L1.3

Chromatin Structures and composition

Chromatin is composed of

- DNA (genetic information)
- variously modified histone proteins (epigenetic info)
- DNA binding factors, enzymes, other proteins and noncoding RNAs (regulatory).



ENS, LYON

(Proteins in this cartoon are just examples)





Nucleosome DNA structure topology (supercoiling) topoisomerases



b







Molecular organization of heterochromatin versus euchromatin

Features distinguishing HC from EC

- Accessibility
- Nucleosome positioning
- Post-translational Modifications to Histones (PTM)
- Histone-binding proteins and modifying enzymes
- Histone variants
- DNA methylation
- Noncoding RNA
- Transcription factors

nucleosome

Are condensed (heterochromatic) and noncodensed (euchromatic) chromatin fragments (loci) distinguishable by <u>simple biochemical assay</u>?

General principle: enzyme accessibility.

The classical assay to detect the gross organization of chromatin at a specified locus: the **DNase I Hypersensitivity** Assay



Keratin K14 gene, analysis of the 5'-flanking region



Tratto da: Sinha et al., (2000), Mol Cell Biol, 20: 2543-2555.

Keratin K14 gene, analysis of the 5'-flanking region



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Keratin K14 gene, analysis of the 5'-flanking region



Tratto da: Sinha et al., (2000), Mol Cell Biol, 20: 2543-2555.

If accessibility is not the same in all parts of the chromatin, it may depend (also) from different distribution of nucleosomes ?

Questions that are relevant to genome organization:

Do nucleosome display preferred positions in the genome ? (are they just arrayed regularly spaced as they appear in pictures?)

Do nucleosomes display any kind of **specificity** for nucleotide sequence?

Are nucleosome mobile ? Can they translate position over DNA ? (are they stable on the same sequence they were assembled to ?)

Is nucleosome position **functional** to gene/genome regulation ?

«Passive» positioning

Protein-DNA contacts are mainly due to basic aminoacids – backbone phosphates



«Passive» positioning

Nevertheless, there is a level of sequence specificity: in general, A/T rich sequences are in touch, whereas C/G rich stay outside. This is due to intrinsic bending of DNA axis at different sequences.



«Assisted» nucleosome positioning:

- ATP-dependent chromatin remodeler
- Presence of DNA-binding factors, active transcription and other factors

How can nucleosome position on a given locus be assessed ?

1. In the test tube: accessibility to restriction enzymes



2. In vivo: Micrococcus Nuclease digestion

Micrococcal Nuclease I is a mixed endo-exonuclease.

MNase induces single-strand breaks and subsequently double stranded ones by cleaving the complementary strand in close proximity to the first break. MNase continues to digest the exposed DNA ends until it reaches an obstruction, such as a nucleosome or a very stably bound Transcription Factor. In appropriate condition, then, MNase releases fragments of approximately one nucleosome length (~147 bp), which are typically selected for sequencing in MNase-Seq experiments.

Nucleosome positioning analysis



2 different cases: mobile nucleosomes *versus* positioned nucleosomes

↓ MNase cuts

R.E. cuts





Is nucleosome position functionally relevant ?





Figure 1. Chromatin Structure at the PHO5 Promoter

Nucleosomes -1, -2, -3, and -4 are remodeled upon activation of the promoter by phosphate starvation conditions (Almer et al., 1986). The small circles mark UASp1 (open) and UASp2 (solid), which are Pho4-binding sites found by in vitro (Vogel et al., 1989) and in vivo (Venter et al., 1994) footprinting experiments. The positions are listed relative to the coding sequence (solid bar). T denotes the TATA box (Rudolph and Hinnen, 1987). The location of a Clal site at -275 relative to the coding region is shown.



FIG. 5. In vivo chromatin remodeling of episomal *PHO5*. Spheroplasts from the indicated strains were treated with micrococcal nuclease (MNase), and the DNA was purified and Southern blotted. (A) The blot was probed with probe A (Fig. 2). Data from each sample were quantified, and the distance from the top of the gel was graphed against the signal density. (B) The blot shown in panel A was stripped and reprobed with probe B (Fig. 2).



Nucleosome - histones

post-translational modifications (PTMs)

Table 1. Different Classes of Modifications Identified on Histones		
Chromatin Modifications	Residues Modified	Functions Regulated
Acetylation	K-ac	Transcription, Repair, Replication, Condensation
Methylation (lysines)	K-me1 K-me2 K-me3	Transcription, Repair
Methylation (arginines)	R-me1 R-me2a R-me2s	Transcription
Phosphorylation	S-ph T-ph	Transcription, Repair, Condensation
Ubiquitylation	K-ub	Transcription, Repair
Sumoylation	K-su	Transcription
ADP ribosylation	E-ar	Transcription
Deimination	R > Cit	Transcription
Proline Isomerization	P-cis > P-trans	Transcription

Overview of different classes of modification identified on histones. The functions that have been associated with each modification are shown. Each modification is discussed in detail in the text under the heading of the function it regulates.



Figure 4. The Types of Posttranslational Modifications Observed on the Core Histones

(A) The histone octamer portion of the nucleosome core particle is shown. The sites of modifications on marked. For clarity, the modifications are shown on one copy of each protein.

(B) The covalent modifications of the amino acids are shown.

В

Nomenclature





• the histone code





• the histone code



H3



This view was derived from single-gene studies.

PTMs are context-specific and quite dynamic

- Some PTMs always go together
- Other PTMs are mutually exclusive (e.g. Kac/Kme)
- Specific PTMs display effects on other PTMs (see next figure)

Question: are the two copies of each histone modified in the same way in the individual nucleosome?



Figure 2 Histone modification cross-talk. Histone modifications can positively or negatively affect other modifications. A positive effect is indicated by an arrowhead and a negative effect is indicated by a flat head (updated from reference [53]).

Textbook: Bannister & Kouzarides, 2011

PTMs are not only «grossly» distributed to HC/EC

They show finely distinct distributions in functional parts of chromatin, for examples in genes:





Figure 3 Distribution of histone modifications. Distributions of six modifications with respect to genes are schematically illustrated. TSS, transcription start site; TES, transcription end site. H3K4me3 is enriched around TSSs. H3K4me1 is nriched around enhancers and more downstream. H3K27ac is enriched around active enhancers and TSSs. In undifferentiated stem cells, both H3K4me3 and H3K27me3 (active and inactive marks, respectively) are enriched around TSSs on many genes. H3K27me3 is enriched around inactive TSS in somatic cells. H3K9me3 is broadly distributed on inactive regions. H3K27me3 and H3K9me3 are usually not colocalized. TSSs are generally devoid of nucleosomes.

interacting proteins
Interacting proteins are:

Enzymes introducing modificationsPTM writersEnzymes reversing modificationsPTM erasersProteins binding to specific PTMPTM readers

PTM = Post-Translational Modification

histone-modifying enzymes:

HAT- histone acetyltransferases HDAC – histone deacetylases HMT – histone methyltransferases histone demethylases histone kinases histone ribosyltrabsferases ubiquitin-transferases

(ATP-dependent remodelling enzymes)

Modifications are dynamic and reversible

Enzymes that Modify Histones	Residues Modified
Acetyltransferase	
HAT1	H4 (K5, K12)
CBP/P300	H3 (K14, K18) H4 (K5, K8) H2A (K5) H2B (K12, K15)
PCAF/GCN5	H3 (K9, K14, K18)
TIP60	H4 (K5, K8, K12, K16) H3 K14
HB01 (ScESA1, SpMST1)	H4 (K5, K8, K12)
ScSAS3	H3 (K14, K23)
ScSAS2 (SpMST2)	H4 K16
ScRTT109	H3 K56

Other enzymes exist with reduced specificity

Enzymes that Modify Histones	Residues Modified		
Deacetylases			
SirT2 (ScSir2)	H4 K16		

histone deacetylases (low specificity) HDAC: two classes

Class III: NAD-dependent deacetylases (Sir-Sirtuins)

Lysine Methyltransferase	writers	
SUV39H1		нзкэ
SUV39H2		НЗК9
G9a		H3K9
ESET/SETDB1		H3K9
EuHMTase/GLP		H3K9
CLL8		НЗК9
SpClr4		H3K9
MLL1		H3K4
MLL2		H3K4
MLL3		H3K4
MLL4		H3K4
MLL5		H3K4
SET1A		H3K4
SET1B		H3K4
ASH1		H3K4
Sc/Sp SET1		H3K4
SET2 (Sc/Sp SET2	2)	H3 K36
NSD1		H3K36
SYMD2		H3 K36
DOT1		H3K79
Sc/Sp DOT1		H3K79
Pr-SET 7/8		H4K20
SUV4 20H1		H4K20
SUV420H2		H4K20
SpSet 9		H4K20
EZH2		H3K27
RIZ1		НЗК9

Enzymes that Modify Histones	erasers	Residues Modified
Lysine Demethyla	ISES	
LSD1/BHC110		H3K4
JHDM1a		H3K36
JHDM1b		H3K36
JHDM2a		НЗК9
JHDM2b		НЗК9
JMJD2A/JHDM3/	4	H3K9, H3K36
JMJD2B		НЗК9
JMJD2C/GASC1		H3K9, H3K36
JMJD2D		НЗК9

Arginine Methlytransferases

CARM1	H3 (R2, R17, R26)
PRMT4	H4R3
PRMT5	H3R8, H4R3
Serine/Thrionine Kinases	
Haspin	НЗТЗ
MSK1	H3S28
MSK2	H3S28
СКІІ	H4S1
Mst1	H2BS14
Ubiquitilases	
Bmi/Ring1A	H2AK119
RNF20/RNF40	H2BK120
Proline Isomerases	
ScFPR4	H3P30, H3P38

readers



Figure 1. Recruitment of Proteins to Histones (A) Domains used for the recognition of methylated lysines, acetylated lysines, or phosphorylated serines.

readers



Figure 1. Recruitment of Proteins to Histones

(B) Proteins found that associate preferentially with modified versions of histone H3 and histone H4.

Causes and effects of PTMs

PTM are introduced in nucleosomes when Transcription Factors bind to regulatory elements through recruitment of Co-activatory complexes



PTMs influence nucleosome stability





In some cases, the PTM itself guides recruitment of multisubunit complexes, thus explaining «spreading» and maintenance of the chromatic status



How are chromatin interacting proteins studied?

Same as for Histone PTMs:

Chromatin Immunoprecipitation (ChIP)

Chromatin immunoprecipitation (ChIP)



histone isoform







H2AX LLGGVTIAQGGVLPNIQAVLLPKKTSATVGPKAPSGGKKATQASQEY H2Abbd LFNTTTISO--VAPGED Sequence Alignment of Variants of Histones H3 and H2A with the Known Secondary Structures of H3 and H2A Depicted on Top.

Upper:

The sequences of the conserved H3.3 and CENP-A variants. H3.3 differs by only a few residues. The arrows above the H3 Nterminal tail indicate the sites that form strands upon binding to chromodomains.

Lower

The sequences of the conserved H2A.Z, macroH2A, H2AX, and H2ABbd variants of H2A. The sequence of H2ABbd is most divergent, while others are closely related with some changes in the turn regions connecting the helices. CENP-A is an H3-like histone and is found only at centromeres over a stretch of 300-500 Kbp

CENP-A, unlike all other histones, is not replaced by protamines in sperm: chromatin status inheritance.

H3.3 is a variant of H3 showing only 4 aminoacids variation.H3.3 is deposited in chromatin also outside S-phaseH3.3 replaces H3 carrying H3K9me in re-activated genes

H2A.Z in S. cerevisiae is incorporated near silenced regions and inhibits the spread of heterochromatin.

If interested, see review by Talbert 2010

Modification of DNA: 5-methyl-cytosine

- **a. Cytonine-5-methylation** (+hydroxymethylation) is the only epigenetic modification concerning the DNA in higher organisms
- b. It occurs mostly at **CpG** dinucleotide (also at CpNpG in some cases, e.g. in plants and *see later in next chapter*)
- c. Methylation of CpG is observed at regulatory regions of **silenced** genes
- d. Hypermethylation is observed through the inactive X chromosome
- e. Housekeeping genes (constitutively expressed) show unmethylated CpG islands at promoters

Important: Cytosine methylation is common in Mammals and Plants, but is not used in the same way in *S. cerevisiae* and *C. elegans*

CpG methylation





- 1. Oxidative pathway (Tet enzymes)
- 2. Dilution (DNMT1 off during repeated cell division)



Figure 2. Mechanisms of DNA-methylation-mediated repression.
(a) DNA methylation in the cognate DNA-binding sequences of some (*not many, ndr*) transcription factors (TF) can result in inhibition of DNA binding.
By blocking activators from binding targets sites, DNA methylation directly inhibits transcriptional activation

From Klose & Bird, 2006



Figure 2. Mechanisms of DNA-methylation-mediated repression.
(b) Methyl-CpG-binding proteins (MBPs) directly recognize methylated DNA and recruit co-repressor molecules to silence transcription and to modify surrounding chromatin

From Klose & Bird, 2006



A family of **methyl-CpG-binding proteins** (MBPs). Six mammalian MBPs have been characterized so far:

- 1. MBD1 uses its methyl-binding domain (MBD) to bind methylated DNA sequences. In addition, MBD1 contains three zinc-binding domains (CxxC), one of which binds specifically to non-methylated CpG dinucleotides, and a C-terminal transcriptional repression domain (TRD).
- 2. MBD2 possesses an MBD that overlaps with its TRD domain, and a GR repeat at its N terminus.
- 3. MBD3 contains a well-conserved MBD domain that does not recognize methylated DNA owing to crucial amino acid changes.
- 4. MBD4 binds methylated DNA through an MBD domain and has a C-terminal glycosylase domain that is important for is function in DNA repair.
- 5. MeCP2 is the founding member of the MBD protein family and contains a conserved MBD domain and an adjacent TRD domain.
- 6. The sixth protein, Kaiso, is unrelated and is not shown here.

From Klose & Bird, 2006, TiBS 31: 89-97.

How to measure CpG methylation

- 1. Methylation-sensitive restriction enzymes
 - 1. Hpall*/Mspl (CCGG)
 - 2. Smal*/Xmal (CCCGGG)
 - 3. McrBC recognize 2 methylated (G/A)pC (50-1000bp apart)

- 2. Bisulfite conversion (Methyl-C is not modified , C is converted to U)
 - 1. Enzymatic analysis (e.g. Cobra-next slide)
 - 2. Cloning alleles and Sanger sequencing

Most common methods to detect CpC methylation are based on **bisulfite**, which converts C (but not methyl-C) to Uridine. Uridine in DNA is then replaced by T in the following PCR.

Then one can sequence frament(s) to detect changed bases.

In some cases, when a single CpG in a sequence has to be detected, we can use also bisulfite conversion followed by simple restriction analysis + PCR. Example:



Heterochromatin

- Facultative
- Constitutive

Facultative contains genes that are activated at some phases, then silenced.

Constitutive at centromeres, telomeres

The leading fact here is represented by **repetitive sequences**

Genome structure and composition