Lab on a Chip



View Article Online

PAPER



Cite this: Lab Chip, 2025, 25, 714

Received 13th September 2024, Accepted 23rd December 2024

DOI: 10.1039/d4lc00765d

rsc.li/loc

Introduction

Antimicrobial resistance represents an incessant global health threat that necessitates optimal approaches to therapy. Antimicrobial susceptibility testing (AST) is the current gold standard clinical microbiological test for predicting antimicrobial effect against the causative pathogen by determining the minimum inhibitory concentration (MIC) of antimicrobials *in vitro*. Traditional AST requires high inoculum of isolated colonies that is grown in culture with various concentrations of antimicrobials for 16–20 hours, followed by detection of cell proliferation. MIC results are then interpreted

Precision single cell analysis to characterize host dependent antimicrobial response heterogeneity in physiological medium[†]

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Antimicrobial stewardship plays an essential role in combating the global health threat posed by multidrugresistant pathogens. Phenotypic antimicrobial susceptibility testing (AST) is the gold standard for analyzing bacterial responses to antimicrobials. However, current AST techniques, which rely on end-point bulk measurements of bacterial growth under antimicrobial treatment in a broth solution, have limitations in resembling the physiological working environment and resolving heterogeneity in response kinetics within the population. In this study, we investigate the responses of uropathogenic bacteria under antimicrobial treatment in individual urine. Our results demonstrate substantial heterogeneity in time-kill kinetics in response to antimicrobials in a host-dependent manner. We also establish a microfluidic gel encapsulation platform for single cell imaging to rapidly resolve heterogeneous subpopulations in response to antimicrobials. The platform captures both bacterial growth and killing within the gel and enables medium exchange to assess the ability of surviving cells to resume growth after antimicrobial removal. Our study lays the foundation for a new generation of precision single cell analysis for personalizing antimicrobial treatment.

> against valid clinical breakpoints to determine susceptibility and predict clinical therapeutic outcome. Clinical breakpoints used in AST interpretation is based on achievable unbound antimicrobial concentration in serum instead of the target site. However, many antimicrobials achieve much higher concentrations in other body fluids. For example, through glomerular filtration and tubular secretion, antimicrobial concentrations in urine have found to be 100-1000 times that achieved in serum.¹ There is poor correlation between serum antimicrobial level and bacterial clearance from urine.² In addition, AST is traditionally performed in standardized nutrient-rich media such as Mueller-Hinton II (MH2), where bacteria grow optimally, and antimicrobial activities are often highest. These conditions do not represent the working condition of antimicrobials in the urinary system. The physiology of the host infection environment has been shown to alter growth and expression of essential genes in various pathogens and consequently influence antimicrobial susceptibility.3-5 Moreover, AST, a growth based measurement, does not resolve antimicrobial tolerance. Tolerance is the ability of a bacterial population to survive a transient exposure to antimicrobials, even at concentrations far exceeding its MIC, and has been associated with treatment failure and relapse of many bacterial infections.⁶⁻⁸ A tolerant bacterial strain can have the same MIC as a susceptible strain but requires longer exposure to an antimicrobial, rather than higher concentration,

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[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d4lc00765d

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to produce the same level of killing as a susceptible strain. Thus, traditional AST based on a static *in vitro* MIC value cannot provide insight into the kinetics of antimicrobial activity and does not mimic *in vivo* exposure dynamics at the effect site. Rapid accurate prediction of the pathogen's response to antimicrobial therapy to tailor treatment regimens that can provide optimal bacterial killing and prevent the amplification of resistance remains a critical gap in clinical care.

Time-kill curves measure the bacterial response to antimicrobials over time, providing detailed information on both growth and killing kinetics, and offering more meaningful pharmacodynamic insights compared to static MIC values.9 Importantly, intercellular heterogeneity in antimicrobial response phenotypes can exist within a bacterial population. Despite a lack of consensus definitions, tolerance refers to a phenotypically, but non-genetically heritable, subpopulation of surviving cells, typically comprising less than 1% of the population. This subpopulation, which includes persister cells and viable but non-culturable (VBNC) cells, has been suggested to serve as evolutionary steppingstones towards resistance due to their increased mutation rate.¹⁰⁻¹⁴ Persister cells are phenotypic variants that survive antimicrobial exposure without replicating but resume growth after the antimicrobial is removed.15 In contrast, VBNC cells do not replicate postantibiotic removal.¹⁶ The presence of these tolerant variants, implicated to cause treatment failure from relapse and emergence of antimicrobial resistance,¹⁷⁻²⁰ can be characterized by a bimodal (or multimodal) time-kill curve that deviates from the simple decay expected from a uniform population.

In this study, we explored precision single cell analysis to capture both growth and killing kinetics in physiological medium, aiming to optimize antimicrobial selection and treatment duration. Time-kill curves offer a valuable tool for characterizing antimicrobial kinetics, distinguishing tolerant or variant subpopulations from susceptible phenotypes. We assessed their utility in evaluating the kinetics of common pathogens causing complicated urinary tract infections (UTIs) in response to frontline intravenous (IV) antimicrobials. Given the substantial differences in physiological conditions between urine and the standard MH2 broth typically used, we conducted our analyses in individual urine samples at physiologically relevant antimicrobial concentrations. Our aim was to examine inter-individual heterogeneity, dependent on the drugpathogen-host combination, in both susceptible and resistant strain responses. Furthermore, to streamline the labor-intensive plating process involved in bulk time-kill analysis, we developed a microfluidic gel encapsulation platform for single cell microscopy, enabling the rapid resolution of intercellular response heterogeneity within a microbial population. Leveraging recent advancements in single cell imaging and microfluidic techniques for bacterial analysis,²¹⁻²⁹ this platform offers a cost-effective solution for analyzing growth and killing kinetics at the single cell level. Additionally, the micrometerthick gel layer is designed to facilitate rapid medium exchange, enabling the assessment of regrowth following antimicrobial removal to identify persistent subpopulations.

Materials and methods

Bacterial strains and healthy volunteer human urines for bulk time-kill curves

To investigate the bacterial responses to antimicrobials in human urines, Escherichia coli, Klebsiella pneumoniae, and Enterococcus faecalis were used as representative UTI causing species.³⁰ K. pneumoniae strains KP10 (AR 0010), KP12 (AR_0012), KP16 (AR_0016), KP120 (AR_0120), and KP153 (AR 0153) were obtained from The CDC & FDA Antimicrobial Resistance Isolate Bank. E. coli strains EC10789 (ATCC10789) and EC1427 (ATCC BAA-1427) and E. faecalis strains EF19432 (ATCC19432) and EF47077 (ATCC47077) were purchased from American Type Culture Collective (ATCC). MICs of antimicrobials to each strain and antimicrobial combination were measured by the broth microdilution method based on the CLSI guideline.³¹ Midstream urine was collected from healthy volunteers, who had not taken any antibiotics in the prior two weeks, after informed consent as per the IRB approved protocol (70759). Collected urine was then aliquoted and stored at -80 °C until further testing. Growth in MH2 broth was used as a control. Susceptible strains were tested in 2 healthy volunteers (urine 1 and urine 2) and resistant strains were tested in 3 healthy volunteers. Urine from individuals 1 and 2 were collected at two independent dates (urines 1 and 4, and 2 and 5). All healthy volunteer urines were confirmed to be sterile by culturing 50 µL of each urine on MH2 agar plate overnight at 37 °C.

Bulk time-kill curves to assess bacterial response kinetics to antimicrobials in urine

Overnight cultures of K. pneumoniae and E. coli strains on MH2 agar, and E. faecalis strains on brain heart infusion (BHI) agar were used for estimating the time-kill kinetics. Colonies were suspended in phosphate-buffered saline (PBS) to achieve a turbidity of 0.5 McFarland standard and further diluted 10-fold in PBS to result in 1.5×10^7 CFU mL⁻¹. Meropenem, ciprofloxacin, gentamicin, and ampicillin were each dissolved in PBS to concentrations of 320, 120, 220, and 5120 µg mL⁻¹, respectively. 10 µL of bacterial suspension was inoculated in 80 µL of urine or MH2 broth and 10 µL of antimicrobial solution to reach the physiologically minimum concentrations of the antimicrobials in human urine under treatment. After 2, 4, 6, 8, and 24 hours of culturing at 37 °C, cells were taken, diluted in PBS buffer, and spotted on MH2 agar (K. pneumoniae and E. coli) or BHI agar (E. faecalis), and incubated at 37 °C overnight to determine the number of viable cells. The minimum urinary concentrations of meropenem,³² ciprofloxacin,³³ and gentamicin³⁴ in patients under antibiotic treatment for UTI were estimated based on previous studies. The concentration of ampicillin was between the thresholds of urinary concentration 3 to 4 h after 500 mg intramuscular injection, which is much higher than the lowest urinary concentration during treatment.35

Microfluidic gel encapsulation

A microfluidic gel encapsulation technique was developed to embed bacteria in thin layers of micropatterned gel pads for evaluating individual bacterial responses to antimicrobials. We used a VLS 3.5 Desktop Series laser machining system to make an array of small wells with a diameter of 0.5 mm on a 5 µm ultra-thin double-sided tape (Dwell GM4095T, PETbase). We then attached the micropatterned tape to glass coverslips as a substrate for single cell microscopy. To implement gel encapsulation, 3% low-melting-point agarose (Fisher BioReagents, BP165-25) was dissolved in 1× PBS and heated to 65 °C to ensure complete dissolution. The gel was then carefully cooled to 37 °C or below before mixing with bacteria to prevent thermal damage to the cells. This mixture was then rapidly applied to the micropatterns. A microscope glass slide was gently used to scrape across the surface of the template to create thin micropatterned gel pads. Following this, we attached a laser-machined acrylic plate with a diameter of 1 cm to the coverslip to create a multiwell platform that is compatible with standard imaging setups and liquid handling equipment. Alternatively, thin layers of gel with embedded bacteria can be produced by centrifugation using a PDMS mold on a multiwell array (Fig. S1[†]). This method is particularly suitable for experiments requiring a large number of samples and diverse experimental conditions.

Clinical samples

The microfluidic gel encapsulation platform was tested using clinical isolates from Penn State Milton S. Hershey Medical Center. These isolates, which included *E. coli, Klebsiella oxytoca, K. pneumoniae*, and *Pseudomonas aeruginosa*, were extracted from urine samples from patients with UTI or positive blood tube samples. All procedures were approved by the Institutional Review Boards of Penn State (STUDY00003415). The clinical isolates were subjected to varying concentrations of ciprofloxacin.

We examined a total of 16 clinical isolates. Initially, we used a centrifuge operating at 5000 rpm for five minutes to effectively isolate the bacterial cultures from the samples. After isolation, the bacteria were pre-cultured in MH2 broth for one hour. Bacterial samples were then prepared to a concentration of $\sim 1.5 \times 10^8$ CFU ml⁻¹ and quantified using a NanoDrop spectrophotometer. Antibiotics were diluted to the desired concentrations using MH2 broth. For the 96-well plate experiments, 100 µl of the appropriate growth medium was added to each well. The antibiotic solutions were dispensed into the corresponding wells, typically using a twofold serial dilution to achieve a gradient of antibiotic concentrations. We inoculated each well with 10 µl of the standardized microbial suspension, ensuring that the final volume in each well was 200 µl. To prevent evaporation and contamination, we sealed the plates with gas permeable parafilm. The plates were incubated at 37 °C for 16-24 hours to allow for bacterial growth. After incubation, we examined

the wells for microbial growth; the presence of a visible pellet or turbidity indicated growth. The MIC was determined as the lowest concentration of antibiotics that completely inhibited the growth of the organism. When using the NanoDrop spectrophotometer for quantification, we followed the same protocol as that used for the 96-well experiments. To establish a baseline, we recorded the initial measurements at 0 hours and compared them with those at 2 hours to determine the fold change. Significant changes in optical density were indicative of the response to each antibiotic concentration.

Single cell imaging and data analysis

The bacteria were pre-cultured in MHB broth for one hour prior to the experiment to ensure they were harvested during the exponential (log) growth phase, providing actively dividing cells for subsequent microfluidic experimentation. The microfluidic gel encapsulation platform was mounted on a Leica DMi8 epifluorescence microscope, equipped with an Okolab UNO-T-H temperature chamber to maintain a constant temperature at 37 °C for optimal bacterial growth. Both bright-field and fluorescence microscopy were performed using a Leica HCX PLAN APO $40\times/0.85$ objective lens. Bright-field images were captured with an exposure time of 15 ms. Fluorescence images for PI staining were captured using a Leica Y3 filter set, with an exposure time of 0.5 s.

Automated image analysis was implemented using an ImageJ Macro to analyze the images, which identified the position of bacteria and captured morphological features, including area and perimeter. For precise calculations, images were converted to a 32-bit format, and background noise was subtracted using a rolling ball algorithm with a 10-pixel radius, assuming a lighter background. Image segmentation was performed using the particle analysis function in ImageJ by setting the threshold using the default method, and particles were analyzed by setting a minimum particle size of 0.02 pixels to exclude background noise. From this analysis, we extracted features for each bacterium, including area, mean intensity, XY coordinates, perimeter, aspect ratio, and solidity. We also analyzed the ratio of bacteria whose perimeter exceeds specific thresholds to compare the fold change of features relative to the initial time point. Additionally, individual cells were tracked using their XY coordinates to analyze single-cell growth and killing kinetics. The position of a cell between two time points was calculated using the Euclidean distance formula:

$$d = \sqrt{(x_2 - x_1)^2 + (y_2 - y_1)^2}$$

where:

- *d* is the distance between the two points,

- (x_1, y_1) are the coordinates of the cell at the initial time point,

- (x_2, y_2) are the coordinates of the cell at the subsequent time point.

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This method enabled accurate tracking of individual cells across different time points. Regions of interest (ROIs) were generated from bright-field images to enhance the analysis of fluorescence images and extract critical viability information indicated by the dye.

Statistical analysis

All assays were performed in triplicate unless specified otherwise. Data are presented as mean \pm standard deviation. For our single cell growth analysis, we acquired three images per measurement, each capturing approximately 3000 bacteria.

Results

Killing kinetics of susceptible strains depends on growth media

We first performed bulk time-kill analyses of bacteria. We compared the time-kill curves for carbapenem and/or ciprofloxacin susceptible *K. pneumoniae* and *E. coli* strains grown in MH2 broth and healthy volunteers' urines with

physiologically minimum concentrations of meropenem (32 $\mu g mL^{-1}$) and ciprofloxacin (12 $\mu g mL^{-1}$) (Fig. 1), a concentration that is more than 64× the MIC for the strains tested. For meropenem, most bacteria were completely killed within 2-4 hours of treatment, except for KP10. This strain exhibited a tolerant subpopulation (<1%) that could survive even after 8 hours of treatment, while the majority of the population was killed within 2 hours of treatment. In urine 2, the surviving population after 8 hours of treatment was larger than in MH2 or urine 1. When prolonging the meropenem treatment of KP10 to 20 hours, similar to the standard AST, no survivors were observed in replicates with MH2 (0/14)(Table S1[†]). In contrast, 3 out of 28 (10.7%) replicates in urine samples treated with meropenem and 2 out of 24 (8.3%) replicates in urine samples treated with ciprofloxacin showed colony formation after treatment. These results suggest that the working media, particularly urine in a hostdependent manner, can contribute to heterogeneity in bacterial susceptibility to antimicrobials.





Fig. 1 Time-kill curves of susceptible *E. coli* and *K. pneumoniae* strains in MH2 and healthy volunteer urines supplemented with meropenem or ciprofloxacin. Slower clearance rates in urines compared to MH2 for EC10789, KP10, and KP16 for both antimicrobials ($32 \mu g m L^{-1}$). Solid lines represent time-kill curves for ciprofloxacin and dashed lines represent those for meropenem. KP12 and KP13883, which were resistant to ciprofloxacin, were tested with meropenem only. Smoothed curves across three replicates at each time point depict the time-kill curves along with their 95% confidence intervals. The starting points of the curves are jittered to represent curves with similar shapes of rapid killing. 1.5×10^6 CFU mL⁻¹ of bacteria were incubated with the antimicrobial for 2, 4, 6, 8, and 24 h in each healthy volunteer urine or MH2 at 37 °C, following which cell survival was determined by plate counting at each time point.

Heterogeneities in susceptibility were also observed in other antimicrobials in these strains. When treated with ciprofloxacin, the time-kill curves revealed bacterial responses distinct from those observed with meropenem. KP10, which exhibited a tolerant subpopulation to meropenem, was killed sharply by ciprofloxacin under all tested conditions. In contrast, KP16 and EC10789, which were killed sharply by meropenem, exhibited surviving subpopulations against ciprofloxacin after 6–8 hours treatment. Notably, urine 2 exhibited a substantially larger surviving subpopulation compared to urine 1 and MH2. In contrast, ECBAA1427 was completely killed by both agents within 4 hours.

We further performed time-kill analysis of ampicillin- and gentamicin-susceptible strains of E. faecalis, EF19432 and EF47077, against these antimicrobials in healthy urine. Interestingly, both strains with MICs of 1 μ g mL⁻¹ ampicillin showed incomplete killing after 24 hours of treatment with 512 μ g mL⁻¹ of ampicillin (Fig. 2). In contrast, both strains had MICs of $<0.25 \ \mu g \ mL^{-1}$ for gentamicin and were killed when treated with 22 $\mu g m L^{-1}$ of the antimicrobial. There was distinguishable heterogeneity in the killing kinetics of EF47077 in the two urines and MH2, with fastest killing, under 2 hours, in urine 2 and slowest in MH2. Subjecting the two strains to both ampicillin and gentamicin had no additive effect on EF19432 but hastened the killing of EF47077 to under 2 hours, highlighting the value of time-kill analysis in resolving the combinatorial effects of the antibiotics on the bacteria. These results further support the notion that the working media can influence the heterogeneity in bacterial susceptibility to antimicrobial.

Killing kinetics of resistance strains depends on growth media

Heterogeneity was also evident in the resistant strains, KP120 and KP153 with known MICs of 32 µg mL⁻¹ for meropenem (Fig. 3). As expected, both strains grew in all urines and MH2 without antimicrobial. However, resistance to meropenem was delayed in urines 1 and 2. The majority (99%) of KP120 was killed within 4 hours in urine 3, but growth resumed after 8 hours. Remarkably, at the strain's MIC of 32 μ g mL⁻¹, KP120 was completely cleared within 4 hours in urines 4 and 5, and KP153 was completely cleared in urines 1, 2, and 3, suggesting that the antimicrobial would be effective against this resistant strain in these individuals on those specific dates. Notably, the strain grew normally in urines 4 and 5 in the absence of the antimicrobial, suggesting the presence of unknown urine factor(s) that may synergize with the potency of meropenem. Such intraindividual and interindividual heterogeneity, and discrepancy in responses between MH2 and urine, underscore the complexity of host-pathogen-antimicrobial interactions and warrant precision measurement techniques for guiding antimicrobial treatment.

Microfluidic gel encapsulation platform

Bulk time-kill analysis relies on plate counting to quantify colony-forming units, but the process is cumbersome and slow due to the colony-forming step. Moreover, bulk time-kill analysis is unable to detect VBNC subpopulations and distinguish between persisters and VBNC. To overcome these limitations, we devised a microfluidic gel encapsulation



Fig. 2 Time-kill curves of susceptible *E. faecalis* against ampicillin, gentamicin, and their combination. The bacteria were not completely eradicated by ampicillin after 24 hours but were completely killed by gentamicin. Time-kill curves provided a quantitative measure of the accelerated killing effect of the combination of ampicillin and gentamicin. Smoothed curves across three replicates at each time point depict the time-kill curves along with their 95% confidence intervals. The starting points of the curves are jittered to represent curves with similar shapes of rapid killing. 1.5×10^6 CFU mL⁻¹ of bacteria were incubated with the antimicrobial for 2, 4, 6, 8, and 24 h in each healthy volunteer urine or MH2 at 37 °C, following which cell survival was determined by plate counting at each time point.



Fig. 3 Time-kill curves of resistant strains KP120 and KP153 treated with meropenem. KP120 was completely killed within 4 hours in two urines only with meropenem at 32 μ g mL⁻¹ (MIC), showed delayed resistance in two others, and had >99% killing in 4 hours in one sample, though the population recovered after 8 hours. KP153 was completely killed with meropenem in three urine samples. Smoothed curves across three replicates at each time point depict the time-kill curves along with their 95% confidence intervals. 1.5×10^6 CFU mL⁻¹ of bacteria were incubated with the antimicrobial for 2, 4, 6, 8, and 24 h in each healthy volunteer urine or MH2 at 37 °C, following which cell survival was determined by plate counting at each time point.

platform for single cell growth and killing analysis (Fig. 4A). The design concept focuses on trapping bacteria in a thin layer of low-temperature agarose gel, which enables single cell timelapse microscopy. The thin gel structure is created by using a glass slide to spread and scrape the gel into laser-machined micropatterns, forming isolated thin gel pads that facilitate image analysis.

The fabrication process begins with laser machining ultrathin tape (5 μ m) to create a micropatterned array consisting of circles with a diameter of 0.5 mm on a coverslip (Fig. 4B). The minimal thickness of the tape ensures that all bacteria are on the same focal plane for imaging. Additionally, the thin gel layer allows for easy reagent loading, washing, and medium exchange. Fig. 4C illustrates the process of spreading the gel with a glass slide. After gel loading, a laser-machined acrylic plate is attached to the coverslips to form a multiwell array for parallel processing. Each well has dimensions of 3 mm in height and 10 mm in diameter. The acrylic well can hold up to 235 μ L of culture medium, while the volume of a single gel cylinder in the micropatterned well is 0.98 nL. Fig. 4D shows the assembled multiwell platform, which is designed for measuring single cell growth and killing kinetics. To accommodate a larger number of samples, we also developed an alternative method to create thin gel layers with embedded bacteria using centrifugation (Fig. S1[†]).

Microfluidic gel encapsulation platform for single cell AST

We first evaluated the microfluidic gel encapsulation platform for measuring bacterial growth. We tested 16 clinical isolates obtained from the Clinical Microbiology Laboratory (Fig. 5A). These clinical isolates were simultaneously tested against different concentrations of ciprofloxacin, taking advantage of the multiplexity of the microwell design. The cells were imaged using brightfield microscopy and analyzed with ImageJ to identify bacteria and automatically extract growth-dependent features, such as area or perimeter. In this experiment, the increase in perimeter after one hour was normalized to the initial value to quantify bacterial growth. For comparison, we also conducted the experiment using a 96-well plate format with both two-hour and overnight culture, resembling standard bulk AST. The growth of bacteria after two-hour culture was detected using a NanoDrop spectrophotometer (Fig. 5B and C). Among the clinical isolates tested, the observed trends showed consistency across the gel encapsulation platform, 96-well plate with NanoDrop, and overnight culture approaches. The MIC values were in general agreement within one to two-fold dilutions of each other, supporting the use of the microfluidic gel encapsulation platform for rapid single cell analysis.

Microfluidic gel encapsulation platform for precision single cell analysis

The microfluidic gel encapsulation platform can resolve the growth kinetics of individual cells. We analyzed the response of a uropathogenic *E. coli* (EC137) exposed to ciprofloxacin. The strain is susceptible to ciprofloxacin with an MIC of 2 μ g mL⁻¹ (Fig. 6A). In this experiment, the clinical isolate was cultured in MHB and harvested at the logarithmic phase. A bacterial sample of ~1 × 10⁸ CFU ml⁻¹ (measured using a NanoDrop) was prepared by centrifuging the culture and then resuspending it in media. As shown in Fig. 6B, notable bacterial growth was



Fig. 4 Microfluidic gel encapsulation platform for time-kill analysis of single cells. (A) Schematic diagram of the gel-based single cell analysis system. Agarose embedded with bacteria is scraped into thin gel layers within micropatterns laser-machined onto ultra-thin tapes. (B) Micropatterns were formed on glass coverslips to facilitate single cell imaging. (C) A glass slide is used to spread the agarose gel on the micropatterns, resulting in thin agarose gel layers. (D) After spreading, a laser-machined acrylic plate is bonded to the chip to create the multiwell platform with micropatterned agarose gel.

observed in the absence of ciprofloxacin. At 2 μ g mL⁻¹ (1× MIC), most bacteria exhibited filamentous morphology and ceased replication. Bacterial filamentation is a marker of cellular stress and is indicative of an induced SOS response to DNA damage or interrupted DNA replication.^{36,37} At concentrations of 4 and 8 mg L⁻¹, the growth of bacteria were largely inhibited. Some cells were stained with propidium iodide (PI), despite ciprofloxacin not directly affecting the bacterial cell membrane integrity.³⁸ Fig. 6C depicts the fold increase in the perimeter and area of single cells across different concentrations of ciprofloxacin, underscoring the intercellular heterogeneity in response to the antimicrobial. Similar results were also obtained by *K. pneumoniae* (Fig. S2†).

In addition to analyzing a fixed time point, similar to traditional AST, the microfluidic gel encapsulation platform also allows time-lapse microscopy to study the growth and killing kinetics of single cells (Fig. 7A and S3–S5†). To demonstrate this capability, we expose bacteria to varying concentrations of ciprofloxacin while tracking them within the same imaging field at four different time points. In the absence of ciprofloxacin, substantial cell growth with large increases in both perimeter and area was observed (Fig. 7B and C). At 2 μ g mL⁻¹ (1× MIC), a small percentage of cells duplicated within 60 minutes; however, beyond this point, only slight elongation without replication was observed. At higher concentrations (20 μ g mL⁻¹), bacterial growth was effectively inhibited,

with minimal changes in morphology. These results demonstrate that the microfluidic gel encapsulation platform can reveal concentration-dependent kinetics of bacterial responses to antibiotics at the single cell level.

Under optimal growth conditions (i.e., rich media at 37 °C), the bacteria exhibited a maximum doubling time of ~30 minutes during the exponential phase. In the microfluidic gel encapsulation platform, we observed a slight delay in the growth rate at the beginning of the experiment. This initial lag phase lasted approximately one duplication cycle (~30 minutes) and may have resulted from the transition to the new solidphase environment and potential thermal or physical stress introduced during the gel loading procedure. However, following the lag phase, the bacterial growth rate returned to the normal rate of approximately 30 minutes per doubling. This observation supports that the thin micropatterned gel pads in the gel encapsulation platform do not impose inherent limitations (e.g., nutrient diffusion or oxygen availability) on bacterial growth and are well-suited for analyzing bacterial responses to antimicrobial treatment.

Microfluidic gel encapsulation platform for fluid exchange

To assess the ability of surviving bacteria to resume growth after the removal of antibiotics, a fluid exchange step is necessary. Given that the gel encapsulating the bacteria is



Fig. 5 AST of clinical isolates using the microfluidic gel encapsulation platform. (A) Growth of *E. coli, K. oxytoca, K. pneumoniae,* and *P. aeruginosa* clinical isolates measured by the microfluidic gel encapsulation platform. Bacteria were exposed to different concentrations of the antibiotic ciprofloxacin. Bacterial growth was quantified by normalizing the perimeter increase after one hour to the initial value. (B and C) Bacterial inoculum was 7.5×10^6 CFU mL⁻¹, and the growth of clinical isolates was measured by optical density in 96-well plates after 2 hours and overnight culture.

only 5 µm thick, fluid exchange can be easily performed on the microfluidic gel encapsulation platform. To demonstrate this capability, we conducted a single cell analysis experiment using *K. pneumoniae* (ATCC 11296) exposed to meropenem at the MIC of the strain (0.03 µg mL⁻¹) for one hour. After antibiotic treatment, the broth containing meropenem was replaced with fresh medium, and viable cells were monitored during an additional hour of recovery incubation period (Fig. 8A). At the beginning of the experiment, PI was introduced to detect dead cells. As expected, the majority of bacteria were killed by meropenem at the MIC and stained by PI. In the analyzed field of view, we observed 95 bacteria, out of which 82 were stained by PI, indicating cell death. Conversely, 13 bacteria were not stained by PI and showed no signs of growth or elongation. These cells appeared to tolerate meropenem during the duration of antimicrobial treatment (dotted circles in Fig. 8B).

To investigate if these viable bacteria could resume growth after the removal of the antimicrobial, a fluid exchange step was performed. Utilizing time-lapse microscopy and automated image analysis, we monitored the growth curve representing changes in the area of individual bacteria (Fig. 8C). Among the cells that were not stained by PI, three bacteria resumed growth within one hour of antimicrobial removal, consistent with the behavior of persister cells, while the remaining 10 cells could represent either VBNC cells or

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Gel-based AST 90 Min



Fig. 6 Single cell response of uropathogenic *E. coli* to different concentrations of ciprofloxacin. (A) Growth measurements obtained using the microfluidic gel encapsulation platform under the same cultivate condition as Fig. 5. The bacteria were incubated in varying concentrations of ciprofloxacin dissolved in MHB at 37 °C. (B) Brightfield and fluorescence images illustrating the response of *E. coli* (EC137) exposed to ciprofloxacin. (C) Changes in the perimeter and area of individual bacteria after 90 minutes. Red dots indicated dead cells, as determined by PI staining. Values were normalized to the initial measurement.

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Fig. 7 Single cell response kinetics to different concentrations of ciprofloxacin. (A) Normalized changes in the perimeter of individual bacteria at 60, 90, and 120 min. The bacteria were inoculated at 0, 2, 20, and 128 mg L^{-1} of ciprofloxacin dissolved in MHB at 37 °C. Values were normalized to the initial measurement. Red dots indicate dead cells as shown by PI staining. Values were normalized to the initial measurement. (B and C) Growth kinetics of bacterial perimeter and area. Values were normalized to the initial measurement.



Fig. 8 Medium exchange in the microfluidic gel encapsulation platform for resolving the ability of bacteria to replicate after antibiotic removal. (A) Schematic timeline of the experiment. *K. pneumoniae* embedded in gel was first treated with meropenem for one hour, after which the medium was replaced to remove the antimicrobial. (B) Fluorescence and brightfield images of *K. pneumoniae* with PI staining, illustrating bacterial death. Dotted circles indicate viable bacteria without PI staining, while green circles indicate bacteria that resumed growth after meropenem removal. (C) Analysis of individual cell growth kinetics. Three cells resumed growth within one hour after medium exchange. Values were normalized to the initial measurement.

persister cells requiring longer recovery times. These results highlight the capability of the microfluidic gel encapsulation platform for single cell time-kill curve analysis and medium exchange, enabling the distinction of different cell fates. Future studies analyzing a larger number of bacteria and a broader range of conditions will be valuable for deciphering the differences between VBNC cells, persisters, and other dormant cells.

Microfluidic gel encapsulation platform for resolving bacterial response to antimicrobial in urine samples

We performed single cell analyses of uropathogenic E. coli (EC 137), a ciprofloxacin-susceptible strain, at 37 °C in MHB and in urine samples from three healthy volunteers (Fig. 9 and S7[†]). Using the MIC of ciprofloxacin (2 μ g mL⁻¹) as a reference, we compared the response of bacteria grown to log phase to the antibiotic across these different culture conditions. In MHB, the bacteria exhibited filamentous elongation, a stress response to antibiotic exposure, with no replication observed after 90 minutes. Several cells stained positive for PI, indicating compromised viability (Fig. 9A). In contrast, in all three urine samples, a substantial portion of cells survived and continued to replicate, with fewer PIstained cells compared to MHB. The percentage of surviving cells varied among the urine samples, potentially reflecting individual differences in urine composition. This observation was generally consistent with the bulk analysis (see Fig. 1). Fig. 9B illustrates the fold increase in the area of individual cells across different media, highlighting intercellular variability in antibiotic responses depending on the medium. These results further support that the choice of medium, particularly host urine, can influence bacterial susceptibility

and contribute to heterogeneity in antibiotic responses. Overall, the microfluidic gel encapsulation platform offers a rapid technique for capturing host-dependent variability in response to antibiotics at the single cell level.

Discussion

In this study, we assessed the variability in bacterial killing kinetics at physiologically relevant concentrations of antibiotics in MH2 and urine and developed a fluid-exchange-enabled microfluidic device to track these response kinetics at the single cell level. While previous studies have compared the MIC of bacteria in MH2 and pooled physiological fluid or adapted media,^{39–43} none have systematically investigated the interindividual heterogeneity in the killing kinetics. Our results highlight the host-dependent heterogeneity of bacterial response to antimicrobials. It is often assumed that if a bacterium can be killed by antibiotics in a broth culture medium, which is favorable for bacterial growth, it can be killed even more effectively by antibiotics in body fluids, which are less ideal for their growth. In contrast, we found that some strains could survive under antibiotic pressure (32× MIC) in urine better than in broth, consistent with previous findings.³⁹⁻⁴³ Nutrient limitation, immunological stress, or variations in pH might increase the persistence of certain subpopulations under antibiotic pressure or reduce the efficacy of the antibiotic, contributing to the observed effect in urine.44-47 While some susceptible strains were completely killed by the antibiotics within 2 hours, others included subpopulations that survived for a longer time. On the contrary, both resistant strains tested were completely eradicated by the antibiotics in certain urines but not in MH2, corroborating previous findings of bacterial resistance reversal to antibiotics

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Fig. 9 Single cell response of *E. coli* to ciprofloxacin across different growth media. (A) Merged bright-field and fluorescent images illustrating the response of uropathogenic *E. coli* (EC137) exposed to 2 mg mL⁻¹ ciprofloxacin at 37 °C in different growth media (MHB and urine samples from three healthy individuals) at 0 minutes and 90 minutes. Red indicates dead cells, green circles indicate duplicated cells, and red arrows highlight filamentation of cells in urine sample 3. (B) Changes in the area of individual bacteria after 90 minutes. Red dots represent dead cells, as determined by PI staining. The area is measured in μ m² while the area change is dimensionless.

under physiological conditions.^{39,43} Thus, bacterial responses to the antibiotics varied based on strains, antibiotics, and the individual urine samples. Characterizing this variability in bacterial responses to antibiotics may inform appropriate treatment choice and duration for effective eradication of infection.

In addition to resolving interindividual heterogeneity in physiological media, the ability to resolve intercellular timekilling heterogeneity in antimicrobial responses is critical for identifying minority subpopulations of response variants, which may have significant clinical implications. As proof of concept, we demonstrate a microfluidic gel encapsulation platform for analyzing single cell bacterial growth and killing kinetics in response to antimicrobials. The platform enables time-lapse, single cell imaging, resolving heterogeneity within the sample. Compared to the traditional time-kill curve assay with colony counting, the platform eliminates laborious procedures and time-consuming steps, which are critical for future clinical translation. The imaging platform is also compatible with advanced image analysis workflows based on computer vision and supervised learning algorithms. For instance, we have succeeded in predicting the susceptibility and MIC of antibiotic-treated bacteria within an hour by using machine learning analysis of morphological changes in bacterial single cells under the microscope.²¹

The microfluidic gel encapsulation platform offers several advantages over other microfluidic approaches, such as the microfluidic mother machine, which has been used to study the killing kinetics of bacteria against various antibiotics.24,25 Importantly, the platform enables single cell analysis in the context of intercellular interactions and quorum sensing, which are critical factors in the regulation of tolerance and resistance.48-50 Its imaging capabilities can incorporate viability indicators to assess both bacterial growth and killing. Furthermore, the platform supports medium exchange to evaluate the ability of surviving bacteria to resume growth after antibiotic removal, enabling differentiation between persisters and VBNCs. Unlike other microfluidic systems, the gel encapsulation platform does not require fluid connections or pumps and is compatible with standard liquid handling and imaging equipment, making it suitable for automation. Additionally, it involves simple procedures and can be multiplexed to test multiple conditions and samples in parallel, offering a cost-effective solution with potential for future clinical translation.

Time-kill analysis provide valuable information for the management of bacterial infections. For instance, the time-kill curve can suggest the most effective antibiotics (*i.e.*, those that achieve fast killing) for the patient among the multiple agents recommended as effective by AST. *K. pneumoniae* and

E. coli strains susceptible to meropenem, ciprofloxacin, and gentamicin used in this study showed substantial variability in the time-kill kinetics, which depends on bacterial strains, host urines, and antibiotic agents. As demonstrated in our results, time-kill curve can also offer quantitative measures of the synergistic effect between different antimicrobial combinations, enabling clinicians to implement antibiotic combination therapy with rapid killing effects that cannot be informed by MIC alone. Currently, combination therapy is chosen based on disease severity, bacterial species, or the prevalence of antimicrobial resistance in the community to avoid the development of resistance and/or subsequent failure with monotherapy.⁵¹

Precision time-kill analysis may also enhance antimicrobial stewardship by optimizing treatment duration. Currently, the duration of antimicrobial therapy is not personalized in clinical practice. Beyond reducing the emergence of antimicrobial resistance, shorter treatment course can reduce adverse effects and cost, improve patient compliance, and minimize collateral damages from gut microbiota dysbiosis. Antimicrobial stewardship has promoted a "less is better" approach to antimicrobial use in hospitals, including daily review of the continued need for intravenous antimicrobials to ensure the shortest effective treatment duration. However, in primary care or outpatient settings, where 90% of fixed-duration oral antimicrobial prescriptions are written for common infections such as uncomplicated UTI, no such ongoing assessment is routinely performed. Treatment durations for common uncomplicated UTI are determined based on comparative studies of treatment success rates over different fixed periods.52-56 While multiple studies report single-dose therapies having similar efficacy to longer-term therapies for uncomplicated UTI, these findings have not been adopted into clinical guidelines.⁵⁷⁻⁶¹ Guiding antimicrobial treatment based on precision time-kill analysis may open new opportunities to improve antimicrobial stewardship.

Conclusions

We have demonstrated that time-kill curve testing of uropathogens based on single cell analysis performed on individual urine samples can provide additional pharmacodynamic insights beyond conventional static MIC, including better prediction of physiological antimicrobial response in vitro, resolution of interindividual and intercellular response phenotypes, and guidance for single or combinatorial therapy based on effective killing kinetics. With further studies to correlate with clinical outcomes, our precision single cell analysis may offer the opportunity to individualize antimicrobial selection and treatment duration.

Data availability

All data generated are included in the manuscript and ESI.† Images generated in this study are available at: https://doi. org/10.5281/zenodo.14171622.

Author contributions

Conceptualization: S. Y., P. K. W., R. A., J.-H. L., S. M. C., N. R.-M., which was refined by input from J. C. L. and K. E. M. Methodology: S. Y., P. K. W., R. A., J.-H. L., S. M. C., N. R.-M. Investigation: R. A., J.-H. L., S. M. C. Data analysis: J.-H. L., S. M. C., N. R.-M. Funding acquisition: S. Y., P. K. W., J. C. L. Project administration: S. Y., P. K. W., J. C. L. Supervision: S. Y., P. K. W., J. C. L. Writing – original draft: P. K. W., S. Y., R. A., N. R.-M. All authors discussed the results and contributed to reviewing and editing the manuscript.

Conflicts of interest

None to declare.

Acknowledgements

This work was partly supported by NIH R01AI153133, R01AI137272, and R21GM147838.

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