



BIMM 143

Genome Informatics I

Lecture 14

Barry Grant
UC San Diego

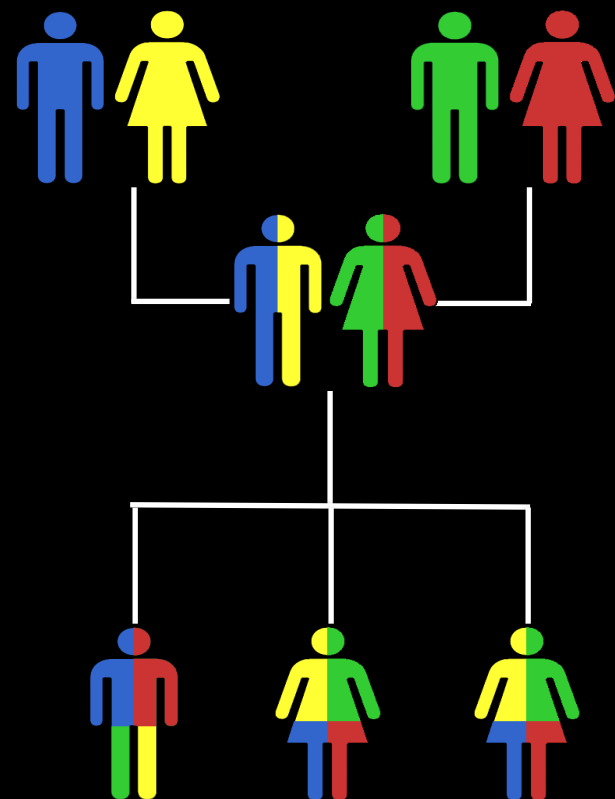
<http://thegrantlab.org/bimm143>

Todays Menu:

- **What is a Genome?**
 - Genome sequencing and the Human genome project
- **What can we do with a Genome?**
 - Compare, model, mine and edit
- **Modern Genome Sequencing**
 - 1st, 2nd and 3rd generation sequencing
- **Workflow for NGS**
 - RNA-Sequencing and Discovering variation

What is a genome?

The total genetic material of an organism by which individual traits are encoded, controlled, and ultimately passed on to future generations

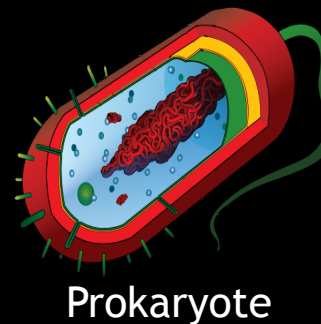


Genetics and Genomics

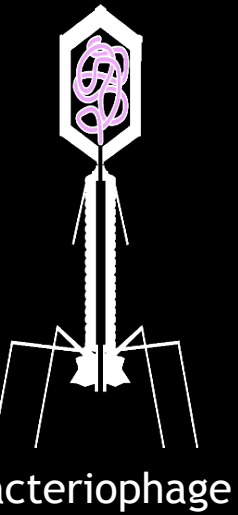
- **Genetics** is primarily the study of *individual genes*, mutations within those genes, and their inheritance patterns in order to understand specific traits.
- **Genomics** expands upon classical genetics and considers aspects of the *entire genome*, typically using computer aided approaches.

Genomes come in many shapes

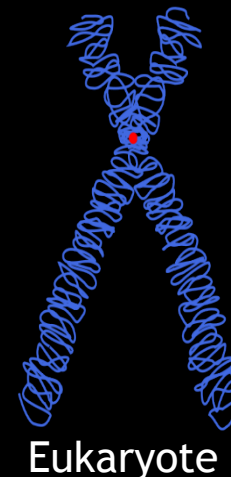
- Primarily DNA, but can be RNA in the case of some viruses
- Some genomes are circular, others linear
- Can be organized into discrete units (chromosomes) or freestanding molecules (plasmids)



Prokaryote



Bacteriophage

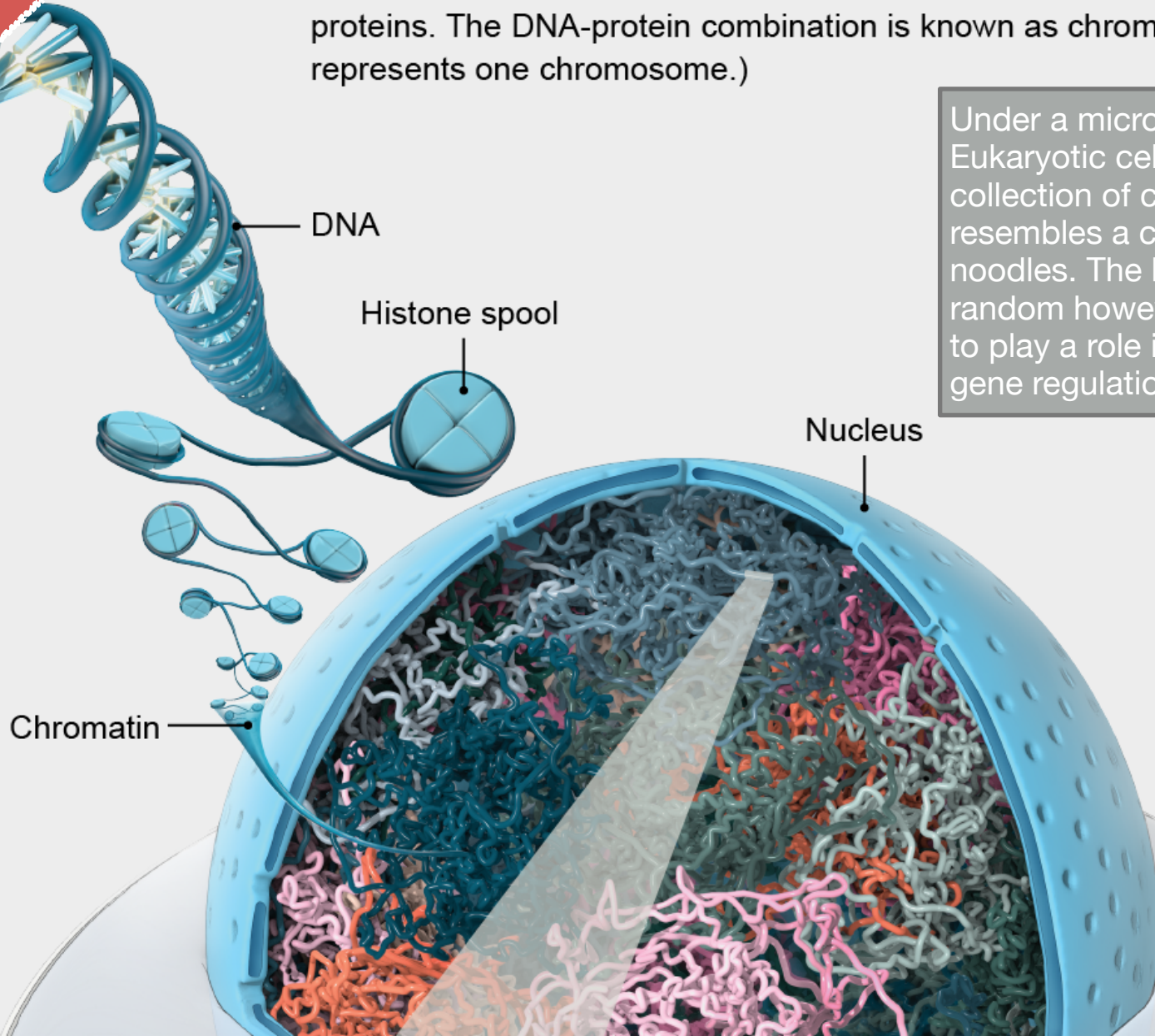


Eukaryote

Side note!

CHROMOSOMES CLOSE-UP

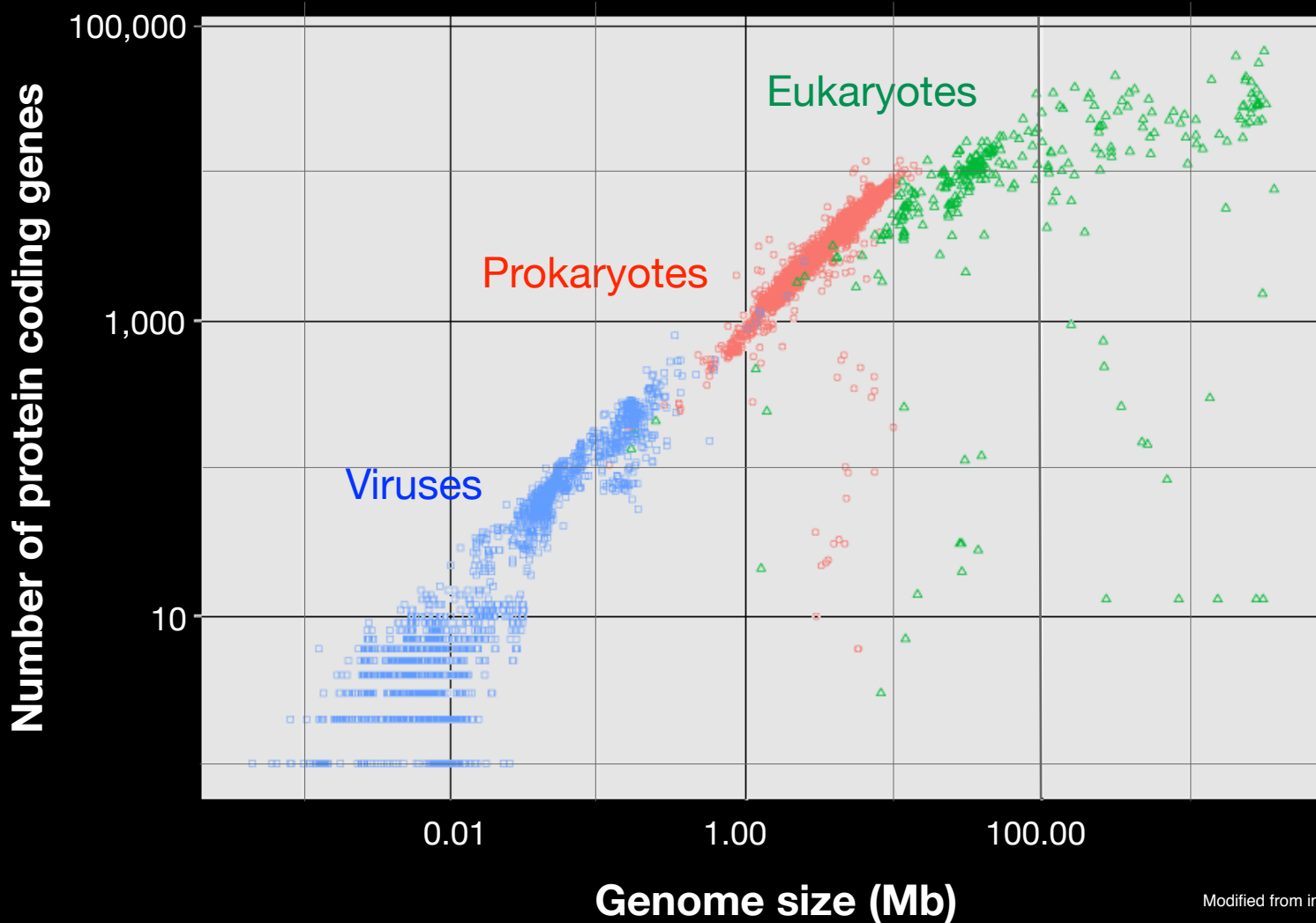
Chromosomes consist largely of double-helical DNA. Cells package the DNA into the nucleus by wrapping it around “spools” composed of histone proteins. The DNA-protein combination is known as chromatin. (Each color represents one chromosome.)



Under a microscope, a Eukaryotic cell's genome (i.e. collection of chromosomes) resembles a chaotic jumble of noodles. The looping is not random however and appears to play a role in controlling gene regulation.

Image credit:
[Scientific American](#)
March 2019

Genomes come in many sizes



Genome Databases

NCBI Genome:

<http://www.ncbi.nlm.nih.gov/genome>

NCBI Resources ▾ How To ▾

Sign in to NCBI

Genome

Search

Limits Advanced

Help

Genome

This resource organizes information on genomes including sequences, maps, chromosomes, assemblies, and annotations.

Using Genome

- Help
- Browse by Organism
- Download / FTP
- Download FAQ
- Submit a genome

Custom resources

- Human Genome
- Microbes
- Organelles
- Viruses
- Prokaryotic reference genomes

Other Resources

- Assembly
- BioProject
- BioSample
- Map Viewer
- Protein Clusters

Genome Tools

- BLAST the Human Genome
- Microbial Nucleotide BLAST
- TaxPlot (3-way Genome Comparison)

Genome Annotation and Analysis

- Eukaryotic Genome Annotation
- Prokaryotic Genome Annotation
- PASC (Pairwise Sequence Comparison)

External Resources

- GOLD - Genomes Online Database
- Ensembl Genome Browser
- Bacteria Genomes at Sanger
- Large-Scale Genome Sequencing (NHGRI)

You are here: NCBI > Genomes & Maps > Genome

Write to the Help Desk

GETTING STARTED

- NCBI Education
- NCBI Help Manual
- NCBI Handbook
- Training & Tutorials

RESOURCES

- Chemicals & Bioassays
- Data & Software
- DNA & RNA
- Domains & Structures
- Genes & Expression
- Genetics & Medicine
- Genomes & Maps
- Homology
- Literature
- Proteins
- Sequence Analysis
- Taxonomy
- Training & Tutorials
- Variation

POPULAR

- PubMed
- Bookshelf
- PubMed Central
- PubMed Health
- BLAST
- Nucleotide
- SNP
- Gene
- Protein
- PubChem

FEATURED

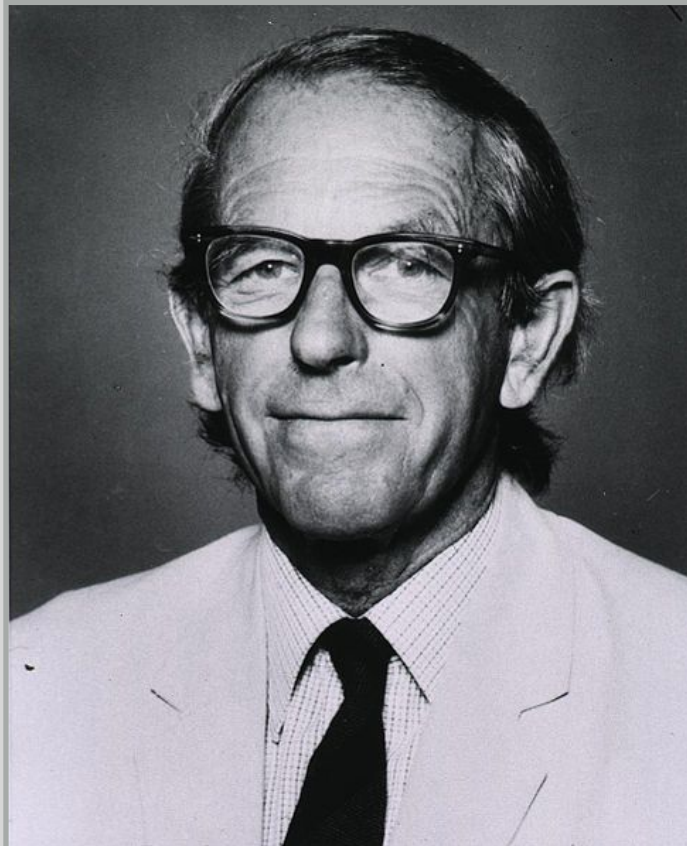
- Genetic Testing Registry
- PubMed Health
- GenBank
- Reference Sequences
- Gene Expression Omnibus
- Map Viewer
- Human Genome
- Mouse Genome
- Influenza Virus
- Primer-BLAST
- Sequence Read Archive

NCBI INFORMATION

- About NCBI
- Research at NCBI
- NCBI News
- NCBI FTP Site
- NCBI on Facebook
- NCBI on Twitter
- NCBI on YouTube

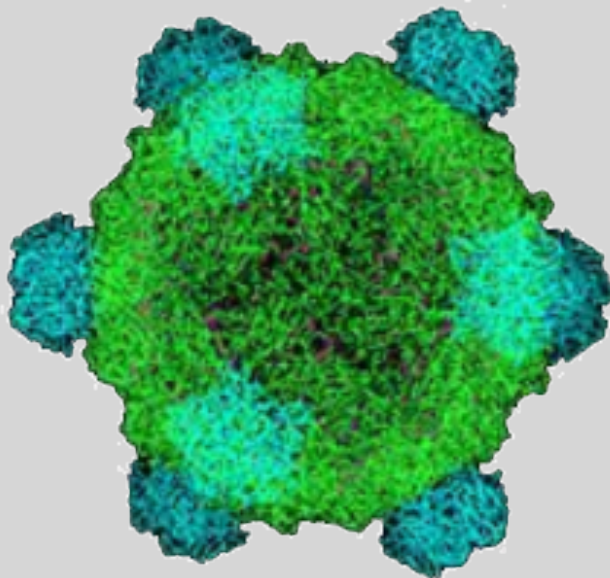
Copyright | Disclaimer | Privacy | Browsers | Accessibility | Contact
National Center for Biotechnology Information, U.S. National Library of Medicine
8600 Rockville Pike, Bethesda MD, 20894 USA

Early Genome Sequencing



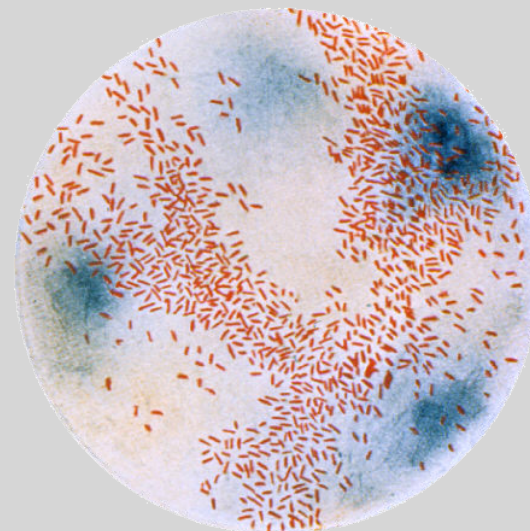
- Chain-termination “Sanger” sequencing was developed in 1977 by Frederick Sanger, colloquially referred to as the “Father of Genomics”
- Sequence reads were typically 750-1000 base pairs in length with an error rate of ~1 / 10000 bases

The First Sequenced Genomes



Bacteriophage φ-X174

- Completed in 1977
- 5,386 base pairs, ssDNA
- 11 genes



Haemophilus influenzae

- Completed in 1995
- 1,830,140 base pairs, dsDNA
- 1740 genes

The Human Genome Project

- The Human Genome Project (HGP) was an international, public consortium that began in 1990
 - Initiated by James Watson
 - Primarily led by Francis Collins
 - Eventual Cost: \$2.7 Billion
- Celera Genomics was a private corporation that started in 1998
 - Headed by Craig Venter
 - Eventual Cost: \$300 Million
- Both initiatives released initial drafts of the human genome in 2001
 - ~3.2 Billion base pairs, dsDNA
 - 22 autosomes, 2 sex chromosomes
 - ~20,000 genes

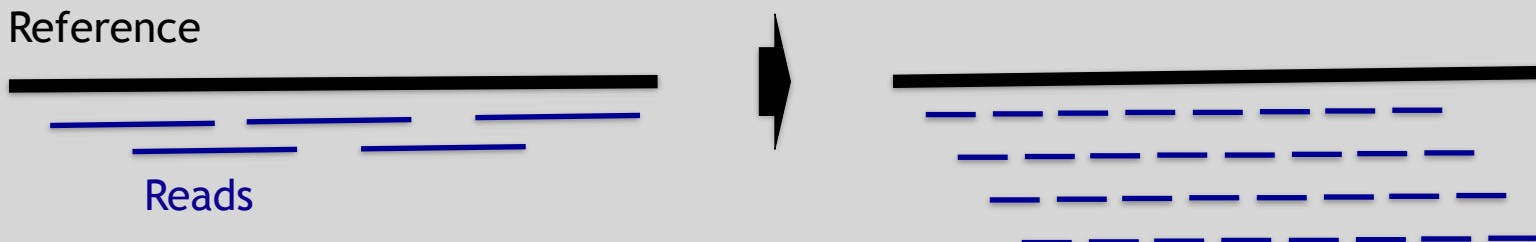




HHMI

Modern Genome Sequencing

- Next Generation Sequencing (NGS) technologies have resulted in a paradigm shift from long reads at low coverage to short reads at high coverage
- This provides numerous opportunities for new and expanded genomic applications



Rapid progress of genome sequencing

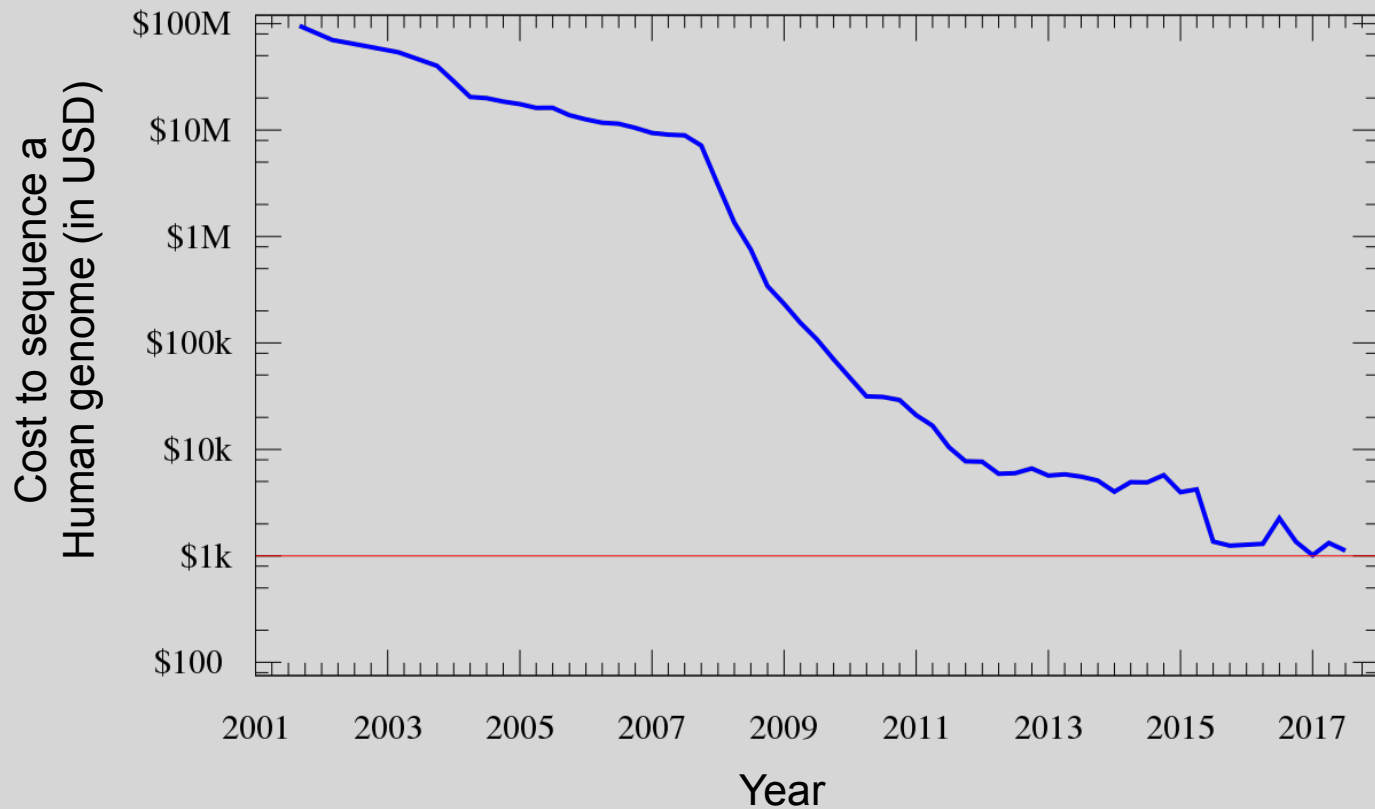


Image source: https://en.wikipedia.org/wiki/Carlson_curve

Rapid progress of genome sequencing

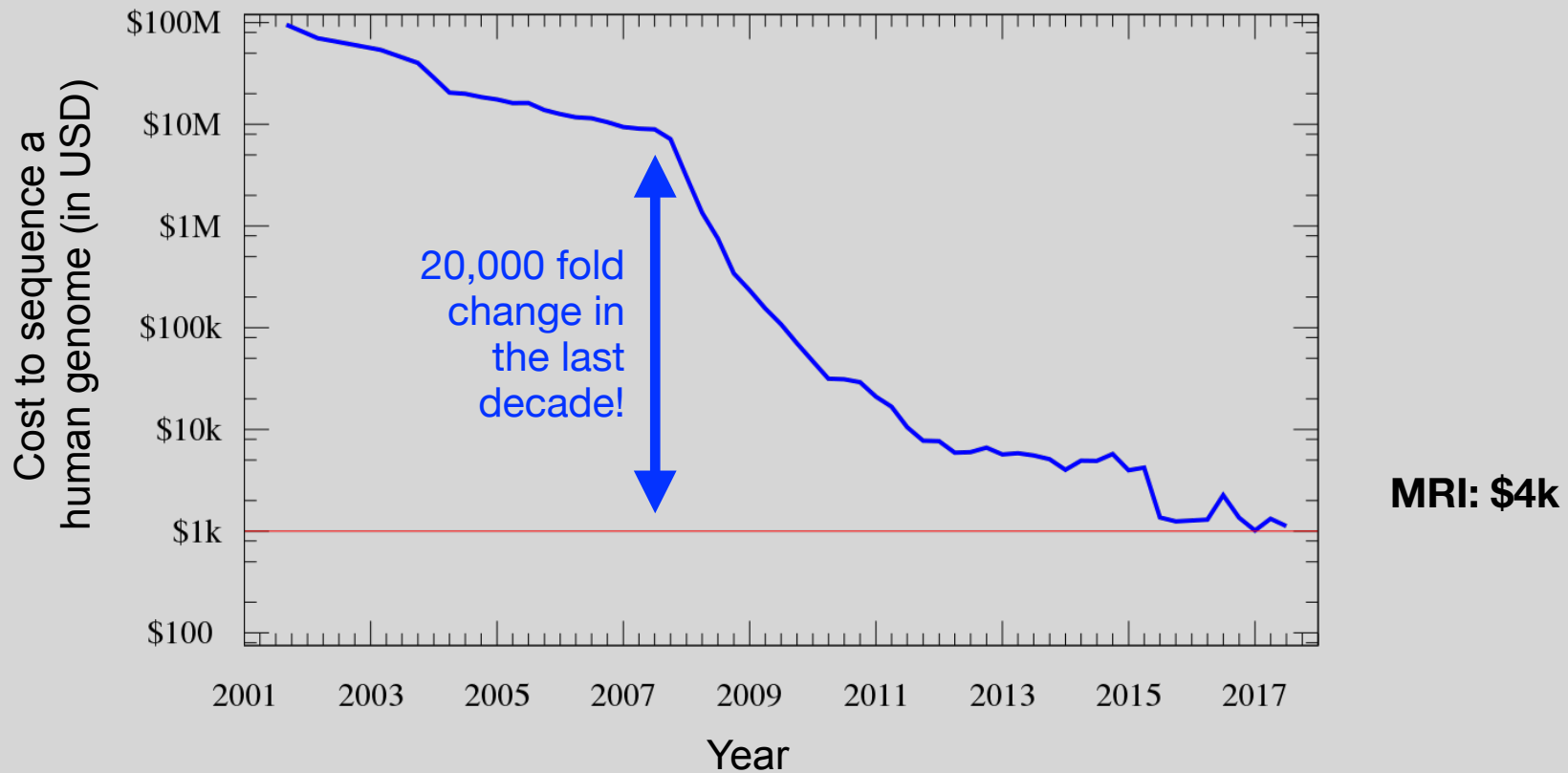
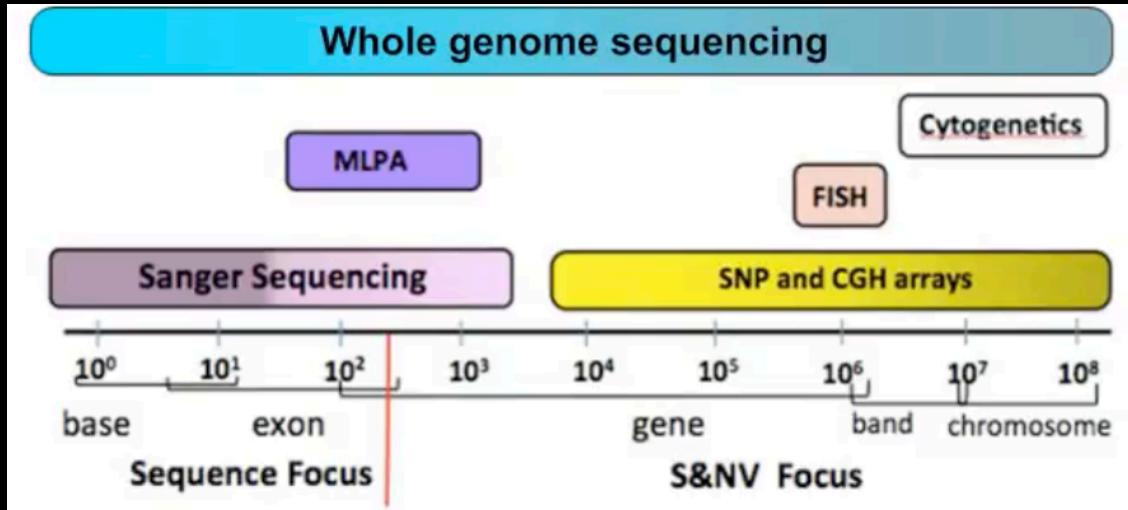


Image source: https://en.wikipedia.org/wiki/Carlson_curve

Whole genome sequencing transforms genetic testing



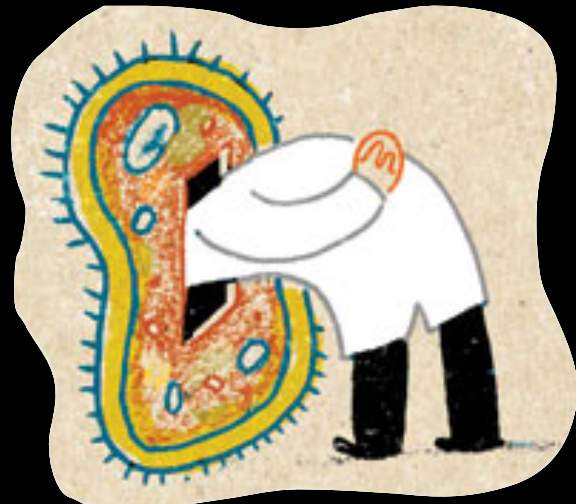
- 1000s of single gene tests
- Structural and copy number variation tests
- Permits hypothesis free diagnosis

Major impact areas for genomic medicine

- **Cancer:** Identification of driver mutations and drugable variants, Molecular stratification to guide and monitor treatment, Identification of tumor specific variants for personalized immunotherapy approaches (precision medicine).
- **Genetic disease diagnose:** Rare, inherited and so-called ‘mystery’ disease diagnose.
- **Health management:** Predisposition testing for complex diseases (e.g. cardiac disease, diabetes and others), optimization and avoidance of adverse drug reactions.
- **Health data analytics:** Incorporating genomic data with additional health data for improved healthcare delivery.

Goals of Cancer Genome Research

- Identify changes in the genomes of tumors that drive cancer progression
- Identify new targets for therapy
- Select drugs based on the genomics of the tumor
- Provide early cancer detection and treatment response monitoring
- Utilize cancer specific mutations to derive neoantigen immunotherapy approaches



What can go wrong in cancer genomes?

| Type of change | Some common technology to study changes |
|-----------------------------|---|
| DNA mutations | WGS, WXS |
| DNA structural variations | WGS |
| Copy number variation (CNV) | CGH array, SNP array, WGS |
| DNA methylation | Methylation array, RRBS, WGBS |
| mRNA expression changes | mRNA expression array, RNA-seq |
| miRNA expression changes | miRNA expression array, miRNA-seq |
| <i>Protein expression</i> | Protein arrays, mass spectrometry |

WGS = whole genome sequencing, WXS = whole exome sequencing

RRBS = reduced representation bisulfite sequencing, WGBS = whole genome bisulfite sequencing

DNA Sequencing Concepts

- **Sequencing by Synthesis:** Uses a polymerase to incorporate and assess nucleotides to a primer sequence
 - 1 nucleotide at a time
- **Sequencing by Ligation:** Uses a ligase to attach hybridized sequences to a primer sequence
 - 1 or more nucleotides at a time (e.g. dibase)

Modern NGS Sequencing Platforms

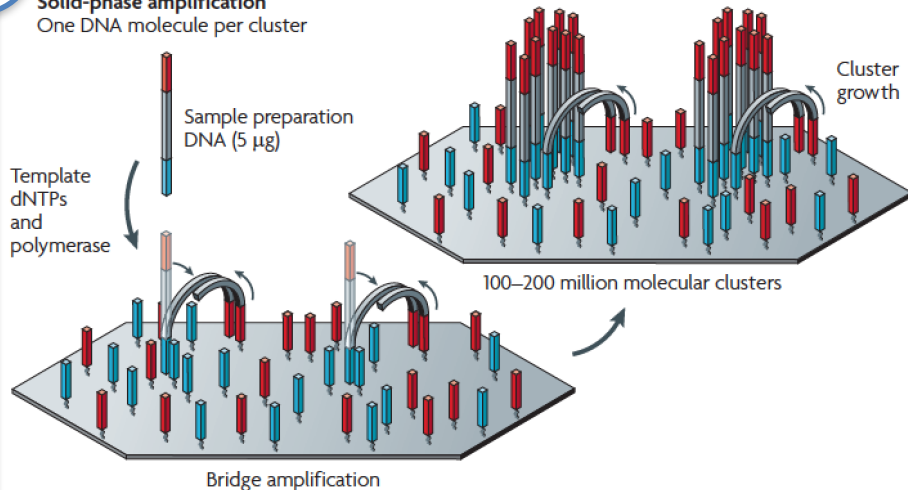
| | Roche/454 | Life Technologies SOLiD | Illumina Hi-Seq 2000 |
|--------------------------------------|---|---|---|
| Library amplification method | emPCR* on bead surface | emPCR* on bead surface | Enzymatic amplification on glass surface |
| Sequencing method | Polymerase-mediated incorporation of unlabelled nucleotides | Ligase-mediated addition of 2-base encoded fluorescent oligonucleotides | Polymerase- mediated incorporation of end-blocked fluorescent nucleotides |
| Detection method | Light emitted from secondary reactions initiated by release of PPi | Fluorescent emission from ligated dye-labelled oligonucleotides | Fluorescent emission from incorporated dye-labelled nucleotides |
| Post incorporation method | NA (unlabelled nucleotides are added in base-specific fashion, followed by detection) | Chemical cleavage removes fluorescent dye and 3' end of oligonucleotide | Chemical cleavage of fluorescent dye and 3' blocking group |
| Error model | Substitution errors rare, insertion/deletion errors at homopolymers | End of read substitution errors | End of read substitution errors |
| Read length (fragment/paired end) | 400 bp/variable length mate pairs | 75 bp/50+25 bp | 150 bp/100+100 bp |

Illumina - Reversible terminators

1

Enzymatic amplification on glass surface

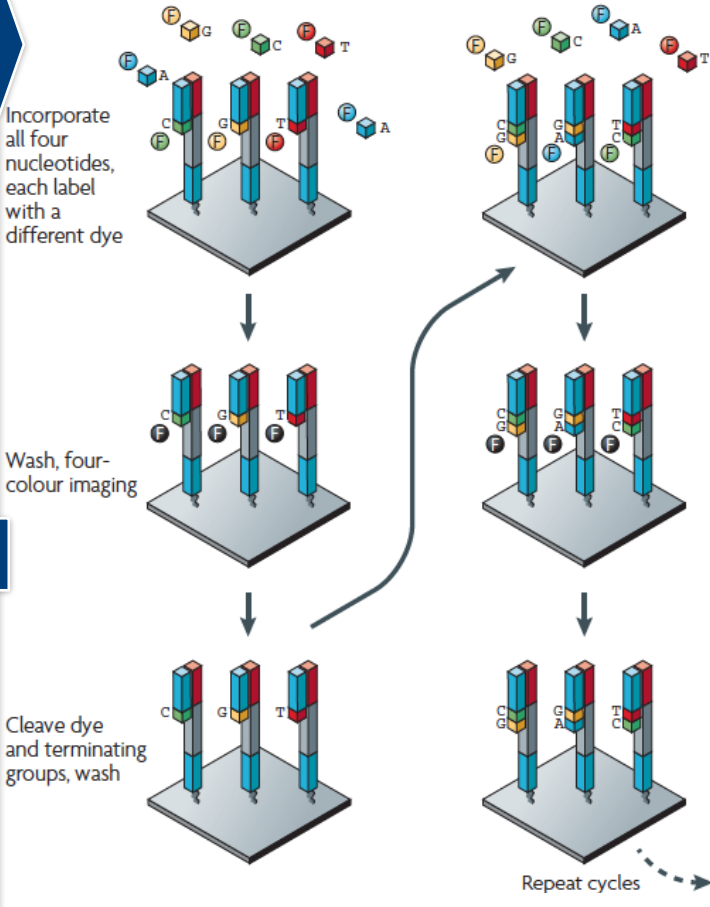
Illumina/Solexa
Solid-phase amplification
One DNA molecule per cluster



2

Polymerase-mediated incorporation of end blocked fluorescent nucleotides

Illumina/Solexa — Reversible terminators



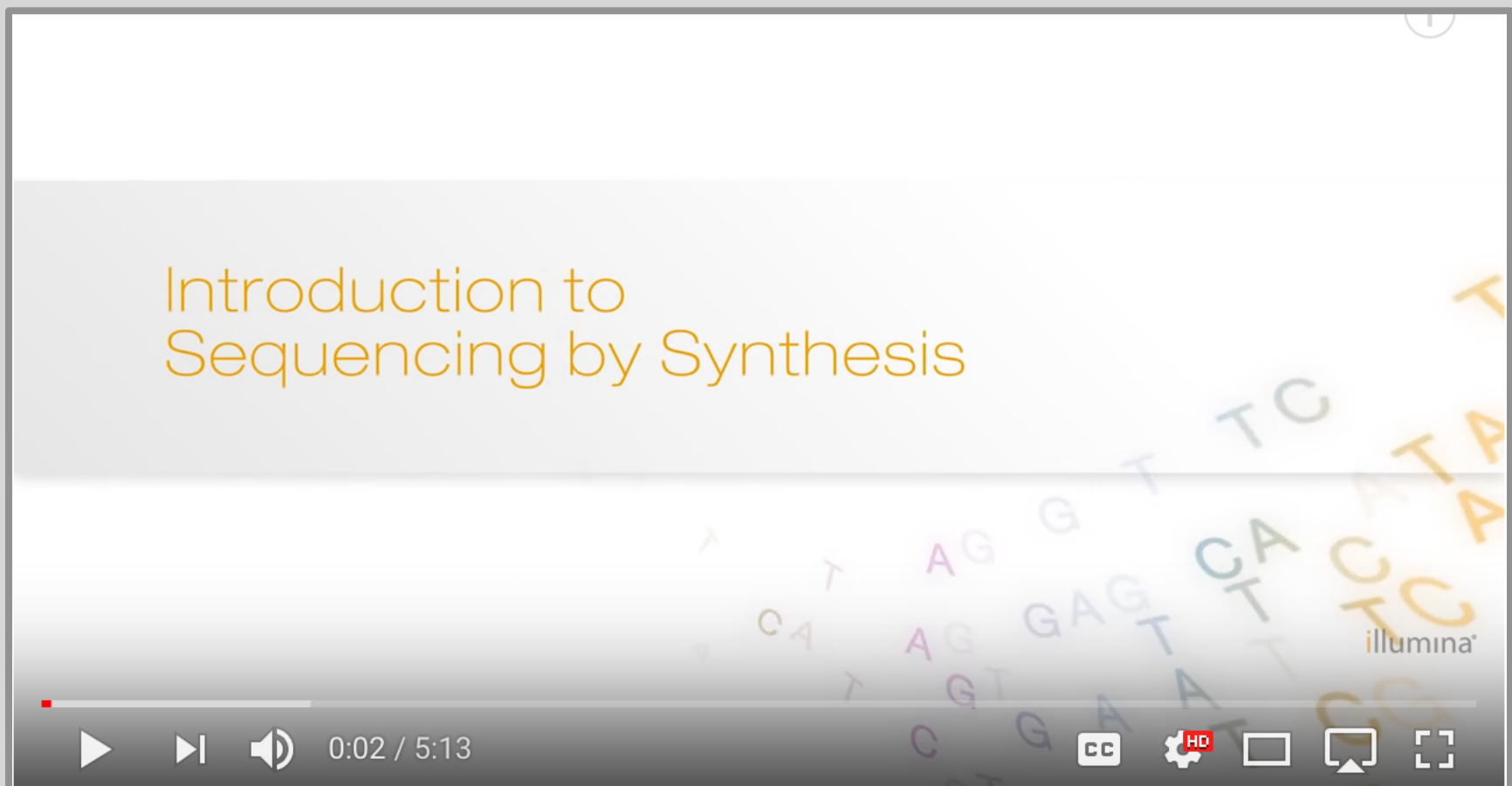
3

Fluorescent emission from incorporated dye-labeled nucleotides



Top: CATCGT
Bottom: cccccc

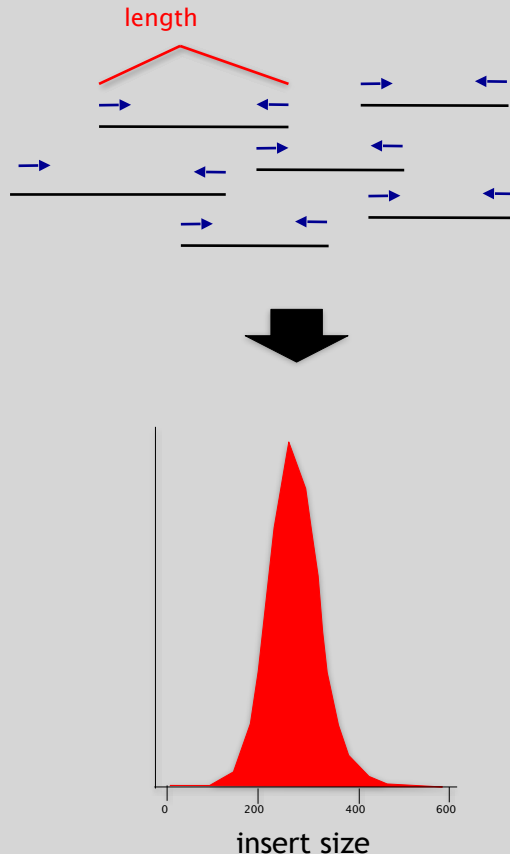
Illumina Sequencing - Video



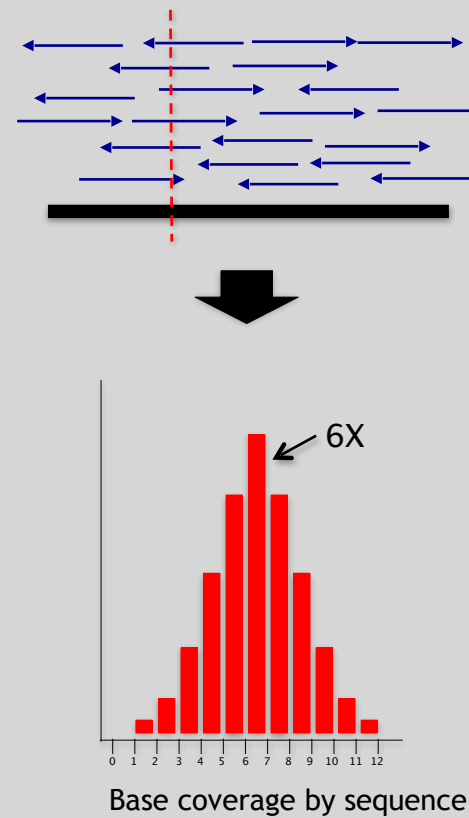
https://www.youtube.com/watch?src_vid=womKfikWlxM&v=fCd6B5HRaZ8

NGS Sequencing Terminology

Insert Size



Sequence Coverage



Summary: “Generations” of DNA Sequencing

| | First generation | Second generation ^a | Third generation ^a |
|--|---|--|--|
| Fundamental technology | Size-separation of specifically end-labeled DNA fragments, produced by SBS or degradation | Wash-and-scan SBS | SBS, by degradation, or direct physical inspection of the DNA molecule |
| Resolution | Averaged across many copies of the DNA molecule being sequenced | Averaged across many copies of the DNA molecule being sequenced | Single-molecule resolution |
| Current raw read accuracy | High | High | Moderate |
| Current read length | Moderate (800–1000 bp) | Short, generally much shorter than Sanger sequencing | Long, 1000 bp and longer in commercial systems |
| Current throughput | Low | High | Moderate |
| Current cost | High cost per base Low cost per run | Low cost per base High cost per run | Low-to-moderate cost per base Low cost per run |
| RNA-sequencing method | cDNA sequencing | cDNA sequencing | Direct RNA sequencing and cDNA sequencing |
| Time from start of sequencing reaction to result | Hours | Days | Hours |
| Sample preparation | Moderately complex, PCR amplification not required | Complex, PCR amplification required | Ranges from complex to very simple depending on technology |
| Data analysis | Routine | Complex because of large data volumes and because short reads complicate assembly and alignment algorithms | Complex because of large data volumes and because technologies yield new types of information and new signal processing challenges |
| Primary results | Base calls with quality values | Base calls with quality values | Base calls with quality values, potentially other base information such as kinetics |

Third Generation Sequencing

- Currently in active development
- Hard to define what “3rd” generation means
- Typical characteristics:
 - Long (1,000bp+) sequence reads
 - Single molecule (no amplification step)
 - Often associated with nanopore technology
 - But not necessarily!

The first direct RNA sequencing by nanopore

- For example this new nanopore sequencing method was just published!
<https://www.nature.com/articles/nmeth.4577>
- "Sequencing the RNA in a biological sample can unlock a wealth of information, including the identity of bacteria and viruses, the nuances of alternative splicing or the transcriptional state of organisms. However, current methods have limitations due to short read lengths and reverse transcription or amplification biases. Here we demonstrate nanopore direct RNA-seq, a highly parallel, real-time, single-molecule method that circumvents reverse transcription or amplification steps."

Side-Note:

SeqAnswers Wiki

A good repository of analysis software can be found at
<http://seqanswers.com/wiki/Software/list>



Page Discussion Read View source View history Go Search

Software/list

< Software

Below is (one of many possible) dynamic tables of software data, created from pages in the wiki. To add a package to the list, use the following form:

new package name

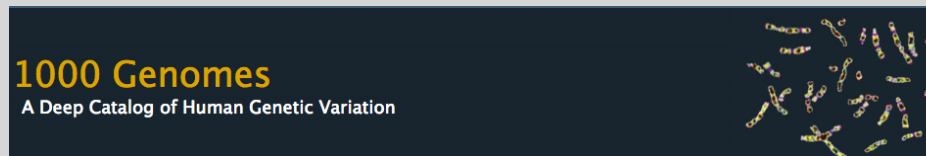
[CSV](#) [JSON](#)

| Name | Summary | Bio Tags | Meth Tags | Features | Language | Licence | OS |
|---------------------|--|----------------------------|--------------------------|------------|----------|-----------------------|----------------------|
| 4peaks | Allows viewing sequencing trace files, motif searching trimming, BLAST and exporting sequences. | Sequencing | Sequence analysis | | | Freeware | Mac OS X |
| AB Large Indel Tool | Identifies deviations in clone insert size that indicate intra-chromosomal structural variations compared to a reference genome. | InDel discovery Sequencing | Mapping | | Perl | GPL | Linux 64 |
| AB Small Indel Tool | The SOLiD™ Small Indel Tool processes the indel evidences found in the pairing step of the SOLiD™ System Analysis Pipeline Tool (Corona Lite). | InDel discovery Sequencing | Mapping Alignment | | Perl C++ | GPL | Linux 64 |
| ABBA | Assembly Boosted By Amino acid sequence is a comparative gene assembler, which uses amino acid sequences from predicted proteins to help build a better assembly | Genomic Assembly | Assembly Scaffolding | | | Artistic License | Linux |
| ABMapper | Maps RNA-Seq reads to target genome considering possible multiple mapping locations and splice junctions | Genomics Transcriptomics | Mapping Alignment | | C++ Perl | GPLv3 | Linux |
| ABySS | ABySS is a de novo sequence assembler designed for short reads and large genomes. | De-novo assembly | Assembly De Bruijn graph | MPI OpenMP | C++ | Free for academic use | POSIX Linux Mac OS X |
| Adaptor Removal | Removes adaptor fragments from raw short read | General | Adaptor Removal | Trimming | Java | Custom licence | Linux R4 |

What can we do with all
this sequence information?

Population Scale Analysis

We can now begin to assess genetic differences on a very large scale, both as naturally occurring variation in human and non-human populations as well somatically within tumors



<https://www.genomicsengland.co.uk/the-100000-genomes-project/>

“Variety’s the very spice of life”

-William Cowper, 1785

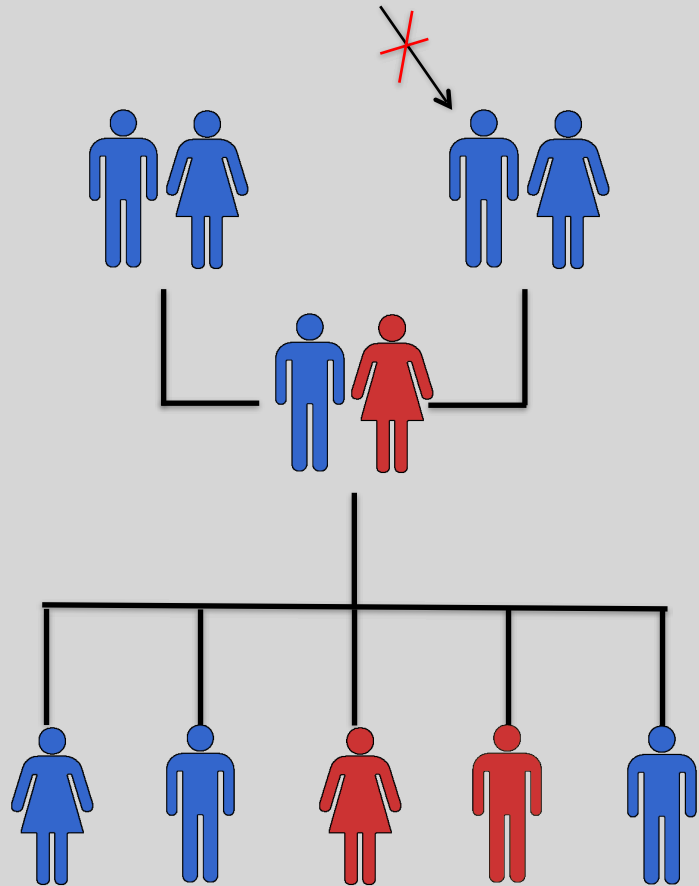
“Variation is the spice of life”

-Kruglyak & Nickerson, 2001

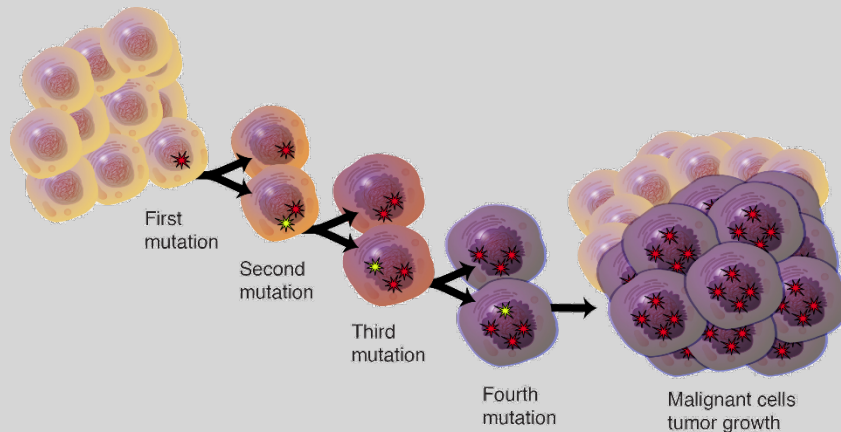
- While the sequencing of the human genome was a great milestone, the DNA from a single person is not representative of the millions of potential differences that can occur between individuals
- These unknown genetic variants could be the cause of many phenotypes such as differing morphology, susceptibility to disease, or be completely benign.

Germline Variation

- Mutations in the germline are passed along to offspring and are present in the DNA over every cell
- In animals, these typically occur in meiosis during gamete differentiation



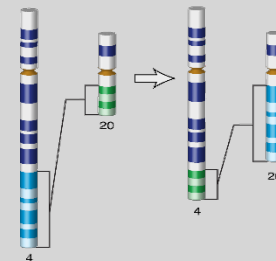
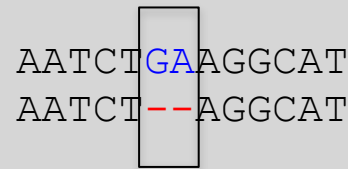
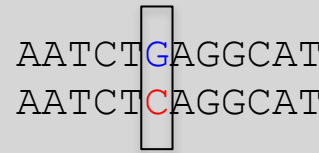
Somatic Variation



- Mutations in non-germline cells that are not passed along to offspring
- Can occur during mitosis or from the environment itself
- Are an integral part in tumor progression and evolution

Types of Genomic Variation

- **Single Nucleotide Polymorphisms (SNPs)** - mutations of one nucleotide to another
- **Insertion/Deletion Polymorphisms (INDELs)** - small mutations removing or adding one or more nucleotides at a particular locus
- **Structural Variation (SVs)** - medium to large sized rearrangements of chromosomal DNA



Differences Between Individuals

The average number of genetic differences in the germline between two random humans can be broken down as follows:

- 3,600,000 single nucleotide differences
- 344,000 small insertion and deletions
- 1,000 larger deletion and duplications

Numbers change depending on ancestry!

Discovering Variation: SNPs and INDELS

SNP

sequencing error or genetic variant?

ATCCTGATTCGGTGAACGTTATCGACGATCCGATCGA
ATCCTGATTCGGTGAACGTTATCGACGATCCGATCGA
CGGTGAACGTTATCGACGATCCGATCGAACTGTCAGC
GGTGAACGTTATCGACGTTCCGATCGAACTGTCAGCG
TGAACGTTATCGACGTTCCGATCGAACTGTCATCGGC
TGAACGTTATCGACGTTCCGATCGAACTGTCAGCGGC
TGAACGTTATCGACGTTCCGATCGAACTGTCAGCGGC
GTATCGACGATCCGATCGAACTGTCAGCGGAAGCT
TTATCGACGATCCGATCGAACTGTCAGCGGAAGCT

ATCCTGATTCGGTGAACGTTATCGACGATCCGATCGAACTGTCAGCGGAAGCTGATCGATCGATCGATGCTAGTG

reference genome

sequencing error or genetic variant?

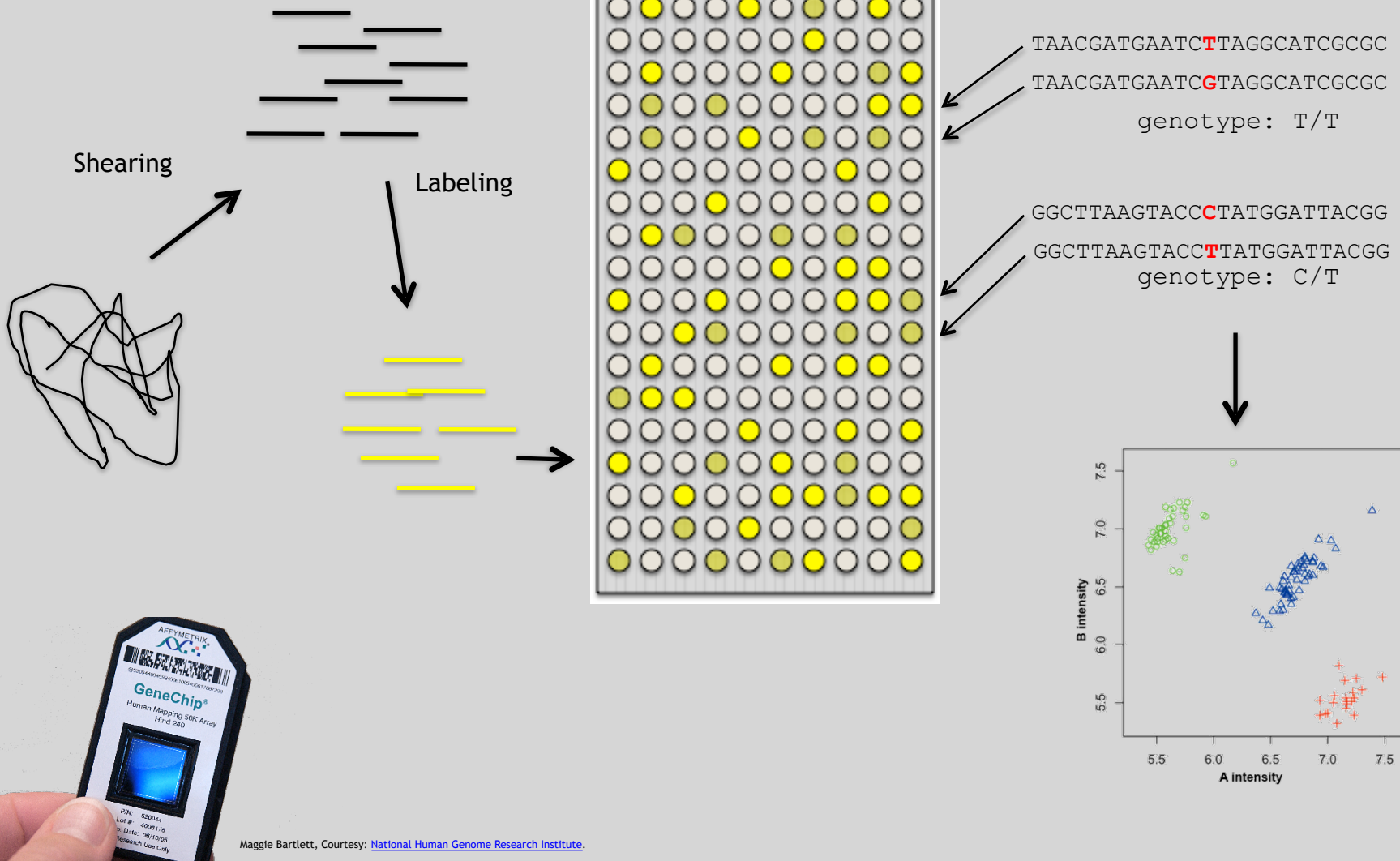
TTATCGACGATCCGATCGAACTGTCAGCGGAAGCT
TCGACGATCCGATCGAACTGTCAGCGGAAGCTGATCG
ATCCGATCGAACTGTCAGCGGAAGCTGATCGCGATCG
TCCGAGCGAACTGTCAGCGGAAGCTGATCGCGATCG
TCCGATCGAACTGTCAGCGGAAGCTGATCGATCGA
GATCGAACTGTCAGCGGAAGCTGATCGCGATCGA
AACTGTCAGCGGAAGCTGATCGCGATCGATGCTA
TGTCAAGCGGAAGCTGATCGATCGATCGATGCTAG
TCAGCGGAAGCTGATCGATCGATCGATGCTAGTG

INDEL

Genotyping Small Variants

- Once discovered, oligonucleotide probes can be generated with each individual allele of a variant of interest
- A large number can then be assessed simultaneously on microarrays to detect which combination of alleles is present in a sample

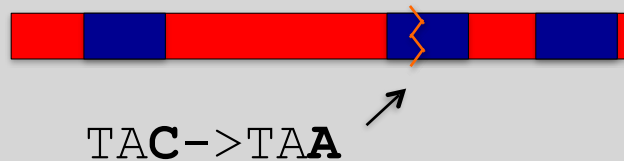
SNP Microarrays



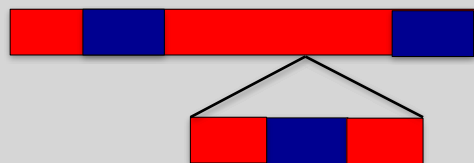
Impact of Genetic Variation

There are numerous ways genetic variation can exhibit functional effects

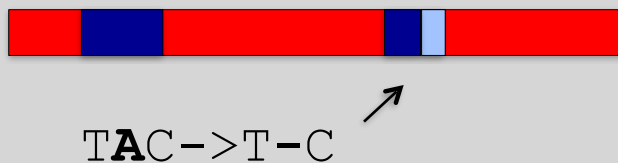
Premature stop codons



Gene or exon deletion

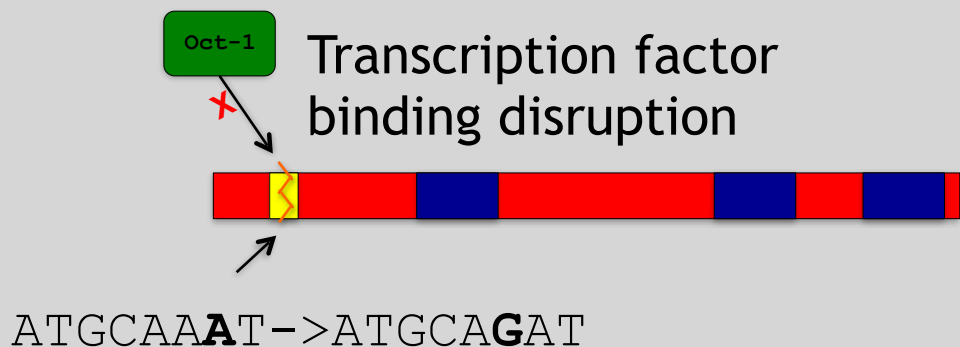


Frameshift mutation



Oct-1

Transcription factor binding disruption



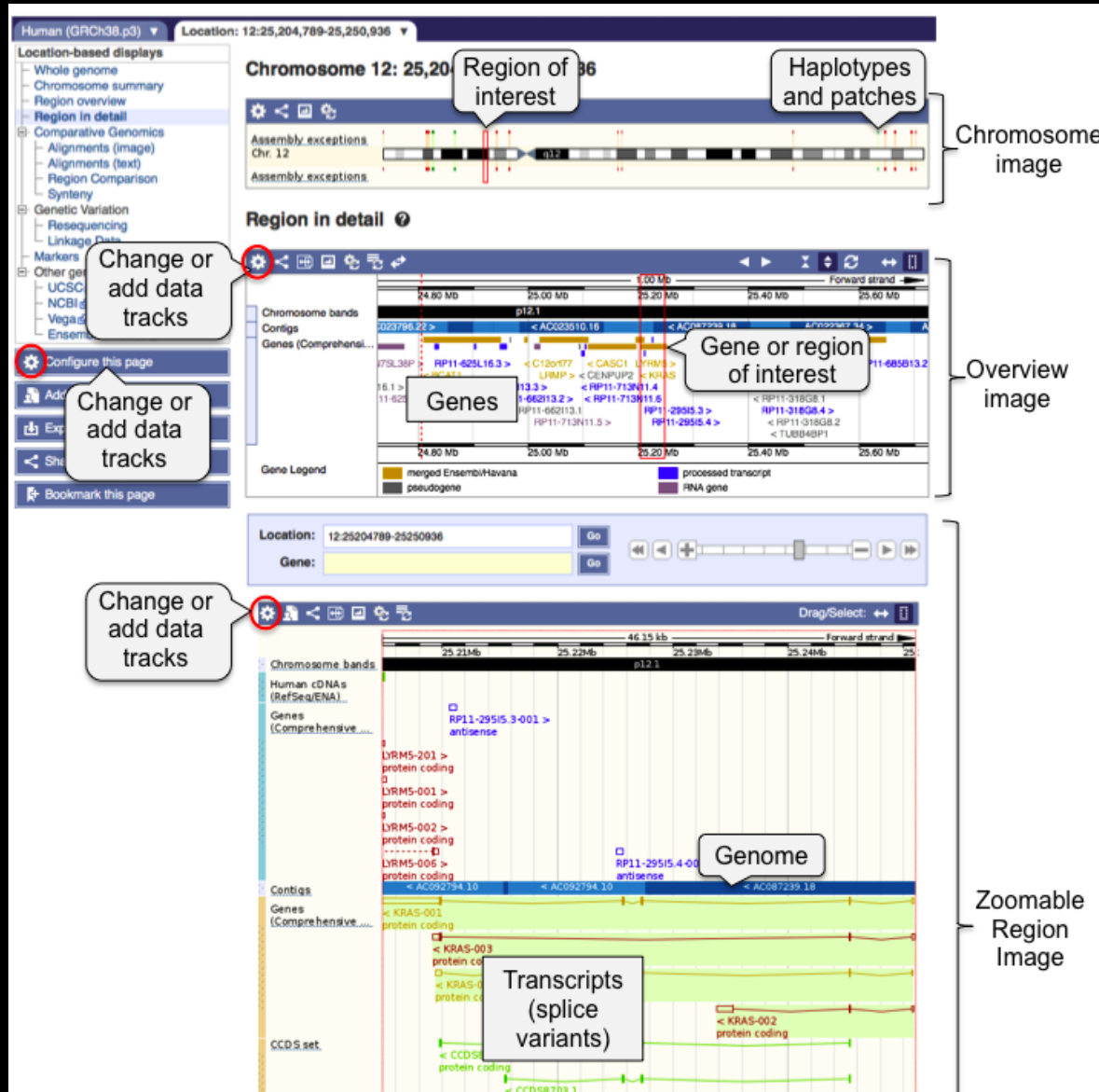
Hand-on time!

https://bioboot.github.io/bimm143_W19/lectures/#14

Sections **1** to **3** please (up to running Read Alignment)

See IP address on website for **your** Galaxy server

<http://uswest.ensembl.org/Help/View?id=140>



Access a jetstream galaxy instance!

Use assigned IP address

Do it Yourself!

The screenshot shows a Galaxy web interface running on port 149.165.169.186. The main content area displays the 'Bowtie2 - map reads against reference genome' tool configuration. The left sidebar contains a list of available tools categorized under 'Collection Operations', 'Text Manipulation', 'Filter and Sort', 'Join, Subtract and Group', 'Convert Formats', 'Extract Features', 'Fetch Sequences', 'Fetch Alignments', 'Statistics', 'Graph/Display Data', 'FASTA manipulation', 'NGS: QC and manipulation', 'NGS: DeepTools', 'NGS: Mapping', 'Lastz map short reads against reference sequence', 'Map with Bowtie for Illumina', 'Map with BWA for Illumina', 'Map with BWA for SOLID', 'Megablast compare short reads against htgs, nt, and wgs databases', 'Parse blast XML output', 'Map with BWA-MEM - map medium and long reads (> 100 bp) against reference genome', 'Map with BWA - map short reads (< 100 bp) against reference genome', 'Bowtie2 - map reads against reference genome', and 'NGS: RNA Analysis'. The central panel shows the specific parameters for the Bowtie2 tool, including FASTQ file selection, write options, reference genome selection, and analysis mode. The right panel shows a history of completed jobs, such as 'htseq-count' and 'Cufflinks' runs, along with their details and preview links.

Raw data usually in FASTQ format

```
@NS500177:196:HFTTTAFXX:1:11101:10916:1458 2:N:0:CGCGGCTG  
ACACGACGATGAGGTGACAGTCACGGAGGATAAGATCAATGCCCTCATTAAAGCAGCCGGTGTAA  
+  
AAAAAAEEEEEEEEEEE//AEEEAEeeeeeee/EE/<<EE/AEAAEE//EEEAEAAEA<
```

1

2

3

4

Each sequencing “read” consists of 4 lines of data :

- 1 The first line (which always starts with ‘@’) is a unique ID for the sequence that follows
- 2 The second line contains the bases called for the sequenced fragment
- 3 The third line is always a “+” character
- 4 The forth line contains the quality scores for each base in the sequenced fragment (these are ASCII encoded...)

ASCII Encoded Base Qualities

```
@NS500177:196:HFTTTAFXX:1:11101:10916:1458 2:N:0:CGCGGCTG  
ACACGACGATGAGGTGACAGTCACGGAGGATAAGATCAATGCCCTCATTAAAGCAGCCGGTGTAA  
+  
AAAAAAEEEEEEEEEEE//AEEEAEeeeeeee/EE/<<EE/AEAAEE//EEEAEAAEA< 4
```

- Each sequence base has a corresponding numeric quality score encoded by a single ASCII character typically on the 4th line (see ④ above)
- ASCII characters represent integers between 0 and 127
- Printable ASCII characters range from 33 to 126
- Unfortunately there are 3 quality score formats that you may come across...

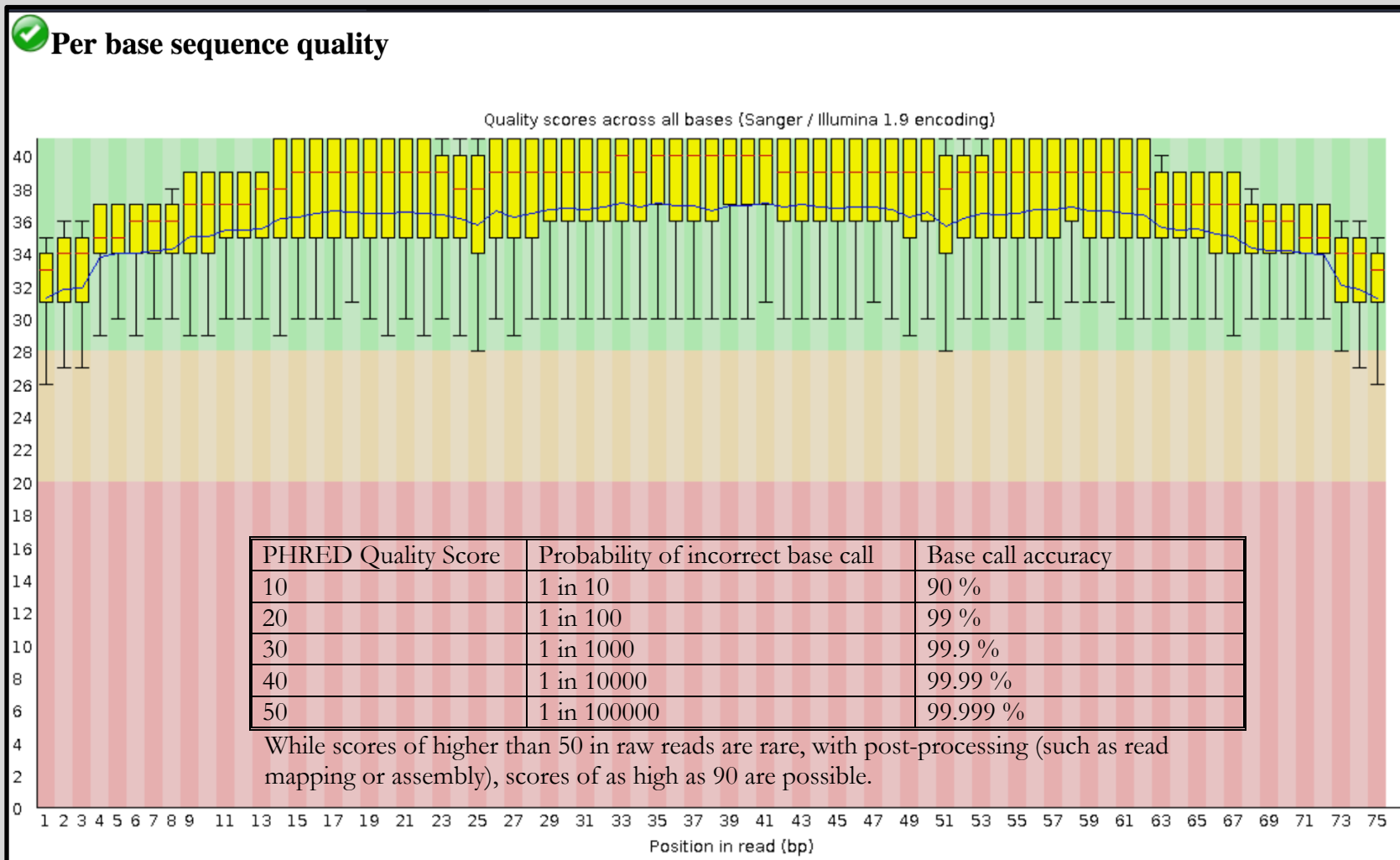
Interpreting Base Qualities in R

| | | ASCII Range | Offset | Score Range |
|---------------------------------|---------------|-------------|--------|-------------|
| Sanger, Illumina (Ver > 1.8) | fastqsanger | 33-126 | 33 | 0-93 |
| Solexa, Illumina (Ver < 1.3) | fastqsolexa | 59-126 | 64 | 5-62 |
| Illumina (Ver 1.3 -1.7) | fastqillumina | 64-126 | 64 | 0-62 |

```
> library(seqinr)
> library(gtools)
> phred <- asc( s2c("DDDDCDEDCCDDDBBDDCC@") ) - 33
> phred
## D D D D C D E D C D D D D B B D D D C C @
## 35 35 35 35 34 35 36 35 34 35 35 35 35 33 33 35 35 35 34 34 31

> prob <- 10**(-phred/10)
```

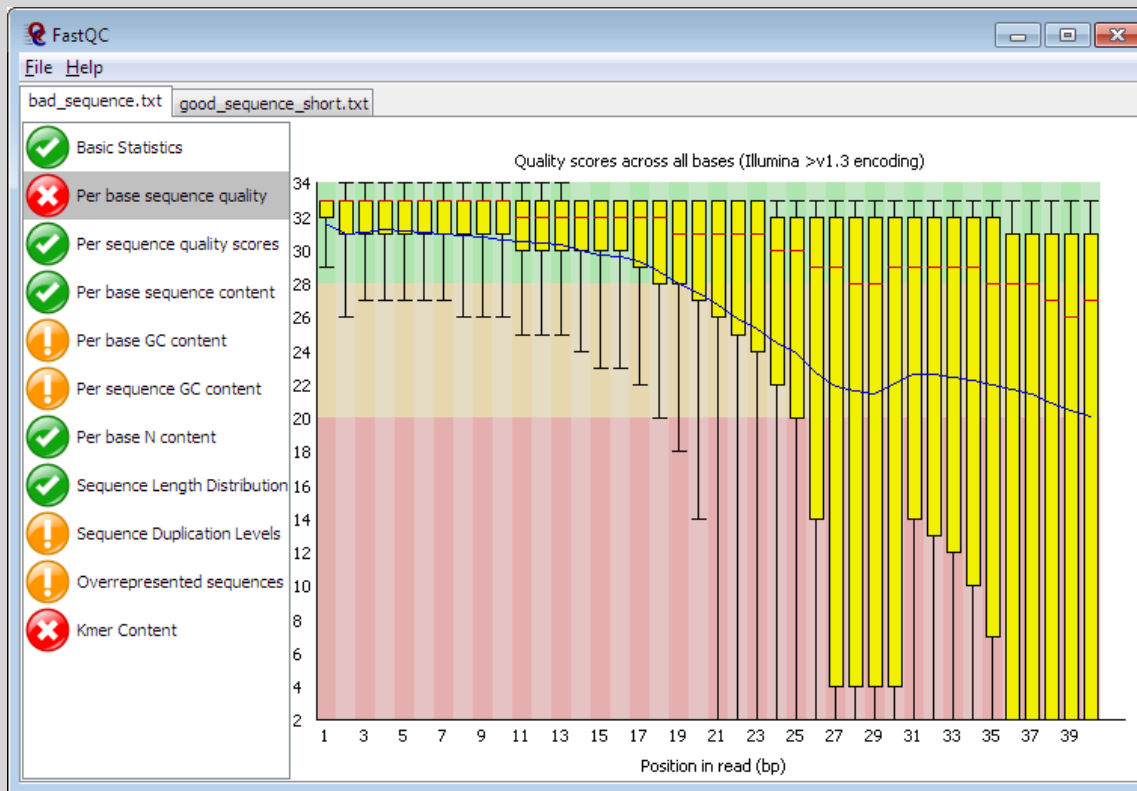
FastQC Report



FASTQC

FASTQC is one approach which provides a visual interpretation of the raw sequence reads

- <http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>



Sequence Alignment

- Once sequence quality has been assessed, the next step is to align the sequence to a reference genome
- There are *many* distinct tools for doing this; which one you choose is often a reflection of your specific experiment and personal preference

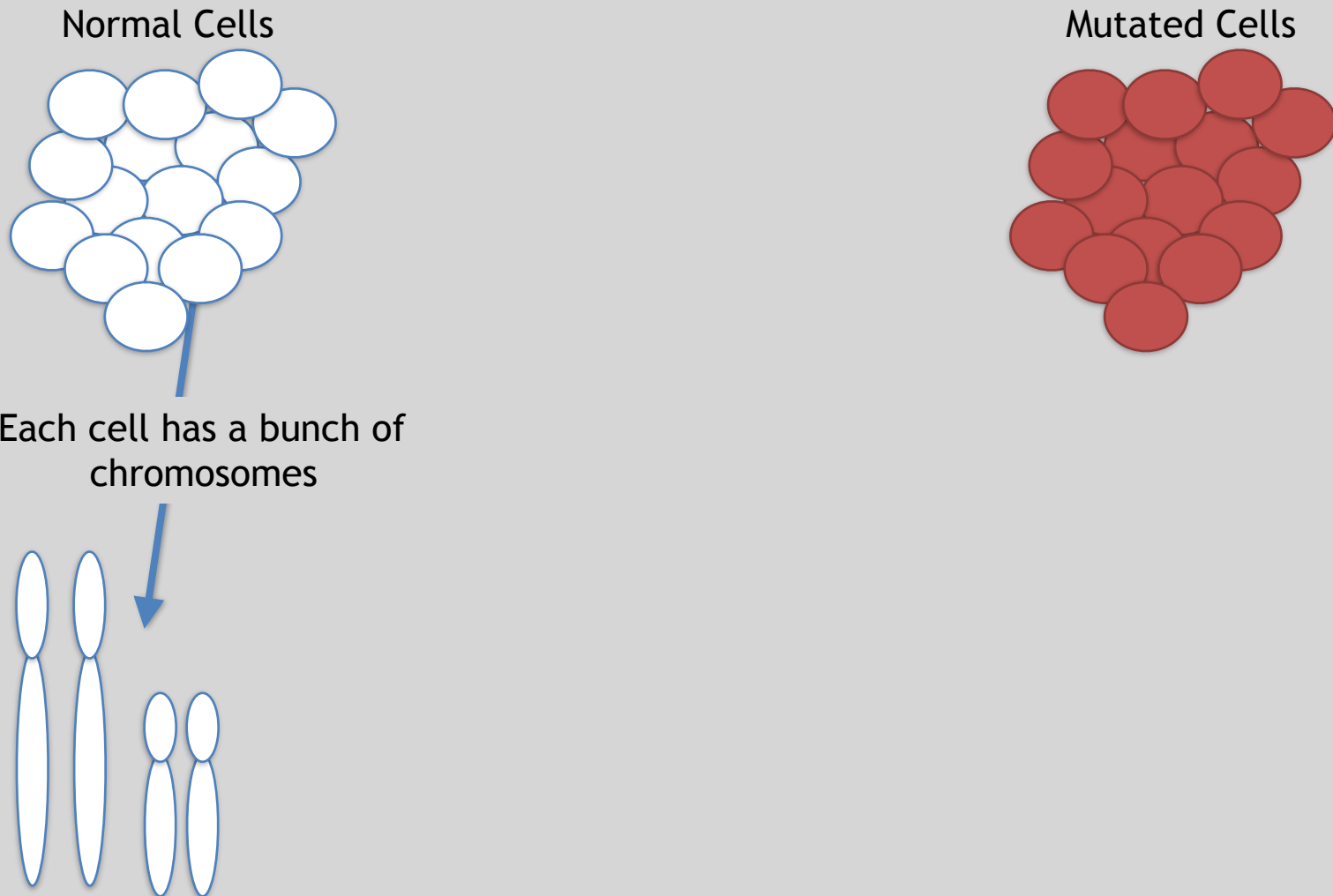
| | | |
|------------|-----------|-------|
| BWA | BarraCUDA | RMAP |
| Bowtie | CASHx | SSAHA |
| SOAP2 | GSNAP | etc |
| Novoalign | Mosiak | |
| mr/mrsFast | Stampy | |
| Eland | SHRiMP | |
| Blat | SeqMap | |
| Bfast | SLIDER | |

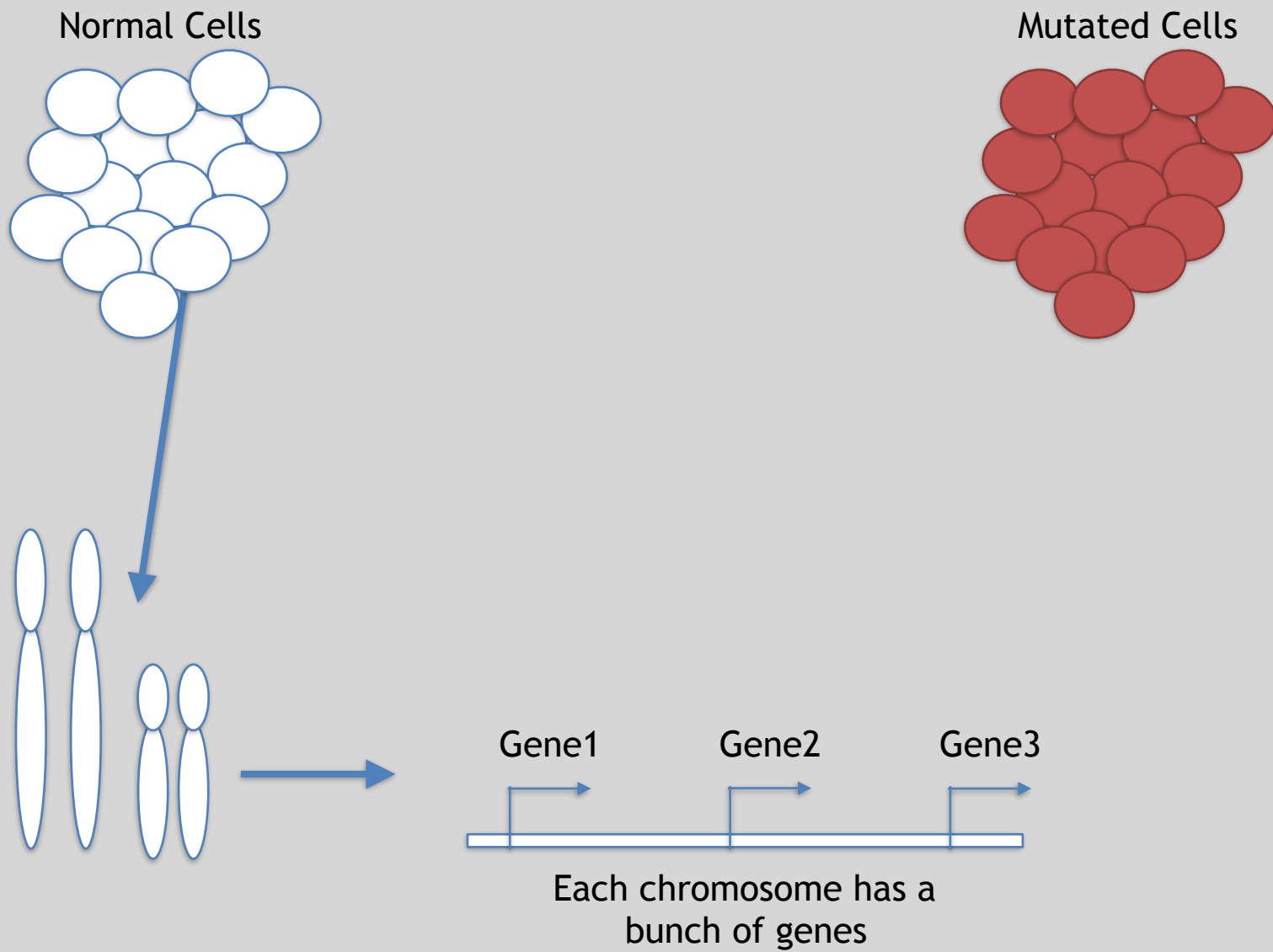
RNA Sequencing

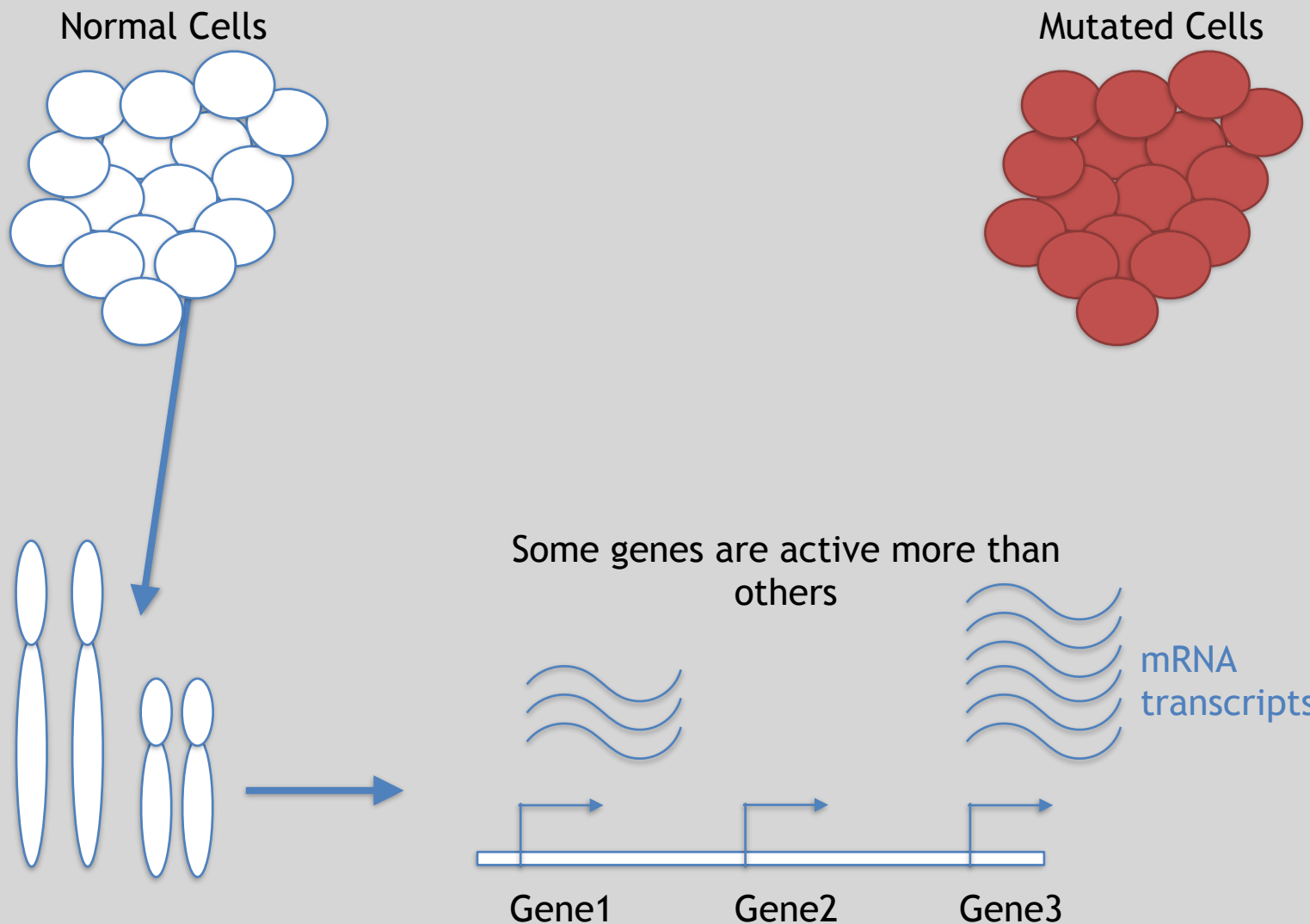
The absolute basics

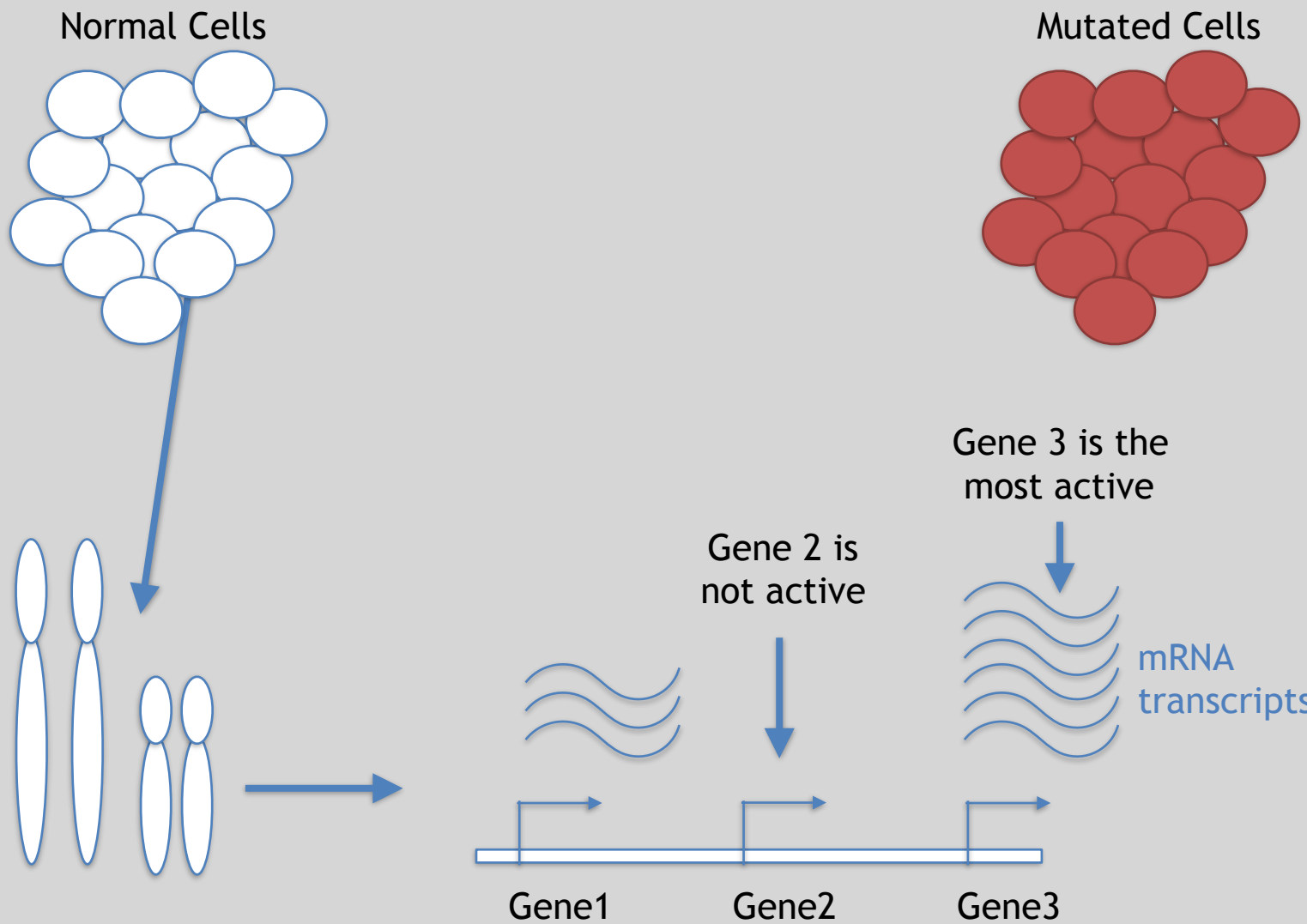


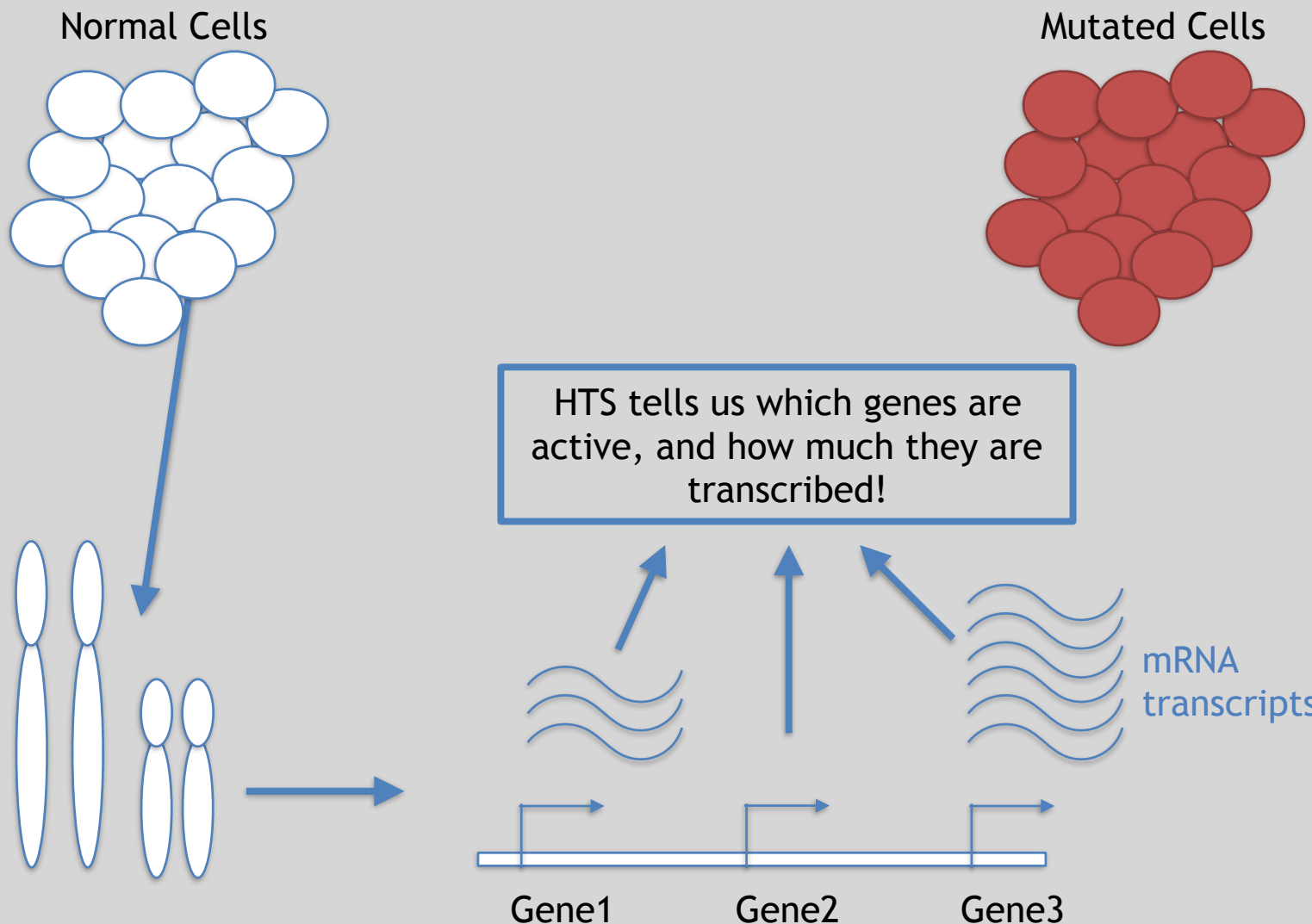
- The **mutated cells** behave differently than the **normal cells**
- We want to know what genetic mechanism is causing the difference
- One way to address this is to examine differences in gene expression via RNA sequencing...

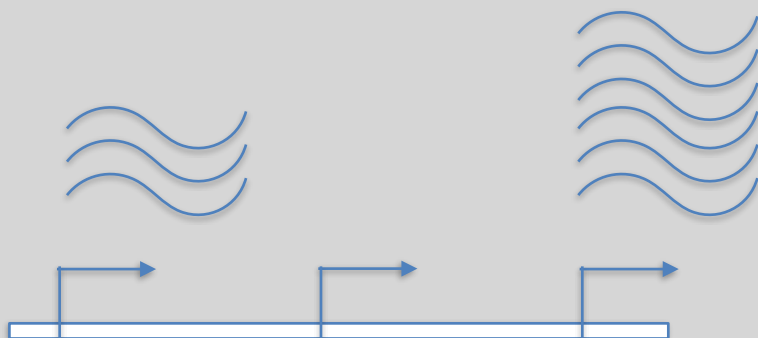
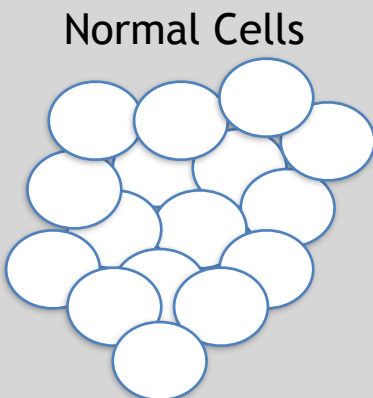




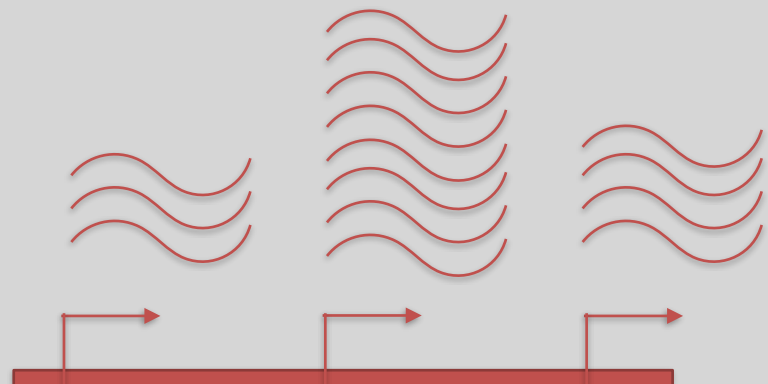
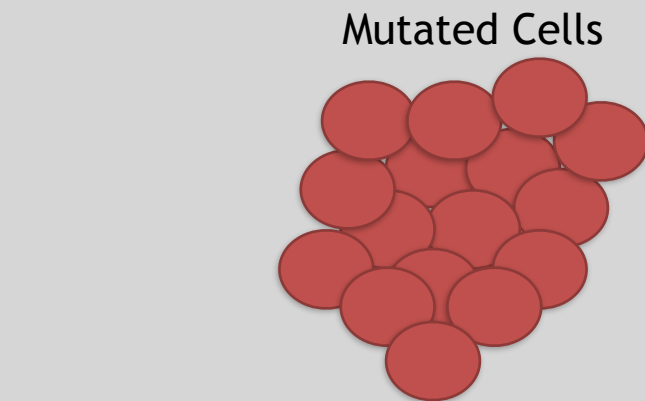




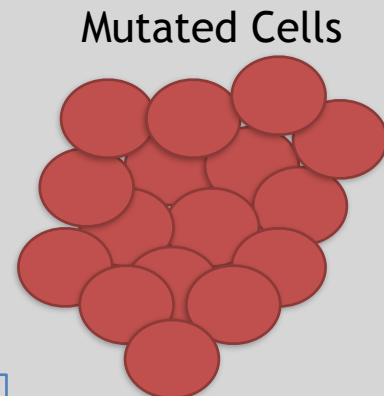
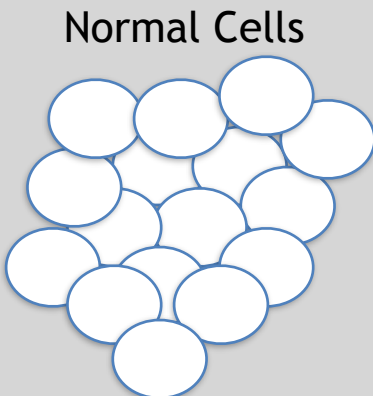




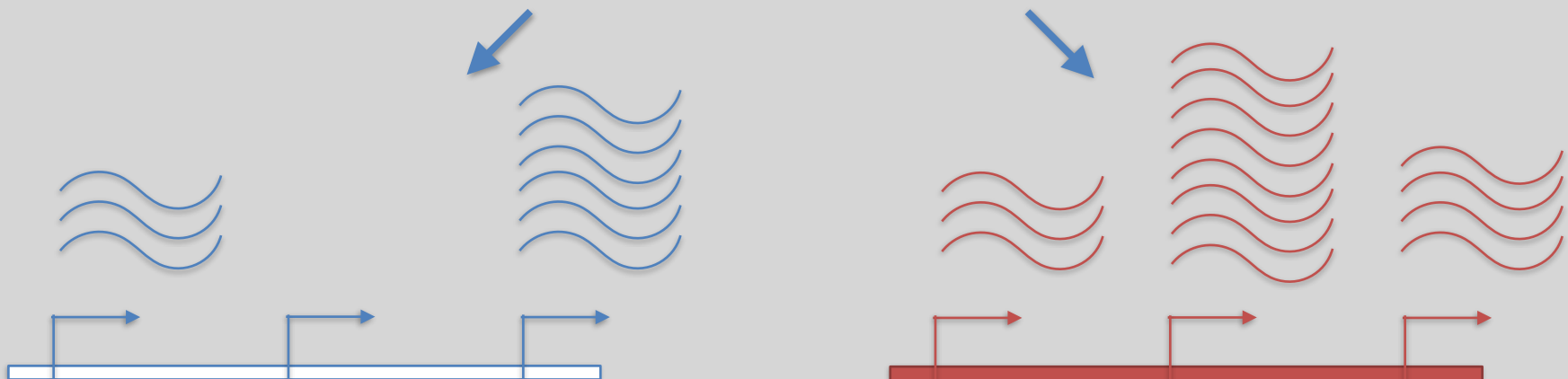
We use RNA-Seq to measure gene expression in normal cells ...

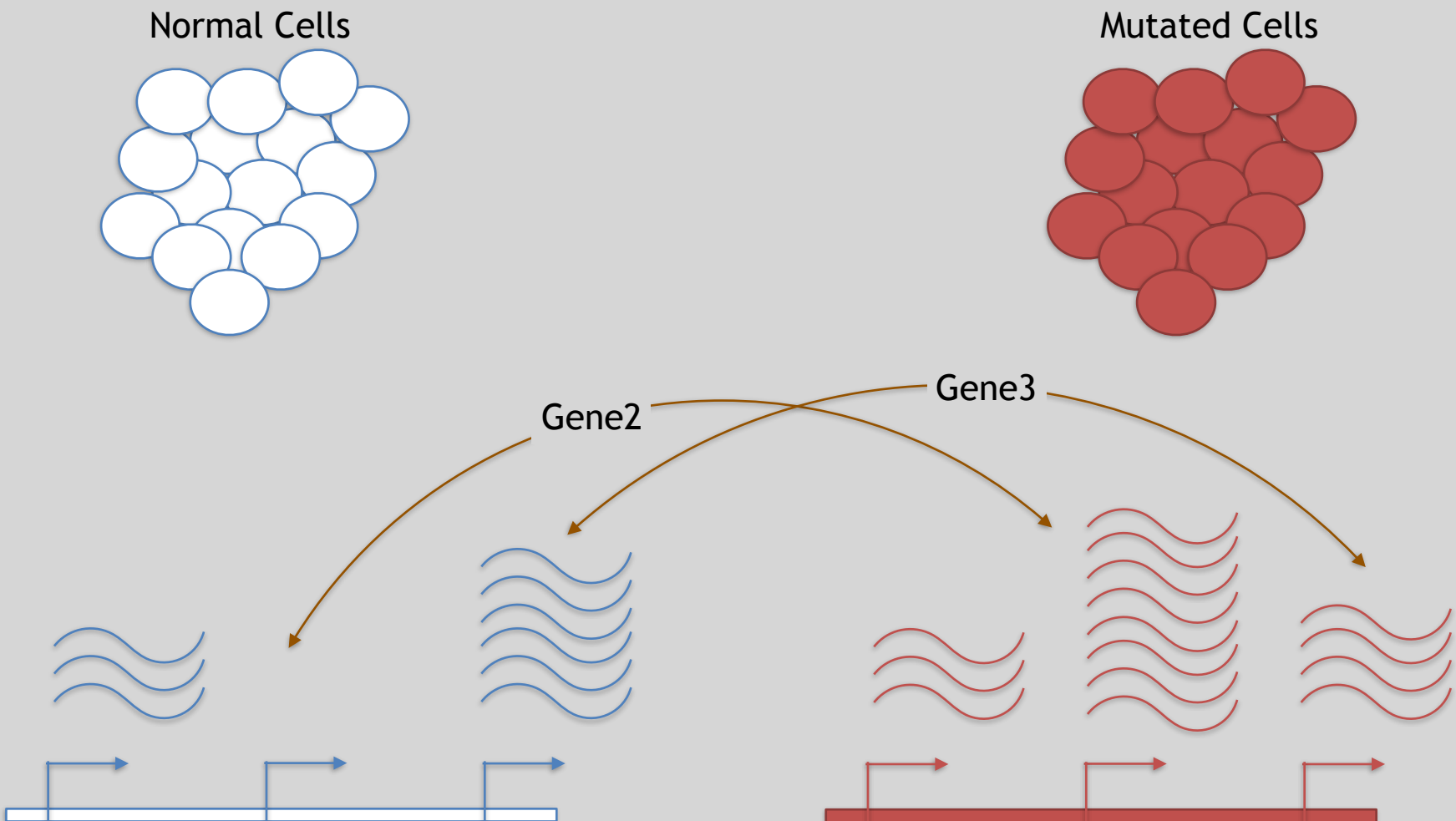


... then use it to measure gene expression in mutated cells



Then we can compare the two cell types to figure out what is different in the mutated cells!





3 Main Steps for RNA-Seq:

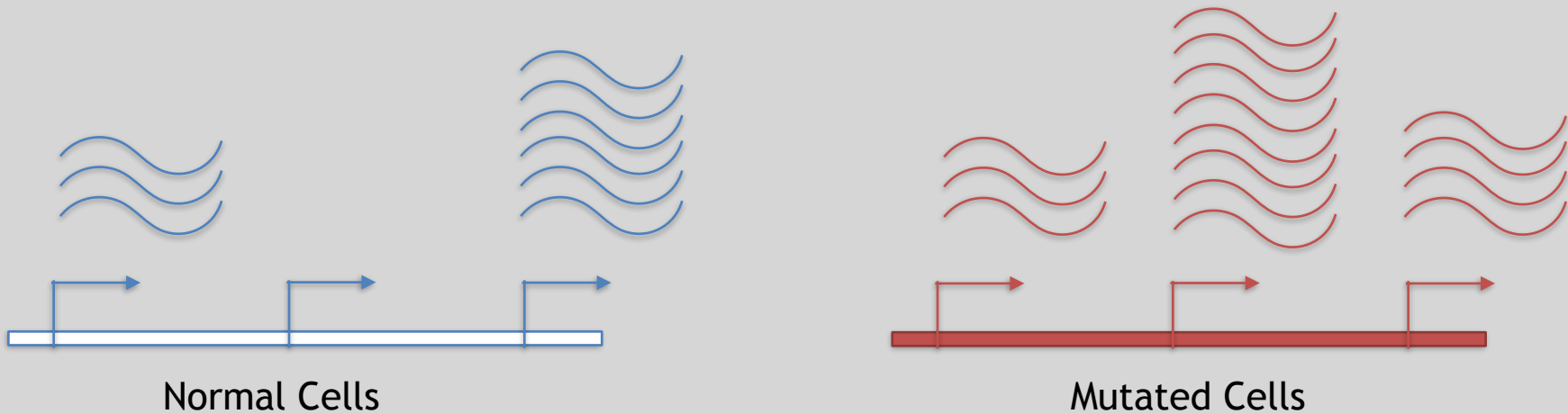
- 1) Prepare a sequencing library**
(RNA to cDNA conversion via reverse transcription)
- 2) Sequence**
(Using the same technologies as DNA sequencing)
- 3) Data analysis**
(Often the major bottleneck to overall success!)

We will discuss each of these steps in detail
(particularly the 3rd) next day!

Today we will get to the start of step 3!

| Gene | WT-1 | WT-2 | WT-3 | ... |
|------|------|------|------|-----|
| A1BG | 30 | 5 | 13 | ... |
| AS1 | 24 | 10 | 18 | ... |
| ... | ... | ... | ... | ... |

We sequenced, aligned, counted the reads per gene in each sample to arrive at our data matrix



Hand-on time!

https://bioboot.github.io/bimm143_W19/lectures/#14

Focus on **Sections 4** please
(After your Alignment is finished)

Feedback:
[Muddy Point Assessment]

Additional Reference Slides

on SAM/BAM Format and
Sequencing Methods

Sequence Alignment

- Once sequence quality has been assessed, the next step is to align the sequence to a reference genome
- There are *many* distinct tools for doing this; which one you choose is often a reflection of your specific experiment and personal preference

| | | |
|------------|-----------|-------|
| BWA | BarraCUDA | RMAP |
| Bowtie | CASHx | SSAHA |
| SOAP2 | GSNAP | etc |
| Novoalign | Mosiak | |
| mr/mrsFast | Stampy | |
| Eland | SHRiMP | |
| Blat | SeqMap | |
| Bfast | SLIDER | |

SAM Format

- Sequence Alignment/Map (SAM) format is the almost-universal sequence alignment format for NGS
 - binary version is BAM
- It consists of a header section (lines start with '@') and an alignment section
- The official specification can be found here:
 - <http://samtools.sourceforge.net/SAM1.pdf>

Example SAM File

- Because SAM files are plain text (unlike their binary counterpart, BAM), we can take a peek at a few lines of the header with head, See:

https://bioboot.github.io/bimm143_F18/class-material/sam_format/

Header section

```
@HD VN:1.0 SO:coordinate
@SQ SN:1 LN:249250621 AS:Ncbi37 UR:file:/data/local/ref/GATK/human_glk_v37.fasta M5:1b22b98cdeb4a9304cb5d48026a85128
@SQ SN:2 LN:243199373 AS:Ncbi37 UR:file:/data/local/ref/GATK/human_glk_v37.fasta M5:a0d9851da00400dec1098a9255ac712e
@SQ SN:3 LN:198022430 AS:Ncbi37 UR:file:/data/local/ref/GATK/human_glk_v37.fasta M5:fd1d811849cc2fadecb929bb925902e5
@RG ID:UM0098:1 PL:ILLUMINA PU:HWUSI-EAS1707-615LHAXXX-L001 LB:80 DT:2010-05-05T20:00:00-0400 SM:SD37743 CN:UMCORE
@RG ID:UM0098:2 PL:ILLUMINA PU:HWUSI-EAS1707-615LHAXXX-L002 LB:80 DT:2010-05-05T20:00:00-0400 SM:SD37743 CN:UMCORE
@PG ID:bw VN:0.5.4
```

Alignment section

```
1:497:R:-272+13M17D24M 113 1 497 37 37M 15 100338662 0
CGGTCTGACCTGAGGAGAACACTGTGCTCCGCCCTTCAG 0;====9,>>>>=>>>>>>=>>>>>>> XT:A:U NM:i:0 SM:i:37 AM:i:0 X0:i:1 X1:i:0
XM:i:0 XO:i:0 XG:i:0 MD:Z:37
19:20389:F:275+18M2D19M 99 1 17644 0 37M = 17919 314
TATGACTGCTAATAATACCTACACATGTTAGAACCAT >>>>>>>>>>>>>>>><>><>>4:>>:<9 RG:Z:UM0098:1 XT:A:R NM:i:0 SM:i:0 AM:i:0 X0:i:0 X0:i:4
X1:i:0 XM:i:0 XG:i:0 MD:Z:37
19:20389:F:275+18M2D19M 147 1 17919 0 18M2D19M = 17644 -314
GTAGTACCAACTGTAAGTCCTTATCTTCATACTTGT ;44999;499<8<8<<8<<<<<<<7<,<<<><< XT:A:R NM:i:2 SM:i:0 AM:i:0 X0:i:4 X1:i:0
XM:i:0 XO:i:1 XG:i:2 MD:Z:18^CA19
9:21597+10M2I25M:R:-209 83 1 21678 0 8M2I27M = 21469 -244
CACCACATACATACCAAGCCTGGCTGTCTTCT <,<9<<5><<<><<><>><>><>>><> XT:A:R NM:i:2 SM:i:0 AM:i:0 X0:i:5 X1:i:0
XM:i:0 XO:i:1 XG:i:2 MD:Z:35
```

SAM header section

- Header lines contain vital metadata about the reference sequences, read and sample information, and (optionally) processing steps and comments.
- Each header line begins with an @, followed by a two-letter code that distinguishes the different type of metadata records in the header.
- Following this two-letter code are tab-delimited key-value pairs in the format **KEY:VALUE** (the SAM format specification names these tags and values).

https://bioboot.github.io/bimm143_F18/class-material/sam_format/

SAM Utilities

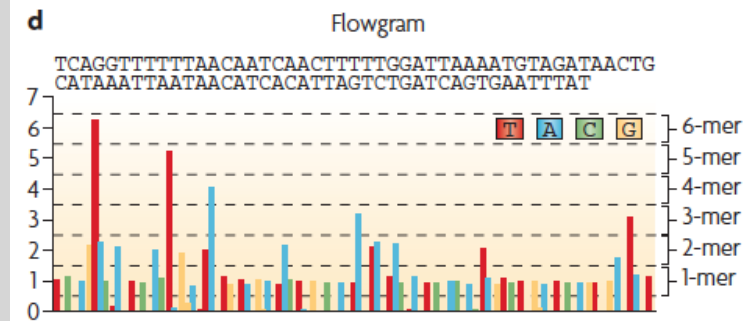
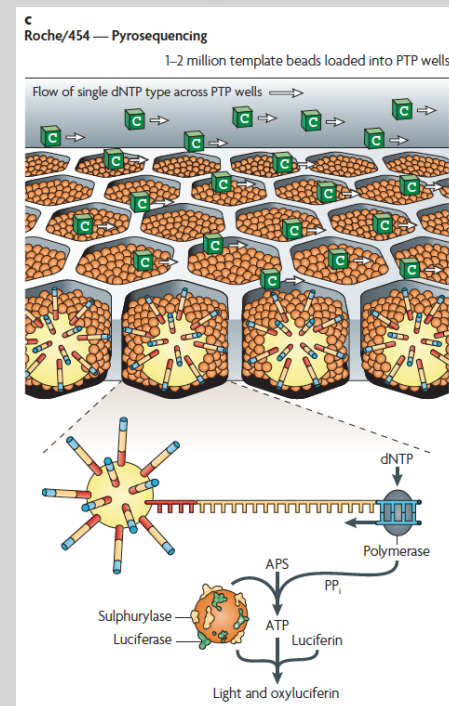
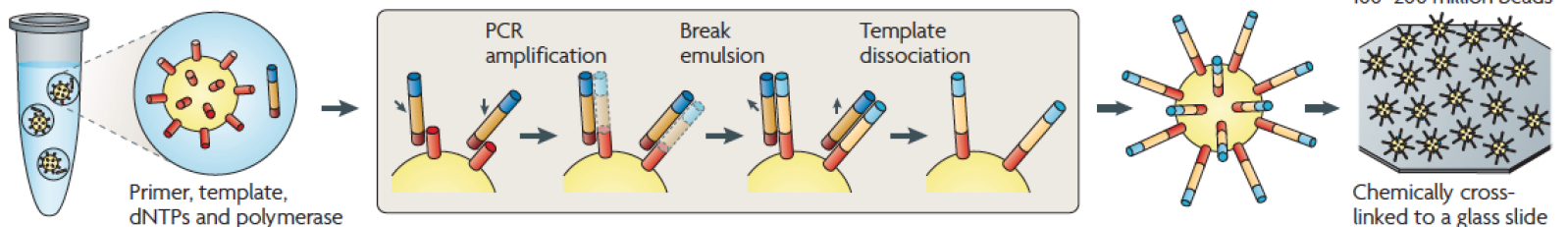
- **Samtools** is a common toolkit for analyzing and manipulating files in SAM/BAM format
 - <http://samtools.sourceforge.net/>
- **Picard** is another set of utilities that can used to manipulate and modify SAM files
 - <http://picard.sourceforge.net/>
- These can be used for viewing, parsing, sorting, and filtering SAM files as well as adding new information (e.g. Read Groups)

Additional Reference Slides on Sequencing Methods

Roche 454 - Pyrosequencing

a Roche/454, Life/APG, Polonator Emulsion PCR

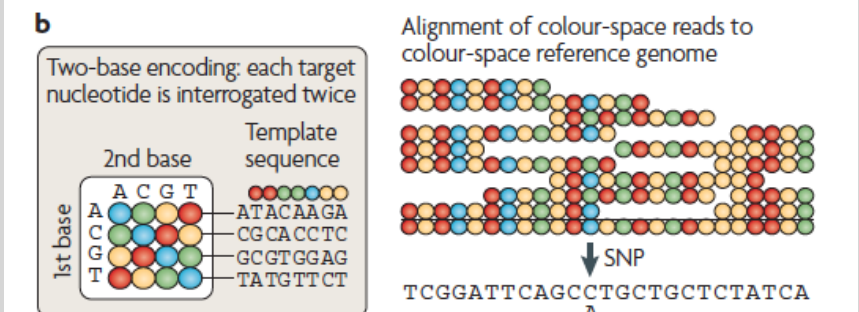
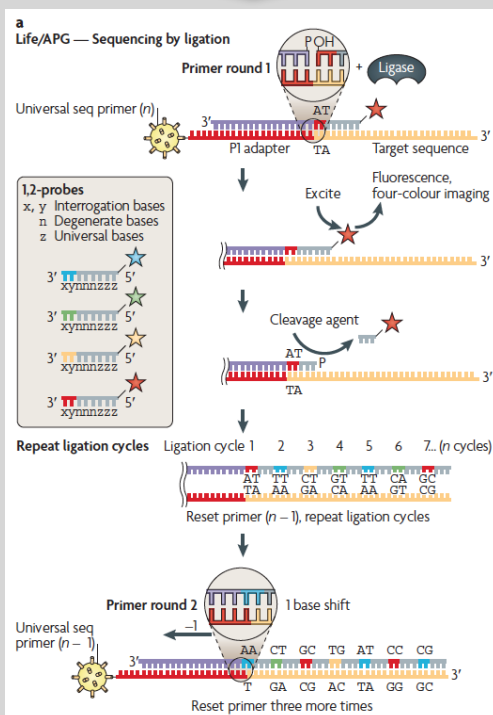
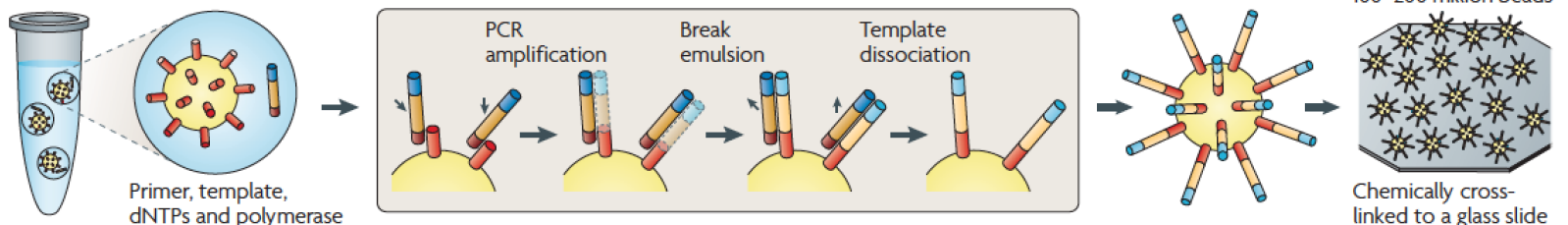
One DNA molecule per bead. Clonal amplification to thousands of copies occurs in microreactors in an emulsion



Life Technologies SOLiD - Sequence by Ligation

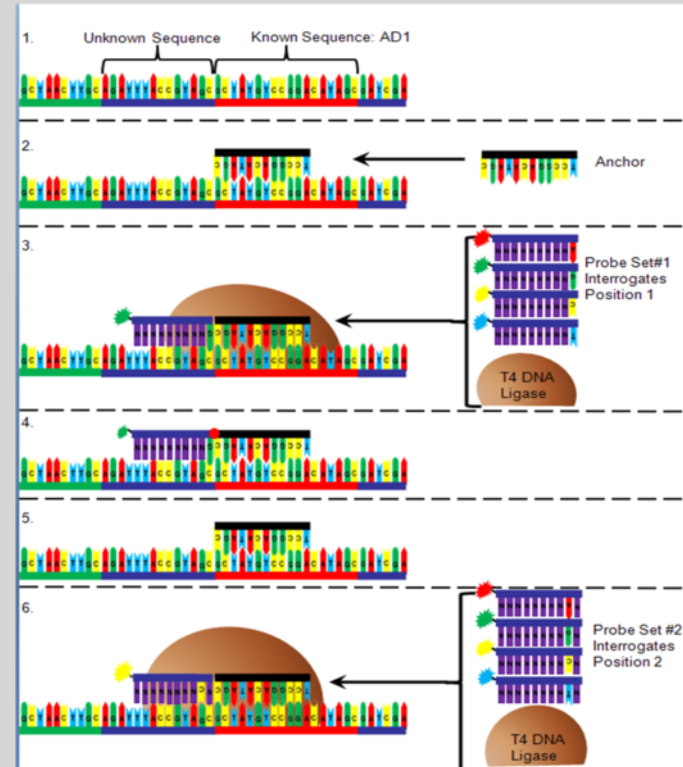
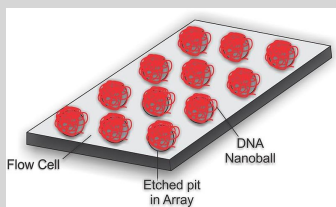
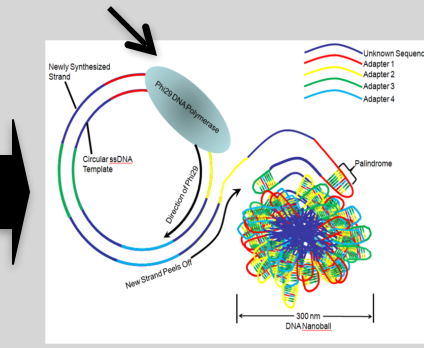
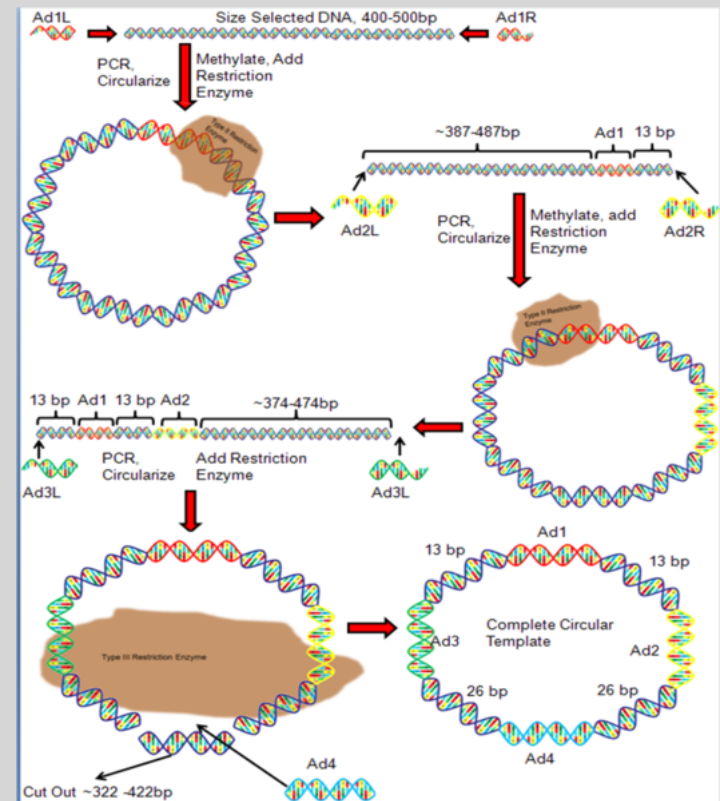
a Roche/454, Life/APG, Polonator Emulsion PCR

One DNA molecule per bead. Clonal amplification to thousands of copies occurs in microreactors in an emulsion



Complete Genomics - Nanoball Sequencing

Has proofreading ability!



“Benchtop” Sequencers

- Lower cost, lower throughput alternative for smaller scale projects
- Currently three significant platforms
 - Roche 454 GS Junior
 - Life Technology Ion Torrent
 - Personal Genome Machine (PGM)
 - Proton
 - Illumina MiSeq

| Platform | List price | Approximate cost per run | Minimum throughput (read length) | Run time | Cost/Mb | Mb/h |
|-------------------------------|-------------------------|--------------------------|----------------------------------|----------|---------|-------|
| 454 GS Junior | \$108,000 | \$1,100 | 35 Mb (400 bases) | 8 h | \$31 | 4.4 |
| Ion Torrent PGM (314 chip) | \$80,490 ^{a,b} | \$225 ^c | 10 Mb (100 bases) | 3 h | \$22.5 | 3.3 |
| (316 chip) | | \$425 | 100 Mb ^d (100 bases) | 3 h | \$4.25 | 33.3 |
| (318 chip) | | \$625 | 1,000 Mb (100 bases) | 3 h | \$0.63 | 333.3 |
| MiSeq | \$125,000 | \$750 | 1,500 Mb (2 × 150 bases) | 27 h | \$0.5 | 55.5 |

Loman, NJ (2012), *Nat. Biotech.*, 5, pp. 434-439

PGM - Ion Semiconductor Sequencing

