

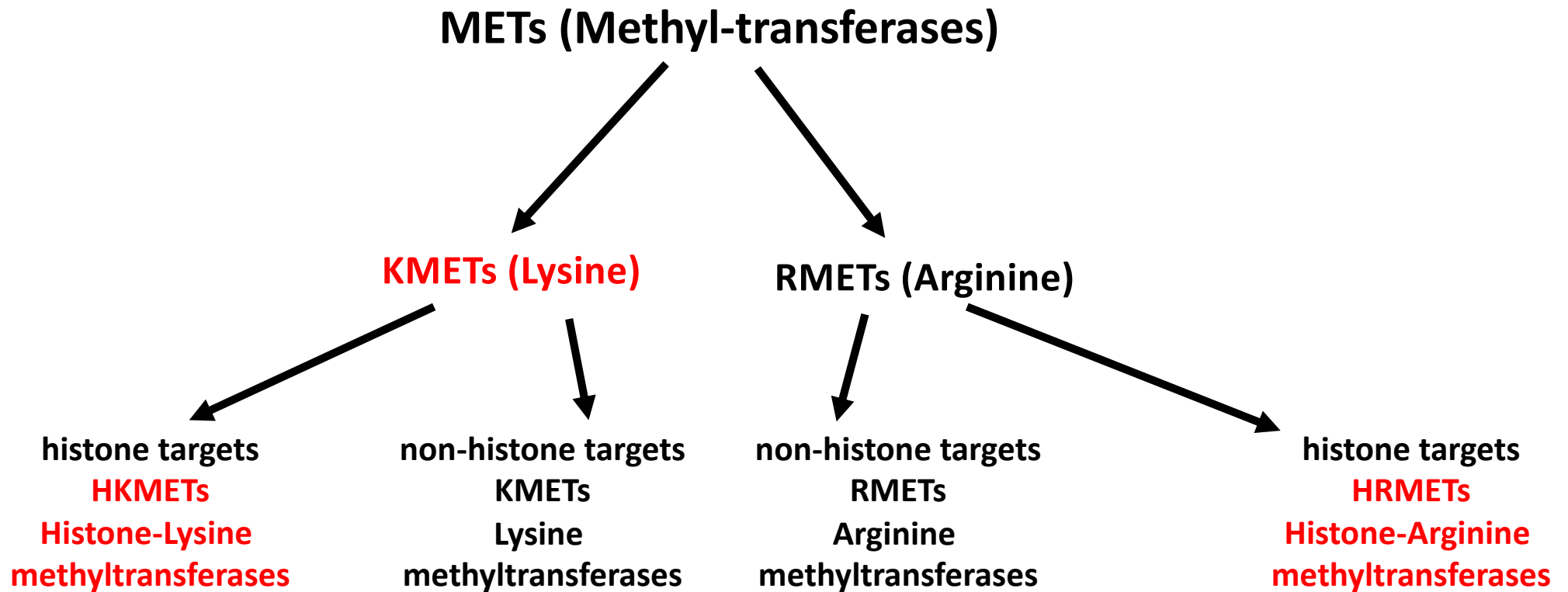
LECTURE 4

HISTONE METHYLATION AND DNA METHYLATION

LECTURE 4

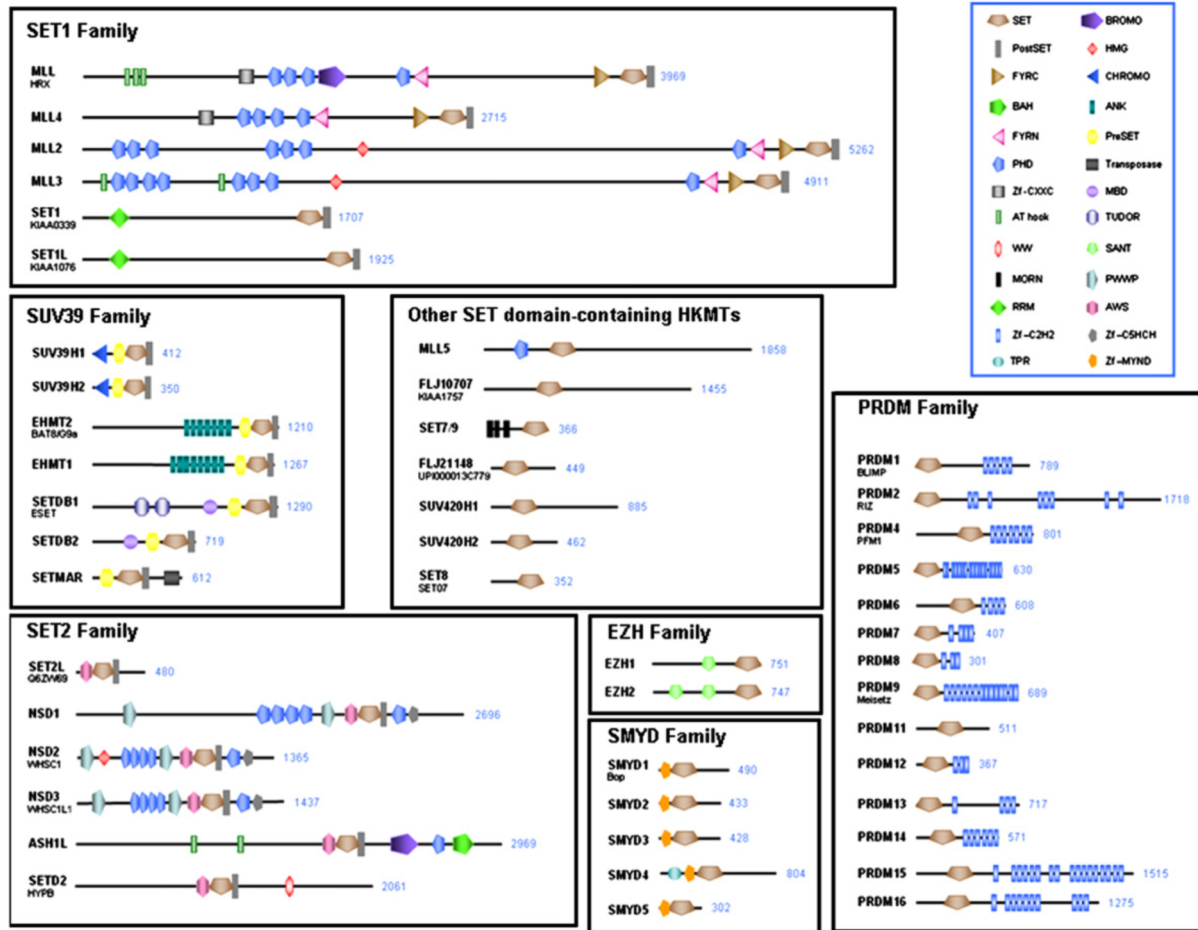
HISTONE METHYLATION MECHANISMS

HISTONE LYSINE AND ARGININE METHYL TRANSFERASES (HKMETs and HRMETs))



HISTONE LYSINE METHYL TRANSFERASES (HKMETs)

all HKMETs contain a conserved SET domain that catalyzes the methylation of Lysines (K)
(exception Dot1 – no SET domain)



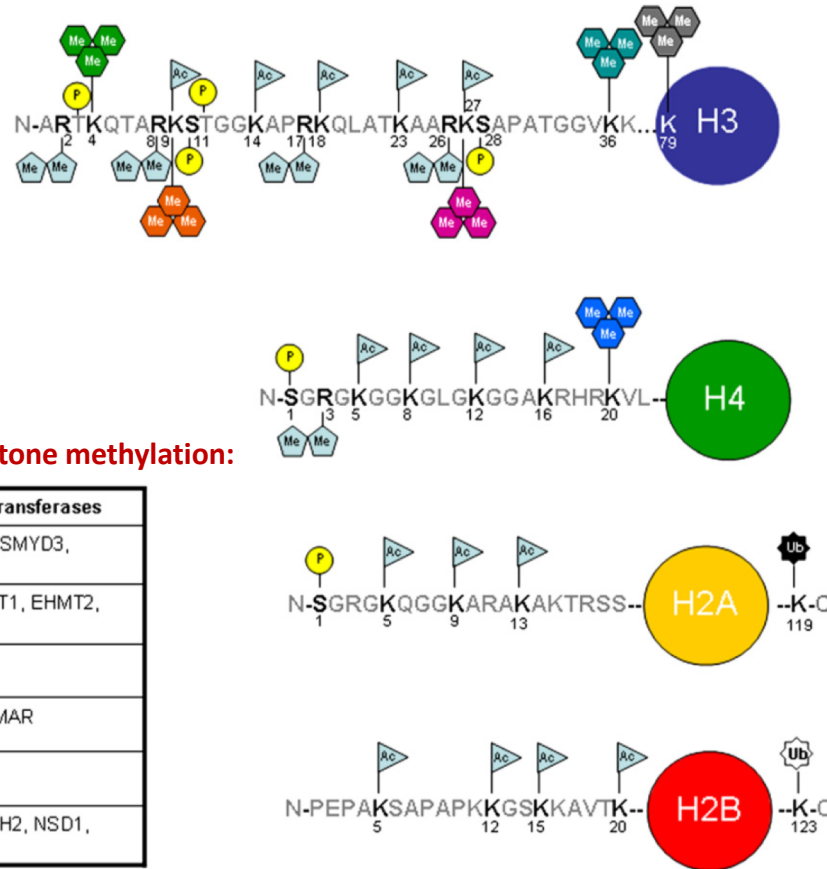
50 SET domain proteins are
Categorized according to sequence
homology
into 6 HKMET subfamilies

1. SET1 family
2. SET2 family
3. SUV39 family
4. EZH family
5. SMYD family
6. PRDM family
7. other SET domain HKMETs

50 SET domain proteins contain many
other protein domains
→ Interaction with other proteins or
DNA

HKMET HRMET SUBSTRATES ON HUMAN HISTONES

A



Effect on gene activity: Best studied examples of histone methylation:

- activation
- repression
- repression
- activation
- activation
- repression

Substrate	Histone lysine methyltransferases
H3K4	SET9, SET1, MLL, ASH1L, SMYD3, PRDM9, SETMAR
H3K9	SUV39H1, SUV39H2, EHMT1, EHMT2, SETDB1, PRDM2, ASH1L
H3K27	EZH2, EHMT2
H3K36	NSD1, SETD2/HYPB, SETMAR
H3K79	DOT1L
H4K20	SET8, SUV420H1, SUV420H2, NSD1, ASH1L

HKMETS epigenetic writers are substrate specific and can result in gene repression but also gene activation →→→

The epigenetic reader that binds to the modified histone K residue at the individual histone tail makes the difference

Fig. 1. Histone modifications. (A) The modifications on human histones include methylation (Me) on arginine and lysine residues, acetylation (Ac) on lysine residues, phosphorylation (P) on serine and threonine residues and ubiquitination (Ub) on lysine residues. (B) The enzymes responsible for methylation of human histone lysine residues are listed according to their target sites. Histone lysine methyltransferases (HKMTs) are very specific but redundant in several cases.

HISTONE MODIFICATIONS AND EPIGENETIC READERS

Protein domains that bind to histone modifications

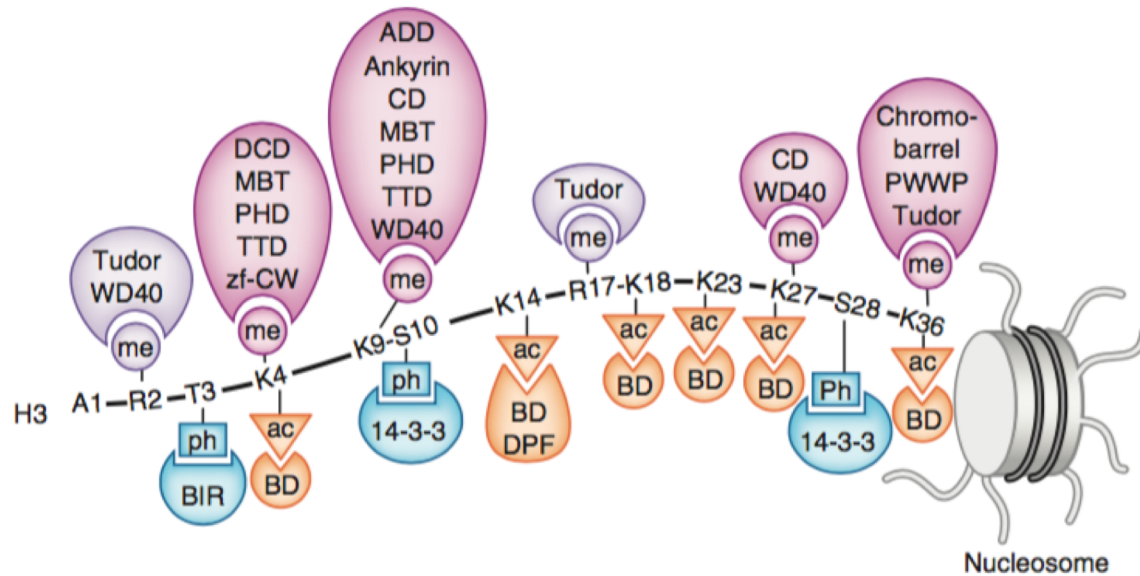


Figure 1 Readers of histone PTMs. Recognition of the methylated (me) lysine, methylated (me) arginine, acetylated (ac) lysine and phosphorylated (ph) serine and threonine residues of the N-terminal histone H3 tail by indicated readers.

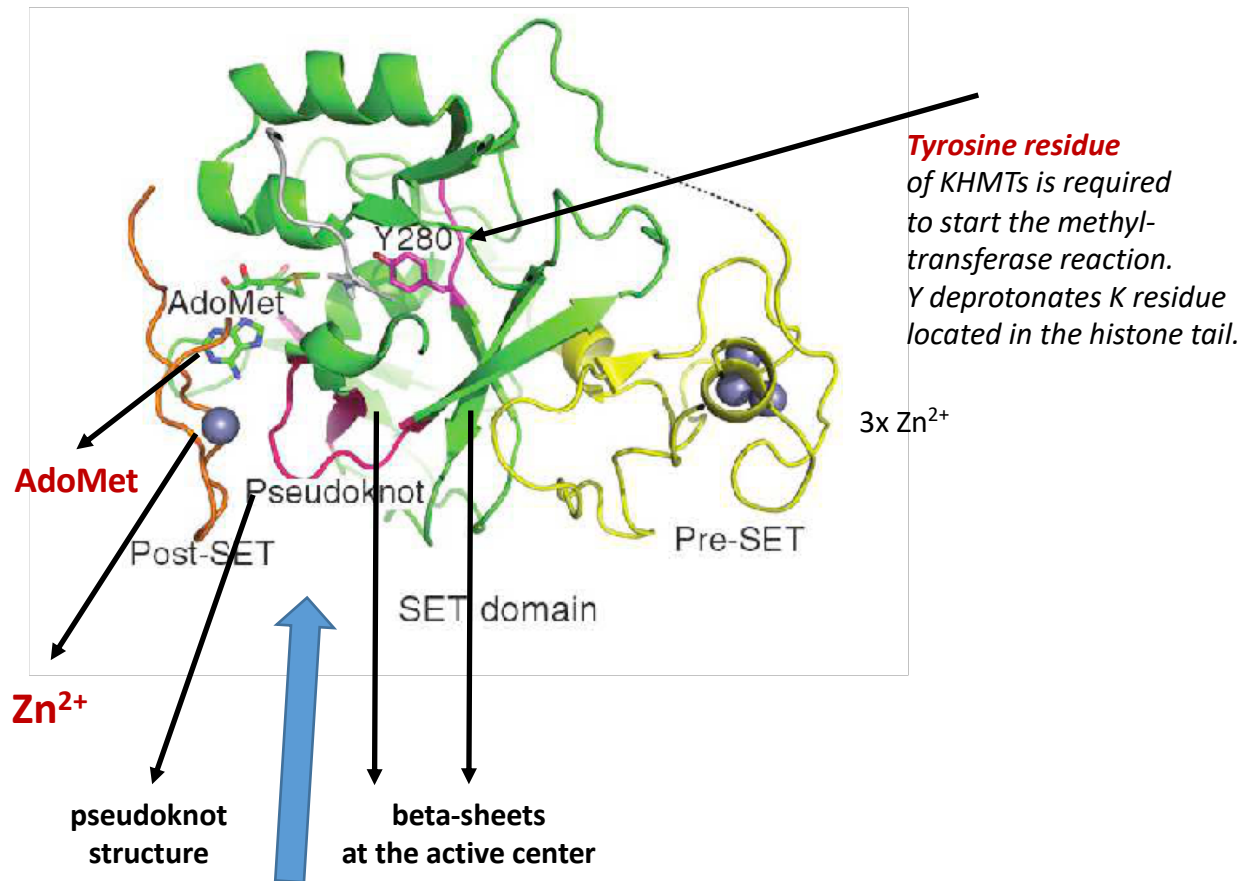
A large number of proteins contain these protein domains:
 → High complexity in gene regulation that
 → Creation of large numbers of EPIGENOMES

Table 1 Histone readers and their target PTMs

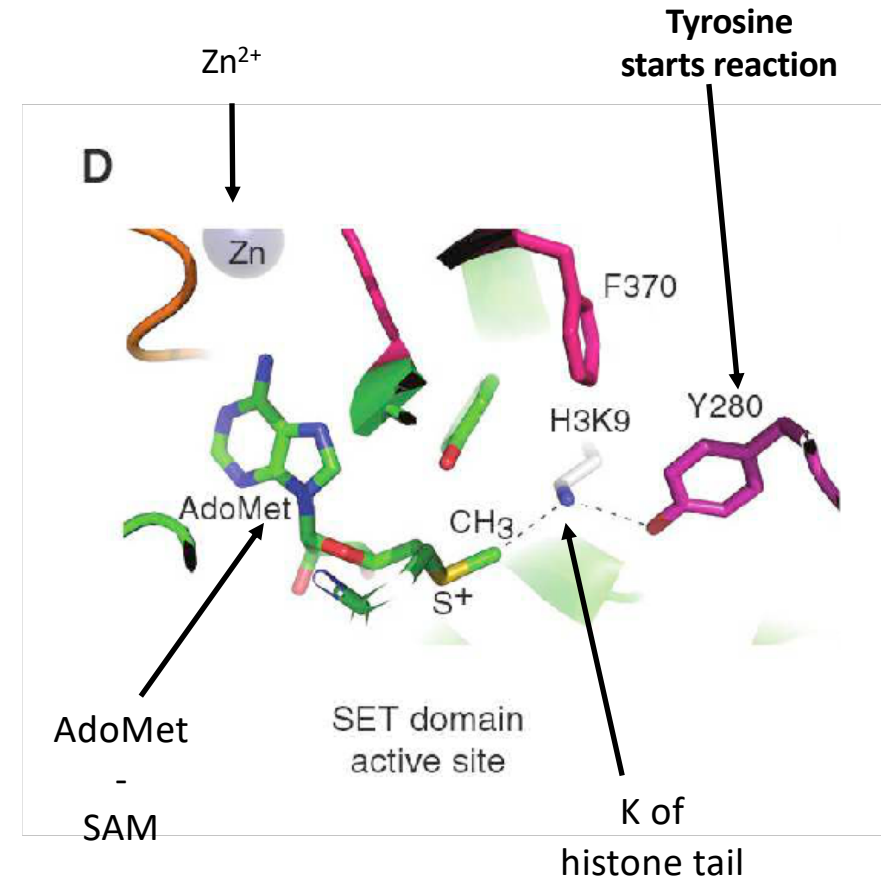
Recognition of	Reader	Histone PTM
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	BAH	H4K20me2
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	Chromodomain	H3K9me3, H3K9me2, H3K27me3, H3K27me2
	DCD	H3K4me3, H3K4me2, H3K4me1
	MBT	H3Kme1, H3Kme2, H4Kme1, H4Kme2
	PHD	H3K4me3, H3K4me2, H3K9me3
	PWWP	H3K36me3, H4K20me1, H4K20me3, H3K79me3
	TTD	H3K4me3, H3K9me3, H4K20me2
Methylarginine	Tudor	H3K36me3
	WD40	H3K27me3, H3K9me3
	zf-CW	H3K4me3
Acetyllysine	ADD	H4R3me2s
	Tudor	H3Rme2, H4Rme2
	WD40	H3R2me2
	Bromodomain	H3Kac, H4Kac, H2AKac, H2BKac
Phosphoserine or phosphothreonine	DBD	H3KacKac, H4KacKac
	DPF	H3Kac
	Double PH	H3K56ac
Unmodified histone	14-3-3	H3S10ph, H3S28ph
	BIR	H3T3ph
Unmodified histone	Tandem BRCT	H2AXS139ph
	ADD	H3un
	PHD	H3un
	WD40	H3un

THE SET DOMAIN – EXCLUSIVELY IN KMETS

THE SET DOMAIN



THE ACTIVE SITE IN THE SET DOMAIN



THE BIOCHEMISTRY OF HISTONE LYSINE METHYLATION

The source of the methyl group is S-adenosyl-l-methionine (AdoMet) or (SAM), which is converted to S-adenosyl-l-homocysteine (AdoHcy) in the reaction.

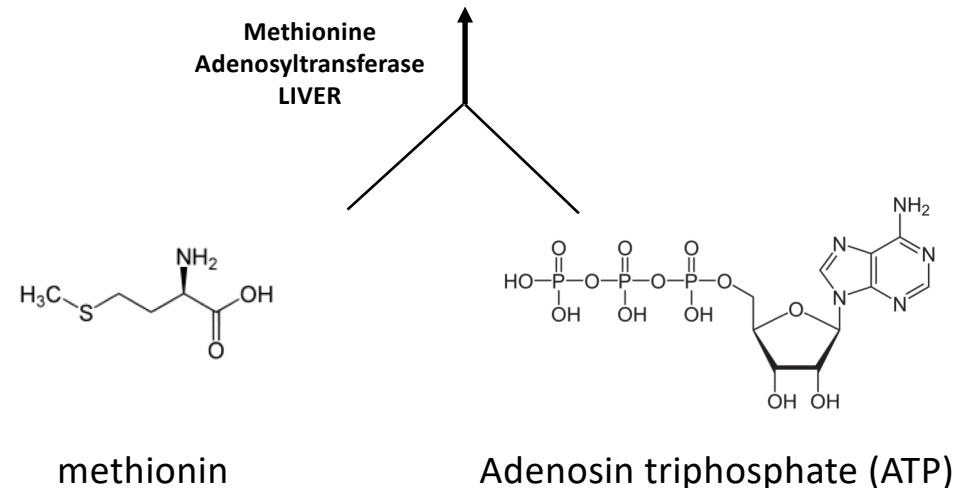
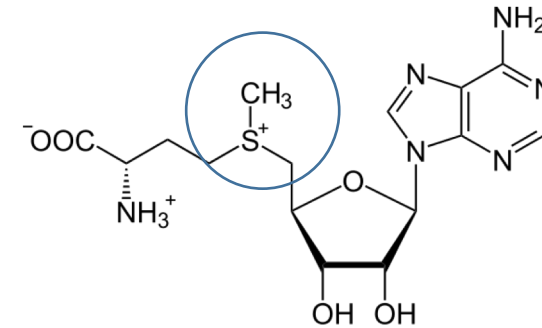
S-Adenosyl methionine is a common cosubstrate involved in methyl group transfers, transsulfuration, and aminopropylation.

SAM = enzymatic cofactor

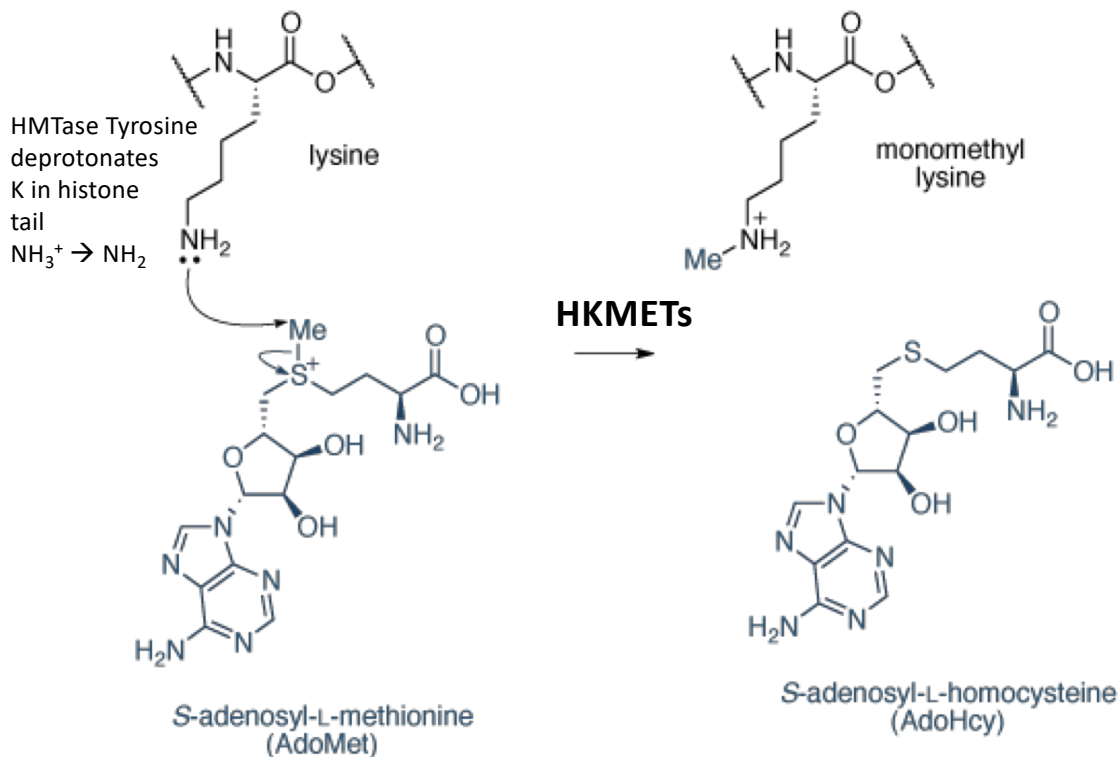
SAM is after ATP the most commonly used cofactor used by the cell

Although these anabolic reactions occur throughout the body, most **SAM is produced and consumed in the liver**. More than 40 methyl transfers from SAM are known, to various substrates such as nucleic acids, proteins, lipids and secondary metabolites. It is made from adenosine triphosphate (ATP) and methionine by methionine adenosyltransferase. SAM was first discovered in Italy by Giulio Cantoni in 1952.

S-adenosyl-l-methionine (AdoMet) or (SAM),



THE BIOCHEMISTRY OF HISTONE LYSINE METHYLATION



Catalytic mechanism

In order for the reaction to proceed, S-Adenosyl methionine (SAM) and the lysine residue of the substrate histone tail must first be bound and properly oriented in the catalytic pocket of the SET domain. Next, a **nearby tyrosine residue deprotonates the ϵ -amino group of the lysine residue.** The lysine chain then makes a **nucleophilic attack on the methyl group on the sulfur atom of the SAM molecule, transferring the methyl group to the lysine side chain.**

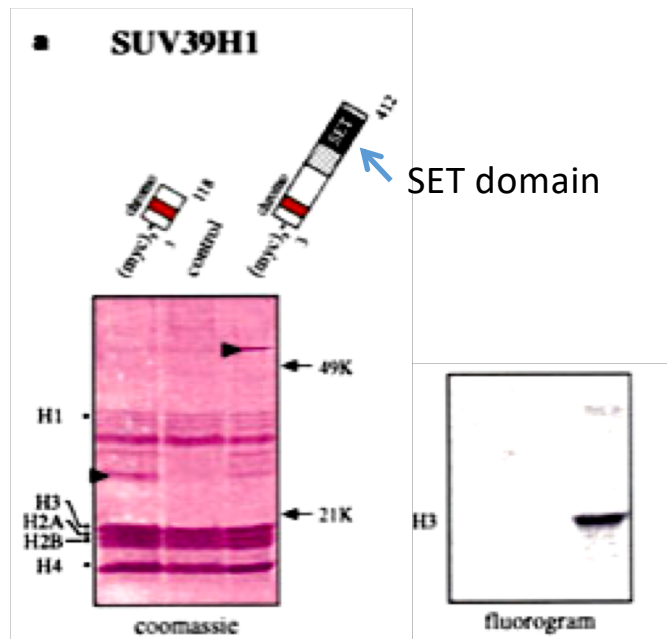
ENZYMATIC ASSAY TO DETECT KMTase ACTIVITY

Experiment:

Overexpression of **myc-tagged-SUV39H1 KMT** in HeLa cells

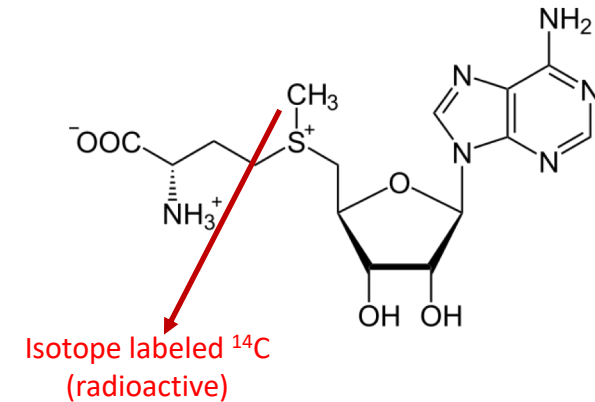
Use an antibody to immunoprecipitate SUV39H1 → high concentration of SUV39H1

Incubate Immunoprecipitate with purified histones and S-adenosyl-[methyl-¹⁴C]-L-methionin as methyl donor



SET – domain is required for histone methyl transferases activity

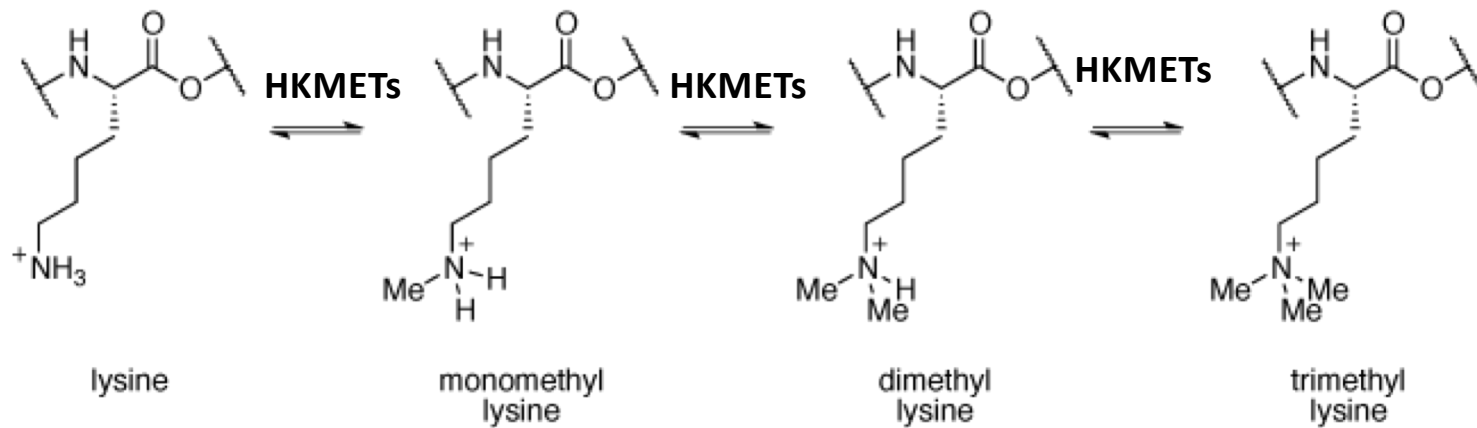
S-adenosyl-L-methionine (AdoMet) or (SAM),



- The SET domain of the SUV39H1 is required for histone methyltransferase activity and this enzyme methylates H3 at Lys9

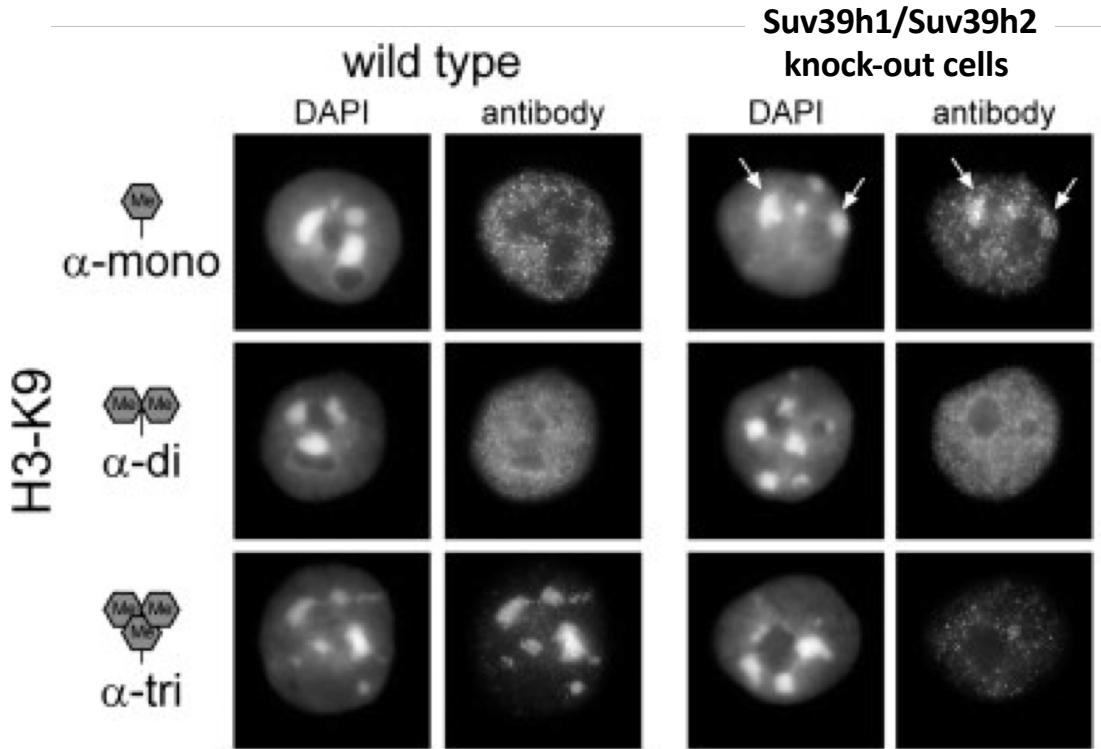
HISTONE LYSINES CAN BE MONO- DI- AND TRI-METHYLATED

lysine methylation



**ARE THERE KMTs THAT CREATE SPECIFIC METHYLATION LEVELS
(mono-methylation, di-methylation, tri-methylation?)**

**SUBSTRATE SPECIFICITY OF HISTONE METHYL TRANSFERASES:
AN EXAMPLE: THE HKMT SUV39H1**

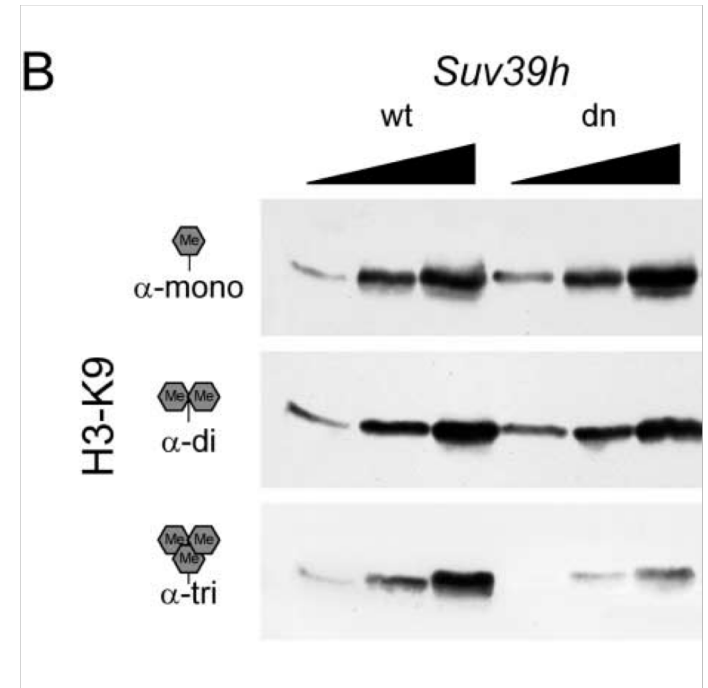


Suv39dn cells

H3K9me1: increased and pattern similar to wt H3K9me3 (chromocenter)

H3K9me2: similar to wt

H3K9me3: strongly reduced; lost at chromocenters



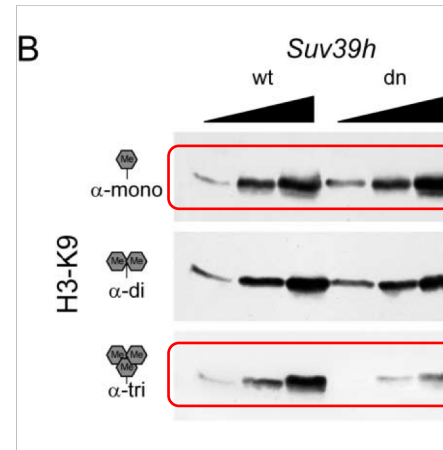
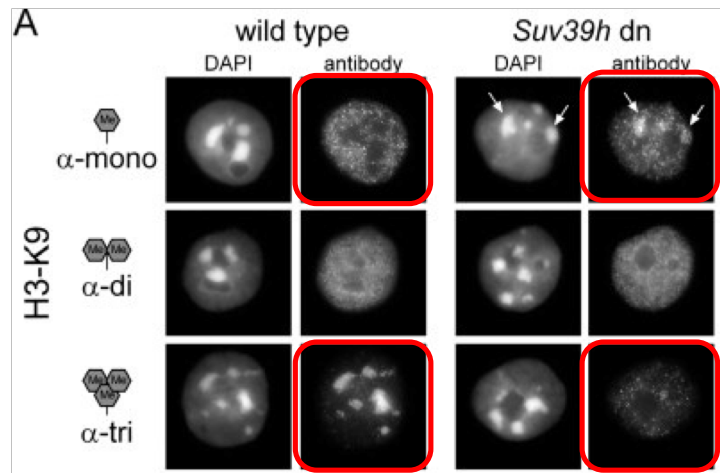
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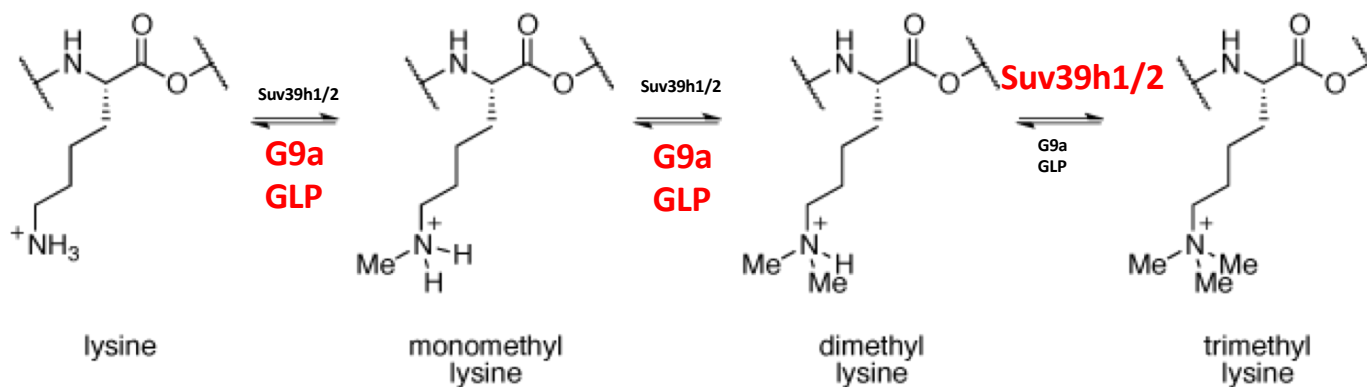


The H3K9 specific KMTs G9a and GLP are the major H3K9me1 and H3K9me2 methyltransferases

The H3K9 specific KMTases Suv39h1 and Suv39h2 are the major H3K9me3 methyltransferases

Suv39h1 and Suv39h1 work best on H3K9me2

H3K9 methylation



EPIGENETIC READERS AN EXAMPLE: H3K9me3 and HP1

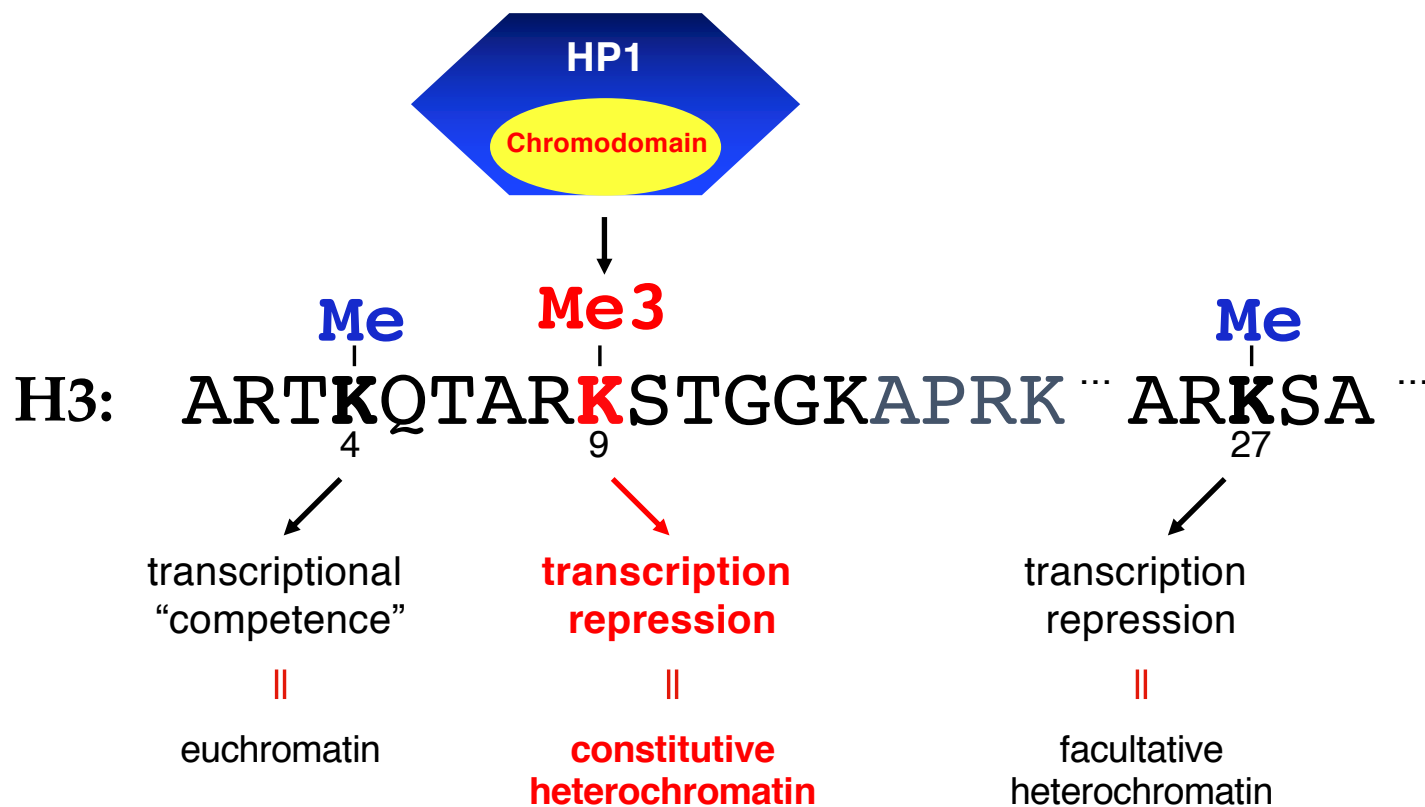
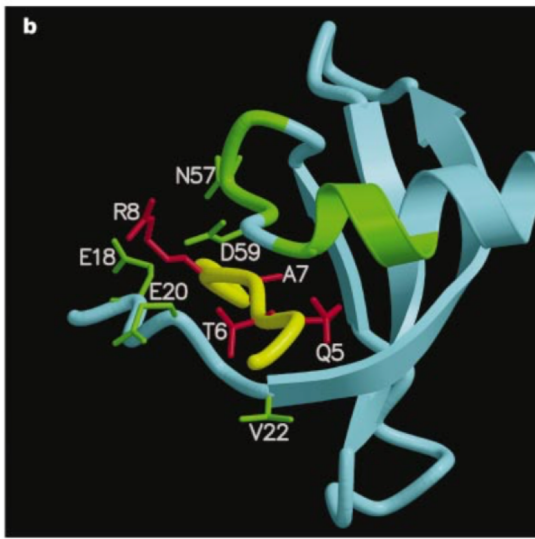


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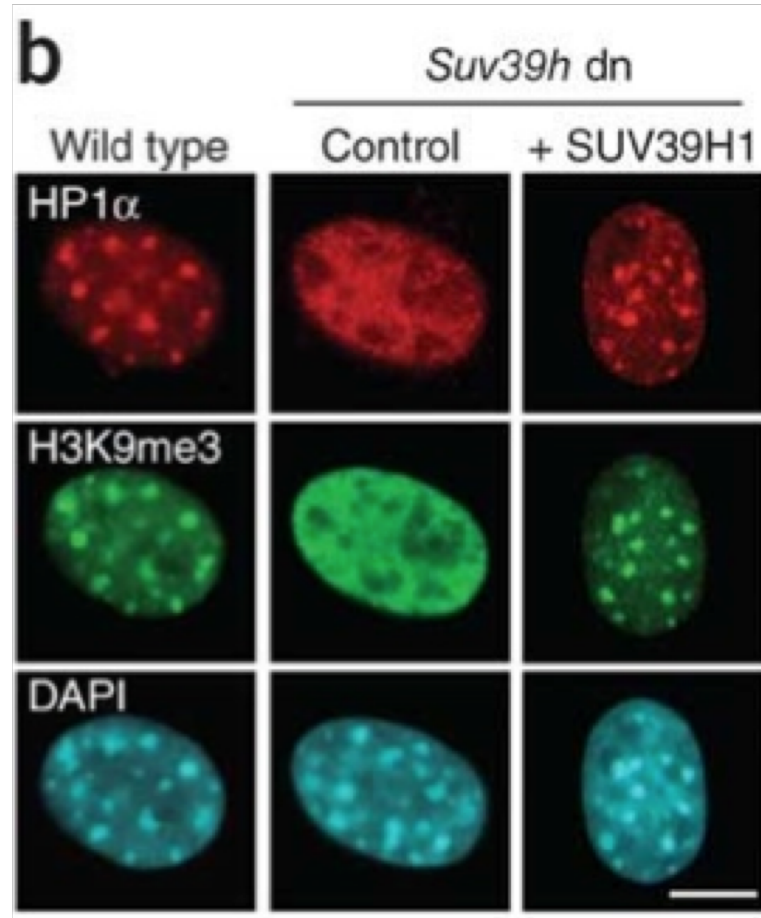
EPIGENETIC READERS – IN VIVO EVIDENCE
AN EXAMPLE: HP1 has high affinity for H3K9me3



A chromodomain (chromatin organization modifier) is a protein structural domain of about 40-50 amino acid residues commonly found in proteins associated with the remodeling and manipulation of chromatin. The domain is highly conserved among both plants and animals, and is represented in a large number of different proteins in many genomes, such as that of the mouse. Chromodomain-containing proteins also bind methylated histones and appear in the RNA-induced transcriptional silencing complex.

YELLOW: histone tail

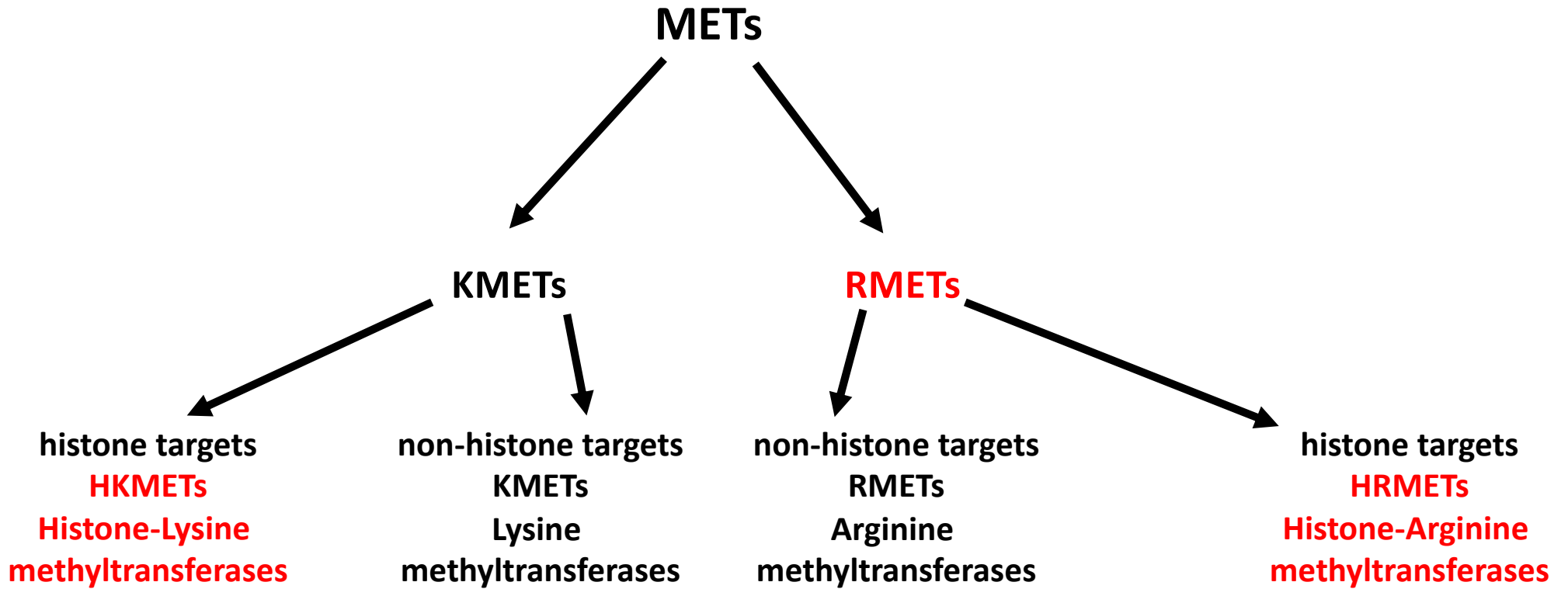
RODS: Interacting aminoacids of HP1



Loss of Suv39h1/2:
reduced
 (2 slides earlier)
and delocalized
 (this slide) **H3K9me3.**

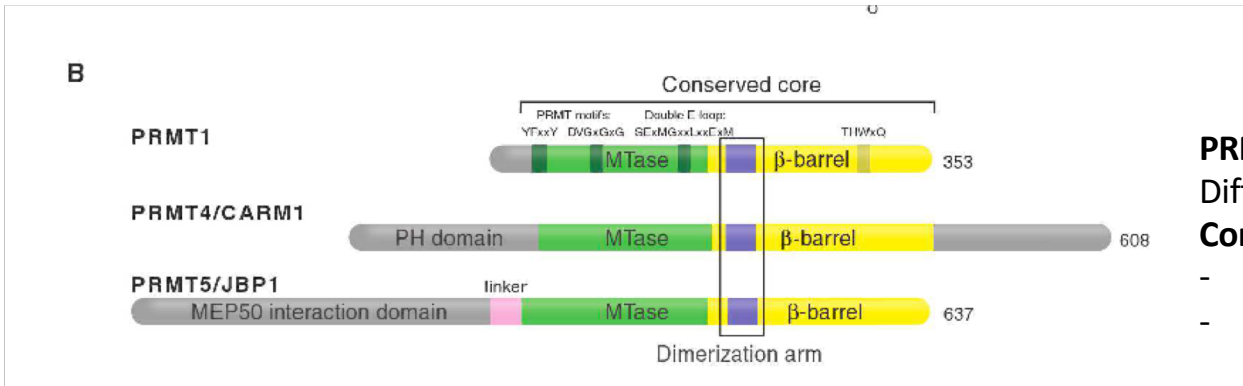
Consequence:
HP1 is also
delocalized!!!!
= binding to low-
affinity targets

HISTONE LYSINE AND ARGININE METHYL TRANSFERASES (HKMETs and HRMETs))



HISTONE ARGININE METHYL TRANSFERASES (HRMETS)

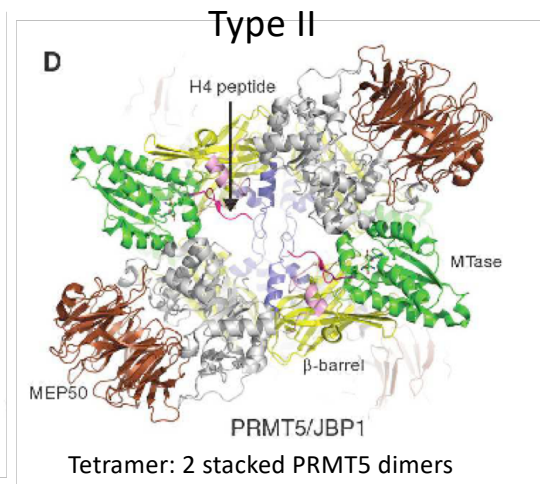
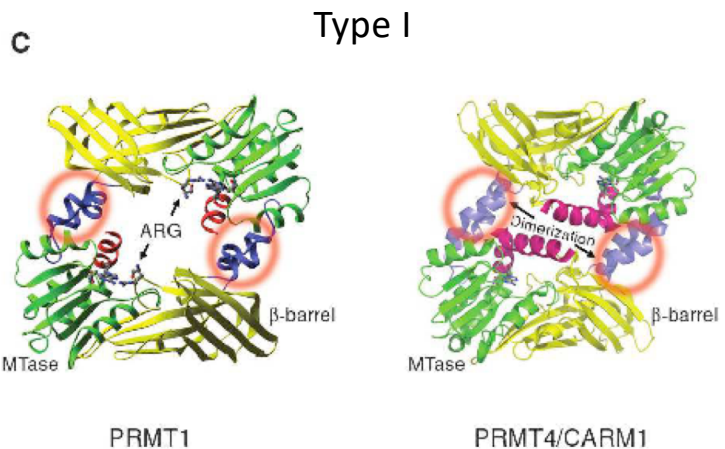
Family of PRMTs: Protein Arginine (R) methyl-transferases



PRMTs have a MTase domain that is Different from the SET domain!!!

Conserved core:

- MTase domain: catalyzes methylation of R
- Beta barrel domain: Important for dimerization of PRMTs



PRMTs

- Type I PRMTs: need to dimerize to be functional
- Type II PRMTs: form larger complexes – dimers interact to form tetramers, other proteins can interact

THE BIOCHEMISTRY OF HISTONE ARGININE METHYLATION

Methyl transfer reactions catalyzed by AdoMet-dependent PRMTs.

Example: PRMT1

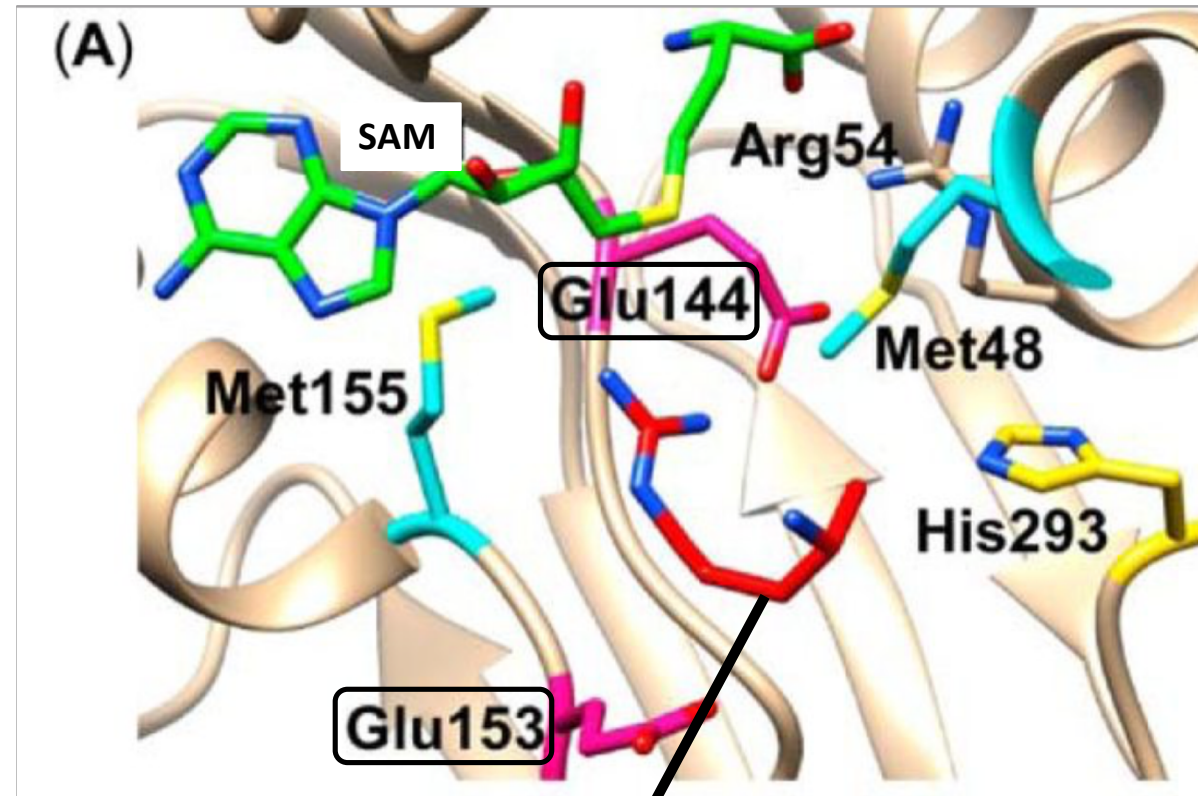
The reacting arginine substrate acts by nucleophilic attack on the methyl group present on SAM (S-AdoMet).

The reaction has been proposed to involve 3 key conserved residues in the active site of PRMT1: Arg-54, Glu-144, and Glu-153.

Arg-54 and **Glu-144** help to properly position the substrates for the nucleophilic attack

Glu-153 is hypothesized to play a role in increasing the nucleophilicity of the guanidinium moiety of the substrate via enhanced electronic effects.

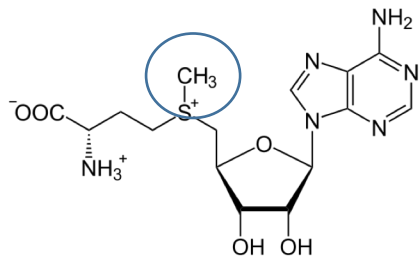
Glu-144 has also been postulated to act as the active site base, abstracting a proton from the reacting arginine.



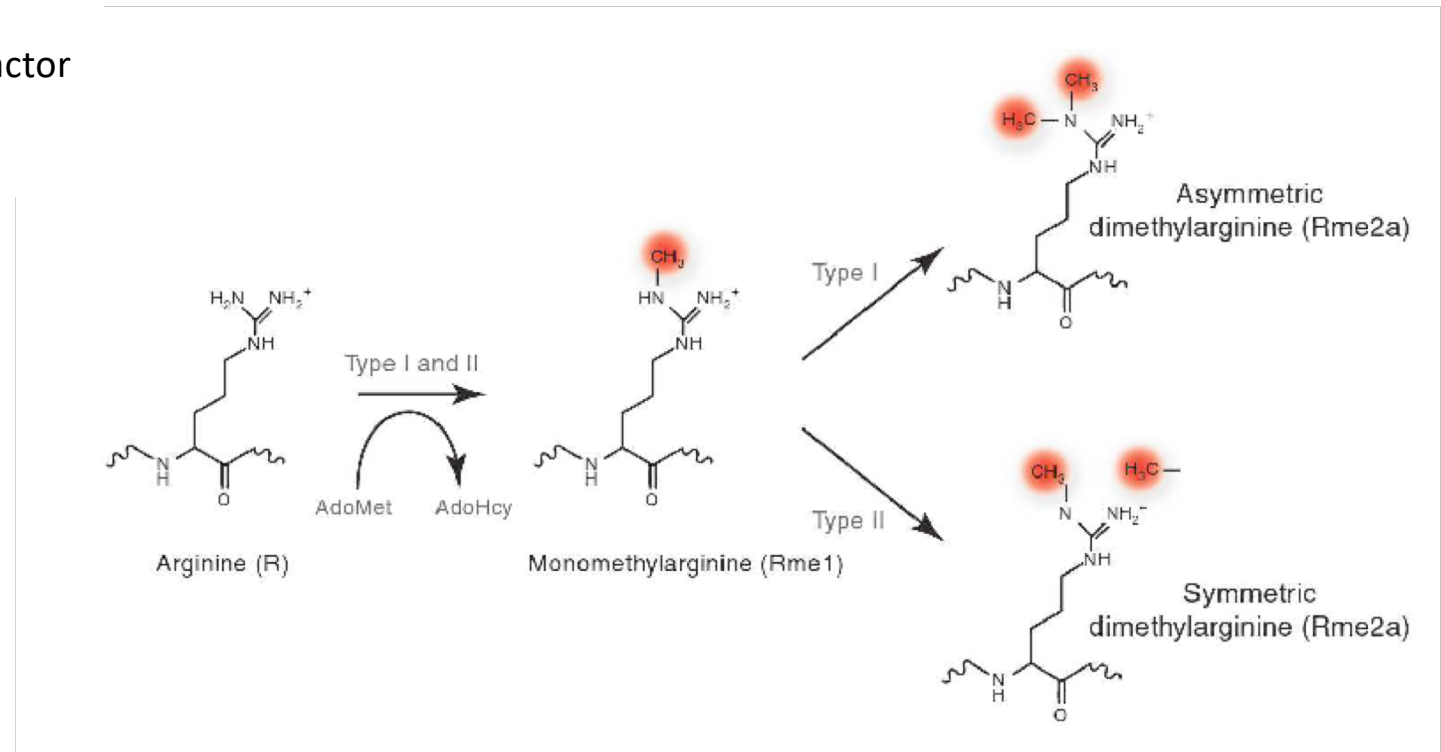
Substrate – Arginine (R)

THE BIOCHEMISTRY OF HISTONE ARGININE METHYLATION

PRMT activity requires :
substrate containing R,
AdoMet (SAM) as enzymatic cofactor
PRMT



S-adenosyl-L-methionine (AdoMet) or (SAM),



PRMTs CATALYZE MONO and DIMETHYLATION
- Not trimethylation -

PRMT SUBSTRATES AND BIOLOGICAL ACTIVITY

PRMTs can act as activators and repressors of gene expression

PRMTs:		Type	Histone substrate	Biological Function
PRMT1		I	H4R3	<u>NR, chromatin dynamic, transcription activation</u>
PRMT2		?		Coactivator for ER, Cellular proliferation
PRMT3		I		ribosomal biosynthesis
PRMT4		I	H3R2, H3R17 (Rare)	<u>NR, transcription activation, epigenetic reprogram in embryos</u>
PRMT5		II	H4R3; H3R8	<u>Stem cell function, transcription repression, repressive chromatin</u>
PRMT6		I	H3R2	<u>Repressive chromatin, supression of H3K4 methylation</u>
PRMT7		II	H2A, H4R3	Potentiating DNMT3 binding, regulation of imprinting genes
PRMT8		I	H4?	?
PRMT9 Isoform 4		II	H4, H2A	?
PRMT10		?		?
PRMT11		?		?

PRMTs epigenetic writers, are substrate specific and can result in gene repression but also gene activation

→→→

The epigenetic reader that binds to the modified histone R residue at the individual histone tail makes the difference

HISTONE MODIFICATIONS AND EPIGENETIC READERS

Protein domains that bind to histone modifications

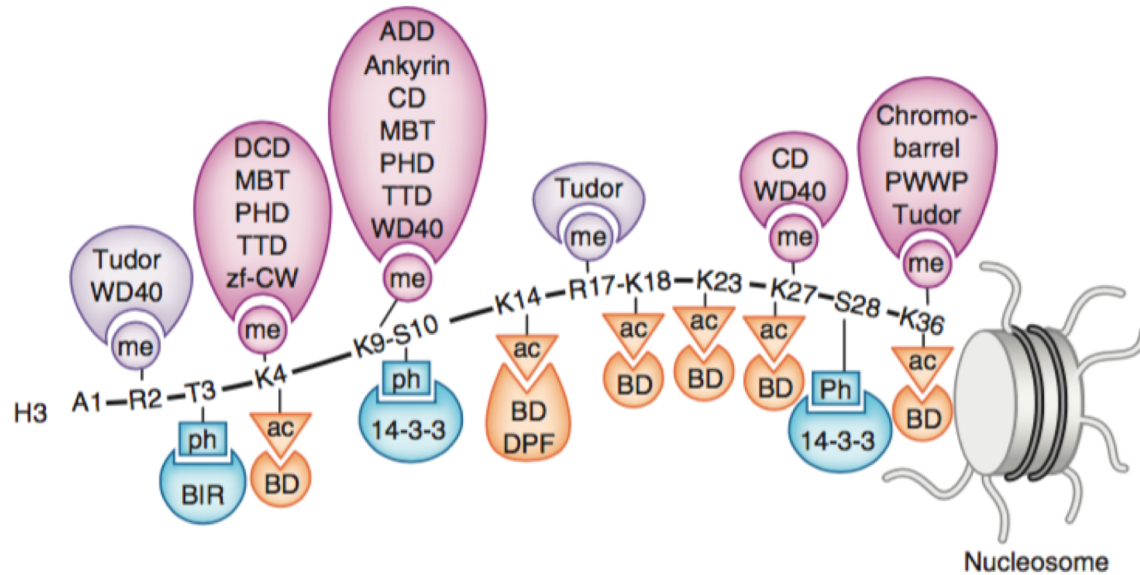


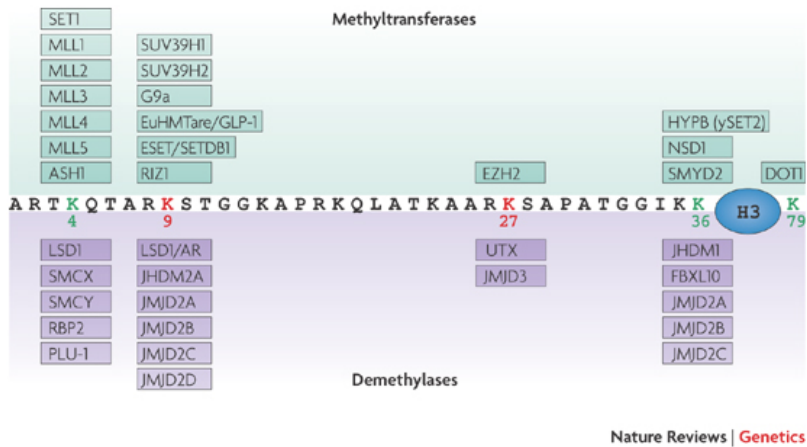
Figure 1 Readers of histone PTMs. Recognition of the methylated (me) lysine, methylated (me) arginine, acetylated (ac) lysine and phosphorylated (ph) serine and threonine residues of the N-terminal histone H3 tail by indicated readers.

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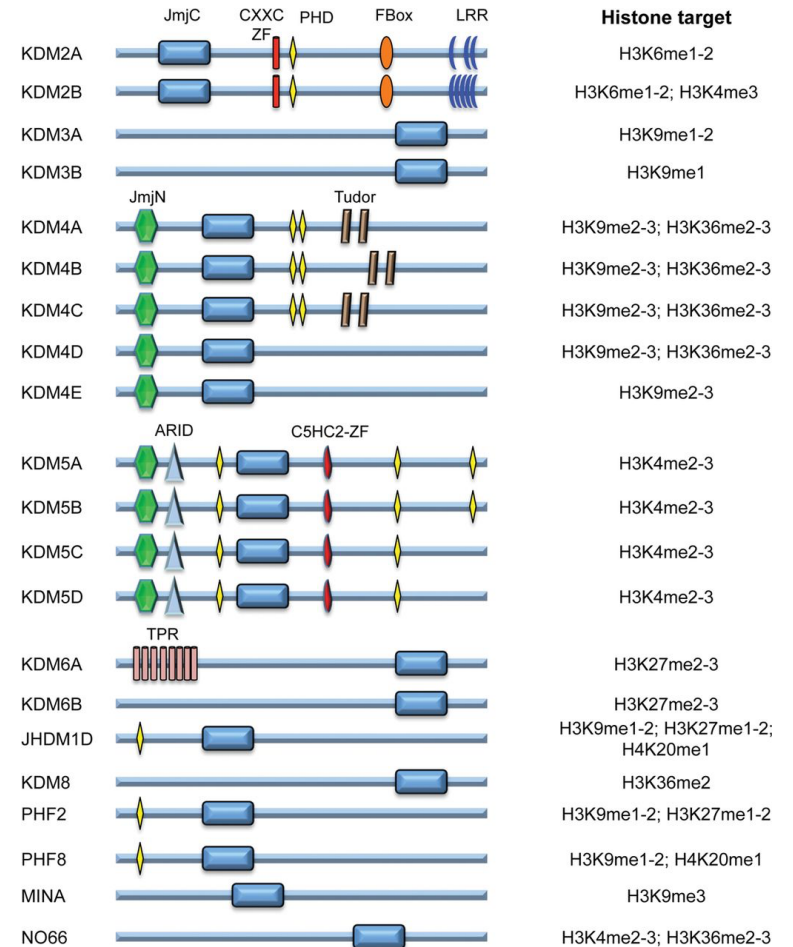
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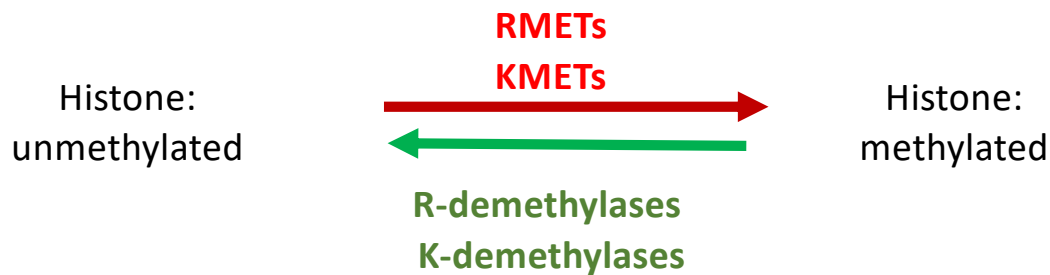
LYSINE AND ARGININE METHYLATION IN HISTONES IS REVERSIBLE



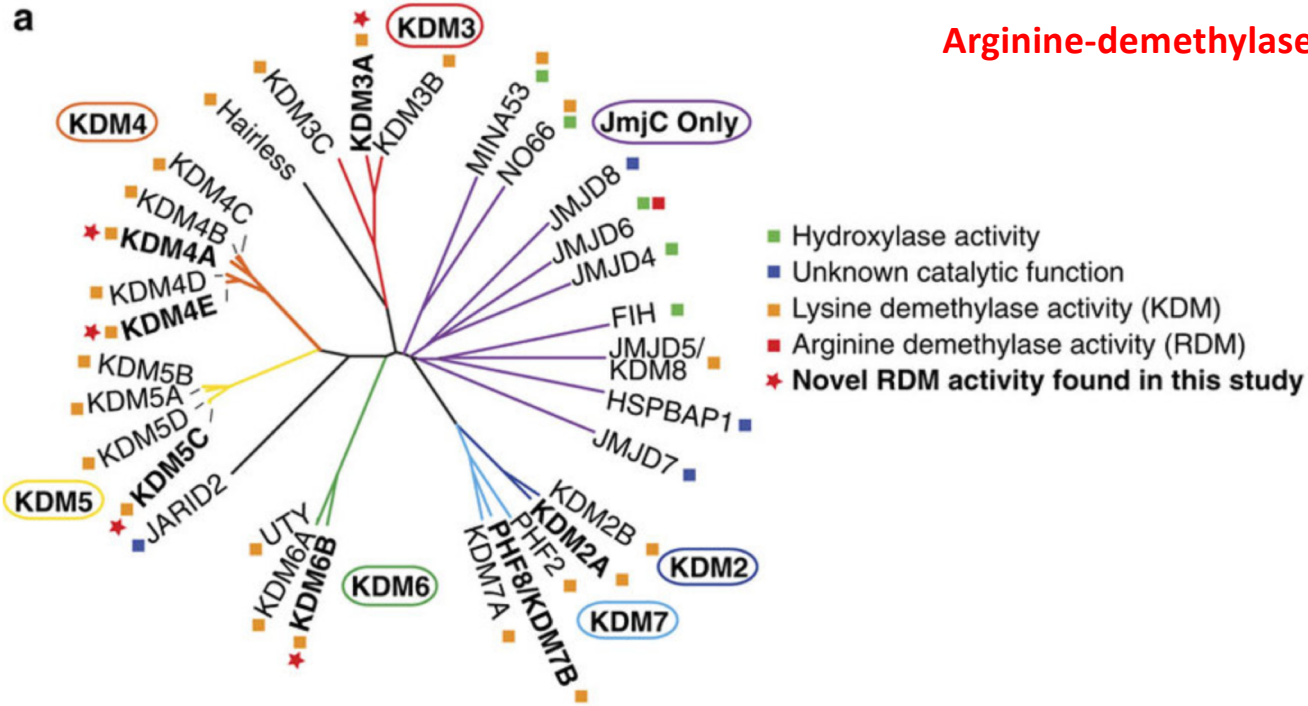
The **Jumonji N (JmjN)** and **Jumonji C (JmjC)** domains are two non-adjacent domains which have been identified in the jumonji family of transcription factors. Although it was originally suggested that the JmjN and JmjC domains always co-occur and might form a single functional unit within the folded protein, the JmjC domain was latter found without the JmjN domain in organisms from bacteria to human. The JmjC domain is the best studied domain that mediated histone demethylation - is conserved from yeast to human



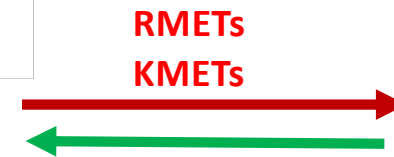
1. LSD1 (KDM1A): demethylation by oxidation
2. Big family of Jumonji domain containing proteins: hydroxylation



LYSINE AND ARGININE METHYLATION IN HISTONES IS REVERSIBLE



Histone:
unmethylated



Histone:
methylated

LECTURE 4

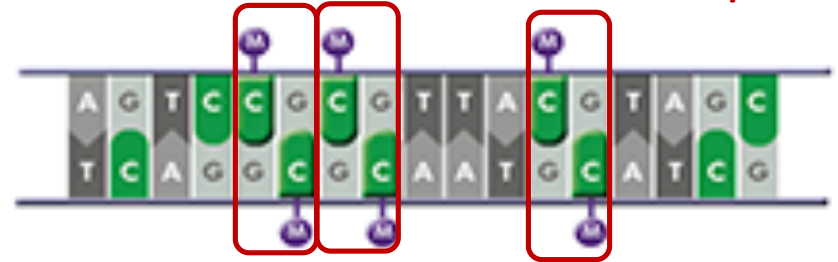
DNA METHYLATION

DNA METHYLATION CONTROLS GENE EXPRESSION

FACTS:

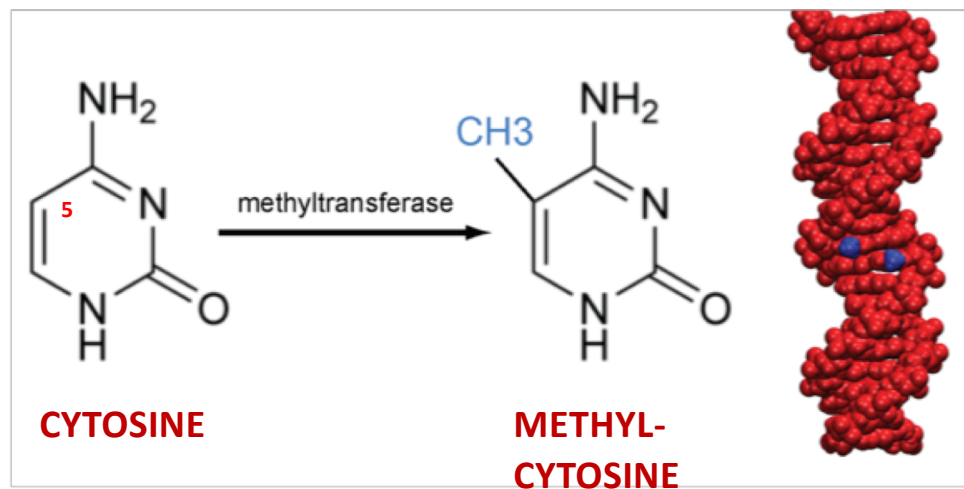
1. DNA methylation is created at CpG di-nucleotide motifs
2. An accumulation of CpG is called "CpG island" (CGI)
3. CpG islands are enriched at promoters and sequence elements that are important for gene expression control. In some cases, CpG islands can be also located in distant locations.
4. *CpG methylation (=“DNA methylation”) is directly linked with stable, inheritable gene silencing*

ADVANTAGE OF DNA METHYLATION AT CpG



CpGs are self-complementary
Di-nucleotide in paired stand also contains methylation
Methylation patterns can be maintained during DNA replication

Epigenetic
modification
that is imposed
on genomic **DNA**

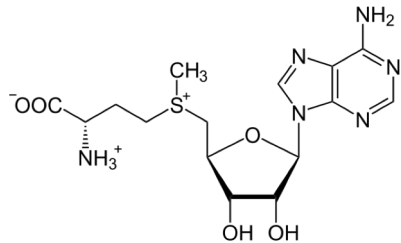


CYTOSINE

METHYL-
CYTOSINE

DNA METHYLTRANSFERASES CATALYZE DNA METHYLATION

DNA methyltransferases (DNMTs) transfer a methyl-group from AdoMet (SAM) to Cytosine located in a CpG dinucleotide



S-adenosyl-L-methionine (AdoMet) or (SAM),

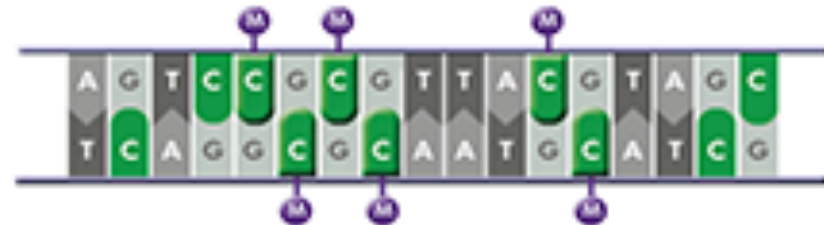
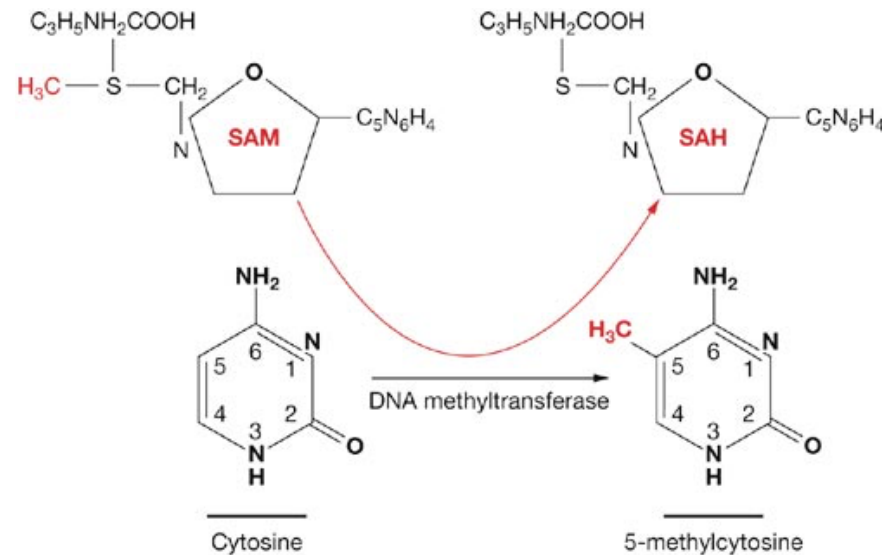
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S-Adenosyl methionine is a common cosubstrate involved in methyl group transfers, transsulfuration, and aminopropylation.

SAM = enzymatic cofactor

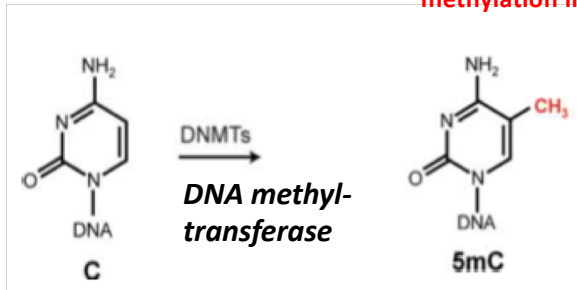
SAM is after ATP the most commonly used cofactor used by the cell

Although these anabolic reactions occur throughout the body, most SAM-e is produced and consumed in the liver. More than 40 methyl transfers from SAM-e are known, to various substrates such as nucleic acids, proteins, lipids and secondary metabolites. It is made from adenosine triphosphate (ATP) and methionine by methionine adenosyltransferase. SAM was first discovered in Italy by Giulio Cantoni in 1952.



Mapping DNA methylation at CpG islands BISULFITE SEQUENCING

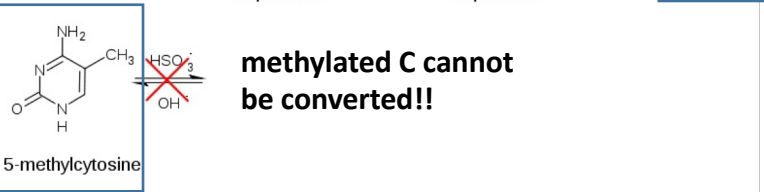
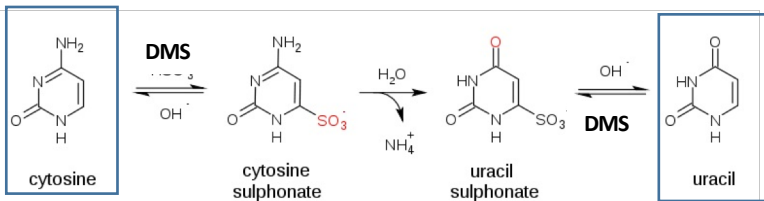
Methylation of cytosine at CpG dinucleotides is an important epigenetic regulatory modification in many eukaryotic genomes. DNA methylation was found to be located genome-wide with a pattern of low methylation in proximity to promoters and high gene body-methylation in highly-expressed genes → methylation pattern can identify transcribed DNA (gene)



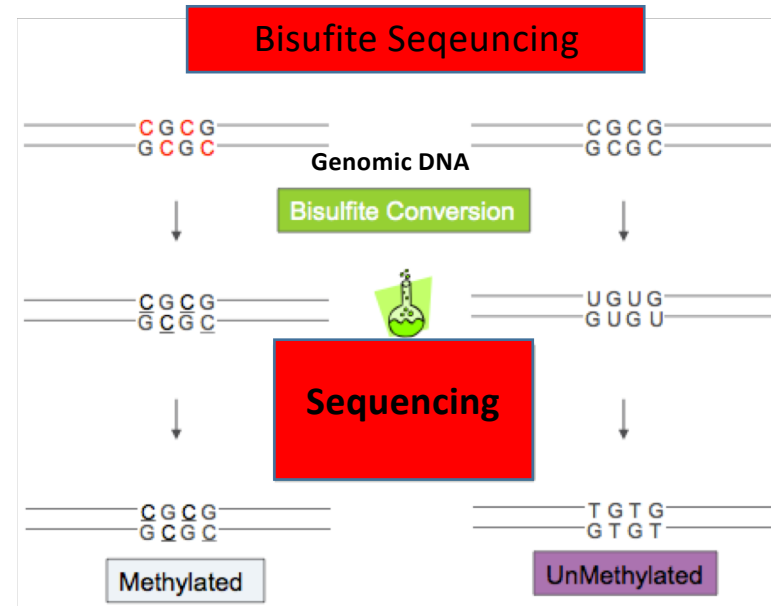
active gene

silenced gene

Bisulfite conversion: C → U conversion using dimethyl sulfate



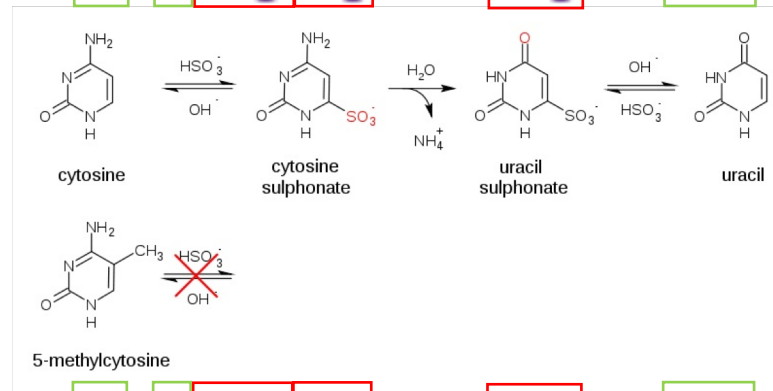
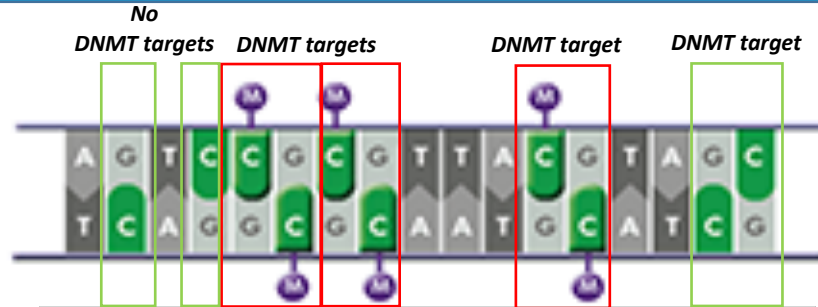
DMS = Dimethyl sulfate



Treatment of DNA with bisulfite converts cytosine residues to uracil, but leaves 5-methylcytosine residues unaffected. Thus, bisulfite treatment introduces specific changes in the DNA sequence that depend on the methylation status of individual cytosine residues, yielding single-nucleotide resolution information about the methylation status of a segment of DNA.

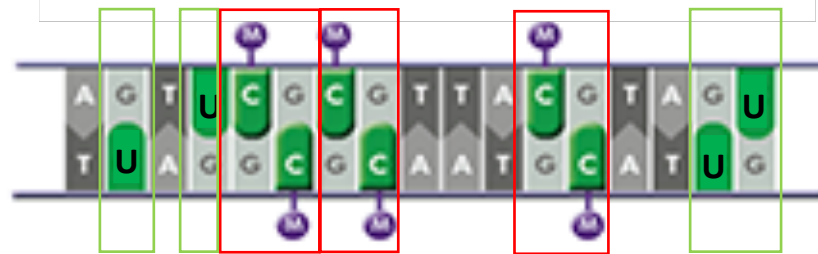
Mapping DNA methylation at CpG islands BISULFITE SEQUENCING

Genomic DNA



Bisulfite conversion

DNA for Sequencing



Sequencing of both strands reveals C → U (T) transition

Compare with genomic sequence
 C → U sequence change = DNA methylation
 C → C no sequence change = no DNA methylation