification of quantitative genetic interactions from high-throughput, genome-scale screen. The genetic screen evaluated 3985 single gene deletions of all non-essential gene mutants of the Keio collection, resulting in the discovery of 225 mutants to be sensitive to the exposure of the 100μ M AgNO₃ challenge and 177 mutants to display a resistant phenotype. The Sensitive ('sicker') or resistant ('healthier') mutants were selected based on a difference of 2 standard deviations of colony size from the normal, which was determined through photographed images that were processed using software that measures colony areas in terms of pixels. Of the total mutants, 20 sensitive and 4 resistant mutants were found to be involved in cell envelope homeostasis. The genes were then analyzed through bioinformatic and literature analysis, which evaluated their cellular function and involvement in cell envelope homeostasis. From these analysis of the 24 genes, one resistant (*damX*) and five sensitive mutants (*rodZ*, *minC*, *sanA*, *ybhO*, *and tolB*) where chosen to evaluate their collective phenotype in the presence of Ag⁺(Table 2.0).

3.2 Chromosomal disruptions

As a comparable biological analysis, mutations of genes rodZ, ispG, mreB, damX, minC, sanA, ybhO, and tolB in the genomic background of *E*. coli K-12 MG1655 were attempted in order to evaluate the discrepancy in Ag⁺ toxin response phenotypes when compared to the same mutants in Keio collection BW25113 strain.

3.2.1 MG1655*A*rodZ knockout mutant and Ag⁺ exposure-response

Utilizing λ Red-recombinase mediated gene replacement (section 2.2), and a standard protocol and methodology similar to Datsenko *et al.*, (2000), the $\Delta rodZ$ chromosomal deletion was generated by using several pairs of 56-to70-nt-long primers that included 18-to50-nt homology extensions and 20-nt priming sequences for helper plasmids pKD13 or pKD32 as template for amplification of their respective antibiotic cassettes (Table 2.1). Figure 3.0 demonstrates the respective ~1.29-kbp PCR product/replacement allele (gel lane: 6-10), which

represented successful annealing of the knockout primers (Table 2.2) to the Cm^{R} cassette from the template plasmid pKD32, which was further purified and transformed into a *E. coli* MG1655 cell already carrying pKD46, which is a low copy number easily curable plasmid encoding the Red-recombinase. This plasmid facilitated homologous recombination through a double crossover event with the ORF of the gene of interest (*rodZ*) and the replacement allele (Figure 3.0) effectively replacing the gene of interest with the FRT- Cm^{R} -FRT. Once transformed, successful recombinants with targeted chromosomal disruptions can be selected for by being plated on LB agar containing the appropriate selection marker; 1.25% (w/v) LB agar + 10 μ g/ml chloramphenicol. The Red recombinase plasmid pKD46 contains a temperature-sensitive origin of replication and is cured at growth at the non-permissive temperature of 37°C. The resulting transformants were then transformed with the temperature sensitive helper plasmid (pCP20) that transiently expressed the yeast recombinase, Flp. The successful removal of the antibiotic cassette facilitated by the flipase enzyme, is illustrated in Figure 3.1. MG1655 $\Delta rodZ$ cells were confirmed via PCR, using the OS and OAS confirmation primers (Table 2.2). mutants produced a band at approximately ~412bp (180 OS and ORF + 146 OAS and ORF). Unsuccessful deletion mutants generated a banding pattern equal to about 1014bp – about the size of the undeleted *rodZ* gene or the intact Cm^{R} antibiotic resistance cassette (1293bp). The mutants were further validated through Sanger sequencing.

1 Kb MW

1 Kb MW



Figure 3.0: Agarose gel to confirm amplification of the gene replacement allele using the template plasmids pKD13 and pKD32 and *rodZ* KO and RC primers (Table 2.2). The primers have 5' 18-50 nt-homology extension, which recognize *E. coli* K-12 MG1655 chromosome, as well having 3' primers with sequence homology that matches the template plasmid. Successful reactions (Lanes:6-10) produce a single clean PCR product that is ~1.29 (Cm^R) or ~1.4 kbp (Kn^R) (lane 5) in size relative to the antibiotic cassette from the template plasmids.

1 Kb MW



Figure 3.1: Confirmation of Flp-mediate excision of the FRT-flanked Cm^R antibiotic resistance cassette. MG1655 $\Delta rodZ$:: FRT- Cm^R -FRT insertion mutants were subjected to Flpmediated excision of the antibiotic resistance cassette by electroporating pCP20 into the positive insertion mutants. Flp excision of the antibiotic resistance (Cm^R) cassette was accomplished by incubating the transformants at 30°C overnight and then at 43°C following antibiotic selection. The deletion mutants were confirmed via PCR using the OS and OAS primers (Table 2.2). Clean mutants should produce a band at approximately 412bp (180 OS and ORF + 146 OAS and ORF). Unsuccessful deletion mutants will have a banding pattern equal to about 1113bp – about the size of the undeleted gene or the intact antibiotic resistance cassette (1293bp).

Following the creation of MG1655 $\Delta rodZ$ mutant *E. coli* cells, a comparative biological analysis of $\Delta rodZ$ mutants in the Keio collection strain background and the newly created MG1655 $\Delta rodZ$ was conducted to evaluate differences in pre-exposure-response (grown in the presence) to three metal challenges. The optical density at 600nm was utilized to evaluate differences in culture turbidity and cell morphology (Section 2.5.1). From Table 3.0, it is evident that $\Delta rodZ$ mutant cells were more susceptible to the three metal challenges compared to the wildtype of each strain based of its respective minimum inhibitory concentration (MIC). Moreover, the $\Delta rodZ$ mutant in the K-12 MG1655 strain was a full dilution more sensitive to AgNO₃ compared to the $\Delta rodZ$ Keio collection counterpart.

				-			
E. coli Strain	AgNO ₃	AgNO ₃	AgSD	AgSD	CuSO ₄	CuSO ₄	
Keio Collection	(µM/µL) MBC	(μΜ/μL) MIC	(µM/µL) MBC	(μM/μL) MIC	(µM/µL) MBC	(μΜ/μL) MIC	
BW25113							
BW25113	50.0	6.25	>50.0	6.25	>50.0	12.5	
BW25113ArodZ	25.0	6.25	>50.0	6.25	>50.0	12.5	
D ((20110A)002	23.0	0.25	20.0	0.25	20.0	12.5	
M01/75	25.0	10.5	. 50.0	2.12	. 50.0	2.12	
MG1055	25.0	12.5	>50.0	3.13	>50.0	3.13	
MG1655∆rodZ	12.5	6.25	50.0	3.13	>50.0	3.13	

Metal Challenge

 Table 3.0: Minimal Bactericidal Concentrations (MBC) and Minimal Inhibitory

 Concentrations (MIC)

*These values are averages calculated from three biological trials containing three technical replicates.

Regrettably, with the unsuccessful chromosomal disruptions of both *ispG* and *mreB*, resulted in a change of coarse for analyzing the collective phenotype of the selected genes. Reformulating the original comparative biology question that aimed to evaluate the changes exposure-response to metal challenges in different genomic backgrounds, I instead focused on characterizing the 6 Ag-responsive Keio collection mutants (Table 2.0) and their phenotypic responses to three metal antimicrobials.

3.2.2 MG1655*\DispG knockout mutant*

Following the same protocol, which generated the MG1655 $\Delta rodZ$ cells, *ispG* the downstream gene in the same operon as *rodZ* was targeted for chromosomal disruption. This gene was targeted to give insight into whether the exposure response phenotype of was a MG1655 $\Delta rodZ$ cells direct response from the chromosomal deletion or the from alteration in expression of downstream genetic elements. However, transformation and targeted replacement of *ispG*, resulted in unsuccessful generations of MG1655 $\Delta ispG$ transformants. Probing the

literature, it was suggested that *ispG* is an 'essential' gene that produces a [4Fe–4S] clustercontaining protein that is involved in isoprenoid synthesis in *E. coli* (Loiseau *et al.*, 2007). Thus, when comparing the validation that *ispG* is 'essential' under other experimental conditions, our experimental methods too suggested that in our growth conditions resulted in *ispG* being unable to be deleted from the chromosome.

<u>3.2.3 MG1655AmreB knockout mutant</u>

Utilizing the same Red recombinase protocol for rodZ and ispG, the gene, mreB, an associated genetic factor whose protein product functions in close association to RodZ was targeted. However, transformation and targeted replacement of mreB, resulted in unsuccessful generations of MG1655 $\Delta mreB$ transformants. Investigating literature, mreB is described as a quasi-essential (meaning can be removed is substituted nutrient is provided) gene in (Baba *et al.*, 2006) and is crucial in determining cell shape in rod-shaped bacteria (Bacilli) (Curtis, 2016). In comparison here it was suggested, that in under our experimental conditions it too was concluded mreB was unable to be deleted from the *E. coli* chromosome.

3.3 Plasmid based complementation and comparative analysis

Following the successful generation of the chromosomal gene disruption mutant K-12 MG1655 $\Delta rodZ$, plasmid based complementation of K-12 MG1655 $\Delta rodZ$::pMS119EH_A::rodZ was conducted through the use of restriction enzymes and T4 ligase (sub-cloning). This method aimed to restore the original wildtype phenotype in response to metal challenges. Utilizing molecular cloning primers (Table 2.2) rodZ was successfully amplified from genomic DNA and ligated into the Amp-resistant expression vector pMS119EH_A generated by Beketskaia *et al.*, 2014, through the use of restriction digestions, T4 Ligase, and antibiotic selection plates.

Following successful plasmid based complementation, the pMS199EH_A::*rodZ* plasmid was transformed into K-12 MG1655 Δ *rodZ* cells and subjected to metal exposure-response assay aimed to generate a comparative analysis between wildtype, Δ *rodZ*, and pMS199EH_A::*rodZ* complemented cells when grown in the presence of AgNO₃, AgSD, and CuSO₄. When analysing the resulting phenotypes, it was determined from Figure 3.3, that in the presence of AgNO₃ challenge, both Δ *rodZ* mutant and complemented strain were more susceptible to AgNO₃ by two dilutions. Evaluating the phenotypic response of each strain to the AgSD metal antimicrobial, all three strains recorded similar optical densities, however, the complemented MG1655 $\Delta rodZ$::pMS119EH_A::rodZ cells achieved similar turbidity to that of the wildtype strain when compared to the mutant MG1655 $\Delta rodZ$. Finally, assessing the CuSO₄ toxin response curves, which served as a comparison antimicrobial metal, resulted in all three cultures achieving similar optical densities at 600nm. Furthermore, at the experimental conditions implemented during the metal exposure-response assay, it appears that at the given antimicrobial 2-fold concentration gradient, CuSO₄ reduced culture density of the three *E. coli* cultures the least as evident from the evaluated turbidity of the samples.



Figure 3.2: Showing agarose gel containing the products of the ligation reaction of *rodZ* insert (1,014 bp) into the low copy number Amp-resistant expression vector pMS119EH_A (3,967bp). Successful ligation (A) produced a product band roughly ~4,981 bp (3,967bp pMS119EH_A + 1,014 bp *rodZ*) in size (lane 5 and 6). The subsequent digestion of the successful ligation products in lanes 5 and 6 in the top gel can be visualized in the (B) gels lane 2. After digestion from restriction enzymes HindIII and XbaI, two bands representing pMS119EH_A (3,967bp) and (1,014 bp) *rodZ* are present.



Figure 3.3: Metal exposure-response curves of the gene deletion mutant K-12 MG1655*ArodZ*, the plasmid based complementation transformant

MG1655 Λ rodZ::pMS119EH_A::rodZ, and wildtype K-12 MG1655 *E. coli* cells. The three bacterial strains were subjected to 16-hour growth in the presences of three different metals antimicrobial challenges AgNO₃, AgSD, and CuSO₄ (Table 2.2). The average OD₆₀₀ of three biological trials each containing three technical replicates was recorded utilizing a SpectraMax Plus 384 Microplate Reader. The solid black line represents K-12 MG1655 cultures, the dotted black line represents K-12 MG1655 Δ rodZ, and dotted red line represents MG1655 Δ rodZ::pMS119EH_A::rodZ.

<u>3.4 Bacterial growth curves</u>

As detailed in section 3.2.1, with the unsuccessful chromosomal deletions of ispG and *mreB* in the genomic background of K-12 MG1655, I reformed my approach and aimed to characterize the Ag⁺ phenotypic responses of the targeted 6 cellular envelope associated genes in the genomic background of the Keio collection, BW25113 strain. Before conducting metal exposure-response analysis of the Keio collection cellular envelope associated Ag-responsive genes towards three chosen metal antimicrobials growth curves for the individual K-12 BW25113 mutant strains ($\Delta rodZ$, $\Delta damX$, $\Delta minC$, $\Delta sanA$, $\Delta ybhO$, $\Delta tolB$) (Table 2.0) and wildtype E. coli cells was conducted. The 7 strains were inoculated into sterile M9 media and subjected to optical density readings by SpectraMax Plus 384 Microplate Reader (section 2.1.7) at 1, 2, 4, 6, 8, 10, 12, 16, 18, 20, 24, and 48-hour time intervals. Each strain was subjugated to three biological trials, having three technical triplicates when recording their changes in cellular density over time. This methodology aimed to reduce discrepancy and increase accuracy of the culture turbidity readings at each designated time interval. Figure 3.4 illustrates the average OD₆₀₀ of 3 technical trials and 3 biological replicates of each of the 7 K-12 BW25113 strains. From the analysis of the growth curve in Figure 3.4, it was inferred that all strains reached mid log (or exponential) phase at around the 12-hour mark with all cultures transitioning into stationary phase around 16 hours of growth. However, one striking observation that was made from all the mutant growth curve cultures during the assay, was each had a prolonged *lag phase* when compared to the wildtype culture. Thus, it was suggested that with deletion in genes associated with cell envelope homeostasis, that mutant cells have a considerably more challenging time synthesizing the enzymes and factors needed to facilitate cell division and population growth resulting in prolonged *lag phase*. Figure 3.5, further illustrates a comparative analysis of two mutant strains ($\Delta damX$ and $\Delta rodZ$) growth curves being compared to wildtype cells, which both displayed similar cellular density at each time interval, except at the 6-hour mark. With each mutant strain generating similar cultural turbidity through out the assay, the remaining growth curves can be found in the appendix B section.



Figure 3.4: Growth curve average of 3 biological trials each containing 3 technical trails, which measured the OD₆₀₀ by SpectraMax Plus 384 Microplate Reader of the 7 different K-12 BW25113 *E. coli* strains (Table 2.0).



Figure 3.5: Growth curve of the average of 3 biological trials each containing 3 technical trails, which measured OD₆₀₀ readings of BW25113 Δ *rodZ* and BW25113 Δ *damX* being compared to wildtype BW25113 cells. The results confirmed that all strains achieved a similar growth pattern, however, mutants appeared to have a prolonged *lag phase*.

3.5 Cell viability Assay (CFU/mL) for planktonic E. coli cells

In order to determine planktonic E. coli cell viability of the 6 cell envelope associated mutants and wildtype cells after growing in the presence to the three metal antimicrobials in M9 media, viable cell counts were conducted on 1.25% (w/v) LB agar. Utilizing the formula in section 2.5.4, colony forming units (CFU) per mL of bacterial culture was evaluated for three separate biological trails and expressed in logarithmic notation (Log-kill CFU/mL) (Figure 3.6). These CFU/mL calculations accurately depict the number of viable cells in each culture suspension and precisely determined the metal concentrations required to prevent planktonic growth (Minimal Bactericidal Concentration, MBC) of the respective E. coli cells. Moreover, the resulting colonies sizes were also analyzed for discrepancies between the mutants and wildtype cells, which resulted in all 7 strains having comparable colony size (appendix D). Examining the planktonic cell viability results, AgNO₃ proves again to be the most efficacious metal antimicrobial resulting in cellular death at much lower concentrations then the other two comparable metals. Examining the silver resistant phenotypes in the presence of AgNO₃, $\Delta damX$, $\Delta sanA$, and $\Delta ybhO$ were found to be a full dilution more resistant to the Ag⁺ challenge when compared to the wildtype cells. The remaining 3 mutants, had similar MBC values to that of the wildtype cells, which were only permitted to grow at AgNO₃ concentrations less than 16.0 µM. When evaluating the CFU's/mL for the AgSD challenge, it was shown that BW25113 $\Delta damX$ was 3 full dilutions more resistant to the AgSD antimicrobial challenge in comparison to the wildtype culture. At a dose concentration of 256.0 μM, ΔdamX mutant BW25113 E. coli cells were still viable, which far exceeded the MBC value of any other Ag-responsive mutant strain. All the other mutants, however, displayed a resistant phenotype compared to the wildtype cells in the presences of AgSD, except for $\Delta ybhO$, which had the same MBC as the wildtype cells, which were unable to grow at concentrations of 64.0 µM and higher. Lastly, evaluating the CFU's/mL produced in the presence of CuSO₄, $\Delta rodZ$, $\Delta minC$, and $\Delta ybhO$ produced the same MBC phenotype as the wildtype cells, being unable to grow at concentrations of Cu²⁺ higher than 128.0 μ M. The other mutants $\Delta damX$ and $\Delta sanA$ produced a resistant phenotype, while $\Delta tolB$ was a full dilution more susceptible to the toxicity of Cu²⁺ when compared to the wildtype cells. Table 3.1, highlights the metal concentrations of each antimicrobial required to prevent planktonic growth of each strain of E. coli. Furthermore, Figure 3.6, demonstrates a generated log-kill CFU/mL challenge curve from metal exposure-response assay for the damX mutant and

the wildtype BW25113 cells. The remaining cell viability curves for each mutant can be found in the appendix D section, along with photographs of one of the three biological trails for $AgNO_3$ treatment spot plated onto 1.25% (w/v) LB agar.



Figure 3.6: log-kill CFU/mL generated for K-12 BW25113 Δ damX and wildtype *E. coli* planktonic cells exposed to three antimicrobial metals AgNO₃, AgSD, and CuSO₄. The Δ damX cells were exposed to a 256 μ M two-fold antimicrobial gradient (Figure 2.0) for a 24-hour period. These cultures were then diluted in dilution plates and 20 μ L of culture was aliquoted onto 1.25% (w/v) LB agar plates. CFU/mL was calculated by multiplying the number of viable bacteria colonies by its respective dilution factor, divided by the total volume of the culture (180 μ L). The CFU/mL was then expressed using logarithmic notation.

3.5.1 Cell viability (CFU/mL) for E. coli biofilms

To determine *E. coli* biofilm viability of the 6 cell envelope associated mutants and wildtype biofilm grown in Lysogeny broth media after post-growth exposure to the three metal antimicrobials, viable cell counts were conducted on 1.25% (w/v) LB agar. Utilizing the formula in section 2.5.4, colony forming units (CFU) per mL of bacterial biofilm was evaluated for three separate biological trails. These CFU/mL calculations accurately depict the number of viable cells in each culture suspension and precisely determined the metal concentrations required to prevent Biofilm growth (Minimal Biofilm Eradication Concentration, MBEC) of the respective E. coli cells. The biofilms were grown in accordance to section 2.1.9. and sonicated of the polystyrene pegs of the CBD lid in to determine the CFU/mL. From the biofilm viability results, AgNO₃ was demonstrated to be the most efficacious of the three metal challenges. The resistant determinant, $\Delta dam X$, biofilm was a full dilution more resistant to the AgNO₃ challenge, while sensitive mutants, $\Delta rodZ$ and $\Delta tolB$, were one dilution more susceptible when compared to the wildtypes biofilm cells viability. The remaining mutants had similar MBEC values to that of the wildtype culture. Analyzing the MBEC values for the AgSD challenge, $\Delta damx$, $\Delta minC$, $\Delta sanA$, and $\Delta ybhO$ were more resistant to the metal challenge when compared to the wildtype. $\Delta tolB$ was found to be more sensitive to the AgSD challenge compared to the wildtype, while $\Delta rodZ$, had a similar MBEC value. Finally, analysing the comparator metal CuSO₄, only $\Delta tolB$ was found to be more susceptible to the metal challenge in comparison to all other metals and wildtype biofilms. Table 3.1 shows the respective MBEC values generated from the biofilm viability assay, which display similar MBEC values to that of publish literature (Lemire et al., 2017). These MBEC end-point values were utilized for generating a challenge concentration, which was utilized for the different microscopy techniques throughout this thesis.

<i>E. coli</i> Strain Keio Collection BW25113	AgNO3 (µM/µL) MBC	AgNO3 (µM/µL) MBEC	AgSD (µM/µL) MBC	AgSD (µM/µL) MBEC	CuSO4 (µM/µL) MBC	CuSO4 (µM/µL) MBEC
wildtype	16.0	64.0	64.0	64.0	128.0	256.0
∆damX	32.0	128	>256.0	256.0	256.0	256.0
ΔrodZ	16.0	32.0	128.0	64.0	128.0	256.0
∆ <i>minC</i>	16.0	64.0	64.0	128.0	128.0	256.0
AsanA	32.0	64.0	256.0	256.0	256.0	256.0
ΔybhO	32.0	64.0	256.0	128.0	256.0	256.0
ΔtolB	16.0	32.0	128.0	32.0	64.0	128.0

 Table 3.1: Minimal Biocidal Concentrations (MBC) and Minimal Biofilm Eradication

 Concentrations (MBEC)

Metal Challenge

*There was no variability between end-point values (MBC and MBEC) for each mutant and wildtype planktonic and biofilm cells with the established experimental conditions.

3.6 Microscopy

Utilizing three different microscopy techniques (CLSM, TEM, and SEM), planktonic and surface biofilm physiology in the presence and absence of a 50 μ M metal antimicrobial challenge (AgNO₃) were characterized. 50 μ M was chosen based of the representative MBEC and MBC values found from the cell viability assay. However, complication arising from both SEM and CLSM experimental procedures generated inconclusive results

3.6.1 Transmission electron microscopy analysis of treated and untreated *E. coli* biofilms

TEM analysis of 50 μ M pre-treated 1.0% phosphotungstic acid stained *E. coli* bacteria showed the external planktonic morphological features of the cell envelope associated mutant BW25113 strains, wildtype BW25113 and MG1655 cells, and the recombinase generated MG1655 Δ rodZ cells. Images were processed and captured with Gatan Digital Micrograph© and FEI microscope interface© (section 2.6.5) at 80-100 keV. The untreated BW25113 and MG1655 *E. coli* cells of wildtype and mutant strains retained their bacillus morphology and appeared normal in solution. However, MG1655 Δ rodZ mutants, commonly displayed elongated bacillus morphology, which contrasted the wildtype MG1655 cells. However, this change in cell morphology present in TEM imaging of MG1655 Δ rodZ cultures did not influence optical density measurements.

All the *E. coli* strains treated with the $50 \,\mu$ M AgNO₃ challenge appeared to have cells in culture undergo lysis, resulting in the release of cellular contents into the surrounding environment. It was common to find electron-dense particles (Figure 3.7) or precipitates surrounding damaged cells, which were electron translucent in comparison to undamaged cells. When examining the cellular envelope of the treated and untreated *E. coli* strains, significant morphology changes were observed. Untreated cells, seemed to have normal phenotypical characteristics. The internal structures appeared to be normal, with some displaying their multilayered cell envelope consisting of the outer membrane a peptidoglycan layer in the periplasmic space, and the inner cytoplasmic membrane (Figure 3.8). On the contrary, it was routinely observed that treated cells had a compromised cellular envelope (Figures 3.7 and 3.8, Images B and D). Treated cells displayed either localized or complete separation of the cell membrane from the cell wall. Furthermore, plumes of cellular debris with accompanied electron-translucent cytoplasm and cellular disruption was observed in treated cells. Even the Ag⁺

resistant determinant, $\Delta damX$, could not withstand the given Ag⁺ ion toxicity in solution and had similar planktonic morphologies as to that of all the other genes and wildtype treated cells (Figure 3.7). During TEM analysis, it was also noted that mutant *E. coli* strains had a drastic reduction in cellular density, making imaging of full populations of cells difficult to achieve. Even with centrifugation steps aimed to increased cell density, resulted in low planktonic cell concentrations. More TEM photos at different magnifications are present in the appendix E section of this thesis.


$\Delta dam X(-Ag)$

 $\Delta damX (+Ag 50 \mu M)$

Figure 3.7: External cell morphologies of 1.0% phosphotungstic acid stained wildtype and BW25113 $\Delta dam X E.$ coli planktonic cells untreated (A and C) and pre-treated with 50 μ M AgNO₃ treated (B and D) observed by transmission electron microscopy. Images A and B, are wildtype cells imaged at 15000x mag, and images C and C, were BW25113 $\Delta dam X$ cells taken at 9600 x magnification. Red arrows indicated electron-dense particles that were commonly found in treated TEM samples and areas of cell membrane separation. Some treated cells displayed either localized or complete separation of the cell membrane from the cell wall. plumes of cellular debris were also witnessed with accompanied electron-translucent cytoplasm and cellular disruption.



Figure 3.8: Internal cell envelope structures of planktonic of 1.0% phosphotungstic acid stained wildtype and BW25113 *E. coli* cells untreated (A and C) and pre-treated with 50 μ M AgNO₃ (B and D) observed by Transmission electron microscopy.

3.6.2 Silver nanomaterial structure

In Figure 3.9, it is observable that a wildtype BW25113 and MG1655 $\Delta rodZ$ treated cells have synthesized electron-dense nanomaterials in the form of a pyramid or nanoprism (1-100 nm) in close proximity to its cellular envelope. In conjunction to these observations, Raheem *et al.*, (2011), also reported on finding similar Ag (I) nano structures when exposing the *E. coli* strain (ATCC 8739) to (1% v/v) of aqueous AgNO₃ in Luria and Burrous broth solution. Furthermore, other types of silver nanomaterials including spheres, cubes, and nanorods were also observed throughout TEM imaging of the treated *E. coli* cells (Appendix E). It appears that under the imposed procedural conditions, that the *E. coli* cells synthesize triangular silver nanoparticles by chemical reduction method through the reduction of silver nitrate in the presence of a reducing agent.



Figure 3.9: Wildtype and MG1655 Δ *rodZ* cells with an electron-dense nanoparticle in the shape of a pyramid or nanoprism associating with its cellular envelope.

3.7 Disk diffusion assay

In order to provide a comparison to the susceptibility of each cell envelope mutant of E. *coli* to other metal antimicrobials, a disk susceptibility test was conducted. Initial experimental concerns expressed included the formation of metal complexes with the components of the agar media resulting in limited diffusion, nullifying the potential for this simple assay. We confirmed that all metals diffuse through the media agar producing growth inhibition zones, by measuring the resulting translucent areas without noticeable bacterial growth surrounding the antimicrobial infused disks. In the thesis research, we observed that all the bacterial strains were susceptible to the metal-salts in the tested concentrations (section 2.1.4). Figures of one of two biological trials, which were utilized for the measurement of growth inhibition zones (where growth was inhibited by dissociation of metal-salt into its respective antimicrobial ion) can be found in the appendix F section of this thesis. These measurements were averaged in millimeters (mm) and used for comparison between wildtype and mutant strains. Several metals, such as; Zn²⁺, Ni²⁺, and Cu²⁺, appeared to be the most efficacious antimicrobials at the tested concentrations, indicative of the larger zones of growth inhibitions (Table 3.2). BW25113 $\Delta damX$ was less susceptible to Ag⁺, Zn^{2+} , and Ni²⁺ toxicity when compared to wildtype cells, but generally had the same susceptibility to all other metals tested. Moreover, BW25113 $\Delta rodZ$ was found to be more susceptible to Ni^{2+} and Zn^{2+} exposure when compared to all other strains.

Table 3.2: Detection of metal toxicity by use of disk diffusion susceptibility test
illustrating displaying the zones of inhibition

Challenges									
Strains	NaCl	NaNO ₃	NaSO ₄	Ga(NO ₃) ₃ •H2O	ZnSO ₄ •7H ₂ O	AgNO3	NiSO ₄ • 6H ₂ O	Al ₂ (SO ₄) ₃ •H2O	CuSO ₄
BW25113	0	0	0	11.5	24.0	11.0	23.8	15.0	19.1
BW25113∆damX	0	0	0	12.00	22.8±0.8	9.60±0. 1	21.5±0.3	15.0	18.5
BW25113∆rodZ	0	0	0	12.2	26.0±0.2	11.2	26.6±0.4	16.0	20.3
BW25113∆minC	0	0	7.00	13.4±0.8	21.5±0.3	11.0	23.3	14.5	19.8
BW25113∆sanA	0	0	0	11.50	23.0	10.2	23.3	15.3	18.0
BW25113⊿ybhO	0	0	0	13.0±0.1	23.4	10.4	23.1	15.2	18.5
BW25113∆tolB	0	0	0	15.0±0.2	25.0	10.5	23.8	15.5	20.6

* Values are represented as the mean \pm the SD and if no change in SD \pm 0.00, n=2,. † Diameter (mm) of the zone of growth inhibition.

Chapter Four: Discussion

4.1 Metal exposure-response of the respective cellular envelope associated genes.

Silver has been known for its antimicrobial properties throughout antiquity (Akram *et al.*, 2016; Jung et al., 2008; Lemire et al., 2013). However, the mechanism by which Ag⁺ imposes its toxicity is still not fully understood (Clement and Jarret, 1994). Recent literature has demonstrated that *E. coli* and *S. aureus* microorganisms treated with Ag⁺ ions, display a severely compromised cell envelope resulting in cell lysis and separation of the cell membrane from the cell wall (Jung et al., 2008). Investigating this uncharacterized toxicity mechanism, data was collected from Dr. Lemire's chemical genetic screen (unpublished), which provided genetic analysis of the Keio collections ordered mutant library. This assay presented phenotypic analysis of the libraries non-essential genes in the presence of 100 µM of AgNO₃ infused into M9 minimal noble agar media. Analyzing colony size as a proxy for fitness, Dr. Lemire discovered over 225 mutants found to be sensitive to the exposure of the 100 µM AgNO₃ challenge while 177 mutants displayed a resistant phenotype. Of these genetic determinants, 20 sensitive and 4 resistant mutants were found to be involved in cell envelope homeostasis. From these 24 genes, 6 were chosen through further bioinformatic analysis and literary searches for their involvement in cell envelope homeostasis. Of the 6, 1 determinant was resistant in Dr. Lemire's screen, damX, the other 5; rodZ, minC, sanA, ybhO, and tolB, (Figure 4.0) were found to be sensitive.



Figure 4.0: The identified silver resistant and sensitive genes protein product and their relative cellular roles within the organism *E. coli*. Contents of the cell are not to scale. Notation defines genes that are silver sensitive or resistant as determined in this study. PG: Peptidoglycan, OMP: Outer membrane porin.

To confirm the respective sensitive and resistant phenotypes of these genetic determinants, I set out to characterize their collective phenotype when subjected to a silver antimicrobial challenge. Utilizing metal-exposure response I aimed to determine the relative minimal bactericidal concentration and minimal biofilm eradication concentration of each of the mutant's planktonic and biofilm cell cultures against three metals. Before conducting this analysis, certain experimental parameters were considered. Despite the abundance of literature committed to examining the anti-microbial activity of metals, less attention has been paid to determining the susceptibility of bacteria to metals within a defined set of conditions (Gugala et al., 2017). Although the minimal inhibitory concentration, minimal bactericidal concentration and minimal biofilm eradication concentrations for many metals have been determined (Harrison et al., 2004), the lack of consistency between techniques, conditions and media has resulted in difficulties when comparing the susceptibilities of bacterial strains to metal compounds. Here, M9 minimal media was utilized for the growth curve analysis, metal exposure-response assays and microscopy analysis. The decision to utilize minimal media rather a rich medium like Lysogeny broth, was based of ionic bonding chemistry. With the establishment that antimicrobial activity of Ag (I) based compounds being intrinsically dependent on the formation of the Ag⁺ ions (Walker and Parsons, 2014), there was concern that high NaCl concentration associated with LB media, would cause for the dissociating Cl⁻ ions and cause the precipitation of Ag⁺ as AgCl, as well as complexation of silver in diverse soluble chlorocomplexes and chelation with molecules within the LB. This concern was further validated by investigations into literature, which suggested that high Cl⁻ concentrations in the exposure medium indeed have a pivotal influence on the resulting toxicity of a Ag(I) based antimicrobial, appearing to significantly attenuate toxicity (Gadd et al., 1989; Groh et al., 2015). However, when trying to accumulate sufficient biomass of the mutant's strains biofilms for examination during the metal exposureresponse assay, it was found that M9 media did not support measurable optical density. This determination was also supported by literary sources that described the BW25113 E. coli strain to be motility-impaired in minimal media, when compared to the MG1655 strain, due to the differences in the expression of motility loci, qseB, flhD, fliA, fliC, and motA (Wood et al., 2006). Wood et al., (2006) found that qseB expression, which is responsible for the activation of the flagella regulon, was 139-fold higher in the MG1655 strain than in BW25113 (Sperandio et al. 2002; Wood et al., 2006). They further suggested that the differentiation in expression has led

to both reduced motility and deficiency of flagellar apparatuses in the BW25113 strain, making them "poor" biofilm producers. Although measurable amounts of biofilm were unachievable through spectroscopy methods, when the biofilms grown in M9 media and were visualized during different microscopy techniques, it was confirmed that each mutant could grow a thin confluent biofilm. This was important finding, for this determination allowed for viable cell counts of BW25113 biofilms to be conducted and respective MBEC values to be determined for each mutant (Table 3.2).

The resulting MBC and MBEC findings from the metal exposure-response assay, suggested that the single gene deletions of $\Delta damX$, $\Delta sanA$, and $\Delta ybhO$, in E. coli K-12 BW25113 strain withstood a higher concertation of Ag(I) metal challenge when compared to the wildtype cultures. This finding proposes the idea that E. coli cells lacking these genes and their respective transcribed protein products are more resistant to Ag⁺ toxicity. To understand why the deletions of these genes conferred relative Ag⁺ resistance resulted in analysing their cellular function and possible association with Ag⁺ ions. The cell, which showed the highest levels of resistance had a *damX* gene deletion. This gene is predicted to code for a bitopic inner membrane protein, which is involved in cell division; as it contains a C-terminal SPOR domain, which targets the protein to the septal ring (Gerding et al., 2009). Utilizing both YFP-DamX and GFP-DamX, Gerding and colleges confirmed that the DamX protein localizes sharply to division sites in wildtype E. coli cells. Further, DamX has been shown through a two-hybrid system to interact strongly to essential cell division proteins FtsQ and FtsN, who are required for septum formation and mid-cell localization of the Z-ring (Arends et al., 2010). Moreover, that same assay also suggested that DamX has weak interactions with FtsZ, which is responsible for forming the contractile ring Z-ring structure (Arends et al., 2010). Investigating into the amino acid composition of the DamX protein resulted in no thiol groups (R-SH) being evident, which have been demonstrated to be favorable binding targets for silver and the energetically favorable bonding of SH-Ag has been shown to result in protein deactivation (Klueh et al., 2000; Rai et al., 2012). However, the DamX protein contains 22 charged amino acid residues of arginine (R). Recent literature has suggested that of all the amino acids, arginine has the highest affinity towards silver ions (Shoeib et al., 2002), and binds silver via chelation and complexation mechanisms (Agnihotri et al., 2015). A study supporting this finding conducted by Agnihotri et al., 2015, demonstrated that ZnO nanorods served as nano-reactors for in situ synthesis of silver

nanoparticles concurrently providing a template for their subsequent immobilization using arginine as a linker (Agnihotri *et al.*, 2015). Thus, from evidence proposed in the literature it is suggested here that the presence of the arginine in the DamX protein could be immobilizing free Ag⁺ onto its surface, which could inhibit DamX's role in cellular division leading to silver sensitivity in cells containing the $\Delta damX$ gene deletion.

With the findings from two-hybrid protein interactions in literature, it is suggested that DamX could also directly link the peptidoglycan layer to the Z-ring. If confirmed, and if Ag⁺ can be shown to be sequestered and immobilized on the DamX protein by its arginine residues, then silver ions could impose their toxicity on the Z-ring and the formation of the septum in dividing cells. The Z-ring is essential step required for cellular division, with any interruption of its formation process leading to the interruption of cellular growth (Hwang and Lim, 2015). Consequently, *E. coli* cells that have the chromosomal deletion $\Delta damX$, have their cellular division and growth less effected by Ag⁺ binding to DamX, making them more tolerant to the silver challenge than the wildtype *E. coli* cells.

The DamX amino acid sequence was further subjected for analysis utilizing Phyre2 (Protein Homology/AnalogY Recognition Engine) a web-based service for protein structure prediction (Kelley *et al.*, 2015). The resulting 3-D model was generated based of techniques that utilize homology modeling, resulted in the SPOR domain being identified with 99.9% confidence and 97% identity. The model further provided secondary structures of the predicated biotic inner membrane protein, that hypothesized the presence of arginine residues being present in both the cytosol and periplasmic space. Thus, adding more speculation to DamX possibly having silver ions immobilized on its periplasmic surface by arginine residues.

The other two resistant determinants validated by MBC and MBEC values, $\Delta sanA$ and $\Delta ybhO$, were found to be more resistant to the exposure of the silver-based antimicrobials when compared to the wildtype cultures. These findings contradict the initial Ag-sensitive phenotype that was observed from the chemical genetic screen. Possible explanations for this discrepancy in phenotype, could have resulted from the difference in the growth mediums utilized. The chemical screen was conducted on solid M9 minimal noble agar infused with 100 μ M of AgNO₃, where as the metal exposure-response experiments in this thesis were carried out in M9 liquid broth medium, present in 96 well plates with cultures grown in the presence of 256 μ M of silver

metal challenge, then subsequently plated on to LB agar plates for colony viability analysis. Expositions from literature, have suggested that the rate of diffusion of silver ions in agar gel is much slower than that of liquid broth medium (Kant, 1962). Thus, the agar infused AgNO₃ plates could have different Ag⁺ saturation throughout the plate, complexation, Ag⁺ dissociation and diffusion rates, that resulted in the reporting of a sensitive MBC for cells containg the gene deletions $\Delta sanA$ and $\Delta ybhO$, when grown on the solid media in comparison to liquid or aqueous mediums. Similar susceptibility findings have been reported in literature where discrepancy between concentration of metals in media can lead to different susceptibility phenotypes. Kumar *et al.*, (2013), reported that the Gram negative rod-shaped *S. marcescens* organisms displayed higher MIC values to transition metals on solid Mueller-Hinton agar then in comparison to liquid medium. Therefore, the possible unequal distribution of Ag⁺ throughout the M9 noble Agar could be the contributing factor for the different Ag- phenotypic responses observed.

Nonetheless, the gene *sanA* has been identified to code for the sole member of the vancomycin-sensitivity protein family in the Transporter Classification Database (Saier et al., 2016). SanA is predicated to be a bitopic inner membrane transport protein, roughly 239 amino acids in size. Unlike DamX, SanA does contain cysteine amino acid residues within its protein sequence, which have been demonstrated empirically through the Hard-soft acid base theory, that soft acids like Ag(I) have electron-sharing affinities that can result in the energetically favourable formation of covalent bonds between soft bases, such as the sulphhydryl groups present on the amino acid, cysteine (Gordon et al., 2010; Harrison et al., 2007; Leung et al., 2013). Thus, Ag^+ ions could be binding to the sulphhydryl groups present within the SanA protein inhibiting its function. Another observation from literature that could possibly support this associated Ag-resistant phenotype in $\Delta sanA E$. coli cells, was the determination that silver enhances the antimicrobial efficacy of the antibiotic, vancomycin (Esmaeillou et al., 2017; QingShan et al., 2006). QinShan and colleagues investigated vacomycin-capped silver nanoparticles and examined their antibacterial activities in vitro against E. coli. They reported from microscopy TEM images, that single bacteria treated with the vacomycin-capped silver nanoparticles, had plenty of aggregates in the cell wall of the E. coli organism (QingShan et al., 2006). QinShan et al., (2006) proposed that a possible antibacterial mechanism of the silver nanoparticles was to destroy or destabilize the membrane, allowing for vancomycin to easily bind to the terminal d-Ala-d-Ala dipeptide inhibiting peptidoglycan biosynthesis (Ge et al.,

1999). In addition to this proposed mechanism, other work conducted by Mitchell *et al.*, (2017) predicted that SanA is involved in OM permeability to sodium dodecyl sulfate (SDS) in carbonlimited cells. Thus, it is proposed if Ag^+ can bind and inhibit SanA function, there would be subsequent changes in membrane permeability. This would account for $\Delta sanA$ mutants having different membrane properties that could account for relative resistance to the Ag^+ challenge in respect to the wildtype cells.

The other resistant genetic determinant, ybhO, codes for an enzyme involved in cardiolipin synthesis. Cardiolipin is a unique phospholipid which was historically identified as a major lipid in the inner mitochondrial membrane of cardiac muscle cells (Paradies et al., 2014). Considerable progress has recently been made in understanding the role of cardiolipin in mitochondrial function and bioenergetics (Paradies et al., 2014). It has been shown that cardiolipin is associated with membranes designed to generate an electrochemical gradient that is used to produce ATP, such as bacterial plasma membranes and inner mitochondrial membrane (Schlame et al., 2000). Literary inferences have suggested that cardiolipin seems to participate in the structural organization and stabilization of the respiratory chain complexes (Electron transport chain, ETC or oxidative phosphorylation) in high order structure of functional importance and play a significant role in the formation of the membrane curvature (Pfeiffer et al., 2003; Schägger, 2002). Its role in ETC functionality, however, has been shown to have deleterious effects if in the presences of reactive oxygen species (ROS)-induced oxidation. Interestingly, this determination that ROS induces the depletion of cardiolipin in the membrane, relates to Ag⁺ induced toxicity. With the establishment that Ag⁺ ions have the ability to disrupt iron-sulfur clusters [4Fe-4S] in proteins (Xu et al., 2009; Xu and Imlay, 2012) by binding to the sulphhydryl groups, it has been suggested that this binding induces the release of Fe²⁺ from these proteins. It has thus been suggested that this release of Fe²⁺ might account for observations that the Fenton-inactive Ag, could generate ROS within a bacterial cell (Park et al., 2009; Xu et al., 2009). Therefore, if Ag⁺ ions are causing the generation of ROS within E. coli cells, then cardiolipin would be subjected to its toxicity, resulting in deleterious effects upon the ETC. Thus, cells that have the chromosomal deletion $\Delta ybhO$, could have a reduction in cardiolipin synthesis, possibly causing the resistant phenotype in comparison to wildtype cells, for the ROS has less of a lethal effect on the $\Delta y bhO E$. coli cells for their membrane physiology has changed and their ETC are not being directly targeted.

In contrast to the to the Ag-resistant determinants, the gene deletions of *rodZ* and *tolB*, resulted in BW25113 E. coli cells to become more susceptible to the silver and copper metal antimicrobial challenges. Thus, suggesting that the protein products of these genes aid in generating relative tolerance to the toxicity of the metal challenges in the wildtype cells. The rodZ gene codes for a bitopic inner membrane protein that has been identified to be involved in cell wall synthesis and retaining cellular morphology (Morgenstein et al., 2015). Bacterial cell shape is structurally determined by the rigid peptidoglycan built outside the cytoplasmic membrane by a series of cell wall assembling enzymes (Typas *et al.*, 2012). In many bacillus, or rod-shaped species, the enzymes have been showing to be coordinated by an actin-like protein, MreB (Morgenstein et al., 2015), with its polymeric nature being identified as the key determinant in retaining the rod shape. However, it has remained difficult to explain the mechanistic link between cell wall growth and MreB located in the cytosol of the bacteria cell. Morgenstein and colleagues however, provided substantial evidence through functional GFP-RodZ fusion, demonstrating that RodZ and MreB are colocalized within E. coli cells and located together around the cells circumferences. They further inferred that RodZ has a periplasmic tail that is important for cell shape and could interact with penicillin binding proteins and a cytoplasmic tail that directly interacts with MreB, thus linking the MreB to protein(s) in the periplasm (Bendezú et al., 2009; Morgenstein et al., 2015; van den Ent et al., 2010). Therefore, it is suggested that cells having the $\Delta rodZ$ gene deletion have a reduction in cell wall biosynthesis and stabilization. Without RodZ linking MreB to the periplasm, proteins could be assembling in a non-orderly manner resulting in decreased cell wall integrity. With compromised structural integrity of the cell wall, it is thought that these $\Delta rodZ E$. coli cells would have a lower MBC and MBEC values than wildtype cells, which was confirmed from the metal exposure-responses experiments within this thesis.

The other Ag-sensitive genetic determinant, *tolB*, codes for a periplasmic protein, which participates in the trimeric porin assembly of OmpF, OmpC, PhoE, and LamB (Rigal *et al.*, 1997) and is involved in outer membrane stability (Lazzaroni *et al.*, 1999). The interaction between TolB proteins and outer membrane stability and porin assembly may be the direct cause for the resulting Ag⁺ sensitive phenotype. Recent literature, conducted by Randall *et al.*, 2015, demonstrated that in *E. coli* endogenous resistance arose to silver as a consequence of two-point mutations in *ompR* and *cusS* resulting in loss of the OmpC/F porins. Thus, confirming that silver

resistance in *E. coli* can be associated with the depressed expression of the OmpC/F outer membrane porins (Randall et al., 2015). This silver resistance phenotype was also reported by Radzig et al., 2009, who demonstrated that E. coli MC4100 strain cells with chromosomal deletions for the genes encoding OmpF and OmpC, were found to be three to four times more resistant to silver ions than the wild-type strain. Radzig and colleagues further suggested, that their data indicates that these porins may be involved directly or indirectly with the movement of silver ions into the cell. Furthermore, literary insight provided by Lazzaroni et al., (1999) has also described that *tolB* mutants in *E. coli* cells demonstrated an increase in OmpC porins (Lazdunksi et al., 1998; Lazzaroni et al., 1999; Misra and Reeves, 1987). Thus, it is suggested here that increased number of OmpC porins from the gene deletion of $\Delta tolB$, could increase the amount of intracellular Ag⁺ accumulation, resulting in the sensitive phenotype that was observed throughout the metal exposure-response experiments in this thesis. Moreover, this explanation provides further insight into the increased susceptibility of BW25113\[Delta to the Cu²⁺] antimicrobial challenge. According to Elger et al., (2005), who investigated the pathways of copper uptake by *E. coli* cells, they have suggested that intracellular accumulation of Cu^{2+} is facilitated through the OmpC and OmpF porins. They further showed that increased expression of OmpC can lead to increased intracellular concentration of metals. Therefore, it is suggested that $\Delta tolB$ genetic deletions generate E. coli cells that are both more susceptible to copper and silver antimicrobials as a result of the amplified OmpC porin protein production and through destabilizing of the OM causing 'leakiness' of Ag⁺ through the OM.

The last genetic determinant, *minC*, is involved in ensuring the correct placement of the septum during cell division, suppressing formation of the septa in aberrant loci within the cell (Pichoff and Lutkenhaus, 2001). Evaluating the resulting MBC and MBEC values for the BW25113 Δ *minC* cells, saw only the biofilm cells being more resistant to the AgSD challenges when compared to the wildtype. Furthermore, BW25113 Δ *minC* cells achieved similar MBC and MBEC values to that of the wildtype cells for the other two metal antimicrobials. One potential cause for the resulting resistant phenotype, was the presence of cysteine residues within the MinC protein. As discussed, Ag⁺ ions have been shown to bind sulphhydryl groups with in proteins, resulting in their inactivation (Xu *et al.*, 2009; Xu and Imlay, 2012).

Comparing the toxicity of Ag^+ ions to the planktonic and biofilm cells of each mutant strain, it is evident that biofilms offered cells increased tolerance to the Ag^+ and Cu^{2+} challenges. Bacterial biofilms are sessile microbial communities frequently imbedded in a EPS matrix (Garret *et al.*, 2008). These robust structures offer bacterial cells protection from persistent environmental pollutants (Teitzel and Parsek, 2003). Teitzel and Parsek (2003), demonstrated that when *P. aeruginosa* (PAO1) strain cells were assayed with heavy metals, free-swimming planktonic cells were more susceptible to the metal challenge in comparison to biofilms cells. This similar trend was found in this thesis, where generated planktonic cells MBC values for each mutant, were lower then the compared biofilms cells MBEC values. Moreover, the MBC and MBEC values were similar to values found in the literature, where Harrison *et al.*, (2004) assayed *E. coli* JM109 strain cells against silver ions in Lysogeny broth medium, generating a MBC and MBEC values of 0.09 ± 0.04 and 0.07 ± 0.02 mM respectively.

4.2 Effects of Ag⁺ on E. coli planktonic cell morphology

Although the mechanisms underlying the antibacterial actions of silver are still not fully understood, several previous inferences from the literature (McDonnell and Russell, 1999; Pal et al., 2007) have suggested that the interaction between silver ions and the constituents of the bacterial cell envelope caused structural changes, damage to the membranes and intracellular metabolic activity which might be the cause or consequence of cell death. Thus, suggesting that genetic determinants involved in cell envelope homeostasis could be direct or indirect targets involved in Ag⁺ ions-imposed toxicity mechanism. Investigating the morphological changes of the mutant BW25113 planktonic cells in the presence of a 50 µM AgNO₃ challenge, was observed through the utilization of TEM. Each of the E. coli strains resulted in similar phenotypes with the unchallenged cells conforming to normal bacillus cellular morphology, while treated cells had aberrant cellular morphology and extensive cell envelope damage and apparent cell lysis. It was also observed that untreated cells had visible intact internal structures with both the IM, periplasm, and OM visible through the use of TEM (Figure 3.18). This observation too has been made in the literature where both Jung et al., (2008) and Yamanaka et al., (2005) both reported that intracellular structures of E. coli were observable utilizing TEM. Jung et al., (2008) further showed that both E. coli and S. aureus cells treated with 0.2 ppm Ag⁺

solution displayed either localized or complete separation of the cell membrane from the cell wall. Jung *et al.*, (2008), further concluded that electron-dense particles or precipitates were also observed around damaged bacterial cells, which were also observed in the TEM treated cells within this thesis. Anlysisng the TEM images, however, concluded that there was no difference in phenotype of treated cells in both wildtype and mutant cells. Thus, suggesting that other genetic determinants or cellular mechanisms must be involved in Ag^+ cell envelope toxicity mechanism.

The TEM images from my study here further lead to the observations of the BW25113 and MG1655 E. coli cells synthesizing Ag nanoparticles (AgNP) in the shape of spheres, pyramids or nanoprisms. Comparable observations have also been recorded in the literature where Shanshoury et al., (2011) observed E. coli strain ATCC 8739 cells, treated with aqueous AgNO₃ in nutrient broth could biosynthesize Ag-nanoparticles that appeared spherical in shape and appeared to be reasonably monodispersed. Shanshoury et al., (2011), further concluded that the nanoparticles ranged in size of 5-25 nm in diameter. These findings have important implications for possible 'biosynthesis' of Ag nanoparticles, which has been actively pursued in recent years as an alternative, efficient, inexpensive, and environmentally safe method for producing nanoparticles with specified properties. Currently, physical and chemical processes are widely used to synthesize metal nanoparticles, which allow one to obtain particles with the desired characteristics (Okitsu et al., 2007). However, these production methods are usually expensive, labor-intensive, and are potentially hazardous to the environment and living organisms (Narayanan and Skthivel, 2010). Thus, investigations into "Green" nanoparticle development that is environment friendly, cost effective and easily scalable is becoming highly sought after. Thus, the mutants and strains of E. coli evaluated here may turn out to be excellent biofactories to produce AgNPs. Optimization of culturing and exposure conditions could lead to control over nanomaterial shapes and characteristics.

4.3 Conclusion

MDR bacterial pathogens have become a serious challenge for healthcare professionals (Bartlett et al., 2013). With the reduction in new investment, focused on developing new novel antibiotic medicines, the subsequent 'drying up' of the development pipeline for new antibiotics, has begun to exacerbate the problem of MDR bacteria (Bartlett *et al.*, 2013). Research scientists

must now begin to investigate the efficacy of alternative antimicrobial compounds to be considered as methods of treatment or to be used synergistically alongside other antibacterial medicines in the treatment of these MDR infections (Allen *et al.*, 2014). These new investigations are now focusing on uncovering the complex toxicity mechanisms by which these alternative antimicrobial agents function.

This thesis utilized data from a chemical genetic screen, which was utilized to investigate genetic determinants, to provide an extensive genomic analysis of possible cellular targets by which silver imposes its toxicity onto *E. coli* cells. It was demonstrated that the growth of planktonic and biofilm cells of both BW25113 and MG1655 *E. coli* strains were affected when grown in the presence of Ag^+ and Cu^{2+} , with cells being susceptible to silver toxicity at micro molar concentrations. The exposure to Ag^+ resulted in morphological changes in planktonic cells, causing extensive cell envelope damage. Biofilm morphology could not be characterized due to the presence of artifacts and complication arising from both SEM and CLSM experimental procedures generated inconclusive results.

Many different mechanisms of Ag(I) toxicity have been suggested in the literature, from its ability to induce oxidative stress through the generation of reactive oxygen species and causes damage to cellular components including DNA damage, activation of antioxidant enzymes, depletion of antioxidant molecules (i.e., glutathione), and binding and disabling of proteins (McShan *et al.*, 2014). Moreover, it is also apparent from many observations from the literature and here from this thesis work, that Ag^+ ions cause extensive damage to the cellular envelope of Gram-negative bacteria. We see that functions around the cell wall and transport across the cell membranes have roles in both silver resistance and sensitivity (summarized in Figure 4.0). This data provides support to the idea that silver appears to be a multi-target antimicrobial by imposing its toxicity on multiple biochemical pathways such as those involved in cell wall homeostasis within the *E. coli* organism.

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Appendix A: Plasmid Based complementation

Includes intermediate steps necessary to carry out plasmid-based complementation of the gene *rodZ*.

<u>A.1 Complementation: genomic DNA isolation, ligation to the InsTAclone cloning vector, and restriction enzyme digestion.</u>

Following the successful generation of the chromosomal gene disruption mutant K-12 MG1655 $\Delta rodZ$, plasmid-based complementation was conducted through the use of restriction enzymes and T4 ligase. The first step was to isolate and amplify the rodZ gene from E. coli k-12 MG1655 genomic DNA with designed PCR primers (Table 2.2). Utilizing gel electrophoresis, the amplified PCR product was identified through size comparison with a 1 Kb molecular DNA ladder. Successful amplification produced a band relative to the size of the rodZ gene (1014bp). Figure B-1, illustrates the 0.7% (w/v) agarose gel generated with the respective rodZ amplified PCR product in well 3. Well 2 acted as the negative control, containing only the PCR primers and no genomic DNA added to the PCR reaction. Well 3 contained digested pMS199EH_A, which was stored at -20 °C for later ligation. The rodZ PCR product was purified and ligated into the InsTAclone cloning vector, pTZ57R/T. This vector offered one-step cloning of rodZ PCR product by exploiting terminal transferase activity of *Taq* DNA polymerase (Zhou, 2000). *Taq* polymerase has non-template dependent activity, which preferentially adds a single adenosine to the 3'-ends of a double stranded DNA molecule, and thus most of the molecules PCR amplified by Taq polymerase possess single 3'-A overhangs (Zhou, 2000). The use of a linearized " pTZ57R/T -vector", which has single 3'-T overhangs on both ends allowed for direct, highefficiency cloning of the *rodZ* PCR product, facilitated by complementarity between the PCR product 3'-A overhangs and vector 3'-T overhangs. The pTZ57R/T vector, also offered for easy quantitative analysis for recognizing successful ligation. Since successful ligation of desired gene fragment with pTZ57R/T lead to the disruption of lacZ gene activity resulting in lack of ability to cleave X-gal and when transformed into DH5a E. coli cells and grown on 1.25% (w/v) LB +/-Amp +/- X-gal at 37°C, 20-50 single white colonies were produced, which were picked and qualitatively examined (Figure B-2). Figure B-3, demonstrates the successful ligation of the purified rodZ PCR product to the pTZ57R/T-vector. The pTZ57R/T::rodZ was isolated from transformed DH5α E. coli cells and using a 1Kb molecular DNA ladder the PCR product band sizes were visualized. Analyzing the gel in Figure B-3, wells 3, 4, and 5 correspond to the successful ligation between insert (rodZ) and the TA vector, which resulted in a band shift relative to the negative control well 2, which only contained the circular pTZ57R/T vector from the Thermo Fischer Scientific kit and no insert. The successful ligation should result in a product

band around ~3,900bp (2,886bp TA vector +1,014bp *rodz* ORF). The PCR products in wells 3,4, and 5 were purified (section 2.1.7) and subjected to digestion by restriction enzymes XbaI and HindIII . Figure B-4, demonstrates the successful digestion of the pTZ57R/T::*rodZ* resulting in two bands (Well 3) corresponding to the TA vector (2,886bp) and *rodz* (1,014bp) ORF. The negative controls in well 2 and 5 are the circularized TA vector and the undigested pTZ57R/T::*rodZ* respectively. The *rodz* (1014bp) ORF was purified and utilized for ligation into the Amp- resistant expression vector pMS119EH_A generated by Beketskaia *et al.*, (2014). The presence of *rodZ* was again confirmed by running the purified 2nd band from well 3 of Figure B-4 on a 0.7% (w/v) agarose gel to ensure *rodZ* ORF was successfully isolated (Figure B-5).



Figure A-1: A agarose gel illustrating the successful amplification of the *rodZ* **gene through PCR from the genomic DNA of K-12 MG1655** *E. coli.* Well 2 contains the negative control where no genomic DNA was added to the PCR reaction, while well 3 contains the successful amplified product, which corresponds to the size of the *rodZ* gene (1,014bp). Well 4 contains digested (XbaI and HindIII) Amp- resistant expression vector pMS119EHA, which was utilized for plasmid-based complementation.


Figure A-2 Illustrates the 1.25% (w/v) LB +/-Amp +/- X-gal plates used for the identification of successful ligation of pTZ57R/T::*rodZ* plasmids, which was transformed into DH5*a E. coli* cells. Plates were grown overnight at 37°C and plasmids were isolated using commercial kit (section 2.1.8) These plasmids were then subjected to gel electrophoresis (section 2.1.9) for qualitatively analysis (Figure B-3).



Figure A-3: A agarose gel demonstrating the successful ligation of the *rodZ* insert to the InsTAclone (Thermo Fischer Scientific) cloning vector pTZ57R/T. Well 2 consists of the negative control circularized pTZ57R/T and corresponds to a band size of 2887bp. Wells 3, 4, and 5 represent successful ligation for their corresponding product band is roughly band around \sim 3,900bp (2,886bp TA vector +1,014bp *rodz* ORF).



Figure A-4: Demonstrates the digestion of pTZ57R/T:*rodZ* by restriction enzymes XbaI and HindIII. Well three product bands highlight the TA (2,886bp) vector (top band) and the *rodz* (1,014bp) ORF (bottom band). Wells 2 and 5 consist of the negative controls used in the comparison of the digested pTZ57R/T::*rodZ* product.



Figure A-5: The successful transformation of pMS119EH_A::*rodZ* into K-12 MG1655 Δ *rodZ E. coli* cells. K-12 MG1655 Δ *rodZ* cells are unable to grow in the presence of +100 µg/mL *Amp*, unless having acquired the Amp-resistant expression vector pMS119EH_A::*rodZ*. These cells were cultivated and utilized for cryogenic stock, which was stored at -80°C.



Figure A-6: A agarose gel containing the purified *rodZ* inserts (1014bp) extracted from the digestion reaction of pTZ57R/T::*rodZ* by restriction enzymes XbaI and HindIII. This gel confirmed that infact the lower band corresponded to the *rodZ* ORF and that ligation of the insert into the Amp- resistant expression vector pMS119EH_A could continue.



Figure A-7: Plasmid map of the InsTAclone cloning vector pTZ57R/T (from Thermo Fisher Scientific). The pTZ57R/T cloning vector is linearized and ddT tailed for direct use in cloning of PCR products (section 2.3.6), generated with *Taq*, or other DNA polymerases or polymerase mixtures, which add extra adenines to the ends of PCR products. The map and the MCS region of the vector are presented in Figure 2.1 showing where XbaI and HindIII restriction enzyme sites are located. (Prathyumnan *et al.*, 2011).



Figure A-8: Plasmid map of *Amp***-resistant expression vector pMS119EH**_A (Beketskaia *et al.*, 2014) utilized for plasmid complementation analysis.

Appendix B: ΔOD₆₀₀ vs Time

Includes the remaining four *E. coli* BW25113 cell envelope mutant's growth curve analysis.



Figure B-1: Growth curve average of three biological each containing 3 technical trails, which measured OD₆₀₀nm readings of BW25113 $\Delta minC$, $\Delta sanA$, $\Delta ybhO$, and $\Delta tolB$.

Appendix C: Metal exposure response optical density values

Contains the remaining metal exposure-response data for the *E. coli* BW25113 mutants subjected to two-fold dilution challenge plates of three metal antimicrobials. the accumulation of cellular density was measured through optical density measurements and was proxy for observing sensitivity and resistance and change in cellular morphology.



Figure C-1: Demonstrates the exposure-response relationship between K-12 BW25113 mutant's and K-12 BW25113 wild type planktonic cells, subjected to a 50 µM two- fold dilution antimicrobial gradient prepared on 96 well microtiter challenge plates (Figure 2.3) containing the three metal antimicrobials: AgNO₃, AgSD, and CuSO₄.



Figure C-2: Demonstrates the exposure-response relationship between K-12 BW25113 mutant's and K-12 BW25113 wild type planktonic cells, subjected to a 256 µM two- fold dilution antimicrobial gradient prepared on 96 well microtiter challenge plates (Figure 2.4) containing the three metal antimicrobials: AgNO₃, AgSD, and CuSO₄.



Figure C-3: Demonstrates the crystal violet assay of the BW25113 mutant's and wildtype biofilm cells grown for 24 hours in M9 media. The $\Delta damX$ cells were exposed to a 256 μ M two-fold antimicrobial gradient (Figure 2.4) of the three antimicrobial metals.

Appendix D: Viable cell counts

Contains images of one biological trial of the viable cell counts for the *E. coli* BW25113 mutant's planktonic cells. Moreover, the log CFU/mL data representing planktonic cell viability in the presences of three metal antimicrobials is also present.



Figure D-1: One of three biological trails of viable cell counts for the 7 *E. coli* strains challenged by AgNO₃ that were spot plated (20 μ L) onto 1.25% (w/v) LB agar.



Figure D-2: One of three biological trails of viable cell counts for the 7 *E. coli* strains challenged by AgSD that were spot plated (20 μ L) onto 1.25% (w/v) LB agar.



Figure D-3: One of three biological trails of viable cell counts for the 7 *E. coli* strains challenged by CuSO₄ that were spot plated (20 μ L) onto 1.25% (w/v) LB agar.



Figure E-4: log CFU/mL values generated for K-12 BW25113 mutant and wildtype *E. coli* cells exposed to three antimicrobial metals AgNO₃, AgSD, and CuSO₄. The $\Delta rodZ$ cells were exposed to a 256 μ M two-fold antimicrobial gradient (Figure 2.4). These cultures were then diluted in dilution plates and 20 μ L of culture was aliquoted onto 1.25% (w/v) LB agar plates (Figure 2.5). The CFU/mL was calculated by multiplying the number of viable bacteria colonies

by its respective dilution factor, divided by the total volume of the culture (180 μL). The CFU/mL was then expressed using logarithmic notation.

Appendix E: TEM Microscopy

Contains the comparison of TEM images of BW25113 wildtype and mutant's planktonic cells subjected to $\pm 50 \ \mu M \ AgNO_3$ challenge.



Figure E-1: External planktonic morphologies of 1.0% phosphotungstic acid stained wildtype BW25113 *E. coli* cells untreated (A, C, and E) and 50 µM AgNO₃ treated (B, D,

and F) observed by TEM. Images A and B, are imaged at 5000x mag, images C and D, were taken at 7000x mag, and images E and F, were taken at 15000x mag. Red arrows indicated electron-dense particles that were commonly found in treated TEM samples and areas of cell membrane separation. Some treated cells displayed either localized or complete separation of the cell membrane from the cell wall. plumes of cellular debris were also witnessed with accompanied electron-translucent cytoplasm and cellular disruption.



Figure E-2: External planktonic morphologies of 1.0% phosphotungstic acid stained BW25113 $\Delta minC$ E. coli cells untreated (A) at 7800x magnification and 50 µM AgNO₃ treated (B) at 9600x magnification, observed by TEM. In untreated sample BW25113 $\Delta minC$ confines to normal bacillus morphology. In contrast, treated BW25113 $\Delta minC$ appear to have complete disruption and compromise of the cellular envelope.



Figure E-3: External planktonic morphologies of 1.0% phosphotungstic acid stained BW25113 Δ sanA E. coli cells untreated (A) and 50 μ M AgNO₃ treated (B) observed by TEM at 7800 x magnification. In untreated sample BW25113 Δ minC confines to normal bacillus morphology. In contrast, treated BW25113 Δ sanA appear to have complete disruption of the cellular envelope.



Figure E-4: External planktonic morphologies of 1.0% phosphotungstic acid stained BW25113 Δ ybhO E. coli cells untreated at (A) and 50 µM AgNO₃ treated (B) observed by TEM at 7800 x magnification. In untreated sample BW25113 Δ ybhO confines to normal bacillus morphology. In contrast, treated BW25113 Δ ybhO appear to have localized disruption of the cellular envelope with plumes of cellular debris being released from cell (red arrow).



Figure E-5: External planktonic morphologies of 1.0% phosphotungstic acid stained BW25113 $\Delta tolB~E.~coli$ cells untreated (A) and 50 μ M AgNO₃ treated (B) observed by TEM, at 19000 x magnification. In untreated sample BW25113 $\Delta tolB$ confines to normal bacillus morphology. In contrast, treated BW25113 $\Delta tolB$ appear to have complete disruption of the cellular envelope.



Figure E-6: External planktonic morphologies of 1.0% phosphotungstic acid stained wildtype K-12 MG1655 *E. coli* cells untreated (A) and 50 µM AgNO₃ treated (B) observed **by TEM, at 4000 x magnification and 7000X magnification respectively.** In the untreated sample, wildtype K-12 MG1655 cells appear to have a normal phenotype. In contrast, the treated cells displayed either localized or complete separation of the cell membrane from the cell wall.



Figure E-7: External planktonic morphologies of 1.0% phosphotungstic acid stained K-12 MG1655 $\Delta rodZ$ *E. coli* cells untreated (A) and 50 μ M AgNO₃ treated (B) observed by TEM, at 5000 x magnification and 6500X magnification respectively. In the untreated cells, some of the mutant MG1655 $\Delta rodZ$ cells displayed a mutant elongated bacillus phenotype, which was commonly found throughout the sample. In contrast the treated cells displayed either localized or complete separation of the cell membrane from the cell wall.

Appendix F: Disk diffusion assay

Contains images of one biological trial of the high throughput toxicological assay aimed to determine the sensitivity or resistance of the *E. coli* BW25113 ag-responsive mutants to various antimicrobial compounds.

Disk Number	Corresponding Challenge
1	NaNO ₃
2	Na_2SO_4
3	Ga(NO ₃) ₃ ·H ₂ O
4	CuSO ₄
5	$ZnSO_4$
6	Saline (NaCl)
7	$Al_2(SO_4)_3$
8	$NiSO_4 \cdot 6H_2O$
9	AgNO ₃

 Table F-1: Indicating the appropriate metal infused into sterile Sterile antimicrobial disks

*Metals were prepared following section 2.1.4.



Figure F-1: Illustrates the disk diffusion assay conducted on wildtype BW25113 cells with the metals present in Table J-1.



Figure F-2: Illustrates the disk diffusion assay conducted on wildtype BW25113 Δ *damX* cells with the metals present in Table J-2.



Figure F-3: Illustrates the disk diffusion assay conducted on wildtype BW25113 Δ *rodZ* cells with the metals present in Table J-2.



Figure F-4: Illustrates the disk diffusion assay conducted on wildtype BW25113 Δ *minC* cells with the metals present in Table J-2.



Figure F-5: Illustrates the disk diffusion assay conducted on wildtype BW25113 Δ SanA cells with the metals present in Table J-2.



Figure F-6: Illustrates the disk diffusion assay conducted on wildtype BW25113 Δ *ybhO* cells with the metals present in Table J-2.


Figure F-7: Illustrates the disk diffusion assay conducted on wildtype BW25113 Δ *tolB* cells with the metals present in Table J-2.