Molecular Methods in Microbial Ecology

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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

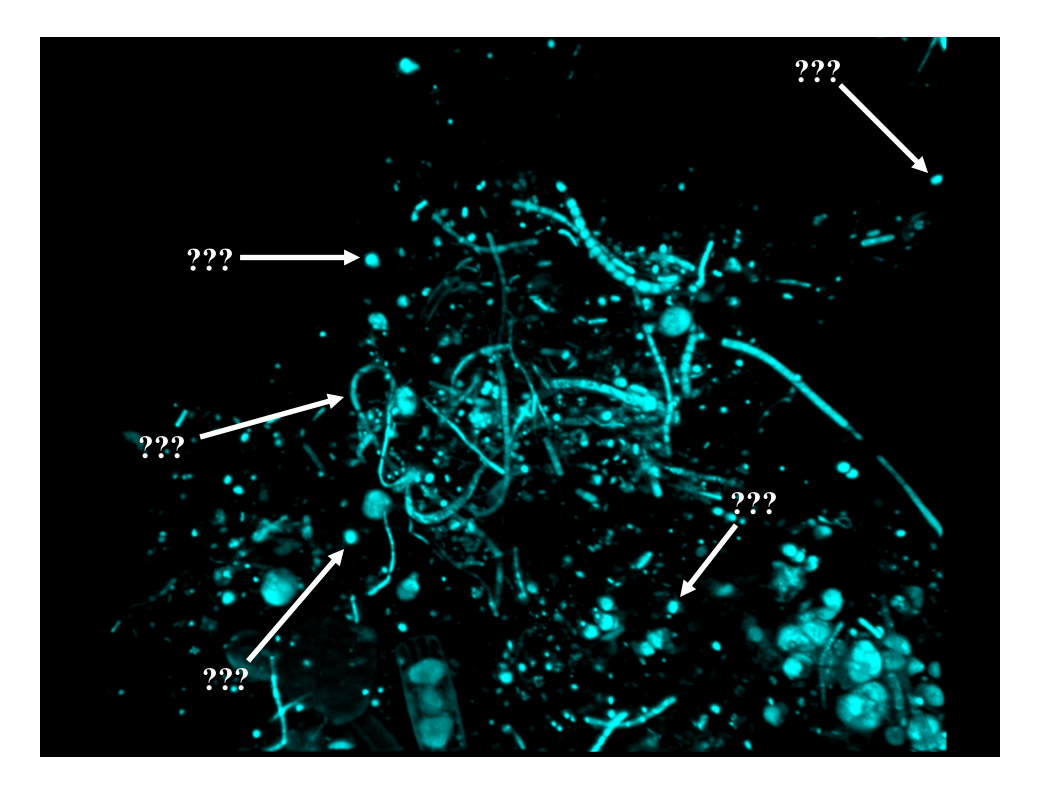
Tuesday 10/30/17 Analyze PCR results, Lecture

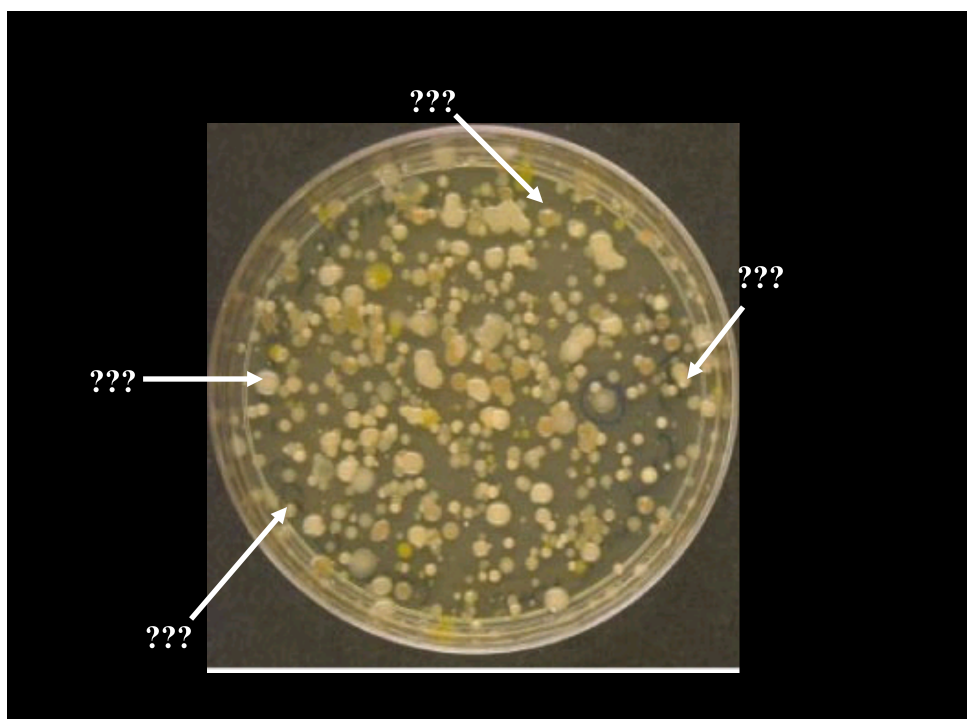
Readings: Head et al. 1998. Microbial Ecology 35: 1-21.

Day 1

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







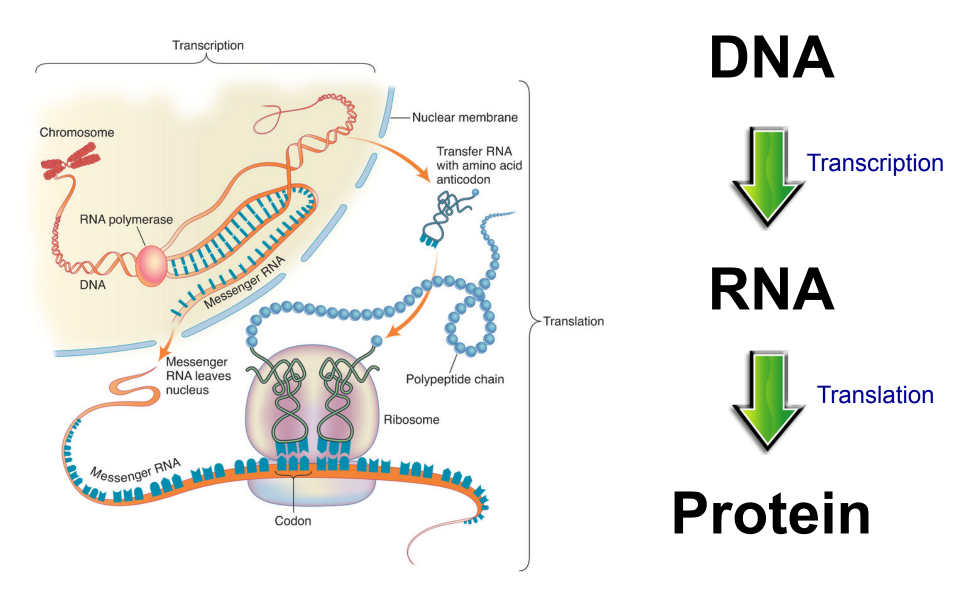
The Challenge for Microbial Ecology

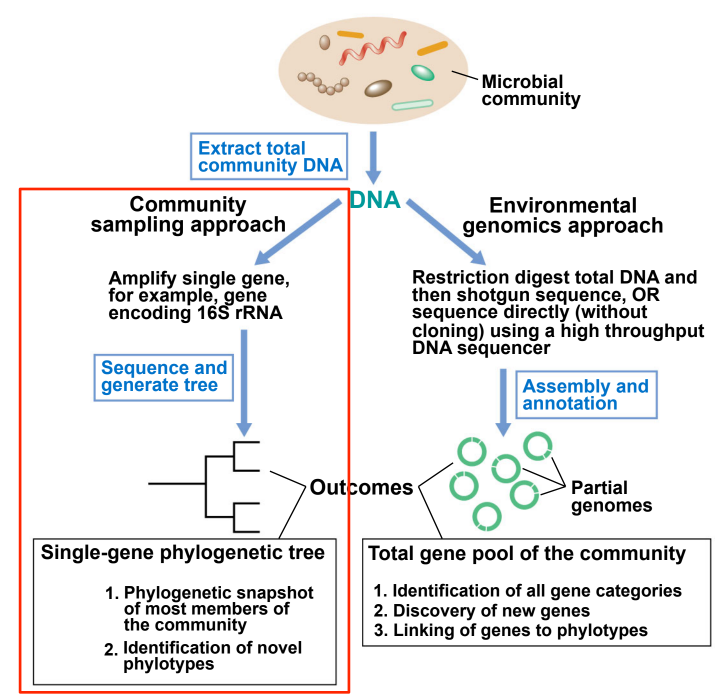
Habitat	Culturability (%)			
Seawater	0.001-0.1			
Freshwater	0.25			
Sediments	0.25			
Soil	0.3			

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

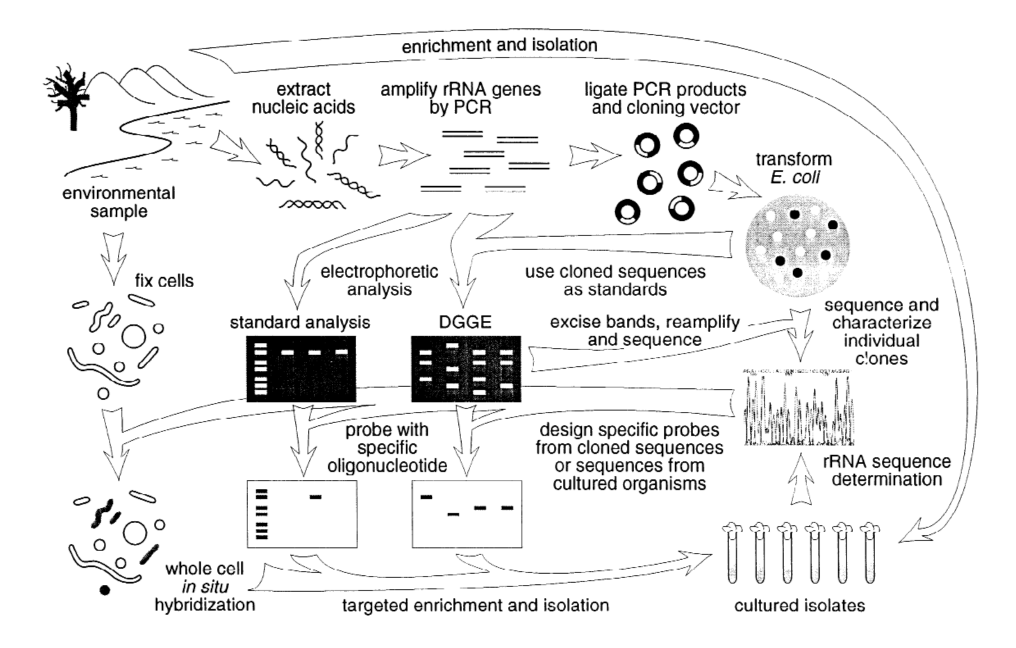
From Amann et al. 1995 Microbiological Reviews

The Central Dogma

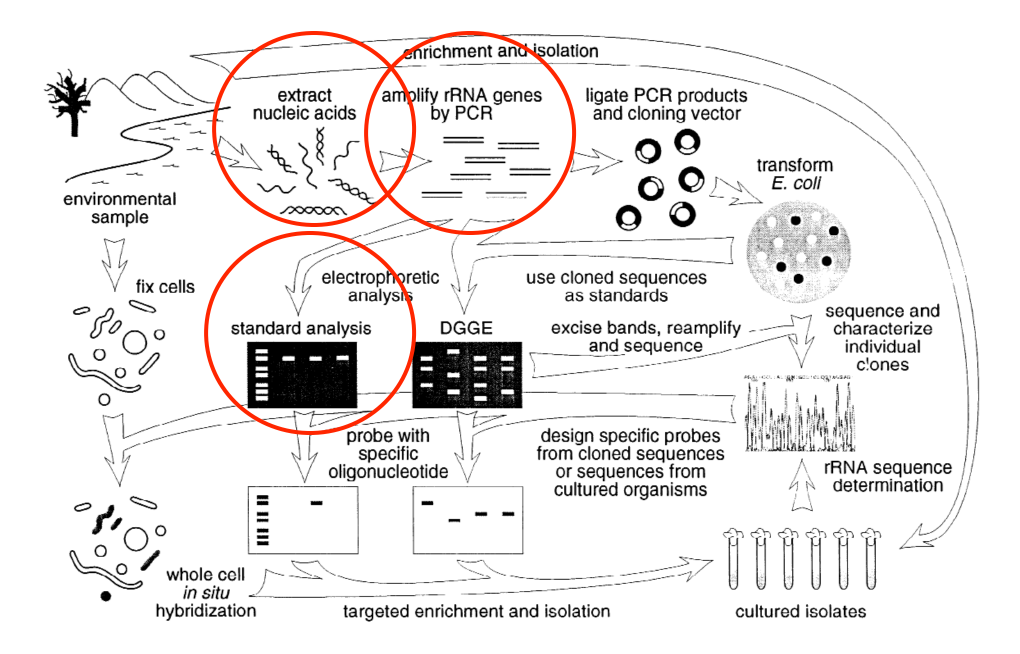




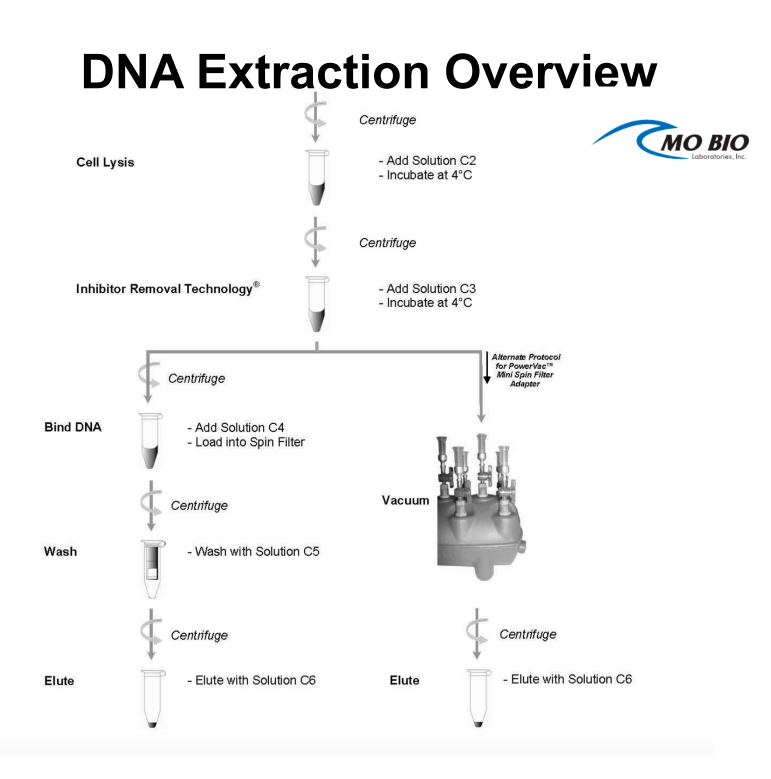
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Head et al. 1998



Head et al. 1998



DNA Extraction

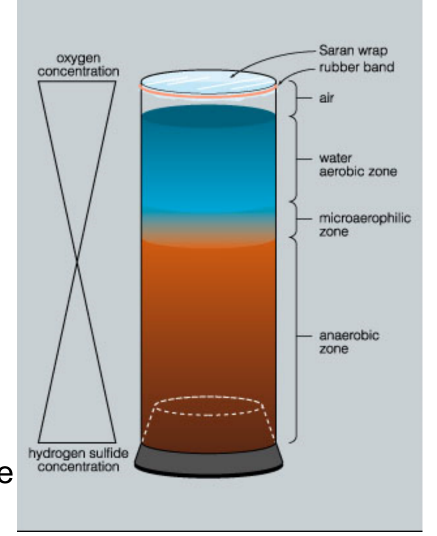
- 1. Lyse cell membrane
 - a. Chemically \rightarrow detergent
 - b. Physically \rightarrow bead beating
- 2. Pellet cell membrane, proteins and other cell parts while DNA stays in solution
- 3. Remove other inhibitors from DNA
- 4. Mix DNA with acid and salt \rightarrow stick to filter
- 5. Wash filter-bound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

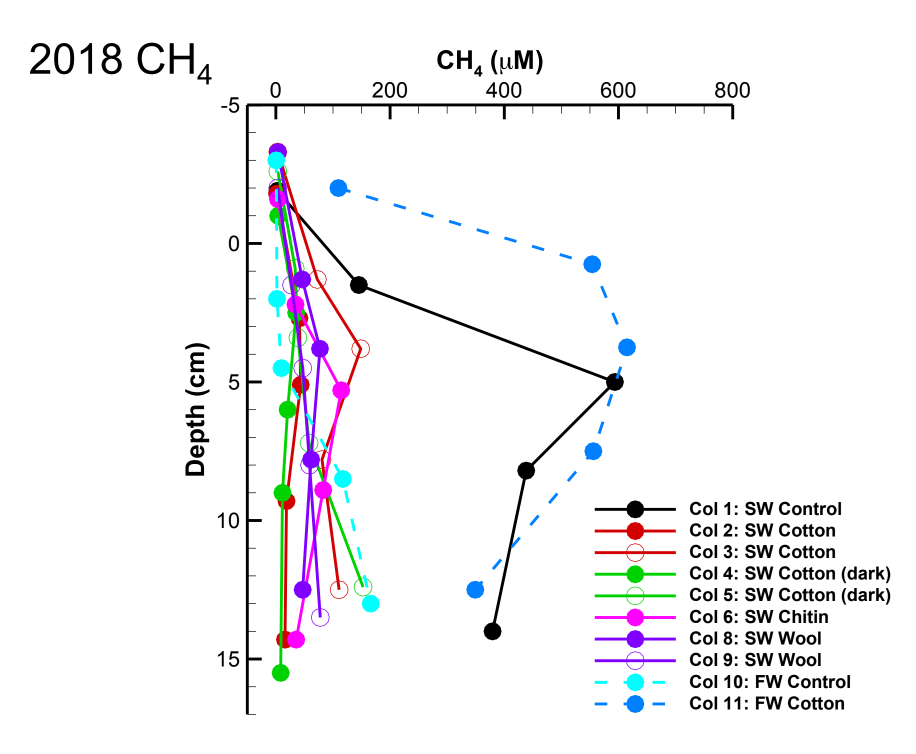
Choosing a Depth Horizon

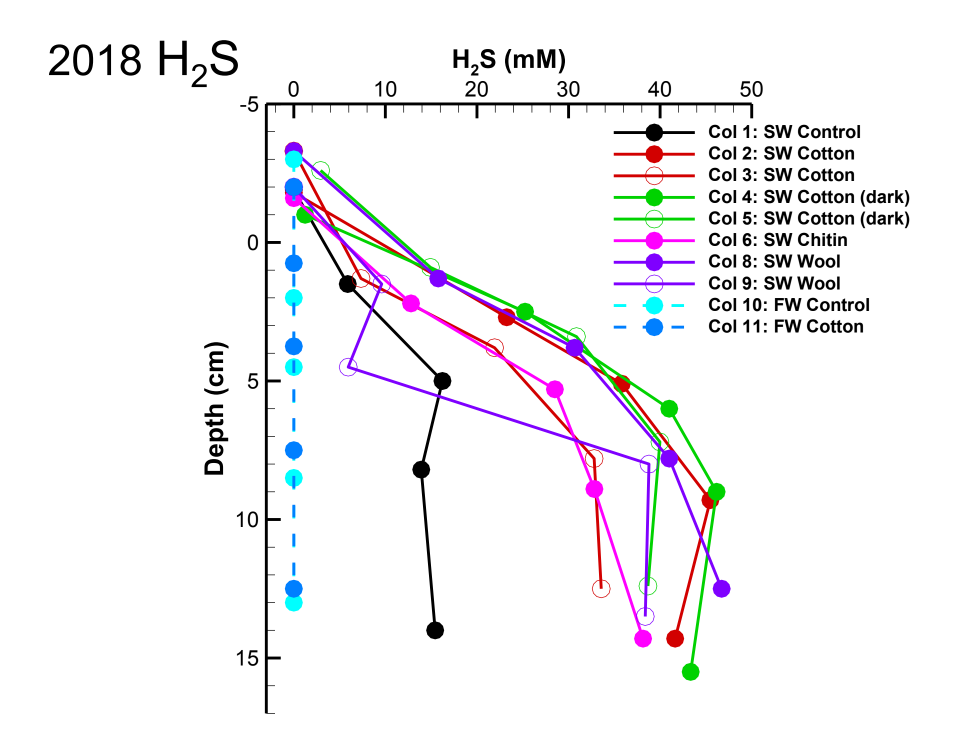
- 16S rRNA Bacteria
- 16S rRNA Archaea
- *mcr*A Methanogens

 Methyl coenzyme M reductase
- *dsr*B Sulfate reducers

 Dissimilatory bisulfite reductase

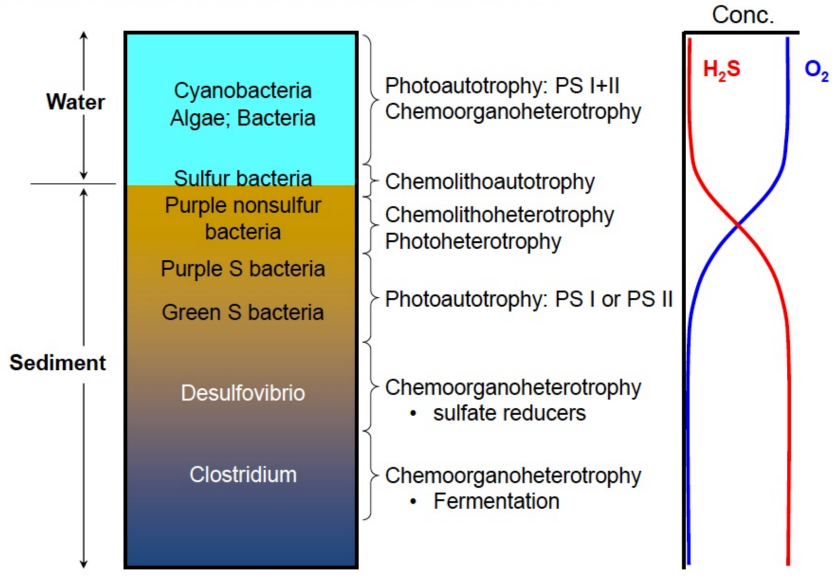






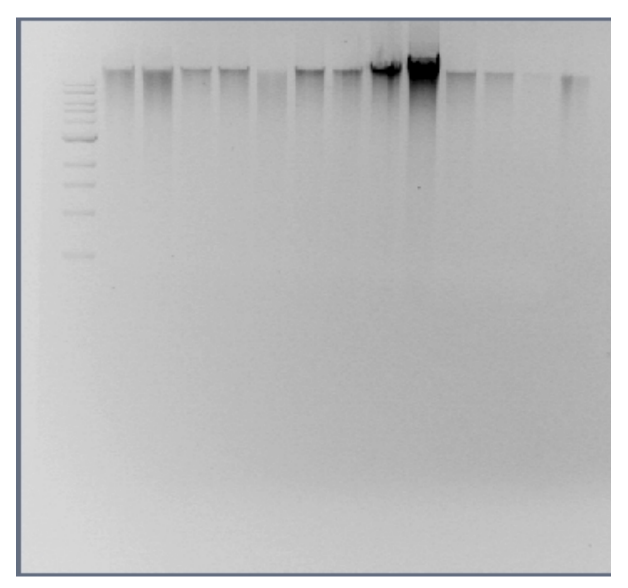
Winogradsky Column

Microenvironments generated by chemical gradients.



Day 1, Part II

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



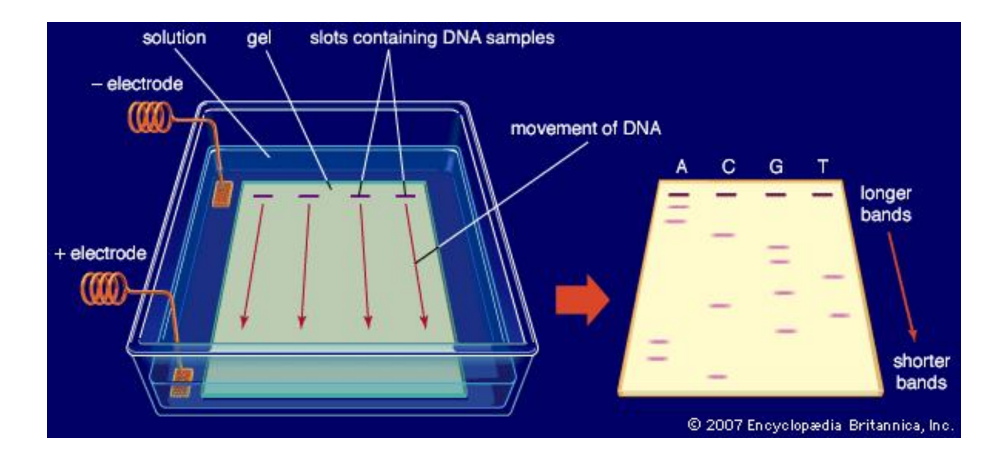
Genomic DNA

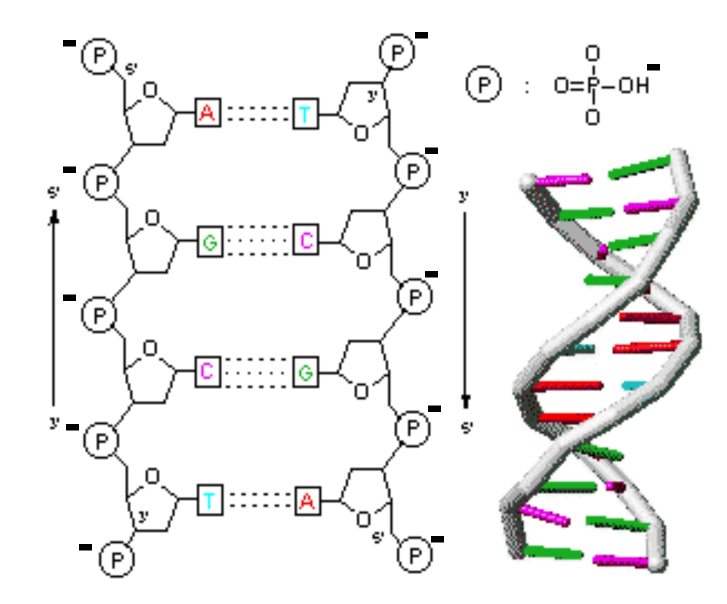
The sum total of all DNA from an organism or a community of organisms

Basics of Gel Electrophoresis

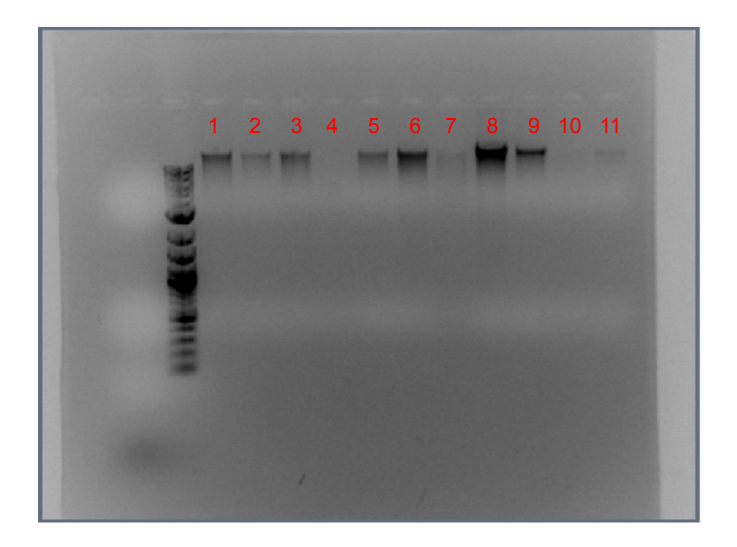
- The gel is a matrix (like jello with holes)
- DNA is negatively charged- will run to positive
- Smaller fragments run faster than larger ones
- Gel contains Ethidium Bromide, which binds to DNA and fluoresces when hit with UV light (WEAR GLOVES!!!)

Gel electrophoresis



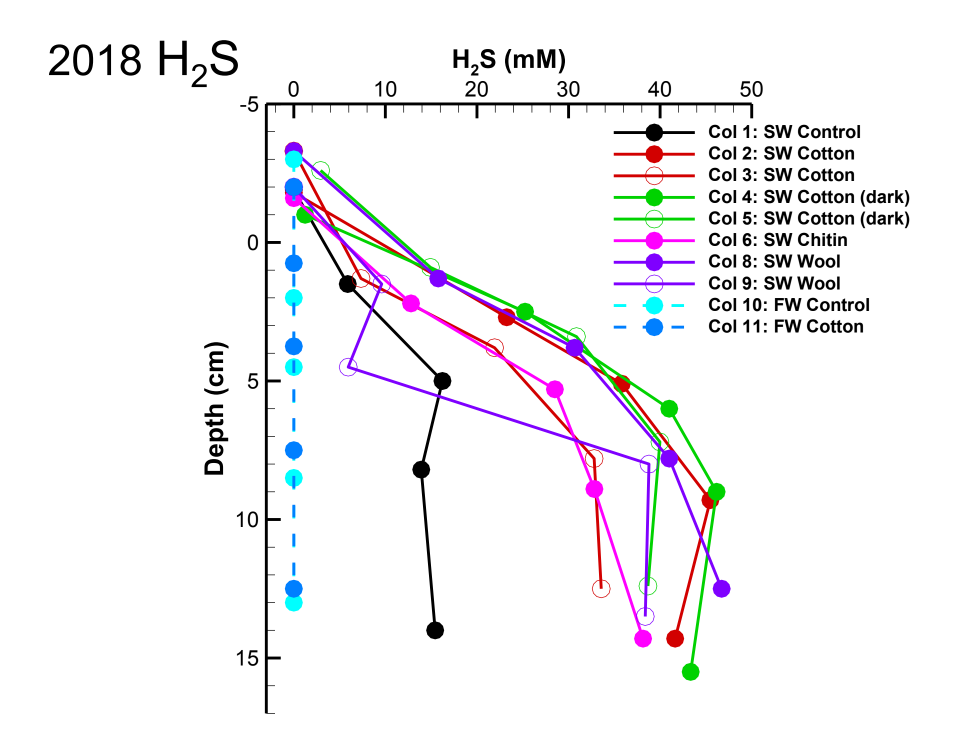


Gel results



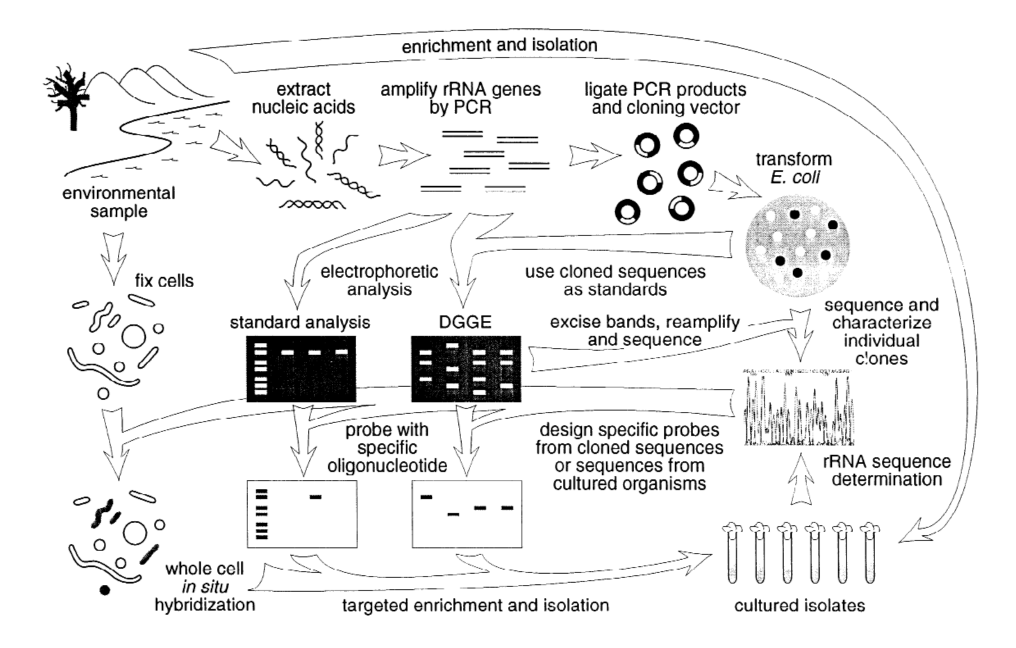
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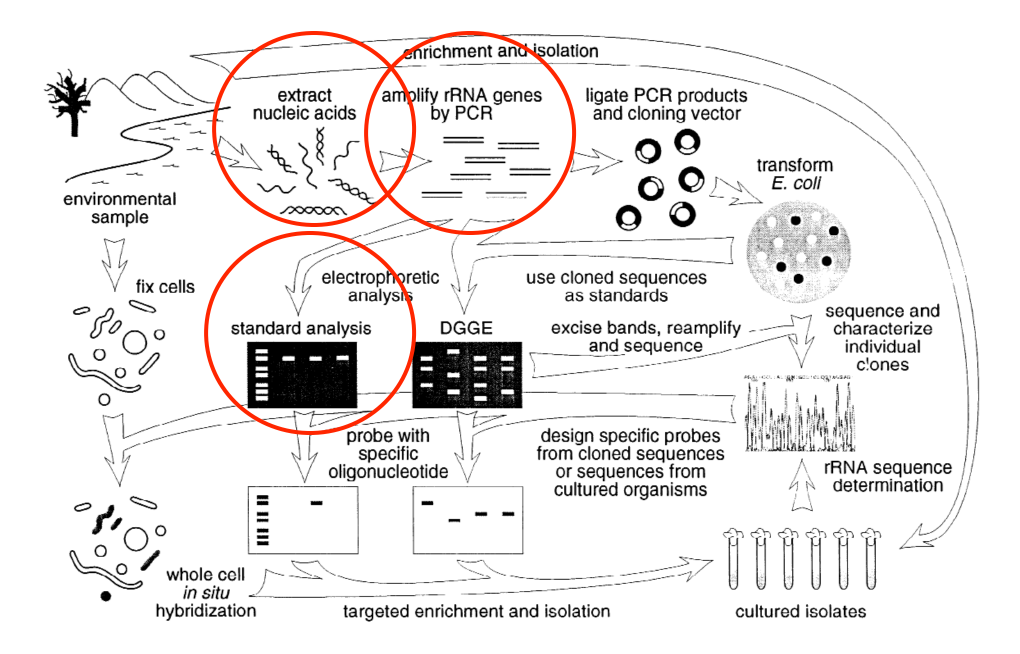


Day 2

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



Head et al. 1998



Head et al. 1998

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

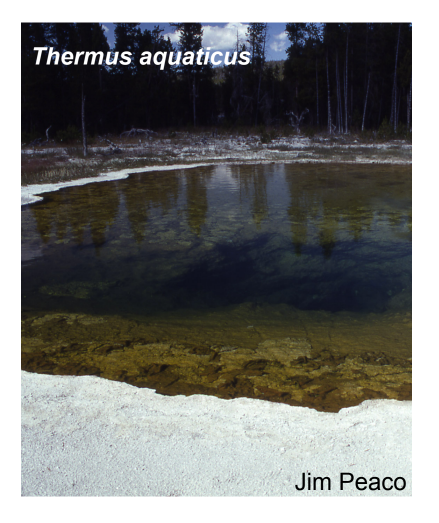


Michael Smith Prize share: 1/2

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".

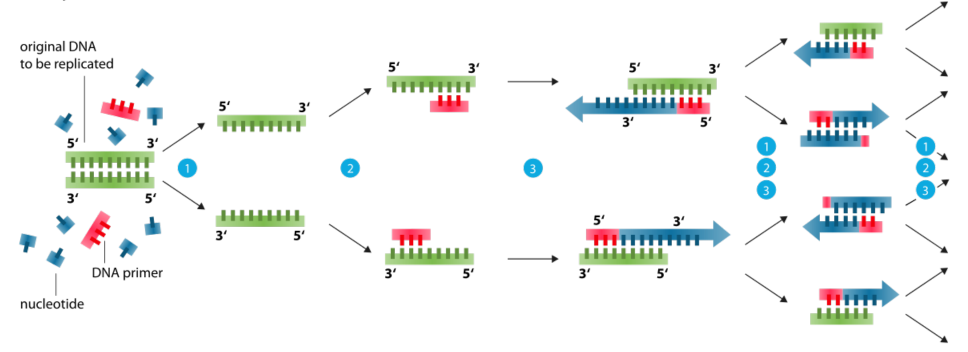
Photos: Copyright © The Nobel Foundation

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





- 2 Annealing at ~68°C
- 3 Elongation at ca. 72 °C

Every PCR contains:

- A DNA Polymerase (most common, Taq)
- Buffer (salt, MgCl₂, etc)
- A set of primers, one Forward, one Reverse
- Various chemicals to minimize inhibition
- Template DNA

Typical PCR Profile

Temperature	Time	Action		
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

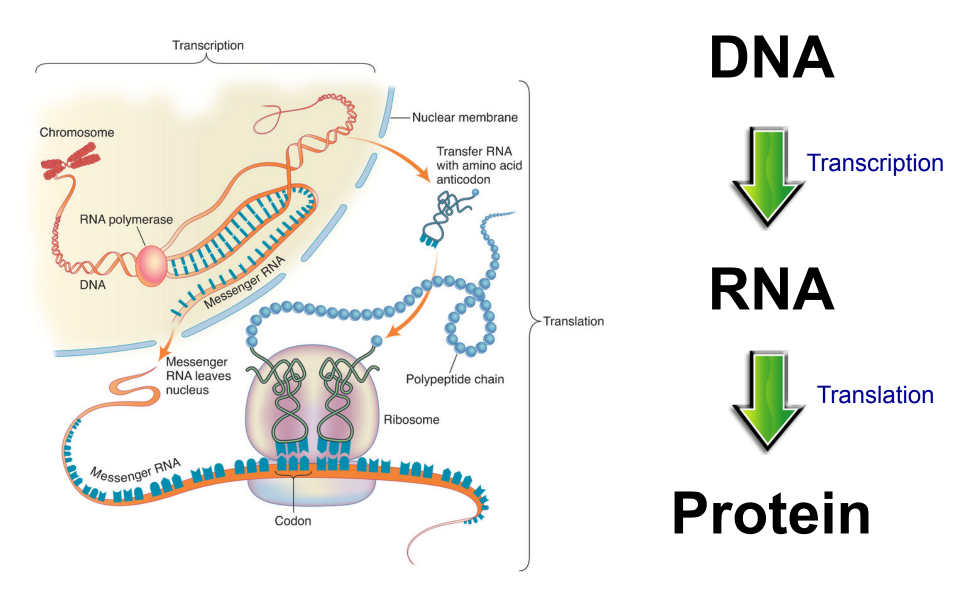
- Temperature and time to activate Taq polymerase
- Temperature and time to allow primer annealing
- Temperature and time for extension
- Concentration of reagents, especially primers, dNTPs, and MgCl₂
- Concentration of template DNA
- Number of replication cycles
- Etc...

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

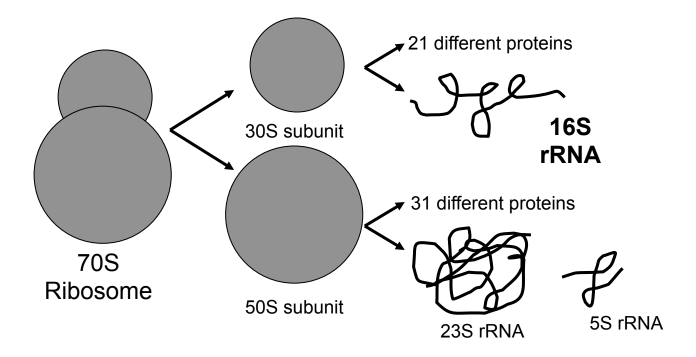


The Central Dogma

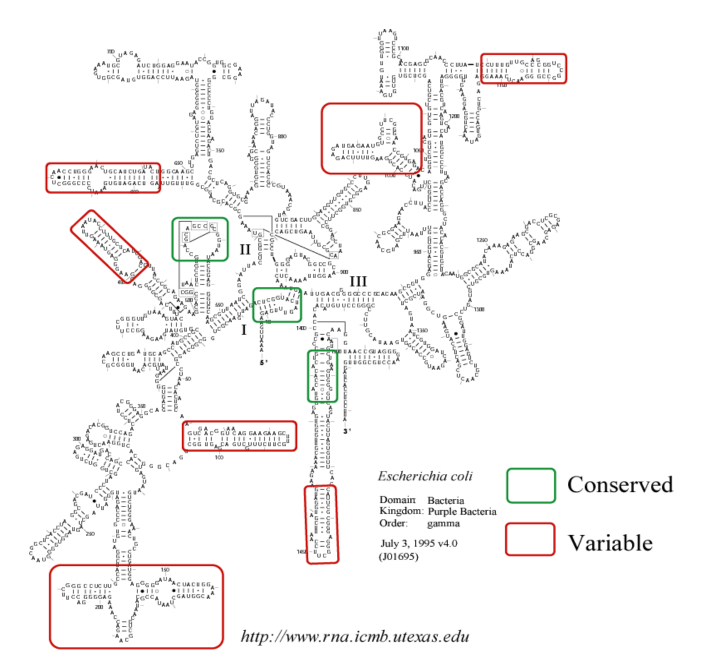


Ribosomes

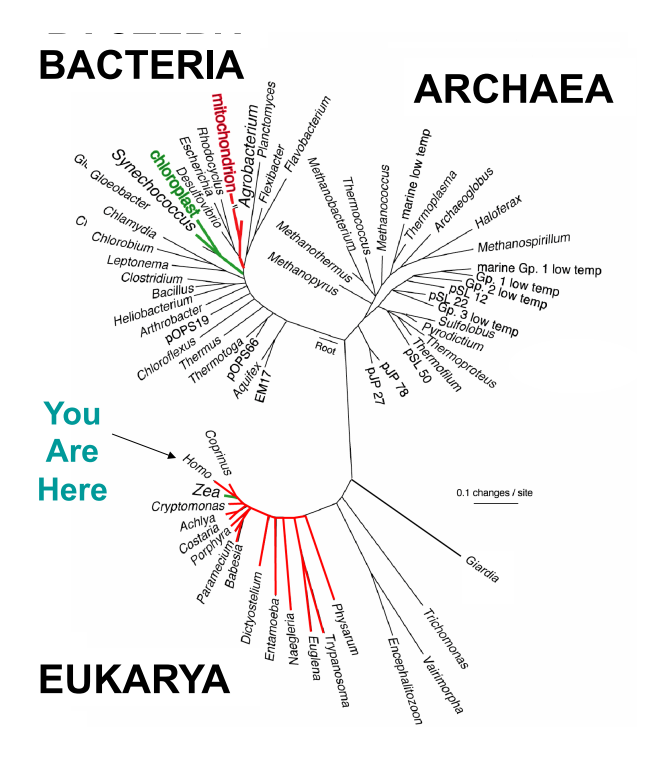
- Make proteins
- rRNA is transcribed from rDNA genes



SSU rRNA

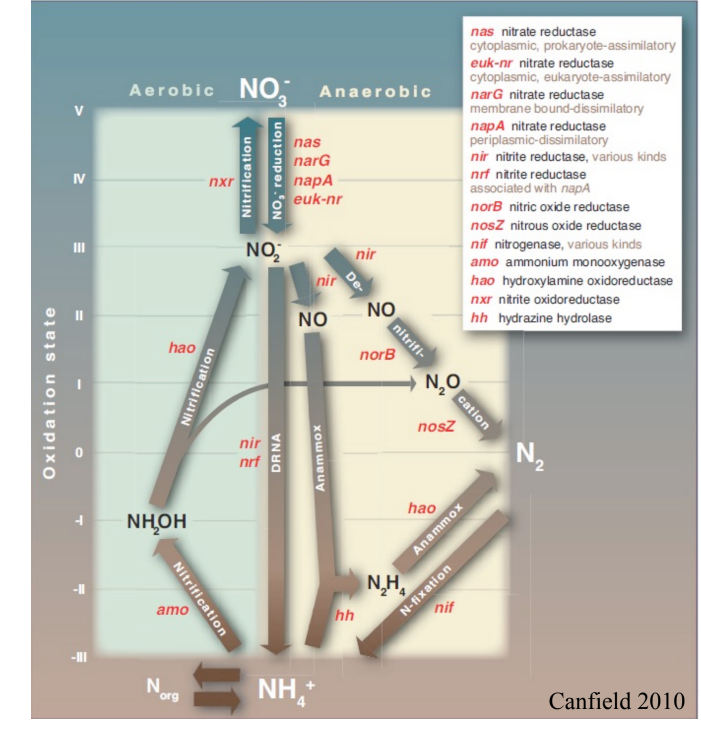


Universal Tree of Life



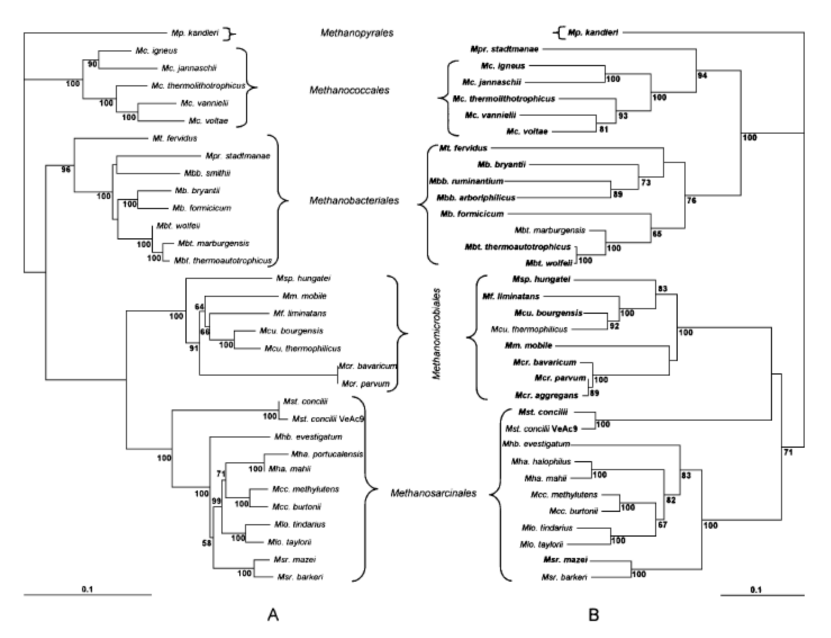
Beyond 16S rRNA:

What is the functional potential of microbes?



16S rDNA

mcrA

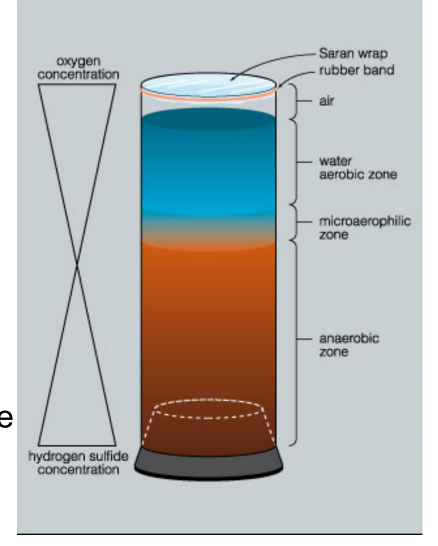


Luton et al. 2002

Our 4 Targets

- 16S rRNA Bacteria
- 16S rRNA Archaea
- *mcr*A Methanogens
 Methyl coenzyme M reductase
- *dsr*B Sulfate reducers

 Dissimilatory bisulfite reductase



What To Do: Make Master Mix

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

What To Do: Set up PCR mix with DNA and specific primers

Tube	Master mix	Target	Template	V ol	F primer	V ol	R primer	V o1
	μ1			μ1		μ1		μ1
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