

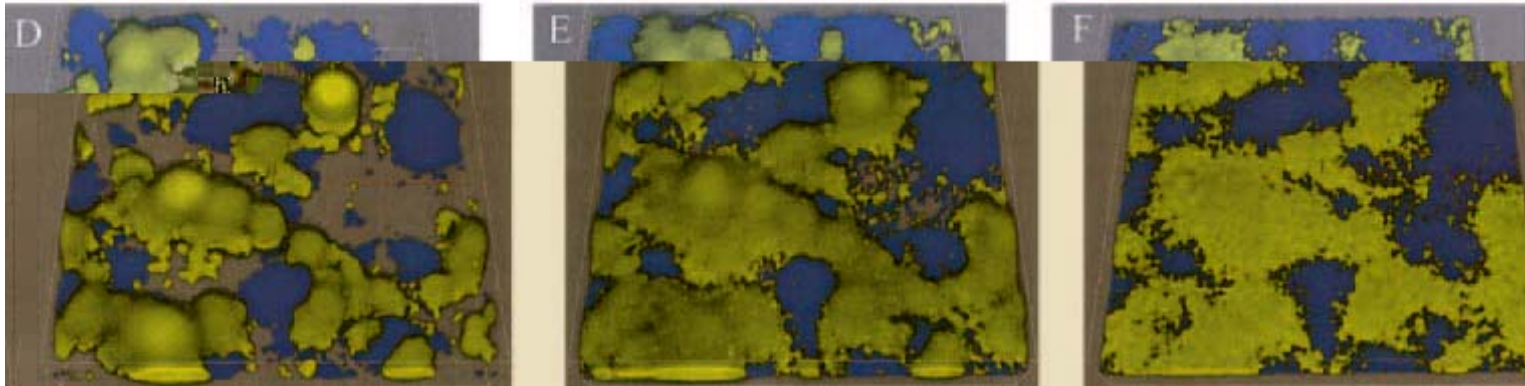
# Structure in biofilms: How does it develop, and what roles does it play?

**Vernita D. Gordon**

Department of Physics, Center for Nonlinear  
Dynamics, Institute for Cellular and Molecular  
Biology, University of Texas, Austin

Talk given at Beijing University, May 24, 2012

# Why study biofilms?



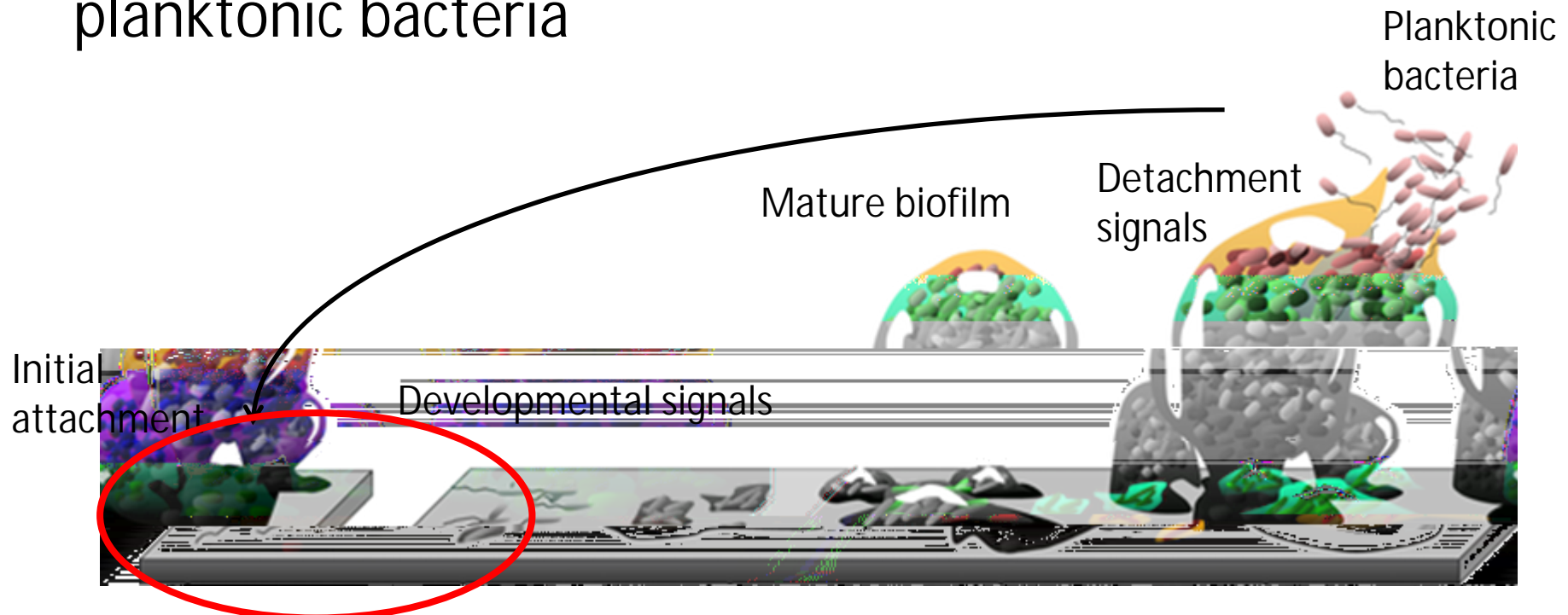
Klausen, M.

48, 1511–1524 (2003).

- Biofilms are multicellular communities of single celled organisms that form at surfaces
- Very common! Most wild bacteria are found in biofilms
- Important in both medical and industrial settings
  - Increased antibiotic resistance and virulence
  - Biofouling of medical devices, pipes, ship hulls
- Model system for multicellularity
  - Simple, easy to tweak

# Biofilms development involves several stages

- Early stages include attachment to a surface and production of extra cellular polysaccharides (EPS)
  - Pel and Psl are two main EPS elements for
- Complex mature biofilms structured by EPS
- Distinct phenotypes (gene expression) from planktonic bacteria



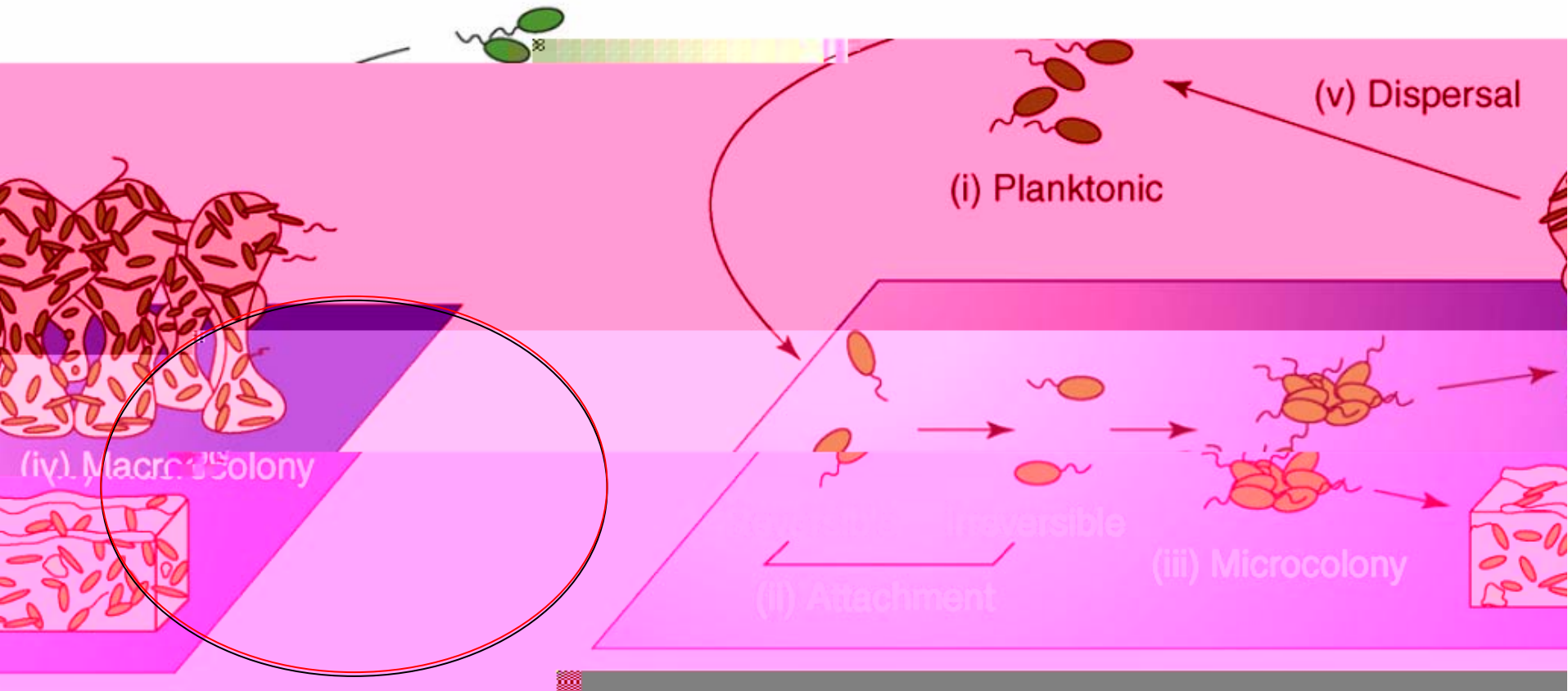


- Ubiquitous bacteria: found in/on water, soil, skin, etc.
- Opportunistic human pathogen, common in hospitals
- Causes serious lung infection in cystic fibrosis patients
  - Most common genetic disease in U.S.
  - Life expectancy ~30 years
- Gram negative, rod shaped bacteria ( $\sim 1 \text{ } \mu\text{m} \times \sim 2 \text{ } \mu\text{m}$ )
- Single polar flagellum, type IV pili
- Readily forms biofilms

Question 1

**WHAT ARE THE TYPES OF SURFACE  
MOTILITY LEADING TO BIOFILMS?**

# Canonical Picture of Biofilm Formation



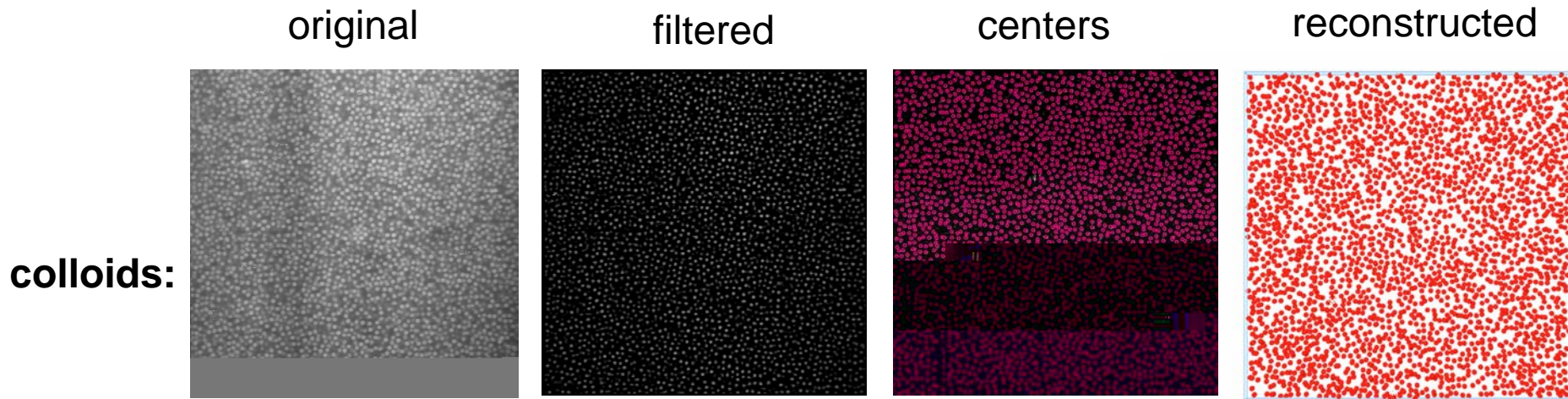
These cells are motile.

Figure from Monds and O'Toole,  
Trends in Microbiology 2009

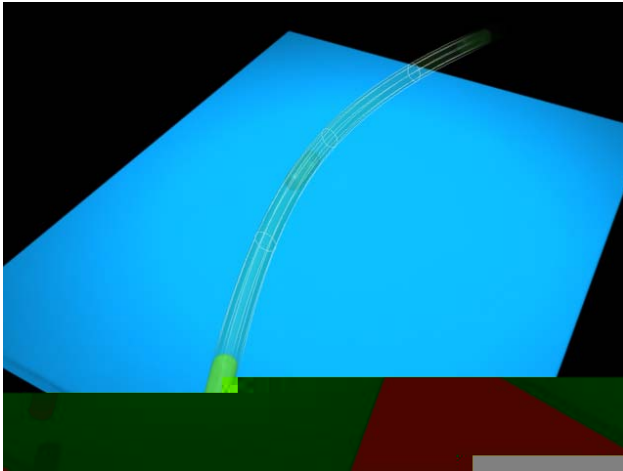


# High throughput tracking and biometric analysis of bacterial surface motility

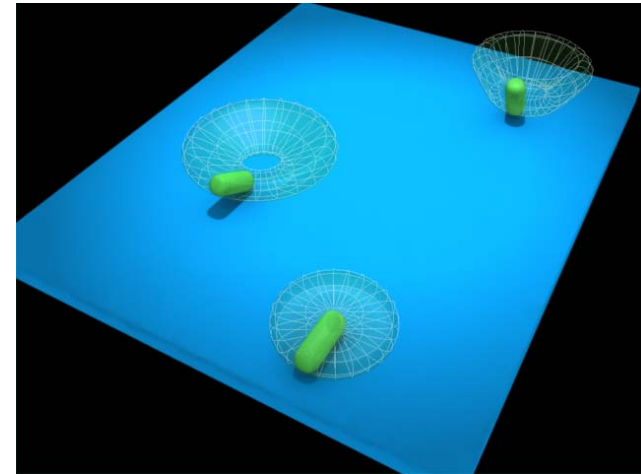
- Codes developed for colloid physics :
- Find centers (& characteristics - orientation, aspect ratio, etc.)
- Link coordinates and characteristics to form trajectories.
- Trajectories reconstruct the original movie's moving bacteria



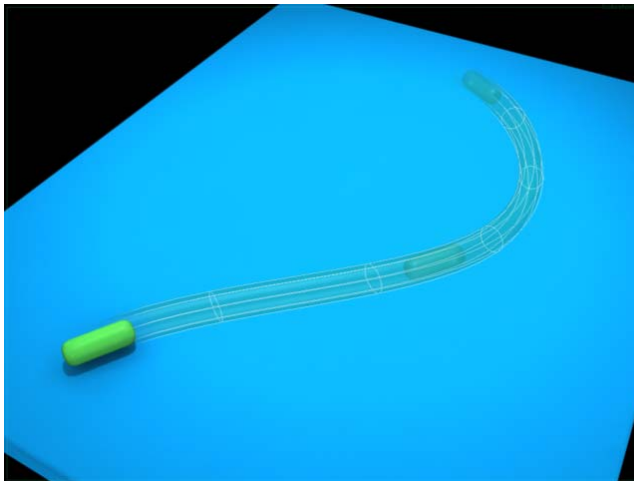
# Tracking identifies distinct motility modes



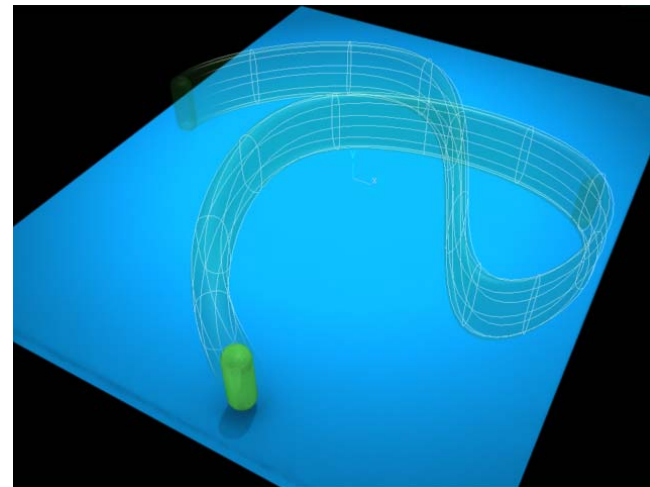
Flagellum-based “**skimming**”



Flagellum-based “**Spinning**”



Pili “**Crawling**” motility



Pili “**Walking**” motility

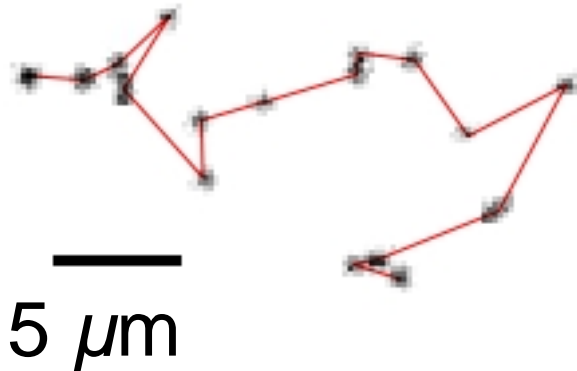


# “Walking” motility

Oriented perpendicular to the surface.

No preferred direction of motion.

trajectories

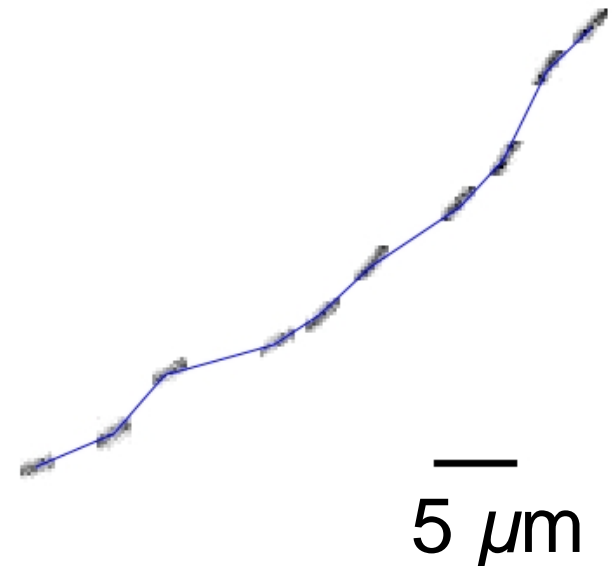


# “Crawling” motility

Oriented flat on the surface.

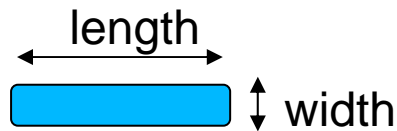
Move along their body axis.

trajectories.

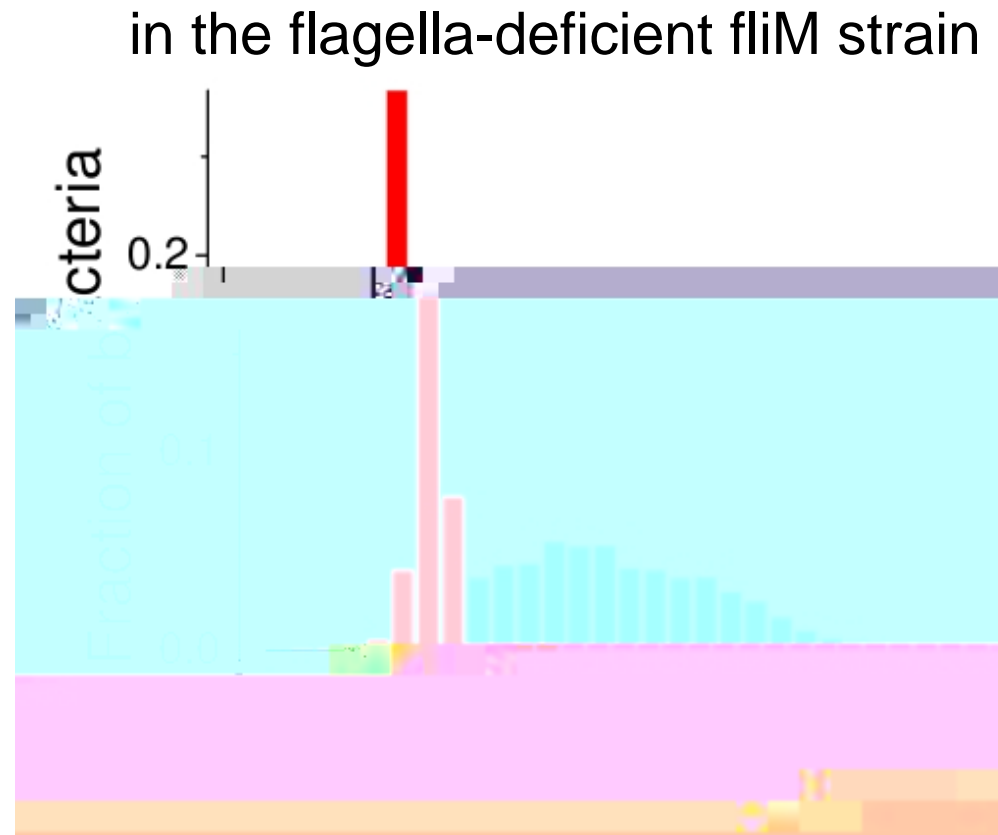


# Motility modes have signature orientations as well as trajectories

- Two peaks in the X Y projected length
  - correspond to the average width and length of a bacterium.

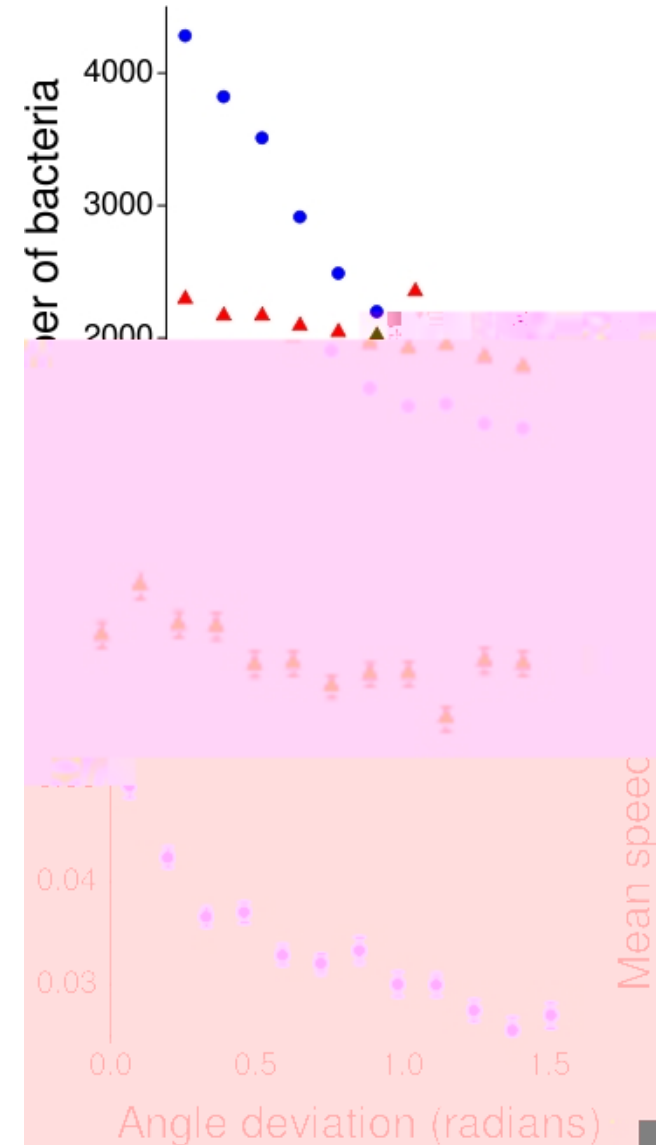
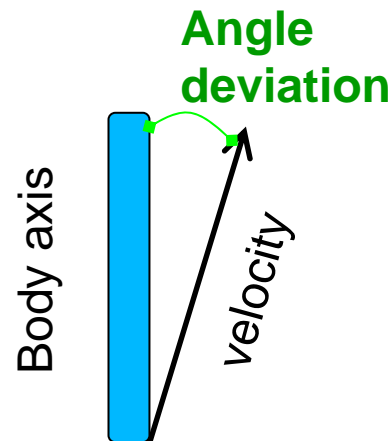


- Up to 50% of bacteria are “walking.”

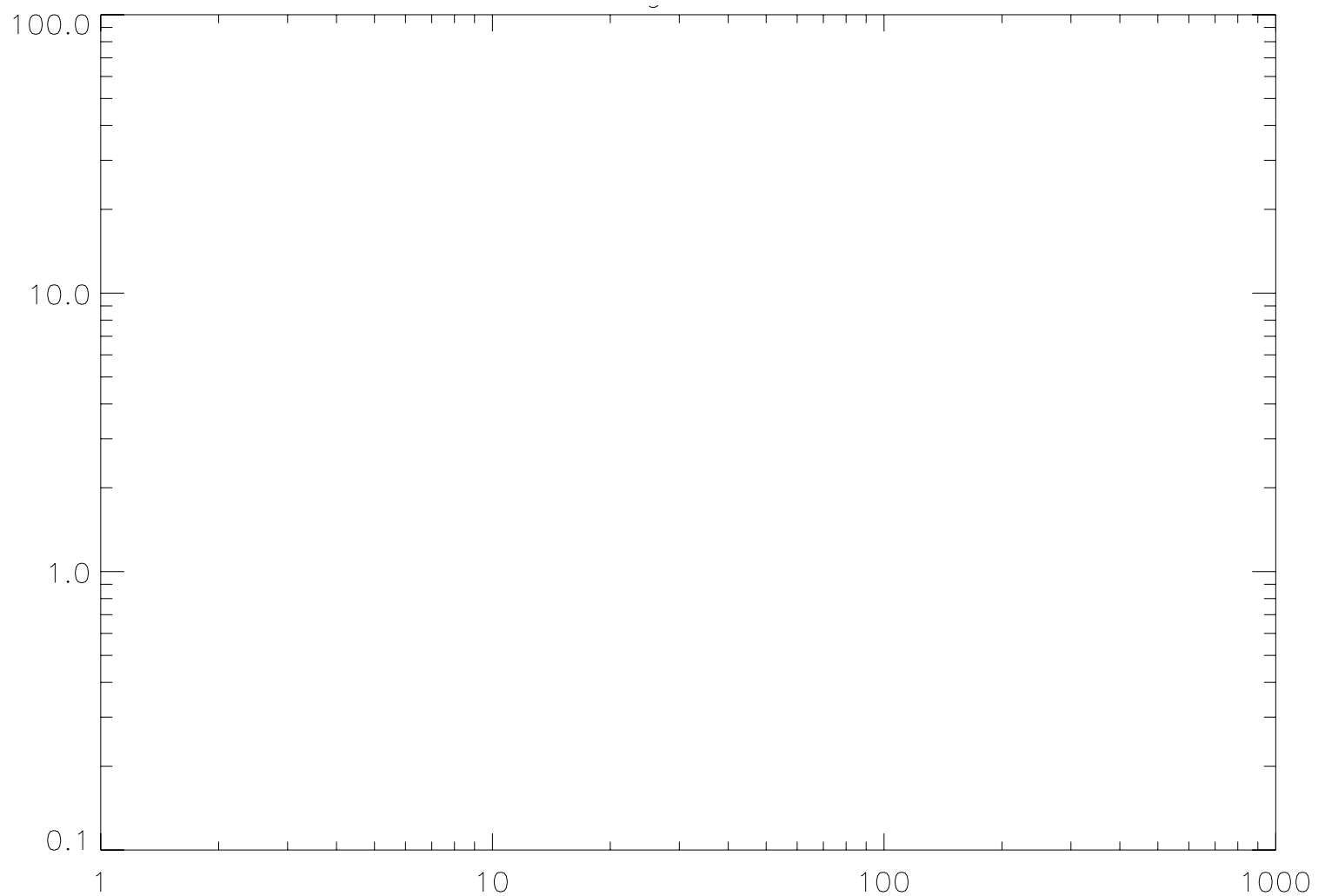


# Motility comparison

- “Crawling” has a preferred direction
- “Walking” has a higher average instantaneous velocity



# Walking, Crawling both superdiffusive



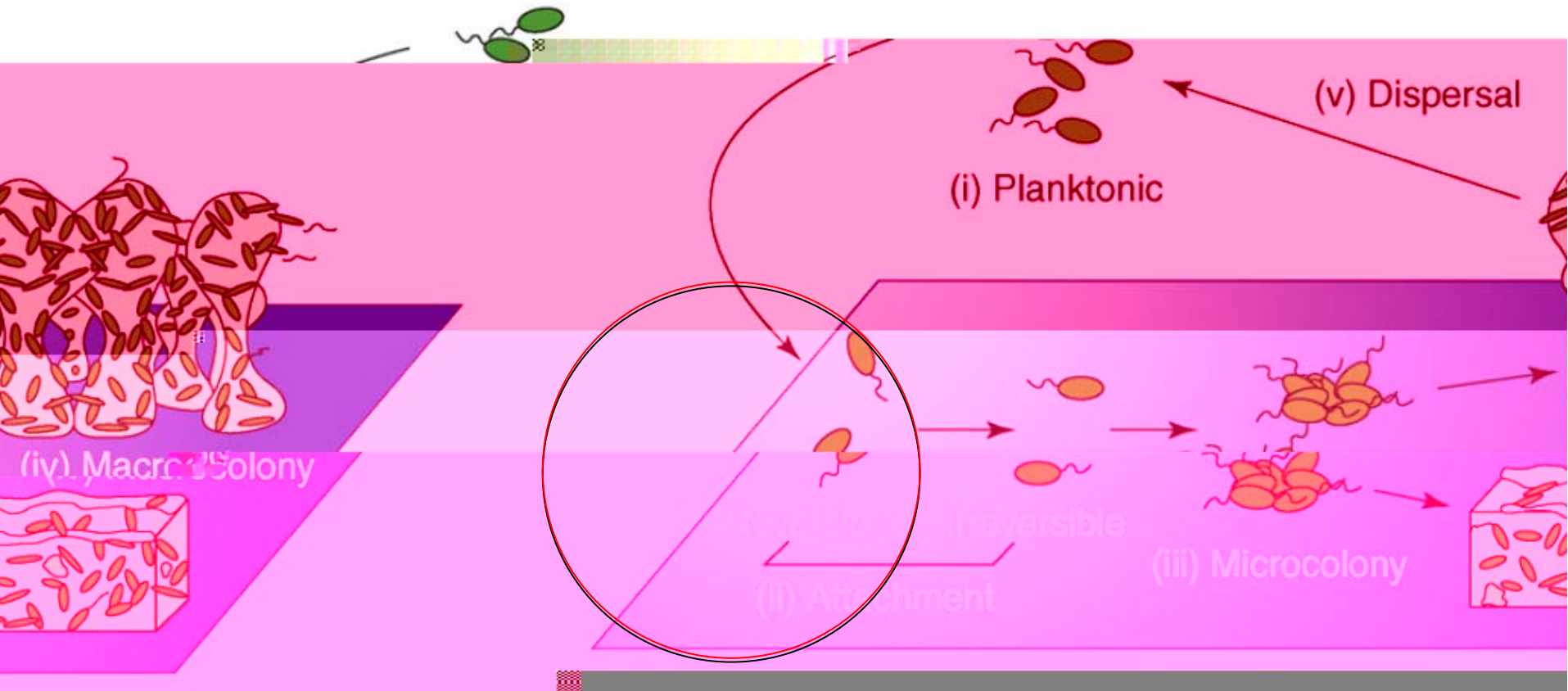
## What we've learned:

- There are two pili driven surface motility modes, flat “crawling” and vertical “walking”.
- “Walking” is not directional (short persistence length), and allows the bacterium to explore its local environment.
- “Crawling” has a preferred direction.

# WHAT ARE THE ROLES OF EXTRACELLULAR POLYSACCHARIDES



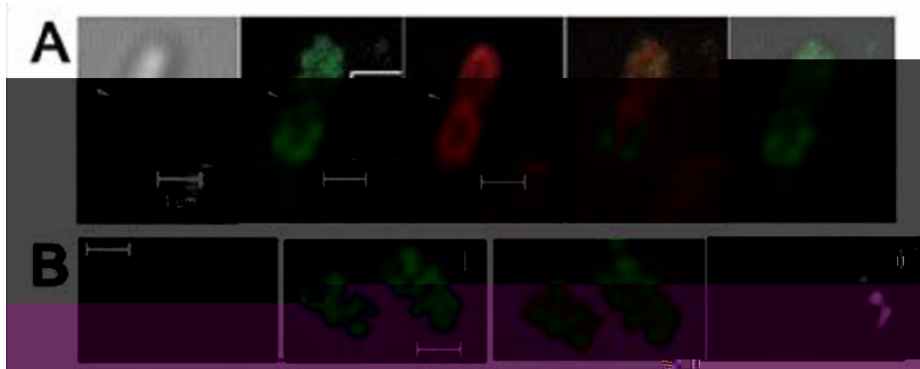
# Canonical Picture of Biofilm Formation



Cells in microcolonies  
stick to each other.

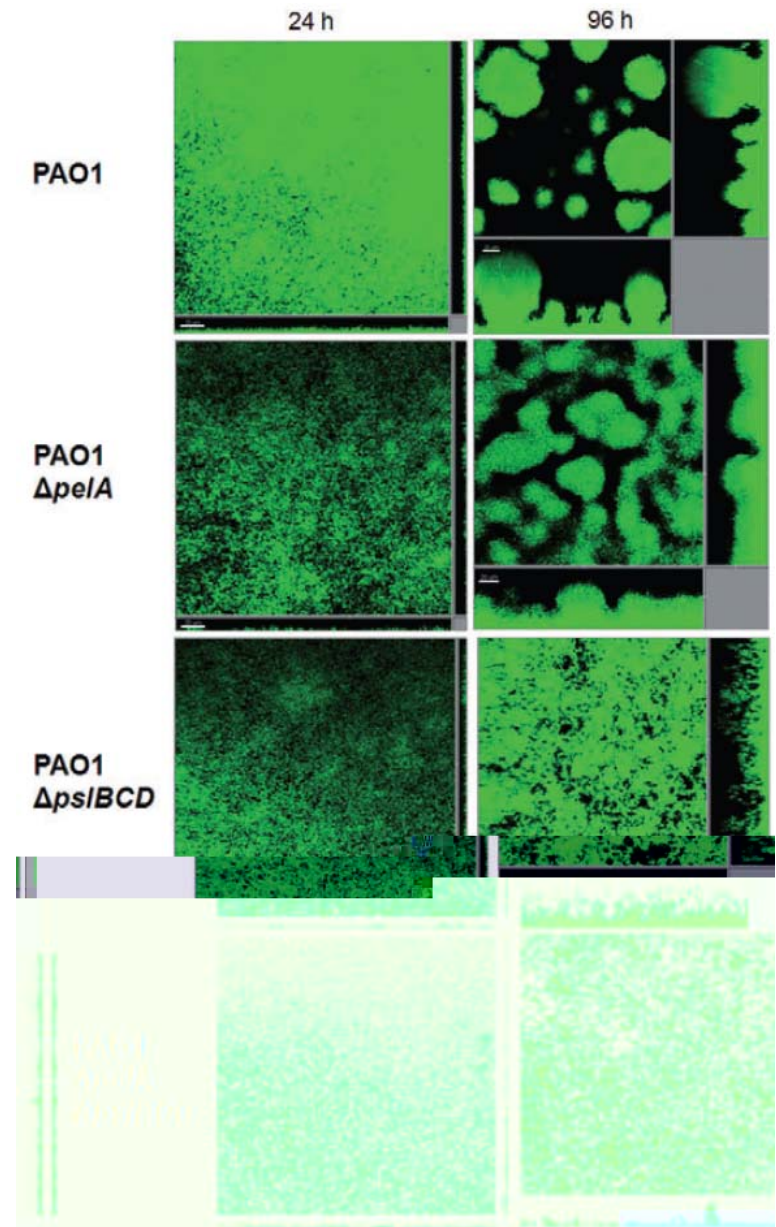
Figure from Monds and O'Toole,  
Trends in Microbiology 2009

# Previous work: Psl > surface adhesion, Pel > self cohesion



Ma, , PLOS Pathogens 5, 1000354 (2009)

- Psl (above) forms helical structures around surface of bacteria
- Structure Pel makes is unknown
- Previous studies showed two distinct roles for Pel and Psl in biofilm formation



Yang, , Environmental Microbiology 13, 1705 (2011)

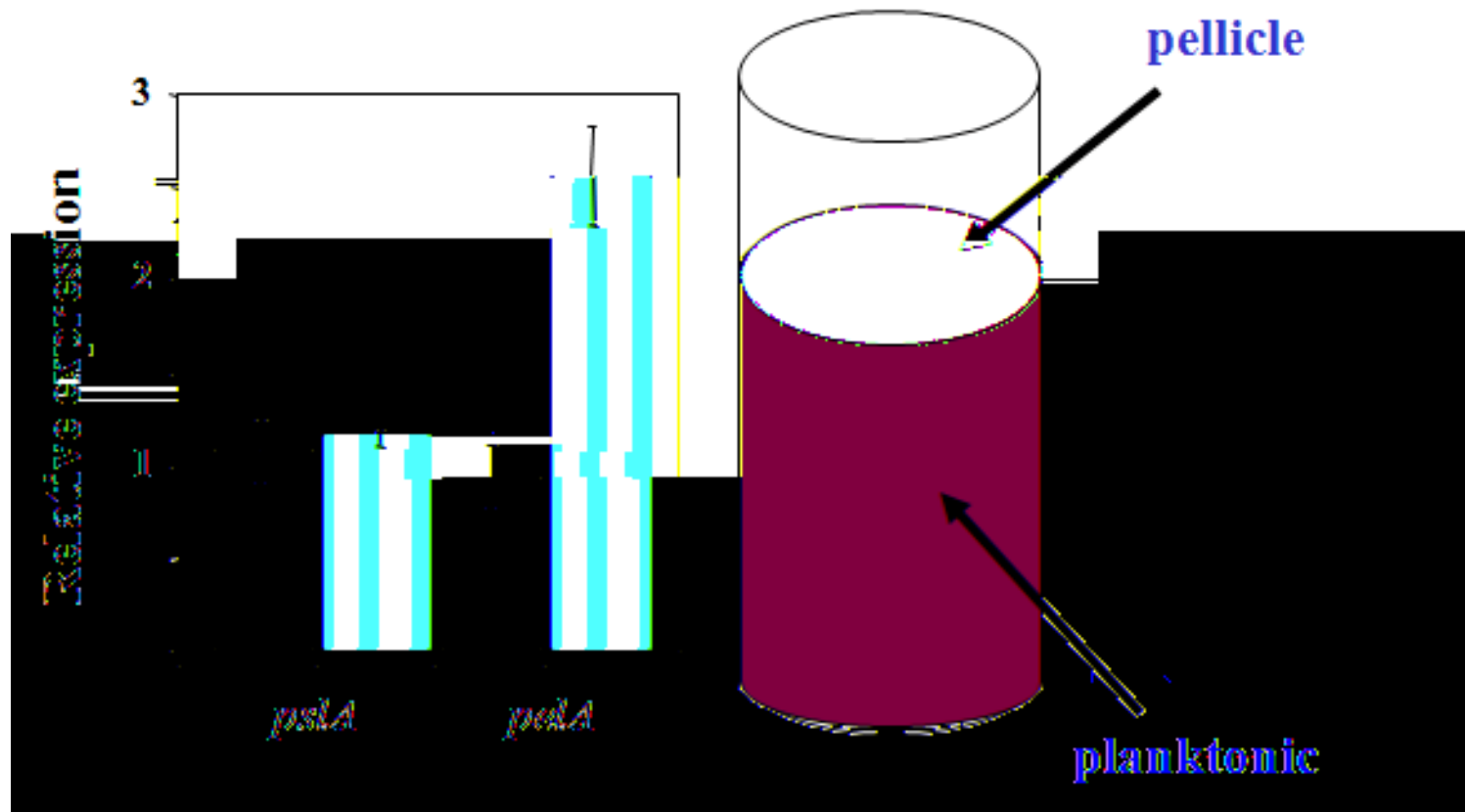
## **Open question:**

**What are the key initial steps for microcolony formation and biofilm initiation?**

**Bacteria must**

- **sense they are at a surface**
- **initiate production of some EPS**  
many possible candidates
- **interact specifically with other bacteria**

# *pel* expression is induced in pellicles formed in standing liquid cultures



Data from Borlee and Parsek, University of Washington, Seattle

# *pelA* expression induced after surface adhesion

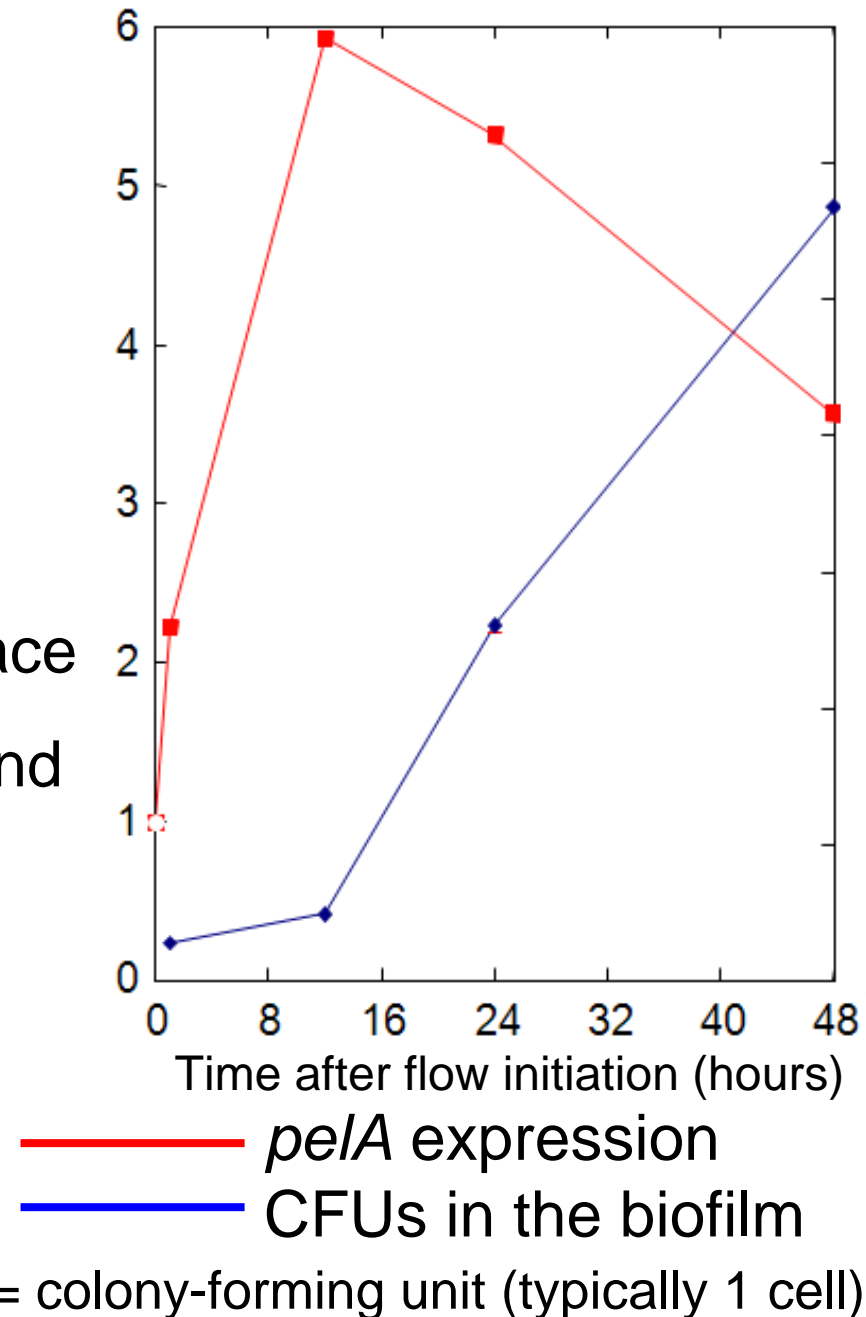
*P. Aeruginosa* biofilm grown in a silicone tube:

- incubate statically for 30 min
- begin flowing fresh medium
- adherent cells harvested off surface

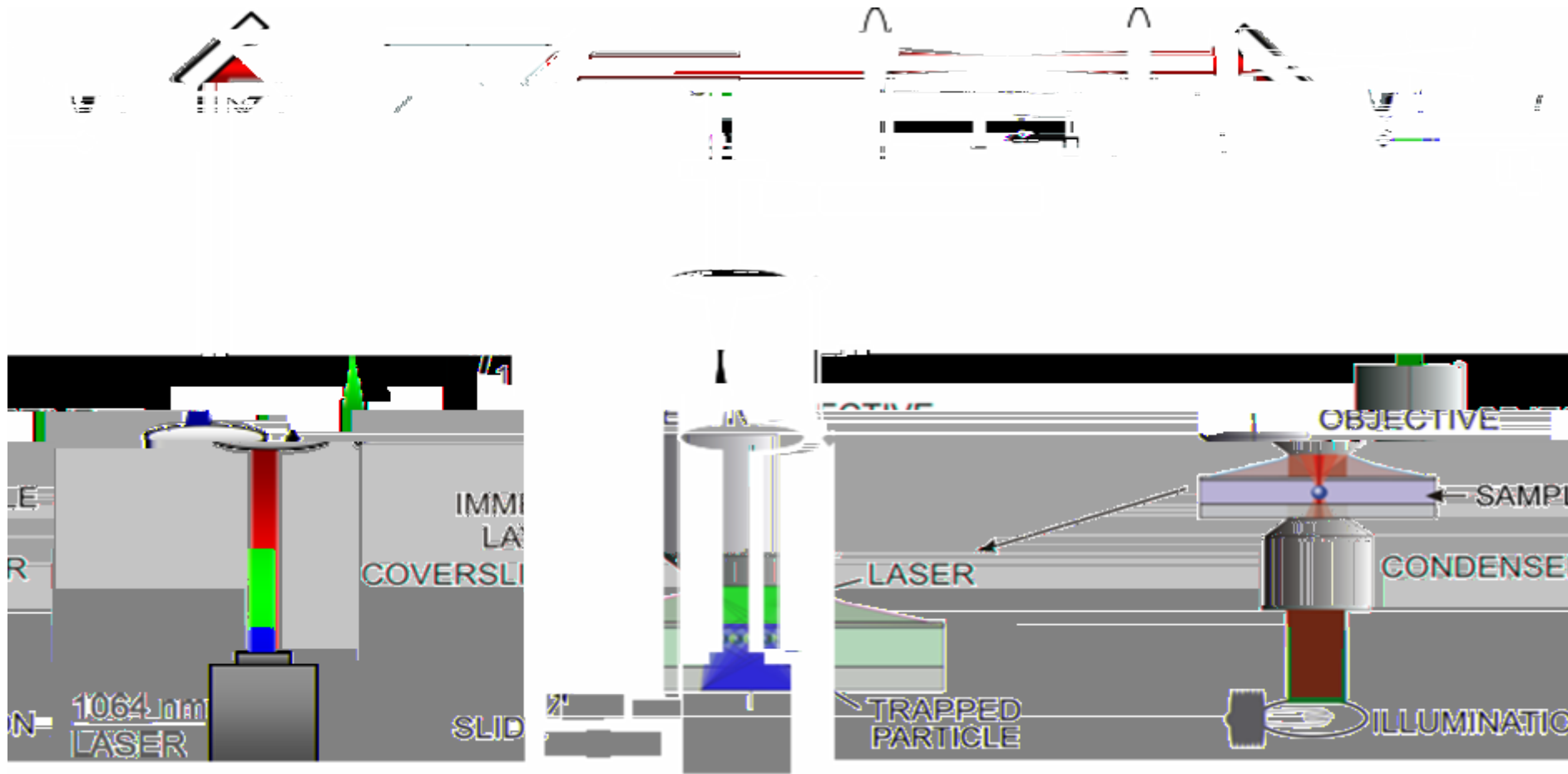
monitor gene transcription levels and viability of cells in biofilm:

*Pel turns on early in biofilm development, but turns off as the biofilm matures.*

Data from Borlee and Parsek,  
University of Washington, Seattle



# Laser-trapping setup

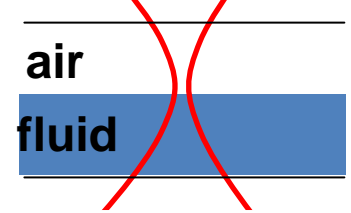


- Built on inverted microscope
- Simultaneous trapping and imaging in brightfield transmission or fluorescence



# Laser directed aggregation

Making Pel is essential for bacteria aggregation on short timescales!

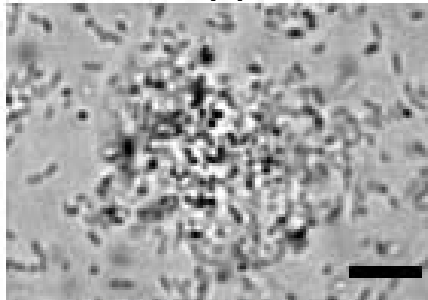


40x LWD objective

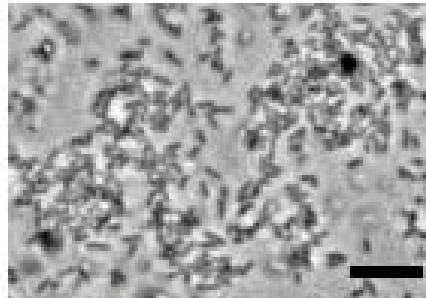
trapped

released

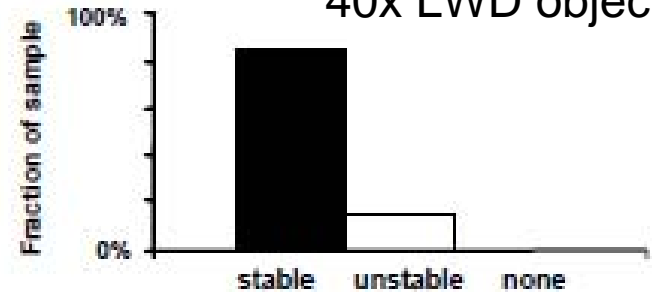
Pel



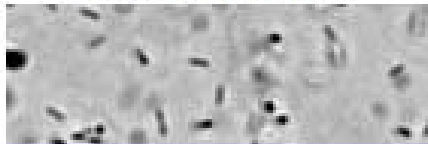
PA14 trapped 20 min



PA14 released 5 min

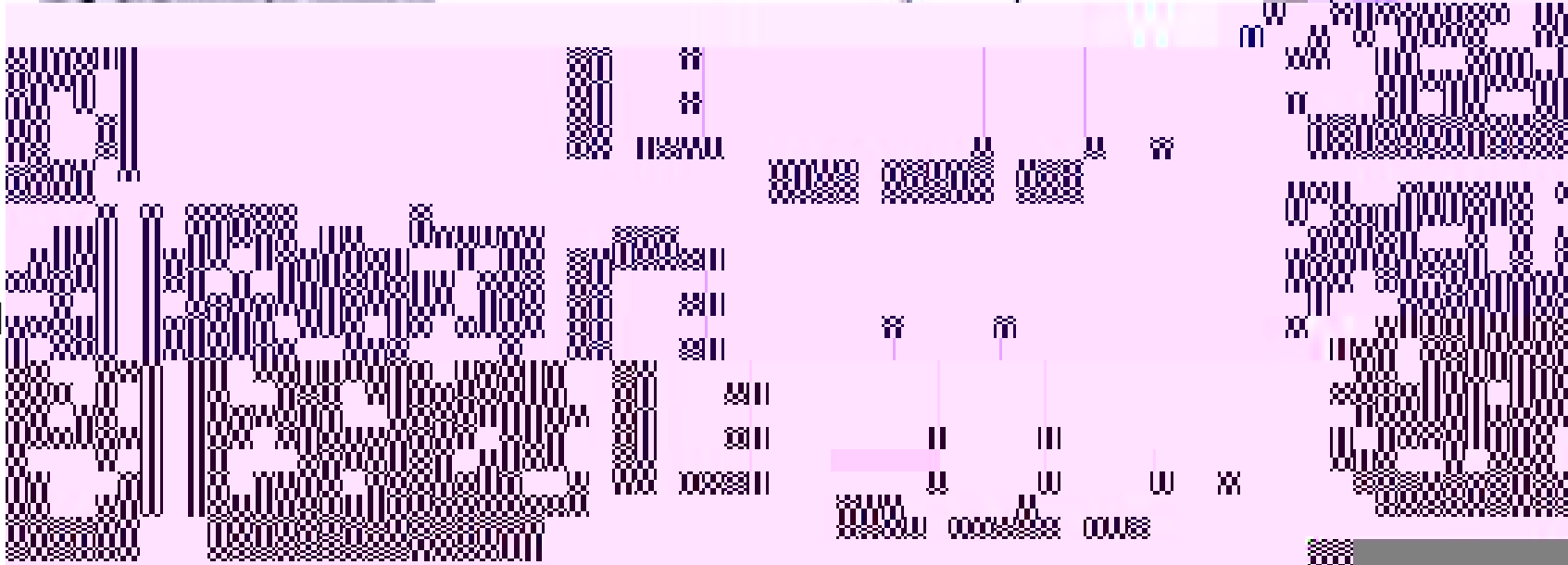


No Pel



ample

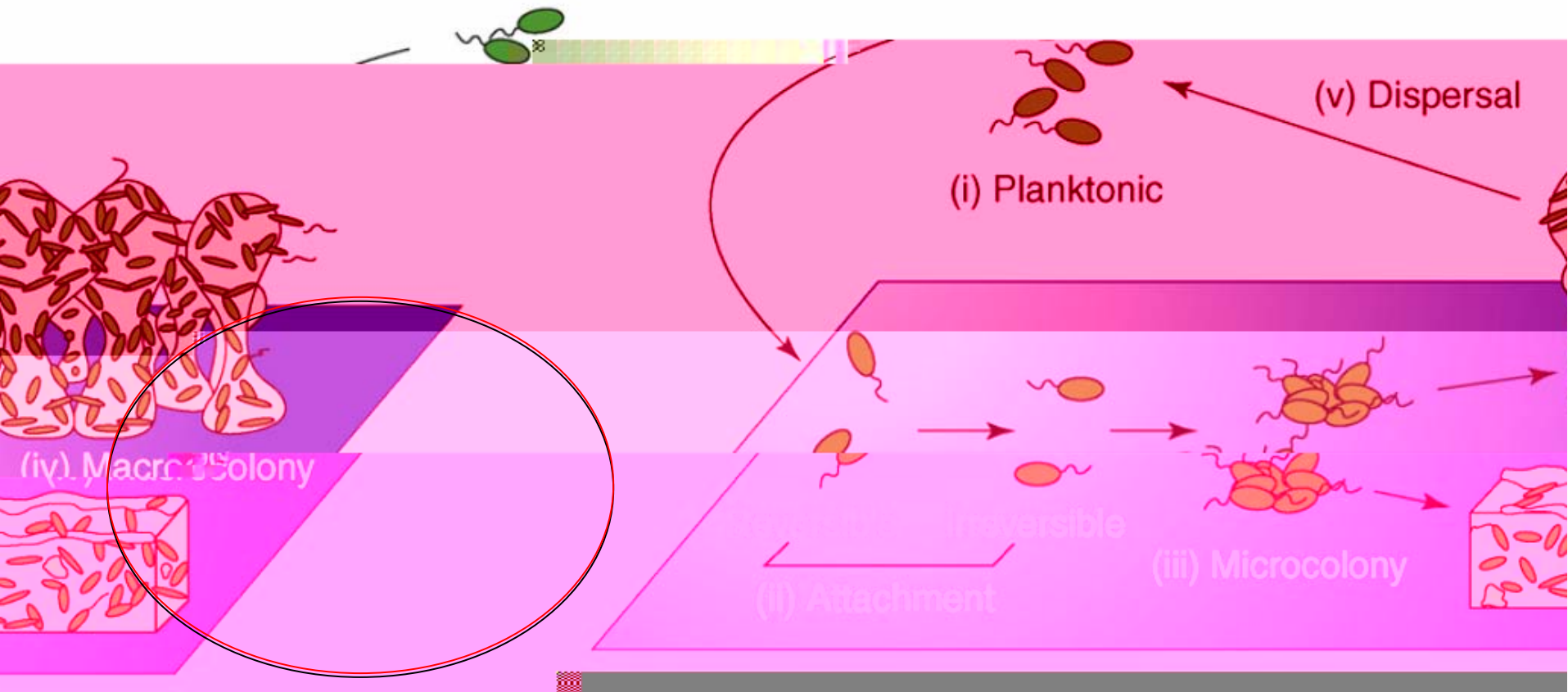
No Pel



# What we've learned:

- is the molecular glue first activated
- is responsible for inter bacterial adhesion early in biofilm development

# Canonical Picture of Biofilm Formation



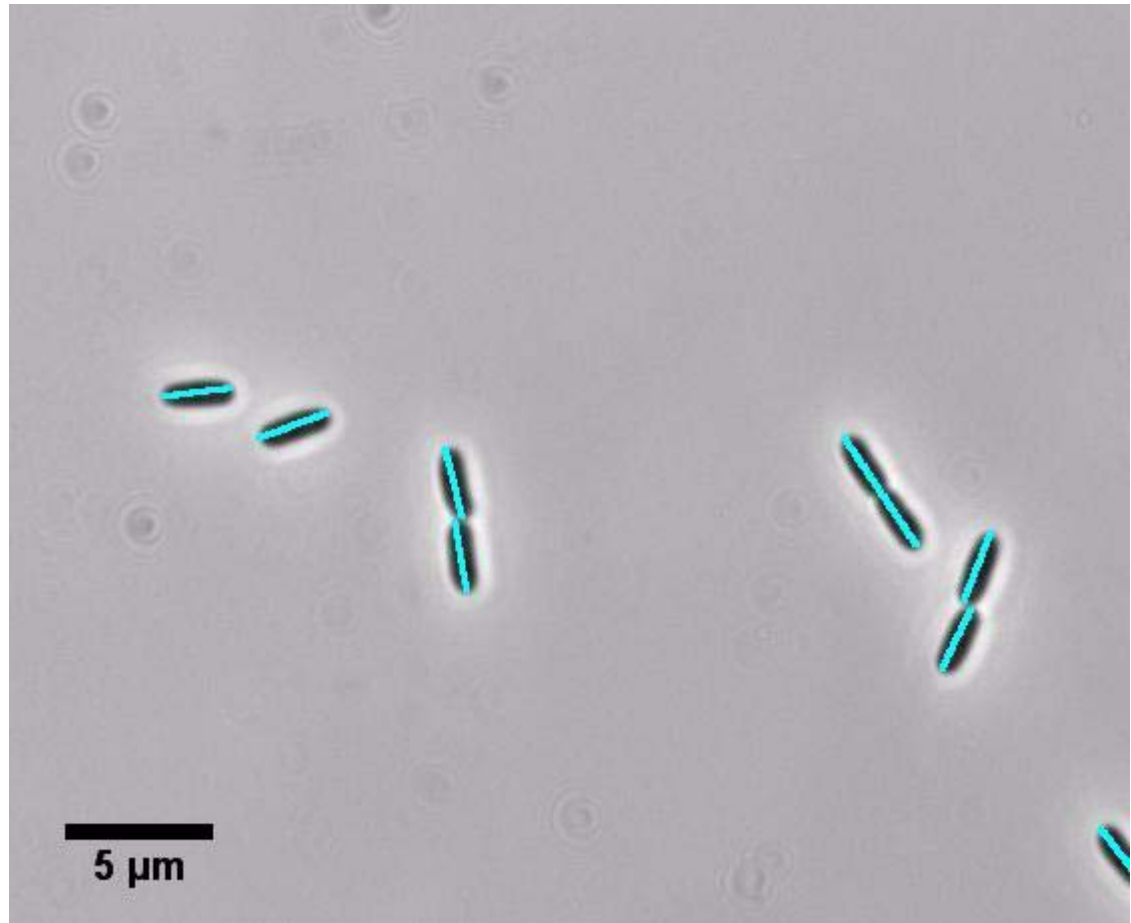
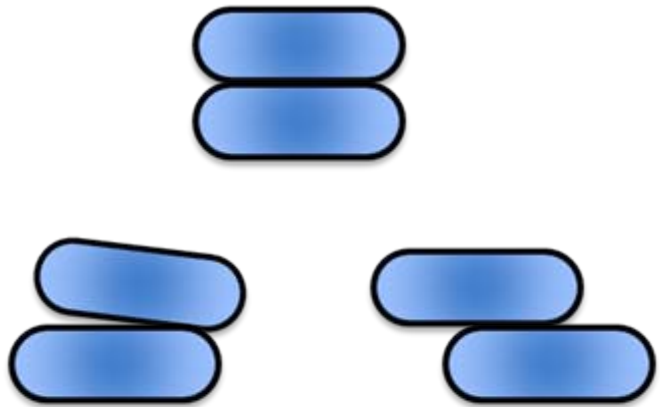
Cells land end on, and lie down flat  
as part of irreversible attachment.

Figure from Monds and O'Toole,  
Trends in Microbiology 2009

# Measuring effects of EPS in very early biofilms

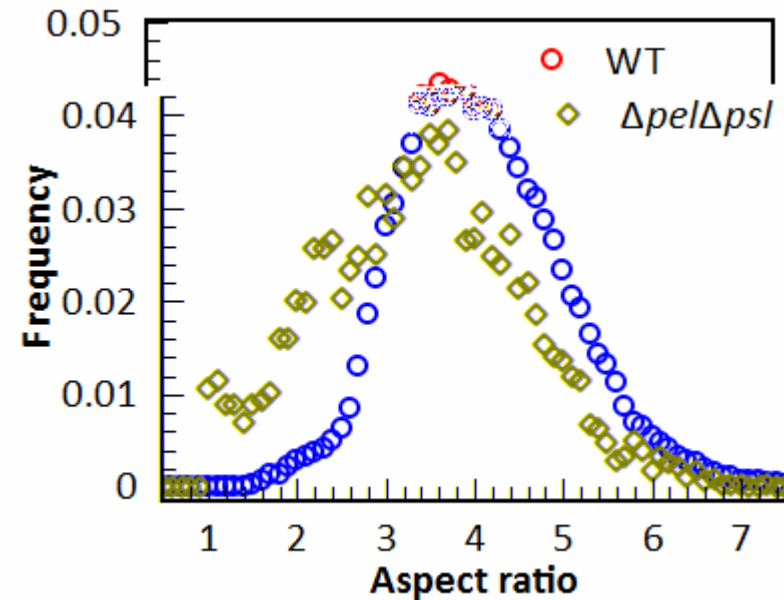
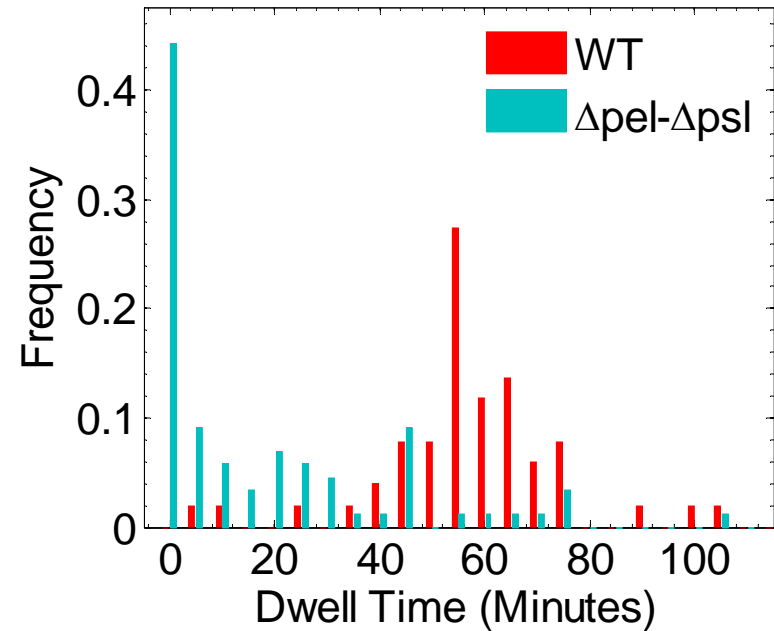
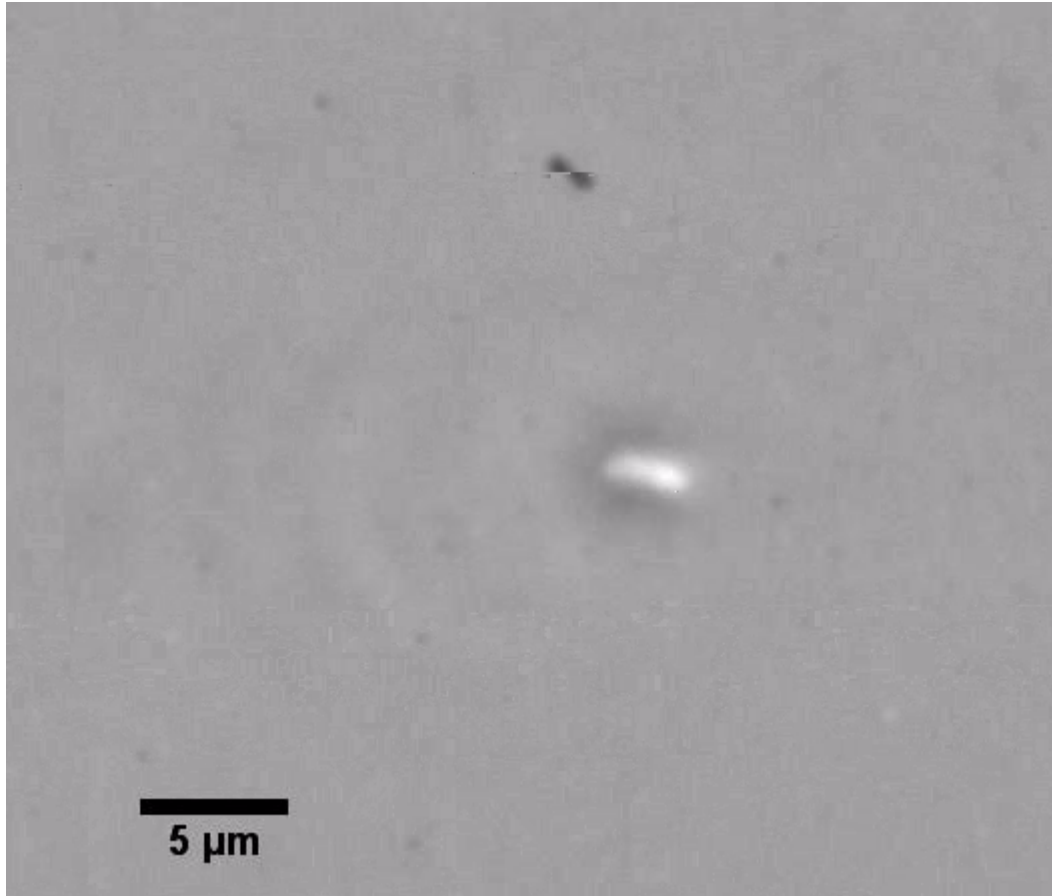
- Tracking code identifies individual bacteria and outputs position, speed, direction, length, aspect ratio

Identify self cohesive bacteria (side by side)



# $\Delta$ $\Delta$ has severely impaired surface adhesion

- Agrees with previous results



–  $W$

–  $\Delta$

- Number  
greater

–  $W$

–  $\Delta$

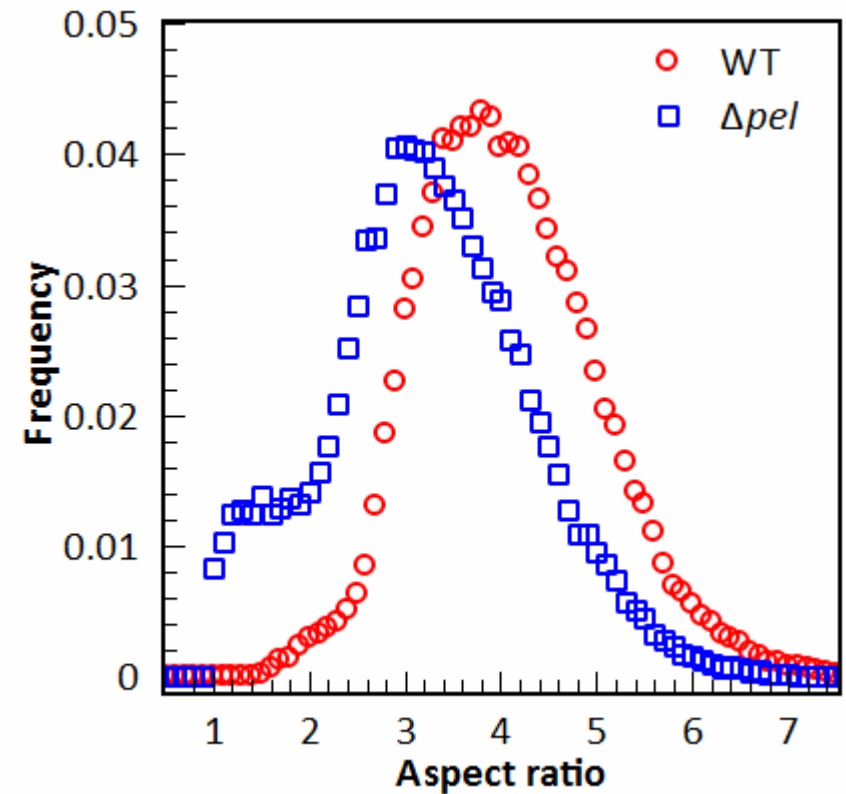
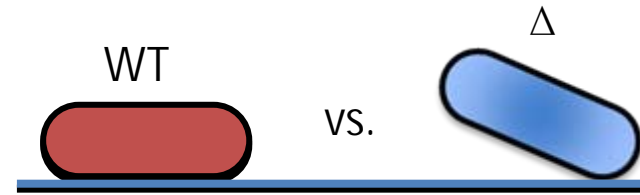
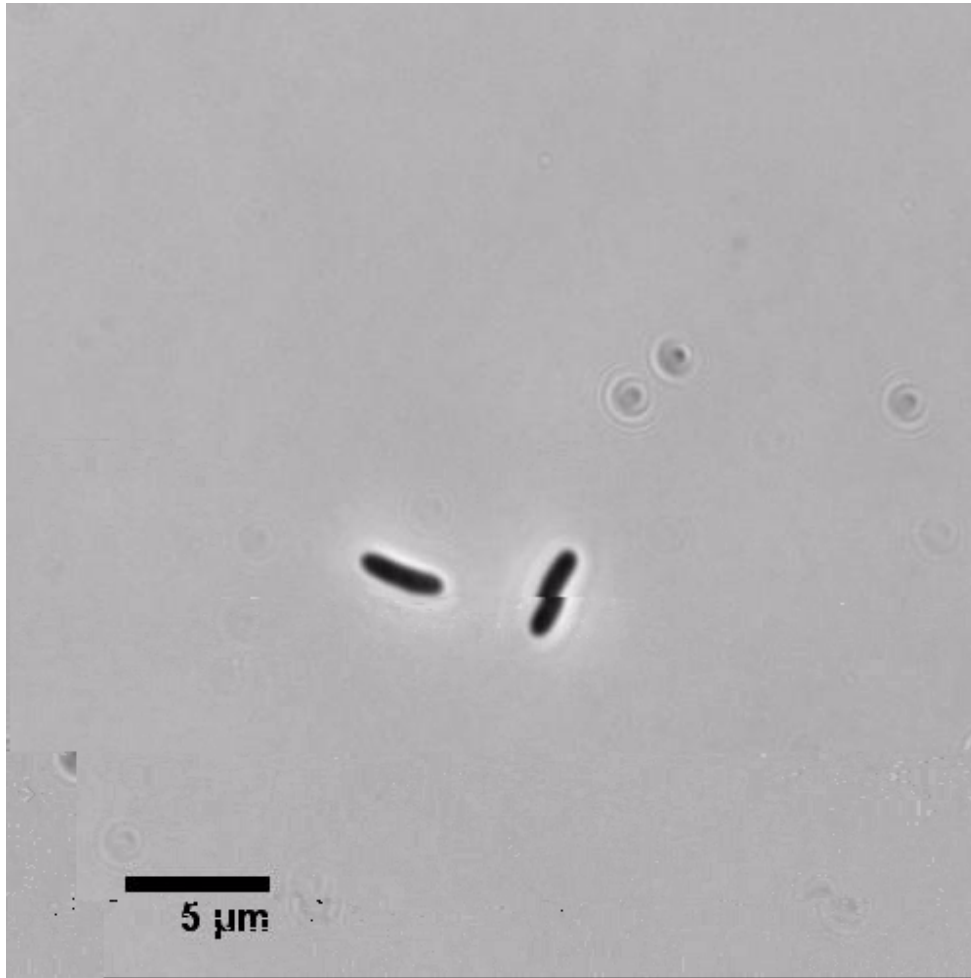
- Percentage  
longer

–  $W$

–  $\Delta$

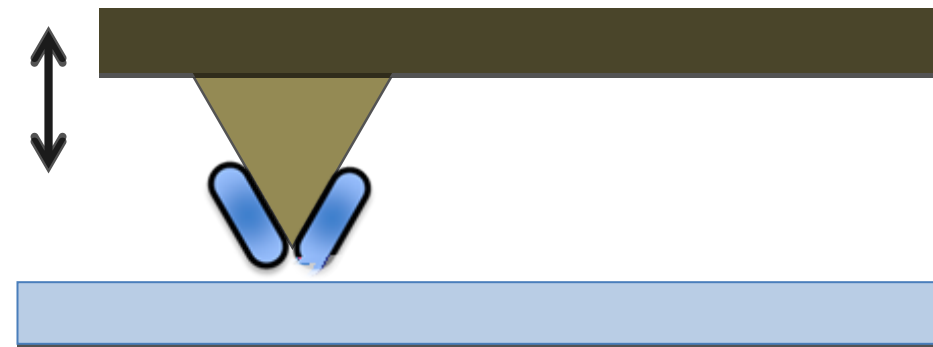
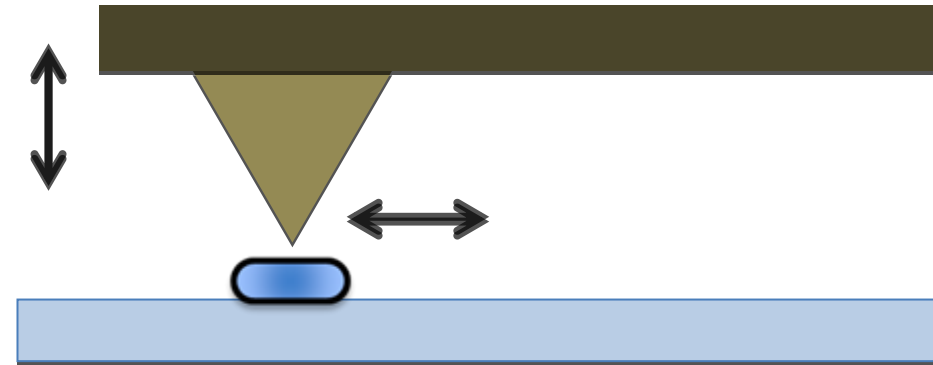


# Surprise! Pel also mediates surface adhesion!



# AFM adhesion force measurements

- Directly measure difference in adhesion between WT and mutants
- Two methods
  - Attach bacteria to surface
  - Attach bacteria to tip
- All measurements done in liquid with live bacteria



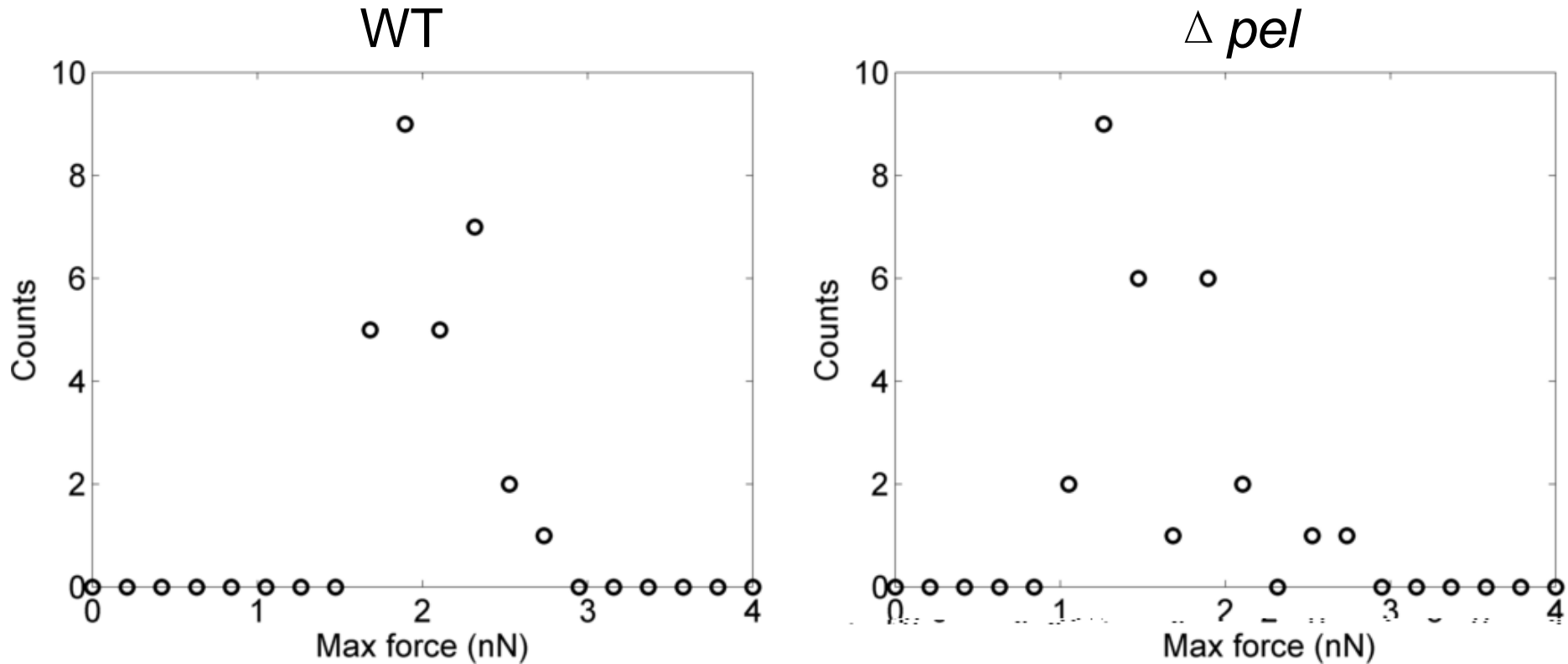
# Our method: bacteria attached to tip



# AFM measurements support inferred roles of EPS

- First time to

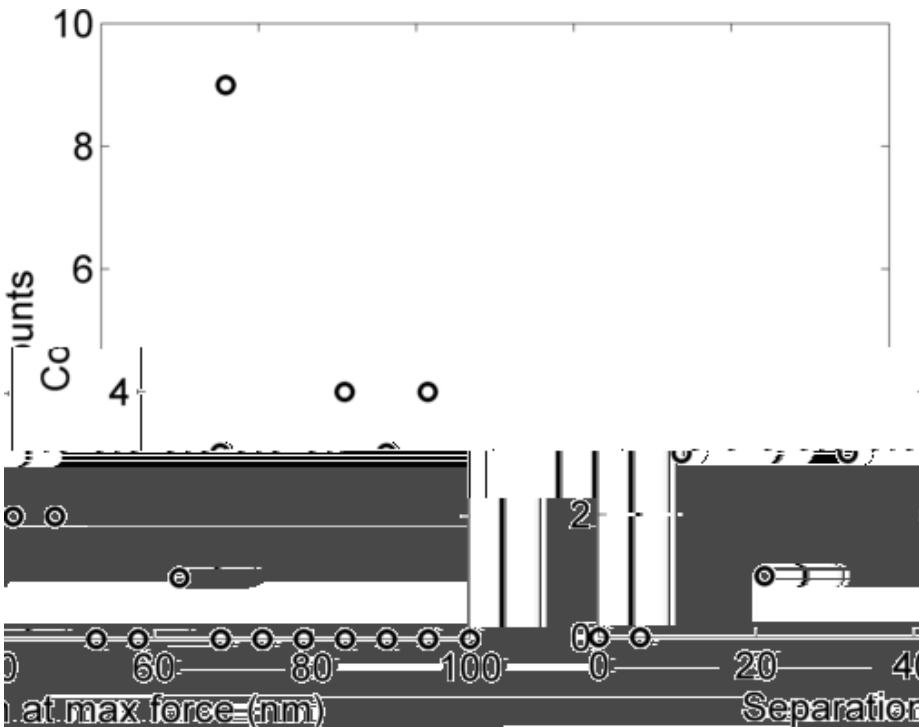
# Peak force measurements



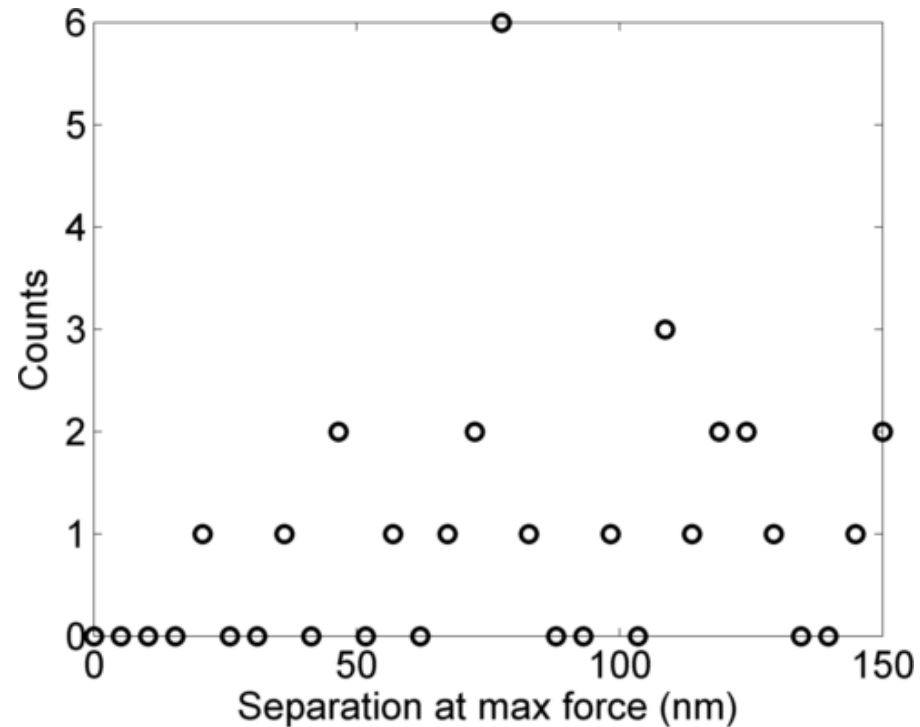
Pel contributes about 25% of the maximum adhesion force.

# Peak force location measurements

WT



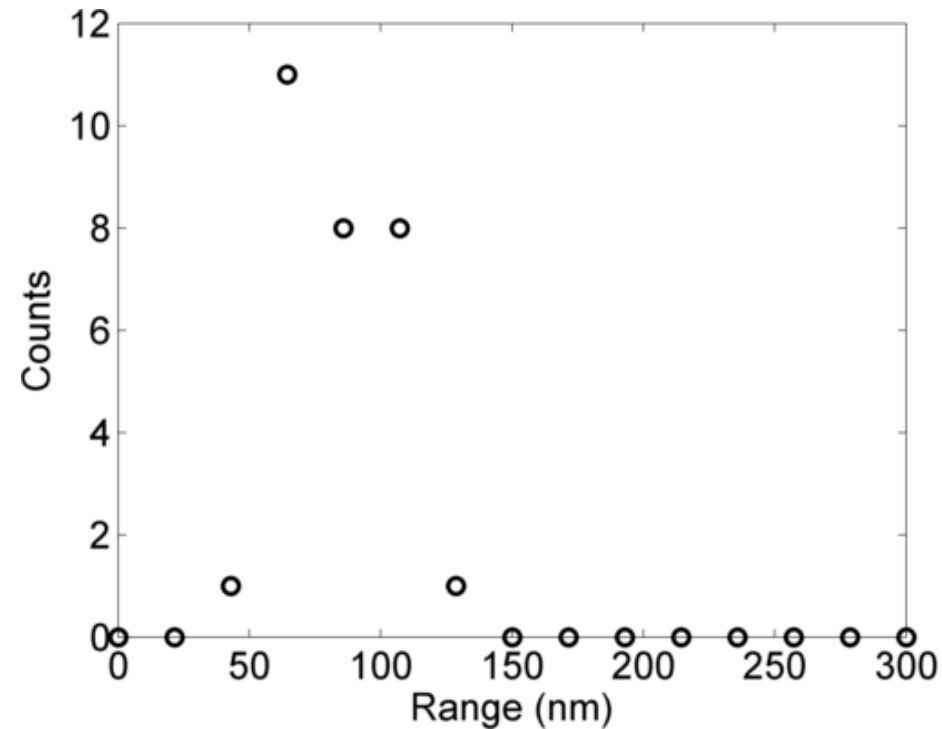
$\Delta pel$



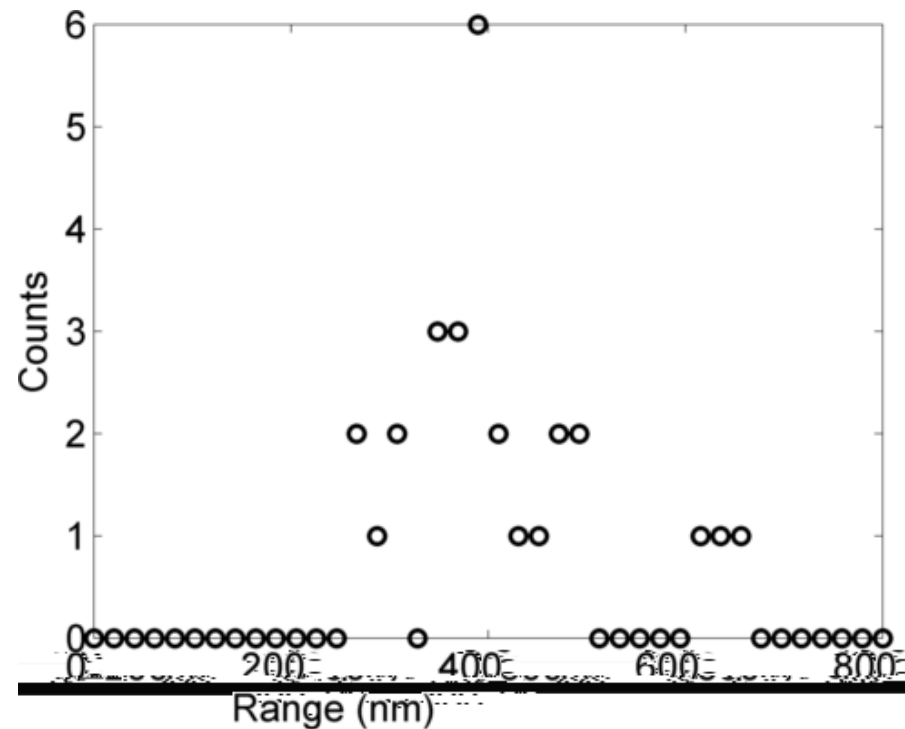
Pel makes the maximum adhesion force location ~4x more short ranged.

# Force range measurements

WT



$\Delta pel$



Pel decreases the extent of the adhesion force ~4x.

# What we've learned:

- Pel helps mediate the lying down associated with irreversible attachment
  - Pel symmetrizes bacterial attachment to surfaces
- Quantitative measurements of EPS mediated adhesion force.
  - Pel makes adhesion short ranged.
- (Implicit: Psl mediates non symmetric attachment – why?)



# Summary

- Bacterial biofilms are important medically, and good model systems for multicellularity.
- Distinct surface motility modes allow bacteria to explore space differently.
- Specific molecular glues mediate surface attachment and intercellular cohesion in distinct ways.

# Acknowledgements

ashmi

Wong

m Gibansky

E DAME  
a Shrout

ERSITY OF HOUSTON  
a Conrad

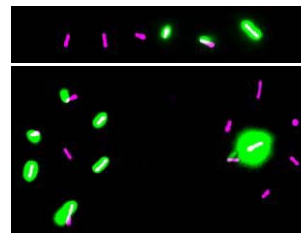
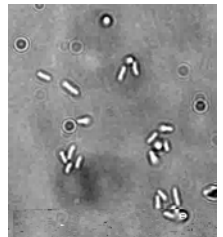
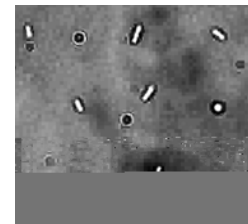
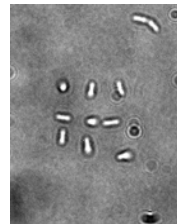
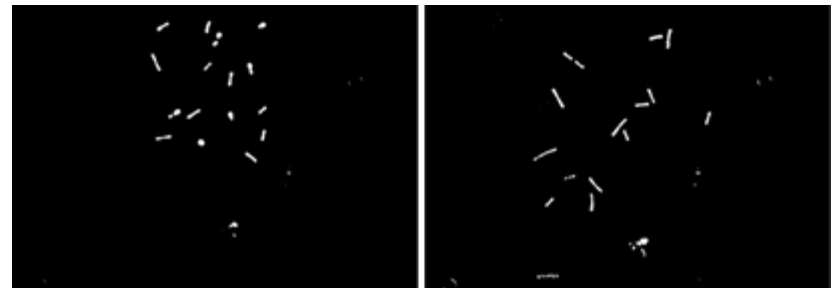
ERSITY OF WASHINGTON  
ew Parsek  
y Borlee  
olvin

FIBROSIS FOUNDATION

# Advertisement

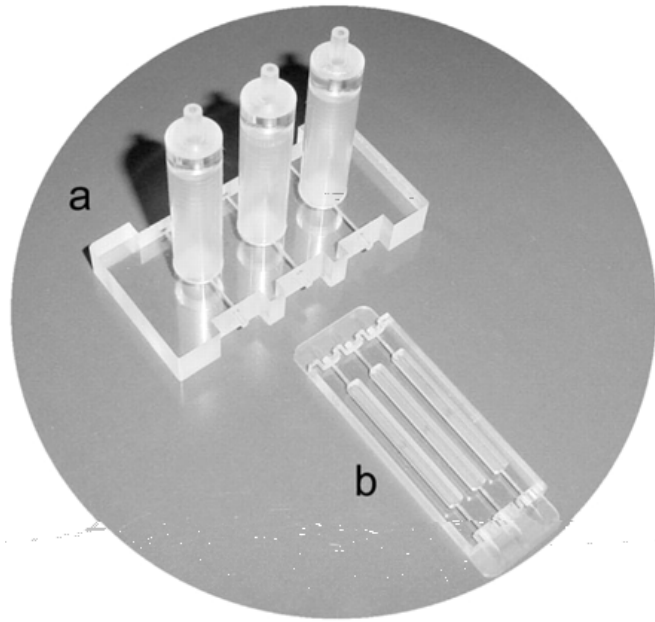
- **Postdoc** to work on a bacteria experiment: how does spatial structure develop in biofilms, and how does this impact cooperation?
  - This 4 investigator collaboration is funded by the Human Frontiers Science Project and is a great opportunity to train across disciplines.

- [gordon@chaos.utexas.edu](mailto:gordon@chaos.utexas.edu)

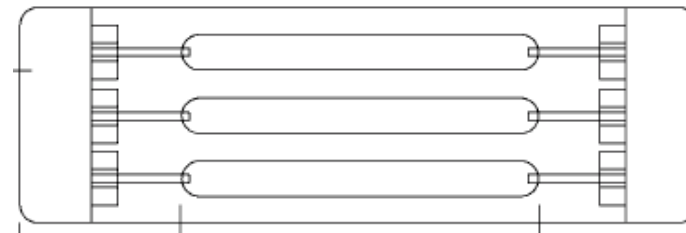


# Flow cell experiment

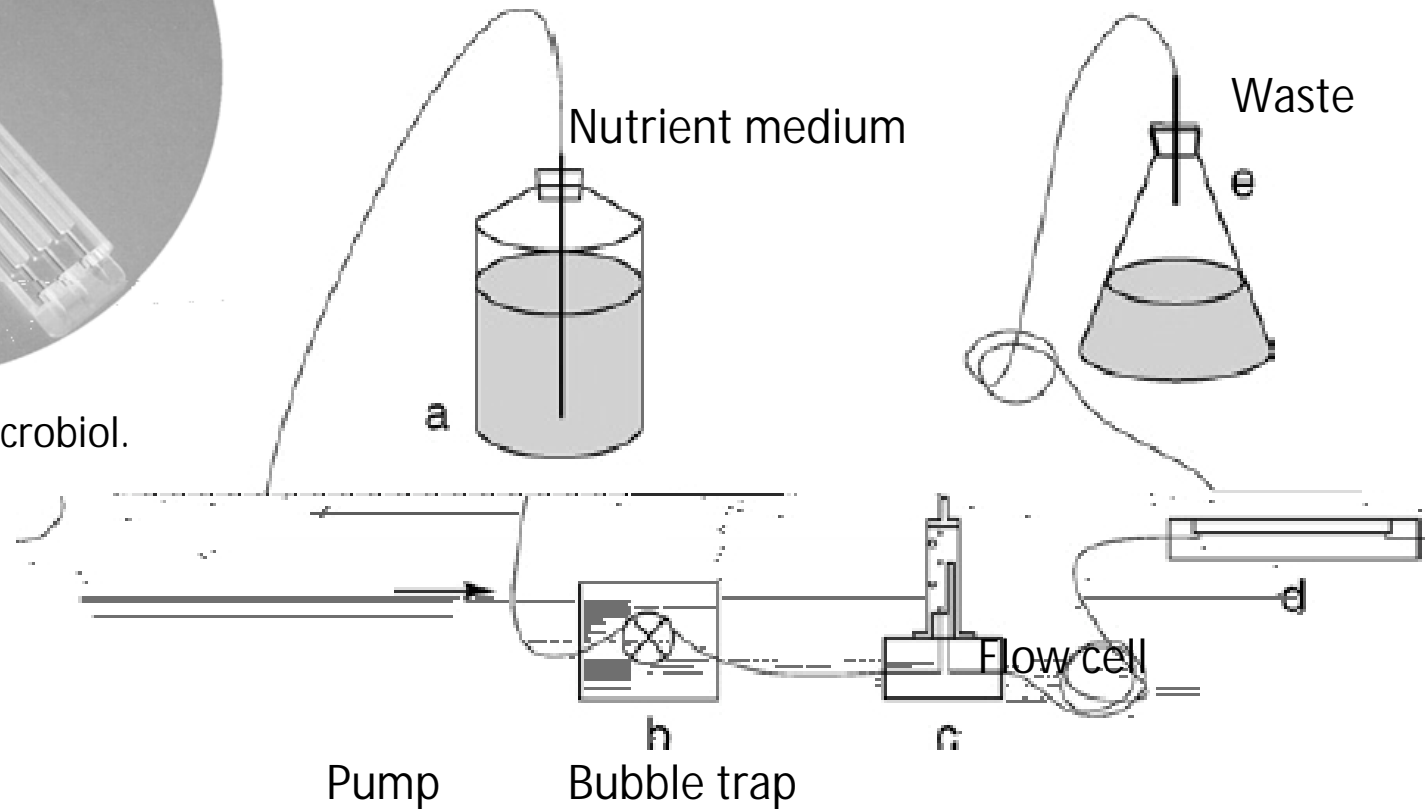
- Static sample chamber useful, but time limited
- Flow cell provides constant nutrient and oxygen supply



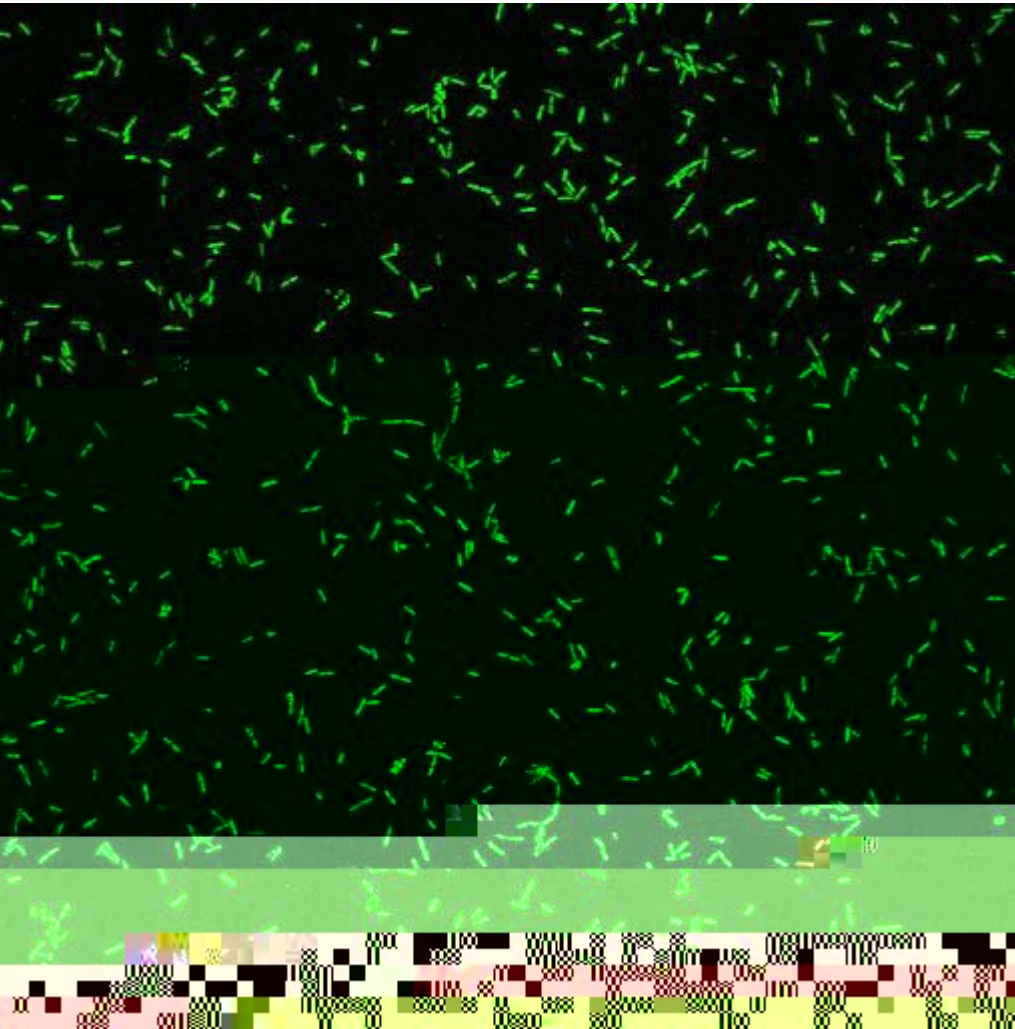
Stapper, ., J. Med. Microbiol.  
53(7): 679–690 (2004)



Sternberg & Tolker Nielsen,  
Curr. Protocols in Microbiol.  
1B.2.1–1B.2.15 (2005)

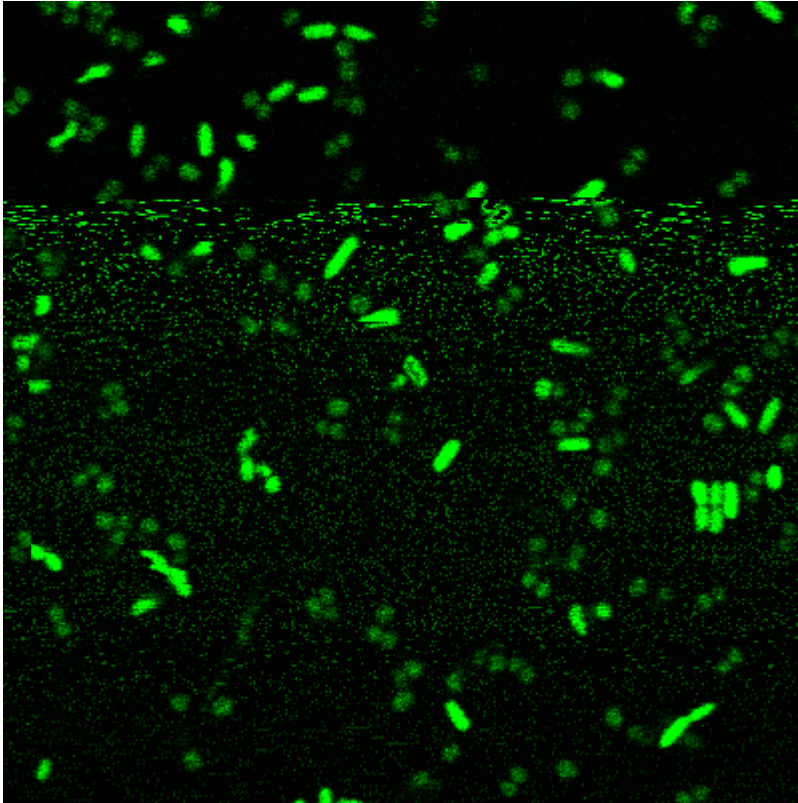


# Flow cell plans



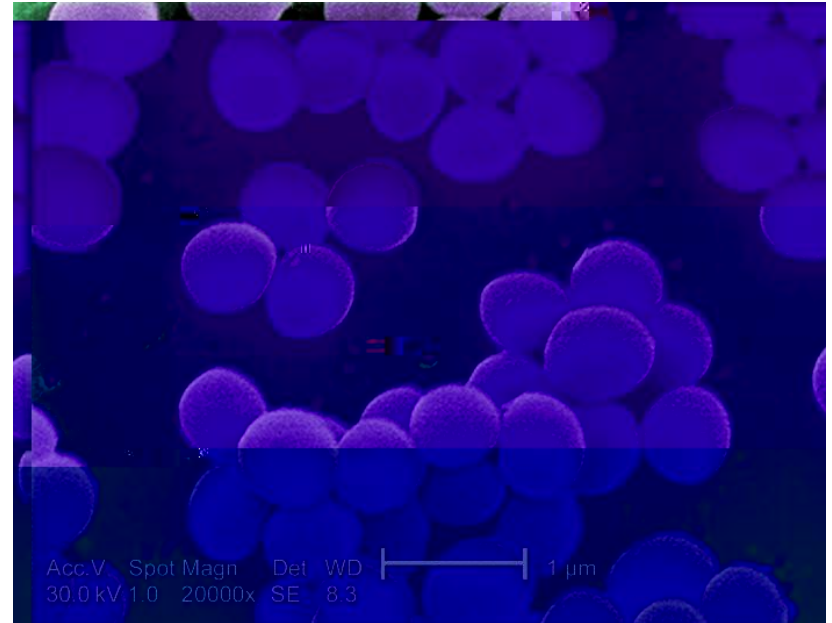
- Use confocal microscopy
- ~18 hour runs (not oxygen limited)
- Start with denser culture than static experiments
- Initial idea: look for similarities to colloid condensation transition
- New ideas and techniques

# coculture



CDC Public Health Image Library

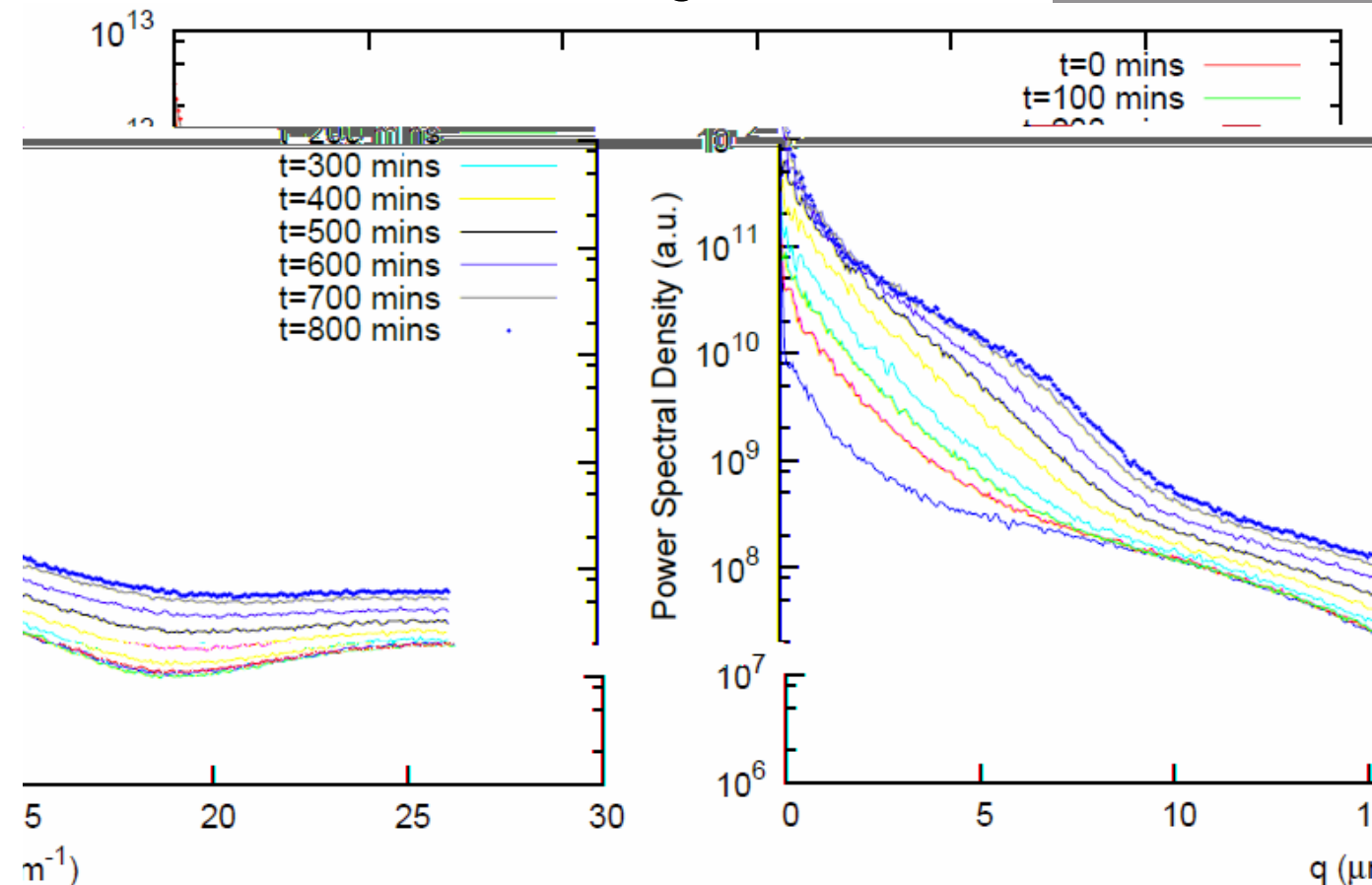
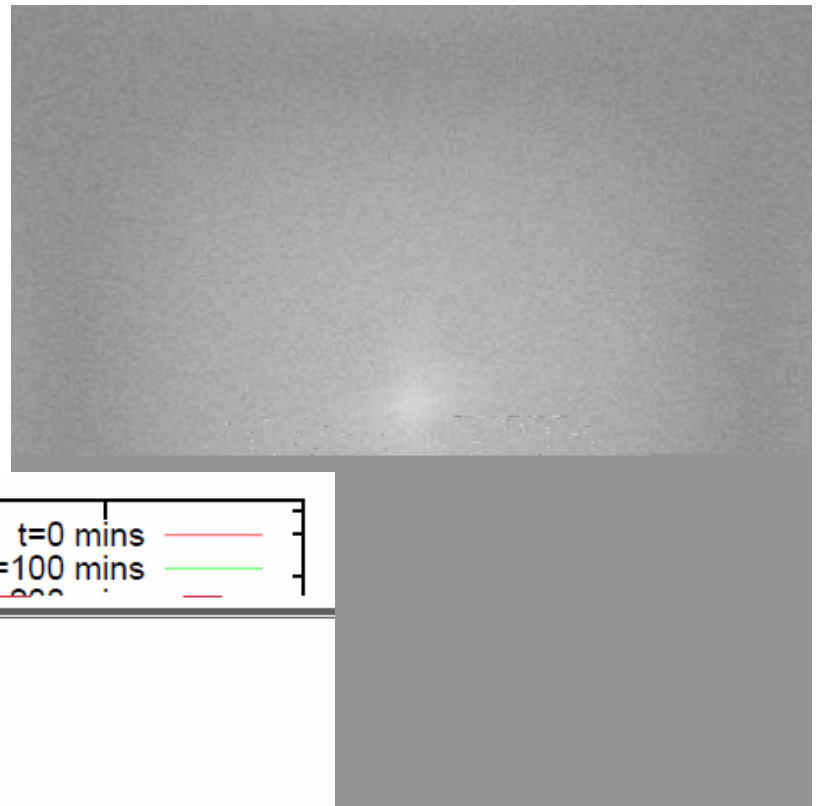
<http://phil.cdc.gov/phil/details.asp?pid=11157>



- *S. aureus* and *P. aeruginosa* both present in CF lung
- Evidence that *P. aeruginosa* can lyse Staph for iron
  - Mashburn, et al. 187, 554–566 (2005)
- How does *P. aeruginosa* biofilm growth change in the presence of Staph?

# New analysis

- Power spectrum of each frame
- Azimuthal avg shows features related to cluster growth



With Laurence  
Wilson, Rowland  
Institute at Harvard

# Flow cell plans

- Testing strains for use in coculture experiments
- Learn to grow Staph.
- Work on analysis (old & new)



# Surprise #2: adhesion leads to faster growth

- Faster doubling on surface vs. liquid culture

