Histones

Replication dependent, canonical histones

- **- Gene organization**
- **Gene expression control mechanisms in S-phase**
- **Histone chaperons**

Replication independent histones – histone variants

- **- A list of histone variants**
- **- Histone variant at centromere**
- **Histone H3.3 a variant with multiple function**
- **Histone gammaH2AX and DNA damage**
- **Histone variants with chromosome specific accumulation**

Coordinated control of histone gene expression and incorporation into DNA

1. Histone and cell cycle regulation:

Maintaining a stable and balanced histone pool is of vital importance for appropriate gene regulation, cell cycle progression and genome stability. Excess of free histones has damaging effect \rightarrow precise control of histone gene expression required \rightarrow cell cycle

2. Controlled production of nucleosomes:

A. REPLICATION DEPENDENT REGULATION: **Canonical histone proteins**: mRNA non-polyadenylated; defined RNA processing mechanisms; expression regualted in histone locus body. Histone mRNA levels increase ca 40-fold increase at the level of histone mRNAs during S phase. At the end of S phase or DNA synthesis is interrupted, cells turned off histone transcription and histone mRNA levels declined rapidly. Replication dependent histone chaperons are linked to DNA polymerase and generate nucleosomes in S-Phase

B. REPLICATION INDEPENDENT REGULATION: histone variants: Classic polyadenyled RNAs Nucleosome need also be formed in G1, G2 phase when nucleosome arrays get disturbed (transcription, DNA damage, eviction, etc...). The second class of histones (non canonical histones) is composed of **histone variants that are expressed at a relatively** low level throughout the cell cycle, and are therefore regulated in a DNA replication-independent **manner. Histone variants have specialized functions!!**

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Histone gene clusters in vertebrates – Canonical histones

Histone genes are organized in histone gene clusters

Human:

Chr.6: major histone gene locus: HIST1 cluster: 45 **core histone genes ; 6 Histone H1 genes**

*Chr.1***:** minor histone gene locus HIST2 cluster: 6 **histone genes**

Chr.1: minor histone gene locus HIST3 cluster: 3 **histone genes**

Yeast:

2 copies for each core histone

The human and mouse major histone gene cluster. (A) The histone gene cluster on human chromosome 6p21-p22 is shown. The position and direction of transcription of the 55 histon genes in this region are indicated, with the genes for the five histone pro- teins indicated in the box. Only "real" genes are shown (defined as genes that contain the expected 3 end of histone mRNA). The portion of chromosome 6 is going (left to right) from the cen- tromere to telomere. HISTH4A is the first H4 gene starting from the right and the same is true for the other core histone genes. The numbers are nucleotides from the arbitrary start of the cluste rat 0. The regions where there are tightly grouped clusters of histone genes have been expanded. The scale of each section is indicated in kilobases (kb). The position of each of the histone H1 histone gene is indi-cated with the nomenclature H1a–e, H1t, and the symbols for the core his- tone genes are in the inset. The aster- isks indicate the position of genes pres- ent in human and not in mouse.

Histone genes are organized in clusters in vertebrates

Gene group: H3 histones (H3) @

A subgroup of @: "Histones"

The mapped domains of P68431, encoded by the H3C1 gene, an example gene within the group. [Source: Pfam & UniProt] Gene: H3C1, UniProt: P68431 136 amino acids

In vertebrates, there are a total of 10–20 genes encoding each of the core histone proteins.

Each of these genes encodes a unique mRNA, with a **distinct 5 - and 3 -UTR**, as well as **nucleotide changes** in the coding region.

All the histone **H4 genes encode the same protein**, **histone H3 genes** encodes variants with aa changes, there **10–12 different H2a and H2b** proteins are known

Canonical H3

Histone H3 variants with special function

https://www.genenames.org/cgi-bin/genefamilies/set/864

Coordinated control expression of replication dependent histones

The core histone gene promoters contain specialized DNA elements that enable cisregulation of histone gene expression: UAS (upstream activating sequence) and NEG (negative regulation of expression) **Divergent arrangement** of histone promoters allows coordinated gene expression to get equal amount of all four core histones.

elements that required for histone gene expression. Histone H2B promoter contains an octamer element (5'-ATTTGCAT-3'), which is bound by transcription activator Oct-1 (octamerbinding factor 1). Histone H4 promoter contains subtype-specific regulatory elements(SSREs)

S. cerevisiae S. cerevisiae

Structure of yeast canonicalhistone genes

Structure of yeast

canonical histone genes

Structure of yeast histone variants H2AZ and CenH3.

H2AZ and CenH3.

Structure of yeast histone variants

Structure of

H. sapiens

CONTROL OF HISTONE EXPRESSION IN S-PHASE (replication dependent histones)

Histone synthesisis limited to S-Phase

The transcription of histone gene takes place in a subnuclear organelle termed the **histone locus body** (HLB), containing factors required for the processing of histone pre-mRNAs which have an *unusual mRNA* **structure, with a 3'UTR that forms a stem-loop structure instead of a polyA tail**

G1 à **S transition**:

- Transcription increases 5-fold during the G1 to S phase transition
- RNA processing enhanced RNA levels 8-fold
- Total increase: 30–40-fold increase of mature histone mRNAs.

S**→**G2 transition:

At the end of S phase, the half-life of these histone mRNAs drops dramatically in response to a destabilization of the specific 3' ends.

Human cells in S-phase IF staining for NPAT combined with

CONTROL OF HISTONE EXPRESSION IN S-PHASE – TRANSCRIPTIONAL REGULATION

NPAT – TRANSCRIPTIONAL ACTIVATION

Histone synthesisis limited to S-Phase

Activation of histone H2B. Oct-1 binds to octamer elementsin H2B promoter. During S phase, activated cyclin E/CDK2 complex phosphorylates NPAT. In combination with NPAT, Oct-1 recruits OCA-S to H2B promoter to activates the expression of H2B.

Activation of histone H4. HiNF-P binds to SSRE within H4 promoters and recruits NAPT and RNA polymerase II to activate gene transcription. NPAT recruits the Tip60 histone acetyltransferase complex to acetylate histone H4 at the G1/S-phase transition. At the end of S phase, the tyrosine kinase WEE1 is recruited to histone promoters to phosphorylate H2B tyrosine 37, which evicts NPAT and RNA polymerase II and instead recruits HIRA to repress histone gene expression.

CDK2 and the cell cycle:

Entry into S-phase is triggered by the activity of the G1-S Cyclin complex. CyclinE/Cdk2.

After CyclinE/Cdk2 activity has reached its peak in early S-phase, CyclinE/Cdk2 activity drops due to the degradation of the essential CyclinE component

NPAT, Nuclear Protein Ataxia-Telangiectasia Locus; RNAPII, RNA polymerase II; TRRAP, transformation/transa ctivation domain-associated protein; SSRE, subtype-specific regulatory elements; OCA-S, Oct-1 co-activator in S-phase; HiNF-P, histone nuclear factor P; TBP, TATA-box binding protein.

CONTROL OF HISTONE EXPRESSION IN S-PHASE – REPLICATION DEPENDENT HISTONE RNA PROCESSING (RHP)

Schematic view of the replication-dependent histone RNA processing **complex**

2 conserved mRNA processing signals 50 nucleotides downstream of the histone ORF stop codon:

- stem-loop element: recognized by stem-loop binding protein (**SLBP**)
- purine-rich histone downstream element (HDE): paired by the U7 snRNP

A 100 kDa zinc finger protein (ZFP-100) forms a connection between these two factors and stabilizes the complex.

The U7 snRNP contains a ring-like structure composed of five Sm proteins and the U7- specific Sm-like proteins, Lsm10 and Lsm11.

A N-terminal extension of Lsm11 additionally binds the 68 kDa subunit of mammalian cleavage factor I (CFIm68) and FLASH. Together, Lsm11 and FLASH provide a docking platform for the **Heat Labile Factor (HLF**) which consists of proteins that are also involved in cleavage/polyadenylation (CPA): symplekin (SYMPK), Cleavage Stimulation Factor 64 kDa subunit (CstF64) and all six subunits of Cleavage and Polyadenylation Factor (CPSF). **CPSF-73** is the endonuclease that cleaves the pre-mRNA after a CA dinucleotide.

While SLBP binds to the 50 side of the hairpin, the 30 side is occupied by 30 **hExo**, an exonuclease that trims the 30 end of the mRNA after cleavage and may also be involved in histone mRNA degradation.

If the canonical histone processing site is not recognized, a downstream **polyadenylation signal** can be used by the CPA machinery, resulting in polyadenylated histone transcripts. Poly-adenylated transcripts remain nuclear and get rapidly degraded by the nuclear exosome. Only a small fraction can be detected on polysomes.

CONTROL OF HISTONE EXPRESSION IN S-PHASE – REPLICATION DEPENDENT HISTONE RNA PROCESSING (RHP)

HBP (SLBP) - REGULATION OF RNA METABOLISM

HBP (histone RNA binding protein; also called SLBP) protein itself is cell cycle regulated.

SLBP mRNA is synthesized constantly throughout the cell cycle, but **HBP becomes translated just prior to S-phase entry and the protein is degraded at the end of S- phase**

Histone mRNA 3'-end processing requires the RNA-binding protein HBP (also called SLBP), which binds to the conserved hairpin structure in histone pre-mRNA, and the U7snRNP, which binds to a sequence element downstream of the cleavage site.

Together with other factors they position the nuclease CPSF73 for cleavage to produce histone mRNA ending immediately after the stem loop.

After nuclear export. HBP interacts with SLIP1 and other translation initiation factors to form a closed-loop structure for efficient translation.

This structure is disrupted, presumably when histone mRNA decay is initiated, for example at the end of S-phase.

Addition of an oligo(U) tail by the terminal uridylyl transferase ZCCHC11 is an early step in decay, which involves decapping followed by $5' \rightarrow 3'$ degradation by Xrn1 or $3' \rightarrow 5'$ decay by the exosome.

FORMATION OF SUBNUCLEAR STRUCTURE = HISTONE LOCUS BODY

Histone locus bodies (HLBs) are subnuclear sites containing the clustered RD histone genes and components of the corresponding gene expression and processing machinery.

A. In G1, only two HLBs corresponding to the major histone cluster on chromosome 6 can be stained by antibodies against HLB marker proteins such as FLASH. As NPAT is not phosphorylated, the interaction with FLASH is weak. The level of SLBP is low, and the U7 snRNP, interacting with hnRNP UL1, inhibits histone transcription. Transcription is minimal and RNA 3' processing is inefficient so that most transcripts from the Replication dependent histone mRNAs become polyadenylated at cryptic polyA signals downstream from their regular cleavage sites.

B. In S phase, NPAT gets phosphorylated by

cyclinE/Cdk2 which stabilizes the interaction with FLASH. The concentration of factors in HLBs increases, and two more HLBs corresponding to the minor cluster on chromosome 1 can be detected. SLBP is upregulated, and FLASH interacts with the U7 snRNP, hence providing a platform for the recruitment of the now activated HLF.

CONTROL OF HISTONE EXPRESSION IN S-PHASE

DROSOPHILA: HERS and Su(var)3-9

Drosophila: The histone gene-specific epigenetic repressor in late S-phase (HERS) protein becomes phos- phorylated by the late S-G2 Cyclin complex CyclinA/Cdk1, which localizes it to the histone genes where it acts to silence histone genes after S-phase .

Cdk-activated HERS silence histone gene expression in late S phase through recruitment of Su(var)3-9/HP1 repressor complex.

Drosphila: also presence of HLB

Ito et al. 2012 Mol Cell. Epigenetic Silencing of CoreHistone Genes by HERS in Drosophila

Figure 4. Distribution of old and new nucleosomes at a replication fork. Old nucleosomes (gray disks) are randomly distributed behind the replication fork and new nucleosomes (cyan disks) are deposited in the gaps. CAF-1-mediated nucleosome assembly is depicted on the leading and lagging strand in magnification. DNA polymerase (green); replication processivity clamp, PCNA (gray ring); histone H3-H4 tetramers (cyan); newly synthesized DNA (red lines).

REPLICATION COUPLED HISTONES

REPLICATION COUPLED (RCs) HISTONES: H2A, H2B, H3, H4 Are incorporated into new and old DNA strand during DNA replication

Chromatin assembly factor 1 (CAF-1) is a HISTONE CHAPERON that is associated with PCNA. \rightarrow Facilitates the formation of new nuclosomes The assembly of a nucleosome consists of the loading of an (H3-H4)2 tetramer (tetrasome) that is followed by the addition of 2 H2A-H2B dimers.

REPLICATION INDEPENDENT (RIs) HISTONES: Are incorporated independently of DNA replication

RI histones require the displacement of a preexisting nucleosome unit (active displacement or "loss")

RI histones can reset the epigenetic state of a pre-exising nucelosome

Histone variants with high similarity to "normal" histone are also incorporated into cannonical octamers during replication. However concentration of variants is very low \rightarrow no big effect on chromatin structure

HOWEVER: HISTONE CHAPERONS EXIST THAT ENSURE CONCENTRATED INCORPORATION OF HISTONE VARIANTS AT DEFINED SITES \rightarrow **CONTROLLED** LOCAL CONCENTRATION \rightarrow ALTERATION OF CHROMATIN STRUCTURE

Incorporation of replication coupled histones by CAF-1 histone chaperone

Humans CAF-1 subunits: p150, p60, and p48, Budding yeast CAF-1 subunits: Cac1, Cac2, and Cac3

CAF-1 p150 interacts with PCNA

- (a) The nucleosome assembly mechanism of CAF-1 is activated by H3-H4 binding, which releases the WHD domain from an intramolecular interaction with the acidic region on Cac1.
- (b) DNA binding promotes the association of two CAF-1•H3-H4 complexes to join the histones into a (H3–H4)2 tetramer
- (c) In the presence of DNA of sufficient length, the (H3–H4)2 histones are directly sequestered from CAF-1.
- (d) (H3-H4)2 are transferred to the DNA to form the **tetrasome**, and the WHD rebinds to the now free acidic region, resulting in its dissociation from DNA.

H2A-H2B can spontaneously associate with tetrasomes in vitro and because CAF-1 itself has significantly lower affinity for H2A-H2B compared to H3-H4, it appears that the **primary role of CAF-1** is to promote **the formation of an ordered (H3-H4), •DNA** complex, the tetrasome

Mattiroli F: Elife. 2017: PMID: 28315523

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REPLICATION INDEPENDENT (RIs) HISTONES: - HISTONE VARIANTS

Figure 1. Histone variants. Protein domain structure for the core histones (H3, H4 H2A, and H2B), linker histone H1, and variants of histones H3 and H2A. The histonefold domain (HFD) is where histone dimerization occurs. Regions of sequence variation in histone variants are indicated in red. WHD, winged-helix domain.

REPLICATION INDEPENDENT (RIs) HISTONES: Are incorporated throughout the cells cycle and independently of **DNA** replication

RI histones require the displacement of a pre-existing nucleosome unit

RI histones can reset the epigenetic state of a pre-exising nucelosome

Assure nucleosome organization in interphase cells or terminally differentiated cell **(transcription, remodeling)**

Used in defined biological process

Histone variants with high similarity to "normal" histone can also incorporated into cannonical octamers during replication. However concentration of variants is very low \rightarrow no big effect on chromatin structure

HOWEVER: HISTONE CHAPERONS EXIST THAT ENSURE CONCENTRATED INCORPORATION OF HISTONE VARIANTS AT DEFINED SITES \rightarrow **CONTROLLED** LOCAL CONCENTRATION \rightarrow ALTERATION OF CHROMATIN STRUCTURE

REPLICATION INDEPENDENT (RIs) HISTONES: - HISTONE VARIANTS

Loop

 $H2A$

 $H2A$

Figure 1. Histone variants. Protein domain structure for the core histones (H3, H4, H2A, and H2B), linker histone H1, and variants of histones H3 and H2A. The histonefold domain (HFD) is where histone dimerization occurs. Regions of sequence variation in histone variants are indicated in red. WHD, winged-helix domain.

1. H3 VARIANTS AT THE CENTROMERE: CENP-A: A HISTONE VARIANT THAT DEFINES EUKARYOTIC CENTROMERES

Centromere protein **C** 1 is a centromere autoantigen and a component of the inner kinetochore plate. The protein is required for maintaining proper kinetochore size and a timely transition to anaphase.

Centromere protein B is a highly conserved protein that facilitates centromere formation. It is a DNA-binding protein that is derived from transposases of the pogo DNA transposon family. It contains a helix-loop-helix DNA binding motif at the N-terminus, and a dimerization domain at the C-terminus. The DNA binding domain recognizes and binds a 17-bp sequence (CENP-B box) in the centromeric alpha satellite DNA. This protein is proposed to play an important role in the assembly of specific centromere structures in interphase nuclei and on mitotic chromosomes. It is also considered a major centromere autoantigen recognized by sera from patients with anti-centromere antibodies.

CENP-A: RI histone variant, incorporated by the cenH3 specific histone HJURP histone chaperone

Kinetochore assembles on CENP-A (Mechanism unknown) **CENP-C** and **CENP-B** are non histone centromere founder proteins

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H3.3: MAINTAINING NCULEOSOME STRUCTURE IN NON S-PHASE CELLS

H3.3 INCORPORATION INTO GENOME IS HIGHLY DYNAMIC: AN EXAMPLE

H3.3 is essential to fill nucleosome gaps in terminally differentiated cells

Gene expression is not altered \rightarrow H3.3 compensates for H3

Histone Chaperons: Pathways if RI incorporation of H3.3 into chromatin

Figure 2. During RC assembly, the ASF1 chaperone is thought to transfer newly synthesized soluble H3.1–H4 dimers to CAF-1 through direct interactions with its p60 subunit (Tyler et al. 2001; Mello et al. 2002). CAF-1 would then facilitate the assembly of a central (H3.1-H4)₂ tetramer to which two H2A-H2B dimers are juxtaposed by other chaperones to complete a core nucleosomal unit. Similarly, during RI chromatin assembly, ASF1-bound H3.3-H4 would be transferred to HIRA for tetramer formation (Green et al. 2005). Note, however, that the deposition of H3.3 in the Drosophila male pronucleus is HIRA-dependent, but ASF1-independent (Bonnefoy et al. 2007) in the histone-rich fertilized egg. Novel alternate pathways for H3.3-H4 deposition include the DAXX chaperone coupled to the ATRX ATP-dependent chromatin remodeler (Drané et al. 2010; Goldberg et al. 2010), as well as targeted H3.3 deposition to regulatory elements by DEK (Sawatsubashi et al. 2010). Boxes highlight queries to consider in future studies.

elements

1. H3 VARIANTS: H3.3 A HISTONE VARIANT MARKING, TRANSCRIBED, ACTIVE CHROMATIN

Figure 7. H3.3 preferentially localizes to actively transcribed regions of Drosophila polytene chromosomes. DAPI staining (red) shows the DNA banding pattern (left), and H3.3-GFP (green) localizes to interbands (middle), which are sites of RNA Pol II localization. The merge (Schwartz and Ahmad 2005) is shown on the right. In each image, the shorter arrow points to a decondensed interband that is enriched in H3.3, and the longer arrow points to a condensed band that lacks H3.3.

H3.3 can form heterodimer with $HA \rightarrow H3.3-H4$ is incorporated by RI assembly

Chromatin is in flux during in G1/G2 and DNA is transcribed \rightarrow frequent RI chromatin reassembly \rightarrow enrichment of H3.3 chromatin in active genes

Active genes are defined by epigenetic writers (i.e. H3K4 HMTse) that impose active histone marks on H3.3.

Additional positive effect: eventual repressive modifications (H3K9me3) are eliminated by exchanging H3-mod for H3.3 unmodified

H3.3 carries same modification like classic H3

Gene induction triggers histone replacement of H3 with H3.3

Brian E. Schwartz, and Kami Ahmad Genes Dev. 2005;19:804-814

Gene induction triggers histone replacement. Heat-shock induction triggers rapid puffing and transcription of the HSP70 genes at polytene bands 87A and 87C. Polytene chromosomes from larvae that contain a constitutively expressed H3-GFP construct (A–D,J,K), or a constitutively expressed H3.3core-GFP construct (E–I) were induced for the indicated times (in minutes) and stained with antibodies against **phosphorylated RNA polymerase II (blue**), which marks activated puffs. Arrows in A and E indicate the positions of the HSP70 loci at 87A and 87C before induction, while the puffs (B–D,F–H) are bracketed. Asterisks indicate chromosome bands that were used as internal standards for quantifying the summed intensities of histone-GFP signals in puffs. (B,C) **Puffs contain some H3 in the first 5 min of induction, but have much less by the time puffs reach their maximal size (D). (F–H) H3.3core-GFP in expanding puffs rapidly increases. (I) After 20 min of heat-shock induction, many active HSP puffs (arrows) have large amounts of H3.3core-GFP, while staining throughout the arms is undiminished.** (J,K) Histone H3 modifications that mark heat-shock puffs do not overlap with H3-GFP**. DNA is in red, and histone-GFP in green.**

(A) Fertilization induces the transition from metaphase II to anaphase II, leading to completion of oocyte meiosis and emission of a second polar body (which usually degenerates). The sperm nucleus decondenses, so the fertilized egg (zygote) contains two haploid nuclei (male and female pronuclei). In mammals, the pronuclei replicate DNA as they migrate toward each other. They then initiate mitosis, with male and female chromosomes aligning on a common spindle. Completion of mitosis and cytokinesis thus gives rise to a twocell embryo, with each cell containing a diploid genome.

Early development of the embryo. Division of the fertilized egg results in the formation of daughter cells called blastomeres, which form a ball of cells called a morula. After six or so divisions, a cavity appears at one end of the morula to form the blastocyst. Until this stage, each of the cells divides evenly to produce two daughter cells, each of which retains the same characteristics, that is, they are totipotent. The inner cell mass of the blastula is the source of ESCs (see below). Invagination of one pole of the blastula leads to formation of the primitive germ cell layers, with formation of the gastrula. During gastrulation and later formation of the fetus, the daughter cells lose potential as they gain specialized function. This process is known as determination

EPIGENTIC REPROGRAMMING Gene activation

Representative pictures of MII oocytes prior to (A , F , and K) and after insemination using ICSI (Day 1: B , G , and L ; Day 3: C , H , and M ; Day 5: D , I , and N ; Day 7: E , J , and O). After insemination, the presence of two pronuclei **(PN) and two polar bodies (PB) was checked to confirm fertilization (B , G , and L). Some embryos arrested at very early stages(e.g., four cell; D and E), whereas others continued** cleavaging until the morula stage (J and O), **when they arrested.**

https://embryology.med.unsw.edu.au/embryology/index.php/Oocyte_Development

Polar Body: The breakdown of the germinal veside indicates a resumption of meiosis and the extrusion of the first polar body (1 PB) indicates completion of the first meiotic division in human oocytes. The polar body is a small cytoplasmic exclusion body formed to enclose the excess DNA formed during the oocyte (egg) meiosis and following sperm fertilization. There are 2-3 polar bodies derived from the oocyte present in the zygote, the number is dependent upon whether polar body 1 (the first polar body formed during meiosis 1) divides during meiosis 2. This exclusion body contains the excess DNA from the reductive division (the second and third polar bodies are formed from meiosis 2 at fertilization). These polar bodies do not contribute to the future genetic complement of the zygote, embryo or fetus. Recent research in some species suggest that the space formed by

the peripheral polar body (between the oocyte and the zona pellucia) can influence the site of spermatozoa fertilization.

MII oocytes injected with flag-tagged H3.1 or H3.2 or H3.3 variants \rightarrow In vitro fertilization \rightarrow Allow 1-cell and 2 cell development \rightarrow Anti-flag IF

P: paternal pronucleus M: maternal pronucleus

H3.2: canonical H3 inserted into pronuclei $+ 2$ cell embryo H3.3 RI histone: inserted into into pronuclei $+2$ cell embryo

2 cell stage \rightarrow blastocyst

H3.1: incorporation into blastocyst chromatin (eu+heterochromatin)

H3.2: incorporation in 2 cell and blastocyst chromatin (eu+heterochr.)

H3.3: incorporation in 2 cell embryos (hetero + euchromatin) **incorporation in blastocyst euchromatin** (DNA not condensed)

 \rightarrow ACTIVATION OF GENE EXPRESSION PARALLELED BY H3.3 INCORPORATON

Reprogramming by dynamic, replication dependent replacement of histone H3 variants with H3.3

MII oocytes fertilized \rightarrow Develop to 1 cell stage \rightarrow injected with siRNAs \rightarrow Injection of Flag tagged variants In 1 blastomere of 2 cell embryo \rightarrow analysis of morula cells

H3.3 incorporation not affected (increases)

 \rightarrow H3.3 is an RI histone (incorporated by HIRA; DAXX or DEK)

2. H3.3 is enriched at repeat elements and contributes to genomic stability

CHROMATIN REGULATION OF REPETITIVE DNA

In addition to genes, recent studies have shown that H3.3 is enriched at repetitive regions such as

- **Telomeres**
- **Pericentric repeats**
- **Endogenous retroelements**

Deposition at these regions is mediated by the **ATRX/DAXX complex**.

This complex is mutated in many human cancers, with links to a process known as alternative lengthening of telomeres, or ALT.

DAXX - ATRX insert H3.3 at repeat elements: at heterochromatic telomeres and centromeric repeats

Daxx: histone chaperon

Figure 8. Model for maintenance of histone modifications by the concerted action of multiple chromatin regulators via RI replacement with H3.3. We address the question of how a histone modification can be inherited when a nucleosome is lost and replaced. (A) The Suv39h H3K9 methyltransferase (an ortholog of fly Su(var)3-9) is recruited by HP-1 protein, which binds specifically to methylated H3K9. To perpetuate this mark when the nucleosome turns over, we speculate that the ATRX ATPase is recruited to the site via its ATRX-DNMT3-DNMT3L (ADD) domain, which binds with high specificity to methylated H3K9 on tails that entirely lack H3K4 methylation (because there are no H3K4 methyltransferases in this region of the genome). (B) ATRX provides the energy of ATP and works together with the H3.3-specific DAXX histone chaperone complex to incorporate the new nucleosome (Goldberg et al. 2010), or half-nucleosome in the case of partial eviction (Xu et al. 2010). The high local concentration of Suv39h results in a new nucleosome with the same H3K9 methylation as the nucleosome that was lost.

H3.3 also enriched at tandem repeat containing sequences (i.e telomere or centromere, ...)

ATRX binds to H3-tails that are H3K4me0 and H3K9me3

ATRX is enriched at telomeres and centromeric repeats

ATRX brings in DAXX \rightarrow H3.3 RI assembly

There is transcription at telomeres an centromeres \rightarrow nucleosome turnover

A model for DAXX-ATRX mediated suppression genomic instability: telomeres as an example

Loss of ATRX/DAXX function \rightarrow lack of H3.3 incorporation \rightarrow abnormal chromatin \rightarrow breaks \rightarrow induction of homologous recombination

= ALT (alternative lengthening of telomeres)

A model for DAXX-ATRX mediated suppression genomic instability: telomeres as an example

G-quadruplex DNA formed on lagging strand

Figure 6 | Model for ATRX-mediated suppression of the ALT pathway. ATRX together with DAXX deposits histone H3.3 at telomeres, which in turn may facilitate DNA replication through G-quadruplex sequences. The presence of G-quadruplex structures in an ATRX null turnour cell leads to replication fork stalling and collapse, providing a substrate for MRN-dependent homologous recombination and maintenance of telomere length through ALT. ATRX additionally interacts with the MRN complex, facilitating its distribution away from PML bodies and telomeres, further limiting HR.

The MRN complex (MRX complex in yeast) is a protein complex consisting of Mre11, Rad50 and Nbs1 (also known as Nibrin in humans and as Xrs2 in yeast). In eukaryotes, the MRN/X complex plays an important role in the initial processing of double-strand DNA breaks prior to repair by homologous recombination or non-homologous end joining.

ROLE OF H3.3 IN TUMORFORMATION

ROLE OF H3.3 IN TUMORFORMATION

PanNETs: human pancreatic neuroendocrine tumors: 40% have loss of function mutations in ATRX or Daxx

Loss of ATRX or Daxx is associated with the induction of recombination at telomeres \rightarrow telomerase independent telomere elongation

Pediatric glioblastoma; Mutations in H3.3 G34R/V associated with ATRX loss \rightarrow tumor formation

ROLE OF ATRX/DAXX AND H3.3 IN Pancreatic neuroendocrine tumors (PanNETs)

Pancreatic neuroendocrine tumors (PanNETs, PETs, or PNETs), often referred to as "islet cell tumors", or "pancreatic endocrine tumors" are neuroendocrine neoplasms that arise from cells of the endocrine (hormonal) and nervous system within the pancreas. PanNETs are a type of neuroendocrine tumor, representing about one third of gastroenteropancreatic neuroendocrine tumors (GEP-NETs). Many PanNETs are benign, while some are malignant. Aggressive PanNET tumors have traditionally been termed "islet cell carcinoma".PanNETs are quite distinct from the usual form of pancreatic cancer, the majority of which are adenocarcinomas, which arises in the exocrine pancreas. Only 1 or 2% of clinically significant pancreas neoplasms are PanNETs.

ROLE OF ATRX/DAXX AND H3.3 IN PanNETs

Figure 1.

Representative examples of PanNETs assessed by DAXX and ATRX IHC and telomere-specific FISH. A, PanNET with preserved nuclear expression for both DAXX (B) and ATRX (C) and absence of the ALT phenotype (D). E, PanNET with DAXX loss (F), but preserved expression for ATRX (G). The loss of DAXX expression correlated with the presence of large, ultrabright intranuclear foci by telomere-specific FISH, consistent with ALT (H). I, PanNET with preserved expression for DAXX (J), but ATRX loss (K) and ALT positive (L) by telomere-specific FISH.

ROLE OF ATRX/DAXX AND H3.3 IN PanNETs

^alndicates that the value in question is statistically significantly better than the relevant control, where significance is defined by $P < 0.05$.

ALT and LOSS OF DAXX/ATRX mediates PanNET aggressivenss

disease-free survival

In cancer, the length of time after primary treatment for a cancer ends that the patient survives without any signs or symptoms of that cancer. In a clinical trial, measuring the disease-free survival is one way to see how well a new treatment works. Also called DFS, relapse-free survival, and RFS.

Histones

Replication dependent, canonical histones

- **- Gene organization**
- **Gene expression control mechanisms in S-phase**
- **Histone chaperons**

Replication independent histones – histone variants

- **- A list of histone variants**
- **- Histone variant at centromere**
- **Histone H3.3 a variant with multiple function:**
- **Histone H2AX and DNA damage**
- **Histone variants with chromosome specific accumulation**

ROLE OF H2A.X IN DNA DAMAGE RESPONSE

HISTONE H2A VARIANTS

DNA double-strand breaks (DSBs) are cytotoxic damages that can be repaired either by the homologous recombinational repair (HR) pathway or by the non-homologous end-joining (NHEJ) pathway. NHEJ, although faster than HR, is less accurate. The early divergent step between the two pathways is end resection, and this step is regulated by numerous factors. In particular, BRCA1 and 53BP1 play a role in determining the balance between the two pathways. 53BP1 restricts resection and promotes NHEJ.

The MRN complex binds avidly to double-strand breaks both in vitro and in vivo and may serve to tether broken ends prior to repair by non-homologous end joining or to initiate resection prior to repair by homologous recombination. Recruits ATM. The MRN complex also participates in activating the checkpoint kinase ATM in response to DNA damage.

 γ -H2AX

Cells cut with laser \rightarrow **DNA** breaks across nucleus \rightarrow **phosphorylation of H2AX at Ser 139 by ATM, ATR** and DNA-PK = γ H2AX

ACTIVATION OF DNA DAMAGE RESPONSE

H2AX is distributed throughout the genome. *DNA damage -> phosphorylation of H2AX* by ATM/ATR/DNA PK that spreads max 1 Mb from DNA damage site

=DNA DAMGE FOCI

DNA Damage \rightarrow Exposure of DNA breaks \rightarrow activation of PI3K family kinases (ATR/ATM/DNA PK) \rightarrow phosphorylation cascade

DNA Damage Foci in genomic DNA of cell nucleus

Histone H2AX is a substrate of several phosphoinositide 3-kinase-related protein kinases (PIKKs), such as ATM (ataxia teleangiectasia mutated), ATR (ATM and Rad3-related), or DNA-dependent protein kinase (DNA-PK). Serves as a platform to recruit DNA damage response factors

ROLE OF H2A.X IN DNA DAMAGE RESPONSE

Spatial organization of DDR protein accumulation at DNA DSBs.

(*A*) DDR signal spreading. DDR proteins initially accumulate at DSB sites and then spread at distance via a positive feedback loop involving MDC1, which binds γH2AX, the MRN complex, and ATM kinase, which phosphorylates additional H2AX molecules further away from the break site.

(*B*) Regional distribution of DDR proteins around DSBs. Factors involved in ATR signaling accumulate proximal to the break site on ssDNA generated by DNA end resection, while ATM signaling factors localize on flanking chromatin regions.

ROLE OF H2A.X IN DNA DAMAGE RESPONSE

γ**H2AX** 53BP1 MRE11 P-ATM

Several lines of evidence suggest the critical role of H2AX phosphorylation at DSB sites for nuclear foci formation and induction of DSB repair.

- (i) H2AX-knockout cells manifested impaired recruitment of NBS1, 53BP1, and BRCA1 to irradiation-induced foci
- (ii) H2AX loss of function mouse thymocytes show an increase in chromosomal aberrations. Mouse embryonic stem (ES) cells deficient in H2AX phosphorylation have alterations in efficiency of DNA repair by NHEJ or HR. As a result of these defects in DNA damage repair, cells show increased sensitivity to DNA damage.
- (iii) H2AX knock-out mice show male-specific infertility and reduced levels of secondary immunoglobulin isotypes, suggesting defects in class switch recombination (CSR). It was shown that efficient resolution of DSBs induced during CSR in lymphocytes requires H2AX, and its absence is associated with chromosome abnormalities involving the immunoglobulin locus.

These facts suggest that H2AX might serve as a docking site for DNA damage/repair proteins and functions to promote DSB repair and genome stability.

 v -H2AX

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ROLE OF macroH2A AND H2A.B ON THE X-CHROMOSOME

MacroH2A and H2A.B and X inactivation

X inactivation is associated with the exchange of H2A variants

A B \overline{c} H₂A.B is excluded from Xi

Figure 11. H2A variants and the inactive X chromosome of human females. (A) macroH2A (red) stains discrete regions of the inactive X chromosome that alternate with a marker for heterochromatin (histone H3K9me3). (B) H2A.B (green) is excluded from the inactive X chromosome (red dot with arrowhead pointing to it). (C) Same nucleus as in B, but stained with DAPI to show chromatin. (A, Reprinted, with permission, from Chadwick and Willard 2004, @ National Academy of Sciences; B,C, reprinted, with permission, from Chadwick and Willard 2001, @ 2001 The Rockefeller University Press. Originally published in Journal of Cell Biology 152: 375-384. doi: 10.1083/jcb.152.2.375.)

macroH₂A is enriched at the Xi

ROLE OF H2A.Z IN CHROMATIN DYNAMICS

Different heterodimers possible:

- 1. $H2A.Z/H2B + H2A.Z/H2B = ZZ$
- 2. H2A.Z/H2B + H2A/H2B
- 3. $H2A/H2B + H2A/H2B$

Inserted of RI H2AZ into chromatin by:

- INO80 and SWR-C complexes (**chromatin remodelers with ATPase domain**);
- **histone chaperon**: human YL1; yeast: Chz1

H2A.Z shows less adherence to DNA and is frequently localized **aroud** the transcriptional start site:

Yeast: ZZ: 32%

 $7A.24%$

 $AA: 44%$

\rightarrow >50% of genes have a H2A.Z at transcription start sites! WHY?

NUCLEOSOME DEPLETED REGIONS (NDRs)

Figure 2 The different physiological outcomes of nucleosome sliding. (a) Nucleosome-remodeling enzymes can introduce and maintain regular spacing of nucleosomes. (b) The phasing of nucleosomal arrays with respect to a nucleosome-depleted region (NDR), a prominent feature of promoters, depends on nucleosome-remodeling enzymes. (c) Nucleosomesliding activity regulates the accessibility of DNA sequences by positioning individual nucleosomes.

NUCLEOSOME DEPLETED REGIONS (NDRs)

Nucleosome-depleted regions at the 5' and 3' ends of transcripts.

As observed in eukaryotes, Both 5' and 3' end profiles are overlaid in this figure for comparison. The 5' NDR is, on average, more depleted and longer. DOI:http://dx.doi.org/10.7554/eLife.00078.005

ROLE OF H2A.Z IN CHROMATIN DYNAMICS

Figure 9.3 DNA strands separate to form a transcription bubble. RNA is synthesized by complementary base pairing with one of the DNA strands.

At promoter and +1 position high turn-over of nucleosomes

The initiation site of transcription is nucleosome-free

Nucleosomes containing H2A.Z (or H3.3) **are more instable**; leaving the promoter and +1 position more flexible

During G1, S, G2, M phase nucleosome at +1 position (transcriptional start site) are **not present.**

Around these position H2A.Z is abundant! In this way the TSS acquires an open configuration - Also when cell pass trough M-Phase

=TRANSCRIPTIONAL MEMORY TO MARK START SITE IN ACTIVE GENES!!!

"normal" histone modifications do not precisely mark the start site of transcription

MECHANSIMS THAT DECIDES HOW H2A.7 IS INCORPORATED (RI assembly) IS UNKNOWN

ROLE OF H2A.Z IN CHROMATIN AND EPIGENTIC INHERITANCE IN STEM CELLS

Bar width indicates expression level

 (b)

EMBRYONIC STEM CELLS: H2A.Z at bivalent promoters (colocalization with PcG/Trx)

Depletion of H2A.Z:

- **- More stabile association of nucleosomes** with promoters and enhance
- **Reduced activation of expression**
-

H2A.Z IS CELL CYCLE REGUALTED AT THE TSS

EXPERIMENTS: Trophoblasts in G1, S, M Phase \rightarrow gene expression array $+$ → ChIP H2A.Z and ANALYSIS OF OCCUPANCY OF GENES THAT ARE ACTIVE IN G1, S, M PHASE

H2A.Z around transcriptional start site is cell cycle regulated

Nekrasov et al 2012, Nature Structural and Molecular Biology