Analyzing Inhibitory Effects of Reagents on *Mycoplasma* Gliding and Adhesion

Taishi Kasai and Makoto Miyata*

Department of Biology, Graduate School of Science, Osaka City University, Osaka, Japan

*For correspondence: miyata@sci.osaka-cu.ac.jp

[Abstract] Dozens of *Mycoplasma* species bind to solid surfaces and glide in the direction of the membrane protrusion at a pole. In gliding, *Mycoplasma* legs catch, pull and release sialylated oligosaccharides fixed on a solid surface. The analyses of inhibitory effects of sialylated compounds on gliding of *Mycoplasma* can determine the target structure of *Mycoplasma* for gliding and adhesion.

**Materials and Reagents**

1. *M. mobile* 163K strain
2. *M. pneumoniae* M129 strain
3. Horse serum (Life Technologies, catalog number: 16050-122)
4. Interested reagents (for example, 0.05-1 mM 3’-sialyllactose, Dextra Laboratories, catalog number: SL302)
5. Heart infusion broth (Becton, Dickinson and Company, catalog number: 238400)
6. Yeast extract (Becton, Dickinson and Company, catalog number: 212750)
7. Amphoterin B (Sigma-Aldrich, catalog number: A2942)
8. Ampicillin (Nacalai Tesque, catalog number: 02739-32)
9. NaCl
10. Sodium phosphate (pH 7.3)
11. Glucose
12. Aluotto medium (see Recipes)
13. PBS-G buffer (see Recipes)

**Equipment**

1. Test tubes
2. Tissue culture flask
3. Scraper (Greiner Bio-One GmbH, catalog number: 541070)
4. 0.45-μm-pore size filter (EMD Millipore, catalog number: SLHVX13NK) (Polyvinylidene Diffuoride, or equivalent)
5. 25 °C incubator (Yamato, model: IL62)
6. 37 °C incubator (Yamato, model: IS62)
7. Centrifuges (Sigma Zentrifugen, model: SIGMA 1-14)
8. Optical microscope (Olympus, model: BX51)
9. Stage heater (for *M. pneumoniae*) (Minitube, model: HT200)
10. Lens heater (for *M. pneumoniae*) (Tokai Hit, model: MATS-LH)

**Software**

1. ImageJ

**Procedure**

A. *M. mobile*

1. Cultivate *M. mobile* cells in a growth medium (Aluotto medium) at 25 °C incubator to mid log phase.
2. Collect the cells in mid log phase at 0.03-0.08 of OD$_{600}$, by centrifugation at 13,000 x g for 4 min at room temperature (RT) and suspend them in the Aluotto medium to be OD$_{600}$ = 1.0.
3. Wash the cells three times by centrifugation followed by suspension, and resuspend the cells with PBS-G buffer by the same volume with the Aluotto medium used in A-2.
4. Prepare a tunnel chamber (5 mm interior width, 18 mm length, 60 µm wall thickness) composed of a coverslip, a glass slide, and double-sided tapes (see Figure 1).

![Mycoplasma Suspension](image)

**Figure 1. Tunnel chamber**
5. Insert 10-50 μl of the cell suspension into the tunnel chamber to fill the tunnel.

6. Observe the cells bound to the coverslip with microscope with a 100x phase-contrast objective lens with video recording (see Figure 2 left).

7. Replace the PBS-G buffer with the buffer containing the interested reagents (i.e. sialylated compounds, monoclonal antibody, etc.). Put the buffer on one side of tunnel and suck the buffer from the other side using a filter paper.

8. Count the number of bound cells by a command “analyze > analyze particles” of ImageJ, an image analyzing software. Unbound cells are in Brownian motion and cannot be counted by this Image J command, owing to the insufficient image density.

**B. M. pneumoniae**

1. Cultivate *M. pneumoniae* cells in Aluotto medium at 37 °C incubator to mid log phase. The density of cells is not detectable because the cells are not floating in the medium. However, the density in mid log phase should be 0.02-0.05 at OD$_{600}$, if the cells are suspended into the medium.

2. Replace the medium by 2-5 times smaller volume of PBS containing 10% horse serum.

3. Scrape the bottom of culture flask to release *Mycoplasma* cells into the solution, because the cells adhere to the flask bottom tightly.

4. Recover the cell suspension.

5. Filter the suspension through a membrane filter unit with a 0.45 μm-pore size.

6. Prepare a tunnel chamber.

7. Insert 10-50 μl of the cell suspension into the tunnel chamber to fill the inside.

8. Incubate the tunnel chamber for 60 min, with facing of coverslip-side to the stage heater. The water evaporates only slightly in Japanese climate, but the tunnel chamber should be cover by a lid of Petri dish.

9. Observe the cells bound to the coverslip by the microscope with a 100x phase-contrast objective lens heated by the lens heater, with video recording (see Figure 2 right).
10. Replace the buffer with the buffer containing the interested reagents. Put the buffer on one side of tunnel and suck the buffer from the other side using a filter paper.

11. Count the number of bound cell by a command “analyze > analyze particles” of ImageJ, an image analyzing software.

Recipes

1. Aluotto medium
   2.1% heart infusion broth
   0.56% yeast extract
   10% horse serum
   0.025% amphotericin B
   0.005% ampicillin

2. PBS-G buffer
   68 mM NaCl
   75 mM sodium phosphate (pH 7.3)
   10 mM glucose

Acknowledgments

This protocol is adapted from Kasai et al. (2013).

References