ADAPTIVE MECHANISMS UNDERLYING MICROBIAL RESISTANCE TO DISINFECTANTS

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Adaptive Mechanisms Underlying Microbial Resistance to Disinfectants

This project was performed to determine if clinical pathogens evolve and acquire resistance to Lysol, an all-purpose cleaner and disinfectant (U.S. Environmental Protection Agency regulation no. 777-89 [ready-to-use (RTU) 1:16 dilution]). A clinical surrogate, *Escherichia coli*, was used in these studies. *E. coli* cells were grown in the absence or presence of Lysol. The parent strain (PS) was sensitive to the presence of 1.6% RTU strength Lysol. LR50, a resistant strain that showed resistance to 50% of the RTU strength Lysol, was derived through progressive subculturing. (A 30-fold increase in resistance to Lysol illustrates genome plasticity and adaptation of bacterial cells.) LR50 was subcultured in Tryptic soy broth five times, and then its resistance phenotype was confirmed in the presence of 50% Lysol. Biochemical characterization revealed the presence or absence of specific polypeptides unique to LR50. Genomic sequencing was done, and some single nucleotide polymorphisms were observed to be unique to LR50. Finally, altered antibiotic resistance was determined for LR50. In a separate set of experiments, the adaptive resistance of *E. coli* cells to 17% Germ-X (GR17), a hand sanitizer, was also observed. GR17 PS, a resistant strain, which was capable of growing in the presence of GR17, was derived during this study.
PREFACE

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1. INTRODUCTION

The emergence of a new resistance mechanism in the “superbug” and its spread across bacterial species illustrate microbial adaptability to sublethal exposures of antimicrobials. The increasing use of disinfectants in clinical and household settings has raised serious concerns for the development of resistance mechanisms in bacterial cells (McDonnell et al., 1999). The mechanisms underlying each of these two phenomena may be quite different. A clearer understanding of biochemical and genetic adaptation is expected to offer new insight into countermeasure development.

Penicillin was first used to treat bacterial infections in the 1940s. By the 1950s, a penicillin-resistant strain of Staphylococcus aureus became common. Methicillin was introduced in 1961 to treat infections with such cells. Within one year, Methicillin-resistant S. aureus (MRSA) strains were observed. MRSA strains are resistant to a host of antibiotics, including vancomycin. Bacterial cells evolved and developed resistance to antimicrobial use, and the short generation times and large population size of bacteria helped boost this evolution.

Disinfectants are commonly used in water treatment plants and for cleaning surfaces in medical treatment facilities. In addition, household use of disinfectants has increasingly grown as these products typically kill 99.999% of pathogens within 5–10 min. Chlorine and quaternary ammonium compounds (quats) are the common key ingredients in these disinfectants. The general functions of disinfectants include destruction of cell membranes, interference with key biochemical functions, blockage of nutrient uptake, and decontamination of waste products. The increasing use of disinfectants raises the prospect of improper use (i.e., sublethal concentrations of disinfectants result in the adaptation of bacterial populations, leading to derivation of resistant isolates).

Mechanisms for disinfectant resistance of bacterial cells include the secretions of gelatinous exopolysaccharide, which some species can use to form biofilms. The use of an efflux pump to selectively export the disinfectant is another possible mechanism for resistance. Some bacteria may alter their gene expression of novel transporter proteins for protection.

This study compared disinfectant-resistant populations of Escheria coli with control sets of E.coli for physiological, biochemical, and genetic differences in an attempt to understand resistance mechanisms. Bacterial cells with increased resistance to disinfectants were analyzed for their resistance to antibiotics to investigate a possible correlation between these characteristics.
2. MATERIALS AND METHODS

2.1 Materials

2.1.1 Bacterial Strains Used

We used *E. coli*, a clinical surrogate, for the disinfectant-resistant tests. The *E. coli* was obtained from the American Type Culture Collection ([ATCC] Manassas, VA [ATCC number 11775]). *E. coli* is a gram-negative, facultative anaerobic, and rod-shaped bacteria commonly found in warm-blooded animals.

2.1.2 Disinfectants Used

We used Lysol brand disinfectant, four-in-one all-purpose cleaner (U.S. Environmental Protection Agency registration number 777-89). This disinfectant consists of alcohols and quaternary ammonium chlorides. Lysol contains the following ingredients: alcohols C12–16, ethoxylated (2.5–10%), Alkyl (50%C14, 40%C12, 10%C16) dimethyl benzyl ammonium chlorides (1–2.5%), and ethanol (0.1–1%). Alkyl dimethyl benzyl ammonium chloride attaches to the bacteria and causes the cytoplasmic membrane to leak, which damages and then kills the bacterial cells. The manufacturer recommended a ready-to-use (RTU) concentration of 1:16 of Lysol/Tryptic soy broth (TSB), which was regarded as 100% effective.

2.1.3 Reagents Used

TSB was used as the diluent in all the experiments. The RTU Lysol was further diluted in the ratio of 1:10 in a working stock.

Tryptic soy agar was used for plating the *E. coli* cells in all of the experiments.

2.2 Methods

2.2.1 Growth of *E. coli*

*E. coli* was grown in TSB overnight, and a micro-titer plate assay was used to screen the parent strain (PS) of *E. coli* for sensitivity. Sensitivity percent was then determined.

*E. coli* was grown over a 6 month period in increasing concentrations of Lysol. Growth curves were conducted for the PS *E. coli* cells, the Lysol-resistant (LR) strain in TSB, and another strain that showed resistance to 50% of the RTU strength Lysol (LR50). Growth curves were completed to determine the doubling time of the bacteria.
2.2.2 Biochemical Tests

Protein extraction was performed from the bacterial cells. SDS-PAGE* gels were run to compare the proteins in the PS and LR strains. The two strains were also examined using microscopy to compare the cell types.

2.2.3 Confirmation of the Disinfectant-Resistant Phenotype

The genetic basis of the disinfectant phenotype of the LR strain was confirmed. The LR50 cultures were grown in TSB and subcultured five times. They were then grown in the presence of 50% Lysol.

2.2.4 Antibiotic Testing

Nineteen antibiotics were tested against PS and LR50 strains, respectively. The following antimicrobial susceptibility test discs were tested: 15 µg of azithromycin, 15 µg of clarithromycin, 5 µg of rifampin, 5 µg of tetracycline, 2 µg of ampicillin, 10 µg of amoxicillin, 2 IU of penicillin, 10 IU of penicillin, 30 µg of tetracycline, 5 µg of cefdinir, 30 µg of cefaclor, 5 µg of novobiocin, 10 µg of streptomycin, 10 µg of gentamicin, 20 µg of amoxicillin, 5 µg of ciproflaxin, 10 µg of bacitracin, 5 µg of vancomycin, and 100 µg of carbenicillin. The appearance of a zone of inhibition was proportional to bacterial resistance. Therefore, the zone of inhibition was measured in side-by-side tests of LR50 and PS samples. The zone of inhibition on the growth of the LR50 strain was caused by each antibiotic and compared with that of the growth on the PS test discs.

2.2.5 Genomic Sequencing

The samples were prepped for genome sequencing using a Nextera (Illumina, Inc., San Diego, CA) sequencing kit and sequenced on an Illumina HiSeq 2000 sequencing system. A 2 × 100 run configuration was used (i.e., reads were produced in pairs, and each pair was 100 bases long). Each sample produced >5 gigabases. Data were analyzed using Bowtie sequencing analysis and SAMtools and CLC Bio genomics workbenches (CLC Bio, Qiagen, Aarhus, Denmark). DNA was isolated from the PS and LR50 strains. The genome sequences of both strains were analyzed and compared.

3. RESULTS

3.1 Growth Curves

Figures 1–3 show the growth curves and doubling times of the PS and LR50 strains in different cultures.

*SDS-PAGE is sodium dodecyl (lauryl) sulfate-polyacrylamide gel electrophoresis.
Figure 1. Growth curve of LR50 strain in TSB. The doubling time is 60 min.

Figure 2. Growth curve of LR50 strain in 50% Lysol. The doubling time is 65 min.
Figure 3. Growth curve of PS in TSB. The doubling time is 60 min.

Figure 4 shows that the LR strain grown in the 50% Lysol is missing a polypeptide of ~130 kDa, which is present in the LR and control strains grown in TSB. A polypeptide of ~100 kDa is uniquely present in the LR50 strain after it was grown in Lysol. Alterations in gene expression appeared to be the basis for the LR50 phenotype.

3.2 **Biochemical Tests**

![Parent Strain in TSB](image)

![Protein gel with both LR strains in 50% Lysol, LR strain in TSB, and control strain in TSB](image)

Figure 4. Protein gel with both LR strains in 50% Lysol, LR strain in TSB, and control strain in TSB.
3.3 **Antibiotic Resistance**

LR50 was resistant to two antibiotics, whereas the PS was not. LR50 had no zone of inhibition around 5 µg of rifampin and 10 IU of penicillin. On the other hand, the PS showed a zone of inhibition around these two antibiotics. Therefore, only LR50 was observed to be resistant to these two antibiotics.

Figures 5 and 6 show the antibiotic-resistant profiles of LR50 and PS.

![Figure 5. Antibiotic-resistant profiles of LR50 and PS for 9 antibiotics.](image1)

![Figure 6. Antibiotic-resistant profiles of LR50 and PS for 10 antibiotics.](image2)
3.4 Genomic Sequencing

DNA was isolated from the PS and LR50 strains, and the genome sequences of both strains were compared. Certain single nucleotide polymorphisms (SNPs) are unique to LR50, including **acrB** and **mdtB** (multidrug efflux system proteins), subunit B (**yfjW**, inner membrane protein), and **yghB** (inner membrane protein). The fact that the multidrug efflux system protein was altered in the LR50 strain, but not in the PS, could be the likely cause of resistance because multidrug efflux pumps are transport proteins that eliminate toxins from within bacterial cells.

4. DISCUSSION

The growth and subculture of LR *E.coli* cells were continued for a 6 month duration in increasing concentrations of Lysol until the cells were able to grow in the presence of 50% of the RTU Lysol. This is consistent with findings in other studies. In one study (Hoff et al., 1986), *E. coli* was shown to have enhanced resistance to chlorine after repeated exposures. The phenotype of LR50 (i.e., LR *E.coli* cells) was confirmed by subculturing five times in TSB (in the absence of Lysol) and then growing cells in the presence of 50% RTU Lysol. Frozen glycerol stocks were prepared and the LR50 phenotype was confirmed by once again culturing cells in the presence of 50% Lysol. In another study (Moen, B. et al., 2012), *E.coli* cells were grown in the presence of quats and each generation of cells showed resistance to it.

Growth curves were performed with PS *E. coli* cells grown in TSB and 50% Lysol, LR50 cells grown in TSB and in 50% Lysol (Figures 1–3). The doubling times for PS and LR50 strains in TSB were about 60 min. The doubling time for LR50 in 50% Lysol was 65 min. As expected, PS *E. coli* cells did not grow in 50% Lysol.

The preliminary results showed differences in the protein levels of PS and LR50 strains, which suggested alterations in gene expression as a basis for LR50 growth in 50% Lysol (Figure 4). The LR strain grown in 50% Lysol was missing a polypeptide of ~130 kDa, which was present in the PS and LR strains grown in TSB. The LR50 strain grown in 50% Lysol had a polypeptide of ~100 kDa that was missing in both the PS and LR strains grown in TSB. This polypeptide could be the reason for the disinfectant resistance, but more studies need to be conducted to verify the identity of the polypeptide.

Nineteen antibiotics were tested against the PS and LR50 strains. The LR50 strain had no zone of inhibition around the 5 µg of rifampicin and the 10 IU of penicillin, whereas the PS had a zone of inhibition around these two antibiotics (Figures 5 and 6). Only the LR50 strain was observed to be resistant to 5 µg of rifampicin and 10 IU of penicillin. These data disagree with one study, which proposed that no correlation existed between antibiotic and disinfectant resistance (Beier et al., 2013).

DNA was isolated from the PS and LR50 strains, and the genome sequences of both strains were compared. The following SNPs were unique to LR50: **acrB** (multidrug efflux...
system protein), *mdtB* (multidrug efflux system), and subunit B, *yfjW*, and *yghB* (inner membrane proteins).

The LR50 strain likely showed resistance because the multidrug efflux system protein altered in the LR50 strain and not in the PS. Multidrug efflux pumps are transport proteins that eliminate toxins from within bacterial cells. In one study, Sulavik et al. (2001) found that the multidrug resistance pumps contributed to the resistance of *E. coli* cells to antibiotics and different chemicals. Our hypothesis is that alteration in the multidrug efflux pump enabled the LR50 strain to grow in the presence of Lysol.

5. **CONCLUSIONS**

In this study, bacterial cells were shown to acquire resistance to increasing concentrations of 50% RTU Lysol. This adaptive phenotype appeared to change in gene expression (i.e., specific proteins were gained and lost). The genome sequencing of the PS and resistant strains showed SNPs in the genome of the Lysol-resistant strain, corresponding to the multidrug efflux pumps. The efflux pumps may have cycled the disinfectants causing this resistance. Finally, this study addressed the very fundamental question of whether disinfectant strains could show altered antibiotics resistance. The results summarized in this report strongly support the idea that cells resistant to disinfectants also display altered resistance to antibiotics.

More studies need to be performed, especially studies addressing the adaptive potential of pathogenic strains, such as *Acinetobacter baumannii*, *Pseudomonas diminuta*, and *S. aureus*, against sublethal doses of disinfectants like Germ-X (a hand sanitizer) and quats. Further work on SNPs is expected to shed light on the specific gene or set of genes that can be considered for phenotype study.


### ACRONYMS AND ABBREVIATIONS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
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<tbody>
<tr>
<td>ATCC</td>
<td>American Type Culture Collection</td>
</tr>
<tr>
<td>LR</td>
<td>Lysol resistant</td>
</tr>
<tr>
<td>LR50</td>
<td>Lysol resistant in 50% Lysol</td>
</tr>
<tr>
<td>MRSA</td>
<td>Methicillin resistant <em>Staphylococcus aureus</em></td>
</tr>
<tr>
<td>PS</td>
<td>parent strain</td>
</tr>
<tr>
<td>quats</td>
<td>quaternary ammonium compounds</td>
</tr>
<tr>
<td>RTU</td>
<td>ready-to-use</td>
</tr>
<tr>
<td>SDS-PAGE</td>
<td>sodium dodecyl (lauryl) sulfate-polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>SNP</td>
<td>single nucleotide polymorphism</td>
</tr>
<tr>
<td>TSB</td>
<td>Tryptic soy broth</td>
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