

Chapter 1

Experimental Background

In this chapter, experimental techniques that are used in my research will be described briefly as a practical guide for those who want to test or perform some parts of the experiments in this thesis. The following aspects will be covered:

- *Escherichia coli* (*E. coli*) bacterial suspensions are the model throughout the whole thesis, so I will start talking about the preparation of motile bacterial sample in Sec. 1.1.
- A key approach I have been using to investigate the properties of bacterial suspensions is optical microscopy. It is used throughout all the experiments in this thesis as well, along with necessary image analysis techniques. Video microscopy and image analysis will be detailed in Sec. 1.2.
- When investigating the rheology of bacterial suspensions, we adopted a homemade microfluidic viscometer device. Details of the fabrication are shown in Sec. 1.3.
- A light-powered *E. coli* strain is used in the giant number fluctuations study and the emergence of active turbulence study (Chap. 3 and Chap. 4). This special strain was obtained by transforming a wild-type strain with an exogenic plasmid

which encodes a light-harvesting membrane protein. The discovery and working principles of the light-powering feature has been well documented by earlier works [?, ?, ?, ?, ?]. Following these works, I constructed a plasmid containing the gene and successfully transformed the wild-type *E. coli* strain. In Sec. 1.4, I will present the details on the materials and procedures I used to construct the mutant as a practical guide to those who need to further modify or trouble shoot the strain I made.

1.1 Motile Bacteria Sample Preparation

Peritrichous *E. coli* bacteria have been widely used as model micro-swimmers for active fluid studies [?, ?]. By bundling and unbundling their flagella, they achieve a so called “run-and-tumble” motion, allowing them to more efficiently explore their surrounding environment and to search for supplies. Fig. 1.1a shows a simplified model of a swimming *E. coli* bacterium model with a 2 μm rod-shape body and a helical-shape flagellum of around 8 μm . When swimming, all the flagella bundle together behind the cell and propel it forward [?]. Fig. 1.1b-c show the bundled state and unbundled state of the flagella, respectively. A swimming *E. coli* bacterium can generate nontrivial fluid flow, which can lead to hydrodynamic attraction to boundaries, alignment with other bacteria and other consequences [?]. It had long been assumed in theoretical works that the effective flow generated by microswimmers like *E. coli* is dipolar, with one force pushing forward from the head and another force pushing backward from the flagella [?, ?, ?, ?]. This assumption was then experimentally verified by Drescher et al. in 2011 [?], by reconstructing the flow field from many tracer particle trajectories. Fig. 1.1d-e show the flow field they measured and the best-fit force dipole flow. As I will show later, the swimming-induced flow plays a key role in the novel properties and collective motions in the bacterial active fluids.

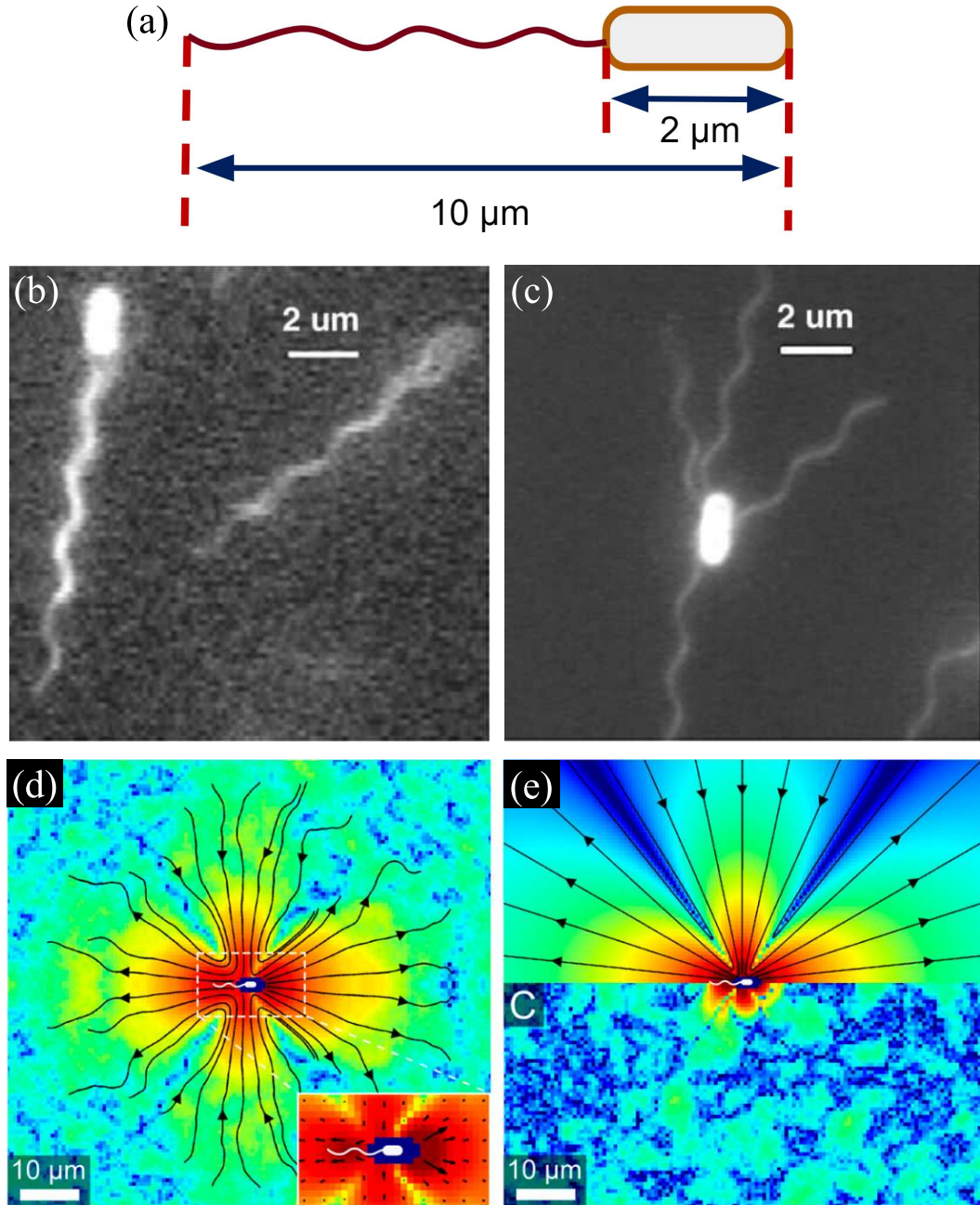


Figure 1.1: **Model swimmer *Escherichia coli* and its flow field.** (a) A schematic of a swimming *E. coli* bacterium. (b) Fluorescence microscopic image of swimming *E. coli* with bundled flagella. (c) Fluorescence microscopic image of tumbling *E. coli* with unbundled flagella. (d) Flow field around a swimming *E. coli*, measured with suspending microspheres. (e) Best-fit force dipole flow for the flow field shown in (d). Image sources: (b) and (c) are reproduced from Fig. 4a and 2a in Ref. [?] with permission from XXX. (d) and (e) are reproduced from Fig. 1a and 1b in Ref. [?] with permission from XXX.

There are quite a few research groups over the world that are using *E. coli* suspensions to study active fluids. To name a few, Yodh and Arratia at University of Pennsylvania, Wu at Cornell University, Poon at the University of Edinburgh and Clement at ESPCI all have published experimental works using *E. coli* [?, ?, ?, ?, ?]. Although the protocols of preparing motile *E. coli* samples are similar across different groups' protocols, they have subtle differences from each other, which may be attributed to the specific strain of *E. coli*, ingredients of media and specific instrument conditions. Schwarz-Linek et al. proposed a sample preparation protocol based on standard bioscience manuals [?] and Berg's *E. coli* protocol. If one wants to learn how to prepare motile *E. coli* from scratch, it is recommended that he/she follows the protocol in Ref. [?].

When I joined the Cheng group at the University of Minnesota in 2015, before Ref. [?] was published, there was already a protocol in our lab that worked pretty well for us. I learned the protocol, and have made some modifications over the years to include the additional procedures for preparing light-powered *E. coli* and to optimize the motility and concentration of samples. Below I describe the protocol that works the best in Cheng lab.

1.1.1 Background Information

Bacterial strains We primarily work on two *E. coli* strains: *AW804* and *BW25113*.

AW804 is light-sensitive. *BW25113* is a wild type strain carrying a plasmid encoding green fluorescence protein, thus it is used when fluorescence / confocal microscopy is needed. Both strains have ampicillin resistance marker and thus require supplementing ampicillin to culturing media.

Antibiotics Bacteria are ubiquitous in the environment and can easily contaminate our bacterial culture. In order to ensure the fidelity of the culture, we add antibiotic resistance markers to the bacteria we want to grow and meanwhile add antibiotics

to the medium. The antibiotics inhibit the growth of contaminating species and allow our desired bacteria to grow normally.

Medium Various types of media (terrific broth, Luria broth, 2XYT and M9, etc.) are commonly used for bacterial culture. We use terrific broth. The recipe can be found in the protocol section.

1.1.2 Protocol

1. Prepare a 2-ml *E. coli* overnight culture.

- (a) Prepare liquid terrific broth (TB). For example, to make 1 L TB, weigh out the following into a 1 L glass bottle:

- 23.6 g Yeast extract (Sigma-Aldrich)
- 11.8 g Tryptone plus (Sigma-Aldrich)
- 4 ml Glycerol (XXX)
- Add dI water to 1 L

Loosely close the cap on the bottle (do NOT close all the way or the bottle may explode!) and then loosely cover the top of the bottle with autoclave tape (stick cap and bottle body together to avoid cap popping off). Autoclave and allow to cool to room temperature. Now screw on the top of the bottle and store the TB at room temperature.

- (b) Using a sterile 10 ml pipette, transfer 2 ml TB to a sterile glass test tube.
- (c) Using a sterile pipette, add 2 microliter (0.1% v/v) antibiotic solution to the TB in test tube.
- (d) Using a sterile pipette tip, pick a small chunk from our bacterial frozen stock (stored in the -80 °C freezer in 251) and carefully transfer the small chunk into the liquid TB + antibiotic.

- (e) Loosely cover the culture with sterile cap that is not air tight.
 - (f) Incubate bacterial culture at 37 °C for 12-18 h in a shaking incubator.
 - (g) After incubation, check growth, which is characterized by a cloudy haze in the media. This is the overnight culture.
2. Dilute overnight culture and harvest motile bacteria at mid-late log phase.
- (a) Using a sterile 10 ml pipette, transfer 3 ml TB to a sterile glass test tube.
 - (b) Using a sterile pipette, add 2 microliter (0.1% v/v) antibiotic solution to the TB in test tube.
 - (c) Transfer 30 microliter (1% v/v) overnight culture into the liquid TB + antibiotic.
 - (d) Incubate bacterial culture at 30 °C for 6-6.5 h in a shaking incubator.
 - (e) After incubation, check for growth, which is characterized by a cloudy haze in the media. This is the log phase bacteria.
3. Centrifuge for better motility and higher concentration bacterial sample.
- (a) Prepare motility buffer (MB), the following recipe is from Ref. [?].
 - 0.01 M potassium phosphate (combine monobasic and dibasic solutions, Sigma-Aldrich)
 - 10^{-4} M EDTA (Sigma-Aldrich)
 - 0.002% weight fraction Tween 20 (Sigma-Aldrich)
 - Adjust pH to 7.0
 - (b) Take out the log phase bacteria from the shaking incubator, centrifuge for 5 min at 800 rcf.

- (c) Discard the supernatant quickly and transfer the left-over liquid to a new centrifuge tube.
- (d) Add 500-1000 ul MB (or water) to resuspend the bottom pellet (avoid bottom pellet) and centrifuge for a second time (5 min, 800 rcf).
- (e) Discard the supernatant and let the tubes sit for two minutes. The remaining left-over liquid should be now filled with the active *E. coli*. Take the left-over solution in another capsule and use it for experiments.
- (f) To measure the concentration, transfer 10 microliter of the suspension into a 1 ml plastic cuvette and dilute 100 times (by adding 990 microliter water). Put the cuvette in the spectrophotometer in 251 and use the OD600 program. The resulting number times 100 will be the number density of your suspension in the unit of n_0 (8×10^8 cells/ml).

1.2 Video Microscopy and Image Analysis: PIV and PTV

1.3 Micro-fabrication and Microfluidics

1.4 Light-controlled E. coli: Genetic Modification, Culturing and Trouble Shooting

Chapter 2

Rheology of Bacterial Suspensions under Confinement^{*}

2.1 Introduction

2.2 Methods

2.3 Results

2.4 Discussion and Conclusion

^{*}Reproduced in part with permission from (Zhengyang Liu, Kechun Zhang and Xiang Cheng, “Rheology of bacterial suspensions under confinement”, *Rheologica Acta*, Springer).

Chapter 3

Giant Number Fluctuations in 3-Dimensional Space

3.1 Introduction

3.2 Methods

3.3 Results

3.4 Discussion and Conclusion

Chapter 4

The Emergence of Active Turbulence^{*}

4.1 Introduction

4.2 Methods

4.3 Results

4.4 Discussion and Conclusion

^{*}Reproduced in part with permission from (Yi Peng, Zhengyang Liu and Xiang Cheng, “Imaging the emergence of bacterial turbulence using light-powered *Escherichia coli*”, *arXiv e-print*).