L02b

Recap L02a

Understanding structure and function, II

Interaction with the environment at the microscale

- Carbon sources

- Energy sources
- Defence and Offence

Capsules and Slime Layers

- Polysaccharide and glycoproteins outside the wall and outer membrane (different charge)
- Capsule poly-CHO extremely diverse in composition and structure (*E. coli* strains ~80)
- Can greatly increase effective cell volume
- May hold hydrolytic enzymes and scavenge metals and radionuclides
- May help attachment to surfaces; biofouling
-

Pili & Fimbriae

Pili & Fimbriae are thin (2–10 nm in diameter) filamentous structures made of proteins, pilin, that extend from cell surface

https://www.easybiologyclass.com/bacterialcell-surface-structures-and-appendagesflagella-fimbriae-and-pili/

Secretion systems, I

- Bacterial secretion is the process by which bacteria **release substances** to their **surroundings**, including **other cells**
- Bacteria achieve this using dedicated secretion systems that **transport molecules**
- Such as factors involved in **bacterial pathogenesis**, so called **effectors**
- Specialized macromolecular nanomachines that secrete a wide range of substrates, i**ncluding small molecules, proteins and DNA , important in host cell adherence as an initial step in colonization and pathogenesis**

Secretion systems, II

Specialized macromolecular

nanomachines that **secrete** a wide range of substrates, including small molecules, proteins and DNA , important in host cell adherence as an initial step in colonization and pathogenesis

All double-membrane-spanning secretion systems (T1SS, T3SS, T4SS and T6SS) use a one-step mechanism, such that substrates are transported directly from the bacterial cytoplasm into the extracellular space or into a target cell

T2SS, T5SS, Chaperon-usher and curli twostep translocases because they depend on either the Sec or Tat system; no ATP

T7SS, *Mycobacterium* (Gram positive)

None secretion systems is constitutively active

Hyp.: secretion may be triggered by the recognition of host receptors by specialized adhesion molecules called adhesins

Gunasinghe et al., 2017 Costa et al., 2015

- **• Resistance–nodulation–division (RND) pumps, antibiotics and small exogenous compounds**
- **• Type I secretion system (T1SS), ATPase (nutrient acquisition and virulence)**
- **• Type 2 secretion system (T2SS), Tat and Sec (folded and unfolded), ATPase (enzyme, toxins)**
- **• Type 3 secretion system (T3SS), effectors, form a pore in host, proteins to help the process**
- **• effectors modulate or subvert specific host cell functions, thereby promoting bacterial invasion and colonization**

- **• Type 4 secretion system (T4SS) mediates the translocation of DNA and proteins into bacterial or eukaryotic target cells**
- **• T4SS in Bacteria and Archaea, conjugation of plasmid DNA, ATP based**
- **• Type 6 secretion system (T6SS) cell envelopes panning machine that translocates toxic effector proteins into eukaryotic and prokaryotic cells**
- **• T6SS, injection mechanisms similar to phage**

- **• Type 5 secretion system (T5SS), chaperone-usher and curli OM only**
- **• Sec traslocase mediates proteins into periplasmic space**
- **• T5SS, substrate and its secretion pore are fused to form a single polypeptide —> a single polypeptide can drive its own secretion (autotransporter)**
- **• Chaperone-usher, used to assemble and secrete multisubunit appendages pili or fimbriae, —> mediate host cell recognition and attachment pathogenicity and biofilm formation**
- **• Curli biogenesis system, Curli are extracellular protein fibres, are functional amyloid**
- **• Curli protect bacteria from hostile environments by contributing to biofilm formation and facilitating interactions with the host immune system**

Self-assembling curli, by AFM

Amyloids A class of thread-like protein aggregates that self-assemble into insoluble toxic nanofibers. In bacteria, the accumulation of such fibers promotes the formation of a protective biofilm, whereas in humans they are involved in neurodegenerative diseases

Type IV pili in Gram negative & positive

T4P retraction brings

DNA to cell surface

 $Cytoplasm$ - Periplasm

structure formed by pilins

DNA binds to T4P or alternative

Type IV pili are **dynamic**: filaments polymerize and depolymerize, leading to rapid cycles of extension and retraction that generate considerable mechanical force

Type IV pili

- 1. Pull adherent bacteria along mucosal surfaces into close association with host cells and other bacteria
- 2. Exert forces on host cells
- 3. Pull bound substrates like DNA and bacteriophages into the periplasm
- 4. Export exoproteins across the outer membrane

Type IV pili extend and retract at rates of \sim 1,000 subunits per second, requiring a complex protein machinery that spans both membranes of Gram-

Chaperone-dependent

DNA uptake

Pili as conducting nanowires

- *Geobacter sulfurreducens* **Pili are made of proteins**
	- Pili form networks
	- γ-Proteobacteria, Cyanobacteria,

Methanogens

- Reduction of ferric oxide by touching via pili
- Cytochromes are involved in electron conduction
- **Strictly anaerobic conditions/ low O2**:
- A. growth medium supplemented with fumarate (40 mM) as the electron acceptor and with acetate (10 mM) as the electron donor for G.s.
- B. Fe(III) citrate (50 mM) as the electron acceptor and lactate (20 mM) as the electron donor

Bacteria Flagella (gross structure)

- For motility; Not always present
- 20 nm, hollow, very long (10-20 body lengths)
- Single protein, flagellin
- If broken they can regenerate
- Flagellar arrangement; polar, peritrichous
- Wavy; wavelength constant for a species
- Rigid, do not make wave-like motion like sperm

Bacteria Flagella (gross structure)

- Grow at free end
- Self assembly from transported flagellin

- Microenvironment viscosity modulate expression of swarmer cells (many lateral flagella)
- *Vibrio*s have sheathed polar flagellum

– Periplasmic flagella in *Borrelia burgdorferi*

Filament

Kim et al. 2017

Bacteria Flagella (ultra-structure and function)

- Filament
- Hook
- Basal body (rod and rings)
	- Filament rotates at base like propeller
	- Basal body acts like a motor
	- Most studied marine bacteria have H+/Na+ driven flagella motors
	- 1700 rps/400 μ m s⁻¹

Proton turbine model of flagella movement 17

Bacteria decision-making system for motility

Gliding Motility

- Movement of cytoplasmic membrane proteins (gliding protein)is driven by proton motive force
- This somehow transmits energy to move to outer membrane proteins
- This (hypothetically) pulls the cell against a **solid surface**

Archaea Flagella (ultra-structure and function)

- \sim half the diameter of flagella, measuring about 10–13 nm in width
- Archaellum can be considered a rotating type IV pilus capable of both CW and CCW rotation
- In flagellum, whose energy requirement is met by **dissipation of the proton motive force**, **achaellum**'s rotation is driven by **ATP hydrolysis**
- In Bacteria a single type of protein makes up the filament
- In Archaea several different filament proteins

Prokaryotic cytoskeleton

- Similar to eukaryotic cytoskeletal actin, tubulin and intermediate filaments
- Cytoskeletal proteins may dictate shape during peptidoglycan synthesis

Proteins involved

- **1.** *FtsZ:* **Cell division protein; related to tubulin; assembles as ring at cell division site, recruits other proteins to form contractile septal ring constricting cell during cell division**
- 2.*MreB:* Member of actin superfamily; in rod-shape, filamentous and helical bacteria; encircles cell as spiral under cell membrane along longitudinal axis contributing to shape of non-spherical bacteria
- *Crescentin:* In *Caulobacter crescentus*; gives vibroid shape; helical filament along cell membrane
- *Spiroplasma:* Move in viscous media; single protein ribbon, stretch & release by conformational change
- *Mycoplasma*: Some motile on solid surface; internal cytoskeletal fibers for movement and attachment

Intracytoplasmic membrane (ICM), I

- ICM as a strategy to concentrate and stabilize functions within the cell
- ICM promote more efficient reactions
- ICM is a solution to the fight for diffusion
- **• Phototrophs: site of photosynthetic apparatus**
- **• Cyanobacteria**

Tomographic reconstruction of a Cyanothece 51142 cell

T, Thylakoid membrane; C, carboxysome; G, glycogen granule; P, polyphosphate body

Blue gray, Plasma membrane (rendered partially transparent for clarity); white, lipid bodies; blue, carboxysomes; green, thylakoid membranes; gray, polyphosphate bodies

Bar = 1,000 nm

Intracytoplasmic membrane (ICM), II

Methylobacterium organophilum

Patt & Hanson 1978

Nitrifk et al., 2004

Fig. 4. Postulated anaerobic ammonium oxidation coupled to the anammoxosome membrane in anammox bacteria resulting in a proton motive force and subsequent ATP synthesis via membrane-bound ATPases. HH: hydrazine hydrolase; the hydrazine-forming enzyme, HZO: hydrazine-oxidizing enzyme, NIR: nitrite-reducing enzyme.

ICM present in methanotrophs, N_2 fixers, nitrifiers and phototrophs *(see also: magnetosomes, gas vacoule, minicompartments, anammoxosome)*

- *Methanotrophs*: ICM is the site of methane oxidation
- $N₂$ fixers: Increases respiratory activity to provide ATP for $N₂$ fixation and remove $O₂$ near nitrogenase
- *Nitrifiers*: Site of enzymes catalyzing ammonia and nitrate oxidation

Gas Vacuoles

- •In aquatic bacteria; for buoyancy
- •Means of motility (float up and down)
- •In Cyanobacteria, some purple and green phototrophic bacteria, some Archaea
- •Spindle shaped hollow, rigid, 300-1000 nm, few to hundreds per cell
- •Membrane only protein (rigid), 2 nm thick, impermeable to water, permeable to gases
-

Storage products

Common storage products- enclosed in thin lipid layer, also in periplasm:

- Poly-ß-hydroxybutyrate (PHB); sulfur globules in S oxidizing bacteria and archaea
- Glycogen
- **Polyphosphate**
- Nitrogen store?
- Carboxysomes (RuBP carboxylase in cyanobacteria); crystal-like, 120 nm; membrane bound
- Mobilized when needed
- Advantageous in nutritionally fluctuating environments

Magnetosomes

- **Cytoplasmic membranes** forms an internal invagination and **recruit proteins for Fe precipitation**
- Magnetotactic bacteria usually mineralize either **iron oxide** magnetosomes, which contain crystals of magnetite (Fe3O4), or **iron sulphide** magnetosomes, which contain crystals of greigite (Fe3S4)
- Mainly in aquatic bacteria; some algae
- Morphology species-specific
- **Permanent magnetic dipole to the cell, for N-S** *orientation* in environment ("**magnetotaxis-aerotaxis**") efficient swimming, passive cell alignment to geomagnetic field lines
- Microaerophilic bacteria may use them to **stay in low oxygen at oxic/anoxic interface**
-

Bacterial microcompartments

Bobik, 2007 Chowdhury et la., 2014

- Sophisticated protein-based organelles used to **optimize** specific metabolic pathways
- **• Metabolic enzymes encapsulated within a protein shell, increase specificity and yield**
- Widely distributed and functionally diverse
- **• Compartmentalization creates an ideal environment for catalysis and facilitates the channeling of toxic/volatile intermediates to downstream enzymes**
- Structurally resemble viral capsids
- Carboxysome channels CO₂
- Pdu microcompartment channels propionaldehyde, *Salmonella enterica*
- Eut microcompartment channels acetaldehyde

Bacterial microcompartments: carboxysome

RuBisCO is the CO2-fixing enzyme of the Calvin-Benson-Basham Cycle, CBB

- 1. RuBisCO catalyses the conversion of $CO₂$ and ribulose bisphosphate into two 3phosphoglycerate
- 2. RuBisCO reacts with O2 in a nonproductive process known as **photorespiration**, which can drain away up to 50% of the carbon fixed by the CBB cycle—> competition with carboxylation
- 3. The carboxysome is essential part of a **carbon dioxide concentrating mechanism** (CCM) that improves the efficiency of CO₂ fixation by RuBisCO
- 4. CCM starts with the concentration of $HCO₃$ in the cytoplasm of the cell by active transport
- 5. Equilibrium with $CO₂$ is not reached due to a lack of carbonic anhydrase (CA)
- 6. Carboxysomal CA converts HCO_3 to CO_2 and releases it within the microcompartment
- 7. The protein shell of the microcompartment impedes $CO₂$ diffusion
- **8. Consequently, CO2 is concentrated in RuBisCO vicinity —> increase in CO2 fixation and suppression of photorespiration**

Spores

Ca2+ crosslinking dipicolinic acid

•Differentiated cells within bacteria (endo-) •Very resistant (heat, desiccation, chemicals); persist long •Gram Positive (e.g. Bacillus, Clostridium)

•LM: Seen as refractile bodies; TEM structure, very different from vegetative cell:

•Many layers: thin (protein) exosporium; spore coat (protein) cortex (PG), core (protoplast)

•Characteristic: Dipicolinic acid (in core); high Ca²⁺ (most in Ca-DPA) ~10 w/w endospore

•Core: 10-30% water; Ca-DPA; cytoplasm is a gel; enzyme inactive; lower pH; SASPs (small Acid soluble proteins)- bind and protect DNA & C/e source during outgrowth the contract of the \sim 29 $\,$

Endospore formation

TABLE 2.2 Differences between endospores and vegetative cells

- Initiated in response to nutrient stress
- Many steps; in B. subtilis 8h; ~200 genes
- Activation of a number of spore specific genes (spo, ssp; encoding SASPs)
- Germination: Activation, germination, outgrowth
- Ca-DPA and cortex lost; SASPs degraded; spore swells (takes in water); new RNA, protein, DNA synthesis; cell emerges from the broken coat 30

Periplasm-periplasmic space

Gram-negative cell wall

a Gram-negative bacteria

- Active metabolic site (reduced in Gram-positive)
- Very viscous —> high concentration of extracellular proteins (via cytoplasmatic protein-exporting system)
- Outer membrane is impermeable to proteins and very large molecules—> prevents extracellular proteins from diffusing away from the cell
- Width of periplasmic space 15 nm
- Major periplasmic proteins:
	- Hydrolytic enzymes (degradation of polymeric substances)
	- Binding proteins (transporting substrates)
	- Chemoreceptors (Chemotaxis response)
	- Structural proteins (peptidoglycan, outer & cytosolic membrane)

Nucleoid

Madigan et al. 2018 **Madigan et al. 2018**

- No nuclear membrane
- Naked DNA
- Generally 2 4 x 10⁹ Daltons (but Vibrios--2 chromosomes; Myxobacteria)
- One mm long (1000 5000 body lengths) supercoiled (histone-like proteins)
- Plasmid, common carrier of antibiotic resistance and metal resistance genes
- *E. coli* nucleoid occupies 0.07 µm3
- In diverse environments the degree of supercoiling is different and coupled with surface/volume ratio

Ribosomes

- • **Fill cytoplasm in fast growing cells (20 000 cell-1 in rapidly growing** *E.coli)*
- **Number depends on the physiological state of the cell**
- • **0.02 µm diameter**
- • **Site of protein synthesis**
- • **Simpler body plan of procaryotes allows simultaneous transcription and translation as well as fine regulation of protein synthesis**
- • **Perhaps only 20 -100 ribosomes in marine bacteria with growth rate <1 d**
- • **RNAs are structural molecules**
-

Protein

Water

- 55% of dry weight in *E. coli* and in marine bacteria
- Together with DNA makes the cytoplasm a thick gel
- Occupies 13% of the cell volume in *E. coli*, but up to 50% in marine bacteria
- L and D aa
- ~90% of the cell volume in *E. coli*
- ~50-90% of cell volume in marine/aquatic bacteria depending on the cell size
- Dryness may be an adaptation for rapid response to nutrient supply
- Just add water to become bigger

Adaptive strategies of Bacteria for nutrient uptake

Azam and Malfatti, 2007 Nature Reviews Microbiology 10:782

- • **Motility, environmental sensing, permeases and cell-surface hydrolases**
- • **Adapted fine biochemical strategies to interact with organic matter natural and human-created**

Transport Mechanisms

- **• Passive diffusion**
- **• Facilitated diffusion**
- **• Active transport**
	- **• Energy coupling for active transport**
	- **• Primary and secondary active transport**
	- **• Binding protein dependent active transport**
- **• Group translocation**

Microbial adaptations to increase uptake of molecules

Microbial interfaces, the membranes as hotspots of activities

External concentration of solute

Figure 2.8 The importance of transport in membrane function. In transport, the uptake rate shows saturation at relatively low external concentrations. Both high-affinity and low-affinity transport systems are depicted.

Passive Diffusion

Concentration dependence of uptake by passive diffusion

- Net flux only until $C_{inside} = C_{outside}$ (no accumulation)
- No metabolic energy required
- No specific interaction with cell membrane component
- If environment, $C_{\text{outside}} \ll C_{\text{inside}}$, not useful for nutrient uptake
- Used for the uptake of O_2 , CO_2 and H_2O
- Through the phospholipid bilayers, small and non-polar

Facilitated Diffusion

- **• Large or polar molecules**
- No metabolic energy required
- Substrate specific interaction with a **membrane permease or carrier**
- **Stereospecific** (D and L amino acids completely discriminated)
- Substrate binds to carrier outside cell and is released inside the cell
- Not effective in dilute solutions unless C_i is kept low by utilization

Active Transport

- **Accumulation against a concentration gradient**
- **Requires metabolic energy**
- Carrier-substrate complex formed outside of the membrane
- Structural specificity and stereospecificity
-

Uniporters, symporters and antiporters

Uniporter: Cause unidirectional transport (through membrane spanning protein)

Symporter: Transport substrate along with H+ (or Na+)

Antiporter: Substrate and H+ (or Na+) transported in opposite directions

[Require PMF, Proton Motive Force]

Simple Transport

Periplasmic space

Cytoplasm

Transport is linked to dissipation of the proton motive force (PMF)

H+ goes into cell

Same or opposite direction

Transmembrane transporters are composed of a polypeptide that forms 12 a-helices —> a channel

PMF is generated by e- transport, **H+ ions are extruded to the outer surface of the membrane**

Inside of cell has net - charge and outside net +

ABC Transporters

• ABC (ATP Binding Cassette) systems: Gram- (periplasm); G+ & Archaea BP membrane-anchored •Used for some amino acids, peptides, sugars, organic acids, sulfate, other ions

•Requires metabolic energy as **ATP** *(not PMF*)

•Mechanism:

–Substrate binds to a high-affinity binding protein in periplasm

–Complex interacts with a membrane-embedded multimeric carrier

–ATP hydrolysis changes subunit interactions; creates a transport channel

•Multiple systems w/different K_m , V_{max} (3 for glucose in E . *coli*; scavenging system)

Periplasmic space **Periplasmic space**

Group Translocation

(e.g. Phosphoenolpyruvate-PEP- phosphotransferase system)

- **Substrate modified** during transport —> phosphorylated
- **• Energy derived from metabolic compound PEP (glycolysis)**
- PEP donates \sim P for phosphorylation (PEP PTS system)
- **Sugar-phosphate is 'trapped**' (membrane is impermeable to it)
- Conserves energy. Transport and phosphorylation with a single \sim P
- EII are sugar specific; EIIb lies @inner membrane face; EIIc: integral
- Examples: sugars (glucose, fructose, mannose), NAG
- **• Generally found in facultative anaerobes and anaerobes**