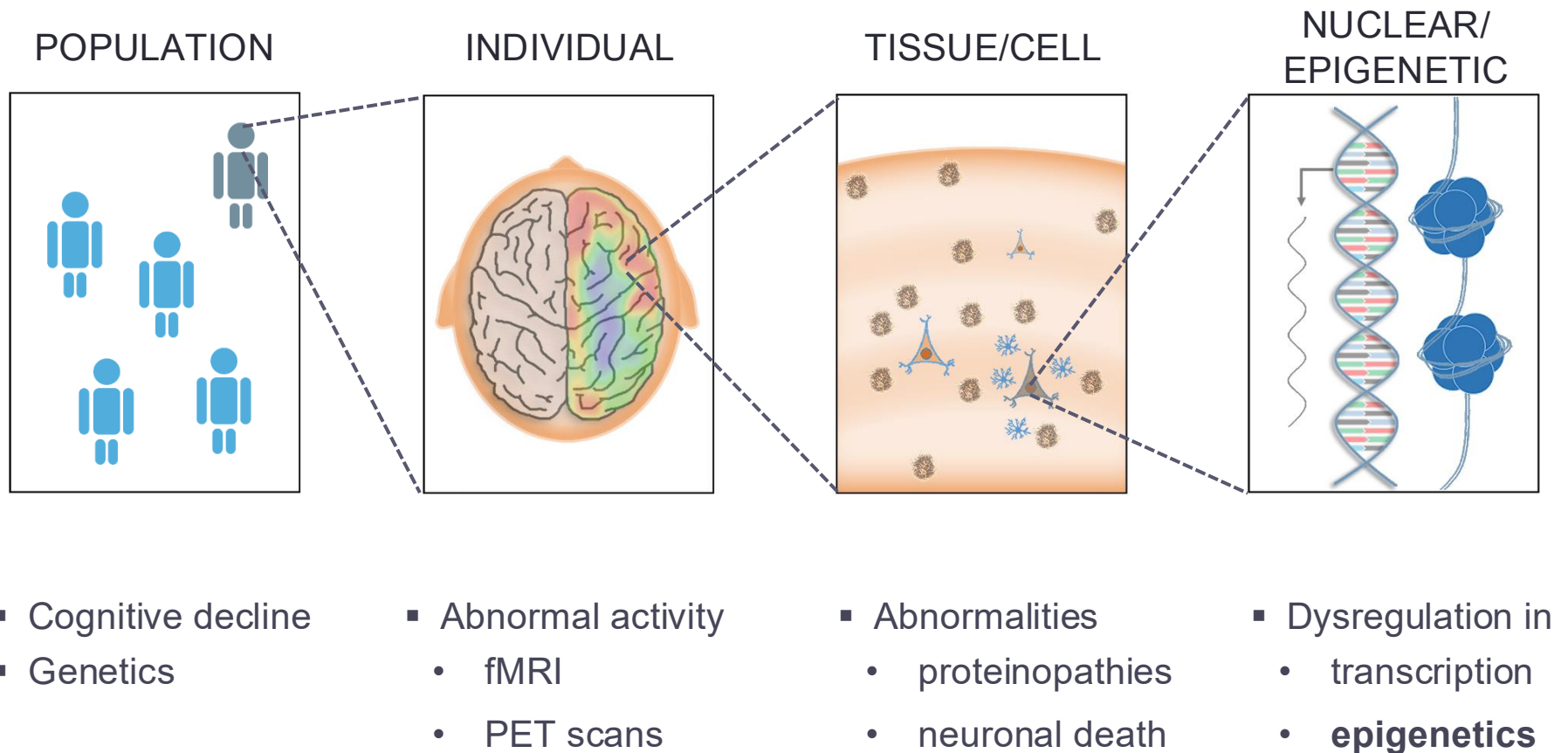


WEEK 11

NEUROEPIGENETICS

Levels of investigation in neurodegenerative disorders



Neuroepigenetics

1) The chromatin – Epigenetic basics (Lecture 1)

- Chromatin condensation
- Regulation of chromatin structure
- Environmental influences on epigenetics
- Epigenetic inheritance

2) Epigenetic dysregulation (Lecture 2)

- in AD

Learning objectives

At the end of this week you will be able to

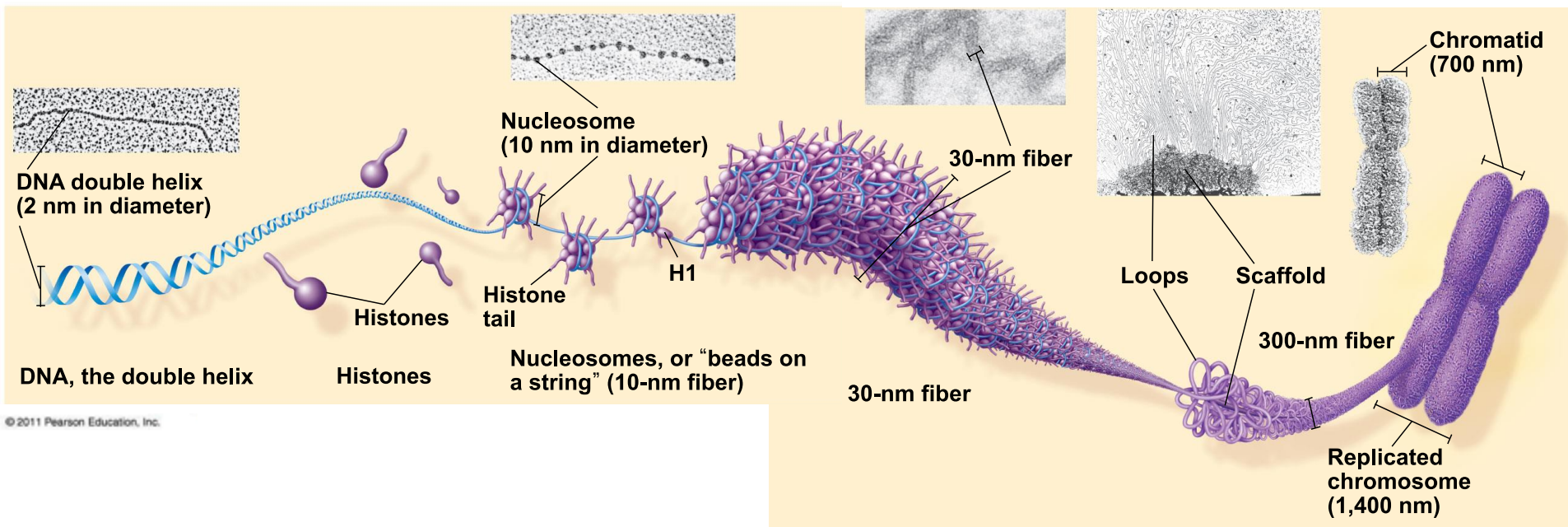
- Know about the core components of chromatin
- Define core epigenetic mechanisms of gene expression
- Know examples of how epigenetics can be influenced by the environment
- Differentiate between genetic and epigenetic modes of inheritance
- Explain how early life stress can alter adult behavior via epigenetic modifications

Insulator
Nucleosome
Barrier
Histone
Phosphorylation
MBD
Promoter
Chromatin
Acetylation
Methylation
Ubiquitination
Intergenerational
Housekeeping
CpG
H1
Transgenerational
DNA
Euchromatin
H2AX
Heterochromatin
Nr3c1

Some definitions:

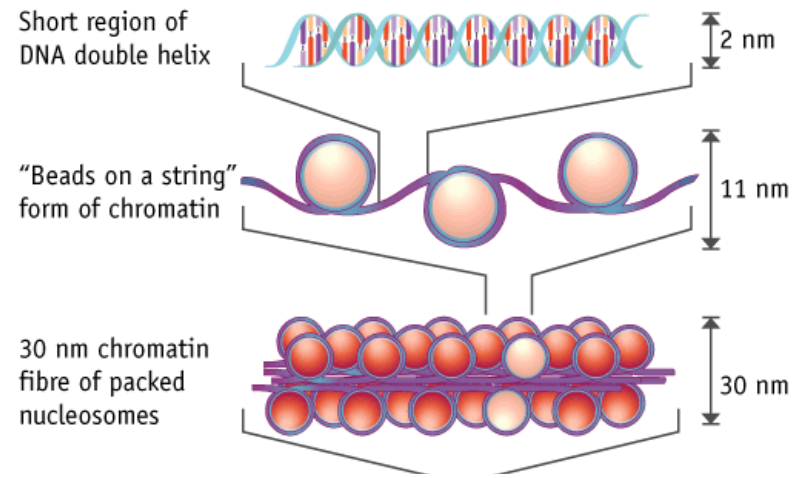
- **Chromatin** = DNA + histones and nonhistone chromatin binding proteins
- **Epi-genetic** = “On” or “above” the genes
 - “The structural adaptation of chromosomal regions so as to register, signal or perpetuate altered activity states” Adrian Bird
- **3 main types of structural adaptations:**
 - Posttranslational modifications of histone proteins
 - DNA methylation
 - Non-coding RNAs

Chromatin condensation:

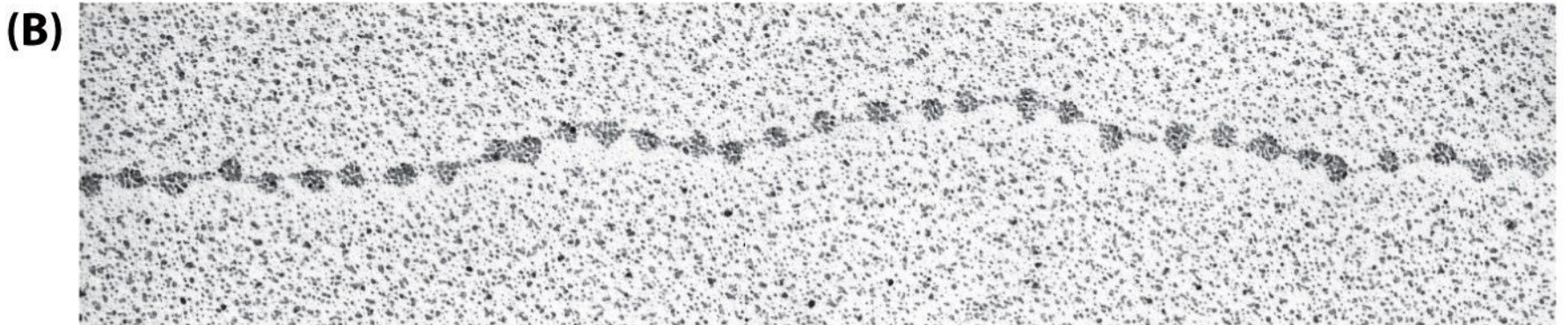
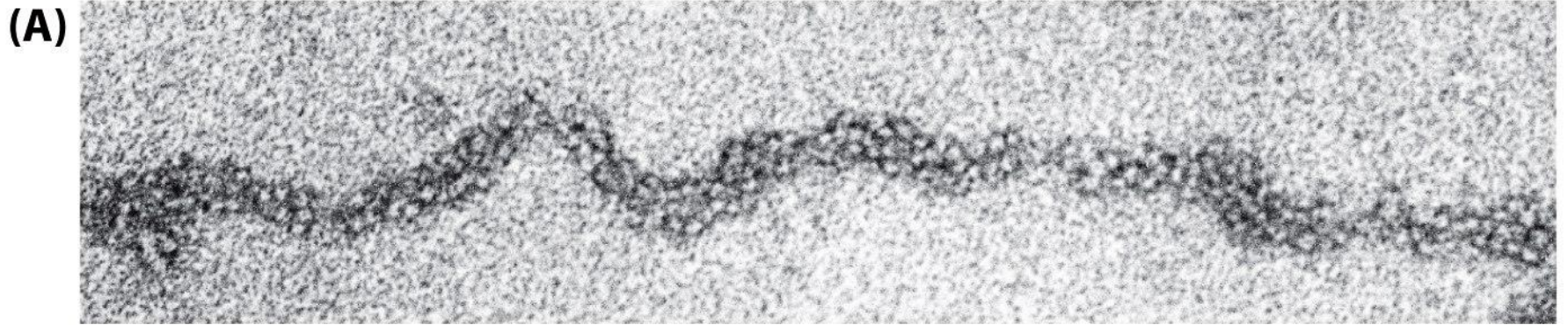


Chromatin condensation:

- 2nm: double helix
- 11nm: beads on a string (“collier en perles”)
- 30nm: fibre of condensed nucleosomes



Chromatin condensation:



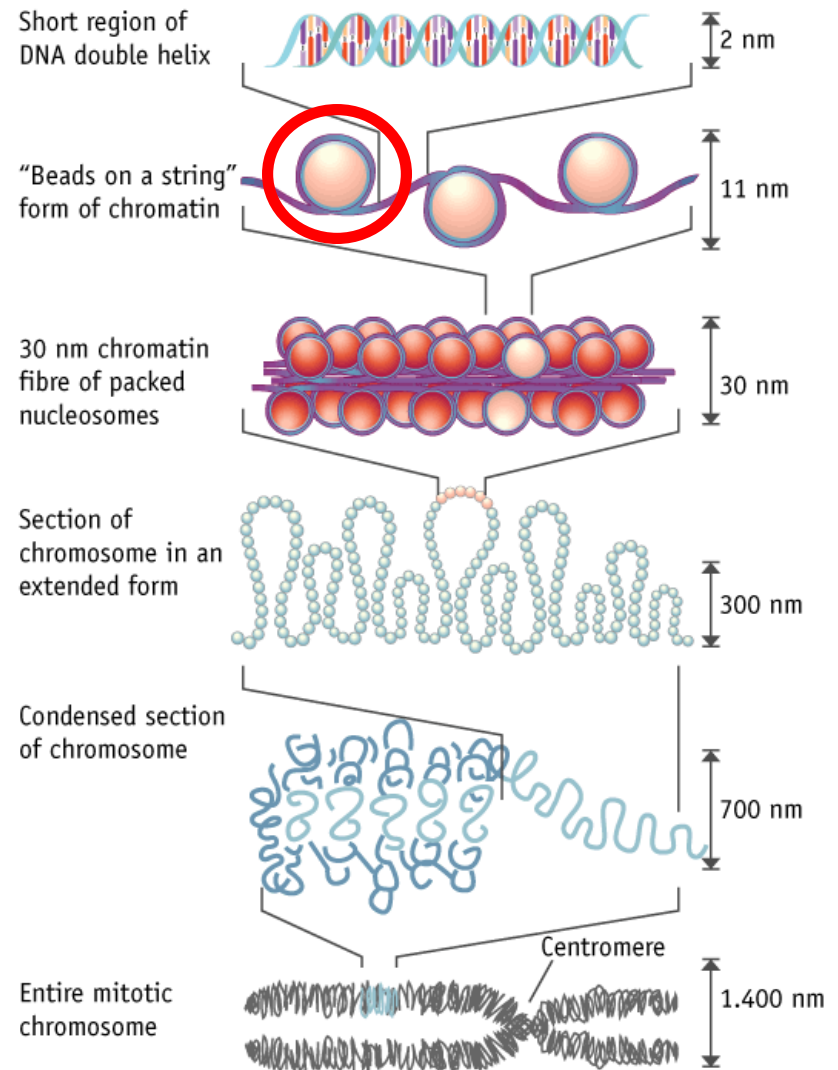
50 nm

(A): 30nm fibre

(B): 11nm beads on a string

Chromatin condensation:

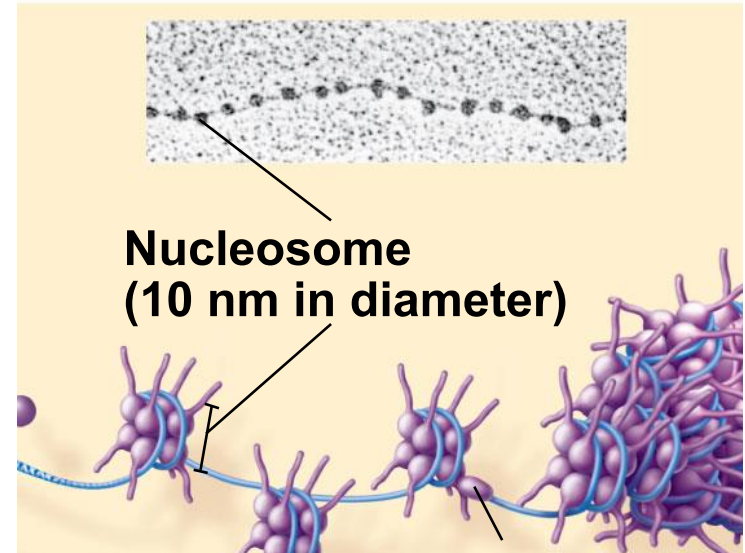
- 2nm: double helix
- 11nm: beads on a string (“collier en perles”)
- 30nm: fibre of condensed nucleosomes
- 300nm: chromatin loops



Net result: 10'000x condensation in length

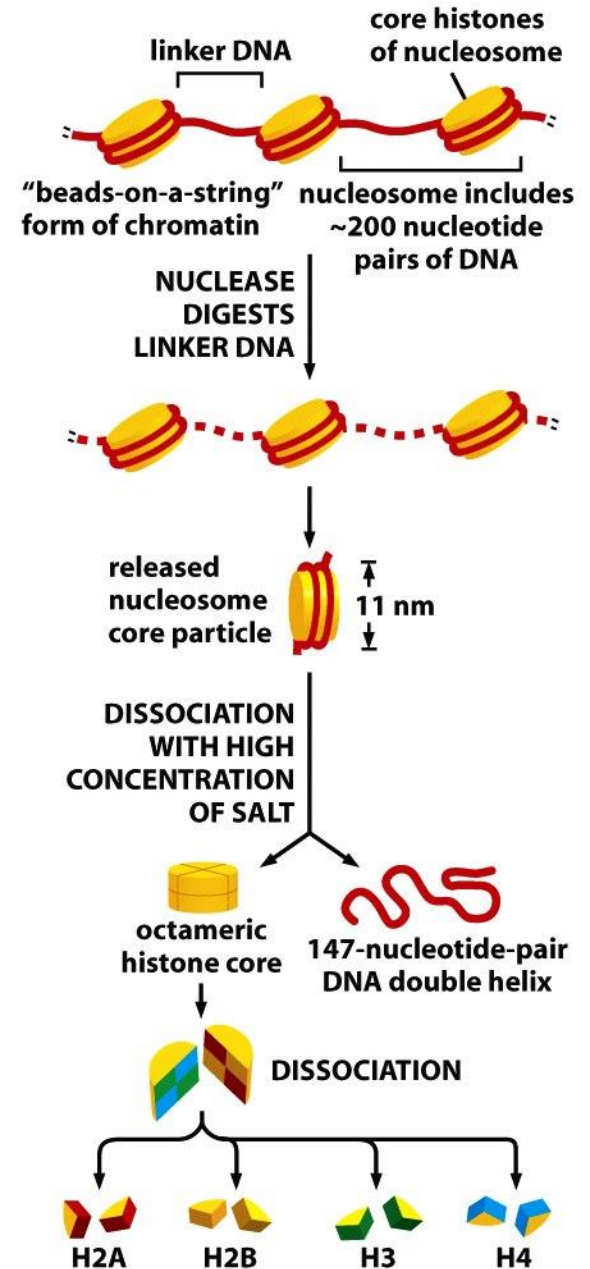
The nucleosome

- Contains about 200bp of DNA
- Nucleosome core particle:
 - 147bp DNA
 - 8 core histone proteins (histone octamer)
 - H2A (2)
 - H2B (2)
 - H3 (2)
 - H4 (2)
- Linker DNA and linker histone

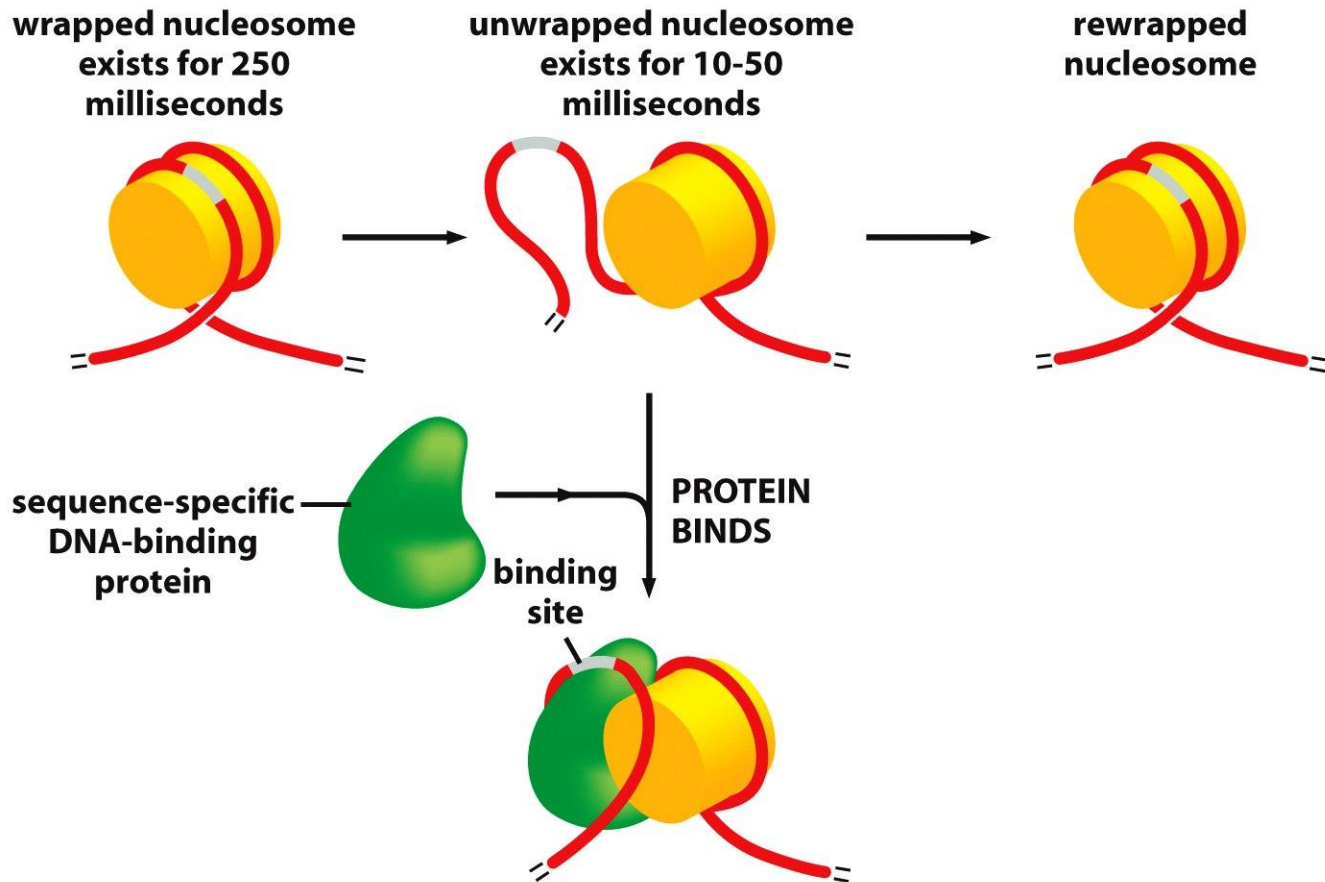


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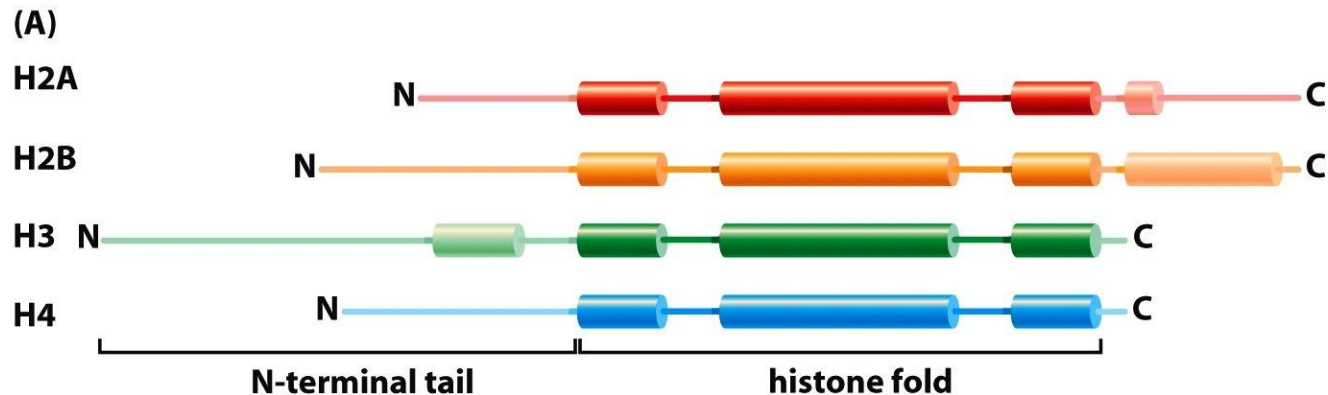


Dynamic nature of the nucleosome



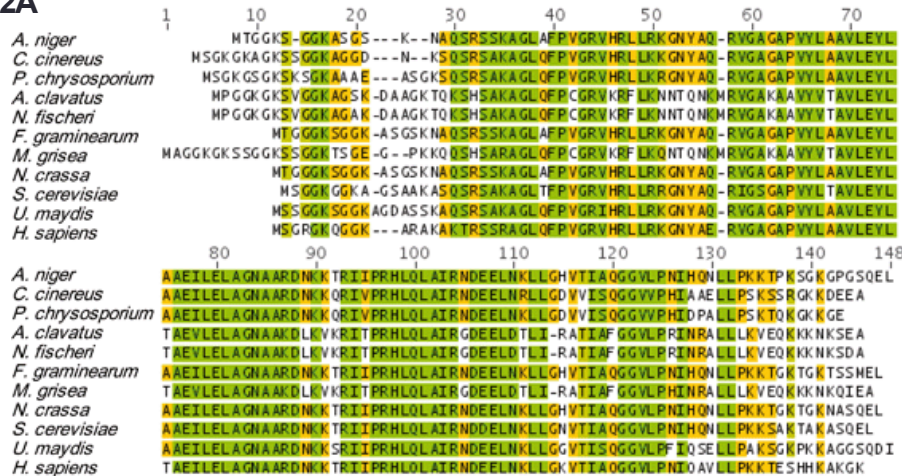
The core histones

- Histone fold: Important for dimerization
- N-terminal domain: Important for posttranslational modifications

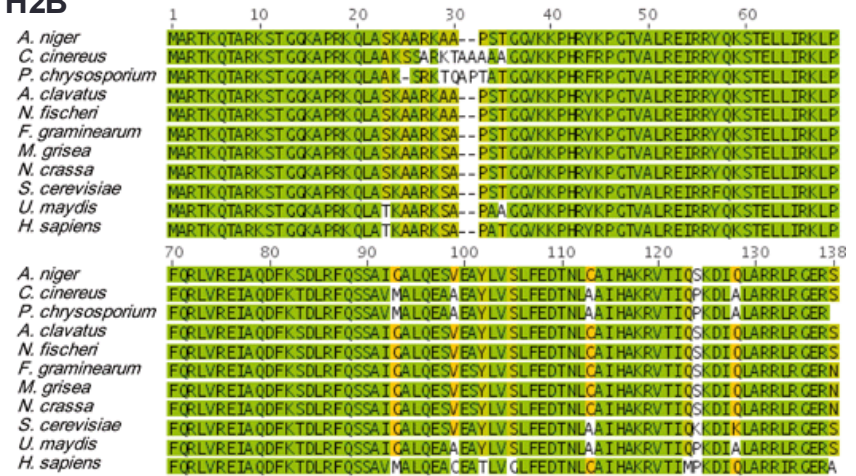


Evolutionary conservation of the core histones

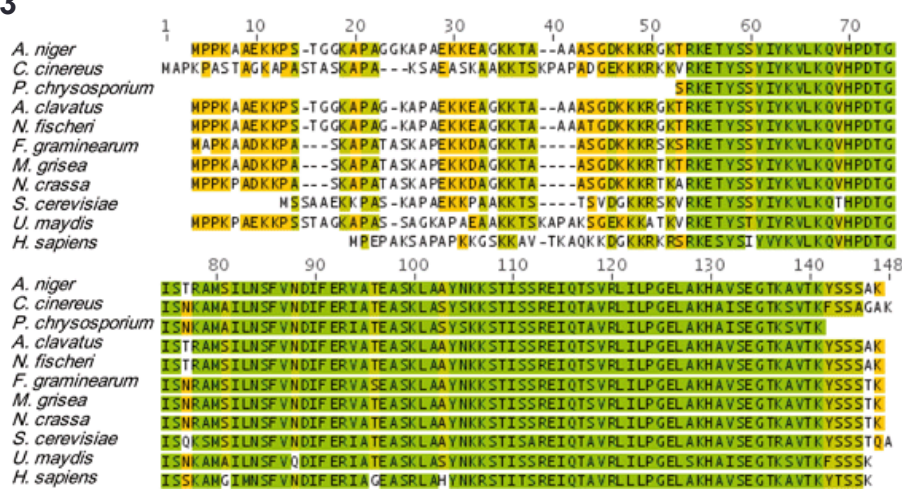
H2A



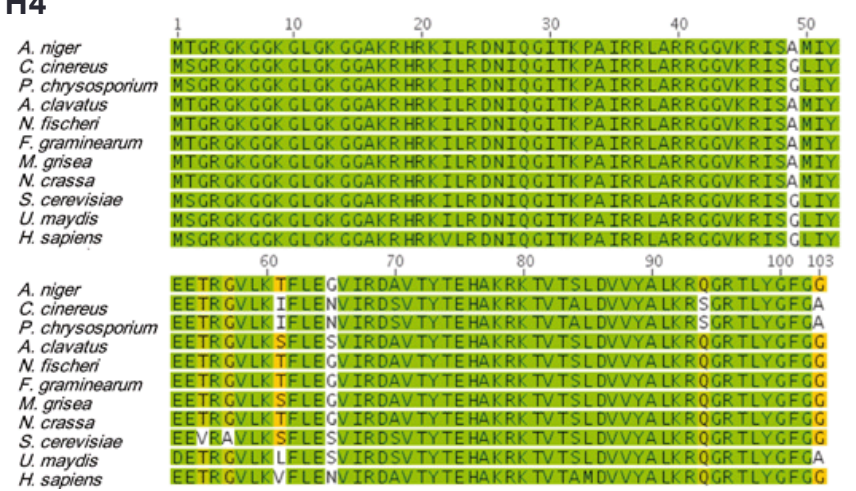
H2B



H3



H4



Neuroepigenetics

1) **The chromatin – Epigenetic basics (Lecture 1)**

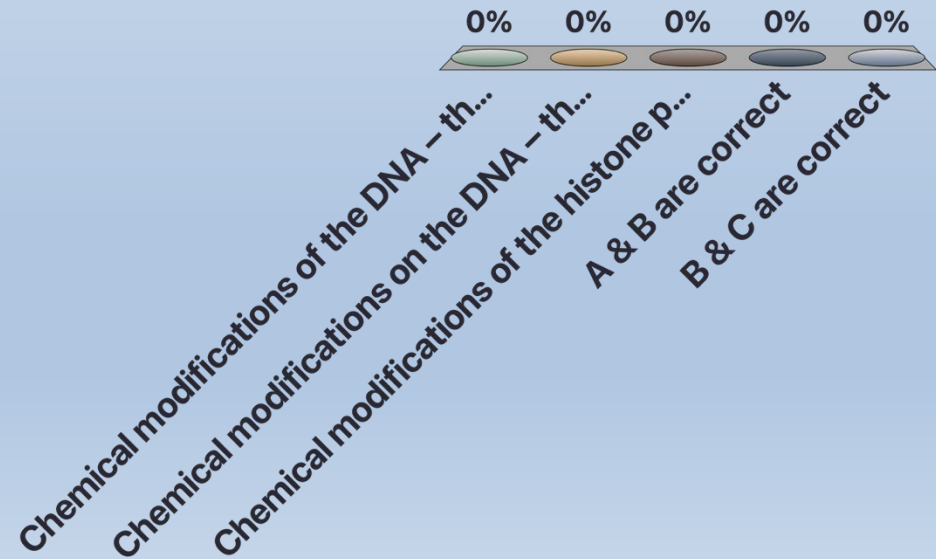
- Chromatin condensation
- **Regulation of chromatin structure**
- Epigenetic inheritance
- Environmental influence on epigenetics

2) **Epigenetic dysregulation (Lecture 2)**

- in AD

What is epigenetics?

- A. Chemical modifications of the DNA – the sequence changes
- B. Chemical modifications on the DNA – the sequence rests intact
- C. Chemical modifications of the histone proteins
- D. A & B are correct
- E. B & C are correct



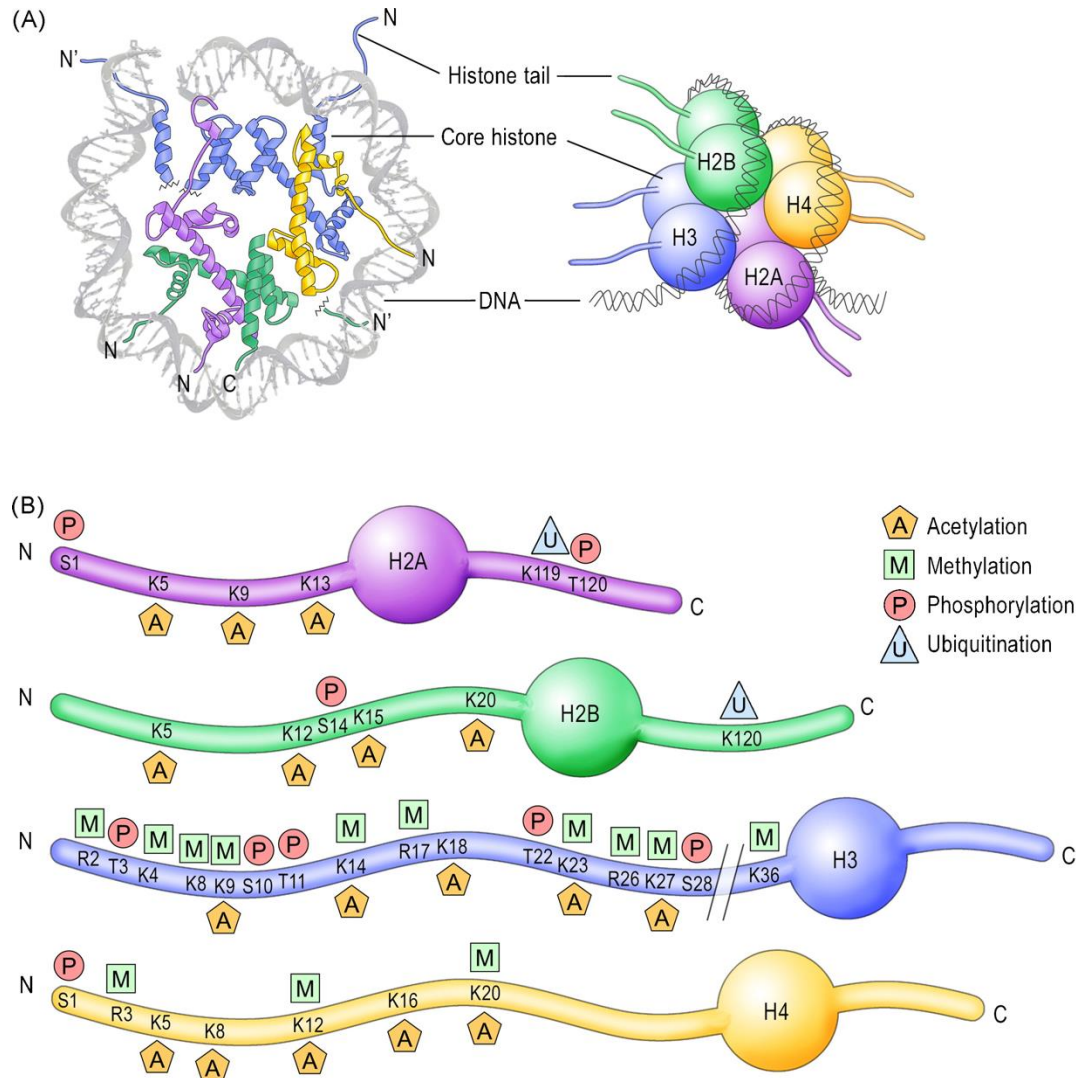
Regulation of chromatin structure:

- 1) Posttranslational histone modifications
- 2) Histone variants
- 3) DNA Methylation

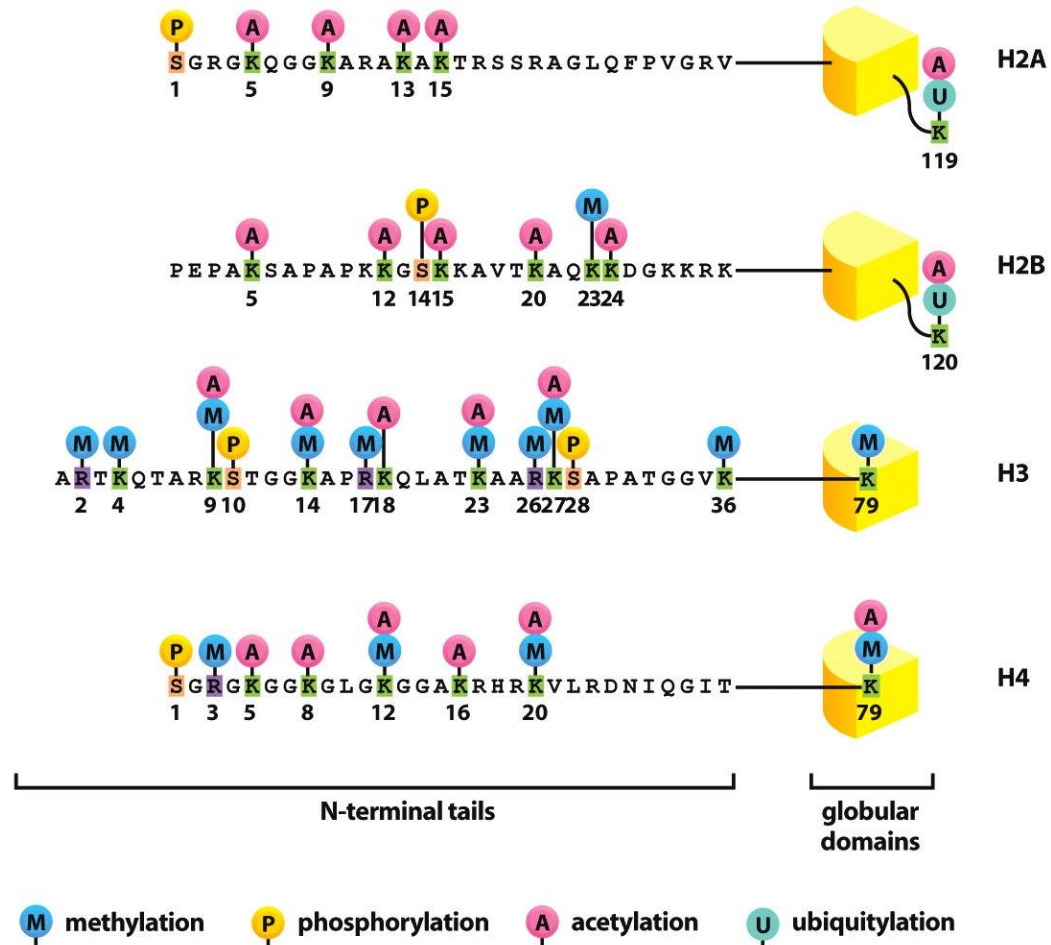
Regulation of chromatin structure:

- 1) **Posttranslational histone modifications**
- 2) Histone variants
- 3) DNA Methylation

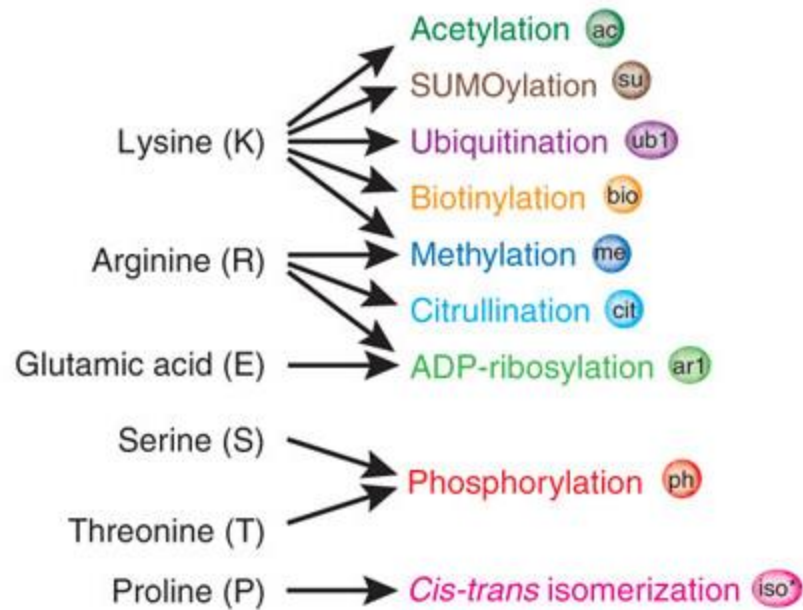
2) Posttranslational modifications (PTMs) of histones



- **Histone PTMs predominantly occur on the N-terminal tail**



- **Amino acids in histones that can be modified:**

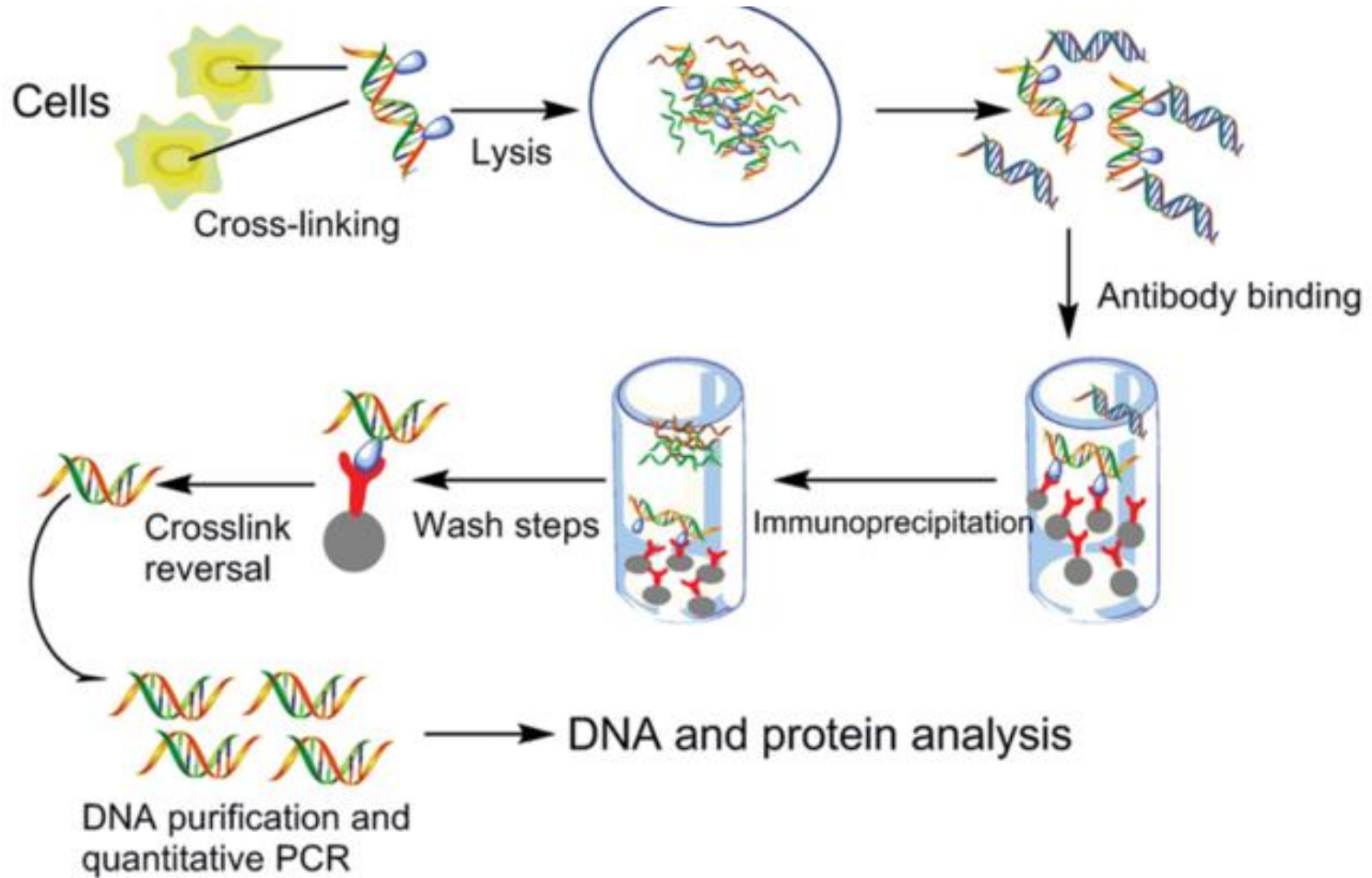


- **2 groups of histone PTMs**

- Group 1: Small chemical groups
- Group 2: Large chemical groups

	Role in transcription	Histone-modified sites
GROUP 1		
Acetylation	activation	H3 (K9,K14,K18,K56) H4 (K5,K8,K12,K16) H2A H2B (K6,K7,K16,K17)
Phosphorylation	activation	H3 (S10)
Methylation	activation	H3 (K4,K36,K79)
	repression	H3 (K9,K27) H4 (K20)
GROUP 2		
Ubiquitylation	activation	H2B (K123)
	repression	H2A (K119)
Sumoylation	repression	H3 (?)
		H4 (K5,K8,K12,K16)
		H2A (K126)
		H2B (K6,K7,K16,K17)

- **How to experimentally measure histone PTMs: Chromatin Immunoprecipitation (ChIP)**



• Epigenetic mechanisms and their effect on transcription

- Mechanisms regulating the compaction of the chromatin
- Regulate **gene transcription** by regulating chromatin compaction
- Different types:

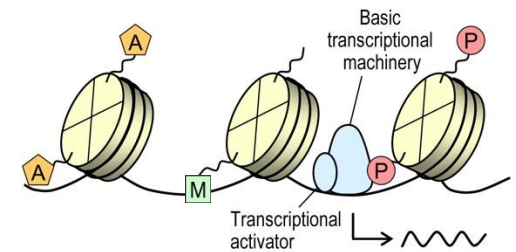
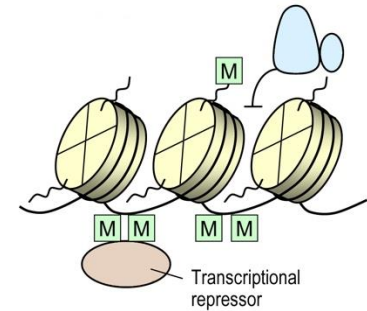
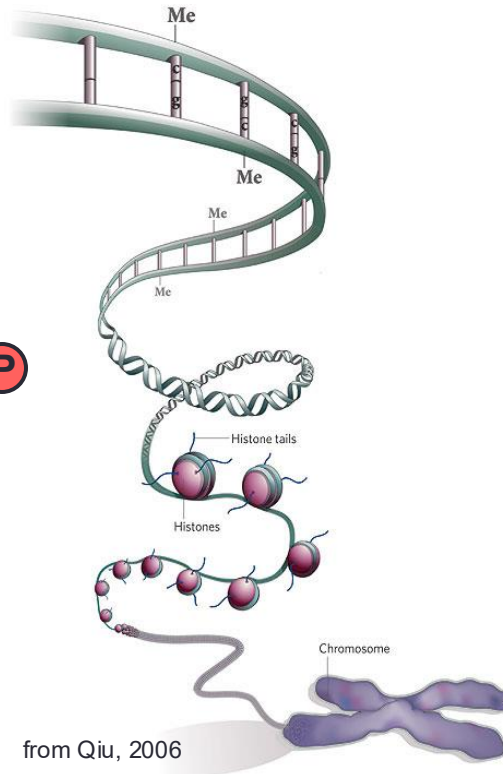
- DNA methylation 

- Histone modifications

- Phosphorylation 

- Methylation 

- Acetylation 



Histone PTMs can occur in a gene's promoter region or in the coding region!

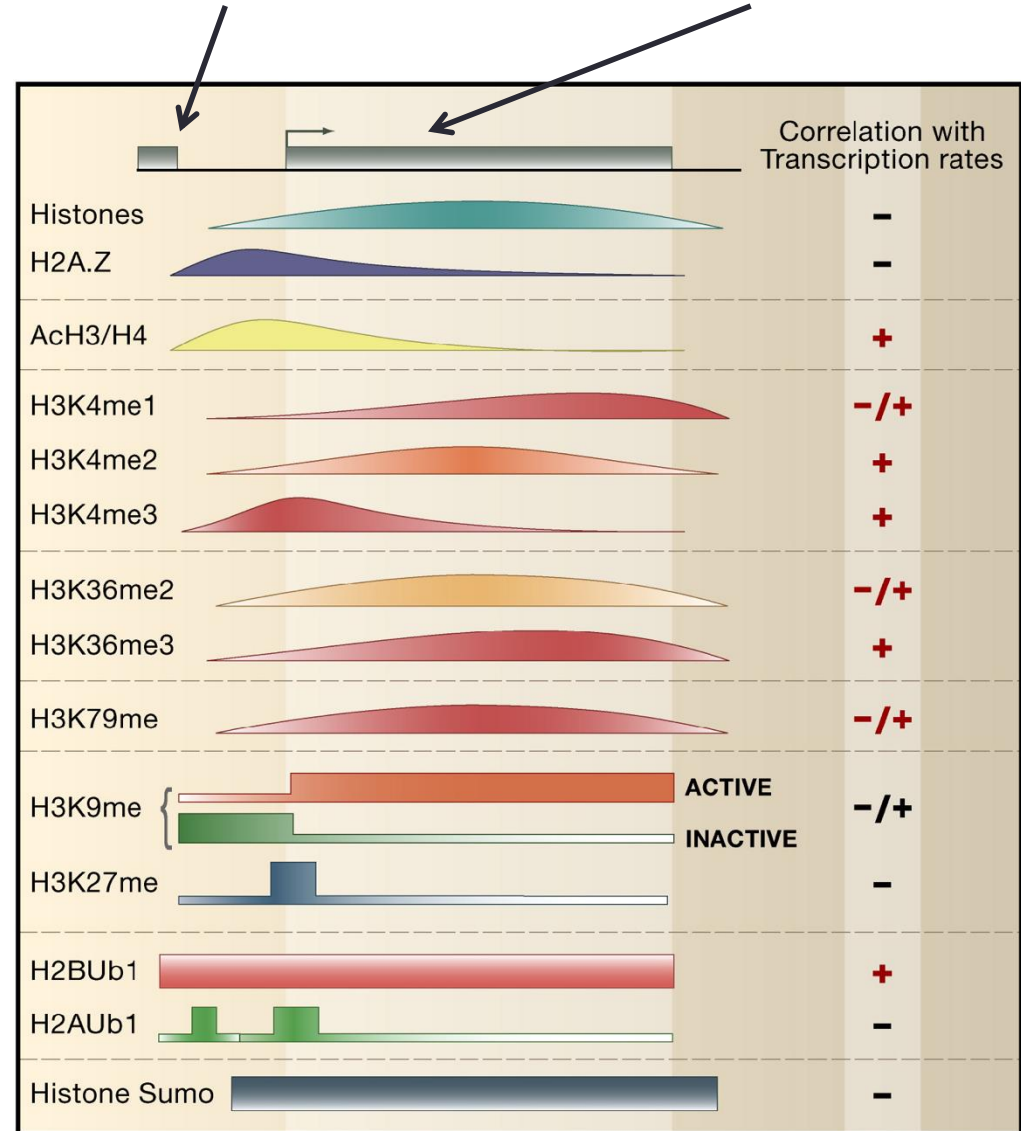
- Effect on transcription**

- Transcription ↗

- Acetylation
- Phosphorylation
- H2B ubiquitination

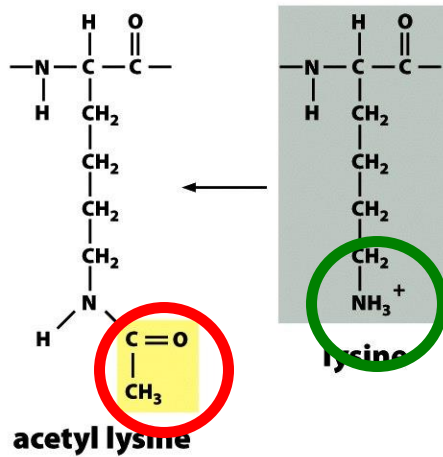
- Transcription ↘

- H2A ubiquitination
- Sumoylation

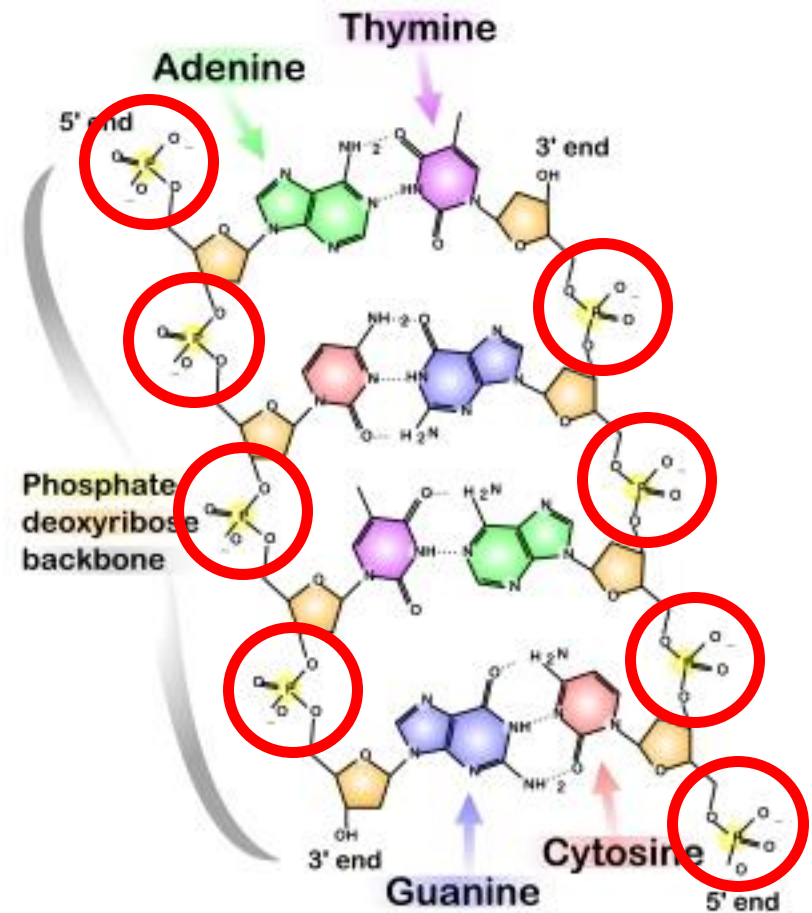
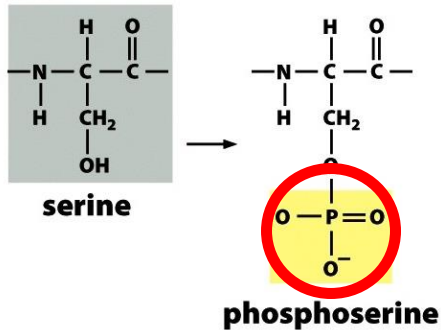


- Effect on transcription – acetylation and phosphorylation

(A) LYSINE ACETYLATION

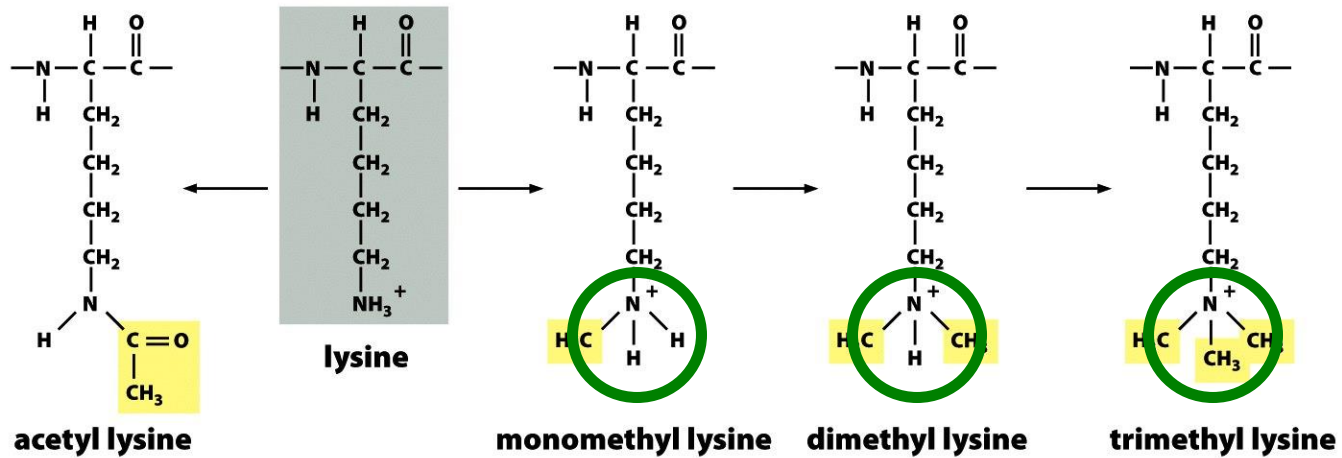


(B) SERINE PHOSPHORYLATION

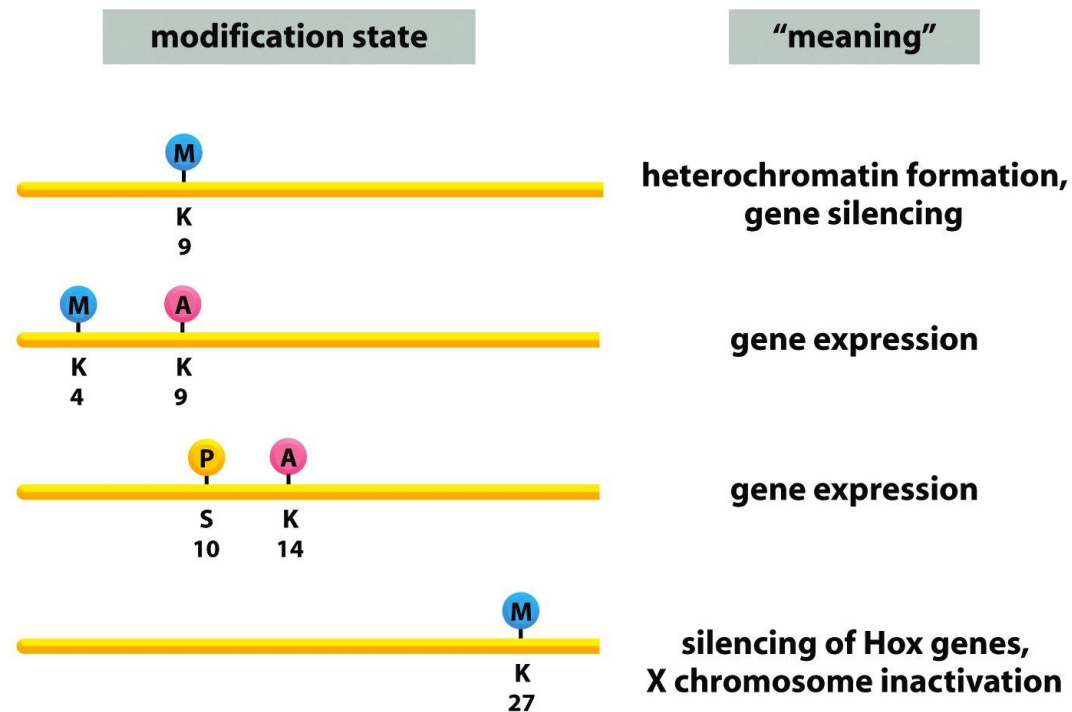


- Effect on transcription – acetylation and methylation

(A) LYSINE ACETYLATION AND METHYLATION ARE COMPETING REACTIONS



- **Co-occurrence of histone PTMs can influence the rate of gene transcription**



- **Enzymes regulating histone PTMs:**

- Acetylation

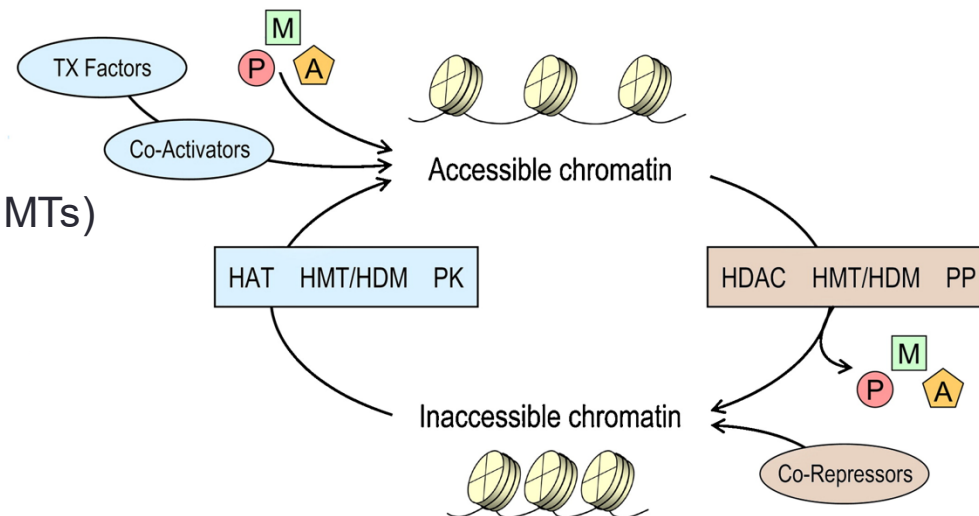
- Histone acetyl transferases (HATs)
- Histone deacetylases (HDACs)

- Methylation

- Histone methyl transferases (HMTs)
- Histone demethylases (HDMs)

- Phosphorylation

- Protein kinases (PKs)
- Protein phosphatases (PPs)



- **New histone PTMs keep being discovered**

LETTER

<https://doi.org/10.1038/s41586-019-1024-7>

Histone serotonylation is a permissive modification that enhances TFIID binding to H3K4me3

Lorna A. Farrelly¹, Robert E. Thompson², Shuai Zhao^{3,4}, Ashley E. Lepack¹, Yang Lyu¹, Natarajan V. Bhanu⁵, Baichao Zhang^{3,4}, Yong-Hwee E. Loh¹, Arathi Ramakrishnan¹, Krishna C. Vadodaria⁶, Kelly J. Heard⁶, Galina Erikson⁶, Tomoyoshi Nakadaï⁷, Ryan M. Bastle¹, Bradley J. Lukasak², Henry Zebroski III⁸, Natalia Alenina⁹, Michael Bader⁹, Olivier Berton¹, Robert G. Roeder⁷, Henrik Molina⁸, Fred H. Gage⁶, Li Shen¹, Benjamin A. Garcia⁵, Haitao Li^{3,4}, Tom W. Muir² & Ian Maze^{1,10*}

28 MARCH 2019 | VOL 567 | NATURE | 535

Article

Metabolic regulation of gene expression by histone lactylation

<https://doi.org/10.1038/s41586-019-1678-1>

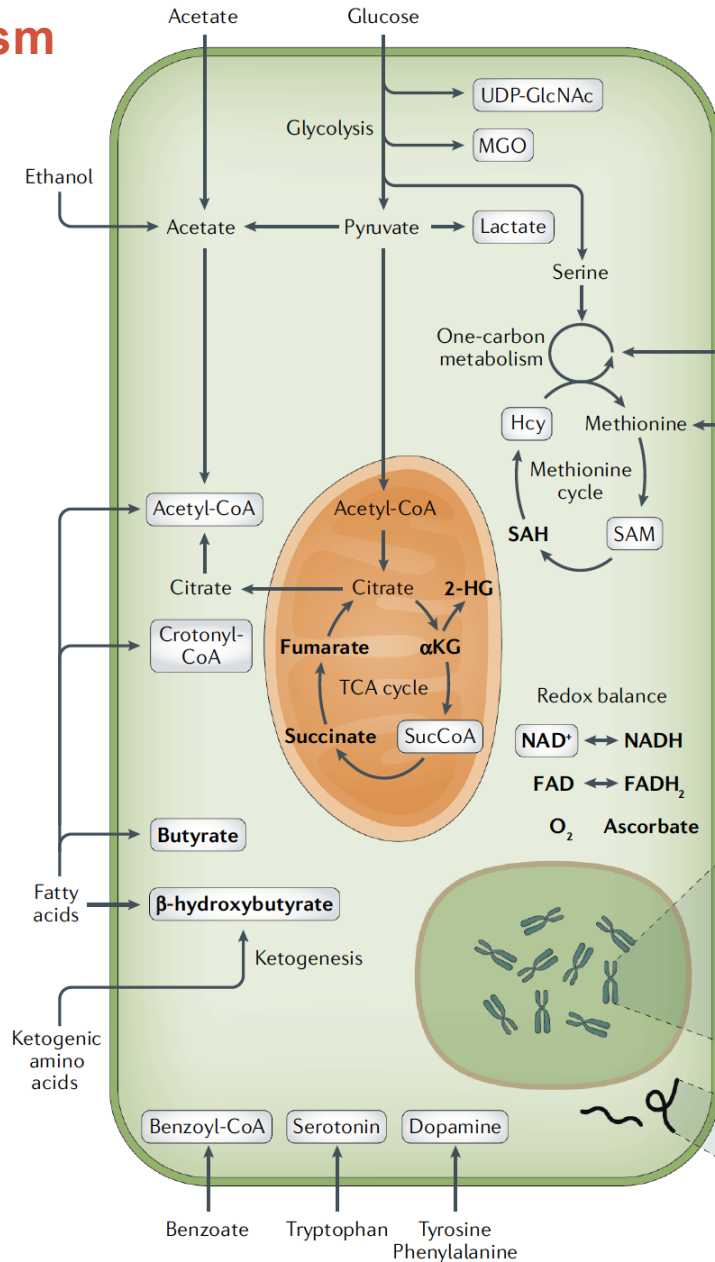
Received: 21 June 2018

Accepted: 13 September 2019

Published online: 23 October 2019

Di Zhang^{1,12}, Zhanyun Tang^{2,12}, He Huang^{1,9}, Guolin Zhou¹, Chang Cui¹, Yejing Weng¹, Wenchao Liu¹, Sunjoo Kim³, Sangkyu Lee³, Mathew Perez-Neut¹, Jun Ding¹, Daniel Czyz⁴, Rong Hu^{5,6}, Zhen Ye^{5,6}, Maomao He⁷, Y. George Zheng⁷, Howard A. Shuman⁴, Lunzhi Dai^{1,10}, Bing Ren^{5,6}, Robert G. Roeder², Lev Becker^{1,8,11*} & Yingming Zhao^{1,8*}

Metabolism



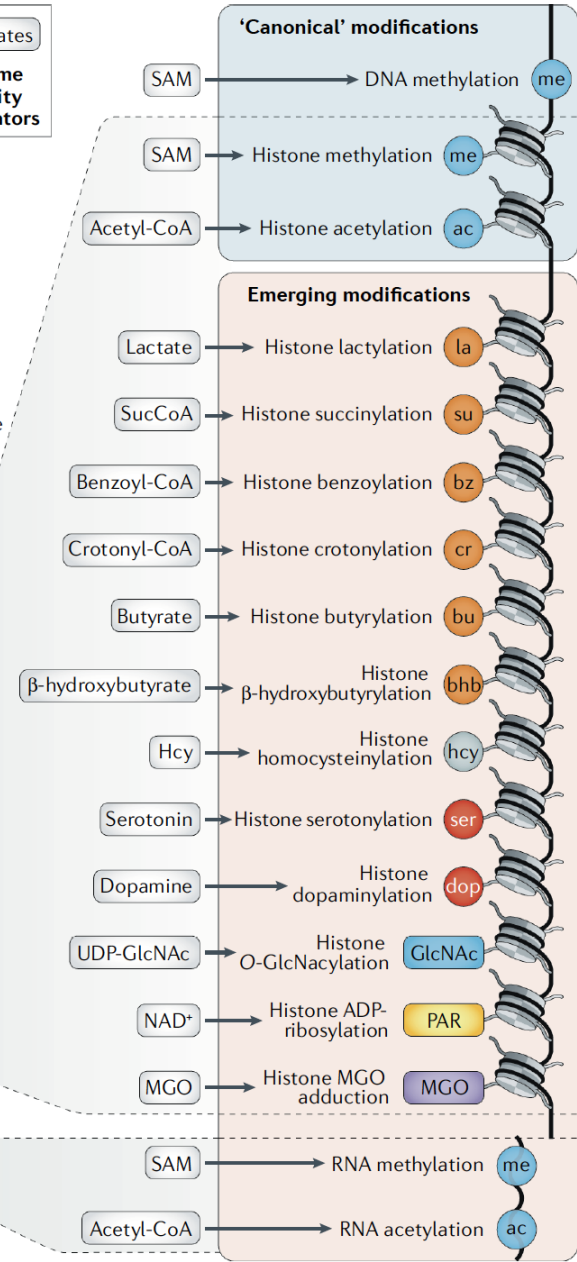
Substrates
Enzyme activity modulators

Folate
Vitamin B₁₂
Choline
Serine
Glycine
Threonine

Methionine cycle
Methionine
Hcy
SAM
SAH

Redox balance
 $NAD^+ \leftrightarrow NADH$
 $FAD \leftrightarrow FADH_2$
 O_2 Ascorbate

Histone PTMs



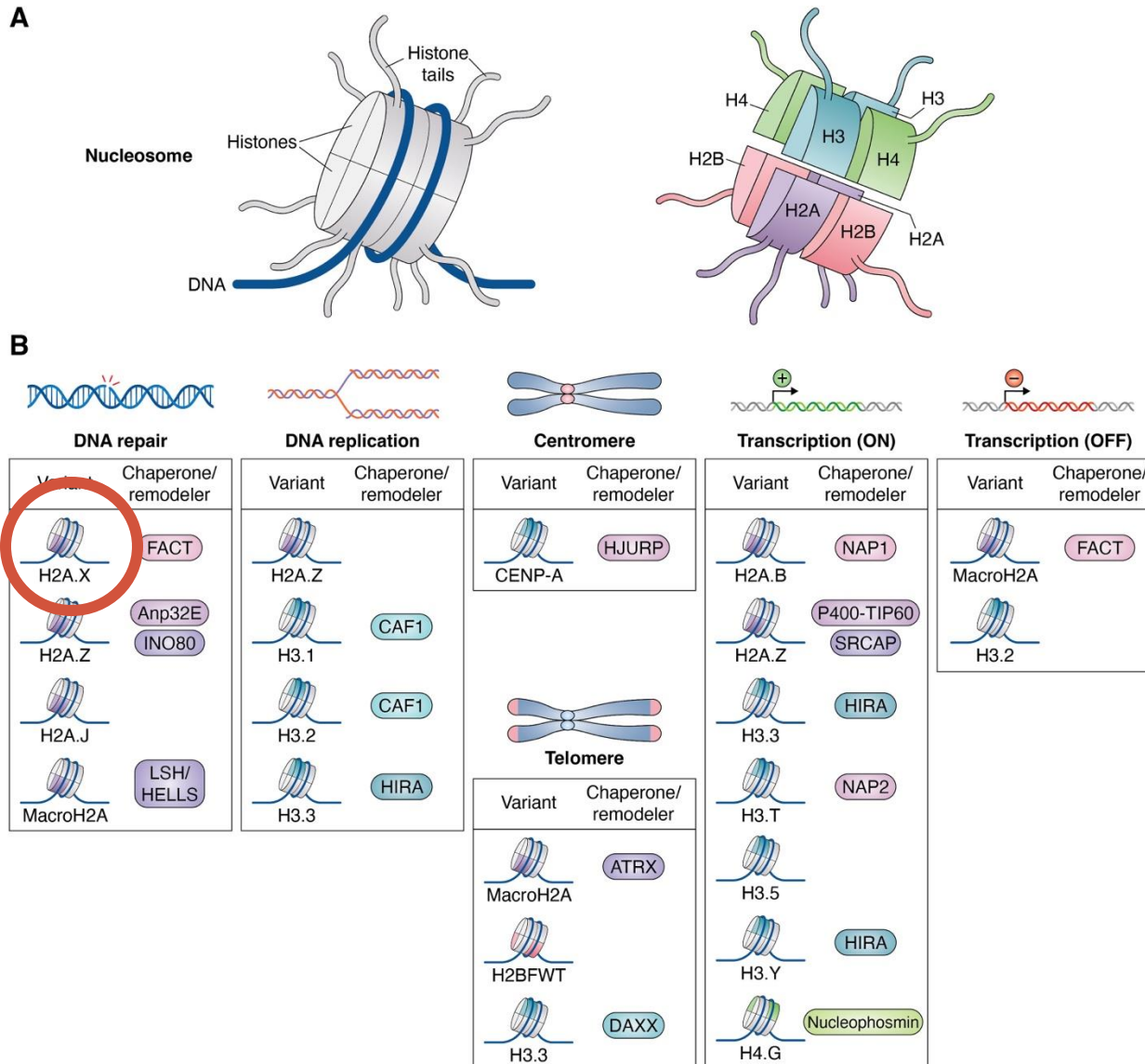
Regulation of chromatin structure:

- 1) Posttranslational histone modifications
- 2) **Histone variants**
- 3) DNA Methylation

Histone variants:

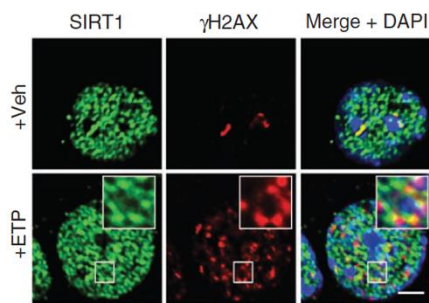
- Structural variants of the core histones
- Specialized function
- Inserted by histone-exchange process

- Histone variants – structure and function**



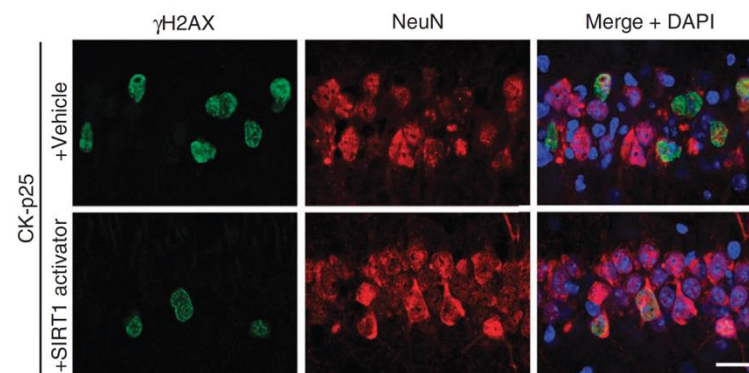
- Histone variants – structure and function**

- H2AX (aka γ H2AX), a marker of DNA damage and repair

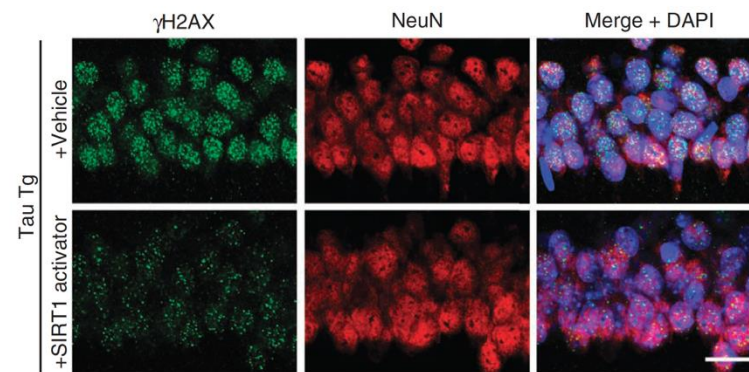


ETP, etoposide, induces DNA double strand breaks

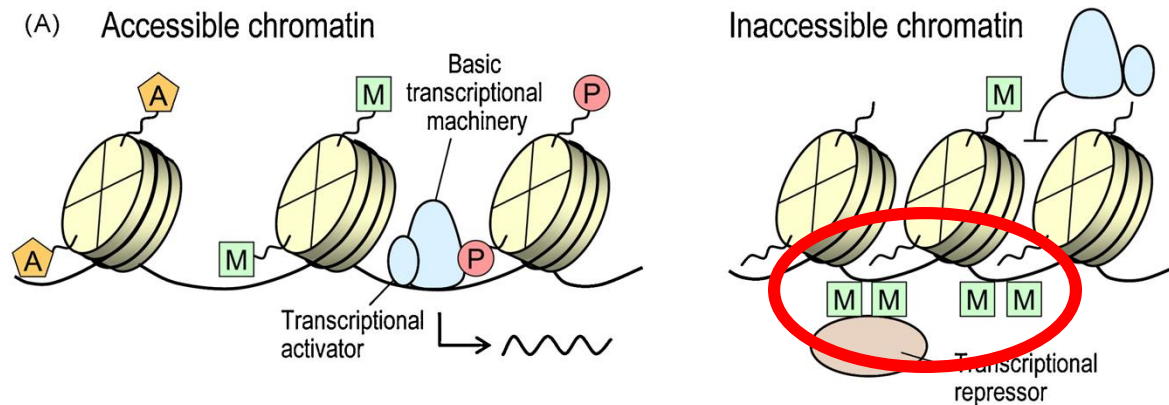
A mouse model of AD (amyloid)



A mouse model of AD (tau)



- **A histone code regulating the accessibility of the chromatin for transcription:**



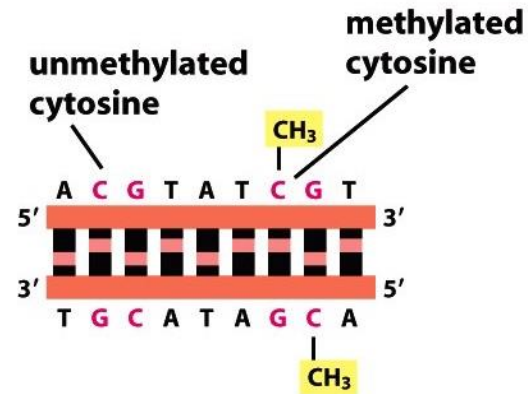
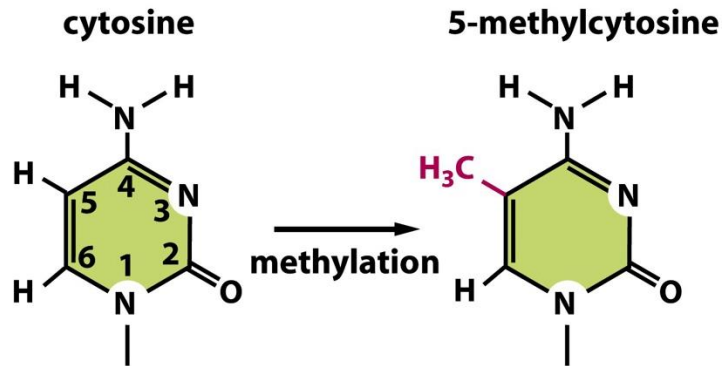
- **Together with DNA methylation, this forms an epigenetic code regulating the accessibility of the chromatin for transcription**

Regulation of chromatin structure:

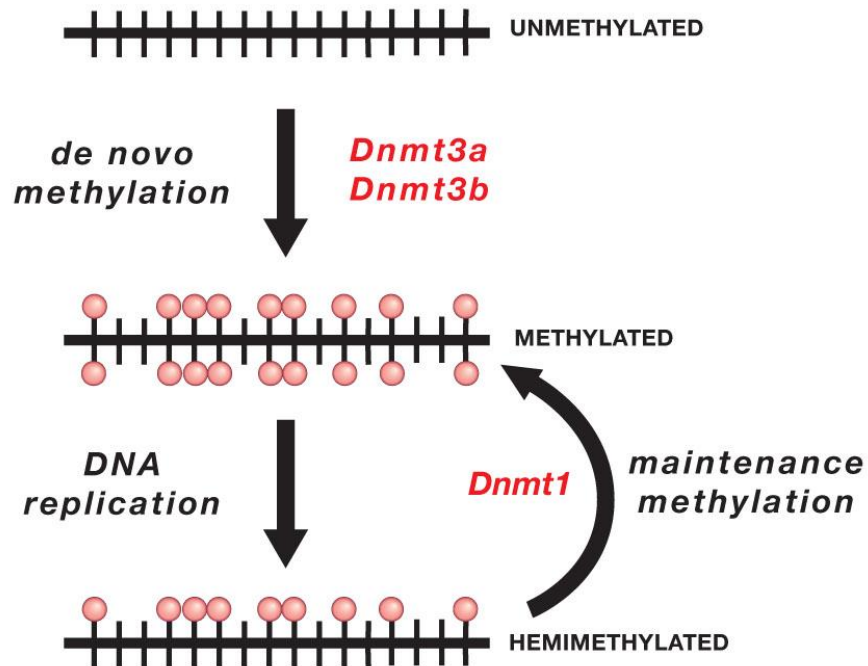
- 1) Posttranslational histone modifications
- 2) Histone variants
- 3) **DNA Methylation**

DNA methylation

- On cytosines
- ... followed by guanine
 - “CG island”
 - “CpG island”



- **2 types of DNA methylation-inducing enzymes**
 - De novo DNA methyltransferases
 - Maintenance DNA methyltransferases

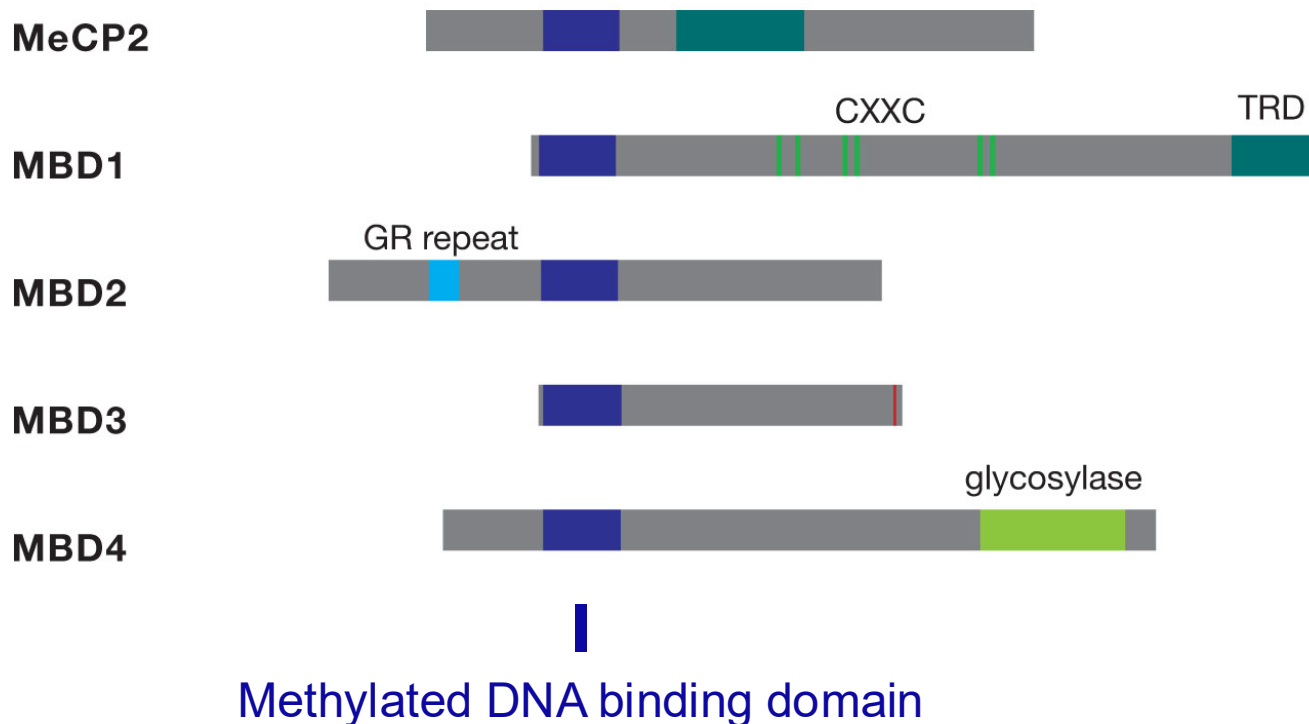


- **2 types of DNA methylation-inducing enzymes**

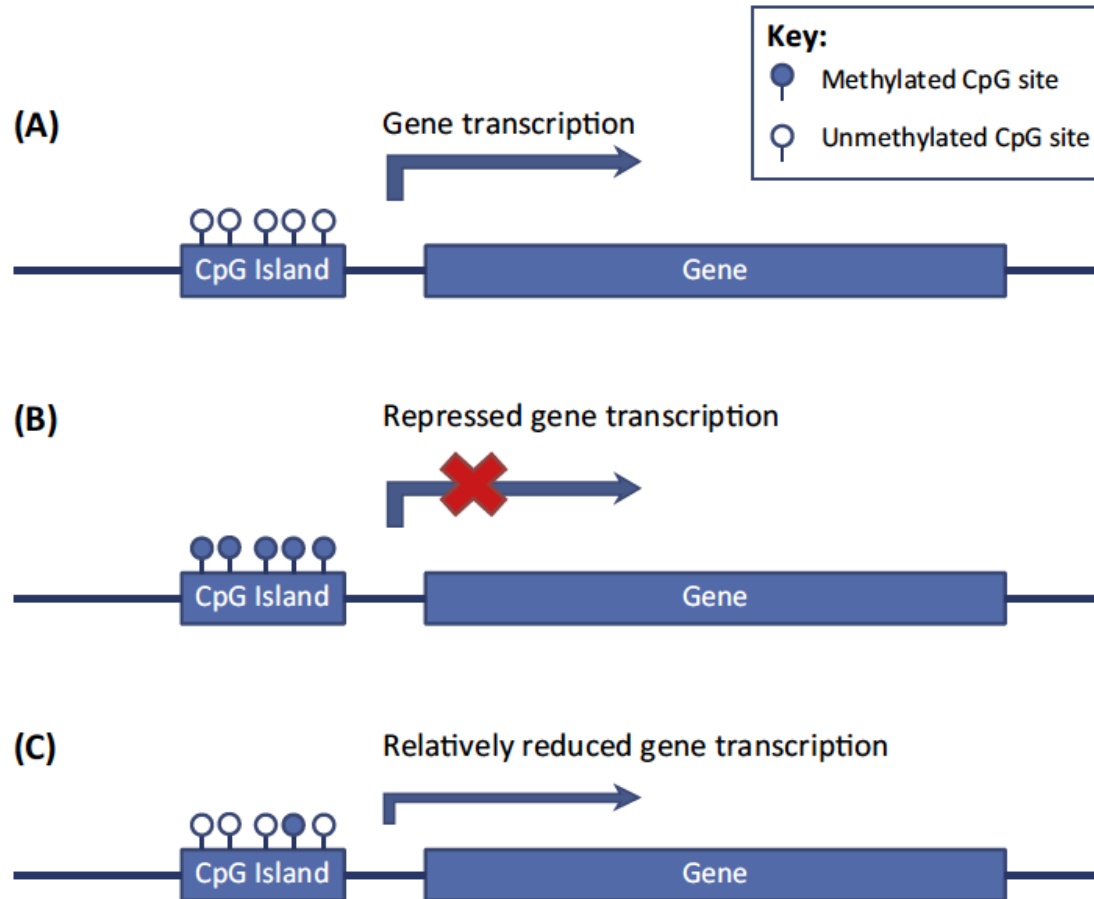
Table 1. Functions of mammalian DNA methyltransferases

DNA methyltransferase	Species	Major activity	Major phenotypes of loss-of-function mutations
Dnmt1	mouse	maintenance methylation of CpG	genome-wide loss of DNA methylation, embryonic lethality at embryonic day 9.5 (E9.5), abnormal expression of imprinted genes, ectopic X-chromosome inactivation, activation of silent retrotransposon
Dnmt2	mouse	weak activity	no change in CpG methylation, no obvious developmental phenotypes
Dnmt3a	mouse	de novo methylation of CpG	postnatal lethality at 4–8 weeks, male sterility, and failure to establish methylation imprints in both male and female germ cells
Dnmt3b	mouse	de novo methylation of CpG	demethylation of minor satellite DNA, embryonic lethality around E14.5 days with vascular and liver defects (embryos lacking both Dnmt3a and Dnmt3b fail to initiate de novo methylation after implantation and die at E9.5)
DNMT3B	human	de novo methylation of CpG	ICF syndrome: immunodeficiency, centromeric instability, and facial anomalies; loss of methylation in repetitive elements and pericentromeric heterochromatin

- Methylated portions of the DNA attract **Methyl-Binding-Domain** containing Proteins (MBD), which act as co-activators or co-repressors

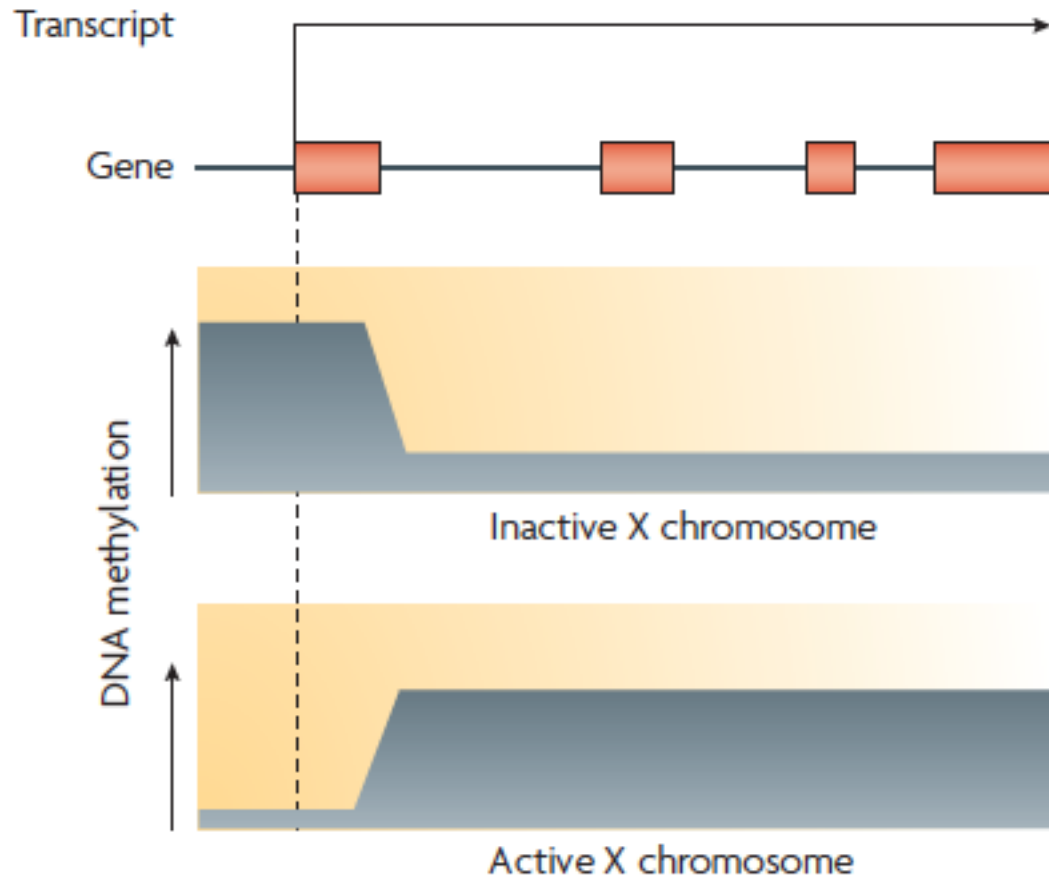


- **Function of DNA methylation: Repression (in general)**



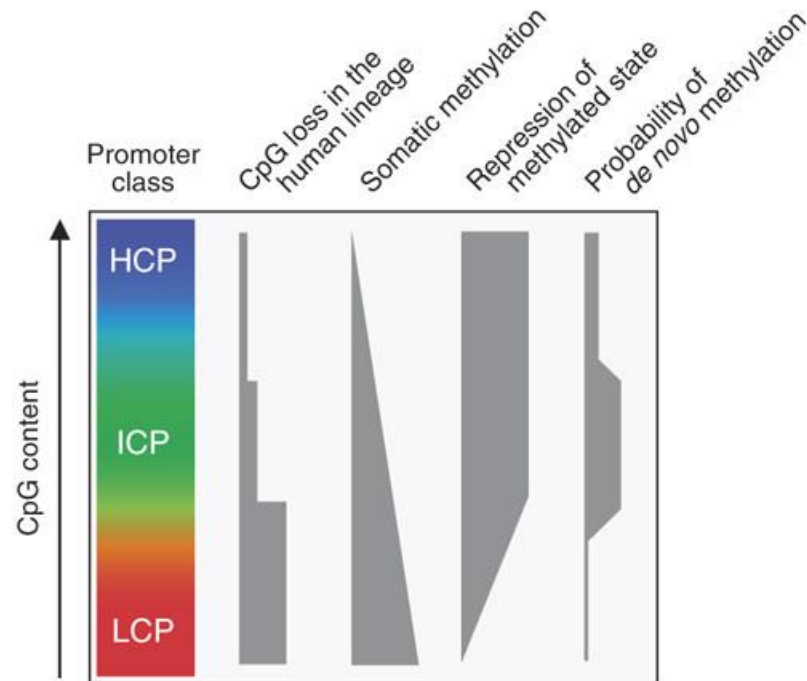
- **Function of DNA methylation: Repression or Activation**

- Depends on the genetic context



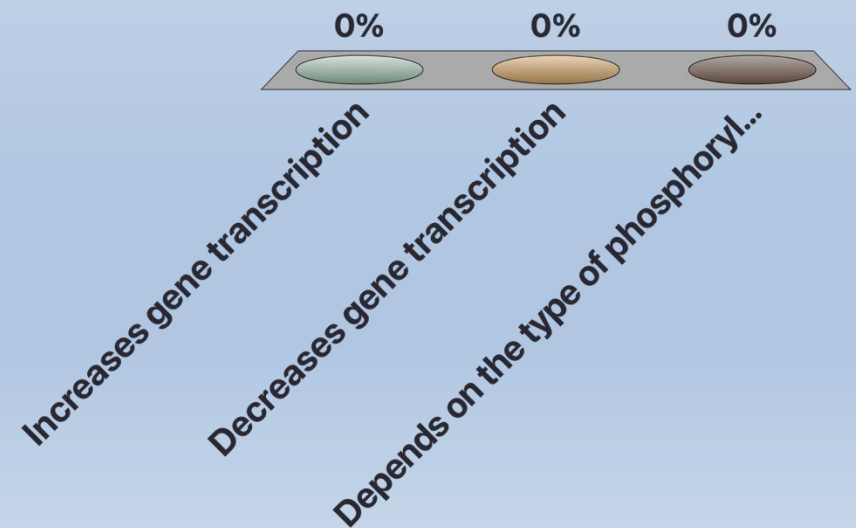
- **Function of DNA methylation: Repression or Activation**

- Depends on the CpG content



What is the effect of histone phosphorylation on transcription?

- A. Increases gene transcription
- B. Decreases gene transcription
- C. Depends on the type of phosphorylation



The transcriptional effect of DNA methylation depends on...

Neuroepigenetics

1) The chromatin – Epigenetic basics (Lecture 1)

- Chromatin condensation
- Regulation of chromatin structure
- **Epigenetic inheritance**
- Environmental influence on epigenetics

2) Epigenetic dysregulation (Lecture 2)

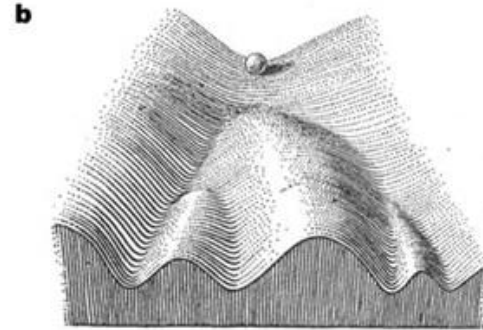
- in AD

Epigenetic modifications can be...

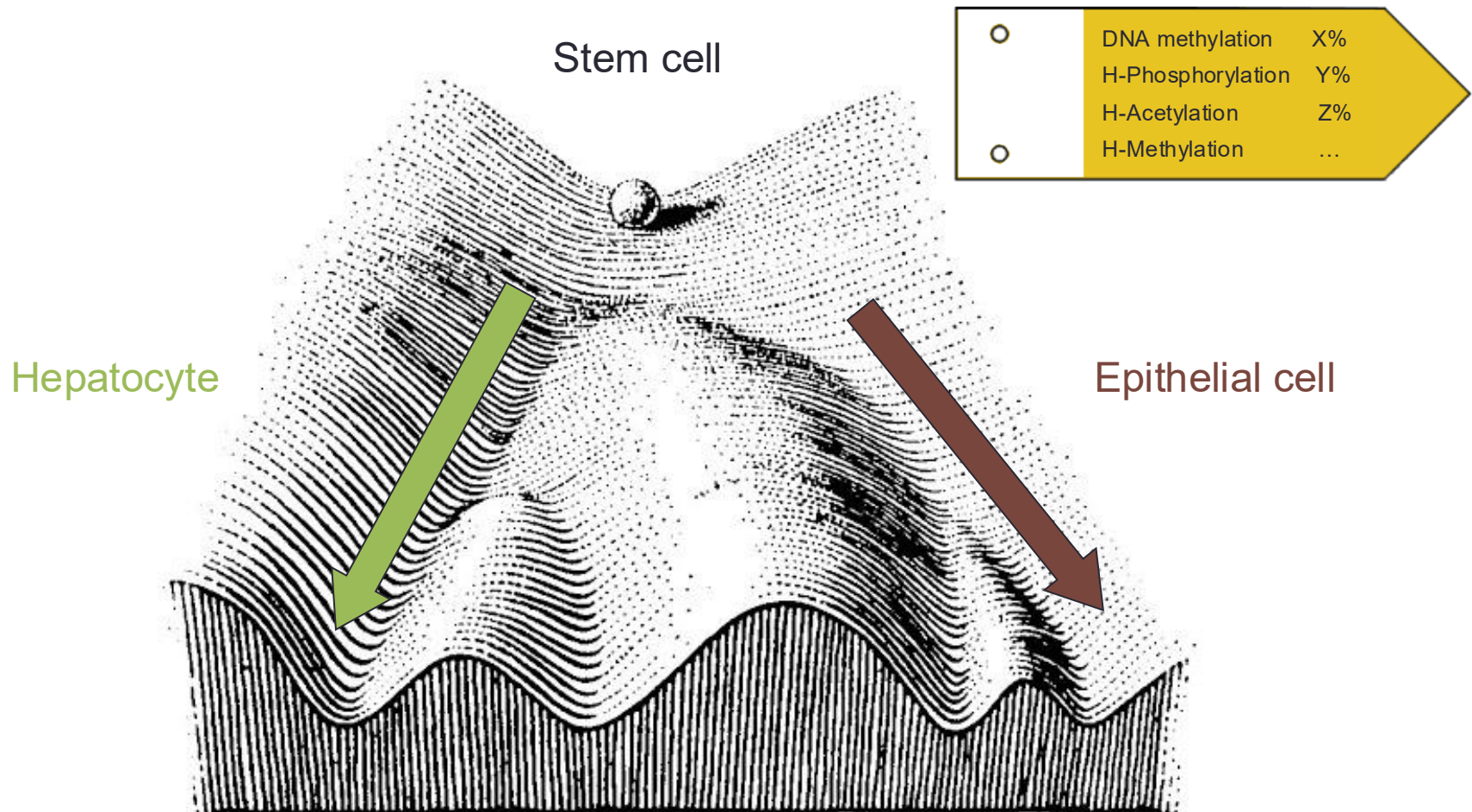
- **inherited** through mitosis
 - development/differentiation
 - **inherited** through meiosis
 - transgenerational epigenetic inheritance
- In both cases, they can be influenced/installed by the environment

1) Epigenetic inheritance during development

- The developmental epigenetic landscape *sensu* Waddington (1905-1975)



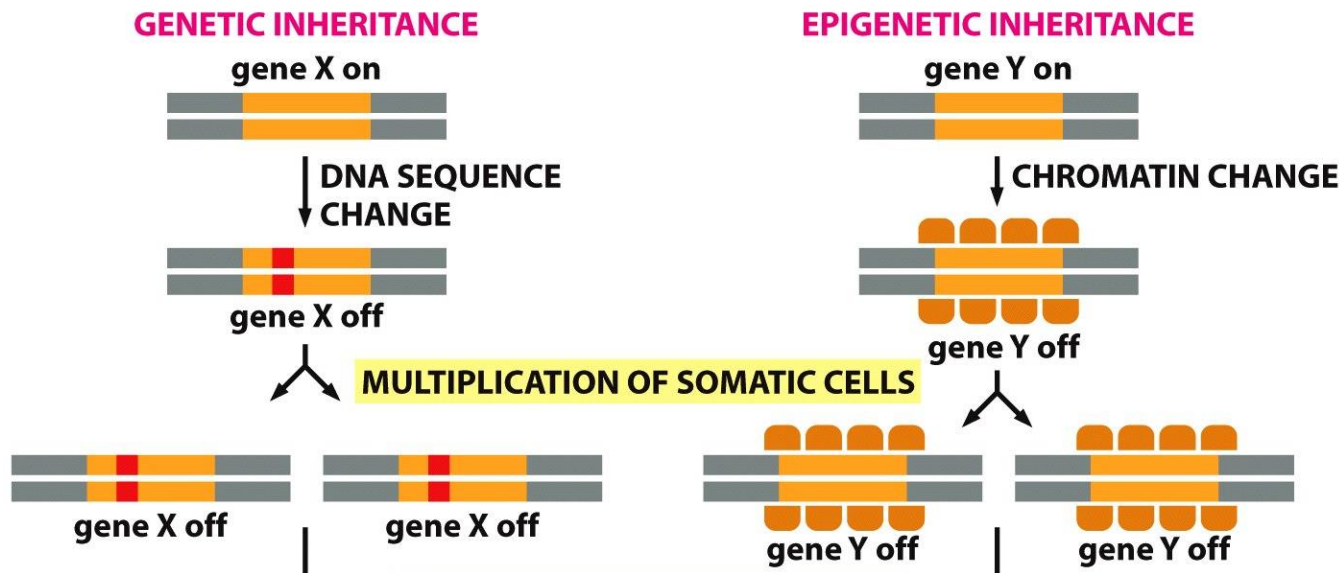
1) Epigenetic inheritance during development



Waddington's epigenetic landscape (1957):
The trajectory paths (the valleys) a cell can take
are defined by epigenetic borders (the hills)

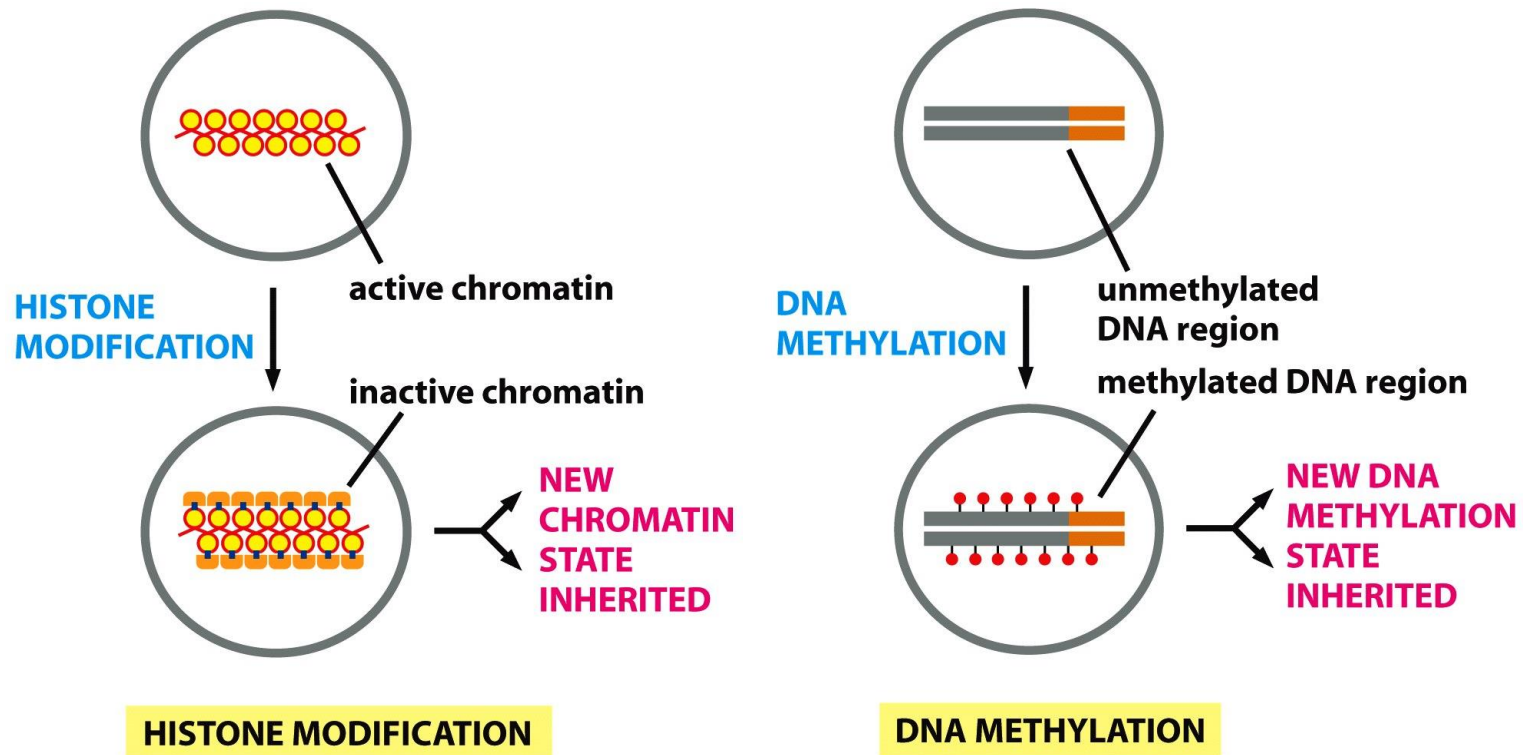
1) Epigenetic inheritance during development

- Epigenetic vs genetic inheritance



On a locus-specific level

- Both histone modifications and DNA methylation changes can be inherited

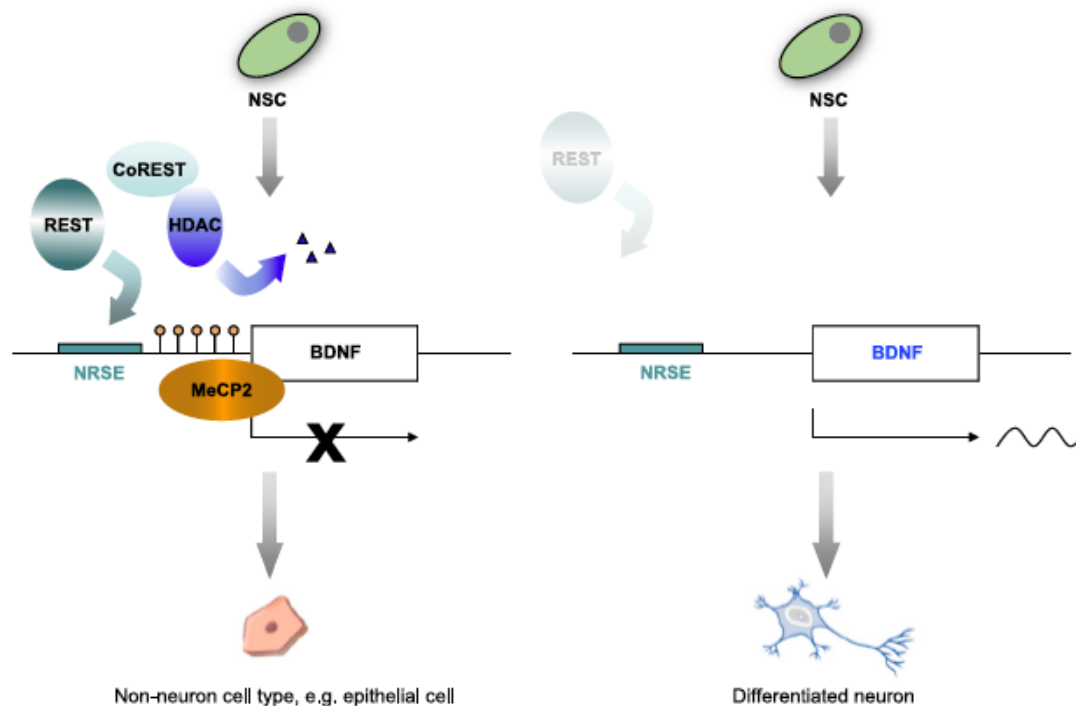


On a locus-specific level

- Differentiation of neurons during development

NSC=Neural stem cell; NRSE=Neuron restrictive silencer element; REST=Repressor; BDNF=brain derived neurotrophic factor

REST is a neuron-specific repressor

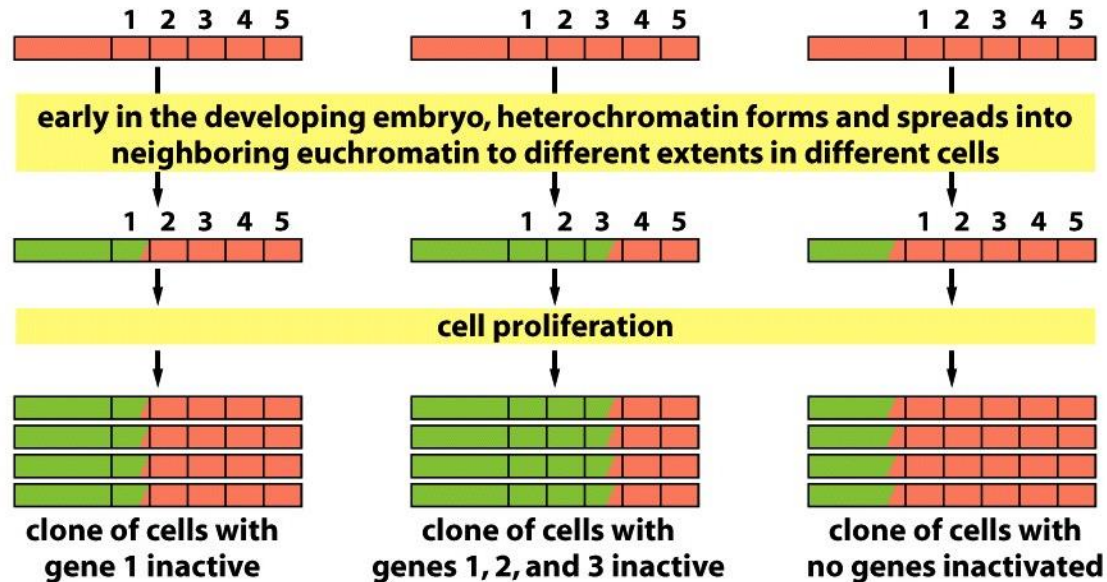


On a “regional” level:

transcriptionally silent

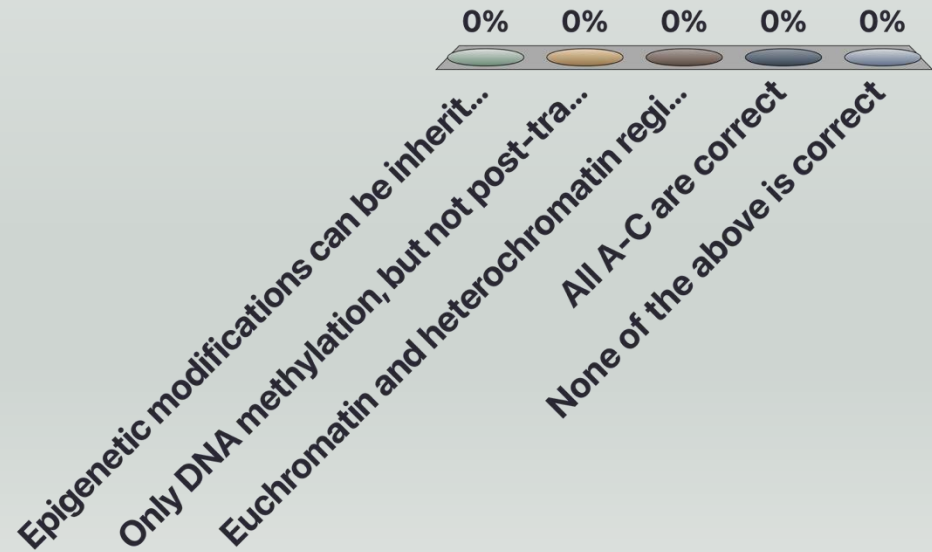
transcriptionally active

- Distinction between **hetero-** and **euchromatin** is inherited through development



Which of the following is correct?

- A. Epigenetic modifications can be inherited through mitosis, but not meiosis
- B. Only DNA methylation, but not post-translational histone modifications can be inherited through mitosis
- C. Euchromatin and heterochromatin regions cannot be changed during development
- D. All A-C are correct
- E. None of the above is correct



Neuroepigenetics

1) The chromatin – Epigenetic basics (Lecture 1)

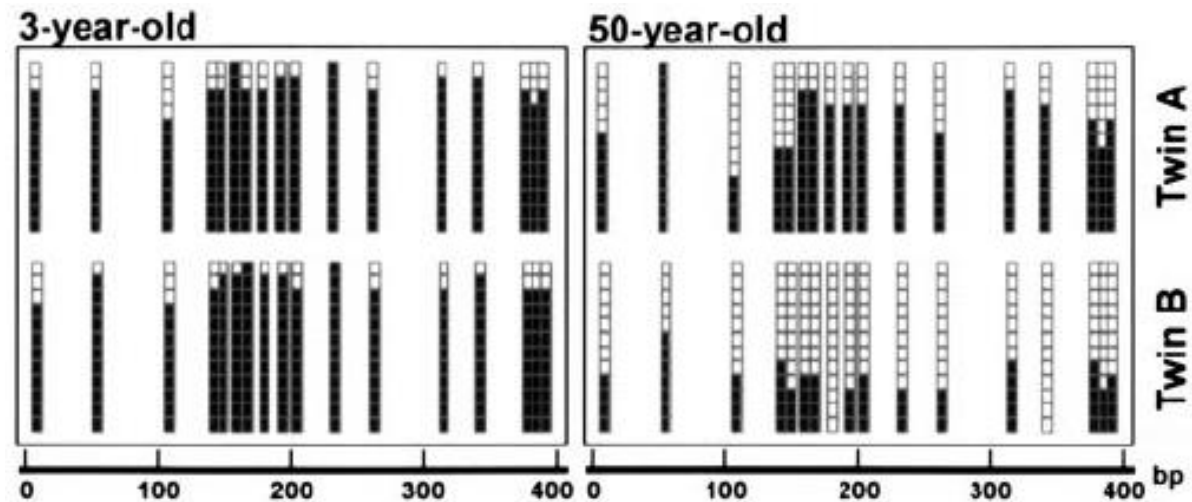
- Chromatin condensation
- Regulation of chromatin structure
- Epigenetic inheritance
- **Environmental influence on epigenetics**

2) Epigenetic dysregulation (Lecture 2)

- in AD

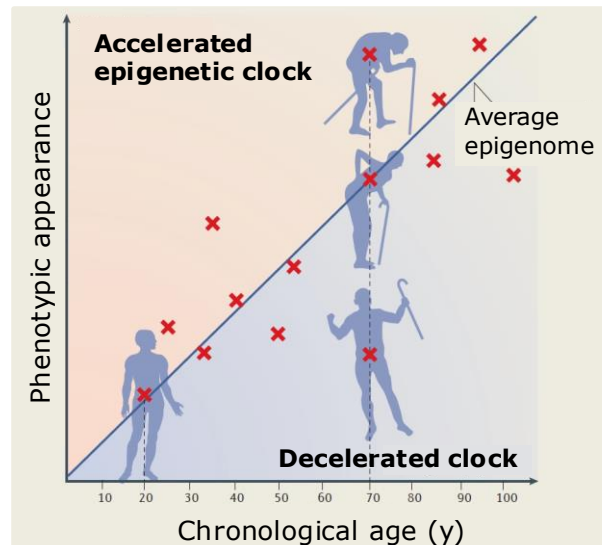
1) Epigenetic inheritance during development

- Influence of the environment I/II – the case of monozygotic twins



1) Epigenetic inheritance during development

- Influence on epigenetic clocks



1) **Epigenetic dysregulation following stress**

- Influence of the environment II/II – The case of early-life stress

2) **Epigenetic inheritance during development can be influenced by the environment**

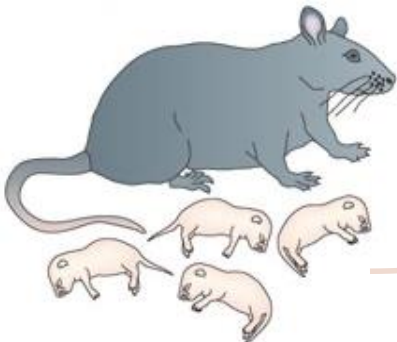
- Maternal care

“Good mothers”



stress resistant
low levels of anxiety
normal behavior

“Bad mothers”



stress susceptible
high levels of anxiety
depressive-like behavior

2) Epigenetic inheritance during development can be influenced by the environment

- Maternal care

“Good mothers”



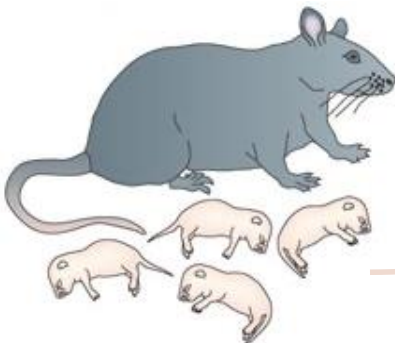
Reduced DNA methylation

Nr3c1 (coding for GR) can be expressed



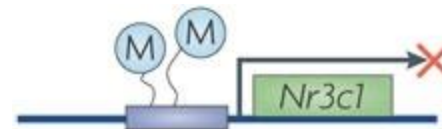
Normal stress response

“Bad mothers”



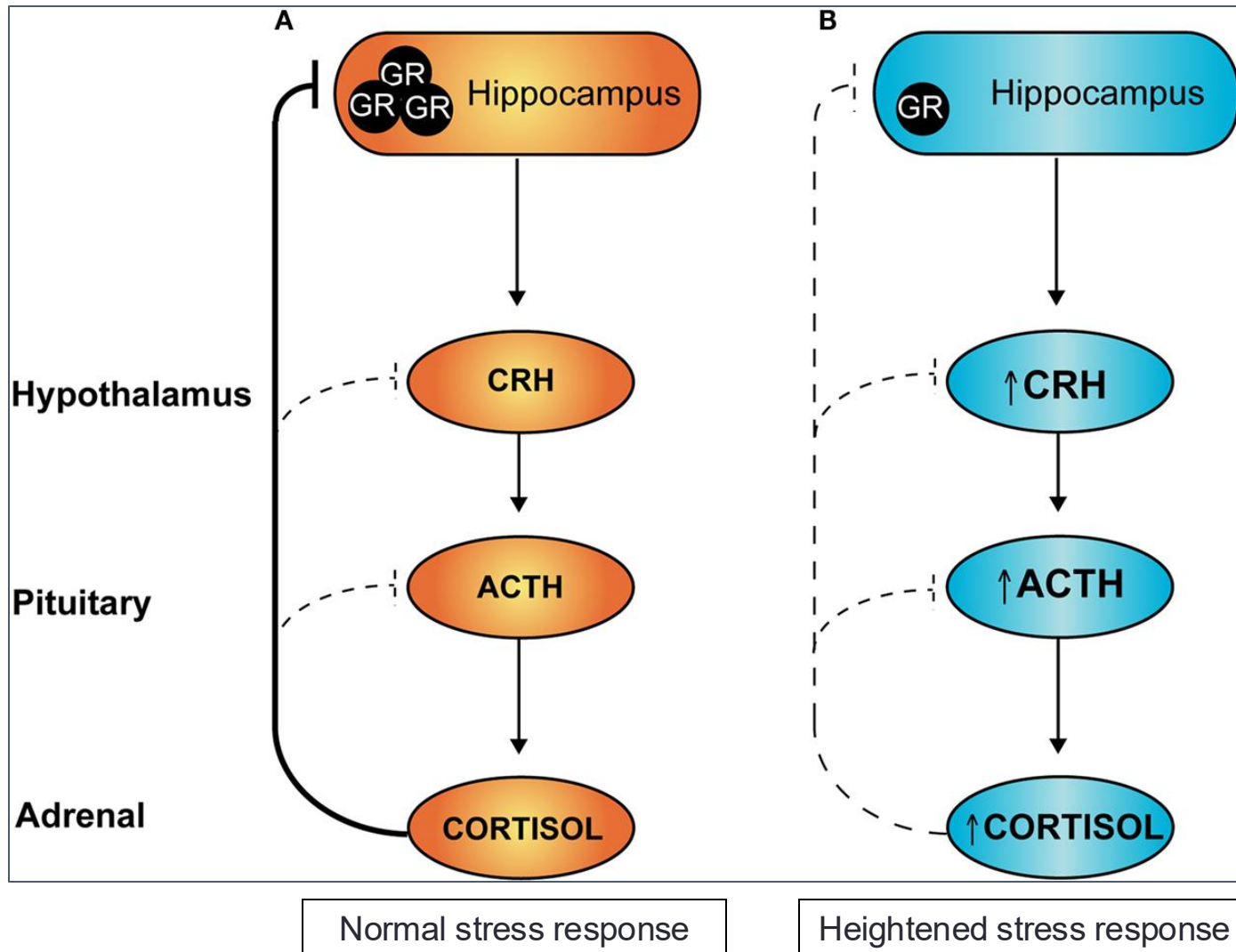
Increased DNA methylation

Nr3c1 is not expressed

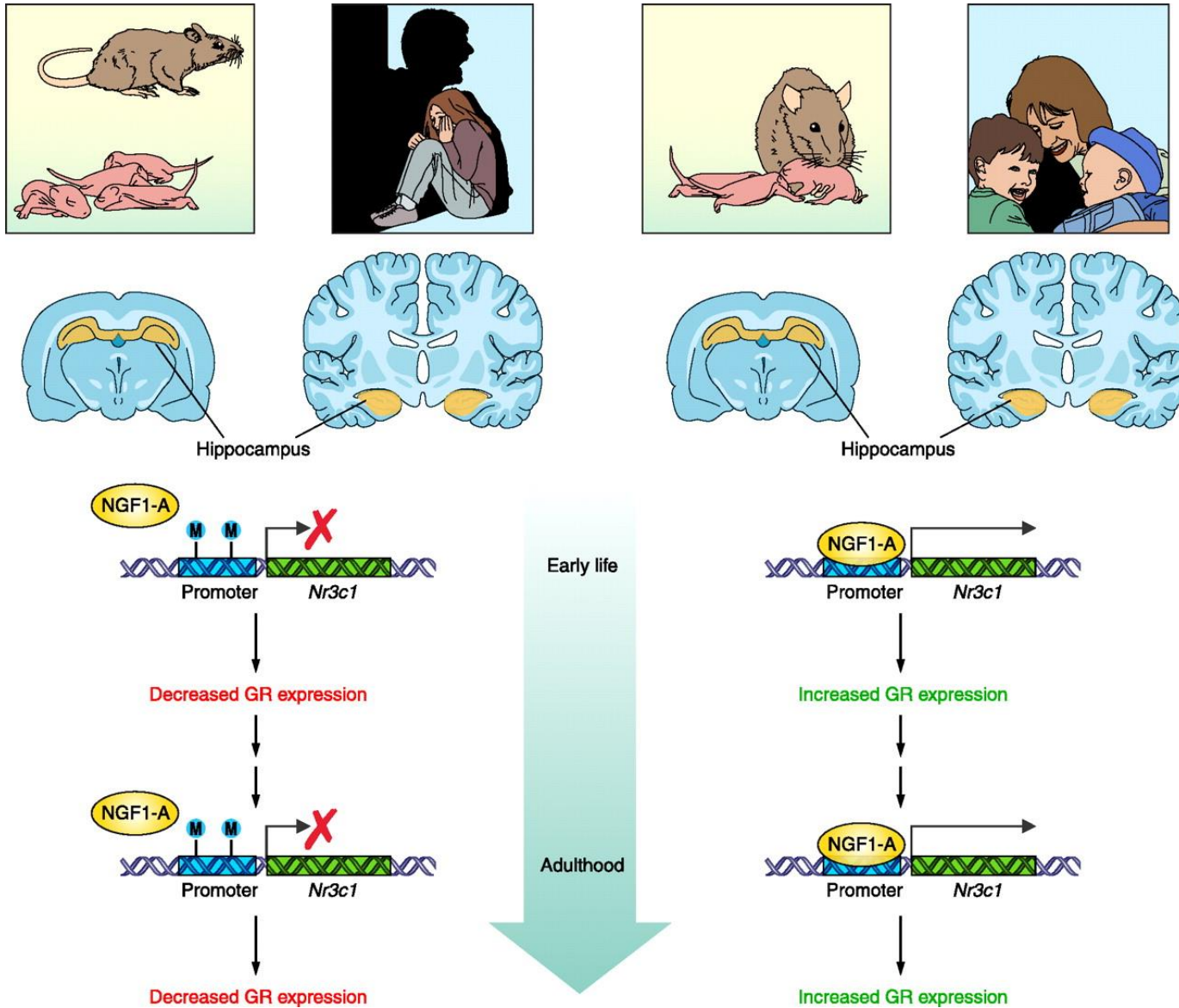


Elevated stress response

Glucocorticoids, glucocorticoid receptors and the stress response



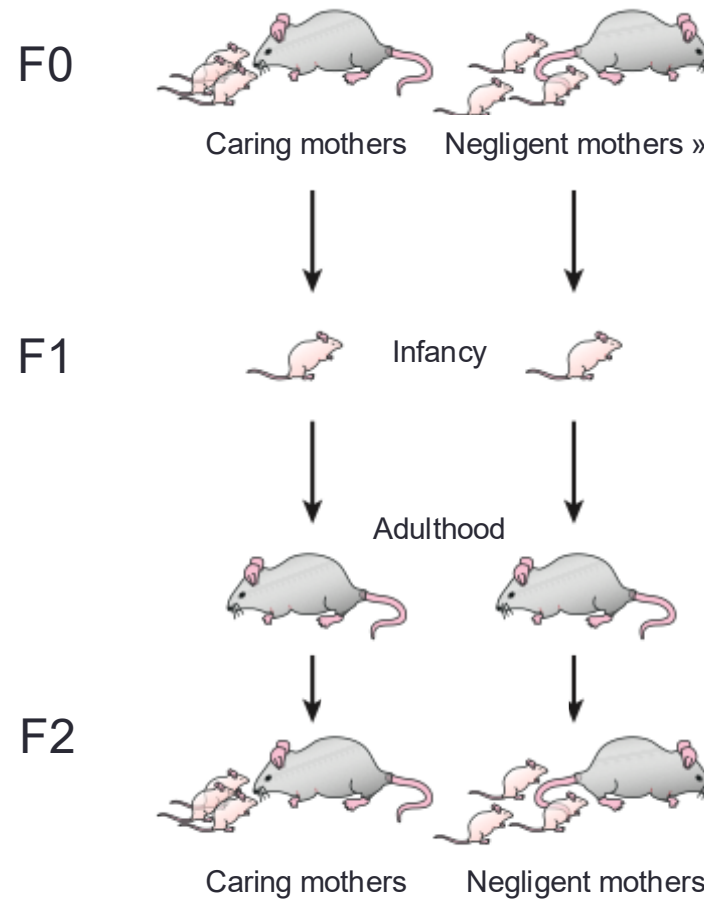
Epigenetic Inheritance



Nr3c1, the gene coding for glucocorticoid receptor 1 (GR); NGF1-A, a transcription factor

2) Intergenerational epigenetic inheritance:

- The quality of maternal care received is passed on...



A word cloud of biological and genetic terms. The words are arranged in a roughly triangular shape, pointing to the right. The colors range from dark brown to light tan. The terms include: Promoter, MBD, Histone, Phosphorylation, Barrier, Nucleosome, Insulator, Chromatin, Acetylation, Methylation, Ubiquitination, Intergenerational, H1, Housekeeping, CpG, Transgenerational, DNA, Euchromatin, Heterochromatin, Nr3c1, and H2AX.

Promoter
MBD
Histone
Phosphorylation
Barrier
Nucleosome
Insulator
Chromatin
Acetylation
Methylation
Ubiquitination
Intergenerational
H1
Housekeeping
CpG
Transgenerational
DNA
Euchromatin
Heterochromatin
Nr3c1
H2AX