

SevenBridges

# Epigenetic control

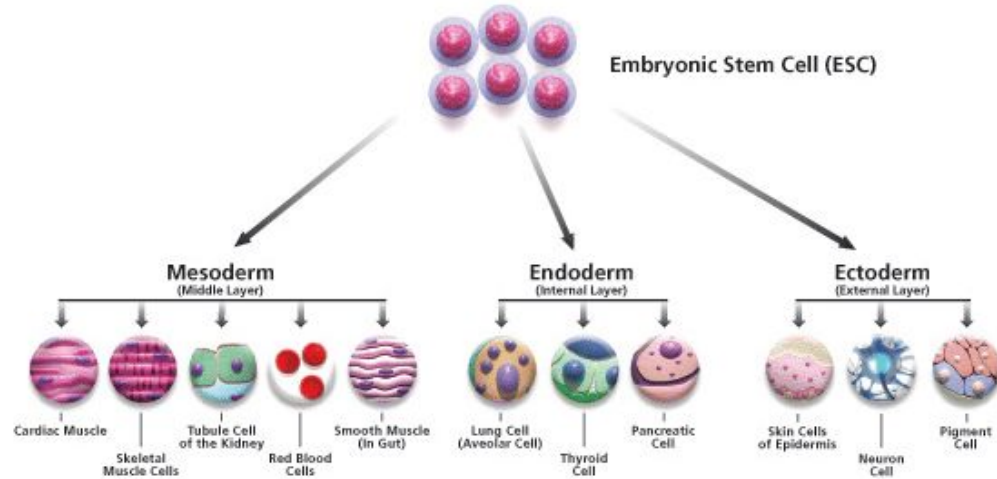
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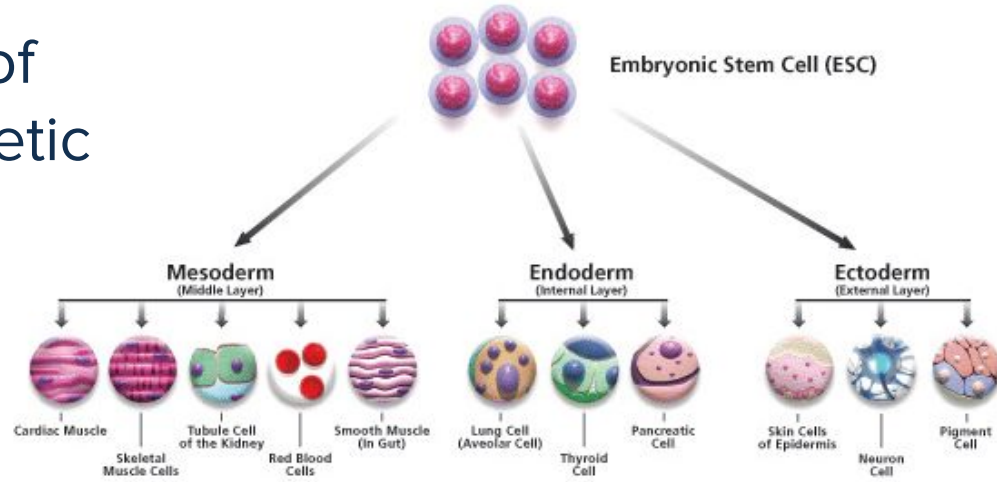
# Epigenetic control

- Same genotype - different phenotypes
- How?



# Epigenetic control

- **Epigenetics** - extra layer of information on top of genetic information
- Each cell type is defined by genes that are **expressed** in the cell



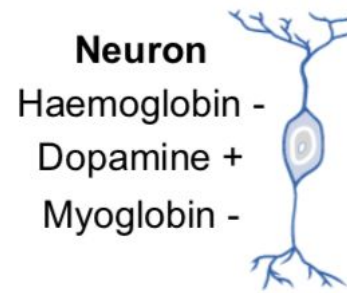
Gene expression

DNA → RNA → protein

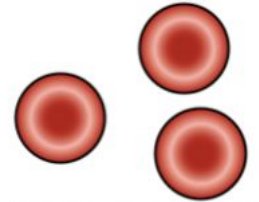
Single cell fertilised egg  
1 cell type



Each cell type only expresses a restricted  
subset of genes.



**Neuron**  
Haemoglobin -  
Dopamine +  
Myoglobin -



**Red blood cells**  
Haemoglobin +  
Dopamine -  
Myoglobin -



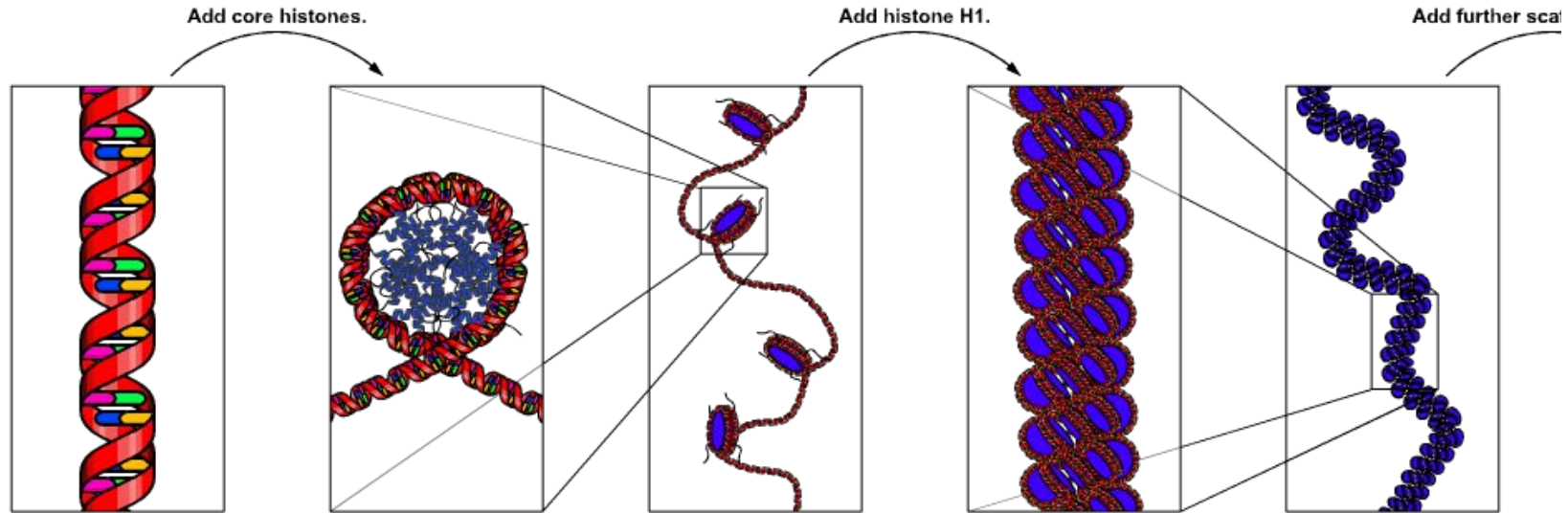
**Muscle cells**  
Haemoglobin -  
Dopamine -  
Myoglobin +

# Epigenetic modifications

- Demarcate the start and end of genes
- Provide structure to the chromosome
- Alter how we read each and every gene
  - genes being expressed (active)
  - genes being not expressed (silent)

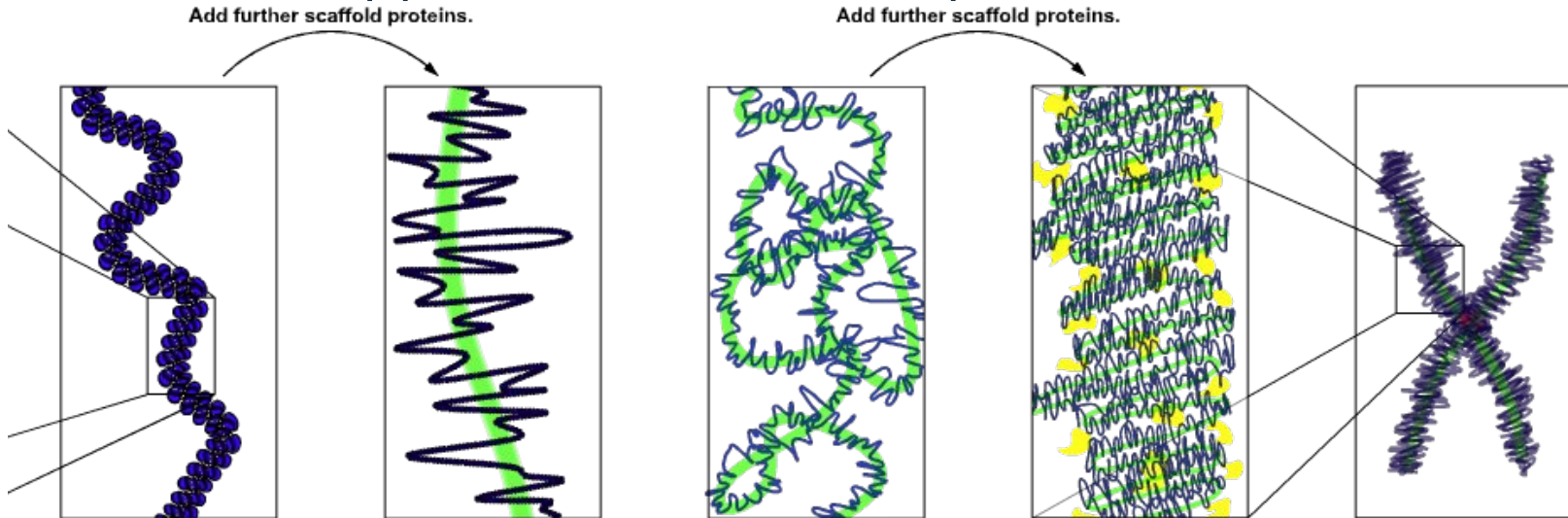
# DNA structure refresher

- DNA is compacted into chromatin
- DNA is wrapped around histone proteins



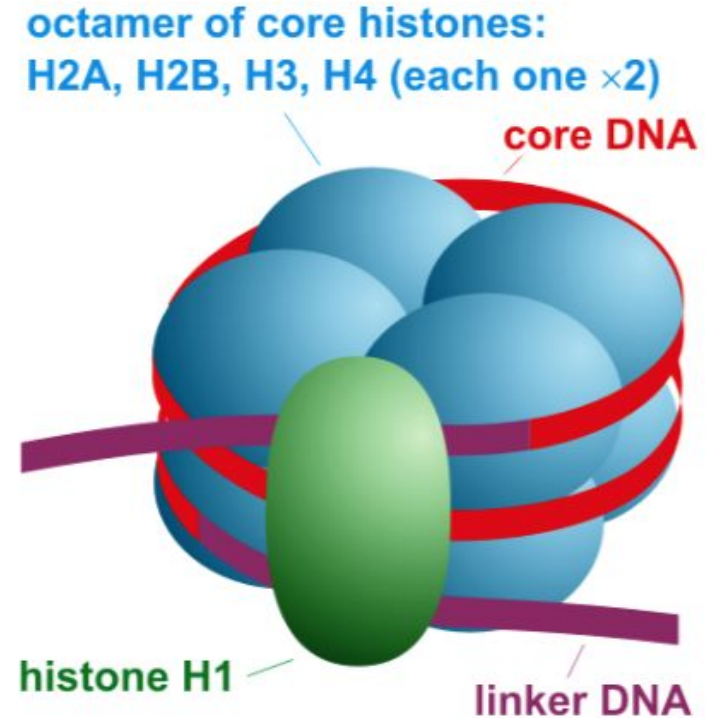
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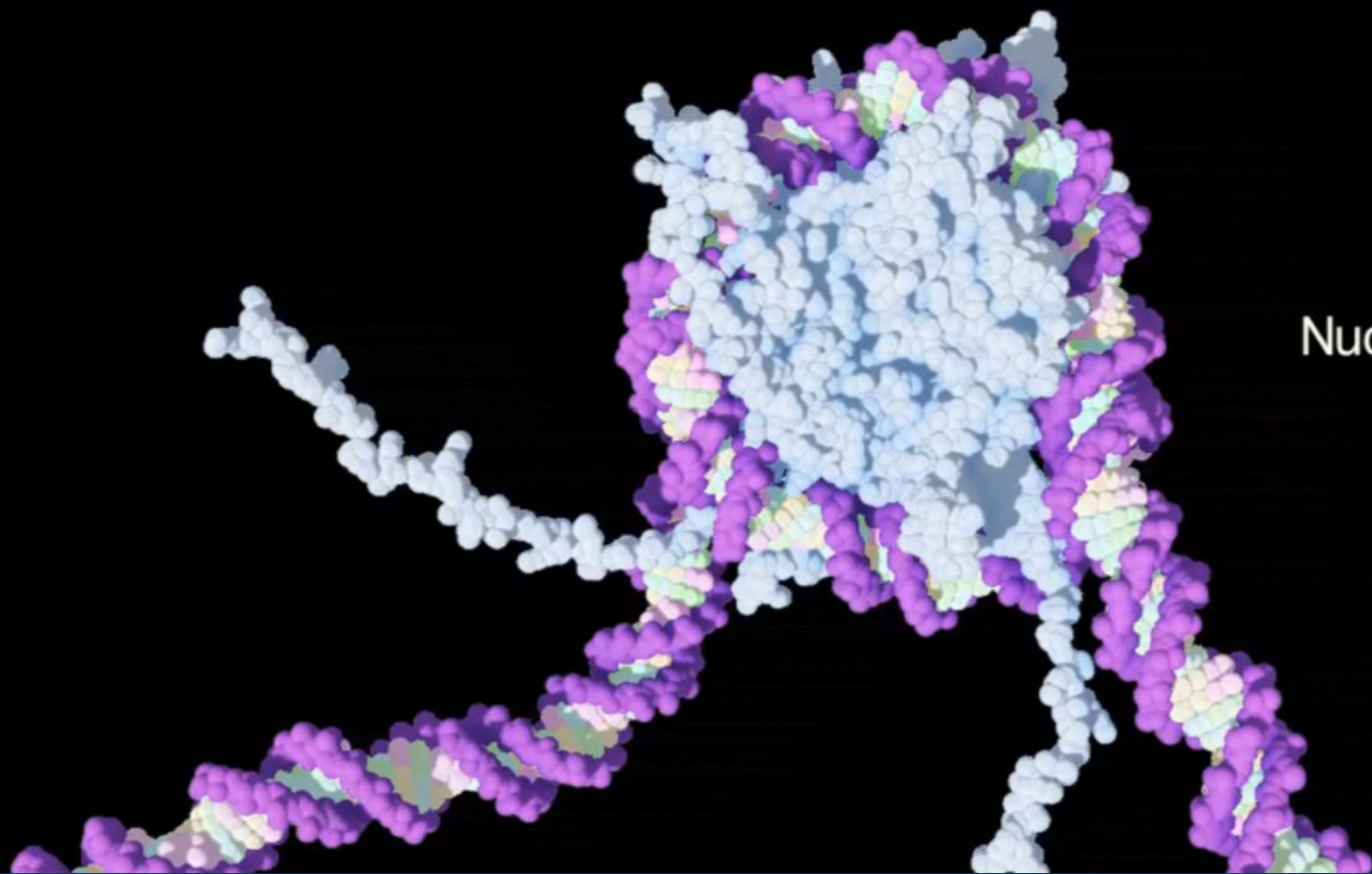


# Nucleosome

- 146bp of DNA wrapped around a histone octamer
- Positively charged histones bind to negatively charged DNA







Nucleosome

# Heterochromatin versus euchromatin

- Heterochromatin - closed chromatin
  - **Facultative** - can differ by cell type or time
    - tissue specific genes
    - parts of X chromosome
  - **Constitutive** - same in all cell types - structural role
    - Centromeres
    - Telomeres
    - Parts of sex chromosomes
- Euchromatin - open chromatin
  - DNA accessible for transcription

# Specific epigenetic modifications

- **DNA methylation**
- **Histone modifications**
- Chromatin remodelling
- Histone variation
- Noncoding RNAs

# DNA methylation

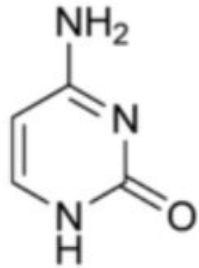
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Changes to DNA amino acids

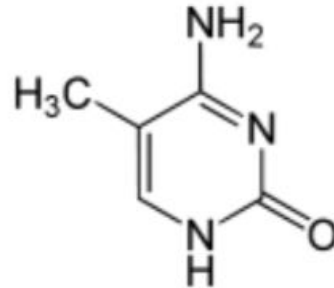


# DNA methylation

- Methylation of cytosine nucleotide
- Almost exclusively on CpG dinucleotides



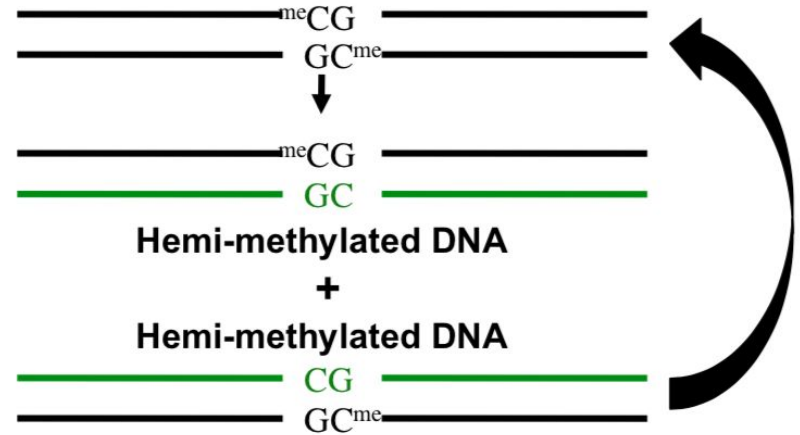
Cytosine



5- methyl cytosine

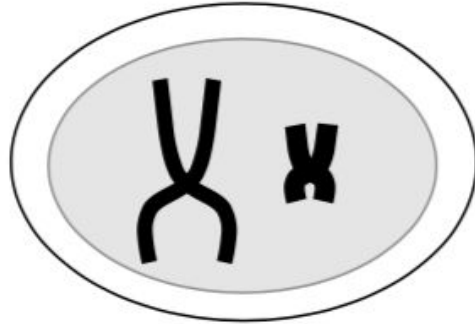
# CpG site

- CpG - just a G following a C in sequence
- **CpG islands**
  - Why are they rare?
- CpG islands at gene promoters
  - Methylated → gene expression silenced

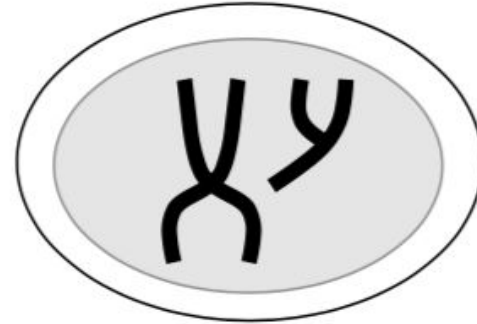


# X inactivation

- Epigenetic dosage compensation mechanism in mammals -males and females have the same dose of genes on the X chromosome
- Inactive X chromosome shows DNA methylation of CpG islands



Female XX



Male XY

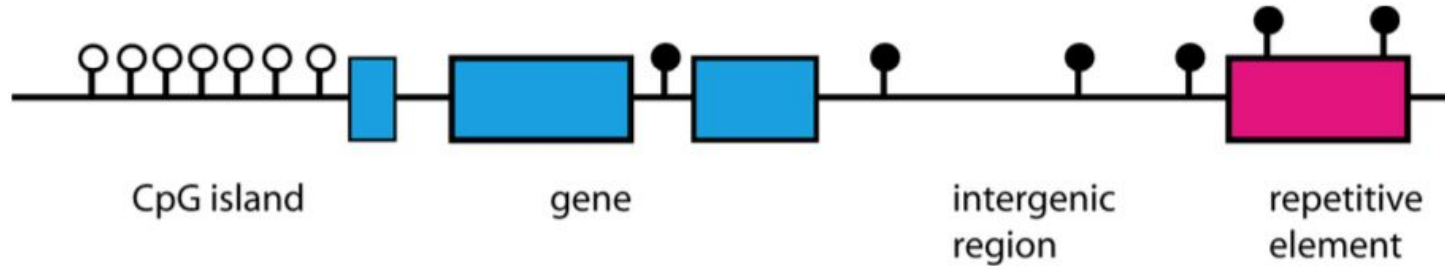
# X inactivation





# DNA methylation - where does it occur?

- CpG islands – usually unmethylated
- Intergenic regions – usually methylated
- Repetitive elements – usually methylated

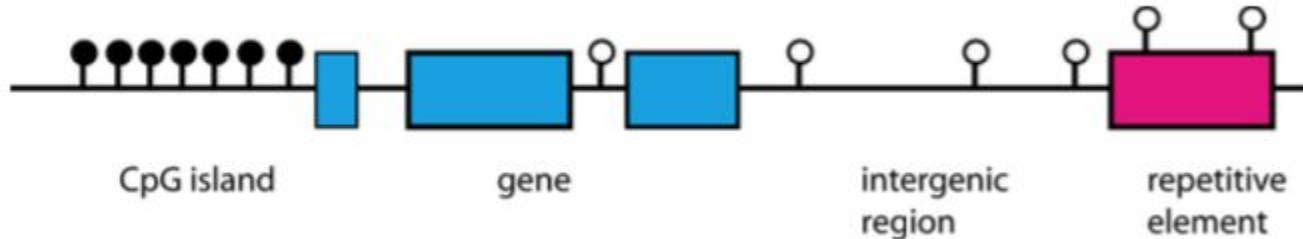


# DNA methylation - function

- Intergenic regions - Maintain genomic stability
- Repetitive regions
  - Silence repeats to prevent transposition
  - Mutate transposable elements
- Oncogenes
  - Silence expression

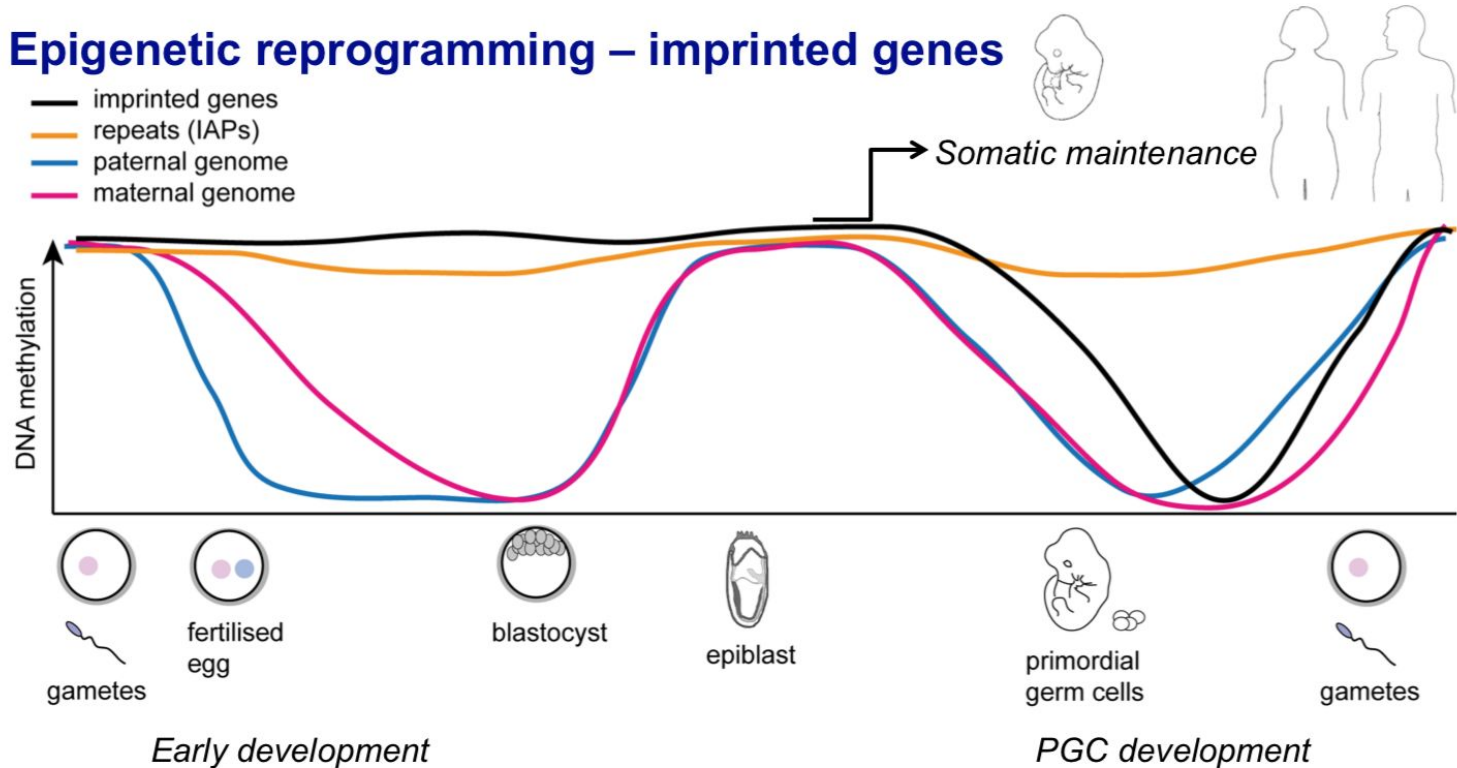
# DNA methylation in cancer

- **Hypermethylation** of CpG islands
  - Disables tumor suppressors
- **Hypomethylation** of genome
  - Leads to genomic instability



# DNA methylation through aging

## Epigenetic reprogramming – imprinted genes



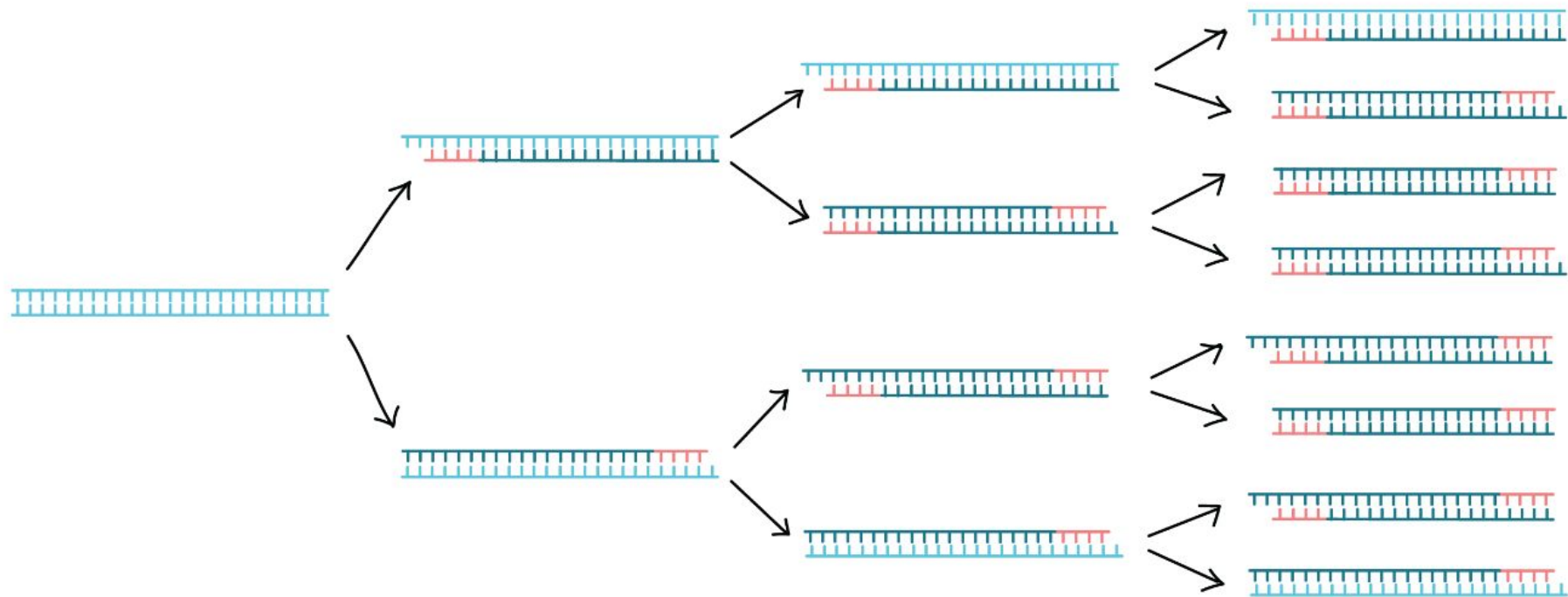
# Bisulfite sequencing

- Bisulfite modification converts non-methylated cytosines to uracils
- PCR amplification results in replacement of uracils with thymines

# PCR

Recap





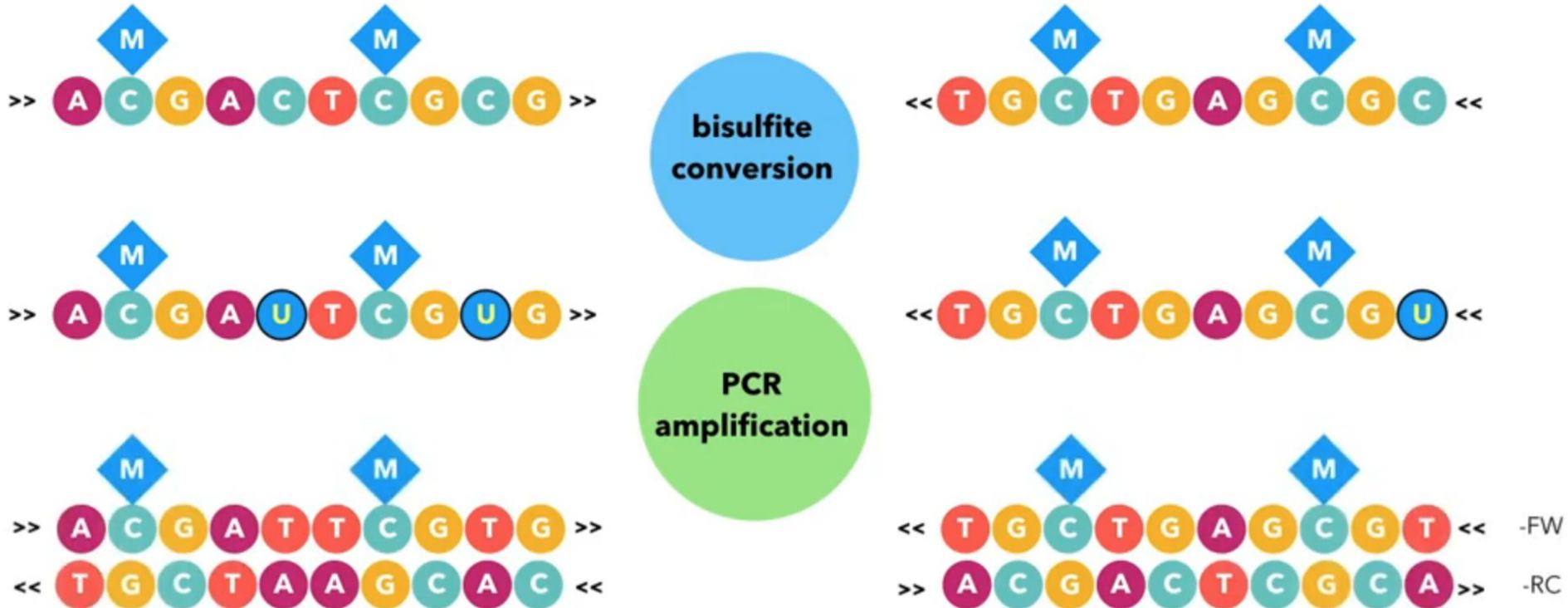
cycle:

1

2

3

## Bisulfite sequencing





# Bisulfite sequencing - alignment

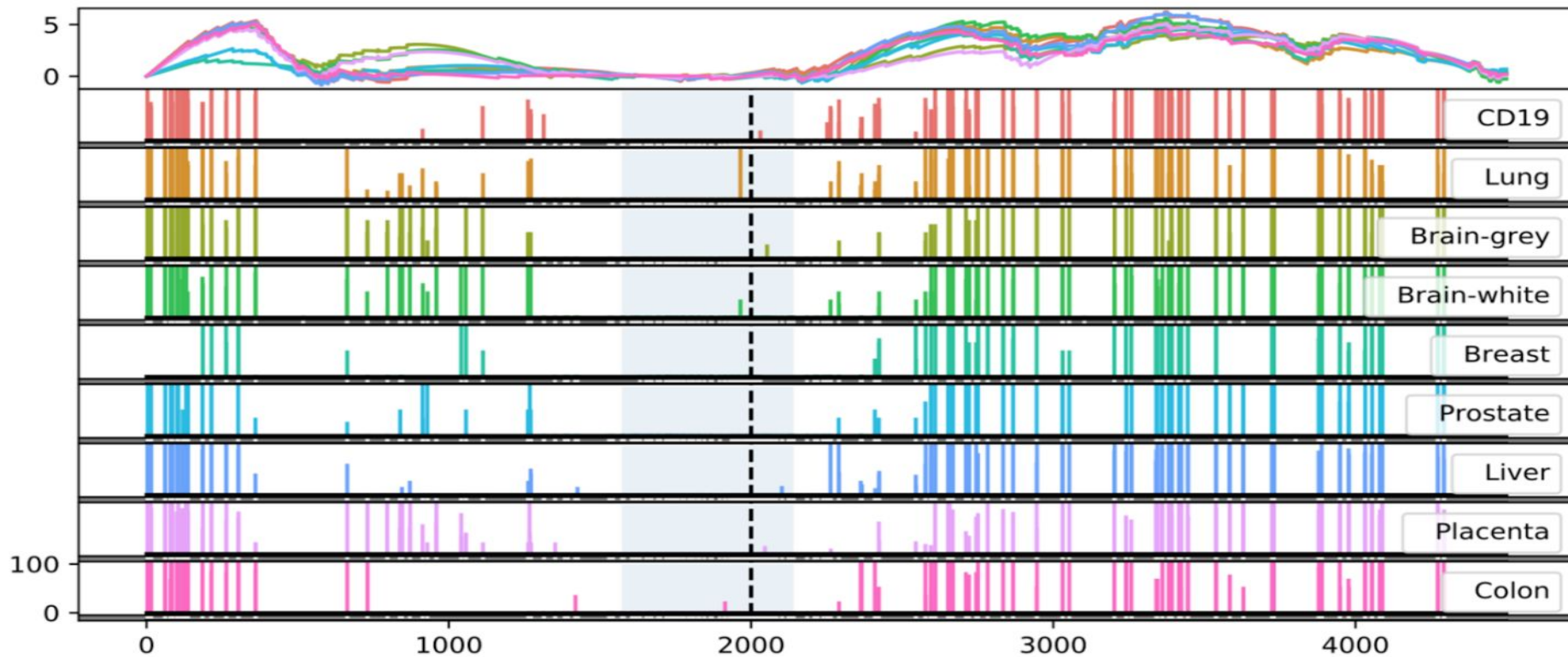
- Similar to regular alignment
- Differences?

# Bisulfite sequencing - alignment

- Dependant on library preparation protocol
  - Case when single strand is used
  - Case when both strands are used

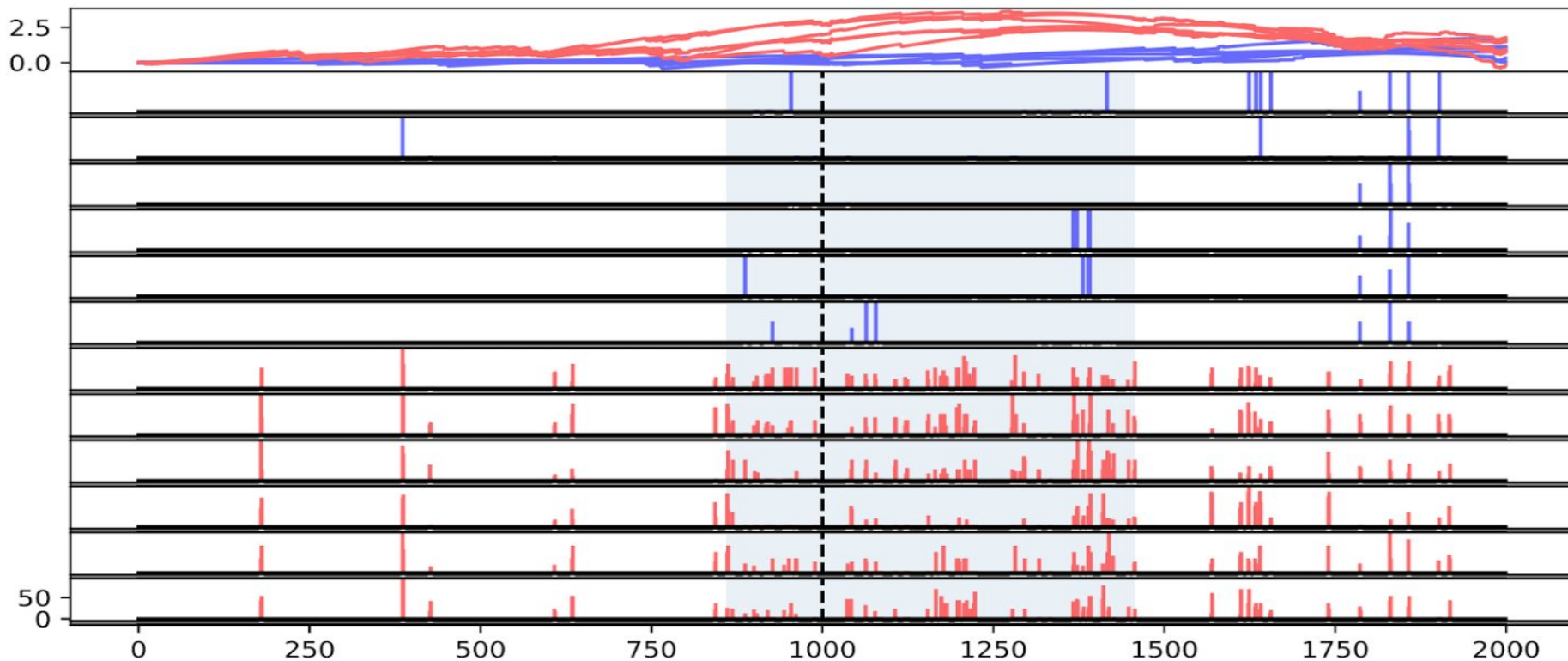
# Differential methylation - between cell types

GUSB



# Differential methylation - in tumor

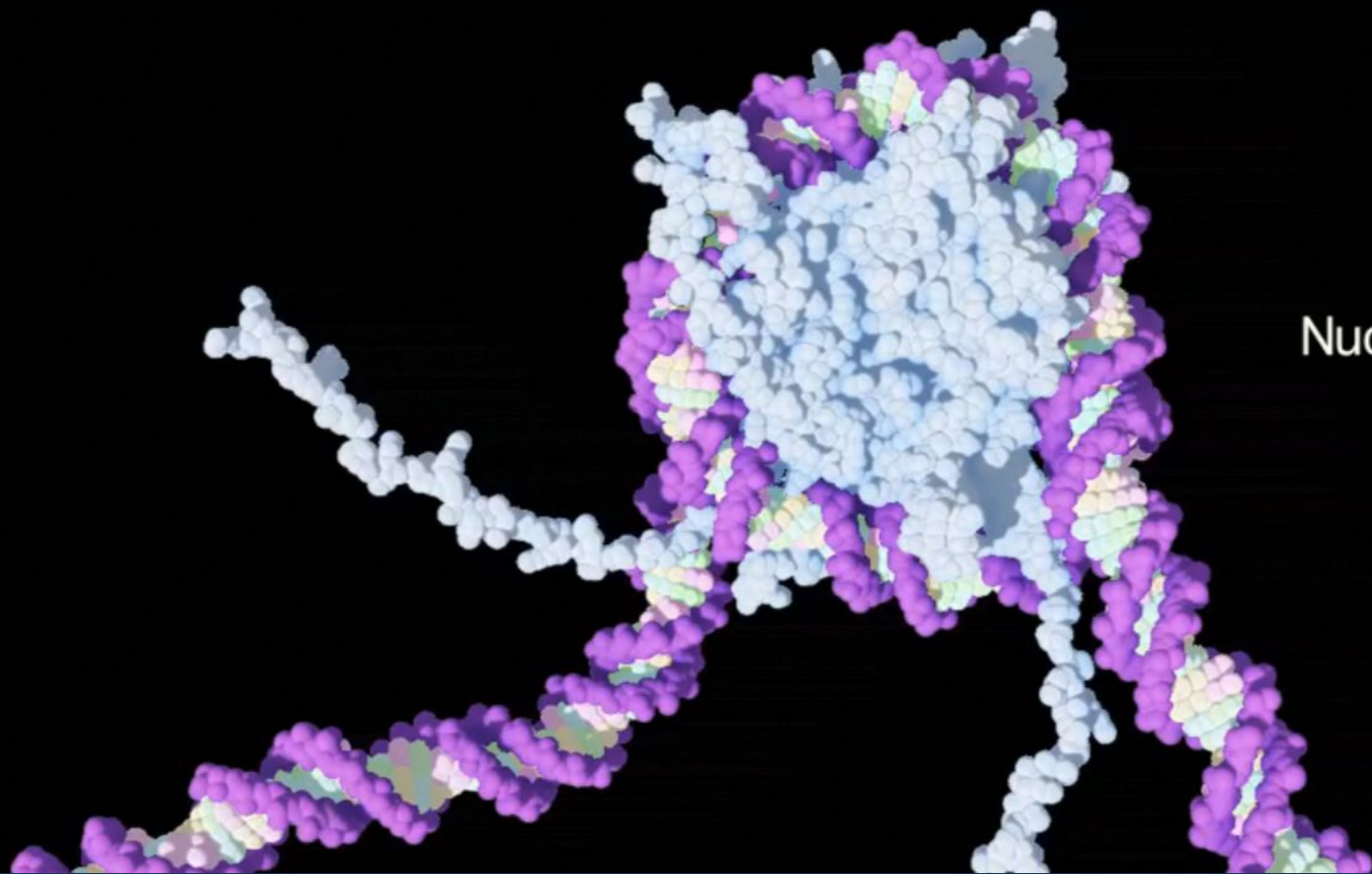
PCDHGB6\_1



# Histone modification

Modifications to structure holding DNA

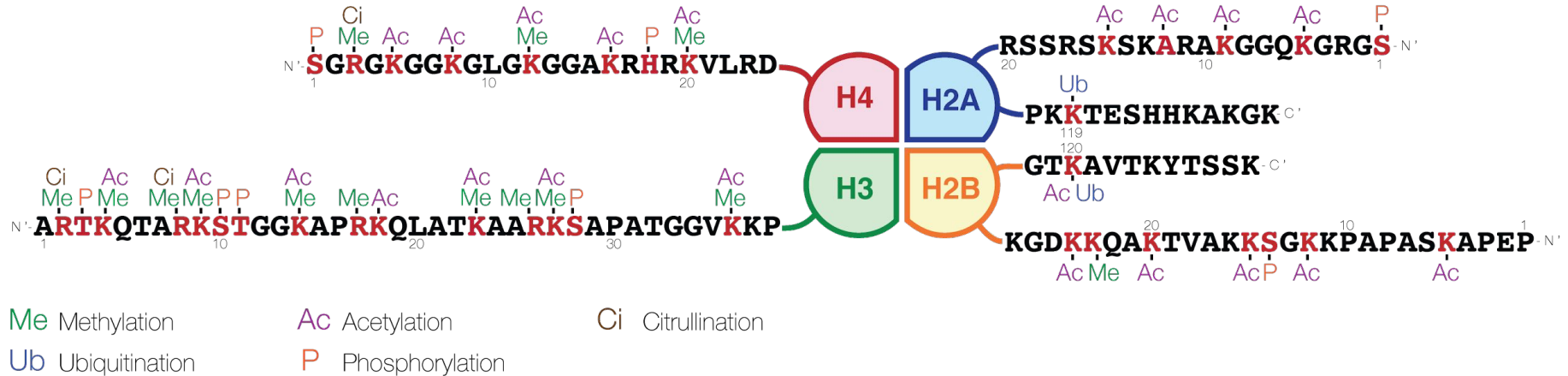




Nucleosome

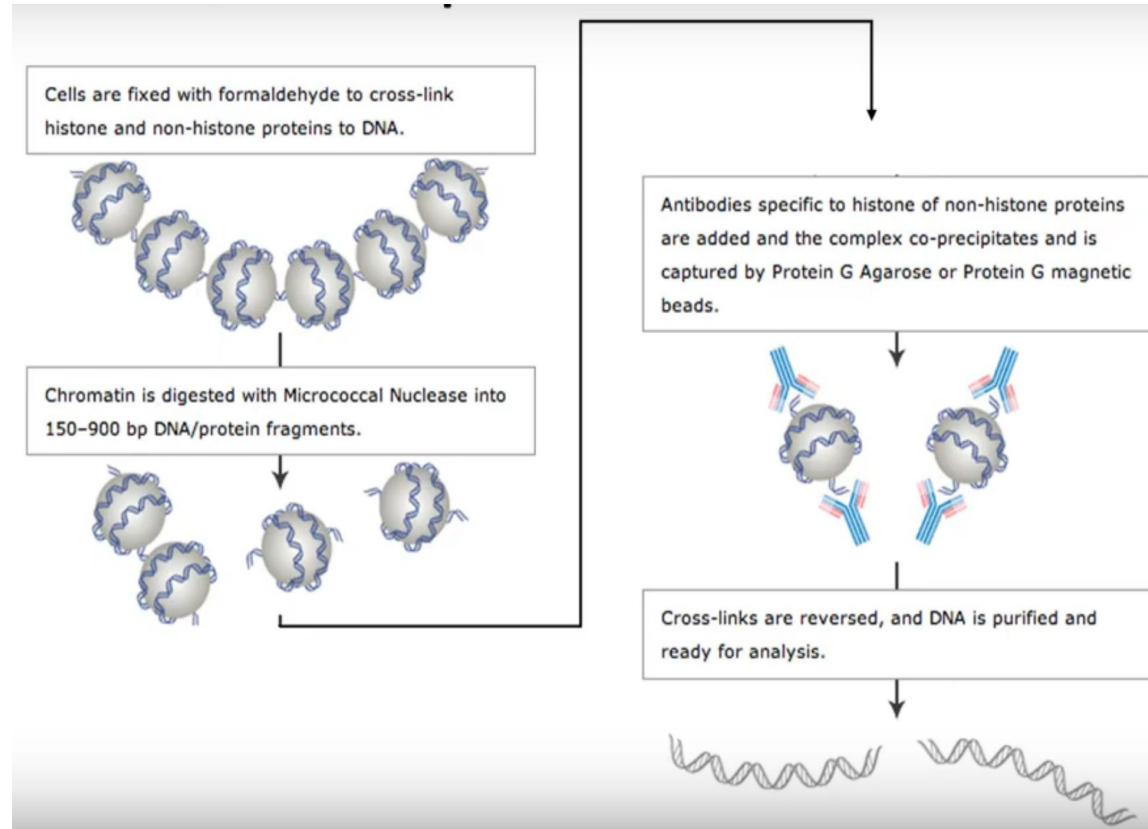
# Histone modifications

- Chemical modifications of histone tails
- More than 50 different modifications



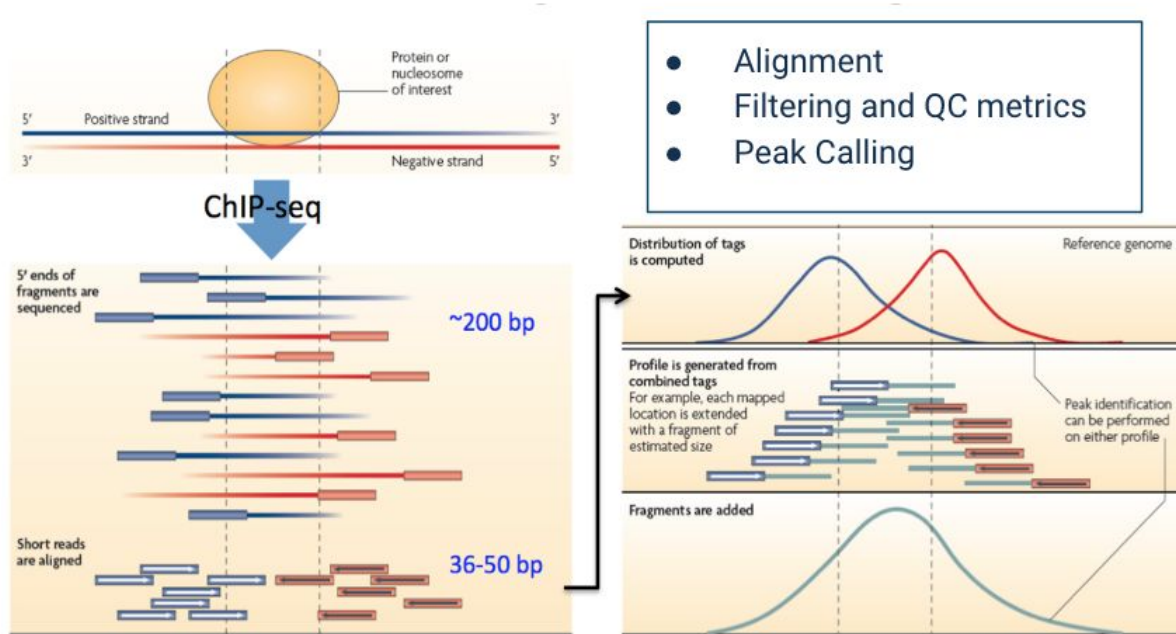
- Chromatin ImmunoPrecipitation

## ChIP-seq





# ChIP-seq



# Other epigenetic modifications

- Chromatin remodeling - dependent on ATP
- Histone variants
- Noncoding RNAs
  - piRNAs
  - siRNAs
  - lncRNAs
- RNA modifications

# Literature

- [Epigenetics review](#)
- [Bisulfite sequencing](#)
- [Bismark - bisulfite-seq aligner](#)
- [Differential methylation](#)
- [Chip-seq](#)

# Differential methylation exercise

```
import re
from Bio import SeqIO

reference = SeqIO.parse(open("/sbgenomics/project-files/example_human_reference.fasta"), 'fasta')
fasta_1 = SeqIO.parse(open("/sbgenomics/project-files/example_1.fasta"), 'fasta')
fasta_2 = SeqIO.parse(open("/sbgenomics/project-files/example_2"), 'fasta')
ref_chromosome = list(reference)[0]
f1_chromosome = list(fasta_1)[0]
f2_chromosome = list(fasta_2)[0]

ref_sequence = str(ref_chromosome.seq)
f1_sequence = str(f1_chromosome.seq)
f2_sequence = str(f2_chromosome.seq)
```