Real-time detection of antibiotic activity by measuring nanometer-scale bacterial deformation

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Abstract. Diagnosing antibiotic-resistant bacteria currently requires sensitive detection of phenotypic changes associated with antibiotic action on bacteria. Here, we present an optical imaging-based approach to quantify bacterial membrane deformation as a phenotypic feature in real-time with a nanometer scale (~9 nm) detection limit. Using this approach, we found two types of antibiotic-induced membrane deformations in different bacterial strains: polymyxin B induced relatively uniform spatial deformation of Escherichia coli O157:H7 cells leading to change in cellular volume and ampicillin-induced localized spatial deformation leading to the formation of bulges or protrusions on uropathogenic E. coli CFT073 cells. We anticipate that the approach will contribute to understanding of antibiotic phenotypic effects on bacteria with a potential for applications in rapid antibiotic susceptibility testing.

Keywords: cell membrane deformation; Escherichia coli; antibiotics; polymyxin; ampicillin.

1 Introduction

Antibiotic resistance poses serious threats to humans with multi-drug-resistant bacteria now commonly known as “superbugs.” 1-4 To combat antibiotic-resistant bacteria, one must develop more effective and rapid antibiotic susceptibility testing (AST) methods. 5 Current AST technologies used in clinic labs rely on cell culture, which can take 1 to 14 days, depending on the growth rate of the bacteria. 6,7 A critical task in AST is to quickly detect if an antibiotic at a given concentration has a phenotypic effect on the bacteria. One commonly monitored phenotypic feature is the bacterial cell morphology visualized via optical microscopy. 8 Other phenotypic features explored for rapid AST include bacterial nanomotion associated bacterial metabolic activities, 9 measuring bacterial growth with magnetic bead rotation sensors, 10 nanometer (nm)-scaled bacterial cell motions measured using atomic force microscopy, 11 changes in plasmic imaging responses, 12 and detection of chemical signatures with electrochemical methods. 13

In the present work, we report an optical method to track nm-scale membrane deformations in individual bacterial cells with the goal to detect antibiotic action at the earliest possible stage. The method detects and quantifies subtle bacterial membrane deformations that are difficult to detect using traditional optical imaging analysis methods. 14,15 We study the membrane deformations of two different Escherichia coli strains, E. coli O157:H7 and uropathogenic E. coli CFT073, 16 associated with antibiotics, polymyxin B (PMB) and ampicillin, respectively.

2 Materials and Methods

2.1 Overview

E. coli O157:H7 and uropathogenic E. coli (UPEC) strain CFT073 cultures were prepared by diluting overnight cultures into fresh Luria broth (LB) and continuing growth for 2 h at 37°C with gentle rotary mixing until the cells reached the mid-exponential growth phase. The individual bacterial cells were tethered on a glass slide via antibody coupling. For antibiotic testing, bacterial cells were incubated in LB or phosphate-buffered saline (PBS) prior to the addition of ampicillin or PMB, respectively. Optical images of the antibiotic effects on the bacteria were obtained with an inverted optical microscope with a 100x objective (numerical aperture of 1.49) and recorded with a CCD camera [Fig. I(a)]. A key task of our method is to track local cellular membrane deformation, including expansion and contraction, associated with antibiotic action. We describe the local membrane deformation with displacement vectors [Fig. I(b)]. A displacement vector points in the direction normal to the local cell edge with magnitude representing the amount of deformation.

2.2 Differential Detection Method

To track subtle cellular deformation, we used a differential detection method, 17 which can track deformations as small as ~9 nm (described in the Sec. 3). The method establishes a linear relation between cell deformation and the intensity profile of a region of interest (ROI) that includes the cell edge. The ROI is divided into two halves with one half inside and the other halve outside of the cell. The total pixel intensities of...
these two halves are denoted by $I_1$ and $I_2$, respectively [Fig. 2(a)]. When the cell expands, $I_1$ decreases and $I_2$ increases. In contrast, when the cell shrinks, $I_1$ increases and $I_2$ decreases. The edge displacement $d$ is given by

$$d = k \frac{I_2 - I_1}{I_1 + I_2}$$ (1)

where $k$ is a constant obtained through a calibration process. The calibration is performed in the first image frame by moving the ROI normal to the cell edge [Fig. 2(b)]. The differential intensity change versus the ROI movement in terms of pixels reveals a linear relation within a certain range of pixel shift [magnified region of Fig. 2(b)]. To obtain a smooth curve and achieve subpixel sensitivity, the images are interpolated by 5 pixels (without interpolation), which corresponds to an actual cellular deformation range of 300 nm.

The linear regime is typically within 4 pixels (or 20 pixels after interpolation), which corresponds to an actual cellular deformation range of 300 nm.

2.3 Materials

*E. coli* O157:H7 (ATCC 43888) and UPEC CFT073 were purchased from Fisher Scientific and ATCC, respectively. Affinity-purified goat anti-*E. coli* O157:H7 IgG polyclonal antibodies, purchased from Kirkegaard and Perry Laboratory, Inc. (Gaithersburg, Maryland), were added to 1-mL PBS (1×) and stored at −20°C prior to use. PMB and ampicillin were purchased from Sigma-Aldrich, dissolved in PBS (1×) at stock concentrations of 10 and 2 mg/mL, respectively, and stored in dark at 2°C to 8°C. 3-Triethoxysilylpropylamine 99% (APTES), N-Hydroxysuccinimide 98% (NHS), N-Ethyl-N′-(3-dimethylaminopropyl) carbodiimide 97% (EDC), and sodium acetate (NaOAc) were also purchased from Sigma-Aldrich.

2.4 Bacterial Preparation

All bacterial stocks were maintained in 5% glycerol at −80°C in small aliquots (20 μL). After overnight incubation at 37°C, *E. coli* cells were diluted into fresh LB and grown for ~2 h at 37°C with gentle rotary mixing until the cells reached the midexponential growth phase. Bacterial cells were pelleted via centrifugation at 2000 g for 30 min and resuspended in 1 mL of PBS (1×).

2.5 Surface Preparation

Glass coverslips (22 × 22 mm) were cleaned with deionized water (DW) and ethanol and then dried with nitrogen gas. A small aliquot of 1% of APTES in 5% ethanol was applied to the glass slides and allowed to dry at room temperature for 50 to 60 min. After the slides were cleaned and dried, a mixture of 0.1 M NHS and 0.4 M EDC in 1 mL of DW was applied to the slides to produce NHS ester receptors. After 10-min incubation, the NHS/EDC slides were again cleaned thoroughly with water and ethanol. The NHS/EDC-activated surface was
blow dried with nitrogen. After the addition of 30 μg/mL of antibody in 20-mM sodium acetate, pH 5.5 to the slides for 60 min, the slides were cleaned again thoroughly with water and ethanol followed by blow drying with nitrogen, generating an antibody-activated surface for imaging experiments.

2.6 Optical Imaging System

The optical imaging system included an Olympus IX-81 inverted microscope with a 100x oil immersion and a Pike 032B CCD camera (Allied Vision Technologies, Newburyport, Massachusetts) to record videos, typically at 50 frames per second (fps) for the PMB experiments and 11.8 fps for the ampicillin experiments. A fast frame rate was used for the PMB experiments to maximize data collection. Since the ampicillin experiments required long video recording to capture bacterial cell bulge/protrusion events, a slower frame rate was used to reduce the data size.

2.7 Bacterial Immobilization

flexiPERM® (Sarstedt Inc.) reusable wells (0.5 mL) were mounted and filled with 1x PBS or LB for the PMB or ampicillin experiments, respectively, and 10 μL of bacterial cells (~10⁸ cells). After 30-min incubation at 37°C to allow bacterial binding to the surface-attached antibodies, unattached bacteria were removed using the flow setup (described below).

![Fig. 3 Schematic for the drug perfusion system. As PBS or PMB was injected into the well, the suction pump concomitantly removed excess liquid from the well, thereby maintaining a constant solution volume.](image)

![Fig. 4 Length change results for noise analysis. (a) A dust spot in the camera was selected to assess the noise of the algorithm without external factors. The signal was pure noise with standard deviation σ = 1.66 nm. (b) A silicon bead of 1 μm was placed on the chip surface. Scale bar: 200 nm. The silicon bead showed some small fluctuations and long-term noise, caused by external factors, such as small focus changes due to instabilities of the microscope stage. The short-term noise for a window of 10 s showed σ = 3 nm. The limit of detection was ~3σ = 9 nm.](image)

![Fig. 5 Length change for a silicon bead when submitted to a 50 mHz and 500 nm peak-to-peak modulation of the stage position in the z-direction (vertical), creating controlled focus changes. The modulation was realized by a piezo microscope stage (Physik Instrumente L.P., Auburn, Massachusetts, model P-545.3C7). From this experiment, there was evidence of long-term decay, due to stage drift, and an oscillation that responds nearly linearly to the modulation, with a peak of ~25 nm. These results indicated that a 10-nm stage change would correspond to a 1-nm (250/25 = 10) measured length change. While short-term measurement windows would not be significantly affected, stage drift could affect interpretation of long-term measurement results.](image)
2.8 Flow Setup

LB, 1× PBS, and PMB (diluted in PBS) were added to the wells through a gravity-based multichannel drug perfusion system with the excess liquid removed by an air pump. A schematic for the flow setup is shown in Fig. 3. The drug perfusion system delivered solutions at a flow rate of 330 μL/min with the transition time between different flow solutions ranging from 1 to 2 s. For the ampicillin experiment, the antibiotics were manually pipetted into the mounted reusable wells atop the antibody surface with immobilized bacteria.

2.9 Image Collection and Processing

All image sequences were collected in transmitted mode at a pixel resolution of 640 × 480. We chose an appropriate exposure time to maximize image intensity and avoid over exposure.

2.10 Data Analysis from Images

Images were processed using custom-written MATLAB programs and the methods described previously in the paper.

Fig. 6 (a) Cell length change of a bacterium (E. coli O157:H7) exposed to PMB (0.5 μg/mL) (red line) compared with PBS (blue line) over time. The arrow indicates the initial expansion shortly after antibiotic introduction. (b) Examples of ROIs and the corresponding displacement vectors for the ROIs. Length changes were determined by summing the two opposite vectors of the red/blue ROIs, and width changes were determined by summing the opposite vectors of the green/yellow ROIs. Scale bar: 200 nm.

Fig. 7 Cell length (blue) and width (red) changes for three E. coli O157:H7 cells. (a–c) Three different cells incubated in PBS buffer only. (d–f) Three different cells exposed to PMB (0.5 μg/mL). The vertical line in each plot at 2 min marked the introduction of PMB. The optical images of each cell at the end of the experiment are shown at the bottom of each response plot (Scale bar: 500 nm).
All recorded images were used for analysis, and all displacement plots were smoothed by averaging 20 frames.

3 Results and Discussion

3.1 Detection Limit

To estimate the detection limit of the method, we performed experiments on dust particles and 1-μm silicon beads on the glass slide (Fig. 4). We estimated a detection limit of ~9 nm (defined as three times the standard deviation). We also tested and measured the system’s response to drift changes in the focus plane (Fig. 5).

3.2 Polymyxin B Results

Using the optical tracking technique described in the previous section, we obtained the local displacement vectors of an E. coli O157:H7 cell before and after exposure to PMB from which we determined the corresponding length changes of the bacterial cell [Fig. 6(a)]. Before exposure to PMB, the bacterial cell incubated in 1× PBS buffer exhibited minimal changes in cell length [blue curve, Fig. 6(a)]. Upon introducing PMB, cell length initially expanded by ~40 nm [arrow, Fig. 6(a)], followed by a large (~200 nm) shrinkage [red curve, Fig. 6(a)]. The PMB antibiotic is a cationic peptide, which elicits bactericidal activity against most gram-negative bacilli, including E. coli O157:H7. The observed PMB-induced bacterial cell deformation arises because PMB binds to the negatively charged lipopolysaccharide layer of the cell and destabilizes the bacterial outer membrane, as the fatty acid portion of PMB dissolves in hydrophobic region of cytoplasmic membrane and disrupts the membrane integrity. Next, we measured PMB-induced membrane deformation (edge displacement via length and width changes) in 40 E. coli O157:H7 cells exposed to a high

![Fig. 8 Histogram analyses of length changes for 40 cells in PBS control (a) and PMB (b) experiments at different time points (after introduction of antibiotics for PMB plots). After antibiotic addition, cell shrinkage increased over time. Comparison of average values between the control and antibiotic experiments for all time points revealed statistically significant differences (\( p = 3.43 \times 10^{-8}; \) Student t-test).]
concentration of PMB (0.5 mg/mL) [Figs. 6(b) and 7]. Length and width changes were calculated by summing opposite displacement vectors, thereby excluding effects caused by changes in cell motion. We intentionally chose a high bactericidal PMB antibiotic concentration, significantly higher than the determined PMB minimum inhibitory concentration (MIC) of 8 μg/mL, to quickly observe deformation in most of the cells, as opposed to using low concentrations where only a few cells showed a significant response. Nearly all cells showed a small, initial expansion in length, followed by a significant decrease in length, which was captured within seconds after PMB addition. Width change in individual cells was less obvious with a few cells displaying decreased width while other cells increased in width. Although there was also variability in the total length decreases between cells, these variations are expected, given the phenotypic heterogeneity of bacterial cells. Bacterial heterogeneity and diversified phenotypic responsiveness underscore the importance of analyzing individual cells with the imaging-based method. Detailed statistical analysis and histograms are shown in Figs. 8 and 9. Compared with traditional AST methods that measure an averaged feature (e.g., turbidity) of the entire sample, single cell analysis could identify a few antibiotic resistant cells among a large population of susceptible cells by tracking the phenotypic responses of the resistant cells to the antibiotic, which could lead to faster AST.

Although the detection limit of the method itself is very small (∼9 nm), external factors, such as the microscope stability (Fig. 3) and the attachment of the bacterial cells to the surface, contribute to the accuracy of the measurement. Microscope focus drift can lead to systematic errors in the measurement of the membrane deformation. Micromotion of cells weakly attached to the surface may also affect the deformation tracking, especially when the cell moves out of the focal plane. Moreover,
cells in the solution may occasionally move on top of a cell on the surface, thereby blocking the surface-immobilized cell from the view of the microscope. These external factors may be reduced by stabilizing the microscope and developing improved cell attachment methods.

### 3.3 Ampicillin Results

We have also studied UPEC CFT073 exposed to the antibiotic, ampicillin, and analyzed the local cellular deformation of individual cells (Fig. 10). Unlike PMB-treated *E. coli* O157: H7, ampicillin causes the UPEC cell to bulge and elongate until cell lysis. Ampicillin targets enzymes that are critical in forming bacterial peptidoglycan, a thin cell wall layer in gram-negative bacteria composed of long glycan chains, which serve to maintain cell structure and balance turgor pressure. By inhibiting peptidoglycan transpeptidases, ampicillin inhibits cell wall synthesis and bacterial cell division. Therefore, ampicillin causes the cells to bulge, producing a cytosolic protrusion surrounded by both the cytoplasmic and outer membranes, near the cell septum, and induces expression of autolysins, subsequently leading to cell lysis. Figure 10 shows different stages of a UPEC cell exposed to ampicillin, with an initial normal appearance, an initial membrane-bound cytoplasmic bulge formed within a few minutes, followed by a large bulge and subsequent lysis.

![Fig. 10 Membrane deformation of a UPEC cell exposed to ampicillin (1 mg/ml). (a) Normal cell, (b) beginning of bulge formation, (c) large bulge, and (d) cell lysis. Scale bar: 200 nm.](image)

**Fig. 10** Membrane deformation of a UPEC cell exposed to ampicillin (1 mg/ml). (a) Normal cell, (b) beginning of bulge formation, (c) large bulge, and (d) cell lysis. Scale bar: 200 nm.

![Fig. 11 Displacement measurements of UPEC cells exposed to ampicillin (1 mg/ml). Displacement vectors (a, c, and e) and corresponding cell plots (b, d, and f) are shown for different regions of different cells. The bulge region is shown with magenta vectors and marked as “1”. The circled regions (b, d, and f) indicate the bulge membrane deformation while the arrows indicate cell lysis. Scale bar 200: nm.](image)

**Fig. 11** Displacement measurements of UPEC cells exposed to ampicillin (1 mg/ml). Displacement vectors (a, c, and e) and corresponding cell plots (b, d, and f) are shown for different regions of different cells. The bulge region is shown with magenta vectors and marked as “1”. The circled regions (b, d, and f) indicate the bulge membrane deformation while the arrows indicate cell lysis. Scale bar 200: nm.
Figure 11(a) shows the image of a UPEC cell captured 20 min after exposure to a high concentration of ampicillin (1 mg/mL). Similar to the PMB experiments, we exposed cells to an ampicillin concentration significantly higher than the determined MIC (4 μg/mL) to rapidly induce cellular responses. The local displacements associated with cell membrane deformation at different locations reveal detailed time characteristics linking the formation of a bulge to cell lysis [Fig. 11(b)]. The bulge region (magenta, and marked by “1”) gradually increases the displacement as the protrusion forms (marked by a circle in the plot). A second location (marked by “2”) next to region “1” also shows a displacement increase (cyan curve), indicating that the observed displacement increase in location “1” is not due to noise. The displacements in other locations (e.g., regions 3 to 5) of the cell display minimal changes. Although the initial bulge formation is small, it can be resolved within ~9 nm, demonstrating the advantage of the present method over other optical imaging and cell segmentation approaches.24 After the cellular bulge expands over time, cell expansion slows and is followed by a sudden decrease as cell lysis initiates (marked by thick black arrows). This characteristic cellular response was observed in other UPEC cells [Figs. 11(c)–11(f)], but the magnitude and initiation of cellular deformation vary due to heterogeneity of different cells.24

We have demonstrated a sensitive optical method to track nanometer-scale cell deformation and growth as a phenotypic feature for potential AST using two antibiotics with distinct action mechanisms on the cell membrane and cell wall, PMB and Ampicillin, as examples. Other antibiotics with different mechanisms may not have the same rapid effects on cell deformation. To further develop it into technology for AST, substantial validation experiments will be needed. Examples include testing of other common antibiotics over clinically relevant concentration ranges, both susceptible and resistant strains, and also real clinical samples.

4 Conclusion
We have developed an optical imaging method to track antibiotic activity on individual bacterial cells in real-time. The method measures bacterial cellular deformations with ~9-nm detection limit, allowing sensitive detection of antibiotic-induced changes in size, shape, and local membrane deformations of single bacterial cells. Using this capability, we have investigated the effects of PMB on E. coli O157:H7 and ampicillin on UPEC. PMB caused E. coli O157:H7 to shrink along the cell edge, whereas ampicillin caused the cellular bulge to form in UPEC. Based on tracking membrane deformation changes of single cells, our method will help identify individual cells in a sample that are susceptible to antibiotics in real-time. The current detection limit is mainly determined by the stability of the microscope and the attachment of the bacterial cells to the surface. This detection limit may be improved for more accurate tracking of antibiotic-induced bacterial membrane deformations, which may lead to a sensitive AST method. The present work focused on developing a sensitive optical tracking method to detect cell deformation associated with antibiotic activity. Further validation experiments must be performed with additional antibiotics and bacteria, including antibiotic-resistant strains, to establish MIC in conjunction with traditional AST methods.

Disclosures
The authors have no relevant financial interests in this article and no potential conflicts of interest to disclose.

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References

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