LECTURE 4

HISTONE METHYALTION AND DNA METHYLATION

LECTURE 4

HISTONE METHYALTION 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

Α





Effect on gene activity Best studied examples of histone methylation:

	Substrate	Histone lysine methyltransferases
activation	H3K4	SET9, SET1, MLL, ASH1L, SMYD3, PRDM9, SETMAR
repression	H3K9	SUV39H1, SUV39H2, EHMT1, EHMT2, SETDB1, PRDM2, ASH1L
repression	H3K27	EZH2, EHMT2
activation	H3 K36	NSD1, SETD2/HYPB, SETMAR
activation	H3K79	DOT1L
repression	H4 K20	SET8, SUV420H1, SUV420H2, NSD1, ASH1L



HKMETS epigenetic writers are substrate specific and can result in gene repression but also gene activation $\rightarrow \rightarrow \rightarrow$

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 EPIGENTIC 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:
- \rightarrow 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
/	Methyllysine	ADD	H3K9me3
		Ankyrin	H3K9me2, H3K9me1
		BAH	H4K20me2
		Chromo-barrel	H3K36me3, H3K36me2, H4K20me1, H3K4me1
		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
		Tudor	H3K36me3
		WD40	H3K27me3, H3K9me3
		zf-CW	H3K4me3
	Methylarginine	ADD	H4R3me2s
		Tudor	H3Rme2, H4Rme2
		WD40	H3R2me2
	Acetyllysine	Bromodomain	H3Kac, H4Kac, H2AKac, H2BKac
		DBD	H3KacKac, H4KacKac
		DPF	H3Kac
		Double PH	H3K56ac
	Phosphoserine or	14-3-3	H3S10ph, H3S28ph
	phosphothreonine	BIR	H3T3ph
		Tandem BRCT	H2AXS139ph
	Unmodified histone	ADD	H3un
		PHD	H3un
		WD40	H3un

THE SET DOMAIN



THE BIOCHEMISTRY OF HISTONE LYSINE METHYLATION

The source of the methyl group is S-adenosyl-lmethionine (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 <u>cofactor used by the cell</u> 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-I-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 \rightarrow high concentration of SUV39H1

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

a SUV39H1







• The SET domain of the SUV39H1 is required for histone methyltransferase activity

and this enzyme methylates H3 at Lys9

Rea et al, Nature, 2000

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

lysine methylation



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

SUBSTRATE SPECIFICITY OF HISTOME 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



Suv39dn cells

H3K9me1: increased compared to wt H3K9me2: similar to wt H3K9me3: strongly reduced

SUBSTRATE SPECIFICITY OF HISTOME METHYL TRANSFERASES: AN EXAMPLE: THE HKMT SUV39H1



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

EPIGENTIC READERS AN EXAMPLE: H3K9me3 and HP1



EPIGENTIC 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



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



HISTONE ARGININE METHYL TRANSFERASES (HRMETs)

Family if 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 AdoMetdependent PRMTs.

Example: PRMT1

The reacting arginine substrate acts by nucleophil attacking the methyl group present on 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.

<u>**Glu-144**</u> has also been postulated to act as the active site base, abstracting a proton from the reacting arginine either during or immediately after methyl transfer.



THE BIOCHEMISTRY OF HISTONE ARGININE METHYLATION



PRMTs CATALYZE MONO and DIMETHYLATION - Not trimethylation -

PRMT SUBSTRATES AND BIOLOGICAL ACTIVITY

PRMTs can act as activators and repressors of gene expression

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

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

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

HISTONE MODIFICATIONS AND EPIGENTIC READERS

Protein domains that bind to histone modifications



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	TTD	H3K4me3, H3K9me3, H4K20me2
	Tudor	H3K36me3
	WD40	H3K27me3, H3K9me3
	zf-CW	H3K4me3
Methylarginine	ADD	H4R3me2s
	Tudor	H3Rme2, H4Rme2
	WD40	H3R2me2
Acetyllysine	Bromodomain	H3Kac, H4Kac, H2AKac, H2BKac
	DBD	H3KacKac, H4KacKac
	DPF	H3Kac
	Double PH	H3K56ac
Phosphoserine or	14-3-3	H3S10ph, H3S28ph
phosphothreonine	BIR	H3T3ph
	Tandem BRCT	H2AXS139ph
Unmodified histone	ADD	H3un
	PHD	H3un
	WD40	H3un

AN EXAMPLE FOR PRMT FUNCTION IN TRANSCRIPTIONAL ACTIVATION



- H4R3me stimulates subsequent acetlylation of H4
- p53 binds transcription factor binding site in promoter and brings p300.
- p300 acetylates promoter regions
- PRMT4 has affinity for acetylated histones H3R17me
- (note: PRMT1, 4 do not contain a bromo domain \rightarrow other proteins mediate affinity to acetylated histones)

\rightarrow FULL TRANSCRIPTIONAL ACTIVATION

LYSINE AND ARGININE METHYLATION IN HISTONES IS REVERSIBLE



The Jumonji N (JmjN) and Jumon ji 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 mia ht form a single functional unit within the folded protein, the JmiC domain was latter found without the JmjN in organisms domain from bacteria to human. The JmjC domain is the best studied domain that mediated histone demethylation - is conserved from yeast to human

LSD1 (KDM1A): demethylation by <u>oxidation</u> Big family of Jumonji domain containing proteins: hydroxylation





LYSINE AND ARGININE METHYLATION IN HISTONES IS REVERSIBLE



Louise J. Walport¹, Richard J. Hopkinson¹, Rasheduzzaman Chowdhury¹, Rachel Schiller¹, Wei Ge¹, Akane Kawamura^{1,2} & Christopher J. Schofield¹

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 promotes 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 gene silencing



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



DNA METHYLTRANSFERASES CATALYZE DNA METHYLATION

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



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

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-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 genebodymethylation in highly-expressed genes → methylation pattern can identify transcribed DNA (gene)



Bisulfite conversion: $C \rightarrow U$ conversion



Treatment of DNA with bisulfite converts cytosine residues to uracil, but leaves 5methylcytosine 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.



DMS = Dimethyl sulfate

Mapping DNA methylation at CpG islands BISULFITE SEQUENCING



Mapping DNA methylation at CpG islands BISULFITE SEQUENCING

An example:

hTERT encodes the telomerase gene hTERT elongates telomeres thereby protecting cancer cells from replicative senescence

- → Cancer cells do not age and can proliferate forever!!
- → Therefore 90% of human cancers express telomeres; but telomerase is silenced by DNA methylation in differentiated cells.
- → Bisulfite sequencing showed that cancer cells have a de-methyalted CpG island located at the hTERT promoter. CpG islands can overlap with the 1st intron of the gene!!!!





Mapping DNA methylation at CpG islands of individual genes BISULFITE SEQUENCING



Mapping DNA methylation at CpG islands BISULFITE SEQUENCING

Prepare DNA from normal cell and cancer cell

Purify DNA and perform bi-sulfite conversion

Amplify your region of interest = CpG islands in the TERT promoter

Purify DNA fragment obtained by PCR

Clone fragment into Plasmid

Transform bacteria with plasmid

Purify amplified plasmids

Sequence inserts using a primer that anneals to the vector DNA, adjacent the insertion site of the PCR product REMEMBER: only a single type of plasmid is sequenced This refers to a single type of molecule. To have a good representation, you need to sequence at least 10 clones

HOW DO I HAVE TO DESIGN THE PRIMER FOR PCR OF BISULFITE CONVERTED DNA??



C Cytosine not in CpG site C G CpG site

Compare with genomic sequence Bisulfite conversion: $C \rightarrow U$ sequence change = DNA methylation $C \rightarrow C$ no sequence change = no DNA methylation

Mapping DNA methylation at CpG islands BISULFITE SEQUENCING



Mapping DNA methylation at CpG islands METHYLATED DNA IMMUNOPRECIPITATION: METHYL-DIP

Methyl-DIP works similar to ChIP: **2** experimental samples:

- Control (normal cells)
- Experimental sample (cancer cells)

Prepare DNA - carries DNA methylation marks of CpG

Sonicate DNA

Immunoprecipitation using an antimethyl-CpG specific Antibody (monoclonal, discriminates between CpG and met-CpG)

Washing of precipitate

DATA ACQUISITION

Main methods to analyse DNA methylation

- 1. PCR on specific CpG islands of interest
- 2. Differential labelling (Cy3-control; Cy5-cancer methyl-DNA) followed by hybridization to genome array
- 3. Massive parallel sequencing (different approaches)





Reduced representation bisulfite sequencing (RRBS) is an efficient and high-throughput technique used to analyze the genome-wide methylation profiles on a single nucleotide level. This technique combines restriction enzymes and bisulfite sequencing in order to enrich for the areas of the genome that have a high CpG content. Due to the high cost and depth of sequencing needed to analyze methylation status in the entire genome. The fragments that comprise the reduced genome but still includes the majority of promoters, as well as regions such as repeated sequences that are difficult to profile using conventional bisulfite sequencing approaches.

Mapping DNA methylation genome wide: REDUCED REPRESENTATION BISULFITE SEQEUNCING (RRBS)



The color key from light blue to dark blue indicates the DNA methylation level from low to high, respectively. The white regions in the left panels indicate a lack of DNA methylation information. The red bars in the right panel represent the average DNA methylation level of the corresponding genomic region. The DNA methylation levels were calculated and presented based on 30 kb windows, only if these windows have more than 5 CpG sites covered.

Mapping DNA methylation genome wide: REDUCED REPRESENTATION BISULFITE SEQEUNCING (RRBS)



Enzyme digestion: First, genomic DNA is digested using a methylation-insensitive restriction enzyme. It is integral for the enzymes to not be influenced by the methylation status of the CoGs (sites within the genome where a cytosine is next to a guanine) as this allows for the digestion of both methylated and unmethylated areas. Mspl is commonly used. This enzyme targets 5'CCGG3' sequences and cleaves the phosphodiester bonds upstream of CpG dinucleotide. When using this particular enzyme, each fragment will have a CpG at each end. This digestion results in DNA fragments of various sizes. End repair and A-tailing: Due to the nature of how Mspl cleaves double stranded DNA, this reaction results in strands with sticky ends. End repair is necessary to fill in the 3' terminal of the ends of the strands. The next step is adding an extra adenosine to both the plus and minus strands. This is referred to as A-Tailing and is necessary for adapter ligation in the subsequent step. End repair and A-Tailing is done within the same reactions, with dCTP, dGTP and dATP deoxyribonucleotides. In order to increase the efficiency of A tailing, the dATPs are added in excess in this reaction. Sequence adapters: Methylated sequence adapters are ligated to the DNA fragments. The methylated adapter oligonucleotides have all cytosines replaced with 5'methyl-cytosines, in order to prevent the deamination of these cytosines in the bisulfite conversion reaction. For reactions to be sequenced using Illumina sequencers, the sequence adapters are used to hybridize to the adapters on the flow cell. Fragment purification: The desired size of fragments is then selected to be purified. The different sizes of the fragments are separated using gel electrophoresis and are purified using gel excising. According to Guetal., DNA fragments of 40-220 base pair are representative of the majority of promoter sequences and CpG islands[2]Bisulfite conversion: The DNA fragments are then bisulfite converted, which is a process that deaminates unmethylated cytosine into a uracil. The methylated cytosines remain unchanged, due to the methyl group protecting them from the reaction.PCR amplification: The bisulfite converted DNA is then amplified using PCR with primers that are complementary to the sequence adapters.PCR purification: Before sequencing, the PCR product must be free of unused reaction reagents such as unincorporated dNTPs or salts. Thus, a step for PCR purification is required. This can be done by running another electrophoresis gel or by using kits designed specifically for PCR purification. Sequencing: The fragments are then sequenced. When RRBS was first developed, Sanger sequencing was initially used. Now, next generation sequencing approaches are used. For Illumina sequencing, 36-base single-end sequencing reads are most commonly performed.Sequence alignment and analysis: Due to the unique properties of RRBS, special software is needed for alignment and analysis. Using Mspl to digest genomic DNA results in fragments that always start with a C (if the cytosine is methylated) or a T (if a cytosine was not methylated and was converted to a uracil in the bisulfite conversion reaction). This results in a non-random base pair composition. Additionally, the base composition is skewed due to the biased frequencies of C and T within the samples. Various software for alignment and analysis is available, such as Mag, BS Seeker, Bismark or BSMAP. Alignment to a reference genome allows the programs to identify base pairs within the genome that are methylated.

DNA methyltransferases methylate DNA

FACTS: 1. DNA methylation is created at CpG di-nucleotide motifs

2. An accumulation of CpG genes is called CpG island

3. CpG islands are enriched at promoters and other, more distant sequence elements that are important for gene expression control

4. CpG methylation (="DNA methylation") is linked with Stable gene silencing



Methylation patterns can be maintained during DNA replication





Figure 1. Cytosine methylation in DNA. (A) Addition of a methyl group, CH₃ (red), at the five position of the cytosine pyrimidine ring (black arrow) does not sterically interfere with GC base pairing (blue lines). DNA methyltransferases associate covalently with the carbon 6 position (straight green arrow) during methyl group transfer. (B) A model of B-form DNA methylated at cytosines in two self-complementary CpG sequences. The paired methyl moieties (magenta and yellow) lie in the major groove of the double helix.

De-novo DNMTs \rightarrow place new DNA methylation

Maintenance DNMTs \rightarrow propagate methylation after replication

DNA methyl transferases methylate DNA

Biochemical discovery of maintenance DNMT1:

Cell extract + DNA containing CpG repeats + 14 C labelled -CH3 in AdoMet (SAM) \rightarrow radioactive -CH3 transferred to DNA

NOTE:

This enzyme is 7– to 100-fold more active on hemimethylated DNA as compared with unmethylated substrate *in vitro*

Next step: Purification of enzymatic activity: 200kDa complex containing **DNMT1**

Discovery of de novo DNMTs:

Sequence of DNMT1 was used to look For genes with similar sequence (sequence homology)

- → Discovery of de-novo DNMTs that work efficiently work on un-methylated DNA (DNMT3a, 3b)
- → De-novo DNMTs cannot efficiently methylate hemi-methylated DNA







LOSS OF DNA METHYLTRANSFERASES IS LETHAL DURING EMBRYONIC MOUSE DEVELOPMENT

Table 1. Function of mammalian DNA methyltransferases

DNA	methyl-		Major	
transferas	e	Species	activity	Major phenotypes of loss of function
Dnmt1		Mouse	Maintenance C methylation of CpG	Genome-wide loss of DNA methylation, embryonic lethality at embryonic day 9.5 (E9.5), abnormal expression of im- printed genes, ectopic X-chromosome inactivation, activation of silent retro- transposon. In cancer cell lines, it leads to cell cycle arrest and mitotic defects.
Dnmt3a		Mouse	De novo P methylation of CpG	ostnatal lethality at 4–8 wk, male sterility, and failure to establish methy- lation imprints in both male and female germ cells
Dnmt3b		Mouse	De novo E methylation of CpG	Demethylation of minor satellite DNA, embryonic lethality around E14.5 days with vascular and liver defects. (Em- bryos lacking both Dnmt3a and Dn- mt3b fail to initiate de novo methylation after implantation and die at E9.5.)
DNMT3B		Human	De novo lo methylation of CpG	CF syndrome: immunodeficiency, centromeric instability, and facial anomalies. Loss of methylation in re- petitive elements and pericentromeric heterochromatin.



Figure 4. Mammalian DNA methyltransferases. The catalytic domains of Dnmt1, Dnmt2, and the Dnmt3 family members are conserved (the signature motifs, I, IV, VI, IX, and X, are most conserved in all cytosine methyltransferases), but there is little similarity among their amino-terminal regulatory domains. Domain abbreviations: PCNA, PCNA-interacting domain; NLS, nuclear localization signal; RFT, replication foci-targeting domain; CXXC, a cysteine-rich domain implicated in binding DNA sequences containing CpG dinucleotides; BAH, bromo-adjacent homology domain implicated in protein–protein interactions; PWWP, a domain containing a highly conserved "proline-tryptophan-tryptophan-proline" motif involved in heterochromatin association; ATRX, an ATRX-related cysteine-rich region containing a C2-C2 zinc finger and an atypical PHD domain implicated in protein–protein interactions.

DNA METHYLATION IS ABUNANT IN THE GENOME AND ISSUBJECTED TO DRAMATIC ALTERATIONS DURING EMBRYOGENESIS



70%- 80% of CpG dinucleotides are methylated in the genome

DNA methylation levels are high in fertilized Oocytes that contain the paternal and maternal genome (carries characteristic methylation patterns)

Paternal and maternal methylation patterns are rapidly erased (exception: imprinted genes maintain paternal and maternal methylation information). \rightarrow the paternal and maternal methylation epigenome is cancelled

DNA methylation levels remain low during the first cell division events until the blastocyst stage

In the blastocyst stage cell differentiation programs are activated and genes need to be regulated on the epigenetic level \rightarrow DNA methylation is increasing (loss of DNMT1, DNMT3a or DNMT3b is lethal \rightarrow establishment and maintenance of DNA methylation is impaired)

70%- 80% of CpG di-nucleotides are methylated in the human genome! Remember only 2% of the genome encode for mRNAs 98% is noncoding DNA that contains a large proportion of transposable elements, repeat sequences, etc... Lecture 4 Histone methylation and DNA methylation **ON THE SINGLE GENE LEVEL:** CpG islands (CGIs) are short sequences stretches with variable DNA methylation that regulate promoter activity

NOTE: single CpGs are generally hyper methylated (60-90%) CpG islands are differentially methylated, but are generally demethylate



CpG islands (CGIs) have a length of ca. 1kb

60% of human genes are controlled by CGIs containing promoters that allow tissue/cell specific gene expression

CpG islands can overlap with the first exon (methylation level in 1st exon is good predictor of gene expression

shores

TRANSCRIPTIONAL REGULATION BY METHYL-DNA BINDING PROTEINS Interference with transcription factor binding



Figure 1. Cytosine methylation in DNA. (A) Addition of a methyl group, CH_3 (red), at the five position of the cytosine pyrimidine ring (black arrow) does not sterically interfere with GC base pairing (blue lines). DNA methyltransferses associate covalendly with the carbon 6 position (straight green arrow) during methyl group transfer. (B) A model of B-form DNA methylated at cytosines in two self-complementary CpG sequences. The paired methyl moieties (magenta and yellow) lie in the major groove of the double helix.

Methylated DNA obtains different structure:
Transcription factors cannot bind anymore
→ DNA methylation sensitive transcription factors



Example: CTCF

Unmethylated DNA CTCF binds \rightarrow activation of expression Methylated DNA: CTCF does not bind \rightarrow no activation

Note: CTCF is a major epigenetic regulator that is involved in controlling genomic imprinting, enhance activation,...

EPIGENTIC READERS OF DNA METHYLATION

Transcriptional regulation by methyl-dna binding proteins

Table 2. Functions of methyl-CpG binding proteins

MADD	Major activity Spoc	Major phenotypes of loss-of-
MeCP2	Binds mCpG with Mous adjacent run AT-rich run Transcriptional repressor	e Delayed onset neurological defects including inertia, hind- limb clasping, nonrhythmic breathing, and abnormal gait. Postnatal survival ~10 wk.
MECP2	Binds mCpG with Hum adjacent AT run Transcriptional re- pressor	an Heterozygotes suffer from Rett syndrome, a profound neuro- logical disorder characterized by apraxia, loss of purposeful hand use, breathing irregulari- ties, and microcephaly
Mbd1	Binds mCpG via MBD; Mous a major splice form is also able to bind CpG via a CxxC do- main	e No overt phenotype, but subtle defects in neurogenesis de- tected
Mbd2	Binds mCpG Mous Transcriptional re- pressor	e Viable and fertile, but show reduced maternal nurturing be- havior. Defective gene regula- tion in T-helper cell differentia- tion leading to altered response to infection. Highly resistant to intestinal tumorigenesis.
Mbd3	Core component of Mou NuRD corepressor complex Does not show strong binding to mCpG	se Early embryonic lethal
Mbd4	DNA repair protein that Mou binds mCpG and T:G mismatches at mCpG sitesThymine DNA glycosylase that excises T from T:G mismatches	se Viable and fertile. three- to fourfold increase in mutations at CpG sites. Increased suscep- tibility to intestinal cancer cor- relates with C to T transitions within the Apc gene. Mbd4 functions to minimize the mu- tability of 5-methylcytosine.
Kaiso	Binds mCGmCG and Mou CTGCNA Transcrip- tional repressor	se No overt phenotype. Small but significant delay in tumorigen- esis on Min background.

Several proteins were identified to have affinity to methylated CpG but do no have affinity to un-methylated CpG \rightarrow mediate transcriptional silencing \rightarrow CpG METHYL BINDING DOMAIN PROTEIN (MBD) FAMILY : MeCP1, MeCP2, Mbd1,

- Mbd2, Mbd2, Mbd4
- → Kaiso (unrelated protein)



Figure 8. Proteins that bind methyl-CpG. Five members of the MBD protein family are aligned at their MBD domains (purple). Other domains are labeled and include TRD; CXXC domains, which are zinc fingers, some of which are implicated in binding to nonmethylated CpG; GR repeats that may bind; a T:G mismatch glycosylase domain that is involved in repair of 5mC deamination. Kaiso lacks the MBD domain, but binds methylated DNA via zinc fingers (orange) and possesses a POB/ BTB domain that is shared with other transcriptional repressors. Domain abbreviations: MBD, methyl-CpG binding domain; TRD, transcriptional repression domain; POZ, poxvirus and zinc finger, a protein–protein interacting domain.

How does MeCP2 effect the brain function?

- Through it's job as a reader of epigenetic bookmarks
- The wide array of functions that MeCP2 performs ALL contribute to Rett syndrome.
- The different mutations have different effects on the presentation of the disease.
- In addition since each person is different based on their personal epigenetics, the disease will be individual as well.



Exon 2 -

R255X

TRD

NLS

R270X

histone deacetylases

transcriptional

TFIIB

co-repressors

YB1 (RNA splicing)

Exon 3 - Exon 4

R306C

CTDa

methylated DNA

in chromatin

CTDB

WW domain

proteins

R294X

Composition of MeCP2: gene structure, splicing patterns and putative functional domains Figure 1

(A) Splicing patterns generating the two mRNA isoforms of MECP2, _e1 and _e2. The two isoforms generate two protein isoforms of MeCP2 with differing N-termini due to the use of alternative translation start sites (bent arrows) and the absence or presence of exon 2 in the transcript. (B) Apart from the N-terminus, both MeCP2 isoforms are identical and contain several functionally distinct domains: NTD. N-terminal domain: MBD, methylated DNA-binding domain: ID, interdomain; TRD, transcription repression domain; CTD, C-terminal domain; NLS; nuclear localization signals. Locations of seven of the most common point mutations in RTT are indicated (\blacklozenge). Below each domain are indicated major (bold) and other (grey) interactors and functions. HP1, heterochromatin protein 1; TFIIB, transcription factor IIB; YB1, Y-box-binding protein 1.

EPIGENTIC READERS OF DNA METHYLATION Transcriptional regulation by methyl-dna binding proteins

Table 2.	Functions of methyl-CpG bindin	g proteins
MBP	Major activity Species	Major phenotypes of loss-of- function mutations
MeCP2	Binds mCpG with Mouse adjacent run AT-rich run Transcriptional repressor	Delayed onset neurological defects including inertia, hind- limb clasping, nonrhythmic breathing, and abnormal gait. Postnatal survival ~10 wk.
MECP2	Binds mCpG with Human adjacent AT run Transcriptional re- pressor	Heterozygotes suffer from Rett syndrome, a profound neuro- logical disorder characterized by apraxia, loss of purposeful hand use, breathing irregulari- ties, and microcephaly
Mbd1	Binds mCpG via MBD; Mouse a major splice form is also able to bind CpG via a CxxC do- main	No overt phenotype, but subtle defects in neurogenesis de- tected
Mbd2	Binds mCpG Mouse Transcriptional re- pressor	Viable and fertile, but show reduced maternal nurturing be- havior. Defective gene regula- tion in T-helper cell differentia- tion leading to altered response to infection. Highly resistant to intestinal tumorigenesis.
Mbd3	Core component of Mouse NuRD corepressor complex Does not show strong binding to mCpG	Early embryonic lethal
Mbd4	DNA repair protein that Mouse binds mCpG and T:G mismatches at mCpG sitesThymine DNA glycosylase that excises T from T:G mismatches	Viable and fertile. three- to fourfold increase in mutations at CpG sites. Increased suscep- tibility to intestinal cancer cor- relates with C to T transitions within the <i>Apc</i> gene. Mbd4 functions to minimize the mu- tability of 5-methylcytosine.
Kaiso	Binds mCGmCG and Mouse CTGCNA Transcrip- tional repressor	No overt phenotype. Small but significant delay in tumorigen- esis on Min background.

Methyl-CpG binding proteins are present in transcriptional co-repressor complexes

MeCP2: component of the Sin3A HDAC complex Mbd3: component of the NuRD HDAC complex Mbd1: interacts with HDAC3. Mbd1 and HDAC3 are recruited by the PML-RARalpha hybrid protein to silence gene expression in Acute promyelocytic leukemia MBD1: interacts with the H3K9 HMTase SETDB1



Collaboration to repress genes

TRANSCRIPTIONAL REGULATION BY METHYL-DNA BINDING PROTEINS RECRUITMENT OF Methyl-CpG binding proteins and co-repressor complexes



Silencing by DNA methylation and H3K9 methylation MeCP2: components of the Sin3A HDAC complex Mbd3: component of the NuRD HDAC complex Mbd1: interacts with HDAC3.

Example: Mbd1 and HDAC3 are recruited by the PML-RARalpha hybrid protein (specialized transcription factor) to silence gene expression in cancer MBD1: interacts with the H3K9me3 HMTase SETDB1

DNA methylation collaborates with other chromatin modifying complexes to repress gene expression



Figure 9. Recruitment of corepressors by methyl-CpG binding proteins. A hypothetical transition between an active, nonmethylated gene promoter and a repressed promoter whose silence is attributable to DNA methylation, as mediated by complexes containing an MBD protein such as MeCP2 (gray shading). The transition phase represents an intermediate step during which transcription is silenced and DNA methylation occurs. MeCP2 is envisaged to recruit the NCoR histone deacetylase (HDAC) complex and histone lysine methyltransferase (HKMT) activity to the methylated sites. I

DNA METHYLATION IS ABUNANT IN THE GENOME AND ISSUBJECTED TO DRAMATIC ALTERATIONS DURING EMBRYOGENESIS



70%- 80% of CpG dinucleotides are methylated in the genome

DNA methylation levels are high in fertilized Oocytes that contain the paternal and maternal genome (carries characteristic methylation patterns)

Paternal and maternal methylation patterns are rapidly erased (exception: imprinted genes maintain paternal and maternal methylation information). \rightarrow the paternal and maternal methylation epigenome is cancelled

DNA methylation levels remain low during the first cell division events until the blastocyst stage

In the blastocyst stage cell differentiation programs are activated and genes need to be regulated on the epigenetic level \rightarrow DNA methylation is increasing (loss of DNMT1, DNMT3a or DNMT3b is lethal \rightarrow establishment and maintenance of DNA methylation is impaired)

70%- 80% of CpG di-nucleotides are methylated in the human genome! Remember only 2% of the genome encode for mRNAs 98% is noncoding DNA that contains a large proportion of transposable elements, repeat sequences, etc...

DNA METHYLATION IS REVERSIBLE: DNA DEMETHYLATION BY Tet-family proteins



Figure 6. Model of Tet-initiated DNA demethylation pathways. DNA methylation (5mC) is established and maintained by DNMT. 5mC can by oxidized by Tet family of dioxygenases to generate 5hmC, 5fC, and 5caC. Because the oxidized 5mC derivatives cannot serve as substrates for DNMT1, they can be lost by replication-dependent passive demethylation. 5hmC can be deaminated by AID/ APOBEC to become 5hmU, which together with 5fC and 5caC can be excised by glycosylases such as TDG, followed by DNA repair to generate C. Alternatively, a putative decarboxylase may convert 5caC to C.

Tet-family proteins mediate DNA demethylation neo-synthesized DNA template DNA strand DNMT1: GC-m3 5mC: 5-methylcytosine m3-CG Tet family oxidation proteins 5hmC: 5-hydroxymethylcytosine GC 5hmCG Tet family oxidation proteins 5fC: 5-formylcytisine GC 5fcCG Tet family oxidation proteins GC 5caC: 5-carboxylcytosine 5caCG

5mC, 5hmC and 5fC are abundant in the cell 5caC is present only at very low abundance

DNMT1 has exclusive specificity for 5mC

DNA METHYLATION IS REVERSIBLE: DNA DEMETHYLATION BY Tet-family proteins



Figure 6. Model of Tet-initiated DNA demethylation pathways. DNA methylation (5mC) is established and maintained by DNMT. 5mC can by oxidized by Tet family of dioxygenases to generate 5hmC, 5fC, and 5caC. Because the oxidized 5mC derivatives cannot serve as substrates for DNMT1, they can be lost by replication-dependent passive demethylation. 5hmC can be deaminated by AID/ APOBEC to become 5hmU, which together with 5fC and 5caC can be excised by glycosylases such as TDG, followed by DNA repair to generate C. Alternatively, a putative decarboxylase may convert 5caC to C.



The protein encoded by this gene belongs to the TDG/mug DNA glycosylase family. Thymine-DNA glycosylase (TDG) removes thymine moieties from G/T mismatches by hydrolyzing the carbon-nitrogen bond between the sugar-phosphate backbone of DNA and the mispaired thymine. With lower activity, this enzyme also removes thymine from C/T and T/T mispairings. TDG can also remove uracil and 5-bromouracil from mispairings with guanine. Interestingly, TDG knockout mouse models showed no increase in mispairing frequency suggesting that other enzymes, like the functional homologue MBD4, may provide functional redundancy. This gene may have a pseudogene in the p arm of chromosome 12.Additionally, in 2011, the human thymine DNA glycosylase (hTDG) was reported to efficiently excises 5-formylcytosine (5fC) and 5carboxylcytosine (5caC), the key oxidation products of 5-methylcytosine in genomic DNA. Later on, the crystal structure of the hTDG catalytic domain in complex with duplex DNA containing 5caC was published, which supports the role of TDG in mammalian 5methylcytosine demethylation.

Check textbooks: glycosilases cleave off bases from sugar \rightarrow apyrimidic/apurinic site \rightarrow BER pthway

DNA METHYLATION IS REVERSIBLE: ACTIVE AND PASSIVE DNA DEMETHYLATION



PASSIVE DNA DEMETHYLATION

Successive rounds of DNA methylation reduce the amount of 5mC. In this situation DNMT1 is excluded from the

Nucleus! (only transient presence of oocyte specufic version of DNMT1 at the 8 cell stage) MATERNAL GENOME: slow de-methylation of DNA

ACTIVE DNA DEMETHYLATION

Enzymatic activity rapidly de-methylates 5mC PATERNAL GENOME: fast de-methylation of DNA

- ightarrow In zygotes Tet3 is localized to the PATERNAL nucleus
- ightarrow Paternal DNA is demethylated
- → High levels of 5hmC: 5-hydroxymethylcytosine, 5fc: 5formylcytisine and 5caC: 5-carboxylcytosine were detected at high levels in the paternal nucleus
- → BER machinery concentrated in pronucleus

