Histone Variants and Epigenetics

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SUMMARY

Histones package and compact DNA by assembling into nucleosome core particles. Most histones are synthesized at S phase for rapid deposition behind replication forks. In addition, the replacement of histones deposited during S phase by variants that can be deposited independently of replication provide the most fundamental level of chromatin differentiation. Alternative mechanisms for depositing different variants can potentially establish and maintain epigenetic states. Variants have also evolved crucial roles in chromosome segregation, transcriptional regulation, DNA repair, and other processes. Investigations into the evolution, structure, and metabolism of histone variants provide a foundation for understanding the participation of chromatin in important cellular processes and in epigenetic memory.

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Histones package DNA by assembling into nucleosome core particles, whereas the double helix wraps around. Over evolutionary time, histone-fold domain proteins have diversified from archaeal ancestors into the four distinct subunits that comprise the familiar octamer of the eukaryotic nucleosome. Further diversification of histones into variants results in differentiation of chromatin that can have epigenetic consequences. Investigations into the evolution, structure, and metabolism of histone variants provides a foundation for understanding the participation of chromatin in important cellular processes and in epigenetic memory.

Most histones are synthesized at S phase for rapid deposition behind replication forks to fill in gaps resulting from the distribution of preexisting histones. In addition, the replacement of canonical S-phase histones by variants, independent of replication, can potentially differentiate chromatin. The replacement of a canonical histone by a noncanonical variant is a dynamic process that changes the composition of chromatin. The differentiation of chromatin by a histone variant is especially conspicuous at centromeres, in which the H3 variant, CENP-A, is assembled into specialized nucleosomes that form the foundation for kinetochore assembly. A centromeric H3 (cenH3) counterpart of CENP-A is found in all eukaryotes. In plants and animals, the faithful assembly of cenH3-containing nucleosomes at centromeres does not appear to require centromeric DNA sequences, a spectacular example of epigenetic inheritance. Some cenH3s have evolved adaptively in regions that contact DNA, which suggests that centromeres compete with each other, and cenH3s and other centromere-specific DNA-binding proteins have adapted in response. This process could account for the large size and complexity of centromeres in plants and animals.

Chromatin can also be differentiated outside of centromeres by incorporation of a constitutively expressed form of H3, called H3.3, which is the substrate for replication-independent nucleosome assembly. Replacement with H3.3 occurs at active genes, a dynamic process with potential epigenetic consequences. Differences between H3 and H3.3 in their complement of covalent modifications might underlie changes in the properties of chromatin at actively transcribed loci.

Several H2A variants can also differentiate or regulate chromatin. H2A.X is defined as a variant by a four-amino-acid carboxy-terminal motif whose serine residue is the site for phosphorylation at sites of DNA double-stranded breaks. Phosphorylation of H2A.X is an early event in double-strand break repair, in which it is thought to concentrate components of the repair machinery. H2A.X phosphorylation also marks the inactive XY bivalent during mammalian spermatogenesis and is required for condensation, pairing, and fertility.

H2A.Z is a structurally diverged variant that has long presented an enigma. Studies in yeast have implicated H2A.Z in establishing transcriptional competence and in counteracting heterochromatic silencing. The biochemical complex that replaces H2A with H2A.Z in nucleosomes is an ATP-dependent nucleosome remodeler, providing the first example of a specific function for a member of this diverse class of chromatin-associated machines.

Two vertebrate-specific variants, macroH2A and H2A.B (also called H2A.Bbd), display contrasting features when packaged into nucleosomes in vitro, with macroH2A impeding and H2A.B facilitating transcription. These features are consistent with their localization patterns on the epigenetically inactivated mammalian X chromosome: macroH2A showing enrichment and H2A.B showing depletion.

The emerging view from these studies is that histone variants and the processes that deposit them into nucleosomes provide a primary differentiation of chromatin that might serve as the basis for epigenetic processes.
1 DNA IS PACKAGED BY ARCHITECTURAL PROTEINS IN ALL ORGANISMS

The enormous length of the DNA double helix relative to the size of the chromosome that contains it requires tight packaging, and architectural proteins have evolved for this purpose. The first level of packaging shortens the double helix and protects it from damage while still allowing DNA polymerase to gain full access to each base pair every cell cycle. In addition, these architectural proteins facilitate higher-order folding to further reduce the length of a chromosome. Perhaps because of stringent requirements for higher-order folding in histone variants are indicated in red. WHD, winged-helix domain.

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 histone-fold domain (HFD) is where histone dimerization occurs. Regions of sequence variation in histone variants are indicated in red. WHD, winged-helix domain.

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2 EUKARYOTIC CORE HISTONES EVOLVED FROM ARCHAEAL HISTONES

The eukaryotic nucleosome is a complex structure, consisting of an octamer of four core histones wrapped nearly twice by DNA, with histone tails and linker histones mediating a variety of packaging interactions outside the core particle (Arents et al. 1991; Wolffe 1992; Luger et al. 1997). Archaeal nucleosomes are much simpler, and it is evident that they resemble the ancestral particle from which eukaryotic nucleosomes evolved (Malik and Henikoff 2003). An archaean nucleosome consists of histone-fold domain proteins that lack tails and form a tetrameric particle that is wrapped only once by DNA. The genomes of all ancient archaean lineages encode histones (Fig. 2), which implies that the eukaryotic nucleosome evolved from an as-yet unidentified archaean ancestor. The kinship between archaean and eukaryotic nucleosomes can be seen by comparing their structures: The backbone of the archaean tetramer nearly superimposes over that of the (H3-H4)2 tetramer. When archaean nucleosomes are reconstituted to form chromatin, the resulting fiber behaves similarly to “tetrasomes” of (H3-H4)2, and when mapped in vivo,
they show phasing downstream from transcriptional start sites analogous to what is seen for eukaryotic nucleosomes (Ammar et al. 2012). Therefore, it is thought that eukaryotic nucleosomes evolved from an archaeal ancestor by doubling the number of subunits to allow for a second DNA wrap, and by acquisition of histone tails. In addition, DNA wraps into a right-handed superhelix around archaeal cores, but into a left-handed superhelix around eukaryotic cores.

Further insight into the origin of the eukaryotic nucleosomes comes from examination of the subunit structures of archaeal nucleosomes. Whereas most archaeal histones are undifferentiated monomers or are differentiated into structurally interchangeable variants that come together to form a tetramer, some are head-to-tail dimeric fusions that come together to form a dimer of fused dimers. When two of these fused dimers assemble into a nucleosome particle, each member of the fused pair is in a structurally distinguishable position. By occupying distinct positions in the particle, each member of the archaeal fused dimer will evolve independently, allowing it to adapt to a single position in the nucleosome particle. In contrast, monomers that occupy interchangeable positions are not free to adapt to particular positions. Indeed, the two members of archaeal dimers have diverged from one another in both independent lineages in which they are found. This process provides a possible scenario for the differentiation of an ancestral histone-fold domain protein into four distinct subunits that occupy distinct positions in the eukaryotic nucleosome. Like their presumed archaeal ancestors, eukaryotic histones form dimers, where H2A dimerizes with H2B, and H3 with H4 (which also stably tetramerizes in solution). The structural backbone of an archaeal histone dimer superimposes with those of H2A-H2B and H3-H4 at 2 Å resolution, with the first member of the dimeric repeat superimposing on H2A or H3 and the second member superimposing on H2B or H4. So, although all four eukaryotic histones lack significant sequence similarity to one another and to archaeal histones, the striking structural superposition of dimeric units suggests that eukaryotic histones evolved and differentiated from simpler archaeal ancestors.

The asymmetry of H2A-H2B and H3-H4 dimers, which appears to have originated from archaeal tandem dimers, could have led the way to subsequent diversification of eukaryotic histone variants. Both H2A and H3 correspond to the first member of archaeal tandem histone dimers and both have subsequently diversified multiple times in eukaryotic evolution. In contrast, H2B and H4 correspond to the second member and have shown little (H2B) or no (H4) functional diversification. Both H3 and H2A make homodimeric contacts in the octamer (Fig. 3), whereas H4 and H2B only contact other histones. As a result, changes in the residues involved in homodimerization of either H2A or H3 can potentially resist formation of mixed octamers, allowing nucleosomes containing an H2A or H3 variant to evolve independently of parental nucleosomes. In general, structural features that facilitated independent evolution of subunits may have been prerequisites for diversification of nucleosome particles.

Although we can rationalize the descent of the eukaryotic core histones from archaeal tandem dimers, other basic questions remain. Where did histone tails come from? Did (H3-H4)_2 tetrasomes evolve before acquiring flanking H2A-H2B dimers, or perhaps the four core histones evolved first as (H2A-H2B-H4-H3) “hemisomes” before doubling to form the octamer, or was there some other evolutionary progression from tetramer to octamer? Did these events occur before, during, or after the evolution of the eukaryotic nucleus? Did the emergence of octameric nucleosomes with two DNA wraps allow for the tight packaging of mitotic chromosomes, a eukaryotic-specific invention? Perhaps the sequences of more archaea or of primitive eukaryotes will reveal intermediate forms that can answer these questions.

![Archaeal lineages](http://cshperspectives.cshlp.org/)

**Figure 2.** Archaeal cladogram indicating the presence of histones in all ancestral clades. Losses are attributable to horizontal transfer of HU proteins from bacteria. (Modified, with permission, from Brochier-Armanet et al. 2011, © Elsevier.)
3 BULK HISTONES ARE DEPOSITED AFTER DNA REPLICATION

The packaging of essentially all DNA in a eukaryotic cell into nucleosomes requires that chromatin is duplicated when DNA replicates. Thus, canonical histones are produced during the DNA synthesis (S) phase of the cell cycle. S-phase coupling of histone synthesis to DNA synthesis is under tight cell-cycle control (Marzluff and Duronio 2002). This is especially evident in animals, in which special processing of histone transcripts by the U7 small nuclear ribonuclear protein complex, and messenger RNA (mRNA) stabilization by the stem-loop-binding protein (SLBP) contributes to the tight coordination of histone synthesis with DNA replication. The need for rapid and massive production of histones during S phase is very likely responsible for the fact that replication-coupled (RC) histones in animals are encoded in clusters that comprise many histone genes. For example, there are 14 H4 genes in the human genome, most of which are found in two major clusters, where these H4 genes are interspersed with other RC histone genes (Marzluff et al. 2002). In animals, RC histones are recognizable by the presence of a 26-bp 3′ sequence that forms a stem-loop for recognition by SLBP when transcribed into histone mRNA. Canonical plant histones are also encoded by multiple genes and are deposited during S phase, although plant histone transcripts are polyadenylated and there does not appear to be a counterpart to SLBP.

To the extent that epigenetic inheritance results from inheritance of a chromatin “state,” the process of RC nucleosome assembly has been of intense interest. The biochemistry of the process was elucidated with the development of in vitro systems that could assemble nucleosomes onto replicating DNA. These studies revealed that a three-subunit complex, CAF-1 (chromatin assembly factor 1), acts as a histone chaperone that facilitates the incorporation of H3-H4 as a first step in nucleosome assembly (reviewed in Loyola and Almouzni 2004). CAF-1 was shown to interact with the replication processivity clamp, PCNA, which implies that DNA replication and RC assembly occur in close proximity (Fig. 4). Work in budding yeast revealed that none of the subunits of complexes involved in RC assembly in vitro is essential for growth, suggesting that in vivo, there are redundant mechanisms for RC assembly. The fact that much of yeast chromatin is assembled in a replication-independent (RI) manner (Altheim and Schultz 1999) provides a rationale for this evident redundancy. As we shall see, histone variants are typically deposited by RI nucleosome assembly.

RC assembly is not completely redundant in budding yeast. An intriguing finding is that absence of the large CAF-1 subunit leads to loss of epigenetic silencing at telomeres (Loyola and Almouzni 2004). Moreover, in human cells, depletion of CAF-1 results in the deposition of H3.3 at sites of DNA replication (Ray-Gallet et al. 2011). The connection between RC assembly and epigenetic silencing has been extended to Arabidopsis, in which loss of CAF-1 subunits results in a variety of defects attributable to loss of epigenetic memory (Kaya et al. 2001). Although the mechanistic basis for these observations is unknown, it seems clear that the proper deposition of new nucleosomes behind the replication fork is important for maintaining an epigenetically silenced state. An example of the importance of replication-coupled assembly for maintaining a devel-
Opmental state is the observation that the identity of one of two sister neurons in Caenorhabditis elegans is disrupted by mutating either the H3-H3 dimerization interface or CAF-1 (Nakano et al. 2011).

A prerequisite for epigenetic inheritance of a nucleosome state is that preexisting nucleosomes must be distributed to daughter chromatids following replication. Indeed this is the case; classical studies have shown that old nucleosomes are inherited intact and evidently at random to daughter chromatids (Annunziato 2005). However, more recent studies of particular cell types and particular loci have challenged this dogma. Germline stem cells in Drosophila male early embryos show asymmetric inheritance of RC H3, but not RI H3.3, whereby old nucleosomes remain in the stem cell and new nucleosomes segregate to the differentiating daughter cell (Tran et al. 2012). Moreover, although RC H3 was not found to split at replication (Xu et al. 2010), the small amount of RI H3.3 that splits was shown to be enriched at active human genes and cell-type-specific enhancers (Huang et al. 2013). Taken together, these two findings suggest that RC nucleosome assembly is profoundly involved in initiating developmental decisions and RI assembly in epigenetic inheritance, resuscitating ideas proposed long before these pathways were elucidated (Weintraub et al. 1976). More work is needed to test these exciting possibilities and to explore the importance of nucleosome deposition pathways in establishing and maintaining epigenetic states (Jenuwein 2001; Henikoff and Ahmad 2005).

4 Variant Histones are Deposited Throughout the Cell Cycle

As we have seen, core histones can be classified based on their ancestral sequence and position in the nucleosome. Linker histones are characterized by a winged-helix domain, rather than a histone-fold domain, and bind to the linker DNA that separates nucleosomes (Wolfe 1992). Although minor variants of these canonical histones exist, they appear to be interchangeable with the major form. For example, mammalian H3.1 and H3.2 differ by a single amino acid that is not known to impart different functional properties to the two isoforms. The existence of multiple genes that produce large amounts of canonical histones for S-phase deposition is typical of eukaryotic genomes. The near ubiquity and overwhelming abundance of canonical S-phase histones has resulted in relatively little attention being paid to histone variants until recently.

The renaissance of interest in histone variants came in part from the realization that they differ from canonical S-phase histones in ways that can lead to profound differentiation of chromatin. One way that they differ is in their mode of incorporation into chromatin. RC assembly incorporates new nucleosomes into gaps between old nucleosomes genome-wide, whereas RI assembly involves local replacement of an existing nucleosome or subunit (Marzluff et al. 2002). RI assembly therefore has the potential of switching a chromatin state by replacing a canonical histone with a variant. Replacing one histone with another also could erase or alter the pattern of posttranslational modifications. Therefore, RI assembly can potentially reset epigenetic states that are thought to be mediated by histones and their modifications. Recent progress in studying histone variants and the processes by which they are deposited has led to new insights into the basis for epigenetic inheritance and remodeling. Below we discuss features of particular histone variants that contribute to chromatin differentiation and might be involved in propagating epigenetic information.

5 Centromeres are Identified by a Special H3 Variant

A defining feature of the eukaryotic chromosome is the centromere, which is the site of attachment of spindle microtubules at mitosis. The first centromeres to be described in molecular detail were those of budding yeast (S. Henikoff and M.M. Smith

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).
myces cerevisiae), in which a 125-bp sequence is necessary and sufficient for centromere formation (Amor et al. 2004b). However, centromeres of plants and animals are very different, typically consisting of megabase arrays of short tandem repeats. Unlike the situation for budding yeast, the role of DNA sequence at these complex centromeres is uncertain because fully functional human neocentromeres are known to form spontaneously at ectopic sites that entirely lack sequences resembling centromeric repeats (Fig. 5A). These and other observations argue against a direct role of DNA sequence in determining the location of centromeres (see Allshire and Ekwall 2014).

A key insight into the basis for centromere identity and inheritance came from the identification of a histone H3 variant, CENP-A which was found to localize specifically to centromeres and be incorporated into nucleosomal particles in place of H3 itself (Palmer et al. 1991). Remarkably, CENP-A remains associated with centromeres during the transition from histones to protamines during spermatogenesis, when essentially all other histones are lost (Palmer et al. 1990). This early observation in the study of CENP-A suggested that CENP-A contributes to centromere identity of the male genome. The generality of this insight was not fully appreciated until it was realized that CENP-A is a much better marker for centromeres than DNA sequence (Fig. 5) (Amor et al. 2004b), and that counterparts of CENP-A (cenH3s) can be found in the genomes of all eukaryotes (Talbert and Henikoff 2010). So, although budding yeast centromeres are determined by a 125-bp consensus sequence, this is also the site of a centromeric nucleosome that contains the cenH3 variant. In fission yeast (Schizosaccharomyces pombe), an array of cenH3-containing nucleosomes occupies the central core region of the centromere flanked by H3-containing nucleosomes that display heterochromatic features (see Fig. 3 of Allshire and Ekwall 2014; Amor et al. 2004b). In flies and vertebrates, cenH3s are present in arrays that alternate with H3-containing arrays that display a unique pattern of histone modifications (Sullivan and Karpen 2004). Alternation can account for the fact that centromeres occupy only the outside edge of the centromeric constriction of metaphase chromosomes. This is consistent with the observation that in worm “holokinetic” chromosomes, microtubules attach throughout the length of each anaphase chromosome, and cenH3 occupies the leading edge all along its length (Fig. 5C) (Malik and Henikoff 2003). Indeed a unique cenH3 variant is found to precisely mark the centromere in nearly all eukaryotes (Fig. 6A). This apparent ubiquity, and the presence of centromeres to perform mitosis in all eukaryotes, raises the possibility that the first canonical H3 evolved from a cenH3.

Genetic experiments in a variety of eukaryotes have confirmed the essentiality of cenH3 for formation of the kinetochore and for chromosome segregation (Amor et al. 2004b). Because they remain in place throughout the cell cycle, cenH3-containing nucleosomes form the foundation for assembly of other kinetochore proteins during mitosis and meiosis (see Allshire and Ekwall 2014). An outstanding question in chromosome research is just how these proteins interact to provide a linkage between the centromere and spindle microtubules that can hold up to the strong pulling forces exerted on kinetochores at anaphase. Several dozen kinetochore-specific proteins have been identified in yeast (for more detail, see Allshire and Ekwall 2014), although exactly how they interact with cenH3-containing nucleo-

Figure 5. cenH3s at centromeres of eukaryotes. (A) Human neocentromeres (indicated by an arrow) lack centromeric α-satellite DNA, but have CENP-A and heterochromatin. Anti-CENP-A staining in green and Anti-CENP-B staining in red (which marks α-satellite DNA) identifies a Chromosome 4 neocentromere that lacks α-satellite (main panel). This Chromosome 4 is otherwise normal, having been transmitted for at least three meiotic generations in normal individuals. Inset shows anti-HP1 staining, which indicates that despite the lack of satellite DNA, heterochromatin forms around active neocentromeres (indicated by arrow). (Reprinted, with permission, from Amor et al. 2004a, © National Academy of Sciences.) (B) Drosophila melanogaster anti-cenH3 antibody (red) stains centromeres in metaphase chromosomes and throughout interphase. (Image courtesy of Suso Platero.) (C) C. elegans anti-cenH3 antibody (green) stains the end-to-end holocentromeres of prophase chromosomes (red). (Image courtesy of Landon Moore.)
Figure 6. Histone variant phylogenies. Histone sequences from selected species were multiply aligned and neighbor-joining trees were produced using the EBI server (http://www.ebi.ac.uk/Tools/phylogeny). (A) Histone H3s. (B) Histone H2As. Note that there are no clear phylogenetic distinctions between RC H3 and RI H3,3, and between RC H2A and RI H2A.X. (C) H1 variants from diverse eukaryotes show a “star” phylogeny, which suggests that they are functionally interchangeable. (C, Modified from Talbert et al. 2012.)
somes and other foundation proteins, such as CENP-C, remains unclear.

The evolution of cenH3s is unlike that of any other histone class. Whereas histone H3 is almost invariant in sequence, which reflects extraordinarily strong purifying selection on every residue, cenH3s are evolving rapidly, especially in plant and animal lineages (Talbert and Henikoff 2010). This is most evident from the amino-terminal tails, which differ in length and sequence to such an extent that they cannot be aligned between the cenH3s of different taxonomic groups. Even the histone-fold domain of cenH3 is evolving orders of magnitude faster than that of H3.

What is the reason for this striking evolutionary difference between an H3 that functions at centromeres and an H3 that functions everywhere else?

Rapidly evolving regions of Drosophila and Arabidopsis cenH3 genes display an excess of replacement nucleotide substitutions over what would be expected from the rate of synonymous substitutions (Malik and Henikoff 2009). This excess is a hallmark of adaptive evolution. Adaptive evolution in plants and animals is also seen for another major centromere foundation protein, CENP-C (Malik and Henikoff 2009). Although adaptive evolution is well documented for genes involved in genetic conflicts such as arms races between hosts and parasites, these are the only known essential single-copy genes that are adaptively evolving in any organism. In the case of cenH3 and CENP-C, the regions of adaptive evolution correspond to regions of DNA binding and targeting. This suggests that the major centromere-binding proteins are adapting to the evolving centromeric DNA, thus allowing centromeric chromatin to interact with the conserved kinetochore machinery that connects the centromere to spindle microtubules. It has been proposed that centromeres compete during female meiosis to be included in the egg nucleus rather than being lost as polar bodies (Malik and Henikoff 2009). An arms race would develop leading to expansion of centromeres, probably by unequal crossing-over between sister chromatids. Host suppression of this meiotic drive process by cenH3 and CENP-C would lead to an excess of replacement changes in regions that interact with DNA. Organisms in which there is no opportunity for centromeres to compete, such as budding yeast, would not undergo centromere drive, and this might account for the fact that they have small centromeres, and their cenH3 and CENP-C proteins are under strong purifying selection.

Thus, we see that a special region of the genome, the centromere, is distinguished by a single histone variant class, whose sequences reveal remnants of an arms race that may have led to the extraordinary complexity of centromeres. The RI assembly process that targets new cenH3-containing nucleosomes to centromeres every cell cycle has been elucidated by the discovery and detailed characterization of the related Scm3 (yeast) and HJURP (mammalian) cenH3-specific chaperones (Stoler et al. 2007; Dunleavy et al. 2009; Foltz et al. 2009). Detailed biochemical and structural characterization of Scm3/HJURP complexes (Shuaib et al. 2010; Cho and Harrison 2011; Hu et al. 2011) indicates a role in CenH3 nucleosome assembly that parallels that of other nucleosome assembly complexes, discussed further in Almouzni and Cedar (2014). Centromeric nucleosomes show a remarkable lack of sequence specificity in that they not only can faithfully localize to neocentromeres that are completely unlike native centromeres (Fig. 5A), but also the yeast homolog Cse4 can functionally replace human CENP-A (Wieland et al. 2004). It is extraordinary that our centromeres have remained in the same positions for tens of millions of years without any evident sequence determinants involved in the process that maintains them. To the extent that epigenetics refers to inheritance that does not depend on DNA sequence, the inheritance of centromeres on a geological timescale is the most extreme form imaginable. Yet, we are still seeking a mechanism to explain how they have maintained themselves for even a single cell cycle.

There is now general agreement in the centromere field that the cenH3 nucleosome is the key to understanding the epigenetic inheritance of centromeres (Black and Cleveland 2011; Henikoff and Furuyama 2012). It is not only necessary for recruitment of the other structural components of the centromere, in some experimental systems it is also sufficient (Guse et al. 2011; Mendiburo et al. 2011). However, its molecular structure has been the subject of controversy for several years. In vivo evidence from flies, humans, and yeast is most consistent with a right-handed hemisome (Henikoff and Furuyama 2012), whereas several groups have shown that reconstitution of cenH3-containing particles generally results in the formation of a partially unwrapped left-handed octameric nucleosome (Black and Cleveland 2011). Indeed, since 2007, the authors of this article coauthored the studies that provided the first evidence for nonoctameric cenH3 particles (Dalal et al. 2007; Mizuguchi et al. 2007), but the composition and structure of these proposed particles were completely different! In light of the continuing controversy, we leave a final resolution of this important issue to the future.

6 THE REPLACEMENT HISTONE VARIANT H3.3 IS FOUND AT ACTIVE CHROMATIN

Like centromeres, transcriptionally active chromatin is thought to be maintained epigenetically and is enriched in an H3 variant, H3.3, which is the substrate for RI dep-
osition (Filipescu et al. 2013). H3.3 is very similar in sequence to the canonical forms of H3, differing by only four amino acids. With so few differences, it might have been assumed that these two forms are interchangeable. However, H3.3 is deposited exclusively by RI nucleosome assembly, whereas H3 is deposited only at replication foci in an RC manner. This difference between the two variants is encoded in the protein itself, with three of the four differences between H3 and H3.3 evidently involved in preventing H3 from being deposited by an RI pathway (illustrated in the α-helix 2 of Fig. 3). Purification of soluble nucleosome assembly complexes confirmed that these two forms participate in distinct assembly processes: H3.1 copurifies with CAF-1 for RC assembly, and H3.3 copurifies with other components, including the HirA and Daxx histone chaperones, and participates in RI assembly.

Although the four-amino-acid difference might seem practically insignificant, when one considers that humans, flies, and clams have precisely the same H3.3 sequence, these differences from H3 stand out. Phylogenetic analysis reveals that the H3/H3.3 pair evolved at least four separate times during eukaryotic evolution, in plants, animals, fungi, ciliates, and apicomplexans (Fig. 6A) (Talbert and Henikoff 2010). Despite having a separate origin from animals and fungi, the animal H3/H3.3 pair and the pair from plants (called H3.1 [RC] and H3.2 [RI]—to avoid confusion, we will refer to all RC isoforms as H3 and all RI isoforms as H3.3) are strikingly similar. The same cluster of amino acids (positions 87–90) that prevents RI deposition of H3 in Drosophila are found to differ in plants, and the remaining difference in animals (position 31 is Ala for H3 and either Ser or Thr for H3.3) is also found in plants. Fungi are especially interesting. Ancestrally, they have both H3 and H3.3; however, ascomycetes, which include yeasts and molds, have lost the H3 form. Thus, the obligate RC form of histone 3 that has received the most attention in animals is not even present in yeast.

Studies of H3.3 in bulk chromatin showed that it is enriched in transcriptionally active chromatin fractions (Filipescu et al. 2013). However, various factors contributed to the obscurity of this potential “mark” of active chromatin during a time of great excitement in the chromatin field when it was realized that histone modifications can distinguish active from silent chromatin. For one thing, no antibodies were available that could effectively distinguish H3 from H3.3 in chromatin (positions 87–90 are blocked by the DNA gyres in the nucleosome), whereas excellent antibodies against many different posttranslational modifications were readily available. Also, the seemingly slight sequence differences between H3 and H3.3 did not suggest any fundamental distinctions in chromatin, whereas histone modifications were mostly on tail lysines that were known to affect chromatin interactions or to bind chromatin-associated proteins. This perception that the two histone 3 forms should be interchangeable was confirmed by the finding in Tetrahymena and Drosophila that the S-phase form can in general substitute for its replacement counterpart. Finally, the influential “histone code” hypothesis envisioned nucleosomes as fixed targets of modification enzymes during chromatin differentiation (Jenuwein and Allis 2001). However, it has become increasingly evident that chromatin is highly dynamic, and even heterochromatin-associated proteins bind with residence times of a minute or less (Phair et al. 2004). It appears that the chromatin of actively transcribed genes is in constant flux, characterized by continual histone replacement (Dion et al. 2007). The three core amino acid differences that distinguish H3 and H3.3 make H3.3-H4 dimers the substrate for RI assembly, and RI assembly itself profoundly changes chromatin. As a result of this process, actively transcribed regions become marked by H3.3 (Fig. 7), and evidence for this process comes from the observation of RI replacement of H3 methylated on lysine-9 (H3K9me) with tagged H3.3 at RNA polymerase I and II (RNA Pol I and II) transcribed loci (Schwartz and Ahmad 2005).

The dynamic nature of chromatin at active loci results in the erasure of preexisting histone modifications, and yet histone modification states persist through multiple rounds of cell division. Therefore, the enzymes responsible for modifying histones must be targeted to their sites of action. For histone modifications that are typically associated with transcriptionally active chromatin, this is achieved by association with the carboxy-terminal domain (CTD) of RNA Pol II, a tandem array of YSPTSPS

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.
This implies that all of the components necessary for main-

enzyme that performs the modification, the modifica-
tion-specific-binding module on the machine that uses 

ATP to provide energy for the replacement process, and 

the fresh unmodified H3.3 substrate that becomes incor-

porated into the new nucleosome (Fig. 8). It seems likely 

that a similar process occurs at other sites of short-period 
tandem repeats in DNA where nucleosomes frequently turn 

over, insofar as ATRX is also abundant in mammals at C+ 

G-rich sites (Law et al. 2010), which are found at most 

promoters, and the Drosophila ortholog of ATRX, XNP, is 

abundantly present at a single site of a (GATA)ₙ repeat 

where H3.3 is actively incorporated (Schneiderman et al. 

2009). The incorporation of H3.3 nucleosomes at sites of 
telomeric heterochromatin belies the common notion that 

H3.3 is a “mark” of active chromatin. As the general sub-

strate for replacement of nucleosomes wherever they are 

lost, the finding that H3.3 is mostly incorporated at active 
genes rather implies that these are sites where nucleosome 
turnover is most intense. This generic replacement func-

tion of H3.3 has important relevance to human disease as 
described in Section 15.

7 H3.3 FUNCTIONS IN THE GERMLINE

When cells exit the cell cycle and differentiate, they no 

longer produce or incorporate S-phase histones, and 

H3.3 accumulates as a result. For example, H3.3 accumu-

lates in rat brains to a level of 87% of the histone 3 by the 
time that rats are 400 days old (Pina and Suau 1987). This 
classical observation suggested that replacement by H3.3 

has no functional significance except to prevent holes in the
nucleosomal landscape. Consistent with this view, H3.3 has been found to be nonessential for *Drosophila* development, as flies lacking both H3.3 genes develop normally to the pupal stage, with occasional adult escapers that die shortly after eclosion but show no specific morphological defects (Hodl and Basler 2009; Sakai et al. 2009). Moreover, H3.3 can functionally substitute for H3 and allow at least some developmental decisions to be made in *Drosophila* embryos, further suggesting that RC and RI substrates are largely interchangeable (Hodl and Basler 2012). It appears that histone replacement in H3.3 null proliferating cells can be accomplished by incorporating H3 nucleosomes using the RI pathway. However, there is no germline development in flies lacking key RI pathway components, such as HirA (e.g., Fig. 9). Females that lack the ChD1 ATP-dependent nucleosome remodeler protein are also sterile, evidently because ChD1 is required in the zygote for sperm nuclei decondensation and the replacement of protamines by maternally encoded RI histones (Orsi et al. 2009). This essential germline function of the RI pathway is conserved in mammals, in which H3.3 is required for remodeling of both maternal and paternal gametes (Santenard et al. 2010; Akiyama et al. 2011). Similar RI processes have been documented in both *C. elegans* and *Arabidopsis*, in which maternal H3.3 is incorporated into the paternal genome of the zygote (Ooi et al. 2006; Ingouff et al. 2007). Therefore, germline remodeling via RI assembly of H3.3 nucleosomes is a universal process that has evolved in both animals and plants, most likely to “reset” the chromatin landscape to a totipotent state.

Remodeling events in the zygote, where RI assembly of H3.3 nucleosomes plays a key role, is analogous to nuclear reprogramming, which can be artificially induced in *Xenopus* eggs, mouse embryonic stem cells, and induced pluripotent cells. In *Xenopus*, transfer of an embryonic nucleus into an enucleated egg can result in the production of mostly normal embryos that nevertheless sometimes (mis)express genes that were active in the differentiated donor nucleus (Ng and Gurdon 2008). The lack of observable gene expression during the intervening 12 embryonic

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**Figure 9.** Models for RI replacement or exchange. A large molecular machine (either the SWR1 complex or RNA polymerase) partially or completely unravels a nucleosome during transit. The result is either retention of heterodimeric subunits, such as the FACT-facilitated transfer of H2A-H2B from in front of RNA polymerase to behind (Formosa et al. 2002; Belotserkovskaya et al. 2003) or loss of a heterodimer. In the latter case, chromatin repair replaces the lost heterodimer with either H3.3-H4 (left) or H2A.Z-H2B (right).
divisions implies that the persistence of an epigenetic mark maintains the memory of prior gene activity. Overexpression of H3.3 in developing embryos improved epigenetic memory and mutation of H3.3K4 to glutamine abolished memory of the active state, whereas no effect was seen using a general DNA methyltransferase inhibitor. Further evidence for the importance of H3.3 in nuclear reprogramming comes from the finding that the shift from somatic to oocyte transcription does not require replication, but does require transcription and the H3.3-specific chaperone, HirA (Jullien et al. 2012). It is attractive to think of H3.3 and its histone partners as general mediators of totipotency both in the zygote and during nuclear reprogramming, insofar as histones are likely to be most accessible to the enzymes that posttranslationally modify their tails during nucleosome assembly (e.g., Fig. 8).

8 PHOSPHORYLATION OF H2A.X FUNCTIONS IN DNA DOUBLE-STRAND BREAK REPAIR

The H2A histones also comprise a family of distinct variants found throughout eukaryotes (Fig. 6B). The H2A.X variant is defined by the presence of a carboxy-terminal amino acid sequence motif, SQ(ε or D)Ø, in which Ø indicates a hydrophobic amino acid. The serine in this sequence motif is the site of phosphorylation, producing a modified protein designated “γH2A.X.” The dynamic nature of chromatin, and H2A.X phosphorylation, is especially evident when double-strand breaks (DSBs) occur in DNA (Morrison and Shen 2005). The lethality of even a single double-stranded (ds) break requires immediate action to repair the lesion and restore the continuity of the double helix. The detection of a ds break normally occurs within a minute or so of its formation and this, in turn, triggers the rapid phosphorylation of H2A.X in the immediate vicinity of a break site. This phosphorylation is performed by members of the phosphoinositol 3-kinase-like kinase family. Following this initial event, H2A.X phosphorylation then spreads quickly along the chromosome marking a relatively large chromatin domain surrounding the break. Finally, the ds break is eventually repaired by either homologous recombination or nonhomologous end-joining and the phosphorylation mark is removed.

Phosphorylation of H2A.X is not essential for detection or repair of DSBs because deletion of the gene or mutation of the target serine residue does not abolish repair. However, H2A.X is not just a marker of damage because such mutants have reduced efficiency of repair and are hypersensitive to radiation damage and genotoxic agents. Currently, H2A.X is thought to function in ds break repair in at least two ways. First, it may help recruit or retain proteins required for repair at the site of the break (Morrison and Shen 2005). Second, it may stabilize the chromosome surrounding the broken ends through the recruitment of cohesin, the protein complex responsible for keeping sister chromatids together (Lowndes and Toh 2005).

The evolution of H2A.X is unlike that of other histone variants. Although a gene for H2A.X is found in nearly all eukaryotes, it has had multiple relatively recent divergences from H2A (Fig. 6B) (Malik and Henikoff 2003; Talbert and Henikoff 2010). For example, the version of H2A.X found in Drosophila is different from that found in another dipteran insect, Anopheles. Some organisms, such as yeasts, have an H2A.X but lack an H2A, consistent with the possibility that most present-day canonical H2As have evolved from ancestral H2A.Xs (Talbert and Henikoff 2010). Presumably, the ability to evolve either a canonical H2A from an H2A.X or vice versa is a consequence of the simplicity of the SQE motif. Depending on selective constraints, the loss or gain of such a simple motif at the carboxyl terminus of a protein might occur repeatedly over evolutionary time. Occasional loss of an existing H2A.X with a newly minted version might be fueled by the need for H2A.X to be very uniformly distributed because DSBs can occur anywhere in the genome. If mutations occur in an existing H2A.X gene that reduce its similarity to the canonical H2A in such a way that its assembly becomes less efficient or uniform, then there will be strong selection to replace it with a version that is more similar to canonical H2A. This rationale could help account for the exceptional case of Drosophila H2A.X, which unlike other eukaryotes, is not derived from its canonical H2A, but rather from the distant H2A.Z variant lineage (Baldi and Becker 2013). If all that is necessary to be an H2A.X is to be in the H2A position in a nucleosome and to have the carboxy-terminal motif for phosphorylation, then an H2A.Z can evolve this capability.

DSB repair is clearly the universal function of H2A.X phosphorylation, and there would seem to be no stable epigenetic aspect to this process. However, H2A.X null mice are sterile, and cytotogical examination of mammalian spermatogenesis has revealed a striking epigenetic feature in which H2A.X is specifically phosphorylated on the XY bivalent (Fig. 10) (Fernandez-Capetillo et al. 2003). This chromosome pair occupies a distinct “sex body” during meiotic prophase, which has been implicated in silencing of sex-linked genes during male meiosis. H2A.X phosphorylation is essential for normal sex-body formation, and H2A.X-deficient spermatocytes fail to pair or condense and fail to inactivate X and Y genes during meiosis. H2A.X phosphorylation of the XY bivalent is distinct from the process that occurs at DSBs. XY phosphorylation comes from the finding that the shift from somatic to oocyte transcription does not require replication, but does require transcription and the H3.3-specific chaperone, HirA (Jullien et al. 2012). It is attractive to think of H3.3 and its histone partners as general mediators of totipotency both in the zygote and during nuclear reprogramming, insofar as histones are likely to be most accessible to the enzymes that posttranslationally modify their tails during nucleosome assembly (e.g., Fig. 8).
most conspicuously at unpaired regions of the chromosomes. The mechanisms whereby H2A.X phosphorylation is targeted to unpaired chromosomes and how this event leads to condensation, pairing, and silencing are currently unknown. However, it is interesting to speculate that this role may be related to its ability to interact with and recruit cohesin.

9 H2A.Z PLAYS DIVERSE ROLES IN CHROMATIN REGULATION

Histone variant H2A.Z is found throughout most eukaryotic lineages and it has been intensely studied for its structure and function in chromatin biology (Zlatanova and Thakar 2008; Draker and Cheung 2009; Marques et al. 2010; Talbert and Henikoff 2010). H2A.Z diverged from an ancestral H2A early in evolution and shares only ~60% similarity with its major histone H2A counterpart. Consistent with this separate lineage, genetic experiments in yeast, plants, flies, and mammals have shown that histones H2A and H2A.Z have evolved to play separate nonoverlapping functions. H2A.Z is an essential histone in most organisms, from ciliated protozoans to mammals. However, in budding and fission yeasts, cells that carry a deletion of the single-copy H2A.Z gene are viable, although the null mutants show a variety of conditional lethal phenotypes. Plants, as exemplified by Arabidopsis thaliana, have three closely related H2A.Z genes, HTA8, HTA9, and HTA11, with roughly 90% identity and a more distantly related HTA4 gene (Talbert and Henikoff 2010). There appears to be functional redundancy among these genes as organisms with single deletions of either HTA8, HTA9, or HTA11 are normal, whereas the hta9 hta11 double mutant shows developmental defects. Interestingly, vertebrates have evolved two closely related H2A.Z variants, H2A.Z.1 and H2A.Z.2, which differ at only three amino-acid residues (Dryhurst et al. 2009; Matsuda et al. 2010; Mehta et al. 2010). These two variants show different patterns of chromatin distribution (Dryhurst et al. 2009) and are apparently nonredundant as the single deletion of H2A.Z.1 in the mouse is lethal (Faast et al. 2001).

Evidence for functional diversification of H2A.Z comes from the discovery of an alternatively spliced form of human H2A.Z.2 mRNA that results in a protein with a reduced CTD. This shorter H2A.Z isoform destabilizes nucleosomes and is most enriched in brain (Bonisch et al. 2012), in which H2A.Z, like H3.3, is known to be especially abundant (Pina and Suau 1987). Although this particular isoform appears to be limited to primates, evidence for a longer alternatively spliced isoform with potential for nucleosome destabilization was reported in carp brain (Simonet et al. 2013). With the increasing popularity of RNA-seq for identification of alternatively processed variants, we expect that other examples of potential H2A.Z functional diversification will be discovered.

The high-resolution structure of an H2A.Z-containing nucleosome reveals several unique properties of the variant (Suto et al. 2000). Compared with H2A nucleosomes, H2A.Z presents an extended acidic patch domain on the surface of the nucleosome and mutational studies have shown this to have functional significance. The acidic patch is then part of a larger “docking domain,” an essential part of the protein necessary for interaction with H3 in the nucleosome. Like other histones, H2A.Z is subject to a variety of posttranslational modifications, including acetylation, ubiquitylation, and sumoylation. There is strong genetic and biochemical evidence that these modifications affect the localization, dynamics, and function of H2A.Z nucleosomes (Talbert and Henikoff 2010).

H2A.Z has been linked to a wide variety of different and sometimes contradictory nuclear functions, including transcriptional activation, transcriptional repression, RNA Pol II elongation, heterochromatin, antisilencing, cell-cycle control, DNA replication, DNA damage repair, chromosome segregation, and genome integrity (Zlatanova and Thakar 2008; Altaf et al. 2009; Marques et al. 2010; Talbert and Henikoff 2010; Xu et al. 2012; Adkins et al.
H2A.Z has been mapped genome-wide in a variety of eukaryotes. In budding yeast, nematodes, and plants, H2A.Z occupancy around the promoter is correlated with nontranscribing genes “poised” for activation (Zhang et al. 2005; Mavrich et al. 2008; Whittle et al. 2008; Kumar and Wigge 2010). However, in flies and mammals, promoter H2A.Z occupancy appears to correlate more with actively transcribing genes (Barks et al. 2007; Mavrich et al. 2008; Hardy et al. 2009; Hardy and Robert 2010; Kelly et al. 2010), similar to the situation for H3.3. Although preferentially found at promoters and regulatory sites, H2A.Z nucleosomes can also be found at lower frequency in gene bodies and elsewhere (Hardy et al. 2009; Weber et al. 2010; Santistebe et al. 2011). Enrichment of H2A.Z over gene bodies closely corresponds to that of chromatin that is extracted with low salt, suggesting that H2A.Z changes the physical properties of nucleosomes (Weber et al. 2010).

H2A.Z is also specifically deposited near or within heterochromatin. In budding yeast, H2A.Z is enriched near telomeres where it serves as an antisilencing factor. Deletion of the H2A.Z gene results in extended spreading of silent chromatin inward from the telomeres and this defect can be suppressed by the additional deletion of genes encoding the silencing factors themselves (see Grunstein and Gasser 2013 for more detail). Indeed, this function may act globally, in parallel with the Set1 histone H3 methyltransferase, to prevent large-scale aberrant distribution of silencing factors (Venkatasubrahmanyam et al. 2007). In metazoans, H2A.Z is also localized in facultative and constitutive heterochromatin, the inactive X chromosome, transposable elements, and pericentric heterochromatin (Greaves et al. 2007; Draker and Cheung 2009; Boyarchuk et al. 2011; Zhang and Pugh 2011).

In contrast to the chromosomal features that are correlated with H2A.Z, there is a remarkable anticorrelation between histone H2A.Z nucleosome occupancy and DNA methylation (Zilberman et al. 2008; Kobor and Lorincz 2009; March-Diaz and Reyes 2009; Conerly et al. 2010; Edwards et al. 2010; Zemach et al. 2010; see Li and Zhang 2014 for a discussion of DNA methylation). There is strong evidence that this mutual antagonism is causal and not simply correlational. Mutants in A. thaliana with decreased DNA methylation show an increase in H2A.Z occupancy at loci where it is normally not found, independent of transcriptional activity. Conversely, mutants defective in H2A.Z deposition show increased DNA methylation over gene bodies normally occupied by H2A.Z nucleosomes. While the precise molecular pathways that account for this mutual exclusion remain to be elucidated, this functional relationship has important implications for development and carcinogenesis. For example, the stochastic
or environmentally effected loss of H2A.Z from the promoter of a tumor-suppressor gene could well contribute to locally increased DNA methylation and heritable epigenetic repression.

11 H2A.Z NUCLEOSOME OCCUPANCY IS DYNAMIC AND CHANGES THE PROPERTIES OF CHROMATIN

The dynamic exchange of H2A.Z nucleosomes in chromatin appears to be an important part of its function. (See Becker and Workman 2013 for an in-depth discussion of histone exchange.) Unlike the major core histones, H2A.Z expression is not restricted to S phase and it can be incorporated into chromatin independent of DNA replication. The deposition of H2A.Z into nucleosomes is performed by multisubunit protein complexes, which have been conserved throughout the eukaryotic kingdom (Lu et al. 2009; March-Diaz and Reyes 2009; Morrison and Shen 2009).

First identified in budding yeast, the SWR1 complexes contain, as their catalytic subunits, homologs of the protein S. pombe SNF family of ATP-dependent chromatin remodelers. The substrate for SWR1 is an H2A.Z-H2B dimer, which is used to replace one of the existing H2A-H2B dimers in the nucleosome in an ATP-dependent exchange reaction (Fig. 9). This reaction is step-wise and unidirectional, in vitro, resulting in the complete replacement of H2A-H2B dimers with H2A.Z-H2B dimers (Luk et al. 2010). In vivo, unidirectional replacement of H2A with H2A.Z by SWR1 is enforced by acetylation of H3K56, which allows the reverse reaction to occur, resulting in local reduction in H2A.Z incorporation, thus modulating transcription (Watanabe et al. 2013). SWR1 is likely dedicated to the task of replacing H2A with H2A.Z because the effects of eliminating SWR1 function are similar to the effects of deleting the gene encoding H2A.Z itself. Indeed, the activity of the SWR1 complex is actually deleterious to the cell in the absence of its H2A.Z-H2B substrate (Halley et al. 2010; Morillo-Huesca et al. 2010).

The removal of H2A.Z from chromatin proceeds by at least two pathways. Nucleosome exchange and eviction occurs in many contexts such as the remodeling and eviction of complete nucleosomes that can happen at promoters. Any H2A.Z that is a part of those nucleosomes will be removed as well. However, there is also evidence that H2A.Z-H2B dimers may be specifically removed from nucleosomes by the INO80 complex, a close relative of SWR1 (Morrison and Shen 2009; Papamichos-Chronakis et al. 2011). In budding yeast, the elimination of INO80 results in the global mislocalization of H2A.Z and a decrease in its apparent exchange rate. Additional mutational results are consistent with this interpretation. In vitro, purified INO80 is reported to catalyze the replacement of a nucleosomal H2A.Z-H2B dimer with a canonical H2A-H2B dimer, that is, the reverse of the SWR1 reaction (Luk et al. 2010; Papamichos-Chronakis et al. 2011), and this reaction may also be regulated by acetylation of H3K56 (Watanabe et al. 2013).

The factors that determine where H2A.Z is deposited are incompletely understood. At present, there is little evidence that H2A.Z templates its own deposition epigenetically (Viens et al. 2006). In budding yeast, a DNA sequence related to the binding site of transcription factor Reb1 is able to target the enrichment of H2A.Z at ectopic sites, independent of Reb1 (Raisner et al. 2005). In other cases, transcription factors themselves are implicated in targeting H2A.Z deposition, although how this recruits H2A.Z is not currently known (Hartley and Madhani 2009). Interestingly, SWR1 complexes often contain homologs of yeast Brd1, a protein containing dual bromodomain motifs capable of binding acetylated lysines. Thus, SWR1 may be recruited to, or stabilized at, chromatin neighborhoods rich in acetylated histones. H2A.Z deposition may also be blocked at specific loci. The S. pombe SWR1 complex contains a regulatory subunit, Msc1, which is dispensable for the loading of H2A.Z at promoter nucleosomes, but which is required to prevent H2A.Z deposition within the chromatin of the inner centromere and subtelomeric regions (Buchanan et al. 2009; Zofall et al. 2009). It has also been suggested that an additional pathway directing localized occupancy by H2A.Z may involve its random deposition and then specific eviction, perhaps as a consequence of transcription (Hardy and Robert 2010).

In any case, the consequence of H2A.Z deposition and replacement can be complex. Because of the exchange reactions, nucleosomes in cellular chromatin may be “ZZ,” “ZA,” or “AA,” containing two, one, or zero H2A.Z-H2B dimers, respectively (Luk et al. 2010; Weber et al. 2010). In Drosophila, the pattern of ZZ and ZA nucleosome occupancy is different, and homotypic H2A.Z nucleosomes are enriched over the bodies of active genes, perhaps as a consequence of transcription elongation. In mouse trophoblast cells, there are distinct changes in the promoter content of ZZ and ZA nucleosomes during G1, S, and M phase, independent of actual transcription activity, suggesting a major remodeling pathway dependent on the cell-division cycle (Nekrasov et al. 2012). Differential post-translational modifications further enrich the situation as H2A.Z acetylation has been linked with both SWR1 and INO80 functions (Millar et al. 2006; Papamichos-Chronakis et al. 2011).
A quantitative evaluation of the +1 nucleosome in yeast revealed that at steady state the relative abundance of ZZ, ZA, and AA nucleosomes is roughly 32%, 24%, and 44% respectively (Luk et al. 2010). How can H2A.Z drive chromatin function with this level of heterogeneity? One explanation is that H2A.Z nucleosomes are in dynamic exchange at these sites. In budding yeast, nucleosomes with rapid turnover were identified by the kinetic incorporation of newly synthesized histone H3. These “hot” nucleosomes preferentially map to promoter regions, including the nucleosomes at the TSS, which are enriched in H2A.Z histones (Dion et al. 2007). Remarkably, in budding yeast H2A.Z appears to increase the global turnover of nucleosomes and not simply those into which it is most abundantly incorporated (Dion et al. 2007; Santisteban et al. 2011). The mechanistic basis for this influence is currently unknown.

The stability of H2A.Z-containing nucleosomes in vitro has been examined in many studies with contrasting results (Zlatanova and Thakar 2008; Talbert and Henikoff 2010). In vivo it is clear that not all H2A.Z nucleosomes are created equal and that the lability of H2A.Z nucleosomes is affected by posttranslational modifications and the presence of other variant histones. Nucleosomes containing both H2A.Z and H3.3, especially H2A.Z/H2A nucleosomes with H3.3, are particularly unstable and sites occupied by these double-variant nucleosomes may be erroneously scored as entirely free of nucleosomes, depending on how the chromatin is isolated (Jin et al. 2009; Nekrasov et al. 2012). H2A.Z acetylation generally destabilizes nucleosomes in vitro and in vivo is associated with gene activation (Tanabe et al. 2008; Wan et al. 2009; Halley et al. 2010).

A particularly striking example of the differential properties of H2A.Z nucleosomes is the transcriptional response of cells to temperature. Plants possess a signaling pathway that senses ambient temperature and regulates gene expression (see Baulcombe and Dean 2014 for a discussion of plant responses to environmental factors). The developmental program for flowering, for example, is accelerated at higher temperature. To identify factors that regulate this response, *A. thaliana* was screened for mutants with a constitutive high temperature expression pattern. The pathways regulating these cell-cycle behaviors of H2A.Z are currently unknown.

The location of a gene within the nucleus can be important for its expression. Positioning at the nuclear periphery is often associated with gene inactivation with the exception of localization at nuclear pores, which is a positive factor in expression (reviewed in Akhtar and Gasser 2007). Several genes in budding yeast, such as *GAL1* and *INO1*, have been found to move to the nuclear periphery on activation and remain there for multiple mitotic cell cycles even after repression. It has been argued that this localization contributes to “transcriptional memory,” which is the observation that recently expressed genes are more rapidly reactivated than genes that have experienced long-term repression (Brickner 2009). The molecular mechanisms responsible for transcriptional memory are currently controversial (Zacharioudakis et al. 2007; Halley et al. 2010; Kundu and Peterson 2010). Nevertheless, H2A.Z appears to have a role in maintaining the localization of recently repressed genes at the nuclear periphery for multiple mitotic generations. In the case of *INO1*, this pathway requires a cis-acting DNA element in the *INO1* promoter and interaction with the nuclear pore protein NUP100 to direct the loading of H2A.Z (Light et al. 2010). If the H2A.Z gene is deleted, recently repressed *INO1* fails to remain at the nuclear periphery and becomes nucleoplasmic (Brickner 2009).

H2A.Z is also linked to the epigenetic specification of cell fates in mammalian stem cells (Creyghton et al. 2008;
see Reik and Surani 2014 for more on stem cell epigenetics). Whole genome-mapping studies in mouse embryonic stem cells revealed that H2A.Z preferentially occupies the promoters of genes that are poised to direct development and differentiation when activated (Ku et al. 2012; Li et al. 2012; Hu et al. 2013). Indeed, the distribution of H2A.Z is coincident with that of Suz12, a component of the Polycomb group complexes, which have primary roles in development (see Grossniklaus and Paro 2014 for a detailed discussion of Polycomb group proteins). Inhibiting H2A.Z expression results in increased and more stable nucleosome occupancy at regulatory regions, decreased methylation of H3K4 and H3K27 at promoters and enhancers, the derepression of developmental target genes, and aberrant differentiation of embryoid bodies. These results are consistent with a model in which H2A.Z variant nucleosomes impart a dynamic instability to chromatin structure increasing access to chromatin-modification factors such as mixed lineage leukemia and Polycomb repressive complex 2 (PRC2), and to lineage-specific factors such as the transcription factor FoxA2 for endoderm/hepatic differentiation (Li et al. 2012) or RARo for neuronal development (Hu et al. 2013).

In theory, stem cells are at a higher risk of mutation because of their continued proliferation during the life of the organism. Cairns first proposed that stem cells might protect against this risk by ensuring the inheritance of the “oldest” DNA strands exclusively to the stem cell daughter as opposed to the differentiating daughter cell (Cairns 1975). There is experimental support for this “immortal-strand model” in a number of systems in which asymmetric self-renewal can be experimentally manipulated. Interestingly, histone H2A.Z expression is a biomarker for this immortal-strand inheritance (Huh and Sherley 2011). First, H2A.Z mRNA is down-regulated in the differentiating sister cell during asymmetric cell division. This observation is consistent with the fact that the H2A.Z gene is a target of Oct4 regulation and is down-regulated following differentiation of both mouse and human stem cell populations (Du et al. 2001; Shaw et al. 2009; Huh and Sherley 2011). More striking, however, is the apparent asymmetric distribution of H2A.Z entirely to the immortal DNA strands segregating to the stem cell sister, as detected by immunofluorescence (Huh and Sherley 2011). This turns out to be because of the “cloaking” of the H2A.Z histones on the mortal-strand chromatids inherited by the differentiating sister cell. That is, H2A.Z is actually present on both sets of chromosomes, but it is only accessible to detection by the anti-H2A.Z antibody on the set segregating to the stem cell sister. Interestingly, mild acid treatment before immunostaining removes whatever is blocking detection and reveals H2A.Z on both sets of chromosomes. The nature of the cloaking mechanism is currently unknown, but these observations place H2A.Z quite proximal to the mechanism of immortal-strand inheritance.

13 OTHER H2A VARIANTS DIFFERENTIATE CHROMATIN, BUT THEIR FUNCTIONS ARE AS YET UNKNOWN

Further diversification of H2A has occurred in vertebrates. In mammals, macroH2A and H2A.B represent unique lineages that appear to play roles in the epigenetic phenomena of dosage compensation (discussed in detail in Brockdorff and Turner 2014). macroH2A is so-called because, in addition to the histone-fold domain and amino- and carboxy-terminal tails, it contains a more than 200-amino-acid carboxy-terminal globular domain (Ladurner 2003). macroH2A is enriched in discrete regions of the facultatively inactive X chromosome of human females that alternate with regions of constitutive heterochromatin (Fig. 11A) (Chadwick and Willard 2004). However, the role of macroH2A must be more diverse, insofar as macroH2A orthologs are present in many nonmammalian clades, and appear to be ancestral in the animal kingdom (Talbert and Henikoff 2010). In vitro, macroH2A reduces transcription factor access and excludes histone H1. The macro domain itself binds to the enzyme PolyADP ribose polymerase 1 and inhibits its activity in vitro (Nusinow et al. 2007), although exactly what is occurring in vivo to mediate gene repression is unclear.

In contrast to macroH2A, H2A.B seems to be undetectable on the Barr body, but otherwise ubiquitous throughout the nucleus (Fig. 11B) (Chadwick and Willard 2001).

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.)
The short wrap of DNA around H2A.B nucleosomes (Bao et al. 2004) and its enrichment at TSSs (Soboleva et al. 2012) is consistent with it playing a role in facilitating transcription. H2A.B and its closest relative, H2A.L (Govin et al. 2007), are rapidly evolving relative to other H2A variant classes, perhaps related to the fact that they are both testes-specific variants (Talbert and Henikoff 2010). The roles that H2A.B and H2A.L play in mammalian male germ cell development remain to be elucidated.

14 MANY HISTONES HAVE EVOLVED TO MORE TIGHTLY PACKAGE DNA

When it is no longer necessary to gain access to DNA for replication and transcription, chromatin typically becomes further condensed, and this often involves replacement of canonical histones. This is obviously the case for sperm, and in some lineages, histone paralogs have evolved specialized packaging roles. For example, sea urchin sperm contains H1 and H2B variants with repeated tail motifs that bind to the minor grooves of DNA (Talbert and Henikoff 2010), presumably an adaptation to tightly package chromosomes for inclusion into sperm heads. A similar adaptation is found in pollen-specific H2A variants in flowering plants. In vertebrates, a sperm-specific H2B specialized histone variant is found in mammalian testes, including an H2B paralog (subH2Bv) that localizes to the acrosome and a testes-specific H3 variant (Witt et al. 1996).

The replacement of histones during sperm maturation by protamines and other proteins provides a potential means of erasing epigenetic information in the male germ line. However, evidence for transgenerational inheritance (Rakyan and Whitelaw 2003; discussed in more detail in Blewitt and Whitelaw 2013), especially in animals that lack DNA methylation, raises the possibility that a subset of nucleosomal histones survive this transition and transmit epigenetic information. As already pointed out, this is just what occurs for CENP-A at centromeres (Palmer et al. 1990), and it is possible that a small fraction of other variants, such as H3.3, remain with sperm for epigenetic inheritance of gene-expression information. Although our understanding of the process that replaces histones during sperm development is rudimentary, we expect that much more can be learned by understanding how CENP-A survives this transition.

Increased compaction also occurs in somatic cells that have finished dividing and undergo differentiation. In some cases, compaction involves quantitative and qualitative changes in linker histones. The stoichiometry of histone H1 relative to nucleosomes determines the average spacing within nucleosome arrays in vivo (Fan et al. 2003). In addition, the presence of H1 in chromatin promotes higher-order chromatin structure that generally inhibits transcription (Wolffe 1992). Linker histones are much more mobile than core histones in vivo. Residence times for H2A and H2B are hours in length, and cannot even be measured for H3 and H4, whereas the residence time of H1 is a few minutes (Phair et al. 2004). As a result, the incorporation of variant linker histones is unlikely to differentiate chromatin in a heritable manner. Rather, the role of H1 variants is thought to change the bulk properties of chromatin that can affect overall compaction (Wolffe 1992).

H1 variants share with core histones a distinction between RC and RI forms (Marzluff et al. 2002). RC variant forms of H1 appear to be interchangeable with one another, based on the fact that knockout mice lacking one or two of the five RC H1 variants are phenotypically normal (Fan et al. 2003). The functional interchangeability of histone variants is implicit in the “star” phylogeny of the H1 family, in which there is little if any evidence for evolutionary conservation of H1 branches in the tree (Fig. 6C). Rather, the greater divergence of H1 relative to core histone variants is likely to reflect weaker structural constraints, and the existence of multiple H1 genes in many lineages might represent an adaptation for regulating levels of linker histone. For example, in birds, the H1.0 linker histone variant (previously referred to as H5) is deposited during erythrocyte maturation, which accompanies extreme compaction of the nucleus. The mammalian H1.0 ortholog (previously referred to as H1*) is deposited at high levels in nondividing cells. Overexpression of H1.0 renders chromatin less accessible to nucleases than similar overexpression of a canonical form. The natural accumulation of H1.0 in nondividing cells might be a general mechanism for chromatin compaction as cells become quiescent.

15 HISTONE VARIANTS AND HUMAN DISEASE

The diverse roles of histone variants in basic epigenetic processes leads to the expectation that their loss or misexpression can result in disease, and indeed some examples of this have recently come to light. For example, ATRX, which is the DNA translocase component of the ATRX-Daxx-H3.3 pathway (Fig. 8), was originally identified as causing α-thalassemia mental retardation on the X (ATRX) syndrome. In this syndrome, loss of ATRX results in defects at the CpG island of the α-globin gene promoter, and can now be understood as resulting from frequent nucleosome loss at CpG islands (Law et al. 2010). Importantly, mutations causing the ATRX syndrome are most frequently found within the ADD histone tail-binding domain (Eustermann et al. 2011), strongly implicating H3/H3.3 tail interactions as being key causal factors in the syndrome. The study of ATRX syndrome has also led to the realization
that another histone variant, macroH2A, is also depleted from specific sites, including the α-globin CpG island (Rattanakumar et al. 2012).

A remarkable relationship has recently emerged between the ATRX-Daxx-H3.3 nucleosome assembly process and cancer that provides important insights into both the process itself and oncogenesis. Genomic sequencing of human pancreatic neuroendocrine tumors (PanNETs) revealed that ∼40% harbored likely loss-of-function mutations in either Daxx or ATRX (Jiao et al. 2011). Notably, all such tumors displayed a distinct phenotype called “ALT” (alternative lengthening of telomeres) in which telomeres show massive lengthening without induction of telomerase (see Fig. 1 of Liu et al. 2014; Heaphy et al. 2011a). Based on examination of more than 6000 human tumors, 3.73% showed ALT occurring in cancers of all different types (Heaphy et al. 2011b). ATRX or Daxx are mutated in the vast majority of cases of ALT examined by sequencing. Although the precise molecular basis for telomere lengthening in ALT is not well understood, lengthening is likely initiated by recombinational strand invasion between telomeres, which suggests that failure to replace a nucleosome at a simple sequence array can result in DNA breakage and strand invasion. In the case of PanNETs, ALT predicts a more favorable outcome than telomerase induction, as if the ALT pathway is less efficient than the telomerase pathway in preventing the senescence that occurs when telomeres erode (Jiao et al. 2011).

Sequencing of tumors has also revealed the very surprising presence of specific mutations in H3.3 itself in a large fraction of pediatric glioblastomas (the topic of Liu et al. 2014; also Schwartzzentuber et al. 2012; Wu et al. 2012). One of the two H3.3 genes in these tumors is found to encode either K27M or G34R/V, and is most likely to be a gain-of-function mutation that drives accelerated tumorigenesis (see Fig. 1 of Liu et al. 2014). G34R/V is associated with ATRX loss and probably enhances ALT, whereas K27M can occur on an H3 as well, so would appear to be independent of the ATRX-Daxx-H3.3 pathway. Rather, K27 is a key substrate for methylation and acetylation involved in Polycomb silencing, and the presence of a nonmodifiable residue at position 27 of the H3/H3.3 tail causes gain-of-function inhibition of methylation by the EZH2 subunit of PRC2 (Lewis et al. 2013). It would then appear that the resulting global reduction in H3K27 trimethylation contributes to this aggressive pediatric tumor expression.

Misregulation of histone variant expression has also been implicated in cancer. In malignant melanoma cells, macroH2A levels are sharply reduced and restoration of macroH2A levels suppresses metastasis (Kapoor et al. 2010). Reduced levels of macroH2A result in up-regulation of many genes, including the CDK8 oncogene. macroH2A is also reduced in more rapidly proliferating lung cancers, and is elevated in cells undergoing senescence, which suggests that reduced senescence of tumor cells is a mechanism whereby loss of macroH2A contributes to oncogenesis (Sporn et al. 2009). It is possible that other defects in the expression of histone variants and in pathways that deposit them can contribute to cancer and other human diseases, but there is as yet insufficient evidence of causality as opposed to mere correlation. For example, excessively high levels of H2A.Z are found to correlate with poor outcomes in estrogen-positive breast cancer (Hua et al. 2008). In addition, CENP-A is found to be overexpressed and ectopically localized in a variety of cancers, suggesting that transient or permanent neocentromere formation might result in some of the aneuploidies that are hallmarks of cancer cells (Dalal 2009). We anticipate that as genome-wide technologies improve, other possible roles of histone variants incorporation pathways in human diseases will be uncovered.

16 CONCLUSIONS AND FUTURE RESEARCH

Histone variants provide the most fundamental level of differentiation of chromatin, and alternative mechanisms for depositing different variants can potentially establish and maintain epigenetic states. Histones H2A, H2B, H3, and H4 occupy distinct positions in the core particle as a result of an evolutionary process that began before the last common ancestor of eukaryotes. Key evolutionary innovations remain uncertain, including the emergence of an octamer from an ancestral tetramer, and we look forward to the sequencing of more archaeal and primitive eukaryotic genomes that might provide missing links. Subsequent elaborations of the four core histones into distinct variants have provided the basis for epigenetic processes, including development and chromosome segregation. For a full understanding of epigenetic inheritance, we need a better understanding of the processes that incorporate variants by replacing canonical histones. An important recent development is the initial characterization of RI assembly pathways dedicated to particular variants.

Centromeres are the most conspicuous examples of profoundly different chromatin that is attributable to special properties of a histone variant. Although it is clear that cenH3-containing nucleosomes form the foundation of the centromere, just how they are deposited in the same place every cell generation without any hint of sequence specificity is a major challenge for future research.

It is becoming evident that histone variants are also involved in epigenetic properties of active genes. Both H3.3 and H2A.Z are enriched at transcriptionally active loci, and understanding the assembly processes that are
responsible for their enrichment is an exciting area of current research. The dynamic behavior of chromatin leads to the realization that transcription, chromatin remodeling, and histone modification might be coupled to nucleosome assembly and disassembly. The study of dynamic processes coupled to histone turnover is only at an early stage, and we look forward to technological advances in molecular biology, cytogenetics, biochemistry, and structural biology that can be harnessed to better understand the dynamic nature of chromatin.

In addition to these universal processes, histone variants are also involved in particular epigenetic phenomena. In the case of the mammalian X chromosome, three different H2A variants, phospho-H2A.X, macroH2A, and H2A.B have been recruited to participate in silencing or activation of genes for purposes of germline inactivation or dosage compensation. Understanding the function of these variants in epigenetic processes remains a major challenge for the future.

The availability of the first high-resolution structure of the nucleosome core particle (Luger et al. 1997) was a seminal advance in elucidating the properties of chromatin. By elaborating this basic structure in a way that has biological consequences, histone variants provide an opportunity to deepen our understanding of how these fascinating architectural proteins have evolved to play diverse roles in epigenetic processes.

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WWW RESOURCES

http://www.ebi.ac.uk/Tools/phylogeny EBI server
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