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The Role of Lipid Droplets in Host-pathogen Interactions of Intracellular and Extracellular Bacteria

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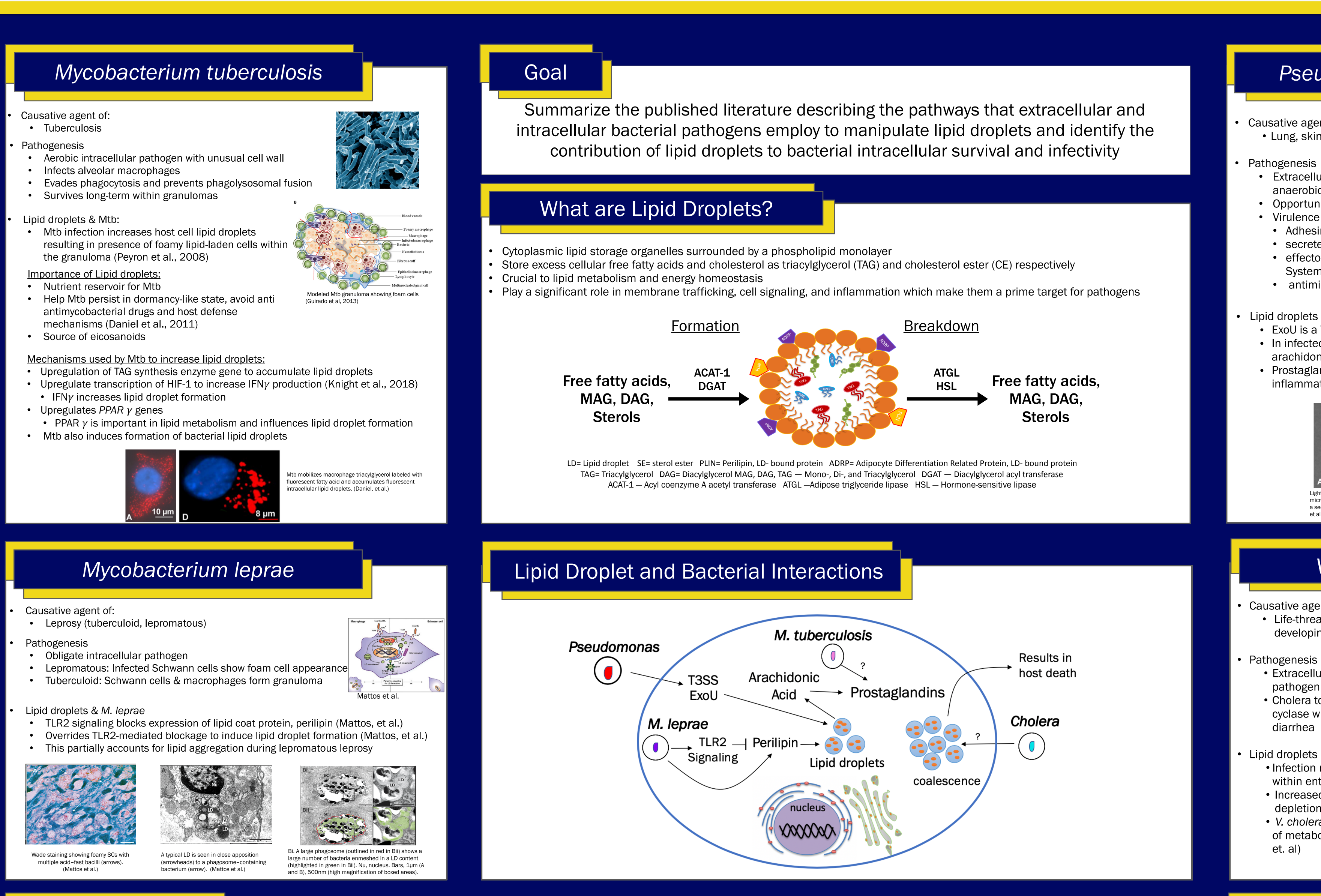
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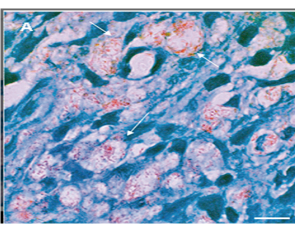
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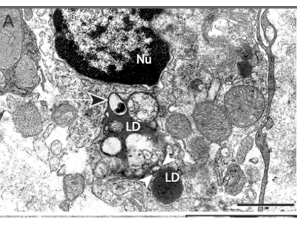


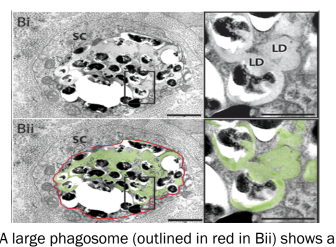
Indianapolis —— ®

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Conclusion

- *M. tuberculosis* and *Pseudomonas* promote infection in part by increasing prostaglandin production to increase inflammation Pseudomonas exoU and T3SS increase arachidonic acid, a precursor for prostaglandins
- *M. leprae* increases LD formation and creates foam cells to increase infectivity
- *V. cholera* causes LD coalescence through a poorly understood mechanism

The Role of Lipid Droplets in Host-pathogen Interactions of Intracellular and Extracellular Bacteria

Adam McDevitt, Ahila, Minal Mulye

M. tuberculosis increases prostaglandins through a poorly understood mechanism

• Intracellular and extracellular pathogens utilize lipid droplets by varied mechanisms to promote their survival in the host.

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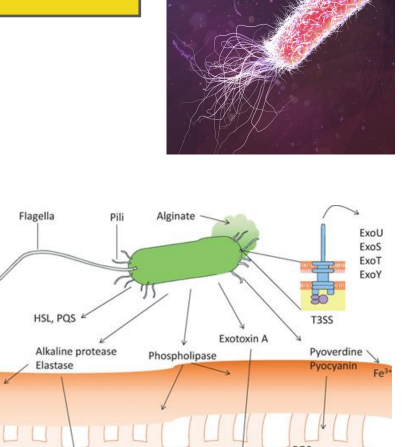
Pathogenesis

Pseudomonas aeruginosa

Causative agent of: • Lung, skin, urinary tract, ear and eye infection

• Extracellular, obligate aerobic or facultative anaerobic bacterium Opportunistic pathogen • Virulence factors:

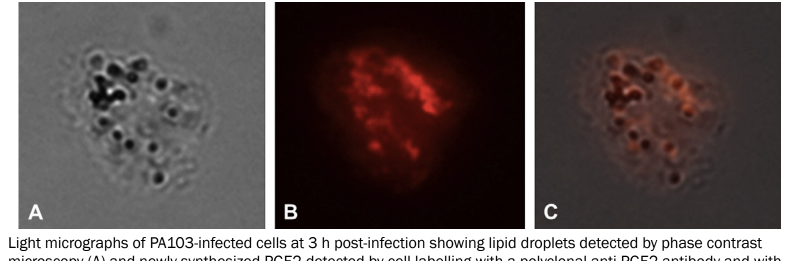
• Adhesins, secreted toxins, • effectors secreted via the Type 3 Secretion System (T3SS), • antimicrobial resistance



Gellatly et al

• Lipid droplets & P. aeruginosa

• ExoU is a T3SS effector which exhibits phospholipase activity • In infected cells, *P. aeruginosa* ExoU mobilizes lipid droplets to release arachidonic acids, a source of lipid immune mediator prostaglandin. • Prostaglandin is important in survival of *P. aeruginosa* by increasing inflammation. (Phillips et al.)



microscopy (A) and newly synthesized PGE2 detected by cell labelling with a polyclonal anti-PGE2 antibody and with a secondary antibody-Texas Red complex (B). Merged image shows localization of PGE2 in lipid bodies (C). (Phillips

Vibrio cholerae

• Causative agent of:

pathogen

diarrhea

et. al)

• Life-threatening diarrheal disease, particularly in developing countries

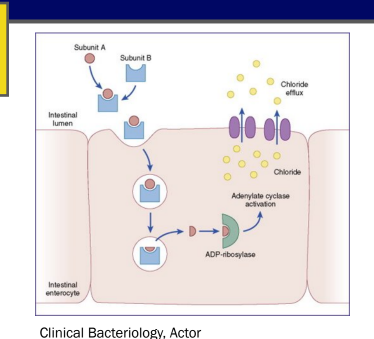
• Extracellular, gram-negative, facultative anaerobic

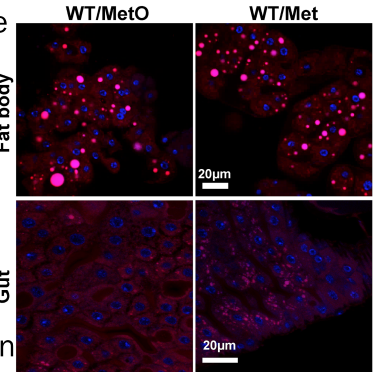
 Cholera toxin which continually stimulates adenylate cyclase within enterocytes \rightarrow profuse, watery

• Lipid droplets & V. Cholerae

• Infection results in coalescence of lipid droplets within enterocytes of small intestine in fruit flies • Increased lipid droplet coalescence results in depletion of lipids resulting in host death. • V. cholerae manipulates this process by degradation

of metabolic methionine sulfoxide (MetO) (Vanhove,





Nile red staining of neutral lipids in the fat body and ntestine of flies fed LB broth or LB broth inoculated vith wild-type V. cholerae(WT) supplemented with methionine (Met) or methionine sulfoxide (MetO) Vanhove, et al.)

The lipid droplet dream team: Adam McDevitt, Ahila, Cassie Libbing, Minal Mulye, Rea Azcueta