Lecture 4 Nucleic Acids & Immunology

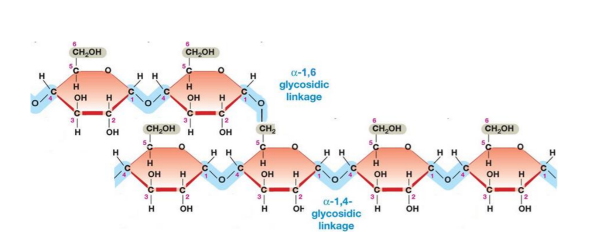
- Nucleic Acid Technologies
- Immunotherapies

Please view the posted video on Enzyme Kinetics before our next class.

Polysaccharides as Energy Storage – Glycogen Storage Disease

Glycogen and is made entirely of glucose units and is used for glucose storage.

Branch point α-1,6- glycosidic



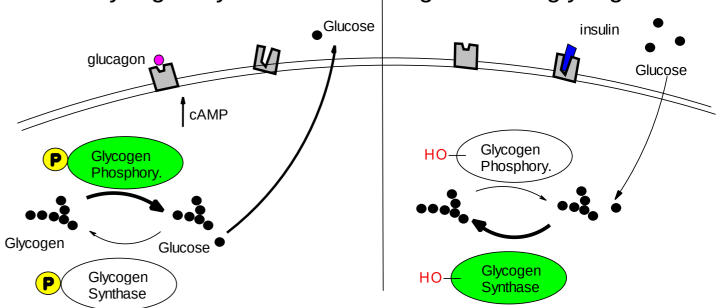
α-1,4- glycosidic linkage

Glycogen Levels are regulated by hormones secreted due to blood glucose levels.

- Glucagon low blood sugar
- Insulin high blood sugar

Two enzymes degrade or synthesize glycogen

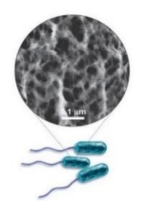
- Glycogen phosphorylase releases glucose from glycogen
- Glycogen synthase stores glucose in glycogen



linkage

Polysaccharides as Structural Molecules

Peptidoglycan (protein + sugar) in bacterial cell wall





Peptidoglycan (Bacterial Cell Wall)

Many antibiotics interfere with cell wall synthesis (e.g. penicillin)

Summary and Expectations for Carbohydrates

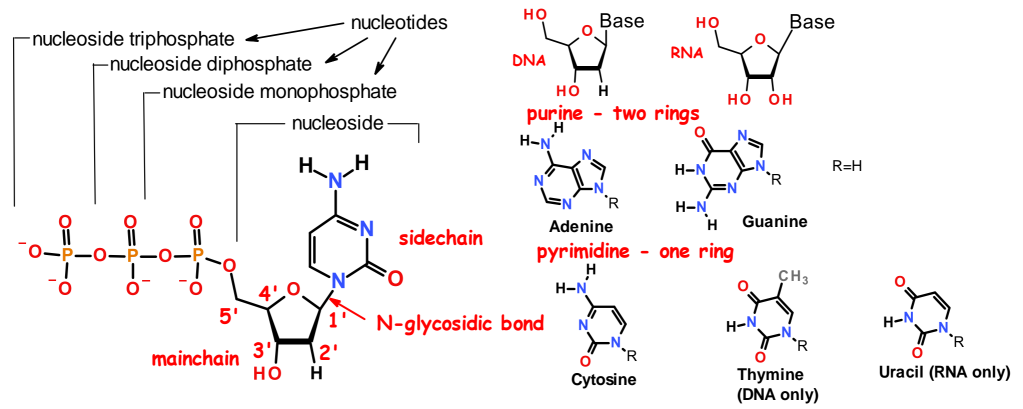
Key Points:

- General structure of monosaccharides be able to distinguish between aldose and ketose (and identify compounds that are not sugars).
- Know how to number carbons on aldoses and ketoses.
- Be able to describe the linkage between two monosaccharides (configuration at the anomeric carbon, atoms linked)
- Treatments for lactose intolerance
- Be able to describe the linkage between glucose molecules in:
 - Glycogen (glucose storage)
- Be able to describe the overall structure of the peptidoglycan in bacterial cell walls.

Nucleic Acid Technologies

- Review of DNA Structure
- Review of DNA Polymerase activity
- Nucleic Acid Technologies PCR & Sequencing

Nucleic Acid Structure



Monomeric Units

- a) Nucleoside triphosphates are the building blocks of nucleic acids (dNTP = dATP, dGTP, dCTP, dTTP)
- b) The base ("sidechain") is connected to the C1' of the sugar ("mainchain") by an **N-linked glycosidic** bond.

Base + sugar + p phosphates - pucleat

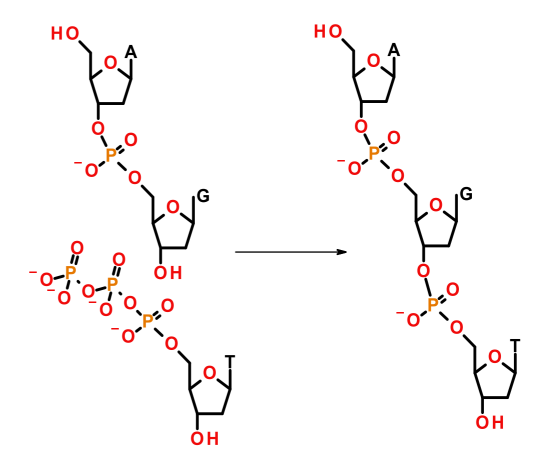
Base + sugar + n-phosphates = **nucleotide**

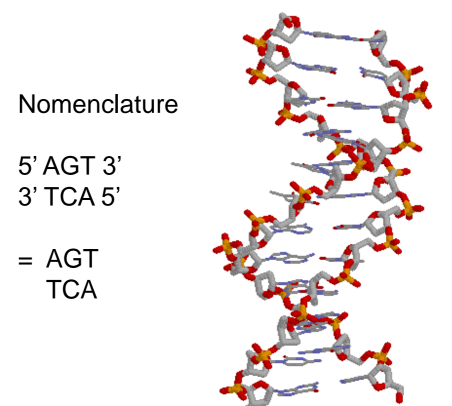
- c) The carbon atoms on the sugar are numbered 1' to 5'. The primes distinguish the atoms on the sugar from those on the base.
- d) DNA differs from RNA in the sugar (deoxyribose versus ribose) and one base.
- e) Four different monomers, A, G, C, T in DNA. U replaces T in RNA.

DNA and RNA are Polynucleotides:

- Two phosphates are lost during polymer formation.
- The phosphodiester backbone is comprised of deoxyribose (DNA) or ribose (RNA) sugars bridged by one phosphate between the 3' and 5' positions of the sugars. Be able to draw this structure.
- The phosphates are always ionized (pK_a~1), nucleic acids are polyanions. The negative charge is important for protein interactions (and electrophoresis).
- Note the polarity: 5' → 3'. Be able to identify the 5' and 3' ends:
 - Start at the end atom and move down the chain.
 The first carbon you find defines the end.

Sequence of nucleotide bases is written in the 5'-3' direction.





https://www.andrew.cmu.edu/user
/rule/jsmol/nucleic.html

Double Helical Structures: B-DNA

- a) The helix is right-handed; the chains are antiparallel.
- b) 10 bp/turn.
- c) The helix interior is filled with stacked base, phosphates and deoxyriboses on the outside.
- d) T pairs with A via two "Watson-Crick H-bonds"
- e) C pairs with G via three "Watson-Crick hydrogen bonds"
- f) Opposite strand termed "complimentary strand". Top strand is always written 5'->3', lower strand 3' -> 5'.

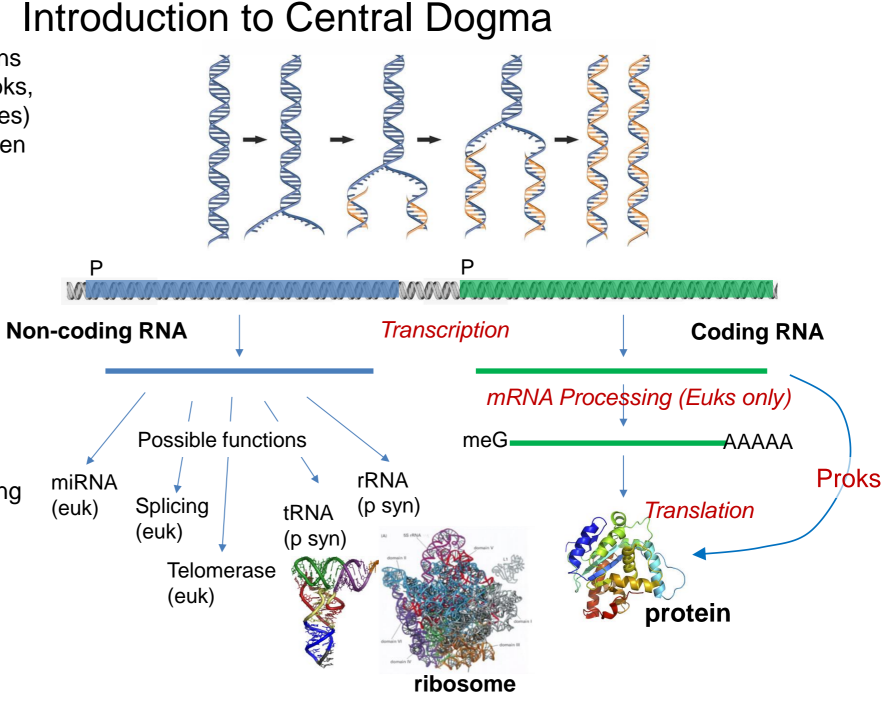
Genome: Entire DNA content of an organism, contains all of the instructions for life. Single circular molecule in Proks, multiple linear molecules (chromosomes) in Euks. The genome is *replicated* when cells divide.

Gene – a segment of DNA that is converted (*transcribed*) to RNA. A *promoter* (P) sequence on the DNA is the minimal requirement for the production of RNA.

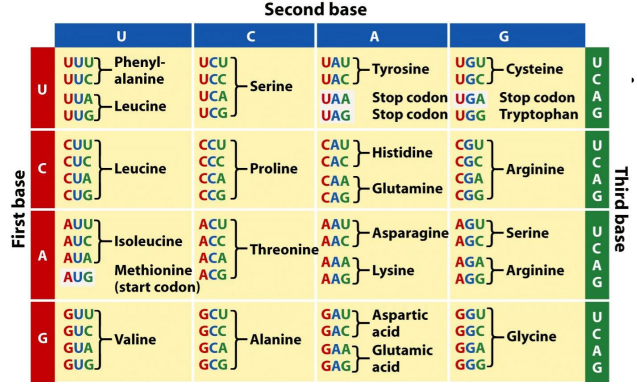
RNA molecules are processed in **Eukaryotic cells** before they are functional. Processing includes:

- Splicing removal of introns, joining exons to generate protein coding region.
- 5' capping
- 3' polyA tail

Many RNAs are functional on their own mRNA are *translated* to a protein.



The Genetic Code – Converting a DNA/RNA Sequence to a Protein



Codon = 3 bases that code for an amino acid

...ATATGCCCATGTGGTAA..
(DNA Sequence)
...AUAUGCCCAUGUGGUAA..

..U-AUG-CCC-AUG-UGG-UAA

(Punctuated RNA sequence – how the ribosome interprets the sequence)

(mRNA Sequence)

- Each codon codes for one amino acid.
- Many amino acids are coded by more than one codon.
- Most organisms use the same codon table some codons have different meanings in some organisms.

Special Codons:

AUG = Is used to begin almost all proteins that are synthesized on the ribosome, codes for methionine when found internally. UAA, UAG, UGA = stop codons, terminate synthesis

(Protein Sequence)

DNA Polymerases – Used in DNA Sequencing and PCR

- DNA polymerases utilize a template to direct the order of added bases,
- The enzyme will continue to the end of the template.
- Require a basepaired primer with a 3'OH. Primer can be DNA or RNA, DNA is used for laboratory work, RNA is used by the cell during replication
- New dNTP added to the 3' hydroxyl of the existing polymer, elongation in the 5' to 3' direction.
- Pyrophosphate (PP) is released and hydrolyzed to two inorganic phosphates.

Expectations: Know the features of this reaction.

DNA Polymerase – Fundamental Activity.

A short 4 base primer (ATCA) is added to a template, and the temperature is lowered to allow annealing (basepairing) of the primer to the template.

- 1. Where (what position) will this primer anneal?
- 2. What is the first base added by the polymerase? A G C T
- 3. What is the last base added by the polymerase? A G C T



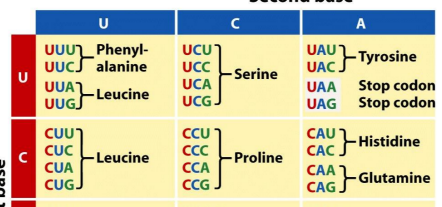
Handbook of Clinical Neurology Volume 147, 2018, Pages 105-123

The state of the s

Repeat Expansion Diseases – Errors in DNA Replication

Second base

Chapter 9 - Repeat expansion diseases				
Henry Paulson ≗ 🖾				
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- CAG at least 10 diseases (Huntington disease, spinal and bulbar muscular atrophy, dentatorubral-pallidoluysian atrophy and seven SCAs)
- CGG fragile X, fragile X tremor ataxia syndrome, other fragile sites (GCC, CCG)
- CTG myotonic dystrophy type 1, Huntington disease-like 2, spinocerebellar ataxia type 8, Fuchs corneal dystrophy
- GAA Friedreich ataxia

https://doi.org/10.1016/B978-0-444-63233-3.00009-

- GCC FRAXE mental retardation
- GCG oculopharyngeal muscular dystrophy
- CCTG myotonic dystrophy type 1
- ATTCT spinocerebellar ataxia type 10
- TGGAA spinocerebellar ataxia type 31
- GGCCTG spinocerebellar ataxia type 36
- GGGGCC C9ORF72 frontotemporal dementia/amyotrophic lateral sclerosis
- CCCCGCCCCGCG EPM1 (myoclonic epilepsy)

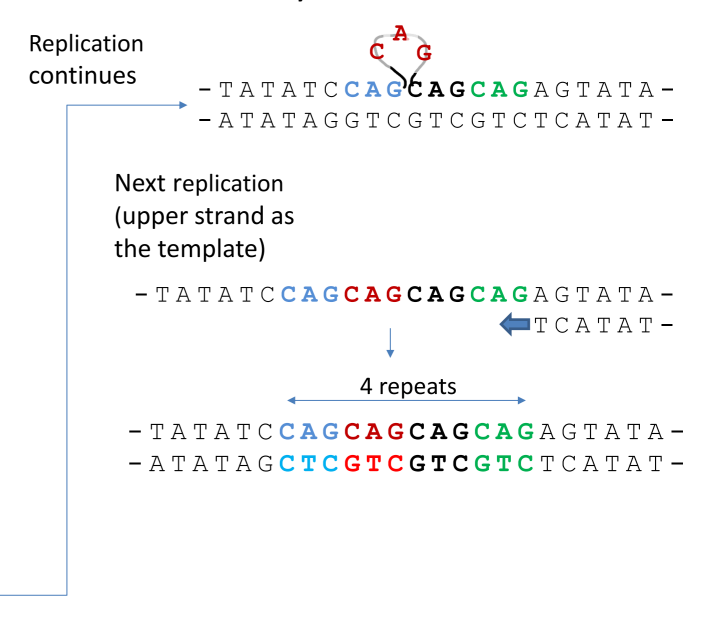
 Repeats in coding regions of genes will generate long stretches of the same amino acid.

CAGCAGCAG = GluGluGlu

- Repeats outside of coding regions can affect gene expression by changing binding of transcription factors.
- These repeats can grow due to slippage of primer during replication
- More repeats = more chance of developing disease.
- The number of repeats can be detected by:
 - DNA sequencing
 - PCR

Repeat Expansions – How Do They Grow?

Original Sequence - 3 repeats (CAG) - T A T A T C C A G C A G C A G A G T A T A --ATATAGGTCGTCGTCTCATAT-During Replication in the cell - T A --ATATAGGTCGTCGTCTCATAT-- TATATCCAGCAGCAG - ATATAGGTCGTCGTCTCATAT -3' end comes loose (primer slippage) -TATATCCAGCAGCAG - ATATAGGTCGTCGTCTCATAT -Looped out DNA - TATATCCA - ATATAGGTCGTCGTCTCATAT -



DNA Sequencing – Sanger (dideoxy) Sequencing

DNA Sequencing - Determining the Order of Bases in the DNA.

Sanger Sequencing:

- Second method to generate long (~1000 base) sequence information (an earlier chemical method developed by Gilbert proved to be impractical for most laboratories (hydrazine = rocket fuel was required)
- Sanger was awarded his 2nd Nobel prize for this work in 1980, shared with Gilbert.







Determine the position of all four bases in a DNA strand = Sequence (video)

Sanger Sequencing:

Primer

$$5'C-A-T-A-T-G^{OH}$$

Known Seq (plasmid) Unknown sequence (insert)

Sequenced region (~1000 bases)

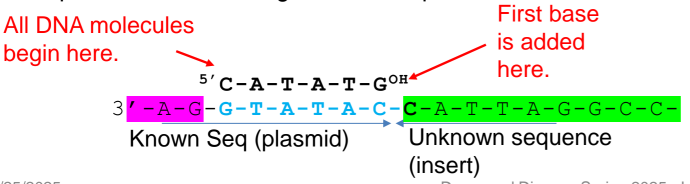
5' C-A-T-A-T-G-G-T-A-A-T-C-C-G-G-T-A-C-G-T-G-C-A---
Template

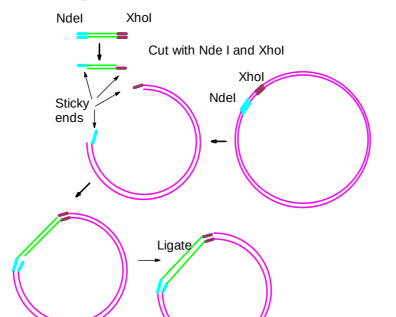
3'G-A-A-G-T-C-G-A-A-G-G-T-A-T-A-C-C-A-T-T-A-G-G-C-C-A-T-G-C-A-C-G-T-----

DNA Sequencing - Determining the Order of Bases Added by DNA Polymerase

- The DNA to be sequenced is inserted into a circular piece of double stranded DNA called a plasmid. The DNA sequence of the plasmid is known.
- The insertion is often accomplished using restriction enzymes that generate single stranded overhangs that allow DNA molecules to be efficiently joined.
- Restriction sites can be added to any DNA fragment using a number of techniques:
 - Addition of a short linker (same site on both ends)
 - PCR (different sites on each end)

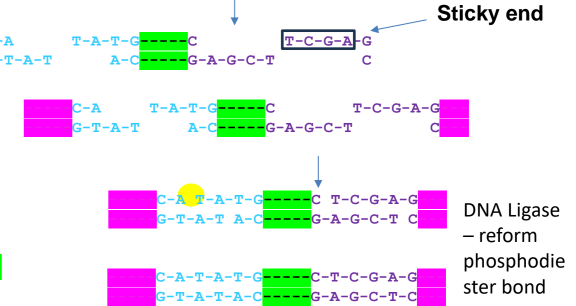
Key Point 1. Start sequencing at known location with primer that anneals at a *unique* location on the plasmid, "upstream" from the region to be sequenced.





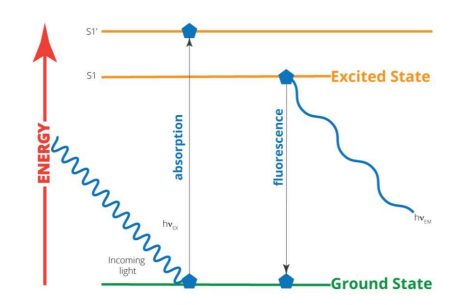
Restriction Enzymes

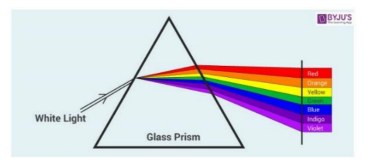
- Recognize a specific sequence in the DNA
- Sequence has 2-fold symmetry same on the top and bottom strand
- Cuts both strands, most generate single-stranded DNA (sticky ends).
- Complementary sticky ends can bind to each other.



DNA Sequencing Methods Use Fluorescent Bases - What is Fluorescence?

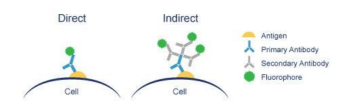
- When molecules absorb light an electron goes from a lower shell to a higher shell. This is where the energy from the light goes.
- In most molecules the electron goes back down to its original shell with the release of heat.
- Fluorescent molecules emit the energy as light of a longer wavelength (different color).
- The color that is emitted depends on the molecule.

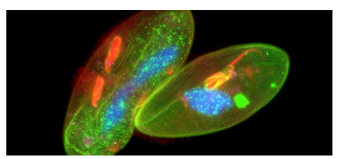






Fluorescently tagged antibodies can be used to stain components of cell with fluorophores.

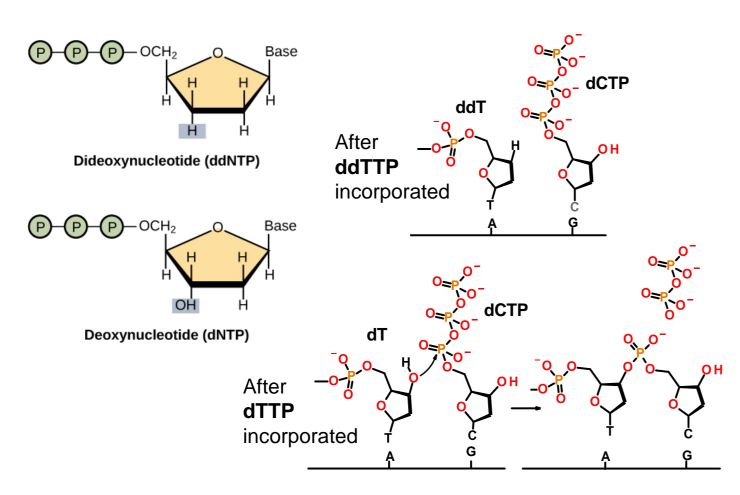




DNA Sequencing - Determining the Order of Bases Added by DNA Polymerase

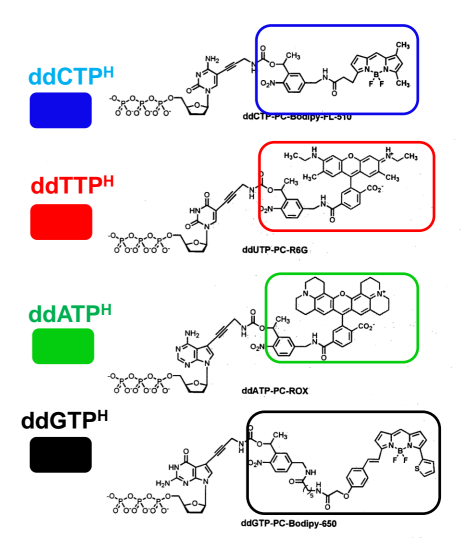
Key point 2. Use a mixture of normal bases (dNTPs) and dideoxy bases (ddNTP) for polymerization. Ratio of dNTP to ddNTP is (100:1), *most of the time elongation occurs.*

•ddNTPs can be added to the DNA since they have a 5'-triphosphate but *terminate* the chain due to the lack of a 3'-OH. ~ 1 in 100 chains terminate at each base addition



Key point 3. The ddNTPs are color coded by different fluorescent emission wavelengths.

The ddNTP that terminated the chain is known from its fluorescent color.



TemplatePrimerDNA PoldTNP, ddNTP

DNA Sequencing – Generation of Fluorescent Fragments

Length=7, Black fluor.

(10)
$$5' - C - A - T - A - T - G - G^H$$

 $3' - A - G - G - T - A - T - A - C - C - A - T - T - A - G - G - C$
(990) $5' - C - A - T - A - T - G - G^{OH}$

3'-A-G-G-T-A-T-A-C-C-A-T-T-A-G-G-C

Length=8, Red fluor.

$$_{\mathbf{z}}$$
(10) $_{\mathbf{5'}}$ $_{\mathbf{C-A-T-A-T-G}}$ $_{\mathbf{G-T^H}}$ $_{\mathbf{3'}}$ $_{\mathbf{A-G-G-T-A-T-A-C-C-A-T-T-A-G-G-C}}$

Length=9, Green fluor

$$(10)_{5'}$$
 - C-A-T-A-T-G-G-T-A^H
3'-A-G-G-T-A-T-A-C-C-A-T-T-A-G-G-C

(990 molecules)

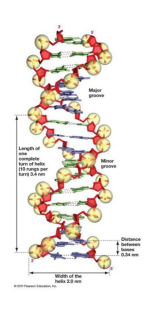
(980 molecules)

All Possible Fragments are Made:

- 1. Each begins with the primer
- 2. Each ends with a known ddNTP, based on the color of the fluorescence.
- 3. Each is one longer than the previous.

Primer Added by Pol.

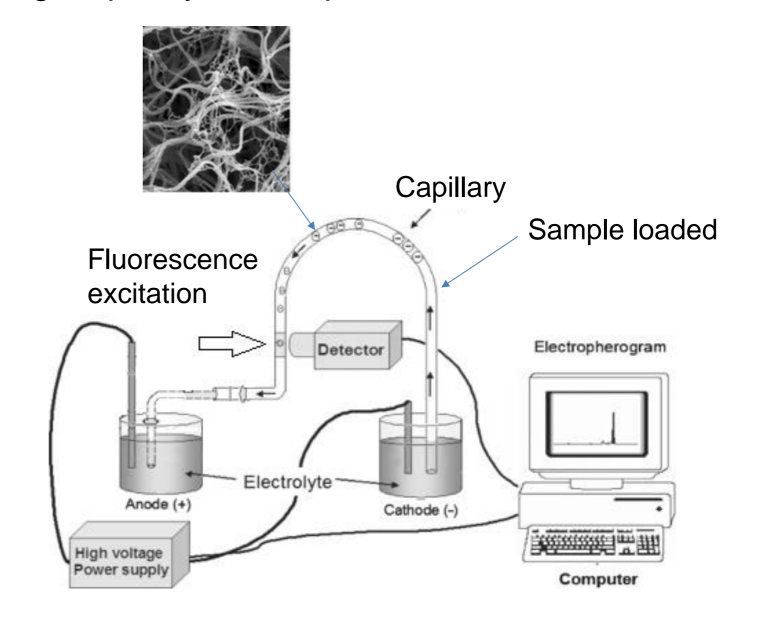
Size Determination of Fragments from DNA Sequencing Capillary Electrophoresis



DNA has a negative charge. It will migrate towards the anode.

Capillary is filled with a gel that causes separation by size.

DNA molecules that are smaller migrate _____.

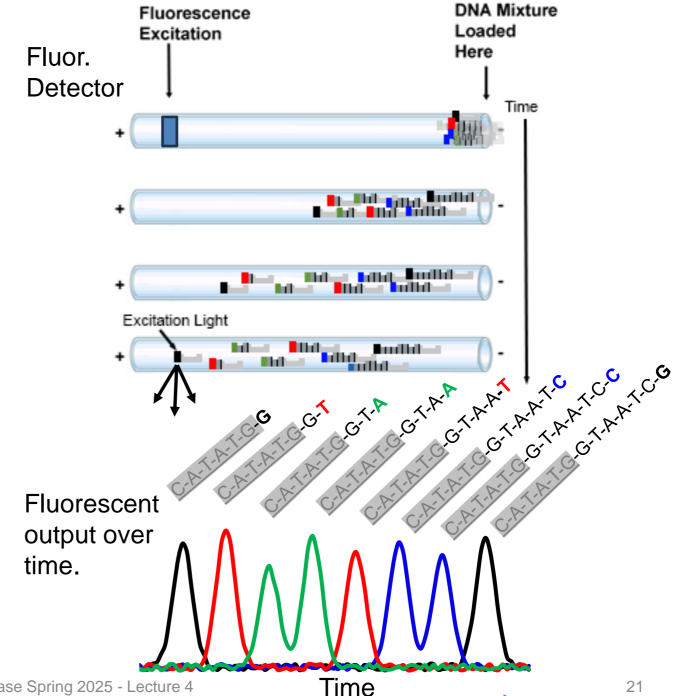


DNA Sequencing – Analysis of Fragments to Determine Order of Addition

4. Capillary Gel Electrophoresis

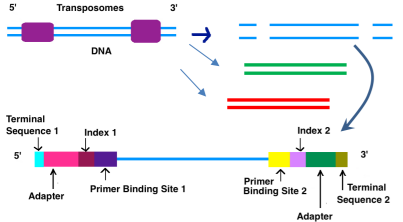
- Migration due to the voltage because of the neg. charge on DNA phosphates
- Separation of DNA molecules by size, smaller travel through gel faster.
- Fragments reach the detector in the order of their length: primer+1 first, primer+2 second, etc.
- At the detector, a laser excites the fluorescence.
- Only fluorescent DNA molecules (terminated) with ddNTP) give a signal.
- The color of the emitted fluorescence gives the dideoxy base at the 3' end of the DNA fragment.
- The order of peaks gives the sequence that is complementary to the template (= strand with primer).

5'-C-A-T-A-T-G G-T-A-A-T-C-C-G 3'-A-G-G-C-T-A-T-A-C-C-A-T-T-A-G-G-C



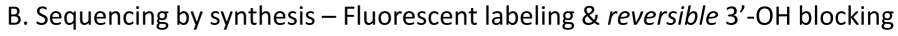
Newer Sequencing Methods-Illumina Dye Sequencing – Next Generation High Throughput

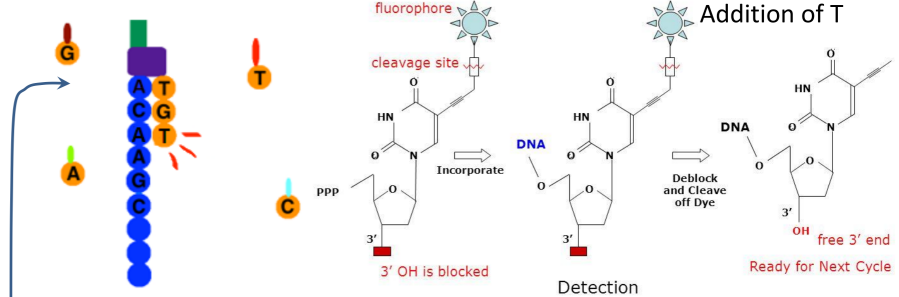
A. Obtaining the DNA



- The entire genome can be sequenced.
- The DNA is fragmented into small 100 base pieces.
- Synthetic DNA is added to the ends (sites for primers for sequencing)
- Different fragments are bound to different location on a solid surface (chip).
- All fragments are sequenced at the same time on the chip.

Cluster formation





- 1. Only one base is added at a time (3'-OH is blocked)
- 2. The base that is added is determined by the color of the fluorescent base.
- 3. 3'-OH blocking group and the fluorescent group are removed prior to the next addition. ~100 cycles can be performed.

Next Generation - Data

HN

fluorophore A

cleavage site

- 1. Primer anneals
- 2. Add dNTPs (3'blocked) + Polymerase
- 3. Wash to remove unincorporated dNTPs
- Detect base added to each cluster using fluorescent emission

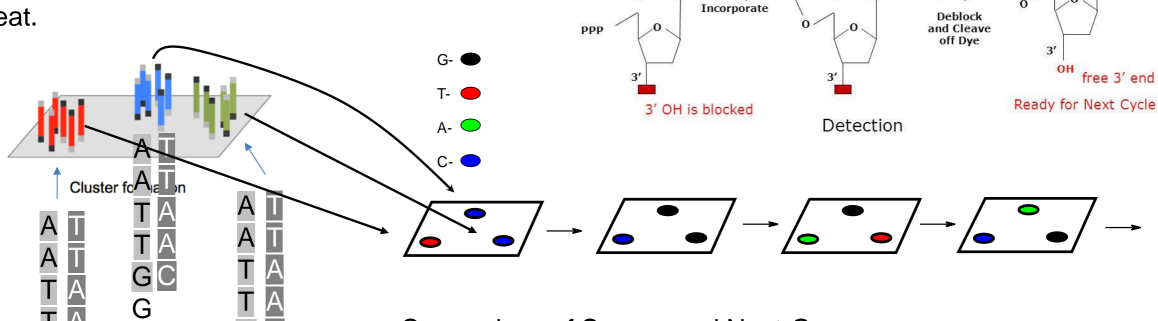
G

G

Α

5. Unblock 3'-OH

Repeat.



Comparison of Sanger and Next-Gen

Method	Read Length	Samples Processed
Sanger	~1000	1
Illumina	~200	~10,000s

DNA

DNA

Α

G

Genotyping at the Molecular Level with DNA Sequencing.

- Sickle cell anemia is caused by a single mutation in the beta chain of hemoglobin
- This mutation causes the hemoglobin to form long polymers that distort the shape of the red blood cell.
- Determining whether someone has the mutation can be useful for treatment.

The 5' end of the Hb gene is shown on the right (ATG=start). Using GGTGCCAG as a sequencing primer gives the following sequences for the normal and mutant genes:

Healthy red blood cell

Sickle red blood cell

Unrestricted blood flow

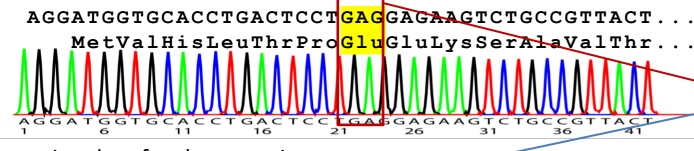
Blood flow blocked by sickle cells

First dd-base added by polymerase

GGTGCCAGAGGATGGTGCACCTGACTCCTGAGGAGAAGTC..

CCACGGTCTCCTACCACGTGGACTGAGGACTCCTCTTCAG..

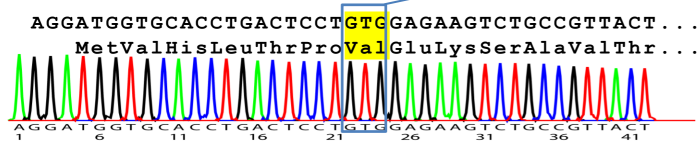
Sequencing data for the normal beta chain is:







Sequencing data for the mutation:



False color code:

A=Green G=Black

T=Red

C=Blue



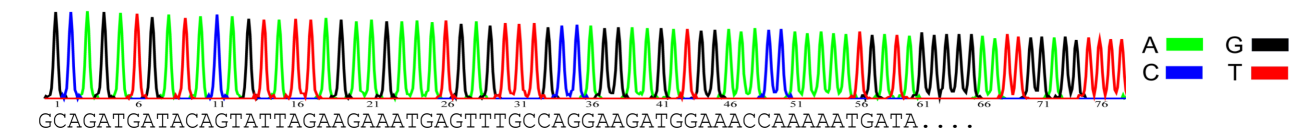
Sequencing Summary

Sanger Sequencing:

- Gives the sequence that is complementary to the template strand = "top" strand, same strand at the primer.
- The start of the sequencing information is defined by a primer that anneals to the template (therefore some of the sequence has to be known, how this is done will be described later)
- Dideoxy sequencing is carried out by adding both dideoxynucleotide triphosphates (ddNTPs) and deoxyribonucleotide triphosphates (dNTPs) to the synthesis reactions, at a ratio of 1:100. Most growing chains do not terminate.
- ddNTPs are identical to dNTPs except that they lack the 3' hydroxyl group. Because of the missing 3'-OH, DNA polymerization stops once one ddNTP is added to a growing strand.
- The type of the added base is determined by "color coding" each base.
- The location of added bases is determined by measuring the size of the DNA fragment that was terminated by the ddNTP.
- It is possible to sequence approximately 1000 bases by this method.

Next Gen-Sequencing:

- Simultaneous sequencing of a large number of fragments
- Shorter "reads" 100 versus 1000 bases/template



Polymerase Chain Reaction - PCR

- In 1983, Kary Mullis developed the molecular biology technique that has since revolutionized genetic research, earning him the Nobel Prize in 1993.
- PCR had an impact on four main areas of biotechnology: gene mapping, cloning, DNA sequencing, and gene detection (e.g. coronavirus).
- PCR is now used as a medical diagnostic tool to detect specific mutations that may cause genetic disease, in criminal investigations and courts of law to identify suspects on a molecular level, and in the sequencing of the human genome.

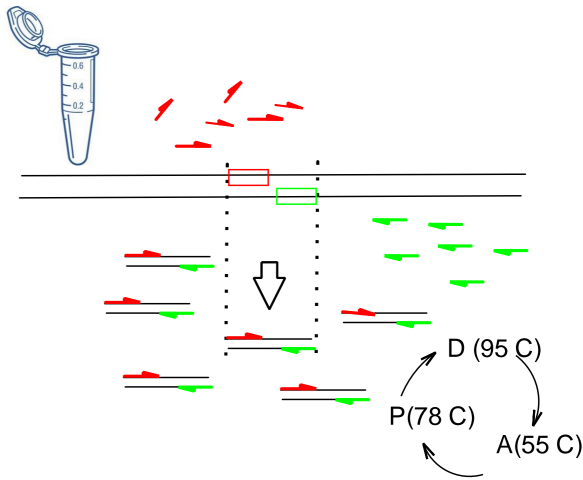


Expectations:

- 1. Be able to explain how PCR works to amplify a segment of DNA.
- 2. Be able to give the left and right primers.
- 3. Apply PCR approaches to determine genotype and detection of viruses.

Polymerase Chain Reaction

- PCR is an in vitro DNA synthesis reaction in which a specific section of DNA is replicated over and over generating exponentially large amounts of a specific piece of DNA from trace amounts of starting material (template).
- Template can be trace amounts of DNA from a drop of blood, a single hair follicle, or a cheek cell.
- The region of DNA that is copied is specified by the sequence of two primers, which are short ssDNA that initiate polymerase activity. The primers are in vast excess over the DNA.
- The location of the amplified segment is defined by two primers (left = upstream, right = downstream):
 - they anneal to their templates according to Watson-Crick pairing rules (A-T, G-C),
 - initiate polymerization from those sites,
 - they are incorporated into the final PCR product.
 - Left primer = sequence of top strand at left boundary
 - Right primer = sequence of bottom strand at right boundary
- The primers are DNA and are synthesized chemically, they can be any desired sequence.
- If there is no homology between the primers and the input DNA, then no PCR product will be formed.



Each PCR cycle consists of three steps:

- 1. Denaturation of the DNA to make it single stranded (2 min at 98 C)
- 2. Lowering of temperature to let the primers form double-stranded DNA (1 min at 55 C)
- 3. Elongation by DNA polymerase (1 min/kb at 78 C)

PCR – Primer Design

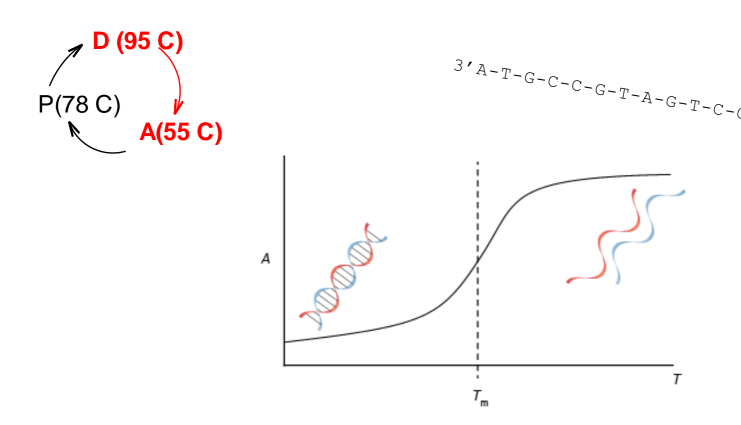
- Before a region of DNA can be amplified, one must identify and determine the sequence of a piece of DNA upstream and downstream of the region of interest.
- These areas are then used to determine the sequence of oligonucleotide primers that will be synthesized and used as starting points for DNA replication.
- Primers are complimentary to the up- and downstream regions of the sequence to be amplified, so they stick, or anneal, to those regions.
 - Left primer = sequence of top strand on the left. This primer will anneal to the bottom strand.
 - Right primer = sequence of bottom strand on the right. This primer will anneal to the top strand.
- Primers are in large excess over the template DNA, they are never used up.
- The primers are incorporated into the final PCR product.

(a) PCR primers must bind to sequences on either side of the target sequence, on opposite strands. Primer annealing site Primer Region of DNA to be amplified by PCR annealing site (b) When target DNA is single stranded, primers bind and allow DNA polymerase to work. Amplified region Amplified region 5'--AAGCTGACTAGTCGATGCGAATGTGCGGTGC-3' 3'--TTCGACTGATCAGCTACGCTTACACGCCACG-5' Know these rules! <mark>CTGAC</mark>TAGTCGATGCGAATGTGC

GACTGATCAGCTACGCTTACACG

Note: Actual primer lengths are 20-30 bases, in the illustrations here and on problem sets, much shorter primers are used.

PCR Step 1 - Thermal Stability of Double Stranded DNA (dsDNA)



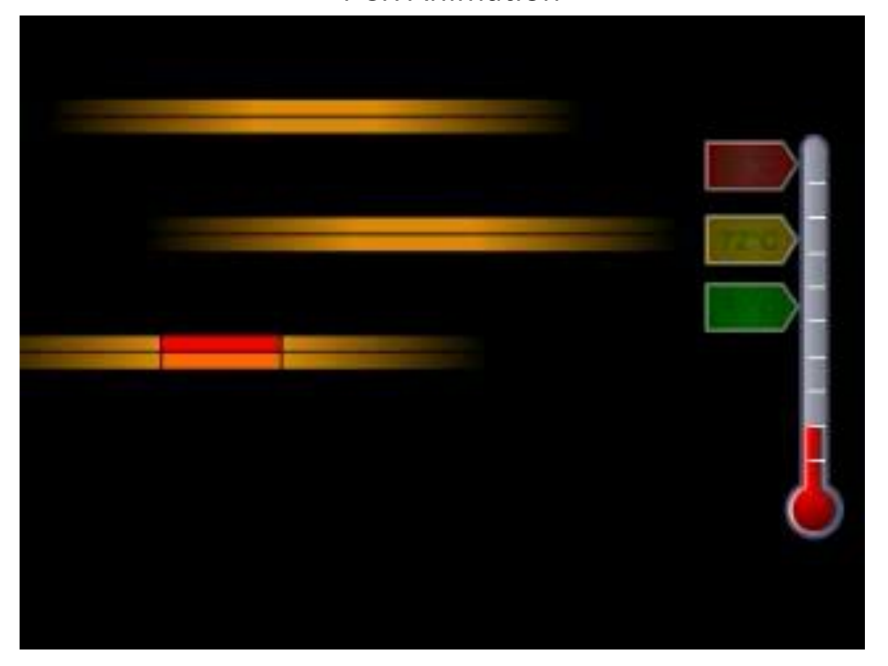
Polymerase Characteristics

- Since the first step of each cycle (D) requires heating to high temperature, a thermostable polymerase is required.
- The first polymerase, Taq, was isolated from Thermus Aquaticus, a bacterial living in hot springs (Yellowstone National Park)
- A number of different polymerases with improved properties have been developed.

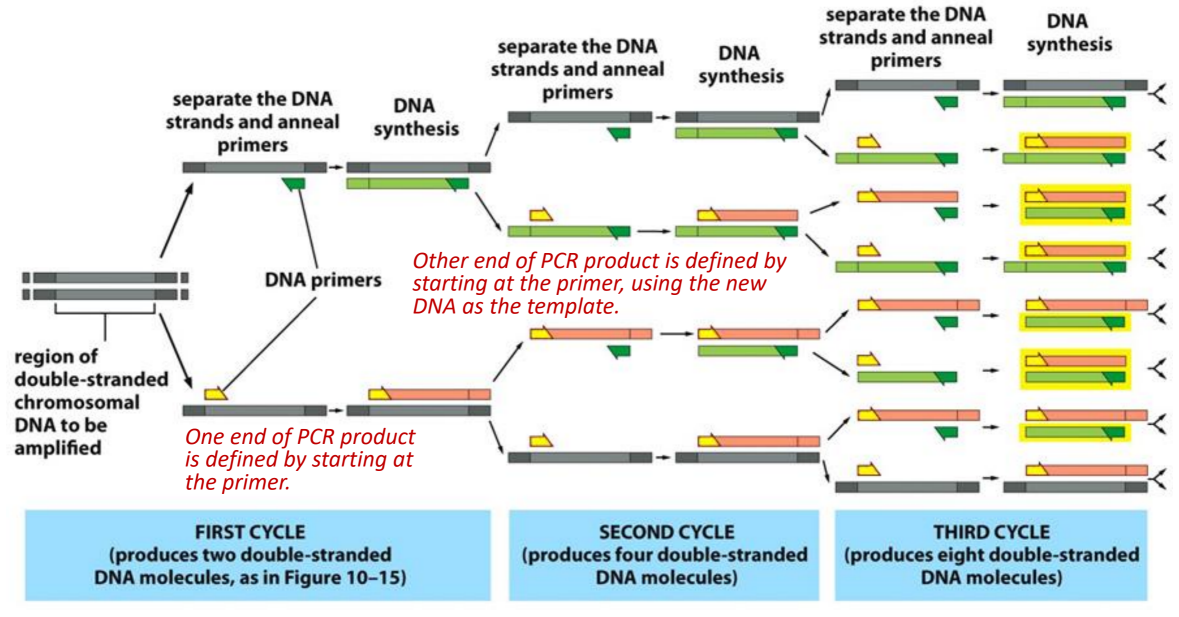
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PCR Animation

Watch Me!

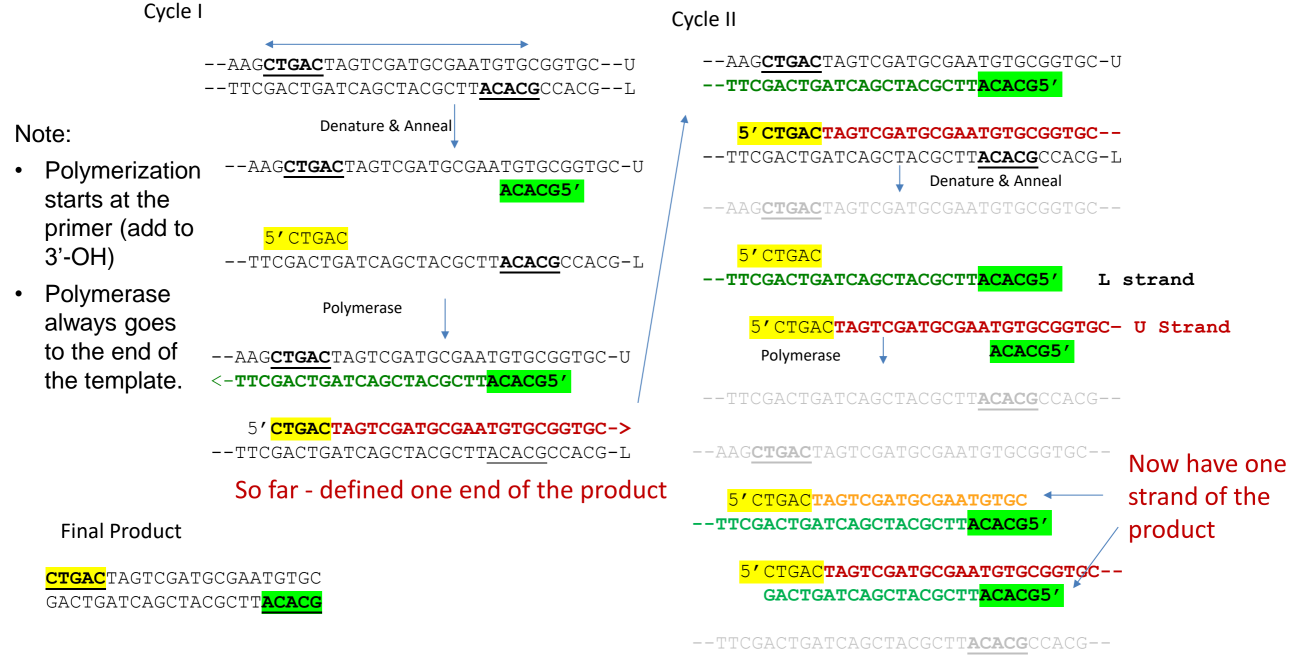


Three PCR Cycles



After 30 cycles there will be 2³⁰, or over 1 billion times more copies than at the beginning!!!

Detailed Events During First Three PCR Cycles



Detailed Events during first Three PCR Cycles

Cycle 3

5'CTGACTAGTCGATGCGAATGTGC --TTCGACTGATCAGCTACGCTTACACG5 5'CTGACAGTCGATGCGAATGTGCGGTGC--GACTGTCAGCTACGCTTACACG5' **Denature & Anneal** 5'CTGACTAGTCGATGCGAATGTGC ACACG5 --TTCGACTGATCAGCTACGCTTACACG5 <mark>5 'CTGAC</mark>AGTCGATGCGAATGTGCGGTGC--5'CTGAC GACTGTCAGCTACGCTTACACG5 **Polymerase** 5'CTGACTAGTCGATGCGAATGTGC GACTGATCAGCTACGCTTACACG5 --TTCGACTGATCAGCTACGCTTACACG5 TGACAGTCGATGCGAATGTGCGGTGC--5'CTGACAGTCGATGCGAATGTGC GACTGTCAGCTACGCTTACACG5

Example – follow the PCR cycles for the following template with primers 5' AATT (left) and 5' GGCC (right)

---AATT-----GGCC----

Now have complete PCR product.
The product will double in each of the following cycles.

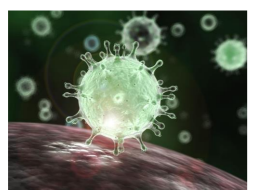
Note that the primers are the first bases at the ends of each strand of the PCR product.

AATT-----GGCC TTAA-----CCGG

PCR Applications - Detection of Viruses

Sequence of Covid-19 (top strand only)

Coronavirus



1	attaaaggtt	tataccttcc	caggtaacaa	accaaccaac	tttcgatctc	ttgtagatct
61	gttctctaaa	cgaactttaa	aatctgtgtg	gctgtcactc	ggctgcatgc	ttagtgcact
121	cacgcagtat	aattaataac	taattactgt	cgttgacagg	acacgagtaa	ctcgtctatc
181	ttctgcaggc	tgcttacggt	ttcgtccgtg	ttgcagccga	tcatcagcac	atctaggttt

gaacaaact aaaatgtctg ataatggacc ccaaaatcag cgaaatgcac cccqcattac

28321	gtttggtgga	ccctcagatt	caactggcag	taaccagaat	ggagaacgca	gtggggcgcg
28381	atcaaaacaa	catcaacccc	aaggtttacc	caatáatact	gcatcttagt	teaccgctct
28441	cactcaacat	ggcaaggaag	accttaaatt	ccctcgagga	caaggcgttc	caattaacac
29701	gggaggactt	gaaagagcca	ccacattttc	accgaggcca	cgcggagtac	gatcgagtgt
29761	acagtgaaca	atgctaggga	gagctgccta	tatggaagag	ccctaatgtg	taaaattaat
29821	tttagtagtg	ctatccccat	gtgattttaa	tagcttctta	ggagaatgac	aaaaaaaaa

CDC Recommended PCR Primers

2019-Novel Coronavirus (2019-nCoV) Real-time rRT-PCR Panel Primers and Probes						
Name	Description	Oligonucleotide Seq	Label ¹	Working Conc.		
2019-nCoV_N1-F	2019-nCoV_N1 Forward Primer	5'-GAC CCC AAA ATC AGC GAA	None	20 μΜ		
2019-nCoV_N1-R	2019-nCoV_N1 Reverse Primer	5'-TCT GGT TAC TGC CAG TTG	AT CTG-3'	None	20 μΜ	

dsSeq of above bold & circled region

28271 aaaatgtctgataatg<mark>GACCCCAAAATCAGCGAAAT</mark>gcaccccgcattacgtttggtggaccctcagattcaactggcagtaaccagaatggagaacgca ttttacagactattacctggggttttagtcgctttacgtggggcgtaatgcaaaccacctgggaa<mark>GTCTAAGTTGACCGTCATTGGTCT</mark>tacctcttgcgt

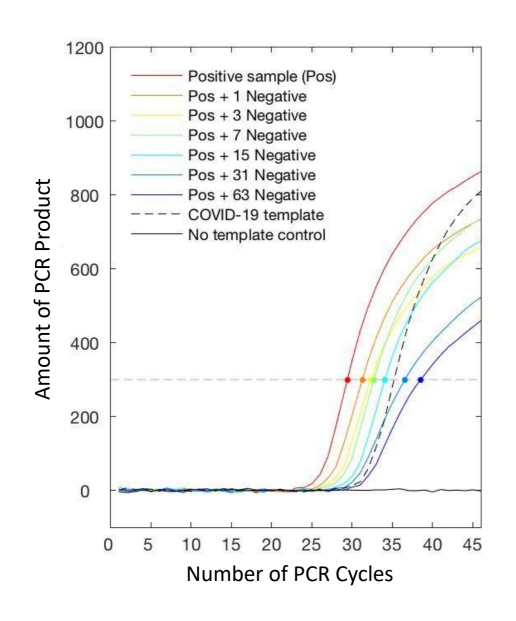
29881 aaaaaaaaaa aaaaaaaaa aaa

PCR Product

GACCCCAAAATCAGCGAAATGCACCCCGCATTACGTTTGGTGGACCCTCAGATTCAACTGGCAGTAACCAGACTGGGGGTTTTAGTCGCTTTACGTGGGGGCGTAATGCAAACCACCTGGGAGTCTAAGTTGACCGTCATTGGTCT

Will PCR generate products if the viral DNA is not present?

Covid 19 PCR Test: Detection of the PCR Product.



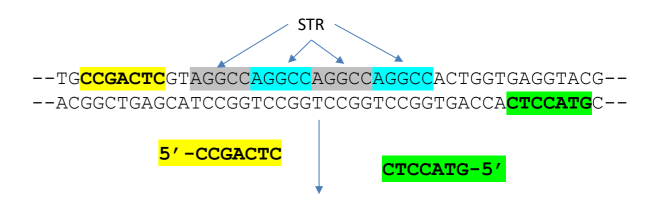
https://www.medrxiv.org/content/10.1101/2020.03.26.20039438v1

- Production of PCR products (double stranded DNA) causes an increase in signal (fluorescence)
- Signal above 300 considered to be positive (dashed gray line)
- Dots represent when a sample crosses the fluorescence threshold.
- Red curve (Positive sample) shows a threshold level of PCR product after 27 cycles.
- Next 6 samples are the positive sample mixed with up to 63 negative samples, showing that it is possible to test pooled samples.
- --- is a *positive control* amount of Covid template. It shows that you can detect a PCR product if the covid genome is present.
- Solid black line is a *negative control*, no Covid DNA. It shows that addition of covid template will lead to a signal.

PCR Applications – Identification of Individuals

- Regions of DNA have variable numbers of repeated DNA sequences (Short tandem repeats, STR). The number of STR can differ from one person to the next.
- Individuals will inherit one copy of the repeat from each parent. The length of the inherited DNA can be the same or different.
- PCR Primers are designed to be outside the repeated region, so that they will anneal to a single location on the chromosome and then amplify the region containing the STR
- PCR Product length = primer lengths

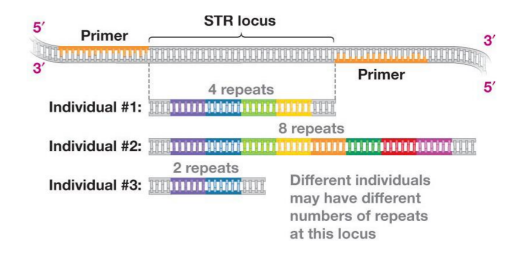
 number of tandem repeats (+ any DNA between the primers and the repeats). Individuals can be differentiated by the length of the PCR product if they have different numbers of STR



CCGACTCGTAGGCCAGGCCAGGCCAGGCCACTGAGGTACGGCTGAGCATCCGGTCCGGTCCGGTCCGGTGACCACCATG

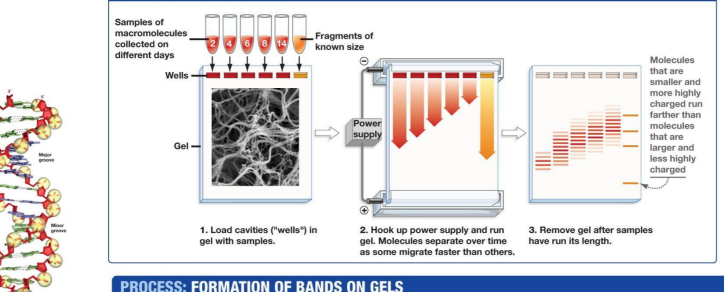
Which individual has the shortest PCR product?

Which has the longest?

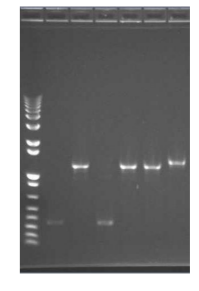


Size Determination of PCR products - Agarose Gel Electrophoresis.

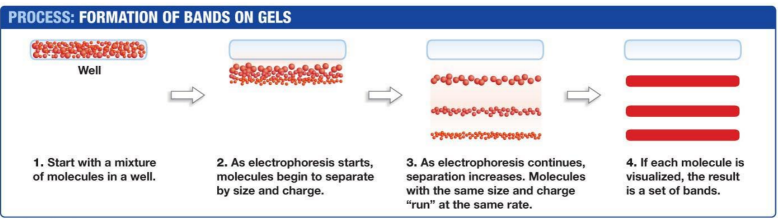
https://dnalc.cshl.edu/resources/animations/gelelectrophoresis.html



PROCESS: GEL ELECTROPHORESIS

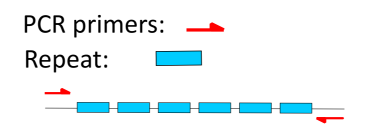


Which are the smallest PCR fragments?



Short Tandem Repeats to Test Paternity

- 1. DNA samples (blood, cheek cells) would be obtained from:
 - Mother
 - Child
 - Candidate fathers.
- 2. PCR would be preformed using primers that amplify a segment of the chromosome containing repeats.
- 3. Each individual would show 2 bands on the gel, corresponding to the PCR product from each chromosome (we have two copies of each chromosome).
- 4. The child would inherit one copy from the mother and the other from the father:
 - One of the child's PCR product would match one of the mothers.
 - The other PCR product from the child would match one of the PCR products from the father.



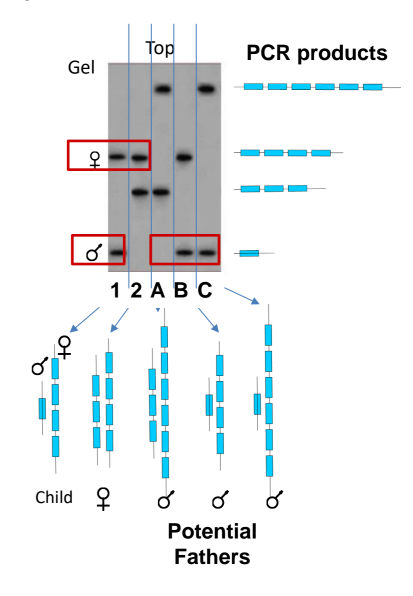
Lane 1: Child

Lane 2: Mother

Lanes A, B, C: Possible Fathers

- 1. Which PCR product is from the mother? From the father?
- 2. Who is **not** the father?

3. Who **may** be the father?



Introduction to Immunology

- 1. Branches of the immune system (Innate and acquired)
- 2. Properties of antibodies (Quaternary structure, antigen recognition)
- 3. How antibodies are produced:
 - Genome DNA changes
 - mRNA splicing
- 4. How antibodies eliminate pathogens

Key Questions:

- 1. Why is the innate system important?
- 2. What is the origin of diversity in acquired immunity?
- 3. How are antibodies made.

The Nobel Prize in Physiology or Medicine 2018





James P. Allison Prize share: 1/2

Tasuku Honjo

Prize share: 1/2

The Nobel Prize in Physiology or Medicine 2018 was awarded jointly to James P. Allison and Tasuku Honjo "for their discovery of cancer therapy by inhibition of negative immune regulation."

Some Important Definitions:

Antigen = something that is recognized by the immune system, e.g. bacteria, virus, pollen.

Epitope = the part of the antigen that is contacted by the antibody.

Antibody (Ab) = Y-shaped protein that recognizes antigens, found on the surface of B-cells or secreted by plasma cells. When bound to antigen, it can initiate a process that results in the destruction of the antigen. Specificity is high due to AA sequence in the variable segments.

Immunoglobulin (Ig) = antibody.

B-cell = involved in antibody production and recognition of pathogen. Has antibody molecule on its surface (as part of the B-cell receptor). Develops into plasma cells after activation by T_H cells. Called B-cells because they are generated in the organ called the Bursa in birds.

Plasma cell = derived from B-cell after activation of the B-cell, produces secreted antibodies with the same specificity as the original B-cell.

 T_H cell = T-helper: Required to activate both B and T_C cells, as well as other cells in the immune system. Called T-cells because they mature in the thymus.

 T_c cell = T-cellular: Involved in defense against viruses and cancer.

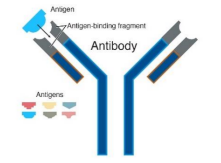
TCR = \underline{T} - \underline{c} ell \underline{r} eceptor – found on the surface of T-cells, recognizes MHC proteins + bound peptide, RTK.

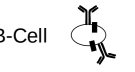
- **T**_c **cell** = recognizes MHC I + peptide
- T_H cell = recognizes MHC II + peptide

MHC = \underline{m} ajor \underline{h} istocompatibility \underline{c} omplex – required for acquired immunity (basis of transplant rejection)

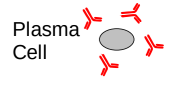
- MHC I = protein found on the surface of *all* cells, "presents" peptides derived from the proteins that were made by the cell. The MHC-peptide complex is recognized by T_c cells. *Only foreign* peptides produce a response.
- MHC II = on the surface of B-cells, macrophages, and dendritic cells. Presents external peptides to T_H cells, leading to activation of the cell by T_H cells. *Only foreign peptides produce a response*.











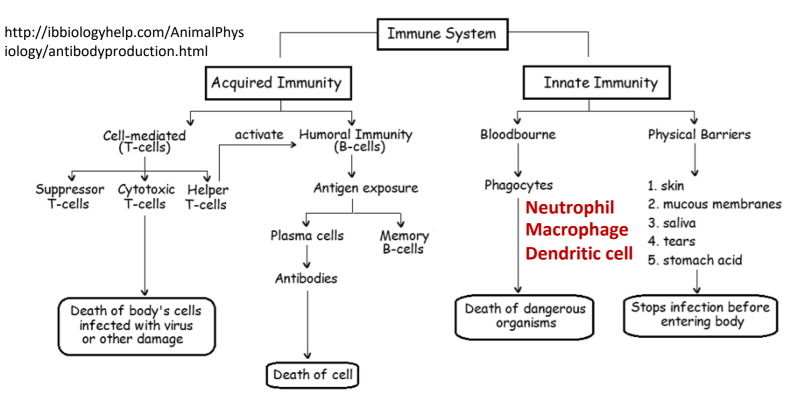
T cell







Branches of the Immune System:

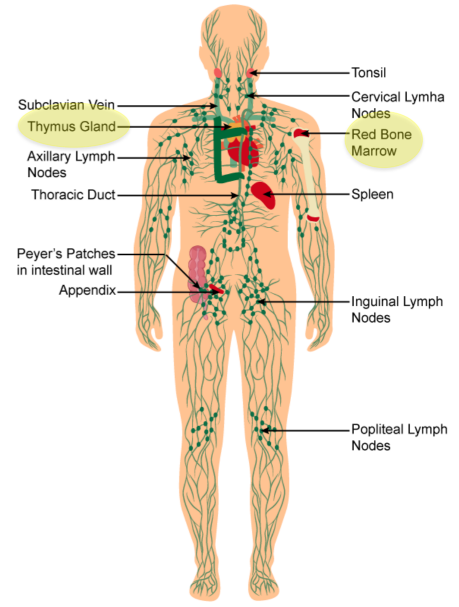


Why is the innate system essential?

- A pathogen doubles every hour.
- It takes 7 days to produce antibody (after 1st exposure)
- Uncontrolled growth would produce many bacteria: $2^{24 \times 7} = 3.7 \times 10^{50}$ (~10³⁰ kg)

Important **primary** lymphatic organs: bone marrow (B), thymus (T)-Generate all immune cell.

Important **secondary** lymphatic organs: lymph nodes, spleen, Peyer's patches – Activation of immune cells.



https://www.topperlearning.com/

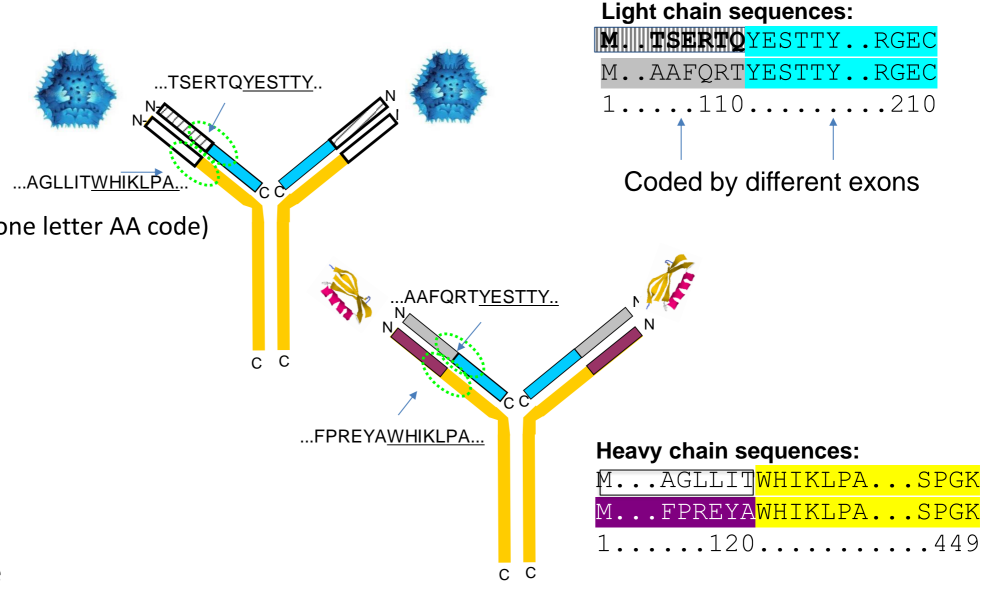
Each Antibody:

- Two identical light chains
- Two identical heavy chains
- First ~100 Amino acids on each chain are ...AGLLITWHIKLPA... called the variable region and differ from antibody to antibody.
- Unique sequence for variable region of both heavy and light chains

 defines specificity –
 different antibodies
 bind different

 antigens.
- Constant regions same protein sequence for all.

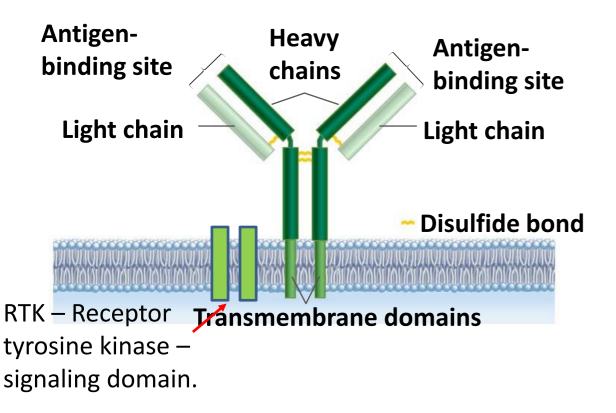
Antibody Structure and Diversity



Production of Antibodies by B-cells & Plasma Cells

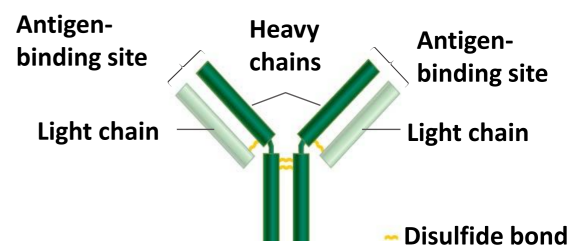
B- Cells & B-cell Receptor (BCR)

- Each B-cell has only one type of antibody as part of its BCR (B-cell receptor), i.e. the 10⁵ BCRs are homogeneous on the same cell.
- Approximately 10⁸ different specificities at any one time. i.e. 10⁸ different B-cells!



Plasma Cells:

- After activation, a B-cell develops into a plasma cell.
- The antibody is secreted.
- The same light chains are produced.
- The heavy chains differ only in the absence of the transmembrane domains.

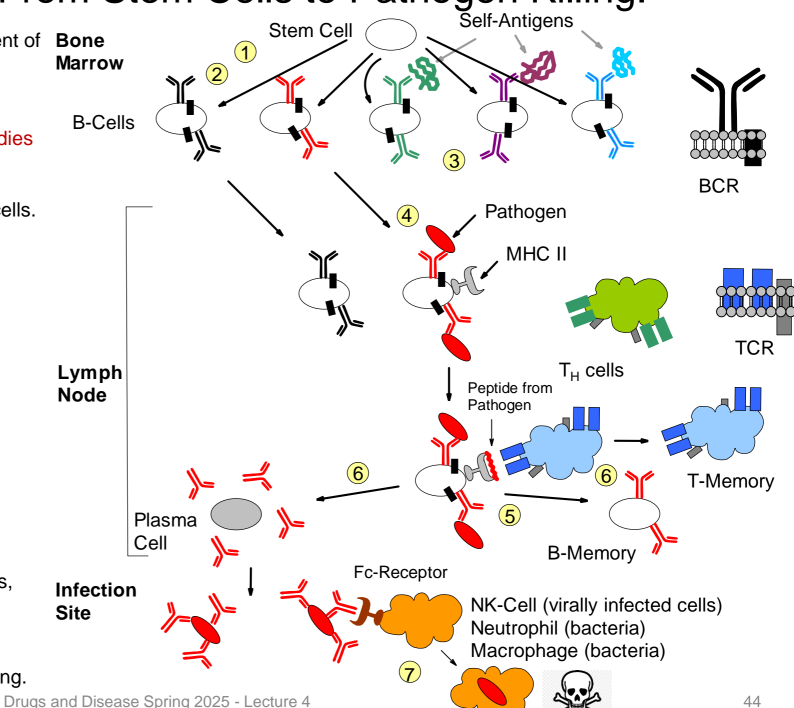


mRNA that codes for antibodies contains two types of sequences:

- Exons contain codons for the amino acids
- Introns removed before translation
 Different exons are used to produce membrane bound or soluble antibodies.

B-Cell Biology - From Stem Cells to Pathogen Killing.

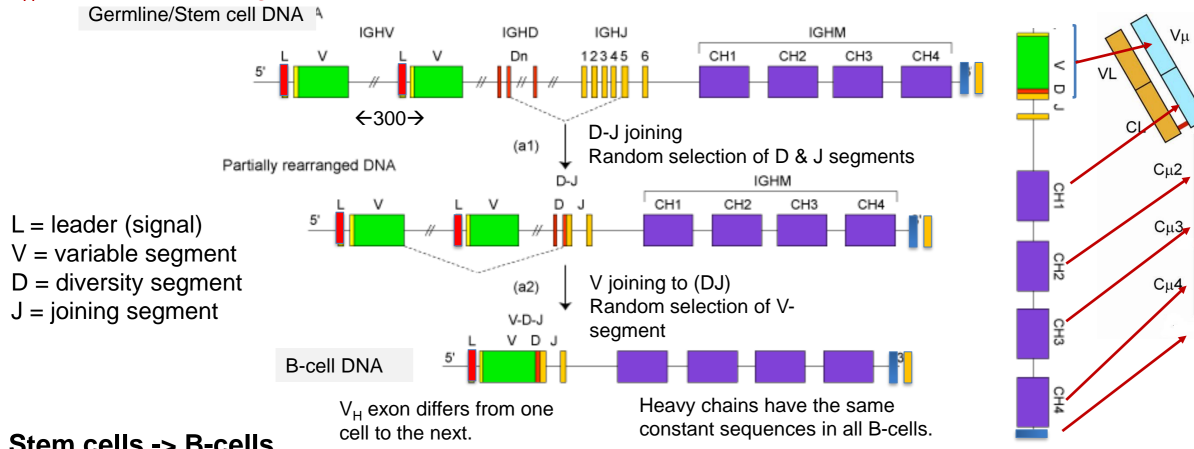
- **1.** Generation of high diversity of chains during development of stem cells to B-cells in bone marrow.
- DNA rearrangements to generate functional exons for variable segments of both light and heavy chain.
- 2. Molecular & cellular biology of membrane bound antibodies on cell surface = B-cell receptor (BCR)
- Transcriptional enhancers, mRNA splicing
- Light chain and heavy chain exported to surface of B-cells.
- **3.** Self tolerance test to prevent autoimmune diseases, autoreactive B-cells eliminated.
- 4. Encounter and capture of antigen in lymph nodes
- **5.** Activation of B-cells by T_H cells
- Peptides from pathogen presented on major histocompatibility proteins (MHC II).
- T-cell activation by tyrosine kinase receptors (T-cell Receptor, TCR), secretion of signaling molecules.
- **6.** Development of
- Plasma cells Production of soluble antibodies of the same specificity as the parent B-cell.
- B-memory cells (basis of immunity)
- T-memory cells (basis of immunity)
- 7. Destruction of Pathogens
- Fc region of antibody binds to Fc Receptor on NK cells, neutrophiles, macrophages
- Pathogen internalized and destroyed.
- **BCR** B-cell receptor = antibody + signaling chains.
- **TCR** T cell receptor = MHC-peptide recognition + signaling.



Antibody Genes are Assembled From DNA Segments: Giving many different sequences.

Production of Heavy Chain Gene: V_H exon = V+D+J segment (selected at random)

The mRNA coding for antibodies contains 5 exons.



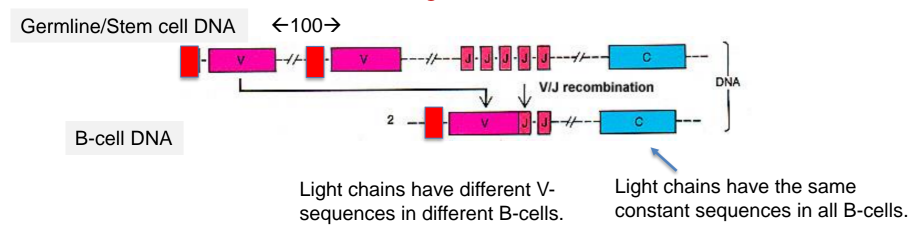
Stem cells -> B-cells

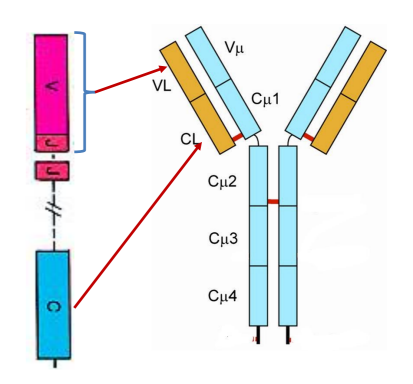
- The exon that codes for the variable region of the heavy chain is generated by the random joining of a V, D, and J DNA segments.
- Each B-cell will generate a unique sequence for its heavy and light chain DNA.
- This is a permanent change to the DNA (*genome*) of the B-cell.

1. If there are 300 possible V-heavy segments, 10 possible D segments, and 6 possible J segments, how many different heavy chains can be made?

Light-chain Genes are Assembled From DNA Segments: Giving many different sequences.

Production of Light Chain Gene





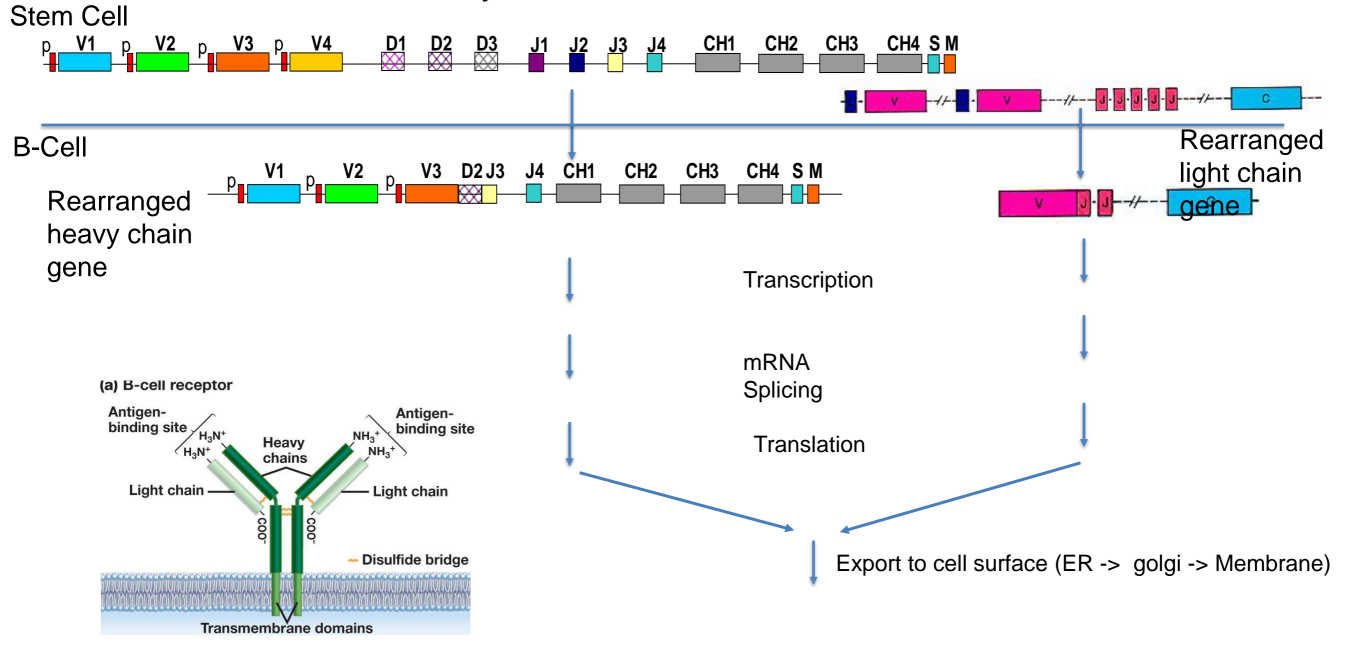
Stem cells -> B-cells

- In the case of the light chain, the variable region is generated by VJ joining.
- Each B-cell will generate a unique sequence for its heavy and light chain DNA.
- This is a permanent change to the DNA (genome) of the B-cell.

Antibody Diversity

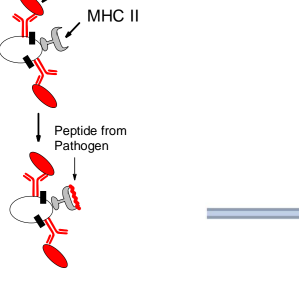
- 1. If there are 100 possible V-heavy segments and 5 possible J segments, how many different light chains can be made?
- 2. If any possible heavy chain can pair with any possible light chain, how many different antibodies can be generated, assuming there are 10,000 possible heavy chains and 500 different light chains?

Antibody Production – From Stem Cells to B-Cells



Antigen Capture by B-Cells - Endocytic Pathways

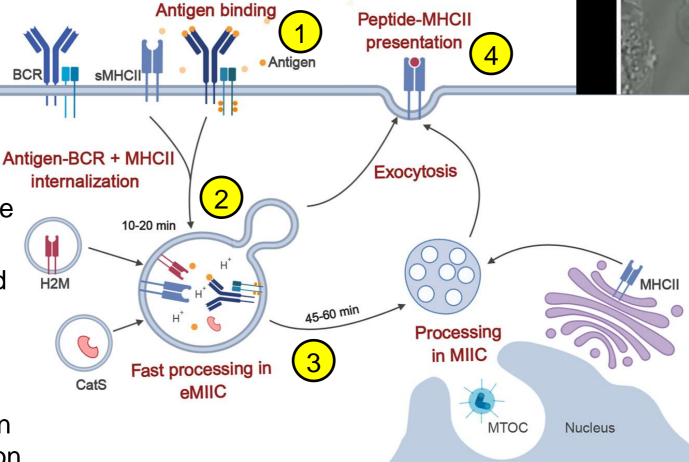
Endocytosis of bacteria by a B-cell



Pathogen

1. Antigen binds to variable domains of antibody on the BCR (B-cell receptor)

- 2. Antigen is internalized and digested into peptides
- 3. Peptides are loaded on to class II MHC
- 4. Peptide-MHC displayed on membrane for presentation to T-cells



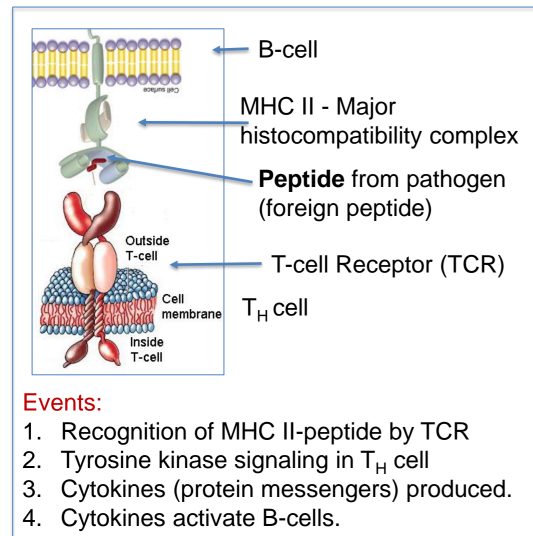
Journal of Cell Science doi: 10.1242/jcs.235192

Bacteria labeled with Green fluorescent protein.

- I. Capture of the bacteria
- Internalization (endocytosis)
- III. Degradation of the bacterial proteins, producing peptides.

Activation of B cells by Antigen - Lymph Node

B-cells develop into



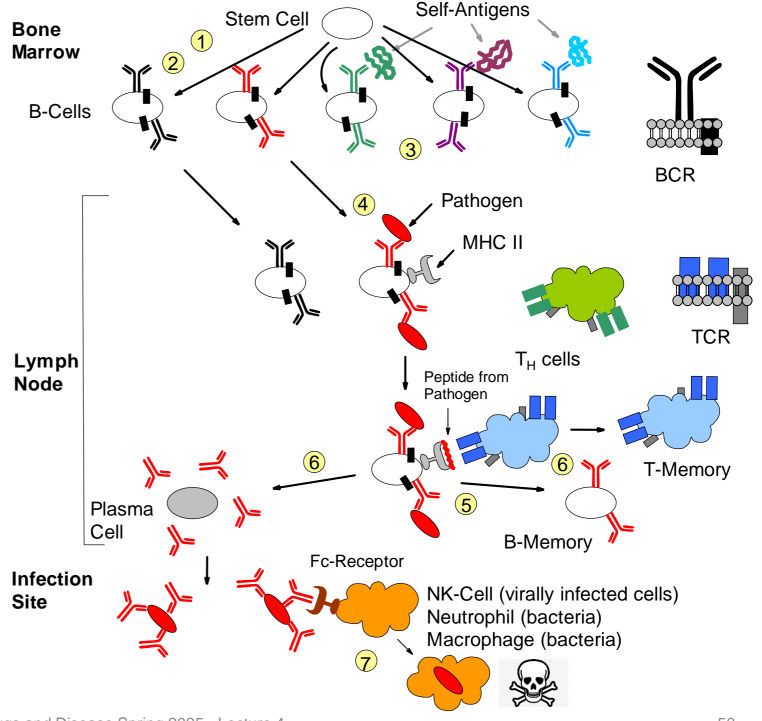
antibody secreting plasma cells. Pathogen B and T-helper cells MHC II develop into memory cells, that are longlived and are quickly activated by the same pathogen. This is the T_H cells basis of vaccination. Peptide from Pathogen 6 T-Memory (5)Plasma Cell

- Soluble antibody from plasma cells has the same light and heavy chains as the original B-cell.
- Membrane anchors are missing, so antibody is secreted outside the cell.

B-Memory

Can you:

- Describe how the genes for the heavy and light chain are generated, and how this give rise to many different antibodies?
- Do you understand the process of B-cell activation, including presentation of foreign peptides on MHC II and the role of the T-helper cell.
- Describe how antibodies inactivate pathogens?



Cell Based Immunology

Key Questions:

- 1. How does your immune system fight viruses?
- 2. How does your immune system detect and destroy cancer cells?
- 3. How can the immune response be engineered to fight cancer?

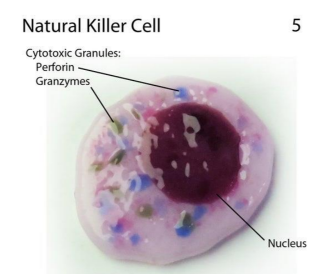
Cell Types:

Innate

 Natural Killer (NK) cell

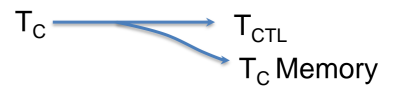
Acquired

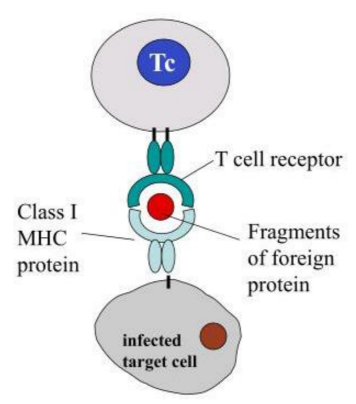
- T_H
- T_C, T_{CTL}



NK: Innate

- Kill virally infected cells
- Kill cancer cells





Activation of Tc cells requires:

- T_C Memory 1. Recognition of *foreign* peptide on MHC I.
 - 2. Assistance from Thelper cells.

Activated Tc cell becomes a cytotoxic T-lymphocyte T_{CTL}

$\mathsf{T}_{\mathsf{CTL}}$

- Kill virally infected cells
- Kill cancer cells

Activation of Tc-Cells

A. Dendritic Cells Acquire Antigen from Viruses and Cancerous Cells

PROCESS: MHC ANTIGEN PRESENTATION

fragment

Peptide fragment Major , histocompatibility **Endosome** (MHC) protein **Dendritic cell** MHC protein Peptide

1/25/2025

Antigen 1. Dendritic cell ingests antigen via phagocytosis (intact virus, cell debris from cancer cell).

2. Enzymes break antigen proteins into peptide fragments.

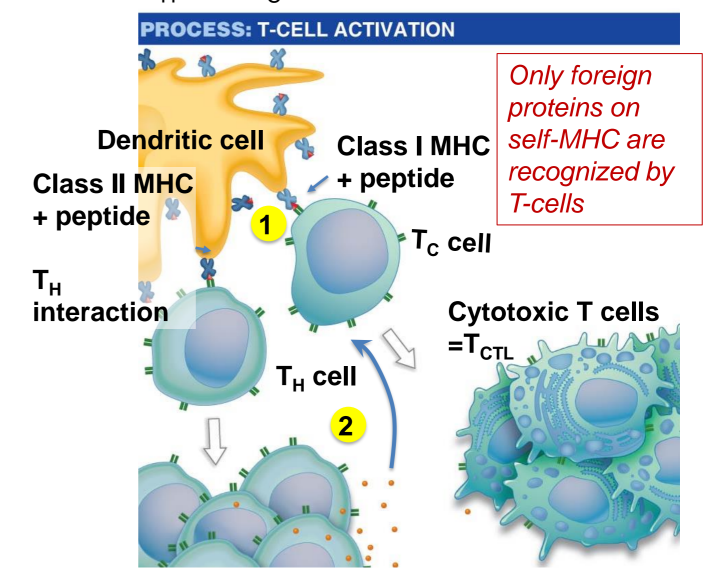
3. Peptide fragments are loaded onto **both** class I and class II MHC proteins in endosomes.

- 4. MHC I & II –peptide complex is transported to cell surface.
- 5. MHC protein presents peptide fragment on cell Surface to T-H and T-C cells.

Activation of Tc-Cells B. Dendritic Cells Activate T_H and T_C cells.

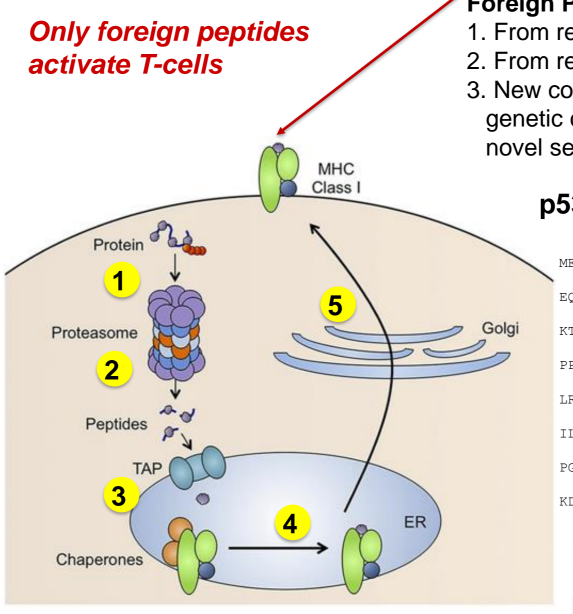
Activation of Tc cells requires:

- Recognition of foreign peptide on MHC I by TCR on Tc cell
- Assistance from T-helper cells via secreted messengers (small proteins called cytokines)



T_c Detection of Diseased/Cancer Cells - Role of MHC I

- MHC I present peptides
- Peptides are generated from of all of the proteins that are made in the cell.
- Steps:
- protein targeted for degradation by ubiquitin
- 2. Protein digested by proteasome
- 3. Peptides transported into ER
- 4. Peptides loaded on to MHC I
- 5. Peptide/MHC complex transported to cell membrane.



Foreign Peptide Source:

- 1. From replication of viruses in the cell
- 2. From replication of intracellular bacteria (e.g. TB)
- 3. New coding sequences in cancer cells due to genetic changes (e.g. mutations in p53 lead to novel sequences).

p53 Protein Sequence

		Zn Fingers (DiyA binding)		
10	20	30	40	50
MEEPQSDPSV	EPPLSQETFS	DLWKLLPENN	VLSPLPSQAM	DDLMLSPDDI
60	70	80	90	100
EQWFTEDPGP	DEAPRMPEAA	PPVAPAPAAP	TPAAPAPAPS	WPLSSSVPSQ
110	120	130	140	150
KTYQGSYGFR	LGFLHSGTAK	SVTCTYSPAL	NKMFCQLAKT	CPVQLWVDST
160	170	80	190	200
PPPGTRVRAM	AIYKQSQHMT	EVV <mark>RRC</mark> PH <mark>H</mark> E	RCSDSDGLAF	PQHLIRVEGN
210	220	230	240	250
LRVEYLDDRN	TFRHSVVVPY	EPPEVGSDCT	TIHYNYM <mark>C</mark> NS	S <mark>C</mark> MGGMNRRP
260	270	280	290	300
ILTIITLEDS	SGNLLG R NSF	EVRVCA.CPGR	DRRTEEENLR	KKGEPHHELP
310	320	330	340	350
PGSTKRALPN	NTSSSPQPKK	KPLDGEYFTL	QIRGRERFEM	FRELNEALEL
360	370	380	390	
KDAQAGKEPG	GSRAHSSHLK	SKKGQSTSRH	KKLMFKTEGP	DSD



Normal seq., ignored by TCR

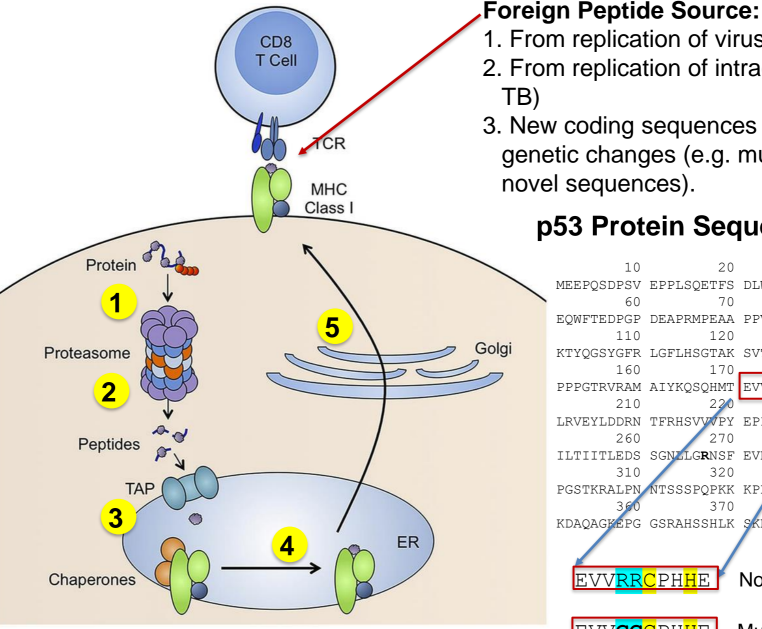
7n Einaara (DNA hindina)



Mutant seq. in cancer, detected by TCR

T_c Detection of Diseased/Cancer Cells - Role of MHC I

- MHC I present peptides
- Peptides are generated from of all of the proteins that are made in the cell.
- Steps:
 - protein targeted for degradation by ubiquitin
- Protein digested by proteasome
- Peptides transported into FR
- Peptides loaded on to MHC I
- Peptide/MHC complex transported to cell membrane.



1. From replication of viruses in the cell

2. From replication of intracellular bacteria (e.g.

3. New coding sequences in cancer cells due to genetic changes (e.g. mutations in p53 lead to novel sequences).

> p53 Protein Sequence 7n Finance (DNA hinding)

		Zn Fingers (DiyA binding)		
10	20	30	40	50
MEEPQSDPSV	EPPLSQETFS	DLWKLLPENN	VLSPLPSQAM	DDLMLSPDDI
60	70	80	90	100
EQWFTEDPGP	DEAPRMPEAA	PPVAPAPAAP	TPAAPAPAPS	WPLSSSVPSQ
110	120	130	140	150
KTYQGSYGFR	LGFLHSGTAK	SVTCTYSPAL	NKMFCQLAKT	CPVQLWVDST
160	170	80	190	200
PPPGTRVRAM	AIYKQSQHMT	EVV <mark>RRC</mark> PH <mark>H</mark> E	RCSDSDGLAF	PQHLIRVEGN
210	220	230	240	250
LRVEYLDDRN	TFRHSVVVPY	EPPEVGSDCT	TIHYNYM <mark>C</mark> NS	S <mark>C</mark> MGGMNRRP
260	270	280	290	300
ILTIITLEDS	SGNLLG R NSF	EVRVCA.CPGR	DRRTEEENLR	KKGEPHHELP
310	320	330	340	350
PGSTKRALPN	NTSSSPQPKK	KPLDGEYFTL	QIRGRERFEM	FRELNEALEL
360	370	380	390	
KDAQAGKEPG	GSRAHSSHLK	SKKGQSTSRH	KKLMFKTEGP	DSD

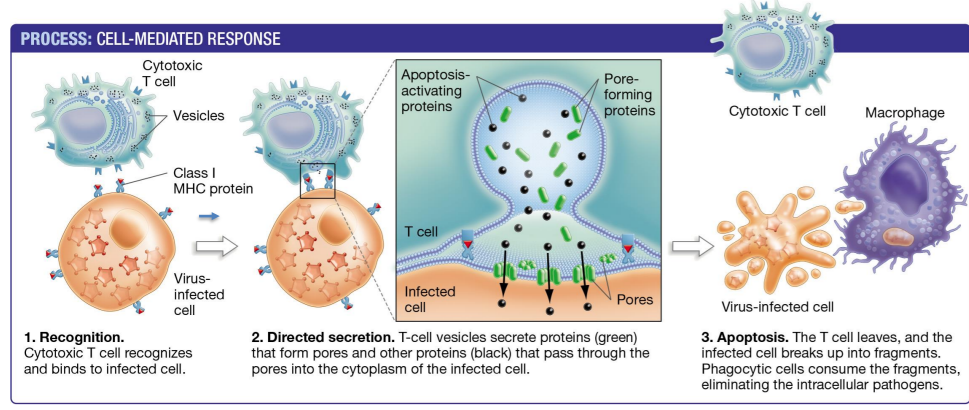


Normal seq., ignored by TCR



Mutant seq. in cancer, detected by TCR

T_C Cells: Detection and Killing of Virally Infected or Cancer Cells



Cytotoxic T-Lymphocyte Killing Target

S James A. Sullivan Quill Graphics Charlottesville, VA USA

Cancer cell or Infected cell

 Granzymes enter through perforin pore and cause cell undergo programmed cell death (apoptosis)

Summary Questions for Immunology:

- 1. What are the two major branches of the immune system? Why are both important?
- 2. What are the roles of different cell types in each system, e.g. what would happen if T_H-cells disappeared?
- 3. What is the quaternary structure of an antibody? Can you sketch an antibody and indicate where the antigen binds?
- 4. What defines the specificity of antibodies?
- 5. What are the steps in the production of antibody genes, at the molecular level:
 - a) How do DNA rearrangements produce functional heavy and light chain genes
 - b) How are is the mature mRNA generated in B-cells and Plasma cells.
 - c) What is the difference between the heavy chain export process for B-cells and plasma cells.
- 6. Can you describe how antibodies kill/inactivate pathogens
- 7. How are virally infected cells and tumor cells recognized by Tc cells?
- 8. How does the Tc cell kill those cells?