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Survival in amoeba- a major selection pressure on the presence of bacterial copper and zinc resistance determinants? Identification of a ‘copper pathogenicity island’

Xiuli Hao^a, Freja L. Lüthje^b, Yanan Qin^b, Sylvia Franke McDevitt^c, Nataliya Lutay^d, Jon L. Hobman^e, Karishma Asiani^e, Fernando C. Soncini^f, Nadezhda German^g, Siyu Zhang^{b,h}, Yong-Guan Zhu^{a,h} and Christopher Rensing^{a,b*}

^aKey Lab of Urban Environment and Health, Institute of Urban Environment, Chinese Academy of Sciences, Xiamen, People’s Republic of China

^bDepartment of Plant and Environmental Science, University of Copenhagen, Frederiksberg, Denmark

^cDepartment of Biology, Skidmore College, Saratoga Springs, NY U.S.A.

^dDivision of Dermatology and Venereology, Department of Clinical Sciences, Lund University, Lund, Sweden

^eSchool of Biosciences, The University of Nottingham, Sutton Bonington Campus, Sutton Bonington, LE12 5RD, United Kingdom

^fInstituto de Biología Molecular y Celular de Rosario (IBR-CONICET, UNR) 2000 Rosario, Argentina

^gDepartment of Pharmaceutical Sciences, School of Pharmacy, Texas Tech University Health Sciences Center, Amarillo, Texas

^hState Key Lab of Urban and Regional Ecology, Research Center for Eco-Environmental Sciences, Chinese Academy of Sciences, Beijing 100085, People’s Republic of China.

*to whom correspondence should be addressed to:

Christopher Rensing

Department of Plant and Environmental Science

University of Copenhagen

1871 Frederiksberg, Denmark

chres@plen.ku.dk

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Abstract

The presence of metal resistance determinants in bacteria usually is attributed to geological or anthropogenic metal contamination in different environments, or associated with the use of antimicrobial metals in human healthcare or in agriculture. While this is certainly true, we hypothesize that protozoan predation and macrophage killing are also responsible for selection of copper/zinc resistance genes in bacteria. In this review we outline evidence supporting this hypothesis, as well as highlight the correlation between metal resistance and pathogenicity in bacteria. In addition, we introduce and characterize the “copper pathogenicity island” identified in *E. coli* and *Salmonella* strains isolated from copper and zinc fed Danish pigs.

Introduction

Essential metals such as iron and copper can cycle between different oxidation states and are used in metalloenzymes that catalyze electron transport reactions. Zinc also plays a major structural and catalytic role in metalloenzymes, and has been reported to counter oxidative stress. But in excess all of these metals are deleterious to cells. To ensure their own survival prokaryotes have developed mechanisms of maintaining cellular Zn^{2+} and Cu^+ homeostasis, while eukaryotes invented very original Zn and Cu binding structures not present in prokaryotes. Such structures allow accumulation of Zn^{2+} in the intracellular organelles followed by its utilization in biological processes specific for a given cell type. In particular, macrophages employ Zn^{2+} and Cu^+ to attack Fe-S clusters essential for bacterial survival (Dupont *et al.*, 2011; Festa and Thiele 2012; Macomber and Imlay 2009; Neyrolles *et al.*, 2015; Subashchandrabose *et al.*, 2014, Xu and Imlay 2012). We hypothesize that such a mechanism, where bacterial killing occurs through accumulation of Zn^{2+} and Cu^+ in the phagosome/vacuole, originated in protozoa long before multicellular life arose and that it later evolved in eukaryotic phagocytes. Our hypothesis is supported by the presence of the homologous copper transporter 1 (Ctr1) in macrophages and P80 in *Dictyostelium discoideum* and *Acanthamoeba castellanii* – proteins, both of which are involved in Cu^+ uptake upon phagocytosis. In addition, amoebae are known to contain P-type ATPases (German *et al.*, 2013; Burlando *et al.*, 2002), and, similar to macrophages, at least one of these P-type ATPases in *A. castellanii* could be pumping Zn^{2+} or Cu^+ into the phagosome of amoeba (Figure 1). Importantly, our hypothesis explains selection of genes involved in conferring copper and zinc resistance not only by the presence of these metals in the environment, but by protozoan predation as well. Since these determinants would aid survival in both protozoans and macrophages one could expect a higher occurrence of additional copper and zinc resistance determinants in virulent bacteria.

A copper silver resistance cluster- or an ancestral defense to phagosomal killing using copper?

Genome sequencing projects have revealed that several strains of *Salmonella enterica* subspecies *enterica* harbor a ca. 12 kb copper-resistance locus. This cluster is shown to either form part of a Tn7-like transposon inserted at the 3' end of the gene that encodes a NAD-utilizing dehydrogenase on the chromosome (Peters *et al.*, 2014), as found in isolates from serovars Heidelberg, Montevideo, Senftenberg or Tennessee; or to be a part of a larger integrating conjugative element inserted at the *pheV* phenylalanine tRNA, as present in strains of the serovars Senftenberg, Ohio or Cubana. Although there is a history of copper and zinc resistant bacteria being isolated from feces of animals fed with a metal supplement containing diets, presence of this resistance cluster has just recently been recognized.

Recently, we have sequenced the genomes of two *Escherichia coli* and three *Salmonella enterica* serovar Typhimurium strains isolated from copper and zinc fed Danish pigs, hence, displaying high level copper resistance (Lüthje *et al.*, 2014; Qin *et al.*, 2014). One of the *E. coli* strains and all three of the *Salmonella* strains contained a specific 19-gene mobile genetic element that we have named as the “Copper pathogenicity island”. In the *E. coli* isolate we have identified this island as a part of Tn7-like transposon, while in the *S. Typhimurium* strains it forms part of an approximately 80-kbp chromosomal element inserted at the *pheU* phenylalanine tRNA, similar to that identified in Heidelberg, Montevideo, Senftenberg or Tennessee isolates (Qin *et al.*, 2014). This genetic cluster is comprised of two previously reported metal ion resistance determinants, neither of which was realized until recently to be part of a single contiguous gene cluster (Crossman *et al.*, 2010, Hobman and Crossman 2015). One, the *pco* determinant was first isolated from plasmid pRJ1004 from an Australian pig *E. coli* isolate (Brown *et al.*, 1995), and confers copper resistance. The other, the *sil* determinant originally located on *Salmonella* Typhimurium plasmid pMG101, but later shown to have transferred into the chromosome of the host *E. coli* K-12 J53 strain (Randall *et al.*, 2015), is associated with silver resistance (McHugh *et al.*, 1975; Gupta *et al.*, 1999). Later sequencing of pRJ1004 (NCBI accession # KC146966) has identified two new genes among the entire 19-gene cluster –

pcoF encoding a putative copper-binding protein, and *pcoG* encoding a putative M23B metallopeptidase, an enzyme that has been implicated in pathogenicity (Bonis *et al.*, 2010) (Figure 2). We have identified this arrangement of *pco/sil* genes in a number of different genome and plasmid sequences.

Similarly, the *sil* determinant has been associated with pathogenicity in the *Enterobacter cloacae* complex, where presence of the genes conferring silver resistance was increased in isolates from hospital settings vs. strains associated with plants (Kremer and Hoffmann 2012). Although identification of the *pco* genes was not part of that study, their presence within the isolates harboring the full *sil* determinant is very likely based on the high rate of co-representation (Mourão *et al.* 2015) (Table 1).

Previous studies and genomic analysis have shown that the copper pathogenicity island is often plasmid associated. Transfer of such plasmids has resulted in a nosocomial outbreak of *Klebsiella pneumonia* (Sandegren *et al.*, 2012). Moreover, the *pco/sil* cluster has been identified on pAPEC-O2-R plasmids from avian pathogenic *E. coli* (Johnson *et al.*, 2005), R478 from *Serratia marcescens* (Gilmour *et al.*, 2004), plasmids pK2044 and pLVPK from *Klebsiella pneumonia* strains (Chen *et al.*, 2004; Wu *et al.*, 2009), and on the chromosome or plasmids of many pathogenic enteric bacteria such as ETEC H10407 (Crossman *et al.*, 2010) and EHEC O104:H4 (Hobman and Crossman 2015) and *Enterobacter cloacae* subsp. *cloacae* strain ATCC 13047 (Ren *et al.*, 2010). For *E. coli* and *Enterobacter* strains the copper pathogenicity island was identified in close vicinity to Tn7-like transposons with *tnsABCD* being present (Peters *et al.*, 2014). At the same time, in *Klebsiella* strains it was often associated with IS4-related elements and genes encoding a HNH endonuclease. These data together with the fact of a similar arrangement of a copper/silver resistance

island has been found in *Salmonella* (Moreno Switt *et al.*, 2012) indicate that the gene cluster behaves like a typical pathogenicity island.

Yersiniabactin- not just for iron

Our recent sequencing of *E. coli* strains isolated from copper-fed pigs allowed us to identify another determinant conferring increased copper resistance – a 10 gene yersiniabactin synthesis cluster (Lüthje *et al.*, 2014). The yersiniabactin determinant is a well-known virulence factor responsible for copper binding (Chaturvedi *et al.*, 2014) that can be present in pathogens such as *Klebsiella pneumonia* (Fodah *et al.*, 2014), *Salmonella* (Aviv *et al.*, 2014), *E. coli* (Schubert *et al.*, 2004), including EHEC O104:H4 outbreak strain, and highly virulent *Yersinia pestis* (Rakin *et al.*, 2012). Interestingly, and several strains of *Klebsiella* and *Escherichia coli* appear to have the *sil/pco* determinant in addition to the yersiniabactin synthesis cluster (Table 1).

Copper and zinc resistance in Gram-positive bacteria

Pathogenicity of Gram-positive bacteria such as *Enterococcus faecium*, *Enterococcus faecalis*, *Staphylococcus aureus* and *Staphylococcus haemolyticus* is also linked to transition metal resistance. Currently, we have sequenced the genomes of three highly copper-resistant *E. faecium* and three *E. faecalis* strains isolated from copper-fed pigs in Denmark (Zhang *et al.*, in press). As a result, we have identified additional copper resistance determinants characteristic for many pathogenic Enterococci, *tcrYAZB*, encoding a negative transcriptional regulator, a copper chaperone and two P_{1B}-Type ATPases flanked by mobile elements (Hasman 2005). In *E. faecalis* this determinant has often been found in the vicinity of a gene encoding a multicopper oxidase resembling CueO, an adjacently encoded two-component system and possibly CopY. Whether CueO is regulated by the adjacent two-component system or CopY is not known. In this genome region there are also several putative copper chaperones and a prolipoprotein diacylglycerol transferase, which has been associated with virulence (Cho *et al.*, 2013; Reffuveille *et al.*, 2012) (Figure 3).

In addition to copper, zinc resistance has also been linked to virulence and increased survival rates of pathogens. For example, in group A *Streptococcus* *czcD* and *gczA* deletion mutants characterized by higher zinc sensitivity had shown much lower survival rates in the presence of neutrophils compared to wild type strains (Ong *et al.*, 2014). Certain correlation between the presence of the zinc resistance gene *czrC*, methicillin resistance and virulence has been found in many *S. aureus* strains (Slifierz *et al.*, 2014; Aarestrup *et al.*, 2010). The gene *czrC* encodes a Zn²⁺-translocating P-type ATPase and is located next to a gene encoding a possible transcriptional regulator of the ArsR/SmtB family and a gene encoding a putative iron/zinc permease. According to genomic similarities the latter might be a distant homolog of the zinc/iron importer ZupT. Sequencing of the *S. haemolyticus* SH32 clinical strain has identified two incomplete Staphylococcal cassette chromosome (SCC) elements, with one of them, Ψ SCC*mec*(SH32), encoding a Cu(I)-translocating P-type ATPase (Yu *et al.*, 2014). This strain was also shown to contain a putative cadmium resistance determinant *cadXD*, encoding for a Cd(II) transporter as well as a transcription regulator of the ArsR family (Yu *et al.*, 2014). A recent study has reported that plasmid SAP078A in methicillin-resistant *Staphylococcus aureus* CC22 SCC*mec*IV (EMRSA-15) contains *cadCA*, *mco* and *copB* in addition to an *ars* operon conferring resistance to cadmium/zinc, copper and arsenic, respectively (Loeffler *et al.*, 2013). It was also shown that plasmid SAP078A is wide spread among both human and animal isolates of *S. aureus* (Loeffler *et al.*, 2013). Moreover, given the role of transition metals in the mammalian immune response, the presence of *cadCA* and *copB/mco* provides corresponding strains with an advantage (Hood and Skaar 2012). Interestingly, the epidemic ST22-IV has been replacing other MRSA clones from hospitals possibly due to enhanced virulence. Detailed studies revealed that ST22-IV has a significantly higher capacity to invade the A549 cells and a higher virulence in a murine model of acute lung infection

causing severe inflammation and determining death in all the mice within 60 hours (Baldan *et al.*, 2012; Baldan *et al.*, 2015). We suggest that severe pathogenicity of ST22-IV might be partially attributed to increased transition metal resistance of this strain.

Cadmium and Silver resistance- mutation plus selection equals evolution to new resistance?

Due to much lower environmental distribution of cadmium and silver they are not essential micronutrients for bacteria and no data has been published on their involvement in bacteria/host interactions. Therefore, it is unlikely for bacteria to develop specific resistance mechanisms to these metals to a large extent. At the same time, most of the metal resistance determinants described to date are involved in detoxification of multiple substrates, e.g. conferring resistance to both copper and silver as well as to both zinc and cadmium (Rensing *et al.*, 1999; Rosenzweig and Arguello 2012). Moreover, both *E. coli* and *Salmonella* contain detoxification systems for zinc and copper in addition to the possible plasmid encoded resistance determinants on their chromosomes. In other words, it is quite unlikely that the evolutionary pressure comes exclusively from metal contaminated environments. Rather, resistance to silver is a by-product of copper resistance and cadmium a by-product of zinc resistance. In fact, resistance studies with *S. aureus* were unable to produce silver-resistant strains even after 42 days of continuous passage in the presence of AgNO₃ (Randall *et al.*, 2013) Similar results were observed for some Gram-negative organisms, whereas in *E. coli* strains silver resistance arises as a result of mutations in both *ompR* and *cusS*, or mediated through *sil* system (Randall *et al.*, 2015).

Metal homeostasis in bacteria is a delicate balance, especially since metals like zinc and iron have been found to be essential for the pathogenicity of these organisms. The fact that protozoan grazing might be a strong force on keeping or gaining resistances against copper or zinc in nature together with the data suggesting that the concentrations necessary to maintain resistance plasmids within a population are well below the minimal inhibitory concentration (MIC) of the non-plasmid containing susceptible stain (Gullberg *et al.*, 2014) might explain the prevalence of resistance mechanisms. Increasing metal contamination

caused by anthropogenic metal use in agriculture and other sectors, might also result in increased ability of protozoans to utilize copper and zinc to kill phagocytized bacteria. This, in turn, can cause higher level of resistance in bacteria – mechanism that has been primarily selected to avoid protozoan killing. In addition, copper is known to induce viable but non-culturable (VBNC) state in some bacteria, causing increase in their survival rates (Ordax *et al.*, 2006). It has been reported that cells of the VBNC state might have been responsible for the recent *E. coli* O104:H4 outbreak (Aurass *et al.*, 2011).

It is worth noting here that copper-induced resistance combined with high toxicity and non-selectivity of redox processes induced by copper presence in cells present major challenges for developing copper-based antimicrobial therapy. Recently, Festa *et al.* have published a novel approach that allows accumulation of copper in pathogen cells without activating its Cu-resistant mechanisms and significantly increases selectivity of the treatment (Festa *et al.*, 2014). In other strategies Cu (II) ions have been either utilized as a carriers for known antibiotics, allowing them to bypass existing efflux-mediated resistance to drugs (Manning *et al.*, 2014; Lopes *et al.*, 2013; Shams *et al.*, 2014), or as chelators that upon binding to ligand change its conformation to the “active” mode (Haeili *et al.*, 2014). Several potent copper chelators with activity against MRSA and *M. tuberculosis* strains have been identified through drug screening assays specifically designed for identification of copper-dependent antimicrobial compounds (Speer *et al.*, 2013) with, potentially, more discoveries on the way.

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Figure Captions

Figure 1. A schematic overview of Zn and Cu involvement in phagosomal killing of bacteria.

Macrophages and amoeba can exploit similar molecules for Zn²⁺ and Cu⁺ trafficking. ZIP family transporters allow Zn²⁺ uptake into the cytoplasm, and cation diffusion facilitator proteins (CDF) could deliver Zn²⁺ to the phagosome and other organelles, like mitochondria, Golgi and Endoplasmatic Reticulum (ER). Cu⁺ uptake and delivery to phagosomes occur due to copper transporter 1 (Ctr1, in amoeba known as P80), antioxidant 1 copper chaperone (Atox1) and in human macrophages the P-type ATPase ATP7A. H⁺-ATPase causes acidification of the phagosomal milieu while natural resistance associated macrophage protein 1 (NRAMP1) removes Fe²⁺ and Mn²⁺, which are needed to protect (Mn²⁺) and rebuild degraded Fe-S clusters of bacteria. In addition, Cu⁺ amplifies toxicity of ROS (hydroxyl radical ($\cdot\text{OH}$) and hydroxid anion (OH^-)). *E. coli* express genes encoding ZntA for Zn²⁺ efflux, CopA for Cu⁺ efflux and the CusCBA complex for periplasmic Cu⁺ efflux but virulent strains have additional copper resistance systems.

Figure 2. Pco and Sil Mechanisms in Action

Proposed genes and protein products forming the molecular mechanisms of Pco and Sil mediated copper and silver detoxification and control in the cell. The bottom line indicates the genes and their transcriptional and translational directions, with the open circles representing potential promoter regions/transcript start sites. The illustrated function of each *sil* and *pco* gene product within the operon (Gram negative) is deduced from homology modelling. The transcription of Pco proteins PcoABCDEFGG appears to be regulated by PcoRS (left). The roles of PcoFG have not been elucidated. In addition to the oxidised catechol siderophores, copper may be detoxified from Cu⁺ to Cu²⁺ by the suggested multi-copper oxidase PcoA. PcoB possibly functions as the outer membrane transporter; whilst sitting in the inner

membrane PcoD drives the transport of Cu^+ from the periplasm to the cytoplasm, with periplasmic PcoC chaperoning/delivering the Cu^+ to PcoD. PcoE is an additional chaperone that binds copper in the periplasm and probably shuttles it to PcoA and/or PcoS. Similarly, the Sil system (right) contains a homolog to PcoE - periplasmic protein SilE. SilE is predicted to bind and chaperone Ag^+ , Cu^+ and Cu^{2+} to the three-polypeptide, transmembrane, chemiosmotic RND exchange system (SilCBA), exporting the metal ions out of the cell. Likewise, SilF acts as a chaperone to SilCBA too. The other putative efflux pump mediating the mechanism is a P-type ATPase – SilP. Although conserved within the *sil*-determinant, a role of SilG has not yet been determined. While the expression of *silCFBAGP* is thought to be governed by the two-component membrane sensor and transcriptional responder SilRS; the expression of *silE*, just like homolog *pcoE*, is thought to be regulated/co-regulated by the Cus system (Zimmerman et al., 2012) and therefore SilE perhaps is involved in the activation of regulators SilRS (proposed by dotted arrow).

Figure 3. Copper fitness island in *Enterococcus faecalis*

Proposed genes and protein products forming the molecular mechanisms of copper detoxification in *E. faecalis*. The bottom line indicates the genes and their transcriptional and translational directions. TcrY regulates expression of *tcrYAZB*, encoding for the repressor, a cytoplasmic chaperone (TcrZ) and two P-type ATPases (TcrA and TcrB) responsible for Cu^+ export. In close proximity to *tcr* genes encoding a two-component regulatory system (CueRS), a multicopper oxidase (CueO) a predicted metal chaperone (no homology to TcrZ and labeled “C”) and transcriptional repressor (CopY) have been identified. CueO is predicted to oxidize Cu^+ to Cu^{2+} . It is not clear to what extend the predicted chaperon (C) might be involved in copper detoxification. It has not yet been established if transcription of these genes is controlled by the two-component regulatory system (CusRS) responding to external copper concentration, CopY, as a response to internal copper concentrations, or both. Adjacent to and separating the two copper resistance determinants genes encoding prolipoprotein diacylglyceryl transferase (A), integral membrane protein (B),

hypothetical proteins (D), transposase (E), disrupted P-type ATPase (F), integrase (G), adenylate kinase (I) and resolvase (I) have been identified. The extent to that some of these proteins might be involved in copper detoxification has not been analyzed.

Tables

Table 1 Distribution of copper/silver resistance cluster and yersiniabactin biosynthesis amongst Enterobacteriaceae

Genus ^{a)}	Number of sequences analyzed ^{b)}	occurrence of copper/silver tolerance determinants				
		<i>pco</i> ^{c)}	<i>sil</i> ^{d)}	<i>pco/sil</i> ^{e)}	yersiniabactin synthesis ^{f)}	<i>pco/silP</i> and yersiniabactin synthesis ^{g)}
<i>Citrobacter</i>	4 genomes	2	2	2	1	0
	5 plasmids	0	0	0	0	0
<i>Cronobacter</i>	6 genomes	1	1	1	0	0
	5 plasmids	3	3	3	0	0
<i>Enterobacter</i>	16 genomes	5	6	5	1	0
	12 plasmids	3	3	3	0	0
<i>Escherichia</i>	74 genomes	10	10	10	42	6
	71 plasmids	2	4	2	0	0
<i>Klebsiella</i>	32 genomes	0	0	0	17	6
	33 plasmids	29	29	29	0	0
<i>Raoultella</i>	2 genomes	0	0	0	1	0
	1 plasmid	0	0	0	0	0
<i>Salmonella</i>	252 genomes	5	5	5	0	0
	77 plasmids	1	1	1	1	0
<i>Serratia</i>	18 genomes	0	0	0	0	0
	8 plasmids	1	1	1	0	0
<i>Yersinia</i>	33 genomes	0	0	0	27	0
	33 plasmids	0	0	0	0	0

^{a)} genera of Enterobacteriaceae harboring *pco*, *sil* and/or *ybt*

^{b)} number of completed genomic and plasmid sequences of respective genera available for Microbial Genome BLAST® (<http://blast.ncbi.nlm.nih.gov>; accessed 05/18/2015)

- ^{c)} analysis (blastn) using *pco* from pRJ1004 (accession # X83541.1; Brown *et al.*, 1995) as query
- ^{d)} analysis (blastn) using *sil* from pMG101 (accession # NG_035131.1; Gupta *et al.*, 1999) as query
- ^{e)} analysis (blastn) using *pco* (accession # X83541.1; Brown *et al.*, 1995) and *sil* (accession # KC146966.1; Staehlin *et al.*, 2013 – direct submission) from pRJ1004 as query
- ^{f)} analysis (tblastn) using Ybt peptide/polyketide synthetase HMWP1 (accession # AAC69588.1; Gehring *et al.*, 1999) as query
- ^{g)} number of strains harboring *pco/sil* and *ybt* with determinants being located on chromosome and/or plasmid, respectively

Figures

Figure 1

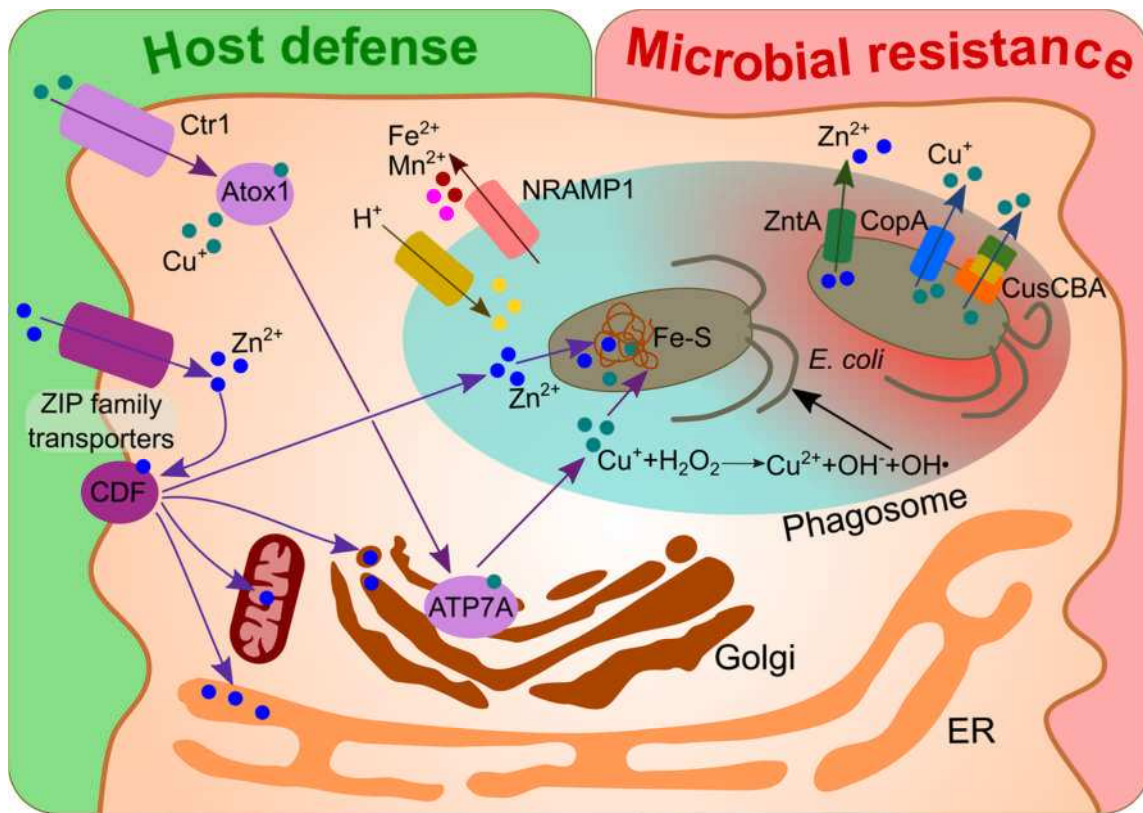


Figure 2

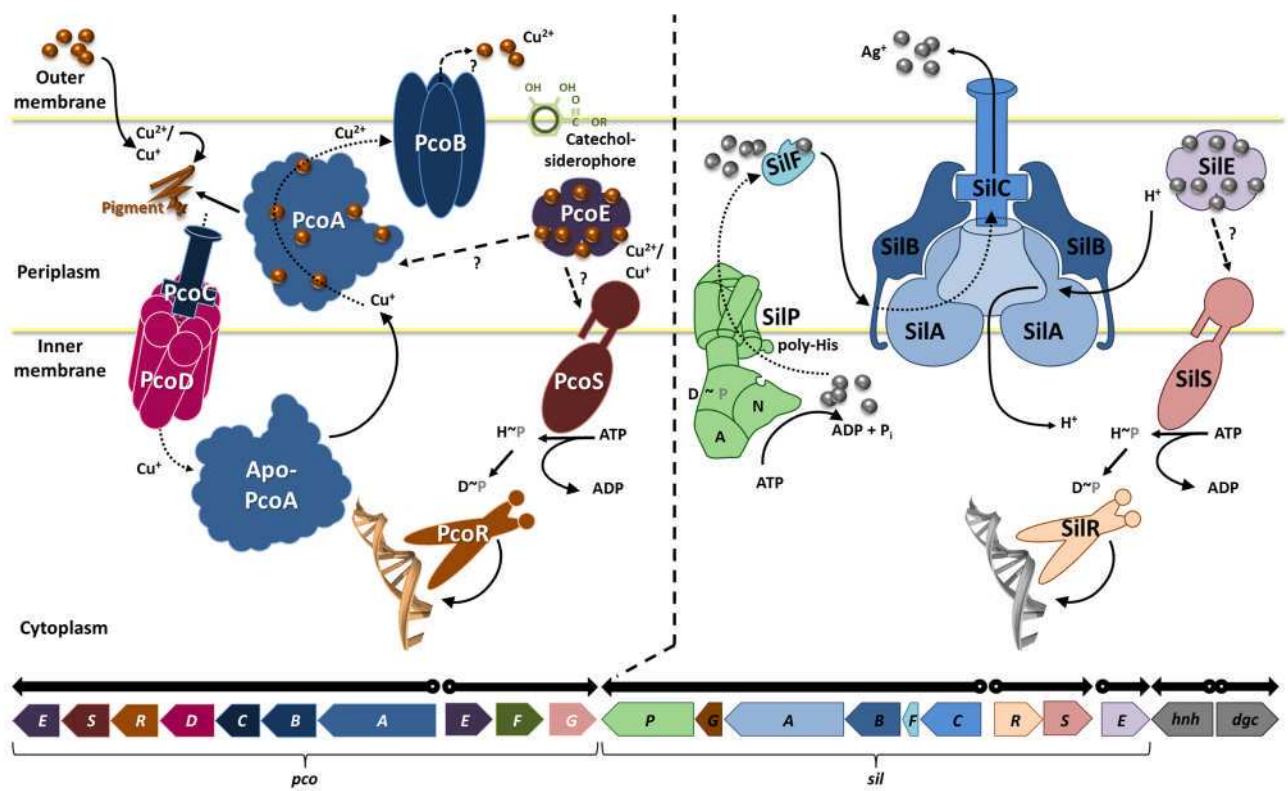


Figure 3

