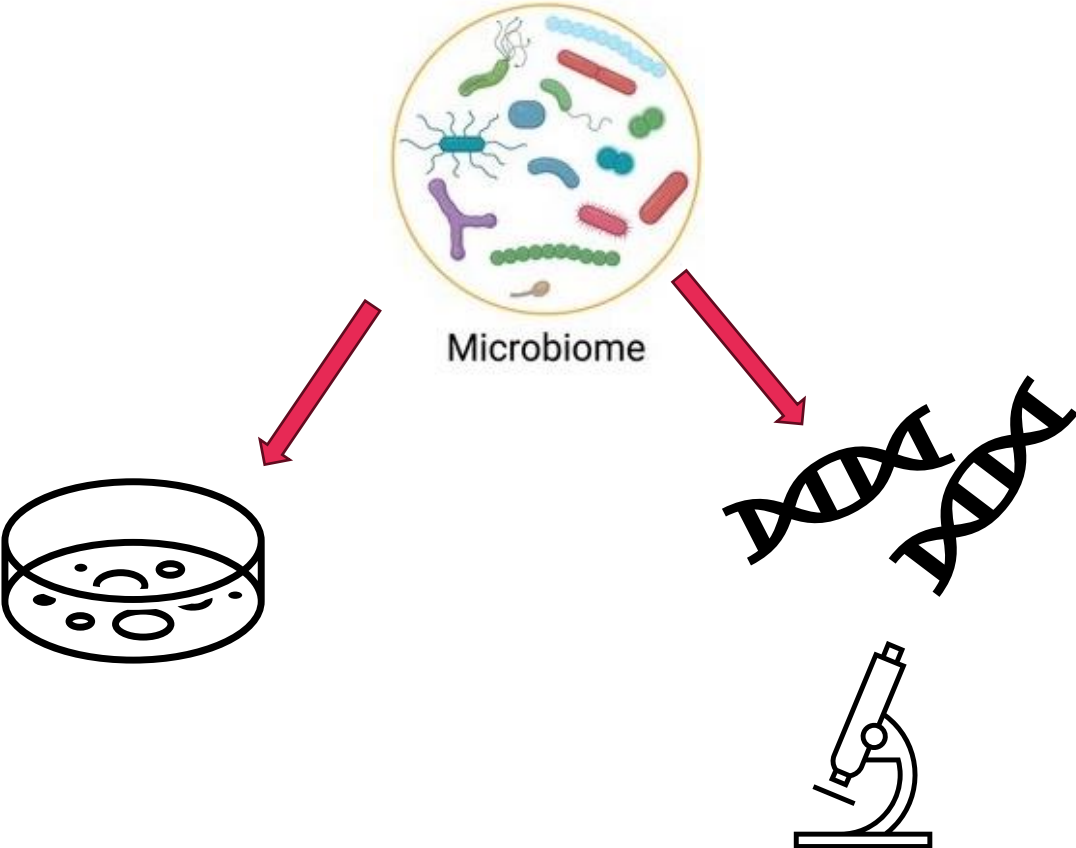




TAKING THE MEASURE OF MICROBIAL SYSTEMS

Microbial Genomics and Other Omics

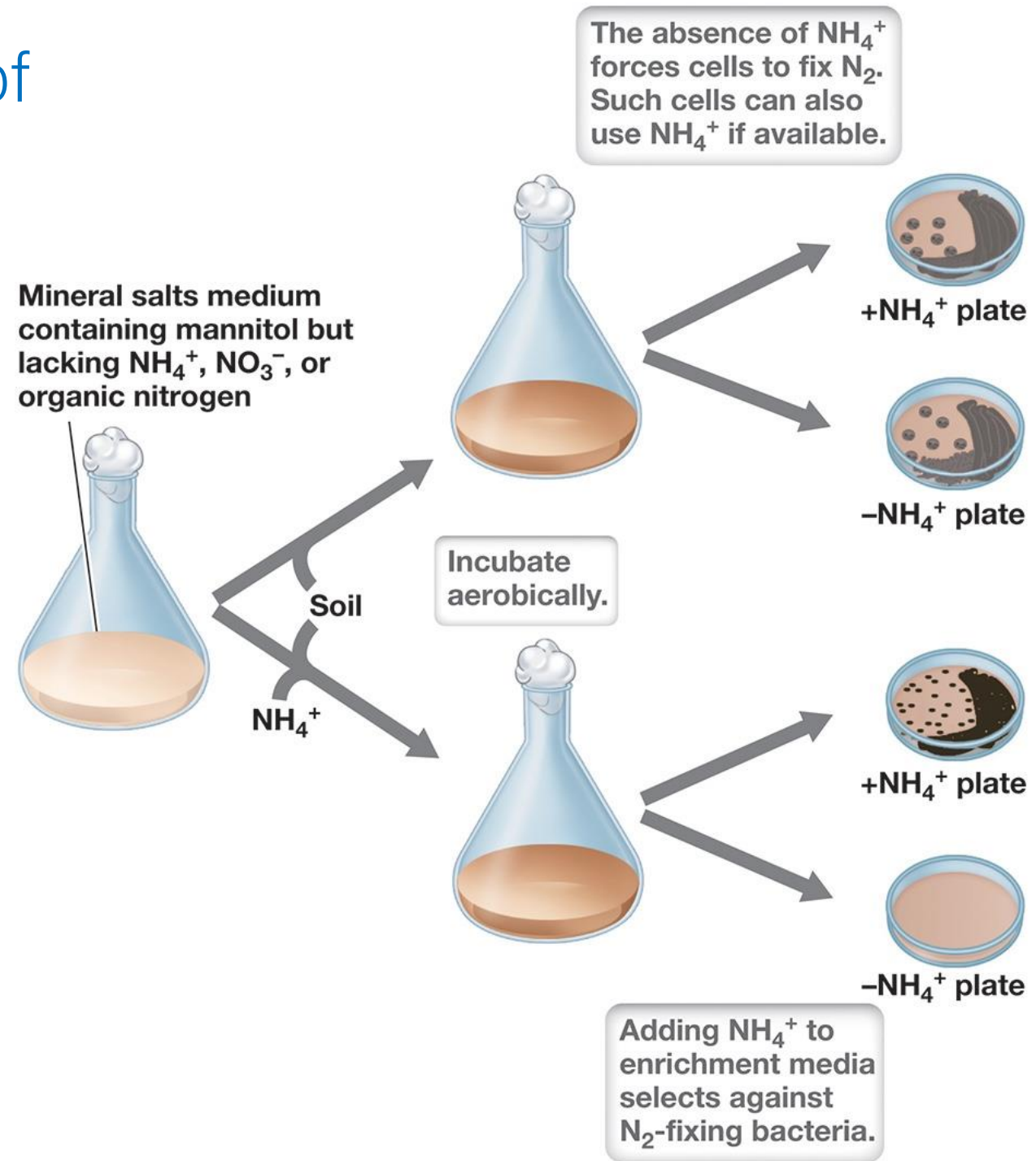
CULTURE DEPENDENT VS. CULTURE INDEPENDENT



ENRICHMENT CULTURE MICROBIOLOGY

- **Inoculum**
 - The sample from which microorganisms is isolated
 - Enrichment cultures are used to isolate bacteria from diverse and densely populated samples, such as feces or soil
- **Isolation**
 - The separation of individual populations from the mixed community
- **Enrichment cultures**
 - Select for desired organisms through manipulation of medium and incubation conditions. This means favoring the growth of target organisms while inhibiting the growth of non-target organisms.

The Isolation of *Azotobacter*



ENRICHMENT CULTURE MICROBIOLOGY

- Enrichment Culture Outcomes
 - Successful enrichment cultures have appropriate resources (nutrients) and conditions (temperature, p H, oxygen, osmotic considerations) that are needed for the target organisms to grow
 - Enrichment cultures can demonstrate the presence of an organism in a habitat
 - They cannot prove that an organism does not inhabit an environment
- Note: The ability to isolate an organism from an environment says nothing about its ecological importance or relative abundance in nature

Some Enrichment Culture Methods for Phototrophic Bacteria (Main C Source - CO₂)

Incubation in air

Incubation condition	Organisms enriched	Inoculum
N ₂ as nitrogen source	Cyanobacteria	Pond or lake water; sulfide-rich muds; stagnant water; raw sewage; moist, decomposing leaf litter; moist soil exposed to light
NO ₃ ⁻ as nitrogen source, 55°C	Thermophilic cyanobacteria	Hot spring microbial mat

Anoxic incubation

Incubation condition	Organisms enriched	Inoculum
H ₂ or organic acids; N ₂ as sole nitrogen source	Purple nonsulfur bacteria, heliobacteria	Same as above plus hypolimnetic lake water; pasteurized soil (heliobacteria); microbial mats for thermophilic species
H ₂ S as electron donor	Purple and green sulfur bacteria	
Fe ²⁺ , NO ₂ ⁻ as electron donor	Purple bacteria	

ENRICHMENT CULTURE MICROBIOLOGY

- Enrichment bias
- Microorganisms cultured in the lab are frequently only minor components of the microbial ecosystem
 - Reason: quantity of nutrients available in the lab culture are typically much higher than in nature
 - Dilution of inoculum may be used to eliminate rapidly growing, but quantitatively insignificant, “weed” species

CLASSICAL PROCEDURES FOR ISOLATING MICROBES

Pure cultures contain a single kind of microorganism

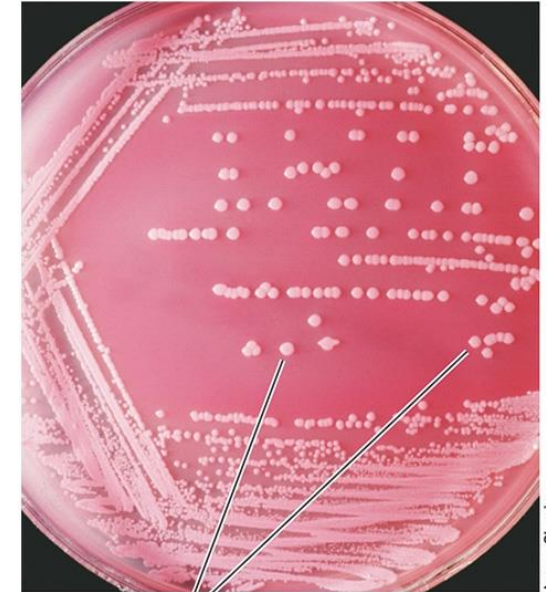
- Can then be used for molecular and physiological experiments

Streak plate

- A well-isolated colony is selected and restreaked several successive times in order to obtain a pure culture

Agar dilution tubes are mixed cultures diluted in molten agar

- Useful for purifying anaerobic organisms



(a) Colonies Paraffin-mineral oil seal



(b)

CULTURE-INDEPENDENT ANALYSES OF MICROBIAL COMMUNITIES

Microscopic Analyses



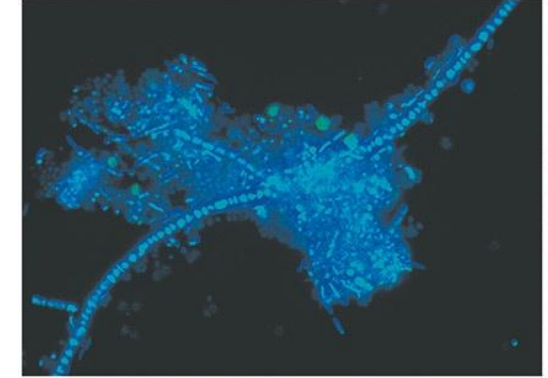
Genetic Analyses



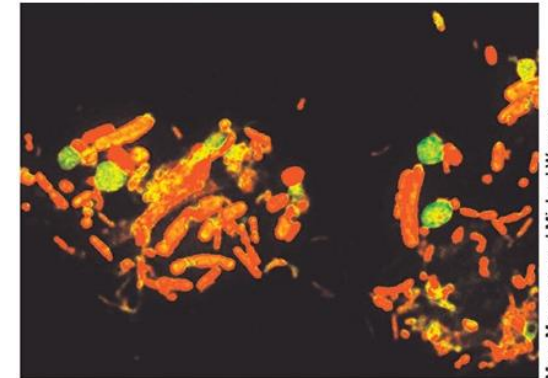
MICROSCOPY ANALYSES OF MICROBIAL COMMUNITIES

Fluorescent staining

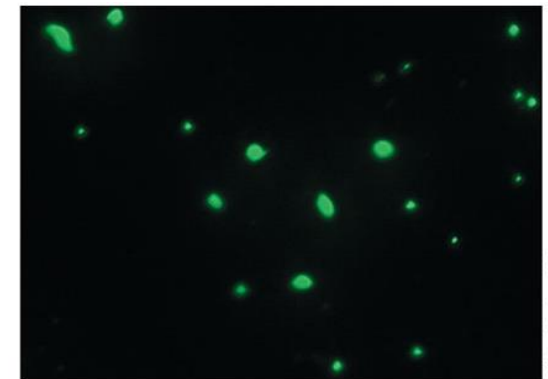
- DAPI-stained cells fluoresce bright blue (DNA)
 - AO-stained cells fluoresce orange (RNA) or green (DNA)
 - SYBR-stained cells fluoresce green (DNA)
-
- Fluoresce under UV light
 - Enumeration of microorganisms in samples
 - Nonspecific and stain nucleic acids, such as DNA
 - Stain nonspecifically
 - cannot differentiate between live and dead cells



Marc Mussman and Michael Wagner



Marc Mussman and Michael Wagner



Willm Martins-Haberna

VIABILITY STAINS DIFFERENTIATE BETWEEN LIVE AND DEAD CELLS

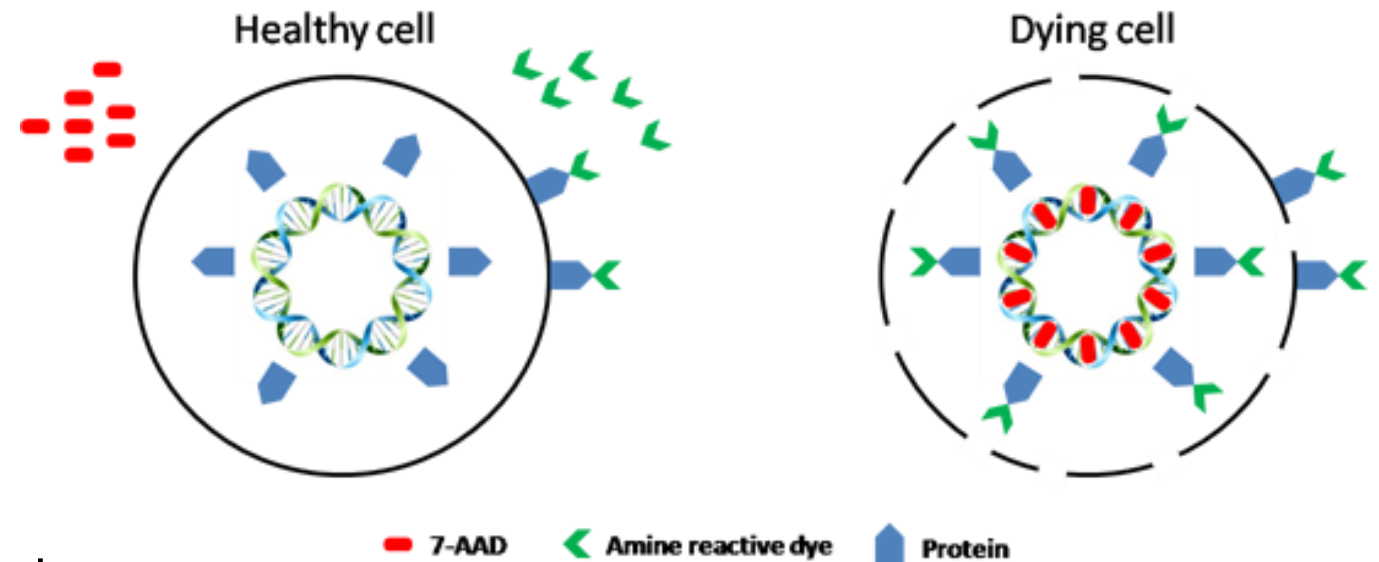
Two dyes are used

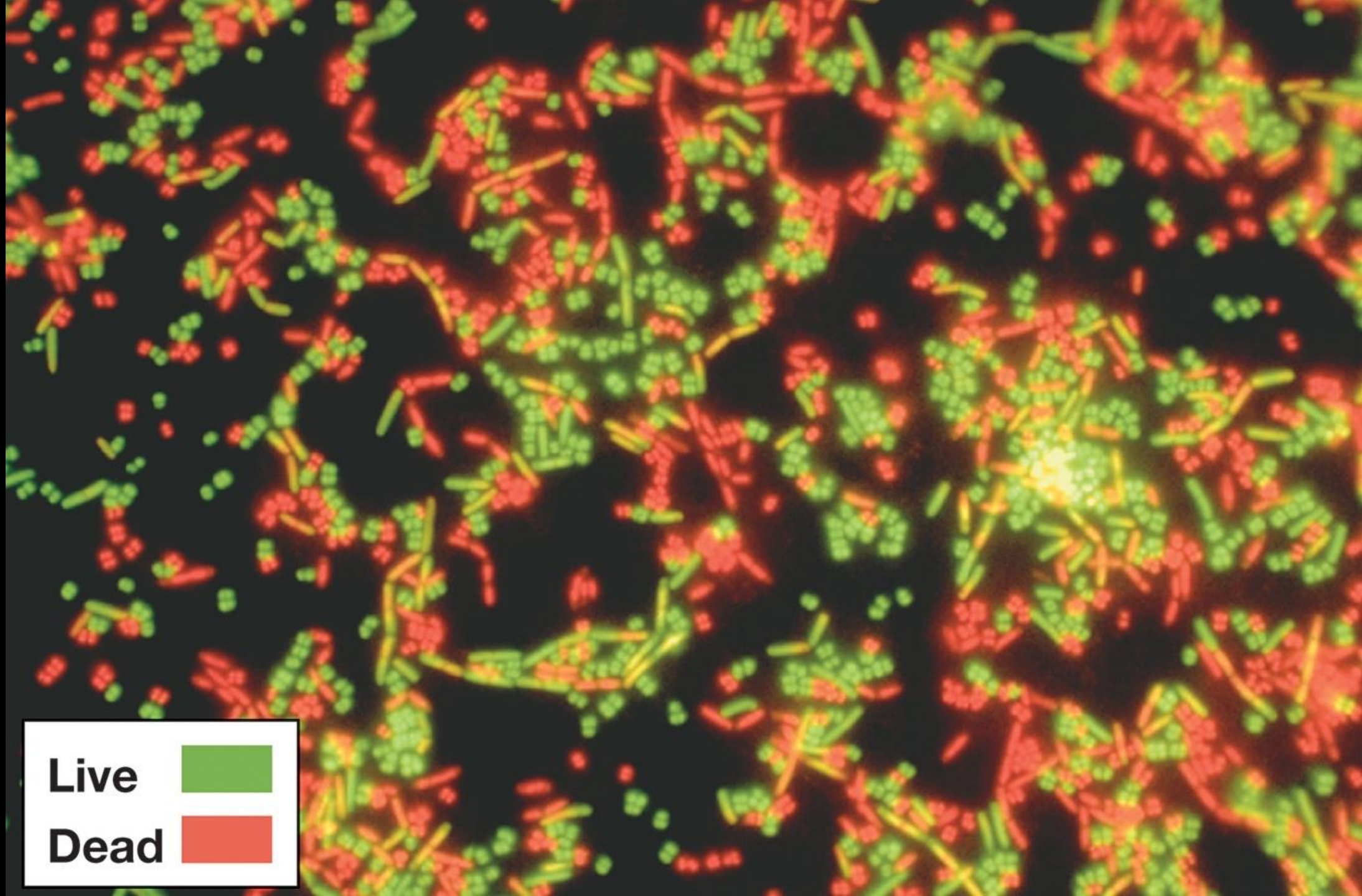
one that stains live cells and the other that stains dead cells, based on integrity of cell membrane

Green cells are live

Red cells are dead

This method may stain background nonspecifically with environmental samples





Live



Dead



CULTURE-INDEPENDENT ANALYSES OF MICROBIAL COMMUNITIES

Microscopic Analyses



Genetic Analyses



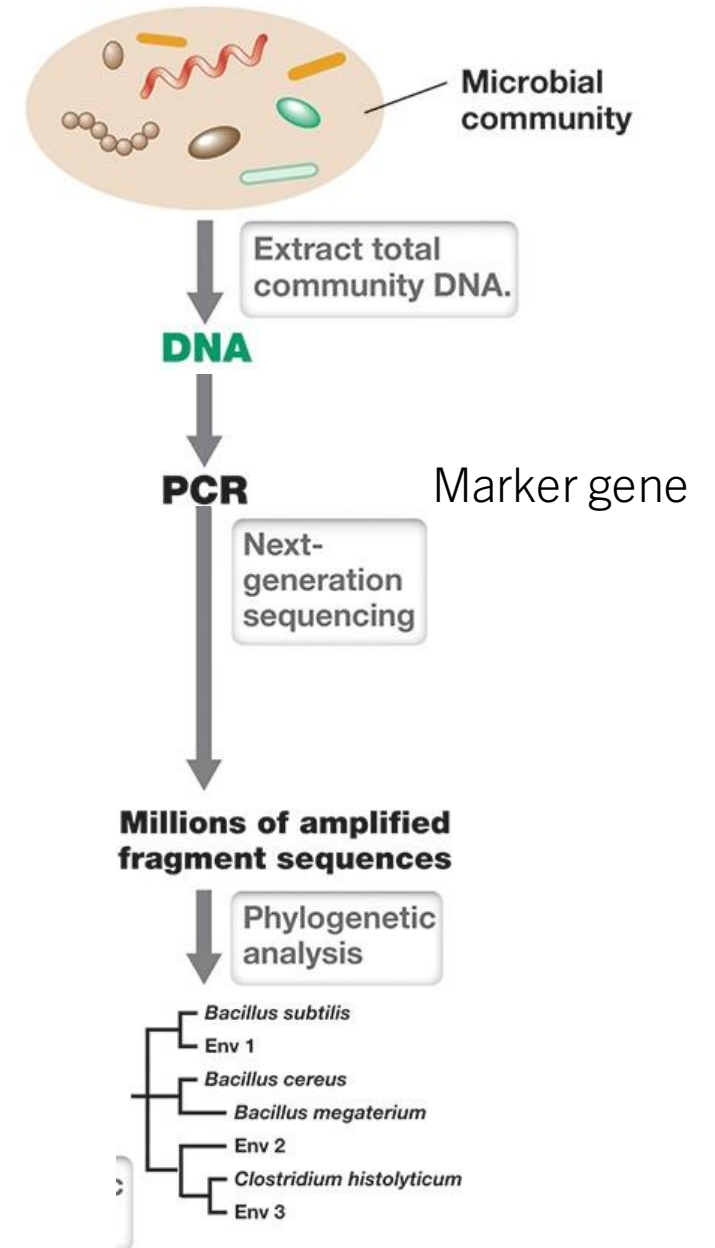
Culture-Independent Genetic Analyses of Microbial Communities

- PCR Methods of Microbial Community Analysis
- Environmental Multi-omics

PCR Methods of Microbial Community Analysis

Specific genes can be used as a measure of diversity: **marker genes**

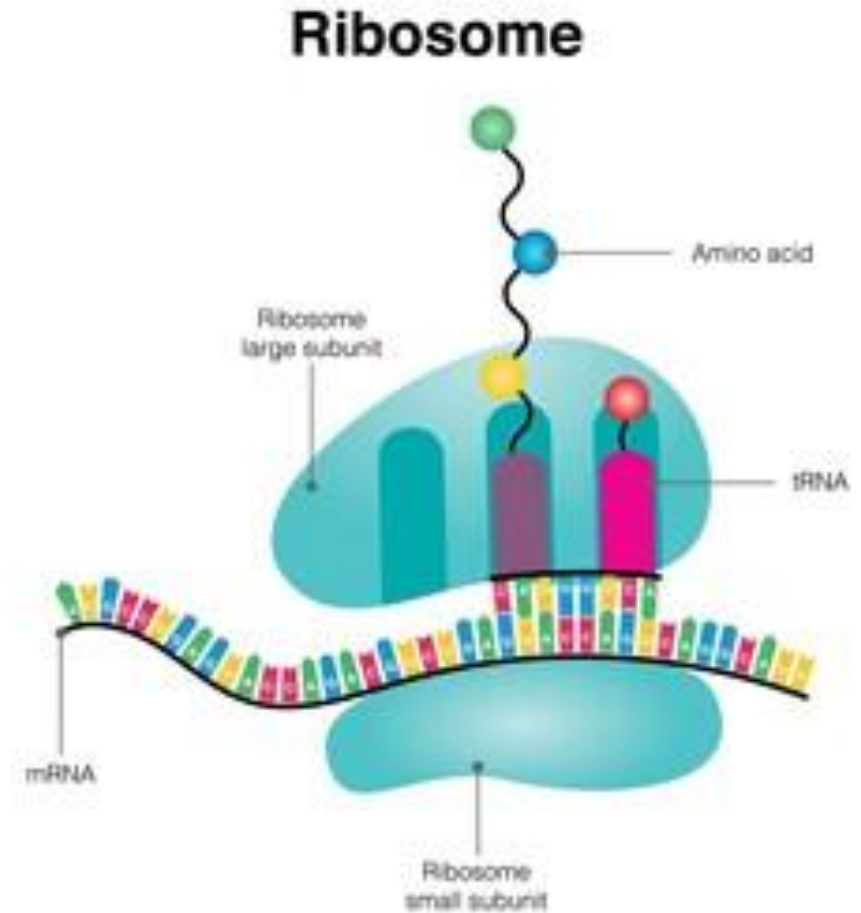
- 16S – bacteria and archaea
- ITS – fungi
- Functional genes – antibiotic production, biogeochemical cycles, etc.



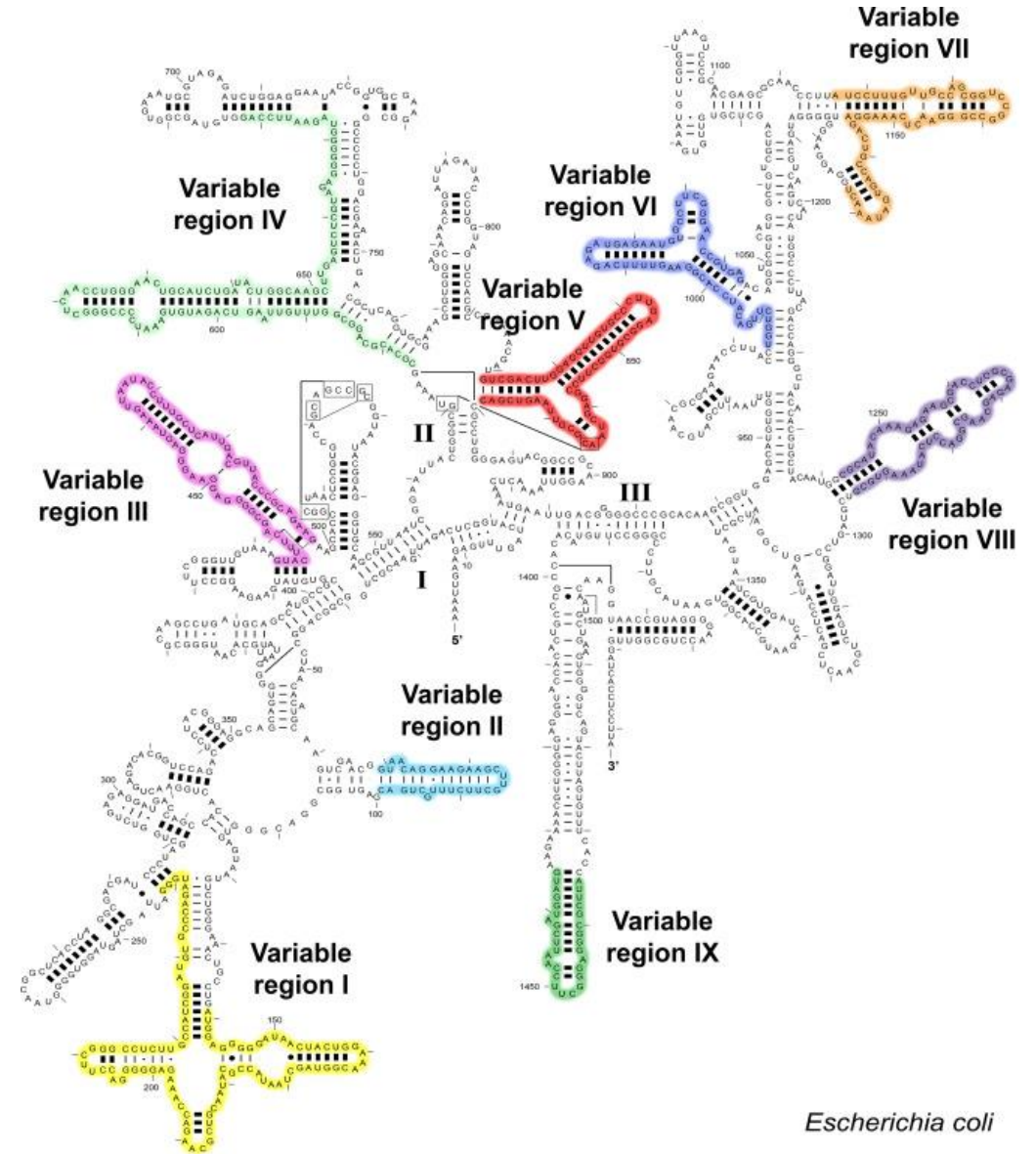
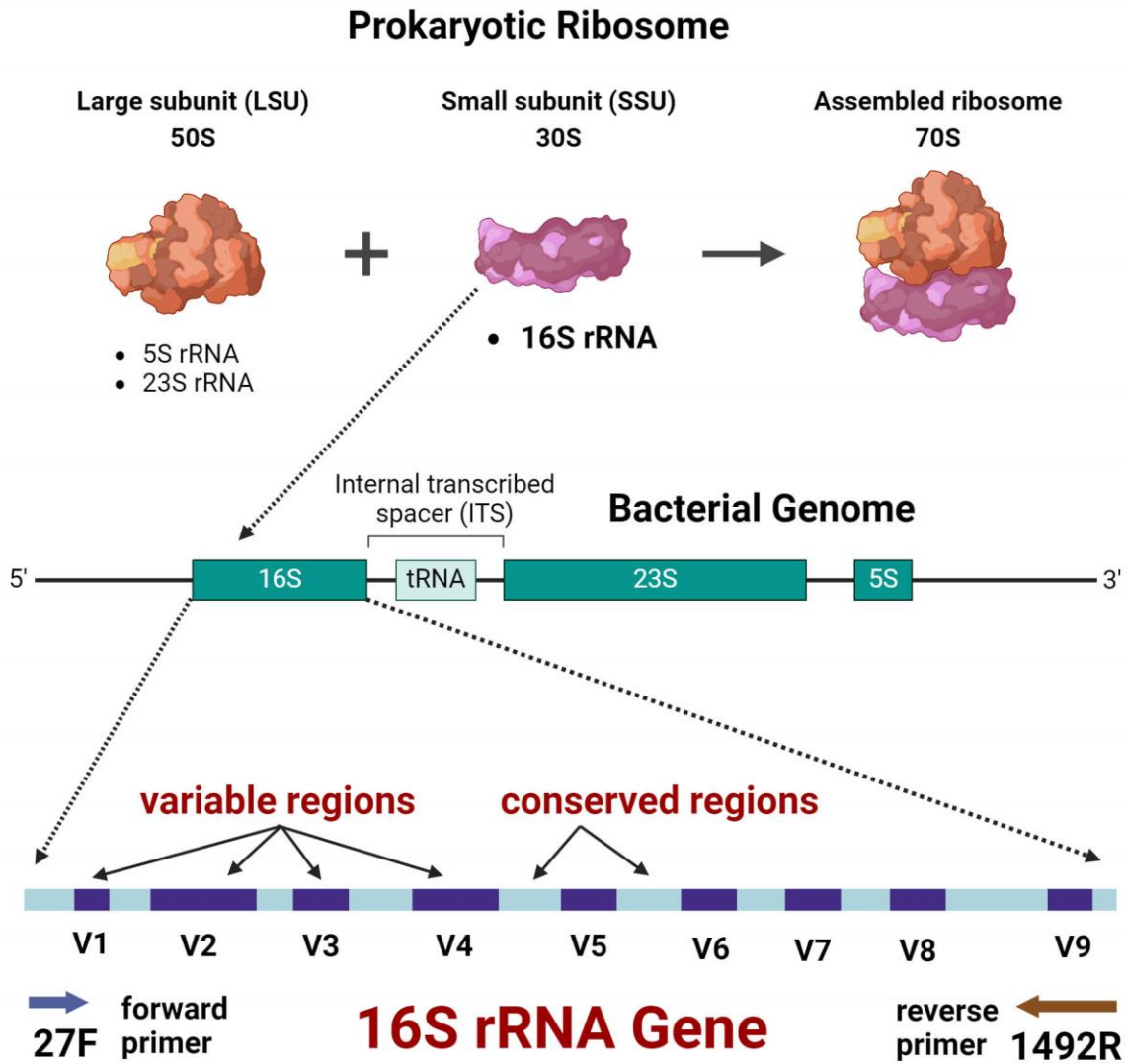
Ribosomal RNA

Ribosomal RNA (rRNA) – RNA molecules that associate with ribosomal proteins to form the ribosomes

Ribosomes are responsible for protein synthesis in the cell



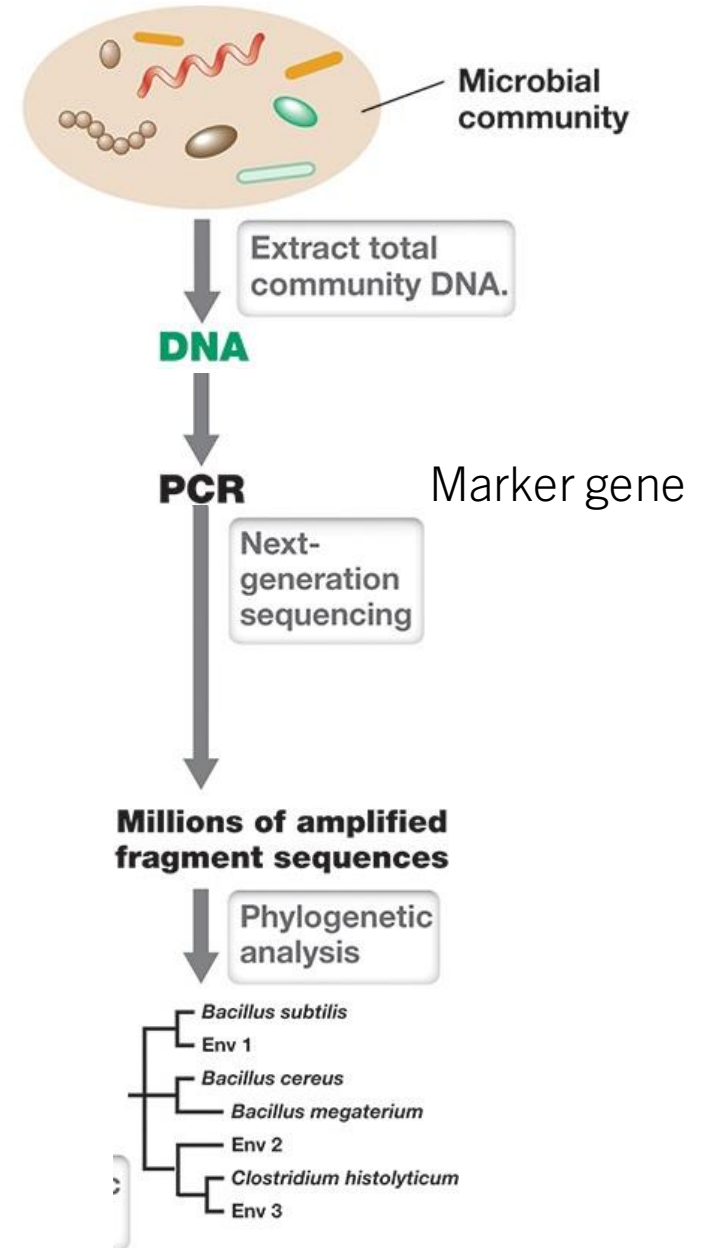
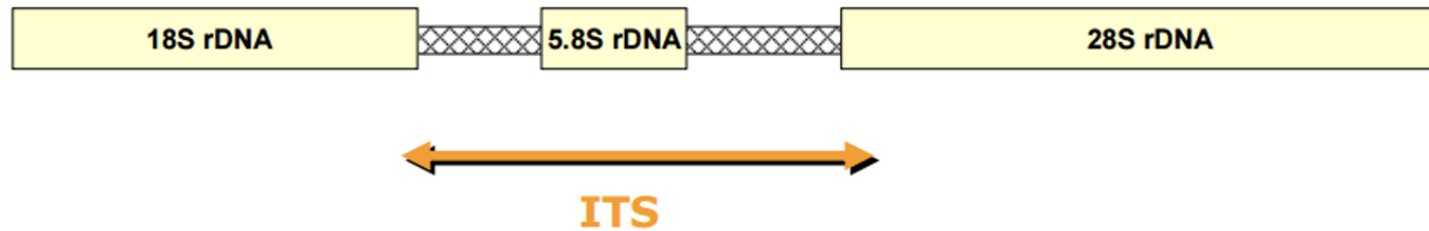
What is 16S rRNA gene?



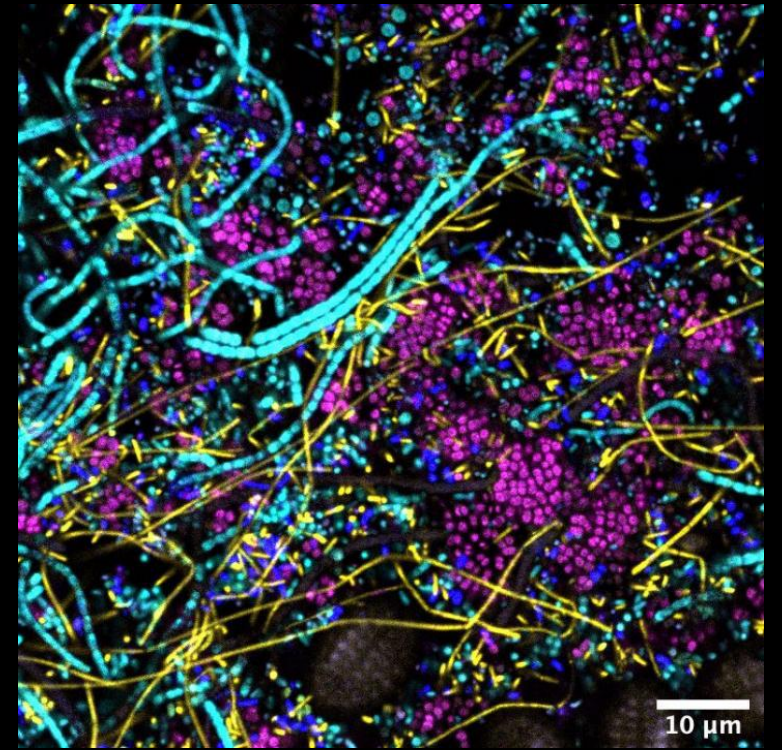
Marker gene for bacteria and archaea

PCR Methods of Microbial Community Analysis

- ITS – fungi



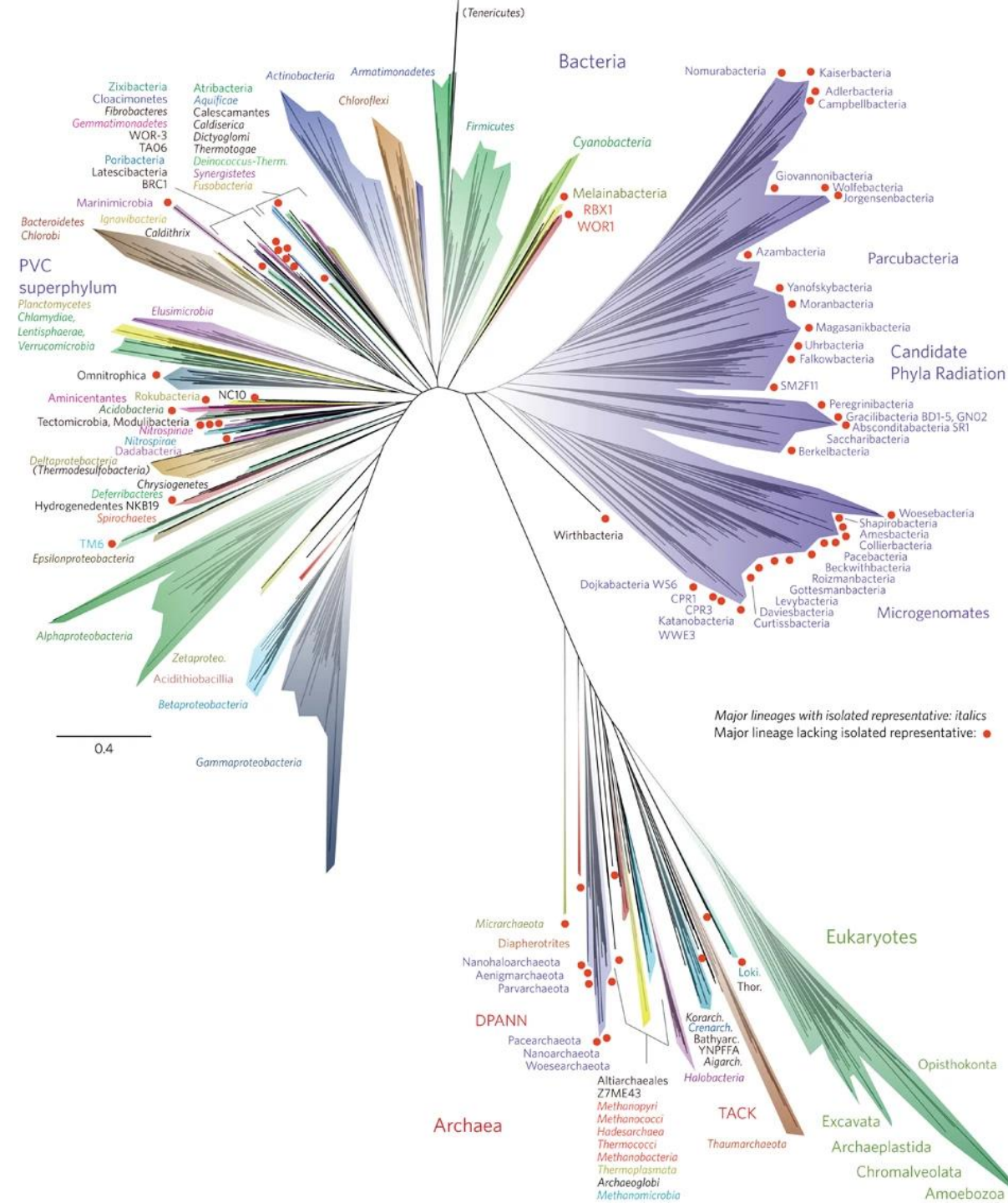
rRNA sequences in environmental samples differ from those of all known laboratory cultures



- fewer than 0.1% of bacteria have been cultured
- enrichment bias is a real problem to culture-based methods

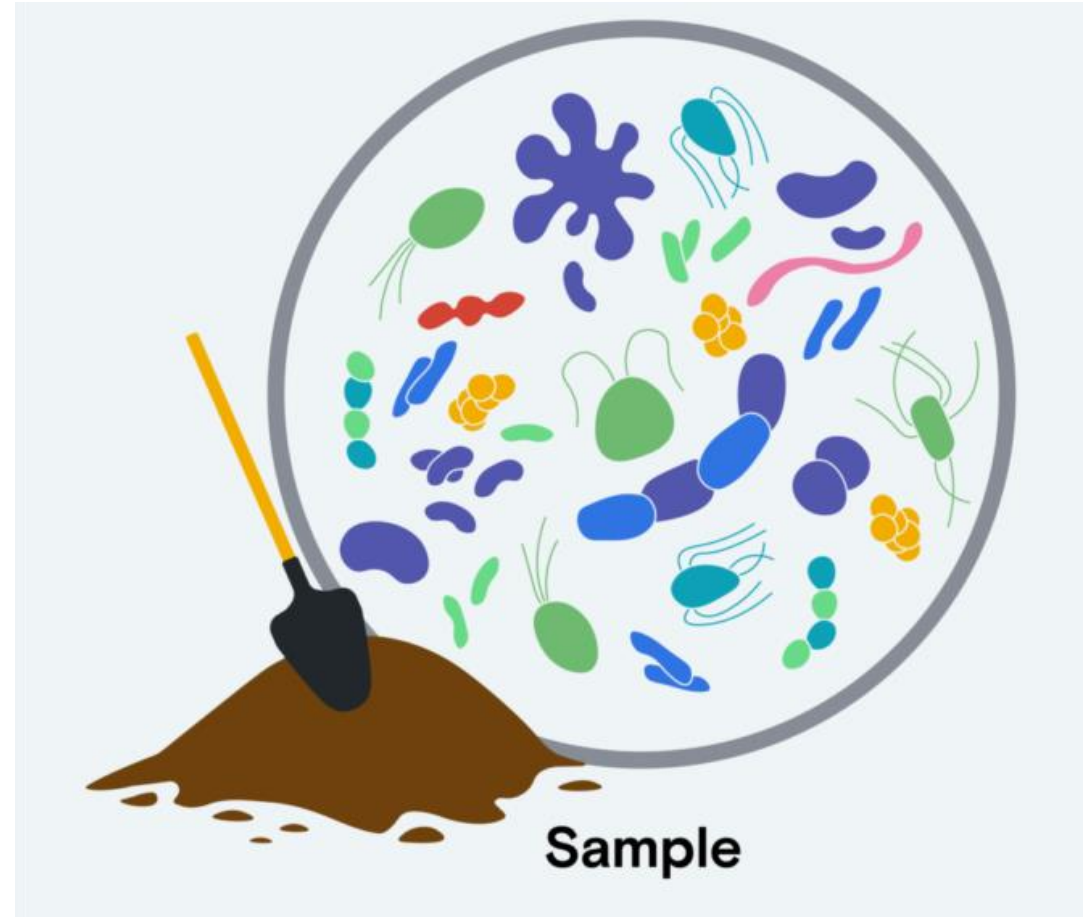


A new view of the tree of life



Metagenomics

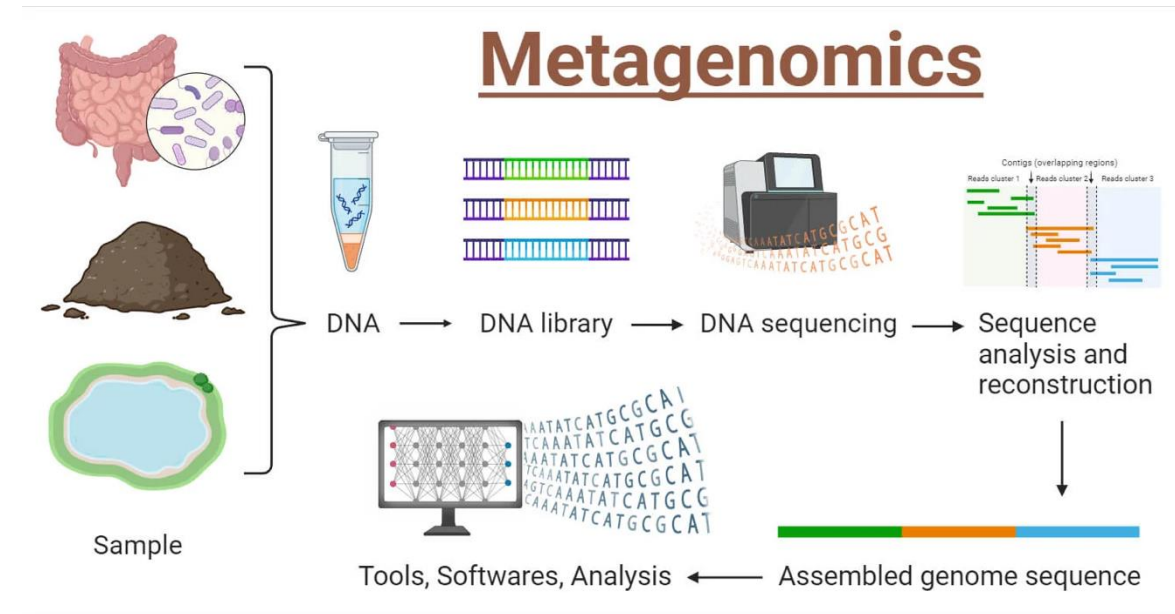
- Metagenomics – Environmental genomics
- The study of the structure and function of **all nucleotide sequences** from all the organisms in a bulk sample.
 - structure of the microbial community (who is there)
 - function of the microbial community (what can they do)



Metagenome: total gene content of microbial community

Metagenomics

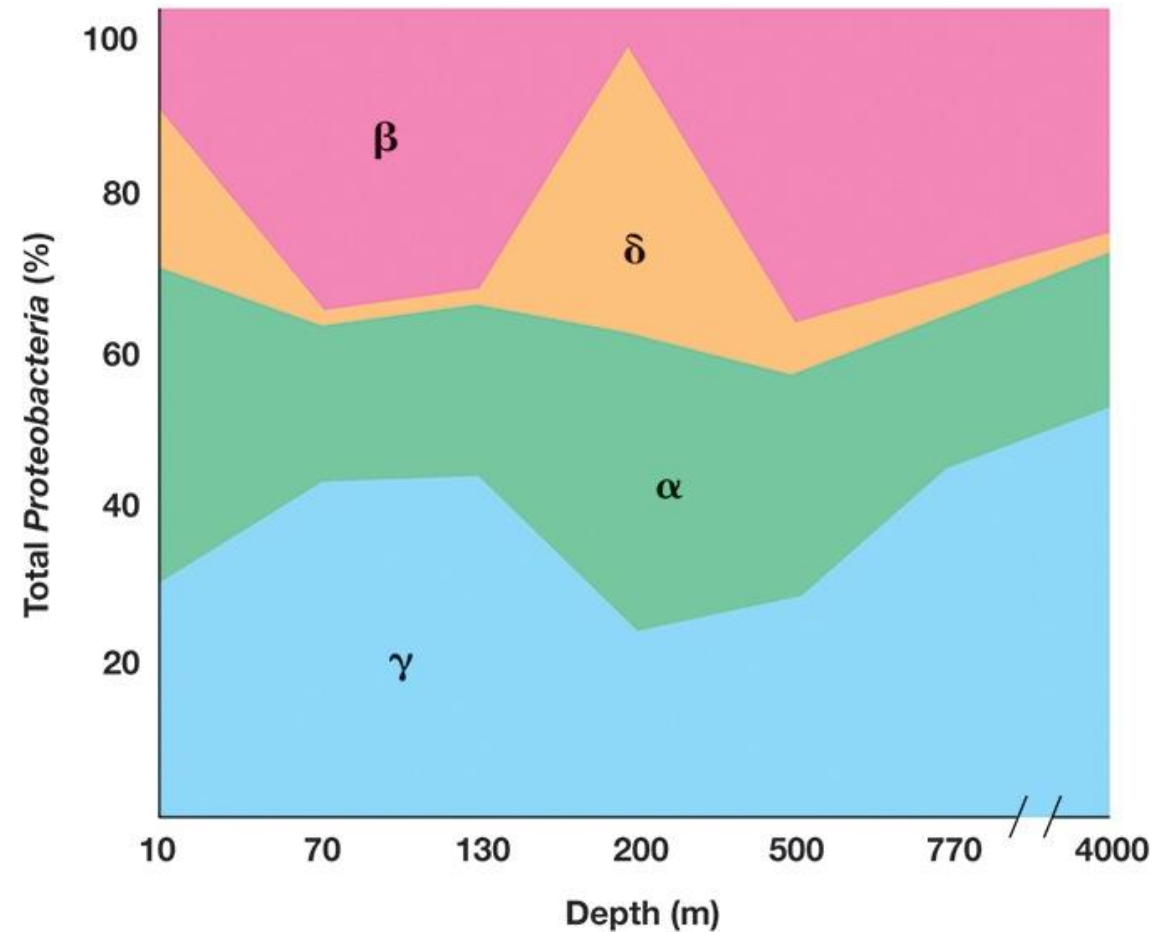
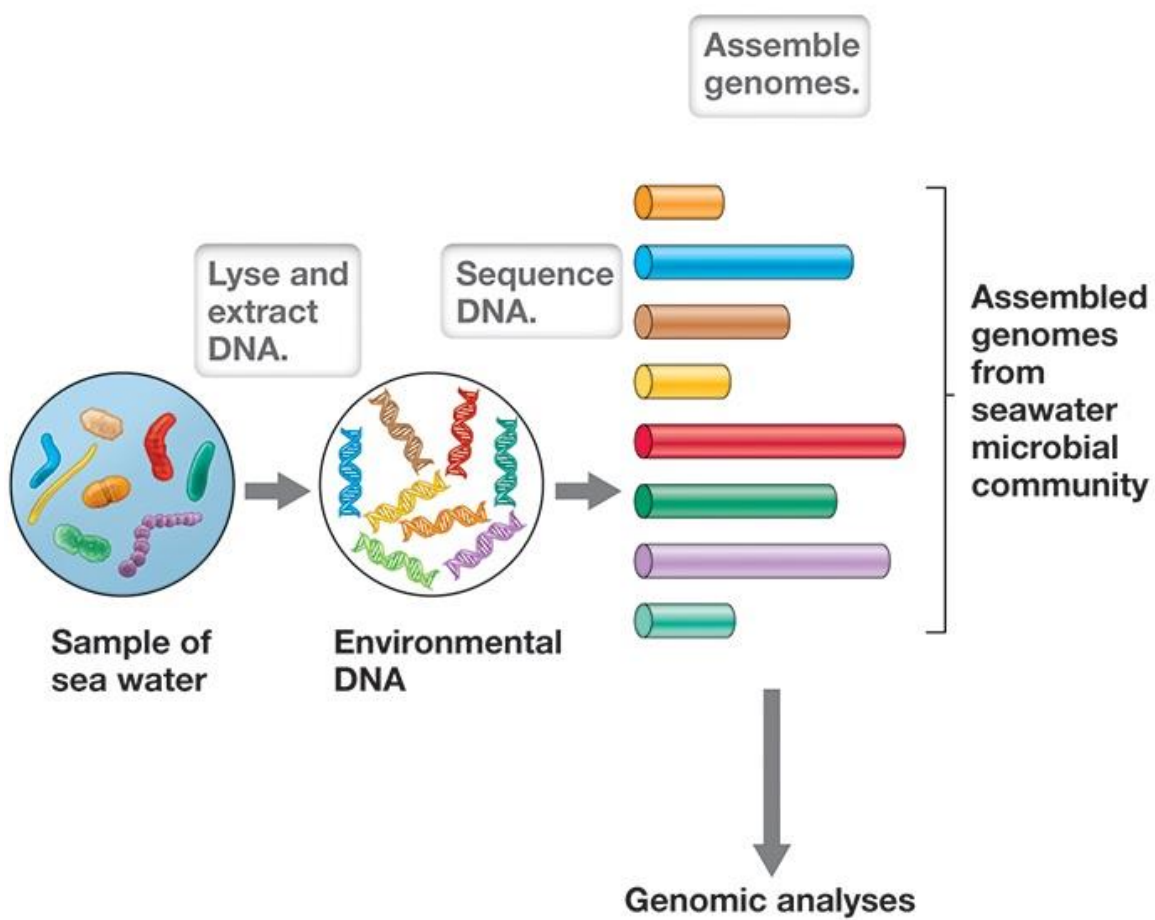
- DNA is isolated from the entire microbial community and sequenced
- Detects as many genes as possible
- Yields picture of gene pool in environment
- Can detect genes that are not amplified by current PCR primers
- Powerful tool for assessing the phylogenetic and metabolic diversity of an environment



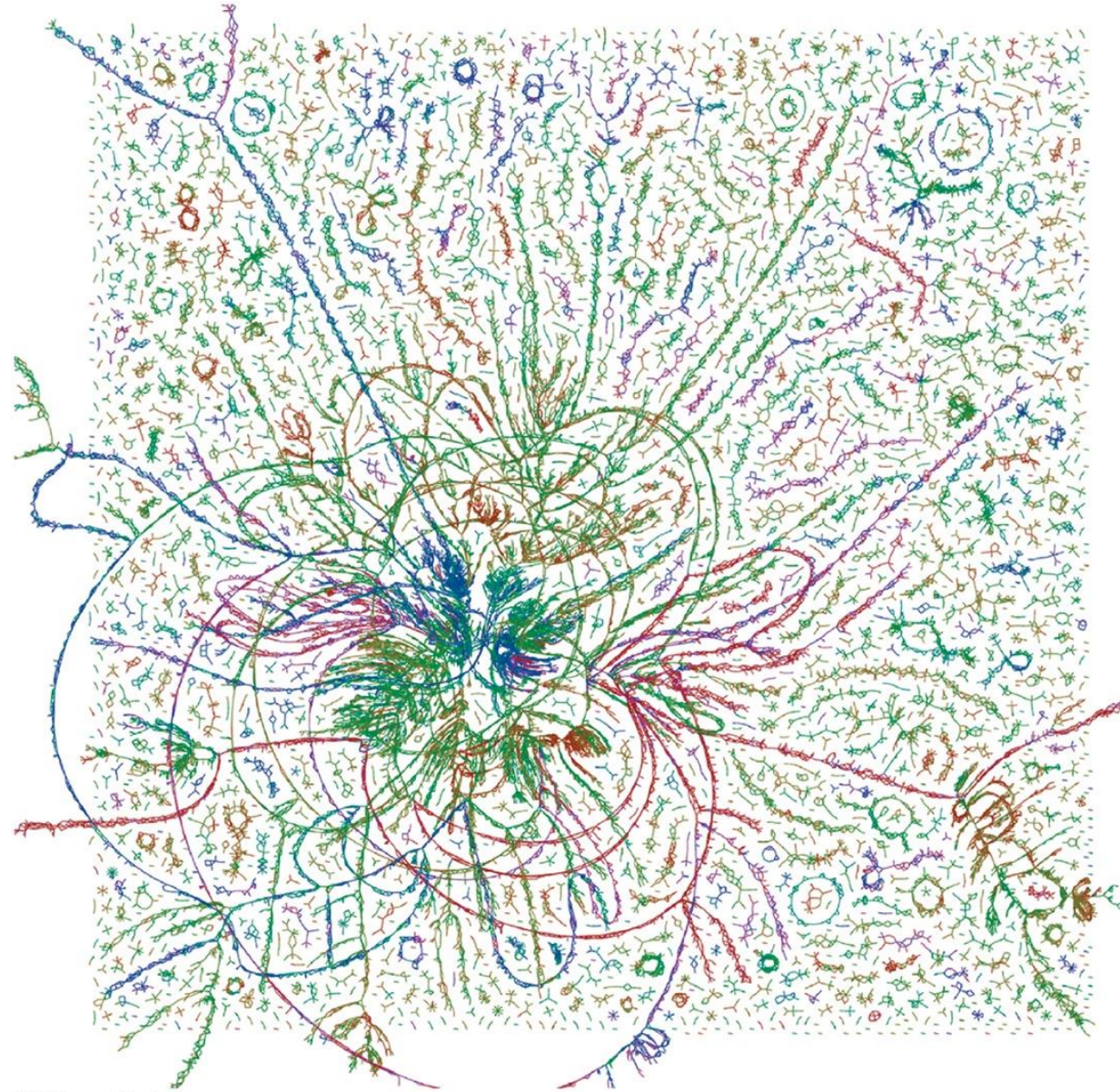
Metagenomics and the Microbiome



Metagenomics and the Microbiome

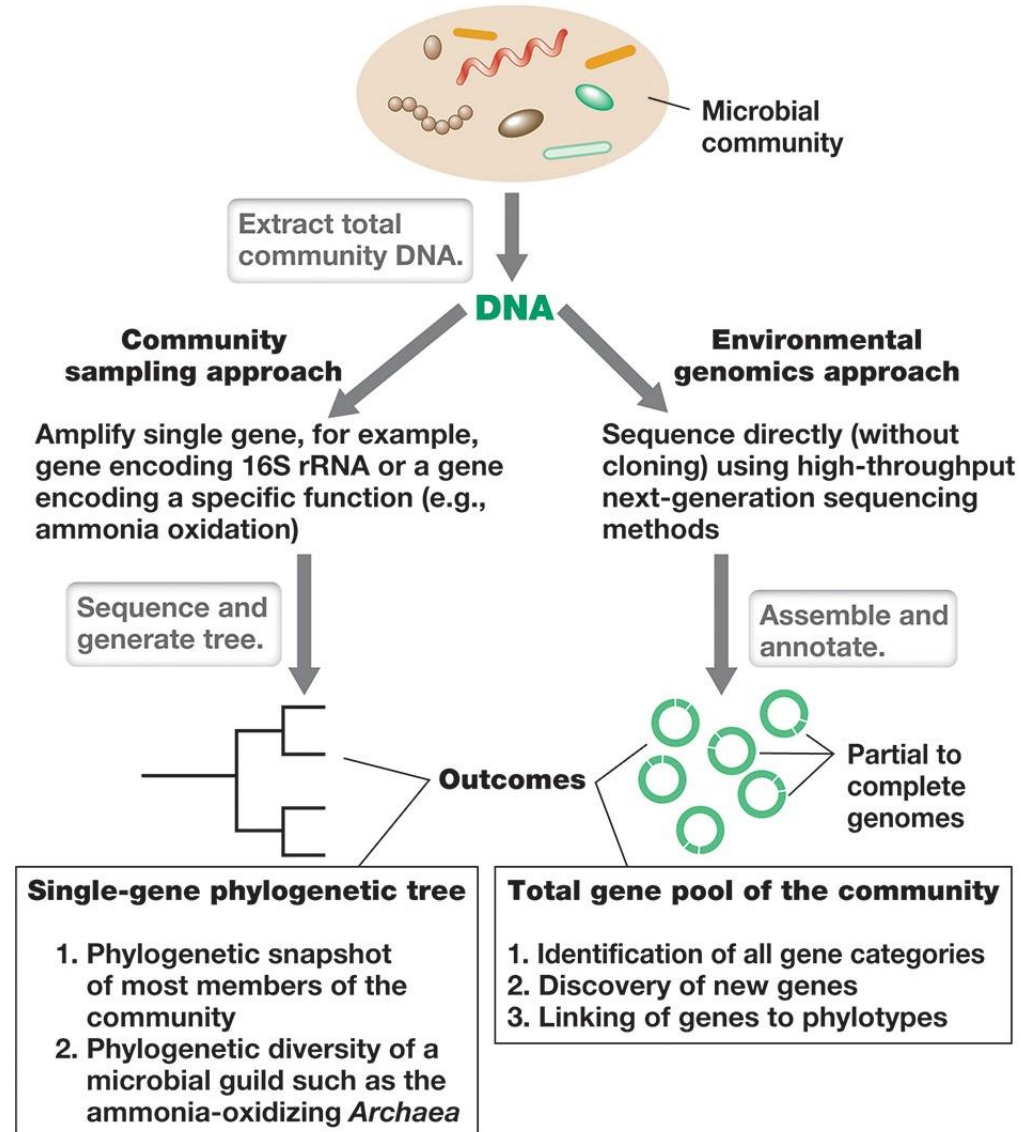


Genome Assembly From a Coastal Marine Metagenome Consisting of 58.5 Billion Nucleotides of Sequence



Vaughn Iverson and Ginger Armbrust

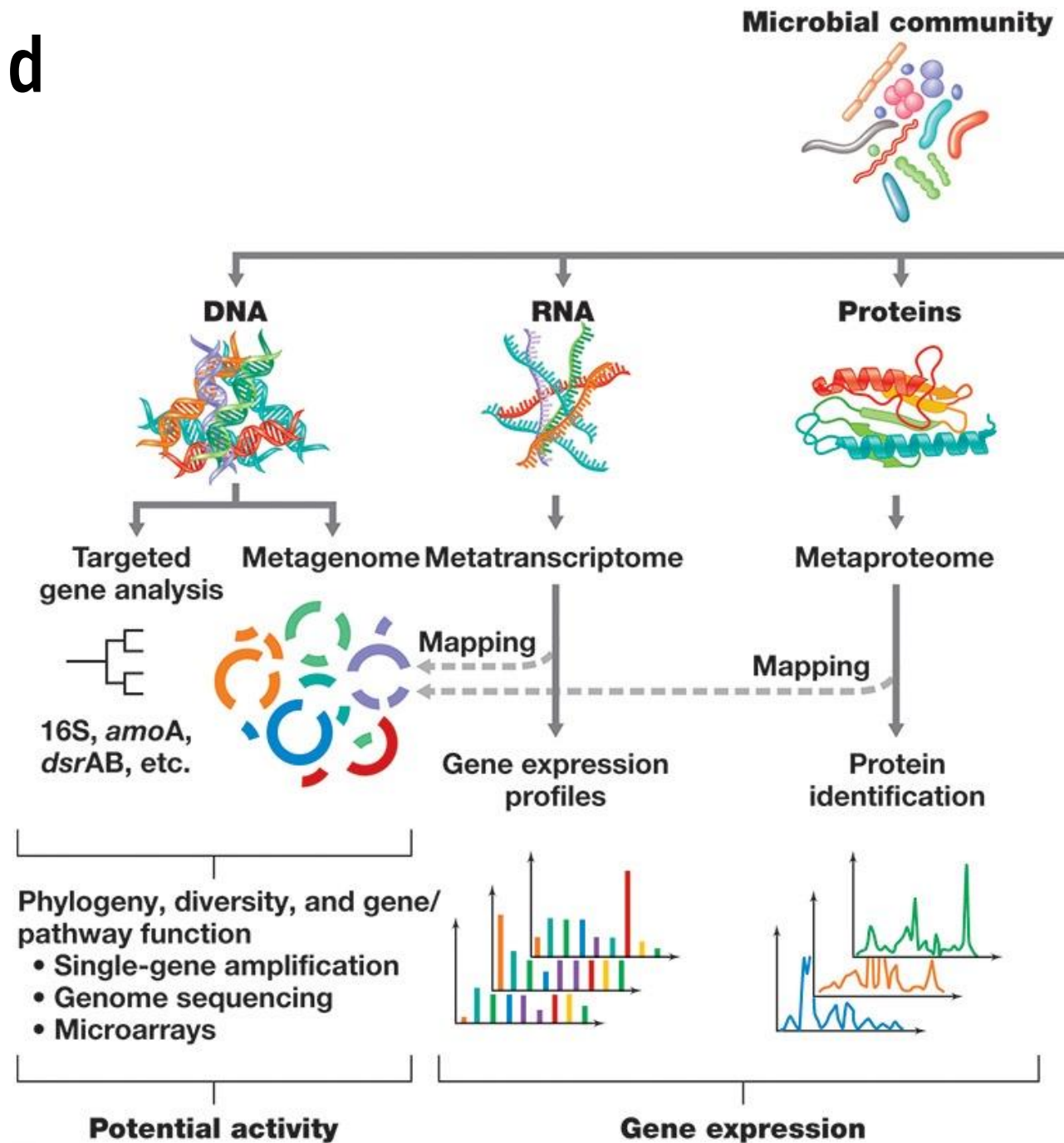
Single-Gene Versus Environmental Genomic Approaches to Microbial Community Analysis



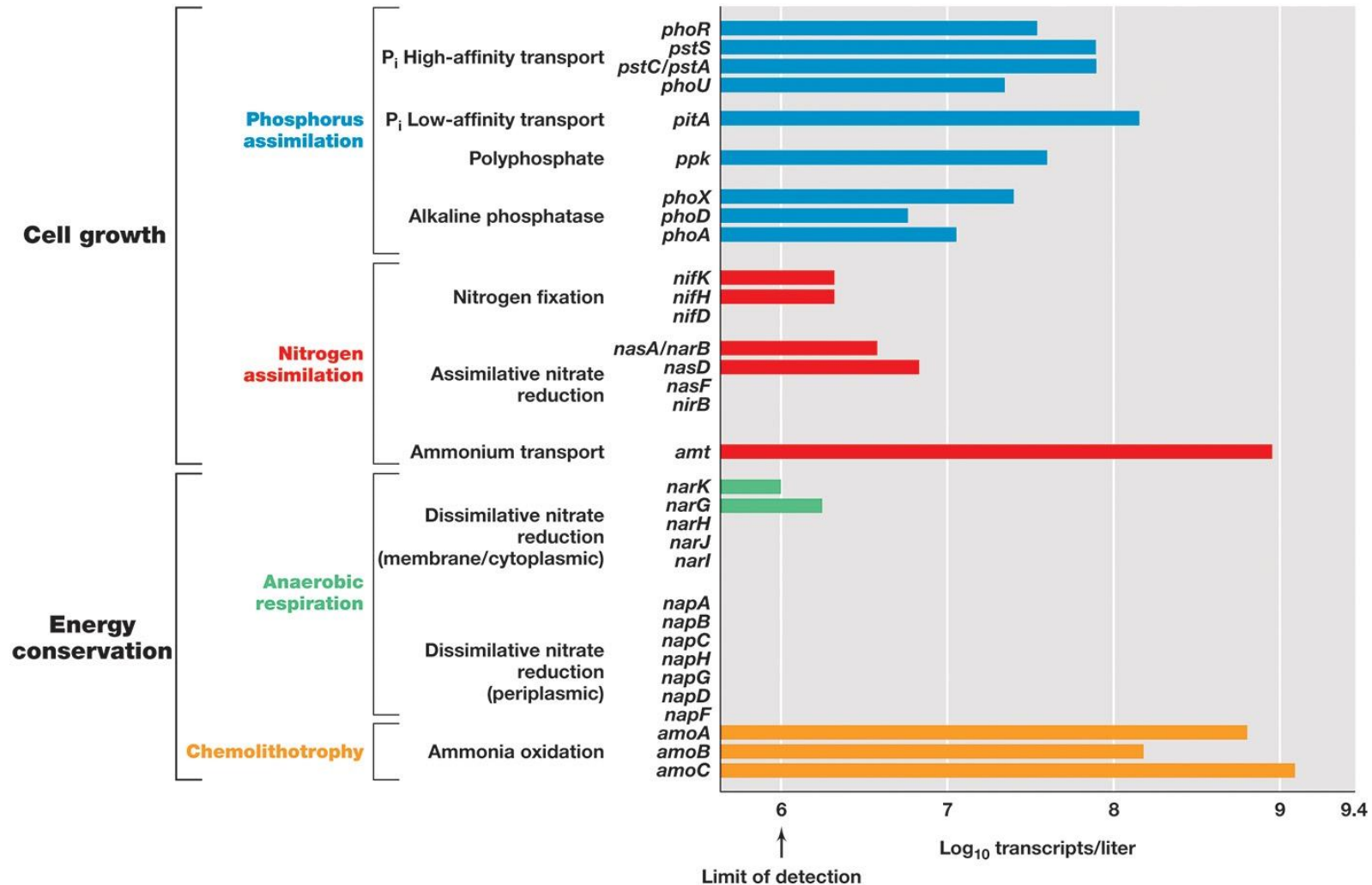
Metatranscriptomics and Metaproteomics

Functionally active portion of the community

- Analyzes community RNA and proteins
 - (who are the active members)
- Reveals genes/proteins in a community that are actually expressed
 - (what they are actually doing)

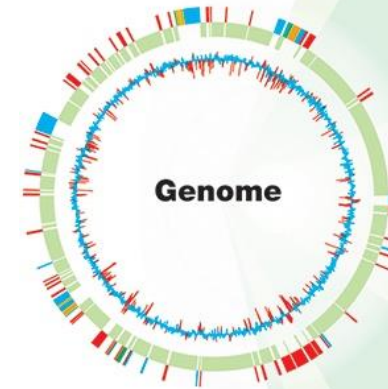


Metatranscriptomic Analysis of Coastal Marine Surface Waters



INTRODUCTION TO GENOMICS

- “Omic”: Broad discipline integrating different methodologies to characterize and quantify large pools of biomolecules
- **Genome:** entire complement of genetic information includes genes encoding proteins, RNA, and regulatory sequences, and noncoding DNA
- Major omic themes: genomics, transcriptomics, proteomics, metabolomics, metagenomics
- **Genomics:** discipline of mapping, sequencing, analyzing, and comparing genomes



Metagenomics

Assessing the entire gene content of a microbial community



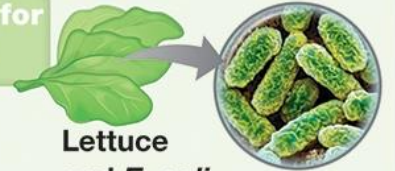
Transcriptomics Proteomics Metabolomics

Interactive mapping and predictive modeling



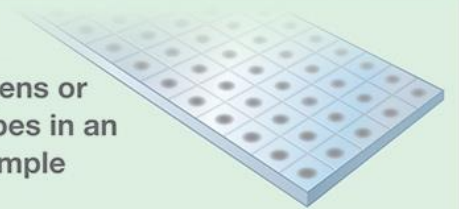
Specific sequences for strain identification

Monitoring disease outbreaks



Diagnostics

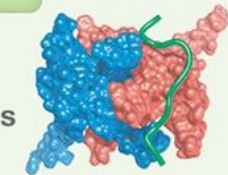
Detecting pathogens or identifying microbes in an environmental sample



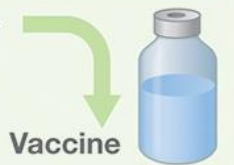
Proteomics

Identifying potential drug targets

Immune response



3D structure



Vaccine development

Drug design

Insights into metabolism or virulence

Revealing nutrients required for culturing and clues regarding pathogenicity



INTRODUCTION TO GENOMICS

- Advances rely on improvements in molecular technologies and computing power
- Number of sequenced genomes has grown rapidly
- Bottleneck is vast amounts of nucleic acid sequence data
- Genomics: Then and Now
 - First genomes sequenced were small viruses
 - First bacterial genome published in 1995
 - Today, DNA sequences from 125,000+ *Bacteria*, *Archaea*, and viruses are available publicly
 - Many eukaryotic genomes sequenced, including human genome

Whole-Genome Random Sequencing and Assembly of *Haemophilus influenzae* Rd

Robert D. Fleischmann, Mark D. Adams, Owen White, Rebecca A. Clayton, Ewen F. Kirkness, Anthony R. Kertavage, Carol J. Bult, Jean-Francois Tomb, Brian A. Dougherty, Joseph M. Merrick, Keith McKenney, Granger Sutton, Will FitzHugh, Chris Fields,* Jeannine D. Gocayne, John Scott, Robert Shirley, Li-Ing Liu, Anna Glodek, Jenny M. Kelley, Janice F. Weidman, Cheryl A. Phillips, Tracy Spriggs, Eva Hedblom, Matthew D. Cotton, Teresa R. Utterback, Michael C. Hanna, David T. Nguyen, Deborah M. Saudek, Rhonda C. Brandon, Leah D. Fine, Janice L. Fritchman, Joyce L. Fuhrmann, N. S. M. Geoghagen, Cheryl L. Gnehm, Lisa A. McDonald, Keith V. Small, Claire M. Fraser, Hamilton O. Smith, J. Craig Ventert†

An approach for genome analysis based on sequencing and assembly of unselected pieces of DNA from the whole chromosome has been applied to obtain the complete nucleotide sequence (1,830,137 base pairs) of the genome from the bacterium *Haemophilus influenzae* Rd. This approach eliminates the need for initial mapping efforts and is therefore applicable to the vast array of microbial species for which genome maps are unavailable. The *H. influenzae* Rd genome sequence (Genome Sequence DataBase accession number L42023) represents the only complete genome sequence from a free-living organism.

A prerequisite to understanding the complete biology of an organism is the determination of its entire genome sequence. Several viral and organellar genomes have been completely sequenced. Bacteriophage ϕ X174 [5386 base pairs (bp)] was the first to be sequenced, by Fred Sanger and colleagues in 1977 (1). Sanger *et al.* were also the first to use strategy based on random (unselected) pieces of DNA, completing the genome sequence of bacteriophage λ (48,502 bp) with cloned restriction enzyme fragments (1). Subsequently, the 229-kb genome of cytomegalovirus (CMV) (2), the 192-kb genome of vaccinia (3), and the 187-kb mitochondrial and

Homo sapiens (11). These projects, as well as viral genome sequencing, have been based primarily on the sequencing of clones usually derived from extensively mapped restriction fragments, or λ or cosmid clones. Despite advances in DNA sequencing technology (12) the sequencing of genomes has not progressed beyond clones on the order of the size of λ (~40 kb). This has been primarily because of the lack of sufficient computational approaches that would enable the efficient assembly of a large number (tens of thousands) of independent, random sequences into a single assembly.

The computational methods developed to create assemblies from hundreds of thou-

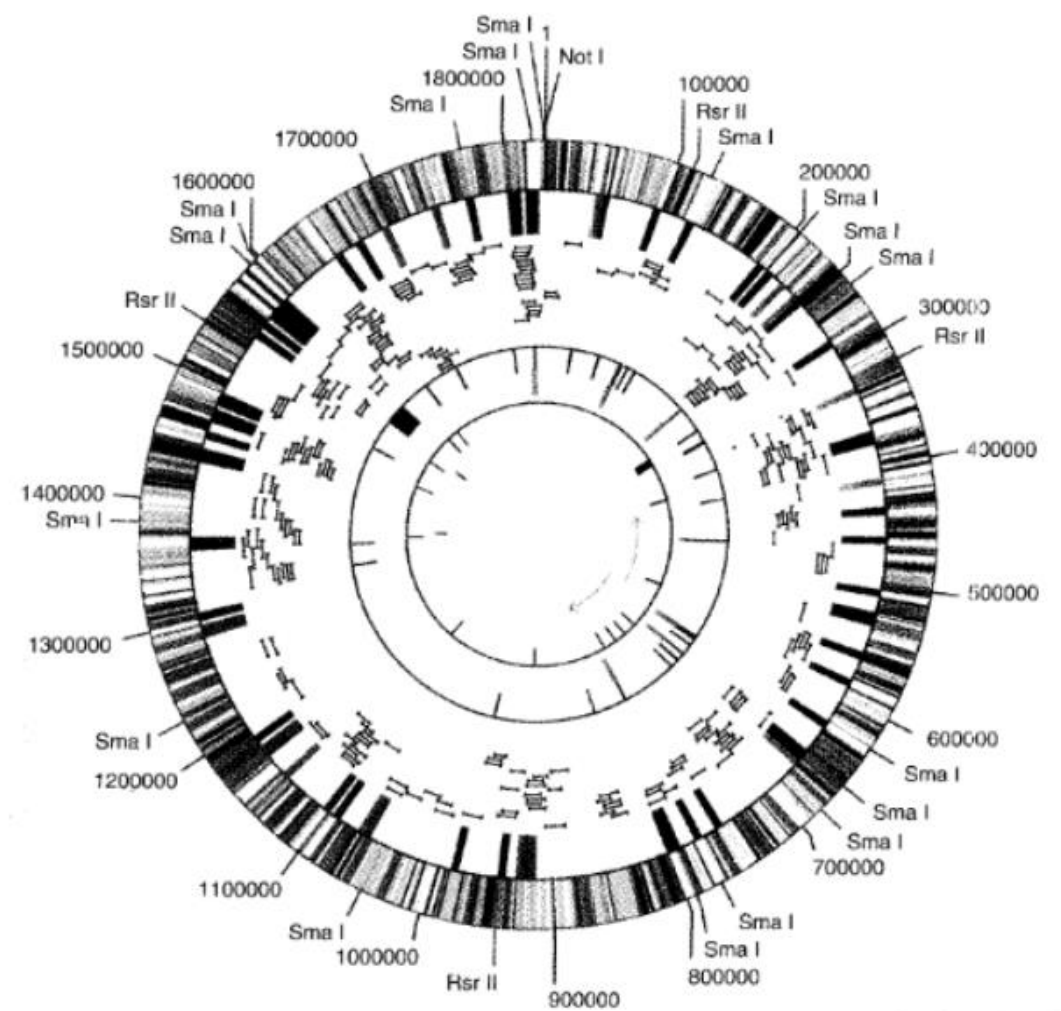
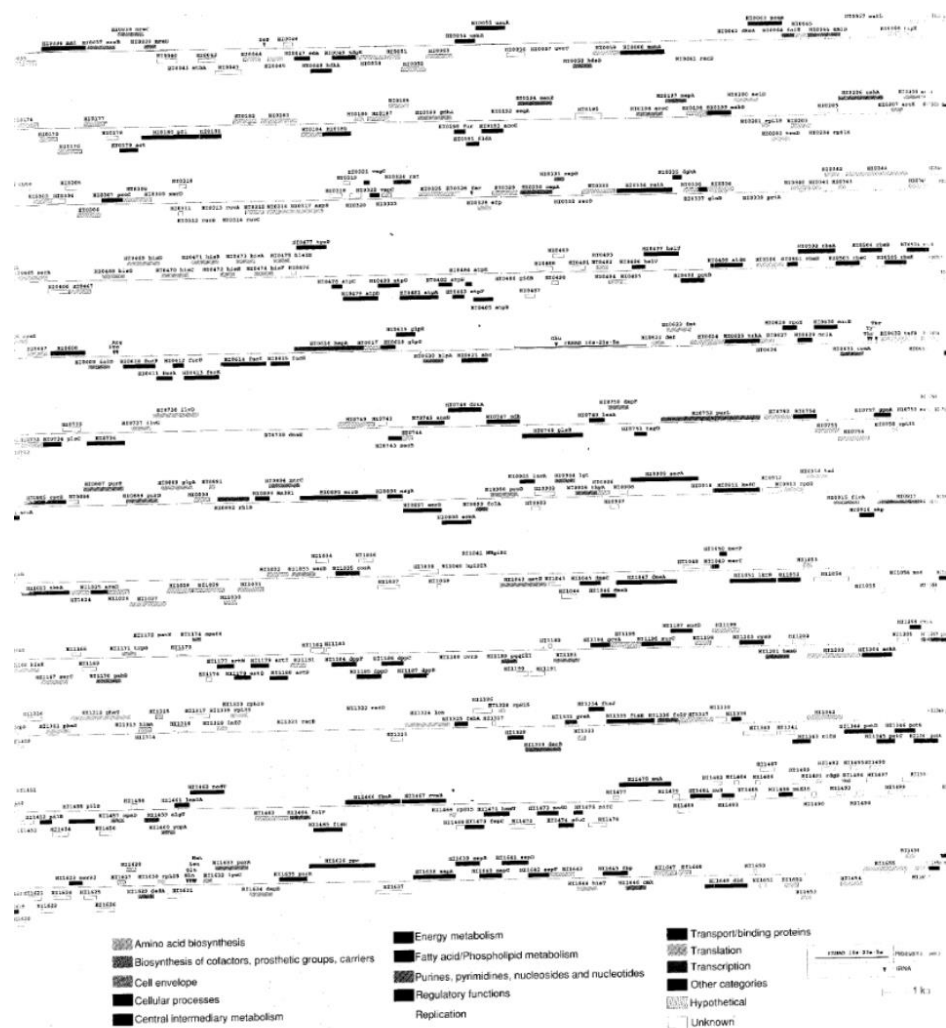


Fig. 1. A circular representation of the *H. influenzae* Rd chromosome illustrating the location of each predicted coding region containing a database match as well as selected global features of the genome. Outer perimeter: The location of the unique Not I restriction site (designated as nucleotide 1), the Rsr II sites, and the Sma I sites. Outer concentric circle: Coding regions for which a gene identification was made. Each coding region location is classified as to role according to the color code in Fig. 2. Second concentric circle: Regions of high G+C content (>42 percent, red; >40 percent, blue) and high A+T content (>66 percent, black; >64 percent, green). Third concentric circle: Coverage by λ clones (blue). More than 300 λ clones were sequenced from each end to confirm the overall structure of the genome and identify the six ribosomal operons. Fourth concentric circle: The locations of the six ribosomal operons (green), the tRNAs (black) and the cryptic μ -like prophage (blue). Fifth concentric circle: Simple tandem repeats. The locations of the following repeats are shown: CTGGCT, GTCT, ATT, AATGGC, TTGA, TTGG, TTTA, TTATC, TGAC, TCGTC, AACC, TTGC, CAAT, CCAA. The putative origin of replication is illustrated by the outward pointing arrows (green) originating near base 603,000. Two potential termination sequences are shown near the opposite midpoint of the circle (red).



Fig. 3. A comparison of the region of the *H. influenzae* chromosome containing the eight genes of the fimbrial gene cluster present in *H. influenzae* type b and the same region in *H. influenzae* Rd. The region is flanked by *pepN* and *purE* in both organisms. However, in the noninfectious Rd strain the eight genes of the fimbrial gene cluster have been excised. A 172-bp spacer region is located in this region in the Rd strain and continues to be flanked by the *pepN* and *purE* genes.

Genomes of Select Species of *Bacteria* and *Archaea*

Organism	Lifestyle ^b	Size (bp)	ORFs ^c	Features
<i>Nasuia deltocephalinicola</i>	E	112,091	137	Degenerate sap-feeding insect endosymbiont
<i>Tremblaya princeps</i>	E	138,931	121	Degenerate mealybug endosymbiont
<i>Hodgkinia cicadicola</i>	E	143,795	169	Degenerate cicada endosymbiont
<i>Mycoplasma genitalium</i>	P	580,070	525	Smallest nonsymbiotic bacterial genome
<i>Rickettsia prowazekii</i>	P	1,111,523	834	Obligate intracellular parasite, causes epidemic typhus
<i>Treponema pallidum</i>	P	1,138,006	1,041	Spirochete, causes syphilis
<i>Methylophilaceae</i> family, strain HTCC2181	FL	1,304,428	1,354	Marine methylotroph, smallest free-living genome
<i>Thermotoga maritima</i>	FL	1,860,725	1,877	Hyperthermophile

Genomes of Select Species of *Bacteria* and *Archaea*

Organism	Lifestyle ^b	Size (bp)	ORFs ^c	Features
<i>Deinococcus radiodurans</i>	FL	3,284,156	2,185	Radiation resistant, multiple chromosomes
<i>Bacillus subtilis</i>	FL	4,214,810	4,100	Gram-positive genetic model
<i>Mycobacterium tuberculosis</i>	P	4,411,529	3,924	Causes tuberculosis
<i>Escherichia coli</i> K-12	FL	4,639,221	4,288	Gram-negative genetic model
<i>Escherichia coli</i> O157:H7	FL	5,594,477	5,361	Enteropathogenic strain of <i>E. coli</i>
<i>Bradyrhizobium japonicum</i>	FL	9,105,828	8,317	Nitrogen fixation, nodulates soybeans
<i>Sorangium cellulosum</i>	FL	14,782,125	11,559	Forms multicellular fruiting bodies
<i>Minicystis rosea</i>	FL	16,040,666	14,018	Forms multicellular fruiting bodies

Genomes of Select Species of *Bacteria* and *Archaea*

Archaea

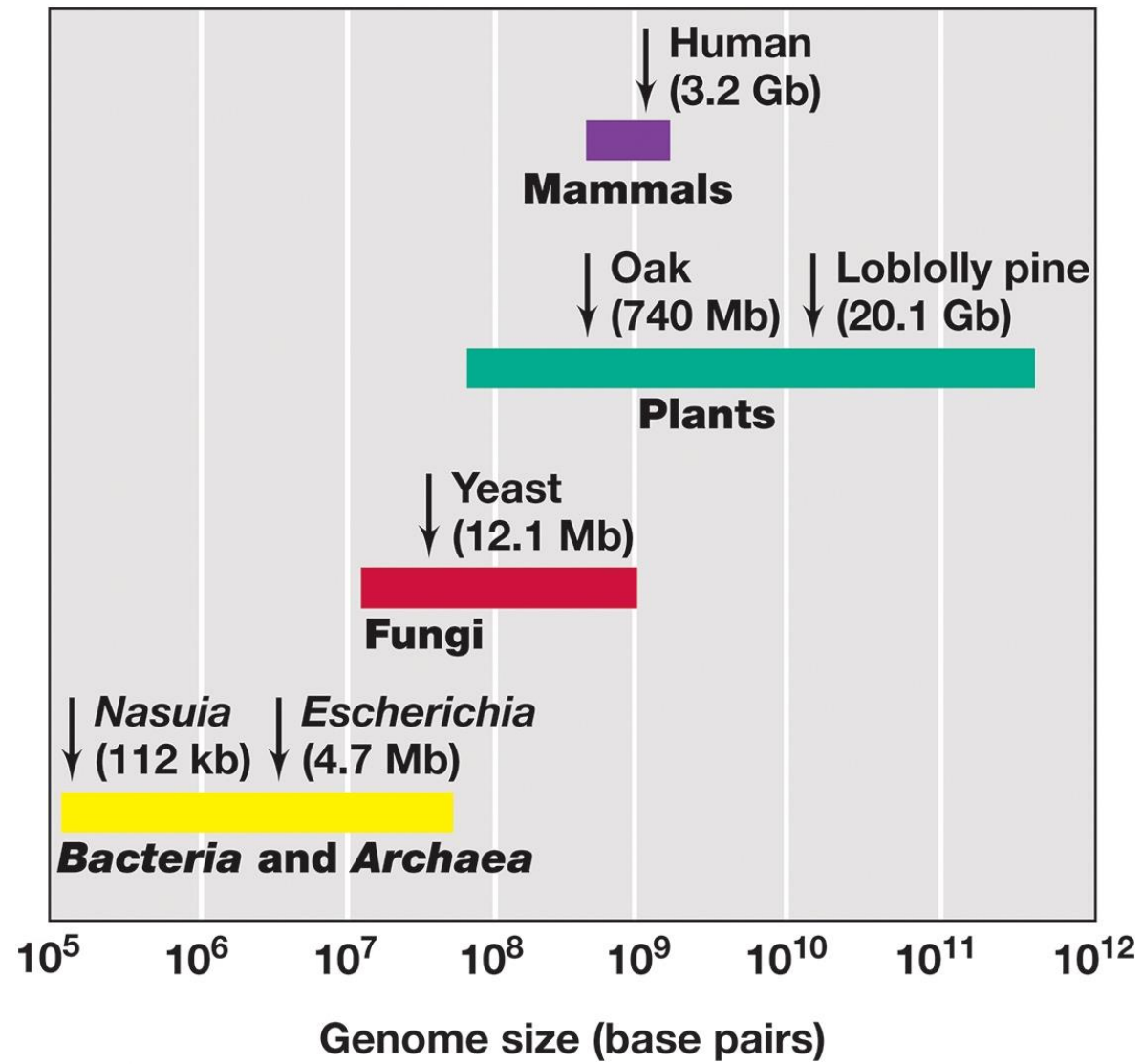
Organism	Lifestyle ^b	Size (bp)	ORFs ^c	Features
<i>Nanoarchaeum equitans</i>	P	490,885	552	Smallest nonsymbiotic cellular genome
<i>Methanocaldococcus jannaschii</i>	FL	1,664,976	1,738	Methanogen, hyperthermophile
<i>Pyrococcus horikoshii</i>	FL	1,738,505	2,061	Hyperthermophile
<i>Sulfolobus solfataricus</i>	FL	2,992,245	2,977	Hyperthermophile, sulphur chemolithotroph
<i>Haloarcula marismortui</i>	FL	4,274,642	4,242	Extreme halophile, bacteriorhodopsin
<i>Methanosarcina acetivorans</i>	FL	5,751,000	4,252	Acetate using methanogen

^a Information on prokaryotic genomes can be found at

^bE, endosymbiont; P, parasite; FL, free-living.

^cOpen reading frames. Genes encoding known proteins are included, as well as ORFs that could encode a protein greater than 100 amino acid residues.

Genome Sizes of Microbial Cells and Higher Organisms



Genomics

What Can Genomes Tell Us?

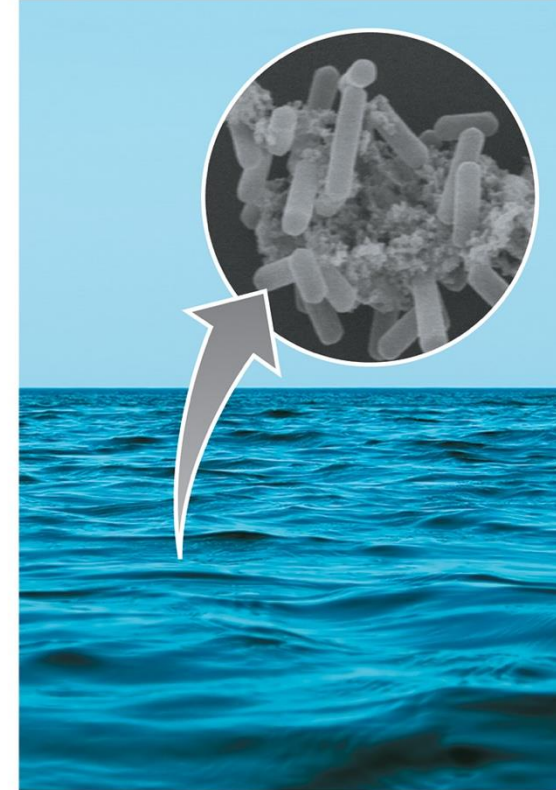
- Modern microbiology thrives on sequences
- Heat-stable enzymes to virulence factors
- Studying gene expression, detecting horizontal transfer, monitoring and diagnosing disease outbreaks, anti-bacteriophage systems, understanding metabolism, and determining growth requirements.....

- Solving medical mysteries (e.g., cause of “Black Death”)

- Identifying new microbial phyla



Yersinia pestis

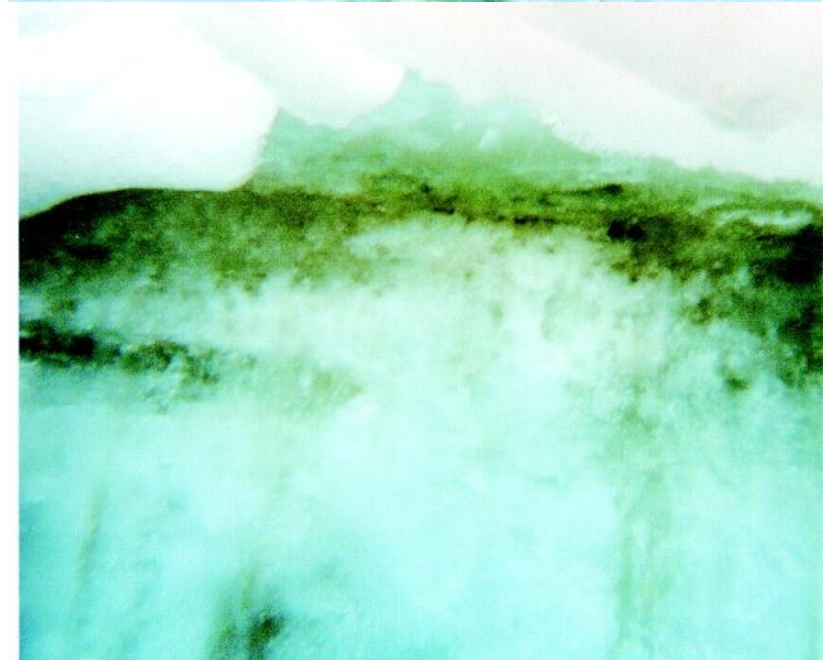


marine ammonia-oxidizing archaeon
Nitrosopumilus assigned to a new phylum:
Thaumarchaeota

Genomics

What Can Genomes Tell Us?

- How do certain organism are able to exist in their environments
- Determine adaptation strategies
- How do certain organism contribute to biogeochemical cycles



Sequencing and Annotating Genomes

- **Sequencing:** determining the precise order of nucleotides in a DNA or RNA molecule
- **Genome annotation:** converting raw sequence data into a list of genes and other functional sequences present in the genome
- **Bioinformatics:** storing and analyzing sequences and structures of nucleic acids and proteins
- Annotation is “bottleneck” in genomics

Sequencing and Annotating Genomes

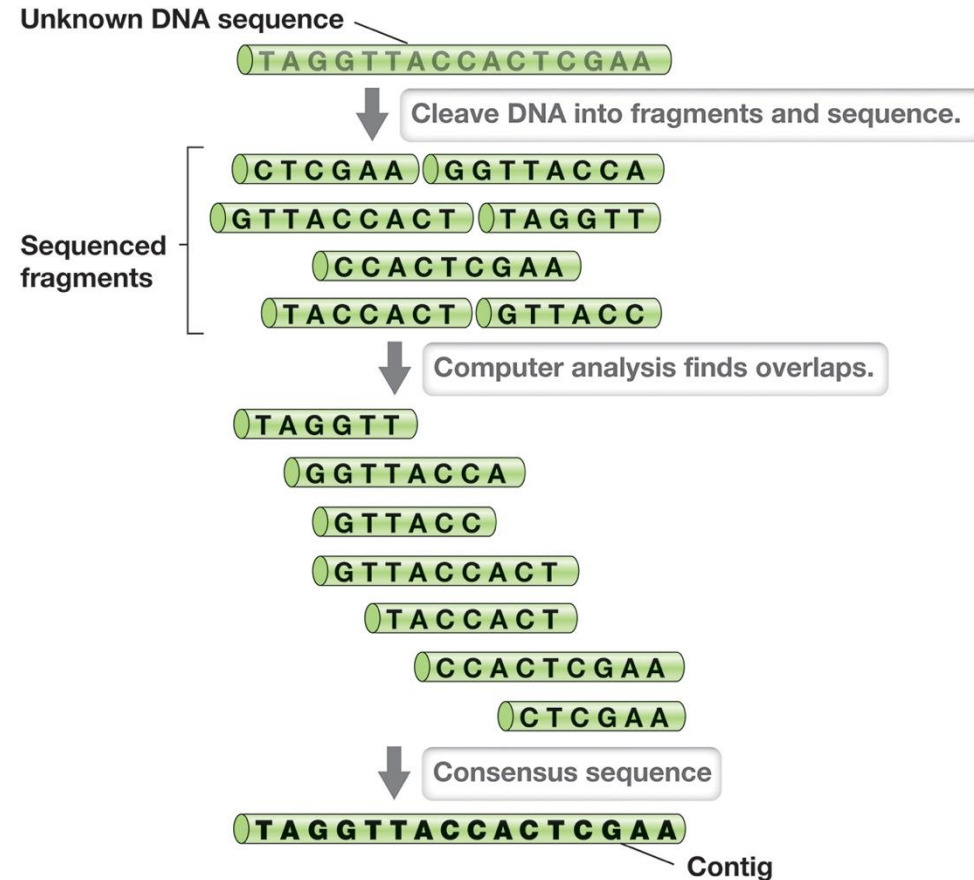
Genome Assembly and Annotation

Genome assembly consists of putting fragments in the correct order and eliminating overlaps

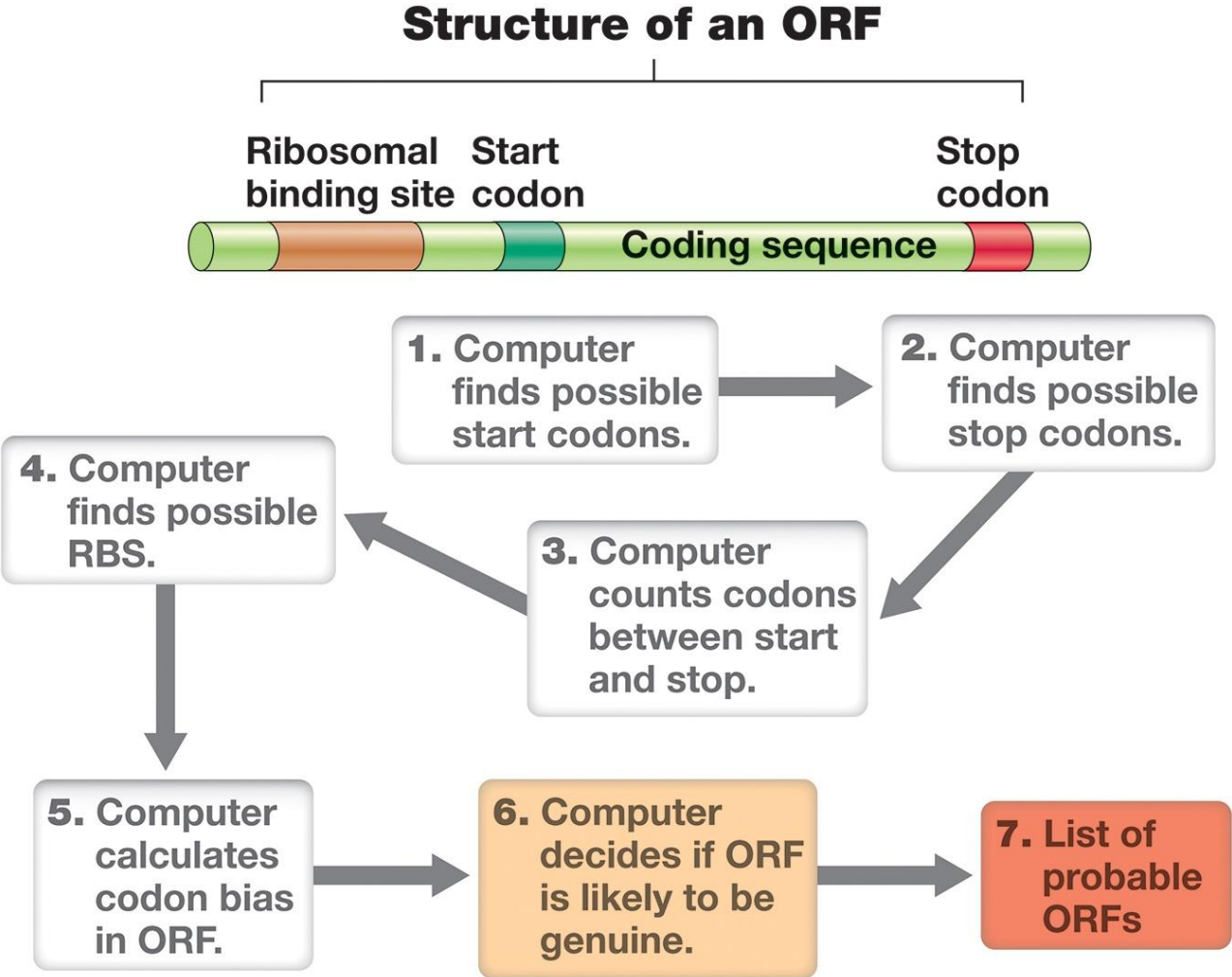
Annotation identifies genes

Highly computational

- Map of complete genome is generated
- Bacterial and archaeal genomes are a series of open reading frames (ORFs)
- Functional ORF: encodes a protein



Computer Identification of Possible ORFs



Sequencing and Annotating Genomes

Never 100 percent identification; many genes encode proteins of unknown function

Number of genes with role that can be clearly identified in a given genome is ~70% or less of total ORFs detected

Hypothetical proteins: uncharacterized ORFs; proteins that likely exist but whose function is currently unknown

- lack sufficient amino acid sequence homology with known proteins for identification
- assign to family or general function

Sequencing and Annotating Genomes

Some genes encode RNA that is not translated (noncoding RNA)

- lack start codons and may have multiple stop codons
 - transfer RNA (tRNA)
 - ribosomal RNA (rRNA)
 - noncoding regulatory RNA molecules

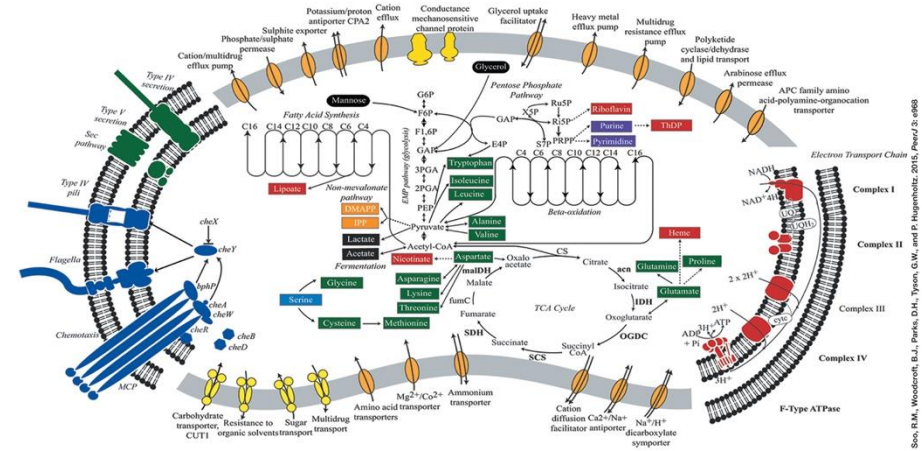
Gene Content of Bacterial Genomes

Complement of genes in a particular organism defines its capabilities

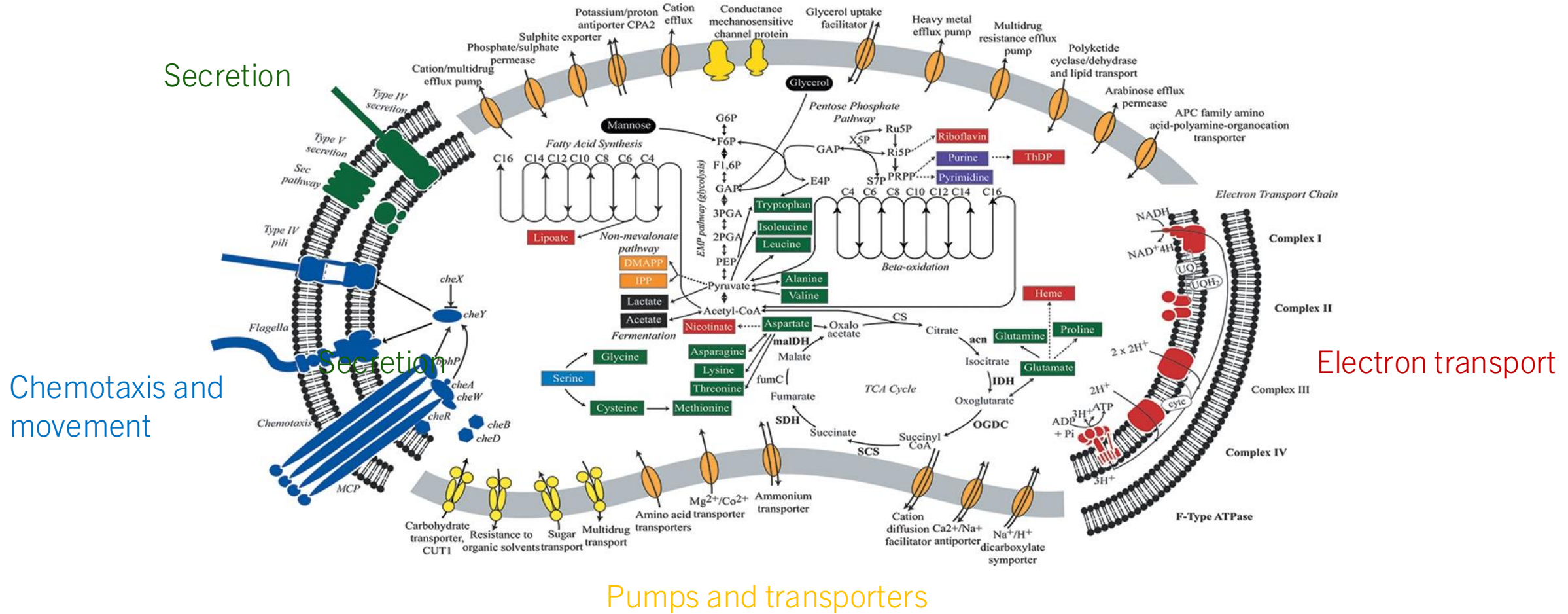
Genomes are also influenced by an organism's lifestyle (or vice-versa?)

Enzymes in the genome can be related to lifestyle of the microorganism

Metabolic genes typically most abundant in bacterial genomes



Functional and Metabolic Predictions for *Vampirovibrio chlorellavorus* Based on Genomic Annotation



Gene Function in Some Genomes of *Bacteria*

Functional categories	Percentage of genes <i>Escherichia coli</i> (4.64 Mbp) ^a	Percentage of genes <i>Haemophilus influenzae</i> (1.83 Mbp) ^a	Percentage of genes <i>Mycoplasma genitalium</i> (0.58 Mbp) ^a
Metabolism	21.0	19.0	14.6
Structure	5.5	4.7	3.6
Transport	10.0	7.0	7.3
Regulation	8.5	6.6	6.0
Translation	4.5	8.0	21.6
Transcription	1.3	1.5	2.6
Replication	2.7	4.9	6.8
Other, known	8.5	5.2	5.8
Unknown	38.1	43.0	32.0

^aChromosome size, in megabase pairs. Each organism listed contains only a single circular chromosome.

PROJECT REPORT

- Research paper

Based on data collected during the course

Intro, Methods, Results, Discussion, Conclusion, References

- 25 % of the grade
- In groups or individual

PROJECT REPORT

- 25 % of the grade
- In groups or individual

Isolated microorganism

What is its taxonomy?

What kind of metabolism does it have?

How does it acquire Nitrogen and Carbon?

Can it produce greenhouse gases? Which?

What kind of metabolites does it make/excrete?

Can it detoxify metal contamination?

Does it have antibiotic resistance genes?

How can it potentially interact with other organisms?

Field Data

Use the data gathered in the field to contextualize the analysis of your isolate.

- Does the chemistry/temperature/gas flux of the environment where you isolated your organism from make sense with what you know about your organism microorganism (from your results and literature)?

Use published literature and your results.
Compare results to published data in the Discussion