**O-specific polysaccharide confers lysozyme resistance to extraintestinal pathogenic *Escherichia coli***

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**ABSTRACT**

Extraintestinal pathogenic *Escherichia coli* (ExPEC) is the leading cause of bloodstream and other extraintestinal infections in human and animals. The greatest challenge encountered by ExPEC during an infection is posed by the host defense mechanisms, including lysozyme. ExPEC have developed diverse strategies to overcome this challenge. The aim of this study was to characterize the molecular mechanism of ExPEC resistance to lysozyme. For this, 15,000 transposon mutants of a lysozyme-resistant ExPEC strain NMEC38 were screened; 20 genes were identified as involved in ExPEC resistance to lysozyme—of which five were located in the gene cluster between *galF* and *gnd*, and were further confirmed to be involved in O-specific polysaccharide biosynthesis. The O-specific polysaccharide was able to inhibit the hydrolytic activity of lysozyme; it was also required by the complete lipopolysaccharide (LPS)-mediated protection of ExPEC against the bactericidal activity of lysozyme. The O-specific polysaccharide was further shown to be able to directly interact with lysozyme. Furthermore, LPS from ExPEC strains of different O serotypes was also able to inhibit the hydrolytic activity of lysozyme. Because of their cell surface localization and wide distribution in Gram-negative bacteria, O-specific polysaccharides appear to play a long-overlooked role in protecting bacteria against exogenous lysozyme.

**Introduction**

Lysozyme is a key player in the innate immune system, secreted by various tissues [1] and cells—including neutrophils and macrophages [2]—and found at high concentrations in the mucosal surface fluids such as the serum, saliva, sweat, and tears. Lysozyme functions as an N-acetylmuramidase glycanhydrolase to cleave 1,4-beta-linkages between N-acetylmuramic acid (NAM) and N-acetylglycosamine (NAG) in the bacterial peptidoglycan, which leads to the lysis of the bacterial cell wall [3]. Lysozyme may also exert a bactericidal effect, via a non-enzymatic mechanism, with its cationic antimicrobial peptide activity and hydrophobic properties inducing cell death via membrane perturbation [4-8]. In addition, the muramidase activity of lysozyme may be involved in the modulation of the immune response and inflammation.

Every attack triggers a counter-attack in the world of microorganisms, hence, it is not surprising that bacteria have developed sophisticated strategies to resist the activity of lysozyme. Gram-positive bacteria, such as *Streptococcus pneumoniae* and *Staphylococcus aureus*, can increase the resistance to lysozyme by modifying their peptidoglycan [9,10]. Well-known modifications include N-deacetylation of NAG, O-acetylation of NAM, N-glycolylation of NAM, and, more recently, O-acetylation of NAG [11,12]. Gram-negative bacteria are surrounded by a double cell wall that renders them naturally impermeable to lysozyme, and therefore, are usually thought to be insensitive to its effects; however, chemical modifications that render the peptidoglycan lysozyme-resistant have also been reported. Additionally, Gram-negative bacteria produce highly specific and potent proteinaceous lysozyme inhibitors [10]. The first known lysozyme inhibitor, Ivy (inhibitor of a vertebrate lysozyme), was identified in *Escherichia coli* strain MG1655 in 2001 [13]. Later, additional periplasmic and/or membrane-bound lysozyme inhibitors were identified.
and characterized, including PliC, MliC \[14\], PliG \[15\], PliI \[16\], and Ts3 \[17\].

In the course of exploratory investigations in our laboratory, we have discovered that ExPEC strains are more resistant to lysozyme than the \textit{E. coli} strain MG1655. The currently known mechanisms of lysozyme resistance cannot explain this difference, since the proteinaceous lysozyme inhibitors characterized to date are encoded by both pathogenic and commensal \textit{E. coli}. In addition to those identified proteinaceous inhibitors, it has been also reported that \textit{E. coli} lipopolysaccharide (LPS) can bind to lysozyme and inhibit its enzymatic activity \[18\]; however, the underlying mechanisms are still elusive. We addressed the molecular basis of our initial observations in the current study. We determined that LPS from \textit{E. coli} MG1655 is truncated and lacks the O-specific polysaccharide. The O-specific polysaccharide of ExPEC inhibited the hydrolytic activity of lysozyme. Further, the O-specific polysaccharide was also required by the complete lipopolysaccharide (LPS)-mediated protection of ExPEC against the bactericidal activity of lysozyme.

## Results

### ExPEC strains are more resistant to lysozyme than nonpathogenic \textit{E. coli} K-12

We first established a rapid method to determine the resistance of \textit{E. coli} to lysozyme to facilitate strain screening. \textit{E. coli} cells (1 $\times$ 10\(^8\)) were washed three times with phosphate-buffered saline (PBS) , re-suspended in 1 mL of PBS, incubated with different concentrations of lysozyme (0–50 mg/mL) at 37 $^\circ$C, and then the minimal lytic concentration (MLC) was determined 24 h later (Fig. 1A). Notably, the four tested ExPEC strains were more resistant to lysozyme than nonpathogenic \textit{E. coli} K-12 MG1655 and the laboratory strain BL21. The lysozyme MLC values of the ExPEC strains (NMEC18, NMEC38, NMEC87, and NMEC58) ranged from 6.25 to 12.5 mg/mL, while the lysozyme MLC values of BL21 and MG1655 strains ranged from 0.049 to 0.78 mg/mL (Fig. 1A). This suggested that ExPEC strains were 8- to 16-fold more resistant to lysozyme than \textit{E. coli} K-12 MG1655. These \textit{E. coli} strains were further evaluated by an \textit{in vitro} lysozyme killing assay, where their sensitivity levels to lysozyme were calculated by $N_0/N$, where $N_0$ and $N$ were the colony counts before and 24 h after incubation. The lysozyme susceptibility test confirmed that all four tested ExPEC strains were significantly ($P < 0.01$) more resistant to lysozyme than MG1655 and BL21 strains (Fig. 1B).

#### Identification of NMEC38 strain genes involved in the resistance to lysozyme

A transposon mutant library was constructed using the mini-Tn5 transposon system in strain NMEC38, one of the lysozyme-resistant ExPEC strains; the library contained 15,000 individual transposon mutants. The mutant library was screened as described in the Materials and methods section, to identify mutants with decreased resistance to lysozyme. In total, 25 mutant strains that showed a reproducible and substantial decrease in lysozyme resistance were identified in the screening assay. The insertion sites of mini-Tn5 in the selected 25 mutants were determined by amplifying their flanking DNA regions in arbitrarily primed polymerase chain reactions (PCR), followed by sequencing of the amplified DNA products (between 150- and 750-bp long). For sequence analyses, BLASTX or BLASTN (https://blast.ncbi.nlm.nih.gov/Blast.cgi) hits with

![Figure 1. Lysozyme sensitivity of clinical ExPEC isolates and laboratory \textit{E. coli} strains.](image)

(A) The densities of six \textit{E. coli} strains in the late log phase of growth (OD\(_{600}\) = 2.0) were adjusted to 10\(^8\) CFU/mL, and the cells incubated with different lysozyme concentrations (0–50 mg/mL) in a 96-well microtiter plate. The lysozyme sensitivity was determined based on MLC, which was the lowest concentration of lysozyme to lyse \textit{E. coli} cells following a 24-h incubation at 37 $^\circ$C. (B) The \textit{in vitro} lysozyme killing assay. The degree of bacterial lysozyme sensitivity was calculated by dividing the CFU number prior to treatment by the CFU number after a 24-h exposure to lysozyme ($N_0/N$). Data represent the mean $\pm$ standard deviation (SD) from three independent experiments. **$P < 0.01$ by one-way ANOVA.
the highest scores and lowest $e$ values were identified. Of the 25 mutants, 20 harbored transposon insertions in different genes (Table 3). The identified genes included genes encoding enzymes involved in LPS biosynthesis, central metabolism, and prophage function; genes with putative regulatory functions and Tripartite ATP-independent periplasmic transport (TRAP); and genes with unknown function.

**Novel genes involved in O-antigen biosynthesis contribute to ExPEC resistance to lysozyme**

In eight mutant strains, the transposon disrupted either of five genes homologous to ECOK1_2260, ECOK1_2261, ECOK1_2263, ECOK1_2264, and ECOK1_2265 [19]; these mutant strains were among the most sensitive to lysozyme (Table 3). Sequence analysis of these five genes revealed that they are located on the O-antigen island (between the galF and gnd genes; Fig. 2A) [19], suggesting that they may be novel genes involved in the O-antigen biosynthesis. To determine their functions, five non-polar mutant strains and their respective complemented strains were constructed. LPS of the five mutant strains, five complemented strains, the wild-type strain (NMEC38), and the laboratory strain MG1655 were isolated, and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and silver-staining. Compared with the wild-type strain, all mutant strains showed altered banding patterns in the O-antigen regions (Fig. 2B, 2C), suggesting that all five genes play a role in the O-antigen biosynthesis. These analyses also demonstrated that the O-antigen chain in E. coli K-12 strain MG1655 was truncated. When the genes were reintroduced into the respective mutant strains, the LPS of four complemented strains showed a similar banding pattern as the wild-type parent, confirming that genes ECOK1_2260, ECOK1_2261, ECOK1_2263, and ECOK1_2264 play a role in the O-antigen biosynthesis.

Wild-type, mutant, and complemented strains were then evaluated by an in vitro lysozyme killing assay to confirm the role of these genes in lysozyme resistance (Fig. 3A). In the absence of lysozyme, all mutant strains survived very well in 1 mM Tris-HCl (pH 7.2), with $N_0/N$ values ranging from 0.82 to 1.37, which were not significantly different from those for the wild-type and complemented strains. After a 24-h incubation with lysozyme (5 mg/mL), all mutants showed a substantial reduction in viability, with an almost 10-fold reduction compared to the wild type, while the complemented strains recovered to the wild-type level. These results confirmed that these novel genes involved in O-antigen biosynthesis contributed to the ExPEC resistance to lysozyme.

**Membrane integrity is weakened in ExPEC mutants with defects in O-antigen biosynthesis**

Propidium iodide (PI) staining was employed to examine whether the deletion of newly identified genes involved in the O-antigen biosynthesis would affect E. coli

| Strain or plasmid | Genotype or description | Source |
|-------------------|------------------------|--------|
| **Wild type strains** | | |
| NMEC38 | Serotype: O18 Laboratory stock [32] |
| NMEC18 | Clinical isolate Laboratory stock [32] |
| NMEC58 | Clinical isolate Laboratory stock [32] |
| NMEC87 | Clinical isolate Laboratory stock [32] |
| S17-1 zipr | RK2 tra regulon, pir, host for pir-dependent plasmids |
| MG1655 | F-, λ, galK, rfb-50, rpi-1 |
| **E. coli DH5α** | Cloning host for maintaining the recombinant plasmids |
| **E. coli BL21** | Cloning host for maintaining the recombinant plasmids |
| **Mutant strains** | | |
| N380 | NMEC38 Δ ECOK1_2260 This study |
| N381 | NMEC38 Δ ECOK1_2261 This study |
| N383 | NMEC38 Δ ECOK1_2263 This study |
| N384 | NMEC38 Δ ECOK1_2264 This study |
| N385 | NMEC38 Δ ECOK1_2265 This study |
| N386 | NMEC38 Δ D4D This study |
| N387 | NMEC38 Δ ECOK1_2261–2264 |
| **Complemented strains** | | |
| N380C | N380 with plasmid p380 This study |
| N381C | N381 with plasmid p381 This study |
| N383C | N383 with plasmid p383 This study |
| N384C | N384 with plasmid p384 This study |
| N385C | N385 with plasmid p385 This study |
| N386C | N386 with plasmid p386 This study |
| N387C | N387 with plasmid p387 This study |
| **Plasmids** | | |
| pKD3 | Template for λ-Red Chl cassette [35] |
| pKD4 | Template for λ-Red Kan cassette [35] |
| pCP20 | Encodes FLP recombinase for the removal of resistance cassette [33] |
| pKD46 | λ-Red recombinase expression [35] |
| pGEN-MCS | Low-copy plasmid for complementation [42] |
| pGEN/pbla | MCS was replaced by the promoter region of ampicillin [36] |
| pGEN/pcm | MCS was replaced by the promoter region of chloramphenicol |
| p380 | pGEN/pcm carrying 2260 coding region |
| p381 | pGEN/pbla carrying 2261 coding region |
| p383 | pGEN/pbla carrying 2263 coding region |
| p384 | pGEN/pbla carrying 2264 coding region |
| p385 | pGEN/pcm carrying 2265 coding region |
| p386 | pGEN/pcm carrying rfbD coding region |
| P387 | pGEN/pbla carrying coding regions from 2261 to 2264 This study |
membrane permeability [20,21]. Bacterial cells with intact membranes are impermeable to charged fluorescent dyes, such as PI; however, if the membrane integrity is compromised, PI can enter the cell and, by binding to the nucleic acid, render the cell fluorescent. The PI uptake in all mutant strains was increased in comparison with that in the wild type, but the difference was only significant in the mutant strain ΔECOK1_2261 (Fig. 3B, P < 0.05). A protein leakage assay was performed to further evaluate membrane integrity of the wild-type and O-antigen gene mutant strains [20,21]. Similar to the results of the PI staining assay, all mutant strains showed increased protein leakage; however, the difference with the wild type was significant only in the mutant strain ΔECOK1_2261 (Fig. 3C, P < 0.05). Taken together, these results suggested that the deletion of the O-antigen synthesis genes affected E. coli membrane integrity, but might not have been the major mechanism of the reduced lysozyme resistance of mutants. 

\[ \text{O-specific polysaccharide is necessary for the LPS-mediated protection of ExPEC from the bactericidal activity of lysozyme} \]

To investigate the mechanism(s) whereby the O-antigen synthesis genes contribute to lysozyme resistance of

| Primer | Sequence (5’–3’) | Target gene, locus, or application |
|--------|------------------|-----------------------------------|
| P6     | CCTAGGCGGCCGACGATC     | For transposon identification |
| P9     | CGCCGCGCTTATGATCC     | For transposon identification |
| Arbi2  | GCCGCCGGCTCAGTAC       | For transposon identification |
| Arbi5  | GCCGCCGCGCTGACTAGTAC(N)   | For transposon identification |

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**Table 2. Primers used in this study.**

| Primer (for gene deletion) | Sequence (5’–3’) | Target gene, locus, or application |
|---------------------------|------------------|-----------------------------------|
| ΔD380F                    | TTTCGACTCATATTAATATTGAGTGTAGTGTGTTTGAACCGGTAACATATTGtgtaggcctggcgctggagctgcttcga | ECO1_2260 |
| ΔD380R                    | TTATTTATTTTACCTTTGAGAATGTTGTTTATGAGAAAATAAAATGgtgtaggcctggcgctggagctgcttcga | ECO1_2261 |
| ΔD381F                    | AAGATGCTAATATTGCTTATTGTAATTTGTAATTTGGAATAAAATGgtgtaggcctggcgctggagctgcttcga | ECO1_2262 |
| ΔD381R                    | TTATTTATTTTACCTTTGAGAATGTTGTTTATGAGAAAATAAAATGgtgtaggcctggcgctggagctgcttcga | ECO1_2263 |
| ΔD383R                    | AACTCCCGATGCCATAAAAAAAATGgtgtaggcctggcgctggagctgcttcga | ECO1_2264 |
| ΔD384F                    | AACTCCCGATGCCATAAAAAAAATGgtgtaggcctggcgctggagctgcttcga | ECO1_2265 |
| ΔD384R                    | AACTCCCGATGCCATAAAAAAAATGgtgtaggcctggcgctggagctgcttcga | ECO1_2266 |
| ΔD386F                    | GGTAGTTGGTGGTCAGTGgtgtaggcctggcgctggagctgcttcga | rfbD |
| ΔD386R                    | GGTAGTTGGTGGTCAGTGgtgtaggcctggcgctggagctgcttcga | ECO1_2267 |

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**Primers** for gene deletion:

| Primer | Sequence (5’–3’) | Target gene, locus, or application |
|--------|------------------|-----------------------------------|
| ΔC380F | GTGTTTTCGGAATCGTGAGC | ECO1_2260 |
| ΔC380R | CTTTCGATGTTGAGCGCGAG | ECO1_2261 |
| ΔC381F | ACTTCCGGATGCGATTAA | ECO1_2262 |
| ΔC381R | ACTTCCGGATGCGATTAA | ECO1_2263 |
| ΔC383F | CTGAAACCGCTAGTAACGA | ECO1_2264 |
| ΔC383R | CTGAAACCGCTAGTAACGA | ECO1_2265 |
| ΔC384F | GGTAGTTGGTGGTCAGTG | ECO1_2266 |
| ΔC384R | GGTAGTTGGTGGTCAGTG | ECO1_2267 |
| ΔC385F | GGTAGTTGGTGGTCAGTG | Km |
| ΔC385R | GGTAGTTGGTGGTCAGTG | cm |
| ΔC386F | GGTAGTTGGTGGTCAGTG | ECO1_2268 |
| ΔC386R | GGTAGTTGGTGGTCAGTG | ECO1_2269 |

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**Primers for gene deletion confirmation**

| Primer | Sequence (5’–3’) | Target gene, locus, or application |
|--------|------------------|-----------------------------------|
| ΔC380F | GTGTTTTCGGAATCGTGAGC | ECO1_2260 |
| ΔC380R | CTTTCGATGTTGAGCGCGAG | ECO1_2261 |
| ΔC381F | ACTTCCGGATGCGATTAA | ECO1_2262 |
| ΔC381R | ACTTCCGGATGCGATTAA | ECO1_2263 |
| ΔC383F | CTGAAACCGCTAGTAACGA | ECO1_2264 |
| ΔC383R | CTGAAACCGCTAGTAACGA | ECO1_2265 |
| ΔC384F | GGTAGTTGGTGGTCAGTG | ECO1_2266 |
| ΔC384R | GGTAGTTGGTGGTCAGTG | ECO1_2267 |
| ΔC385F | GGTAGTTGGTGGTCAGTG | Km |
| ΔC385R | GGTAGTTGGTGGTCAGTG | cm |
| ΔC386F | GGTAGTTGGTGGTCAGTG | ECO1_2268 |
| ΔC386R | GGTAGTTGGTGGTCAGTG | ECO1_2269 |

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**Primers for complementation experiments**

| Primer | Sequence (5’–3’) | Target gene, locus, or application |
|--------|------------------|-----------------------------------|
| p380F  | ACGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2260 |
| p380R  | TGGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2261 |
| p381F  | ACGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2262 |
| p381R  | TGGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2263 |
| p383F  | ACGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2264 |
| p383R  | TGGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2265 |
| p384F  | ACGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2266 |
| p384R  | TGGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2267 |
| p385F  | ACGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2268 |
| p385R  | TGGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2269 |
| p386F  | ACGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2260 |
| p386R  | TGGCTGGCAGATGACAGCTGAAACACTAA | ECO1_2261 |
| pPro384F | ACGCGCTCAAGGCGACGATGACAGCTGAAACACTAA | ECO1_2260 |

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1. The underlined sequences are homologous to the pKD3 or pKD4 vector sequence flanking the Cm' or Km' genes.
2. Iitalicized sequences are homologous to the target gene flanking sequence.
3. The underlined sequences are the enzyme digestion sites.

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These results suggested that the deletion of the O-antigen synthesis genes affected E. coli membrane integrity, but might not have been the major mechanism of the reduced lysozyme resistance of mutants.
| Recombinant class       | Recombinant | MLC* (mg/mL) | Ref. strain | Gene/locus tag | Accession No. b | Identity (%) | Putative function                                                                 |
|-------------------------|-------------|--------------|-------------|----------------|-----------------|--------------|----------------------------------------------------------------------------------|
| LPS synthesis           | F72         | 5            | IHE3034     | rfaJ           | ADE89653        | 97           | Lipopolysaccharide 1,2-glucosyltransferase                                        |
|                         | M62         | 6.25         | IHE3034     | rfbD           | ADE92608        | 100          | Glycosyl transferase, group 1 family protein                                      |
|                         | U424        | 3.125        | IHE3034     | ECOK1_2260     | ADE93999        | 97           | dTDP-4-dehydroxamino reductase                                                   |
|                         | U475        | 6.25         | IHE3034     | ECOK1_2263     | ADE92801        | 91           | Glycosyl transferase, group 2                                                    |
|                         | k262        | 1.56         | IHE3034     | ECOK1_2264     | ADE96668        | 97           | Putative membrane protein                                                        |
|                         | F172        | 0.3125       | IHE3034     | ECOK1_2261     | ADE91992        | 98           | Glycosyl transferase, group 1 family protein                                      |
|                         | J399        | 1.56         |             |                |                 | 98           |                                                                                  |
|                         | U446        | 0.78         |             |                |                 | 98           |                                                                                  |
|                         | T358        | 6.25         | IHE3034     | ECOK1_2265     | ADE9127         | 87           | Putative O-antigen transporter                                                   |
|                         | V66         | 6.25         |             |                |                 | 98           |                                                                                  |
|                         | U504        | 6.25         | IHE3034     | rfaI           | ADE88840        | 99           | Lipopolysaccharide 1,3-galactosyltransferase                                     |
|                         | V501        | 6.25         |             |                |                 | 98           |                                                                                  |
|                         | A2          | 6.25         | IHE3034     | ECOK1_4065     | ADE9362         | 94           | Lipid A-core surface O-antigen polymerase                                           |
|                         | L208        | 3.125        |             |                |                 | 98           |                                                                                  |
| Metabolism              | F185        | 5            | IHE3034     | pepA           | ADE89150        | 98           | Leucyl aminopeptidase                                                            |
|                         | G295        | 5            | IHE3034     | allD           | ADE93999        | 99           | Glycosyl transferase, group 2 family protein                                      |
|                         | T244        | 6.25         | IHE3034     | Pgi            | ADE90626        | 95           | Ureidoglycolate dehydrogenase                                                   |
|                         | K240        | 3.125        | IHE3034     | dsbA           | ADE9344         | 98           | Glucose-6-phosphate isomerase                                                   |
|                         | T352        | 6.25         | IHE3034     | galU           | ADE90279        | 90           | Thiol/disulfide interchange protein DsbA                                          |
|                         | V153        | 6.25         | IHE3034     |                |                 |              |                                                                                  |
| Prophage                | F199        | 5            | CFT073      | intC           | AE016769        | 92           | Putative prophage integrase                                                      |
| Regulator               | B54         | 6.25         | IHE3034     | dhaR           | ADE9325         | 91           | PTS-dependent Dihydroxyacetone kinase operon regulatory protein                  |
| Transporter             | Q159        | 1.56         | IHE3034     | ECOK1_4026     | ADE92303        | 92           | TRAP transporter, DctM family                                                   |
| Unknown                 | G413        | 6.25         | pECOS88     | ECS88_p0136    | CAQ7198         | 98           | Hypothetical protein                                                             |
|                         | H157        | 6.25         | E.coli W    | WFL_20590      | AFH13760        | 96           | Putative repressor protein C                                                     |

*MLC, minimal lytic concentration
ExPEC, we tested whether ExPEC LPS could inhibit the bactericidal activity of lysozyme. Complete LPS was extracted from NMEC38 (wild type) by a hot phenol-water method [18,22]. The mutant strain DECK1_2265, which was the most lysozyme-susceptible mutant strain, was cultured to the late exponential phase (10⁹ CFU/mL) and used in in vitro lysozyme killing assay. The mutant strain DECK1_2265 was incubated with 5 mg/mL of lysozyme and different concentrations of wild-type LPS in 1 mM Tris-HCl (pH 7.2); the survivors were enumerated after 24 h. In the absence of wild-type ExPEC LPS, the lysozyme resulted in an approximately 16-fold bacterial reduction and killed most mutant cells; in contrast, upon the addition of 0.12 mg/mL of LPS, the lysozyme resulted in only a 10.8-fold bacterial reduction (Fig. 4A). When more than 0.3 mg/mL of LPS was added, most mutant cells survived the lysozyme treatment, showing a 1.57- to 2.3-fold bacterial reduction (Fig. 4A), suggesting that wild-type LPS could indeed inhibit the bactericidal activity of lysozyme.

Since LPS consists of a lipid A unit, core oligosaccharide and an O-specific polysaccharide chain, the ability of LPS lacking the O-specific polysaccharide and the O-specific polysaccharide alone to inhibit the bactericidal activity of lysozyme was next tested. LPS lacking the O-specific polysaccharide was extracted from mutant strains DECK1_2260, DECK1_2261, DECK1_2263, DECK1_2264, and DECK1_2265; and the O-specific polysaccharide was purified from NMEC38 wild type. The ability of LPS lacking the O-specific polysaccharide and the O-specific polysaccharide alone to inhibit the bactericidal activity of lysozyme was significantly (P < 0.001) reduced compared to complete LPS (Fig. 4B). These results indicated that the O-specific polysaccharide alone cannot inhibit the bactericidal activity of lysozyme but is necessary for the LPS-mediated protection of ExPEC against this activity.

**O-specific polysaccharide from NMEC inhibits the hydrolytic activity of lysozyme**

LPS from NMEC38 wild type was further examined by a turbidity assay described by Callewaert [14] to determine whether it could inhibit the hydrolytic activity of lysozyme. When *Micrococcus lysodeikticus* ATCC 4698 was incubated with 4 μg/mL of lysozyme...
in the absence of LPS, the OD600 value of the reaction mixture dropped close to zero after 2 h, suggesting that *M. lysodeikticus* ATCC 4698 cells were almost entirely lysed (Fig. 4C). When LPS (>50 μg/mL) was included in the reaction mixture, the OD600 value of the reaction remained constant, with a turbidity similar to that of the negative control incubated in the absence of lysozyme (Fig. 4C), suggesting that the lytic activity of lysozyme was completely inhibited. These results indicated that LPS from NMEC38 could inhibit the hydrolytic activity of lysozyme.

We next investigated whether LPS lacking the O-specific polysaccharide or the O-specific polysaccharide alone could inhibit the hydrolytic activity of lysozyme. The experiment (Fig. 4D) revealed that the ability of LPS that lacked the O-specific polysaccharide (purified from ExPEC mutant strains) to inhibit the hydrolytic activity of lysozyme was significantly reduced compared to the complete LPS. Similarly, LPS purified from *E. coli* K-12 strain MG1655 was unable to inhibit that activity. The O-specific polysaccharide from NMEC38 wild-type almost entirely inhibited the hydrolytic activity of lysozyme, emphasizing the pronounced differences between LPS lacking the O-specific polysaccharide and LPS from the *E. coli* K-12 strain MG1655.

**LPS and the O-specific polysaccharide from ExPEC can directly interact with lysozyme**

The direct interaction of LPS with lysozyme was examined by gel filtration chromatography. In this assay, LPS and/or lysozyme were pre-incubated for 30 min in a binding buffer and resolved on an Enrich SEC 650 (10/300) gel filtration column, as specified in the Materials and methods section. The elution profile was then analyzed by measuring absorbance at 215 nm and western

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*Figure 3.* The *in vitro* lysozyme killing assay and membrane integrity tests. (A) Lysozyme sensitivity of the wild-type, mutant, and complemented strains was determined by an *in vitro* killing assay. The strain compromised viability levels (N0/N) under the indicated conditions were analyzed as in Fig. 1B. (B) Membrane integrity of the wild-type, mutant, and complemented strains determined by PI staining. (C) Membrane integrity determined by a protein leakage assay. Data represent the mean ± standard deviation (SD) from three independent experiments. *P < 0.05, **P < 0.01 by one-way ANOVA.
In the absence of LPS, lysozyme was eluted in fractions 13–19 (Fig. 5A). In the absence of lysozyme, LPS eluted as two peaks; however, only the first peak contained LPS, as revealed by western blotting (fractions 3–5, Fig. 5B). Similar observations have been reported by Miki et al [22]. Nevertheless, co-incubation of lysozyme with LPS resulted in the occurrence of lysozyme in both peaks in fractions A4–A7 and 13–19 (Fig. 5C). The lysozyme-LPS complex in the pronouncedly forward-shifted peak A4–A7 was confirmed by western blotting (with anti-lysozyme antibody) and LPS (by silver-staining) (Fig. 5C). The results strongly suggested that LPS bound lysozyme directly.

The interaction of the O-specific polysaccharide to lysozyme was also analyzed by gel filtration chromatography. The O-specific polysaccharide isolated from NMEC38 was pre-incubated in the presence or absence of lysozyme, and then loaded onto the Enrich SEC 650 (10/300) column. The elution profile was analyzed by monitoring A215. In the lysozyme-only sample, lysozyme was eluted in fractions number 13–18 (Fig. 6A). As shown in Fig. 6B, in the absence of lysozyme, the O-polysaccharide was eluted in fractions 14–18. Co-incubation of lysozyme with the O-polysaccharide resulted in only one peak, which was pronouncedly forward-shifted compared to the peak of the purified O-polysaccharide, confirming the interaction of O-polysaccharide to the lysozyme (Fig. 6C).

**LPS from ExPEC strains with different O serotypes inhibit the hydrolytic and bactericidal activities of lysozyme**

To examine if LPS from strains with different O serotypes could inhibit the hydrolytic activity of lysozyme, additional 27 ExPEC strains with different O serotypes (including O1, O2, O18, O8, and untypeable serotypes) isolated at different times and from
different regions [23], were examined. LPS from 24 strains (89.7%) was able to completely inhibit the hydrolytic activity of lysozyme (Fig. 7A); however, this effect was almost abolished in the case of LPS from three ExPEC strains (10.3%) with untypeable O serotypes (DE005, DE207, and DE477; Fig. 7A). SDS-PAGE and silver-staining revealed that similar to that in *E. coli* K-12 strain MG1655, LPS from those three strains was truncated and lacked the O-polysaccharide (Fig. 7B). Furthermore, the purified LPS from ExPEC with O1, O2, and O18 could protect mutant strain D*ECOK1*2265 from the bactericidal activity of lysozyme (Fig. 7C). To rule out influence of other surface antigens for example capsular antigen, we deleted the capsule genes (*ECOK1*3365-neuD) and found no significant difference between wild type and the capsule mutant in lysozyme in vitro killing assay. As the control, deletion of O antigen biosynthesis gene (*ΔECOK1*2265) significantly reduced ExPEC’s resistance to lysozyme (Fig. 7D). We further purified LPS from capsule deleted mutant strain (*ΔECOK1*3365-neuD) and showed that capsule polysaccharides were not necessary for the LPS-mediated protection of ExPEC from the bactericidal activity of lysozyme (Fig. 7E).

**Discussion**

LPS consisting of a lipid A unit, core oligosaccharide and an O-specific polysaccharide chain is known to interact with lysozyme and inhibit its enzymatic activity [18]. Although the binding of lipid A and a synthetic monosaccharide lipid A analogue to lysozyme has been experimentally confirmed [24], it remains unknown whether lipid A and/or the O-specific polysaccharide indeed inhibit lysozyme activity. In the current study, we identified several genes involved in the synthesis of the O-specific polysaccharide, which contributed to bacterial lysozyme resistance. Deletion of these genes resulted in truncated LPS that lacked the O-polysaccharide, which substantially decreased the resistance of ExPEC mutants to the bactericidal activity of lysozyme. Correspondingly, LPS that lacked the O-specific polysaccharide purified from the constructed mutant strains was characterized by a significantly lower ability to inhibit the bactericidal and hydrolytic activities of lysozyme than the complete

![Figure 5](image-url)
LPS purified from the ExPEC wild type. LPS was likely to self-aggregate into supramolecular structures and directly interact with lysozyme, thus inhibit lysozyme’s activity [25]. Furthermore, our results also indicated that the purified O-specific polysaccharide alone could inhibit the hydrolytic activity of lysozyme by a direct interaction with the enzyme. To the best of our knowledge, this is the first-ever report that O-specific polysaccharide from ExPEC contributes to bacterial lysozyme resistance.

Lysozyme kills Gram-positive bacteria by cleaving the cell wall peptidoglycan. In addition, accumulating evidence suggests that lysozyme can kill both Gram-positive and -negative bacteria also independently of its enzymatic muramidase activity [4,5,7], with its cationic antimicrobial peptide activity and hydrophobic properties playing an important role [8]. We demonstrated that the complete LPS from ExPEC wild type was able to inhibit the bactericidal activity of lysozyme, while neither LPS lacking the O-specific polysaccharide nor the O-specific polysaccharide alone showed this activity. These observations suggest that the O-specific polysaccharide is required for LPS-mediated inhibition of the bactericidal activity of lysozyme, but by itself, the O-specific polysaccharide lacks this activity. On the other hand, the O-specific polysaccharide alone was able to entirely inhibit the enzymatic and lytic activities of lysozyme, independently of the remaining portion of LPS. The different mechanisms whereby LPS inhibits the various activities of lysozyme require further investigation.

Gram-negative bacteria are generally thought to be more resistant to lysozyme than their Gram-positive counterparts. This difference in resistance is primarily
ascribed to the different cell envelope architecture of these two groups of bacteria. Gram-positive bacteria have a thick cell wall composed of up to 40 layers of peptidoglycan, which is very sensitive to lysozyme [26], while Gram-negative bacteria typically have only a single layer of peptidoglycan surrounded by an asymmetric membrane bilayer, which is thought to render Gram-negative bacteria naturally impermeable to lysozyme, and thus resistant to its effects [26]. However, recent evidence indicates that lysozyme can interact with the negatively charged membrane lipid bilayers leading to protein aggregation and membrane fusion [27], and can permeabilize the outer and inner membranes of an E. coli mutant ML-35p by inducing the formation of large pores [6,28]. This suggests that the physical barrier afforded by the outer membrane of Gram-negative bacteria might not be the main reason responsible for their greater resistance to lysozyme. LPS, with the O-specific polysaccharide, is a unique and common feature of all Gram-negative bacteria, and our findings suggest that this structure might be responsible for the elevated lysozyme resistance of Gram-negative bacteria. The observation that the O-specific polysaccharide-related lysozyme inhibition activity is independent of the E. coli serotype...
(Fig. 7) and that Salmonella LPS that contains the O-specific polysaccharide can also inhibit lysozyme hydrolytic activity (Yinli Bao, data not published) further support this hypothesis.

In addition to the novel O-specific polysaccharide biosynthesis genes and other LPS biosynthesis genes, we also identified metabolism, regulatory, and transport genes involved in bacterial resistance to lysozyme. However, the genes for proteinaceous lysozyme inhibitors, such as Ivy [13], MliC (membrane bound lysozyme inhibitor of c-type lysozyme) [14], and PliG (periplasmic lysozyme inhibitor of g-type lysozyme) [15,29], were not identified in our screen. The performed deletion and lysozyme inhibition assays indicated that these “other” genes did not appear to play a significant role in the ExPEC resistance to lysozyme in our in vitro model (Yinli Bao, data not shown). Recently, Ivy and its homolog were shown to be potent inhibitors of lytic transglycosylases involved in the biosynthesis and maintenance of peptidoglycan [30]. In addition, mliC or its homolog is adjacent to the annmK gene in both E. coli and Salmonella, which encodes an anhydro-NAM kinase involved in the recycling of murein [14,31]. These findings suggest that the true physiological functions of Ivy, MliC, and PliG might be to control excessive activity of endogenous bacterial autolysins, with the inhibition of exogenous lysozyme as a simply fortuitous coincidence. The localization of the proteinaceous inhibitors to the periplasm (MliC is bound to the luminal side of the outer membrane) [13-15,29] rather than to the external milieu further suggests that the O-specific polysaccharide at the bacterial surface may play a much more important role in protecting bacteria against exogenous lysozyme than proteinaceous inhibitors.

In summary, the present study demonstrated that the O-specific polysaccharide from ExPEC contributes to the LPS-mediated inhibition of the bactericidal activity of lysozyme, and the O-specific polysaccharide alone is able to inhibit the hydrolytic activity of lysozyme through a direct interaction with the enzyme. While these findings were only acquired in a single strain, the observed localization to external milieu and wide distribution of O-specific polysaccharide in Gram-negative bacteria suggests that this surface structure might play an under-appreciated role in protecting bacteria against exogenous lysozyme.

**Materials and methods**

**Strains and plasmids**

Bacterial strains and plasmids used in this study are listed in Table 1. The transposon mutagenesis library and mutants were created in the ExPEC NMEC38 background [32]. Additional 27 ExPEC strains with different O serotypes [23] were used to evaluate the correlation between the structure of LPS and bacterial resistance to lysozyme. All E. coli strains were grown at 37 °C in Luria-Bertani (LB) broth and agar. Antibiotics, including nalidixic acid (30 μg/mL), kanamycin (50 μg/mL), ampicillin (50 μg/mL) and chloramphenicol (30 μg/mL), were added when necessary.

**Mutant library construction, screening, and transposon insertion site identification**

More details about crucial steps in the genetic studies were provided in supplementary materials. Briefly, a transposon library of 15,000 mutants was generated using the transposon delivery vector pUTmini-Tn5 (Km), as described previously [33]. All mutants were inoculated in 96-well, U-bottom plates containing 1 mL of sterile LB medium and grown at 37 °C to late exponential phase (OD

**Recombinant DNA techniques, SDS-PAGE, and western blotting**

PCR, DNA ligation, electroporation, and gel electrophoresis were performed according to Sambrook and Russel [34], unless indicated otherwise. All oligonucleotide primers were purchased from Integrated DNA Technologies and are listed in Table 2. All restriction and DNA-modifying enzymes were purchased from New England Biolabs and were used as per the supplier’s recommendations. Recombinant plasmids, PCR products, and restriction fragments were purified using QIAquick PCR purification kit or MinElute gel extraction kit (Qiagen), as recommended by the supplier. DNA sequencing was performed at the DNA Facility, Iowa State University (Ames, IA).
Deletion mutants were constructed using the λ red mutagenesis method [35]; all primers used for mutant construction are listed in Table 2. To construct the plasmids for mutant complementation, the target genes and their native promoters were amplified (the primers are listed in Table 2) from ExPEC wild-type strain (NMEC38) and digested by SalI; they were then ligated into SalI-digested plasmid pGEN/pbla [36]. The resulting plasmids were used to complement the corresponding mutants.

SDS-PAGE and silver-staining of LPS were performed according to the protocol described by Tsai and Frach [37]. Western blotting was performed according to Sambrook and Russel [34], using a semidry blotting apparatus, on polyvinylidene difluoride membranes (Thermo Fisher Scientific). The rabbit anti-\(E.\ coli\) O18 antibody (1:1000), horseradish peroxidase-conjugated goat anti-rabbit secondary antibody (1:5000, #31466) and the membrane developing DAB kit were purchased from Thermo Fisher Scientific.

**Lysozyme in vitro killing assay**

The \textit{in vitro} killing assay was performed as described previously [38]. The killing activity was calculated by \(N_0/N\), where \(N_0\) and \(N\) were the colony counts before and 24-h after incubation, respectively. To block lysozyme bactericidal activity, 100 \(\mu\)L of serial dilutions of LPS, LPS lacking the O-specific polysaccharide, or the purified O-specific polysaccharide were used. The samples without LPS or lysozyme were used as controls.

**Membrane integrity determinations**

Fluorescence PI staining measurement and protein leakage assays were performed to determine the integrity of the bacterial membrane. PI staining assay was performed as described by Garcia-Gonzalez et al. [20,21], and the protein leakage assay was performed as described previously [20,21]. Protein concentrations in reaction supernatants were determined by the BCA protein assay Kit (Pierce).

**LPS purification and preparation of the O-specific polysaccharide**

LPS was extracted by the hot phenol-water method as described previously [18,22], with minor modifications. Briefly, 100 mL of bacterial culture was grown to late logarithmic phase (\(OD_{600} = 2\)), centrifuged, washed twice with PBS (pH 7.2), and re-suspended in 20 mL of ddH\(_2\)O. An equal volume of hot 90\% (v/v) phenol (68 °C) was slowly added to the mixtures, followed by a vigorous shaking at 68 °C for 30 min. Suspensions were then cooled on ice and centrifuged at 2,851 \(\times\) g for 45 min at 10 °C. Supernatants were transferred to 50-mL conical tubes; ten volumes of cold absolute ethanol and sodium acetate (to final concentration of 0.3 M) were added, and mixed thoroughly. The reactions were stored at -20 °C overnight and then centrifuged at 2,000 \(\times\) g for 10 min at 4 °C. The LPS pellets were finally re-suspended in 1 mL of ddH\(_2\)O. The residual phenol, nucleic acids, and proteins were removed by dialysis against ddH\(_2\)O and treatment by Dnase I, RNase A, or proteinase K, respectively. SDS-PAGE and agarose gel electrophoreses were performed to examine protein and/or nucleic acid contamination, respectively. The purified LPS was re-suspended in ddH\(_2\)O, and its concentration was determined by using the anthrone-sulfuric acid method [39].

The O-specific polysaccharide from LPS was prepared by hydrolyzing LPS with 1\% acetic acid at 100 °C for 2 h. The precipitated lipid A was removed by low-speed centrifugation at 524 \(\times\) g for 30 min [40,41]. Water-soluble fractions were then cooled on ice and centrifuged for 10 min at 4 °C. The LPS pellets were finally re-suspended in 1 mL of ddH\(_2\)O. The residual phenol, nucleic acids, and proteins were removed by dialysis against ddH\(_2\)O and treatment by Dnase I, RNase A, or proteinase K, respectively. SDS-PAGE and agarose gel electrophoreses were performed to examine protein and/or nucleic acid contamination, respectively. The purified LPS was re-suspended in ddH\(_2\)O, and its concentration was determined by using the anthrone-sulfuric acid method [39].

**Inhibition of the hydrolytic activity of lysozyme**

The inhibition of lysozyme activity by LPS or the O-specific polysaccharide was determined as described by Callawaert [14] using \(M.\ lysodeikticus\) ATCC 4698 (Sigma) and lysozyme from chicken egg white (Sigma).

**Gel filtration chromatography**

Gel filtration chromatography was carried out using an Enrich SEC 650 (10/300) column (Bio Rad) [18,22]. For analysis, 500 \(\mu\)L of lysozyme (500 \(\mu\)g/mL) and 500 \(\mu\)L of LPS (500 \(\mu\)g/mL) were mixed and then incubated for 30 min at 37 °C at room temperature, and applied to the column at a flow rate of 1.0 mL/min. The elution profile was analyzed by monitoring A215, and by SDS-PAGE and western blot analyses of the fractions using an affinity-purified anti-lysozyme antibody (1:500 dilution, ab391, Abcam).

**Statistical analysis**

Statistical analyses were performed using Prism 5.0 (GraphPad). One-way ANOVA was performed to analyze the \textit{in vitro} killing assay data and for membrane integrity determination. Differences were deemed to be statistically significant at \(P < 0.05\).
Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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