Validation of Anti-Oxidative Stress Genes from Genome-Wide Screening of Escherichia coli

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Validation of Anti-Oxidative Stress Genes from Genome-Wide

Screening of *Escherichia coli*

An Honors Thesis submitted in partial fulfillment of the requirements of Honors Studies in Chemistry with Biochemistry Concentration

By

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Abstract

The primary purpose of this project is to evaluate the genes that play a role in the oxidative stress response in *Escherichia coli*. In doing so, the entire genome of *E. coli* was subject to throughput in which individual genes were determined to have a role in the bacteria’s oxidative stress response. Moreover, this project focused on the validation of the genes that were able to pass the initial throughput stage. The genes were subject to two forms of validation. In the first validation technique, candidate genes were overexpressed and minimum inhibitory concentrations of hypochlorous acid were taken. Following, a second validation technique tested the minimum inhibitory concentrations of hypochlorous acid utilizing knock-out mutants. These experimental results will enable for better understanding of a common pathogen as well as ways to defend against it. Additionally, *E. coli* has often been dubbed as a model bacterium and is utilized and studied in many scientific investigations. Furthermore, this experiment may lead to a greater general understanding of the bacterium, which could assist in future studies.
Introduction

When an organism undergoes oxidative stress, the organism’s cells experience an excess concentration of reactive oxygen intermediates, such as superoxide (O$_2^•$), hydroxyl radicals (HO$^•$), or hydrogen peroxide (H$_2$O$_2$). These intermediates are capable of being produced naturally, primarily as a byproduct of electron transfer in the respiratory chain in the mitochondria. Although these reactive oxygen species are produced endogenously within the cell, the reactive species can become detrimental to cells at both unusually high and unusually low concentrations. At high concentrations, reactive oxygen intermediates are capable of oxidizing a multitude of macromolecules, including proteins and DNA. This ultimately may lead to the cessation of normal cellular function and subsequent cell death. Additionally, within humans, an excess of intermediates correlates with higher rates of cancer, cardiovascular, and neural diseases.

While high levels of reactive oxygen species are harmful to various organisms, low levels of endogenous reactive oxygen species can also prove disadvantageous. Abnormally low level of these intermediates can coincide with a lower levels of cell growth, development, and differentiation. Thus, maintaining favorable levels of reactive oxygen species is vital for cells to maintain normal functions and efficiency.

To combat potential pathogens or unwanted bacteria, many pharmaceutical drugs and household cleaners attempt to target undesired organisms by introducing compounds, such as hydrogen peroxide and hypochlorous acid (HOCl), which attempt to generate high levels of reactive oxygen species within the pathogen or bacteria. In addition, bacteria are also subject to intermediates in other hostile environments. Plants and
animals regularly produce reactive oxygen intermediates in efforts to protect against pathogens as well\(^5\).

Previously, it was thought that oxidative stress induced by reactive oxygen species had been well studied in *Escherichia coli*, the model bacterium. However, in more recent studies, it was found that these bacteria have other tools at its disposal to combat high concentrations of reactive oxygen species. More specifically, it was discovered that noncoding RNAs and small proteins, less than 50 amino acids, also played a role in the bacteria’s oxidative stress response and have yet to be studied\(^6\). Prior studies, instead, tended to focus on *E. coli*’s use of enzymes to scavenge the reactive intermediates, including catalase, peroxidase, mutase\(^7\). The studies placed emphasis on regulatory genes, OxyR, SoxR and SoxS in particular, that are involved in the activation of detoxifying enzymes when a bacterium is undergoing oxidative stress as well. Some of these inducible detoxifying enzymes include alkyl hydroperoxide reductase and superoxide dismutase\(^8\).

This evidence points to the fact that the knowledge of the oxidative stress response utilized by *E. coli* is still relatively limited. Therefore, this project’s goal is to screen the entire genome of *E. coli* without reliance on previously established knowledge. Moreover, this project would be able to provide a complete set of genes that allow *E. coli* to survive toxic levels of reactive oxidative intermediates. By identifying all of the genes *E. coli* uses to resist oxidative stress, new ways to combat the bacteria may emerge and a greater understanding of the bacteria itself may be grasped. Since hypochlorous acid, more commonly known as bleach, is routinely used as an antiseptic, this project aimed to identify the genes in *E. coli* that allow for resistance to HOCl.
Prior to my experimentation, the growth effects of varying concentrations of HOCl on the AG1 cells containing the plasmid were measured by others in the Fan Lab. To limit the effects of deviation within the experiment, four duplicates of each plate were made. Candidates were only able to pass the initial testing if three of the four copies were able to effectively grow in 6 mM HOCl, which is three times the minimal inhibitory concentration of *E. coli*.


To ensure these 23 genes actually did play a role in the bacteria’s stress response, two validation tests were conducted. The first form of validation was conducted by use of overexpression experiments. In this, *E. coli* cells were made to overexpress 1 of the 23 candidate genes listed previously. Thus, if the genes were truly able to assist in the bacteria’s oxidative stress response, the minimum inhibitory concentration of HOCl would be predicted to increase. Conversely, the second form of validation was done so through knock-out experimentations. In doing so, minimum inhibitory concentrations of *E. coli* cells missing 1 of the 23 candidate genes were taken. As opposed to the overexpression test, the minimum inhibitory concentrations would be predicted to decrease if the candidate gene truly did allow for survival at toxic levels of HOCl.
Experimental Methods

1. Validation of Identified Genes Through Overexpression

Due to the selection pressure of hypochlorous acid, mutated genes may have occurred in *E. coli* cells that enabled cell growth despite the unfavorable conditions brought on by HOCl. Another potential problem may have resulted due to the fact that the AG1 strain of *E. coli* is not a wild-type strain. To remedy this, gene candidates that were able to withstand the initial round of screening were overexpressed in wild-type MG1655 *E. coli* K-12 strain. Additionally, the MG1655 *E. coli* K-12 is known to be highly accurate strain and is widely utilized in other *E. coli* physiology experiments.\(^\text{12,13}\)

Genes were overexpressed by employing a pCA24N plasmid vector. Minimal inhibitory concentrations of HOCl for the various overexpressed MG1655 *E. coli* strains were determined and can be found in Figure 1. Figure 1 also displays the minimal inhibitory concentrations for a control in which a MG1655 *E. coli* strain was void of any overexpressed genes. The displayed genes mirror the genes that passed the first round of screenings.
Figure 1. Minimum inhibitory concentrations (MIC) of HOCl in overexpressed gene MG1655 mutants.

2. Validation of Identified Genes Through Knock-Out Mutations

As priorly stated, the *E. coli* cells may be subject to random mutation or selective pressure due to HOCl. Thus, the genes that were able to pass the screening were subject to an additional form of validation. A second test was used to further verify the results of the previous overexpression test and initial throughput screening. Instead of the ASKA Library, the Keio Collection was used in this section of experimentation. The Keio Collection contains a set of single-gene, knockout mutants for all nonessential genes in *E. coli*\textsuperscript{14,15}. Thus, the Keio Collection allows for robust analysis of *E. coli* cells with loss of specific gene function.

Much like the previous validation test, minimal inhibitory concentrations of HOCl were determined. These results are presented in Figure 2. Figure 2 also exhibits the minimal inhibitory concentration of the BW25113 strain, the wild-type control of the Keio Collection. The results of Figure 2 confirm a higher sensitivity to HOCl when
compared to the wild-type BW25113 strain. Furthermore, the removal of the gene candidates from the overexpression portion of testing resulted in a high sensitivity to HOCl as well. This data substantiates the idea that the genes identified do play a role in the bacteria’s defense against oxidative stress.

Figure 2. Minimum inhibitory concentrations (MIC) of HOCl in gene knockout BW25113 mutants.
Discussion

From Table 1, which displays the tested genes’ functions and subcellular location, one can see that most of the genes that played a role in \textit{E. coli}’s oxidative stress defense are predominately located in the cytosol or within the cellular membrane. More specifically, 12 of the 23 gene products that passed through screening and validation are found in the cytosol. Contrastingly, the remaining 11 gene products are observed in the bacteria’s cell membrane. However, many of these genes have not been the subject of any studies in regard to oxidative stress defense. The localization and function of candidate genes were determined using the UniProt-GOA Database, STRING Database, and EcoCyc Database.

Table 1. Brief description of identified genes for HOCl resistance.

<table>
<thead>
<tr>
<th>Genes</th>
<th>Encoded Protein Function</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{prpE}</td>
<td>Propionate-CoA ligase; Catalyzes the synthesis of propionyl-CoA from propionate and CoA.</td>
<td>cytosol</td>
</tr>
<tr>
<td>\textit{frmB}</td>
<td>S-formylglutathione hydrolase; Hydrolyzes S-formylglutathione to glutathione and formate.</td>
<td>cytosol</td>
</tr>
<tr>
<td>\textit{rutC}</td>
<td>Aminoacrylate peracid reductase RutC.</td>
<td>cytosol</td>
</tr>
<tr>
<td>\textit{sucC}</td>
<td>Succinate-CoA synthetase; Succinyl-CoA synthetase functions in the citric acid cycle.</td>
<td>cytosol</td>
</tr>
<tr>
<td>\textit{ycbB}</td>
<td>L, D-transpeptidase; Involved in peptidoglycan biosynthesis and exhibits transferase activity.</td>
<td>plasma membrane</td>
</tr>
<tr>
<td>\textit{yccJ}</td>
<td>PF13993 family protein YccJ.</td>
<td>cytosol</td>
</tr>
</tbody>
</table>
hyaC  Probable nickel-dependent hydrogenase b-type cytochrome subunit.

hyaE  Hydrogenase-1 operon protein HyaE; appears to be a non-essential chaperone that is specific for the beta subunit of hydrogenase isoenzyme 1.

taP   Chemotaxis methyl-accepting receptor.

yoaE  Integral membrane protein TerC; Involved in flavin adenine dinucleotide (FAD) binding.

motB  Motility protein B; MotA and MotB comprise the stator element of the flagellar motor complex which is required for rotation of the flagella.

rnC   Ribonuclease 3; Involved in rRNA processing and digestion of double-stranded RNA formed within single-stranded substrates.

elaA  N-acetyltransferase ElaA; Involved in transference of acyl groups other than amino-acyl groups.

yfdY  Protein YfdY; Involved in colanic acid synthesis and biofilm development.

exbD  Biopolymer transport protein ExbD; ExbD is used in the TonB energy system\(^{26}\).

ilvA  Threonine deaminase; Catalyzes the anaerobic formation of alpha-ketobutyrate and ammonia from threonine in a two-step reaction.
DNA-methylase; Methylates the second adenine of the Nsil recognition sequence.

Uncharacterized membrane protein YtfI.

Encodes probable multidrug ABC transporter permease.

Catalyzes the alpha, beta-elimination reaction of D-cysteine and of several D-cysteine derivatives; May be used as a defense mechanism against D-cysteine\textsuperscript{27}.

MarC is an inner membrane protein with six predicted transmembrane domains.

Multi-copy expression of sanA can complement permeability defects to vancomycin-susceptible mutants\textsuperscript{28}.

Pseudouridine synthase A that is responsible for pseudouridylation of 16S rRNA at position 516\textsuperscript{29}.

Among the genes brought forth by this study, ycbB, yedQ, sanA, marC, ybhR, exbD, yfdY, motB, yoaE, hyaC, and taP, encode proteins that are vital for the maintenance of the membrane system and its functions. ycbB, a gene that encodes for L, D-transpeptidase and aids in peptidoglycan synthesis, was discovered to be allow for resistance to beta-lactam antibiotics, which targets the outer membrane of the bacteria\textsuperscript{16}. The gene, yfdY, has been shown to be involved in the synthesis of colonic acid, an extracellular polysaccharide that aids in \textit{E. coli}'s ability to form biofilms\textsuperscript{17}. These biofilms can provide the bacteria added protection against antibiotics as well as give the bacteria significant adhesive properties to hard surfaces and other bacteria that allow
them to potentially evade host defense mechanisms\textsuperscript{18}. Additionally, \textit{taP} and \textit{motB} were found to encode proteins that assist in \textit{E. coli}'s ability to move, with \textit{taP} mediating chemotaxis and \textit{motB} partially encoding a protein that is necessary for flagellar motion\textsuperscript{18}. Genes, \textit{hyaC} and \textit{yoeA}, play roles in bioenergetics within the cell. Additionally, the genes assist in the complex transport system that exists within the cell, specifically \textit{exbD} and \textit{ybhR}. The remaining of the suggested genes located within the plasma membrane, \textit{yfl}, \textit{yedQ}, \textit{marC}, and \textit{sanA}, participate in various barrier functions of the plasma membrane.

Furthermore, Figure 1 demonstrates that the greatest increase of minimum inhibitory concentration coincided with overexpression of the gene \textit{yoeA}. Although not studied within \textit{E. coli}, the gene \textit{yoeA} has been more closely analyzed inside a different bacterium, \textit{Salmonella enterica}. \textit{S. enterica} and \textit{E. coli} are closely related as they are both facultative anaerobes, Gram-negative, and rod-shaped\textsuperscript{19}. Moreover, it was observed that \textit{yoeA}, a gene that codes for a membrane protein, was upregulated by CpxR when \textit{S. enterica} were put into chicken egg whites\textsuperscript{20}. An egg white presents a very complex and unique environment that aides in a chicken’s defense against pathogens, including its high alkalinity, the large number of lysozymes present, and its high viscosity\textsuperscript{21,22}. Huang et. al concluded that, although it is unknown which stress factor the upregulation is in response to, \textit{yoeA} was necessary for the bacteria’s survival within the egg white environment. Therefore, due to the two bacteria’s structural and genetic comparability, \textit{E. coli} may function similarly and require upregulation in similar hostile environments.

Moving from genes localized in the membrane, the remaining suggested genes, \textit{prpE}, \textit{frmB}, \textit{rutC}, \textit{sucC}, \textit{yccJ}, \textit{hyaE}, \textit{rnC}, \textit{elaA}, \textit{ilvA}, \textit{yhdJ}, and \textit{rsuA}, are localized within the cytosol. These genes encode proteins with a wide variety of functions. Among these
many functions included proteins involved in rRNA production ($rnC$) as well as DNA and RNA modification ($yhdJ$). Furthermore, $ilvA$ contributes to isoleucine synthesis and $rsuA$ is responsible for pseudouridine synthesis. Pseudouridine, one of the most common RNA modifications, has been exhibited to have important roles in various aspects of gene regulation, such spliceosomal small nuclear ribonucleoprotein biogenesis and translation fidelity\textsuperscript{23}. Finally, genes $prpE$ and $sucC$ play roles in cellular metabolism, specifically propionate synthesis and succinyl-CoA synthesis, respectively.

From the study, we have found that there is still much to learn about $E. \textit{coli}$’s ability to combat oxidative stress generated by HOCl, a common household disinfectant. While many of the genes have been previously studied, they were not done so in regard to their ability to allow for the bacteria to survive toxic levels of reactive oxygen species. Thus, future studies may focus the ways two or more genes interact with one another when the genes are overexpressed or deleted. Additionally, further experimentation may be necessary to determine the bacteria’s response to other defense mechanisms that seek to generate excess reactive oxygen species, such as superoxide which is commonly produced by mammalian phagocytes in efforts to destroy potential pathogenic cells.
References


