Evolution of programmed cell death in bacteria is a poorly understood phenomenon in biology. A critical limitation is the lack of high-throughput technology to examine dynamics of small bacterial populations. To address this limitation, we proposed to develop a droplet-based microfluidic technology to generate population ‘bottleneck’.

This platform will serve as a critical foundation for our long-term goal to develop a quantitative understanding of microfluidics, systems biology.

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**Abstract**

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**Subject Terms**

Microfluidics, systems biology

**Security Classification**

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**Number of Pages**

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**Supplementary Notes**

The views, opinions and/or findings contained in this report are those of the author(s) and should not contrived as an official Department of the Army position, policy or decision, unless so designated by other documentation.
ABSTRACT

Evolution of programmed cell death in bacteria is a poorly understood phenomenon in biology. A critical limitation is the lack of high-throughput technology to examine dynamics of small bacterial populations. To address this limitation, we proposed to develop a droplet-based microfluidic technology to generate population ‘bottleneck’. This platform will serve as a critical foundation for our long-term goal to develop a quantitative understanding of PAD evolution by using engineered bacteria. For the proposed pilot study, we will focus on the technology development. In particular, our proposed work consists of following specific tasks, focusing on proof-of-concept demonstration of this device and its use for analyzing engineered bacteria

Enter List of papers submitted or published that acknowledge ARO support from the start of the project to the date of this printing. List the papers, including journal references, in the following categories:

(a) Papers published in peer-reviewed journals (N/A for none)

11/03/2015 1.00 Jaydeep K. Srimani, Shuqiang Huang, Anna J. Lee, Ying Zhang, Allison J. Lopatkin, Kam W. Leong, Lingchong You. Dynamic control and quantification of bacterial population dynamics in droplets, Biomaterials, (08 2015): 0. doi: 10.1016/j.biomaterials.2015.05.038

TOTAL: 1

(b) Papers published in non-peer-reviewed journals (N/A for none)

Number of Papers published in non peer-reviewed journals:

(c) Presentations
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Patents Awarded

Awards

Graduate Students

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Names of Post Doctorates

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### Student Metrics

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- The number of undergraduates funded by your agreement who graduated during this period and will pursue a graduate or Ph.D. degree in science, mathematics, engineering, or technology fields: ...... 0.00
- Number of graduating undergraduates who achieved a 3.5 GPA to 4.0 (4.0 max scale): ...... 0.00
- Number of graduating undergraduates funded by a DoD funded Center of Excellence grant for Education, Research and Engineering: ...... 0.00
- The number of undergraduates funded by your agreement who graduated during this period and intend to work for the Department of Defense: ...... 0.00
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Sub Contractors (DD882)
Inventions (DD882)

Scientific Progress

Technology Transfer

See attachment

Not applicable
Development of a microfluidic platform to analyze evolution of programmed bacterial death

PI. Lingchong You, PhD
Duke University

Statement of the problem
Our long-term goal is to address the challenges mentioned above in two aspects. On one hand, we plan to use engineered gene circuits as well-defined model systems to develop a quantitative understanding of programmed altruistic death (PAD). On the other, we plan to develop a droplet-based microfluidic technology to generate population ‘bottleneck’. This platform will serve as a critical foundation for our long-term goal to develop a quantitative understanding of PAD evolution by using engineered bacteria. For the proposed pilot study, we focused on the technology development.

Summary of the most important results
We have successfully accomplished the proposed objectives in the pilot project. The results of this research have been published in leading biomaterials journal, Biomaterials:

Below, we summarize the key achievements described in detail in the paper.

1. Generating stable droplets to culture and quantify engineered bacteria
To validate the capability of our proposed droplet technology, we fabricated a microfluidic device to produce droplets, and used the droplets to quantify two different population dynamics.

Programmed population control by an ePop circuit: We first use the droplet platform to quantify the population dynamics of an engineered E. coli MC4100Z1 carrying the ePop circuit that we previously engineered. The ePop cells generated oscillatory dynamics in droplets similar to previous observation in bulk cultures (Figure 1A)

Figure 1. (A) Quantification of multiple droplets with oscillating behaviors. Each line represents one subpopulation in droplet started with low cell density (1~3 cells per droplet). Images are the representative time points of ePop oscillation. (B) Investigation of IE in droplets. The engineered bacteria with GFP reporter were encapsulated in droplets with varied concentrations of antibiotics. The curves indicate mean fluorescence intensity of sampled droplets (n>20) versus time, and shades refer to the standard deviation.
Inoculum effect in response to antibiotics: We also investigated the inoculum effect (IE) of a lab bacterial strain within droplets. The sensitive bacterial strain BW25113 with a GFP reporter was encapsulated in the droplets at a high and low initial cell densities, respectively. When the antibiotic concentration was below 4µg/ml, droplets with both high and low initial cell densities grew to a high final density. When the antibiotic concentration was above 4µg/ml, neither of them could grow (Figure 1B top row). Importantly, when the antibiotic concentration was 4µg/ml, the droplets with a high initial density survived, while those with a low initial density did not. This density-dependent survival is the defining character of IE. In contrast, the antibiotics that do not induce HSR and rapid ribosome degradation will not cause IE (Figure 1B bottom row). Droplets with both high and low cell densities either grew ([Cm] ≤ 4µg/ml) or were inhibited ([Cm]>4µg/ml).

2. Demonstrating a droplet injection method to control the droplet environment.

As the original proposal, we introduced a droplet merging method to control the growth dynamics within droplets. Herein, we used an electrode-free injection technology to manipulate the droplet environment that was controlled by a DC power source. When the power source is OFF, the generated droplet is stabilized by surfactant molecules and the injection does not occur. However, when the power source is ON, the electricity field destabilizes the surfactant layer of each droplet as it passes the orifice. The injection channel can then deliver a small amount of liquid into the droplet, forming a larger one (Figure 2).

![Figure 2. Mechanism of droplet injection. A layer of surfactant molecules stabilizes the droplets after production. An electric field disrupts this protecting layer when power is turned ON, and the injection phase is injected into the original droplets when they move across the orifice. Real-time images depict the difference between power OFF and ON. Orange arrows indicate the individual droplets from injection phase without being injected.](image)

3. Applying optimized droplet injection method to precise control growth dynamics in droplets.

We next tested the PAD dynamics in the droplets. We modulated the final concentrations of 6-APA and IPTG in droplets by varying their concentrations in the injection phase. At 25µg/ml 6-APA, the population without any induction of BlaM production initiated growth for the first 510min but stopped growing thereafter (green line in Figure 4A); in comparison, when the population was induced by 1mM IPTG, the cells grew to a higher density; this is due to the degradation of 6-APA by BlaM (red line in Figure 4A). Similarly, when the concentration of 6-APA was increased to
50µg/ml, the discrepancy between un-induced (green line in Figure 4B) and induced populations (red line in Figure 4B) was more significant. For the population without IPTG induction, the inflection point of population growth was slightly shifted to shorter time due to higher stress compared to 25µg/ml 6-APA. However, if the concentration of 6-APA was further increased to 100µg/ml, the discrepancy between the two populations appeared to decrease (Figure 4C). This could be explained by the fact that the stress was too high, and the released BlaM could not efficiently remove the antibiotic. The stress level was also reflected by the inflection point of population growth without IPTG, which shifted to 250min (green line in Figure 4C).

**Figure 3.** (A) 25µg/ml (B) 50 µg/ml and (C) 100µg/ml 6-APA was injected into droplets with (red) and without (green) 1mM IPTG rescuing. Black dash line indicates inflection point when the population stops growing without IPTG.