

# Molecular Methods in Microbial Ecology

**Contact Info:** Kristin Gribble, [kgribble@mbi.edu](mailto:kgribble@mbi.edu)

## **Schedule:**

Tuesday	10/23/18	Introduction, Extraction of DNA from Winogradsky columns Run DNA products on gel
Thursday	10/25/18	Lecture on PCR, Prepare PCR reactions
Tuesday	10/30/17	Analyze PCR results, Lecture

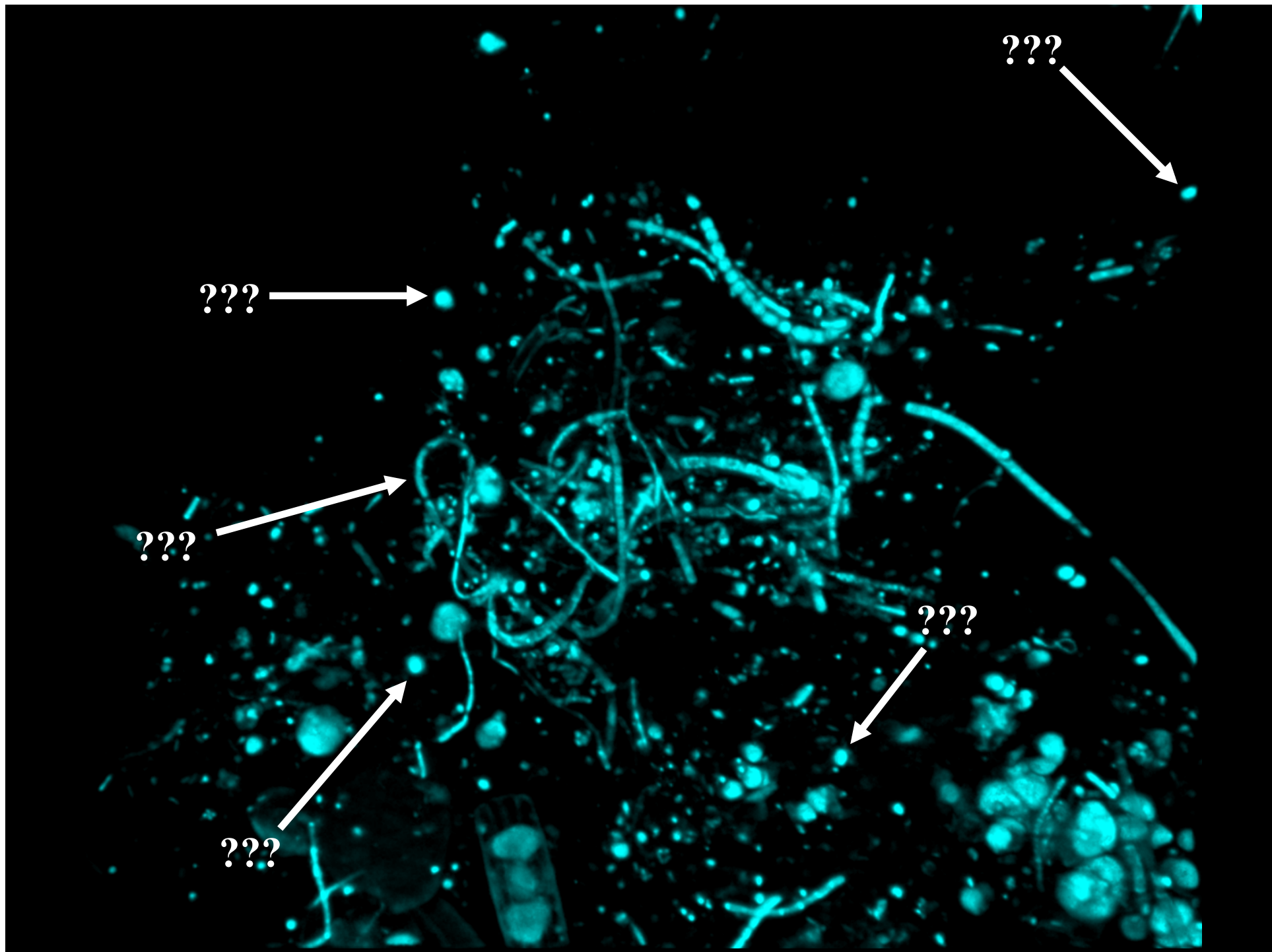
**Readings:** Head *et al.* 1998. *Microbial Ecology* 35: 1-21.

# Day 1

- Introduction to molecular methods in microbial ecology
- Extract DNA from Winogradsky Columns
- Run DNA on agarose gel

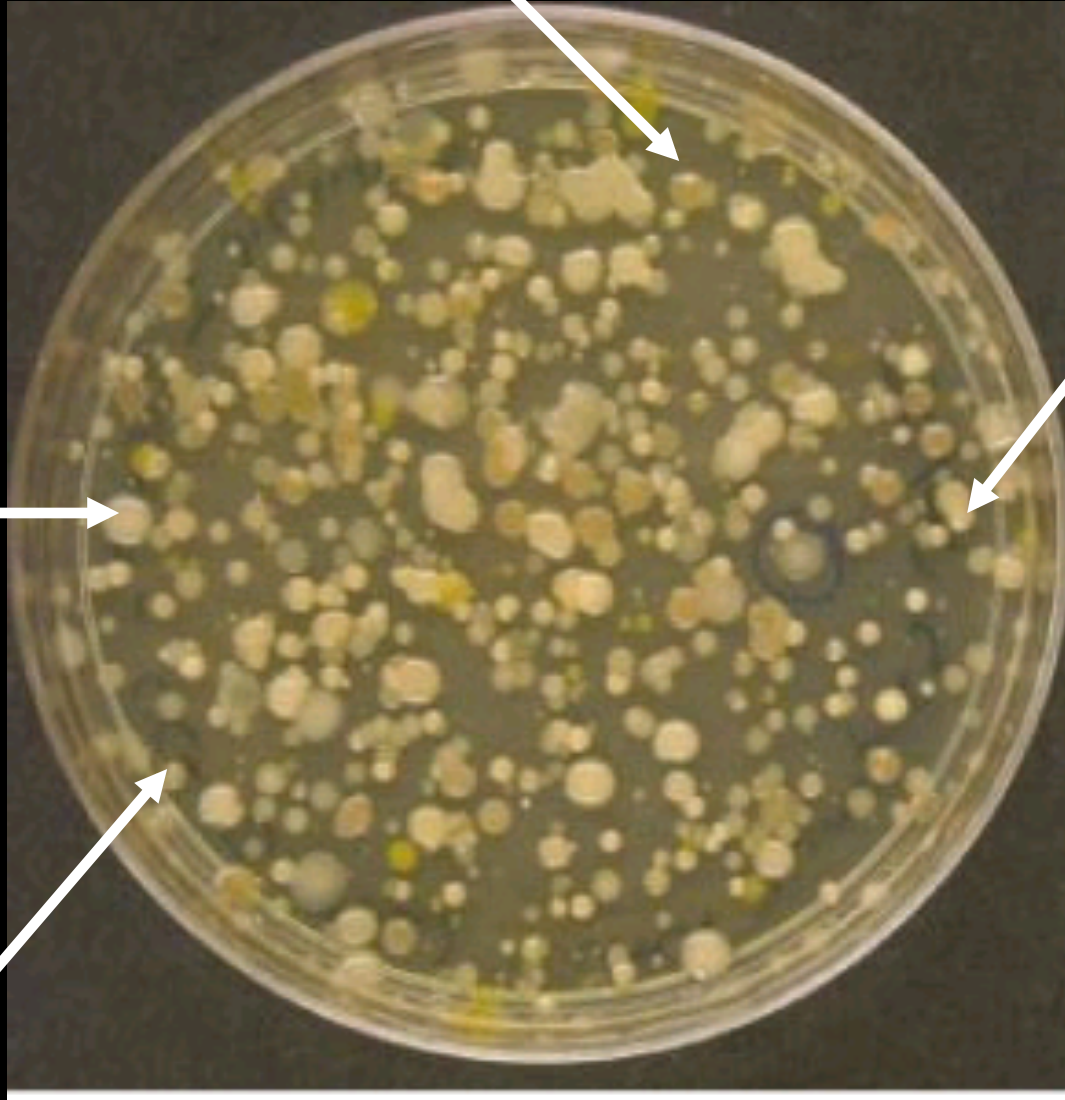








???



???

???

???

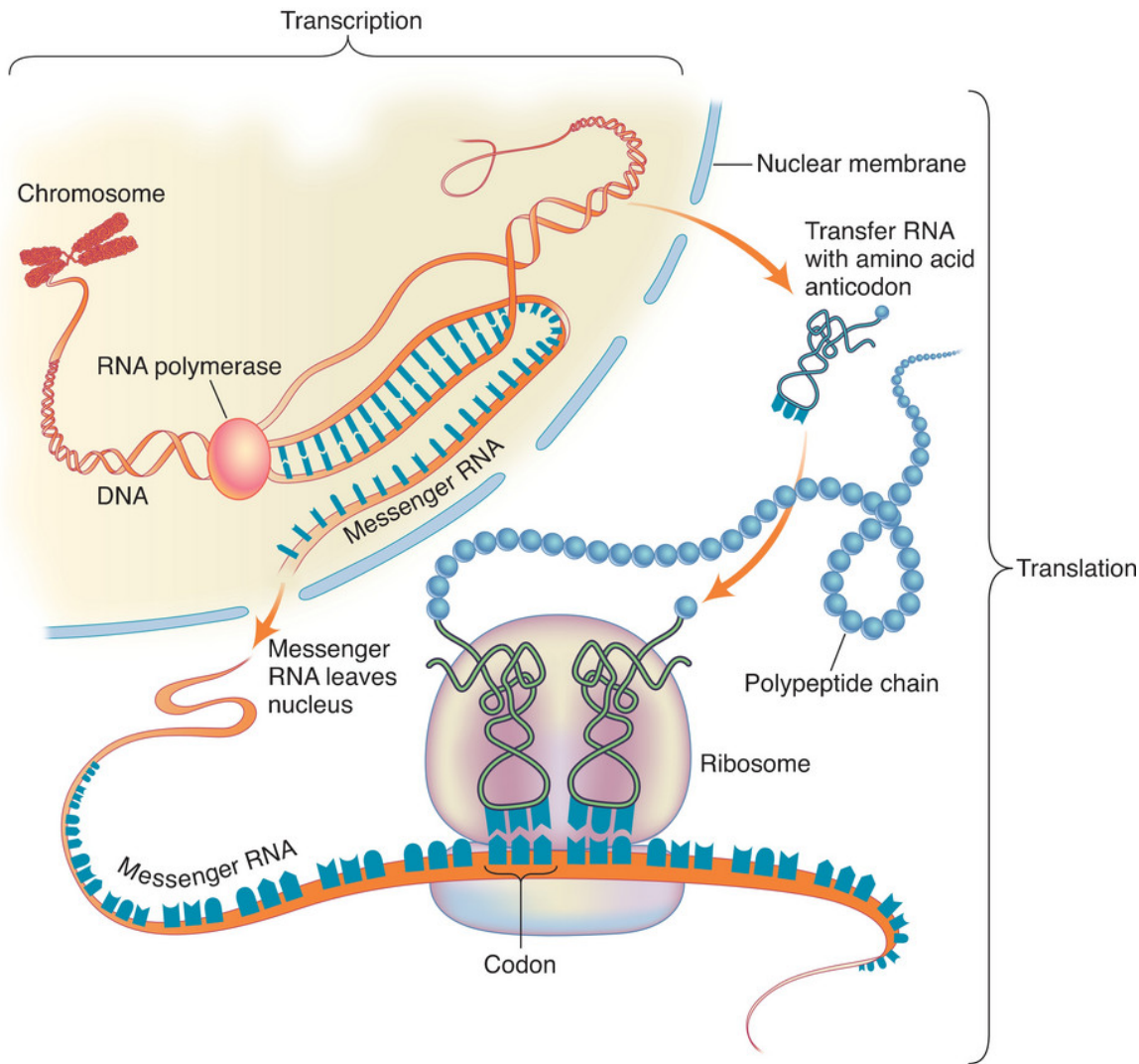
# The Challenge for Microbial Ecology

<b>Habitat</b>	<b>Culturability (%)</b>
Seawater	0.001-0.1
Freshwater	0.25
Sediments	0.25
Soil	0.3

**How do you study something  
you can't grow in the lab?**

From Amann *et al.* 1995 Microbiological Reviews

# The Central Dogma



**DNA**



Transcription

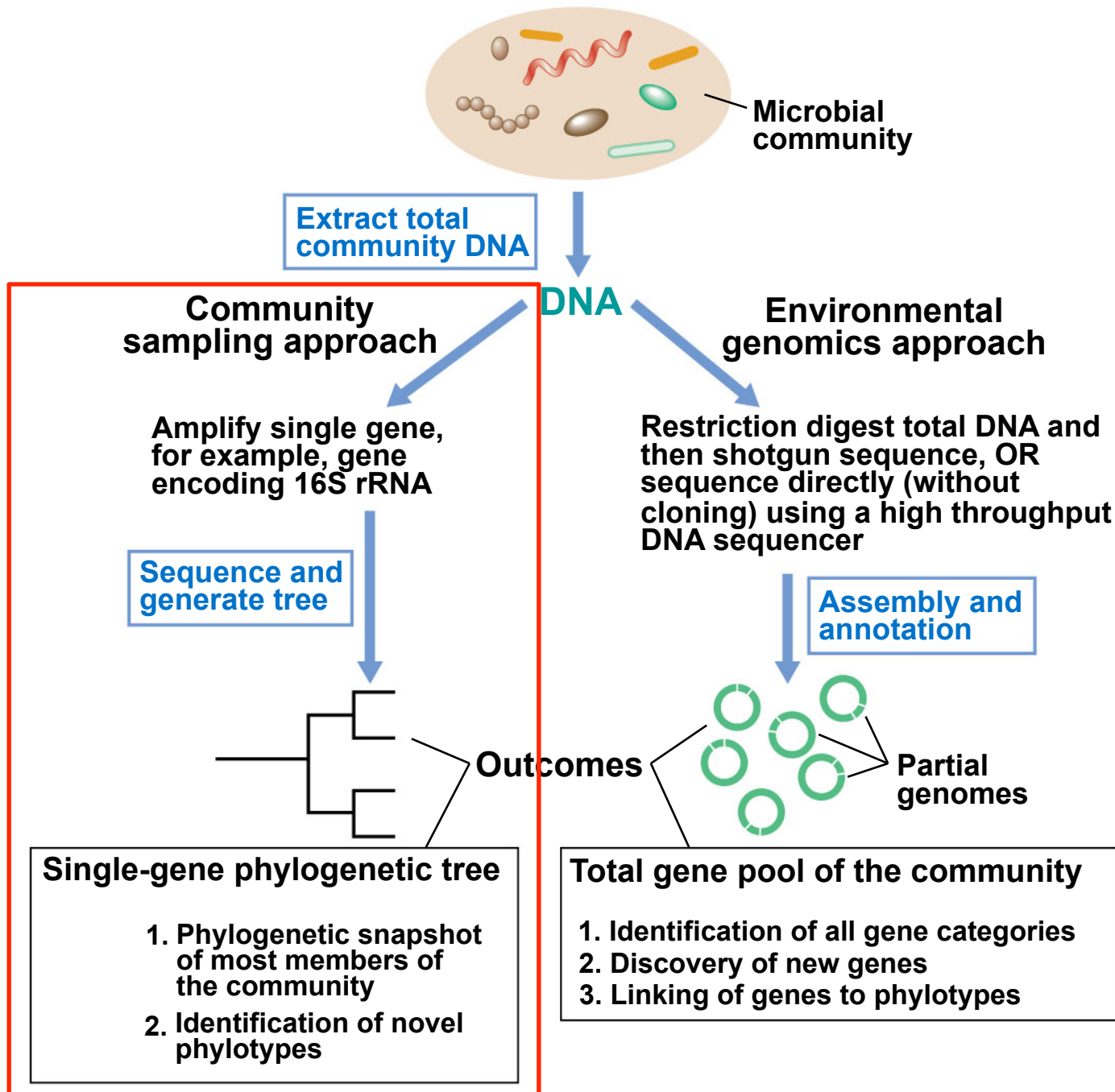
**RNA**

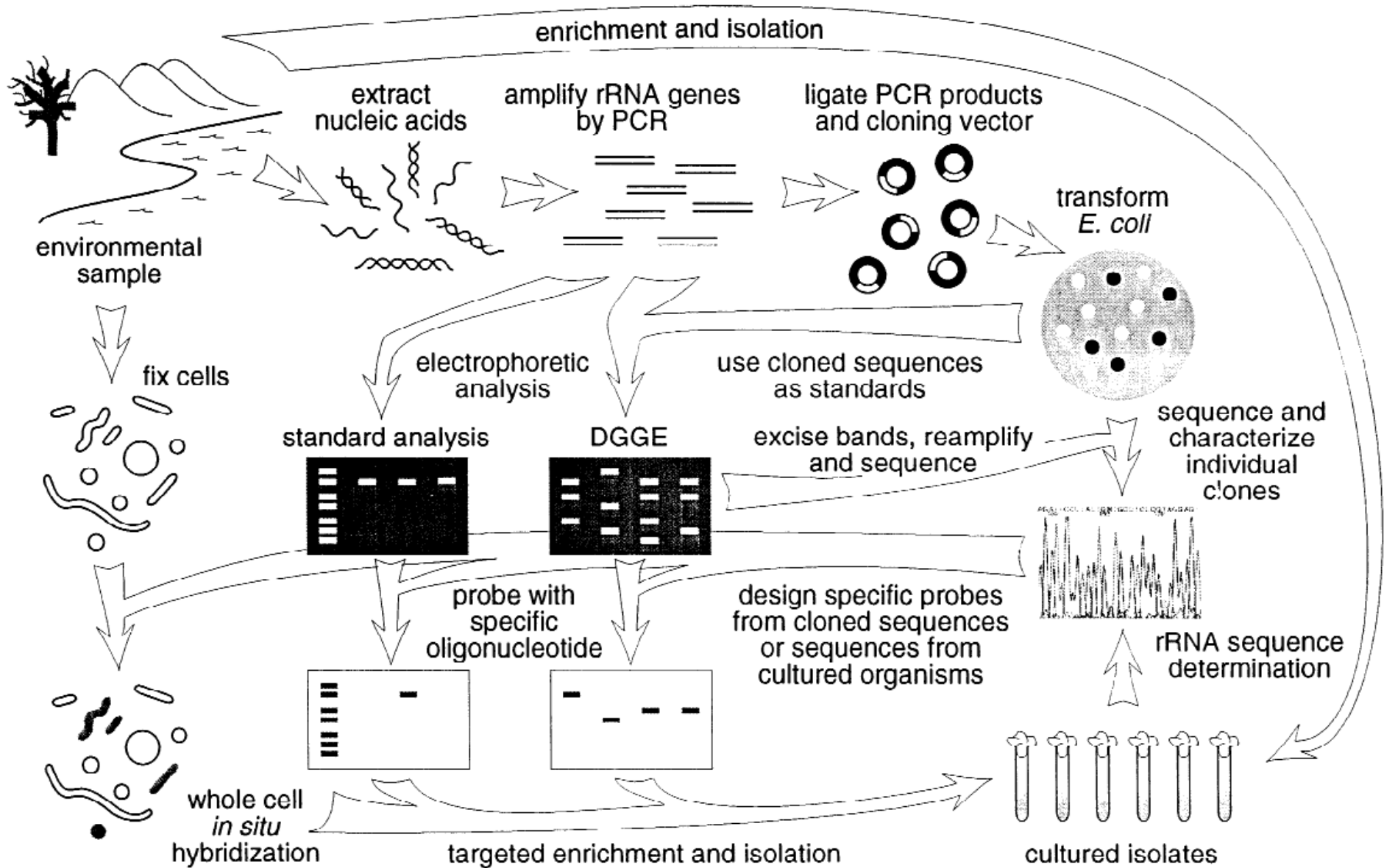


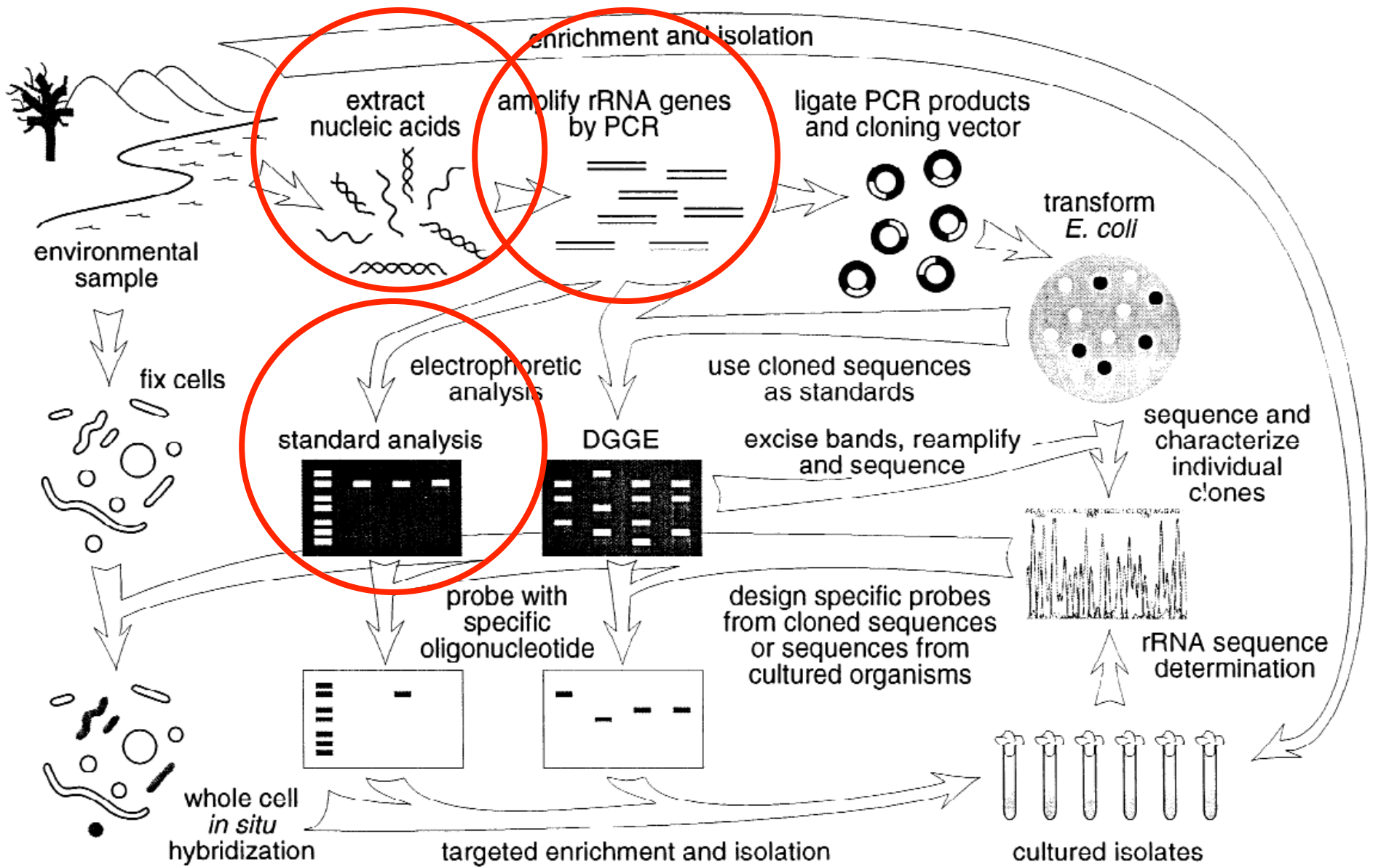
Translation

**Protein**



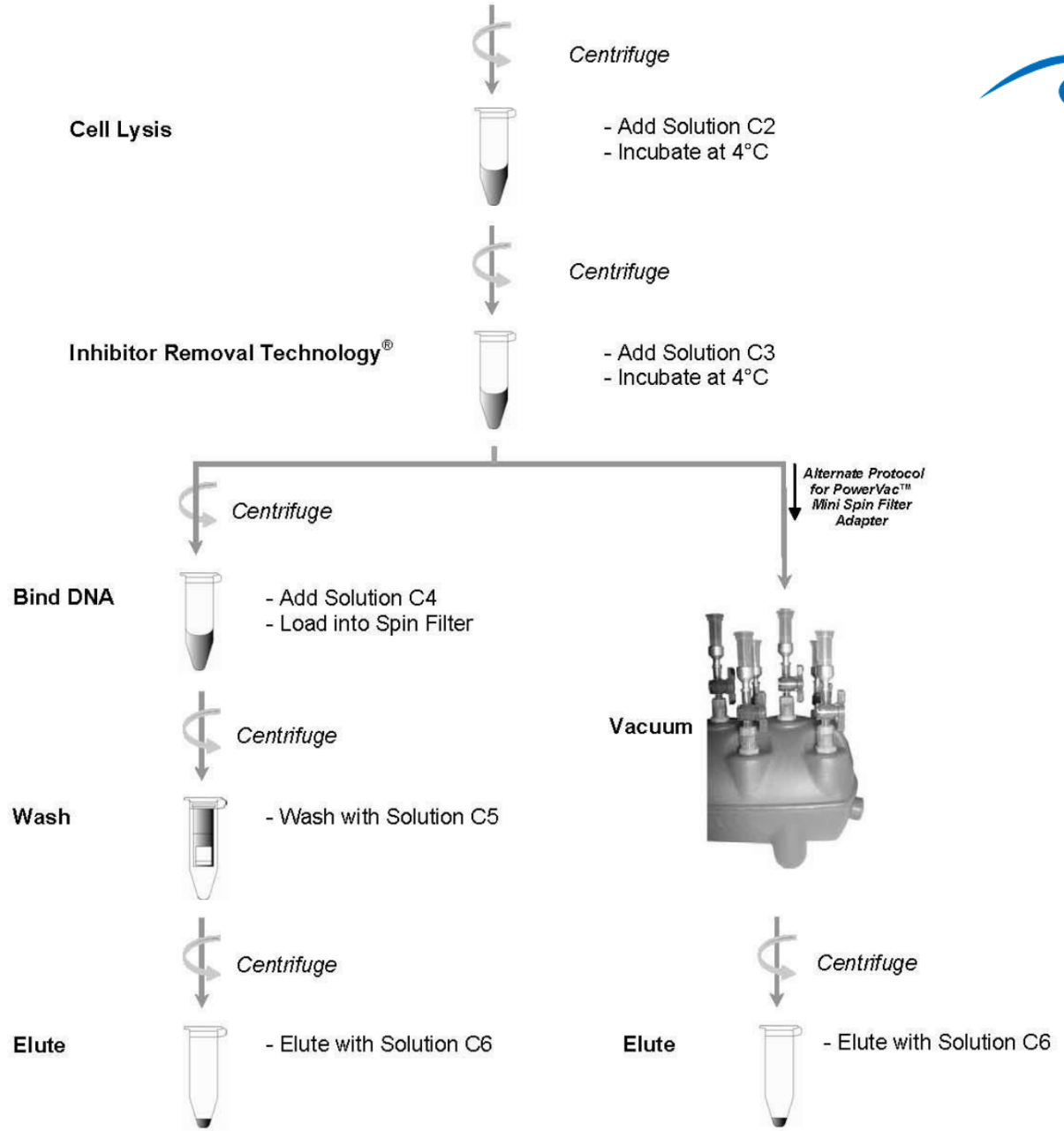








# DNA Extraction Overview

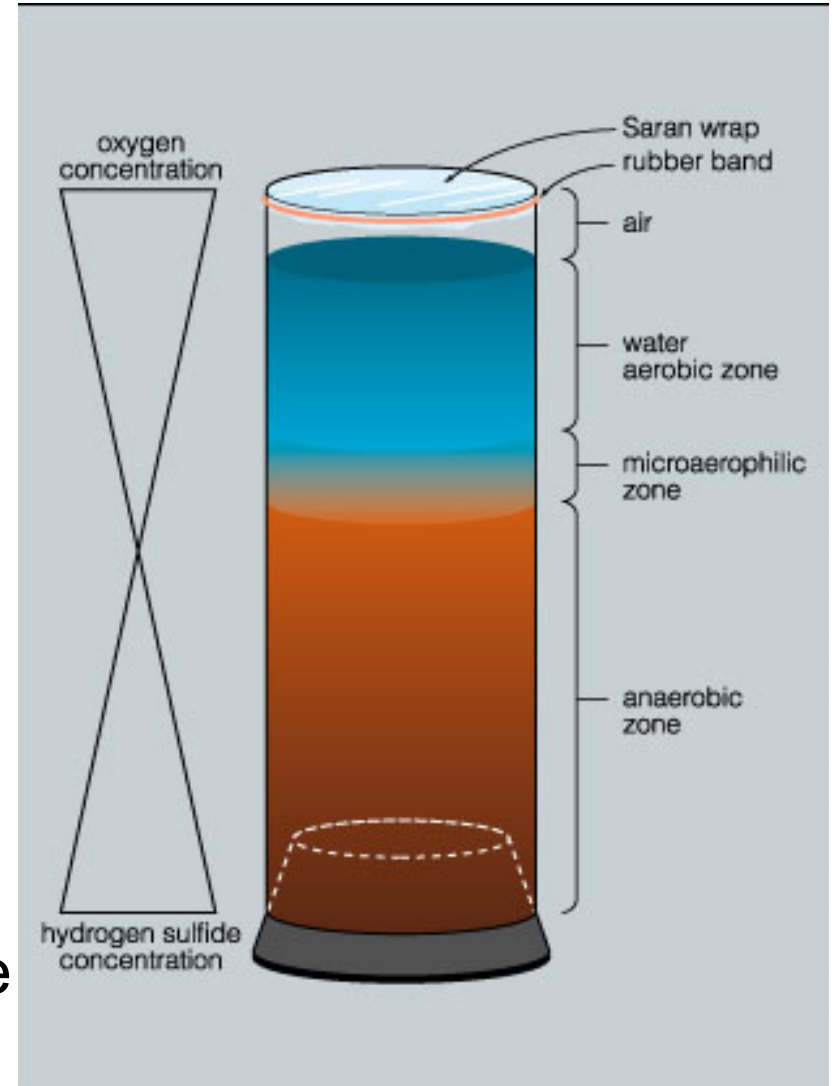


# DNA Extraction

1. Lyse cell membrane
  - a. Chemically → detergent
  - b. Physically → bead beating
2. Pellet cell membrane, proteins and other cell parts while DNA stays in solution
3. Remove other inhibitors from DNA
4. Mix DNA with acid and salt → stick to filter
5. Wash filter-bound DNA several times with alcohol
6. Elute DNA off membrane with pH 8, low-salt buffer

# Choosing a Depth Horizon

- 16S rRNA Bacteria
- 16S rRNA Archaea
- *mcrA* Methanogens
  - Methyl coenzyme M reductase
- *dsrB* Sulfate reducers
  - Dissimilatory bisulfite reductase



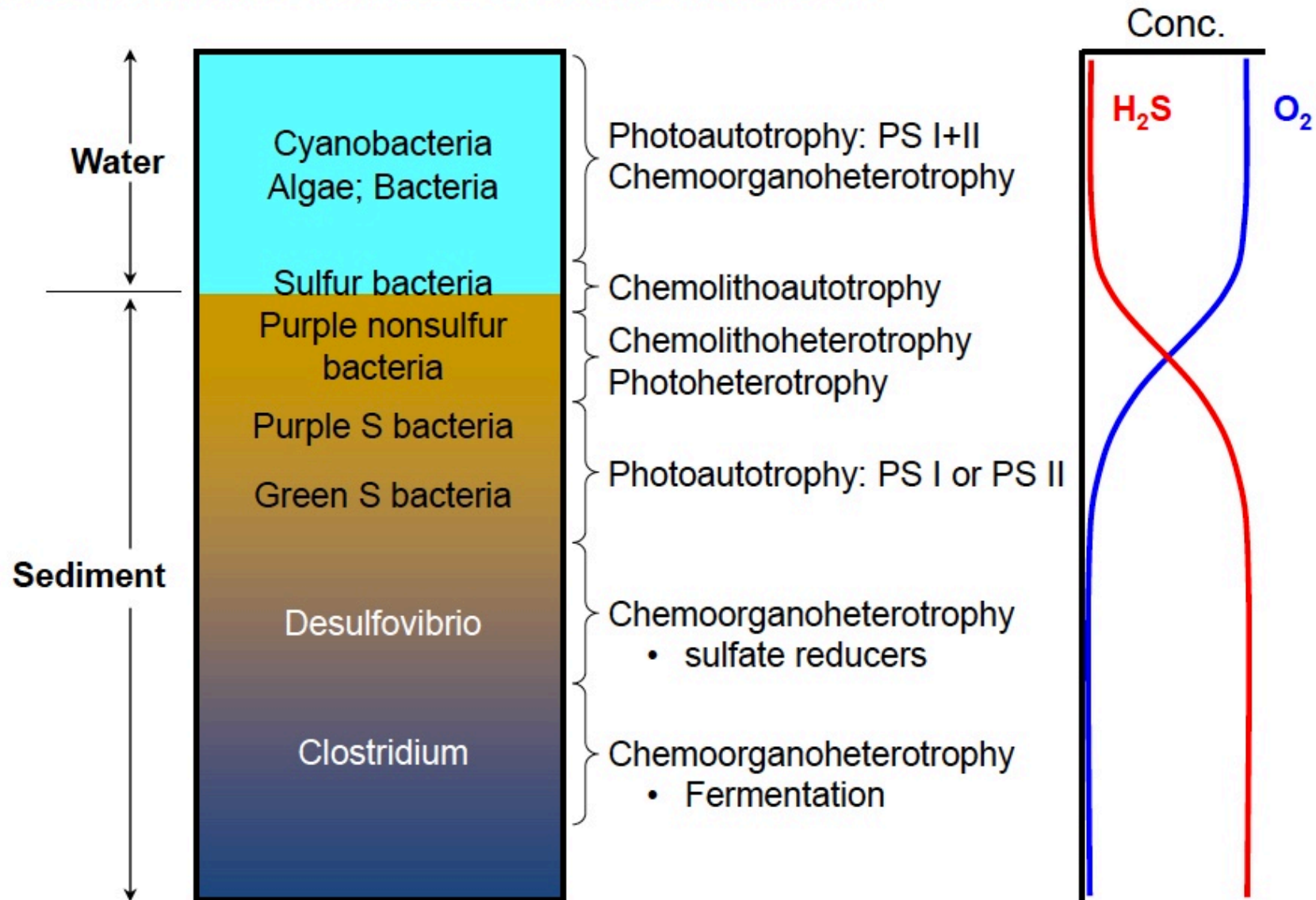






# Winogradsky Column

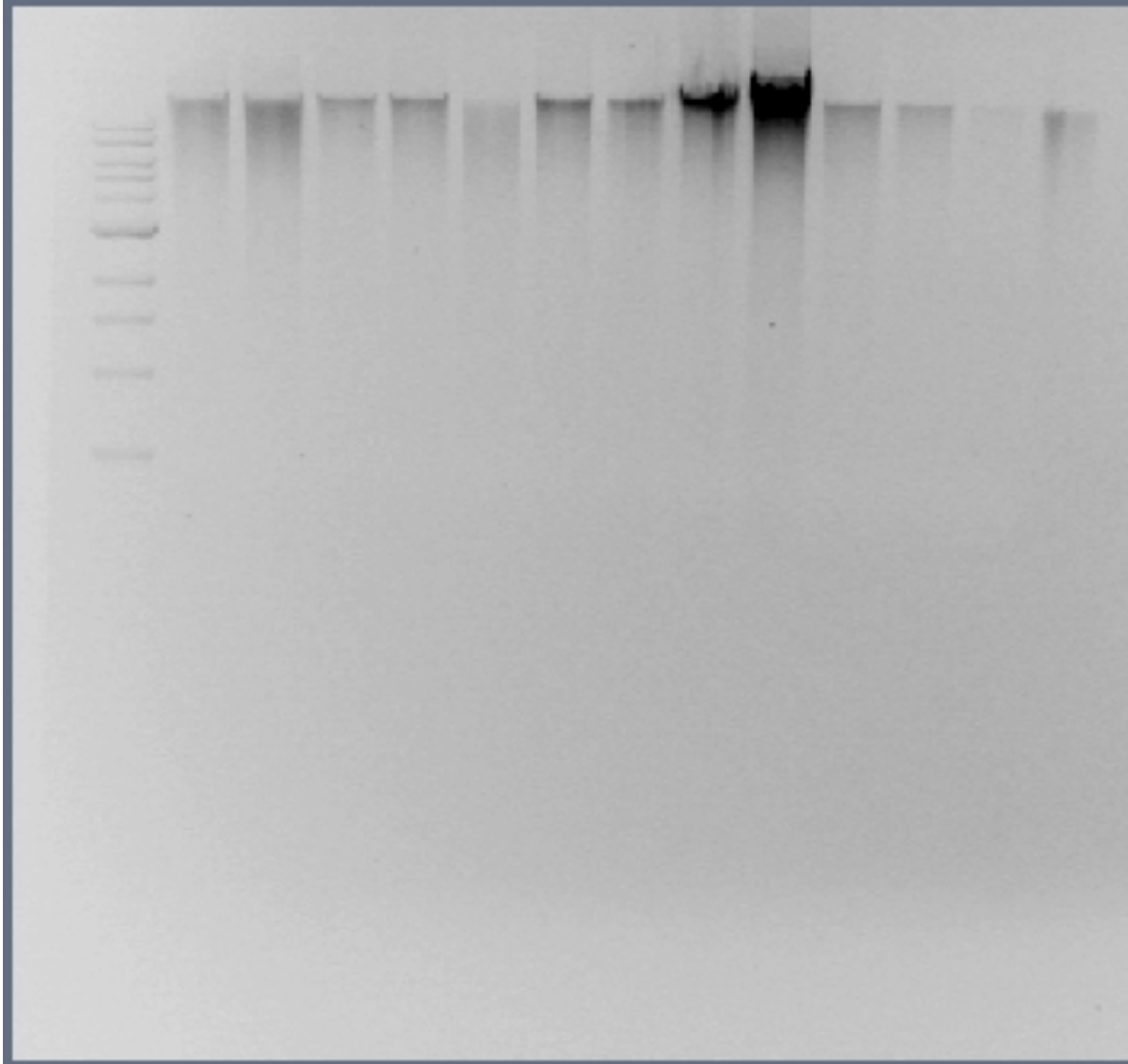
Microenvironments generated by chemical gradients.





# Day 1, Part II

Run an electrophoresis gel of the DNA products extracted from your columns



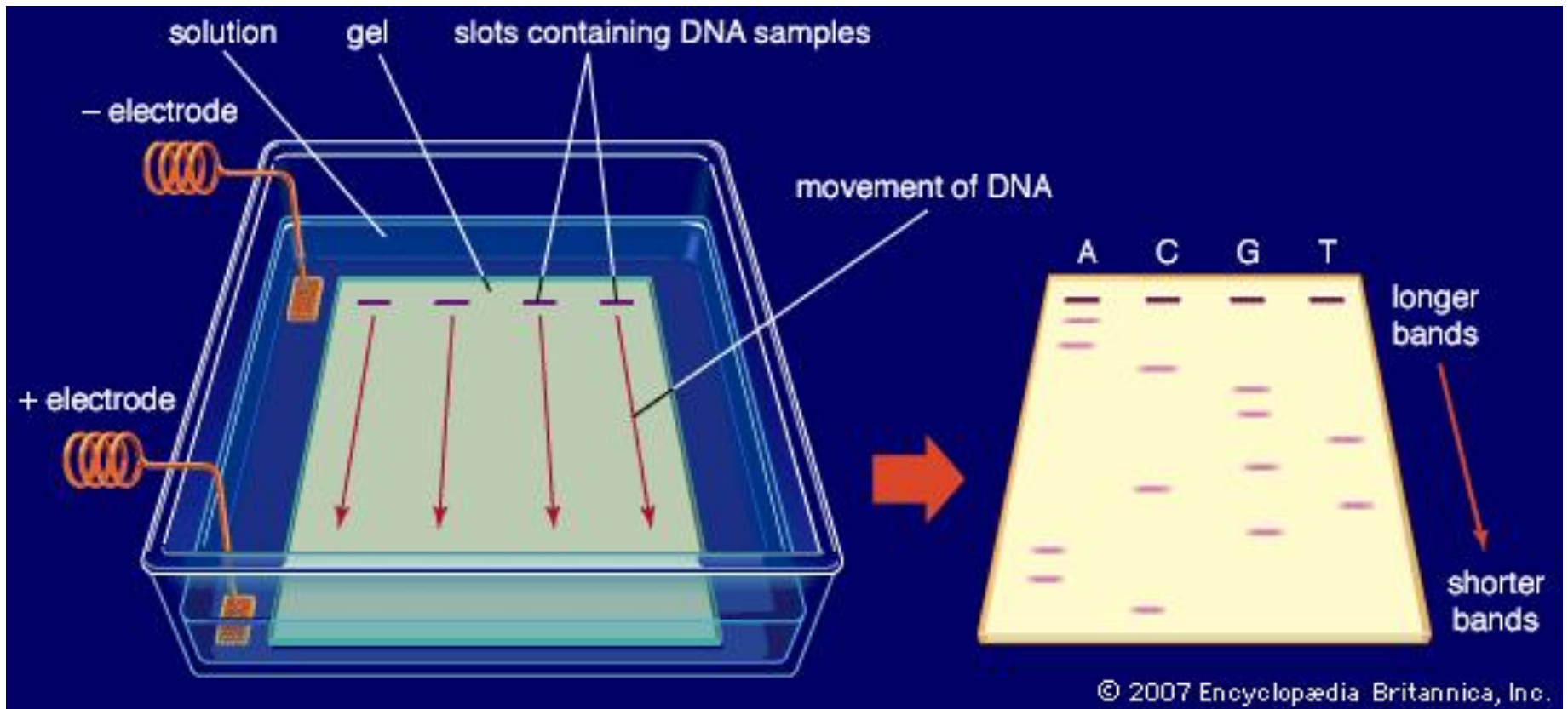
## **Genomic DNA**

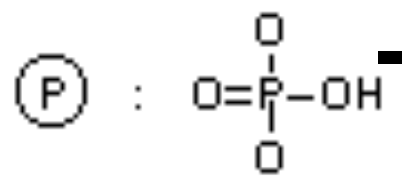
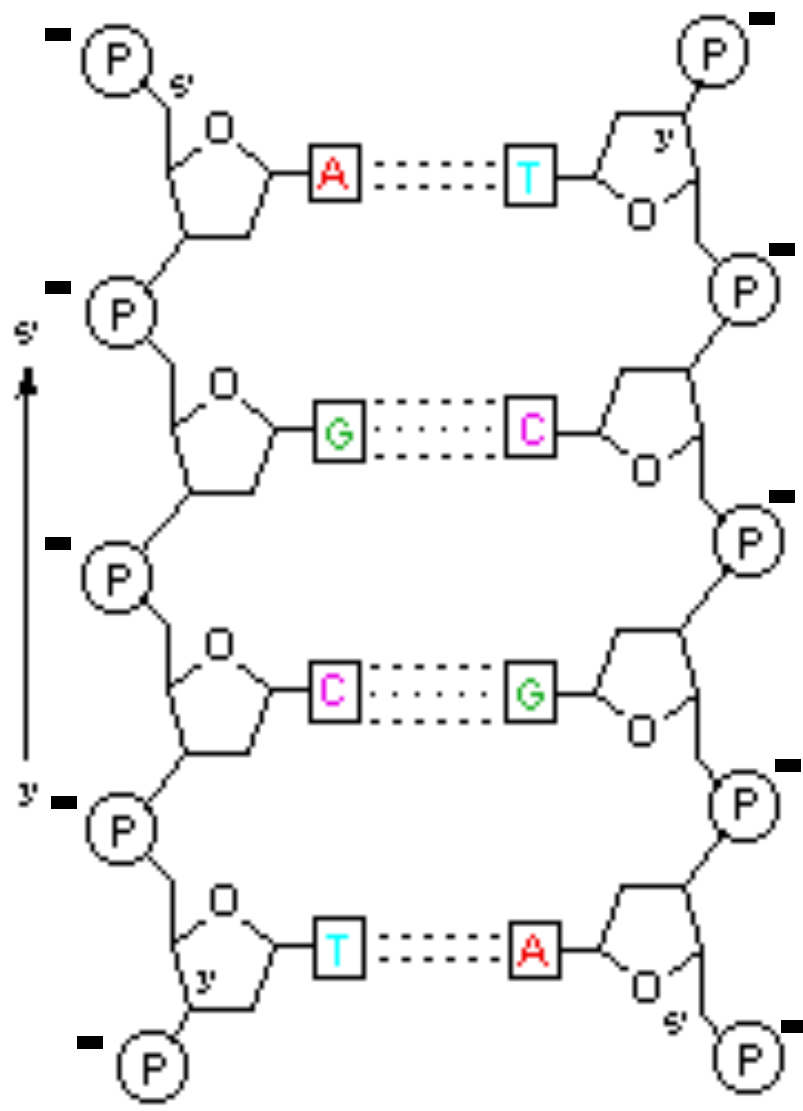
The sum total of all DNA from an organism or a community of organisms

# Basics of Gel Electrophoresis

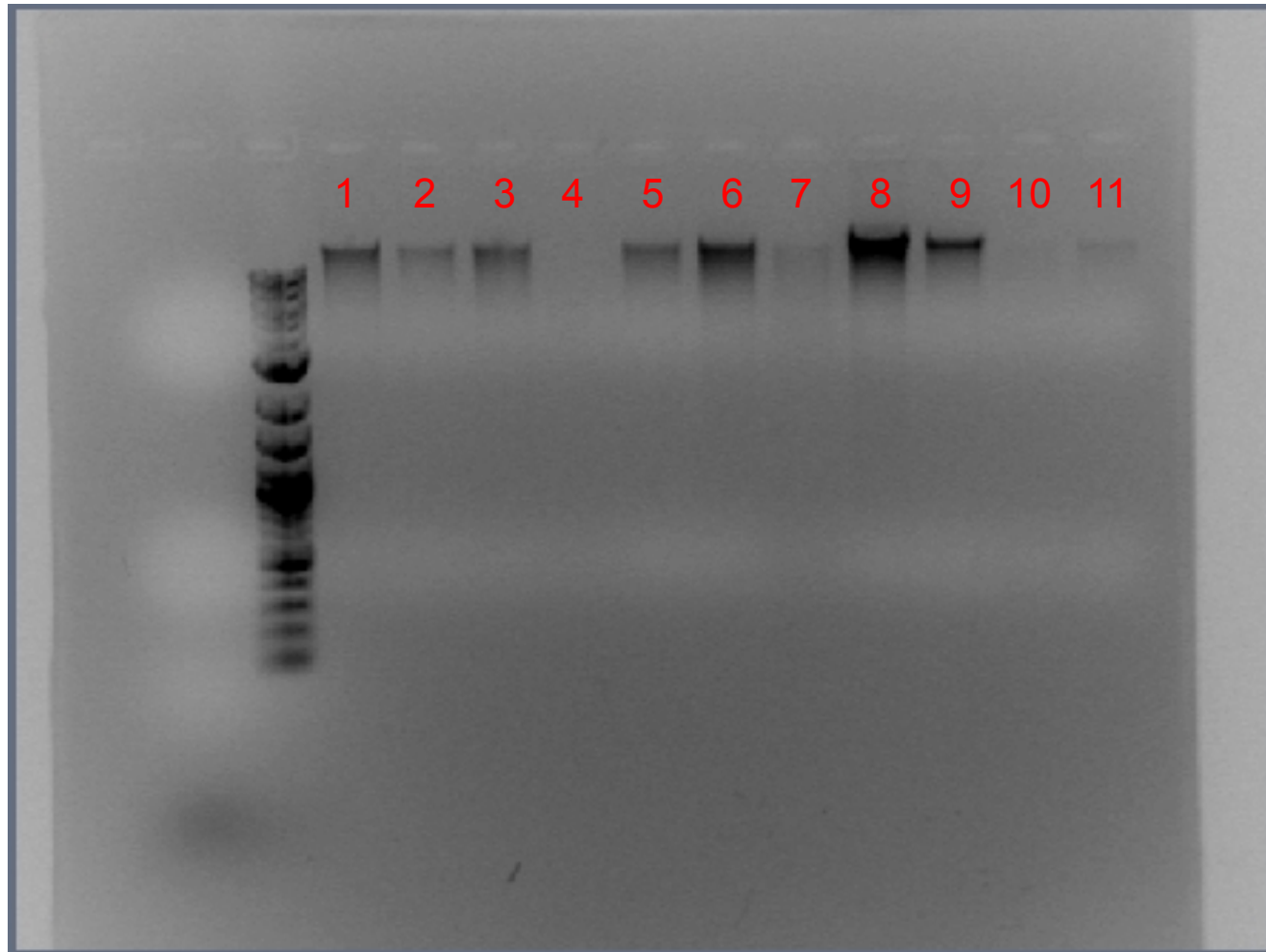
- The gel is a matrix (like jello with holes)
- DNA is negatively charged- will run to positive
- Smaller fragments run faster than larger ones
- Gel contains Ethidium Bromide, which binds to DNA and fluoresces when hit with UV light (WEAR GLOVES!!!)

# Gel electrophoresis





# Gel results . . . .



## **Genomic DNA**

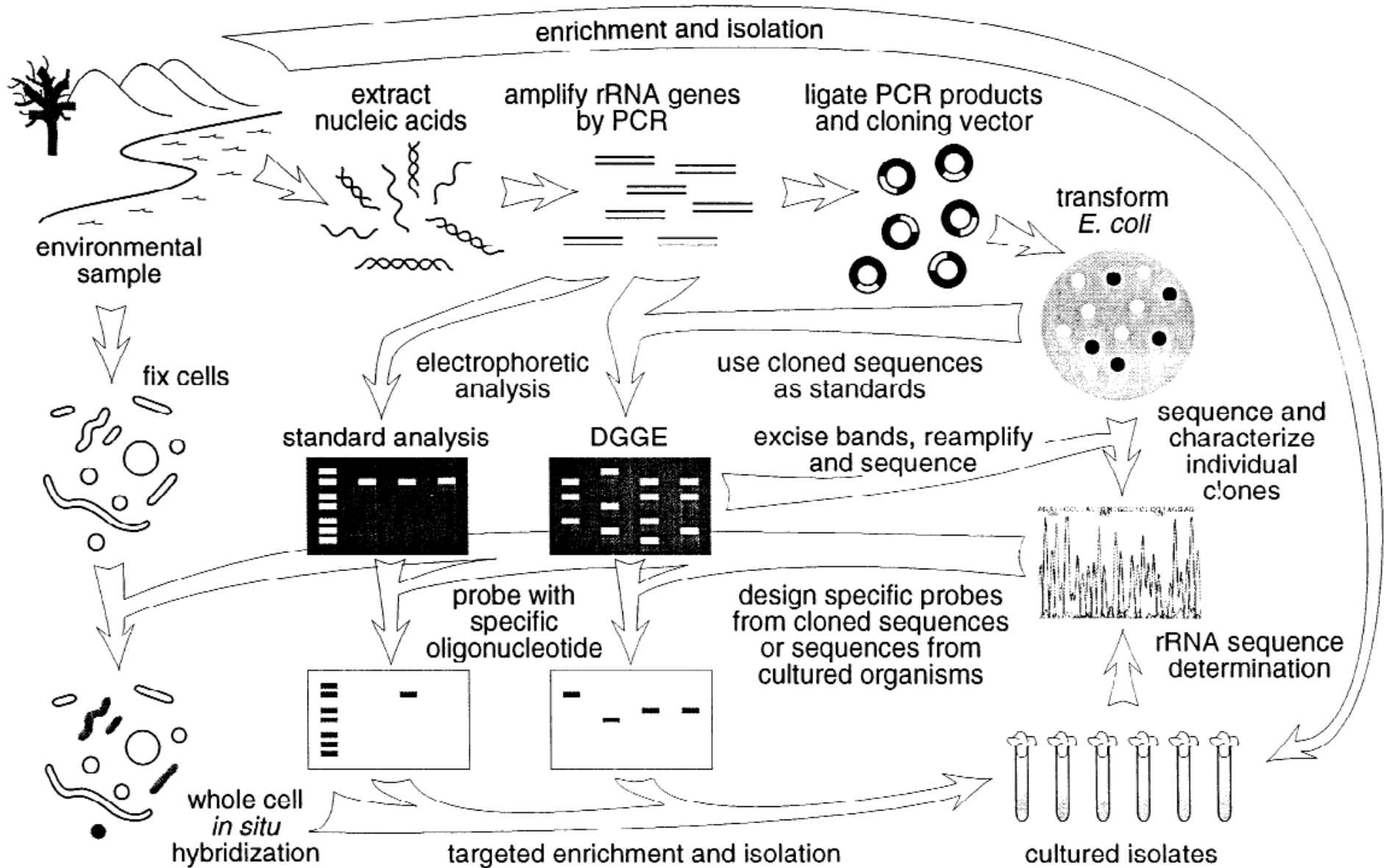
The sum total of all DNA from an organism or a community of organisms

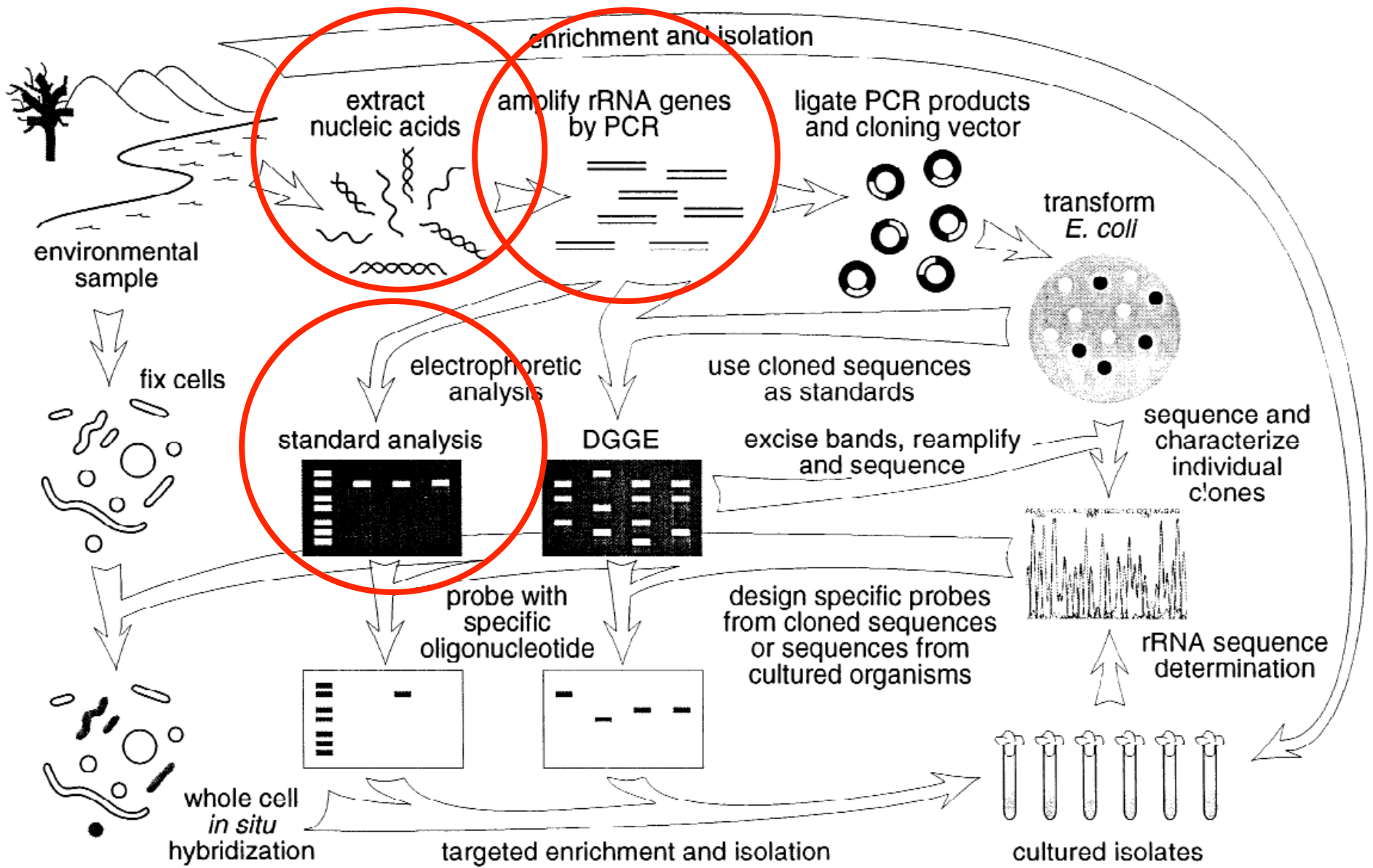




# Day 2

- Basics of PCR
- Set up PCR reactions using the DNA from your extractions and an assortment of primers





# Polymerase Chain Reaction (PCR)

- Rapid, inexpensive and simple way of making millions of copies of a gene starting with very few copies
- Does not require the use of isotopes or toxic chemicals
- It involves preparing the sample DNA and a master mix with primers, followed by detecting reaction products

# The Nobel Prize in Chemistry 1993



Kary B. Mullis

Prize share: 1/2



Michael Smith

Prize share: 1/2

The Nobel Prize in Chemistry 1993 was awarded *"for contributions to the developments of methods within DNA-based chemistry"* jointly with one half to Kary B. Mullis *"for his invention of the polymerase chain reaction (PCR) method"* and with one half to Michael Smith *"for his fundamental contributions to the establishment of oligonucleotide-based, site-directed mutagenesis and its development for protein studies"*.

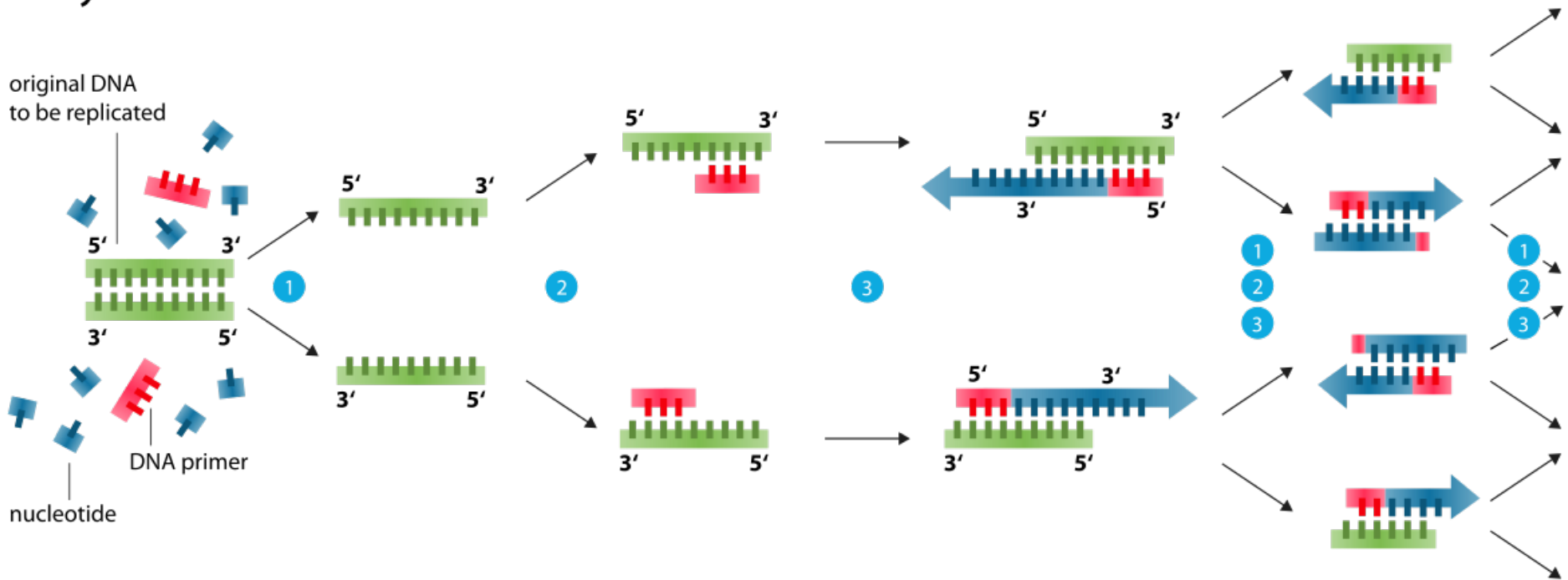


# Polymerase Chain Reaction (PCR)



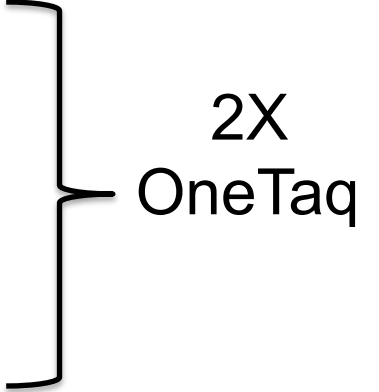
- Takes advantage of properties of Taq DNA polymerase to amplify (make copies of) a selected gene region
- Requirements
  - You must know the sequence flanking the region to be amplified

# Polymerase chain reaction - PCR



- 1 **Denaturation** at 94-96°C
- 2 **Annealing** at ~68°C
- 3 **Elongation** at ca. 72 °C

# Every PCR contains:

- A DNA Polymerase (most common, Taq)
  - Deoxynucleotide Triphosphates (A, C, T, G)
  - Buffer (salt,  $MgCl_2$ , etc)
  - A set of primers, one Forward, one Reverse
  - Various chemicals to minimize inhibition
  - Template DNA
- 
- 2X  
OneTaq

# Typical PCR Profile

<b>Temperature</b>	<b>Time</b>	<b>Action</b>
95°C	5 minutes	DNA Taq polymerase activation
35 cycles of: 95°C 54°C 72°C	1 minute 1 minute 1 minute	DNA denaturization Primer annealing Extension creation
72°C	10 minutes	Final extension created

# Things you can optimize

- Temperature and time to activate Taq polymerase
- Temperature and time to allow primer annealing
- Temperature and time for extension
- Concentration of reagents, especially primers, dNTPs, and  $MgCl_2$
- Concentration of template DNA
- Number of replication cycles
- Etc...

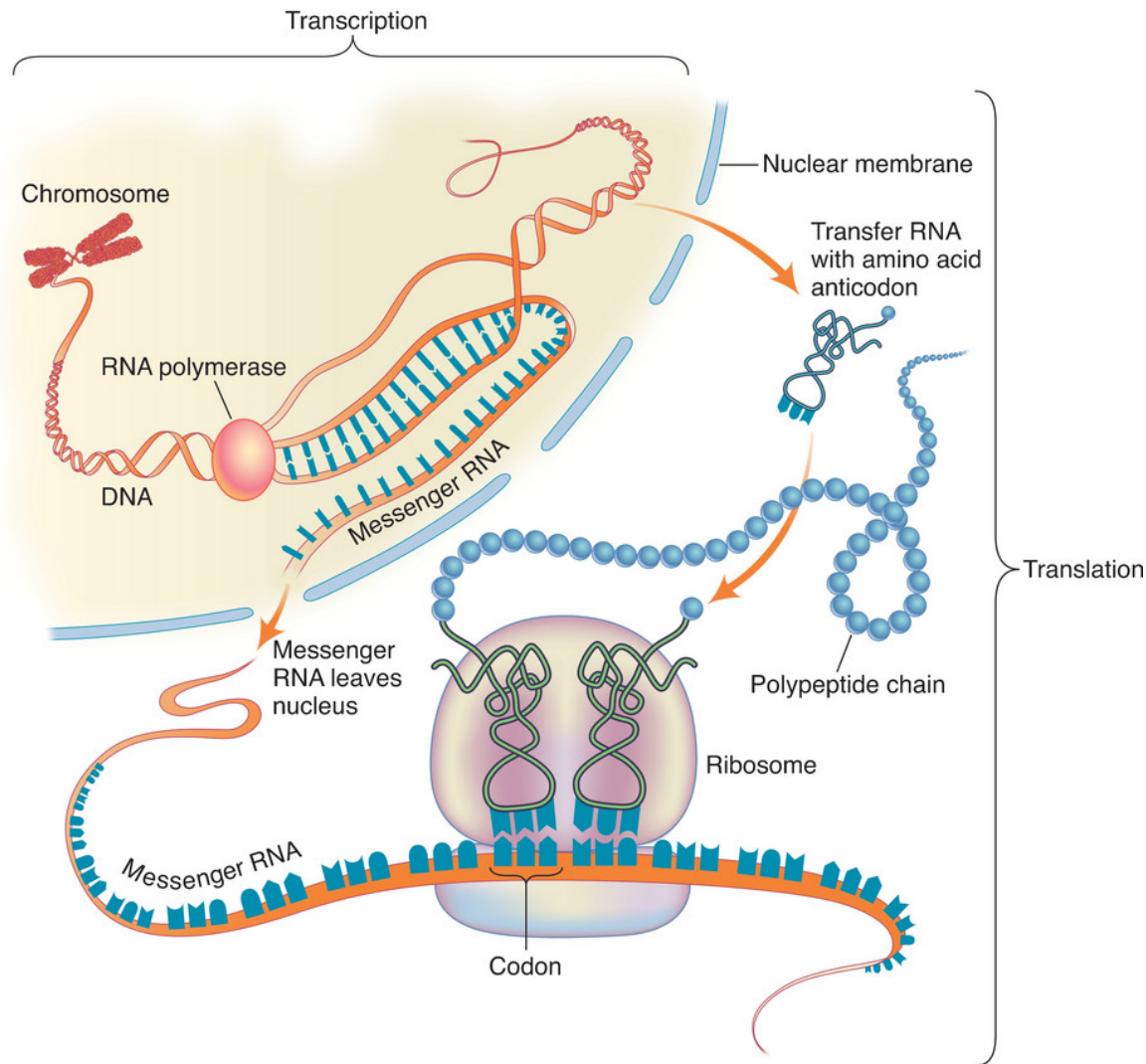
# The Star of the Show: SSU rRNA

- Everybody has it
- Contains both highly conserved and variable regions
  - allows making comparisons between different organisms over long periods of time (evolutionary history)
- Not laterally transferred between organisms
- HUGE and growing database





# The Central Dogma



**DNA**



Transcription

**RNA**

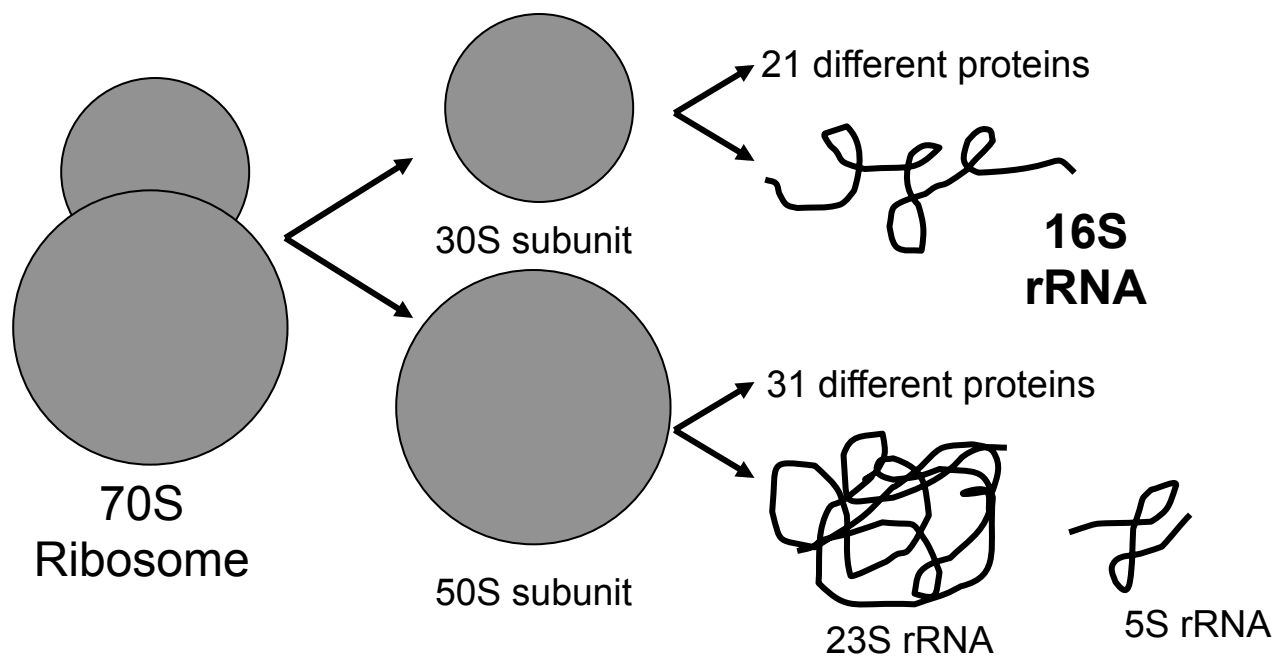


Translation

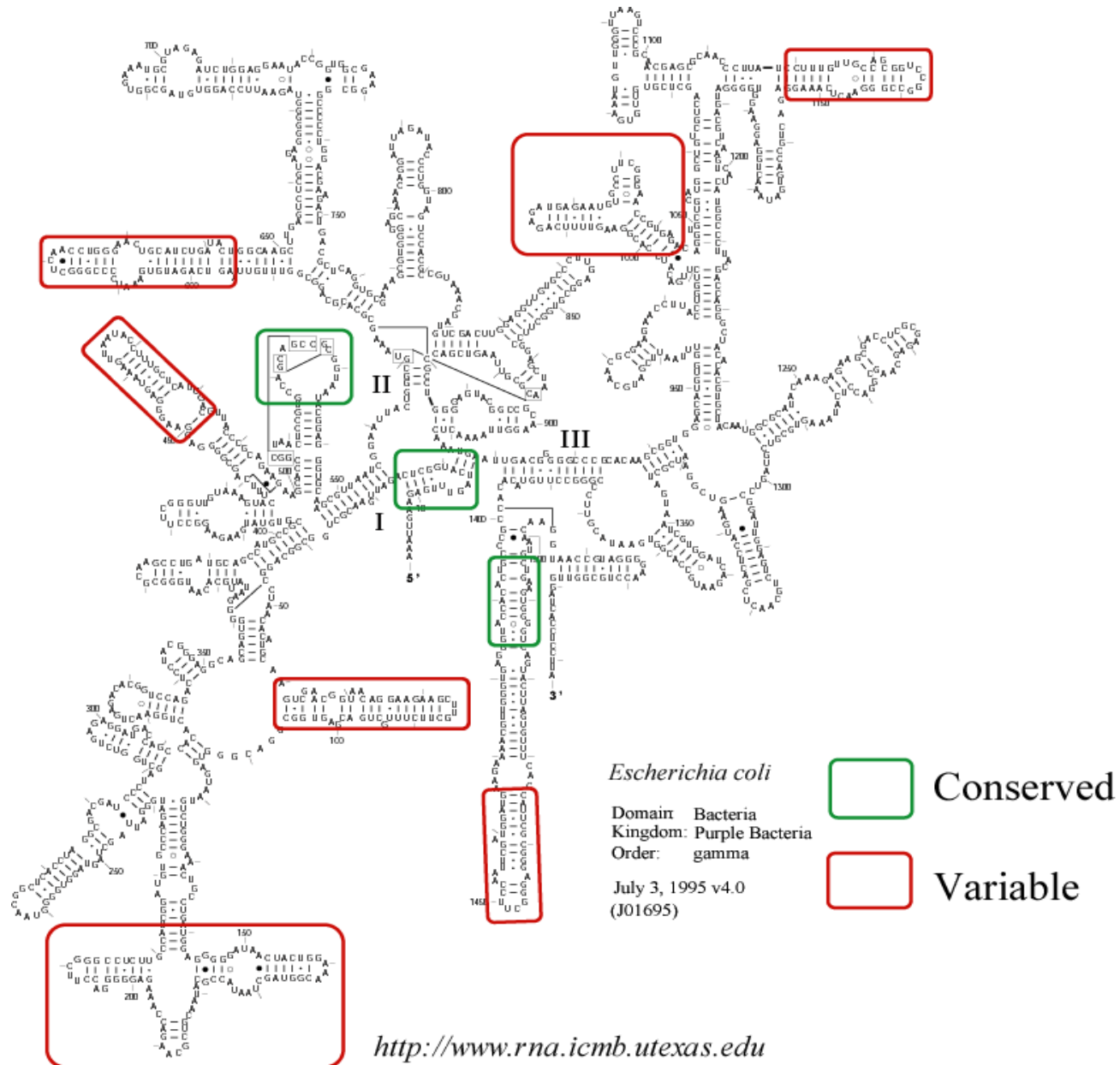
**Protein**

# Ribosomes

- Make proteins
- rRNA is transcribed from rDNA genes



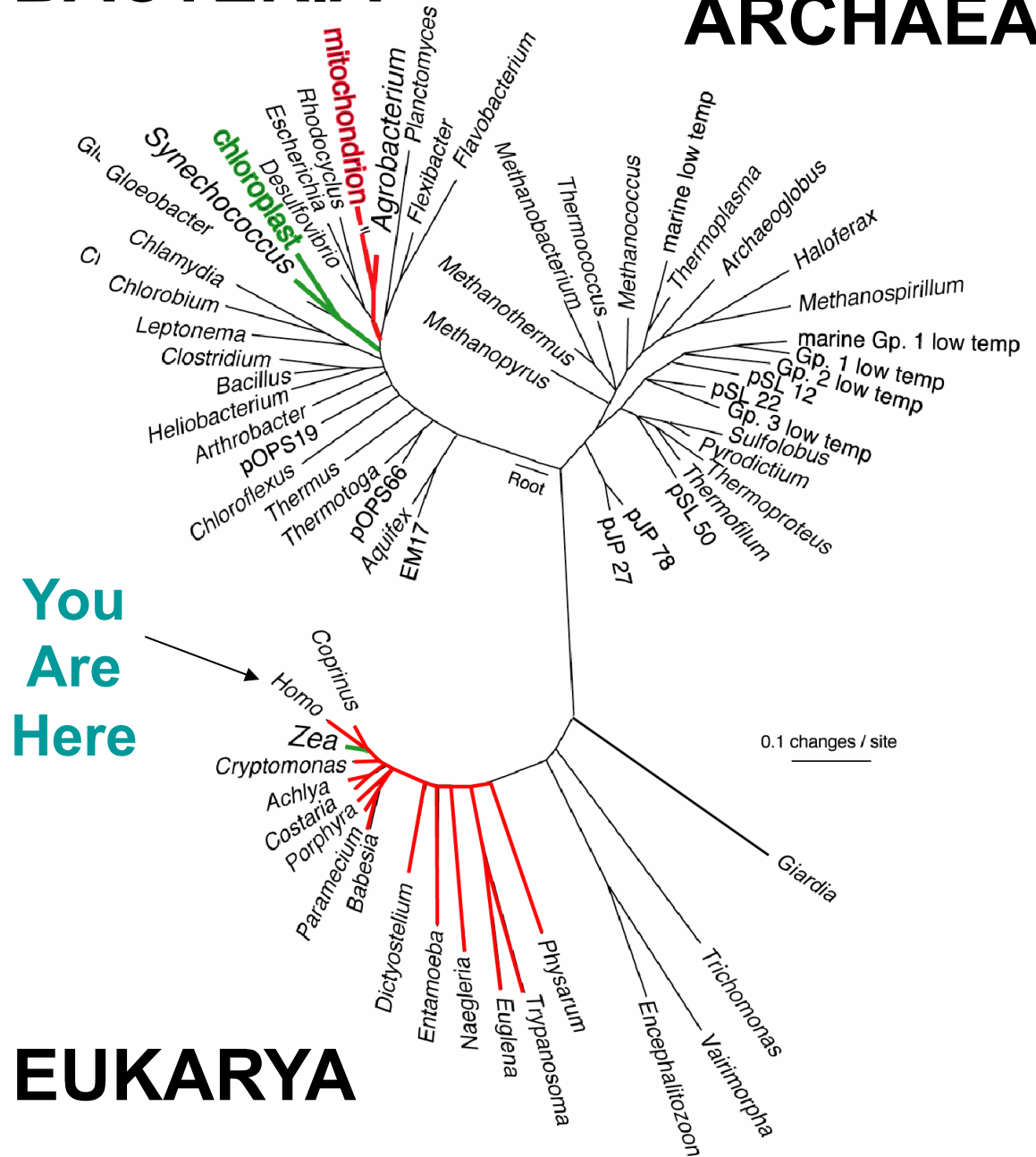
# SSU rRNA



# Universal Tree of Life

## BACTERIA

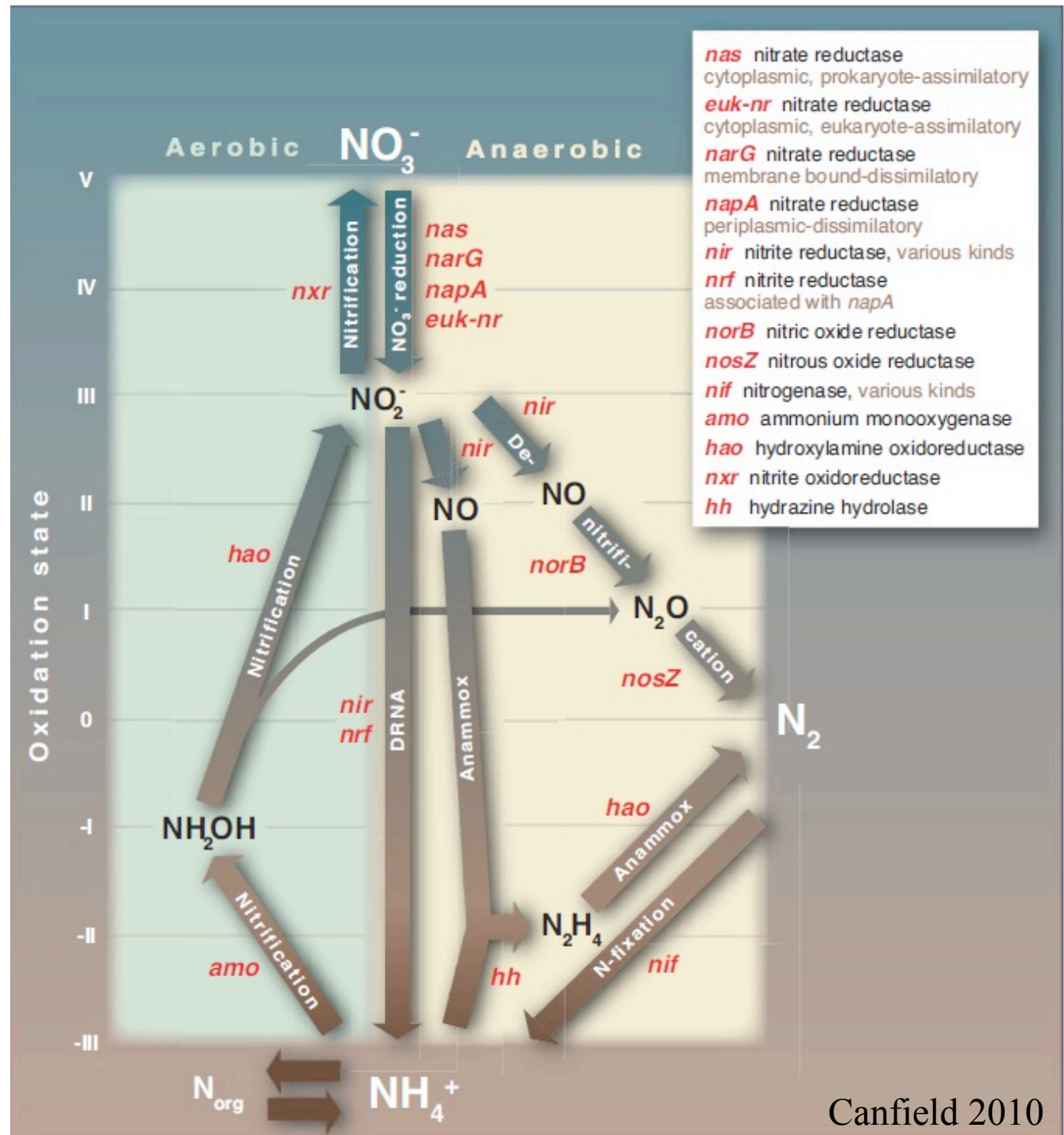
## ARCHAEA



Modified from Norman Pace

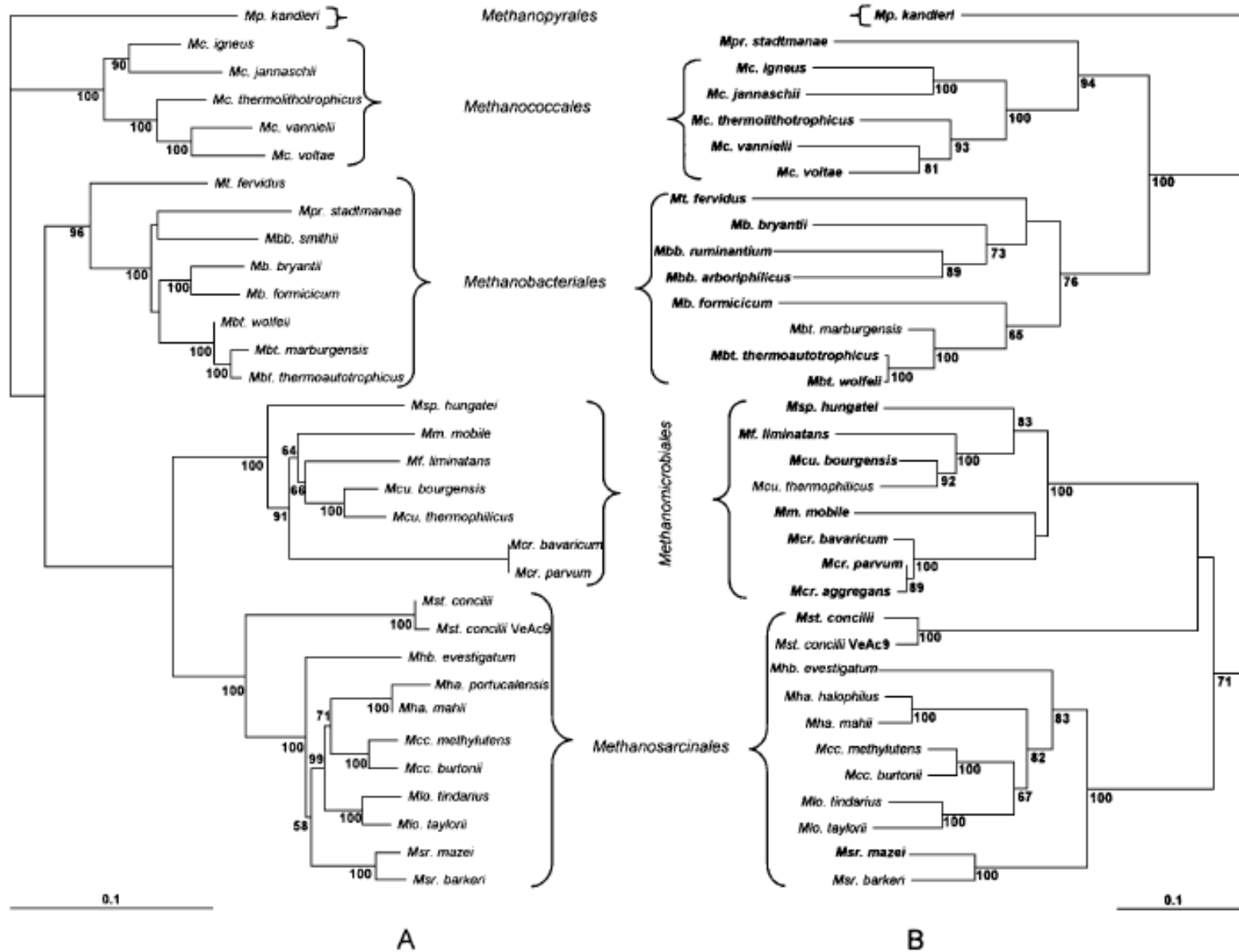
# Beyond 16S rRNA:

## What is the functional potential of microbes?



# 16S rDNA

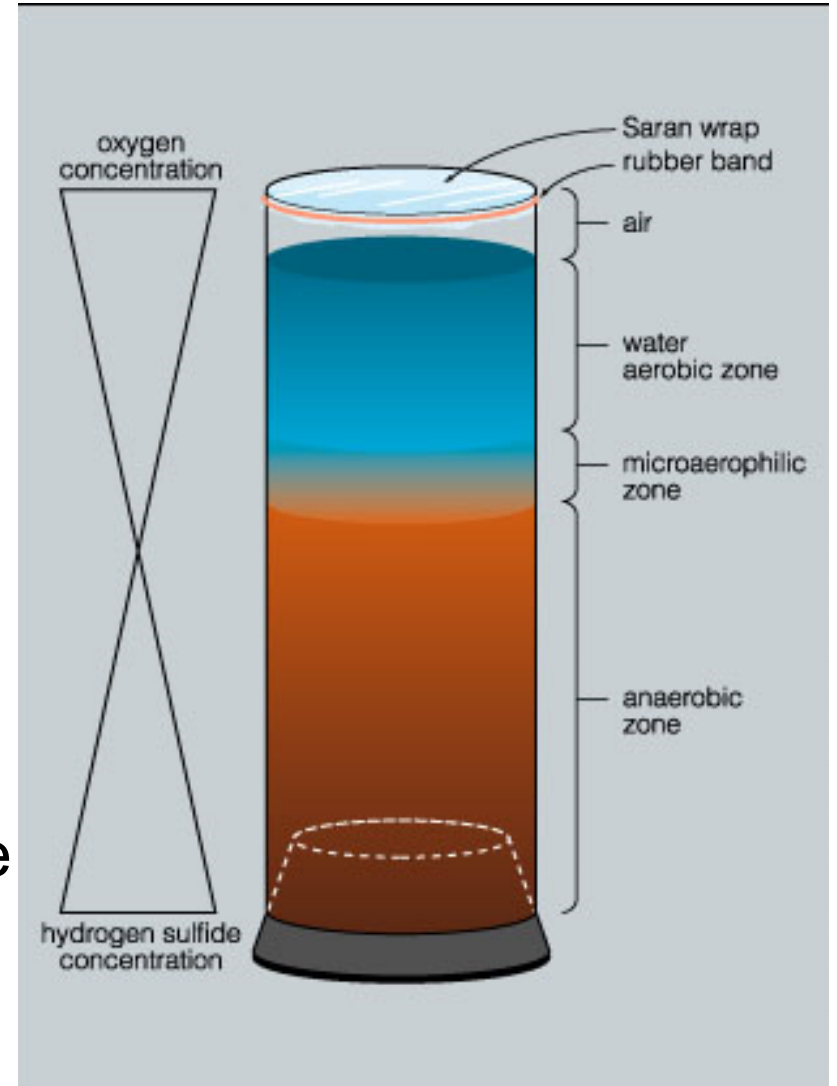
# mcrA





# Our 4 Targets

- 16S rRNA Bacteria
- 16S rRNA Archaea
- *mcrA* Methanogens
  - Methyl coenzyme M reductase
- *dsrB* Sulfate reducers
  - Dissimilatory bisulfite reductase



# What To Do: Make Master Mix

Reagent	1X (25 $\mu$ l rxn)	X 7
Water	5.5 $\mu$ l	
OneTaq 2X Master Mix	12.5 $\mu$ l	
0.4% BSA	4 $\mu$ l	
Total	22 $\mu$ l	

---

## What To Do:

### Set up PCR mix with DNA and specific primers

Tube	Master mix	Target	Template	Vol	F primer	Vol	R primer	Vol
	$\mu\text{l}$			$\mu\text{l}$		$\mu\text{l}$		$\mu\text{l}$
1	22	Sulfate reducers	Sediment DNA	1	dsr1F	1	dsr4R	1
2	22	Methanogens	Sediment DNA	1	ME1	1	ME2	1
3	22	Bacteria	Sediment DNA	1	8F	1	1492R	1
4	22	Archaea	Sediment DNA	1	21F	1	958R	1
5	22	Archaea	+ control	1	21F	1	958R	1
6	22	Water	- control (water)	1	21F	1	958R	1