### **Bioinformatics Lab**

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### **Objectives**

- Hands on introduction to bioinformatics programming
- Review basic biological/computational aspects
  - 1. basics of molecular biology
  - 2. basics of sequencing
  - 3. basics bioinformatics problems
    - short sequences read alignment
    - gene expression quantification
    - single cell approaches
    - computational epigenetic



### **Objectives**

- Introduction to Bioinformatics Frameworks/Tools
  - 1. biological sequence data formats/handling
    - Biopython, Pysam, R/bioconductor
  - 2. bioinformatics tools
    - BWA (aligner), Seurat, Cell Ranger, ...



### **Grading/Online material**

#### **Evaluation:**

- 20% prototypes
- 60% final project
- 20% presentation

#### **Extra-work for media informatics:**

research report

#### References/Courses Online

http://costalab.org/teaching/bioinformatics-software-lab-2024/



## Introduction to Molecular Biology



#### **Understanding Live in a Molecular Level**

How is genetic information inherited?

How the genetic information influence cellular processes?

How genes work together to promote particular molecular functions?



#### **Genetic Information - DNA**

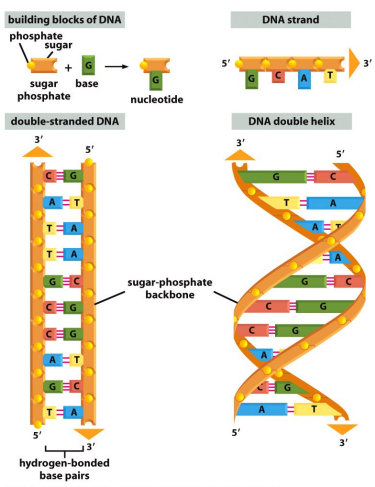
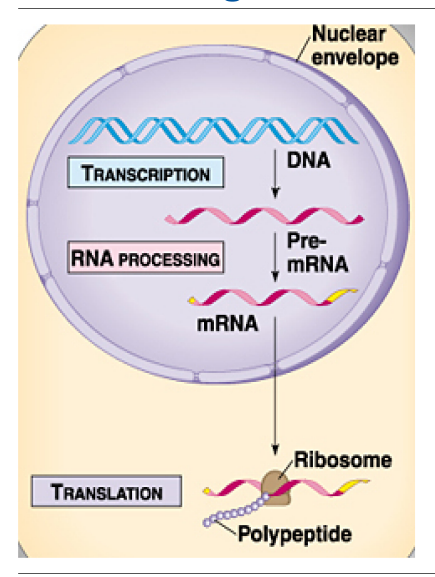


Figure 4-3 Molecular Biology of the Cell 5/e (© Garland Science 2008)

#### DNA (Deoxyribonucleic)

- chain of nucleic acids
- 4 bases: A;C;G;T
- forms DNA duplexes with paring A = T e C = G

#### **Central Dogma - Transcription**



#### **Transcription**

- DNA to RNA RNA (ribonucleic acid)
  - single stranded
  - 4 bases: A;C;G;U
  - unstable
  - transport of information from nucleus to cytoplasm



### **Central Dogma - Transcription**

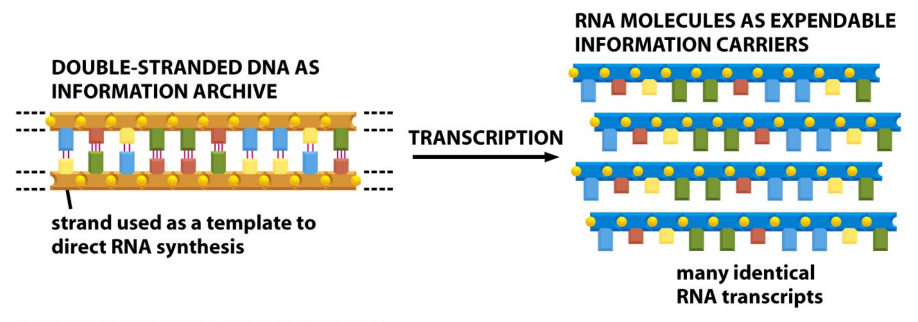
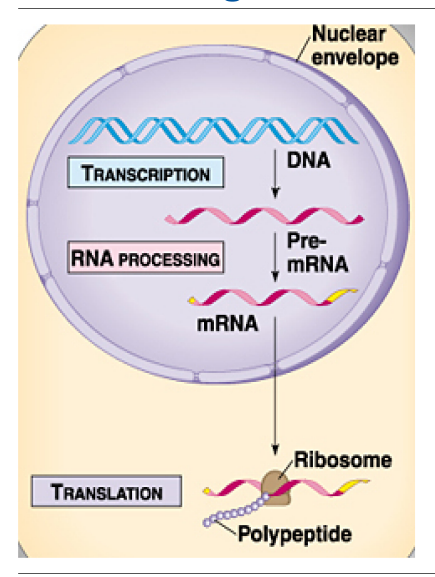


Figure 1-5 Molecular Biology of the Cell 5/e (© Garland Science 2008)

#### Transcription - copy of DNA information to RNA (T to U)



### **Central Dogma - Translation**



#### **Translation**

- RNA to Protein
- performed by the ribosome
- follows the genetic code

#### **Proteins**

- single stranded chain
- 20 amino acids
- assumes 3D structure
- main functional entities in the cell



#### **Genetic Code - Translation**

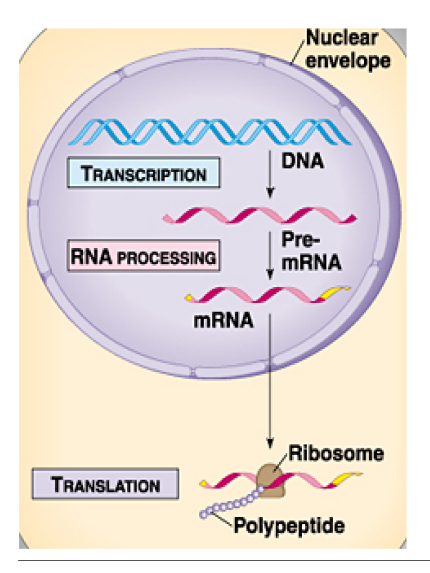
GCC GCG	AGA AGG CGA CGC CGG CGU								AUC		AAA	AUG		CCC	AGC AGU UCA UCC UCG UCU	ACC ACG	UGG		GUA GUC GUG GUU	UAA UAG UGA
Ala	Arg	Asp	Asn	Cys	Glu	Gln	Gly	His	lle	Leu	Lys	Met	Phe	Pro	Ser	Thr	Trp	Tyr	Val	stop
Α	R	D	N	C	E	Q	G	Н	1	L	K	M	F	Р	S	Т	W	Υ	V	

Figure 6-50 Molecular Biology of the Cell 5/e (© Garland Science 2008)

#### triples of RNA bases encodes a amino acid



### **Central Dogma**



- Dogma: information flux
   DNA -> mRNA -> Proteins
- Gene: DNA segment coding a protein.
- Transcript: RNA segment associated to a gene.
- Genes is associated to one proteins and one function\*

\* Genes might be associated to many proteins



### **Control of Gene Expression**

How is the expression of genes controlled?

enhancer
(binding site for activator protein)

BINDING OF transcription
GENERAL TRANSCRIPTION
FACTORS, RNA POLYMERASE,
MEDIATOR, CHROMATIN REMODELING
COMPLEXES, AND HISTONE ACETYLASES

Certain proteins
(transcription factors)
bind to DNA and
initiate transcription

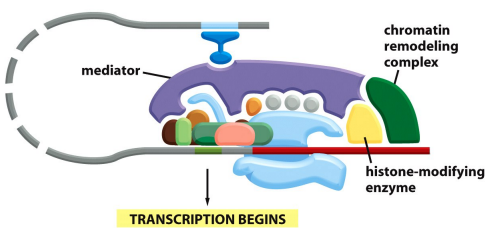
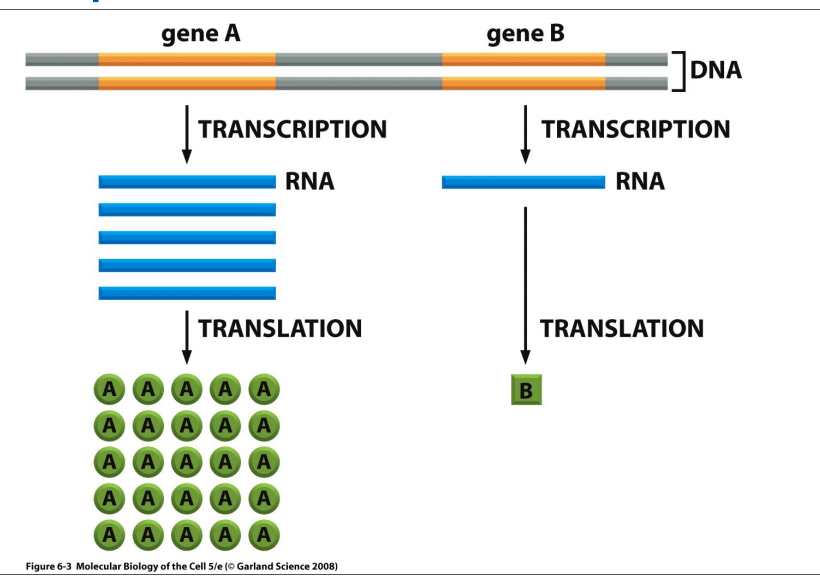


Figure 6-19 Molecular Biology of the Cell 5/e (© Garland Science 2008)



### **Gene Expression**



### **Cellular Complexity**

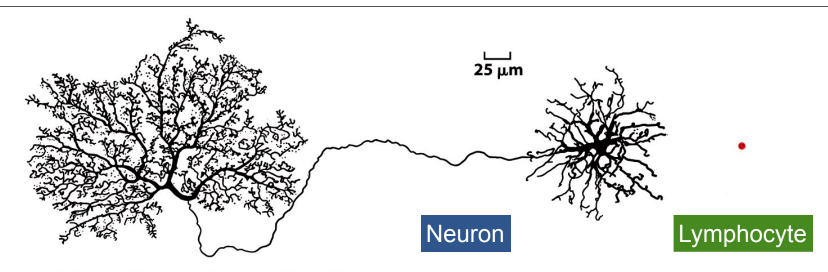


Figure 7-1 Molecular Biology of the Cell 5/e (© Garland Science 2008)

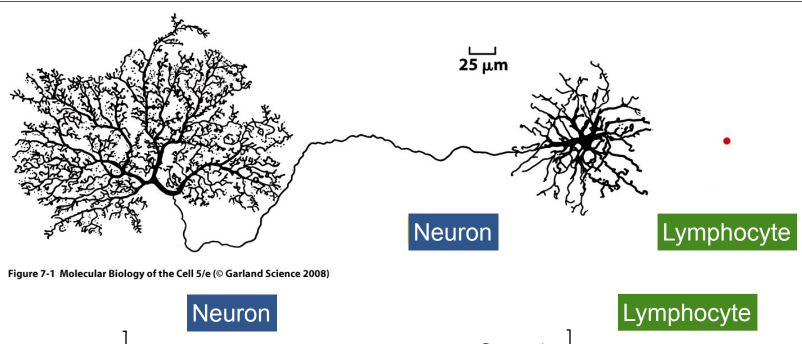
#### Two cells of a organism have exactly\* the same DNA

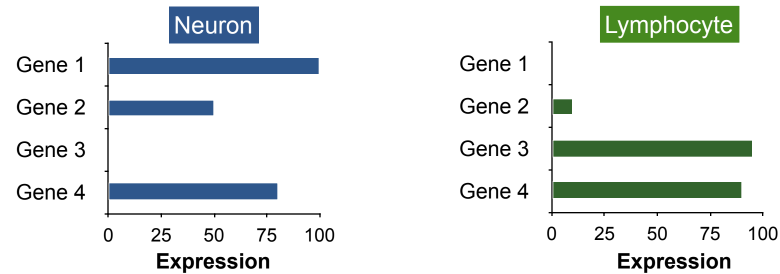
# How does this differences arise? How is cell fate remembered?

\* with exception of somatic mutations and rearrangements of immunological loci



### Cellular Complexity & Gene Expression







# Sequencing

### Sequencing

#### Read the bases of a particular DNA/RNA sequence

#### **Applications:**

- sequence DNA of known and unknown organism
- detect variants on patients
- sequence the RNA of a cell
- detect location of proteins interacting with DNA or open chromatin

#### **Problem:**

- only short DNA sequences (<1.000 bs) can be read

#### **Solution:**

break DNA in several small pieces and use bioinformatics



### **Next Generation Sequencing**

- NGS take advantage of parallelization
  - reads millions/billions of reads for a time
  - short reads (50-300 bps)
  - moderate error rates (0.1%)
- commercial products:
  - **454**
  - SOLID
  - Solexa (Illumina)



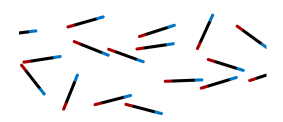


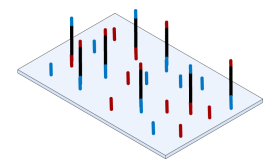
### Illumina Flow Cell - NGS Sequencing

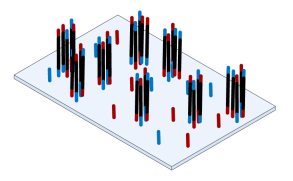
1- fragment sample DNA, insert adapters, attach to flow cell

2- use (bridge) PCR to copy fragments (close to origin)

3- clusters of single stranded DNA (200m clusters with 2k DNA strands





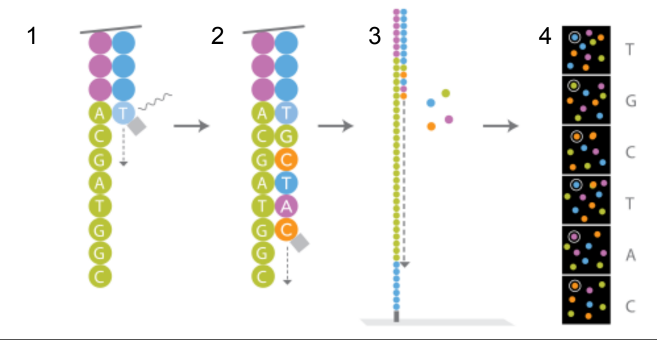


See video <a href="http://www.wellcome.ac.uk/Education-resources/Education-and-learning/Resources/Animation/WTX056051.htm">http://www.wellcome.ac.uk/Education-resources/Education-and-learning/Resources/Animation/WTX056051.htm</a>

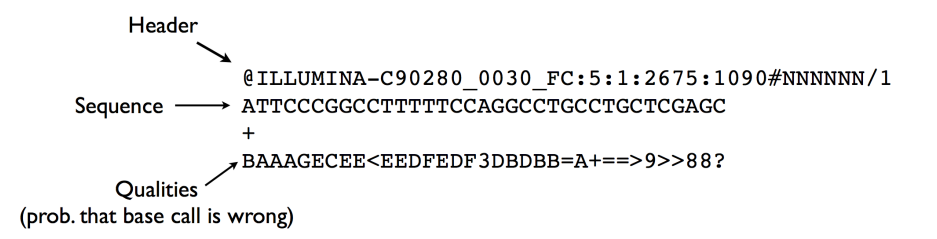


### Illumina Flow Cell - NGS Sequencing

- Iterative evaluation process:
  - 1. add RT-bases, polymerases integrate them
  - 2. wash away all not integrated elements
  - 3. take picture of flow cell to determine current base by dye
  - 4. derive reads from pictures



### **Sequencing Results**



One character encodes a number using ascii table (0-255)

This number (Q) can be converted to P

Phred-scale

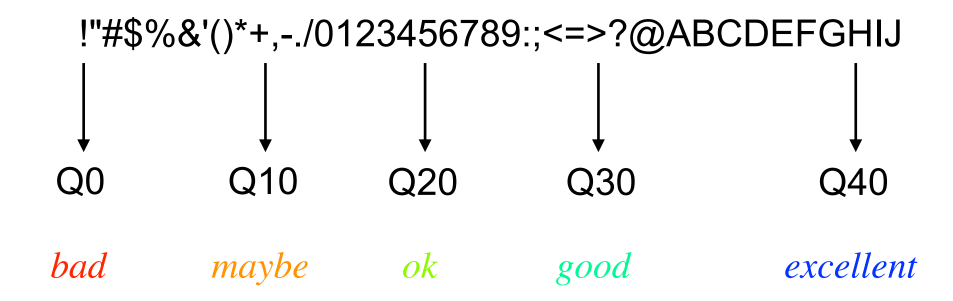
Q = -10 \* log 10 P

 $P = 10^{(-Q/10)}$ 



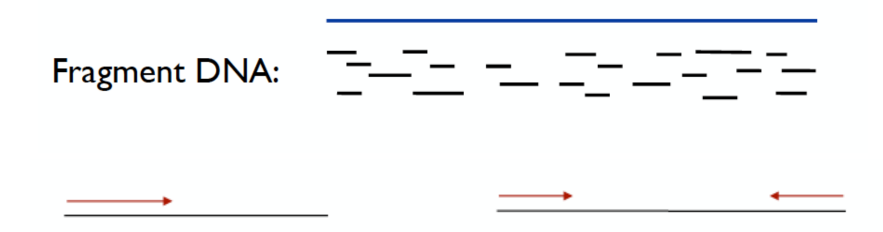
#### Sequencing Results / Phred scores

Uses letters/symbols to represent numbers:





### Read Types



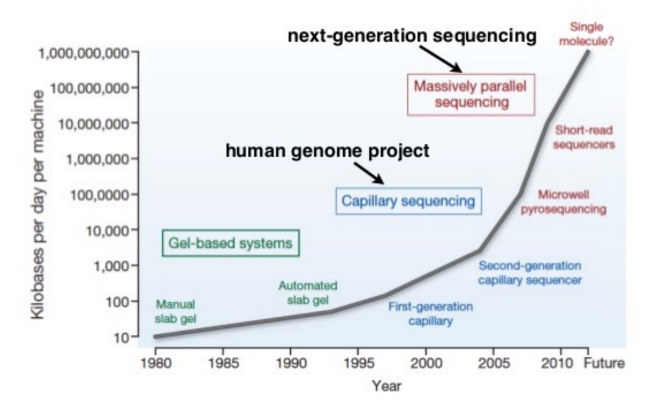
Single end

Paired end Ins: 200-800 bp



### **Next Generation Sequencing**

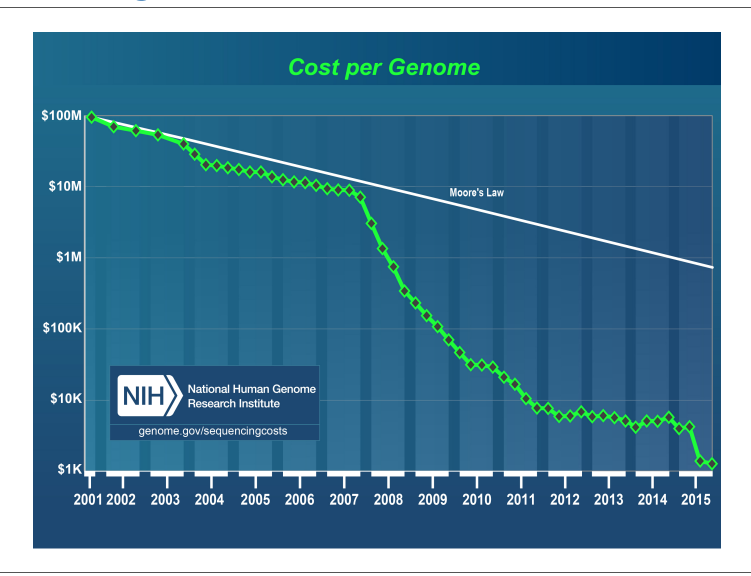
#### Improvements in the rate of DNA sequencing over the past 30 years



Stratton, M. R., Campbell, P. J. & Futreal, P. A. The cancer genome. Nature 458, 719-724 (2009).



### **Sequencing Costs**





# **Sequence Alignment**



### **Sequence Alignment**

#### NGS

- reads from DNA fragments
- position in genome is unknown
- solution: alignment

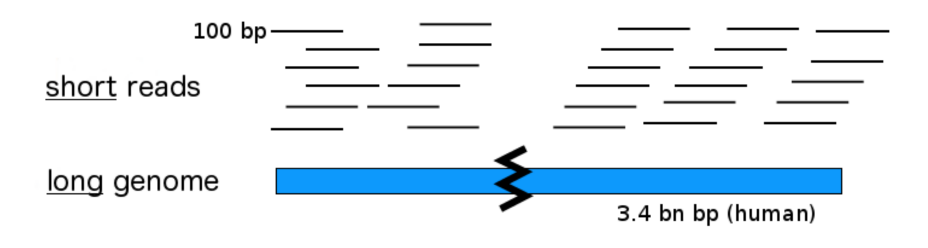
#### **DNA Sequencing**

- de-novo assembly
  - construct unknown reference sequence from scratch
- resequencing / mapping
  - reference sequence given (applies to human- and mousestudies)
  - build sequence that is similar but not necessarily identical to reference sequence



### **Alignment Problem**

- a large reference sequence is given (genome)
  - up to billions of base pairs
- millions of short reads (<200bps)</li>
- find most probable position of the read in the genome (by inexact string matching)





#### **Pitfals**

- (Unknown) divergent of sample and reference genome
- Repeats in the genome (larger than read size)
- Recombinations
- Poor genome reference quality
- Sequencing/read errors



Alignment/Mapping is a typical inexact string match problem

**Algorithmic Solutions: ?** 



Alignment/Mapping is a typical inexact string match problem

#### **Algorithmic Solutions:**

 Smith & Waterman - dynamic programming (quadratic time/memory)



Alignment/Mapping is a typical inexact string match problem

#### **Algorithmic Solutions:**

- Smith & Waterman dynamic programming (quadratic time/memory)
- Blast k-mer search for seeding followed by dynamic programming
  - large memory requirement
  - local alignment



Short read alignment is a special problem

- reference sequence is large and fixed
- query sequence (reads) are short and many

**Solution: ?** 



#### Short read alignment is a special problem

- reference sequence is large and fixed
- query sequence (reads) are short and many Solution: ?
- 1. Use a data structure to represent reference
  - k-mer hash table (>40GB for k=8)
  - suffix trees (> 4GB)

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- 2. Find candidate (k-mer) hits on genome (>100)



#### Short read alignment is a special problem

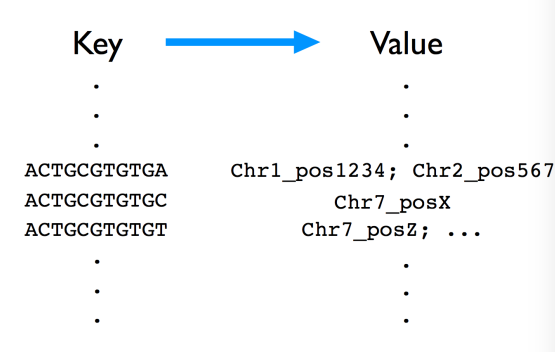
- reference sequence is large and fixed
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   Solution: ?
- 1. Use a data structure to represent reference
  - k-mer hash table (>40GB for k=8)
  - suffix trees (> 4GB)
- 2. Find candidate (k-mer) hits on genome (>100)
- 3. Improve alignment with Smith-Waterman Methods work on linear time (query sequence)



### Hash based algorithm

#### Lookups in hashes are fast!

- I. Index the reference using *k*-mers.
- 2. Search reads vs. hash k-mers
- 3. Perform alignment of entire read around seed
- 4. Report best alignment

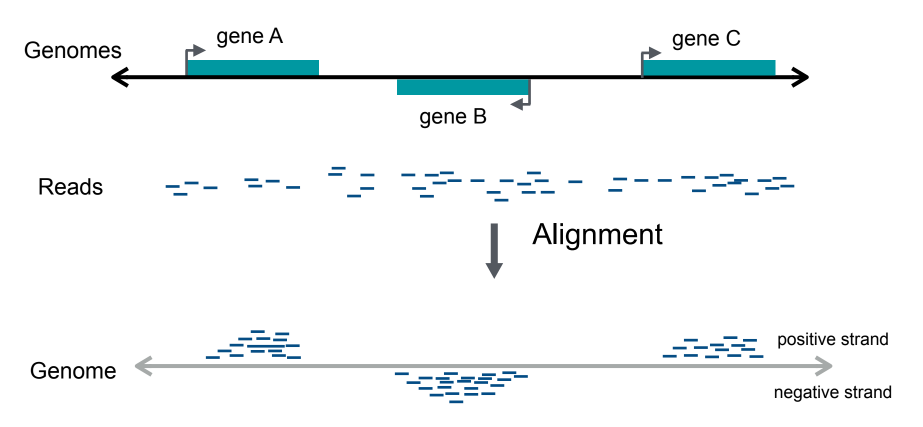


Also known as Seed and extend



### RNA sequencing / Alignment Results

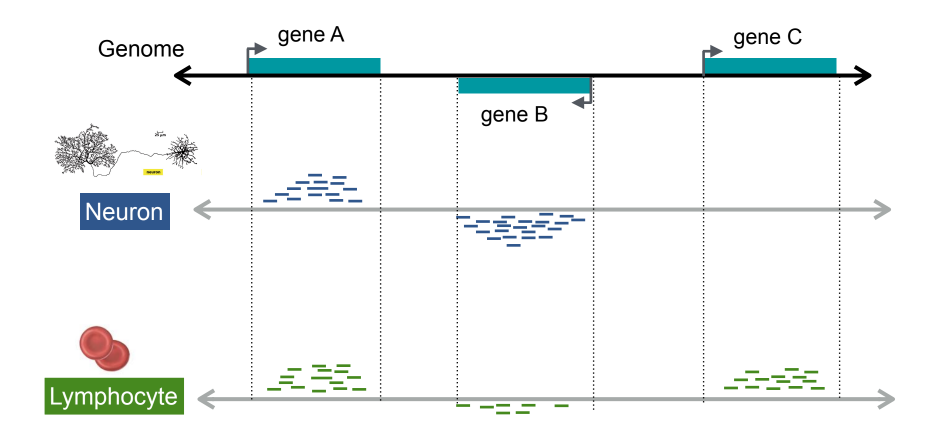
- Position and strand of reads aligned to the genome





#### **Gene Quantification**

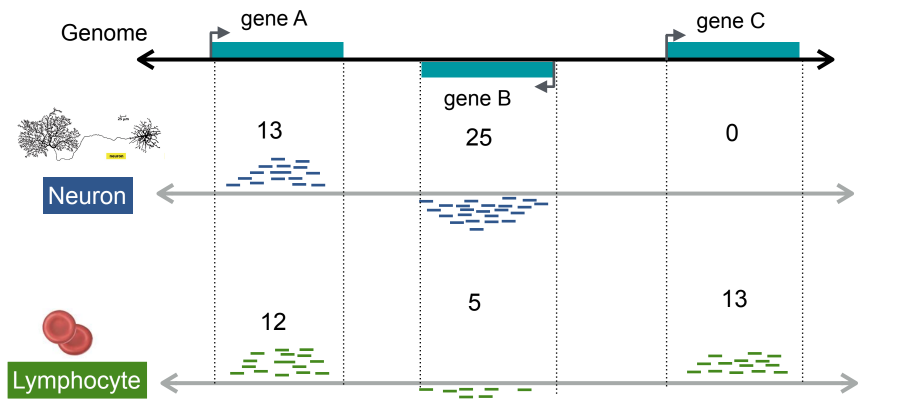
- Perform sequencing for each cell (neuron, lymphocyte)
- Align reads to genome





#### **Gene Quantification**

- Perform sequencing for each cell (neuron, lymphocyte)
- Align reads to genome
- Count number of reads inside genes (using known genes annotation)





#### **Quantification - Normalization**

#### Correct for:

- Genes having distinct size
- Sequencing efficiency differs between cell (usually same RNA quantity provided for sequencing)

	Cell A	Cell B	
GeneA (1kb)	20	15	30
GeneB (2kb)	100	300	10
GeneC (1.5kb)	10	20	100
Gene D (3kb)	300	200	100
Total Library	430	535	240

Reads per kilobase million (RPKM) = #reads \* gene size \* total library

1.000 1.000.000



#### Resume

- Review basic biological/computational aspects
  - 1. basics of molecular biology
  - 2. basics of sequencing
  - 3. basics bioinformatics problems
    - short sequences read alignment
    - gene expression quantification
    - single cell sequencing (next week)

