

Why study bacterial genetics?

- Its an easy place to start
 - ◆ history
 - ◆ we know more about it
 - systems better understood
 - ◆ **simpler genome**
 - ◆ good model for control of genes
 - build concepts from there to eukaryotes
 - ◆ **bacterial genetic systems are exploited in biotechnology**



Bacteria

- Bacteria review
 - ◆ one-celled organisms
 - ◆ **prokaryotes**
 - ◆ reproduce by splitting
 - binary fission
 - ◆ **rapid growth**
 - generation every ~20 minutes
 - 10⁸ (100 million) colony overnight!
 - ◆ dominant form of life on Earth
 - ◆ incredibly diverse

Bacteria as Pathogens

- Disease-causing microbes
 - ◆ plant diseases
 - wilts, fruit rot, blights
 - ◆ animal diseases
 - tooth decay, ulcers
 - anthrax, botulism
 - plague, leprosy, "flesh-eating" disease
 - STDs: gonorrhea, chlamydia
 - typhoid, cholera
 - TB, pneumonia
 - lyme disease



Bacteria as Beneficial (& necessary)

- Life on Earth is dependent on bacteria
 - ◆ **decomposers**
 - recycling of nutrients from dead to living
 - ◆ **nitrogen fixation**
 - only organisms that can fix N from atmosphere
 - ◆ needed for synthesis of proteins & nucleic acids
 - ◆ plant root nodules
 - ◆ help in digestion (E. coli)
 - digest cellulose for herbivores
 - ◆ cellulase enzyme
 - produce vitamins K & B₁₂ for humans
 - ◆ produce foods & medicines
 - from yogurt to insulin

Bacterial Diversity

Borrelia burgdorferi
Lyme disease

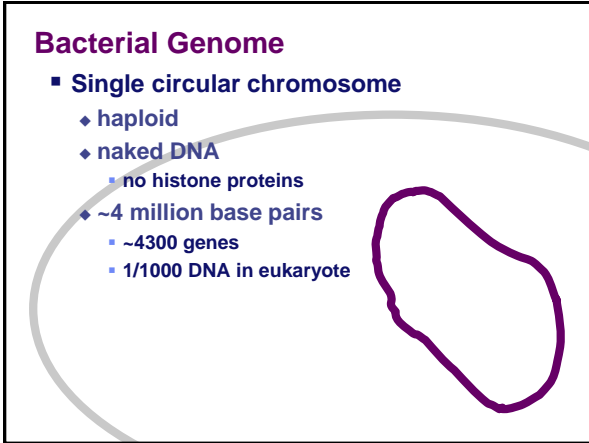
Treponema pallidum
Syphilis

Escherichia coli O157:H7
Hemorrhagic E. coli

Enterococcus faecium
skin infections

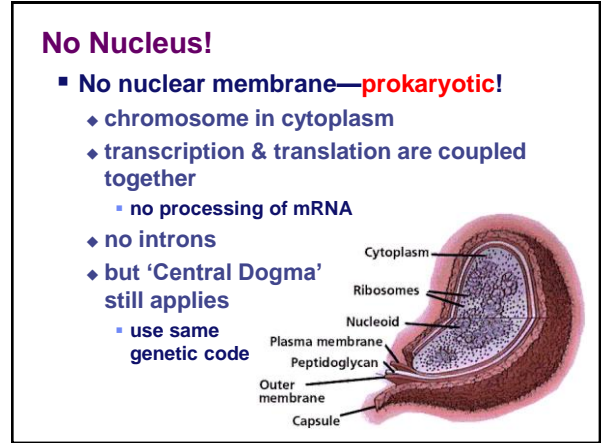
Bacterial Genome

- Single circular chromosome
 - haploid
 - naked DNA
 - no histone proteins
 - ~4 million base pairs
 - ~4300 genes
 - 1/1000 DNA in eukaryote



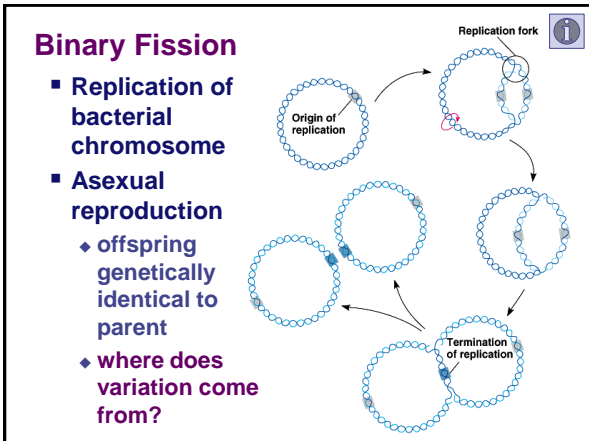
No Nucleus!

- No nuclear membrane—**prokaryotic!**
 - chromosome in cytoplasm
 - transcription & translation are coupled together
 - no processing of mRNA
- no introns
- but 'Central Dogma' still applies
 - use same genetic code



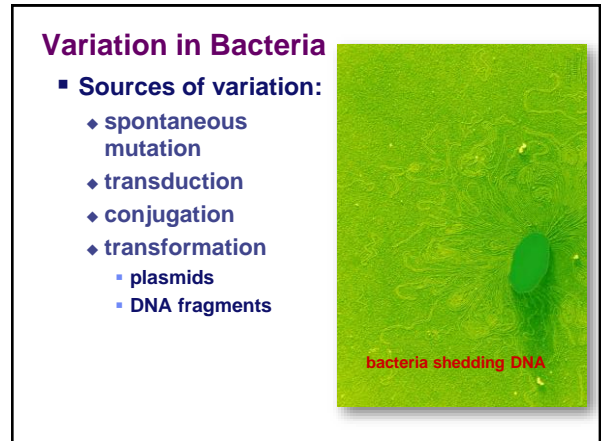
Binary Fission

- Replication of bacterial chromosome
- Asexual reproduction
 - offspring genetically identical to parent
 - where does variation come from?



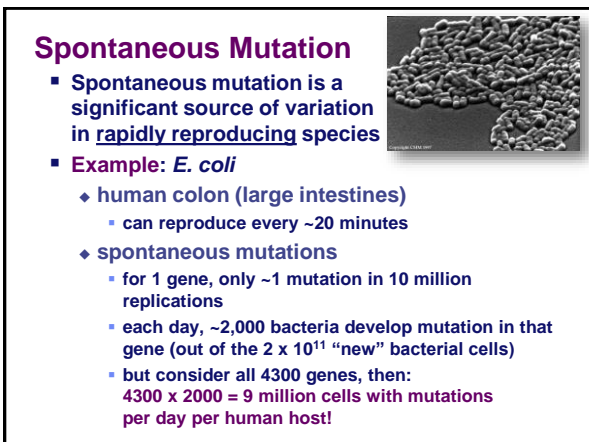
Variation in Bacteria

- Sources of variation:
 - spontaneous mutation
 - transduction
 - conjugation
 - transformation
 - plasmids
 - DNA fragments



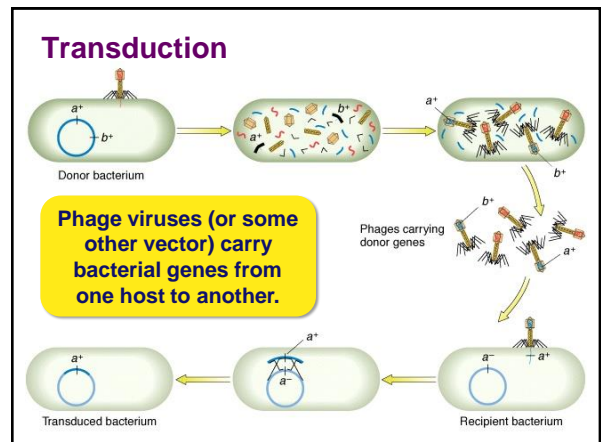
Spontaneous Mutation

- Spontaneous mutation is a significant source of variation in rapidly reproducing species
- Example: *E. coli*
 - human colon (large intestines)
 - can reproduce every ~20 minutes
 - spontaneous mutations
 - for 1 gene, only ~1 mutation in 10 million replications
 - each day, ~2,000 bacteria develop mutation in that gene (out of the 2×10^{11} "new" bacterial cells)
 - but consider all 4300 genes, then: $4300 \times 2000 = 9$ million cells with mutations per day per human host!



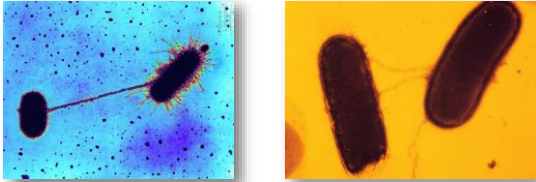
Transduction

Phage viruses (or some other vector) carry bacterial genes from one host to another.



Conjugation

- Direct transfer of DNA between 2 bacterial cells that are temporarily joined
 - ◆ results from presence of F plasmid with F factor
 - F for "fertility" DNA
 - ◆ E. coli "male" extends sex pilli, attaches to female bacterium
 - ◆ cytoplasmic bridge allows transfer of DNA

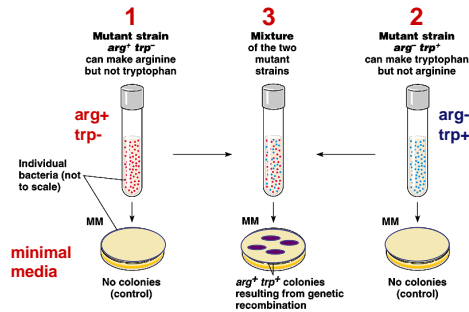


Transformation

- some bacteria are opportunists
 - ◆ pick up naked foreign DNA wherever it may be hanging out
 - have surface transport proteins that are specialized for the uptake of naked DNA
 - ◆ import bits of chromosomes from other bacteria
 - ◆ incorporate the DNA bits into their own chromosome
 - express new gene
 - form of recombination

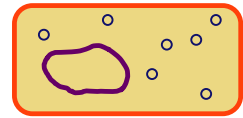
Swapping DNA

- Genetic recombination by trading DNA



Plasmids

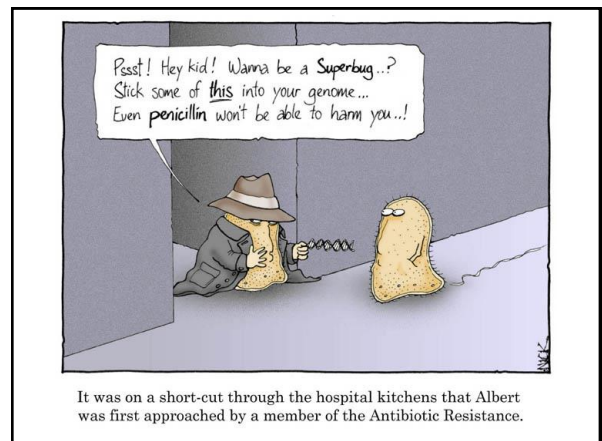
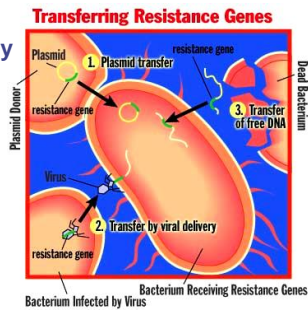
- Plasmids
 - ◆ small supplemental circles of DNA
 - 5000 - 20,000 base pairs
 - self-replicating
 - ◆ carry extra genes
 - 2-30 genes
 - ◆ can be exchanged between bacteria
 - rapid evolution
 - antibiotic resistance
 - ◆ can be imported from environment



Plasmids & Antibiotic Resistance

- Resistance is futile?

- ◆ 1st recognized in 1950s in Japan
- ◆ bacterial dysentery not responding to antibiotics
- ◆ worldwide problem now
 - resistant genes are on plasmids that are swapped between bacteria



Biotechnology

- Used to insert new genes into bacteria
 - example: pUC18
 - engineered plasmid used in biotech

antibiotic resistance gene on plasmid is used as a selective agent

Recombinant Plasmid

- Antibiotic resistance genes as a **selectable marker**
- Restriction sites for splicing in gene of interest

Selectable marker

- Plasmid has both "added" gene & antibiotic resistance gene
- If bacteria **don't** pick up plasmid then **die** on antibiotic plates
- If bacteria pick up plasmid then survive on antibiotic plates
- selecting for successful transformation**

Selection for Plasmid Uptake

- Ampicillin becomes a selecting agent
 - only bacteria with the plasmid will grow on **amp** plate

all bacteria grow only transformed bacteria grow

LB plate LB/amp plate

Copy DNA

- Plasmids
 - small, self-replicating circular DNA molecules
 - insert DNA sequence into plasmid
 - vector** = "vehicle" into organism
 - transformation**
 - insert **recombinant** plasmid into bacteria
 - bacteria make lots of copies of plasmid
 - grow recombinant bacteria on agar plate
 - clone of cells = lots of bacteria
 - production of many copies of inserted gene

DNA → RNA → protein → trait

Development of GFP 1961, 1994 | 2008

- Shimomura, Chalfie, Tsien
 - discovery, isolation, and purification of GFP and many fluorescent analogs

Osamu Shimomura Martin Chalfie Roger Tsien



